LECTURE PRESENTATIONS For CAMPBELL BIOLOGY, NINTH EDITION Jane B. Reece, Lisa A. Urry, Michael L. Cain, Steven A. Wasserman, Peter V. Minorsky, Robert B. Jackson

Chapter 12

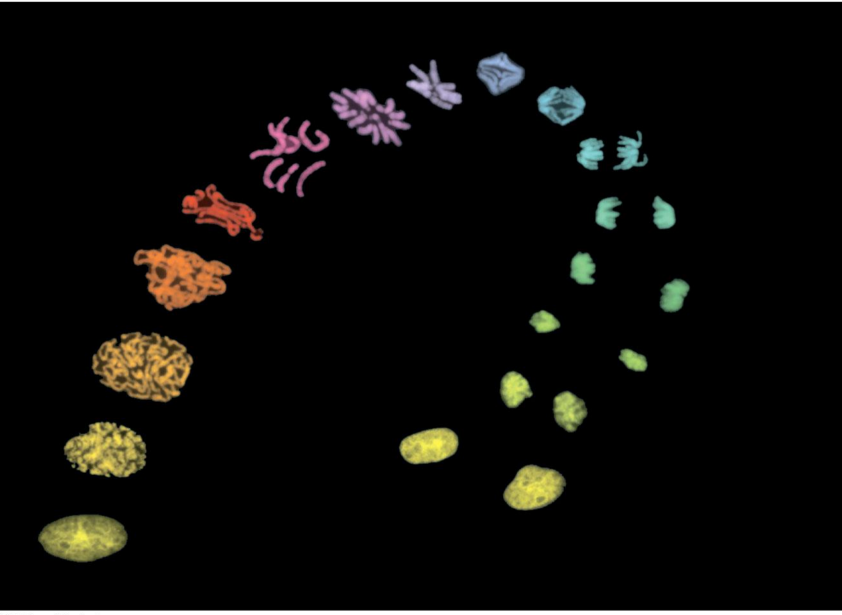
The Cell Cycle

Lectures by Erin Barley Kathleen Fitzpatrick

Overview: The Key Roles of Cell Division

- The ability of organisms to produce more of their own kind best distinguishes living things from nonliving matter
- The continuity of life is based on the reproduction of cells, or cell division

Figure 12.1



- In unicellular organisms, division of one cell reproduces the entire organism
- Multicellular organisms depend on cell division for
 - Development from a fertilized cell
 - Growth
 - Repair
- Cell division is an integral part of the cell cycle, the life of a cell from formation to its own division

Figure 12.2

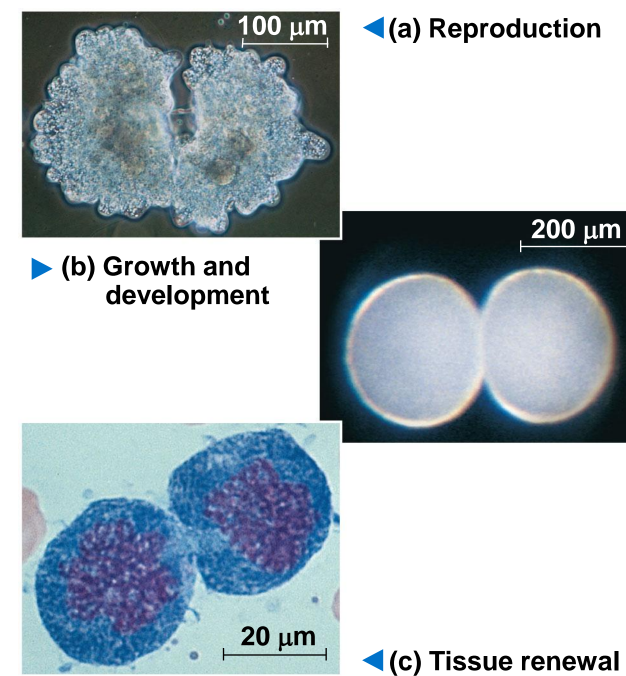
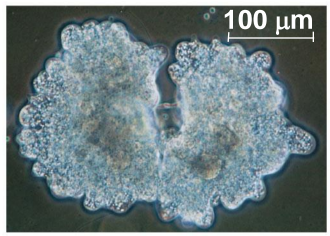


Figure 12.2a



(a) Reproduction



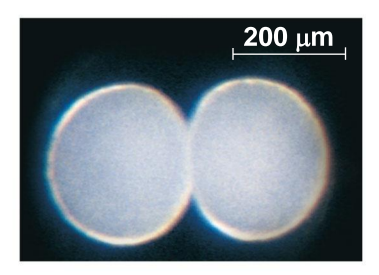
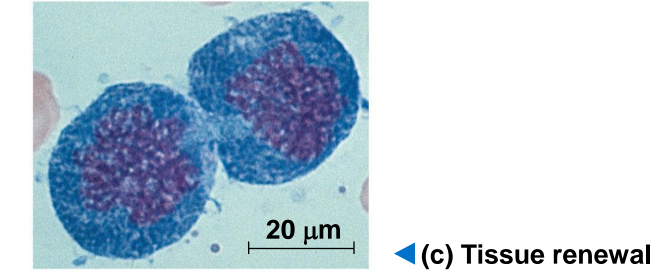


Figure 12.2c



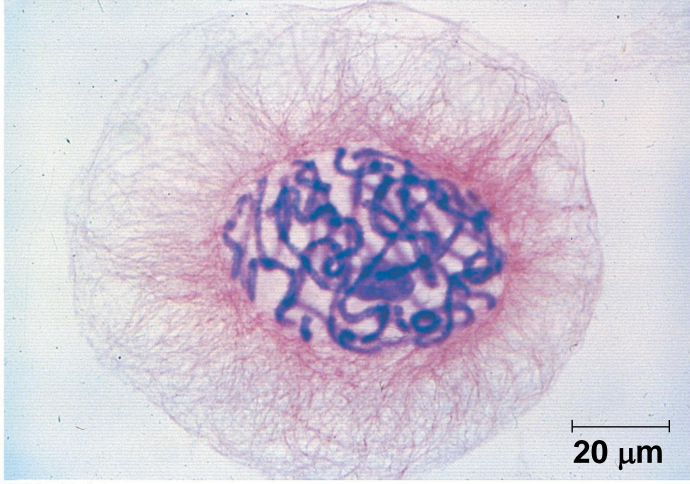
Concept 12.1: Most cell division results in genetically identical daughter cells

- Most cell division results in daughter cells with identical genetic information, DNA
- The exception is meiosis, a special type of division that can produce sperm and egg cells

Cellular Organization of the Genetic Material

- All the DNA in a cell constitutes the cell's genome
- A genome can consist of a single DNA molecule (common in prokaryotic cells) or a number of DNA molecules (common in eukaryotic cells)
- DNA molecules in a cell are packaged into chromosomes

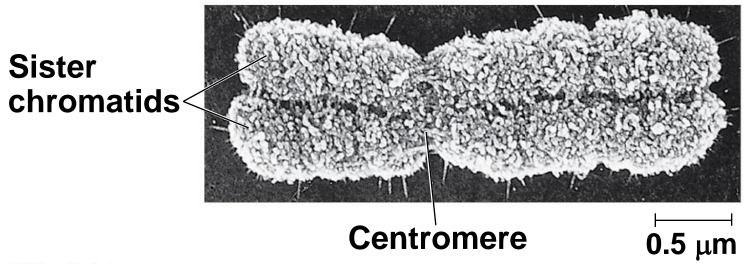
Figure 12.3



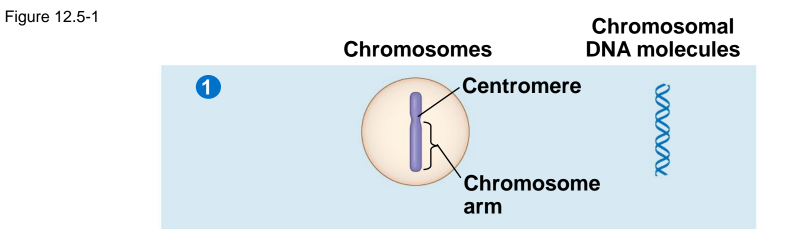
- Eukaryotic chromosomes consist of chromatin, a complex of DNA and protein that condenses during cell division
- Every eukaryotic species has a characteristic number of chromosomes in each cell nucleus
- Somatic cells (nonreproductive cells) have two sets of chromosomes
- **Gametes** (reproductive cells: sperm and eggs) have half as many chromosomes as somatic cells

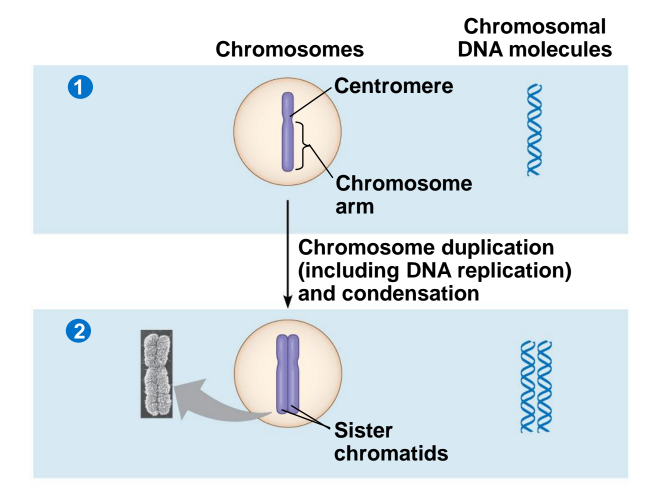
Distribution of Chromosomes During Eukaryotic Cell Division

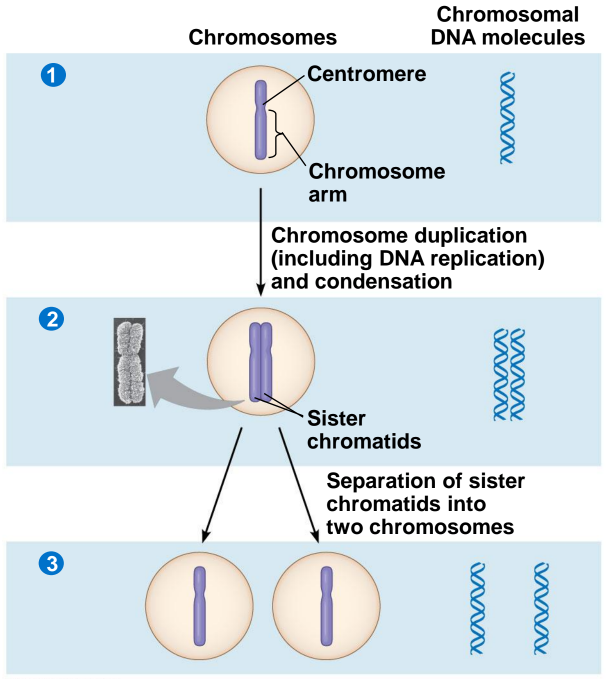
- In preparation for cell division, DNA is replicated and the chromosomes condense
- Each duplicated chromosome has two sister chromatids (joined copies of the original chromosome), which separate during cell division
- The centromere is the narrow "waist" of the duplicated chromosome, where the two chromatids are most closely attached



- During cell division, the two sister chromatids of each duplicated chromosome separate and move into two nuclei
- Once separate, the chromatids are called chromosomes







- Eukaryotic cell division consists of
 - Mitosis, the division of the genetic material in the nucleus
 - Cytokinesis, the division of the cytoplasm
- Gametes are produced by a variation of cell division called meiosis
- Meiosis yields nonidentical daughter cells that have only one set of chromosomes, half as many as the parent cell

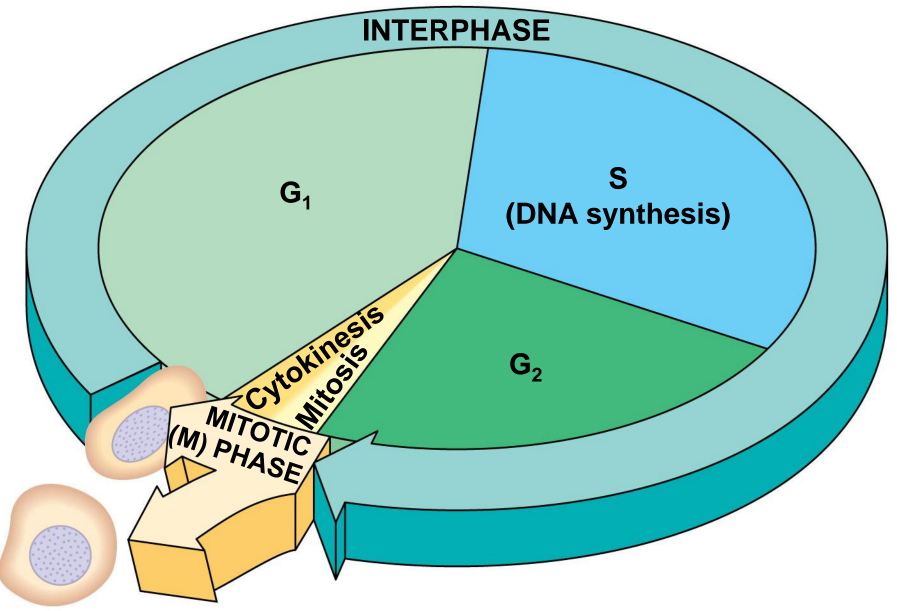
Concept 12.2: The mitotic phase alternates with interphase in the cell cycle

 In 1882, the German anatomist Walther Flemming developed dyes to observe chromosomes during mitosis and cytokinesis

Phases of the Cell Cycle

- The cell cycle consists of
 - Mitotic (M) phase (mitosis and cytokinesis)
 - Interphase (cell growth and copying of chromosomes in preparation for cell division)

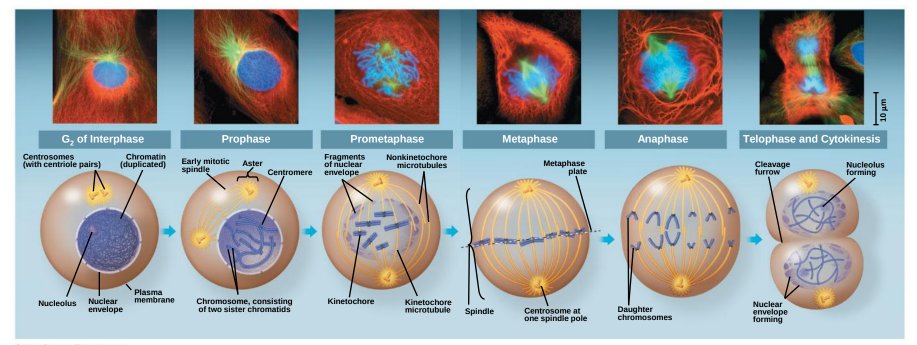
- Interphase (about 90% of the cell cycle) can be divided into subphases
 - G₁ phase ("first gap")
 - S phase ("synthesis")
 - G₂ phase ("second gap")
- The cell grows during all three phases, but chromosomes are duplicated only during the S phase

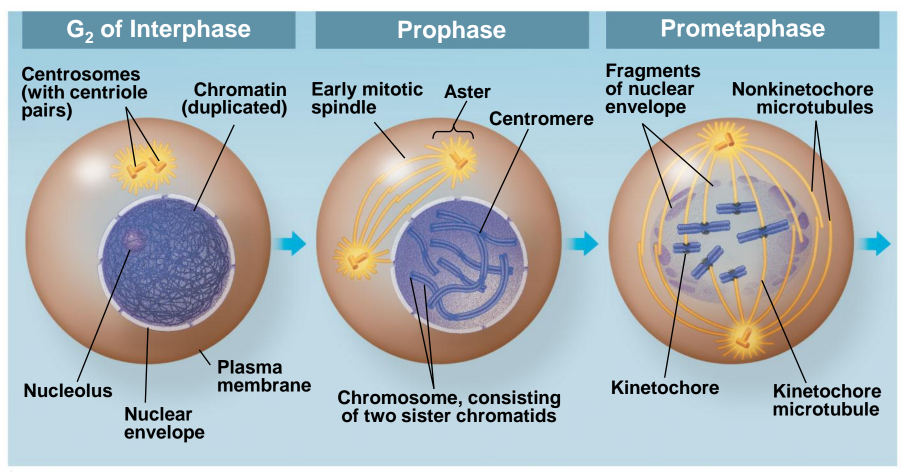


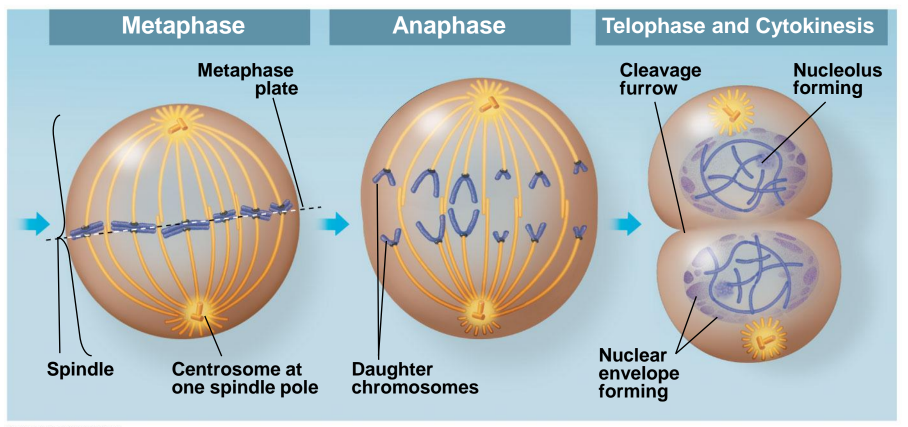
• Mitosis is conventionally divided into five phases

- Prophase
- Prometaphase
- Metaphase
- Anaphase
- Telophase
- Cytokinesis overlaps the latter stages of mitosis











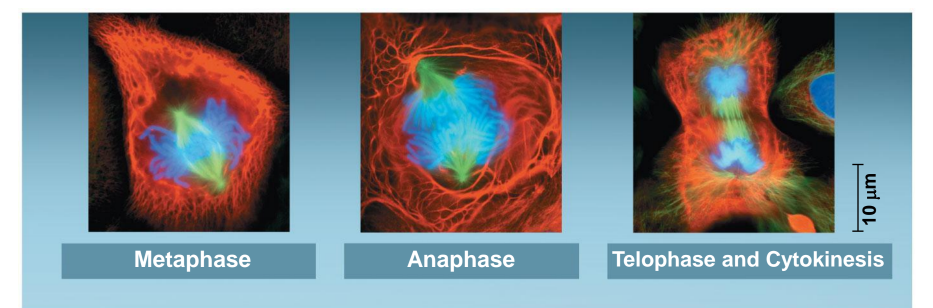
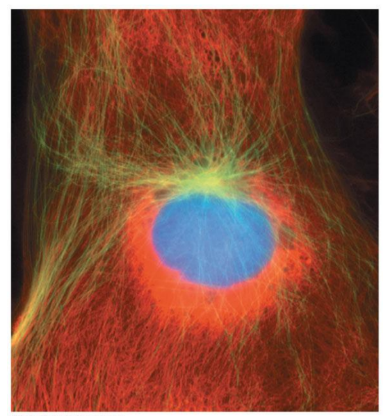
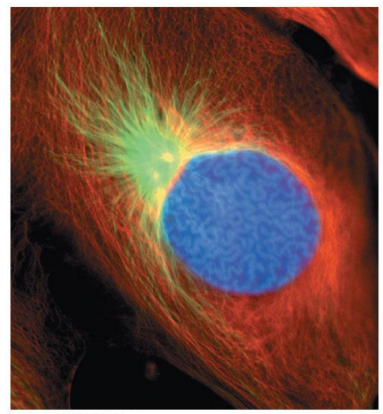


Figure 12.7e



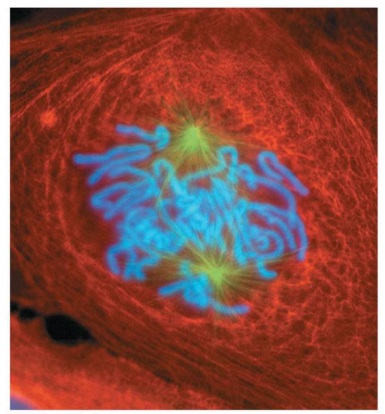
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Figure 12.7f



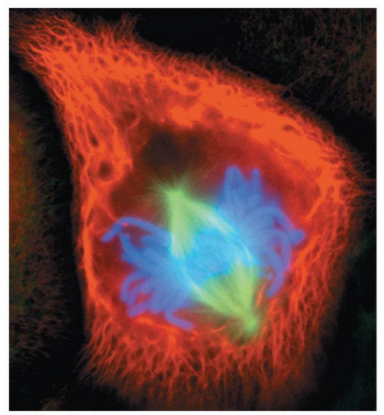
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Figure 12.7g



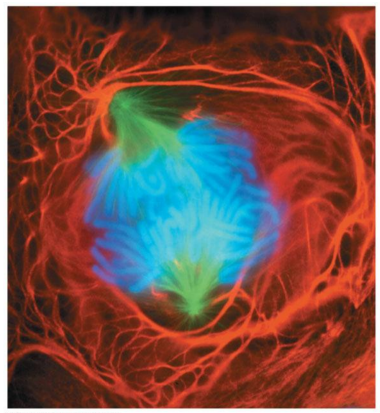
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Figure 12.7h



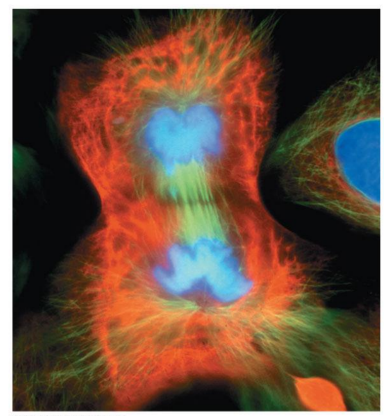
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Figure 12.7i



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Figure 12.7j



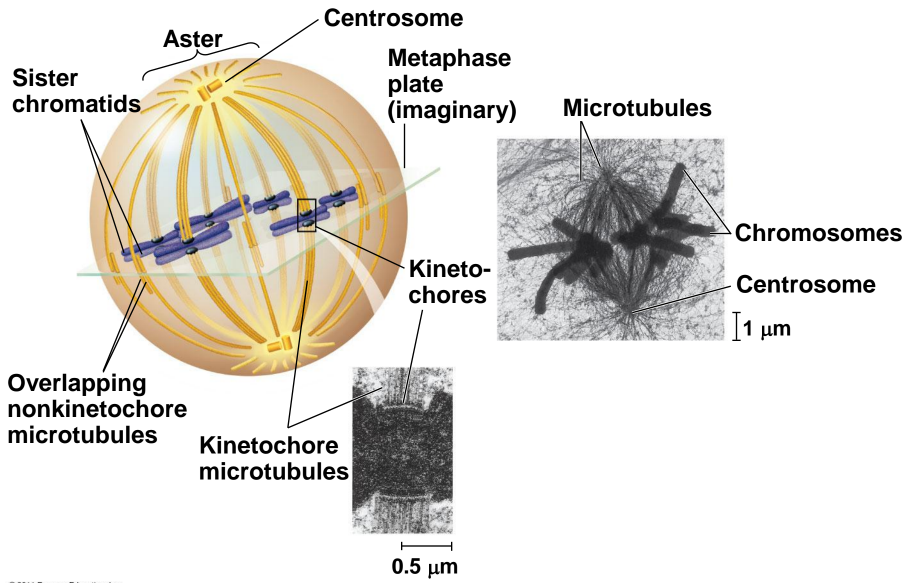
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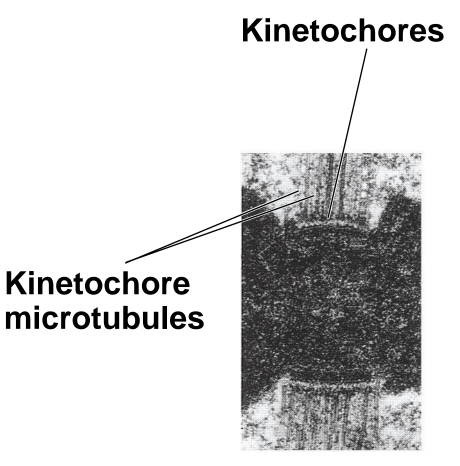
The Mitotic Spindle: A Closer Look

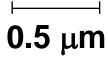
- The mitotic spindle is a structure made of microtubules that controls chromosome movement during mitosis
- In animal cells, assembly of spindle microtubules begins in the centrosome, the microtubule organizing center
- The centrosome replicates during interphase, forming two centrosomes that migrate to opposite ends of the cell during prophase and prometaphase

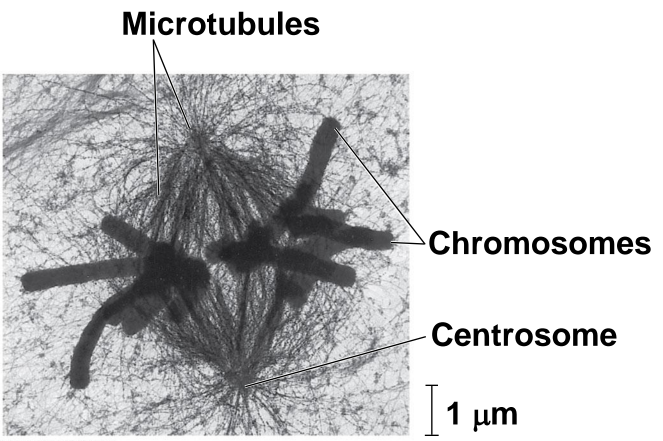
- An aster (a radial array of short microtubules) extends from each centrosome
- The spindle includes the centrosomes, the spindle microtubules, and the asters

- During prometaphase, some spindle microtubules attach to the kinetochores of chromosomes and begin to move the chromosomes
- Kinetochores are protein complexes associated with centromeres
- At metaphase, the chromosomes are all lined up at the metaphase plate, an imaginary structure at the midway point between the spindle's two poles





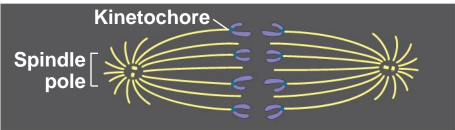


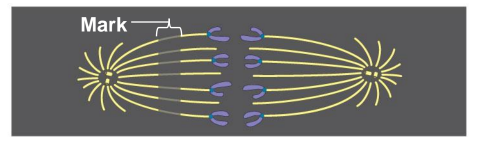


- In anaphase, sister chromatids separate and move along the kinetochore microtubules toward opposite ends of the cell
- The microtubules shorten by depolymerizing at their kinetochore ends

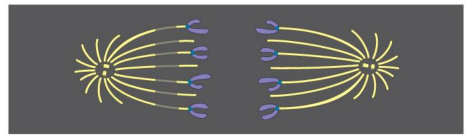
Figure 12.9

EXPERIMENT





RESULTS



CONCLUSION

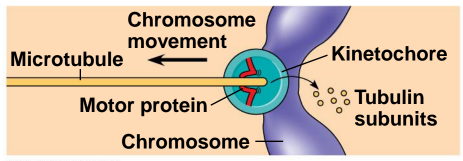
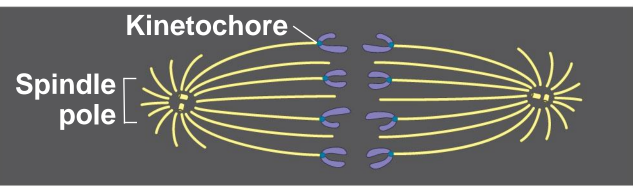
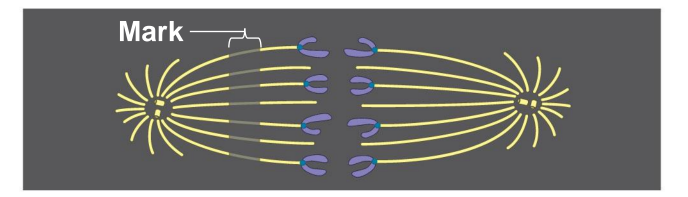


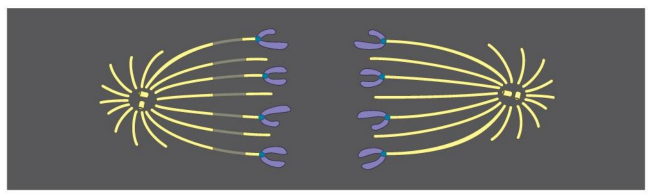
Figure 12.9a

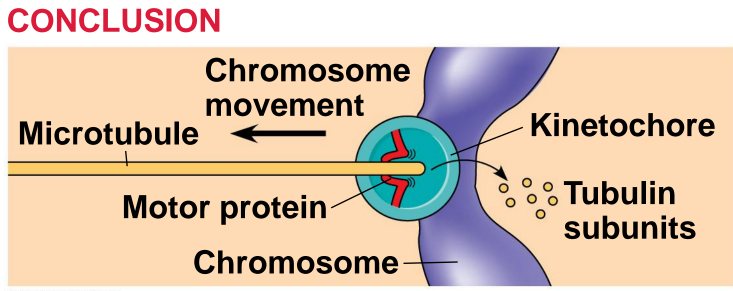
EXPERIMENT





RESULTS



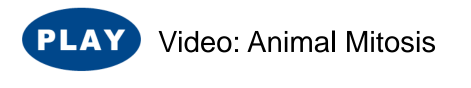


- Nonkinetochore microtubules from opposite poles overlap and push against each other, elongating the cell
- In telophase, genetically identical daughter nuclei form at opposite ends of the cell
- Cytokinesis begins during anaphase or telophase and the spindle eventually disassembles

Cytokinesis: A Closer Look

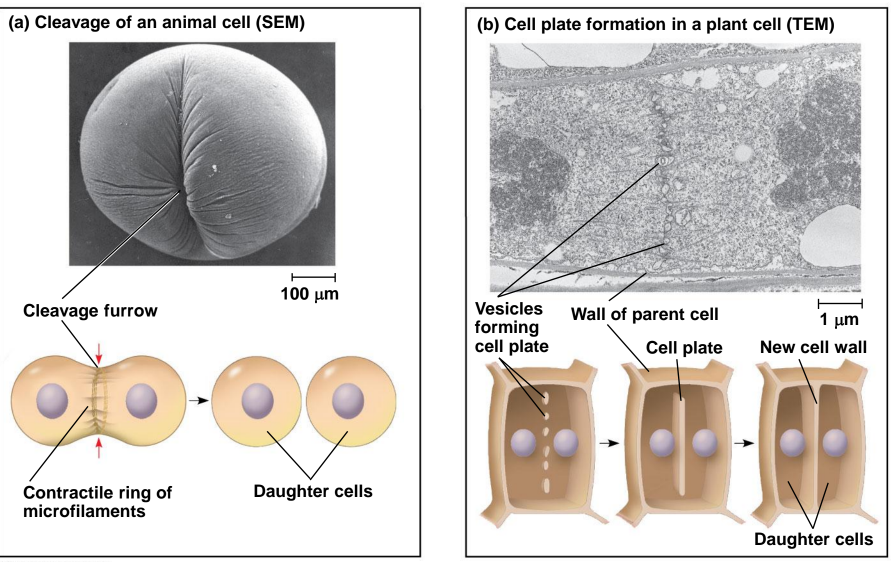
- In animal cells, cytokinesis occurs by a process known as cleavage, forming a cleavage furrow
- In plant cells, a cell plate forms during cytokinesis







PLAY Video: Sea Urchin (Time Lapse)



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Figure 12.10a

(a) Cleavage of an animal cell (SEM)

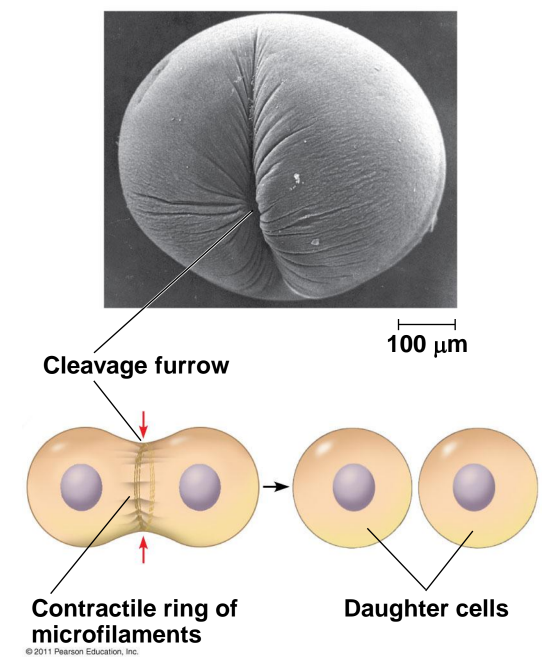
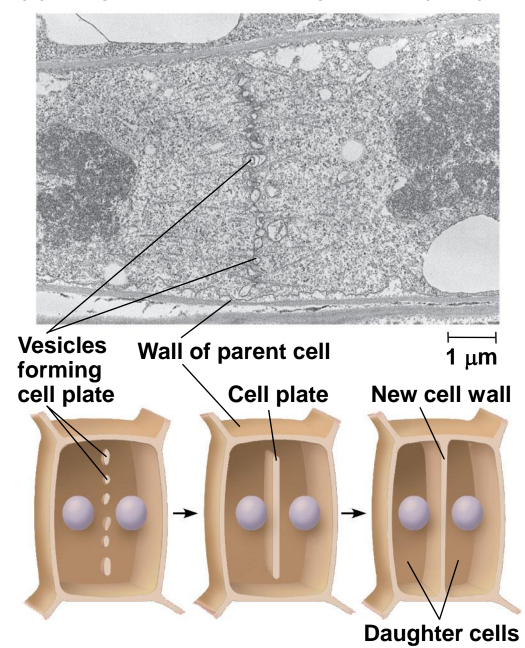


Figure 12.10b

(b) Cell plate formation in a plant cell (TEM)



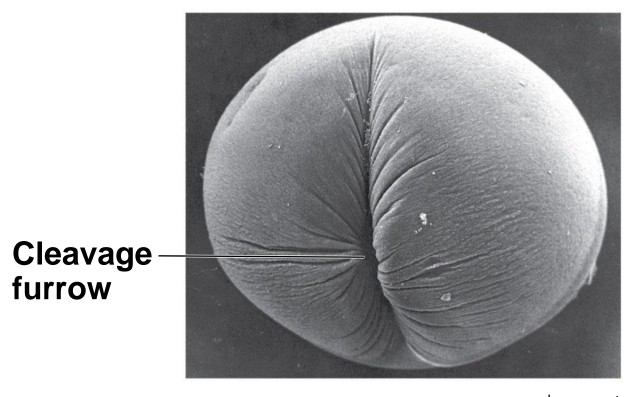
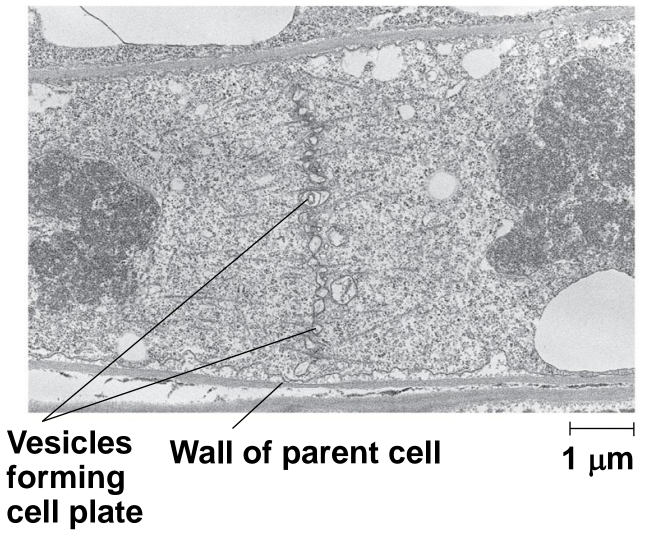
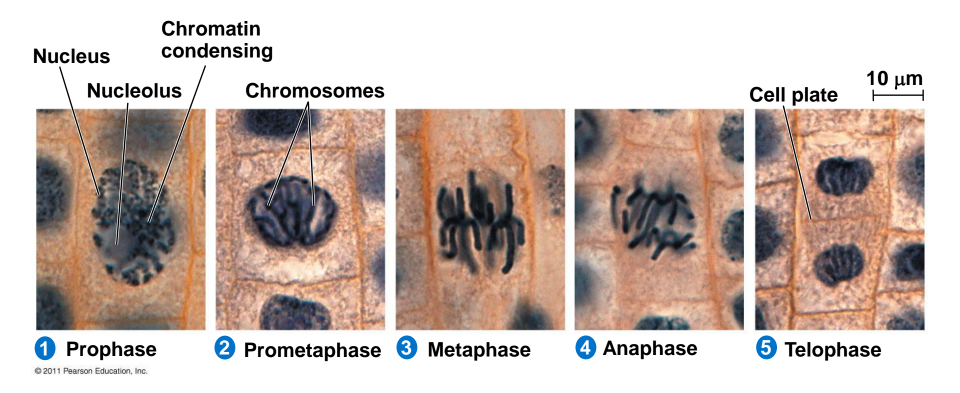
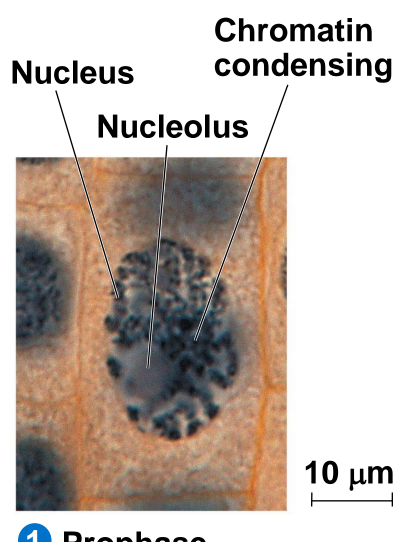




Figure 12.10d









Chromosomes

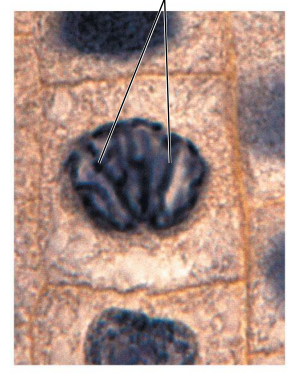






Figure 12.11c

10 µm



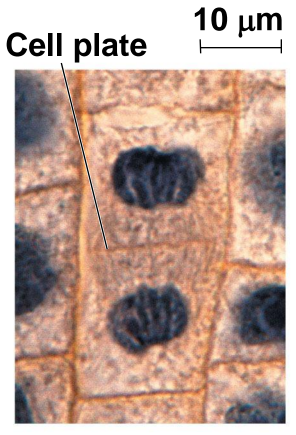


Figure 12.11d

10 μm





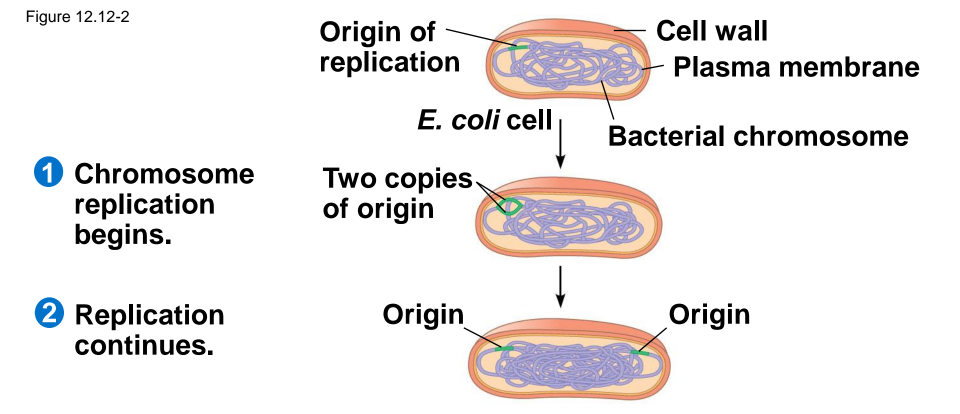


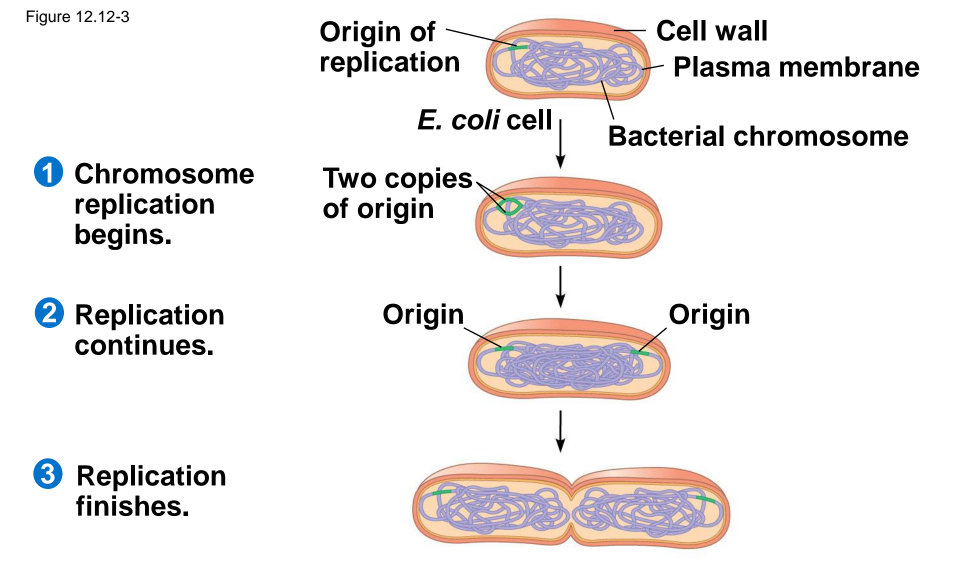


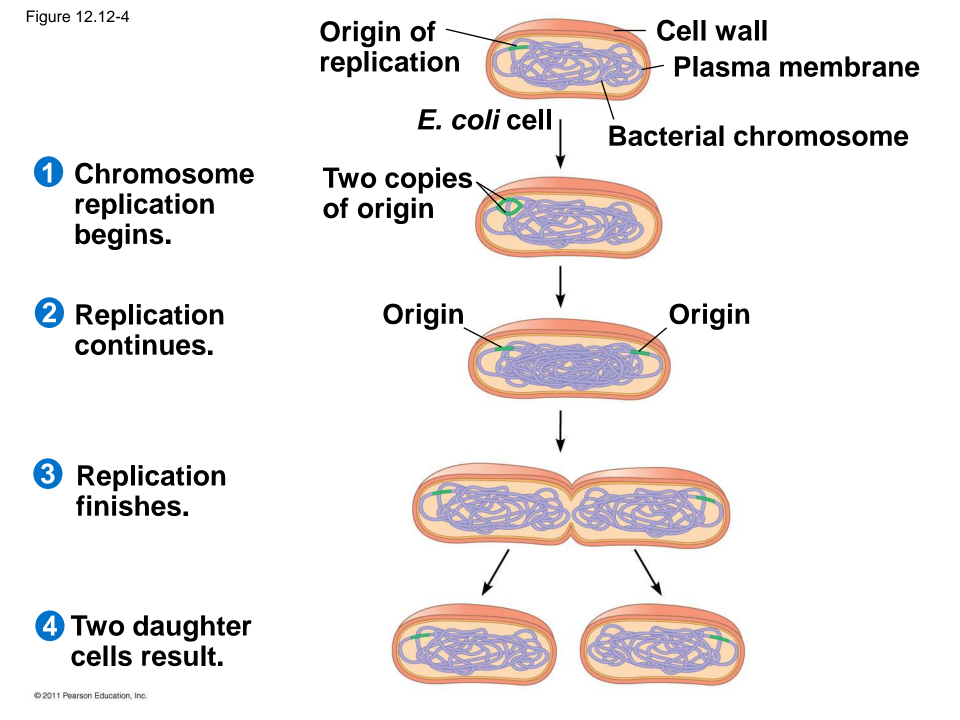
Binary Fission in Bacteria

- Prokaryotes (bacteria and archaea) reproduce by a type of cell division called binary fission
- In binary fission, the chromosome replicates (beginning at the origin of replication), and the two daughter chromosomes actively move apart
- The plasma membrane pinches inward, dividing the cell into two

Figure 12.12-1
Origin of cell wall Plasma membrane
E. coli cell
Bacterial chromosome
Two copies of origin







The Evolution of Mitosis

- Since prokaryotes evolved before eukaryotes, mitosis probably evolved from binary fission
- Certain protists exhibit types of cell division that seem intermediate between binary fission and mitosis

Figure 12.13

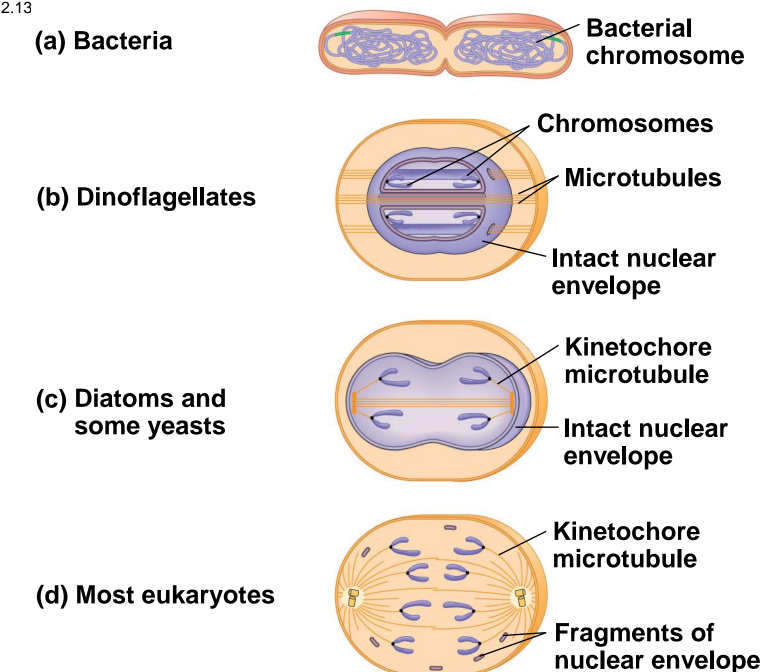
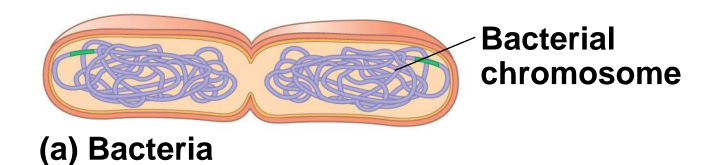


Figure 12.13a



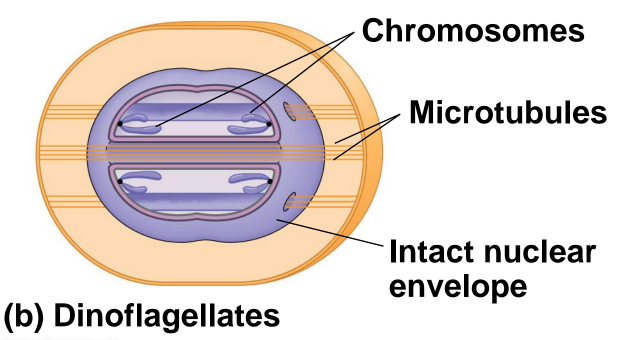
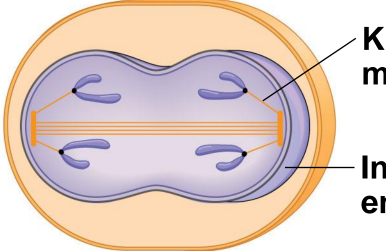


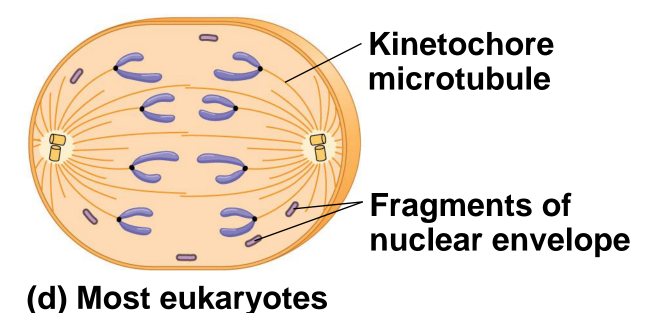
Figure 12.13b



 Kinetochore microtubule

Intact nuclear envelope

(c) Diatoms and some yeasts

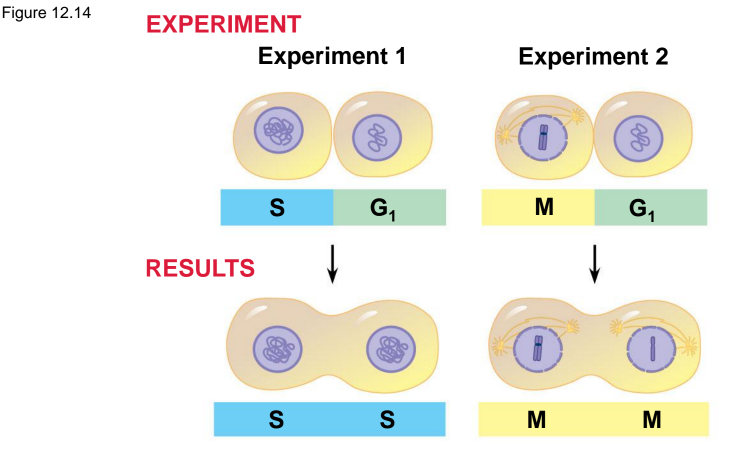


Concept 12.3: The eukaryotic cell cycle is regulated by a molecular control system

- The frequency of cell division varies with the type of cell
- These differences result from regulation at the molecular level
- Cancer cells manage to escape the usual controls on the cell cycle

Evidence for Cytoplasmic Signals

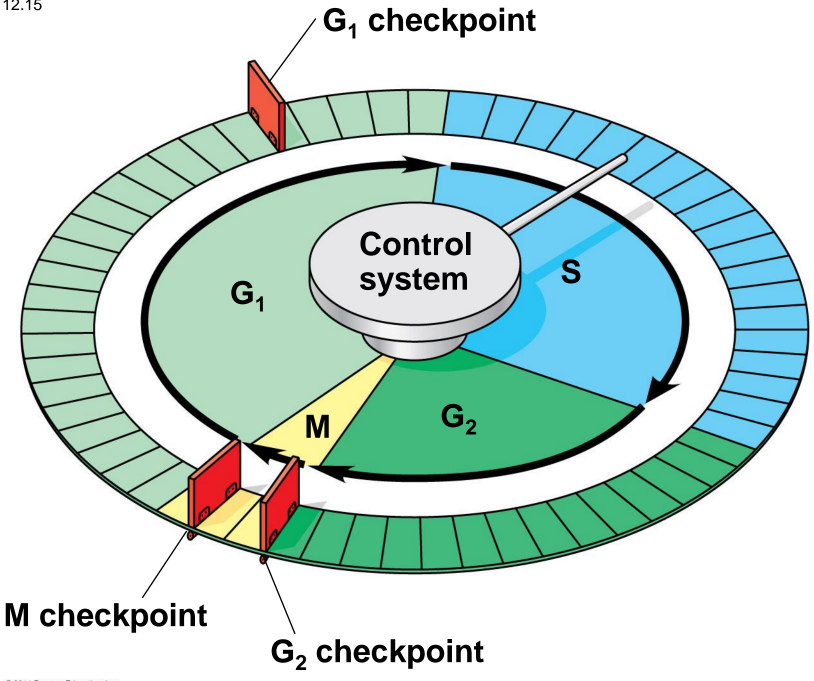
- The cell cycle appears to be driven by specific chemical signals present in the cytoplasm
- Some evidence for this hypothesis comes from experiments in which cultured mammalian cells at different phases of the cell cycle were fused to form a single cell with two nuclei



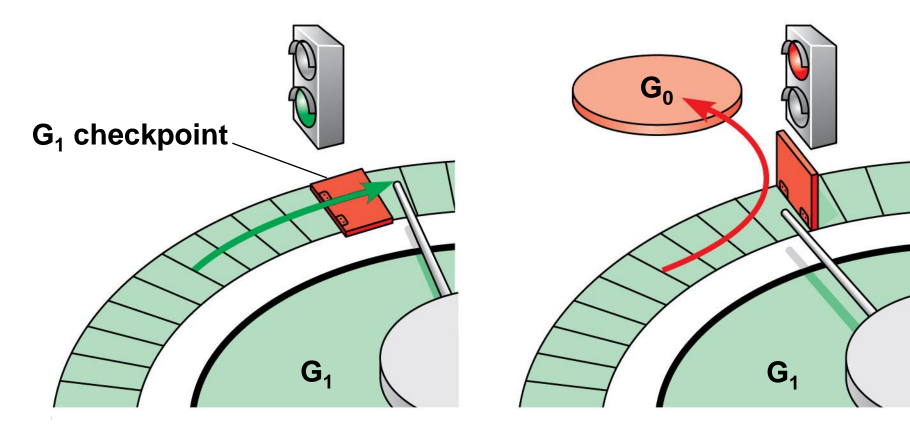
When a cell in the S phase was fused with a cell in G_1 , the G_1 nucleus immediately entered the S phase—DNA was synthesized. When a cell in the M phase was fused with a cell in G_1 , the G_1 nucleus immediately began mitosis—a spindle formed and chromatin condensed, even though the chromosome had not been duplicated.

The Cell Cycle Control System

- The sequential events of the cell cycle are directed by a distinct cell cycle control system, which is similar to a clock
- The cell cycle control system is regulated by both internal and external controls
- The clock has specific checkpoints where the cell cycle stops until a go-ahead signal is received



- For many cells, the G₁ checkpoint seems to be the most important
- If a cell receives a go-ahead signal at the G₁ checkpoint, it will usually complete the S, G₂, and M phases and divide
- If the cell does not receive the go-ahead signal, it will exit the cycle, switching into a nondividing state called the G₀ phase



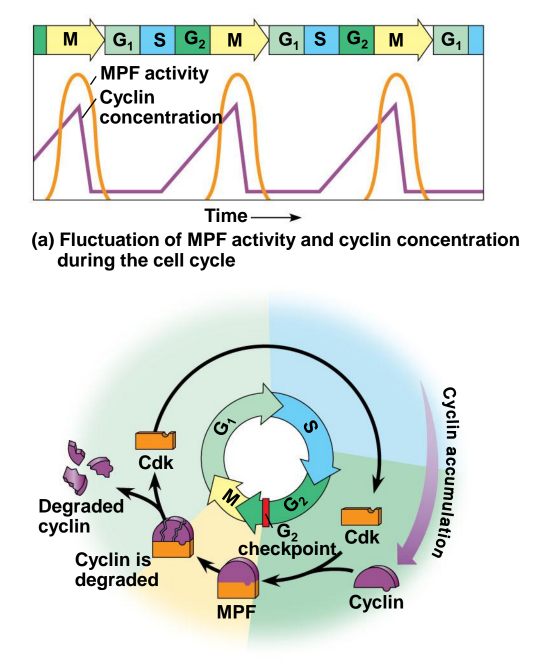
(a) Cell receives a go-ahead signal.

(b) Cell does not receive a go-ahead signal.

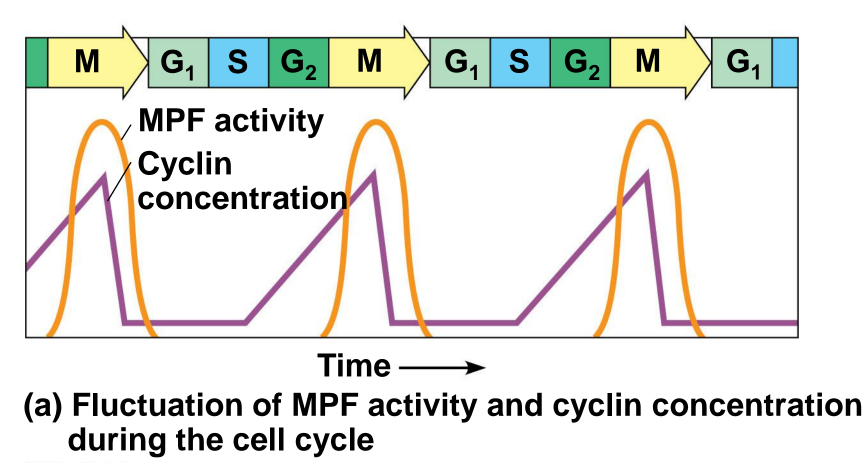
The Cell Cycle Clock: Cyclins and Cyclin-Dependent Kinases

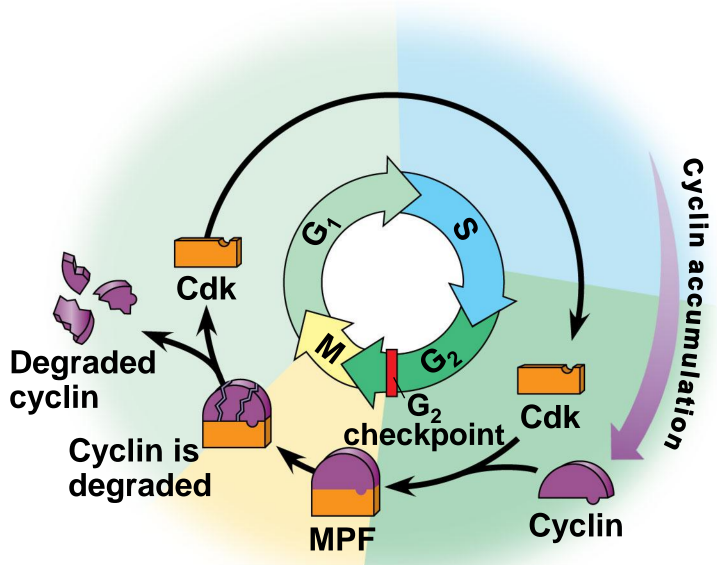
- Two types of regulatory proteins are involved in cell cycle control: cyclins and cyclin-dependent kinases (Cdks)
- Cdks activity fluctuates during the cell cycle because it is controled by cyclins, so named because their concentrations vary with the cell cycle
- MPF (maturation-promoting factor) is a cyclin-Cdk complex that triggers a cell's passage past the G₂ checkpoint into the M phase

Figure 12.17



(b) Molecular mechanisms that help regulate the cell cycle © 2011 Pearson Education, Inc.





(b) Molecular mechanisms that help regulate the cell cycle

Stop and Go Signs: Internal and External Signals at the Checkpoints

- An example of an internal signal is that kinetochores not attached to spindle microtubules send a molecular signal that delays anaphase
- Some external signals are growth factors, proteins released by certain cells that stimulate other cells to divide
- For example, platelet-derived growth factor (PDGF) stimulates the division of human fibroblast cells in culture

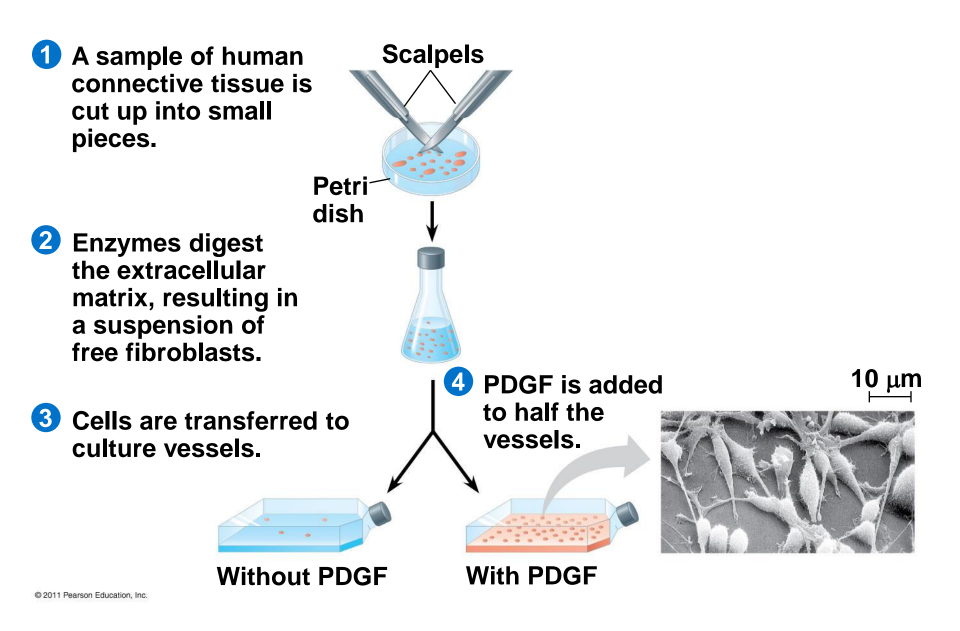
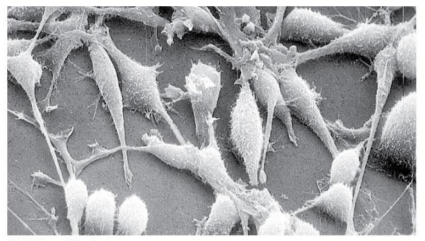


Figure 12.18a

10 μm



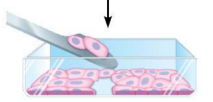
- A clear example of external signals is densitydependent inhibition, in which crowded cells stop dividing
- Most animal cells also exhibit anchorage dependence, in which they must be attached to a substratum in order to divide
- Cancer cells exhibit neither density-dependent inhibition nor anchorage dependence



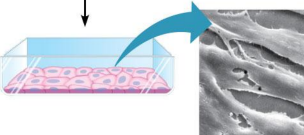
Anchorage dependence

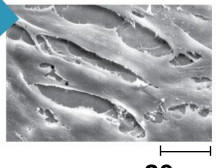


Density-dependent inhibition



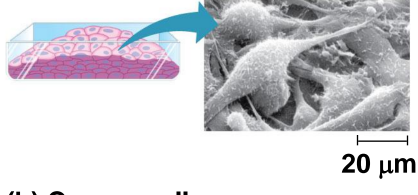
Density-dependent inhibition





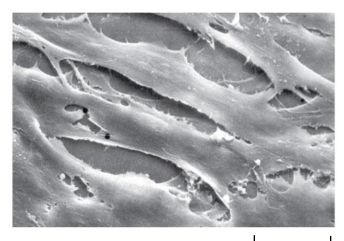
20 µm





(b) Cancer cells

Figure 12.19a



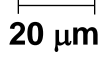
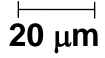


Figure 12.19b

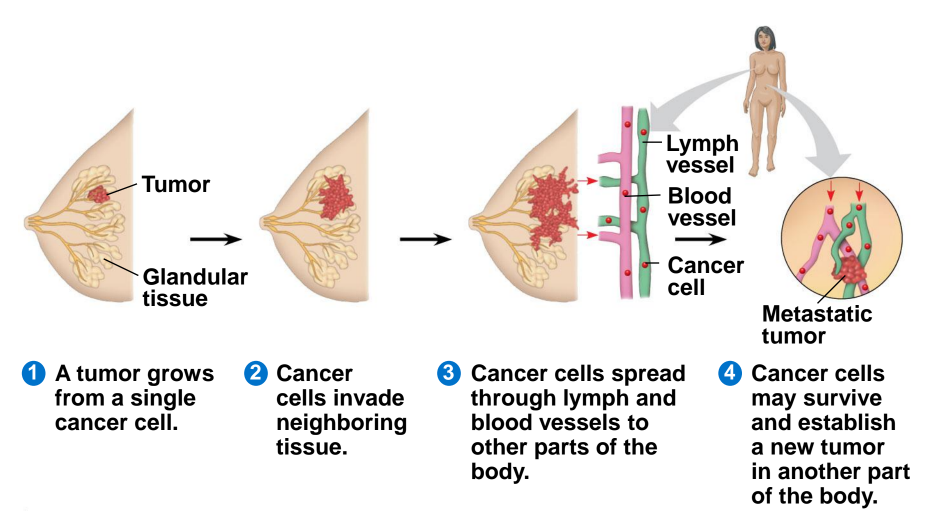




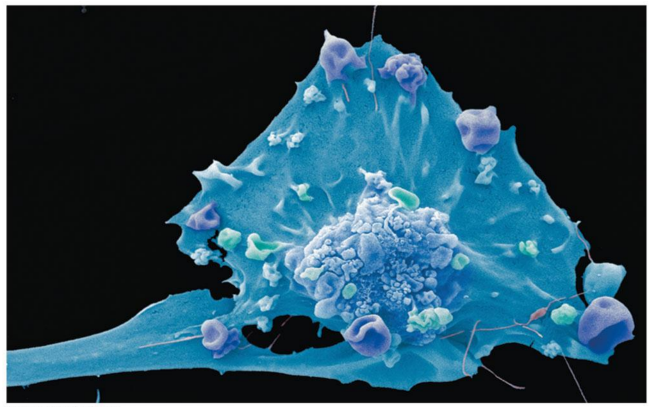
Loss of Cell Cycle Controls in Cancer Cells

- Cancer cells do not respond normally to the body's control mechanisms
- Cancer cells may not need growth factors to grow and divide
 - They may make their own growth factor
 - They may convey a growth factor's signal without the presence of the growth factor
 - They may have an abnormal cell cycle control system

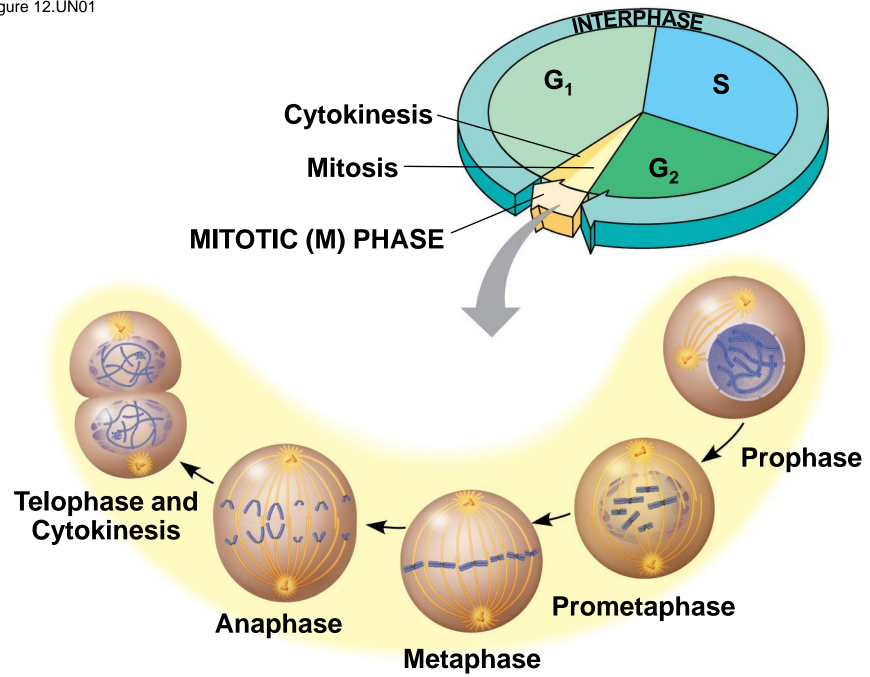
- A normal cell is converted to a cancerous cell by a process called transformation
- Cancer cells that are not eliminated by the immune system, form tumors, masses of abnormal cells within otherwise normal tissue
- If abnormal cells remain at the original site, the lump is called a benign tumor
- Malignant tumors invade surrounding tissues and can metastasize, exporting cancer cells to other parts of the body, where they may form additional tumors

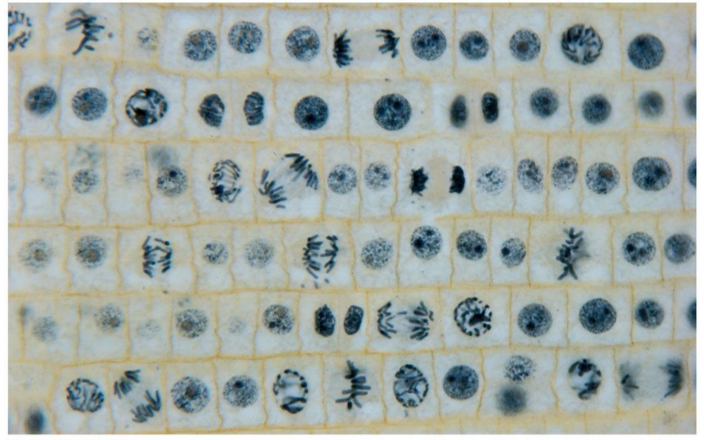


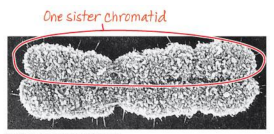
 Recent advances in understanding the cell cycle and cell cycle signaling have led to advances in cancer treatment











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