

***Brock Biology of Microorganisms 15th edition
Madigan Solutions Manual***

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Summary

Chapter 2 is an excellent introductory overview of the structure and function of both prokaryotic and eukaryotic cells. For courses designed for nonscience majors, this chapter provides general details on each topic that, if supplemented with material from related chapters later in the text, may be sufficient background for most students. However, it is recommended that Chapter 2 be used to set the stage for more detailed coverage later in the course.

2.1 | Cell Morphology

Using Figure 2.1, point out the three major morphologies of prokaryotic cells (*coccus*, *rod*, and *spirillum*). Inform your students that, in some species, the cells remain attached following cell division, giving rise to different arrangements that are often genus-specific. For example, coccus cells may exist as short chains (*Streptococcus*) or grapelike clusters (*Staphylococcus*). Less common cell morphologies also exist, such as *spirochetes*, *appendaged* (budding) bacteria, and *filamentous* bacteria (Figure 2.1). Stress to students that these morphologies are only *representative* of those found in nature. Other unusual shapes have also been described in rare cases (for example, square and star-shaped cells!).

Before the molecular era, morphological and physiological properties were used to classify bacterial species. However, we now know that these criteria are poor predictors of evolutionary relationships. For example, certain species of *Archaea* may appear identical in size and shape to species of *Bacteria* under the microscope, but these organisms are of different phylogenetic domains and thus are not closely related to one another on an evolutionary basis. The cell morphology of a particular species is primarily a result of selective pressures in a given habitat that favored a particular cell shape for enhanced reproductive success.

2.2 | The Small World

The presentation in the text of *the significance of being small* is an important concept for students to internalize as they progress in their study of microbiology. Table 2.1 shows the wide size range variability of bacterial cells, which range from a diameter of about 0.2 μm to over 700 μm . Use the two examples of unusually large bacteria discussed in this section to illustrate the current upper limit of bacterial cell size: (1) the surgeonfish gut symbiont *Epulopiscium fishelsoni* (>600 μm in length; Figure 2.2a), and (2) the sulfur chemolithotroph *Thiomargarita namibiensis* (750 μm ; Figure 2.2b). The evolutionary “rationale” for the existence of unusually large-celled bacteria is a mystery when one considers that the metabolic rate of a cell varies inversely with the square of its

size. Ask your students for ideas and/or hypotheses that might explain the selective advantage of large cell size in these two prokaryotes.

The fact that bacteria can live independently as single cells (unlike an individual cell of a multicellular organism) suggests that they must possess some capabilities that provide a selective advantage over their multicellular counterparts that ensure their survival on the planet. Small cells have more *surface area* to *volume* (i.e., a higher *surface-to-volume ratio*), and this alone confers many of the evolutionary advantages of being small, including the following:

- Rapid nutrient and waste transport into and out of the cell allows for faster metabolic rates and growth rates.
- Rapid growth rates result in the rapid production of large populations of cells. These populations, in turn, can greatly affect the physiochemical conditions of an ecosystem within a short time period.
- Transport rates are a function of the surface area of the cytoplasmic membrane relative to cell volume. Use Figure 2.3 to mathematically demonstrate to students that the *surface area* of a sphere is a function of the square of the radius, whereas the *volume* of a sphere is a function of the cube of the radius. This means that the surface-to-volume ratio of a spherical cell can be expressed as $3/r$, where r equals the radius of the cell. Therefore, a coccus cell having a smaller radius has more surface area per volume, and thus more efficient transport capabilities, than a coccus cell having a larger radius.
- Rates of evolutionary change are higher in smaller, faster growing haploid cells than in larger, slower growing diploid cells. This allows for greater adaptive potential through rapid selection for advantageous mutations and counterselection against deleterious mutations.

The theoretical lower limit of size for a living cell is likely near 0.2 μm in diameter. This limit is dictated by the amount of volume required to contain cellular components that are crucial for maintaining life, such as (1) the presence of essential genes on the chromosome; (2) having a sufficient number of ribosomes; and (3) containing a minimal number of metabolic, structural, and transport proteins within the cell. Challenge students to list these and other molecular components that a cell would have to contain to maintain life. Remind students that some cells are parasitic in nature. Inform them that, much like viruses, such microorganisms often have streamlined genomes that lack important genes and may make them dependent upon their hosts for growth. Can such organisms truly be considered living? *This might make a good outside project for group debate, requiring students to view the cell as a three-dimensional physical structure constrained in space and to research a problem that is currently being debated.*

2.3 | The Cytoplasmic Membrane

The structure of the cytoplasmic membrane, a phospholipid bilayer, should be discussed in considerable detail because it plays a critical role in establishing and maintaining the cell's internal environment. Students must understand that the cytoplasmic membrane is the *selectively permeable* boundary between the cytoplasm of the cell and the cell's immediate environment. If the integrity of the membrane becomes compromised, then essential cellular components can leak out of the cytoplasm and into the environment, thereby destroying the cell. Convey to students that the cytoplasmic membrane generally does *not* confer a specific shape and provide rigid support to the cell (these are roles of the cell wall, to be discussed later), but rather the membrane has a fluid nature that allows for a degree of lateral movement of phospholipids and proteins (Figures 2.4 and 2.5). Proteins embedded in the membrane consist of both hydrophobic regions that are situated within the lipid portion of the phospholipid bilayer and hydrophilic regions that are oriented toward either the external environment or the aqueous cytoplasm of the cell.

In contrast to eukaryotic cells, which contain rigid sterol molecules to strengthen and stabilize membranes (especially those of animal cells, which lack cell walls), most prokaryotic membranes instead contain planar molecules called *hopanoids* that serve a similar function. Exceptions to this generalization include methanotrophic bacteria, which contain large amounts of sterols in internal membranes, and the mycoplasmas, a group of parasitic bacteria that lack cell walls.

While members of the *Bacteria* and *Eukarya* contain *ester* linkages that bond the fatty acids to glycerol in their membranes (Figure 2.4a), *Archaea* contain *ether* linkages between the glycerol and lipid portions of their membranes. In addition, archaeal membrane lipids are not composed of fatty acids but instead consist of repeating five-carbon isoprene units that combine to form 20-carbon *phytanyl* side chains (Figure 2.6a and d). Together, the glycerol and phytanyl form a *glycerol diether*. In some *Archaea*, glycerol diethers are joined at their hydrophobic ends to create a lipid monolayer of diglycerol tetraethers (Figure 2.6b and e). This structural conformation provides superior thermostability of the membrane, and indeed lipid monolayers are most commonly found in hyperthermophilic archaeal species. Finally, members of the *Crenarchaeota* often contain *crenarchaeol*, a unique monolayer membrane lipid having four cyclopentyl rings and one cyclohexyl ring (Figure 2.6c). Despite the molecular differences between archaeal membranes and bacterial/eukaryotic membranes, their basic structural properties are the same in that each possesses hydrophobic interior hydrocarbon chains attached to polar (hydrophilic) glycerophosphate molecules.

Although molecular adaptations of membranes to high and low temperatures are discussed in some detail in Chapter 5, this may be a good opportunity to introduce the topic of saturated versus unsaturated hydrocarbon chains and discuss how they relate to membrane fluidity under high and low temperature extremes (e.g., why vegetable shortening is a solid at room temperature, and vegetable oil is a liquid under the same conditions).

The major functions of the cytoplasmic membrane are summarized in Figure 2.7 and include its role as (1) a *permeability barrier*, (2) a *protein anchor*, and (3) a means of *energy conservation*. With respect to acting as a permeability barrier, impress upon students that even extremely small ions do not freely pass through the hydrophobic interior of the membrane due to their charges. While water molecules do diffuse through membranes (due to their small size and only weak polarity) in a process called *osmosis*, the movement of water across membranes is greatly accelerated by water transport proteins called *aquaporins*. These transport proteins have been identified in the membranes of organisms from all domains of life but are perhaps best

studied in the bacterium *Escherichia coli*.

Introduce students to the concept that a membrane can function much like a battery in that it can store potential energy. By separating protons to the outside of the membrane from hydroxyl ions on the inside, the membrane becomes “energized” (i.e., polarized), and this energized state is referred to as the *proton motive force (PMF)*. The dissipation of this force results in the conversion of potential energy to kinetic energy. When protons stored outside of the membrane return to the inside of the cell through an ATP synthase enzyme complex, ADP and P_i are converted to ATP, the cell’s chemical energy currency. This concept will be discussed in detail in Chapter 3.

Discuss with your students the necessity for membrane-bound transport proteins by comparing the rate of simple diffusion of a solute across a membrane to the greatly accelerated rate of carrier-mediated transport of a solute across a membrane (Figure 2.8). Transport proteins allow for the accumulation inside a cell of a solute that may be in very low concentration in the environment. Point out that carrier-mediated transport proteins may show high or low specificity for a given solute, but in either case, the movement of molecules across the membrane is typically faster with a transporter than by simple diffusion alone.

2.4 | Bacterial Cell Walls: Peptidoglycan

The bacterial cell wall warrants extensive coverage in the classroom. Research on its structure and function can be traced back to the early history of microbiology. It began with the early observation of the differential reaction of various bacterial cells to the Gram stain. This stain distinguished two types of bacteria based on the composition of the cell wall: gram-positive and gram-negative. Research proceeded with the discovery that both lysozyme and penicillin induced cell lysis and with the realization that some of the bacterial cell wall constituents (diaminopimelic acid and N-acetylmuramic acid) were unique. These discoveries were exciting. They increased our understanding of prokaryotic cells and helped to obtain better chemotherapy with which to combat bacterial diseases. The mechanisms of peptidoglycan biosynthesis, cell division (covered in Chapter 5), osmotic lysis, and the activity of penicillin are important topics of discussion because they provide striking examples of the interrelationship of basic knowledge and practical applications of great significance.

Figure 2.9 provides an excellent summary of the differences in structure and appearance of gram-positive versus gram-negative cell walls. Point out the fundamental repeating structure of peptidoglycan (the *glycan tetrapeptide*; Figure 2.10), which consists of alternating N-acetylglucosamine (NAG) and N-acetylmuramic acid (NAM) residues. Indicate that the latter of these two sugars is connected to a short peptide chain consisting of four amino acids, including in many bacteria the unique lysine analog diaminopimelic acid (DAP). The peptide chains provide structural rigidity to peptidoglycan by cross-linking the polysaccharide layers such that tensile strength is conferred on the cell wall in both the X and Y directions (Figure 2.11). Although there is some variation in the amino acid composition of these peptide cross-linkages, there is great unity within the bacteria regarding the presence of N-acetylmuramic acid, DAP (which may be replaced by lysine), and D-amino acids (D-alanine and D-glutamic acid) rather than the usual L stereoisomers found in proteins. However, in contrast to gram-negative bacteria, some NAM residues in the peptidoglycan of gram-positive bacteria contain covalently bound teichoic acids,

polyalcohols joined by phosphate esters (Figure 2.12). Teichoic acids contribute to the overall negative charge of the gram-positive cell surface and help to sequester cationic micronutrients (e.g., Ca^{2+} and Mg^{2+}) from the environment.

At this time you may want to foreshadow the structure of the gram-positive bacterial cell wall in the context of antibiotic therapy and design. Emphasize that examples of unique cell chemistry often provide targets for successful chemotherapy without the problems of host toxicity (Figures 2.10–2.14). The mechanisms of action of both lysozyme and penicillin are good examples of how a chemical agent can destroy peptidoglycan, resulting in bacterial cell lysis.

Finally, note that there are also prokaryotic cells that lack a cell wall, including the mycoplasmas, a group of pathogenic bacteria (see Chapter 16), and species of the archaeal genera *Thermoplasma* and *Ferroplasma* (see Chapter 17). As previously noted, the mycoplasmas are unusual among bacteria in that they contain sterols in their cytoplasmic membranes. The structural rigidity provided by these molecules presumably helps to maintain cell integrity during mild osmotic stress.

2.5 | LPS: The Outer Membrane

The outer membrane of gram-negative bacteria is obvious in TEM sections, where it is seen as a wavy lipid bilayer outside of a thin layer of peptidoglycan. In the outer membrane, *lipopolysaccharide* (LPS) (Figure 2.13) replaces most of the phospholipids in the outer leaflet, whereas lipoproteins in the inner leaflet function to anchor the outer membrane to peptidoglycan (Figure 2.14). Depending upon the chemistry background of your students, discuss the chemical components of the LPS: (1) *Lipid A*, the phosphoglycolipid portion of the LPS; (2) the *core polysaccharide*, consisting of ketodeoxyoctonate (KDO), heptoses (7-carbon sugars), hexoses, and *N*-acetylglucosamine; and (3) the *O-polysaccharide*, consisting of repeating sequences of hexoses that form long chains and may be branched (Figure 2.13).

Although the purpose of the outer membrane is structural, it is toxic to animals due to the lipid A component of the LPS. Toxins that are part of the cell wall of gram-negative bacteria are called *endotoxins*. Provide examples for your students of endotoxic human pathogens (e.g., *Shigella*, *Salmonella*, and *Escherichia*) that elicit ill effects in the host, including fever, gastrointestinal distress, and septic shock.

In contrast to the cytoplasmic membrane, the outer membrane is permeable to small molecules due to membrane channels called *porins* (Figure 2.14a and c), which vary in specificity from nonspecific to highly specific. Students should be made aware that the *periplasm* of gram-negative bacteria contains binding proteins that are not present in gram-positive bacteria. The periplasm contains a number of different classes of enzymes, some of which facilitate transport (Section 3.2) or chemotaxis (Section 2.13).

2.6 | Archaeal Cell Walls

Cell walls of *Archaea* do not contain peptidoglycan, but they do possess diverse chemistries that include proteins, polysaccharides, and glycoproteins. Some methanogens (Chapter 17) produce a polysaccharide similar to peptidoglycan called *pseudomurein* (Figure 2.15). Point out that the β -1,3 glycosidic linkage in pseudomurein is different from the β -1,4 linkage in peptidoglycan, thus making the former insensitive to the action

of lysozyme. There are no known human pathogens from the *Archaea*, and thus the evolution of lysozyme most probably arose from the interactions of *Bacteria* with animal hosts over time.

Although not all *Archaea* contain pseudomurein, nearly all contain a cell wall of some type (exceptions were noted in Section 2.4). Extremely halophilic *Archaea* have sulfate (SO_4^{2-}) incorporated into their cell walls to bind excessive sodium (Na^+) and help shield the cell from its extremely salty environment. Other *Archaea* (and some *Bacteria*) have a paracrystalline surface layer, the *S-layer*, composed of protein or glycoprotein (Figure 2.16). Discuss the potential functions of S-layers, which are varied and may include protecting the cell from osmotic lysis; preventing the access of larger particles, such as viruses, to the cytoplasmic membrane; and retaining secreted proteins near the cell surface.

2.7 | Cell Surface Structures

Cell surface structures produced by bacteria that are not an integral part of the cell wall are generally not considered essential to cell survival. However, the presence of *capsules* and *slime layers* (Figure 2.17), *fimbriae* (Figure 2.18), and *pili* (Figure 2.19) on many prokaryotes suggests that such structures play important ecological roles for these organisms, including the establishment of pathogenic associations with host cells. The attachment of one cell to another is a specific molecular interaction between host and pathogen, and this contact often initiates changes in the host cell resulting in internalization of the pathogen and continuation of its life cycle.

Although details of host–pathogen interaction are not part of the material presented here, you could pique student interest by showing a specific example of the role played by these cell surface structures in a specific pathogenesis (e.g., *Streptococcus pneumoniae*, *Yersinia pestis*, and *Listeria monocytogenes*). Clearly define the structural and functional differences of fimbriae, pili, and flagella for students so that they do not equate these structures based on their similar microscopic appearance. Lastly, point out the unique attachment structures, called *hami*, of the SM1 group of *Archaea* (Figure 2.20). These molecular grappling hooks, which resemble type IV pili in *Bacteria*, facilitate the formation of a dense biofilm network that may help trap nutrients in their deep subsurface habitat.

2.8 | Cell Inclusions

Many bacterial cells contain inclusions, such as the storage granules *polyhydroxyalkanoate* (*PHA*, of which *polyhydroxybutyrate* [*PHB*] is one type; Figure 2.21) and *glycogen*, both of which serve as carbon and energy reserves. Additional nutrient inclusions include polyphosphate and elemental sulfur granules (Figure 2.22). The latter of these inclusions serves as an important secondary energy source for a variety of phototrophic and chemolithotrophic bacteria that oxidize sulfide (H_2S) and other reduced sulfur compounds as electron donors (see Chapter 14).

Other storage inclusions are not necessarily for nutritional purposes. Many prokaryotes catalyze biomineralization, the process of mineral formation by microorganisms. Figure 2.23 shows a beautiful example of benstonite granule accumulation inside a cell of the cyanobacterium *Gleomargarita*; the function of these structures is unknown but it may be to provide ballast for the cell in its aqueous environment. A discussion of magnetosomes and magnetotactic bacteria should be of interest to students, who will likely find the idea of

“magnetic bacteria” intriguing. Although the function of magnetosomes is also unknown, they are most certainly important to species that form them (Figure 2.24). One hypothesis is that the magnetite in these inclusions acts like a compass, pulling the microaerophilic aquatic bacteria that contain them downward toward the Earth’s magnetic poles and into the sediments where dissolved oxygen (O₂) concentrations are lower. Unlike polyphosphate and elemental sulfur granules, both magnetosomes and PHA inclusions have a “nonunit” (single layer) phospholipid membrane.

2.9 | Gas Vesicles

Gas vesicles are rigid, hollow structures in the cytoplasm of some cells that allow vertical migration in a water column. They are therefore considered a mechanism of motility (Figures 2.25 and 2.26). The proteinaceous shell is permeable to gases but not to water and solutes. At the molecular level, the shell contains two proteins: GvpA, a rigid β -sheet that makes up 97% of the shell; and GvpC, a cross-linking protein made up of α -helices (Figure 2.26b). These structures are found mostly in aquatic phototrophs, allowing them to regulate their position in the water column where the light intensity required for photosynthesis is optimal. Some nonphototrophs, including some species of *Archaea*, also contain gas vesicles.

2.10 | Endospores

Introduce your discussion of endospores by reminding students of Ferdinand Cohn’s discovery of the endospore-forming genus *Bacillus* and his research demonstrating the incredible heat resistance of these structures (Chapter 1). Because of Cohn’s efforts, important new methods of sterilization were developed that are still used by the food and medical industries. To spark student interest, note that many endospore-forming bacteria are also pathogenic and cause some of the most serious diseases known. For example, pathogenic members of the genera *Bacillus* and *Clostridium* often produce potent toxins that cause fatal diseases if not treated within a short time. Examples include botulism (*C. botulinum*), tetanus (*C. tetani*), gas gangrene (*C. perfringens*), gastroenteritis (*C. difficile*), and anthrax (*B. anthracis*).

Depending on the level of your course, discuss the structure of endospores and the endosporulation and germination processes in some detail (Figures 2.27–2.32). Key points to stress include the following:

- Describe the unique nature of the core, stressing the functions of the dipicolinic acid (DPA) and calcium (Ca²⁺) complexes (Figure 2.31), the low water content and low pH, and the role of small acid-soluble spore proteins (SASPs) in protecting the DNA and in serving as a carbon and energy source for the cell during germination.
- Discuss endospore formation as an example of cellular differentiation in prokaryotes, using *Bacillus subtilis* as a model (Figure 2.32). To impress upon students the remarkable complexity of the differentiation process, mention that more than 200 genes are involved in sporulation, and many details of the process are still being investigated in laboratories around the world.
- Point out the key differences between vegetative cells, which are metabolically active and dividing cells, and endospores, which are essentially dormant cell forms that exhibit minimal enzymatic activity (Table 2.2).

Finally, students should show interest in a discussion concerning how long endospores can remain viable. The debate on the longevity of these structures has now pushed their life

span to millions of years. If experimental evidence from independent research laboratories repeatedly supports these claims, this would indeed be an extraordinary testament to the life-preserving design of these structures.

2.11 | Flagella, Archaeella, and Swimming Motility

The ability of *Bacteria* to move via flagella (or *Archaea* to move via archaeella) is intimately connected to their ability to sense and respond to environmental signals through complex signal transduction pathways. Flagella are arranged in a variety of ways on the cell surface (Figures 2.34 and 2.35). Rather than moving in a whip-like motion like the flagella of eukaryotes, bacterial flagella and archaeella *rotate* to propel the cell (Figure 2.36, 2.37, and 2.39). For many species, this rotation is reversible, and the direction of the rotation dictates whether the cell moves forward (called a *run*), backward, or tumbles in place; the latter occurs for cells having peritrichous flagellation (multiple flagella arranged all around the cell).

The flagellar structure is complex, and its synthesis and assembly involve more than 40 genes in *E. coli* (Figures 2.37 and 2.38). Rotation of bacterial flagella requires significant energy directly from a *proton motive force* (PMF; Section 2.3 and Chapter 3). In fact, a single rotation requires the translocation of about 1000 protons across the cytoplasmic membrane through the Mot complex (Figure 2.37b). Although bacterial flagella do not rotate at a constant speed, up to 300 revolutions per second are possible, resulting in extremely fast movement of about 60 cell lengths per second. When measured as the number of body lengths moved per second, a bacterium swimming at full speed would be moving nearly 2.5 times faster than a cheetah can run! As impressive as this is, the fastest known cells are actually *Archaea* of the genus *Methanocaldococcus*, which are capable of swimming nearly *500 cell lengths per second*. Thus, relative to its size, this species is easily the fastest organism on Earth.

Flagella from the different domains of life exhibit significant structural and operational differences. For example, bacterial flagella are about twice as thick as archaeella. In addition, the filament portion of all bacterial flagella is composed of a single type of flagellin protein, whereas the number of proteins that compose archaeellar filaments varies depending on the species. Because of their structural similarity, the archaeellum has been likened to a rotating version of the type IV pilus of bacteria, an ATP-powered appendage that allows for twitching motility (Section 2.7). As such, archaeellar rotation is powered directly by the hydrolysis of ATP rather than by a PMF, as is the case for bacterial flagella. The fundamental differences between bacterial flagella, eukaryotic flagella, and archaeella suggest that these mechanisms of motility have arisen independently as a result of convergent evolution rather than from a common origin.

2.12 | Gliding Motility

Gliding motility (motility without flagella) in bacteria is a relatively underrepresented phenomenon in microbiology texts, and this is probably because of the lack of knowledge of the molecular mechanisms involved in the process. Consequently, it is a good puzzle to present to students following your discussion of flagellar locomotion, about which much is known. Gliding motility has never been observed in *Archaea*, but several species of gliding *Bacteria* are known, with key examples including species of cyanobacteria and the

genera *Myxococcus*, *Cytophaga*, and *Flavobacterium* (Figure 2.40). Movement by gliding requires a solid surface and is considerably slower than flagellar motility. Several mechanisms of gliding motility have been described. Some bacteria secrete a polysaccharide slime that adheres to a surface and pulls the cell forward. Others exhibit the aforementioned twitching motility, in which cell movements are carried out by the repeated extension and retraction of type IV pili in a grappling hook style of motion. Cells of *Flavobacterium johnsoniae* appear to glide via the coordinated ratcheting of cytoplasmic membrane proteins with outer membrane proteins, where the latter move along a surface in the opposite direction of the cell itself, much like the movement of a tank on its tracks (Figure 2.41).

2.13 | Chemotaxis and Other Taxes

The ability of bacteria to exhibit taxes (i.e., directed movement) confers a selective advantage depending upon environmental conditions. Students should understand that, unlike larger organisms, microorganisms sense gradients in a *temporal* (an effect lasting for only a short time) rather than a *spatial* (a lingering effect) manner. In other words, they must continually compare their current external conditions with those of a few moments before. The studies of chemotaxis in *E. coli* provided the first genetic model of the process in swimming bacteria. Chemotaxis will be discussed in Chapter 6 in the context of two-component signal transduction systems, but use Figures 2.42 and 2.43 to show the run and tumble “directed” response and capillary assay system used to evaluate and identify signal molecules that act as attractants or repellants.

Many of the protein components that function in chemotactic pathways are also activated during phototaxis, and flagellar rotation is controlled accordingly. The response of *Rhodospirillum centenum* to light is a fascinating example of phototaxis. Mention to students that *scotophobotaxis* (movement away from dark) is not the same as true phototaxis, which involves movement up a light gradient (Figure 2.44). *R. centenum* is also unusual in that an entire colony of cells on solid media will move toward an infrared light source (the wavelengths absorbed by their photosynthetic pigments) and away from fluorescent light. If one observes the cells within the colony as the colony moves in one direction, the individual cells appear to be moving more or less randomly, suggesting that there must be some intercellular communication occurring to generate a net directional movement.

Other taxes have been observed in microorganisms, including directed movement toward or away from oxygen (*aerotaxis*), toward a specific osmotic condition (*osmotaxis*), or toward water (*hydrotaxis*).

2.14–2.16 | Eukaryotic Microbial Cells

Students should be familiar with the organelles and general structure of the eukaryotic cell from general biology courses, but you should still present a brief overview of the topic, focusing first on the nucleus and chromosome organization (Figures 2.45 and 2.46). Remind students that eukaryotic DNA is packaged in the nucleus by being wound around positively charged *histone* proteins. Note that transport of proteins and nucleic acids through *nuclear pores* requires the energy of GTP. The *nucleolus* is the site of ribosomal RNA (rRNA) synthesis and assembly of the large and small subunits of the ribosome. Also

review *mitosis* (Figure 2.47) and *meiosis* with your students, reminding them that these processes, which occur only in eukaryotes, are mechanisms by which a cell divides to create two diploid daughter cells or four haploid gametes (or spores), respectively.

Review mitochondrial structure (Figure 2.48) and function with your students, and allow this to lead into a discussion of the *hydrogenosome* (Figure 2.49). The latter organelle is found in certain eukaryotes that lack mitochondria, the present-day representatives of which arose early after the divergence of *Eukarya* from *Archaea* (see Chapter 13). Unlike mitochondria, hydrogenosomes usually lack cristae and do not contain citric acid cycle enzymes. Therefore, organisms possessing hydrogenosomes are strictly fermentative. Several are parasites, such as the sexually transmitted pathogen *Trichomonas vaginalis*, and all are either aerotolerant or obligate anaerobes. Figure 2.49 shows the pyruvate oxidation scheme carried out by the hydrogenosome for ATP synthesis.

Chloroplasts are the structures found in plant cells and many microbial eukaryotes (e.g., dinoflagellates, euglenids, diatoms, and various algae) that carry out photosynthesis and allow for photoautotrophic growth (Figure 2.50). The relationship of chloroplasts, mitochondria, and hydrogenosomes to *Bacteria* is well documented and is discussed briefly here as the *endosymbiotic hypothesis*. Although this topic will be addressed in more detail in future chapters (see Sections 13.4 and 18.1), you may wish to point out key features supporting endosymbiosis to your students at this time:

- All three structures (mitochondria, chloroplasts, and hydrogenosomes) contain their own DNA in covalently closed circles. This DNA encodes rRNAs, transfer RNAs (tRNAs), and some respiratory enzymes.
- These organelles contain their own ribosomes that are bacterial in structure and are sensitive to antibiotics that affect bacterial ribosomes.
- The nuclear DNA of eukaryotic cells contains bacterially derived genes, lending additional evidence to support the endosymbiotic hypothesis (Section 18.1).
- 16S rRNA gene sequence analyses show the evolutionary relatedness of these structures to *Bacteria*.

Finally, review the function of other eukaryotic cell structures (e.g., the endoplasmic reticulum, Golgi complex, ribosomes, and cytoskeletal elements) mentioned in Section 2.16, which should already be familiar to most students (Figures 2.51–2.53). Point out that eukaryotic flagella are powered by ATP hydrolysis and propel the cell via a whip-like motion rather than by rotation, as seen in bacterial flagella and archaella (Section 2.11).

Answers to Review Questions

1. The major prokaryotic morphologies are coccus, rod, spirillum, spirochete, appendaged, and filamentous. Several variations and subcategories exist.
2. Cell sizes of bacteria typically range from about 0.4 μm in length to several hundred micrometers, with most cells averaging 1–3 μm . The maximum size of a bacterial cell appears to be in excess of 750 μm , and the smallest is likely to be about 0.2 μm . We are better able to know the lower limit of size for a bacterial cell than the upper limit because we know the minimum amount of volume required for genetic material, ribosomes, proteins, and other cellular constituents necessary to carry out metabolism. An *E. coli* cell is 1–2 μm in length and 0.5–0.6 μm wide.
3. Because phospholipids, the major structural component of unit membranes, contain both hydrophobic and hydrophilic moieties, their aggregation in aqueous solution leads to the

formation of a bilayer in which hydrophobic lipids face each other in the internal region and polar glycerophosphates communicate with the external environment. In contrast to *Bacteria* and *Eukarya*, which use ester linkages to bond fatty acids to the glycerol molecule, lipids from *Archaea* have ether linkages between the glycerol and their hydrophobic side chains. Moreover, archaeal lipids lack fatty acids and instead have side chains composed of repeating units of the C₅ hydrocarbon isoprene that combine to form the C₂₀ molecule phytanyl. Glycerol diethers, diglycerol tetraethers, and crenarchaeol are the major classes of lipids present in species of *Archaea*. The phytanyl side chains of diglycerol tetraether are covalently bonded together at their ends, thus forming a phospholipid monolayer instead of a bilayer. Small, uncharged molecules, such as gases, certain solvents, and to some degree water, are able to diffuse through the cytoplasmic membrane in a direction congruent with their concentration gradient. Ionized molecules, which have a net charge, are hydrophilic and are therefore repelled by the hydrophobic interior of the membrane. Such molecules are brought into the cell through membrane-spanning transport proteins.

4. Because the rigid layer is a polymer composed of sugars linked by β -1,4 glycosidic bonds and polypeptides that cross-link *N*-acetylmuramic acid residues, the term *peptidoglycan* (“protein” + “sugar”) is appropriate. Together, the covalent glycosidic and peptide bonds of peptidoglycan provide tensile strength in both the X and Y directions, respectively.
5. Functions of the outer membrane of gram-negative bacteria include antigenicity, adhesion to surfaces, toxicity for animals, provision of membrane channels and porins for the influx and efflux of low molecular weight substances, and a permeability barrier to the passage of high molecular weight substances. The chemical composition of the outer membrane is a phospholipid bilayer containing LPS. The endotoxin lipid A is the lipid portion of the LPS. It consists of fatty acids connected through amine groups to a disaccharide composed of glucosamine phosphate.
6. Peptidoglycan is a polysaccharide present in *Bacteria*, but absent in *Archaea*. S-layers consist of protein or glycoprotein in a paracrystalline array. They are found in both *Bacteria* and *Archaea* and may serve as an outermost selective sieve around the cell to prevent access of large particles, such as viruses, to the cytoplasmic membrane. Methanogenic *Archaea* have cell walls composed of pseudomurein, which is similar in structure to peptidoglycan but contains lysozyme-insensitive β -1,3 glycosidic bonds. Extremely halophilic *Archaea* have polysaccharide cell walls containing large amounts of sulfate, which buffers the cell from its very salty environment by binding Na⁺.
7. Polysaccharide layers produced by bacteria are referred to as *capsules* and/or *slime layers*, and they have several potential functions. They play a major role in the attachment of cells to a surface (a property that allows for biofilm formation); they prevent phagocytosis by immune system macrophages; and they function to prevent desiccation.
8. Cytoplasmic inclusions include carbon polymers (e.g., poly- β -hydroxybutyrate and glycogen), inorganic phosphate (polyphosphate), elemental sulfur, and magnetite. Poly- β -hydroxybutyrate granules serve as a cell energy reserve and consist of a polymeric fatty acid, whereas magnetosomes consist of magnetite (Fe₃O₄) and serve as small magnetic compasses. Aquatic cells may use magnetosomes as a means to align themselves within zones of favorable oxygen concentration.
9. Because of their buoyancy, gas vesicles provide a means of motility for aquatic cells. These

structures are water impermeable because their membranes consist of highly hydrophobic, rigid, cross-linked proteins. Gas-filled vesicles decrease the density of the cell. Therefore, the number of inflated vesicles in a cell determines its buoyancy and position in the water column.

10. An endospore differs from a vegetative cell for several reasons. The cytoplasm of endospores is considerably dehydrated compared to vegetative cells, and endospores have a unique cell surface consisting of multiple layers of proteins. In addition, endospores are “shut down” in the sense that they are essentially metabolically dormant. All of these factors permit significant resistance to heat, radiation, desiccation, toxic chemicals, and extremes of pH, and they allow for remarkable longevity of these structures.
11. The bacterial flagellum is a long, rigid, protein filament of polymerized flagellin that extends from the cytoplasmic membrane and cell wall. It generates cellular motility by rapidly rotating, much like a miniature propeller. A wider hook at the base of the flagellum serves to connect the flagellum to the motor portion (basal body), which is anchored to the cell wall and cytoplasmic membrane. In *Bacteria*, the proton motive force provides the energy for rotation via the flow of protons across the membrane. Archaeella are thinner by comparison and may be composed of several proteins that show homology to the type IV pilus of *Bacteria*. In addition, unlike bacterial flagella, archaeellar rotation is driven by ATP hydrolysis.
12. The gliding motility exhibited by species of *Flavobacterium* differs from the flagellar motility of *E. coli* in several respects. Gliding motility, which requires a surface for the cells to move across, is considerably slower than the swimming motility of flagellated cells. The mechanism of gliding in *Flavobacterium* appears to be by a ratcheting movement between proteins anchored in the cytoplasmic membrane and the outer membrane. However, gliding motility in species of *Flavobacterium* and swimming motility in *E. coli* are both powered by energy from the proton motive force.
13. Motile bacteria move toward attractants by comparing the strength of stimuli (using chemoreceptors) obtained over different moments in time, during which they have changed their position (i.e., the response is temporal). If an environmental stimulus is desirable, flagellar activity results in fewer tumbles and longer runs. If a stimulus is not desirable, then the opposite will occur. Although the new direction following a tumble is random, adjusting the duration of runs and the frequency of tumbles allows the cell to ultimately move toward the attractant, albeit in a somewhat erratic manner. Chemotaxis, therefore, resembles smell rather than sight. In the experiment shown in Figure 2.43, the control is a capillary that contains neither an attractant nor a repellent. The control is essential because it provides an unmanipulated point of comparison and, therefore, serves to validate the results obtained when a variable (i.e., the attractant or repellent) is introduced to the capillary.
14. Three features that clearly differentiate eukaryotic from prokaryotic cells are the following: (1) Almost all eukaryotes possess a membrane-bound nucleus containing their genomic DNA (although, interestingly, there are a few nucleated prokaryotes among the *Planctomycetes*, described in Chapter 16); (2) unlike nearly all prokaryotes, eukaryotic cells contain various membrane-bound organelles (e.g., mitochondria and the Golgi complex); and (3) because prokaryotes have no nucleus, they carry out coupled transcription and translation; these processes are separate in eukaryotes because of the presence of a nuclear membrane. Histones are positively

charged protein complexes that bind and compact DNA in the nucleus of eukaryotic cells.

15. The hydrogenosome and mitochondrion are about the same size, and they both function to catabolize pyruvate. In mitochondria, pyruvate is oxidized completely using enzymes of the citric acid cycle. Carbon dioxide (CO₂) and water (H₂O) are the products of this aerobic respiration, and ATP is synthesized by oxidative phosphorylation using a proton motive force established by the mitochondrial electron transport chain. The hydrogenosome is found only in certain anaerobic eukaryotes. Because it lacks citric acid cycle enzymes and cristae containing electron transport chain components, pyruvate is fermented to acetate, CO₂, and H₂, and ATP is synthesized via substrate-level phosphorylation. The chloroplast is the site of photosynthesis. The light-induced reaction center and electron transport chain generate energy (ATP) and reducing power (NAD[P]H) required to reduce CO₂ to organic carbon (e.g., phosphoglyceric acid) via the Calvin cycle for biosynthetic and metabolic requirements. The important Calvin cycle enzyme RubisCO comprises over 50% of the total chloroplast protein. Mitochondria, chloroplasts, and hydrogenosomes contain circular DNA that is more similar to bacterial DNA than eukaryotic DNA in sequence composition. In addition, these organelles contain their own ribosomes that resemble those of bacteria and, therefore, are sensitive to many of the same antibiotics that target bacterial ribosomes.
16. The endoplasmic reticulum (ER) is a network of membranes continuous with the outer leaflet of the nuclear membrane that is involved in the synthesis of lipids and carbohydrates (smooth ER) and glycoproteins and new membrane material (rough ER). The Golgi complex is a stack of membranes that works in concert with the ER in the production, sorting, and modification of membrane and secretory proteins. Such modifications help direct these products to their final destinations. Lysosomes contain digestive enzymes to break down various macromolecules and recycle them for biosynthesis. The eukaryotic cytoskeleton is composed of (1) microfilaments, which consist of a pair of intertwined actin filaments; (2) intermediate filaments, which are composed of keratin proteins; and (3) microtubules, which are hollow tubes composed of α - and β -tubulin proteins.

Answers to Application Questions

1. The surface-to-volume ratio for a cell having a diameter of 15 μm is 0.4. The surface-to-volume ratio for a cell with a diameter of 2 μm is 1.5. As the diameter of the cell *decreases*, there is more surface area per unit volume. Thus, with a higher surface-to-volume ratio, there is an increased capacity for the cell to interact with its environment per unit of cytoplasm. This generally leads to higher levels of metabolic activity in the cell and, therefore, a faster growth rate.
2. Listed below are a few ways to distinguish *Bacteria* and *Archaea*:

<u>Bacteria</u>	<u>Archaea</u>
Ester links from side chains of glycerol	Ether links from side chains of glycerol
Fatty acid side chains from glycerol	Polyisoprene side chains from glycerol
<i>N</i> -acetylmuramic acid in cell wall	<i>N</i> -acetylglucosaminuronic acid in cell wall
Peptidoglycan in cell wall	Other polysaccharides in cell wall
Lysozyme-sensitive	Lysozyme-resistant
3. Sixty cell lengths per second for an *E. coli* cell that is 2 μm long equals a speed of 120 $\mu\text{m}/\text{sec}$. A 3-cm capillary tube = 30,000 μm , so at this speed it would take 250 seconds, or nearly 4.2 minutes, for the cell to travel the length of the tube.