

Chapter 6

Microbial Growth

Growth

- **increase in cellular constituents that may result in:**
 - **increase in cell number**
 - e.g., when microorganisms reproduce by budding or binary fission
 - **increase in cell size**
 - e.g., **coenocytic** microorganisms have nuclear divisions that are not accompanied by cell divisions
- **microbiologists usually study population growth rather than growth of individual cells**

The Growth Curve

- **observed when microorganisms are cultivated in batch culture**
 - culture incubated in a closed vessel with a single batch of medium
- **usually plotted as logarithm of cell number versus time**
- **usually has four distinct phases**

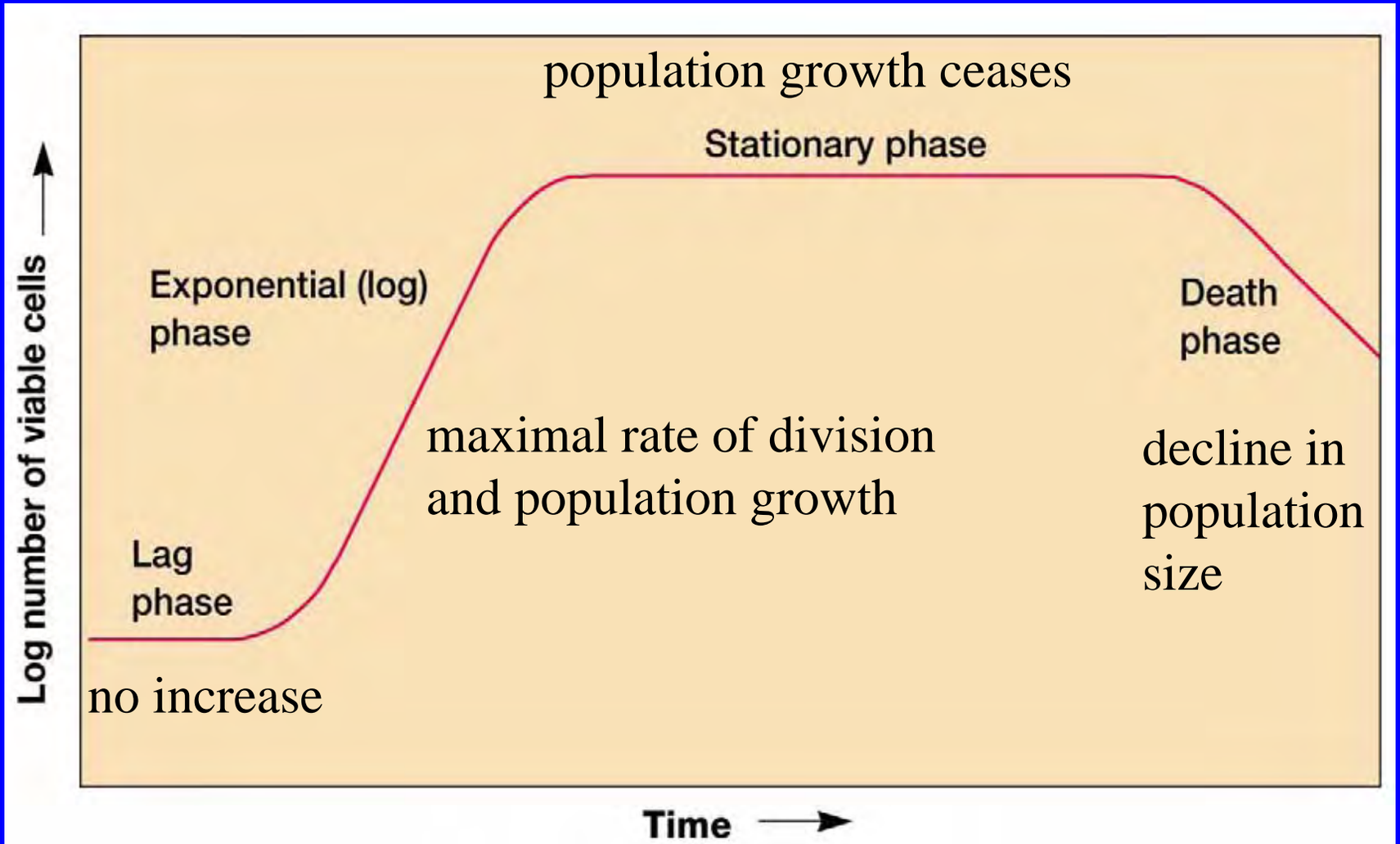


Figure 6.1

Lag Phase

- **cell synthesizing new components**
 - e.g., to replenish spent materials
 - e.g., to adapt to new medium or other conditions
- **varies in length**
 - in some cases can be very short or even absent

Exponential Phase

- also called **log phase**
- **rate of growth is constant**
- **population is most uniform in terms of chemical and physical properties during this phase**

Table 6.1 An Example of Exponential Growth

Time^a	Division Number	2^n	Population ($N_0 \times 2^n$)	$\log_{10} N_t$
0	0	$2^0 = 1$	1	0.000
20	1	$2^1 = 2$	2	0.301
40	2	$2^2 = 4$	4	0.602
60	3	$2^3 = 8$	8	0.903
80	4	$2^4 = 16$	16	1.204
100	5	$2^5 = 32$	32	1.505
120	6	$2^6 = 64$	64	1.806

^aThe hypothetical culture begins with one cell having a 20-minute generation time.

cells are dividing and doubling in number at regular intervals

each individual cell divides at a slightly different time

curve rises smoothly rather than as discrete steps

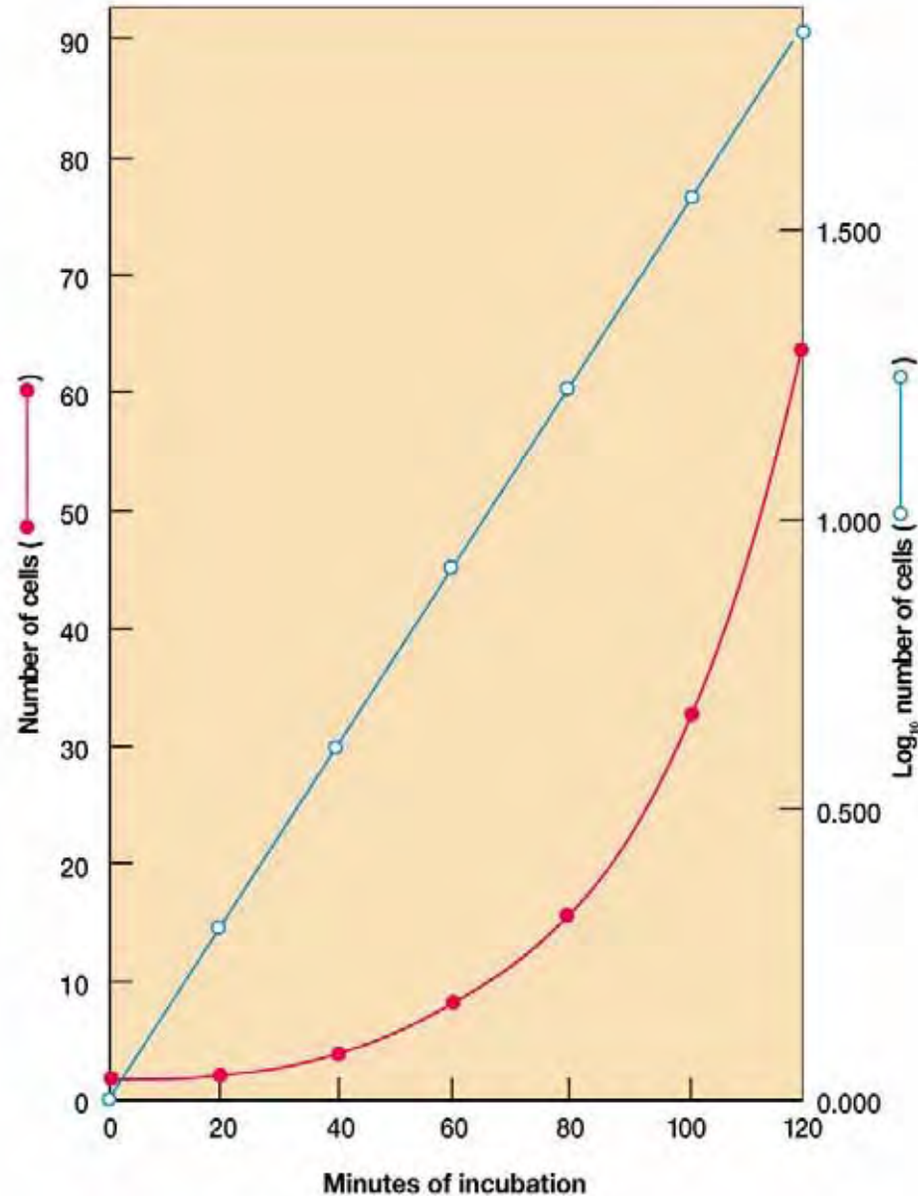


Figure 6.3

Balanced growth

- during log phase, cells exhibit **balanced growth**
 - cellular constituents manufactured at constant rates relative to each other

Unbalanced growth

- **rates of synthesis of cell components vary relative to each other**
- **occurs under a variety of conditions**
 - **change in nutrient levels**
 - **shift-up (poor medium to rich medium)**
 - **shift-down (rich medium to poor medium)**
 - **change in environmental conditions**

Effect of nutrient concentration on growth

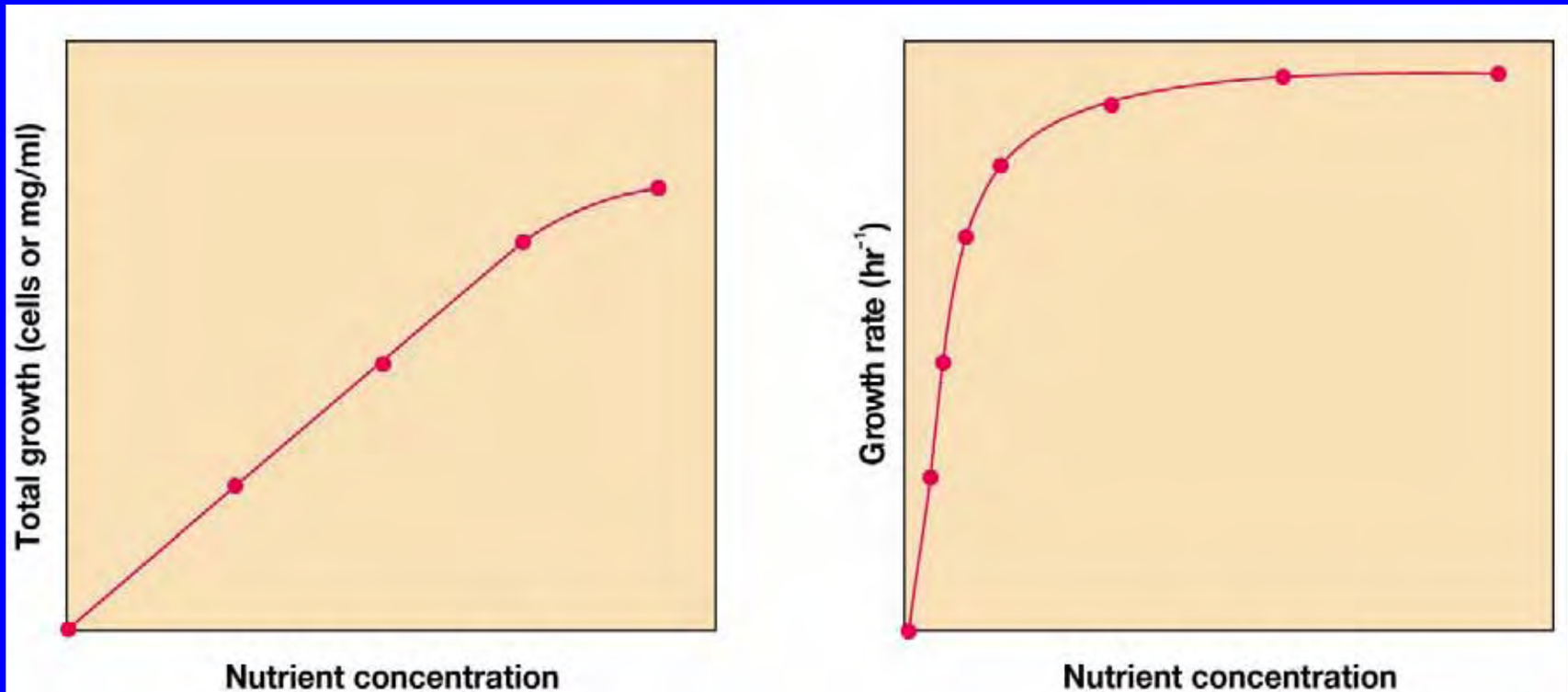


Figure 6.2

Stationary Phase

- **total number of viable cells remains constant**
 - **may occur because metabolically active cells stop reproducing**
 - **may occur because reproductive rate is balanced by death rate**

Possible reasons for entry into stationary phase

- **nutrient limitation**
- **limited oxygen availability**
- **toxic waste accumulation**
- **critical population density reached**

Starvation responses

- **morphological changes**
 - e.g., endospore formation
- **decrease in size, protoplast shrinkage, and nucleoid condensation**
- **production of starvation proteins**
- **long-term survival**
- **increased virulence**

Death Phase

- **cells dying, usually at exponential rate**
- **death**
 - **irreversible loss of ability to reproduce**
- **in some cases, death rate slows due to accumulation of resistant cells**

The Mathematics of Growth

- **generation (doubling) time**
 - time required for the population to double in size
- **mean growth rate constant**
 - number of generations per unit time
 - usually expressed as generations per hour

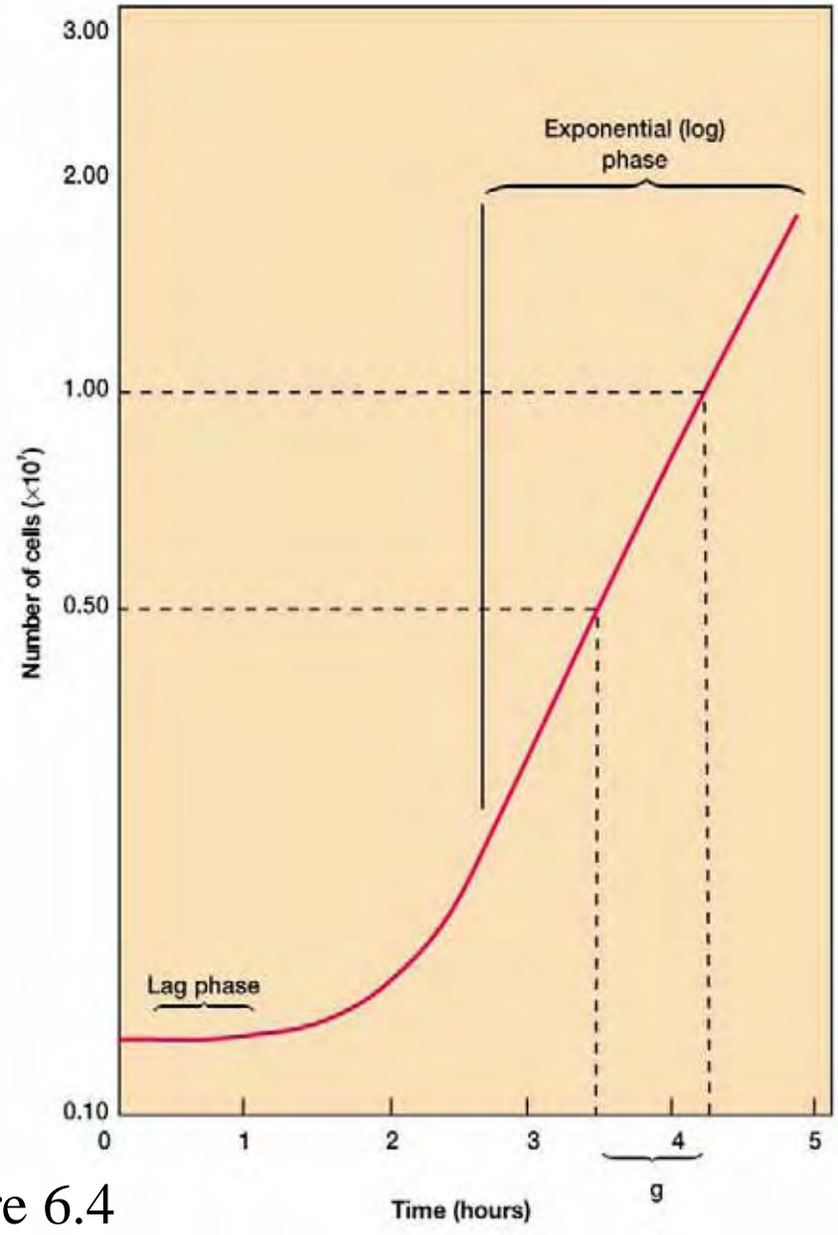


Figure 6.4

Table 6.2 Generation Times for Selected Microorganisms

Microorganism	Temperature (°C)	Generation Time (Hours)
Bacteria		
<i>Beneckeia natrigens</i>	37	0.16
<i>Escherichia coli</i>	40	0.35
<i>Bacillus subtilis</i>	40	0.43
<i>Staphylococcus aureus</i>	37	0.47
<i>Pseudomonas aeruginosa</i>	37	0.58
<i>Clostridium botulinum</i>	37	0.58
<i>Rhodospirillum rubrum</i>	25	4.6–5.3
<i>Anabaena cylindrica</i>	25	10.6
<i>Mycobacterium tuberculosis</i>	37	≈12
<i>Treponema pallidum</i>	37	33
Algae		
<i>Scenedesmus quadricauda</i>	25	5.9
<i>Chlorella pyrenoidosa</i>	25	7.75
<i>Asterionella formosa</i>	20	9.6
<i>Euglena gracilis</i>	25	10.9
<i>Ceratium tripos</i>	20	82.8
Protozoa		
<i>Tetrahymena geleii</i>	24	2.2–4.2
<i>Leishmania donovani</i>	26	10–12
<i>Paramecium caudatum</i>	26	10.4
<i>Acanthamoeba castellanii</i>	30	11–12
<i>Giardia lamblia</i>	37	18
Fungi		
<i>Saccharomyces cerevisiae</i>	30	2
<i>Monilinia fraa</i>	25	30

Measurement of Microbial Growth

- **can measure changes in number of cells in a population**
- **can measure changes in mass of population**

Measurement of Cell Numbers

- **Direct cell counts**
 - counting chambers
 - electronic counters
 - on membrane filters
- **Viable cell counts**
 - plating methods
 - membrane filtration methods

Counting chambers

- **easy, inexpensive, and quick**
- **useful for counting both eucaryotes and procaryotes**
- **cannot distinguish living from dead cells**

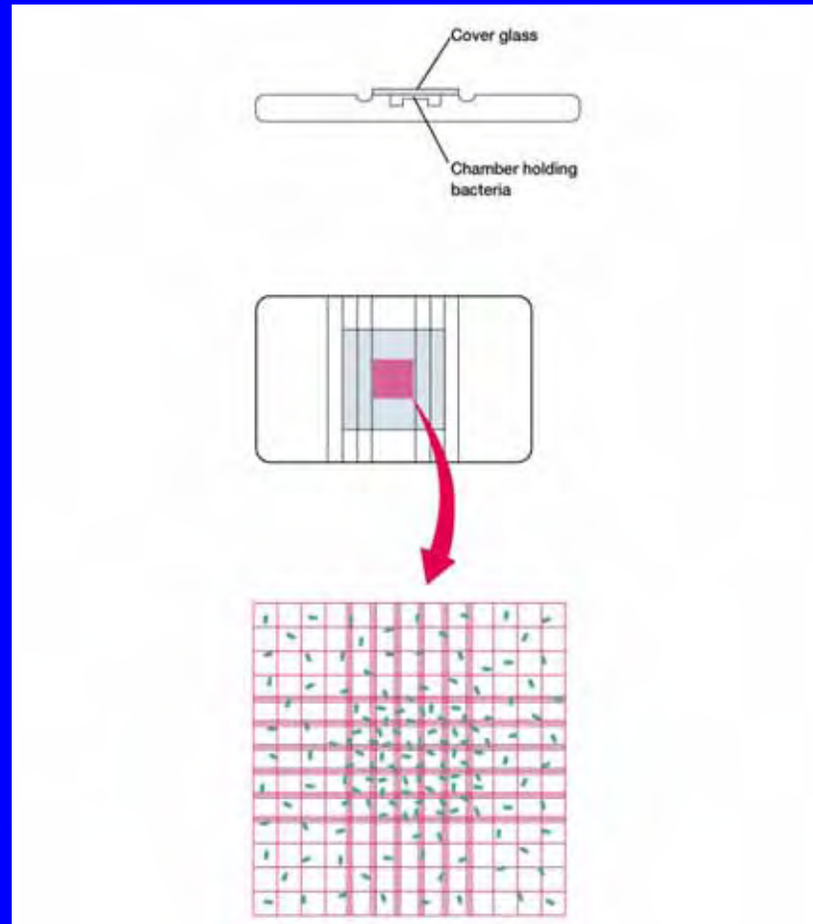


Figure 6.5

Electronic counters

- **microbial suspension forced through small orifice**
- **movement of microbe through orifice impacts electric current that flows through orifice**
- **instances of disruption of current are counted**

Electronic counters...

- **cannot distinguish living from dead cells**
- **quick and easy to use**
- **useful for large microorganisms and blood cells, but not procaryotes**

Direct counts on membrane filters

- **cells filtered through special membrane that provides dark background for observing cells**
- **cells are stained with fluorescent dyes**
- **useful for counting bacteria**
- **with certain dyes, can distinguish living from dead cells**

Plating methods

- **measure number of viable cells**
- **population size is expressed as colony forming units (CFU)**

plate dilutions of population on suitable solid medium



count number of colonies



calculate number of cells in population

Plating methods...

- **simple and sensitive**
- **widely used for viable counts of microorganisms in food, water, and soil**
- **inaccurate results obtained if cells clump together**

Membrane filtration methods

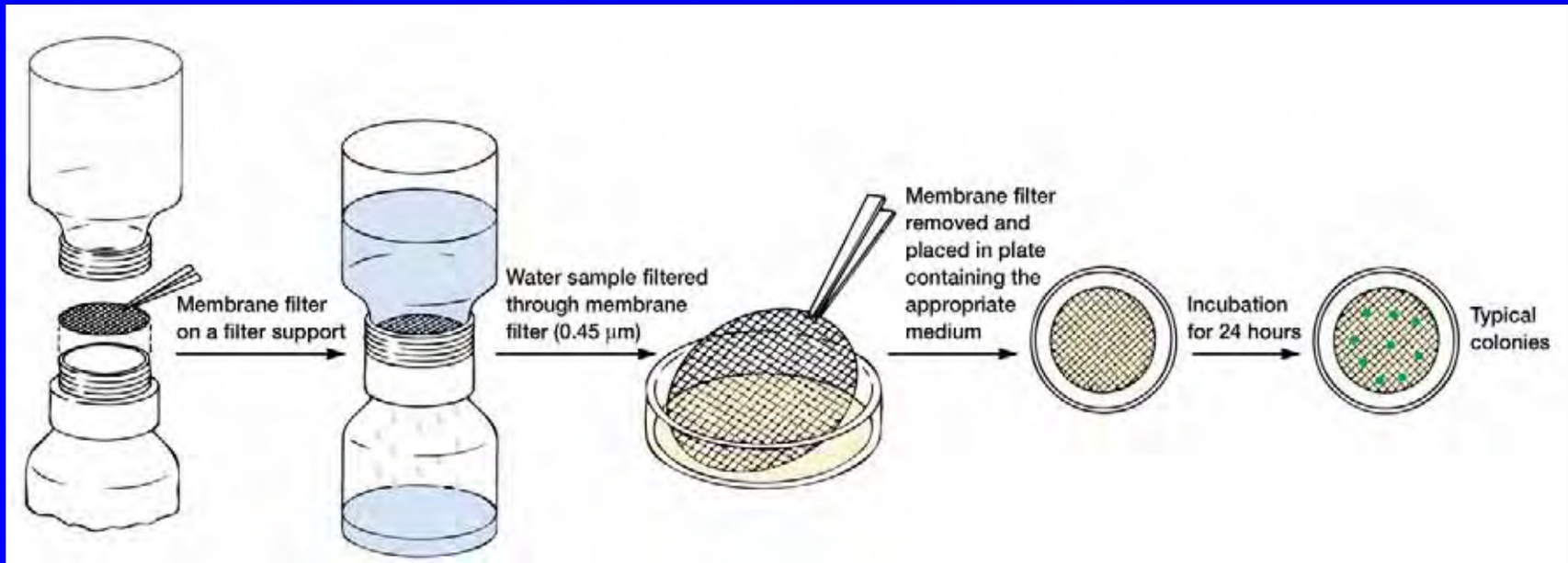
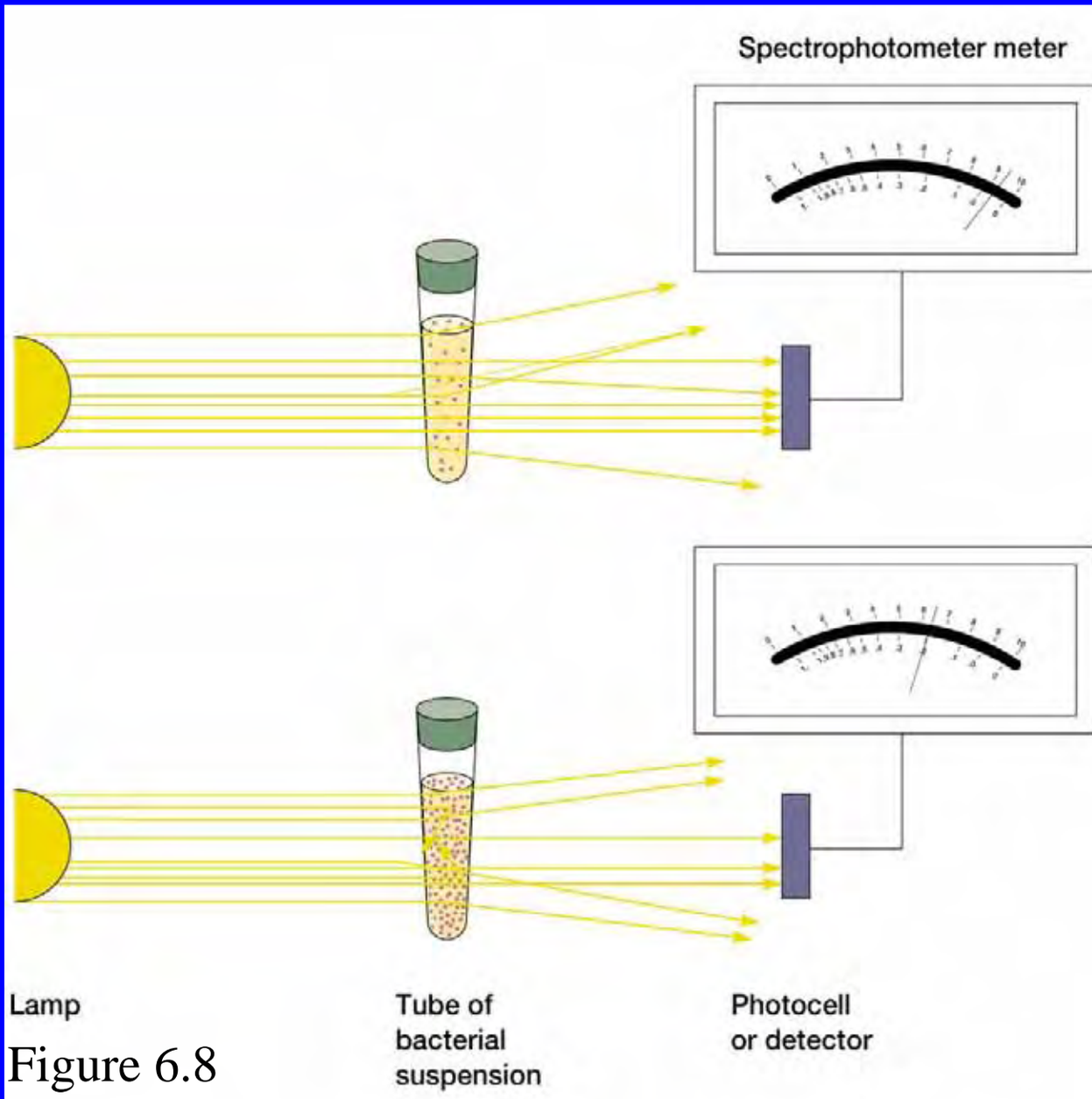


Figure 6.6

especially useful for analyzing aquatic samples

Measurement of Cell Mass

- **dry weight**
 - time consuming and not very sensitive
- **quantity of a particular cell constituent**
 - e.g., protein, DNA, ATP, or chlorophyll
 - useful if amount of substance in each cell is constant
- **turbidometric measures (light scattering)**
 - quick, easy, and sensitive



more cells
↓
more light scattered
↓
less light detected

Figure 6.8

The Continuous Culture of Microorganisms

- **growth in an open system**
 - **continual provision of nutrients**
 - **continual removal of wastes**
- **maintains cells in log phase at a constant biomass concentration for extended periods**
- **achieved using a continuous culture system**

The Chemostat

- rate of incoming medium = rate of removal of medium from vessel
- an essential nutrient is in limiting quantities

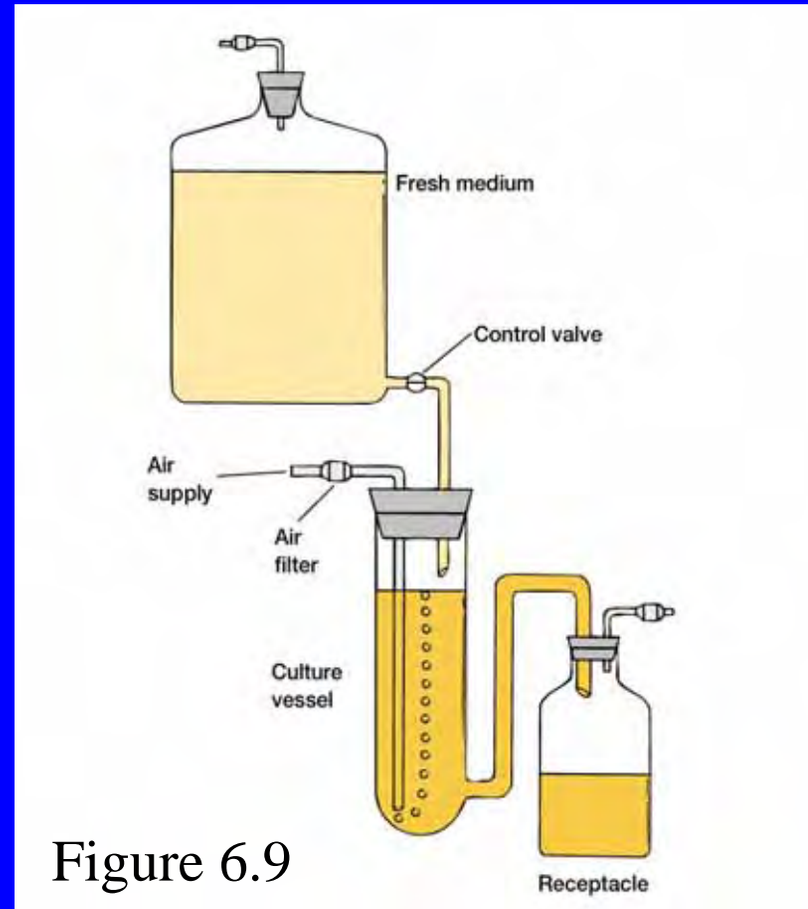


Figure 6.9

Dilution rate and microbial growth

dilution rate – rate at which medium flows through vessel relative to vessel size

note: cell density maintained at wide range of dilution rates and chemostat operates best at low dilution rate

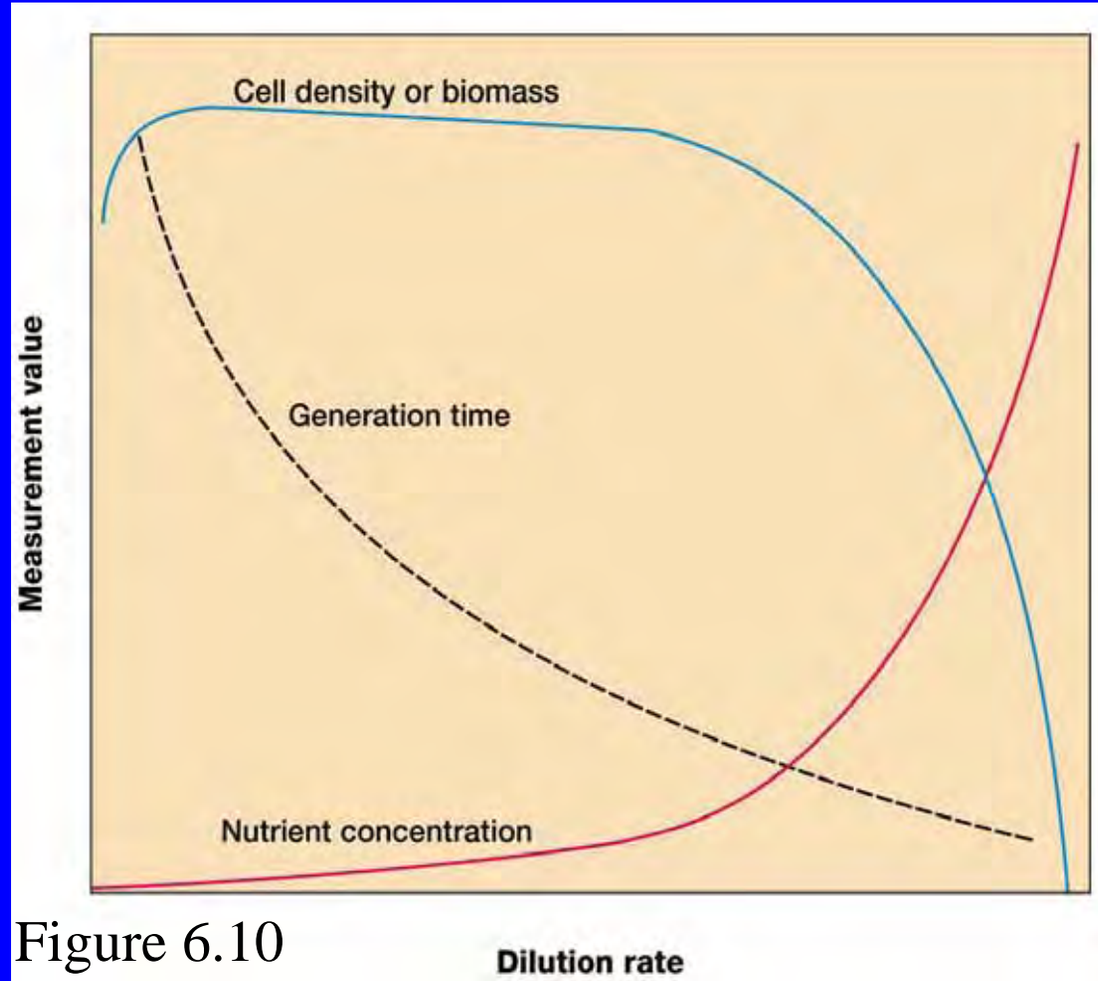


Figure 6.10

The Turbidostat

- **regulates the flow rate of media through vessel to maintain a predetermined turbidity or cell density**
- **dilution rate varies**
- **no limiting nutrient**
- **turbidostat operates best at high dilution rates**

Importance of continuous culture methods

- **constant supply of cells in exponential phase growing at a known rate**
- **study of microbial growth at very low nutrient concentrations, close to those present in natural environment**
- **study of interactions of microbes under conditions resembling those in aquatic environments**
- **food and industrial microbiology**

The Influence of Environmental Factors on Growth

- most organisms grow in fairly moderate environmental conditions
- **extremophiles**
 - grow under harsh conditions that would kill most other organisms

Solutes and Water Activity

- **water activity (a_w)**
 - amount of water available to organisms
 - reduced by interaction with solute molecules (osmotic effect)
 - higher [solute] \Rightarrow lower a_w
 - reduced by adsorption to surfaces (matric effect)

Table 6.4 Approximate Lower a_w Limits for Microbial Growth

Water Activity	Environment	Bacteria	Fungi	Algae
1.00—Pure water	Blood Plant wilt Seawater	Most gram-negative nonhalophiles		
	{ Vegetables, meat, fruit			
0.95	Bread	Most gram-positive rods	<i>Basidiomycetes</i>	Most algae
0.90	Ham	Most cocci, <i>Bacillus</i>	<i>Fusarium</i> <i>Mucor</i> , <i>Rhizopus</i> Ascomycetous yeasts	
0.85	Salami	<i>Staphylococcus</i>	<i>Saccharomyces rouxii</i> (in salt)	
0.80	Preserves		<i>Penicillium</i>	
0.75	Salt lakes Salted fish	<i>Halobacterium</i> <i>Actinospora</i>	<i>Aspergillus</i>	<i>Dunaliella</i>
0.70			<i>Aspergillus</i>	
0.60	Cereals, candy, dried fruit		<i>Saccharomyces rouxii</i> (in sugars)	
	Chocolate Honey Dried milk		<i>Xeromyces bisporus</i>	
0.55—DNA disordered				

Adapted from A. D. Brown, "Microbial Water Stress," in *Bacteriological Reviews*, 40(4):803–846 1976. Copyright ©1976 by the American Society for Microbiology. Reprinted by permission.

Osmotolerant organisms

- grow over wide ranges of water activity
- many use **compatible solutes** to increase their internal osmotic concentration
 - solutes that are compatible with metabolism and growth
- some have proteins and membranes that require high solute concentrations for stability and activity

Effects of NaCl on microbial growth

- **halophiles**
 - grow optimally at >0.2 M
- **extreme halophiles**
 - require >2 M

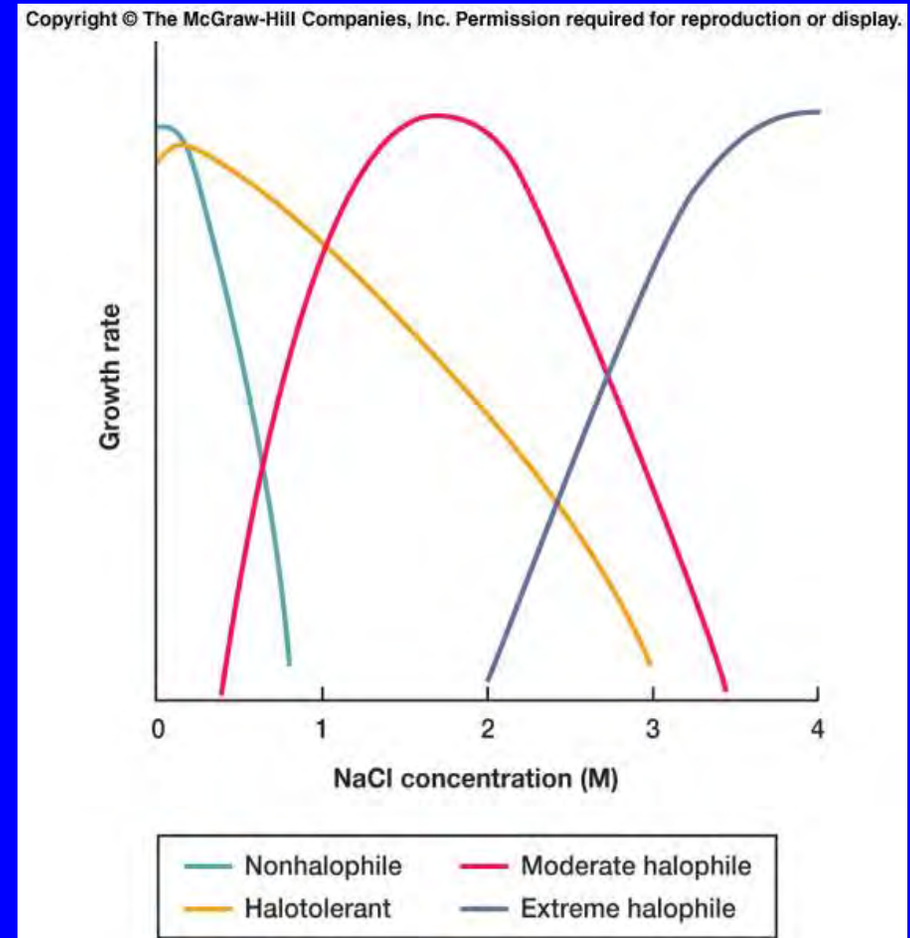


Figure 6.11

pH

- negative logarithm of the hydrogen ion concentration

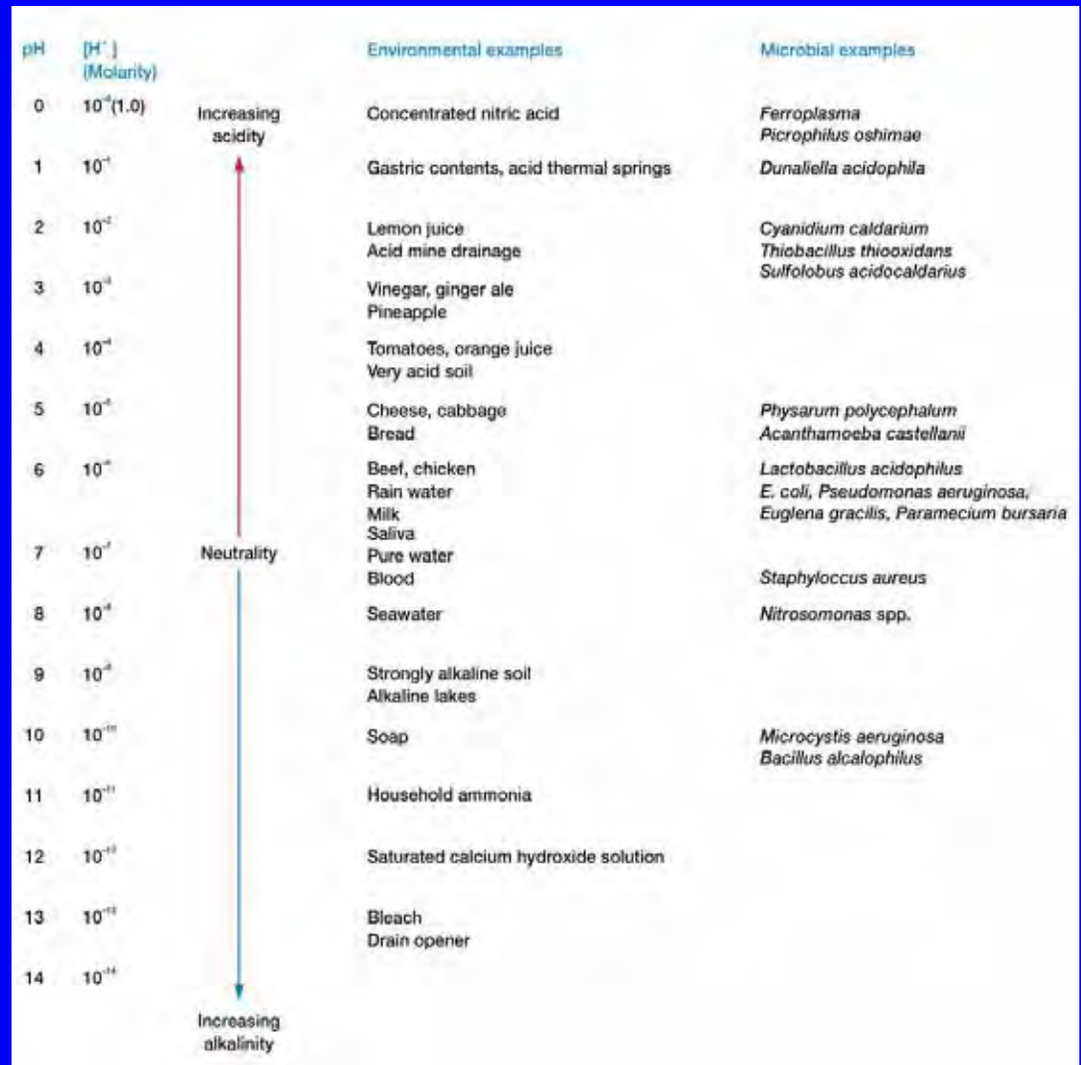


Figure 6.12

pH

- **acidophiles**
 - growth optimum between pH 0 and pH 5.5
- **neutrophiles**
 - growth optimum between pH 5.5 and pH 7
- **alkalophiles**
 - growth optimum between pH 8.5 and pH 11.5

pH

- **most acidophiles and alkalophiles maintain an internal pH near neutrality**
 - some use proton/ion exchange mechanisms to do so
- **some synthesize proteins that provide protection**
 - e.g., acid-shock proteins
- **many microorganisms change pH of their habitat by producing acidic or basic waste products**
 - most media contain buffers to prevent growth inhibition

Temperature

- organisms exhibit distinct cardinal growth temperatures
 - minimal
 - maximal
 - optimal

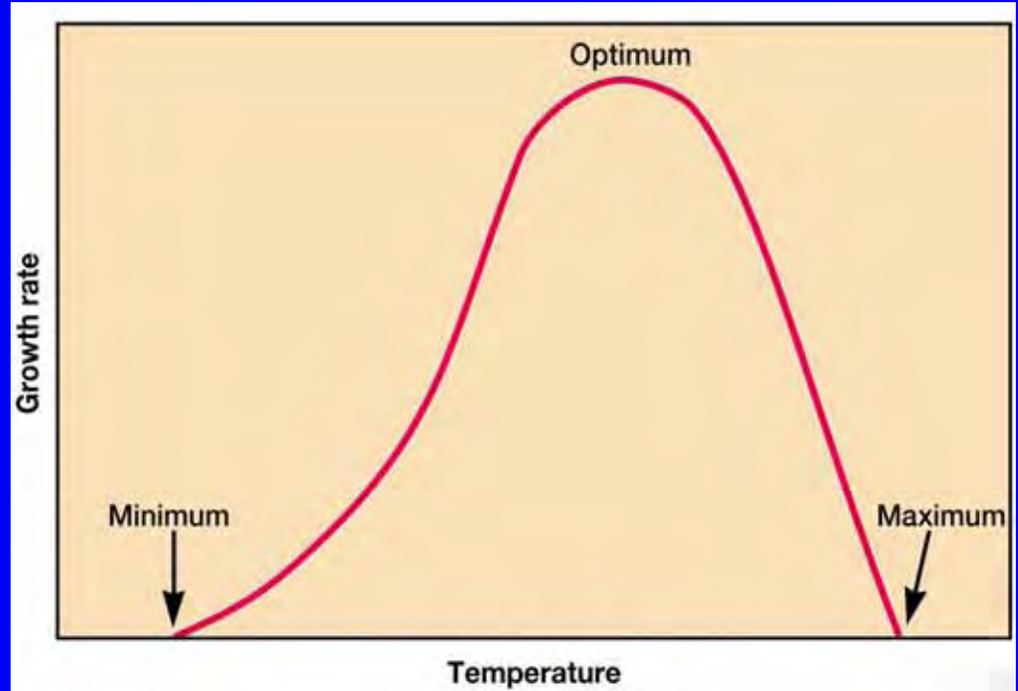


Figure 6.13

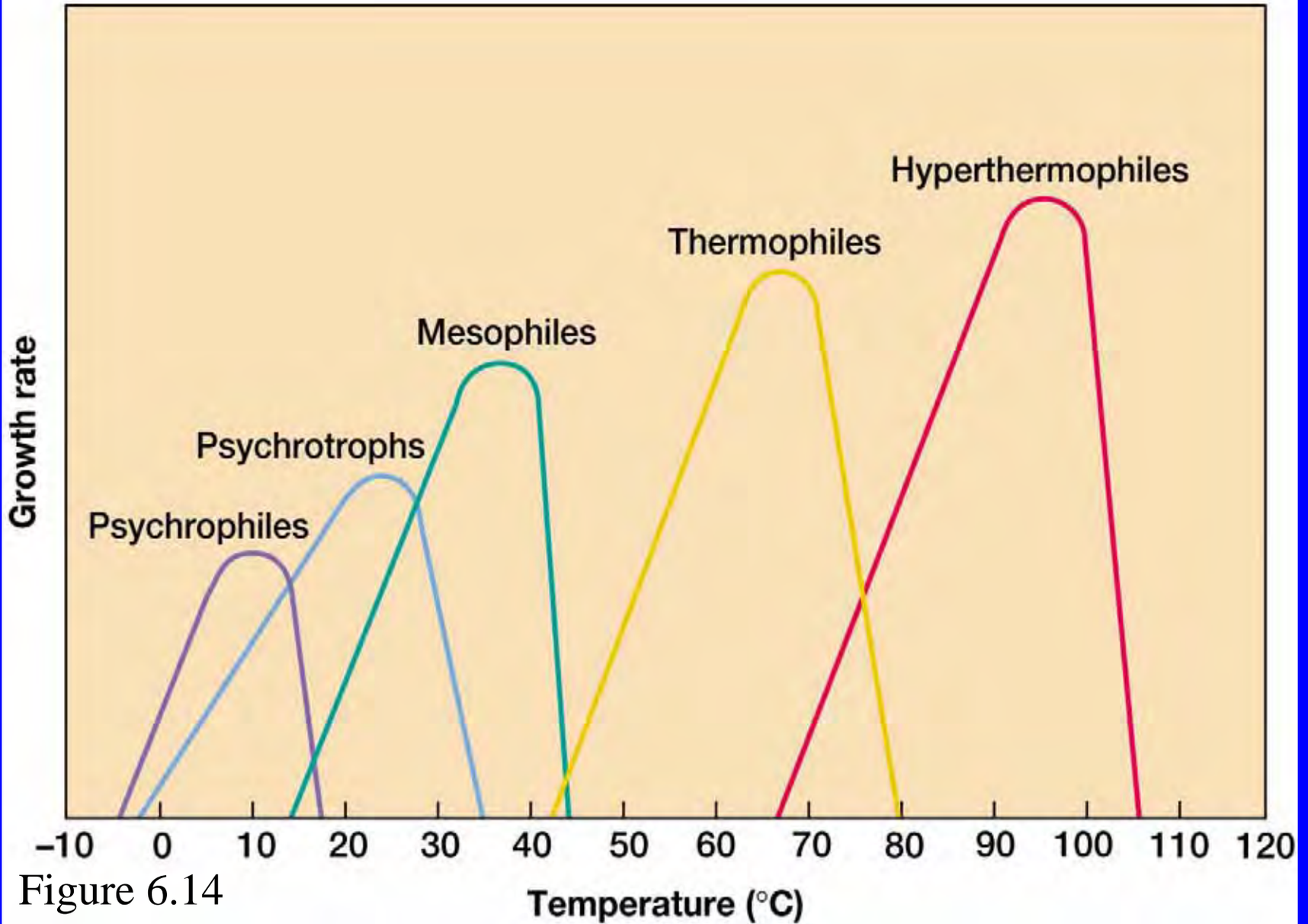


Figure 6.14

Adaptations of thermophiles

- **protein structure stabilized by a variety of means**
 - e.g., more H bonds
 - e.g., more proline
 - e.g., chaperones
- **histone-like proteins stabilize DNA**
- **membrane stabilized by variety of means**
 - e.g., more saturated, more branched and higher molecular weight lipids
 - e.g., ether linkages (archaeal membranes)

Oxygen Concentration

need
oxygen

prefer
oxygen

ignore
oxygen

oxygen is
toxic

< 2 – 10%
oxygen

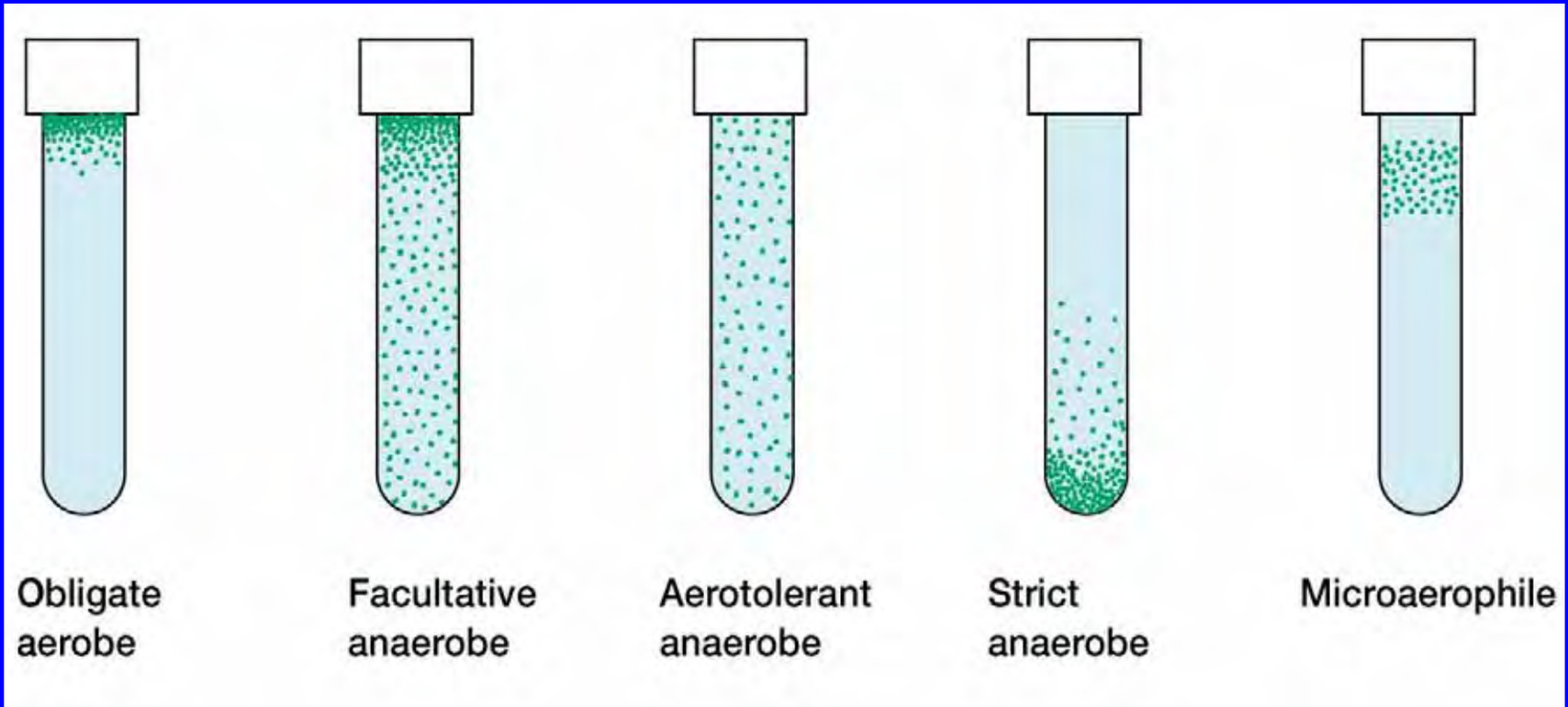


Figure 6.15

Basis of different oxygen sensitivities

- oxygen easily reduced to toxic products
 - superoxide radical
 - hydrogen peroxide
 - hydroxyl radical
- aerobes produce protective enzymes
 - superoxide dismutase (SOD)
 - catalase

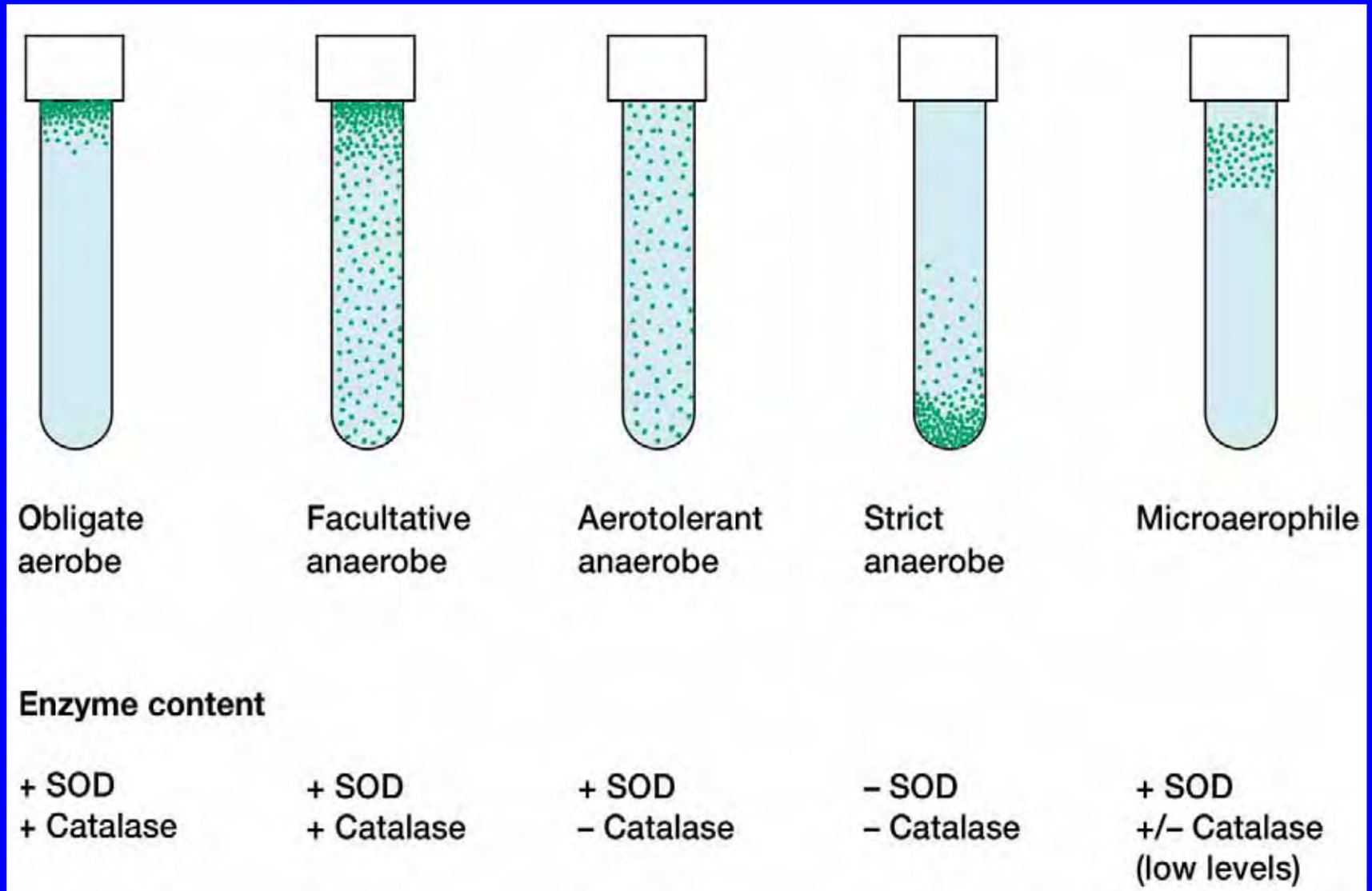


Figure 6.14

Pressure

- **barotolerant organisms**
 - adversely affected by increased pressure, but not as severely as nontolerant organisms
- **barophilic organisms**
 - require or grow more rapidly in the presence of increased pressure

Radiation

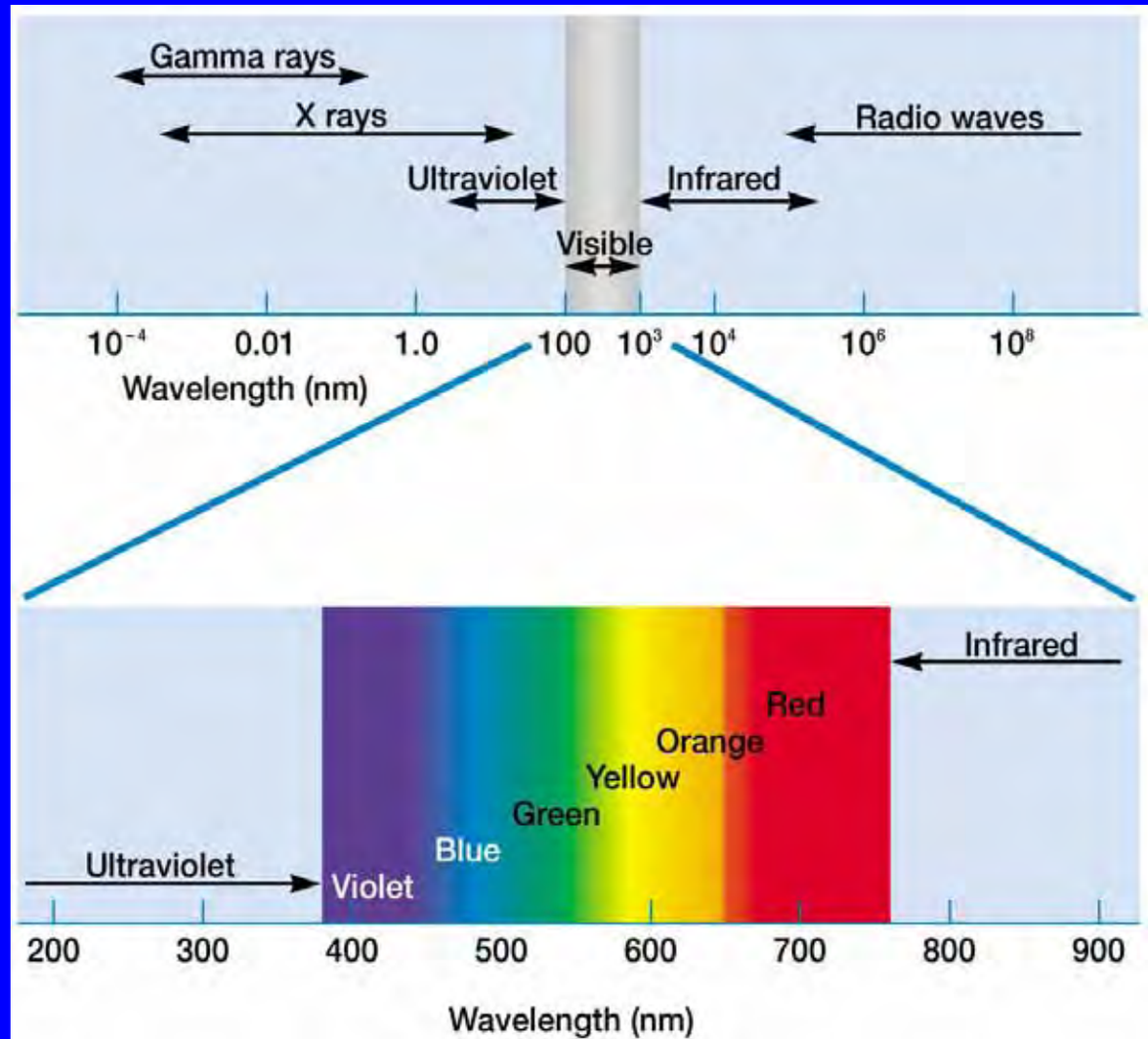


Figure 6.18

Radiation damage

- **ionizing radiation**
 - **x rays and gamma rays**
 - **mutations → death**
 - **disrupts chemical structure of many molecules, including DNA**
 - **damage may be repaired by DNA repair mechanisms**

Radiation damage...

- **ultraviolet (UV) radiation**
 - mutations → death
 - causes formation of thymine dimers in DNA
 - DNA damage can be repaired by two mechanisms
 - **photoreactivation** – dimers split in presence of light
 - **dark reactivation** – dimers excised and replaced in absence of light

Radiation damage...

- **visible light**
 - at high intensities generates **singlet oxygen** ($^1\text{O}_2$)
 - powerful oxidizing agent
 - **carotenoid pigments**
 - protect many light-exposed microorganisms from photooxidation

Microbial Growth in Natural Environments

- **microbial environments are complex, constantly changing, and may expose a microorganism to overlapping gradients of nutrients and environmental factors**

Growth Limitation by Environmental Factors

- **Leibig's law of the minimum**
 - total biomass of organism determined by nutrient present at lowest concentration
- **Shelford's law of tolerance**
 - above or below certain environmental limits, a microorganism will not grow, regardless of the nutrient supply

Responses to low nutrient levels

- **oligotrophic environments**
- **morphological changes to increase surface area and ability to absorb nutrients**
- **mechanisms to sequester certain nutrients**

Counting Viable but Nonculturable Vegetative Procaryotes

- **stressed microorganisms can temporarily lose ability to grow using normal cultivation methods**
- **microscopic and isotopic methods for counting viable but nonculturable cells have been developed**

Quorum Sensing and Microbial Populations

- **quorum sensing**
 - **microbial communication and cooperation**
 - **involves secretion and detection of chemical signals**

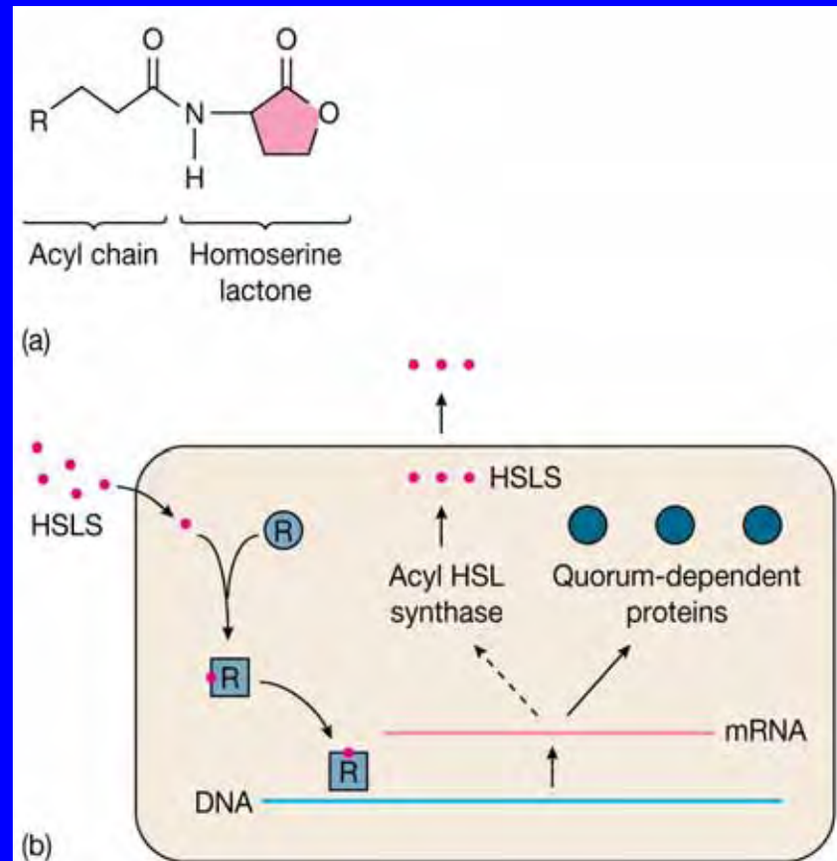


Figure 6.20

Processes sensitive to quorum sensing: gram-negative bacteria

- bioluminescence (*Vibrio fischeri*)
- synthesis and release of virulence factors (*Pseudomonas aeruginosa*)
- conjugation (*Agrobacterium tumefaciens*)
- antibiotic production (*Erwinia carotovora*, *Pseudomonas aureofaciens*)
- biofilm production (*P. aeruginosa*)

Quorum sensing: gram-positive bacteria

- often mediated by oligopeptide pheromone
- processes impacted by quorum sensing:
 - mating (*Enterococcus faecalis*)
 - transformation competence (*Streptococcus pneumoniae*)
 - sporulation (*Bacillus subtilis*)
 - production of virulence factors (*Staphylococcus aureus*)
 - development of aerial mycelia (*Streptomyces griseus*)
 - antibiotic production (*S. griseus*)