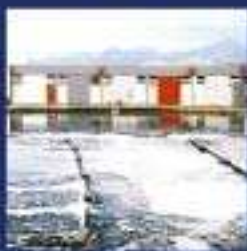
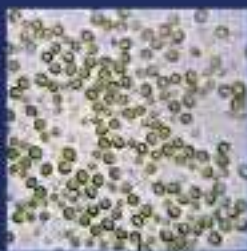


The Handbook of

Water and Wastewater Microbiology



Edited by Duncan Mara and Nigel Horan



Handbook of Water and Wastewater Microbiology

Edited by

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and

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Preface

Some 2300 years ago Hippocrates wrote: ‘My other topic is water, and I now wish to give an account both about waters that cause disease and about those that are healthy, and what bad things arise from water and what good things. For water contributes very much to health.’ So our appreciation of a relationship between the water we use and our health has been with us for a very long time. Hippocrates was unlikely to have been the first person to realise the existence of this relationship and we have probably known since our species evolved that water of adequate quantity and quality is essential for our survival and our health.

We now know (and have known for just over 100 years) that water quality is governed by (but, of course, not only by) microorganisms – the viruses, bacteria and parasites that can infect us and may (and very often do) make us ill. Microorganisms are also central to wastewater treatment and the reuse of treated wastewaters – we exploit them to treat our wastes biologically (actually, microbiologically), and we must ensure that pathogenic microorganisms are removed in the treatment processes to a level at which they do not cause any excess disease resulting from wastewater use in agriculture or aquaculture.

Water disinfection, usually with chlorine, has been practised in many parts of the world (but regrettably not all) for over 100 years. Water chlorination is a very efficient process: it kills bacteria very quickly (but viruses more slowly, and protozoa such as *Giardia* and *Cryptosporidium* hardly at all). Faecal bacterial numbers, in particular, are reduced to zero, and thus early water engineers judged the quality of chlorinated water supplies quite simply on

whether faecal indicator bacteria – principally coliform bacteria – were present in the disinfected water or not. Zero coliforms, and zero faecal coliforms, quickly became *the* microbiological goal of drinking water quality. No-one would really question the general sense of this goal – chlorinate your water and you get zero coliforms per 100 ml, so everything’s OK. End of story.

Life is rarely this simple, and water and wastewater microbiology is no exception. Emerging water-borne pathogens (*Cryptosporidium*, for example) require us to have a deeper understanding of water microbiology. Optimizing (really, maximizing) microbiological wastewater treatment also requires a knowledge of microbiology greater than that possessed by many design engineers. Structural engineers have a pretty good understanding of concrete, for example – so why shouldn’t those who design activated sludge plants or waste stabilization ponds have an equal appreciation of the microorganisms whose activities are essential to the treatment process they are designing?

The purpose of this Handbook is to provide an introduction to modern water and wastewater microbiology, especially for water and wastewater engineers. The study of water and wastewater microbiology is very rewarding: better water treatment, better wastewater treatment, safer wastewater reuse, and thus healthier people – in all parts of our world.

Duncan Mara
Nigel Horan

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Part I Basic Microbiology

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Microbial nutrition and basic metabolism

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1 INTRODUCTION

In nature, of all living organisms, microorganisms are the most versatile and diversified with regard to their nutritional requirements. For example, microorganisms, such as bacteria, can be found that represent the entire spectrum of nutritional types. Some microbes have an unconditional need for preformed complex organic compounds while others can thrive with just a few inorganic substances as their sole nutritional requirements. Most microorganisms fall between these two extremes.

Although great variation is found in the specific requirements for growth of the diverse species of microorganisms, in general, the nature and functions of growth substances are common for all cells. In part, this is because the chemical composition of microbial cells is more or less similar. For instance, all microbes require a carbon source because carbon is a component of protoplasm; nitrogen is a component of many major macromolecules, such as protein and nucleic acids. Over 95% of a cell's dry weight is made up of a few major elements, such as C, O, H, S, P, K, Ca, Mg, and Fe. All these substances must be put together by biosynthesis to form cellular material. However, not all growth substances are incorporated into cell material. Some are used instead as energy sources. For example, carbon compounds are frequently used as energy sources by many microorganisms; inorganic sulphur compounds are used in energy metabolism by some microbes.

Assimilated nutrient substances need to be metabolized. *Metabolism* refers to the totality of organized biochemical activities carried out by an organism. Such activities, usually catalysed by enzymes, are of two kinds: those involved in generating energy; and those involved in using energy. Some microorganisms use nutrients absorbed by the cell as the source of energy in a process called *catabolism*, which is the reverse of biosynthesis (*anabolism*). In catabolism, large molecules are degraded in sequential stepwise reactions by enzymes and a portion of the energy released is trapped in the form of chemical energy. Other microorganisms derive their energy from the trapping of light and also convert it into chemical energy. This chemical energy is then harnessed to do work for the cell. A microorganism must perform many different types of work, such as synthesis of physical parts of the cell, repair or maintenance of cell components, and growth. Thus metabolism may be viewed as the coupling of energy generation and energy utilization. The kinds of nutrients and how they are assimilated and fed into the various metabolic pathways for energy production and utilization in microorganisms form the subject matter of this chapter.

2 CHEMICAL ELEMENTS AS NUTRIENTS

It may be generalized that there are three essential nutritional needs of a cell: water,

energy, and chemical compounds, which may be used as building blocks. Most microorganisms, except for the phagocytic protozoa, have an absorptive type of nutrition. Thus chemical energy sources and chemical compounds must be dissolved in water. Water carries the solutes by transport mechanisms into the cell. Within the cell, water is the solvent in which the cell's biochemical reactions occur. It is also the medium for elimination of soluble waste substances from the cell. It is not surprising then that 70–80% of the microbial cell is water and that it constitutes the major portion of a cell's weight.

Required nutritional chemical compounds include chemical elements. These elements are necessary for both the synthesis of cell material and the normal functioning of cellular components such as enzymes. In order to grow, microorganisms must have proper essential chemical elements. The main chemical elements for cell growth include C, N, H, O, S and P – the same elements that form the chemical composition of the cells.

2.1 Macroelements

The major elements (macroelements), like C, H, N, O, S and P, are used in large amounts by microorganisms. The minor elements (microelements), like K, Ca, Mg, Na and Fe, are used in smaller quantities, while the trace elements, such as Mn, Zn, Co, Mo, Ni, Cu, are used in relatively very much smaller amounts. The amount of an element used by a microbe does not correlate with its relative importance; even one used in a trace amount may be essential to the growth or life of a microbial cell.

2.1.1 Carbon

Carbon is one of the most important chemical elements required for microbial growth. Fifty per cent of the dry weight of any cell is carbon; thus all organisms require carbon in some form. Carbon forms the backbone of three major classes of organic nutrients: carbohydrates, lipids and proteins. Such compounds provide energy for cell growth and serve as building blocks of cell material.

Microorganisms that use organic compounds as their major carbon source are called *heterotrophs*. Heterotrophs obtain such organic molecules by absorbing them as solutes from the environment. Some phagotrophic heterotrophs obtain organic molecules by ingestion of other organisms. Microorganisms that use carbon dioxide (the most oxidized form of carbon) as their major or even sole source of carbon are called *autotrophs*. They can live exclusively on relatively simple inorganic molecules and ions absorbed from the aqueous and gaseous environment.

2.1.2 Nitrogen

All organisms require nitrogen in some form. It is an essential part of amino acids that comprise cell proteins. Nitrogen is needed for the synthesis of purine and pyrimidine rings that form nucleic acids, some carbohydrates and lipids, enzyme cofactors, murein and chitin. Many prokaryotes use inorganic nitrogen compounds such as nitrates, nitrites, or ammonium salts. Unlike eukaryotic cells, some bacteria (like the free-living *Azotobacter* and the symbiotic *Rhizobium* of legume plants) and some archaeons (like the methanogens *Methanococcus* and *Methanobacterium*) can use atmospheric or gaseous nitrogen for cell synthesis by a physiological process called *nitrogen fixation*. Some microbes require organic nitrogen compounds such as amino acids or peptides. Some microorganisms use nitrate as an alternative electron acceptor in electron transport.

2.1.3 Hydrogen, oxygen, sulphur and phosphorus

Other elements essential to the nutrition of microorganisms are hydrogen, oxygen, sulphur and phosphorus. Hydrogen and oxygen are components of many organic compounds. Because of this, the requirements for carbon, hydrogen, and oxygen often are satisfied together by the availability of organic compounds. Free oxygen is toxic to most strict anaerobic bacteria and some archaeons, although aerobic microorganisms use oxygen as a terminal electron acceptor in aerobic respiration. Sulphur is needed for the biosynthesis of the amino acids cysteine, cystine,

homocysteine, cystathione and methionine, as well as the vitamins biotin and thiamine. Phosphorus is essential for the synthesis of nucleic acids and adenosine triphosphate. It is a component of teichoic acids and teichuronic acids in the cell walls of Gram-positive bacteria as well as a component of various membrane phospholipids. Reduced forms of sulphur may serve as sources of energy for *chemotrophs* or as sources of reducing power for *phototrophs*. Sulphate may serve as a terminal electron acceptor in electron transport.

2.2 Microelements and trace elements

Many other essential elements, the microelements and trace elements, are required in smaller amounts than the macroelements by microorganisms in their nutrition. Some of their functions in supporting the growth of microorganisms are summarized in Table 1.1.

For example, sodium is required by the permease that transports the sugar melibiose into the cells of the colon bacterium *Escherichia coli*. Sodium is required by marine microorganisms for maintaining cell integrity and growth. Some 'salt-loving' prokaryotes, the red extreme halophiles, cannot grow with less than 15% sodium chloride in their environment. Essential elements are often required as cofactors for enzymes. Because iron is a key component of the cytochromes and electron-carrying iron-sulphur proteins, it plays a key role in cellular respiration. However, most inorganic iron salts are highly insoluble in

water. Thus, many microbes must produce specific iron-binding agents, called siderophores, in order to utilize this element. Siderophores are chelating agents that solubilize iron salts and transport iron into the cell. Many enzymes, including some involved in protein synthesis, specifically require potassium. Magnesium functions to stabilize ribosomes, cell membranes and nucleic acids and is needed for the activity of many enzymes.

Trace elements, needed in extremely small amounts for nutrition by microorganisms, include manganese, molybdenum, cobalt, zinc, and copper. For instance, molybdenum is required by nitrogenase, the enzyme that converts atmospheric nitrogen to ammonia during nitrogen fixation. Manganese aids many enzymes to catalyse the transfer of phosphate groups. Cobalt is a component of vitamin B₁₂ and its coenzyme derivatives.

3 NUTRITIONAL TYPES OF MICROBES

Microbes can be grouped nutritionally on the basis of how they satisfy their requirements for carbon, energy, and electrons or hydrogen. Indeed, the specific nutritional requirements of microorganisms are used to distinguish one microbe from another for taxonomic purposes.

Microorganisms may be grouped on the basis of their energy sources. Two sources of energy are available to microorganisms. Microbes that oxidize chemical compounds (either organic or inorganic) for energy are

TABLE 1.1 Functions of some microelements and trace elements in the nutrition of microorganisms

Element	Major functions in some microorganisms
Sodium	Enzyme activator. Transport across membranes. Maintenance of cell integrity. Facilitates growth. Salt form of some required organic acids
Potassium	Cofactor for enzymes. Maintenance of osmotic balance
Iron	Component of cytochromes, haem-containing enzymes, electron transport compounds and proteins. Energy source
Magnesium	Enzyme activator, particularly for kinase reactions. Component of chlorophyll. Stabilizes ribosomes, cell membranes, and nucleic acids
Calcium	Enzyme activator, particularly for protein kinases. Component of dipicolinic acid in bacterial endospores
Cobalt	Component of vitamin B ₁₂ and its coenzyme derivatives
Manganese	Enzyme activator, particularly for enzymes transferring phosphate groups
Molybdenum	Enzyme activator for nitrogen fixation

called *chemotrophs*; those that use light as their energy sources are called *phototrophs*. A combination of these terms with those employed in describing carbon utilization results in the following nutritional types:

1. *Chemoautotrophs*: microbes that oxidize inorganic chemical substances as sources of energy and carbon dioxide as the main source of carbon.
2. *Chemoheterotrophs*: microbes that use organic chemical substances as sources of energy and organic compounds as the main source of carbon.
3. *Photoautotrophs*: microbes that use light as a source of energy and carbon dioxide as the main source of carbon.
4. *Photoheterotrophs*: microbes that use light as a source of energy and organic compounds as the main source of carbon.

Microorganisms also have only two sources of hydrogen atoms or electrons. Those that use reduced inorganic substances as their electron source are called *lithotrophs*. Those microbes that obtain electrons or hydrogen atoms (each hydrogen atom has one electron) from organic compounds are called *organotrophs*.

A combination of the above terms describes four nutritional types of microorganisms:

1. *Photolithotrophic autotrophy*
2. *Photo-organotrophic heterotrophy*
3. *Chemolithotrophic autotrophy*
4. *Chemo-organotrophic heterotrophy*.

The characteristics of these types with representative microorganisms as well as other organisms are shown in Table 1.2.

Photolithotrophic autotrophs are also called *photoautotrophs*. The cyanobacteria, algae and green plants use light energy and carbon dioxide as their carbon source but they employ water as the electron donor and release oxygen in the process. The purple and green sulphur bacteria use inorganic compounds as electron donors (e.g., H_2S , S^0) and do not produce oxygen in the process. Thus they are described as *anoxygenic*. Chemo-organotrophic heterotrophs are also called *chemoheterotrophs*. They use organic compounds for energy, carbon and electrons/hydrogen. The same organic nutrient compound often satisfies all these requirements. Animals, most bacteria, fungi, and protozoa are chemoheterotrophs. Photo-organotrophic heterotrophs are also called in short *photoheterotrophs*. The purple and green non-sulphur bacteria are photoheterotrophs and use radiant energy and organic compounds as their electron/hydrogen and carbon donors. These common microorganisms, found in polluted lakes and streams, can also grow as photoautotrophs with molecular hydrogen as electron donor. The chemolithotrophic autotrophs are also called *chemoautotrophs* in brief. They include the nitrifying, hydrogen, iron and sulphur bacteria. They oxidize reduced inorganic compounds, such as nitrogen, iron or sulphur molecules, to derive both energy and electrons/hydrogen. They use carbon dioxide as their carbon source. A few of them, however,

TABLE 1.2 Nutritional types of microbes and other organisms

Nutritional type	Energy source	Electron or hydrogen source	Carbon source	Examples of organisms
Photolithotrophic autotrophy	Light	Inorganic compounds, water	Carbon dioxide	Purple and green sulphur bacteria; algae; plants; cyanobacteria
Photo-organotrophic heterotrophy	Light	Organic compounds	Organic compounds	Purple and green non-sulphur bacteria
Chemolithotrophic autotrophy	Inorganic compounds	Inorganic compounds	Carbon dioxide	Nitrifying, hydrogen, iron, and sulphur bacteria
Chemo-organotrophic heterotrophy	Organic compounds	Organic compounds	Organic compounds	Most bacteria, fungi, protozoa, and animals

can make use of carbon from organic sources and thus become heterotrophic. Such bacteria that use inorganic energy sources and carbon dioxide, or sometimes organic compounds, as carbon sources can be called *mixotrophic*, because they combine autotrophic and heterotrophic processes. Chemotrophs are important in the transformations of the elements, such as the conversion of ammonia to nitrate and sulphur to sulphate, that continually occur in nature.

Even though a particular species of microorganism usually belongs to only one of the four nutritional types, some show great metabolic flexibility and can alter their nutritional type in response to environmental change. For example, many purple non-sulphur bacteria are photoheterotrophs in the absence of oxygen but become chemoheterotrophs in the presence of oxygen. When oxygen is low, photosynthesis and oxidative metabolism can function simultaneously. This affords a survival advantage to the bacteria when there is a change in environmental conditions.

The specific nutritional requirements of bacteria are used extensively for taxonomic purposes. Specific identification tests have been designed for particular groups of bacteria, such as the Gram-negative intestinal bacilli, to determine the nature of water pollution.

4 GROWTH FACTORS

Some microorganisms have good synthetic capability and thus can grow in a medium containing just a few dissolved salts. The simpler the cultural medium to support growth of a species of microbe, the more complex or advanced is the microbe's nutritional synthetic capability. Thus, the photoautotrophs are the most complex in their nutritional physiology. With the addition of one organic compound, such as the addition of glucose, a glucose-salts medium can support the growth of many chemoheterotrophs; an example is the bacterial indicator of faecal contamination, *Escherichia coli*. However, many microorganisms lack one or more essential enzymes and therefore cannot

synthesise all their nutritional requirements. They must obtain these preformed or supplied in the environment or medium.

Organic compounds required in the nutrition of microorganisms, because they cannot be synthesized specifically, are called *growth factors*. The three major classes of growth factors are *amino acids*, *purines* and *pyrimidines*, and *vitamins*. Proteins are composed of about 20 amino acids. Some bacteria and archeons cannot synthesize one or more of these and require them preformed in the medium. For example, *Staphylococcus epidermidis*, the normal resident on the human skin, requires proline, arginine, valine, tryptophan, histidine and leucine in the medium before it can grow. Requirements for purines and pyrimidines, the nucleic acid bases, are common among the lactic acid bacteria. Vitamins are small organic compounds that make up all or part of the enzyme cofactors (non-protein catalytic portion of enzymes). Only very small, or catalytic, amounts suffice to support growth of the cells. Lactic acid bacteria, such as species of *Streptococcus*, *Lactobacillus* and *Leuconostoc*, are noted for their complex requirements of vitamins and hence many of these species are used for microbial assays of food and other substances. Vitamins most commonly required by microorganisms are thiamine (vitamin B₁), biotin, pyridoxine (vitamin B₆) and cyanocobalamin (vitamin B₁₂). The functions of some vitamins for the growth of microorganisms are summarized in Table 1.3.

5 ENERGY TRAPPING IN MICROORGANISMS

Like all living things, microorganisms require energy to live. The ability of a microorganism to maintain its life processes and to reproduce its own kind depends on its ability to trap energy and to use it to drive the endergonic reactions of the cell. Like all living forms, microbes trap or obtain energy in one of two ways: by *high-energy molecules* or by a *proton motive force* (*proton gradient*) across a cell membrane.

TABLE 1.3 Functions of some vitamins in the growth of microorganisms

<i>Vitamin</i>	<i>Function(s)</i>
Folic acid	One-carbon transfers; methyl donation
Biotin	Carboxyl transfer reactions; carbon dioxide fixation, β -decarboxylations; fatty acid biosynthesis
Cyanocobalamin (B ₁₂)	Carries methyl groups; synthesis of deoxyribose; molecular rearrangements
Lipoic acid	Transfer of acyl groups
Nicotinic acid (niacin)	Precursor of NAD ⁺ and NADP ⁺ ; electron transfer in oxidation–reduction reactions
Pantothenic acid	Precursor of coenzyme A; carries acyl groups
Riboflavin (B ₂)	Precursor of FMN, FAD in flavoproteins involved in electron transport; dehydrogenations
Thiamine (B ₁)	Aldehyde group transfer; decarboxylations
Pyridoxal-pyridoxamine group (B ₆)	Amino acid metabolism, e.g. transamination and deamination
Vitamin K group; quinones	Electron transport; synthesis of sphingolipids
Hydroxamates	Iron-binding compounds; solubilization of iron and transport into cell
Haem and related tetrapyrroles	Precursors of cytochromes

5.1 High-energy molecules

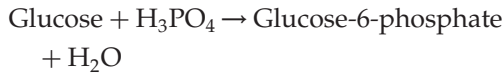
The energy liberated by an exergonic reaction can be used to drive an endergonic reaction if there is a reactant common to both reactions. This common reactant in the trapping of energy is called an energy-rich or energy-transfer compound. The energy-transfer compounds of greatest use to a cell are able to transfer large amounts of free energy and are called *high energy-transfer compounds* or *high-energy molecules*. Of these, adenosine triphosphate (ATP) is the most important. Such compounds are found in the cytosol or soluble part of the cell.

High-energy molecules are important because they drive biosynthesis in the cytoplasm, including the synthesis of nucleic acids, proteins, lipids and polysaccharides. They also are involved in the active transport of certain solutes into the cell. High-energy molecules, such as ATP, have bonds that have a high free energy of hydrolysis. In the case of ATP, a large amount of energy is needed to link the third phosphate group to adenosine diphosphate (ADP) because of the electrostatic repulsion of the negative charges on the other adjacent phosphate groups. Reactions in which the phosphate is removed from ATP will be favoured since the electrostatic repulsion is decreased as a result of hydrolysis. Thus, this energy is liberated if ATP is hydrolyzed back to ADP.

There is a reason why the phosphoryl group is a common chemical group that is transferred between molecules. The phosphorus in all phosphate groups carries a positive charge. This is because phosphorus forms double bonds (P=O) poorly so that the phosphorus–oxygen bond exists as the semi-polar bond P⁺–O[−]. The electrons in the bond are shifted toward the electron-attracting oxygen. The positively charged phosphorus is attacked by the electronegative oxygen in the hydroxyl of a water molecule in a hydrolytic reaction resulting in the release of inorganic phosphate. One can compare the tendency of different molecules to donate phosphoryl groups by comparing the free energy released when the acceptor is water, i.e. the free energy of hydrolysis. That is, a scale is used, where the standard nucleophile is the hydroxyl group of water, and the phosphoryl donors are all compared with respect to the tendency to donate the phosphoryl group to water. This gives a free energy change, or energy available to perform useful work and designated ΔG , per mole of substrate hydrolysed. The *standard free-energy change* under standard conditions (reactants and products at 1 M concentration and the reactions take place at 25°C at pH 7) is designated ΔG° .

The general role of ATP in providing energy to drive an endergonic reaction may be shown below.

Consider the following reaction:



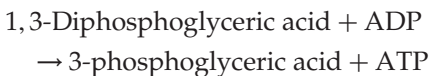
The ΔG° is +13.8 kJ/mol; it is an endergonic reaction and will not proceed spontaneously.

However, if ATP is provided as a reactant, the reaction becomes exergonic since the $\Delta G^\circ = -16.7$ kJ/mol (-30.5 (from ATP hydrolysis) + $13.8 = -16.7$ kJ/mol) and proceeds spontaneously:



The compound ADP is also sometimes used by cells as a high energy-transfer compound since its hydrolysis also liberates an equally large quantity of energy as ATP ($\Delta G^\circ = -30.5$ kJ/mol). However, adenosine monophosphate (AMP) is a low-energy molecule; its hydrolysis yields only a small amount of energy ($\Delta G^\circ = -8.4$ kJ/mol).

Table 1.4 lists some high energy-transfer compounds with their standard free-energy values upon hydrolysis. Each of them can transfer its energy of hydrolysis directly or indirectly to ATP synthesis, as in the following example:



The synthesis of high-energy-transfer compounds, such as ATP, involves phosphorylation. ATP is formed by phosphorylation of ADP, with energy for the phosphorylation being provided by an exergonic reaction.

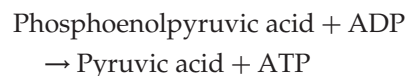
TABLE 1.4 Some high energy-transfer compounds with their standard free energy release upon hydrolysis

Compound	ΔG° , kJ/mol
Adenosine triphosphate (ATP)	-30.5
Guanosine triphosphate (GTP)	-30.5
Uridine triphosphate (UTP)	-30.5
Cytidine triphosphate (CTP)	-30.5
Acetyl phosphate	-42.3
1,3-Diphosphoglyceric acid	-49.4
Phosphoenolpyruvic acid (PEP)	-61.9

There are two general ways in which this phosphorylation of ADP can occur:

- *Substrate-level phosphorylation*, a reaction in which the phosphate group of a chemical compound is removed and directly added to ADP
- Phosphorylation by a membrane-bound enzyme called a proton-translocating ATPase, which uses the energy of an energy-trapping system called the proton motive force (described later).

In substrate-level phosphorylation, the rearrangement of atoms within chemical compounds derived from nutrients may result in a new compound that contains a high-energy phosphate bond. Such rearrangements can occur when cells dissipate nutrients. The phosphate group involved in the high-energy phosphate bond then can be transferred directly to ADP, forming ATP, which now contains the high-energy phosphate bond. (Note that bond energy is the energy required to break a bond, e.g. by hydrolysis, and it is equal to the energy released when the bond is formed. It is not the energy released when a bond is broken. For example, the P–O bond energy is about +413 kJ, while the hydrolysis energy is -35 kJ.) One example of substrate level phosphorylation is as follows:



It should be added that any chemical group that is electronegative (such as the hydroxyl groups in sugars) can attack the electropositive phosphorus resulting in phosphoryl group transfer if the proper enzyme is present to catalyse the reaction. In this manner, ATP can phosphorylate many different compounds. Enzymes that catalyse phosphoryl group transfer reactions are called *kinases*.

5.2 The proton motive force

The *chemiosmotic theory* was proposed by the biochemist Peter Mitchell in 1961. He received the Nobel Prize in 1978 for this proposal.

The theory described how organisms can trap energy by a means other than the synthesis of high-energy molecules. In brief, the chemiosmotic theory states that energy transducing membranes, such as bacterial cell membranes, mitochondrial membranes, and chloroplast membranes, pump protons across the membrane resulting in the generation of an electrochemical gradient of protons across the membrane. In the bacteria and the Archaea, the protons are pumped across the cytoplasmic membrane to the outside of the cell. In the non-phototrophic Eukarya, the cytoplasmic membrane is not involved; instead the protons are pumped outward across the inner membrane of a mitochondrion into the space between the inner and outer mitochondrial membrane. In the phototrophic Eukarya, the protons are pumped inward across the thylakoid membrane of the chloroplast. This gradient, also known as the *proton potential* or *proton motive force*, can be harnessed to do useful work when the protons return across the membrane through special proton conductors to the lower potential. Some membrane proton conductors are solute transporters, others synthesise ATP, and others are motors that drive bacterial flagella rotation. The proton potential also provides the energy for other membrane activities, such as gliding motility and reversed electron flow. The chemiosmotic theory brings together the principles of physics and thermodynamics to bear on the biological problem of membrane bioenergetics.

There are two components to the proton motive force:

1. The cell can pump or translocate protons (i.e. hydrogen ions, or H^+) across a membrane without also pumping a compensating anion such as OH^- or Cl^- at the same time. The protons are pumped out of the cell by exergonic driving reactions (energy-producing reactions), which are usually biochemical reactions of respiration, photosynthesis, or ATP hydrolysis. Once protons have been pumped across the membrane, they cannot pass back across to equalize the concentrations, because the membrane is impermeable to protons.
2. The protons that pass to the outside of the cell cannot pass back in because the membrane is impermeable to protons. The protons accumulate to the point where they decrease the external pH (higher acidity) relative to the pH (lower acidity) inside the cell. The concentration of protons may be 100 times greater on one side of the membrane than on the other (a 2-unit pH difference). This proton gradient, i.e. the greater concentration of protons on one side of the membrane and a lower concentration of protons on the opposite side, is referred to as a ΔpH .

The unequal distribution of protons and electrical charges across the membrane represents a form of potential energy called the proton motive force, or Δp . It is the resultant of both the $\Delta\Psi$ and the ΔpH and can be calculated at 30°C in terms of millivolts by the following relationship:

$$\Delta p = \Delta\Psi - 60\Delta pH$$

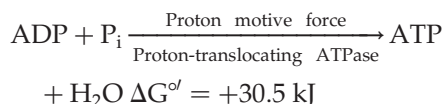
For example, in a bacterial cell, if the ΔpH is 2, and the $\Delta\psi$ is -160 mV (negative since the inside of a bacterial cell would be negatively charged with respect to the outside), the Δp will be $-160 + (-60)(2) = -160 - 120 = -280$ mV. Either $\Delta\psi$ or ΔpH alone can constitute the proton motive force; the existence of both factors is not necessary.

When the protons return to the inside of the cell, such as the bacterial cell, moving down the concentration gradient and towards the more negative side of the membrane, the membrane potential is dissipated, i.e. energy is given up and work can be done. Since the membrane is impermeable to protons, there are special conditions under which the protons can pass back across. To do this, the protons must carry out some type of work.

In bacteria, specific channels exist in the cytoplasmic membrane by which protons can pass back across the membrane, and the proton flow through these channels is harnessed by the cell to do various types of work, which includes synthesizing ATP from ADP, powering the rotation of bacterial flagella, active transport of nutrients into the cell, secretion of proteins out of the cell, and reversed electron transport.

Any agent that allows protons to pass freely across the cytoplasmic membrane will destroy the proton gradient and hence the proton motive force. *Uncoupling agents* carry protons freely across the membrane. One such agent is CCCP or carbonylcyanide-chloro-phenylhydrazine. In their presence, there is no accumulation of protons outside the cell, there is no proton motive force, and the bacterial cell dies.

One function of the proton motive force is to power the synthesis of ATP from ADP. This is catalysed by an enzyme complex called a proton-translocating ATPase, or simply ATPase synthase, in the following reaction:



The reaction catalysed by the ATPase is reversible. Not only can an influx of protons drive the synthesis of ATP but, in the absence of a proton motive force, the hydrolysis of ATP can drive the export of protons to create a proton motive force that can be used for energy-requiring activities, such as nutrient accumulation and motility.

The proton motive force is usually generated by one of the two main mechanisms: the *oxidation reactions in electron transport systems* (leading to ATP formation) and the *hydrolysis of ATP* (as just stated above). Other mechanisms exist but they are less common.

6 OXIDATION-REDUCTION SYSTEMS IN ELECTRON TRANSPORT

Generation of the proton motive force by the oxidation reactions in electron transport systems depends on the principle that oxidation

reactions liberate energy. *Oxidation* is the loss of electrons from an atom or molecule. It can also be the loss of hydrogen atoms from a molecule, since each hydrogen atom contains an electron in addition to its proton. The opposite of oxidation is *reduction*, i.e. the gain of electrons or hydrogen atoms. Unlike oxidation reactions, reduction reactions do not liberate energy but instead require energy in order to proceed. The reverse of any oxidation is a reduction and, it follows, that the reverse of any reduction is an oxidation. In an oxidation-reduction reaction, a pair of substances is involved: one is the oxidized form and the other the reduced form, e.g. Fe^{3+} and Fe^{2+} . Each pair of such substances is called an oxidation-reduction (O/R) system. One O/R system may tend to accept electrons from another system, i.e. the first system will be reduced while the second system will be oxidized. The capacity of an O/R system to be oxidized or reduced depends on the relative oxidizing power of each O/R system.

The oxidative power of any O/R system is called its E_h value. The 'h' means that it is measured electrically in relation to the H^+/H_2 system, which is the standard system, and is expressed in volts. The more positive the E_h of an O/R system, the greater the oxidizing ability of the system. Knowledge of the relation of each O/R system to the standard H^+/H_2 system permits the comparison of one system with another. To do this, the concentrations of the oxidized and reduced forms in an O/R system, as well as the pH and temperature, must be taken into account. These variables affect the E_h value of an O/R system. When this is done, a value called E_O , the *standard oxidation potential*, is obtained and used. It is the particular E_h value when O and R are at the same concentration ($[\text{O}] = [\text{R}]$), the temperature is 25°C , and the pH is 7. Under these particular conditions, any system can oxidize any other system having an E_O that is less positive but not more positive. These relationships are very important in recognizing the sequence in which biological oxidations occur, especially in an electron transport system, where the oxidized and reduced forms of the reactants are in approximately a 1:1 ratio.

When one O/R system oxidizes another, energy is released (the $\Delta G^{\circ'}$ value is negative). The amount of energy released or the standard free energy change, the $\Delta G^{\circ'}$ is directly proportional to the difference in $E_{O'}$ values of the two O/R systems:

$$\Delta G^{\circ'} = -nF\Delta E_{O'}$$

Here, n = the number of electrons transferred per molecule, usually 1 or 2; F is the Faraday (96 500 coulombs). Application of this equation to substances whose standard oxidation potentials are known, allows prediction of the direction of the reaction and the amount of energy released from it. Furthermore, the equation allows determination of whether or not the energy released from a particular oxidation is sufficient to allow formation of a molecule of ATP. The following example is instructive. Consider the oxidation of NADH by FAD. The standard oxidation potential ($E_{O'}$) value for NAD is -0.32 , while that for FAD is -0.22 . The process involves the transport of two electrons. Substituting these values into the above equation, the amount of energy released per mole of NADH oxidation is:

$$\begin{aligned}\Delta G^{\circ'} &= \frac{-2 \times 96\,500(-0.22 - [-0.32])}{4.18} \\ &= -4,617 \text{ cal} = -4.62 \text{ Kcal}\end{aligned}$$

where 4.18 is the conversion factor between coulomb-volts (Joules) and calories. In this example, 4.62 kcal of energy are released per mole of NADH oxidized, an amount insufficient to produce an ATP molecule. The formation of an ATP molecule requires 7.5–8 kcal per mole. It should be remembered, however, that the actual amount of energy released from oxidative processes in cells may be different from that obtained by the kind of calculation as above using values for compounds obtained under standard conditions. Such calculations should be used with caution.

7 ELECTRON TRANSPORT SYSTEMS

An *electron transport system* is a sequence of oxidation reactions in which electrons from a source of reducing power pass from one O/R

system to another. The source of reducing power may be as diverse as electrons derived from organic compound oxidation, oxidation of an inorganic ion, or radiant energy. In the sequence, each successive O/R system has a greater ability to gain electrons (i.e. has a greater $E_{O'}$ value) than the one preceding it. As the electrons flow through the series of oxidation reactions, much of the free energy of these reactions is trapped in the form of the proton motive force, which can then power the synthesis of ATP, a process called *oxidative phosphorylation*, as distinguished from substrate level phosphorylation. Electron transport systems are always associated with membranes. In prokaryotes they are in the cytoplasmic membrane; in eukaryotes they are in mitochondrial or chloroplast membranes.

Electron transport through the successive O/R systems is finally terminated in one of the two ways. In cyclic electron transport, present in *anoxygenic* (non-oxygen-forming) photosynthetic prokaryotes, the electron removed from the initial chlorophyll molecule is returned to the chlorophyll molecule from which it arose in a cyclic manner. In chemotrophic electron transport, oxidation of the last electron carrier in the chain is used to reduce a *terminal electron acceptor*. In aerobic microorganisms, the terminal electron acceptor is oxygen. The resulting compound is water, formed by the reduction of a half of a molecule of molecular oxygen with two protons and two electrons derived from the transport process. In some anaerobic microorganisms, organic compounds, like fumarate, may serve functions analogous to molecular oxygen resulting in reduced products, like succinate. In addition, some microorganisms that are facultative use nitrate or sulphate as terminal electron acceptors in the absence of oxygen, leading to the formation of reduced forms of nitrogen and sulphur.

7.1 Carriers in chemotrophic electron transport systems

A generalized scheme of the electron transport system in *chemotrophic* microorganisms is depicted in Fig. 1.1. In this system, the first

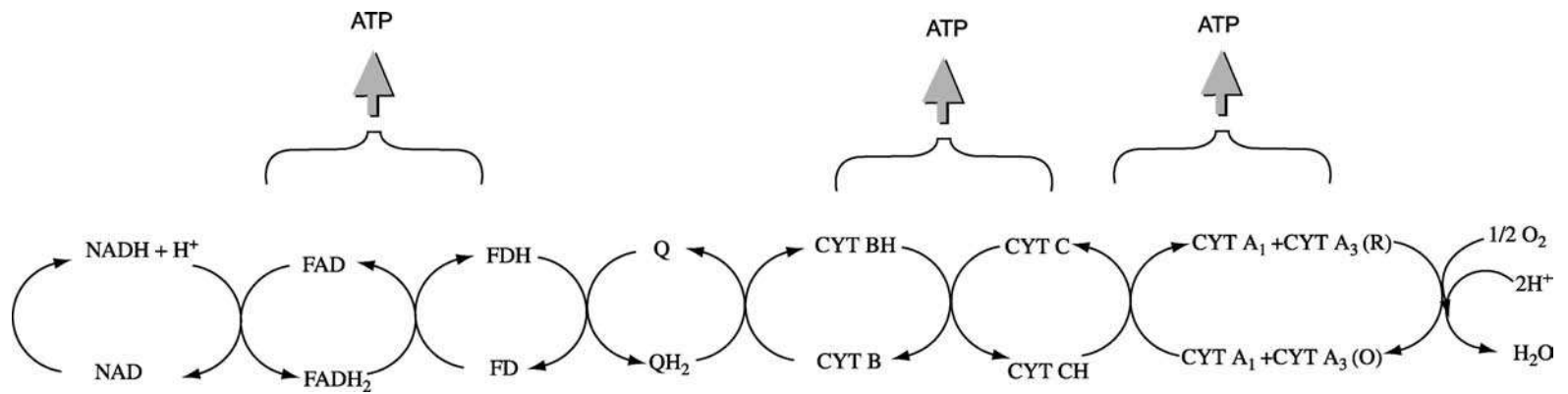
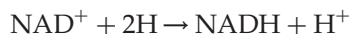


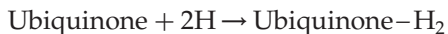
Fig.1.1 Generalized scheme of electron transport in chemotrophic microorganisms.

molecule in the electron chain is typically NAD or NADP. Enzymes, called dehydrogenases, remove electrons and hydrogen ions from substrates and many dehydrogenases use the pyridine nucleotides (NAD) nicotinamide adenine dinucleotide or NADP as their coenzyme. Some dehydrogenases are specific for NAD or NADP, whereas others are non-specific and show activity with either compound. NAD is composed of an ADP, a ribose phosphate and niacinamide. Its structure is shown in Fig. 1.2. NADP is similar to NAD but has an extra phosphate group. NAD can accept two electrons and one proton and thus exists in the reduced form:



Reduced forms of these pyridine nucleotides are normally reoxidized by transfer of their electrons to a molecule of a flavin-containing compound, typically a flavoprotein.

Flavoproteins contain either flavin adenine dinucleotide (FAD) or flavin mononucleotide (FMN) as prosthetic groups (see Fig. 1.2). The vitamin riboflavin is part of the structure of these prosthetic groups. FAD can accept two hydrogen atoms to exist in a reduced form, FADH_2 . Similarly FMN can be reduced to FMNH_2 . The flavoprotein is reoxidized by electron transfer to the quinines, ubiquinone and menaquinone. These are water-soluble compounds, not coenzymes, that move freely within the non-polar matrix of membranes and function as acceptors of two hydrogen atoms, e.g.



In some systems, this transfer is mediated by a non-haem-containing iron sulphur protein, most often a ferredoxin or a structurally similar compound. Transfer of electrons from reduced quinone compounds to a series of cytochromes is the next step in electron transport.

A diversity of cytochromes is found in electron transport systems. All of the cytochromes are haem proteins that differ in the nature of both their proteins and the side chains of their haem moieties. Cytochromes are enzymes with a prosthetic group that is a derivative of haem. This prosthetic group has a

single iron atom that is responsible for the oxidative or reductive properties of the enzyme (see Fig. 1.2). On the basis of differences in light absorption spectra, cytochromes can be divided into four main categories designated cytochromes a, b, c, and d. There is also cytochrome o, which belongs to the cytochrome b class. Each of these groups has a different function in a respiratory chain and can be further subdivided on the basis of minor differences in light absorption spectra, such as cytochromes c and c_1 or cytochromes a and a_3 . Each cytochrome type can exist in either an oxidized or reduced form, depending on the state of the iron atom contained in their structure. Since electrons are usually transferred in pairs, two cytochrome molecules are needed for each electron pair. Certain cytochrome complexes, such as cytochromes a and a_3 (called cytochrome aa_3) and cytochromes b and o (called cytochrome bo) are called cytochrome oxidases, because they can transfer electrons directly to O_2 to form H_2O .

Various electron transport systems may use slightly different chemical substances, but the functions of related substances are physiologically equivalent. It follows that the precise nature of the electron transport systems of prokaryotes differs substantially among microorganisms, but the carrier molecules involved in most electron systems are both structurally and functionally similar.

It will be recalled that energy is released as the electrons move along the carriers in the electron transport system. If the energy release is large, this can be coupled with the movement of H^+ across the membrane, generating a proton motive force resulting in the production of ATP. The number of sites where this can occur depends on the difference between the redox potential of the substrate and the final electron acceptor.

When $\text{NADH} + \text{H}^+$ is the electron donor and oxygen is the electron acceptor, proton movements associated with large free energy changes occur at three sites (see Fig. 1.1). The sites are at NADH-FAD, cytochrome bc_1 complex, and the terminal oxidase. In microorganisms with shorter electron transport systems or where electrons enter at other sites in the system

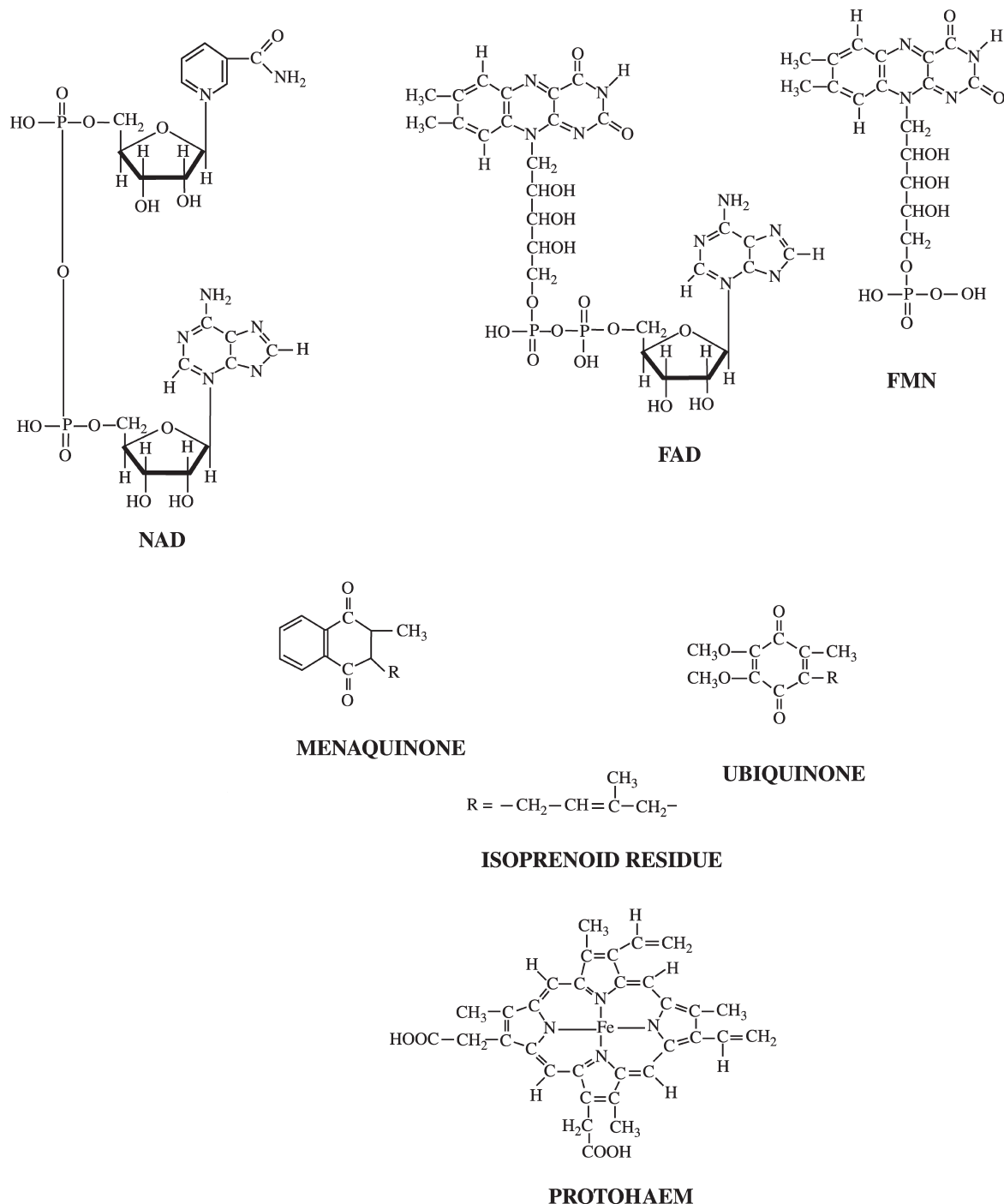


Fig.1.2 Structures of prosthetic groups and compounds in the electron transport system of chemotrophic microorganisms. The various quinones and menaquinones differ from each other by the number of R units attached to the carbonyl-containing rings. Protohaem, or haem B, is a prototype for all of the prosthetic groups of the cytochromes. Haems associated with cytochromes other than cytochromes of the *b*-type have side chains other than those shown for haem B.

or chain (having a more positive redox potential than that of the NADH/NAD⁺ couple), less ATP is produced.

In environments devoid of oxygen, many microorganisms can utilize electron transport mechanisms for ATP synthesis, but they use alternate terminal electron acceptors other than oxygen. This process is called *anaerobic respiration*. The alternate acceptors used include nitrate, sulphate, CO₂, and specific organic compounds. (Facultative microbes will use oxygen in aerobic respiration when it is available.) Microorganisms that use nitrate as an electron acceptor reduce nitrate to nitrite and ammonia by the activity of nitrate and nitrite reductase. Methanogens use CO₂ as the terminal electron acceptor and produce methane in the process. Some sulphur bacteria, like *Desulfovibrio*, reduce sulphate to sulphide when using the former as terminal electron acceptor.

Anaerobic respiration is much less energy-efficient than aerobic respiration. A large amount of substrate is required to produce sufficient ATP for the cell. This is because the pathways for this metabolic process yield less energy for the proton motive force than do pathways with oxygen as the terminal electron acceptor because the E_0' value of the terminal electron acceptor system is more negative than that of the 1/2O₂/H₂O system—i.e. it has less oxidizing power.

7.2 Carriers in phototrophic electron transport systems

In *phototrophic* electron transport, the original light-absorbing substance is one or more chlorophyll and bacteriochlorophyll molecules, in tandem with carotenoid pigments and phycobiliproteins. The chlorophylls differ both structurally and in their light-absorbing properties. All of these molecules have a large number of double bonds. This property is essential for their ability to transfer energy from one molecule to another. Briefly, light is absorbed by these pigments, which pass the

energy to the photosynthetic reaction centres, which then create the electron transport systems.

Thus, phototrophs have the amazing ability to use light energy to create an electron transport system that provides them with a proton motive force. They do not obtain the terminal electron acceptor for this electron transport system from the environment or the culture medium; instead they use their own oxidized chlorophyll or bacteriochlorophyll molecules.

There are two types of photosynthesis in microorganisms: the *anoxygenic* type does not form oxygen as a product of photosynthesis; the *oxygenic* type forms oxygen. The cyanobacteria carry out oxygenic photosynthesis. Photosynthesis in algae and green plants is similar to that of cyanobacteria. Green and purple bacteria differ from the cyanobacteria and carry out anoxygenic photosynthesis.

7.2.1 Oxygenic photosynthesis

Photosynthetic electron transport in the cyanobacteria is carried out by localized intracytoplasmic membranous organelles called thylakoids. Thylakoids are covered with phycobilisomes containing the light-harvesting pigments. Light-harvesting pigments can be divided into chlorophylls that absorb light in the red (>640 nm) and blue (<440 nm) regions of the light spectrum, and the carotenoids, phycobiliproteins, phycoerythrin and phycocyanin (termed accessory pigments because they absorb light at wavelengths where chlorophylls do not function efficiently (470–630 nm)). The pigments are arranged in a highly organized manner to maximize energy transfer from light. Pigments in Photosystem I (PS I), or Reaction Centre I (RC I), absorb light of 700 nm, while pigments in Photosystem II (PS II), or Reaction Centre II (RC II), absorb light at 680 nm.

In oxygenic photosynthesis carried out by cyanobacteria, algae and plants, light energy is used not only to generate a proton motive force but also simultaneously to reduce NADP⁺ to NADPH. They accomplish this by two kinds of reaction centres, PS I and PS II. The Chl *a* in PS I is energized by light at 700 nm, whereas

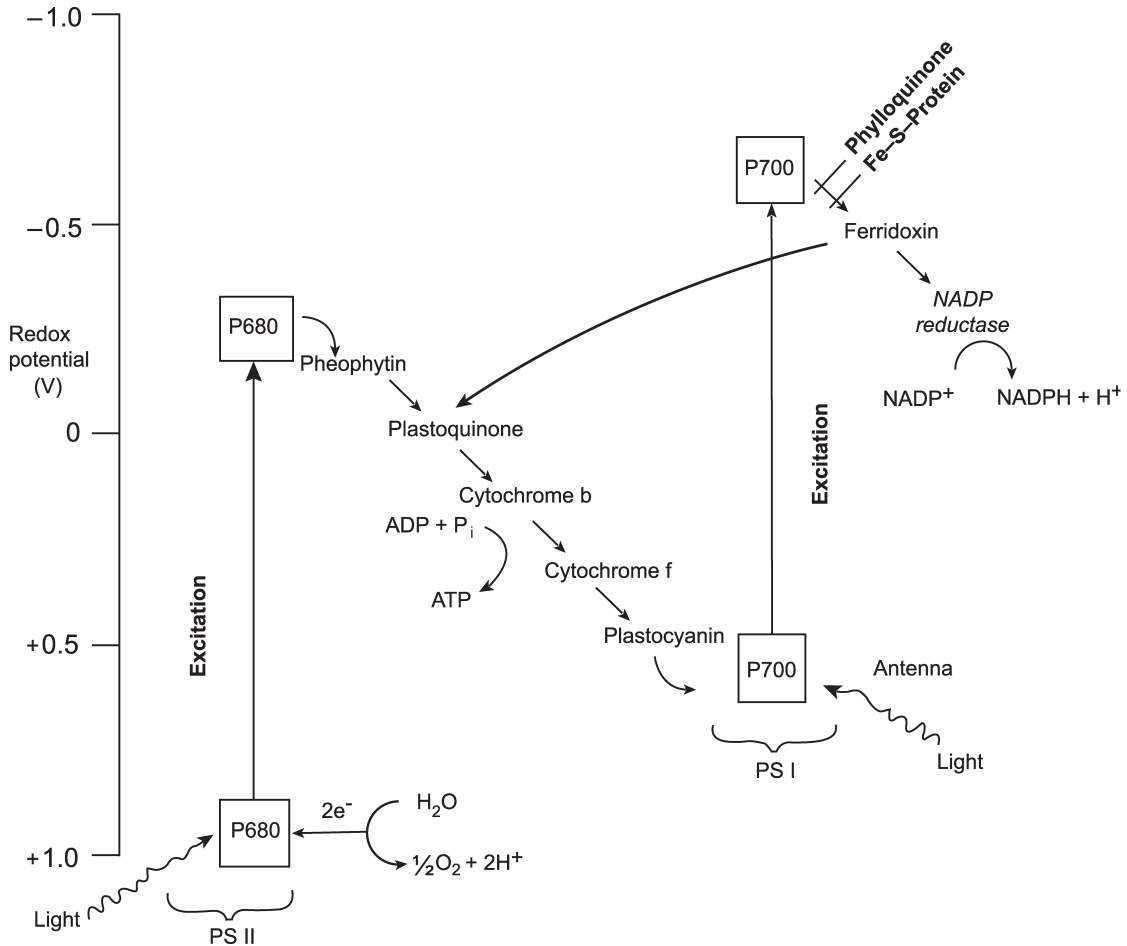


Fig. 1.3 Electron flow in oxygenic photosynthesis carried out by photosynthetic cyanobacteria, algae and green plants.

the Chl *a* in PS II is energized by light at 680 nm. The electrons from the excited reaction centres can then travel through one of two routes as shown in Fig. 1.3.

In one route, light-energized Chl *a* is oxidised by phylloquinone, which then passes the electron to Fe-S proteins and then to ferredoxin. The ferredoxin reduces NADP^+ to NADPH. The ferredoxin can also pass the electron to plastoquinone (arrow) instead of NADP^+ and generate a proton motive force. In PS II, the light energized Chl *a* is oxidized by pheophytin (a Mg-deficient form of Chl *a*), which passes the electron to plastoquinone and then to a cytochrome bf complex to generate a proton motive force. The electron

finally passes to plastocyanin and then to the electron-deficient Chl *a* in PS I. These reactions leave Chl *a* in PS II deficient in electrons. The electrons are replaced in PS II by a process called photolysis of water, in which water serves as an external electron donor. Cyanobacteria, algae and green plants are responsible for producing almost all of the oxygen in the Earth's atmosphere.

7.2.2. Anoxygenic photosynthesis

Green and purple sulphur bacteria differ from cyanobacteria because most of them are strict anaerobes and do not use water as an electron source. The green and purple sulphur bacteria

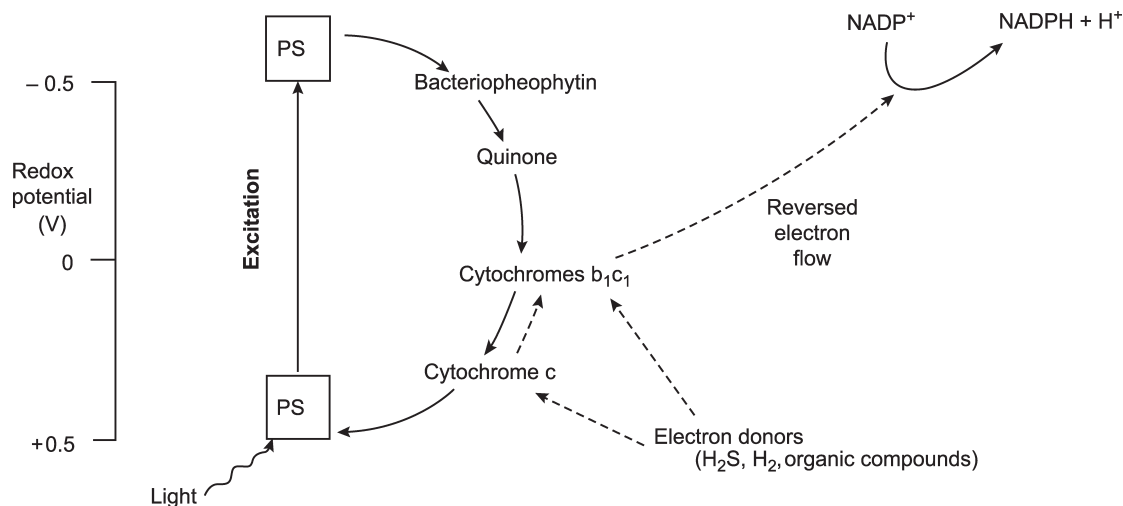


Fig. 1.4 Electron flow in anoxygenic photosynthesis carried out by the purple bacteria.

use H_2 , H_2S and elemental sulphur as electron donors and possess different light-harvesting pigments called bacteriochlorophylls. They absorb light at longer wavelengths. Bacteriochlorophyll *a* has an absorption maximum at 775 nm, while bacteriochlorophyll *b* has a maximum at 790 nm. The Photosystem (PS) or Reaction Centre in purple bacteria is called P870, while that in green bacteria is called P840 (denoting the absorption wavelength maxima associated with them). Both bacterial groups exhibit cyclic electron transport which can be used to generate ATP. They are unable to synthesize $\text{NADPH} + \text{H}^+$ directly by photosynthetic electron movement. The generalized photosynthetic electron flow in anoxygenic photosynthesis is shown in Fig. 1.4. Briefly, an electron is removed from the PS by bacteriopheophytin and passed along to ubiquinone. Electrons pass from ubiquinone through an electron transport system (creating a proton motive force, Δp) and return to the PS restoring it to the reduced state. Purple bacteria generate $\text{NADPH} + \text{H}^+$ by *reversed electron flow* to drive electrons from organic compounds or inorganic compounds to NADP^+ and energized by the proton motive force. Alternatively, in the presence of H_2 , $\text{NADPH} + \text{H}^+$ can be produced directly as H_2 has a reduction potential more negative than

NAD^+ . In the green sulphur bacteria, the primary acceptor of electrons from the PS is not bacteriopheophytin but an isomer called bacteriochlorophyll 663. The subsequent electron acceptor is not a quinone but ferredoxin. Green sulphur bacteria also exhibit a form of non-cyclic photosynthetic electron flow in order to reduce NAD^+ (not shown in Fig. 1.4). They oxidize sulphide to sulphur, with donation of electrons to the PS, to bacteriochlorophyll 663 and then to an iron sulphur-cytochrome *b* complex with ferredoxin serving as the immediate donor of electrons to NAD^+ . The elemental sulphur accumulates as refractile granules outside of the bacterial cells.

8 CATABOLIC PATHWAYS

Besides being the most nutritionally versatile of all living organisms, bacteria are also the most versatile in their ability to extract energy from oxidation of chemical compounds or from phototrophic processes. They exhibit all the metabolic sequences found in other life forms and, as well, possess metabolic sequences that are uniquely bacterial.

Bacteria and other microorganisms use a wide variety of chemical compounds as energy sources. Sometimes these compounds are large

molecules like proteins, lipids or polysaccharides that must first be broken down to smaller molecules before they can be dissimilated or used to supply energy. Microorganisms use enzymes to break down proteins to amino acids, fats to glycerol and fatty acids, and polysaccharides to monosaccharides.

Chemotrophic microorganisms derive energy by a series of consecutive enzyme-catalysed chemical reactions called a *dissimilatory pathway*. Such pathways serve not only to liberate energy from nutrients but also to supply precursors or building blocks from which a cell can construct its structure.

Although highly diverse in their metabolism, microorganisms, such as bacteria do not have individually specific pathways for each of the substances they dissimilate. Instead, a relatively few number of *central pathways* of metabolism that are essential to life are found in bacteria and, indeed, in all cellular forms. The ability of bacteria to oxidize a wide spectrum of compounds for energy reflects their ability to channel metabolites from unusual substances into the central metabolic pathways.

The central metabolic pathways are those pathways that provide precursor metabolites, or building blocks, for all of the biosynthetic pathways of the cell. They also generate ATP and reduced coenzymes, such as NADH, that can enter electron transport systems and generate a proton motive force. The most important central metabolic pathways are the Embden-Meyerhof-Parnas pathway (also called the glycolytic pathway), the pentose phosphate pathway, and the Entner-Doudoroff pathway. The last pathway has been found only among the prokaryotes.

8.1 Glycolysis

Glycolysis ('splitting of sugar') is the most common dissimilatory pathway; it occurs widely and is found in animal and plant cells as well as in microorganisms. The majority of microbes utilise the glycolytic pathway for the catabolism of carbohydrates such as glucose and fructose. This series of reactions occurs in the cytosol of microbes and can operate either

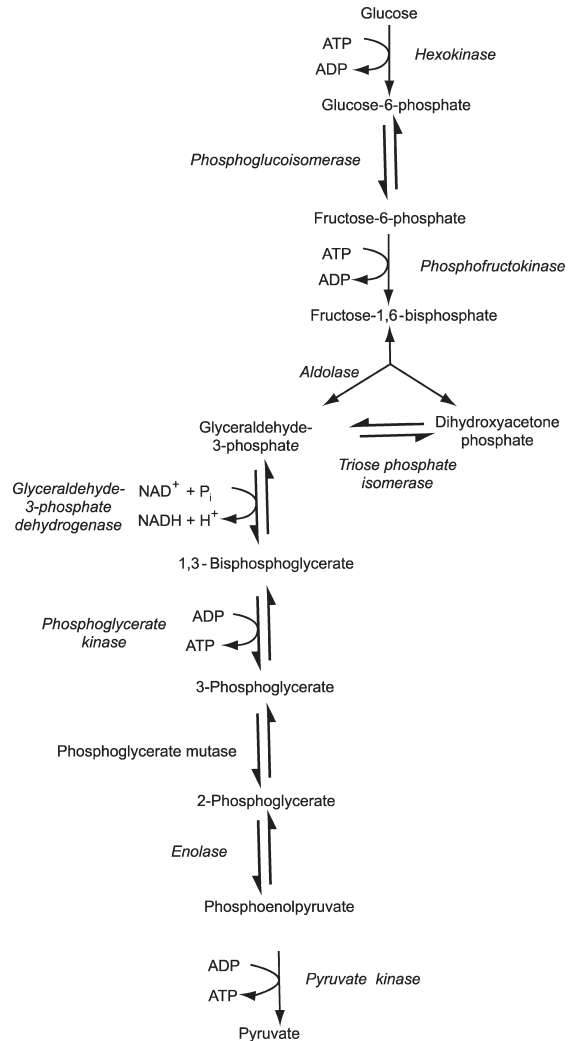


Fig. 1.5 The glycolytic pathway.

aerobically or anaerobically. Fig. 1.5 shows the steps in the glycolytic pathway.

Important features of glycolysis are discussed as follows:

- An enzyme called hexokinase uses the energy of ATP to add a phosphate group to glucose to form glucose-6-phosphate. Similarly, an enzyme, called phosphofructokinase, uses ATP to add a phosphate group to fructose-6-phosphate to form fructose-1,6-diphosphate. This 6-carbon compound is then split by an enzyme,

called aldolase, into two 3-carbon moieties, dihydroxyacetone phosphate and glyceraldehyde-3-phosphate. These two compounds are interconvertible by the action of triose isomerase.

- The oxidation of glyceraldehyde-3-phosphate results in the removal of a pair of electrons by NAD^+ and the addition of a phosphate group to form 1,3-diphosphoglyceric acid. This high-energy compound can be used for ATP synthesis by substrate-level phosphorylation. Similarly, the removal of H_2O from 2-phosphoglycerate results in the high-energy compound phosphoenolpyruvate, which also can be used to synthesise ATP by substrate-level phosphorylation.
- For each molecule of glucose metabolized, two molecules of ATP are used up and four molecules of ATP are formed. Therefore, for each molecule of glucose metabolized by glycolysis, there is a net yield of two ATP molecules.

The overall equation for the glycolytic pathway is:



In the absence of oxygen, the electrons removed from glyceraldehyde-3-phosphate can be used to reduce pyruvic acid to lactic acid or ethanol or other products. In organisms having electron transport systems, the electrons can be used to generate a proton motive force. That is, $\text{NADH} + \text{H}^+$ can be used to produce energy via oxidative phosphorylation.

8.2 Pentose phosphate pathway

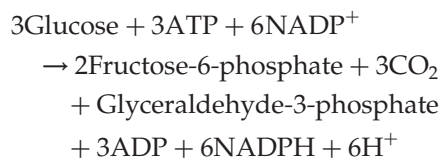
This pathway is also called the oxidative pentose pathway and the hexose monophosphate shunt. It has been called the latter because it involves some reactions of the glycolytic pathway and therefore has been viewed as a shunt of glycolysis. It exists in both prokaryotic and eukaryotic cells. It may operate at the same time as glycolysis or the Entner-Doudoroff pathway. Like glycolysis, it

can operate either in the absence or presence of oxygen.

The pentose phosphate pathway is an important source of energy in many microorganisms. Glucose can be oxidized with the liberation of electron pairs, which may enter an electron transport system. However, its major role appears to be for biosynthesis. The pathway does provide reducing power in the form of NADPH, which is required for many biosynthetic reactions instead of NADH, and it provides 4C and 5C sugars for use in nucleotide and aromatic amino acid synthesis. However, the pentose phosphate pathway cannot serve as the sole carbohydrate pathway in an organism since it cannot provide many of the required intermediates for growth. Hence, as mentioned, it usually operates in conjunction with other pathways or sequences. For instance, glyceraldehyde-3-phosphate can be used to generate energy via the glycolytic and the Entner-Doudoroff pathways.

The basic outline of the pathway is depicted in Fig. 1.6.

Glucose is initially phosphorylated to form glucose-6-phosphate. This is oxidized to 6-phosphogluconic acid with the simultaneous production of NADPH. An enzyme, called 6-phosphogluconate dehydrogenase, catalyses the decarboxylation of 6-phosphogluconic acid, yielding NADPH and a 5-carbon monosaccharide, ribulose-5-phosphate. Epimerization reactions yield xylulose-5-phosphate and ribose-5-phosphate (not shown in figure). If not used for RNA and DNA synthesis, these two compounds can be the starting point for a series of transketolase and transaldolase reactions leading subsequently to glyceraldehyde-3-phosphate and fructose-6-phosphate, which can feed into the glycolytic pathway, thereby generating ATP by substrate-level phosphorylation. The overall reaction of the pentose phosphate pathway is as follows:



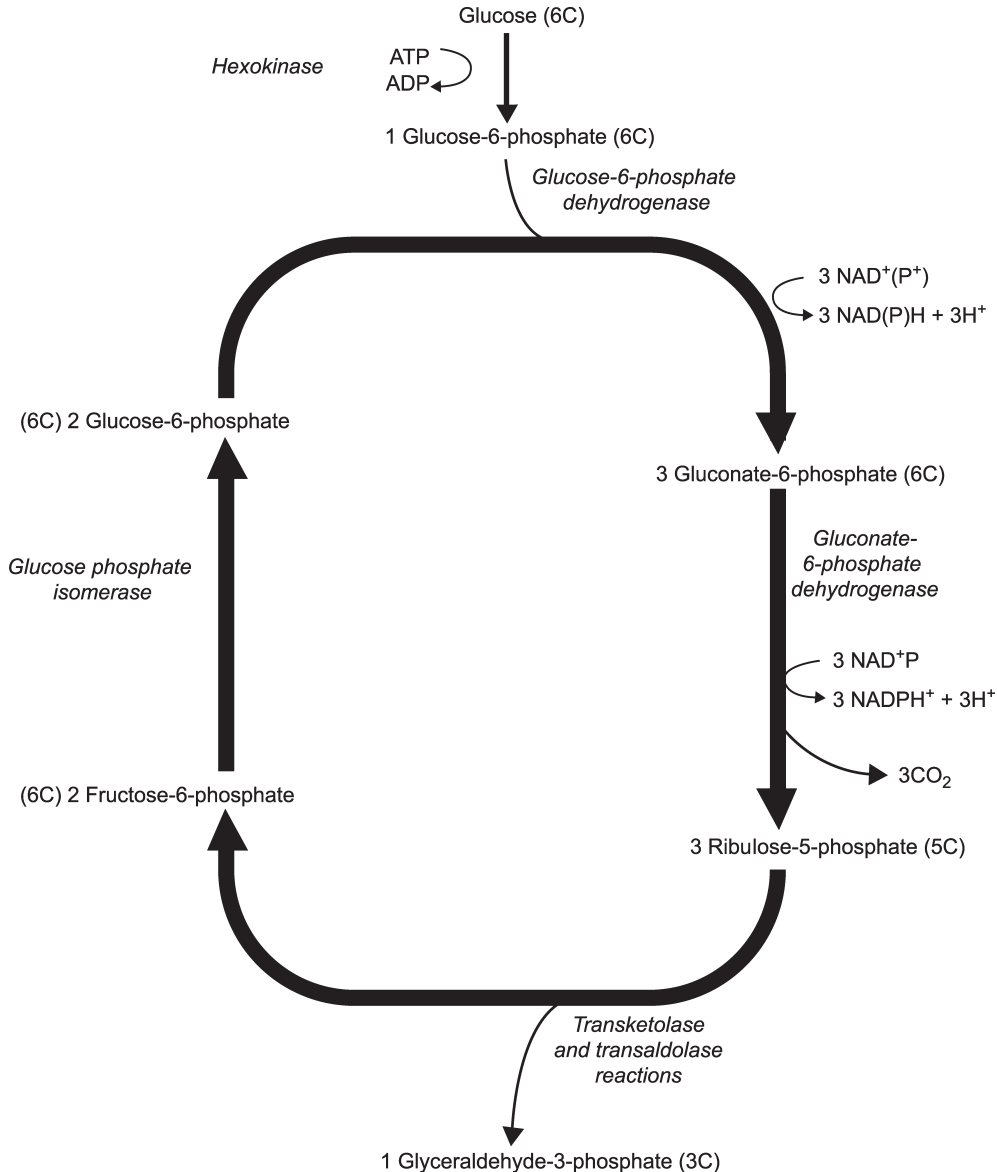


Fig. 1.6 Basic scheme of the pentose phosphate cycle.

8.3 The Entner–Doudoroff Pathway

The *Entner-Doudoroff pathway* occurs in a small number of aerobic Gram-negative bacteria including *Pseudomonas*, *Rhizobium* and *Agrobacter*. It is used mainly by bacteria that are unable to grow on glucose by glycolysis because they lack certain glycolytic enzymes but can grow on gluconic acid. Glucose, however, can be used via the Entner-Doudoroff pathway by being first phosphorylated to

glucose-6-phosphate and then oxidized to 6-phosphogluconic acid. The sequence of reactions of the pathway is shown in Fig. 1.7. If gluconic acid is used, it is phosphorylated directly to 6-phosphogluconic acid by an enzyme called glucokinase (not shown in Fig. 1.7). Then 6-phosphogluconate dehydratase catalyses the removal of a molecule of water from 6-phosphogluconic acid to yield 2-keto-3-deoxy-6-phosphogluconic acid (KDPG). KDPG aldolase then catalyses the cleavage of KDPG

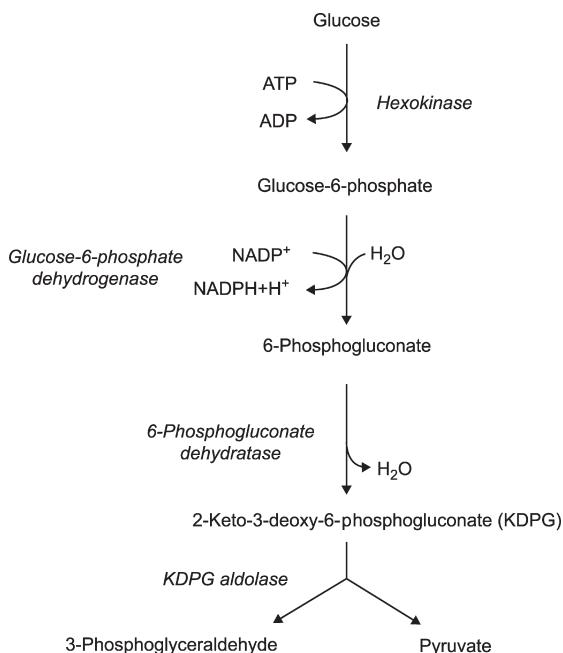
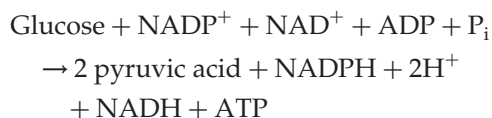


Fig. 1.7 The Entner-Doudoroff pathway.

to pyruvic acid and glyceraldehyde-3-phosphate. The latter is metabolized via glycolysis reactions to produce a second molecule of pyruvate. In many aerobic bacteria the dissimilation is completed via acetyl CoA and the citric acid cycle. The overall reaction of the Entner-Doudoroff pathway is:



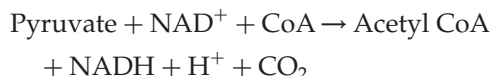
8.4 The citric acid cycle

Although the previous pathways discussed provide energy, some of the critical intermediates, and reducing power for biosynthesis, none of them provides all of the critical biosynthetic intermediates. Furthermore, the energy yield obtainable from the dissimilation of pyruvate by one of the previous pathways is limited; a significantly greater yield can be attained in the presence of oxygen from the further oxidation of pyruvate to carbon dioxide via the *citric acid cycle*, also known as the *Krebs*

cycle or the *tricarboxylic acid cycle*. This cycle provides the remaining critical intermediates and large amounts of additional energy.

Although the citric acid cycle in its complete form is found in aerobic microorganisms, even those that do not use oxygen in energy metabolism display most, if not all, of the citric acid cycle reactions. For instance, obligate anaerobes typically contain all of the reactions of the citric acid cycle with the exception of the alpha-ketoglutarate dehydrogenase. This is a physiological necessity since the critical intermediates for biosynthesis must be supplied for all organisms, irrespective of the way in which they obtain energy. Such a cycle is termed *amphibolic* because it participates in both dissimilative energy metabolism and in the production of biosynthetic intermediates.

The sequence of reactions of the citric acid cycle is shown in Fig. 1.8. Pyruvate does not enter this pathway directly; it must first undergo conversion into acetyl coenzyme A (acetyl CoA). This reaction is catalysed by a three-enzyme complex called the *pyruvate dehydrogenase complex*. This complex contains three kinds of enzymes: pyruvate dehydrogenase, dihydrolipoic transacetylase and dihydrolipoic dehydrogenase. The overall reaction is as follows:



In the citric acid cycle, initially each acetyl CoA molecule is condensed with oxaloacetic acid to form the 6-C citric acid. For every acetyl CoA entering the cycle:

- 3 molecules of NADH are generated
- One molecule of FADH_2 is generated in the oxidation of succinic acid to fumaric acid. (FAD is used here because the E_0' of the NAD^+/NADH system is not positive enough to allow NAD^+ to accept electrons from succinate)
- One molecule of guanosine triphosphate (GTP) is generated by substrate-level phosphorylation. The GTP is energetically equivalent to an ATP ($\text{GTP} + \text{ADP} \rightarrow \text{GDP} + \text{ATP}$).

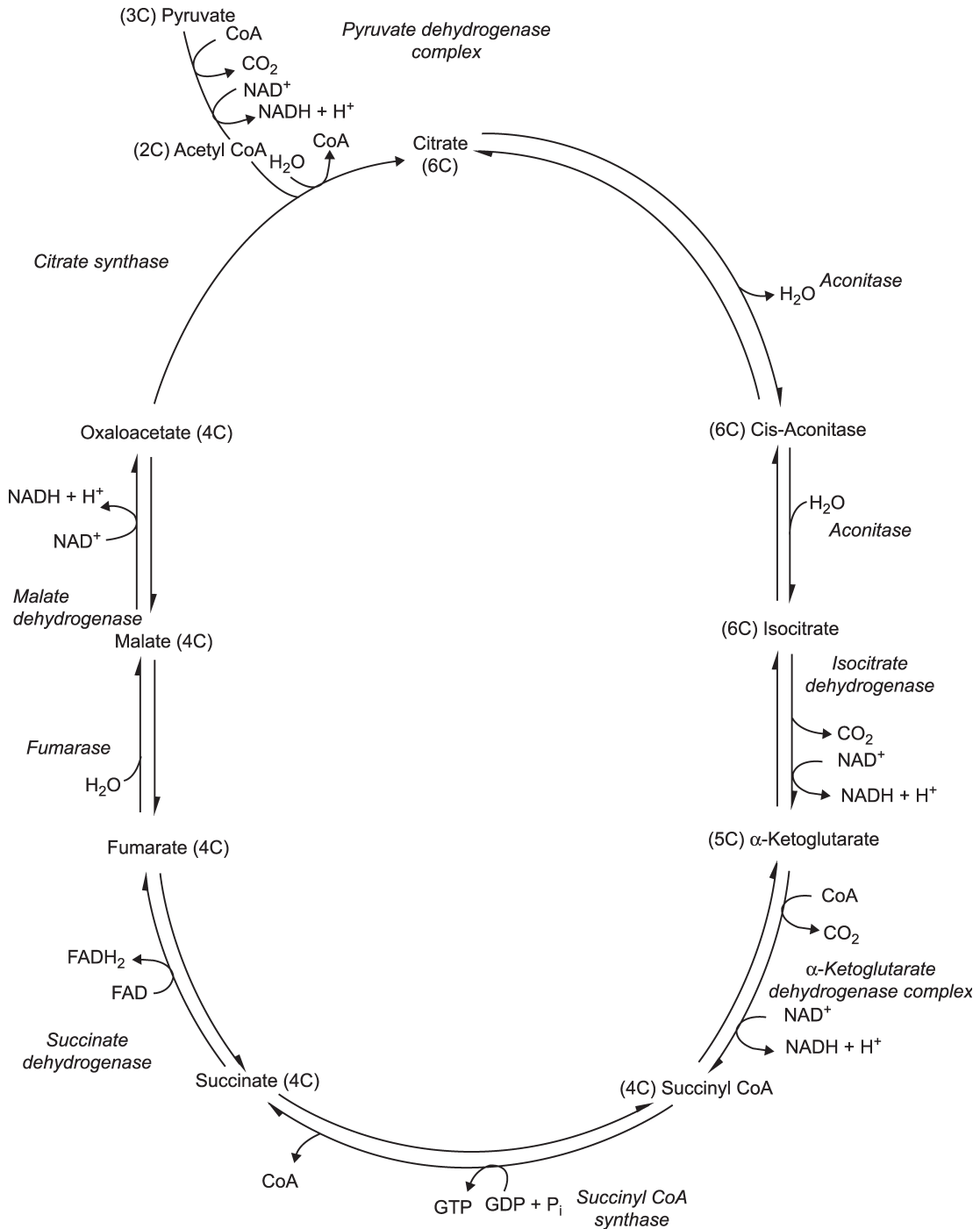
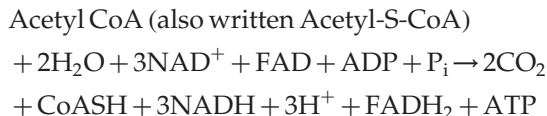


Fig. 1.8 The citric acid cycle (also called the Krebs cycle and the tricarboxylic acid cycle).

The overall reaction of the citric acid cycle is as follows:

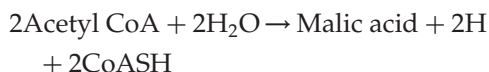


8.5 The glyoxylate cycle

The *glyoxylate cycle* or *glyoxylate bypass* is employed by aerobic bacteria to grow on fatty acids and acetate as the sole carbon source. This pathway does not occur in animals because they are never forced to feed on 2-C molecules alone. It occurs also in plants and protozoa.

The glyoxylate cycle resembles the citric acid cycle except that it bypasses the two decarboxylations in the citric acid cycle. For this reason, the acetyl CoA is not oxidized to CO_2 . A scheme of sequences of the glyoxylate cycle is shown in Fig. 1.9. The glyoxylate

cycle shares with the citric acid cycle the reactions that synthesize isocitrate from acetyl CoA. The two pathways diverge at isocitrate. In the glyoxylate cycle the isocitrate is cleaved to succinate and glyoxylate by the enzyme isocitrate lyase. The glyoxylate condenses with acetyl CoA to form malate, in a reaction catalysed by malate synthase. The malate is then used to replenish the oxaloacetate that was used up at the beginning of the cycle. The overall reaction of the glyoxylate cycle is:



Enzymes such as isocitrate lyase and malate synthase, which carry out replenishment reactions such as this, are known as *anaplerotic enzymes*; they function to maintain a pool of essential intermediates for biosynthesis.

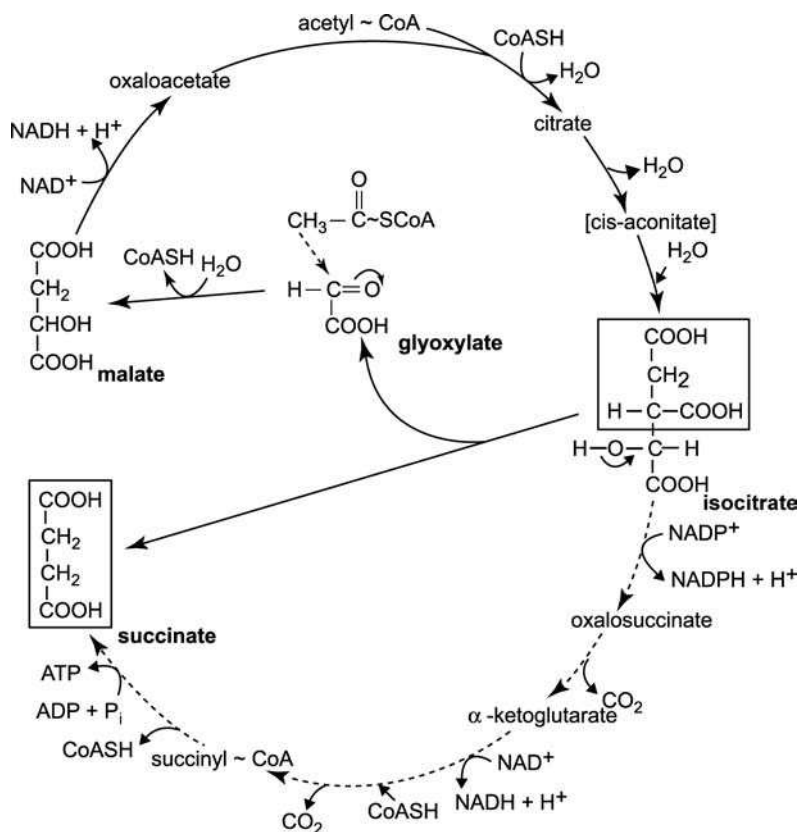


Fig. 1.9 The glyoxylate cycle or bypass. The dotted arrows represent the reactions of the citric acid cycle that are bypassed.

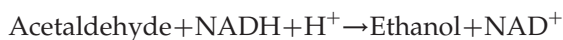
9 FERMENTATION

A major product in the catabolic pathways discussed above is the reduced pyridine nucleotides. A cell contains only a very limited amount of NAD^+ and NADP^+ . A means for continuously regenerating these from the reduced forms must exist in order for the central metabolic pathways of dissimilation to continue. Microorganisms do this either by *fermentation* or by *respiration*.

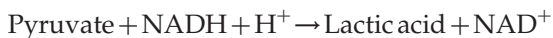
Fermentation is a metabolic process in which the reduced pyridine nucleotides produced during glycolysis or other dissimilatory pathways are used to reduce an organic electron acceptor that is synthesized by the cell itself, i.e. an endogenous electron acceptor. Many microbes utilize derivatives of pyruvate as electron and H^+ acceptors and this allows $\text{NAD(P)H} + \text{H}^+$ to be reoxidized to NAD(P) . For example, when yeast cells are grown under anaerobiosis, they carry out an *alcoholic fermentation*. After making pyruvate by glycolysis, they remove a molecule of CO_2 from pyruvate to form acetaldehyde:



The acetaldehyde is the acceptor for the electrons of the $\text{NADH} + \text{H}^+$ produced during glycolysis and becomes reduced to ethanol, thus regenerating NAD^+ :



Other microorganisms use different fermentations to regenerate NAD^+ . Lactic acid fermentation is a common type of fermentation characteristic of lactic acid bacteria and some *Bacillus* species. For instance, *Lactobacillus lactis* carries out a lactic acid fermentation by using pyruvic acid itself as the electron acceptor:



Just as the alcoholic fermentation is of great importance to the alcoholic beverage industry, the lactic acid fermentation is important for the dairy industry. The many other types of fermentations carried out by bacteria lead to various end products such as propionic acid, butyric acid, butylene glycol, isopropanol, and acetone.

Knowledge of the kinds and amounts of products made by particular microorganisms is often helpful in identifying that organism. For example, formic acid fermentation is associated with the bacterial family *Enterobacteriaceae*. Formic acid fermentation can be divided into mixed acid fermentation, which results in the production of ethanol, and a mixture of acids such as acetic, lactic and succinic. This is a characteristic of the human intestinal bacteria *Escherichia coli* and *Salmonella* spp. found in polluted water. The second type is butanediol fermentation where 2,3-butanediol, ethanol, and smaller amounts of organic acids are formed. This is characteristic of *Serratia* spp. A different type of bacterial fermentation is that of amino acids by the anaerobic clostridia. These reactions can produce ATP by oxidizing one amino acid and using another as an electron acceptor in a process called the *Stickland reaction*.

Fermentation is an inefficient process for extracting energy by the cell because the end products of fermentation still contain a great deal of chemical energy. For example, the high energy content of the ethanol produced by yeasts is indicated by the fact that ethanol is an excellent fuel and liberates much heat when burned.

10 RESPIRATION

Respiration is the other process for regenerating NAD(P)^+ by using NAD(P)H as the electron donor for an electron transport system (which was described previously). It is much more efficient than fermentation for yielding energy. Not only is NAD(P)^+ regenerated but the electron transport system generates a proton motive force that can be used to power the synthesis of additional ATP molecules (see Fig. 1.1). For instance, when yeast cells are grown aerobically with glucose, the NADH molecules produced during glycolysis can donate their electrons to an electron transport system that has oxygen as the terminal electron acceptor (aerobic respiration). This system allows not only regeneration of NAD^+ , but also the generation of enough of a proton motive force to drive the synthesis of an additional 6 molecules of ATP. Further breakdown

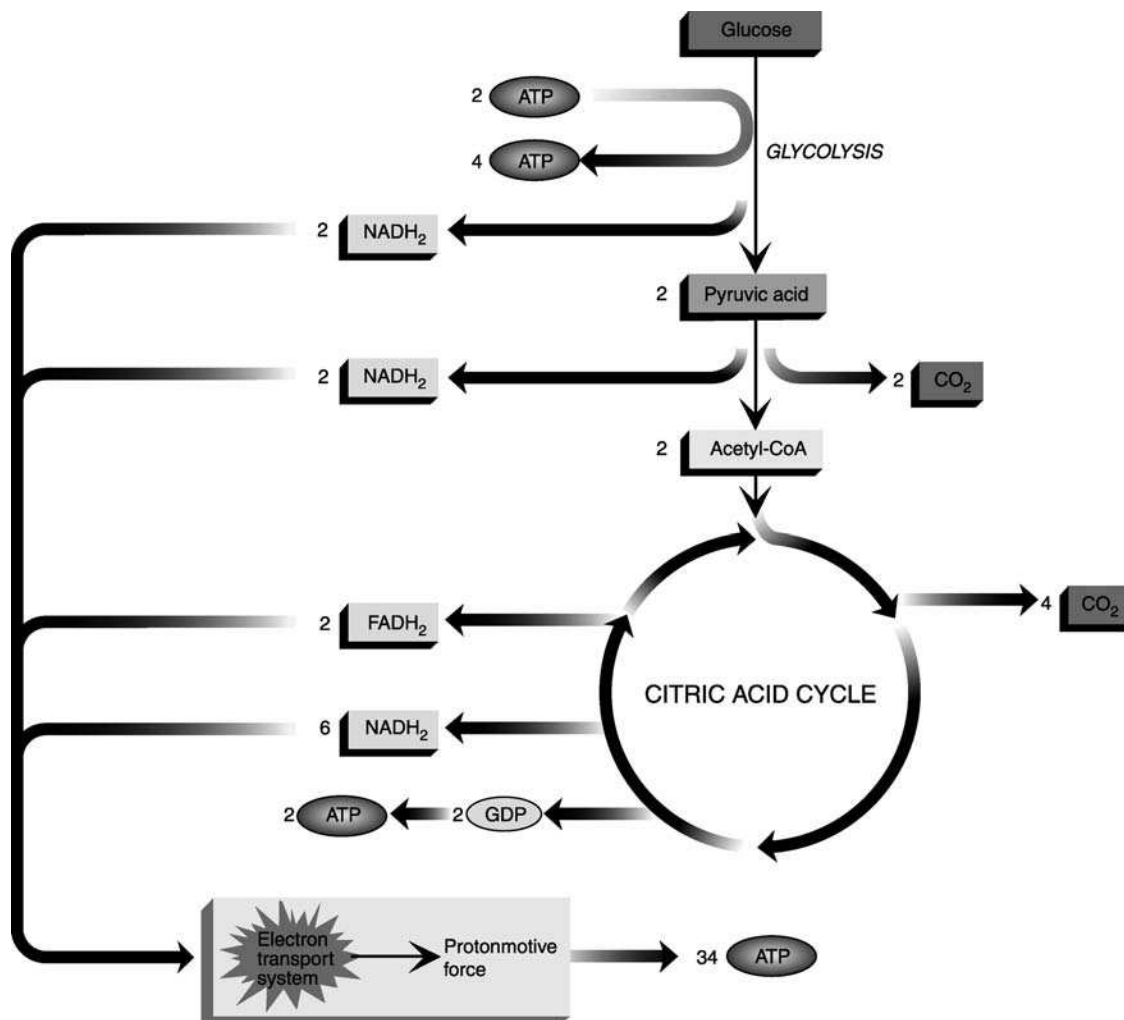


Fig. 1.10 ATP production by aerobic respiration. The complete breakdown of glucose to 6 molecules of CO₂ results in a net yield of 38 ATP molecules.

occurs when pyruvic acid is oxidized to acetyl CoA by pyruvate dehydrogenase. Each of the two molecules of NADH so formed can serve as the electron donor for an electron transport system, creating a proton motive force that can be used for synthesizing 6 molecules of ATP. Oxidation of the acetyl CoA by the citric acid cycle yields 6 more NADH which can be used to make 18 ATP. In addition, two molecules of FADH₂ are produced that can provide enough energy for the synthesis of 4 molecules of ATP. Two additional ATPs are made by substrate level phosphorylation. Therefore, the net yield of ATP from complete

dissimilation of one glucose molecule is 38 molecules (Fig. 1.10). This is in sharp contrast to the yield of ATP from fermentation when yeast cells are grown anaerobically, where the yield is only 2 molecules of ATP per molecule of glucose.

11 ENERGY-REQUIRING METABOLIC PROCESSES

How microorganisms trap chemical and light energy within high energy transfer compounds and the proton motive force has

been discussed up to this point. This chapter concludes with a brief survey of how this energy is used to fuel the various endergonic processes essential to the life of the cell. No attempt is made to provide a comprehensive catalogue of metabolic reactions requiring energy. By intent as well as by the demands of space limitation, only some important energy-requiring processes will be introduced to illustrate how trapped energy is used by microorganisms to maintain themselves and to grow.

Some energy-requiring metabolic processes are biosynthetic ones by which the complex chemical constituents of a cell are constructed. Energy in the form of ATP is needed to power the biosynthesis of many of the various chemical components of the cell, such as DNA, RNA, enzymes, bacterial cell wall peptidoglycan, and cell membrane phospholipids. In addition to being used for biosynthetic processes, energy is also used for several non-biosynthetic functions of the cell, including the inward transport and accumulation of nutrients, the export of certain proteins from the cell, and the powering of the mechanism for flagellar motility.

12 BIOSYNTHETIC PROCESSES

An autotrophic bacterium that can synthesize all its cellular constituents from simple inorganic compounds obviously has great biosynthetic ability. In the same way, a heterotrophic bacterium that can grow in a medium containing only glucose as the carbon and energy source, ammonium sulphate as the nitrogen and sulphur source, and a few additional inorganic compounds, also has great biosynthetic capability. From these nutritional substances, the microbe can synthesize:

1. nitrogenous substances, including proteins (such as enzymes) and nucleic acids
2. carbohydrates, including complex polysaccharides (such as capsules and the peptidoglycan of the cell wall)
3. phospholipids (such as components of the cytoplasmic membrane).

Details of the various specific biosynthetic pathways may be found in biochemistry and microbial physiology textbooks (such as those cited in the references to this chapter). Suffice it here to make some generalizations because all these pathways have fundamental features in common:

- A biosynthetic pathway begins with the synthesis of the biochemical building blocks that are needed to make more complex substances. Such building blocks are small molecules.
- The building blocks are then energized, usually with the energy of ATP molecules. This energy is needed to establish the covalent bonds that subsequently will link the building blocks. In a very real sense, it is these small molecules that constitute the link between energy generation and energy utilisation.
- The energized building blocks are joined together to form complex substances that become structural or functional parts of the cell.

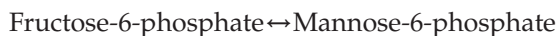
12.1 Biosynthesis of carbohydrates

A heterotrophic bacterium, such as *Escherichia coli*, can convert glucose (supplied as the major carbon source) to various other monosaccharides. For instance, it makes ribose phosphate (needed to synthesise nucleotides) from glucose by the pentose phosphate pathway. However, many bacteria, such as *Campylobacter jejuni*, cannot use glucose or any preformed sugar, probably because sugars are impermeable to the cytoplasmic membrane. Nor do they fix carbon dioxide. Such microorganisms must synthesize all of their sugars from non-carbohydrate precursors by a process called *gluconeogenesis*. The pathway involved is the reverse of glycolysis except for three sites where the glycolytic reaction is irreversible. At these sites the glycolytic enzyme is replaced by another enzyme to bypass the irreversible reaction:

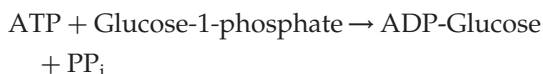
1. Conversion of pyruvate into phosphoenolpyruvate is catalysed by phosphoenolpyruvate carboxykinase (glycolytic enzyme is pyruvate kinase)

2. Conversion of fructose-1,6-bisphosphate into fructose-6-phosphate is catalysed by fructose bisphosphatase (glycolytic enzyme is phosphofructokinase)
3. Conversion of glucose-6-phosphate into glucose is catalysed by glucose-6-phosphatase (glycolytic enzyme is hexokinase).

Fructose can also be synthesized by this route. The synthesis of other sugars can be by simple rearrangements. For example, the enzyme mannose-6-phosphate isomerase catalyses



In synthesizing polysaccharides from monosaccharides, a cell cannot merely glue sugars together. Energy is required to establish the necessary covalent linkages. This energy is usually in the form of ATP and is expended only inside the cell to energize the monosaccharides. For example, an energized form of glucose is uridine diphosphate glucose (UDPG). Nucleoside diphosphate (ADP-linked) sugars are required for the synthesis of polysaccharides as shown by the following equations:



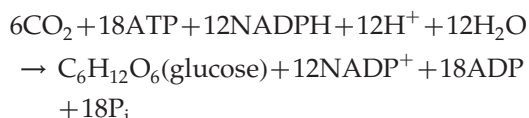
The biosynthesis of bacterial cell wall peptidoglycan is an endergonic process. Although the polymer is located in the cell wall, most of the chemical energy needed for its synthesis is expended inside the cell. The process is highly complex and the resulting macromolecule is a rigid cross-linked material that maintains the shape of the bacterial cell.

12.2 Carbon dioxide fixation by autotrophs

Autotrophic microorganisms are capable of using CO₂, the most oxidized form of carbon, from the atmosphere as their major source of carbon and reducing it to organic compounds. This process is called *carbon dioxide fixation* and is an endergonic process. Autotrophs use ATP

as the energy source for CO₂ fixation and NADH or NADPH as the electron source for the reduction.

The main method of CO₂ fixation in autotrophic organisms is the *Calvin cycle*. The reactions of the Calvin cycle are also recognized as the *dark reactions* of photosynthesis because green plants, algae, cyanobacteria, and the purple bacteria use the Calvin cycle for CO₂ fixation. The Calvin cycle is depicted in Fig. 1.11. In the initial reaction of the Calvin cycle, CO₂ is added to the 5-C ribulose bisphosphate resulting in the formation of two molecules of the 3-C phosphoglyceric acid. Then NADPH or NADH, aided by the energy of ATP, provides electrons to reduce the phosphoglycerate to glyceraldehyde-3-phosphate. All of the organic compounds needed by the cell are synthesized subsequently from the phosphoglycerate and the glyceraldehyde-3-phosphate. Each turn of the cycle results in the fixation of one molecule of CO₂. To synthesize one glucose the cycle must operate six times. The overall stoichiometry of the cycle is as follows:



For a long time, the Calvin cycle was believed to be the sole means of CO₂ fixation in living organisms. Recently, additional modes of CO₂ fixation have been discovered. The *reductive TCA cycle* is known in the green bacterium *Chlorobium limicola* and certain members of the sulphate-reducing bacteria. The *acetyl CoA pathway* is widely distributed among the chemolithotrophic anaerobes including the sulphate-reducing and methanogenic microorganisms.

12.3 Nitrogen fixation and assimilatory nitrate reduction

Seventy-eight per cent of air in the atmosphere is nitrogen, but it cannot be used as a nutrient source of nitrogen by most living organisms. However, some prokaryotes, like the free-living *Azotobacter* and the legume plant symbiont *Rhizobium*, are able to use it by a process

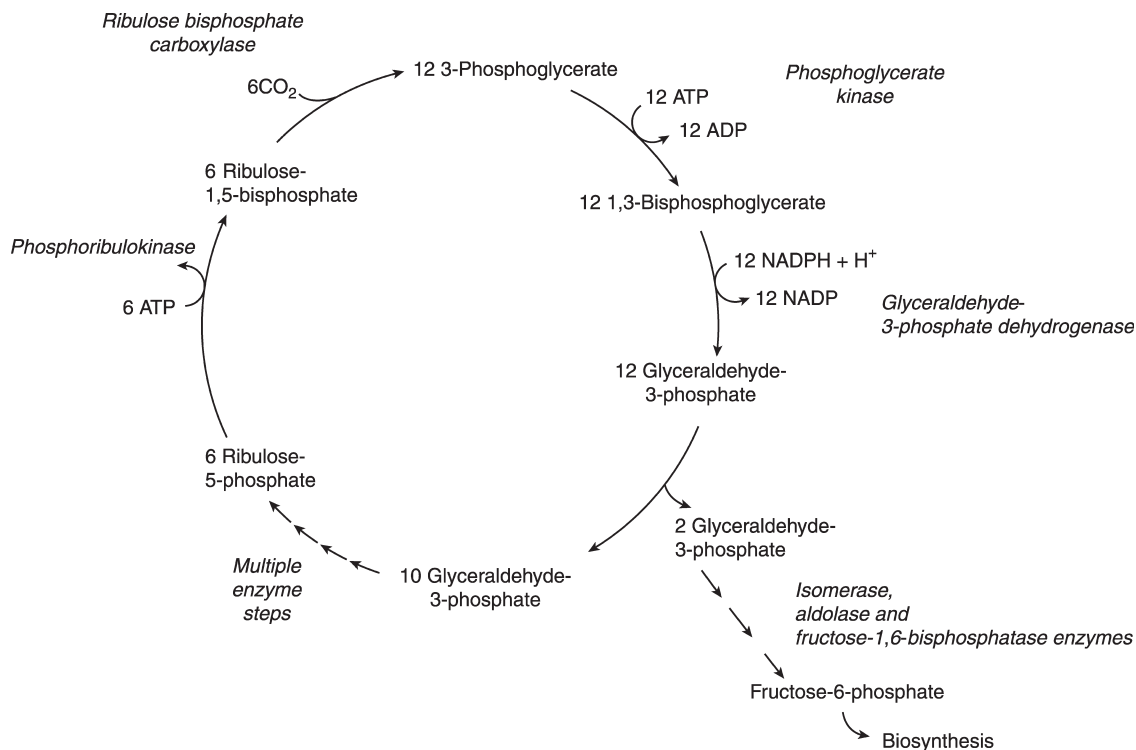


Fig. 1.11 The Calvin cycle.

called *nitrogen fixation*. The process is catalysed by a cytoplasmic nitrogenase complex consisting of two enzymes: one enzyme is dinitrogenase, which contains molybdenum and iron (a MoFe protein); the other enzyme is dinitrogenase reductase, an iron-containing enzyme (an Fe protein). The source of the electrons for N_2 reduction is usually the reduced form of the Fe-S protein ferredoxin (Fd_{red}) which has a very negative $E_{\text{O}'}$ value. Anaerobic or microaerophilic bacteria can provide Fd_{red} from oxidation of pyruvic acid by pyruvate:ferredoxin oxidoreductase. Aerobic bacteria reduce NAD^+ to NADH during pyruvic acid oxidation and thus must use the proton motive force to power reversed electron transport allowing NADH to reduce Fd. The mechanism of nitrogen fixation is shown in Fig. 1.12. The overall reaction for the process is:



Many microorganisms cannot fix nitrogen and assimilate nitrate in a process called

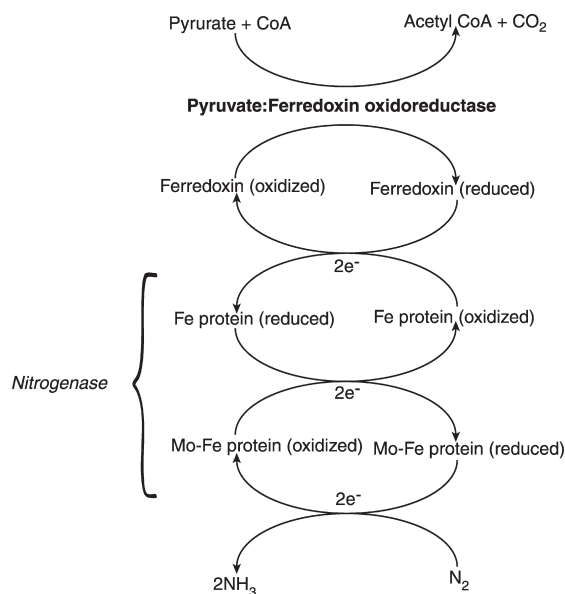
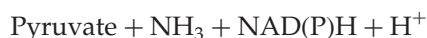


Fig. 1.12 Mechanism of nitrogen fixation.

assimilatory nitrate reduction thereby obtaining ammonia. Assimilatory nitrate reduction uses two cytoplasmic enzymes, nitrate reductase and nitrite reductase. Nitrate reductase catalyzes the reduction of NO_3^- to NO_2^- , and nitrite reductase catalyses a multistep reduction: NO_2^- to NOH (nitroxyl), a postulated intermediate that has never been isolated), NOH to NH_2OH (hydroxylamine), and NH_2OH to NH_3 .

Ammonia can be incorporated directly as it is more reduced than other forms. The simplest mechanisms found in all microorganisms involve the formation of alanine and glutamate as shown in the following reactions:



Glutamic acid occupies a key position in amino acid biosynthesis because it is involved in the synthesis of many other amino acids. Microorganisms can use alanine and glutamate to synthesize a number of new amino acids via *transamination reactions* (transfer of α -amino groups). Such amino acids include aspartic acid, serine, valine, isoleucine, leucine, arginine, histidine, phenylalanine and tyrosine. Another way by which glutamic acid can be used to make other amino acids is by the alteration of its molecular structure; proline is made in this manner.

Amino acids need to be energized before they can be linked together to make proteins. Cells energize amino acids by using the energy of ATP:



A microorganism synthesizes hundreds of different proteins, each protein having its own unique sequence of amino acids.

12.4 Biosynthesis of nucleotides

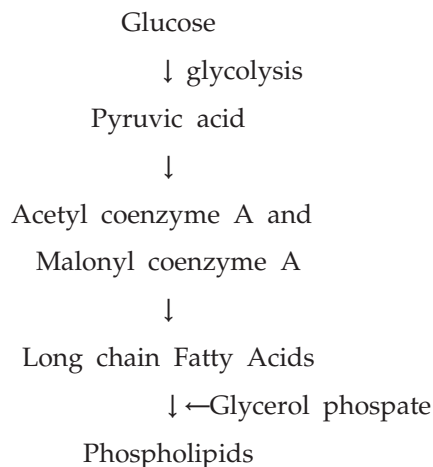
A nucleotide consists of a nitrogenous base-pentose-phosphate. When the sugar is ribose, a nucleotide is a ribonucleotide and is used for

the biosynthesis of RNA. Similarly, when the sugar is deoxyribose, a nucleotide is a deoxy-ribonucleotide and is used for the biosynthesis of DNA. Ribonucleotides and deoxyribonucleotides that have adenine or guanine as the nitrogenous base are purine nucleotides. Those that have cytosine, thymine or uracil as the nitrogenous base are pyrimidine nucleotides.

The energized form of a nucleotide is its diphosphate derivative. For instance, if the nucleotide is cytidine monophosphate (CMP), the energized form would be cytidine triphosphate (CTP)—the extra phosphate groups supplied by ATP. Prior to energization, the synthesis of pyrimidine nucleotides requires ribose phosphate (from the pentose phosphate cycle), ATP, aspartic acid and glutamine. The synthesis of purine nucleotides requires ribose phosphate, ATP, GTP and the amino acids glycine, aspartic acid and glutamine. DNA and RNA are subsequently assembled from pyrimidine and purine diphosphates.

12.5 Biosynthesis of lipids

The major lipids of microbial cells are the phospholipids that, together with proteins, form the structure of the cytoplasmic membrane. Phospholipids are made from long chain fatty acids. The general way in which microorganisms synthesise phospholipids may be depicted in this manner:



As may be noted, two important building blocks for long chain fatty acids are acetyl

coenzyme A (acetyl CoA or acetyl-S-CoA) and malonyl coenzyme A (malonyl CoA or malonyl-S-CoA). Acetyl CoA is made by the oxidation of pyruvic acid. Malonyl CoA is an energized building block made from acetyl CoA. The malonyl group contains three carbon atoms; the extra carbon comes from CO_2 via the bicarbonate ion, HCO_3^- , and energy in the form of ATP is required for the addition.

The synthesis of fatty acids is shown in Fig. 1.13. After malonyl CoA is made, the coenzyme group on each malonyl CoA and acetyl CoA is replaced by a large protein molecule called an acyl carrier protein that serves to anchor the compounds so that appropriate enzymes can act on them. Then a malonyl ACP complex is complexed with an acetyl ACP complex. In this reaction, two of the three carbon atoms of the malonyl group are added to the acetyl group to

make a 4-carbon acetoacetyl ACP molecule; the third carbon atom of the malonyl group is liberated as CO_2 . The acetoacetyl ACP is eventually converted to butyryl ACP (as shown in Fig. 1.13).

If the butyryl group were to be released from the ACP protein, it would become butyric acid, a 4-C fatty acid. To make a longer fatty acid, the cell must combine the butyryl ACP with another malonyl ACP to add another 2-carbon unit, resulting in a 6-carbon fatty acid ACP. This process of adding two carbons at a time to elongate the fatty acid chain is continued until a fatty acid of the required length is attained (usually 16 to 18 carbon atoms long).

After the fatty acids of the required length are obtained, the cell uses them to synthesize phospholipids. Glycerol phosphate is needed for its synthesis. It is made from dihydroxyace-

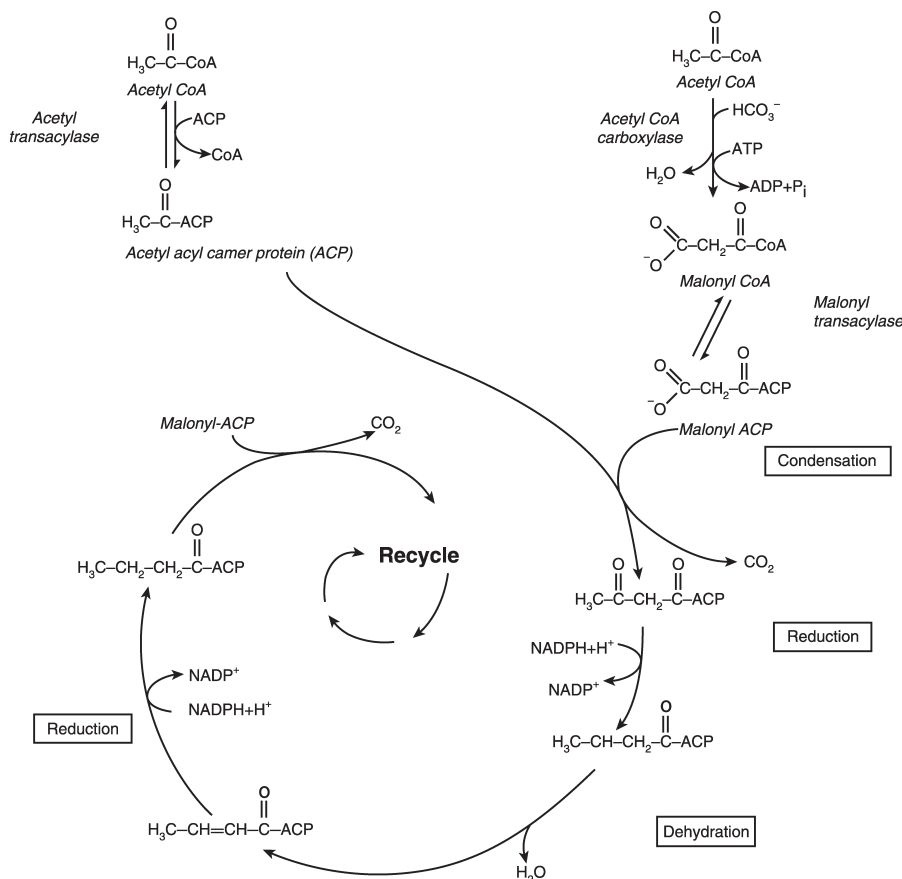
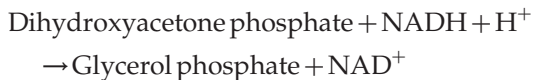


Fig. 1.13 Fatty acid synthesis. The cycle is repeated until the specific chain length is obtained.

tone phosphate, an intermediate in glycolysis:



Two fatty acid molecules are linked to one molecule of glycerol phosphate to form a molecule of phosphatidic acid, a simple phospholipid. The cell can then link other chemical groups to the phosphate group of the phosphatidic acid to make other phospholipids. For example, the amino acid serine can be added to phosphatidic acid to make phosphatidylserine. Energy in the form of cytidine triphosphate (CTP) is required for this reaction.

13 NON-BIOSYNTHETIC PROCESSES

There are non-biosynthetic processes in microbial cells that require energy. These processes include transport of nutrients into cells, protein export and motility in prokaryotic cells.

13.1 Transport of nutrients

Transport of the nutrients into cells essentially means transport of nutrient substances across the cytoplasmic membrane. When substances move across a membrane without interacting specifically with a carrier protein in the membrane, the process is called *passive diffusion*. Gases and water, and some non-ionized organic acids, go in and out of the cell by passive diffusion and are driven only by concentration gradients, i.e. they move down the gradient from high to low concentrations.

The cytoplasmic membrane is an effective barrier to most polar molecules such as the nutrient sugars and amino acids. Transport of these molecules requires the participation of *membrane transport proteins*, or *carrier proteins*, which span the membrane. Three classes of carrier proteins have been identified: *uniporters* move a single type of compound across the membrane; *symporters* move two types of compounds simultaneously in the same direction; *antiporters* move two compounds in opposite directions.

Even with carrier proteins, no metabolic energy is needed for the mere 'downhill' entry

into the cell of a solute molecule, i.e. from a greater concentration outside to a lower concentration inside the cell. Such gradient-driven carrier-mediated transport is called *facilitated diffusion*. Carrier proteins exhibit specificity for molecule binding as well as saturation kinetics.

Microbial cells can also concentrate nutrient molecules 'uphill' against a gradient. Consequently, if a bacterium lives in a dilute aqueous environment, with very low levels of nutrients, it can concentrate these nutrients inside the cell to the point where they can be readily used for metabolism. Such internal concentration of nutrients requires metabolic energy and the process is called *active transport*. The energy is used to change the affinity (strength of interaction) of the carrier protein for the particular solute. On the outer surface of the membrane the carrier protein exhibits high affinity for the substrate; during transport, the affinity of the carrier must change to a low one so that the substrate can be released into the cell. The energy for the process to change the conformations of the carrier proteins, and hence their affinities, comes from the proton motive force or ATP hydrolysis.

In the *PEP-phosphotransferase system* of nutrient transport, the solute is chemically altered by acquiring a phosphate group as the result of an enzymatic reaction during its transport across the cytoplasmic membrane. The process is also called *group translocation*. It is the only process in which there is chemical alteration (phosphorylation) of the solute during transport. The carrier has only a low affinity for the phosphorylated form of the solute. The energy for the alteration of the solute molecule comes from the high-energy compound phosphoenolpyruvic acid (PEP), which is an intermediate in glycolysis.

13.2 Export of proteins

The translocation process of proteins synthesized by cytosolic ribosomes into or through the membranes is referred to as *protein export*. The proteins end up in the cytoplasmic membrane, periplasmic space, outer envelope, cell wall, glycocalyx, cell surface appendages and extracellularly into the surrounding

environment. Examples of cytoplasmic membrane proteins include cytochromes and other respiratory enzymes as well as carrier proteins. Periplasmic proteins include binding proteins, lipoproteins and various degradative enzymes such as ribonuclease and alkaline phosphatase. Porins are found in the outer membrane. Examples of secreted proteins include exotoxins and enzymes (e.g. penicillinase, amylases, proteinases).

As to be expected, most research work on protein export has been performed with *Escherichia coli*, primarily because of the ease of genetic manipulation. This has led to the formulation of a model for the export of protein; it is called the Sec system.

Proteins destined for export are distinguished by having an extra 20 or so amino acid sequences, called the *signal sequence*, at their NH₂-terminal end when they are being synthesized within the cytoplasm by ribosomes. Signal sequences vary with the particular export protein. They are removed during transport across the cytoplasmic membrane. The protein chain to be exported must not be allowed to fold into its mature form before export. Upon leaving the ribosome, the nascent pre-protein binds to a *chaperone protein* that prevents it from assuming a tightly folded configuration that cannot be translocated. The signal sequence, or peptide, retards the folding of the pre-protein, giving the chaperone protein time to bind. The chaperone protein then delivers the exportable protein to a *docking protein*. Energy is expended when ATP initiates the transfer of the exportable protein across the cytoplasmic membrane and the proton motive force drives translocation of the rest of the protein chain.

13.3 Motility in prokaryotic cells

A microbial cell is of a very small dimension. As a consequence, water is a very viscous medium to such a cell. Microscopic fins, flippers, or fishtails would be useless for propelling such a cell in such a liquid. The analogy would be that

of a human trying to swim or row a boat on a body of thick molasses. However, by rotating its flagellum, a prokaryotic cell can move readily through water in much the same way a corkscrew can easily penetrate a piece of cork. The discs in the basal body of the flagellum are responsible for rotating the appendage. The motor apparatus that causes a flagellum to rotate is associated with the bottom-most disc in the basal body of the flagellum. The motor is a proton motor, driven by a flow of protons from the proton motive force.

14 EPILOGUE

This chapter has provided a comprehensive overview of current information on how microorganisms fill their nutritional needs and extract energy to carry out their life metabolic processes. To this end, the author has drawn freely from his experience in writing textbooks with his co-authors, Dr Michael J. Pelczar, Jr (Professor Emeritus, University of Maryland) and Dr Noel R. Krieg (Professor, University of Virginia and Technical Institute). Their latest combined contribution is *Microbiology: concepts and applications*. This chapter contribution is a grateful acknowledgement of their collaborative efforts through the years, but the author is solely responsible for the accuracy and acceptance of the assembled information in this chapter.

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Introduction to Microbes of Sanitary Importance

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Viruses

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1 INTRODUCTION

The bacteriological examination of water is routinely used to detect the presence of faecal indicators such as *Escherichia coli*, *Clostridium perfringens* and enterococci, particularly detection of *Escherichia coli*. These bacteria are easy to manipulate in the laboratory and the bacteriological examination of water is easily performed. In contrast, the virological examination of water involves more specialized techniques and relatively few laboratories undertake these duties. The viruses that can be found in water cause infections that range from those that do not cause noticeable symptoms to those that cause life-threatening infections. It is essential, therefore, that testing water for the presence of viruses is only carried out by personnel who have been properly trained in virology and, where appropriate, have proper vaccination cover.

Because some of the waterborne viruses that will be cultured during the examination of water samples are capable of causing serious disease and also because they may be present in culture extracts in large numbers, proper containment facilities must be used. The pathogens that may be cultivated require Level 2 containment facilities for safe handling. Many laboratories use simian cells in which to cultivate viruses. These must also be handled with caution. All materials used for the virological examination of water must be sterilized before and after the testing procedure to minimize the contamination of samples during testing and to prevent release of

infectious viruses from the materials after use. The need for safe handling of viruses and of materials that are potentially contaminated with viruses cannot be overemphasized.

As with the examination of water for faecal bacteria, the virological examination of water depends upon the detection of *indicator viruses*. These viruses are typically found in waters contaminated with human faeces. Not all waterborne viruses are capable of growth in tissue culture and hence the need for exploiting indicators. The most common viruses used as indicators of faecal contamination of water are the enteroviruses, particularly the poliomyelitis virus and the Coxsackie viruses. These will grow relatively easily in tissue culture. Other enteroviruses grow less well in the laboratory. These viruses can cause life-threatening infections. The reference laboratories devoted to water virology help to control such infections and literally play a vital role in the monitoring and maintenance of public health.

2 THE NATURE OF VIRUSES

Viruses are relatively simple structures. They are made up of a nucleic acid core surrounded by a protective protein coat, or capsid, made up of subunits called capsomeres. The nucleic acid of a virus may be either RNA or DNA but virus particles do not carry both nucleic acids. The RNA or DNA carries the virus genome: all the genetic information necessary to complete the virus life cycle. Retroviruses are unusual. One is HIV, the human immunodeficiency

virus. Infection with this virus is the ultimate cause of AIDS. The retrovirus particle contains an RNA genome but when the virus infects a host cell, a DNA copy is made of the virus genome. This subsequently becomes integrated into the genome of its host cell. This DNA copy of the virus RNA is made by an enzyme known as reverse transcriptase used in many molecular biological techniques.

The virus genome encodes the structural proteins that make up the capsid and all the enzyme functions necessary for virus replication and cell infection. Some viruses have a lipid envelope but this is not the case with all viruses. The envelope material may be closely associated with the capsid, or it may be loose fitting. Although very simple in their construction, viruses can display an astonishing variety of shapes. Amongst the most complex of virus particles is the T-even bacteriophage that infects *Escherichia coli*. These have

been said to resemble a lunar landing module (Fig. 2.1).

Viruses are obligate intracellular parasites. This means that to complete their life cycles, viruses depend absolutely on functions provided by cells of a host organism. Not all viruses can be grown in artificial culture. If this can be achieved, the growth of viruses in laboratories requires exploitation of tissue culture technology. When outside a host cell, the only thing a virus can do is to infect a new host. Once cellular infection has been accomplished, however, viruses become highly active and subvert the normal metabolic activity of the cell to produce new virus particles.

Following attachment to and penetration of a new host cell, all viruses show the same general life cycle (Fig. 2.2). Viruses induce the biosynthesis of virus proteins and replication of the virus genome. Typically, the enzymes that are required for virus replication are made

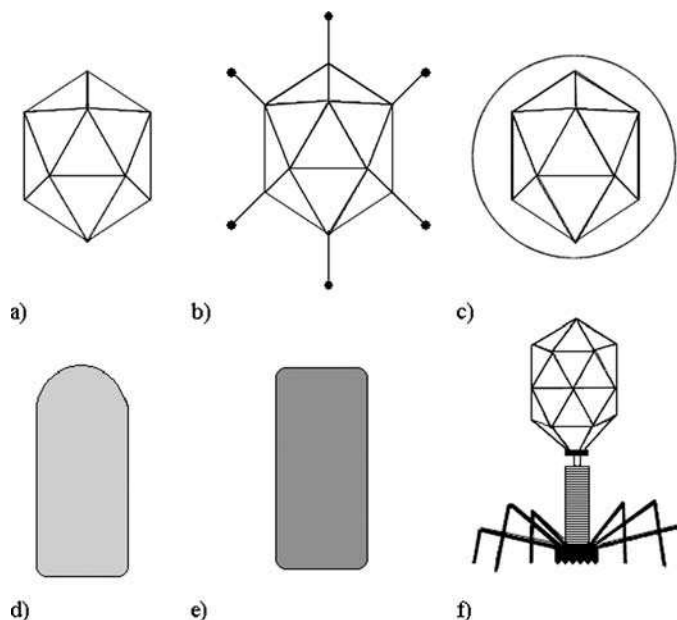


Fig. 2.1 Representative virus particles. Human viruses (a–e) are relatively simple when compared with the T-even bacteriophage that infects *Escherichia coli* (f). Many virus types are icosahedral (a). Icosahedral viruses include enteroviruses and rotaviruses, used in the virological examination of water. Adenoviruses (b) are icosahedral viruses that carry spikes on their surfaces. Some viruses, such as the herpes virus (c) have an envelope around an icosahedral core. Rabies viruses are bullet-shaped (d) and pox viruses are brick-shaped (e). These pictures are not drawn to scale; the smallest human viruses are less than 20 nm in size while the largest are over 400 nm.

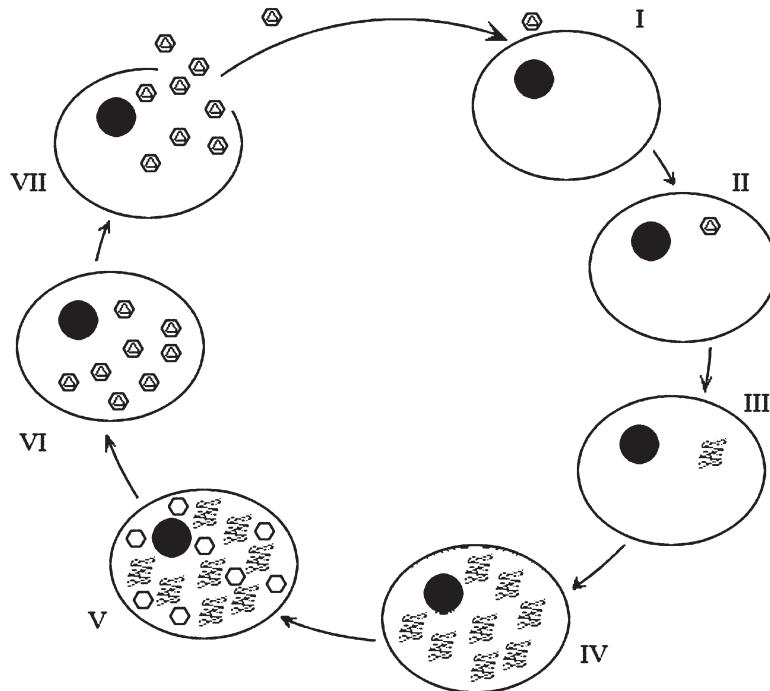


Fig. 2.2 Representative virus life cycle. The enterovirus virion approaches the surface of the cell to be infected (I). It penetrates the cell to infect it (II), and virus RNA passes into the cytoplasm as the virus is uncoated (III). Non-structural proteins are formed during early translation. The virus genome is replicated (IV) and new virus structural proteins are formed (V). New virus particles are assembled (VI) and released by lysis of the infected cell (VII). Reproduced with kind permission from Heritage, J., Evans, E.G.V. and Killington, R.A. (1996). *Introductory Microbiology*, Cambridge University Press.

soon after the virus infects a new host cell and these are referred to as 'early proteins'. The structural proteins that make up the virus capsid are produced later after infection and are thus referred to as 'late proteins'. The assembly of new virus particles and the subsequent release of progeny follows the biosynthesis of virus components. This may result from the complete destruction or lysis of the host cell. Alternatively, new virus particles may 'bud out' from their infected host cell. Viruses that have an envelope sometimes use this strategy, the envelope material being acquired as the newly emerging particles pass through the cell membrane. Not all virus envelopes are derived from the cell membrane. If virus replication occurs within the nucleus of the cell, the envelope material may be derived from the nuclear membrane within the infected cell.

Although the overall pattern of attachment, penetration, replication and release is constant

for all viruses, a wide array of strategies has evolved so that viruses can exploit their hosts. For example, enteroviruses can cause the synthesis of messenger RNA from RNA rather than DNA genome templates. The human immunodeficiency virus can make a DNA copy of its RNA genome. The vaccinia virus used for smallpox vaccination can cause the replication of DNA in the cytoplasm of its host cell rather than in its nucleus. There are many other variations of the virus life cycle theme.

All classes of organism have been subject to virus infection. The viruses that infect bacteria are known as bacteriophage and these have recently been exploited as biological markers, for example to trace the source of polluted water. Plant viruses may cause effects that horticulturists find appealing. For example, the striped appearance of tulip flowers prized by many gardeners results from a virus infection.

Animal viruses cause a spectrum of diseases. These range from sub-clinical, where the host is unaware of the infection, through to fatal conditions.

3 VIRUS INFECTIONS THAT SPREAD VIA THE FAECAL–ORAL ROUTE

Viruses that cause waterborne infections are of devastating importance. Viruses are the commonest cause of gastrointestinal infection world-wide and rotaviruses are the most important cause of life-threatening diarrhoea in children under 2 years old. About one-third of all children have experienced an episode of rotavirus diarrhoea by the time they are 2 years old and nearly all children are infected by this age, even though a significant minority show no clinical symptoms. Rotaviruses alone account for 20% of infectious diarrhoea cases in developing countries. In the developed world, rotaviruses are a significant problem for hospitalized patients of all ages. Over half of children suffering from dehydration who are admitted to hospital in the UK are suffering from virus gastroenteritis. Outbreaks of rotavirus diarrhoea are not uncommon in hospitals in the UK. There have been notorious cases associated with cruise liners where a very large number of passengers have developed diarrhoea and vomiting during their holiday. These viruses are particularly infectious. Volunteer studies have shown that fewer than 10 virus particles may cause clinical disease. This makes elimination of the infectious agent very difficult.

Although rotaviruses are a significant cause of gastrointestinal symptoms, there are a number of other viruses that are associated with diarrhoea and vomiting. These include caliciviruses, astroviruses and certain adenoviruses. These viruses are more often associated with infections in older children and adults than in infants. Infections caused by these viruses may show a remarkable seasonal variation in temperate climes. This is reflected in the folk name given to the virus-mediated 'winter vomiting disease'. The most notorious of the caliciviruses causing gastrointestinal

disease is the Norwalk agent, named after the Ohio town from which it was first recognized. It causes muscle pain, headache and nausea as well as profuse vomiting and diarrhoea, but the symptoms generally resolve very quickly.

Most of the viruses that cause diarrhoea cannot be grown in tissue culture. They may be visualized in infected faeces by electron microscopy but this is an expensive and specialized technique that is not routinely performed. Latex agglutination techniques can also be used to detect viruses that cause diarrhoea. Latex beads are coated in antibodies specific to the virus to be identified. These are then mixed with a sample of faeces to be tested. If virus particles are present, these will react with the antibodies, causing the latex particles to clump. More frequently, the diagnosis of virus gastroenteritis is made by excluding other causes for the symptoms.

Not all waterborne viruses cause symptoms of gastroenteritis. There are two other important groups of viruses that cause waterborne infections: hepatitis viruses and enteroviruses. A number of viruses are known to cause hepatitis. Clinically, the disease processes seem indistinguishable but the viruses that cause hepatitis have different structures, life-cycles and modes of transmission. Two hepatitis viruses are particularly associated with waterborne infection – hepatitis A and hepatitis E virus. Hepatitis A virus, a small RNA virus, causes acute infective hepatitis. Symptoms of the disease are of rapid onset but the condition generally resolves within a relatively short time. Cases often occur in large clusters. Patients become overwhelmingly tired in the early stages of the illness. They develop a fever and may suffer diarrhoea. Appetite is suppressed and patients complain of feeling awful. About one-third of adults infected with the hepatitis virus experience no further symptoms. The remaining two-thirds go on to develop jaundice, lasting from 1 to 3 weeks. Once the first sign of jaundice appears patients start to feel much better. Progression of the illness is much less common in children with only about one in twelve developing jaundice. Faeces from a patient with acute

infective hepatitis caused by hepatitis A are highly infectious. Hepatitis E virus, another RNA virus, is a very rare cause of hepatitis. Like hepatitis A, infection with hepatitis E is associated with exposure to contaminated water and it is most common in the Far East. It, too, causes a relatively mild self-limiting illness, except in pregnant women. The hepatitis E virus can cause a life-threatening infection in such cases.

The hepatitis A virus was at one time classified together with the enteroviruses. It is now considered to be sufficiently different from the enteroviruses to be classified separately. The enteroviruses are a large family of viruses that are spread by water and that cause illness with extra-intestinal symptoms. In tropical countries enterovirus infections are seen throughout the year but in countries with a temperate climate the incidence of enterovirus infection shows a marked seasonal variation with most cases occurring in summer and autumn; at a time when exposure to contaminated water is most likely. The most important of the enteroviruses is the poliomyelitis virus. As well as causing waterborne infections this virus may be transmitted by the respiratory route. Infected faeces are, however, a very significant source of poliomyelitis virus. In most people who are infected with this virus, it causes a mild sore throat and, perhaps, a headache. In about 5% of patients, these symptoms progress to meningitis. The symptoms of severe headache and intolerance of bright light can last from 2 to 10 days but the meningitis then spontaneously resolves. In a very few patients, probably about 0.1%, the infection causes paralytic poliomyelitis. This is an infection that affects the primary motor neurons, causing paralysis and muscle wasting. This damage is not a result of infection of the muscle cells but is secondary to the effects of virus infection of the motor neurones. If the respiratory muscles are infected then the victim is condemned to a life in an iron lung. An effective vaccination programme has almost eradicated poliomyelitis from the globe.

Two other important groups of enterovirus exist. The ECHO viruses cause meningitis. This

condition is not uncommon but is much less severe than bacterial meningitis and patients typically make a full recovery from enterovirus meningitis. The Coxsackie viruses infect muscles. Coxsackie B viruses have a predisposition for infecting heart muscle, causing progressive damage. Many heart transplants have been made necessary as a consequence of Coxsackie B infection. These viruses have been associated with a number of other extra-intestinal infections, including respiratory infections and conjunctivitis, although these are generally self-limiting.

Sewage is the main source of virus contamination of water. Raw sewage contains a very large number of viruses and even treated effluent may carry some viruses. Groundwater has only occasionally been found to be contaminated with human viruses. Treatment of potable water removes infectious viruses and the presence of viruses in tap water is an indication of the failure of water treatment processes. Lake and pond water may contain viruses. This depends, at least in part, on exposure to sewage. The occurrence of human viruses in marine and estuarine waters reflects the load of the rivers that feed these bodies of water. Viruses may be found in seawater in the absence of bacterial faecal indicators. If rivers have a large effluent input then the waters they feed will have a correspondingly high virus load.

Shellfish are an important potential source of human virus infection. Many shellfish that we consume are filter feeders. They extract their microscopic food from water by pushing vast quantities of water across their filter gills. There the food particles become trapped in mucus and the beating of cilia on the surface of gills propels food into the gut proper. If filter feeding shellfish live in water that is contaminated with human sewage, then they can very easily concentrate virus particles that are present in the water. In 1998, in a single incident, 61 people in Florida were infected with the hepatitis A virus after eating shellfish that had been exposed to human sewage at very low levels. Although the bacterial examination of shellfish is routine, there is currently

no reliable and reproducible method for assaying shellfish for the presence of human viruses.

4 VIRUSES USED AS FAECAL INDICATORS

As with faecal indicator bacteria, viruses chosen to demonstrate faecal contamination of water are chosen because of their resilient properties and their association with human infections. Faecal indicator viruses can survive for long periods in contaminated water. Two groups of viruses are routinely sought as faecal indicators, providing evidence that water is contaminated with human faeces and is thus unsafe. These are the **enteroviruses** and the **rotaviruses**. Although representatives of these two groups may be responsible for mild or even subclinical infection, the enteroviruses are associated with serious and even life-threatening infections and rotaviruses are the most common agents responsible for life-threatening diarrhoea in children under 2 years old worldwide. These infections can be spread by the faecal–oral route, with contaminated water a frequent vector of infection. Rotaviruses are responsible for infections that display the typical gastrointestinal symptoms of vomiting and diarrhoea. In contrast, enterovirus infections are often characterized by extra-intestinal symptoms. The enteroviruses and rotaviruses thus represent useful markers of human faecal contamination of water. Enteroviruses are, in general, much easier to culture than are rotaviruses. There are currently four groups of rotaviruses recognized by their serological reactions. There are over seventy serotypes of enterovirus.

If both enteroviruses and rotaviruses are found in a water sample, this is taken as evidence of gross faecal contamination. When natural waters are examined it is more likely that enteroviruses rather than rotaviruses will be isolated. This is, in part, a reflection of the relative ease with which each of the virus families can be recovered. It also indicates

the relative prevalence of these viruses in human faeces and also their differential survival properties.

5 TISSUE CULTURE TECHNIQUES USED IN THE EXAMINATION OF WATERBORNE VIRUSES

Since viruses are obligate intracellular parasites, they must be provided with living cells if they are to replicate. In the laboratory, viruses are often cultivated in tissue cultures. Cells can be removed from animal tissues and can be grown in one of a number of growth media at a suitable temperature and under appropriate atmospheric conditions. Cells in tissue cultures attach to the glass or plastic surface of their culture vessels and spread to form a cell monolayer. In many tissue cultures, cell contact within the monolayer causes an inhibition of further growth. Cells harvested from embryonic tissues tend to grow better than do cells from adult tissues. Once a monolayer is formed, its cells may be harvested and diluted to seed subsequent monolayers. This process is known as passaging. Tissue cultures can typically be maintained for a limited number of passages. Embryonic cells may be passaged up to 100 times. At high passage numbers, such cell lines gradually lose the ability to become infected with viruses. Cancer cells are particularly well suited to tissue culture. If the cells of tumours are maintained in tissue culture, they can be grown indefinitely. They are also not subject to the contact inhibition that causes the formation of monolayers. Once a single layer of tumour cells has developed, the culture will continue to accumulate in layers as cells pile up on top of one another.

Animal cells have exacting requirements for their growth media. In essence, however, the purpose of the growth medium is to provide an environment in which the tissue culture cells can flourish and divide. To do this they must be in an isotonic state where the osmotic pressure of the medium matches that within the cells. The pH must be maintained by a buffer system so that the acid balance suits the cells in culture. Tissue culture media include neutral

red as a pH indicator. This is why healthy tissue cultures have a pink colour, whereas infected tissue cultures turn bright yellow. It is important to maintain tissue cultures in the dark since the neutral red indicator becomes toxic to cells if exposed to the light.

It is necessary to provide cells in culture with all their nutritional requirements. The osmotic balance of culture media is provided by a solution of different inorganic salts, referred to as the basal salts solution. If tissue culture cells are to be rested, then they may be maintained in a basal salts medium, provided that it is appropriately buffered. Growth media must contain all the nutrients necessary for cell development. These include vitamins as well as glucose and amino acids. Growth media for tissue cultures include a serum component if cells are to be maintained through a number of passages. Sera may be derived from a number of sources. These include human serum, rabbit serum, calf serum and fetal calf serum. The serum component provides cells in tissue culture with numerous growth factors necessary for the long-term maintenance of the culture. Tissue cultures are typically incubated at 37°C and under an atmosphere where the concentration of carbon dioxide is elevated to between 5 and 10%. This is necessary to provide an environment that models that inside the mammalian body.

Tissue culture media meet the very exacting growth demands of animal cells. Microbes may be much less demanding in their growth requirements. Hence microbial contamination of tissue cultures is a significant problem. Consequently, great attention must be applied to aseptic technique when setting up and maintaining tissue cultures. The risk of microbial contamination may be further reduced by the addition of antibiotics to the tissue culture medium. The antibacterial antibiotics penicillin and streptomycin are routinely added to media used for tissue culture. Together, these two antibiotics have a very broad antibacterial spectrum. They also have a synergistic effect. This means that their combined activity is greater than the additive action of each drug. Care must be employed when using antimicrobial supplements in tissue

culture media. They may have an adverse effect on the growth of the cells in culture or they may be detrimental to the yield of virus from the supplemented culture. Although these antibiotics are antibacterial, it is necessary to use them at concentrations that are not toxic to the animal cells in culture. Typically 100 Units of penicillin G and 0.1 mg of streptomycin per millilitre of medium are used. Antifungal agents, such as nystatin at a concentration of 25–50 µg/ml, are also commonly employed in tissue cultures.

Viruses can be grown in a wide array of vessels, depending upon the requirements of particular protocols. If small volumes of cells are required for experiments that need many replicates, then microtitre trays are frequently the container of choice. These are plastic trays measuring approximately 10 cm by 12 cm and containing eight rows of twelve wells: 96 wells in total. Each well will hold slightly more than 300 µl. Alternatively, if large numbers of cells are required, for example to propagate a stock culture of virus, then a Winchester bottle can be used. These are large round bottles that have a 2.5 litre capacity. When such vessels are used, a relatively small volume of culture medium is placed in the seeded bottle. During incubation the bottle is slowly rotated so that its walls will be regularly bathed in the medium. Cells grow as a monolayer on the glass of such bottles.

Viruses can only infect a limited range of cells. This phenomenon is referred to as tropism. It is a consequence of the nature of the infection process. For a virus to infect and penetrate a cell it must first attach to the surface of its target host. To do this, specific receptors on the cell surface must interact with anti-receptors on the virus surface. Infection of the host will only occur if an appropriate interaction between receptor and anti-receptor has occurred, leading to a specific attachment and subsequent penetration of the target host cell. Tropism is a phenomenon that is associated with the attachment and penetration of viruses into host cells. If this step is experimentally by-passed, then cells that cannot normally act as hosts to particular viruses have been shown to support virus replication. For example,

poliomyelitis virus is only capable of infecting primate cells under normal circumstances. If, however, virus particles are introduced into mouse cells using microinjection techniques, then the virus will replicate as if it were in a primate cell.

In tissue culture, enteroviruses undergo a complete infectious cycle. Viruses infect cells in the culture, they replicate to form new virus particles and then the infected cell bursts to release new virus particles. The enteroviruses can thus easily be detected using a simple plaque assay. In contrast, the rotaviruses are not capable of undergoing a complete cycle of infection in tissue cultures. Rather, these fastidious viruses infect cells to produce viruses that lack an outer capsid. For this reason, cells in tissue culture that have been infected with rotaviruses are detected using an immunological assay to detect the presence of virus antigens associated with the VP6 antigen of the inner capsid.

Liquid culture can be used for the detection of enteroviruses. Cells in suspension are exposed to samples that may contain virus particles. Enteroviruses will infect cells to cause characteristic changes in the cell morphology. If a number of cultures are infected with aliquots of a sample then reference to

published tables can be used to determine the most probable number of virus particles present in the medium. More commonly, simple plaque assays are used to enumerate the viruses in water. This permits slower growing viruses to be detected as well as those that can grow easily and rapidly in tissue culture.

6 PREPARATION OF WATER SAMPLES FOR VIROLOGICAL EXAMINATION

The procedures described in the rest of this chapter relate to the examination of water samples as carried out in the UK (Fig. 2.3). Other methodologies are used elsewhere but no single method has been shown to be superior to other methods. The methods described below are technically demanding and are relatively expensive. Unlike bacteriological testing of water, the examination of water samples for viruses is not undertaken as a routine procedure on a large number of samples. Care must therefore be taken when interpreting the results of such assays.

Water samples should ideally be processed as soon as possible after sampling. It is recommended that samples should be processed within 24 hours of collection and after 36

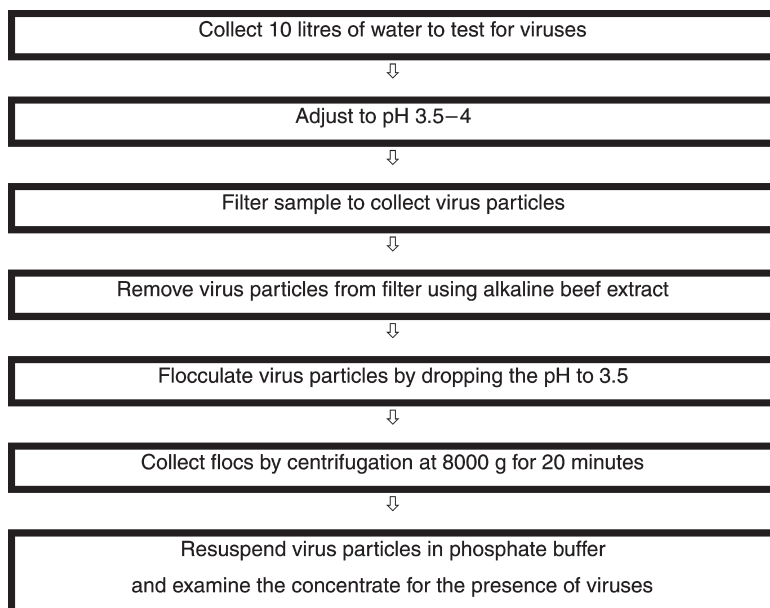


Fig. 2.3 Flow chart for the concentration of virus particles in water samples.

hours if a sample has not been processed it must be discarded as worthless. If they cannot be processed immediately, water samples should be kept cool and in the dark. Preferably they should be maintained at temperatures below 10°C. They should not, however, be frozen.

Samples of water for virological examination may contain very small numbers of virus particles. It is thus necessary to treat the samples to concentrate any viruses that may be present so that they may be detected more easily. If they cannot be tested immediately, sample concentrates may be stored frozen. Concentrates may be frozen at -20°C but if lower refrigeration temperatures are available so much the better. All samples must be properly labelled throughout the examination procedure. If the water to be tested has been previously chlorinated – for example mains water in the UK – then the residual chlorine must be inactivated by the addition of sodium thiosulphate to a final concentration of 18 mg/l.

Typically, 10 litre samples are collected and the pH of the sample is adjusted to lie between pH 3.5 and 4. This is achieved by adding 2 M hydrochloric acid to the sample and monitoring the pH with an accurate pH meter. The suggested method is to take a small subsample of known volume and calculate the volume of hydrochloric acid needed to lower its pH to 3.5. The volume of hydrochloric acid required to treat the whole sample may thus be calculated. At a pH value of 3.5, protein material, including virus particles, will adhere to nitrocellulose. The entire sample is then passed through a nitrocellulose filter with a 45 micron pore size to trap any virus particles that may be present in the sample.

To remove the trapped virus particles, the test filter is treated with 3% (w/v) alkaline beef extract solution (pH 9.5). Not all beef extract can form flocs in acid so it is essential to test each batch that may be used in this procedure for its ability to form flocs. A satisfactory batch will produce a visible precipitate when the pH is lowered to pH 3.5. The beef extract is sterilized by autoclaving prior to use and the pH is raised to pH 9.5 only after autoclaving.

For each 10 litre water sample, 300 ml of alkaline beef extract is reacted with the filter for 2 minutes. After agitation the virus particles become resuspended in the alkaline beef extract but they require further concentration before they may be easily detected. Skimmed milk made up as a 0.5% (w/v) solution in sterile water may be used in place of beef extract. The pH of the beef extract virus mixture is reduced to pH 3.5 by the stepwise addition of 2 M hydrochloric acid. Under these conditions, virus particles within the sample will flocculate and the flocs can be collected by centrifugation at 8000 g for 20 minutes. The supernatant beef extract is decanted from the floccular material containing the virus particles. These are resuspended in 10 ml of a 0.15 molar sodium phosphate buffer at pH 8. This procedure concentrates virus particles 1000-fold and the concentrate is then ready for use in subsequent virus assays. This may be used immediately. Alternatively the concentrate may be frozen and stored for examination in the future. Approximately one-third of the concentrate may be used to test for the presence of rotaviruses; the remainder is available to assay for enteroviruses.

Concentration of water samples using the method described above inevitably leads to the loss of virus particles from the sample. Loss of virus particles may occur at each step of the concentration process. When examining water for its virus content, each experimental run includes a control in which a sample of sterile water is seeded with a known number of virus particles. For the examination of enteroviruses, control samples are typically seeded with 10^6 particles of an attenuated strain of poliomyelitis virus. Attenuated viruses are capable of normal growth in tissue culture but they are incapable of causing clinical disease in humans. The Sabin poliomyelitis vaccine, for example, exploits a live, attenuated virus to confer immunity. The control sample is put through the identical concentration procedure to the test samples. The number of particles recovered from the control sample is then compared with the number that was used in the seeding. From this

comparison the percentage recovery of virus particles can be calculated:

$$\frac{\text{Virus particles recovered}}{\text{Virus particles input}} \times 100\%$$

When using the concentration procedure described above the recovery of virus particles is typically between 60 and 70% (Fig. 2.4). When reporting data from the virological examination of water it is often the practice to report the raw data together with the results from the control experiment. This is so that the person commissioning the test can calculate the actual number of viruses in the sample. Alternatively, the tester may calculate the projected number of particles in the original sample.

Whichever approach is used it is essential to make clear in the final report whether the count is adjusted or not.

Water samples typically require concentration if viruses are to be recovered, since virus particles are not usually present in water in large numbers. For sludge and samples that contain solid particulate material it is important to dilute samples so that there is no more than 4% (w/v) solid material in the sample. Aluminium chloride, added to a final concentration of 0.5 mM, may be added to aid the flocculation of virus particles in the sample.

Enteroviruses and rotaviruses are almost universally present in sewage. These are both very common virus families and their numbers in raw sewage can be very high. The virological

A 10 litre sample of sterile water is seeded with 10^6 particles of poliomyelitis virus. The control is then subjected to the same concentration protocol as test water from a reservoir suspected of being contaminated with human faeces. A serial 100-fold dilution of the resultant concentrate was made and these dilutions were used in a plaque assay for enteroviruses (poliomyelitis virus is an enterovirus). The following numbers of plaques were observed:

Dilution	Plaque forming units (pfu) per plate
100	Too many to count
10^{-2}	Too many to count
10^{-4}	69

The number of particles recovered is thus the number of plaques seen multiplied by the dilution factor:

$$\begin{aligned} &69 \times 10^4 \\ &= 6.9 \times 10^5 \end{aligned}$$

As 10^6 particles were used to seed the control the percentage recovery in this experiment is:

$$\frac{6.9 \times 10^5}{10^6} \times 100\%$$

$$= 69\%$$

Fig. 2.4 Sample calculation for the recovery of virus particles from a control sample of water.

examination of sewage is not often undertaken and is probably only important when studying the progress of vaccination programmes, e.g. to monitor the eradication of poliomyelitis virus or if studying the efficacy of novel treatment programmes. Once a sewage sample has been homogenized it is not necessary to subject it to a concentration procedure to isolate viruses of faecal origin. Rather, it is more common practice to use a dilution procedure to detect faecal viruses in samples of sewage. Bacteria and other microbes in the sewage will flourish when inoculated onto a tissue culture. These will outgrow the animal cells and will thus destroy the tissue culture. If microbial growth is not contained, viruses will be unrecoverable from sewage. Dilutions of sewage material are typically made in phosphate buffered saline and the diluted samples are treated with antimicrobial agents. These generally include a β -lactam antibiotic, such as one of the penicillins, together with an aminoglycoside, such as streptomycin. These will kill bacteria in the samples. An antifungal agent is also used to treat the diluted sewage sample before it is used to inoculate a suitable tissue culture.

7 EXAMINATION OF WATER FOR THE PRESENCE OF ENTEROVIRUSES

Enteroviruses are detected in water samples using a tissue culture plaque assay. Primate cells are used as hosts for these viruses. BGM cells or human diploid fibroblasts are the cell lines most commonly used in the laboratory culture of enteroviruses. Other cell lines may be used at the preference of the local virologist. These include PMK, Vero, FL HeLa and Hep-2 cells. Using more than one cell line can improve the detection of human viruses in water concentrates. Tissue cultures are initiated in 75 cm² tissue culture flasks and cells are passaged every 5 days. After five passages, the tissue cultures are transferred from stationary flasks to roller bottle cultures. In the examination of water samples for enteroviruses BGM cells are only used between passages 75–120. During this time they have maximal susceptibility to enterovirus infection. With all virus

assays, it is important to include both positive and negative controls in the experimental protocol.

Cells from a roller bottle culture are harvested by replacing the normal culture medium with a medium containing 0.05 mg/ml trypsin and incubating for one hour. This loosens the cells from the glass and makes them more susceptible to virus infection. Once the cells have been harvested, the culture yield is determined by counting the cells. This permits a calculation of the volume of cells necessary to initiate a plaque assay. The cell count is undertaken using a counting chamber such as the Improved Neubauer counting chamber shown in Fig. 2.5.

Plaque assays are conventionally carried out in Petri dishes. For enterovirus assays of water samples, 90 mm Petri dishes are used and each dish is seeded with approximately 3×10^7 cells mixed with 15 ml of growth medium. A sample calculation to determine the number of cells in a tissue culture and to determine the volume of cells is shown in Fig. 2.6. This calculation assumes that an Improved Neubauer Counting Chamber is used to determine the cell number. This counting chamber was originally developed for haematology laboratories. Other counting chambers have been developed for different purposes. They all operate on the same principle: determination of the number of particles in a fixed volume. The calculations will, however, require substitution of different factors, depending upon the fixed volume in which particles are counted.

In a plaque assay, a cell monolayer is infected with virus and the culture is overlaid with a semisolid medium to prevent the lateral spread of virus particles. The virus particles within the sample to be tested infect cells within a cell monolayer. Infected cells then produce new virus particles and lyse to release their progeny. These newly released virus particles then infect adjacent cells that subsequently lyse to release even more virus particles. These cannot spread through the tissue culture since they are restrained by a semisolid overlay or by agar within the growth medium, depending upon the restraining method chosen. Consequently, discrete holes appear within the cell

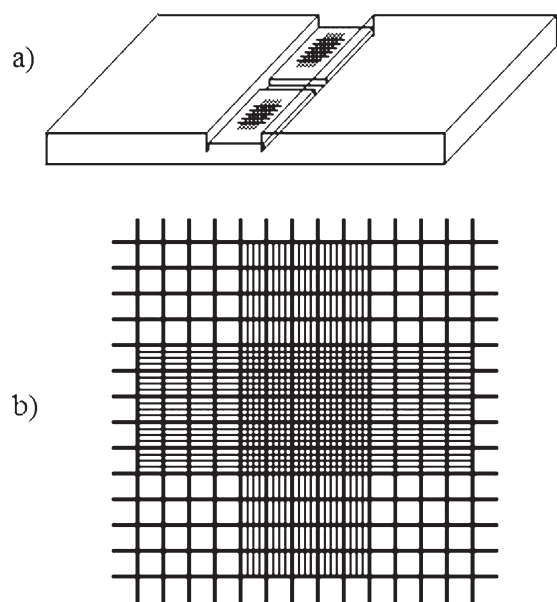


Fig. 2.5 The Improved Neubauer Counting Chamber. This counting chamber (a) is slightly larger than a conventional microscope slide and has indentations for two specimens. In the centre of each chamber is an accurately etched grid (b). The total area of the 25 larger squares in the centre of the grid is 1 mm^2 . The sample to be counted is introduced into the chamber and a coverslip is placed over the sample. When placed correctly, the contact between the clean glass of the chamber and the coverslip causes the formation of 'Newton's rings': a rainbow effect between the coverslip and the glass on either side of the counting chamber. If Newton's rings are seen, the height of the coverslip from the grid is 0.1 mm . The volume of fluid trapped over the central squares of the grid is thus 0.1 mm^3 . This is equivalent to 10^{-4} ml . By counting the number of objects within the 25 central squares and multiplying by 10^4 , the number of objects per ml can be obtained after taking into account any dilution factor.

monolayer, each originating from one infectious virus unit. These holes are known as plaques and the infectious virus units are referred to as plaque-forming units.

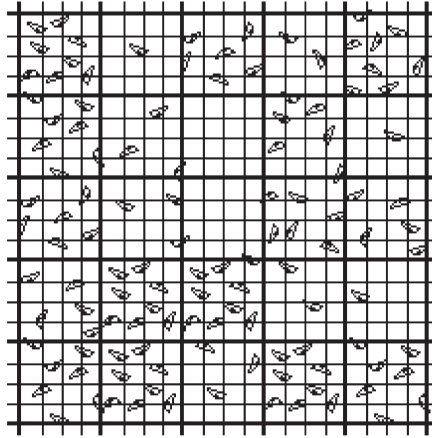
When assessing the number of enterovirus particles in a concentrated water sample, in each assay dish 3×10^7 cells are mixed with 15 ml of growth medium containing 2.4% agar that has been previously cooled to 45°C . The mixture is then used to seed a sterile 90 mm tissue culture quality triple-vented Petri

dish. The purpose of the agar is to prevent the spread of newly released virus particles. This causes plaques to develop as discrete entities. A total of 6.5 ml of the concentrated water sample is split between three assay plates. The contents of each of the Petri dishes are then carefully mixed and the tissue cultures are incubated at 37°C in an atmosphere of 5% carbon dioxide in air for 48 hours. The counts resulting from each of the three assays should be of the same order. If one of the three tests differs significantly from the others, then the validity of that assay must be questioned and the sampling must be repeated. This acts as an internal control for the assay procedure. By counting the number of plaques present, the number of virus particles present in the original sample can be calculated and the results presented. A sample calculation and reports are given in Fig. 2.7.

It is necessary to confirm that the plaques that are seen are due to virus activity rather than to a toxic reaction within the tissue culture. Plaques are subcultured to ensure the presence of live virus. Normally, no attempt is made to differentiate viruses isolated from plaque assays. No rapid, inexpensive method is available to identify enteroviruses to serovar level and the presence of any enterovirus is evidence of the faecal contamination of the original sample. It may, however, be desirable to determine which viruses are present in a given sample. For example, if a particular body of water is epidemiologically linked with a particular virus infection, then it may be beneficial to identify the virus isolated from the sample to subfamily level. There are a number of techniques available to differentiate enteroviruses.

A variety of serological methods of identifying enteroviruses may be used but, as there are over 70 distinct serovars of enterovirus, this method requires over 70 antisera to identify fully the isolate in question. In practice, isolates can be identified to group level with a much smaller panel of antisera. Serological techniques used in enterovirus identification include indirect immunofluorescence; enzyme linked immunosorbent assays (ELISA) and neutralization tests. ELISA tests work on a similar

The picture below is a representation of the central squares of an Improved Neubauer Counting chamber with cells from a tissue culture to be used to seed an enterovirus plaque assay. This was derived from a 1:50 dilution of the original cell harvest



The cell counts in each square may be tabulated thus:

8	1	6	2	8
7	3	0	6	1
6	3	0	6	5
5	8	9	2	1
7	8	3	8	7

There are a total of 120 tissue culture cells within the grid. In the Improved Neubauer Counting Chamber, this grid entraps a volume of 10^{-4} ml. Hence the sample as counted represents a cell count of 120×10^4 cells/ml. The sample counted, however, was a 1:50 dilution of the original cell suspension.

Hence, the number of cells in the original suspension would have been $50 \times 120 \times 10^4$ cells/ml. This is equivalent to a count of 6000×10^4 cells/ml. This is more properly written as 6×10^7 cells/ml.

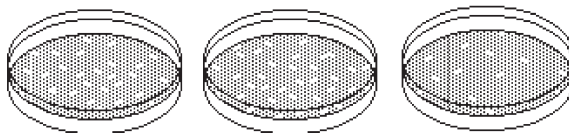
As 3×10^7 cells are required to seed the enterovirus plaque assay, each 90 mm Petri dish requires 0.5 ml of the original cell suspension.

$$\frac{6 \times 10^7}{3 \times 10^7} = \frac{1}{2}$$

Fig. 2.6 Sample calculation to decide how many cells to seed for an enterovirus plaque assay.

A party had enjoyed an outward-bound weekend at the Chaldicotes Centre on the River Ouse. This involved participation in water sports, including the practise of a capsized drill. Inevitably, most of the group ingested some of the river water and subsequently a number of the party became unwell. The main symptoms of the illness were a fever and gripping chest pains. One of the party was a microbiologist who suspected that they were suffering from a Coxsackie B virus infection. She considered faecal contamination of the River water to be the common epidemiological link. A virological examination was commissioned and enteroviruses were, indeed, recovered from the water sample examined.

The sample of river water was concentrated and 6.5 ml of concentrate was used in a plaque assay for enteroviruses. A control sample was examined in parallel, as described. This control established that 69% of enterovirus particles were recovered after the concentration procedure. The 6.5 ml of concentrated water sample taken from the River Ouse was divided between three plaque assay plates. The first plate yielded 27 plaques, the second 35 and the third plate 13 plaques. The total count was thus 75 plaque forming units.



As 6.5 ml from a total of 10 ml of concentrate was used, there would be

$$\frac{75}{6.5} \times 10$$

= 115 virus particles in the whole of the concentrated sample. Since there was a 69% recovery of the control virus, the report may be presented as follows:

There was 115 plaque forming units of enterovirus per 10 litres isolated from the water sample at Chaldicotes on the River Ouse.

Recovery of enteroviruses in a control experiment was 69%.

This is indicative of a total enterovirus load of 167 pfu/ 10 litre, allowing for loss of virus particles during processing.

Fig. 2.7 Sample calculation and specimen report having determined the number of enteroviruses in a water sample.

principle to immunofluorescence but use antibodies tagged with an enzyme rather than a fluorescent dye. The enzyme catalyses a reaction that yields a coloured product. The intensity of the colour produced indicates the strength of the ELISA reaction. In neutralization assays, virus particles are reacted with an array of antisera before being exposed to tissue culture. The antiserum that prevents plaque formation is specific for the virus being investigated. There are currently no standard typing systems based upon the analysis of the virus genome. Enteroviruses readily undergo

mutation and the instability of the genome makes nucleic acid-based typing too unreliable for the unambiguous identification of an isolate.

8 EXAMINATION OF WATER FOR THE PRESENCE OF ROTAVIRUSES

In contrast to the plaque assay used to enumerate enteroviruses, when rotaviruses are detected, a cell monolayer is infected and virus particles are allowed to grow within

the tissue culture and the cells are subsequently fixed and subjected to indirect immunofluorescence. LIC-MK2 or MA-104B cells are most often used for the culture of rotaviruses. Rabbit serum containing antibodies that react specifically with rotaviruses is then added to the tissue culture. If there are virus particles present within the fixed cells, these will react with the antibodies in the rabbit serum and will trap the antibodies within the fixed cells. The culture is thoroughly washed to remove any unbound antibody and is then flooded with fluorescently labelled anti-rabbit antiserum. The tagged antibodies in this antiserum will react with the rabbit antibodies that have attached to the rotavirus particles. When viewed under ultraviolet light, infected cells will be able to fluoresce and appear brightly coloured. This is because the rotavirus particles will trap the rabbit antibodies that in turn will trap the fluorescently labelled anti-rabbit antibodies. Uninfected cells are unable to fluoresce, as they have no means of trapping the rabbit antibodies (Fig. 2.8). This two-step indirect immunofluorescence technique is used rather than using direct fluorescent microscopy where tagged anti-rotavirus antibodies are used because it permits amplification of the signal. With direct fluorescence each virus particle is quickly saturated with antibody. When an indirect immunofluorescence is used, each virus particle will have numerous anti-rotavirus antibodies attached to it. In turn, each of these antibodies will provide binding sites for the anti-rabbit antibodies.

Human rotaviruses are more difficult to culture than are enteroviruses and they do not undergo a full infectious cycle in tissue culture. Trypsin treatment of the cells prior to infection is essential for the efficient propagation of rotaviruses in tissue culture. At one time, rotaviruses were grown in large-scale tissue cultures but now microtitre plates are used and small volumes of material may be examined.

Rotavirus assays are carried out in microtitre trays. These trays have eight rows of twelve wells each: a total of 96 wells per plate. Each well holds just more than 300 μl . A total of 1.5×10^5 cells in 100 μl of growth medium are

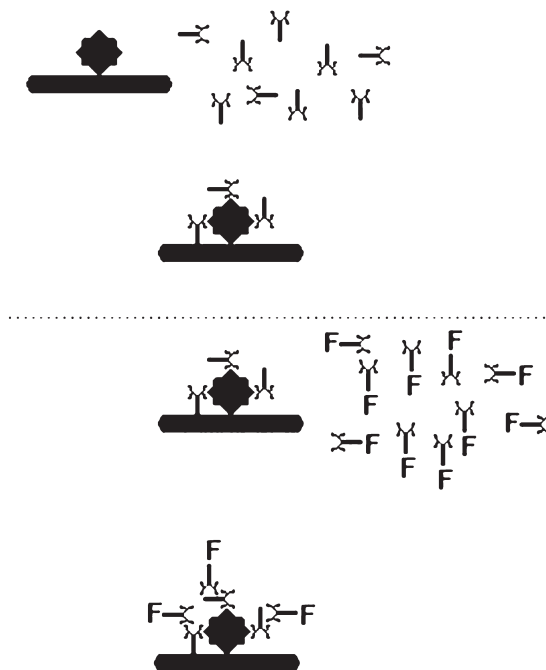


Fig. 2.8 Indirect immunofluorescence used to detect rotavirus particles. The rotaviruses within infected cells provide binding sites for anti-rotavirus antibodies within rabbit antiserum. These are allowed to react and the preparation is thoroughly washed to remove any unbound antibody. The preparation is then reacted with fluorescently tagged anti-rabbit antibodies. These bind to the antibodies that are already attached to the virus particles, thus amplifying the signal and making virus-infected cells easier to see when viewed in a fluorescence microscope.

added to each of the wells of a microtitre tray. The cells are incubated for one hour at 37°C under 5–10% carbon dioxide. This allows the cells to attach to the wells of the microtitre plate. After this, the growth medium is decanted off and is replaced with 100 μl of cell maintenance medium with 0.05 mg/ml of trypsin added. The trypsin is added to make the cells in the culture susceptible to rotavirus infection. After incubation at 37°C in elevated carbon dioxide for a further hour, thirty-two 100 μl aliquots of the concentrated water sample are added to individual wells in the microtitre tray. The water samples and trypsin-containing medium are incubated with the cells for another hour. The liquid in each

well is replaced with 100 μ l of maintenance medium and the microtitre tray is incubated for another hour to rest the infected cells. Finally, the maintenance medium is replaced with 100 μ l of growth medium and the cells are incubated overnight.

On the following day growth medium is removed from the tissue cultures and the cells are fixed. This is achieved by washing the cells thoroughly with three changes of phosphate buffered saline then treating with ice-cold methanol or with absolute ethanol for at least 10 minutes. After ethanol fixation the cells are rehydrated with three washes of 250 μ l of phosphate buffered saline. The cells are then ready for virus detection using an indirect immunofluorescence technique.

Fifty microlitres of a 1:500 dilution of anti-rotavirus antiserum raised in rabbits is then added to each well in the microtitre tray. This allows anti-rotavirus antibodies in the antiserum to attach to virus particles within the fixed tissue culture cells. After incubation at 37°C for one hour the rabbit antiserum is removed from each well and the contents of the wells are washed with three washes of phosphate buffered saline. Fifty microlitres of fluorescently tagged anti-rabbit antiserum is added and the tray is incubated for a further hour. The antibodies in this antiserum will attach to the rabbit antibodies that are, in turn, attached to the virus particles. Using this indirect immunofluorescence technique the signal from each virus particle is amplified and the particles can be detected by fluorescence microscopy. The wells are washed with a further three changes of phosphate buffered saline before microscopic visualization. Plates may be stored at 4°C in the dark for up to 6 months without apparent loss of signal.

Rotaviruses only replicate in the cytoplasm of infected cells. Cells with fluorescent nuclei are thus not counted in this assay. Fluorescent cells with a fluorescent cytoplasm are assumed to be infected with rotavirus. As with the enterovirus assay a control sample is seeded with a known concentration of rotavirus so that the percentage loss during the concentration procedure can be calculated. The virus counts are reported as the number of fluorescent foci

per 10 litres of water. It is again important that the report makes clear whether or not allowance has been made for the loss of particles during concentration of the water sample.

9 THE POLYMERASE CHAIN REACTION (PCR) AND DETECTION OF VIRUS PARTICLES IN WATER SAMPLES

There have been many advances in the use of molecular biological techniques in recent years. One of the most significant has been the development of the polymerase chain reaction, known as PCR for short. DNA is the molecule in which almost all genetic information is stored. Four different 'bases' code for genetic information. Groups of three bases on the coding strand of DNA specify which amino acids will become incorporated into proteins. These triplets also provide the punctuation of the code, indicating where protein translation should start and stop. DNA molecules have two strands. This enables it to be accurately replicated as each base can only form a stable pair with one other base: adenine pairs with thymine, guanine with cytosine.

DNA replication is carried out by enzymes known as DNA polymerases and is a chain reaction. Each of the strands of a DNA molecule acts as a template for replication. Thus one molecule of two strands is replicated to produce two molecules with a total of four strands. These are both replicated to give rise to four molecules with a total of eight strands, and so on. DNA replication is initiated when the two DNA strands become separated. A very short sequence of DNA known as an oligonucleotide primer, or just primer for short, then anneals to the strand that is to be replicated. Primers have a sequence that is complementary to their target strand. If the target has adenine, the primer will have thymine, and so on. The primer serves as the starting point for DNA replication and nucleotides are added to the new strand by the action of a DNA polymerase enzyme. DNA replication can take place *in vitro* when single stranded DNA

is mixed with appropriate primers in a buffer containing all four nucleotides and magnesium ions together with DNA polymerase.

DNA molecules can easily be separated into their component strands by heating. At temperatures of about 95°C the bonds that maintain the base pairs that hold the two DNA strands together are broken and the strands separate. The strands may re-anneal if the temperature is dropped. The precise temperature of re-annealing depends upon the nucleotides present in the DNA strands. At about 60°C, however, melted strands of DNA can re-anneal to regenerate the familiar double helix structure.

In PCR amplifications, the template DNA is melted in a mixture that contains two primers: one for each DNA strand, and designed to anneal at either end of a target sequence. Primers are generally between 10 and 20 nucleotides long. This is sufficiently long to overcome the problem of primers binding at random to sequences other than the target DNA at the appropriate annealing temperature. This gives PCR a high degree of specificity. Lowering the annealing temperature sufficiently may permit non-specific binding and this may be exploited, for example in PCR typing techniques. This makes PCR a very versatile tool as well as potentially a very specific technique. The temperature of the reaction is lowered to allow the primers to anneal to the target DNA strands. DNA polymerase from *Escherichia coli* was used in the earliest PCR amplifications. DNA replication was thus carried out at 37°C. This was unsatisfactory as for each amplification step new enzyme had to be added to the reaction. This allowed the opportunity for contamination with DNA that could act as a target for amplification, giving rise to false positive results. It was also expensive in its use of DNA polymerase. A typical PCR amplification has between 20 and 30 rounds of replication. The technique was revolutionized when a thermostable DNA polymerase, isolated from a hot water bacterium, *Thermus aquaticus*, was introduced. This enzyme is stable at 95°C and replicated DNA at 72°C. Currently, PCR amplifications involve 20–30 cycles where

DNA is melted at 95°C, primers are annealed at an appropriate temperature, typically about 60°C, and DNA is replicated by the *Taq* polymerase at 72°C. This leads to the production of a DNA amplicon of a predicted size, dictated by the positions of the primer pair. The first three cycles of the reaction are illustrated in Fig. 2.9.

PCR amplicons are DNA molecules and can be subjected to the same technologies as DNA isolated from organisms. They are most often detected by agarose gel electrophoresis. DNA molecules are charged and so will migrate when they are placed in an electric field. DNA samples can be loaded into wells in an agarose gel across which an electrical potential difference is applied. The DNA molecules will move out of the well and into the gel. The smaller the DNA molecule, the easier it will pass through the gel. In this way DNA molecules are separated according to size with the smallest molecules migrating the furthest. Having carried out an electrophoretic separation, the DNA may be visualized by soaking the gel in a very dilute solution of ethidium bromide. This dye can intercalate between the base pairs of a DNA molecule, causing it to fluoresce bright orange when illuminated by ultraviolet light. DNA amplicons can be identified and the distance they run can be compared to size standards. More sophisticated techniques may occasionally be applied to verify the nature of an amplicon. DNA carries restriction endonuclease recognition sites. These are points where restriction endonucleases cut DNA into fragments of predictable sizes. Restriction digest analysis can be used to confirm the structure of an amplicon if its nucleotide sequence is known. If the sequence of an amplicon is not known then the amplicon itself may be subjected to nucleotide sequence determination.

As well as amplifying DNA, the polymerase chain reaction may be adapted to detect the presence of RNA in a sample. This is very useful when trying to detect viruses with an RNA genome. In such cases, nucleic acid is extracted from the sample being examined and the RNA is acted upon by reverse

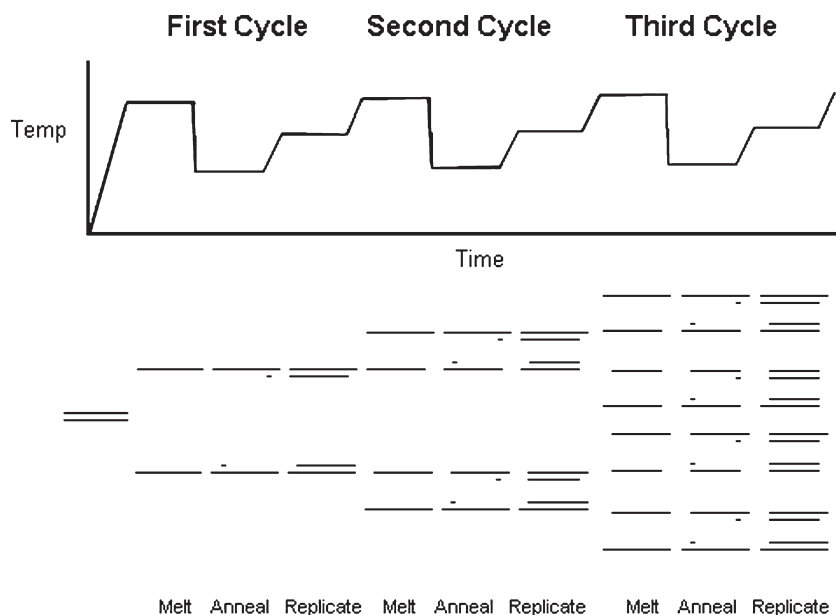


Fig. 2.9 The first three cycles of the polymerase chain reaction. The sample is heated to allow the target DNA to melt into its two separate strands. The reaction temperature is then dropped and primers bind to either end of target sequence, one primer on each strand of DNA. The reaction temperature is raised once more to permit DNA replication mediated by the thermostable *Taq* polymerase. By the end of the third cycle the reaction is dominated by DNA molecules of a size specified by the limits of the primer binding sites. Hence, when the PCR amplification is complete, the product overwhelmingly contains DNA molecules of this defined size.

transcriptase. This enzyme makes a DNA copy of an RNA template. The cDNA produced in this reaction can then subsequently be used as a template in a conventional PCR amplification. The polymerase chain reaction can also be used to amplify DNA from microorganisms that cannot be cultivated. In such cases, providing a unique protein can be characterized and a partial amino acid sequence can be determined, then a set of 'degenerate' primers can be constructed. More than one triplet can code for a single amino acid. Hence the degenerate primers must be designed to anneal with every possible nucleotide sequence that could encode the amino acids in the target protein. Having used degenerate primers to yield an amplicon, the nucleotide sequence of the amplified DNA can be determined in a standard sequence analysis. Specific primers may then be designed to amplify DNA from an organism that cannot be cultivated. The polymerase chain reaction is thus a highly versatile and sensitive technique for the detec-

tion of DNA, and, by implication, specific organisms. In theory it is capable of detecting single molecules and in practice it has been found reliable to detect microbes present in very small numbers within a sample. The only problem with using PCR technology to detect viruses is that it detects the total number of virus particles in a sample. It gives no indication of the proportion of virus particles that are potentially infectious. It may thus seriously overestimate the risk of exposure to a water sample.

10 THE USE OF BACTERIOPHAGE IN WATER TESTING

The examination of water for the presence of viruses need not necessarily imply a threat to human health. Humans may harness viruses in the examination of water. Every major group of organisms is susceptible to infection by viruses. This includes bacteria and the viruses that can

infect bacteria are known as bacteriophage. These viruses have a very limited host range. Indeed, the susceptibility to infection by a panel of bacteriophage is a very useful method of typing some important bacteria to subspecies level. This specificity of host range can be exploited in the examination of water.

Serratia marcescens is a Gram-negative bacterium, strains of which are susceptible to bacteriophage MS2. If this virus is mixed with a suspension of bacteriophage-sensitive bacteria that are then plated as a lawn onto nutrient agar, the bacteriophage causes the appearance of plaques in the bacterial lawn after overnight incubation of the culture. The bacterial lawn is punctuated with holes, each representing an infectious bacteriophage particle. These plaques are analogous to the plaques seen in tissue culture monolayers such as are counted in the enterovirus assay described in Section 7. This makes for the easy detection and enumeration of bacteriophage in samples.

Large quantities of bacteriophage can be cultivated by inoculating a broth culture of bacteria that are growing in late exponential phase. Bacteriophage will infect bacterial cells and thus they become replicated. The infected cells lyse to release new bacteriophage particles that can go on to infect other bacteria. The cloudy bacterial culture will become clear as all the bacterial cells become infected and are subsequently lysed to release large numbers of bacteriophage. The culture is centrifuged at low speed to remove cell debris and then at high speed to harvest virus particles.

Bacteriophage MS2 has been used to study the flow of water and other fluids. For example, silage production involves the fermentation of grasses to provide feed for cattle. This fermentation is a bacterial process and it produces acid. This can cause corrosion of the silo and consequently leakage of silage material and its bacteria. This may lead to serious

contamination of watercourses. To establish whether leakage of silo contents has occurred and if this has subsequently caused contamination of a particular watercourse, the silo in question can be seeded with MS2 bacteriophage particles. Water samples are then taken at regular intervals and mixed with *Serratia marcescens*—the indicator bacterium. These are then plated onto nutrient agar and incubated so that plaques may subsequently be detected. The appearance of plaques is evidence of bacteriophage in the water sample. Their occurrence after but not before seeding of the silo establishes that the silo is contaminating the watercourse. By taking samples at regular intervals not only can the source of contamination be confirmed, but also the time it takes for the silage material to reach the water may be established. The MS2 bacteriophage marker can also be used, for example, to track the source of water through subterranean courses. It is much more satisfactory than using tracking dyes – the old method for tracking water.

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Bacteria

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1 INTRODUCTION

Bacteria are indeed the lowest, or at least the simplest, form of life. Their prokaryotic cell structure is significantly different from higher forms such as the single celled algae and protozoa, invertebrates, plants and animals. Only the *Archaea* have a cell structure of similar simplicity. Bacteria are the most numerous of living organisms on earth in terms of number of species, number of organisms, and total mass of organisms. Each cell is small, typically 1–2 μm in diameter and length, and has a total mass of 1–10 pg. One thousand bacteria occupy approximately 10^{-12} ml of volume. Because bacteria are most commonly found in aqueous environments, their size is particularly important. A concentration of 1000 organisms per ml is very significant in terms of protecting health, but is invisible to the naked eye. Bacterial growth or replication is in numbers and concentration rather than mass of a single organism. Under optimal conditions of temperature, pH and nutrient availability, some bacterial species have generation times of less than 30 minutes. Such short generation times account for the rapid progression of infectious diseases.

Bacteria cause many human, animal, and plant diseases. Pathogenic bacteria, those that cause disease, are transmitted through direct contact with an infected host, by ingestion of contaminated food or water, or by the action of an intermediate host or disease vector. An example of a disease vector is a type of leafhopper, the glassy wing sharpshooter, which carries the bacterium *Xylella fastidiosa*, the cause of Pierce's disease in grapes and

other woody perennials. Most pathogenic bacteria are relatively host specific and pathogens of non-human animal and plant pathogens generally do not cause diseases in humans. However, human pathogens are often carried by asymptomatic non-human hosts. Such pathogens are termed *zoonoses* and include the organisms causing the most important waterborne diseases. Bacteria of the genus *Salmonella*, which are normal habitants of birds, reptiles and mammals, but are thought to be uniformly human pathogens, are examples of zoonoses.

In this chapter, the focus is on pathogenic bacteria transmitted through water or associated with water in some manner. All waterborne or water-washed diseases are also associated with food and food preparation. In some cases the pathogenic bacteria are normal members of the intestinal flora of food animals and contamination results during processing. Infection results from improper handling during preparation or from inadequate cooking. In other cases organisms may be transferred from a processor to the food directly or by washing the food with contaminated water. In most cases proper cooking and washing of preparation surfaces with soap and hot water will prevent infection. With the exception of cholera and legionellosis, most of the diseases discussed here are more commonly transmitted through food, as indicated in Table 3.1; note that the period 1976–1990 was a period in which water supplies in Israel were greatly improved with respect to public health protection but that food protection lagged considerably.

TABLE 3.1 Incidence of waterborne salmonellosis, typhoid fever and shigellosis in Israel from 1976 to 1997 (Source: Tulchisky *et al.*, 2000)

Disease	76–80	81–85	86–90	91–95	96–97
<i>Salmonellosis</i>					
waterborne	979	157	244	260	0
total	10 101	12 386	17 127	28 986	11 481
<i>Typhoid fever</i>					
waterborne	112	76	0	0	0
total	596	629	216	0	0
<i>Shigellosis</i>					
waterborne	6557	10 180	1524	260	0
total	32 839	44 152	29 070	25 874	7274

2 INFECTIOUS DOSE

Infection normally occurs only when a sufficient number of pathogens has been ingested. The number of bacteria required to produce an infection ranges greatly. Based on infections in volunteers, the required dose for *Shigella* is approximately 10 organisms while *Campylobacter* infections require several hundred organisms. Cholera infection requires approximately 10^6 organisms under normal conditions. However, if antacids are taken to raise the stomach pH the infectious dose drops as low as 10^4 (Madigan *et al.*, 2000). High infectious dose requirements make transmission through water more difficult because pathogen concentrations are generally low in natural waters, even when contaminated with human wastes, and because natural waters are, with few exceptions, not good growth mediums for human pathogens.

3 MORBIDITY AND MORTALITY STATISTICS

Morbidity is the incidence of disease, usually stated as cases per 10 000 population per year. Mortality is the rate of deaths from the disease, also commonly stated in units of 100 000 population per year. For most waterborne diseases the morbidity statistics are probably understated. Many cases of all of the diseases are not reported because the symptoms are relatively minor, because the ill person does not see a physician, or because of poor reporting

practices within the medical community. Public health agencies in many poor countries have difficulty in maintaining databases and even more difficulty in carrying out investigations of the causes of infection and death. When epidemics occur, the World Health Organization (WHO) and other agencies provide support that often results in improved investigation and record keeping. Thus data for large epidemics are probably more reliable than data for endemic problems.

4 PATHOGENIC BACTERIA OF CONCERN

Several million bacterial species are estimated to exist, with only a few thousand having been identified and to some extent characterized. At present, bacteria are classified by their 16S ribosomal RNA characteristics into 14 kingdoms, each of which is made up of many genera, which in turn include many species (Madigan *et al.*, 2000). Human pathogens are scattered among several of the kingdoms. New pathogens appear from time to time and it is not completely clear whether the organisms are new strains or if their pathogenic characteristics had not been observed previously. However, most of the principal bacterial pathogens have been recognized for over 50 years and these organisms account for nearly all of the serious bacterial infections that have been observed.

The principal waterborne diseases and the associated pathogenic bacteria are listed in Table 3.2. The first five diseases listed are all similar, except for severity, in that diarrhoea is the principal symptom. For all diarrhoeal diseases dehydration is a major complication. Even without treatment, keeping patients hydrated results in a high rate of recovery. The sixth disease, haemolytic uraemic syndrome, is also diarrhoeal but a toxin produced by the pathogen attacks the intestinal lining, destroys red blood cells and often results in kidney failure. Legionnaire's disease (also called legionellosis), tularaemia, and bacterially caused peptic ulcer are not normally classed as waterborne diseases. However, water

TABLE 3.2 Principal waterborne diseases and the associated pathogenic bacteria

Disease	Bacteria	Infectious dose	Characteristics
Cholera	<i>Vibrio cholerae</i>	10 ⁶	Severe diarrhoea, dehydration
Salmonellosis	<i>Salmonella</i> spp.		Watery diarrhoea often with abdominal cramping, nausea, vomiting fever and chills
Typhoid fever	<i>Salmonella typhi</i>	< 1000	Fatigue, headache, abdominal pain, elevated temperature. Approximately 4% death rate
Gastroenteritis or campylobacteriosis	<i>Campylobacter jejuni</i>	< 500	Watery diarrhoea often with abdominal cramping, nausea, vomiting, fever and chills
Dysentery or shigellosis	<i>Shigella</i> spp.	≈ 10	Bloody diarrhoea, abdominal cramps, rectal pain. Most virulent species, <i>S. dysenteriae</i> , produces toxin causing haemo uraemic syndrome
Haemorrhagic colitis and haemolytic uraemic syndrome	<i>Escherichia coli</i> O157:H7	Unknown ^a	Severe, systemic condition that occurs principally in children under 10 years of age
Leptospirosis	<i>Leptospira</i> spp.	Unknown	Fever, kidney infection, may result in kidney failure. In some cases there is internal bleeding, including pulmonary haemorrhage
Legionellosis	<i>Legionella pneumophila</i>	Unknown	Acute pneumonia, high fever, headache, cough, little sputum
Peptic ulcer and gastric cancer	<i>Helicobacter pylori</i>	Unknown	Sore or hole in the lining of the stomach or duodenum. Cancer

^a Probably similar to *Shigella*.

is or may be a mode of transmission in each case and all three diseases are widespread throughout the world. Hence they will be discussed here.

4.1 Cholera

Cholera, caused by *Vibrio cholerae*, a Gram-negative, curved and flagellated rod, is perhaps the clearest example of a waterborne disease. Although the disease can be contracted in other ways, by far the most common mode of transmission is ingestion of contaminated water. The infectious dose is very large, at least 10⁶ organisms, under normal circumstances due to the sensitivity of *V. cholerae* to low pH environments. Increasing the stomach pH with antacids lowers the infectious dose to about 10⁴ organisms. Ingestion of the organisms with food results in a similar lowering of the infectious dose. *V. cholerae* attach to the intestinal lining and secrete an enterotoxin that causes a high rate of discharge of a watery mucus. In addition to severe diarrhoea, symptoms include

abdominal cramps, nausea, vomiting, and shock. Fluid and electrolyte loss result in death rates of up to 60% without treatment. Symptoms may begin within 6 hours of ingestion of the organisms, probably because of the high infectious dose requirements.

V. cholerae was first discovered in 1854 by Filippo Pacini. However, the relationship of cholera to contaminated water was elucidated 30 years earlier in 1855 by John Snow in a study of the sources of a cholera epidemic in London. Two private water companies provided water to a London district on a competing basis. The water from the two companies was distributed somewhat randomly, much as long distance telephone service is in the USA today. Snow found that the incidence of cholera in residents of the company taking water from the Thames river near the city centre was approximately 10 times that of the company taking water from the Thames upstream of London.

Several strains of *V. cholerae* cause the disease. Prior to 1960, the strain isolated by Koch (O1) was dominant. In 1960 the *El Tor* strain became the more commonly reported

organism. In 1993, *V. cholerae* serogroup O139 (Bengal) became prominent in Bangladesh and eastern India and has since become endemic.

Cholera occurs world-wide but is far more prevalent in areas with inadequate protection of water supplies. The disease is endemic in over 80 countries and is responsible for approximately 10 000 deaths annually. Virtually absent from the Western Hemisphere for most of the 20th century, a cholera epidemic began in Peru in 1991 that gradually moved northward through Central America and Mexico. Over one million cases and 10 000 deaths were reported by 1995. The South American cholera strain has not crossed into the USA other than with infected travellers and through some imported seafood. A snapshot of the worldwide presence of cholera is provided

in Table 3.3, which contains incidence and death data for 1999. The total reported cases of cholera in 1999 were 254 310 with 9175 deaths. Of these, five cases were in Australia and New Zealand and 16 were in Europe. Note that 81% of the cases and 95% of the deaths were in sub-Saharan Africa. A number of Latin American nations had infection rates similar to countries in sub-Saharan Africa. Death rates were generally low in Latin America and Asia for those countries reporting.

Cholera is virtually absent in the USA, Europe, Australia and New Zealand where water supplies have been filtered and disinfected since the early part of the 20th century. Of those cases reported in the USA and Europe, a large fraction is associated with travellers from areas where cholera is endemic.

TABLE 3.3 Reported incidence and death due to cholera in 1999 (Source: World Health Organization, 1999)

<i>Africa</i>	<i>Cases</i>	<i>Deaths</i>	<i>America</i>	<i>Cases</i>	<i>Deaths</i>
Benin	855	25	Belize	12	0
Burkina Faso	93	6	Brazil	3233	83
Burundi	3440	63	Colombia	42	0
Cameroon	326	35	Ecuador	90	0
Chad	217	18	El Salvador	134	0
Comoros	1180	42	Guatemala	2077	0
Congo	4814	20	Honduras	56	3
Dem. Rep. of Congo	12 711	783	Mexico	9	0
Ghana	9432	260	Nicaragua	545	7
Guinea	546	44	Peru	1546	6
Kenya	11 039	350	USA	6	0
Liberia	215	0	Venezuela	376	4
Madagascar	9745	542	American total	8126	103
Malawi	26 508	648	<i>Asia</i>		
Mali	6	3	Afghanistan	24 639	152
Mozambique	44 329	1194	Brunei Darussalam	93	0
Niger	1186	85	Cambodia	1711	130
Nigeria	26 358	2085	China	4570	0
Rwanda	217	49	Hong Kong SAR	18	0
Sierra Leone	834	5	India	3839	6
Somalia	17 757	693	Iran	1369	21
South Africa	68	2	Iraq	1985	30
Swaziland	7	0	Japan	40	0
Tanzania U. R. of	11 855	584	Malaysia	535	0
Togo	667	31	Philippines	330	0
Uganda	5169	241	Singapore	11	0
Zambia	11 535	535	Sri Lanka	108	5
Zimbabwe	5637	385	Vietnam	169	0
African total	206 746	8728	Asian total	39 417	344

Non-virulent *V. cholerae* has been isolated from Chesapeake Bay, particularly Baltimore Harbor, quite routinely (Jiang *et al.*, 2000a,b). Similar results have been reported for near shore waters in Peru (Patz, 2000). Their presence was correlated to temperature and increasing salinity. The fact that a virulence factor was missing, together with the temperature dependence, suggests that the population was resident. Increases in cholera cases were correlated with temperature in Mexico (Patz, 2000) and in Peru (Checkley *et al.*, 2000). Isolation of *V. cholerae* from sewage in Lima was determined to have a threshold temperature of 19.3°C and correlated with hospital admissions (Speelmon *et al.*, 2000). In the 1991 epidemic in Peru, initial cases occurring at virtually the same time were geographically scattered. Considering there were no cases in 1990, the geographical scatter may suggest an environmental factor being involved (Seas *et al.*, 2000).

4.2 Gastroenteritis/campylobacteriosis

Gastroenteritis is a general term to describe watery diarrhoea that may be accompanied by abdominal cramping, nausea, vomiting, fever and chills. Among the waterborne and water-washed diseases the most commonly associated bacterium is *Campylobacter*. In the USA 45% of the people complaining of diarrhoea are found to have *Campylobacter* infections. Usually these symptoms occur within 24 hours of ingestion. Illness is usually self-limited and lasts 3 days. Although campylobacteriosis is not a reportable disease in the USA, the Centers for Disease Control and Prevention receive about 10 000 reports of the disease each year, approximately 1 per 100 000 population. The disease generally occurs sporadically and large outbreaks are unusual. In the USA an estimated 2 million cases of campylobacteriosis occur each year with 500 deaths. However, the disease is generally not considered life threatening unless the organisms get into the bloodstream or the infected person has a weakened or underdeveloped immune system.

There are 16 species of *Campylobacter*, all of which are pathogenic to humans or animals.

The species most commonly responsible for human infections is *Campylobacter jejuni*. Food, particularly poultry, is probably the most common mode of transmission, although waterborne infections are also common. *Campylobacter* are Gram-negative, motile spirilla and have been placed in the ϵ -subdivision of the proteobacteria. They are microaerophilic organisms and do not grow well at oxygen tensions greater than 10% of atmospheric. *Campylobacter* produces an enterotoxin similar to that of *V. cholerae* and the resulting diarrhoea is often bloody (Madigan *et al.*, 2000).

4.3 Salmonellosis

Depending on the source, 1000–2000 strains of the genus *Salmonella* have been identified. The discrepancy results from the change in identification methods from phenotyping to genotyping. Many organisms that were considered separate species through phenotyping have been grouped together as a result of information about their genetic makeup. In other cases, genotyping has resulted in organisms being placed in different genera. The recataloguing of large genera, such as *Salmonella*, is far from complete. Bacteria of the genus *Salmonella* are members of the γ -subgroup of the Proteobacteria. Close genetic relatives include the genera *Escherichia*, *Shigella*, *Proteus* and *Enterobacter*, and the group is commonly referred to as enteric bacteria. The World Health Organization considers all members of the genus *Salmonella* to be human pathogens. However, three species, *S. typhi*, *S. enteritidis* and *S. typhimurium* are the most important. *S. typhi* is the cause of typhoid fever and will be discussed in the following section. The other *Salmonella* infections are relatively mild, similar in their characteristics and severity to campylobacteriosis, and like campylobacteriosis, are more often associated with food than water. The ratio of waterborne to total infections in Israel dropped from nearly 10% in 1976 to less than 1% in 1995 as the result of improvements in water supply systems (Tulchisky *et al.*, 2000). Approximately 30% of the people with diarrhoea who see a physician in the USA are found to have salmonellosis. Whether

this statistic is valid in other countries is questionable because of differences in foods, food handling and water supply.

Symptoms of salmonellosis begin from 12 to 72 hours following ingestion. Duration of symptoms is usually less than one week and most people do not require treatment. As with campylobacteriosis, hospitalization is sometimes required. Unlike campylobacteriosis, the disease is notifiable in the USA. Thus statistics are maintained by the US Public Health Service on cases in which the organism is identified (Centers for Disease Control and Prevention, 1994). However, it is likely that only a small minority of infections are treated by a physician and within that group only severe cases would receive a full laboratory work-up. Thus the statistic of 40 000 cases of salmonellosis per year in the USA is probably very much lower than the actual number. Approximately 1000 people die each year in the USA due to salmonellosis. This is a large number for a disease generally considered a mild infection and would lead to speculation that the situation in developing nations is much worse.

Salmonella are zoonoses and are particularly associated with poultry, wild birds and reptiles. An outbreak estimated at over 200 000 infections in the USA in 1994 resulted from using non-pasteurized eggs in ice cream (World Health Organization, 1996).

4.4 Typhoid fever

Typhoid fever, caused by *Salmonella typhi*, is similar to cholera in severity but has much greater incidence and causes approximately 60 times as many deaths. Worldwide, the annual number of cases of typhoid fever is over 16 million, many times that of cholera, of whom nearly 4%, or 600 000, die. Death rates are between 12 and 30% when the disease is not treated and 1 to 4% when treatment is given. As in the other diarrhoeal diseases, the incidence and death rate is far greater in developing countries. There are approximately 400 cases in the USA and a little over 100 cases in the UK per year. Typhoid epidemics have not occurred in developed countries since the 1920s. However,

epidemics are relatively frequent in the developing world. There were over 7500 cases in Tajikistan in August 1996. Over 900 cases occurred in January 1996 in Ain Taya, an Algerian city of 45 000.

Typhoid fever is different from salmonellosis in that the responsible organism, *S. typhi*, after initially colonizing the intestines enters the bloodstream causing a systemic infection. Symptoms generally do not occur for at least a week and may take up to 3 weeks to develop. High fever, fatigue, abdominal pain and either constipation or diarrhoea are typical symptoms. Large outbreaks are generally associated with water contamination, as in both the Algerian and Tajikistan cases noted above, or street vendors. Typhoid fever is unusual in that up to 5% of those infected continue to shed infective cells for up to a year following recovery from the disease.

Three typhoid fever vaccines are currently licensed. One is a dead cell variety, one is based on polysaccharides from the cell wall, and the third is a live vaccine. The three vaccines have similar effectiveness but somewhat different periods of protection and side effects.

4.5 Shigellosis (dysentery)

Dysentery is most easily defined as bloody diarrhoea. Although bloody diarrhoea can result from several types of bacterial infection, the most common form is shigellosis, caused by bacteria of the genus *Shigella*. Shigellosis, like campylobacteriosis and salmonellosis, is more commonly associated with direct contact or contaminated food than water, as indicated in Table 3.1. Four species of the genus *Shigella*, *S. dysenteriae*, *S. flexneri*, *S. boydii*, and *S. sonnei*, are most commonly isolated from active cases. Endemic shigellosis is associated with *S. flexneri* and epidemic dysentery is associated with *S. dysenteriae*. Twelve serotypes of *S. dysenteriae*, 13 serotypes of *S. flexneri*, 18 serotypes of *S. boydii* and 1 serotype of *S. sonnei* have been identified on the basis of the O-specific polysaccharide of the lipopolysaccharide. Shigellosis is estimated to cause over 600 000 deaths worldwide each year (World Health Organization, 1998). In the USA,

the majority of infections are by *S. sonnei* and direct contact, such as occurs in day care centres, is a major mode of transmission (Centers for Disease Control and Prevention, 2001).

The disease results from growth of *Shigella* spp. in the epithelial cells of the colon. Invaded cells die and the infection spreads to adjacent cells. The result is ulceration, inflammation and bleeding. Classic symptoms of shigellosis include severe diarrhoea, a bloody stool, cramping, and tenesmus. Infection by *Shigella dysenteriae* Type 1, the principal cause of epidemic dysentery, may result in haemolytic uraemic syndrome (HUS), seizures, sepsis and toxic megacolon. A factor in the particular severity of *Shigella dysenteriae* Type 1 is the production of an enterotoxin that inhibits protein synthesis. Haemolytic uraemic syndrome will be discussed below in connection with *Escherichia coli* O15:H7. Up to 15% of *Shigella dysenteriae* Type 1 infections are fatal. As might be expected the most severe problems with dysentery, particularly that associated with *Shigella dysenteriae* Type 1, are in areas where public health facilities and sanitation are poor. Shigellosis in developed countries is a serious but less deadly problem, as indicated by the fact that most cases in the USA are caused by *S. sonnei*. Epidemic dysentery is widespread. Between 1968 and 1972, over 500 000 cases and 20 000 deaths occurred in Central America. The disease surfaced in Congo in 1979 and has been found throughout sub-Saharan Africa since then.

Bacteria in the genus *Shigella* are classified as enteric bacteria and are morphologically similar to other enteric bacteria such as *Escherichia*, *Salmonella* and *Klebsiella*. All are Gram-negative, motile rods and are non-sporulating and facultatively anaerobic. Enteric bacteria are oxidase negative and reduce nitrate only to nitrite (Madigan *et al.*, 2000). *Shigella* and *Escherichia* are closely related genetically with DNA homology generally greater than 70% between members of the two genera.

Shigella are commonly present in soil and water but are not always infective. For infection to occur the organisms must contain the invasion plasmid that allows it to bind to

the epithelial cells of the intestine and a smooth lipopolysaccharide somatic antigen. *Shigella* found in natural environments are often unable to initiate the disease.

Members of the genus *Shigella* have shown an impressive ability to develop antibiotic resistance. *Shigella dysenteriae* has shown a particular ability in this regard and often develops resistance to a new antibiotic within 2 years. Thus treatment of *Shigella* outbreaks is difficult and problems increase where endemic conditions exist.

4.6 Haemorrhagic colitis and haemolytic uraemic syndrome

Haemorrhagic colitis and haemolytic uraemic syndrome (HUS) are diseases that can be caused by a number of bacteria. However, the principal cause is six strains of the usually non-pathogenic species *Escherichia coli*. The strains are termed enteropathogenic (EPEC), verotoxigenic (VTEC), enterotoxic (ETEC), entero-invasive (EIEC), diffusely adhesive (DAEC), or enterohaemorrhagic (EHEC), depending on the type of toxin produced. The toxin produced by *E. coli*, Type O15:H7, the EHEC type, is indistinguishable from that of *Shigella dysenteriae* Type 1.

Escherichia coli is one of the predominant species present in the gut of warm-blooded animals and as such is used as an indicator of the possible contamination of food, water or surfaces with human or animal faeces. Pathogenic strains of *E. coli* were not reported prior to 1982 and it is unclear if the diseases are new or simply had not been detected previously. The severity of HUS is such that lack of identification of pathogenic *E. coli* as the cause would seem surprising. However, HUS is caused by other bacteria, most notably *Shigella dysenteriae* Type 1, and the ability to identify strains of species has increased rapidly over the past 50 years.

All of the pathogenic *E. coli* strains cause haemorrhagic colitis that results in abdominal pain and loose stools that may progress to bloody diarrhoea. Severity of the disease varies with minor cases lasting 1–2 days and more

severe cases lasting up to 2 weeks. The disease is not distinguishable from shigellosis or salmonellosis except by isolation of the organisms.

Haemolytic uraemic syndrome, caused by *E. coli* Type O15:H7, is a systemic disease that occurs principally in children under 10 years old. Approximately 10% of children with *E. coli* Type O15:H7 infections will develop HUS. The first stage of the disease begins with symptoms similar to gastroenteritis and haemorrhagic colitis and may last up to 2 weeks. Type O15:H7 is pathogenic because of the ability to produce the *Shigella* enterotoxin that inhibits protein production and induces fluid loss from epithelial cells. When the toxins enter the bloodstream red blood cells and the platelets that contribute clotting cells are destroyed. Kidney damage is common and generalized fluid accumulation is a result. Treatment consists of maintenance of fluids and salts and, in some cases, blood transfusions. There is no known way to stop or manage the progress of the disease. Approximately 90% of the children receiving supportive treatment survive the disease but up to 30% have permanent kidney damage that leads to kidney failure.

Outbreaks of *E. coli* Type O15:H7 infections have occurred world-wide. The most common source of infections is incompletely cooked meat, particularly minced (ground) beef. However, outbreaks have been traced to contaminated water. The most notable waterborne outbreak to date occurred in May, 2000 in the small community of Walkerton, Ontario where the community water supply system was contaminated. Six deaths and several hundred illnesses are attributed to the outbreak. The infection route is believed to have been poorly sealed wells, some of which were located downslope from cattle pastures.

4.7 Leptospirosis

Leptospirosis, caused by several species of the genus *Leptospira*, is a widespread disease with its principal reservoir in non-human animals. At least 100 000 cases of leptospirosis occur worldwide annually. Most infections are mild and do not require treatment. However, severe

cases may result in kidney failure, internal bleeding and pulmonary haemorrhage. Incidence in developed countries is low – there were 389 cases in the USA between 1985 and 1998. In the UK the rate is about 8 cases per year. In the Caribbean Islands the annual infection rate is between 5 and 30 cases per 100 000 population per year and the death rate is nearly 10%.

Leptospira invades the body through mucous membranes or through broken skin. Infections occur in the central nervous system, kidneys and liver but are most persistent in the kidney tubercles. The organism is discharged in the urine and contact with animal urine or with water contaminated with animal urine is the principal mode of transmission. Although pathogenic *Leptospira* species are commonly found in both wild and domestic organisms, the organism is endemic in rat populations and rats are usually implicated in outbreaks. Individual infections occur among swimmers, fishermen, cavers, rice field workers and abattoir workers. Outbreaks of leptospirosis usually occur following storms and it is presumed that the organisms are washed into watercourses where they come into contact with people. Diagnosis is difficult and the disease is often confused with others having similar symptoms. The incidence of leptospirosis is much greater in developing, particularly tropical, countries than in developed nations. Severe outbreaks occurred in the Azores in 2001, Orissa, India in 1999 and Nicaragua in 1995.

4.8 Legionellosis (Legionnaire's disease)

Legionellosis is a form of pneumonia caused by *Legionella pneumophila*. The name is derived from the first observation of the disease at a convention of the American Legion, a military veterans' organization, in Philadelphia, Pennsylvania in 1976 (Fraser *et al.*, 1977). Over 220 people contracted legionellosis, of whom 34 died, in this first epidemic. The source of the *Legionella pneumophila* was found to be condensate in the air-conditioning system.

Legionella pneumophila is a Gram-negative rod typically measuring from 2 to 20 μm in length. Specific growth requirements are L-cysteine and ferric iron. The organism is a normal inhabitant of natural waters and is commonly found in cooling towers, hot tubs (it grows well up to 45°C), hot water tanks, and sprays in vegetable sections of supermarkets (Madigan *et al.*, 2000; Winn, 1988). *Legionella pneumophila* is a facultative intracellular parasite. It does not multiply in sterile tap water but grows well when free-swimming amoeba cultures are added (Breiman *et al.*, 1990). In the lungs, macrophages engulf *Legionella pneumophila*, but the organisms are able to grow by blocking the fusion of lysosomes with the phagosome.

Legionella pneumophila infections are relatively common. Approximately 1000 confirmed cases of legionellosis occur in the USA each year. However, the Centers for Disease Control and Prevention estimate that the actual rate is 10 000–15 000 cases annually. Although the disease is reportable, most hospitals do not normally check for the organism. Most infections result in subclinical symptoms. However, immunocompromised individuals, young children and older adults are considerably more susceptible to development of severe symptoms. Epidemics, such as the one from which the disease derived its name, are unusual. Most cases occur one or two at a time and rarely are reported in the general media. To date the most common sources have been air-conditioning systems and cooling towers. Not surprisingly, legionellosis is largely a disease of the developed world. Very few data exist on the incidence of the disease in developing countries.

4.9 Peptic ulcer, gastritis and stomach cancer

In the late 1970s Marshall and Warren (1984, 1989) observed *Helicobacter pylori*, a motile Gram-negative rod that is oxidase, catalase and urease positive, consistently to overlay inflamed gastric tissue. They suggested that gastritis and peptic ulcers might be a bacterial disease rather than the result of stress and

excessive acid production. The suggestion was received with little enthusiasm from the medical community and probably even less from the pharmaceutical industry. However, Marshall fulfilled Koch's postulates by ingesting a culture of *H. pylori*, developing ulcers, and then curing the disease by taking antibiotics and bismuth and their suggestion began to be taken more seriously (Marshall *et al.*, 1988). Epidemiological studies have established a linkage of *H. pylori* infection to non-cardia gastric adenocarcinoma as well as peptic ulcers. However, removal of *H. pylori* appears to increase the risk of gastro-oesophageal reflux disease, Barrett's oesophagus and adenocarcinoma of the oesophagus (Blaser, 1992, 2000).

H. pylori appear to create relatively high pH microregions by secreting a urease with an unusually low Michaelis constant and by suppressing production of ascorbic acid (Moore, 1994). The organism secretes cytotoxins, proteases and phospholipidases that cause inflammation and damage to epithelial cells and decreasing protection from gastric acids. A suggestion has been made that *H. pylori* obtains sustenance from epithelial cell secretions and therefore irritation of the tissue is beneficial to growth (Blaser, 2000).

Human infection by *H. pylori* is very common. About 2% of children under age nine are infected. Infection rates increase with age until about age 50 when over 50% of all individuals world-wide are infected. Infection is significantly greater in lower economic levels and in developing countries where the rate is as high as 80% of all people. In the developing world, infection rates seem to be decreasing. The incidence of disease in individuals infected with *H. pylori* is quite high. In an Italian study of over 1000 adults, 42% were positive for *H. pylori* antibodies. Of the 420 positive individuals 121 submitted to endoscopy. Only 14 of the 121 were disease free (Vaira *et al.*, 1994).

The mode of *H. pylori* infection is unknown but assumed to be through contaminated food and water. Thus there is a possibility that peptic ulcers and gastric cancer are waterborne diseases. Given the prevalence of these diseases, transmission through water should be investigated.

5 INDICATOR ORGANISMS

Waterborne diseases pose very serious threats to society because of their potential to infect large numbers of individuals over a very short time. The common waterborne diseases are all incapacitating, many have high rates of death for untreated victims, and are particularly difficult problems for children, the elderly and those with challenged immune systems. Testing water for each of the possible pathogens, including viruses, protozoans, and worms is impractical. Early in the development of sanitary practice the need for *indicators of human contamination* was perceived. The issue presented was that if recent human contamination were established then there would be a certain probability of the presence of human pathogens. Accuracy of the relationship would be greater in urban than rural areas because there would be a greater probability of sick people in the general population. Thus the question asked was what could be monitored in water to determine if recent human contamination had occurred. In the early 1900s chemical analysis was nowhere near sensitive enough for chemical compounds to be considered as indicators. However, the ability to identify bacteria characteristic of the colon had been developed and methods were developed to estimate the concentration of these *coliform* bacteria. Consequently coliform bacteria of various types have become the standard method of establishing the biological quality of water (see Chapter 7).

5.1 Drinking water

There is little question that application of the coliform test (APHA, 1998) has been a major factor in the improvement and maintenance of drinking water quality. Whether monitoring a broader group of organisms would have had equally good results is unknown. High concentrations of any bacteria in drinking water would generally be discouraged by public health authorities even if there was no evidence of *E. coli* or enterococci. Water filtration and disinfection result in near sterile water and one usually devoid of coliforms. Microbial films

grow in water distribution lines but as long as the system is kept isolated there is not a problem with introduction of pathogens or of coliforms (Camper *et al.*, 1991). Thus samples taken at the tap are almost always free of coliforms in well-maintained and operated systems. However, recent experiences have shown that the lack of coliforms does not provide evidence of the absence of human pathogens, most notably protozoans and viruses. Some protozoans and viruses are much more difficult to remove by filtration and are more resistant to disinfection methods now in use than coliforms, or bacteria in general. The most interesting example is cryptosporidiosis, caused by the protozoon *Cryptosporidium parvum*. Cryptosporidiosis, a normally mild, self-limiting diarrhoeal disease was first recognized as a human disease in 1976, although it had undoubtedly been present for some time. The first report of cryptosporidiosis in association with a water supply that met all standards for coliforms was in 1987 in Carrollton, Georgia, where 13 000 people became ill. In 1993 the largest outbreak of waterborne disease recorded in the USA occurred when an estimated 400 000 people became ill with cryptosporidiosis in Milwaukee, Wisconsin. Again the water supply met all standards for coliform bacteria. Since that time cryptosporidiosis has become the most common known source of waterborne disease outbreaks in the USA.

5.2 Recreational and labour water quality

Individual contact with water occurs through recreational activities such as wading, swimming, boating and water skiing in a variety of venues, pools, ponds, lakes and rivers. Workers come in contact with water in agriculture, food processing and activities requiring washing facilities. Swimming pool water is usually filtered and disinfected but the density of use is greater than in other venues. In more natural venues, such as beaches, there is much less ability to control contamination. For example, beaches typically receive storm runoff from

adjacent land and often have one or more small streams or storm drains running across them. In many cases the streams disappear in sand prior to reaching the receiving waters during dry periods. Contaminants mobilized by storm runoff are carried into the streams and storm drains, onto beaches and into adjacent water. Thus animal wastes can be carried into natural recreational waters on a continual basis. In areas where human waste disposal is on land or in cesspools or septic systems there may be direct contamination of recreational waters following storms, also.

Workers contact water in fields during irrigation, in food processing where water is used for washing and for carriage of both products and wastes, and in general commercial activities where water is used to clean surfaces and equipment. There is potential for direct ingestion of contaminated water as well as for infection through the skin or through cuts and abrasions. In areas where water supplies are limited, water recycling is often practised. Without appropriate treatment, water recycling increases the potential for infections to occur.

Problems with using coliforms as indicator organisms in judging the quality of recreational and labour waters are quite different than for judging drinking-water quality. Infections of concern in recreational waters include skin, throat, and ear infections and parasitic diseases as well as the gastrointestinal diseases of concern in drinking water. Sources of contamination are much more diverse and far less controllable than is the case for drinking water and the types of contact with recreational and labour water are also quite diverse. Moreover, natural runoff, including irrigation water from both agriculture and parks, is commonly high in coliforms, *E. coli* and *Enterococcus* concentrations even when it is clear that human contamination is unlikely (Solo-Gabriel et al., 2000; Schroeder et al., 2002). Suspected sources of *E. coli* and *Enterococcus* are generally listed and include domestic, feral and wild animals, although it is apparent that these organisms can survive and grow in soil.

The microbial population in the guts of warm-blooded animals varies considerably, at

least in relative numbers of the various species (Madigan et al., 2000). Considerable effort has been invested in developing methods of determining the source of indicator organisms through identification of strains specific to certain animal species, ratios of *E. coli* and *Enterococcus* concentrations, and application of tools such as neural network analysis to microbial populations. To date a widely accepted method of determining the source of indicator organisms in a stream has not been developed. However, because of the number of zoonoses, the need to differentiate between contamination by humans and animals appears questionable except where epidemics are involved and the source is known.

At present biological standards of quality of recreational waters are based on coliforms, *E. coli* and *Enterococcus* concentrations (USEPA, 2000). The near ubiquitous presence of indicator organisms in both storm and dry weather drainage presents severe problems and would suggest that virtually all drainage needs to be disinfected if current standards are to be met.

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Protozoa

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1 INTRODUCTION

Protozoa are a large collection of organisms with considerable morphological and physiological diversity. They are all eukaryotic and considered to be unicellular. The majority of them employ chemoheterotrophic nutrition, but as certain species possess chloroplasts they can also practise photoautotrophy. Protozoa are found in almost every aquatic environment and are widely distributed. They play an important role in all aspects of sanitary microbiology ranging from their health impacts as human pathogens through to their role in the treatment of potable waters and wastewaters. With around 35 000 species they demonstrate a wide diversity in form and mode of life and occupy a range of ecological niches. The number of species together with the number of individuals of each species provides an indicator value as to the nature of the habitat in which they are observed. They play an important role in many communities where they occupy a range of trophic levels. As predators upon unicellular or filamentous algae, bacteria and microfungi, protozoa can be both herbivores and consumers in the decomposer link of the food chain. They are also a major food source for microinvertebrates, thus they have an important role in the transfer of bacterial and algal production to successive trophic levels.

Protozoa are defined as single-celled eukaryotic organisms that feed heterotrophically and exhibit diverse mechanisms of motility. As they are unicellular they lack tissues and organs, thus each protozoan cell is able to carry out all the processes that are

essential for life. The cell achieves this by means of organelles that are able to carry out specialized functions. Thus there are organelles for feeding (cell mouths), excreting (cytopyge) and locomotion (cilia, flagella, pseudopodia).

2 CLASSIFICATION OF PROTOZOA

In order to understand fully the contribution of protozoa to aquatic ecosystems and to exploit these same properties in engineered systems, it is essential to be able to identify and then classify them. Classification is an aid to identifying similar characteristics in behaviour and response, it proves very useful in the field as it removes the necessity of repeating experiments on each and every species.

Protozoa are classified on the basis of their morphology, in particular as regards their mode of locomotion. However, their classification is not simple as they are not a natural group but simply a collection of single-celled eukaryotes gathered together for convenience. In addition, classification is a fluid area as more information becomes available and ideas about evolution are refined. Classification of protozoa, in particular, is subject to regular revision and this can be an emotive and controversial issue. A complete revision of the protozoa took place in 1980 and this placed them in the Kingdom Protista, sub-Kingdom the Protozoa. They are then further divided into seven Phyla: I Sarcomastigophora, II Labyrinthomorpha, III Apicomplexa, IV Microspora, V Ascetospora, VI Myxozoa, VII Ciliophora. Of these, Phyla I and VII form the free-living protozoa whereas the remainder

are parasitic protozoa which parasitize plants, animals and algae. The free-living protozoa are aquatic species and are ubiquitous and widely distributed world-wide.

A much simpler scheme, proposed by Jahn (1979) and described in Ross (2000) is used in this article, which recognizes four major groups of protozoa and gives them the status of a Class. The first three Classes are the free-swimming protozoa and the last one comprises the parasitic protozoa.

1. *Mastigophora*: or flagellated protozoa (*Euglena*)
2. *Sarcodina*: or amoeba-like protozoa (*Amoeba*)
3. *Ciliophora* or ciliated protozoa (*Paramecium*)
4. *Sporozoa* or parasitic protozoa (*Plasmodium* – malaria)

Each of the four classes is discussed in turn in the following sections and the full scheme is illustrated in Fig. 4.1.

2.1 Mastigophora – the flagellated protozoa

The mastigophora are characterized by possession of one or more flagella, which are used

both for locomotion and feeding. Some flagellates have a characteristic arrangement of their flagella, e.g., *Dinoflagellates*. In common with the amoebae they usually multiply by longitudinal binary fission. They are the most primitive of the protozoa, believed to be the ancestors of the animals. Although some of them are free living, most are parasitic living in or on other organisms. These include *Trypanosoma gambiense* (the agent of sleeping sickness), *Trichonympha* (found in the guts of termites) and *Giardia lamblia* (causing acute diarrhoea).

Many flagellates are able to feed autotrophically as well as heterotrophically and the dinoflagellates are important primary producers (photosynthesizers) in lakes and oceans, yet they can also ingest prey and feed in an animal-like fashion. This makes them difficult to classify and they are often divided into two classes. The phytomastigophorea usually contain chloroplasts and are capable of autotrophic as well as heterotrophic nutrition. Consequently, many organisms classified as phytomastigophorea are also classified as algae, this includes such organisms as *Euglena*, *Volvox*, *Oicomonas* and the dinoflagellates. The remaining class, zoomastigophora, are heterotrophic

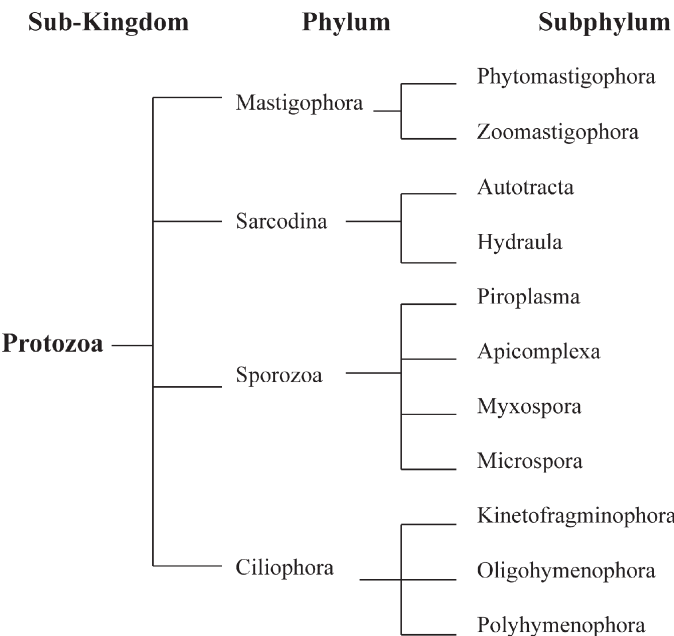


Fig. 4.1 Classification of protozoa which considers them as a sub-kingdom (from Ross, 2000).

and have oval-shaped bodies. They are generally the most numerous protozoa in terms of their total numbers in wastewater treatment plants such as activated sludge and trickling filters (Table 4.1).

2.2 Sarcodina – the amoebae

The Sarcodina possess pseudopodial structures that are used for movement and also for feeding, by means of protoplasmic flow. They demonstrate considerable diversity in that some of them lack any skeletal structure (naked amoeba), whereas others have elaborate shells known as tests (testate amoeba). The shells are composed of proteinaceous, siliceous or carbonaceous material and a wide range of structures have evolved, e.g. *Foraminiferans* contain shells of calcium carbonate which has been extracted from sea water. The shells do not cover the whole body, however, and naked pseudopodia are still used for feeding and locomotion.

Amoebae move by means of flowing cytoplasm, usually with the production of pseudopodia. The pseudopodial morphology has been used in classification of amoebae and the Sarcodina are divided into two major groups or classes. The Rhizopoda have unsupported pseudopodia, whereas the Actinopoda have radiating microtubule-supported axopodia.

TABLE 4.1 Representative protozoa from the three phyla found in activated sludge and trickling filters

Species	Subclass	Phylum
(a) <i>Bodo caudatus</i>		
(b) <i>Oicomonas termo</i>		Zoomastigophora
(c) <i>Amoeba proteus</i>		
(d) <i>Arcella vulgaris</i>		Sarcodina
(e) <i>Chilodenella uncinata</i>		
(f) <i>Colpoda cucullus</i>	Holotrichia	Ciliophora
(g) <i>Aspidisca costata</i>		
(h) <i>Euplotes moebiusi</i>	Spirotrichia	Ciliophora
(i) <i>Opercularia microdiscum</i>		
(j) <i>Vorticella microstoma</i>	Peritrichia	Ciliophora
(k) <i>Popodphyra collini</i>		
(l) <i>Acineta tuberosa</i>	Suctoria	Ciliophora

2.3 Ciliophora – the ciliates

The ciliates form an extremely large group and are the most specialized and complicated of the protozoans. They are characterized by the arrangement of cilia in an ordered fashion over the surface of the cell, and the presence of two kinds of nuclei, a macronucleus and a micronucleus. They also divide by transverse fission. Cilia, which serve to provide locomotion, are usually arranged in rows called kineties and the cilia beat with their effective stroke in the same direction. In addition, cilia are also distributed around what is the protozoal equivalent of a mouth, namely the cytosome. Here they provide an aid to feeding by the production of feeding currents.

The Ciliophora is the largest of the three phyla in terms of the number of species it represents, with over 7000 described in nature. The Ciliophora also provide the greatest species diversity in wastewater treatment plants, although not necessarily the largest number of individuals. Four distinct types of Ciliophora may be identified, based on their locomotion and arrangement of cilia:

1. Holotrichia – free-swimming protozoa which have cilia arranged uniformly over their whole bodies. Typical species are *Tracheophyllum* and *Litonotus*.
2. Spirotrichia – these possess a flattened body with locomotory cilia found mainly on the lower surface. The cilia concerned with feeding are well developed and wind clockwise to the cytosome. They are represented by *Aspidisca*, *Stentor* and *Euplotes*.
3. Peritrichia – these are immediately recognizable by their inverted, funnel- or bell-shaped bodies which are mounted on a stalk. The other end of this stalk is attached to particulate material such as a sludge floc, and serves to anchor the protozoan. In certain species the stalk is contractile. The wide end of the bell acts as an oral aperture in the Peritrichia and they have cilia arranged around this as an aid in feeding. Typical peritrichs are *Vorticella* and *Opercularia*.
4. Suctoria – the final type of ciliophora is ciliated only in early life, when the cilia

TABLE 4.2 Diseases associated with protozoal infection of humans

Niche	Protozoa	Condition
Skin	<i>Leishmania</i>	Cutaneous leishmaniasis
Eye	<i>Acanthamoeba</i>	Corneal ulcers
Gut	<i>Giardia</i> , <i>Entamoeba</i> , <i>Cryptosporidium</i>	Giardiasis, Cryptosporidiosis
Bloodstream	<i>Plasmodium</i> , <i>Trypanosoma</i>	Malaria, African sleeping sickness
Spleen	<i>Leishmania</i>	Visceral leishmaniasis
Liver	<i>Entamoeba</i> , <i>Leishmania</i>	Visceral leishmaniasis
Muscle	<i>Trypanosoma</i>	Chaga's disease

enable the young suctoria to disperse from their parents. After this they lose their cilia and develop a stalk and feeding tentacles. The stalk is non-contractile and attaches to particulate material, whereas the feeding tentacle is capable of capturing, and then feeding, on other protozoa. This is achieved by piercing them and sucking in organic material from the cytoplasm to form food vacuoles. Suctoria found in wastewaters include *Acineta* and *Podophyra*.

2.4 Sporozoa

All species of the sporozoa are spore producing and have no apparent means of locomotion (in other words they lack cilia or flagella). They are all parasitic protozoa which require the presence of a host organism (such as humans, animals and fish) to complete their life cycle. They generally obtain nutrients by absorbing organic molecules from the host organism.

Sporozoans often have very complicated life cycles and are able to exploit a range of ecological niches in the human body, including the skin, eye, mouth, gut, blood, spleen, liver and muscle, with an associated medical condition (Table 4.2).

3 MOTILITY

Motility plays an important role in protozoal survival. Most protozoa are required to move in order to find food or to avoid unfavourable

environmental conditions. The nutrition of protozoa also depends on the use of locomotory appendages to capture or collect food and direct it to the feeding apparatus. As a consequence, all free-living protozoa have some motility and can move freely in the environment at some time during their life cycle.

The different free-living groups have evolved diverse mechanisms for motility and these vary from the pseudopodia, utilized by the Sarcodina, through to the cilia and flagella employed by the ciliated and flagellated protozoa. Flagella and cilia are the most important organelles associated with movement in the protozoa and structurally they are very similar. However, whereas a cell will have only one or two flagella, ciliated cells have large numbers of cilia and this requires a complex system to coordinate their ciliary beat.

The motion of flagella is an undulatory wave beginning at the base of the flagellum and most flagella move only in a planar mode, although for some species, such as *Euglena*, movement is helical.

The pattern of waves generated by cilia is more difficult to discern. The numbers of cilia on each cell are more numerous, they are in close proximity to each other and the nature of the beat they generate is more complex than flagella. A cilium is characterized by a stroke which involves the cilium bending at the base while the rest of the cilium is straight. The cilium is then drawn back to its initial position close to the surface of the cell. This movement of flagella and cilia serves either to propel the protozoa through the water by the viscous forces which act on the cilia, or to generate water currents which permit feeding. The speed at which a protozoan moves is relatively constant and appears to be independent of the cell size. For ciliates this velocity is around 1 mm/s and for flagellates it is about 0.2 mm/s.

Many ciliated species can be observed not swimming, but walking or sliding along a solid surface and this is particularly evident in samples of activated sludge or trickling filter slime. It is thought that van der Waals forces help to hold the protozoa to these surfaces and

that the cilia aid in their movement along the surface. Often with these species, the cilia are organized in dense bundles known as cirri.

The Sarcodina have a very different motion involving frontal contraction. The protozoa extend a pseudopodium, which then attaches to the surface. This pseudopodium retracts strongly and pulls the testate form forward. It appears that when two pseudopodia are extended by the same cell, but at 180° to each other, they will compete against each other with the one which is most firmly attached determining the direction of movement.

4 PROTOZOAL NUTRITION

Protozoa demonstrate a wide range of feeding strategies of which four types are represented by the protozoa found in wastewater treatment systems. Certain members of the Phytomastigophorea are primary producers and capable of photoautotrophic nutrition, in addition to the more usual chemoheterotrophic nutrition.

Heterotrophy among the flagellated protozoa contributes to the process of biochemical oxygen demand (BOD) removal, and uptake of soluble organic material occurs either by diffusion or active transport. Protozoa that obtain their organic material in such a way are known as saprozoic, and are forced to compete with the more efficient heterotrophic bacteria for the available BOD. Amoebae and ciliated protozoa are also capable of forming a food vacuole around a solid food particle by a process known as phagocytosis. The organic content of the particle may then be utilized after enzymic digestion within the vacuole, a process which takes from 1 to 24 hours. This is known as holozoic or phagotrophic nutrition, and does not involve direct competition with bacteria, which are incapable of particle ingestion.

The final nutritional mode practised by the protozoa is that of predation. These predators are mainly ciliates, some of which are capable of feeding on algae (and are thus herbivores), as well as other ciliate and flagellate protozoal forms.

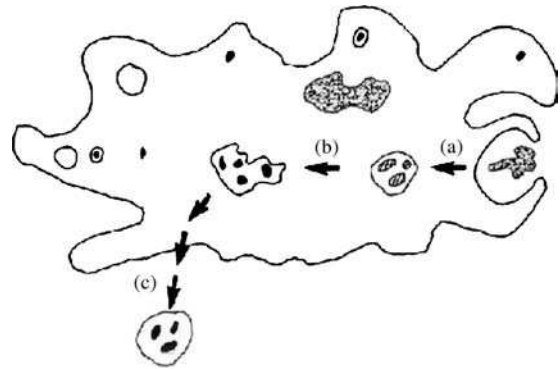


Fig. 4.2 Degradation of a food particle by phagocytosis. (a) Food particle engulfed by pseudopodia and a vacuole formed; (b) enzymic digestion occurs within the vacuole and digestion products released to the cytoplasm; (c) undigested remains ejected from the body.

All protozoa rely on phagocytosis for their energy and carbon for building cellular material (Fig. 4.2). This involves the enclosure of a solid food particle in a vacuole, which is covered with a membrane and in which digestion occurs. Dissolved nutrients are removed from the vacuole leaving the indigestible remains behind. These are removed from the cell by fusion of the vacuole with the cell surface membrane. The typical lifetime of a food vacuole is around 20 minutes, although this time reduces if the cell is not feeding.

In addition to phagocytosis, there are other mechanisms by which a protozoan can obtain energy and cellular building blocks. Some protozoa participate in symbiotic relationships with photosynthetic organisms, whereas others are thought able to take up dissolved nutrients. It is doubtful, however, if this latter mechanism plays any role for the free-living protozoa outside of a laboratory culture.

Although phagocytosis is practised by all the protozoa there are a number of different feeding patterns which are exploited to capture the solids particle and these can be classified into three categories, namely: filter feeders, raptorial feeders and diffusion feeders. Filter feeding involves the creation of a feeding current, which is then passed through a device which acts to

filter out the solids particles in the water. In the flagellates this is a collar of straight, rigid tentacles. For the ciliates the water is passed through an arrangement of parallel cilia. The clearance between the tentacles in the collar and the parallel ciliates, dictates the size of particle that is retained. This is typically between 0.3 and 1.5 μm and helps to explain why the presence of a healthy ciliate population in an activated sludge plant generates such a crystal clear effluent with a reduced number of faecal indicator bacteria (Table 4.3).

Raptorial feeding is practised in small flagellates and amoebae, which use it to feed on bacteria. In this mode, water currents are driven against the cell using a hairy anterior flagellum. Particles which make contact with a lip-like structure on the protozoa are phagocytized (Fig. 4.2). As each particle is captured separately (compared to filter feeders which retain all particles of the correct size), this allows the protozoa some discrimination as to what is ingested. This may be based on prey size or type, such as algae or small flagellates. Protozoa are also able to discriminate between different bacterial species with preferred types being selected.

Diffusion feeding is practised by the sarcodines. The suctorians are common protozoa in activated sludge and they feed by diffusion, largely on other ciliates. The suctorians are attached to a floc particle by a stalk and they have bundles of tentacles supported with an internal cylinder of microtubules. Ciliates which touch these tentacles become attached

and immobilized. The tentacles then penetrate the attached ciliate and draw the contents through the tentacle into the suctorian.

Within the latter two modes of nutrition, the protozoa display a certain degree of selective feeding. Larger forms of amoebae are carnivorous, eating mainly ciliates and flagellates, whereas the smaller amoebae feed primarily on bacteria. The predatory suctorians are found to feed almost exclusively on holotrichous and spiriotrichous ciliates, with hypotrichs, flagellates and amoebae rarely being captured. Peritrichous ciliates are primarily bacterial feeders, but have a limited number of bacterial species upon which they can feed. Certain bacterial species are capable of supporting growth for long periods, whereas others induce starvation after a short time. In addition, many bacteria, in particular the pigmented types, prove toxic to those ciliates which ingest them.

5 PROTOZOAN REPRODUCTION

For the free-living protozoa, their growth curve involves an increase in cell size followed by some form of asexual reproduction such as binary fission. A single cell divides into two and generates two daughter cells. As the protozoan is not symmetrical the two daughter cells are not identical initially, however, the differences soon disappear. Sexual reproduction is only resorted to in times of stress or adversity, for instance if the food supply diminishes. There are many types of binary fission in which the plane of division varies among the different groups. Division is simplest for the naked amoebae as this has no clear division plane. The cells become rounded and division of cytoplasm into two equal daughter amoebae occurs. It is more complex in the testate amoebae as there is both protoplasm and skeletal structure to reproduce. The flagellated protozoa undergo division along a longitudinal plane and the ciliated protozoa divide along a transverse plane. The Suctoria have a sedentary lifestyle spent attached to particulate material. Thus a method of asexual reproduction involving fission would lead to a rapid increase in their population with

TABLE 4.3 The effects of ciliated protozoa on the effluent quality from a bench-scale activated sludge plant (from Pike and Curds, 1971)

Parameter	Without ciliates	With ciliates
BOD (mg/l)	53–70	7–24
COD (mg/l)	198–250	124–142
Organic nitrogen (mg N/l)	14–21	10
Suspended solids (mg/l)	86–118	6–34
OD ₆₂₀	0.95–1.42	0.23–0.34
Viable bacterial count (cfu/ml $\times 10^6$)	106–160	1–9

increased competition for food and space. Consequently they divide by budding in which the new cell appears as a slight protuberance on the parent cell surface. This is evaginated and liberated, but unlike its parent it is motile in its immature stage, for around 30 minutes, which allows it to migrate and reduce crowding.

The most common form of protozoan reproduction is known as binary fission in which the organism divides into two equal-sized daughter cells. In the ciliated protozoa binary fission is usually transverse with the posterior end of the upper organism forming next to the anterior end of the lower one.

Binary fission generally produces daughter cells with genetic material (DNA) identical to that of the parent. It is an efficient way for protozoa to increase in number during periods when environmental conditions are relatively stable. However, when environmental conditions begin to change, sexual reproduction generally becomes more prevalent. Sexual reproduction allows for the mixing of DNA among the various strains (asexual daughters) of a local protozoan population. Although it has proved possible to maintain populations of protozoa for many years (equivalent to many hundreds of protozoan generations), the introduction of genetic exchange by sexual reproduction produces cells which are genetically different from each other, an important characteristic as genetic make-up determines how cells respond to their environment. A genetically diverse population is more able to adapt to changing conditions. In the competitive world of a wastewater treatment plant the more diverse the gene pool for a species, the greater the likelihood that it will persist over a wide range of changing environmental and operating regimes.

6 IMPORTANCE OF PROTOZOA IN WATER AND WASTEWATER TREATMENT

The importance of protozoa in water treatment systems focuses primarily on their ability to cause disease if ingested in a potable water

source. Of particular importance in this respect are the protozoan parasites *Giardia lamblia* and *Cryptosporidium parvum*, which inhabit the gastrointestinal tract and are a major world-wide cause of human morbidity and a significant contribution to mortality. More recent opportunistic pathogens, which infect immunocompromised individuals, include *Cyclospora cayetanensis* and a number of genera from the Phylum Microspora, such as *Encephalitozoon*, *Entereocytozoon* and *Septata*.

In addition to being parasites, protozoan populations are also subject to parasitism by a large number of fungi, bacteria and other protozoa. The opportunistic human pathogen *Legionella pneumophila* is one such bacterium which is able to parasitize the ciliated protozoan *Tetrahymena pyriformis* and it is thought this mechanism may help its distribution and survival in aquatic environments (Fields *et al.*, 1984).

In wastewater treatment, protozoa are known to play a major role but the full extent of their contribution is not fully quantified. Their major role is probably in maintaining a slime layer in trickling filter systems and in aiding flocculation in activated sludge systems. As shown earlier (Table 4.3), they aid in achieving a good effluent quality by their predatory role in removing bacteria and other small particles. The indicator value of protozoa in treatment systems, together with a fuller account of their role in wastewater treatment, is found in Chapter 22. A simplified key to identifying the protozoa of importance in wastewater treatment systems is found in Patterson (1998), which contains a very useful guide to the influence of organic loading rate and cell residence time on the distribution of protozoal species.

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Filamentous fungi in water systems

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1 INTRODUCTION

Fungi are ubiquitous, occupying every conceivable niche and habitat, displaying ecological strategies that range from micropredators and pathogens to more benign saprobes or symbionts (Dix and Webster, 1995). Such a diversity renders a brief circumscription of filamentous fungi near-impossible, but it includes eukaryotic microorganisms which, for at least some part of their life cycle, display a growth habit comprising a mycelium composed of individual hyphal elements, and which obtain nutrition by the absorption of extracellularly digested material. In many cases the mycelium lacks rigid self-support and is vulnerable to physical damage, but this is met in the majority of fungi by close adherence to surfaces or penetration of the substrate and growth occurring within. A detailed discussion on the biology of the fungal organism is given by Wessels and Meinhardt (1994).

As well as occupying a vast array of habitats, fungi utilize many different media and vectors for dispersal of their spores or other propagules such as sclerotia, sporangia or hyphal fragments. This can involve air (Lacey, 1996), water (Dix and Webster, 1995), plants, including their seeds (Richardson, 1996) animals (Andrews and Harris, 1997), and all kinds of organic and inorganic debris. Despite such a wide range of dispersal mechanisms (Ingold, 1971), some of which are highly-targeted, many fungi appear to rely upon high productivity and subsequent chance encounter with a substrate suitable for growth, and given the preponderance of chance

encounter strategies among fungi, it is inevitable that many will find their way into water systems. Moreover, the fungi present are likely to be highly diverse and include more than just those primarily adapted for aquatic habitats.

Water is as essential for the growth of fungi as it is for any other organism and, although fungal dispersal structures are often resistant to desiccation, mycelial hyphae are generally poor at withstanding dry conditions. Most require comparatively high levels of water in the environment to enable vegetative growth (Dix and Webster, 1995). The majority of fungi prefer wet aerobic conditions and most of those that are known in culture can be grown in liquid media, in some cases including unamended water (Kelley *et al.*, 2001). Irrespective of a dependency on high water levels, many fungi excel as oligotrophs, scavenging nutrients from whatever sources that may be available, however dilute or recalcitrant (Parkinson *et al.*, 1989; Wainwright *et al.*, 1992, 1993; Ghariieb and Gadd, 1999; Karunasena *et al.*, 2000). It is unsurprising therefore that fungi are frequently found in waters of all sorts and that their activities contribute to many of the microbial processes which are of concern to the water-related industries, including biodegradation, bioremediation, biofilms, biofouling, as well to health effects and taste and odour issues.

Kelley *et al.* (1997) reviewed the literature on fungi from water distribution systems. They concluded that comparatively few systematic surveys of fungi in water distribution

systems had been carried out and, in many cases, fungi were detected only incidentally during the course of bacterial isolation programmes. Fungi are to be found in raw, surface and groundwaters and may increase in numbers in storage reservoirs. More fungi are present in surface water than in groundwater. More fungi are present in untreated than in (chlorine) treated water. Fungi are less susceptible to chlorine treatment than are bacteria. Sedimentation and flocculation remove many fungi, but rapid filtration is not an effective treatment. There is evidence to suggest that fungi survive and multiply in distribution systems in surface films and in sediments, particularly where conditions are warmer or flow rates are restricted. Fungi are capable of contributing off-tastes, odours and flavours, but no links to ill health had been reported.

Isolation techniques for fungi from water have also been evaluated (Anon., 1996). This has included an assessment of the diversity of fungi that can be detected and the reproducibility and significance of fungal counts. A study of fungal diversity and significance in the two USA distribution systems has also recently been conducted (Kelley *et al.*, 2001), involving samples taken from several points throughout the systems. This included an examination of the ability of selected fungi to grow in water and to produce metabolites of concern to human health, and also looked at the effectiveness of different control measures. Kinsey *et al.* (1999) have summarized many of the main considerations in sampling fungi from water.

2 STUDY METHODS

The main elements are sampling, detection, identification, quantification and enumeration. Implicit throughout all of these must be an awareness that, due to the exceptional diversity of fungi, there is no single approach that will be effective for all fungi. The study of fungi has relied on the development of technologies and techniques that can be adapted for the purpose (for a historical perspective see

Webster, 1996). Increasing sophistication has led to some highly sensitive and specific techniques for particular fungal species (Paterson and Bridge, 1994; Bridge *et al.*, 1998) but, in many cases, these require further development before enabling determinations directly from environmental samples or for resolving mixed populations to species level. Hence, there is still a necessary reliance on more traditional microbiological methods.

2.1 Sampling

Standard procedures for taking water samples for microbiological examination are also adequate for fungi (Anon., 1994). Due to their filamentous nature fungi tend to be less evenly distributed in water than bacteria and so due consideration must be given to representativeness of sample volumes. Ideally, samples should be processed at the place and time of sampling, but, if as is usually the case, storage and transport is unavoidable then samples should be kept cool (2–10 °C and not exceeding the temperature at the site of sampling), and processed as soon as possible, preferably within 6 hours, or exceptionally up to 24 hours and the extent of the delay is recorded. Inadequate storage or delayed processing may mean that fungi present as hyphae are lost, while those present as spores may germinate and proliferate leading to disproportionate recording of the species present. Other factors which may affect fungi present in samples prior to processing include light and vibration. Exposure to light can affect fungal growth and sporulation; hence samples should be stored in the dark. Irregular sudden vibration can cause fragmentation of mycelia and may lead to dispersal of spores; conversely more regular repetitive motion can set up circular rotations which if prolonged can unite the mycelial biomass as a pellet.

There are inevitably interfaces with surfaces of pipework and other objects in any water system, and it is niches such as these where fungi are most likely to be found growing. If access is possible, such surfaces should be sampled by taking swab or sediment samples. Ideally sampling should occur immediately

upon exposure to the site, any delay between exposure and sampling should be recorded. Once taken, the swabs or sediment should not be allowed to dry any further; again cool storage and prompt sample handling will help prevent this occurrence.

2.2 Detection

It is not usually possible to distinguish fungi in water samples with the naked eye and some form of concentration is usually required as an initial step. Filtration is often employed, similar to bacteriological methods (Anon., 1994), whereby samples are passed through 0.4 μm pore size filters and the filters retained for subsequent examination. Alternatively, centrifugation at 1500 g for 3 minutes is sufficient to pellet fungal material in 10 ml sample volumes. Although filtration is very convenient, it is possible that some fungi may be 'lost' on the filter, whereas centrifugation has the advantage of resulting in a pelleted mass that can be resuspended for further examination. Swabs or sediment samples can be handled directly. There are other methods that can be used as part of the initial stages of detection, such as enrichment or baiting techniques, and these are designed with specific fungi in mind (see Diversity below).

2.2.1 Direct observation

In the majority of cases it is too time-consuming to examine water samples directly. However, for rapid confirmation of the presence of fungi, especially in instances of heavy contamination, it may be worthwhile to examine swabs or scrapings by directly making a slide preparation for microscopy. It is also possible to observe filters directly, either by using light or electron microscopy (Anon., 1996). Similarly, pelleted samples can be directly observed by simply resuspending them in a drop or two of water. However, it is largely impractical to attempt subsequent removal of individual spores or hyphal elements for further study. Hence, direct observation is essentially limited to determination the nature of the fungal material (i.e. hyphal fragment or spore) and

describing the morphology of the material (size, shape, colour, etc.). There is an established practice of examining fungal spores from environmental samples, particularly from the air (Lacey, 1996), but it has always been the case that when only spores or small fragments are available, it is rare to be able to identify the fungi with any confidence. The main exceptions are the aquatic and aero-aquatic hyphomycetes (see Diversity below), many of which have complex spores.

2.2.2 Culture methods

Many fungi can be cultured from their spores or hyphal fragments and, given suitable growth media (see entry under 'media' in Kirk *et al.*, 2001; Smith and Onions, 1994) and incubation conditions, will germinate, grow and eventually produce spores and spore-bearing structures (Arx, 1981). It is the details of complete reproductive structures that are required for species-level identification. The main disadvantage of culturing fungi is that it requires the intermediate and often time-consuming step of obtaining each organism in isolation on appropriate media. Nor is this guaranteed to result in an identification as not all fungi will grow in laboratory conditions, some never sporulate and, even more frustratingly, some fungi are identifiable only once – details of their primary host or other substrate specificity are known – data which are never likely to be known for cultures derived from spores present by chance in water samples. Nonetheless, culture-based techniques are the main methods for fungal analysis.

Filters through which samples have been passed can be plated directly onto the surface of nutrient agar media and the resultant fungal colonies can be counted and picked off for subculturing. Passing a range of volumes through separate filters avoids the possible extremes of too much or too little material being present as at least some will contain discrete fungal colonies sufficient for enumeration. However, it is preferable to resuspend fungi from filters by placing them in a tube with a known volume of sterile water. By varying the volumes into which the filters are resuspended it is possible to create a dilution

series which can then be spread-plated or pour-plated onto the agar media. The resultant fungal colonies can be counted and picked off to prepare pure cultures for identification. The range of dilutions required is a trial and error process depending on the density of fungi present in the particular sample, but a range of 10-fold dilutions from 10^0 – 10^{-5} is sufficient for most cases. Fungal isolation programmes are prone to suffer from occasional fast-growing colonies which rapidly spread over the entire plate making subsequent counts and subculturing impossible; the only solution is vigilance and prompt picking-off, even to the point of examining plates with a dissecting microscope to detect colonies before they are visible with the naked eye. Colony counting can be continued if required by marking the Petri dish below each point of removal with a permanent pen.

There are many variations of culture methods for fungi, and one of the perennial areas of debate is choice of growth medium. There is a need to standardize media and associated incubation conditions and to some extent this has begun (Anon., 1996), but at the same time an awareness has to be maintained as to the limitations of individual growth media and culture techniques. A similar situation exists for the examination of fungi in soils, where limitations of media and techniques have been known and discussed for some time (Cannon, 1996).

2.2.3 Biochemical methods

There are emerging biochemical techniques which are showing considerable promise for use in detecting fungal populations in environmental samples.

Detection of fungi by HPLC analysis of ergosterol is one such technique which has been applied to fungi from a range of different environments (Newell *et al.*, 1987; Johnson and McGill, 1990; Martin *et al.*, 1990; Trevisan, 1996; Dales *et al.*, 1999; Pasanen, *et al.*, 1999; Montgomery *et al.*, 2000; Saad *et al.*, 2001), including fungi from water samples (Gessner and Schwoerbel, 1991; Gessner and Chauvet, 1993; Suberkropp *et al.*, 1993; Newell, 1994; Bermingham *et al.*, 1995; Kelley *et al.*, 2001).

Ergosterol is a membrane lipid found in virtually all filamentous fungi (Weete, 1989) but in very few other organisms (Newell *et al.*, 1987). The extraction of ergosterol from a sample is comparatively simple; a microwave-assisted method is among the most convenient (Young, 1995; Montgomery *et al.*, 2000). Using dried filters, this essentially involves the addition of methanol followed by brief (35 s) irradiation in a domestic 750 W microwave oven set at medium power and then extraction by using petroleum ether. Coupled with HPLC this enables rapid and very sensitive detection of any ergosterol present.

Chitin has also been used as an indicator of fungi (Martin *et al.*, 1990), but it is not specific to fungi nor are assays as sensitive or convenient as for ergosterol. There are many other techniques targeted towards gene products and metabolites but, whether employing chromatographic, spectroscopic, substrate-degradation or immunoaffinity methods (Paterson and Bridge, 1994; Kelley *et al.*, 2001), these tend to be specific for individual taxa and, as such, cannot be used for general detection of fungi in a sample. They do, however, have considerable value for analysis where only particular compounds are of concern (see Secondary Metabolites below).

With the continued development of genomic techniques, it is now possible to detect and analyse fungal DNA in environmental samples (Bridge *et al.*, 1998). These techniques are based on the polymerase chain reaction (PCR) (Edel, 1998). A widely used approach is to conduct PCR using oligonucleotide primers specific for the conserved flanking regions of the internally transcribed spacers (ITS) of the fungal rRNA gene (White *et al.*, 1990). The ITS primers are regarded as universal for fungi because rRNA genes appear to be present in most fungi (Gardes and Bruns, 1993). The technique has the potential for very sensitive detection of fungi in environmental samples. However, with environmental samples there is a need for caution as a range of factors can bias the DNA amplification, such as achieving initial DNA extractions for only a subset of the total species present, the presence of compounds

inhibitory to amplification, obtaining primer-primer annealment, or the formation of chimeric co-amplified sequences (Wang and Wang, 1997; Hastings, 1999; Head, 1999). As PCR techniques continue to become more widely used, further development will undoubtedly resolve many of these potential problems (Viaud *et al.*, 2000).

It is usual to incorporate further analysis of the PCR product, and this essentially involves either sequence analysis, restriction fragment analysis, or the use of taxon specific probes or primers (Bridge *et al.*, 1998). In many cases the techniques target specific fungal taxa but some, although originally designed for bacteriological studies (examples in Hurst *et al.*, 1997; Edwards, 1999), are targeted much more broadly and are being adapted for analysing community structures in mixed fungal populations.

Suitable techniques include denaturing gradient gel electrophoresis (DGGE) (Hastings, 1999), temperature gradient gel electrophoresis (TGGE) (Hastings, 1999), and single strand conformational polymorphism (SSCP) (Fujita and Silver, 1994). DGGE involves running PCR product (using ITS primers) through a gel containing a gradient of denaturants; the sequence of the bases in the DNA determines at what point on the gel's gradient the strands denature and hinder further migration, so that on visualization the resulting bands reflect the diversity of fungi present in the original sample. DGGE is now beginning to be used for fungal populations (Kowalchuk *et al.*, 1997; Vainio and Hantula, 2000). TGGE is similar except that a fixed concentration of denaturant is used and the temperature is steadily increased during migration to mediate strand separation, which again is sequence-sensitive. Smit *et al.* (1999) applied TGGE to analyse wheat rhizosphere fungi. Both DGGE and TGGE provide profiles of fungal populations, but it is not possible to identify the fungi thus detected without comparison to the behaviour of test strains on the same gel and which therefore requires *a priori* knowledge of the fungi likely to be encountered, or by recourse to sequencing techniques. SSCP uses pre-denatured PCR product added to a non-

denaturing gel, and as the strands partially re-anneal, the conformation they adopt is sequence-dependent and alters their mobility through the gel. Walsh *et al.* (1995) have used SSCP to discriminate a range of opportunistic fungal pathogens; identification is effectively by comparison with test strains and, as with DGGE and TGGE, requires *a priori* knowledge of the fungi likely to be encountered.

Another promising technique for population analysis is fluorescent *in situ* hybridization (FISH) (Lindrea *et al.*, 1999). This is somewhat different in that it involves intact cells, permeabilized by means of enzymes or detergents, to which species-specific oligonucleotide probes carrying fluorochromes are added; the probes enter the cells and hybridize with the complementary nucleic acid sequence if present, and the unbound probe is washed away prior to observation. Spear *et al.* (1999) have used FISH to analyse *Aureobasidium* populations on leaves but, while FISH has the advantage over DGGE and TGGE in that it enables immediate identification, it is dependent on developing sufficient probes to detect every possible fungal species that might be present, again requiring *a priori* knowledge of the fungi likely to be present.

Where individual fungal species are of interest it is possible to use much more specific PCR-based techniques, many of which are well-established for use with fungi (Bridge *et al.*, 1998), such as amplified fragment length polymorphism (AFLP), which has been used to study *Aspergillus niger*, *Fusarium graminearum* and *Penicillium chrysogenum* isolates from water systems (Kelley *et al.*, 2001).

2.2.4 Assessing activity

In some situations it may be required to detect fungal activity. Direct observation can only provide data on a presence-or-absence basis and does not necessarily even distinguish viable from non-viable. Culture methods only include viable organisms but do not distinguish between resting spores and active hyphae. The molecular methods described above likewise do not give an indication of whole-organism viability or activity. Techniques

which can provide an indication of activity include staining of active regions (Cox and Thomas, 1999) or monitoring respiration and ATP (Suberkropp, 1991; Suberkropp *et al.*, 1993); however, with water systems there is the additional challenge, yet to be addressed, of how to sample representative volumes without altering the activity of the fungi in the process of collecting or concentrating the sample.

2.3 Identification

A consequence of the large range of fungal diversity and the need for an expert taxonomic knowledge is that there is a tendency for microbiological surveys to curtail fungal identifications once the level of genus has been reached. However, only in a very few cases does this provide useful information or enable specific cross-referencing to literature data on ecology, metabolites or health implications (for example Table 5.5 shows literature data on fungi producing taste and odour compounds). Identification of fungi is still heavily dependent upon microscopic observation of the physical appearance of both spores and spore-bearing structures, usually achieved by first obtaining the fungi in pure culture. Examination of just the mycelium rarely enables an identification, nor are spores alone sufficient as there are many species producing very similar spores. The UK has an organised network, the UK National Culture Collection (www.ukncc.co.uk), through which it is possible to obtain appropriate taxonomic advice and services on many microorganisms, including fungi.

2.4 Enumeration and quantification

2.4.1 Fungal counts

Enumeration of fungi usually relies on counts of colony-forming units (CFU) obtained by using dilution plates, and attempts critically to assess such methods have been made (Anon., 1996). The data generated by such means are notoriously difficult to analyse as there is very high variability and low reproducibility. The filamentous habit and the production of clus-

tered spore masses by fungi contribute to causing this, i.e. the individual fungus may comprise a discrete sporulating colony but which, during the course of sampling and plating, can become fragmented into many smaller pieces and/or the spores dispersed. Hence, a single individual could give a count greater than one. The degree of fragmentation or spore dispersal is unlikely to be the same for different species, so that some fungi may be over-represented in relation to others. The fungal CFU count from an environmental sample, while it may give an indication of the potential inoculum carried in the sample, is therefore unlikely to reflect accurately the number of individual fungi initially present. Hence a considerable degree of caution is required when interpreting fungal count data. There are methods for statistical analysis, such as most probable number (MPN) estimation, but while this is well established for bacterial counts (Anon., 1994), it has been little used for fungal organisms (Feest and Madelin, 1985; Cooper, 1993). Despite the problems inherent in obtaining fungal counts, they nonetheless provide valuable information, particularly where an assessment is required of the potential for fungal contamination of materials that come into contact with the water.

2.4.2 Quantification

Determining biomass by dry weight is problematic as quantities in water samples are likely to be very low to the point of negligibility for most laboratory balances (i.e. sub-milligram). Metabolite analysis is a possible way forward and HPLC of ergosterol, as well as being a promising technique for fungal detection, has the potential to be used for quantification, albeit without discrimination of individual species (Newell *et al.*, 1987; Nilsson and Bjurman, 1990; Johnson and McGill, 1990; Schnürer, 1993; Pasanen *et al.*, 1999; Montgomery *et al.*, 2000). As a key membrane component, ergosterol is likely to provide a good indication of fungal surface area present in a sample rather than an accurate measure of biomass (Ruzicka *et al.*, 2000). Other measures have been used to estimate biomass including hyphal length (Schnürer, 1993) and ATP (Suberkropp, 1991;

Suberkropp *et al.*, 1993; Ruzicka *et al.*, 2000), but these require the fungi to be in grown in isolation from other organisms.

3 DIVERSITY

3.1 Higher taxa containing filamentous fungi

Fungi and fungi-like organisms are presently classified across three kingdoms, and those which show some form of filamentous growth are found in two of these kingdoms (Table 5.1) Kirk *et al.* (2001) provide background information and further references for all fungal taxa to the genus level.

3.2 Filamentous fungi in water

The term ‘aquatic fungi’ is somewhat misleading in that many fungi normally considered as terrestrial have been isolated from water samples. In many cases they comprise the majority of fungi detected (see Table 5.2) Similar species have been reported by others with the main genera broadly the same in many other listings of fungi from water

samples (references in Kelley, *et al.*, 1997). These ‘terrestrial’ fungi are predominantly the asexual hyphomycete states (genera such as *Acremonium*, *Alternaria*, *Aspergillus*, *Aureobasidium*, *Cladosporium*, *Penicillium* and *Trichoderma*) and coelomycete states (e.g. *Phoma*) of Ascomycota, and also include Zygomycota (*Absidia*, *Mucor*, *Mortierella* and *Rhizopus*) and Basidiomycota (see Table 5.3). Conversely, substrates such as soil can yield fungi that are interpreted as aquatic (Hall, 1996). Moreover, for the majority of fungi the ability to survive and grow in water has not been examined.

There is usually an implicit restriction of the term ‘aquatic fungi’ to mean freshwater fungi, with marine and brackish water fungi receiving separate treatment (see Kohlmeyer and Kohlmeyer, 1979; Moss, 1986; Kohlmeyer and Volkmann-Kohlmeyer, 1991; Hyde, 1997; Kis-Papo, 2001).

There will always be fungi which remain undetected. For example, it would be reasonable to expect common Urediniomycetes and Ustilaginomycetes (rust and smut fungi) to be present by chance in water samples as spores. However, as highly specific biotrophic plant pathogens, there are a few culture-based

TABLE 5.1 Higher taxa containing fungi and fungus-like organisms, taxa containing filamentous fungi are in bold (following Kirk *et al.*, 2001)

Kingdom: PROTOZOA		
Kingdom: CHROMISTA		
	Phylum: Hyphochytriomycota	
	Phylum: Labyrinthulomycota	
	Phylum: Oomycota	Class: Oomycetes
Kingdom: FUNGI		
	(‘lower fungi’)	
	Phylum: Chytridiomycota	Class: Trichomycetes
	Phylum: Zygomycota	Class: Zygomycetes
	(EUMYCOTA)	
	Phylum: (‘Symbiomycota’ or Glomomycota)	Class: (‘Symbiomycetes’ or Glomomycetes)
	Phylum: Ascomycota	Class: Ascomycetes
		Class: Neoelectomycetes
		Class: Pneumocystidiomycetes
		Class: Schizosaccharomycetes
		Class: Saccharomycetes
		Class: Taphrinomycetes
		Class: Basidiomycetes
		Class: Urediniomycetes
		Class: Ustilaginomycetes
	Phylum: Basidiomycota	

TABLE 5.2 Numbers of samples in which fungi were detected in UK and US surveys (Anon., 1996; Kelley *et al.*, 2001)

<i>Fungus</i>	<i>USA (23 sites sampled)</i>	<i>UK (20 sites sampled)</i>	<i>Where present in treated water samples</i>	<i>Classification</i>
<i>Absidia cylindrospora</i>		3		Zygomycetes
<i>Absidia glauca</i>	2	3		Zygomycetes
<i>Acremoniella atra</i>	1			Ascomycetes
<i>Acremonium cf. strictum</i>	2			Ascomycetes
<i>Acremonium simplex</i>		1	UK	Ascomycetes
<i>Acremonium spp</i>	5	5	UK + USA	Ascomycetes
<i>Actinomucor elegans</i>	1			Zygomycetes
<i>Alternaria alternata</i>	12	3	UK + USA	Ascomycetes
<i>Alternaria infectoria</i>		2		Ascomycetes
<i>Alternaria spp</i>	1	1	USA	Ascomycetes
<i>Apiospora montagnei</i>	3	1		Ascomycetes
<i>Arthrinium phaeospermum</i>	1	4		Ascomycetes
<i>Arthrographis cuboidea</i>	1	2		Ascomycetes
<i>Ascochyta spp</i>	2	3	UK	Ascomycetes
<i>Aspergillus aculeatus</i>		2		Ascomycetes
<i>Aspergillus candidus</i>	1			Ascomycetes
<i>Aspergillus clavatus</i>		3		Ascomycetes
<i>Aspergillus flavus</i>		5	UK	Ascomycetes
<i>Aspergillus fumigatus</i>		3	UK	Ascomycetes
<i>Aspergillus japonicus</i>		1		Ascomycetes
<i>Aspergillus niger</i>	20	1	USA	Ascomycetes
<i>Aspergillus parvulus</i>		1		Ascomycetes
<i>Aspergillus spp</i>	1			Ascomycetes
<i>Aspergillus terreus</i>		2	UK	Ascomycetes
<i>Aspergillus versicolor</i>	6	3	UK + USA	Ascomycetes
<i>Asteroma spp</i>		1		Ascomycetes
<i>Asteromella spp</i>		1		Ascomycetes
<i>Aureobasidium pullulans</i>	2	1	UK	Ascomycetes
Basidiomycete	3		USA	Basidiomycetes
<i>Beauvaria bassiana</i>	2	1		Ascomycetes
<i>Beauvaria brongniartii</i>		1		Ascomycetes
<i>Beauvaria spp</i>	1			Ascomycetes
<i>Botrytis cinerea</i>		8	UK	Ascomycetes
<i>Chaetocladium brefeldii</i>		1		Zygomycetes
<i>Chaetomium bostrychodes</i>	1			Ascomycetes
<i>Chaetomium globosum</i>	1	1	UK	Ascomycetes
<i>Chaetomium robustum</i>	1			Ascomycetes
<i>Chaetomium spp</i>	2		USA	Ascomycetes
<i>Chaetomium sphaerale</i>	1			Ascomycetes
<i>Chaetomium sulphureum</i>	1			Ascomycetes
<i>Cladorrhinum spp</i>	2			Ascomycetes
<i>Cladosporium cf. herbarum</i>	1			Ascomycetes
<i>Cladosporium cladosporioides</i>	14	11	UK + USA	Ascomycetes
<i>Cladosporium herbarum</i>	20	13	UK + USA	Ascomycetes
<i>Cladosporium oxysporum</i>	16		USA	Ascomycetes
<i>Cladosporium spp</i>	8	4	UK + USA	Ascomycetes
<i>Cladosporium sphaerospermum</i>	8	1	UK + USA	Ascomycetes
<i>Cladosporium tenuissimum</i>	7		USA	Ascomycetes
<i>Cochliobolus pallescens</i>	1			Ascomycetes
<i>Colletotrichum spp</i>	1			Ascomycetes
<i>Cryptococcus albidus</i>	1			Basidiomycetes
<i>Cryptococcus spp</i>	1			Basidiomycetes

TABLE 5.2 (continued)

Fungus	USA (23 sites sampled)	UK (20 sites sampled)	Where present in treated water samples	Classification
<i>Cyclothyrium</i> spp		1		Ascomycetes
<i>Cylindrocarpon magnusianum</i>	1	1		Ascomycetes
<i>Cytospora</i> spp	1	4		Ascomycetes
<i>Discosporium</i> spp		1		Ascomycetes
<i>Embellisia</i> spp		1		Ascomycetes
<i>Epicoccum nigrum</i>	13	4	USA	Ascomycetes
<i>Eupenicillium</i> spp		1		Ascomycetes
<i>Exophiala jeanselmei</i>		1	UK	Ascomycetes
<i>Fusarium acuminatum</i>	1			Ascomycetes
<i>Fusarium aquaeductuum</i>		1		Ascomycetes
<i>Fusarium avenaceum</i>	1	6		Ascomycetes
<i>Fusarium</i> cf. <i>oxysporum</i>		2	UK	Ascomycetes
<i>Fusarium</i> cf. <i>solani</i>	1			Ascomycetes
<i>Fusarium culmorum</i>		8		Ascomycetes
<i>Fusarium diamini</i>		1	UK	Ascomycetes
<i>Fusarium equiseti</i>	4	1		Ascomycetes
<i>Fusarium flocciferum</i>		1		Ascomycetes
<i>Fusarium graminearum</i>	7		USA	Ascomycetes
<i>Fusarium moniliforme</i>		1		Ascomycetes
<i>Fusarium oxysporum</i>	2	3	UK + USA	Ascomycetes
<i>Fusarium pallidoroseum</i>	2			Ascomycetes
<i>Fusarium soecheri</i>	2			Ascomycetes
<i>Fusarium solani</i>	1	2	UK	Ascomycetes
<i>Fusarium</i> spp	2	3		Ascomycetes
<i>Fusarium sporotrichioides</i>	2		USA	Ascomycetes
<i>Fusarium torulosum</i>		1		Ascomycetes
<i>Geomyces pannorum</i>	1			Ascomycetes
<i>Geotrichum candidum</i>	3	4		Ascomycetes
<i>Geotrichum</i> spp	3	1		Ascomycetes
<i>Gliocladium roseum</i>	1	3	UK	Ascomycetes
<i>Gliocladium</i> spp		1		Ascomycetes
<i>Gongronella butleri</i>		1		Zygomycetes
<i>Hormonema dematioides</i>	1	1		Ascomycetes
<i>Hormonema</i> spp	1			Ascomycetes
<i>Idriella bolleyi</i>	4			Ascomycetes
<i>Idriella</i> sp.	1			Ascomycetes
<i>Leptodontidium</i> cf. <i>elatus</i>	1			Ascomycetes
<i>Leptodontidium elatus</i>	1		USA	Ascomycetes
<i>Leptodontidium</i> spp	3	1		Ascomycetes
<i>Leptodothiorella</i> spp		1		Ascomycetes
<i>Leptosphaeria coniothyrium</i>		2		Ascomycetes
<i>Leptosphaeria fuckelii</i>		2		Ascomycetes
<i>Mauginiella</i> spp		2	UK	Basidiomycetes
<i>Microdochium nivale</i>	1			Ascomycetes
<i>Microdochium</i> spp	1		USA	Ascomycetes
<i>Micromucor isabellina</i>		1		Zygomycetes
<i>Micromucor ramannianus</i>		3		Zygomycetes
<i>Microsphaeropsis olivacea</i>	2	2	USA	Ascomycetes
<i>Microsphaeropsis</i> spp		1		Ascomycetes
<i>Mortierella alpina</i>	1	1	UK	Zygomycetes
<i>Mortierella elongata</i>		1	UK	Zygomycetes
<i>Mortierella zychae</i>		1		Zygomycetes
<i>Mucor circinelloides</i>	5	3	USA	Zygomycetes
<i>Mucor circinelloide</i> sf. <i>janssenii</i>		2		Zygomycetes

(continued on next page)

TABLE 5.2 (continued)

Fungus	USA (23 sites sampled)	UK (20 sites sampled)	Where present in treated water samples	Classification
<i>Mucor flavus</i>	1			Zygomycetes
<i>Mucor fuscus</i>		1		Zygomycetes
<i>Mucor genevensis</i>	1			Zygomycetes
<i>Mucor hiemalis</i>	6	11	UK + USA	Zygomycetes
<i>Mucor mucedo</i>	1			Zygomycetes
<i>Mucor plumbeus</i>		1		Zygomycetes
<i>Mucor racemosus</i> f. <i>racemosus</i>	5	5		Zygomycetes
<i>Mucor racemosus</i> f. <i>sphaerosporus</i>		2		Zygomycetes
<i>Mucor</i> spp	4	3		Zygomycetes
<i>Mucor strictus</i>		1		Zygomycetes
<i>Myrioconium</i> spp	1			Ascomycetes
<i>Nectria ingoluensis</i>	1			Ascomycetes
<i>Neurospora</i> spp		1		Ascomycetes
<i>Nigrospora sphaerica</i>	2			Ascomycetes
Oomycete		1	UK	Oomycetes
<i>Paecilomyces lilacinus</i>	2	1		Ascomycetes
<i>Penicillium aurantiogriseum</i>	7	5	USA	Ascomycetes
<i>Penicillium brevicompactum</i>	8	9	USA	Ascomycetes
<i>Penicillium chrysogenum</i>	17	14	UK + USA	Ascomycetes
<i>Penicillium citrinum</i>	6	12	UK + USA	Ascomycetes
<i>Penicillium corylophilum</i>	1	2	UK	Ascomycetes
<i>Penicillium crustosum</i>		1		Ascomycetes
<i>Penicillium echinulatum</i>	1	1		Ascomycetes
<i>Penicillium expansum</i>	3	5	USA	Ascomycetes
<i>Penicillium funiculosum</i>	1			Ascomycetes
<i>Penicillium glabrum</i>	9	4	USA	Ascomycetes
<i>Penicillium griseofulvum</i>	1	1		Ascomycetes
<i>Penicillium herquei</i>	2			Ascomycetes
<i>Penicillium hirsutum</i>	1	1		Ascomycetes
<i>Penicillium janczewskii</i>		11	UK	Ascomycetes
<i>Penicillium minioluteum</i>	1	1	UK + USA	Ascomycetes
<i>Penicillium ochrosalmoneum</i>		1		Ascomycetes
<i>Penicillium oxalicum</i>	10		USA	Ascomycetes
<i>Penicillium paxilli</i>	1		USA	Ascomycetes
<i>Penicillium pinophilum</i>	3	1	USA	Ascomycetes
<i>Penicillium purpurogenum</i>		10	UK	Ascomycetes
<i>Penicillium raistrickii</i>		2		Ascomycetes
<i>Penicillium roquefortii</i>	1		USA	Ascomycetes
<i>Penicillium simplicissimum</i>	2	2		Ascomycetes
<i>Penicillium solitum</i>		1		Ascomycetes
<i>Penicillium</i> spp		1		Ascomycetes
<i>Penicillium spinulosum</i>	6	7	UK + USA	Ascomycetes
<i>Penicillium thomii</i>	3		USA	Ascomycetes
<i>Penicillium waksmanii</i>		2		Ascomycetes
<i>Periconia byssoides</i>	1			Ascomycetes
<i>Periconiella</i> spp		1		Ascomycetes
<i>Pestalotiopsis guepinii</i>	1		USA	Ascomycetes
<i>Phialophora</i> cf. <i>malorum</i>	1		USA	Ascomycetes
<i>Phialophora fastigiata</i>		5	UK	Ascomycetes
<i>Phialophora malorum</i>	1		USA	Ascomycetes
<i>Phialophora</i> spp	4		USA	Ascomycetes
<i>Phoma eupyrena</i>	1			Ascomycetes
<i>Phoma exigua</i>		3	UK	Ascomycetes
<i>Phoma fimeti</i>	1			Ascomycetes

TABLE 5.2 (continued)

Fungus	USA (23 sites sampled)	UK (20 sites sampled)	Where present in treated water samples	Classification
<i>Phoma glomerata</i>	5		USA	Ascomycetes
<i>Phoma herbarum</i>	1			Ascomycetes
<i>Phoma jolyana</i>		2	UK	Ascomycetes
<i>Phoma leveillei</i>	1	5	UK	Ascomycetes
<i>Phoma macrostoma</i>	1	2	UK	Ascomycetes
<i>Phoma medicaginis</i>	1	1		Ascomycetes
<i>Phoma nebulosa</i>	1	2		Ascomycetes
<i>Phoma</i> sect. <i>Phyllostictoides</i>		1		Ascomycetes
<i>Phoma</i> spp	16	12	UK + USA	Ascomycetes
<i>Phoma tropica</i>	2			Ascomycetes
<i>Phomopsis</i> spp		3	UK	Ascomycetes
<i>Pichia anomala</i>	1		USA	Saccharomycetes
<i>Pichia</i> spp	1			Saccharomycetes
<i>Pilidium concavum</i>		2		Ascomycetes
<i>Pithomyces chartarum</i>	7		USA	Ascomycetes
<i>Pithomyces sacchari</i>	1	1		Ascomycetes
<i>Preussia fleischakii</i>	1			Ascomycetes
<i>Pseudeurotium zonatum</i>		1		Ascomycetes
<i>Pycnidophora dispersa</i>		1		Ascomycetes
<i>Pyrenochaeta</i> spp		1		Ascomycetes
<i>Rhizopus oryzae</i>	2			Zygomycetes
<i>Rhizopus stolonifer</i>	1	9	UK	Zygomycetes
<i>Robillarda sessilis</i>	1			Ascomycetes
<i>Saprolegnia ferax</i>		1		Oomycetes
<i>Saprolegnia parasitica</i>		3		Oomycetes
<i>Saprolegnia subterranea</i>		1		Oomycetes
<i>Scopulariopsis acremonium</i>		1		Ascomycetes
<i>Scopulariopsis brevicaulis</i>		2		Ascomycetes
<i>Scopulariopsis</i> spp	1		USA	Ascomycetes
<i>Seimatosporium lichenicola</i>		1		Ascomycetes
<i>Sesquicillium</i> spp	1		USA	Ascomycetes
<i>Sordaria fimicola</i>	1			Ascomycetes
<i>Sporothrix</i> spp		2	UK	Ascomycetes
<i>Sporotrichum pruinosum</i>		1		Basidiomycetes
<i>Stereum</i> spp		1	UK	Basidiomycetes
<i>Syncephalastrum racemosum</i>		2		Zygomycetes
<i>Trichoderma harzianum</i>	9	11	UK + USA	Ascomycetes
<i>Trichoderma koningii</i>	3	7		Ascomycetes
<i>Trichoderma piluliferum</i>	3			Ascomycetes
<i>Trichoderma polysporum</i>	1	2		Ascomycetes
<i>Trichoderma pseudokoningii</i>	1	4		Ascomycetes
<i>Trichoderma</i> spp	5	2		Ascomycetes
<i>Trichoderma strigosum</i>	1			Ascomycetes
<i>Trichosporiella sporotrichoides</i>	1			Ascomycetes
<i>Trichosporon</i> spp	1			Ascomycetes
<i>Truncatella angustata</i>		1	UK	Ascomycetes
<i>Ulocladium atrum</i>		1		Ascomycetes
<i>Ulocladium botrytis</i>		1		Ascomycetes
<i>Verticillium lecanii</i>	1			Ascomycetes
<i>Verticillium</i> sp.		1		Ascomycetes
Zygomycete	2			Zygomycetes
<i>Zygorhynchus heterogamus</i>	4			Zygomycetes
<i>Zygorhynchus moelleri</i>	4	3		Zygomycetes

(continued on next page)

TABLE 5.2 (continued)

Fungus	USA (23 sites sampled)	UK (20 sites sampled)	Where present in treated water samples	Classification
Number of species present	UK = 136	USA = 141	Total species = 213	
Number of species in common	64		Species in common/total species = 0.3	

methods likely to detect them, but there is no reason to discount the possibility of their presence.

Fungi that are usually regarded as aquatic are often grouped informally by habitat preference and the nature of the spores.

3.2.1 Aquatic hyphomycetes

This is a grouping of asexual states of fungi which typically occur on submerged leaf litter and woody debris in rapidly flowing, well-aerated, non-polluted streams, and produce water-dispersed spores of elaborate appearance. It is thought the spore forms are adaptations to increase the likelihood of attaching to underwater surfaces in flowing waters (Dix and Webster, 1995). Sampling techniques for these fungi include filtration and baiting methods (Shearer and Lane, 1983; Shearer and Webster, 1985). The connected sexual states are ascomycetes and some basidiomycetes; these often occur on exposed plant debris near to streams (Webster, 1992; Shearer, 1993). There has been considerable interest in the ecology of aquatic hyphomycetes and it is known that they have a significant role, interacting with invertebrates, in the degradation of leaf litter and similar debris in streams and other bodies of water (Suberkropp, 1991; Barlocher, 1992; Suberkropp *et al.*, 1993; Dix and Webster, 1995; Bermingham, 1996; Gessner *et al.*, 1997).

3.2.2 Aero-aquatic hyphomycetes

These are asexual states of fungi found on plant debris in still and slow-flowing habitats, typically at the edges of ponds and ditches where sporulation occurs on newly-exposed leaf litter which is still moist but exposed to air (Dix and Webster, 1995). The conidia are complex, often helicoid, trapping air as an aid to buoyancy and hence dispersal. As with the aquatic hyphomycetes, the connected sexual states are ascomycetes and basidiomycetes. These fungi grow best under aerobic conditions but can survive prolonged periods of anaerobic conditions and of drought, suggesting that these may be among the most significant decomposer fungi in waters prone to fluctuating levels. Isolation methods include baiting and damp-chamber incubation of litter (Fisher, 1977).

3.2.3 Zoosporic fungi

Filamentous fungi producing zoospores belong to the Oomycetes (sometimes referred to as 'water moulds') and include well-known genera such as the plant-pathogenic genera *Pythium* and *Phytophthora*, pathogens of aquatic vertebrates such as species of *Achlya* and *Saprolegnia*, as well as members of the 'sewage fungus' complex such as *Leptomitius lacteus* and allied genera such as the fermentative *Aquilinderella*. Methods for isolation and study have

TABLE 5.3 Classification of fungi detected in UK and US water surveys (Anon., 1996; Kelley *et al.*, 2001)

UK and US survey data combined					
Ascomycetes	Basidiomycetes	Oomycetes	Saccharomycetes	Zygomycetes	Total
173	6	4	2	28	213
81	3	2	1	13	%

recently been discussed by Hall (1996) and Dix and Webster (1995) and involve specialized baiting and other techniques in order to detect their presence.

There are other fungi-like organisms, such as chytrids, that also produce zoospores but these are non-filamentous and are not considered here (for references see Kirk *et al.*, 2001).

3.2.4 Other fungi from water

In some cases fungi are still found which are not easily related to existing descriptions (e.g. Willoughby, 2001), indicating that there is continuing scope for discovery of new fungi in aquatic environments, subject to the development of sampling and culture techniques.

3.3 Structure of populations

The above-mentioned uncertainties surrounding the enumeration of fungi make it difficult to describe mixed fungal populations quantitatively. Often it is only possible to indicate which fungal species are detected; however, as more surveys are completed, which provide detailed species information, it becomes possible to gain an idea of which species and genera are most frequently seen. Tables 5.2 and 5.3 show the fungi detected in two different surveys and, in particular, highlight those fungi present in both the US and UK post-treatment water samples. These include: *Acremonium* spp., *Alternaria alternata*, *Aspergillus versicolor*, *Cladosporium cladosporioides*, *C. herbarum*, *Cladosporium* spp., *C. sphaerospermum*, *Fusarium oxysporum*, *Mucor hiemalis*, *Penicillium chrysogenum*, *P. citrinum*, *P. minioluteum*, *P. spinulosum*, *Phoma* spp. and *Trichoderma harzianum*. While this list is unlikely to be exhaustive, it is interesting to note, that although these fungi belong to large multispecies genera, only a small subset of the known species for each genus is represented. In many cases therefore, species-level differences may be more important than generic or higher level differences in determining persistent or frequent occurrence in water systems.

TABLE 5.4 Species numbers in samples from UK and US surveys (Anon., 1996; Kelley *et al.*, 2001)

Country	Mean number of species present per sample \pm standard error	
	Untreated water	Treated water
UK	30.7 \pm 1.1 (n = 10)	8.5 \pm 1.2 (n = 10)
USA	29.6 \pm 5.1 (n = 10)	9.9 \pm 1.1 (n = 13)

3.3.1 Comparing samples

It is often required to analyse the diversity of species present. Samples can be compared in terms of overall numbers of species present, as exemplified in Table 5.4, which shows that samples of treated water in both the US and UK surveys contain approximately one third of the numbers of untreated waters. In order to compare species composition, samples can be analysed in terms of the proportion of the species in common between sites relative to the total number of species detected for both sites; comparing the US and UK species shown in Table 5.2, this ratio is 0.3 (sometimes known as the Jaccard coefficient; see Anon., 1990). Given that the US and UK surveys yielded 141 and 136 species respectively (and the similarity of these numbers of species is also worth noting), this indicates a moderately high overlap with just under 50% of all species present in the US survey also appearing in the UK survey and *vice versa*. There are other numerical treatments, which include weightings for frequency or abundance of individual species, such as calculating species richness estimators (Dighton, 1994; Colwell, 1997), but with fungi the difficulties of enumeration often render this an uncertain exercise.

3.4 Environmental factors

Dix and Webster (1995) and Magan (1997) have reviewed the tolerance of fungi to extreme environmental conditions. Many of the fungi detected in water systems are likely to be allochthonous, with the majority transiently passing through as spores. In which case, the main concern is simply in the ability of the fungi to withstand a range of conditions in

order to maintain viability until subsequent deposition. However, there is also an indication that fungi can grow in water systems (Kelley *et al.*, 1997, 2001), and in this case it is important to appreciate the range of conditions that are favourable for fungal growth.

3.4.1 Habitats and niches within distribution systems

Fungi have been reported from all types of water systems from raw waters to bottled drinking waters and from heavily polluted waters to distilled or ultra-pure water (Kelley *et al.*, 1997). They may form floating mats of thin finely branched hyphae in static water, or may form pelleted colonies in sustained circular flows. They may also grow attached to a substrate, forming part of microbial biofilms on pipework inner surfaces, debris or sediments, and are particularly likely to become established where there are cracks, pitting or dead ends.

3.4.2 Temperature

The majority of fungi are mesophilic, capable of growth between 5 and 35 °C, with the optimum often between 25 and 30 °C (Smith and Onions, 1994; Dix and Webster, 1995; Magan, 1997), which covers the temperatures found in most water systems including the tropics (Yuen *et al.*, 1998). In terms of extremes, there are psychrotolerant fungi, such as some species of *Cryptococcus*, which are capable of growth at or close to 0°C. Psychrophilic fungi, including some species of *Mucor*, are unable to grow above 20°C. However, many fungi are capable of withstanding temperatures down to 0°C or lower, with survival depending on the cell water content and the rate of freezing (Smith and Onions, 1994); generally spores survive better than hyphae. In terms of upper temperatures, the maximum for growth of thermotolerant fungi, such as *Aspergillus fumigatus*, is in the region of 50°C and for some thermophiles, such as *Thermomyces lanuginosus*, approximately 60°C. These thermotolerant and thermophilic species have been reported from heated aquatic habitats such as cooling waters at power stations (Ellis, 1980;

Novickaja–Markovskaja, 1995). The majority of fungi are readily killed by temperatures at or above 80 °C, although a few such as *Talaromyces flavus* can withstand this temperature for an hour; survival again depends on the cell water content, with spores surviving better than hyphae (Magan, 1997).

3.4.3 pH

Fungi have been detected in extremely acidic and alkaline waters and many fungi when tested can grow over a wide range of pH, often between 2 and 11, although how this correlates to the ability to grow in environments subject to extremes of pH is in many cases still unclear, as other environmental factors are often closely linked (Magan, 1997). An example is the River Tinto in Spain which has been associated with mining for over 5000 years; it presently has a pH of 2–2.5 and contains ferric iron at 2.5 g/l and other heavy metals including Cu, Zn, Co, Mn, Ni as well as sulphate at 10–15 g/l and has high redox potential and conductivity. Nonetheless, a study has yielded 350 fungal isolates belonging to *Bahusakala*, *Heteroconium*, *Mortierella*, *Penicillium*, *Phialophora* (as *Lecythophora*), *Phoma* and *Scytalidium*. Of the isolates, 44% were capable of growth in the conditions present in the river (López–Archilla *et al.*, 1996).

3.4.4 Aeration

The majority of fungi are aerobic with a wide range of genera showing the capacity to grow under low oxygen conditions (Dix and Webster, 1995; Magan, 1997). While fungal microaerophilic capabilities have been studied in soils and plant materials it appears that there have been no studies of fungi in water systems in which the oxygen levels have been accurately measured (Kelley *et al.*, 1997). Many fungi, provided appropriate nutrient sources are present (such as fatty acids, sterols and vitamins), are facultative anaerobes. This includes species of *Fusarium*, *Mucor*, *Penicillium* and *Trichoderma* as well as *Saccharomycetes*. Other facultatively anaerobic fungi found growing in stagnant waters include members of the Oomycete genera *Aqualinderella*, *Mindeniella*, *Rhipidium*

and *Sapromyces*. In the case of *Aqualinderella fermentans*, it is incapable of aerobic respiration. Interestingly, the related *Leptomitius lacteus*, although found in heavily polluted waters, is not a facultative anaerobe. The aero-aquatic hyphomycetes mentioned earlier are capable of surviving prolonged anaerobic conditions.

3.4.5 Nutrients

Many fungi have the ability to grow in apparently nutrient-free conditions, including unamended water, scavenging nutrients from whatever sources that may be available, however dilute or recalcitrant, even the atmosphere (Parkinson *et al.*, 1989; Wainwright *et al.*, 1992, 1993; Kelley *et al.*, 1997, 2001; Gharieb and Gadd, 1999; Karunasena *et al.*, 2000). The utilization of simple sugars as C-sources occurs in virtually all fungi. Many are also well adapted to make use of more complex C polymers of plant and animal origin, especially cellulose-based materials (see Gessner *et al.*, 1997). The main exceptions are many of the Zygomycetes and saprobic Oomycetes, which appear to be restricted to simple sugars. In terms of N utilization, many fungi can grow over a wide range of concentrations and on a wide range of N-sources. Some show adaptations to high N levels, as in the case of coprophilous fungi, or to very low levels as in wood-decay fungi (Dix and Webster, 1995).

3.4.6 Flow rates

For those fungi transiently passing through a system, water flow rate is largely immaterial, although the consequent length of time spent in the system may affect viability. Whether attached to surfaces or otherwise, many other fungi appear to grow equally in either standing or moving water (Kelley *et al.*, 1997, 2001). Extremely high flow rates or turbulence could conceivably cause cellular damage or inhibit colonization, but there is a lack of detailed information for fungi. Aquatic hyphomycetes favour flowing water over standing water and there is an indication that spore shape may in some species be an adaptation to assist anchorage in such conditions (Webster and Davey, 1984).

3.4.7 Light

Growth of many species occurs equally in the dark or in daylight. Near-ultraviolet light of wavelengths between 300 and 380 nm is routinely used to induce sporulation in many fungi, while others respond better to darkness or to diurnal cycles (Smith and Onions, 1994). Ultraviolet light has also been used as a treatment against fungi in water (see references in Kelley *et al.*, 1997).

3.4.8 Pollution and global environmental change

Wainwright and Gadd (1997) have reviewed the effects of industrial pollutants on fungi, particularly acid rain, heavy metals and radionuclides. Apart from the well known 'sewage fungus' complex (the component organisms include unrelated fungi such as *Leptomitius lacteus* and species of *Fusarium* and *Geotrichum*) found in low pH water heavily contaminated with organic matter, much of the other data concerns terrestrial fungi, especially mycorrhizal species and those which produce large fruitbodies. While there is a strong indication that many fungi are capable of accumulating pollutants, the distinction between survival and tolerance is not always clear, nor yet are the longer-term effects of either toxicity or eutrophication on the composition of fungal communities understood. For example, Bermingham (1996) has observed aquatic hyphomycete communities in a river subject to contamination by heavy metals. An effect of the contamination was a reduction in leaf litter decomposition. Experimental tests showed that sporulation was inhibited by the metals but mycelial growth continued. Since identification of aquatic hyphomycetes is reliant on determination of spore morphologies, it is therefore unclear if the change in community function actually represented a loss of species.

4 CLEANING AND DISINFECTION TREATMENTS

The effectiveness of disinfection treatments in removing fungi from water has been reviewed by Kelley *et al.* (1997). The most commonly used treatment is chlorination, to which fungal

spores are more resistant than hyphae. It was suggested that an initial residual level of 1 mg/l chlorine is required for use against fungi. However, they also pointed out examples showing that much higher concentrations can be required for control of certain species such as 129 mg/l for *Botrytis cinerea*. The survey data shown in Table 5.2 indicate that some fungi can pass through treatment systems into distribution systems.

Kelley *et al.* (2001) examined the effectiveness of different treatment strategies to control fungi. This included physical treatments (clarification, rapid gravity filtration) and chemical disinfection (monochloramine, chlorine, chlorine dioxide and ozone), as well as mains cleaning (flushing, air-scouring, swabbing, pigging and flow-jetting). They concluded that physical treatments performed well, with clarification removing over 70% of fungi, while filtration removed over 90% of fungi, irrespective of whether sand or granular activated carbon was used. Efficacy of chemical disinfectants varied from species to species but, overall, chlorine dioxide and ozone were the most powerful of those tested. However, monochloramine appeared more stable in distribution systems and may therefore have a greater long-term efficacy. It was also found that, of the disinfectants, chlorine was the most affected by variation in temperature. Of the mains cleaning treatments, none gave particularly good results. Flushing and air-scouring were least effective, albeit also among the least damaging to the pipe. Swabbing only gave good results on smooth-surfaced plastic pipes but not on rough-surfaced cast iron pipes. Pigging, although very aggressive, also gave poor results due to the limited contact that can be achieved between the bristles and the pipe surface. Flow-jetting was reported as the most effective and was not affected by pipe materials.

5 SIGNIFICANCE OF FUNGAL ACTIVITIES

5.1 Biofilms and biofouling

There have been very few reports of fungi in biofilms (Kelley *et al.*, 1997). In many cases, this

is probably a consequence of the use of methods which do not target fungi rather than an absence of fungi. Nagy and Olson (1985) reported fungi belonging to *Acremonium*, *Alternaria*, *Epicoccum*, *Candida*, *Cryptococcus*, *Penicillium*, *Rhodotorula* and other species from pipe surfaces. More recently, Dogget (2000) reported fungi from biofilms in a municipal water distribution system, including members of *Acremonium*, *Alternaria*, *Aspergillus*, *Aureobasidium*, *Candida*, *Cladosporium*, *Cryptococcus*, *Mucor*, *Penicillium*, *Phoma* and *Stachybotrys* as well as other fungal species. In addition, during the surveys reported by Kelley *et al.* (2001), swab samples from pipes at sites of reported taste and odour incidents yielded fungi belonging to *Aspergillus*, *Cladosporium*, *Cryptococcus*, *Pichia*, *Rhodotorula* and *Yarrowia*. Fungi, such as *Aspergillus*, *Fusarium*, *Mucor*, *Penicillium* and *Trichoderma* species, have been recovered from biofouled membranes of reverse osmosis systems (Dudley and Christopher, 1999). At present there is little information on the role of fungi in causing the formation and stabilization of biofilms in water systems.

5.2 Biodegradation and bioremediation

There are many proposed strategies for biological treatment of wastewater and run-off including constructed wetlands such as reedbeds (Haberl *et al.*, 1997). Fungi are widely recognized as being important organisms for many biodegradative and bioremediative processes (Gadd *et al.*, 1987; Wainwright *et al.*, 1987; Frankland *et al.*, 1996), but there is a surprising lack of information on their activities or potential for use in treatment of water. For example, one of the main macrophytes used in reedbeds is *Phragmites* and about one hundred fungal species have been recorded in, on or around *Phragmites* worldwide (as shown in the databases of the herbaria of CABI Bioscience, incorporating the former International Mycological Institute, and the Royal Botanic Gardens Kew). With such a known list of associated fungal species it is difficult to

TABLE 5.5 Compounds produced by fungi which cause off-tastes or odours (from Kelley *et al.*, 1997a,b)

Fungus	Compound produced	Relative production	Reference
<i>Acremonium</i> spp ¹	oct-1-en-3-ol	major	Kaminski <i>et al.</i> (1980)
<i>Alternaria</i> spp	oct-1-en-3-ol	major	Kaminski <i>et al.</i> (1980)
<i>Aspergillus</i> spp	2-methylisoborneol	minor	Anderson <i>et al.</i> (1995)
<i>Aspergillus niger</i>	oct-1-en-3-ol	major	Kaminski <i>et al.</i> (1980)
<i>Aspergillus ocraceus</i>	oct-1-en-3-ol	major	Kaminski <i>et al.</i> (1980)
<i>Aspergillus oryzae</i>	oct-1-en-3-ol	major	Kaminski <i>et al.</i> (1980)
<i>Aspergillus parasiticus</i>	oct-1-en-3-ol	minor	Kaminski <i>et al.</i> (1980)
<i>Basidiobolus ranarum</i>	Geosmin	no data	Lechevalier (1974)
<i>Botrytis cinerea</i>	furfural, benzaldehyde, phenylacetaldehyde, benzyl cyanide	no data	Kikuchi <i>et al.</i> (1983)
<i>Chaetomium globosum</i>	geosmin, 2-phenylethanol	no data	Kikuchi <i>et al.</i> (1983)
<i>Cladosporium</i> spp	2-methylisoborneol	major	Anderson <i>et al.</i> (1995)
<i>Fusarium</i> spp	oct-1-en-3-ol	major	Kaminski <i>et al.</i> (1980)
<i>Geotrichum</i> spp	2,4,6-TCA	minor	Anderson <i>et al.</i> (1995)
<i>Mucor</i> spp	2,4,6-TCA/2,3,4-TCA	minor	Anderson <i>et al.</i> (1995)
<i>Penicillium aethiopicum</i>	Geosmin	minor	Larsen and Frisvad (1995)
<i>Penicillium aurantiogriseum</i>	oct-1,3-diene	minor	Larsen and Frisvad (1995)
<i>Penicillium camembertii</i>	2-methylisoborneol, oct-1-en-3-ol	minor, minor	Larsen and Frisvad (1995)
<i>Penicillium chrysogenum</i>	oct-1,3-diene, oct-1-en-3-ol	minor, minor	Larsen and Frisvad (1995), Kaminski <i>et al.</i> (1980)
<i>Penicillium citrinum</i>	oct-1-en-3-ol	major	Kaminski <i>et al.</i> (1980)
<i>Penicillium clavigerum</i>	Geosmin	major	Larsen and Frisvad (1995)
<i>Penicillium commune</i>	2-methylisoborneol, oct-1,3-diene, oct-1-en-3-ol	minor, minor, minor	Larsen and Frisvad (1995)
<i>Penicillium coprobium</i>	oct-1,3-diene	minor	Larsen and Frisvad (1995)
<i>Penicillium crustosum</i>	2-methylisoborneol, geosmin, dimethyldisulphide	minor, minor, minor	Larsen and Frisvad (1995)
<i>Penicillium decumbens</i>	oct-1-en-3-ol	not given	Halim <i>et al.</i> (1975)
<i>Penicillium discolor</i>	2-methylisoborneol, geosmin	major, major	Larsen and Frisvad (1995)
<i>Penicillium echinulatum</i>	Geosmin	minor	Larsen and Frisvad (1995)
<i>Penicillium expansum</i>	Geosmin	no data	Mattheis and Roberts (1992), Dionigi and Champagne (1995)
<i>Penicillium formosanum</i>	Geosmin	major	Larsen and Frisvad (1995)
<i>Penicillium freii</i>	oct-1,3-diene	minor	Larsen and Frisvad (1995)
<i>Penicillium funiculosum</i>	oct-1-en-3-ol	major	Kaminski <i>et al.</i> (1980)
<i>Penicillium glabrum</i>	oct-1,3-diene	minor	Larsen and Frisvad (1995)
<i>Penicillium hirsutum</i> var. <i>venetum</i>	Geosmin	minor	Larsen and Frisvad (1995)
<i>Penicillium polonicum</i>	2-methylisoborneol	minor	Larsen and Frisvad (1995)
<i>Penicillium raistricki</i>	oct-1-en-3-ol	major	Kaminski <i>et al.</i> (1980)
<i>Penicillium roqueforti</i> var. <i>carneum</i>	Geosmin	minor	Larsen and Frisvad (1995)
<i>Penicillium solitum</i>	2-methylisoborneol	minor	Larsen and Frisvad (1995)
<i>Penicillium</i> spp	2,4,6-TCA	no data	Nyström <i>et al.</i> (1992)
<i>Penicillium</i> sp. #1	2-methylisoborneol	minor	Anderson <i>et al.</i> (1995)
<i>Penicillium</i> sp. #2	2,4,6-TCA/2,3,4-TCA	minor	Anderson <i>et al.</i> (1995)
<i>Penicillium tricolor</i>	oct-1,3-diene, oct-1-en-3-ol	minor, major	Larsen and Frisvad (1995)
<i>Penicillium viridicatum</i>	oct-1-en-3-ol*	minor/major	Larsen and Frisvad (1995); Kaminski <i>et al.</i> (1980)
<i>Penicillium vulpinum</i>	2-methylisoborneol, oct-1,3-diene	minor, major	Larsen and Frisvad (1995)
<i>Phialophora</i> spp	2,4,6-TCA	no data	Nyström <i>et al.</i> (1992)
basidiomycete fungi	oct-1-en-3-ol, oct-1-en-3-one, 1,3-octadiene	high	Jüttner (1990)

¹ as *Cephalosporium* sp.

Note: Oct-1-en-3-ol has also been found in *Agaricus bisporus* and *Boletus edulis* (Kaminski *et al.*, 1980) but it is virtually certain that these fungi would not be found in drinking water.

imagine that fungi do not play a significant role in the functioning of reedbeds, or in other biological treatment systems.

5.3 Secondary metabolites

Fungi produce a wide range of secondary metabolites including toxic compounds (Moss, 1996; Scudamore, 1998), many of which have implications for human health (e.g. Richard *et al.*, 1999). Paterson *et al.*, (1997b) compiled a listing of the mycotoxins known for many of the fungi isolated from water samples. Fungi also produce many compounds liable to cause off-tastes and odours (Table 5.5). There is also concern over oestrogenic compounds and fungi, such as some species of *Fusarium*, are known to produce zearalanone, an oestrogenic compound (Aldridge and Tahourdin, 1998). Kelley *et al.* (2001) have demonstrated production of secondary metabolites by fungi in water including zearalanone by *Fusarium graminearum*, and have detected aflatoxin in a water storage tank contaminated by *Aspergillus flavus* by using immunoaffinity columns (see also Paterson *et al.*, 1997b).

5.4 Water as a source of fungal inoculum

The majority of fungi detected in water samples are plurivorous saprobes of decaying plant materials. Some of these, however, are also secondary/opportunistic pathogenic organisms of plants or animals, including humans (Hoog, 1996; Warnock and Campbell, 1996).

5.4.1 Pathogenic fungi

Human pathogenic fungi can be transported in water. Of those listed in Table 5.2, many are also reported as potential secondary pathogens (such as *Aspergillus* species, many of the Zygomycetes, as well as species of *Alternaria*, *Cladosporium*, *Phoma*, and others), for detailed listings and further references see Smith (1989). In most instances these fungi cause no problems for healthy adults, although there may be a potential risk for immunocompromised individuals. In critical

environments water can be a source of fungal infection. For example, Warris *et al.* (2001) examined water samples taken from the paediatric bone marrow transplantation (BMT) unit of the National Hospital University of Oslo, Norway; 168 water samples and 20 surface-related samples from taps and showers in the BMT unit and from the main pipe supplying the paediatric department were taken on 16 different days over an 8-month period. Filamentous fungi were recovered from 94% of all the water samples taken inside the hospital. *Aspergillus fumigatus* was recovered from 49% and 5.6% of water samples from the taps and showers respectively. More than one-third of water samples from the main pipe revealed *A. fumigatus* and it was suggested that the source of contamination was located outside the hospital.

Plant pathogenic fungi can also be transported in water, and many of those listed in Table 2.2 are also capable of acting as secondary plant pathogens (such as members of *Alternaria*, *Cladosporium*, *Fusarium*, *Phoma* and others). Untreated water used in irrigation systems may deliver an inoculum of potential plant pathogenic fungi directly to crop plants under ideal conditions for infection (i.e. creating wet conditions). Hydroponic systems are most vulnerable, but any irrigated system, especially in glasshouses, are also liable to be at heightened risk of disease unless appropriate measures are taken (Jarvis, 1992).

6 CONCLUSIONS

Fungi are a significant part of the microbial population in water systems of all types. Their activities influence both water quality and human health. At present there is a need better to understand and quantify the composition and significance of fungal populations.

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Microbial flora of the gut

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1 INTRODUCTION

The importance of the intestinal bacteria in the context of water and wastewater derives from the significance of water as a vehicle for the spread of infection. Most important in this context are intestinal infections causing diarrhoea.

Diarrhoea is a major symptom of many intestinal infections. Some gut pathogens may also invade the body from the gut and cause either generalized or localized infections. Such invasion is characteristic of typhoid and some other infections caused by salmonellae. During infections restricted to the gut, bacteria will be excreted in the faeces. When invasion has occurred, bacteria may be passed in the urine as well and can also be found in the bloodstream.

When considering the faeco-oral transmission of an intestinal pathogen by water it is seldom possible to demonstrate directly the pathogen in the water supply. The reasons for this are various but the relatively small number of a pathogen excreted and the limited time in which the excretion occurs are contributory factors. Cholera provides an instructive example of the problem. Patients with severe cholera will be excreting *Vibrio cholerae* in large numbers for a limited time, terminated by death or isolation. Patients with mild disease, or a convalescent carrier, by contrast may look relatively healthy and be mobile while still excreting vibrios in their faeces. Such mobile persons might be thought more likely to be a risk to water supplies, but it must be remembered that the numbers of *V. cholerae* excreted would be small.

A consequence of the vagaries of pathogen detection is the use of normal faecal bacteria as indicators of water pollution. These bacteria are ubiquitous and numerous in the faeces of healthy people. The most widely used indicator has been the faecal coliform *Escherichia coli*, but faecal enterococci are also used as indicators (see Chapter 7). Anaerobic bacteria also, such as *Clostridium*, *Bacteroides* and *Bifidobacterium*, have served as indicators, and each has potential value (Evison and James, 1977).

The bacterial microflora of humans is the most intimate portion of their biological environment and mediates many of their interactions with the chemical environment. This development of our knowledge of the composition of the intestinal microflora can be traced in a series of books (Drasar and Hill, 1974; Clarke and Bauchop, 1977; Hentges, 1983; Hill, 1995). However, an explanation of the effects of the gut bacteria on the body or of the mechanism by which they are controlled remains possible only in outline. Intestinal physiology and host defence mechanisms must play an important role in preventing the microflora from overrunning the host and in determining the distribution of the microflora within the intestine. It seems likely that environmental factors, such as diet, also influence the microflora. The ecological interactions between bacterial species are probably significant in the large intestine. However, their exact involvement is largely obscure.

2 THE STOMACH

The stomach receives bacteria from the mouth and the external environment. In the context of the faeco-oral transmission of disease it functions as a barrier to infection. The bactericidal action of gastric juice results from secretion of hydrochloric acid. Immediately after a test meal, about 10^5 bacteria per ml of gastric juice can be isolated but, as the pH falls, the bacterial count declines and few viable cells can be recovered after a pH of about 3 has been attained (Drasar *et al.*, 1969). Lactobacilli, which are particularly resistant to acid, persist longer than other bacteria. The achlorhydric stomach, whatever the cause, is usually heavily colonized by bacteria, indicating that gastric acid is an important factor in reducing the numbers of pathogenic bacteria entering the small bowel. Experimental evidence for this is afforded by studies of the infective dose of *Vibrio cholerae* in human volunteers, where neutralization of gastric acid with a solution of sodium bicarbonate enables an infection to be initiated by oral administration of a much smaller challenge dose compared to that needed for untreated subjects (Hornick *et al.*, 1971; Gianelia *et al.*, 1973).

3 THE SMALL INTESTINE

Although the restriction on bacterial proliferation in the small gut stems partly from the efficiency of gastric acid in reducing the bacterial load entering the bowel, small intestinal factors also play a role. Samples of small gut contents obtained from fasting European and North American residents contain few if any cultivable bacteria. The viable count seldom exceeds 10^4 per ml of digesta, but for a short period after a meal, an increased number of bacteria, that are probably transients from the mouth, can be isolated from jejunal samples. Although relatively few bacteria can be cultured, microscopy of jejunal juice reveals large numbers of cells, usually exceeding 10^6 per ml. The importance of these bacteria that can be seen but not grown has not yet been determined.

In healthy persons living in Western Europe and North America, the number of cultivable bacteria is lowest in the upper small bowel and greatest in the terminal ileum. Gram-positive species predominate among those bacteria isolated and those observed microscopically. Streptococci and lactobacilli are most often isolated, though bacteroides and enterobacteria may be found in small numbers. The organisms seen but not identified are Gram-positive rods and cocci.

These results differ markedly from data obtained during studies of the small intestinal flora in residents of developing countries which show that they have a richer and more permanent flora. The relationship of this flora to nutritional status, nutrient utilization and subclinical disease may have important practical consequences.

The flow rate of gut contents contributes to control of bacterial colonization, being greatest at the top of the small intestine where microbial multiplication usually does not exceed the rate at which organisms are removed. Animal experiments have shown that bacteria are cleared from the small intestine wrapped in intestinal mucus. Among the roles that have been suggested for intestinal antibody is the binding of bacteria to this material. The anatomically normal small intestine can only be colonized by adhesion to the epithelium, as occurs with some enteropathogenic serotypes of *E. coli*. In the ileum, water is absorbed, reducing the flow rate, which allows bacterial multiplication. The lower small intestine, the distal and terminal ileum, contain many more cultivable bacteria than the proximal ileum. Although lactobacilli and streptococci are still prominent, bacteroides and enterobacteria occur more constantly. In the terminal ileum viable bacterial counts of 10^5 – 10^7 per ml are not uncommon, and the flora here qualitatively begins to resemble that of faeces (Gorbach *et al.*, 1967).

4 THE LARGE INTESTINE

In the UK, the adult colon contains approximately 220 g of contents. Bacteria are a major component and about 18 g consist of bacterial

dry matter. Bacterial numbers, as estimated by direct microscope counts, increase progressively from the caecum to the rectum. However, most studies on the colonic flora are made on faeces, which can provide some useful information, since the few investigations where viable counts of bacteria were carried out with gut contents have indicated that, with respect to the major taxonomic groups, the gut flora is qualitatively similar to faeces (Bentley *et al.*, 1972). There is some evidence, however, for the existence of specific mucosa-associated populations, whose functional association with the mucus layer and intestinal wall may be of ecological and metabolic significance to the host (Croucher *et al.*, 1983).

In colonic contents and faeces, bacterial counts usually exceed 10^{11} per g dry weight. Several hundred different species are present, but some 30–40 species account for about 99% of the bacterial mass, and most of these are strict anaerobes (Finegold *et al.*, 1983). An important factor that facilitates the establishment of such a luxuriant flora is the increase in transit time of gut contents from about 3–4 hours in the small intestine to about 70 hours in the colon (Cummings, 1978).

Although bacteria are distributed throughout the gastrointestinal tract, the vast majority occurs in the large bowel. The major influences of the gut flora on host metabolism arise from bacterial metabolism of various substances in the colon. Of particular significance in the normal healthy bowel is the fermentation of proteins and carbohydrates. Much recent research has focused on the metabolic significance of the gut bacteria (Hill, 1995; Macfarlane and Cummings, 1999).

5 FAECAL BACTERIA

Many different types of bacteria, representing most bacterial groups, have at some time or other been isolated from faeces. About 400 species are thought to be present usually (Holdeman *et al.*, 1976). Those detected most frequently can be considered as members of the resident flora or as regular contaminants from the environment. The number of bacterial

groups that are found is related to the methods used for their isolation and characterization. Few researchers have ever attempted a systematic investigation to determine the composition of the faecal flora, and any list of species present must be incomplete. The scale of the problem, in terms of the numbers of bacteria to be isolated and identified, makes investigations based on classical bacteriological techniques problematic.

A few studies have relied on the identification of large numbers of isolates from non-selective media (Moore and Holdeman, 1974; Holdeman *et al.*, 1976). Other investigators have combined this approach with the use of selective media (Peach *et al.*, 1974; Finegold *et al.*, 1975, 1977). More recently the use of molecular techniques has rekindled interest in the problem and provide long-term solutions (Jansen *et al.*, 1999).

Table 6.1 shows the major groups of commonly isolated faecal bacteria. It can be seen that bacterial counts in different individuals range over several orders of magnitude. Most of the bacteria growing in the colon are non-sporing anaerobes and include members of the genera *Bacteroides*, *Bifidobacterium* and *Eubacterium* among many others. Clostridia are also represented, though they are outnumbered by the non-sporing anaerobes, as are facultative anaerobes, such as streptococci and enterobacteria. Quantitatively, the most important genera of intestinal bacteria in

TABLE 6.1 Numbers (\log_{10} /g dry wt) of bacteria reported in human faeces (Finegold *et al.*, 1983)

Bacteria	Description	Mean	Range
Bacteroides	Gram-negative rods	11.3	9.2–13.5
Eubacteria	Gram-positive rods	10.7	5.0–13.3
Bifidobacteria	Gram-positive rods	10.2	4.9–13.4
Clostridia	Gram-positive rods	9.8	3.3–13.1
Lactobacilli	Gram-positive rods	9.6	3.6–12
Ruminococci	Gram-positive cocci	10.2	4.6–12.8
Peptostreptococci	Gram-positive cocci	10.1	3.8–12.6
Propionibacteria	Gram-positive rods	9.4	4.3–12.0
Actinomyces	Gram-positive rods	9.2	5.7–11.1
Streptococci	Gram-positive cocci	8.9	3.9–12.9
Fusobacteria	Gram-negative rods	8.4	5.1–11.0
Escherichia	Gram-negative rods	8.6	3.9–12.3

animals and man are the *Bacteroides* and *Bifidobacterium*, which can account for up to 30 and 25% of the total anaerobic counts respectively (Macy and Probst, 1979; Mitsuoka, 1984; Scardovi, 1986). Among the Gram-positive non-sporing rods, several genera are numerically significant; these include *Eubacterium* and *Bifidobacterium*. Species such as *B. bifidum* and *B. infantis* have been isolated from the faeces of breast-fed infants but are probably less common in adults. The genus *Lactobacillus* contains many species that occur in the guts of most warm-blooded animals.

Several types of spore-forming rods and cocci are also normal inhabitants of the gut. The genus *Clostridium* is probably the most ubiquitous: *C. perfringens*, *C. bifermentans* and *C. tetani* are regularly isolated, albeit in relatively low numbers, from the lower gut of man and animals, and are of significance in human and veterinary medicine. The presence of aerobic members of the genus *Bacillus* is thought to result from contamination from the environment. Facultative and obligately anaerobic Gram-positive cocci are also numerically important. The strict anaerobes include *Peptostreptococcus*, *Ruminococcus*, *Megasphaera elsdenii* and *Sarcina ventriculi*. The facultatively anaerobic streptococci are well represented by many species from Lancefield group D, including *Enterococcus faecalis*, *S. bovis* and *S. equinus* and some from group K such as *S. salivarius*, which is usually associated with the mouth. Gram-negative anaerobic cocci include *Veillonella* and *Acidaminococcus*.

Although they are not numerous, the Gram-negative facultative anaerobic rods include a number of very important pathogens. Members of the Enterobacteriaceae, particularly *Escherichia coli*, are usually thought of as characteristic intestinal bacteria and have historically been the most important indicators of faecal pollution.

6 FAECAL BACTERIA AND INFECTION

Most of the bacteria isolated from faeces do not cause infection and in the present context are important only in so far as they act as surrogate

TABLE 6.2 Rank order of major bacterial indicators (Moore and Holdeman, 1974)

Rank	Species	Approximate number per gram faeces
37	<i>Escherichia coli</i>	10^7
64	<i>Enterococcus faecalis</i>	10^7
Unranked	<i>Clostridium perfringens</i>	10^5

indicators of infective hazard. This is not a straightforward relationship as there are few data on the relationship of the major faecal organisms to the numbers of pathogens or to traditional indicator organisms.

There have been very few systematic attempts to determine the numbers of a pathogen excreted by an infected person, however, in most cases the number of bacteria must be very many fewer than that contributed by the normal flora. A working estimate would be 10^7 – 10^9 pathogens per gram of faeces during acute infection. These numbers are very similar to those achieved by the usual indicator bacteria, but such organisms can be regarded, for practical purposes, as always present (Tables 6.1 and 6.2). Among the numerically dominant bacterial species (Table 6.3), only *Bifidobacterium adolescentis* has been subject to continuing investigation as an indicator (Resnick and Levin, 1981; Rhodes and Kator, 1999). Studies of other groups such as *Bacteroides* and *Peptostreptococcus* have been disabled by the complexity of the methods needed for their identification.

TABLE 6.3 The 10 most common bacterial species in faeces (Moore and Holdeman, 1974)

1.	<i>Bacteroides vulgatus</i>
2.	<i>Fusobacterium prausnitzii</i>
3.	<i>Bifidobacterium adolescentis</i>
4.	<i>Eubacterium aerofaciens</i>
5.	<i>Peptostreptococcus productus</i> – 2
6.	<i>B. thetaiotaomicron</i>
7.	<i>E. eligens</i>
8.	<i>Peptostreptococcus productus</i> – 1
9.	<i>E. biforme</i>
10.	<i>Eubacterium aerofaciens</i> – 3

Thus, in conclusion, it will be clear that the human intestine is the source of a diverse range of bacteria, but that from the practical viewpoint many can be ignored at present in the context of water testing. However, it must be realized that such testing considerably underestimates the actual bacterial load that arises from faecal pollution.

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Faecal indicator organisms

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1 INTRODUCTION

Snow (1855) first demonstrated that gastrointestinal diseases such as cholera were able to be transmitted via faecally-contaminated water. His work, now rightly regarded as a classic in the field of public health medicine in general and water science and engineering in particular, was all the more remarkable as it was done without any knowledge of the causative microorganism, *Vibrio cholerae*. Smith (1895) showed that the presence of the bacterium, then known as *Bacillus coli communis*, could be regarded 'as a valuable indication or symptom of pollution', and he was the first bacteriologist to promote what later became known as the coliform group of bacteria as indicators of faecal pollution. Water bacteriology was then a rapidly advancing field: of the 26 papers presented at the first meeting of the Society for Bacteriology (now the American Society for Microbiology), five were concerned with *B. coli communis* and water and wastewater (Dixon, 1999). The first edition of *Standard Methods* appeared in 1905 and included recommendations for the bacteriological examination of waters (APHA, 1905). In contrast, the first edition of *Report 71*, the UK standard on the bacteriological examination of drinking waters, was not published until 1934 (Ministry of Health, 1934). Emphasis in these early years was on the detection of coliform bacteria using the bile salt–lactose medium developed by MacConkey (1908), although tests for the faecal enterococci and *Clostridium perfringens* were also included.

Faecal indicator organisms remain at the forefront of water and wastewater

microbiology. They represent our first and foremost technique for detecting and quantifying aquatic pollution. The American Society for Microbiology has recently declared that 'Microbial pollution of water in the United States is a growing crisis in environmental and public health' (ASM, 1999). The role of faecal indicator organisms is central to the reduction of this crisis which is occurring in all parts of the world.

2 THE IDEAL FAECAL INDICATOR BACTERIUM

Bonde (1962), the World Health Organization (WHO, 1993), Grabow (1996) and Godfree *et al.* (1997) list the following properties of the ideal faecal indicator bacterium:

1. suitable for all categories of water
2. present in wastewaters and polluted waters whenever pathogens are present
3. present in greater numbers than pathogens
4. having similar survival characteristics as pathogens in waters and water and wastewater treatment processes
5. unable to multiply in waters
6. non-pathogenic
7. able to be detected in low numbers reliably, rapidly and at low cost.

Of course, such an ideal organism does not exist. Some, such as faecal coliform bacteria, come close, but even so, difficulties remain with the indication of non-bacterial pathogens (viruses, protozoa and helminths).

3 COLIFORM BACTERIA

Early water microbiologists defined coliform bacteria as those bacteria able to grow at 37°C in the presence of bile salts (used to inhibit non-intestinal bacteria) and able to produce acid and gas from lactose. Faecal coliforms were considered to be those coliforms which were of exclusively faecal origin and consequently able to grow and ferment lactose at 44°C, and to produce indole from tryptophan. However, the last edition of *Report 71* (Department of the Environment, 1994) defines coliform bacteria as those members of the Enterobacteriaceae possessing the gene coding for the production of β -galactosidase, the enzyme which cleaves lactose into glucose and galactose. This new definition is not dependent on laboratory techniques (acid and gas production from lactose), and recognizes that gene expression may be influenced by environmental (including laboratory) conditions – i.e. under different conditions a coliform bacterium may or may not ferment lactose with the production of acid and gas.

The coliform group, as defined above, includes species of the genera *Citrobacter*, *Enterobacter*, *Escherichia*, *Hafnia*, *Klebsiella*, *Serratia* and *Yersinia*. Some members of the genus *Aeromonas* may mimic coliforms in the standard laboratory tests for the latter, but they are oxidase-positive, whereas all coliforms are oxidase-negative.

Faecal coliform bacteria are principally *Escherichia coli*, but some non-faecal coliforms may also grow at 44°C. *E. coli* was found to be a better indicator of the risk of diarrhoeal disease from tropical drinking waters than faecal coliforms (Moe *et al.*, 1991). However, for routine purposes, faecal coliforms, thermotolerant coliforms and *E. coli* may be regarded, if not exactly the same, as generally equivalent indicators of faecal pollution. Total coliforms (i.e. faecal and non-faecal coliforms) are routinely used as indicators of the general bacteriological quality of treated drinking waters, and only waters that have tested positive for this group are examined for the presence of faecal coliforms. Those workers who favour *E. coli* over faecal coliforms

(e.g. Gleeson and Grey, 1991) do so on the grounds that some non-faecal coliforms are thermotolerant; however, since humans and many animals excrete around 10^{11} *E. coli* per day and over 95% of excreted thermotolerant coliforms are *E. coli*, in practice this distinction is but a fine one.

Total coliforms may grow in certain non-intestinal environments, e.g. in biofilms in water distribution pipes, but such regrowth of faecal coliforms is rare. Nevertheless, Payment *et al.* (1993) considered regrowth within the distribution system as a major potential cause of gastrointestinal illness attributable to the consumption of treated drinking waters meeting current microbiological water quality standards. The potential for regrowth in distribution systems is best assessed by the numbers of *Pseudomonas* spp.; this group was found to be a better indicator than the usual faecal bacteria or *Aeromonas* spp. (Ribas *et al.*, 2000). Recently *E. coli* has been found to regrow in subtropical coastal environments (Solo-Gabriele *et al.*, 2000).

Despite these certain limitations, faecal coliforms, and particularly *E. coli*, remain the best overall indicators of faecal pollution of waters (Edberg *et al.*, 2000). They are of course especially good indicators of bacterial pathogens, less good indicators of viral pathogens, but very poor indicators of protozoan and helminthic pathogens.

4 FAECAL ENTEROCOCCI

The faecal enterococci comprise species of two genera, *Enterococcus* and *Streptococcus*, which occur in the intestines, and hence faeces, of humans and many animals (Leclerc *et al.*, 1996). Until recently they were all described as faecal streptococci, and the term 'intestinal enterococci' has been proposed (Hernandez *et al.*, 1995); however, the original description (Thiercelin in 1899) was 'enterococcus' and it seems sensible to keep the familiar epithet 'faecal'. The species of *Enterococcus* and *Streptococcus* that are of faecal origin are listed in Table 7.1.

TABLE 7.1 Species of the genera *Enterococcus* and *Streptococcus* of faecal origin

Species	Intestinal origin
<i>Enterococcus</i>	
<i>faecium</i>	Man, cattle, pigs, birds
<i>faecalis</i>	Man, cattle, pigs, birds
<i>durans</i>	Man, pigs, birds
<i>hirae</i>	Man, pigs, birds
<i>avium</i>	Man, cattle, pigs, birds
<i>gallinarum</i>	Man, birds
<i>cecorum</i>	Cattle, pigs, birds
<i>columbae</i>	Cattle, pigs, birds
<i>Streptococcus</i>	
<i>bovis</i>	Man, cattle, pigs
<i>equinus</i>	Man, cattle, pigs
<i>alactolyticus</i>	Pigs, birds
<i>hyointestinalis</i>	Pigs
<i>intestinalis</i>	Pigs
<i>suis</i>	Pigs

Source: Godfree *et al.* (1997).

The faecal enterococci fulfill most of the requirements of an ideal indicator – the species listed in Table 7.1 are exclusively associated with the intestinal tract of man or other animals; they do not multiply in the environment; they occur in much higher numbers than bacterial and viral pathogens (around 10^6 per 100 ml of raw domestic wastewater); and they are around twice as resistant to disinfection as faecal coliforms (Godfree *et al.*, 1997).

While the taxonomy of the faecal enterococci has improved recently, as shown in Table 7.1, methods for their isolation from faecal and environmental samples has not – these are essentially tests for the enumeration of the faecal streptococci, and the use of these can lead to erroneous results and misleading conclusions concerning the pollution source (Pinto *et al.*, 1999). Clearly there is a need to evaluate and modify media and methods for faecal streptococci, so that reliable techniques exist for the isolation and enumeration of faecal enterococci (and preferably as a group, i.e. all the species of both genera in Table 7.1, although methods for individual species identification and enumeration would be epidemiologically useful).

5 CLOSTRIDIUM PERFRINGENS

Clostridium perfringens (formerly *Cl. welchii*) is an anaerobic sulphite-reducing spore-forming bacterium. It is present in human and animal faeces, but in lower numbers than faecal coliforms or faecal enterococci. If it can be cultured from a water sample, but faecal coliforms and faecal enterococci cannot be, its presence indicates:

either 'remote' faecal pollution, i.e. pollution that has occurred at sometime past, since when faecal coliforms and faecal enterococci have died off, but not the spores of Cl. perfringens,
or faecal pollution in the presence of industrial wastewater toxicants which have killed or inhibited faecal coliforms and faecal enterococci, but not the spores of Cl. perfringens.

It is thus generally only used when faecal coliforms and faecal enterococci cannot be detected, or when industrial toxicants are present (or suspected to be present).

6 BACTERIAL INDICATORS OF HUMAN AND ANIMAL FAECAL POLLUTION

Geldrich and Kenner (1969) advocated the use of faecal coliform/faecal streptococci (FC/FS) ratios to differentiate between faecal pollution of human and animal origin. Ratios > 4 were held to be indicative of human pollution and ratios < 0.7 of animal pollution. McFetters *et al.* (1974) questioned the general validity of this interpretation, and Oragui (1978) and Wheeler *et al.* (1979) demonstrated unequivocally its invalidity: FC/FS ratios > 4 were found in faecal samples obtained from sheep, pigs, cats, dogs, chickens, ducks, pigeons and seagulls, as well as from humans.

Oragui (1983) found that, in the UK, sorbitol-fermenting strains of *Bifidobacterium breve* and *B. adolescentis* were present only in human faeces, whereas *Streptococcus bovis* and *Rhodococcus coprophilus* were only present in

animal faeces (see also Mara and Oragui, 1981, 1983; Oragui and Mara, 1981, 1984). This was confirmed by Mara and Oragui (1985) in Nigeria and Zimbabwe, although very small numbers of *S. bovis* were found in human faeces.

However, these 'new' specific indicators of human and animal faecal pollution have several disadvantages:

1. sorbitol-fermenting bifidobacteria are extremely sensitive in the extra-intestinal environment, disappearing much more rapidly than faecal coliforms (Oragui *et al.*, 1987; Pourcher, 1991), and they are much more sensitive to chlorine (Resnick and Levin, 1981)
2. the incubation period for *R. coprophilus* is very long, 17–18 days at 30°C
3. *S. bovis* does occur in some human faeces, albeit in very small numbers.

Nonetheless, these new indicators have, in practice, proved useful (Jagals and Grabow, 1996; Rhodes and Kator, 1999), and they are included in the current edition of *Report 71* (Environment Agency, 2002).

Recently, Carson *et al.* (2001) reported successful identification of *E. coli* from humans and individual animals (cattle, pigs, horses, chickens, turkeys, dogs and migratory geese) by ribotyping. This extends earlier work on ribotyping and antibiotic resistance profiles to distinguish between human and non-human (i.e. animals collectively) pollution (Parveen *et al.*, 1999). Antibiotic resistance patterns in faecal enterococci were used by Wiggins (1996) to distinguish human and animal faecal pollution in surface waters.

7 VIRUSES AND BACTERIOPHAGES

Viral pollution, especially of ground and coastal waters, is of concern, particularly in the absence of bacterial indicators, due to the high infectivity and virulence of viral pathogens (Friedman-Hoffman and Rose, 1998). Bacteriophages have been used as indicators of enteroviral pollution of waters. The IAWPRC

Study Group (1991) recommended the use of male (F⁺)-specific bacteriophages, rather than somatic coliphages, as enteroviral indicators (see also Calci *et al.*, 1998). They occur in polluted coastal waters where they are concentrated in shellfish (Chung *et al.*, 1998) and they are found in large numbers in wastewater sludges (Lasobras *et al.*, 1999) and also in aerosols emanating from activated sludge plants (Carducci *et al.*, 1999). Leclerc *et al.* (2000) found that all types of bacteriophage were limited as indicators of enteric viruses in groundwaters due to their low concentration in such waters. They also noted that there are very few field data that permit an understanding of the ecology and survival of phages in the natural environment. Grabow (2001) lists a number of major shortcomings of the use of bacteriophages as indicators of, or surrogates for, enteric viruses in water; thus they cannot be satisfactorily regarded as *absolute* indicators or surrogates, although they can be used in combination with other indicators in the assessment of faecal pollution.

However, Oragui *et al.* (1987) found that faecal coliform numbers in treated tropical wastewaters were good indicators of the level of viral contamination (Table 7.2).

8 FUNGI AND YEASTS

Fungal infections are of increasing concern, particularly in immunocompromised individuals, and an important focus of infection is communal bathing facilities. Research in Portugal (Mendes *et al.*, 1993, 1997, 1998) has led to the fungus *Scopulariopsis* and the yeast *Candida* being proposed as indicators of the microbiological quality of sand on bathing beaches, and to recommendations for maximum levels of these organisms on beaches (10 *Scopulariopsis* and 10 *Candida* per g sand).

9 PROTOZOA AND HELMINTHS

There are no satisfactory indicator organisms for the important protozoa (such as *Giardia lamblia*, *Cryptosporidium parvum* and *Entamoeba*

TABLE 7.2 Geometric mean bacterial and viral numbers^a and percentage removals in raw wastewater (RW) and the effluents of five waste stabilization ponds in series (P1–P5)^b in northeast Brazil at a mean mid-depth pond temperature of 26°C

Organism	RW	P1	P2	P3	P4	P5	Percentage removal
Faecal coliforms	2×10^7	4×10^6	8×10^5	2×10^5	3×10^4	7×10^3	99.97
Faecal streptococci	3×10^6	9×10^5	1×10^5	1×10^4	2×10^3	300	99.99
Clostridium perfringens	5×10^4	2×10^4	6×10^3	2×10^3	1×10^3	300	99.40
Total bifidobacteria	1×10^7	3×10^6	5×10^4	100	0	0	100.00
Sorbitol-positive bifids	2×10^6	5×10^5	2×10^3	40	0	0	100.00
Campylobacters	70	20	0.2	0	0	0	100.00
Salmonellae	20	8	0.1	0.02	0.01	0	100.00
Enteroviruses	1×10^4	6×10^3	1×10^3	400	50	9	99.91
Rotaviruses	800	200	70	30	10	3	99.63

^a Bacterial numbers per 100 ml, viral numbers per 10 litres.

^b P1 was an anaerobic pond with a mean hydraulic retention time of 1 day; P2 and P3–5 were secondary facultative and maturation ponds respectively, each with a retention time of 5 days. Pond depths were 3.4–2.8 m.

Source: Oragui *et al.* (1987).

histolytica). Examination of water and wastewater samples thus has to be for these organisms themselves. However, the cost of routine monitoring outweighs potential benefits (Fairley *et al.*, 1999).

The same is true for helminthic pathogens. When treated wastewaters are used for crop irrigation, the World Health Organization recommends that they contain no more than one human intestinal nematode egg per litre (eggs of *Ascaris lumbricoides*, *Trichuris trichiura*, and the human hookworms) (WHO, 1993), and treated wastewaters are analysed directly for these (Ayres and Mara, 1996).

10 WASTEWATERS

Feachem and Mara (1979) reviewed the usefulness of faecal indicator bacteria in wastewaters, the faecal origin of which is not generally in doubt. Faecal coliform bacteria are regarded as the most useful indicator of the microbiological purifications achieved by wastewater treatment and disinfection. Standards and guideline values for the microbiological quality of recreational waters subject to the discharge of treated wastewaters, and of treated wastewaters used for crop irrigation and fish culture, are commonly expressed in terms of maximum permitted numbers of faecal coliforms (FC). For example, bathing waters in the European

Union should contain no more than 2000 FC per 100 ml (Council of the European Communities, 1976), and treated wastewaters used to irrigate vegetables eaten raw no more than 1000 FC per 100 ml (WHO, 1989; see also Mara, 1995; Shuval *et al.*, 1997).

Oragui *et al.* (1987) showed that faecal coliforms were indeed good indicators of bacterial and viral pathogens in wastewater treatment in waste stabilization ponds (WSP; see Chapter 26): when FC numbers were 7000 per 100 ml, salmonellae and campylobacters were absent and faecal viruses present in only very small numbers (see Table 7.2). Oragui *et al.* (1993) found that *Vibrio cholerae* was eliminated in a series of WSP when FC numbers were 60 000 per 100 ml.

11 THE FUTURE?

Haas *et al.* (1999) present order-of-magnitude calculations for waterborne infections in the USA, where there is an estimated number of waterborne illnesses of 1 million. Taking the population of the USA as 280 millions, this is equal to an annual occurrence rate of 0.0036, equivalent to a daily rate of 9.8×10^{-6} . Using dose-response parameters for rotavirus, *Salmonella*, *Giardia* and *Cryptosporidium*, the daily intake of organisms corresponding to this level of daily risk can be determined, as

TABLE 7.3 Daily microbial intakes per person corresponding to an estimated daily rate of waterborne disease in the USA of 9.8×10^{-6} per person

Organism	Dose-response model and parameter values	Daily microbial intake per person corresponding to daily risk of 9.8×10^{-6} per person ^a ($= d$) ^{b,c}
Rotavirus	Beta-Poisson ^b $N_{50} = 6.17$ $\alpha = 0.263$	1.78×10^{-5}
Salmonella	Beta-Poisson $N_{50} = 23\,600$ $\alpha = 0.313$	0.091
Giardia	Exponential ^c $r = 0.0199$	4.92×10^{-4}
Cryptosporidium	Exponential $r = 0.0042$	2.33×10^{-3}

^a 1 million waterborne illnesses per year in a population of 280 millions, equivalent to a daily risk of waterborne illness of $(1/280 \text{ million})/365$, i.e. $= 9.8 \times 10^{-6}$ per person.

^b $P_1(d) = 1 - \{1 + [(d/N_{50})(2^{1/\alpha} - 1)]\}^{-\alpha}$ where $P_1(d)$ = probability (risk) of infection per day (= fraction of persons ingesting d pathogens per day who become infected), d = average daily dose, N_{50} = median infectious dose, and α = a species constant.

^c $P_1(d) = 1 - \exp(-rd)$, $\approx rd$ for $rd \ll 1$ where r = a species constant.

Source: Haas *et al.* (1999), pp. 383 and 435.

shown in Table 7.3. The computed numbers are small and, given the immense difficulties in measuring low numbers of pathogens in treated drinking waters, it is highly unlikely that these numbers are measurable. For example, in the case of *Cryptosporidium*, assuming all exposure is from drinking water and that water consumption is 2 litres per adult per day, the daily intake of 0.00233 oocyst per person translates into approximately 0.001 oocyst per litre of treated drinking water; this would require the examination of at least one thousand 1-litre samples for a substantial probability of detection (and this assumes that oocyst counting methods are 100% efficient, which they are not; typically they are <50%).

Haas *et al.* (1999) conclude by noting that 'the general inability to measure pathogens in water as [it is] being consumed is not necessarily inconsistent with the estimated preva-

lence of waterborne infectious diseases.' Given this, it makes little sense to examine treated drinking waters routinely for the presence of pathogens (some of which have long incubation periods – 5–28 days in the case of *Cryptosporidium* and 30–60 days for hepatitis A virus) (see Fairley *et al.*, 1999). This is true also during epidemics of waterborne disease since, as noted by Haas *et al.* (1999), even for the largest known outbreak of waterborne disease (that in Milwaukee in March–April 1993 in which over 400 000 persons became ill with cryptosporidiosis), local public health authorities were unaware of the outbreak during its first 4 days.

The order-of-magnitude calculations of Haas *et al.* (1999) and the above conclusions drawn from them mean, in practice, that *either* microbial indicator organisms must continue to be used for the routine assessment of faecal contamination of treated drinking waters, at least until pathogen detection methods improve to the point that they can reliably detect, e.g. *Cryptosporidium* levels of 1 oocyst per 1000 litres (perhaps through the use of gene probes), *or* a train of water treatment processes are specified which can reliably achieve such low pathogen levels. The latter approach is favoured by the USEPA (2002) in its Long Term 1 Enhanced Surface Water Rule, which also includes watershed control and reservoir protection. A useful supplement to this would be Hazard Analysis Critical Control Point (HACCP), which has been widely used in the food industry but only recently proposed for drinking water (Deere and Davison, 1998).

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Detection, enumeration and identification of environmental microorganisms of public health significance

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1 INTRODUCTION

Detection and enumeration of environmental bacteria is a vast and exciting subject area and one that undergoes continual change as new environments are examined and new methods are developed. Recent years have witnessed an ongoing revolution in detection methods based on techniques from molecular biology. These methods hold the potential for reduced analytical time and improved sensitivity and specificity when compared to the traditional cultural methods that now constitute the majority of contemporary detection and enumeration techniques. While molecular methods, based on specific gene sequences, offer the possibility of simultaneous detection of multiple pathogens, thereby addressing a classically vexing problem in the sanitary microbiology of natural waters, their application also raises important questions related to target viability and infectivity. Because organisms must be isolated for study and method development, culture-based techniques will undoubtedly remain important tools in the environmental microbiologist's arsenal of methods. In the near future, Sartory and Watkins (1999) envisioned a melding of conventional culture with biochemical methods for detection of viable cells over shorter time intervals.

This chapter will focus primarily on detection and enumeration topics related to the sanitary microbiology of receiving waters. Problems of surface water sanitation remain significant world-wide and advances in approved method development have been surprisingly limited given the history and continued scope of disease problems encountered. Microbial contamination of groundwater used for drinking purposes is a dominant concern in many countries. The material presented is intended as a general overview, with specific detailed discussion of selected environmental microorganisms of interest used in public health and water quality assessment. It is not intended as a comprehensive literature review.

Microorganisms of sanitary importance present in natural waters are derived from domestic sewage and non-point runoff containing the excreta of humans and animals. Rivers, lakes, and coastal waters serve to receive, dilute and disperse domestic sewage and untreated human and animal waste. The degree of domestic waste treatment varies considerably (Chambers *et al.*, 1997) and its influence on water quality and health remain important problems throughout the world. Improperly operating septic systems, poor well maintenance, surface application of wastewaters and direct injection have led to contamination of

groundwater. Eutrophication, ecosystem perturbation, toxicities, and disease, in both human and animal communities, are consequences of waste discharge. From a human disease perspective, domestic wastes contribute bacterial (salmonellas, vibrios and aeromonads), viral (Norwalk, polio, hepatitis, enteroviruses), and protozoan (*Cryptosporidium*, *Giardia*) pathogens. Large numbers of indicator organisms (e.g. coliforms, enterococci, streptococci, bacteriophage, *Bacteroides fragilis*, *Bifidobacterium* spp, clostridia) and potential 'marker' compounds (proteins, enzymes, sterols, whitening agents) are also found in these wastes.

2 DETECTION AND ENUMERATION

Microorganisms, especially those normally associated with warm-blooded animals, are likely to become stressed in natural environments from exposure to unfavorable temperatures and abiotic factors such as light, osmotic stress, availability of carbon, and chemical stressors (e.g. disinfection). Importantly, there is a direct link between the efficiency of recovery and degree of stress using traditional media and procedures (Kator and Rhodes, 1991). Many investigators have also observed changes in culturability that can also affect recoverability (Roszak *et al.*, 1984; Rollins and Colwell, 1986; Lopez-Torres *et al.*, 1987; Munro *et al.*, 1987; Roth *et al.*, 1988; Martinez *et al.*, 1989; Desmonts *et al.*, 1990; Garcia-Lara *et al.*, 1991; Cappelier and Federighi 1998; Lleò *et al.*, 1998). Moreover, while most methods listed in compendiums such as Standard Methods (APHA, 1998) can be applied to detection and enumeration of target microorganisms from saline waters there is no reason to assume that detection methods developed and verified for freshwaters will be as effective in saline waters. Various studies, including our own, have proven such an assumption unwise.

2.1 Sampling considerations

Detection and enumeration are final procedures in a chain of events that commence

with sample collection and sample storage during transport. Sample collection and sample storage are thus important initial elements of detection/enumeration protocols. Lapses in these activities can affect recovery and enumeration of microorganisms of interest. Improperly prepared or contaminated sampling containers or gear, inadvertent contamination of samples, poor or inappropriate sample storage conditions, excessive transport/storage time, are all factors that can influence ultimate detection and affect quantitative validity. The most careful analyses may be compromised by poor sampling and/or adverse sample storage during transport. Sample volume must be adequate for the organism being sought. True sample replication as opposed to analytical replication might be necessary if the spatial distribution of the microorganism is highly variable.

Sampling locations should be chosen to reflect the desired characteristics of the environment being evaluated. Source evaluation surveys are useful and provide information on which to base a station location process. Consider pollution sources, delivery mechanisms and hydrographic parameters relevant to the aquatic or sediment environment being evaluated. Hydrographic considerations might include factors such as water flow and direction, the existence of vertical or horizontal stratification, local circulation patterns, presence of feeder streams or tributaries, and climatological events such as precipitation and wind speed. Microbiological data should be collected and analyzed with regard to known or potential pollutant sources to confirm the validity of station selection. Advanced modeling techniques, land use data, historical hydrographic data sets, and GIS information systems may be of value in this process.

2.1.1 Sample collection methods

General aseptic principles as outlined in Standard Methods (APHA, 1998) should be followed. Collect water samples for bacteriological examination in clean, sterile containers. Protect the container fully against contamination before, during, and after collection.

Surface water samples can be collected without the aid of special sampling devices. Keep the sample container unopened until immediately before filling. During sampling protect the container stopper or closure from contamination. To collect a sample by hand, hold the bottle near the base and plunge it neck downward below the surface when the water depth permits. Then tilt it with the neck pointing slightly upward and into the direction of flow. During filling, push the container horizontally forward in a direction away from the hand to avoid contamination. If sampling from a vessel, take care to avoid contamination from effluents and sample 'upstream' of the vessel motion. In shallow waters, avoid sampling from waters that are disrupted by sediment suspended by boat activity. An ordinary laboratory clamp affixed to a length of steel or aluminum tubing can be used to hold a bottle for sampling as directed above.

Commercial bottle holding devices are available, but it is very easy to fabricate one's own. Appropriate personal safety precautions should be taken if sampling untreated sewage or similar wastewaters to avoid direct contact. To facilitate sample mixing prior to enumeration, leave a small air space in the bottle before sealing. Precautions against chlorine residuals should be considered when human point source impacted waters or drinking waters are sampled (APHA, 1998).

A variety of sampling devices are available for collection of subsurface samples. Ideally, a subsurface sampler should be of simple (i.e. reliable) mechanical design and able to hold a sterile container that is non-metallic and non-bactericidal, with mechanical devices to open and shut the container at the desired sampling depth. The container should remain sealed until the desired depth is reached. Sampling depth can affect recovery of bacterial populations, especially in dynamic environments where currents vary temporally as a function of tides or winds and bed resuspension may be an important factor. Using *Vibrio* spp. as a target group, Koh *et al.* (1994) found significant variation in cell concentrations occur as functions of both depth and tidal cycle.

A number of subsurface sampling devices are available commercially. Water sampling 'bottles' that remain open prior to triggering and closing at depth are common. However, their use must be considered carefully if contamination during descent will affect results. Bottles should be cleaned between uses with a suitable, non-toxic liquid if contaminant adsorption is a concern. DC-powered peristaltic pumps and sterile tubing can also be used for sampling at depth provided the head capacity of the pump is not exceeded and the tubing is adequately flushed prior to sample collection. Fresh sterile tubing should be used for each new sample. Peristaltic pumps of this type can also be used routinely to sample shallow groundwater wells in the field.

An area that has received little attention is the use of continuous or integrating *in situ* samplers for microbiological sampling. In highly dynamic environments or in those with low densities of bacteria, unattended samplers that collect large multiple sample volumes over selected time intervals 'integrate' water conditions and may provide better means to assess sanitary water quality when compared to traditional grab sampling.

Various studies illustrate that bacteria are concentrated at interfaces such as the surface microlayer (e.g. Hatcher and Parker, 1974; Marshall, 1976; Dutka and Kwan, 1978). The importance of this microhabitat is now recognized as a region regulating air/water gas exchange and the capture of organic compounds of low solubility in water (Sodergen, 1993). There are no standard methods for collection of bacterioneuston, which can be sampled using a variety of techniques ranging from glass plates to mesh screens (van Vleet and Williams, 1980). Care must be taken to use sterile sampling devices to recover microorganisms from microlayers.

Methods for concentration of viruses, chemical indicators and other non-approved indicators may require special techniques. These methods should be evaluated and validated prior to actual field studies. Soil and sediment samples can be collected using sterile spatulas or coring devices. An inexpensive surface corer can be made by cutting off

the luer end of a 50-ml autoclavable plastic syringe. Similarly prepared smaller volume syringes can be used for replicate sampling of single core samples. Sampling of groundwater and subsurface soils may require special equipment and great attention must be given to avoid sample contamination.

2.1.2 Sample storage

Sample storage is often not considered a vital part of the enumeration/detection process. Simple icing is usually recommended for field samples by many sources. The degree of sample cooling required can vary depending on the activity of a sample. Highly enriched samples with active heterotrophic microbial communities (e.g. domestic sewage, eutrophic lake waters) may change in composition if temperature is not reduced quickly. On the other hand, icing water samples can produce sublethal stress expressed as reduced recovery when target faecal organisms collected from cold waters are placed directly in warm media without an intermediary resuscitation or adaptation step (Rhodes *et al.*, 1983). Samples collected from chlorinated effluents must be treated with sodium thiosulfate to neutralize residual chlorine following recognized protocols (APHA, 1998). In general, it is always best to process samples as soon as possible after collection and never to exceed the specified time limits for a given technique. These and other concerns are described in Standard Methods (APHA, 1998). For non-regulatory methods, sample storage parameters should be evaluated empirically using spiked samples and controls for sample/microorganism interactions.

2.1.3 Accountability

A good sampling program must be supported by a consistently applied laboratory and field quality control plan. Thoroughly identify sample containers using an easily understood coding scheme and use field record sheets or water resistant notebooks to provide sample accountability ('chain of custody') from field to laboratory personnel. Use pencils and marking pens that will remain legible under field and laboratory conditions. The chain of custody

may be important for quality control or for epidemiological or legal purposes. Insure that all field observations include date, time, and place of collection; identity of person responsible for collection; weather and water conditions at time of collection; and comments on unusual conditions or events observed.

2.2 Relevant physical and chemical parameters

Interpreting field microbiological data requires an understanding of the physical and chemical characteristics of the environment sampled. Routine parameters measured or described during sampling of surface waters can include temperature, conductivity or salinity, pH, and oxygen concentration. Non-routine parameters, such as stage and current of the tide, Secchi depth or water clarity, amount of precipitation, total suspended solids, selected nutrients, chlorophyll *a* and Eh may be required for specific studies. For sediments, parameters such as sediment dry weight, total organic concentration, or grain size distribution are common variables. Approved methods and instrument specifications for some physical and chemical parameters can be found in Standard Methods (APHA, 1998). References for saline waters (Strickland and Parsons, 1972) and soils (e.g. Weaver, 1994; Parks, 1996) are very useful sources of analytical methods. Many routine parameters can now be measured easily using lightweight portable datasondes, which usually possess comparatively large data storage capability. Salinity can be determined by conductivity or with a hand refractometer if the reduced accuracy of the latter instrument is acceptable. Rainfall often can be site-specific and it is recommended that a simple rain gauge or recording rain gauge be installed and monitored for each study site. Antecedent precipitation should be measured at regular intervals prior to sampling. Remote areas may necessitate installation of a battery-operated recording rain gauge. In all instances, instrumentation should be calibrated using

appropriate standards and verified to be within specifications prior to and during sampling, and in some cases, after sampling is completed.

3 METHODS OF ENUMERATION AND DETECTION

The basis of much of environmental microbiology over its brief history has been learned through the removal and isolation of microorganisms under artificial laboratory conditions from natural environments to which they have adapted. While this approach has been functionally necessary, naturally occurring microorganisms belong to natural assemblages where they often exist as synergistic members of mixed populations and their isolation in a defined medium is an obvious oversimplification. Given this background it is not surprising that we can culture but a minor proportion of the total microbial community from a given habitat. Future researchers will no doubt become more adept in developing more sophisticated recovery methods. The recovery of microorganisms from environments to which they are poorly adapted, i.e. pathogens and commensals of warm-blooded animals in surface waters, has been the focus of much enumeration/detection method development for obvious reasons. Even within this group, recognition of the necessity for resuscitation, attention to stress, and even the effects of starvation, has now become part of enumeration/detection methods. The following general discussion contains information related to the application of cultural methods.

3.1 Non-culturability

An important concept influencing the detection of certain enteric pathogens and indicator organisms following environmental exposure was first described by Xu *et al.* (1982), who demonstrated that cells starved in seawater rapidly enter a 'non-recoverable' dormant stage when assayed with standard media and conditions, but remain viable by a direct viable count assay (Kogure *et al.*, 1979). Reduced

culturability has been observed by others as well (Roszak *et al.*, 1984; Rollins and Colwell, 1986; Lopez-Torres *et al.*, 1987; Munro *et al.*, 1987; Roth *et al.*, 1988; Martinez *et al.*, 1989; Desmonts *et al.*, 1990; Garcia-Lara *et al.*, 1991; Cappellet and Federighi, 1998; Lleò *et al.*, 1998) and reported to occur over a variety of microbial genera (e.g. Oliver, 1993; Caro *et al.*, 1999). Moreover, the viable-but-non-culturable (VBNC) state for certain species is characterized by a sequence of demonstrable physiological and biochemical changes (Oliver, 1993). If entry into the VBNC state represents a physiologically-programmed survival response to adverse environmental conditions (Morita, 1993), VBNC cells should be capable of resuscitation under appropriate circumstances. Resuscitation attributed to nutrient addition is difficult to demonstrate unequivocally because the complete absence of culturable cells in a population of VBNC cells may be difficult to show. Bogosian *et al.* (1996) discounted entry of *E. coli* into a VBNC state concluding that non-culturable cells were non-viable. Moreover, the extent of entry into a non-culturable state is affected by prior growth conditions (Munro *et al.*, 1995). Contemporary readings on starvation and non-culturability can be found in Colwell and Grimes (2000).

A practical consequence of the VBNC phenomenon is that, although variable numbers of metabolically active cells may remain in a sample, because of non-culturability the 'true' numbers (and consequences) of indicator (e.g. *E. coli*) or pathogenic microorganisms may be underestimated. Thus, the effectiveness of public health standards based on quantitative recovery of microorganisms susceptible to non-culturability could be compromised. *Enterococcus faecalis*, an organism identified as an indicator of risk associated with swimming-associated illnesses has been reported to enter the VBNC state (Lleò *et al.*, 1998). The extent to which the VBNC response affects detection and enumeration of bacteria in natural populations, in wastewater facilities, in low-nutrient environments such as groundwater, and in environmental biofilms remains to be described and evaluated.

3.2 Cultural considerations

3.2.1 Media components and cultural conditions

Cultural methods are based upon providing a combination of nutritional and physicochemical conditions that will support the growth of the microorganisms of interest. The range of media developed to culture bacteria reflects their nutritional versatility and range of growth conditions tolerated. The diversity of unique microbial physiologies found in domestic sewage, fecal wastes, and aquatic environments necessitates methods that span a wide range of recovery conditions. Most of the bacteria dealt with in sanitary microbiology are chemoorganotrophs (also called chemoheterotrophs), i.e. bacteria that require organic compounds for energy and as a carbon source. The range of organic compounds an individual species can degrade varies from simple carbohydrates to complex molecules such as polycyclic aromatic hydrocarbons and synthetic halogenated molecules. Bacteria with very specific substrate trace organic and physicochemical chemical requirements are termed 'fastidious'.

Basic components of isolation media include carbon sources to enable growth, an appropriate ionic environment, trace elements, vitamins, a buffer system to stabilize pH as metabolic products accumulate or substrates are removed from the medium, electron acceptors, and nutrients. A general discussion of factors involved in formulation of culture media can be found in Tanner (1996) and Cote and Gherna (1994).

Organic components commonly used in media are classified as either complex or chemically defined. Components such as peptones (products of hydrolytic reactions) or yeast extract are complex organic mixtures whose exact composition will vary from biological source to source. Chemically-defined media contain analytical-grade chemicals of known composition and purity such as glucose, inorganic salts, buffers, metals, dyes or inhibitors.

In addition to a carbon source culture media may contain trace compounds such as vitamins and trace metals required as enzyme cofactors. Salts are commonly added for osmotic balance,

buffering of pH, and as sources of nitrogen and phosphorus. Frequently, various compounds are added which are indicators of pH, oxidation/reduction potential or specific metabolism. Biological dyes may be added to enhance contrast between bacteria and the medium. Examples are phenol red as a pH indicator; tetrazolium salts, which change color, are used as redox or electron transport indicators, and a variety of substrates conjugated to chromogens are available to assess species-specific metabolic activities. Examples of the latter include carbohydrate analogs which are conjugated to fluorescent or chromogenic molecules. Enzymatic hydrolysis of specific bonds leads to release of the reporter molecule signaling the presence of the target microorganism. An example of a common method to determine if *Escherichia coli* is associated with positive tubes in the fecal coliform MPN test (APHA, 1998) is based on hydrolysis of a compound called methyl-umbelliferyl- β -glucuronide. Use of this compound is based on the specific presence of the enzyme β -glucuronidase (GUD) in *E. coli* and is the basis for a variety of detection methods (Feng and Hartman, 1982; Watkins *et al.*, 1988; Rice *et al.*, 1990; Sartory and Howard, 1992). A new medium, mTEC (modified mTEC, USEPA, 2000) contains a chromogenic compound (5-bromo-6-chloro-3-indoyl- β -D-glucuronide) which is the basis for enumeration and detection of *E. coli* in one step (USEPA, 2000). The critical assumption in using chromogenic or fluorogenic analogs, or any other substrates to detect a translational product, is that the enzyme is produced in all target microorganisms in the sample. This assumption appears generally correct, but false negative and false positive reactions can occur. For example, Watkins *et al.* (1988) reported that some *Citrobacter* and *Enterobacter* spp. produced GUD.

Some bacteria may also require very low oxygen levels or none at all in order to grow. Bacteria that require oxygen at atmospheric concentrations are called aerobes. Bacteria that require oxygen at very low levels are termed microaerophiles. Bacteria that are intolerant of oxygen are called anaerobes. Coliforms and fecal coliforms are facultative anaerobes,

meaning they can grow under aerobic or anaerobic conditions through a metabolic process known as fermentation.

pH is an important variable that must be maintained within a comparatively defined range for most bacteria. Some bacteria require relatively narrow pH ranges. Commercially available media should be checked for appropriate pH values. Media made from individual components for recovery of unique or specialized groups of bacteria must always be checked for pH. Many bacterial genera found in fecal wastes and domestic sewage are facultative anaerobes and require no special considerations for oxygen or Eh. Obligate or aerotolerant anaerobes used as faecal indicators, e.g., *Bacteroides* spp. or *Bifidobacterium* spp., may require complete elimination or reduction of gaseous oxygen, the presence of CO₂ and low Eh values. These incubation conditions can be achieved using water-activated gas generators and portable plate chambers, or more complex and difficult to maintain anaerobe chambers using gas mixtures and catalysts to remove oxygen. Methods for the culture of anaerobes have been described in detail in Holdeman *et al.* (1977).

Salt content and ionic composition is normally not a major issue with recovery of bacteria of sanitary significance and salts are present in media components as contaminants. Environmental exposure to salt has been shown to affect recovery of indicator bacteria from estuarine or marine waters, but resuscitation from salt exposure is not a part of accepted enumeration procedures.

3.2.2 Media preparation

Media preparation is an important aspect of the culture-based approaches. The importance of following a reasonable and consistent QA/QC program designed to evaluate reagent water quality, sterilization procedures and effectiveness, pH, temperature, gravimetric measurements, chemical components and other aspects of media preparation cannot be overemphasized. Storage of prepared media should be considered with regard to published shelf life and exposure to light. Bolton *et al.* (1984) demonstrated the production of labora-

tory light-mediated photochemical products in prepared agar media. Chelala and Margolin (1983) documented that a medium containing riboflavin, aromatic compounds such as tryptophan or phenylalanine, and MnCl₂ produced bactericidal products when exposed to visible light. Media should never be stored near windows where exposure to sunlight is possible. Carlsson *et al.* (1978) reported the formation of bactericidal hydrogen peroxide and superoxide radicals in anaerobic broths containing glucose and phosphate when autoclaved in the presence of oxygen. Inhibitors can be avoided using PRAS or prereduced media prepared by removal of oxygen prior to and after autoclaving.

3.2.3 The recovery process and the design of detection media

Recovery of bacteria from environmental samples poses several practical problems. First, the medium/incubation conditions should be designed to recover all of the target microorganisms present. Second, these same conditions should not favor the growth of interfering microorganisms (the background microbiota) that are always present. From a practical perspective, for many chemo-organotrophic bacteria, these two objectives are mutually exclusive and ideal recovery is unrealized for many bacterial groups.

Consequently, recovery approaches for environmental bacteria may employ an initial non-selective enrichment phase designed to recover and support the growth of the target organism. This enrichment step may or may not incorporate conditions designed to limit the growth of interfering microorganisms found in environmental samples. If the enrichment medium incorporates compounds inhibitory to such non-target microorganisms, the enrichment is said to be selective. Enrichment and selectivity can be combined in one medium and one step, such as the modified medium for recovery of enterococci by membrane filtration (mEI; USEPA, 2000). The presumptive multiple tube coliform test (APHA, 1998) uses a relatively benign initial enrichment phase (lactose broth) followed by transfer to a harsher, more selective medium and

incubation temperatures. Improved recovery can sometimes be obtained from the selective phase because of the primary, benign enrichment step.

Medium components used to provide selectivity by inhibiting growth of unwanted species include organic compounds such as bile salts and sodium deoxycholate, inorganics such as sodium selenite and sodium tetrathionate, and dyes such as brilliant green, malachite green and crystal violet. An example of the influence of inhibitors on recovery of *Salmonella* spp. in various culture media is discussed in Arroyo and Arroyo (1995). A large variety of media has been developed for recovery of fecal streptococci. Audicana *et al.* (1995) evaluated the effect of sodium azide concentration and antibiotic replacement on the selectivity of several agars. Depending upon the organisms being isolated and the environment sampled, inhibitors may be used singly or synergistically.

Inhibitors listed above are commonly employed in media designed to isolate enteric bacteria from environmental water samples. Bile salts, sodium deoxycholate and sodium selenite are used to inhibit Gram-positive organisms in coliform detection media. However, the use of inhibitors is not without consequences. Thus, inhibitors affect the growth of both interfering background and sublethally injured target organisms and can result in reduced recoveries of target bacteria (e.g., Rhodes *et al.*, 1983).

Isolation by streaking or plating samples on solid media differentially supporting the growth of the target organism generally follows and allows for colony isolation and subsequent identification using biochemical, physical characterization, immunological or molecular methods.

Indicator bacteria and some pathogens, especially total and fecal coliforms, can become stressed or altered physiologically in natural waters from exposure to sunlight (Rhodes and Kator, 1990; Sinton *et al.*, 1994; Pommepuy *et al.*, 1996) lack of appropriate substrates (Lopez-Torres *et al.*, 1987), dissolved substances salts or chlorine (McFeters *et al.*, 1986; Munro *et al.*, 1987) and their detection may be compromised. Resuscitation procedures, such as an initial

period of incubation at a lower temperature, have been formally incorporated into several methods (e.g. mTEC; USEPA, 2000).

The development and application of microbiological tests for enumerating environmental bacteria should be accompanied by a rigorous evaluation of the method's performance characteristics. Performance characteristics are empirically determined experimental values used to assess the effectiveness of a given enumeration method to recover the target microorganisms from a selected environment. Important performance characteristics include method specificity, selectivity, precision and recovery (ASTM, 1993). In some instances the upper and lower boundaries of counts are also determined. Method specificity is a measure of a procedure's capability to detect all target microorganisms present in a sample with a minimum of false positives. Selectivity refers to a method's ability to primarily support only the growth of the target microorganism and to minimize interfering background growth. Selectivity is conferred through the use of chemical inhibitors, antibiotics or restrictive physical conditions, such as elevated temperature. Method recovery is simply the number of target organisms recovered from a spiked sample compared to a recovery from this same sample using a 'reference' method. Understanding the performance characteristics of a method is important with regard to method application and the quality of quantitative data. For example, using an extremely precise method with low selectivity might be a waste of time when sampling grossly polluted or eutrophic systems because high background counts would defeat the value of high precision.

Detection of target microorganisms obtained through cultural methods can involve a complete variety of techniques available to the modern microbiologist. These include isolation, purification, and biochemical characterization; use of immunologies including fluorescein-labeled antibodies (Wright, 1992; McDermott, 1996), use of enzyme-specific substrates (Toranzos and McFeters, 1996; Sartory and Watkins, 1999), and molecular techniques such as colony lifts followed by

hybridization with oligonucleotide probes (Wright *et al.*, 1993).

3.3 Qualitative/quantitative cultural methods

Bacteria are enumerated for a variety of purposes that include detection of pollution, detection of specific phenotypic or phylogenetic groups, and to insure compliance with water quality regulations. Cultural methods provide but a single opportunity to detect and count cells that occur in a freshly collected sample. Beyond maximum holding periods sufficient changes occur and results may be unreliable. The application of appropriate QA/QC programs is essential to enumeration methods. While many methods frequently incorporate verification of isolates randomly selected from samples few, if any, prescribe use of pure cultures to assess responses of recovery media or use of bacterial suspensions of known density to assess recovery efficiency. Donnison *et al.* (1999) consider use of standard bacterial suspensions as a quality control approach to routinely monitor method performance.

3.3.1 Most probable number (MPN)

This is an historically common and important enumeration method used to estimate numbers of viable bacteria in surface waters, soils and sediment, animal scat and shellfish. As a quantitative technique it is extremely versatile and its usage extends well beyond its more common application for estimation of total and fecal coliforms in drinking and surface waters. It has been used to enumerate petroleum-degrading bacteria (Mills *et al.*, 1978), phenol and toluene-degrading bacteria (Fries *et al.*, 1997) and *Staphylococcus aureus* (Bennett and Lancette, 1998). When combined with isotopically-labeled substrates, an MPN method can be used to measure densities of populations with specific degradation potential (Lehmicke *et al.*, 1979). The MPN test is a statistical estimate of culturable units and assumes such units are randomly distributed within a sample (Garthright, 1998). An MPN may or may not represent densities of individuals should the cells occur in groups adsorbed to particles.

Functionally, the MPN test is based on dilution of a target microorganism to extinction, i.e. at some dilution, usually by factors of ten, a sample will not contain a single culturable unit. Actual dilutions required are based on experience and MPN tables are commonly based on inoculation of media with 3, 5, 10 or even 12 sample portions at each dilution. The MPN is useful when cells are present at low densities in natural samples. As a statistical estimate based on the binomial distribution, the precision of an MPN value is a function of the number of sample portions assayed for each dilution. In the format generally used in water quality laboratories, the 3 or 5 tube configuration, the 95% confidence intervals of an MPN value are relatively large (APHA, 1998) and significantly better precision can be obtained using a direct count method or increasing the number of portions to 12, 15 or higher values. For improved precision, the MPN method has been miniaturized using microplates, which provide for high portion numbers. Hernandez *et al.* (1991, 1993) used 96-well microtitration plates to enumerate *E. coli* and presumptive enterococci. Gamo and Shoji (1999) evaluated the use of multiple BIOLOG substrate plates to obtain MPN estimates of a given microbial community's capability to utilize selected substrates.

Limitations with the MPN method result from departures from assumptions that cells are randomly distributed, interference or inhibition effects, which can reduce the occurrence of positive results, false positives, and the poor precision when small numbers of portions are used. MPN-based methods are much less precise than direct plating or membrane filtration methods. A clear discussion of the MPN technique with useful formulas and tables can be found in Garthright (1998).

3.3.2 Colony or plate counts using solid media

Colony counts on solid media offer significantly better precision when compared to MPN methods and provide for the isolation of individual clones for identification and study. A general reference supplying details concerning the uses of the methods identified below and others can be found in Koch (1994).

Pour plate. This is a comparatively simple enumeration procedure applicable to samples where densities of target organisms are sufficient to yield appropriate numbers of culturable elements for counting purposes. In this procedure, a small sample volume (approximately 0.1–1 ml) containing suspended bacteria or dilutions thereof is mixed with melted or so-called ‘tempered’ agar (an agar medium is heated or autoclaved to dissolve the agar and then held at a temperature of about 46°C in a water bath to prevent gelling). A known sample volume is placed in a petri dish followed by the tempered agar, carefully mixed to disperse the sample uniformly, and allowed to solidify. Petri dishes should be turned over as soon as feasible to avoid condensation on the lid. Colonies, which will grow both on the agar surface and within the agar, are counted to yield a number of cells per unit sample volume. Limitations of this technique apply to environmentally stressed cells or those sensitive to elevated agar temperatures, which may be killed leading to underestimates of population densities.

Spread plate. A second method to avoid sublethal stress owing to elevated temperatures used in the pour plate is the spread plate. In this technique a small volume of cell suspension, 0.1 ml, is placed on the surface of a slightly ‘dried’ agar plate and rapidly spread over the surface using a sterile bent rod. If the agar water content is correct, the sample will be rapidly taken up as it is being distributed over the surface. Distribution of the sample can be done by rotating the plate with one hand while continuing to spread the sample using the other. Manual or motorized spread plate ‘turntables’ are also available for this purpose.

3.3.3 Membrane filtration

Membrane filters prepared from a variety of chemical stocks are available for recovery of microorganisms from natural waters where their concentrations are often too low for direct plating and large volume MPN plants would be cumbersome. Membrane filters of various functional pore size distributions can be

coupled with appropriate media for the selective recovery of target organisms. The filter pore size commonly used for microbiological methods is 0.45 µm. Depending on the filter, cells are retained owing to mechanical exclusion because of physical pore size or electrochemical forces. Care should be taken to avoid damage to susceptible target microorganisms by unduly high vacuum. Membranes can be placed directly upon solid media or pads saturated with liquid medium. Membranes offer an additional advantage of *in situ* biochemical testing where a membrane can be removed and placed on a second medium or saturated pad containing test reagents to detect specific properties or to calculate a confirmed count. An example of this approach is the urease test used to confirm colonies as *E. coli* on mTEC medium (USEPA, 2000). In this *in situ* test the membrane filter with colonial growth is transferred to a pad saturated with a medium called *urease substrate medium*. After a short incubation period target colonies remain yellow, yellow-green or yellow brown; non-target or urease positive colonies will turn purple or pink. Limitations to the membrane filter method relate to the comparatively narrow range within which method precision applies, e.g. 20–80 colony forming units cfu/plate in the case of USEPA’s membrane methods for enumeration of *E. coli* (USEPA, 2000) and interference in colony development and/or counting due to high suspended solids or toxics in a sample.

3.4 Non-cultural methods

3.4.1 Direct counts

Direct counts differ from culturable counts in that cells are generally counted by microscopic evaluation and thus do not require the use of complex media to facilitate cell growth to yield densities visible to the unaided eye. Direct counts can be determined on sample suspensions prepared from water, soil or sediment. Cells can be visualized using phase contrast microscopy for unstained cells or by staining cells caught on membrane filters using one of a large variety of fluorescent stains now commercially available. Direct counting methods

are eminently amenable to automated counting using image processing hardware and software. The inherent value of direct methods is to reduce the time required for conventional cultural analysis.

In its simplest form quantitative estimates of cell numbers can be determined in a known volume of sample using a special microscope slide/cover slip combination known as a Petroff-Hausser counting chamber. While this technique is simple it does not differentiate live from dead cells, motile cells are difficult to count and should be fixed prior to counting, and sample concentration requires a separate step.

Membrane filter-based assays are now very common and continually expanding in application as new fluorogenic dyes and their conjugates are being developed. Using membrane filters dyed black, common fluorogenic dyes such as acridine orange, 4',6-diamidino-2-phenylindole (DAPI), Hoechst 33258, or proflavine allow for counting total cells fluorescing against a black background. Kepner and Pratt (1994) provide a comprehensive review of direct enumeration methods, preservation, fluorochromes available, procedures and problems with different kinds of samples.

Specificity can be obtained using fluorescent dyes conjugated to specific target antibodies (Tanaka *et al.*, 2000) or oligonucleotide probes. Langendijk *et al.* (1995) and Franks *et al.* (1998) used 16S rRNA targeted fluorescent oligonucleotide probes to quantify *Bifidobacterium* spp. and major anaerobic groups in suspensions prepared from human fecal samples.

Microscope-based techniques to count rapidly and detect total viable cells, the direct viable count, have been developed that are based on incorporation and chemical modification of dyes or biochemicals that combine the advantages of direct counts with the discrimination of viable from non-viable cells. Autoradiographic techniques have also been used (Meyer-Reil, 1978; Tabor and Neihof, 1982; Roszak and Colwell, 1987) but modern safety concerns and disposal costs associated with use of radioisotopes render their use for

routine work prohibitive. Determination of viability can be based on reduction of redox dyes such as tetrazolium salts (Søndergaard and Danielsen, 2001), incorporation of biochemicals that interfere with cell division such as nalidixic acid (Kogure *et al.*, 1979; Roszak *et al.*, 1984; Roszak and Colwell, 1987) and use of conjugated substrates whose hydrolysis by enzymes (e.g. esterase, phosphatase) active in viable cells releases a fluorescent or colored compound that can be detected visually or with photometric techniques (e.g. Gaudet *et al.*, 1996). Nalidixic acid prevents cell division by inhibiting DNA gyrase and, in the presence of appropriate nutrients, cells which can be resuscitated become enlarged and can be separated from 'non-viable' cells using epifluorescence microscopy. Yokomaku *et al.* (2000) improved the Kogure *et al.* (1979) technique by promoting spheroplast formation selectively in viable cells followed by a lysis step to discriminate viable from non-viable cells. Double staining with a fluorogenic ester and a dye that penetrates inactive membranes can be used to separate active from inactive cell populations (Yamaguchi and Nasu, 1997; Tanaka *et al.*, 2000). The dye, 5-cyano-2,3-ditolyl tetrazolium chloride (CTC), provides advantages of direct counts and discrimination of cells capable of respiring versus non-respiring cells (Søndergaard and Danielsen, 2001).

3.4.2 Flow cytometry

Significant progress has been made in adapting flow cytometry to environmental microbiological uses in recent years. Although expensive, flow cytometers offer benefits of individual cell counting, high sample throughput, identification and cell sorting. Advances in direct count techniques now include the use of flow cytometry to quantify total bacteria or when coupled with specific fluorescent dyes can be used to count physiologically active cells (Porter *et al.*, 1995; Yamaguchi and Nasu, 1997; Grégori *et al.*, 2001). CTC has been used with flow cytometry to determine a total viable count (Sieracki *et al.*, 1999). Flow cytometry can be combined with molecular

methods through the use of fluorochrome-conjugated oligonucleotide probes (Amann *et al.*, 1990).

3.4.3 Immunological methods

Use of fluorescent antibodies for detection and enumeration of specific species or groups of microorganisms has been widely used in environmental microbiology (Wright, 1992). Microorganisms or products thereof have been used to produce polyclonal and more recently, monoclonal antibodies. The latter are considered more specific and potentially more sensitive but require considerably more time, expense and skill to produce. Immunological techniques are useful to confirm the identity of a target microorganism in a purified form or in a mixed cultured population. They can also be used for direct counting when conjugated with fluorochromes, such as fluorescein isothiocyanate (FITC). Direct counting can be accomplished on cells immobilized on glass slides, on membrane filters, or by flow cytometry. Some workers consider routine immunological methods, such as immunofluorescence assays, easier to use and less exacting than molecular methods. Issues of concern with immunological methods commonly encountered when applied to intact environmental samples are those of non-specific staining or background fluorescence. Species in a sample may share reactive sites or epitopes or reactive material derived from the target cells may be present throughout the sample. Immunofluorescence assays have been used to detect VBNC *E. coli* and *Vibrio cholerae* in water (Xu *et al.*, 1982) to study the survival of *Campylobacter* in biofilms (Buswell *et al.* 1998) and to enumerate *V. vulnificus* using a combined MPN-enrichment/selective medium method in samples of estuarine water, sediment and oysters (Tamplin *et al.*, 1991). Fiksdal and Berg (1987) evaluated a fluorescent antibody technique for enumeration of *Bacteroides fragilis* group organisms in water.

3.4.4 Molecular and biochemical methods

The evolution and continual development of molecular methods renders a contemporary

review of this material a task fundamentally beyond the scope of this chapter. The reader is referred to current literature for specific details. There are many articles covering basic techniques and applications of molecular methods to environmental microbiology (e.g. Hurst *et al.*, 1996; Gerhardt *et al.*, 1994).

Generalized approaches to detection and identification of specific phylogenetic groups using molecular methods can include recovery of target DNA or RNA from environmental or cultured samples, extraction and removal of inhibitors, cloning or amplification of DNA or RNA by PCR, followed by hybridization using specific probes. Direct methods include colony lift (e.g. Wright *et al.*, 1996), fluorescent *in situ* hybridization (FISH), and most recently, flow cytometric methods where DNA probes conjugated to specific dyes (Amann *et al.*, 1990) have been used. Probes have been designed to discriminate specific taxonomic groups using 16S rRNA-targeted and functional genes. Many examples of the former are now found in the literature. Probe design, regardless of the target, requires sequence information and extensive validation. Again, numerous examples of probes, design and verification approaches to avoid hybridization with non-target sequences are found in the literature. Quantitative recovery or enumeration of selected targets remains difficult and is subject to uncertainties related to efficiency of recovery of nucleic acids from environmental samples. Approximate quantitative estimates of organism densities in water samples can be achieved by analysis of diluted samples until no further hybridization is detected.

Fluorescent in situ hybridization (FISH). Whole-cell fluorescent *in situ* hybridization or FISH is a non-cultural technique combining attributes of nucleic acid hybridization with various direct detection techniques. For example, fluorochrome-conjugated oligonucleotide probes, designed to target 16S rRNA sequences, can be used to identify individual bacterial cells when visualized by epifluorescence microscopy. Probes can also be designed to detect

selected genes and, in the case of ribosomal RNA, to be species genus or group-specific with detection accomplished using epifluorescence microscopy or flow cytometry (Amann *et al.*, 1990). Limitations inherent in direct count microscopy, i.e. the density of the target microorganism must be approximately $\geq 10^4$ cells/ml to see one organism per field, can be overcome using flow cytometry or increasing the volume of sample filtered. Other problems encountered include background autofluorescence, which obscures the target signal, poor signal intensity, non-specific binding of the probe, probe hybridization to unidentified species, or loss of cells during the hybridization process.

To perform FISH by epifluorescence microscopy, a water sample or suspension is filtered through a black polycarbonate membrane, using the same procedure for a membrane filter direct count. The cells on the filter are then hybridized to a selected fluorochrome-conjugated probe following protocols for *in situ* hybridization. Importantly, FISH detects all target cells including those that may be non-culturable or dead. Maruyama and Sunamara (2000) demonstrated that FISH could be combined with direct microscopic counting (called FISH-DV) by coating filters with poly-L-lysine to promote cell retention during the hybridization process. Following hybridization steps and washings to remove non-hybridized probe, salts and reagents, the cells are stained with a non-specific fluorescent dye such as DAPI and washed. Using an epifluorescent microscope total direct counts are obtained using UV illumination and probe-specific counts with excitation wavelengths specific for the fluorochrome-labeled oligonucleotide probe used, e.g. FITC. Multiple probes conjugated to different dyes can be used simultaneously, providing specific multiple and total direct counts. More recently, Maruyama and Sunamara (2000) suggest FISH can be used in a quantitative fashion. DeLong *et al.* (1999) increased the fluorescent intensity of FISH using polyribonucleotide probes instead of singly labeled oligonucleotide probes.

Detection of selected genes. To avoid the use of cultural methods, which are intrinsically variable in recovery and specificity because of shared physiological requirements, investigators have targeted genes to detect selected species, genera or functionally-defined groups. As with direct counting, these methods usually do not provide information on the viability or even expression of the targeted gene in the cells detected. Examples in the recent literature relevant to environmental microbiology include efforts to use the glucuronidase gene (*uidA* gene) to detect and identify *Escherichia coli*. Green *et al.* (1991) used a fragment of the *uidA* gene as probe to detect *E. coli* after a short incubation period on membranes through which water samples had been filtered. Fricker and Fricker (1994) used PCR primers to detect *E. coli* (*uidA*) and coliforms (*lacZ*) isolated from water samples but encountered specificity problems with the latter primer set. All biochemically verified *E. coli* strains tested were positive for the *uidA* gene and a relatively low false positive rate for the *E. coli* primer set occurred. Bej *et al.* (1994) used PCR for improved sensitivity to amplify a segment of the *himA* gene to detect *Salmonella* spp. in oysters. The cytolysin gene of *V. vulnificus* was the target for an enumeration method based on a colony lift procedure (Wright *et al.*, 1993). Following direct inoculation of a sample on a non-selective solid medium, a piece of absorbent paper is placed over the resulting colonial growth, treated to release and retain the DNA from the lifted colonies, and then hybridized with an alkaline phosphatase-labeled oligonucleotide probe. After hybridization and development, areas on the filter showing alkaline phosphatase activity correspond to *V. vulnificus* colonies.

Lipid analysis. Lipid biomarkers have been applied as a direct detection technique to environmental samples from a variety of environments including soil, aquatic sediments, groundwater, and deep subsurface sediment. These methods are based on the assumption that phospholipid fatty acids and fatty acid methyl esters can be used to

discriminate microorganisms by group (e.g. Gram-negative versus Gram-positive, eukaryotes versus prokaryotes), based on unique fatty acid structure, chain length, double bond position and number and degree of substitution. Other applications include estimates of biomass, detection of specific physiological groups (Parkes *et al.*, 1993) community structure, and physiological status expressed through phospholipid fatty acid profiles (Nickels *et al.*, 1979). (For details related to the principles, methods, and applications see Frostegaard *et al.* (1991) and White *et al.* (1997).) Commercial systems making specific use of fatty acid methyl esters for identification of microorganisms are available. Gas chromatographic analysis of mycolic acids is a powerful tool for identification of environmental mycobacteria (Butler and Guthertz, 2001).

Enzymatic methods. In addition to targeting specific genes to detect bacterial groups on the basis of characteristic enzymes, the presence of enzymes *in situ* can be assayed by addition of substrate to water samples. In such studies, fluorogenic or chromogenic substrates are added and their hydrolysis products detected, providing the basis for a rapid assay. Berg and Fiksdal (1988) and Fiksdal *et al.* (1994) used several fluorogenic substrates including MUG to detect total and fecal coliform bacteria as an alternative to cultural counting. Van Poucke and Nelis (1977) used chemiluminescent substrates in a presence/absence test to detect β -galactosidase- and β -glucuronidase-containing cells in a variety of water samples over relatively short incubation times. Results suggested significant improvements in sensitivity came at the expense of unfavorable false-positive and false-negative errors. George *et al.* (2000) discuss aspects of the current of the literature concerning enzymatic methods for rapid enumeration and notes such methods detect VBNC as well. Enzymatic methods are uniquely suited to remote sensing applications and may be valuable for this purpose.

4 SELECTED ENVIRONMENTAL MICROORGANISMS OR INDICATORS OF PUBLIC HEALTH SIGNIFICANCE AND EXAMPLES OF METHODS USED FOR THEIR DETECTION/ENUMERATION

4.1 Coliform and enterococcal indicators

There is a large and growing body of literature on coliform and enterococcal indicators and whose size is well beyond the scope of this chapter. Readers are referred to such sources as Standard Methods for the Examination of Water and Wastewater (APHA 1998); Manual of Environmental Microbiology (Hurst *et al.* 1996), the USEPA ICR Microbial Laboratory Manual (USEPA, 1996). The USEPA National Exposure Research Laboratory, Microbiological and Chemical Exposure, maintains a web site (<http://www.epa.gov/microbes>) which contains the publications of the most recent approved methods for detection and enumeration of microorganisms of public health significance. Toranzos and McFeters (1996) summarize methods for detection and enumeration of indicators in fresh and drinking waters.

4.2 Alternate indicators and selected environmental pathogens

Although, historically, attention has been focused on the indicators within the Enterobacteriaceae and enterococci, recent exploratory research has focused on the potential use candidate indicators such as *Clostridium*, *Bacteroides*, *Bifidobacterium*, and of bacteriophages as model viral indicators. In recent years other environmental pathogens such as mycobacteria, *Campylobacter* and have been the cause for public health concern. The remainder of this chapter provides examples of alternate indicators and pathogens of interest and methods for detection and enumeration.

4.2.1 *Clostridia*

Clostridia are obligately anaerobic bacteria commonly found in soils and the gastrointestinal tracts of warm-blooded animals including man. Clostridia form very resistant structures called endospores, which enable these microorganisms to survive in air, at elevated temperatures and in the presence of chemical disinfectants. Spores of these organisms can persist for extended periods of time in soils, water and aquatic sediments. Although clostridia are fermentative saprophytes, they are commonly associated with food-borne disease in humans owing to their ubiquity, the high temperature tolerance of their spores, and their ability to elaborate extremely potent endotoxins. Typically, spores germinate in food products that are inadequately processed. *Clostridium perfringens* and *C. botulinum* are common causes of food-borne illnesses associated with cooked and stored meats or poultry and seafoods that are inadequately heated before eating. *Clostridium botulinum* toxin is also associated with mass mortalities of waterfowl (Fay *et al.*, 1965; Wobeser, 1997).

The presence of *C. perfringens* in receiving waters has been linked to contamination by faeces and wastewaters (Cabelli, 1977; Bisson and Cabelli, 1980). Bisson and Cabelli (1980) suggested *C. perfringens* spores have value as an indicator of chlorination efficiency and the presence of unchlorinated sources of fecal contamination, and as a 'conservation tracer' delineating the impact and transport of wastewater effluents. Because *Cryptosporidium* oocysts are also resistant to chlorination, it has been suggested that *C. perfringens* be used as an indicator of drinking water treatment (Payment and Franco, 1993). Detection of vegetative cells in environmental waters reflects fresh and untreated fecal matter because the persistence of vegetative cells is very short (Bisson and Cabelli, 1980). Fujioka and Shizumura (1985) and Roll and Fujioka (1997) used *C. perfringens* to detect wastewater discharge into fresh water streams and brackish waters in Hawaii. Hill *et al.* (1993) confirmed the utility of *C. perfringens* to delineate the extent and movement of sewage

sludge dumped at the deep ocean '106-Mile Site' off the northeast coast of the USA.

Bisson and Cabelli (1980) developed a membrane filtration method for *C. perfringens* recovery applicable to saline waters based on a highly selective medium (mCP). The composition of this medium was later modified to reduce the concentration of a costly biochemical component without compromising its selectivity (Armon and Payment, 1988). Sartory (1986) compared recovery of *C. perfringens* on mCP and egg yolk-free tryptose-sulphite-cycloserine (TSC) agar as membrane based tests. For a variety of sample types, which included polluted river water and groundwater, egg yolk-free TSC was found as selective as mCP and more efficient. Egg yolk-free TSC was recommended because of its ease of preparation, lower cost, and availability of a simple confirmation scheme. Hill *et al.* (1996) reported a false-positive rate of about 5% using mCP to recover *C. perfringens* from deep ocean dump sites. Compared to TSC, Sartory *et al.* (1998) found mCP to possess a significantly higher false positive rate, to be less efficient, and generally more difficult to use with environmental samples.

4.2.2 *Campylobacter spp*

C. jejuni is the species most frequently associated with campylobacteriosis infections in humans and some consider this organism to be the leading cause of bacterial gastroenteritis in the developed world (Skirrow, 1991; Allos and Blazer, 1995). This microorganism is commonly found in the gastrointestinal tracks of domestic pets (dogs, cats), farm animals (cattle, pigs, poultry), and wild animals such as rodents and birds (Kapperud and Rosef, 1983). Campylobacters can be isolated from sewage and are most likely carried to surface receiving waters in runoff following periods of rainfall. Humans and animals can be asymptomatic carriers. Campylobacters discharged into surface waters used for drinking, contaminated and improperly cooked poultry or meat products, and non-pasteurized milk products, have all been implicated as the cause of campylobacteriosis in humans. Campylobacters

have also been found in shellfish beds and market shellfish (Abeyta *et al.*, 1993), but the human health significance of shellfish as a vector remains uncertain.

Campylobacters are microaerophilic organisms with an optimum growth temperature of about 42°C and require complex media supplemented with non-selective constituents, such as whole blood and other compounds (Bolton *et al.*, 1984). Recovery of cells from natural waters is hindered by slow growth rates, competitive background microbiota low numbers, and possible non-culturability (Rollins and Colwell, 1986; Cappelier and Federighi, 1998). Given the complexities for detection of *Campylobacter* spp. from environmental samples, many applications of molecular methods based on PCR exist for its detection in poultry, feces, environmental waters and sewage (e.g. Waage *et al.*, 1999). Other investigators have used immunological-based methods for detection (Buswell *et al.*, 1998). These tools will be valuable in continued efforts to study the persistence of *Campylobacter* cells in natural waters and to understand the health significance of viable and non-culturable states.

4.2.3 Bacteriophages

Human enteroviruses have been implicated as causative agents of waterborne disease transmitted by drinking water and shellfish (IAWPRC, 1991). In the USA, cases of Norwalk virus transmitted through fecally-contaminated shellfish are a recurring concern (Centers for Disease Control, 1995). Routine methods for direct detection of infective enteroviruses, such as Norwalk, from environmental samples remain to be developed and refined. Lacking such methods there has been ongoing interest in the use of bacterial viruses, or bacteriophages, as surrogates for pathogenic enteroviruses, because of the inadequacies of bacteria as indicators of virus. Resistance to chlorination, in particular, is a desirable property of a bacteriophage indicator of treated sewage effluent because hepatitis A and rotavirus are also relatively resistant to chlorination and UV radiation (IAWPRC, 1991). Havelaar *et al.* (1993) reported that concentrations of enteric

viruses and F-specific coliphages were highly correlated in contaminated river and lake waters.

Detection of bacteriophages ideally requires a host whose susceptibility range is restricted to a target bacteriophage associated only with the fecal source. In reality this discrimination is difficult to achieve using wild-type hosts because of the diversity of lytic bacteriophages in the environment.

Bacteriophages most recently studied and proposed as viral indicators of fecal contamination are the F-specific or male-specific coliphage and *Bacteroides fragilis* bacteriophage. The former bacteriophages selectively adsorb to the sex pilus which is usually encoded by a plasmid. Superior hosts sensitive to F-specific coliphages and resistant to antibiotics to reduce the background growth of interfering microorganisms have been developed and tested using a variety of environmental waters (Havelaar and Hogeboom, 1984; Debartolomeis and Cabelli, 1991). These hosts differ somewhat in their host range and avoid lysis by somatic phages using different strategies. WG49 (Havelaar and Hogeboom, 1984) is a *Salmonella typhimurium* selected for lack of susceptibility to somatic coliphage and because there are relatively low numbers of salmonella phage in sewage. It was constructed using an F-plasmid which enables the strain to produce sex pili. Debartolomeis and Cabelli (1991) developed an *Escherichia coli* (HS(pFamp)R) host naturally resistant to somatic coliphage. Using relatively uncomplicated procedures, these host strains are capable of routine detection and enumeration of male-specific coliphage from domestic wastewaters and have also been applied to receiving waters (Havelaar and Hogeboom, 1984; Debartolomeis and Cabelli, 1991). Their application to detect male-specific coliphage in environmental waters and other sample types must be carefully monitored to detect lysis by environmental phage (Rhodes and Kator, 1991; Handzel *et al.*, 1993) or FDNA male-specific coliphage. This can be achieved through parallel processing of plates incorporating Rnase. Alterations to the conventional double-agar-overlay assay have been proposed to

allow for sample concentration and promote ease of use (Sobsey *et al.*, 1990; Sinton *et al.*, 1996).

Owing to the labor and time associated with cultivation of *Bacteroides* spp. and its presumed poor survival in the environment, the use of phages to *B. fragilis* as indicators of human fecal pollution has received considerable attention. *Bacteroides* phages occur in human feces and sewage (Booth *et al.*, 1979; Tartera and Jofre, 1987; Tartera *et al.*, 1989; Grabow *et al.*, 1995) and are unable to replicate in natural waters and sediments due to their reliance on a metabolically active host (Jofre *et al.*, 1986; Tartera *et al.*, 1989). *B. fragilis* phages appear to be highly specific to humans and have not been detected in a variety of domestic animals or captive primates (Tartera and Jofre, 1987; Grabow *et al.*, 1995). A strong correlation between *B. fragilis* phages and enteroviruses in sediments and treated wastewater has been reported (Jofre *et al.*, 1989; Gantzer *et al.*, 1998; Pina, 1998; Torroella, 1998).

Resistance of these phages to disinfectants and environmental factors (Jofre *et al.*, 1986; Bosch *et al.*, 1989; Sun *et al.*, 1997) has also promoted their use as indicators. The frequency of recovery of *B. fragilis* phages from chlorinated drinking water suggests these phages are more resistant than somatic and F-specific phages (Jofre *et al.*, 1995; Armon *et al.*, 1997). Although *B. fragilis* phages are more persistent in seawater under dark conditions than F-RNA phages (Chung and Sobsey, 1993) the former are more susceptible to ultraviolet (Bosch *et al.*, 1989) and sunlight inactivation (Sinton *et al.*, 1999). We have observed similar results for *B. fragilis* and F-RNA phages exposed *in situ* in estuarine water but noted prolonged persistence of both phages in estuarine sediments (Kator and Rhodes, 1992). Quantification of phages to enteric bacteria in freshwater environments indicates that *B. fragilis* phages are more resistant to natural inactivation than male-specific or somatic coliphages (Araujo *et al.*, 1997).

Enumeration of *B. fragilis* bacteriophages involves more complex media and methods than used for coliphage assays. *B. fragilis* phages in sewage and fecal samples can be

enumerated by conventional plaque assay but less contaminated samples may require concentration or enrichment. The double-agar layer (DAL) assay is more efficient than the MPN technique, but both methods are subject to conditions of the host culture, composition of the host medium, presence of divalent cations and decontamination method used (Cornax *et al.*, 1990). Tartera *et al.* (1992) describe an enumeration procedure based on sample decontamination by filtration through polyvinylidene difluoride membrane filters with subsequent filtrate assay by DAL using a rich medium, *Bacteroides* phage recovery medium (BPRM). Kator and Rhodes (1992) combined the use of antichaotrophic salts to promote phage adsorption to nitrocellulose membrane (Farrah, 1982) with a protracted elution process (Borrego *et al.*, 1991) for application to phage recovery from estuarine waters. Lucena *et al.* (1995) describe concentration or enrichment (presence/absence tests) for use with samples containing low phage densities such as drinking water. Samples are concentrated by filtering a large volume (1 litre) through a beef extract-treated membrane of inorganic material with a honeycomb pore structure and low protein binding activity with subsequent elution of phages and assay of the eluate by DAL or, direct DAL assay of phages retained on the membrane. Enumeration by this concentration technique was significantly affected by sample turbidity. Enrichment-based tests provide good recoveries but require removal of oxygen and are non-quantitative (Armon and Kott, 1993; Lucena *et al.*, 1995; Armon *et al.*, 1997).

A critical shortcoming of using *B. fragilis* phages as indicators is the susceptibility range of the host used for recovery. *B. fragilis* HSP40 (Tartera and Jofre, 1987; Tartera *et al.*, 1992; Cornax *et al.*, 1990; Lucena *et al.*, 1995) used to detect phages in Mediterranean seawater has been ineffective in similar studies conducted in temperate climates (Kator and Rhodes, 1992; Bradley *et al.*, 1999). However, in both of the latter studies, *B. fragilis* hosts recovered from local waters were used successfully to isolate bacteriophage. Also problematic is the high specificity of bacteriophages such that host

range among *B. fragilis* strains is restricted (Tartera and Jofre, 1987; Kator and Rhodes, 1992; Bradley *et al.*, 1999).

Infrequent recovery and low densities of *B. fragilis* phages in environmental waters except those proximate to sewage pollution (Cornax *et al.*, 1991; Bradley *et al.*, 1999) may negate the use of this indicator in studies of non-point pollution. However their tendency to adsorb to sediments (Jofre *et al.*, 1986), recovery from sediment samples (Tartera and Jofre, 1987; Jofre *et al.*, 1989; Bradley *et al.*, 1999), and persistence characteristics (Kator and Rhodes, 1992) suggest that sediment analysis might be valuable in determining fecal sources in waters of low to moderate pollution. Lucena *et al.* (1996) demonstrated that persistence of *Bacteroides* phages in sediments contributed to their use as indicators in areas subject to persistent, albeit remote, fecal pollution. These investigators (Lucena *et al.*, 1994) also report that the extended persistence characteristics of *Bacteroides* phages compared with FRNA phages contributes to their usefulness as indicators of human enteric viruses in shellfish.

Some workers suggest that somatic coliphage may be of value (Cornax *et al.*, 1991) as sewage indicators based on observations that somatic coliphages appear to persist longer than FRNA coliphages in natural waters. However, both the early and more recent literature (e.g. Leclerc *et al.*, 2000) suggest there are problems with use of somatic coliphages as fecal or viral indicators. These include significant heterogeneity, broad host range, ability to replicate in the environment, and inconsistent relationships to contaminant sources. In particular, the ability of somatic coliphages to replicate *in situ* or to experience host lysis from indigenous virus in the environment are stumbling blocks to the use of somatic coliphages. Croci *et al.* (2000) compared a variety of indicators, including somatic and FRNA coliphage with enterovirus burden in mussels. They concluded that only direct viral detection is an acceptable method to assess the risk associated with consumption of mussels from the Adriatic Sea.

4.2.4 *Bacteroides* spp

Bacteroides spp. are Gram-negative, anaerobic, pleomorphic rods and represent a major constituent of the colonic flora in normal humans (Jousimies-Somer *et al.*, 1995). In recent years the genus has undergone major taxonomic revisions. The *Bacteroides fragilis* group originally included isolates considered to be subspecies, which have since been assigned species rank based on DNA–DNA homology analyses (Sinton *et al.*, 1998). Consequently early studies referring to the '*B. fragilis* group' as fecal indicators actually may include different species and it is probably more appropriate to use the term *Bacteroides* spp. Interest in *Bacteroides* spp. as fecal indicators is based on their presence at high concentrations in human feces and either absence or occurrence at significantly lower concentrations in most animals (Allsop and Stickler, 1984, 1985; Kreader, 1995). Furthermore as obligate anaerobes, *Bacterioides* spp. are unlikely to multiply in receiving waters and are less persistent in natural waters than *E. coli* (Allsop and Stickler, 1985; Fiksdal *et al.*, 1985). Despite these advantages, complexities associated with cultivation have limited their use as indicators.

The use of prereduced anaerobically sterilized (PRAS) media is superior to prereduced aerobically prepared media for culturing *Bacteroides* spp. (Mangels and Douglas, 1989). A selective and differential medium (WCGP) has been developed that relies on detection of aesculin hydrolysis by *Bacteroides* spp. in the presence of gentamicin and penicillin G, which inhibit other bacteria (Allsop and Stickler, 1984). Recovery of *Bacteroides* spp. by membrane filtration and culturing on WCGP is enhanced by resuscitation initially at a lower temperature. A PRAS agar medium selective for *B. vulgatus* (BVA) has been developed based on this organism's resistance to kanamycin, vancomycin, colistin, ability to grow in the presence of bile, and inability to hydrolyze esculin (Wadford *et al.*, 1995). Isolation of *B. vulgatus* from shellfish meeting the market standard for fecal coliforms suggests it may be a more reliable indicator of fecal contamination

(Wadford *et al.*, 1995) although not necessarily specific for humans since high levels occur in house pets (Kreader, 1995).

Difficulties associated with culturing *Bacteroides* spp. can be overcome by use of alternate detection methods. Culturing has been avoided by enumerating *Bacteroides* spp. with a rapid (2–3 h) fluorescent antibody technique (Fiksdal and Berg, 1987). Species-specific DNA hybridization probes have been used for detection of *Bacteroides* in feces (Kuritzin and Salyers, 1985; Kuritzin *et al.*, 1986). Kreader (1995) increased the sensitivity of species-specific assays using PCR amplification of 16S rRNA gene sequences followed by detection of specific PCR products through hybridization. Although human feces generally yielded high signals, non-humans generally had lower signals with the exception of house pets. The significance of large fecal outputs from farm animals exhibiting weak signals on distinguishing sewage from farm runoff is not known. Subsequent study showed that *Bacteroides distasonis* DNA, detectable by PCR-hybridization assay, disappeared as a function of temperature and predation with detection possible after 2 weeks exposure of human feces in river water (Kreader, 1998). Analysis of 16S rDNA with PCR amplification has been used to detect *Bacteroides* in coastal lagoons (Benlloch *et al.*, 1995). Bernhard and Field (2000a,b) used host-specific 16S rDNA genetic markers and PCR to discriminate human and ruminant feces and for identification of non-point sources of fecal pollution.

4.2.5 Fecal streptococci

The term 'fecal streptococci' has been used to describe a group of taxonomically distinct streptococci that are Gram-positive, catalase negative, non-spore-forming, facultative anaerobes recoverable from the gastrointestinal tracts of humans and animals. This group has been considered as an indicator of fecal contamination in environmental waters. They are present in relatively high densities in human and animal feces, in sewage, and it is generally believed they do not grow and multiply in environmental waters and soils. Recent revisions in the taxonomy of this

'group' have resulted in some members of the group being placed into a new genus called *Enterococcus*. Members of this genus include *E. avium*, *E. casseliflavus*, *E. durans*, *E. faecalis*, *E. faecium*, *E. gallinarum*, *E. hirae*, *E. malodoratus*, and *E. mundtii*. These species usually grow at 45°C in 6.5% NaCl, and at pH 9.6; most grow at 10°C. *S. bovis* and *S. equinus*, fecal streptococci generally associated with animal feces, which are negative for two or more of these characteristics, are retained in the genus *Streptococcus*. Consequently, some species previously considered as fecal streptococci are now enterococci.

Early studies identified the ratio of fecal coliforms to fecal streptococci as useful for differentiation of human from animal fecal pollution sources (Geldreich and Kenner, 1969) but this assumption now appears largely unwarranted. Although it appears that animal gastrointestinal contents and feces have proportionately more fecal streptococci to fecal coliforms than humans, the ratio is not stable and varies with exposure to natural environments. Differential survival of both enterococcal and streptococcal components of fecal streptococci in aquatic environments will occur and further complicates a straightforward interpretation of 'fecal streptococcal' indicator levels. Source specificity is also an issue with this indicator because some fecal streptococcal strains are associated with non-fecal plant, insect, reptile and soil habitats. The presence of these 'atypical' strains in the environment can complicate a straightforward interpretation of the sanitary significance of this group (Mundt *et al.*, 1958; Mundt, 1963; Geldreich *et al.*, 1964; Geldreich and Kenner, 1969). Recent interest in the enterococci (a 'subgroup' of fecal streptococci) as an indicator derives from the USEPA adoption of an epidemiological-based enterococcus criterion for recreational marine and freshwaters waters (USEPA, 1986). Excellent discussions surrounding the use of faecal streptococci as an indicator enumeration and associated problems can be found in Sinton *et al.* (1993a,b; 1994).

Generally, the fecal streptococcal habitat is the intestinal content of both warm- and

cold-blooded animals, including insects. None of the enterococci can be considered as absolutely host specific, although some species evidence a degree of host specificity. Although certain genera (e.g. *E. faecalis*, *E. faecium*) have been considered specific to feces from human or other warm-blooded animals, phenotypically similar strains and biotypes can be isolated from other environmental sources (Mundt, 1982; Clausen *et al.*, 1977; Beaudoin and Litsky, 1981). Strains and biotypes of *E. faecalis* and of *E. faecium* can be isolated from a variety of plant materials, reptiles and insects. In comparison, the distribution and viability of *S. bovis* and *S. equinus* in extra-enteral environments appears restricted.

Functionally, 'fecal streptococci' as an indicator are, in effect, defined on the same operational basis as fecal coliforms, i.e. organisms recovered using a given method. The selectivity and specificity of available methodologies both define and complicate the meaning of the term 'faecal streptococci' because not all streptococcal and enterococcal species with a fecal habitat are recovered using any one technique.

Several media have been suggested for the selective isolation and/or enumeration of the fecal streptococci or the enterococci. For the most part, the media and methods that are available presently lack selectivity, differential ability, quantitative recovery, relative ease of use, or a combination of these deficiencies. Some strains of fecal streptococci from anaerobic environments, for example, initiate growth only in the presence of elevated levels of CO₂ until the cultures have been adapted to an aerobic environment. *E. faecalis* and *E. faecium* are the most common enterococci encountered. This undoubtedly reflects the rationale of employing KF streptococcal agar for the estimation of enterococci in foods. m Enterococcus agar is used most often for water.

Validation of the enterococci as an indicator of public health risk in marine waters will require improved recovery methods. The range of selectivity and specificity characteristics of methods for enumeration of enterococci has been mentioned. Current recovery methods require confirmatory testing of presumptive

isolates (Ericksen and Dufour, 1986), increasing the time and cost of analysis. Although the mE-based method has been applied to marine and estuarine waters (Cabelli *et al.*, 1983), its utility in non-point source impacted marine and shellfish growing areas is undetermined. The relative occurrence and densities of non-fecal biotypes of *E. faecalis* and *E. faecium* in these waters should be determined. Rapid methods for confirmation of selected enterococcal species, based on serological or biochemical characteristics are needed. Bosley *et al.* (1983) and Facklam and Collins (1989) describe useful and rapid (4 h) identification schemes to separate enterococci from group D non-enterococci based on hydrolysis of L pyrrolidonyl- β -naphthamide (PYR). A colony hybridization method, employing oligonucleotide probes synthesized for specific sequences of 23S rRNA of selected enterococci, was used successfully to detect and identify *E. faecalis*, *E. faecium* and *E. avium* in mixed culture (Betzl *et al.*, 1990).

4.2.6 Non-tuberculosis mycobacteria

In addition to the well-known obligate pathogens, *Mycobacterium tuberculosis* and *M. leprae*, the genus *Mycobacterium* also includes free-living saprophytes that for almost a century received little recognition with regards to public health (Covert *et al.*, 1999). These non-tuberculosis mycobacteria (NTM) are ubiquitous, occurring in a variety of soil and water environments as well as in diseased animals and humans. During the last several decades the clinical significance of these species has become widely acknowledged (Wolinsky, 1992; Falkinham, 1996; Horsburgh, 1996) and water recognized as an important vehicle of transmission (Collins *et al.*, 1984; Covert *et al.*, 1999; Dailloux *et al.*, 1999). Ecological studies of NTM in diverse natural waters show that densities and distribution vary geographically and according to physicochemical characteristics (Kirschner *et al.*, 1992; Iivanainen *et al.*, 1993, 1999; Dailloux *et al.*, 1999). In addition to exposure to mycobacteria indigenous to natural aquatic habitats, individuals are exposed to these organisms in engineered aquatic environments such as treated drinking water systems.

Mycobacteria, relatively resistant to numerous disinfectants including chlorine (von Reyn *et al.*, 1993, 1994; Covert *et al.*, 1999) are particularly persistent in drinking water systems (Dailloux *et al.*, 1999) and are more frequently isolated in water supplies in developed countries (von Reyn *et al.*, 1993).

During recent years, non-tuberculosis mycobacterial infections have increased in actual numbers and in the proportion of total mycobacterial disease, with the highest incidence attributed to the *M. avium* and *M. intracellulare* (MAI) complex (Wolinsky, 1992). Disseminated infections frequently occur in patients with acquired immunodeficiency syndrome and can dramatically affect survival (Horsburgh and Selik, 1989; Horsburgh, 1996). The environment is considered the prime source of these free-living strains and ingestion of contaminated water is a mechanism that could explain severe intestinal infections in AIDS patients (Wallace, 1987; Wolinsky, 1992). One study definitively showed that a human infection by *M. avium* occurred via contaminated potable water (von Reyn *et al.*, 1994). However, overall specific reservoirs and factors affecting human exposure leading to infection have not been defined (Horsburgh, 1996).

The MAI complex can produce a variety of chronic non-tuberculosis mycobacteriosis in both animals and humans, including infections characterized by intestinal colonization. Swine have been implicated as vehicles of infection to humans based on the occurrence of MAI strains of the same serotypes (Ikawa *et al.*, 1989) and similar genotypic characteristics (Bono *et al.*, 1995). *M. avium* and *M. avium* subsp. *paratuberculosis* infections in ruminants and ungulates such as cattle, goats, sheep, deer, and elk cause chronic granulomatous enteritis (Cocito *et al.*, 1994; Essey and Koller, 1994) and result in massive numbers of mycobacteria, e.g. 10^8 organisms per g feces (Whittington *et al.*, 2000). Recently the isolation of *M. avium paratuberculosis* from feces of patients with Crohn's disease and ulcerative colitis has implicated this organism as the etiologic agent (Del Prete *et al.*, 1998). Although members of the MAI complex are also the causative agents of avian tuberculosis in wild birds and

domestic poultry, recovered strains differ significantly from those isolated from pigs and humans (Grange *et al.*, 1990; Bono *et al.*, 1995).

Despite the recognition that animals are potential vectors for transmitting mycobacteriosis to humans, concerns have been focused primarily on food-borne or aerosol transmission from animals (Falkinham, 1999; Yoder *et al.*, 1999). Waterborne transmission of mycobacterial infections, primarily *M. avium*, has focused on the aerolization of natural waters and treated water as well as ingestion of the latter (Falkinham, 1999). Public health risks associated with animal slaughterhouse wastewater, stormwater associated with feedlots and intensive animal holding facilities have received little attention.

Recovery of mycobacteria from water samples typically involves three steps: concentration, decontamination and cultivation. Many mycobacteria grow slowly, requiring weeks or months to form visible colonies in culture and their detection can be compromised by background overgrowth. A variety of disinfectants, e.g. 2% HCl, 2% NaOH, and 0.3% Zephiran have been used (Jaramillo and McCarthy, 1986; Schulze-Robbeke *et al.*, 1991; Kamala *et al.*, 1994; Neumann *et al.*, 1997) to eliminate background microbiota, although these chemicals frequently are detrimental to mycobacterial recovery. Our laboratory studies have shown that treatment of aqueous suspensions of pure cultures with widely used decontaminants result in several log reductions after exposure periods routinely used. Observations that mycobacterium species were differentially affected indicates that a particular chemical decontaminant would 'select' for specific species and thus 'bias' the composition of the original sample. A variety of culture media and incubation conditions exist which, likewise, will also affect the types of mycobacteria recovered. It is generally accepted that reported mycobacterial densities are underestimates due to these methodological factors.

Although molecular techniques have been used to detect mycobacteria from diseased fish (Knibb *et al.*, 1993; Talaat *et al.*, 1997) and clinical samples (Crawford, 1999) detection of mycobacteria in natural water samples has

been pursued primarily by cultivation (Dailloux *et al.*, 1999). Application of molecular methods developed for detection of bacterial DNA in environmental waters (Belas *et al.*, 1995) to detection of mycobacteria is needed to facilitate ecological and epidemiological studies to increase our understanding of these microorganisms and their transmission via the water route.

4.2.7 *Streptococcus bovis*

Streptococcus bovis is a non-enterococcal, group D streptococcus. High densities occur in feces of ruminants (Wheater *et al.*, 1979) and other domestic animals including dogs, cats, horses and pigs (Clausen *et al.*, 1977; Kenner *et al.*, 1960). Recent investigations demonstrated that feral mammals typically resident in tidal and freshwater marsh habitats (deer, muskrat, raccoon) are sources of both *E. coli* and *S. bovis* (Kator and Rhodes, 1996 unpublished results). Because of its association with animals, *S. bovis* has been proposed as a specific indicator of animal pollution (Cooper and Ramadan, 1955). However, this organism is not unique to animals and is present in the intestines of healthy (11%) and immunocompromised (up to 56%) humans, particularly those with neoplasias of the gastrointestinal tract (Klein *et al.*, 1977; Beebe and Koneman, 1995). *S. bovis* can cause disease, e.g. endocarditis, bacteremia septic arthritis and meningitis in compromised hosts and less frequently in healthy individuals (Klein *et al.*, 1977; Muhlemann *et al.*, 1999; Grant *et al.*, 2000). Bacteremia in patients with colonic neoplasia and bacterial endocarditis has been attributed primarily to *S. bovis* biotype I (Ruoff *et al.*, 1989), although biotype II also is associated with pathological conditions (Cohen *et al.*, 1997; Grant *et al.*, 2000).

Oragui and Mara (1981) developed a selective medium for recovery by membrane filtration, membrane-bovis agar, for the recovery of *S. bovis* based on its ability to utilize NH_4^+ as the sole source of nitrogen and the absence of a requirement for exogenous vitamins when cultured under anaerobic conditions. The original medium formulation was modified (Oragui and Mara, 1984) to contain less sodium azide because of inhibitory effects on *S. bovis*

strains from different geographical regions. Although the specificity of the modified medium was 96% when applied to sewage and surface water samples (Oragui and Mara, 1984) lower specificity, 35–47% was observed for samples from freshwater feeder streams and estuarine shellfish growing areas in a different geographic region (Kator and Rhodes, 1991, 1996). The modified medium was not sufficiently selective to inhibit growth of *Enterococcus faecalis* (Oragui and Mara, 1984; Kator and Rhodes, 1991).

Other selective techniques have incorporated fluorogenic substrates to detect *S. bovis*. A medium selective for fecal streptococci provides for the differentiation of *S. bovis* based on its ability to hydrolyze both a fluorogenic galactoside and a chromogenic starch substrate (Little and Hartman, 1983). However strains unable to hydrolyze starch (Oragui and Mara, 1981; Kator and Rhodes, 1991) could lead to an underestimation of *S. bovis* densities using this medium. A novel miniaturized method for the specific detection of *S. bovis* as well as *E. coli* and enterococci is based on the ability of the former to hydrolyze a fluorogenic substrate and its inability to reduce triphenyltetrazolium chloride (TTC) (Pourcher *et al.*, 1991). Enumeration is based on the most-probable-number calculation using microtiter plates (16 wells per dilution).

Distinguishing ruminal from human strains of *S. bovis* has been problematic, relying on biochemical characteristics such as mannitol and melibiose fermentation, growth on starch and amylase activity (Knight and Shales, 1985; Nelms *et al.*, 1995). Recently a PCR technique has been published (Whitehead and Cotta, 2000) to differentiate between human and ruminal strains of *S. bovis* based on 16S rRNA gene sequences (Nelms *et al.*, 1995). Application of molecular techniques to cultural methods could provide for rapid and specific enumeration of *S. bovis* and avoid reliance on time consuming and laborious biochemical verification. For example, related organisms such as *Streptococcus alactolyticus*, *Streptococcus saccharolyticus* and *Enterococcus columbae* would not be identified readily using routine procedures (Knudtson and Hartman, 1992) and

require additional biochemical testing, DNA–DNA hybridization studies or 16S rRNA analysis (Farrow *et al.*, 1984; Devriese *et al.*, 1990; Rodrigues and Collins, 1990). Furthermore, reliance on conventional and commercially available miniaturized identification techniques may be misleading. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis analysis of whole-cell proteins of isolates previously identified as *S. bovis* showed that those from human and animal infections were predominantly *Streptococcus gallolyticus* (Devriese *et al.*, 1998), a newly described species originally detected in the feces of koalas (Osawa *et al.*, 1995). Recently ruminal tannin-tolerant streptococci have been isolated from a variety of animals and, on the basis of 16S rRNA, are most closely related to ruminal strains of *S. bovis* and *S. gallolyticus* (Nelson *et al.*, 1998). Such complexities underscore the advantages of using molecular probes to identify and distinguish specific *S. bovis* strains unique to human and animal sources (Whitehead and Cotta, 2000).

Indirect detection of ruminal *S. bovis* by using bacteriophage has been proposed but does not appear promising. Although restriction endonuclease digestion patterns of the PCR amplified 16S rRNA gene of *S. bovis* strains isolated from a variety of ruminants indicated genetic homogeneity isolated lyso-genic phages were highly strain specific (Klieve *et al.*, 1999). Previous investigations by others had shown that the host range of lytic *S. bovis* phages was limited (Klieve and Bauchop, 1991; Tarakanov, 1996).

Relative to coliforms and enterococci, *S. bovis* dies off more rapidly in aquatic environments (Geldreich and Kenner, 1969; McFeters *et al.*, 1974; Wheeler *et al.*, 1979). *In vitro* experiments conducted at 6°C and 25°C using an isolate of *S. bovis* demonstrated poor survival in filtered fresh and estuarine water, particularly at the higher temperature (Kator and Rhodes, 1991). Persistence experiments (Kator and Rhodes, unpublished data) using non-filtered fresh and estuarine water and conducted *in situ* at 10°C using light transmitting plastic bags, exposed to ambient sunlight immediately below the water surface, showed reduction of four orders

of magnitude to undetectable levels within 3 days. Corresponding survival in the dark was enhanced with less than one and two orders of magnitude decrease in fresh and estuarine waters, respectively. Despite poor persistence in environmental waters, we observed prolonged survival of *S. bovis* in deer and muskrat scat exposed in mesh bags at the estuarine intertidal zone with little or no change in concentrations after 1 month of weathering. These observations suggest that animal fecal matter can serve as reservoir of *S. bovis* (and other enteric microorganisms) in watersheds and that, owing, to its poor survival in environmental waters, its detection indicates recent input.

4.2.8 Iron oxidizing bacteria

The *Sphaerotilus-Leptothrix* group of organisms, also known informally and collectively as ‘sewage fungus’, are sheathed bacteria whose filamentous appearance in contaminated surface waters appears ‘fungus-like’. In reality, the filaments are sheaths containing single rod-shaped cells which are Gram-negative, obligate aerobes that divide by binary fission, and do not branch. These bacteria also tend to accumulate poly-hydroxybutyrate which can be seen as inclusions in the cells. The sheaths are composed of protein-polysaccharide-lipid complex (van Veen *et al.*, 1978) and are covered with an exopolysaccharide or slime layer that is associated with deposits of iron or manganese oxides. Whether the oxidized iron associated with *Sphaerotilus natans* represents the outcome of an energy-deriving process remains unconfirmed. This group of microorganisms is important in the cycling of manganese and iron in natural waters.

Waters polluted by sewage contain high concentrations of organic matter and can support indicative populations of filamentous bacteria belonging to the genera *Sphaerotilus* or *Leptothrix*. These bacteria oxidize reduced iron and deposit its oxidized form in conjunction with their mucilaginous filaments. van Veen *et al.*, (1978) recommended isolation by collecting samples of the gray-brown sheaths from a colonized surface and mixing to separate the sheaths so that small pieces of the sheathed

cells can be transferred to a dried agar plate containing a low-nutrient medium. Various enrichment media are available for enrichment from water samples (e.g. APHA, 1998). Microscopic examination of water samples, cultures, flocs scraped from rocks or pipes, or material captured on filter surfaces can facilitate identification of these organisms based on their characteristic morphology and encrustations. From a culture-independent perspective, Siering and Ghiorse, (1997) applied 16S rRNA-targeted fluorescein-labeled oligonucleotide probes for detection of *Leptothrix* and *S. natans* by FISH.

4.2.9 *Staphylococci*

Disease symptoms frequently associated with exposure to recreational waters, including swimming pools, are complaints of ear, eye, nose, throat and skin conditions which are of non-enteric etiology. Consequently, indicators such as the staphylococci (Favero, 1985; Borrego *et al.*, 1987; Charoenca and Fujioka, 1993) and *Pseudomonas aeruginosa* (Seyfried and Cook 1984) have been proposed for use as indicators of health risk in recreational waters. Borrego *et al.* (1987) and Charoenca and Fujioka (1993) evaluated a variety of media for detection of *Staphylococcus* spp. Charoenca and Fujioka (1993) subsequently developed a selective medium for use with membrane filtration. The medium was Vogel-Johnson agar and the authors were able to relate numbers of staphylococci to swimmer density in marine waters. Moreover, Charoenca and Fujioka (1993) detected staphylococci in beach waters at night when swimmers were absent implying an ability of these organisms to persist in saline waters. The incidence of staphylococci in beach waters, their association with clinical conditions, and the possibility of persistence, supports the need for continued study of the staphylococci as pathogens in recreational waters.

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Fundamentals of biological behaviour and wastewater strength tests

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1 INTRODUCTION

1.1 Definition of pollution

One definition for pollution is, 'an undesirable in change in the physical, chemical or biological characteristics of our air, land and water that may or will harmfully affect human life or that of desirable species, our industrial processes, living conditions, and cultural assets; or that may waste or deteriorate our raw material resources' (Odum, 1971). However, whatever definition of *pollution* one may choose, all imply that pollution is, (1) implicitly connected with life, and (2) disrupts an existing life order. Nature knows no pollution. Rather, it has a built-in propensity to respond to any stimulation and dispassionately absorbs it. From our human perspective, the response may be beneficial or adverse: if adverse we call the stimulation pollution. Although it is hard in this first decade of the 21st century to conceive a stimulation that is not somehow connected to human activity, the stimulation need not arise from mankind's activities but may be completely natural.

1.2 The role of energy and matter

From a temporal point of view, life requires a continuous throughput of *energy* and *matter* to

maintain itself. For any system (e.g. living organism or ecosystem), the energy is received at a high level and leaves at a lower level, the difference being the energy utilized. Any inefficiency in energy utilization leads to energy being lost as heat. The required continual input of energy arises from the laws of the created order: during any natural process, in the absence of life an isolated system will move naturally towards a state of disorder, where the molecules, atoms and elementary particles are arranged in the most random manner. This is called an increase in *entropy*. However, the living cell (or any other living system, e.g. organism, ecosystem) is anything but random, being a highly ordered structure.

To counter the natural tendency towards disorder, the living cell must continually take in energy from the surroundings. If no energy is available the cell will deteriorate and die. Thus, the cell is constantly tending towards degradation and energy is constantly needed to restore structures, as they are degraded; cells need to maintain structure in order to function. Furthermore, energy is also required for cell reproduction and the various cell functions (e.g. motility). Thus to maintain and reproduce life, a continuous energy flow is required. Matter is inextricably bound up with this energy flow. The matter passing through living systems is principally the macronutrients

carbon (C), hydrogen (H), oxygen (O), nitrogen (N) and phosphorus (P), but a variety of organic and inorganic micronutrients also take part. If the sources of energy, or matter, diminish, the quantity of life in the sense of mass of living organisms that can be supported diminishes correspondingly as happens in famine and drought, and vice versa.

1.3 Categorization of organisms

Energy is principally derived from three sources and carbon from two sources, and these form convenient criteria to categorize organisms, in particular the microorganisms involved in water quality control and wastewater treatment:

1.3.1 Energy

1. Sunlight radiation – *phototrophs*
2. Organic compounds – *heterotrophs*
3. Inorganic compounds – *lithotrophs*

1.3.2 Carbon

1. Organic compounds – *heterotrophs*
2. Inorganic compounds – *autotrophs*

In the biosphere, the basic source of energy is solar radiation. The photosynthetic autotrophs, e.g. algae, are able to fix some of this solar energy into complex organic compounds (cell mass). These complex organic compounds then form the energy source for other organisms – the heterotrophs. The general process whereby the sunlight energy is trapped, and then flows through the ecosystem, i.e. the process whereby life forms grow, is a sequence of reduction–oxidation ('redox,' reactions). In chemistry the acceptance of electrons by a compound (organic or inorganic) is known as reduction and the compound is said to be reduced, and the donation of electrons by a compound is known as oxidation and the compound is said to be oxidized. Because the electrons cannot remain as entities on their own, the electrons are always transferred directly from one compound to another so that the reduction–oxidation reactions, or electron

donation–electron acceptance reactions, always operate as couples. These coupled reactions, known as redox reactions are the principal energy transfer reactions that sustain temporal life. The link between redox reactions and energy transfer can be best illustrated by following the flow of energy and matter through an ecosystem.

1.4 Redox reactions

1.4.1 Phototrophs

The photosynthetic autotrophic organisms, e.g. algae, grow and thereby fix some of the solar energy into organism mass comprising complex organic compounds; oxygen is a byproduct of this process. Of the mass requirements for the photosynthetic process, the hydrogen and oxygen are obtained from water, the carbon from carbon dioxide and phosphorus and nitrogen from dissolved salts of these elements. While carbon dioxide and water are readily available in the biosphere, the remaining two macronutrients, nitrogen and phosphorus, may be limited. Nitrogen (N) is useful to organisms in the nitrate or ammonia forms that may be limited and phosphorus (P) does not readily occur in its useful soluble phosphate form. The restricted availability of the N and P compounds usually limits the mass of photosynthetic autotrophic life a body of water can sustain. For this reason N and P are called eutrophic¹ or life nourishing substances and the excessive growth of algae or water plants in response to the N and P nutrients is called eutrophication. Whereas phosphorus is usually limited by the mass available that enters an ecosystem, certain microorganisms can generate ammonia from dissolved nitrogen gas (called N fixation). For this reason availability of phosphorus in a body of water is usually more crucial in eutrophication than

¹ From the Greek 'eu' meaning life and 'trophikos' meaning nourishment. Trophic and tropic should not be confused: they are distinctly different: tropic (from Greek 'tropikos' meaning turning towards) means attracted to, which is the reason why domestic flies are called phototropic, i.e. attracted to light, an important feature in the design of VP pit latrines: phototrophic means nourished by light.

the availability of nitrogen. Limitations in the availability of phosphorus will limit the amount of life (algae, plants, etc.) the ecosystem can support through photosynthesis. Accordingly, strategies to control eutrophication are usually focused more on limiting the discharge of P than N to the aqueous environment. Thus in photosynthesis, the simple low energy building blocks carbon dioxide and water are transformed to high energy complex organic compounds (algae, plants), the energy for this provided by sunlight radiation. Although there are some photosynthetic organisms, known as photosynthetic heterotrophs, that can use simple organic molecules as carbon source instead of carbon dioxide this is not common. The photosynthetic process is accomplished by redox reactions in which water is oxidized, producing oxygen as a by-product and the electrons (and protons) are used to reduce carbon dioxide to form complex organic compounds.

1.4.2 Heterotrophs

The complex organic compounds formed through photosynthesis (i.e. plant matter) are the source of matter and energy for the heterotrophs. This is a diverse group of organisms, which, given sufficient time and appropriate environmental conditions, will utilize every type of organic material. The group is ubiquitous and in any given situation those members of the group that derive maximum benefit from the specific organic matter and environmental conditions will develop. As the organic source or environmental conditions change, so associated changes in the heterotrophic organism species take place.

To obtain energy from complex organic compounds, the heterotrophs degrade these in a sequence of biochemical pathways. In the redox reactions accompanying this degradation, electron transfer proceeds in the opposite direction to that for photosynthesis. The electrons originally captured in photosynthesis are now donated by complex organic compounds and are passed to an electron acceptor. For example, if the reaction is aerobic the electrons are passed from the complex organic compound to oxygen, forming water. In the

process, the original sunlight energy captured in the complex organic molecules by the phototrophs is released and becomes available for use by the particular organism executing the degradation reaction. A number of different types of degradation reactions occur.

1.5 Electron donors and acceptors

In an aquatic environment, if dissolved oxygen is available in the water, the heterotrophic organisms will be aerobic and degrade some of the organic compounds to carbon dioxide and water and consume some of the available oxygen. In this process, more heterotrophic organisms are formed so that some of the energy (and matter) contained in the high-energy organic compounds (algae) is transformed to heterotrophic organism mass and retained as different organic compounds in the ecosystem. The remainder of the energy (about one-third) is lost as heat. In these processes, the organic compound supplies the electrons and is oxidized and the function of the oxygen is to accept the electrons released and thus become reduced. For this reason the organic compound is called the electron donor and the oxygen is called the terminal electron acceptor. Should the dissolved oxygen become depleted and anoxic conditions develop (no dissolved oxygen present), then some of the heterotrophic organisms can utilize nitrate instead of oxygen as the terminal electron acceptor, reducing the nitrate to nitrogen gas. In this way these organisms continue their redox reactions and grow under anoxic conditions. Should both dissolved oxygen and nitrate become depleted and sulphate is present, then certain microorganisms, the sulphate reducers, utilize sulphate as the terminal electron acceptor. The sulphate is reduced to sulphide and hydrogen sulphide gas, which not only has a foul smell but also is toxic to many aquatic life forms. Fortunately, sulphides are relatively insoluble ($pK_a \sim 7.1$) and form a black precipitate with many metals. However, the odours and black appearance in a natural water body are aesthetically unpleasant. The organisms that are unable to reduce sulphate have no external electron acceptors available.

If high-energy organic compounds are still available in the water, the organisms have to generate their own electron acceptor internally in order to gain energy through redox reactions.

1.6 Defining aerobic and anaerobic processes

As a general definition, with organic compounds as the electron donor and when the terminal electron acceptor is available externally, the redox reactions are called respiration. The nature of the terminal electron acceptor defines the kind of respiration; if this is oxygen, it is called aerobic respiration, if not oxygen it is called anaerobic respiration². When the terminal electron acceptor needs to be generated internally to the organisms, the redox reaction is called fermentation. In fermentation both the electron donors and acceptors are organic compounds; e.g. in fermentation of glucose to lactate, the glucose is first oxidized to pyruvate, the electrons then being used to reduce the pyruvate to lactate.

1.7 Predation

The mass of heterotrophic organisms generated via the different degradation reactions detailed above, in turn form the matter and energy sources (prey) for other organisms that live on them (predators), and these in turn become prey to yet other predators. Each predator–prey transformation is accompanied

by a substantial loss of energy, lost as heat, due to inefficiencies in energy transfers. In this fashion, through the sequential chain of life there is a continuous reduction of the total organic energy originally fixed by the photosynthetic autotrophs, which in turn is dependent on the quantity of the available macronutrients nitrogen (N) and phosphorus (P). When the organic energy reduces to zero, heterotrophic life ceases. If the quantity of organic energy is small, it can be reduced to zero while the conditions remain aerobic; however, if the quantity is large, the conditions in the water body can become anaerobic.

1.7.1 Lithoautotrophs

The last of the three groups of organisms categorized by energy source and termed the lithotrophs are, like the heterotrophs, a diverse group. Of importance in wastewater treatment are the lithoautotrophs, more commonly called autotrophs. Interesting species in this group are the sulphur, iron, nitrifying and hydrogenotrophic methanogenic organisms. The sulphur and iron organisms are important in sewer and pipe corrosion. They are all obligate (strict) aerobes, e.g. *Thiobacillus* (*T. thiooxidans* and *T. ferrooxidans*) and have found wide biotechnological application in the bioleaching of ores such as iron from pyrites (FeS). In bioleaching, these organisms utilize the S from iron sulphide (FeS) as electron donor and oxygen as electron acceptor, with the S being oxidized to sulphate (SO_4^{2-}). The same reaction occurs in the condensate in the top gas space of sewers except in this case the electron donor is hydrogen sulphide (H_2S) gas which is oxidized to sulphate. The sulphates so produced generally form sulphuric acid (H_2SO_4) resulting in a very low pH between 1 and 2 which corrodes the top of the sewers. Surprisingly, the low pH values are not detrimental to the *Thiobacillus* organisms provided the pH value > 1; indeed it is their preferred pH. The H_2S gas is generated from sulphate originally present in the wastewater. The sulphate is used as electron acceptor by anaerobic heterotrophic sulphate reducers (e.g. by *Desulfovibrio desulphuricans*) with organics in the wastewater

² These definitions are those generally used in bacteriology and microbiology. The definitions conventionally used in sanitary engineering are somewhat different in that respiration in an absence of dissolved oxygen but a presence of oxidized nitrogen, nitrite or nitrate, is called anoxic respiration and that in an absence of oxygen and oxidized nitrogen but presence of sulphate is called anaerobic respiration. Furthermore, in biological nutrient removal (BNR) activated sludge systems, the term anaerobic has a different meaning from that used in microbiology. In BNR it means an absence of *both* dissolved oxygen and oxidized nitrogen and not just absence of oxygen. Actually, in BNR, the definitions of anoxic and anaerobic go further in that they include zero input of dissolved oxygen (anoxic) and zero input of dissolved oxygen and oxidized nitrogen (anaerobic).

serving as electron donors; the sulphate is reduced to hydrogen sulphide in the liquid wastewater phase, some of which comes off as a gas and redissolves in the condensate of the top gas space.

Interestingly, both lithoautotrophic *Thiobacillus* species are also capable of utilizing iron as electron donor, oxidizing ferrous (Fe^{2+}) to ferric (Fe^{3+}) while reducing oxygen to water, but the reaction requires the presence of sulphate. This feature of these organisms is therefore an added danger for metal pipes transporting water and wastewaters. Chlorination of municipal water supplies assists in providing protection against these bioreactions. A third *Thiobacillus* species, *T. denitrificans*, can utilize reduced sulphur, such as S or sulphides (H_2S , S^{2-}), as electron donor and nitrate or nitrite as electron acceptor, with the reduced sulphur being oxidized to sulphate and the oxidized nitrogen being reduced to nitrogen gas – a lithoautotrophic denitrification.

While the *Thiobacillus* autotrophs are important in water and wastewater transport systems, only the nitrifying and hydrogenotrophic methanogen lithoautotrophs are of particular importance in municipal wastewater treatment systems. The nitrifiers are strict (obligate) aerobes and the organisms of importance in wastewater treatment systems are generically called Ammonia Oxidizers (AO) and Nitrite Oxidizers (NO). Being autotrophs, the carbon they require to form cellular material is obtained from carbon dioxide like the algae, but their energy requirements are obtained from oxidizing ammonia to nitrite and nitrite to nitrate, respectively – reactions collectively called nitrification. The ammonia oxidizers, e.g. *Nitrosomonas* and *Nitrosococcus* spp., ammonia (NH_4) as an electron donor, and oxygen as an electron acceptor. The ammonia is oxidized to nitrite and the oxygen is reduced to water. The nitrite oxidizers, e.g. *Nitrobacter* and *Nitrospira* spp., which are usually encountered together with the ammonia oxidizers, use nitrite as electron donor, oxidizing it to nitrate with oxygen serving as the terminal electron acceptor. These nitrifying organisms can only execute these redox reactions while dissolved oxygen is present and as a consequence they

are obligate aerobes. In contrast, the hydrogenic methanogens are obligate anaerobic organisms (oxygen is toxic to these organisms). To obtain energy they use hydrogen as electron donor and carbon dioxide as an electron acceptor to form methane. Being autotrophs, carbon requirements to form cell material are obtained from carbon dioxide. The organisms are of importance in anaerobic digestion systems, e.g. in digestion of primary and waste activated sludges generated in the treatment of municipal wastewater and in upflow anaerobic sludge bed (UASB) systems.

1.8 Anabolism, catabolism and yield

From the above discussion, a fundamental difference between the heterotrophic organisms is that the former uses organic compounds as electron donor, whereas the latter uses inorganic compounds like sulphides, ammonia or nitrite. Since for the lithotrophs the electron donor does not include carbon, they have to obtain the carbon for building the organic compounds of cell material from molecules separate from the electron donor. Of importance are the lithoautotrophs that obtain carbon from carbon dioxide dissolved in the water. As a consequence, for the lithoautotrophs the energy requirements are derived from reducing the electron donor and different inorganic compounds supply the material requirements. The redox reactions that supply the energy requirements are called *catabolism* and those that supply the material requirements are called *anabolism*. Catabolism and anabolism together are called *metabolism*. In contrast to the lithoautotrophs, the heterotrophs obtain the energy requirements (catabolism) and material requirement (anabolism) from the same organic compounds irrespective of the type of external terminal electron acceptor. This difference in the metabolism of the autotrophic and heterotrophic organisms is the principal reason why the cell yield, (i.e. the organism mass formed per electron donor mass utilized) is different. It is low for autotrophs (e.g. 0.10 mg VSS/mg NH_4 – N nitrified for the nitrifiers) and high for heterotrophs (e.g. 0.45 mg VSS/mg COD utilized). Not only is

the energy requirement in anabolism to convert carbon dioxide to cell mass far greater than that required to convert organic compounds to cell mass, but also the energy released (catabolism) in oxidizing inorganic compounds is less than that in oxidizing organic compounds.

2 OBJECTIVES OF WASTEWATER TREATMENT

From the foregoing description of biological behaviour, the consequences of discharging municipal wastewater to a water body can be appreciated. From a chemical point of view, municipal wastewater or sewage contains (1) organic compounds, such as carbohydrates, proteins and fats, (2) nitrogen, principally in the form of ammonia, and (3) phosphorus, which is principally in the form of phosphate from human waste and detergents. Municipal wastewater of course, has many other constituents of a particulate and dissolved nature, such as pathogens, plastics, sand, grit, live organisms, metals, elements, anions and cations. All these constituents have to be dealt with and considered at wastewater treatment plants, but not all of these are important for wastewater treatment plant modelling and design. Rather, attention is focused on the carbonaceous (C), nitrogenous (N) and phosphorus (P) constituents because these are the main ones that influence biological activity and eutrophication in the receiving water.

When municipal wastewater is discharged to a water body, the organic compounds stimulate the growth of the heterotrophic organisms causing a reduction in the dissolved oxygen. While oxygen is present, the ammonia, which is toxic to many higher life forms such as fish and insects, will be converted to nitrate by the nitrifying organisms causing a further demand for oxygen. Depending on the volume of wastewater discharged and the amount of oxygen available, the water body can become anoxic. If the water does become anoxic, nitrification of ammonia to nitrate by the autotrophs will cease. However, some of the heterotrophs now will use nitrate instead of oxygen as terminal electron acceptor and continue their metabolic

reactions. Depending upon the relative amount of organics and nitrate, the nitrate may become depleted. In this case the water will become anaerobic with associated sulphate reduction and fermentation. Irrespective of whether or not all of these steps happen, when the organic energy of the wastewater has been depleted, the water body in time recovers, clarifies and again becomes aerobic. But most of the nutrients, nitrogen (N) and phosphorus (P), remain, and these with sunlight penetration into the water body, stimulate aquatic plants like algae to grow in much the same way as farmers fertilize their lands with nitrogen and phosphates to stimulate crop growth. The algae reintroduce organic material into the water body and the pollution cycle starts over again. Only when the nutrients N and P are depleted and the organic energy sufficiently reduced, can the water body be said to be eutrophically stable again.

From these considerations, the objectives of wastewater treatment can be seen to be three-fold:

1. reduce organic bound energy to a level where it will no longer sustain heterotrophic growth and thereby avoid deoxygenation
2. oxidize ammonia to nitrate to reduce its toxicity and deoxygenation effects
3. reduce eutrophic substances, ammonia, nitrate and particularly phosphates, to levels where photosynthetic microorganisms are limited in their growth and therefore are limited in their capacity to fix solar energy as organic bound energy.

3 BASIC BIOLOGICAL BEHAVIOUR

3.1 Organism types in wastewater treatment

It was shown above that the nature of the electron donors and electron acceptors, which the various organism types utilize in their biological activities, form a useful means of categorizing organisms. In municipal wastewater treatment incorporating biological nutrient removal (BNR), two basic categories of

organisms are of specific interest: the heterotrophic organisms and the lithoautotrophic nitrifying organisms including the ammonia and nitrite oxidizers. The former group utilizes the organic compounds of the wastewater as electron donor and either oxygen or nitrate as terminal electron acceptor depending on whether or not the species is obligate aerobic or facultative and the conditions in the plant are aerobic or anoxic respectively. Irrespective of whether obligate aerobic or facultative the heterotrophic organisms obtain their catabolic (energy) and anabolic (material) requirements from the same organic compounds. In contrast the latter group, being lithoautotrophs, obtain their catabolic and anabolic requirements from *different inorganic* compounds, the catabolic (energy) from oxidizing free and saline ammonia in the wastewater to nitrite and nitrate and the anabolic (material) from dissolved carbon dioxide in the water. Being obligate aerobic organisms, only oxygen can be used as electron acceptor and therefore the nitrifying organisms require aerobic conditions in the plant.

In activated sludge plants irrespective of whether or not biological N and P removal is incorporated, the heterotrophic organisms dominate the biocenosis (mixed culture of organisms) and make up more than 98% of the active organism mass in the system. If the organism retention time (sludge age) is long enough, the nitrifiers may also be sustained but, because of their low specific yield coefficient compared to the heterotrophs (see Section 1.8 above) and relatively small amount of ammonia nitrified compared to organic material degraded, they make up a very small part of the active organism mass (<2%). Therefore, insofar as sludge production and oxygen or nitrate utilization is concerned, the heterotrophs have a dominating influence on the activated sludge system. Three different groups of heterotrophic organisms grow in the nutrient removal activated sludge system: (i) ordinary aerobic, (ii) facultative and (iii) polyphosphate accumulating heterotrophs, designated OHO, FHO and PAO respectively. While these three subgroups have some distinct behavioural characteristics, which result in the particular biological reactions required

to achieve the wastewater treatment objectives, their overall behaviour is similar and governed by their basic heterotrophic nature. An understanding of basic heterotrophic organism behaviour from a bioenergetic point of view, will therefore form a sound foundation on which many important principles in biological wastewater treatment are based, such as (1) growth yield coefficient and its association with oxygen or nitrate utilization, and (2) energy balances and its association with wastewater strength measurement.

3.2 Bioenergetics of heterotrophic organism behaviour

Bioenergetics is the study of material pathways occurring within organisms (and biosystems) as a result of an energy flow through a system. In *aerobic* systems the two principal pathways by means of which organic material is processed (metabolized) by organisms are catabolism and anabolism.

3.2.1 Catabolism and anabolism

1. *Catabolic pathway*: this is the pathway by which organisms obtain energy in a useful form for carrying out biological work, *inter alia* the work involved in protoplasm (organism mass) synthesis (termed anabolism see below). A fraction of the organic molecules taken up by an organism is enzymically oxidized to CO₂ and water, and a large amount of energy is released. A part of this energy is captured by the organism and is available for performing useful biological work; the part the organism is unable to 'capture' is lost as heat to the surroundings.
2. *Anabolic pathway*: this is the pathway by which organisms synthesize protoplasm (construct new cell mass). A fraction of the organic molecules taken up by the organism is enzymically modified to form part of the biological protoplasm. This synthesis process not only requires an input of organic molecules, but also inorganic molecules (e.g. ammonia (NH₄), phosphorus

(P), and micronutrients), energy, protons and electrons.

In biological aerobic treatment of organic compounds in wastewaters, the two material cycles set out above together constitute the *metabolism* of the organism and are of paramount importance. This is because one of the principal objectives of wastewater treatment is removal of energy (contained in the organic compounds) from the wastewater. Viewed over the biological treatment system as a whole, this objective is attained biologically in two ways:

1. through energy losses (as heat) to the surroundings resulting from the partial capture of energy in catabolism and the partial loss in the use of this energy for anabolism (both due to inefficiencies in energy transfers)³
2. through transformations of the organic compounds (dissolved or suspended) in the wastewater to organic molecules incorporated in biological protoplasm (anabolism).

The biomass so formed is settleable and is removed from the treated wastewater by physical separation of the biomass solids and wastewater liquid phases, leaving a clarified effluent virtually free of the original organic compounds. In most wastewater treatment plants, the biomass that is collected from the solid/liquid separation stage (usually a secondary settling tank) is returned to the biological reactor to inoculate the reactor with biomass for treating the continuously incoming wastewater (Fig. 9.1). Should the wastewater contain particulate organic material that is

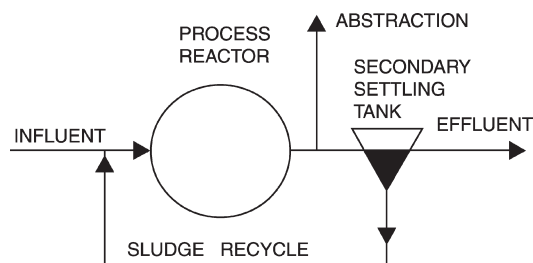


Fig. 9.1 Schematic layout of the activated sludge system in its simplest form showing biological reactor, secondary settling tank, sludge return and waste sludge abstraction from the biological reactor for hydraulic control of sludge age.

degraded slowly, this material will become enmeshed with the biomass, separated from the clarified effluent and returned to the reactor for further treatment. Slowly biodegradable particulate material therefore does not influence the effluent organic concentration. If there is slowly biodegradable organic material in a dissolved form it will escape with the effluent. However, in municipal wastewater treatment, most dissolved organics are readily biodegradable so low effluent organic material concentrations are usually easily achieved.

By means of a simplified bioenergetic model based on thermodynamic principles, an approximate estimate can be obtained of the proportion of the energy in the original wastewater organic compounds that is lost as heat or is transformed into biomass. This proportion is known as the active heterotrophic mass yield coefficient (Y_h , mass of organism formed per mass of organic material degraded). Apart from producing theoretical yield coefficients that can be usefully compared with experimentally measured values, the bioenergetic model also provides insight for many of the fundamental principles accepted as 'laws of nature' in activated sludge design models and procedures.

3.2.2 Organism electron and energy transport mechanisms

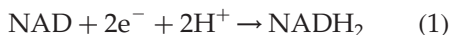
Energy becomes available to organisms principally through internally mediated reduction-oxidation (redox) reactions involving electron

³ In municipal wastewater treatment, the volumes of liquid undergoing biological treatment are too great and the heat losses from the biological reactions too small to make a significant difference to the temperature of the activated sludge mixed liquor. However, in autothermal sludge treatment where the organics are much more concentrated (by >100 times) and special precautions taken to minimize heat losses, the bio-heat released in the metabolism of the organisms is sufficient to raise the sludge liquor to greater than 60°C to affect sludge pasteurization. This is done in forced air composting, thermophilic aerobic digestion and dual digestion of sewage sludge.

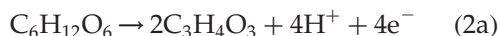
(and proton) transfer under controlled conditions. For heterotrophic organisms, the electrons are transferred from the organic compound, the electron donor (substrate or 'food' for the organisms), via a number of intermediate compounds, to some final or terminal electron (and proton) acceptor. In the course of the electrons moving from the electron donor to the electron acceptor via the various intermediate steps, energy is released progressively, some of which is captured by the organism. Fundamental to the means by which organisms harness the energy released in the redox reactions, are two types of compounds which couple into the redox reactions. The first is a compound called nicotinamide adenine dinucleotide (NAD). The oxidized (NAD) and reduced (NADH₂) forms act as *intermediate* electron acceptor and donor respectively. The second is a compound that in the form adenosine diphosphate (ADP) acts as an energy acceptor and in the form adenosine triphosphate (ATP) acts as an energy donor.

3.2.3 Electron transport molecule, NADH₂

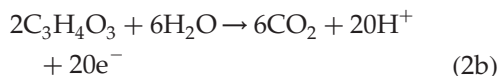
The organism electron acceptor molecule, NAD, is able to accept two electrons (e⁻) and two protons (H⁺) thereby becoming reduced, and in the reduced state is depicted as NADH₂, (equation 1):



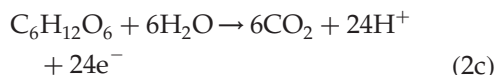
For the above half reaction to occur the organic compounds provide the electrons and protons in a concomitant second half reaction. Overall the organic compound is oxidized enzymatically to CO₂, protons (H⁺) and electrons (e⁻). Taking oxidation of glucose as an example, this reaction takes place in two stages. In the first stage the glucose is processed in a biochemical pathway called the Embden–Meyerhof pathway to pyruvate (equation 2a):



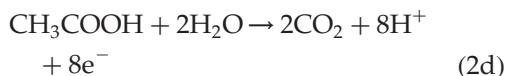
The pyruvate is then converted to acetyl CoA which enters the main oxidation pathway, called the Krebs or tricarboxylic acid (TCA) cycle, and is completely oxidized as follows (equation 2b):



The above two reactions combined give the net reaction for complete oxidation of the glucose (equation 2c):

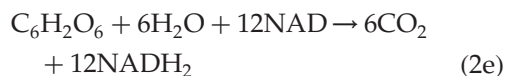


If the organic compound is acetate, this enters the Krebs or TCA cycle directly to give (equation 2d):

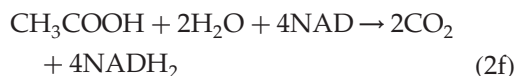


The electrons and protons from the organic material are captured by the organism in the NAD/NADH₂ reaction (equation 1). Balancing the electrons, the sum of the two half reactions in equations 1 and 2c, 2d is (equations 2e and 2f):

Glucose:



Acetate:

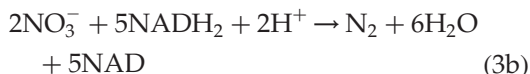


Once the organism has captured the electrons (and protons) in the form of NADH₂, the NADH₂ is available for generating new cell mass (anabolism) or energy (catabolism). In the latter case the electrons are donated to terminal electron acceptors either oxygen or nitrate, which are themselves reduced to water and nitrogen gas respectively (equations 3a and 3b).

Oxygen:



Nitrate:

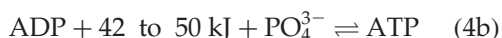
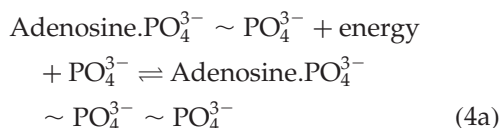


The energy released in these half reactions is partially captured by the organism via the formation of ATP, its energy transport molecule (see 3.2.4). Thus, the molecule NADH₂ provides the link between the substrate oxidation

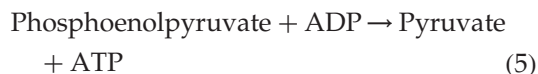
pathway, where it captures the electrons (and protons) released in oxidation of the organic substrate and the energy generation reactions, where it donates the electrons to the terminal electron acceptor. In this process some of the energy released is captured by the organism.

3.2.4 Energy transport molecule, ATP

The formation of ATP from ADP + and phosphate (PO_4^{3-}) requires an energy input of about 42 to 50 kJ/mol (10 to 12 kcal/mole) ATP formed under physiological conditions. The energy is conserved in the high-energy phosphoryl bond (equation 4a):

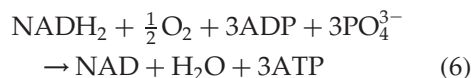


Two types of processes are used by organisms to form ATP from ADP and so capture the energy released in reactions, substrate and oxidative phosphorylation. In substrate phosphorylation, the energy capture couples directly into the transformation of organic compounds and ATP is formed without transfer of electrons to the terminal electron acceptor. This energy release/capture is associated with the direct transfer of a high energy phosphoryl group from the substrate to ADP to form ATP. This occurs during the partial breakdown of the organic compound to simpler organics like pyruvate that can enter the TCA (Krebs) cycle (equation 2a). Substrate phosphorylation does not involve the net release of electrons and therefore does not require an electron acceptor (Fig. 9.2). For example, in one step of the Embden-Meyerhof pathway, equation 5:



In oxidative phosphorylation, the energy capture reaction couples into the transfer of electrons to the terminal electron acceptor in the electron transport chain (ETC). In the ETC, NADH_2 is oxidized and the electrons are released and passed via a series of enzyme complexes to the terminal electron acceptor

(equation 3); in this process energy is released and some of it is captured by the organism in ATP. Noting that for aerobic conditions, passing 2 electrons from NADH_2 to the terminal electron acceptor oxygen, ideally gives rise to the formation of 3 ATPs (a feature of the ETC) as follows (equation 6):

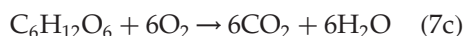


From equation 6 it can be concluded that:

1. Each molecule of NADH_2 releases a pair of electrons and protons which are captured by the electron acceptor oxygen.
2. Oxidation of 1 molecule of NADH_2 with oxygen ideally yields 3 ATP, but this ATP yield will vary depending on the electron acceptor.
3. Thus, each half molecule of oxygen reduced to water indicates that two electrons and protons have been transferred to it and 3 ATP formed.

3.2.5 Efficiency of energy capture in catabolism

Taking the oxidation of glucose with oxygen as an example (equations 7a, b and c):



By considering the free energies of formation (ΔG_f^0) of the reactants ($\text{C}_6\text{H}_{12}\text{O}_6$ and O_2) and products (CO_2 and H_2O), the free energy of the reaction (ΔG_R^0) is -2893 kJ/mol (-689 kcal/mole) glucose (where $-ve$ indicates a release of energy) (equation 8):

$$\begin{aligned} \Delta G_R^0 &= \Delta G_f^0(\text{products}) - \Delta G_f^0(\text{reactants}) \\ &= 6\Delta G_f^0(\text{CO}_2) + 6\Delta G_f^0(\text{H}_2\text{O}) \\ &\quad - \Delta G_f^0(\text{C}_6\text{H}_{12}\text{O}_6) - 6\Delta G_f^0(\text{O}_2) \\ &= 6(-395) + 6(-239) - (-611) - 6(0) \\ &= -2893 \text{ kJ/mol glucose or} \\ &= -689 \text{ kcal/mol glucose} \end{aligned} \quad (8)$$

In aerobic breakdown of 1 mole of glucose by heterotrophic organisms, 12 moles of NADH_2

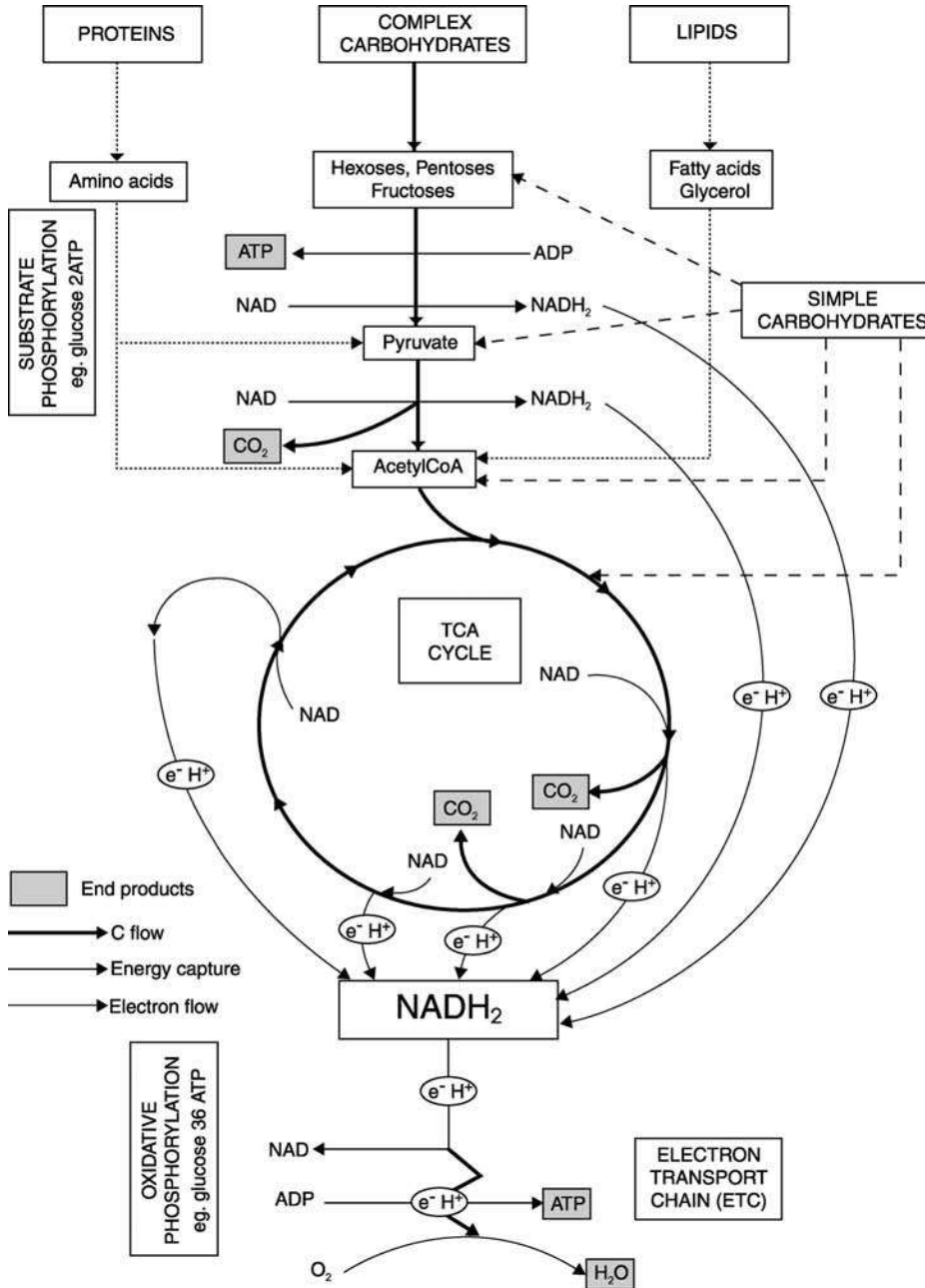


Fig. 9.2 Simplified schematic representation of heterotrophic organism catabolism of organics showing carbon flow from organics (substrate) to carbon dioxide, electron flow from donor (organics) to acceptor (oxygen) and energy capture as ATP in substrate phosphorylation (transformation – no electron release) and oxidative phosphorylation (electron transfer to terminal electron acceptor).

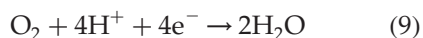
are produced (equation 2e). From equation 5, each mole of NADH, ideally produces 3 moles ATP by oxidative phosphorylation. This gives $12 \times 3 = 36$ moles of ATP produced by oxi-

dative phosphorylation in oxidation of 1 mole of glucose. In addition, 2 moles ATP are produced by substrate phosphorylation in the Embden-Meyerhof pathway (equation 2a).

This gives a total of 38 moles ATP produced for each mole glucose oxidized. Accepting that with each mol ATP formed, 42 kJ (10 kcal) is captured, then the energy capture is 1596 kJ/mol (380 kcal/mole) glucose. From the above, in complete oxidation of one mole of glucose, the free energy released is 2893 kJ/mol glucose. Thus, the energy captured in ATP on oxidation of glucose represents a fraction of $1596/2893 = 0.55$ of the free energy released. The organism is therefore able to capture about 55% of the free energy released and this energy is obtained via substrate phosphorylation (2 out of 38 ATP) in the Embden-Meyerhof pathway and via oxidative phosphorylation (36 out of 38 ATP) in passing the 24 electrons donated by the glucose to the electron acceptor oxygen in the transport chain (ETC) where 3 ATP are formed per electron pair transferred.

3.2.6 Organic material oxidation and free energy release

Taking the oxidation of glucose with oxygen as an example, from the section above, the free energy of the reaction (ΔG_R^0) is -2893 kJ/mol glucose where $-ve$ indicates a release of energy. Because $24e^-$ equivalents (e^- eq, or moles of e^-) are released in the reaction the free energy released per electron equivalent (e^- eq) is $-2893/24 = 120.5$ kJ/ e^- eq (-28.7 kcal/ e^- eq). Also, because (equation 9):



so that $1e^-$ eq = 8 gO (see equation 3a), the free energy released per gO is $-2893/(24 \times 3) = -15.1$ kJ/gO (-3.59 kcal/gO). For acetate, the free energy released is -853 kJ/mol (-203 kcal/mole). Noting that 1 mole acetate yields $8e^-$ eq and each e^- eq is equivalent to 8 gO, yields for acetate a free energy release of $853/(8 \times 3) = 13.3$ kJ/gO (-3.17 kcal/gO). With 4.2 J/cal, this is equal to -13.3 kJ/gO.

The free energy per gO is termed the free energy of combustion and represents the amount of available energy (energy that can be used to do work) released in the complete oxidation of the organic. It is important to note that the free energy of combustion for many different organic compounds when oxidized to

CO_2 and H_2O , is closely constant and ranges from 13.3 kJ/gO for the short-chain fatty acids and alcohols to 15.1 kJ/gO for the sugars and carbohydrates. Even the gaseous non-physiological fuels like methane and the liquid petroleum gases (LPG) give free energies of combustion in this range. For example for methane (CH_4) it is 13.86 kJ/gO and octane (C_8H_{18}) it is 13.65 kJ/gO. Clearly the free energy of combustion (useful energy released) is closely linked and almost proportional to the mass of oxygen consumed by the compound, or equivalently the number of electrons it can donate in its oxidation to CO_2 , and water. This is irrespective of whether it is done rapidly such as in combustion or slowly such as in organism respiration.

Experimental support for the proportionality between the free energy released and the electron donating capacity (EDC, i.e. kJ/ e^- eq), or equivalently through equation (9), the theoretical chemical oxygen demand (Th.COD, i.e. kJ/gO), was presented by Servizi and Bogan (1963, 1964) and Burkhead and McKinney (1969). Servizi and Bogan (1963) identified two groups of organics, a high energy group comprising carbohydrates, glycolysis products and TCA cycle intermediates and a low energy group comprising alcohols, aromatics, hydrocarbons (benzene ring compounds) and aliphatic acids (short or long chain saturated or unsaturated fatty acids). The difference in the free energy released per Th.COD between the high and low energy groups of organics was only 5% of the two groups mean value. McCarty (1972) found that the range of free energy released per Th.COD for naturally occurring organic compounds was bounded by methane, which yields the lowest, and formate, which yields the highest. The difference between formate and methane is only 10% of their mean value. Clearly there is substantive evidence that the free energy release in the oxidation (respiration to end products carbon dioxide and water) of organic compounds is proportional to the EDC or equivalently the Th.COD, and remains relatively constant for the spectrum of organic compounds.

The conclusion that the free energy released is proportional to the EDC for most organics is

fundamental not only to bioenergetics and tracing energy pathways in organisms, but also for measuring the energy strength of organic wastewaters. The application of this principle to both these aspects is discussed below.

3.2.7 Free energy release and energy capture by the organism

It has been concluded above that in complete oxidation of organics the free energy released is proportional to the EDC. In biological oxidation of the organics the proportion of this free energy captured by the organism is approximately constant provided the terminal electron acceptor (e.g. oxygen) is the same. This implies that the energy that becomes available to the organism in biological oxidation of organics is proportional to the EDC of the organic.

In considering biological oxidation, two separate phases can be identified. In the first phase, irrespective of which particular biochemical pathway or combination of biochemical pathways is used, the organic is oxidized to CO_2 , electrons and protons. The electrons and protons are captured by the organism in NADH_2 . In the second phase the electrons and protons from NADH_2 are donated to the terminal electron acceptor (via the ETC) and some of the energy released is captured by the organism in the formation of ATP (oxidative phosphorylation). *Since for a specific terminal electron acceptor the number of ATP molecules formed per pair of electrons passing to the terminal electron acceptor is constant, the energy captured by the organism in oxidative phosphorylation is proportional to the EDC of the organic.*

In Section 3.2.6 above, it was noted that there were two groups of organics; one comprising complex organics, such as carbohydrates and sugars, with about 14% higher heats of combustion than the other group comprising simple organics, such as acetate and pyruvate. There are also some minor variations in the total number of ATPs formed per e^- donated by different organics. These minor differences in the energy captured by the organism are due to the relative contribution to ATP production by substrate and oxidative phosphorylation. In oxidative phosphorylation with the same terminal electron acceptor, the same amount of

ATP is generated by the organism per electron donated by the organic material. However, the amount of ATP generated by substrate phosphorylation will vary depending on the nature of the organic compound and the biochemical pathways used in its transformation prior to entering the TCA (Krebs) cycle (see Fig. 9.2). For example, glucose will give 2 ATP via substrate phosphorylation and acetate none. However, substrate phosphorylation plays a relatively minor role in ATP generation – for glucose only 2 ATP out of a total of 38 ATP are generated by substrate phosphorylation and the balance by oxidative phosphorylation. The dominating effect of oxidative over substrate phosphorylation implies that the EDC or Th.COD will provide a good estimate of the energy that will become available to the organism. This dominating effect of oxidative phosphorylation is also the reason why the difference in the heats of combustion between the different organic groups is so small. It is also the reason why energy changes, both free energy released and that captured by the organism in the form of ATP, can be monitored via the changes in EDC or Th.COD.

3.3 Energy utilization in heterotrophic organisms – organism yield and oxygen utilization

In utilization of an organic compound by the organism, both the energy and the electrons must be conserved, i.e. we must be able to trace the fate of both the energy and electrons. The energy is conserved in that, of the free energy available in the organic compound, some is retained as organic compounds of a different form, i.e. the organism mass and the rest is 'lost' as heat. The electrons are conserved in that, of those available in the organic compound, some are retained in the organic compounds of the organism mass formed and the rest are passed on to the terminal electron acceptor (Fig. 9.3). The proportion of the electrons (or energy) retained by the organisms defines the specific organism mass yield coefficient Y_h , i.e. mass of organism mass formed per mass of organic compound

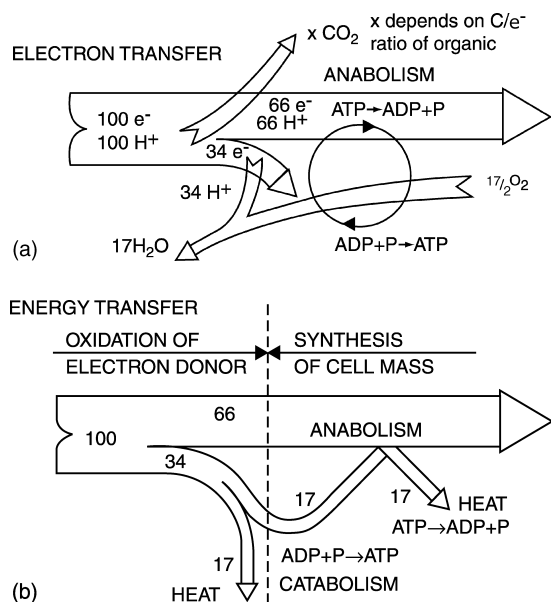


Fig. 9.3 Electron and energy transfer in anabolism and catabolism, showing electron and energy capture in organism mass synthesized (anabolism) and electron 'loss' by transfer to the electron acceptor (a, top) and energy 'loss' as heat to drive the anabolic processes (catabolism) (b, bottom).

oxidized, all in terms of electron equivalents (equation 10):

$$Y_h = \frac{e^- \text{ (or energy) conserved in organism mass}}{e^- \text{ (or energy) in substrate utilized}} \quad (10)$$

Loewenthal (1986) developed a procedure from bioenergetic and thermodynamic principles with which it can be shown that the specific yield coefficient (Y_h) for heterotrophic respiration reactions involving oxygen as the terminal electron acceptor (i.e. in which 3 ATPs are formed per pair of electrons transferred) is 0.42 g cells/gTh.COD. This procedure is briefly outlined below in which the anabolic and catabolic pathways are considered separately and then combined to determine the yield.

3.3.1 Anabolic pathways

The organic components of organism mass are highly complex and very numerous,

but principally are proteins, fats (lipids) and carbohydrates. The formation of these compounds (anabolism) follows a wide variety of biochemical pathways too complex to trace. Consequently, the procedures to estimate heterotrophic yield are developed from global considerations only.

The first requirement for a global approach is a stoichiometric formula representative of organism cell mass. A number of these have been proposed (Table 9.1) each comprising different quantities of C, H, O and N. Of these formulae, $C_5H_7O_2N$ seems the most widely accepted and consequently will be adopted for this discussion. An important requirement of the stoichiometric formula for organisms is its free energy of formation (ΔG_f^0). This value is not reported in the literature, but has been deduced by Loewenthal (1986) to be +40 kcal/mole protoplasm as defined by $C_5H_7O_2N$. With this value it is possible to calculate from thermodynamic principles the fraction of energy or EDC transformed to cell mass and the fraction lost as heat. For such calculations the ΔG_f^0 protoplasm can be accepted to remain constant for different electron donors and acceptors provided the redox reactions are those of respiration.

Recognizing that the anabolic synthesis reactions follow a myriad of pathways too complex to model, the reactions are simplified by assuming they take place in two steps. The first of these is the formation of protoplasm ($C_5H_7O_2N$) from CO_2 , NH_3 and electrons (and protons). The second step is the oxidation of substrate to supply the CO_2 and electrons

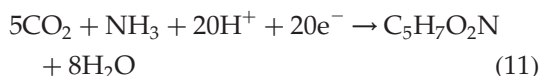
TABLE 9.1 Theoretical COD values for various empirical formulations for microbial sludge (after McCarty, 1965)

Bacterial formula	Mol. mass	Th. COD (g)			Ref
		Per mol	Per g vss	Per g C	
$C_5H_7O_2N$	113	160	1.42	2.67	1
$C_5H_9O_3N$	131	160	1.22	2.67	2
$C_7H_{10}O_5N$	156	232	1.48	2.76	3
$C_5H_8O_2N$	114	168	1.47	2.80	4

1. Hoover and Porges (1952) 2. Speece and McCarty (1964) 3. Sawyer (1956) 4. Symons and McKinney (1958).

(and protons) for the first step. In terms of these two steps, the function of the anabolism reactions is to transform electrons and protons in the substrate to electrons and protons in the organism mass.

Formation of protoplasm from CO₂, NH₃ and electrons. To form 1 mole of protoplasm (which has a mass of 113 g – Table 9.1) from CO₂ and NH₃ requires 20 electrons (and proton) (equation 11):



As noted earlier, organisms transfer electrons (and protons) via the NADH₂/NAD molecules. Coupling the NADH₂ → NAD half reaction into equation 11 yields equation 12:



Equation 12 is *not* intended to show that the organisms are chemical autotrophs obtaining their cell material from carbon dioxide and ammonia. The organisms considered are heterotrophs and so the cell material is obtained from the organic compound. It is only for the purposes of establishing a datum for the free energy calculations that the organic compound is assumed to be broken down to CO₂ and H₂O and the protoplasm built up from these basic products.

Oxidation of substrate to supply electrons and protons. To supply the electrons and protons necessary for the first step above, the organism must oxidize substrate (note that the balance is made on supply of electrons, not carbon since the ratio of moles of CO₂ produced per mole of substrate oxidized varies between substrates). From equation 11 for protoplasm formation, the organism needs to obtain 20 e⁻eq (10 moles NADH₂). If the organic substrate is glucose or acetic acid, which yield 24 or 8 electrons per mole of substrate respectively (equations 2c and d), then 20/24 = 0.83 moles of glucose or 20/8 = 2.50 moles of acetate are required for anabolism. The anabolism reaction (equation 12) is an uphill one and energy must be

supplied to drive it. The anabolic energy requirement is calculated from the free energy of reaction for the formation of protoplasm (equation 12). Subtracting the standard free energies of formation (ΔG_f⁰) of the reactants (left hand side of equation 12) from those of the products (right hand side of equation 12), yields the standard free energy of reaction (ΔG_r⁰) for the formation of protoplasm. In other words, + 465 kJ/mol (+110.6 kcal/mole) protoplasm with the +ve sign indicating that it is an energy requiring reaction. The energy transfers by the organism are not 100% efficient and it can be assumed that the efficiency of the energy transfer in the anabolic reaction is the same as calculated earlier for ATP formation at only about 55%. Hence 465/0.55 = 844 kJ/mol (201 kcal/mole) protoplasm are required. This energy is supplied by ATP generated in the catabolic pathways.

3.3.2 Catabolic pathways

The catabolic pathways supply the energy (ATP) requirements for the anabolic reactions. As mentioned earlier each ATP has an available free energy of 42 kJ/mol (10 kcal/mole) ATP. Thus to supply the energy requirements for anabolism (944 kJ/mol protoplasm), 844/42 = 20.1 mol ATP must be generated in catabolism. Remembering that ideally 3 ATP are formed per pair of electrons transferred to the terminal electron acceptor oxygen, the catabolic electron equivalent requirement is 20.1/3 = 13.4 e⁻eq. To supply these electron equivalents, organic molecules are oxidized by the organism. If the organic is acetic acid or glucose then 13.4/8 = 1.67 or 13.4/24 = 0.56 moles of organic substrate/mol protoplasm respectively are required.

3.3.3 Yield

The combined anabolic and catabolic electron equivalent (e⁻eq) requirements to synthesize 1 mole of protoplasm are therefore 20 + 13.4 = 33.4 e⁻eq. At 8 and 24 e⁻eq/mole for acetate and glucose respectively, this gives 2.5 + 1.67 = 4.17 moles acetic acid and 0.83 + 0.56 = 1.39 moles glucose respectively. Now 1 mole of protoplasm is 113 g mass and because its constituents C₅H₇O₂N are all organic, its

volatile solids mass, VSS, is also 113 g (see Table 9.1). Therefore the theoretical specific organism yield is $113/4.17 = 27.1$ gVSS/mole acetic acid or $113/1.67 = 67.7$ gVSS/mole glucose. Because the organisms respond to the free energy or equivalently the e^- eq, it is better to specify the yield coefficient in terms of e^- eq, i.e. $113/33.4 = 3.38$ gVSS/ e^- eq. To express e^- eq in terms of oxygen, $1 e^-$ eq is equal to 8 gO (equation 9), so that the theoretical specific organism yield in terms of the Th.COD is $3.38/8 = 0.42$ gVSS/gTh.COD. Using the COD test (Standard Methods, 1985) as an estimate of the Th.COD, Marais and Ekama (1976) measured the yield coefficient for activated sludge grown on municipal wastewater as 0.43 mgVSS/mg COD, very close to the theoretically estimated value. In activated sludge models, a Y_h value for heterotrophs of 0.45 mgVSS/mgCOD is used.

From Table 9.1, it can be seen that the Th.COD of the $C_5H_7O_2N$ protoplasm formed is 1.42 gO/gVSS. This was obtained by dividing the Th.COD of the protoplasm in its oxidation to CO_2 and water (i.e. $20 e^-$ eq or 5 moles of O_2 which is equal to $5.32 = 160$ gO) by the mass of protoplasm i.e. $160/113 = 1.42$ mgTh.COD/mgVSS. With a yield coefficient of 0.42 mgVSS/mgTh.COD, the Th.COD of the protoplasm formed is $0.42 \times 1.42 = 0.60$ mgTh.COD/mgTh.COD. This means that of the original organic material Th.COD, 60% is transformed to new organism mass. The balance of 40% is electrons passed on to the terminal electron acceptor oxygen. The same proportion (60/40) is obtained by considering the electron equivalents directly, i.e. of the $33.4 e^-$ eq required to form 1 mole (113 g) of protoplasm, $20 e^-$ eq (60%) were conserved in protoplasm and $13.4 e^-$ eq (40%) were passed on to oxygen. The e^- eq passed on to oxygen manifests as oxygen utilization by the organism.

Experimental support for Loewenthal's (1986) approach to estimate theoretically the yield coefficient, Y_h from bioenergetics and thermodynamic data is provided by Bauchop and Elsdén (1960). They found that organism yield is linearly related to the number of ATP moles produced in the complete oxidation of

organics, i.e. $Y_h \propto K_1 N_{ATP}$ where N_{ATP} is the number of ATP moles generated per mole of organic and Y_h the gVSS generated per mole of organic. The constant K_1 was found to be 2.12 gVSS/mole ATP. Furthermore, for the known aerobic substrate pathways, it can be shown that the number of ATP moles produced per kJ free energy released is constant at about 0.013 moles ATP/kJ, i.e. $N_{ATP} = K_2 \Delta G_f^0 = 0.013 \Delta G_f^0$. For example, in the breakdown of glucose, 38 moles of ATP are generated and -2894 kcal of free energy released making the ratio $K_2 = 38/2894 = 0.013$ mole ATP/kJ. Therefore, through K_1 and K_2 the yield coefficient for heterotrophs, Y_h , is proportional to the free energy released by the organic compound, i.e. $Y_h = K_1 K_2 \Delta G_f^0$. Defining Y_h and ΔG_f^0 in terms of the Th.COD, Y_h for glucose is $0.013 \times 2.12 \times (2894/192) = 0.41$ gVSS/gTh.COD, which is very close to Loewenthal's theoretical value of 0.42 gVSS/gTh.COD. Experimental data from the aerobic metabolism of a wide variety of organic substrates presented by Eckenfelder (1965) give an average coefficient of 0.39 gVSS/gCOD, which is also close to the theoretically estimated value (Fig. 9.4).

3.3.4 Nitrate as terminal electron acceptor

When nitrate serves as electron acceptor, approximately the same quantity of free energy

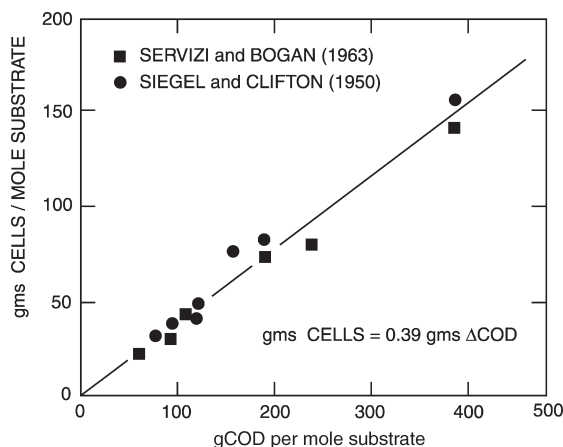


Fig. 9.4 The direct relationship between COD and Yield (Y_h) for a wide variety of substrates (Eckenfelder, 1965).

TABLE 9.2 Comparison of measured free energy transferred by different organics with oxygen and nitrate as terminal electron acceptor (McCarty, 1972)

<i>Energy source electron donor</i>	<i>Energy transfer (kcal/e⁻)</i>		
	<i>Electron acceptor</i>		<i>Ratio</i>
	<i>Oxygen</i>	<i>Nitrate</i>	
1. Methanol	27.6	26.1	0.94
2. Ethanol	26.3	24.7	0.94
3. Domestic WWV	26.3	24.7	0.94

is available (94%) to the organisms as when oxygen serves as electron acceptor (Table 9.2). However, the number of ATP moles formed per electron transferred is reduced. In the calculations above with oxygen as terminal electron acceptor, 3 ATP are formed per pair of electrons transferred. If nitrate is used instead of oxygen, only 2 ATP are formed per pair of electrons transferred to the nitrate (Payne, 1981). Accordingly with nitrate, the catabolic electron equivalent requirement is $20.1 \times 2/2 = 20.1 \text{ e}^- \text{eq}$. To supply these electron equivalents, organic molecules are oxidized by the organism. If the organic is acetic acid or glucose then $20.1/8 = 2.51$ or $20.1/24 = 0.84$ moles of acetic acid or glucose respectively are required.

The combined anabolic and catabolic $\text{e}^- \text{eq}$ requirements to synthesize 1 mole of protoplasm with nitrate as electron acceptor is therefore $20 + 20.1 = 40.1 \text{ e}^- \text{eq}$, which for 8 and $24 \text{ e}^- \text{eq}$ for acetate and glucose respectively, gives $2.5 + 2.51 = 5.01$ moles acetic acid and $0.83 + 0.84 = 1.67$ moles glucose respectively. Converting the units as described above, this gives the theoretical specific organism yield in terms of the Th.COD as $0.35 \text{ gVSS/gTh.COD}$. Thus the yield with nitrate as terminal electron acceptor ($0.35 \text{ gVSS/gTh.COD}$) is lower than that with oxygen ($0.42 \text{ gVSS/gTh.COD}$). This theoretical reduction in organism yield has not been incorporated in steady state activated sludge models and design procedures nor is it recognized in the earlier general kinetic activated sludge simulation models. These include UCTOLD (Dold *et al.*, 1991), UCTPHO (Wentzel *et al.*, 1992) and ASM Nos 1 and 2 of IAWO

(Henze *et al.*, 1987, 1994). In these models, it is accepted that the bioenergetics of aerobic and anoxic respiration are the same in every respect, the only difference is that the electrons are passed to oxygen for aerobic conditions and nitrate for anoxic conditions. More recent simulation models like ASM2d and 3 include a reduction in yield coefficient under anoxic conditions but the differences are relatively small.

3.4 Conclusions from bioenergetics

Two important principles flow from the above discussion of bioenergetics.

3.4.1 The biological oxygen utilization is directly linked to substrate utilization for synthesis

The oxygen equivalent (Th.COD) of the organic compound is not the same as the oxygen demand for synthesis of protoplasm. The former is a measure of the free energy released in the complete oxidation of all the organic compound to CO_2 and H_2O . This is equivalent to the heat of combustion where all the free energy released is lost as heat. The oxygen utilization for synthesis forms part of the heterotrophic biological oxygen utilization (the other part is endogenous respiration required for cell maintenance) and represents only a part (40%) of the original Th.COD of the organic compounds. The balance of the Th.COD is transformed into the new cell mass synthesized. Thus, in a hypothetical 1 m^3 bioreactor wherein only synthesis of heterotrophic biomass is taking place on glucose, in the degradation of 1 mole of glucose (180 g), which has a Th.COD of 192 gO, $0.42 \times 192 = 80.6 \text{ gVSS}$ organism mass ($\text{C}_5\text{H}_7\text{O}_2\text{N}$) is formed, comprising $80.6 \times 1.42 \text{ gTh.COD/gVSS} = 114.5 \text{ gTh.COD}$. The biological oxygen utilization (OU) is the difference between the Th.COD of the glucose and the organism VSS mass formed, i.e. $\text{OU} = 192 - 114.5 = 77.5 \text{ gO}$, which is 40% of the original Th.COD of the glucose.

Thus, in general heterotrophic organism synthesis, if the Th.COD concentration of

the organic compound is denoted S and it is completely degraded, i.e. $\Delta S = S - 0$, then the concentration of organic material degraded, the concentration of organisms formed and the biological OU are directly linked and given by equations 13a, 13b and 13c.

$$X_a = Y_h \cdot \Delta S \quad (13a)$$

where:

X_a = concentration of organisms formed (gVSS/m³)

Y_h = yield coefficient (gVSS/gTh.COD)

ΔS = substrate utilized (gTh.COD/m³)

$$Z_a = f_{cv} \times Y_h \Delta S \quad (13b)$$

where:

Z_a = Th.COD of X_a formed (gCOD/m³)

f_{cv} = COD/VSS ratio of organisms = 1.42
organism VSS

$$\begin{aligned} \text{OU} &= \Delta S - Z_a = \Delta S - f_{cv} Y_h \Delta S \\ &= (1 - f_{cv} Y_h) \Delta S \end{aligned} \quad (13c)$$

where:

OU = Biological oxygen utilized for synthesis of the organism concentration (gO/m³)

3.4.2 Energy balances can be made over biological treatment systems in terms of electron equivalents

By measuring the OU, which represents the electrons passing to oxygen in organism mass synthesis, an energy balance over the biological system is possible in terms of e^- eq or O. If the organic compounds are measured as Th.COD, then for electrons to be conserved, the mass of Th.COD entering the reactor must be equal to the mass of Th.COD leaving the reactor. The former is the mass of Th.COD entering the reactor via the influent flow and the latter is given by the sum of the Th.COD of the undegraded organics in the effluent flow, the OU, and the Th.COD of the organisms removed from the system, all over a fixed period of time. The ability to build e^- eq or Th.COD mass balances is a powerful feature of the Th.COD approach and forms the basis not only of the steady-state activated sludge

model (Marais and Ekama, 1976; WRC, 1984), but also of the activated simulation models such as UCTOLD (Dold *et al.*, 1991), UCTPHO (Wentzel *et al.*, 1992) and IWA ASMs Nos 1 and 2 (Henze *et al.*, 1987, 1994).

To build from the above two principles in activated sludge kinetic modelling requires a means whereby the EDC or theoretical COD of the organics in wastewater can be measured in a practical way. The issues associated with this and energy measurement in general are discussed in the next section.

4 ENERGY MEASUREMENT

If the organic compounds are defined in terms of both type and concentration, then there is no difficulty in determining the free energy release, EDC or Th.COD. These can be calculated directly from thermodynamic and chemical data as illustrated for glucose and acetate (see Section 3.4). The EDC or Th.COD can even be calculated for a mixture of organics, if each is defined in type and concentration. However, when the mixture of organics is not defined in type and concentration, as is the usual case in wastewater treatment, then the free energy release, EDC or Th.COD cannot be calculated from thermodynamic and chemical data. It is here that the significance of the above discussion comes to the fore because (1) the free energy release in the oxidation of most organics is related to the EDC or Th.COD of the organics, and (2) organism metabolic behaviour is deeply rooted in electron transfer redox reactions. If a method of experimentally determining the EDC or Th.COD of a mixture of organics could be established, then a method of tracing the free energy changes in biological treatment systems would be available.

Currently there are four tests in wide use to estimate the energy content of the organics in wastewaters. Three of these are oxygen based tests and one is a carbon based test, namely: biochemical oxygen demand (BOD), chemical oxygen demand (COD), total oxygen demand (TOD) and total organic carbon (TOC). Each is briefly reviewed to evaluate their advantages and disadvantages for measuring the energy content of organics in wastewaters.

4.1 Biochemical oxygen demand (BOD) test

This test provides a measure of the oxygen consumed in the biological oxidation of organic (carbonaceous) material in a sample at 20°C over a prescribed period of time, usually 5 days. According to Gaudy (1972), the origins of this test can be traced back about a hundred years to Frankland who appears to have been among the first to recognize that the observed oxygen depletion in a stored sample is due to the activity of microorganisms. The Royal Commission on Sewage Disposal appeared to have developed the concept of using the oxygen depletion as a measure of the strength of pollution. Over the past century extensive research towards elucidating the meaning of the test, quantifying it and integrating it in theoretical descriptions of wastewater treatment processes, has been undertaken. To date the biochemical oxygen demand (BOD) test is still the most popular parameter for assessing the organic material strength of influents and effluents and for describing the behaviour of treatment systems. Its continued popularity seems to stem mainly from the body of experience that has built up in its use. While there is some justification for retaining it for setting effluent quality standards for regulatory purposes, in so far as present day modelling, design and operation of biological treatment processes is concerned, few cogent arguments for its continued use over that of the COD test can be advanced.

The BOD test is an empirical one performed under strictly specified conditions and procedures and any deviation may give rise to very uncertain results. In the 5-day BOD test (BOD₅), which is the one commonly employed, the sample of wastewater is suitably diluted with well-oxygenated water and an inoculum of microorganisms, adapted to the wastewater if necessary, is introduced. The initial oxygen concentration is measured and the sample stored in darkness at 20°C for 5 days. Over the 5-day period, the oxygen concentration is measured (which is done automatically with modern BOD apparatus) at regular intervals and at the end of the 5-day period. The differ-

ence in oxygen concentration between the beginning and the end of the 5-day test period, taking due account of dilution, gives the BOD₅. For the ultimate BOD test (BOD_u) the sample is stored for about 20 days at 20°C to provide sufficient time for the complete utilization of the biodegradable organic energy (Standard Methods, 1985). Of the two, the BOD_u test probably gives the most stable relative measure of the biodegradable organic energy, but the time factor of 20 days makes the test impractical for routine use. If the BOD_u/BOD₅ ratio remains constant for different wastewaters and different stages of primary (sedimentation) and secondary (biological) treatment, then the BOD₅ test would serve as a satisfactory relative measure. However, this is different between wastewaters and also at different stages of treatment for the same wastewater.

The BOD tests are also very sensitive to test procedure and inoculum preparation. Low BOD values are obtained if the inoculum is not suitably adapted to the wastewater. Some wastewater constituents like heavy metals and toxic organics inhibit organism activity. Also nitrifying organisms may develop in the test sample unless specifically suppressed by chemical additives (usually allyl thiourea, ATU). In particular, nitrification may occur in effluent samples from nitrifying biological treatment plants, or in influent samples where some treatment plant operational practice introduces nitrifying secondary sludge to the influent stream, or in tests where the inoculum is obtained from a nitrifying system. Nitrifying organisms in the wastewater or effluent sample convert ammonium (NH₄⁺) to nitrite (NO₂⁻) and nitrate (NO₃⁻) and utilize oxygen in the process (about 4.6 mgO₂/mgNH₄ - N nitrified to nitrate, NO₃ - N) to give an inflated value for the organic 'energy' content. Test procedure, inoculum adaptation, inhibition and nitrification always need to be carefully considered in evaluating BOD test results.

Despite the deficiencies listed above, if the BOD₅ value constituted a consistent estimate it could still have provided a relative measure of practical usefulness, but even this cannot be guaranteed. Gaudy (1972), in a thought provoking review on the BOD, gave a summary of

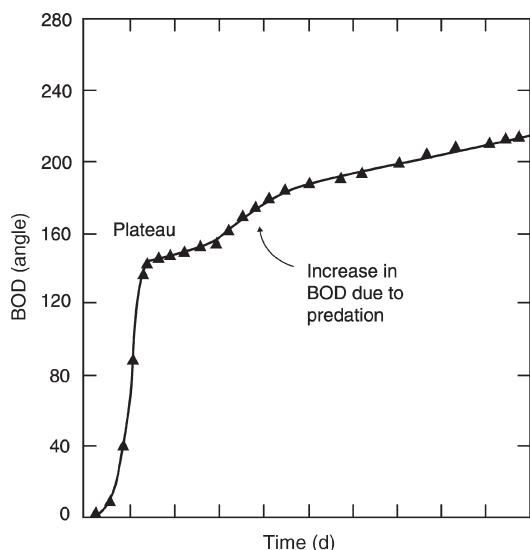


Fig. 9.5 Typical experimental BOD time curve on wastewater with heterogeneous seed clearly showing 'plateau' behaviour after about 1.5 days, followed by an increase in oxygen utilized, considered to be due to predation (Copcutt, 1993).

the work on the BOD test (in which he took a leading part). He concluded that the standard approach to formulating the BOD time curve as a first order reaction is quite invalid. The actual curve arises from two main effects: (1) bacterial synthesis from the biodegradable input – a reaction usually complete in 1–2 days, and (2) predator growth using the synthesized bacteria as substrate. The second phase, being dependent on the first, usually lags the first so that a distinct 'pause' or 'plateau' in the time curve is exhibited (Fig. 9.5). The occurrence and duration of this pause is dependent on the bacteria–predator relationship initially present in the sample, the 'strength' of the organic material, presence of inhibitory substances and so on (Hoover *et al.*, 1953). These compound effects certainly are not reflected by a first order formulation. Furthermore, should nitrification occur during the 5 days, a second plateau in the time curve might arise. This may not be apparent unless the test results are compared with a test in which procedures to suppress nitrification are imposed. Evidently the BOD₅

value is subject to a number of influences, which can be present in varying degrees, resulting in unknown effects on the observed value, which indicates the importance of measuring the complete BOD-time curve in the test to enable a meaningful interpretation. For routine use this is onerous and time consuming.

The main deficiency of the BOD tests is that they do not measure the electron donating capacity (EDC) of the organics in the wastewater. From the discussion of biological behaviour above, the oxygen utilized is only the catabolic energy requirement of the organisms; the greater part (~60%) of the energy or electrons are transformed to new organic compounds in the form of organism mass, the anabolic requirement of the organisms. Even though these compounds become organics for other organisms to utilize in repeated predator–prey cycles with a catabolic oxygen requirement in each cycle, some of the organism mass formed is not degraded within the time period of the test and also some of the organism mass is unbiodegradable and accumulates as unbiodegradable organism (endogenous) (Fig. 9.6). Therefore in the BOD₅ test organics not only is the utilization of the wastewater organics incomplete to some unknown degree, but also some of the electrons (energy) of the wastewater organics become bound up with unbiodegradable organism residue. Even in the BOD_u where the utilization of the wastewater organics may be complete, an unknown fraction of the biodegradable energy originally present in the wastewater does not manifest as oxygen utilization and remains as unbiodegradable residue. Consequently, it is not possible to perform an electron (or energy) balance on the biodegradable organics of the wastewater with the BOD tests. This is a severe deficiency and places the BOD outside the major developments that have taken place in biological treatment process modelling, design and simulation, all of which are based on electron (COD) mass balance principles (Marais and Ekama, 1976; Dold *et al.*, 1980; Van Haandel *et al.*, 1981; WRC, 1984; Henze *et al.*, 1987, 1994; Dold *et al.*, 1991; Wentzel *et al.*, 1992).

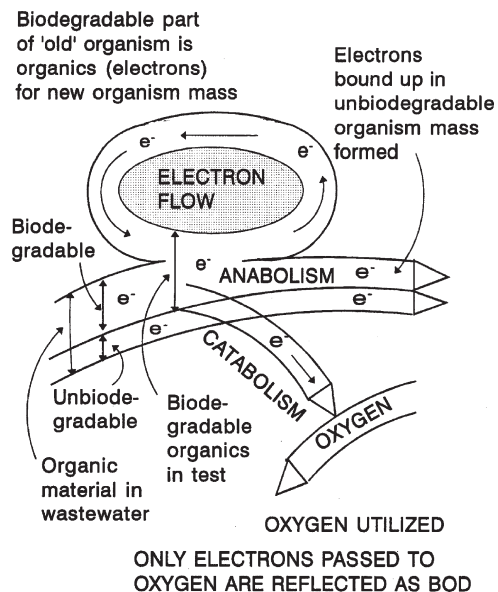


Fig. 9.6 Electron flow in the BOD test showing (1) oxygen utilized (BOD) as a consequence of organism catabolic action only, (2) organic material recycling through predation, and (3) electrons bound in unbiodegradable material from the influent or formed biologically in the test do not manifest as oxygen utilized (BOD).

Many organizations, in particular the regulatory agencies, regard the fact that the (nitrification inhibited) BOD tests measure only the strength of the biodegradable organics in the sample an advantage. For the wastewater treatment plant effluent that may be so, but for the influent this is a considerable shortcoming. Most wastewaters include biodegradable and unbiodegradable organics of a soluble and particulate nature. The soluble unbiodegradable organics (in terms of the time spent in the wastewater treatment system) escape with the effluent, but the particulate fraction becomes enmeshed in the biomass in the biological reactor and manifests as a part of the volatile solids concentration of the biomass. The BOD tests therefore do not incorporate a procedure related to the test whereby an assessment of this accumulation process can be obtained. In BOD₅-based design procedures, this is taken account of by measuring the suspended solids concentrations with 0.45 μm membrane filters and assuming a certain

fraction of these solids are not biodegradable. In the COD test (discussed below), because the total electron donating capacity of the organics is measured, whether it is biodegradable or unbiodegradable, it is possible to estimate the soluble and particulate unbiodegradable organic fractions from the COD test results and the kinetic behaviour (sludge production and oxygen utilization) of the activated sludge system.

4.2 Chemical oxygen demand (COD) test

This test gives the electron donating capacity (in terms of oxygen, see equation 7) of practically all the organic compounds in the sample, biodegradable or unbiodegradable and soluble or particulate. A strong oxidizing agent, a hot mixture of dichromate and concentrated sulphuric acid making chromic acid, oxidizes the organic compounds to CO_2 , the same end product as biological activity. In the test, a known mass of potassium dichromate (volume times concentration) and measured volumes of wastewater and sulphuric acid are added to the test and refluxed (boiled) for 2 hours. After the sample has cooled, it is titrated with ferrous ammonium sulphate (FAS) and ferrous indicator (Ferroin). The dichromate is the electron acceptor and the organic compounds and FAS the electron donors. During refluxing, the organic compounds are oxidized to CO_2 , and their electrons pass to the dichromate, which is reduced. After all the organic compounds are oxidized, some dichromate electron acceptor remains and it is this remaining amount that is measured in the titration with the FAS. The ferrous (Fe^{2+}) in FAS donates electrons to the remaining dichromate and is oxidized to ferric (Fe^{3+}) while the dichromate is reduced. When the dichromate has all been reduced, it can accept no more electrons and the ferrous of the FAS remains as Fe^{2+} . At this point the Ferroin indicator detects the presence of the Fe^{2+} ions and changes colour (yellow to red), which defines the end point of the titration. The difference between the initial electron accepting capacity of the dichromate (measured by

titrating a blank in which distilled water instead of sample is refluxed) and number of electrons donated by the FAS titration, gives the number of electrons donated in the oxidation of the organic compounds. This electron donating capacity expressed as O_2 is the COD. The COD test is thus rooted in redox chemistry and therefore is directly related to the Th.COD discussed earlier. But how close is the COD test result to the Th.COD?

An important and time saving feature of the COD test is that ammonia is not oxidized. For most municipal wastewaters the COD result is therefore virtually exclusively associated with the organic material. Nitrite is oxidized to nitrate in the test, a reaction which exerts a COD. However, this COD can be suppressed in the test by adding sulphamic acid to the dichromate, but usually nitrite concentrations are so low in comparison with the organic contents and the oxygen requirement so low ($1.2 \text{ mgO per mgNO}_2^- - \text{N}$ oxidized to $\text{mgNO}_3^- - \text{N}$) that this influence is negligible. With industrial wastewaters, which have significant concentrations of reduced inorganic compounds such as sulphides, the electrons donated in the oxidation of these inorganics can contribute significantly to the COD result, and therefore caution should be exercised with such wastewaters.

In practical application, some organic material is not oxidized by the COD test under any circumstances but microorganisms do utilize it. This organic material includes aromatic hydrocarbons (e.g. benzene, phenols) and pyridines (water soluble benzene rings with an N in place of one of the carbon atoms). They are mainly encountered in petrochemical and plastic industry wastewaters and are not likely to be encountered in municipal wastewater in significant concentration. A poor estimate for COD can also be obtained with lower fatty acids (the C_2 to C_5 acids) such as acetate. These compounds are in the unionized form at the low pH at which the test is performed – the unionized molecules can be extremely difficult to oxidize. The addition of the catalyst silver sulphate overcomes this problem to a large degree to give virtually 100% oxidation of unionized short-chain fatty

acids. However, silver sulphate reacts with the halides (chloride, bromide and iodide) to produce precipitates which do not act as catalysts. This difficulty can be largely overcome by adding mercuric sulphate to complex the halides, to an upper limit of 2000 mg Cl/l .

In performing a COD test it should be remembered that the test involves a time dependent reaction and the specified refluxing period of 2 hours must be adhered to to ensure complete or near complete oxidation. Furthermore, the degree of oxidation is subject to mass action effects – equivalent masses of the same organic compound refluxed at different final test volumes yield different results for the COD. The temperature at which the refluxing step takes place also affects the oxidation rate. The refluxing temperature in turn is dependent on the concentration of sulphuric acid in the test. These factors have all been considered in the development of the COD test procedure. Hence, the COD test will give reliable results only if the test is done in strict accordance with the set procedures, e.g. those in Standard Methods (1985) or later editions.

If the specified procedures for the COD test are strictly followed, experience has shown that it accurately reflects the electron donating capacity (EDC) of a mixture of organic compounds. Over the past 15 years application of COD test results to organic material balances on fully aerobic and anoxic/aerobic activated sludge plants indicate that, with careful experimental work, electron balances (in terms of oxygen) between 95 and 102% are obtainable. In such balances, taking due account of nitrification in the oxygen utilization, the daily influent COD mass can be accounted for by the sum of the daily COD masses in the waste sludge and effluent, the carbonaceous oxygen demand and, if anoxic conditions exist, the COD utilized in the denitrification of nitrate to nitrogen gas. Experience has indicated that where poor COD mass balances are obtained, invariably the causes can be traced to errors in operation of the experimental activated sludge system. When the COD test is conducted in accordance with Standard Methods (1985), very seldom has the error been in the COD test as a measure of the EDC.

4.3 Total oxygen demand (TOD) test

In the TOD test, the sample is oxidized at high temperature in a combustion oven housed in a sophisticated electronic instrument called a TOD meter. In the TOD meter, a very small (10–20 μl) known volume of sample is injected by means of a Hamilton syringe into a gas stream consisting of nitrogen to which oxygen is dosed at a constant rate. The nitrogen-oxygen-vaporized liquid sample mixture is passed through a high temperature (900°C) combustion column with a platinum catalyst. The oxidizable matter is oxidized, consuming oxygen from the gas stream. The reduction in the oxygen content of the gas is detected electronically and is measured, also electronically, as the mass of oxygen required to restore the oxygen content of the gas mixture to its original value. In older instruments the test result is obtained in about 5 minutes but in more modern instruments, automatic injection and faster sample throughput is possible.

The TOD is the mass of oxygen required for the oxidation of practically all oxidizable material contained in a unit volume of wastewater sample. Both carbonaceous and nitrogenous compounds are oxidized. A problem with the test is that the amount of oxygen available in the combustion oven influences the degree of oxidation of ammonia and organic nitrogen. When oxygen is in excess there is a conversion of nitrogenous material to nitric oxide, whereas when oxygen is not in excess, oxidation of the nitrogenous material may be only partial. Furthermore, nitrate, nitrite and dissolved oxygen in the sample influence the TOD test result. Because the test determines both the carbonaceous and nitrogenous oxidation potential, an additional nitrogen test is required to isolate the carbonaceous fraction of the TOD. With the large proportion of particulate matter in municipal wastewater, difficulty is found in obtaining representative and reproducible readings; samples must be thoroughly homogenized to obtain representative reading because the sample volume is extremely small (10–20 μl). However, sample homogenization increases

the dissolved oxygen contamination. The difficulties of applying the TOD test to wastewater can be overcome, but the resulting test procedure and the expensive sophisticated instrument required reduces its attraction for general routine and field use.

4.4 Total organic carbon (TOC) test

The TOC test gives a specific and absolute measure of the carbon in the sample. It operates on the same basic principle as the TOD meter except that carbon dioxide, the product of combustion (oxidation), is detected. In the TOC analyser, a small volume (10–20 μl) is injected with a syringe into a catalysed combustion furnace at 900 °C. The organic compounds are oxidized to carbon dioxide and water and the total carbon dioxide concentration is measured. The dissolved carbon dioxide (inorganic) in the sample is therefore included in the result. For high organic strength wastewaters this normally is not a problem because of the low dissolved CO_2 concentrations. If the organic carbon content only is required, then it is necessary to have a pretreatment stage in sample preparation to remove inorganic carbon, by acidification and CO_2 purging by sparging, or, to do an additional test in which the inorganic carbon only is measured. The older instruments can only measure the carbon content of soluble organic material but newer models can deal with particulate organic material.

As a procedure to estimate the electron donating capacity (or free energy release) the TOC test is not useful. This is because the organic carbon content of an organic compound is not proportional to the number of electrons it can donate. This can be demonstrated by comparing glucose and glycerol. Glucose ($\text{C}_6\text{H}_{12}\text{O}_6$) with a carbon content of $6 \times 12 = 72 \text{ gC/mol}$ can donate $24\text{e}^-/\text{mol}$ which is equivalent to a COD of $24/4 \times 32 = 192 \text{ gCOD/mol}$, which gives a COD/TOC ratio of $192/72 = 2.67 \text{ gCOD/gTOC}$. Glycerol ($\text{C}_3\text{H}_8\text{O}_3$) with a carbon content of $3 \times 12 = 36 \text{ gC/mol}$, can donate $14\text{e}^-/\text{mol}$ which is equivalent to a COD of $14/4 \times 32 =$

112 gCOD/mol and yields a COD/TOC ratio of $112/36 = 3.11$ gCOD/gTOC which is 17% higher than glucose. Therefore 17% more electrons (and therefore energy) are released per gram of C in the utilization of glycerol than in the breakdown of glucose. The organism yield coefficient in terms of TOC (i.e. gVSS synthesized per gTOC oxidized) can therefore be expected to be higher for glycerol than for glucose. However, in terms of COD, the yield coefficient can be expected to be the same for glucose as glycerol because the free energy release per electron transfer is the same. For the short-chain fatty acids $[\text{CH}_3(\text{CH}_2)_n\text{COOH}]$ the COD/TOC ratio increases as the chain gets longer (n increases from 0 to 9) starting at 2.67 for acetic ($n = 0$) and increasing to 3.1 for propionic ($n = 1$), 3.33 for butyric ($n = 2$), 3.47 for valeric ($n = 3$) up to 3.73 for capric acid ($n = 8$).

Servizi and Bogan (1963) contend that the TOC has similar advantages as the COD with regard to its application to biological growth kinetics. However, according to McCarty (1963), this is only true if the COD/TOC ratio of the organic compounds oxidized is equal to the COD/TOC ratio of the organism mass formed; generally this is not the case, so that the TOC is not a good substitute parameter for COD.

The above discussion demonstrates that, fundamentally, it is inadvisable to use the TOC test to describe the energy based behaviour of biological wastewater systems. While carbon balances over activated sludge systems are possible, provided the CO_2 gas production by the system can be measured, these require more complex measuring and analysis techniques than oxygen (or electron) based mass balances. However, the test is inferior to the COD principally because TOC is not proportional to the free energy changes that take place in biological redox reactions. This does not mean that the test does not have other uses; it can serve a valuable function as a control parameter or as an analytical method in tertiary water treatment and water reclamation where the carbon content in the water is of major significance to assess water quality.

4.5 Advantages of the COD test

From the above discussion, it is clear that the COD is the most appropriate test for assessing organic wastewater strength from a theoretical and practical point of view, in that:

- it measures the electron donating capacity (EDC) which is directly related to the free energy changes in the oxidation of organics
- it does not require sophisticated analytical equipment – it is a simple wet chemical titration procedure and, as such, is appropriate for practice in the field
- it gives a result in a short time
- it allows electron balances to be made over the biological system
- it does not oxidize ammonia so it gives only the electron donating capacity of the organic compounds
- it includes the unbiodegradable organics.

Many opponents of the COD test, particularly the proponents of the BOD test, regard the last point as a disadvantage, not an advantage. For surface water quality this may be true, but for wastewaters to be biologically treated it is important to have a measure that embraces both the biodegradable and unbiodegradable organics. This is because even though the unbiodegradable material is inert in the biological reactor, the particulate unbiodegradable material accumulates in the biological reactor and adds to the sludge production of the plant and the soluble unbiodegradable will appear in the effluent. One cannot of course *ab initio* determine what the unbiodegradable components of the COD are, but with simple kinetic models based on the COD and materials balances (Marais and Ekama, 1976) and the response of the activated sludge system to the particular wastewater, it is possible to define the unbiodegradable fractions of organics in the wastewater.

4.6 Relationships between wastewater strength tests

Over the past 10 years a large number of activated sludge kinetic theories and models

have been developed. These are practically all based on the COD as the organic energy measure and the electron mass balance feature of the COD has allowed these models to be formulated into computer simulation programs. These programs are capable of simulating the activated sludge system with remarkable accuracy over a wide variety of configurations and applications, including biological N removal. These kinetic models have also led to the development of simpler steady-state design procedures which assist with the determination of the important system parameters required for the simulation models (e.g. WRC, 1984) is one of the steady-state design procedures based on COD for activated sludge in general and biological N and P removal in particular.

The BOD₅ test has been in use for more than a century and consequently remains deeply entrenched in the practice and experience of biological wastewater treatment. However, as a test it is clearly lacking in many respects compared to the COD test. These shortcomings have caused it to fall outside the major developments that have taken place over the past two decades in biological wastewater treatment design procedures and modelling. Indeed, for biological wastewater treatment (as distinct from surface water quality standards) there remain very few cogent reasons for continuing to use the BOD₅. Nevertheless, in order to make the theories and developments in activated sludge biological wastewater treatment based on COD accessible to those continuing to use the BOD, or TOC, some information is given below allowing conversion of wastewater strength to COD.

4.6.1 BOD₅ and COD

From data in the literature (Table 9.3) there is an approximate ratio between the COD and BOD, for influent municipal wastewater (equation 14):

$$\text{COD} \cong 2\text{BOD}_5 \quad (14)$$

The data in Table 9.3 are for settled wastewater which has had a significant proportion of the

TABLE 9.3 COD:BOD₅ ratio of settled sewage

<i>Reference</i>	<i>Mean</i>	<i>Range</i>	<i>No. of data</i>
Smith and Eilers (1969)	2.06	1.90–2.27	12
Boon and Burgess (1972)	1.88	1.73–2.23	17
Dixon <i>et al.</i> (1972)	2.04*	1.95–2.16	

* Average of four years operation, 1967–1970.

settleable solids removed. The settleable solids contain a significant proportion of unbiodegradable particulate organics so the COD/BOD₅ ratio can be expected to be lower for settled wastewater than for raw (unsettled) wastewater. With biological growth and endogenous respiration kinetics based on COD and the first order BOD curve, it is possible to devise theoretically a COD/BOD₅ ratio (Hoover *et al.*, 1953; Ekama and Marais, 1978). For settled wastewater (with a low unbiodegradable particulate COD fraction), the theoretical ratio is between 1.8 and 1.9. For raw wastewater (high unbiodegradable particulate COD fraction) the ratio is 2.0 to 2.1. These approximate ratios between COD and BOD₅ are very useful, but caution should be exercised in using them as they can vary between wastewaters depending on the wastewater characteristics, i.e. type and quantity of industrial wastewater contribution, permissibility of garbage grinding and other community practices, etc. The ratios given are strictly for influent municipal wastewater principally of domestic origin. The ratio COD/BOD₅ changes as the wastewater passes through the biological treatment plant, usually increasing. The ratio increases because, as the biodegradable organics are degraded, so the oxygen utilization for this reduces, while the unbiodegradable organics, which exert a COD but not a BOD₅, remain unchanged in their passage through the plant. The COD:BOD₅ ratio for biological wastewater treatment plant effluents ranges between 4 and 12 depending on the unbiodegradable soluble COD fraction, effluent suspended solids concentration and nitrification in the BOD test.

For the effluent of a biological wastewater treatment plant, the BOD₅ gives a relative but

uncertain measure of the energy still available for rapid (5 day) degradation in the receiving stream. Unless the BOD_u/BOD_5 ratio for the effluent is known, the 5-day test value is no more than a rough estimate of the pollution potential (i.e. oxygen demand) of the effluent discharge. The BOD gives no estimate of the 'unbiodegradable' (in terms of the time under treatment in the plant) organics. The COD test includes the unbiodegradable organics but the biodegradable and unbiodegradable cannot be determined separately in the test. However, with the aid of kinetic models of the biological treatment process, it is possible to obtain estimates of the unbiodegradable and biodegradable effluent COD fractions (Marais and Ekama, 1976; WRC, 1984). For long sludge age activated sludge systems (>5 days) treating normal municipal wastewater, the effluent biodegradable COD fraction is very small, less than 2% of the influent COD, and usually can be ignored.

4.6.2 TOC and COD

The TOC has found very little application in biological wastewater treatment, although there are some instances of its use in research. A ratio TOC to COD is therefore unlikely to be required. In the discussion of energy measurement above the ratio electron donating capacity (EDC) per carbon was mentioned. This is the same as the COD/TOC ratio and therefore for the various organics like glucose, short-chain fatty acids, glycerol and others, ratios between 2.7 and 3.3 were derived. For influent municipal wastewater, which comprises mostly relatively simple organics, a ratio of 3.0 is usually employed. For municipal wastewater effluents, the ratio can change substantially because the unbiodegradable organics, which generally are more complex in structure, form the major constituents.

4.6.3 COD/VSS ratio

The great advantage of the COD, which became apparent in the discussion of bioenergetics above, is that it gives a direct measure of the electrons (or energy) built into the organism

mass formed. On the basis of the stoichiometric formula assumed for organism mass ($C_5H_7O_2N$) its COD was calculated above to be 1.42 gCOD/g organism mass (see Table 9.1). For a pure culture system growing on a completely biodegradable soluble substrate in a high dilution rate chemostat (i.e. short sludge age), the volatile suspended solids (VSS) formed can be accepted to be virtually entirely organism mass. Therefore the COD/VSS ratio for such pure systems can be measured by measuring the COD and the VSS concentrations of the suspended solids material that accumulate in the reactor. Measured values on such systems give values close to 1.42 mgCOD/mgVSS (Eckenfelder and Weston, 1956).

For municipal wastewater the situation is more complex. The biomass or VSS that accumulates in the activated sludge reactor (aerobic or anoxic-aerobic) comprises three components:

1. the active heterotrophic organisms (nitrifier active mass is too low (<2%) to contribute significantly to the VSS)
2. the unbiodegradable particulate material originating from the organisms themselves called endogenous residue and
3. the unbiodegradable particulate material from the influent wastewater called inert solids.

The proportion of the VSS that these component fractions attain depends on the wastewater characteristics (fraction of unbiodegradable particulate organics in the wastewater) and the sludge age of the system. Generally, the active organism fraction decreases as the sludge age increases, while the two unbiodegradable particulate VSS components increase at different rates with sludge age. Consequently, if each component part of the VSS had a different COD/VSS ratio then it would be expected that the COD/VSS ratio of the mixed liquor would change with sludge age. This was found not to be the case (Fig. 9.7 and Table 9.4). Consequently, Marais and Ekama (1976) concluded that the COD/VSS ratio of the different components of the VSS can be accep-

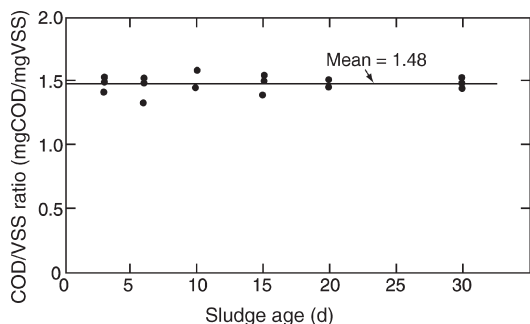


Fig. 9.7 Relationship of COD/VSS ratio of the mixed liquor with sludge age from laboratory scale aerobic completely mixed activated sludge systems. The constancy of the ratio with sludge age indicates that the ratio of the constituent fractions of the mixed liquor are the same and equal to 1.48 mgCOD/mgVSS.

ted for practical purposes to be the same, allowing a global measured value of 1.48 mgCOD/mgVSS to be used. This value is acceptably close to the theoretical 'organism VSS' value of 1.42 mgCOD/mgVSS. The global COD/VSS ratio of 1.48 mgCOD/mgVSS has been found to apply to all activated sludge systems in which heterotrophic organisms dominate the biocenosis, i.e. fully aerobic, anoxic/aerobic and also biological excess P removal systems. Even though the biochemical behaviour of the polyphosphate accumulating organisms is different to that of the

aerobic and facultative ones, a very similar COD/VSS ratio (f_{cv}) and yield coefficient (Y_g) have been found applicable to them (Wentzel *et al.*, 1989).

5 CLOSURE

By considering the bioenergetics of heterotrophic organisms, the most important group in biological wastewater treatment, it is demonstrated that the chemical oxygen demand (COD) test is the most appropriate and convenient for measuring wastewater organic content (strength). The COD measures the electron donating capacity of the organics and this is not only closely linked to the energy content of the wastewater, but also is intimately associated with the reduction-oxidation (redox) reactions whereby organisms degrade organics and grow. This association establishes two important principles in biological wastewater treatment system modelling:

1. the organism growth yield coefficient and oxygen utilization are directly linked and quantified in terms of biological COD (organic) utilization, and
2. electron, or COD, balances can be conducted over the biological treatment system in which the influent COD mass is reconciled with the effluent COD mass, oxygen mass utilized and COD mass of sludge produced over a defined unit of time.

TABLE 9.4 Experimental activated sludge COD:VSS ratios (Marais and Ekama, 1976)

Sludge source	COD/VSS ratio	96% Confidence interval	No. of tests
Bellville full scale plant	1.43	± 0.01	230
Laboratory full aerobic units 2.5–30 days (see Fig. 9.1)	1.48	± 0.08	120
Aerobic digested sludge from laboratory system above	1.54	± 0.17	80
Raw sewage	1.45	± 0.04	180
Wine distillery raw waste	1.43	± 0.01	13

Combining these principles with the practical simplicity of the test and the rapidity with which it gives a result, makes the COD test an almost ideal test for monitoring the progress of biological reactors in wastewater treatment research and practice. The above two principles form the basis of the steady state and general kinetic activated sludge models of Marais and Ekama (1976), Dold *et al.*, (1980), Van Haandel *et al.* (1981) and WRC (1984). They have paved the way to transform activated sludge design procedures and models from an empirical approach,

using a disconnected but associated assemblage of formulae, to an integrated process modelling approach based on mass balance and continuity principles. Modern design procedures are based on these principles and are essentially simplifications for steady-state conditions of the more complex formulations in the general kinetic simulation models such as those of the UCT (UCTOLD – Dold *et al.*, 1991; UCTPHO – Wentzel *et al.*, 1992) and IAW (ASM No. 1 – Henze *et al.*, 1987; ASM No. 2 – Henze *et al.*, 1994).

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Part II Water and Excreta Related Diseases

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Microorganisms and disease

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1 INTRODUCTION

Many aspects of daily life are dependent upon the activities of the smallest of organisms (microorganisms): from the preparation of food (e.g. yoghurt, bread, cheese) and drink (e.g. wine and beer), the production of antibiotics (e.g. penicillin), the natural breakdown of decaying material (composting), sewage treatment (activated sludge, filtration beds) – the list is endless. Equally, they can also be a nuisance to mankind causing, for instance, the corrosion of concrete pipes, spoilage of food and, of course, disease. Indeed, the history of mankind is inextricably dominated by a close interaction with the microbial world. However, the most effective manifestation of the activities of microorganisms is the range of diseases which they may give rise to in susceptible hosts.

2 HISTORY OF DISEASE

It is impossible to pinpoint exactly when man became aware of the significant effect that microorganisms had on his activities, but it was certainly before the establishment of the science of microbiology. Milk, butter and cheese were used by the Egyptians as early as 3000 BC, while between 3000 and 1200 BC the Jews used salt from the Dead Sea to prevent the deterioration of foods (Jay, 1992). Wines are known to have been prepared by the Assyrians by 3500 BC and fermented sausages were prepared and consumed by the Babylonians and Chinese as far back as 1500 BC.

The recognition of disease goes back at least 5000 years, although the concept of a microorganism was unknown at that time. Smallpox was familiar to the Chinese in 1000 BC and the mummy of Rameses V (who died in 1157 BC) showed clear signs of the disease (Morris and Waite, 1981) before it became a scourge of Western Europe in the 18th century. Yellow fever (a viral disease) has been known ever since European sailors first visited West and Central Africa. Influenza has been recognized in Europe since the 16th century and has shown a remarkable tendency to cause periodic pandemics interspersed with long periods of relatively little infection.

Over 4000 years ago, the Egyptians realized that contamination of drinking water supplies by dead dogs, rats etc. was a cause of illness among the populace and took measures to protect their drinking water supply. Such action was based entirely on the necessity rather than any recognition of a microbial cause. Similarly the Romans recognized the risks associated with drinking polluted water and developed the habit of throwing silver coins into the water which, although they did not understand why, rendered many of the harmful bacteria innocuous.

Girolamo Fracastoro, a 16th century Italian scholar, proposed a disease theory long before the existence of microorganisms was demonstrated. He believed that disease occurs when 'seeds too small to be seen pass from one thing to another' (Atlas, 1995). However, it was not until the late 19th century that Robert Koch showed that specific microorganisms could be related to specific diseases. By the development

TABLE 10.1 Koch's Postulates

1	The microorganism must be present in all hosts suffering from the disease and absent from those without disease
2	The microorganism must be capable of being grown outside the diseased host as a pure culture
3	When such a pure culture is administered to a healthy host then the symptoms of the disease must develop
4	The microorganism must be capable of re-isolation from the experimentally infected host and shown to be identical to the original microorganism

of his *Postulates*, Koch argued that a microorganism isolated from a disease case, when administered to a healthy host, would initiate the same disease in that host. Koch also demonstrated the possibility of growing disease-causing organisms outside the host which, when introduced into a healthy host, could result in disease. Koch's Postulates are summarized in Table 10.1.

Disease is particularly obvious when populations come under stress, e.g. warfare and natural disasters. Such events are usually correlated with the breakdown of social structure and the failure of hygiene measures. Wars, for instance, often lead to a concentration of people, the intermixing of populations and the diversion of resources. These lead to a decrease in hygiene and medical care, with an increase in malnutrition and even famine. All lead to social disintegration and the stressed individual is particularly susceptible to disease. As an example: before the First World War, malaria and typhus (both with insects as vectors) occurred in Europe, albeit at low levels. However, the conditions endured during the war caused these two diseases to erupt into epidemics (Zinsser, 1934). Similarly, two million Greeks contracted malaria during the Second World War after German occupation of the country despite malaria-free conditions beforehand (Peterson, 1995).

3 WHAT ARE MICROORGANISMS?

The term 'microbe' or 'microorganism' is generally taken to refer to those small biological

entities that can only be seen with the aid of magnification – although this is not always the case. They include the viruses, bacteria, fungi, algae and protozoa. (For the purpose of this review, the larger parasitic helminths etc. have not been included.) They can vary in size from less than 20 nm (some viruses) to several microns (protozoal parasites). Nearly all microorganisms show complex organizational structure, possessing the means to reproduce (DNA and RNA) and carry out complex biochemical processes. The exceptions are the viruses, which possess either DNA or RNA (never both) and do not have the means to carry out biochemical processes, i.e. they are obligate intracellular parasites. The bacteria, in contrast, are capable of independent growth and multiplication given a suitable source of nutrients.

4 INFECTION VERSUS DISEASE

The entry of a microorganism into a host body in itself does not necessarily lead to infection, let alone disease. Many inhabit the gut, skin and mucosal surfaces without any apparent invasion of the host tissue. These are essentially commensal organisms, which utilize the available nutrients in their immediate surrounding environment to sustain their viability. When this benign relationship is disrupted it is possible that such organisms may invade the host tissue – however, organisms such as *Escherichia coli* (used as an indicator of faecal pollution) may remain in the host gut without any apparent ill effects on the host.

Some microorganisms actively invade host tissue, i.e. they are pathogenic – this is the process of infection. In such cases the organism multiplies and is released into the environment to infect other susceptible hosts. When the activity of such an infection is such that it causes a reaction in the host, this is termed disease. In most cases, infection at best leads to a mild disease often not recognized. However, in extreme cases the disease that develops may vary from mild, through a recognized sequence of symptoms and may even result in host death.

Manifestation of an infection as a disease may result in stomach upsets, skin rashes, headache, pulmonary problems, muscular paralysis, blindness, encephalitis, arthritis and many other clinical conditions.

The relationships between microorganisms, hosts and disease are dynamic. Usually the host is healthy and environmental conditions, e.g. social factors, genetic, lifestyle etc., do not allow pathogens to become a problem. In theory, virtually any microorganism can lead to infection (and possibly disease) under suitable conditions. The infectious microorganism (because of inherent properties or failure of the host defences) is able to establish itself within the host. Provided sufficient numbers of microorganisms can penetrate into the host and reach the target tissues, then the process of pathogenesis may begin.

Thus, infection indicates the penetration into and the multiplication of an agent within the host and is determined largely by factors governing exposure to the agent and the susceptibility of the host. Disease represents the host's response to infection when it is severe enough to evoke a recognizable pattern of clinical symptoms. The factors influencing the occurrence and the severity of this response vary with the particular pathogen involved and their portal of entry, but the most important determinants for many common infections lie within the host itself. Of these, the age of the host at the time of infection is most crucial, with the young and old being particularly susceptible. Those who have diminished host defences, i.e. are immunocompromised (e.g. transplant patients and AIDS sufferers) are also at great risk of acquiring infections.

5 INFECTIVE DOSE

The number of pathogens that are necessary to enter the host to establish an infection is the *minimum infective dose* (MID). The level necessary to initiate infection varies widely, being dependent upon the pathogen itself, the susceptibility (i.e. the immune status) of the host,

route of infection, environmental factors and even the way in which the infective dose was experimentally determined.

It has already been pointed out that in cases of stress (such as war and famine), hosts are particularly vulnerable to infection, often initiated by low levels of microorganisms and depressed body defences. In other cases, ingestion of high levels of organisms may be necessary to initiate infection because of the effect of acid in the stomach – protection from such physiological conditions, e.g. by encasement in food particles, may allow much lower levels of pathogens to initiate infection. Such is the case with salmonellosis, where it is generally accepted that 10^4 microorganisms may need to be ingested. However, *Salmonella typhimurium* was capable of causing disease in 361 people after consumption of contaminated chocolate which had bacterial levels as low as 10/100 g (Kapperud *et al.*, 1990).

That erroneous infective dose values can easily be arrived at by the use of an insensitive assay procedure is illustrated by a study of the MID for Echovirus 12 (Schiff *et al.*, 1984). The report found an apparent low MID but a reassessment of the virus preparation used in the volunteer studies using a different assay technique showed that the administered dose was at least a 100-fold higher! (Ward and Akin, 1984).

Thus, the infective dose is the minimum intake of a pathogen that will elicit a response from the host (this may be a serological conversion with no evidence of disease or may lead to a whole variety of medical conditions). However, infective doses must be viewed with some caution as:

- they are often extrapolated from epidemiological investigations
- they are obtained by feeding of the pathogen to healthy young adult volunteers
- they are only best estimates based on a limited database from outbreaks
- they are worse case estimates
- because of variables of the microorganism and the host, they cannot be directly used to assess risk.

TABLE 10.2 Factors affecting derivation of infective dose

Pathogen variables	Variability of gene expression of multiple pathogenic mechanisms Potential for damage or stress of the pathogen Interaction of pathogen with food menstruum and environment Susceptibility to pH Immunological 'uniqueness' Interaction with other organisms
Host variables	Age, general health, pregnancy, occupation Malignancy Metabolic disorders, alcoholism, cirrhosis, gastric acid variation (antacids, natural variation, dilution) Nutritional status, amount of food eaten Immune competence Surgical history Genetic disturbances

Some factors affecting the derivation of MID_s are shown in Table 10.2.

The understanding of the lack of certainty of the infective dose is essential if the microbiologist is to carry out meaningful epidemiological investigations of disease outbreaks and assessment of risk of populations to disease.

6 IMMUNITY

The continual exposure to microorganisms through contact in the environment, consumption of contaminated foods and water, inhalation of aerosols etc. does not result in continuous disease in the individual. This is because of a complex network of overlapping defence systems that protect the host from potentially pathogenic microorganisms. This immunity determines whether the host is resistant to pathogens or susceptible to infection and disease. Disease may, therefore, be regarded as a failure of this defence system.

Defences against pathogens are non-specific and immunospecific responses. Non-specific responses are especially important in preventing infections. Physical barriers include the skin, mucosal membranes and fluid flow

(e.g. tears from the eyes), which block the entry of the pathogen into the host. Probably the most important of the chemical defences is acidity, as many microbes are sensitive to low pH conditions – thus the stomach is a particularly important barrier to prevent invasion of the small intestine. Other chemicals in the host also play an important role in preventing infection, including enzymes, interferons, iron-binding proteins and complement (a blood component).

The presence of phagocytic cells, macrophages, which occur throughout much of the host body and are capable of engulfing, killing and digesting microorganisms, is an additional non-specific defence to infection by pathogens. Phagocytosis is stimulated by an abnormal rise in body temperature (fever) which increases the rate of enzymatic reactions that lead to degradation of microorganisms and repair of damaged tissue, intensifies the action of interferons and causes a reduction in blood iron concentration (needed by many microorganisms for replication). In some situations, an inflammatory response may result, e.g. in skin abrasions. Inflammation is a generalized response to infection or tissue damage, aiming at localization of the invading pathogen and arresting the spread of infection. Inflammation is usually characterized by reddening of the localized area by blood vessel dilation, swelling, pain and elevated temperature.

The non-specific defences are augmented by a second mechanism – the specific immune response. This is a learned response that recognizes specific substances which trigger the formation of antibodies. This physiological response is especially important against infection and for protection against disease. The system is so essential that failure, such as with infection by organisms affecting the immune system directly (e.g. the human immunodeficiency virus, HIV), can rapidly result in severe disease and mortality caused by secondary infections. The adaptive immune response is characterized by specificity, memory and the acquired learning ability to detect foreign substances. For example, exposure to measles and diphtheria results in life-long immunity. In other cases, e.g. viral gastroenteritis, immunity

may be relatively short-lived, perhaps only a few months, allowing many episodes of the disease during a lifetime.

The immune system responds to antigens, which are specific components of each type or group of microorganisms. These activate the immune response ('learning') followed by a rapid and specific secondary response. Thus, the host only acquires immunity after exposure to the pathogen and full immunity is only gained after the secondary immunological response. Acquiring immunity protects the host against future infections by pathogens that the body has learned to recognize. This acquired immunity results in exposure to many pathogens resulting in disease only once in a lifetime – this response has been the basis for the development of vaccines against the more serious disease-causing pathogens. (The body's defence mechanisms against pathogens are described in Atlas, 1995).

7 EPIDEMIOLOGY

The idea that environmental factors can influence the occurrence of disease is not a new one. Over 2000 years ago, Hippocrates and others postulated that this was the case, inadvertently setting the scene for the science of *epidemiology* (Beaglehole *et al.*, 1993). However, it was not until the 19th century that the relationship between disease, human population and the environment in which they lived was studied.

Perhaps the most famous founding father of epidemiology was, appropriately, John Snow, a physician in London who was trying to deal with an outbreak of cholera in 1854. His observation that the distribution of cholera cases correlated with the use of water from the Broad Street Pump, together with determination that the water supply was indeed contaminated with faecal material, led to a most dramatic containment of an outbreak by the simple expediency of removing the pump handle! Cholera cases declined markedly (but not altogether) as a consequence. Snow's studies (Snow, 1855) were only one aspect of

a wide-ranging series of investigations being undertaken at the time which involved a consideration of physicochemical, biological, sociological and political factors (Cameron and Jones, 1983).

Snow was able to arrive at his conclusions by painstakingly documenting cholera cases and correlation of the comparative incidence of cholera among subscribers to two of the city's water supply companies. He found that cholera occurred more frequently in customers of one water company, the Southwark & Vauxhall (Table 10.3). This company, he observed, drew its water from the lower reaches of the River Thames where it was contaminated with London's sewage. In contrast, the Lambeth Company's supply came from the relatively unpolluted River Thames further upstream.

The modern day concept of epidemiology has developed considerably since Snow's monumental conclusions. The epidemiology of infectious disease is now concerned with the circumstances under which both infection and disease occur in a population and the factors that influence their frequency, spread and distribution. Such approaches are now regularly applied in the investigation of outbreaks of disease under all situations, e.g. the waterborne outbreak of cryptosporidiosis at Milwaukee in 1993 (MacKenzie *et al.*, 1996) and the *E. coli* O157 outbreak associated with meat products in Scotland in 1994 (Davis and Brogan, 1995).

The excellent book on basic epidemiology published by the World Health Organization is recommended for further reading (Beaglehole *et al.*, 1993). How society has developed the means to monitor, assess and control infectious

TABLE 10.3 Cholera in London 8 July to 26 August 1854 – relationship to water supply (Snow, 1855)

Water supply company	Number of deaths	Death rate/100 000
Lambeth	18	94
Southwark & Vauxhall	844	503

diseases is described by Williams (1985) who details the development of the UK's Public Health Laboratory Service.

8 TRANSMISSION ROUTES

An infectious disease is an illness caused by transmission of the pathogen from an infected host to a susceptible host, either directly or indirectly. Communicable disease is the most important of health problems in all countries. In most developing countries, communicable diseases are the greatest cause of morbidity and mortality in all age groups, whereas in developed countries, diseases associated with the respiratory tract, in particular, are the commonest cause of mortality in the young and old. The emergence of previously unrecognized diseases has greatly stimulated the need to gain epidemiological data, which can lead to risk assessment and to appropriate control programmes.

One of the most important epidemiological factors that needs to be understood before control measures can be undertaken to prevent or control disease outbreaks, is the means by which the pathogens move from infected to susceptible hosts. Some routes of transmission are summarized in Table 10.4.

All routes can be interrupted, so offering a means of disease control and prevention. For example, water- or food-borne disease can effectively be prevented by sound hygiene established through educational programmes.

Transfer of diseases, such as AIDS and hepatitis, by the use of contaminated needles by drug addicts can be reduced by using use-once needles and the avoidance of needle sharing.

To be successfully transmitted, the pathogen needs to be able to survive outside the host body without loss of viability until a susceptible host is infected. For some routes of transmission, e.g. genital, the transfer time between hosts can be very short, whereas with others, e.g. those pathogens excreted in the faeces, there is a need to be able to survive for considerable lengths of time. In the latter case, the pathogen has to be able to survive a host of environmental factors, some of which may be designed to prevent their transmission. The level of intervention (whether it be water or food production, personal hygiene, hospital practices etc.) will depend upon the resources available, what is being controlled and the level of protection desired.

9 THE ECONOMIC AND SOCIOLOGICAL COST OF DISEASE

Diseases cause suffering among all populations on the planet. It also carries with it a substantial cost in economic, as well as sociological, terms. During 1995 in Sweden, 25% of a 1400 population reported gastroenteritis caused by *Campylobacter* contamination of the drinking water supply (Andersson *et al.*, 1997). An assessment of the cost of the outbreak was attempted (taking into account

TABLE 10.4 Examples of routes for disease transmission in humans (Beaglehole *et al.*, 1993)

Exit route	Transmission route	Entry route	Example
Respiratory	Bite	Skin	Rabies
	Aerosol	Respiratory	Legionella
	Mouth to hand	Oropharyngeal	Herpes simplex
Gastrointestinal tract	Faeces to hand	Mouth	Salmonella
	Faeces to water	Mouth	Cryptosporidium
Skin	Air	Skin	Warts
Blood	Insects	Skin	Typhus
	Blood transfusion	Skin	Hepatitis B
	Needles	Skin	HIV
Genital	Intercourse	Genital	Syphilis
Placental	Vertical to embryo	Blood	Rubella

the cost of rectifying water treatment failure, medical costs, time off work costs, use of bottled water etc.), with the finding that a conservative final economic burden was in the order of \$1 million. The cost of the much larger Milwaukee outbreak of cryptosporidiosis remains to be finalized, but is likely to run into several hundreds of millions of dollars.

Outbreaks in developing countries have little value in an economic sense. Here the cost is predominantly one of human misery with death being all too common.

10 CONCLUSIONS

Infectious diseases continue to be an important public health problem world-wide. Access to sufficient water and food of suitable quality is a prime requirement for the improvement and maintenance of public health. Education of susceptible populations in good hygienic practices (especially personal) is particularly important. Avoidance of situations which would lead to the ideal conditions for disease outbreak, e.g. warfare, famine, floods, social deprivation, would also contribute greatly to safeguarding public health. However, all these require financial and physical resources, as well as a willingness to address public health matters on a personal and a political level.

The benefit of sound public health policies is self-evident as the pioneering work of John Snow adequately demonstrated. An openness of mind to recognize that 'emerging pathogens' may result in a serious threat to public health (e.g. *Cryptosporidium*, *E. coli* O157, Ebola, antibiotic-resistant tuberculosis – causing *Mycobacterium*) is equally essential. Complacency about the importance of disease to Man's welfare is reprehensible.

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Unitary environmental classification of water- and excreta-related communicable diseases

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1 INTRODUCTION

In the 30 years since Bradley developed his environmental classification of water-related diseases (White *et al.*, 1972), and the 19 years since Feachem *et al.* (1983) presented their environmental classification of excreta-related diseases, several new water-related and excreta-related pathogens have been identified (Table 11.1). Given the recent and continuing concern with the risks to global public health posed by these and other newly emerging (and re-emerging) pathogens (see, for example Ewald, 1994; Wilson *et al.*, 1994; Scheld *et al.*, 1998), it is appropriate to reconsider these earlier classifications of water- and excreta-related diseases and, due to the fact that many (but not all) water-related communicable diseases are also excreta-related, combine them into a unitary classification which is both comprehensive and up-to-date, while remaining useful to engineers in the design of water supply and sanitation systems, especially those for low-income communities in developing countries.

2 ENVIRONMENTAL CLASSIFICATION OF WATER- AND EXCRETA-RELATED COMMUNICABLE DISEASES

An environmental classification of disease groups, such as water-related and excreta-related

diseases, is more useful to environmental engineers than one based on biological type (virus, bacterium, protozoon or helminth) as it groups the diseases into categories of common environmental transmission routes. Thus an environmental engineering intervention designed to reduce the transmission of pathogens in a particular category is likely to be effective against all pathogens in that category, irrespective of their biological type.

We present our unitary classification of water- and excreta-related communicable diseases in Table 11.2. Category I comprises the faeco-oral diseases that are transmitted from person to person either by faecally-contaminated water (the classical waterborne transmission route) or through the unavailability (or lack of use) of a sufficient quantity of water for personal and domestic hygiene (Bradley's water-washed route). Category II contains the skin and eye infections which are not faeco-oral but transmitted by the water-washed route.

Bradley's concept of water-washed disease transmission (White *et al.*, 1972) represents the single most important development in our understanding of water-related disease since Snow's (1849, 1855) demonstration of the waterborne transmission of cholera (see Longmate, 1966). In low-income communities in developing countries with inadequate water supplies, the direct person-to-person water-washed

TABLE 11.1 Major water- and excreta-related pathogens recognized since 1973^a

Year	Pathogen	Type	Disease
1973	Rotavirus	Virus	Diarrhoea
1976	<i>Cryptosporidium parvum</i>	Protozoon	Acute enterocolitis
1977	<i>Legionella pneumophila</i>	Bacterium	Legionnaires' disease
1977	<i>Campylobacter jejuni</i>	Bacterium	Diarrhoea
1982	<i>Escherichia coli</i> O157:H7	Bacterium	Haemorrhagic colitis, haemolytic uraemic syndrome
1983	<i>Helicobacter pylori</i>	Bacterium	Gastric ulcers, stomach cancer
1985	<i>Enterocytozoon bienusi</i>	Protozoon	Diarrhoea
1986	<i>Cyclospora cayetanensis</i>	Protozoon	Diarrhoea
1988	Hepatitis E	Virus	Enteric hepatitis
1991	<i>Encephalitozoon hellem</i>	Protozoon	Conjunctivitis
1992	<i>Vibrio cholerae</i> O13	Bacterium	Cholera
1992	Hepatitis F	Virus	Enteric hepatitis

^a Adapted from Satcher (1995), Grabow (1997) and Lederberg (1997).

transmission of faeco-oral diseases is responsible for a much greater proportion of these diseases than their more indirect waterborne transmission. Thus engineering interventions in water supply for these communities should concentrate first on supplying them with adequate volumes (around 30 litres per person per day) of water to minimize water-washed transmission, rather than with high quality treated water to reduce waterborne transmission (Feachem, 1975) – in an ideal world both would be done, but the expense of the latter generally precludes its consideration, especially in the rural tropics.

The water-washed transmission route is also likely to be important even in areas with an adequate water supply but where personal and/or domestic (including food) hygiene is poor – *Operation Clean Hands*, launched by the American Society for Microbiology (Cassell and Osterholm, 1996), is an example of the recognition of the need to reduce water-washed disease transmission in a highly developed society.

As noted by Feachem (1975), 'the term "waterborne disease" has been, and still is, greatly abused by public health and water engineers who have applied it indiscriminately so that it has almost become, synonymous with "water-related disease".' Regrettably this is true today – for example, the WHO Regional Office

for Europe's recent report entitled *Eradication of water-related diseases* (WHO (1997)) refers exclusively to waterborne diseases, and so ignores water-washed diseases in particular, even though these are a significant health problem in Europe (Letrilliart *et al.*, 1997). Water engineers need to be aware that water-related diseases are not just simply waterborne diseases, and that other categories of water-related disease, especially water-washed diseases, are generally of greater epidemiological significance, particularly in developing countries. This is not to deny the importance of truly waterborne disease (for example, Payment *et al.* (1997) found that between 14 and 40% of gastrointestinal illnesses were attributable to the consumption of tap water meeting current North American microbiological quality requirements), but to emphasize the greater importance of water-washed diseases.

Category I includes infection with *Helicobacter pylori*, which is transmitted faeco-orally (and also oro-orally) and which causes gastritis and stomach cancer (Blaser, 1998); its prevalence is high (30–90% in industrialized countries and 80–90% in developing countries), and it is the only bacterium designated as a known human carcinogen (IARC, 1994; see also Mara and Clapham, 1997). Category I also includes several other 'new' pathogens (see Table 11.1): rotavirus and *Campylobacter jejuni*

TABLE 11.2 Unitary environmental classification of water- and excreta-related diseases

Category	Environmental transmission features	Examples	Control strategies
I. Faeco-oral waterborne and water-washed diseases	Non-latent (except <i>Ascaris</i>) No intermediate host Infectivity: medium to low (bacteria), high (others) Persistence: medium to high (bacteria), low to medium (others, except <i>Ascaris</i> : very high) Able (bacteria) and unable (others) to multiply outside host	<i>Viral</i> : Hepatitis A, E and F Poliomyelitis Rotaviral diarrhoea Adenoviral diarrhoea <i>Bacterial</i> : Campylobacteriosis Cholera <i>Helicobacter pylori</i> infection Pathogenic <i>Escherichia coli</i> infection Salmonellosis Typhoid and paratyphoid Yersiniosis <i>Protozoan</i> : Amoebiasis Cryptosporidiasis <i>Cyclospora cayetanensis</i> diarrhoea <i>Enterocytozoon bienusi</i> diarrhoea Giardiasis <i>Isospora belli</i> diarrhoea <i>Helminthic</i> : Ascariasis Enterobiasis Hymenolepiasis	Improve water quantity, availability and reliability (water-washed disease control) Improve water quality (waterborne disease control) Hygiene education
II. Non-faeco-oral water-washed diseases	Non-latent No intermediate host High infectivity Medium to high persistence Unable to multiply	Skin infections (scabies, leprosy, yaws) Eye infections (trachoma, conjunctivitis, including that caused by <i>Encephalitozoon hellem</i>) Louse-borne fevers	Improve water quantity, availability and reliability. Hygiene education
III. Geohelminthiasis	Latent Very persistent Unable to multiply No intermediate host Very high infectivity	Ascariasis; trichuriasis; hookworm infection	Sanitation. Effective treatment of excreta or wastewater prior to reuse. Hygiene education.
IV. Taeniasis	Latent Persistent Able to multiply Very high infectivity Cow or pig intermediate host	Beef and pork tapeworm infections	As III above, plus proper cooking of meat and improved meat inspection

(continued on next page)

Table 11.2 (*continued*)

<i>Category</i>	<i>Environmental transmission features</i>	<i>Examples</i>	<i>Control strategies</i>
V. Water-based diseases	Latent Persistent Able to multiply High infectivity Intermediate aquatic host(s)	<i>Bacterial:</i> Leptospirosis Tularaemia Legionellosis <i>Helminthic:</i> Schistosomiasis Clonorchiasis Fasciolopsiasis Guinea worm infection <i>Fungal:</i> Pulmonary haemorrhage due to <i>Stachybotrys atra</i> infection	Decrease contact with contaminated water. Improve domestic plumbing. Public education Decrease contact with contaminated waters. Sanitation. Treatment of excreta or wastewater prior to reuse. Public education Drying of flood-damaged homes. Public education
VI. Insect-vector diseases		<i>Water-related:</i> Malaria Dengue Rift Valley fever Japanese encephalitis Yellow fever African sleeping sickness Onchocerciasis Bancroftian filariasis <i>Excreta-related:</i> Fly-borne and cockroach-borne excreted infections ^a Bancroftian filariasis	Decrease passage through breeding sites. Destroy breeding sites. Larvicide application. Biological control. Use of mosquito netting and impregnated bed nets Improve stormwater drainage Public education
VII. Rodent-vector diseases		Rodent-borne excreted infections ^a Leptospirosis Tularaemia	Rodent control. Hygiene education Decreased contact with contaminated water Public education

^a The excreted infections comprise all those diseases in Categories I, II and IV and the helminthic diseases in Category V.

(globally the most common viral and bacterial causes of diarrhoea, respectively); new strains or types of known pathogens, such as hepatitis E and F viruses, *Escherichia coli* O157 and *Vibrio cholerae* O139; and two new protozoa – *Cryptosporidium parvum* (responsible for the largest recorded outbreak of waterborne disease in the USA, that in Milwaukee in 1993 when over

400 000 people were infected (Mackenzie *et al.*, 1994), and recently reported to be also disseminated by waterfowl (Graczyk *et al.*, 1998); and *Cyclospora cayentanensis*, which may be waterborne (Benenson, 1995) or foodborne (Majkowski, 1997) and is unusual in that it is not immediately infective upon excretion but is latent for a few days to a few weeks. Diarrhoea

due to the protozoon *Isospora belli* is also included, an example of a re-emerging pathogen, especially (but not only) in immuno-compromised persons (Marcial-Seoane and Serrano-Olmo, 1995; Goodgame, 1996; Marshall *et al.*, 1997). Two other new protozoa, both mainly affecting the immunocompromised, are also included for completeness: *Enterocytozoon bienusi*, which causes diarrhoea (Collins, 1997), in Category I; and *Encephalitozoon hellem*, which causes conjunctivitis (Didier *et al.*, 1991), in Category II.

Categories III, IV and V are the geohelminthiases, taeniasis and water-based diseases, respectively. These are all helminthic infections, although Bradley's original category of water-based disease is broadened to include water-based bacterial diseases such as legionellosis (which can be transmitted in a sauna; Den Boer *et al.*, 1998), leptospirosis (see Mara and Alabaster, 1995), and Buruli ulcer due to *Mycobacterium ulcerans* in swamp waters (Wright, 1998). Category V also includes pulmonary haemorrhage/haemosterois due to the toxigenic fungus *Stachybotrys atra*, which occurs in infants living in homes subjected to flood-induced water damage which promotes the growth of this fungus (Dearborn, 1997).

Categories VI and VII comprise the water- and excreta-related insect vector and excreta-related rodent vector diseases, respectively. Category VI includes important mosquito-borne diseases such as malaria and Bancroftian filariasis, and also dengue fever and dengue haemorrhagic fever which are now the most important human insect-vector viral diseases; the dengue pandemic has intensified over the last two decades, and currently there are 50–100 million cases of dengue fever and several hundred thousand cases of dengue haemorrhagic fever each year (Gubler and Kuno, 1997). In category VI Bancroftian filariasis appears under both the water-related and the excreta-related subcategories as in some parts of the world its mosquito vector is *Aedes aegypti* which breeds preferentially in 'clean' water, whereas in others it is *Culex quinquefasciatus* which prefers to breed in 'dirty', i.e. excreta- or wastewater-contaminated) water, although

this distinction between clean water and dirty water breeding mosquitoes is now less clearcut than it used to be.

3 GLOBAL BURDEN OF WATER- AND EXCRETA-RELATED DISEASES

Murray and Lopez (1996b) present data on the incidence and prevalence of several major water- and excreta-related diseases in 1990. These are listed in Table 11.3. While these data are awesome in their sheer magnitude, they tell us little about disease *burden*. WHO (1980) claimed that as much as 80% of all morbidity in developing countries was due to water- and excreta-related disease, but the epidemiological evidence for this figure was not given. More recently, Murray and Lopez (1996a) give data on the global burden of disease in 1990, including that fraction attributable to poor water supply, sanitation and personal and domestic hygiene (Table 11.4). Globally, 5.3% of all mortality in 1990 was due to poor water supply, sanitation and hygiene. In the developing world as a whole the corresponding percentage was 6.7, but much higher in the Middle Eastern Crescent (8.3), India (9.0) and sub-Saharan Africa (10.7), and in developing countries this risk factor was second in importance only to malnutrition (which was responsible for 14.9% of all deaths).

When the burden of disease is expressed in disability-adjusted life years (DALYs), which equal the sum of discounted years of life lost due to premature death and discounted years lived with disability (see Murray and Lopez, 1996a), a similar picture emerges (Table 11.4) since, with water- and excreta-related diseases, the principal component (>90%) of DALYs is years of life lost, rather than years lived with disability. The percentage of total DALYs lost (i.e. DALYs lost due to all causes) attributable to poor water supply, sanitation and hygiene is 6.8 globally and 7.6 in developing countries as a whole, and again higher in the Middle Eastern Crescent (8.8), India (9.5) and sub-Saharan Africa (10.1).

Of the total number of deaths attributable to poor water supply, sanitation and hygiene, i.e.

TABLE 11.3 Global water- and excreta-related disease statistics for 1990^a

<i>Disease</i>	<i>Number</i>	<i>Remarks</i>
Diarrhoea	4 073 920 110 episodes	56% in children aged 0–4 94% in developing countries
Malaria	213 743 000 episodes	All in developing countries 87% in sub-Saharan Africa
African sleeping sickness	267 000 persons infected	All in sub-Saharan Africa
Schistosomiasis	208 276 000 persons infected	87% in sub-Saharan Africa
Onchocerciasis	5 802 000 persons infected 478 000 persons with impaired vision 356 000 blind persons	> 99% in sub-Saharan Africa > 99% in sub-Saharan Africa > 99% in sub-Saharan Africa
Leprosy	2 434 000 persons infected	70% in Asia
Dengue	415 000 episodes	92% in Asia
Japanese encephalitis	58 000 000 episodes	All in Asia
Trachoma	292 000 persons with impaired vision 192 000 blind persons	34% in sub-Saharan Africa; 34% in China; 24% in MEC ^b 34% in sub-Saharan Africa; 34% in China; 24% in MEC ^b
Ascariasis	61 847 000 persons with high-intensity infection	73% in children aged 5–14 All in developing countries
Trichuriasis	45 421 000 persons with high-intensity infection	79% in children aged 5–14 All in developing countries
Human hookworm infection	152 492 000 persons with high-intensity infection 36 014 000 persons with anaemia	84% in adults aged 15–59 All in developing countries 72% in adults aged 15–44 All in developing countries

^a The world population in 1990 was 5.3 billions, of which 3.9 billions (74%) were in developing countries.

^b Middle East Crescent, covering North Africa, the Middle East, Pakistan and the Central Asian republics of the former Soviet Union.

Source: Murray and Lopez (1996b).

attributable to water- and excreta-related diseases, 99.9% occurs in developing countries. For total DALYs lost, the corresponding percentage is 99.8.

4 APPROPRIATE CONTROL STRATEGIES

Table 11.2 lists appropriate control strategies to reduce the incidence of water- and excreta-

related communicable diseases. Generally, and especially in developing countries, these need only to be relatively simple, low-cost interventions that can, in many cases, be done by the communities involved or at least with their participation, and there is much information available on these, e.g. water supplies (Feachem, 1975; Hofkes, 1983; Morgan, 1990), and sanitation (Morgan, 1990; Mara, 1996). Vector control is discussed by Curtis (1990, 1991).

TABLE 11.4 Burden of disease attributable to poor water supply, sanitation and personal and domestic hygiene in 1990

Region ^a	Deaths	Percentage of all deaths	DALYs ^b	Percentage of all DALYs
EME	1100	<0.1	101 000	0.1
FSE	2400	0.1	128 000	0.2
IND	839 900	9.0	27 463 000	9.5
CHN	81 400	0.9	4 231 000	2.0
OAI	354 300	6.4	13 192 000	7.4
SSA	875 600	10.7	28 870 000	10.1
LAC	135 300	4.5	5 183 000	5.3
MEC	378 200	8.5	13 224 000	8.8
World	2 668 200	5.3	93 392 000	6.8
Developed regions	3500	<0.1	229 000	0.1
Developing regions	2 664 700	6.7	93 163 000	7.6

^a EME, established market economies; FSE, formerly socialist economies of Europe; IND, India; CHN, China; OAI, other Asia and islands; SSA, sub-Saharan Africa; LAC, Latin America and the Caribbean; MEC, Middle Eastern Crescent.

^b DALYs, disability-adjusted life years.

Source: Murray and Lopez (1996a).

Hygiene education is of paramount importance, and this is discussed by Nyamwaga and Akuma (1986), Boot (1990, 1991), Boot and Cairncross (1993) and Hubley (1993).

5 CONCLUDING REMARKS

The unitary environmental classification of water- and excreta-related diseases is a comprehensive categorization of these diseases into seven categories of common environmental transmission patterns. This unitary classification is likely to be useful to tropical public health engineers and other professionals as it highlights the facts that many of these diseases are both water and excreta related, and that these diseases are best controlled in the long term by sustainable (and indeed sustained, i.e. well operated and maintained) improvements in both water supply and sanitation, which are supplemented by effective programmes of hygiene education. Although convincing arguments can be made for broadening the disease concept from water- and excreta-related diseases to housing-related diseases (Mara and Alabaster, 1995) or, more generally, to cover urban or peri-urban health (Harpham and Tanner, 1995), the fact remains that water- and excreta-related diseases are

the most important of these diseases. It is in this context that the unitary environmental classification of these diseases is most useful.

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Emerging waterborne pathogens

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1 INTRODUCTION

The English dictionary defines the word 'emerging' to mean, 'to rise out of a state of depression or obscurity; to come to notice; to reappear after being eclipsed'. Thus is the state of infectious diseases throughout the world in the twenty-first century. Not only are new microorganisms being identified every year but some of the old established diseases are again gaining attention. More specifically, emerging diseases have been defined as those 'whose incidence in humans has increased in the past two decades or threatens to increase in the near future' (NIM, 1998).

The reason for this 'emergence' of infectious diseases has been attributed to many factors:

- Sensitive or susceptible populations: there is an increasing number of elderly and immunocompromised individuals (transplant patients, AIDs patients) in our communities, in addition to diabetics, infants, and pregnant women, all of whom may be more susceptible to severe outcomes (Gerba *et al.*, 1996a).
- Global transportation: the food supply comes from all over the world, thus leading to a distribution of pathogens (i.e. *Cyclospora*) and people are much more mobile and can transverse the globe in less than 24 hours, bringing infections with them.
- Antibiotic resistance: due to the widespread application of antibiotics in medicine and in agriculture, antibiotic resistance is spreading. The World Health Organization (WHO) has recently reported that

increasingly drug-resistant infections throughout the world are threatening to make once-treatable diseases incurable (<http://www.who.int/emc/amr.html>).

- Zoonotic transmission: more animals and changes in agricultural practices may lead to greater chances for microbial transmission and spread from animals to humans.
- Evolution of pathogens: RNA viruses, e.g. during replication lack repair mechanisms and can evolve quickly.
- Improved diagnostic tools: diseases previously not recognized or microorganisms that are not culturable can now be studied through the use of powerful molecular tools.

The factors describe above also apply to the growing list of waterborne pathogens. Table 12.1 describes some of the microorganisms that can be spread through contaminated water. In the last three decades, 1970 to 2000, numerous species of bacteria, parasites and viruses have been described that have caused important waterborne outbreaks throughout the world. This chapter will briefly discuss these pathogens. Because most of the microorganisms that are waterborne are spread by the faeco-oral route, their occurrence and control in wastewaters, animal wastes and drinking waters are important. Key in the future to understanding and controlling emerging pathogens in drinking water and wastewater is the application of new tools for monitoring and studying these microorganisms.

In the past, the direct detection of pathogenic microorganisms in the water environment

TABLE 12.1 Examples of ‘emerging’ waterborne pathogens of concern

<i>Pathogen</i>	<i>First described: as human pathogen/ waterborne transmission</i>	<i>Methods of detection</i>
Bacteria		
<i>Escherichia coli</i> 0157:H7 Associated with bloody diarrhea (hemorrhagic colitis) and Hemolytic uremic syndrome	1982*, Riley <i>et al.</i> , 1983; Mead and Griffin, 1998. 1991 [†] , Centers for Disease Control, 1991	Cultural methods Molecular methods (AWWA Research Division Microbiological Contaminants Research Committee, 1999)
<i>Helicobacter pylori</i> Diarrhea, peptic and duodenal ulcer disease, gastric carcinoma	1982*, Warren and Marshall, 1983. 1991 [†] , Klein <i>et al.</i> , 1991.	Cultural methods (microbiological methods) Molecular methods Autoradiography ATP bioluminescence (Velázquez and Feirtag, 1999)
<i>Mycobacterium avium complex</i> Diarrhea and respiratory disease	1982*, Greene <i>et al.</i> , 1982 1994 [†] , Singh and Lu, 1994	Cultural methods Molecular methods Chemical methods (AWWA Research Division Microbiological Contaminants Research Committee, 1999)
<i>Vibrio cholerae</i> Profuse watery diarrhea ‘rice water stools’, and vomiting.	1600* [†] , Pollitzer, 1959	Cultural methods Molecular methods (Baumann <i>et al.</i> , 1984; Sharma <i>et al.</i> , 1997; Basu <i>et al.</i> , 2000)
Parasites		
<i>Cryptosporidium</i> Profuse watery diarrhea (cholera-like), fluid loss, fever and abdominal pain	1976*, Meisel <i>et al.</i> , 1976. 1984 [†] , D’Antonio <i>et al.</i> , 1985.	Immunofluorescence assay Tissue culture methods Molecular methods (Gasser and O’Donoghue, 1999)
<i>Cyclospora</i> Explosive, watery diarrhea, fatigue, anorexia, weight loss, nausea	1979*, Ashford, 1979, 1994 [†] , Rabold <i>et al.</i> , 1994	Epifluorescence (UV illumination), acid-fast staining, molecular techniques. (Long <i>et al.</i> , 1991; Yoder <i>et al.</i> , 1996)

TABLE 12.1 (*continued*)

<i>Pathogen</i>	<i>First described: as human pathogen/ waterborne transmission</i>	<i>Methods of detection</i>
Microsporidia Gastrointestinal, pulmonary, nasal, ocular, muscular, cerebral and systemic infections	1959*, Matsubayashi <i>et al.</i> , 1959 1997†, Sparfel <i>et al.</i> , 1997	Chromotrope staining, chemofluorescent agents, Giemsa stain, immunodetection, molecular methods, tissue culture methods (Weber <i>et al.</i> , 1999; Dowd <i>et al.</i> , 1998a, 1999)
<i>Toxoplasma</i> (Flu-like symptoms). Painful swollen lymph glands in the cervical, supraclavicular and inguinal regions. Fever, headache, muscle pain, anemia, lung complications. Fetus at greatest risk	1979†, Benenson <i>et al.</i> , 1982	Mouse bioassay (Aramini <i>et al.</i> , 1998) ICR US. EPA cartridge filtration (Isaac-Renton <i>et al.</i> , 1998).
Enteric viruses		
Coxsackie virus Aseptic meningitis, herpangina, paralysis, exanthema, hand, foot and mouth disease, common cold, hepatitis, infantile diarrhea, acute hemorrhagic conjunctivitis	1947*, Dalldorf and Sickles, 1948	Tissue culture methods Molecular methods (Hurst <i>et al.</i> , 1997; Hurst, 1999; Murrin and Slade, 1997)
Hepatitis viruses Fever, nausea, abdominal pain, anorexia and malaise, associated with mild diarrhea, arthralgias, scleral icterus. Cytologic damage, necrosis and inflammation of the liver (HAV)	1950†, Bradley, 1992	Tissue culture methods Molecular methods Immunofluorescence (Smith, 2001, Sobsey, 1999, Jothikumar <i>et al.</i> , 1993, Pina <i>et al.</i> , 1998)
Norwalk-like viruses Diarrhea, vomiting, abdominal pain, cramping, low fever, headache, nausea, tiredness (malaise), and muscle pain (myalgia)	1968*, 1968†, Gerba, 1996c	Transmission electron microscopy; enzyme-linked immunosorbent assay Molecular methods (Hurst, 1999).
Rotavirus Vomiting, abdominal distress, diarrhea, dehydration, fever,	1973*, Bern <i>et al.</i> , 1992	Tissue culture methods Molecular methods (Gratacap-Cavallier, 2000)

* Pathogen.

† Waterborne transmission.

had been tedious and often required many biochemical analyses to identify the genus and/or species present. In fact, no method for cultivating the microorganism in question existed at all. The use of polymerase chain reaction (PCR, the genetic copying method) now allows for a more rapid and specific detection of microorganisms and their associated public health risks. The advantages of using this method are tremendous and the field is moving forward very quickly. Almost all the emerging pathogens discussed in this chapter have been studied in the water environment using this powerful technique.

2 BACTERIAL PATHOGENS

Early in the 1800s, when it became apparent that disinfection of drinking water could prevent waterborne disease epidemics caused by bacterial pathogens such as *Salmonella typhi* (typhoid fever), most people thought that the days of waterborne bacterial epidemics were gone forever. Unfortunately, this was not to be the case. During the recent decade, there has been a number of waterborne disease outbreaks caused by bacteria such as *Escherichia coli* and *Vibrio cholerae*. Advanced techniques that combine standard cultural methods with enhanced PCR detection methods have revealed the presence of bacteria previously not associated with waterborne disease transmission such as *Mycobacterium avium* and *Helicobacter pylori*.

2.1 *Escherichia coli* O157:H7

Escherichia coli O157:H7 is an enteropathogenic strain of *E. coli* that was first identified as a human pathogen in 1982 (Mead and Griffin, 1998). Infection with the organism can cause severe bloody diarrhea with abdominal cramping. In small children and the elderly, fluid replacement is of the utmost importance for a full recovery. A common, more serious, complication of infection with *E. coli* O157:H7 is hemolytic uremic syndrome (HUS) where

there is loss of red blood cells and kidney failure. In severe cases, HUS can cause permanent kidney damage or death.

Escherichia coli O157:H7 has been shown to survive similarly to typical *E. coli* strains under routine drinking water conditions. There have been numerous documented outbreaks of waterborne disease caused by *E. coli* O157:H7. The most recent drinking-water outbreak of *E. coli* O157:H7 took place in Walkerton, Canada (90 miles west of Toronto) in May 2000. It is estimated that there were more than 1000 cases of illness and nine deaths associated with the outbreak. It is believed that the organisms entered the town's drinking water supply when animal waste from nearby farms washed into wells during a flood earlier in the month. The town had been using chlorine to disinfect the drinking water, however, during the time of the outbreak the chlorine disinfection system was malfunctioning.

In the USA there have been several documented outbreaks of waterborne *E. coli*. In 1990, a drinking-water outbreak of pathogenic *E. coli* took place in Cabool, Missouri, with 243 cases, 32 hospitalizations and four deaths (Geldreich *et al.*, 1992; Swerdlow *et al.*, 1993). The source of the water for Cabool was deep groundwater wells with no disinfection in the distribution system. Disturbances in the distribution system included the replacement of 43 water meters and two line breaks. Disturbances to distribution systems can provide a pathway for infiltration from stormwater run-off or sewage contamination. In August 1999, unchlorinated well water, used to prepare beverages and ice at the Washington County Fair near Albany, New York, was contaminated with *E. coli* and caused more than 900 cases of illness with 65 hospitalizations and two deaths. Also in 1999, in Petersburg, Illinois more than 200 people became ill after eating and/or drinking contaminated food and water at a festival that was being held in a cow pasture. In 1996, in Atlanta, Georgia, 26 children playing at a water park became ill after ingesting pool water that had been contaminated with fecal material containing the bacteria (Barwick *et al.*, 2000). Another documented

waterborne outbreak of *E. coli* O157:H7 took place in Scotland with 496 cases (272 laboratory confirmed cases) and 19 deaths (Dev *et al.* 1994).

The largest outbreak of *E. coli* O157:H7 occurred in 1996, in Sakai City, Japan. The outbreak was due to the consumption of radish sprouts that had been washed with contaminated water. Approximately 6000 people were infected and three children died as a result of the infection (Michino *et al.*, 1999).

2.2 *Helicobacter pylori*

Helicobacter pylori has been cited as a major etiologic agent for gastritis and has been implicated in the pathogenesis of peptic and duodenal ulcer disease (Taylor and Blaser, 1991). It has also been associated with the development of gastric carcinoma (Eurogast Study Group 1993). The mode of transmission of *H. pylori* is not well characterized. Recent studies suggest that gastrointestinal dissemination may be due to vomiting in childhood (Axon, 1995). Persons living in low socioeconomic conditions have consistently been shown to have a high prevalence of *H. pylori* and the organism has also been found routinely in the feces of children living in endemic areas (Thomas *et al.*, 1992). Klein *et al.* (1991) reported that, in Peru, the water source might be a more important risk factor than socioeconomic status in acquiring *H. pylori* infection. Children whose homes had external water sources were three times more likely to be infected than those whose homes had internal water sources. Among families with internal water sources, there was no difference in *H. pylori* infection associated with income. Children from high-income families whose homes were supplied with municipal water were 12 times more likely to be infected than were those from high-income families whose water came from community wells. These findings show that substandard municipal water supplies may be important sources of *H. pylori* infection.

2.3 *Mycobacterium avium* complex (MAC)

MAC has been isolated from natural water and drinking water distribution systems in the USA and is likely present throughout the world. Members of the MAC are considered opportunistic human pathogens. There has been a dramatic increase in the number of AIDS patients infected with MAC, with 25–50% of late stage AIDS patients now infected. Evidence for environmental transmission of MAC includes: increased frequency of MAC in gastrointestinal tract of advanced AIDS patients, higher frequency of isolation of MAC from the gastrointestinal tract than from the respiratory tract, and gastrointestinal symptoms (i.e. nausea, vomiting and diarrhea) common with MAC infections (Singh and Lu, 1994). *M. avium* strains from infected AIDS patients in Los Angeles have been shown to be genetically related to isolates recovered from water to which the patients were exposed through drinking or bathing (von Reyn *et al.*, 1994; Glover *et al.*, 1994). Control of MAC can be difficult as they are extremely resistant to conventional disinfection methods and have been shown to survive 10 mg/l of free chlorine (duMoulin and Stottmeier, 1983).

2.4 *Vibrio cholerae*

Epidemics of cholera have devastated Europe and North America since the early 1800s. A lack of sanitation and an increasing population, often with limited access to clean water, has brought about numerous disease outbreaks. A total of 1,041,422 cases and 9642 deaths due to cholera in the Americas were reported by 1995 (Morbidity and Mortality Weekly Report, 1995). In 1994 non-O1 cholera was detected for the first time from the Bug River (freshwater) in Poland and Hong Kong has recently reported two outbreaks of cholera (Lee *et al.*, 1996). Although the cause of the Hong Kong outbreaks was not clearly identified, increasing pollution of coastal waters has been implicated. Further concern over the cholera epidemic stems from the discovery of a new strain of *V. cholerae*, O139, which has

shown an increased mortality rate (Lee *et al.*, 1996). Inadequate disinfection or the lack of disinfection has contributed significantly to the spread of cholera throughout Africa and Latin America.

3 PROTOZOAN PARASITES

The emerging protozoa associated with waterborne disease all produce an oocyst or spore-like structure, which are the environmental as well as the infectious stage. These life stages are very resistant to routine disinfection in drinking water and little is known about wastewater treatment and how it may affect their survival and removal. Most are associated with animals (with the exception of *Cyclospora*) and therefore the control of animal wastes is as equally important as the control of human wastes. Microscopy had been the most common method for detection of protozoa in clinical specimens. However, in the water environment, while microscopy is often used, future methods are focusing on PCR and cell culture.

3.1 *Cryptosporidium*

Cryptosporidium was first diagnosed in humans in 1976. Since that time, it has been well recognized as a cause of severe watery diarrhea, lasting several days to a week (Dubey *et al.*, 1990). Reported incidence of *Cryptosporidium* infections in the population range from 0.6 to 20% depending on the geographic locale. While there is a greater prevalence in the populations of Asia, Australia, Africa and South America, in the USA and the UK, *Cryptosporidium* has been described as the most significant cause of waterborne disease today. Detection of this organism has been reported in waters throughout the world.

Oocysts have been detected in 4 to 100% of surface waters tested at concentrations ranging from 0.1 to 10 000/100 l depending on the impact from sewage and animals (Lisle and Rose, 1995). Groundwater, once thought to be a more protected source, has shown between 9.5 and 22% of samples positive for *Cryptosporidium* (Hancock *et al.*, 1998). The largest

documented drinking water outbreak of *Cryptosporidium* in the USA occurred in Milwaukee, Wisconsin in 1993 where 400 000 people were ill and 100 died due to contamination of the water supply (MacKenzie *et al.*, 1994).

New methods are available for the detection and characterization of *Cryptosporidium*. In particular, immunomagnetic separation techniques for improved recovery (Bifulco and Schaeffer, 1993; Deng *et al.*, 1997; Bukhari *et al.*, 1998); cell culture techniques for evaluating the infectivity of the oocysts (Slifko *et al.*, 1997, 1999; DiGiovanni *et al.*, 1999) and finally, PCR methods for distinguishing the human-human genotype from the animal-human genotype (Wiedenmann *et al.*, 1998). PCR is an attractive diagnostic procedure as it is rapid, sensitive, and pathogen specific (Johnson *et al.*, 1995; Rochelle *et al.*, 1997; Kostrzynska *et al.*, 1999; DiGiovanni *et al.*, 1999). Xiao *et al.*, (1999) evaluated several PCR techniques to determine whether they detected *C. parvum* in environmental samples. The authors concluded that two nested-PCR-restriction fragment length polymorphism (RFLP) based on the small-subunit rRNA (Xiao *et al.*, 1999) and dihydrofolate reductase genes (Gibbons *et al.*, 1998) were more sensitive than single-round PCR or PCR-RFLP protocols for detection and *Cryptosporidium* species differentiation (Xiao *et al.*, 1999).

3.2 *Cyclospora cayetanensis*

Cyclospora (previously termed cyanobacterium-like body) is a single cell coccidian protozoan that has been implicated as an etiologic agent of prolonged watery diarrhea in humans (Ortega *et al.*, 1993). The organism was first described as early as 1977 (Ashford, 1979) and has been reported with increased frequency since the mid-1980s. *Cyclospora* has been described in patients from North, Central and South America, Europe, Asia and North Africa, however, the true prevalence of this parasite in any population is unknown (Soave and Johnson, 1995).

Cyclospora is now known to be an obligate parasite of immunodeficient and immunocompetent humans (Ortega *et al.*, 1993). In an

immunocompromised person the parasite can cause profuse, watery diarrhea lasting several months. The infection is much less severe in immunocompetent patients. Symptoms may range from no symptoms to abdominal cramps, nausea, vomiting and fever lasting from 3 to 25 days (Goodgame, 1996).

While *Cryptosporidium* appears to be predominantly waterborne, *Cyclospora* has been related more often to transmission through contaminated produce from a world market and the possible contamination of irrigation waters with human wastes. The differences between the protozoa and their transmission may be due to their biology and structure, size of the oocysts, need for sporulation and presence of animal reservoirs (Table 12.2). The treatment of human wastes in North America compared to lack of treatment in other parts of the developing world may also contribute to the transmission of the organism.

Two waterborne outbreaks have been reported. In June 1994, several cases of diarrhea were detected among British soldiers and dependants stationed in a small military detachment in Pokhara, Nepal (Rabold *et al.*, 1994). The drinking water for the camp was a mixture of river and municipal water that was treated by chlorination. A candle filtration system was also used to remove particulates, but was not guaranteed to filter *Cyclospora*-sized particles (8–10 μm). *Cyclospora* was detected in 75% of the diarrhea samples examined, and a water sample processed by membrane filtration taken from the camp also revealed the presence of *Cyclospora* oocysts.

Twenty-one cases of prolonged diarrhea in employees and staff physicians were noted on July 9, 1990 in a Chicago hospital (Huang *et al.*, 1995). Upon investigation, *Cyclospora* oocysts were identified in the stools and epidemiological investigations implicated the tap water in the physicians' dormitory. It was speculated that the storage tank had become contaminated. Although this outbreak has been identified as a waterborne outbreak, a plausible scenario for the contamination of the water has not been developed.

In 1996, a total of 1465 cases of cyclosporiasis occurred in 20 states in the USA and Canada. Some 55 events were identified as a part of this nationwide outbreak. The suspected vehicles of transmission for these outbreaks were Guatemalan raspberries. *Cyclospora* reappeared in 1997 (Rose and Slifko, 1999). Twenty-five clusters of outbreaks have occurred resulting in 1450 cases in nine states. The foods associated with these outbreaks included raspberries, basil, and lettuce. Some of these foods were traced to Central Guatemala. In 1998, few cases were identified in the USA; however, an outbreak did occur in Canada.

The predominant method for detecting and identifying *Cyclospora* has been by microscopy. There are currently no antibodies available for specifically staining the oocyst. However, PCR will likely turn out to be the better method for screening for the presence of this pathogen in wastewaters and potentially surface waters used for irrigation that are receiving human sewage. The amplification reaction for *Cyclospora cayetanensis* has

TABLE 12.2 Comparison of *Cryptosporidium* and *Cyclospora*

Attribute	<i>Cryptosporidium</i>	<i>Cyclospora</i>
Taxonomy	Intestinal coccidian	Intestinal coccidian
Infective unit	Oocysts 4–5 μm Immediately infectious upon excretion	Oocysts 8–10 μm Requires sporulation* in the environment, not immediately infectious upon excretion
Animal reservoir	<i>C. parvum</i> found in most mammals, can cross species barriers	<i>C. cayetanensis</i> documented only in humans
Waterborne disease	12 waterborne outbreaks in North America since 1985	1 outbreak in Chicago, 1 in Nepal

* Sporulation is a process by which the oocysts undergo maturation in the environment before becoming infectious.

been based in a nested PCR reaction that included forward and reverse oligonucleotide primers that hybridize to sequences encoding 18S rRNA (Yoder *et al.*, 1996). An oligo-ligation assay (OLA) for the differentiation between *Cyclospora* and *Eimeria* spp. (an animal enteric protozoan which is related to *Cyclospora*) has also been included in the procedure (Jinneman *et al.*, 1999).

3.3 Microsporidia

The microsporidia are obligate intracellular spore-forming protozoa that are capable of infecting both vertebrate and invertebrate hosts. Their role as an emerging pathogen in immunosuppressed hosts is being increasingly recognized. The prevalence of microsporidiosis in studies of patients with chronic diarrhea ranges from 7 to 50% world-wide (Bryan, 1995). It is unclear whether this broad range represents geographic variation, differences in diagnostic capabilities or differences in risk factors for exposure to microsporidia. Microsporidia are single-celled, spore-forming, obligate intracellular parasites that belong to the phylum Microspora. The microsporidia are considered emerging pathogens because new species have been identified as causes of disease in humans during the last 20 years. Because species of microsporidia that were recognized causes of disease in animals are now causing infections in humans, microsporidia are also considered re-emerging pathogens (Didier and Bessinger, 1999).

In the summer of 1995, in France, a waterborne outbreak of microsporidia occurred with approximately 200 cases of disease. The species identified was *E. bienewsi* (Sparfel *et al.*, 1997). While fecal contamination of the drinking water was never detected, contamination from a nearby lake was suspected. Typical symptoms of infection include chronic diarrhea, dehydration and significant weight loss (>10% body weight). Other symptoms include keratitis, conjunctivitis, hepatitis, peritonitis, myositis, central nervous system infection and renal disease. Treatments are

available for certain species of microsporidia, however, some species remain resistant to therapy (Cotte *et al.*, 1999).

Microsporidia spores have been shown to be stable in the environment and remain infective for days to weeks outside their hosts (Shadduck and Polley 1978; Waller 1979; Shadduck, 1989). Because of their small size (1–5 µm) they may be difficult to remove using conventional filtration techniques and there is a concern that these organisms may have an increased resistance to chlorine disinfection similar to *Cryptosporidium*. Initial studies using cell culture suggest that the spores may be more susceptible to disinfection (Wolk *et al.*, 2000).

In the USA there are minimal data on the occurrence of human strains of microsporidia in surface waters. There are more than 1000 species of microsporidia, of which 13 are presently known to infect humans (e.g. *Encephalitozoon cuniculi*, *Enterocytozoon bienewsi*, *Encephalitozoon intestinalis* and *Encephalitozoon hellem*). There are two species of microsporidia associated with gastrointestinal disease in humans: *E. bienewsi* and *E. intestinalis* (Dowd *et al.*, 1998a). Dowd *et al.* (1998b) described a PCR method for detection and identification of the microsporidia (amplifying the small sub-unit ribosomal DNA of microsporidia). They found isolates in sewage, surface waters and ground waters. The strain that was most often detected was *Enterocytozoon bienewsi*, which is a cause of diarrhea and excreted from infected individuals into wastewater.

3.4 *Toxoplasma gondii*

Toxoplasma is considered a tissue protozoan of cats and other felines who become infected mainly from eating infected rodents or birds, or from feces of infected cats. The symptoms of infection include flu-like symptoms, with swollen glands in the neck, armpits or groin area and most people recover without treatment. The cat is known as the definitive host, which means it is the only mammal that is known to excrete oocysts.

Humans and other mammals acquire the disease through ingestion of the oocysts (or ingestion of contaminated meat which has

been undercooked and contains the tissue stage). In a healthy adult the illness is a febrile illness of varying severity. However, in the immunocompromised the infection may cause severe disease. The greatest risk is to the fetus from infection of the woman during pregnancy, which can lead to fetal infection, chronic chorioretinitis or death.

Two outbreaks of toxoplasmosis have been associated with contaminated surface water. In 1979, 600 USA soldiers attended a 3-week training course in a jungle in Panama. Within 2 weeks of their return to the USA, 39 out of 98 soldiers in one company came down with a febrile illness. Serological testing revealed 31 confirmed cases of acute toxoplasmosis. The outbreak was attributed to the ingestion of contaminated water while on maneuvers in the jungle (Benenson *et al.*, 1982).

In March 1995, the Capital Regional District of Victoria, British Columbia identified 110 cases of toxoplasmosis (Bowie *et al.*, 1997). The outbreak was attributed to a single drinking water source for the area. The number of newly identified cases of toxoplasmosis declined sharply after the drinking water reservoir suspected of contamination was shut down. An estimated 3000 people (1% of the population) may have been infected by municipal drinking water contaminated with toxoplasmosis. Detection methods for oocysts in water have not been developed, however, the presence of infectious *T. gondii* oocysts in cougars as well as domestic cats living in the affected area was confirmed using mouse bioassays and serological surveys (Aramini *et al.*, 1998; 1999, Isaac-Renton *et al.*, 1998).

4 ENTERIC VIRUSES

There are several hundred enteric viruses that are potentially important agents of waterborne disease. In the past, it was thought that viruses were very species specific for the most part. Thus human sewage and wastes were considered the only source of viruses in water. However, there is new concern about the ability of RNA viruses, in particular, to evolve quickly and potentially jump the species

barrier. Such is the case with hepatitis E virus found in pigs. There is limited information regarding the incidence of virus infections in the USA population as well as throughout the world. Bennett *et al.* (1987) have reported 20 million cases of enteric viral infections and 2010 deaths per year.

Contamination of groundwater with viruses is of great concern due to the resistant nature of the viral structure and the colloidal size (20 nm), which makes this group of microorganisms easily transported through soil systems. Viruses also survive up to months in groundwater and are more resistant to water disinfection than are the coliforms (Yates and Yates, 1988; Gerba and Rose, 1989). Studies in the USA have found viruses in 20–30% of the groundwater where coliforms were not predictive of viral contamination (Abbaszadegan *et al.*, 1993). New techniques using PCR (Table 12.3) have shown that there is much more contamination than previously recognized.

The most frequently identified viruses in water or associated with waterborne outbreaks are hepatitis, Norwalk, coxsackie, rotavirus and the echoviruses. The cultivatable enteroviruses, which include the poliovirus, echovirus and coxsackie virus, make up only a small percentage of what is in wastewater and the tried and true cell culture methods used, which elicit a cytopathic effect when the virus is present, have been shown to be insensitive to many viruses, which grow but do not destroy the cells. Thus, PCR is now being utilized in cell culture to detect the non-cytopathic

TABLE 12.3 Virus detection in groundwater in the USA by various methods (Abbaszadegan *et al.*, 1993)

<i>Virus</i>	<i>Method</i>	<i>Percentage of samples positive</i>
Culturable enteric viruses	Cell culture	6.8 (12/176)
Enteroviruses	RT-PCR	30
Hepatitis A virus	RT-PCR	7
Rotavirus	RT-PCR	13
Total viruses	RT-PCR	39.3 (53/135)

RT-PCR (reverse transcription, polymerase chain reaction) is nucleic acid amplification for detection of the internal components of the virus.

viruses. In addition, new viruses are constantly being identified, which are non-culturable causes of gastroenteritis. These human caliciviruses (of which Norwalk is one) will need to be investigated in water using RT-PCR (reverse-transcription, polymerase chain reaction) methods.

4.1 Coxsackie viruses

Diarrhea has been one of the risks associated with many of the enteric viruses, but the recognition is emerging that more serious chronic diseases are now being associated with viral infections and these risks need to be better defined. Studies have reported that coxsackie virus B is associated with myocarditis (Klingel *et al.*, 1992). This could be extremely significant given that 41% of all deaths in the elderly are associated with diseases of the heart. In recent studies, enteroviral RNA was detected in endomyocardial biopsies in 32% of the patients with dilated cardiomyopathy and 33% of patients with clinical myocarditis (Kiode *et al.*, 1992). In addition, there is emerging evidence that coxsackie virus B is also associated with insulin dependent diabetes (IDD) and this infection may contribute to an increase of 0.0079% of IDD (Wagenknecht *et al.*, 1991).

Coxsackie virus B can be cultivated in cell culture and do cause a destruction of the cells, the cytopathic effect (Payment, 1997). They are commonly isolated from sewage, surface waters and groundwaters. Dahling *et al.* (1989) reported enterovirus concentrations in the discharges from several poorly operated activated sludge and trickling filter plants that were higher than 100,000 plaque-forming units per liter. At one of the plants, the discharge outlet was within one km of a drinking water treatment plant. On average, 95% of the enteroviruses detected were coxsackie virus type B5. Other viruses, like reoviruses and polioviruses, were detected but were found at lower levels. Krikelis *et al.* (1985, 1986) reported that almost 30% of the isolates from raw sewage from central Athens were coxsackie virus types B2, B4 and B5 with an estimated coxsackie virus concentration of 35.8 to 172.8

cytopathic units (CPU) per liter. Approximately 23% of the enteroviruses detected were of the echovirus group.

Between 1970 and 1979, four laboratories in different areas of the German Democratic Republic analyzed 1908 surface water samples from 30 sites for the presence of enteric viruses. Coxsackie virus (particularly types B3 and B4) was isolated every year during the sampling period (Walter *et al.*, 1982). Treated sewage discharges into the river were not disinfected. Lucena *et al.* (1985) collected samples from the Liobregat River and the Besos River and found an average coxsackie virus type B concentration of 0.107 and 0.60 most probable number cytopathic units, respectively.

Payment *et al.* (1985) sampled seven drinking water treatment plants twice a month for 12 months and detected coxsackie viruses in 7% (11 of 155) of the finished water samples. Coxsackie virus types B3 and B5 were also detected in well water during an outbreak in Texas in 1980 with an average concentration of 0.31 PFU/liter detected (Hejkal *et al.*, 1982). Compared to other viruses, including those in the enterovirus group, coxsackie viruses are commonly isolated from sewage and surface waters. Their role as a transmitter of waterborne disease is significant.

The cell culture methods for isolation and identification are cumbersome and lengthy, taking several weeks. New methodologies using cell-culture integrated with RT-PCR (reverse transcription, polymerase chain reaction) will provide faster more efficient and specific identification (Reynolds *et al.*, 1996; Grabow *et al.*, 1999; Lewis *et al.*, 2000).

4.2 Hepatitis viruses

Hepatitis A virus (HAV) is the second most commonly reported infection in the USA and hepatitis A is the most commonly reported type of viral hepatitis (Melnick, 1995). The hepatitis A virus is considered to be endemic in most Latin American and Caribbean countries (Craun, 1996). The symptoms of hepatitis A include fever, nausea, anorexia and malaise, often with mild diarrhea. The liver cells are

ultimately infected causing cytologic damage, necrosis and inflammation of the liver. Illness usually lasts from 1 to 2 weeks but may last several months. While HAV is an old disease, more recently outbreaks have occurred as a result of contaminated produce (Hutin *et al.*, 1999). Thus the control of HAV in human wastewaters and their elimination from waters used for irrigation of crops is a worldwide concern.

Hepatitis E virus (HEV) is an enteric RNA virus and causes jaundice and clinical symptoms very much like HAV. HEV has caused devastating waterborne disease outbreaks, particularly in tropical and subtropical countries with inadequate sanitation (Aggarwal and Naik, 1997; Balayan, 1997). In Kanpur, India in 1991, there were 79 000 cases of HEV due to sewage contamination of the drinking water. The earliest confirmed outbreaks occurred in the 1950s in India (Bradley, 1992).

Children are often asymptomatic and the mortality rate is between 0.1 and 4% (Grabow *et al.*, 1994). In pregnant women in their third trimester, the mortality rate can exceed 20% (Hurst *et al.*, 1996). There has been speculation that HEV is endemic in various parts of the world and subclinical cases may be contributing to the spread of the disease.

HEV is found in wild and domestic animals. Studies using genetic sequencing have found that human and pig HEV are very similar genetically in the USA as well as in Nepal (Meng *et al.*, 1997; Tsarev *et al.*, 1999). Thus zoonotic transmission seems very possible (Mushahwar *et al.*, 1998; Smith, 2001).

Although the cultivation of HEV has been reported (Smith, 2001), RT-PCR is the method of choice in water and has been used to detect HEV RNA in sewage (Jothikumar *et al.*, 1993; Pina *et al.*, 1998).

4.3 Norwalk-like caliciviruses viruses

The Norwalk-like caliciviruses (NLV) are enteric viruses (also known as small round structured viruses) and have been the cause of numerous food and waterborne outbreaks, associated with fecally-contaminated shellfish (Lipp and Rose, 1997), drinking water (Kaplan

et al., 1982; Lawson *et al.*, 1991; LeGuyader *et al.*, 1994) and recreational water (Kappus *et al.*, 1982). A diverse group of RNA viruses, they are a common cause of gastroenteritis, with diarrhea and/or vomiting, fever and respiratory symptoms lasting approximately 2 days. They were first identified with use of the electron microscope and include Norwalk virus, Snow Mountain agent, astroviruses and caliciviruses. There are extensive data showing these viruses are a major cause of shellfish-associated disease and may be the most significant group of viruses causing adult gastroenteritis (Lipp and Rose, 1997). Symptoms include vomiting, diarrhea, fever and, in some cases, respiratory illness. The viruses are heat stable and more resistant to chlorine, which is used to disinfect wastewaters, than the bacteria.

It has been suggested that these viruses are much more prevalent than previously thought. Reported in sewage at 10^7 RNA-containing particles per liter (Lodder *et al.*, 1999), a variety of genotypes could be detected, including those specific to outbreaks in the Netherlands.

4.4 Rotavirus

Rotavirus was first identified in 1973 and is believed to be responsible for the deaths of 4–5 million persons annually world-wide (Bern *et al.*, 1992). It is the major cause of viral gastroenteritis throughout the world and several waterborne outbreaks have been documented. In the USA, it is estimated that over one million cases of severe diarrhea in the 1–4 years age group are caused by rotaviruses annually, with up to 150 deaths (Ho *et al.*, 1988). In Third World countries it has been estimated that over 125 million cases of gastroenteritis due to rotavirus occur in children every year, with 18 million of these cases classified as moderately severe to severe. The very young, elderly, transplant patients and immunocompromised persons appear to be at greater risk of severe disease and mortality from rotavirus. The failure of a vaccine trial for young children continues to highlight the need to understand and prevent rotavirus transmission.

All of the rotavirus outbreaks have been associated with direct fecal contamination of a water supply or suboptimal drinking water treatment. Rotaviruses have been detected in surface waters world-wide with average concentrations ranging from 0.66 to 29 per liter (Gerba *et al.*, 1996b). The highest concentrations have been reported in surface waters receiving untreated sewage discharges. Rotavirus has also been detected in drinking water wells in Mexico and in Colombia (Deetz *et al.*, 1984; Toranzos *et al.*, 1988). The potential source of contamination was septic tank discharge into the groundwater.

While cell culture methods are available for detection of rotaviruses, they are cumbersome. These methods do not necessarily distinguish between human and animal rotaviruses. Throughout the animal kingdom there are many animal rotaviruses and natural reassortants arising from different species may appear including in humans (Vonsover *et al.*, 1993). Thus, RT-PCR is being used to detect and differentiate animal from human rotaviruses in water (Gratacap-Cavallier *et al.*, 2000).

5 FINAL COMMENTS

No doubt this chapter on emerging waterborne pathogens will need to be rewritten in another few years as new microorganisms are discovered or rediscovered and added to the list of concern. The detection of the bacterium *Tropheryma whippelii*, for example in sewage, using PCR (Maiwald *et al.*, 1998) makes one wonder if the faeco-oral and waterborne transmission of Whipple's disease (an intestinal lipodystrophy disorder) is likely; this needs to be fully investigated. The recent terrorist act associated with the Klingerman virus, which causes dysentery with a high mortality, means that genetically engineered enteric organisms may now enter the waste-stream and the water cycle. Only through continued monitoring studies and the application of new technologies can one address the emerging microorganisms that we are likely to see in this next century in the water environment.

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Health effects of water consumption and water quality

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The subject of microbiological safety of water will very soon become an international priority as travel across national boundaries and the sheer numbers of human citizens increase in the decade ahead, placing Promethean demands on water resources.

(Ford and Colwell, 1996)

1 INTRODUCTION

It was only at the beginning of the 19th century with the advent of water filtration, wastewater disposal, disinfection of drinking water with chlorine, pasteurization of milk and food, and by a general enhancement of hygiene, that waterborne pathogenic microorganisms and their diseases were finally controlled to an acceptable level in most countries. Waterborne diseases should not be seen as an independent part of the infectious disease cycle, but as a vehicle for their transmission. While we assume that a significant proportion of these gastrointestinal illnesses may be waterborne, we have no data to estimate the proportion of the overall burden of disease they represent (Figure 13.1). In a holistic approach, a reduction of waterborne disease should also result in a measurable reduction of the overall rate of gastrointestinal illnesses (Mara and Cairncross, 1989). This is the result of limiting person-to-person transmission and also the result of reducing the risk of the contamination of foods by contaminated water or individuals that have been infected through drinking water (Figure 13.2).

At the onset of the third millennium, what do we know of the health effects of drinking water and their impact on our so-called

modern societies? There is a pleiad of reports on the impact of waterborne diseases in countries world-wide, revealing thousands of outbreaks due to bacterial, viral, and parasitic microorganisms associated with the consumption of untreated or improperly treated drinking water (WHO, 1993, 1996; Ford and Colwell, 1996; Fewtrell and Bartram, 2001). These reports emphasize the fact that access to water and water quality are in direct relationship to life expectancy and child mortality, as countries with poor access to drinking water have the highest morbidity rates in children under 5 years of age.

2 WATERBORNE OUTBREAKS

Numerous enteric pathogens have been involved in waterborne outbreaks and recognized as such. Others are just emerging as being implicated in diseases transmitted by the water route (LeChevallier *et al.*, 1999a,b). Urbanization, ageing of water treatment plants, increasing numbers of immunocompromised individuals and ageing populations are potential causes for an increased risk of waterborne infectious diseases (Ford and Colwell, 1996; Fewtrell and Bartram, 2001). As we will see later, the endemic level of disease due to

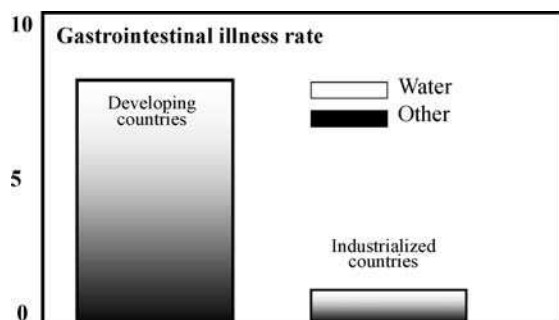


Fig. 13.1 The rate of gastrointestinal illnesses in developing countries is much less than in industrialized countries, but we still know very little about the proportion that can be assigned to water compared to other routes of transmission.

drinking-water consumption has also been shown to be quite significant and it contributes to maintaining these pathogens in the affected populations. The dilemma of balancing microbial and cancer health risks is also a difficult one to resolve, but it should not result in a reduction of treatment efficiency. The low risk level for cancer is insignificant when compared to the risk of waterborne infectious disease in absence of adequate water treatment (Payment and Hartemann, 1998; Fewtrell and Bartram, 2001).

The microorganisms implicated are described elsewhere in this book as well as in many recent publications on clinical and environmental microbiology (Murray *et al.*,

1995; Hurst *et al.*, 2001). Waterborne diseases are usually described in terms of outbreak reporting in the various countries and the USA has produced most of the available data through the decades-long effort of Gunther Craun (Craun, 1984, 1986, 1990) and a continuous effort to collect data (Herwaldt *et al.*, 1991, 1992). In other countries, data gathering is often performed very poorly because of lack of resources to identify the water-related events as well as the lack of centralized data gathering official authorities (Andersson and Bohan, 2001). Several methods for the detection and investigation of waterborne outbreaks have been described, but are still not widely used (Craun, 1990; Andersson and Bohan, 2001) as resources and funds are critically lacking, even in industrialized countries (Fewtrell and Bartram, 2001). An enormous effort is needed to educate all on the importance of water in the dissemination of disease. All levels of society, from consumer to politicians, must be educated to the necessity of improving water quality (Ford and Colwell, 1996) as a major step in improving the quality of life and health.

3 THE LESSONS OF THE RECENT DECADES

In all countries, a steady decline in waterborne disease was evidenced by the virtual elimination of cholera and the reduction of waterborne outbreaks to very low levels in most countries.

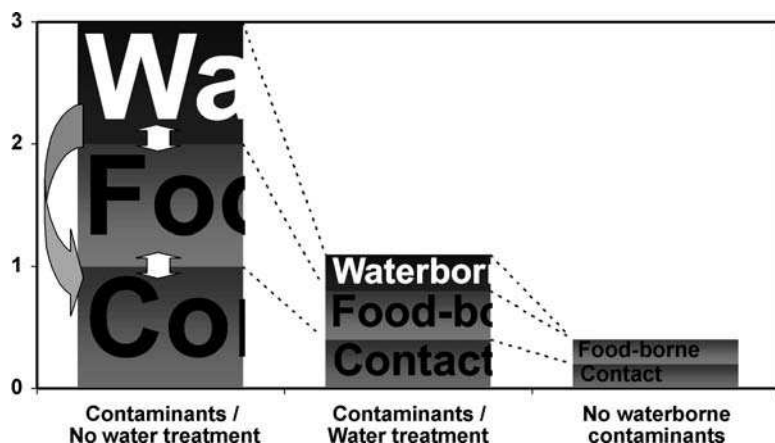


Fig. 13.2 Interrelations between waterborne disease and other routes of infection: reducing the role of water as a vehicle of transmission will significantly reduce the level of infectious disease in the community.

Most bacterial waterborne pathogens have been eliminated by the simple use of chlorine disinfection. There are still reports of bacterial disease outbreaks, mainly attributable to untreated water (non-disinfected groundwater), failure of disinfection when it is used as the sole barrier against microorganisms and finally re-contamination of the water in the distribution system. Major outbreaks such as the Walkerton outbreak of *Escherichia coli* in Canada (Anonymous (2000)) act as a reminder that the barrier must be maintained at all times.

However, we are finding strains of *Vibrio cholerae* that are more resistant to disinfection, *Legionella* has been found in water heaters and the *Mycobacterium avium* complex is now on the list of potential pathogens (LeChevallier *et al.*, 1999b; AWWA, 1999).

Since the 1950s, with the development of methods to detect and identify viruses, many outbreaks of waterborne gastrointestinal illness that would otherwise have been classified as of non-bacterial origin were attributed to enteric viruses. Hepatitis A and E, Norwalk, small round structured viruses, astroviruses, caliciviruses and many others are now well-known names in the water industry (LeChevallier *et al.*, 1999a; AWWA, 1999).

Parasites are also being identified as pathogens of importance, even in industrialized countries (Ortega, 1993; Chiodini, 1994; LeChevallier *et al.*, 1999a; AWWA 1999). The recognition that giardiasis was waterborne, with numerous outbreaks in the USA (Craun, 1986) and several outbreaks of cryptosporidiosis in England (Badenoch, 1990a,b) and in many other countries (Payment and Hunter, 2001) have now shifted the focus to these parasites, which are extremely resistant to the water disinfection process. Dozens of outbreaks of cryptosporidiosis have now been reported world-wide, but most are small compared to the explosive outbreak experienced in Milwaukee (USA) in the spring of 1993 (Edwards, 1993; Mackenzie *et al.*, 1994). Following what appears to be a failure in treatment, water that met the United States Environmental Protection Agency water quality guidelines may have caused gastrointestinal

illnesses in over 400 000 people or one-third of the population of this city over a period of one month. Most of these illnesses were cryptosporidiosis, but many were probably of viral origin and the number of illnesses reported may have been overestimated by 100-fold (Hunter and Syed, 2001). The most surprising aspect of this event is that cryptosporidiosis was probably occurring even before it was detected following a report from an inquisitive pharmacist (Morris *et al.*, 1998). This suggests that unless an effort is made to identify waterborne diseases they will remain undetected, buried in the endemic level of illness in the population (Payment and Hunter, 2001).

4 WATERBORNE DISEASE IS NOT ONLY DIARRHOEA

While the most often reported disease associated with drinking water remains gastroenteritis, this is probably due to the very apparent nature of the symptoms and the fact that the attack rates for these infections can often reach over 50% of the exposed population. Even infectious disease specialists often forget that enteric microorganisms are associated with a wide range of symptoms and diseases (Table 13.1). Protozoan parasites, such as amoebae, can cause severe liver or brain infections and contact lens wearers are warned of the dangers of eye infections. Bacteria can cause pneumonia (*Legionella*) and some are suggesting the possibility that *Helicobacter pylori*, which has been associated with gastric ulcers, could be transmitted by the water route. Among the 72 known enteroviruses, many can induce illnesses other than gastrointestinal, such as hepatitis (viral type A or E), poliomyelitis, viral meningitis, enteroviral carditis, epidemic myalgia, diabetes, ocular diseases. Some viruses can also induce abortions, stillbirth and fetal abnormalities (Payment, 1993a,b). Who can say that a death due to myocardial failure is not the result of a waterborne coxsackie virus infection months or years earlier?

TABLE 13.1 Enteric viruses and associated diseases or symptoms

<i>Symptoms/diseases</i>	<i>Viruses</i>
Minor malaise, asymptomatic diseases	All
Paralysis, poliomyelitis and polio-like	Poliovirus, coxsackie virus, enteroviruses 70–71
Aseptic meningitis and mild paresis	Poliovirus, coxsackie virus (B1–6, A7, A9)
Eye diseases, conjunctivitis	Echoviruses, coxsackie virus (B2, B4, A24), enterovirus 70,
Exanthems (Hand-foot-and-mouth)	Enterovirus 71, coxsackie viruses A, echoviruses
Cardiac diseases (myocarditis, chronic)	Coxsackie virus A and B, echoviruses
Pleurodynia and epidemic myalgia	Coxsackie virus A and B
Meningoencephalitis	Enterovirus 70–71
Respiratory illnesses	Coxsackie virus, echoviruses, rotaviruses
Viral gastroenteritis	Enteroviruses, calicivirus, astrovirus, rotavirus, coronavirus, adenovirus 40–41, etc.
Neonatal diseases	Coxsackie viruses A and B
Viral hepatitis	Hepatitis A and E viruses, coxsackie viruses A and B
Diabetes	Coxsackie virus B4
Post-viral fatigue syndrome	Enteroviruses, coxsackie viruses B

5 ENDEMIC WATERBORNE DISEASE IN INDUSTRIALIZED COUNTRIES: EARLY EPIDEMIOLOGICAL STUDIES

The fact that waterborne disease is still a major problem in developing countries does not need to be reaffirmed. In this chapter, we will thus emphasize the current state of knowledge of waterborne diseases in countries that have historically been assuming that their populations were not exposed to risks from their drinking water, as this was assumed to be adequately treated.

In the absence of evident acute health effects (i.e. epidemics or outbreaks), epidemiological studies have been centred on long-term effects of potentially carcinogenic chemicals (Crump and Guess, 1982). Birth defects or abortions have also been associated with the consumption of tap water (Swan *et al.*, 1992; Wyndham *et al.*, 1992), even though much of this effect might have been due to recall bias during retrospective and matched-control studies (Hertz-Picciotto *et al.*, 1992). Such epidemiological studies are extremely difficult to assess and are often never confirmed because there are too many confounding variables in studies that attempt to analyse lifetime exposure to various factors.

Batik *et al.* (1979), using hepatitis A virus cases as an indicator, could not establish a

correlation with water quality and could not find a correlation between current indicators and the risk of waterborne outbreaks (Batik *et al.*, 1983).

In France, the group of Collin *et al.* (1981) prospectively studied the gastrointestinal illnesses associated with the consumption of tap water using reports from physicians, pharmacists and teachers. Their results were based on more than 200 distribution systems of treated or untreated water and they reported five epidemics (more than 1000 cases) associated with poor quality water. This study is typical of most studies which relied on the detection of epidemics to assess the level of water quality: they do not address the endemic level of gastrointestinal illnesses which may be due to low level contamination of the water. The same group, in a prospective follow-up study on 48 villages for 64 weeks, has evaluated the gastrointestinal illnesses associated with untreated ground water and found a relationship between faecal streptococci and acute gastrointestinal disease (Ferley *et al.*, 1986; Zmirou *et al.*, 1987). Faecal coliforms did not appear to be related to acute disease. Total coliforms and total bacteria had no independent contributions to disease but, even in the absence of all measured indicator germs, one-fourth of the cases were still observed.

In Israel, Fattal *et al.* (1988) addressed the health effects of both drinking water and aerosols. Their studies on kibbutz water quality and morbidity were performed in an area with relatively high endemicity of gastrointestinal disease and did not show a relationship between health effects and total or faecal coliforms. This study was, however, based only on morbidity reported to physicians, data that are considered to represent only 1% of the actual cases in a population.

In Windhoek (Namibia), Isaäckson and Sayed (1988) conducted a similar study over several years on thousands of individuals served by recycled wastewater as well as normal drinking water. They did not observe an increased risk of reported acute gastrointestinal illness associated with the consumption of recycled waters. The populations compared had higher incidence rates than the one observed in North America and they were subjected to a high endemicity level due to other causes, thus masking low levels of illnesses.

6 THE CANADIAN EPIDEMIOLOGICAL STUDIES

Two major epidemiological studies were conducted in Canada in 1988–89 and in 1993–94. The results of these studies suggest that a very high proportion of gastrointestinal illnesses could still be attributable to tap water consumption even when water met the current water quality guidelines (Payment *et al.*, 1991a,b, 1997).

6.1 First study

The first study was carried out from September 1988 to June 1989. It was a randomized intervention trial carried out on 299 randomly selected eligible households which were supplied with domestic water filters (reverse osmosis(RO)) that eliminated microbial and most chemical contaminants from their tap water, and on 307 randomly selected households which were left with their usual tap

water without treatment. The gastrointestinal symptomatology was evaluated by means of a family health diary maintained prospectively by all study families. The estimated annual incidence of gastrointestinal illness was 0.76 among tap-water drinkers as compared with 0.50 among RO-filtered-water drinkers ($P < 0.01$). Because participants in the RO-filter group were still exposed to tap water (i.e. about 40% of their water intake was tap water), it was estimated that about 50% of the illnesses were probably tap water-related and preventable. The remaining illnesses were probably attributable to the other possible causes such as endemic infectious illnesses, food-related infections, allergies, etc.

The fact that the participants in the study, while randomized into two groups, were not blinded to the type of water they consumed was considered a confounding factor that could have affected the results. However, the rate of illness increased with the amount of water consumed (i.e. a dose–response effect was demonstrable), suggesting that the effect observed was probably not due to bias.

Attempts were made to correlate microbiological data obtained on water samples from the water distribution system. There was no correlation between the number of episodes and total or faecal coliforms, chlorine or heterotrophic bacteria in the tap water. There was an association between the duration of illness and heterotrophic plate count (HPC) bacteria at 20°C (Payment *et al.*, 1993). However, due to the large number of statistical analyses performed this could have been a spurious result due to chance only.

In the RO-filtered water study group, there was a significant correlation between the rate of illness and the HPC bacteria growing at 35°C on R2A medium, but no relationship with the amount of water consumed (Payment *et al.*, 1991b). The rate of water-related illness increased with distance from the plant and bacterial regrowth was suggested as an explanation. Studies of the virulence of bacteria isolated from the tap water revealed that a small fraction of these bacteria could be considered as potential pathogens (Payment

et al., 1994a) and it was suggested that their multiplication to high numbers could be a health risk (Payment, 1995).

Attempts were also made to determine the aetiology of the observed illnesses. Sera had been collected on four occasions from volunteers and they were tested for antibodies to various pathogens. There was no indication by serology for water-related infections by enteroviruses, hepatitis A virus and rotavirus (Payment, 1991) or Norwalk virus infections (Payment *et al.*, 1994).

6.2 Second study

The second study (Payment, 1997) was more complex because it attempted not only to re-evaluate the level of waterborne illness, but also to identify the source of the pathogens responsible for them. It was conducted from September 1993 to December 1994 and it compared the levels of gastrointestinal illness in four randomly selected groups of 250 families, which were served water from one of the following sources:

- Tap water (normal tap water)
- Tap water with a valve on the cold water line (to examine the effect of home plumbing)
- Plant effluent water as it left the plant and bottled (i.e. not influenced by the distribution system)
- Plant effluent water further treated and bottled (to remove any contaminants).

The site was selected for the high level of microbial contamination of the river water that it treats and for the quality of operation of the water treatment plant. Raw water entering the plant was contaminated with parasites, viruses and bacteria at levels found throughout the world in faecally-contaminated waters. The water treatment plant produced water that met or exceeded current Canadian and US regulations for drinking-water quality. The distribution system was in compliance for coliforms but residual chlorine was not detectable at all times in all parts of the distribution system.

The rates of highly credible gastrointestinal illnesses (HCGI) were within the expected

range for this population at 0.66 episodes/person-year for all subjects and 0.84 for children 2 to 12 years old. The rate of illness was highest in autumn–winter and lowest in summer. Overall, there were more illnesses among tap-water consumers than among subjects in the purified (bottled) water group, suggesting a potential adverse effect originating from the plant or the distribution system. Children were consistently more affected than adults and up to 40% of their gastrointestinal illnesses were attributable to water. The rates of gastrointestinal illness among consumers of water obtained directly at the treatment plant were similar to the rate of illness among consumers of purified water. Two periods of increased tap-water-attributable illnesses were observed respectively in November 1993 and in March 1994.

Subjects in the two bottled water groups (i.e. purified water and water obtained directly at the water treatment plant) still consumed about one-third of their drinking water as tap water. They were thus exposed to some tap water and its contaminants; as a result the risks due to tap water are underestimated.

Consumers of water from a continuously running tap had a higher rate of illness than any other group during most of the observation period. This was completely unexpected, since the continuously running tap was thought to have been able to minimize the effects of regrowth in home plumbing. Although there are several unsubstantiated theories as to the cause of this effect, it remains unexplained.

The data collected during these two epidemiological studies suggest that there are measurable gastrointestinal health effects associated with tap water meeting current standards and that contaminants originating from the water treatment plant or the distribution system could be the source of these illnesses. Short-term turbidity breakthrough from individual filters at the water treatment plant might explain the observed health effects. Potential follow-up research should further examine the relationship of turbidity breakthrough and should investigate the role of the continuously running tap in the occurrence of gastrointestinal illness.

7 OTHER EPIDEMIOLOGICAL STUDIES

Following the example of these studies in Canada, the Australians (Hellard *et al.*, 2001) and the Americans (Hayward, 2000; Colford *et al.*, 2002) have initiated similar prospective intervention studies.

The Australian study reported no health effects due to unfiltered clean surface waters. The design was modelled on the Canadian studies but participants were blinded to the type of device they had and one group received a sham device. The results suggested that when source water quality is excellent, drinking water that is disinfected does not constitute a significant health risk (Hellard *et al.*, 2001).

Initial results of the first of several ongoing American studies appear to confirm the Canadian studies. This American study also models the Canadian studies and also made efforts to insure that participants were effectively blinded to the type of device installed in their house. This study is closer to the conditions described in the Canadian studies: the treatment plant is a well-operated complete conventional treatment plant that treats water from a river that is significantly polluted (Colford *et al.*, 2002). Preliminary results are very similar to the results of the Canadian studies.

8 ENDEMIC WATERBORNE DISEASES: POSSIBLE CAUSES

In the Canadian studies, it has not been possible to assign the observed effects to a single cause. While in the first study the source of illnesses was a complete mystery, the authors suggested three explanations: low level or sporadic breakthrough of pathogens at the water treatment plant, intrusions in the distribution system (repairs, breaks, etc.) and finally bacterial regrowth in the mains or in the household plumbing. While no single explanation has been given, many answers have been provided in recent years.

8.1 Health significance of bacterial regrowth

Bacterial regrowth is common in water and has been observed even in distilled water. In water distribution systems, the heterotrophic plate count can occasionally be elevated and there have been concerns that this flora could still contain opportunistic pathogens. Data from epidemiological studies involving reverse-osmosis units suggested that there was a correlation between gastrointestinal illnesses and heterotrophic plate counts at 35°C (Payment *et al.*, 1991b). However, a few outliers in the dataset drove the correlation and the study would have to be repeated in order to confirm the relationship. Furthermore, this observation could be limited to such water purification devices in which a rubber bladder is used to accumulate the purified water. Bacteria or fungi growing on rubber could have been selected for virulence factors increasing their potential for initiating disease.

Heterotrophic bacteria isolates from tap water were subjected to a number of tests to determine the presence of virulence factors (Payment *et al.*, 1994). Several isolates contained numerous virulence factors, but these bacteria were found in numbers generally too low to significantly affect human health. For most bacterial pathogens, even frank pathogens, thousands to millions of bacteria are needed for an infection to occur. This is a rare occurrence in drinking water. It was concluded that these bacteria did not present a significant health risk (Payment, 1995; Edberg *et al.*, 1996; WHO, 2002).

The health significance of pigmented bacteria found in drinking water was reviewed in a recent paper (Rusin *et al.*, 1997). The species investigated included the opportunistic pathogens *Flavobacterium*, *Pseudomonas*, *Corynebacterium*, *Nocardia*, *Mycobacterium*, *Erwinia*, *Enterobacter*, *Serratia* and *Micrococcus*. The authors also concluded that, although a number of these bacteria are opportunistic human pathogens, the available data show them to be infrequent causes of disease.

8.2 Health significance of turbidity

Subsequent to the investigation of the main Milwaukee outbreak mentioned earlier, Morris *et al.* (1996) carried out an analysis of hospital records and water turbidity readings over a period of 16 months before the recognized outbreak. They found that attendance of children with gastrointestinal illness at hospital emergency departments showed a strong correlation with rises and falls in turbidity, but did not describe any specific time lag relationships.

Schwartz *et al.* (1997) presented evidence that rises in rates of gastroenteritis in children were preceded by rises in turbidity of the treated drinking water supply. The authors looked for correlations between the number of children attending the emergency department at the Children's Hospital of Philadelphia (CHOP) for gastroenteritis with the turbidity of treated water in the preceding 14 days. The 1.2 million people of Philadelphia receive their drinking water through three water treatment plants supplied by rivers considered to be 'highly contaminated'. The finished waters tested in all these studies met the commonly accepted definition of safe drinking water in terms of turbidity and chlorine treatment and this raised serious questions about their wholesomeness (Franco, 1997).

These findings complement each other. The Canadian studies concluded that a fraction of gastrointestinal illness attributable to drinking water arises from microbiological events in the distribution system but did not discount the treatment plant as a source of pathogens. Schwartz *et al.* (1997) and Morris *et al.* (1996) suggested that variations in rates of illness were due to changes in the numbers of pathogens (carried in or on small suspended particles) coming through the distribution system from the treatment plant. A water treatment plant is a continuously working system and it changes minute by minute according to the demands. Rapid sand filters are not homogeneous and their overall performance varies considerably. The distribution system is subjected to numerous challenges

and the intrusion of pathogens is now a highly debated subject.

9 COSTS TO SOCIETY

The societal cost of the so-called 'mild gastrointestinal illnesses' is several orders of magnitude higher than the costs associated with acute hospitalized cases (Payment, 1997). In the USA, it was estimated that the annual cost to society of gastrointestinal infectious illnesses is \$9500 million dollars (1985 US dollars) for cases with no consultation by physician, \$2750 million dollars for those with consultations, and only \$760 million dollars for those requiring hospitalization (Garthright *et al.*, 1988, Roberts and Foegeding, 1991). These estimates do not even address the deaths associated by these illnesses, particularly in children and older adults. Furthermore, this mode of contamination acts as a constant reinfection source for the population, maintaining at an endemic level illnesses that could be further reduced.

From the data collected during the Canadian studies, the economic costs of endemic waterborne diseases were calculated based on reported symptom and behaviour rates between unexposed and tap-water exposed groups (Payment, 1997). The rate of gastrointestinal illness in the unexposed group was taken as the baseline, then the excess illness in the tap water groups was 50% in the first study and 20% in the second study – a difference attributable to tap water. As even the control groups drank some tap water, the difference attributable to tap water is likely to be higher than the observed figures at a range of 25–50%. These estimates were then combined with published figures for the cost of gastrointestinal infectious diseases in the USA (Garthright *et al.*, 1988; Roberts and Foegeding, 1991). Assuming a population of 300 million individuals, the estimate of the cost of waterborne illness ranges from US\$269 to \$806 million for medical costs and US\$40 to \$107 million for absences from work. Such figures can only underscore the enormous economic costs of endemic gastrointestinal illnesses, even in societies believed to be relatively free of it.

10 THE FUTURE: RISK ASSESSMENT

Risk assessment in microbiology was almost non-existent until recently (Eisenberg *et al.*, 1996). It has now become an important part of the preparation of guidelines for water quality in all countries (Fewtrell and Bartram, 2001). In their current rule-making process, the USA have incorporated an important place to microbial risk assessment (Regli *et al.*, 1991; Sobsey *et al.*, 1993) and all agencies world-wide are now debating the balance to be maintained between chemical risks and microbial risks (Fewtrell and Bartram, 2001). It is now recognized that the chemical risk reduction objective should never result in a decrease in water disinfection (Guerra de Macedo 1993; WHO 1996; Fewtrell and Bartram, 2001). This could have catastrophic implications as was experienced during the 1991 cholera outbreak in South America. It is important to remember that differentiation between developed and developing countries is an arbitrary process when drinking water safety is considered. Problems and solutions vary from country to country, a fact that must be taken into account when formulating solutions.

The spread of cholera in Peru during the last outbreak may have been attributable to the fact that disinfection may have been abandoned or reduced in response to the report from the US EPA on risk assessment of the carcinogenic potential of disinfection byproducts (Anderson, 1991).

Water is becoming a commodity whose value will grow in the coming years. Wars have been fought for water and will probably continue to occur if the current use of water resources continues. The global decline in water quality due to global climatic changes can easily upset the balance in many parts of the world and the first observable effect is often a microbiological degradation that can affect the health of millions of individuals.

The subject of microbiological safety of water will very soon become an international priority as travel across national boundaries and the sheer numbers of human citizens increase in the decade ahead, placing Promethean

demands on water resources. (Ford and Colwell, 1996, preface by Rita R. Colwell).

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Drinking-water standards for the developing world

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1 INTRODUCTION

In 2000 it was estimated that in excess of 1 billion persons still lacked access to improved drinking water sources (WHO and UNICEF, 2000). Most of these are the rural and urban poor living in developing countries who remain disadvantaged in terms of access to basic services and whose health and well-being is put at risk from poor and contaminated environments. Drinking-water supply and quality both have important impacts upon health and socio-economic development and therefore remain important components of the poverty cycle.

Approaches to the management of drinking water and of drinking-water quality vary widely between countries in response to factors such as environmental and climatic conditions, technical capacity, level of economic development and cultural and societal norms and practices.

In countries where access to improved water sources or supply is low, the role of legislation, regulation and standards is likely to be best oriented towards encouraging extension of access to higher levels of service, ensuring effective use of existing infrastructure and ensuring that the minimum safeguards are in place to prevent outbreaks of waterborne disease. Improving access to improved drinking-water supply that present limited risks to health and fulfil basic rights to a 'safe' and

'adequate' water supply implies making best use of limited resources towards these priorities. Legislation, regulation and standards may promote this directly and may themselves be more or less cost-effective in terms of the resources demanded for their implementation and the impact of their application.

In terms of drinking-water quality, most regulations and standards are dominated by health concerns. The World Health Organization publishes *Guidelines for Drinking-water Quality* to assist countries at all levels of economic development in establishing national approaches to drinking water that are effective in the protection of public health (WHO, 2003a).

In December 2000 formal recognition was given to the human right to water through the publication of a 'General Comment' on this theme. The General Comment creates an obligation upon governments to support *progressive realisation* of universal access to sufficient, safe, acceptable, physically accessible and affordable water and to *respect, protect* and *fulfil* the right to water (WHO, 2003b).

1.1 Water supply and quality – differences between developed and developing countries

Challenges and problems in drinking water supply are generally more complex and daunting in poorer countries which are at the same time typically less well equipped (both financially and with technical and policy expertise) to deal with the complexity.

The opinions expressed in this article are those of the author and do not necessarily represent the policies or views of WHO or University of Loughborough.

Industrially developed nations typically have established drinking-water supply infrastructure for all except some rural areas and small community supplies. The principal technical challenges therefore relate to efficient management of that infrastructure, including rehabilitation and replacement strategies that require significant recurrent investment and to the pursuit of improved water quality. The problems of fecal contamination are largely under control, although outbreaks of disease may occasionally be recognized and it is likely that other outbreaks go unrecognized. Increasing attention will be dedicated to chemical hazards in drinking water, some of which, while attracting significant public concern, may be of limited health significance.

Less industrially developed nations have incomplete water supply infrastructures such that even middle-income countries may have significant populations with limited access to piped drinking water supply at higher levels of service. As a result, significant proportions of their populations utilize water supplies of poor quality and reliability (Table 14.1). Institutions responsible for both supply and oversight of water supply may be less 'mature' than elsewhere and supply of trained personnel of all types may be inadequate. As a result the efficiency of the water supply sector may be low.

In less developed countries a complex juggling act is therefore required to make the best use of limited resources for public health benefit. This requires trade-offs between extension of supply, basic water quality (largely management of fecal contamination) and other water quality issues.

In both more and less industrially developed nations ongoing changes, especially in public and private sector roles, are creating new challenges to established structures and procedures, with the expectation of improved future efficiency.

In rural and poor urban areas of developing countries direct community management of water supply is common and indeed is seen as desirable by many in the sector. Development of effective legislative and regulatory approaches to support effective community management is especially complex.

TABLE 14.1 Examples of countries with significant populations without access to improved sources of drinking water in 2000

<i>Country</i>	<i>Percentage without safe drinking water</i>		
	<i>Urban</i>	<i>Rural</i>	<i>Total</i>
Afghanistan	81	89	87
Central African Republic	20	54	40
Chad	69	74	73
Papua New Guinea	12	68	28
Madagascar	15	69	53
Angola	66	60	62
Mozambique	14	57	40
Sierra Leone	77	69	72
Uganda	28	54	50
Vietnam	19	50	44
Mali	26	39	35
Myanmar	12	40	32
Lao PDR	41	–	10
Nigeria	19	61	43
Iraq	4	52	15
Nepal	15	20	19
Zambia	12	52	36
Malawi	5	56	43
Sri Lanka	9	20	17
Benin	26	45	37
Sudan	14	31	25

Source: based on data from WHO/UNICEF/WSSCC, 1996.

In relation to water quality, microbiological hazards figure highly in rational analysis of health priorities in both developed and developing countries. This is a result of their contribution to both outbreaks of disease, which may be large in scale, and of their contribution to background rates of disease. Diarrheal diseases related to sanitation, water supply and personal hygiene account for 1.73 million deaths annually and are ranked third in importance in disability adjustment life years in the global burden of disease (World Health Report, 2002). Other diseases are related to poor water supply, sanitation and hygiene such as trachoma, schistosomiasis, ascariasis, trichuriasis, hookworm disease, guinea worm disease, malaria and Japanese encephalitis and contribute to an additional burden of disease. In some circumstances chemical hazards such as arsenic, fluoride, lead and nitrate in drinking water may also be of significance.

1.2 Regulating water supply and quality in developing countries

Overall, the management of drinking water supply and quality for public health benefit is more complex in developing countries than within industrially developed nations.

In many industrially developed nations, water quality standards may be developed and, indeed, implemented in isolation from considerations related to the need to extend supplies to unserved populations. This greatly simplifies the development of legislation and of regulations. Legislation in the established market economies, oriented principally toward the regulation of established water supplies administered by recognizable entities, may therefore have little relevance in less industrially developed nations.

While a significant proportion of the world's population does not have access to piped water supply, and many more do not have access to reliable, safe, piped drinking-water supply, most legislation and regulation is oriented towards this means of provision.

Ensuring an adequate piped water supply is dependent upon proper activity throughout a chain that begins with securing adequate volumes of water and promoting maintenance of high quality water resources from which drinking-water can be abstracted. It also depends on management of drinking-water sources, on effective treatment and distribution and on proper household care. The latter means that basic elements such as household water-handling and domestic plumbing may have a significant impact upon effective water supply. The overall approach to regulation may be similar in more industrially developed nations than less-developed nations where water supply coverage is incomplete and more flexible legislative approaches are required to enable trade-offs to be made.

Legislation and regulation supportive of improved non-piped water sources is relatively scarce, although many principles are similar. The tendency to regulate piped but not other forms of water supply may be one of the factors that draws additional resources to the further improvement of already improved supplies

rather than towards the extension of improvements to the unserved.

Overall, for many less developed countries, the role of regulation might be argued to be to support the best use of available resources in the provision of incrementally improved service standards to the population as a whole. Available experience and guidance derived from regulatory bodies in the more developed nations may be inappropriate for direct application in developing nations. As a result, countries may be reluctant to adopt foreign practice in legislation because of its limited relevance to their national situation (Caponera, 1992). This is especially valid in the case of drinking water supply.

In all cases, regulation tends to be more effective when it is driven by processes of negotiation and the inclusion of civil society within the debate about both standards and the structure of the water supply services. The use of a range of regulatory tools and instruments will also be commonly required in order for balanced decisions to be made based on priorities within the sector.

1.3 Urban areas of developing countries

While there is a common perception that urban populations receive high-quality water supplies at higher service levels, in reality there is great inequity in access to drinking water supplied on-plot (i.e. at least a yard level of service). In most developing countries a minority of wealthier households enjoy water supplies of high quality and service level, while the majority of the poor rely on communal piped and point water sources (Howard, 2002). Where piped water supplies are used, it is not uncommon to find that the urban poor experience greater unreliability in supply.

Not only are the inequities immense in terms of service level, the consequences for the urban poor of utilizing point sources of water is increased exposure to pathogens and therefore increased risk to health. In addition, there is an increased risk of re-contamination of the water during transport and storage. Furthermore, in most developing countries

the urban poor pay proportionately more for poorer water services than those in wealthier areas and are often more likely to be disconnected from utility supplies.

The improvement of water supplies in urban areas, while in some regards simpler than for rural areas, still retains fundamental problems. Sustained access and operation and maintenance is reliant on payment of utility water tariffs and therefore the costs of water become a significant factor in determining access and, in particular, whether charges meet the ability and willingness to pay by poor households and communities. Furthermore, the tariff structure and payment methods, with an emphasis on payment on bills presented periodically (typically monthly or quarterly), are often at odds with the income patterns of the urban poor where incomes may be highly unstable and abilities to save are limited (Howard *et al.*, 1999).

The regulation of the quality, reliability and cost of utility supplies that form a major component of the urban water sector can be effectively achieved utilizing simple and transparent approaches, however, a careful balance must be maintained between competing demands of affordability and access and concerns over quality (Ince and Howard, 1999). More significant problems are found in regulating other aspects of water supply, such as social provision and leakage control. In the case of social provision, there may remain significant conflicts between the demands for financial sustainability of the utility. Where leakage-control programs are implemented, sustaining rehabilitation and replacement strategies (which require significant expense and planning) may be difficult. Consultation with both users and water suppliers may be essential in establishing targets for water quality and other aspects of water supply, in order to ensure that required improvements are made.

1.4 Rural areas of developing countries

There remain particular problems in the development of water supplies in rural areas

where 80% of the presently unserved population live. Overall, coverage of the rural population with functioning water supplies remains low in many countries. The rural water supply sector outside the established market economies typically comprises diverse means of supply and with diverse types of infrastructure, many of which fail either permanently or temporarily soon after construction and a high percentage of the accumulated infrastructure out of commission at any given time.

In addition to the frequent functional failure of many small water supplies, water quality has often proved difficult to sustain at acceptable levels as sanitary protection measures deteriorate through a lack of maintenance. Such problems are not unique to developing countries. There is substantial evidence that small, effectively community-managed, supplies in Western Europe and North America fail to meet national and regional water-supply regulations. For instance, in the UK, 47% of small supplies gave at least one unsatisfactory result in a recent assessment (Fewtrell *et al.*, 1998).

Approaches to improving drinking water supply and quality have increasingly tended to emphasize the role of the community as water managers. However, as early as 1980, community participation was referred to as the 'mythology' of the International Water Supply and Sanitation Decade (the 1980s) (Feachem, 1980). In a review of the results of the massive investments in the Decade, it was noted that while community management of water supplies can be effective, it is unlikely to offer more than a medium-term solution to overcome current inadequacies in government and the private sector in meeting rural water demands (Carter *et al.*, 1993).

The special problems of the rural sector are recognized by those specializing in water law (Caponera, 1992), both because of the specific relationship of rural populations with water resources and in terms of water supply and sanitation and their relationship to health and disease transmission. In a book otherwise couched in relatively specific legal terms, Caponara (op. cit.) indicates that:

In such areas legislation by itself is not sufficient to abate the occurrence. It is necessary to educate the users of water points in proper operation and maintenance practices. This often requires a minimum training background in public hygiene and health, in operation and maintenance of the facilities, and in other basic aspects relating to water management. Water legislation should contain the necessary provisions, as well as provisions tending to promote the organization of water users' associations.

For rural water supply it is especially important that ownership and responsibilities related to operation and maintenance of systems constructed by one agency and operated by another (often local) one (such as a water users' association) are clear and unambiguous. Thus, Appleton (1995) refers to the need for legislative changes to provide the authority and autonomy for decentralized operations to function successfully. Attempts have been made to promote and support community operation and maintenance through monitoring, and evidence is accumulating that this can be effective. These approaches suggest that far more resources and time need to be spent in developing capacity with communities to be able to recognize the need for preventative maintenance and identify early signs of major faults and thus be able to rectify these.

1.5 Water supply and quality in international trade

The issues arising from the process of 'globalization' are complex. Of particular relevance to drinking-water supply is international trade in drinking-water supply services. Within the past two decades, the role of international companies in managing water supply to 'foreign' cities has emerged as a significant issue. Unlike other areas of trade, such as food, internationally recognized points of reference for specification or performance evaluation are often lacking. Exceptions include the WHO *Guidelines for Drinking-water Quality*.

There remain unresolved key issues about international trade in water services. While the

increased use of such approaches is advocated by many donor agencies and governments, public perception may be far less favorable. Important considerations concern quality standards to be met, their independent surveillance and the pursuit of public health priorities in a privately managed system. The latter may imply significant relaxation in quality standards to enable effective extension of supply into under- or unserved areas. The acceptability of such relaxations may be low despite their evident logical need.

2 DRINKING-WATER LEGISLATION, REGULATION AND STANDARDS

Regulation in general is primarily designed to protect the public interest against undue exploitation or the provision of substandard goods and services that either infringe 'rights' or raise the risks to health. However, an important subsidiary objective is to ensure that there is a transparent system of management where responsibilities and liability are clearly defined, an approach that could be said to be providing a 'free and fair playing field'. Thus effective regulation is often as much in the best interest of water suppliers (or at least those of a reputable nature) as it is for the public at large.

The principal concerns of drinking-water legislation and regulation are which functions must be exercised and what administrative arrangements are required in order that they are exercised effectively and efficiently. While the functions vary relatively little between countries, the exact means of their implementation and, in particular, administrative arrangements, vary widely and these require adaptation to take account of wide differences in basic circumstances.

2.1 Functions

The principal functions that must be exercised include:

- those relating to the management of drinking-water sources and the resources from which drinking water is abstracted – these

typically include volume management (conservation of resources), licensing of abstractions and pollution control (effectively water resource quality management)

- measures relating to the provision of supply services to the population – these often vary in response to administrative arrangements. Essential are those measures that define minimum service standards (continuity, quality of water, tariff controls, leakage targets) and which seek to pursue the objective of increasing population access and service level to a reasonable minimum target such as a single tap within the house or yard
- drinking-water quality standards and their implementation through actions by both the supply agency and others, including information sharing and public reporting
- the regulation of ancillary services, which impact directly upon supply or quality (including for example, those which may impact upon both water security through leakage and safety through proper practice against contamination)
- public health functions (including ongoing public health oversight and response to occurrences endangering public health)
- licensing of materials and chemicals used in the production and distribution of drinking water (such as pipes and treatment chemicals) and of household devices such as taps.

These different functions can be translated into a set of regulations and standards that would typically be incorporated into a regulatory framework. The areas of regulation and their principal features and means of monitoring are outlined in Table 14.2.

2.2 Basic law

It is generally preferred that all provisions concerning water are contained in a single law and so the basic law will typically be far wider than drinking-water supply alone. The basic law that relates to water may take diverse forms (constitution, act, code etc.) and would normally include a statement of purpose (policy) which would refer to the intention to protect human health and welfare and define

priorities among different uses. Aspects such as public health protection and consumer rights may be dealt with in other parts of the basic law but be applicable to water supply.

The basic law generally refers to the jurisdictions, responsibilities and authority of specified competent agencies in relation to water and to their relationships to one another and, in particular, to the development of subordinate legislation. This should include water users (consumers), those constructing and administering water supply systems, those regulating different aspects of water and having responsibilities in related domains and agencies involved in data and information management and sector planning. Kandiah (1995) refers to the legal and policy frameworks that assign responsibility to agencies to implement and coordinate their programs as one of the key factors affecting the suboptimal performance of all types of water schemes.

In general, while the principles of drinking-water regulation are contained within the basic law, the actual tools of regulation are more effective when established as statutory instruments. Provisions for the development of such instruments would typically be included within the basic law, which would outline the aspects of drinking water where such statutory instruments can be set up, who the responsible minister will be, the frequency of revision and what procedures and expert advice is required for such revisions. In the case of drinking-water quality for example, standards may be set by the Minister of Health, with:

1. A review of standards and new information undertaken on an annual basis by an expert review body appointed by the minister composed of relevant sector stakeholders (including water suppliers).
2. The review to take into account new evidence of health impact.
3. The review to take into account impacts of new standards on investment and other requirements.

In Ghana, a review of standards undertaken in light of a private sector lease operation recommended that the regulatory instruments

TABLE 14.2 Regulations and standards for drinking-water supply and quality

<i>Area of regulation</i>	<i>Typical major features of regulations and standards</i>	<i>Means of verification/monitoring</i>
Water sources and resources	<ul style="list-style-type: none"> • Abstraction licensing • Source protection norms and practice • Polluter-pays principle • Monitoring of land-use, control of encroachment 	<ul style="list-style-type: none"> • Permits and bulk flow meters; annual visits and periodic assessment • Sanitary inspection • Pollution monitoring and enforcement
Continuity of supply	<ul style="list-style-type: none"> • Set and enforce targets for continuity of supply • Investment requirements for improvement and maintenance of reliability 	<ul style="list-style-type: none"> • Sanitary inspection/community interview • Investment plans and expenditure; evidence of rehabilitation strategies
Drinking-water quality	<ul style="list-style-type: none"> • Place obligation continuously to supply safe water upon the supply agency • Require supply agency to establish a water safety plan and exercise due care • Define what is 'safe' in terms of acute or long-term exposure • Establishment of limit values, objectives and guidelines • Development of statutory instruments, negotiation and enforcement procedures • Independent verification 	<ul style="list-style-type: none"> • Routine surveillance and monitoring of water quality • Audit of monitoring programs • Best practice approaches and international evidence • Standards set through consultative process • Instruments developed and linked to primary law • Surveillance/audit
Monitoring requirements	<ul style="list-style-type: none"> • Divide monitoring obligations among a number of agencies or sectors (public sector oversight of water resource management; public health authorities; drinking water suppliers; including local government) • Require information-sharing among these agencies • Define the extent of monitoring required of each agency and define the conditions under which monitoring should be undertaken • Define the technical requirements against which monitoring results would be compared 	<ul style="list-style-type: none"> • Allocation of responsibility in primary law and through statutory instruments • Legal instruments • Limits of responsibility and definition of duty of care • Standards for water quality, process control, source protection and distribution management

(continued on next page)

TABLE 14.2 *(continued)*

<i>Area of regulation</i>	<i>Typical major features of regulations and standards</i>	<i>Means of verification/monitoring</i>
Treatment requirements	<ul style="list-style-type: none"> • Require supply agency to treat water to achieve the required standards (may be implicit in the requirement of the basic law to supply safe water) • Require that newly constructed or rehabilitated supply systems reach minimum treatment capabilities (these may be formulated in terms of performance criteria or processes required, generally in relation to the characteristics both of the source water and of the supplied population) • Require the upgrading of existing systems to reach the standards required of newly constructed or rehabilitated systems • Define technical details of construction or operating practice 	<ul style="list-style-type: none"> • Minimum treatment requirements for source types and source quality; process control standards; drinking-water quality standards. Verification through surveillance • Minimum treatment requirements for source types and source quality; process control standards; and drinking-water standards. Verification through surveillance • Rehabilitation and upgrading strategies in place with firm investment plans • Standard operating procedures, verification through audit
Contingency plans	<ul style="list-style-type: none"> • Oblige government agencies to establish contingency plans • Define the conditions under which contingency planning is required, and the types of contingency to be planned for • Describe the basic components of an 'adequate' plan 	<ul style="list-style-type: none"> • Monitoring through regional organization, government watchdog bodies and non-government organizations to ensure plans developed that meet agreed criteria of the basic plan for supply quality under extreme conditions

be set and related to the new water act. The responsibility for developing and applying the regulatory instruments would fall under a public utilities regulatory commission and where consultation in the derivation of standards would include water suppliers, the health sector and local government.

The use of statutory instruments is important for a number of reasons. The setting of water quality standards is a dynamic process (Ince and Howard, 1999). However, basic or primary law tends to be relatively inflexible since its change is generally a long process, usually involving an Act of Parliament. Provisions in primary law cannot therefore reflect the rapidity of change in the available knowledge about, for instance, the health impacts of drinking water, changing socio-economic conditions and society's demand for protection. For similar reasons, penalties to be applied in enforcement actions should not be incorporated into basic law. In many developing countries where such penalties were traditionally incorporated into the basic law, changing economic conditions have reduced fines (originally used as a deterrent) to insignificant amounts which water suppliers themselves may be willing to pay, rather than invest in the upgrading of water supplies.

2.3 Subsidiary legislation

Among the details that form part of the subsidiary legislation on water are standards of various kinds. For example, there may be standards on minimum treatment requirements linked to source type and quality, and for drinking-water quality. However, standard-setting should not be restricted to drinking-water quality, but should also extend to water resource protection and water supply service more generally. Indeed, it should extend beyond the simple assessment of quality and include the adequacy of structures and systems, such as the definition of safe facilities and practices, minimum standards specification, and minimum standard operating systems.

There may also be schedules or annexes to statutory instruments, consisting of even more detailed subject matter. In some countries,

drinking water service standards are contained in a sanitary code or code of good practice. It is worth noting that such standards are most likely to receive support in their implementation if consultation with the affected entities (e.g. supply agencies and professional bodies) has occurred.

One of the most important criticisms of current approaches to standards and norms relates to microbiological quality of water and, in particular, the over-reliance on a set of end-product indicator standards that have well-recognized limitations. Such criticisms apply equally to utility and alternative supplies. This has led to a change in approach within the third edition of the WHO *Guidelines for Drinking-Water Quality* to a quality assurance approach based on Water Safety Plans (Davison *et al.*, 2002; WHO, 2003). The Water Safety Plan approach emphasizes control of water safety from the catchment to the consumer through the use of the multiple barrier principle. Water Safety Plans focus on effective process control linked to simple monitoring activities and periodic internal audit. These are accompanied by health-based water quality targets set at national level and periodic independent surveillance/audits.

In the case of community-managed supplies, enforcement of regulations that would typically govern aspects related to ongoing operation and maintenance, such as water quality and continuity, is unlikely to be a workable mechanism to achieve improvements in water supply. Given inherent problems in applying regulatory procedures in circumstances where there is no differentiation between consumers and suppliers, different approaches are required to support water supply improvement. One approach is to develop water supply surveillance programs to act as a supportive role in promoting improvements. Critical to this process is the routine use of sanitary inspection as a measure of operation and maintenance and the support to communities to address such weaknesses. There remains some potential for regulation during the construction phase of such supplies, provided an organization or contractor external to the community is

undertaking the work through a variety of mechanisms.

2.4 Institutional aspects

There are diverse approaches to drinking water management – varying between the extremes of centralized state ownership and decentralized private ownership and management. The ongoing trends towards private sector participation and towards community management are attempts to move towards a more demand-driven approach, which may offer better possibilities for regulation and greater public demand for protection.

Other significant trends have been towards *decentralization* (the principle of devolving decision-making to the lowest appropriate level); and the recognition at governmental level that effective policy development and management requires *multisectoral cooperation*. There is general acceptance of the principle that central government should retain responsibility for public health and maintaining quality standards (see for example, Appleton, 1995; IRC, 1995).

As the use of water for drinking is effectively universal, it is perceived as a high priority for effective independent oversight. The oversight functions will vary according to legal, administrative and technical arrangements. There is extreme diversity in approaches to this independent oversight between exercise of direct control by a governmental surveillance agency and auditing approaches requiring relatively limited public sector intervention. However, most 'models' for organization of monitoring, including that proposed by WHO (1976, 1985), envisage two complementary roles: service quality control by the supply agency and an independent surveillance function.

WHO has indicated that 'the water supplier and surveillance agency should be separate and independently controlled' (WHO, 1976; WHO, 1997). Whilst such a model is both reasonable and often viable in urban areas worldwide, a rigid division between surveillance and quality control is rarely applicable for small community supplies in developing countries (Helmer, 1987), except where privatization has

occurred to a definable responsible agency or where a small community is supplied by a formal utility as sometimes occurs near major cities or where several rural communities are supplied through a single centralized system.

Within the urban sector, the viability of strict division of roles depends, in part, on the water supply management arrangements. Where water supplies are provided by a utility, surveillance by local public health departments under the overall guidance of a national health body, would generally be the preferred model (Howard, 2002). Where local government provides water, this may become more problematic, although experience in Uganda indicates that provided separate departments within local government undertake the two functions, effective separation of roles can still be achieved (Howard *et al.*, 1999).

In both urban and rural settings, the use of surveillance to promote better hygiene during the collection, transport and storage of water is important. This is particularly true in any setting where the water supply is primarily through communal water supplies.

In all cases duplication of functions may be wasteful of limited resources and procedures for information sharing. Appropriate auditing and quality control are increasingly accepted to enable the independent surveillance agency to focus energies on high-risk areas and population groups.

It is not inevitable that the surveillance function should fall under the direct supervision of the health sector (WHO, 1985, 1996). For example, in the United Kingdom regulation of water quality in utility supplies falls under the remit of the Department of Environment. In many cases, actual implementation of monitoring and surveillance functions is more effective when de-centralized to local levels. This approach is reflected in UK practice in relation to private water supplies and similar models are found in Yemen.

Given the broad range of issues in the water sector that require some form of regulatory practice, there are a number of institutions with an interest in ensuring that this is implemented (see Table 14.3).

TABLE 14.3 Functions and examples of potentially interested agencies concerned with drinking-water supply

Abstraction licensing	Water resources management agency, agricultural authorities, abstraction licensing body, river basin authorities, navigation boards, hydroelectricity generators, water supply agencies
Water resource quality	Water resources management agency, pollution control boards, river basin authority, health, water supply agencies, local governments
Drinking water supply service quality	Water supply oversight body, consumer protection bodies, water supply agencies, health, local government
Ancillary services (e.g. plumbing)	Professional associations, trade associations, water supply agencies, local government (through environmental health)
Public health protection	Health, local government, water supply agencies, consumer protection bodies
Licensing of materials and chemicals	National licensing authority/certification agency; health, water supply agencies

NB: representatives of civil societies including NGOs and professional associations operate at each level

Aspects of the administrative arrangements that may be put in place to ensure that the functions outlined above are exercised effectively may relate, for example, to:

- Arrangements for multisectoral participation and integrated management
- Provisions for participation in management and decision making (including community participation and evolving public–private sector roles)
- Provisions related to the sharing of information
- Approaches to and responsibility for monitoring and assessment in general and independent oversight in particular.

3 STANDARD SETTING AND IMPLEMENTATION

Conflicts may exist between standards and ensuring optimal health effects (Holland, 1991). Resources are finite and, in practice, a supply agency may have to choose, for example, between investing in fulfilling water quality standards despite relatively good quality, and extension of coverage. The choice between these two may have a profound influence on the resulting impact on the health of the population (Esrey *et al.*, 1985, 1991). There has been a change in approach in recent decades among public health professionals to water supply provision from ‘all or nothing’ to ‘something for all’ (IRC, 1995).

While regulation often focuses upon direct drinking-water *quality* standards, its scope includes regulation concerning:

- construction standards
- plumbing standards
- service quality standards (cost, continuity)
- water quality standards
- coverage (e.g. when an authority has responsibility for a defined geographical area or population)
- licensing of materials and chemicals used in drinking water supply.

Standards need to be developed to take account of national circumstances, such as health burden, current water qualities (source and distributed), availability of resources (infrastructure, finance, willingness to pay and expertise) to meet standards and implications for upgrading and analytical capacity. Key to the setting of standards is therefore the evaluation of health priorities related to water, of the current ability to meet existing or anticipated standards (reflecting technical capacity, water source qualities, abilities to pay) and the available resources to implement improvement required.

Standards for different components are highly inter-related and the establishment of a standard in one aspect, for instance water quality, has direct impacts on the achievability of other standards. The tariff set, for instance, may determine the ability to meet water quality standards and may directly influence

the ability of the supplier to extend services or to be able to reduce unreliability. In the same way, the establishment of a water quality standard will typically have direct influences on the cost of producing water and therefore the tariff that will be applied. Experience suggests that integrated approaches to standard derivations are required.

3.1 Scientific basis for microbiological standards

The scientific basis for establishing standards for microbial safety are the same in all settings, although the final targets established may vary. Overall approaches to establishing effective standards for the microbial safety of water, taking into account the range of exposure routes (drinking water, recreational water and re-use of wastewater), has been recently reviewed: a 'harmonized' framework for future application to developing standards has been proposed (Fewtrell and Bartram, 2001).

Typically, microbiological hazards are of greatest concern and constitute the causative agents of disease (pathogens) that may be transmitted through consumption of contaminated water. They may be protozoa, viruses or bacteria. There are a wide variety of microbiological agents that may be transmitted through drinking water (Table 14.4).

For all of the agents listed in Table 14.4, a single exposure may be significant for public health and, for example, sufficient microbes to cause disease may be consumed in a single glass of apparently innocuous water. Furthermore, this 'infectious dose' may be extremely small – potentially as little as a single viable cyst or virion for some protozoa and viruses.

Analytical methods are available for some of these, but unavailable for many others. Where analytical methods exist they may or may not be quantitative and problems may be encountered in their application to water as an analytical medium.

The principle that no pathogens should be present in drinking water has therefore become widely accepted. This principle, alongside the lack of analytical methods and the fact that almost all of the pathogens of interest are

primarily derived from human excreta (feces), led to the development of the concept of 'fecal indicators'. The value of quantitative estimates of fecal indicator bacteria in water was recognized early in the history of sanitary microbiology. The definition of microbiological quality now used in most regulatory and non-regulatory monitoring world-wide is therefore based on the premise that fecal contamination of drinking water is unsafe and that assessing the presence of indicators of fecal contamination provides an indication of the safety of drinking water.

The use of fecal indicators has made a significant contribution to the protection of human health over a sustained period and it continues to be valuable and popular. It is, nevertheless, imperfect both in conception and in application. Its principal limitations relate to well-recognized shortcomings of the principal available indicators and their ability to meet the basic criteria presented in Box 1. The majority of currently used indicators are bacteria and this has important implications regarding their use and the information they provide in relation to non-bacterial pathogens.

BOX 1: *Characteristics of an ideal microbial indicator*

- Should be present in wastewater and contaminated water when pathogens are present
- Should be present when there is a risk of contamination by pathogens
- Should be present in greater numbers than pathogens
- Should not multiply in environmental conditions under which pathogens cannot multiply
- Should correlate with the degree of fecal contamination
- Survival time in unfavorable conditions should exceed that of pathogens
- Should be more resistant to disinfection and other stresses than pathogens
- Should present no health risk
- Should be easy to enumerate and identify by simple methods
- Should have stable characteristics and give consistent reactions in analysis

(Source: WHO Teaching Pack on Drinking-Water Quality.)

TABLE 14.4 Pathogenic microorganisms that may be found in water

<i>Pathogen</i>	<i>Health significance</i>	<i>Persistence in water supply</i>	<i>Resistance to chlorine</i>	<i>Relative infective dose</i>	<i>Important animal reservoir</i>
Bacteria					
<i>Campylobacter jejuni</i> , <i>C. coli</i>	High	Moderate	Low	Moderate	Yes
Pathogenic <i>E. coli</i>	High	Moderate	Low	High	Yes
<i>Salmonella typhi</i>	High	Moderate	Low	High	No
Other salmonellae	High	Long	Low	High	Yes
<i>Shigella</i> spp.	High	Short	Low	Moderate	No
<i>Vibrio cholerae</i>	High	Short	Low	Moderate	No
<i>Yersina enterocolitica</i>	High	Long	Low	High (?)	No
<i>Pseudomonas aeruginosa</i>	Moderate	May multiply	Moderate	High (?)	No
<i>Aeromonas</i> spp.	Moderate	May multiply	Low	High (?)	No
Viruses					
Adenoviruses	High	?	Moderate	Low	No
Enteroviruses	High	Long	Moderate	Low	No
Hepatitis A	High	?	Moderate	Low	No
Enterically transmitted non-A, non-B hepatitis viruses, hepatitis E	High	?	?	Low	No
Norwalk virus					
Rotavirus	High	?	?	Low	No
Small round viruses	High	?	?	Moderate	No (?)
	Moderate	?	?	Low (?)	No
Protozoa					
<i>Entamoeba histolytica</i>	High	Moderate	High	Low	No
<i>Giardia intestinalis</i>	High	Moderate	High	Low	Yes
<i>Cryptosporidium parvum</i>	High	Long	High	Low	Yes
Helminths					
<i>Dracunculus medinensis</i>	High	Moderate	Moderate	Low	Yes

Source: WHO (1993) Guidelines for Drinking-water Quality Volume 1.

There has been increasing evidence of presence of pathogens in water meeting guidelines and standards for the principal fecal indicator bacteria, *E. coli*. In some cases this has led to outbreaks of infectious disease related to the consumption of contaminated water. As a result, greater attention has been placed on alternative methods of defining microbiological quality, including testing for pathogens, identification of alternative indicators for non-bacterial pathogens (e.g. phage as an indicator of potential viral contamination) and other risk assessment approaches, such as sanitary inspection. For instance, in the case of *Cryptosporidium*, monitoring efforts may be better focused on ensuring adequate sanitary completion of groundwater sources and control of turbidity during treatment.

It is increasingly recognized that zero risk is unachievable and its pursuit inhibits the application of risk–benefit approaches. As noted above, it is unlikely that any approach will provide the degree of certainty required to define a ‘safe’ water supply in all circumstances, while current indicators can be taken as indicative of gross fecal pollution. This then leads to change in the way we view microbiological contamination, which is that no water is ‘safe’, but rather may be low, intermediate or high risk – a ‘tolerable disease burden’.

The idea of tolerable disease burden (TDB) is especially important in drinking-water microbiological safety because of the variable health outcomes from different exposures. Most studies have addressed diarrheal diseases which, although they account for a significant global burden of disease, are often self-limiting and, in some circumstances, of limited concern to the general public. They may be contrasted with more severe health outcomes. Different pathogens may produce diseases of varying public health importance, e.g. enteric hepatitis viruses (hepatitis), *Vibrio cholerae* (cholera), *Cryptosporidium* spp. (cryptosporidiosis), or *Salmonella typhi* (typhoid). Some workers have suggested specific figures for a TDB for diarrhea, but this has not as yet appeared in a form translatable to other disease outcomes (such as disability-adjusted life years or DALYs) and proposals for TDBs for more

severe health outcomes have not yet been made.

Increasing attention is therefore being paid to the characteristics of individual pathogens and the concept of TDB as the future basis for definition of microbiological safety (quality) for drinking water. Substantial ongoing work will see this theme develop rapidly in future years. However, translating TDB into practical descriptions of quality is complex. The distribution of microbes in water (individual or clustered on particulates) may have an effect on the probability of infection or of developing disease. The relationship between exposure, infection and disease, especially at low dose exposures, remains poorly understood for most pathogens and inter-relates with external factors such as immunity and the form of exposure (e.g. from food, aerosols etc.) (Fewtrell and Bartram, 2001; Prüss *et al.*, 2001).

Approaches to ensuring microbial safety are increasingly based on a preventive management approach, taking account of factors from source through to consumer. The quality of drinking water may be controlled through a combination of protection of water sources, control of treatment processes and management of the distribution and handling of water.

The WHO Guidelines for drinking-water quality outlines a framework for safe drinking water that comprises five key components (Davison *et al.*, 2002; WHO, 2003):

1. Water quality targets based on critical evaluation of health concerns
2. System assessment to determine whether the water supply chain (from source through treatment to the point of consumption) as a whole can deliver water of a quality that meets the above targets
3. Monitoring of the control measures in the supply chain which are of particular importance in securing drinking-water safety
4. Management plans documenting the system assessment and monitoring; and describing actions to be taken for normal and incident conditions; including upgrade and improvement and documentation and communication

5. A system of independent surveillance that verifies that the above are operating properly.

Component 1 would normally be undertaken by the national authority responsible for public health. Components 2–4 collectively comprise a water safety plan and are the responsibility of the water supplier, whether large or small. Component 5 may be the responsibility of several independent agencies with local and national capacity to undertake auditing and/or compliance testing.

3.2 Scientific basis for chemical standards

There are a few chemicals in drinking water that have been clearly associated with large-scale health effects. These include arsenic and fluoride. In order to direct resources to best effect, identifying those chemicals that may be of public health concern nationally or locally, is important. Guidance on practical approaches employing diverse types of information in addition to water quality analysis is available in documents accompanying the WHO *Guidelines for Drinking-Water Quality*.

The principal focus for controlling chemical safety of drinking water is through establishing numerical standards for substances against which the results of water quality analysis can be compared. An exception to this is where chemicals are either added during treatment or may leach from pipe and pipe joining materials. These would typically be regulated through product specifications.

In order to establish standards, it is necessary to identify the concentration of a substance in drinking water that will represent 'safety'. Two approaches are adopted depending on whether the chemical has a 'threshold effect' or not. A threshold effect is where a concentration can be defined below which no adverse effect on health is seen over a lifetime consumption. Non-threshold chemicals are largely genotoxic carcinogens where any level of exposure represents a theoretical risk.

For threshold chemicals, the concentration at which no adverse effect is detected forms the basis of the standards. Corrections are made for exposure from other sources, such as food. In practice, data from real human populations are rarely available and reliance must be placed on the studies undertaken over short periods on animals. An uncertainty factor is therefore often incorporated into the Guideline Value.

To establish a numerical standard for non-threshold chemicals, a level of 'negligible' risk is defined (often 10^{-5} or 10^{-6} lifetime risk of cancer). In order to accommodate variation in the human population, the concentration derived is in fact an upper bound estimate.

A number of practical considerations are also taken into account when establishing guidelines for standards for chemical substances in water. These include preventing standards that:

1. are below the limit of detection
2. are not technically or economically achievable; and/or
3. would counteract other aspects of water safety (such as discouraging disinfection).

A comprehensive explanation of the derivation of Guidelines Values is presented in the WHO *Guidelines for Drinking-Water Quality* (2003).

3.3 Social/cultural, economic and environmental factors

While risk management is often seen as a highly rational process, it should be recognized that the scientific basis for many of its elements is actually often weak. Furthermore, scientific assessment will not support effective risk management if it fails to address the perceptions and priorities of the society concerned. One schematic explaining the process of risk management is presented in Fig. 14.1. This describes the process as circular showing the feedback between policy evaluation into revised (improved) policy. It places communication at the centre of the decision-making process that is designed to both ensure that information from each stage is communicated

Risk Management

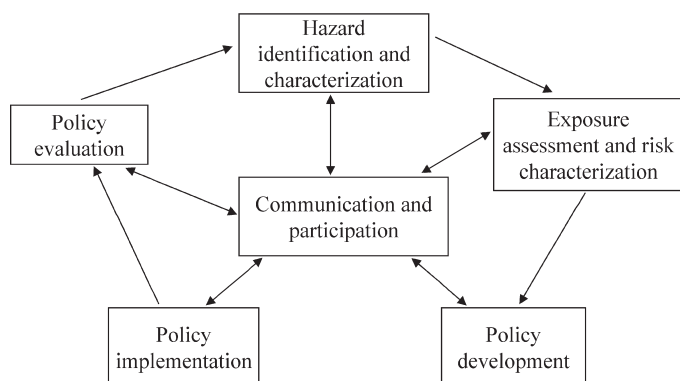


Fig. 14.1 The Risk Management Cycle (adapted from Chorus and Bartram, 1999 and Sobsey *et al.* 1993)

with stakeholders, and that the needs for communication are taken into account when undertaking each stage of the risk assessment and management process.

As well as the scientific aspects of standard development, adaptation is therefore required to take account of social, cultural and economic factors. In some circumstances, societal concerns will lead to the adoption of standards that are stricter than can be justified by simple health risk assessment alone.

A critical component in the development of standards and the balancing of numerous demands for improved water supply, is the systematic assessment of costs of meeting objectives and the implications that this will have for tariffs and access. This is further complicated by the very different forms of water supply commonly found in many developing countries. Unlike their wealthier counterparts in the developed world, a large proportion of supplies will not be expected to meet stringent water quality standards, primarily in the rural sector, while the more sophisticated urban piped water supplies should be able to achieve a quality of water similar to those in developed countries. This, therefore, raises difficult ethical and political problems regarding the application of standards and the dangers of the already significant inequity in the water sector being exacerbated. When standards are set, the process includes negotiation on the quality to

be achieved. Such negotiation already often occurs between regulators and suppliers. However, in many countries, this needs to be expanded to include more public consultation to take into account social demands for protection and public perception of risks.

3.4 Progressive implementation

In many developing countries, the need to develop and apply drinking-water standards has increased in importance as the nature of government roles in the sector has changed. The result, in part in response to public pressure effectively to 'control' private sector supply agencies, may be 'strict' standards. However, where water supply legislation is 'utopian' it will tend to promote the perfection of existing water supplies at perhaps great cost despite minimal health risk and at the expense of providing an improved, but perhaps imperfect, supply to (perhaps many) more individuals. In particular, the emphasis on meeting advanced water quality standards may be counter-productive in situations where costs of investment in new infrastructure to meet particular standards may result in reduced access by the poor to services that are yet to be developed or become too expensive.

The use of regulatory instruments, in particular the application of relaxations and exemptions, the development of interim standards and definition of priorities for control,

are therefore important if regulation is to be a positive force for change. Essential to this is the identification of priorities with regard to water supply and an emphasis placed on meeting key standards before attention is paid to less significant problems. Often this will lead to an emphasis on microbiological before chemical quality.

A relatively frequent need is to adopt less stringent 'interim standards', either to encourage progressive improvement or to ensure that limited financial and technical resources are not diverted from issues of high to low relative priority for public health.

This is common where the resolution of water quality problems may require significant upgrading of water infrastructure that will take an extended period of time or large-scale investment or enforcement of other legislation (For instance, prevention of encroachment). In such cases, an interim standard for a substance may be set at a higher level to allow improvements to be made progressively. Examples might include requiring immediate implementation of chlorination and thereafter progressively improved treatment where surface waters are supplied without treatment.

In addition to the delay in implementing standards fully or developing interim standards is the use of two specific tools – exemption and relaxation.

In many cases, a failure in water supply may not be easy to resolve in a short period of time without large-scale investment that will eventually be passed onto the consumer. In some cases, this may be supply and substance-specific, in which case an exemption can be applied. This would typically be time-constrained, linked to an agreed and defined program of action and an allowable deviation from the standard set. Such approaches would usually require a case to be made by the water supplier for an allowed deviation.

Relaxations would typically be much broader and cover all supplies, or particular types of water supply. They may in some cases also apply to particular parameters of more limited health concern. Relaxations would usually be phrased in terms of an acceptable

percentage of samples where analysis shows that a standard has been exceeded, often with an upper limit of allowed deviation also specified.

3.5 Enforcement

Standards are intended to ensure that the consumer enjoys a reliable supply of safe, potable water; not to shut down deficient systems (WHO, 1984; Wheeler, 1990; Bartram, 1996). However, the most frequently articulated reason for routine monitoring of water supplies is to detect failures in compliance with water supply standards and regulations to be employed to enforce improvement (daSilva and O'Kane, 1989; Ward, 1995). Thus Le Moigne (1996, p. 20) suggests that 'the best water management strategy would mean little without laws and regulations that can be enforced'; while Lloyd and Helmer (1991, p. 58) have gone so far as to claim that in rural areas 'it is only worth considering planning the development of surveillance programs if the political will to implement water legislation exists'.

Many water supply regulations are therefore designed to assess measures of water supply service quality (and especially water quality), to compare results with a standard and to take some form of action upon detection of failure. However, because 'reality' is so far removed from the ideal situation proposed in many water supply standards and legislation (Okun and Lauria, 1992), standards and legislation have lost credibility in many countries (Jensen, 1967). In order to optimize the benefits of their use it is therefore appropriate that explicit schemes for enforcement implementation are provided for and pursued (Jensen, 1967).

3.6 Source protection and minimum treatment requirements

The establishment of drinking-water quality standards alone is not sufficient to provide confidence in the water supply in achieving an adequate level of water quality throughout the system. In addition, best practice norms need to be established for source protection and treatment requirements. Within the emerging

paradigm for water safety promoted within the third edition of the WHO *Guidelines for Drinking-Water Quality* ensuring adequate source protection and treatment are given higher profile and are linked to specific control measures.

As noted in Section 2 above, source protection norms would normally be included within the statutory instruments linked to the primary law that governs water. Typically, such norms will make provisions for the requirements at several levels. These would usually include the basic sanitary completion measures designed to prevent direct contamination of the source, a broader area to protect the source from more widespread microbiological contamination, as well as broader measures protection for sources against chemical contamination and broader watershed management. In general, source protection norms have been better developed and enforced for groundwater than surface water, largely because the natural microbiological quality of groundwater is much higher, control of pollution is often simpler and an acceptance that all surface water should be treated prior to consumption. Such a simplistic approach is being increasingly challenged in recognition of the chemical hazards found in groundwater sources of drinking water that pose a significant risk to public health. The need for protection of surface water sources should also not be minimized, as ineffective source protection leads to increasing loss of resources available for drinking-water supply within reasonable levels of treatment costs.

Minimum treatment requirements are often specified in relation to specific sources. However, the basis of most treatment requirements is that the processes should be proven to be able effectively to remove contaminants and that a multiple barrier principle should be applied. This principle is designed to ensure that water will pass through several stages of treatment, thus reducing the risks of supply of untreated water. In some cases, failures in critical processes may lead to automated shut-down. The different treatment processes are not reviewed here, but it should be noted that increasing attention is being paid to 'older'

processes that are primarily biological and physical as these may represent more cost-effective approaches than chemical processes.

The final component in most treatment processes is disinfection and this may also be applied to groundwater. Disinfection serves two purposes: it provides the final stage of treatment and is designed to remove any remaining microorganisms that have survived the preceding processes; it may also provide partial protection against re-contamination within distribution. The latter is an important function as water quality failure post-production remains common within many water supplies, particularly those that are older or prone to discontinuity or high leakage. However, it is increasingly recognized that there are some pathogens for which commonly used disinfectants are not effective. Furthermore, relying on disinfectant residuals to control the presence of pathogens in drinking water during distribution is ineffective.

The disinfectant of choice worldwide remains chlorine, although alternatives such as ozone are attracting increasing attention. Disinfection with chlorine is commonly applied as a final treatment step for surface water. In this application a minimum 'contact time' is normally engineered into the treatment plant configuration. In the case of groundwater, the practice has, in general, been only to provide in-line terminal disinfection that has limited contact time and therefore may not inactivate microorganisms entering the distribution system. There is increasing evidence of pathogen survival greatly in excess of those previously thought which indicates that this approach may need to be re-considered.

4 CONCLUSION

The development of standards for drinking-water quality is an important component in water quality control. However, this needs to be considered in light of other needs and, in particular, the extension of supply to those not served. The establishment of standards needs to consider not just health evidence and technical ability, but also impacts on costs

and tariffs and social considerations. Routine monitoring and enforcement are essential if standards are to have meaning. In addition to standards governing water quality, complementary standards should address source protection and minimum treatment requirements in order to ensure adequate water quality.

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Control of pathogenic microorganisms in wastewater recycling and reuse in agriculture

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1 INTRODUCTION

As presented in detail, in Chapter 11, raw domestic wastewaters normally carry the full spectrum of pathogenic microorganisms – the causative agents of bacterial, virus and protozoan diseases endemic in the community and excreted by diseased and infected individuals. While recycling and reuse of wastewater for agriculture, industry and non-potable urban purposes can be a highly effective strategy for developing a sustainable water resource in water short areas, nutrient conservation and environmental protection, it is essential to understand the health risks involved and to develop appropriate strategies for the control of those risks. This chapter will concentrate on the control of pathogenic microorganisms from wastewater in agricultural reuse since this is the most widely practised form of reuse on a global basis. However, more and more water specialists, natural resource planners and economists see water as an economic good and, as time goes on, there will be an increased motivation to divert recycled wastewater from low income agriculture to areas where the added value of water is greater, such as industrial and non-potable urban uses including public parks, green belts, golf courses, football fields. As time goes on and water

shortages in arid areas increase, there will undoubtedly be an expansion of the reuse of purified wastewater for industrial and a wide variety of urban/non-potable purposes. There are some specific health guidelines and standards for such uses, but they will not be reviewed in this chapter.

The control measures for agricultural reuse include establishing and enforcing microbial guidelines for effluent quality, regulation of the types of crops to be irrigated, minimizing the potential for crop contamination by various irrigation techniques and the treatment of the wastewater to an *appropriate degree* so as to control potential health risks, both to the farmers and the consumers of crops, from pathogenic microorganisms in the wastewater stream. This chapter will deal with the above questions.

2 PATHOGENIC MICROORGANISMS CAN BE TRANSMITTED BY WASTEWATER IRRIGATION

Pathogenic microorganisms in the wastewater stream can be transmitted to healthy individuals and cause disease if improper regulation and control methods in wastewater irrigation are practised. In order for disease causing microorganisms or pathogens in the wastewater stream flowing from a community to infect a susceptible individual they must be

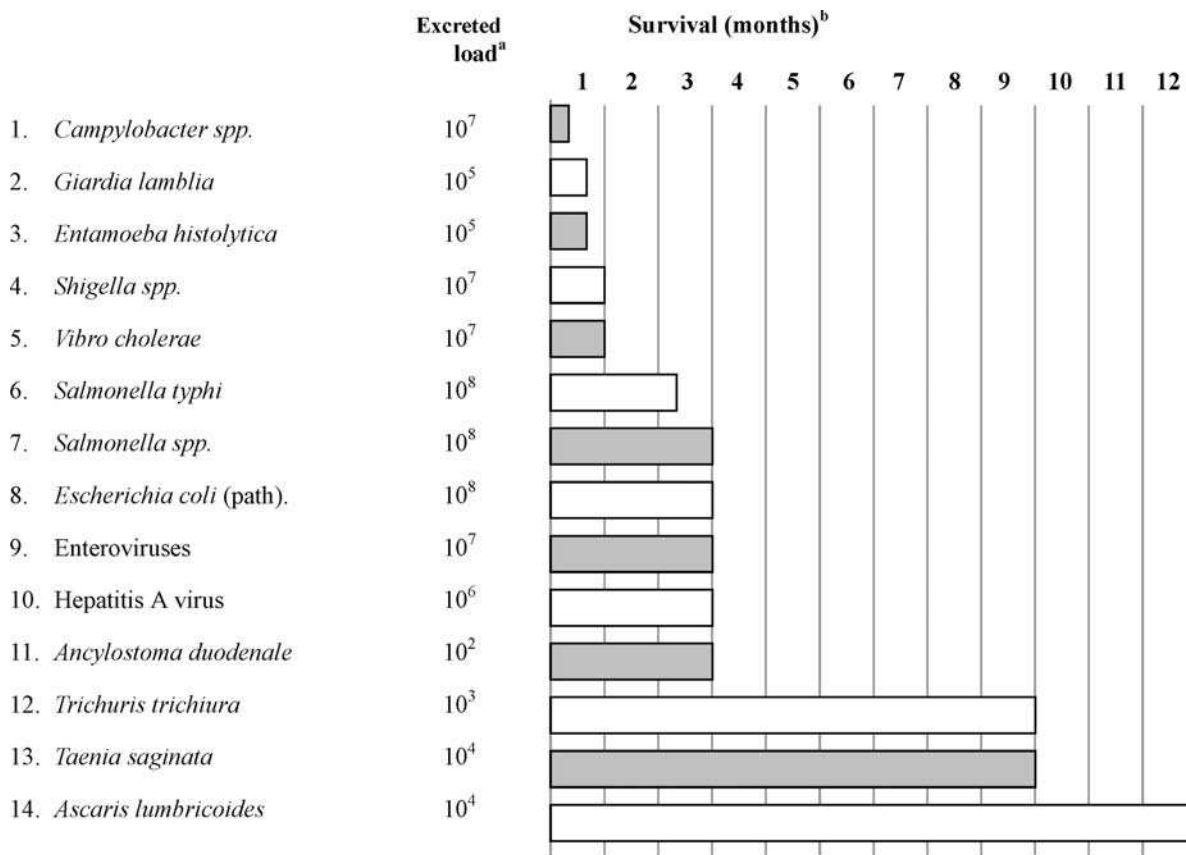
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able to survive in the environment (i.e. in water, soil, or food) for a period of time and they must be ingested in a sufficiently high number. Factors that affect the survival of pathogens in soil include antagonism from soil bacteria, moisture content, organic matter, pH, sunlight and temperature. Excreted enteric pathogens such as bacteria, viruses, protozoa, and helminth eggs do not usually penetrate undamaged vegetables but can survive for long periods in the root zone, in protected leafy folds, in deep stem depressions, and in cracks or flaws in the skin.

Data from numerous field and laboratory studies have made it possible to estimate the persistence of certain enteric pathogens in water, wastewater, soil, and on crops. These

survival periods in the environment are presented in summary graphic form in Fig. 15.1. For example, it appears that *Campylobacter* may survive in soil or on crops for only a few days, whereas most bacterial and viral pathogens can survive from weeks to months. The highly resistant eggs of helminths, such as *Trichuris*, *Taenia* and *Ascaris*, can survive for 9–12 months, but their numbers are greatly reduced during exposure to the environment.

Field studies in Israel have demonstrated that enteric bacteria and viruses can be dispersed for up to 730 m in aerosolized droplets generated by spray (sprinkler) irrigation, but their concentration is greatly reduced by detrimental environmental factors such as sunlight and drying (Teltsch *et al.*, 1980; Applebaum



^a Typical average number of organism/g feces

^b Estimated average life of infective stage at 20–30°C.

Fig. 15.1 Survival times of enteric pathogens in water, wastewater, soil and on crops (from Shuval *et al.*, 1986 - based partially on data from Feachem *et al.*, 1983).

et al., 1984; Shuval *et al.*, 1988, 1989a). Thus, most excreted pathogens can survive in the environment long enough to be transported by the wastewater to the fields and to the irrigated crops. The contaminated crops eventually reach the consumer, although by then the concentration of pathogens is greatly reduced. The rapid natural die-away of pathogens in the environment is discussed in a later section as it is an important factor in reducing the health risks associated with wastewater reuse.

Theoretical analysis suggests that a number of epidemiological factors determine whether various groups of pathogens will cause infections in humans through wastewater irrigation. We have developed a model to evaluate the empirical epidemiological data and to formulate control strategies (Shuval *et al.*, 1986).

Table 15.1 summarizes the epidemiological characteristics of the main groups of enteric pathogens as they relate to the five factors that influence the transmission and degree of infections and disease resulting from wastewater irrigation. This summary provides a simplified theoretical basis for ranking the groups of pathogens according to their potential for transmitting disease through wastewater irrigation. On this basis, it appears that the helminthic diseases are the ones most effectively transmitted by irrigation with raw wastewater because they persist in the environment for relatively long periods; their minimum infective dose is small; there is little or no immunity against them; concurrent infection in the home is often limited; and latency is long and a soil development stage is required for transmission.

In contrast, the enteric viral diseases should be least effectively transmitted by irrigation

with raw wastewater in developing countries with low levels of sanitation in the home, despite their small minimum infective doses and ability to survive for long periods in the environment. Due to poor hygiene in the home, and the prevalence of concurrent routes of infection in some areas, most of the population has been exposed to, and acquired immunity to, most of the enteric viral diseases as infants. Most enteric viral diseases impart immunity for life, or at least for very long periods, so that they are not likely to re-infect individuals exposed to them again, e.g. through wastewater irrigation. The transmission of bacterial and protozoan diseases through wastewater irrigation lies between these two extremes. In developed countries with higher levels of home sanitation and little concurrent disease transmission due to poor hygiene practice, there will be lower levels of immunity to diseases which could be transmitted by wastewater irrigation or vegetables eaten uncooked.

3 REVIEW OF RESEARCH FINDINGS ON DISEASE TRANSMISSION BY WASTEWATER IRRIGATION

This section will provide an extensive review and evaluation of the research findings on disease transmission by wastewater irrigation based on available scientific papers published in recognized journals and in numerous unpublished government reports, university theses, and private papers obtained during an intensive world-wide search carried out with the help of international and national

TABLE 15.1 Epidemiological characteristics of enteric pathogens *vis-à-vis* their effectiveness in causing disease through wastewater irrigation

Pathogen	Persistence in environment	Minimum infective dose	Immunity	Concurrent routes of infection	Latency/soil development stage
Viruses	Medium	Low	Long	Mainly home contact, food and water	No
Bacteria	Short/medium	Medium/high	Short/medium	Mainly home contact, food and water	No
Protozoa	Short	Low/medium	None/little	Mainly home contact, food and water	No
Helminths	Long	Low	None/little	Mainly soil contact outside home and food	Yes

agencies and individuals. Over 1000 documents, some more than 100 years old, were examined in the course of this study, but few offered concrete or reliable epidemiological evidence of health effects. Most of them based their conclusions on inference and extrapolation. Nonetheless, about 50 of these reports provided enough credible evidence based on sound epidemiological procedures to make a detailed analysis useful. Those studies are reviewed in detail in the UNDP-World Bank report on which this chapter is partially based (Shuval *et al.*, 1986). Our general conclusions of some of the more pertinent studies are presented below.

One of the goals of our studies for the UNDP/World Bank study (Shuval *et al.*, 1986; Shuval, 1990), as described in this chapter, was to re-evaluate all the credible, scientifically valid and quantifiable epidemiological evidence of the real human health effects associated with wastewater irrigation. Such evidence is needed to determine the validity of current regulations and to develop appropriate technical solutions for existing problems.

3.1 Illness associated with wastewater irrigation of crops eaten raw

In areas of the world where the helminthic diseases caused by *Ascaris* and *Trichuris* are endemic in the population, and where raw, untreated wastewater is used to irrigate salad crops and/or other vegetables generally eaten uncooked, the consumption of such wastewater-irrigated salad and vegetable crops may lead to significant levels of infection. Khalil (1931) demonstrated the importance of this route of transmission in his pioneering studies in Egypt. Similarly, a study in Jerusalem (Shuval *et al.*, 1984) provided strong evidence that massive infections of both *Ascaris* and *Trichuris* may occur when salad and vegetable crops are irrigated with raw wastewater. These diseases almost totally disappeared from the community when raw wastewater irrigation was stopped (Fig. 15.2). Two studies from Darmstadt, Germany (Krey, 1949; Baumhogger, 1949) provided additional support for this conclusion.

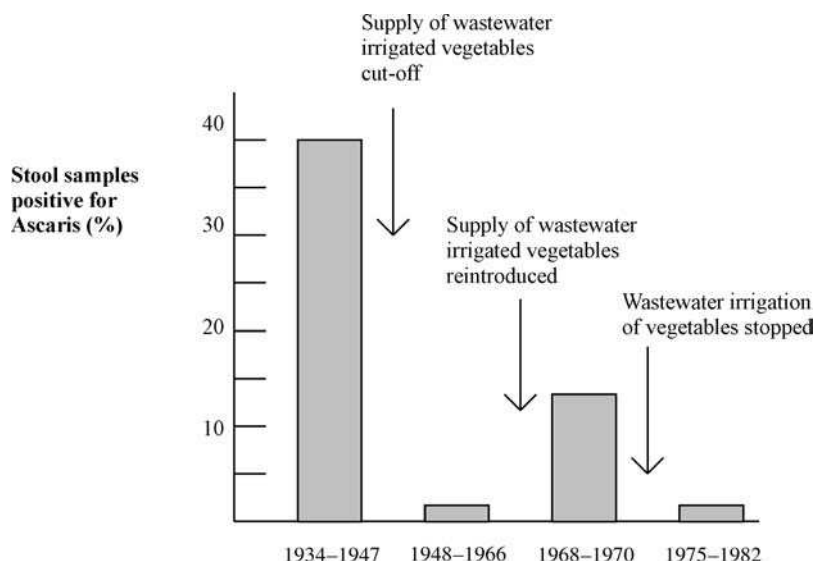


Fig. 15.2 Relationship between *Ascaris*-positive stool samples in population of western Jerusalem and supply of vegetables and salad crops irrigated with raw wastewater in Jerusalem, 1935-1982 (from Shuval *et al.*, 1986 – based on partially on data from Ben-Ari, 1962; Jjumba-Mukabu and Gunders, 1971; Shuval *et al.*, 1984).

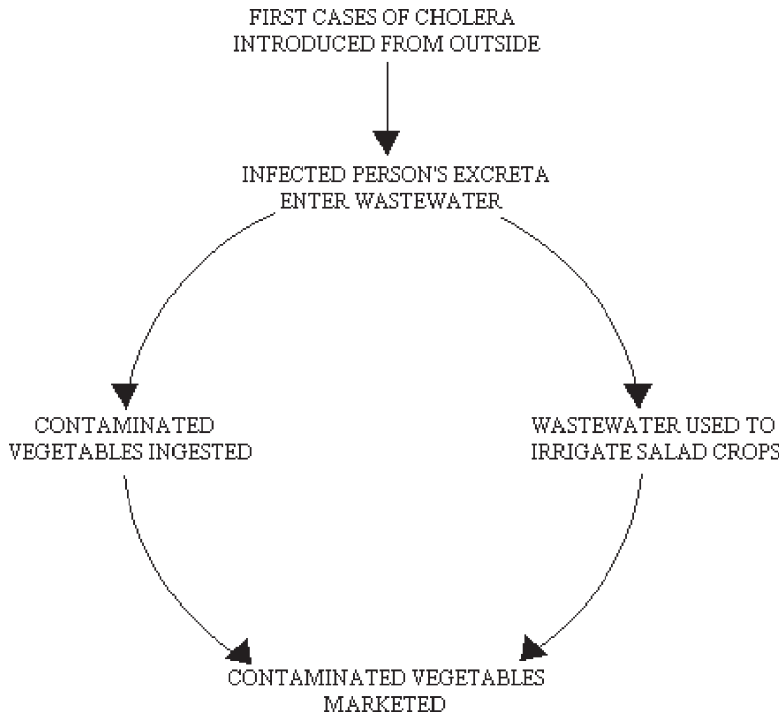


Fig. 15.3 Hypothesized cycle of transmission of *Vibrio cholerae* from first cholera carriers introduced from outside the city, through wastewater-irrigated vegetables, back to residents in the city (from Fattal *et al.*, 1986b).

These studies also indicate that regardless of the level of municipal sanitation and personal hygiene, irrigation of vegetables and salad crops with raw wastewater can serve as a major pathway for continuing and long-term exposure to *Ascaris* and *Trichuris* infections. Both of these infections are of a cumulative and chronic nature, so that repeated long-term re-infection may result in a higher worm load and increased negative health effects, particularly among children.

Cholera can also be disseminated by vegetable and salad crops irrigated with raw wastewater if it is carrying cholera vibrios. This possibility is of particular concern in non-endemic areas where sanitation levels are relatively high, and the common routes of cholera transmission, such as contaminated drinking water and poor personal hygiene, are closed. Under such conditions, the introduction of a few cholera carriers (or subclinical cases) into a community could lead to massive infection of the wastewater stream and subsequent transmission of the disease to the consumers of the vegetable crops irrigated with the raw wastewater, as occurred in

Jerusalem in 1970. The hypothesized cycle of transmission from the first imported case from outside the city through wastewater irrigated vegetables back to the residents is shown in Fig. 15.3 (Fattal *et al.*, 1986b).

Similarly, our study from Santiago, Chile (Shuval, 1984), strongly suggests that typhoid fever can be transmitted by fresh salad crops irrigated with raw wastewater. The number of typhoid fever cases in Santiago rose rapidly *annually* at the beginning of the irrigation season, after 16 000 ha of vegetables and salad crops (usually eaten uncooked) had been irrigated with raw wastewater. The relatively high socioeconomic level, good water supply, and good general sanitation in the city supports the hypothesis that wastewater irrigation can become a major route for the transmission of such bacterial disease.

3.2 Cattle grazing on wastewater irrigated pastures

Wastewater is often used to irrigate pasture for cattle and sheep. What are the health risks

associated with this practice? There is only limited epidemiological evidence to indicate that beef tapeworms (*Taenia saginata*) have been transmitted to populations consuming the meat of cattle grazing on wastewater-irrigated fields or fed crops from such fields. However, there is strong evidence from Melbourne, Australia (Penfold and Phillips, 1937), and from Denmark (Jepson and Roth, 1949) that cattle grazing on fields freshly irrigated with raw wastewater or drinking from raw wastewater canals or ponds can become heavily infected with the disease. This condition can become serious enough to require veterinary attention and may lead to economic loss. Irrigation of pastures with raw wastewater from communities infected with tapeworm disease may provide a major pathway for the continuing cycle of transmission of the disease to animals and humans.

3.3 Exposure of wastewater farmers to disease

Obviously the individuals most intensely exposed to the wastewater stream at the farms where wastewater irrigation is practised, are the farmers themselves. Some of the studies with the clearest epidemiological evidence relates to the health of such farm workers. Sewage farm workers exposed to raw wastewater in areas of India, where *Ancylostoma* (hookworm) and *Ascaris* infections are endemic, have much higher levels of infection than other agricultural workers (Krishnamoorthi *et al.*, 1973). The risk of hookworm infection is particularly great in areas where farmers customarily work barefoot, because the broken skin of their feet is readily penetrated by the motile hookworm larva. Sewage farm workers in this study also suffered more from anaemia (a symptom of severe hookworm infestation) than the controls. Thus, there is evidence that continuing occupational exposure to irrigation with raw wastewater can have a direct effect on human productivity and, thus, on the economy.

Sewage farm workers are also liable to become infected with cholera if the raw

wastewater being used for irrigation is from an urban area experiencing a cholera epidemic. This situation is particularly likely to arise in an area where cholera is not normally endemic and where the level of immunity among the sewage farm workers is low or non-existent. This proved to be the case in the 1970 cholera outbreak in Jerusalem (Fattal *et al.*, 1986b). In a related study (Fattal *et al.*, 1985), we have shown that irrigation workers exposed to aerosols from spray irrigation of both fresh water and wastewater had significantly higher rates of serum positivity to antibodies of *Legionella pneumophila*, the causative agent of Legionnaires' disease, as compared to the non-exposed control group of farmers and their families. This study does show that virulent pathogens can be transmitted by aerosols from spray irrigation and can infect highly exposed workers. However, it does not suggest that Legionnaires' disease is transmitted by wastewater irrigation any more than by fresh water, where the organisms are very often found.

Studies from industrialized countries have thus far produced only limited, and often conflicting, evidence of the incidence of bacterial and viral diseases among wastewater irrigation workers exposed to partly or fully treated effluent, or among workers in wastewater treatment plants exposed directly to wastewater or wastewater aerosols. Most morbidity and serological studies have been unable to give a clear indication of the prevalence of viral diseases among such occupational exposed groups.

It is hypothesized that many sewage farmers or treatment plant workers have acquired relatively high levels of permanent immunity to most of the common enteric viruses endemic in their communities at a much younger age. Thus, by the time they are exposed occupationally, the number of susceptible workers is small and not statistically significant. Presumably this is also the case among infants and children in developing countries, because they are exposed to most endemic enteric viral diseases by the time they reach working age. Although this is not the case for some bacterial

and protozoan pathogens, multiple routes of concurrent infection with these diseases may well mask any excess infection among wastewater irrigation workers in developing countries.

3.4 Exposure of residents in the vicinity of wastewater farms

A number of studies have evaluated the potential negative health effects that might result from living in the vicinity of farms where wastewater irrigation, particularly sprinkler irrigation is practised. There is little evidence linking disease and/or infection among population groups living near wastewater treatment plants or wastewater irrigation sites with pathogens contained in aerosolized wastewater. Most studies have shown no demonstrable disease resulting from such aerosolized wastewater, which is caused by sprinkler irrigation and aeration processes. Researchers agree, however, that most of the earlier studies have been inadequate.

Recent studies in Israel suggest that aerosols from sprinkler irrigation with poor microbial quality wastewater can, under certain circumstances, cause limited infections among infants living near wastewater-irrigated fields. The studies, however, also concluded that these were negligible and could be controlled by better treatment of the wastewater (Fattal *et al.*, 1986a, 1987; Shuval *et al.*, 1988, 1989b).

These findings support the conclusion that, in general, relatively high levels of immunity against most viruses endemic in the community block additional environmental transmission by wastewater irrigation. Therefore, the additional health burden is not measurable. The primary route of transmission of such enteroviruses, even under good hygienic conditions, is through contact infection in the home at a relatively young age. As already mentioned, such contact infection is even more common in developing countries, so that a town's wastewater would not normally be expected to transmit a viral disease to rural areas using it for irrigation.

3.5 Epidemiological evidence of beneficial effects from wastewater treatment

When raw wastewater is used for irrigation there is no doubt that the wastewater stream carries very high concentrations of pathogens. Conventional wastewater treatment plants were not normally designed to reduce the concentration of pathogenic microorganisms, however, such treatment can nonetheless provide a degree of removal up to about 85–95% reduction in coliform bacteria and pathogens. Some epidemiological studies have provided evidence that negative health effects can be reduced when wastewater is treated for the removal of pathogens. For example, Baumhogger (1949) reported that, in 1944, residents of Darmstadt who consumed salad crops and vegetables irrigated with raw wastewater experienced a massive infection of *Ascaris*; but the residents of Berlin, where biological treatment and sedimentation were applied to the wastewater prior to the irrigation of similar crops, did not.

Another study on intestinal parasites was conducted on school children near Mexico City (Sanchez Levya, 1976). The prevalence of intestinal parasites in children from villages that used wastewater irrigation did not differ significantly from that in children from the control villages, which did not irrigate with wastewater. The lack of significant difference between the two groups may have resulted from long-term storage of the wastewater in a large reservoir for weeks or months prior to its use for irrigation. It is assumed that sedimentation and pathogen die-away during long-term storage were effective in removing the large, easily settleable protozoa and helminths, which were the pathogens of interest in this study. This study provides the first strong epidemiological evidence of the health protection provided by microbial reductions achieved in wastewater storage reservoirs.

Furthermore, the absence of negative health effects in Lubbock, Texas (Camann *et al.*, 1983) and in Muskegon, Michigan (Clark *et al.*, 1981),

appears to be associated with the fact that well-treated effluents from areas of low endemicity were used for irrigation.

Data from these field studies strongly suggest that pathogen reduction by wastewater treatment, including long-term storage in wastewater reservoirs, can have a positive effect on human health. In all the above studies, this positive effect was achieved despite the use of effluent which had not been disinfected and which contained a few thousand of fecal coliform bacteria per 100 ml. These data agree with water quality data on pathogen removal and suggest that appropriate wastewater treatment resulting in effective reduction of coliforms to the level of a few thousand/100 ml, but not total removal, can provide a high level of health protection.

3.6 Conclusions from the analysis of the epidemiological studies

It is possible to draw certain conclusions from the series of epidemiological studies on the health effects of wastewater reuse in agriculture. The studies from both developed and developing countries indicate that the following diseases are occasionally transmitted via raw or *very poorly* treated wastewater:

1. The general public may develop ascariasis, trichuriasis, typhoid fever or cholera by consuming salad or vegetable crops irrigated with *raw wastewater*, and probably tapeworm by eating the meat of cattle grazed on wastewater-irrigated pasture. There may also be limited transmission of other enteric bacteria, viruses and protozoa.
2. Wastewater irrigation workers may develop ancylostomiasis (hookworm), ascariasis, possibly cholera and, to a much lesser extent, infection caused by other enteric bacteria and viruses, if exposed to raw wastewater.
3. Although there is no demonstrated risk to the general public residing in areas where wastewater is used in sprinkler irrigation, there may be minor transmission of enteric

viruses to infants and children living in these areas, especially when the viruses are not endemic to the area and raw wastewater or very poor quality effluent is used.

Thus, the empirical evidence on disease transmission associated with *raw wastewater* irrigation in developing countries strongly suggests that helminths are the principal problem, with some limited transmission of bacterial and viral disease. The above ranking, based on empirical data, agrees with that predicted in our model.

In interpreting the above conclusions, one must remember that the vast majority of developing countries are in areas where helminthic and protozoan diseases such as hookworm, ascariasis, trichuriasis, and tapeworm are endemic. In some of these areas, cholera is endemic as well. It can be assumed that in most developing countries, in populations with low levels of personal and domestic hygiene, the children will become immune to the endemic enteric viral diseases when very young through contact infection in the home.

In conclusion, epidemiological evidence of disease transmission associated with the use of *raw wastewater* in agriculture in developing countries indicates that the pathogenic agents may be ranked in the following order of declining importance:

1. *High risk*: helminths (*Ancylostoma*, *Ascaris*, *Trichuris* and *Taenia*)
2. *Lower risk*: enteric bacteria (cholera, typhoid, shigellosis and possibly others); protozoa (amebiasis and giardiasis)
3. *Least risk*: enteric viruses (viral gastroenteritis and infectious hepatitis).

As pointed out earlier, these negative health effects were all detected in association with the use of *raw or primarily treated* wastewater. Therefore, wastewater treatment processes that effectively remove all, or most, of these pathogens, according to their rank in the above list, could reduce the negative health effects caused by the utilization of raw wastewater. While helminths are very stable

in the environment, bacteria and viruses rapidly decrease in numbers in the soil and on crops.

Thus, the ideal treatment process prior to wastewater recycling and reuse, should be particularly effective in removing helminths, even if it is somewhat less efficient in removing bacteria and viruses. Wastewater treatment technologies that can be used to achieve this goal are discussed later in this chapter. In general, the above ranking of pathogens will not apply to the more developed countries or other areas in which helminth diseases are not endemic. In those areas the negative health effects, if any, resulting from irrigation with raw or partly treated wastewater will probably be associated mainly with bacterial and protozoan diseases and, in a few cases, with viral diseases. Whatever the country or the conditions, however, the basic strategies for control are the same – the pathogen concentration in the wastewater stream must be reduced and/or the type of crops irrigated must be restricted.

Overall, our studies have demonstrated that the extent to which disease is transmitted by wastewater irrigation is much less than was widely believed to be the case by public health officials in the past. Moreover, this study does not provide epidemiological support for the use of the much-copied California standard requiring a coliform count of 2/100 ml for effluent to be used in the irrigation of edible crops and even less support for the more recent USEPA/USAID recommended guideline of zero fecal coliforms. No detrimental health effects were detected or reported when well-treated wastewater with much higher coliform counts was used.

4 THE DEVELOPMENT OF HEALTH STANDARDS AND GUIDELINES FOR WASTEWATER REUSE

4.1 The importance of health guidelines and standards for reuse

One of the most important and widely practised administrative methods for protecting the public health from the risks of uncontrolled

wastewater irrigation, particularly of vegetables and salad crop consumed uncooked, is the establishment of guidelines or legally binding standards for the microbial quality of wastewater used for irrigation. This section will review the scientific basis and historical and social forces that influenced the evolution of microbial standards and guidelines for wastewater reuse for agricultural purposes. This analysis will draw extensively on World Bank and World Health Organization studies and reports whose goal was a cautious re-evaluation of the credible scientific evidence which could provide a sound basis for establishing safe and feasible health guidelines for wastewater reuse (Engelberg Report, 1985; Shuval *et al.*, 1986; WHO, 1989; Shuval, 1990).

The strict health regulations governing wastewater reuse that have been developed in the industrial countries over the past 60 years, such as those of the Department of Health of the State of California, which requires an effluent standard of 2 coliforms/100 ml for irrigation of crops eaten uncooked, and even the more recent USEPA/USAID (1992) recommended guidelines for unrestricted effluent use in agriculture of zero fecal coliforms, have been based to a great extent on early scientific data indicating that most enteric pathogens can be detected in wastewater and that they can survive for extended periods in wastewater-irrigated soil and crops (see Fig. 15.1). Many health authorities have erroneously concluded that, because pathogens can survive long enough to contaminate crops, even if their numbers are very low and below the minimum ineffective dose level, they still pose a serious risk to public health. However, these regulations were formulated at a time when sound epidemiological evidence was rather scanty. As a result, policy makers used the cautious 'zero risk' approach and introduced very strict regulations that they hoped would protect the public against the potential risks thought to be associated with wastewater reuse. Most industrial countries were not concerned that these regulations were overly restrictive because the economic and social benefits of wastewater reuse were of only marginal interest.

Many of the current standards restrict the types of crops to be irrigated with conventional wastewater effluent to those not eaten raw. Regulations like those in California, requiring the effluent used for the irrigation of edible crops to have a bacterial standard approaching that of drinking water (2 coliforms/100 ml), are usually not technically feasible or sustainable without very highly skilled operators and a high-tech service infrastructure. This is particularly true for developing countries, but even applies to many developed countries. In reality, a standard of 2 coliforms/100 ml for irrigation is superior to the quality of drinking water for the majority of urban and rural poor in developing countries (where fecal coliforms are generally in excess of 10/100 ml of drinking water).

In developed countries, where these crop restrictions can normally be enforced, vegetable and salad crops are not usually irrigated with wastewater. In the developing countries, many of which have adopted the same strict regulations, public health officials do not approve of the use of wastewater for irrigation of vegetable and salad crops eaten raw. However, when water is in short supply such crops are widely irrigated illegally with raw or poorly treated wastewater. This usually occurs in the vicinity of major cities, particularly in semi-arid regions.

Since the official effluent standards for vegetable irrigation are not within the obtainable range of common engineering practice and for economic considerations, new projects to improve the quality of effluent are not usually approved. With the authorities insisting on unattainable, expensive and unjustifiable standards, farmers are practising widespread uncontrolled and unsafe irrigation of salad crops with raw wastewater. The highly contaminated vegetables are supplied directly to the nearby urban markets, where such horticultural products can command high prices. This is a classic case in which official insistence on the 'best' prevents cities and farmers from achieving the 'good'.

Some inconsistency exists between the strict California standards, which require edible crops to be irrigated with wastewater of drinking water quality, and the actual agricultural irrigation with normal surface water as prac-

tised in the USA and other industrialized countries with high levels of hygiene and public health. There are few, if any, microbiological limits on irrigation with surface water from rivers or lakes, which may be polluted with raw or treated wastewater. For example, the US Environmental Protection Agency's water quality criteria for unrestricted irrigation with surface water is 1000 fecal coliforms/100 ml (USEPA, 1972). A WHO world survey of river water quality has indicated that most rivers in Europe have mean fecal coliform counts of 1000–10 000/100 ml. And yet none of these industrialized countries has restrictions on the use of such river water for irrigation.

A number of microbial guidelines have been developed for recreational waters considered acceptable for human contact and swimming. In the USA, for example, microbial guidelines for recreational water have, in the past, ranged from 200 to 1000 fecal coliforms/100 ml, although they currently are at about 10 fecal coliforms/100 ml. In Europe, guidelines vary from 100 coliforms/100 ml in Italy to 20 000 coliforms/100 ml in Yugoslavia. The European Community has recommended a guideline of 2000 fecal coliforms/100 ml for recreational waters (Shuval *et al.*, 1986).

It is difficult to explain the logic of a 2 coliforms/100 ml standard for effluent irrigation when farmers all over the USA and Europe can legally irrigate any crops they choose with surface water from free-flowing rivers and lakes, which often have fecal coliform levels of over 1000/100 ml. It is even more difficult to explain the epidemiological rationale of the 2 coliforms/100 ml standard for effluent irrigation, while in Europe recreational water for bathing is considered acceptable at 2000 fecal coliforms/100 ml.

4.2 The World Bank/World Health Organization initiative to re-evaluate wastewater reuse guidelines

Because of the questions raised in Section 4.1 above, and the fact that the strict coliform standards adopted by many countries were rarely enforced and often found to be unfeasible for

economic reasons and due to the lack of adequate technical infrastructure, in 1981, the World Bank and the World Health Organization initiated an extensive multidisciplinary study on the health effects of wastewater irrigation. The primary goal was to obtain an up-to-date scientific evaluation of the public health justification and validity of existing standards and guidelines and to develop alternatives if this was deemed to be justified. These studies were carried out by teams of epidemiologists, engineers, agronomists and environmental specialists simultaneously and independently at three different environmental sciences and public health research centers – the London School of Hygiene and Tropical Medicine, the International Reference Centre on Wastes Disposal, Zurich and the Division of Environmental Sciences of the Hebrew University of Jerusalem, Israel. These groups, working independently, prepared reports summarizing their findings, analysis and recommendations (Feachem *et al.*, 1983; Shuval *et al.*, 1986; Blum and Feachem, 1985).

In July 1985, a group of environmental experts, including engineers and epidemiologists, meeting at Engelberg, Switzerland, under the auspices of UNDP, World Bank, WHO, UNEP and IRCWD, reviewed the preparatory scientific studies which provided new epidemiological data and insights and from them formulated new proposed microbiological guidelines for treated wastewater reuse in agricultural irrigation (Engelberg Report 1985). The group accepted the main findings and recommendations of the UNDP-World Bank study (Shuval *et al.*, 1986) and concluded that 'current guidelines and standards for human waste use are overly conservative and unduly restrict project development, thereby encouraging unregulated human waste use'. The new guidelines recommended in the Engelberg Report were later accepted and approved by a WHO Meeting of Experts (WHO, 1989). However, before the WHO Executive approved those recommendations for publication as an official WHO document, the report was sent out for review and approval by a panel of some 100 epidemiologists, public health officials and environmental

engineers. Thus, these new recommended guidelines for the microbial quality of effluent used for wastewater irrigation of edible crops carry the stamp of approval of the highest international authority on public health and environmental matters. Other important international technical assistance agencies joined the WHO in supporting the new recommended guidelines for wastewater irrigation including the World Bank, The Food and Agricultural Organization (FAO), The United Nations Environment Program (UNEP) and the United Nations Development Program (UNDP). Meanwhile, a number of governments in developing and developed countries have adopted the new WHO recommended guidelines, including France.

There were a number of innovations in the new recommended guidelines. Since the possibility of transmitting helminth disease by wastewater irrigation of even non-edible crops was identified as the principal health problem, a new, stricter approach to the use of raw wastewater was developed. The new WHO guidelines recommend effective water treatment in all cases to remove helminths to a level of *one or fewer helminth eggs per liter*. The main innovation of the WHO guidelines is: for crops eaten uncooked, an effluent must contain one or fewer helminth eggs per liter, with a *geometric mean of fecal coliforms not exceeding 1000/100 ml*. This is a much more liberal coliform standard than the early California requirement of 2 total coliforms/100 ml.

An attractive feature of the new WHO (1989) effluent guidelines is that they can be readily achieved with low cost, robust waste stabilization pond systems and wastewater storage and treatment reservoirs that are particularly suited to developing countries. The high levels of pathogen removal that can be achieved by such low-cost stabilization pond systems are shown in Fig. 15.4. In conjunction with alternating wastewater storage reservoirs even higher degrees of treatment with an added safety factor can be achieved. Most studies indicate that the critical design parameter to achieve the WHO microbial guidelines in the effluent is a long detention period of up to 25–30 days in maturation ponds or long

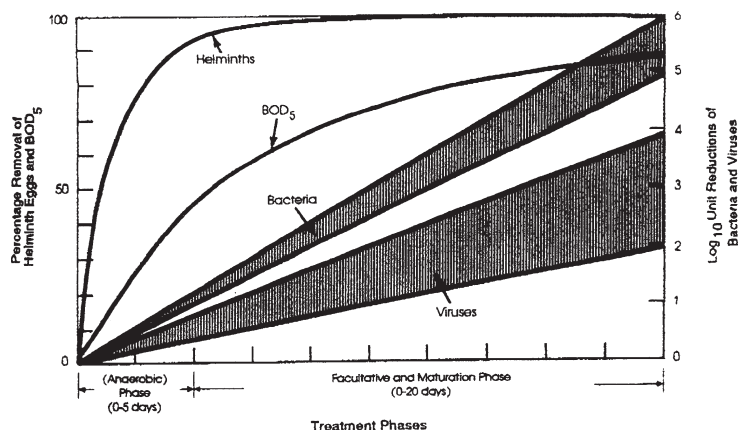


Fig. 15.4 Generalized removal curves for BOD, helminth eggs, excreted bacteria and viruses in wastewater stabilization ponds at temperatures above 20°C (from Shuval *et al.*, 1986).

detention times of several months in batch-operated storage reservoirs prior to the irrigation season (see Chapter 26).

4.3 The USEPA/USAID initiative for wastewater reuse guidelines

The US Environmental Protection Agency (USEPA), with the support of the US Agency for International Development (USAID), established their own rigorous recommended guidelines in 1992 of zero fecal coliforms/100 ml, a BOD of 10 mg/l, a turbidity of 2 NTU and a free chlorine residual of 1 mg/l. This quality of wastewater effluent can only be achieved in very costly high-tech wastewater treatment plants that require a high level of technological infrastructure for operation and maintenance, so that they can continuously meet such very rigorous standards. These guidelines were drafted by one of the leading American consulting engineering firms under contract to USAID. Such consulting engineering firms often tend to favor such high-tech treatment processes. Again, these new American guidelines are essentially as strict as those required for drinking water and reaffirm the 'no risk' or 'fail safe' approach that has been taken by the Americans in setting wastewater reuse guidelines and standards. The fact that little if any natural river water or water at approved bathing beaches in the USA or elsewhere could meet these recommended irrigation guidelines did not seem to bother those who drafted and approved the new American guidelines. No one

has suggested that such river water should not be allowed for irrigation purposes, nor has any health risk from such irrigation been reported.

5 THE DEVELOPMENT OF A RISK ASSESSMENT/COST-EFFECTIVENESS METHOD FOR EVALUATION OF GUIDELINES

The WHO Guidelines are supported by numerous international technical assistance agencies and have been adopted by France and a number of other developed and developing countries. However, some groups have favored the stricter 'no risk' USEPA/USAID recommended guidelines and have questioned whether the WHO guidelines provide a sufficiently high level of safety and health protection. The debate over the appropriateness of the various guidelines has so far been on a qualitative level.

We have carried out a study aimed at developing a quantitative risk assessment and cost-effectiveness approach based on a mathematical model and experimental data, to arrive at a comparative risk analysis of the various recommended microbiological guidelines for unrestricted irrigation of vegetables normally eaten uncooked (Shuval and Fattal, 1996; Shuval *et al.*, 1997). The guidelines that were compared are those of the World Health Organization (<1000 fecal coliforms/100 ml) and those recommended by the USEPA/USAID (zero fecal coliforms/100 ml).

5.1 Risk assessment model

For the purposes of this study, the risk assessment model, estimating the risk of infection and disease from ingesting microorganisms in drinking water, developed by Haas *et al.* (1993) was selected. However, certain modifications were required since we are estimating the risk of infection associated with eating vegetables irrigated with wastewater of various microbial qualities. The basic model of Haas *et al.* (1993) for the probability of infection (P_I) from ingesting pathogenic microorganisms in water is:

$$P_I = 1 - [1 + N/N_{50}(2^{1/\alpha} - 1)]^{-\alpha} \quad (1)$$

where

- P_I = risk of infection by ingesting pathogens in drinking water
- N = number of pathogens ingested
- N_{50} = number of pathogens that will infect 50% of the exposed population
- α = the ratio N/N_{50} and P_I

Various studies have shown a wide variation in the probability of infected persons becoming ill, with morbidity rates varying between 1 and 97%, depending on the virulence of the pathogen, and the age, nutritional and general health status of the subjects. Since not every person infected by the ingestion of pathogens becomes ill, an independent estimate is made of P_D , the probability of contracting a disease:

$$P_D = P_{D:I} \times P_I \quad (2)$$

- P_D = the risk of an infected person becoming diseased or ill
- $P_{D:I}$ = the probability of an infected person developing clinical disease.

5.2 Determining the number of pathogens ingested

In order to make an estimate of the number of pathogens ingested from eating wastewater-irrigated vegetables, we first had to determine, through laboratory experiments, the amount of liquid that might cling to vegetables irrigated with wastewater. It was then possible to estimate the concentration of indicator organisms and pathogens that might remain on

such wastewater irrigated vegetables and be ingested by subjects eating such vegetables. In doing this we assumed that any microorganisms contained in the residual wastewater remaining on the irrigated vegetables would cling to the vegetables even after the wastewater itself evaporated.

Based on laboratory determinations, we have estimated that the amount of wastewater of varying microbial qualities that would cling to the outside of wastewater irrigated cucumbers would be 0.36 ml/100 g (or one large cucumber) and 10.8 ml/100 g on long leaf lettuce (about three lettuce leaves).

Based on these measurements, we estimated the amount of indicator organisms that might remain on the vegetables if irrigated with raw wastewater and with wastewater meeting the WHO guidelines. In the case of irrigation with raw wastewater, we estimated that the fecal coliform (FC) concentration was 10^7 /100 ml. In the case of irrigation with wastewater meeting the WHO guidelines the FC concentration would be 10^3 /100 ml. We then estimated that the enteric virus:FC ratio in wastewater, based on various studies (Schwartzbrod, 1995), is $1:10^5$. For the purposes of this preliminary risk estimate we have assumed that all of the enteric viruses are a single pathogen, such as the virus of infectious hepatitis, so that it will be possible to make certain assumptions as to median infectious dose and infection to morbidity ratios. This errs on the conservative side.

We also assumed that under actual field conditions there would be a certain degree of indicator and pathogen die-away and/or removal from the wastewater source and irrigated vegetables until the final ingestion by the subject in the home. These factors include settling, adsorption, dessication, biological competition, UV irradiation from sunlight, and a degree of removal and/or inactivation by washing of the vegetables in the home. A number of studies have indicated that there is rapid die-off or removal of bacterial indicator organisms as well as pathogenic bacteria and viruses in wastewater irrigated soil and on crops by as much as 5 logs or 99.999% in 2 days under field conditions (Rudolfs *et al.*, 1951; Bergner-Rabinowitz, 1956; Sadovski *et al.*, 1978;

Armon *et al.*, 1995). Armon *et al.* also suggest that there is the possibility of the regrowth of bacteria on vegetables contaminated with wastewater, but presented no data to support this hypothesis. In any event, *human enteric viruses cannot ever multiply under environmental conditions*. Asano and Sakaji (1990) have determined virus die-off in the environment under field conditions of wastewater reuse, which indicate that in 2 weeks the total virus inactivation reaches some 99.99%, while in 3 days there is a 90% reduction of virus concentration. Even superficial washing of vegetables in the home can remove an additional 99–99.9% of the virus contamination. Schwartzbrod (1995) has estimated that there would be as much as a 6 log reduction (99.9999%) of virus concentration between irrigation with wastewater and consumption of the crops if the total time elapsed reached 3 weeks. To be on the conservative side, we have estimated that the total virus inactivation and/or removal from the wastewater source until ingestion results in a reduction in virus concentration by 3 logs, or 99.9%, although a 99.99% loss is not unreasonable and might occur in most cases.

5.3 Estimates of risk of infection and disease

Based on the above tests results and assumptions we now can estimate the number of pathogens ingested by a subject who eats a 100 g of cucumber or 100 g (three leaves) of long lettuces irrigated with wastewater of various qualities. In this preliminary risk estimate we have selected the enteric virus infectious hepatitis, which can result in serious disease sequelae and which has had a clear epidemiological record indicating the possibility of it being environmentally transmitted and waterborne (Schwartzbrod, 1995). We have assumed that the median infectious dose for 50% of the exposed subjects to become infected (N_{50}) could range between 30 and 1000 PFU. We have also assumed that while the ratio of infections to clinical disease is often as low as 100:1, we shall estimate, as a worst case, that 50% of those infected will

succumb to clinical disease ($P_{D:I} = 5$). We also assumed, based on vegetable consumption patterns in Israel, that on an annual basis a subject would consume 100 g of lettuce or cucumbers/day for 150 days ($P_{D:I} = 25$). We have assumed that $\alpha = 0.5$, however, even assuming $\alpha = 0.2$ does not lead to a significantly increased risk.

First, as a positive control test of the model, we have estimated the risk of infection and disease from consuming vegetables irrigated with raw wastewater with an estimated initial fecal coliform level of 10^7 . Based on the above assumptions, including a 3 log die-away, we have estimated that under such conditions a 100 g cucumber or 100 g (three leaves) of lettuce irrigated with raw wastewater will have a final level of contamination of FC of 10^{-2} . Based on that level of FC contamination and a virus:FC ratio of $1:10^5$, it can be estimated that there is a probability, in the case of irrigation with raw wastewater, that one cucumber in 10 000 will carry a single enteric virus and that one leaf of lettuce in 1000 will carry a single enteric virus. Based on these estimates of ingesting enteric viruses, more specifically the infectious hepatitis virus, we have estimated the risk of infection and disease that might result. The rate of disease, from eating 100 g lettuce, among a population eating such vegetables irrigated with *raw wastewater* is between 10^{-3} to 10^{-4} or about one case per 1000 to one case per 10 000 individuals exposed to risk.

This rate of infection has been found to correlate well with the disease rates detected in our field studies of the cholera outbreak in Jerusalem (Fattal *et al.*, 1986b) and typhoid fever in Santiago, Chile (Shuval, 1984), which were clearly associated with the irrigation of vegetables normally eaten uncooked with raw wastewater. Thus our estimates based on the laboratory data, our assumptions and using the Haas mathematical model provided a reasonably close approximation of the degree of risk of disease that occurs in real world situations. This is a vital step in the validation of the reliability and usefulness of such a predictive simulation model and helped assure us that our predictions of risk under various conditions have a reasonably sound basis.

We then proceeded to evaluate the risk of disease if crops were irrigated with wastewater that meets the WHO guidelines. If the effluent is treated to meet the WHO guidelines for irrigation of vegetables to be eaten uncooked of 1000 FC/100 ml, the risk of hepatitis infection and disease estimates for lettuce is reduced to about 10^{-7} – 10^{-8} , which means that the chances of becoming infected and diseased from such a low level of exposure is somewhere around one person per million people exposed per year or less. We have also calculated the annual risk of disease from the more infectious, but less severe, rotavirus as 10^{-5} – 10^{-6} /year.

Are these a high level of risk or a low level of risk? In order to shed some light on what are considered reasonable levels of risk for communicable disease transmission from environmental exposure it should be noted that the USEPA has determined that microbiological quality guidelines for drinking water microbial standards should be designed to ensure that human populations are not subjected to *risks of infection by enteric disease greater than 10^{-4} (or 1 case per 10 000 persons per year)* (Regli *et al.*, 1991). Thus, compared with the USEPA estimates of reasonable acceptable risks for water-borne disease acquired from drinking treated drinking water, the WHO wastewater reuse guidelines appear to be safer by some one to two orders of magnitude.

5.4 Cost/effectiveness analysis

At this stage of our study we have made only some very preliminary estimates of the cost/-effectiveness associated with meeting the various wastewater effluent guidelines. As an example we shall present the hypothetical case of a Third World city of 1 million population about to build a wastewater treatment plant to assure safe utilization of the effluent for agricultural irrigation of vegetable crops, including those eaten uncooked, which would serve the population of the city. It is assumed that they have opted for a waste stabilization pond system that will meet the WHO guidelines. They want to compare the cost and risks at that level of treatment to the cost and risks if they had adopted the USEPA/USAID guidelines for

treatment for vegetables eaten uncooked. We have assumed, for the purposes of this illustration only, that the unit cost of wastewater treatment to meet the various guidelines can roughly be estimated as follows:

WHO guidelines – 1000 FC/100 ml
(in stabilization ponds) \$0.125/M³
or the cost/person/year (assuming
100 M³/person/year) \$12.5/p/year
US-EPA/US-AID guidelines – 0 FC/100 ml
\$0.40/M³
or the cost/person/year (assuming 100 M³/
person/year) \$40.00/p/year.

The estimate of treatment costs for meeting WHO guidelines does not necessarily apply to all situations but is generally illustrative of a situation that may apply in hot sunny climates in developing countries where low cost land is available for effective stabilization pond treatment. According to this estimate, the additional cost for that city to meet the US-EPA/US-AID guidelines would be \$25 500 000/year.

We have assumed that half the population consumes wastewater irrigated vegetables on a regular basis and that the degree of annual risk of contracting a case of infectious hepatitis associated with the use of irrigated vegetables eaten uncooked with wastewater meeting the WHO guidelines is in the worst case some 2×10^{-6} (or about 1 case per year per 500 000 exposed persons) as estimated in this study. If we assume that the USEPA/ USAID guidelines, which require no detectable FC/100 ml, will achieve an essentially zero risk of disease, then we can estimate that the one case of infectious hepatitis per year would have been prevented. Thus the additional cost of treatment would result in a cost of about \$25 000 000 for each case of disease prevented. In the case of rotavirus disease, the cost would be some \$2 500 000 per case prevented. If, however, the true level of risk associated with the WHO guidelines is closer to the 10^{-7} level, estimated by the less conservative interpretation of the results of this study, then no detectable reduction of risk would

be gained by the additional investment of \$25 000 000 required to meet the strict and expensive USEPA/USAID guidelines.

It is questionable whether this level of additional treatment associated with major additional expenditures is justified to reduce further, the already negligibly low levels of risk of infection and disease that our estimates indicate are associated with the new WHO Guidelines.

Our conclusion is that the new WHO guidelines, which are based on extensive epidemiological evidence and can be achieved with low-cost wastewater treatment technology, provide a high degree of public health protection at a reasonable cost. There is little or no justification, in our opinion, for the unreasonably restrictive 'zero risk' USEPA/USAID guidelines, which are exceedingly expensive to achieve and require costly high-tech wastewater treatment technology, which provides little if any measurable increase in health protection.

Thus, after well over a century, health guidelines for wastewater reuse have gone through a complete cycle from no regulation or control in the 19th century to unreasonably strict standards in the earlier part of this century to what now appears to be a scientifically sound and rational basis with a less restrictive approach as recommended by the WHO guidelines. It is hoped that this new approach will encourage the development of controlled wastewater reuse for the benefit of mankind, while providing an appropriate level of health protection.

6 CONTROL OF CROPS AND IRRIGATION METHODS TO REDUCE HEALTH RISKS

Early in the development of wastewater irrigation for agriculture, methods of reducing health risks by controlling the type of crops grown or the methods of irrigation have been proposed and in some cases used effectively. The risk of transmission of communicable disease to the general public by irrigation with raw or settled wastewater can be reduced by a number of agronomic techniques. Some of these restrict

the types of crops grown, and others, through modification and/or control of irrigation techniques, prevent or limit the exposure of health-related crops to pathogens in the wastewater.

6.1 Regulating the type of crops

One of the earliest and still most widely practised remedial measures is to restrict the type of crops irrigated with raw wastewater or with the effluent of primary sedimentation. Since there is ample evidence that salad crops and other vegetables normally eaten uncooked are the primary vehicles for the transmission of disease associated with raw wastewater irrigation, forbidding the use of raw effluent to irrigate such crops can be an effective remedial public health measure. Although such regulations have been effective in countries with a tradition of civic discipline and an effective means for inspection and enforcement of pollution control laws, they will likely be of less value in situations where those preconditions are absent.

In many arid and semi-arid areas near major urban centers, where subsistence farmers irrigate with raw wastewater, the market demand for salad crops and fresh vegetables is very high. Thus, governmental regulations forbidding farmers to grow such crops would be little more than a symbolic gesture. Even under the best of circumstances, it is difficult to enforce regulations that work counter to market pressures; to enforce regulations that prevent farmers from obtaining the maximum benefit from their efforts under conditions of limited land and water resources would be impossible.

6.2 Controlling irrigation methods

Basin irrigation of salad and vegetable crops usually results in direct contact of the crops with wastewater, thus introducing a high level of contamination. Sprinkler irrigation of salad crops also results in the deposit of wastewater spray on the crops and their contamination. The level of contamination may be somewhat less than basin irrigation. Many vegetables

that grow on vines (i.e. tomatoes, cucumbers, squash, and the like) can be partially protected from wastewater contact if properly staked and/or grown hanging from wires that keep them off the ground, although some of these vegetables will inevitably touch the ground.

Well-controlled ridge-and-furrow irrigation reduces the amount of direct contact and contamination. These methods cannot completely eliminate direct contact of the wastewater with leafy salad crops and root crops.

Drip irrigation causes much less contamination of the crops than any other irrigation method. In fact, our studies indicate the use of drip irrigation tubes under polyethylene plastic surface sheeting used as a mulch can vastly reduce or totally eliminate crop contamination. In our studies, we determined that the level of enteroviruses and bacterial indicator organisms on cucumbers grown under such protective drip irrigation systems was negligible during the first 24 hours after the introduction of a massive seeding of microorganisms into the wastewater effluent stream used for irrigation. After 24 hours no contamination of irrigated crops was detectable (Sadovski *et al.*, 1976, 1978). Drip irrigation is the most costly form of irrigation, but its hygienic advantages make it attractive as a safe method of wastewater irrigation of sensitive vegetable and salad crops, even when the microbial quality of the effluent is not up to the strictest standards.

Fruit orchards do well with basin or ridge-and-furrow irrigation, but normal overhead sprinkler irrigation leads to direct contamination of the fruit. With low-level low-pressure sprinkler irrigation, however, the main spray is below the level of the branches, and the fruit is less likely to be contaminated. In all cases, windfalls picked from the ground will have been in contact with wastewater-contaminated soil. Another possible control measure is to discontinue irrigation with wastewater at a specified period, such as 2 weeks, before harvesting the crop. This option is feasible for some crops, but the timing of a vegetable harvest is difficult to control. In addition, some types of vegetables are harvested over long periods of time from the same plot.

Some of the above irrigation control techniques can help reduce the danger of crop contamination, but they are feasible only in fairly advanced and organized agricultural economies. Health regulations dependent upon any of the above procedures to protect certain high-risk crops from contamination must be enforced by legal sanctions and frequent inspections. If well-organized inspection and law enforcement systems are not present, as in some developing countries, the value of these options as a major remedial strategy may be limited. However, in the case of large centrally operated sewage farms, managed by the government or large well-organized companies, such procedures can be of value.

7 WASTEWATER TREATMENT

7.1 What are the goals of wastewater treatment for recycling and reuse?

Obviously, the degree of wastewater treatment, particularly as it relates to the effective removal and inactivation of pathogenic microorganisms, will have a critical effect in controlling any possible health risks associated with wastewater irrigation. The microbiology of wastewater treatment is reviewed in depth in Chapters 19–28. Thus it is beyond the scope of this chapter to review all possible wastewater treatment technologies suitable for wastewater recycling and reuse. However, we shall attempt to review some general principles and to emphasize low cost treatment technologies particularly suited to warm climates in developing countries.

In areas with plentiful rainfall, wastewater has traditionally been disposed of or diluted in large bodies of water, such as rivers and lakes. High priority has been given to maintaining the oxygen balance of these bodies of water to prevent serious detrimental effects, such as anaerobic conditions and odors from wastewater pollution. Most of the conventional processes used to treat wastewater in industrial countries have been designed primarily to remove the suspended and dissolved organic

fractions, which decompose rapidly in natural bodies of water. The organic matter in wastewater, usually measured as biochemical oxygen demand (BOD), provides rich nutrients to the natural microorganisms of the stream, which multiply rapidly and consume the limited reserves of dissolved oxygen (DO) in the streams. If oxygen levels drop too far, anaerobic conditions may develop, serious odors may evolve and fish may die. However, for recycling and reuse of wastewater for agricultural irrigation, a high degree of BOD removal in the effluent is not directly relevant since, with land disposal, the soil is not harmed, but benefits from high level of organic matter and plant nutrients deposited on it. However, removal of organic matter, as represented by BOD, may be necessary when chemical disinfection processes are required, after the biological treatment stage, to achieve the microbial effluent standards.

A secondary goal of conventional wastewater treatment has been to reduce pathogenic microorganisms in order to protect the quality of the sources of drinking water used by downstream communities. However, conventional biological wastewater treatment systems, such as biological filtration and activated sludge, are not particularly efficient in removing pathogens. Thus, communities that draw their drinking water from surface sources cannot depend upon upstream wastewater treatment plant systems to reduce pathogens to a safe level. Therefore, they must remove the pathogens with their own drinking water treatment plants using a series of highly efficient, technical, and costly processes (i.e. coagulation, sedimentation, filtration and chemical disinfection). The most effective conventional wastewater treatment system is activated sludge, which removes 90–99% of the viruses, protozoa and helminths and 90–99.9% of the bacteria. Conventional processes cannot achieve higher levels of pathogen removal without additional expense for chemical disinfection, such as chlorination, and additional sand filtration. Further research and development are needed to improve the removal of helminths by conventional methods. However, the new micro/nanofiltration

processes hold much promise in the effective removal of helminths and all other pathogenic microorganisms from wastewater.

7.2 Low cost wastewater stabilization ponds provide high quality effluent

In contrast to conventional treatment systems, studies have shown that well-designed multi-cell stabilization ponds allowing 20–30 days of retention can remove almost 100% of the helminth eggs (Yanez *et al.*, 1980; Feachem *et al.*, 1983; Mara and Silva, 1986; Fattal *et al.*, 1998). Bacteria, viruses and protozoa are often attached to larger fecal particles that settle out in pond systems. At best, however, only 90% can be removed by sedimentation.

The most effective process for removing bacteria and viruses in stabilization ponds is natural die-off, which increases with time, pH and temperature. Many developing countries have hot climates in which stabilization ponds are exposed to the direct rays of the sun and may reach temperatures up to 40°C. The pH at midday is commonly 9 or higher due to the photosynthetic activity of the algae. Predatory or competing microorganisms may also affect die-off by attacking or damaging pathogens directly or indirectly. Exposure to the ultraviolet rays of the sun may also play a role in killing pathogens in ponds. Long retention times, however, appear to be the most important factor in reducing bacterial concentrations in pond systems.

In warm climates with temperatures in excess of 20°C, a pond system with 4–5 cells and a 20- to 30-day retention time usually reduces the fecal coliform concentration by 4–6 log orders of magnitude – i.e. by 99.99–99.9999%. Thus, if the initial concentration of fecal coliform bacteria in the raw effluent is approximately $10^7/100$ ml, the effluent will contain 10^3 or 1000/100 ml. The same pond system will reduce enteric viruses by 2–4 log orders of magnitude (i.e. from an initial concentration of about 1000/100 ml to 10 or fewer/100 ml). Helminth eggs will be removed almost completely, while the BOD will be reduced by about 80%. Fig. 15.4 shows the generalized removal curves for BOD, helminth

eggs, bacteria, and viruses in a multicell stabilization pond system in a warm climate.

Our cooperative studies with Egyptian colleagues in the city of Suez on waste stabilization pond treatment of wastewater for irrigation and aquaculture have demonstrated that with well-designed and well-operated multicell ponds with 30 days of detention, an effluent of 10–100 fecal coliforms/100 ml was consistently achieved (Mancy, 1996). In our studies at a waste stabilization pond pilot plant system, treating primary effluent from the Jerusalem Municipal treatment plant, an effluent of 1000 fecal coliforms/100 ml was achieved with 25–30 days of detention (Fattal *et al.*, 1997).

Waste stabilization ponds are therefore highly suitable for treating wastewater for irrigation. They are more efficient in removing pathogens, particularly helminths, than are conventional wastewater treatment systems. In addition, they produce a biologically stable, odorless, nuisance-free effluent without removing too many of the nutrients. Thus, ponds should be the system of choice for wastewater irrigation in warm climates, especially if land is available at a reasonable price. Ponds are particularly attractive for developing countries because they cost little to build and maintain and are robust and fail-safe. They should never be considered a cheap substitute. In reality they are superior to conventional methods of treatment in almost all respects. Although ponds require relatively large land areas, land costs are, in many cases, not a serious obstacle.

When wastewater will be used to irrigate crops for human consumption, the goals of treatment are the reverse of the goals of conventional treatment. The primary goal for treatment of wastewater to be used for irrigation must be removal of pathogenic microorganisms in order to protect the health of the farmers and consumers. Removal of the organic material, however, which contains valuable agricultural nutrients is neither necessary nor desirable, although aerobic conditions should be maintained because a black, highly odorous, anaerobic wastewater

effluent would probably be an environmental nuisance to farmers and nearby residents.

7.3 Public health advantages of wastewater treatment and storage reservoirs

Since wastewater is generated by the community 365 days a year and the irrigation season in most areas is limited to a number of months per year, a means must be found to handle wastewater flows during non-irrigation periods. If allowed to flow unrestricted, the effluent will contaminate the region's natural bodies of water.

A suitable solution is the storage and treatment reservoirs pioneered in Israel (Juanico and Shelef, 1994; Pearson *et al.*, 1996). Such reservoirs are designed to upgrade the quality of the effluent during the long residence time in the reservoirs and to store up to 8–10 months of wastewater flow in the rainy winter months to be used for irrigation during the dry summer season. They are often preceded by settling ponds and/or by conventional stabilization ponds, and may also be designed to catch surface runoff. The operational regimes of such reservoirs vary. Some are operated to change between non-steady-state flow and batch while others are operated sequentially, receiving and storing influent for extended periods, after which the inflow is stopped for a period of stabilization and bacterial decay prior to the discharge for irrigation. Log mean concentration of microorganisms in often partially treated typical wastewater influent at the entrance of the reservoir per 100 ml are: heterotrophic bacteria 10^7 – 10^8 , fecal coliforms 10^6 – 10^7 , enterococci 10^4 – 10^6 and F^+ bacteriophages 10^5 – 10^6 .

The results at the outlet in such reservoirs show that, in general, the quality of the effluents is much improved and is best at the beginning of the irrigation season when the reservoir is full of old effluent (long detention time), but sharply deteriorates when the water level drops and new wastewater continues to be pumped into the reservoir

(Juanico and Shelef, 1994; Pearson *et al.*, 1996). The die-off rate of microorganisms in the summer is higher than in the winter. We have shown that there is a significant correlation between the mean hours of sunshine in the month and the rate of bacterial die-off. Thus with greater sunshine duration in the summer months the bacterial die-off is highest (Fattal *et al.*, 1996). However, in single reservoirs during the summer the relative volume of the influent, raw or partially treated is higher, leading to poorer bacterial quality of the effluent since the pond levels are at their lowest during the intensive irrigation season. Coliform removal is high in the epilimnion, where high pH values occur due to algal activity, and low in the hypolimnion where pH values are low (Liran *et al.*, 1994). The logs mean reduction of heterotrophic bacteria, *E. coli*, enterococci and F⁺ bacteriophages at the outlet of the reservoir are: 1, 3, 2.5 and almost 4 logs, respectively (Fattal *et al.*, 1993). The mean reduction of BOD is 76%, COD 72% and TSS 45% (Fattal *et al.*, 1993).

In order to improve the quality of the effluent, innovative changes were made in Israel in the design of such stabilization/storage reservoirs (Liran *et al.*, 1994; Juanico and Shelef, 1994; Juanico, 1996; Friedler and Juanico, 1996). The storage of the effluent was changed from seasonal to multiseasonal and/or from single reservoir to two or more reservoirs used sequentially as batch reservoirs, supplying effluents for irrigation only from reservoirs which no longer receive fresh influent. By doing so, the input of the effluent from treatment units into the reservoir is stopped before reservoir effluent is released for irrigation. Another improvement was made by implementing better treatment of the wastewater before entering into the reservoir. The performance of the improved batch stabilization reservoirs, when properly designed and operated, showed that they are able to remove fecal coliform bacteria by up to five orders of magnitude (i.e. 99.999%). Such improved alternating reservoirs systems can normally produce an effluent which can easily meet the WHO guidelines for unrestricted irrigation of 1000 fecal coliforms/100 ml or less.

7.4 Remedial environmental methods assures greater health protection

The history of public health progress has proven that the broadest and most effective public health benefits are obtained from remedial or preventive measures taken by a central authority and involving environmental interventions that lower the level of exposure of large populations to environmentally transmitted disease. Such measures as central plants for the purification of drinking water supplies, pasteurization of milk, and area-wide campaigns for reducing the breeding sites of malaria-carrying mosquitoes are well-known examples of success using this strategy. Any remedial action based on changing personal behavior and lifestyle through education, law enforcement, or both, is a much slower process and, in general, has succeeded only in areas with relatively high educational levels and living standards.

The wastewater storage and treatment/storage reservoir option reviewed above offers this type of centrally managed and engineered form of remedial environmental intervention. It is the only remedial measure that will simultaneously reduce the negative health effects for sewage farm workers and for the public that consumes wastewater-irrigated vegetables. It is also the only measure that can bring about health benefits in a short time without massive changes in personal behavior or restrictive regulations that depend on complex inspection and law enforcement procedures. However, it does require central organizational and management capacity, availability of major financial resources and availability of land.

Although it may be appropriate in some situations to restrict the type of crops grown or to control wastewater irrigation practices, such regulations are difficult to enforce where there is great demand for salad crops and garden vegetables. In arid and semi-arid zones (as well as some humid areas), where irrigation is highly desirable, many economists and agricultural authorities consider it economically prudent to allow unrestricted wastewater irrigation of cash crops in high demand. That

goal can only be achieved with an effective high level of wastewater treatment as suggested in this chapter.

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Developing risk assessments of waterborne microbial contaminations

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1 INTRODUCTION

Pathogens can and do gain entry into drinking water supplies even in industrialized countries. Break-through during treatment and ingress through cracked pipes are well-documented causes of waterborne outbreaks of cryptosporidiosis (Craun *et al.*, 1998) and *Escherichia coli* O157 (Swerdlow *et al.*, 1992), respectively. Microbiological risk assessment (MRA) is the emerging method to predict the risks to public health from these pathogens. Models have been developed for a range of waterborne pathogens in drinking water, including *Cryptosporidium parvum* (Teunis *et al.*, 1997), *Giardia* (Regli *et al.*, 1991) and enteric viruses (Haas *et al.*, 1993). Most of the models represent non-outbreak conditions and model the endemic levels of infection through drinking water. Recently, a model for *C. parvum* has been developed for conditions representative of a waterborne outbreak of cryptosporidiosis (Gale and Stanfield, 2000).

The risk assessment approach can be used to answer questions ('what if?' scenarios), such as how many more people will be infected if part of the drinking water treatment plant fails and by how much will public health be jeopardized if disinfection is eliminated. One application of MRA is to provide a defensive position for new and emerging agents in the absence of epidemiological evidence. For this reason, a risk assessment for the transmission of bovine

spongiform encephalopathy (BSE) to humans through groundwater supplies was developed (Gale *et al.*, 1998). In the absence of epidemiological data, the primary objectives of the MRA were therefore to determine whether the BSE agent in the environment could conceivably present a risk to those drinking water consumers supplied by water sources near to rendering plants and to determine whether drinking water supplies were adequately protected. The objective was to develop order of magnitude estimates of risks of infection.

Over the last few years the application of MRA has expanded greatly to cover other environmental routes, including pathogens in:

- sewage sludges applied to agricultural land; e.g. BSE (Gale and Stanfield, 2001), salmonellae (Gale, 2001b) and *E. coli* O157
- manures and animal slurries applied to agricultural land
- composted domestic wastes applied to land
- rendering plant condensates discharged to surface waters.

2 THE SOURCE-PATHWAY-RECEPTOR APPROACH

Environmental risk assessments are based on the source-pathway-receptor approach. The 'source' term defines the amount of infectivity in the environment, and the 'receptor' term

comprises the various human and animal groups exposed to infectivity. The 'pathway' term describes the routes by which receptors may be exposed. These are best modelled by event trees. Central to environmental MRAs are pathway barriers and biomedical barriers (Gale, 2001b). Pathway barriers include drinking-water treatment processes (chemical coagulation, sand filtration), the hydrogeological substrata above aquifers, sewage sludge treatment processes, decay of the agent in the environment, and dilution. The pathway barriers serve to reduce exposure. The biomedical barriers include the species barrier in the case of BSE (Gale *et al.*, 1998), acquired protective immunity in *C. parvum* infection (Chappell *et al.*, 1999) and natural gut microbiota in salmonella and *Cryptosporidium* infection (Meynell, 1963; Harp *et al.*, 1992). The biomedical barriers serve to protect the receptors from infection after exposure. In general, there is greater uncertainty in the biomedical barriers than the pathway barriers.

2.1 The use of the arithmetic mean exposure

Gale and Stanfield (2000) used a Monte Carlo approach to simulate the variation in doses of *Cryptosporidium* oocysts ingested by individual drinking water consumers under conditions representative of a waterborne outbreak (Table 16.1). Log-normal distributions for the oocyst densities in 100–l volumes of raw and treated waters are shown in Fig. 16.1 The simulation is based on the Poisson-log-normal distribution and takes into account:

- the variation in oocyst densities in the raw water, in this case a river with agricultural drainage (Hutton *et al.*, 1995)
- the variation in removal of oocysts at the drinking-water treatment works based on particle removal data for 67 works (Fig. 16.2)
- the variation in the volumes of tap water ingested daily by drinking water consumers (0.1–3 l/person/day; Roseberry and Burmaster, 1992).

TABLE 16.1 Daily exposures to *Cryptosporidium* oocysts simulated for individual consumers under conditions consistent with an outbreak (Gale and Stanfield, 2000). Doses ingested are compared accommodating the variation predicted using the Poisson-log-normal distribution and simply using the arithmetic mean exposure in the Poisson distribution

Dose (oocysts/ person/day)	Percentage of population exposed to dose	
	Poisson-log-normal distribution* (Monte Carlo simulation)	Arithmetic mean exposure (0.37 oocysts/person/day) in Poisson distribution
0	88.43	68.9
1	6.76	25.7
2–5	3.57	5.4
6–10	0.61	0.00027
11–50	0.57	<10 ⁻¹⁰
51–100	0.05	0
>100	0.01	0

* The arithmetic mean of the simulated exposures = 0.37 oocysts/person/day.

According to the Poisson-log-normal distribution of exposures simulated in Table 16.1, the majority of consumers do not ingest any oocysts each day, even during an outbreak. This is consistent with the finding that 66% of 1000 litre volumes did not contain oocysts during the Farmoor (UK) outbreak of cryptosporidiosis (Richardson *et al.*, 1991) and that oocysts are often not detected in outbreaks (Joseph *et al.*, 1991; Craun *et al.*, 1998). The simulation predicts that about half of those consumers who were exposed ingested just a single oocyst each day. However, a small proportion of consumers (1.2%) were exposed to between 6 and 50 oocysts/day. Furthermore, 1 in 10 000 consumers was exposed to very high daily doses, which exceeded 100 oocysts. These high count exposures reflect a combination of temporally high raw water loadings (e.g. after heavy rain or agricultural discharge) and poor oocyst removal efficiencies by treatment, e.g. because of failure or bypass (Craun *et al.*, 1998).

The arithmetic mean for the simulated oocyst exposures was 0.37 oocysts/person/day.

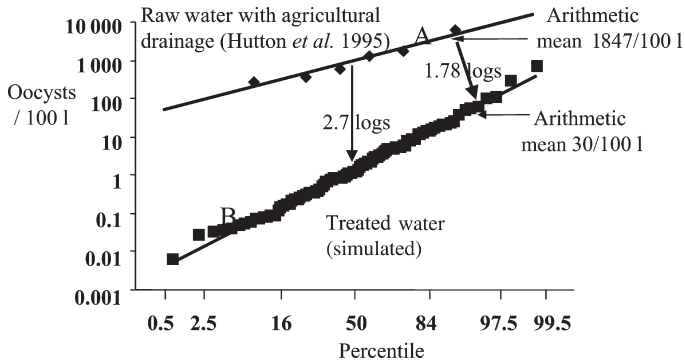


Fig. 16.1 Simulated log-normal distribution of densities of *Cryptosporidium* oocysts in treated water during a waterborne outbreak (Gale and Stanfield, 2000).

Exposures calculated using this single point average and assuming a Poisson distribution are also presented in Table 16.1. MRA models, which use this single point average with the Poisson distribution, and ignore the large variation in the simulated Poisson-log-Normal exposures, would predict that a much higher proportion of the population is actually exposed daily to oocysts (31% compared to

11%), but that those who are exposed never ingest more than about 5 oocysts each day. By ignoring the temporal/spatial heterogeneity in exposures, MRA would never predict that a consumer would be exposed to high pathogen doses in a single exposure.

The expected numbers of *Cryptosporidium* infections across the population for the two exposure scenarios in Table 16.1 are very similar. Indeed the arithmetic mean exposure of 0.37 oocysts/person/day may be used directly in the dose–response curve, even though it represents a fraction of a pathogen. Thus, Haas (1996) demonstrated mathematically that the appropriate statistic for risk assessment is the arithmetic mean exposure, even for log-normal distributions of exposures, which show great variation in magnitude across the population (Table 16.1). This ‘rule’ only holds if the pathogens act independently and breaks down for the more highly infectious agents such as rotavirus.

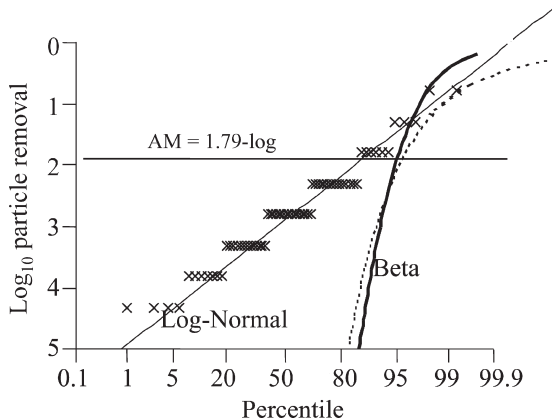


Fig. 16.2 Distribution of particle (>3 μm) removal ratios by drinking water treatment. Data for 67 drinking water treatment plants from LeChevallier and Norton (1995). Log-normal distribution ($\mu = -2.70$; $\sigma = 0.89$) as fitted by Gale and Stanfield (2000) and beta distribution (solid line) fitted to data ($\alpha = 0.0162$; $\beta = 0.986$). Arithmetic mean (AM) removal is same for both models. In addition the beta distribution ($\alpha = 0.025$; $\beta = 4.10$) for removal of spores of sulohite-reducing bacteria as presented by Teunis *et al.* (1997) is shown as dashed line.

2.2 Presentation of risks

A Monte Carlo simulation will give a range of predicted risks for individual consumers within the population. Thus, in the outbreak simulation, consumers are exposed to between 0 and >100 oocysts/person/day (see Table 16.1). This raises the question of what value should be quoted for the risk. Typically 5, 50 and 95 percentiles are quoted. However, the actual risk values for these percentiles will depend critically on the distribution of pathogens within the medium. This is shown with an example in

TABLE 16.2 Three exposure scenarios for 50 *C. parvum* oocysts distributed within 100 litre of drinking water and 50 consumers each ingesting 2 litres

Exposure model	Percentile exposure (oocysts/person)			Number of infections * per 50 persons
	5	50	95	
Arithmetic mean:	1	1	1	
All 50 persons ingest 1 oocyst each				0.209
Poisson distribution of exposures:	0	1	3	0.208
18 persons ingest 0 oocysts				
19 persons ingest 1 oocyst				
9 persons ingest 2 oocysts				
3 persons ingest 3 oocysts				
1 person ingests 4 oocysts				
Extreme clustering:	0	0	0	0.189
49 persons ingest 0 oocysts				
1 person ingests all 50 oocysts				

* Calculated using dose-response relationship in Equation (1) where $r = 0.00419$ (Haas *et al.*, 1996).

Table 16.2 which considers 50 *C. parvum* oocysts in 100 litre of drinking water, and 50 consumers each drinking 2 litre of that water. Thus, in the case of the extreme clustering distribution scenario, the 5, 50 and 95 percentile risks are all zero. This questions the appropriateness of quoting these percentile values and, in particular, the median (or 50% percentile) risk. Indeed, quoting a 5, 50 or 95 percentile risk merely gives information on the least exposed portion of the population and raises the question of what risks are the remaining proportion exposed to. The 'bottom line' is how many infections are there across the whole population. For all three scenarios the group risk across the 50 individuals is similar (i.e. between 0.19 and 0.21 infections per 50 persons). In this respect, the appropriate value is the arithmetic mean risk. This, in effect, represents the number of infections across the population as a whole. The median risk is typically in the region of an order of magnitude less than the arithmetic mean risk for a log-normal model, and may thus give too optimistic a representation (Fig. 16.1).

2.2.1 The arithmetic mean is appropriate for each of the steps in the source-pathway-receptor approach

The source, pathway and receptor terms used by Gale and Stanfield (2000) to simulate *Cryptosporidium* exposures during an outbreak

are presented in Table 16.3. Gale and Stanfield (2000) demonstrated by Monte Carlo simulation that the arithmetic means are all that is required for each of the source, pathway and receptor terms. Indeed, providing the dose-response relationship is linear, then the variation in the source, pathway and receptor terms (Table 16.3) may be accommodated simply by using the arithmetic mean. This provides a 'quick and simple' alternative to Monte Carlo simulations. Thus, the arithmetic mean *Cryptosporidium* oocyst exposure in Table 16.3 is calculated as:

$$18.47 \div 61.3 \times 1.26 = 0.38 \text{ oocysts/person/day}$$

which is very similar to the value of 0.37 oocysts/person/day calculated from the simulated Poisson-log-normal distribution of exposures (Table 16.1). This exposure may then be used directly in the dose-response curve (see the nature of the dose-response relationship, p. 267) to calculate the risk of infection (see Integrating pathogen exposures, p. 271). The main sources of uncertainty are underestimating the arithmetic mean oocyst loading in the raw water (or treated water by monitoring programmes) and overestimating the net oocyst removal by drinking water treatment. Both contribute to underestimating the arithmetic mean exposure and hence the risk of infection.

TABLE 16.3 Source, pathway and receptor terms used to simulate exposures to *Cryptosporidium* oocysts through drinking water (Gale and Stanfield, 2000)

Term	Data	Arithmetic mean*	Variation	Uncertainty
Source	Oocyst concentrations in raw water	18.47 oocysts/l	Natural fluctuations in oocyst concentrations, e.g. after heavy rainfall	Monitoring protocol missing high count spikes and recovery efficiency of analytical method contribute to underestimating net oocyst loading [†]
Pathway	Removal of oocysts from raw water by drinking water treatment	61.3-fold (arithmetic mean removal ratio)	Fluctuations in removal efficiency between and within individual treatment works	Monitoring protocol may cause overestimation of net removal efficiency [‡]
Receptor	Volume of tap water ingested daily by individual consumers	1.26 l/person/day	Individual habits, age groups, seasonal effects, amount imbibed after boiling	Method of data collection, e.g. self-assessment
Exposure	Simulated	0.37 oocysts/person/day		

* Calculated from 10 000 samples 'drawn' from log-normal distributions by Monte Carlo simulation.

[†] Uncertainty–Monitoring may underestimate the arithmetic mean pathogen exposure and hence risk (p. 273).

[‡] Uncertainty–Estimating the removal of pathogens by treatment processes (p. 274).

2.3 The nature of the dose–response relationship

Dose–response data are critical for quantitative MRA. Dose–response data from human studies have been obtained for several water-borne pathogens, including rotavirus (Ward *et al.*, 1986) and *C. parvum* (DuPont *et al.*, 1995; Chappell *et al.*, 1999). Dose–response parameters have also been estimated for BSE in humans (Gale, 2001a).

2.3.1 *Cryptosporidium parvum*

In feeding trials, 29 healthy immunocompetent adult volunteers were each given a single dose of between 30 and 10^6 oocysts of a calf strain of *C. parvum* (DuPont *et al.*, 1995). The proportion infected at each dose is plotted in Fig. 16.3. Exposure simulations presented in Table 16.1 suggest that, even under outbreak conditions, most drinking-water consumers ingest fewer than 30 oocysts daily and half of those consumers exposed to oocysts ingest only a single oocyst. Therefore, the doses administered by DuPont *et al.* (1995) are generally too high for the application of modelling risks through

water. To estimate the risk from ingestion of just a single *C. parvum* oocyst, a process of low dose extrapolation of fitted mathematical dose response curves is used.

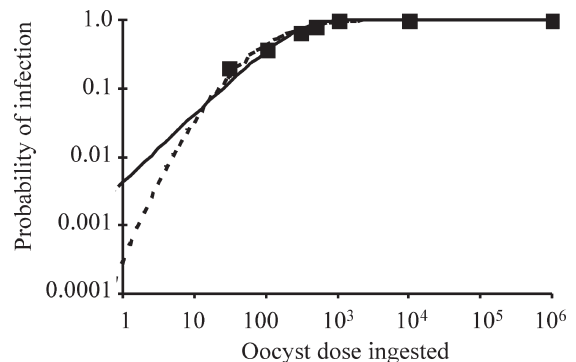


Fig. 16.3 Independent or co-operative action. Negative exponential dose response curve (Equation 1; $r = 0.00419$) fitted by Haas *et al.* (1996) to *C. parvum* human infectivity data (N) obtained for volunteers selected on the basis of having no serological evidence of the past infection with *C. parvum* (DuPont *et al.*, 1995). The log-probit curve (dashed) ($\mu = 2.119$, $\sigma = 0.614$; \log_{10}) is shown for comparison.

Haas *et al.* (1996) fitted a negative exponential dose–response curve to the *C. parvum* human infectivity data (Fig. 16.3). According to this model, the probability, P_i , of being infected by ingesting a dose of N pathogens is expressed mathematically as:

$$P_i = 1 - e^{-rN} \quad (1)$$

where r is a parameter specific for the pathogen and the host population and represents the fraction of the ingested pathogens that survive to initiate infections (Regli *et al.*, 1991). This dose–response relationship assumes that the oocysts act completely independently during infection and do not act cooperatively, for example, in overcoming the host protective barriers. Indeed at low doses, Equation (1) transforms to:

$$P_i = rN \quad (2)$$

indicating a direct linear relationship between the ingested dose, N , and risk of infection. The risk of infection from ten oocysts, for example, is tenfold that from the ingestion of just a single oocyst. If pathogens acted cooperatively during infection, then at low doses, the probability of infection would be non-linearly related to the dose. One mathematical model that describes a non-linear dose–response relationship is the log-probit curve (Haas, 1983). This model is defined by two parameters and is also shown in Fig. 16.3. It is apparent that the negative exponential and the log-probit curves do not differ markedly over the range of doses administered in the trial (30 to 10^6 oocysts). However, more considerable differences are observed on extrapolation to the low doses of oocysts to which drinking water consumers are likely to be exposed (see Table 16.1). The risk of infection predicted by the log-probit model from ingestion of a single oocyst is about 15-fold smaller than that predicted by the negative exponential model (Fig. 16.3). This reflects the fact that, if oocysts acted cooperatively, then low doses (e.g. one oocyst) would present greatly diminished risks. According to the log-probit model, the risk of infection from a dose of 10 oocysts is 120-fold higher than from ingestion of only a single oocyst.

2.3.2 The term ‘infective dose’ is meaningless

The extreme scenario for a non-linear dose–response relationship is the threshold or minimum infective dose. The concept of a minimum infective dose or threshold is misleading in MRA because it suggests zero risk of infection at very low pathogen doses. This does not appear to be the case for *C. parvum*. Teunis (1997) has demonstrated that fitted models assuming threshold doses of 2, 3, 4, or 5 oocysts do not fit the infectivity data in Fig. 16.3 and lead to increasingly steep dose–response curves. There is no experimental evidence for a minimum infective dose and no reason why a *single* oocyst or other waterborne pathogen should not be able to initiate infection, albeit with very low probability in some cases. Indeed, individual bacterial cells and virions contain all the macromolecular machinery and genes for infection. The diverse pathogenic strategies of enteric bacteria are based on a limited array of macromolecular systems (Donnenberg, 2000). Blewett *et al.* (1993) concluded that the minimum infective dose for gnotobiotic lambs is just one oocyst of *C. parvum*. According to Blewett *et al.* (1993), *C. parvum* is very infectious to gnotobiotic lambs and the ID₅₀ is less than 10 oocysts. The ID₅₀ for *C. parvum* in healthy human adults (with no acquired protective immunity) is 165 oocysts (Fig. 16.4), and single oocysts appear to present much lower risks to healthy human adults than to gnotobiotic lambs. The reason for this may lie in the nature of the host protective barriers and, in particular, the indigenous microbiota. Harp *et al.* (1992) presented data supporting the hypothesis that resistance of adult mice to *C. parvum* infection is mediated by non-specific mechanisms associated with the presence of intestinal microbiota.

In terms of dose–response curves, there is no such quantity as an ‘infective dose’. Indeed, each dose has a specific probability of initiating infection according to the dose–response curve (see Fig. 16.3). The ‘minimum infective dose’ is one pathogen.

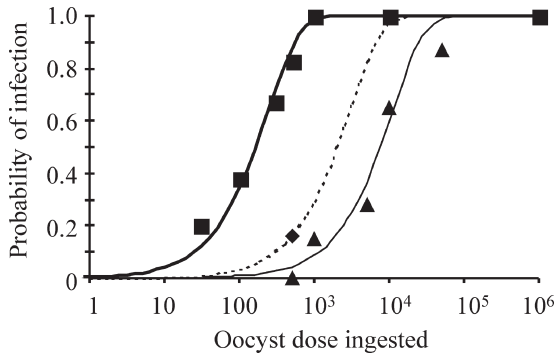


Fig. 16.4 Acquired protective immunity for *C. parvum* (Iowa isolate) in humans. Negative exponential dose response curve ($r = 0.00419$) fitted by Haas *et al.* (1996) to human infectivity data (*N*) obtained for volunteers selected on the basis of having no serological evidence of past infection with *C. parvum* (DuPont *et al.*, 1995). On rechallenge one year later (Okhuysen *et al.*, 1998), 16% of those volunteers were infected with a dose of 500 oocysts (*U*). The negative exponential curve ($r = 0.00035$) through this point is presented as the dashed line. Negative exponential curve ($r = 0.000091$) for confirmed infection (Chappell *et al.*, 1999) in human volunteers selected with pre-existing anti-*C. parvum* IgG (*S*).

2.3.3 Assessing the evidence that pathogens act independently and do not cooperate during infection

Meynell (1957) demonstrated that fatal infections from *Salmonella typhimurium* given to mice by mouth arose from a very small number of organisms, or possibly just a single organism. He concluded that inoculated bacteria act independently, not cooperatively. However, it could be argued that, if infection through the oral route were a two-stage mechanism, then his conclusions are only applicable to the second stage and that the initial stage *could* involve events more consistent with a cooperative model. The oral route presents many barriers to infection by salmonellae, including pH and the combined inhibitory effects of a low oxidation-reduction potential and short-chain fatty acids produced by the normal gut microbiota (Meynell, 1963). Indeed Meynell (1955) reported that when the gut microbiota were killed by streptomycin, the oral LD₅₀ for a

streptomycin-resistant strain of *S. typhimurium* was reduced from greater than one million organisms to less than five organisms.

Bacteria contain a variety of mechanisms for overcoming host barriers, including endotoxins to damage the mucosa, neutralization of antibacterial agents, IgA proteases and acid suppression. In addition, microorganisms produce many substances which give them competitive advantage, or a means of defence, in establishing themselves in natural environments. For example, the bacteriocins are polypeptide antibiotics produced by one species of bacteria, which cause damage to the cellular membranes of other bacteria. It could be speculated that in an oral challenge comprising thousands of salmonellae, some of those salmonellae could use these mechanisms to lower the host barriers, so increasing the chances of a subsequent single bacterium successfully establishing infection. Essentially this would involve a two-stage process for microbial infection. Meynell and Maw (1968) presented results with salmonellae in mice suggesting that infection passed through an initial stage lasting a few hours, in which a varying proportion of the inoculum is killed, followed by a second stage. The initial stage is 'decisive'. Could it be that bacteria act cooperatively during that initial stage, so helping subsequent bacteria to survive and progress to the second stage in which a single bacterium multiplies and causes infection? If so, then exposure to low pathogen doses would present lower risks than those predicted by the negative exponential or beta-Poisson dose-response curves and a non-linear model perhaps resembling the log-probit curve would be more appropriate (see Fig. 16.3, for example). There are currently no experimental data on whether *C. parvum* oocysts act independently or cooperatively during infection. Indeed, oocysts may not even have the biochemical mechanisms necessary. It is generally assumed for MRA that oocysts act independently and that the negative exponential dose-response curve is appropriate (Haas *et al.*, 1996). For the purpose of public health protection, this assumption is acceptable because it gives an added margin of safety for exposures to low

doses. However, if there were some cooperative mechanism for oocysts then MRA could overestimate the risks by as much as 15-fold (see Fig. 16.3), which would be unacceptable for water company operators.

The recent discovery of small molecules for pathogen communication demonstrates a potential mechanism for pathogen cooperation, although whether this could occur in the human gut is not known. Thus, cell-to-cell communication is used by bacterial cells to sense their population density (Dong *et al.*, 2001). This is known as quorum-sensing and controls expression of certain genes, which control biological functions such as virulence. The best characterized example is autoinduction of luminescence in the marine bacterium *Vibrio fischeri* (Fuqua *et al.*, 1996). Quorum-sensing signals, such as acyl-homoserine lactones (AHLs), are central to the communication system. The quorum-sensing regulation of virulence is a common strategy that many bacterial pathogens have adopted during evolution to ensure their survival in host–pathogen interactions. The expression of virulence genes in the Gram-negative bacterium, *Pseudomonas aeruginosa*, for example, requires cell-to-cell communication (Passador *et al.*, 1993). Among the extracellular virulence factors produced by *P. aeruginosa* is a protease, elastase, which cleaves elastin, human immunoglobulins, collagens and components of the complement system. These are all components of the host defence system. Passador *et al.*, (1993) speculate that when carried out in response to cell density or other environmental and nutritional stimuli, cell-to-cell communication could result in a concentrated attack on the host. Cellular communication in pathogenic bacteria is an area where discoveries in molecular biology may input to the future development of quantitative MRA. Whether quorum-sensing is important in botulism toxin production by *Clostridium botulinum* spores, or toxin production by *Bacillus anthracis* spores would be an important factor in MRA, and would necessitate the development of models to take into account the spatial heterogeneity of the spores (see Within-batch or spatial variation, p. 275) in making exposure assessments. There is evi-

dence that campylobacters at high concentrations protect each other. The same would be true for an aggregate of virus particles. Furthermore, there could be cooperative effects of viruses during infection of target cells. Thus binding of one virus to receptors on the cell membrane could facilitate binding of further virus particles, such as in the formation of clathrin-coated pits.

2.3.4 Acquired protective immunity for *C. parvum*

The 29 adults used in the study of DuPont *et al.*, (1995) were selected on the basis of having no serological evidence of past infection with *C. parvum*. Information on the extent of seroprevalence in various drinking-water communities and the quantitative protective effect is necessary for development and validation of MRA models for *C. parvum* in drinking water. Anti-*C. parvum* IgG is typically found in 25–33% of persons in the USA (Chappell *et al.*, 1999). Okhuysen *et al.*, (1998) investigated if infection of humans with *C. parvum* is protective 1 year after exposure. The subjects were 19 of those 29 adults used in the study of DuPont *et al.*, (1995). Rechallenge with a dose of 500 oocysts resulted in infection of three of the 19 adults (16%). Constructing a negative exponential dose–response curve through this one result (the dashed curve in Fig. 16.4) suggests that primary exposure increased the ID₅₀ from 165 oocysts to about 1900 oocysts and that the risk of infection (r in Equation (1)) from a single oocyst is decreased from 0.0042 to 0.00035. In further human infectivity studies, groups of volunteers with anti-*C. parvum*-specific serum antibody were challenged with doses between 500 and 50 000 oocysts (Chappell *et al.*, 1999). These points are also shown in Fig. 16.4. The negative exponential dose–response curve ($r = 0.000091$) is shifted to the right, and the ID₅₀ is 7600, i.e. 46-fold higher than for volunteers with no serum antibody.

2.3.5 Human rotavirus

The beta-Poisson dose–response curve assumes that r in Equation (1) is actually not a constant but is itself described by a beta

probability distribution. The probability of infection is written as:

$$P_i = 1 - [1 + \frac{N}{N_{50}}(2^{1/\alpha} - 1)]^{-\alpha} \quad (3)$$

where α and N_{50} characterize the dose–response (Regli *et al.*, 1991). As α increases so the beta-Poisson model becomes closer to the negative exponential model (Equation (1)). N_{50} represents the ID_{50} .

Ward *et al.*, (1986) obtained infectivity data for human rotavirus in 62 healthy adult volunteers. Subjects ingested doses ranging from 0.009 to 90 000 focus-forming units (ffu). The proportions of adults infected at each dose are plotted in Fig. 16.5. The beta-Poisson dose–response curve fitted by Haas *et al.* (1993) is plotted, together with a two-component negative exponential curve, which is written as:

$$P_i = f_1(1 - e^{-r_1 N}) + f_2(1 - e^{-r_2 N}) \quad (4)$$

where f_i represents the fraction of the population represented by the negative exponential model defined by r_i . The sum of f_1 and f_2 is always equal to 1.0 in a two-component model. The two-component model assumes there are two human populations which differ fundamentally in their susceptibility to rotavirus infection. This could reflect molecular differences, e.g. in the rotavirus receptor on the

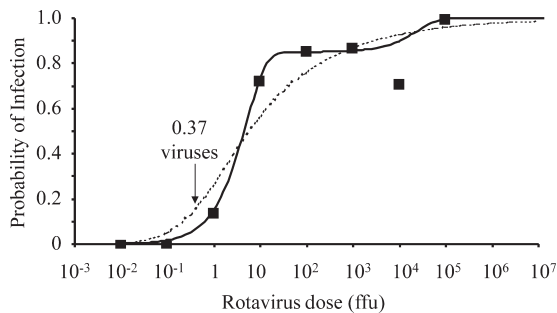


Fig. 16.5 Beta-Poisson dose-response curve ($\alpha = 0.265$; $N_{50} = 5.6$) (dashed line) fitted by Haas *et al.* (1993) to rotavirus infectivity data (N) of Ward *et al.* (1986). Two component negative exponential model (solid line) fitted assuming $r_1 = 0.20$ for 85% (f_1) of the population and $r_2 = 0.00004$ for the remaining 15% (f_2) in Equation 4.

human host cells, although there is no evidence currently for this. The sequencing of the human genome and the science of bioinformatics may shed light on this in the future (International Human Genome Sequencing Consortium, 2001). In the case of BSE infection in humans, epidemiological data for vCJD and genetic differences (polymorphisms) in the human prion protein gene at codon 129 provide a theoretical basis for a two-component dose–response curve (Gale, 2001a). The two-component negative exponential model fits the rotavirus data better (Fig. 16.5) than the beta-Poisson model, particularly at the lower doses. The beta-Poisson curve predicts risks 3.5-fold higher for the lower doses.

3 INTEGRATING PATHOGEN EXPOSURES AND DOSE–RESPONSE CURVES

The effect of the variation in the magnitude of the pathogen doses ingested by individual drinking-water consumers (see Table 16.1) on risk prediction is illustrated using the dose–response curves for *C. parvum* (Fig. 16.4) and rotavirus (Fig. 16.5) to predict the number of infections across the population (Table 16.4).

3.1 Risk prediction for *C. parvum*

Ignoring the log-normal variation in oocyst exposures and just using the Poisson distribution of exposures with the log-probit dose–response curve underestimates the risk of infection by a factor of about ninefold. This is because the Poisson distribution of pathogen exposures does not predict exposures to high pathogen doses approaching the ID_{50} (see Table 16.1). It only predicts exposures to low doses which have a greatly diminished risk according to the log-probit curve (see Fig. 16.3). It is concluded that modelling the log-normal variation in pathogen exposures would be critically important for MRA if pathogens were to act cooperatively during infection. Indeed, if pathogens acted cooperatively then dispersion within the drinking-water supply (i.e. going from Poisson-log-normal to Poisson)

TABLE 16.4 Predicted number of infections (per 10 000 persons/day) using simulated pathogen exposures in Table 16.1 with *C. parvum* (Iowa isolate) and rotavirus dose–response curves

Pathogen	Dose–response curve	Exposure model		
		Poisson-log-normal ($\mu = -1.66$; $\sigma = 1.04 \log_{10}$ oocysts/person/day)	Poisson distribution (mean = 0.37 oocysts/person/day)	Arithmetic mean (0.37 oocysts/ person/day)
<i>C. parvum</i> (Fig. 16.3 and Fig. 16.4)	Negative exponential (no immunity)	14.97*	15.44*	15.47
	Log-Probit (no immunity)	9.65*	1.70*	0.17
	Negative exponential (volunteers with anti- <i>C. parvum</i> IgG in serum)	0.336*	0.336*	0.336
Rotavirus (Fig. 16.5)	Beta-Poisson	410*	886*	1488
	Two component negative exponential model	331*	551*	605

* The risk calculation is described fully in Gale and Stanfield (2000).

would reduce the risk by a factor of fivefold according to Table 16.4. However, it is never appropriate to use the arithmetic mean in the log-probit model. Furthermore, if there were morbidity and mortality thresholds, as suggested by Williams and Meynell (1967), then MRA models need to accommodate the log-normal variation in exposures to cover the possibility of some consumers ingesting high doses which exceed those thresholds.

In contrast, the risk predicted using the negative exponential dose–response curve is little affected by whether the log-normal variation in exposures is accommodated or just the single point arithmetic mean exposure is used in the Poisson distribution (see Table 16.4). For both the Poisson-log-normal and Poisson distributions of exposure, a risk of 15 *Cryptosporidium* infections/10 000 persons/day is predicted. Indeed, the same risk is predicted simply by using the arithmetic mean of 0.37 oocysts/person/day directly in Equation (1). This is because the dose–response relationship is essentially linear (Equation (2)). In effect, the risk of infection is directly related to the total number of oocysts in supply and is not influenced by their spatial distribution (Gale, 2001a).

It should be noted that acquired protective immunity reduces the risk of infection by a factor of 46-fold (see Table 16.4).

3.2 Risk prediction for rotavirus

Using the arithmetic mean exposure of 0.37 virus/person/day directly in the beta-Poisson dose–response curve (arrow in Fig. 16.5) predicts 1488 infections/10 000 persons/day (see Table 16.4). In contrast, accommodating the log-normal variation in exposures in Table 16.1 predicts 410 viral infections/10 000 persons/day. Thus, ignoring the log-normal variation in virus exposures through drinking water would, in the case of the virus risk assessment, over-predict the risk by 3.6-fold. This is partly due to the fact that ignoring the log-normal variation predicts that more consumers would be exposed to highly infectious virus each day (see Table 16.1). In effect, dispersion of the virus across the population increases the number of infected persons. The effect is less with the two-component dose–response curve.

3.3 Dispersion of pathogens serves to increase the risk

In the event of pathogens cooperating during initiation of infection, dispersion of the pathogens within the medium would naturally lower the risk (see Table 16.4). However, assuming pathogens act independently, then dispersion serves to increase the number of infections in

situations where the exposure/ ID_{50} ratio is high. This is apparent for rotavirus in Table 16.4 (and also to a lesser degree for *C. parvum* in Table 16.2). Thus, dispersing the virus particles (i.e. going from Poisson-log-normal to Poisson) doubles the number of infections in the population predicted by the beta-Poisson dose–response curve. Using the arithmetic mean directly increases the predicted number of infections still further. In this respect, using the arithmetic mean gives the worst case. In general, pathogen exposures to individuals through drinking water are low relative to the ID_{50} (see Table 16.1) and it is acceptable to use the arithmetic mean. Only in the case of highly infectious agents such as rotavirus does the arithmetic mean over-predict the risk. However, consider a 100 g porcine kidney infected with 100 oral ID_{50} of foot and mouth disease virus and discarded into pig swill. Dispersion of the kidney tissue in the swill would increase the number of infected pigs considerably.

4 UNCERTAINTY – MONITORING MAY UNDERESTIMATE THE ARITHMETIC MEAN PATHOGEN EXPOSURE AND HENCE THE RISK

The counts of *Cryptosporidium* oocysts in 100-litre volume ‘spot’ samples of treated water in an outbreak simulation are presented in Table 16.5. Taking single 100-litre volume spot samples from the drinking-water supply would, in general, underestimate the arithmetic mean oocyst concentration and hence the risk of *Cryptosporidium* infection. Indeed, almost nine out of every ten (88%) 100-litre samples contain fewer oocysts than the arithmetic mean of 29.75 oocysts/100 litre. Furthermore, 54.75% of the samples contained 0, 1, or 2 oocysts, i.e. >10-fold less than the arithmetic mean (Table 16.5). Further simulations showed that even taking the arithmetic mean of the counts for ten 100-litre random spot samples underestimated the ‘true’ arithmetic mean (29.75 oocysts/100 litre) by more than fivefold some 25% of the time. Thus a quarter of all sets of ten samples

TABLE 16.5 Simulated counts of *Cryptosporidium* oocysts (Poisson-log-normal distribution) in 100-litre volumes under conditions consistent with an outbreak (Gale and Stanfield 2000)

Oocyst counts (100 litre)	Percentage of samples
0	32.91
1	13.82
2	8.04
3–5	12.62
6–9	7.87
10–29	12.40
30*–50	4.14
51–100	3.47
101–200	2.17
201–1000	2.05
1001–5000	0.47
5001–16 458	0.02

* Arithmetic mean = 29.75 oocysts/100 litre.

(on taking the average) will underestimate the risk by more than fivefold. In the case of sets of just five samples, the means of 37% of the sets underestimated the true arithmetic mean by more than fivefold. This is why the *Cryptosporidium* Regulations in the UK require continuous monitoring of treated drinking water as it leaves the works (Gale and Stanfield, 2000).

4.1 Risk assessment complements epidemiological observations

Craun *et al.* (1998) reviewed oocyst count data collected during 12 waterborne outbreaks and concluded that there was no clear association between the risk of illness and the *Cryptosporidium* oocyst concentrations measured in the drinking-water supply. This would appear to challenge the very foundations of MRA for *Cryptosporidium* in drinking water. However, it can be explained by the fact that any association will tend to be masked by the large temporal-spatial heterogeneity in oocyst concentrations in drinking water because data from spot sampling in general underestimate the risk of cryptosporidiosis.

4.2 A zero oocyst reading provides little reassurance even during an outbreak

Haas and Rose (1995) proposed an action level of 10–30 oocysts/100 litre in finished water within and above which there is the possibility of a waterborne outbreak of cryptosporidiosis. Craun *et al.* (1998) dismissed this approach on the basis that many samples collected during outbreaks contained fewer counts than the action level. Indeed, in four of the 12 outbreaks studied, no oocysts were ever detected by sampling. According to the model for an outbreak in Table 16.5, a third of 100-litre samples contained zero oocysts, while a small proportion of 100-litre samples contained hundreds and even thousands of oocysts. Thus, a zero reading from a Poisson-log-normal distribution gives very little reassurance. Indeed, the heterogeneity explains why oocysts are often not detected during waterborne outbreaks of cryptosporidiosis. If, on the other hand, all the oocysts were evenly distributed in the treated water during an outbreak, then the oocyst concentration would equal the arithmetic mean (29.75 oocysts/100 litre) at all points in space and time. The probability of any 100-litre volume spot sample containing zero oocysts would be remote at 10^{-13} (assuming the Poisson distribution), and most would contain at least 19 oocysts, which is well within the proposed action level for an outbreak of cryptosporidiosis (Haas and Rose, 1995). A single spot sample would therefore be adequate to give a representative picture of the level of contamination and the risk of illness across the supply. A reading of 0 oocysts/100 litre would be very reassuring and representative of very low numbers of oocysts across the whole supply.

5 UNCERTAINTY – ESTIMATING THE REMOVAL OF PATHOGENS BY TREATMENT PROCESSES

Treatment removal efficiencies may vary greatly for a particular process due to 'between-batch' and 'within-batch' variation.

As shown in the section on the arithmetic mean exposures, p. 266, the arithmetic mean pathogen removal by treatment may be used directly in the risk assessment. Thus increasing the net pathogen removal by 10-fold will decrease the risk of infection by 10-fold. The problem is how to determine experimentally whether a treatment process has been improved by 10-fold.

5.1 Experimental designs may overestimate the net pathogen removal by treatment

Determining the arithmetic mean pathogen removal is complicated by the large variation in the treatment removal efficiencies, which gives rise to large variations in pathogen densities in the treated water (see Table 16.5). A key assumption of the *Cryptosporidium* outbreak model in Fig. 16.1 is that drinking water treatment not only removes oocysts but also increases the variation in the concentrations of those oocysts remaining in the treated water relative to the raw water. Therefore, monitoring programmes based on spot sampling will tend to underestimate the net pathogen loading in the treated water to a greater degree than in the raw water (Gale and Stanfield, 2000). The result of this is that, without intensive monitoring of the treated water samples, experiments to measure the net pathogen removal by treatment will tend to overestimate the arithmetic mean removal and hence underestimate the risk.

5.2 Modelling 'between-batch' or temporal variation

The variation in particle removal efficiencies by drinking water treated as used for the model in Fig. 16.1 are described by a log-normal distribution (see Fig. 16.2). Although particle removal ratios varied between 1 and 5 logs and the median (50 percentile) removal was 500-fold (2.7 logs), the arithmetic mean or net removal was 61-fold (1.79 logs). It is apparent that 50% of removals are more than eightfold more efficient than the arithmetic mean. Therefore half of the single 'one-off' experiments

undertaken to measure the removal efficiency will overestimate the arithmetic mean efficiency by more than eightfold. As an example, consider an experiment in which a single raw water sample is taken at Point A and a single treated water sample at Point B in Fig. 16.1. The result would suggest a >4 -log removal, when in fact the net removal is only 1.79-logs. This demonstrates the importance of understanding the variation in removal efficiencies when designing experiments to estimate the net removal by a treatment process.

Beta distributions are typically used to model the variation in the proportions of pathogens breaking through a treatment process (Teunis *et al.*, 1999). The beta distribution does not fit the data in Fig. 16.2 too well. Indeed, data for plate counts in drinking water support the application of the log-Normal distribution (Gale, 1996). Teunis *et al.* (1997) fitted a beta distribution to model the variation in removal efficiencies for spores of sulphite-reducing bacteria by drinking-water treatment. This is also plotted in Fig. 16.2. It is interesting to note how the log-normal and beta models differ at the higher removal efficiencies. Thus, according to the beta distribution, 83% of removals are greater >5 log, while for the log-normal distribution, only 0.5% are >5 log. Indeed, the beta distribution predicts 50% of removals are >18.5 logs. These high removal efficiencies have little effect on the magnitude of the arithmetic mean removal. Indeed, the log-normal and beta distributions fitted to the data in Fig. 16.2 have the same arithmetic mean of 1.79 logs. This is because, irrespective of whether a high removal value is 5 log (log-normal distribution) or 18.5 log (beta distribution), the proportion breaking through, i.e. 10^{-5} or $10^{-18.5}$ respectively, is so small as to tend to zero and has little impact on the arithmetic mean. However, the poor removal efficiencies (i.e. between 0 log and 1 log) have a great influence on the arithmetic mean removal (Gale and Stanfield, 2000; Gale, 2001b). Indeed, the beta distribution of Teunis *et al.* (1997) plotted in Fig. 16.2 has fewer poor removal efficiencies in the 0 to 1 log range, and consequently has a better net removal efficiency, the arithmetic mean being 2.2 logs.

5.3 'Within-batch' or spatial variation

The effect of drinking-water treatment on the statistical distribution of spores of aerobic spore-forming bacteria within 100-litre volume samples has been studied (Gale *et al.*, 1997). This can be considered as 'within-batch' or spatial variation. The overall conclusion was that operational drinking water treatment not only removed 94–98% of the spores, but also promoted the spatial association of the remaining spores. Further data from studies undertaken at pilot scale suggested a more complicated picture (Gale *et al.*, 2002). Thus, spore counts were not over-dispersed in all treated water volumes investigated and coliform counts were, in general, Poisson-distributed in the treated water volumes. Such findings contribute to explaining why no 'ideal' surrogate has been identified for treatment plant performance (Nieminski *et al.*, 2000; Gale *et al.*, 2002).

6 THE DEVELOPMENT AND APPLICATION OF EVENT TREES

Event trees simplify the process of modelling the various pathways and visualizing how infectivity in the source term partitions through the various routes to the receptors.

6.1 *Escherichia coli* O157 in sewage sludge

6.1.1 Source term

Gale (2001b) presented an event tree to model partitioning of salmonellae into sewage sludge at a sewage treatment works based on the plentiful data available for salmonella concentrations in raw sewage. There is currently little information on *E. coli* O157 concentrations in raw sewage or in sewage sludge. An alternative approach is to estimate the arithmetic mean pathogen loading in the sewage sludge, as developed by Gale and Stanfield (2001) for BSE in sewage sludge. The main source of *E. coli* O157 in raw sewage is from slaughter of cattle and sheep at abattoirs. The model uses UK Government data that 3.13 million cattle and 15.86 million sheep are slaughtered annually at abattoirs in England and Wales and assumes, on

the basis of expert advice, that 5% of faecal material in animals slaughtered at abattoirs enters the sewage treatment works, either directly into sewers or by tankering. Shere *et al.* (1998) report counts of *E. coli* O157:H7 in infected cattle ranging from 200 to 87 000/g. Since the arithmetic mean is nearer to the maximum in magnitude for a log-normal distribution (see Fig. 16.1), this value of 87 000 cfu/g is used as an estimate of the arithmetic mean for the MRA. Assuming 15.7% of cattle are infected (Chapman and Siddons, 1997) and that each bovine contains 10 kg of faeces at time of slaughter, then 2.1×10^{13} *E. coli* O157/year enter sewage works across England and Wales from the slaughter of cattle. The arithmetic mean *E. coli* O157 concentration in faeces from sheep and lambs during the New Deer (Scotland) outbreak may be estimated from data of Strachan *et al.*, (2001) as 365 500/g. (This assumes that the count of $> 10^6$ /g recorded in one lamb was 10^7 /g.) Assuming 2.2% of sheep flocks are infected (Chapman *et al.*, 1997) and each sheep has 1 kg of faeces at point of slaughter, then the arithmetic mean *E. coli* O157 loading across sewage works in England and Wales from slaughter of sheep in abattoirs is 6.3×10^{12} cfu/year. In England and Wales, 967 000 tonnes dry solids (tds) of sewage sludge are produced annually (Gale and Stanfield, 2001). On the basis of the salmonella model (Gale, 2001b), 82.9% of *E. coli* O157 in raw sewage would partition into the raw sludge at the works. Thus, the arithmetic mean loading from cattle and sheep combined in raw sewage sludge in England and Wales is 2.4×10^7 *E. coli* O157/t (dry solids). This is three orders of magnitude lower than the arithmetic mean of 2.0×10^{10} salmonellae/t (dry solids) of raw sewage sludge predicted by the salmonella model (Gale 2001b) based on monitoring data for raw sewage.

6.1.2 Pathway term

The pathway for transmission to root crops of *E. coli* O157 in sewage sludge applied to agricultural land is shown in Fig. 16.6. The model assumes that conventional sludge treatment (e.g. anaerobic digestion) destroys 2 logs

of *E. coli* O157 according to the UK Code of Practice for Agricultural Use of Sewage Sludge (Department of the Environment, 1996). Strachan *et al.*, (2001) allow for a T_{90} (time for a 1-log decay) of 16 days in a model for an environmental outbreak of *E. coli* O157, and Bolton *et al.* (1999) demonstrated a linear 5-log decay over 50 days on grazing land during the winter months. The model (Fig. 16.6) allows a 5-log decay, which is conservative for the 12-month time interval specified by the Safe Sludge Matrix (www.adas.co.uk/matrix/) between application of treated sludge and harvesting of root crops. Dilution in soil is modelled as the probability of a root crop colliding with a sludge particles compared to the much greater probability of colliding with a soil particle (Gale and Stanfield, 2001). At point of harvest, 0.02 tonnes of soil/sludge is transferred to each tonne of root crops. This is the origin of the 0.02 in Fig. 16.6. Following the event tree through, the arithmetic mean incremental exposure to root crops at point of harvest is calculated as:

$$2.4 \times 10^7 \times 0.01 \times 10^{-5} \times 0.0018 \times 0.02 \\ = 8.3 \times 10^{-5} \text{ } E. coli \text{ O157/root crops.}$$

The salmonella model predicted 70 salmonellae/root crops at point of harvest from application of conventionally-treated sewage sludge, although this assumed only a 2-log decay in the soil (Gale, 2001b).

6.2 *Escherichia coli* O157 risk assessment for a rendering plant discharging treated effluent to a surface water

Gale (2001b) estimated the risks of *E. coli* O157 from a rendering plant discharging treated effluent to a river, which is used for recreational water sport activities (including water skiing) and also for municipal drinking water abstraction. The source term is based on the processing of 619 cattle carcasses/day with the same *E. coli* O157 loadings assumed above. The pathway from carcass to raw effluent is set out as an event tree in Fig. 16.7.

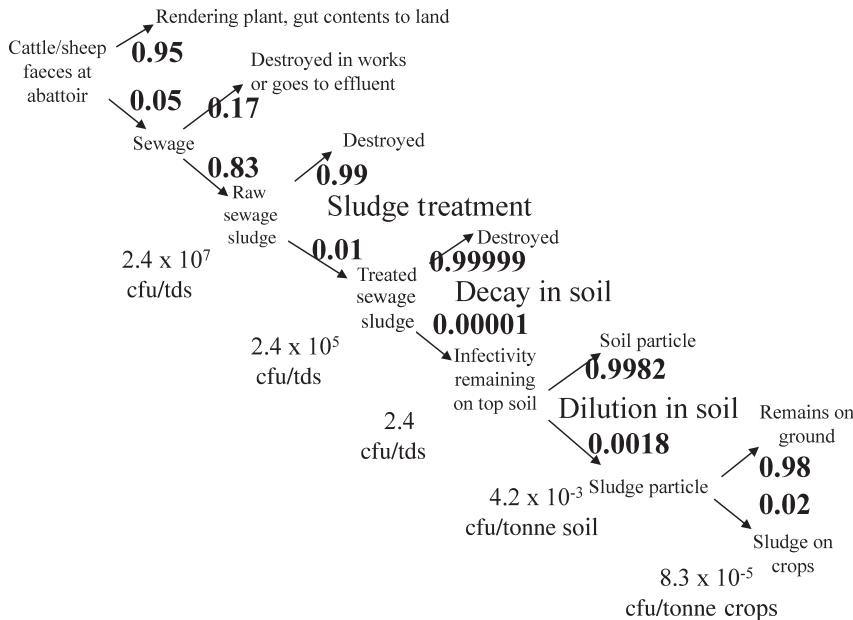


Fig. 16.6 Event tree for transmission of *E. coli* O157 in faeces at abattoirs to root crops via application of treated sewage sludge to agriculture land.

6.2.1 Uncertainty in the fate of the lorry washings has little impact

The model assumes that only 1% of faeces leaking out in the lorry goes to the raw effluent in the form of lorry/yard washings. This may be rather optimistic. A 'What if?' scenario is modelled in Fig. 16.7 to assess the impact of 10% of the faeces leaking out in the lorry going to the raw effluent as washings. The total loading of *E. coli* O157 in the raw effluent increases from 6.5×10^8 to 11.7×10^8 /day. Therefore the risk is not even doubled.

6.2.2 By-pass of treatment has a huge impact

The raw effluent is treated on site prior to discharge to the river. The net removal of *E. coli* O157 by treatment (which includes DAF, activated sludge treatment and UV disinfection) is 4.6 logs, i.e. only 0.0025% of the *E. coli* in the raw effluent breakthrough treatment. However, if there is a 1% by-pass of treatment such that for 3.6 days of the year the effluent receives no treatment, then the net removal is reduced to just 1.999-logs, i.e. 1.002% of *E. coli*

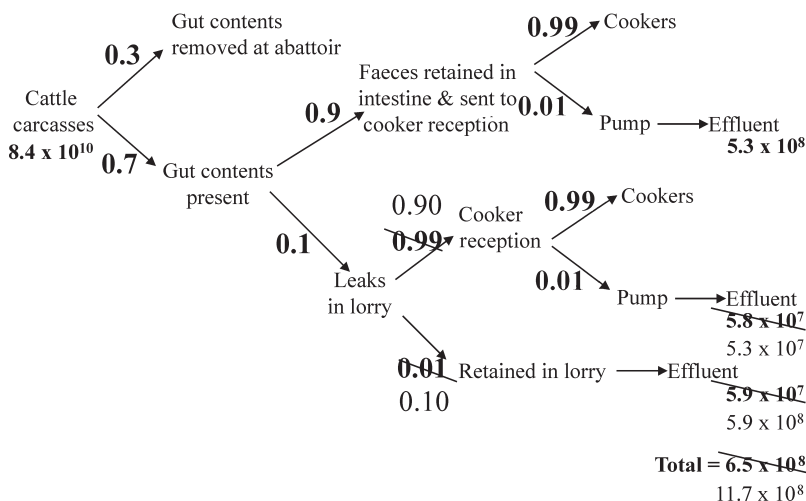


Fig. 16.7 Event tree for modelling partition of *E. coli* O157 in cattle carcasses into the effluent each day at a rendering plant. The model demonstrates a 'What If?' analysis.

break-through. This increases the *E. coli* O157 loading in the treated effluent, and hence the risk, by 400-fold.

7 BSE IN RENDERING PLANT EFFLUENTS

Gale *et al.*, (1998) estimated the risks of BSE transmission to consumers' drinking water from a chalk aquifer above which a rendering plant discharges treated effluent. The rendering plant processes 286 cattle carcasses/day. Assuming the human oral ID₅₀ is 1 g of BSE-infected bovine brain and that 0.54% of the cattle are in the final stages of infection, it may be calculated that 1544 human oral ID₅₀ enter the plant, on average, daily. An event tree to model breakthrough of BSE infectivity into the aquifer is presented in Fig. 16.8. Rendering destroys 98% of the BSE infectivity (Taylor *et al.* 1995) and 0.6% of the meat and bone meal (MBM) product breaks through into the condensate as suspended solid. The effluent treatment processes include DAF and activated sludge, which together remove 99.7% of particulate material, and hence any BSE agent. The model allows for a 93% decay of BSE agent in the soil (over a period of 2 years) and a 90%

attenuation in the chalk substrata. The rendering plant produces 120 000 litre of effluent/day, which is diluted into 10 Ml of groundwater in the aquifer. Dilution is modelled in the event tree as the probability of a consumer ingesting effluent compared to the much greater probability of ingesting (uncontaminated) ground water. The concentration of BSE in the drinking water is calculated from Fig. 16.8 as 4.7×10^{-8} ID₅₀ in 120 000 litre. A person ingesting 2 litre of tap water/day for a year would therefore ingest 2.8×10^{-10} ID₅₀. Assuming a negative exponential dose-response curve (Gale, 1998), this translates into a risk of 2.0×10^{-10} /person/year.

7.1 'What if' 1% of the raw material by-passes the rendering process?

According to the model in Fig. 16.8, the rendering process not only destroys 98% of the BSE agent but also partitions 99.4% of the remaining infectivity away from the condensate and into the MBM product. The net break-through of BSE agent into the condensate from the rendering process is therefore calculated as $0.02 \times 0.006 = 0.00012$. According to the *E. coli* O157 model in Fig. 16.7, some 1% of the raw material in the cooker reception by-passes

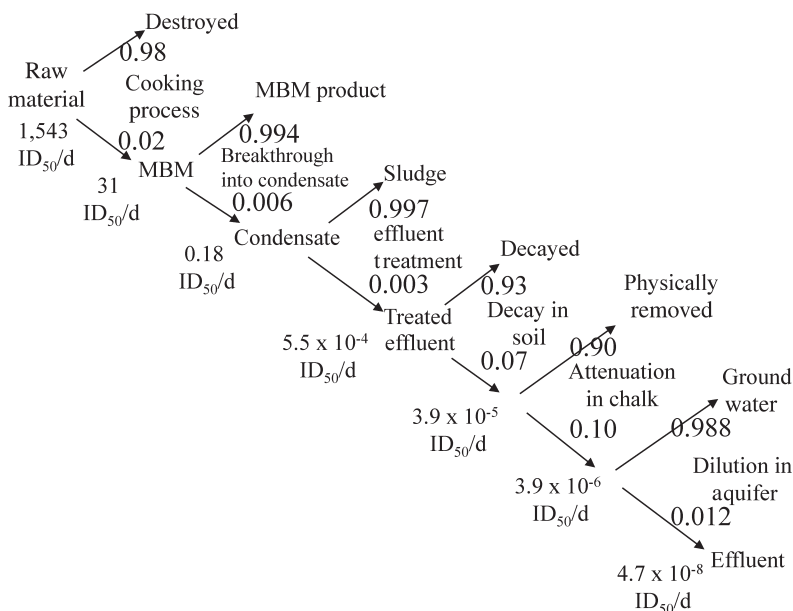


Fig. 16.8 Event tree for modelling the partitioning of BSE infectivity from cattle at a rendering plant into an aquifer (based on Gale *et al.*, 1998).

the cookers and is pumped to the raw effluent. It is extremely unlikely that as much as 1% of the brain and spinal cord from each bovine bypasses the cookers because the head is removed intact and the carcass is no longer butchered up the spinal column at abattoirs (see Gale and Stanfield 2001). However, skulls may be cracked by the Archimedes screws which lift the material from the sump to the cookers. By-pass of 1% of the raw brain/spinal cord from each bovine would mean that the net breakthrough of rendering would be $0.01 \times 1.0 + 0.99 \times 0.00012 = 0.01012$. Thus, the net breakthrough of rendering would be increased 84-fold from 0.00012 to 0.01012. The risks to drinking-water consumers would also be increased 84-fold, to 1.6×10^{-8} /person/year.

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Health constraints on the agricultural recycling of wastewater sludges

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1 INTRODUCTION

The objective of sewage treatment is to remove solids and to reduce its biochemical oxygen demand (BOD) before returning the treated wastewater to the environment. Sewage sludge, increasingly referred to as biosolids, is an inevitable product of wastewater treatment. Conventional wastewater treatment processes comprise separate process streams for the liquid and solid fractions (sludge). The overall aim of treatment (in terms of solids) is: (i) to reduce to the minimum the amount of solids in the treated effluent in order to achieve discharge standards; and (ii) maximize the level of solids in the sludge in order to minimize the volume requiring further treatment and disposal.

Sludge is produced at various stages within the wastewater treatment process (Fig. 17.1). Usually, these solids are combined and treated as a whole. Dedicated sludge treatment may not be available at all works, particularly smaller plants. In these circumstances it is normal practice to transport the sludge to a larger works for subsequent treatment.

2 BASIC PRINCIPLES OF WASTEWATER TREATMENT

2.1 Preliminary treatment

Preliminary treatment consists of screening through bar screens to remove coarse solids

and buoyant materials, such as plastics or rags, which may become trapped in pumps or other mechanical plant. The screenings are usually removed from the process stream and disposed of separately by landfilling or incineration. Occasionally, screenings may be shredded (comminuted) to reduce their size and returned to the process stream. This may cause problems with downstream processes and, as a consequence, is rarely practised at works which incorporate secondary (biological) treatment. The other component of preliminary treatment is grit removal, which is accomplished in chambers (or channels) or by centrifugation, taking advantage of the greater settling velocities of these solids. The material is largely inorganic in nature and is usually disposed of to landfill.

2.2 Primary treatment

Primary treatment is designed to reduce the load on subsequent biological (secondary) treatment processes. Although the design of primary sedimentation tanks differs, they achieve the removal of settleable solids, oils, fats and other floating material, and a proportion of the organic load. Efficiently designed and operated primary treatment processes should remove 50–70% of suspended solids and 25–40% of the BOD (organic

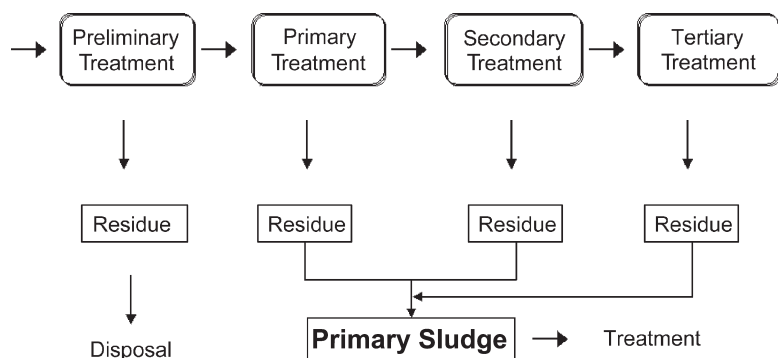


Fig. 17.1 Schematic representation of wastewater treatment showing sludge production.

load). The separated solids have a high organic content and are usually treated in order to stabilize the material prior to disposal.

2.3 Secondary treatment

Biological processes are used to convert dissolved biodegradable organic substances and colloidal material into inorganics and biological solids (biomass). There are several secondary treatment processes, but these may be divided into fixed film (e.g. trickling filters, rotating biological contactors) and suspended processes (e.g. activated sludge). Solids separation is the final stage of many of these treatment systems, which produces a sludge, the nature of which will depend on the upstream treatment process. Sludges arising from secondary treatment are usually combined with primary sludge and treated as a whole.

2.4 Tertiary treatment

Tertiary treatment will only be required for treatment works subject to specific discharge conditions. Many of these processes will not produce sludge and those that do are likely to generate only small amounts requiring dedicated treatment. Solids removed by granular media filtration will be passed to the primary sludge treatment process.

3 SLUDGE TREATMENT

The sludge obtained from the various stages is usually in the form of a liquid containing

between 0.5 and 6% dry solids. The typical composition of raw (untreated) and anaerobically digested sludge is shown in Table 17.1. The nature and extent of any treatment depends on the means of final disposal or beneficial use. The aim of treatment is to reduce the water and organic content of the sludge and render it suitable for disposal or reuse. There are several commonly used methods of sludge treatment, including long-term storage (in lagoons), lime stabilization, digestion (aerobic or anaerobic), air drying, thermal drying, and incineration or gasification (for energy recovery). Detailed descriptions of these processes are outside the scope of this handbook. However, the effect of the various treatments on pathogens is described below.

Sewage sludge contains valuable amounts of plant nutrients (nitrogen and phosphorus) and trace elements (Table 17.2). For this reason sludge has historically been applied to agricultural land as part of an integrated farm management plan. Other options for disposal include energy recovery and land reclamation activities. In Europe, North America and elsewhere, the disposal of sewage sludge is subject to strict controls designed to protect soil quality while encouraging the use of sludge in agriculture. Codes of Practice, such as those published by the UK Department of the Environment (DoE, 1996) and the UK Ministry of Agriculture, Fisheries and Food (MAFF, 1998a,b), provide advice on practical aspects of utilizing sewage sludge in agriculture.

Strict limits are set on the amounts of potentially toxic elements permitted in sludge which may be used in agriculture. Application

TABLE 17.1 Chemical composition and properties of untreated and digested sludge

<i>Constituent</i>	<i>Untreated primary sludge</i>		<i>Digested primary sludge</i>	
	<i>Range</i>	<i>Typical</i>	<i>Range</i>	<i>Typical</i>
Total dry solids (TS) %	2.0–8.0	5.0	6.0–12.0	10.0
Volatile solids (% of TS)	60–80	65	30–60	40
Grease and fats (% of TS)				
Ether soluble	6–30		5–20	18
Ether extract	7–35			
Protein (% of TS)	20–30	25	15–20	18
Nitrogen (N, % of TS)	1.5–4	2.5	1.6–6.0	3.0
Phosphorus (P ₂ O ₅ , % of TS)	0.8–2.8	1.6	1.5–4.0	2.5
Potassium (K ₂ O, % of TS)	0.0–0.1	0.4	0.0–3.0	1.0
Cellulose (% of TS)	8.0–15.0	10.0	8.8–15.0	10.0
Iron (not as sulphide)	2.0–4.0	2.5	3.0–8.0	4.0
Silica (SiO ₂ , % of TS)	15.0–20.0		10.0–20.0	
pH	5.0–8.0	6.0	6.5–7.5	7.0
Alkalinity (mg/l as CaCO ₃)	500–1500	600	2500–3500	3000
Organic acids (mg/l as HAc)	200–2000	500	100–600	200
Energy content (Kj/kg)	23 000–29 000	25 500	9300–14 000	11 500

Source: Metcalf and Eddy (1991).

rates are controlled to minimize the accumulation in the soil of toxic metals. Due to the low levels of metals in sludge, application rates are governed in practice by maximum nitrogen application rates (250 kg/ha y⁻¹ or 500 kg/ha y⁻¹) and to balance phosphorus addition with crop off-take.

Information on the amounts of sewage sludge produced and its disposal is collected by a number of countries, principally the USA and the member states of the European Union. Annual sludge production in the USA is in the region of 6.8 million tonnes dry solids (M tds) of which 54% is applied to land (Bastian, 1997). The figures for the EU are 5.1 M tds and 48% respectively (CEC, 1999). Within the EU amounts of sludge produced vary considerably, with Germany producing the largest amount of treated sludge followed by the

UK and France (Fig. 17.2). The proportion of treated sludge used in agriculture varies across the European Union, with just over 10% of sludge production in Ireland being applied to land compared with 66% in France (Fig. 17.3). Factors affecting the amount of sludge applied to agricultural land include topography, land use, climatic conditions, and the availability of alternative means of disposal. In the UK, sludge production is increasing, principally as a result of the EU Directive on the treatment of urban wastewater (CEC, 1991). The cessation of sea disposal has resulted in a greater proportion of sludge being used in agriculture (Table 17.3), a trend which is projected to continue in the medium term (Fig. 17.4).

4 REGULATIONS GOVERNING THE USE OF SLUDGE IN AGRICULTURE

4.1 USA

The treatment and ultimate disposal of sewage sludge, including domestic septage, derived from the treatment of domestic sewage, is governed by 40 CFR Part 503 rule (US EPA, 1993). The regulations were developed over

TABLE 17.2 Nutrient content of sewage sludge (% dry weight)

<i>Constituent</i>	<i>Range</i>	<i>Typical</i>
Nitrogen	<0.1–17.6	3.0
Phosphorus	<0.1–14.3	1.5
Sulphur	0.6–1.5	1.0
Potassium	0.02–2.6	0.3

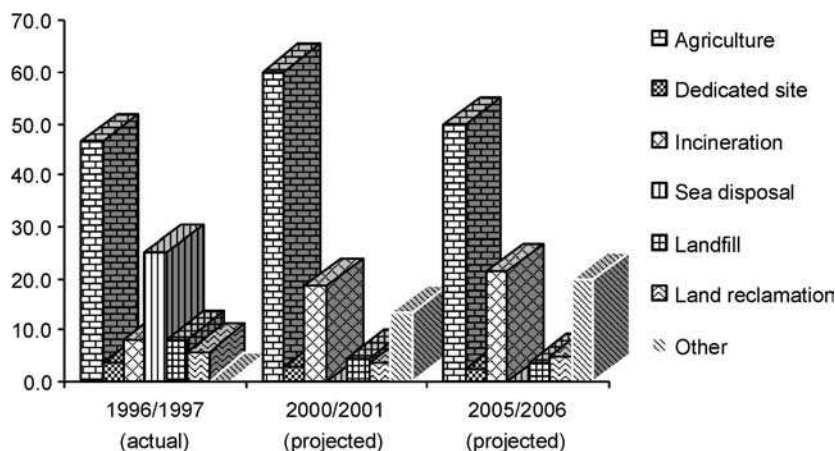
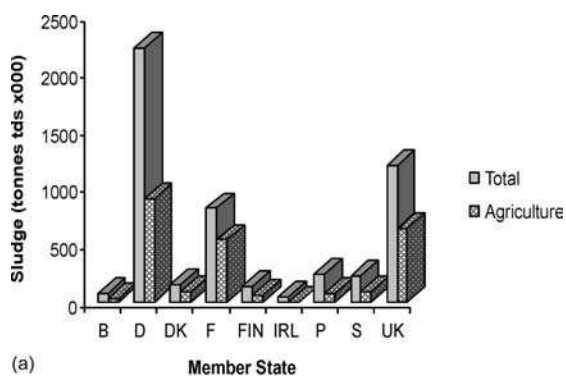
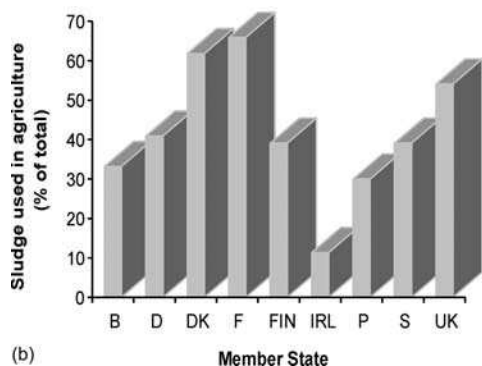


Fig. 17.2 Sludge disposal routes in the UK (Environment Agency, 1999).

a number of years and are designed for uniform application across the country over a wide range of geographic and climatic conditions, from Alaska to Hawaii. For this reason



(a)



(b)

Fig. 17.3 (a) Sludge production within the European Union and amounts recycled to agricultural land; (b) Proportion of EU sludge recycled to agricultural land.

the regulations are extensive and complex. Only treated sludge is permitted to be applied to land of any type. Sludges are categorized as Class A or Class B depending on the level of treatment intended to reduce pathogens.

In order to attain Class A status sludges must have been treated by 'a process to further reduce pathogens' (PFRP). Such a process is considered capable of reducing the number of pathogens to those normally present in the soil. Provided that the treated sludge complies with end product microbiological standards (Table 17.4), it may be applied without restriction to a wide range of land types, including that intended for agricultural or horticultural

TABLE 17.3 Sludge disposal outlets in the UK

Outlet	Quantity (%) ($\text{tds/y}^{-1} \times 10^3$)	
	1990/91	1996/97
Agriculture	465 (42)	520 (47)
Dedicated site	25 (2)	39 (3)
Sea disposal	334 (30)	280 (25)
Incineration	77 (7)	91 (8)
Landfill	88 (8)	91 (8)
Land reclamation		64 (6)
Forestry		1 (<1)
Horticultural compost		13 (1)
Storage (on site)	50 (5)	15 (1)
Other	68 (6)*	1 (<1)
Total	1107 (100)	1115 (100)

* More general category of 'Beneficial' used which included activities classified separately in 1996/7 survey.

Source: WRc (1998).

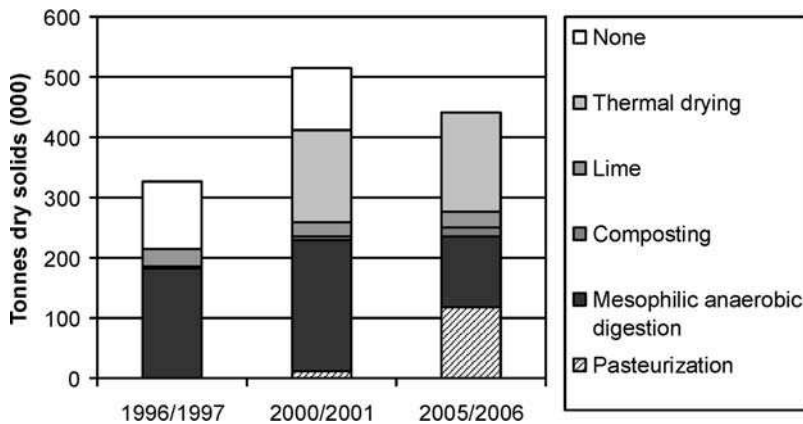


Fig. 17.4 Sludge treatment processes used in the UK.

use. Class B sludges are required to have been treated by 'a process to significantly reduce pathogens' (PSRP). With restrictions, Class B sludge may be applied to agricultural land. The sludge is required to meet end product microbiological standards (Table 17.4). The restrictions on application are:

- No grazing or harvesting of fodder crops with 30 days (of application)
- No harvesting of crops grown above ground within 14 months
- No harvesting of crops grown below ground for 20 months if the sludge remains on the soil for 4 months or longer; 38 months if the sludge remains on the soil for less than 4 months
- No harvesting of turf within 12 months
- No public access within 12 months (parks, playing fields etc.)

The implicit goal of the requirements for Class A biosolids is to reduce the number of

TABLE 17.4 End product microbiological standards for Class A and Class B sludges (US EPA, 1993)

Standard	Class A	Class B
Faecal coliforms/g ds	Less than 1000	Less than 2 000 000*
Salmonellae 4/g ds	Less than 3	
Enteroviruses pfu 4/g ds	Less than 1	
Parasite ova 4/g ds	Less than 1	

ds Dry solids.

* Geometric mean of seven samples.

pathogens in sewage sludge to below the level of detection (<3 MPN salmonella, <1 PFU enteric viruses, and <1 viable helminth ova – all per 4-g dry weight). The goal for the production of Class B biosolids is the reduction in the number of pathogens to levels that are unlikely to pose a public health risk (US EPA, 1999).

4.2 European Union

The controls on the application of sewage sludge to agricultural land within member states derive from Council Directive 86/278/EEC published in 1986 for implementation within 3 years (CEC, 1986). The principal rationale of the Directive was to minimize the accumulation in the soil of heavy metals or other potential toxic elements (PTE) with the objective of protecting soil fertility and public health. However, the Directive included measures for controlling transmissible disease by introducing constraints on the use of sludge. Article 7 of the Directive requires Member States to prohibit the use of sludge or the supply of sludge for use on:

- grassland or forage crops if the grassland is to be grazed or the forage crops to be harvested before a certain period has elapsed. This period, which shall be set by the Member States, taking particular account of their geographical and climatic situation, shall under no circumstances be less than 3 weeks:

TABLE 17.5 Examples of effective sludge treatment processes as defined in the UK Code of Practice

Process	Conditions
Pasteurization	Minimum 30 min at 70°C; or Minimum 4 h at 55°C
Mesophilic anaerobic digestion	Followed in all cases by mesophilic anaerobic digestion Mean retention of at least 12 days at 35°C ± 3°C; or Mean retention of at least 20 days at 25°C ± 3°C. Followed in each case by secondary digestion with a mean retention time period of at least 14 days
Thermophilic aerobic digestion	Mean retention of at least 7 days. All sludge to be subjected to a minimum of 55°C for at least 4 h
Composting (windrows or aerated piles)	Compost must be retained at 40°C for at least 5 days including a period of 4 h at a minimum of 55°C. Followed by a period of maturation
Alkaline stabilization (with lime)	pH to be 12 or greater for a period of at least 2 h
Liquid storage	Storage for at least 3 months.
Dewatering and storage	Dewatering and storage for at least 3 months. Storage at least 14 days if sludge previously subjected to mesophilic anaerobic digestion

Source: DoE (1966).

- soil in which fruit and vegetables are growing, with the exception of fruit trees
- ground intended for the cultivation of fruit and vegetable crops which are normally in direct contact with the soil and normally eaten raw, for a period of 10 months preceding the harvest of crops and during the harvest itself
- sludge shall be treated before being used in agriculture¹. Member States may nevertheless authorize, under conditions laid down by them, the use of untreated sludge if it is injected or worked into the soil.

In the UK, The Sludge (Use in Agriculture) Regulations 1989 directly implement the provisions of the Directive (Anon, 1989). This was accompanied by a Code of Practice (DoE, 1996) which provided practical guidance on how the requirements of the Directive could be met. It recognizes that pathogens may be present in untreated sludges and that their numbers can be reduced significantly by appropriate treatment. Examples of effective treatment processes are given in the Code (Table 17.5).

¹ Treated sludge is defined in Article 2(b) of the Directive as 'sludge which has undergone biological, chemical or heat treatment, long-term storage or any other appropriate process so as significantly to reduce its fermentability and other health hazards resulting from its use.'

At the time that the Code was prepared the pathogens of concern were considered to be salmonellae, *Taenia saginata* (human beef tapeworm), potato cyst nematodes (*Globodera pallida* and *Globodera rostochiensis*) and viruses.

The guidance was based on the concept of multiple barriers to the prevention of transmission of pathogens when sludge was applied to agricultural land. The barriers are:

- Sludge treatment, which will reduce pathogen content
- Restrictions on which crops may be grown on land to which sludge has been applied
- Minimum intervals before grazing or harvesting.

The scientific and public health principles which underpin this concept are valid. They recognize that for certain crops the risk of disease transmission is unacceptable, i.e. salad items which have a short growing period and which are to be consumed raw. For other crops the combination of treatment and a suitable period of no harvesting will result in the numbers of pathogenic microorganisms being reduced below a minimum infective dose (MID). The concept of MID is important – it relates to the number of organisms which must be ingested to cause

TABLE 17.6 Minimum infective dose (MID) for a range of gastrointestinal pathogens

Organism	Minimum infective dose
<i>Salmonella</i> spp.	10 ⁴ –10 ⁷
<i>Salmonella typhi</i>	10
<i>Escherichia coli</i> O157:H7	10–10 ²
<i>Vibrio cholerae</i>	10 ³
<i>Giardia intestinalis</i>	10–10 ²
<i>Cryptosporidium parvum</i>	10–10 ²
<i>Entamoeba histolytica</i>	10–10 ²
Hepatitis A virus	1–10 PFU

PFU, plaque forming unit

disease. It varies widely depending not only on the particular pathogen but also on the susceptibility of the host (Table 17.6). For example in the young, elderly, pregnant or those whose immunity is reduced the minimum number of organisms required to initiate disease is much smaller.

Despite the current concerns surrounding the risks to food safety, it is important to recognize that there have been no instances documented in which disease transmission to man or animals has occurred where the provisions of the relevant UK Regulations and Codes of Practice were followed.

5 PATHOGENS

Pathogens are microorganisms that are capable of causing disease in the host species (man, animals or plants). All the major groups of microorganisms contain species which are pathogenic including viruses (e.g. hepatitis virus), bacteria (e.g. salmonellae), fungi (e.g. *Aspergillus*), protozoa (e.g. *Cryptosporidium*) and helminths (e.g. *Taenia*). Although there are several plant pathogens which may potentially be present in sewage sludge (e.g. brown rot, potato root eelworm and beet rhizomania), this review will deal with pathogens affecting man and animals. Many of these are described as zoonotic, i.e. directly transmissible to man from animals. Examples of zoonotic infections include salmonellosis and cryptosporidiosis. This is a particularly important factor when considering the risks to human health arising from the use of sludge in agriculture.

The type and number of pathogens that are likely to be present in untreated sewage will depend on the inputs to the sewerage system. The spectrum of human pathogens will mirror the incidence of infection in the community. People suffering from diseases of the gastrointestinal tract will excrete large numbers of the pathogen in their faeces. Industrial sources of pathogens include meat processing plants, abattoirs and livestock facilities. The World Health Organization in its review of health risks arising from sewage sludge applied to land described a wide range of pathogens that could be present in sludge (WHO, 1981). This was subsequently updated and expanded by Strauch (1991) and the United States EPA who collated the data shown in Table 17.7 (USEPA, 1989).

The list is extremely comprehensive and, in reality, the risk from many of these microorganisms is very small. The organisms shown in bold are those identified by the US EPA as posing a significant risk to human health and which were taken into account in the development of the current Part 503 Regulations (US EPA, 1992). It is interesting to note that at that time they did not consider *Escherichia* as posing a significant risk to health. It is now known that certain shiga toxin-producing strains, such as *E. coli* O157², are capable of being transmitted by contaminated foodstuffs (Armstrong *et al.*, 1996; Tauxe, 1997; Mead and Griffin, 1998; Parry and Palmer, 2000).

In practice, the list of microorganisms that we need to be concerned with is relatively small. The pathogens of concern will vary from region to region depending on the nature and prevalence of endemic infectious intestinal disease within the indigenous population. For example, data for England and Wales collated by the PHLS Communicable Disease Surveillance Centre reveal that over half of all notified infections are due to *Campylobacter* (Fig. 17.5).

In contrast, intestinal parasites, particularly *Ascaris*, are a major disease burden in the developing world. These parasites form cysts or ova which are especially robust and resistant to environmental conditions, attributes which contribute to high levels of re-infection.

² Previously referred to as verotoxigenic *E. coli* or VTEC.

TABLE 17.7 Pathogenic microorganisms which may be present in sewage sludge

<i>Bacteria</i>	<i>Viruses</i>	<i>Protozoa</i>	<i>Yeasts</i>
Salmonella	Hepatitis A	Cryptosporidium	Candida
Shigella	Enteroviruses	Entamoeba	Cryptococcus
Yersinia	Poliovirus	Giardia	Trichosporon
Escherichia	Coxsackie viruses	Balantidium	
Pseudomonas	Echoviruses	Toxoplasma	Fungi
Clostridia	Rotavirus	Sarcocystis	Aspergillus
Bacillus	Adenovirus		Phialophora
Listeria	Reovirus	Cestodes	Geotrichum
Vibrio	Astrovirus	Taenia	Trichophyton
Mycobacterium	Calicivirus	Diphyllobothrium	Epidermophyton
Leptospira	Coronavirus	Echinococcus	
Campylobacter	Norwalk and Norwalk-like viruses		
Staphylococcus		Nematodes	
Streptococcus		Ascaris	
		Toxocara	
		Trichuris	
		Ancylostoma	
		Necator	
		hymenolepsis	

Source: US EPA (1999).

The literature contains numerous reports of surveys and investigations into the occurrence of pathogens and indicator bacteria in wastewater and sludge treatment plants. Because of methodological and geographical differences and, in some instances, lack of details regarding treatment processes, many of these data are not directly comparable. However, the data shown in Table 17.8 provide an indication of the likely numbers of indicators and pathogens in domestic wastewater and sludges.

6 EFFECTS OF SLUDGE TREATMENT ON PATHOGENS

It must be recognized at the outset that treatment is designed to stabilize sewage

sludge and reduce its putrescence. Pathogens may be inactivated as a consequence of the particular treatment applied. It has not been normal practice to optimize sludge treatment processes for pathogen reduction. Indeed to do so may reduce the effectiveness of the stabilization process.

A review of the literature by Ward and colleagues (1984) showed that the range of pathogen inactivation reported was large, depending on the extent of the treatment process and variation between operating conditions, even for the same generic treatment process (Table 17.9).

There are limited data on the effect of some of these processes on certain pathogens. In practice, research in this area has been restricted to

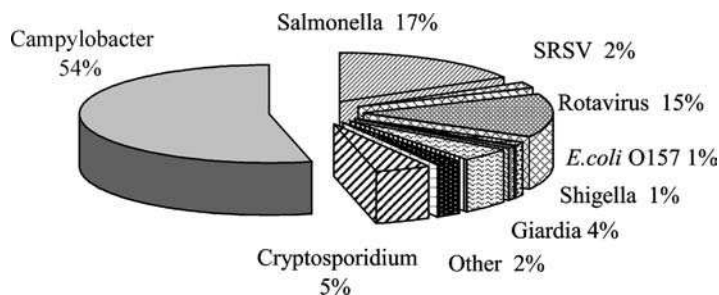


Fig. 17.5 Infectious intestinal disease in England and Wales by aetiological agent.

TABLE 17.8 Typical numbers of microorganisms found in various stages of wastewater and sludge treatment

Microorganism	Number per 100 ml			Number per gram		
	Crude sewage	Primary treatment	Secondary treatment	Tertiary treatment ^a	Raw	Treated ^b
Faecal coliforms	10 ⁸	10 ⁷	10 ⁶	<2	10 ⁷	10 ⁶
Salmonellae	10 ³	10 ²	10	<2	10 ³	10 ²
Shigella	10 ³	10 ²	1	<2	10 ²	3
Listeria	10 ⁴					10 ³
Campylobacter	10 ⁵				10 ⁴	10
Enteric virus	5 × 10 ⁴	1 × 10 ⁴	10 ³	0.002	10 ³	10 ²
Helminth ova	8 × 10 ²	10	0.08	<0.08	10	10
Giardia cysts	10 ⁴	5 × 10 ³	2.5 × 10 ³	3	10 ²	10

^a Including coagulation, sedimentation, filtration, disinfection.^b Mesophilic anaerobic digestion.Source: Jones *et al.* (1990); Metcalf and Eddy (1991); National Research Council (1996); De Luca *et al.* (1998); Watkins and Sleath (1981).

those pathogens with a high prevalence and likely to cause disease (e.g. salmonellae) and those that are more likely to exhibit resistance to the sludge treatment process (e.g. *Ascaris*). In comparison little is known about the pathogens which have only recently emerged as public health issues, most notably *E. coli* O157. The recent emergence of this pathogen means that there is little information available on the fate of *E. coli* O157 (and other shiga toxin-producing *E. coli* (STEC)) during the treatment of wastewater and sewage sludge. Despite the highly infectious nature of STEC and the presence of multiple virulence factors there is evidence that it is no more resistant to inactivation during sludge treatment than the indigenous populations of *E. coli* in sewage and sludge (Horan, personal communication).

TABLE 17.9 Summary of pathogen reduction during sludge treatment

Treatment	Log reduction		
	Bacteria	Viruses	Parasites
Mesophilic anaerobic digestion	0.5–4	0.5–2	0
Aerobic digestion	0.5–4	0.5–2	0
Composting	2–> 4	2–> 4	2–> 4
Air drying	0.5–4	0.5–> 4	0.5–> 4
Lime stabilization	2–> 4	> 4	0

Source: Ward *et al.* (1984)

The inactivation of indigenous *E. coli* in full-scale sludge treatment processes was investigated during a 3-month study, which looked at nine different sludge treatment processes at 35 sites in the UK (UKWIR, 1999). All of the processes surveyed reduced the numbers of *E. coli*. So-called ‘enhanced’ treatment processes, such as composting, lime addition and thermal drying, were capable of reducing numbers of *E. coli* to the detection limit of the analytical method. For all of these methods, over 90% of results showed bacterial reductions of 6 log or greater. Lagooning of sludge was capable of significantly reducing numbers of *E. coli* and, depending on the method of operation, reductions in the order of 5 log were observed. Mesophilic anaerobic digestion (MAD), the process carried out at the majority of sites surveyed, reduced numbers of *E. coli* by, on average, between 1.4 and 2.3 log depending on the solids content of the product. For sites producing a liquid product (2–4% ds) 78% of all reductions for were in the range 1 to 2 log. Where digested sludge was subsequently dewatered to produce a cake, 89% of results showed reductions in the range 2 to 4 log. The one vermiculture site in the survey showed results intermediate between MAD and the ‘enhanced’ treatment processes (Table 17.10).

Cryptosporidium parvum is another pathogen of increasing importance. Wastewater discharges and run off from agricultural land are

TABLE 17.10 Effect of sludge treatment processes on numbers of *E. coli*

Treatment	n	Log reduction in <i>E. coli</i>		Log reduction in <i>E. coli</i> treated sludge (100/g dry wt)	
		Mean	95%ile	Mean	95%ile
Lagooning	36	2.65	6.00	5.93	8.32
MAD, liquid	208	1.39	2.36	7.41	8.27
MAD, cake	93	2.29	3.64	6.65	7.46
Vermiculture	14	5.12	6.54	4.50	5.07
Composting	31	6.71	9.10	2.43	4.70
Lime addition	32	7.10	9.05	1.45	3.00
Thermal drying	70	7.14	8.90	1.67	3.56

Source: UKWIR (1999).

an important source of *Cryptosporidium* oocysts found in watersheds. The transmission of cryptosporidiosis is zoonotic and the possibility exists of foodborne infection arising from the use of sewage sludge in agriculture. Stadterman *et al.* (1995) found that a laboratory activated sludge plant removed 98.6% of seeded *Cryptosporidium parvum* oocysts. In a comparison of different treatment regimes, activated sludge and anaerobic digestion were found to be the most effective means of removing oocysts, the latter destroying 99.9% in 24 hours.

Studies of anaerobic mesophilic digestion under laboratory conditions showed that oocysts added to the contents of a digester operating at 35°C rapidly lost viability (as measured by excystation), decreasing to 17% after 3 days from an initial 81% viability (Whitmore and Robertson, 1995). Losses of viability in distilled water and anaerobic sludge at 35°C were similar, amounting to 90% after 18 days, indicating that the principal effect on viability was temperature. Oocysts exposed to mesophilic anaerobic digestion for 3 days and then stored for a further 14 days were completely inactivated. Aerobic digestion or pasteurization, both at 55°C, caused 92% loss of viability in 5 minutes. Thermophilic anaerobic digestion at 50°C resulted in complete inactivation within the first 24 hours (Whitmore and Robertson, 1995).

7 ROUTES OF TRANSMISSION

Recently expressed concerns over the use of sewage sludge in agriculture have focused on the risks to human health arising from the production of foods on land to which sludge has been applied (Anon, 1998a). A number of exposure pathways whereby foodstuffs become contaminated with pathogens can be envisaged. The exposure pathways relevant to sewage sludge in agricultural production are:

1. Sludge → soil → plant → human
2. Sludge → soil → animal → human
3. Sludge → soil → plant → animal → human
4. Sludge → soil → drinking water → human
5. Sludge → soil → irrigation water
→ plant → human.

These may be developed into a conceptual model which describes the framework for a microbiological risk assessment (Fig. 17.6). Other routes of exposure, which do not involve the food chain, can be identified. These include direct contact with sludge-treated soil or indirect contact via companion animals. The risk of exposure by this route (direct contact) is probably greatest among children.

It can be seen that there are several pathways that are unrelated to the use of sewage sludge, probably the most important of which is the application to land of organic wastes such as animal slurries and manures. The use of such materials in agriculture is less regulated than for sewage sludge and accounts for the majority of organic waste spread to land (Table 17.11). Despite this, the focus of attention has been on the human health risks via the food chain from the application of sewage sludge to agricultural land (RCEP, 1996; Anon, 1998a).

As previously mentioned, at the time that existing controls on the agricultural use of sewage sludge were being formulated, the pathogens of concern were salmonellae and *T. saginata*. In the intervening period, pathogens such as shiga toxin-producing *Escherichia coli* and *Cryptosporidium* have been recognized as important causes of intestinal infectious disease in humans. Any assessment of health

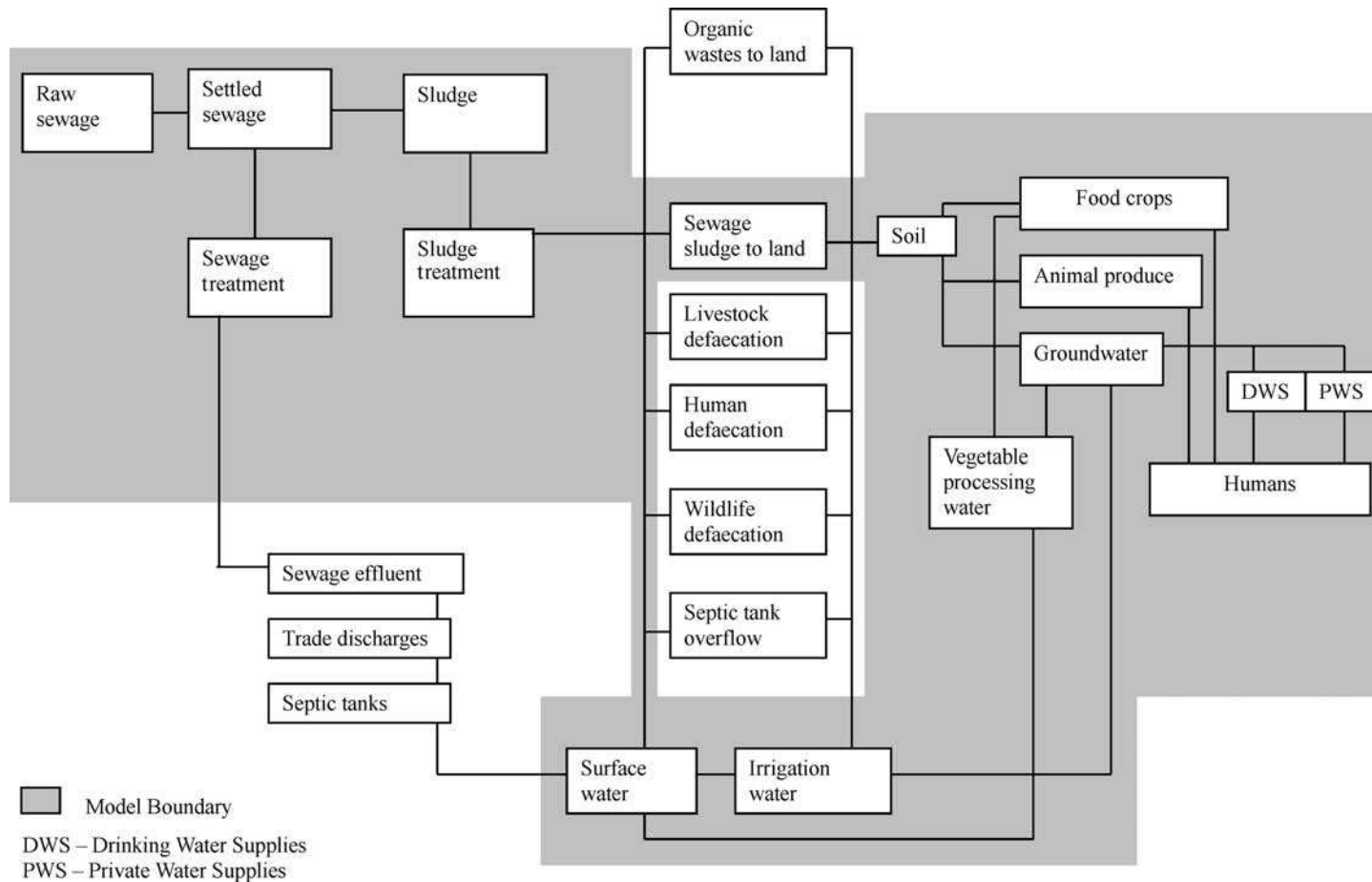


Fig. 17.6 Conceptual model for a microbiological risk assessment into the application of sludge to agricultural land (Pollard, personal communication).

TABLE 17.11 Estimates of the quantities of organic materials applied to land in the UK

Origin	Quantity (tonne $\times 10^3$ dw)
Farm animal	21 000
Sewage sludge	430
Paper industry	520
Food industry	600
Sugar industry	200
Others	150

dw: Dry weight.

Source: WRc (1998).

risks associated with the beneficial use of sewage sludge in agriculture must consider these two pathogens.

7.1 Shiga toxin-producing *E. coli*

Shiga toxin-producing *E. coli* (STEC) are now recognized as an important group of enteric pathogens. Although there are many serotypes capable of producing shiga toxins, *E. coli* O157:H7 is the most widely known. This organism was first described in 1982 following an outbreak of haemorrhagic colitis in the USA (Riley *et al.*, 1983). Outbreaks have been associated with the consumption of foodstuffs, drinking water, and swimming in natural surface waters. Zoonotic infections have also been reported. The majority of cases are believed to be foodborne, with an estimated 85% of cases ($n = 110\,220$) in the USA suspected to be food-related (Mead *et al.*, 1999).

There is very little information concerning the presence of STEC in sewage and sewage sludge. It is reasonable to assume that domestic sewage will contain STEC, if not continuously then intermittently, reflecting the incidence of infection in the community. The likelihood of STEC being present will be greater for those wastewater treatment works receiving wastes from animal-handling facilities, such as markets, abattoirs and meat processing plants. Surveillance of *E. coli* O157 in animals presented for slaughter carried out in northern England revealed that 15.7% of cattle, 2.2% of sheep and 0.4% of pigs were positive for the organism (Chapman, 2000).

Survival of *E. coli* O157 in the environment has been investigated by a number of workers. Maule (1997, 2000) showed that survival of the organism was found to be greatest in soil cores containing rooted grass. Under these conditions viable numbers were shown to decline from approximately 10^8 /g soil to between 10^6 and 10^7 /g soil after 130 days. When the organism was inoculated into cattle faeces it remained detectable at high levels for more than 50 days. In contrast, the organism survived much less readily in cattle slurry and river water where it fell in numbers from more than 10^6 /ml to undetectable levels in 10 and 27 days, respectively. Survival of *E. coli* O157:H7 in bulk manures may be prolonged, with the organism being detected for more than one year in static piles of ovine manure (Kudva *et al.*, 1998). Survival was reduced if the manure piles were aerated.

The fate of *E. coli* O157 present in animal slurry applied to pasture was investigated by Fenlon and colleagues (2000). Following application, numbers of both *E. coli* and *E. coli* O157 declined steadily with greater than 2 log reduction within 29 days. Relatively few cells (2% of total) were transported away from the soil surface and into the deeper layers of the soil. Run-off following heavy rainfall resulted in a loss from the soil of 7% of the *E. coli* applied in the slurry. A recent ecological study on predation of *E. coli* O157 by *Acanthamoeba polyphaga* has shown that the bacterial cells are capable of surviving and even replicating within this common environmental protozoan (Barker *et al.*, 1999). This may be important in the dissemination and survival of STEC within the environment.

7.2 *Cryptosporidium*

Human infection with *Cryptosporidium* was first reported in 1976 (Fayer *et al.*, 2000). It is now apparent that *Cryptosporidium* is a significant cause of infectious intestinal disease (IID), accounting for about 5% of IID in which the causative organism is identified. In England and Wales, during 1999, there were nearly 5000 laboratory confirmed cases of cryptosporidiosis ($n = 4759$) (Anon, 2000a); in the USA

the estimated number of cases is 300 000 of which 10% are believed to be foodborne (Mead *et al.*, 1999).

Oocysts of *C. parvum* are environmentally hardy and, under certain conditions, may remain viable for many months. Survival studies in microcosms containing untreated river water showed that oocysts are extremely persistent, the times for 10 log reductions in viability being 160 days at 15°C and 100 days at 5°C (Medema *et al.*, 1997). Investigations into the survival in soils treated with sewage sludge showed that viability declined by 20–40% at 20°C over 44 days. Temperature was the principal factor affecting oocyst survival (Whitmore and Robertson, 1995).

Little is known about the movement of oocysts through the soil. The most relevant data were reported by Mawdsley and colleagues (1996a) who studied the transport of *Cryptosporidium* oocysts through soil following the application of slurry to a poorly draining silt clay loam soil. Bovine slurry seeded with 5×10^9 *C. parvum* oocysts was applied to the surface of soil blocks (80 cm \times 56 cm \times 20 cm) removed from a perennial ryegrass ley at an application rate equivalent to 50 m³/ha. The blocks were irrigated 24 h following slurry application and periodically thereafter. Samples of run-off (at 4 cm depth) and leachate (at 20 cm depth) were collected and the number of oocysts enumerated. After 70 days the blocks were destructively sampled and examined for the presence of oocysts. Experiments were carried out in triplicate. Numbers of oocysts leaching from the blocks declined from 8.4×10^6 on day 1 to 2.3×10^4 at day 70. Oocysts levels were consistently lower in run-off compared with the leachate from the base of the soil blocks. Numbers fell below the limit of detection after 21 days and 28 days in two blocks, but were detectable in the third block for the duration of the experiment (70 days). These results suggest that oocysts tend not to become associated with soil particles, being either transported away in run-off or moving vertically downwards through the soil column. The majority of oocysts were retained in the top 2 cm of soil (Mawdsley *et al.*, 1996b).

8 ASSESSMENT OF HUMAN HEALTH RISKS

The process of microbiological risk assessment is now considered to comprise three phases: problem formulation, analysis, and risk characterization (ILSI, 2000). The analysis phase consists of two elements: characterization of exposure and characterization of human health effects (Fig. 17.7).

Characterization of exposure requires an evaluation of the interaction between the pathogen, the environment and the human population. Factors that need to be considered include the virulence of the pathogen, survival in the environment, route of infection, numbers of pathogen present, effectiveness of control/treatment processes, infectious dose, severity of illness and size of exposed population.

Unlike the field of chemical toxicology, microbiological risk assessment is in its infancy. There exist major gaps in our knowledge about the organisms of concern. This is particularly the case for the emerging pathogens such as STEC and *Cryptosporidium*. There have been attempts to assess the risks posed by *Cryptosporidium* in drinking water supplies (Gale, 1996, 1999; Haas, 2000). However, the foodborne route has not been modelled, probably because of the received view that cryptosporidiosis is primarily waterborne, despite evidence demonstrating the potential for foodborne transmission. *Cryptosporidium* oocysts have been found on the surface of fresh, raw vegetables obtained from retail markets (Ortega *et al.*, 1991; Monge *et al.*, 1996). In the UK, an outbreak of cryptosporidiosis affecting 50 school children was linked to the consumption of improperly pasteurized milk (Gelletli *et al.*, 1997). In the USA, outbreaks have been associated with the drinking fresh-pressed apple juice (non-alcoholic cider) (Millard *et al.*, 1994). Outbreaks of cryptosporidiosis associated with infected food-handlers demonstrate clearly the potential for significant foodborne transmission of *Cryptosporidium* (Besser-Wiek *et al.*, 1996; Quiroz *et al.*, 2000).

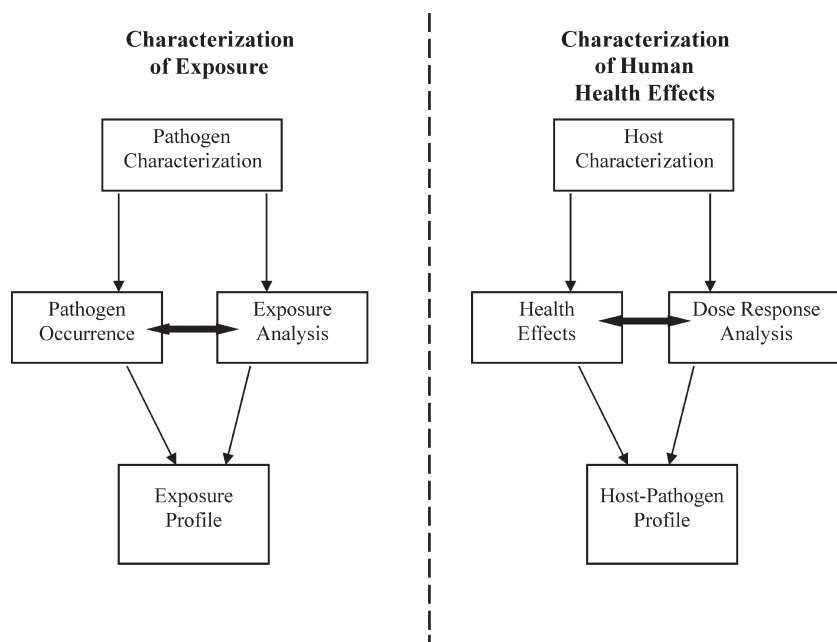


Fig. 17.7 Analysis phase of a microbiological risk assessment for foodborne pathogens (ILSI, 2000).

On the other hand, it is clear that STEC infection is primarily foodborne (Mead *et al.*, 1999). Data are required on the levels of STEC in treated sludge, their survival following land application and the potential for transfer to food crops before a microbiological risk assessment can be performed. Research is currently being undertaken to address these issues.

9 PRECAUTIONARY PRINCIPLE

Against a background of concern over methods of food production in the UK, the water industry, under the auspices of Water UK, and representatives of the food suppliers agreed a set of guidelines matching the level of sewage treatment with the crop under cultivation (Anon, 1998b).

The Safe Sludge Matrix (Anon, 2000b, Tables 17.12 and 17.13) forms the basis of the agreement and consists of a table of crop types, together with clear guidance on the minimum acceptable level of treatment for any sewage sludge (biosolids) based product, which may be applied to that crop or rotation. The agreement was driven by the

desire to ensure the highest possible standards of food safety and to provide a framework that gives the retailers and food industry confidence that sludge reuse on agricultural land is safe. The matrix enables farmers and

TABLE 17.12 The safe sludge matrix

Crop group	Untreated sludges	Treated sludges	Enhanced treated sludges
Fruit	X	X	✓ [†]
Salads	X	X	✓ [†]
		(30 month harvest interval applies)	
Vegetables	X	X	✓ [†]
		(12 month harvest interval applies)	
Horticulture	X	X	✓ [†]
Combinable and animal feed crops	X	✓	✓
Grass – grazing	X	X [†]	✓ [†]
		(Deep injected or ploughed down only)	
Grass – silage	X	✓ [†]	✓ [†]
Maize – silage	X	✓ [†]	✓ [†]

[†] 3 week no grazing and harvest interval applies.

[‡] 10 months harvest interval applies.

TABLE 17.13 Cropping categories within the Safe Sludge Matrix

<i>Fruit</i>	<i>Salad (e.g. ready to eat crops)</i>	<i>Vegetables</i>	<i>Horticulture</i>	<i>Combinable and animal feed crops</i>	<i>Grassland and maize</i>	
					<i>Silage</i>	<i>Grazing</i>
Top fruit (apples, pears, etc.)	Lettuce	Potatoes	Soil based glasshouse and polythene tunnel production (including tomatoes, cucumbers, peppers, etc.)	Wheat	Cut grass	Grass
Stone fruit (plums, cherries, etc.)	Radish	Leeks	Mushrooms	Barley	Cut maize	Forage Swedes/turnips
	Onions	Sweetcorn	Nursery stock and bulbs for export	Oats	Herbage	Fodder mangolds/beet/kale
	Beans (including runner, broad and dwarf French)	Brussels sprouts	Basic nursery stock	Rye	Seeds	Forage rye and triticale
Soft fruit (currants and berries)	Vining peas	Parsnips		Triticale		Turf production
Vines	Mange tout	Swedes/turnips	Seed potatoes for export	Field peas		
Hops	Cabbage	Marrows	Basic seed potatoes	Field beans		
Nuts	Cauliflower	Pumpkins	Basic seed production	Linseed/flax		
	Calabrese/broccoli	Squashes		Oilseed rape		
	Courgettes	Rhubarb		Hemp		
	Celery	Artichokes		Sunflower		
	Red beet			Borage		
	Carrots			Sugar beet		
	Herbs					
	Asparagus					
	Garlic					
	Shallot					
	Spinach					
	Chicory					
	Celeriac					

growers to continue to utilize the beneficial properties in sewage sludge as a valuable and cost effective source of nutrients and organic matter.

The main impact was the cessation of raw or untreated sewage sludge being used on agricultural land. As from the end of 1999, all untreated sludges have been banned from application to agricultural land used to grow food crops. Treated sludge³ can only be applied to grazed grassland where it is deep injected into the soil. The regulations require that there will be no grazing or harvesting within 3 weeks of application. Where grassland is reseeded, sludge must be ploughed down or deep injected into the soil.

More stringent requirements apply where sludge is applied to land growing vegetable crops and, in particular, those crops that may be eaten raw (e.g. salad crops). Treated sludge can be applied to agricultural land which is used to grow vegetables provided that at least 12 months has elapsed between application of the sludge and harvest of the vegetable crop. Where the crop is a salad, which might be eaten raw, the harvest interval must be at least 30 months. Where enhanced treated sludges⁴ are used, a 10-month harvest interval applies.

10 CONCLUSIONS

From an environmental perspective there is a persuasive argument that, of the disposal options available, recycling nutrients by means of applying sewage sludge to land, with appropriate safeguards, is the Best Practicable Environmental Option (BPEO) (CEC, 1986; RCEP, 1996). The risks to human, animal and plant health were taken into account when developing the current regulations and codes

³ There is a range of different treatment processes used to reduce the fermentability and possible health hazards associated with sewage sludge. These rely on biological, chemical or heat treatment. The most common form of treatment is anaerobic digestion.

⁴ Enhanced treatment, originally referred to as 'Advanced Treatment', is a term used to describe treatment processes which are capable of virtually eliminating any pathogens which may be present in the original sludge.

of practice. The fundamental principle (for reducing disease transmission risk) implicit to these controls on the use of sludge in agriculture is the concept of imposing multiple barriers to the recycling of pathogens from sludge to their hosts. The effectiveness of this approach is borne out in practice as noted by the Royal Commission on Environmental Pollution (RCEP, 1996) which concluded that, 'There are no instances in the UK in which a link has been established between the controlled application of sewage sludge and occurrence of disease in the general population through water or food contamination'. However, it is the case that the current controls predate the emergence of pathogens such as *Cryptosporidium* and STEC and may not sufficiently reduce the risk associated with these microorganisms.

More data are required on the numbers and fate of these emergent pathogens before a meaningful microbiological risk assessment can be carried out. Research is being undertaken in the UK, USA and elsewhere to generate these data. The likelihood is that the controls on the use of sewage sludge in agriculture will be strengthened as results of this research and in the light of public perception and expectations about food safety and environmental risks.

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Effluent discharge standards

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1 INTRODUCTION

From the Middle Ages until the early part of the nineteenth century the streets in many cities of Europe were fouled with excreta and garbage to the extent that people often held clove-studded oranges to their nostrils in order to tolerate the atmosphere. The introduction of water-carriage systems of sewage disposal in the early nineteenth century merely transferred the filth from the streets to the rivers. The problem was intensified by the Industrial Revolution with the establishment of factories on the banks of the rivers where water was freely available for power, process manufacturing and effluent disposal. As a consequence, the quality of most rivers deteriorated to the extent that they were unable to support aquatic life and some were little more than open sewers. In much of today's developed world there followed a period of slow recovery when many rivers were restored to balanced ecosystems with good fish stocks. However, in most of the developing world there has been no such recovery; urban watercourses still substantially act as open sewers and the position is often one of continuing deterioration, rather than marked improvement.

In countries where there has been recovery it has not been easy nor has it been cheap. It has been based on the application of good engineering supported by the passing and enforcement of necessary legislation and the development of suitable institutional capacity to finance, design, construct, maintain and operate the required sewerage and sewage treatment systems. Such institutional and technical systems not only include provisions

for the disposal of domestic sewage, but also for the treatment and disposal of industrial wastewaters and for the integrated management of river systems. A key element in all successful systems has been the establishment and enforcement of appropriate and affordable effluent discharge standards.

In the developing world, improvements to the current situation are often hampered by poor legislation, lack of appropriate institutional arrangements for regulation, lack of resources, insufficient finances and, frequently, lack of political will to enforce control measures. The position is complicated by the fact that many developing countries are at a loss on how to set standards and resort to importation of those from the developed world without either the means of achievement or the faintest idea of the costs of regulation and control. While no doubt well intentioned, the consequent problems exemplify the 'law of unintended consequences' (Cairncross, 1991) and result in slow or non-existent progress.

The objective of this chapter is to describe the general basis of setting standards and to discuss institutional arrangements for their regulation and enforcement. It is aimed primarily at discharges from domestic sewage treatment plants, including those accepting industrial effluents. It is not aimed at setting standards for the direct discharge of industrial effluents to rivers or to other watercourses, but it does give a brief account of the control of such effluents when discharged to public sewers and hence conveyed to a sewage treatment plant. A short historical review is followed by an overview of standard setting and a discussion on the problems

of implementation. There is also a discussion on the problems of regulating and controlling the private sector, especially where penalties are imposed for failure to comply with contractual standards, and where many of the problems are due to a lack of understanding of sampling procedures and the statistical nature of sewage effluents.

2 HISTORICAL REVIEW

Every country, whether designated as developed or developing, has its own historical perspective on the evolution of environmental control measures. Parallel progress in a number of developed world countries has led to different ways of describing environmental quality objectives and the standards deemed necessary to achieve them. It is impossible, but fortunately unnecessary, to describe each of these different ways: it is sufficient to outline some key developments, to discuss trends that have influenced standards over the years, and to describe significant differences in approach.

Much of the pioneering work was carried out in the UK (see Johnstone and Horan, 1996) where many watercourses had been devastated by the effects of the industrial revolution and had become a serious menace to public health, the extent of which was demonstrated in two great epidemics of cholera in 1866 and 1872. Following a number of studies and Commissions, the government of the day passed the Public Health Act 1875, which is regarded as one of the foremost sanitary measures of its time. Its prime importance was that it recognized, for the first time, that care of public health was a national responsibility. This Act was immediately followed by the Rivers Pollution Prevention Act 1876, which formed the basis of all legal action with regard to river pollution until 1951. A significant element of the 1876 Act was that it prohibited the discharge of solid or liquid sewage matter into a river and determined that *it was no defence to argue that the river had already been polluted upstream*. The significance of this

simple statement is profound and today forms a cornerstone of much of the world's environmental legislation.

Although the 1876 Act was comprehensive, it was in many ways ahead of its time. It specified duties with respect to sewage treatment at a time when such treatment was still in the development stage and many technical and scientific issues had not been resolved. In order to resolve these issues and determine a way forward, the government of the day set up a Royal Commission on Sewage Disposal. The Commission commenced its work in 1898 and completed the last of its nine voluminous reports 17 years later in 1915.

By far the most famous, and arguably the most important findings, were presented in the eighth report published in 1912. It dealt with the question of standards and tests to be applied to sewage and sewage effluents, and it introduced the BOD (biochemical oxygen demand) test. However, the first attempt at establishing effluent standards came in the fifth report published in 1908. Preliminary, empirical standards were proposed, based on an analysis of data from existing plants. Thus it was expected that 'a well-operated treatment works effluent':

1. should have no more than 30 mg/l suspended solids and
2. should, after filtering, absorb no more than:
 - 5 mg/l dissolved oxygen in 24 hours,
 - 10 mg/l dissolved oxygen in 48 hours, and
 - 15 mg/l dissolved oxygen in 5 days.

The eighth report was produced *inter alia* to ensure that money was not spent on sewage treatment when the circumstances of the area did not warrant it. This involved gathering data on local stream conditions (volume, depth, velocity and degree of aeration), together with chemical analyses of water quality. The report recommended that the old permanganate value be replaced by the BOD test (or dissolved oxygen absorption test, as it was then known). It recommended that the test be carried out at 65°F (18°C), which was the maximum river temperature during the hottest month in the UK. It also recommended that

a 5-day incubation period be used as this was found to give a smaller experimental error than tests carried out at shorter incubation periods. Contrary to popular belief, the 5-day period had nothing to do with the retention period of rivers in the UK.

In the course of the investigations, a number of rivers, and reaches of rivers, were classified as follows:

Very clean	1 mg/l BOD
Clean	2 mg/l BOD
Fairly clean	3 mg/l BOD
Doubtful	5 mg/l BOD
Bad	10 mg/l BOD

Although the Commissioners realized that it was logical to provide individual discharge standards for each sewage treatment works, depending on the nature and condition of the receiving water, this course of action was not adopted because of objections that:

- it would be difficult to administer
- it would place unequal burdens of purification between authorities discharging to the same watercourse
- it would allow a relaxation in discharge quality in some cases.

The final outcome was a general standard based on a mass-balance that assumed an eightfold dilution with river water of 2 mg/l BOD. It was felt that, if the river was not to deteriorate with serious consequences, the water after the discharge point should not absorb more than 4 mg/l BOD. On this basis, a mass balance shows that the sewage effluent should not absorb more than 20 mg/l BOD. Superficially, this would appear to have been a relaxation of the 15 mg/l standard provisionally set in the fifth report. However, the standard of 20 mg/l BOD was to be applied to an unfiltered sample as opposed to the filtered sample proposed in the fifth report.

The Commission retained the 30 mg/l value for suspended solids recommended in the fifth report and thus was born the famous

(or even infamous) Royal Commission 20/30 Standard still adopted today as a base standard in much of the world. Thus, by the year 1915, the Royal Commission on Sewage Disposal had established the first standards for effluent discharges and had introduced an elementary river classification system. In a separate report it also laid the basis for the control and disposal of industrial wastewaters.

Over the following 80 years there were many developments in procedures for setting effluent standards and determining environmental quality objectives. In the UK there was a series of, often fragmented, approaches to developing river classifications based on mathematical modelling and the use of river water quality objectives. River quality classification systems developed over a number of years and were changed as and when considered necessary. For example, as a result of a River Quality Survey in 1975, it was felt that the classification then in use should be replaced by a more objective one that would take account of potential uses of river water, and during a subsequent Review of Consent Conditions in 1978, there were moves towards linking effluent standards to the concept of water quality objectives. There were also technical changes, such as the introduction of the inhibited BOD test using allyl thiourea and ammoniacal nitrogen became an important standard for some rivers, such as the Thames, where there is widespread use of the river water for potable supply.

Improvements in the knowledge of sewage treatment processes meant that it was practical for standards to include limits on nitrogen and phosphorus and many countries introduced limits on bacteria, usually specified in terms of faecal coliforms. In 1989 the World Health Organization published health guidelines (Mara and Cairncross, 1989) for use of effluents in agriculture and aquaculture that specified limits on faecal coliforms and helminth eggs. More recently, WHO published a comprehensive set of guidelines on water quality, including those for sewage effluents; it deals, *inter alia*, with risk, risk management, economic

evaluation and practical aspects of implementation (WHO (2001) *Water Quality: Guidelines* (2001)).

Over the years there has been much debate over the advantages and disadvantages of establishing a system based on 'fixed emissions', where all discharges have to comply with one set of standards, and a system based on 'water quality objectives', in which an individual discharge has standards based on the quality needs of the receiving watercourse. Some countries have included a long list of parameters based on 'the precautionary principle', while the BATNEEC principle is used elsewhere, especially in conjunction with the philosophy of integrated pollution control (IPC). The BATNEEC principle recognizes the cost of environmental pollution control and attempts to choose mitigation measures using the 'best available technology not entailing excessive costs'.

For much of the twentieth century, discharge standards were based on numerical limits on chemical parameters, sometimes accompanied by some descriptive limits. In recent years there have been considerable advances on the use of toxicity testing and it is highly likely that it will become an increasingly useful tool in controlling discharges, especially from sources such as the chemical industry or from sewage treatment works that include complex industrial wastewaters. Control of individual chemicals by concentration does not consider the possible synergistic effects such that 'cocktails' may have on the aquatic environment, whereas toxicity testing provides an indication of potential damage. At present the application of toxicity techniques is in its infancy and suitable protocols and procedures need to be formulated.

Today it has become a fact of life that ever-increasing environmental standards have proliferated throughout the world (Johnstone and Horan, 1994). Even when the standards are superficially simple, such as in the European Directive on Urban Waste Water Treatment, their application can vary widely, leading to difficult comparisons and misleading interpretation, as illustrated by Jacobsen and Warn (1999). In the developed world, there has been

a major increase in the number of determinands included in effluent standards. Garber (1992) states that the wastewater from Los Angeles wastewater treatment plant in the USA has to be analysed for 140 priority pollutants before it is discharged to sea. Many of the limits are based on a precautionary approach and the achievement of 'zero risk'.

Unfortunately, many of the very high standards are very expensive to achieve and some have been imposed without any logical attempt at a cost-benefit analysis. It is even more unfortunate that many such standards, set for the developed world, have been adopted by some developing countries without consideration of the cost of implementation or the costs or the means of enforcement and regulation. Many developing countries now find that they cannot tackle problems of gross pollution in a phased and affordable way, due to the adoption of inappropriate standards, a subject that is discussed in greater detail later.

3 SETTING STANDARDS

Essentially there are two approaches to setting standards, although there are various options within each. One approach uses Fixed Emission Standards (FES) that are applicable to all discharges, while the other sets Water Quality Standards (WQS) to achieve some predetermined Water Quality Objective (WQO). Water quality standards therefore vary from discharge to discharge depending on the particular objective. There are advantages and disadvantages to both.

A number of issues need to be considered when setting standards, irrespective of the general approach used and whose understanding is important when designing treatment facilities. The most important are as follows:

- the chemical or other intensive parameters to be used as control measures
- the average and maximum daily flows
- the instantaneous rate of flow
- the possibility of using mathematical models

- whether to use concentration or minimum percentage removal
- whether the sampling regime is to be based on 'snap' samples or 'composite' samples
- errors in sampling and analysis
- whether standards are to be based on absolute values or on statistical compliance
- the period of judgement of compliance. Whether it is to be, for example, a single day's sample, a monthly average, a yearly average, or on a percentile value measured over a certain period
- the required frequency of sampling and analysis, for example daily, weekly, monthly, etc.
- the contractual nature of the 'permit', the 'licence' or the 'consent' that encompasses the standards
- the legal and regulatory basis of the 'permit', the 'licence' or the 'consent'
- penalties for failure to achieve the standards
- the cost and affordability of achieving and regulating the standards
- the possibility of phasing standards to suit affordability
- the need to control industrial wastewaters that discharge to the public sewer.

3.1 Fixed emission standards (FESs)

Fixed emission standards are relatively simple to apply and to regulate and can provide an effective way of achieving basic environmental objectives. They can be used as a first step in a phased approach to achieving some long-term environmental objective. They are particularly useful in countries where there is virtually no sewage treatment and where it is difficult or impossible to afford strict standards in the short term. Fixed emission standards usually take either the form of a percentage removal of one or a number of contaminants, or limits based on concentration.

There are some dangers in using this approach if the standards are not set appropriately. If they are set too low, there is a danger that they might not achieve noticeable improvement in the short term, which can lead to problems arising from the public's perception of benefit, especially

where expectation of improvement is high. On the other hand, if generally applicable fixed emission standards are set too high they could lead to unnecessary over-expenditure or could be deemed unaffordable with the result that nothing is done.

Fixed emission standards need not be the same for all discharges. It is not unusual to set one FES for inland rivers, another for lakes and another for saline waters. The European Directive on Urban Waste Water Treatment (CEC, 1991) is a good example of an effective fixed emission standard approach. For discharges to inland watercourses there are two fixed standards, a basic one that provides a minimum degree of treatment and a more rigorous one for discharges to 'sensitive areas'. The basic standard is given in terms of BOD or COD, and suspended solids, the latter being optional, while the more rigorous one includes limits on nitrogen and phosphorus. Both standards can be achieved in terms of concentration or minimum percentage removal. An attractive feature of the European standards is that they contain only a few determinands and they are simple. The Directive is meant to establish a minimum degree of treatment in all member states, upon which individual countries can build and impose stricter standards if not already in place. This is a sensible approach in Europe where countries vary from ones with virtually no treatment to ones with high standards and high levels of treatment. A danger, however, is that those with high standards could be tempted to reduce them to the basic ones.

While the EU Directive can be considered as a sensible application of FES, there are examples of poor application in other parts of the world. A major criticism is that they often contain long lists of parameters with concentration limits that could not be achieved in an affordable manner. In many developing countries, urban watercourses are no more than open sewers while the standards in place are aimed at maintaining high-class rivers. Brazil is a good example, especially since it covers such a vast area with rich communities towards the south and very much poorer communities towards the northeast.

The National Council for the Environment passed a Resolution (No. 20) in 1986 (CONAMA, 1986) that established FESs for all discharges to all watercourses. The standards, described in Article 21 of the Resolution, are federal ones, but states are free to adopt their own. However, states either tend to adopt the federal ones or use them as a basis for their own. There are logical limits on such measures as pH and temperature but some restrictions, such as the one that restricts flows to 1.5 times the daily average, are without logical foundation. Strangely, the standards do not include either BOD or COD or any other measure of oxygen demand. The main problems arise from the limits set on 31 individual contaminants that range from heavy metals to organophosphates and chlorinated hydrocarbons. A major problem is the limit of 5 mg/l placed on ammoniacal nitrogen. If this were to be strictly applied, it would completely exclude the ability of the country to develop a phased approach to the improvement of environmental conditions and provision of basic sanitation. The only way to achieve the ammonia limit would be to use aerobic treatment. It could not be achieved with simple primary settlement nor with anaerobic treatment, yet these are both very cost-effective methods of providing significant initial improvements.

Fixed emission standards that contain long lists of determinands are often adopted in an attempt to cover all circumstances, but they sometimes turn out to have negative benefits. In many situations they are impossible to regulate effectively; they cause design and contractual problems and they give rise to application of inappropriate or unaffordable technologies and sometimes nothing is built for fear of prosecution and no progress is made.

3.2 Water quality standards (WQSs)

Much of the world sets its discharge standards to meet some water quality objective (WQO) usually based on the use that would be made of the water in a watercourse or in a particular stretch of a watercourse. WQOs are usually determined in the form of a classification system. In the UK there are four classes, with

class 1 divided into 1A and 1B. Brazil has eight classes, four for rivers, two for saline waters and two for brackish waters. There are similar classification systems in other countries but it is not within the scope of this chapter to describe or discuss in detail any such classification. It is sufficient to recognize that standards can be set to meet the individual WQOs. This, of course, means that standards could vary from discharge to discharge, depending on individual water classification.

For discharges to inland rivers, the historical method for determining an effluent quality was to use a simple mass balance that determined the discharge load by balancing the sum of the upstream load and the discharge load with the needs of the downstream quality objectives. Although the Royal Commission Standards are often used as fixed emission ones, they are actually based a WQO approach, using a simple mass balance in which the model had an eight-to-one dilution factor and the downstream quality objective was that the river should not deteriorate by more than 2 mg/l BOD.

Over the years, limitations of the simple mass balance approach have been recognized and more sophisticated methods developed to describe WQOs on a statistical basis, where the value of some determinand would not be exceeded for more than a specified percentage of time, usually 95%. The mass balance to determine the discharge standard is then determined statistically using a combined distribution method such as Warn-Brew or Monte Carlo simulations (Warn and Brew, 1980). The result is that an effluent standard is produced on a percentile basis. While the use of such techniques is logical and environmentally sound, it can cause difficulties in enforcement and a full understanding of the significance of the statistical approach is vital in the design of treatment facilities.

Some legal and institutional systems find it difficult to cope with a statistical discharge standard since it, in effect, 'allows' failures to occur. There is sometimes a fear that significant pollution could occur during the 'allowed' failure days, while the discharge limit is still achieved in overall statistical

terms. A way to overcome this problem is to introduce, in addition to the percentile standard, an upper 'maximum admissible concentration', never to be exceeded. Another problem with the approach is that, to be effective, many samples have to be taken to give statistical significance, and this is often beyond the capacity of some countries.

3.3 Sample types and sampling frequency

Discharge standards can be set using either 'snap' samples or 'composite' samples. A 'snap' (also referred to as 'grab' or 'spot') sample is simply one taken at random at a single moment in time. A 'composite' sample is one that attempts to measure the effect of an effluent over a period of time, usually a day. Composite samples can be produced manually by mixing individual snap samples in proportion to the volume that passes in the time between consecutive samples. However, it is more common to use an automatic sampler that can produce a composite sample weighted in proportion to the wastewater flow. Most automatic samplers are also capable of taking a number of discrete samples that could be analysed individually or mixed to produce a composite.

'Snap' samples are simple to take, do not involve the use of expensive apparatus and are particularly useful for audit purposes. However, they only reflect the position at a single instant and not at other times. Flow-weighted composite samples, on the other hand, better reflect the total pollution load but tend to mask individual 'spikes' in concentration. They also require expensive apparatus. The most comprehensive way to monitor a discharge would be to use a flow weighted 'composite' sample to measure the daily pollution load and to analyse a number of discrete 'snap' samples over the period of a day. However, this is prohibitively expensive in many cases. A reasonable compromise for large, significant discharges is to use composites along with occasional diurnal surveys based on 12 or 24 individual 'snap' samples.

It can be argued that, for one-dimensional free-flowing rivers, snap samples are better

TABLE 18.1 Log-normal distribution of BOD and suspended solids

	95%		Absolute	
	Composite	Snap	Composite	Snap
<i>BOD (mg/l)</i>	25	31	50	65
	20	25	45	60
	15	19	40	55
	12	15	35	50
<i>Suspended solids (mg/l)</i>	50	70	115	160
	45	63	110	140
	40	56	90	125
	30	42	75	105
	25	35	65	90

than composite since the important environmental criterion is concentration and not load. For tidal estuaries or tidal rivers, where the flow direction changes with tide, load may be more appropriate and a composite sample a better measure.

The description of an effluent quality using a series of snap samples is not the same as the description of the same effluent using a series of composite samples. The matter is further complicated when snap and composite samples are used to describe effluents based on 'absolute' values or on 'percentile' values. Ross (1994) used the information given in Table 18.1 to indicate the nature of the relationship based on a log-normal distribution of BOD and suspended solids.

Thus, an effluent with a composite BOD of 25 mg/l is equivalent to an effluent with a snap of 31 mg/l. The same effluent, if it were judged on an absolute (99.9%) basis, would have BOD values of 50 mg/l and 65 mg/l. These figures are not presented to indicate absolute relationships; rather they indicate the nature of the relationships and the difficulties of both specifying and understanding effluent discharge standards. When setting numerical limits, it is important to be aware of the significance of the sampling regime to be used, as well as the approach used to determine the standards. It is a complex subject far beyond the scope of this chapter.

3.4 Industrial wastewaters

Industrial effluent control is of vital importance, particularly in developing countries where control is often very poor or non-existent. There is an old axiom which states that 'good sewage treatment starts with good control of industrial effluents'. Non-existent control could mean an inability to treat domestic sewage effectively.

The prime reasons for the control of industrial wastewaters are:

- to protect workers working within the sewerage system
- to protect the fabric of the sewer and the fabric of any downstream treatment works
- to prevent fires and explosions due to inflammable or explosive chemicals
- to prevent surcharging or blockages within the sewerage system
- to protect the physical operation of downstream treatment works
- to ensure that industrial discharges do not affect the performance of downstream biological treatment processes
- to protect the environment where sewers eventually discharge to a watercourse or to the sea
- to ensure that industrial discharges do not affect disposal of sludges from treatment works.

The organization responsible for the sewerage system and sewage treatment works must set standards for individual sewer discharges, based upon the following factors:

- the type and nature of the industrial wastewater
- the capacity of the sewerage system to accommodate the wastewater
- the potential of the wastewater to cause surcharging or blockages of the sewerage system
- the potential effect on the capacity of the treatment plant
- the potential effect on biological treatment processes.

For any given sewerage system, the total daily flow, suspended solids, BOD or COD, and ammoniacal nitrogen determine the size and capacity of the required sewage treatment works. Pretreatment of industrial wastewaters may be required to maintain flows and loads within treatment capacities, or extension of the treatment works may be necessary. Substances that affect biological processes must be limited to concentrations below those that would cause problems and the limits need to be set taking account of quantities arising from other sources.

If the standard for the sewage treatment discharge has limits on certain substances, the only recourse may be to control them at the industrial source. Standards for substances such as sulphides, cyanides and heavy metals should be set at very low limits, whereas some, such as the 'red list substances', should be banned altogether. Others including many volatile organic compounds could also be considered for prohibition. Toxic metals arising from industry need to be controlled as they become incorporated into the biosolids (sludge) and can seriously affect disposal to agricultural land. In some countries (e.g. USA) pretreatment standards apply to specific industrial categories, so that all are treated with some degree of equity, but this is not the case in other countries.

4 MEETING THE STANDARDS

Significant developments in process technology in recent years mean that there are now many ways of treating sewage to achieve a wide range of discharge standards. However, the availability of a process is not by itself sufficient; it should be cost-effective, appropriate to the local conditions and affordable. In this sense there needs to be a differentiation made between the developed and developing worlds.

Most of the developed world has extensive sewerage networks, a relatively high degree of sewage treatment and some form of industrial wastewater control. Discharge standards are now aimed at improving conditions in which

gross pollution is generally absent. Targets differ from place to place, but the general tendency is towards improving the aquatic environment and minimizing risk to humans using that environment. In general, developed world countries can afford to pay for these improvements, although there are some who may be reluctant. By contrast, most of the developing world has inadequate sewerage systems, very little sewage treatment and almost no effective industrial wastewater control. Urban watercourses are no more than open sewers whose condition is further reduced by large amounts of garbage arising from inadequate collection facilities. Many developing countries either do not have discharge standards or use inappropriate ones imported from the developed world.

Although the conditions in the two worlds are different, the problems are really similar but out of time sequence. The conditions in many developing world cities today are the same as those of the developed world cities 50 to 100 years ago. There are, however, important differences. On the one hand, the developing world's financial problems are likely to make it more difficult to bring about improvements at a rapid rate but, that on the other hand, they can benefit from some of the technological advances made during the intervening years and, hopefully, learn from previous mistakes of the developing world.

4.1 Sewerage and sewage treatment

Critical to the improvement of environmental conditions is the need for effective collection and transportation of sewage for treatment. Most of the developed world has elaborate networks based on either separate or combined sewerage systems with sewage transported to treatment plants by a system of interceptor sewers and pumping stations. When properly installed, operated and controlled the separate system is the more cost-effective as it reduces the amount of sewage to be treated and avoids the problems of discharges from combined sewer overflows (CSOs). In fact, the issue of treating the flows from CSOs is one of the principal problems facing the sanitation

industry in the developed world today; the solutions are not easy and tend to be very expensive.

Following the lessons of the developed world, the sewerage systems of many developing world cities were designed and built as separate systems. However, unlike the developed world cities, they have often not been well operated since the control of connections is virtually non-existent. As a result, so-called separate systems have many connections illegally made to the surface water sewers and not to the foul or sanitary sewers as intended. There are also many cross connections and, in many cases, separate systems are effectively combined systems. This causes great problems when collecting sewage in interceptor sewers. If only discharges from recognized sanitary sewers are collected, much of the sewage will continue to be discharged through the surface water system, so diminishing the benefit of collection. Separating out the two systems is extremely difficult and prohibitively expensive and could delay effective collection by many years. An option could be to collect all discharges containing sewage, transport everything for treatment and install storm-water facilities at the treatment plant, but this would completely remove the cost effectiveness of a separate system. It could also cause some urban watercourses to dry up.

It is beyond the scope of this chapter to debate the problems of competing sewerage systems, interception and transportation. However, for towns and cities with such conditions, it should be recognized that the introduction of high discharge standards for treatment plant effluents would be of no avail if a large proportion of the sewage were still discharged directly through stormwater sewers. Under these circumstances it would be much more effective to phase discharge standards over a long period to take account not only of affordability, but also of prolonged delays in getting adequate quantities of sewage to the treatment plant.

With regard to the treatment of sewage, there is a wide range of methods and processes available to cover all of the world's needs

provided the selection is appropriate to the environmental conditions, that it is affordable and that the community is willing to pay. Within today's treatment technology it is possible to design plants that can do anything from simple removal of gross solids to ones that would produce high quality water for reuse. The standard should therefore reflect the objectives, and the objectives of the developed world are not the same as those of the developing world, at least not in the short term. The use of developed world standards in the developing world is generally a retrograde step.

While a standard should be set to meet an environmental objective, the way that it is written can have a profound effect on the design and cost of treatment; a subject that should be fully understood by both the standard setter and the treatment plant designer. This is particularly true if the standard is written in statistical terms. The difference in design for an absolute standard and a statistical standard (see Section 3.2) can be very significant and the interpretation of the standard is very important. This was illustrated by Johnstone and Norton (2000) who showed the difference in cost of an aeration system to produce a 50 mg/l BOD standard based on four different ways of describing 50 mg/l, – as an average composite, a 95 percentile composite, an absolute composite and an absolute snap. Designing on an absolute composite basis produced the lowest cost. The design based on a 95 percentile composite was 31% more expensive, while that based on an absolute (99.9 percentile) composite was 92% more expensive. The design based on an absolute (99.9 percentile) snap was 230% more expensive. These are very significant cost differences, which become even greater when the numerical limits are reduced (i.e. when the standards become higher), not only in absolute terms but also in relative terms.

An appreciation of the statistical nature of sewage effluents and a full understanding of the nature of the standard is particularly important in the development of BOOT (Build, Own, Operate and Transfer) schemes for the provision of a sewage treatment plant

by the private sector, especially where financial penalties are to be imposed for failures (see Section 5.2). Dunn *et al.* (1998) made studies on the statistical nature of sewage effluents that attempt to predict the risk on the design of new treatment facilities.

4.2 The developed world

In much of the developed world, sewage treatment processes now meet ever-improving standards and the tendency is towards the application of improved aerobic secondary treatment or the addition of tertiary or advanced processes. There is greater use of nitrogen and phosphorus removal techniques, of UV disinfection rather than chlorine and, in many places, of processes that can be contained within buildings or can be built underground. Communities demand better odour control from treatment plants and no longer tolerate disposal of non-sterilized sewage sludges to agricultural land. In addition, more attention is being placed on treating combined sewer overflows.

Although some process developments have been driven by the need to reduce costs, many more in the future will be driven by the need to meet ever-improving discharge standards (Johnstone and Norton, 2000). It is likely that more use will be made of advanced treatment techniques, such those based on membrane technology or granulated activated carbon, as attention is further drawn towards removal of specific compounds. Toxicity testing of effluents will lead to greater control of industrial effluents at source. Whether or not many of the standards, especially those based on the precautionary principle and desire for zero risk, can be justified is a matter of debate (Garber, 1992; Johnstone and Horan, 1994). The cost of meeting them will no doubt increase; whether or not the community will continue to pay for them, or not, is another matter for debate.

4.3 The developing world

Meeting discharge standards in the developing world is often by application of processes at

the other end of the technological scale to those used in the developed world, although there are exceptions. In most large cities, land availability is limited and often expensive, so that some of the more basic processes are not applicable. There is no hard-and-fast rule to selecting a cost-effective process, but the decision is sometimes complicated in the developing world. On the one hand, inappropriate standards imported from the developed world would drive the selection towards processes that are inappropriate and unaffordable. On the other hand, there are many advocates of so-called 'appropriate technology' (meaning in this case low technology) who demand that basic technologies be applied everywhere in the developing world. There is a tendency in some quarters to promote the use of processes, such as waste stabilization ponds in large cities, where they may not be practicable and more conventional processes may be more appropriate.

While waste stabilization ponds may not always be appropriate for large cities, they can find great use in many other circumstances, although care must be taken in specifying standards based on BOD. Often waste stabilization ponds discharge algae with the final effluent that exerts a BOD but with little environmental damage. This was recognized by the European Union in the Urban Waste Water Directive by basing the BOD standard for effluents from waste stabilization ponds on filtered samples and not unfiltered samples as is required for all other determinands; the Directive also requires a suspended solids standard for pond effluent of only 150 mg/l.

Another problem arises when some standard setters say that BOD, suspended solids and some of the other basic sanitary parameters, are not important in the developing world and that the only parameter of importance is faecal bacteria. While there is little doubt that reduction in faecal bacteria is important, statements like this underline a lack of understanding of the underlying principles of wastewater treatment systems, and indeed often a lack of understanding of BOD. BOD is a measure of sewage strength; it is a measure of readily degradable organic matter and not just

a measure of oxygen demand. Outside the field of waste stabilization ponds, treatment plants do not remove substantial quantities of bacteria, but by removing BOD and suspended solids they can make an effluent amenable to disinfection. To purport that effluents can be efficiently disinfected without first adequately reducing the BOD and suspended solids displays ignorance of the principles of process technology.

A sensible approach to improving environmental conditions in the developing world is to use a strategy based on phasing process units, a strategy given appropriate emphasis by von Sperling and Fattal (2001) in the newly published WHO guidelines. In many cases, the provision of simple preliminary screening could do much for the aesthetics of the area, but it is more usual to have a first stage based on primary treatment or an anaerobic process such as an USAB. However, phasing is only suitable provided that the standards in force are flexible enough to allow for this type of approach. Anaerobic processes can be particularly attractive as they use very little power and they produce minimal quantities of sludge. However, they tend not to be capable of dealing with large flow variations and can be adversely affected by high concentrations of heavy metals, which can be a problem in places with poorly controlled industrial effluents.

Advanced primary sedimentation using chemicals could also provide an initial stage, or it could be used as a first upgrade for a basic sedimentation system. There are a number of proprietary advanced primary treatment processes available.

The addition of secondary treatment or the enhancement of first phase treatment can take place at any time, but it should be introduced in the full knowledge of pollution from other sources. For example, if there is still major pollution arising from contaminated surface water sewers, the benefit from secondary treatment may be marginal and funds would be better spent in tackling the sewerage problem. There is no simple answer but, for developing countries, it is important to consider the problem in the long term and phase treatment facilities according to benefit and affordability. To do so also

requires a phased approach to the introduction of standards and, if necessary, to have interim standards and long-term targets.

5 REGULATION

There is little doubt that the setting of appropriate environmental standards is important. However, their effectiveness is only as good as the subsequent enforcement and regulation. Enforcement is principally a matter of political will. In some parts of the world, standards abound without the political will to enforce them and this achieves nothing. Sometimes there are ostensibly good reasons, as the expenditure needed to meet environmental standards must take its place beside other priorities.

Regulation is often seen as being too expensive and it can be if not done effectively. For example, consider the monitoring of discharges and the frequency of sampling. It is reasonable to argue that where society correctly requires a high standard, the frequency of sampling should reflect the importance of that high standard and the costs of achieving it. Equally, when a low standard is required, it seems reasonable to allow a relatively low sampling and monitoring regime. Yet there is little evidence of this being common practice and even when it is practised, it is doubtful if the monitoring frequency is high enough to monitor high standards with justifiable statistical significance. Surely if society deems it appropriate to impose a high standard, then the cost of adequate monitoring should be taken into account as part of the cost of achieving it.

The issue of regulation and enforcement highlights another marked distinction between the developed and developing worlds. Countries in North America, Europe and Scandinavia, for example, have mature and sophisticated environmental agencies, most of which are well funded. They generally have political support and society can afford to pay for them. By contrast much of the developing world is characterized by poor or inadequate environment agencies that are badly funded and without real political support.

Traditionally, sewage treatment works have been built and operated by the public sector, with the discharges regulated and controlled by other public sector organizations, such as an environment agency. This would normally be done by the issue of a licence or some other legal instrument and controlled through a regular monitoring programme. The consequence of failing to comply with a standard would range from virtually none, through reprimands to financial fines.

Over recent years there has been a greater use of private sector financing of treatment plants by way of some form of BOOT (Build, Own, Operate and Transfer) scheme, of which there many variations. Regulation and control of discharges is also through a licence that is usually no different from that for a public sector works. However, regulation and monitoring tend to be much stricter and the consequence of failure much more severe. It is not unknown for daily fines of around US\$25 000 to be imposed for failure to achieve the discharge standard. Under these circumstances, the nature and wording of a standard and its interpretation can have profound effects on the design and operation of the plant. Indeed, it can have a profound effect on the success of the whole project, a subject discussed below.

5.1 Permit, licence or consent

Legal instruments for setting out discharge standards and conditions differ from place to place, but they generally take the form of a 'permit', 'licence' or 'consent'. It is a contract between the discharger and, usually, a regulator empowered by government to enforce standards. Although there is a wide variation in such documents, they usually comprise:

- the standards to be achieved; these may be descriptive or numeric
- specific conditions relating to the effluent
- general conditions relation to the institutional requirements.

Descriptive standards usually relate to things that are difficult to measure and could include general statements such as:

- 'there shall be no sign of visible signs of solid matter downstream of the discharge point' and/or
- 'there shall be no signs of visible oil downstream of the discharge point'.

Numerical standards specify limits on physical and chemical parameters, such as daily flow rate, maximum flow rate, pH, temperature, BOD, COD, suspended solids and heavy metals, etc. They might also include toxicity limits but, at present, protocols for setting such limits are not fully developed.

Specific conditions may relate to the exact location of the discharge point and might describe the exact nature of a required outlet, which could be a pipe or spillway. Materials of construction may also be specified. It is often a requirement that a flow metre is installed and that flow records be kept and made available for inspection by the regulator. It would be usual to specify that a suitable sampling point be constructed at a designated position.

General conditions may specify regulations in the law under which the legal instrument was issued, and the powers of entry that allow regulators to inspect premises and to take samples. The frequency of audit sampling could also be specified, as could the penalties for infringement and the steps that have to be taken during infringement. For example, it may specify the exact procedures to be followed when taking samples, especially those to be used in evidence of a breach. There could also be general conditions that absolutely exclude the discharge of certain substances, or groups of substances.

It is important that, no matter what the form of the instrument, it should specify the exact requirements and lay out unambiguously the performance, duties and obligations of the discharger.

5.2 The polluter pays principle

In some areas of the world, such as in Scandinavia and the Baltic States, there are systems for charging for pollution, which is an additional means of regulation and control. In

one Baltic country, charges are imposed on the loads of BOD, suspended solids, total nitrogen, phosphorus and on 'oil and grease'. There are separate tariffs per tonne of each parameter and the charge is imposed four times per year on the total load for the period. As an encouragement to meet the standard, the tariff is reduced to 50% if the effluent complies with the standard for the whole period. If it does not, the full tariff is imposed for all days during which the effluent remains within the standard; and for those days during which the effluent fails the standard the tariff is increased tenfold. If the tariffs are set at proper levels, this can prove to be a very effective means of regulation as it can force operators to look towards optimizing operational control. However, if set too low, it merely becomes another form of taxation. It can also be an expensive form of control, since samples of effluent have to be taken every day of the year and, eventually, this cost and the cost of the fines has to be borne by the community.

5.3 Standards, regulation and the private sector

There are many ways that the private sector can participate in the treatment of sewage, one of which is the BOOT contract. In this, the private sector, usually a consortium of contractors and operators, raises the finance to fund the construction of a treatment plant. After construction the operating arm of the consortium operates the plant for some 20 to 30 years, during which time they are paid per m³ of sewage treated. Payment on a volume basis is usually the only source of income and must be sufficient to pay for loan repayments, dividends and all operational and other expenses. In some contracts, failure to comply with the discharge standard could incur substantial financial penalties.

There are many risks in this type of venture and there are two that are relevant here, which if not recognized and handled properly could cause the failure of this type of scheme. The first is concerned with payment on a

volumetric basis. Due to the problems with sewerage and bringing sewage to the treatment plant (see Section 4.1), there could be prolonged delays in achieving sufficient volumes to pay for treatment, which, in combination with failures to meet a suitable standard, could cause the project to fail. If not properly addressed, the private sector may not be prepared to bid for such a project in the first place.

The problems with discharge standards can arise from the nature of the standards, the limits set, type of monitoring to be undertaken, errors in sampling and analysis, and generally the question of 'what constitutes a failure?' When the penalties for failure are high, these issues are of great concern, especially since there is often mistrust between public sector regulators and private sector operators. Often problems arise due to unclear or unspecified rules.

If a standard is set as an absolute one (never to be exceeded) there will inevitably be failures and fines, and the higher the standard, the more frequent the fines. However, it is more practical to have the standards imposed as (for example) a monthly average, or a monthly average with an upper limit, or a percentile compliance over a given period. Whatever is chosen, it needs to be specified unambiguously in the contract and understood clearly by both the operator and regulator. At present, there are many cases where it is not and where problems are inevitable.

Another issue is concerned with what actually constitutes a failure and should this incur a fine. If there are, for example five parameters in the standard where four pass and one fails, is this a failure that should incur a fine? Also, what allowance should be made at the margin of analytical and sampling error? For example, if a BOD limit is 20 mg/l and a sample analysis is 21 mg/l, is this a failure bearing in mind that analytical error may be around 20%? It seems inequitable that a private company should suffer a large daily fine if four out of five determinands are within limits and the fifth is outside only by the margin of analytical error. The environmental consequences of this would be zero. It may appear

that such ambiguities would not be real but they are, and they arise from inexperienced regulators setting inappropriate rules for regulation and uninformed contractors agreeing to them without understanding the consequences. The problems would not be so great if disputes were left to reasonable people to resolve, but this is rarely the case. The problems become acute when lawyers and prosecutors get involved, as the law tends to see things as black and white, whereas they are usually grey. However, the regulation of the private sector is still in its infancy and no doubt these issues will be resolved with time and experience.

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Part III Microbiology of Wastewater Treatment

Introduction to Microbiological Wastewater
Treatment

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Fixed film processes

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1 INTRODUCTION

Raw wastewaters have long been discharged directly to receiving waters (and still are, even in some industrialized countries), hoping their self-purification capacity would take care of our waste. Unfortunately, most receiving bodies failed to do so as they were overcharged with organic and nitrogen pollution resulting in a chronic state of degradation. In most cases, wastewater treatment is therefore needed before effluent discharge. One of the most popular types of treatment is the biological one which more or less mimics some of the natural processes found in a self-purifying receiving body, mainly organic degradation and nitrogen conversion through bacterial action. These can be done by attached or suspended microorganisms, giving rise to two main families of wastewater treatment processes (Fig. 19.1): (a) the fixed film processes, discussed here and represented by the trickling filter, and (b) the suspended growth processes, such as the activated sludge, discussed in Chapter 21.

However, one has to remember that to engineer these natural processes into a compact biological sewage treatment processes, one must satisfy the following criteria (Mudrack and Kunst, 1986):

- The biomass concentration in the reactor must be increased

- The increased oxygen demand must be catered for
- An optimal contact between the biomass, the sewage constituents and dissolved oxygen must be ensured
- Inhibitory or toxic substances or inhibiting factors must not be allowed to reach critical levels in the reactor.

These points may seem trivial, but they are in fact not that easy to achieve.

2 FIXED FILM PROCESSES IN WASTEWATER TREATMENT

2.1 Biofilm development

Fixed film processes are based on the capacity of different microorganisms to grow on surfaces. They tend to attach to solid surfaces due to various reasons:

1. substrate availability
2. protection from a harmful environment, particularly at high-velocity water currents
3. interaction of physical forces like attraction, adsorption and adhesion (Senthilnathan and Ganczarczyk, 1990).

Fundamental mechanisms of microbial attachment are reviewed in a subsequent section,

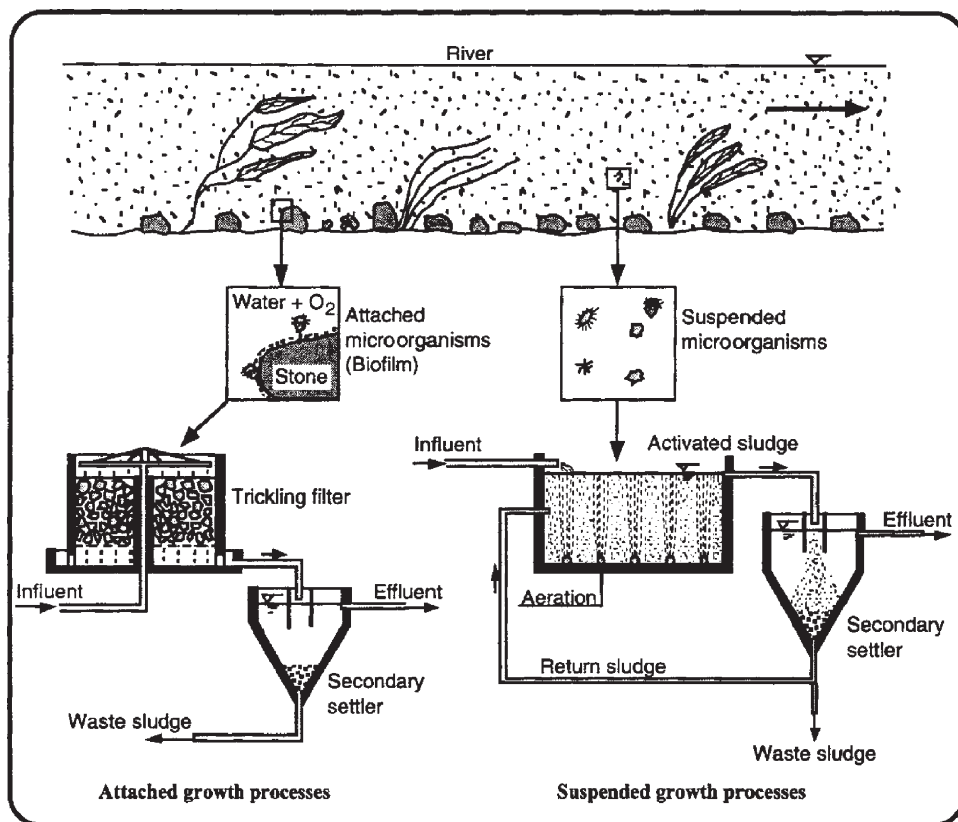


Fig. 19.1 Transfer of self-purification process and organisms from a river to a biological sewage treatment plant (adapted from Mudrack and Kunst, 1986).

but it is necessary here to describe minimally the biofilm formation process before discussing technologies.

Bacterial adhesion is, in general, a three-step process (adsorption, adhesion and adherence) which takes approximately from 20 to 30 minutes under optimal environmental conditions (Brisou, 1995). Fixation mechanisms will depend on the type of bacteria and on the type of surface available for them to colonize.

Adsorbed microorganisms (specifically bacteria) grow, reproduce and produce extracellular polymeric substances, which frequently extend from the cell forming a gelatinous matrix called a 'biofilm' (Bryers and Characklis, 1990). These exopolymers consist mainly of a variety of heterogeneous polysaccharides depending on the type of

microorganisms involved (Lazarova and Manem, 1995). A schematic view of a biofilm is shown in Fig. 19.2.

The substrate removal mechanism within a fixed film process is very complex. The phenomena that take place when a biofilm is brought into contact with a wastewater-containing substrate and oxygen are as follows (WPCF, 1988):

- Transport of the substrate and oxygen from the wastewater to the surface of the biofilm
- Internal transport of the substrate and oxygen through the biofilm by diffusional processes
- Oxidation of the substrate within the biofilm
- Diffusion of by-products back to the wastewater.

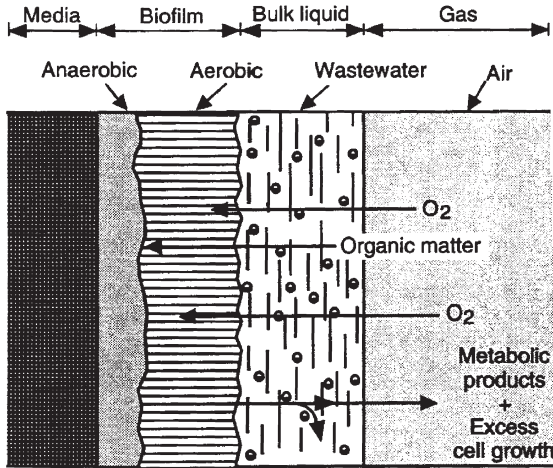


Fig. 19.2 Schematic view of a biofilm.

Typically, after some microorganisms have adhered to a surface medium they start growing, producing the biofilm, which is aerobic. As growth continues, oxygen diffusion across the biofilm may become limiting and an anaerobic layer develops along the surface media. Thus removal of organic matter implies a continuous growth of bacteria and consequently a thicker biofilm. This accumulation process is generally balanced by the biofilm sloughing off. The following conditions are important for the sloughing off to occur and are used technologically, intentionally or unintentionally, to control the biofilm (Henze *et al.*, 1995):

- Hydraulic erosion acts continually on the surface of the biofilm and leads to a steady sloughing off on the outer side
- Degradation of bacteria in endogenous phase at the bottom of the biofilm may cause a weakening of the adhesion
- Gas formation within the biofilm (e.g. methane or nitrogen) may destroy the adhesion.

Development of a bacterial biofilm for substrate degradation is thus a complex phenomenon that is influenced by many factors such as: wastewater characteristics (nature of substrate, nutrients ratio and environmental conditions), operational factors (inoculation, organic loading rate, hydrodynamics) and the support medium.

The development of technologies based on the concept of attached growth therefore necessitates a good knowledge, not only of process behavior but also of microbiology and biofilm formation.

2.2 Usual processes

A possible classification of biological growth systems is shown in Fig. 19.3. Some authors have proposed more specific classifications for fixed film processes (e.g. Crites and Tchobanoglous, 1998; Lazarova and Manem, 1994), but a general one for all types of fixed film processes found in wastewater treatment is difficult to establish as many factors can be considered: type of medium (organic versus inorganic), type of reactors (fixed bed versus moving bed), environmental conditions (aerobic versus anaerobic), wastewater characteristics (municipal versus industrial) and types of pollutants to be removed (C, N, P, xenobiotics). It is difficult to cover in this chapter all possible fixed film technologies for wastewater; only the applications of fixed film processes related to municipal wastewaters are described, which means mainly aerobic processes in packed bed reactor or hybrid systems, and carbon, nitrogen and phosphorus removal.

Fixed-film processes in municipal wastewater treatment are now over a hundred years old, with the introduction in the late nineteenth century of the trickling filter (Peters and Foley, 1983). Since then other processes have appeared, but until the 1980s, trickling filter (TF) and rotating biological contactors (RBC) were almost the only two processes used on a regular basis. Biological aerated filters (BAF) appeared on an industrial scale in the 1980s, while the 1990s saw the arrival of hybrid systems consisting of biomass support systems (BSP) immersed in AS reactors and used mainly for plant upgrading.

2.2.1 Trickling filters

The trickling, or percolating, filter, was the first fixed film process that was developed. It is a packed bed reactor (2–3 m deep) filled with a medium consisting of 5–10 cm stones

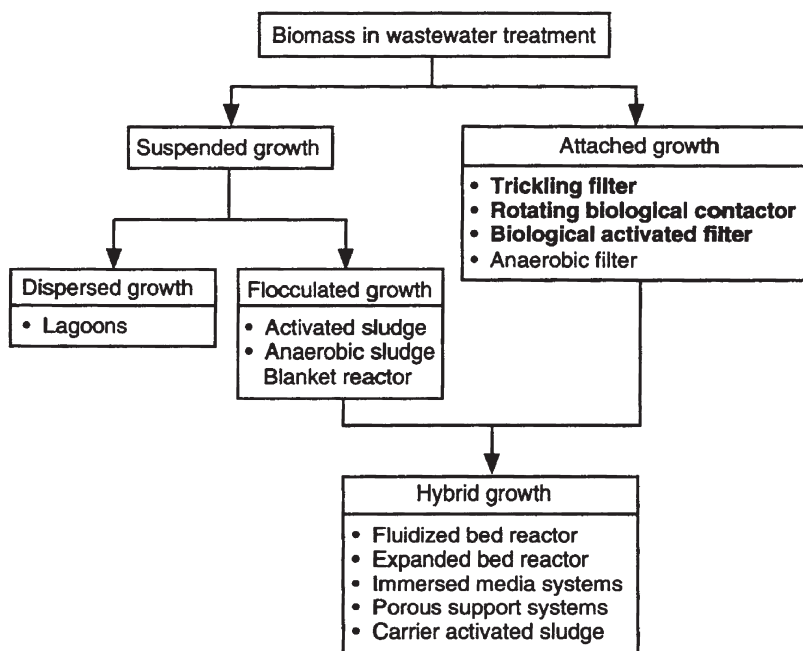


Fig. 19.3 Systematics of biomass forms in wastewater treatment systems (adapted from Senthilnathan and Ganczarczyk, 1990).

with a specific surface area of $40\text{--}100\text{ m}^2/\text{m}^3$. The trickling filter is very efficient with respect to adhesion of bacteria, contact between water and biofilm and reaeration of the water (Henze *et al.*, 1995).

After a compulsory primary sedimentation step, the wastewater is distributed over the gravel bed, trickles down to be collected under the filter and flows to a secondary settling tank, also called a humus tank. Aeration is provided through natural drafts resulting from temperature differences between the ambient and the internal air. A typical trickling filter is shown in Fig. 19.4, and its advantages and disadvantages are summarized in Table 19.1.

Trickling filters can be operated in different modes: single pass, alternating double-filtration

and recirculation mode (Bitton, 1994). Depending mainly on the organic and hydraulic loading rate, trickling filters will be classified as low, standard (or intermediate) and high rate. Design criteria are given in Table 19.2, and more information can be found in standard textbooks (Metcalf and Eddy, 1991; WEF and ASCE, 1992; Henze *et al.*, 1995; Crites and Tchobanoglous, 1998; Grady *et al.*, 1999). However, according to Parker (1999), engineering practice in trickling filter design has been influenced by the propagation of myths as much as by the analysis of data and factual determinations.

Effluents from well-operated trickling filters range around 20 mg/l for BOD_5 and SS (Crites and Tchobanoglous, 1998). At weak organic and hydraulic loadings (around $0.2\text{ kg BOD}/\text{m}^3\text{day}$

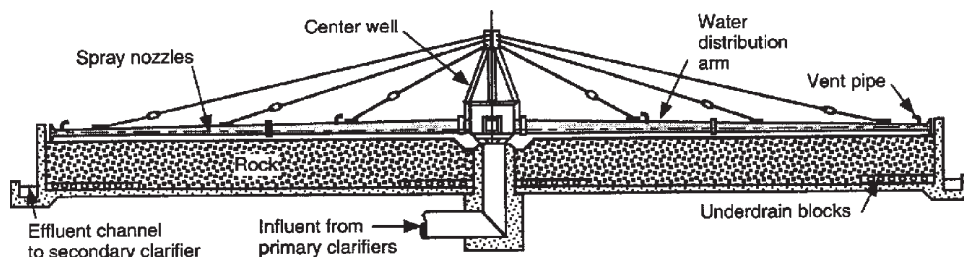


Fig. 19.4 Typical trickling filter.

TABLE 19.1 Some advantages and disadvantages of trickling filters

<i>Advantages</i>	<i>Disadvantages</i>
Ease of operation	Filter clogging for high organic loads results of excessive slime bacteria
Low maintenance and energy costs	Potential odor problems
Reliability	Flies
Sloughed biofilm easily removed by sedimentation	No real possibilities for operational control
Able to treat industrial wastewaters	Pretreatment and primary sedimentation necessary
Withstand shock loads	

and $0.66 \text{ m}^3/\text{m}^2 \text{ day}$) trickling filters can nitrify fully (Henze *et al.*, 1995). The process is thus capable of compliance with effluent discharge consents of $2 \text{ mg NH}_3\text{-N/l}$, as shown by Pearce and Williams (1999).

The use of plastic media, with a specific surface area around $200 \text{ m}^2/\text{m}^3$, has made it possible to create a large volumetric surface in low-weight filters, and these can therefore be built as towers. Higher organic and hydraulic loading rates can be applied: $1\text{--}5 \text{ kg BOD}/\text{m}^3 \text{ day}$ and $36\text{--}72 \text{ m}^3/\text{m}^2 \text{ day}$. Such filters are, however, typically used for the treatment of industrial wastewaters.

2.2.2 Rotating biological contactors

Rotating biological contactors (RBCs) consist of a series of circular plastic disks mounted on a

horizontal central shaft and distanced at $1.5\text{--}2.5 \text{ cm}$. Normally 40% of the disks' surface is submerged and they are rotated in a tank containing the wastewater. Attached microorganisms rotate into the water, where organic matter is adsorbed onto the biofilm, and out of the wastewater, where the oxygen necessary for the conversion of organic matter is obtained by adsorption from the air. Thus, the rotation provides aeration and the shear force that causes sloughing off of the biofilm from the disk surface. The peripheral speed on the disks should not be less than 0.3 m/s (Henze *et al.*, 1995). A typical RBC is shown in Fig. 19.5, and its advantages and disadvantages are given in Table 19.3.

Some authors suggest that the suspended biomass found in the reactor may also contribute to some 4 to 10% of pollution removal (Edeline, 1993). However, due to the short retention time, these additional removals can be neglected (Benefield and Randall, 1980).

Design criteria are given in Table 19.4, and more information is given in standard textbooks (Metcalf and Eddy, 1991; WEF and ASCE, 1992; Henze *et al.*, 1995; Crites and Tchobanoglous, 1998; Grady *et al.*, 1999).

Good quality effluents (approximately 15 mg BOD/l) can be obtained with organic loadings in the range given in Table 19.4. However, according to the Bavarian Water Authority, the organic loading should not exceed $10.5 \text{ g}/\text{m}^2 \text{ day}$, if an effluent BOD less than 25 mg/l is desired (Edeline, 1993). Nitrification can be achieved in RBCs, provided a second or third stage of disks is added.

2.2.3 Biological aerated biofilters

Over the last 20 years biological aerated filters (BAF) have been developed not only as competitors for other secondary treatment systems, such as trickling filters and activated sludge plants, but also for tertiary treatment (Mann *et al.*, 1999). These biofiltration processes are popular in Canada (Pineau and Lessard, 1994) and France (Canler and Perret, 1994), and they are gaining interest in other countries as well. The possibility of achieving carbon oxidation with nitrification/denitrification in

TABLE 19.2 Design criteria for trickling filters

<i>Design criteria</i>	<i>Intermediate rate*</i>	<i>Reference</i>
Hydraulic loading ($\text{m}^3/\text{m}^2 \text{ day}$)	3.52–9.39	Crites and Tchobanoglous (1998)
	14.40–28.80	Henze <i>et al.</i> (1995)
	–	Agences de l'eau (1994)
	4.5–25	Edeline (1993)
Organic loading ($\text{kg BOD}/\text{m}^3 \text{ day}$)	0.24–0.48	Crites and Tchobanoglous (1998)
	0.45–0.75	Henze <i>et al.</i> (1995)
	–	Agences de l'eau (1994)
	0.40–1.0	Edeline (1993)

* Some call those normal, standard.

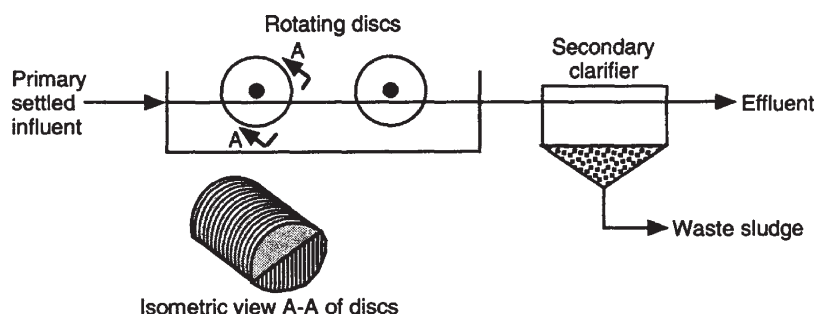


Fig. 19.5 Typical RBC system.

the same reactor is probably a major reason for its increasing usage.

This technology was developed for retention of suspended solids and for oxidation of carbonaceous soluble pollution in the same reactor. The water flows through a packed bed media which is generally inorganic with a particle size around 3–5 mm and a specific surface area of 300–500 m²/m³. Aeration for oxidation is done at the base of the biofilter and water flows upwards or downwards, giving two main types of biofilters: upflow or downflow biofilters (Fig. 19.6).

Given that the biofilm grows and suspended solids are retained within the biofilters, clogging occurs and backwashing of the process is necessary. A filtration cycle length is typically between 12 and 48 hours and backwash with air and water takes around one hour (Agences de l'Eau, 1994). The advantages and disadvantages of BAFs are given in Table 19.5.

Many types of BAFs exist and different organic loading rates can be found in the literature (Stephenson, 1997). However, the Agences de l'Eau (1994) suggested the follow-

ing design criteria for biofiltration processes: for hydraulic loading, 2–5 m³/m² h; and organic loading, 6–8 kg COD/m³ day. Nitrogen removal (nitrification) is possible with BAFs (Lacamp *et al.*, 1992; Stephenson, 1997; Canler *et al.*, 2002) together with denitrification (Peladan *et al.*, 1997).

The second generation of submerged filters involves a floating support made from a synthetic material with a density lower than that of water. This overcomes some of the disadvantages of the original systems, especially in the filter regeneration phase, for which large amounts of fluid (water and air) and power can be consumed to loosen the bed and wash away the excess sludge (Capdeville and Rols, 1992). Floating media have been found to perform better than sunken media for SS, COD_t and ammonia removal and at higher flow rates under shock loading conditions (Mann *et al.*, 1998, 1999).

The trend in BAFs is now to look at more compact filters able to perform at high water velocities (10–33 m³/m² h) and to achieve nitrification/denitrification within the same biofilter (Peladan *et al.*, 1997; Puznava *et al.*, 2001). Studies are also being undertaken to determine the feasibility of removing phosphorus biologically through sequencing of

TABLE 19.3 Some advantages and disadvantages of RBCs

Advantages	Disadvantages
Short residence time	No operational flexibility
Low operation and maintenance costs	Has to be covered for cold climates
Low energy costs	Need of primary sedimentation
Good settling characteristics of sloughed materials	Possible dryness of unsubmerged biofilm portion (warm climates)

TABLE 19.4 Design criteria for RBCs

Design criteria	Criteria	Reference
Hydraulic loading (m ³ /m ² day)	0.04–0.15	Edeline (1993)
Organic loading (kg BOD/m ² day)	0.005–0.026	Henze <i>et al.</i> (1995)
	0.015–0.060	Edeline (1993)

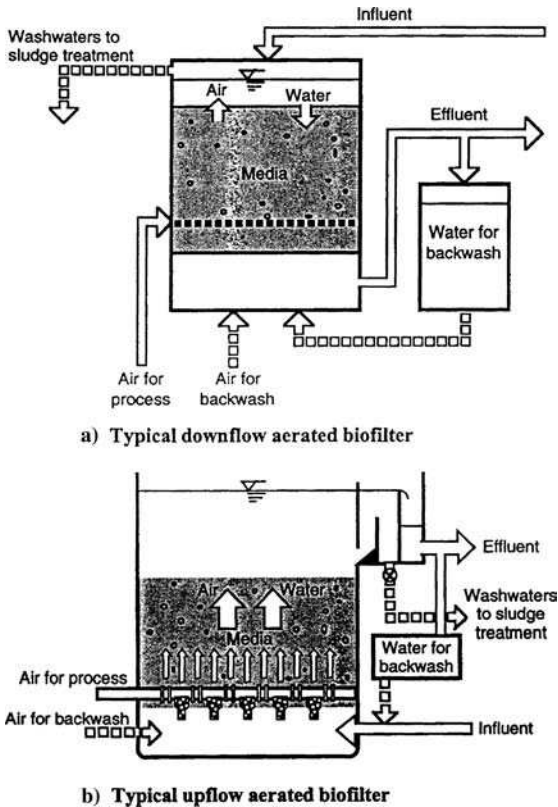


Fig. 19.6 Aerated biofilters. (a) Typical downflow aerated biofilter; (b) typical upflow aerated biofilter.

the filter operation (Morgenroth and Wilderer, 1999).

2.3 Microbiology of standard processes

For all processes, the continuous addition of substrate and microorganisms present in the wastewater encourages the formation of a

TABLE 19.5 Advantages and disadvantages of biological aerated biofilters

Advantages	Disadvantages
Good quality of treatment	Transient operation
Compact process, no secondary settlers required	High energy consumption
Fast recovery after problems	Large volumes of washwaters
Start up quick even after several months stopped	

complex biofilm composed of bacteria, fungi, protozoa, macroinvertebrates (larvae, worms) and sometimes algae (Bruce and Hawkes, 1983). Fig. 19.7 shows in which order the colonizing microorganisms appear in a biofilm, which already contains several cellular waste and inert particles from the wastewater being treated.

The microbiology of attached growth processes is somewhat similar from one process to another. Possible differences are attributable to the maturity of the biofilm, which varies between processes, thus influencing microbial colonization.

2.3.1 Bacteria

Bacteria are unicellular prokaryotic organisms. Besides being the most abundant organisms in the fixed film treatment processes ($\pm 10^{10}$ CFU/g VSS), they are the first to colonize the biofilm (Mack, 1975). Their small size and large surface with relation to their volume, gives them a clear advantage for the assimilation of the substrate in the liquid phase. Under optimal growth conditions, the doubling time of heterotrophic bacteria is about 20 minutes, which helps them out-compete any other organism in the process.

Generally, for fixed film processes, the dominant bacterial genera are very similar to those found in activated sludge. During the aerobic treatment of domestic wastewater, these genera are often Gram-negative heterotrophic rod-shaped organisms, including *Zooglea*, *Pseudomonas*, *Chromobacter*, *Achromobacter*, *Alcaligenes* and *Flavobacterium*. There are also many coliforms in the treatment processes, but they are not considered indigenous members of the microbial community (Gray, 1989), rather coming in with the *influent* and retained by the process (Tremblay *et al.*, 1996). Filamentous bacteria such as *Beggiatoa*, *Thiotrix* and *Sphaerotilus* genera are also found in the biofilm (Vedry, 1996), as are nitrifying organisms like *Nitrosomonas* and *Nitrobacter* (Bitton, 1994). For a thick biofilm, certain facultative or obligate anaerobic species are susceptible to being buried deep in the anoxic or anaerobic zone of the biofilm. These bacteria include sulfate-reducing bacteria, facultative aerobic bacteria

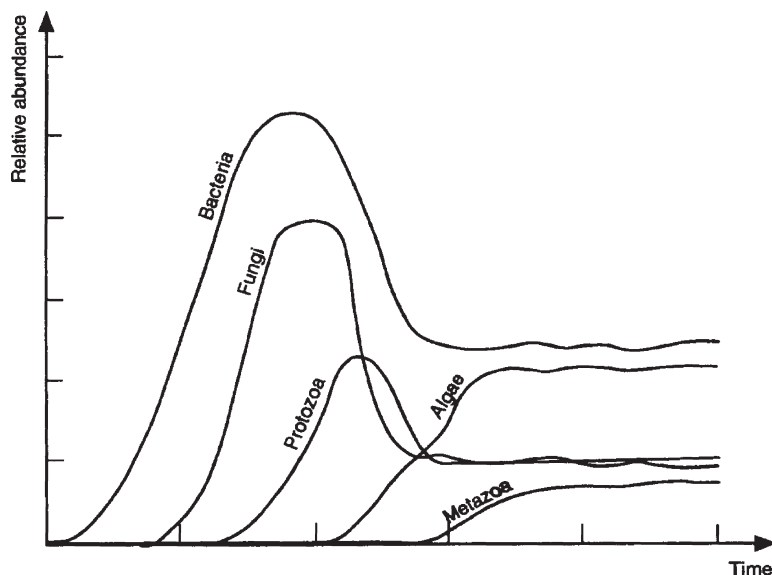


Fig. 19.7 Relative succession of microbes constituting microbial film (adapted from Iwai and Kitao, 1994).

(fermentative) and methanogenic bacteria (Arvin and Harramoës, 1990). However, these anaerobic bacteria are not, of course, dominant in the aerobic processes (Crowther and Harkness, 1975).

Role and nuisance of bacteria. Most bacteria in fixed film processes take part in the removal of the soluble pollutant matter. The colloidal part is adsorbed on the biofilm and hydrolyzed by extracellular enzymes and used later (Bitton, 1994). Certain pH, temperature, aeration and low loading conditions ($0.1\text{--}0.4\text{ kg BOD}_5/\text{m}^3/\text{day}$) can encourage the growth of autotrophic nitrifying organisms. Denitrification can take place in the anoxic zone of the biofilm and methanization, carried out by biofilm organisms, can occur in the anaerobic zone, i.e. close to the support medium.

Generally, bacteria are the dominant organisms in the biofilm, and few species are known to hinder process performance. However, certain species such as *E. coli* or *Nocardia* spp. perform less well than others in removing carbonaceous pollution (James, 1964).

Under certain conditions, filamentous bacteria can develop as a thick biofilm, resisting sloughing or shearing, thus encouraging biofilter clogging (Vedry, 1996). For processes using rotating biological contactors, the presence of organisms, like *Beggiatoa*, increases

biofilm accumulation, thus favoring a biomass surcharge at the disk surface, and a decrease in oxygen diffusion into the biofilm (Metcalf and Eddy, 1991); this situation may generate some operational problems (WEF and ASCE, 1992).

2.3.2 Fungi

Fungi are single- or multi-cellular eukaryotic organisms. In fixed film processes, they play a role of purifier, important for the carbonaceous substrate. In lower quantities than bacteria, about 10^5 and 10^7 CFU/g VSS for fungi and yeast respectively (Le Bihan and Lessard, 1998). Their high hydrolytic potential makes them very competitive against bacteria. Their growth is favored over that of bacteria by cold temperatures, an acid pH (<5) and an industrial or toxic *influent*, especially one that is highly concentrated or contains carbohydrates (Tomlinson and Williams, 1975). Except for yeasts, most fungi are obligate aerobes. The average doubling time of fungi varies between 2 and 14 hours.

The most frequent fungi colonizing fixed film processes are: *Sepedonium* spp., *Subarromyces splendens*, *Ascoidea rubescens*, *Fusarium aquaeductuum*, *Geotrichum candidum* and *Trichosporon cutaneum* (Tomlinson and Williams, 1975). Cooke (1963) identified over a hundred species of fungi in wastewater and in the biomass of biological processes.

Role and nuisance of fungi. Fungi take part in the removal of the carbonaceous substrate in a similar manner to heterotrophic bacteria. However, the conversion rate, much higher than for bacteria, encourages a significant production of biomass for a given quantity of substrate. Too great a proportion of fungi in the biomass favors the formation of a highly resistant biofilm that is difficult to slough. As oxygen can diffuse through fungal protoplasm deep into the biofilm, there is an accumulation of fungal biofilm (Gray, 1989). This characteristic helps the fungal biofilm maintain a greater thickness of aerobic conditions, thus limiting the sloughing effect favored by the deep anaerobic zone.

For trickling filter processes, such as biological aerated filters, these growth characteristics help the development of an abundant biofilm, which could accelerate clogging of the process (Tomlinson and Williams, 1975).

It seems fungi have very specific growth cycles based on the season. The maximum number of fungi is reached in winter and in early spring; they become rarer in midsummer (Tomlinson and Williams, 1975). During the warm period (25°C), only *Furasium*, *Geotrichum* and *Trichosporon* spp. have a doubling time short enough (7–8 hours) to compete with zoogeal bacteria with a doubling time of 10–25 hours (Gray, 1989). As their affinity for the substrate is lower than that of the bacteria, their growth is favored when the *influent* is concentrated (> 280 mg BOD₅/l) and generally at the surface of the filter, where the substrate is more concentrated (Gray, 1989). Thus, for cold temperatures and a concentrated substrate, the fungal biomass is clearly favored in relation to the bacteria.

2.3.3 Protozoa

Protozoa are unicellular eukaryotic organisms widespread in fixed film processes. More than 218 species were identified (Gray, 1989). These organisms, larger than bacteria, are mostly aerobic. They mainly feed on organic particulate matter, bacteria, algae or other *Protozoa*. Certain genera of Mastigophora flagellates are

able to use a soluble substrate, but are less competitive than bacteria and fungi (Bruce and Hawkes, 1983).

Protozoan growth is slower than bacterial growth, the latter have an observed average doubling time of 3 to 22 hours (Pike, 1975; Iwai and Kitao, 1994). Their quantity varies with the type of *influent* and aeration conditions but there are generally between 10^6 and 10^8 organisms/g VSS. Protozoa are divided into three branches: Mastigophora (phytomastigina and zoomastigina), Rhizopoda (heliozoa and amoebae), and Ciliophora (ciliated). A few studies have identified ciliated Protozoa as being the most numerous in the biomass (Curds and Cockburn, 1970; Hoag *et al.*, 1983; Hull *et al.*, 1991). The most abundant ciliated protozoa (Ciliophora) in trickling filter processes are fixed organisms of the genera *Vorticella*, *Opercularia*, *Carchesium* and swimmers like *Aspidisca* and *Chilodonella* (Curds and Cockburn, 1970). These authors also showed that *Rhizopoda* (amoebae), which are slightly less abundant than the ciliates, are frequently represented by the *Arcella* (testate amoebae) genus. Naked amoebae are rather rare in fixed film processes. Mastigophora mostly include *Paranema*, *Bodo* and *Trepanomas* spp.

Role and nuisance of protozoa. Protozoa take on the role of predators of bacteria and take part in maintaining bacterial growth by reducing their population density (Gray, 1989). Certain Mastigophora species are able to assimilate particulate and dissolved organic matter. The absence of Mastigophora in a biomass is an indicator of toxic conditions (Vedry, 1996).

The Rhizopoda found in the biomass of fixed film processes are mainly composed of Thecamibae (testate amoeba). These shelled organisms do not swim, but move by using their pseudopod on the biomass. They feed on mainly fixed, agglomerated, filamentous bacteria or unicellular algae. Ciliates feed on suspended particles, loam and free bacteria. All are very sensitive to a lack of oxygen. Some swimming ciliate species also feed on protozoa and flocculated bacteria. Other species (e.g. *Chilodonella cucullulus*) have an oral morphology, which permits them to feed

on filamentous bacteria and fungi. Fixed ciliates (peritricha) require a support on which to fix, and they feed mainly on free bacteria.

Only a few studies deal with the nuisance caused by protozoa. However, Vedry (1996) shows the inconveniences of an invasion by fixed ciliates, specifically, colonial peritricha. When they invade a biological aerated filter, they can hardly be removed from the support, even with a wash. This fastening capacity increases their number to a point of equilibrium with the turbulence in the environment. According to this author, accumulation of fixed ciliates increases oxygen consumption and head loss of the process. Other studies also mention that the presence of protozoa and other predators could affect the development of microorganisms with a low growth rate, such as the bacteria responsible for nitrification (Lee and Welander, 1994).

2.3.4 *Algae and cyanobacteria*

Several photosynthetic organisms, algae and cyanobacteria, can grow exposed to light at the surface of a biofilm. The difference between algae and cyanobacteria is that algae are single or multicellular photosynthetic eukaryotes, while cyanobacteria (blue-green) are unicellular photosynthetic prokaryotes. The frequently found genera include *Ulothrix*, *Phormidium*, *Anacystis*, *Euglena*, *Stigeoclonium*, *Chlorella* and *Oscillatoria* (Cooke, 1959). Their doubling time is similar to that of protozoa, varying from 7 to 25 hours, depending on the species (Iwai and Kitao, 1994).

Roles and nuisance of algae and cyanobacteria.

Algae and cyanobacteria only play a small role in fixed film processes (Vedry, 1996). Some species, especially *Phormidium cyanobacterium*, develop into multilayers at the surface of the trickling filter and decrease the efficiency of the process, eventually clogging the filter. This layer of photosynthetic organisms also serves as a favorable environment for fly breeding that could cause nuisance during their emergence (Bruce and Hawkes, 1983).

2.3.5 *Metazoa*

Metazoa are multicellular eukaryotic organisms that encompass the phyla of worms (rotifers, nematodes, gastrotricha and oligochaetes) and arthropods (arachnids, tardigrades, crustacea, myriapods and insects). With their relatively high doubling time, between 47 and 238 hours (Iwai and Kitao, 1994), these organisms are only present in an aged biomass. Most of them are aerobic; however, some organisms can survive under microaerophilic conditions (nematodes, oligochaetes).

Roles and nuisance of Metazoa. These predators and detritivorous organisms feed on the biofilm, embrittle it, encourage loosening, decrease biomass volume and accelerate mineralization. Their grazing capacity also helps avoid clogging of trickling filters through biofilm consumption. With their low activity in cold water (winter), trickling filter clogging is often observed during this period, especially in the case of a growing fungal biofilm (Hawkes, 1983). Nematodes are often observed where voluminous, potentially anoxic, bacterial masses with low oxygen permeability are found. Their role in biomass mixing allows the oxygen to reach the anoxic zones inside these bacterial masses.

Metazoa are frequently observed in trickling filter processes, but rarely in aerated biological filter processes. The frequent washings of these filters control the age of the biomass. Thus the presence of oligochaetes, rotifers and nematodes in an aerated biological filter process is an indicator of poor operation of the washing process, which allows the undesirable accumulation of biomass with a high sludge age (Vedry, 1996). Moreover, since they are light, oligochaete dejects could find their way into the effluent of upflow biofilters, thus increasing the concentration of suspended solids.

For arthropods, the *Psychoda* and *Sylvicola* larvae are also excellent biofilm consumers (Bruce and Hawkes, 1983). They are mainly found in non-submerged filters. Their development and emergence into adult insects (filter flies) can cause invasion and nuisance

problems for the trickling filter operators. Flow increase or sporadic immersion of the filter helps control and reduce larval development. However, there are very few data regarding the problems caused by arthropods in wastewater treatment processes.

2.3.6 Environmental effects on biofilm ecology

Some microorganism species, favoring or not proper process operation, will develop with relation to various operational parameters such as: substrate, nutrients, dissolved oxygen, temperature, pH, and hydraulic loading.

Substrate. Substrate composition imposes a selective strain on biomass species (e.g. heterotroph versus autotroph). Normally, soluble compounds such as sugars, acids and amino acids are much more easily assimilated by a large quantity of heterotrophic organisms than are insoluble or partially soluble compounds such as cellulose, lignin or fatty acids with high molecular weights. These substrates are used as secondary substrates by specialized organisms. Moreover, sugars are consumed by many microorganisms, whereas proteins and fats are restricted to more specialized organisms.

Substrate concentration also has an effect on biofilm nature. Generally, growth of an abundant, easy-to-slough biofilm is obtained using a readily oxidizable substrate (Heukelekian and Crosby, 1956). However, according to these authors, more biofilm is accumulated in the process when fed by a weak influent. Seek Park *et al.* (1998) observed that the biofilm at the surface of a rotating biological contactor was thicker, less dense, filamentous and contained more extracellular polymers (EPS) when the process was fed by a concentrated influent. For a diluted *influent*, the biofilm was thin, but very dense.

From the selection aspect, an easily bioavailable substrate (sugar) favors the growth of filamentous bacteria such as *Sphaerotilus natans* (Gray, 1989). A very concentrated influent favors the growth of fungi.

Nutrients. Generally, a C/N ratio value of 18 or less is required for adequate growth and

treatment efficiency. Values above 22 decrease performance and favor the development of filamentous organisms (Hatting, 1963). A fungal biofilm can also develop with an influent with a very high C/N ratio (Tomlinson and Williams, 1975). These conditions are commonly found in the treatment of industrial effluents.

When using a soil to treat wastewater, a high C/N ratio favors the production of extracellular polymers (EPS) and decreases the soil's hydraulic conductivity (Magesan *et al.*, 1999). It seems that similar clogging phenomena could be observed in aerated biological filter processes with a small-sized medium (2–3 mm).

For phosphorus, a C/P ratio lower than 90 to 150 is required to maintain treatment efficiency (Gray, 1989). However, no specific problems related to phosphorus deficiency have been reported in the literature.

Dissolved oxygen. Oxygen is used as a final electron acceptor by aerobic microorganisms and their growth rate increases with increasing dissolved oxygen concentration. In biofilm processes, the anoxic zone normally starts 0.1–0.2 mm below the biofilm surface; in the case of a fungal biofilm, this zone can reach down to 2 mm due to the diffusion of oxygen through the fungal protoplasm (Gray, 1989). In this case, the aerobic layer is thicker and the sloughing caused by the development of the anaerobic zone is delayed and the biofilm thickens. Applegate and Bryers (1991) have shown that a biofilm limited by oxygen sloughs more easily than a carbon-limited biofilm.

Temperature. On average, domestic wastewater temperatures vary between 10 and 20°C and, in general, trickling filter processes are more sensitive to temperature than activated sludge (Bruce and Hawkes, 1983). Influent temperature affects the performance of fixed film processes since this factor has an impact on microbial populations. The first effects are the slowdown of microbial metabolism. Under normal circumstances, the activity doubles with each temperature increase of 10°C between 5 and 30°C. Temperature also has an inhibiting effect on nitrification, which is

drastically reduced at temperatures below 10°C (Bruce and Hawkes, 1983).

In addition to decreasing metabolism, cold temperatures favor the accumulation of biofilm in the process (Heukelekian, 1945). Cold winter temperatures decrease the activity of grazing fauna on the biofilm and favor an accumulation of solids in the process (Bruce and Hawkes, 1983), a process critical for biofilm clogging (Hawkes, 1983). Moreover, at cold temperatures, several fungi grow more quickly than bacteria and thus can dominate in the biofilm.

pH. pH effects microbial growth. Fungi develop between a range of pH (2.5–9) wider than that of bacteria, and are favored at pH < 6.5. Bacteria dominate under neutral pH conditions (Gray, 1989). The optimal pH for oxidation of carbonaceous compounds is about 6.5–8.5, which corresponds to the pH of domestic wastewater. For nitrification, an optimal pH of 7.5 and 8.5 has been reported (Bock *et al.*, 1986).

Hydraulic loading. The shear forces generated by the water flow favors sloughing of the biofilm, although this depends on the nature and type of biofilm. A fungal biofilm is more resistant to shear than a bacterial biofilm (Tomlinson and Williams, 1975). Nitrifying biofilms are thinner and denser, but slough more easily than heterotrophic biofilms (Oga *et al.*, 1991).

In rotating biological contactor processes, sloughing is influenced by rotational speed. A rapid flow rate delays initial biofilm formation, but favors biofilm development later on (Heukelekian and Crosby, 1956). In aerated biological filter processes, the flow rate also favors biomass distribution deep into the filter and accelerates clogging (Visvanathan and Nhlen, 1995).

2.4 Comparison between attached and suspended growth

It is important at this stage to make a comparison between suspended and attached

growth processes before discussing hybrid systems. The main characteristics of both types of processes are given in Table 19.6. To summarize briefly the difference between the two types of processes: fixed film processes (TF and RBC) seem to be less efficient than suspended growth processes (activated sludge), but they are more stable. (The main difficulty with AS is to grow a biomass that will separate well.)

Attached microorganisms possess some advantageous properties compared to suspended microorganisms. Often they exhibit:

1. Increased persistence in the system
2. Faster growth rate
3. Increased metabolic activity
4. Greater resistance to toxicity (Senthilnathan and Ganczarczyk, 1990).

An ecological comparison between the process activated sludge and a trickling filter is made in Fig. 19.8. Biomass diversity is greater for the trickling filter process than for the activated sludge, which means a process that is more stable and less subject to environmental conditions.

Most importantly, the two forms of biomass, fixed and suspended, differ in their main parameter, the solids retention time Θ_x , and this may be very advantageous in cases where fast and slow degradable components have to be combined, e.g. organic removal with nitrification (Wanner *et al.*, 1988).

2.5 Hybrid systems

The optimal growth conditions differ for each type of microorganism involved in biological treatment (nitrifiers, heterotrophs, etc.), and several compromises have to be made to enable all the bacteria to achieve as high a degree of activity as possible. The use of hybrid systems can help to achieve that degree of activity. Hybrid systems can be defined by two types of processes, i.e. systems that use: (1) a biofilm reactor, usually a trickling filter (TF), and an activated sludge (AS) process in series, and (2) a biomass support system (fixed or mobile) immersed in an activated sludge reactor.

TABLE 19.6 Biofilm systems vs suspended-growth systems in wastewater treatment (adapted from Shieh (1987) and Bryers (1987))

<i>Biofilm systems</i>	<i>Suspended-growth systems</i>
Microorganisms are retained within the biofilm attached to the media.	Microorganisms are suspended in the wastewater by means of mixing and/or aeration.
Fair protection against the adverse effects of toxicants in the influent.	Sensitive to the adverse effects of toxicants in the influent.
Inter- and intra-phase mass transfer is significant because the biofilm structure tends to retard the rate of transport of substrate through it. Liquid-biofilm interface forms another resistance to the transport of substance across it. Therefore, the biofilm systems are heterogeneous systems.	Inter- and intra-phase mass transfer is insignificant. Vigorous mixing and/or aeration reduces the thickness of the liquid film surrounding the flocs and the size of the flocs which, in turn, reduce the effects of mass transfer. Therefore, the suspended-growth systems are treated as homogeneous systems.
The system performance is not affected by the performance of the secondary clarifier. The biomass in the reactor is maintained through the attachment of microorganisms on the media surface and the subsequent growth of biofilm.	The system performance is intimately linked to the performance of the secondary clarifier because the maintenance of the desirable biomass concentration in the reactor depends upon the recirculation of concentrated microbial solids from the secondary clarifier with proper thickening function.
Elimination of wash-out restrictions.	Possible wash-out of biomass.
Secondary clarification in some cases may be eliminated because the suspended solids level in the system effluent is very low. This is possible because most of the biomass is retained within the biofilms rather than suspended in wastewater.	Secondary clarification is required to reduce the effluent suspended solids concentration to the acceptable level.
Unified design approaches are not fully developed yet because the biofilm systems are generally complex.	Unified design approaches, such as F/M approach and SRT approach are now well developed and widely employed for practical applications.

2.5.1 Systems in series

Secondary treatment processes combining fixed growth (trickling filter) and suspended growth (activated sludge) systems in series are popular, especially in the USA. These processes offer simplicity of operation and process stability of fixed film processes with the high quality effluent associated with suspended growth processes, and they may also offer economic advantages in comparison to other treatment options (Harrison *et al.*, 1984). Besides this stable performance, sludge with good settling properties is produced with those processes (Harrison *et al.*, 1984), which could be explained as follows (Wanner *et al.* 1988):

- large particles of biofilm, in which filamentous microorganisms are fixed in a matrix of extra cellular polymers, are generated in the preceding biofilm reactor;
- according to a conventional selection theory, preceding biofilm reactor can act as selector.

Various combinations exist, and typical hybrid systems in series are shown in Fig. 19.9. Design information can be found in standard textbooks (Metcalf and Eddy, 1991; WEF and ASCE 1992; Crites and Tchobanoglous, 1998).

These processes have proved to be adaptable to a variety of situations including both large and small plants, cold and warm climates, and a wide variety of effluent requirements (Parker *et al.*, 1993, 1998; Parker and Bratby, 2001). Well-designed and properly operated combined TF and AS plants of all modes are capable of producing secondary effluent quality (30 mg/l BOD and SS) and many can produce even better quality effluents (Harrison *et al.*, 1984). An economic trade-off exists between the more energy-efficient TF/SC and ABF processes and the less capital-intensive RF/AS and BF/AS processes (Grady *et al.*, 1999). The microbiology of such systems is thought to be similar to that of trickling filters and activated sludge processes.

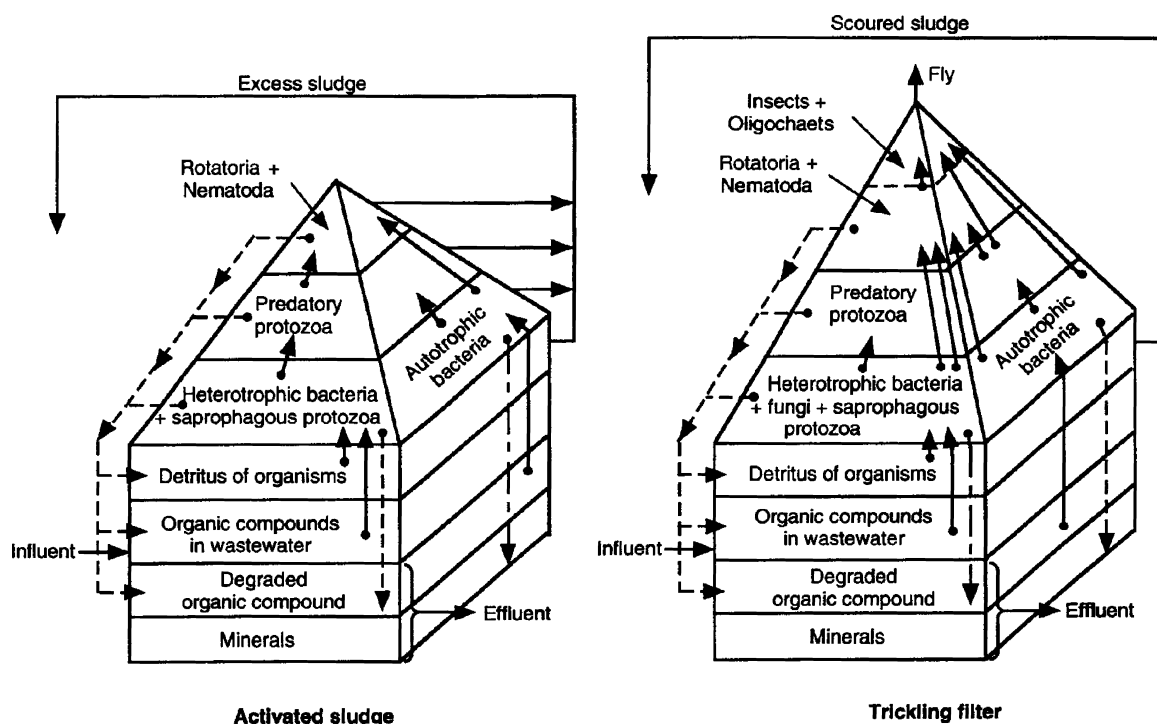


Fig. 19.8 Comparison of biotas between activated sludge and microbial film (adapted from Iwai and Kitao, 1994).

2.5.2 Biomass support systems

Biomass support systems consist of immersing various types of support media in an activated sludge reactor to favor the growth of fixed bacteria. The support can be fixed in the reactor or can consist of mobile media such as foam pads, small carriers, etc. (Fig. 19.10).

These hybrid systems should allow a reduction in the aeration tank volume following the introduction of biomass support to meet a certain objective, and thus an increase in the treatment system stability and performance (Gebara, 1999). The main advantages of these systems are improved nitrification and an increase in sludge settleability (Wanner *et al.*, 1988; Muller, 1998). Nicol *et al.* (1988), based on mathematical simulation, also suggested the following advantages: ability to resist failure from large hydraulic surges; stable nitrification under transient inhibitory conditions resulting from temperature changes, hydraulic surges, and/or toxic chemicals; and the ability to

establish stable operating conditions with respect to both carbon oxidation and nitrification at short hydraulic retention times and low sludge ages.

Kinetic analysis of such systems have been made and models proposed (Nicol *et al.*, 1988; Hamoda, 1989), but no clear design guidelines exist for such systems. However, one point that must always be remembered when considering an activated sludge process with biomass supports is that, because of a higher biomass concentration, the volumetric oxygen requirement will be greater in these systems than in conventional activated sludge. Hence, reductions in aeration tank volume proposed for these systems will be limited by the aeration system's ability to handle higher volumetric oxygen demands (Nicol *et al.*, 1988).

These types of hybrid systems are mainly used for the upgradation of activated sludge systems, either to increase the effective biomass (higher treatment capacity) or to implement nitrification. Many examples can be found in the literature (Morper, 1994; Randall and Sen,

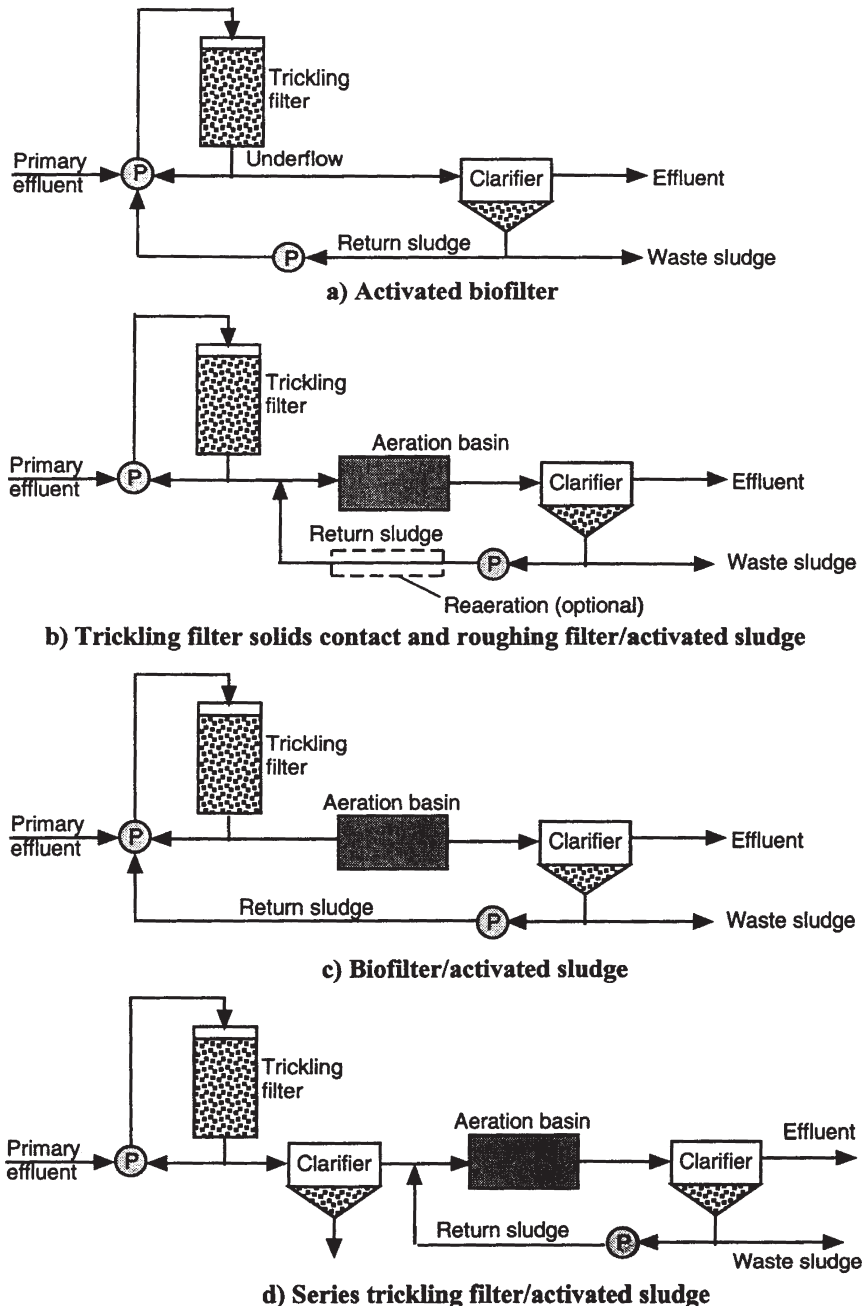
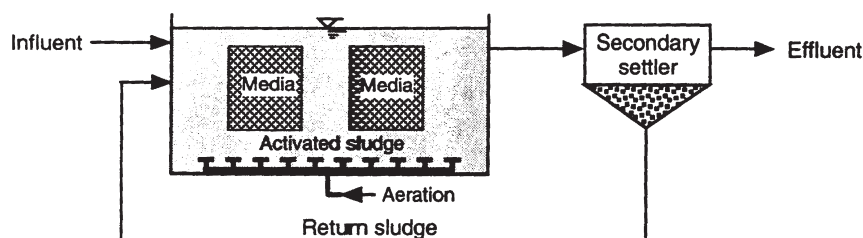


Fig. 19.9 Typical combined attached- and suspended-growth aerobic treatment system flow diagrams. (a) Activated biofilter; (b) trickling filter solids contact and roughing filter/activated sludge; (c) biofilter/activated sludge; (d) series trickling filter/activated sludge (adapted from Crites and Tchobanoglous, 1998).

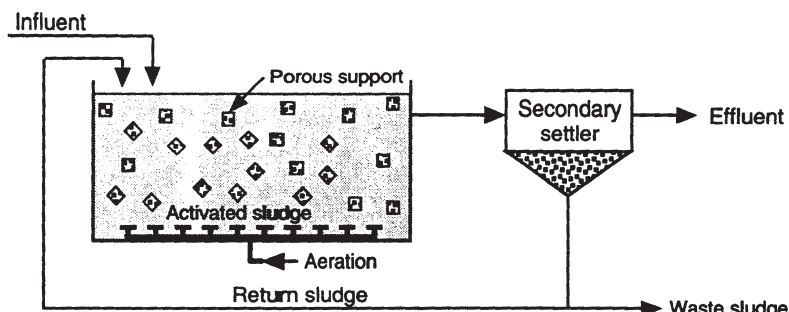
1996; Jones *et al.*, 1998; Muller, 1998; Rusten and Neu, 1999). Moreover simultaneous denitrification, probably in the deeper layers of the biofilm, has been observed in plants with stable nitrification (Muller, 1998). These types of systems can also be operated in sequencing

batch mode to permit phosphorus removal (Garzon-Zuniga and Gonzalez-Martinez, 1996; Wang *et al.*, 1998).

Unfortunately, not much has been written on the microbiology of such systems. Most studies rapidly qualify the biomass. For example,



a) Immersed media systems



b) Porous biomass support systems

Fig. 19.10 Hybrid biomass support systems.

Gonzalez-Martinez and Duque-Luciano (1992) described the biomass adhering to their support as a mixture of biofilm and filamentous growth, with length between 1–5 cm, attached to the plastic material from one end. However, Reddy *et al.* (1994) drew an interesting comparison between the organisms in the activated sludge liquor and those on the media of their process (Table 19.7). It is to be observed that rotifers and

free swimming ciliates are abundant in the biofilm which could explain the low SS and BOD in the process effluent (Muller, 1998).

3 CONCLUSION AND PERSPECTIVES

Growth of biofilms is a natural process that has been engineered in many ways to treat wastewater. Fixed film processes, developed mainly for the treatment of municipal wastewaters, have been presented here along with their microbiology.

Attached growth processes are often compared to, and seen as competitors of, suspended growth processes, but it is interesting to note that currently there is a tendency to couple both processes to make better use of each technology. This in a sense makes the point that there is no such thing as *the technology*, but that each technology has its advantages and disadvantages and that it is applicable in some cases but not in others.

So far as future trends are concerned, it is always difficult to predict exactly what is going to happen. However, increasing standards for

TABLE 19.7 Organisms found in a hybrid system (from Reddy *et al.*, 1994)

Organisms	Activated sludge MLSS		CAPTOR media MLSS	
	Abundance	Rank	Abundance	Rank
Rotifers	Few	4	Abundant	1
Free swimming ciliates	Abundant	1	Very common	2
Stalked ciliates	Very common	2	Common	3
Flagellates	Common	3	Rare	5
Nematodes	Rare	5	Rare	5

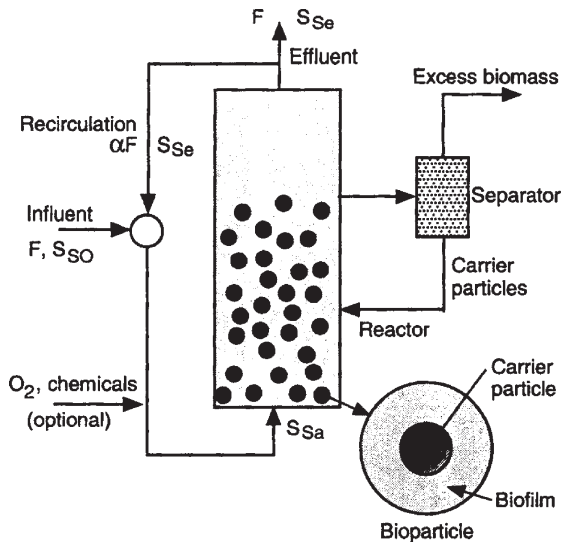


Fig. 19.11 Schematic diagram of a fluidized bed reactor (from Grady *et al.*, 1998).

nutrient and xenobiotic removal, along with more rigorous noise and odor control, have stimulated the development of new intensive advanced technology processes (Lazarova and Manem, 1994).

One of these intensive processes that is attracting increasing attention is the fluidized bed reactor (FBR), which consists of a biofilm growing on small carriers kept in suspension in the fluid by the drag forces associated with the upward flow of water (Fig. 19.11). FBR are a more sophisticated version of the immersed mobile biomass system presented earlier. Their main advantages are:

1. high removal efficiency of carbon and nitrogen through large amounts of fixed biomass with a low hydraulic retention time
2. no clogging
3. better oxygen transfer
4. reduced sludge production (Lazarova and Manem, 1994).

However, the following problems are met at large scale fluidized-bed facilities (Chuboda *et al.*, 1998):

1. a problematic media separation and biofilm control system

2. a lack of suitable bed height control
3. a very high sensitivity of the inlet distribution system to clogging
4. a relatively uncertain control of a stable fluidization.

Despite these problems, many systems have been built, but mainly as two-phase fluidized beds for the treatment of industrial wastewaters (Cooper and Wheeldon, 1980; Sutton and Mishra, 1991, 1994). Time may come when they will be used for municipal wastewater treatment.

Finally, the development of intensive processes may be of interest, but there is a growing need for the development of extensive processes (low cost and low operational requirements) for on-site and small community wastewater treatment. In that perspective, packed-bed reactors using organic media, such as peat or compost, are being studied and used on site. Couillard (1994) carried out a thorough literature review on this subject, and he concluded that the peat bed does have some disadvantages, such as color, high COD and low pH in the effluent, but that, despite these, the effluent quality satisfies criteria for discharge into most receiving waters at a cost outcompeting most processes. However, their low hydraulic loading rate makes them at the moment only applicable for small flows. Laboratory scale and field studies have demonstrated the good performance of organic media (Lens *et al.*, 1994; Talbot *et al.* 1996), and it is likely that they will be used more and more frequently in the future.

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Biofilm formation and its role in fixed film processes

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1 BIOFILMS: CONCEPT AND RELEVANCE

A biofilm is a surface-attached gelatinous matrix composed of microorganisms, the extracellular polymers they excrete (EPS) and foreign substances such as adsorbed molecules and small abiotic particles. Such a matrix is highly hydrated, often containing more than 90% water (mass percentage) and its properties depend on three kinds of factors: biological (microbial species), chemical (fluid composition in contact with the biofilm), and physical (hydrodynamic and thermal conditions under which the biological layer is formed). The thickness of biofilm layers can reach a few millimetres, or even centimetres, but typically ranges from 10 micron to 1 mm. Cell concentrations inside biofilms are commonly in the range of 10^7 – 10^9 cells/cm². EPS often represent more than 50% of the total organic mass of the biological layer. A great diversity of microbial species is found in biofilms in quite different environments, as illustrated in Table 20.1. As can be seen, bacteria predominate in biofilms, but algae and fungi are also present.

Microbial films can have beneficial or detrimental effects, depending on where they build up. Examples of the former can be found in fixed biomass reactors used in wastewater treatment (Harremöes, 1978) or in the production of ethanol (Dempsey, 1990) and citric acid (Briffaud and Engasser, 1979). Unwanted biofilm formation (biofouling) occurs on the surfaces of heat exchangers (Bott, 1995), cooling

water towers, valves, tubes and ships' hulls, increasing the resistance to flow and to heat transfer and inducing/enhancing corrosion (Geesey, 1991). Some biofilms can also become a serious problem for human health when they attach on lungs, teeth, urinary catheters, etc. (Costerton *et al.*, 1987). The present chapter will focus essentially on biofilm growth and characteristics in industrial equipment.

2 BIOFILMS VERSUS DISPERSED CELLS

Microorganisms live in dispersed suspensions, in flocs and in attached films. The aggregated forms are clearly predominant in nature (Lappin-Scott and Costerton, 1995) and they probably constitute a more efficient way of surviving in many environments. Two words can be used to explain why microbes may choose to form biofilms or other aggregates, instead of living as dispersed cells: proximity and protection.

The distances between microorganisms in biofilms are considerably smaller than in suspension and this leads to the creation of microenvironments with specific nutrient concentration gradients, pH's, electrical charges distributions, proton concentrations, etc., which affect the metabolism of microorganisms. Natural selection will lead to growth of certain species as a result of the specific local conditions created within the biological matrix. Microbial growth rates may sometimes be lower in biofilms than in suspensions, but

TABLE 20.1 Examples of microorganisms present in biofilms

Processes	Microorganisms
Cooling water systems	<i>Pseudomonas</i> spp., <i>E. coli</i> , <i>Staphylococcus</i> , <i>Desulfovibrio</i> , clostridia, green algae
Biological wastewater treatment	<i>Nitrosomonas</i> , <i>Nitrospira</i> , <i>Nitrobacter</i> , <i>Pseudomonas</i> , <i>Methanotrix</i> , <i>Methanosarcina</i> , <i>Clostridium</i> , <i>Desulfovibrio</i> , <i>Staphylococcus</i>
Food processing	<i>Legionella</i> , <i>Salmonella</i> , <i>Listeria</i>
Chemical and extractive industry	<i>Acetobacter</i> , <i>Thiobacillus ferrooxidans</i> , <i>Thiobacillus thiooxidans</i>
Lungs	<i>Pseudomonas aeruginosa</i>
Urinary catheters	<i>Providencia stuarti</i> , <i>E. coli</i> , <i>Pseudomonas aeruginosa</i> , <i>Klebsiella pneumoniae</i>
Teeth and mouth	<i>Prevotella intermedia</i> , <i>Actinobacillus</i> , <i>Streptococcus sanguis</i> , <i>Candida albicans</i>

the production of extracellular polymers is normally higher, resulting in a protective network within which the cells are entrapped.

This leads us to the second specificity of biofilm matrices: protection. The exopolymers, often composed of polysaccharides and glycoproteins, contribute decisively to both the adhesion of microorganisms to the initially 'clean' surface and the internal cohesion of the film. Simultaneously, these biopolymers act as a protective network that makes diffusion of toxic substances more difficult and attenuates the effects of chemical and hydrodynamic aggression from the environment. Additionally, polymers may enhance the adsorption of nutrient molecules and the accumulation of particles in the inner zones of the biofilms. Organic particles, for instance, can be retained long enough inside the matrix in order to be biologically degraded.

It should be stressed that biofilms grown in nature and industrial equipment usually contain a great variety of microbial species and that such variety is stimulated by the existence of zones with distinct microenvironments. One example is the creation of anaerobic zones far away from the water-biofilm interface and the development of aerobic species near this interface. Another example is the simultaneous presence of nitrifying and anoxic denitrifying bacteria in the same biofilm, allowing nitrogen removal (from ammonium to nitrogen gas) in the reactor which otherwise would be quite difficult to achieve. The corrosion of metallic surfaces is favoured by the development of

sulphate-reducing bacteria underneath the aerobic layers adjacent to the aqueous environment. These cooperative mechanisms constitute clear advantages over the planktonic growth mode from the point of view of the microbial world.

The protective biopolymer network also acts as a diffusion barrier to the transport of substrates, which explains in part the smaller microbial growth rates sometimes found in biological films.

Thiobacillus ferrooxidans showed growth rates 70 times lower in biofilms on glass particles than in suspended cultures (Karamanev, 1991). However, this is not a general rule, particularly when surfaces are prone to adsorb substrates: Davies and McFeters (1988) proved that *Klebsiella oxytoca* growth rate was much higher (10 times) when attached to activated carbon particles than in liquid suspension when the substrate was glutamate; this did not happen when glucose was used as substrate, because glutamate adsorbs easily to the activated carbon, opposite to glucose.

An important factor that affects the behaviour of cells in microbial films as compared to suspended systems is the attachment process. The presence of a solid surface was found to increase the production of exopolymers (e.g. Vandevivere and Kirchman, 1993). Some authors concluded that adhesion seems to trigger the expression of genes that control the excretion of the biofilm polysaccharides (McCarter *et al.*, 1992; Davies *et al.*, 1993). Adhesion may in fact modify the phenotype

expression of a very significant fraction of the cell proteins. An increase in the liquid viscosity was found to stimulate the production of lateral flagella in *Vibrio parahaemolyticus* (McCarter *et al.*, 1992), as a result of special gene expression induced by the presence of a solid surface.

3 BIOFILM PROPERTIES: ARCHITECTURE AND COMPOSITION

The internal structure of biofilms is characterized by a marked heterogeneity: the biological material is organized in clusters containing cells and the excreted polymeric network, while channels and pores filled with the ambient liquid occupy the free spaces between the clusters. This liquid is not necessarily stagnant and, in fact, convective flows were measured within the channels (Stoodley *et al.*, 1994). Each cluster can contain layers with different microbial species, different polymer compositions, different densities of active cells, etc. Often, particularly in aerobic heterotrophic biofilms, filamentous structures ('streamers') protrude out of the film into the external liquid. Stacks of cells have been often found to extend from a thin basal biofilm attached to the solid surface (Keevil and Walker, 1992).

The physical structure of microbial films changes with time and, in 'older' biofilms, biopolymers were observed to form bridges between clusters and increase the density of the fixed biomass (Fig. 20.1).

Nowadays, new analytical tools, based on the use of molecular probes together with confocal laser scanning microscopy, allow detailed studies of microbial population distributions inside biofilm matrices (Wobus *et al.*, 2000; Neu, 2000) and contribute to establish sound relations between the operating conditions of reactors and the biofilm properties.

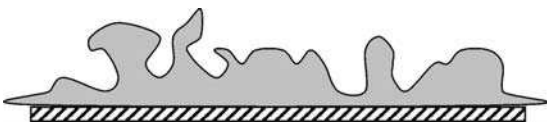


Fig. 20.1 Schematic representation of biofilm structure.

For example, in a combined nitrification and organic carbon removal process, the use of FISH techniques (fluorescence *in situ* hybridization) confirmed that heterotrophic bacteria developed on the top of the nitrifying biofilm under oxygen limiting conditions (Nogueira *et al.*, 2002). In this work, curious interactions between microbial growth and physical phenomena were found: the nitrifying efficiency of the biological reactor decreased significantly when the reactor was operated at high values of the hydraulic residence time, contrary to the low residence time operation. This could be due to the presence of greater amounts of heterotrophs in suspension at higher residence times and therefore to a higher liquid viscosity; ultimately, it led to a decrease in the liquid shear stress and to the formation of a thicker layer of heterotrophs on top of the nitrifying biofilm, inhibiting the activity of the ammonium and nitrite-reducing bacteria located in the inner zones.

Similar techniques were applied to observe how the density of cells varies from the substratum to the biofilm-liquid interface. The results were not always coincident: in some cases, higher cell densities were found near the solid surface (Lawrence *et al.*, 1987; Lawrence and Korber, 1993; Kuehn *et al.*, 1998), whereas in other studies the reverse situation was detected (Neu and Lawrence, 1997).

4 HOW BIOFILMS GROW

The main processes involved in the build-up of biofilms are:

1. formation of the so-called 'conditioning film', a very thin (mono-) layer of organic molecules and ions on the adhesion surface
2. transport of microorganisms (and other particles) to the surface
3. microbial attachment and growth (see also process 5)
4. substrate transport through the external liquid medium and through the biofilm matrix
5. biological reaction at the 'active sites' inside the biofilm, i.e. at the active microbial cells

- that consume the substrate and produce not only new cells but also exopolymers
6. mass transfer (from the biofilm back to the liquid) of the products resulting from the biological reactions within the biolayer
 7. detachment of parts of the biofilm by erosion and abrasion (continuous processes) and sloughing off (sporadic events).

The last process (detachment) causes loss of fixed biomass while the other six processes contribute directly or indirectly to the growth of the biofilm layer (Fig. 20.2).

The initial conditioning film is established in a few minutes following the transport and adsorption of organic molecules contained in the liquid, some resulting from the decomposition of organisms such as algae, bacteria, small plants, etc. These molecules may form polymeric chains that interact with the exopolymers on the surface of microorganisms and form strong bridges that stabilize cell adhesion. The formation of the first microbial layer facilitates the subsequent adhesion of other cells and abiotic material.

The main substrate transport mechanisms from the liquid to the biofilm surface are molecular diffusion (in stagnant or laminar flowing media) and eddy diffusion (when turbulent flow prevails). Within a mature biofilm, molecular diffusion is predominant, but convective flows contribute to enhance mass transport in the biofilm channels.

Once the substrate molecules reach the cells inside the matrix, the production of cellular

biomass and of extracellular polymers leads to the growth of the dry mass of the biofilm while its thickness also increases. Usually, the effect of the shear stress (and other detachment mechanisms) tends to level out the growth rate of the fixed biomass and a final 'steady-state' thickness is reached, although subject to periodic fluctuations. Sometimes, the production of gases as a result of microbial metabolism creates pockets of higher pressure inside the matrix and eventually causes the detachment of large portions of the attached biomass. This may happen, for instance, in biofilms containing a consortium of nitrifying and denitrifying bacteria: the latter tend to build up on the top of the biofilm in order to utilize the oxygen dissolved in the water, while the anoxic denitrifiers tend to be located near the initial adhesion surface. In this case, the conversion of ammonium to nitrate by the nitrifiers, followed by the reduction of nitrate to nitrogen gas creates bubbles inside the biofilm that break and disrupt the biomass structure.

Abrasion due to particle collisions is considered a determining mechanism in the detachment of fixed biomass from carrier particles in turbulent air-lift reactors (Gjaltema, 1996).

It should be stressed that the first six processes mentioned above are consecutive (in series), meaning that the slowest step will control the rate of the growth mechanisms. More often than not, one of the following three processes is the controlling one: mass transfer in the liquid, mainly in laminar flow situations;

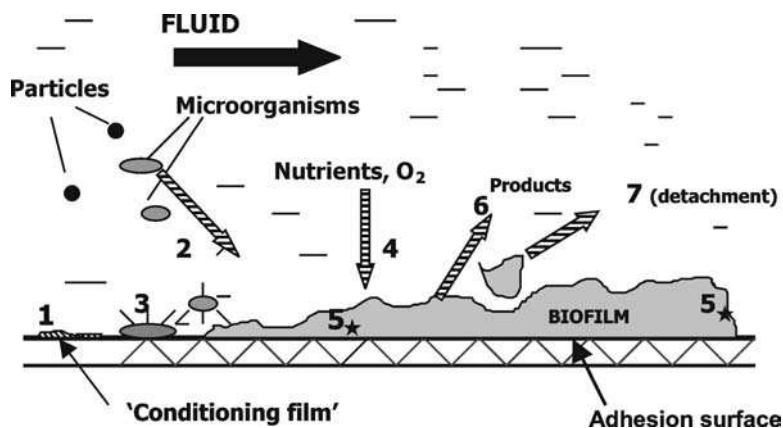


Fig. 20.2 Phenomenology of biofilm formation.

mass transfer through the biofilm; biological kinetics within the biofilm. However, detachment is acting simultaneously, as a parallel process. Therefore, the rate of biofilm formation results from the competition between the overall growth process (comprising the first 6 processes, controlled by one of them) and the detachment process. The faster of these two overall processes will condition the rate at which biofilm builds up. In turbulent flows, for example, the detachment rate is frequently controlling the growth rate of the attached layer.

A typical curve of biofilm growth over time is shown in Fig. 20.3.

The curve in Fig. 20.3 can be modelled by a simple overall phenomenological equation:

$$m_f = m_f^\infty [1 - \exp(-b \cdot t)] \quad (1)$$

where:

m_f = mass of attached biofilm per unit surface area at time t

m_f^∞ = maximum mass of attached film (for $t = \infty$, pseudo-steady state)

b = empirical parameter proportional to the hydrodynamic forces acting upon the biofilm surface and to the amount of attached biomass.

The reciprocal of b ($1/b$) increases with the cohesiveness of the biofilm, which means that $1/b$ is a measure of the biofilm resistance to detachment. The above expression can be derived by considering a few assumptions (Melo and Vieira, 1999; Melo and Oliveira, 2001), one of them being that the fraction of cellular biomass in the biofilm decreases as the biofilm grows. This assumption is justified by the fact that the cells in each layer or cluster

use the substrate to produce living material (new cells) and inert material (polymers) and that these exopolymers are not able to create any new mass (Pereira *et al.*, 2002).

This simple overall model may be used if the parameters m_f^∞ and b are experimentally correlated with the environmental (chemical, biochemical and physical) conditions under which the biofilm is grown. In recent years, structured multidimensional models using powerful computer tools based on 'cellular automata' techniques were developed to describe the growth, morphology and composition of biofilms over time (Picioreanu *et al.*, 1999; Noguera *et al.*, 1999; Hermanowics, 1999). Such models, as many others, including the very simple one presented above, still require basic experimental information on the values of diffusion and kinetic parameters or related variables.

Several factors affect the formation and properties of microbial films. Among others, the following should be emphasized:

1. characteristics of the microbial species and strains
2. composition and roughness of the surface material where the microorganisms attach
3. liquid composition, its pH, temperature, ionic strength
4. hydrodynamic features of the fluid, such as velocity and turbulence.

Microbial species that produce greater amounts of extracellular polymers can more easily attach to solid surfaces. Incorporation of small inorganic particles, such as clays, in the biofilm structure can make them more resistant to external aggressions (Vieira and Melo, 1995). Surface electrical charges of microbes and solid surfaces are affected by pH and, as such, increase or reduce repulsive interactions between them. Naturally, pH has a marked influence on microbial metabolism, which is thus dependent on the specific microenvironments created inside the biofilm matrix. The carbon/nitrogen ratio in the medium affects the ability of microorganisms to attach to solid surfaces, since it conditions the production of extracellular polymers (Veiga *et al.*, 1992).

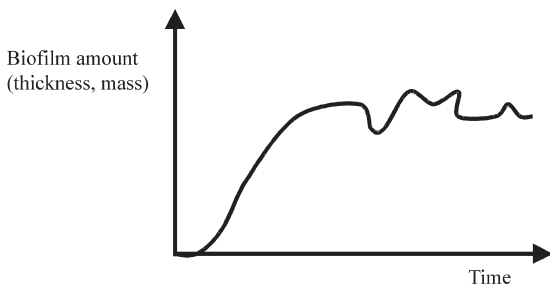


Fig. 20.3 Biofilm growth curve.

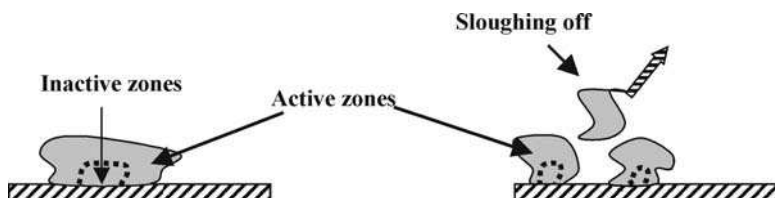


Fig. 20.4 Sloughing off of attached biomass.

Fluid velocity causes a variety of effects:

1. substrate transport rates increase with fluid velocity and turbulence
2. on the other hand, detachment rates are enhanced by higher liquid velocities resulting in thinner biofilms
3. additionally, higher fluid velocities increase the compactness of the biological matrix, which in turn affects internal substrate diffusion (Melo and Vieira, 1999; Vieira and Melo, 1999).

It is not easy to predict which will be the overall effects of the velocity field on the properties and activity of biofilms. Fig. 20.4 is an illustrative example of what may happen when a biofilm is subject to low fluid velocities and high substrate concentrations: it will probably become too thick, and the substrate will not be able to reach its inner zones favouring the appearance of biologically inactive mass. This will enhance the sloughing off of attached biomass, which will produce instability in biofilm performances. In recent years, design of biofilm reactors has favoured turbulent, high shear stress processes that minimize such instabilities (Tijhuis et al., 1995; Lazarova and Manem, 1997).

5 BIOFOULING IN INDUSTRIAL WATER SYSTEMS

Problems caused by biofouling in industry are particularly relevant in water systems, such as cooling water circuits (cooling towers, heat exchangers), paper mill operations, reverse osmosis membranes (Melo and Bott, 1997; Flemming, 1997). Biofilms create additional pressure drops (resistance to flow), increased thermal resistances in heat exchangers and

reduced permeability in reverse osmosis membranes. Water used in industry (taken from rivers, lakes, bore holes) contains various macromolecules that result from the breakdown of living material and these substances not only adsorb onto the equipment surfaces, but also act as nutrients to the living cells, thereby starting the biofilm growth process. In cooling water systems, temperatures in heat exchangers are usually quite suitable for microbial growth (from 10–15°C at the inlet to 30–40°C at the outlet).

Biofilm control is an important aspect of industrial equipment where water flows at moderate temperatures. Use of appropriate velocities (above 1 m/s), temperatures and surface conditions can reduce biofouling effects, but often will not be enough, and mechanical and chemical methods will have to be considered to mitigate biofouling effects (see detailed analysis in Bott, 1995). The former include the physical removal of biofilm from the surface, which can be applied on-line by periodically circulating sponge rubber balls (or similar devices) through the tubes during normal operation, or off-line by shutting down the equipment and cleaning it with high pressure water jets or manual procedures when feasible (e.g. in plate heat exchangers). Use of modified solid surfaces (well polished metals or coated with specific polymeric molecules) also reduces the build-up of biofouling layers.

Another way of minimizing biofilm development is to apply disinfectants (biocides) to kill or inactivate the microorganisms. Chlorine is still the preferred biocide, but its use has been under attack because of the carcinogenic by-products of chlorine reactions with organic matter. Hydrogen peroxide and ozone, although more expensive, may prove to be more environmentally acceptable alternatives.

Other techniques based on the use of surfactants and phages (biological predators) are important contributions to biofouling mitigation, but some are still under scientific, technological and economical scrutiny. A methodology based on the use of a mild biocide (carbamate solution), which simultaneously has an aggregative action, was proposed in order to reduce biofouling problems in paper and pulp plants (Pereira *et al.*, 2001). The idea in this specific situation was to aggregate the bacteria before they had time to form slimy EPS layers that eventually incorporate in the cellulose pulp and cause paper machine shutdowns and affect paper quality.

6 BIOFILMS IN WASTEWATER TREATMENT

Biofilm reactors contain particles of the carrier or support material where the film is attached. The particles are sometimes porous and the biofilm also develops within the pores. The wastewater to be treated flows in direct contact with the biofilm allowing nutrients and metabolic products to be exchanged between the film and the liquid. Supports are made of plastic material of a great variety of shapes and dimensions, and also of natural materials such as sand and basalt particles. The specific area of the carriers has increased in the last decades from around $100 \text{ m}^2/\text{m}^3$ to more than $500 \text{ m}^2/\text{m}^3$.

For the cells to attach to the carrier surfaces in significant amounts, the residence time of the fluid in the reactor should be smaller than the replication time of the cells. In such cases, the cells will tend to adhere to the supports (provided they are good exopolymer producers) to avoid being washed out of the reactor. Most of the biofilm reactors also contain suspended biomass, some of it resulting from the detachment of attached biomass (which has to be purged periodically).

There is a wide range of biofilm reactors available for wastewater treatment (Melo and Oliveira, 2001). They are mostly continuous reactors, the main exception being the sequencing batch biofilm reactor (SBBR) consisting of a tank, which is periodically filled with the feed

liquid and discharged (Wilderer, 1995). Continuous reactors include:

1. trickling filters, where the liquid is split into fine streams and percolates downwards through the biofilm while the air flows upwards
2. rotating disc contactors, with the biofilm attached on the surface of vertical discs rotating in the liquid medium
3. submerged beds with biofilm particles immersed in the liquid (upflow or downflow)
4. fluidized beds, where the biofilm layer forms around small solid particles which are kept under fluidization conditions
5. moving beds, that consist of an expanded bed of particles circulating throughout the equipment together with the fluid (often with gas bubbles too), such as the air-lift reactor and the circulating bed reactor
6. membrane reactors, with microbial films attached to microporous membrane surfaces allowing the supply of nutrients and oxygen to both sides of the film.

These reactors are described in a number of publications (Metcalf and Eddy, 1987; Tijhuis *et al.*, 1994; Wilderer, 1995; Lazarova and Manem, 1997; Nogueira *et al.*, 1998). Reactors containing dense microbial granules without support particles, such as UASB (upflow anaerobic sludge blanket) are usually treated as biofilm reactors as well (Letting and Hulshoff-Pol, 1992; Brito and Melo, 1997).

Fig. 20.5 schematically presents examples of some biofilm reactors.

Biofilm reactors used in wastewater treatment are still often designed on the basis of an empirical parameter (the 'eliminated load'), the values of which are known from previous practical experience, without taking into account any phenomenological approaches. Basically, the following mass balance is used:

$$V_R = \frac{Q \cdot (S_1 - S_2)}{B_v} \quad (2)$$

V_R is the reactor volume, Q the volumetric flow rate of wastewater, S_1 and S_2 are the inlet

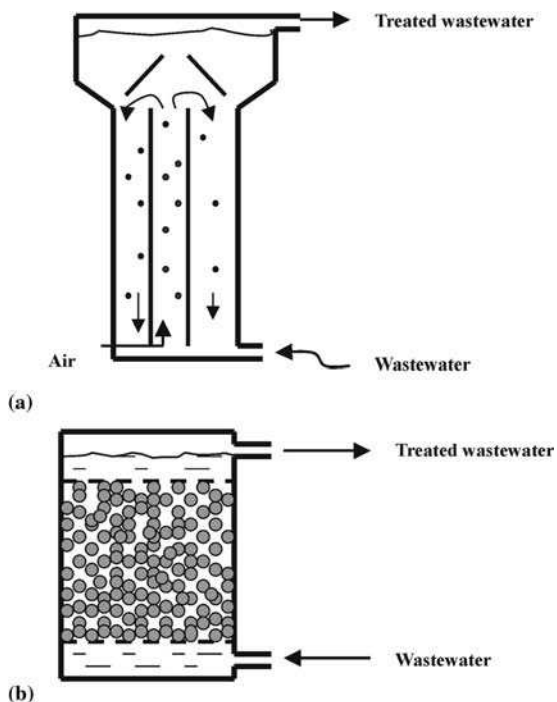


Fig. 20.5 Examples of two biofilm reactor configurations: (a) air-lift reactor; (b) submerged up-flow fixed bed.

and outlet substrate concentrations and B_v is the mass of substrate consumed per unit time and unit volume of the reactor (eliminated load).

A more rational approach to estimate B_v or an analogous rate parameter, is to model it in terms of diffusion-reaction concepts similar to the ones developed in heterogeneous catalysis (Harremöes and Henze, 1995). Integrated equations for these models are available for zero order and first order biological kinetics, and are summarized in Table 20.2. They relate the rate of substrate consumption per unit area of biofilm (r_A) to substrate concentration (S), external mass transfer coefficient (k_m), substrate diffusivity within the biofilm (D_f), intrinsic reaction constants (k_{1f} -first-order and k_{0f} -zero order) and biofilm thickness (L_f).

Parameter B_v (eliminated load) is related to the substrate consumption rate (r_A) by:

$$B_v = r_A \frac{A_f}{V_R} \quad (3)$$

where A_f is the surface area of the biofilm, i.e. the interfacial mass transfer area. If the microbial film is attached to a spherical particle (e.g. in a fluidized bed or an air-lift reactor) and is not too thick, it can be considered as a flat biofilm and its surface area will be:

$$A_f = \frac{3V_R(1 - \varepsilon)}{L_f + r_p} \quad (4)$$

where L_f is the thickness of the biofilm and ε is the bed porosity (ratio of the liquid volume over the reactor volume). For thin biofilms, $L_f \ll r_p$ and the denominator can be reduced to r_p .

Taking into account Equations 2–4, together with Equations 5–12 in Table 20.2, the reactor volume can be estimated as indicated in Table 20.3.

The external mass transfer coefficient can be calculated for many types of reactors by appropriate correlations (see textbooks on Transport Phenomena and Reaction Engineering). However, there is not yet an established methodology for the prediction of the following variables *as a function of the reactor operating conditions*:

1. reaction constants (the reaction constants depend also on the knowledge of the *mass of active cells* and of the *biomass yield* inside the microbial film)
2. internal diffusivities
3. biofilm thickness.

The joint efforts of experimental researchers and mathematical modellers are required to obtain this type of information in different practical situations.

Harremöes (1978) reported that in many wastewater treatment reactors biofilms were partially penetrated and the apparent reactions were of half order. Values of the 'apparent half-order constant' ($k_{1/2}^{app}$) were presented for several pairs of microbial species-substrates, although the specific hydrodynamic and chemical conditions were not fully known. Table 20.4 shows a list of some of those practical values.

The 'apparent reaction constants' differ from the 'intrinsic' ones because the former

TABLE 20.2 Main equations of the diffusion-reaction model for two limiting cases: zero order and first order intrinsic biological kinetics

First order intrinsic kinetics	Zero order intrinsic kinetics
$r_A = \frac{S}{\frac{1}{k_m} + \frac{1}{\eta_i k_{1f} L_f}} \quad (5)$	$r_A = \frac{k_m S}{2\lambda^2} \left[\sqrt{1 + 4\lambda^2} - 1 \right] \quad (8)$
$\eta_i = \frac{\tanh \phi}{\phi} \quad (6)$	$\lambda = \frac{k_m \sqrt{S}}{\sqrt{2k_{of} D_f}} \quad (9)$
$\phi = \sqrt{\frac{k_{1f} L_f^2}{D_f}} \quad (7)$	<p>Special cases When external mass transport is not the limiting step:</p> <p>(a) Biofilm fully penetrated by the substrate:</p> $r_A = k_{of} L_f = \frac{(k_{1/2})_{app}^2}{2D_f} \quad (10)$ <p>(b) Biofilm only partially penetrated by the substrate:</p> $r_A = k_{of} L_f \beta = \sqrt{2k_{of} D_f S} = \quad (11)$ <p>or : $r_A = (k_{1/2})_{app} \sqrt{S}$</p> $\beta = \frac{1}{\phi} = \sqrt{\frac{2D_f S_i}{k_{of} L_f^2}} \quad (12)$

TABLE 20.3 Evaluation of reactor volume based on the expressions of the diffusion-reaction model

Ideal continuous well stirred reactor	Ideal plug flow reactor
$V_R = \frac{Q(S_1 - S_2)(L_f + r_p)}{3r_A(1 - \varepsilon)} \quad (13)$ <p>(For these reactors: $S_2 = S$)</p>	<p>First order reaction:</p> $V_R = \frac{Q(L_f + r_p)}{3(1 - \varepsilon)k_{1f} L_f \eta} \ln \left(\frac{S_1}{S_2} \right) \quad (14)$ <p>with:</p> $\frac{1}{\eta} = \frac{1}{\eta_i} + \frac{k_{1f} L_f}{k_m} \quad (15)$ <p>Zero order reaction with full substrate penetration and no external mass transfer limitations:</p> $V_R = \frac{Q(L_f + r_p)}{3(1 - \varepsilon)k_{of} L_f} (S_1 - S_2) \quad (16)$ <p>Zero order reaction with partial substrate penetration (apparent half-order) and no external mass transfer limitations:</p> $V_R = \frac{Q(L_f + r_p) \sqrt{2S_1}}{3(1 - \varepsilon) \sqrt{k_{of} L_f}} \left[1 - \sqrt{\frac{S_2}{S_1}} \right] \quad (17)$

TABLE 20.4 Apparent half-order constants in the diffusion-reaction model

<i>Process, microbial species and limiting biological factor</i>	<i>Reactor type</i>	$K_{1/2app} 10^5 \text{ (kg}^{1/2}/\text{m}^{1/2}/\text{s)}$
Aerobic, heterotrophic biomass Oxygen (Grasmick <i>et al.</i> , 1982)	Fixed bed	0.12
Aerobic, heterotrophic biomass Toluene (Pederson and Arvin, 1996)	Waste gas trickling filter	0.07–0.11
Nitrification, autotrophic biomass Oxygen (Çeçen and Gönenç 1994)	Submerged filter	0.05–0.10
Nitrification, autotrophic biomass Oxygen (Nogueira <i>et al.</i> , 1998)	Circulating bed reactor	0.15
Nitrification, autotrophic biomass Ammonium Gönenç and Harremões, 1985	Rotating disc	0.06
Anaerobic, methanogenic biomass Acetate (Hamoda and Kennedy, 1987)	Downflow filter	0.12
Anaerobic, methanogenic biomass Molasses (Gönenç <i>et al.</i> , 1991)	Upflow filter	0.04–0.48
Anoxic, denitrifying biomass Nitrate (Jansen, 1982)	Rotating drum	0.02–0.14
Anoxic, denitrifying biomass Nitrate (Watanabe and Ishiguro, 1978)	Rotating disc	0.11

encompass both the biological reaction and the internal mass transfer in the biofilm. The definition of the 'half-order apparent reaction constant' is:

$$(k_{1/2})_{app} = \sqrt{2k_{of}D_f} \quad (18)$$

As regards the prediction of the effective diffusivity in the biofilm matrix (D_f), advances have been achieved in recent years, but their applicability to reactor design is still limited. Published values of the ratio between the effective diffusivity in biofilms and the diffusivity of the same component in water range from 15 to 120% (Melo and Oliveira, 2001). Some authors related biofilm density to diffusivity (Fan *et al.*, 1990; Stewart, 1998). The effect of velocity on biofilm structure and on the effective diffusivity of nutrients was reported by a few authors (Vieira *et al.*, 1993; van Loosdrecht *et al.*, 1995; Brito and Melo, 1999; Casey *et al.*, 2000). The results do not yet allow a unifying theory because, in general, higher velocities produce two opposite effects (thinner but more compact biofilms) on the mass transfer resistance offered by the biological layer. The particular hydrodynamic patterns present around biofilms (laminar versus turbulent flow) seem to have a determinant role not yet duly understood.

Although the above unstructured model and other more sophisticated structured models help in understanding the role of the different variables in the efficiency of substrate removal in biofilm reactors, they are not yet able to provide engineers with values for practical application to reactor design. Alternatively, empirical expressions were developed over the years by the practitioners. For the treatment of domestic wastewaters in trickling filters, the following is commonly used:

$$C_2 = C_1 \exp \left[0.11^{(T-20)} Z \cdot A_v \cdot \frac{A_s}{Q} \right] \quad (19)$$

with C_1 and C_2 being the inlet and outlet substrate concentrations expressed in mg/l of BOD₅ (biochemical oxygen demand during 5 days), Z the height of the filter, A_v the surface area of the carrier per unit reactor volume (m^2/m^3), A_s the cross-sectional area of the reactor (m^2), Q the volumetric flow rate (m^3/s) and T the wastewater temperature ($^{\circ}\text{C}$).

For rotating biological contactors:

$$C_2 = C_1 - \frac{A_f}{Q} \frac{p \cdot C_2}{(k + C_2)} \quad (20)$$

where A_f is the surface area of the discs covered by biofilm and p and k are empirical parameters, which have to be estimated from

the operation of similar reactors at pilot-plants or full-size plants.

NOMENCLATURE

A_f	surface area of biofilm (m^2)
A_s	cross-sectional area of the filter (m^2)
A_v	specific area of support per volume of reactor (m^2/m^3)
b	reciprocal of the resistance to detachment (s^{-1})
C_1	substrate concentration at reactor inlet, as soluble BOD ₅ (mg/l^{-1})
C_2	substrate concentration at reactor outlet, as soluble BOD ₅ (mg/l^{-1})
D_f	effective diffusion coefficient or effective diffusivity (m^2/s^{-1})
k_m	external mass transfer coefficient (m/s^{-1})
k_{0f}	zero order biofilm reaction rate ($\text{kg}/\text{m}^3/\text{s}^{-1}$)
k_{1f}	first order biofilm reaction rate ($/\text{s}^{-1}$)
$k_{1/2ap}$	apparent half order constant ($\text{kg}^{1/2}/\text{m}^{1/2}/\text{s}^{-1}$)
L_f	thickness of the microbial layer (m)
m_f	mass of attached biofilm per surface area ($\text{kg}/\text{m}^2/\text{s}^{-1}$)
m_f	maximum mass of biofilm at steady state
Q	volumetric flow rate (m^3/s^{-1})
r_A	reaction rate per unit area of biofilm or surface reaction rate ($\text{kg}/\text{m}^2/\text{s}^{-1}$)
r_p	radius of the bare carrier particles (m)
S	bulk substrate concentration in the solution (kg/m^3)
S_1	substrate concentration at reactor inlet (kg/m^3)
S_2	substrate concentration at reactor outlet (kg/m^3)
t	time (s)
T	temperature ($^{\circ}\text{C}$)
V_R	reactor volume
Z	depth of the trickling filter (m)

Greek symbols

β	degree of substrate penetration in the biofilm
ε	reactor porosity

ϕ	Thiele modulus
η_i	biofilm internal efficiency
η	biofilm efficiency based on external substrate concentration
λ	external mass transfer rate/internal coupled diffusion-reaction rate

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Suspended growth processes

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1 INTRODUCTION

Suspended growth systems comprise aggregates of microorganisms generally growing as flocs in intimate contact with the wastewater they are treating. The aggregates or flocs are responsible for the removal of polluting material and comprise a wide range of microbial species. The most prevalent and important of these microorganisms are the bacteria, the protozoa and the metazoa. Fungi and viruses are also found, but probably contribute little to the treatment of the wastewater. Suspended growth treatment systems permit the exploitation of the full range of microbial metabolic capabilities. The full spectrum of redox environments from aerobic, through anoxic to anaerobic can be found within the floc itself, but they can also be created by appropriate process reactor design. These environments allow the growth of both the organoheterotrophs (which oxidize organic carbon and remove BOD) and the lithoautotrophs (which are responsible for ammonia oxidation). Indeed, in waste stabilization pond systems, phototrophic organisms, which utilize a range of electron acceptors, can be exploited to achieve good treatment with negligible energy input.

The food source for all the above organisms comes either in the soluble form dissolved in the wastewater, or as particulate material that is first solubilized by microbial action.

Treatment of wastewaters in suspended growth environments offers many process advantages, which has led to the proliferation

of this type of treatment system. Waste stabilization ponds, aerated lagoons and activated sludge are all examples of treatment options that rely on the actions of microorganisms growing in a suspension of the wastewater under treatment.

A suspended growth wastewater treatment processes is a biological reactor which has been engineered to encourage the growth of specific types of microorganisms that are able to undertake the reactions necessary to achieve purification of the influent wastewater. Their successful design requires the provision of:

1. a reactor (or series of reactors) of sufficient capacity to retain the wastewater long enough for the microorganisms to undertake the biological interconversions necessary to achieve the required effluent standard
2. facilities to ensure that the microorganisms are retained in the reactor long enough to grow and divide, thus maintaining a stable population
3. the correct redox environment to achieve the required biological reactions.

The most effective and innovative reactor design thus requires a full understanding of the biochemical basis for the growth of microorganisms using wastewater as the substrate. It is apparent from factors 1 to 3 above that the key design issues for a suspended growth process are the requirements of the treatment system in terms of its effluent quality.

2 EFFLUENT QUALITY REQUIREMENTS

The sole role of a wastewater treatment process is to protect the receiving watercourse from environmental degradation and ensure that the impact of the effluent from the plant is minimized. This is normally achieved by means of standards, which are set by the appropriate regulatory authority, to describe a minimum quality that the effluent must attain. Ideally these standards should be evaluated based on the characteristics of the receiving watercourse, the likely benefits that will accrue from maintaining a healthy watercourse and the ability of the community to finance treatment. Standards always contain a requirement for the treatment plant to achieve a certain effluent quality in terms of the biochemical oxygen demand (BOD) and nearly always have a standard for suspended solids. Depending on the quality of the receiving watercourse they

TABLE 21.1 The minimum treatment standards for sewage effluents that must be achieved by EC member states

<i>Parameter</i>	<i>Concentration (mg/l)</i>	<i>Minimum % of reduction^a</i>
(a) Requirements for discharges from urban waste water treatment plants		
BOD ₅ at 20°C without nitrification	25	70–90
COD	125	75
Total suspended solids	35 ^b	90
	(>10 000 pe)	
	60	70
	(2000–10 000 pe)	
(b) Discharges to sensitive areas which are subject to eutrophication		
Total phosphorus	2 mg P/l (10 000–100 000 PE) 1 mg P/l (>100 000 PE)	80
Total nitrogen ^c	15 mg N/l (10 000–100 000 PE) 10 mg N/l (>100 000 PE)	70–80

^a Reduction in relation to the load of the influent.

^b Optional requirement.

^c Organic and ammonical nitrogen and nitrite-nitrogen.

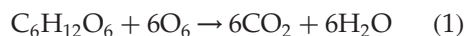
may then contain requirements for ammonia-nitrogen, total nitrogen and phosphorus. The minimum standards that apply throughout the European Community are summarized in Table 21.1, however, member states may impose stricter standards than this where it is thought appropriate.

3 ACHIEVING EFFLUENT QUALITY STANDARDS – MICROBIAL REACTIONS IN SUSPENDED GROWTH SYSTEMS

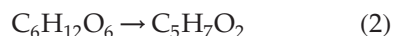
For each of the parameters listed in Table 21.1(a) and (b) there is one or more species of bacteria which can be exploited to metabolize the parameter using it either as a source of electrons or as a terminal electron acceptor. These reactions are summarized briefly below but are covered in more detail in other chapters in this volume.

3.1 BOD removal

The biochemical oxygen demand measures the amount of oxygen required to oxidize the organic carbon present in a wastewater according to the equation:



This oxidation reaction, referred to as respiration or catabolism, provides the energy necessary for bacterial growth and reproduction and there is a vast range of heterotrophic species which will carry it out in suspended growth processes. In addition to energy, the microorganisms need a source of carbon to build new cell material. This is also provided by the organic material measured in the BOD test according to Equation (2), where $\text{C}_5\text{H}_7\text{O}_2$ represents a new bacterial cell.



All biological wastewater treatment systems are designed to remove BOD, both in its particulate and soluble form. However, as much as 40% of the total BOD of a wastewater is particulate, consequently, it is generally most cost effective to remove this fraction by

sedimentation. That is the role of the primary sedimentation tank, septic tank or anaerobic pond. The fraction that remains after sedimentation is a mixture of colloidal and soluble BOD that can only be removed biologically. The soluble BOD is usually biodegraded very rapidly, generally in less than one hour. The colloidal fraction is entrapped in the sludge floc and is degraded more slowly. It is apparent from Equation (1) that breakdown of BOD to CO_2 and H_2O , removes BOD without the generation of new bacteria. Thus, if this reaction were encouraged, sewage treatment would take place without the generation of sludge. However, a lot of oxygen would be required and thus power consumption would be high. By contrast, Equation (2) shows that BOD can be removed by conversion to new bacteria without the need for oxygen. Thus, if this reaction were encouraged, treatment would take place without the need for aeration, however, the sludge generated would be very high. In practice there is a balance between the two reactions and treatment plants are operated with some sludge generated and some oxygen required.

It is worth bearing in mind Equations (1) and (2) when evaluating processes that claim a reduced sludge yield or a reduced power requirement. It is easy to reduce sludge yield by increasing the amount of oxygen provided and increasing the sludge age (extended aeration and aerobic digestion are examples of this). Similarly, it is easy to reduce the power requirement by a high rate process which generates more sludge. It is not possible to reduce both the sludge yield and the oxygen requirement!

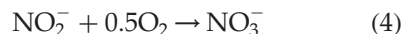
3.2 Ammonia removal

Ammonia is removed by the action of two groups of bacteria, collectively termed the nitrifying bacteria, which catalyse the reactions of nitrification. Nitrification is the process of ammonia oxidation in which ammonia is oxidized ultimately to nitrate in two reactions carried out by distinct groups of obligately aerobic bacteria. The first intermediate is nitrite and this reaction is catalysed by the genus

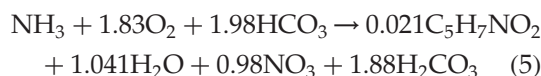
Nitrosomonas:



Nitrite is further oxidized to nitrate by *Nitrobacter*:



Nitrosomonas and *Nitrobacter* are both autotrophic genera which reduce carbon dioxide (in the form of bicarbonate or carbonate) as a source of cellular carbon. Assuming a gross cell composition for a typical nitrifying bacteria of $\text{C}_5\text{H}_7\text{NO}_2$, then the overall reaction for the oxidation of ammonia, coupled to the synthesis of new nitrifying bacteria, can be represented as:



The energy expenditure required to achieve the reduction of bicarbonate is relatively high, yet the nitrifying bacteria can achieve only a low yield of energy from the oxidation of their chosen substrates, ammonia and nitrite. Consequently, these organisms demonstrate very low growth yields and require a long retention time in the aeration basin to ensure they can divide and maintain a stable population. They are also very susceptible to temperature changes and below 20°C their reaction rate slows dramatically. This is illustrated in Fig. 21.1 which shows the sludge age required at a range of temperatures, in order to ensure that nitrification is achieved.

3.3 Nitrogen removal

Nitrate itself is able to act as a terminal electron acceptor. In the absence of a supply of dissolved oxygen, the utilization of oxygen for respiration (Equation (1)) cannot take place. However, certain chemo-organotrophs are capable of replacing O_2 with NO_3^- as an oxidizing agent and respiration can proceed with the reduction of nitrate to nitrite, nitric oxide, nitrous oxide or nitrogen. Equation (6) demonstrates the stoichiometric reaction for the reduction of nitrate using methanol as a source of electrons, with the production of

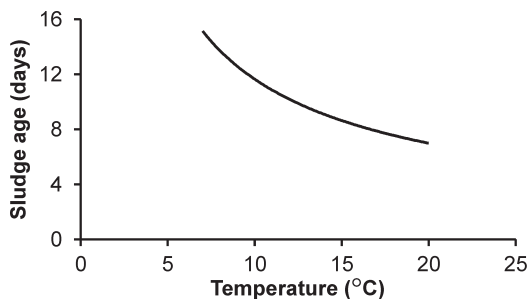
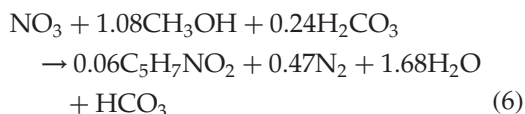


Fig. 21.1 The minimum design sludge age required at a range of sewage temperatures.

nitrogen gas and new cell material. This process is known as anaerobic or nitrate respiration and is carried out by a variety of bacteria such as *Alcaligenes*, *Achromobacter*, *Micrococcus* and *Pseudomonas*. Not all these genera are capable of complete oxidation to nitrogen and a variety of gaseous products can be produced.



As denitrification is an oxidation reaction, it needs an electron donor and an electron acceptor. Nitrate is the electron acceptor and organic carbon generally provides the source of electrons. This is usually from the BOD present in the sewage, or as a supplementary carbon source such as methanol or ethanol. It generally requires 8 mg COD to achieve the removal of 1 mg nitrate and it is advisable to aim for a nitrate concentration <12 mg/l in the final effluent to prevent sludge settlement problems.

3.4 Phosphate removal

Of the nutrients that are capable of supporting luxuriant growths of algae in a receiving water, phosphorus is rate limiting, and a concentration of 10 µg/l is required before algal growth will occur. It has been argued, therefore, that control over phosphorus-containing compounds in aquatic ecosystems presents a means of controlling the deleterious effects of

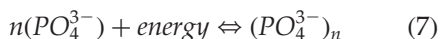
eutrophication. Consequently, if the small concentration of phosphorus that is present in sewage effluents can be removed, then algae will not be able to flourish, regardless of the nitrogen concentration. Phosphorus load control has been demonstrated as one of the most effective ways of dealing with cultural eutrophication. A typical phosphorus standard is 1 mg/l dissolved orthophosphate (as phosphorus). The major sources of phosphorus in domestic wastewaters are from human excreta (50–65%) and synthetic detergents (30–50%) and typical concentrations are in the range 10–30 mg/l as phosphorus.

4 ENHANCED BIOLOGICAL PHOSPHORUS REMOVED IN ACTIVATED SLUDGE SYSTEMS

The uptake and removal of phosphorus from a wastewater by activated sludge followed from the observation that if an activated sludge is allowed to become anaerobic, then the amount of phosphorus (as phosphate, PO_4^{3-}) in the supernatant increases. Upon resumption of aeration, however, there is a rapid uptake of phosphate by the sludge which is in excess of that released during anaerobiosis. This 'luxury uptake' of phosphate results in a phosphate depleted mixed liquor and a phosphate-rich sludge. The ability to release or store phosphorus in a sludge by manipulating the prevailing oxygen concentration, has been exploited in several processes, often referred to as enhanced nutrient removal processes.

The mechanism by which phosphorus is stored and released helps in understanding the operation of phosphorus-removal activated sludge plants. Removal and release of phosphorus within a sludge is thought to be the result of a single genus of bacteria known as *Acinetobacter calcoaceticus*. They are only able to take up and metabolize compounds contributing to the BOD under aerobic conditions. Under anaerobic conditions they cannot metabolize compounds but they can take up volatile fatty acids, in particular acetate, and store them intracellularly in the

form of a compound known as phb. Phosphate can also be stored intracellularly as polyphosphate granules known as volutin (Equation (7)):



Thus, if a wastewater is allowed to become anaerobic, *Acinetobacter calcoaceticus* will start to take up the volatile fatty acids that are generated under anaerobic conditions. The energy required to achieve this is obtained from the degradation of polyphosphate according to Equation (7), which releases soluble phosphate into the wastewater. The volatile fatty acids are then converted to phb and stored within the bacterial cell.

If the wastewater is now aerated the *Acinetobacter calcoaceticus* reverses this procedure and degrades the phb to volatile fatty acids. However, as oxygen is now present, aerobic metabolism is possible and the bacterium can break down the acetate to generate energy. This energy is used to take up phosphate and generate polyphosphate, but far more phosphate is taken up than is released in the anaerobic phase. The key to biological phosphate removal is thus to achieve anaerobic conditions and generate phosphate release. The more anaerobic the wastewater becomes, the more phosphate released and thus the more taken up in luxury phosphate uptake. In order to achieve the necessary anaerobiosis a lot of BOD is needed in the influent wastewater (around 12 mg BOD for every mg P removed). Often this is not available, particularly during the winter months (or during periods of heavy rainfall), and consequently additional chemical dosing capacity must be provided to ensure consent is met.

5 MANIPULATING REDOX ENVIRONMENTS

It is quite straightforward in a suspended growth wastewater treatment process to vary the electron source and the terminal electron acceptor and in such a way control the redox potential within the reactor. By doing so this allows a wide range of reactions to take place (Table 21.2). Up to three reactor types are exploited; these are defined, based on the source of electrons, as aerobic (or oxic), anoxic and anaerobic.

6 AEROBIC REACTORS

An aerobic reactor exploits oxygen as the terminal electron acceptor and is able to sustain a large number of important reactions that use a number of different electron donors. Principal among these are: oxidation of organic material, which reduces the BOD; oxidation of ammonia to nitrate, which reduces the ammonia concentration; and the luxury uptake of phosphate with the synthesis of polyphosphate, which reduces effluent phosphate concentrations.

A variety of options are available to achieve aerobic conditions in wastewaters. The strength of the incoming wastewater dictates the amount of oxygen that must be introduced into the reactor and guideline values are around 1.2 mg oxygen for every mg of BOD to be removed and 4.8 mg of oxygen for every mg ammonia to be removed. Thus, for a wastewater which has a BOD of 180 mg/l and ammonia of 25 mg/l and which must achieve an effluent quality of 20 mg/l BOD and 1 mg/l ammonia, it is necessary to introduce

TABLE 21.2 The reactions that can be achieved by manipulating the redox environment in a suspended growth reactor

Reaction	Electron donor	Electron acceptor	Redox potential
BOD removal	Organic material	Oxygen	> +200 mV
Nitrification	Ammonia	Oxygen	> +300 mV
Denitrification	Organic material	Nitrate	– 100 mV to + 150 mV
Phosphorus release	Polyphosphate	phb	< – 300 mV
Phosphorus uptake	Acetate	Phosphate	> – 150 mV

TABLE 21.3 Key design parameters for suspended growth processes

Reactor type	Biomass (mg/l)	Hydraulic retention time (days)	Volumetric loading (kg BOD/m ³)	Effluent quality ^a			
				BOD	NH ₄	P	SS
Waste stabilization pond	200	>20	0.002–0.008	30	4	5	^b
Aerated lagoon	800	2–4	0.04–0.08	30	10	5	30
Activated sludge	3500	0.3–0.5	0.3–0.6	10	<1	<1	10
Membrane bioreactor	>12 000	0.4–0.6	0.6	2	<1	<1	<2

^a Measured at a 95%ile.^b Pond effluents are high in suspended solids, but as this is largely algal biomass it is permitted up to a concentration of 150 mg/l.

307 mg O₂/l. Natural processes such as atmospheric diffusion and photosynthesis are able to introduce oxygen to a wastewater, although very slowly and thus the wastewater must be retained for a long time to achieve the necessary oxygen transfer. As a consequence, natural treatment options, such as waste stabilization ponds, have long hydraulic retention times, typically between 15 and 40 days, with large reactor volumes. By contrast as the rate of oxygen transfer is increased by mechanical means, the retention time of the wastewater can be reduced to as little as a few hours in the activated sludge process. However, as aeration intensity increases, so does the cost of the treatment process and high intensity aeration is both capital and operating cost intensive.

When wastewaters are aerated with a residual dissolved oxygen >1 mg/l, BOD removal occurs at a rate that is proportional to the amount of microorganisms held in the reactor. The more microorganisms that can be retained in the system the more rapid the rate of BOD removal will be and, consequently, the smaller the reactor that is required. However, there are two physical limits to the mass of microorganisms that can be retained, the first of these is the ability to transfer oxygen fast enough to match the oxygen uptake of the microorganisms and the second is the ability to remove the microorganisms by flocculation and sedimentation (section 10). This latter is generally the rate-limiting step in treatment. A new generation of suspended growth systems, known as membrane bioreactors, has eliminated this stage. Instead of a sedimentation step, the microorganisms are retained in the

system by a membrane, generally with a pore size of around 0.4 µm, which permits the treated effluent to exit while retaining the microbial biomass. Such reactor systems permit a large increase in the amount of solids that can be retained (Table 21.3).

7 ANOXIC REACTORS

The major anoxic reaction of importance in wastewater treatment is the reduction of nitrate to nitrogen gas. This is important for two reasons. First of all, if the nitrate concentration is high in the final sedimentation tank of the activated sludge process, denitrification can occur in the solids that have settled to the base of the tank. The nitrogen gas generated will buoy the settled sludge carrying it to the surface of the sedimentation tank and over the weir into the final effluent. This problem, termed a rising sludge, can be eliminated by ensuring the nitrate concentration in the final effluent does not rise above 12 mg/l.

A second reason is that some treatment plants have effluent discharge standards for total nitrogen (see Table 21.1b), in which case nitrate removal is mandatory.

Anoxic conditions are achieved by eliminating all the residual dissolved oxygen and ensuring an adequate supply of electrons. The final effluent from a nitrifying process has high concentrations of nitrate but lacks a supply of electrons. Thus, if it is directed to a reactor to undergo denitrification, the electron supply needs augmenting. This is achieved either by adding external carbon (usually as methanol or

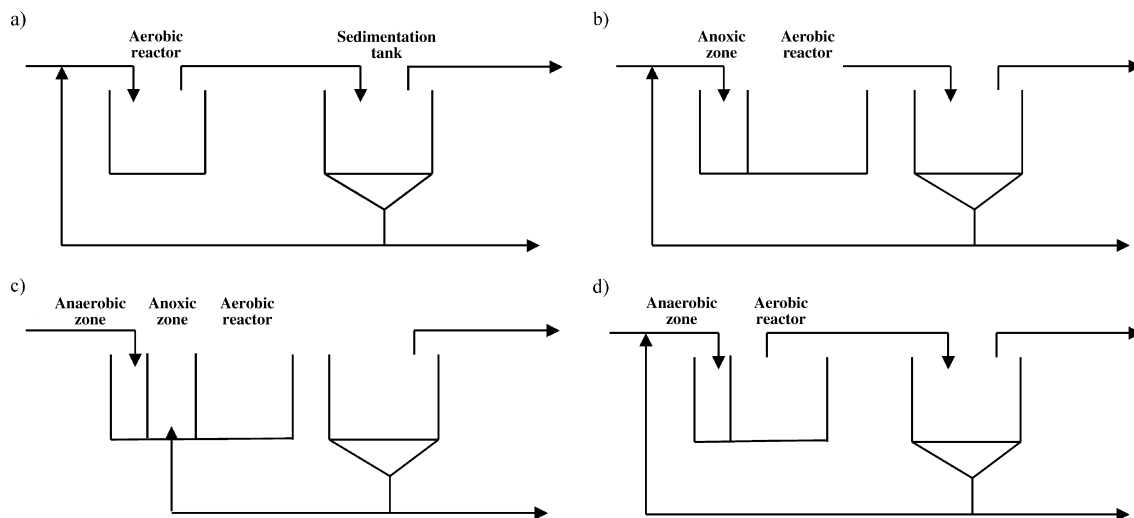


Fig. 21.2 Reactor configurations for manipulating redox conditions. (a) Carbonaceous removal only: an aerobic basin with sludge return direct to the basin; (b) Carbonaceous removal and nitrification: an aerobic basin with sludge return to an anoxic zone which also receives the incoming settled sewage; (c) Carbonaceous removal, nitrification, denitrification and phosphorus removal: an anaerobic zone to permit phosphate release, an anoxic zone which denitrifies the return sludge from the aeration basin and recycles it to the anaerobic zone, and an aerobic basin which achieves carbonaceous removal, nitrification and luxury phosphate uptake; (d) Carbonaceous removal and phosphorus removal. An anaerobic zone which receives return sludge from the aeration basin and permits phosphate release, followed by an aeration basin which achieves carbonaceous removal and luxury phosphate uptake.

a secondary industrial waste such as molasses) or directing a fraction of the settled sewage to the anoxic tanks. The most cost effective way of achieving denitrification is simply to remove the aeration from a small section at the head of the reactor and ensure that both the return sludge and the settled sewage are fed to this section. It requires a retention time of around 1 hour together with gentle mixing to ensure solids do not deposit in the basin (Fig. 21.2).

8 ANAEROBIC ZONES

An anaerobic reactor is necessary to promote the release of phosphate and uptake of acetate by the phosphate-accumulating bacteria. The more anaerobic the conditions then the more acetate is generated. This in turn leads to more phosphate release. Luxury phosphate uptake in the aerobic zone is always proportional to the amount of phosphate released and thus there is less phosphate in the final effluent. In order to ensure anaerobiosis within a reactor, it

is essential that there is no aeration device present and that the reactor is highly loaded with BOD such that any dissolved oxygen is removed rapidly. Anaerobic reactors are thus always sited at the head of the reactor treatment train (Fig. 21.2). If the aerobic stage of the treatment plant removes only BOD with no nitrification, then both the return sludge and the settled sewage can be fed to the anaerobic reactor. However, if the aerobic stage achieves nitrification, the nitrate in the return sludge will reduce the extent of anaerobiosis. Under such conditions the return sludge is fed to a separate anoxic reactor to undergo denitrification, and the nitrate free mixed liquor is recycled to the anaerobic reactor (Fig. 21.2).

9 MAINTAINING MICROBIAL GROWTH IN REACTORS

A typical wastewater has a BOD of around 200 mg/l of which 90% is removed during treatment to achieve an effluent quality of

20 mg/l. Using Equation (1), which is based on glucose as the carbon source, each mg of glucose requires 1.06 mg oxygen to oxidize it fully to CO_2 . Extending this to a typical wastewater, the microorganisms would have access to only $(200-20)/1.06 = 170$ mg glucose/l. When growing bacteria in laboratory cultures, it is common to use a glucose concentration of around 10 000 mg/l and thus it is apparent that a wastewater is a very dilute bacterial food source.

In order to determine the value of the three design factors described in the Introduction, it is necessary to know how quickly BOD is metabolized and how fast the bacteria can grow and divide. The rate of BOD removal can be described by the simple equation:

$$-\frac{dS}{dt} = qX \quad (8)$$

which says that the rate of BOD removal (dS/dt) is proportional to the number of microorganisms in the reactor (X in mg/l). The proportionality constant q is the specific BOD uptake rate (/day) and more usually referred to as the Food: Microorganism ratio or F/M .

In a similar way the rate at which bacteria grow can be described by Equation (9):

$$\frac{dX}{dt} = \mu X \quad (9)$$

where the rate of bacterial growth (dX/dt) is proportional to numbers of bacteria present in the reactor. The proportionality constant (μ) is known as the specific growth rate. There is a link between the specific growth rate and the amount of food source available and this is described by the Monod equation, when the food source is the rate limiting substrate (Equation (10)):

$$\mu = \mu_m \frac{S}{k_s + S} \quad (10)$$

where:

μ is the bacterial growth rate (/day)
 μ_m is the maximum specific growth rate (/day)

k_s is the saturation coefficient (mg/l), and
 S is the amount of rate limiting substrate in the effluent.

For a typical wastewater where the required effluent BOD is 20 mg/l, the value for μ_m is around 0.47/day and k_s is 100 mg/l. This gives a value for μ of 0.42/day (Equation (11)):

$$\mu = 2.5 \times \frac{20}{100 + 20} = 0.42/d \quad (11)$$

The reciprocal of the specific growth rate will have units of days and it is referred to as the microbial retention time. In other words, it is the amount of time the microorganism must remain in the reactor to ensure that they divide, based on the amount of food available to the microorganisms. Equation (11) can thus be expressed as:

$$\frac{1}{\mu} = \frac{V}{Q} \quad (12)$$

where V is the volume of the reactor (m^3) designed to hold the microorganisms and Q is the flow of sewage (m^3/day) into the reactor.

Substituting the calculated value of μ from Equation (11) into Equation (12) gives:

$$\frac{1}{0.42} = \frac{V}{Q}; \quad V = 2.4Q \quad (13)$$

Thus, in order to achieve an effluent BOD of 20 mg/l, it is necessary to construct a reactor with a retention time of 2.4 days in order to allow the microorganisms time to divide and grow. This is the principle behind a number of simple suspended growth treatment systems such as aerated lagoons and waste stabilization and they are characterized by large reactors with long residence times.

However, it is possible both to reduce the size of reactors for wastewater treatment, yet at the same time meet the requirements of Equation (10) in terms of bacterial growth. It was noted as long ago as 1912 that:

five weeks continuous aeration was required in order to completely nitrify an average sample of Manchester sewage. If, however, the resultant solid matter was allowed to deposit, and the purified sewage removed by

decantation and replaced by a further sample of crude sewage, complete nitrification of this second dose of sewage ensued within a reduced period of time. It was shown that accumulated deposit resulting from the complete oxidation by prolonged aeration of successive quantities of sewage, which were termed 'activated' sludge, had the property of enormously increasing the purification effected by the simple aeration of sewage.

This observation forms the basis of the activated sludge process and introduced the concept of recycling settled biomass from the sedimentation tank back to the aeration basin. In this way the hydraulic retention time is separated from the solids retention time and the requirements of Equation (10) for a long sludge age (measured in days) can be achieved in an aeration basin that has a short hydraulic retention time (measured in hours).

10 FLOCCULATION AND SEDIMENTATION – THE WEAK LINK IN THE PROCESS

The efficiency of the activated sludge process depends on the fact that, after repeated deposition of solid matter, the microorganisms that comprise these solids begin to change their community structure, and this is most marked for the protozoan and metazoan communities. The new communities which are so formed, acquire flocculating properties such that the sludge rapidly agglomerates and flocculates once aeration has ceased and quiescent conditions are applied. The ability of a floc to settle and thus provide a clear interface layer permits the discharge of a clarified effluent and the recycle of thickened solids back to the aeration basin. It is thus the basis on which the activated sludge process functions.

Floc settlement relies on the formation of large, compact flocs that are able to flocculate into larger aggregates, which have high settling velocities. This is the bottleneck in the treatment process and, although it is essential for ensuring efficient treatment, it is unreliable,

unpredictable and uncontrollable. A huge research effort has been expended into understanding the microbiological basis of activated sludge flocculation but negligible progress has been made in our ability to predict the onset of flocculation problems or alleviate them when they do occur.

Our current understanding of flocculation sees extracellular polymeric substances (EPS) as major components of the floc matrix with polymer bridging the mechanism by which flocs are formed. The major components of the EPS are polysaccharide and proteins excreted by bacteria and protozoa. Numerous authors have undertaken correlations that relate the EPS content of the sludge to the settleability as measured by SVI or SSVI, but it has been more difficult to correlate plant operating conditions (sludge age, loading rate, dissolved oxygen concentration etc.) with either EPS or settleability. It would appear that the amount of EPS generated is not as important as the properties of the EPS (namely hydrophobicity, surface charge and composition) in controlling sludge flocculation (Liao *et al.*, 2001).

However, the major settlement problem, which plagues almost all suspended growth processes, is that of filamentous foaming and bulking. This phenomenon is thought to be caused by the proliferation of filamentous bacteria and whenever bulking and foaming do occur these bacteria are almost always observed. Foaming results in the production of voluminous quantities of foam on top of the aeration basin, which can carry over to the secondary sedimentation tank and thus to the final effluent, where it will cause a solids consent failure. It is generally associated with the filament types *Microthrix parvicella* and *Nocardia* spp. Currently, there are no control strategies which will eliminate these filaments while still maintaining a high degree of wastewater treatment. However, it is possible to operate the treatment plant effectively by containing the foam within the aeration basin and preventing its carryover to the sedimentation tank.

Sludge bulking is a very different problem and it is thought that this is caused by the outgrowth of filamentous bacteria from

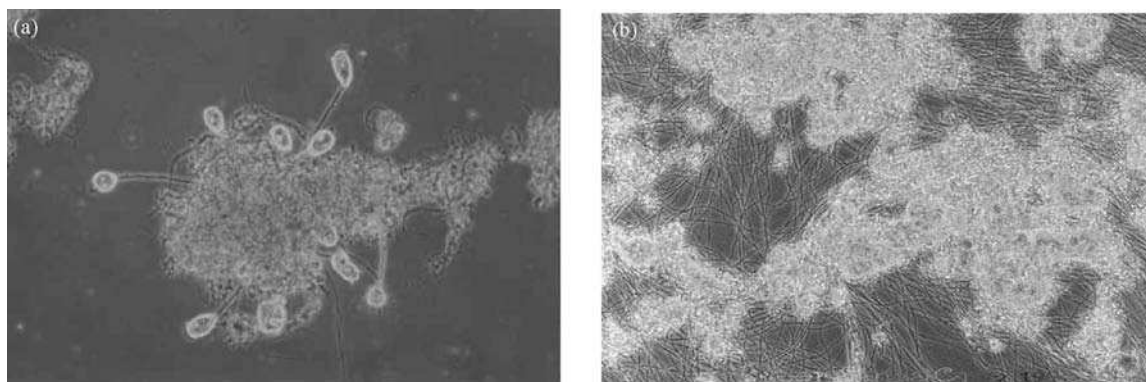


Fig. 21.3 (a) Well-settling sludge with a diverse protozoal community but showing clearly the rigid backbone of filaments supporting the floc biomass; (b) poor settling sludge with a large filament population of Type 021N.

the sludge floc. It is apparent from Fig. 21.3 that these outgrowths will prevent the flocs from approaching close to each other to form compact, well-settling structures. The effect is that the sludge occupies a much larger volume in the sedimentation tank, the sludge blanket is therefore much higher and the recycled sludge has a much lower suspended solids concentration. Where the final sedimentation tank is large with adequate capacity, this is a minor inconvenience which simply requires an increased recycle rate to return adequate solids back to the aeration basin and compensate for the thinner recycled sludge. However, if the sedimentation tank is working close to its capacity (and this is often the case at industrial treatment plants), the increased blanket height may cause solids to be lost over the outlet launder. In extreme cases the whole blanket may overtop the launder with a major loss of solids.

The provision of techniques to identify the causes of filamentous foaming and bulking and strategies to control their proliferation remains one of the challenges in activated sludge research. This problem, which has plagued the process since its inception, is one of the reasons why the membrane bioreactor (MBR) is such an attractive process option as it does not

require a sedimentation stage. It is likely that MBRs will become the suspended growth system of choice in the future, as membrane costs reduce and the operating experiences from those plants that have been constructed are disseminated.

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Protozoa as indicators of wastewater treatment efficiency

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1 PROTOZOA AS A COMPONENT OF THE ACTIVATED SLUDGE ECOSYSTEM

Today, most wastewater treatment processes make use of the natural self-purification capacity of aquatic ecosystems which results from the presence and action of microbial communities. Thus, biological sewage-treatment plants may be regarded as constructed ecosystems subjected to extreme conditions. As in every other biological system, the community living in the aeration basin of an activated-sludge plant has a precise structure and follows exact dynamics. In activated sludge, abiotic components are represented by the plant and by the sewage, while biotic components are represented by decomposers (bacteria, fungi) which utilize the dissolved organic matter in the wastewater, and by consumers (heterotrophic flagellates, ciliates, rhizopods, and small metazoans) which feed on dispersed bacteria and other organisms. Physical factors that act on biological sewage treatment systems are the climate, temperature and the turbulence to which mixed liquor is subject during the aeration phase, while chemical factors are represented by the composition of the wastewater and by oxygen dissolved by means of the aerators.

Regarding the biotic components, it is well known that activated sludge develops specific communities of protists which are sustained by large populations of bacteria. Ciliated protists are numerous in all types of aerobic biological-treatment systems. They are commonly found

at densities of about 10 000 cells/ml of activated sludge mixed liquor and constitute approximately 9% of the dry weight of suspended solids in mixed liquor (Madoni, 1994a). More than 200 species of protists (of which 33 flagellates, 25 rhizopods, 6 actinopods and 160 ciliates) were observed in the various types of aerobic treatment systems. Less than half of them, however, have been observed frequently (Curds and Cockburn, 1970a; Madoni and Ghetti, 1981; Al-Shahwani and Horan, 1991; Amann *et al.*, 1998). This number is evidently only a small part of the thousands of species of freshwater protists that could theoretically be observed in these environments.

The majority of ciliates present in biological wastewater treatment facilities feed upon dispersed populations of bacteria. The bacterivorous ciliates can be subdivided into three functional groups on the basis of behaviour:

- *Free-swimming*: swim in the liquor phase and remain evenly dispersed in the sedimentation tank
- *Crawling*: although they are free forms, inhabit the surface of sludge flocs or biofilm
- *Attached*: firmly fixed to the substrate by means of a stalk. They are strictly associated with the sludge flocs or biofilm and thus settle during sedimentation.

Some ciliates, however, are predators of other ciliates or are omnivorous, feeding upon a variety of organisms including small ciliates, flagellates and dispersed bacteria (Fig. 22.1).

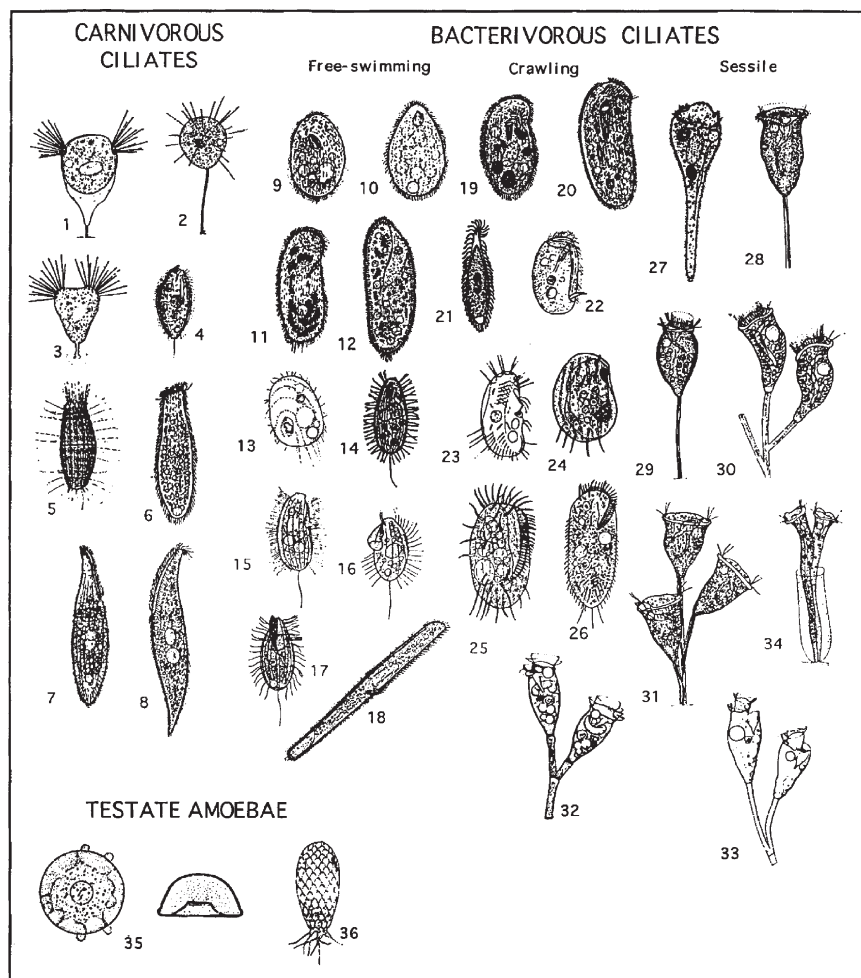


Fig. 22.1 Some common ciliates and testate amoebae of sewage treatment plants and their food habit. 1–8 Carnivorous or omnivorous ciliates: (1) *Acineta* spp.; (2) *Podophrya* spp.; (3) *Tokophrya* spp.; (4) *Plagiocampa rouxi*; (5) *Coleps hirtus*; (6) *Spathidium* spp.; (7) *Litonotus* spp.; (8) *Amphileptus* spp. 9–18 Free-swimming bacterivorous ciliates: (9) *Glaucoma* spp.; (10) *Tetrahymena* spp.; (11) *Colpidium* spp.; (12) *Paramecium* spp.; (13) *Cinetochilum margaritaceum*; (14) *Dextrotricha* spp.; (15) *Uronema* spp.; (16) *Cyclidium* spp.; (17) *Pseudocohnilembus pusillum*; (18) *Spirostomum teres*. 19–26 Crawling bacterivorous ciliates: (19) *Chilodonella* spp.; (20) *Trithymostoma* spp.; (21) *Acineria uncinata*; (22) *Trochilia minuta*; (23) *Drepanomonas revolute*; (24) *Aspidisca* spp.; (25) *Euplotes* spp.; (26) *Stylonychia* spp. 27–34 Sessile bacterivorous ciliates: (27) *Stentor* spp.; (28) *Vorticella convallaria*; (29) *V. microstoma*; (30) *Carchesium* spp.; (31) *Zoothamnium* spp.; (32) *Epistylis* spp.; (33) *Opercularia* spp.; (34) *Vaginicola* spp. 35–36 Testate amoebae: (35) *Arcella* spp.; (36) *Euglypha* spp.

All bacterivorous ciliates rely upon ciliary currents to force suspended bacteria to the oral region. The subclass Peritrichia is the most important group of ciliates in sewage treatment plants. All peritrichs found in aerobic processes are attached forms. It should be remembered that both activated sludge and attached growth processes rely on the presence of

surfaces (sludge or biofilm) upon which the microorganisms can grow. Thus organisms with the ability to attach to, or remain closely associated with, sludge or biofilm have a distinct advantage over organisms that swim freely in the liquid phase and are subject to washout in the effluent. So, while free-swimming and attached ciliates are in competition

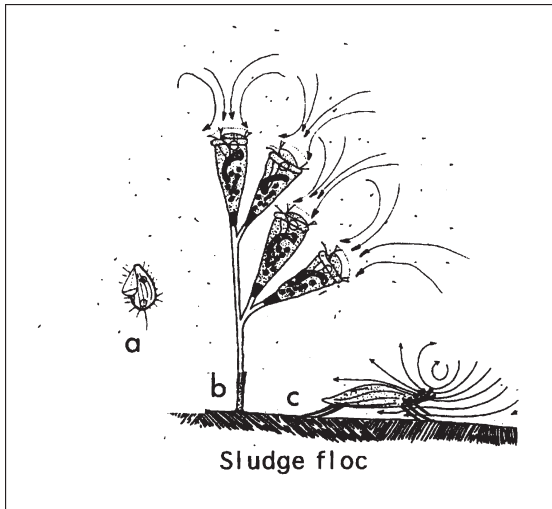


Fig. 22.2 Feeding mechanisms of filter-feeding ciliates in the activated sludge process. Swimming (a) and attached (b) forms filter out dispersed bacteria in the mixed liquor, while crawling forms (c) scrape bacteria from the surface of the flocs.

for bacteria dispersed in the liquid phase, crawling forms that are in close proximity to surface growths feed upon particles that only lightly adhere to the sludge and that are dislodged very easily by the feeding currents (Fig. 22.2).

However, frequent reference to 'scavenger' hypotrichs such as *Aspidisca* and *Euplotes* and ciliophorids such as *Chilodonella* and *Trochilia* implies that these crawling ciliates can scrape bacteria from surfaces because their mouth is located in the ventral part of the cell.

2 THE ROLE OF PROTISTS IN THE ACTIVATED SLUDGE PROCESS

Large numbers of ciliates are present in all types of aerobic treatment systems and play an important role in the purification process, as well as the overall regulation of the entire community. Ciliated protists improve the quality of the effluent because of their involvement in the regulation of the bacterial biomass by removal, through predation, of the major part of the bacteria dispersed in the mixed liquor (Curds *et al.*, 1968). It is generally assumed that

their primary role in the wastewater treatment is the clarification of the effluent. In the absence of ciliated protists, in fact, the effluent from the system has an elevated BOD and is highly turbid due to the presence of many dispersed bacteria.

Ciliates, moreover, feed on pathogenic and faecal bacteria. In the effluent from systems lacking ciliates, the presence of *Escherichia coli* is, on average, equal to 50% of that observed in the sewage entering the aeration tank. This is reduced to 5% when ciliates are present (Curds and Fey, 1969).

3 PROTOZOA DYNAMICS IN ACTIVATED SLUDGE

In the aeration tank of biological processes a true trophic web is established. A simplified diagram of this is illustrated in Fig. 22.3. The biological system of these plants consists of populations in continuous competition with each other for food. The growth of decomposers, prevalently heterotrophic bacteria, depends on the quality and quantity of dissolved organic matter (DOM) in the mixed liquor. For predators, on the other hand, growth depends on the available prey. Dispersed bacteria are thus food for heterotrophic flagellates and bacterivorous ciliates which, in turn, become the prey of carnivorous organisms. The relationships of competition and predation create oscillations and successions of populations until dynamic stability is reached. This is strictly dependent of plant management choices based on design characteristics aimed at guaranteeing optimum efficiency.

Although the ciliates are widely distributed, and many species are able to tolerate precarious environmental conditions, the particular conditions found in sewage treatment processes limit their presence to a restricted number of species (Madoni and Ghetti, 1981). Because of this protists have come to be considered as potential indicators of wastewater treatment plant performance. Curds and Cockburn (1970b) were probably the first to relate the sludge loading to the various forms of ciliated protists. According to these authors,

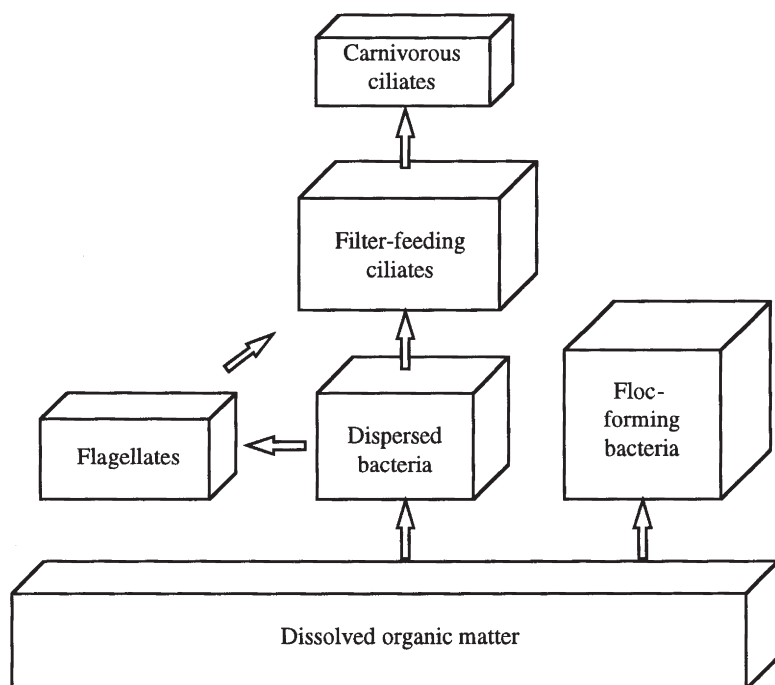


Fig. 22.3 The trophic web in the activated sludge of sewage treatment plants.

crawling ciliates (hypotrichs) decrease with increasing loading (no hypotrichs would be observed in sludge loaded above 0.6 kg BOD/kgMLSS/day), while sessile ciliates (peritrichs) are able to grow throughout a large range of sludge loadings. Nevertheless, at low loadings (0.1–0.3 kg BOD/kgMLSS/day) a wider range of ciliates, in terms of numbers of species with a more uniform distribution between the three functional groups, would be observed. Klimowicz (1970) also found that some ciliated protist species were most abundant on occasions at high sludge loadings (*Paramecium caudatum*, *Vorticella microstoma*, *Opercularia coarctata*, *O. microdiscus*), while some other species occurred in great numbers at medium loadings (*Acineria uncinata*, *Amphileptus claparedei*, *Litonotus fasciola*) and some other species were most numerous at low loadings (*Vorticella picta*, *Zoothamnium*, *Coleps hirtus*). Such species as *Aspidisca cicada* and *Vorticella convallaria* were equally abundant with various loadings of sludge. Studies on both colonization and population succession in activated sludge emphasized the role of protists as plant performance indicators and demonstrated the effect of

environmental conditions in the aeration tank on determining the established ciliate community (Curds, 1971; Madoni, 1982).

Another important result was the identification of three phases in the time span from the beginning to stabilization of the system (Madoni and Antonietti, 1984). The plant starting phase is characterized by the presence of species typical of raw sewage (Fig. 22.4). These 'pioneer' species are represented chiefly by free-swimming bacteriophagous ciliates and flagellates and are thus not linked to the presence of sludge so they cannot be considered typical components of these environments. With the formation of activated sludge, they compete with species better adapted to an aeration tank environment and rapidly decline in numbers. The second phase is one of transition, and is characterized by the strong growth of ciliates typical of the aeration tank habitat. In this phase, the community has a wealth of species, but rapid substitution of the species takes place with the progressive formation of activated sludge. The free-swimming forms, above all the hymenostomes, are involved only during the first days of

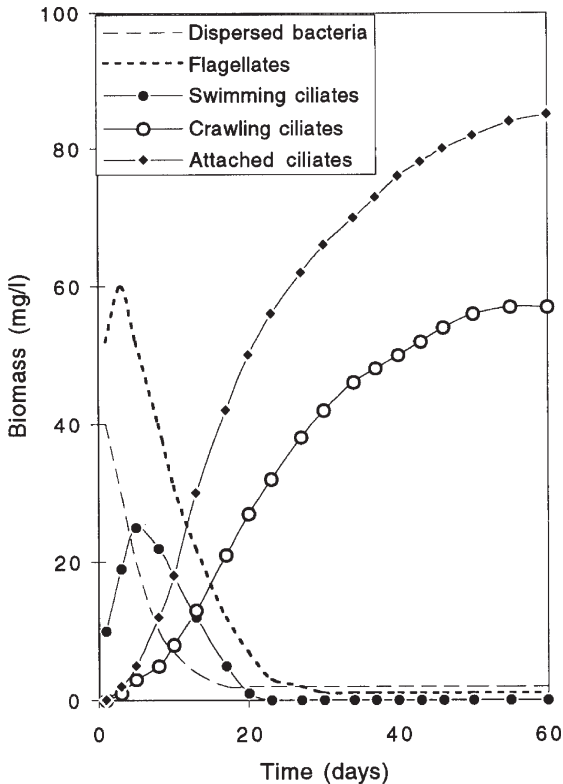


Fig. 22.4 Microorganism succession during colonization of activated sludge.

colonization and are gradually substituted by the attached and crawling forms. The steady-state phase is characterized by a ciliate community whose structure reflects the stable condition of the aeration tank environment with a balance between the organic loading and the sludge that is produced, removed and recycled. Each of the three phases is characterized by a typical species structure. Free-swimming bacterivorous ciliates such as *Colpidium*, *Cyclidium* and *Paramecium* are linked to the first phases of colonization of the plant, while attached peritrichs such as *Vorticella*, *Epistylis*, *Opercularia* and crawling ciliates such as *Aspidisca* are typical of the third phase.

A fully functioning plant need not host species characteristic of one of the colonization phases unless dysfunctions cause regression in the environmental conditions, such as the amount of sludge, the degree of aeration, sewage retention time, and organic loading at input.

The species structure of the microfauna is thus a diagnostic instrument which serves to integrate the parameters on which the evaluation of plant performance is based. By following the general criterion of the biological indicators, the presence of particular species, as well as the overall composition of the microfauna, can be taken as indicators of the performance of an activated sludge plant. With regard to ciliated protists, the groups of species take on particular importance both in systematic and functional terms (free-swimming, crawling, attached).

4 MICROFAUNA AS INDICATOR OF ACTIVATED-SLUDGE SYSTEMS

The numerous studies undertaken have been able to ascertain that the number of ciliated protists living in a normally functioning plant is about 10^6 individuals per litre. When the number falls below 10^4 per litre it indicates insufficient purification. In this case, there is a proliferation of dispersed bacteria which render the effluent turbid and consequently cause a greatly increased BOD in the output water. A high number of ciliates ($>10^7$ per litre) instead, almost always indicates good purification and optimum plant performance. The microfauna of a normally functioning system is almost always highly diversified. That is, it is composed of different groups of organisms, and each group is made up of several species. No group or species is every numerically dominant over the other components, even though the ratios between various groups or species can differ. However, a microfauna that is dominated by one species or group is almost always an index of trophic imbalances due to the existence of limiting factors impeding development of most of the other species and favouring the growth of forms more tolerant to these factors. The chief limiting conditions are generally the presence of a shock load of toxic discharge, under- or overloading, excessive sludge wastage or lack of aeration.

The structure of the microfauna is indeed a valid indicator of plant purification perform-

ance and an efficient activated sludge plant has the following characteristics:

- High numbers of microfauna cells ($\geq 10^6$ organisms/l)
- Microfauna composed chiefly of crawling and attached ciliates, with almost no flagellates or swimming ciliates
- The species and ciliates groups are highly diversified and none dominates numerically by a factor greater than 10.

When this is not the case, the identification of the dominant group of the microfauna allows diagnosis of the particular state of functionality of the plant.

Just as observation of the microfauna, identification of flagellates, free-swimming, crawling and attached ciliates and the analysis of their abundance ratios is a useful tool in the diagnosis of plant performance, so more complete information can be obtained by a knowledge of the different species of the protistan community. Thus, if it is true that a microfauna rich in attached and crawling ciliates indicates improved performance with respect to that found when free-swimming ciliates dominate, it is also true that different species of crawling and attached ciliates can be associated with various conditions. Thus, each species is able to add more detailed information than that obtained through simple identification of functional groups. For example, among attached ciliates, *Vorticella*

convallaria and *V. microstoma* characterize the first phase of colonization but the latter are then substituted by *Vorticella convallaria* which may reach high numbers during the growth phase of the sludge. In the case of a drastic reduction in the oxygen content in the mixed liquor, the alternation of these two species is observed, due to their different degree of tolerance to the lack of oxygen (Fig. 22.5). Large quantities of *Vorticella microstoma* thus indicate a poorly aerated sludge.

5 THE KEYGROUPS

Quantifying the indicator value of the microfauna is a difficult task because there are groups that are more or less tolerant of a wide range of environmental factors, while the plant performance results from the simultaneous action of many of these external and operational conditions. Nevertheless, investigating the relationship between the various groups of the microfauna and the main operational conditions of the plant in a wide survey, it was possible to select and group microfauna organisms into positive and negative keygroups (Madoni, 1994b). The positive keygroups – positively correlated with high plant performance – are crawling and attached ciliates and testate amoebae. The negative keygroups – negatively correlated with high plant performance – are small flagellates, swimming bacterivorous ciliates, and the sessile ciliates *Vorticella microstoma* and *Opercularia* spp. Density and species richness of the microfauna, moreover, appear to be highly correlated to plant performance (Table 22.1).

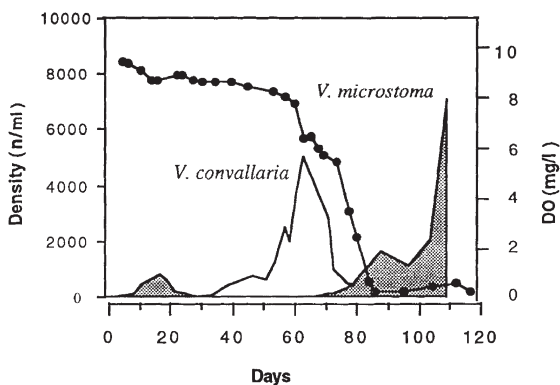


Fig. 22.5 Dynamics of two species of *Vorticella* related to dissolved oxygen (DO) in the aeration tank.

5.1 Small flagellates

Small heterotrophic flagellates continuously enter the plant in the sewage influent where they are very numerous. Flagellate species such as *Bodo*, *Polytoma* and *Tetramitus*, normally dominate the microfauna during the starting phase of the plant when floc-forming bacteria are scarcely present. They feed on dispersed bacteria and, in time, are substituted by bacterivorous ciliates. In a normally-functioning

TABLE 22.1 Positive and negative correlation between protozoans and plant operational conditions (* $P < 0.01$, ** $P < 0.001$)

	DO	Nitrifying ability	BOD removed	Effluent colour
Small flagellates	— *	— **	— **	+ **
Swimming ciliates	— **	— **	— **	+ **
Crawling ciliates	+ **	+ **	+ **	— **
Sessile ciliates	+	—	+	— **
<i>V. microstoma</i>	— **	— **	— **	+ **
<i>Opercularia</i> spp	— **	— **	— **	+ **
Testate amoebae	+ **	+ **	+ **	— **
Microfauna density	+ **	+	+ **	— **
Number of species	+ **	+ **	+	— **

activated sludge, in fact, these protists are outstripped by the bacterivorous ciliates and, in addition, they are strongly subjected to predatory activity of other protists; so their presence in the activated sludge is limited to few individuals (>10 individuals counted along the diagonal in a Fuchs-Rosenthal chamber). By contrast, the massive presence of these protists in a mature activated sludge is associated with a poor performance in the biological purification, due to the following causes: poorly aerated sludge; overloading; fermenting substances involved (Drakides, 1978; Madoni, 1986). The dominance of flagellates becomes apparent when they reach a density of more than 100 individuals along the diagonal in a Fuchs-Rosenthal chamber. In this case the effluent leaving the system has an elevated BOD and is highly turbid due to the presence of these microorganisms. Flagellates become the only protistan form present in highly-loaded sludge (>0.9 kg BOD/kgMLSS/day) (Curds and Cockburn, 1970b).

5.2 Swimming bacterivorous ciliates

Free-swimming bacterivorous ciliates are more abundant in the early phases of a developing plant when sludge flocs are still scarce and consequently attached ciliates are absent. Nevertheless, they are soon substituted by sessile ciliate species owing to competition for bacteria dispersed in the mixed liquor. Sessile forms in fact are filter-feeders and are more

efficient than free-swimming ciliates in forcing suspended bacteria into the cell by means of ciliary currents. Small free-swimming ciliates (such as *Uronema*, *Dexiostoma* and *Tetrahymena*) sometimes dominate the microfauna of plants operating at sludge age which is too low or at both high sludge loadings and a lack of oxygen (Martín-Cereceda *et al.*, 1996). These bacterivorous ciliates require high concentrations of dispersed bacteria but survive better than other protists when there is toxicity in the influent or a lack of oxygen. Swimming ciliates dominate the aeration basin of plants characterized by high sludge loadings (0.6–0.9 BOD/kgMLSS/day).

5.3 Crawling and attached ciliates

These two functional groups normally codominate the protistan community in activated sludge plants. This is due to their different food habits preventing their competition. Nevertheless, the ratio between the two groups tends to change with sludge loading. Crawling ciliates reduce their presence as sludge loading increases, so that above 0.6 kg BOD/kgMLSS/day, most species of this group disappear (Curds and Cockburn, 1970b; Klimowicz, 1970).

Sessile ciliates are normally codominant in the activated sludge. Nevertheless, a massive increase in their number (>80% of the whole microfauna) occurs on occasions in transient situations that reduce the plant performance (Drakides, 1978; Madoni, 1981). Such transient conditions are a rapid increase in the sludge load due to a loss of sludge and a discontinuous input of organic load from the influent. Sessile ciliates are able to grow throughout a large range of sludge loadings; nevertheless, at values ranging from 0.3 to 0.6 kg BOD/kgMLSS/day, these protists dominate and, for sludge loadings of 0.6–0.9, sessile ciliates some forms, such as *Vorticella microstoma* and *Opercularia* spp., can survive and grow in activated sludge subjected to severe conditions (lack of oxygen, presence of toxins). When these species are present in high numbers, they must be considered as separate keygroups.

5.3.1 *Vorticella microstoma*

The peritrich ciliate *V. microstoma* is quoted in the saprobic system as a polysaprobic species (Foissner, 1988). This ciliate is frequently present in the plant during the first phase of colonization but is substituted by other sessile ciliates (*V. convallaria*) during stable conditions when there is a drastic and prolonged reduction in the dissolved oxygen concentration in the mixed liquor. An alternation of the two species can be observed owing to their different degrees of tolerance to the lack of oxygen (Madoni and Antonietti, 1984). *V. microstoma* thus indicates a lack of dissolved oxygen in the aeration tank. Massive growth of this sessile ciliate was also observed at times of high wastewater flow to the sewage plant and low values of mass loading (Esteban *et al.*, 1990).

5.3.2 *Opercularia* spp.

Low numbers of *Opercularia* often occur in activated sludge, where three species are commonly observed: *O. articulata*, *O. coarctata*, and *O. microdiscus*. These ciliates are quite useful as bioindicators, because their numbers increase when the activated sludge is of poor quality. *Opercularia* spp. are associated with high final effluent BOD concentrations, and are among the most abundant forms at high loadings (Curds and Cockburn, 1970b; Klimowicz, 1970; Esteban *et al.*, 1991; Salvadó *et al.*, 1995). These ciliates moreover can survive in stressed environments better than other protists; in fact, large numbers of *Opercularia* were found in sludge of plants receiving industrial waste containing toxic substances (Antonietti *et al.*, 1982; Becares, 1991; Becares *et al.*, 1994). Moreover, *Opercularia* spp. were associated with high effluent BOD and ammoniacal N concentrations (Poole, 1984; Madoni *et al.*, 1993). *O. coarctata* may be the only component of the microfauna in sludges of plants treating industrial waste containing metal salts. *O. microdiscus* is able to survive the severe lack of oxygen found in sedimentation tanks with a low return-sludge ratio. *Opercularia* spp. are often associated with *V. microstoma*.

5.3.3 *Testate amoebae*

Two genera of testate amoebae, *Arcella* and *Euglypha*, are commonly present in activated sludge. These protists are found normally in the aeration basins of N-removal plants (Poole, 1984). Testate amoebae are more abundant or dominant in sludges characterized by low loading, a long retention time and high DO in aeration tanks that enable complete nitrification (Madoni *et al.*, 1993). Under these conditions, the quality of the effluent is excellent and a high biological performance of the plant is reached. Testate amoebae only colonize plants with a long sludge age since these protists have low growth rates. They are often seasonal, being more common in summer when temperature and growth rates increase. They may be more abundant in activated sludge plants at breweries with a low sludge load and a good quality effluent. Nevertheless, when the sludge load reaches high values (>1 kg BOD/kgMLSS/day) and the COD of the effluent is elevated, these protists are replaced by the peritrich ciliate *Opercularia* and by free-swimming ciliates (Sasahara and Ogawa, 1983).

5.3.4 *Density and diversity*

The number of ciliates living in a normally functioning plant is about 10^3 individuals/ml, and when the number falls below 10^2 /ml it indicates insufficient purification (Drakides, 1980; De Marco *et al.*, 1991). In this case, there is a proliferation of dispersed bacteria which make the effluent turbid and consequently causes a greatly increased BOD in the output water. On the other hand, a high number of ciliates ($\geq 10^4$ /ml) indicates good purification and optimum biological performance of the plant. The microfauna of a normally functioning plant is almost always highly diversified, namely composed by a high number of species (>10). In this case no species is ever numerically dominant over the other organisms, even if the ratios between various species differ. A microfauna that is dominated by one species is an index of trophic imbalances due to the existence of limiting factors impeding the development of most of the other species and favouring the growth of forms more tolerant

TABLE 22.2 Two-way table to determine the Sludge Biotic Index (modified from Madoni, 1994b). Horizontal entrance in the table on the basis of both keygroup and density. Vertical entrance in the table. Total number of taxa of the microfauna and number of small flagellates F counted along the Fuchs-Rosenthal chamber diagonal

Dominant keygroup	Density (ind/l)	> 10		8–10		5–7		< 5	
		F > 10	10 < F < 100	F > 10	10 < F < 100	F > 10	10 < F < 100	F > 10	10 < F < 100
Crawling + sessile ciliates ^a and/or testate amoebae	≥10 ⁶	10	8	9	7	8	6	7	5
	<10 ⁶	9	7	8	6	7	5	6	4
Sessile ciliates ^a >80%	≥10 ⁶	9	7	8	6	7	5	6	4
	<10 ⁶	8	6	7	5	6	4	5	3
<i>Opercularia</i> spp.	≥10 ⁶	7	5	6	4	5	3	4	2
	<10 ⁶	6	4	5	3	4	2	3	1
<i>Vorticella microstoma</i>	≥10 ⁶	6	4	5	3	4	2	3	1
	<10 ⁶	5	3	4	2	3	1	2	0
Swimming ciliates	≥10 ⁶	5	3	4	2	3	1	2	0
	<10 ⁶	4	2	3	1	2	0	1	0
Small flagellates (> 100) ^b	≥10 ⁶		4		3		2		1
	<10 ⁶		3		2		1		0

Conversion of SBI values into four quality classes

SBI Value	Class	Judgement
8–10	I	Very well colonized and stable sludge; excellent biological activity; very good performance.
6–7	II	Well colonized and stable sludge; biological activity on decrease; good performance.
4–5	III	Insufficient biological purification in the aeration tank; mediocre performance.
0–3	IV	Poor biological purification in the aeration tank; low performance.

^a *Opercularia* and *V. microstoma* not dominant.

^b along the Fuchs-Rosenthal chamber diagonal.

to these factors. The number and diversity of ciliate communities change according to the quality of the settled sewage and operating conditions of the plant (Esteban *et al.*, 1991; Esteban and Tellez, 1992). The most common limiting conditions are generally the presence of a shock load of toxic material, the lack of aeration, and the excess sludge wastage.

6 THE SLUDGE BIOTIC INDEX (SBI)

The performance of the plant has to be constantly monitored and is subjected to strict regulation. Nevertheless, malfunctions resulting in decreased purification efficacy are frequent. The SBI, an objective index based on the protistan community, has been devised to monitor activated-sludge plant performance (Madoni, 1994b). This method is based on two principles. First, the dominance of protistan keygroups changes in relation to environmental and operational conditions of the plant. Second, cell density and number of taxa diminish as the efficiency of the plant drops. The SBI enables the operator to define the biological quality of the sludge by means of conventional numerical values (from 0 to 10) that are grouped into four quality classes (Table 22.2). The identification of the various species of protista is important, in order to obtain an accurate SBI value; some keys written specifically about the protista found in sewage treatment processes and polluted waters are available (Bick, 1972; Foissner *et al.*, 1991–95).

Since the SBI was set up specifically for the evaluation of the biological reactor performance, this index is unable to reveal any dysfunction in the final sedimentation tank such as sludge bulking or rising sludge.

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The microbiology of phosphorus removal in activated sludge

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1 INTRODUCTION

Eutrophication is a natural process which usually occurs in lakes and other quiescent bodies of water through the introduction of the plant nutrients, phosphorus (P) and nitrogen (N), to the impoundment. Without human intervention, the process takes place over hundreds of years, but is greatly accelerated by various human activities in sensitive areas.

Eutrophication results in many undesirable effects, the primary effect being profuse algal blooms and excessive growth of nuisance aquatic plants. These include several species of blue-green (*Cyanobacteria*) and green (*Chlorophyta*) algae, the diatoms and flagellates, water grasses, rooted broad-leaved plants and floating water plants (hyacinth) (Rudd, 1979). Secondary effects, which are a direct result of weed and algal growth, include rapid oxygen uptake from the water causing the lower water to become anaerobic; production of methane and sulphides, which may result in fish kills (especially in winter) or impart a strong rotten egg smell to the water; and the negative aesthetic appeal of the dam due to overturning of the water and the appearance that the water is black; and the water can no longer be used for potable purposes as certain species of the algae are toxic (Rudd, 1979). Costs of purification of such eutrophied waters escalate dramatically.

Many limnological studies have been conducted concerning eutrophication, primarily

with its causes and effects and the results have conclusively indicated that eutrophication is promoted if phosphate (P) and nitrogen (N) are released into a reservoir or catchment area (Walmsley and Thornton, 1984; Chutter, 1990; Dillon and Molot, 1996). These studies have also shown that eutrophication can be effectively controlled if the P load to receiving waters is controlled. Gross eutrophication becomes marked when the inorganic soluble N and P concentrations in waters are in excess of 0.3 mgN/l and 0.015 mgP/l, respectively (Lilley *et al.*, 1997) (Table 23.1). It is virtually impossible to control eutrophication by limiting nitrogen due to the ability of the causative agents (algae) to fix and assimilate atmospheric N. Assimilated N is then made available to other aquatic life forms when these cells die and assimilated N is released. The increased awareness that P is the limiting nutrient has led to the introduction of more stringent legislation governing the discharge of P to receiving waters.

To prevent eutrophication, phosphate removal from effluents is necessary, whether it is by chemical and/or biological means (Toerien *et al.*, 1990). The activated sludge process comprises a complex and enriched culture of a mixture of generalists and specialist organisms. The lack of fundamental understanding of the enhanced biological phosphate removal (EBPR) processes, and the requirement to improve processes designed to remove nutrients from the influent have compelled researchers to examine and attempt to optimize the biological

TABLE 23.1 Relationship of lake trophic status (productivity) to average concentrations of N and P (Wetzel, 1983)

Lake productivity	Inorganic N ($\mu\text{g/l}$)	Organic N ($\mu\text{g/l}$)	Total P ($\mu\text{g/l}$)
Ultra-oligotrophic	<200	<200	<5
Oligo-mesotrophic	200–400	200–400	5–10
Meso-eutrophic	300–650	400–700	10–30
Eutrophic	500–1500	700–1200	30–100
Hypereutrophic	>1500	>1200	>100

components of the mixed liquor of activated sludge plants. Many different configurations of activated sludge processes exist. We will discuss only one of these, representing the most common denominators.

1.2 The activated sludge process

Activated sludge systems are based on suspended-growth processes and have become an integral part of municipal wastewater treatment. The process relies on the dense growth of microorganisms in a reactor where air is continuously supplied to allow for carbonaceous oxidation. The term 'activated sludge' refers to an aerobic slurry of microorganisms which can be removed from the process through sedimentation and returned, in quantifiable amounts, to the wastewater stream (Grady and Lim, 1980a, b). All activated sludge systems operate with the following characteristics in common: utilization of a flocculent slurry of microorganisms to remove organic matter from the surrounding wastewater; prior to effluent discharge from the plant, microorganisms are removed through sedimentation thereby reducing outgoing solids loads; settled microorganisms are recycled to the biological reactor via a clarifier underflow; and dependency of plant performance on the mean cell residence time (MCRT or sludge age) in the system (Grady and Lim, 1980a, b). Microbial metabolism of the organic matter present results in the production of oxidized end-products such as carbon dioxide, nitrates, sulphates and phosphates, as well as the biosynthesis of new microbial biomass (Gray, 1989; Horan, 1990; Bitton, 1994; Muyima *et al.*, 1997).

The basic ecological unit of activated sludge is the floc. Microbial floc formation is essential to the success of activated sludge processes as it allows for rapid and efficient separation of sludge from treated wastewater in the sedimentation tank. Although the exact mechanism of floc formation is not well understood, it seems to be almost entirely bacterially mediated (Muyima *et al.*, 1997). As extracellular polysaccharide production gradually continues, other microorganisms and colloidal material become entrapped in the matrix and the floc diameter increases. Surface charges on the microbial cells and bridge formation by polyvalent cations also contribute to flocculation (Gray, 1989). This very rigid floc structure has impeded the quantitative analysis of the activated sludge microbial community structure, as complete dispersion of the floc is extremely difficult. Clumping of cells in the floc leads to an underestimation of the number of active cells present in the mixed liquor when using viable plate count techniques. Problems attributable to poor floc formation can be one of two causes:

1. non-filamentous bulking which describes flocs that do not settle well due to excessive production of extracellular polysaccharides and the formation of loose flocs, i.e. excess of zoogloeal organisms;
2. filamentous bulking, caused by excessive growth of filamentous bacteria (Bitton, 1994). These undesirable characteristics will have negative economic consequences to the plant in question.

1.2.1 Anaerobic zone

The principal function of the anaerobic zone is to establish a facultatively anaerobic microbial community as indicated by the fermentation pattern. During anaerobiosis this bacterial community produces compounds such as ethanol, acetate and succinate, which serves as carbon sources for phosphate-accumulating bacteria (Cloete and Muyima, 1997).

In the anaerobic zone dissolved oxygen and oxidized nitrogen (nitrate or nitrite) are absent (Barnard, 1976; Buchan, 1984). Sludge from the clarifier flows together with the influent wastewater into this zone. The anaerobic zone is

essential for the removal of phosphate, because the bacteria in the activated sludge passing through this zone are preconditioned to take up excess phosphate under aerobic conditions (Cloete and Muyima, 1997). The release of a certain quantity of phosphate from the biomass into the solution indicates that the bacteria have been suitably conditioned (Pitman, 1984). The retention time (about 1 h) of the influent wastewater is of extreme importance. Nitrates and dissolved oxygen discharge into the zone must be zero or as near to zero as possible at all times (Cloete and Muyima, 1997). The presence of nitrate in an anaerobic zone interferes with the phosphate removing capability of the activated sludge during aerobiosis (Barnard, 1976; Nicholls and Osborn, 1979; Marais *et al.*, 1983). The reason may be due to competition for substrate between phosphate-accumulating and denitrifying organisms. In the presence of nitrate, the redox potential is too high to produce fatty acids for the release of phosphate (Cloete and Muyima, 1997). However, the use of unsettled influent and the presence of sludge from the sludge treatment in the primary clarifiers, probably producing lower fatty acids, have a positive effect on the phosphate removal (Mulder and Rensink, 1987). The degree of nitrate feedback that can be tolerated depends on the strength of the sewage feed to the anaerobic zone and its readily biodegradable chemical oxygen demand (COD) concentration (Pitman, 1984).

The microorganisms found in the anaerobic zone are normally living in soil and water and are capable of fermentation (species of *Aeromonas*, *Citrobacter*, *Klebsiella*, *Pasteurella*, *Proteus* and *Serratia*). They accumulate and produce organic compounds such as lactic acid, succinic acid, propionic acid, butyric acid, acetic acid and ethanol during fermentation. These organic compounds serve as electron donor and acceptor, but cannot be utilized under anaerobic conditions. These organic compounds will only be consumed in the anoxic and aerobic zones. Therefore it seems as though the anaerobic zone provides substances for the proliferation of aerobic phosphate-accumulating bacteria (Fuhs and Chen, 1975; Buchan, 1984).

Phosphate release or uptake can be induced by certain carbon sources. The addition of acetate to the medium and the lowering of the pH, as well as phosphate starvation under anaerobic conditions, resulted in phosphate release (Fuhs and Chen, 1975; Barnard, 1976; Buchan, 1983). Nicholls and Osborn (1979) confirmed that bacteria relieved from stress conditions rapidly take up phosphate in an overplus reaction immediately on entering the aerobic zone, where stress conditions are relieved and phosphate together with an abundant source of energy is available. Phosphate seemed to be released from the acid-soluble fraction of cells and to a minimal degree from the RNA and DNA (Cloete and Muyima, 1997).

1.2.2 Primary anoxic zone

Anoxic refers to the presence of nitrates and the absence of dissolved oxygen (Buchan, 1984; Pitman, 1984; Streichan *et al.*, 1990). The primary anoxic zone is the main denitrification reactor in the activated sludge process. It is fed by effluent from the anaerobic zone and by mixed liquor recycled from the aerobic zone (Cloete and Muyima, 1997). The absence of oxygen and the presence of nitrate and nitrite lead to the enrichment of denitrifying bacteria, which reduce nitrate or nitrite to molecular nitrogen. Soluble and colloidal biodegradable matter is removed in the primary anoxic zone. It was found that phosphate release was induced in the anoxic zone in the presence of low fatty acids or their salts such as acetate, formate and propionate (Gerber *et al.*, 1986). Phosphate release was not only affected by the substrate but also by the dosage of the substrate concentrations of soluble, readily biodegradable carbon substrates (Cloete and Muyima, 1997).

1.2.3 Primary aerobic zone

The main function of the primary aerobic zone is to oxidize organic material in the sewage, to oxidize ammonia to nitrite and then to nitrate (by chemoautotrophs), and to provide an environment in which the biomass can take up all the phosphate released in the anaerobic zone, plus all the phosphate that enters the

process in the feed sewage (Cloete and Muyima, 1997). Ammonia is oxidized to nitrite by *Nitrosomonas*, *Nitrospira* and *Nitrosolobus* spp. Nitrite is oxidized to nitrate by *Nitrobacter*, *Nitrospira* and *Nitrococcus* spp. (Buchan, 1984). The principal operational determinant of the efficiency of phosphate removal seems to be the aeration rate, which should be sufficient to promote the rapid uptake of released and feed phosphate. The aeration rate should ensure the oxidation of the carbon compounds and ammonia and suppress the growth of filamentous microorganisms that produce poorly settling sludge (Pitman, 1984). High rate aeration resulted in maximum uptake of phosphate within 2 h (Levin and Shapiro, 1965; Shapiro, 1967, Carberry and Tenny, 1973). and colleagues Wentzel *et al.*, (1985) indicated that the excessive phosphate uptake in the aerobic stage was associated directly with the degree of phosphate release during the previous anaerobic phase, therefore, more phosphate released leads to more phosphate uptake. There is a linear relation between phosphate release and uptake (Wentzel and colleagues *et al.*, 1985).

Other parameters also play a role in enhanced phosphate removal such as: the presence of readily biodegradable compounds, especially volatile fatty acids (VFA), produced by fermentative bacteria from organic compounds in the influent. These volatile fatty acids are removed during the anaerobic phase and polymerized at the expense of energy obtained from the breakdown of polyphosphate (Cloete and Muyima, 1997). Poly- β -hydroxybutyric acid (PHB) plays an important role in the mechanism of phosphate uptake and release (Nicholls and Osborn, 1979; Deinema *et al.*, 1980; Lawson and Tonhazy, 1980; Comeau *et al.*, 1986). The presence of PHB in biopolyphosphate bacteria helps them to grow and rebuild their own polyphosphate by taking up soluble phosphate from the solution (Comeau *et al.*, 1986). PHB and polyphosphate will therefore play a mutually interdependent role to assist aerobic bacteria to survive through an anaerobic period (Nicholls and Osborn, 1979).

1.2.4 Secondary anoxic zone

The function of this zone is the removal of excess nitrates that were not removed in the primary anoxic zone. Denitrification in this zone is very slow and therefore the quantity of nitrate removed is small. Due to the low COD, the retention time in this anoxic zone is relatively long (Cloete and Muyima, 1997).

1.2.5 Secondary aerobic zone and clarifier

The main function of the secondary aerobic zone is to increase the dissolved oxygen to a level between 2 and 4 mg/l in the mixed liquor before it enters the clarifier (Barnard, 1976), and to refine the final effluent by the removal of additional phosphate and the oxidation of residual ammonia (Cloete and Muyima, 1997). Mixed liquor must be aerated for at least 1 h before it passes into the clarifier to promote phosphate uptake and maintain good aeration conditions (Cloete and Muyima, 1997). Excess aeration should be prevented as it would result in the conversion of organically bound nitrogen to nitrate and cause the slow aerobic release of phosphate from the solids (Keay, 1984; Pitman, 1984). Ammonia nitrification as well as phosphate removal which has not been completed, will continue in the secondary aerobic zone (Buchan, 1984).

The function of the clarifier is to produce a clear effluent free of suspended solids, and a thickened sludge for recycling to the inlet of the process. The quality of the underflow sludge should be such that nitrate is not recycled to the anaerobic zone (Ekama *et al.*, 1984).

The biomass of activated sludge is the active agent of biological wastewater treatment, responsible for carbonaceous material oxidation and nutrient removal. To date, process engineering has received the greatest attention and, in the interim, has practically become optimized. Yet systems based on EBPR principles still regularly fail to achieve the desired end result. This is due to limitations in our current understanding of the complexities of microbiological interactions occurring within the sludge, as well as our inadequate knowledge of microbial community structure-function correlations (Wagner *et al.*, 1993).

To describe and control these microbial processes and mechanisms, Wanner (1997) suggests that activated sludge should be characterized from the following viewpoints:

- characterization and quantification of microbial constituents according to metabolic activities
- identification and classification of microorganisms
- activated sludge quality, i.e. settleability, de-waterability.

Microbial–ecological studies of activated sludge are integral to creating a complete and more definitive understanding of the process, diversity and various functions performed by the constituent microflora. A review of our knowledge about the microbiology of biological phosphorus removal will now follow.

2 MICROBIAL POPULATION DYNAMICS OF BIOLOGICAL PHOSPHORUS REMOVAL

The microbial community of activated sludge consists of bacteria, protozoa, fungi, algae and filamentous organisms (Bux *et al.*, 1994; Muyima *et al.*, 1997). Since organic carbon is the most important energy source entering these systems, it can be expected that heterotrophic bacteria will dominate the community structure (Kämpfer *et al.*, 1996). A variety of methods have been used to study the microbial community in activated sludge. The results obtained using these methods today form the basis of our understanding of the microbiology of biological phosphorus removal and will hence be discussed in more detail.

2.1 Culture based techniques

Since enhanced phosphate removal was first postulated to be mediated by a biological mechanism (Srinath *et al.*, 1959; Fuhs and Chen, 1975), much research has centred around identifying and elucidating those organisms responsible for the process (Barnard, 1976; Hart and Melmed, 1982; Buchan, 1983; Cloete and

Steyn, 1988a, b; Bosch and Cloete, 1993; Kavanaugh and Randall, 1994; Nakamura *et al.*, 1998).

Some time ago, *Acinetobacter* came to be regarded as the model organism in biochemical models describing the mechanism of EBPR (Wentzel *et al.*, 1986) due mainly to its presence in high numbers and its favourable physiological characteristics when isolated from biological phosphorus removal plants using conventional plating techniques. Bacteriological studies of biological nutrient removal (BNR) systems have emphasized the functional role of *Acinetobacter* in enhanced P removal (Lötter, 1985). Development of enhanced cultures of polyphosphate accumulating organisms (PAOs) has shown tremendous specificity in the population structure when one considers that more than 90% of the organisms cultured aerobically from laboratory-scale UCT and three-stage Bardenpho systems were identified as *Acinetobacter* (Wentzel *et al.*, 1988, 1989).

Since Fuhs and Chen (1975) first implicated *Acinetobacter* spp. as having an important role in EBPR, most subsequent studies have focused on this bacterial genus. The reasons for this attention have not been entirely unjustified. Culture-dependent methods consistently indicated that *Acinetobacter* spp. were the numerically dominant members of EBPR systems. Furthermore, some, but not all, *Acinetobacter* strains isolated from activated sludge accumulated excessive amounts of polyphosphate in pure culture, suggesting their importance in the EBPR process (Bond *et al.*, 1995). However, researchers have difficulties reconciling the carbon and phosphorus transformations in pure cultures of *Acinetobacter* strains with the biochemical model for EBPR. In recent years, serious doubts have been raised as to the significance of *Acinetobacter* spp. in the EBPR process. The most compelling evidence for this change of view has been the recent non-culture-dependent studies of phosphate-removing communities. In all cases, *Acinetobacter* spp. was found to represent only a small portion of the total EBPR microbial population. Instead, other bacterial groups such as the Gram-positive bacteria and the beta-subclass groups of the Proteobacteria were numerically domi-

nant (Bond *et al.*, 1995). While the use of these non-culture-dependent methods reduces the significance of *Acinetobacter* spp. in EBPR processes, the resolution of the methods has not been sufficient to propose alternative EBPR candidate genera (Bond *et al.*, 1995).

A number of heterotrophic organisms have been isolated from EBPR systems using culture dependent methods. These organisms include *Acinetobacter*, *Moraxella*, *Pseudomonas*, *Micrococcus*, *Achromobacter*, *Aeromonas*, *Vibrio*, *Citrobacter*, *Pasteurella*, *Enterobacter*, *Proteobacter*, *Klebsiella*, *Bacillus* and coliforms (*Escherichia coli* and *E. intermedium*) (Hart and Melmed, 1982; Brodisch and Joyner, 1983; Cloete *et al.*, 1985a, b; Lötter, 1985; Lötter and Murphy, 1985; Lötter *et al.*, 1986; Cloete and Steyn, 1988a, b; Kavanaugh and Randall, 1994; Wagner *et al.*, 1994a, b; Momba and Cloete, 1996a, b; Ubukata and Takii, 1998).

We know that there is a large discrepancy between the total direct microscopic counts and viable plate counts (usually less than 1% of the former) for many ecosystems (Cloete and Steyn, 1987; Wagner *et al.*, 1993). Recoveries from activated sludge systems, even with optimized media, are only between 5 and 15% (Wagner *et al.*, 1993). The exact ecological significance of these organisms in EBPR is not understood. This is due to the fact that these organisms make up part of the culturable activated sludge microbial community. Consequently, researchers have been investigating the use of non-culture dependent techniques to obtain a better understanding of EBPR microbiology.

2.2 Fluorescent *in situ* hybridization (FISH)

All life on earth is comprised of three domains, i.e. the *Bacteria* (e.g. *Acinetobacter*, *Pseudomonas*), the *Archaea* (microorganisms thought to live in extreme environments, i.e. anaerobic, heat and salt) and the *Eukarya* (animals, fungi, plants, protists) (Woese *et al.*, 1990). This phylogenetic system is based on comparative analyses of sequences of the small subunit (16S) rRNA gene.

The domain (super-kingdom) *Bacteria* presently consists of eleven characterized bacterial phyla (divisions) (Woese, 1987; Wagner and

Amann, 1997). It is evident that when using the 16S rRNA molecule phylogenetically to structure the domain *Bacteria*, the arrangement within the various phyla becomes somewhat confusing, i.e. photosynthetic species grouped with non-photosynthetic species; anaerobes are paired with aerobes and heterotrophs with chemolithotrophs. Most of the taxa are defined by unique, conserved oligonucleotide sequences or signatures, i.e. rRNA sequence positions that are not found in the rRNA of other groups (Manz *et al.*, 1992). Many of the traditional Gram-negative bacteria are contained in the *Proteobacteria* class (Woese, 1987; Manz *et al.*, 1992). At least four distinct subdivisions exist in the purple bacteria, designated alpha (α), beta (β), gamma (γ) and delta (δ) (Woese, 1987).

Each rRNA-targeted oligonucleotide is chemically linked to a fluorochrome molecule which allows cells hybridized with the fluorescently-labelled oligonucleotide to be directly visualized using epifluorescent microscopy or scanning confocal laser microscopy (Amann, 1995; Amann *et al.*, 1995). The process usually incorporates a total cell count using the DNA intercalating dye 4,6-diamidino-2-phenylindole (DAPI) followed by hybridization with the universal bacterial probe, EUB (the fluorescing stain 3,6-tetramethyl diaminoacridine or acridine orange can also be used to visualize microscopically DNA molecules, but there are a number of difficulties inherent to its application; refer to Porter and Feig (1980) for a brief review). DNA-DAPI complexes fluoresce bright blue when visualized at wavelengths of > 390 nm, while unbound DAPI and non-DNA-DAPI complexes fluoresce a weak yellow which enhances the visibility of DAPI fluorescence (Porter and Feig, 1980).

The EUB/DAPI ratio gives an indication as to the bacterial composition of the sample. Dual EUB/DAPI staining of activated sludge samples (Wagner and Amann, 1997) has revealed that approximately 80% of the microbial cells present in the samples were metabolically active bacteria, of which only 3–19% could be cultivated on optimized media. The specificity of oligoprobes is freely adjustable, dependent upon the requirements of the

user. Different phylogenetic levels ranging from kingdom, e.g. *Bacteria* (EUB), to subclass, e.g. gamma (GAM) subclass of *Proteobacteria*, to genus, e.g. *Acinetobacter* (ACA), species and subspecies can be probed, the degree of organization required being the only limitation (Wagner *et al.*, 1994a) (Table. 23.2).

In situ identification of the organisms in a batch-type EBPR sludge has revealed that the four major bacterial groups present were the alpha and beta subclass of *Proteobacteria*, Gram-positive bacteria with a high G + C content (GPBHGC) and bacteria belonging to the *Cytophaga-Flavobacterium* cluster of the *Cytophaga-Flavobacterium-Bacteroides* phylum (Kawaharasaki *et al.*, 1999). However, Wagner *et al.*, (1994a), when characterizing the *Proteobacterial* microbial consortia in municipal mixed liquor from Hirblingen, Germany, found that the beta subclass dominated over the alpha and gamma subclasses. Simultaneous plating of the sludge on nutrient-rich medium showed dominance of the gamma subclass of *Proteobacteria* emphasizing the bias introduced with cultivation techniques.

In situ hybridization of mixed liquor samples at the family level of organization from the anaerobic and aerobic zones of the same plant in Germany have revealed the following trends (Snaidr *et al.*, 1997):

anaerobic basin	α - <i>Proteobacteria</i>	= 11%
	β - <i>Proteobacteria</i>	= 24%
	γ - <i>Proteobacteria</i>	= 5%
	GPBHGC	= 24%
	<i>Cytophaga-Flavobacterium</i>	= 9%
aerobic basin	α - <i>Proteobacteria</i>	= 9%
	β - <i>Proteobacteria</i>	= 26%
	γ - <i>Proteobacteria</i>	= 10%
	GPBHGC	= 19%
	<i>Cytophaga-Flavobacterium</i>	= 8%

These results show conclusively that there is no markable population shift between zones in an EBPR process when characterized at family level. Whether a shift occurs at species level requires verification, but regardless of any shifts which may occur, the significance of the *Proteobacteria* to efficient EBPR operations is evident.

TABLE 23.2 Phyla and subdivisions (including representative examples) of the domain *Bacteria* (shaded rows indicate those subdivisions known to be implicated in EBPR)

<i>Proteobacteria</i> (Purple bacteria)
α subdivision (<i>Nitrobacter</i>)
β subdivision (<i>Alcaligenes</i> , <i>Pseudomonas testoteroni</i> , <i>P. cepacia</i> , autotrophic nitrifiers)
γ subdivision (<i>Acinetobacter</i> , <i>Aeromonas</i> , fluorescent pseudomonads, Enterics, <i>Vibrio</i>)
δ subdivision (sulphate-reducing bacteria and myxobacteria)
<i>Firmicutes</i>(Gram-positive bacteria)
A. <i>Actinobacteria</i> (high G + C species) (<i>Arthrobacter</i> , <i>Micrococcus</i> , <i>Microtholunatus</i>)
B. <i>Bacillus</i> / <i>Clostridium</i> group (low G + C species) (<i>Clostridium</i> , <i>Bacillus</i>)
C. Photosynthetic species (<i>Heliobacterium</i>)
D. Species with Gram-negative walls (<i>Megasphaera</i>)
Cyanobacteria and chloroplasts
<i>Nostoc</i> , <i>Oscillatoria</i>
Spirochetes and relatives
A. Spirochetes (<i>Spirochaeta</i>)
B. Leptospiras (<i>Leptospira</i>)
Green sulphur bacteria
<i>Chlorobium</i>
Bacteroides, flavobacteria and relatives
A. Bacteroides (<i>Bacteroides</i>)
B. Flavobacterium group (<i>Flavobacterium</i> , <i>Cytophaga</i>)
Planctomyces and relatives
A. Planctomyces (<i>Planctomyces</i>)
B. Thermophiles (<i>Isocystis</i>)
Chlamydiae
<i>Chlamydia</i>
Radioreistant micrococci and relatives
A. Deinococcus group (<i>Deinococcus radiodurans</i>)
B. Thermophiles (<i>Thermus aquaticus</i>)
Green non-sulphur bacteria and relatives
A. Chloroflexus group (<i>Chloroflexus</i>)
B. Thermomicrobium group (<i>Thermomicrobium roseum</i>)
Thermotogoa

Probing activated sludge with fluorescently-labelled oligonucleotide probes specific for the alpha, beta and gamma subclasses of the *Proteobacteria* (see section 2.5) has revealed that the microbial consortia are dominated by the *Proteobacteria* (approximately 80%), a phylum containing the majority of the traditional Gram-negative bacteria (the majority of which are heterotrophic) (Wagner *et al.*, 1993).

Staining activated sludge samples with DAPI at elevated concentrations to those used for DNA staining – referred to as the polyphosphate-probing concentration – results in the fluorescence of intracellular volutin and lipid inclusions (Nakamura *et al.*, 1998; Streichan *et al.*, 1990, as cited by Kawaharasaki *et al.*, 1999). Bacteria that accumulate large quantities of poly-P are easily distinguished by colour and intensity of fluorescence due to the following DAPI stain characteristics:

1. DNA-DAPI fluorescence is blue-white
2. polyphosphate-DAPI fluorescence is bright yellow
3. lipid-DAPI fluorescence is weak yellow and fades rapidly (in the space of seconds) (Kawaharasaki *et al.*, 1999).

Dual staining of samples with DAPI at elevated concentrations and EUB will result in the determination of the PAO population in EBPR sludges, i.e. PAO/EUB ratio. *In situ* identification of those bacteria exhibiting strong poly-P accumulation (identified through DAPI staining) can also be achieved through dual staining with class, i.e. *Proteobacteria*, and subclass, i.e. alpha, beta or gamma, oligonucleotide probes which totally negates culture-dependent methods of isolation and/or Neisser (poly-P) staining procedures.

For all its promise and potential, however, there are still technical problems inherent to FISH when applying the technology to microbial systems such as activated sludge. Qualitatively, the protocol of probe hybridization and detection has been optimized at all levels of organization, but due to the nature of activated sludge, quantitative results are often difficult and limited, i.e. complete dispersion of sludge flocs remains a technical problem which

limits the application. Other problems include deoxyribonucleic acid (DNA) retrieval for sequence determination, polymerase chain reaction (PCR) biases when amplifying the sequence of interest and an imposed selection of the retrieved or target sequences (Hiraishi *et al.*, 1998). One of the possible solutions directed towards these problems is the combination of molecular and biomarker methods.

2.3 Immunological techniques

The immunofluorescence approach was introduced as an *in situ* identification technique, prior to FISH, in an attempt to avoid culture-dependent techniques and has been used effectively to identify *Acinetobacter* in activated sludge samples (Cloete *et al.*, 1985a, b; Lötter and Murphy, 1985; Cloete and Steyn, 1987, 1988b). Cloete and Steyn (1987) found that less than 10% of microscopic cell counts of activated sludges could be accounted for in agar plate enumerations. Although the technique is highly specific for the bacterium in question, there are a number of limitations associated with it (Wagner and Amann, 1997). First, the presence of extracellular polymeric substances in activated sludge flocs can inhibit the penetration of antibodies to the target cells; the method of raising antibodies in host animals requires initially culturing the bacterium of interest; and cross-reaction of antibody with contaminants does occur, resulting in high levels of background fluorescence (Wagner and Amann, 1997).

Monoclonal antibody production is the method of choice to obtain immunoglobulins against proteins that cannot be purified or that are available in low amounts (Drenckhahn *et al.*, 1993). Antibodies have been used in investigations of samples from activated sludge systems, although such studies are not numerous. Cloete and Steyn (1987) used a fluorescent antibody technique for the identification and enumeration of *Acinetobacter* in activated sludge (Cloete and Steyn, 1988b). Monoclonal antibodies specific for *Nitrobacter* and *Nitrosomonas* respectively were produced and used for estimation of cell numbers in activated sludge (Sandén *et al.*, 1994).

Conventional as well as focused immunization strategies were also used to maximize the possibility of producing antibodies with the required ability to discern between the antigen preparations. A significant finding emerging from this study was based on the observation that antigenic differences were clearly detected between the aerobic zones of two activated sludge systems with differing phosphate removal ability. Characterization of the antigen recognized suggested a protein nature. The antibody showed five compact bands on the blot, with M_r appearing to be multiples of 18 kDa. The epitope recognized probably occurred on an 18 kDa proteinaceous monomer, represented on the blot in incremental steps of subunit association. The *raison d'être* of the unique antigen in the phosphate removing system may be speculated upon: first, did the specific conditions prevailing in the phosphate removing system as opposed to the non-phosphate removing system induce its expression, or did the anaerobic zone select for a unique bacterial population in the phosphate removing system which is recognized by the antibody, but is not necessarily correlated with phosphate removal? Could the state of oligomerization of the 18 kDa protein correlate to the phosphate removing ability of the sludge? All these and other questions still need to be answered, before any valid conclusions can be made using this technique.

2.4 Quinone profiles

Respiratory or isoprenoid quinones are a class of lipids that are constituents of bacterial plasma membranes. They play important roles in electron transport, oxidative phosphorylation and active transport across the membrane (Collins and Jones, 1981). The numerical analysis of lipoquinone profiles has offered an effective method for monitoring population shifts and for classifying bacterial communities in wastewater sludges (Hiraishi *et al.*, 1991). Quinones are usually extracted from an environmental sample using an organic solvent. After evaporation and re-extraction, the concentrated quinone is applied to column chromatography to separate menaquinone and

ubiquinone (Hiraishi *et al.*, 1998). Municipal sludges are usually characterized according to their menaquinone and ubiquinone components (Hiraishi *et al.*, 1989, 1998). Quinone components are then identified and quantified using spectrochromatography and mass spectrometry. Numerical analyses of quinone profiles can enhance the information regarding bacterial community dynamics in wastewater ecosystems. The strength of the technique lies not only in its ability to assess taxonomic structure of bacterial communities, but also in that variations in bacterial population structure over space and time can be quantified (Hiraishi *et al.*, 1991). The use of respiratory quinone profiles to characterize the bacterial population structure of the anaerobic-aerobic activated sludge system showed that *Acinetobacter* species were not important in the system (Hiraishi *et al.*, 1989). However, it was shown that *Acinetobacter*, as detected by the biomarker diaminopropane, was the dominant organism only in wastewater treatment plants with low organic loading (Auling *et al.*, 1991).

2.5 Microautoradiography

Autoradiography has classically been used in the medical field but has recently been introduced to environmental sample analyses to determine microbial community structures. Typically, a radio-labelled compound appears in the cell or biological structure of interest through adsorption of a tracer or labelled substrate uptake. The radio-labelled sample is then placed in contact with a radiosensitive emulsion and the emissions from the radioactive sample interact with silver bromide crystals in the emulsion. The emulsion is then developed using standard photographic procedures and the silver grains appear on top of the radioactive structure, which can then be viewed microscopically (Nielsen *et al.*, 1998, 1999). Although autoradiography can successfully be applied to study the *in situ* physiology of various microorganisms, it is limited by its lack of proper identification of the organisms in question. However, Nielsen *et al.*, (1998), through simultaneous use of autoradiography and FISH, were able to correlate function/

activity with identification, which is a tremendous breakthrough for activated sludge identification-diversity-functional studies.

2.6 SDS-PAGE of proteins used for fingerprinting microbial communities

The extraction of proteins directly from environmental samples is desirable for multiple reasons:

1. Analysis of proteins extracted from environmental samples may help characterize the response of microbial communities to stressful conditions such as contamination with toxic chemicals, starvation, heat or oxygen levels
2. Analysis of total proteins extracted from an environmental sample can be employed as a 'fingerprint' to type the diversity in the sample, in a way similar to grouping of bacteria according to enzyme polymorphisms and immunological reactions. Such fingerprints may eventually be used to monitor the deterioration or enrichment of species diversity in microbial communities
3. The abundance of proteins to which specific antibodies are available can be directly measured in total proteins extracted from complex ecosystems and used as an index for monitoring the progress of a biocatalytic reaction *in situ* (Ogunseitan, 1993).

The main advantages of numerical analysis of electrophoretic patterns of large numbers of bacteria are:

1. rapid grouping
2. allocation of unknown microorganisms to a group and its possible identification
3. storage of large numbers of patterns in data banks for reference
4. a quick decision on whether two colony types in a culture are due to variation or contamination
5. information on epidemiology, spreading of animal and plant pathogens
6. DNA:DNA homology determinations of large numbers of strains can be reduced to hybridization of DNAs from the typical

representative of each group, previously established by gel electrophoresis (Kerstens and De Ley, 1975).

Similarities are calculated between each sample using a suitable similarity coefficient, the Pearson product moment correlation coefficient (r). The resultant matrix is clustered using the average linkage algorithm (also known as unweighted pair group method of arithmetic averages, UPGMA) to provide a sorted similarity matrix or dendrogram.

Valuable information concerning the bacterial population structure of activated sludge was obtained when sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) was used. The results confirmed previous studies performed by Cloete and Steyn (1987), which indicated that the bacterial population of activated sludge stayed the same throughout the system (Ehlers and Cloete, 1999a). Furthermore, no specific protein pattern due to seasonal changes or between different zones or between N- and P-removing and N-removing systems were observed using SDS-PAGE (Ehlers and Cloete, 1999a, b, c). The main drawback of this technique was that it was not sensitive enough to determine the difference in protein profiles of P-removing and non-P-removing bacterial populations.

2.7 Community-level carbon source utilization

Garland and Mills (1991) introduced the use of community-level carbon source utilization patterns for comparison of microbial communities from different habitats. The Biolog system (Biolog Inc., Hayward, USA) is based on the different utilization of a large number of organic compounds by the test organisms.

The Biolog system has already been used for characterization of naturally occurring bacteria and for classification of bacterial communities of different environments (Frederickson *et al.*, 1991; Garland and Mills, 1991; Verniere *et al.*, 1993; Winding, 1994; Zak *et al.*, 1994). The comparisons of Biolog results with other test systems, i.e. API and Biotype-100, have also been performed (Frederickson *et al.*, 1991; Amy

et al., 1992; Klinger *et al.*, 1992; Verniere *et al.*, 1993). Community-level physiological profiles (CLPP) have been used to characterize microbial communities in freshwater, coastal and lagoon areas, as well as soil, rhizosphere, bioreactors and gnotobiotic mixtures (Garland and Mills, 1991; Gorlenko and Kozhevin, 1994; Winding, 1994; Zak *et al.*, 1994; Bossio and Scow, 1995; Ellis *et al.*, 1995; Haack *et al.*, 1994; Lehman *et al.*, 1995; Vahjen *et al.*, 1995; Garland, 1996a, b; Glimm *et al.*, 1997; Engelen *et al.*, 1998). Environments such as groundwater, phyllosphere, activated sludge and compost were also studied (Ellis *et al.*, 1995; Heuer *et al.*, 1995; Lehman *et al.*, 1995; Insam *et al.*, 1996; Victorio *et al.*, 1996). For rapidly assessing the dynamics of autochthonous microbial communities, the CLPP techniques have been used (Wünsche and Babel, 1995). Diversity can be calculated using univariate indices which do not capture all potential differences in the CLPP (Garland, 1997).

The purpose of using Biolog was therefore not to try to detect each and every metabolic reaction of all the species in the community, but the collective utilization pattern for a specific community. A high species diversity should lead to a higher relative number of substrates utilized, because there are more possibilities and upon dilution, some organisms will be lost (causing a decrease in species diversity) from the community, depending on their abundance and the relative contribution (perhaps only one metabolic reaction in the system), reducing the number of possibilities. The extent of the reduction of the possibilities upon dilution, should theoretically reflect something about the community structure. The key, therefore, lies in the interpretation of the results.

Differences in microbial community structure in activated sludge systems may exist, but this had no bearing on the effectivity with which these systems removed phosphate. If PO_4^{3-} removal was related to the microbial community composition, a high correlation of the Biolog patterns would be expected among PO_4^{3-} removing systems. However, this was not the case. All the different zones of systems tested indicated a high initial diversity (10^{-1} to 10^{-2} dilutions), due to the high number of substrates utilized. No specific patterns could,

however, be identified for PO_4^{3-} removing systems, indicating that PO_4^{3-} removal was not community structure specific. This agrees with previous studies (Dold *et al.*, 1980; Kersters *et al.*, 1997).

2.8 Biomass

Current research in wastewater treatment has been directed towards mathematical modelling of basic design and operational procedures. One important parameter in such models is the amount of viable biomass. For this reason attempts have been made to find simple and reliable methods to determine the biomass in wastewater and activated sludge. The simplest and most often used method is to measure suspended solids or volatile suspended solids. Such methods, however, do not distinguish between living cells and debris of either organic or inorganic origin. Using traditional plate techniques, the problem is normally an underestimation of the biomass due to selectivity of the media (Jørgensen *et al.*, 1992).

One key component of mixed liquor suspended solids (MLSS) is the heterotrophic active biomass. This component mediates the biodegradation processes of COD removal and denitrification. Thus the rates for these processes are directly related to the heterotrophic active biomass present, and the specific rates should be expressed in terms of this parameter to allow a meaningful comparison of the rates measured in different systems. However, the heterotrophic active biomass parameter has been only hypothetical within the structure of these models; it has not been measured directly, primarily due to the lack of suitable simple measurement techniques. Ubisi *et al.*, (1997) have, however, reported close correlation between measured heterotrophic active biomass concentration with those calculated theoretically, thus promoting confidence in the application of the models for design, operation and control of activated sludge systems.

It was indicated that biomass was related to phosphorus removal (Bosch, 1992; Momba, 1995; Muyima, 1995). The higher the biomass, the better the P removal. This suggested that the main difference between P-removing and

non-P-removing systems was biomass related and not due to the microbial community structure. An increase in biomass led to an increase in P removal (Momba and Cloete, 1996a). When calculating the quantity of P removal per g of sludge no significant difference was observed, indicating that there was a direct relationship between P removal and MLSS for a specific system. The effects of growth phase and initial cell concentration of *Acinetobacter* spp. on P release and uptake have also been investigated (Momba and Cloete, 1996a, b; Rustrian *et al.*, 1997). As can be expected, Momba and Cloete (1996b) found a linear relationship existed between initial biomass and phosphate uptake and that at low cell densities (10^2 – 10^5 per ml) there was a net release of P rather than uptake. The physiological state of the cells also determines the response of the biomass to P accumulation. Bosch and Cloete (1993) have observed that P is accumulated at the end of log and during stationary phase once active growth has ceased and concluded that a maximum number of cells in the stationary phase of growth should be accumulated in the aerobic zone to optimize bio-P removal. Rustrian *et al.* (1997) found that *Acinetobacter* released more P under anaerobic conditions when in the stationary phase of

growth yet P uptake was equally efficient with cells in the log and stationary phase. These results can have a direct implication on EBPR operations as growth conditions for the polyphosphate accumulating organisms (PAO) community can be better optimized to improve and promote the mechanism.

2.8.1 *Microthrix phosphovorus* – the new model organism?

Due to the controversy surrounding the role of *Acinetobacter* in P removal operations (Table 23.3), microbiologists have, for the past decade, attempted to isolate other bacteria in activated sludge upon which the mechanism can be modelled.

Recently, a new P-removing bacterium from a laboratory-scale activated sludge system in Japan was isolated and identified as *Microthrix phosphovorus* (*M. phosphovorus*) (Nakamura *et al.*, 1995, as cited by Ubukata and Takii, 1998). The bacterium shows all the physiological traits characteristic of PAOs, i.e. P accumulation mechanism is only induced in sequential anaerobic/aerobic systems, and has the propensity to accumulate P to a maximum of 23% dry weight (luxury uptake). Its carbon and phosphorus transformation patterns coincide with those of EBPR sludge and, as such, the bacterium has been considered as a candidate for the model PAO in EBPR processes. *M. phosphovorus* is a Gram-positive coccus (diameter = 1.7–2.1 μm) and sequencing shows a high G + C genomic DNA content (65.6 mol%; phylogenetically belongs to the Gram-positive high GC (HGC) bacteria) (Ubukata and Takii, 1998).

The excess P accumulation mechanism is inducible in the bacterium, a useful feature when examining and attempting to define bio-P removal processes. However, the bacterium's dominance in EBPR processes has yet to be demonstrated and, because it cannot readily be isolated from activated sludge (as opposed to *Acinetobacter*), its application to BPR studies may be limited.

2.8.2 The 'G' bacteria

When assessing EBPR efficiency, cognisance must be taken of the 'G' bacteria, a group of

TABLE 23.3 Numbers of *Acinetobacter* spp. in activated sludge using different enumeration techniques

Number of <i>Acinetobacter</i> cells (% of total community)	Enumeration/identification technique	Reference
54–66%	API 20E	Hart and Melmed, 1982
90%	total plate counts and API 20E	Lötter <i>et al.</i> , 1986a
5–11%	MPN and API 20E	Kavanaugh and Randall, 1994
<10%	fluorescent antibodies	Cloete and Steyn, 1987, 1988b
3–6%	quinone profiles	Hiraishi <i>et al.</i> , 1989
<1%	16S rRNA	Bond <i>et al.</i> , 1997
	oligonucleotide probes	
6%	16S rRNA	Wagner <i>et al.</i> , 1994b
	oligonucleotide probes	

normal inhabitants found in activated sludge mixed liquor. There is a microbial community in activated sludge which is capable of organic substrate uptake and assimilation in the anaerobic zone with subsequent metabolism of these storage granules in the aerobic zone. This community is composed of two distinct groups of organisms with distinctly different modes of substrate uptake and synthesis of storage granules. The first group are the PAOs, the metabolism of which has been previously discussed. The second group are the 'G' or glycogen accumulating organisms (GAOs). In conventional systems GAOs are involved in normal organic oxidation processes but, in selector systems such as BPR, their presence and impact to system efficiency can become more prominent. These Gram-negative cocci grow as tetrads and are able to out-compete PAOs in anaerobic/aerobic systems by accumulating polysaccharide and not polyphosphate in the aerobic zone (Cech and Hartman, 1993; Cech *et al.*, 1994; Maszenan *et al.*, 1998). The 'G' bacteria are thought to be able to compete effectively with fermentative organisms for RBCOD and PAOs for VFA in the anaerobic zone as they are able to obtain the necessary reducing power and energy required for uptake through glycolysis, i.e. the Embden-Meyerhof pathway. Their proliferation in an EBPR system will eventually lead to a decline so far as phosphate removal is concerned. The 'G' bacteria were able to dominate in an anaerobic-anoxic system when the influent consisted of an acetate-glucose mixture yet, when acetate was used as substrate alone, the PAOs were able to dominate their competitors (Cech and Hartman, 1993). Influent P/COD (P^{ti}/S^{ti}) ratios also affects the microbial community structure of EBPR activated sludge (Lui *et al.*, 1997). A low P^{ti}/S^{ti} ratio enriches for GAOs, while high P^{ti}/S^{ti} ratios promote the growth of PAOs and suppress the proliferation of GAOs in the process (Lui *et al.*, 1997). Competing with the non-PAOs, i.e. fermentative bacteria, for RBCOD in the anaerobic zone reduces VFA production which ultimately influences P release by the PAOs. With the advent of molecular techniques for elucidating bacterial identification and relatedness, it has been

shown that all the Gram-negative tetrad cocci in activated sludge are closely related taxonomically, belonging to the alpha subdivision of the *Proteobacteria* (see section 2.5) (Maszenan *et al.*, 1998). According to Blackall *et al.*, (1997), independently isolated 'G' bacterial strains from the Cech and Hartman (1993) study and from full-scale plants in Italy showed near identical homogeneity to one another and 93% homogeneity to *Rhodobacter* (alpha subdivision of *Proteobacteria*), based on 16S rDNA sequences. They proposed the generic and species name, *Tetracococcus cechii*, for the two strains. Maszenan *et al.*, (1998) proposed a new genus (*Amaricoccus*) to house four 'G' bacterial isolates from Australia, Italy and Macau. Phylogenetic classification of these isolates in the *Proteobacteria* phylum of the domain *Bacteria* shows their relatedness to many of the PAOs, indicating that some relationship between the two competing bacteria does exist. This implies that an identical or similar mode of metabolism between the two bacterial types must have existed at some stage of their evolution.

3 CONCLUDING REMARKS

The microbiology of nitrification and denitrification in the activated sludge process is well understood. Nitrification is the process where ammonia, the reduced form of nitrogen is oxidized by autotrophic nitrifying bacteria to nitrite and nitrate. Ammonia is oxidized to nitrite by *Nitrosomonas*, *Nitrosospira* and *Nitrosolobus* spp. (Buchan, 1984). Nitrification is a chemolithotrophic process where microorganisms utilize energy generated from the oxidation of inorganic compounds (Cloete and Muyima, 1997). When reduced nitrogen is incorporated into newly synthesized biomass the process is termed assimilative nitrate reduction (Cloete and Muyima, 1997). When nitrate nitrogen is reduced to elementary nitrogen and serves as an electron acceptor, the process is known as denitrification (Cloete and Muyima, 1997). A wide range of heterotrophic bacteria can accomplish denitrification under anoxic conditions. Nitrite is oxidized to nitrate by *Nitrobacter*, *Nitrospira* and *Nitrococcus* spp. (Buchan, 1984).

Culture dependent methods have yielded a variety of microbial isolates. However, due to the limitations of culture-based techniques, the ecological role of these microorganisms is not well understood.

The immunofluorescence approach was introduced as an *in situ* identification technique, prior to FISH, in an attempt to avoid culture-dependent techniques and has been used effectively to identify *Acinetobacter* in activated sludge samples (Cloete *et al.*, 1985a, b; Lötter and Murphy, 1985; Cloete and Steyn, 1987, 1988b). Cloete and Steyn (1987) found that less than 10% of microscopic cell counts of activated sludges could be accounted for in agar plate enumerations. Although the technique is highly specific for the bacterium in question, there are a number of limitations associated with it (Wagner and Amann, 1997). *In situ* identification of the organisms using FISH in a batch-type EBPR sludge has revealed that the four major bacterial groups present were the alpha and beta subclass of *Proteobacteria*, Gram-positive bacteria with a high G + C content and bacteria belonging to the *Cytophaga-Flavobacterium* cluster of the *Cytophaga-Flavobacterium-Bacteroides* phylum (Kawaharasaki *et al.*, 1999). However, Wagner *et al.*, (1994a), when characterizing the *Proteobacterial* microbial consortia in municipal mixed liquor from Hirblingen, Germany, found that the beta subclass dominated over the alpha and gamma subclasses. Simultaneous plating of the sludge on nutrient rich medium showed dominance of the gamma subclass of *Proteobacteria* emphasizing the bias introduced with cultivation techniques.

The use of respiratory quinone profiles to characterize the bacterial population structure of the anaerobic-aerobic activated sludge system showed that *Acinetobacter* species were not dominant in the system (Hiraishi *et al.*, 1989). However, it was shown that *Acinetobacter*, as detected by the biomarker diaminopropane, was the dominant organism in wastewater treatment plants with a low organic loading (Auling *et al.*, 1991).

Due to the controversy surrounding the role of *Acinetobacter* in P removal operations microbiologists have, for the past decade, attempted to isolate other bacteria in activated sludge

upon which the mechanism can be modelled. Recently, a new P-removing bacterium from a laboratory-scale activated sludge system in Japan was isolated and identified as *Microcystis phosphovorans* (*M. phosphovorans*) (Nakamura *et al.*, 1995, as cited by Ubukata and Takii, 1998). The bacterium shows all the physiological traits characteristic of PAOs, i.e. P accumulation mechanism is only induced in sequential anaerobic/aerobic systems, and has the propensity to accumulate P to a maximum of 23% dry weight (luxury uptake). However, the bacterium's dominance in EBPR processes has yet to be demonstrated and, because it cannot readily be isolated from activated sludge (as opposed to *Acinetobacter*), its application to BPR studies may be limited.

The 'G' bacteria are thought to be able to compete effectively with fermentative organisms for RBCOD and PAOs for VFA in the anaerobic zone as they are able to obtain the necessary reducing power and energy required for uptake through glycolysis, i.e. the Embden-Meyerhof pathway. Their proliferation in an EBPR system will eventually lead to a decline as far as phosphate removal is concerned.

Bacterial community composition studies using SDS-PAGE, Biolog and biomass indicated that there were no differences in the bacterial community structure between different zones in the same systems, as well as between different activated sludge systems. These results indicated that EBPR was, therefore, not due to a specific bacterial community structure but rather due to the amount of biomass in the system. The higher the biomass, the higher the phosphorus removal capacity of an activated sludge system.

In conclusion the ultimate method to determine the bacterial community structure and function of environmental samples still has to be developed. Therefore, each possible method should be investigated, until one or a combination of methods is found, that can assist in the better understanding of microbial community structure and function in activated sludge and specifically biological phosphorus removal.

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Anaerobic treatment processes

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1 INTRODUCTION

The term anaerobic digestion is used to describe many different anaerobic waste treatment processes. The basic principle common to them all is the fact that the biological degradation of the organic components in the waste is achieved with no requirement for molecular oxygen (air). In most cases, anaerobic digestion processes are also methanogenic, i.e. most of the carbon atoms originating in the waste material are reduced to methane (CH₄), the ultimate product of biological metabolism in anaerobic environments. The economic value of methane produced by anaerobic digestion can be a major consideration in the selection of this treatment technology, allowing significant levels of energy recovery for space heating, lighting and power generation. Although the anaerobic degradation of organic pollutants may be achieved without methane formation, e.g. when denitrifying and sulphate-reducing bacteria utilize nitrate and sulphate as terminal electron acceptors, the term anaerobic digestion generally refers to the methanogenic process in which carbon dioxide acts as the terminal electron acceptor.

The feasibility of anaerobic digestion has been demonstrated with many different types of waste-stream, however, only a relatively small proportion of these have shown consistent advantages over aerobic treatment technologies when implemented at the commercial scale. As a result, the range of proven applications is presently small compared to the number of potential applications and anaerobic digestion is generally restricted to the treatment of high strength wastes having a relatively

consistent composition. These include food and paper processing wastewaters, sewage sludge and farm manures, crop residues, and several other specific industrial wastewaters. The maximum organic loading rate employed in these cases is often in excess of an order of magnitude higher than that in an equivalent aerobic process. Furthermore, many organic pollutants, such as pesticides, halogenated compounds and the complex chemical cocktails present in landfill leachates, are not particularly amenable to aerobic degradation pathways but may be readily treated anaerobically.

As an emerging technology, the success of anaerobic digestion was initially hampered by unsatisfactory reactor designs that had been inherited from conventional (aerobic) processes. These failed to optimize treatment efficiency and were often the cause of process failure. Recent developments in anaerobic reactor design have been instrumental in allowing the full potential of anaerobic digestion to be achieved and these are discussed later in more detail.

Until recently, the complex microbial ecology of the sludge (biomass) in anaerobic digestion systems was poorly understood, and this has undoubtedly played its part in the slow move towards the exploitation of this technology in effluent treatment. These issues have now been fully elucidated over the last two decades which has allowed modern full-scale treatment plants to be designed, monitored and controlled with a high degree of confidence in their performance characteristics.

This chapter begins with an overview of the microbiology and biochemistry unique

to the anaerobic digestion process before describing the key physiological and environmental factors that affect microbial activity and system performance. The final section outlines the range of reactor designs currently employed in the anaerobic digestion of wastewaters.

2 MICROBIOLOGY OF ANAEROBIC DIGESTION

Methane production is the final step in a cascade of biochemical reactions taking place within anaerobic digesters treating the organic components of sludges and wastewaters. However, no individual microorganism is capable of carrying out all of these reactions independently. Consequently, anaerobic treatment processes are complex ecosystems comprising several diverse microbial guilds (organisms that have the same specialist role in their community) that work together in a coordinated manner to convert the organic components to methane and carbon dioxide. Early concepts of anaerobic digestion recognized that the transformation process would involve at least two stages, initial acidification of the complex organic matter followed by gas formation from simpler intermediates (Fig. 24.1).

As a greater understanding of the process developed, alternative new schemes were proposed that detailed both substrate conversion and flux in the conversion of organic materials to methane and carbon dioxide. A well-accepted scheme is presented in Fig. 24.2 (Gujer and Zehnder, 1983) and according to this, six distinct processes may be identified in an anaerobic digester:

1. Hydrolysis of biopolymers:
 - (i) hydrolysis of proteins
 - (ii) hydrolysis of carbohydrates
 - (iii) hydrolysis of lipids

2. Fermentation of amino acids and sugars
3. Anaerobic oxidation of long chain fatty acids and alcohols
4. Anaerobic oxidation of intermediary products such as volatile acids, except acetate
5. Conversion of acetate to methane
6. Conversion of hydrogen to methane.

Since none of the above steps occurs through spontaneous chemical reactions to any significant degree, the individual steps are catalyzed by separate groups of microorganisms, each having its own specific role within the overall process. Such microbial consortia effect complete breakdown of complex organic substrates through series metabolism, a process involving the sequential flow of substrates and products from one microbial guild to another. The following classification illustrates the bacterial consortium of anaerobic digestion and highlights the relationships between each of the bacterial groups concerned.

The first microbial group is composed of hydrolytic bacteria. These break down lipids, complex polymeric molecules (e.g. protein and carbohydrate) and particulate organic matter into simpler soluble components such as short-chain fatty acids, glycerol, peptides, amino acids, oligosaccharides and sugars.

The second group is designated the acid-forming bacteria and consists of both acidogenic (organic acid forming) and acetogenic (acetate forming) bacteria. These acid-forming bacteria convert the end-products from the first group into both the key substrates of methanogenesis, namely acetate, hydrogen, carbon dioxide, and a number of minor intermediary products such as formate, propionate, butyrate, valerate etc.

The sequence is completed by the third group of bacteria, the methanogens, which consume the end-products from the second group of bacteria and convert them to the final

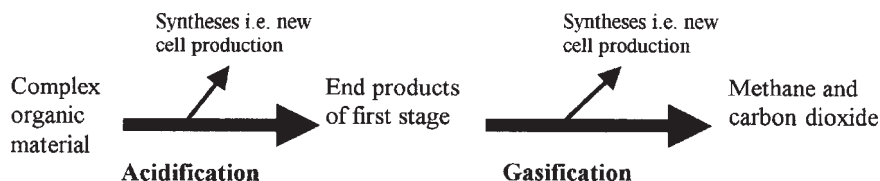


Fig. 24.1 Simple schematic representation of the anaerobic digestion process.

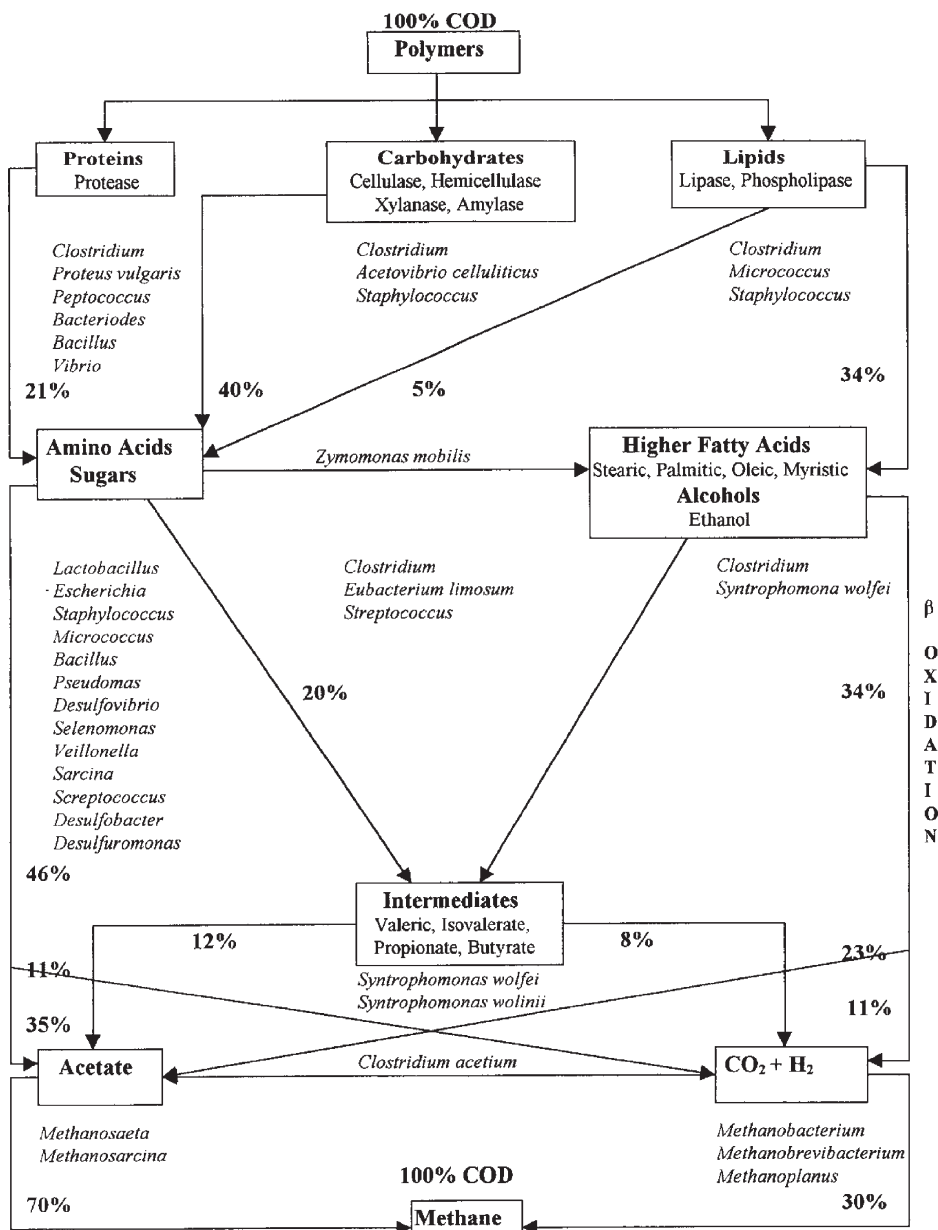


Fig. 24.2 Carbon flow to methane in anaerobic digesters with the microorganisms responsible for each step. Adapted from Gujer and Zehnder, 1983.

end-products (methane and carbon dioxide). The methanogens comprise two physiologically distinct groups of methane-forming bacteria, namely acetoclastic methanogenic bacteria and hydrogen-utilizing methanogenic bacteria. Each of the stages outlined above will now be described in more detail.

2.1 Hydrolytic bacteria

Since methanogenic and acetogenic bacteria are generally incapable of utilizing complex polymeric substrates directly, these materials must first be broken down into soluble monomers before the process of methanogenesis can

proceed. Consequently, hydrolysis is the first essential step in the anaerobic degradation of complex biopolymers. A number of extracellular hydrolytic enzymes that are capable of initiating the attack on these complex substrates may be produced within anaerobic digesters by hydrolytic genera such as *Clostridium*, *Peptococcus*, *Vibrio*, *Micrococcus* and *Bacillus*. These include protease, lipase, cellulase, pectinase, amylase, chitinase etc. and their relative composition and activity will reflect the prevalence of their respective substrates in the digester feed. Being extracellular, these enzymes are able to access large substrate molecules that are incapable of crossing the bacterial cell wall due to their size. Anaerobic digesters contain between 10^8 – 10^9 hydrolytic bacteria per ml comprising both facultative and obligate anaerobes.

In addition to bacteria, protozoa and fungi have been observed in anaerobic digesters; however, on the basis of observed numbers, protozoa are not thought to have an important role in the anaerobic digestion process. The protozoa that have been observed include flagellates belonging to the genera *Trepomonas*, *Tetramitus* and *Trichomonas*, amoebae belonging to genera *Vahlkampfia* and *Hartmanella*, and ciliates belonging to the genera *Metopus*, *Trimyema* and *Saprodinium*.

On account of similarly low cell densities, fungi are also believed to play a relatively minor role within anaerobic digesters and much of the recorded fungal biomass probably enters in the digester feed. However, fungi have been shown to be capable of reproduction within operational digesters and to this extent are taking part in the digestion process by consuming nutrients for growth. Genera of fungi observed include *Phycomycetes*, *Ascomycetes* and fungi imperfecti.

2.2 Acid-forming bacteria

The monomers produced by the hydrolytic bacteria during the first stage of the digestion process are fermented during the second acid-forming stage to produce several intermediate products, namely acetate, propionate, butyrate and hydrogen. Each acidic end-product

contains a carboxylic acid group that was introduced during the fermentation, and hence the microorganisms responsible for these fermentations are collectively called the acid-forming bacteria. There are two groups of acid-forming bacteria, namely the acidogenic bacteria and the acetogenic bacteria.

2.2.1 Acidogenic bacteria

Acidogens provide important substrates for acetogens and methanogens. They metabolize amino acids and sugars to the intermediary products, acetate, hydrogen and carbon dioxide. The acidogenic stage includes many different fermentative genera and species; among them are *Clostridium*, *Bacteroides*, *Ruminococcus*, *Butyribacterium*, *Propionibacterium*, *Eubacterium*, *Lactobacillus*, *Streptococcus*, *Pseudomonas*, *Desulfobacter*, *Micrococcus*, *Bacillus* and *Escherichia*. The facultative members of this group also help protect the oxygen-sensitive methanogens by consuming traces of oxygen that may enter in the feed. Typical cell counts of acidogens in anaerobic digesters range from about 10^6 – 10^8 per ml (Archer and Kirsop, 1990).

2.2.2 Acetogenic bacteria

The main function of acetogenic bacteria in anaerobic digestion is the production of acetate, carbon dioxide and hydrogen, as these are the only substrates that can be metabolized efficiently by the methanogens in the final stage of anaerobic digestion. Two distinct groups of acetogenic bacteria can be distinguished on the basis of their metabolism.

The first group, the obligate hydrogen-producing acetogens (OHPA), also called proton-reducing acetogens, produce acetic acid, carbon dioxide and hydrogen from the major fatty acid intermediates (propionate and butyrate), alcohols and other higher fatty acids (valerate, isovalerate, stearate, palmitate and myristate via β -oxidation). OHPA species are particularly important in the β -oxidation of longer-chain fatty acids arising from lipid hydrolysis and are also involved in the anaerobic degradation of aromatic compounds.

Thermodynamic considerations of the free energy (ΔG) available from fatty acid oxidation

and hydrogen production, predict that the OHPA will be capable of growth only in environments that maintain a low concentration of the metabolic product hydrogen. Such environments will exist where hydrogen-consuming species like methanogens are able to thrive. This mutualistic interaction between OHPA and hydrogen-removing species (e.g. methanogens and sulphate-reducing bacteria) has been termed *syntrophy*, literally 'eating together', and is a critical requirement for efficient methanogenic anaerobic digestion. The syntrophic relationships in a working anaerobic digester are held in a fairly fragile state of equilibrium because even small perturbations may lead to positive feedback of the inhibitory effects. Specifically, methanogens are inhibited by fatty acids (substrates of the OHPA) and the OHPA are inhibited by hydrogen (a substrate of the methanogens); any significant increase in the level of either of these substrates will eventually lead to the inhibition of both groups of bacteria (Fig. 24.3).

Most methanogenic environments maintain a hydrogen concentration below 10^{-4} atm, which is sufficiently low to stimulate the OHPA and prevent the accumulation of fatty acids. So far, only a limited number of OHPA species have been isolated and identified, namely *Syntrophomonas wolfei* and *Syntrophobacter wolnii*, which oxidize butyrate and propionate, respectively. Mesophilic sludges contain approximately 4.5×10^6 *S. wolfei* per gram of digester sludge (Toerien and Hattingh, 1969).

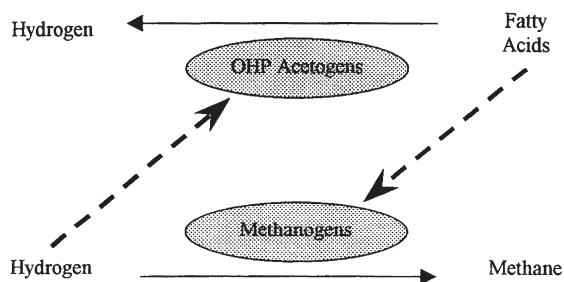


Fig. 24.3 Schematic representation of feedback inhibition (broken lines) among syntrophic bacteria (methanogens) and obligate hydrogen-producing (OHP) acetogens. Substrates and products are shown incompletely.

The second group of acetogenic bacteria are the homoacetogens, which are strictly anaerobic microorganisms catalyzing the formation of acetate from hydrogen and carbon dioxide. Homoacetogens are known in the genera *Acetobacterium*, *Acetoanaerobium*, *Acetogenium*, *Butribacterium*, *Clostridium* and *Pelobacter*. Homoacetogenic bacteria are also syntrophs because they participate in the interspecies hydrogen transfer process which maintains the low hydrogen concentrations required by the OHPA. However, their importance in this respect, relative to that of the methanogens, is still not clear although the number of homoacetogenic bacteria in anaerobic digesters is considerably lower, at around 1×10^5 per ml (Toerien and Hattingh, 1969), than that of the methanogens, suggesting a relatively minor role.

2.3 Methanogenic bacteria

The methanogens are strict anaerobes and form methane gas as the end-product of their metabolism. They are known to be truly distinct from the typical bacteria (*Eubacteria*) and are classified in a separate kingdom, the *Archaea*. A recent taxonomic scheme of methanogens is given in Table 24.2. They are the key organisms in the production of methane from acetate, hydrogen and carbon dioxide. Without methanogens, the ultimate breakdown of an organic material would not take place due to the accumulation of the end-products of the acid-producing bacteria. Importantly, methane production is considered to be the slowest (rate limiting) step in the anaerobic digestion process. Furthermore, since the methanogens are most active in the pH range of 6.5–8.0, they will be sensitive in environments poorly buffered against acidification caused by the products of the acidogenic and acetogenic bacteria. A limited range of substrates can be utilized by methanogens with acetate, hydrogen and carbon dioxide being the most important. According to their substrate specificity, methanogens are divided into two groups: acetoclastic methanogens and hydrogen-utilizing methanogens. The methanogenic

population in anaerobic digesters is typically present at a level of 1×10^6 to 1×10^8 per ml (Toerien and Hattingh, 1969).

2.3.1 Acetoclastic methanogens

Of the various end-products produced by acid-forming bacteria, acetate is regarded as the most important precursor of methane production and the source of up to 70% of methane evolved in digesters. In spite of this fact, only two methanogenic genera contain species that are able to utilize acetate (acetoclastic), and these are *Methanosaeta* (formerly known as *Methanothrix*) and *Methanosarcina*. In addition to this acetoclastic activity, *Methanosarcina* spp. are also capable of using methanol, methylamines and sometimes H_2 and CO_2 as growth substrates, while *Methanosaeta* spp. are restricted to growth only on acetate.

Methanosarcina spp. exhibit a higher maximum specific growth rate (shorter doubling times) on acetate than *Methanosaeta* spp., however, the latter has the higher substrate affinity (lower K_s) for acetate. Consequently, *Methanosaeta* will be the dominant acetoclastic species at acetate concentrations below 1 mM, whereas higher acetate concentrations favour *Methanosarcina* due to its faster growth. With acetate as the growth substrate, doubling times of *Methanosarcina* spp. and *Methanosaeta* spp. are 24 hours and 3.5–9.0 days, respectively. The two genera also exhibit different growth physiology; *Methanosaeta* are filamentous organisms, whereas *Methanosarcina* usually grow in aggregates consisting of large numbers of individual cells, each surrounded by a thick cell wall.

2.3.2 Hydrogen-utilizing methanogenic bacteria

A significant quantity of the methane production within anaerobic digesters, up to 30% of the total, is produced by hydrogen-utilizing methanogenic bacteria. These methanogens reduce carbon dioxide, formate, methanol and methylamines, using the hydrogen produced fermentatively by the hydrolytic and acid-forming bacteria earlier in the digestion

process. In 1906, Soehngen was the first to record the reduction of CO_2 by hydrogen and the subsequent yield of methane.

When utilizing only hydrogen and carbon dioxide, the methanogens grow as chemolithotrophic autotrophs because they derive both their energy and cellular carbon from inorganic chemicals. Interestingly, such energy metabolism does not involve conventional cytochromes for electron transport. Instead, a complex seven-step process has evolved with specific cofactors, such as coenzyme M (CoM), which are unique to the methanogens. CoM is the smallest coenzyme known and is exceptional in its high sulphur content and acidity. The other implication of using only carbon dioxide or other one-carbon (C_1) substrates for growth is the need to generate two-carbon (C_2) building blocks for anabolic processes. Methanogens achieve this in a manner similar to that of the homoacetogenic bacteria (Zeikus *et al.*, 1985). These issues are discussed further in Section 4.1.

3 LABORATORY METHODS FOR ANAEROBIC BACTERIA

3.1 Isolation and cultivation methods

Most anaerobic bacteria are extremely sensitive to oxygen and therefore require stringent anaerobic techniques. In order to maintain an anaerobic environment during handling and incubation periods, oxygen exclusion techniques should be employed. Holland *et al.* (1987) stated that two types of oxygen exclusion techniques can be used in anaerobic microbiology. The first method employs all possible precautions to ensure that oxygen is excluded from every step of the handling procedure, including sampling, media preparation, transfer and incubation. The second technique is a less systematic approach in which only the incubation step is maintained oxygen-free. These workers also noted the advantages and disadvantages of each method. For example, the former is slow, time consuming and demanding on the operator, however, when performed correctly it is the only method that allows successful handling of the strict

anaerobes. The latter is relatively quick and easy to carry out, but has an unacceptably high failure rate with sensitive organisms.

Removal of oxygen by chemical means can be achieved by the addition of oxygen-reactive compounds directly into culture media (e.g. cysteine, sodium sulphides) or into traps within the culture vessel in order to absorb molecular oxygen (e.g. the mixture of pyrogallol acid and sodium hydroxide, the use of iron wool dipped in acidified copper sulphate, the combination of chromium and sulphuric acid, and the burning of yellow phosphorus). However, the use of the latter chemicals can be associated with toxicity and inhibition of anaerobic bacteria and may also represent an unacceptable risk to the worker.

Biological methods for establishing low oxygen concentrations in culture media involve the inoculation of aerobic organisms together with anaerobic bacteria in order to reduce any oxygen to carbon dioxide. However, the aerobic organism should be a non-fermentative strain so as not to interfere with the fermentation reactions of the anaerobic bacteria. Organisms such as *Acinetobacter*, *Pseudomonas aeruginosa* and *Bacillus subtilis* may be suitable for this purpose.

When more sophisticated physical methods of oxygen exclusion are desired, the need for specialized equipment will be inevitable. Typical equipment includes anaerobic jars and anaerobic cabinets that are filled with an inert gas such as nitrogen or helium to maintain an oxygen-free environment. Such equipment has been widely used in the isolation and detailed study of anaerobic microorganisms.

Exclusion of oxygen from culture media by chemical, biological or physical methods has been detailed by several authors (Willis, 1977, 1990; Holland 1987 *et al.*, Levett, 1990). The methodology for culturing strict anaerobic bacteria in anaerobic jars was first developed at the beginning of this century. The standard anaerobic jars can hold 12 petri dishes and contain a sachet holding 1 g deoxo cold catalyst for oxygen removal in the presence of excess hydrogen. Anaerobic conditions develop in the jar when the air is vacuumed out and replaced with a mixture of hydrogen (90%) and carbon

dioxide (10%). It is advisable to put an indicator into the jar in order to verify the establishment of anaerobic conditions; resazurin, methylene blue or biological indicators (either a strict aerobe or a strict anaerobe) are frequently used.

Anaerobic cabinets were first employed in isolation work during the 1960s to eliminate the problems encountered with other devices. An anaerobic cabinet, equipped with glove ports and a rigid air lock for transfer of materials into the working area, provides an oxygen-free environment in which conventional aseptic bacteriological techniques can be applied to the isolation and manipulation of obligate anaerobes under conditions of strict and continuous anaerobiosis (Willis, 1990). They offer the microbiologist a large working area with control over temperature, pressure and gas composition. The advent of the anaerobic cabinet greatly facilitated use of petri dish plate cultures, subculturing of anaerobic bacteria, and culture inspection without the continual risk of oxygen exposure. Prolonged culture incubations, handling of multiple sample replicates and the storage of prereduced media are all feasible with this technique. However, microscopic examinations within an anaerobic cabinet remain problematic.

There are several alternative methods to achieve anaerobic culture conditions in the laboratory. These include the Hungate or roll-tube technique (Holland *et al.*, 1987; Levett, 1990), the serum tube method (Balch *et al.*, 1979) and shake-flask cultures and fluid cultures (Willis, 1977, 1990).

3.2 Examination and identification of anaerobic bacteria

The anaerobic bacteria are a diverse group of microorganisms and examples of the major generic groups are given in Table 24.1. However, in the context of anaerobic digestion processes, our interest is focused on those bacteria associated with the mechanism of methane production. Therefore, this section will be limited to the enrichment, identification, enumeration and taxonomy of only

TABLE 24.1 Types of anaerobic bacteria and their generic names (Holland *et al.*, 1987)

1. Phototrophic bacteria	
A. Purple sulphur bacteria (Family Chromatiaceae):	<i>Amoebobacter</i> , <i>Chromatium</i> , <i>Lamprocystis</i> , <i>Thiocystis</i> , <i>Thiodictyon</i> , <i>Thiopedia</i> , <i>Thiosarcina</i> , <i>Thiospirillum</i> , <i>Ectothiorhodospira</i>
B. Green sulphur bacteria (Family Chlorobiaceae):	<i>Ancalochloris</i> , <i>Chlorobium</i> , <i>Pelodictyon</i> , <i>Prosthecochloris</i> (Family Chloroflexaceae): <i>Chloroflexus</i> , <i>Chloronema</i> , <i>Oscillochloris</i>
2. Spirochaetes: <i>Spirochaeta</i> , <i>Cristispira</i> , <i>Treponema</i> , <i>Borrelia</i>	
3. Anaerobic Gram-negative bacteria: <i>Bacteroides</i> , <i>Fusobacterium</i> , <i>Butyrivibrio</i> , <i>Leptotrichia</i> , <i>Succinivibrio</i> , <i>Succinomonas</i> , <i>Anaerobiospirillum</i> , <i>Wolinella</i> , <i>Anaerovibrio</i> , <i>Pectinatus</i> , <i>Acetovibrio</i> , <i>Selenomonas</i> , <i>Lachnospira</i> , <i>Desulfovibrio</i> and other sulphate-reducers	
4. Methanogenic bacteria: <i>Methanobacterium</i> , <i>Methanobrevibacter</i> , <i>Methanothermus</i> , <i>Methanococcus</i> , <i>Methanomicrobium</i> , <i>Methanogenium</i> , <i>Methanospirillum</i> , <i>Methanosarcina</i> , <i>Methanosaeta</i> , <i>Methanoplanus</i>	
5. Anaerobic cocci	
A. Gram-positive:	<i>Streptococcus</i> , <i>Peptococcus</i> , <i>Peptostreptococcus</i> , <i>Gaffkya</i> , <i>Ruminococcus</i> , <i>Sarcina</i> , <i>Coprococcus</i>
B. Gram-negative:	<i>Veillonella</i> , <i>Megasphaera</i> , <i>Gemmiger</i> , <i>Acidaminococcus</i>
6. Anaerobic Gram-positive non-sporing bacteria: <i>Bifidobacterium</i> , <i>Eubacterium</i> , some <i>Lactobacillus</i> , some <i>Actinomyces</i> , <i>Corynebacterium</i> (<i>Bacterionema</i>), <i>Propionibacterium</i>	
7. Anaerobic endospore-forming bacilli: <i>Clostridium</i> , <i>Desulfotomaculum</i>	
8. Cell-wall deficient anaerobes: <i>Anaeroplasm</i> and L-forms	
9. Microaerophilic bacteria: <i>Beggiatoa</i> , <i>Campylobacter</i> , <i>Rhodopseudomonas</i> , <i>Spirillum</i>	

the methanogenic bacteria. For a comprehensive treatise on the other anaerobic bacteria, the reader is referred to Holland *et al.* (1987).

3.3 Taxonomy and identification of methanogens

With increasing knowledge of the physiology and structure of methanogens and with the advent of newer techniques for establishing the interrelationship between bacteria, it has been confirmed that the methanogens are a coherent group. Indeed, they are now known to be truly distinct from the typical bacteria (*Eubacteria*) and are classified in a separate kingdom, the *Archaea* (Kirsop, 1984). The phylogenetic evidence for the taxonomic distinction between *Archaea* and *Eubacteria* was derived from 16S ribosomal (rRNA) sequence characterization two decades ago. In addition to this phylogenetic difference, there are also a number of phenotypic differences between them (Kirsop, 1984; Levett, 1990):

1. Eubacterial cell walls contain muramic acid, whereas those of Archaeobacteria do not
2. the occurrence of unique transfer RNA and 5S ribosomal RNA in the translation mechanisms of protein synthesis

3. membrane lipids in Eubacteria are glycerol esters of fatty acids, whereas Archaeobacterial lipids are diethers of glycerol and isoprenoids
4. Archaeobacteria possess tRNA devoid of ribothymidine in the T ψ C loop
5. Archaeobacterial RNA polymerases have distinct subunit structures.

As the basis for any taxonomic work, an identification method should be employed to distinguish the species of methanogenic bacteria. In the early studies, cell shape was used as a primary determinant for taxonomic assignment of methanogenic genera, while physiological and nutritional properties were used to form the basis of species designation. A recent attempt by Kasapgil (1994) to organize the taxonomy of methanogens using cell morphology and nutritional capability is tabulated in Table 24.2. However, an adequate taxonomic classification of methanogens could not be established using only these parameters. A more comprehensive elucidation of the taxonomy of the methanogens was provided using comparative cataloging of 16S rRNA (Balch *et al.*, 1979). In this technique RNA is extracted from the cell and the 16S rRNA is purified. This material is digested by ribonuclease T₂ into oligonucleotides that can then be separated

TABLE 24.2 Methanogenic classification (Kasapgil, 1994)

Order	Family	Genus	Species	Gram reaction	Morphology	Substrate
Methanobacteriales	Methanobacteriaceae	<i>Methanobacterium</i>	<i>M. formicicum</i>	+	Long rods, filaments	H ₂ , CO ₂ , formate
			<i>M. bryanti</i>	+	Short long rods	H ₂ , CO ₂ , formate
			<i>M. thermoautotrophicum</i>	+	Long rods, filaments	H ₂ , CO ₂ , formate
			<i>M. wolfei</i>	+	Rods	H ₂ , CO ₂
			<i>M. alcaliphilum</i>	+	Rods	H ₂ , CO ₂
			<i>M. uliginosum</i>	–	Rods	H ₂ , CO ₂
			<i>M. thermoformicum</i>	+	Rods	H ₂ , CO ₂ , formate
		<i>Methanobrevibacter</i>	<i>M. arbophilus</i>	+	Short rods	H ₂ , CO ₂ , formate
			<i>M. ruminantium</i>	+	and short	H ₂ , CO ₂ , formate
			<i>M. smithii</i>	+	chains	H ₂ , CO ₂ , formate
	Methanothermaceae	<i>Methanothermus</i>	<i>M. fervidus</i>		Short rods	H ₂ , CO ₂
			<i>M. sociabilis</i>		Rods	H ₂ , CO ₂
Methanococcales	Methanococcaceae	<i>Methanococcus</i>	<i>M. vannielli</i>	–	Irregular	H ₂ , CO ₂ , formate
			<i>M. voltae</i>	–	cocci	H ₂ , CO ₂ , formate
			<i>M. maripaludis</i>	–	single or pairs	H ₂ , CO ₂ , formate
			<i>M. thermolithotrophicus</i>	–		H ₂ , CO ₂ , formate
			<i>M. halophilus</i>	–		Methanol, methylamines
			<i>M. jannaschi</i>	–	Irregular cocci	H ₂ , CO ₂ , formate
			<i>M. deltae</i>	–		H ₂ , CO ₂ , formate
			<i>M. frisisus</i>	–	Irregular cocci	H ₂ , CO ₂
			<i>M. mobile</i>	–	Short rods single	H ₂ , CO ₂ , formate
			<i>M. paynter</i>	–	Short rods single	H ₂ , CO ₂

(continued on next page)

TABLE 24.2 (continued)

Order	Family	Genus	Species	Gram reaction	Morphology	Substrate
		<i>Methanogenium</i>	<i>M. cariaci</i>	–	Irregular	H ₂ , CO ₂ , formate
			<i>M. marisnigri</i>	–	cocci, single or	H ₂ , CO ₂ , formate
			<i>M. olentangyi</i>		pairs	H ₂ , CO ₂ , formate
			<i>M. thermophilicum</i>	+	Irregular cocci	H ₂ , CO ₂
			<i>M. aggregans</i>	+	Irregular cocci	H ₂ , CO ₂ , formate
			<i>M. bourgense</i>	+	Irregular cocci	H ₂ , CO ₂ , formate
			<i>M. tationis</i>		Irregular cocci	H ₂ , CO ₂ , formate
		<i>Methanospirillum</i>	<i>M. hungatei</i>	–	Spirillum, regular rods and filaments	H ₂ , CO ₂ , formate
	Methanoplanaceae	<i>Methanoplanus</i>			Plated shape	H ₂ , CO ₂ , formate
	Methanosarcinaceae	<i>Methanosarcina</i>	<i>M. limicola</i>	–	Pseudosarcina	H ₂ , CO ₂ , formate
			<i>M. barkeri</i>	+	Irregular cocci in	
			<i>M. mazei</i>	+	large aggregates	
			<i>M. thermophila</i>		Pseudosarcina	Acetate
			<i>M. acetivorans</i>	+	Pseudosarcina, coccoid	Methylamines
			<i>M. vacuolate</i>		Pseudosarcina	
		<i>Methanococcoides</i>	<i>M. methylutents</i>	+	Irregular cocci	Methanol, methylamines
		<i>Methanotherix</i>	<i>M. soehngenii</i>	+	Irregular cocci sheath forming long filament	Acetate
			<i>M. concilli</i>		Sheathed rod	Acetate
		<i>Methanolobus</i>	<i>M. tindarius</i>	+	Irregular cocci single or loose	Methanol, methylamines

TABLE 24.3 Taxonomic treatment of methanogens by Balch *et al.* (1979) based on 16S rRNA comparative cataloguing

Order	Family	Genus	Species
Methanobacteriales	Methanobacteriaceae	<i>Methanobacterium</i>	<i>M. formicum</i> <i>M. bryantii</i> <i>M. thermoautotrophicum</i>
		<i>Methanobrevibacter</i>	<i>M. ruminantium</i> <i>M. orbariphilus</i> <i>M. smithii</i>
Methanococcales	Methanococcaceae	<i>Methanococcus</i>	<i>M. vanmielii</i> <i>M. voltae</i>
Methanomicrobiales	Methanomicrobiaceae	<i>Methanomicrobium</i>	<i>M. mobile</i>
		<i>Methanogenium</i>	<i>M. cariaci</i> <i>M. marisnigri</i>
	Methanosarcinaceae	<i>Methanospirillum</i> <i>Methanosarcina</i>	<i>M. hungatei</i> <i>M. barkeri</i>

and sequenced. Sequences of oligonucleotides with six or more units are compared between pairs of bacteria to show their identity (Taylor, 1982). Using this approach, a new taxonomic treatment for the methanogenic bacteria (Table 24.3) based on 16S rRNA comparative cataloguing was proposed (Balch *et al.*, 1979). Other techniques that have been used to identify and classify the taxonomy of methanogens include:

1. morphology and staining
2. cell wall structure and composition
3. colonial appearance
4. growth inhibition and stimulation
5. gas liquid chromatography
6. distribution and molecular weight of polar lipids
7. antigenic relationships
8. polyamine content.

For more information on the above techniques, the reader is referred to Willis (1977); Balch *et al.* (1979); Whitman *et al.* (1992).

3.4 Enrichment

Enrichment culture is the technique that is used to enhance the population density of a particular group of microorganisms within the total microbial population of a sample. This is achieved by preferentially stimulating the growth of the target group of microorganisms

by judicious manipulation of the physiological conditions during the enrichment phase. Methanogenic enrichments are usually performed in media that has a nutrient composition, environmental pH value, temperature and oxygen-free conditions, similar to those of natural methanogenic environments.

Confirmation of methanogens in the enrichment can be achieved by microscopy or gas chromatographic analysis of the headspace gas for methane. There are a number of specialized media used in anaerobic enrichment. Some allow greater initial selectivity for the methanogens against the background populations and are best suited to isolation work, while others have been developed to optimize growth of the isolated strains and are used for routine culture maintenance. Balch *et al.* (1979) described a standard medium suitable for the enrichment of methanogenic bacteria.

Early work on the isolation of pure cultures and the selection of anaerobic and methanogenic bacteria was reported by Toerien and Hattingh (1969). Further details on the isolation and selection of specific anaerobic bacteria are given by Holland *et al.* (1987).

As an example of the nutritional requirements of methanogens, Table 24.4 lists an anaerobic growth medium based on that reported by Balch *et al.* (1979). For more information on other media preparations, the reader is referred to the reports of Willis (1977); Kirsop (1984); Whitman *et al.* (1992); Levett (1990).

TABLE 24.4 Composition of an enrichment media (Balch *et al.*, 1979)

Sodium formate	1200 mg
Sodium acetate	1200 mg
Yeast extract	3000 mg
Sodium carbonate (8% w/v)	15 ml
Resazurin (0.01% w/v)	0.6 ml
Ferrous sulphate (1% w/v)	0.6 ml
Mineral solution 1	15 ml
Mineral solution 2	15 ml
Trace minerals	6 ml
Vitamins	6 ml
Reducing solution	9 ml

Mineral solution 1: Dipotassium hydrogen orthophosphate (0.6 (w/v)); *Mineral solution 2:* Potassium dihydrogen orthophosphate (600 mg), Ammonium sulphate (12000 mg), Sodium chloride (12000 mg), Calcium chloride (2400 mg), Magnesium sulphate (2500 mg); *Trace mineral solution:* Nitrilotricetic acid (1500 mg), Magnesium sulphate (3000 mg), Manganese sulphate (500 mg), Sodium chloride (1000 mg), Ferrous sulphate (100 mg), Cobalt sulphate (100 mg), Calcium chloride (100 mg), Zinc sulphate (100 mg), Copper sulphate (10 mg), Aluminium potassium sulphate (10 mg), Boric acid (10 mg), Sodium molybdate (10 mg), Nickel chloride (10 mg), Sodium selenate (10 mg); *Vitamins:* D-Biotin (2 mg), Folic acid (2 mg), Pyridoxine hydrochloride (10 mg), Thiamine hydrochloride (5 mg), Riboflavin (5 mg), Nicotinic acid (5 mg), D-Pantothenic (5 mg), Vitamin B₁₂ (0.1 mg), p-Aminobenzoic acid (5 mg), DL-6 8 Thiotic acid (5 mg); *Reducing solution:* Cysteine hydrochloride/sodium sulphide (2.5%)*.

* Dissolve 2.5 g cysteine hydrochloride in 40 ml of distilled water. Adjust pH of the solution to 10 using NaOH pellets. Dissolve 2.5 g sodium sulphide in 40 ml of distilled water. Mix the solutions and make up to 100 ml. Heat and cool under O₂ free nitrogen gas. Dispense into Hungate tubes. Store at 4°C until required.

3.5 Bacterial enumeration

The treatment capacity in any anaerobic digester is primarily determined by the concentration of active biomass retained within the system. Maintaining a high biomass concentration in full-scale anaerobic digesters makes anaerobic treatment more stable and the improved performance also makes the process more attractive economically (Morgan *et al.*, 1991). Therefore, it is important to monitor anaerobic reactors from a microbiological point of view in order to understand the operational conditions that promote high levels of active biomass within the system.

Supervision of full-scale and laboratory-scale anaerobic reactors can be greatly assisted

by bacterial enumeration of the digester sludge, as this provides an indication as to whether or not:

1. the reactor is overloaded by an excessive feed rate, and/or
2. there are any inhibitory materials in the system (and the significance and magnitude of their effect).

Since the enumeration of anaerobic bacteria requires strict anaerobic conditions, it will be necessary to use some of the techniques mentioned above to exclude oxygen while handling anaerobic bacteria.

Enumeration of anaerobic bacteria can be achieved by the agar plate technique, most probable number (MPN) methods and direct counts by microscopy (utilizing the fluorescent properties of oxidized factor F₄₂₀ and specific staining techniques to show particular groups of bacteria). The basic procedures for the direct count and MPN methods are described in more detail below.

3.5.1 Microscopic method (direct count)

Enumeration of the total bacteria and the total fluorescent methanogenic bacteria can be made by using an Epifluorescent microscope (e.g. Zeiss D-7082) with illumination by either a halogen/ultraviolet lamp at 376 nm or with a 50 W high pressure mercury lamp (Kasapgil, 1994). Methanogenic and non-methanogenic bacteria can be differentiated on the basis that only methanogenic bacteria will be seen to emit blue-green fluorescence when the ultraviolet lamp is used.

The following protocol may be used to enumerate methanogens and the acidogenic bacteria. In a sample (Ince *et al.* 1997):

1. Dilute and homogenize the sample to give a count of between 50 and 400 bacteria per field of view
2. Place the sample onto a Neubauer Counting Chamber (graduated and calibrated glass slide)
3. Microscopically count several chambers using (i) visible light illumination to record the total bacteria, and (ii) visible plus

ultraviolet light illumination to record the number of fluorescent cells (methanogenic bacteria)

4. calculate the number of bacteria per ml of sample with the following formula:

$$\text{Count of bacteria (ml}^{-1}\text{)} = Y \cdot D / V$$

where:

Y = mean count per chamber,

D = dilution factor giving counts between 50 and 400 per field of view and

V = volume of a chamber (ml).

Estimate the number of acidogenic bacteria by difference:

Acidogenic bacteria = Total bacteria – Methanogenic bacteria.

3.5.2 Multiple tube method

The most probable number (MPN) technique can be used to estimate the number of viable methanogenic bacteria by serial dilution of the sample and inoculation into culture media containing a specific carbon source. This method was described by Siebert and Hattingh (1967, 1987) and used by Morgan *et al.* (1991) and Kasapgil (1994). The most probable number of the methanogens can be calculated by using the probability tables described by Greenberg *et al.* (1985). For the MPN enumeration, the media described by Balch *et al.* (1979) can be used. The media and samples are placed into Hungate tubes for incubation. After inoculation, the head space of each tube should be replaced with an oxygen-free mixture of hydrogen and carbon dioxide (4:1) and tubes should be incubated statically for 4–6 weeks at 37°C. Positive tubes can be identified by the presence of methane gas in the head-space using gas chromatography and each should be confirmed by epifluorescence microscopy.

3.5.3 Nucleic acid hybridization techniques

Recently, the molecular methods of rRNA analysis, widely used for identifying phylogenetic relationships between microorganisms, have been developed into a number of

emerging techniques. These are capable of both quantifying and visualizing individual groups of microorganisms within complex communities, such as anaerobic granular sludge (Harmsen *et al.*, 1996). The two main techniques both use oligonucleotide probes (or primers) that are specific for conserved regions of rRNA in their target organisms. The probes can be tailored to detect individual species, genera or wider groups of phylogenetically-related microorganisms. Quantification involves extracted rRNA from the sample material being hybridized on a membrane with a series of radio-labelled probes for each target organism (blot technique). The intensity of the radio-label signal is proportional to the concentration of the target group and results can therefore be used to quantify the dynamics of the microbial composition in a sample.

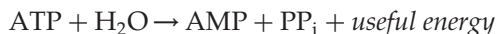
A powerful variation of this technique is fluorescent *in situ* hybridization (FISH) in which fluorescently-labelled probes, similar to those above, are made to react directly with intact microbial cells in the sludge sample (Raskin *et al.*, 1994). This allows fluorescently-stained cells to be visualized microscopically showing the numbers and location of the target cells and their proximity to other cells in the sample. The latter is proving to be a powerful tool in determining the syntrophic relationships between community members in anaerobic sludge granules (Rocheleau *et al.*, 1999).

4 BIOCHEMISTRY OF ANAEROBIC DIGESTION

4.1 Energy conservation

All microorganisms are endowed with a metabolic system that transforms the chemical energy of their feed substrate into biologically useful energy, which is then utilized to perform work for the cell (Thauer *et al.*, 1977). The energy taken up by cells is first used to drive the endergonic synthesis of ATP (adenosine triphosphate) from ADP (adenosine diphosphate) and P_i (inorganic orthophosphate). This ATP

then drives anabolic metabolism and other cellular processes (ion transport, motility, etc.).



In anaerobic cells, ATP is produced through the same two metabolic routes that occur in aerobic metabolism. The first of these is substrate-level phosphorylation, in which the chemical bond energy is transferred directly to ADP from key intermediates in the fermentation pathway, with concomitant ATP generation. Anaerobic fermentation is equivalent to the process of glycolysis in aerobic metabolism, with the exception that fermentation fails to bring about any net oxidation of the substrate during the formation of fermentation products. The second metabolic route is electron transport phosphorylation (ETP), in which electrons flow down an electrochemical gradient between redox carriers (respiratory chain) and the energy in this redox potential drives the translocation of protons (or sodium) across the cell membrane, creating a chemiosmotic proton imbalance. The resulting proton motive force (PMF) is coupled to the phosphorylation of ADP as protons channel back into the cell through a membrane-integrated proton-translocating ATP-ase (Thauer *et al.*, 1977; Hawkes, 1980; Madigan *et al.*, 1997).

Despite the enormous range of electron donors (e.g. H_2 , NADH, NH_3 , Fe^{2+} , $\text{S}_2\text{O}_3^{2-}$) and electron acceptors (CO_2 , NO_3^- , SO_4^{2-} , Fe^{3+} , O_2) that may donate and accept reducing equivalents (electrons) to the respiratory chains of both aerobic and anaerobic metabolism, many of the membrane redox carriers appear to be ubiquitous (e.g. cytochromes). In contrast, major differences exist in the enzymes and cofactors that are involved in the first and last steps of electron transfer both to and from the respiratory chain. Some examples of these are discussed below for the organisms involved in the final steps of methanogenic anaerobic digestion.

The maximum free energy that is available to the chemolithotrophic autotrophs of anaerobic digestion (i.e. sulphate-reducing bacteria (SRB), methanogens and acetogens) can be deter-

mined from thermodynamic consideration of the electron donors and acceptors according to the Gibbs free energy relationship:

$$\Delta G^{0'} = -nF\Delta E^{0'}$$

where:

$\Delta G^{0'}$ is the free energy change under standard conditions, i.e. 25°C and a pressure of 1 atm, at pH 7;

n is the number of electrons transferred;

F is the Faraday constant (96.48 kJ/V);

$E^{0'}$ is the electrochemical oxidation potential;

$\Delta E^{0'}$ is $E^{0'}$ (electron accepting couple) minus $E^{0'}$ (electron donating couple).

Negative values of $\Delta G^{0'}$ indicate an exergonic reaction, i.e. energy is available for the synthesis of ATP. Free energy values for the stoichiometric equations that represent some of the common substrates and intermediates in anaerobic digestion are given in Table 24.5.

The energy available from such reactions is further influenced by the actual concentrations of reactants and products, reactions becoming more energetically favourable when product concentrations are maintained at low levels (see Madigan *et al.*, 1997). Consequently, during anaerobic digestion, the energy yield from methane production and acetogenesis will be considerably lower than the quoted values (−135.6 and −104.6 kJ/mol respectively) because the concentration of hydrogen is usually extremely low in most anaerobic digesters.

4.2 Growth substrates

In practice, most wastes that are treated by anaerobic digestion seldom contain the compounds shown in Table 24.5 in their raw state. More typically, they comprise complex or polymeric organic materials such as lipids, cellulose and protein. In order for these to be converted to the immediate substrates of the methanogens (hydrogen, acetate and CO_2), an

TABLE 24.5 Estimated free energy changes of some biological reactions under standard conditions, i.e. 25°C and a pressure of 1 atm

<i>Dehydrogenations i.e. electron donating, oxidation, reactions:</i>	$\Delta G^{0'}$ (J/mol)
Butyrate \rightarrow Acetate $\text{CH}_3\text{CH}_2\text{CH}_2\text{COO}^- + 2\text{H}_2\text{O} \rightarrow 2\text{CH}_3\text{COO}^- + 2\text{H}_2 + \text{H}^+$	+48.1
Lactate \rightarrow Acetate $\text{CH}_3\text{CHOHCOO}^- + 2\text{H}_2\text{O} \rightarrow \text{CH}_3\text{COO}^- + \text{HCO}_3^- + 2\text{H}_2 + \text{H}^+$	-4.2
Propionate \rightarrow Acetate $\text{CH}_3\text{CH}_2\text{COO}^- + 3\text{H}_2\text{O} \rightarrow \text{CH}_3\text{COO}^- + \text{HCO}_3^- + 3\text{H}_2 + \text{H}^+$	+76.1
Ethanol \rightarrow Acetate $\text{CH}_3\text{CH}_2\text{OH} + \text{H}_2\text{O} \rightarrow \text{CH}_3\text{COO}^- + 2\text{H}_2 + \text{H}^+$	+9.6
Formate \rightarrow HCO_3^- $\text{HCOOH} + \text{H}_2\text{O} \rightarrow \text{HCO}_3^- + \text{H}_2$	+1.3
Acetate \rightarrow Methane $\text{CH}_3\text{COO}^- + \text{H}_2\text{O} \rightarrow \text{HCO}_3^- + \text{CH}_4$	-31.0
<i>Hydrogenations, i.e. electron accepting reactions:</i>	
Methanol \rightarrow Methane $\text{CH}_3\text{OH} + \text{H}_2 \rightarrow \text{CH}_4 + \text{H}_2\text{O}$	-112.5
Formate \rightarrow Methane $\text{HCOOH} + 3\text{H}_2 + \text{H}^+ \rightarrow \text{CH}_4 + 3\text{H}_2\text{O}$	-134.3
$\text{HCO}_3^- \rightarrow$ Methane $\text{HCO}_3^- + 4\text{H}_2 + \text{H}^+ \rightarrow \text{CH}_4 + 3\text{H}_2\text{O}$	-135.6
$\text{HCO}_3^- \rightarrow$ Acetate $2\text{HCO}_3^- + 4\text{H}_2 + \text{H}^+ \rightarrow \text{CH}_3\text{COO}^- + 4\text{H}_2\text{O}$	-104.6

initial partial degradation is carried out by non-methanogenic populations (see Table 24.1) and involves the following major steps.

4.2.1 Hydrolytic degradation

Hydrolases cleave their substrate by insertion of a water molecule. They do not require stoichiometric quantities of cofactors or an energy supply and must be able to function extracellularly as their substrates are too large to cross the cell membrane. Examples include protease, which hydrolyses proteins to peptides and amino acids, amylase, cellulase, pectinase and xylanase, which hydrolyse their respective carbohydrate substrates to dextrins and hexose sugars and lipase which hydrolytically releases the glycerol and fatty acids from the parent lipid.

4.2.2 Fermentation pathways

The products of hydrolysis, along with other low molecular weight compounds present in the effluent, are further degraded intracellularly by a number of different fermentation pathways. These involve enzymic steps that may require cofactors (e.g. dehydrogenases) or energy (kinases) to be supplied, and unless reducing equivalents are consumed by a respiratory chain oxidation process, or converted to molecular hydrogen (H_2) by a hydrogenase, the fermentation product will have the same oxidation state as the substrate. Typical fermentation products from sugars and amino acids are acetate, propionate, butyrate and formate. Long chain fatty acids are degraded by β -oxidation, a process that is coupled to hydrogen formation as means of recycling the adenine nucleotide cofactors (NAD and FAD).

4.2.3 Hydrogen production

It is clear from the above considerations that hydrogen is an important intermediate in anaerobic digestion, being produced during the degradation of all major classes of complex substrate. During anaerobic digestion, approximately 30% of all methane production is generated from the reduction of carbon dioxide by hydrogen. In addition, the concentration of hydrogen also affects the pH value in anaerobic reactors. A rise in the level of H_2 leads to a lowering of pH in the digester. Low pH can selectively inhibit the methanogenic bacteria and this in turn may result in a further and catastrophic acidification of the digester as volatile acids accumulate (Archer and Kirsop, 1990).

Hydrogen may also affect the substrate conversion potential of many major anaerobic bacteria. A build-up of hydrogen can inhibit the growth of hydrogen-producing organisms and alter electron flow, resulting in the formation of more reduced products such as lactate, butyrate or ethanol instead of acetate. Inhibition of hydrogen-producing bacteria occurs because hydrogen inhibits the hydrogenase of these bacteria (Oremland, 1988), preventing them from recycling their reduced cofactors.

Briefly, hydrogen imparts two major effects on anaerobic degradation. The first involves the selection process that determines the chemical pathway that is used to degrade organics, and the second is associated with inhibition of syntrophic bacteria responsible for C_3 and C_4 acid oxidation (Harper and Pohland, 1986).

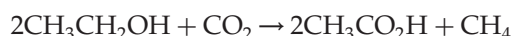
Partial pressure of hydrogen (1 ppm (v/v) H_2 produces 10^{-6} atm pressure at pH 7) has a vital role in methane formation and substrate flow. In other words, thermodynamically, the conversion of VFA to methane depends on the partial pressure of H_2 . For example, if the reactions for ethanol, butyrate, propionate are to be thermodynamically favourable methanogenic substrates, the partial pressure of H_2 must be maintained below 10^{-1} , 10^{-3} and 10^{-4} atm, respectively. On the other hand, for H_2 to be converted to methane, a concentration of H_2 greater than 10^{-6} atm is required. Therefore, there is a limited range of hydrogen concentrations that satisfies both these requirements and these are shown in Fig. 24.4.

The partial pressure of H_2 is regulated not only by hydrogen-producing bacteria but also by hydrogen-consuming bacteria, e.g. homo-acetogens and methanogens (see Interspecies hydrogen transfer).

Hydrogen level in anaerobic digesters can be used for process control and in the diagnosis of process problems. It can be measured by several methods, such as gas chromatography, spectrometry (Archer and Kirsop, 1990) and polarographic methods, such as the exhaled hydrogen monitor (Collins and Paskins, 1987).

4.3 Interspecies hydrogen transfer

The discovery that *Methanobacterium omelianskii* really consisted of a syntrophic association of two different bacteria helped to reveal the subtle role of H_2 as an intermediate. Originally *M. omelianskii* was believed to be a methanogen that catabolized ethanol according to:



Later this was shown not to be the case and resulted from a syntrophic association between an acetogen and a methanogen. The acetogen converts ethanol to acetate and hydrogen while the methanogen uses the hydrogen to reduce CO_2 :

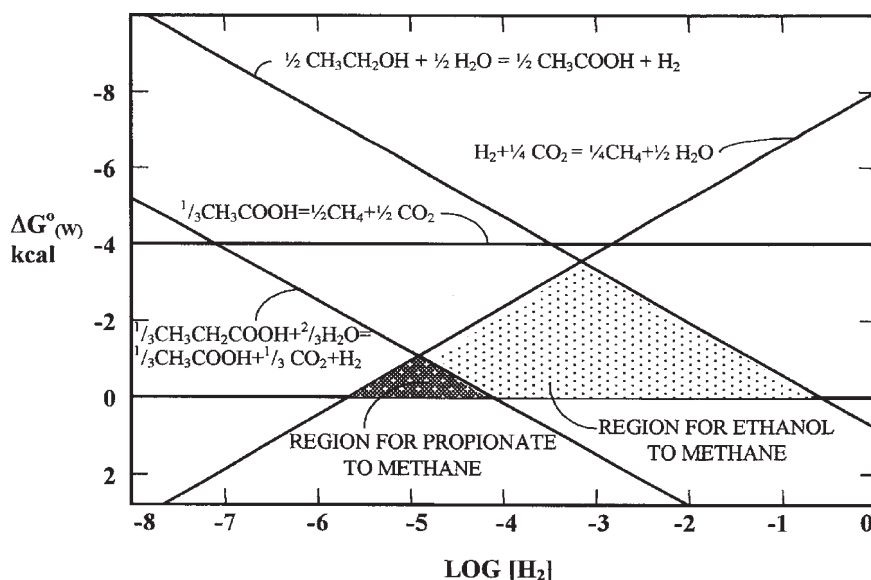
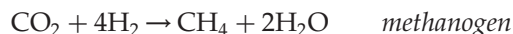
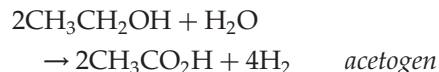


Fig. 24.4 Effect of hydrogen partial pressure on the free energy changes ($\Delta G^{\circ}_{(w)}$) (McCarty, 1981, with permission).

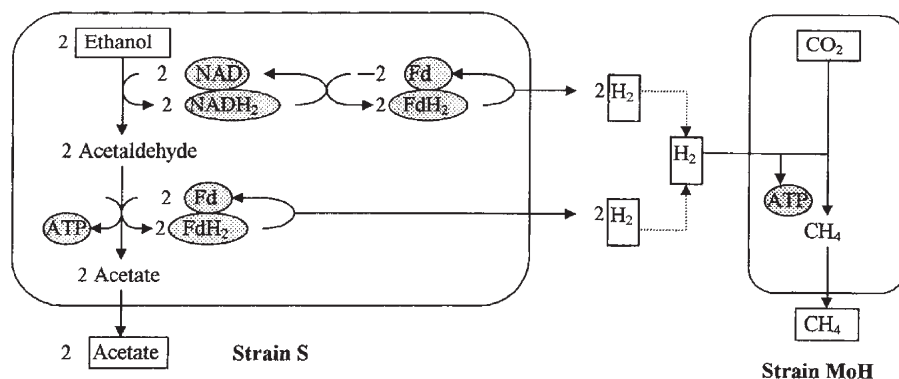


Fig. 24.5 Inter-species hydrogen transfer between two bacteria, Strain S and Strain MoH (Schlegel, 1993, with permission).

It has now been established that other long chain fatty acids are also catabolized by similar associations (Klass, 1984).

In anaerobic digesters, the above mechanism of 'interspecies hydrogen transfer' is usually an obligatory syntrophic association between the anaerobic bacteria forming hydrogen and those converting it to methane. Each member benefits from, or is dependent on, the activity of the other. This is shown schematically in Fig. 24.5.

4.4 Methanogenesis

The final stage in anaerobic digestion is methanogenesis. The production of methane, the most fully reduced form of carbon, is the terminal product since it can undergo no further degradation anaerobically. Methane production brings two major economic benefits to the anaerobic digestion process. First, it is a valuable resource in its own right having a high calorific content, and second, its dissipation from the treatment system prevents a major part of carbon load in the wastewater from being incorporated anabolically into biomass, ensuring low sludge production rates.

The methanogens have evolved as a specialized group of *Archaea* that utilize the CO₂, hydrogen and acetate generated as end-products by the fermentative bacteria. This capability depends on several enzymes and coenzymes found in no other living organisms. By definition coenzymes exist in cells in catalytically small amounts (Hawkes, 1980), however, because of their importance in the energy-yield-

ing pathway and relatively low energy yield, many of the coenzymes are present at high levels in methanogens.

4.4.1 Carrier coenzymes of methanogenesis

The pathway of methane production from carbon dioxide and hydrogen is an intriguing process. First, synthesis of methane appears to offer little energetic reward and second it is intimately linked to the generation of acetate within the cell, the starting intermediate for anabolic pathways in methanogens, SRBs and homoacetogens. The general scheme of these two processes is described below with reference to the key biochemical carriers.

C₁-carriers. During methanogenesis, carbon dioxide is subjected to a series of reductive steps in which hydrogen is the ultimate supplier of reducing electrons, however, the exact nature of many of the immediate electron carriers involved in these steps is still unknown. Several specific coenzymes (carriers) have been elucidated which function to stabilize the carbon atom of carbon dioxide as it is successively reduced to formyl-carbon, methylene-carbon, and methyl-carbon before being finally released as methane.

Methanofuran. Initially, carbon dioxide binds to methanofuran (Fig. 24.6), is reduced to the formyl-level and is then transferred to methanopterin, the second carrier in the pathway. Methanofuran is a low-molecular weight coen-

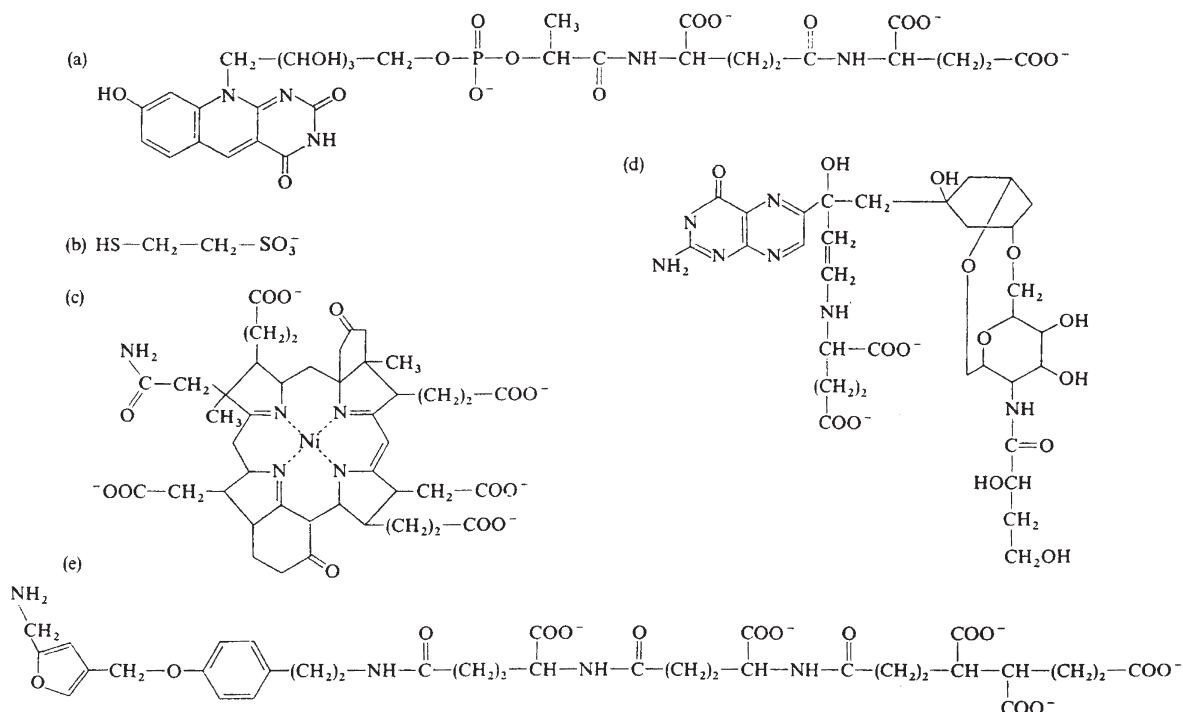


Fig. 24.6 Unique coenzymes of the methanogenic pathway. (a) Factor 420; (b) coenzyme M; (c) Factor 430; (d) methanopterin; (e) methanofuran (Stanier *et al.*, 1986).

zyme which consists of phenol, two glutamic acid molecules, an unusual long chain dicarboxylic fatty acid, and a furan ring.

Methanopterin. Methanopterin then carries the C₁-atom through the next two steps. First, during reduction to the methylene-level by reducing electrons supplied from coenzyme F₄₂₀ and second as it is further reduced to the methyl-level. At this point the methanopterin-bound methyl group may enter the acetyl-CoA biosynthetic pathway (see below) or continue along the methanogenic pathway. The latter involves its transfer to the final carrier, coenzyme M (CoM).

Coenzyme M (CoM). Coenzyme M (CoM) is the simple molecule, 2-mercaptoethanesulphonic acid (HS-CH₂.CH₂.SO₃⁻). During the final step, the CoM-bound methyl group is reduced to free methane in a complex reaction involving the enzyme methyl reductase. Apart from releasing the C₁-unit as a free form (CH₄)

once again, this final step is also of note because it is the position of energy conservation in the methanogenic pathway, creating a PMF through the proton translocation that occurs concomitantly with reduction of the methyl group.

CoM has been detected in all methanogenic bacteria at an internal concentration of 0.3–50 nmol/mg protein. It is also used as a growth factor (Stanier *et al.*, 1986). The overall scheme in which the C₂ of carbon dioxide is successively reduced to methane is shown schematically in Fig. 24.7.

4.4.2 Other cofactors of methanogenesis

Methyl-CoM and factor₄₃₀ (F₄₃₀). Methyl-CoM (CH₃-S-CH₂.CH₂.SO₃⁻) is probably the central intermediate in methane formation from CO₂, formate, methanol, methylamines, and acetate. In cell extracts of methanogenic bacteria

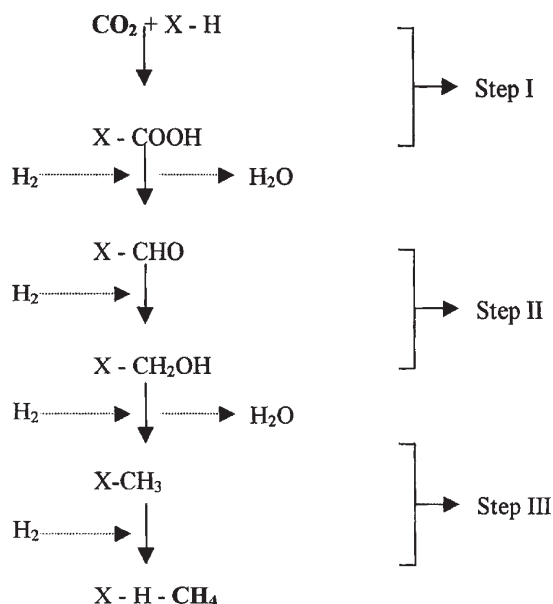
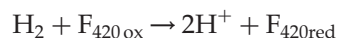


Fig. 24.7 The pathway of methane production from CO_2 . X represents the various cofactor carriers that bind the C_1 unit during each successive reduction step. (See text for details.)

methyl-CoM is reduced to methane and CoM. The methyl reductase catalysing this reaction has been purified and shown to contain the cofactor F_{430} , which is a yellow, non-fluorescent compound with an absorption maximum at 430 nm.

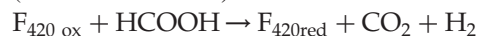
Coenzyme F_{420} . Coenzyme F_{420} is an electron carrier and has a blue-green fluorescence with an absorption peak at 420 nm. The fluorescence ability of F_{420} can be used to identify and enumerate methanogenic bacteria. F_{420} interacts with a number of different enzymes in methanogens, including hydrogenase, formate dehydrogenase, NADP reductase, carbon monoxide dehydrogenase, and pyruvate dehydrogenase.



(hydrogenase)



(NADP reductase)



(formate dehydrogenase)

Coenzyme HS-HTP. Coenzyme HS-HTP is a phosphorylated derivative of the amino acid threonine containing a fatty acid side chain with a terminal SH group. It is the electron donor involved in the last step of CO_2 conversion to CH_4 :



CoM and HS-HTP are regenerated by reduction with H_2 :



4.5 Biosynthesis of acetate

The pathway of methane formation in methanogens is intimately linked to their biosynthetic pathway for acetate synthesis, as all chemolithotrophic autotrophs must produce this as the starting intermediate for their anabolic pathways. Specifically, the methanopterin-bound methyl group in the methanogenic pathway (see above) may be transferred to a C_1 -carrier (a corrinoid protein) which is part of the acetyl-CoA (Ljungdahl-Wood) pathway. This methyl group is then attached to a second C_1 -unit, also originating from carbon dioxide, in a complex series of reactions involving the enzyme carbon monoxide dehydrogenase (Madigan *et al.*, 1997). The C_2 -acetyl group formed by the fusion of these two C_1 -fragments is the precursor of all subsequent cellular biosynthesis.

4.6 Alternative electron acceptors

4.6.1 Sulphate and nitrate

Although methanogenesis is the major route for hydrogen oxidation in anaerobic digesters, the terminal electron acceptor of this process, carbon dioxide, is not the only compound likely to exist in such environments with the capacity to support chemolithotrophic metabolism. The most common alternative electron acceptors are sulphate and nitrate, terminal electron acceptors used by sulphate reducing bacteria (SRBs) and denitrifying bacteria respectively. The former are often present at

significant concentrations in sewage sludge and industrial effluents, leading to competition for hydrogen, lower methane productivity and the formation of hydrogen sulphide, a potentially toxic compound (Colleran *et al.*, 1995).

4.6.2 Xenobiotics as electron acceptors

One of the main advantages of anaerobic digestion over aerobic treatment technologies is the ability of anaerobic bacteria to reductively transform a number of oxidized carbon compounds of environmental significance. These remain largely unaffected as they pass through activated sludge plants because their chemically oxidized structure is difficult to degrade further through oxidative metabolism. Typical examples usually have highly oxidized substituents, like chlorine, on an aromatic nucleus and include pesticides, dyes, solvents, explosives and numerous speciality chemicals. As highly reduced environments are able to promote both the chemical and biological reduction of these xenobiotic compounds, anaerobic digestion will often be the most practical option for their destruction in a treatment process.

It appears that in certain groups of anaerobic bacteria, such compounds may act directly as terminal electron acceptors during growth on electron donors such as VFA. Furthermore, this type of metabolism can yield similar amounts of energy as metabolism with typical electron acceptors like nitrate (Mackiewicz and Wiegel, 1998), making their destruction ecologically feasible in the competitive environment of an anaerobic digester.

5 ENVIRONMENTAL FACTORS AFFECTING ANAEROBIC DIGESTION

As discussed above, effective anaerobic degradation of organic matter requires healthy populations of the relevant bacterial groups to be working in synergy. In all biological wastewater treatment processes, the effective removal of pollutants and contaminants depends not only on the metabolic potential

of the microorganisms but also on the existence of suitable environmental conditions to support these activities. In anaerobic treatment processes, mainly as a result of the critical nature of the syntrophic relationships, environmental conditions require stringent monitoring and control if process failure is to be avoided. Consequently, factors such as nutrient composition, temperature, pH, mixing, toxicity and inhibition have all been thoroughly investigated, and the salient points are summarized below.

5.1 Nutrients

The nutritional requirements of anaerobic bacteria are of paramount importance because nutrients supply the basic cellular building blocks for growth and ensure the cell is able to synthesize the enzymes and cofactors that drive the biochemical and metabolic reactions. Nutrients can be divided into two groups, the macronutrients and micronutrients, according to the relative quantities required by the cell. It is essential for both types of nutrients to be present in an available form in the growth environment to allow effective uptake. Ideally, nutrient levels should be in excess of the optimum concentration required as anaerobic bacteria can be severely inhibited by even slight nutrient deficiencies. However, many essential nutrients can become toxic when present in high concentrations (Gunnerson and Stuckey, 1986), a fact that precludes the use of excess amendments. A rough estimate of the theoretical amount of macronutrients, nitrogen (N), phosphorus (P) and sulphur (S), that are required in digesters can be derived from the elemental composition of bacterial cells within the anaerobic sludge (Lettinga, 1995).

Alternatively, Speece (1996) has suggested that since the empirical formula for biomass is $C_5H_7O_2N$, then the nitrogen requirement would be 3–6 kg N/1000 kg of COD consumed or 0.5–10 kg N/60 m³ of methane produced. However, some workers have cited an optimum C:N ratio of 30:1 (Gunnerson and Stuckey, 1986). Others have preferred to differentiate further the N requirement of a digester

according to whether the system is highly loaded or lightly loaded with COD. Using this approach, Henze and Harremoes (1983) recommended that the COD:N ratios of 350:7 and 1000:7 are required for highly loaded and lightly loaded systems, respectively. Nevertheless, the COD:N ratio most commonly recommended and practised in anaerobic digestion is 100:2.5. The above ratios are all consistent in predicting lower nitrogen requirements for anaerobic digestion than those typically used in aerobic processes (COD:N of 20–30:1). It should be noted that true nutrient balances in both aerobic and anaerobic treatment systems are not fixed relationships and will vary according to operating conditions (organic load, temperature, solid retention time (SRT) etc.).

Nitrogen can occur in a great variety of inorganic forms, the most common being ammonia (NH_3), nitrate (NO_3^-), nitrite (NO_2^-), and nitrogen gas (N_2). Ammonia is the most readily utilized of the inorganic forms of nitrogen, existing in the reduced state that is required for anabolic metabolism and an uncharged state that facilitates cellular uptake.

Several values for the phosphorus requirement have been reported on the basis of batch experiments with pure cultures and by estimating the value from the empirical formula for biomass (Archer and Kirsop, 1990; Lettinga, 1995; Speece, 1996). Although most of these estimates predict an N:P ratio of 7:1, some propose a COD:P ratio varying from 80:1 to 200:1. The usual forms taken by phosphorus in aqueous solution include orthophosphate, polyphosphate, and organic phosphate. The orthophosphates, e.g. PO_4^- , HPO_4^{2-} , H_2PO_4^- , H_3PO_4 , are immediately available for biological metabolism without further modification, organic phosphates must generally be hydrolysed by the cell to release inorganic phosphate before use.

In addition to the two main macronutrients (N and P), the sulphur (S) requirements of anaerobic bacteria should also be satisfied and this can be supplied as sulphur, sulphide, sulphite, thiosulphate, sulphate or amino acids (cysteine and methionine). Optimum

TABLE 24.6 Typical elemental composition (mg/l) of methanogenic bacteria

C	370 000	–	440 000
H	55 000	–	65 000
N	95 000	–	128 000
Na	3000	–	40 000
K	1300	–	50 000
S	5600	–	12 000
P	5000	–	28 000
Ca	85	–	4500
Mg	900	–	5300
Fe	700	–	2800
Ni	65	–	180
Co	10	–	120
Mo	10	–	70
Zn	50	–	630
Cu	<10	–	160
Mn	<5	–	25

Adapted from Scherer *et al.* (1983).

digester concentrations of S have been reported between 0.001 and 1.0 mg/l (Speece, 1996).

As for micronutrients, anaerobic bacteria require a range of trace elements for metabolism and growth. These requirements can be inferred from the elemental composition of anaerobic bacteria (Table 24.6). In addition, the physiological effects and working concentrations of individual micronutrients that have been reported to improve process performance are listed in Table 24.7.

5.2 Temperature

Temperature is one of the most influential environmental factors as it controls the activity of all microorganisms through two contrasting effects. Generally, a rise in temperature leads to an increase in the rate of biochemical and enzymatic reactions within cells, causing increased growth rates. However, above a specific temperature, which is characteristic of each species, this phenomenon gives way to one of inhibition, and then mortality, as the proteins and structural components of the cell become irreversibly denatured.

Although methane formation is biologically feasible at all temperatures between 0°C and 100°C, there are two distinct temperature

TABLE 24.7 Effects of micronutrients on some physiological and operational processes of anaerobic digestion (after Ince, 1993; Kasapgil, 1994)

Nutrient	Concentration required (mg/l)	Effects on digestion
Ca	100–200	Granulation and increase in activity
Mg	75–150	Granulation and increase in activity
Na	100–200	Increase in activity
Fe	20–100	Increase in activity and precipitation of sulphide
K	200–400	Increase in activity
Ba	0.01–0.1	Divalent cation effect hence good granulation
Co	20	Vitamin B ₁₂ dependent
W	–	Formate dehydrogenase
Se	0.8	Formate dehydrogenase, glycine reductase, hydroxylase, and dehydrogenase dependent
SO ₄	0.1–10	Sulphur source of cell synthesis

ranges associated with anaerobic digestion. These are defined as mesophilic, with a temperature optimum at 30–37°C and thermophilic, with an optimum at 55–60°C, the former being most commonly employed in engineered processes of anaerobic treatment. A third range, favoured by psychrophilic organisms has a temperature optimum at 15–20°C (ambient). Although it is not as efficient as high rate mesophilic and thermophilic digestion, it may still have desirable economic trade-offs for the anaerobic treatment of wastewaters in temperate climates.

It is generally believed that thermophilic reactors are more efficient than mesophilic reactors and bench-scale experiments reveal methane production rates in thermophilic reactors can be double that of mesophilic reactors. Thermophilic reactors can also accept higher organic loading rates and produce lower quantities of sludge. However, a number of disadvantages have been observed for full-scale thermophilic digesters which include:

1. they are often less stable than mesophilic reactors

2. they require more energy to heat the reactor
3. they produce high concentrations of VFA in their effluent.

Nevertheless, thermophilic anaerobic digestion is an attractive option for treating warm industrial effluents and slurries of relatively constant composition (Lettinga, 1995).

Compared to many aerobic processes which are relatively robust to temperature variations, anaerobic digestion is sensitive to sudden temperature fluctuations; changes as small as 1–2°C having significant adverse effects on process performance particularly when changes occur rapidly (<2 hours). Should the bacteria become adversely affected by digester temperature variations, several days or even weeks may be required to restore a healthy population once again.

5.3 pH

Anaerobic bacteria, especially methanogens, exhibit a characteristic sensitivity to extremes of pH. Therefore, maintaining a suitable and stable pH within the digester should be a major priority for ensuring efficient methanogenic digestion. This is because the hydrogen ion concentration has a critical influence on the microorganisms responsible for anaerobic digestion, the biochemistry of digestion, alkalinity buffering and several other chemical reactions affecting the solubility and availability of dissolved ions. The best pH range appears to be around neutrality, while the range between 6.5 and 7.8 is generally believed to be optimal. However, there are exceptions for which no satisfactory explanation has been given, such as anaerobic digestion occurring under conditions as low as pH 3 (Zehnder *et al.*, 1982) and as high as pH 9.7 (Oremland, 1988).

There are four major types of chemical and biochemical reaction that influence the pH of a digester. These reactions are:

1. ammonia consumption and release
2. volatile fatty acid production and consumption
3. sulphide release by dissimilatory reduction of sulphate or sulphite

4. conversion of neutral carbonaceous organic carbon to methane and carbon dioxide (Anderson and Yang, 1992).

In an effective working digester, pH reduction can be countered by natural processes such as bicarbonate alkalinity and the consumption of volatile fatty acids by methanogens. However, the latter is dependent on the equilibrium between acidogens and methanogens and this can be easily upset by changes in the operational or environmental conditions (Anderson and Yang, 1992). Should this occur, there are two options to rectify the situation. The first approach is to stop feeding the reactor, giving the methanogens sufficient time to consume excess fatty acids and raise the pH value to an acceptable level. The second option is to dose the reactor with alkali, i.e. NaOH or Na₂CO₃, in order to raise the pH or provide additional buffering capacity. In some cases both options may be used simultaneously (Gunnerson and Stuckey, 1986).

An appreciation of bicarbonate alkalinity and its relationship with the biochemistry of anaerobic digestion must be gained if the pH balance is to be maintained. The alkalinity in water is due principally to salts of weak acids and strong bases, (e.g. bicarbonate ions). Such substances act as buffers when the pH of the environment is approximately equal to their pK_a value, resisting changes in the hydrogen ion concentration that result from acid production or consumption (Sawyer *et al.*, 1994). Although the amount of alkalinity required to accommodate VFA increases in a reactor depends on many factors (Speece, 1996), well-established anaerobic reactors treating typical organic loads are likely to contain alkalinity in the range 2000 to 3000 mg/l as CaCO₃. This level of alkalinity will impart an improved resistance to acidification caused by short-term fluctuations in feed composition.

5.4 Mixing

Methanogenic anaerobic digestion comprises an inherent degree of mixing from the continuous rise of methane bubbles within the reactor, however, this natural mixing is usually con-

sidered to be rate limiting for efficient mass transfer. Consequently, contact between the organic matter and the microorganisms can be improved by enhanced mixing, leading to higher reactor performance. The level and type of mixing also affects the growth rate and distribution of microorganisms within the sludge, substrate availability and utilization rates, granule formation, and gas production. Mixing can be enhanced using:

1. mechanical devices (paddles, turbines and propellers)
2. hydraulic shear force (feed recycle)
3. gas recirculation.

Stafford (1981) reported that so long as adequate mixing was achieved, the method of mixing had little bearing on the digestion rate. However, it was also reported that excessive mixing could actually lead to a reduction in reactor performance. Smith *et al.* (1996) stated that although plug-flow conditions enable feed components to remain in the reactor for one complete hydraulic retention time (HRT), giving maximum theoretical contact time, the potentially high concentration of substrates and their fermentation products that can occur at the reactor inlet may inhibit the biomass. On the other hand, although excessive mixing alleviates this problem, it may result in short-circuiting of the reactor, leading to unconverted substrate appearing in the reactor effluent. In practice, an intermediate degree of mixing appears to give the best substrate conversion by striking a balance between these adverse effects. The relationship between performance and mixing can be quantified using spikes of lithium tracer in the reactor feed to indicate mixing efficiency (Uyanik *et al.*, 2002a).

5.5 Toxicity and inhibition

The definitions of toxicity and inhibition within the context of anaerobic digestion have been defined by Speece (1996) and will be adopted here. According to these, toxicity is an adverse effect (not necessarily lethal) on bacterial metabolism, while inhibition is an impairment

themselves toxic. For example, depending on pH, propionic acid concentrations in the order of grams per litre can be tolerated with a minimal degree of toxicity. However, at low pH values much more of the propionic acid exists in the undissociated HPr form which is much more toxic than propionate ion, Pr^- , due to its greater membrane permeability. In a well-operating digester running with lightly loaded feed, VFA concentration is typically less than 100 mg/l (Hajarnis and Ranade, 1994).

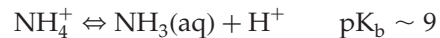
5.8 Oxygen

Strict anaerobes are very sensitive to oxygen exposure. Oxygen can cause an irreversible dissociation of some enzymes and cofactors, such as F_{420} . Another adverse effect of oxygen stems from its capacity to increase the standard redox potential (E_h). Optimum methane production occurs with a redox potential of between -520 and -530 mV with a limiting value of around -350 mV. Therefore, highly reduced environments (absence of oxygen) should be maintained to promote obligate anaerobic bacteria (Pfeffer, 1979). However, methane production is possible even in the presence of oxygen because many of the fermentative bacteria involved in the initial stages are facultative, and will consume the oxygen that may be present in a reactor feed. Furthermore, anaerobic bacteria often exist in structured communities (e.g. granules, biofilms and flocs) where the outer layers of cells are responsible for creating anoxic or anaerobic core microenvironment suitable for the sensitive methanogens.

5.9 Ammonia

Ammonia is released by the fermentation of amino acids and proteins and the breakdown of methylamine and other nitrogenous compounds (Anderson and Yang, 1992). Although ammonia acts to buffer some of the acidity generated by anaerobic digestion, and will be beneficial to anaerobic bacteria at low concentrations, high concentrations can lead to process failure. Depending on the pH of anaerobic reactors, ammonia can be present either in the

form of ammonium ion (NH_4^+) or as ammonia gas in solution (NH_3). The equilibrium between them can be expressed as:



At pH values around neutrality, most ($>99\%$) of the $\text{NH}_3\text{-N}$ will be present as NH_4^+ , which is much less toxic than dissolved ammonia. However, at the higher pH values that are compatible with anaerobic digestion (pH 8), the equilibrium shifts towards the more toxic free ammonia, with concentrations of the latter almost an order of magnitude higher than at pH 7. The maximum concentration of free (dissolved) ammonia should not exceed the inhibitory threshold of 150 mg/l, quoted by Kasapgil (1994), although depending on the operating pH, reactors may be acclimatized reliably to $\text{NH}_3\text{-N}$ concentrations of several thousand mg/l.

5.10 Sulphide

Inorganic forms of sulphur present in reactor feeds, mainly sulphate, are rapidly converted by sulphate-reducing bacteria (SRB) to the reduced forms, sulphide (S^{2-}), and hydrogen sulphide (H_2S), which are ranked as important inhibitors of anaerobic digestion (Anderson *et al.*, 1982). Sulphides in anaerobic treatment can result from other sulphur-containing compounds in the feed, and will be prevalent during anaerobic protein degradation (McCarty, 1964). It is thought that sulphide (S^{2-}) inhibition of anaerobic digestion arises from:

1. competitive consumption of the methanogenic substrates, acetate and hydrogen, as SRB reduce sulphate to sulphide, lowering methane production
2. its toxic, even fatal, effect on anaerobic microorganisms (it reacts chemically with their cytochromes and other iron-containing compounds)
3. Precipitation of essential trace metals (micronutrients) as insoluble metal sulphides.

The degree of sulphide inhibition is largely dependent on pH, temperature, metal ion concentration and operation type. However, inhibition of acetate-utilizing methanogens has been observed at concentrations of 250 mg S/l (Koster *et al.* 1986). Maillacheruvu and Parkin (1996) showed that the H₂S toxicity of the hydrogen-utilizing methanogens ($K_i = 625$ mg S/l) is relatively weaker than for other microbial groups, explaining why methanogenesis from complex substrates can occur even at high concentrations of sulphide.

Hydrogen sulphide inhibition may be mitigated by increasing the pH (H₂S converted to the less toxic HS⁻ form), chemically scrubbing and recycling the reactor gas, or the addition of metals (e.g. Fe precipitates the sulphide as iron sulphide and Mo inhibits the SRB that generates the H₂S).

In addition to those mentioned above, Oremland (1988) listed several other chemical and organic compounds which can cause inhibition in anaerobic reactors, e.g. chlorinated methanes, 2-bromoethanesulphonic acid, iodopropane, monensin, unsaturated carbon-carbon bonds, halogenated aromatic compounds, and some aromatic chemicals.

In toxicity control, it is important to diagnose the inhibition first, and then take the necessary actions. This requires close process monitoring. Monitoring should be implemented in either the liquid/slurry phase or the gas phase. The former should include measurements of pH, total and individual VFA, alkalinity, COD, BOD and solids. The latter usually involves measurements of gas production rate and gas composition (methane and carbon dioxide). In addition to these chemical analyses, microbiological analyses (enumeration of anaerobic bacteria by microscopy, agar plate count and MPN), and biochemical analyses (ATP, coenzyme F₄₂₀, specific methanogenic activity (SMA) and dehydrogenic activity) have been undertaken to give an early indication of metabolic inhibition (Kasapgil, 1994).

Gunnerson and Stuckey (1986) have proposed the following measures to control reactor toxicity:

1. remove toxic substances from the feed
2. dilute the feed to bring levels below the toxic threshold value
3. add chemicals to form a non-toxic complex or insoluble precipitate
4. add an antagonistic substance.

Application of one or more of these measures may attenuate the apparent toxicity of an industrial effluent, making it more amenable to anaerobic treatment.

6 REACTOR CONFIGURATIONS

Anaerobic treatment of wastewaters has existed as a practical technology for over 100 years. It gradually evolved from a simple uncontrolled septic tank system to the high rate, completely controlled reactors now used for treating complex industrial effluents. In anaerobic systems, the key microbial populations generally have a lower reproductive growth rate than their counterparts in aerobic reactors. This requires that a longer sludge retention time (SRT) be provided in order to allow a stable equilibrium to be achieved between the diverse microbial community members in the anaerobic sludge. The loading rates permissible in an anaerobic waste treatment process are, therefore, primarily dictated by the concentration of active biomass within the digester. Consequently, the maintenance of a high SRT has been, at least until recently, the major point of interest in the practical application of anaerobic processes (Lettinga *et al.*, 1980; Anderson and Saw, 1986). As the anaerobic technologies advanced, it became evident that high rate anaerobic treatment could be achieved by employing efficient biomass retention methods. These allowed the solids retention time (SRT) to be maintained well in excess of the hydraulic retention time (HRT) resulting in higher biomass densities within the system. Consequently, improved conservation rates per unit volume of reactor were possible and the high biomass densities also conferred greater resistance to any inhibitory substances in the feed. To accomplish the higher treatment efficiency and reliability associated with a long SRT, a number of novel anaerobic reactor configurations have been developed. During the last two decades, a

succession of small modifications to existing reactor designs has enabled engineers to develop completely new reactor configurations. Most of these, however, have only been evaluated at bench-scale and pilot-scale. An illustration of anaerobic reactor configurations covered in this section is given in Fig. 24.8.

6.1 Conventional or completely mixed anaerobic digester

The concept of the conventional anaerobic digester goes back to 1881 when the first

anaerobic digestion process was developed by the French inventor, M. Louis Mouras. The reactor, called the 'Mouras automatic scavenger', was an air-tight chamber and was used to liquidify the solid components of sewage. In 1891, the first septic tank to retain solids was constructed in England by W.D. Scott Moncrieff. Later, William Travis developed the 'Travis' tank at Hampton in 1904 in order to separate and ferment septic sewage sludge in a separate reaction chamber. This was followed by the development of the 'Imhoff' tank in Germany by Karl Imhoff in 1905. These simple

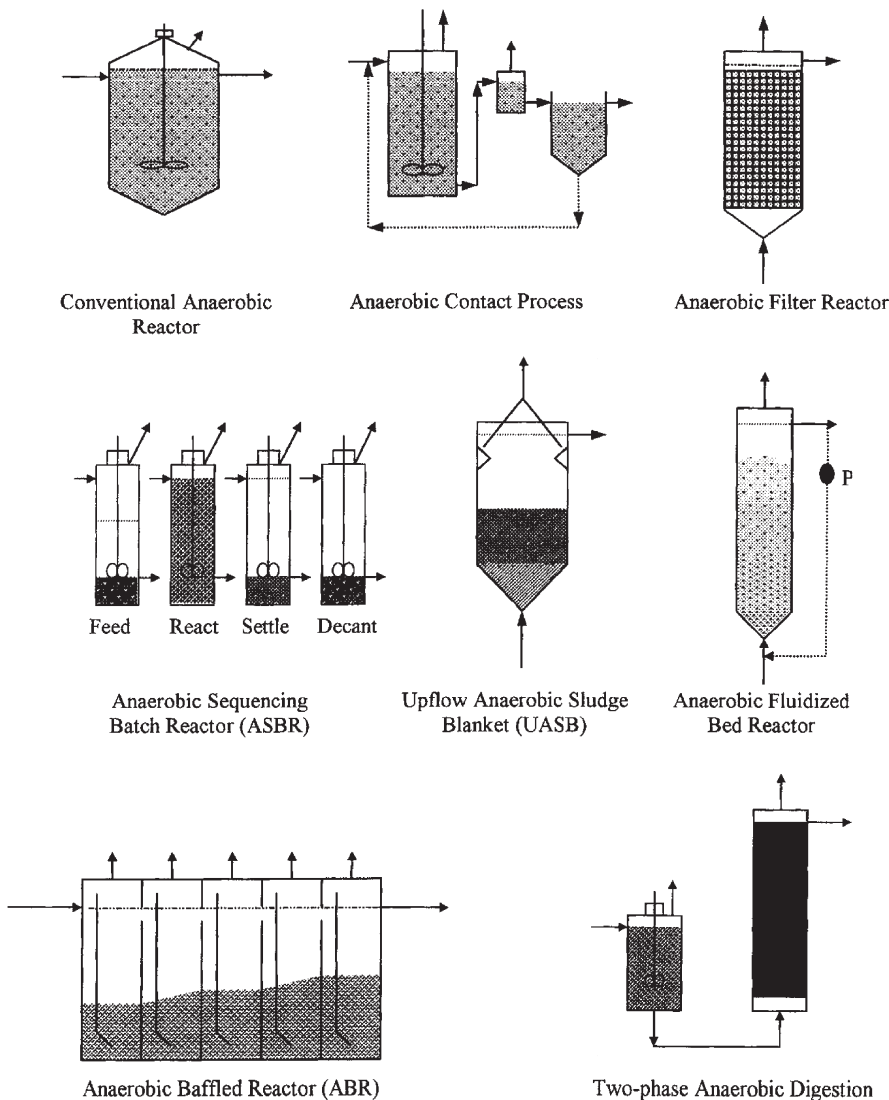


Fig. 24.8 Anaerobic reactor configurations used in wastewater treatment.

unmixed chambers provided a means of enhanced sludge retention within a reactor chamber. Later on, the digesters were also mixed and heated to improve digestion rate. At this time, only the suspended solids fraction of the sewage was subjected to anaerobic treatment and it was not until the development of the anaerobic contact tank by Schroepfer in 1955 that anaerobic digestion was used for anaerobic treatment of dilute wastewaters. Thereafter, conventional anaerobic digesters have been used for the degradation of organic matter in both sewage sludges and industrial wastewaters (McCarty, 1985).

The conventional anaerobic digester is a completely mixed reactor with no solids recycle in which the solids retention time equals the hydraulic retention time. In these reactors, wastewater and anaerobic bacteria are mixed together and allowed to react. When the anaerobic bacteria have reduced the organic pollutant load to the desired level, the treated wastewater is then removed from the reactor for disposal. This system can be operated in either batch or continuous mode and depends on the continuous growth of new biomass to replace that lost in the effluent. We know that some key methanogens are slow-growing organisms with doubling times of 5–10 days; hence at least 10 days of solid and hydraulic retention time will be required. This will necessitate a reactor with a very large volume. The time requirement can be reduced by applying phase separation/solid recycle process, i.e. two tanks, one for mixing and reaction, and the other for solids settlement and storage (see Anaerobic Contact Process).

The large volume requirements and wash-out of microorganisms in the effluent pose serious problems downstream and make conventional anaerobic digesters unsuitable for use with most industrial wastewaters. However, they can be used successfully for sludge treatment and for wastewaters that contain high solids and organic matter content.

6.2 Anaerobic contact process

The link between high biomass concentration, greater efficiency and smaller reactor size

seeded the idea of the contact tank process. The principle of the anaerobic contact reactor has parallels with the activated sludge process, i.e. settling of microbial sludge in a settling tank and its return back to the reactor allows further contact between biomass and raw waste (see Fig. 24.8). In anaerobic contact reactors, due to the arrangement for solids recycle, the SRT is no longer coupled to the HRT as in conventional digesters. As a result, considerable improvements in treatment efficiency can be achieved. Good mixing is essential within the anaerobic contact digester so that adequate mass transfer occurs between the feed and active biomass.

The major shortcoming for contact reactors is their reliance on favourable settling characteristics of the anaerobic sludge, a requirement which may not be fulfilled at all times. Poor sludge settlement arises from the growth of filamentous anaerobic bacteria and gas formation by anaerobic bacteria in the settling tank. Anderson and Saw (1986) suggested that filamentous growth may be a result of nutrient deficiency and this can be rectified by careful maintenance of the N/P ratio. The problems of gas formation can be minimized by employing vacuum degasification or thermal shock prior to sedimentation, by using flocculating agents in the settling tank or by incorporating inclined plates into the settler design.

Although simple in concept, the individual component units of the anaerobic contact process make it more complex than other high rate anaerobic digester systems, however, the absence of any internal fittings offers some advantages for the treatment of wastes having a high solids content (Wheatley, 1990). Typical performance of the anaerobic contact process is reported in Table 24.11.

6.3 Anaerobic sequencing batch reactor (ASBR)

The anaerobic sequencing batch reactor (ASBR) process was developed by Dague's group at Iowa State University. It is a batch-fed, batch-decanted, suspended growth system and is operated in a cyclic sequence of four stages: feed, react, settle and decant (Wirtz and Dague,

TABLE 24.11 Typical operating conditions of various anaerobic digester configurations

Reactor type	Load (kg COD/m ³ day)	HRT (hour)	COD removal (%)
Conventional anaerobic reactor	1–5	240–360	60–80
Anaerobic contact reactor	1–6	24–120	70–95
Anaerobic sequencing batch reactor	1–10	6–24	75–90
Anaerobic filter	2–15	10–85	80–95
Fluidized bed	2–50	1–4	80–90
UASB	2–30	2–72	80–95
Anaerobic baffled reactor	3–35	9–32	75–95
Two phase anaerobic digestion	5–30	20–150	70–85

1996). It comprises a single tank in which all events take place, and since a significant part of the cycle-time is spent settling the biomass from the treated wastewater, the reactor volume requirement is higher than for continuous flow processes. However, this disadvantage is largely offset by its simplicity (it requires no additional biomass settling stage or solids recycle) and the absence of feed short-circuiting which often occurs in continuous flow systems. It has also been reported that biomass granulation (see the next section) can occur in an ASBR after long periods (300 days) of operation. Wirtz and Dague (1996) claimed that the time required for granulation could be shortened by approximately 2 months when granulation enhancements such as granular active carbon (GAC), silica, polymers and ferric chloride were utilized. Operational cycle-times for the ASBR can be as short as 6 hours (Wirtz and Dague, 1996; Banik *et al.*, 1997).

6.4 Anaerobic packed bed or anaerobic filter

Introduced by Coulter *et al.* in 1957 and developed by Young and McCarty in 1967, the anaerobic filter is a fixed-film biological wastewater treatment process in which a fixed matrix (support medium) provides an attachment surface that supports the anaerobic microorganisms in the form of a biofilm. Treatment occurs as the wastewater flows upwards through this bed and the dissolved pollutants are absorbed by the biofilm (Young, 1983). Anaerobic filters were the first anaerobic systems that eliminated the need for solids

separation and recycle while providing a high SRT: HRT ratio. Various types of support material can be used, such as plastics, granular activated carbon (GAC), sand, reticulated foam polymers, granite, quartz and stone. These are usually particulate, moulded or sheet materials that have been produced specifically for the task, often having exceptionally high surface area to volume ratios (400 m²/m³) and low void volumes. The simplicity and robustness of the anaerobic filter, i.e. its resilience to shock loads and operational perturbations (e.g. pH, flow rate inhibitors etc.), make it ideally suited to treatment of both dilute soluble wastewaters and higher-strength soluble wastewaters that can be diluted by recycling (Wheatley, 1990).

The limitations of the anaerobic filter are mostly physical ones related to the deterioration of the bed structure through a gradual accumulation of non-biodegradable solids. This leads eventually to channelling and short-circuiting of flow, and anaerobic filters are therefore unsuitable for wastewaters with a high solids content. Additionally, there is a relatively high cost associated with the packing materials. To overcome some of these problems, packed bed reactors can be operated in a downflow mode which forces the non-biodegradable solids out of the system.

6.5 Anaerobic fluidized and expanded bed reactors

The concerted research efforts of the mid-1970s greatly improved the performance of existing anaerobic technologies and resolved many of the early operational problems, e.g. the diffi-

culties of biomass separation experienced with conventional and contact anaerobic digesters and the blocking problems encountered in anaerobic filters. One such development was the fluidized bed reactor, originally proposed by Jeris in 1974 during denitrification studies (van Haandel and Lettinga, 1994). Adaptation of this system to anaerobic digestion was made by Jewell and his colleagues in the 1970s in an attempt to develop a biological reactor that would accumulate a maximum active attached biomass yet still handle fine suspended solids without blocking (Jewell, 1985). They suggested that if the surface area available for microbial attachment was maximized and the volume occupied by the media minimized, then this should achieve a maximum specific activity of attached biomass for a given reactor volume. In practice, a filter containing extremely small particles (0.5 mm) provides adequate surface area to achieve these benefits. In order to achieve fluidization of the biomass particles, units must be operated in an upflow mode. This simplistic design led to the development of a unit similar to fluidized bed reactors called the 'anaerobic attached film expanded bed reactor' (Jewell, 1980). In fact, it is only the rate of liquid flow and the resulting degree of bed expansion that determines whether the reactor is termed a fluidized bed or expanded bed system (van den Berg and Kennedy, 1983). Expanded bed reactors have a bed expansion of 10% to 20% compared to 30% to 90% in fluidized beds.

In the fluidized bed reactor, the biomass is attached to the surface of small, low specific gravity particles (such as anthracite, high density plastic beads, sand etc.) which are kept in suspension by the upward velocity of the liquid flow (Anderson and Saw, 1986). Effluent is recycled to dilute the incoming waste and to provide sufficient flow-rate to keep particles in suspension. The large surface area of support particles and high degree of mixing that results from the high vertical flows enable a high concentration of biomass to develop and efficient substrate uptake kinetics, respectively. Biomass concentrations between 15 000 and 40 000 mg/l have been reported in these systems (Metcalf and Eddy, 1991).

The greatest risk with the fluidized bed and expanded bed reactors is the loss of biomass particles from the reactor following sudden changes in particle density, flow rate or gas production. If flow is interrupted and the bed allowed to settle, there is a tendency once flow is restarted for the entire bed to move upward in plug-flow rather than fluidizing. Furthermore, in practical applications, considerable difficulties have been experienced in controlling the particle size and density of the flocs due to variable amounts of biomass growth on the particles. Fluidized beds have therefore acquired the reputation for being difficult to operate (Wheatley, 1990; van Haandel and Lettinga, 1994). Such operational drawbacks are often considered to outweigh the inherent performance benefits of these reactors.

6.6 Upflow anaerobic sludge blanket (UASB) reactor

The problem associated with packed bed anaerobic filters and fluidized or expanded bed reactors has led to the development of unpacked reactors that still incorporate an immobilized form of particulate biomass. One of the most successful high-rate unpacked anaerobic digester configurations was developed during the 1970s by Lettinga *et al.* (1980) in the Netherlands, and was descriptively termed the 'upflow anaerobic sludge blanket (UASB) reactor'. This reactor was very similar to the first upflow sludge blanket reactor, the 'biolytic tank', used by Wilson and Phelps in 1910. Lettinga *et al.* (1980) have modified this design and applied it to wastewater treatment in a number of studies; however, recent emphasis has been on its application for warm and more highly concentrated industrial wastewaters and it has been successfully commercialized in this respect (Jewell, 1985). In fact, the UASB reactor is by far the most widely used high-rate anaerobic system for domestic and industrial wastewater treatment (van Haandel and Lettinga, 1994).

The UASB reactor is based on the premise that anaerobic sludge exhibits inherently good settling properties, provided the sludge is not

exposed to heavy mechanical agitation. Adequate mixing between sludge and wastewater is provided by an even flow-distribution combined with a sufficiently high upflow velocity, and by the agitation that results from gas production (Lettinga, 1995). The biomass in this type of reactor is retained as a blanket or granular matrix, and is kept in suspension by controlling the upflow velocity. The wastewater flows upwards through an expanded bed of active sludge located in the lower part of the reactor, while the upper part contains a three-phase (solid, liquid, gas) separation system. The latter facilitates the collection of biogas and also provides internal recycling of the sludge by disengaging adherent biogas bubbles from rising sludge particles (Anderson and Saw, 1986). van Haandel and Lettinga (1994) considered the three-phase separation device to be the most characteristic feature of the UASB. This device, which is located at the top of the reactor, creates three separate internal zones: the digestion zone, the settling zone and the gas and liquid separation zone. Typical performance of the UASB is presented in Table 24.11. Although UASB reactors have been built and operated since 1971, it was not until 1974–1976 that a particularly desirable variant of the anaerobic sludge, granular sludge, was observed inside a pilot plant treating sugar beet factory wastewater. The superior settling characteristics of these sludge granules allowed higher sludge concentrations to be retained and consequently permitted the system to achieve much higher organic loading rates. Granular sludge development has now been observed in UASB reactors treating many different types of wastewater (de Zeeuw, 1987).

The phenomenon of granulation is a process in which a non-discrete flocculent biomass begins to form discrete well-defined pellets, or granules. These vary in dimension and appearance depending on the wastewater and reactor conditions, but generally have a flattened spherical geometry with a diameter of 1–3 mm. The mechanism of biomass granulation has been widely studied the objective being that the rate and extent of granule formation could be manipulated, particularly in waste-

waters that show little intrinsic propensity to granulate (e.g. fat and oil containing effluents).

The consequences of granulation and its influence on the physiology of anaerobic bacteria and UASB reactor performance have been reviewed by Calleja *et al.* (1984). These are summarized as follows:

1. leads to internal physicochemical gradients within the aggregates
2. leads to heterogeneous structured populations of syntrophic microorganisms
3. affects overall stoichiometry, rates of growth and metabolism
4. allows the manipulation of growth rate independent of the dilution rate
5. allows the manipulation of biomass as a single phase
6. generates a reactor effluent with low suspended solids
7. allows high biomass concentrations in continuous reactors
8. allows reactors to be operated continuously beyond normal washout flow rates.

There are several hypotheses to explain the development of granulation, although it is likely that a complex combination of physical, chemical and biological interactions between the dissolved and suspended solids and the microorganisms is required, rather than a single condition.

The phenomenon of granulation has many aspects in common with biofilm formation, a process that has been studied extensively, and it has been postulated that granule formation is probably dependent on the microorganisms participating in the same critical events (Schmidt and Ahring, 1996). These are:

1. transport of microbial cells to the surface of an uncolonized inert material or other microbial cells (termed 'substratum' in the following points)
2. initial reversible adsorption to the substratum by physicochemical forces
3. irreversible adhesion of the cells to the substratum by microbial appendages and/or polymers attaching the cell to the substratum

4. replication of the cells and the development of the granules.

This conceptual chain of events does not address the more complex and interesting issue of how the microbiological and biochemical processes of substrate (pollutant) utilization are linked spatially within the granule. It is probable that the microbial distribution within the internal architecture of the granules is a highly ordered arrangement, with interdependent syntrophic cells existing in juxtaposition to maximize substrate transport efficiency. Such a spatial organization of microorganisms would in part explain the high specific activity of granular sludge relative to flocculent biomass.

6.7 Anaerobic baffled reactor (ABR)

The anaerobic baffled reactor was developed by Bachmann *et al.* (1983), and evolved from initial studies with an anaerobic rotating biological disc reactor. They described the concept of the reactor as essentially a number of upflow sludge blanket reactors connected in series. The design consists of a series of staggered vertical baffles which the wastewater passes over and under as it flows from inlet to outlet (Grobicki and Stuckey, 1991). In the last 20 years much research has been carried out with the ABR, mostly at laboratory-scale. This work has identified some of the advantages and disadvantages of using anaerobic baffled reactors. The main advantages are:

1. it combines the advantages of the anaerobic filter, which has a high stability and reliability due to attachment of the biological solids onto and between the filter media, and the upflow anaerobic sludge process, in which the microbial mass itself functions as the support medium for microorganism attachment, leading to a high void volume (Bachmann *et al.*, 1983)
2. its unique baffled design enables the system to reduce biomass washout, hence retain a high active biological solids content (Polprasert *et al.*, 1992)
3. the reactor can recover remarkably quickly from hydraulic and organic shock loads e.g.

9 hours (Grobicki and Stuckey, 1991; Nachaiyasit and Stuckey, 1997)

4. it has a simple design and requires no special gas or sludge separation equipment (Bachmann *et al.*, 1983; Polprasert *et al.*, 1992; Yu *et al.*, 1997)
5. owing to its compartmentalized configuration, it may function as a two-phase anaerobic treatment system with some spatial separation of acidogenic and methanogenic biomass (Nachaiyasit and Stuckey, 1997; Uyanik *et al.*, 2002b)
6. the fraction of dead space in the ABR is low compared to other designs of anaerobic digesters (Grobicki and Stuckey, 1992)
7. the ABR can be used for almost all soluble organic wastewater from low to high strength (Polprasert *et al.*, 1992)
8. considering its simple structure and operation, it could be considered a potential reactor system for treating municipal and domestic wastewater in tropical and subtropical areas of developing countries (Yu and Anderson, 1996).

There are few significant drawbacks to the ABR mentioned in the literature. The only real disadvantages that can be levelled against this reactor is that it has not been widely used at full-scale. The ABR has been reviewed extensively by Barber and Stuckey (1999).

6.8 Two-phase anaerobic digestion

Two-phase anaerobic digestion implies a process configuration employing separate reactors for acidification and methanogenesis. These are connected in series, allowing each phase of the digestion process to be optimized independently since the microorganisms concerned have different nutritional requirements, physiological characteristics, pH optima, growth and nutrient uptake kinetics, and tolerances to environmental stress factors (Cohen, 1983). The idea of a two-phase anaerobic digestion process was originally proposed by Pohland and Ghosh (1971). It aims to enhance anaerobic biodegradation through a controlled separation of the major reactions, with the hydrolysis, fermentation and acidification reactions

contained within the first phase and the acetogenic and methanogenic reactions predominating in the second phase (Fox and Pohland, 1994). Owing to the fact that bacteria associated with the former reactions have the highest growth and activity rates, the acidogenic reactor will always be smaller than the methanogenic phase reactor.

Several different methods of achieving two-phase separation have been suggested. These include selective inhibition of certain groups of microorganisms, kinetic control, pH control and membrane separation (Anderson and Saw, 1986).

A number of advantages and disadvantages of the two-phase digestion process have been noted by several authors (Cohen, 1983; van den Berg, 1984; Anderson and Saw, 1986; van Haandel and Lettinga, 1994; Kasapgil, 1994; Fox and Pohland, 1994). These are:

Advantages:

1. improvement in process control
2. disposal of excess fast growing acidogenic sludge without any loss of slow growing methanogenic bacteria
3. degradation and attenuation of toxic materials in the first phase (protects the sensitive methanogens)
4. precise pH-control in each reactor
5. higher methane content in the biogas from the methanogenic phase
6. increased loading rate possible for the methanogenic stage
7. balancing tanks in existing treatment plants might be readily converted to acidification tanks for two-phase operation.

Disadvantages:

1. possible disruption of syntrophic relationships
2. high sludge accumulation in the first phase
3. lack of process experience and so more difficult to operate
4. difficulty maintaining a balanced segregation of the phases.

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The nitrogen cycle and its application in wastewater treatment

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1 THE BIOLOGICAL NITROGEN CYCLE

Nitrogen is a necessary element for all living organisms. Living cells contain up to 14% nitrogen, which forms an essential part of several key cell components, such as protein and DNA.

The biological nitrogen cycle can be summarized as in Fig. 25.1. Various forms of nitrogenous compounds exist in nature with oxidation states ranging from -3 to $+5$. Even though (di)nitrogen gas makes up around 79% of the atmosphere, it is chemically extremely inert and generally unavailable to living organisms. Only a select group of microorganisms (*Rhizobium*, *Azotobacter*, *Clostridium*) are capable of fixing nitrogen gas directly into a biological usable form (Postgate, 1998). The product of such nitrogen fixation is ammonia, and this is the essential 'gateway' for the formation of organic nitrogenous compounds, such as plant proteins. In the various assimilatory pathways, nitrogenous compounds of higher oxidation states (such as nitrate and nitrite) can also be reduced to organic nitrogen via ammonia.

Degradation and decay of plant and animal material returns nitrogen to an inorganic form, the principal component of which again is ammonia (via ammonification). In nitrification, another select group of microorganisms is capable of oxidizing free ammonia to nitrogenous compounds of higher oxidation states. The end product of nitrification is nitrate, while

in denitrification, nitrate is reduced by a group of respiratory microorganisms to nitrogen gas.

1.1 Imbalance within the nitrogen cycle

As can be seen in Fig. 25.1, the balance of the nitrogen cycle depends on the combined activities of nitrogen fixation, nitrification, denitrification, nitrogen assimilation and dissimilation. Nitrogen fixation and assimilatory pathways form that part of the cycle (moving from left to right in Fig. 25.1) that is most useful for agriculture, given that the success of commercial plant growth is usually limited by the source of nitrogen. Industrial processes (such as the Haber process) have, thus, been used for the manufacture of nitrogen-based fertilizers, which can be used to improve agricultural performance.

Increased urbanization and industrialization, however, lead to intense cultivation practices that can locally distort the natural balance of the nitrogen cycle and may result in the accumulation of nitrogenous compounds (such as ammonia and nitrate) in rivers and lakes. Typical nitrogen-rich wastewater includes municipal sewage, as well as landfill leachate and wastewater from abattoirs and livestock farms. Table 25.1 summarizes typical concentrations of nitrogenous compounds in these forms of wastewater.

Ammonia, nitrite and nitrate are toxic to aquatic life if present at sufficiently high

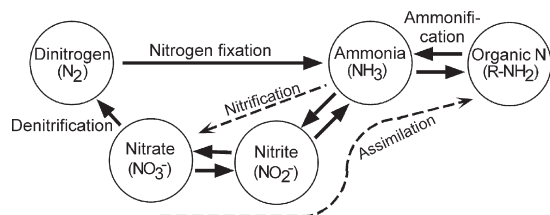


Fig. 25.1 The biological nitrogen cycle.

concentrations. In addition, such nitrogenous material is also known to stimulate growth of algae and other photosynthetic aquatic life, which can lead to eutrophication via excessive loss of available oxygen, and other undesirable changes in the aquatic ecological system. Hence, it is necessary to prevent the accumulation of such nitrogenous compounds in aquatic systems.

To restore the balance of an ecological system, it is natural to exploit the biological mechanisms within the nitrogen cycle to eliminate any accumulation of ammonia and nitrate. More specifically, nitrogen removal from wastewater is achieved via nitrification and denitrification, converting harmful ammonia and nitrate to (essentially) harmless nitrogen gas (i.e. by moving from right to left in Fig. 25.1).

2 BIOLOGICAL TRANSFORMATIONS OF NITROGEN IN WASTEWATER TREATMENT

Nitrogen in municipal and agricultural sewage is usually present in the form of organic nitrogen or ammonia. Fig. 25.2 depicts the

TABLE 25.1 Typical nitrogen concentrations in municipal and agricultural wastewater

Source	Concentration N (mg/l)	Reference
Municipal sewage	20–85	Metcalf and Eddy (1991)
Landfill leachate	1825–2985	Inanc <i>et al.</i> (2000)
Livestock (swine)	390–690	Edgerton <i>et al.</i> (2000)
Abattoir run-off	160–280	Pochana and Keller (1999)

fate of such nitrogen within a wastewater treatment plant. In this figure, bold solid lines indicate that the reaction is coupled to the generation of biomass (anabolism). The rectangular boundary signifies the physical limits of the wastewater treatment plant, while the horizontal dotted line divides the wastewater treatment plant into solid and liquid phases.

Solid hydrolysis converts organic nitrogen (in both soluble and insoluble forms) to ammonia. Biological removal of ammonia is then achieved by first oxidizing ammonia to nitrite/nitrate (via nitrification) which, in turn, is reduced to nitrogen gas (via denitrification). In summary, then, the role of a wastewater treatment plant is to convert nitrogen-rich influent to nitrogen-lean 'clean' effluent, with solid sludge as a by-product of the overall process.

As shown in Fig. 25.2, several nitrification and denitrification routes are possible, depending on the environmental conditions existing within the treatment plant. These will now be discussed separately.

2.1 Nitrification

Nitrification is the aerobic conversion of reduced nitrogenous compounds such as ammonia, hydroxylamine (NH_2OH) and nitrite to more oxidized products. Instead of the ammonium ion, ammonia is the real substrate for the nitrifiers as it is easier to transport the neutral ammonia molecule across the cell membrane (Wiesmann, 1994). Although heterotrophic nitrification has been reported (Robertson *et al.*, 1988), the dominant mechanism for nitrification in wastewater treatment plants is autotrophic in nature.

Chemo-litho-autotrophic bacteria oxidize ammonia to nitrate in two separate stages. The oxidation of ammonia to nitrite is carried out by *Nitrosomonas*, *Nitrosospira*, *Nitrosovibrio* and *Nitrosococcus*, while nitrite oxidizers include *Nitrobacter*, *Nitrospira* and *Nitrococcus* species (Bock *et al.*, 1986; Burrell *et al.*, 1999).

In this oxidation process, ammonia is first oxidized to hydroxylamine with ammonium monooxygenase as the catalyst. The second, energy generating, step is catalysed by hydroxylamine oxidoreductase, which also delivers

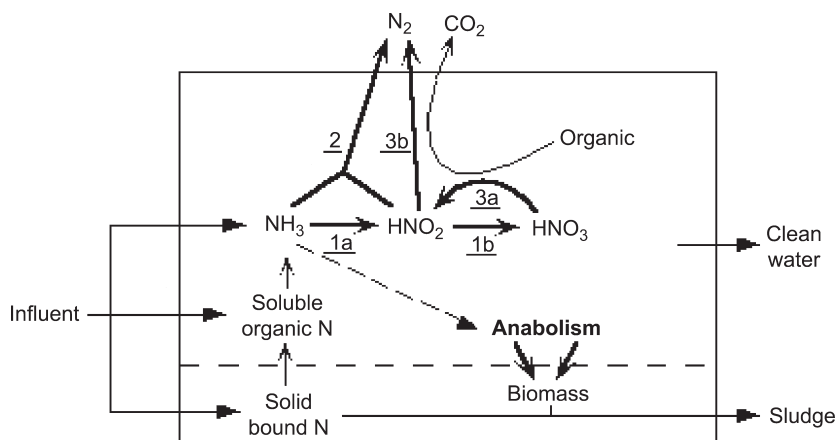
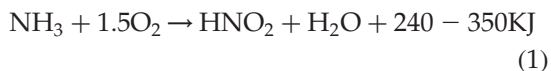
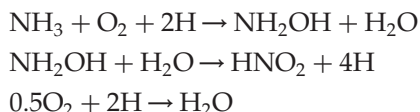
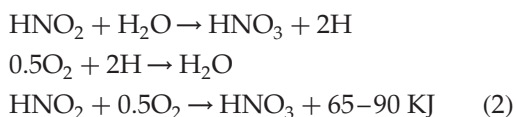


Fig. 25.2 Fate of nitrogen in a sewage treatment plant. (1) Autotrophic nitrification (oxygen consumption is not shown); (2) autotrophic anaerobic oxidation of ammonia; (3) heterotrophic denitrification.

the reducing agents required for the first step. The source of the second oxygen atom in the nitrite is water.



Nitrite is then oxidized to nitrate (by the nitrite oxidizers) with nitrite oxidoreductase catalysing this reaction:

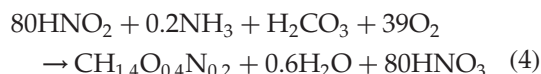
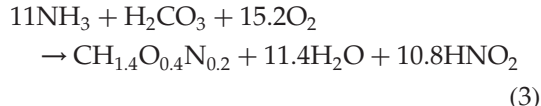


Reducing agents [H] generated in both the oxidation of ammonia and nitrite are oxidized in the bacteria's respiratory chain in order to generate the energy necessary for growth. The terminal electron acceptor for the respiratory chain of both the ammonia and nitrite oxidizers is oxygen, a fact that emphasizes the importance of oxygen to autotrophic nitrification.

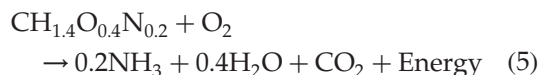
In the case of anabolism, inorganic carbon (such as bicarbonate) is the primary carbon source for the autotrophic nitrifiers. Such carbon fixation is achieved via the Calvin cycle and takes up about 80% of the energy generated from catabolism (Wood, 1986). This energy requirement contributes to the slow

growth rate of both the ammonia oxidizers (about 21 hours doubling time) and the nitrite oxidizers (some 15 hours doubling time).

The overall stoichiometry for nitrification can be derived assuming an average bacteria composition of $\text{CH}_{1.4}\text{O}_{0.4}\text{N}_{0.2}$ with ammonia as the only source of cell nitrogen:



In the absence of substrate, bacteria survive via the process of endogenous respiration:



From Equations (3) and (4), the following yields of biomass from substrate can be derived: 0.15 g-biomass/g-NH₃-N for ammonia oxidizers and 0.02 g-biomass/g-HNO₂-N for nitrite oxidizers. In terms of the yields of biomass on an oxygen basis, the following can also be derived: 0.05 g-biomass/g-O₂ for ammonia oxidizers and 0.02 g-biomass/g-O₂ for nitrite oxidizers.

From the above stoichiometry, it is clear that a significant amount of oxygen is needed for nitrification while the wastewater pH would also decrease as a result of this process.

It should also be noted that the sludge yield from nitrification (0.17 g-biomass/g-NH₃-N) is also low compared to heterotrophic growth (0.40–0.50 g-biomass/g-COD).

2.2 Denitrification

Denitrification is the reduction of oxidized nitrogenous compounds to nitrogen gas (Payne, 1981). This is generally a facultative trait and is carried out by a variety of respiratory bacteria that can utilize oxidized nitrogen compounds (instead of oxygen) as the electron acceptor. Typical denitrifiers include the species *Pseudomonas*, *Alcaligenes* and *Paracoccus*.

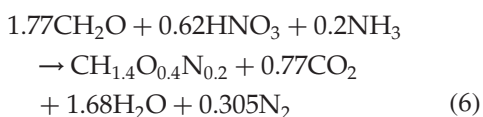
Reduction of nitrate is carried out in two stages. First, nitrate reductase catalyses reduction to nitrite while nitrite reductase completes the reduction to nitrogen gas.

The possible electron donors for denitrification include organic (both extracellular and intracellular) and inorganic ions such as ammonia, sulphide and hydrogen. Each class is discussed below.

2.2.1 Organic denitrification

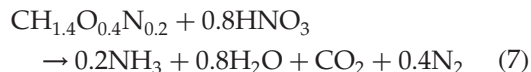
Denitrification with an organic material relies on facultative heterotrophs that are capable of using nitrite and/or nitrate as the terminal electron acceptors in their respiratory chain. In the coupled redox reactions, nitrate/nitrite is reduced to nitrogen gas while the organic material is oxidized to carbon dioxide and water. Virtually all bacteria that are able to reduce nitrate are also able to reduce nitrite. Thus, for simplicity, sometimes a single group of denitrifiers capable of both nitrite and nitrate reduction is assumed.

Organic carbon is the source of cell carbon. Thus, for example, with acetate as the energy source and ammonia as the source of cell nitrogen, the overall stoichiometry for catabolism and anabolism is:



The stoichiometry of endogenous metabolism can be derived as for nitrification,

except that nitrate is the electron acceptor in this case:



The actual growth yield can be derived from Equation (7), considering only one group of denitrifiers, as 0.43-g biomass/g-CH₂O, or 0.40-g biomass/g-COD. The sludge yield from such heterotrophic denitrification is higher than that for nitrification, while denitrification generates alkalinity and, thus, the wastewater pH will increase as a result of this process.

Apart from extracellular substrates, denitrification can also occur using intracellular organic polymer. Heinemann and Müller (1991) co-immobilized denitrifying bacteria with poly-β-hydroxybutyric (PHB) acid in a biopolymeric matrix to denitrify water. The possibility of using PHB for denitrification allows for the combined removal of nitrogen and phosphorus from sewage (Kuba *et al.*, 1996).

2.2.2 Inorganic denitrification

Autotrophic denitrification can be achieved with sulphide, hydrogen and ammonia as the electron donor. In general, however, the concentrations of hydrogen and sulphide in wastewater are too low for them to be regarded as significant electron donors.

Van de Graaf (1997) showed that ammonia could be an electron donor for the reduction of nitrite to nitrogen gas. The ammonia is oxidized under anaerobic (or strictly speaking, anoxic or denitrifying) conditions by nitrite, with both species contributing equally to the nitrogen content in the resultant nitrogen gas.



This process is termed anaerobic ammonia oxidation (Anammox) and is classified as denitrification (rather than nitrification) given that nitrogen gas is generated in the reaction. The bacteria responsible for Anammox are autotrophic, with inorganic carbon as the source of their cell carbon.

2.3 Summary of biological nitrogen cycle in wastewater treatment plants

As shown in Fig. 25.2, removal of ammonia-nitrogen in wastewater treatment plants is dominated by two mechanisms:

1. A conversion pathway to nitrogen gas via the related processes of nitrification and denitrification.
2. An assimilatory pathway for the formation of bacterial biomass.

Sludge production and its subsequent removal is usually a major operating cost in any wastewater treatment plant. To reduce sludge production, endogenous respiratory conditions are often encouraged in a downstream sludge digester. However, sludge digestion means that any assimilated nitrogen is returned as soluble components to the wastewater again (see Equation (5)). Hence, rather than sludge production, the major objective of biological nitrogen removal should be to maximize the conversion of ammonia to nitrogen gas via nitrification and denitrification.

For combined nitrification-denitrification, Fig. 25.2 also shows that there are at least two routes:

- Anaerobic ammonia oxidation (i.e. the Anammox process)
- Autotrophic nitrification followed by heterotrophic denitrification.

In theory, Anammox promises to deliver combined removal of both ammonia and nitrite. In the case of ammonia removal from wastewater, if half of the influent ammonia is first oxidized to nitrite, Anammox could then be used to convert ammonia to nitrogen gas without any aeration and without any organic carbon source. The autotrophic nature of the reaction also promises a low sludge yield. The growth rate of the bacteria responsible for Anammox is, however, extremely low with a doubling time of around 29 days (Van de Graaf, 1997), a fact that translates to a very lengthy start-up time for a full-scale Anammox plant. In addition, for organic-rich wastewater, the

slow growth rate of Anammox microorganisms implies that they would be easily outgrown by organic-consuming heterotrophs.

The application of Anammox has, however, been shown to be feasible given the right operating conditions. For example, for digester effluent at moderately high temperatures (25–35°C, to promote a higher growth rate) and with a low organic concentration, Anammox has been shown to be excellent option for the removal of ammonia (Strous *et al.*, 1997).

Under typical wastewater treatment conditions (10–20°C, and with the presence of organic material), autotrophic nitrification followed by heterotrophic denitrification is the dominant mechanism for biological nitrogen removal. As such, these are the subjects of further discussion for the remainder of this chapter – with ‘nitrification’ referring to autotrophic nitrification while ‘denitrification’ refers to heterotrophic denitrification, unless stated otherwise.

From a pH control perspective, nitrification consumes alkalinity while denitrification generates alkalinity. In theory, combining nitrification with denitrification minimizes the need for pH control. There are, however, several other considerations relating to combined nitrification and denitrification, and these are discussed in the following section.

3 KINETICS OF NITRIFICATION AND DENITRIFICATION

3.1 General considerations

Real bacterial yield is defined as,

$$Y_{X/S}^o = \frac{r_X^g}{r_S} \quad (9)$$

with the specific value given explicitly by the stoichiometry of the reaction. In Equation (9), subscripts ‘X’ and ‘S’ refer to bacteria and substrate, respectively. The specific growth rate is usually assumed to be first-order with respect to the bacterial concentration:

$$r_X^g = \mu^g \cdot X \quad (10)$$

Considering the loss of biomass by endogenous respiration to also be first-order with respect to the bacterial concentration,

$$r_X^e = \mu^e \cdot X \quad (11)$$

the observed yield is, thus, given by:

$$Y_{X/S} = \frac{r_X^g - r_X^e}{r_S} = Y_{X/S}^0 \left(1 - \frac{\mu^e}{\mu^g} \right) \quad (12)$$

Experimentally, the real yield ($Y_{X/S}^0$) cannot be directly observed. It is a 'theoretical upper limit' that can only be estimated for cases where μ^g is significantly larger than μ^e .

The specific growth rate (μ^g) is, however, not a constant. It is a function of the substrate concentration, inhibitor concentration (if any) and other environmental factors such as pH and temperature.

3.2 Autotrophic nitrification

Substrate limitation and inhibition have both been reported for nitrification. Haldane-kinetics are commonly used to describe the inhibition process (Wiesmann, 1994):

$$\mu^g = \mu_{\max}^g \cdot \frac{c(\text{NH}_3 - \text{N})}{K_S + c(\text{NH}_3 - \text{N}) + \frac{c(\text{NH}_3 - \text{N})^2}{K_I}} \times \frac{c(\text{O}_2)}{K_{\text{O}_2} + c(\text{O}_2)} \quad (13)$$

The maximum specific growth rate is given by μ_{\max}^g , while the second factor in Equation (13) describes any oxygen limitation. K_S and K_{O_2} are the half-saturation constants describing substrate and oxygen limitation, respectively, while K_I is a constant related to substrate inhibition.

Fig. 25.3 illustrates the effect of substrate concentration on the relative growth rate (note that the x-axis here is a logarithm scale). The specific growth rate of ammonia oxidizers is, thus, substrate limited at low concentrations and substrate inhibited at high concentrations.

Substrate inhibition is caused by the toxicity of ammonia, which is the reason that the concentration of free ammonia is used in Equation (13). In practice, however, the total ammonium and ammonia concentration,

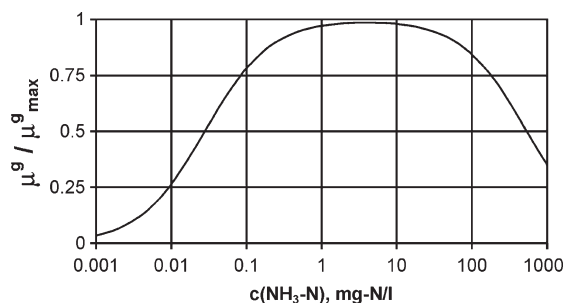


Fig. 25.3 Effect of substrate based on Haldane-kinetics.

$c(\text{NH})$, is usually what is experimentally measured. Given that the ammonia-ammonium equilibrium is pH-dependent,



with an equilibrium constant K_a , we get the following:

$$\begin{aligned} c(\text{NH}) &= c(\text{NH}_3) + c(\text{NH}_4^+) \\ &= c(\text{NH}_3) \cdot (1 + K_a \cdot 10^{-\text{pH}}) \end{aligned} \quad (15)$$

Replacing $c(\text{NH}_3 - \text{N})$ with $c(\text{NH} - \text{N})$, as given by Equation (15), the substrate term in Equation (13) becomes:

$$\frac{c(\text{NH} - \text{N})}{K_S \cdot (1 + K_a \cdot 10^{-\text{pH}}) + c(\text{NH} - \text{N}) + \frac{c(\text{NH} - \text{N})^2}{K_I \cdot (1 + K_a \cdot 10^{-\text{pH}})}} \quad (16)$$

In this case, the new half-saturation constant, $K_S \cdot (1 + K_a \cdot 10^{-\text{pH}})$ is pH-dependent. For the ammonia-ammonium equilibrium, $\text{p}K_a$ equals -9.26 at 25°C . For $\text{pH} < 9$ and $c(\text{NH} - \text{N}) < 65 \text{ mg-N/l}$, the concentration of free ammonia is negligible and the substrate inhibition term can be ignored (Wiesmann, 1994). The effect of the substrate is thus reduced to a limitation at low concentrations and can be modelled using Monod kinetics:

$$\mu^g = \mu_{\max}^g \cdot \frac{c(\text{NH}_3 - \text{N})}{K_S + c(\text{NH}_3 - \text{N})} \cdot \frac{c(\text{O}_2)}{K_{\text{O}_2} + c(\text{O}_2)} \quad (17)$$

A similar analysis can be carried out for nitrite oxidation and the following expression

can be derived:

$$\mu^g = \mu_{\max}^g \frac{c(\text{HNO}_2 - \text{N})}{K_S + c(\text{HNO}_2 - \text{N}) + \frac{c(\text{HNO}_2 - \text{N})^2}{K_I}} \times \frac{c(\text{O}_2)}{K_{\text{O}_2} + c(\text{O}_2)} \quad (18)$$

3.2.1 Effect of other environmental factors

Both ammonia monooxygenase and nitrite oxidoreductase are photosensitive. Thus, under strong light conditions, the nitrifiers will be out-competed for the inorganic carbon source by photosynthetic autotrophs or algae (Wood, 1986). Thiourea or allylthiourea have been used to inhibit nitrification in BOD tests, while the toxicity of other chemicals to nitrification has been summarized by Henze *et al.* (1995).

The effects of temperature and pH could be analysed in terms of an increase or decrease in the value of the maximum specific growth rate (μ_{\max}^g). This rate increases with temperature before the enzymes responsible for nitrification are deactivated over the range 32–35°C. Meanwhile, the optimum pH for nitrification is in the 8–9 range. As already mentioned, the effect of pH on the growth rate is also linked to substrate inhibition, especially at pH values above 9. It has also been reported that pH values lower than about 6.0 are unfavourable for nitrification (Henze *et al.*, 1995).

3.3 Heterotrophic denitrification

Here, the specific growth rate is limited by both the concentration of the substrate and the electron acceptor (HNO_2 or HNO_3), and double Monod-kinetics is usually used. Thus, for the denitrification of nitrate, the specific growth rate is given by,

$$\mu^g = \mu_{\max}^g \frac{c(\text{COD})}{K_S + c(\text{COD})} \times \frac{c(\text{HNO}_3 - \text{N})}{K_{\text{HNO}_3 - \text{N}} + c(\text{HNO}_3 - \text{N})} \quad (19)$$

while a similar expression holds for the denitrification of nitrite:

$$\mu^g = \mu_{\max}^g \frac{c(\text{COD})}{K_{SH} + c(\text{COD})} \times \frac{c(\text{HNO}_2 - \text{N})}{K_{\text{HNO}_2 - \text{N}} + c(\text{HNO}_2 - \text{N})} \quad (20)$$

3.3.1 Effect of other environmental factors

Molecular oxygen has been shown to repress the enzymes responsible for denitrification (Gottschalk, 1979). In the presence of oxygen, the denitrification reaction is inhibited and the denitrifying bacteria switch to using oxygen as their terminal electron acceptor. The inhibition mechanism is, however, reversible and the response of the enzymatic system to the switch of electron acceptor has been shown to only be in the order of minutes (Kornaros and Lyberatos, 1998).

Including the effect of non-competitive inhibition by oxygen, Equations (19) and (20) become, respectively:

$$\mu^g = \mu_{\max}^g \frac{c(\text{COD})}{K_S + c(\text{COD})} \times \frac{c(\text{HNO}_3 - \text{N})}{K_{\text{HNO}_3 - \text{N}} + c(\text{HNO}_3 - \text{N})} \times \frac{K_{\text{O}_2, I}}{K_{\text{O}_2, I} + c(\text{O}_2)} \quad (21)$$

$$\mu^g = \mu_{\max}^g \frac{c(\text{COD})}{K_{SH} + c(\text{COD})} \times \frac{c(\text{HNO}_2 - \text{N})}{K_{\text{HNO}_2 - \text{N}} + c(\text{HNO}_2 - \text{N})} \times \frac{K_{\text{O}_2, I}}{K_{\text{O}_2, I} + c(\text{O}_2)} \quad (22)$$

with $K_{\text{O}_2, I}$ as a constant related to oxygen inhibition.

The effects of temperature and pH on denitrification are similar to those for nitrification. The optimum pH for denitrification is, however, somewhat broader than for nitrification, falling in the range 7–9 (Henze *et al.*, 1995).

3.4 Summary of kinetic data

Average kinetic coefficients for nitrification and denitrification (obtained from the literature) are summarized in Table 25.2. Values of μ_{\max}^g , K_L , K_{O_2} , K_{HNO_x} and $Y_{X/S}^O$ are adapted from Wiesmann (1994), μ^e and K_S are from Wiesmann (1994); and Henze *et al.* (1995), $Y_{O_2/S}^O$ and $Y_{NO_x/S}^O$ are derived from stoichiometry, while $K_{O_2,I}$ are from Gujer *et al.* (1999).

Note that the coefficients related to organic degradation are strongly dependent on the nature of the organic substrate, i.e. μ_{\max}^g and K_S for aerobic and denitrifying organic degradation. Note that the values given here are only representative. For more specific values, these parameters should be experimentally measured for the wastewater of interest.

The data given in Table 25.2 highlight several significant differences between nitrification, aerobic organic degradation and denitrification:

1. Nitrification requires oxygen while denitrification is inhibited by oxygen.
2. Both sludge yield and maximum specific growth rate are significantly lower for nitrification than for organic degradation.

The different effects of oxygen link directly to the aeration pattern in nitrogen-removing plants. Moreover, the differences in sludge yield and maximum specific growth rate

imply that the nitrifiers are usually outgrown by organic-consuming heterotrophs, unless a two-sludge system is used. Design consideration of nitrogen-removing plants will be discussed in the next section.

4 DESIGN OF BIOLOGICAL NITROGEN REMOVAL PLANTS

4.1 Overview

The primary design objective in the design of a nitrogen-removing plant is to devise a system that is capable of reducing the concentrations of nitrogenous compounds to the desired levels, although clearly the capital and operating costs are also important considerations in the final decision. The capital cost is usually dominated by the size of the reactor used and its associated solid-liquid separation unit, while the dominant factors in the operating cost include:

- Aeration
- Sludge removal
- Pumping cost (primarily for any recycle stream required)
- Chemical dosage requirement.

Over the past few decades, a variety of process flowsheets for nitrogen-removing plants have been proposed, with the basic structure shown in Fig. 25.4. For such a flowsheet structure,

TABLE 25.2 Typical kinetic coefficients (20°C and a pH of 8)

	NH_3 oxidation	HNO_2 oxidation	Aerobic degradation of organic	HNO_3 reduction	HNO_2 reduction
μ_{\max}^g	0.77/d	1.08/d	7.2/d	2.6/d	1.5/d
μ^e	0.03/d	0.03/d	0.2/d	0.1/d	0.1/d
K_S	0.028 mg- NH_3 -N/l	$3.2 \cdot 10^{-5}$ mg- HNO_2 -N/l	10–100 mg-COD/l	5–25 mg-COD/l	–
K_I	540 mg- NH_3 -N/l	0.26 mg- HNO_2 -N/l	–	–	–
$K_{O_2,I}$	–	–	–	0.20 mg- O_2 /l	0.20 mg- O_2 /l
K_{O_2}	0.3 mg O_2 /l	1.1 mg O_2 /l	0.08 mg O_2 /l	–	–
K_{HNO_x}	–	–	–	≤ 0.14 mg- HNO_3 -N/l	≤ 0.12 mg- HNO_2 -N/l
$Y_{X/S}^O$	0.147 g-cell/ g- NH_3 -N	0.02 g-cell/ g HNO_2 -N	0.43 g-cell/g-COD	0.40 g-cell/g-COD	0.40 g-cell/g-COD
$Y_{O_2/S}^O$	3.16 g- O_2 / g- NH_3 -N	1.11 g O_2 / g- HNO_2 -N	0.40 g- O_2 /g-COD	–	–
$Y_{NO_x/S}^O$	–	–	–	0.378 g- HNO_3 -N/ g-COD	0.255 g- HNO_2 -N/ g-COD

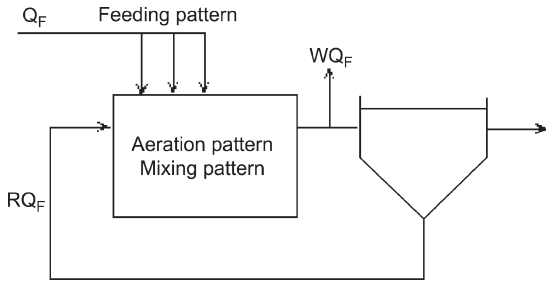


Fig. 25.4 Basic flow sheet for biological nitrogen removal.

the operating decisions available to a design engineer might be summarized as follows:

- Solids retention time
- Reactor mixing pattern
- Recycle ratio
- Feeding pattern to the reactor
- Aeration pattern in the reactor.

The effect of each of these design variables on the overall performance of a nitrogen-removing plant can be analysed from a kinetic point of view.

4.2 Impact of individual design decision

If the effect of endogenous respiration on substrate removal is ignored, then the combination of Equations (9) and (10) shows that the rate of substrate removal is proportional to both the specific growth rate (μ^s) and the bacteria concentration (X):

$$r_S^s = \frac{1}{Y_{X/S}^o} \cdot \mu^s \cdot X \quad (23)$$

X is controlled by the solids retention time. μ^s is a function of the substrate, electron acceptor and inhibitor concentrations, as given by Equations (13) and (18) for nitrification, and Equations (21) and (22) for denitrification. As a result, the average value of μ^s in the reactor is influenced by the recycle ratio, mixing pattern, feeding pattern and aeration pattern. The impact of pH and temperature changes on μ_{\max}^s is often ignored in the first instance, assuming constant values for both.

4.2.1 Solids retention time

The solids retention time or SRT controls the concentrations of bacteria throughout the treatment system. A higher SRT contributes to a higher bacterial concentration in the reactor, which gives rise to:

- Smaller reactor size
- Larger separator size
- Reduced sludge production
- Higher aeration requirements due to the extra oxygen required for endogenous respiration.

Clearly, an optimum SRT exists, resulting from a trade-off between the gains and losses in the various cost terms. For municipal sewage treatment plants performing combined nitrification–denitrification, typical wasting ratios generally fall in the range 0.025–0.10 for a hydraulic retention time of 12–24 hours.

4.2.2 Mixing pattern

Two types of idealized mixing pattern are possible, i.e. plug-flow and mixed-flow. For zero-order kinetics, both mixing patterns give exactly the same reactor size. For first-order kinetics, however, a plug flow reactor is always smaller in volume than a mixed flow reactor carrying out the same amount of reaction (Levenspiel, 1972).

Monod-kinetics are mixed-order, being close to zero-order at high substrate (or electron acceptor) concentration and essentially first-order at low substrate (or electron acceptor) concentration. The defining concentrations are given by the half-saturation constants K_S , K_{O_2} and K_{HNOx} (refer to Table 25.2 for typical values).

For nitrification, K_S is a function of pH – at a pH value of 7 (and a temperature of 25°C), this parameter equals 5.1 mg NH – N/l. This suggests that if an effluent ammonia–ammonium concentration of less than 1 mg NH – N/l is required (as in the case of most municipal sewage treatment plants), a plug-flow reactor is the preferred option. A similar argument holds for organics removal if an

effluent COD concentration of less than 10 mg COD/l is required.

For denitrification, values of K_{HNO_x} generally fall in the range 0.10–0.15 mg HNO_x – N/l, which are low in comparison to typical effluent requirements of 1–10 mg HNO_x –N/l. This suggests that the kinetics will be primarily zero-order with respect to the nitrate concentration, and both plug-flow and mixed-flow reactors give similar performance if organic material is adequate for growth.

4.2.3 Recycle ratio, feeding pattern, aeration pattern

Any recycle stream affects the overall treatment system in two ways – it dilutes the influent concentration, and increases the hydraulic loading on both the reactor and the separator. The first effect reduces the required reactor size, while the second effect reduces the effective hydraulic retention time of the reactor and separator. In addition, pumping costs increase with the recycle ratio.

For modestly loaded municipal sewage treatment plants, the optimum recycle ratio is usually in the range of 0.5–1.0. However, for wastewater with a high ammonia concentration, such as many industrial effluent streams, a higher recycle ratio is required to dilute the influent ammonia concentration to avoid substrate inhibition, which will reduce the effective rate of removal. As shown in Fig. 25.3, the desired influent ammonia concentration in the reactor is in the range 5–30 mg NH_3 –N/l.

As an alternative, step feeding can be used to maximize exposure of the reactor volume to the optimal substrate concentration ranges, and thereby reduce the overall reactor size.

The aeration pattern to be used in the reactor is also strongly related to the wastewater characteristics, as will be discussed in the next section.

4.3 Process flowsheet selection based on wastewater composition

The flowsheet requirements of a treatment system differ quite markedly depending on

the characteristics of the nitrogenous compounds in the wastewater, which can be broadly categorized as follows:

- Nitrate-rich wastewater
- Ammonia-rich wastewater
- Ammonia and organic-rich wastewater.

4.3.1 Nitrate-rich wastewater

Nitrate-rich wastewater includes wastewater from fertilizer industry and metal finishing plants where nitric acid is used as a cleaning agent. For this class of wastewater, an anoxic-aerobic system is commonly used. The source of cell nitrogen in this case is nitrate, which can be achieved through an assimilatory pathway as depicted in Fig. 25.1.

The exact flowsheet of the treatment plant depends on the organic to nitrate ratio of the wastewater. From stoichiometry, 3–5 g of COD is needed for every gram of NO_3 –N reduced (Henze *et al.*, 1995). Based on this ratio, if nitrate is in excess, aeration is not needed, while an organic supplement (methanol is a typical organic supplement for denitrification) may be needed to reduce the nitrate concentration to the desired level (see Fig. 25.5a).

If organic material is in excess, then an anoxic-aerobic system is used first to remove nitrate in the anoxic reactor before any excess organic is oxidized aerobically in the second reactor to meet the desired organic effluent concentration (see Fig. 25.5b).

4.3.2 Ammonia-rich wastewater

For ammonia-rich wastewater that is depleted of organic carbon (such as digester effluent and wastewater from fertilizer industry), Anammox is potentially useful if faster growth rate can be promoted, e.g. if the wastewater temperature is high.

For dilute wastewater, heating up volume of wastewater is cost-prohibitive. In this case, an aerobic-anoxic reactor can be used. In the aerobic reactor, ammonia is oxidized to nitrate while nitrate is reduced in the second anoxic reactor with supplement of organic carbon source (see Fig. 25.5c).

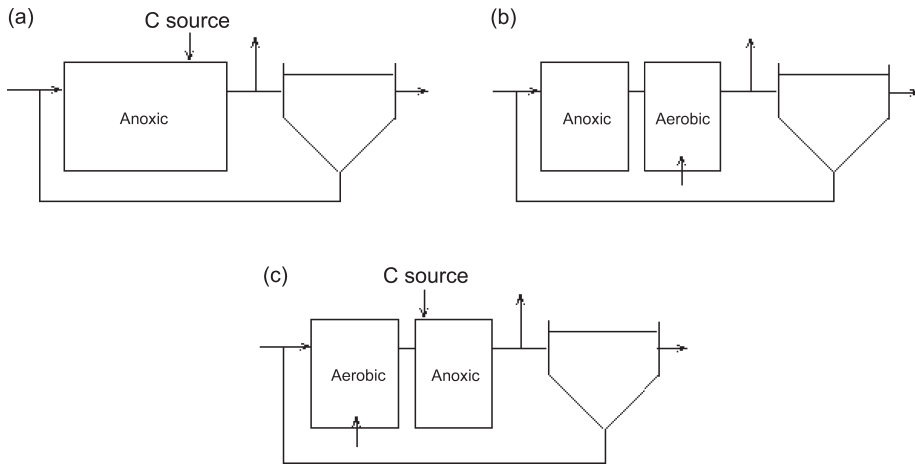


Fig. 25.5 Treatment systems for nitrate-rich and ammonia-rich wastewater.

4.3.3 Ammonia and organic-rich wastewater

Most wastewater (such as municipal sewage, leachate and agricultural run-off) is rich in both ammonia and organic material. For combined removal of organic and nitrogenous material, the conflicting oxygen requirements for nitrification and denitrification usually mean that complete nitrogen removal cannot be obtained. Nitrification produces nitrate that is reduced in the denitrification stage, however, the aerobic nitrifying conditions also promote aerobic degradation of organic material. As a result,

denitrification is usually limited by the organic carbon source, unless an organic supplement is added.

If the organic material in wastewater is in excess of the requirements for denitrification, a single mixed aerobic-anoxic system could be used (see Fig. 25.6a). In such systems, the dissolved oxygen concentration in the reactor is controlled at a low level (typically 0.5–2.0 mg-O₂/l) to allow simultaneous nitrification, denitrification and aerobic organic degradation. Examples of such systems include the 'oxidation ditch' and various attached

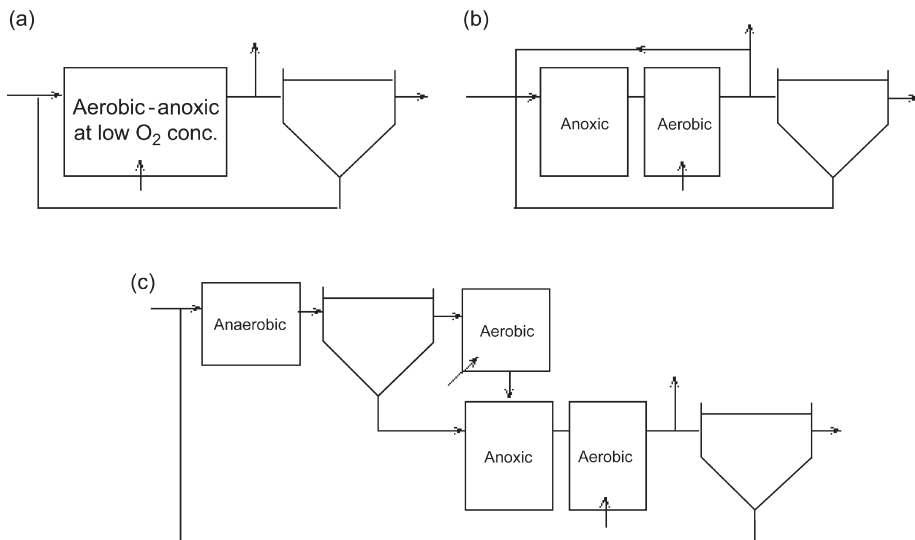


Fig. 25.6 Process flowsheets for ammonia/organic-rich wastewater.

growth systems – the latter often being the preferable option, given that a thicker biofilm layer can be formed which encourages the formation of an anoxic zone for denitrification.

If the ratio of organic to nitrate is not in excess of the stoichiometry required for denitrification, it is necessary to avoid aerobic degradation of the organic material. In this case, an anoxic–aerobic-recycle system can be used (see Fig. 25.6b; note that this is often referred to as the modified Ludzack-Ettinger configuration). Nitrification occurs in the second (aerobic) reactor. Internal recycling of mixed liquor returns nitrate to the front of the reactor system where it is reduced with the influent organic under anoxic conditions. Clearly, more nitrate can be reduced if a higher internal recycle is used, but this can lead to higher pumping costs and an increased hydraulic loading on the reactor (which may in turn require a larger reactor). However, it should be noted that internal recycle ratios in excess of 4–6 have not been reported in the literature for municipal sewage treatment plants.

Internal recycling can be avoided if the organic material can be separated from the ammonia. This may be achieved via sorption of such material onto bacterial flocs (Wanner *et al.*, 1992; Kuba *et al.*, 1996). Subsequent solid–liquid separation allows ammonia-rich supernatant to be nitrified separately, before it is returned as a nitrate-rich stream and reduced with the organic-rich sludge. An aerobic zone is then used to degrade any residual organic material (see Fig. 25.6c). This form of flowsheet is useful for wastewaters with a low organic to nitrogen ratio. Given that there is no internal recycling used, the reactor size is minimal despite the fact that there are several anaerobic/anoxic/aerobic zones. The only (minor) drawback with this configuration is the presence of an additional settler.

5 CONCLUSIONS

Nitrification and denitrification both play important roles in the balance of the nitrogen cycle. In wastewater treatment plants,

nitrification followed by denitrification is commonly used to convert potentially harmful nitrogenous compounds into nitrogen gas. In this chapter the characteristics of the dominant pathways for nitrification and denitrification were reviewed, key kinetic data were summarized (in Table 25.2), before moving on to consider the evaluation of the various design variables for typical nitrogen-removing treatment plants.

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Low-cost treatment systems

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1 INTRODUCTION

Low-cost wastewater treatment systems comprise:

1. Waste stabilization ponds (WSP)
2. Wastewater treatment and storage reservoirs (WSTR)
3. Constructed wetlands.

They are often called 'natural' treatment systems as they do not rely on electromechanical plant, such as aerators, for their operation. The oxygen needed by the heterotrophic bacteria for the oxidation of organic compounds comes mainly from the photosynthetic activity of the algae in WSP and WSTR and of the plants in constructed wetlands. Since this activity depends on sunlight (which arrives at the earth's surface), these processes require a greater surface area of land than electromechanical treatment systems. In developing countries this is often not a problem; in industrialized countries, on the other hand, this generally means that these low-cost treatment processes are used for small rural communities, under around 5000 population.

2 WASTE STABILIZATION PONDS

WSP systems comprise a series of anaerobic, facultative and maturation ponds. The primary function of anaerobic and facultative ponds is BOD removal. Maturation ponds are primarily

for the removal of excreted pathogens, although some pathogen removal occurs in anaerobic and facultative ponds (particularly the removal of *Vibrio cholerae*, helminth eggs and protozoan cysts), and some BOD removal occurs in maturation ponds.

WSP design is detailed in EPA (1983), Mara *et al.* (1991), Mara (1997) and Mara and Pearson (1998). Arthur (1983) presents a methodology for comparative costing of wastewater treatment options, and in his case study (based on the city of Sana'a in the Yemen Arab Republic), WSP were found to be the least-cost technology at land prices of US\$ 50 000–150 000 per ha, depending on the discount rate used (5–15%).

WSP are widely used in France and Germany, where there are around 2500 and 1100 systems, respectively (Bucksteeg, 1987; CEMAGREF *et al.*, 1997), and also in the USA, which has around 7000 systems (EPA, 1983). The UK has, in contrast, only 19 WSP systems (Mara *et al.*, 1998).

2.1 Anaerobic ponds

Anaerobic ponds (2–4 m deep) function much like open septic tanks: they receive such a high organic loading (100–350 g BOD/m³/day, depending on temperature) that they are devoid of both dissolved oxygen and algae (although occasionally a thin surface film of *Chlamydomonas* may be present). In warm climates they are extremely efficient in removing BOD – up to 75% at 25°C and a retention

time of 1 day for wastewaters with a BOD up to 350 mg/l (Silva, 1982; Pearson *et al.*, 1996a). Performance is lower at lower temperatures, but even in European winters BOD removal is 40–50% (Bucksteeg, 1987). Odour (due to H₂S release) is not a problem provided the recommended BOD loadings are not exceeded (see Mara and Pearson, 1998) and the wastewater contains less than 500 mg SO₄²⁻/l (Gloyna and Espino, 1969); the latter should not be a problem with purely domestic wastewater as drinking waters should contain no more than 250 mg SO₄²⁻/l (WHO, 1993), although in coastal areas salt water intrusion into sewers can increase the sulphate concentration of the wastewater.

Vibrio cholerae is very sensitive to sulphide concentrations as low as 3 mg/l, and its removal in anaerobic ponds is consequently very good; Oragui *et al.* (1993) report a reduction from 485 to 28/l.

Helminth eggs and protozoan cysts are removed in WSP by sedimentation. Compliance with the WHO (1989) recommendation for crop irrigation of less than 1 egg per litre is often achieved in a series of anaerobic and facultative ponds, although this depends on the number of eggs in the raw wastewater (which in turn depends on the endemicity of helminthic infections in the community producing the wastewater) and the WSP retention times (see Ayres *et al.*, 1992).

2.2 Facultative ponds

Facultative ponds (1.5–1.8 m deep) (Fig. 26.1) receive organic loads in the range 100–400 kg BOD/ha day and, due to the profuse growth of algae, they appear dark green, although when slightly overloaded they may appear red or purple due to the predominance of anaerobic sulphide-oxidizing photosynthetic bacteria (see Chapter 27). The pond algae and the pond heterotrophic bacteria depend on each other: the algae act as oxygen generators and the bacteria use this algal oxygen for BOD removal, and the bacterially-produced carbon dioxide is fixed by the algae (Fig. 26.2). Some oxygen and some carbon dioxide enter the pond through its surface from the atmosphere,

but most of the oxygen comes from algal photosynthesis and most of the carbon dioxide from bacterial metabolism.

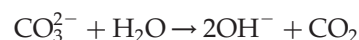
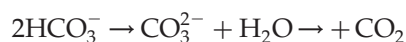
BOD removal in facultative ponds is around 80–90% on a *filtered* basis, i.e. (and as permitted by the European Directive on Urban Wastewater Treatment – Council of the European Communities (CEC), 1991) effluent samples are filtered prior to BOD analysis to remove algal solids (and hence algal BOD). Filtration is through glass fibre filters of the type normally used for suspended solids (SS) determination (e.g. Whatman GF/C). In Europe (CEC, 1991) WSP effluents must contain ≥ 25 mg filtered BOD/l and ≥ 150 mg SS/l, and this is often achieved by facultative pond effluents (see Pearson *et al.*, 1995; CEMAGREF *et al.*, 1997).

2.3 Maturation ponds

The function of maturation ponds (1–1.5 m deep) is the destruction of faecal bacteria, particularly pathogenic members of the Enterobacteriaceae, and faecal viruses. As shown in Table 26.1, this is extremely efficient in warm climates. Faecal coliform (FC) bacteria are commonly used as indicators of faecal bacterial pathogens and, as shown by Marais (1974), FC removal in WSP can be adequately modelled by first order kinetics in a series of completely mixed reactors.

Faecal viruses are removed by adsorption on to settleable solids and subsequent sedimentation, including the adsorption on to algae in facultative and maturation ponds and the subsequent sedimentation of the algae when they die.

Faecal bacterial removal (except, as noted above, that of *V. cholerae*) is largely due to the activities of the pond algae. When the algae are photosynthesizing rapidly, their demand for CO₂ outstrips its supply from bacterial metabolism (see Fig. 26.1) and this leads to carbonate and bicarbonate dissociation:



The resulting CO₂ is fixed by the algae and the hydroxyl ions accumulate to raise the pH to



Fig. 26.1 A facultative pond at Botton Village in North Yorkshire, England.

often above 9, which is rapidly lethal to faecal bacteria (Parhad and Rao, 1974; Pearson *et al.*, 1987). Curtis *et al.* (1992) showed that high levels of dissolved oxygen (due to algal photosynthesis) and high visible light intensities, in the presence of dissolved humic substances, which act as very efficient exogenous photosensitizers, are rapidly fatal to faecal bacteria, a process that is exacerbated by high pH (Fig. 26.3).

A properly designed series of maturation ponds, that is also well operated and maintained (good O & M is principally the removal of vegetation and any floating materials to permit unfettered sunlight penetration for

algal photosynthesis) can easily reduce FC levels to below 1000 per 100 ml, which is the

TABLE 26.1 Management strategy for three WSTR in parallel for an irrigation season of 6 months

Month ^a	WSTR 1	WSTR 2	WSTR 3
January	Rest	Fill (1)	Empty
February	Rest	Fill (1)	Empty
March	Rest	Rest (or Fill (1))	Fill (1) (or Empty)
April	Rest	Rest	Fill (1)
May	Use	Rest	Fill (1)
June	Use	Rest	Fill (1)
July	Fill (1) ^b	Use	Rest
August	Fill (1)	Use	Rest
September	Fill ($\frac{1}{2}$)	Fill ($\frac{1}{2}$)	Use
October	Fill ($\frac{1}{2}$)	Fill ($\frac{1}{2}$)	Use
November	Fill ($\frac{1}{2}$)	Fill ($\frac{1}{2}$)	Empty
December	Fill ($\frac{1}{2}$)	Fill ($\frac{1}{2}$)	Empty
Volume ^c	4	4 (or 5)	4 (or 3)

^a July and August are the hottest months, so WSTR No. 3 has the minimum rest period of two months at this time. The other two WSTR have rest periods of 4 months to ensure FC die-off to <1000 per 100 ml during the cooler months.

^b Proportion of monthly flow discharged into each WSTR.

^c WSTR volume expressed as multiple of monthly wastewater flow. The WSTR need not have equal volumes as the volume of treated wastewater required for irrigation may vary throughout the irrigation season.

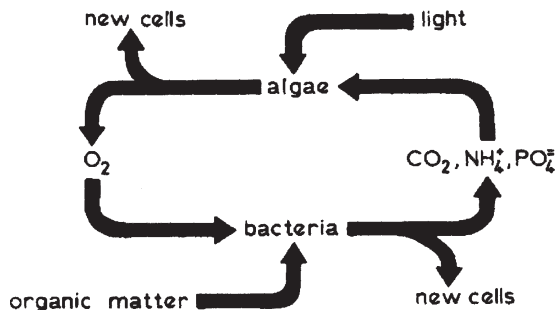


Fig. 26.2 Algal-bacterial mutualism in facultative ponds.

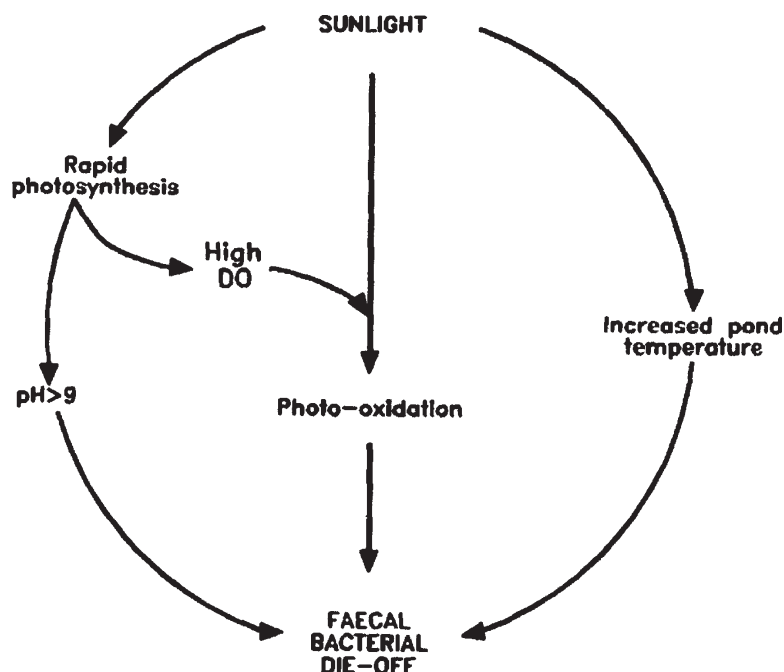


Fig. 26.3 Conceptual mechanisms for faecal bacterial die-off in facultative and maturation ponds.

WHO (1989) recommendation for unrestricted crop irrigation (see Chapter 15). However, WSP systems designed to produce effluents suitable for unrestricted irrigation generally require about twice the area of land required for those designed to produce effluents suitable for restricted irrigation. Thus decisions regarding unrestricted versus restricted irrigation should not be taken lightly and are best discussed with the local farmers.

3 WASTEWATER STORAGE AND TREATMENT RESERVOIRS

In arid or semi-arid areas where agricultural production is limited by the amount of water, including treated wastewater, available for irrigation, it is often sensible to use WSTR as these permit the whole year's wastewater to be used for irrigation, rather than just that produced during the irrigation season. WSTR are 5–15 m deep and function much like facultative ponds (Mara *et al.*, 1996). Single WSTR are used if their contents are used for restricted irrigation as, for example in Israel where WSTR were developed. Israeli practice

has been to treat the wastewater in an anaerobic pond and to discharge its effluent into a single WSTR which is 5–15 m deep (Fig. 26.4a). The irrigation season in Israel is 4–6 months long, and so the single WSTR has a storage capacity equivalent to 6–8 months' wastewater flow. It is full at the start of the irrigation season, and empty at the end of it. In this way two to three times as much land can be irrigated, and two to three times as many crops produced. Further details are given in Juanico and Shelef (1991, 1994). The long retention time in the WSTR ensures the removal of all helminth eggs, so the WSTR contents can be safely used for restricted irrigation.

For unrestricted irrigation, sequential batch-fed WSTR can be used (Mara and Pearson, 1992). This system comprises an anaerobic pond and, depending on the length of the irrigation season, three or four WSTR in parallel (but usually only three) (Mara and Pearson, 1992; Mara *et al.*, 1997) (Fig. 26.4b). Each WSTR is operated sequentially on a cycle of fill-rest-use (see Table 26.1), and faecal coliform reduction to <1000 per 100 ml occurs

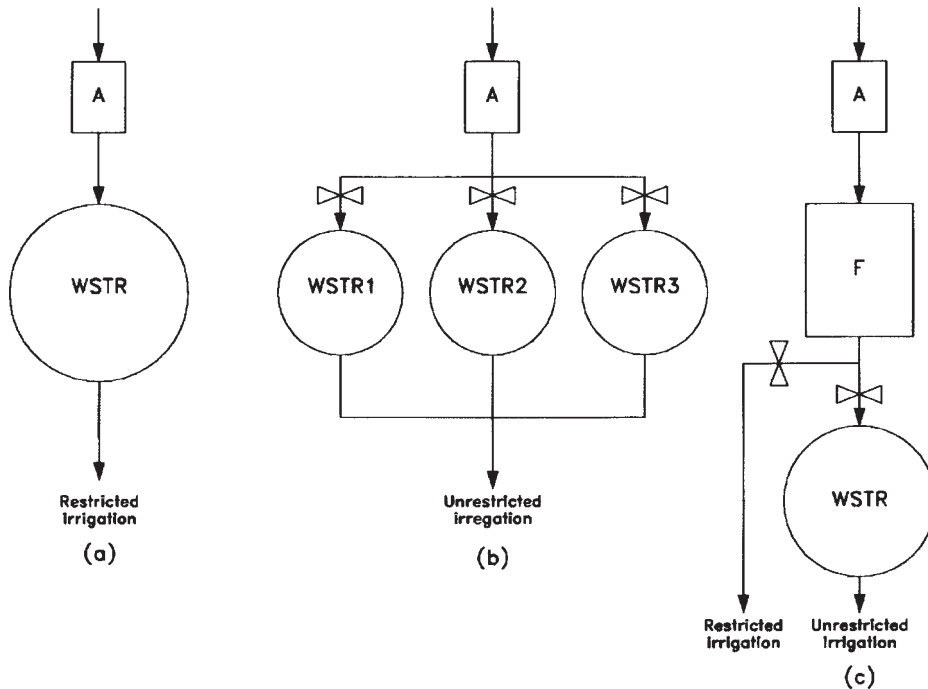


Fig. 26.4 (a) Single WSTR system for restricted irrigation, (b) sequential batch-fed WSTRs in parallel for unrestricted irrigation, and (c) hybrid WSP-WSTR system for both restricted and unrestricted irrigation. A, anaerobic pond; F, facultative pond.

rapidly during the fill and rest phases (Pearson *et al.*, 1996b). Thus the whole year's wastewater is available for unrestricted irrigation during the irrigation season.

WSTR design is detailed in Mara (1997) and Mara and Pearson (1998). Mara and Pearson (1999) describe a hybrid WSP–WSTR system to produce effluents for both restricted and unrestricted irrigation (Fig. 26.4c).

4 CONSTRUCTED WETLANDS

Constructed wetlands generally referred to as rooted macrophyte ponds, e.g. reedbeds, although the term can be used to describe an ecologically more diverse pond system with both rooted and floating macrophytes. The latter is often more to provide a pleasant recreational aquatic habitat (see EPA, 1993; Gearheart, 1996), rather than being specifically for wastewater treatment. This section will therefore address only reedbeds.

4.1 Reedbeds

Usually following septic tanks, anaerobic ponds or rotating biological contactors, reedbeds are long thin beds of reeds (*Phragmites australis*) planted either in the soil or in a gravel bed (alternatively, or in addition, other plants such as bulrushes (*Typha latifolia*) or rushes (*Scirpus lacustris*) can be used) (Fig. 26.5). If planted in the soil, the bed usually has an open water surface and this promotes the breeding of mosquitoes, both culicines (the larvae of which obtain their oxygen by breathing through the water surface) and *Mansonia* spp. (the larvae of which bore into the stem of the plant to obtain air from the air lacunae which transport air from the leaves to the roots). For this reason soil reedbeds are generally not much favoured, especially in developing countries where culicine mosquitoes are the vector of Bancroftian filariasis.

Gravel-bed reedbeds are commonly horizontal flow reactors with the water surface

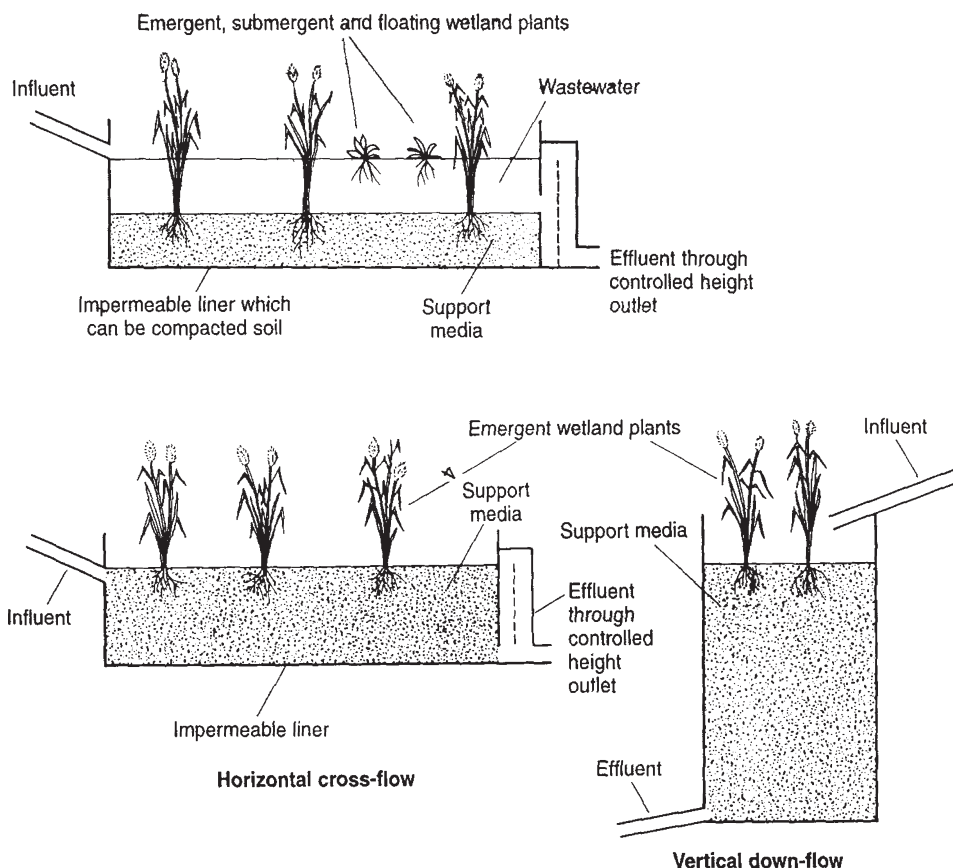


Fig. 26.5 Surface-flow soil-based reedbed with free water surface (top) and horizontal- and vertical-flow gravel-bed reedbeds (bottom) (Nuttal *et al.*, 1997).

below the top of the gravel bed to avoid mosquito breeding. Bed depth is 0.6 m, varying from 0.3 m at the inlet to 0.9 m at the outlet. Gravel size is 5–15 mm. The reeds (or other rooted macrophytes) (4–8 plants/m²) help to oxygenate the bed, and heterotrophic bacterial activity within it reduces the BOD by around 90% (Williams *et al.*, 1995; Cooper *et al.*, 1996; Nuttal *et al.*, 1997). Bed size varies according to application: 3.7–5 m²/person for secondary treatment, 1–2 m²/person for tertiary treatment and 0.3–0.5 m²/person for stormwater treatment (Nuttal *et al.*, 1997).

Vertical-flow reedbeds are used at higher organic loadings than horizontal flow beds, often where space is at a premium. Bed depth is 0.5–1.5 m. An area of around 1 m²/person

is required for 90% BOD removal and 2 m²/person for 70% ammonia removal (Nuttal *et al.*, 1997).

Cooper (2001) should be consulted for an up-to-date review of reedbed design.

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Microbial interactions in facultative and maturation ponds

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1 INTRODUCTION

The key component that sets facultative and maturation ponds apart from other wastewater treatment systems in terms of process microbiology is the presence of microalgae. These microorganisms basically control treatment efficiency and effluent quality and ponds must be designed to optimize both the concentration and species diversity of algae present. This factor is frequently overlooked by sanitary engineers accustomed to designing other types of treatment facility and it is the main cause of poor pond design. Thus an understanding of the dynamics of algal–bacterial interactions in ponds is fundamental both to the design and efficient operation of such systems. Even so, correctly designed pond systems are highly robust and easy to maintain.

2 ALGAL BIOMASS AND ALGAL DIVERSITY

It has been estimated that approximately 80% of the dissolved oxygen in waste stabilization ponds (WSP) results from the photosynthetic activity of the microalgae forming the phytoplankton population and thus the aeration of ponds depends heavily on algal activity rather than on surface reaeration. The maintenance of a healthy algal population is therefore fundamental to the efficient oxidation of organic material by the bacterial population.

This relationship between the phototrophic microalgae and the aerobic chemo-organotrophic bacteria is often illustrated as a mutualistic relationship (see Chapter 26, Fig. 26.2) in which the bacteria benefit from the oxygen produced by algal photosynthesis to metabolize aerobically organic material for growth and energy production. The algae benefit by utilizing the carbon dioxide produced by bacterial respiration along with the released nutrients (N and P) to derive energy and fix carbon for growth via photosynthesis. This view of the role of the microalgae is, however, somewhat simplistic since the algae are now known to be strongly implicated in the natural disinfection processes in maturation ponds that are responsible for destroying viral and bacterial pathogens as will be discussed later in this chapter.

The algal genera and species that predominate in a pond appear to be a function of the surface organic loading the pond receives and, in general, at higher organic loadings species diversity decreases (Konig, 1984; König *et al.*, 1987). Thus facultative ponds have fewer algal genera than maturation ponds and flagellate genera tend to predominate, whereas in maturation ponds diversity is greater and non-flagellates frequently predominate (Table 27.1). This fact serves as a quick on-site means of determining whether the ponds in a series are underloaded or overloaded as the dominant algal species can be quickly identified with a simple field microscope. Algal biomass decreases with increased BOD surface loadings

TABLE 27.1 Examples of algal genera present in facultative and maturation ponds

Algal genus	Facultative ponds	Maturation ponds
Euglenophyta		
<i>Euglena</i>	+	+
<i>Phacus</i>	+	+
Chlorophyta		
<i>Chlamydomonas</i>	+	+
<i>Chlorogonium</i>	+	+
<i>Eudorina</i>	+	+
<i>Pandorina</i>	+	+
<i>Pyrobotrys</i>	+	+
<i>Ankistrodesmus</i>	–	+
<i>Chlorella</i>	+	+
<i>Micractinium</i>	–	+
<i>Scenedesmus</i>	–	+
<i>Selenastrum</i>	–	+
<i>Carteria</i>	+	+
<i>Coelastrum</i>	–	+
<i>Dictyosphaerium</i>	–	+
<i>Oocystis</i>	–	+
Cryptophyta		
<i>Rhodomonas</i>	–	+
<i>Volvox</i>	+	–
Chrysophyta		
<i>Navicula</i>	+	+
<i>Cyclotella</i>	–	+
Cyanobacteria ^a		
<i>Oscillatoria</i>	+	+
<i>Arthrospira</i>	+	+

+ = present; – = absent.

^a photosynthetic prokaryotes.

in facultative ponds (Fig. 27.1) and the impact appears greater in shallow ponds than in deeper ones. At high organic loads the algal

population tends towards monoculture of flagellate genera with *Chlamydomonas* proving to be the most resistant.

The recent work of Athayde (2001), studying algal dynamics in various series of waste stabilization ponds operating at different surface organic loadings at Extrabes in Northeast Brazil, where light intensities remain relatively constant and water temperatures are normally around 25°C, identified 28 different algal genera with the Chlorophyta dominant irrespective of organic loading. The genera *Chlamydomonas*, *Pyrobotrys*, *Phacus* and *Euglena* were among the most resistant algae to high organic loadings, being present at the highest loadings encountered of 770 kg BOD₅/ha day. In contrast, *Ankistrodesmus* and *Scenedesmus* were sensitive to high organic loadings and were most abundant at surface organic loadings below 75 and 50 kg BOD₅/ha day, respectively. High concentrations of ammonia and sulphide are associated with high organic loadings and it is these rather than the organic load (BOD) which control algal biomass and algal species dominance (Abelovich and Azov 1976; König 1984; König *et al.*, 1987; Mills 1987; Pearson *et al.*, 1987a; Athayde 2001). Since it is the non-ionic species of both ammonia (NH₃) (which is the increasingly dominant form at pH above 7) and sulphide (H₂S) (which predominates at neutral to acid conditions) that rapidly enter algal cells leading to toxicity, the prevailing pH of the pond water (which is influenced

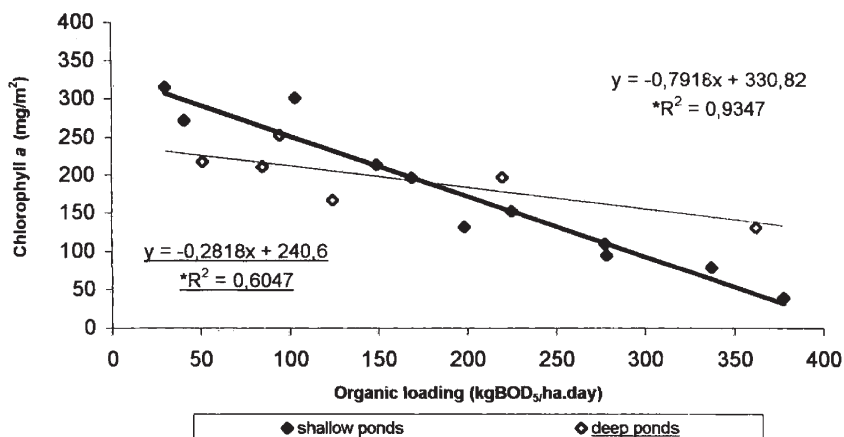


Fig. 27.1 The effect of surface organic loading on chlorophyll *a* values expressed on an area basis in shallow (1.0 m) and deep (2.20 m) pond series in Northeast Brazil (Athayde, 2001).

by algal photosynthetic activity), as well as the overall concentrations of these substances, is important.

The genus *Chlorella* is an enigma in that it is probably the most persistent genus, occurring across a very wide range of organic loadings. However, in terms of maximum biomass concentration it exhibits a preference for low loadings in the region of 28 kg BOD₅/ha day, normally found in maturation ponds. Athayde (2001) concluded that changes in the biomass (cell numbers) of *Chlamydomonas*, as the most pollution tolerant genus, and *Scenedesmus*, which favours low organic loadings, were the best indicator algae to use in monitoring changes in loading conditions in pond systems (Figs 27.2 and 27.3).

There is much still to do in terms of identifying pond algae as most studies have limited algal identification only down to the genus level. This is not surprising since, for example, there are allegedly over 200 species of *Euglena* and, according to Lund and Lund (1995), more than 400 species of *Chlamydomonas*. Only by the use of modern molecular techniques will it be possible to determine whether a species is, for example, miniaturized or modified in form due to prevailing

conditions or represents a different species of the same genus which is proliferating under the changing conditions. Such studies will help considerably in understanding the complex microbiology of the waste stabilization pond environment.

Total algal biomass as determined by chlorophyll *a* concentration is also higher in facultative ponds than in the subsequent maturation ponds of a series. This probably reflects the reduction in available nutrients and also the increased grazing pressures by the zooplankton population which is larger in the more aerobic conditions prevailing in maturation ponds.

The degree of grazing of algae by the invertebrate fauna in ponds can affect effluent quality in two ways, one being beneficial and the other detrimental. A reduction in the algal standing crop in the final maturation pond of a series by grazing will improve effluent quality in terms of BOD since the algal concentration leaving the pond will be diminished. In fact, zooplankton grazing can theoretically reduce algal BOD in the effluent by 80–90% since the energy transfer from algae to the next trophic level, the grazers, will be between 10 and 20%, with 80% energy loss via heat and detritus sedimentation.

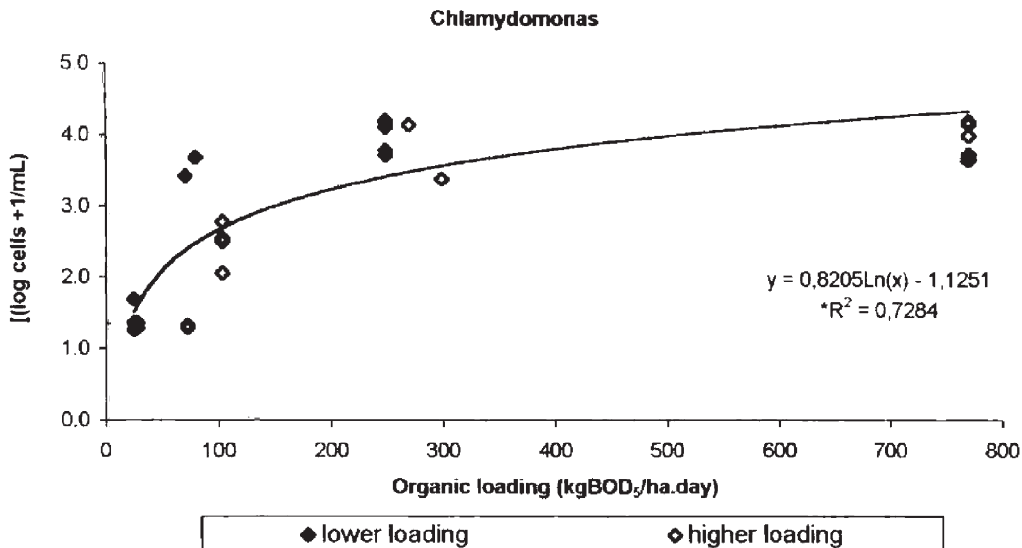


Fig. 27.2 *Chlamydomonas* biomass concentration plotted against surface organic loadings under both lower (250 kg BOD/ha day) and higher (770 kg BOD/ha day) loadings applied to the secondary facultative pond of the 5 pond series which included an anaerobic pretreatment pond (Athayde, 2001).

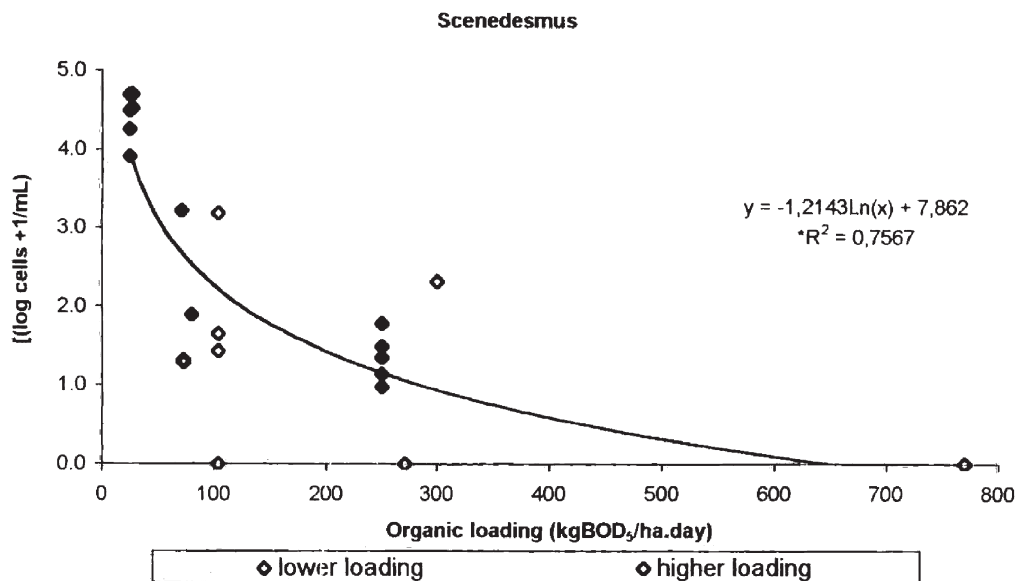


Fig. 27.3 *Scenedesmus* biomass concentration plotted against surface organic loadings under both lower (250 kg BOD/ha day) and higher (770 kg BOD/ha day) applied to the secondary facultative pond of the 5 pond series which included an anaerobic pretreatment pond (Athayde, 2001).

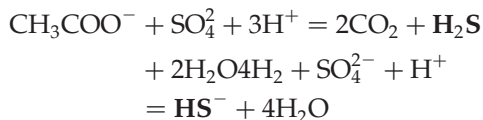
In contrast, excessive zooplankton grazing can reduce the effectiveness of the natural disinfection process which is reliant upon an active algal population to produce the high oxygen concentrations and elevated pH which are fundamental to the overall process (see *Pathogen removal* below).

Several algal genera common in facultative ponds, including *Euglena*, are capable of heterotrophic growth on organic substrates such as acetate released as a result of anaerobic degradation of organic material in the anaerobic sediments (Mills 1987; Pearson *et al.*, 1987a). *Euglena* species have also been observed migrating to the lower anaerobic organic-rich sediments in facultative ponds at night (Konig, 1984). However, the significance of this in terms of overall organic carbon removal in ponds is not clear.

3 SULPHATE-REDUCING BACTERIA AND PHOTOSYNTHETIC SULPHUR BACTERIA

The production of H₂S and thus the risk of bad odour in ponds is the result of the activity of sulphate-reducing bacteria (SRB) of, for

example, the genera *Desulfovibrio* and *Desulfobacter*. These are obligately anaerobic bacteria and are present in the anaerobic layer and sediments of the facultative ponds (as well as in anaerobic ponds). They require organic material (e.g. organic acids) or hydrogen as a source of reductant, and sulphate (or sulphur or sulphite) as the terminal electron acceptor to re-oxidize their electron transport chains under anoxic conditions during the production of energy (ATP) required for growth. Examples of equations for H₂S production by SRB are as follows:



Thus both excess sulphate and excess organic material (organic overloading) will stimulate the growth and activity of SRB. Acidic conditions (pH < 6) or alkaline conditions (pH > 8) will favour SRB proliferation over pH-sensitive methane-producing bacteria, which are also found in the anaerobic sediments since they

compete for similar organic substrates. This results in more H_2S production and greater (or complete) inhibition of methane production.

The photosynthetic purple and green sulphur bacteria utilize hydrogen sulphide (H_2S), generated by SRB activity, as an electron donor for CO_2 reduction in photosynthesis. The sulphide is oxidized to elemental sulphur (S^0) and the sulphur is stored within the cells or, more accurately, in the periplasm (the space between the outer membrane and cytoplasmic membrane) of the cell in the case of the purple forms, but is deposited outside the cells in the case of the greens. If no H_2S is available, the purple forms utilize the stored elemental sulphur oxidizing it to sulphate. They can also utilize other reduced sulphur compounds such as thiosulphate and sulphite. Unlike the cyanobacteria and microalgae they do not utilize water as an electron donor and thus do not liberate oxygen as a by-product of photosynthesis. This non-oxygen-evolving photosynthetic process is termed anoxygenic photosynthesis, as opposed to oxygen-evolving or oxygenic photosynthesis.

The purple sulphur bacteria (e.g. the genus *Thiopedia*) are common members of the microbial flora of properly functioning facultative lagoons and are normally located in the water column at a depth of approximately 50 cm in the anoxic but illuminated zone below the surface microalgal layer. They do not compete for light with the algae as they utilize wavelengths longer than 800 nm, which pass unabsorbed through the algal zone. They normally obtain H_2S by diffusion of the gas through the water column from the lagoon sediments where it has been produced by the activity of sulphate-reducing bacteria as part of the normal anaerobic activity occurring in the base of facultative lagoons. In fact, under normal conditions in ponds, purple and green sulphur bacteria are important components of the natural odour filtration system as they oxidize a proportion of the H_2S before it reaches the aerobic surface waters of the pond where the photosynthetically produced oxygen completes the process. They also protect the algae from photosynthetic inhibition by sulphide (Houghton and Mara, 1992).

On occasions facultative (and indeed maturation lagoons) turn purple as a result of an increase in the purple bacterial population to the exclusion of the algal population. This usually results from the presence of high H_2S concentrations generated by SRB activity in the sediments sufficient to support the photosynthetic requirements of a large population of photosynthetic bacteria and to remove all the dissolved oxygen from the pond water column so providing a larger than normal anoxic illuminated (photic) zone, in which for example, purple photosynthetic bacteria such as *Thiopedia* can develop. Concentrations of H_2S greater than $\approx 8 \text{ mg/l}$ are known to inhibit oxygenic photosynthesis, leading to the death of many pond microalgae (Mills, 1987; Pearson *et al.*, 1987a), so further exacerbating the presence of anaerobic conditions. However, it is interesting to note that in Morocco, WSP systems close to the sea with purple ponds continue to function well in terms of BOD removal. These ponds were dominated by *Thiopedia* as this genus is favoured over other photosynthetic purple sulphur bacteria in illuminated anoxic saline environments. Thus purple lagoons are often an indicator not only of organic overloading or high sulphate concentrations, but also of saline intrusion into coastal sewers.

4 HETEROTROPHIC BACTERIAL POPULATION IN WSP

A wide range of aerobic heterotrophic bacterial genera was found in ponds by Gann *et al.* (1968), including *Pseudomonas*, *Achromobacter*, *Flavobacterium* and *Bacillus*, although much less is known of their activities than those of the photosynthetic organisms. The bacteria present in the aerobic conditions of facultative ponds are deemed to be essentially those saprophytes present in the incoming wastewater and also include *Beggiatoa*, *Sphaerotilus*, and *Alcaligenes*, in addition to those already mentioned (Environmental Protection Agency, 1983). In other words, most aquatic bacterial groups are present and it is assumed that the microbial degradation of organic matter in ponds is

similar to that in other biological wastewater treatment systems, although the biomass concentrations are presumably lower and thus account for the longer hydraulic retention times required for effective treatment.

5 PATHOGEN REMOVAL AND THE NATURAL BIOLOGICAL DISINFECTION PROCESS IN WSP

In comparison with other secondary wastewater treatment processes, ponds have been known to be very effective at removing most types of helminths, protozoa, bacteria and viruses for some years. Indeed, Caldwell (1946) reported on the excellent removal of coliforms in WSP, and Marais produced an engineering design equation for coliform removal in WSP based on time and temperature. It assumes first order kinetics for bacterial removal with the first order removal coefficient based on air temperature and that the ponds are completely mixed (Marais and Shaw, 1961; Marais, 1966, 1974). Several workers have criticized the concept that WSP are completely mixed reactors as demanded by the Marais design approach.

Dissanayake 1980 (Polprasert *et al.*, 1983), assuming first order removal kinetics for bacterial die-off, developed a dispersed flow model which corresponds more closely with the hydraulic regime found in ponds (Mara, 1976; Marecos do Monte and Mara, 1987). Yanez (1984) and Saenz (1992) also favoured the inclusion of a dispersion number in their equations to explain better bacterial removal in WSP and suggested dispersion values based on basic pond geometry. While such an approach may well better explain the kinetics of bacterial removal in ponds, and in this context is of value, it does little to aid pond design directly since a range of variables affects the value of the hydraulic dispersion number of a pond. These include pond shape, the number and positioning of inlets and outlets, shear stresses at the sides and the bottom, wind speed and direction, and thus basically it can only be determined with any accuracy by the use of tracers once the pond has been constructed

(Polprasert *et al.*, 1983; Marecos do Monte and Mara, 1987; Vorkas and Lloyd, 2000). Indeed Shilton and Harrison (2002) suggest that inlet design (including baffles) is a key factor in determining the hydraulic regime in pond systems, and in comparison the influence of outlet design and the relative significance of the wind can be considered to be only of secondary importance.

Despite the criticisms made of the Marais design approach in terms of pond hydraulics, it nevertheless seems to work well in practice in terms of predicting the effluent quality of a WSP series. Pearson *et al.* (1995), studying various series of experimental ponds in Northeast Brazil where mean air and water temperatures vary little throughout the year, calculated first order bacterial removal constants (K_b) for faecal coliforms using the Marais equations and found that in the five facultative ponds studied the K_b values ranged between 2.5 and 4.0/day and were thus lower than the theoretical value of 6.2/day for a temperature of 25°C. In contrast, the tertiary maturation ponds had elevated K_b values in the range 10.0–13.8/day. Despite these discrepancies, when actual log numbers of faecal coliforms in individual pond effluents in a series were plotted against the theoretical effluent values that could be predicted from raw wastewater values using the Marais design approach, linear regressions with highly significant positive correlations ($r^2 = 0.99$) were obtained (Fig. 27.4). Thus, while the Marais approach may have its limitations, it nevertheless works well when there are three or more ponds in series since over- and under-estimations of K_b balance out, giving effluent values very close to predicted values using the Marais pond design equations. Furthermore, there was a good negative correlation between total hydraulic retention time (HRT) and final effluent concentrations of faecal coliforms in the five experimental pond series at depths of 1.0 m and in the deeper 2.2 m studied by Athayde (2001) in Northeast Brazil (Fig. 27.5).

Brissaud *et al.* (2000) found, in tracer tests using sodium iodide on a maturation lagoon in the South of France to determine residence time distribution and the simultaneous determination of the die-off constant in pilot lagoons

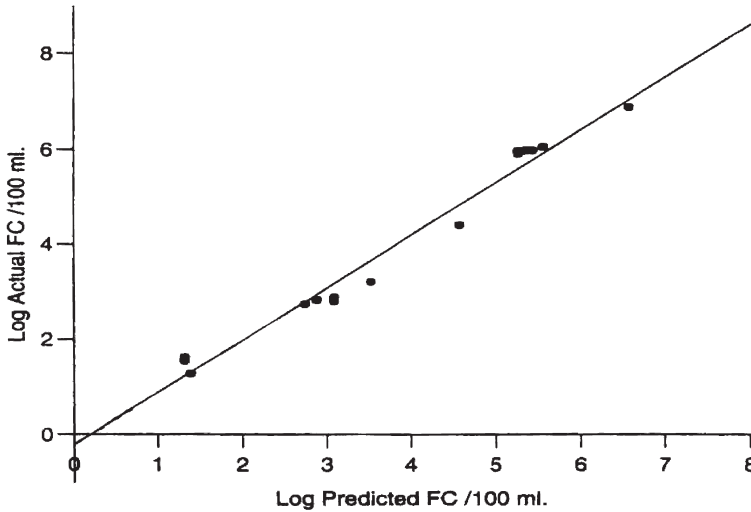


Fig. 27.4 Comparison of log actual and log predicted faecal coliform (FC) numbers in individual pond effluents. Log predicted values were derived using the Marais equation for determining effluent concentrations in a series of ponds and the FC concentration of the raw wastewater.

assuming a first order faecal coliform decay in a completely mixed reactor, that the Marais equations gave a good prediction of the actual faecal coliform numbers in the effluent. The pond in general mixed quickly throughout most of its depth. They thus verified that faecal coliform removal could be predicted from observed residence time assuming first-order faecal coliform die-off constant in a completely mixed reactor.

Despite these useful engineering studies on pond kinetics, the underlying mechanisms involved in the natural disinfection process have only become apparent relatively recently. It was Parhad and Rao (1974) who, on observing high pH values in ponds in India, investigated the role of microalgae on *E. coli* die-off. When algal photosynthesis raised the

pH above 9.4, *E. coli* were removed very rapidly. Buffering the flasks containing *E. coli* and the algal cultures to keep the pH near neutral prevented die-off, and conversely raising the pH above 9.4 with alkali in the absence of algae also caused rapid bacterial die-off. They therefore concluded that high pH conditions brought about by rapid algal photosynthesis were the key mechanism for killing bacterial pathogens in WSP. Pearson *et al.* (1987b) also reported the rapid die-off of various faecal coliform populations isolated from WSP in Portugal at pH values above 8.6. The rate of die-off under laboratory conditions was independent of both light and oxygen concentration (up to 100% saturation), but die-off accelerated with increasing temperatures in the range 10–30°C (however, it should be

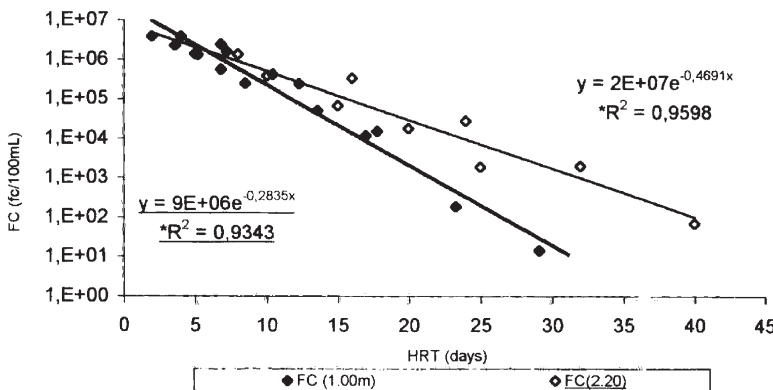
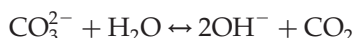
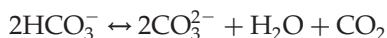


Fig. 27.5 Faecal coliform concentration plotted against the hydraulic retention time (HRT) in shallow (1.0 m deep) and deep pond series (2.2 m deep) (Athayde, 2001).

mentioned that only relatively low light intensities were used from 'Growlux' fluorescent tubes). The effects were independent of the presence of algae and they also concluded that pH was a key factor in the die-off of bacterial pathogens. They also noted that pH values and die-off were generally higher in maturation ponds than in facultative ponds.

The rise in pH is a direct result of rapid photosynthetic carbon fixation by the algae, which removes dissolved CO₂ from the pond water more rapidly than it can be replaced by either bacterial respiration or from the air across the pond-air interface. This results in a shift in the carbonate-bicarbonate equilibrium to produce CO₂ and hydroxyl ions. This CO₂ is fixed by the microalgae and hydroxyl ions accumulate with a consequent rise in pH:



This process can lead to pH values of above 10, especially in maturation ponds.

Smallman (1986) and Mills (1987) working together placed samples of pond water in dialysis bags and suspended them in various maturation and facultative ponds at different depths in the water column and found that when the daytime dissolved oxygen concentration (DO) was less than 20 mg/l and the pH was 8.5 or less the faecal coliform removal rate was roughly uniform at all depths. However, when the pH exceeded 9.0 and DO exceeded 20 mg/l the removal rate was highest at the surface. In further studies on the effluent quality of two maturation ponds in series, one, in which *Daphnia* had consumed the bulk of the algal population and thus had good light penetration but low pH and DO, had an effluent faecal coliform concentration that was higher than that of the other maturation pond which did not have a grazing *Daphnia* population and thus had a high algal concentration, lower light penetration and high pH and oxygen concentrations.

Curtis (1990) concluded that, while light did not seem to be directly responsible for faecal coliform die-off in ponds, high pH was also not an explanation for all situations in which

good faecal coliform die-off rates occurred. Trousellier *et al.* (1986), working on ponds in the south of France, correlated faecal coliform die-off with three parameters, namely light, pH and chlorophyll *a* and provided the first clear evidence that more than a single parameter might be responsible for faecal coliform removal in ponds. Curtis *et al.* (1992a,b), in a series of laboratory field experiments, established that light between 425 and 700 nm (i.e. photosynthetically active radiation) killed faecal coliforms in ponds principally through an oxygen-mediated exogenous photosensitization process. The light is absorbed by dissolved humic substances in the pond water and these then enter an excited state for long enough to damage the bacterial cells. Light-mediated faecal coliform die-off was completely dependent on the presence of oxygen and is greatly enhanced by high pH levels (both of which are products of algal photosynthesis). Curtis *et al.* (1992b) have summarized conceptual mechanisms for faecal coliform die-off in WSP (See Chapter 26, Fig. 26.3) and concluded that any models explaining the bactericidal activity of light in ponds should include dissolved oxygen concentration and pH.

Recently Davies-Colley *et al.* (2000), in a series of experiments using WSP effluent in small, stirred reactors showed that different faecal indicators were inactivated by different components of the solar spectrum, and the rates of sunlight inactivation had differing dependencies on physicochemical conditions. For example, F-specific DNA phage was inactivated only by solar UVB (300–320 nm) at a rate unaffected by other factors, whereas enterococci and F-specific RNA phage were inactivated by a wide range of wavelengths (300–550 nm) by DO-dependent photo-oxidation. Sunlight inactivation of faecal coliforms was particularly complicated: at pH < 8.5 only solar UVB (300–320 nm) caused (slow) inactivation, but at higher pH, the inactivation rate increased and a wider range of wavelengths (300–550 nm) contributed, suggesting photo-oxidation damage to membranes, which sensitizes faecal coliforms to high external pH.

In general therefore it seems likely that the biological disinfection process implicating

light, pH and oxygen is a key mechanism controlling most faecal bacterial and at least some viral removal in ponds, rather than the mechanisms of starvation and predation. A notable bacterial exception appears to be *Vibrio cholerae* O1 which is tolerant of high pH but seems more vulnerable to sulphide inhibition and was more rapidly removed in anaerobic ponds (Arridge *et al.*, 1995). There is therefore a good reason for including anaerobic ponds in the treatment system where cholera is common. Also it would seem that RNA viruses may be less vulnerable to the photo-oxidative process, which accounts for why Oragui *et al.* (1995) found rotavirus removal to be slow in ponds.

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Sulphate-reducing bacteria

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1 INTRODUCTION

The corrosion of sewers and the control of odour are the major operational and maintenance problems in wastewater collection and treatment systems. Significant capital investment is required for extensive sewer rehabilitation as reflected by the fact that the corrosion rate is about 5 mm/year in certain sections of the sewers in Japan (Mori *et al.*, 1992) and Los Angeles County (EPA, 1991). The public demand for an odour-free environment presents one of the major problems in the wastewater field.

The odour problems result from the generation of hydrogen sulphide (H_2S) by a specialized group of anaerobes, collectively called sulphate-reducing bacteria (SRB). Different sulphide species (H_2S , HS^- and S^{2-}) coexist in equilibrium in wastewater and the distribution of each species is a function of pH. For instance, approximately 45% of the total sulphide exists as $\text{H}_2\text{S}_{\text{aq}}$ at pH 7. There is a further equilibrium between aqueous and gaseous H_2S governed by Henry's law; and the escape of $\text{H}_2\text{S}_{\text{gas}}$ depends on the degree of mixing, and surrounding ambient $\text{H}_2\text{S}_{\text{gas}}$ concentration. $\text{H}_2\text{S}_{\text{gas}}$ exhibits a typical rotten-egg smell.

The corrosion problem results from the subsequent biological process in which sulphide is converted to sulphuric acid by aerobic autotrophic bacteria (sulphide oxidizing bacteria, SOB). The H_2SO_4 produced reacts with the various materials in sewer systems, resulting in eventual sewer corrosion. Additionally, H_2S itself, under anaerobic conditions, also causes the so-called depolarization of iron,

hence the corrosion of iron (e.g. Bryant *et al.*, 1991). The formation of different copper sulphides (e.g. CuS and Cu_2S) is further indicative of SRB-induced corrosion of copper and copper alloys (McNeil *et al.*, 1991).

The practical control of corrosion and odour problems essentially involves the prevention of sulphide generation induced by SRB. Obviously, an adequate flow velocity in sewers may prevent sediment deposits which provide the ideal habitat for SRB. Also, adequate dissolved oxygen (DO) in wastewater can inhibit growth of the strictly anaerobic SRB; a few SRB species, however, can survive under aerobic conditions as discussed later. These practices for controlling odour problems, however, may be limited by the sewer system and treatment plant related constraints.

Normally, heavy metals present in the wastewater above certain threshold levels can be toxic to microorganisms, including SRB, thus restricting SRB growth and sulphide generation. Conversely, heavy metals readily react with sulphide to form metal sulphide precipitates, resulting in a lower sulphide concentration and reduced metal toxicity. For example, Dezham *et al.* (1988) used iron salts to form FeS in anaerobic digesters, thereby reducing the available sulphide, to control the H_2S present in the digester gas for both corrosion and air pollution control purposes.

Although the SRB and SOB cause some environmental problems of odour and corrosion, the activities of these bacteria may also alleviate some environmental problems in nature and in man-made engineered systems. Specifically, SRB can minimize metal mobility and toxicity because of insoluble metal

sulphide precipitates. A potential widespread utilization of SRB in removing heavy metals is envisioned for the future. Furthermore, as laboratory experiments are transformed into pilot and full scale works, the microbial transformation of oxyanions such as As(V)/As(III), Se(VI)/Se(IV) and Cr(VI) by SRB may provide a cost-effective means in bioremediation of these hazardous oxyanions.

This chapter presents pertinent information about the microbiology of SRB in general and a brief review of utilizing SRB for removing heavy metals and reducing metal oxyanions in particular.

2 SULPHUR CYCLE

The biological sulphur cycle, similar to the biological nitrogen cycle described in chapter 25, is driven by microbial activities. The sulphate ion (SO_4^{2-}), the most common sulphur species present in wastewater, is biologically reduced to sulphide. H_2S is the most common and prevalent odourous inorganic substance associated with wastewater collection and treatment systems. The threshold odour concentration is extremely low (0.0005 ppm, Leonardos *et al.*, 1969). Adverse physiological effects to humans can occur at concentrations greater than 10 ppm. Finally, H_2S may reach life-threatening levels at concentrations greater than 300 ppm (Mori *et al.*, 1992). The odour of H_2S is so strong that it is readily perceived in the environment long before the toxic level is reached. Unfortunately, human deaths due to sulphide toxicity are not uncommon, especially for sewage workers and others working in enclosed systems where H_2S might build up (Brock and Madigan, 1991). In sewer systems, $\text{H}_2\text{S}_{\text{gas}}$ concentration ranges from 3 to an extremely high 400 ppm near a manhole (Mori *et al.*, 1992). Ambient H_2S levels at a wastewater treatment plant were found to be higher (>1 ppm) around the inlet structure and primary clarifiers (Koe, 1985). The conditions leading to H_2S formation generally favour the production of other malodourous organic compounds. Thus, solving H_2S odour problems can often solve other odour problems as well.

Microbial corrosion of materials is the result of oxidative reactions initiated by the metabolic activity of SOB, of which *Thiobacillus thiooxidans* is the predominant species. The corrosive action of H_2SO_4 results in damage to the concrete walls of reactors and sewer systems as well as steel pipelines. (Refer to Section 4 for biologically mediated corrosion problems in water distribution and treatment systems.)

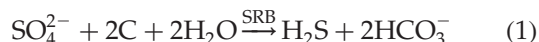
Although SOB are aerobic and SRB are anaerobic microbes, the presence of SRB in aerobic environments as well as that of SOB within the anaerobic microbial community is well documented.

3 SULPHATE-REDUCING BACTERIA

A relatively wide range of genera of SRB has been identified. The major characteristics of the presently classified SRB are summarized by Widdel (1988). All SRB are Gram-negative except *Desulfonema* species. The two most prevalent genera of SRB are *Desulfotomaculum*, comprised of spore-forming, straight or curved rods, and the non-spore-forming genus *Desulfovibrio* with curved motile vibrios or rods (Gibson, 1990).

3.1 Metabolism

SRB are obligate anaerobes which obtain energy for growth by oxidation of organic substrates and use sulphate as the terminal electron acceptor as:



Equation 1 is similar to nitrate reduction in that alkalinity is produced and there is a stoichiometric relationship between mg carbon required per mg SO_4^{2-} reduced. However, the sulphate reducers, unlike the denitrifiers, which are facultative organisms and prefer an aerobic environment, can only use sulphate in the absence of oxygen or nitrate.

3.1.1 Carbon source

The preferred carbon sources for SRB are low-molecular-weight compounds such as organic

acids (e.g. lactate, pyruvate, formate and malate), fatty acid (e.g. acetate), and alcohols (e.g. ethanol, propanol, methanol and butanol). Nearly all of these compounds are fermentation products from anaerobic degradation of carbohydrates, proteins and lipids. Thus, SRB are terminal degraders and their role is analogous to that of methanogenic bacteria that produce methane and CO₂ as final products. The sulphide production rate in sewers has been correlated with dissolved carbohydrate and volatile fatty acids (Nielsen *et al.*, 1998). The well known *Desulfovibrio* genus does not utilize acetate, i.e. only degrading lactate to acetate (commonly referred to as incomplete SRB), while different *Desulfotomaculum* species are acetate-utilizing bacteria (Postgate, 1984; Widdel, 1988; Brock and Madigan, 1991).

A few species can even grow autotrophically using CO₂ as a sole source of carbon. Some strains of *Desulfotomaculum* can utilize glucose, but this is rather rare among sulphate reducers in general. Degradation of aromatic and saturated cyclic organic compounds, as well as long-chain n-alkanes, has also been reported with some sulphate reducers (Widdel, 1988; Caldwell *et al.*, 1998; Häggblom, 1998). SRB metabolic reactions and free energy changes with different carbon substrates are shown in Table 28.1.

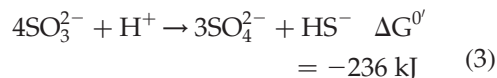
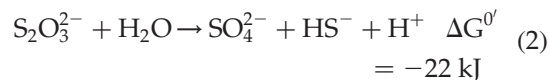
TABLE 28.1 Metabolic reactions and free energy changes for SRB (Hao *et al.*, 1996)

Reactions	Free energy (kJ/mol)
3LA ⁻ → 2PA ⁻ + HA ⁻ + HCO ₄ ⁻ + H ⁺	-165
2LA ⁻ + SO ₄ ²⁻ + H ⁺ → 2HA ⁻ + 2CO ₂ + HS ⁻ + 2H ₂ O	-189
4PA ⁻ + 3SO ₄ ²⁻ → 4HA ⁻ + 4HCO ₄ ⁻ + 3HS ⁻ + H ⁺	-151
HA ⁻ + SO ₄ ²⁻ → 2HCO ₄ ⁻ + HS ⁻	-60
HA ⁻ + 4S + 3H ₂ O → 4H ⁺ + HCO ₄ ⁻ + 4HS ⁻ + CO ₂	-24
4H ₂ + SO ₄ ²⁻ + CO ₂ → 3H ₂ O + HS ⁻ + HCO ₄ ⁻	-152

Note: LA⁻, CH₃COOHCOO⁻ (lactate); PA⁻, CH₃CH₂COO⁻ (propionate); HA⁻, CH₃COO⁻ (acetate).

3.1.2 Sulphur source

In the presence of organic electron donors and thiosulphate or sulphite, SRB may first break down the thiosulphate or sulphite compounds to sulphate (equations 2 and 3) and then oxidize the organic substrates with the newly formed sulphate (Widdel, 1988):



These disproportionation reactions can provide energy for cell synthesis (Bak and Pfennig, 1987; Brock and Madigan, 1991). In fact, some SRB prefer thiosulphate to sulphate. The sulphide production rates from both thiosulphate and sulphite were found to be considerably higher than those from sulphate (Nielsen, 1991). It is assumed that sulphate would not be a limiting factor for sulphide production in most wastewaters at concentrations greater than 20 mg/l (EPA, 1974).

3.2 Environmental factors

A knowledge of the various factors affecting SRB growth is essential to control their microbial reactions in wastewater environments.

3.2.1 Dissolved oxygen

Although sulphate-reducers are obligate anaerobes, they may survive a temporary exposure to oxygen and again become active under anaerobic conditions. Because of the O₂ mass transfer, the critical DO concentration in the wastewater below which sulphate reduction can occur is 0.1–1.0 mg/l (EPA, 1985). If aerobic sediments or waters are rich in organic particles, SRB may be active in anaerobic microniches despite the aerobic surroundings (Widdel, 1988). Several SRB strains, including *Desulfovibrio* and *Desulfobacter* species, could be enriched from aerobic activated sludge. A few strains of SRB have been reported to be able to utilize oxygen without growth (Dilling and Cypionka, 1990). Sulphate reduction occurs

consistently within the oxygenated photosynthetic microbial mats (Canfield and Des Marais, 1991; Frund and Cohen, 1992). In general, DO concentrations above certain levels inhibit SRB; thus, maintenance of relatively high DO levels is recommended in the design and operation of sewer and treatment systems to prevent odour problems.

3.2.2 Temperature

Sulphate reduction rates depend strongly on temperature; sulphate reduction typically increases 2- to 3.9-fold with a temperature increase of 10°C (Nielsen, 1987; Widdel, 1988). The optimum temperature for most pure cultures of SRB ranges between 28 and 32°C; the lower optima among SRB are observed with some *Desulfobacterium* strains and a curved *Desulfobacter* strain at 24–28°C and the highest is around 70°C for *Thermodesulfobacterium commune*. Most species of SRB die rapidly at temperatures above 45°C (Postgate, 1984; Widdel, 1988).

Okabe and Characklis (1992) reported that the maximum specific growth rate (μ_{\max}) of *Desulfovibrio desulphuricans* varied between 0.38 and 0.55/h at 25°C and 43°C, respectively, and the odours dramatically decreased outside this temperature range. A temperature shock of 45°C imposed in a mesophilic (30°C) upflow sludge bed system treating sulphate-containing wastewater did not show any detrimental effect; temperature shocks to 55 and 65°C, however, significantly reduced the treatment efficiency (Visser *et al.*, 1993).

3.2.3 pH

SRB prefer an environment around pH 7 and are usually inhibited at pH values below 5.5 or above 9 (EPA, 1974). Nevertheless, sulphate reduction has been observed in a peat bog and acid mine water that exhibit pH values of about 3 to 4. SRB isolated from these habitats, however, were inhibited below pH 6. Therefore, it was hypothesized that the SRB in acidic environments were present in microniches, where higher and more favourable pH conditions could exist. Recent studies further

demonstrated that two isolated strains of SRB could even grow at pH < 4.5 (Hard *et al.*, 1997), and SRB are capable of sulphate reduction and production of alkalinity at pH values as low as 3.3 (Elliott *et al.*, 1998). The effect of low pH could also be minimized in sulphate-reducing systems, since the sulphate-reduction process generates additional bicarbonate alkalinity.

A *Desulfovibrio* culture was inoculated with lactate and sulphate in the pH range of 5.8–7.0 and both the growth yield coefficient (Y) and μ_{\max} increased with pH; maximum values of 0.15 g cell/g SO_4^{2-} and 0.14/h were obtained at pH 6.6 (Reis *et al.*, 1992). Changing waste pH with acid or alkali has been suggested as a method for diminishing sulphate reduction in industrial plants.

3.2.4 Sulphide

Sulphide is known to inhibit SRB, a classic example of the product inhibition often encountered in chemical and biological processes. Sulphide inhibits SRB probably when anyone of the sulphide species (H_2S , HS^- or S^{2-}) combines with the iron of ferredoxin, cytochromes or other essential iron-containing compounds in the cell, causing the electron transport systems to cease activity (Okabe *et al.*, 1992). However, literature data with respect to inhibitory concentrations of H_2S or total sulphide vary due to different environmental conditions (e.g. pH, temperature), wastewater characteristics (e.g. presence of iron salts and type of carbon source) and reactor systems (e.g. fixed film).

3.3 Ecology

SRB are widespread in sewers and wastewater treatment plants. In sewer systems, SRB are normally present in slime layers (EPA, 1985; Holder, 1986) and sediments (Mori *et al.*, 1991). Sulphate reduction rates increase proportionally with the increase in the SS (suspended solids) concentration; e.g. an increase from 250 to 1500 mg/l SS results in a sixfold increased rate (Attal *et al.*, 1992). Because of their slow

TABLE 28.2 SRB count in sewers, wastewater and sludge samples

Sample type	MPN count (per 100 ml unless otherwise noted)	References
Raw wastewater	$> 2 \times 10^8$	Lish, 1993
Primary effluent	10^7	Morton <i>et al.</i> , 1991
Aerobic fixed film (trickling filter and rotating biological contactor)	$6 \times 10^7 - 10^8$ (per g VS)	Lens <i>et al.</i> , 1995
Activated sludge (aerobic)	$2 \times 10^5 - 10^6$ (per g VS)	Lish, 1993; Yamamoto-Ikemoto <i>et al.</i> , 1994; Lens <i>et al.</i> , 1995
Activated sludge (anoxic)	7×10^5	Lish, 1993
UASB (upflow anaerobic sludge blanket) sludge	$3 \times 10^7 - 5 \times 10^{10}$ (per g VSS)	Harada <i>et al.</i> , 1994; Lens <i>et al.</i> , 1995; Fang <i>et al.</i> , 1997
Anaerobic digester sludge	$6 \times 10^5 - 5 \times 10^7$	Postgate, 1963; Lish, 1993; Harada <i>et al.</i> , 1994
Acidogenic anaerobic sludge	up to 10^{11}	Mizuno <i>et al.</i> , 1998
Enriched sludge for sulphate reduction	10^8	Lish, 1993
Pure SRB culture	up to 8×10^{10}	Postgate, 1963
Stream	6×10^2	Postgate, 1963
Sediment in sewer	$3 \times 10^4 - 10^8$ (per g VSS)	Mori <i>et al.</i> , 1991; Schmitt and Seyfried, 1992
Landfill	8×10^6 (per g)	Fairweather and Barlaz, 1998

growth characteristics and carbon preference, SRB are usually outnumbered by other types of microbes, except in special environments. In anaerobic sludge digesters, there exists a competition between SRB and MPB (methane-producing bacteria), since these two groups of bacteria exhibit a similar affinity for organic substrates. In general, factors such as concentration of sulphide, type of fermented organics, COD/SO₄²⁻ ratios, and type and concentration of other inhibitory substances affect the relative prevalence of the bacterial types.

Despite their strict anaerobic requirement, the presence of SRB has been detected in many aerobic regions (Gibson, 1990), including activated sludge and aerobic fixed film. Table 28.2 presents the results of SRB counts in various environmental samples. The differences in SRB number are obvious, even with the same type of samples. Many factors can contribute to the differences, e.g. depth of sediment samples (a three-order of magnitude difference (7×10^4 to 3×10^7 /g) in sewer trunk sediments (Mori *et al.*, 1991), type of substrate (e.g. benzoate or H₂) used, and system objective (e.g. maximum of MPB) in various SRB studies.

4 METAL REMOVAL

The role of sulphide in preventing heavy metal toxicity under anaerobic conditions has long been recognized (Lawrence and McCarty, 1965). The insoluble metal sulphide precipitates (Table 28.3), due to sulphide generated *in situ*, are responsible for immobilization of heavy metals in natural environments of soils and sediments and in engineering systems as:

**TABLE 28.3** Solubility product of metal sulphide and metal hydroxide precipitates (Zumdahl, 1989)

Metal	K _{sp}	
	Metal sulphide	Metal hydroxide
Cu	8.5×10^{-45}	1.6×10^{-19}
Zn	1.2×10^{-23}	8.5×10^{-17}
Pb	3.4×10^{-28}	1.2×10^{-15}
Cd	3.5×10^{-29}	5.9×10^{-15}
Fe	3.7×10^{-19}	1.8×10^{-15}
Ni	1.6×10^{-16}	1.6×10^{-16}
Cr	—	6.7×10^{-31}

As mentioned earlier, the external addition of iron salts may be used to precipitate sulphide for odour control (Sercombe, 1995), reduce metal/sulphide toxicity (Edwards *et al.*, 1997), reduce H₂S gas in anaerobic digesters (Dezham *et al.*, 1988), and control dissolved sulphide in sewers (Padival *et al.*, 1995). In fact, Jin *et al.* (1998) even added sulphide into a methanogenic system to reduce the effect of copper toxicity. In wastewater, if iron salts are present in large quantities, FeS precipitation normally occurs. The black colour of sediments as well as primary and digested sludge, where sulphate reduction occurs, is due to the accumulation of FeS.

The use of SRB for metal removal has been applied to different industrial wastes, including contaminated groundwater (Barnes *et al.*, 1991), landfill leachate (DeWalle *et al.*, 1979), plating wastes (Bewtra *et al.*, 1995), leachate from contaminated soils (White *et al.*, 1998) and mining effluent (Maree *et al.*, 1987), in a variety of processes including anaerobic filters, sludge blanket reactors, and contact systems. The different metal removal schemes using SRB are shown in Fig. 28.1. Essentially, metal-laden wastes are supplemented with organics and other nutrients to provide adequate sulphate/COD ratios, and other conditions are used, e.g. lower hydraulic detention times and neutral pH, to favour the growth of SRB. Once sulphide is generated *in situ*, it readily reacts with metals to form MeS precipitates. In general, high efficiencies of metal removal can be achieved. For example, an upflow sludge blanket reactor (9 m³) was installed near a zinc refining site in the Netherlands to treat metal-contaminated ground water (Barnes *et al.*, 1991). With external addition of nutrients (N and P) and carbon/energy substrate (ethanol), the system reduced Zn concentrations of 107–1070 mg/l to below 1 mg/l.

Haas and Polprasert (1993) used the recycling stream containing bio-generated sulphide to remove Cu, similar to the scheme shown in Fig. 28.1c. The system produced an effluent with consistently low COD and metal content. Hammack *et al.* (1994) used a similar system (shown in Fig. 28.1d) to treat mine

waste, with a limestone neutralization reactor between the metal precipitator and sulphate reduction reactor. The countercurrent metal precipitator removed metal and the limestone reactor increased pH to 6. Sodium lactate was injected directly into the sulphate reduction reactor. The process removed more than 99% of the initial concentrations of Fe (620 mg/l), Cu (178 mg/l), Zn (530 mg/l), and Al (278 mg/l).

The potential use of SRB for pollution abatement of acid mine drainage (AMD) has long been recognized (Tuttle *et al.*, 1969; Wakao *et al.*, 1979). Since the 1980s, laboratory, pilot-scale, and full-scale evaluations have demonstrated the capability of using SRB to remove sulphate and metals from AMD (Table 28.4). In general, the amount and type of organics added, as well as pH adjustment, are critical for metal removal. Most metals are precipitated as MeS, although some are retained as hydroxides or carbonates.

Because sulphate reduction may be significant in artificial or natural wetlands, there has been some keen interest in using wetlands for the treatment of the AMD (e.g. Webb *et al.*, 1998), due to low cost and low-maintenance requirements. A survey conducted in 1989 indicated that a total of 142 wetlands in the eastern USA were constructed specifically for AMD treatment (Wieder, 1989). However, the effectiveness of these wetland treatments of AMD has been variable and unpredictable (Wieder, 1989).

4.1 Metal toxicity

Toxicity studies for SRB are often reported as a function of the initial metal concentrations. Because of complicated biological processes, some metals may precipitate and/or physically entrap within the biomass. Thus, the residual soluble metal concentrations should be determined to account for the observed toxicity phenomenon. Furthermore, metals present in free form, and not complexed forms due to naturally and biologically generated surfactants, exhibit a toxic effect; and different forms of metals, e.g. As(V) versus As(III), may exhibit

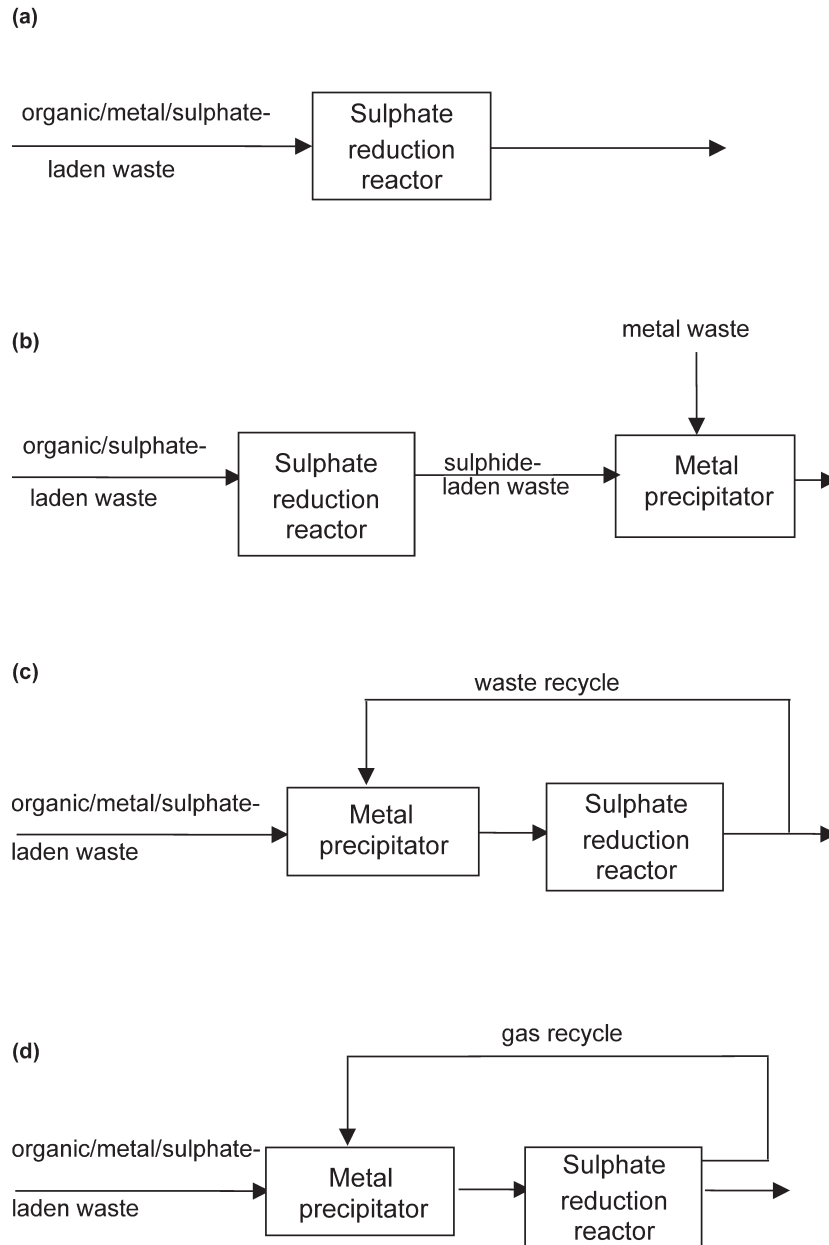


Fig. 28.1 Potential schemes for removal of heavy metals by sulphate-reducing bacteria

different effects. Consequently, highly soluble metal concentrations may not exert inhibitory effects in anaerobic sulphate-reducing systems which generate biosurfactants rendering metals less toxic.

The correlation between metal toxicity and sulphide content in sediments has been extensively studied (e.g. Di Toro *et al.*, 1992)

in terms of the ratio of simultaneously extracted metal (SEM) to the acid volatile sulphide (AVS). Typically, sediment/sludge samples with SEM/AVS ratios of metals (e.g. Cd) greater than one may be toxic to the test organism, whereas samples with ratios less than one are not toxic, due to insoluble MeS.

TABLE 28.4 Treatment of acid mine drainage by SRB

Type of mine wastes	Systems	Efficiency	References
Ni simulated waste supplemented with lactate and sulphate	Columns with mushroom compost	540 mg Ni/kg compost-day	Hammack and Edenborn, 1992
Coal mine dump and smelting residues	3–200 l reactors in series, filled with spent mushroom compost and a single 4500 l reactor	95% removal efficiencies for Al, Cd, Fe, Mn, and Zn	Dvorak <i>et al.</i> , 1992
Copper-zinc mine waste supplemented with whey	Batch treatment in cylinders for 200 days	Significant removal of Cu, Fe, Al and Zn; pH increased and redox decreased	Christensen <i>et al.</i> , 1996
Mine contaminated groundwater	Reactive wall (40% municipal compost, 50% leaf compost and 20% wood chips) installed in the path of waste	Fe decreased from 250–1300 to 10–40 mg/l; pH increased from 5.8 to 7; alkalinity from 50 up to 2000 mg/l	Benner <i>et al.</i> , 1997

5 METALLOID OXYANION REDUCTION

Many strains of SRB are capable of reducing the soluble oxidized oxyanions (e.g. MoO_4^{2-}) to insoluble phases (e.g. MoS_2 or MoO_2). Growth of these microbes may occur with these oxyanions as the sole electron acceptors (Lovley, 1993; Newman *et al.*, 1997; Tebo and

Obraztsova, 1998), although some SRB strains cannot grow with oxyanions as sole electron acceptors (Lovley and Phillips, 1994). It is generally established that the enzymatically mediated reactions are responsible for oxyanion reduction, e.g. both hydrogenase and cytochrome *c* were responsible for Cr(VI) reduction (Lovley and Phillips, 1994) and c_3 cytochrome for the U(VI) reduction (Lovley

TABLE 28.5 Reduction of metalloid oxyanions by SRB strains

Soluble oxyanions	Insoluble species	Comments	SRB species	References
U(VI)	U(IV), $(\text{UCa})\text{O}_2$	From 235 to less than 2 mg U/l	<i>Shewanella putrefaciens</i>	Abdelouas <i>et al.</i> , 1998
	Mo(IV) $\text{MoS}_2(\text{s})$	Either lactate or H_2 as electron donor; $\text{MoS}_2(\text{s})$ formed in the presence of sulphide	<i>Desulfovibrio desulphuricans</i> , <i>Desulfovibrio vulgaris</i>	Tucker <i>et al.</i> , 1997
MoO_4^{2-} , SeO_4^{2-} , $\text{UO}_2(\text{CO}_3)_2^{2-}$, $\text{UO}_2(\text{CH}_3\text{COO})_2$, Cr(VI)	MoS_2 , UO_2 , Se^0	High removal efficiency in an immobilized system; formate or lactate as electron donor	<i>Desulfovibrio desulphuricans</i>	Tucker <i>et al.</i> , 1998a, b
SeO_4^{2-} or SeO_4^{2-}	Se^0	Formate as electron donor, sulphate or fumarate as electron acceptor	<i>Desulfovibrio desulphuricans</i>	Tomei <i>et al.</i> , 1995
U(VI)	U(IV)	Half saturation coefficient = 127 mg/l, yield coefficient = 0.02 g/mol pyruvate	<i>Desulfovibrio desulphuricans</i>	Tucker <i>et al.</i> , 1996
U(VI), Cr(VI), Mo(VI)	U(IV), Cr(III), Mo(IV),	First-order rate constants for U(VI), Cr(VI), and Mo(VI) = 0.8, 0.15, and 0.02/h, respectively	<i>Desulfovibrio desulphuricans</i>	Lovley and Phillips, 1992; Fude <i>et al.</i> , 1994; Tucker <i>et al.</i> , 1997

et al., 1993). The role of sulphide (Fude *et al.*, 1994) and, to a lesser extent, SRB adsorptive capability (Mohagheghi *et al.*, 1985) in oxyanion reduction has, however, been attributed in removing these oxidized oxyanions.

A summary of the reduction of metalloid oxyanions by several SRB strains is shown in Table 28.5.

6 CLOSING REMARKS

The SRB directly and indirectly cause many environmental problems, including odour and sewer corrosion. However, the activities of SRB may also alleviate some environmental problems in nature and in man-made engineered systems, specifically for the minimization of metal mobility and removal of heavy metals because of insoluble metal sulphide precipitates. Furthermore, future usage of SRB for microbial transformation of hazardous oxyanions such as As(V)/As(III), Se(VI)/Se(IV), Cr(VI), Mo(VI), and U(VI) may establish SRB as one of the most important microbial species in bioremediation.

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Behaviour of Pathogens in Wastewater Treatment Processes

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Viruses in faeces

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1 INTRODUCTION

Many viruses are excreted by humans and animals. Some of these are present in very large numbers but cannot be grown, others can only be grown with difficulty in cell cultures. Enteroviruses, hepatitis A virus, rotaviruses, parvovirus-like viruses, astroviruses, caliciviruses, adenoviruses and coronaviruses may be present in human excreta. Rotaviruses are the commonest cause of acute non-bacterial gastroenteritis in infancy and childhood. In children aged 6 months to 10 years, infection by rotaviruses occurs more frequently during winter months in temperate countries. In tropical countries, there is considerable variation in the incidence of rotavirus infections among children.

Enteroviruses include the polio, coxsackie A and B and echoviruses. These belong to the family of Picornaviridae. They contain single-stranded RNA. Poliomyelitis is rare in developed countries, as a result of the widespread use of oral polio vaccines, but continues to circulate in developing countries. Similarly, hepatitis A is an RNA virus and has the physicochemical characteristics of a typical enterovirus. It is transmitted from person to person by the faeco-oral route. Outbreaks may originate from viral contamination of food, water, milk and shellfish. Rotaviruses and parvovirus-like viruses cause gastroenteritis. The role of adenoviruses, astroviruses, caliciviruses and coronaviruses is not well-established. However, it is now recognized that these viruses can cause outbreaks of acute

diarrhoeal disease. Enteric adenoviruses represent a separate serotype distinct from adenoviruses associated with acute respiratory infection and have been associated with acute diarrhoeal disease.

Besides parvovirus-like viruses, other small rounded viruses have been detected in faecal samples from patients with acute diarrhoeal disease. These include:

1. astroviruses, encountered in outbreaks in children's wards (Kurtz *et al.*, 1977), stools of lamb with diarrhoea and calf faeces
2. caliciviruses, detected in stools in children in winter and staff with diarrhoea and also asymptomatic children (McSwiggan *et al.*, 1978)
3. coronaviruses, which cause severe diarrhoea in pigs and calves; there is no conclusive evidence that these viruses cause diarrhoea in humans.

2 VIRUSES IN WASTEWATER

The types and numbers of human enteric viruses in raw wastewater depend on the origin and nature of the wastewater. Enteroviruses, including hepatitis A virus, adenovirus and rotaviruses, are frequently present in domestic wastewater. These viruses reflect those circulating in the community and depend on a number of factors, which include population, climate, seasonal and diurnal fluctuations and the presence of chemical effluents. Raw wastewater contains viral numbers which vary widely from country to country. For example,

in raw wastewater in Israel, higher numbers are generally present than in the USA, although the numbers reported depend on the detection and concentration methods used.

Animal viruses may also be present in raw wastewater from the faeces of household pets and effluents from farms, abattoirs and stormwater. The isolation of animal viruses from wastewater, especially abattoir wastes, has been reported by several workers. However, more emphasis has been on human viruses due to their potential for causing human diseases.

Viruses may be categorized or divided into three main groups:

1. viruses associated with humans
2. viruses associated with higher plants and animals
3. viruses associated with the microbial flora (bacteriophages)

Over a hundred different types of viruses are known (WHO, 1979). Excreta is the commonest source, but by no means the only source. Nose and throat secretions also contribute some viruses to wastewater, and viruses from skin lesions and blood (which may find its way through baths and toilets during acute stages of infection giving rise to viraemia) can also be found in wastewater. These sources, although they do not contribute much, may in certain situations be important.

Viruses associated with plants and animals may be important for economic reasons, especially in countries where the reuse of wastewater for crop irrigation is practised. Some of these viruses come from the wastewater itself, whereas others may enter the sewer via wastewater from farms, slaughterhouses, food-processing factories and stormwater. Faecal materials from birds, dogs and other animals will be washed from roofs and roads by rain and find their way ultimately into the wastewater.

The third group comprises the bacteriophages, which are viruses that parasitize bacteria and, because of the huge variety of bacteria in wastewater, which may all have their own phages, phage numbers in wastewater can be very high.

3 VIRUS RECOVERY FROM WASTEWATERS

One of the major problems encountered in the enumeration of viruses in water and wastewater before and, particularly, after treatment is the large quantity of sample that has to be examined in order to detect and count them. This is made even more difficult by the fact that the enumeration technique usually employed for rotavirus assays utilizes only 25–200 µl of sample. For enterovirus assays, 0.1–1 ml can usually be assayed conveniently. The consequent dilution of viruses in faeces when discharged into receiving waters or sewage means the sample must be concentrated in order to analyse them. Various techniques are available for concentrating viruses (see Oragui and Mara, 1996).

One of the difficulties encountered with concentrated wastewater samples is the toxic effects that they have on the cell lines used for virus propagation and enumeration. Concentrates of raw wastewater and, in particular those obtained from anaerobic waste stabilization ponds, are very toxic to the cells used for rotavirus assay. Toxic effects are indicated by the destruction or detachment of cells during rotavirus assay by the indirect immunofluorescence technique. A simple method for the detoxification of concentrate has been developed and is described by Oragui and Mara (1989).

4 VIRUS REMOVAL AND SURVIVAL IN WASTEWATER TREATMENT

Conventional wastewater treatment plants are not specifically designed to reduce the number of excreted pathogens, including excreted viruses. However, there is usually some incidental removal of viruses – primary sedimentation ranges from 30 to 65% (Berg, 1973; Rao *et al.*, 1977); in activated sludge from 80 to >90% (Clarke *et al.*, 1961; Malina, 1976); and in trickling filters from 10 to 20% (Sherman *et al.*, 1975), although Berg (1973) reported removals of 15–100%. Disinfection processes, principally chlorination and ozonation, for the removal of faecal bacteria, have not been

TABLE 29.1 Ranges and geometric means of faecal coliforms and excreted rotaviruses in raw wastewater and pond effluents, September–December 1988

Sample source	No. of samples	Faecal coliforms (per 100 ml)	Rotaviruses (per litre)	Percentage removal	
				Faecal coliforms	Rotavirus
Raw wastewater	12	3.5×10^7 – 7.8×10^4 (6.12×10^4)	1.06×10^4 – 2.66×10^5 (1.13×10^4)	–	–
Anaerobic pond	12	2.1×10^6 – 4.2×10^6 (3.13×10^6)	1.02×10^3 – 7.27×10^4 (5.87×10^3)	94.9	94.8
Facultative pond	12	1.1×10^5 – 1.4×10^6 (8.91×10^5)	8.5×10^2 – 3.2×10^4 (1.41×10^3)	71.5	76.0
1 st Maturation pond	12	8.5×10^4 – 2.75×10^5 (1.50×10^6)	1.06×10^2 – 6.6×10^3 (1.87×10^2)	83.2	86.7
2 nd Maturation pond	12	2.5×10^4 – 5.85×10^3 (4.37×10^4)	0.9×10^2 – 6.6×10^2 (0.234×10^2)	70.8	87.5
3 rd Maturation pond	11	5.3×10^2 – 2.75×10^3 (1.41×10^3)	0.09×10^2 (3.9)	96.8	83.3

Retention time: 5 days in each pond.

Geometric means are given in parentheses.

Source: Oragui and Mara (1996)

widely adopted other than in a few countries, notably the USA, Israel and South Africa, because of the detrimental effects on the natural fauna of receiving waters and the production and discharge of toxic and carcinogenic chlorination by-products.

In a study on rotavirus removal in a series of five 3 m deep waste stabilization ponds in northeast Brazil, Oragui and Mara (1996) found four log unit removals of both rotaviruses and faecal coliform bacteria (Table 29.1). The highest rotavirus removal was found in the anaerobic pond (95%), with lower removal in the facultative and maturation ponds (76–88%).

4.1 Factors affecting rotavirus removal in ponds

Several factors may explain viral inactivation in waste stabilization ponds, including solar radiation, temperature, pH, adsorption onto solids, heavy metals, algal and bacterial activity, and the action of certain chemicals, notably ammonia and sulphide.

The unionized ammonia (dissolved NH_3 gas) is toxic to bacteria (Pearson *et al.*, 1987), with toxicity increasing with increasing pH. The effect of ammonia toxicity on rotavirus inactivation has also been shown to be pH dependent (Oragui and Mara, 1996): at pH 9.0 there was a reduction of just over half a log unit at all ammonia concentrations (20–80 mg N/l), whereas at pH of 6.9, the log reduction ranged

from 1.3 to 2.2. In contrast to the effect of ammonia on bacteria, ammonia is more toxic to rotaviruses at pH 6.9, when it is essentially all present as the ammonium ion, than at pH 9.

Sulphide is one of the main toxicants present in raw wastewater and anaerobic pond effluents. Sulphide was found to exert the most effect on simian rotaviruses at pH 6.9, with log reductions of 0.1–3.9; at pH 8 reduction was 0.04–0.09, but at pH 9 it was 0.03–0.46. In contrast to ammonia, increasing sulphide concentrations in the range 4–16 mg S/l increased the degree of virus inactivation.

In 2 m deep facultative ponds in-pond rotavirus numbers decreased principally in the top 1 m of the pond, with little difference found in rotavirus numbers in the 1–2 m lower layer. This indicates that, as the surface layers experience algae-induced pH changes, the toxic effects of ammonia and sulphide are likely to be the main mechanism of removal of rotaviruses, and possibly of all excreted virus, although adsorption on to the algae and subsequent settlement when the algae die may be an important secondary mechanism.

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Bacterial pathogen removal in wastewater treatment plants

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1 INTRODUCTION

No widespread outbreak of enteric bacterial disease would be complete without a call for improved wastewater treatment (McCabe, 1970; Nichols 1991, Tauxe *et al.*, 1994). These statements were presumably made in the sincere belief that wastewater treatment systems are generally good at removing pathogenic bacteria and that this makes a significant contribution to public health in the richer countries. Unfortunately, pathogen removal is poor in most wastewater treatment plants. Thankfully, this usually has little impact on public health in most countries as other measures (notably water treatment and supply) protect us from the pathogenic bacteria we excrete.

However, where pathogen removal is important it can be very important indeed. For example, wastewater reuse in agriculture and aquaculture will be vital to balancing the water demand and supply in the 21st century. This will mean removing bacterial pathogens before the water is used for food production. We already know that reuse without treatment can lead to significant excess disease (Shuval *et al.*, 1986; Swerdlow *et al.*, 1992) and (as in Peru in the early 1990s) to significant economic losses. Economic issues may also be an important factor in compliance with recreational water treatment standards. The costs of recreational water quality standards may be borne by relatively small coastal communities who may be ill-equipped to pay for large centralized disinfection schemes.

2 PATHOGEN RANGE AND LOAD DICTATE THE HAZARD AND RISK IN UNTREATED WASTEWATER

The numbers and range of pathogens found in wastewater are significant considerations. The range of bacterial pathogens is important because it dictates the nature of the hazard posed by the wastewater. On the other hand, the numbers of pathogens have a direct bearing on the risk those pathogens pose and the magnitude of the task of reducing that risk.

A knowledge of the range of pathogens also tells us something of the ecology of the organism we are trying to remove (Table 30.1). This table will probably be incomplete within a few years as, at present, it appears that significant new enteric bacterial pathogens are being discovered every decade or so. Some, such as *Helicobacter pylori*, which can cause ulcers, pose unexpected hazards. Others such as the highly infectious *Escherichia coli* 0157 are extraordinarily virulent agents with familiar symptoms (Collier *et al.*, 1998).

The wastewater of any significant community can be assumed to have virtually all the pathogens normally found in excreta at any one time. Even in the wealthiest countries the annual incidence of infectious intestinal disease may exceed 20% (Wheeler *et al.*, 1999). Organisms such as *Vibrio cholerae* O1 or O139, which cause cholera, may be present in non-endemic areas in carriers and the mildly ill can make a substantial contribution to the contamination of wastewater without coming to the attention of the medical authorities (Curtis,

TABLE 30.1 The reported concentrations of bacterial pathogens in untreated domestic wastewater

<i>Organism</i>	<i>Disease</i>	<i>Symptoms</i>	<i>Reported concentrations/ 100 ml wastewater</i>	<i>References</i>
Genus <i>Campylobacter</i>	Campylobacter enteritis	Diarrhoea	70–1630	Oragui <i>et al.</i> , 1987; Stampi <i>et al.</i> , 1992
Enterohaemorrhagic <i>E. coli</i>	Haemorrhagic colitis	Diarrhoea, cramp, loss of kidney function	Ubiquitous but no quantitative reports	Muniesa and Jofre, 2000
Other enterovirulent <i>E. coli</i>	Diarrhoea and bacillary dysentery	Diarrhoea and mild dysentery	Detected in wastewater but no quantitative reports	Ohno <i>et al.</i> , 1997
<i>Helicobacter pylori</i>		Achlorhydria stomach ulcers	No quantitative reports	Morton and Bardhan, 2000
<i>Salmonella typhi</i>	Typhoid	Septicaemia and fever	No quantitative reports	Shuval <i>et al.</i> , 1986
<i>Salmonella enteritidis</i>	Salmonellosis	Diarrhoea	20– >1800	Langeland, 1982; Oragui <i>et al.</i> , 1987; Yaziz and Lloyd, 1979
Genus <i>Shigella</i>	Shigellosis	Diarrhoea and dysentery	No quantitative reports	
<i>Vibrio cholerae</i> O1 and O130	Cholera	Diarrhoea	1–10 ⁷	Curtis, 1996

1996). In England and Wales there are typically over 50 cases of imported cholera reported to the Public Health Authority each year (Communicable Disease Surveillance Centre, 2000). Given the degree of under-reporting of enteric disease, the true number of cases might be an order of magnitude higher.

Our knowledge of the ecology of these bacteria is poor, even though they represent some of the most intensively studied organisms on the planet and the importance of them in environmental transmission is beyond doubt. This gap reflects both cultural and scientific problems. On the technical side, although the entire genome of *V. cholerae* has been sequenced (Heidelberg *et al.*, 2000), there is probably not a scientist alive who could enumerate 2000 such organisms in 100 ml of raw wastewater. The pathogenic organism would be hidden among a much larger number of virtually identical non-pathogenic types of *V. cholerae* that are themselves a numerically insignificant proportion of the bacterial diversity of the wastewater.

Thus the enumeration of pathogenic enteric bacteria is often like to trying to find a needle

in a needlestack that has been thrown into a haystack! These severe technical limitations underlie the paucity of quantitative information on many pathogens in raw sewage. Many ecological studies rely on pure cultures seeded into beakers and buckets (microcosms) from at best, the environment and at worst the laboratory. These studies tend to fall short of the highest standards in microbial ecology (Brock, 1987). Furthermore, such technical shortcomings are compounded by the general perception (with honourable exceptions) that the study of the microbial ecology of bacterial pathogens is not a prestigious area of medicine or ecology (though, in truth, infectious disease epidemiology and biological waste treatment are both branches of microbial ecology).

3 COMPARATIVE ECOLOGY OF BACTERIAL PATHOGENS

Engineers and scientists in the field tend to fall back on a small range of indicator organisms (Table 30.2) because of the difficulties of measuring bacterial pathogens in wastewater.

TABLE 30.2 Typical indicator values in untreated raw sewage

Indicator	Concentration in typical wastewater (/100 ml)	Comments	References
Faecal coliforms	10^6 – 10^7	Gram-negative	Oragui <i>et al.</i> , 1987
Faecal enterococci	10^5 – 10^6	Gram-positive	Feachem <i>et al.</i> , 1983
<i>Clostridium perfringens</i> spores	10^4 – 10^5	Gram-positive	
<i>Bifidobacteria</i>	10^7	Oxygen sensitive	

They typically employ faecal coliforms, *E. coli* or Gram-positive enterococci to monitor bacterial pathogen removal in wastewater treatment. These organisms have been borrowed from the water supply industry where they were used to indicate the presence of faecal contamination. Sewage is, of course, faecally contaminated so nothing can be inferred from the mere presence of the indicator organisms in wastewater. However, these indicator organisms occur in large numbers and are relatively easy to enumerate and are (usually) non-pathogenic. The organisms are therefore monitored on the assumption that they will act as analogues of these numerous and more difficult to count pathogenic bacteria. The term pathogen indicator has been used to describe the use of these organisms to monitor pathogens (Feachem and Mara, 1979).

Ideally, pathogen indicators should be used in situations where there is evidence that the indicator is a valid analogue of the pathogens of concern. However, because the pathogens of concern are often difficult, or impossible, to enumerate, this condition is rarely fulfilled. In the absence of such evidence some indication of the validity of the indicator may be sought by determining how the pathogen and the indicator are related or by comparing the known physiology of the pathogen and the indicator. Faecal coliforms, the most widely used indicators, are usually >95% *E. coli* in temperate and tropical wastewaters which facilitates such comparisons.

E. coli are mesophilic Gram-negative bacteria in the gamma subdivision of the proteobacteria (Woese *et al.*, 1985). They are capable of using oxygen to oxidize carbon and of growing fermentatively, can withstand some oxidative

damage and find pH values of about 7.75 most agreeable. pH values more than 1 unit above or below this value may be considered stressful.

Shigella and (obviously) enterovirulent *E. coli* are essentially subspecies to *E. coli* (Collier *et al.*, 1998) and so are most likely to find an adequate analogue in faecal coliforms. However, we cannot discount the possibility that the extra genetic material associated with pathogenicity may affect the survival of the organism in the environment. For example *E. coli* O157 retains the ability to make toxin on a lysogenic (is capable of lysing the cell) bacteriophage encoded gene (Muniesa and Jofre, 2000). The phage may be induced, and the cell lysed, by exposure to oxidative stress (e.g. sunlight). This may confer a disadvantage on this organism outside the host. *Salmonella* spp. are Gram-negative rods and members of the family Enterobacteriaceae. They are thought to have similar sensitivities to pH, oxygen and temperature to *E. coli*, to which they are relatively closely related (Feachem *et al.*, 1983).

Campylobacter and *Helicobacter pylori* are only very distant relatives of *E. coli*, being from the epsilon subgroup of the proteobacteria. Nevertheless, they are known to grow well at moderate temperatures and salinities favouring *E. coli*. However, they are both microaerophilic (only able to tolerate low oxygen concentrations) and susceptible to oxidative conditions and sunlight. In addition, *Helicobacter* are tolerant of high pH values (Bergey and Holt, 1994).

E. coli and *V. cholerae* are fundamentally different because the latter is not really a gut organism at all. Although the two organisms

have a common evolutionary ancestry, as they both fall within the gamma subdivision of the proteobacteria (Woese *et al.* 1985), they have diverged considerably from their common ancestor. Consequently, *E. coli* and *V. cholerae* are members of distinct and varied bacterial families, the Enterobacteriaceae and the Vibrionaceae (MacDonell *et al.*, 1986). Only two serotypes of *V. cholerae* (O1 and O139) are capable of causing the eponymous disease. There are well over 100 other serotypes, which are relatively harmless and ubiquitous members of the natural bacterial flora of brackish water. Indeed, the disease-causing serotypes of the organism are now considered to be aquatic organisms, which infected humans, rather than a human pathogen that occasionally entered bodies of water (Feachem, 1982; West, 1986). Not surprisingly, the organism survives better in slightly brackish water at elevated pH values and has an absolute requirement for sodium ions (Singleton *et al.*, 1982a; Miller *et al.*, 1984). There is evidence that nutrient levels typical of wastewater may permit growth in fresh water (Singleton *et al.*, 1982b; Nair *et al.*, 1988). The optimum pH for survival was found to increase with decreasing salinity. At salinities equivalent to about 0.1% NaCl the survival is best at pH values between 7.5 and 9.0 (Miller *et al.*, 1984). In addition, *V. cholerae* is thought to have specific relationships with higher organisms, some of which may be found in wastewater treatment plants, including phytoplankton (Tamplin *et al.*, 1990; Islam *et al.*, 1994), water hyacinth (Spira *et al.*, 1981) and freshwater amoebae (Thom *et al.*, 1992). *V. cholerae* are not thought to be particularly sensitive to sunlight (Lema *et al.*, 1979; MacKenzie *et al.*, 1992), a form of oxidative stress. However, it is now realized that the cholera toxin gene is encoded on a filamentous bacteriophage (Faruque *et al.*, 2000) and that the phage may be induced by sunlight. Though the phage does not lyse the cells (they bud off), it seems plausible that the budding phage will impose a metabolic burden on the cell that may place it at a disadvantage in relation to its non-pathogenic competitors in the environment.

In principle, other pathogen indicators could be compared with pathogens in the same way. However, faecal enterococci are more heterogeneous than faecal coliforms and not particularly closely related to any of the known pathogenic organisms present in wastewater. Thus, the most we can say with certainty is that faecal enterococci are different from the organisms that we hope they will mimic and that different species may be different in slightly different ways. Similarly *Clostridium perfringens* is very different from the important excreta-transmitted pathogens. Consequently, there is no *a priori* reason to suppose that either of these indicators will be good analogues of the bacterial pathogens known to be present in faeces.

Although these comparisons represent mere thought experiments in ecology, they do seem broadly to tally with existing data on the comparative survival rates of different indicators, at least in waste stabilization ponds (a high pH, oxidizing environment). The faecal coliforms appear to be adequate indicators of salmonella. Campylobacters appear to die quickly. Faecal enterococci and *Cl. perfringens* spores are indeed different and appear to be more conservative indicators (Fig. 30.1).

The range of pathogen concentration (see Table 30.1) represents snap shots of wastewaters at different times and places. In reality we know that pathogen numbers vary over the day (Yaziz and Lloyd, 1979) because of diurnal patterns in defaecation. In addition, the point prevalence for a particular disease will vary within and between communities because of a myriad of other epidemiological factors. Seasonal changes of prevalence are extremely common in bacterial diseases (Feachem *et al.*, 1983), e.g. in temperate climates faecal oral bacterial infections tend to peak in the summer. In addition, widespread outbreaks of disease can lead to significant (two to three orders of magnitude) and rapid changes in pathogen concentration.

The concentration of pathogens in wastewater may be related to point prevalence (the number of people with a disease on any given day) by equation 1:

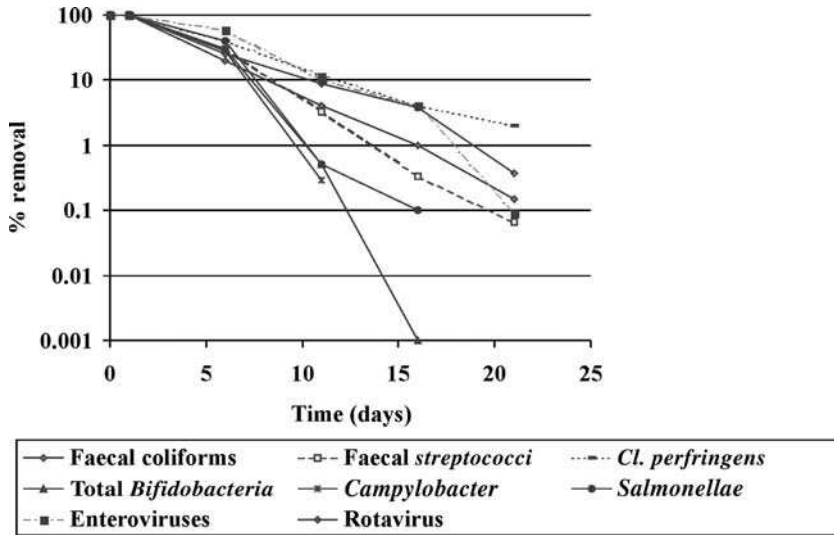


Fig. 30.1 A comparison of the removal of some enteric pathogens and typical indicator organisms in a series of deep waste stabilization ponds in the north-east of Brazil (Oragui *et al.*, 1987).

$$\begin{aligned} &\text{Pathogen/100 ml of sewage} \\ &= ((m \times f)/w \times 10) \times p \end{aligned} \quad (1)$$

where:

m is a weighted mean pathogen/g faeces
 f = mean weight of faeces/person/day
 w = litres mean water used/person/day
 p = the proportion of the population excreting the organism and 10 is a correction factor.

The value of m will depend on the severity of the illness while p reflects the extent of disease in the community.

We can use this formula to consider the likely load of *V. cholerae* on a wastewater treatment plant. The values of the variables

will depend on whether the excretors being considered are carriers or patients. For the carriers: $m = 7 \times 10^4$ organisms/g (Dizon *et al.*, 1967); $f = 100$ g/day (Feachem *et al.*, 1983) and p is unlikely to exceed 0.01 (a point prevalence of 1%), $w = 50$ litres/day. Experimental infections have suggested that most cholera patients have mild symptoms (Levine *et al.*, 1979), for such patients: $w = 10^8$ organisms/g (Dizon *et al.*, 1967), $f = 4000$ g/day (that is 4 litres of diarrhoea, assumed to have a density of 1: Levine *et al.*, 1979), $w = 200$ litres/day and p is unlikely to exceed 0.002 (a point prevalence of 0.2%). Cholera patients with severe symptoms are unlikely to use the toilet and may be discounted in the calculations.

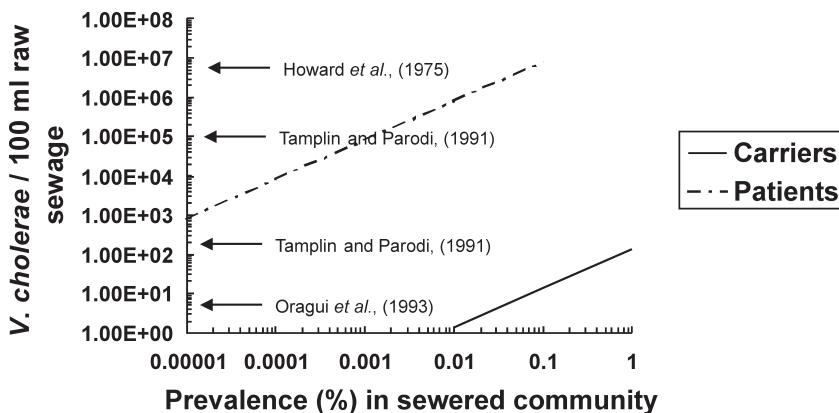


Fig. 30.2 The relationship between prevalence and estimated levels of *V. cholerae* O1 (lines) in raw sewage and reported levels of *V. cholerae* O1 in raw sewage.

These estimates are consistent with published levels of *V.cholerae* in raw sewage (Fig. 30.2). One mildly ill individual in a city of a million could contaminate the wastewater for a whole town without ever coming to the attention of the medical authorities. At the other extreme, a hospital treating cholera victims may have more toxigenic *Vibrio* than *E. coli* in the wastewater.

In certain exceptional circumstances, extremely high pathogen concentration may assume particular significance. Recreational and wastewater reuse standards assume that the pathogens occur at much lower densities than the indicator and that the reduction of the indicator by 3–4 logs results in the elimination of the bacterial pathogens. While this is nearly always true, it is possible that some breakthrough could occur in a severe outbreak.

4 THE FATE OF PATHOGENIC BACTERIA IN WASTEWATER TREATMENT PLANTS

Wastewater treatment can be divided into anaerobic systems (septic tanks, anaerobic ponds and anaerobic digesters), aerobic systems with attached growth (typically trickling filters) and aerobic suspended growth systems (typically activated sludge). A primary sedimentation tank usually precedes these systems. An increasingly popular alternative is the reed bed or wetland. Another well-established alternative technology is the waste stabilization pond (WSP) – a sort of very wetland. Both ponds and wetlands are relatively easy to construct, biologically sophisticated and contain both aerobic and anaerobic zones. Waste stabilization ponds and wetlands can be designed to optimize pathogen removal. In this sense they are different from the other wastewater treatment systems we will consider. In most wastewater treatment systems any pathogen removal that occurs is a fortuitous by-product of the principal design objective (usually organic carbon removal). Pathogen removal cannot usually be refined

in these systems without compromising this objective or at least the costs of meeting this objective.

For any given system, there are essentially two factors in pathogen removal: how long the pathogen stays in the system and how quickly it dies. The former is governed by the hydraulic flow regime and the latter depends on the ecology of the reactor.

The simplest conception of hydraulic flow in a reactor is as a completely stirred reactor in which any material entering into a system is immediately uniformly dispersed within the reaction vessel. The effluent quality of a completely stirred reactor is the same as the reactor quality and therefore some of the pathogens pass almost directly from the inlet to the outlet. However, if a number of completely mixed reactors are placed in series then they can begin to approximate plug flow reactors. In plug flow reactors the material entering into the reactor proceeds through the wastewater treatment plant without mixing with the material that has entered before or after. In reality some mixing does occur, this is known as dispersed flow.

Low rate systems such as ponds and wetlands have long retention times (typically days) and thus more opportunity to remove pathogens. However, they may have inefficient hydraulic flow regimes. Processes such as activated sludge have much lower hydraulic retention times, though some plants may be hydraulically very efficient indeed.

5 PRIMARY SEDIMENTATION TANKS

Most conventional wastewater treatment plants have some form of primary sedimentation tank to remove solids. Although the retention time of such tanks is short (typically 2–6 hours), 30–50% faecal coliform removal and 29–99% (mean 79%) salmonellae removal have been reported (Yaziz and Lloyd, 1979; Wheeler *et al.*, 1980). The salmonellae study found removal to be positively correlated with suspended solids removal which plausibly implies that the organisms are associated with

solids. Since other pathogens are also likely to be associated with solids, we may assume that faecal coliforms will be adequate indicators of bacterial pathogen removal in primary sedimentation tanks. However, indicator removal is quite variable (Feachem *et al.*, 1983) and so the lower figure should be used for design purposes.

6 ACTIVATED SLUDGE AND OTHER SUSPENDED GROWTH SYSTEMS

In suspended growth systems the biomass is formed into flocs that are typically separated from the water by sedimentation (or, more recently, membranes) allowing the retention time of the biomass to be varied independently of the hydraulic retention time. Hydraulic retention times are therefore short, typically 6–12 hours in a conventional plant, rising to 30 hours in certain designs. The opportunities for bacterial pathogen removal are therefore necessarily limited. Between 90 and 99% removal of faecal coliforms (Feachem *et al.*, 1983) and salmonellae (Yaziz and Lloyd, 1979) has been reported. Stampi *et al.* (1993) reported 99% removal of the microaerophilic *Campylobacter*. This is not very different from the removal of aerobic organisms. Clearly, there is not enough oxygen in the activated sludge significantly to prejudice the survival of this microaerophilic organism. There are no reports of the removal of *V. cholerae* in working activated sludge plants. Streeter (1930) reported that Courmont and Rochaix (1922) had seeded a batch of activated sludge and sewage with *V. cholerae* and found that they could not recover the organisms after 5–8 hours aeration. The same review stated that Bruns and Sierp (1927) had conducted analogous experiments and found that 98% of the seeded *V. cholerae* were removed. It is doubtful if even this level of removal could be achieved in practice since most suspended growth systems operate on a continuous flow basis and consequently a proportion of the influent will leave the aeration basin before the mean hydraulic retention time.

The underlying mechanisms for the removal of bacterial pathogens in activated sludge have not been rigorously investigated. However, they are likely to be related to the adsorption to solids and predation. These mechanisms should not confer a particular advantage or disadvantage on any bacterial pathogen or indicator organism. However, it is unlikely that the system can be engineered to enhance the removal mechanisms and predation-based removal is likely to be inherently variable due to the variation inherent in predator–prey systems (Case, 2000). Moreover, if solids retention is poor, pathogen removal will dip below the 90% removal suggested above. Increasing the hydraulic retention time of the system may enhance removal of the pathogens. The impact is likely to be marginal, especially in completely mixed reactors.

Membrane reactors may be very different. A side effect of the use of membranes (rather than secondary sedimentation tanks) to retain biomass is the efficient retention of virtually all bacterial pathogens. The use of membranes will impose significant capital and running costs and, at present, the technology is still relatively new. However, where the costs can be borne they may well prove to be an excellent technology for the removal of bacterial pathogens (Ueda *et al.*, 1996; Ueda and Hata, 1999). In principle, membrane bioreactors should not be effective against viruses which cause the low-grade diarrhoea associated with recreational waters and the hepatitis A occasionally spread by shellfish. A technology that just removes bacteria without removing viruses could allow an effluent to meet a health-based standard expressed in terms of an indicator while not actually protecting health. However, in practice, it appears that model scale membrane bioreactors can remove viruses, once a biofilm has built up on the membrane (Winnen *et al.*, 1996; Ueda and Horan, 2000).

7 FIXED FILM REACTORS

Fixed film reactors are also designs that allow short hydraulic retention times and long biomass retention times. Fixed film secondary

treatment processes work by passing the wastewater over a biofilm, which is capable of removing the oxygen demand. Units in which the water is distributed over a stationary media include tricking filters and biological aerated filters (though no filtration occurs), while units in which the biofilm is passed through the wastewater are called rotating biological contactors.

In trickling filters, the removal of indicator bacteria (Wheater *et al.*, 1980) and salmonellae (Yaziz and Lloyd, 1979) is poor. By contrast, Daniel and Lloyd (1980b) reported a three-log increase in the numbers of *V. cholerae* non-O1 (the non-pathogenic kind) when they passed the effluent from a dual chambered septic tank over an improvised trickling filter. The authors showed that *V. cholerae* non-O1 would reproduce in batch samples of the trickling filter effluent at 20–25°C and suggested that ponding in the filter was permitting the growth of the organism. However, when the survival of *V. cholerae* El Tor was studied in identical batches of effluent, the number of culturable organisms was observed to decline by a factor 10 over a period of 7 hours and then to maintain itself at the reduced level for a further 41 hours. The apparent discrepancy between the two forms of *V. cholerae* could not be readily explained.

Factors likely to affect the fate of bacterial pathogens in fixed film systems include binding to the biofilm, predation by protozoa and competition for nutrients. It would be difficult to engineer such a system to optimize the apparently unimpressive removal mechanisms. The short retention times characteristic of fixed film reactors could not be easily extended. It seems, therefore, that attached growth systems will never be any good at removing bacterial pathogens.

8 ANAEROBIC REACTORS

Four principal kinds of anaerobic system are used to treat domestic wastewaters anaerobically: the septic tank, the anaerobic pond, anaerobic filters and the upward flow anaerobic sludge blanket (UASB). Retention times vary

with the design and with the level of maintenance (which may be very low). However, for septic tanks and anaerobic ponds, retention times are typically a matter of 1–5 days, while for anaerobic filters and UASBs retention time may be only a matter of hours.

A well-designed and well-maintained septic tank (the minority) with a 3-day retention time might remove 50–95% of the indicator bacteria (Feachem *et al.*, 1983). Because septic tanks serve small populations, they cannot be expected reliably to contain pathogens. However, the removal of *V. cholerae* O1 and *Salmonella* (Howard *et al.*, 1975) and *V. cholerae* non-O1 (Daniel and Lloyd, 1980a) have been studied in double chambered septic tanks with a total mean hydraulic retention time of 12–15 days. The performance was relatively poor with an overall removal of 99.8% for *V. cholerae* O1 and 98.8% for salmonellae. The conditions in the tanks were inhospitable (anaerobic, with pH values of 6.3–6.7 and 40–90 mg/l of ammonia), so the poor removal may have been attributable to short-circuiting.

Anaerobic filters have been used as post-treatment systems for septic tanks. Cullimore and Viraraghavan (1994) report that indicator removal in such systems is poor. However, UASB and anaerobic ponds are being used for the domestic treatment of wastewater in a number of countries. Indeed, where the year-round ambient temperatures are high and the wastewater is reasonably strong, anaerobic treatment must be the pre-treatment method of choice. The pre-treatment contributes to pathogen removal in two ways: by removing pathogens directly; and by promoting the aerobic conditions required for removal in the subsequent post-treatment stages.

Dixo (Dixo, 1994; Dixo *et al.*, 1995) reported that a UASB reactor in northeast Brazil removed 67% of the faecal coliforms in the influent. The retention time was short (about 8 hours) and it is likely that the performance probably represents the removal of the solids-associated organisms. This mechanism is not likely to be specific to a particular organism and therefore the UASB can be expected to achieve this level of performance for bacterial

pathogens. Although, superficially disappointing, the apparently low bacterial indicator removal of the UASB must be set against the excellent BOD removal these systems can achieve. As WSP are frequently used as a post-treatment and pathogen removal in ponds is related to the organic load, a UASB can make a substantial indirect contribution to bacterial pathogen removal.

Anaerobic ponds should not have retention times of much less than a day, even in the warmest climates, to avoid the washout of the treatment bacteria. Faecal coliform removals of 80 and 95% have been reported for anaerobic ponds in northeast Brazil with retention times of 1 and 5 days respectively (Oragui *et al.*, 1987; Silva, 1982). Oragui *et al.* (1987) reported similar levels of removal of salmonellae, campylobacter, bifidobacteria and faecal enterococci. The removal observed in anaerobic ponds must, in part, reflect a one-off elimination of solids-associated organisms that would be independent of the organism or the climate and would not occur in a second identical pond in series.

9 WASTE STABILIZATION PONDS

Waste stabilization ponds are a series of one or more hypereutrophic basins, typically 1–2 m deep. Owing to the algal activity, WSP will usually have high dissolved oxygen levels and high pH values. They have very low running costs and are simple to construct, the primary expense usually being the cost of the land. The physical simplicity of WSP belies the complexity of the pathogen removal mechanisms that take place within them. For, unlike the systems we have considered above, WSP can be designed to optimize pathogen removal and the performance of two systems at the same site can vary by 3–4 orders of magnitude. Our increasing appreciation of the mechanism by which this is achieved has yet to be fully incorporated into the design equations and the design of such systems is still comparatively crude. Nevertheless, a proper understanding of these mechanisms is invaluable in the design of WSP.

High rate algal ponds (HRAP) are specially designed waste stabilization ponds in which the algal population is gently stirred mechanically, usually in some form of race-track configuration to optimize algal production. Though more difficult and expensive to operate than algal ponds, HRAP are, from a bacterial pathogens' point of view, not dissimilar to a WSP. HRAP also achieve high (sometimes higher) pH and dissolved oxygen concentrations and have correspondingly high indicator removal rates (Bahlaoui *et al.*, 1997).

Faecal coliforms are the organisms most commonly used to monitor the removal of pathogens from wastewater treatment plants. This indicator has been compared with other indicators and salmonellae, campylobacter (see Fig. 30.1) and with *V. cholerae* 01 (Curtis, 1996) and found to be adequate. The oxygen-sensitive campylobacter presumably die due to the high oxygen concentrations. The removal of *V. cholerae* 01 is slightly more mysterious, because the closely related non-pathogenic *V. cholerae* have been reported to grow in WSP (see Curtis, 1996 for a comprehensive review). It has recently been discovered that the toxin gene that distinguishes the pathogenic organisms from the non-pathogenic form is encoded by a filamentous bacteriophage that is incorporated into the organism chromosome. This phage is induced by sunlight (Faruque *et al.*, 2000). This decreases the growth rate of the organisms and might leave the toxigenic organism at a sufficient disadvantage to inhibit its survival in a WSP.¹

The mechanisms of faecal coliform removal are not surprisingly, multifactorial (Troussellier *et al.*, 1986). However, the factors can be divided into fast and slow. The fast (3–0.1 log/h) factors are photo-oxidation and pH and are associated with light and thus the upper parts of the pond during the hours of daylight.

¹ It does not appear that the engineer has much scope for increasing the inherent removal rates in wetland systems, beyond ensuring a low organic load to encourage predation. However, as with WSP, considerable improvements in the performance may be achieved by improving the hydraulic flow regime in the reed bed with consequent reduction in short-circuiting.

The slow (<0.1 log/h) factors are less well defined, and presumably include starvation, predation and adsorption to solids. The slow processes are independent of light and are thought to function at all times at all depths. Although most of research has focused on the fast factors, their relative importance depends on the depth and location of the WSP. WSP are heterogeneous in space and time. Thus the fate of a pathogenic bacterium in a WSP depends not only how long it remains in that pond, but which part of the pond it passes through and at what time of day it passes through it.

In photo-oxidation light kills an organism when the energy in the light is absorbed by a chemical or sensitizer. The sensitizer enters a short-lived excited state in which it can pass the light energy on to oxygen and form toxic forms of oxygen, including singlet oxygen, hydroxyl radicals, super-oxide radicals or hydrogen peroxide. The toxic forms of oxygen kill the cell by oxidizing vital cellular components (Curtis *et al.*, 1992).

The rate of photo-oxidation increases linearly with oxygen concentration, presumably because the increasing oxygen concentration increases the probability of the excited sensitizer meeting an oxygen molecule. In addition, the process is synergistic with pH, at least for some organisms (Davies-Colley *et al.*, 1999). The basis for the synergism is not clear; it might be because the oxygen radicals damage the membrane, thus rendering the cells sensitive to high pH values. It could also be because high pH values prolong the life of hydroxyl radicals. Since even long wavelength red light has enough energy to create singlet oxygen, any wavelength of visible or ultraviolet light can initiate photo-oxidative damage. Thus the relative importance of a given wavelength is governed by the nature of the sensitizer, the wavelengths of light involved and the relative ability of these wavelengths to penetrate into a pond. In WSP the principal sensitizer for faecal coliforms is the humic substances that occur naturally in wastewater. Other organisms may have other sensitizers, including sensitizers within the cell (Davies-Colley *et al.*, 1999).

High pH values are also thought to be responsible for some rapid removal processes.

I believe that death occurs when the bacterium is no longer able to acidify its own cytoplasm. This would explain why the relationship between removal rate and pH is non-linear, with pH values below a certain threshold value having little or no effect on bacterial indicator removal. Working with real pond water, Parhad and Rao (1974) and Curtis (1991) reported a threshold pH value of about the 9.3, while Pearson *et al.* (1987), working in buffer, report a threshold of pH 8.9. A considerable amount of energy is required for a cell to acidify its cytoplasm (Booth, 1985), thus the threshold probably represents the point at which this homeostatic mechanism is overwhelmed. By using low nutrient conditions, Pearson *et al.* (1987) artificially lowered the threshold pH value. Thus, in real ponds, very rapid pH related removal will only occur above the higher threshold.

More and better work (especially quantitative work and modelling) remains to be done on the fast processes. However, it is clear that the excellent removal seen in WSP in warm climates is intimately linked with the hyper-eutrophic conditions observed in WSP. It follows that algal concentrations really should feature in design for bacterial pathogen removal and that proposals for very clear ponds (James, 1987) would probably not work.

The light independent processes could be driven by a variety of factors, the most obvious of which are starvation, predation and adsorption to algae. Whatever the dark mechanisms are they will certainly be very important wherever pH and photo-oxidation effects are retarded by the depth or location of the pond, e.g. in deep VJSP or WSP in winter in temperate climates. Unfortunately, there has been essentially no systematic study of these factors in ponds. In an excellent paper, Troussellier *et al.* (1986) sought linear relationships between a number of interrelated factors in WSP and faecal coliform removal in the South of France. A weak negative relationship was found between algal concentrations and faecal coliforms but not between indicator organism and the BOD₅ or the rotifer concentration. However, if the effects were non-linear or relatively unimportant at this study site,

compared to the light and pH effects (which were significant), then the study would not find them to be significant.

Of course the fastest removal processes will be ineffective if the bacterial pathogens do not spend very long in the system. Conversely, even a slow rate of removal may be adequate if endured for long enough. There has always been uncertainty about the nature of the hydraulic regime in WSP. Recently, a number of authors have pointed to the importance of short-circuiting in the removal of pathogens in WSP (James, 1987; Fredrick and Lloyd, 1996; Salter *et al.*, 1999). However, it should be noted that short-circuiting is implicit in Marais' (1974) original assumption of completed mixing (some of the influent will instantaneously find itself in the effluent). Moreover, recent survey work has shown that ponds crudely approximate to complete mix reactors (Nameche and Vasel, 1998) and, for the time being, this assumption may be retained and perhaps used in conjunction with a small safety factor.

At present, in practice, the best way to reduce the impact of short-circuiting is to use more than one pond in series (Mara, 1976). Other ways of improving WSP hydrodynamics, e.g. baffles, are conceivable. However, to be usable the improvements must not prejudice other features of the WSP. For example, by promoting the build-up of solids and sulphide generation baffles in facultative WSP, baffles may cause odour and prejudice photosynthesis. In addition to being used in design, the improvements must be predictable. I hope that the current interest in the use of computational fluid dynamics will yield verified models that are sufficiently cheap to employ to justify their widespread use in the optimization of WSP design for pathogen removal.

10 WETLANDS AND REED BEDS

Wetland is a term used to describe a variety of formats of wastewater treatment systems in which wastewater is fed from, through, or over a substrate planted with reeds. Though in many ways analogous to the nineteenth

century land treatment of wastewater, wetlands have become more and more popular. Like ponds, they may be specifically designed to remove pathogenic organisms. Indeed, wetlands have been advocated for use by small communities in temperate climates trying to meet pathogen or pathogen indicator standards (Cooper *et al.*, 1996).

There have been a number of reports of pathogen removal in wetlands in various formats (Coombes and Colett, 1995; Cooper *et al.*, 1996; Green *et al.*, 1997). The general capabilities of wetlands may be inferred from work on an interesting design of intermittently fed shallow (~ 0.3 m), narrow (~ 1.3 m) gravel-filled hydroponics beds fed settled (secondary treatment) or treated sewage (tertiary treatment) in the UK and Egypt. The researchers expressed the performance as a decimal reduction distance or DRD, i.e. the distance required to effect a 90% reduction in pathogen or indicator concentration. The treatment appears to be partially related to organic load with better faecal coliform removal in tertiary treatment than in secondary treatment (DRD values of 51 m and 95 m respectively) and climate, removal was better in Egypt than in the UK (DRD for secondary treatment 30 m). The removal of salmonella (in the UK) was slightly better than the removal of faecal coliforms, however, the removal of *Vibrio cholerae* (presumably non-O1) was twice as slow as the removal of the indicator (Stott *et al.*, 1996).

Cooper *et al.* (1996) have usefully reinterpreted these and other reports and concluded that the rate of removal of faecal coliforms for secondary beds is about 0.02 log/h and about 0.08–0.04 log/h tertiary beds. These removal rates are subject to modest climatic (Stott *et al.*, 1996) and seasonal effects (Rivera *et al.*, 1995) and are comparable to those seen in the dark in WSP.

Speculation that protozoal predation (Green *et al.*, 1997) is important in bacterial pathogen removal has proven to be well founded. Decamp and Oliver (1998) found that observed levels of ciliate bacterivores could account for all the observed removal of faecal coliforms. The effect of organic loading may be accounted for by the reduced level of predation observed at

low oxygen tensions (Decamp *et al.*, 1999) and the climatic and seasonal variation by documented temperature effects (Iriberri *et al.*, 1995). However, Decamp and Oliver (1998) did not discount the possibility of other biotic and abiotic factors (bacteriophage attack, adsorption to solids, starvation and sedimentation) being significant in the removal of bacterial pathogens.

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Fate and behaviour of parasites in wastewater treatment systems

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1 INTRODUCTION

Diseases caused by intestinal parasites (helminths¹ and protozoa) are often a principal cause of human morbidity. Recent estimates suggest that at least 50% of the world's population may be infected with one or more helminth species (Chan, 1997), while prevalence rates of protozoan diseases ranging between 15 and 30% are not uncommon in tropical regions and may range between <1% and 10% in temperate areas (Mata, 1986; Casemore *et al.*, 1997). Transmission of many of the parasitic diseases, particularly helminthic infections, is often as a result of inadequate water supplies, lack of sanitation or disposal of insufficiently treated wastewater (Shuval *et al.*, 1986; Cairncross and Feachem, 1993). However, the association of faecally-contaminated drinking water supplies with recent waterborne outbreaks of emerging protozoan diseases is becoming of increasing concern (Smith and Rose, 1998; Huston and Petri, 2001).

Parasites are considered wastewater-associated pathogens of great public health importance due principally to their environmentally persistent transmissive stages, a low infective dose, limited or transient acquired immunity and morbidity, particularly in immunocompromised hosts (Anderson and May, 1979; Chan *et al.*, 1994; Guerrant *et al.*, 1999;

Stephenson *et al.*, 2000b; Stanley, 2001). It is apparent that while intestinal parasites continue to have a significant worldwide impact on human health, wastewater can be a major contributor to the transmission of parasitic diseases to susceptible hosts, either directly through occupational exposure and consumption of wastewater irrigated crops (Schlosser *et al.*, 1999; Blumenthal *et al.*, 2000), or indirectly through ingestion of contaminated water and exposure to polluted recreational water (Furtado *et al.*, 1998). However, the performance of wastewater treatment systems in removing parasites (thereby reducing the potential risk of disease transmission resulting from wastewater disposal or reuse) is rarely considered.

This chapter reports on current information about the occurrence of parasites in wastewater and their fate and behaviour during wastewater treatment. The removal and destruction of parasites by treatment processes is considered for 'conventional' (mechanical) and 'natural' wastewater treatment systems.

2 PARASITES IN RAW WASTEWATER

Wastewaters can contain a wide variety of excreted parasites depending on the source of wastewater and diseases present in the sewage contributing community (Table 31.1). The principal groups of helminth parasites include nematodes, cestodes and trematodes, while that of protozoa includes coccidia, flagellates,

¹ Helminth is the collective term for 'worm' which can be classified further into three main groups: roundworms, tapeworms and flatworms.

TABLE 31.1 Principal intestinal parasites found in raw wastewaters

<i>Parasite</i>		<i>Species</i>	<i>Common name</i>
HELMINTH	Nematode (roundworms)	<i>Ascaris lumbricoides</i>	human roundworm
		<i>Trichuris trichiura</i>	human whipworm
		<i>Ancylostoma duodenale</i>	hookworm
		<i>Necator americanus</i>	hookworm
		<i>Enterobius vermicularis</i>	pin worm/threadworm
		<i>Strongyloides stercoralis</i>	small roundworm ^a
		<i>Toxocara</i> spp.	
	Cestode (tapeworms)	<i>Taenia solium</i>	pork tapeworm
		<i>Taenia saginata</i>	beef tapeworm
		<i>Hymenolepis nana</i>	dwarf tapeworm
		<i>Hymenolepis diminuta</i>	rodent tapeworm
	Trematode (flatworms/flukes)	<i>Schistosoma</i> spp.	blood fluke
		<i>Clonorchis sinensis</i>	oriental liver fluke
		<i>Fasciola hepatica</i>	liver fluke
		<i>Paragonimus westermani</i>	lung fluke
PROTOZOA	Coccidia	<i>Cryptosporidium</i> spp.	cryptosporidiosis
		<i>Cyclospora</i> spp.	cyclosporiasis
		<i>Isospora belli</i>	isosporiasis
	Flagellate	<i>Giardia</i> spp.	giardiasis
	Amoebae	<i>Entamoeba histolytica</i>	amoebic dysentery
	Ciliate	<i>Balantidium coli</i>	balantidiasis
		<i>Microsporidia</i> spp.	Microsporidiosis

^a may also be called threadworm

amoebae and ciliates. General information about parasites can be found in Smith (1994) and Bogitsh and Cheng (1998).

Intestinal helminth parasites are usually found in wastewaters as eggs, although some species of helminths (notably *Strongyloides*) may be present as the larval form. Protozoan parasites are found in wastewater as cysts or oocysts depending on the species. Disease transmission to humans for the parasites shown in Table 31.1 can be through a variety of routes. In helminth infections, the infective stage may be an embryonated egg containing a larva which is then transmitted directly to other human hosts by ingestion (e.g. *Ascaris* spp.) or the larva hatches, developing to an infective filariform larva in the environment, which then penetrates the skin of unsuspecting hosts (e.g. hookworm). Some helminth infections require transmission to humans via one or more intermediate hosts (e.g. trematodes such as *Schistosoma* spp. or cestodes such as *Taenia* spp.). Protozoan (oo)cysts excreted in faeces are usually infective immediately or within a few hours of being shed in faeces;

a few species may require a longer incubation period of several days or weeks at ambient temperature. Disease transmission occurs through ingestion of (oo)cysts usually from faecally-contaminated water or food.

Humans can also become infected accidentally by some intestinal parasitic diseases normally transmitted between wild or domestic animals or agricultural livestock (Faust *et al.*, 1976; McLauchlin *et al.*, 2000; Robertson *et al.*, 2000a). These 'zoonotic' diseases include helminths such as *Fasciola*, *Paragonimus* and *Toxocara* (eggs of which are also sometimes found in raw municipal wastewaters) and also the protozoa *Giardia duodenalis* and *Cryptosporidium parvum*. Thus effective treatment of animal wastes with removal of excreted parasites can also be an important human health-related treatment objective.

2.1 Parasite occurrence in raw wastewaters

The occurrence and concentration of parasites in raw and influent wastewaters depends on a variety of factors, including the source of

wastewater (e.g. domestic/agricultural), prevalence, intensity and duration of infection in the contributing population, population size, per capita water consumption and proportion of population sewered, fecundity or multiplication rate of the parasite species and survival of the excreted stage in wastewaters. Point prevalence surveys indicate that parasites eggs and (oo)cysts can commonly occur in wastewaters worldwide with species diversity often reflected in sludge surveys (Reimers *et al.*, 1982; Schwartzbrod *et al.*, 1987; Paulino *et al.*, 2001). Eggs of nematodes (roundworms) and cestodes (tapeworms) are typically found in wastewaters (Fig. 31.1) with eggs of *Ascaris* usually predominating in raw wastewaters followed by *Trichuris*, hookworm and *Hymenolepis* spp. (Bouhoum *et al.*, 1997; Stott *et al.*, 1997). *Taenia* eggs are sometimes found as well as eggs of *Toxocara* and *Enterobius*. However, eggs of trematodes (flukes) are seldom reported as they tend to hatch on contact with water.

Protozoan parasites are also reported in wastewaters, with *Giardia* cysts and *Cryptosporidium* oocysts being the most frequently identified (Fig. 31.2). The relative frequency of protozoan species may vary depending on catchment characteristics. In England, the majority of protozoa found in sewage influent to treatment works were that of *Giardia*, while a greater prevalence of *Cryptosporidium* was found in agricultural impacted wastewaters (Bukhari *et al.*, 1997). Generally, although both species often coexist in wastewaters, *Giardia* usually exceeds *Cryptosporidium* in the number of positive samples detected and concentration. Point prevalence surveys carried out in developed countries, and reviewed by Robertson *et al.* (1999), indicated that over 66% of influent samples collected from wastewater treatment sites contained *Cryptosporidium* oocysts in comparison to more than 98% of influent samples containing *Giardia*. Cysts of *Entamoeba histolytica* have been identified in

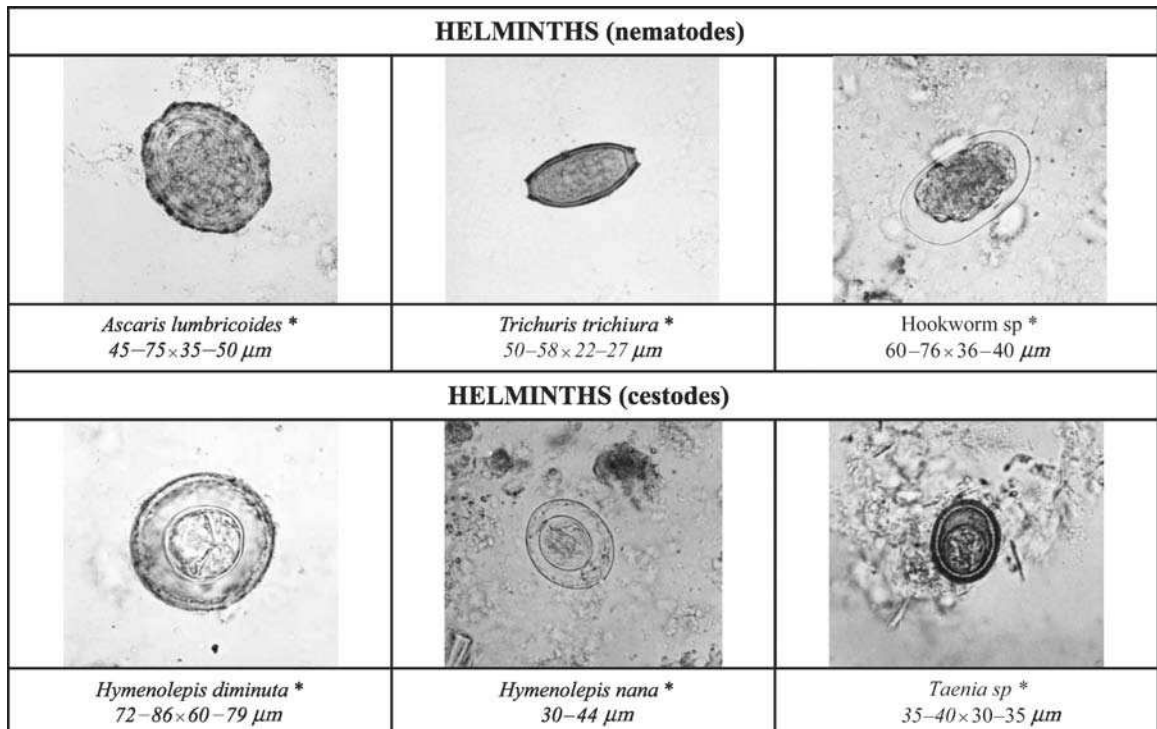


Fig. 1 Helminth parasites frequently found in raw and influent wastewaters.

* Printed with permission by R Ayres. Not to scale.

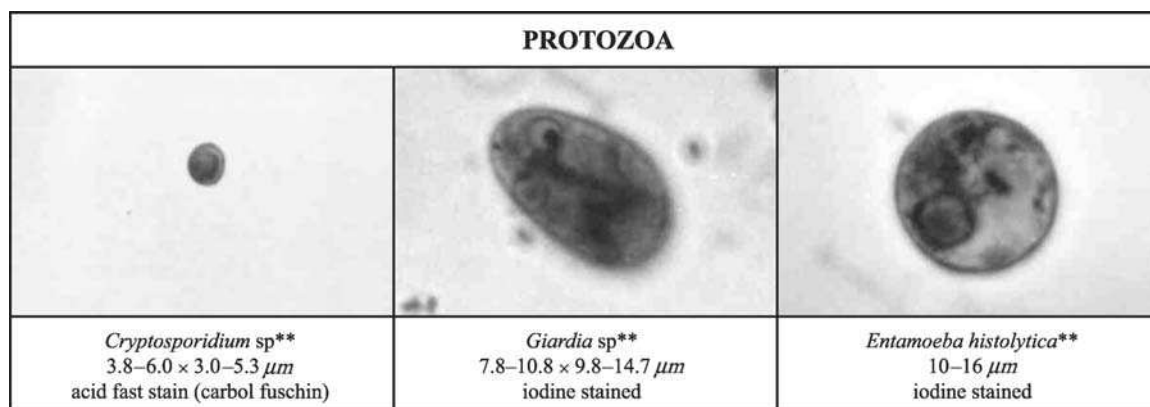


Fig. 2 Protozoan parasites frequently found in raw and influent wastewaters.

** Image source R Stott. Not to scale

NB. Parasite images can also be found on the web pages of the Parasitology Department at Chiang Mai University, Thailand. (www.medicine.cmu.au.th/dept/parasite/official/image.htm)

raw wastewaters from developing countries (Panicker and Krishnamoorthi, 1978a) but have not been reported in developed countries probably due to the low level of amoebiasis in the community. There is limited information on the occurrence of other 'emerging' enteric protozoan parasites such as *Isospora*, *Cyclospora* and microsporidia in wastewater. Sturbaum *et al.* (1998) reported unsporulated (i.e. undeveloped non-infective) oocysts of *Cyclospora* in 64% of wastewater samples and sporulated (i.e. infective) oocysts in 18% of samples collected from a shantytown in Peru where cyclosporiasis was widespread. Spores of microsporidia have been reported in raw wastewater in one study in which the human-specific *Encephalitozoon intestinalis* was identified (Dowd *et al.*, 1998).

2.2 Parasite concentration in raw wastewaters

The concentration of parasites commonly found in raw wastewater and influents to treatment systems can vary greatly. High prevalences and levels of infection for diseases exacerbated by partial sanitation, poverty and low water usage per capita (Cairncross and Feachem 1993; Chan, 1997) and the potentially large number of parasite eggs or (oo)cysts

excreted daily by infected individuals and zoonotic hosts (10^2 – 10^4 eggs/g and 10^5 – 10^7 (oo)cysts/g (Feachem *et al.*, 1983; Smith and Rose, 1998) can all contribute to high parasite loading rates in wastewaters. Dixo *et al.* (1995) reported up to 35 000 helminth eggs/l in wastewaters from an urban slum area of NE Brazil, while Lloyd and Frederick (2000) report concentrations of *Ascaris* of >100 000/l in a high density slum area in Bangladesh. However, maximum concentrations of around 1000 eggs/l are perhaps more typical in urban wastewaters. In India and Africa, concentrations ranging from 120 to around 900 eggs/l have been reported in raw wastewaters (Panicker and Krishnamoorthi, 1981; Ayres *et al.*, 1993). Ayres (1991) also found comparable numbers of eggs in Brazil with relatively high numbers of *Ascaris* eggs (up to 700 eggs/l) and up to 19 *Trichuris* eggs/l, 8 hookworm eggs/l and 20 *Hymenolepis* spp eggs/l. Of all the helminth eggs likely to be present in wastewaters, *Ascaris*, *Trichuris* and hookworm are of particular public health concern because of severe socioeconomic consequences of an estimated 39 million DALYs (disability-adjusted life year) lost to these infections (Chan, 1997; Crompton, 2000; O'Lorcain and Holland, 2000; Stephenson *et al.*, 2000a). There is limited information on protozoan (oo)cysts

in raw wastewaters in developing countries. Concentrations of 25 000 cysts per litre have been found in Africa (Grimason *et al.*, 1996b), although, in general, lower concentrations rarely exceeding 1000 (oo)cysts/l have been reported (Panicker and Krishnamoorthi, 1981; Jimenez *et al.*, 2001). However, it is likely that higher concentrations may be present due to high reported incidences of protozoan infection (exceeding 50% in some communities) in developing countries (Zu and Guerrant, 1993).

Relatively few studies have assessed parasite egg numbers in raw sewage in developed countries. Where reported, helminth eggs occur in lower concentrations, rarely exceeding 100 eggs/l, probably reflecting a generally low incidence of helminthiasis in communities. In contrast, protozoan (oo)cysts have been found in very high numbers in several developed countries with concentrations frequently exceeding 10 000 (oo)cysts/l; up to 44 000 cysts/l have been reported in UK influent wastewaters (Robertson *et al.*, 1995). Significantly higher numbers of *Giardia* than *Cryptosporidium* have been detected in influent domestic wastewaters in the USA, UK and Europe (Mayer and Palmer, 1996; Robertson *et al.*, 2000b; Medema and Schijven, 2001). However, higher numbers of *Cryptosporidium*, more than 10^4 – 10^5 oocysts/l, have been reported in US livestock slaughterhouse and UK holding pen effluents (Robertson *et al.*, 1999).

The magnitude of parasite concentrations in raw wastewaters may also fluctuate temporally. Diurnal and seasonal variations in parasite egg and (oo)cyst numbers have been observed (Casson *et al.*, 1990; Gassman and Schwartzbrod, 1991; Sykora *et al.*, 1991; Capizzi and Schwartzbrod, 1998) possibly reflecting human activity factors and periods of increased incidence of infection in communities (Casemore *et al.*, 1997; Chai *et al.*, 2001). Parasite concentrations in wastewaters can also be considerably affected by rainfall and storm events (Panicker and Krishnamoorthi, 1978a; Atherholt *et al.*, 1998; Capizzi and Schwartzbrod 1998). Parasite loading rates into wastewater treatment systems can thus

be highly variable with short- and longer-term trends in excreted parasite concentrations possible.

In general, raw wastewaters in developed countries may contain helminth egg numbers of up to 10^2 eggs/l and protozoa ranging from 10 to 10^5 (oo)cysts/l. In comparison, concentrations of helminths of 10^2 – $<10^4$ eggs/l and protozoa of 10^2 – 10^3 (or 4) (oo)cysts/l have been reported for developing countries. Since parasites are highly infectious with infective doses as few as 1–30 eggs/(oo)cysts capable of causing infections (Feachem *et al.*, 1983; Okhuysen *et al.*, 1999; Kothary and Babu, 2001), a high degree of parasite removal (at least 2–4 log units, i.e. >99%) may be required by sewage treatment processes for public health protection and compliance with current recommended quality criteria of 0.1–1 egg/l for wastewater irrigation (WHO, 1989; Blumenthal *et al.*, 2000).

2.3 Parasite removal in wastewater treatment systems

The effect of treatment processes on the fate and behaviour of parasites has been evaluated for high rate mechanical processes (conventional) and low rate (natural) treatment systems. Parasite removal is commonly assessed by comparing the microbiological quality of influent and effluent wastewaters. Removal rates reported for treatment systems may thus only refer to the removal of parasite eggs or (oo)cysts from suspension in wastewaters into sediments and sludge, and not to their destruction or inactivation as a result of the treatment process. In addition, non-detection of parasites in wastewaters does not guarantee that effluents are completely free of parasites since none of the enumeration and detection methods for parasitological analysis of wastewaters can guarantee a 100% recovery of parasites in processed samples. In relation to inactivation, reports usually refer to an assessment of viability rather than infectivity of eggs or (oo)cysts. Viability of helminth eggs is commonly determined by a morphological assessment

of egg development to an embryonated stage containing a larva (e.g. *Ascaris*) or the hatchability of eggs (e.g. hookworm). The viability of protozoan (oo)cysts is typically by dye inclusion/exclusion or excystation methods (e.g. *Cryptosporidium*).

The contribution of each treatment process on removal or destruction of parasite is discussed below for the various levels of wastewater treatment, which usually consist of a sequence of physical, biological and sometimes chemical processes generally classified as preliminary, primary, secondary and tertiary or advanced. The fate of parasites in sludge treatment processes is covered comprehensively in Lewis-Jones and Winkler (1991).

2.4 Preliminary treatment – screening and grit removal

Preliminary treatment processes are used to improve downstream operations and typically consist of physical processes for screening and grit separation to remove large solids and rubbish. It is unlikely that parasites will be removed at this stage of treatment. The use of coarse screens with bar spacing usually at >10 mm and even fine screens (0.2–10 mm spacing) will not impede the passage of parasite eggs and cysts which range in size from 0.001 to 0.15 mm. Short detention times and high velocity flow of wastewater through grit chambers are also unlikely to facilitate removal of parasites.

2.5 Primary treatment

Parasite removal is highly variable between primary treatment processes. Generally, helminth eggs are removed more effectively than protozoan (oo)cysts. Performance for primary treatment processes for helminth removal range from 60 to 99% and where reported, primary effluents can contain between 0 and 1500 eggs/l. Removal of protozoa is more variable ranging from 4 to 93% with effluents containing $10\text{--}10^5$ (oo)cysts/l.

2.5.1 Primary settlement tanks

The aim of primary treatment is to remove settleable solids of 0.05–10 mm by gravity in sedimentation tanks. The principal mechanism for parasite removal during primary treatment is also assumed to be by sedimentation, and possibly by adsorption onto solids that are settling out of suspension.

2.5.1.1 Plain/simple (non-assisted) sedimentation.

Parasite sedimentation rates (estimated using Stoke's law for discrete particle settling) will vary between parasite species depending on the specific gravity and dimensions of the parasite and liquid density (and temperature) suggesting that eggs of *Ascaris* and *Trichuris* will be removed more effectively than eggs with slower rates of settling velocity, such as hookworm and (oo)cysts of protozoa (Table 31.2). Panicker and Krishnamoorthi (1978b) reported *Ascaris* and *Trichuris* removal rates of 96% and 90% respectively in comparison to removal rates of 80% for hookworm eggs during primary sedimentation. Stokes' law, however, does not hold for *Giardia*, *Entamoeba histolytica* and *Cryptosporidium* (oo)cysts as their Reynolds' numbers are below 10^{-4} indicating that protozoan (oo)cysts are unlikely to be effectively removed by sedimentation. Nevertheless, protozoan removal has been reported during primary sedimentation in operational works, although removal performance is poor. Removal rates of between 4 and 47% have been reported by Robertson *et al.* (2000b) for protozoa in primary sedimentation tanks with estimated retention times of up to 25 hours. *Cryptosporidium* were less efficiently removed (19% average removal) than *Giardia* (38% removal) probably as a result of their smaller size and slower settlement rates.

A variety of factors can hinder the removal of parasites during primary sedimentation. Free-falling settling velocities of helminth eggs (<0.8 m/h excluding *Schistosoma* spp.) and protozoan (oo)cysts (<0.01 m/h at 20°C) are not much greater than the upward flow velocity of wastewater within tanks (conventionally between 0.5 and 1.5 m/h) which can prevent parasite eggs and (oo)cysts from settling out of suspension unless assisted. Adsorption,

TABLE 31.2 Estimated settling velocities of helminth eggs and protozoan (oo)cysts (in water 5–20°C)

Parasite	Dimensions (μm) average	Specific gravity (g/cm ³)	Settling velocity (m/ h)		
			5°C	15°C	20°C
Helminths					
<i>Ascaris lumbricoides</i> – fertile	60 × 45	1.11	0.51	0.48	0.77
<i>Ascaris lumbricoides</i> – infertile	90 × 40	1.2	2.09		3.15
<i>Trichuris trichiura</i>	50 × 22	1.15	0.48		0.73
Hookworm	60 × 40	1.055	0.26		0.39
<i>Taenia saginata</i>	40 × 30	1.23	0.48	0.83 ^a	0.72
<i>Schistosoma mansoni</i>	150 × 55	1.18	5.24		7.87
Protozoa					
<i>Cryptosporidium parvum</i>	5.0 – 4.5	1.045	0.0014	0.0035 ^b	0.002
<i>Giardia lamblia</i>	12.2 × 9.3	1.036	0.007		0.01
<i>Entamoeba histolytica</i>	12 – 10	1.07		0.018	

References: Thienpoint *et al.*, 1986; Fayer *et al.*, 1997; Medema *et al.*, 1998; David and Lindquist, 1982; Shuval *et al.*, 1986; Pike, 1990

^a based on *Taenia* Sp gravity 1.3

^b based on *Crypto* Sp gravity 1.08

aggregation or attachment to larger or denser particles may increase settling velocities and enhance parasite settling during clarification treatment processes. Whitmore and Robertson (1995) and Medema *et al.* (1998) present evidence that this could be an important removal mechanism for protozoan (oo)cysts. Settling velocities of *Cryptosporidium* oocysts during primary sedimentation in laboratory trials (0.022–0.028 m/h) exceeded rates in water suggesting attachment of oocysts to particles with greater settling. Protozoan (oo)cysts were also found to attach readily to wastewater particles in secondary sedimentation effluents, with maximum velocities of 0.25 m/h observed when (oo)cysts attached to the largest particles (> 200 μm ; Medema *et al.*, 1998). In contrast, no significant association has been found between suspended solids and parasite eggs numbers in wastewaters suggesting a lack of solid-associated removal for helminth eggs (Ayres, 1991; Stott *et al.*, 1999). Interestingly, Casson *et al.* (1990) also reported a lack of correlation between the removal of *Giardia* cysts and suspended solids during primary treatment.

Detention time can also affect the parasite removal performance of primary sedimentation. Sedimentation tanks are usually 1.5–4 m deep with detention times typically 1.5–2.5 h,

although primary settling tanks preceding biological treatment may be designed for shorter detention periods of 0.5–1 h (Metcalf and Eddy, 1991; Gray, 1999). Parasite removal increases with retention time, with settlement times of around 2 h observed for effective removal of parasite eggs. Parasite removal rates after 1.5 h sedimentation of, on average, 52% for helminth eggs and 27% for protozoan cysts, were improved after 2 h detention to 74% and 67% for parasite eggs and cysts respectively (Bhaskaran, *et al.*, 1956; Panicker and Krishnamoorthi, 1978b). Longer retention times thus facilitate the removal of free-falling parasite eggs and also allow a greater degree of flocculation, which may assist in the settlement of protozoan (oo)cysts.

Sedimentation tanks are generally designed to remove all particles which have a terminal velocity equal or greater than the surface overflow rate ($\text{m}^3 \cdot \text{m}^{-2}/\text{day}$, i.e. m/day). Factors such as detention time, water velocity, turbulence (from sewage inflow), wind, and tank design can greatly influence sedimentation rates. Consequently, primary removal of parasites is highly variable. Protozoa removal rarely exceeds 50%, while removal rates for helminth eggs average 70%. Numbers of parasites in primary treated effluents can also be considerable, particularly for protozoa due

to limited removal. In some studies protozoan (oo)cysts concentrations have exceeded 10^4 *Giardia* cysts/l and 10^5 *Cryptosporidium* oocysts/l (Robertson *et al.*, 2000b; Medema and Schijven, 2001). Generally, protozoan (oo)cysts can average 2400/l (general range 17–6370) in comparison to around 10 helminth eggs/l (range 0–15) reported in primary effluents (Wang and Dunlop, 1954; Robertson *et al.*, 1995; Stott *et al.*, 1996; Chauret *et al.*, 1999; Medema and Schijven, 2001).

2.5.1.2 Chemically assisted sedimentation. Chemically assisted sedimentation, using coagulants (ferric chloride, alum or lime) to assist flocculation and enhance solids removal, can also facilitate concomitant enhanced removal of parasites from suspension, particularly protozoa. Pilot studies in Brazil, using lime treatment, reported removal rates of 99.99% for helminth eggs (4.01 log units; Taylor *et al.*, 1994). The viability of eggs was not assessed but the effect of lime treatment on inactivation of helminth eggs is time, temperature and pH dependent. Elevated pH (average of 11.5) may not alone have had a significant detrimental effect, as helminth eggs such as *Ascaris* can survive more than 120 days at pH > 12 (Storm *et al.*, 1981 cited Taylor *et al.*, 1994). In Mexico, pilot studies of advanced primary treatment (APT), using coagulants (typically aluminium sulphate), anionic polymer flocculants and a high rate sedimentation basin with a hydraulic retention time ranging from 15 to 50 min, demonstrated removal rates of 95% for helminth eggs (effluent contained around 1.2 helminth eggs/l) at influent wastewater concentrations of 23 eggs/l (Jimenez *et al.*, 2000). A removal efficiency of 71% was also reported for protozoan cysts with APT effluent containing almost 400 cysts/l when influent wastewater contained around 1300 cysts/l (Jimenez *et al.*, 2001). Comparable protozoan removal rates were reported by Payment *et al.* (2001) for a large operational works in Canada in which physicochemical treatment removed, on average, 76% *Giardia* cysts with effluent containing up to 350 cysts/l, when influent contained around 1500 cysts/l. Oocysts of *Cryptosporidium* were less effectively removed

with removal rates averaging 27% and effluent containing around 10 oocysts/l. Higher removal rates for *Giardia* of 80–93% have been reported for a plant in the USA treating wastewater containing up to 13000 cysts/l (Mayer and Palmer, 1996).

Rates of helminth egg removal during chemical assisted sedimentation are comparable to non-assisted/plain sedimentation but are achieved with shorter detention periods. However, protozoan removal rates of, on average, 40% during non-assisted sedimentation (due mainly to removal of solids) may be improved to around 70% removal efficiency using chemical assisted sedimentation.

2.5.2 Septic tanks and anaerobic digesters

There is limited information available on the removal of parasites from septic tanks. Sedimentation is the most likely removal mechanism and thus, depending on tank conditions and extended retention times, septic tanks are likely to be relatively effective for the removal of parasites. Feachem *et al.* (1983) suggested that septic tanks with a normal retention time of 1–3 days would reduce protozoa and helminths by 0–2 log₁₀ units. These rates are supported by Bhaskaran *et al.* (1956), who reported helminth removal rates of 99.4% in an experimental septic tank system with a 3-day retention and by Lloyd and Frederick (2000) of 99.95–100% helminth egg removal for an OXFAM emergency system of two tanks in series providing anaerobic settling conditions and a hydraulic retention time of 2–3 days. However, Bhaskaran *et al.* (1956) also reported concentrations of *Ascaris* and hookworm eggs of 30 eggs/l in effluent from an operational system. Further treatment of the septic tank effluent through a rock filter removed more than 70% of parasite eggs and reduced the concentration of eggs to 8 and 13 eggs/l for *Ascaris* and hookworm respectively. Chlorination of the rock filter effluent further reduced the egg concentration to around 4 eggs/l and improved overall parasite egg removal to 95%. Lloyd and Frederick (2000) also report disappointing rates of parasite egg removal of 84.5–89.5% in a double vaulted

septic tank indicating poor sedimentation/retention time. However, the system received high numbers of eggs at 4×10^4 – 2×10^5 eggs/l, which were reduced to 2×10^4 after septic tank treatment and reduced further to 5×10^3 after treatment in two ponds (overall reduction of 97.5%).

The anaerobic digestion process is generally less effective in parasite removal. Paulino *et al.* (2001) found removal rates of between 59.7% and 93% for helminth eggs and protozoan cysts during anaerobic digestion. Black *et al.* (1982) reported that less than 25% of *Ascaris* eggs were destroyed in anaerobic digesters, but no effect was found on *Trichuris* or *Toxocara* eggs. Furthermore, the proportion of viable eggs was higher after treatment indicating that non-viable eggs were preferentially removed/destroyed in the process. Other studies have reported that mesophilic digestion does not significantly reduce numbers of protozoan (oo)cysts, although high densities of *Giardia* and *Cryptosporidium* of up to 11 800 cysts/100 g and 3810 oocysts/100 g can be present in sludge (Sykora *et al.*, 1991; Chauret *et al.*, 1999). However, several studies have shown that anaerobic digestion can significantly reduce parasite viability with the effect of anaerobic digestion on parasite inactivation related to temperature, exposure time and parasite species. Cram (1943) reported that the viability of *Ascaris* eggs was unaffected after 3 months mesophilic anaerobic digestion at 20–30°C, but was significantly reduced after 6 months when only 10% of *Ascaris* eggs were still viable. Hookworm eggs were more susceptible surviving only 40 days at 30°C. In contrast, thermophilic digestion (>50°C) can significantly reduce egg viability with 15 minutes at 55°C reportedly required for 99% inactivation of *Ascaris* eggs (Pike *et al.*, 1988 cited by Lewis-Jones and Winkler, 1991). Although mesophilic digestion appears to be ineffective in removing viable helminth eggs, the process has a greater effect on protozoan cyst inactivation. Cram (1943) reported that cysts of *Entamoeba histolytica* did not survive the digestion process. *Giardia* and *Cryptosporidium* (oo)cysts are also readily inactivated. Gavaghan *et al.* (1993) and Stadterman *et al.*

(1995) reported that between 40 and 50% of *Giardia* and *Cryptosporidium* (oo)cysts were inactivated (non-viable) after 4 hours exposure to mesophilic digestion at 37°C and more than 99% of (oo)cysts were non-viable after 24 h exposure in laboratory studies. However, Whitmore and Robertson (1995) reported that 10% of *Cryptosporidium* oocysts were still viable after 18 days exposure to mesophilic anaerobic digestion and suggested that it was likely that viable oocysts would still be present after 3 weeks.

Clearly, mesophilic anaerobic digestion may not completely remove parasite eggs and (oo)cysts from wastewaters. However, with detention times varying from 30 to 60 days for the standard rate anaerobic digestion process and up to 15 days for high rate digestion (Metcalf and Eddy, 1991), significant inactivation of parasites may occur, especially for protozoa and reduction in (oo)cyst viability.

2.5.3 Upflow anaerobic sludge blanket (UASB)

High rate anaerobic processes, such as UASB reactors, have been used as a pretreatment stage for a variety of processes, including activated sludge and waste stabilization ponds. In one study, Dixo *et al.* (1995) reported that a UASB digester reduced parasite egg concentrations in raw wastewater from 17 000 eggs/l to 1740 eggs/l (89.6% removal) after a theoretical hydraulic retention time of 7 hours. With a volumetric egg loading of $5.8 \times 10^7/\text{m}^3/\text{day}$, the UASB performance was better than that of an anaerobic pond loaded at $4.5 \times 10^5/\text{m}^3/\text{day}$ (with a retention time of 10 h), which demonstrated a rate of removal of only 26.6%. The authors suggested that the majority of parasite eggs were removed in the UASB by filtration and aggregation as the influent flowed up through the sludge blanket. Sedimentation is unlikely to play an important role in removal as upflow velocities (0.6–0.9 m/h) used to keep the sludge blanket in suspension (Metcalf and Eddy, 1991) are higher than most parasite settling velocities. A high removal efficiency in the UASB reactor was attributed by Dixo *et al.* (1995) to a hydraulic flow regime

approximating plug flow ensuring all influent was strained through the sludge blanket limiting short-circuiting. The eggs in the UASB effluent were sequentially removed in experimental waste stabilization ponds incorporating a secondary facultative pond (3.5 eggs/l) and completely removed after the first tertiary maturation pond. In another study, a UASB demonstrated removal rates of 70–82% for helminth eggs with effluent containing on average 18 eggs/l (range 14–21 eggs/l) after treatment of raw wastewater containing much lower parasite concentrations of between 47 and 121 eggs/l (Chernicharo *et al.*, 2001).

2.6 Secondary treatment

Primary sedimentation (assisted or non-assisted) is not an effective process for complete removal of parasite eggs or (oo)cysts. However, primary treatment combined with secondary treatment can significantly improve parasite removal efficiency especially for protozoa. Robertson *et al.* (2000b) reported that removal of *Giardia* and *Cryptosporidium* occurred predominantly during secondary treatment (>60% removal) rather than during primary treatment (<40% removal).

2.6.1 Conventional wastewater treatment

Secondary treatment generally consists of an aerobic biological treatment process for the removal of organic material and nutrients. Typical conventional secondary treatment systems utilize mechanical high rate processes such as activated sludge and trickling filters. Generally, conventional treatment systems are not particularly effective in parasite removal, especially for protozoa. In general, secondary treatment plants (STP) achieve 80–100% removal of helminth eggs (<1–60 eggs/l) and 5–99% removal of protozoan cysts (10–< 8000 (oo)cysts/l).

Comparisons between activated sludge and trickling filter plants suggest parasite removal is highly variable. Generally, activated sludge plants (including secondary sedimentation) demonstrate slightly higher removal rates

than trickling filters. *Giardia* removal in activated sludge plants varied from 97 to 99.8% in comparison to 92–< 98.3% removal in trickling filter plants (Casson *et al.*, 2000; Wiandt *et al.*, 2000). Robertson *et al.* (2000b) reported more variable protozoan removal of 15–99% in activated sludge compared to 5–85% removal in trickling filters but no significant difference in efficacy of protozoan removal was found between the two processes. Helminth egg removal rates are comparable for activated sludge and trickling filter plants with overall removal performance typically 75–100% (Bhaskaran *et al.*, 1956; Panicker and Krishnamoorthi, 1978b; Rose *et al.*, 1996).

2.6.1.1 Activated sludge. The activated sludge process is one of the most commonly used for secondary wastewater treatment. As a suspended-growth biological treatment process, activated sludge utilizes a dense microbial culture in suspension to biodegrade organic material under aerobic conditions and form a biological floc for solid separation in the settling units. Diffused or mechanical aeration maintains the aerobic environment in the reactor. Typical retention times are 5–14 hours in conventional units rising to 24–72 in low rate systems (Gray, 1999).

Performance of activated sludge systems in removing parasites varies with parasite species. Bhaskaran *et al.* (1956) reported greater removal of *Ascaris* and *Trichuris* (96–97%) than for hookworm (88%). Similarly, higher rates of removal have been reported for *Giardia* cysts (>97%) than for cysts of *Entamoeba histolytica* (<85%) or *Cryptosporidium* oocysts (around 80%) (Panicker and Krishnamoorthi, 1978b; Feachem *et al.*, 1983; Madore *et al.*, 1987; Casson *et al.*, 1990; Mayer and Palmer, 1996).

Efficacy of removal reported may depend on whether effluents were sampled after aeration and sludge separation or after secondary sedimentation following activated sludge treatment. Parasite removal during activated sludge treatment occurs predominantly during secondary sedimentation with free and assisted sedimentation likely to be occurring in a similar manner to that during primary sedimentation.

Medema *et al.* (1998) reported that 70% of protozoan (oo)cysts attached to particles 0–200 µm in secondary settled sewage after 24 h of activated sludge treatment suggesting enhanced settling of protozoa during secondary sedimentation. Higher removal efficiencies for protozoan (oo)cysts have been reported after activated sludge treatment (including secondary sedimentation) than during primary sedimentation. At two activated sludge plants, primary sedimentation removed 4–42% protozoan (oo)cysts in comparison to 53–98% removal in activated sludge units (including secondary sedimentation) (Robertson *et al.*, 2000b). Overall the activated sludge plants removed <99% of protozoan (oo)cysts (primary and secondary sedimentation included). Similarly, Mayer and Palmer (1996) found that most protozoan removal occurred after activated sludge treatment (84.5–99.5%; lowest rates were that of *Cryptosporidium*); removal performance improved to 98.9–99.9% for the activated sludge plant (complete works including primary sedimentation). In contrast, both primary and secondary sedimentation may be important for helminth egg removal. In an activated sludge plant not preceded by primary sedimentation, removal rates of 75% helminth eggs and 93% protozoan cysts were reported. Where primary and secondary sedimentation was considered, activated sludge plants (complete works) removed 97–100% *Ascaris* eggs and 99.7% *Schistosoma mansoni* eggs (Rowan, 1964). Effective removal of *Ascaris* is most likely due to sedimentation. The high removal rates for *Schistosoma* may have been facilitated by egg hatching since *Schistosoma* eggs have been found to hatch readily in experimental wastewater tanks (Jones *et al.*, 1947).

Relatively high rates of protozoan removal during activated sludge treatment may be due to mixing enhancing attachment of (oo)cysts to flocs for subsequent solid assisted sedimentation in secondary clarifiers and/or flocculated material acting as a settling blanket during secondary sedimentation. In laboratory studies, Stadterman *et al.* (1995) found that *Cryptosporidium* removal was higher for activated sludge effluents compared to trickling filter after secondary sedimentation for 2.8 h

(92% removal activated sludge) and 3.5 h (50% removal trickling filter). The authors suggested that protozoan removal during activated sludge occurred in the hindered (zone) settling region in which flocculated material forms a sludge blanket and filters suspended particles.

Despite relatively high removal rates, parasites can survive the activated sludge process and are usually not completely removed. Helminth eggs of *Taenia saginata*, *Ascaris*, *Trichuris*, hookworm and *Hymenolepis* spp. and protozoan (oo)cysts of *Giardia* spp., *Entamoeba* spp., *Cryptosporidium* spp. as well as microsporidia spores have all been reported in activated sludge effluents (Bhaskaran *et al.*, 1956; Panicker and Krishnamoorthi, 1978b; Mayer and Palmer, 1996; Dowd *et al.*, 1998). The concentration of parasites in activated sludge treated effluents is usually low at less than 5 eggs/l, though higher numbers of around 60 *Ascaris* eggs/l have been reported (Bhaskaran *et al.*, 1956). Numbers of protozoan (oo)cysts in activated sludge effluents can be high despite relatively high removal rates. Concentrations have ranged from 11 to 7600 (oo)cysts/l, while 0.05–< 20 (oo)cysts/l have been specifically reported in activated sludge effluents following secondary sedimentation (Madore *et al.*, 1987; Sykora *et al.*, 1991; Mayer and Palmer, 1996; Chauret *et al.*, 1999; Robertson *et al.*, 2000b).

The activated sludge process has little effect on the viability and infectivity of parasite eggs that pass through the system (Cram, 1943; Newton *et al.*, 1949), but may affect (oo)cysts. Robertson *et al.* (2000b) reported no reduction in viability for *Cryptosporidium* oocysts during activated sludge treatment. Oocyst viability after 7 days exposure was 68% in raw wastewaters and was comparable in primary sedimentation effluent (77% viability), activated sludge effluent (73% viability) and secondary sedimentation effluent (77% viability). In contrast, Carraro *et al.* (2000) reported a presumed reduction in protozoan (oo)cysts' viability from 66 to 70% in raw wastewater to 46–50% in filtered activated sludge effluent. They suggested that activated sludge treatment favours the excystation of (oo)cysts. Infectivity of *Cryptosporidium* oocysts was

also reduced after laboratory simulation of activated sludge, although some oocysts retained infectivity after treatment to initiate infection (Villacorta-Martinez de Maturana *et al.*, 1992). Incidence of infection in mice was variable (5–6% and 74–95% in two trials) but intensity of infection was reduced to 82–99%.

In general, activated sludge systems demonstrate average removal rates of 92% and 87% for parasite eggs and cysts respectively (Panicker and Krishnamoorthi, 1978b; Madore *et al.*, 1987; Schwartzbrod *et al.*, 1989; Casson *et al.*, 1990; Rose *et al.*, 1996; Robertson *et al.*, 2000b). For the activated sludge process itself, 0–2 log₁₀ units of removal may be expected for protozoa and helminths: activated sludge including primary and secondary sedimentation improves the removal performance to 0–3 log₁₀ units of removal for parasites.

2.6.1.2 Trickling filter. Trickling filtration is an attached growth process which utilizes filter beds, consisting of highly permeable media to which microorganisms are attached, to treat wastewater which is percolated or trickled through the bed. Following filtration, effluent is treated in settling tanks where remaining solids and detached biofilm from the filter bed are separated from the treated wastewater. In the trickling filter process, biodegradation occurs due to microbial activities associated with the biofilm attached to the filter media.

Biological filter beds as a unit process generally contribute little to the removal of parasites from wastewaters and parasite eggs and cysts pass through into effluents. Panicker and Krishnamoorthi (1978b) reported low rates of helminth egg removal by trickling filter units ranging from 18 to 35% and trickling filter bed effluent concentrations ranging from 22 to 41 eggs/l. In contrast, higher removal rates of 98–100% for parasite eggs have been found in India after treatment of filter bed effluents in humus tanks (i.e. after secondary sedimentation): all hookworm eggs were removed and final effluents contained less than 1 *Ascaris* egg/l when raw wastewater contained 466 eggs/l (Bhaskaran *et al.*, 1956). Treatment of

wastewater through a trickling filter plant in the Ukraine reduced *Ascaris* egg concentrations by 97%. Raw wastewater containing 60 eggs/l was reduced to 20 egg/l after primary sedimentation (67% removal) to 13 eggs/l after trickling filtration (35% removal) and finally to 2 eggs/l after secondary sedimentation (85% removal) (Vishneuskaya, 1938 cited Feachem *et al.*, 1983).

Protozoan (oo)cysts may not be so efficiently removed. In several trickling filter plants, rates of protozoan (oo)cyst removal averaged 84% after secondary sedimentation in comparison to higher removal rates of 91% for helminth eggs (Kott and Kott, 1967; Panicker and Krishnamoorthi, 1978b): *Ascaris* and *Trichuris* eggs were predominantly removed (95%) followed by hookworm (87%), *Giardia* (85%), *Hymenolepis nana* (84%) and *Entamoeba histolytica* (83%) (Panicker and Krishnamoorthi, 1978b).

Bukhari *et al.* (1997) reported highest removal rates for *Giardia* when wastewater treatment works included biological filtration (26–94%) in comparison to activated sludge treatment systems (26–71.7%). Trickling filter effluents contained <10–50 cysts/l, whereas activated sludge effluents contained <10–210 cysts/l. Feachem *et al.* (1983) reported similar removal rates of 71–94% for *Entamoeba histolytica* cysts in trickling filters. However, oocysts of *Cryptosporidium* may not be so favourably removed. Poor removal rates of 5% have been shown for trickling filters with effluents containing up to 1000 oocysts/l (and also up to 3800 *Giardia* cysts/l; Robertson *et al.*, 2000b).

Parasite removal mechanisms may be similar to those that contribute to the removal of suspended particulate matter in trickling filter beds, such as mechanical straining, sedimentation or inertial impaction, interception, adhesion or flocculation (Metcalf and Eddy, 1991). Biofilms on filter media may also provide attenuation sites for parasites as wastewater percolates through the trickling filter with grazing by predatory fauna associated with biofilms possibly contributing to removal and inactivation (Stott *et al.*, 2001). Panicker

and Krishnamoorthi (1978b) proposed that predation by oligochaetes (which colonize the filter beds) may reduce pathogen survival.

Although the majority of parasites are removed throughout the treatment works up to secondary sedimentation, parasites are rarely completely removed, and parasite cysts and eggs can pass through trickling filters, with filter effluents potentially containing significant parasite concentrations (Nupen and Villiers, 1975; Bukhari *et al.*, 1997). Treated effluents may contain, on average, <5 helminth eggs/l (range 0–13) and up to 4000 protozoan (oo)cysts/l. Trickling filters are generally less effective in removing parasites than activated sludge processes. From the literature, trickling filters (including primary and secondary sedimentation) demonstrate average removal efficiencies of 92% for parasite eggs and 77% for protozoan (oo)cysts. In isolation, trickling filters may demonstrate 0–1 log₁₀ units of reduction for helminth and protozoan (oo)cysts (Feachem *et al.*, 1983). Higher parasite removal rates are reported after secondary sedimentation of trickling filter effluent of around 0–2 log₁₀ units.

2.6.1.3 Aerated lagoons. Fully aerated lagoons or ponds may be considered as a simple modification of the activated sludge process with screened rather than settled sewage aerated in ponds with no sludge return. Retention times are typically 2–6 days in these lagoons followed by further treatment in a settling pond (5–10 days retention) or secondary sedimentation tank (usually >2 hours retention) (Feachem *et al.*, 1983).

Parasites are not completely removed during treatment despite longer retention times in aerated lagoons than in conventional activated sludge plants. Parasite removal rates ranging from 50 to 100% have been reported for lagooning without secondary sedimentation (Panicker and Krishnamoorthi, 1978b, 1981). Average removal rates for helminth eggs were 88% and 85% for protozoan cysts. Smaller eggs with lower specific gravity and hence slower settling velocities (such as hookworm) are usually not removed as effectively as the larger and heavier eggs (such as *Ascaris*). Panicker

and Krishnamoorthi (1981) reported 0–40 hookworm eggs/l, 0–20 *Ascaris* eggs/l and up to 96 *Entamoeba histolytica* cysts/l in lagoon effluents. Settling ponds following aerated lagoons are likely to enhance parasite removal due to extended retention times. Feachem *et al.* (1983) suggested aerated lagoons (including settling ponds) could achieve 1–3 log and 0–1 log removal for helminth and protozoa respectively.

2.6.1.4 Oxidation ditches. Oxidation ditches are also another modification of the activated sludge process. Screened sewage is aerated and circulated around an oval ditch (0.25–0.35 m/s). Oxidation ditches usually operate with long detention and solids retention times (ditch retention time of 1–3 days). Effluent is usually then treated in secondary sedimentation tanks with settled sludge returned to the oxidation ditch (Feachem *et al.*, 1983).

Panicker and Krishnamoorthi (1978b, 1981) reported that under optimum operation, oxidation ditches removed, on average, 92% (80–100%) of parasite ova and cysts; minimum rates of removal were 72% helminths and 60% protozoa, with effluents containing 4–14 helminth eggs/l and up to 120 cysts/l of *Entamoeba histolytica* (Panicker and Krishnamoorthi, 1981). Sedimentation was considered to be the principal removal process despite water turbulence with removal rates enhanced by floc settlement with parasites attached. Rates of removal may vary with helminth species and velocity of water flow.

2.7 Tertiary treatment

Wastewater treatment plants may use tertiary treatment measures for improving the quality of effluents from conventional treatment works. Processes utilized include sand filtration and disinfection. Effects of tertiary treatment on parasite removal are variable. Where tertiary treatment has been employed, the majority of parasite removal occurs in the preceding primary and secondary treatment processes. Overall, tertiary treatment plants (TTP) remove >99% helminth eggs (10 eggs/l) and 95–99% protozoa (0.1–<20 (oo)cysts/l).

2.7.1 Sand filtration

Sand filtration enhances a reduction of parasites, although removal rates may be highly variable depending on grain size and hydraulic loading rates (Logan *et al.*, 2001). In general, sand filtration is effective in removing helminths with almost 100% removal reported for helminth eggs (Rose *et al.*, 1996) and specifically eggs of *Ascaris* and hookworm (Cram, 1943), *Schistosoma japonicum* (Jones *et al.*, 1947) and *Taenia saginata* (Newton *et al.*, 1949). Sand filters have demonstrated a great capacity for helminth egg removal. Schwartzbrod *et al.* (1989) reported 99% removal of helminth eggs (for secondary filtration) with effluent containing 10 eggs/l when high numbers of eggs (<900 eggs/l) were present in influent wastewaters. Sand filtration may also be effective in improving protozoan removal. Rose *et al.* (1996) reported that rapid sand filtration improved upon activated sludge removal by a further 7–10% with removal rates of >98% observed for the unit process. However, protozoa were still present in tertiary filtered effluents of around 4 (oo)cysts/l. Filtration is slightly more effective for *Giardia* removal than for *Cryptosporidium* (Rose *et al.*, 1996; Robertson *et al.*, 2000b).

Rates of removal for tertiary filtration units range from 97.9 to 99% for rapid sand filters (including coagulation/flocculation) and multimedia filtration (gravel, sand and carbon) with filtered effluents containing <1–18(oo)cysts/l and 0.1–2.4 (oo)cysts/l respectively (Madore *et al.*, 1987; Rose *et al.*, 1996; Carraro *et al.*, 2000). However, higher numbers of protozoa were reported in effluents from rapid sand filters treating activated sludge effluents in the UK (<10–435 (oo)cysts/l) and also from sandfilters treating trickling filter effluents (<10–60 (oo)cysts/l) (Bukhari *et al.*, 1997).

2.7.2 Disinfection

Parasitic eggs and (oo)cysts are extremely resistant to various forms of disinfection and few disinfectants are effective in inactivating parasites (Korich *et al.*, 1990). Eggs of *Ascaris lumbricoides* are very resistant to the effects of chemical disinfection (Krishnaswami and Post,

1968), which may be due to its relatively impermeable egg shell membrane (Wharton, 1980). However *Schistosoma* eggs are susceptible to chlorination at concentrations of 3.9–11 mg/l residual chlorine for 30 min (Jones and Hummel, 1947). Chlorination is particularly ineffective for removing protozoan (oo)cysts as free chlorine levels of up to 16 000 mg/l are required to inactivate completely *Cryptosporidium* oocysts (Smith, 1990), levels far in excess of dosages routinely used for wastewater disinfection of primary effluents (5–20 mg/l), secondary effluents (2–15 mg/l) and filtered secondary effluents (1–5 mg/l; Metcalf and Eddy, 1991). Chlorination is also less effective for *Cryptosporidium* removal than for *Giardia*. Rose *et al.* (1996) reported removal rates of 61% and 78% for *Cryptosporidium* and *Giardia*, respectively, after chlorination of filtered tertiary effluents. Disinfection of effluents was not effective since removal was only improved by 0.05–0.1% after chlorination and protozoan (oo)cysts were still present in 17–25% of samples of chlorinated effluents at levels of 1.8 *Cryptosporidium* oocysts/l and 1.0 *Giardia* cysts/l. McHarry (1984) reported similar concentrations of 0.8 *Giardia* cysts/l (0.3–1.2 cysts/l) after tertiary chlorination of trickling filter effluents. Chlorination of filtered effluent following advanced primary treatment demonstrated similar rates of removal of 67% for protozoan cysts after 1 h exposure to 10–14 mg/l (Jimenez *et al.*, 2001). Extended exposure improved removal to 89% after 4 h, but chlorination was not capable of complete cyst removal and removal rates of 94% were observed after 48 h (17 cysts/l). Protozoan (oo)cysts have also been detected in final effluents following UV and chlorination (range <0.26–7 oocysts/l and 0.3–104 cysts/l) and ozone disinfection of tertiary effluents; the viability of cysts was not determined (Rose *et al.*, 1996; Oswald *et al.*, 2000; Liberti *et al.*, 2000; Wiandt *et al.*, 2000). Spores of human microsporidia species also present in tertiary treated filtered and chlorinated effluent indicate that microsporidia can survive wastewater treatment processes including disinfection (Dowd *et al.*, 1998). Microsporidia spores may also be resistant to disinfection inactivation

since spore structure (Weber *et al.*, 1994 cited in Dowd *et al.*, 1998) is reportedly similar to that of *Giardia* spp. cysts and *Cryptosporidium* spp. oocysts, which are already known to be highly resistant to disinfection processes.

Ozone may offer greater potential for parasite inactivation than chlorination. Ozone, at initial levels of 1.1 mg/l, resulted in a 90% loss of viability for *Cryptosporidium* oocysts (Somiya *et al.*, 2000) and at levels of 4 mg/l is able to inactivate *Schistosoma mansoni* eggs (Mercado-Burgos *et al.*, 1975). However, ozone appears to have no effect on *Ascaris* eggs (Burleson and Pollard, 1976 cited by Reimers, 1989). UV irradiation has also shown potential for inactivating protozoan cysts by 90–99% depending on intensity and duration of exposure (Campbell *et al.*, 1995).

2.8 Natural wastewater treatment

Waste stabilization ponds (WSP) and constructed wetlands (CW) utilize physical and ecological processes to remove nutrients and pathogens. These natural wastewater treatment systems have been used as primary, secondary and tertiary systems to treat a variety of wastewaters worldwide.

2.8.1 Anaerobic lagoons

Anaerobic ponds are commonly used for the treatment of high strength wastewaters (e.g. agricultural) and can have high detention times of 20–50 days (Metcalf and Eddy, 1991). They are typically deep earthen ponds and are anaerobic throughout their depth. Partially clarified effluents are passed into other unit processes for further treatment. For domestic wastewater treatment, anaerobic ponds are usually used in series with a facultative pond followed by at least one maturation pond (see waste stabilization ponds below).

Information relating to parasite removal in agricultural anaerobic lagoons is limited. In one study, effective removal of protozoan (oo)cysts was reported for an anaerobic lagoon (60 days detention) treating dairy wastewaters (Karpiscak *et al.*, 2001). Removal rates of 99.99% for *Cryptosporidium* oocysts were

found with almost complete removal of 2.22×10^4 oocysts/l in influent wastewaters reduced to <10 oocysts/l in lagoon effluents. Ayres *et al.* (1992) reported that helminth egg numbers can be substantially reduced in anaerobic ponds (up to 99% removal) with detention times of <7 days. Anaerobic lagoons with retention times typically in excess of 50 days thus offer enhanced opportunities for sedimentation removal of parasite eggs.

2.8.2 Waste stabilization ponds

A variety of studies has shown that waste stabilization ponds (WSP) can effectively remove parasites from wastewaters, although wide variability in removal performance has been reported in the literature. All types of WSP (anaerobic, facultative and maturation) have demonstrated high rates of parasite removal, with the principal removal mechanism considered to be sedimentation facilitated by the long hydraulic retention times of 1–5+ days (anaerobic ponds), 5–40 days (facultative ponds) and 3–10 days (maturation ponds) (Feachem *et al.*, 1983; Ayres *et al.*, 1992; Mara *et al.*, 1992).

In a study of single ponds in India, Veerannan (1977) reported parasite egg removal rates of 62.9%, 88.5% and 93.3% for three systems (unknown retention times) with effluent concentrations of 56, 61 and 181 eggs/l. Protozoan removal rates of 84–100% and 94–100% for cysts of *Entamoeba* spp. and *Giardia* spp. respectively have also been reported for single ponds in India (Panicker and Krishnamoorthi, 1978b). However, generally, a higher efficiency for parasite removal is found for a series of ponds rather than single ponds with comparable overall retention times, although removal rates may still vary with parasite species.

In India, complete removal of parasite eggs and cysts was reported for three ponds in series with a total retention time of 6–7 days, although hookworm larvae (non-infective) were found in final pond effluents (Lakshminarayana and Abdulappa, 1969; Arceivela, 1970 cited Bouhoum *et al.*, 2000). Similarly, parasite eggs were completely removed after 9 days retention in

two ponds in Colombia (Madera *et al.*, 2002) and after 16 days retention in an experimental two pond system in Morocco; protozoan cysts were also completely removed (Bouhoum *et al.*, 2000).

Feachem *et al.* (1983) suggested that a well designed series of three or more ponds with a total retention time greater than 20 days would completely remove all nematode eggs and protozoan cysts. Grimason *et al.* (1993) suggested that for complete removal of protozoan cysts, a minimum retention period of 37 days was required. Mara and Silva (1986) reported that an 11-day, three-pond series comprising an anaerobic pond (1 day retention), and a facultative and maturation pond (each of 5 days retention) would remove most parasite eggs and achieve WHO (1989) reuse irrigation guidelines of ≤ 1 nematode egg/l, while Saqqar and Pescod (1992) suggested that 14 days were required to achieve the WHO quality criteria. Complete removal of parasites or attaining a final effluent quality of less than 1 parasite egg/l has been reported in several studies. Stott *et al.* (2002a) reported pond effluent compliance with WHO criteria after 10 days retention with reductions of parasite egg concentrations from around 1000 eggs/l in raw wastewater to 54 eggs/l after 5 days retention in an anaerobic pond (94.6% removal) and to 0.2 eggs/l after 5 days retention in a facultative pond (99.6% removal); eggs were not completely removed until after the second tertiary maturation pond (total of 20 days retention). Eggs of *Ascaris*, *Trichuris* and hookworm were found in anaerobic pond effluents, but only *Ascaris* eggs were present in effluent from facultative and the first maturation pond. Madera *et al.* (2002) reported that parasite eggs in raw wastewater (440 eggs/l) were completely removed after 12 days in a three-pond series incorporating an anaerobic pond (2.5 days), facultative pond (6.5 days) and maturation pond (3 days).

However, complete removal of parasites or achieving the nematode criterion is not guaranteed in multicelled systems. In a two-pond system in Sudan, only 90% of hookworm eggs were removed after 16.4 days total retention, although no *Ascaris* eggs were detected in the

final pond effluent (Klutse and Baleux, 1995). Hookworm eggs have also been detected in final effluent from a 17-day retention five-pond series (Mara and Silva, 1986), and from a 23-day, three/five-pond series (33–690 eggs/l), although *Ascaris*, *Trichuris* and *Giardia* were effectively removed (Ellis *et al.*, 1993). Nematode eggs were also found (6 eggs/l) in final effluent from a six-pond series with a total retention time of 25 days (Saqqar and Pescod, 1991), while protozoan cysts have been found in final maturation pond effluent at concentrations ranging from 0.1 to 90 cysts/l from a series of three and four ponds with total retention times of 18–40 days (Grimason *et al.*, 1996b).

A number of studies have shown that parasite eggs and cysts are detected with relative frequency in primary pond effluents and to a lesser extent in secondary and tertiary effluents. Grimason *et al.* (1996a) studied parasite egg removal in 10 WSP systems in Kenya where raw wastewater contained between 18 and 133 *Ascaris* eggs/l. Eggs were detected in 30% of primary anaerobic pond effluents which contained 3.6–88.9 eggs/l after 1–6 days estimated hydraulic retention (85.2–93.3% removal), while effluent from an 18.4-day primary facultative pond contained 0.7 eggs/l (98.9% removal). Effluent from a secondary facultative pond contained 2 eggs/l after 9.3 days retention (97.7% removal) but no helminth eggs were detected in any maturation pond effluents from any of the systems. In contrast, primary anaerobic, primary and secondary facultative and first tertiary maturation pond effluents contained protozoan cysts from pond series in France and Kenya (Grimason *et al.*, 1996b). Cysts were reduced from around 4×10^3 /l in raw wastewater to 225 cysts/l (primary anaerobic: 92.5% removal), 13 cysts/l (primary facultative: 99.7% removal), 2–120 cysts/l (secondary facultative: 46.7–84.6% removal), 70 cysts/l (first tertiary maturation: 41.7% removal) and 1–34 cysts/l (final tertiary maturation: 50–51.4% removal). Helminth eggs were completely removed in the Kenya pond system after the primary facultative pond unlike protozoan cysts, indicating that the removal of helminth eggs is not a reliable

indicator for protozoan cyst removal. Pond performance data also show that primary ponds (anaerobic and facultative) remove the greatest numbers of protozoan cysts, smaller oocysts of *Cryptosporidium* are not reliably removed in comparison to *Giardia* cysts and low numbers of (oo)cysts in influents are difficult to remove completely especially in secondary and tertiary ponds. Smith *et al.* (1992, cited Grimason *et al.*, 1993) detected *Cryptosporidium* spp. oocysts and *Giardia* spp. cysts in raw wastewaters in Brazil (60 oocysts/l, 1.1×10^4 cysts/l) in effluents from primary facultative ponds (15 oocysts/l, 75% removal; 490 cysts/l, 95% removal) and in effluents from secondary maturation ponds (25 oocysts/l; 0 cysts/l, 100%). Similar observations have been found for parasite egg loading and performance. Stott *et al.* (2002a) reported that effluent from a 5-day maturation pond still contained 0.1 egg/l despite influent containing only 0.2 eggs/l.

Parasite numbers in pond effluents vary widely in the literature. Helminth eggs in effluents can typically range from 1.5 to 89 eggs/l in effluents from anaerobic ponds, 0–88 eggs/l in facultative ponds, 0–0.1 eggs/l in first and second maturation ponds and 0 eggs/l in final maturation pond effluents (Mara and Silva, 1986; Ouazzani *et al.*, 1995; Grimason *et al.*, 1996a; Stott *et al.*, 2002a). Grimason *et al.* (1993) analysed raw and treated wastewaters from 11 pond systems in Kenya and detected protozoan parasites in raw wastewaters (13–73 *Cryptosporidium* oocysts/l; 213–6213 *Giardia* cysts/l), anaerobic pond effluents (2.3–50 oocyst/l; 133–231 cysts/l), facultative pond effluents (3.3–17 oocysts/l; 1.4–193 cysts/l), first and second maturation pond effluents (2.5–4.5 oocysts/l; 3.1–177 cysts/l) and final maturation pond effluents (0 oocysts; 40–50 cysts/l from one site).

The sequential reduction in parasite numbers with subsequent pond effluents indicates that removal (presumed by sedimentation) and/or destruction of parasite eggs and cysts is related to the cumulative hydraulic retention time. The majority of eggs and cysts are removed in anaerobic ponds, although lower percentage rates of removal are usually seen

probably due to shorter hydraulic retention times in comparison to facultative/maturation ponds. Ayres *et al.* (1992) reported egg removal in anaerobic ponds with 1–4 days retention of 77–98%. In comparison, facultative/maturation ponds with 3–18 days retention demonstrated parasite egg removal rates of 83–99.99%. The primary function of maturation ponds is usually pathogen removal. However, removal rates for parasite pathogens may vary considerably between tertiary maturation pond systems with removal rates for different parasite species ranging between 36 and 100% (Maynard *et al.*, 1999).

Ayres *et al.* (1992) developed a simple empirical model for predicting nematode egg removal as a function of hydraulic retention time (HRT). The lower 95% confidence limits of the model can be used to determine the number and retention times of ponds needed to reduce the parasite egg level in effluents to ≤ 1 nematode egg/l for restricted irrigation based on initial concentrations of nematode eggs in raw wastewaters. There are presently no design guidelines for protozoan removal.

Nematode Egg Removal %

$$= 100 [1 - 0.41 \exp (-0.49 \text{HRT} + 0.0085 \text{HRT}^2)] \text{ (Ayres et al., 1992)}$$

Using the model above, Ayres *et al.* (1992) predicted 1–2 day retention in ponds would achieve 75–84% egg removal, 3–5 days retention 89.8–95.6% removal, ≤ 10 days up to 99.3% removal and ≤ 20 days retention, up to 99.9% egg removal. Overall, 1–4 log units of reduction for protozoan and helminth parasites can be achieved in a well-designed series of WSP.

Multiple pond systems with sequential environments demonstrate higher parasite removal performance than conventional treatment systems (see Section 2.6.1). Parasite removal in ponds is attributed primarily to sedimentation facilitated by long HRT. Gravitational settling of free-falling helminth eggs with relatively fast settling velocities favours their partition from liquid to sludge (see

Section 2.5.1). The predominant removal of protozoa in primary ponds suggests that the principal removal mechanism for protozoan (oo)cysts is the adsorption/attachment of (oo)cysts onto settleable solids (Grimason *et al.*, 1993). Relatively high rates of removal can be achieved for eggs of *Ascaris* and *Trichuris* and sometimes for *Giardia* cysts, but WSP are less successful in removing hookworm eggs and *Cryptosporidium* oocysts. The removal efficiency of ponds can be affected by various factors retarding the sedimentation process and settling velocity of parasites. Ellis *et al.* (1993) reported that resuspension of parasite eggs due to gas production and floating scum passing through the ponds was a likely factor affecting the removal of hookworm eggs in their pond system. However, Lloyd and Frederick (2000) also implicate the use of aerators in the facultative ponds and pronounced short-circuiting (4.5 days hydraulic retention time compared to nominal 23 days) as a factor for the under-performance of the ponds. Yanez *et al.* (1980) also suggested that temperature inversions within ponds could affect protozoan cyst removal. Thermal stratification and accumulation of sludge are also thought to increase the likelihood of short-circuiting, thereby reducing the hydraulic retention time and facilitating the transport of parasites through the ponds (Bartone, 1985). Even if high sedimentation rates are achieved, parasite eggs can survive for at least 2–7 years in sludge (O'Donnell *et al.*, 1984; Schwartzbrod *et al.*, 1987; Nelson and Jimenez, 2000) and thus disturbance of sediments such as high inflow from storm events or wildlife, can potentially release high concentrations of parasites back into suspension.

Optimizing parasite reduction in pond effluents thus requires strategies to maximize sedimentation and minimize parasite resuspension and transport. WSP can achieve very high rates of parasite removal providing that hydraulic retention times are attained and that measures are taken to prevent hydraulic short-circuiting which can cause WSP to fail microbiological quality criteria for wastewater irrigation. Ayres *et al.* (1992) recommend that a larger number of smaller ponds in series is

used as a design approach to removing parasite eggs in WSP in order to minimize hydraulic short-circuiting effects and maximize removal. Other simple approaches suggested include the careful use of baffles to prevent breakthrough of solids and retain floatable scum and to increase hydraulic retention time (Shilton and Harrison, 2002) and non-shading pond windbreaks to reduce wind effects and short-circuiting in maturation ponds (Lloyd *et al.*, 2002).

2.8.3 Hyacinth and duckweed ponds

Macrophyte ponds have been utilized for wastewater treatment and show relatively effective removal of parasite eggs and cysts though removal performance is likely related to retention times of up to 1 week.

Mandi (1994) and Ouazanni *et al.* (1995) reported complete removal of parasite eggs in water hyacinth ponds with 7 days retention in Morocco when influent contained between 0 and 120 eggs/l. Performance compared favourably with that of facultative ponds with 50 days retention (also 100% removal) and was better than that for a 0.4 day retention anaerobic pond achieving 79–100% egg removal.

The removal of protozoa was size related in duckweed ponds treating secondary effluents (Falabi *et al.*, 2002). Cysts of *Giardia* were removed more efficiently than *Cryptosporidium* oocysts from wastewater with removal rates of 98% and 89% reported for a 9-day detention *Lemna gibba* covered pond. Influent concentrations were reduced from 16 *Giardia* cysts/l and 1.6 *Cryptosporidium* oocysts/l to 0.4 cysts/l and 0.2 oocysts/l respectively.

2.8.4 Advanced pond systems and high rate algal ponds

Advanced pond systems (APS) are an adaptation of waste stabilization ponds and optimize conditions for algal photosynthetic oxygen production to enhance treatment efficiency and reduce pond area. APS usually incorporate an advanced facultative pond followed by a high rate algal pond (HRAP) consisting of a shallow raceway around which

pond contents are circulated by a paddle wheel to maintain aerobic conditions (1–3 day retention), an algal settling pond (2–3 day retention) and two to three maturation ponds (3–5 day retention) for further disinfection. Variations of APS have used anaerobic digesters or anaerobic ponds instead of facultative ponds for smaller footprint systems and constructed wetlands instead of maturation ponds.

HRAP systems can demonstrate high rates of parasite removal especially when preceded by settling ponds/basins. El Hamouri *et al.* (1994) reported a removal of 99.3% for helminth eggs in a 36 m long HRAP system (including 24 settling) in Morocco. However, the majority of helminth eggs (96.4%) were removed from raw wastewater (113 eggs/l) in the preceding settling basin and, in isolation, the HRAP achieved a removal of 80% with effluent still containing 0.8 eggs/l despite low influent numbers from the settling basin of 4 eggs/l. Thus the HRAP improved egg removal by around 3%. A variety of eggs detected in the raw wastewater, including *Ascaris* and *Trichuris* (15% and 6% occurrence), were completely removed by sedimentation, but eggs of *Toxocara* spp. and *Enterobius* spp. passed through the system and were also present in the HRAP effluent. In a later study, a 100% removal rate was reported for a HRAP with an increased raceway of 788 m and treating effluent from a 4-day anaerobic lagoon that contained only 2 eggs/l (El-Hamouri *et al.*, 1995). Information relating to helminth egg removal in HRAP receiving high parasite loading rates and protozoan removal in these systems is still required in order to assess the capability of HRAP systems for parasite removal.

Egg removal processes were not investigated by the authors, though sedimentation and die-off were suggested as egg removal mechanisms. Some sedimentation of helminth eggs may occur in the HRAP despite continuous mixing of the pond contents and a surface water flow rate of 0.15 m/s. Settlement of helminth eggs (and protozoan cysts) has been found in wastewater flowing at 0.7 m/s along an 2 km open channel in Morocco (Bouhoum *et al.*, 1997). Parasites were reduced from 145

eggs/l and 1.6×10^5 cysts/l to 33.4 egg/l and 2.7×10^4 cysts/l after 2 km with greatest reductions found for *Ascaris* eggs (85%). Sediment samples showed that *Ascaris* eggs settled first at 80 m, but *Enterobius* eggs remained longer in suspension and were not detected until 560 m. Sediment samples at 80 m contained 9.6 eggs and 1.7×10^4 cysts/g yet, at 2 km, densities in sediments increased to 78 eggs and 1.4×10^5 cysts/g.

Parasite viability in HRAP ponds was investigated by Araki *et al.* (2000, 2001). Viability of helminth eggs (*Parascaris equorum*: horse roundworm) was reduced by 60% and 90% after 4 and 10 days exposure within HRAP ponds respectively. However, the authors also reported a 25% loss of egg viability in sterile water control suspensions after 10 days at 4°C (presumably refrigerated and in the dark) which suggests that the pond inactivation results for helminth eggs may be overestimated. Infectivity of *Cryptosporidium* oocysts retained in HRAP ponds was also significantly reduced by 97–99.9% after 3 and 10 days exposure, but some oocysts remained viable and infective since prevalence rates of infection in mice were 40% and 20% respectively. The authors implicated the effect of pH, ammonia and/or light as factors affecting inactivation (predation factors were excluded from the experimental studies). Ammonia levels in HRAP wastewaters (1.2–1.7N-NH₄ mg/l reported) are unlikely to have a deleterious effect on parasite survival. Biological wastewater treatment had no effect on *Cryptosporidium* oocyst viability (Robertson *et al.*, 2000b), although ammonia concentrations in effluents were likely to range up to 20 mg/l. Furthermore, very high levels of >1000 mg/l (levels far in excess of those reported in HRAP wastewater) are reportedly required for >99.9% inactivation of *Cryptosporidium* oocysts after a similar 8 days exposure (Jenkins *et al.*, 1998).

2.8.5 Constructed wetlands (CW)

Artificially constructed wetlands (or reedbeds) are of two main types – vertical and horizontal. Horizontal systems may be further classified depending on the pathway of water flow as surface and subsurface flow systems. Vertical

wetlands can be used as primary treatment processes, whereas horizontal systems are commonly used for secondary or tertiary treatment. Detention times in wetland systems are comparatively shorter than in WSP and are typically hours to several days.

Constructed wetlands have also demonstrated effective removal of parasites with horizontal subsurface flow (HSSF) gravel based systems usually more effective in parasite removal than equivalent soil based wetland systems (Rivera *et al.*, 1995). Greater rates of parasite removal are typically found for subsurface flow than surface flow wetlands. Gerba *et al.* (1999) reported protozoan removal rates of 69–88% in subsurface flow (SSF) wetlands compared to 58–73% removal in surface flow (SF) systems each with a retention of less than 4 days; SSF wetland effluents contained, on average, 0.01–0.07 (oo)cysts/l and SF effluents 4.9–16 (oo)cysts/l. Higher removal performance of subsurface systems was attributed to greater surface area for adsorption and filtration/straining. Parasite removal in wetland systems may also be related to size since cysts of *Giardia* are more effectively removed than smaller *Cryptosporidium* oocysts. Thurston *et al.* (2001) reported protozoan removal in HSSF receiving tertiary treated effluents of 87.8% for *Giardia* and lower rates of 64.2% removal for *Cryptosporidium*. Helminth removal in HSSF reedbeds in Morocco (1–4 h retention) was related to bed length (Mandi *et al.*, 1996) and season (Mandi *et al.*, 1998). Removal performance was greater with longer bed lengths of 50 m (95%) than 20 m (71%) and higher for all beds 20–50 m in the hot season (89.5%) than in the cold season (77.5%). In Egypt, HSSF reedbeds demonstrated helminth egg removal rates of 90% (0.02 eggs/l) in 50 m beds and 100% in 100 m beds also indicating that parasite egg removal is improved with increasing bed length and concomitant reduction in hydraulic loading (Stott *et al.*, 2002a). When challenged with artificially high numbers of eggs, equivalent to 100–500 eggs/l and a daily loading rate of $0.5\text{--}4 \times 10^4$ eggs/m²/day, 100 m HSSF beds demonstrated a substantial capacity (up to 100%) for helminth egg removal (Stott *et al.*,

1999). All eggs recovered from sediments were considered to be non-viable. Parasite cysts were less effectively removed and a few cysts of *Entamoeba* spp. were detected in 100 m bed effluents (Stott *et al.*, 1997). Generally, subsurface wetlands can achieve between 0.4 and 3 log units of removal in horizontal flow systems. There is currently no information available on parasite removal in vertical flow constructed wetland systems.

Sedimentation is considered to be one of the principal egg removal mechanisms in pond systems with long retention times of days, yet the role of this or other physical processes, such as filtration, in the removal of parasite eggs in CW with a short retention time of <1 day and horizontal water flow is unclear. Parasite eggs and (oo)cysts might be removed in wetland systems by entrapment and sedimentation within the bed and root matrix as wastewater percolates through the bed substrate. Wastewater and sediment profile analysis along 100 m reedbeds in Egypt has shown that helminth egg removal increases with distance along the bed with the majority of eggs removed within the first 10 to 25 m of the bed (Stott *et al.*, 1999). Particulate matter has also been found to accumulate predominantly within the first 10 to 20 m of the 100 m reed beds (Williams *et al.*, 1995) suggesting that parasite eggs (and cysts) in subsurface flow systems might be removed by mechanical straining and sedimentation as well as by similar processes described previously for particulate matter removal in trickling filters (see Section 2.6.1.2). Attenuation and entrapment within wetland biofilms with subsequent grazing by predatory fauna may further improve removal and inactivation, especially of protozoan (oo)cysts (Stott *et al.*, 2001).

The influence of vegetation in constructed wetlands on parasite removal is unclear. In surface flow systems, removal of protozoan parasites was improved in planted beds (95% removal) compared to unplanted beds (88–92% removal) for both *Giardia* and *Cryptosporidium* (Quiñonez-Díaz *et al.*, 2001). In soil based subsurface systems in Mexico, planted soil mesocosms showed a 50% improvement in helminth egg removal compared to unplanted

mesocosms (Rivera *et al.*, 1995). However, in a field-scale gravel based wetland trial in Egypt, vegetation had no apparent effect on helminth egg removal (Stott *et al.*, 1996). Vegetation in surface flow systems probably provides quiescent zones where sedimentation of parasites can occur. While the presence of hydrophytes in subsurface flow gravel beds may not be a significant factor for the removal of parasites, planted beds generally perform better than unplanted beds in removing other pathogens (Gersberg *et al.*, 1987).

Rates of removal in CW compare favourably with that in ponds and CW can provide complementary technology to WSP. CW usually have relative short hydraulic retention times (typically hours to a few days) compared to 20–30 days in ponds. Furthermore, the efficiency of sediment retention in CW is higher and thus there is less chance of resuspension. Slow water velocities and effective entrapment within short distances facilitate high rates of removal. Multicomponent hybrid systems may offer further opportunities for parasite removal and inactivation. Hybrid systems incorporating aquatic ponds and subsurface flow wetlands gave the greatest removal rates of *Cryptosporidium* (95.3%) and *Giardia* (99.94%; Gerba *et al.*, 1999) and hybrid systems can exceed rates of removal for ponds with similar retention times (Bavor *et al.*, 1987).

2.8.6 Overland flow/land application

Overland flow systems have been used as a polishing process for conventional and natural treatment effluents. Information on parasite removal is limited. An overland flow system consisting of three slopes (each 25 m long by 3 m wide and gradient of 4%) and receiving UASB effluent, demonstrated complete removal of parasite eggs when receiving a constant flow of influent containing 14 eggs/l (Chernicharo *et al.*, 2001). In another trial, slope removal efficiency was 99% under intermittent influent flow conditions with all species of helminths removed except hookworm. Slope effluent contained, on average, 0.2 eggs/l (maximum 2 eggs/l) with hookworm eggs detected in 18.5% of effluent samples. The authors conclude that the overland flow process

with application rates of 0.4–0.5 m³/m/h is a satisfactory post-treatment system capable of producing effluents with low concentrations of helminth eggs, although the slope length should be extended to at least 35 m for intermittent inflow operations.

3 PARASITE SURVIVAL IN WASTEWATER TREATMENT SYSTEMS

The survival of parasites in wastewaters depends on exposure to prevailing abiotic and biotic variables. Factors that affect survival include parasite species (structure of egg/(oo)cyst cell wall confers protection), stage of development (developed infective stage may be more resilient than undeveloped stages), temperature (longer survival at lower temperatures), humidity (longer survival at high humidity), pH (extremes reduce survival), sunlight (solar radiation can be detrimental to survival depending on exposure duration and intensity), protection by vegetation (from sunlight mediated disinfection), and presence of predatory fauna. Although survival of parasites can vary widely, typical survival times for protozoa in sewage are less than 30 days, while eggs of helminths (especially *Ascaris*) can reportedly survive for many months in manures, raw wastewaters and sludges (Feachem *et al.*, 1983; Schwartzbrod *et al.*, 1987). Protozoan (oo)cysts and helminth eggs are thus both likely to survive beyond typical retention times for treatment processes.

Sedimentation processes within conventional and natural wastewater treatment plants are unlikely to have much effect on parasite survival. Ambient temperatures in wastewaters (typically 10–25°C) and pH are unlikely to have a detrimental effect on survival of parasite eggs or cysts. *Cryptosporidium* oocysts can retain viability and infectivity after freezing (–15°C) and at temperatures up to 68°C (Fayer, 1994; Fayer and Nerad, 1996). Helminth eggs can also tolerate similar temperature ranges of –27°C to <65°C (Cram, 1943; Feachem *et al.*, 1983). Since helminth eggs and protozoan cysts can survive in wastewaters

for at least 1–2 weeks without loss of viability (Feachem *et al.*, 1983; Robertson *et al.*, 2000b), a retention time of several hours to days within sedimentation tanks will thus have a very limited effect on inactivation for parasites still in suspension. Anaerobic environments may also prolong survival of parasites. Parasites that are removed from suspension can remain in sludges for at least 1 year with very little loss of viability (Gaspard *et al.*, 1995; Gibbs *et al.*, 1995). The development of *Ascaris* eggs is suppressed at low oxygen concentrations (Brown, 1928) and at low temperatures of 4°C at least 50% of *Ascaris* eggs can remain viable after 33 months in sludge (O'Donnell *et al.*, 1984). Therefore, resuspension of sludge from sedimentation reactors and ponds can also release viable parasites back into effluents.

Conventional secondary biological treatment processes do not inactivate parasites. Field studies have shown that trickling filters, intermittent sand filters and activated sludge processes can promote embryonation of helminth eggs and the hatching of hookworm and *Schistosoma* eggs (Cram, 1943; Jones *et al.*, 1947). Protozoan (oo)cysts are also unlikely to be affected during treatment. Robertson *et al.* (2000b) reported no reduction in *Cryptosporidium* oocyst viability during aerobic biological treatment (activated sludge and trickling filters); between 72 and 77% of oocysts remained viable in raw, primary settled and secondary effluents. Robertson *et al.* (1999) also reported higher proportions of viable *Giardia* cysts in treated effluents (>20%) than in influents (<10%), indicating that wastewater treatment processes may selectively remove non-viable cysts.

Treatment in natural systems such as waste stabilization ponds (WSP) may have a limited effect on parasite inactivation, depending on the type and species of parasite and type of pond. Viable eggs of *Ascaris* have been reported in WSP effluents and may embryonate in aerobic conditions (Ayres, 1991). Detection of viable eggs in pond sludges by Nelson and Jimenez (2000) shows that eggs removed from influent wastewaters by WSP are not eliminated and

that pond sludges may serve as parasite reservoirs. Hookworm and *Schistosoma* eggs may also survive anaerobic WSP conditions, although cellular development and hatching may be significantly reduced or inhibited. Hookworm eggs can survive anaerobic conditions for 2 weeks without loss of hatchability (Ayres, 1991) and can pass into aerobic ponds where they may hatch into non-infective free-living forms, although development to infective stage larvae in either anaerobic or aerobic environments was not found in experimental studies. Similarly, *Schistosoma* eggs can hatch in facultative or maturation ponds (Kawata and Kluse, 1966), although ecological factors within facultative ponds can reportedly affect hatchability of *Schistosoma* eggs and infectivity of miracidia (Bunnag *et al.*, 1978). The maximum survival time of hatched *Schistosoma* miracidia is less than 10 hours, which is less than typical retention times in ponds (Kawata and Kluse, 1966). Ponds may therefore provide an effective barrier for schistosomiasis transmission. Information relating to protozoan viability in WSP wastewaters is extremely limited. Sporulated (i.e. potentially infective) oocysts of *Cyclospora cayetanensis* detected in 18% of samples collected from primary ponds indicate that WSP environments do not have a significant detrimental effect on protozoan survival either on development or on the infective stage (Sturbaum *et al.*, 1998). However, information relating to the viability of more ubiquitous protozoa in wastewaters such as *Giardia* and *Cryptosporidium* is needed.

Algae-based shallow pond systems such as HRAP, however, seem to have a deleterious effect on the survival of protozoan oocysts and perhaps to a lesser degree on nematode eggs. Experimental studies suggest that parasite inactivation in these systems is higher than in conventional treatment systems and may be rapid with exposure of only a few days required for significant reductions in protozoan oocyst infectivity. Further work is clearly needed to elucidate the contribution of physicochemical conditions within HRAP/APS systems and the consequences of stirring

pond contents on parasite removal and inactivation. It is interesting to note that activated sludge which also utilizes mixing processes, can remove protozoan (oo)cysts but may not appear to have any effect on parasite viability. All protozoan oocysts in activated sludge effluent from a model system were viable despite removal rates of 80% from influent wastewaters (Villacorta-Martinez de Maturana *et al.*, 1992). However, it has been found that dead oocysts show greater adhesion to each other and to debris (Bukhari and Smith, 1995), which thus may facilitate the removal of non-viable oocysts during secondary sedimentation, while the passage of viable oocysts through the treatment process is unaffected. In HRAP, mixing of the water column and shallow pond depths may facilitate sunlight mediated disinfection processes on parasite inactivation and die-off so that environmentally 'stressed' cells may be more susceptible to removal and destruction. Curtis *et al.* (1992) reported that light mediated destruction of faecal coliforms in WSP was dependent on oxygen and sensitive to elevated pH, physicochemical conditions that are found within APS and WSP systems. Ayres (1991) reported depth dependent effects on hookworm development and larvae die off in WSP with faster embryonation of eggs and significantly shorter survival of larvae at 10–20 cm below water surface of a maturation pond compared to depths of 0.5–1.1 m. pH, temperature and dissolved oxygen varied near the surface (pH 8–9; 6–20 mg/l DO; 23.5–26°C) but were constant and lower at deeper depths (pH 8; 2–6 mg/l DO; 23.5°C).

The effect of UV radiation and sunlight exposure on helminth inactivation can depend on the stage of egg development exposed. Undeveloped eggs and larvae are more susceptible to the effects of UV radiation and sunlight than fully developed eggs (Spindler, 1940; Tromba, 1978). Elevated pH within HRAP (and facultative ponds) is unlikely to have much effect on helminth egg inactivation and removal. *Ascaris* egg development is unaffected in alkaline pH <12 (Lewis-Jones and Winkler, 1991) and eggs of hookworm can hatch and develop to infective stages over the

pH range of 4.6–9.4 (Udonsi and Atata, 1987). The detrimental effects associated with photo-oxidation are likely to be limited for most helminth eggs.

Protozoa survive differently under similar conditions and appear more susceptible to environmental factors such as pH and solar radiation. Sunlight can significantly inactivate *Cryptosporidium* oocysts in comparison to dark environments (Huffman *et al.*, 2000), and UV irradiation and high pH can have a significant impact on viability of *Cryptosporidium* oocysts (Robertson *et al.*, 1992; Campbell *et al.*, 1995). Jenkins *et al.* (1998) suggested that pH indirectly affects oocyst survival by affecting oocyst wall permeability, the effect of which is time dependent. Elevated pH may therefore facilitate synergistic inactivation by other factors such as sunlight and UV exposure. Wiandt *et al.* (1995) reported seasonal variation in protozoan removal in WSP with highest removals of *Giardia* in spring and summer compared to relatively poor removal rates in winter, possibly due to sunlight exposure and inactivation. Further work is needed on elucidating the contribution of light (and dark) removal and die-off processes for parasites, especially protozoa and the influence of environmental factors, wastewater characteristics and process operation in mediating these effects.

Other processes likely to contribute to the removal and inactivation of parasites during wastewater treatment are predation and microbial attack. Ovicidal fungi have been shown to penetrate and destroy helminth eggs (Lysek and Bacovsky, 1979; Sobenina, 1978), while ingestion by insects (Miller *et al.*, 1961) and gastropods (Asitinskaya, 1979) can contribute to egg removal and inactivation. Predation by free-living protozoa and metazoa (surface associated and suspension feeders) may also play a role in the removal of protozoan (oo)cysts (Stott *et al.*, 2002b) where sedimentation and sunlight exposure are similar to that found for virus removal in biological wastewater treatment systems (Kim *et al.*, 1996). The ecological significance of invertebrates and microbial communities and their

grazing activities requires further investigation to determine the role of predation in removal and inactivation of parasites.

4 CONCLUSIONS

Parasite removal can vary considerably in wastewater treatment plants (WTP). Eggs and (oo)cysts of helminth and protozoan parasites have been detected in final effluents from a variety of WTP despite advanced levels of treatment. Factors affecting removal include:

1. the *type* and *level* of treatment (conventional and natural systems; primary, secondary, tertiary treatment)
2. *unit processes* (e.g. activated sludge, trickling filters, WSP and CW)
3. *parasite type and species* (helminth or protozoa; *Ascaris*, hookworm, *Giardia*, *Cryptosporidium* etc.)
4. *parasite loading* in influent wastewaters (Table 31.3).

Natural wastewater treatment systems with longer retention times are generally more effective at parasite removal than 'conventional' (mechanical) systems. In conventional WTP, removal of helminth eggs occurs predominantly during primary treatment. In comparison, primary sedimentation is relatively ineffective in removing protozoan (oo)cysts and (oo)cysts are preferentially removed during secondary treatment (including secondary clarification). Tertiary treatment plants, which usually incorporate filtration and chlorination, may improve removal but, unless influent concentrations are low, they will not completely remove parasites or produce effluents satisfying reuse criteria (0.1–1 egg/l; 1 cyst/40 l; Rose and Gerba, 1990; Blumenthal *et al.*, 2000). Generally, eggs of *Ascaris* and *Giardia* cysts are more readily removed in WTP than other helminth and protozoan species such as hookworm and *Cryptosporidium*, respectively.

The use of multiple stages with long retention times (days) and exposure to different environmental conditions (pH, DO, sunlight

exposure) in natural treatment systems, such as ponds and wetlands, appears to be more effective in removing and inactivating parasites. In waste stabilization ponds, reduction of helminth and protozoan parasites numbers occurs principally in primary ponds (typically anaerobic). Removal can be improved for helminths and, to a lesser degree, for protozoa, in sequential ponds. Parasite removal in constructed wetlands is also size related and wetland systems are especially effective for helminth egg removal. Increasing bed length and reducing hydraulic loading improves removal performance.

Although a variety of processes has been identified in the removal of bacterial and viral pathogens during wastewater treatment, there is a lack of detailed studies of parasite removal and inactivation mechanisms, and qualitative assessment of the relative importance of each. Parasite removal during wastewater treatment is likely due to a combination of physical, chemical and biological factors. Key removal mechanisms include:

1. direct sedimentation
2. solid assisted sedimentation and entrapment in flocs
3. 'filtration' through reactor substrates and attached biofilm or biological flocs
4. sunlight inactivation
5. grazing by predatory protozoa, metazoa and invertebrates.

Short-circuiting and buoyant sludge particles (e.g. in anaerobic or overloaded facultative ponds) will reduce removal rates and lead to passage of parasites into effluents. Enhancing opportunities for sunlight-mediated disinfection and natural 'die-off', e.g. by mixing and/or shallow reactors may offer improved inactivation, especially for protozoa.

Surveys indicate that 25–100% of wastewater effluents may contain protozoan (oo)cysts with densities ranging from <1 to 10³ (oo)cysts/l. Protozoan concentrations in treated effluents indicate that further wastewater treatment may be necessary for effluent discharge into receiving waters sourced for drinking water, especially during low flow in

TABLE 31.3 Summary of reported parasite removal and effects of wastewater treatment^a

Unit process	Parasite removal (%) mean and (range)		Effect
	Helminth eggs	Protozoan (oo)cysts	
Sedimentation:			Removal depends on operating conditions;
Plain	73 (35–96)	38 (4–68)	H & P: No parasite inactivation; H & P: Inactivation may
Chemical assisted	97 (95– >99.99)	70 (27–93)	depend on elevated temperatures, pH, and time
Anaerobic digestion	60 (5–90)	31 (0–50)	Removal and inactivation depend on time, temperature
Mesophilic			and parasite species; H: limited effect: 52% inactivation
			after 20 days; > 90% inactivation after 6 month;
			P: Some inactivation: 40–50% inactivation after 4 h,
			99.9% inactivation after 18–24 h; < 10% still viable
			after 18 days
Septic tanks	94.7 (85–100)		Anaerobic conditions may prolong survival; H: no effect
Imhoff tanks	97		
UASB	80 (70–90)		
Trickling filter	60 (5–90)	77 (5–93)	No effect on inactivation; H: promotes egg development,
(Complete works)			embryonation and hatching; P: promotes development
Activated sludge	92 (75–100)	87 (15–>99.9)	No effect on parasite inactivation; H: promotes egg
(Complete works)			development and hatching; P may promote excystation
Aerated Lagoon	83 (48–100)	85 (84–87)	
Oxidation ditch	93 (72–100)	81 (60–91.3)	
Filtration (Sand/ multimedia)	92 (78–99.6)	72 (40–99)	H: promotes egg development and hatching
Disinfection:	81	69 (61–78)	Removal and inactivation depend on contact time,
Chlorination			dose/intensity and parasite species; H & P: Very little
Ozonation			effect on parasite inactivation; P: >50% non-viable after
UV irradiation			2 min at > 0.3 mg ozone/ l; H limited effect: P: >90%
			non-viable at > 80 mW s/cm ²
Anaerobic lagoons	<99	99.99	Anaerobic conditions may prolong survival
Macrophyte ponds:	100	86 (69–98)	
Hyacinth Duckweed			
Waste Stabilization Ponds			
WSP (complete works)	99 (88–100)	98 (87–100)	Anaerobic conditions may prolong survival
Anaerobic pond	89 (58–100)	71 (32–98)	Aerobic conditions may promote egg development and
			hatching
Facultative pond	92 (50–100)	76 (16–99.9)	
(1–2) Maturation ponds	92 (50–100)	62.4 (8–100)	WSP: H: No effect
Final Maturation pond	99.8 (98.9–100)	83 (72–100)	WSP: P limited effect. Solar disinfection?
HRAP (complete works)	99.7 (99.3–100)		H: limited effect: 60% inactivated after 4 days
HRAP unit only	90 (80–100)		P: >97% inactivation after 3 days
Constructed Wetlands:			
Surface flow (SF)	89 (71–100)	85 (58–99.9)	
Subsurface flow (SSF)		80 (32–99.8)	

^a relates to information in text: H: Helminth; P: Protozoa

In most cases, removal rates may range from 0 to 100%. Parasite removal rates in the table refer to minimum removals where reported.

dry seasons. Furthermore, where wastewater is intended for reuse, it is advised that effluents should contain less than 1 parasite egg/l for public health protection. Restrictions regarding protozoan concentration in reclaimed wastewaters may also apply.

To meet these parasitological treatment objectives for effluent quality, tertiary treated conventional WTP or natural systems that are properly hydraulically loaded and designed should be sufficient. Conventional secondary WTP are unlikely consistently to meet these criteria when treating high influent loadings of $<10^3$ helminth eggs/l and 10^2 – 10^4 protozoan (oo)cysts/l in raw wastewaters and thus additional wastewater treatment may be required. Multicomponent wastewater treatment systems are more effective for parasite removal. Natural wastewater treatment systems have potential for offering flexible wastewater treatment systems in situations unsuitable for conventional plants and/or can be used as a component of hybrid, and modular systems in combination with other natural or conventional treatment processes where parasite removal is required.

Although information on parasite removal in wastewater treatment is available, further work is still needed. Understanding processes that cause parasite inactivation, the effect of mixing wastewaters on parasite removal and inactivation, multifactorial sunlight-mediated disinfection on parasite viability in illuminated surface waters and the role of grazing by predatory fauna would be of value. By fully understanding parasite fate and behaviour during wastewater treatment, the design of treatment systems can be improved and strategies undertaken to minimize the release of excreta-related parasitic diseases and safeguard public health.

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Problems in Wastewater Treatment Processes

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Activated sludge bulking and foaming: microbes and myths

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1 HISTORY OF BULKING AND FOAMING

1.1 Activated sludge bulking

When the activated sludge process was first developed in 1919 the sludge was described by Arden as 'flocculent in character', indicating its natural ability to settle well. This property of good settlement following a period of aeration allowed development of batch experiments into a full-scale process involving a continuous flow of wastewater and a continuous recycle of settled sludge from a subsequent settlement process. Poor settleability of activated sludge does not appear to be a widespread or serious problem during the early years of developing activated sludge to treat domestic wastewater.

However, in the next decade there are early signs of future problems. 'Undesireable filamentous organisms' were identified by Morgan and Beck (1928) as being responsible for poorly settling activated sludge. These filamentous growths were identified as *Sphaerotilus natans* and were stimulated by the presence of excessive carbohydrates in the wastewater.

A few years later, Donaldson (1932) described filamentous growths as 'the weeds of activated sludge' and firmly laid the blame for the cause of poor settleability at their presence. By this time two types of filament were identified, *S. natans* and *Leucothrix*. While this is now known to be an over-simplification, the mechanisms by which filamentous growths

affect activated sludge settleability was becoming known. The filaments physically keep the floc forming bacteria apart leading to a sludge of high water content.

The number of activated sludge plants in the UK increased rapidly after the Second World War, but sludge bulking did not present a serious problem except for plants with a high proportion of industrial wastewater.

The picture had changed dramatically by the 1970s. A survey by Eikelboom (1975) and others of several hundred activated sludge plants showed that 50% of them had problems at one time or another with bulking sludge. By this time identification of the causative filamentous organisms had also improved by Eikelboom's simple microscopic technique. By the 1980s at least 17 different filamentous bacteria had been separately identified in bulking sludges on three continents.

In the 1980s attempts were made to characterize these filaments into groups showing common morphological, physiological and metabolic similarity. This approach, by Wanner and Grau (1989), significantly incorporated operational conditions and problems:

- S *Sphaerotilus*-like oxic zone growers
- C Cyanophyte-like oxic zone growers
- A *Microthrix parvicella* types – all zone growers
- F Foam-forming filaments.

More recently, attempts have been made to characterize further specific filaments in

terms of their preferred conditions for growth and triggers for proliferation. This has finally led to a more scientific approach to reduction or elimination.

1.2 Activated sludge foaming

One of the earliest reports of foam formation is the so-called Milwaukee mystery (Anon., 1969). Surveys in the USA showed that 66% of activated sludge plants expressed some types of foaming as long ago as 1979 (Pitt and Jenkins, 1990). In a French study of some 6000 plants, over half had experienced foaming (Pujol *et al.*, 1991).

Foaming became more widespread in the UK in the early 1980s as reported in the South-East Dorset area of England (Foot, 1992). In this study foaming was not confined to any particular type, size or configuration of activated sludge plant and was liable to foam at any time. Biological foaming is mostly reported in nutrient removal and extended aeration plants (Pujol *et al.*, 1991).

Foaming appears to be connected to the presence of three types of filamentous microorganisms, as identified by Wanner in 1994.

- *Microthrix parvicella*, the most common (Fig. 32.1)



Fig. 32.1 Gram-positive *Microthrix parvicella* filaments coiled in typical 'bootlace' fashion. $\times 1000$ magnification. Courtesy of Dr E. Kocianova, Environmental Consultancy Services, Cornwall.

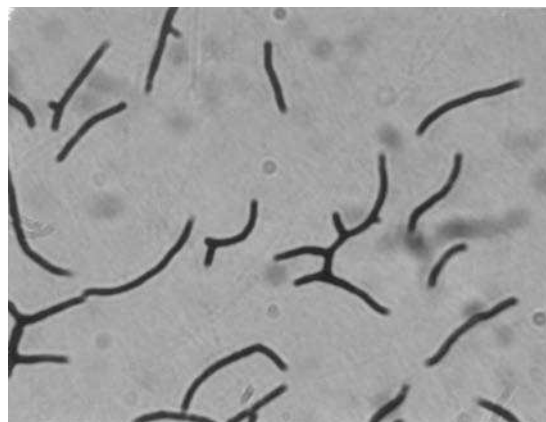


Fig. 32.2 Branched nocardiaforms grown in pure culture. $\times 2000$ magnification. Courtesy of Dr E. Kocianova, Environmental Consultancy Services, Cornwall.

- *Nocardia* (now called *Gordona*) *amarae*-like organisms, including *Rhodococcus* spp. (Fig. 32.2)
- *Nostocoidia limilocola*, Type 0041 and Type 0092 (Fig. 32.3).

Studies have shown that the distribution of filamentous organisms in biological foams at wastewater treatment plants varies from country to country, season to season and is changing with time (Wanner, 1998).

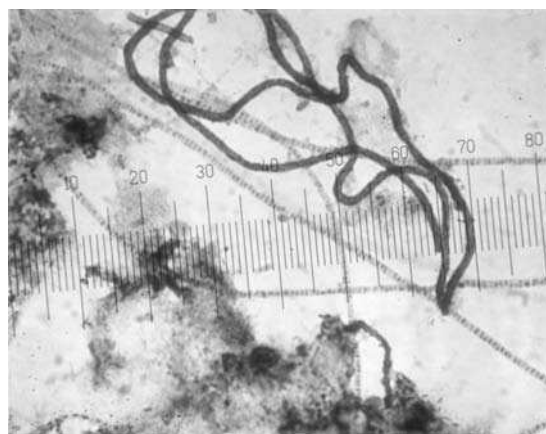


Fig. 32.3 Coiled Neisser positive *Nostocoidia limilocola* and extended filaments of Type O21N (or *Thiothrix* spp.). Courtesy of Dr E. Kocianova, Environmental Consultancy Services, Cornwall.

1.3 Anaerobic digester foaming

Foaming within anaerobic digesters is not uncommon, but is usually associated with physical causes, such as gas recirculation or simple overloading. In the latter, an excess of volatile fatty acids causes foaming. Such foaming is usually short-lived and dependent upon the continued presence of the volatile fatty acids.

However, in recent years, the presence of stable foams at times of acceptable digester loading rate has been observed. When the sludge has been investigated, filamentous bacteria have been found. In Stockholm (Dillner Westlund *et al.*, 1998) identified *Microthrix parvicella* as the causative agent of a stable anaerobic digester foam. Interestingly, the ability to create such a foam was eliminated when the feed sludge was heated for 5 minutes at 70°C. In this study, the source of the *Microthrix parvicella* was waste activated sludge. The same study also found that adding aluminium salts to the feed sludge caused the foam to collapse. An American study (Pagilla *et al.*, 1997) identified *Nocardia* as being responsible for anaerobic digester foaming in Sacramento. Again, the source of the *Nocardia* was waste activated sludge. A later study by the same authors (Pagilla *et al.*, 1998) concluded that chlorination of the waste activated sludge stream was ineffective for controlling such foams, even when dose rates were as high as 200 mg/l. Anaerobic digester foaming caused by filamentous bacteria appears to be a consequence of the presence of the same bacteria in waste activated sludge fed to the digester.

2 DEFINITIONS OF BULKING AND FOAMING

2.1 Activated sludge bulking

'Bulking' is a term referring to the phenomenon of a poorly settling sludge. Typically, suspended solids become visible in the bulk of a final settlement tank and, in more acute cases, will be lost over outlet weirs to pollute the effluent discharge. Ironically, chronic bulking can often be associated with high quality effluent as the sludge blanket captures and retains small floc particles.

Bulking should not be confused with the loss of suspended solids from physical causes. There are four main possible causes as follows:

- Failure of recycle RAS (return activated sludge) pump. This will allow solids to accumulate in the final settlement tank which will eventually be lost *en-masse* in a gross pollution of the final effluent.
- Excessive hydraulic loading, e.g. from failure of inflow control. This will usually be a problem in wet weather conditions only. However, an under-sized final settlement tank will exhibit similar characteristics.
- A high concentration of MLSS (mixed liquor suspended solids) caused by an insufficient wastage rate of SAS (surplus activated sludge). Gross pollution of the final effluent is likely if this is not resolved and is often accompanied by reduced nitrification due to the excess oxygen demand exerted by the excess mixed liquor solids.
- Denitrification within the final tanks, caused, for example by too long a solids retention time in the final settlement tank, e.g. failure of a final tank scraper.

Bulking sludges can be defined by reference to the measurement of sludge settleability or SSVI (stirred specific volume index) as shown in Table 32.1.

Normally settling activated sludge separates quickly from the supernatant liquor which will have an excellent clarity. When the SSVI approaches 120 ml/g then pin-floc solids can be observed in the bulk of the final settlement tank, often migrating to the peripheral weir. Unfortunately, pin-floc can also be caused by shock loads from excess sludge dewatering, trade effluents, or even over-aeration in

TABLE 32.1 Definition of sludge bulking by measurement of SSVI

SSVI (ml/g)	Condition
80–100	Normal
>120	'Solids' appear in final settlement tank
>150	On-set of bulking
>180	On-set of chronic bulking

the aeration tank. The division between blanket level and supernatant liquor becomes ill-defined because the sludge does not flocculate properly and hence does not compact. Often, this process is accompanied by denitrification, due to the prolonged solids retention time, and a surface film forms on the surface of the final settlement tank. Unfortunately, surface films can be caused by other processes, particularly foam-forming microorganisms.

There are two known biological causes of bulking, or poorly settling sludge:

- By far the most common is the presence of excessive growths of filamentous organisms.
- The remaining type is often referred to as non-filamentous bulking or zooglycal bulking. This occurs when common floc-forming bacteria produce extracellular polysaccharides which attract water, resulting in a sludge that has a high degree of hydration.

The incidence of non-filamentous bulking is comparatively rare. Activated sludge bulking has been observed world-wide and is a phenomenon affecting both municipal and industrial wastewater treatment plants. A survey of activated sludge plants in the UK in 1976 by Tomlinson revealed that 63% had experienced bulking.

2.2 Definition of activated sludge foaming

The presence of foam in an activated sludge plant is obvious; frequent points of generation or accumulation of microbiological foams are:

- On the aeration tank surfaces
- On the surface of final settlement tanks
- On the interconnection channels and chambers
- From pipeline chambers.

Biological foaming is often referred to as stable foam, scum or mousse. Biological foaming should not be confused with chemical foaming caused by the presence of excess surfactants

from trade effluent discharges or during the start-up of new plant.

Minor foaming is usually present in all activated sludge plants at one time or another.

Filamentous foaming is caused by the presence of foam-forming filamentous microorganisms in the activated sludge. When these filaments become dominant, foam is produced. The critical factors involved in the formation of foam are cell-surface hydrophobicity coupled with biosurfactant production and the presence of gas bubbles from denitrification or aeration.

These three factors combine to produce a stable three-phase system of air, water and cell material. In extreme cases, foaming can escape from its point of generation and spill into unwanted areas such as effluent channels, electrical equipment and on roads and paths. This is illustrated in Fig. 32.4. It is also a health hazard and risk to safety and effluent quality.



Fig. 32.4 A *Microthrix parvicella* foam illustrating the risk placed upon effluent quality and safety.



Fig. 32.5 A biological (nocardioform) foam similar in appearance to a detergent foam.



Fig. 32.6 Foam can be managed to an acceptable level and retained on the surface of the aeration tank.

One way to differentiate a detergent (surfactant) foam from a biological foam is from an estimate of the half-life (the time for 50% of the foam to degrade). A surfactant foam will have a typically short half-life of a few hours or less, whereas a microbiological foam will have a half-life of between 15 and 55 hours depending on the species. The effects of detergent foams are often spectacular and similar in appearance to the foam illustrated in Fig. 32.5. On this occasion the foam was biological in origin (*Rhodococcus* spp.).

It should be noted that the presence of biological foams is not necessarily related to settlement characteristics of the activated sludge. It is quite possible to have chronic foaming that can be kept in check by good management, without noticing any ill effect on sludge settleability or effluent quality (Fig. 32.6).

Sludge floc particles appear to be necessary to stabilize foam structure and hence give it a long half-life. As foam stabilizes apart from the main body of activated sludge biomass, further foam formation is limited only by the growth of foam-forming microorganisms. These provide a key to their control. De-selection of foam-forming microorganisms and removal or suppression of the foam will eventually prevent its proliferation.

2.3 Definition of anaerobic digester foaming

The presence of stable foam within an anaerobic digester is characterized by a foam layer up to 2.4 m deep on its surface. The foam will usually cause a spill of digested sludge onto walkways and the surrounding ground area if it occurs on a floating dome gas-holder design. The foam will enter gas draw-off pipework and compromise gas mixing systems and heating systems that rely upon biogas combustion.

Over-feeding or reducing the operating temperature can induce a similar, but less stable, foaming. In this case, the presence of volatile fatty acids will cause an unpleasant smell. If foaming is noticed with no unpleasant smell then the presence of filamentous bacteria should be investigated.

3 CAUSES OF BULKING AND FOAMING

Given the nature of the activated sludge process we are unlikely ever to be in the position to define the precise cause of bulking and foaming. The process itself forms the largest biotechnology industry in the world, providing economic treatment to an infinitely variable range of effluents and environmental conditions. It is not surprising, therefore, that for every report identifying a single cause there exists another, suggesting an alternative. The challenge for the microbiologist is first to dispel the myths surrounding activated sludge bulking and foaming and then to present a reasoned case to both operators and engineers for adequate controls and treatment capacity.

3.1 Common causes of activated sludge bulking and foaming

Sludge bulking and foaming is invariably associated with:

- Industrial effluents or domestic waste with a large proportion of industrial waste
- Plant loading
- Plant configuration
- Process control.

Throughout the world the filamentous organisms commonly associated with bulking and foaming are the nocardiaforms and *Microthrix parvicella* (Pitman, 1996).

3.1.1 Industrial and high strength waste

Little needs to be said about industrial waste other than to be aware that filamentous bacteria are capable of exploiting selective substrates. The science is well known and can be applied to new works design or nutrient dosing programme. Non-biological foams are generated by the introduction of surfactants and unused or unsuitable polyelectrolytes in recycle flows.

Even the best designed and operated treatment plant will occasionally suffer the effects of filamentous bacteria, often from external causes such as a sudden discharge of high strength waste (e.g. recycle liquors or a sludge

spill, use of glycol, sugar). When this occurs, filamentous growth is rapid and its effects long lasting.

Process air requirements also need to allow for sewages with variable treatability (alpha factor) caused by salinity, septicity and industrial waste.

3.1.2 Salinity

Saline intrusion into coastal towns can also result in poor sludge settleability. While the organisms in waste treatment are able to adapt to increased levels of chloride, the rate of change in chloride concentration through a spring tide can be more than activated sludge can accommodate. Acceptable limits would be a chloride concentration no greater than 6 g/l and an hourly change no higher than 50% of the average chloride concentration of the previous 24 hours. Large salinity swings cause dispersed floc and poor treatability. Chronic infiltration can cause chloride concentration to go from 0.1 g/l to 17 g/l within a matter of hours. Investing in the sewerage infrastructure gives additional cost benefits by reducing pumping, pump maintenance and overall treatment costs.

3.1.3 Septicity

Increased sulphate concentrations from saline intrusion also contribute to sewage septicity. Stormwater attenuation systems and long pumping mains cause sulphide to develop in the sewage, increasing malodours, concrete

corrosion and sludge production. Oxygen consumption can increase by 30% (Dormoy *et al.*, 1999). Septicity bulking follows the growth of sulphur-oxidizing filamentous bacteria.

3.1.4 Plant loading

Previous work on foaming and non-foaming plants shows that plants with a low aeration capacity (relative to the organic loading rate) are more likely to foam (Foot *et al.*, 1993). In Fig. 32.7, installed aeration capacity (as kg O₂/m³/day) is compared with the organic loading rate of the plant (as kg BOD/kg MLSS/day). Those plants that fall below the regression line are low loaded plants with an ample supply of oxygen, whereas those above represent plants with a high organic load but without a corresponding increase in the amount of oxygen available. Stable foam production occurs on most of the plants which are positioned above the regression line. Therefore, plants with a high F/M (food to mass) to O₂/m³ ratio are more likely to foam than those with a low ratio.

3.1.5 Plant configuration

The relationship between the settling properties of the activated sludge and the configuration of the aeration tanks is well established. As early as the 1960s plug-flow systems with a high substrate gradient were shown to have had better settling characteristics than completely mixed systems with a low substrate gradient (Water Pollution Research Laboratory,

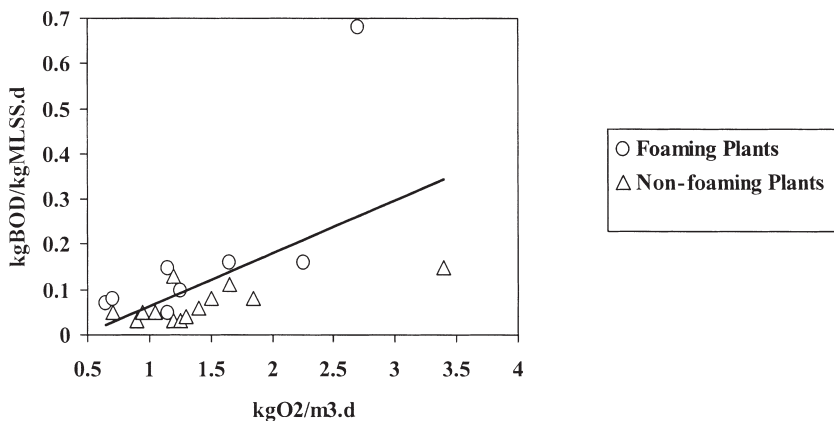


Fig. 32.7 Foam production as a function of F:M ratio and available oxygen. Adapted from Foot *et al.* (1993).

1967). Absence of a substrate gradient, initial contact zone (selector) or longitudinal mixing is invariably linked to poor settlement and foam formation. A substrate gradient can be achieved in both conventional plant and oxidation ditches where:

- The aeration tank is separated into compartments, or
- Flow is channelled.

Compartmentalization is key to control, with the initial compartments exerting the all important species selection pressures. These have been defined as kinetic (based on microbial competition) and metabolic (based on cultivation conditions), the principles of which are combined in the design of modern activated sludge plants to achieve good settling properties (Wanner, 1998).

3.1.6 Process control

Lack of process control leads not only to poor settlement and foam formation but also to inefficient treatment. In turn, inefficient treatment results in poor effluent quality and high energy costs. For example, a plant designed to treat peak flow over a 5-year design horizon could incur an increase in power consumption of 50% if operated without process control. SCADA (supervision control and data acquisition), PLC (process logic controller) or simple timer controls usually control plants' dissolved oxygen levels and sludge recycle rates. Insufficient knowledge of the treatment process will ultimately lead to treatment problems.

3.2 No effect without cause

As discussed previously, solids loss from a final settlement tank, caused by plant failure, can easily be mistaken for a bulking or foaming incident. The effects of a pump failure or blockage can look very similar to that caused by a bulking or foaming sludge (i.e. rising sludge blanket or floating solids on the final tank). Filamentous microorganisms are present in most activated sludges in

varying amounts and their presence does not necessarily mean that they are the cause of the solid loss. Correctly identifying the dominant filaments and getting a measure of their abundance is the first step in establishing this. Furthermore, an understanding of the mechanisms and drivers behind the predominance of filamentous microorganisms in the biomass will lead to control strategies for their eventual elimination.

4 ORGANISMS RESPONSIBLE: MECHANISMS AND DRIVERS

4.1 Taxonomic history

4.1.1 Diagnostics keys

The ecology, or balance of microbial species in activated sludge, has long been held as a significant factor in providing explanations for the attributes of the activated sludge biomass. With attention being drawn to filamentous species and the absence of diagnostic keys to identify these organisms, came an oversimplification of the nature of the problem and the organisms involved. *Sphaerotilus natans* (commonly known as sewage fungus) was aptly described as the 'weeds' of the activated sludge process and was alone blamed for activated sludge bulking. It was not until Eikelboom and van Buijsen (1983) devised an analytical technique for filament identification, that species other than *S. natans* became associated with bulking.

A similar pattern emerged with the arrival of the 'foaming' phenomenon, which also had a microbial origin. Like bulking, this problem is also associated with a dominance of filamentous species and originally it was also considered that a single species, *Nocardia amarae*, was responsible. Armed with additional diagnostic keys (Jenkins *et al.*, 1986), microbiologists set about identifying (and enumerating) a range of filamentous microbes associated with both bulking and foam formation.

Being in a position to identify individual types of filamentous microbe, patterns soon emerged, linking filaments with plant operation conditions (Table 32.2). While many had not yet been taxonomically defined (being

TABLE 32.2 Filamentous microbes in activated sludge grouped (Forster, 2000)

Conditions	Likely filaments
Low loading, extended aeration plants	<i>M. parvicella</i> (winter) 0092 (summer)
Conventional loading; full/partial nitrification	021N, <i>M. parvicella</i> , 1701, 0803, <i>H. hydrossis</i>
Conventional loading, no nitrification	021N, <i>S. natans</i> , <i>H. hydrossis</i>
Low DO	1701, <i>S. natans</i> , <i>H. hydrossis</i>
Septicity	<i>Thiothrix</i> , <i>Beggiatoa</i> , 021N
Nutrient imbalance	<i>Thiothrix</i> , <i>S. natans</i> , 021N
Low F:M	<i>M. parvicella</i> , <i>H. hydrossis</i> , 021N, 0031, <i>Nocardia</i> , 0675, 0092, 0961, 0803

know only by type rather than species or genus), this work provided vital clues to cause and, therefore, control mechanisms.

Comparisons drawn between filament populations associated with bulking and foaming events soon identified a hard-core of filaments. Rankings based on these reports have

shown similar patterns as shown in Fig. 32.8(a) and (b). However, rather than clarifying the situation these data show that (although some species have a unique function), there is a small group of filaments that can induce either bulking or foaming, i.e. they appear in both bulking and foaming sludges. One of these is *Microthrix parvicella*, which is seen to be just as common in bulking sludges as Type 021N, in addition to being dominant in almost half the foaming sludges. While not complete, the data are indicative and show how some species are capable of exploiting more than one set of operating conditions.

4.1.2 Genetic classification

Mirothrix parvicella was one of the first to be defined using genetic techniques (Blackall *et al.*, 1994). Eikelboom type 021N became linked to *Thiothrix* (Howarth *et al.*, 1999) and type 1701 to *Sphaerotilus natans* (Howarth *et al.*, 1998) using 16S rRNA sequencing. Nocardioforms were

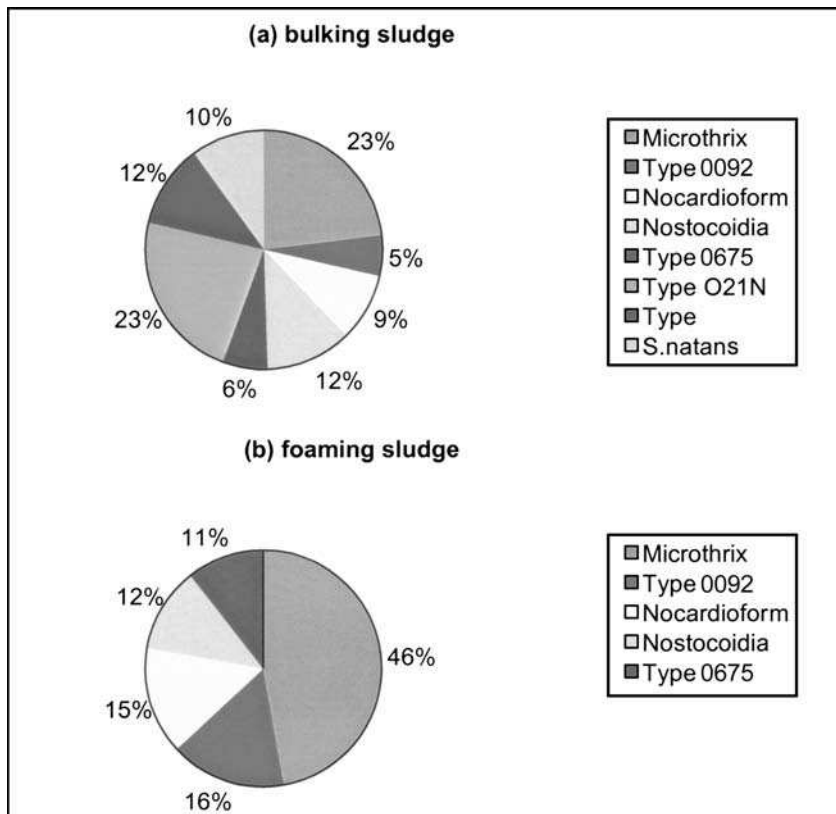


Fig. 32.8 Ranking of filamentous microorganisms in (a) bulking sludge and (b) foaming sludge.

placed under similar scrutiny using cluster analysis of their morphological and physiological characteristics (Soddell and Seviour, 1998).

With this approach comes the use of acronyms (e.g. NALO for *Nocardia amarae*-like organisms, PTLO for pine tree-like organisms and GALO for *Gordona amarae*-like organisms) underlining a desire to become technically correct when describing a species.

While essential for research into bulking and foaming, the diagnostic tools available to water industry practitioners (microscopes and stains) are much less sophisticated, but still fit for purpose. In this application the skill lies in knowing what first to look out for, appreciating the significance of subtle changes and then being able to act on that information.

4.2 Mechanisms and drivers

4.2.1 Selection pressures

The principles set out in Tomlinson and Chambers' (1984) benchmark paper 'Control strategies for bulking sludge' demonstrated how the substrate model developed by Chudoba and his co-workers (1985) worked in practice. Understanding the physiometabolic grouping of filamentous organisms responsible for bulking and foaming (Wanner and Grau, 1989) and the definition of the physicochemical properties of foam-forming sludges (Kocianova *et al.*, 1992) has gone some way to explaining how scum forms and control strategies work. Then, by quantifying design criteria for initial contact zones (Albertson, 1991) came the means to design selectors specific to the organisms involved (Foot and Forster, 1997; Wanner, 1998).

4.2.2 Bubbles and buoyancy

Biological foam on activated sludge plant consists of gas bubbles held in a dense matrix of fibrous material. Filamentous microbes form the basis of this structure which, when compared to sludge of a similar density, have a greater viscosity (Foot *et al.*, 1993). Therefore, the network of intermeshed microbial filaments within the foam layer provides structure, stability and buoyancy to flocs intended for use as a suspension, in a mixed reactor.

The relationship between the sludge ecology and foam formation is described thus:

- The affinity foam-forming nocardioforms have for hydrophobic substrates (Soddell *et al.*, 1998) explains why, once filamentous microbes exploit the foam environment, their domination becomes absolute.
- Subtle changes in the composition of the mixed liquor may provide the initial trigger to stimulate foam formation (Stratton *et al.*, 1998).
- Foams also have the ability to denitrify depending upon the dominating filamentous microbes and the type of substrate (Wanner *et al.*, 1998) and associated nitrogen gas would further increase the separation process.

4.2.3 Bridging

Bridging describes the physical separation of floc by filamentous microbes, which extend beyond the floc into the aqueous phase of the liquor. While surrounded by floc-forming bacteria the filamentous microbes provide structure to the floc but excessive filament growth within the floc leads to a dispersed floc. Bridging also places filamentous microbes in a position of advantage, where they are able to absorb substrate (sewage) in preference with little competition from the floc-forming bacteria.

4.2.4 Exploitation

The selection of filamentous microbes is to a large extent dependent upon the nutrient and metabolic conditions placed upon the biomass by the process itself. Eliminating the controllable variables such as oxygen, mixing patterns and solids retention time leaves us to consider temperature and substrate.

Species such as *M. parvicella* are known to predominate at certain times of the year (Wanner, 1998) but, unfortunately, little can be done about this other than have a strategy to deal with it.

Substrate appears to offer the greatest potential for control as selective substrates have long been associated with microbial selection, in particular surfactants. Recent investigations into micronutrient supplements

(Burgess *et al.*, 1999) returns research to an area better understood by biotechnologists with the potential to explain trigger mechanisms and enhanced treatment performance.

5 TREATMENT STRATEGIES

5.1 Background

While the types of filamentous organisms responsible for bulking and foaming appear not to have changed in recent years, treatment standards and waste treatment processes have. Anticipated problems associated with biological nutrient removal have not yet materialized in the UK as they have done elsewhere in the world because of the general adoption of chemical phosphorus removal. However, achieving compliance with UWWTD (Urban Waste Water Treatment Directive) standards has presented a new challenge when providing full treatment on restricted sites in coastal areas. The further requirement to disinfect final effluents discharging close to bathing waters extends the need for consistent treatment to a high standard. Given the robustness of the process, it is with little surprise that activated sludge (and its variants) remains a popular treatment option.

Treatment plant with suspended biomass, SBRs (sequencing batch-reactors), fixed film biomass BAF (biological aerated filter) plant, SBCs (submerged biological contactors) and more recently MBRs (membrane bioreactors) with a combined fixed and suspended biomass have been used in this application. Table 32.3 explores the relative merits of these plants.

Just as increased urbanization led to the development of conventional activated sludge processes in the middle of the last century so

has the demand for high quality effluent treatment today resulted in the development of more intensive processes. Sewage treatability has also changed. For example, in coastal areas there is a likelihood of increased chloride levels, increased septicity and increased levels of industrial waste. Unless controlled, these factors will increase the tendency to sludge bulking and foaming.

Some plants remain unaffected by the symptoms of bulking and foaming, whether by design or by accident. The industry would suggest that control features now being incorporated into new plant designs avoid features that would have previously made plants vulnerable to bulking and foaming. Our challenge is to ensure that current experience influences future plant design.

5.2 Conventional sludge bulking and foaming control

5.2.1 Managing expectation

Tipping (1995) concluded, in his operator's overview of the activated sludge process, that 'The operational and ecological factors which are responsible for inducing microbial growth, sufficient for foam formation, are largely unresolved and likely to be site specific'. This resulted in the frequent application of physical control measures along with the expectation that the problem would likely recur. While the temptation exists to 'bend the biology' of activated sludge by expecting more from the process than it can deliver, what we now have is a greater awareness of the requirements (and hence limitations) of the organisms involved in waste treatment.

TABLE 32.3 Design features of treatment plant

	<i>Conventional</i>	<i>SBR</i>	<i>BAF</i>	<i>SBC</i>	<i>MBR</i>
Primary treatment required	✓	X	✓	✓	X
Final tanks required	✓	X	X	✓	X
Disinfected effluent	X	X	X	X	✓
Relatively easy to cover	X	✓	✓	✓	✓
Simple controls	✓	X	X	✓	✓

5.2.2 Solution pathway

Conventional wisdom applied to resolve a sludge bulking or foaming event has been to:

- Review operating and control procedures and make required corrections
- Employ short-term remedial action if necessary.

And if the problem persists:

- Review plant design parameters
- Implement design changes.

The first of these two steps requires an initial understanding of the process and the key performance measures. A greater understanding of the process is obtained when both cause and effect are measured, thus leading to an overall improvement in performance. Only when this initial cycle fails does it become necessary to take the second step of reviewing plant design. Improved awareness of the requirements and limitations of activated sludge organisms gives structure to the control measures employed and direction to the recommendations made for plant design.

This approach has formed the basis for improvements to plant design and operation in resolving persistent bulking and foaming problems. However, if significant changes occur in the incoming sewage or the process control, then bulking and foaming will still happen, irrespective of how well the plant is designed or operated. It is therefore important to know the relative efficiency and the cost of available solutions.

5.2.3 Common cures

Given that bulking and foaming can seriously compromise effluent quality, it is essential to reduce the risk of discharge consent compliance failure as soon as possible. Table 32.4 illustrates the most commonly applied bulking and foaming control measures for conventional activated sludge plant along with an estimated response time and cost. Not all solutions apply to every type of plant (e.g. importing MLSS to large plants), nor are some of them sustainable for prolonged periods of time (e.g. increased RAS pumping and chlorine dosing). Response times also vary depending upon the nature of the activity. While control changes can have an immediate effect on solids loss, measures exerting selective pressures on the biomass (selector or MLSS reduction) can take at least one cycle of the solids retention time (sludge age) to take effect.

5.2.4 Best working practice

Building your own knowledge database is relatively easy to do and essential to assure that best practice is employed when first assessing the nature of the problem. Making sure that this approach is transferred through time and to new processes is more difficult to achieve. However, problems can be resolved by different means, sometimes novel and sometimes with little reference to perceived wisdom.

5.2.5 Better by design

Good aeration plant design is only the first step in the elimination and prevention of activated sludge bulking or foaming. Basic principles include:

TABLE 32.4 Summary of commonly used cures for sludge bulking and foaming

	<i>Response period</i>	<i>Cost</i>	<i>Application</i>
Reduce MLSS	1–3 weeks	Nil	all plant
Import MLSS	Immediate	£0.50 per capita	small plant
Increase RAS rate	Immediate	Nil	all plant
Nutrient dose	Immediate	£1.00 per trade p.e./annum	all plant
Chlorine dose	1–10 days	£0.01 per capita	large plant
Sprays	Immediate	£3k per tank	all plant
Selector	1 month	Site specific	all plant

- Attention to the layout of new designs (e.g. avoiding complete-mix systems in favour of plug-flow) to promote the growth of good settling bacteria.
- It has also been observed that plants with insufficient aeration capacity to cope with diurnal load variations or breakdown of aeration equipment are more prone to outbreaks of filamentous bacteria.
- Ability to return a maximum of 150% dry weather flow as RAS (return activated sludge) is also important in counteracting the effects of poorly settling sludge.
- Spray bars fitted to final tank scrapers are also useful for dispersing and avoiding the proliferation of foams.
- Final tanks also need to be large enough to cope with a peak SSVI of 120 ml/g. However, if selector zones are incorporated into the plant, then final tanks can be sized to a lower value of 100 ml/g.
- Selectors designed purely to discourage filamentous growth would have an average retention time of between 10 and 20 minutes (Wanner, 1998), while tanks designed to remove nitrate would have a much longer retention time.

Success rates for selectors vary depending on the application and ability to document accurately the effect due to plant modifications that are invariably linked to construction work. The design favoured by many has a high F:M (food to mass) loading rate of 12 kg BOD/kgMLSS/day in the initial contact zone and 3 kg BOD/kgMLSS/day overall.

5.2.6 Better by operation

While good design will increase the tendency towards a good, settleable activated sludge plant, operators need to be given flexibility to change operating parameters as conditions demand. Adjustable dissolved oxygen targets, avoidance of prolonged over-aeration, variable RAS and SAS rates, control of final tank blanket levels, etc. are all useful tools for operators to employ to reduce the effects of foaming and bulking. Standard operational process changes can be very effective in combating filamentous

growths if they occur. These would normally be to:

- Reduce mixed liquors concentration (to avoid solids loss)
- Increasing/decreasing dissolved oxygen levels (to correct under/over-aeration)
- Increase RAS rate (to reduce final tank blanket levels)
- Taking final tanks out of service (to reduce the effects of denitrification).

There is often the temptation (and pressure) to carry out all these changes at once, but this makes it difficult to evaluate any effects. As these changes act on the biomass slowly, they are often (on their own) insufficient to prevent a series of effluent discharge consent violations.

One fallacy is that chemicals will provide a permanent quick fix. Chemicals employed at wastewater treatment plants to combat the effects of filamentous bacteria include settlement aids such as polymers and coagulants. Examples of the latter include, iron and aluminium compounds and even talcum powder! Bio-augmentation with enzymes and other cultured bacterial mixes have also been on trial, but the results are often inconclusive. Inorganic chemicals, such as bleach and hydrogen peroxide, are also proven in disrupting bacterial filaments. A timely application of chlorine or bleach (Jenkins *et al.*, 1986) encourages a rapid return to the biomass equilibrium associated with a well-designed and operated plant.

Control strategies for foaming in anaerobic digesters follow similar patterns:

- Dosing with poly-aluminium salts and chlorine
- Mechanical disruption
- Decrease feed rate
- Lower reactor level.

5.3 Solution review

Reviewing sludge bulking and foaming control for conventional activated sludge processes, it is apparent that new control measures come from a better understanding of the sludge

TABLE 32.5 Bulking and foaming control for treatment plant

	<i>Conventional</i>	<i>SBR</i>	<i>BAF</i>	<i>SBC</i>	<i>MBR</i>
Reduce MLSS	✓	✓	✓	n/a	✓
Import MLSS	✓	✓	X	n/a	✓
Increase RAS rate	✓	X	X	n/a	X
Nutrient dose	✓	✓	✓	✓	✓
Chlorine dose	✓	✓	X	n/a	✓
Sprays	✓	✓	✓	n/a	✓
Selector	✓	✓	X	n/a	✓

Note: n/a = control not applicable.

microbiology. When all filaments in activated sludge were generically termed 'sewage fungus', little could be done to exploit the metabolic preferences of the organisms comprising that group. Thankfully that is not now the case and the operator has a range of options to consider.

Looking at the new processes now employed in treating sewage from coastal areas, it can be seen that not all of the old solutions employed to combat bulking and foaming are appropriate or even necessary (Table 32.5). While combined fixed film and suspended biomass systems can experience forms of sludge bulking and foaming the absence of some process flows (e.g. sludge recycle) reduces the options available for control.

The activated sludge process has been suitably challenged following tighter effluent quality standards such as N and P (nitrogen and phosphorus) removal and disinfection. Irrespective of how well the plant is designed, problems will occur. Conventional methods for the prevention or elimination of sludge bulking

and foaming have proved adequate provided that:

- Decisions are made in a logical order
- Contingency plans are in place to offer a quick response
- The treatability of sewages are correctly assessed before designing new plant
- Salinity and septicity of sewages from coastal sites are brought within reasonable limits
- The treatment process operating procedures are flexible
- More use is made of existing data to identify problems before they impact on plant performance.

6 CASE STUDIES

Examples of bulking and foaming in the South East Dorset area of the UK demonstrate how these problems can be resolved. Table 32.6

TABLE 32.6 Summary of bulking and foaming case histories in the UK

<i>STW design</i>	<i>Problem</i>	<i>Cause</i>	<i>Solution</i>
Surface aeration	Filamentous bulking	Excessive aeration in anoxic zone	Chlorine dose
Fbda	Microthrix foaming	Excessive aeration in anoxic zone	Reduce aeration
Fbda	Biopolymer production	Low DO levels	Spray bars, intensive aeration
Ditch	Nocardia foaming	Digested brewery waste	Chlorine dosing
Surface aeration	Microthrix bulking	Sludge spillage	Chlorine dosing
Ditch	Filamentous foaming	Overloaded, low DO	Additional aerators
Surface aeration	Non-settling sludge	Metal dust discharge	Re-seed
Package plant	Flow bulking	Hydraulic flushing	Re-seed and tankering
Surface aeration	Filamentous bulking	No known cause	Selector zone and spray bars
Ditch	Filamentous bulking	No known cause	Selector zone and spray bars
SBR	Flow bulking	Poor hydraulic design	Reduce mlss and process review

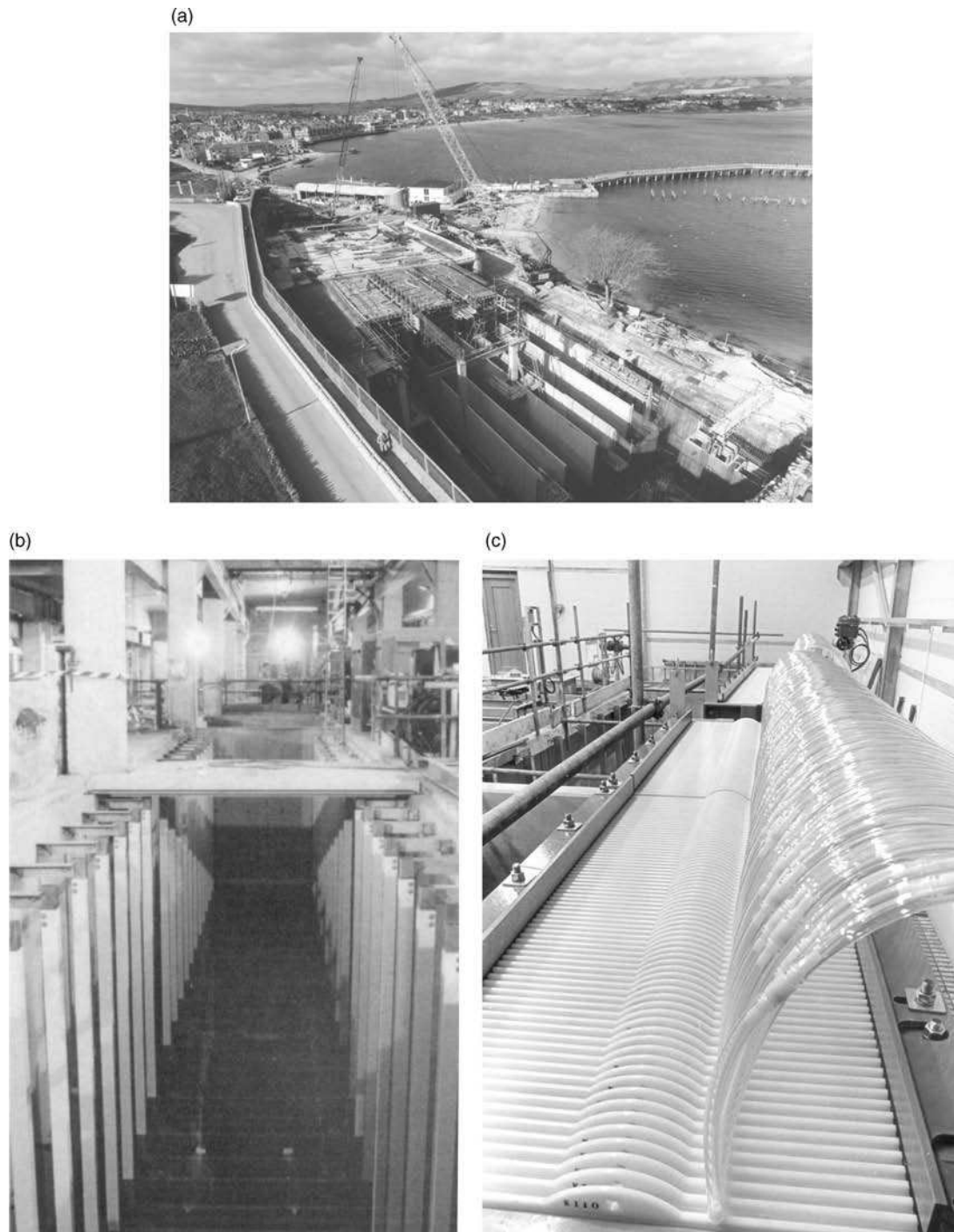


Fig. 32.9 Swanage MBR plant showing, (a) site excavation, (b) empty aeration tank with diffusers but no membranes, (c) membranes and connecting pipework.

summarizes how recent bulking and foaming events have been tackled to ensure compliance with effluent quality standards.

More recent experience with secondary treatment on UWWTD (urban waste water treatment directive) sites has led to an awareness of several important issues not yet fully understood.

Space restrictions for constructing wastewater treatment plant on the coast are illustrated in Fig. 32.9(a). The area under excavation defined the limit of the land available to install secondary treatment to a resort town on the south coast of the UK. Proximity to bathing beaches required the effluent to be disinfected. The MBR plant (19 000 connected population) is now complete and enclosed. Aeration tank detail includes diffusers positioned along the length of the channel (Fig. 32.9(b). Fig. 32.9(c) illustrates the type of membrane used and the associated pipework through which treated effluent would pass.

Compact plant designs make good process control an essential prerequisite. For example, taking tanks out of service for re-seeding an SBR plant with activated sludge is not a viable option as both bioreactor and settlement capacity are lost. Similarly, an MBR and BAF plant require lead-in time to build up sufficient biomass or seed the media. Not that these processes are inflexible rather, remedial controls should be incorporated into the design as a precautionary measure. Installing spray bars and chlorine dosing lines is more easily done during construction, chlorine is often available for odour treatment or chemical cleaning.

7 FURTHER DEVELOPMENT

7.1 Potential options for the future

Understanding the interactions between cause and effect in the activated sludge process is not easy, especially when there may be a dozen or so reasons for a bulking or foaming incident to occur. The process represents the largest biotechnology industry in the world and the combinations of inputs, control variables and

configurations are daunting. Mapping the processes involved is seen as one way of putting structure to the apparent chaos of the operating process. Operating parameters are often documented within procedures (held in site manuals) and the implementation of changes to the plant is controlled by standard work instructions. The discipline of managing these activities is fundamental to the efficient running of treatment plants and control of quality and costs.

7.1.1 Process mapping

By way of an overview, the process map presented in Fig. 32.10 illustrates how the attributes of a plant, such as influent, is linked to operating variables such as flow strength and composition. Bulking and foaming symptoms are then placed against each of these variables along with the suggested solution to resolve the problem. Controls, system configuration, and treatment capacity attributes have also been applied to the same matrix of conditions, symptoms and solutions. While not exhaustive in terms of detail the process map offers a mechanism by which both:

- The myths behind the science can be dispelled and
- The gaps in our knowledge can be filled.

The next logical step would then be to review improvements that have been made, measure them in some way and incorporate them into new working practices or designs.

7.1.2 Process modelling

Process models have been used for many years and models (such as STOAT and GPSX) are readily available and often used for new works designs. Modelling is particularly relevant when an existing plant is to be extended and its performance assessed. However, to minimize the risk of bulking and foaming, process models need to be capable of defining the risk or probability for this occurring.

7.1.3 Performance management

Two areas where advances have been made are in that of process monitoring and information

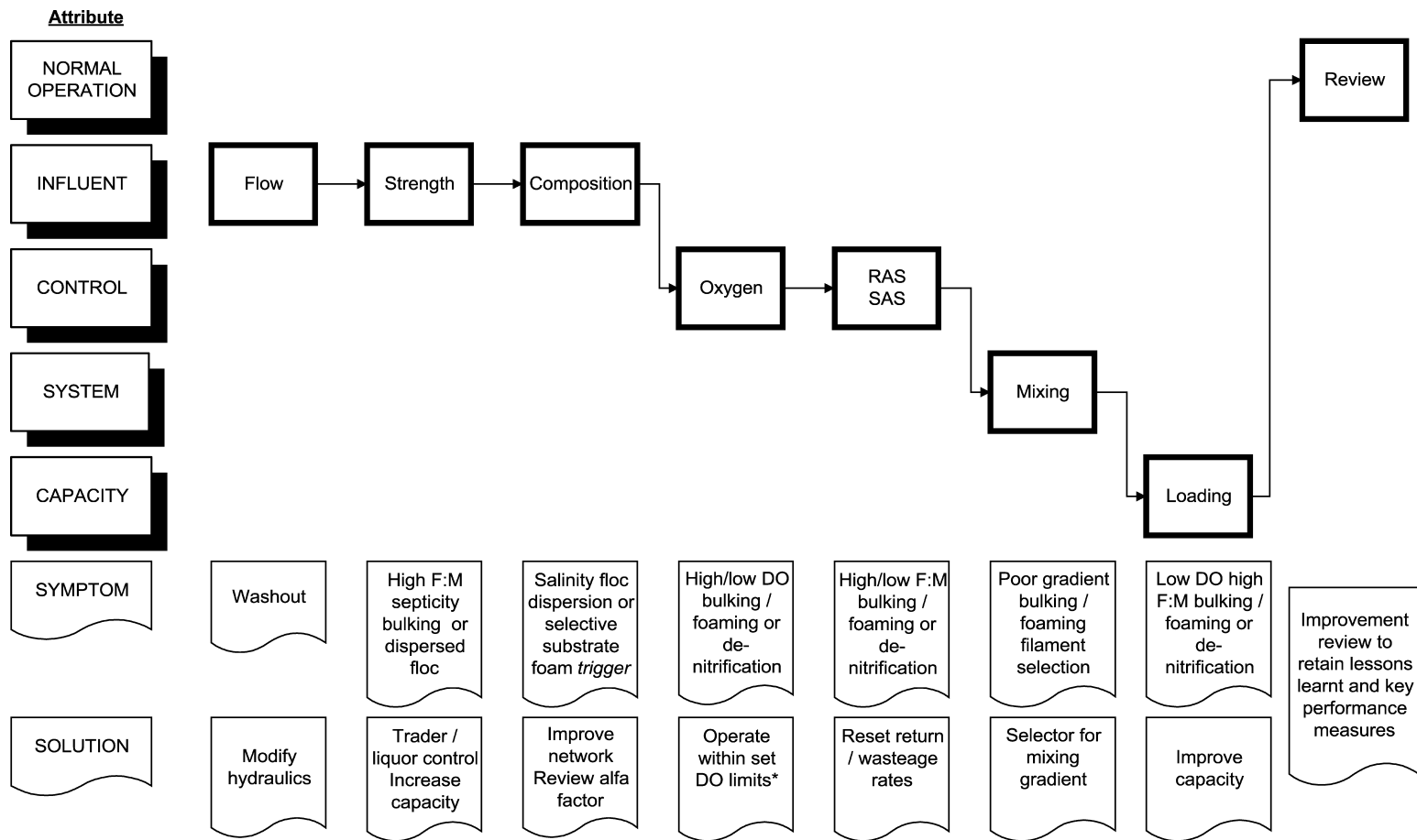


Fig. 32.10 Process flow for the control of bulking and foaming.

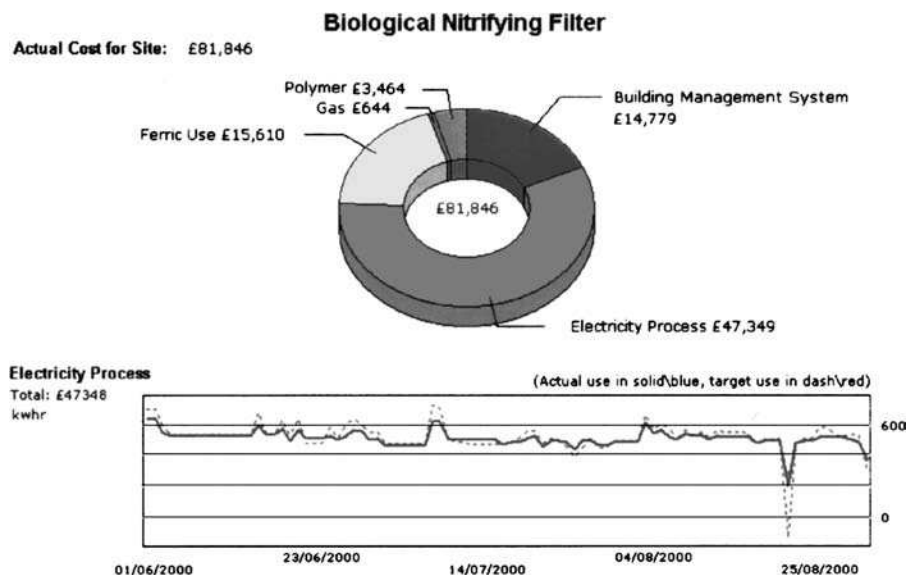


Fig. 32.11 The relative costs of operating a modern waste treatment process. Courtesy of Meniscus, UK.

networks. It is now possible to measure plant performance against benchmark values and treatment quality to provide unit process cost (<http://www.meniscus.co.uk>). This information can then be relayed to plant operators and managers for appropriate action. Activated sludge treatment is basically the same regardless of the type of plant discussed, which is probably why many of the remedies for bulking and foaming in conventional plant now apply to new plant with suspended biomass. Similar pressures are exerted upon the biomass irrespective of the type of process involved and the same filamentous microbes that cause separation problems in conventional plant are present also in the new treatment processes.

Variables include sewage (such as flow, load and substrate type), consumables (power and oxygen requirement), while outputs include waste solids settlement (and disposal) and effluent quality. Broadly speaking it is a case of balancing inputs with outputs and, although the configuration might change the values to be measured remain the same. Fig. 32.11 compares the relative costs of these components in a modern sewage treatment process.

Benchmarking has already indicated that the typical aeration cost for an activated sludge plant is £0.009 per m³ treated (based on an

electricity price of £0.035 per kWh), which is a valuable starting point in evaluating the cost performance of a works. One would expect an optimum unit treatment cost for any given process and that an excessively high or low unit cost would reflect an imbalance in the inputs or outputs. This will help identify conditions that may give rise to sludge bulking or foaming.

By combining expert systems, advanced data management and summary data from telemetry, it is now feasible to identify known operating problems at a site and to suggest appropriate remedial action within minutes of inputting the original raw data.

With advances in internet and mobile phone technology the day is not far off when operators will enter site readings into their mobile phone and, using the Internet, view process and cost exceptions, operating suggestions, trends and telemetry data on their phone while they are still on site.

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Odour generation and control

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1 INTRODUCTION

Fresh sewage has a characteristic musty smell resulting from its components; a mixture of discharges from toilets, baths, sinks, dish-washers and washing machines, and industrial wastes. The mixture of malodours typically contains compounds utilized in cleaning agents used in the home (such as limonene), solvents (such as tetrachloroethylene), petrol derivatives (such as benzene) as well as the odours associated with human waste, such as urea and ammonia from urine and skatole and indole (breakdown products of tryptophan) from faeces.

Although the smell of fresh sewage is not pleasant, the odorous compounds given above rarely give rise to odour problems. It is the strong objectionable malodours, in particular hydrogen sulphide, that develop when sewage and sewage sludges are stored, which give rise to odour problems. These strong objectionable malodours are nearly always associated with the development of anaerobic conditions and the activity of anaerobic bacteria as soon as nitrate (and nitrite) and dissolved oxygen are exhausted. This anaerobic condition of sewage or sludge is often termed septicity. The chemicals commonly responsible for sewage odour are given in Table 33.1. All the compounds given in Table 33.1 are produced by the action of bacteria on components of wastewater. Table 33.1 also gives the concentrations in the air at which they can be detected by the human nose (the threshold odour concentration).

Septicity can occur whenever sewage or sludge is retained for more than a few hours

without replenishment of dissolved oxygen and in the absence of nitrate (or nitrite) salts. Typically sewage becomes septic in pumped (rising-main) sewers, primary settlement tanks, storm storage tanks and flow-balancing tanks. Primary sludges and mixed primary and biological sludges become septic in primary settlement tanks, storage tanks, and during anaerobic digestion. Sludges containing primary sludge are likely to become septic very rapidly because the numbers of microorganisms and the available substrates per unit volume are greater than in sewage.

The consequences of sewage or sludge becoming septic are important for a number of reasons, in addition to the potential for odour nuisance:

1. Hydrogen sulphide in the atmosphere is an extremely toxic gas at concentrations above 300 ml/m³ (ppm) and has been the cause of deaths in enclosed areas. It is a particularly dangerous gas as, although it can be smelt at very low levels (0.5 µl/m³ (ppb)), it cannot be smelt at concentrations above 150–250 ppm, and at the concentrations at which it is toxic (World Health Organization, 1987).
2. Hydrogen sulphide that has been released into the atmosphere can cause corrosion of metals (such as copper (and many of its alloys), silver as well as iron or steel) and is a significant cause of corrosion of concrete, as it may be biochemically oxidized on moist surfaces to sulphuric acid by various autotrophic *Thiobacillus* spp. such as *Thiobacillus concretivorus* (Parker, 1947).

TABLE 33.1 Threshold odour concentrations of most common sewage odour compounds (Woodfield and Hall, 1994; Bonnin *et al.*, 1990)

Class	Compound	Chemical formula	Description	Threshold odour concentration ^a ($\mu\text{l}/\text{m}^3$)
Sulphides	Hydrogen sulphide	H_2S	Rotten egg	0.5
	Methylmecaptan (methanethiol)	CH_3SH	Cabbage, garlic	0.0014–18
	Ethylmercaptan (ethanethiol)	$\text{C}_2\text{H}_5\text{SH}$	Rotten cabbage	0.02
	(Di)methylsulphide	$(\text{CH}_3)_2\text{S}$	Rotten vegetable	0.12–0.4
	(Di)methyl disulphide	$(\text{CH}_3)_2\text{S}_2$	Putrefaction	0.3–11
Nitrogen containing	Ammonia	NH_3	Very biting, irritating	130–15300
	Methylamine	CH_3NH_2	Rotten fish	0.9–53
	Dimethylamine	$(\text{CH}_3)_2\text{NH}$	Fish	23–80
	Indole	$\text{C}_8\text{H}_6\text{-NH}$	Faecal, nauseating	1.4
	Skatole (3-methyl indole)	$\text{C}_9\text{H}_8\text{-NH}$	Faecal, nauseating	0.002–0.06
Acid	Acetic	$\text{CH}_3\text{-COOH}$	Vinegar	16
	Butyric (butanoic)	$\text{C}_3\text{H}_7\text{-COOH}$	Rancid	0.09–20
	Valeric	$\text{C}_4\text{H}_9\text{-COOH}$	Sweat, transpiration	1.8–2630
Aldehydes and Ketones	Formaldehyde	H-CHO	Acrid, suffocating	370
	Butyraldehyde	$\text{C}_3\text{H}_7\text{-CHO}$	Rancid	4.6

^a The threshold odour concentration of a specific compound is the concentration at which it can only just be detected by a panel of at least 8 people. A similar technique, olfactometry, can be used to measure the strength of an unknown odorous gas containing a mixture of compounds. In this case the odour strength (in units of ou/ m^3) is defined as the number of dilutions of an odorous gas required before it can be detected by only 50% of a panel of at least 8 people (CEN (1999) European Committee for Standardization). A range of values is given by Woodfield where quality of data is uncertain.

However great the concentration of odorous compounds that develop in sewage or sludge, they have impact only when they are transferred into the atmosphere, particularly beyond the boundary of the works and to the vicinity of a potential complainant. Therefore, in addition to the factors affecting the production of malodours, the factors leading to transport to locations where they may cause nuisance, and the factors affecting their release also need to be considered when dealing with a potential odour problem. Hydrogen sulphide and organic sulphides that remain dissolved and are subsequently oxidized chemically or biochemically while in solution will not create odour problems.

An important factor in the amount released is the chemical state of the odorous compound within the liquid phase, which is affected by the pH value of the sewage or sludge. Factors affecting the pH value of the sewage or sludge are, therefore, also important in the consideration of the potential for odour nuisance. Low pH values favour the emission of H_2S , mercaptans and volatile fatty acids, while high pH

values favour the emission of ammonia and amines. Although odours at high pH values can be unpleasant, they are generally less intense, and increasing the pH value of sludges by chemical addition has been used as an odour control technique.

The solubility of H_2S in water varies with temperature from 4 ml/ml at 4°C to 2.55 ml/ml at 20°C , although the amount dissolved will also be directly proportional to the total pressure and to the partial pressure of H_2S in the gaseous phase, according to Henry's law. H_2S is a weak dibasic acid which dissociates as follows:



The proportions of H_2S , HS^- and S^{2-} are significantly affected by pH value and will also be slightly affected by temperature and ionic strength (EPA Process Design Manual for Sulphide Control and Sanitary Sewerage Systems, 1974). The proportion of dissolved sulphide, in an associated (non-ionic) form as H_2S , can vary from 75% at pH 6.5 to 50% at pH 7 and only 10% at pH 8. Between pH 9

and 10, the dissolved sulphide would be almost entirely present as HS^- ions. Some of the sulphide may also be present as insoluble metal sulphides, having combined with metals in the sewage. In the UK, the pH value of sewage is normally within the range 6.5 to 8 (depending on the alkalinity of the water) and the formation of volatile fatty acids that will occur under anaerobic conditions can cause the pH value to decrease below 6.

2 DEVELOPMENT OF MALODOURS

Sewage and sludge malodours are predominantly the result of the action of anaerobic and facultative anaerobic bacteria. These bacteria are derived from the original wastewater. Anaerobic bacteria, which grow more slowly and have a lower cell yield than their aerobic counterparts, are selectively encouraged to grow by the conditions in the various stages and processes. The steps of increasing levels of anaerobicity leading to the formation of malodours include:

1. Depletion of dissolved oxygen and nitrate by microorganisms present in sewage and slimes
2. Microbial reduction of organic sulphides and nitrogen compounds by anaerobic and facultative anaerobic proteolytic bacteria under anaerobic conditions, giving rise to dimethyl sulphide, ammonia, amines, mercaptans and other malodorous compounds
3. Production of volatile fatty acids (e.g. butyric, propionic, lactic, acetic) by microbial lipolysis by anaerobic and facultative anaerobic bacteria, also leading to a reduction in pH value and increased release of any sulphur containing malodours
4. Microbial respiration of sulphate by strict anaerobes (the sulphate-reducing bacteria) giving rise to H_2S . This is the most important of the odour-generating processes
5. Microbial breakdown of volatile fatty acids giving rise to methane (anaerobic digestion) in digester gas. In the digester, methanogenesis occurs concurrently with hydrolysis,

acidogenesis, proteolysis and sulphate reduction and biogas will contain hydrogen sulphide and other malodorous sulphur compounds. However, the residual odour of well-digested sludge, after cooling, is less than that of primary sludge.

These steps are accompanied by a decrease in the redox potential of the sewage, as illustrated in Fig. 33.1 (Boon, 1995). Steps 2, 3, 4 and 5 are of particular importance with respect to sludge odours. Step 4 is of particular importance with respect to sewage malodours.

2.1 Depletion of dissolved oxygen and nitrate

Electron receptors for microbial respiration are (in order of decreasing preference) oxygen, nitrate, sulphate and carbonate. Their utilization under increasingly anaerobic conditions produces respectively water, nitrogen, hydrogen sulphide and methane (Fig. 33.1). Anaerobicity, or 'septicity', is considered to occur in sewage or sludge when the microorganisms that are present have utilized all the dissolved oxygen (DO) and any nitrates (or nitrites) that may be present.

The rate at which the DO will be used by the microorganisms (aerobic respiration) depends on the 'age' and temperature of the sewage and the area of the submerged surfaces. The 'age' of sewage affects the respiration rate of the sewage because the numbers of microorganisms will increase while sewage remains in a sewer under aerobic conditions. Under anaerobic conditions, complex soluble, suspended and colloidal organic compounds, such as carbohydrates and protein will be broken down into readily biodegradable and soluble volatile fatty and amino acids. The respiration rate will eventually reach a maximum as readily biodegradable substrates become limiting and it will then start to decline. In practice, the rate of uptake of DO by aerobic sewage has been found to vary from 2 to 3 mg/l/h for 'young' domestic sewage and up to 14 mg/l/h at 15°C, as the sewage 'ages' during its passage through the sewerage system

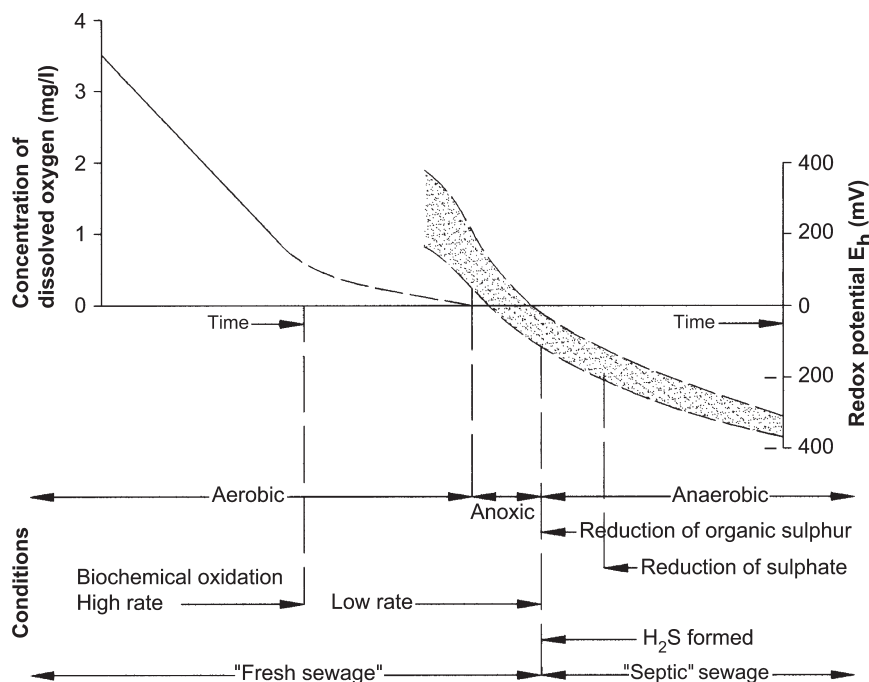


Fig. 33.1 Variations in condition of sewage in relation to concentration of dissolved oxygen and redox potential.

(Boon, 1995). A typical average rate in the UK is about 6 mg/l/h at 15°C. Bacterial slimes that adhere to submerged surfaces in sewers are generally not limited by available substrate; they have been found to consume DO at a rate of about 700 mg/m²/h at 15°C (Boon and Lister, 1975). The consequence of respiration in both the sewage and slimes occurring simultaneously is that the sewage will rapidly become devoid of DO, particularly in a rising-main sewer of small diameter as the surface area per unit volume will be large compared with a sewer of large diameter.

In gravity sewers, the slope of the sewer is designed so that grit and other debris do not accumulate, and under average flow-rate conditions, the velocity of sewage generated by the slope will normally create sufficient turbulence to ensure that the rate at which oxygen dissolves from the atmosphere in the sewer exceeds the respiration rate of the microorganisms growing in the sewage and on the submerged surfaces. Under such conditions, septicity will not occur, although volatile and odorous compounds already present in sewage will be stripped from solution and into the atmosphere. However, such compounds are

unlikely to cause a nuisance, unless the sewage had become septic upstream of the turbulent conditions.

If aerobic sewage (containing DO) enters a rising-main sewer or other process stage, where there is no opportunity for the sewage to be re-aerated, the concentration of dissolved oxygen will decrease due to the respiration of sewage and slimes. The rate that microorganisms consume DO will remain fairly constant (see Fig. 33.1) until the concentration reaches about 0.2 to 0.4 mg/l. At about 0.2 mg/l, the rate will decline and asymptote to zero at zero DO. However, as the DO becomes rate limiting, any oxidized nitrogen present in the sewage will provide an alternative electron acceptor for the dissimilation of organic matter (in the same process as denitrification in the anoxic zone of an activate-sludge process). Under such conditions, the microorganisms will continue to 'respire' and oxidize substrate but at a slower rate of about 40% of the aerobic rate. Under these 'anoxic' conditions, the redox potential will decline from about +50 mV to about -100 mV (E_h). Nitrate levels are rarely high in domestic sewage, but addition of nitrate salts (normally of calcium or magnesium) has been

used as an effective means of septicity control at a large number of sites. When all DO and nitrate is utilized the sewage then becomes septic. Time taken for sewage to cease being aerobic and to become anaerobic will vary according to the concentrations of microorganisms and the substrates present; it is likely to be in the range from 30 minutes to 2–3 hours (Boon and Lister, 1975).

Septicity is most likely to occur during warm conditions, when an increase in temperature of the sewage leads to increased respiration and growth of the microorganisms and rapid depletion of a reduced concentration of dissolved oxygen. As the activity of anaerobic bacteria also increases with temperature, odour problems are most frequently associated with warm summer months.

The causes of septicity developing in primary sludges are similar to those given above for sewage in sewers and storage tanks, although the effect of microorganisms growing on submerged surfaces is likely to be less than for sewage in pipes. In primary sludges, depletion of dissolved oxygen and nitrate occurs very rapidly because the numbers of microorganisms in the sludge are several orders of magnitude higher than in sewage and the availability of substrate per unit volume is also much greater. It is unlikely that primary sludge will contain dissolved oxygen or nitrate.

Surplus activated or humus sludges, derived from aerobic biological treatment processes, may contain dissolved oxygen and nitrate (particularly if the aerobic biological processes achieved complete nitrification). Such sludges become septic less rapidly than primary sludges, unless co-settled with primary sludge. The depletion of nitrate, by biological sludges in settlement tanks, can be a cause of problems of rising sludge, due to the formation of micro-bubbles of evolved nitrogen derived from nitrate. Humus sludge, from a nitrifying biological filter, is unlikely to contain nitrate as the retention period in the humus settling tank is likely to be long and any nitrate in the filter effluent will have been 'respired' by heterotrophic bacteria in the sludge.

2.2 Microbial reduction of organic sulphides and nitrogen compounds under anaerobic conditions

In the absence of DO, proteolytic bacteria (which may be facultative anaerobes or anaerobes) break down proteins in the sewage into amino acids, and subsequently degrade the sulphur-containing amino acids, cysteine, cystine and methionine to produce thioles (mercaptans), and other organic sulphides and disulphides. These compounds smell very obnoxious when released to the atmosphere and have very low odour thresholds (see Table 33.1). The proteolytic bacteria are active at a higher redox potential than those which subsequently reduce sulphate to form H_2S .

Domestic sewage normally contains about 3–6 mg/l of organic sulphur, which is present mainly as proteinaceous matter, and additional organic sulphur in the form of sulphonates (about 4 mg/l), derived from household detergents. The ability to produce sulphide from protein is quite common among bacteria, for instance *Proteus* spp., *Bacteroides* spp. and some *Clostridium* spp., all of which can grow anaerobically (Crowther and Harkness, 1975).

Volatile sulphides produced in this way are a small proportion of the malodours associated with sewage but can be very significant in the odours from sludges where the concentration of the bacteria and substrates are higher. Recent measurements on-site have shown that concentrations of dimethyl sulphide and/or thiols in air and/or biogas in contact with sludge may be occasionally greater than the concentration of hydrogen sulphide in the air and/or biogas.

2.3 Production of volatile acids

In addition to the action of proteolytic bacteria under anaerobic conditions in the breakdown of protein, there are also the bacterial actions of hydrolysis, fermentation and lipolysis on carbohydrates, fats and oils leading to the formation of short-chain volatile fatty acids.

Under aerobic conditions in sewers and primary settlement tanks, the concentration of the volatile fatty acids (VFAs) developing in sewage is relatively low, and is rarely

the main cause of odour nuisance. Under anaerobic conditions, the formation of VFAs may lead to a lowering of the pH value, resulting in the enhancement of release of any hydrogen sulphide formed under such conditions. Significant concentrations of VFAs may develop in process stages which deliberately introduce anaerobic conditions, e.g. the use of a primary sludge fermentation stage to increase the concentration of readily biodegradable BOD to enhance biological phosphorus removal (see Chapter 23), or the use of anaerobic treatment processes for the partial treatment of high BOD wastewaters or sludges.

In sludges, the reduction of carbohydrates (including polysaccharides), fats, oils and greases is the 'acid forming' stage of the anaerobic digestion process as described in Chapter 24. When the process is contained within a heated anaerobic digester, the volatile acids are rapidly converted to methane. However, in primary sludges the presence of volatile fatty acids contributes to the characteristic and obnoxious smell of septicity. The reduction in volatile acids during the anaerobic digestion process can be accompanied by a reduction in the odour level of the sludge of about 90% (Hobson, 1995).

The concentrations of volatile fatty acids that can develop in sludges are much higher than in sewage and the decrease in pH value is more marked with values of about 5.5 having been observed. At such low pH value, hydrogen sulphide and other malodorous sulphur-containing compounds, which will also be present in the sludge, are almost completely associated (non-ionic) and available for release into the atmosphere.

2.4 Reduction of sulphate

Hydrogen sulphide, produced from sewage or sludge by the reduction of sulphates by anaerobic bacteria, is the most significant cause of odour problems related to sewage and sludge treatment. Even when hydrogen sulphide is not the main cause of malodour, it is nearly always present where there are sewage or sludge odours. For this reason,

hydrogen sulphide is often used as a 'marker' for the detection of sewage-related malodours. Factors that affect the generation and control of hydrogen sulphide in sewers have been studied by numerous research workers (Thistlethwayte, 1972; Boon, 1995). In addition, the use of sulphate-reducing bacteria in the treatment of strong wastes has also been the subject of numerous papers, as discussed in Chapter 28.

The bacteria responsible for sulphide formation are in a class which 'respires' sulphate to provide energy for the dissimilation of organic matter. This is dissimilatory sulphate reduction, as opposed to assimilatory reduction of inorganic sulphate which occurs naturally in an analogous process to nitrogen fixation. The process of dissimilatory sulphate reduction is analogous to 'nitrate respiration', associated with denitrification in an anoxic environment. A very small amount of reduced sulphur is assimilated by the bacteria, but most is released into solution as sulphide ion, usually substantially as molecular (non-ionic) H_2S . These bacteria are termed 'sulphate-reducing' (Postgate, 1959, 1984).

The sulphate-reducing bacteria are strict anaerobes and cannot metabolize while dissolved oxygen or nitrate (or nitrite) remain in the sewage. However, they can survive adverse conditions of temperature, aerobicity, salinity and pressure, and are widely distributed in the environment, ready to become active whenever local conditions become anaerobic, such as within sewage or sludges and also in bacterial slimes on the submerged surfaces of sewers and holding-tanks and in the undisturbed sediments of sewers, rivers and estuaries. They are heterotrophic bacteria and grow slowly at a rate which is about 7% of the growth rate of aerobic microorganisms. The relative rates of growth can be calculated from the rate of sulphide formation in a rising-main sewer in comparison with the rate of demand for dissolved oxygen in the same sewer that would be required to prevent septicity (Boon, 1995).

A number of genera and species of sulphate-reducing bacteria have been isolated, as listed in Table 33.2 (after Postgate, 1984).

TABLE 33.2 Sulphate-reducing bacteria (after Postgate, 1984)

Genus	Species	Strains
<i>Desulfomaculum</i>	<i>nigrificans</i>	7
	<i>orientis</i>	2
	<i>ruminis</i>	2
	<i>acetoxidans</i>	1
<i>Desulfovibrio</i>	<i>africanus</i>	3
	<i>desulfuricans</i> subsp. <i>aestuarii</i>	1
	<i>desulfuricans</i> subsp. <i>desulfuricans</i>	18
	<i>gigas</i>	2
	<i>salexigens</i>	7
	<i>vulgaris</i> subsp. <i>oxamicus</i>	1
	<i>vulgaris</i> subsp. <i>vulgaris</i>	13
	<i>Desulfovibrio</i> spp.	7
	<i>baarsii</i>	1
	<i>sapovorans</i>	1
	<i>postgatei</i>	1
<i>Desulfobacter</i>	<i>propionicus</i>	1
<i>Desulfobulbus</i>	<i>multivorans</i>	1
<i>Desulfococcus</i>	<i>limicola</i>	1
<i>Desulfonema</i>	<i>magnum</i>	1
<i>Desulfosarcina</i>	<i>variabilis</i>	1

The best studied is the non-spore forming genus *Desulfovibrio*, of which the main type associated with the production of sulphide from sewage and sludge is *Desulfovibrio desulfuricans* (Crowther and Harkness, 1975). *Desulfovibrio* species are Gram-negative, polarly-flagellated, curved rods (Stanier *et al.*, 1976).

The number of sulphate-reducing bacteria present in sewage or sludges is dependent on the length of time that the sewage or sludge has been retained under anaerobic conditions. Heukelekian (1948) found typical numbers of sulphate-reducing bacteria of 25 000/ml in sludges and 60 to 600/ml in crude sewage, with numbers increasing, under anaerobic conditions, from 100/ml to nearly 100 000/ml over 14 days. Toerian *et al.* (1968) found counts of 27 500 to 53 000/ml in raw sewage sludge. Numbers increased when sulphate rich water was added to the sludge. Schmitt and Seyfried (1992) found concentrations in sewer sediment samples of 2.4×10^7 – 11.4×10^7 per g volatile solids (volatile solid content of the sediment was less than 5% in a cleaned sewer, and between 3 and 27% in two sewers that were not cleaned).

The biochemistry of sulphate-reducing bacteria has been reviewed by Postgate (1984).

Ammonia is the principal source of nitrogen for the bacteria and pre-formed organic matter (acetate in particular) is required for growth of *Desulfovibrio* species. Postgate also indicated an exceptionally high requirement for inorganic iron. The sulphide ion generated by sulphate reduction will also precipitate iron and other heavy metals as insoluble sulphides. This accounts for the darker colour of septic sewage and sludges.

The reduction of sulphate to sulphide by *Desulfovibrio* involves the transfer of four pairs of electrons (Stanier *et al.*, 1976). The process, described in Postgate's review (1984), is a four step process, with sulphite, trithionate and thiosulphate as intermediates. The reduction to sulphide is mediated by five enzymes and the catalyst desulfovibrin with a key intermediate of adenyolphosphosulphate (APS). The process is inhibited by structural analogues of sulphate, including selenate (SeO_4^{2-}), and monofluorophosphate (FPO_3^{2-}). Chromate and molybdate ions are also inhibitors, probably acting by forming an unstable analogue of APS.

The sulphate-reducing bacteria require a redox potential of about -200 mV for the initiation of growth. Sulphate reduction

occurs at a redox potential within the range -200 to -300 mV (E_h), depending on the pH value of the sewage being within the range 6.5–8, respectively. The growth of sulphate-reducing bacteria is accompanied by a further drop in redox potential (Postgate, 1984).

The concentration of sulphate in sewage and its resultant sludges can vary greatly from area to area, depending, for example, on the hardness of the ground water (which may infiltrate the sewerage system) or the potable water supply, the method of treatment of potable water and on the input of industrial wastewaters. In coastal areas, where there may be infiltration of the sewerage system with seawater, and in certain volcanic parts of the world, concentrations may be very much higher and extreme problems with malodours, corrosion and inhibition of anaerobic digestion may be experienced. Sulphate concentrations in sewage in inland areas of the UK are generally over 10 mgS/l, with typically about 20 mgS/l present.

In wastewater, the concentration of sulphate (at about 20 mgS/l) rarely limits the sulphide production and, in the range 10–60 mgS/l, does not affect the rate of sulphide formation (Boon, 1995). The amount of sulphide formed in a rising-main sewer has been found to be directly proportional to the retention time of the sewage in the main. The longer the time, the greater the amount formed. As the pipe remains full of sewage, the amount formed daily is directly related to the volume of the sewage in the pipe, the internal surface area of the pipe, the chemical oxygen demand (COD) and temperature of the sewage and the availability of inorganic and organic sulphur-containing compounds (Boon, 1995; Hvitved-Jacobson *et al.*, 1988; Nielson *et al.*, 1998). Changing the rate or frequency of pumping will have little or no effect on the total amount formed daily, although flow variations and changes in COD will account for variations in concentration of sulphide during the day. Increases in temperature will increase the rate of formation; the rate is likely to double for a 10°C increase in temperature within the range of about 5 – 25°C . At temperatures above 25°C , increases in temperature will

start to have an adverse effect on growth rate of microorganisms, with the rate declining to zero as the pasteurization temperature of 45°C is reached.

A number of empirical relations have been published to estimate the production of sulphide in rising-main sewers and gravity sewers, and to derive the conditions necessary to prevent sulphide formation. These have recently been reviewed (Boon, 1995). Equations based on these relations, together with others which are traditionally used for the hydraulic design of a sewerage system, have been linked together to form a mathematical model (Boon *et al.*, 1998).

In sediments and thick slimes, sulphate concentration may become limiting across the depth (Schmitt and Seyfried, 1992). The rate of sulphate reduction in sediments was found in experiments on sewer sediments in Hannover (Schmitt and Seyfried, 1992) to be up to $0.64\text{ g S/m}^2/\text{h}$. Maximum rates of sulphate reduction were found at sediment depths of between 5 and 7.5 cm.

In sludges, much higher concentrations of dissolved H_2S may develop compared with sewage, particularly where there are high sulphate concentrations. The rate at which septicity develops depends on the solids content and temperature of the sludge and the availability of sulphur-containing organic and inorganic compounds. Total sulphur in sludge can account for between 1 and 2% of the dry matter (Sommers *et al.*, 1977), although the form of the sulphur as inorganic and organic compounds is variable. Storage of primary sludge for more than 1–2 hours will generally result in hydrogen sulphide being produced by sulphate reduction. In addition, other malodorous compounds will be formed by the action of the proteolytic and acid-forming bacteria as described above, producing compounds such as methyl sulphide, dimethyl sulphide, methyl disulphide, thiols and volatile organic fatty acids, such as acetic, propionic and butyric. Lowering of the pH value increases the potential for release of volatile sulphides to the atmosphere.

In the anaerobic digestion process, the sulphate-reducing bacteria operate in parallel,

and in competition, with methanogenic bacteria. Both sulphate-reducing bacteria and methanogenic bacteria are strict anaerobes. The anaerobic digestion process breaks down a wide range of odorous compounds with a reduction in the volatile fatty acid content and an increase in pH value to approximately neutral. However, sulphate reduction during the anaerobic digestion process will lead to H_2S being released in the biogas and being present in the digested sludge. In addition, other malodorous sulphides, formed by the action of proteolytic bacteria, will be present in both digesting and digested sludge.

The release of malodours during the digestion process is controlled because the process of anaerobic digestion is carried out within totally enclosed vessels. Biogas produced by the process is combusted in a boiler, combined heat and power (CHP) plant or flare-stack, with the consequent oxidation of odorous compounds. H_2S concentrations in biogas can be within the range 100 to 3000 ppm (v/v). Severe odour nuisance can occur if biogas is released directly into the atmosphere, through pressure-relief valves or if there are problems with the gas-handling or storage systems. Malodours may also be released when the digested sludge is transferred to a secondary digester, if care is not taken to avoid turbulence of the sludge after digestion. Hydrogen sulphide in the biogas can cause corrosion problems with CHP or boiler equipment.

In addition to exacerbating odour and corrosion problems, high levels of sulphate in digesting sludge and the resultant development of sulphate-reducing bacteria, have been found to have an adverse effect on the methane-producing bacteria. The apparent incompatibility of sulphate-reducing and methane-producing bacteria (Postgate, 1984) is thought to be partly due to the following:

- the toxic effects of sulphide produced (concentrations of soluble sulphide are toxic to methanogenic bacteria in concentrations above 200 mg/l (Lawrence *et al.*, 1964)
- competition by some species of sulphate-reducing bacteria for acetate (e.g. *Desulfotomaculum acetoxidans* and *Desulfobacter postgateii*).

However, *Desulfovibrio* species do not oxidize acetate, and the end-products of carbon oxidation, coupled to sulphate reduction by most *Desulfovibrio* species, are acetate, water and carbon dioxide

- competition by the sulphate-reducing bacteria for molecular hydrogen (Pfennig and Widdel, 1982; Postgate, 1984).

Various methods of control of sulphate-reducing bacteria during digestion have been tried, including the use of specific inhibitors of the sulphate-reducing bacteria and the addition of iron salts to precipitate sulphide as it is formed.

2.5 Summary of common causes of odour nuisance

The main causes of odour problems associated with different stages of wastewater and wastewater sludge handling and treatment are summarized in Table 33.3.

3 EMISSION OF MALODOURS AND THEIR CONTROL

Control of malodours is achieved by a combination of measures ranging from prevention to reduction of septicity, together with minimization of the release and potential impact of malodours. In most circumstances, total prevention of septicity at all times and under all conditions would not be practical or economical.

3.1 Prevention of septicity

Prevention, or a reduction in malodour formation may be achieved by the following:

- Maintaining conditions that prevent the development or action of anaerobic bacteria. This is most effectively achieved by maintaining aerobic or anoxic conditions and by minimizing retention in systems where there is no aeration, such as in holding tanks or rising-main sewers.

TABLE 33.3 Summary of the main causes of odour production at different process stages

<i>Process stage</i>	<i>Main odorous compounds</i>	<i>Bacteria responsible</i>	<i>Possible locations of odour release</i>
SEWAGE			
Rising main sewer	H ₂ S	Sulphate-reducing bacteria in sewage and in slimes on sewer walls	At and downstream of the discharge point
Sediments	H ₂ S	Sulphate-reducing bacteria in sediments	From sewage when sediments are disturbed
Primary, storm storage or balancing tanks	H ₂ S	Sulphate-reducing bacteria in sewage and sludge	Tank weirs and outlet channels
Biological filters	H ₂ S	Sulphate-reducing bacteria in slimes if excessive slime growth (particularly high rate filters)	Ventilation air from the biofilter
SLUDGES AND IMPORTED WASTES			
Primary sedimentation and primary sludge storage tanks	H ₂ S	Sulphate-reducing bacteria	De-sludging wells, discharges to storage tanks, dewatering area, return liquor sumps and discharge points, tanker collection/delivery points
	Organic sulphides	Proteolytic bacteria in sludge	
	Volatile fatty acids which also lower pH value and increase potential release of sulphides	Lipolytic bacteria in sludges	
Anaerobic digesters	Ammonia in the form of ammonium salts	Formed by the hydrolysis of urea and proteolysis and released if lime dosing employed	Downstream of alkaline additions
	H ₂ S	Sulphate-reducing bacteria	Discharge wells, sludge-digestion tanks, pressure relief valves for biogas
	Organic sulphides	Proteolytic bacteria	
	Ammonia	Proteolytic bacteria in well-digested sludge	Ammonia from discharge of liquors after sludge pressing

Aerobic conditions can be maintained in some cases by the addition of air, oxygen, or hydrogen peroxide or anoxic conditions can be achieved by the addition of nitrate salts

- Reduction in the input of sulphate, or modifications to the treatment processes, in order to reduce the formation of sulphide in the treatment and handling of sludges, particularly if there were saline intrusion in the sewerage system or an industrial discharge with a high sulphate content. A reduction in sulphate concentration is unlikely to have a significant impact on sulphide formation in sewage or the malodours associated with septic sewage
- Addition of bactericides (such as chlorine or hypochlorite) has been used to reduce the growth of sewer wall slimes and to inhibit the respiration rate of bacteria in sewage. These have had variable success and may have potentially harmful effects on subsequent biological treatment stages or the quality of effluent discharge to receiving waters
- Addition of specific inhibitors of sulphate-reducing bacteria have also been used (particularly for anaerobic digestion stages), and proprietary products are available, although their use is not widespread in the water industry. However, these tend to be successful in the short term only, as species may be selected which are resistant to the inhibitor.

3.2 Treatment of septicity

Treatment of sewage or sludges that have already become septic can be achieved either by bacterial oxidation of the odorous compounds or by addition of chemicals to oxidize or precipitate sulphides, or to increase the pH value so that they are not emitted to the atmosphere.

- Hydrogen sulphide and volatile organic sulphides in sewage or sludge will be oxidized when conditions become aerobic (such as in a gravity sewer following a rising-main sewer, or in an aerated activated-sludge plant following a primary sedimentation tank) and this may be used as a control method, if release of sulphide prior to the aerobic step can be avoided, e.g. by reducing turbulence or provision of covers for treatment stages upstream of the aerobic stage.
- Hydrogen sulphide may also be controlled by the addition of precipitants, such as iron salts, or oxidizing agents, such as peroxide.

3.3 Prevention of release of malodorous compounds

The amount of hydrogen sulphide and other malodorous compounds released into the atmosphere will depend on the interfacial area between liquid and air and the solubility of the odorous gases, which in turn is affected by temperature, total pressure and partial pressure. Under static enclosed conditions, e.g. under the covers of enclosed storage tanks, gases will be released from solution at a slow rate to maintain chemical equilibrium with the gas phase according to Henry's law. However, significant release of odour is likely to occur where the interfacial area is large, such as settling tanks or sludge lagoons and at points of turbulence, where the flow breaks up into droplets or air is entrained into the liquid (e.g. at the discharge end of a rising-main sewer, at discharge weirs of primary sedimentation tanks or during sludge handling). Malodours may also be released during disposal of sludges to agricultural land.

Odour release can be minimized, for example, by avoiding cascades and turbulence, retaining sludges in pipes (rather than in open channels) and covering distribution chambers, channels and tanks.

In the case of ionic species of volatile organic chemicals, the pH value will also have an important impact on the amount and type of odorous compounds available to be released. Many of the odorous compounds in sewage (including sulphides and some nitrogenous compounds) are present in sewage in unionized (dissolved gas) and ionized forms (see above). Only the unionized form is available for transfer to the atmosphere and release can be prevented by maintaining the compounds in the ionized form. Neutral or acidic conditions (pH 7 or below) favour the release of 'acidic' malodours including organic sulphides and hydrogen sulphide and volatile, organic, fatty acids, such as acetic, propionic and butyric (see Table 33.1). Under alkaline conditions there will be an increase in the release of ammonia and amines, together with other volatile organic compounds such as skatole and indole. These 'alkaline' odours have much higher odour threshold concentrations than acidic sulphides but are more persistent.

Measures to control septicity by addition of alkali should avoid increasing the pH value above about 8.5 or the problem will not be resolved, as malodorous alkaline odours would prevail.

3.4 Treatment of malodorous gases

Where covers are provided for tanks or other odour-emitting stages, odorous gases may need to be vented to an odour treatment unit. A wide range of odour treatment processes are available, including chemical and biochemical scrubbing and absorption processes. One form of control for treated gases employs biological processes in the form of a biofilter or a bioscrubber, in which sulphide-oxidizing bacteria (*Thiobacilli*) reoxidize sulphide, mercaptans, and other malodorous compounds. The *Thiobacilli* are chemoautotrophic and utilize the oxidation of reduced sulphur

compounds to support growth (Sublette, 1992). The end product is sulphate. In this process the autotrophic bacteria, as well as heterotrophic bacteria, are allowed to develop on the surface of a media (e.g. a natural medium such as peat, shells or coir) or in the recirculation liquor and on the exposed surfaces of a plastic medium bioscrubber. In both cases, sufficient moisture must be provided to maintain the biological film and ensure that the sulphide and other soluble odorous compounds transfer from the gas phase. Control is required to ensure that the biofilter does not become too acidic. Recently, two-stage bioscrubbers have been developed and successfully used (Joyce and Leach, 1998) to remove carbon- (and nitrogen-) based odours in the first stage (by the growth of heterotrophic bacteria) and to remove sulphides in the second stage (by maintaining the growth of the slowly-growing autotrophic bacteria). In a development of these methods, the nitrifying activated-sludge process has been used to treat odorous air, by ducting malodorous air to the intake of the blowers that are normally provided for aerating the process (Stillwell *et al.*, 1994; Ostojic and O'Brien, 1994).

An effective and low-cost catalytic hydrated iron oxide chemical filter has been recently developed (Boon and Boon, 1998), which has been successfully used to remove up to 95% of H_2S from air, together with lower percentages (about 30 to 80%) of organic sulphides. The design of this filter is based on the fact that H_2S will react chemically with hydrated iron oxide to form water and sulphur. The conditions that this reaction requires have been investigated and an application for a patent has been submitted to provide commercial benefit for the future design and operation of such filters.

3.5 Summary of control methods

Odour problems can be minimized by:

1. minimizing retention under anaerobic conditions, including avoiding excessive accumulation of debris and grit in pipes, channels and tanks

2. avoiding unnecessary contact of sewage and sludge with the atmosphere and minimizing turbulence
3. retaining malodours dissolved in solution until they can be biochemically oxidized naturally, e.g. in an aerobic stretch of sewer, or in an aerated activated-sludge plant
4. covering units and venting of contained air to an odour treatment unit
5. addition of chemicals such as:
 - oxygen or nitrate to maintain aerobic or anoxic conditions
 - oxidant chemicals, such as hypochlorite, hydrogen peroxide or potassium permanganate to reduce microbial activity and oxidize previously formed sulphides
 - iron salts to precipitate sulphides. Iron salts are also used as a catalyst for the chemical and biochemical oxidation of sulphide
 - alkali to convert H_2S to HS^- and S^{2-} , the pH value should not exceed 8.5, to avoid release of alkaline odours (e.g. ammonia, amines, skatole and indole)
 - specific inhibitors for sulphate reduction, these may be effective in the short term only, as adaptation of microbial species may occur.

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Recalcitrant organic compounds

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1 INTRODUCTION

The question of recalcitrant organic compounds first surfaced as a real public and scientific issue with the publication of *Silent Spring* (Carson, 1962). This book raised awareness of the actual and potential problems caused by the accumulation in the environment of organic compounds which were not readily destroyed by biological activity and so built up to concentrations which were toxic to man or wildlife or had some other unacceptable effects. Perhaps the most infamous of these compounds are the polychlorinated insecticides such as DDT, aldrin and dieldrin, which are subject to bioaccumulation in the food chain and which caused devastating effects on the populations of predatory birds in the 1950s and 1960s. A more visible example was the problem of water pollution by so-called 'hard' detergents during the same period. The realization of the consequences of the release of these chemicals in the environment led to much research, which is still ongoing, into the fate of these materials in the wider environment.

In the 1950s there was a strongly held view among microbiologists, often called the 'principle of microbial infallibility', that all chemicals were susceptible to microbial degradation if the right organism and conditions could be identified. The metabolic capabilities of microbes were 'all powerful'. The growing realization that many chemicals that found their way into the environment were not being degraded and were persisting for many years put an end to such thinking. Half-lives of

polychlorinated organic insecticides in the order of tens of years were reported and there was considerable fear that such compounds could be with us forever. With these rather frightening statistics in view, Martin Alexander reviewed biodegradation in 1965 and raised the possibility that microbes were, in fact, fallible and that, through a combination of a chemical's structure and the environment it found itself in, a chemical might survive in the environment for prolonged periods. Since then there has been intensive research into this problem world-wide in both academia and industry. Although there are still many causes for serious concern, there is now hope that some of the mistakes of the past will not be repeated and, indeed, that some old pollution problems may be cleared up. Particularly important developments have included:

1. An understanding of how different types of recalcitrant chemicals are degraded and what structural factors make them recalcitrant. This has allowed the design of important chemicals with degradability in mind.
2. The discovery of co-metabolism.
3. The realization of the role of microbial communities rather than pure cultures in degradative processes.
4. The realization that degradation of many chemicals can occur in anaerobic environments.
5. The discovery that white rot fungi have the ability to degrade a huge range of recalcitrant chemicals.

2 DEFINITIONS

It is useful in starting this chapter to define and discuss a few important terms that will be used repeatedly.

2.1.1 Recalcitrant

This term means difficult or obstinate, not easy to control. In the current context, recalcitrant means *difficult, but not necessarily impossible, to degrade*. Recalcitrant chemicals may resist biodegradation for a whole range of reasons and some are more resistant than others. Many recalcitrant chemicals are xenobiotic in nature, but by no means all xenobiotics are recalcitrant. Furthermore, many natural compounds and materials (e.g. lignin) are degraded only with difficulty.

2.1.2 Xenobiotic

Xenobiotic means 'foreign to life'. It is a term applied to many recalcitrant organic chemicals. These are synthetic chemicals and are not found in nature. They will contain structural elements that are unknown or rare in nature and/or are assembled in a structure which is not of natural occurrence. Examples of such structural elements are the aromatic sulphonic acids (found for example in alkyl benzene sulphonate surfactants and many dyes); polychlorination of an alkane or aromatic compound; and the diazo bond. Examples are given in Fig. 34.1.

Some researchers also use the term '*xenobiotic concentration*', meaning a concentration that is not found in nature – so we may have a natural compound present at a xenobiotic concentration. Such a concentration might occur as the result of, for example, an oil spill or discharge of an industrial effluent.

2.1.3 'Hard' and 'soft'

The terms '*hard*' and '*soft*' are used quite widely to describe, respectively, recalcitrant chemicals and ones that can easily be degraded.

2.1.4 Biodegradation

Biodegradation is the term given to the breakdown of organic chemicals by the biological action of a living organism. In the environmental context, generally microorganisms are the most important agents of biodegradation. Although extensive degradation of some xenobiotic chemicals can occur in mammals (usually in the liver), they are not particularly important in degradation of environmental pollutants. The heterotrophic bacteria are often considered to be of prime importance, however, the role of fungi is being increasingly recognized. Algae and cyanobacteria can catalyse some biodegradative processes but are probably of only limited importance. Varying degrees of biodegradation can occur according to how much the compound has been modified.

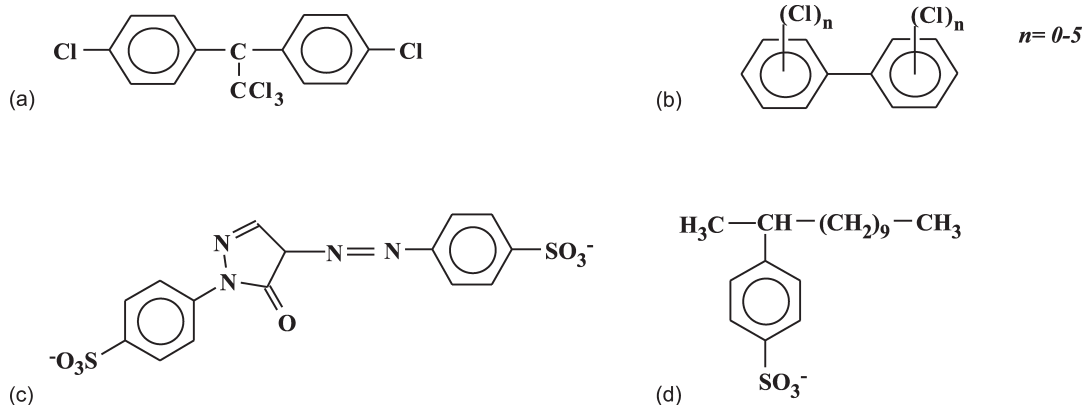


Fig. 34.1 Examples of recalcitrant xenobiotic compounds. (a) DDT (1,1,1-trichloro-2,2-di(chlorophenyl)ethane) an insecticide; (b) PCB (polychlorinatedbiphenyl) used in electrical transformers; (c) Tartrazine, an azo dye used as a food colourant; (d) Linear alkyl benzene sulphonate, a surfactant.

2.1.5 Primary biodegradation

This term is used to describe processes in which a compound loses its characteristic properties but may be little altered in terms of its size or complexity. The compound will have lost some characteristics and may no longer respond in a particular assay. Good examples of primary biodegradation are to be found in the degradation of surfactants (Fig. 34.2). Anionic surfactants are often assayed by the methylene blue active substance (MBAS) test. Positive results are given by compounds that contain a large hydrophobic group linked to a hydrophilic group. If the hydrophobic and hydrophilic groups in a compound are separated then it will no longer give a response in the assay and will no longer have surfactant properties.

Another example (Fig. 34.2) might be the reductive decolorization of an azo dye. This primary biodegradation leads to a change in colour or its removal, but two aromatic amines are released. Depending on which amines they are, these could be much more harmful to the biota than the original dye. Several aromatic amines are, for example, actual or suspect carcinogens.

Thus, a large portion of a molecule may remain intact – this portion may have a BOD and COD and may be toxic or may have other environmental effects, it may also be recalcitrant to further degradation. In conclusion,

primary biodegradation may herald the complete destruction of a pollutant or may cause even more problems.

2.1.6 Mineralization

Mineralization is synonymous with *ultimate biodegradation* or *complete biodegradation*. It describes the degradation of a compound to its mineral components, i.e. carbon dioxide and water. Depending on the compound's composition, other minerals may be released; these might include sulphide, sulphate or sulphite; ammonia, nitrite or nitrate; phosphate or phosphite; chloride; fluoride etc. If the mineralization is anaerobic, methane may be a product. In addition to the carbon dioxide etc. released, some of the mass of the chemical may be converted to biomass (cellular components). In fact this is usually, but by no means always, the case. Thus the complete destruction of a chemical may result in only 50–80% of the carbon being converted to CO₂.

2.1.7 Co-metabolism

This term is used (and often mis-used) to apply to a range of different types of degradative process. Perhaps the best definition is *the degradation of a compound in the obligate presence of another compound(s)*. It is used to describe a situation in which a microbial culture is only able to degrade a substance in the presence

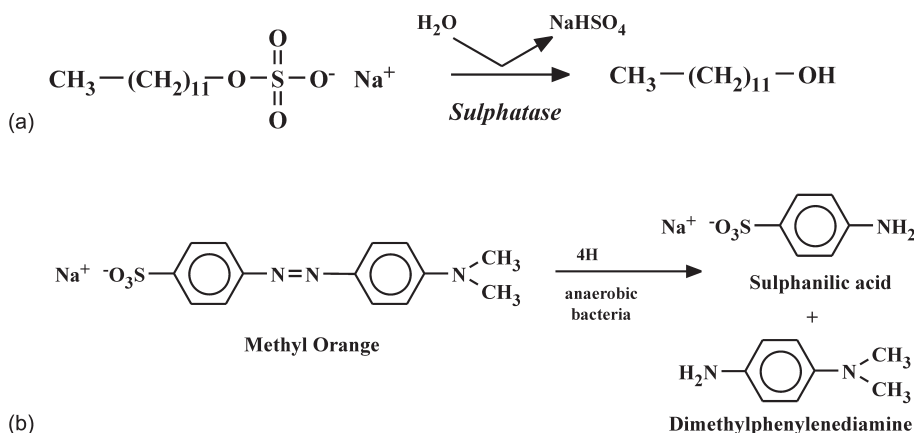


Fig. 34.2 Examples of primary biodegradation. (a) The surfactant sodium dodecyl sulphate (sodium lauryl sulphate) is hydrolysed to dodecanol and sulphate. Removal of the sulphate ester group prevents the molecule acting as a surfactant and giving a positive MBAS test. (b) Reduction of methyl orange by anaerobic intestinal bacteria results in decolorization and production of sulphanilic acid and N,N-dimethyl-4-phenylenediamine.

of other compounds. These compounds may be specific in some cases, while in others a wide variety of *co-substrates* (additional compounds) may be suitable.

Specific co-substrates are required in the case of a process often called *analogue enrichment*. This is where the recalcitrant compound and the co-substrate are structurally related and the co-substrate has a role in inducing the production of some, or all, of the enzymes required to degrade the recalcitrant compound which cannot itself act as an inducer. This process can only work if the degradative enzymes are sufficiently non-specific to degrade compounds other than their natural substrate. The target compound may be mineralized or only suffer partial biodegradation. For example, the addition of biphenyl has been reported to encourage biodegradation of the analogous polychlorinated biphenyls (PCBs).

Non-specific co-metabolism generally occurs in cases in which the degrading organisms derive no obvious benefit from the degradation of the recalcitrant compound. The degrading organisms cannot use energy or carbon resulting from the degradation process and therefore need to be provided with a source of carbon and energy on which they can grow. Such processes are sometimes referred to as *gratuitous degradation* (i.e. it is for free) or alternatively as *fortuitous* or *accidental degradation* (i.e. it occurs by chance). There are many examples of this type of co-metabolism. Occasionally degradative processes are reported to involve co-metabolism when the degradative organism simply requires small amounts of a growth factor (e.g. a vitamin or amino acid) – this is not really an appropriate use of the term.

2.1.8 Microcosms

The term *microcosms* has been used to describe mixed microbial populations which carry out degradative reactions. It really means little more than the terms 'mixed cultures' or consortia. It would be more helpful perhaps to use the term *community degradation*. Some compounds can be degraded completely by single microbial strains in pure cultures, but this is not always the case. It is quite common to find that a compound can only be degraded by a mixed

culture; none of the individual strains present being able to use the recalcitrant compound as a sole source of carbon and energy. In some cases, this requirement for a mixed culture is apparently obligate while in others researchers can sometimes find pure culture degradation and in other circumstances find a need for a mixed culture. There are several possible reasons for the requirement for a mixed culture:

1. *Cross-feeding of growth factors*. In this process one organism may produce a growth factor, such as a vitamin or amino acid, which is required for the primary degradative organism to grow. The process may be mutual, i.e. two organisms will produce growth factors that each promote the other's growth. Cross-feeding between bacterial strains often occurs in nature and also in activated sludge populations.
2. *Removal of accumulated toxic intermediates*. Often in degradation processes intermediates can accumulate in the organism or the growth medium. Such intermediates may be toxic to the degradative organism and so the growth in the mixed culture of another organism that can remove these intermediates, thus relieving toxicity, will be of value. The requirement for such a detoxifying organism will clearly depend on the concentration of the initial substrate and the accumulated toxin. Thus, at high initial substrate concentrations, this interaction may be obligatory while at low concentrations it may not.
3. *Sequential degradation*. It is not uncommon to find that an organism can only catalyse certain initial steps in a degradation or only degrade certain parts of a recalcitrant molecule. In these circumstances, complete degradation of the compound can only be achieved by the action of two or more microbial strains working together and carrying out different parts of the degradative process.

Some consortia/mixed cultures may involve two or more of these interactions. In nature, as opposed to the research laboratory, it is very likely that most biodegradative processes are

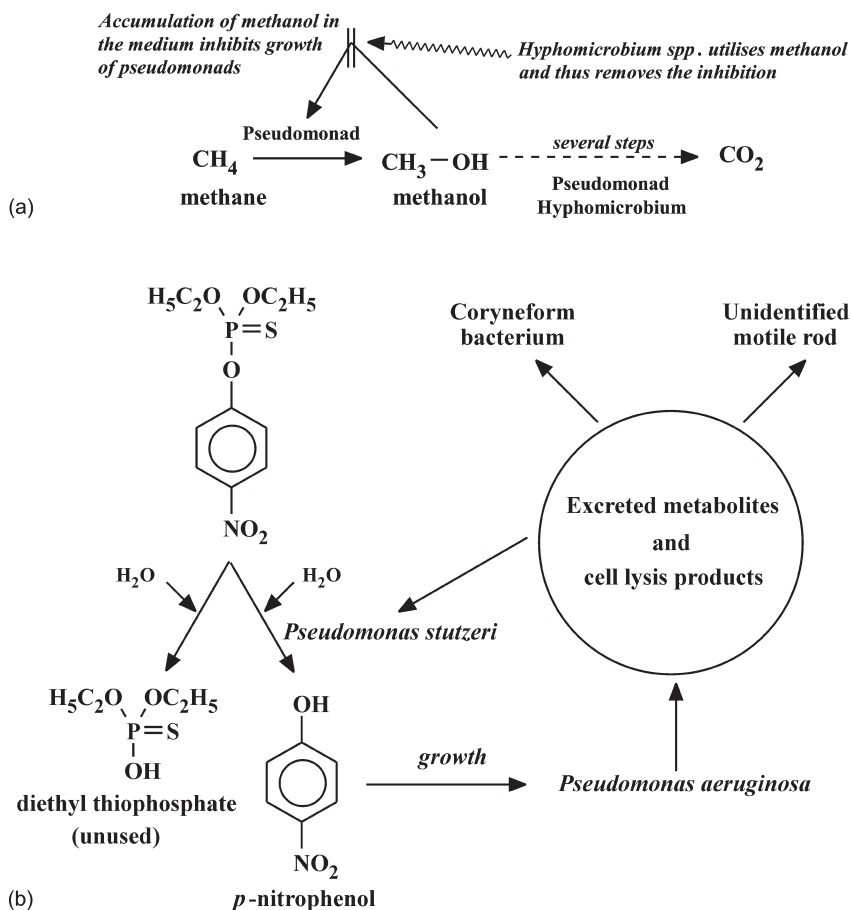


Fig. 34.3 Examples of community degradation processes. (a) Relief of inhibition during growth of a mixed culture on methane. (b) Degradation of parathion by a mixed microbial culture. Initial degradation was catalysed by a strain of *Pseudomonas stutzeri* which could not utilize the hydrolysis products. This organism and the other two were dependent on metabolites of the *p*-nitrophenol-utilizing *P. aeruginosa*.

dependent on mixed cultures. Pure cultures are rarely found outside the laboratory, however, they are of great scientific importance in allowing us to identify the biochemical and genetic mechanisms that are of importance in degradation processes. Some examples of the involvement of mixed cultures in biodegradation are given in Fig. 34.3.

3 FACTORS INFLUENCING THE BIODEGRADATION OF RECALCITRANT COMPOUNDS

In order to understand why some recalcitrant organic compounds can persist for a prolonged time in the environment, or survive effluent

treatment processes, it is essential to study the factors that influence their degradation. These can be crudely separated into factors associated with the compound and factors associated with the environment. They are summarized in Table 34.1.

The failure of a compound to be degraded, or its slow biodegradation, may be due to environment-related factors or to chemical-related factors or to a combination of both. Sometimes several of the factors listed from both categories may be involved to varying degrees. It is perhaps useful to give more details and some examples of the effects of these various factors on degradative processes.

TABLE 34.1 Factors which influence the biodegradation of recalcitrant compounds*1) Chemical-specific factors*

State	Gas, liquid or solid
Solubility	Surface area
Hydrophobicity	Aqueous
Adsorbability	Ability to dissolve in hydrophobic (lipophilic) solvents, hydrophobic compounds will have relatively low water solubility
Size and shape	Ability to adsorb to and complex with organic and inorganic fractions in soil, sediments and water
Charge	Can affect ability to permeate cell membrane and to interact with enzymes
Toxicity	Can affect ability to permeate cell membrane and to interact with enzymes
Detailed molecular structure	May be a specific effect or a general one
	Important factors include:
	i) Presence of an 'easily metabolizable' structural unit (s)
	ii) Presence of a 'difficult to metabolize' structural unit (s)
	iii) Presence of an 'unnatural' (xenobiotic) structural unit (s)
	iv) Degree of branching
	v) Nature of substituents
	vi) Number of substituents
	vii) Position of substituents
Concentration	Too high or too low

2) Environment-specific factors

Biotic factors	Presence of suitable or 'potentially suitable' organisms
Abiotic factors	
Physical	Temperature or pressure
Chemical	Nutrients – minerals, growth factors
	Presence of oxygen as:
	i) terminal electron acceptor
	ii) substrate
	iii) inhibitor
	Presence of alternative electron acceptors:
	i) nitrate
	ii) sulphate
	iii) carbon dioxide
	PH
	Inhibitory materials
	Soil type
	Moisture level
	Type of water – fresh, brackish, saline

3.1 Chemical-specific factors

3.1.1 State/solubility/hydrophobicity

These factors are grouped together since they all influence the availability of a compound. For biodegradation to occur a chemical must either be in intimate contact with an organism or its enzymes (the enzymes may be intracellular or extracellular). Biological, enzyme-catalysed reactions normally occur in an aqueous milieu

with the enzyme and its substrate being dissolved in water. The availability of a compound may be influenced by its physical state. Gases permeate well but may only be present at very low concentrations (see below). Liquids and solids are often present at apparently very high local concentration but, if they are not readily soluble, then the concentration available to microbes may in reality be very low and this may retard degradation. In such cases the rate

at which a compound is metabolized may depend on the surface area available, as this will dictate the rate at which it can dissolve into the aqueous phase and become available to the enzymes which degrade it.

Sometimes biodegradative reactions involve the adhesion of the degrading organisms to their insoluble substrate – this is often the case for lipids and for hydrocarbons, such as mineral oils. When this is so, surface area is again important and if the material is finely divided into small particles, a greater surface area will be available for the attachment of degrading microbes. This will apply in the cases of environmental contamination during ‘oil spills’ – if the oil is dispersed it will degrade faster than if it accumulates in large masses. For solid hydrocarbons, such as polyaromatics, surface area will be important. Materials that aid solubilization (e.g. surfactants) may promote biodegradation. In fact, some microbes can themselves produce surfactants which improve degradation reactions. Evidence on the effects of adding synthetic surfactants on rates of biodegradation is equivocal.

There is a tendency for materials that are of limited water solubility to be relatively resistant to biodegradation, but it must be emphasized that this is not always the case. For example, toluene is only slightly soluble in water but is fairly degradable while some sulphonated aromatic compounds are very soluble but highly resistant to degradation. Hydrophobicity is inversely related to water solubility and, as such, affects the availability of compounds. Also hydrophobic materials tend to coalesce into droplets of low surface area, again restricting their availability.

A further complication is that hydrophobic organic compounds have a tendency to *bioaccumulate* in the fatty tissues of higher organisms. Thus the concentration in the higher organism will be greater than that in the surrounding environment. Once present in these tissues, *biomagnification* or *bioconcentration* may occur, a process by which the concentration of a material increases in organisms higher up the food chain. Sometimes a compound may bioaccumulate in an animal to levels thousands or even millions of times greater than those

found in its environment. In one example, PCBs were present in water at a concentration of 2×10^{-6} ppm, in phytoplankton at 2.5×10^{-3} ppm, in zooplankton at 0.123 ppm, in rainbow trout smolt at 1.0 ppm, in lake trout at 4.83 ppm and in herring gulls at 124 ppm. This represents a biomagnification in the order 10^8 .

A compound may cause little damage at lower trophic levels, but as it increases in concentration the damage increases. The biomagnification of organochlorine pesticides had disastrous effects on the populations of birds of prey and fish-eating birds in the 1950 and 1960s. Populations plummeted and it took many years for these populations to recover after the withdrawal of the pesticides. Another problem with bioaccumulation is that once accumulated in fatty tissues of animals or birds, a recalcitrant compound may be sequestered from microbial degradative processes until the animal dies and thus it may survive for a more prolonged period in the environment. Thus bioaccumulation can increase the persistence of a compound in the environment. The hydrophobicity of a compound and the likelihood of bioaccumulation can be assessed by determination of its octanol:water partition coefficient (often referred to as K_{ow}) – the higher the coefficient the more likely it is that bioaccumulation will occur. Often values of K_{ow} are so high that they are given as logarithm of K_{ow} . The types of compounds that are likely to bioaccumulate are hydrocarbons and organochlorines (e.g. DDT, lindane, aldrin and polychlorinated biphenyls – PCBs).

3.1.2 Adsorbability

Some chemicals have a propensity to adsorb to other materials, e.g. particular fractions of soils. If the adsorption is ‘tight’ then it may reduce the availability of a compound to microbes and enzymes thereby restricting or preventing its biodegradation. A classic example of this are the herbicides paraquat and diquat, which adsorb very tightly to certain clay fractions (notably montmorillonite) in soils. In these circumstances virtually no biodegradation occurs even in the presence of organisms which can degrade the herbicide. The amount of degradation will depend on the amount of

the herbicide and the adsorbent material present in the soil. Some chemicals may adsorb to the organic (humus) fraction rather than to clay minerals.

3.1.3 Size and shape

For a chemical to be degraded it will usually have come into contact with an enzyme. Often, but not always, the degradative enzymes are intracellular, i.e. they are located inside a microbial cell and therefore within the cytoplasmic membrane – a selective permeability barrier that controls access to the cell. Some chemicals, by dint of their size, cannot pass across the membrane and therefore cannot come into contact with the enzymes therein. Thus, if they cannot be attacked by extracellular enzymes they will remain undegraded.

For an enzymic reaction to occur the substrate must enter into intimate contact with the enzyme and form an enzyme/substrate complex. Some recalcitrant compounds may be unable to complex with the active site of the enzyme due to their size, and probably more so, to their shape.

3.1.4 Charge

Highly charged molecules, particularly those with large negative charges, have great difficulty in penetrating cell membranes. Compounds with many sulphonic acid groups, e.g. some azo dyes and surfactants may come into this category.

3.1.5 Toxicity

It is fairly obvious that there may be problems in the biodegradation of chemicals that are toxic to microbes. Their biodegradation will clearly depend on their concentration and the susceptibility of the microbes. There will be similar problems with chemicals that are not generally toxic but inhibit a specific key enzyme in the degradative pathway. Many recalcitrant chemicals are toxic to some extent to some or most microbes; examples include most phenols (especially chloro- and nitro- derivatives) and fungicides. Toxic compounds may kill susceptible microbes (-cidal effects) or merely inhibit their growth (-static effects). The mechanisms of toxicity will vary considerably and can include

inhibition of vital processes (like respiration), protein denaturation and membrane disruption. It may in some cases be possible for cells to evolve to become resistant, or tolerant, to a toxic compound. So, for example, most bacteria are inhibited or killed by toluene, which damages their membrane structure, but some can develop resistance and some bacteria have been reported to survive in the presence of 50% v/v toluene!

3.1.6 Concentration

Concentration is a major factor in the fate of recalcitrant compounds in the environment. If concentration is too high, problems of toxicity may arise (see above). However, concentrations may be too low. For example, in one study on degradation of the 2,4-D in stream water, about 75% mineralization occurred in 8 days when the herbicide was present at 2.2×10^{-2} or 2.2×10^{-4} g/l. However, at 2.2×10^{-6} or 2.2×10^{-8} g/l there was less than 10% mineralization. The lack of biodegradation of compound at very low concentrations can be accounted for in two ways. First, at low concentrations the compound (or its metabolites) may not be present at a sufficient level to induce the formation of the enzymes required to degrade it. Second, if evolution of new metabolic capacities is required, low concentrations of recalcitrant compounds may not provide enough carbon and energy to give newly evolved organisms a selective advantage. It is therefore highly desirable that compounds are degraded when present in relatively high concentrations (e.g. in industrial effluents) rather than escaping undegraded into the wider environment in which they will be greatly diluted and their degradation will be less assured.

3.1.7 Detailed molecular structure

Certain features of a chemical's structure are likely to make it either easier or more difficult for microorganisms to degrade. Ultimately, it is the properties of an entire molecule, not its components, that decides its fate. However, it is important to know how different subunits may influence the degradation process.

1. Easily metabolizable units: certain structures are easily metabolized by microbial enzymes. These include the ester (carboxylic,

sulphate and phosphate) and amide bonds, which are common and widely distributed in nature. Furthermore, all are susceptible to hydrolysis and thus do not need special coenzymes. Fig. 34.4 shows some of these easily metabolizable structures and their degradation.

Another relatively easily degradable bond is the amine bond. However, amines

are much less degradable than amides and, in general, are cleaved oxidatively rather than hydrolytically. Amine-degrading enzymes are usually oxygenases, oxidases or dehydrogenases. As amine cleavage involves a redox reaction, the enzymes often need co-factors and they are therefore less likely to operate extracellularly.

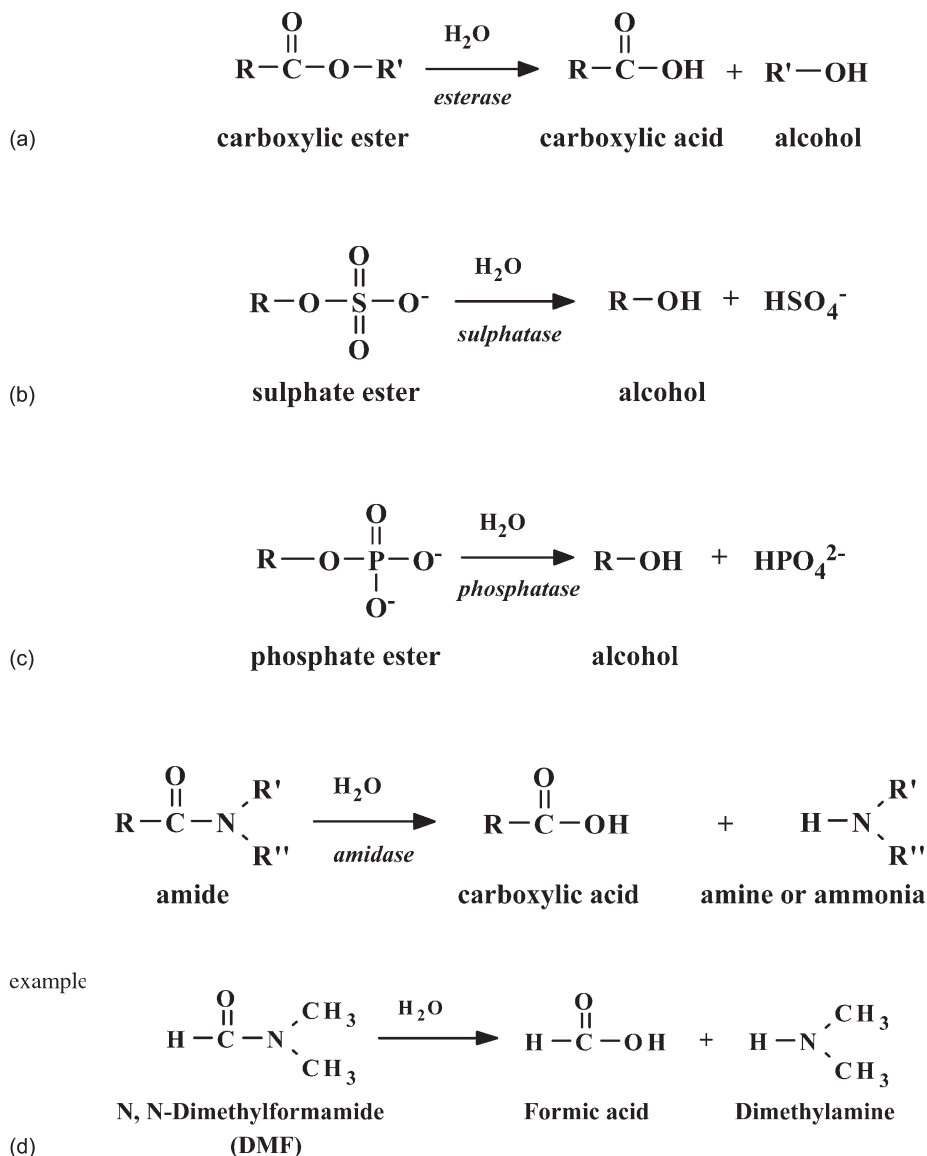


Fig. 34.4 Examples of easily degradable structural units. (a) Carboxylic ester linkage. (b) Sulphate ester linkage (example given in Figure 34.2). (c) Phosphate ester linkage. (d) Amide linkage (where R, R' and R'' can be an alkyl group or hydrogen).

2. Units which are difficult to metabolize: some structural units are difficult to metabolize because they are not amenable to the normal processes of metabolism. A good example is the quaternary carbon atom, i.e. a carbon atom that has attached to it four other carbon atoms. Such structures can occur in alkyl chains and are resistant to degradation because they cannot participate in the process of β -oxidation (they prevent the formation of a carbon-carbon double bond) which is used to degrade alkanes and fatty acid or alcohols. Quaternary carbon structures can occur in, and restrict the biodegradability of, alkyl benzene sulphonate surfactants or hydrocarbons.
3. Presence of xenobiotic structural units: by definition, xenobiotic structural units have not been present in our biosphere for a long period. Such units only became common with the rise of synthetic organic chemistry in the 19th century and so microbes will have only had a relatively short interval in which to adapt to their presence compared with the geological time-scale that has been available for evolution to degrade 'natural chemicals'. Nevertheless, microbes, and particularly bacteria, can evolve very rapidly and some enzymes have evolved to degrade xenobiotic compounds. In other cases organisms may happen to possess enzymes of broad specificity, which can catalyse fortuitous metabolism of xenobiotics (with no certain advantage to the microbe).

It is difficult to be certain whether a particular structure is, or is not, truly xenobiotic. So, for example, carbon-halogen bonds are not common in nature and natural compounds containing them only make up a tiny proportion of the organic matter on earth. Chloramphenicol, trifluoroacetic acid and thyroxine are examples of natural compounds with C-Cl, C-F and C-I bonds. Most organohalogens are truly xenobiotic and polychlorination of organic molecules is not natural. Nitro groups do occur naturally but are uncommon. Probably the best examples of xenobiotic structural units are the diazo linkage, aromatic (but not

aliphatic) sulphonic acids and polychlorinated aromatic or aliphatic compounds; all these features retard biodegradation. Methyl, methoxy, amino and hydroxyl groups are common in nature but may, of course, be part of a xenobiotic compound.

4. Degree of branching: in alkanes or compounds with alkyl chains, the degree of branching is an important factor influencing biodegradability. Simply stated, the more branched a compound the less degradable it will be. Furthermore, branching in certain positions tends to be particularly problematic. So, for example, two branches on the same carbon atom (see quaternary carbon above) or on adjacent carbon atoms are particularly problematical. The effects of branching on biodegradability have been noted particularly for alkanes and for alkyl benzene sulphonate surfactants (see Section 4).
5. Nature of substituents: some substituents of a basic chemical structure have the effect of increasing its biodegradability and others of decreasing degradability. The effects of substituents can be direct or indirect. So, for example, a substituent may indirectly decrease degradability by increasing toxicity, by decreasing solubility, by increasing hydrophobicity, by increasing ability to adsorb to soil or by altering charge. Direct effects tend to be related to changes in electron distribution, steric effects or hydrophobicity, which alter the compound's ability to participate in enzyme reactions.

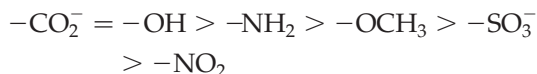
Electron distribution is always an important consideration; enzyme reactions tend to be either electrophilic or nucleophilic and the possibility of a reaction occurring (and its rate) depends, *inter alia*, on the density of negative or positive charges at certain positions in the molecule. Chemical substituents usually either donate or withdraw electrons, thus altering charge density. The magnitude and position of these changes in charge depend on the chemical and its substituent. There is no simple rule that electron donation or withdrawal is beneficial or adverse, it depends entirely on the nature of the enzyme reaction under consideration. Some will be accelerated by

substituents that withdraw electrons and others by substituents that donate electrons.

Steric effects are related to the bulk of substituents. If a substituent is large it may cause *steric hindrance*. This is a phenomenon whereby a substituent may interfere with a reaction simply by dint of its size and position. Large substituents may prevent a potential substrate entering the active site of the enzyme or prevent it taking up the correct orientation at the enzyme's active site. For a reaction to occur, certain parts of the substrate must be aligned with certain regions of the enzyme, if a large structural unit is present this alignment may be impossible and thus the reaction may not occur.

The correct alignment of a substrate at the active site of an enzyme may also depend on hydrophobic or charge interactions or on hydrogen bonding. Changes in hydrophobicity or charge may prevent this alignment and thus prevent formation of the enzyme-substrate complex prior to catalysis.

A good example of the effects of substituents on biodegradation is provided by a study of the aerobic degradation of substituted benzene derivatives by soil microflora. The time taken for decomposition of benzoate, phenol, aniline, anisole, benzene-sulphonate and nitrobenzene was 1, 1, 4, 8, 16, and > 64 days respectively. Thus in terms of their ability to improve degradation:



Clearly nitro- and sulphonic acid groups tend to retard degradation (the same is also true of chloro- and other halogen groups). For aerobic attack on aromatic rings, structural groups that have an electron withdrawing effect (e.g. chloro-, nitro- and sulphonate) often decrease the rate of degradation because in withdrawing electrons they inactivate the aromatic ring to electrophilic attack.

It must be emphasized that electron withdrawing groups do not always have an adverse effect on degradation, so, for example, when the initial attack is nucleophilic, electron withdrawing groups may

increase reaction rates. This is the case, for example, for the anaerobic reduction of the diazo bond in azo dyes.

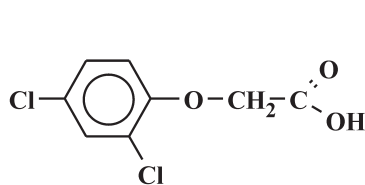
Very good quantitative correlations have been obtained in some instances between the coefficients that describe electronic effects of substituents and biodegradation rates. However, this is not always the case because other factors, like steric effects and hydrophobicity, can complicate the picture. The effects of particular substituents may well be different in aerobic and anaerobic conditions.

6. Number of substituents: as a general rule, although not an absolute one, biodegradability decreases with increasing number of substituents. Aerobic degradation often involves the introduction of hydroxyl groups onto the carbon skeleton. The more substituents there are, the less free carbons there will be for hydroxylation. If an unsubstituted carbon cannot be hydroxylated then a substituent will need to be removed, which may be more difficult.

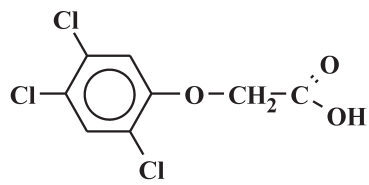
A good example of the effect of number of substituents is to be found in the aerobic degradation of the chlorinated phenoxyacetate herbicides (Fig. 34.5). Many pure microbial cultures have been described which can readily degrade 2,4-D and these can be easily obtained by enrichment culture. 2,4,5-T differs only in the presence of one extra chlorine atom, but it is much more resistant to biodegradation. It takes much longer to isolate 2,4,5-T-degrading bacteria by enrichment culture and only a few pure cultures have been isolated that can degrade it as a sole carbon source.

PCBs (polychlorinated biphenyls) (see Fig. 34.1) were widely used industrial chemicals, but are now banned for most applications. The commercial preparations were mixtures of many congeners (isomers). Several studies have investigated the relationship between the number and pattern of substitution and biodegradability. Aerobic biodegradation is favoured by:

- i) the presence of fewer chlorine atoms
- ii) the presence of chlorine substituents on only one aromatic ring



2,4-D - 2,4-dichlorophenoxyacetic acid
(readily biodegradable)



2,4,5-T - 2,4,5-trichlorophenoxyacetic acid
(highly recalcitrant)

Fig. 34.5 The structure of 2,4-D and 2,4,5-T.

- iii) the absence of chlorine atoms in certain positions (the 2, 6, 2' or 6').

Conversely in anaerobic degradation of chlorinated compounds, particularly aromatics, it often appears that reductive dechlorination is easier with the more highly chlorinated compounds.

7. Position of substituents: as has already been mentioned for branching, the position of substituents can have a major bearing on degradability of a compound. This effect can be due to either steric or electronic factors. Perhaps the best examples of positional effects are found with positional isomers of disubstituted aromatic compounds and of the insecticide lindane.

Disubstituted aromatic compounds have three isomers usually known as *ortho*, *meta* and *para* or 1, 2, and 3, respectively (Fig. 34.6).

A study on the effect of position of substitution on the aerobic degradation of a range of disubstituted benzenes by soil microflora showed large differences in the degradability of *o*, *m* and *p* isomers. In general, *o* and *p* isomers of a compound were of similar degradability to each other, while the *m* isomer differed in its

degradability. This is because the patterns of electron distribution are similar for *o* and *p* isomers but differ in the *m* isomer. Often *o* and *p* isomers are more degradable than the *meta* isomer, but this is not always so and it clearly depends very much on the effects of particular combinations of substituents on the electron density in particular parts of the molecule. A typical example is given in Fig. 34.7. In some cases, while *o* and *p* isomers of substituted benzoic acids are degraded in similar times, the *ortho* isomer is somewhat more recalcitrant. This is probably due to steric effects.

Lindane is the trade name of the γ -isomer of hexachlorocyclohexane (HCCH), formerly, and inaccurately, known as benzenehexachloride (BHC). This recalcitrant compound consists of a cyclohexane ring (in the chair conformation) substituted at each carbon atom with a chlorine atom and a hydrogen atom. At first sight it may appear that all carbon atoms are equivalent to each other and therefore that no isomers would exist. This is not so. HCCH is a planar molecule and the chlorines and hydrogens can be disposed either roughly in line with the plane of the ring or at an angle to it or both and several positional isomers are possible (Fig. 34.8).

It has been shown that there are considerable differences between isomers in terms of their degradability (Fig. 34.9). These differences cannot be accounted for by differences in electron distribution and must be related to steric effects – the distribution of the bulky chlorine atoms presumably interfering with the degradative process in some isomers.

In multisubstituted chloroaromatic compounds, e.g. chlorinated phenols or benzoic

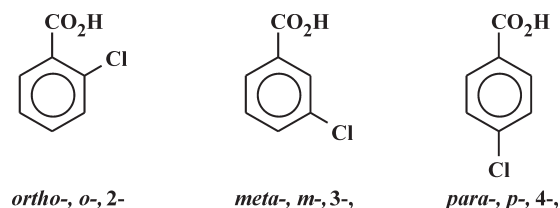


Fig. 34.6 Structural isomers of chlorobenzoic acids. Typical disubstituted derivatives of benzene.

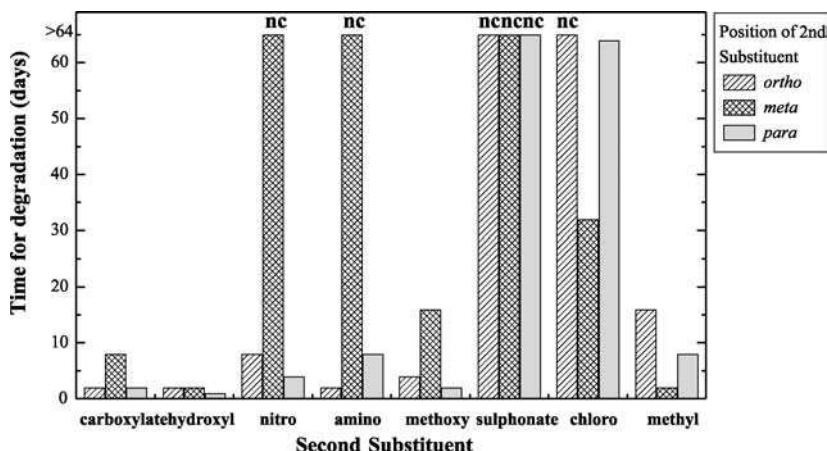


Fig. 34.7 The effect of the position of substitution in benzoic acid derivatives on the rate of biodegradation by soil microorganisms. The time for degradation is the time taken for the total loss of ultraviolet light absorption which equates to cleavage of the aromatic ring. NC indicates that a compound was not completely degraded in 64 days.

acids, it may be possible to predict which bonds will be broken and in which order. Bond strength of carbon-chlorine bonds can be predicted using computer models and, on this basis, the likelihood of particular bonds being cleaved reductively by unacclimated anaerobic microbial consortia has been determined. Very good correlations between predictions and experimental results have been obtained, with the most negatively charged carbon-chlorine bond being the first to be reduced.

In conclusion, it should be emphasized that while the susceptibility of a compound to biodegradation can sometimes be accounted for by one structural factor, more commonly a combination of several factors governs its fate. In some cases a compound may be less degradable than might have been expected on theoretical grounds. Such deviations from expected relationships may be due to factors like steric hindrance and toxicity.

3.2 Environment-specific factors

A whole range of environment-specific factors can influence a compound's fate in the environment. These can act alone or in concert and are discussed in detail below.

3.2.1 Presence of suitable organisms

For a compound to be degraded the presence of suitable organisms is essential. By 'suitable' we mean organisms that already have some ability to degrade the target compound. 'Potentially suitable organisms' (see Section 6 on evolution) are ones which initially cannot degrade a compound but have the potential to evolve to carry out the required biodegradation. Clearly, degradation of a target chemical will occur more rapidly after its exposure to 'suitable' rather than 'potentially suitable' organisms, because the time period during which evolution occurs will not be required. Nevertheless, these 'suitable' organisms are often present in very small numbers and, until a large population of suitable organisms has developed, little degradation will be observed. The 'lag' period before degradation is observed is known as the *acclimation* period and it may be due to the time taken for evolution to occur or the time taken for a large population to develop, or both. The time taken for the microbial population to increase to a 'critical mass' will depend on both the original number of suitable organisms present and on their growth rate. It will also depend on the number of organisms needed to make an observable effect on substrate concentration and clearly

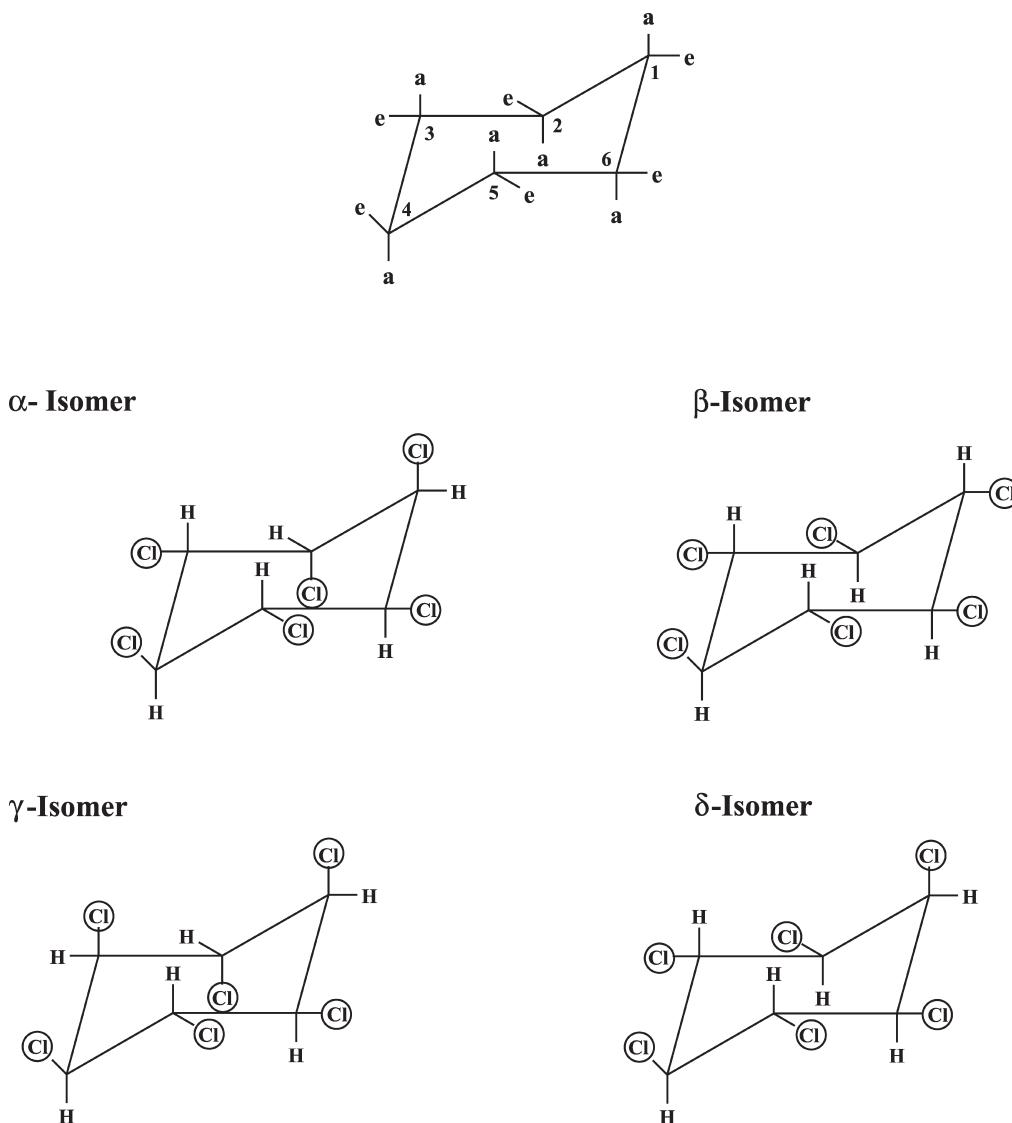


Fig. 34.8 Structural isomers of hexachlorocyclohexane (Lindane). Each carbon atom carries one chlorine atom; the position of the chlorine atom is either approximately in the plane of the ring (equatorial – e) or at an angle to the plane (axial – a). There are several possible isomers. The four most abundant are α , β , γ and δ .

this will be a function of the sensitivity of the analytical methods used. Obviously it is very difficult to generalize about this, but the number needed to make an impact may be in the order of 10^4 – 10^6 bacteria/ml.

Thus, if a chemical enters an environment with very few organisms in it then degradation will either be slow or will not occur. Pristine water in the upper reaches of a stream, ground water (in aquifers) and subsoil all have only

very small microbial communities, thus recalcitrant compounds entering these environments will be degraded slowly, if at all. The presence of acclimated organisms, which have previously been exposed to a target compound, can greatly accelerate biodegradation. This has been shown repeatedly with degradation of herbicides, surfactants, industrial chemicals and mineral oils, in both soil and in river systems. Microbes in river water that has

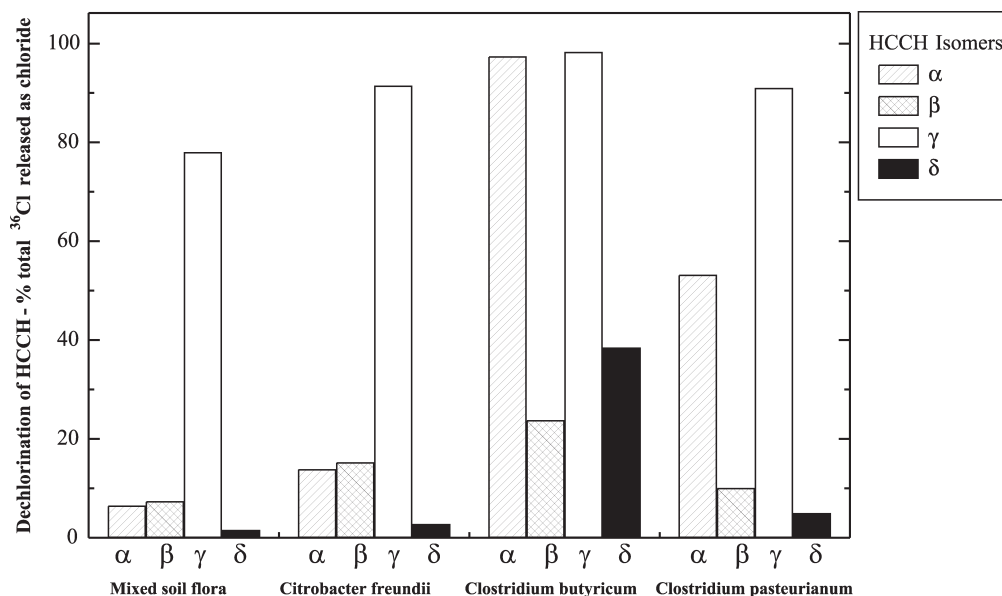


Fig. 34.9 The effect of isomer structure on the anaerobic degradation of hexachlorocyclohexane (HCCH) by microbial cultures. Biodegradation was co-metabolic with glucose as co-substrate. Cultures were incubated for 6 days at 25°C with 10 mg/1 HCCH. Degradation was assessed by measuring the release of radio-labelled ³⁶Cl from HCCH as chloride ions.

suffered industrial pollution will degrade target chemicals faster than those from unpolluted streams.

3.2.2 Physical factors

In general, temperatures that are too high or too low will retard, or even prevent, biodegradation. All organisms have their own optimum, maximum and minimum temperature for growth and degradative reactions. It should be noted that in regions of very high (e.g. hot springs) or low (e.g. polar regions) temperatures, it is likely there will only be a restricted and specialized microflora and thus the chances of organisms with the requisite degradative capabilities being present is small. However, there has been little research specifically on the biodegradation of recalcitrant organics by thermophiles and psychrophiles.

In addition to influencing the composition of the microflora and rates of reaction, changes in temperature may alter the physical state of a compound or its solubility – such changes may be beneficial or adverse to the chances of degradation.

Another physical factor is barometric pressure; high pressures, such as those found in the abyssal depths of the oceans, will restrict the diversity of the microflora with similar consequences to extremes of temperature.

3.2.3 Nutrient availability

In order to grow, microbes need a range of nutrients. The simplest requirements are for nothing but mineral salts and a source for carbon and energy. Most microbes that degrade organic pollutants are heterotrophs and the carbon and energy source can be a single compound (often a target recalcitrant compound). The major elements required are carbon, nitrogen, phosphorus, sulphur, oxygen and hydrogen, with smaller amounts of iron, magnesium, calcium and potassium. Sometimes specific metals are needed for particular enzymic reactions. For example, cobalt will be required if the degradative enzyme contains cyanocobalamin (Vitamin B₁₂) as a prosthetic group (this cofactor is important in some dechlorination reactions in certain organisms). Manganese will be required by white rot

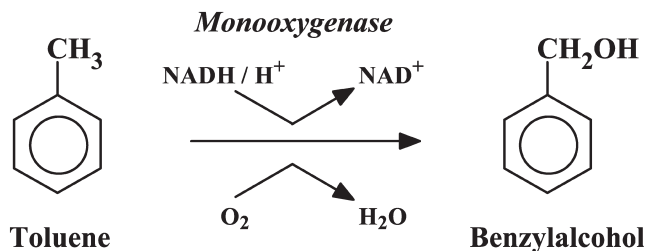
fungi if manganese peroxidase is important. If important elements are missing or in short supply growth will be impossible or unbalanced and degradative reactions may be slow or impossible. Some microbes also need the presence of growth factors; these are organic chemicals which the organism cannot make itself and must have if it is to grow. Growth factors may include specific amino acids, nucleotides or vitamins/cofactors. Some cofactors may not be essential but may, nonetheless, increase the rate of specific degradative reactions. For example, in some cases, the addition of flavine nucleotides (e.g. FAD) can increase the rate of anaerobic reduction of azo dyes.

3.2.4 Presence of oxygen

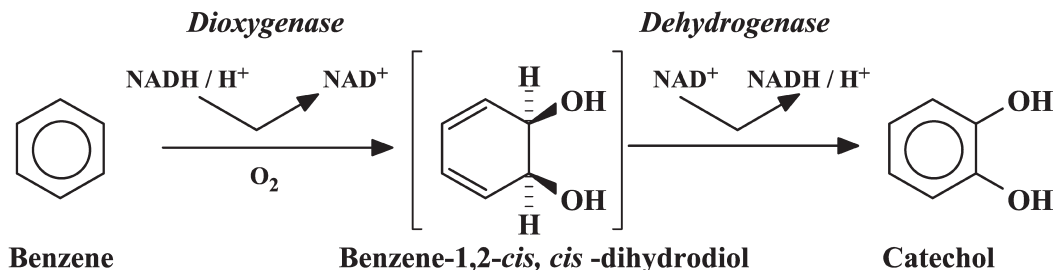
Oxygen has several roles in microbial growth and in each of them its presence can profoundly influence the occurrence and rate of degradative reactions.

1. Quantitatively the main use of oxygen is as a terminal electron acceptor during respiration in aerobic growth. Biological

oxidations often involve the removal of hydrogen and electrons (reducing equivalents), which must be coupled to an oxidizing agent – the terminal electron acceptor. In this process energy is produced and intermediate electron acceptors are regenerated so that more oxidation can occur. Strict aerobes cannot grow without oxygen and if it is in short supply, their growth will be slower than normal and metabolism may be atypical. Thus, in the absence of oxygen, biodegradative reactions that depend on strict aerobes will not occur. At one time it was thought that oxygen was essential for degradation of most recalcitrant xenobiotic compounds, it is now known that this is not the case. Indeed, many compounds are degraded preferentially, if not uniquely, in anaerobic conditions. There are a few examples of compounds whose degradation requires oxygen, e.g. the degradation of lignin and the complex lignosulphonates produced in wood pulping appears to have an obligate requirement for oxygen.



(a) Toluene Monooxygenase



(b) Benzene Dioxygenase

Fig. 34.10 Examples of monooxygenase and dioxygenase reactions.

2. Oxygen may also react directly as a substrate in certain enzymic oxidations. There are many examples of this type of involvement in which molecular oxygen is incorporated into an organic molecule – these are called oxygenations. Such reactions are often used to activate a relatively inert molecule and lead to further degradation, they are important in many processes for biodegradation of recalcitrant organics. Monooxygenations incorporate one atom of oxygen and dioxygenations two. Some examples are given in Fig. 34.10.
3. Finally, oxygen can act as an inhibitor and thus have a deleterious effect on certain biodegradative reactions. It has been observed in recent years that not only can many xenobiotic compounds be degraded very well in the absence of oxygen, but that some biodegradative reactions are inhibited by oxygen or occur only in its absence. Although some such reactions are catalysed by strict anaerobes, this is by no means always the case and degradative processes catalysed by facultative anaerobes or even obligate respiratory organisms (like pseudomonads) can occur preferentially in the absence of oxygen. For example, many polychlorinated organic compounds like DDT or lindane are degraded better anaerobically than

aerobically. This is also the case for cleavage of the diazo bond. While some simple azo dyes can be decolourized in the presence of oxygen, much better results are obtained anaerobically. It seems that what most of these reactions, which are preferentially anaerobic, have in common is that they are reductive reactions. As such they will only be successful if there is an excess of reducing power. In the presence of oxygen, electrons may be diverted into reduction of oxygen rather than the target compound.

3.2.5 Presence of alternative electron acceptors

In the absence of molecular oxygen some organisms can carry out respiratory metabolism using so-called 'alternative electron acceptors'. These are inorganic compounds that can be reduced, thus allowing energy generation by oxidative phosphorylation. The materials that can be used in this way are listed in Table 34.2. In some cases the organisms using these alternate electron acceptors are normally aerobic, only resorting to this type of metabolism in the absence of molecular oxygen. In other cases the organisms are strictly anaerobic and are often very intolerant of oxygen – dying in its presence. There are now many examples of the degradation of

TABLE 34.2 Use of 'alternative' electron acceptors in biodegradation

<i>Electron acceptor and/or process</i>	<i>Reduced to</i>	<i>Organisms' usual relation to oxygen</i>	<i>Degradation of recalcitrant organics in pure culture</i>	<i>Involvement in centre mixed cultures</i>
Nitrate and oxides of nitrogen— <i>denitrification, nitrate reduction</i>	Nitrogen, N ₂ O	normally aerobes	✓ e.g. toluene, phenol	possible
Sulphate, sulphur, and some oxides of sulphur— <i>sulphate reduction, sulphidogenesis</i>	Hydrogen sulphide	Always strict anaerobes	✓ e.g. aniline, phenol	may act as electron sink; <i>Desulfomonile tiedjei</i> can dechlorinate 3-chlorobenzoate
Ferric ion Fe (III)— <i>iron reduction</i>	Ferrous ion Fe (II)	both normal aerobes and strict anaerobes	✓ e.g. toluene	possible
Manganic ion Mn (IV)— <i>manganese reduction</i>	Manganous ion Mn (II)	both normal aerobes and strict anaerobes	✓ e.g. toluene	Possible
Carbon dioxide— <i>methanogenesis</i>	Methane	Always strict anaerobes	no	methanogens act as an electron sink

recalcitrant organics under strictly anaerobic conditions. Sometimes a pure culture can be isolated while, in others, consortia are involved. The role of sulphate reducers and methanogens in consortia may be to act as an electron sink. They are able to remove hydrogen, which is inhibitory to the fermentative organisms that produced it. These fermentative organisms may degrade recalcitrant xenobiotics with production of hydrogen, but often these reactions appear to be thermodynamically unfavourable without the removal of hydrogen. Some sulphidogens can also dechlorinate or even cleave aromatic rings.

For many years it was considered that some compounds, like aromatic hydrocarbons, were only degraded aerobically. It has now been shown that a range of the monocyclic aromatics can be degraded under a range of different types of anaerobic conditions. This gives hope that eventually such materials which have contaminated oxygen-deficient or anoxic environmental niches like ground water may ultimately be removed, especially if suitable electron acceptors are present. When considering bioremediation of hydrocarbon-contaminated aquifers it may be easy to encourage degradation processes by the supply of nitrate (which is highly soluble) rather than sparingly soluble oxygen.

3.2.6 pH

Clearly all organisms have an optimum pH for growth and for any degradative processes that they catalyse. Changes from optimal pH will clearly decrease rates of biodegradation for particular reactions. It has to be said that most research has concentrated on metabolism at near neutral pH (6–8) and there is very little literature on biodegradative reactions at very high or very low pH. The role of acidity in preservation is well known, common examples being preservation of pickled foods with acetic or lactic acids, silage production and yoghurt manufacture. The acid conditions in peat bogs are also a contributory factor to preservation of materials in that environment. Many fungi have acidic pH optima for growth and some can survive and metabolize at very low pHs – cellulolytic and saccharolytic fungi, for

example, will often reduce the pH of their own growth media to between 2 and 3. However, as a group, the fungi are less able to utilize recalcitrant organics as growth substrates than are the bacteria. Having said this, the white rot fungi are known to be able to degrade a huge range of recalcitrant xenobiotics but, in general, these processes are fortuitous degradations from which the fungi derive little obvious benefit from the degradative process. White rots often operate best in the laboratory as agents of biodegradation at around pH 5. Although these organisms are widespread in the environment, they only grow in a restricted ecological niche (dead wood or dying trees).

Although quite a range of fungi can utilize aromatic compounds at subneutral pH, there has been little systematic study of their abilities. Similarly, despite recent research that has established the ability of many organisms, notably bacteria, to grow at very alkaline pHs, the degradative capacities of these alkalophiles has rarely been determined. As with any other extreme environment we can only expect to find a restricted microflora at extremes of pH and so it is likely that the range of biodegradative capacities of these specially adapted organisms will be less than that found with neutralophiles.

3.2.7 Inhibitory materials

Natural environments and man-made ones (like effluent treatment plants) may contain a wide range of toxic, inhibitory materials, which may either kill microorganisms or inhibit their degradative processes. Some toxic chemicals (e.g. phenol, toluene or cyanide) may inhibit a wide range of microbes, others may be much more specific, only killing particular taxa (e.g. fungi), groups of microbes (e.g. nitrifying bacteria) or even species. Some inhibitors are naturally occurring (e.g. plant phenolics derived from decomposing plant tissues, or heavy metals derived from mineral deposits), others are present as the result of environmental pollution. The concentration of these inhibitors is crucial in determining their effects. Phenol and toluene are degradable at low concentrations but generally extremely toxic

at high concentrations. As always with microbes, there is the possibility that given constant exposure some will evolve to become more resistant to the inhibitory materials. This is certainly true for toluene and for many heavy metals, e.g. mercury.

Salinity is of course an important factor, as increasing salinity generally inhibits microbial growth. There are some obligate halophiles, which not only survive in, but actually need, very high salt concentrations, but this is a small group of specialized organisms, with relatively low genetic diversity. The metabolic capabilities of halophiles have been studied to some extent and appear, as expected, to be much more limited than those of organisms living in less extreme conditions.

3.2.8 Soil type

The composition of a soil may profoundly affect the degradation of materials in the soil. Some soil fractions can adsorb and sequester recalcitrant compounds (see Section 3.1.2 on adsorbability). It is also possible for soil fractions (notably some clays) to bind extracellular enzymes, thus inactivating them and allowing their substrates to survive for a longer time in the soil than would otherwise be possible. This has been shown to occur with materials like proteins, which are normally perishable and so greater problems might be expected with recalcitrant organics that require extracellular enzymes for their degradation. Similarly, some soils contain tannins and similar plant phenolics (derived from decomposing plant remains) which can complex with and inactivate enzymes in a non-specific fashion.

4 'DESIGNER DETERGENTS'

Artificial surfactants, the active agents in detergent preparations, first came into widespread use after the Second World War when they replaced soaps. There are four categories of surfactants: anionic, cationic, non-ionic and zwitterionic and within each category there are many variations on the theme. All consist of a hydrophobic (lipophilic) group coupled to a

hydrophilic group; this latter may have a negative or positive charge, no charge or a variable charge according to pH. By far the most important group in terms of amount used is the anionics of which there are two main types, linear alkyl sulphates and alkyl benzene sulphonates (ABS). Alkyl benzene sulphonates are the principal type of laundry surfactants and these gave rise to enormous problems when they were introduced. The type of formulation used was a material called tetrapropylene alkyl benzene sulphonates (TPS), which were made by polymerizing four molecules of propylene. This gave rise to an indeterminate mixture of many different isomers (all branched). A high proportion of these isomers were recalcitrant to degradation and so they passed through effluent treatment plants unscathed by the microbial populations. On reaching the rivers they did what surfactants are best at – they foamed, especially in regions of great turbulence like weirs and waterfalls. Huge layers of foam developed and were carried down the rivers like sheets of meringue! The foam could be a few feet deep, and it would blow off the rivers and into surrounding countryside or riverside towns. Not only was this an eyesore, but there were also problems of the toxic effects of the surfactants to aquatic life. Furthermore, the foam picked up and transported pathogenic microbes from the sewage polluted rivers. Action was called for and an intense research programme was established to determine how a biodegradable surfactant could be produced.

The following observations were made:

1. the presence of a quaternary carbon near the end of the alkyl chain prevented degradation
2. the degree of branching was important; the more branches present, the worse the degradation – a preparation of TPS will typically be very resistant to biodegradation with only a small part of the mixture succumbing easily to degradation
3. in ABS surfactants with straight alkyl chains, the length of the chain affects degradability. Chains with about 12 carbon

atoms being optimal, longer or shorter chain lengths delayed but did not prevent biodegradation

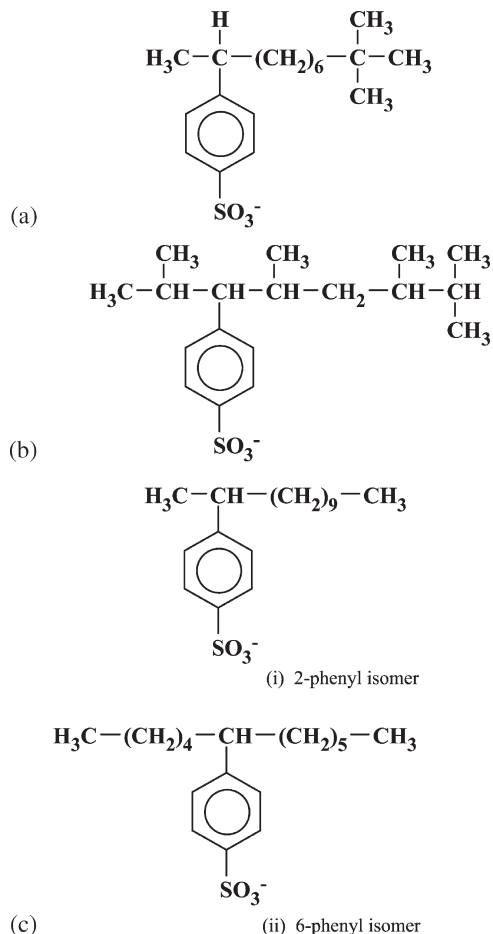


Fig. 34.11 The effect of structure on degradability of alkylbenzene sulphonate detergents. (a) An alkyl benzene sulphonate detergent with a terminal quaternary carbon atom. This is highly resistant to biodegradation since the initial attack is at the end of the alkyl chain and the quaternary carbon prevents β -oxidation. (b) This structure is a typical tetrapropylene alkylbenzene sulphonate (TPS). TPS comprises of a mixture of isomers which differ in the position of the methyl branches and the phenyl group. Most isomers are very resistant to biodegradation. (c) These structures represent typical linear alkylbenzene sulphonates (LABS) the phenyl group can be on any alkyl carbon except carbon-1. The 2-phenyl derivative (i) is more degradable than the 6-phenyl isomer (ii).

4. in straight chain ABS molecules the benzene sulphonic acid group can be attached in several positions. It is generally the case that degradability is enhanced by attachment close to the end of the alkyl chain and decreased by attachment near the middle of the chain.

Thus the ideal ABS surfactant will have a straight alkyl chain about 12 carbon atoms long with the benzene ring attached near the end of the alkyl chain and there will be no branching. Typical structures of these surfactants are shown in Fig. 34.11.

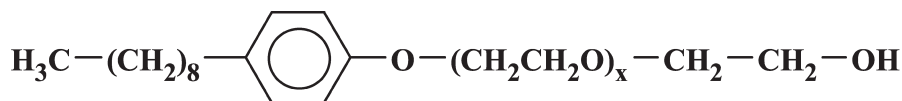
By investigating the effect of the various structural elements on degradability of a surfactant it was possible to work out the optimum structure for a readily degradable alkyl benzene sulphonate surfactant. So-called 'soft' surfactants were produced and subjected to trials in certain parts of the UK with excellent results. 'Hard' ABS surfactants were largely replaced following a voluntary agreement in 1964 between manufacturers and the UK government.

Surfactant pollution is now largely a thing of the past on most rivers in the UK. The only real exception being found on a few rivers where the local textile industry still uses certain types of hard non-ionic surfactants. The problem compounds are commonly known as 'APES' (alkylphenol polyethoxylates). They are still required for certain functions, although for many applications they have been replaced by the 'softer' alkyl ethoxylates (Fig. 34.12).

Among the ethoxylate non-ionic surfactants, degradation is favoured by:

1. a linear hydrophobe rather than a branched one
2. an alkyl rather than an alkyl phenol hydrophobe
3. a relatively short ethoxylate chain.

Non-ionic surfactants can include propoxy or butoxy groups as well as or instead of ethoxylate groups. These larger alkoxy groups tend to decrease degradability.



(a) Nonyl phenol ethoxylate – a typical alkyl phenol ethoxylate surfactant (APE).
The number of ethoxyl units (x) can be in the range 2 to 50.



(b) An alkyl ethoxylate (fatty alcohol ethoxylate surfactant - FAE).
The number of methylene units (y) can be in the range 7 to 16. The number of ethoxyl units (z) can be in the range 2 to 30.

Fig. 34.12 Structures of ethoxylate non-ionic detergents.

5 ACCLIMATION OF MICROBIAL POPULATIONS TO DEGRADE RECALCITRANT ORGANICS

Acclimation is the process by which a microbial population adapts to degrade a compound to which it is exposed. Given a population which, on initial exposure, appears to have no significant degradative ability against a target

compound, acclimation may involve three different processes. The acclimation or 'lag' period can vary greatly in duration. The different processes and possible time-scales are summarized in Table 34.3.

Acclimation to degrade a target compound can be lost as well as gained. Loss of acclimation can occur not only for xenobiotics, but even for central metabolites like glucose.

TABLE 34.3 The mechanisms of acclimation and the processes involved

<i>Mechanism</i>	<i>Processes involved</i>	<i>Relative speed</i>	<i>Possible timescale</i>	<i>Factors influencing rate</i>
Induction of enzymes or transport systems	The organisms already have potential to make the required enzymes etc. but need presence of the target compound to stimulate production	Fastest	Minutes to hours	Rate of metabolism
Growth of population of suitable microbes	A few 'suitable organisms' are present but have to grow until the population is large enough to make an observable change in the concentration of the target compound	Intermediate	Hours to weeks	Growth rate; initial number of organisms
Evolution (see Section 6)	Mutations required to change an organism so that it can now degrade the target compound. One or more mutations may be needed in one or more genes. Once mutations have occurred the size of the mutant population has to increase – see above	Slowest	Weeks to months	Number of organisms with 'potential'; mutation rate; number of mutations needed; growth rate

It can occur for three reasons: absence of degradative enzymes in the absence of inducer (see above); genetic instability (see Section 11); and loss of a specific group of degradative organisms (see Section 14.3).

6 EVOLUTION OF DEGRADATIVE ABILITIES

Assuming that it is not toxic, the ability of an organism to degrade a compound depends on the ability of the compound to come into contact with an enzyme or a series of enzymes which can degrade it.

This is affected by three things:

1. access of the compound to the enzymes
2. ability of the enzyme to catalyse a degradative reaction
3. production of the enzyme in suitable quantities.

All three of these may be subject to evolutionary change.

Evolution may involve the mutation of pre-existing genes and thus the production of new, altered proteins. It can also involve the acquisition of new genetic information from other organisms. This new information may come via plasmid transfer, transposons (jumping genes) or by uptake of DNA from the environment.

Sometimes extracellular enzymes are involved in a biodegradation, in which case the ability of compounds to enter the cell is not a problem. If enzymes are intracellular, then the compound needs to cross the cell membrane. This can involve free diffusion or use of a permease or other transport system. If the compound cannot easily enter the cell, there is the possibility for improvement in access to the cytoplasm, due either to alterations in the structure of the cell membrane (this could involve changes in either proteins or lipids) or to changes in the specificity of the permease proteins which catalyse translocation of compounds across membranes.

Enzymes may already exist that catalyse metabolism of similar compounds. Small changes in the structure of the protein may

alter its substrate specificity such that a new substrate can be metabolized. Such changes can occur either in the regions which catalyse the reaction or in regions responsible for binding of the substrate into the enzyme/substrate complex. It should be noted that some enzymes are naturally of broad substrate specificity (see Section 9).

Where an organism already possesses genes for the production of an enzyme which can metabolize a target compound, degradation may not occur due to a failure of the organism to produce the requisite enzymes. This can be overcome by mutations that lead to constitutive production of the enzyme (i.e. the enzyme is produced all the time) or to a change in the inducer specificity for the enzyme such that the target compound is now an inducer.

It should be noted here that there are often large differences in the specificity of transport systems and enzymes, and also of the control systems which regulate their production. Thus, when a new metabolic activity arises in an organism, it may be due to mutation (followed, of course, by selection) in one or several of the functions mentioned above. Studies on the evolution of new metabolic activities in organisms have shown that often the early stages of evolution involve selection of constitutive mutants or mutants that produce very large amounts of an enzyme and this is then followed by the selection of mutants with altered enzyme activity. Alterations in enzyme activity are more likely to involve changes in the regions of the protein that involve substrate binding than changes in catalytic sites.

One stage in the evolution of an organism to degrade a toxic compound may involve the organism becoming tolerant to the toxic effects. This tolerance may be due to a range of different adaptations depending on the mechanism by which the compound causes inhibition.

7 METABOLIC VERSATILITY

It is interesting to observe that, although some microbes are very specialized in the type of compounds they can degrade or utilize as growth substrates, others may not be. This is

particularly so for bacteria in a few genera which show extreme nutritional versatility, i.e. the ability to degrade many organic compounds. It is not uncommon to find individual bacterial *strains* (not species) that can utilize over 100 defined organic compounds as sole source of carbon and energy. This trait is most famous in the aerobic pseudomonads. The old genus *Pseudomonas* has now been dismantled by taxonomists and split into a range of new genera. Some of the most versatile species include *Pseudomonas aeruginosa*, *P. putida*, *P. fluorescens*, *P. stutzeri*, *Burkholderia cepacia* and *Comamonas testosteroni*. However, the pseudomonads are not the only group possessed of nutritional versatility. Other noted Gram-negative bacteria include *Acinetobacter*, *Alcaligenes*, *Moraxella*, *Achromobacter* and *Flavobacterium* spp. For many years the qualities of the Gram-positives were rather overlooked – possibly because the nutritionally versatile species are often rather slow-growing in comparison to the pseudomonads. The Gram-positives most noted for their ability to degrade recalcitrant chemicals are all in the actinomycete line and include *Mycobacterium*, *Nocardia*, *Rhodococcus* and *Arthrobacter* spp. The organisms most frequently identified as agents of biodegradation tend to belong to the genera named above, although other genera, e.g. *Streptomyces*, are occasionally encountered.

All of the organisms listed above are primarily aerobes, however, with the increasing appreciation of biodegradation in anaerobic conditions, it is likely that there will soon be a considerable number of strict or facultative anaerobic bacteria to add to the list. For example, degradation of aromatic compounds under denitrifying conditions has been demonstrated for members of the genera *Azoarcus* and *Thauera*, organisms which were little known until recently but are likely to prove important. Similarly, the significance of sulphate reducers in biodegradation has only just become apparent with the identification of new organisms like the aromatic-degrading *Desulfobacula toluolica*.

Fungi are to some extent the 'Cinderellas' of biodegradation, their role having been largely ignored. In the last 10 years their abilities have

been more recognized with the realization of the potential of white rots (discussed in more detail in Section 12). However, outside the white rots they are still largely ignored.

8 METABOLIC PATHWAYS

Clearly with the plethora of different recalcitrant chemicals, a very wide range of metabolic pathways are involved in biodegradation. If a compound is to serve as a carbon and energy source, then it has to be converted into a form that can enter central metabolism. Normally this involves converting it into one, or more, low molecular weight intermediates of the tricarboxylic acid (TCA) cycle (otherwise known as Krebs's or the citric acid cycle) or compounds that feed into it. The means by which this is done obviously varies considerably – typical intermediates include acetate (or acetyl CoA), acetaldehyde, pyruvate, succinate or fumarate. The length of metabolic pathways varies enormously according to the complexity of the target compound; the number of steps ranges from one or two to over a dozen. In some cases target compounds require very little modification before they can enter a pre-existing pathway.

It should be stressed that the metabolic pathways used for a degradation may vary not only according to the environmental conditions (e.g. aerobic or anaerobic) but also according to the type of organism. There may be a wide range of pathways available, some of which differ from each other subtly and others markedly. It is quite likely that different strains of the same species will employ different metabolic pathways for the same degradative process. For this reason it is of little value to reproduce here details of lots of metabolic pathways, as no assumptions can be made as to which pathway will be used for a particular degradative process without proper testing being done. Generally, there is no obvious reason why one strain uses one pathway and another strain a different pathway. Indeed, in effluent treatment plants there does not appear to be any evidence as to which pathways are in practice used by the microflora.

One of the best examples of multiple pathways is found with toluene degradation (Fig. 34.13). Two pathways have been proposed for anaerobic degradation and at least five for aerobic degradation. Two of the five are from different strains of *Pseudomonas putida*.

Many recalcitrant chemicals are aromatic and it is perhaps useful to look briefly at how aromatics are degraded aerobically. There are basically three different sets of pathways. To begin with the aromatic compound is converted to a ring cleavage intermediate which normally has two hydroxyl substituents. These may be next to, or opposite, each other. This intermediate is then cleaved by a dioxygenase to give a straight chain intermediate. If the hydroxyl groups are next to each other then the ring is either cleaved between them (*ortho* cleavage) or to one side of them (*meta* cleavage). Where the hydroxyls are opposite each other the ring is cleaved immediately to one side of one of the hydroxyl groups. The straight chain ring cleavage product is degraded to give smaller units, which will enter the tricarboxylic acid cycle. The initial pathway which produces the ring cleavage substrate may be simple, involving only one step (e.g. hydroxylation of phenol to produce catechol) or complex involving many steps. Fig. 34.14 shows the early stages of the three main types of pathway. It should be noted that a range of analogues may act as ring cleavage substrates, but often (not always) distinct, specific enzymes are involved. So, for example, in *Pseudomonas putida*, catechol 1,2-oxygenase employed to cleave catechol is distinct from protocatechuate 3,4-oxygenase which cleaves its carboxylated analogue in the *ortho* pathway.

Anaerobic degradation of aromatics proceeds in a very different way since it is not possible to activate the benzene ring by the addition of oxygen. Instead the ring is activated by converting alkyl side chains, or adding carbon dioxide, to give a carboxylated ring, which is then converted to a coenzyme A ester. Benzoyl CoA or its *para* hydroxy derivative appears to be a common intermediate and this is progressively reduced to a cyclohexyl derivative which is primed for ring cleavage by the addition of water across a double bond next

to the carboxyl group. After ring cleavage the straight chain intermediate is degraded to acetyl CoA by β -oxidation. Thus, in this type of attack the oxygen required is introduced from water and oxidation is accomplished by removal of hydrogen/electrons. These are generally disposed of by the reduction of nitrate or sulphate or the generation of hydrogen, which is utilized in consortia by methanogenic or sulphidogenic.

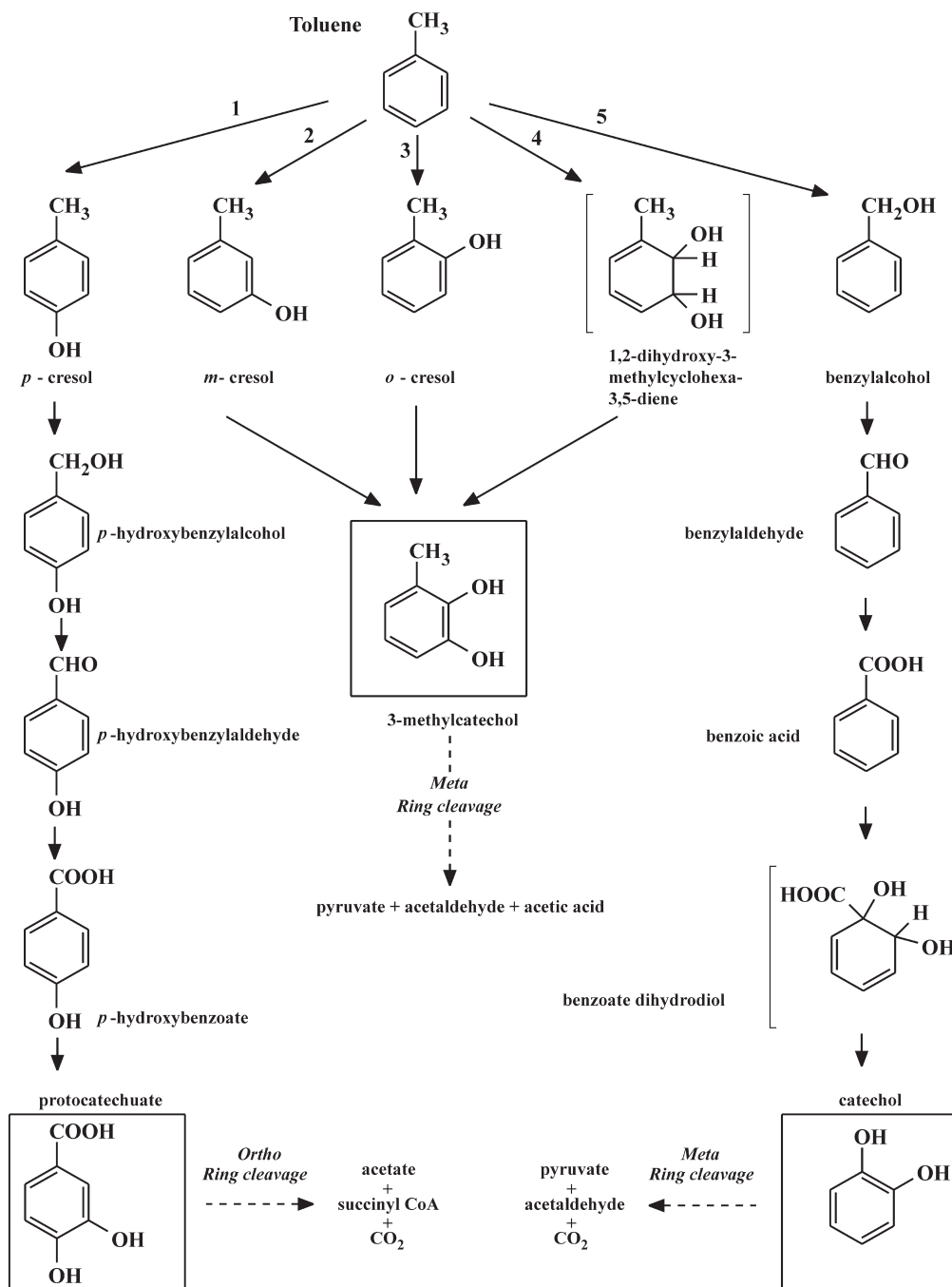
9 NON-SPECIFIC ENZYMES

Some types of co-metabolism depend on the fact that certain enzymes are sufficiently non-specific to catalyse degradation of compounds other than their 'normal' substrate. Several enzymes are well known for the wide range of substrates that they can modify, some of which have little resemblance to their 'normal' substrate. These enzymes include methane monooxygenase, ammonia monooxygenase and toluene dioxygenase. Examples of some of the substrates they modify and reactions they catalyse are given in Table 34.4.

10 REDUCTION BY CO-ENZYMES

Co-enzymes are relatively small organic molecules often derived from vitamins. Some are reactants in enzyme-catalysed reactions while others are essential parts of many enzymes (sometimes called prosthetic groups). Co-enzymes are often important in enzymes carrying out oxidation or reduction reactions and they may participate as electron carriers.

It has been observed that in some cases the co-enzyme on its own can catalyse important biodegradative reactions with recalcitrant compounds. For example, cyanocobalamin and other similar derivatives of vitamin B₁₂ are able to catalyse reductive dechlorination reactions. Compounds dechlorinated in this way include PCBs, hexachlorobenzene, carbon tetrachloride, chloroform, dichloromethane and chlorinated ethanes. Some other co-enzymes involving transition metals, e.g. nickel and



Toluene is degraded via 5 different routes by the following bacteria; *Pseudomonas mendocina* KR1 (1), *P. pickettii* PKO1 (2), *Burkholderia* (formerly *Pseudomonas*) *cepacia* G4 (3), *P. putida* F1 (4), *P. putida* mt-2 (5). Postulated intermediates are shown in parentheses. Intermediates immediately prior to ring cleavage are shown in boxes. -----► indicates several steps.

Fig. 34.13 Different pathways for the aerobic biodegradation of toluene.

Name of pathway	ortho-cleavage intra-diol cleavage; β -ketoadipate pathway	meta-cleavage; extra-diol cleavage	gentisate/ homogentisate pathway
Typical ring cleavage substrate	catechol; protocatechuate; 3 or 4 methyl catechol	catechol; protocatechuate; 3 or 4 methyl catechol	gentisate; homogentisate
Typical ring cleavage product	<i>cis,cis</i> -muconate; β -carboxy- <i>cis,cis</i> -muconate	2-hydroxymuconic semialdehyde; 2-hydroxy-4-carboxymuconic semialdehyde	maleylpyruvate; maleylacetoacetate
Typical degradation products	Acetyl CoA, succinate		fumarate; pyruvate; acetoacetate; acetate

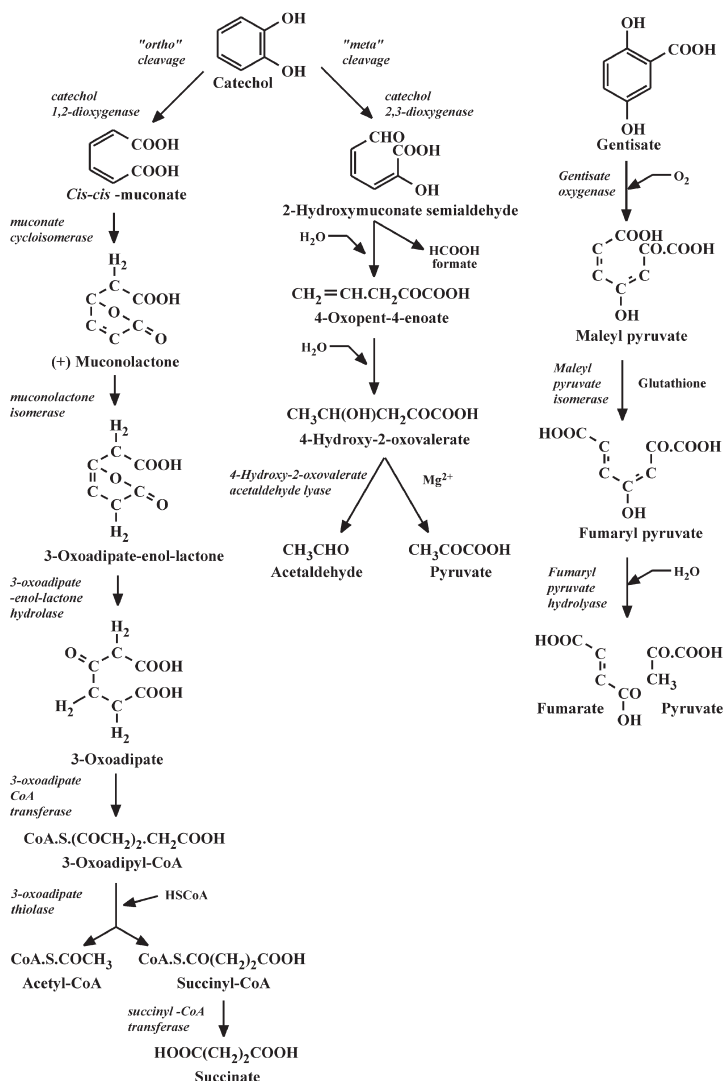


Fig. 34.14 Shows the major metabolites involved in aerobic aromatic degradation pathways.

TABLE 34.4 Examples of 'non-specific' enzymes and the reactions they can catalyse

Enzyme	Organism	Normal reaction	Examples of substrates metabolized or reactions catalysed
Ammonia monooxygenase	<i>Nitrosomonas europaea</i>	$\text{NH}_3 \rightarrow \text{NH}_2\text{OH}$	dichloroethylene; trichloroethylene; vinyl chloride
Methane monooxygenase	Methanotrophs e.g. <i>Methylosinus trichosporium</i> ; <i>Methylococcus capsulatus</i>	$\text{CH}_4 \rightarrow \text{CH}_3\text{OH}$	chloromethane to formaldehyde; chloroform to CO_2 ; dichloromethane to CO ; pyridine to pyridine N-oxide; toluene to benzyl alcohol
P450 _{cam}	<i>Pseudomonas putida</i>	camphor to hydroxy camphor	reductive dechlorination of chlorinated methanes and ethanes
Propane monooxygenase	<i>Mycobacterium vaccae</i>	propane to propanol	cyclohexane to cyclohexanol; dichloroethylene; trichloroethylene; vinyl chloride
Toluene dioxygenase	<i>Pseudomonas putida</i> F1	Toluene \rightarrow 3 methylcatechol	trichloroethylene to glyoxylate, formate and chloride

iron, can have similar effects. Reduced flavin nucleotides (FMN and FAD) are thought in some cases to mediate azo dye reduction.

It is possible that use may be made of some of these processes at least to facilitate primary biodegradation of some pollutants. The co-enzymes may have certain advantages over cell-free enzymes as biocatalysts. Notably they may be more stable. It is possible to immobilize some co-enzymes for use in bioreactors. One factor that has to be borne in mind is that most of the reactions catalysed are redox reactions and that for continuous use it will be necessary to return the co-enzyme to the correct oxidation state. In some cases this can be accomplished using chemical reagents, in others use of microbes is a possibility.

11 GENETIC ASPECTS

The genes that encode for biodegradation of many recalcitrant organic chemicals are often encoded on plasmids. Sometimes only one or two enzymes are encoded on a plasmid with the rest on the bacterial chromosome, while in others, an entire pathway is plasmid encoded.

Plasmids are small genetic elements separate from the chromosome and are often

unstable. There is a range of causes for their instability but often a plasmid will be lost from an organism if it is cultivated in the absence of the substrate whose degradation it encodes. Plasmids tend to remain stable in a population when their presence confers a selective advantage (e.g. ability to utilize a substrate which is present) on individual cells which carry them. When no advantage accrues to the organism, plasmids can be lost because their maintenance in the population represents a drain of resources which can impose (at least in the short term) a selective disadvantage.

Plasmids can often be transferred easily between organisms, even ones that are not closely related, and this transfer offers a means by which genetic recombination can occur. This happens in nature but it can also be made to happen in the laboratory giving scientists the potential to develop new bacterial strains with altered and enhanced degradative ability. Various researchers have genetically engineered so-called 'super bugs', which can degrade a wider range of, for example, hydrocarbons. The potential application of these strains in clearing up oil spills or other pollution incidents has been patented and much 'hyped'. In practice, it seems unlikely that most of these genetically

engineered organisms will really offer much advantage over mixed cultures developed by conventional enrichment culture, or indeed over the native microflora. Such strains are more likely to be of use in the more carefully controlled conditions of special treatment plants.

Genetic engineering may have a role in creating bacterial strains with entirely new metabolic activities by combining genes from different organisms to produce an entirely new metabolic pathway. An example of this is the creation of a new strain of *Pseudomonas* to degrade chlorobenzoates by using part of the *ortho* pathway from one organism with part of the *meta* cleavage pathway from another. It is likely that such strains will also arise as a result of 'natural' genetic recombination.

Table 34.5 gives a list of some of the recalcitrant organics which can, at least in some organisms, be degraded by plasmid-encoded enzymes.

It is important to remember that enzymes that are plasmid encoded in one organism are

not necessarily plasmid encoded in other organisms, or even other strains of the same species.

12 WHITE ROT FUNGI

One of the most interesting areas of study in the field of biodegradation in recent years has been that of degradation by white rot fungi. These basidiomycete fungi are commonly found growing on dead or dying trees. Their main growth substrates are cellulose and hemicellulose, but to gain access to these substrates they have first to remove the complex aromatic polymer lignin, which encases it. The organisms cannot utilize lignin as a growth substrate, and indeed to remove it they need the presence of an easily degraded substrate. White rots have evolved a very interesting mechanism for the removal of lignin, which has been likened to 'enzymic combustion'. Lignin is not a typical polymer, it has several kinds of subunits and these are assembled randomly in a 3D-structure with several different types of bonds. It is not amenable to simple hydrolysis.

White rots produce a variety of extracellular enzymes which, between them, generate, with the aid of hydrogen peroxide (made by the fungus), chemical oxidants that destabilize the lignin and render it liable to oxidation or hydrolysis. The enzymes vary from fungus to fungus but generally include manganese peroxidase and/or lignin peroxidase and sometimes laccase. The chemical oxidants include Mn^{3+} ion chelates and free radicals or radical cations of veratryl alcohol but others are almost certainly involved. This system has evolved to degrade one of nature's most complex and recalcitrant materials and it has to be non-specific to deal with the wide range of structural units and bonds in lignin. It comes as no real surprise that this system is sufficiently non-specific to allow it to degrade a huge range of recalcitrant xenobiotics – many, but not all, of which are aromatic in nature. The list of chemicals degraded by white rots is long and some examples are given in Table 34.6.

TABLE 34.5 Examples of substrates whose degradation is, at least in part, plasmid-encoded

Substrate degraded	Organism
toluate, toluene, xylenes	<i>Pseudomonas putida</i>
<i>p</i> -toluidine	<i>Pseudomonas putida</i>
salicylate	<i>Pseudomonas putida</i>
octane, hexane, decane	<i>Pseudomonas oleovorans</i>
camphor	<i>Pseudomonas putida</i>
nicotine, nicotinate	<i>Pseudomonas convexa</i>
2,4-D/methylchloro phenoxyacetate	<i>Alcaligenes eutrophus</i> , <i>A. paradoxa</i>
aniline	<i>Pseudomonas</i> spp.
naphthalene	<i>Pseudomonas putida</i>
fluoroacetate	<i>Moraxella</i> spp.
chlorinated biphenyls	<i>Alcaligenes</i> , <i>Arthrobacter</i>
S-ethyl N,N-di-isopropyl thiocarbamate	<i>Arthrobacter</i>
parathion	<i>Pseudomonas diminuta</i> , <i>Flavobacterium</i> spp.
geraniol	<i>Pseudomonas putida</i>
phenanthrene	<i>Mycobacterium</i> spp.
morpholine	<i>Mycobacterium chelonae</i>
chloropropionic/ chloroacetic acid	<i>Alcaligenes</i>
chlorobenzoate	<i>Alcaligenes</i>

TABLE 34.6 A list of some of the recalcitrant compounds and materials reported to be degraded by white rot fungi

<i>o</i> -cresol	PCBs
Catechol	DDT
Creosote	Lindane
Chlorophenols, pentachlorophenol	Polycyclic aromatics – e.g. pyrene, benzopyrene, anthracene
2,4,5-T	Trinitrotoluene
Azo dyes – e.g. Orange II	Dioxins
Heterocyclic dyes	Chlorolignins
Triphenyl methane dyes – e.g. crystal violet	Lignosulphonates Kraft pulp effluent
Chloro anilines	Cotton black liquor

The type of information available on these biodegradations varies very much. In some cases it is clear that target compounds are mineralized extensively while in others only primary degradation may be occurring. Some chemicals, notably dyes, can be metabolized at relatively high concentrations (grams per litre) while others have only been studied at microgram per litre concentrations. What is clear is that the range of chemical structures degraded is large and, in terms of their degradative abilities, white rots are among the most versatile of microbes. The type of compounds most studied in this context are the chlorinated hydrocarbons, polyaromatics and dyes and coloured effluents. With coloured effluents, the main aim is to decolorize but, in some cases, COD removal is also possible. There is evidence to suggest that with some materials part of the COD may be metabolized but not removed. However, it may be converted to a form that is susceptible to biodegradation by other organisms such as those in activated sludge. Thus, although the white rots may not always remove COD, they may be able to 'soften' hard COD, thus enabling its ultimate removal.

One drawback to the use of these organisms is their obligate requirement for the provision of an easily degradable co-substrate like glucose or starch. The co-substrate is required to maintain viability and to provide reducing equivalents for hydrogen peroxide production. Some white rots can only work effectively in nitrogen-limited conditions (which promote

synthesis of their ligninolytic enzyme systems), however, this is not the universal requirement it was once thought to be and there is considerable variation between fungal strains. The fungal mycelia once produced can be re-used many times and can be employed without loss of effectiveness in continuous or semi-continuous reactors for several months. Another advantage is that once produced, the mycelium can be kept in cold storage for several months without loss of activity.

There has been much interest in the use of white rot fungi to clean-up land polluted by recalcitrant organics. Studies involve inoculating the soil with a suitable white rot fungus (usually *Phanerochaete chrysosporium*) and providing a readily degradable carbon source to encourage its growth. Trials involving both *in situ* and *ex situ* bioremediation have been performed. The success of these trials has been rather patchy and it seems likely that white rots are not well adapted to the soil environment. They are, after all, normally inhabitants of wood, a more specialized ecological niche. In the soil there will be much more competition for the 'readily degraded nutrients' and it is likely that the relatively slow growing white rotters will be unable to compete effectively. Without their easily degraded carbon source they will be unable to degrade lignin or the recalcitrant target chemicals in the soil. They also work best under pH conditions (e.g. pH 4–5) which are considerably more acidic than those pertaining in normal soils.

One problem in this research field is that there has been too much emphasis on one organism, *Phanerochaete chrysosporium*, which is not a typical white rot. More recently a wider range of organisms, e.g. *Coriolus* (*Trametes*) *versicolor*, *Pleurotus ostreatus* and *Bjerkandera adusta* have been studied and been found to be as good as or better than *P. chrysosporium*. Notably some of the other white rots appear to be able to degrade a wider range of target compounds and to be more resistant to inhibition than is *Phanerochaete*. It may be possible to isolate other ligninolytic fungi (perhaps leaf litter degraders) that are better adapted than the white rots to the soil environment.

White rot fungi are perhaps better adapted to growth in aseptic conditions, without competition. This factor together with their relatively slow growth suggested that they will not be a useful option for treatment of large volumes of dilute effluents. However, they clearly have considerable potential for the pre-treatment of particularly difficult waste streams prior to a further treatment by conventional processes. Some industrial waste streams arise at such high temperature and extreme pH that they are essentially sterile and so the costs of separate sterilization may be avoided.

13 ANAEROBIC-AEROBIC TREATMENT

Many compounds are resistant or highly resistant to biodegradation under aerobic conditions, often because their substituents (e.g. nitro, chloro or diazo units) have a strong electron withdrawing effect making them less susceptible to electrophilic attack under oxidative conditions. Such compounds are often relatively easily metabolized by reductive attack. In some cases this anaerobic attack may only result in primary biodegradation but may produce a product that is readily degradable under aerobic conditions. There are plenty of examples of anaerobic treatment followed by aerobic treatment that can either lead to complete mineralization or to substantial degradation of a molecule which would otherwise be difficult or impossible to degrade. In some of the reported studies the microbial populations are very specific and highly selected while in others no selection is required. In most examples of this phenomenon the anaerobic and aerobic stages are carried out in different reactors or in the same reactor but sequentially. However, in one study a mixed population of bacteria was immobilized in gel beads. Reduction occurred in the anaerobic centre of the beads and oxidative metabolism in the aerobic outer zone. It is thus possible that anaerobic and aerobic reactions can occur at the same time in different regions.

In one promising process, nitrobenzene, which is very recalcitrant to aerobic degradation, is anaerobically reduced to aniline. This

gratuitous reduction is encouraged by the addition of glucose or a mixture of industrial solvents. The resultant aniline is easily degraded in a conventional activated sludge process.

With azo dyes, the ability to cleave the diazo bond reductively is very widely distributed and can probably be found in most activated sludges – the problem is that some, though by no means all, of the resulting aromatic amines are still difficult to degrade. In one study, the azo dye Mordant Yellow 3 is reduced anaerobically by a mixed culture of two pseudomonads to yield 6-aminonaphthalene-2-sulphonate and 5-aminosalicylate, with glucose added to promote reduction. When the culture is made aerobic, *Pseudomonas* BN6 partly degrades the naphthalene derivative, converting it to another molecule of 5-aminosalicylate, this is then completely degraded by *Pseudomonas* BN9. Thus the entire dye is mineralized. This is an encouraging observation. However, Mordant Yellow 3 is a simple dye, and most of the commercially used azo dyes are more complex and highly sulphonated giving rise to aryl amines that are more complex and difficult for bacteria to degrade. This technology may not be universally applicable and its utility has to be established on a case-by-case basis. However, for azo dyes that have at least one easily metabolizable moiety (e.g. aniline) anaerobic reduction may be useful and may lead eventually to a net reduction in effluent COD.

Sequential anaerobic/aerobic treatment has also been demonstrated to be of potential value in the treatment of chlorinated aliphatics, trichlorophenol and polychlorinated benzenes. It appears that for monocyclic chlorinated aromatics reductive dechlorination occurs more readily with more highly chlorinated compounds. These are reduced in a stepwise fashion but, generally, some mono- or dichlorinated compounds are left which are much more readily degraded by aerobic cultures than their polychlorinated precursors.

It has been suggested that for polychlorinated pesticides, like DDT, alternate anaerobic/aerobic degradation may be a means of achieving mineralization. This may be so, but DDT is rarely a problem in effluent treatment as it occurs mainly as a diffuse pollutant, rather

than at point sources. Flooding of soils which are heavily contaminated with polychlorinated compounds may produce anaerobic conditions which could allow reductive primary biodegradation yielding metabolites that can be aerobically degraded when the soil becomes aerobic again. It has been shown that DDT can be converted anaerobically by the common coliform bacterium *Aerobacter aerogenes* to a range of less chlorinated products including 4, 4'-dichloro benzophenone (DBP). DBP is much more amenable to aerobic attack than is DDT.

14 DEGRADATION OF RECALCITRANT ORGANICS IN EFFLUENT TREATMENT PLANTS

Many recalcitrant organic chemicals find their way into effluent treatment plants either from domestic use or from industry, principally from the chemical, petrochemical and textile industries. There is no doubt that it is much better for such chemicals to be degraded in a treatment plant than to enter the wider environment in which degradation is even less certain. In the UK it is still the case that most industrial liquid effluents are treated at plants which treat mixtures of domestic and industrial wastes – special treatment plants dedicated to particular industrial enterprises are a distinct rarity. Often consent conditions for discharge of effluent are very general (COD, BOD etc.) and usually do not include lists of particular chemicals. If specific chemicals are effectively to be removed from effluents then routine monitoring is essential to establish how efficient the treatment plant is and what increases or decreases its efficiency. Limits on COD are becoming more common and will probably soon be imposed on all effluents – if these are to be met it will be necessary to ensure that all potentially degradable materials are in fact degraded. This can be difficult as in many situations the composition of effluents can vary enormously and fluctuations can be irregular or on any time basis (hourly, weekly etc.). The factors that control the possibility of degradation have been spelt out in general

terms in Section 3, but some specific examples in the context of effluent treatment are now given.

14.1 Mean sludge retention time

Morpholine is a simple heterocyclic compound (Fig. 34.15) that has a reputation for recalcitrance and failure to obtain biodegradation of morpholine has been reported regularly.

It has now been clearly established that it is degradable but the only organisms that have been shown to degrade it in pure culture are all *Mycobacterium* or related species and all are intrinsically slow growing. Their growth rates are generally about a quarter to a tenth of those typical of *Pseudomonas* spp., which are often considered the 'classical' degradative bacteria. The slow growth rate of these morpholine degraders means that they can easily be lost from activated sludge treatment systems if sludge is wasted at too great a rate. Sludge wastage is described by the mean solids retention time (MSRT) or sludge age; this is a measurement of the mean length of time a particle of activated sludge is retained within the treatment system. It has been shown for morpholine that if MSRT is less than 8 days then morpholine degradation will be incomplete and if it less than 3 days there will be no morpholine removal at all. Similar findings have been made for some of the non-ionic surfactants.

It is possible to predict the MSRT needed to ensure degradation of a chemical, and indeed to predict the concentration of the chemical in the effluent of an activated sludge plant, if certain parameters, such as the specific growth rate, are known.

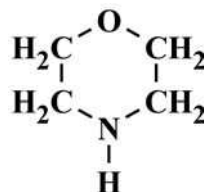


Fig. 34.15 The structure of morpholine.

S_1 is the concentration of the target compound in the treatment plant effluent and it can be obtained from the following equation:

$$S_1 = \frac{K_s(1 - K_d\theta_s)}{\theta_s(\mu_m - K_d) - 1}$$

K_s is the saturation constant for the target compound with the competent organisms; K_d is the decay constant for the microbial population; μ_m is the maximum specific growth rate for the competent microorganisms; θ_s is the mean sludge retention time.

It is notable that the only plant operating parameter in this equation is the MSRT. Both μ_m and K_s are constants only under specified conditions and may (will) change with environmental conditions (see temperature below). Similar relationships between MSRT and activated sludge plant performance have been demonstrated for ammonia oxidation, a similar process in that it relies on growth of a very specific group of slow-growing bacteria (the nitrifying bacteria). There is no doubt at all that the same will be true for the degradation of a great many xenobiotic recalcitrant chemicals. Many of these chemicals are degraded only by a limited range of slow-growing, specialized bacteria. Longer MSRTs may lead to improvement in removal of some of these chemicals, however, it should be noted that MSRTs can be too long and can lead to settling, and other problems.

14.2 Temperature

As detailed above, temperature is an important determinant in biodegradation reactions. In the context of effluent treatment, temperature will control the rates of metabolism and of microbial growth. It has been shown that at higher temperatures shorter MSRTs are required for degradation of ammonia and also for recalcitrant non-ionic surfactants. Low temperature is a real problem in domestic effluent treatment plants in the winter as the effluent is cold after passing through several miles of sewers and heating large volumes of dilute effluent is not economically viable. This can result in large variations in effluent quality over the course of a year, especially with

regards to materials degraded by slow-growing microbes. Dedicated industrial effluent treatment plants are less susceptible to low temperature problems than domestic plants, partly because the effluent is in relatively smaller volume, is more concentrated and is usually warmer. Furthermore, cheap sources of heat may be available to maintain higher temperatures in the treatment plant. Thus seasonal fluctuations in temperature are smaller and less important for industrial activated sludge plants.

14.3 Specialist degraders

Some chemicals can be degraded by a wide variety of different microbes and this is true of compounds like toluene, phenol and benzoate. Quite often, however, a compound is only degraded by a small group of specialist organisms. When this is the case the ability of an activated sludge to degrade the compound will be more precarious and more likely to be lost. The specialists, for example, may be susceptible to a toxic effluent component that may eliminate the whole population or repress its activity. A related example in which specialist populations are involved is that of nitrifying bacteria which, as a group, are susceptible to inhibition by a wide range of organic chemicals.

In the treatment of chemical industry effluents, one potential problem is that products are often made on a campaign basis. Consequently, a recalcitrant chemical may be present in the factory effluent for several months or weeks (or even less) and then absent for a prolonged period. When this happens the microbial ecology of the treatment plant may be affected. If a 'specialist' group of degradative microbes are the agents of degradation, these may suffer if their substrate is no longer present. This has been demonstrated for morpholine degradation. In a laboratory treatment plant the population of potential morpholine degraders declined markedly with the absence of morpholine in the influent. When the activated sludge was re-exposed to morpholine the lag period before its degradation greatly

increased – from nothing to about a day after 7 weeks, and about 2 days after 14 weeks, without morpholine in the feed.

This finding agrees with observations from an industrial activated sludge plant serving a chemical factory where it was noted that if morpholine had been absent from the factory effluent for a time, when it was reintroduced the activated sludge plant had lost the ability to degrade it. To overcome this, when morpholine was absent from the effluent it was added artificially to help maintain the sludge's acclimation to degrade it. The consequences of failure to degrade being that morpholine would pass through into the final effluent undegraded, giving an avoidable increase in COD. We know of another chemical factory that occasionally uses morpholine as a solvent and has developed a special microbiological pre-treatment process for the morpholine-containing waste stream. This has proved essential as, with only occasional presence of morpholine, it was not possible to develop and maintain a morpholine-degrading capability in the main treatment plant.

Specific pre-treatment of minor effluents generated during campaigns may be a useful approach, but is likely to be an expensive option. Another possibility is the addition, as and when required, of specially adapted microbial cultures or activated sludges to the main treatment plant. For this to be successful, storage of large volumes of cultures or sludge would be required; success cannot be guaranteed as it may take a considerable time for the added cultures to become established in the treatment plant. The artificial addition to the effluent of likely problem compounds to help maintain the biodegradative capabilities of the activated sludge between campaigns may be a more reliable and cheaper option, although it may go against the grain actually to add chemicals to an effluent!

14.4 Analysis and monitoring

It is clearly desirable for the recalcitrant organic compounds found in industrial effluents to be degraded in effluent treatment

plants. Indeed, if companies are to meet the ever-tightening restrictions it is essential. One factor that mitigates against the success of such treatment plants is the lack of detailed knowledge about specific aspects of their performance. This problem is particularly bad in treatment plants that principally treat domestic sewage with a variable portion of different industrial effluents. It is important that plant operators have available to them information not only on gross characteristics, like BOD, COD or TOC, but also on the concentrations of specific chemicals in the effluent. It should be pointed out that BOD is not a good indicator of the degree of removal of recalcitrant organics from effluents. It may give an idea of relatively short-term effects of an effluent on a water course but not of the extent to which any recalcitrant chemicals have been removed (often recalcitrant chemicals will not exhibit a BOD using the standard test).

At the end of the treatment process effluents will contain a proportion of 'hard COD' which is difficult to remove. If progress is to be made operators must have detailed knowledge of what this residual material comprises and the extent to which the microflora is able to degrade individual components of it. Such analytical data need to be available from regular time intervals (or composite samples) spread over a considerable period of time. Operators will then be able to look for trends in the degree of degradation of specific problem chemicals. These trends can then be correlated with overall plant performance, variations in operating practice (e.g. sludge wastage) and parameters (e.g. concentration of mixed liquor suspended solids and MSRT), changes in pH or oxygen levels, effluent composition (e.g. concentration of potentially toxic components) and environmental factors (e.g. temperature). Such correlations may allow operators to optimize removal of potential problem compounds. If problem compounds or waste streams are identified then details of their nature, and effects, can be determined and research undertaken to develop strategies for their more effective

treatment. In extreme cases this may involve special pre-treatments (biological or chemical) for particular waste streams.

Detailed chemical analysis has always been expensive but is becoming more and more important. Techniques such as GC/MS and even HPLC/MS are becoming cheaper and more readily available and offer the prospect of relatively easy identification and quantification of a wide range of chemicals without the employment of an army of analysts. Part of the price for the successful treatment of industrial effluents is constant analysis and monitoring of data.

15 MEASUREMENT OF BIODEGRADABILITY

There are many different methods of assessing the biodegradability of a chemical, but there have been efforts to standardize them by transnational bodies such as the Organization for Economic Cooperation and Development (OECD) or the European Community. Pollution, after all, is no respecter of international boundaries and it is important that official tests are consistent between different countries. There is no room to describe all the different tests and readers are directed to the early chapters of Karsa and Porter (1995) which give an excellent analysis of the different methods and their 'pros and cons'. When choosing a test for biodegradability it is important to ensure that the incubation time is sufficiently long. Many biodegradable compounds have in the past been described as completely recalcitrant because incubations for biodegradability tests were too short. To be effective incubations may need to be continued for anything from a few weeks to several months. This is particularly so when un-acclimated seed organisms are used. In the BOD test, use is often made of sewage effluent and, while this is adequate for many compounds, it will not usually be a good seed for recalcitrant chemicals. For example, it proved impossible to demonstrate a BOD for morpholine until a seed was used which had previously been acclimated for over a hundred days! In other studies the lag period prior to

observed degradation of morpholine was often between 20 and 60 days, shorter lag periods were only encountered when the inoculum came from a site likely to have been exposed previously.

It can be seen that in order to assess the inherent, or potential, biodegradability of a compound, considerable persistence may be required on behalf of the scientist! Preferably a range of tests should be performed which can give evidence of the type of biodegradation, as well as its rate; e.g. is the biodegradation merely primary or has mineralization occurred? Is the compound readily degradable or is it degraded only slowly and after a prolonged lag period? All this information will be of value in assessing the likely fate of a compound in an effluent treatment plant or in the wider environment.

16 BIOREMEDIATION

This term means making things better by biological means and is applied to processes used to solve problems of environmental pollution. The term bioremediation is used in a variety of contexts. It is applied, for example, to the 'cleaning-up' of land polluted by industry, attempts to remove pollutants from contaminated aquifers and the biological treatment of contaminated effluents. In our view this term is best reserved for processes that are used to solve past incidents of pollution rather than those used to treat new effluents. There are many problems of 'historic' pollution including old chemical factory sites that have been grossly polluted over decades due to ignorance of the effects of materials which were spilled onto them accidentally or deliberately. The process of coal gasification has also left a legacy of land polluted, for example, with phenols and pyridine derivatives and cyanides. Pollution of land and aquifers by spilling or disposal of solvents (such as the chlorinated aliphatics used for de-greasing and dry-cleaning) or jet fuel is also widespread and threatens water supplies in some areas. The military have also been responsible for contamination of land with explosives like TNT. Acute

problems of recent pollution are also regrettably common, with the spillage of crude oil at sea or on coastlines, due to the sinking of oil tankers, being the best-known example.

Bioremediation processes are generally split into two broad categories – *in situ* and *ex situ* processes. In the former, the pollution problem is tackled where it occurs and in the latter contaminated materials, be they soil or ground water, or the contaminants extracted from them, are removed and treated in a separate location.

Ex situ treatment has the advantage that it is possible to exercise much more control over the process with regard to environmental conditions such as pH, temperature and moisture. It is also possible to ensure more easily that aeration (if necessary) is adequate and that nutrients, if required, are available in the correct amounts. The disadvantage of *ex situ* processes is the expense involved in removing and transporting the contaminated materials to the treatment site. Once at the treatment site, a variety of approaches can be used including the composting of polluted soil, the treatment of the soil as a slurry (in lagoons or reactors) and the extraction of pollutants into water with treatment of the resulting solution by activated sludge. This list is not exhaustive. Contaminated ground water or water from aquifers can be pumped up and treated by a conventional biological treatment processes. Volatile compounds can also be removed from below ground by air stripping and subsequently removed from the effluent gases. Aerobic treatment of soil or water may lead to evaporation of volatiles. It is undesirable simply to volatilize the pollutants and transfer them to the atmosphere and so arrangements may need to be made to collect vapours and to remove volatile organics from them – possibly by biological treatment. Compounds like toluene and trichloroethylene can be treated in the vapour phase by a trickling filter-type apparatus. Care has to be taken not to pollute the surrounding soil and so the treatments have to be conducted on lined, impervious surfaces.

The *ex situ* approach may be impossible, undesirable or unnecessary. *In situ* processes are usually less intensive. They may involve the addition of suitable microorganisms where none exist, the addition of nutrients that are in short supply or the addition of air or oxygen (or an alternate electron acceptor, like nitrate). The simplest of such treatments are processes such as nutrient addition and ploughing of oil-contaminated soils (to improve aeration). More difficult processes involve the injection of oxygen or mineral nutrients into aquifers.

The addition of competent microbial cultures in bioremediation processes ('bioaugmentation') is somewhat controversial. In some situations it has been shown that such additions are unnecessary, for example, this appears to be so in the case of oil pollution. Several studies have shown that the addition of microbial cultures gives little or no benefit. The most effective approach for this type of pollution is the addition of mineral nutrients, notably nitrogen and phosphorus. The metabolism of such a concentrated carbon source as crude oil will rapidly deplete nutrient supplies and, in the case of soil pollution, supplies of oxygen as well. Nutrients have to be provided in an appropriate form and it has been found effective in treating oil spills to use oleophilic fertilizer preparations that will bind to beached oil and are not washed away by the large volumes of water. In bioremediation it is clearly important to identify the rate-limiting factor and address that in whatever way is feasible. Alterations in other factors may be of little benefit. Processes in which conditions for pre-existing organisms are improved by addition of nutrients/O₂ etc. are often referred to as 'biostimulation'.

Often the native microflora will already have the desired capabilities or will be capable of evolving to degrade the added pollutants. The addition of laboratory strains of microorganisms may be of little use. Such strains are often habituated to laboratory life and may have a very limited ability to survive in the more challenging and competitive environments of the soil or seawater. This is particularly true for genetically-engineered organisms which may be rendered uncompetitive in

many situations by the carriage of additional genetic information (which will impose a metabolic cost on the organism). In some situations in which the native microflora is very small or non-existent, e.g. in deep sub-soil or aquifers, the addition of competent microbes may be of use. However, it must be established that the organisms can cope with, and carry out the required metabolism under the prevailing conditions (which may include a shortage, or absence, of oxygen). It has been proposed that white rot fungi may be suitable for the bioremediation of contaminated land – their use in this context is discussed in Section 12.

For a more in-depth treatment of bioremediation readers are referred to the August 1993 edition (Volume 11) of the journal *Trends in Biotechnology*, most of which is devoted to this topic, or to the series of books *Remedial Treatment of Contaminated Land* (see reference list).

17 CONCLUSIONS

After the gloom of impending disaster and environmental calamity that was present in the 1960s, the threat from recalcitrant organic chemicals has diminished to some extent. Extensive microbiological and biochemical research has explained the reasons for recalcitrance and suggested routes by which problem compounds may be degraded. New processes of biodegradation (e.g. by anaerobes and white rot fungi) have come to light and methods of utilizing microbial activity have been developed or improved. An understanding of the effects of chemical structure on biodegradability means that we now have the knowledge to design chemicals that are more amenable to biodegradation in the environment, *if we wish to*. If progress is to be made it will be essential to gain an improved knowledge of the microbiology and biochemistry of the treatment of industrial effluents, and the biodegradation of their components, so that it can become more effective. If tighter environmental protection legislation is to be applied effectively and fairly then research

will be required to ensure that industry can comply with more stringent limits on the COD of effluents and direct toxicity assessments.

Environmental pollution is a transnational phenomenon and Western countries will not be able to escape its consequences by simply manufacturing products in developing countries where legislation may be less strict or less strictly enforced. Although DDT is now less of a problem in developed countries (in which its use has generally been banned) it has not entirely disappeared and is still in use in many poorer tropical countries. It is well known that DDT has been found in regions of the world in which it has never been used. This compound is still effective in malaria control, it is cheaper than, and has some advantages over, alternatives. Although it can be slowly degraded there is no *easy* biological treatment solution for it, especially as it arises as a diffuse pollutant and spreads. Clearly, if the threat of DDT is to be lifted completely then developing countries will need aid, either to buy more expensive but less persistent alternatives or to develop new insecticides. Currently there is a thriving illegal international trade in serious pollutant chemicals (like CFCs) which are banned in the West but are manufactured elsewhere. It is vital that new chemicals being developed are given more extensive testing to ensure their biodegradability before they are put into commercial use. Export of pollution is not morally acceptable.

As we enter the 21st century, further scientific investigation is essential, but its beneficial effects will be nullified without a transnational legal framework to outlaw serious pollutant chemicals and generous financial aid to make it effective across the globe.

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Heavy metals in wastewater treatment processes

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1 INTRODUCTION

The term heavy metals is used to denote a large group of elements. Widely accepted definitions are those of Phipps (1981) and Weast (1984) defining the heavy metals as those elements with a density greater than 6 g/cm^3 and 5 g/cm^3 respectively. From an environmental standpoint, the definition of a heavy metal is fairly broad and includes metalloid substances which exhibit similar properties in the environment; typically high toxicity at low concentrations and with a long residence time in soil. From this standpoint the definition of Weast is the closest.

Some of the heavy metals are micronutrients and are required in trace amounts by living organisms for their normal metabolic function. At elevated concentrations in the environment they exhibit a toxic effect. Elements such as cadmium and mercury have no known metabolic function and are considered to exhibit some toxicity at all concentrations. The toxicity of the heavy metals depends upon the chemical form in which they are present (Rochow, 1964). Organometallic complexes may well show the phenomenon of bioconcentration due to the greater solubility, and hence preferential uptake, of the substance in lipid rather than water.

Besides directly causing non-infectious diseases in man, heavy metals may also have a significant impact when released into the environment. They are a serious and persistent pollutant of terrestrial and aquatic ecosystems

(Bourg, 1995). Unlike various refractory organic compounds there is no possibility of destruction of these elements in the biosphere. Contamination of soil by heavy metals is a long-term problem. Estimation of the half-lives of some of the elements in soil range from 15 to 1100 years for cadmium, 310 to 1500 years for copper and 740 to 5900 years for lead, the wide range of values being due to differing soil conditions (Alloway, 1995).

Heavy metals are present as trace contaminants in most materials; more concentrated sources of these elements are found in fossil fuels and mineral ores. Anthropogenic sources have greatly accelerated biogeochemical cycles of heavy metals (Lombi *et al.*, 1998). Atmospheric release may generally disperse contaminants, such as lead from the, now banned, use of tetraethyl lead as an 'anti-knocking' agent in petrol or pollutants may be more concentrated on a local level, e.g. cadmium from pre-industrial mining and smelting operations. Pollution from heavy metals can generally be traced back to a point source.

Many industrial processes are reliant on the use of heavy metals as process reagents, or they are present as contaminants in their raw materials. Elimination of compounds from many sources of waste is often either impossible or needlessly expensive.

For the operator of a wastewater treatment plant the problems of heavy metals in the wastewater are twofold; their presence may significantly affect the efficiency of the plant, and discharges containing metals may have

a serious effect on the environment or man. This is reflected in the legislation concerning such emissions.

2 LEGISLATION

Global warming and the increasing world population will place an increased stress on the agricultural and water supply systems. Industrial activities pollute these resources and reduce the future capacity. Concern over the impact of the activities of man on the natural environment has led to international agreements limiting resource usage and environmental pollution. With these concerns in mind legislation is now being implemented which not only requires an installation to meet specific discharge criteria, but also to consider to Best Environmental Practice (BEP), Best Available Technology (BAT) and Integrated Pollution Prevention and Control (IPPC). In order to enforce these regulations enforcing bodies require some form of criteria and indicators to measure environmental damage and, to conform to such regulations, the operators of polluting or potentially polluting installations will most probably have to implement some form of environmental management programme. The heavy metals of concern, as have been identified in European Union and US legislation, are those which are present in sufficient quantity in the earth's crust and are of sufficient solubility in water to be determined an environmental hazard. European Union Directives concerning these dangerous substances list the pollutants as being of concern in the aquatic environment.

The use of heavy metals in industry is governed by health and safety, transportation and storage legislation. These will also cover operators of wastewater treatment facilities. They may impose exposure limits to gaseous concentrations, requirements for personal protection equipment and storage, handling and disposal procedures.

The legislation covering discharges of heavy metals into the environment can be split into three broad categories:

1. Discharges to water
2. Discharges to land
3. Release to the atmosphere.

Note that heavy metals in ground water may be considered analogous to contaminated land where there is not a threat to a potable water supply.

2.1 Discharges to water

Limitations on the quantity of heavy metals discharged into water are required to protect potable water supplies, fisheries and, more recently, the aquatic environment. Environmental concerns over the marine environment, once seen as a suitable dumping ground for radionuclides and raw sewerage have seen this route of disposal curtailed.

World Health Organization recommendations on the maximum acceptable concentration of various priority pollutants have been accepted internationally and provide the basis of European Union and US legislation on the protection of potable water supplies. These are designed to ensure that drinking water is fit for human consumption (see Table 35.1 for metal concentration limits on discharge and land loading rates).

Municipal sewerage works have to protect the receiving medium from heavy metals in the aqueous discharge. This has traditionally been achieved by the dilution of metal-bearing industrial waste with less contaminated wastewater from domestic sources. Biological treatment systems as well as their primary role of reducing the BOD of the effluent, also have a great affinity for heavy metals and provide a convenient method of removal. The solid wastes generated from these treatment processes must also be disposed of. Recycling of these wastes is seen as advantageous from an environmental standpoint, but the metal content may preclude this option. Disposing of metals to landfill and incineration are the other two major options available, but are set to become increasingly more expensive. When constructing a wastewater treatment facility, consideration should be given to the disposal of the solid waste generated. The European Union

TABLE 35.1 Limitations on discharges into the environment of heavy metals

<i>Heavy metal</i>	<i>British EQS $\mu\text{g l}^{-1}$</i>	<i>EU limit values for the concentration of metals in sludge $\text{mg kg}^{-1}\text{dm}$</i>	<i>EU annual loading rates for agricultural land receiving MSW sludge^a $\text{g ha}^{-1}\text{y}^{-1}$</i>	<i>US part 501 Ceiling concentration of metals in sludge mg kg^{-1}</i>	<i>Annual pollutant loading rates $\text{kg ha}^{-1}\text{y}^{-1}$</i>
Arsenic	50			75	2.0
Cadmium	5	20–40	150	85	1.9
Chromium	20				
Copper	10	1000–1750	12 000	4300	75
Iron	1000				
Lead	10	750–1200	15 000	840	15
Manganese					
Mercury (total)	1	16–25	30	57	0.85
Molybdenum				75	
Nickel	150	300–400	3000	420	21
Selenium				100	5.0
Vanadium	20				
Zinc	75	2500–4000	30 000	7500	140

Based on a 10 year average.

^a MSW Municipal Sewage Works.

Directives give an indication of future legislative requirements on national governments.

2.2 Discharges to land

The European Union Landfill Directive (EU (1999), implemented in the UK in July 2001, will further increase the pressure to recycle biological waste to land with a commitment to reduce, in stages, the landfill of biological waste to 25% of its current level by 2010.

The application of waste and industrial activated sludge to agricultural land as a soil conditioner and to contaminated land, is regulated under the Waste Management Licensing Regulations 1994 (HMG, 1994). Under these rules the spreading of various wastes, including those from industrial and biological treatment plant origins, is exempt from waste management licensing, provided that agricultural or ecological improvement can be demonstrated. This waste application is limited to a maximum annual application of 250 tonnes/hectare.

The Environment Agency of England and Wales (EA) and the Scottish Environmental Protection Agency (SEPA) have both issued reports expressing concern over the spreading of industrial waste to land (Anon, 1998a,b). In

these reports they have recommended the tightening of regulation to ensure that industrial waste which is spread to land is of a similar quality to that of municipal sewage waste (MSW). The application of potentially toxic elements from MSW to agricultural land is well regulated. The Department of Environment, Transport and Regions (DETR) has recently reviewed the scientific evidence relating to the agricultural use of sewerage sludge, including the scientific basis for the limits of the potentially toxic elements (PTEs) Cd, Cu, Zn, Pb, Hg, Cr, As, Se, Mo, and Fe (DETR, 1998). Application of excess activated sludge from industrial waste treatment that contains PTEs above these limits is unlikely to be acceptable.

The textile industry, despite recent reports in the press, is still a major manufacturing industry in the UK and will continue to be so for the foreseeable future (Barker, 2000). In October 1999 the European Commissions Environment Directorate produced a draft working paper outlining plans to revise the 1986 EC Sludge use in agriculture directive (EC, 2000; EEC, 1986). Included in the plans is a proposal to reduce the limit values for metals in sludge in 2005 and 2010, which may cause 21% and 72% failure of excess

municipal sewerage works (MSW) return activated sludge (RAS) to meet the limits (Anon, 1999). RAS is the suspended solid fraction found in the bottom of a secondary clarifier, composed mostly of microorganisms, which is returned to the aeration treatment pond and mixed with influent waste.

Certain sections of the textile industry, wet processing in particular, engage in practices that may have a significant environmental impact. The wet processing sector of the textile industry is among the top four or five major polluters, together with pharmaceuticals, the paper industry and smokeless fuel production in the UK (Pierce, 1994). Of these, approximately 10% have their own effluent treatment plants. The majority of the remaining effluent is combined with other domestic and industrial sources for treatment at a municipal sewerage works.

Solid waste from water treatment plants must be disposed of or recycled. Three practical routes exist: incineration, applying to land as a fertilizer or soil conditioner, and dumping in landfill. Incineration leaves ash, which requires disposal either by one of the two above methods or its incorporation into building materials.

Biosolids are rich in phosphorus and, as such, are suitable for application to land. Authorities in Australia, for example, limit use to land especially when applied to acidic agricultural land. Bioleaching occurs, transporting the heavy metals to other areas. The metals end up bioleaching in this manner as a result of bioacidification. Heavy metals can therefore, be removed from aqueous effluent wastewaters by bioacidification prior to disposal of biosolids to land. Unfortunately, the nutrient concentration is also reduced significantly in the process (Shanableh and Ginige, 1999).

2.3 Discharges to air

Within the workplace limit values are applied to various hazardous chemicals establishing short- and long-term exposure concentrations, which provide no or an acceptable health risk.

Discharges to atmosphere are also regulated, particularly important in industrialized urban areas.

2.4 Concerns for the future

In the EU, discharges of dangerous substances to water, as listed in EC directives 76/-464EEC (Dangerous Substances Directive) and 86/440/EEC, will be banned.

2.4.1 *Legislative concerns for the future*

Climate change due to global warming is now widely accepted internationally by most industrialized nations. The goal of sustainable development arising and the further transcription of the precautionary and polluter pays principles into law will necessitate more binding environmental legislation than is currently in force. In the European Union, future requirements upon industry can be found in directives which have not yet been transcribed into national law. When considering the disposal of metal-containing sludge an EU policy document advocates the introduction of much more restrictive practices.

2.4.2 *The EU water framework directive*

The Directive 2000/60/EC will integrate existing water quality legislation (EU, 2000a). An obligation is placed on member states to ensure that surface and ground waters reach a good condition by 2010. Polluters will be required to pay the full cost of the damage they cause, rather than leave it to society to bear the cost. This may have important cost implications for industries discharging heavy-metal-bearing industrial wastes to sewerage treatment works.

Member states will be required to establish river basin management plans, overseen by a competent authority. Previous water management regulations will be integrated into the water framework directive. The discharge limits of heavy metals will then be set by environmental quality standards (EQSs) specific to the river basin management plan. The EQSs will be set at a level intended to protect the biota, both terrestrial and aquatic, of the river basin. The EQSs will also have to ensure that naturally occurring dangerous

substances will not be present in the marine environment at levels significantly above the background concentration.

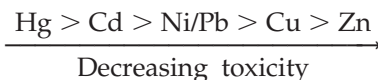
2.4.3 Proposed sludge application to land revisions

The third EU draft working document on sludge contains proposals that will affect most treatment works that recycle excess sludge to land as a fertilizer. A significant proposal is that all operations producing sludge are to be brought under similar regulations to those currently regulating sewerage treatment works. Current limit values on the acceptable sludge metal content, soil metal loading rates and the total acceptable metal loading on the soil will be lowered. Reductions are proposed in the short, medium (≈ 2015) and long term (≈ 2025). If these medium-term values are applied, as suggested, the new limit values would preclude the use of many MSW sludges currently recycled to land. The new tighter threshold limit values for the amount of allowable heavy metal in receiving soils will reduce the available arable land.

Under the proposal, it will not be possible to reduce the concentration of heavy metals in sludge for application to land by combining it with another sludge source. The producers of sludge will also be under obligations to ensure that the sludge they generate is suitable for land application, even if the distribution and application of the sludge is performed by a third party. The producer will also be required to implement a quality system over the whole process, from reduction of pollutants into the waste treatment, to the supply of relevant documentation to the delivery site. The producer of sludge for arable land application, will be responsible for testing for the presence of toxic elements.

There are several outstanding features which may be observed from Table 35.1, the most significant being the relatively high loading rates of these metals. Generally this would be worthy of considerable discussion. However, it is not considered to be appropriate in this text. It is apparent that the relative toxicity of metals common to the

UK, EU and the USA are of the following order:



From Table 35.1 there is general agreement that Hg and Cd are the most highly unacceptable pollutants of the metals considered.

3 CHEMISTRY OF HEAVY METALS IN WATER

Heavy metals consist of a large group of elements. They may be present in water in a variety of states including free metal ions, chelated compounds, organometallic complexes and elemental metal. These substances may be dissolved in the water, associated with colloidal compounds or present in association with particulate matter. The oxidation state of the metal ion may also be of importance; chromium exists in water in two oxidation states. The Cr^{3+} is relatively non-toxic and is required as a trace element by animals. The Cr^{6+} ion, however, exhibits carcinogenic properties. The majority of the heavy metals are cations in solution. At acidic pH they tend to exist as free metal ions. Around neutral pH 6–9 some precipitate as hydroxides or other insoluble species if the appropriate co-ion is available.

4 DETECTION OF HEAVY METALS

The current method of choice for quantitative heavy metal detection is inductively coupled plasma mass spectroscopy (ICPMS), which can simultaneously detect multiple elements, in some cases down to low parts per trillion concentrations for many metals. The equipment required is expensive to purchase and operate. Electrochemical methods of detection, including polarography and anodic stripping voltimetry, are capable of detecting multiple elements. They can also yield valuable information about the availability of metal ions, and are most suitable for research purposes. Where the element of interest is known, then the use of traditional chemical reagents for metal

detection provides a useful tool. These reagents are available as proprietary kits, which may be used onsite.

5 SOURCES OF HEAVY METALS IN WASTEWATER

Industrial waste contains pollutants that may preclude the use of excess activated sludge, derived from industrial waste treatment, as an agricultural fertilizer. Recent legislative changes have considerably reduced the options available for the disposal of solid waste from biological treatment plants (McCann, 1998). Consequently, it is now more economically favourable to recycle organic wastes to land. Heavy metals may gain entry into waste streams from a variety of sources. The presence of metals in some dyestuffs is well known, but these contaminants may also be present on the raw materials and as a contaminant of other process auxiliary chemicals, where it has been involved in the compound's synthesis (Charlton, 1999). The effluent from cotton wet processing typically has a highly alkaline pH. This must be lowered to a pH value generally between 6 and 8 before the effluent is processed by aerobic biological treatment.

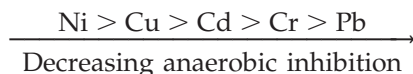
Heavy metals are present in wastewater from industrial and domestic sources. Concentrated highly contaminated wastes can be found in wastes from metal processing industries such as electroplating. Chemical works, textile wet processing, tanneries, photographic industries and mining may also produce wastes with a significant metal content.

Sources of heavy metals in domestic waste may come from metal piping (lead and copper), galvanic corrosion (zinc), cosmetics and household cleaning agents. Surface water runoff from roads and fire water may also contribute to the metal load on a wastewater treatment plant when water from these sources enters the sewer.

6 THE EFFECT OF HEAVY METALS ON BIOLOGICAL TREATMENT PROCESSES

Cu^{2+} , Pb^{2+} , Cd^{2+} , Ni^{2+} , Zn^{2+} and Cr^{6+} are known to inhibit anaerobic digestion (Lin, 1992; Mueller and Steiner, 1992). The

relative degree of anaerobic inhibition in a municipal sludge has been found to be:



The toxicity of the metal is directly related to its solubility in the presence of the sludge. If the affinity of the metal for the sludge is high then the toxicity is reduced (Mueller and Steiner, 1992). Metal toxicity in such systems may be reduced by precipitating the metal out of solution as the insoluble sulphide or hydroxide, etc. Metals such as Ni, Co and Mo, however, are known to promote anaerobic digestion (Shonheit *et al.*, 1979; Whitman and Wolfe, 1980; Murray and van den Berg, 1981; Bitton, 1994).

Toxicity is expressed as 'inhibition of respiratory activity of microorganisms' present in the activated sludge'. Cr^{3+} and Cr^{6+} were used in a study in order to determine their effect on the activated sludge. Oxygen concentration and biological oxygen demand were measured as an indication of inhibition in the presence of the metal. Exposure times and activated sludge biosolids concentration was examined and their effect on EC_{50} recorded. The 1-hour EC_{50} for Cr^{6+} was found to be 40–90 mg/l. This effect was not observed for Cr^{3+} alone, only the combined effect of the two ions (Vanková *et al.*, 1999).

In another study, the three metal ions Cu^{2+} , Zn^{2+} and Ni^{2+} were found to inhibit β -glucosidase enzyme activity. The pH and buffer type were investigated and were found to have considerable effect on the enzyme activity. Under optimal conditions, a metal concentration of 0.6 mM and a pH of 5, the presence of Zn and Ni decreased the enzyme activity by 25 to 30%, whereas Cu reduced it by more than 90%. The citrate buffer was not inhibited at all, even at the higher Cu concentrations. The inhibition by the Zn and Ni was found to be less pH dependent – range 4 to 5.5. Chemical speciation models were used to describe inhibition by buffer and pH for all three metals. Here it was assumed that the enzyme activity depended on protonation of the amino acid at the reactive site complexation by the heavy metal cation (Geiger *et al.*, 1999).

In natural unpolluted environments, heavy metals are present in trace concentrations. The most abundant may have a metabolic function. The inability of most living organisms to cope with the toxic effects of heavy metals is due to lack of evolutionary selection pressures to develop these mechanisms.

Some fungi have shown toxic effects to heavy metals. An example here is the thermochemical study of some heavy metals. Thermogenetic growth curves of *Rhizopus nigricans* have shown how a variety of metals have inhibited growth rate constants and inhibitory ratios, although low Cu concentrations promote these properties. The method of analysis, microcalorimetric analysis bioassay, for acute cellular toxicity is based on metabolic heat evolution from cultured cells (Cheng Nong *et al.*, 1999).

Here the effect of heavy metals in activated sludge was examined when used as a food supplement in Wistar rats. The toxic effect of heavy metals as contaminants from sludge-supplemented diets on male Wistar rats has been studied (Bag *et al.*, 1999). Poultry and cattle feed was supplemented with activated sludge as it is well known that it is high in nitrogenous matter and is inexpensive. The toxic effect of using domestic sewage sludge, which is contaminated with heavy metals, pesticides etc. on male Wistar rats was undertaken, and the metal content of the dried sludge before mixing with the cattle feed was determined using atomic absorption. The toxicity was determined by analysis of enzyme activity in serum, liver, muscle and the brain of the Wistar rats. It was concluded that the mixing with animal feeds should be implemented with caution because of the growing awareness that biomagnification of heavy metals may result in the food chain. This begs the question that, if awareness of the biomagnification effect was not growing, would we implement without caution?

6.1 Effect of heavy metals on treatment efficiency

Heavy metals have a deleterious effect on living organisms if present in concentrations above that which naturally occurs in the organisms'

environment. In general, heavy metals tend to have a bacteriostatic effect. Increasing concentrations lead to mortality. Some microorganisms have developed mechanisms to cope with elevated concentrations of heavy metals in their environment.

A recent study has shown that, at concentrations well below that where toxic effects are observed, and not uncommon in most combined industrial and domestic wastewater treatment works, the presence of heavy metals can considerably affect the efficiency of the plant, reducing the chemical oxygen demand adsorption capacity (CAC). The presence of heavy metals may also affect the settling characteristic of sludge. This may be due to a direct effect upon the extracellular biopolymer or to the relative promotion of filamentous bacteria. Subsequent anaerobic digestion of excess metal contaminated sludge flocs from activated sludge treatment may be inhibited or prevented by the presence of heavy metals. Anaerobic treatment processes are, generally, most sensitive to contamination by heavy metals and the material entering a sludge digester may contain many times the concentrations of heavy metals compared to the influent plant waste stream.

Trace levels of Cu, Cr and Zn have been investigated on the performance of activated sludge. Adsorption onto batch flocs followed both Langmuir and Freundlich isotherms. The presence of the three metals affected organic matter adsorption and the CAC. The metal ions reduced CAC by competition for adsorption sites on bioflocs in a sequence batch reactor (SBR) depending on hydraulic retention time (HRT). Heavy metals were adsorbed quicker in shorter HRTs (2 days) as opposed to longer HRTs (5 days). Again, indicative of the greater mobility of the single metal ion despite its high atomic mass, although this being quite small compared with the organic competitors (Sin *et al.*, 2000).

There is an excellent review on microbial heavy metal resistance, which discusses metabolism of heavy metals to a degree of detail (Nies, 1999). Even though heavy metals form a major part of the elements in the periodic table, as mentioned above, a lot of

them are essential as trace elemental nutrients, although, at high enough concentrations most are toxic. Transport of the most important heavy metals is compared after discussion of the principles of homeostasis for all heavy metal ions.

The uptake and control of heavy metals across cell membranes has been investigated and studies have shown that heavy metals are transported across the outer microbial cell membrane via two types of transport system:

1. Fast unspecific uptake driven by a chemostatic gradient across a cell membrane. These are constitutionally expressed mechanisms of which two general types have been recognized. Divalent metal cations enter through one type of channel that is present to allow the ingress of micronutrient metal cations, e.g. Zn^{2+} . Oxyanions, e.g. chromate, gain entry through unspecific uptake systems allowing the passage of phosphate and sulphate ions.
2. Systems showing high substrate specificity. This may be utilized against a concentration gradient, sometimes use an ATP source as an energy source; they are inducible.

The toxic mode of action of heavy metals is generally within the cell itself. This may be by the substitution of the metal ion for a structural homologue micronutrient or binding to material within the cell, especially heavy metals with a high atomic mass showing a great affinity for thiol constituent groups on amino acids, such as cysteine. Both mechanisms can cause considerable oxidative stress on the cell.

7 NATURAL MICROBIAL RESISTANCE OR DETOXIFICATION MECHANISMS

Three general mechanisms are available to the microbial cell:

1. Reduction to a less toxic form
2. Effusion from the cell generally requiring a specific transport mechanism working

against a concentration gradient coupled to an energy releasing reduction

3. Complexing the heavy metal to form an inert relatively non-reactive species.

Microbial cells generally adopt two or all three of the above mechanisms.

The rate of oxygen uptake inhibition with PACT sludge has been studied in the presence of a variety of inhibitors (Sher, 2000). However, in this particular study, Zn was the only metal used and its inhibitory effect was compared with those of phenol and one of its derivatives. The sludges were unacclimated and their ages were 4 to 12 days old. For the poorly adsorbed Zn, the IC_{50} defined values and sorption did not show any change. However, the 12-day-old sludge showed much more resistance to Zn exposure. Changes in biomass were the only apparent reason for this behaviour. PACT was mixed with the activated sludge and is an activated carbon adsorbent.

7.1 Metabolic activity and metal biotransformations

The metabolic activity of microorganisms can result in solubilization, precipitation, chelation, biomethylation or volatilization of heavy metals (Iverson and Brinckman, 1978). Microbial activity may result in the following:

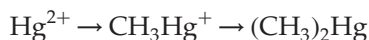
- Strong acid production, such as sulphuric acid by chemoautotrophic bacteria, e.g. *Thiobacillus*, which dissolves minerals
- Weak organic acid production, such as citric acid, which dissolves and chelates metals forming organometallic compounds
- Ammonia production or organic bases, which precipitate heavy metals as the insoluble hydroxides
- Hydrogen sulphide production by sulphate-reducing bacteria, which precipitates heavy metals as insoluble sulphides
- Extracellular polysaccharide production, which can chelate heavy metals and reduce toxicity (Bitton and Freihoffer, 1978)
- Certain bacteria (sheathed filamentous) fix Fe and Mn on their surface in the form of hydroxides or other insoluble metal salts

- Biotransformation of certain bacteria that have the ability biomethylate or volatilize heavy metals.

7.1.1 Metal biotransformation

Heavy metal sources in wastewater treatment plants include mainly industrial discharges and urban storm water runoff. Biological treatment processes such as activated sludge, biological filtration and oxidation ponds remove 24% (e.g. Cd) to 82% (e.g. Cu, Cr) of metals (Hannah *et al.*, 1986). Toxic metals may adversely affect biological treatment processes as well as the quality of receiving waters. They are inhibitory to both aerobic and anaerobic processes in wastewater treatment (Barth *et al.*, 1965).

Biomethylation of Hg was first reported in 1969. Since then, many bacterial, e.g. *Pseudomonas fluorescens*, *E. coli*, *Clostridium* spp., and fungal, e.g. *Aspergillus niger*, *Saccharomyces cerevisiae*, isolates have been shown to methylate Hg^{2+} to methyl mercury. Vitamin B₁₂ stimulates the production of dimethyl mercury $(\text{CH}_3)_2\text{Hg}$ by microorganisms:



Methyl mercury accumulates under anaerobic conditions and is degraded under aerobic conditions. Sulphate-reducing bacteria are also capable of biomethylating mercury (Compeau and Bartha, 1985). Molybdate, an inhibitor of sulphate reducers, decreases mercury methylation in anoxic environments by 95%. Microorganisms can also transform Hg^{2+} or organic mercury compounds, such as $(\text{CH}_3)_2\text{Hg}$, phenylmercuric acetate and ethylmercuric phosphate to the volatile metal Hg^0 . Mercury volatilization is a detoxification mechanism, the genetic control of which has been elucidated. Plasmid genes code resistance to metals such as Hg or Cd. Resistance to Hg is carried out by the *mer* operon, which consists of a series of genes (*merA*, *merC*, *merD*, *merR*, *merT*). The *merA* gene is responsible for the production of mercuric reductase enzyme, which transforms Hg^{2+} to Hg^0 (Silver and Misra, 1988).

Cadmium, Cd^{2+} , can be accumulated by bacteria, e.g. *E. coli*, *B. cereus* and fungi,

e.g. *Aspergillus niger*. This metal can also be volatilized in the presence of vitamin B₁₂. Like Hg, Pb can be methylated by bacteria such as *Pseudomonas*, *Alcaligenes*, *Flavobacterium*, to $(\text{CH}_3)_4\text{Pb}$. Fungi, e.g. *Aspergillus*, *Fusarium*, are capable of transforming As to $(\text{CH}_3)_3\text{As}$, a volatile form with a garlic-like odour. Selenium is another metal which is methylated through the metabolic activity of bacteria, e.g. *Aeromonas*, *Flavobacterium* and fungi such as *Penicillium*, *Aspergillus*.

Cr-rich wastewaters are generated by industrial processes such as leather tanning, metal plating and cleaning. An *Enterobacter cloacae* strain, isolated from municipal wastewater, was able to reduce Cr^{6+} to Cr^{3+} , which precipitates as metal hydroxide at pH 7, thus reducing the bioavailability and toxicity of this metal (Ohtake and Hardoyo, 1992).

8 HEAVY METALS REMOVAL

The methods of metal removal are numerous and selection of the appropriate techniques depend upon factors such as the particular metal(s) involved, the chemical composition of the wastestream, the volume requiring treatment, variability in composition and flow of the wastestream and level of treatment required.

Both physical and chemical processes are employed for the removal of heavy metals. Such processes include ion exchange, oxidation/reduction, precipitation, ultrafiltration and many others. During primary treatment much of the particulate-associated metal is removed by sedimentation.

Microorganisms offer an alternative to physical/chemical methods for metal removal and recovery (Forster and Wase, 1987). Metal removal or immobilization is carried out by the biomass and depends on the biomass solids concentration and sludge age. The older the sludge the higher the biomass solids concentration and therefore the greater the metal removal.

Cation exchanger textiles (CET) have been used for the removal of heavy metals from industrial wastewaters in batch reactors.

The CETs carry carboxylate, sulphonate and phosphate functional groups. The influence of counter-ion and functional group, metal concentration and the presence of different competing ions in solution on the exchange capacity and selectivity coefficients was studied for Cd^{2+} and Cu^{2+} . Comparisons of optimum operating conditions for the CETs in this particular study with conventional ion exchange resins were carried out. Regeneration of the CETs used enables them to be an attractive alternative for wastewater treatment (Lacour *et al.*, 2001).

9 METAL REMOVAL BY WASTEWATER MICROORGANISMS

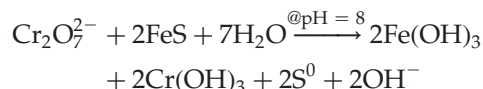
The extracellular polymer produced by microorganisms such as *Zooglea ramigera* and *Bacillus licheniformis*, for example, is known to have great affinity for a range of metals, e.g. Fe, Cu, Cd, Ni and U. The metals form complexes with the extracellular polymer and can readily accumulate. The bioaccumulation behaviour and toxicity of Cd in a protozoan community has also been studied (Fernandes-Leborans and Herrero, 1999). They found that Cd was accumulated by protozoa and bacteria at different Cd concentrations. The protozoa accumulating approximately 15 to 70 times more than the bacteria. The presence of Cd caused the abundance of protozoa to decrease down to a level at which recovery of the community was then suspected to be unrealistic. The abundance of the bacteria was not affected. The metals above, however, can easily be removed by treating with hydrochloric acid. Citric acid was used as a more sustainable acid, because of its inherent biodegradability to remove Cu and Zn and competing metals Ca and Fe from activated sludge prior to composting and use as fertilizer (Veeken and Hamelers, 1999). The metals were increasingly removed at higher temperatures and citric acid concentrations. At pH 3–4 the Cu removal was up to 60–70%, whereas with the Zn as much as 90–100% removal was achieved. The microorganism, *Zooglea ramigera*, was capable of removing Cu by 0.17 g/g

biomass (Norberg and Persson, 1984; Norberg and Rydin, 1984).

The same bacterium when immobilized in alginate beads can accumulate as much as 250 mg/l Cd from solution. The alginate beads themselves actually remove some of the Cd (Kuhn and Pfister, 1990). Heavy metals have been removed from anaerobically digested sewage sludge by isolated indigenous iron-oxidizing bacteria (Xiang *et al.*, 2000). The bacteria were used in an anaerobic batch reactor for the removal of Cr, Cu, Zn, Ni and Pb. Solubilization of the metal ions was promoted by inoculation with the iron-oxidizing bacteria and FeSO_4 . It was confirmed that the technique was effective, although the effectiveness varied between the different metals, e.g. 92% removal for Cu and only 16% for Pb.

RAS has a high metal affinity and may contain hundreds of times the concentration of heavy metals of the initial waste from which it was derived (Tyagi *et al.*, 1993). The secreted products of microbial anabolism (bacterial extracellular polymers) play the major significant role in the uptake of heavy metals from the aqueous medium (Nelson *et al.*, 1981). Binding sites in the secreted anionic and neutral biopolymers may combine with the cationic metal ions in solution (Brown and Lester, 1982). Studies into the removal of heavy metals from activated sludge solids have concentrated on the release of these metal ions from the sludge matrix by lowering the pH of the sludge with mineral and organic acids, typically below pH 4, the metal cations being in competition with H^+ ions (Sreekrishnan *et al.*, 1993; Du *et al.*, 1994). Hence lowering the pH lowers the affinity of the sludge for the metal ions and these are released into solution. Above neutral pH the dominant insoluble form of many heavy metals becomes the metal hydroxide. These are only sparingly soluble in alkaline conditions and lead to the precipitation of the heavy metal hydroxide for aqueous concentrations above their limit of solubility (Apak *et al.*, 1999). The pH of the medium has a significant effect on the solubilities of the metal precipitates. Most metals may also be

precipitated as sulphides as shown in the proprietary Sulfex process (Feigenbaum, 1978):



some as carbonates, phosphates or elemental (S^0) metal precipitation with sodium borohydride as the reducing agent. The use of sodium borohydride as a reagent is limited by its expense. Sulphide precipitation has been used as an effective alternative to hydroxide precipitation. It has the advantage of producing a precipitate even in conditions where the presence of a chelating agent would prevent the use of metal hydroxide precipitation. A serious disadvantage to the precipitation of metal sulphides is the potential for production of toxic hydrogen sulphide as a by-product. In activated sludge the potential number of ligand sites is often in excess of the metal ions present (Nelson *et al.*, 1981). The presence of free soluble heavy metal ions is therefore low except in acidic conditions. However, metals may still be present in the soluble phase attached to soluble ligands. By raising the pH of the activated sludge to favour the formation of metal hydroxides, or other insoluble precipitates, over metal ligand complexes, crystals of metal hydroxide should form within the sludge matrix which proves more amenable to physical separation techniques.

Studies have shown that inactivated microbial biomass in water treatment works takes up heavy metals from solution in a similar manner and concentration to live organisms. The main method of concentration of the heavy metals on to biosolids is by cation exchange with acidic functional groups on the excreted extracellular biopolymer surrounding the bacteria within a sludge floc.

Siderophores are low molecular mass molecules that are able to chelate with heavy metals and allow transport into the microorganism cells. Some microorganisms produce these especially to promote Fe^{3+} uptake from solution, the ferric ion being only sparingly soluble (Lundgren and Dean, 1979). Immobilized microorganisms (bacteria, fungi and

algae), e.g. Algasorb – algae cells embedded in silica gel polymer – is able to remove heavy metals including uranium (Anon, 1991). Fungal mycelia such as *Aspergillus* and *Penicillium* have also been used for metal removal from wastewater (Galun *et al.*, 1982) and are therefore a suitable alternative for detoxifying effluents. Metals are adsorbed from solution by fungi rather than by intracellular uptake, the latter process being much slower and energy dependent. Biosorption column studies have shown that immobilized *Aspergillus oryzae* remove Cd efficiently from solution. It has also been shown that detergent treatment of fungal biomass considerably improves metal removal (Ross and Townsley, 1986). Cd seems to adsorb onto the fungal biomass but active uptake of the metal by the fungus does not occur to any significant extent and is therefore probably not a metabolic process (Kiff and Little, 1986). Artificial wastewater has been used for the removal of the metals Cd^{2+} , Pb^{2+} and Cu^{2+} by biosorption onto dry fungal biomass. The metals were in the 5–500 mg/l concentration range. Biosorption was established after a period of 6 hours. Good agreement between experimental data for the three metals and the Langmuir model was found (Say *et al.*, 2001).

Table 35.2 below gives a list of some microorganisms used for removal and/or recovery of metals from industrial wastewaters (Eccles and Hunt, 1986).

9.1 Intracellular accumulation of metals

Microbial cells can accumulate metals, which gain entry into the cell by specific transport systems.

9.2 Adsorption onto cell surface

Interactions between metal ions and the negatively charged microbe surfaces cause microorganisms to bind metals to the surface. Gram-positive bacteria are particularly suited to metal binding. Fungal and algal cells also have a good affinity for heavy metals (Darnall *et al.*, 1986; Ross and Townsley, 1986).

TABLE 35.2 Some microorganisms involved in metal removal/recovery from industrial wastewaters

Microorganisms	Metal
<i>Zooglea remigera</i>	Cu
<i>Saccharomyces cerevisiae</i>	U and others
<i>Rhizopus arrhizus</i>	U
<i>Chlorella vulgaris</i>	Au, Zn, Cu, Hg
<i>Aspergillus oryzae</i>	Cd
<i>Aspergillus niger</i>	Cu, Cd, Zn
<i>Penicillium spinulosum</i>	Cu, Cd, Zn
<i>Trichoderma viride</i>	Cu
AMT Bioclaim™	Biotechnology-based use of granulated product derived from biomass.

Removal of metals from solution may take place by various mechanisms (Brierley *et al.*, 1989; Sterrit and Lester, 1986; Trevors, 1989; Trevors *et al.*, 1985).

Adsorption of metals by activated sludge biomass has been found to follow Langmuir and Freundlich isotherms (Mullen *et al.*, 1989).

The sublethal effects of the heavy metals Cu, Cr, Pb and Zn have been studied and they were shown to adsorb onto biomass much faster than organic competitors. This is probably due to the increased mobility of the metal ion over the organic molecules due to their relative size and greater charge. The metal ions, being much more mobile, are able to seek out adsorption sites quicker than their larger organic competitors on the bioflocs.

'Biosorption being a process that utilizes inexpensive dead biomass to sequester toxic heavy metals and is particularly useful for the removal of contaminants from industrial effluents' (Kratohvil and Volesky, 1998).

9.3 Complexation

Microorganisms are known to produce weak acids such as citric acid, which may form metal chelates. Metals may also be complexed by carbonyl groups found in microbial polysaccharides and other polymers. This is of importance in wastewater treatment plants, particularly those using the activated sludge process, where industrial wastes are treated (Bitton and Freihoffer, 1978; Brown and Lester,

1979, 1982; Rudd *et al.*, 1984; Sterrit and Lester, 1986; McLean *et al.*, 1990). *Pseudomonas putida* has a cystein-rich protein that readily binds with Cd (Higham *et al.*, 1984).

9.4 Precipitation

Some bacteria promote metal precipitation producing ammonia, organic bases, or hydrogen sulphide, which precipitate metals as hydroxides or sulphides. Sulphate-reducing bacteria transform SO_4 to H_2S , which promotes the extracellular precipitation of metals from solution. *Klebsiella aerogenes* is able to detoxify Cd to CdS, which precipitates as electron-dense granules at the cell surface. This process is induced by Cd (Aiking *et al.*, 1982).

It has also been shown, using ICP, that by using the pyridine-thiol ligand Cu and Cd can be removed from solution at pH values of 4.5 and 6.0 respectively by as much as 99.99%; the starting concentrations being 50 ppm. The multiple bonding sites on the pyridine-thiol ligands are responsible for the metal precipitation (Matlock *et al.*, 2001).

9.5 Volatilization

Some metals are transformed to volatile species as a result of microbial action. Bacterially mediated methylation converts Hg^{2+} to $(\text{CH}_3)_2\text{Hg}$, a volatile compound. Some bacteria have the ability to detoxify Hg by transforming Hg^{2+} to Hg^0 , which again is volatile. This detoxification process is plasmid-encoded and is regulated by an operon consisting of several genes. The most important gene is the *merA* gene, which is responsible for the production of mercuric reductase, the enzyme that catalyses the transformation Hg^{2+} to Hg^0 .

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