CHAPTER 2

Tissue Structure/Unit Cell Processes

2.1 Working/Operational Definitions

2.2 Tissue Composition/Structure
   Cells, insoluble extracellular matrix (ECM), soluble regulators
   2.2.1 Structure of the Cell

2.3 Types/classification of Tissue
   2.3.1 Tissue Classification
   2.3.2 Embryology
   2.3.3 Composition of Certain Tissues

2.4 Unit Cell Processes
   Limitations of the paradigm

2.5 Examples of Unit Cell Processes

2.6 Contraction and Migration

2.7 Summary of Effects of Exogenous Mechanical Forces on Cells and the Generation of Contractile Force by Cells
2.1 WORKING/OPERATIONAL DEFINITIONS

**Tissue**
An aggregation of similarly specialized cells united in the performance of a particular function. Cells serving the same general function and having the same extracellular matrix.

**Connective Tissue**
The matrix-continuous tissue which binds together and is the support of all of the structures of the body. The predominant structural protein comprising the extracellular matrix of connective tissue is collagen.

**Organ**
Two or more tissues combined to form a larger functional unit.

**Unit Tissue Structure**
The smallest repeating unit of a tissue or organ comprising a cell and its surrounding matrix (synthesized by the cell).

**Development**
The process of growth and differentiation of tissues and organs.

**Embryology**
The science of the development of the individual during the embryonic stage (2 weeks after fertilization of the ovum to the end of the eighth week) and, by extension, in several or even all preceding and subsequent stages of the life cycle.

**Regeneration**
The renewal of a tissue or organ at the completion of healing.

**Repair**
The formation of scar at a site of injury at the completion of healing.

**Remodeling/Maintenance/Turnover**
The process by which extracellular matrix is replaced in a process of degradation followed by synthesis.

**Endogenous Mechanical Force**
Force generated within a control volume of the unit cell process (e.g., by cell contraction).

**Exogenous Mechanical Force**
Force applied to a control volume of the unit cell process.

**Autocrine**
Denoting the influence of chemical factors secreted by a cell on itself.

**Paracrine**
Denoting influence of one cell on other cells in the vicinity.

**Endocrine**
Refers to tissues and organs whose function is to secrete into the blood or lymph a substance (e.g., hormone) that has a specific effect on another organ or tissue.
Hyperplasia
   The abnormal multiplication or increase in the number of normal cells in normal arrangement in a tissue.

Hypertrophy
   The enlargement or overgrowth of an organ or part due to an increase in size of its constituent cells.

Atrophy
   A diminution in the size of a cell, tissue, organ, or part.
2.2 TISSUE COMPOSITION/STRUCTURE

Unit Tissue Structure

Cell + Matrix $\rightarrow$ Product + Regulator

$C + M \rightarrow P + R$

TISSUES

For a Tissue "A" comprising cells of the "a" type:

$Tissue_A = \sum_i [C_a + M_a]$

For Example:

Dermis $= \sum_i [\text{Fibroblasts} + \text{Collagen}]$

Epidermis $= \sum_i [\text{Epidermal Cells} + \text{Basement Membrane}]$

Bone $= \sum_i [\text{Osteoblasts} + \text{Bone Matrix}]$

ORGANS

Organ $= T_A + T_B$

Skin $= \text{Epidermis} + \text{Dermis}$

Bone (Organ) $= \text{Bone (Tissue)} + \text{Marrow}$
2.3 TYPES/CLASSIFICATION OF TISSUE

2.3.1 Tissue Classification

Tissues can be classified into four groups:

1) Epithelia
2) Connective Tissue
3) Muscle
4) Nerve

2.3.2 Embryology

The tissues of the body develop from three types of embryonic tissues:

1) Ectoderm
2) Mesoderm
3) Endoderm
2.4 UNIT CELL PROCESSES

2.4.1 Cell Activities
"Packaged" sequences (cascades) of cell activities/functions with an identifiable beginning and an end (Fig. 2.1).

2.4.1.1 Mitosis (Proliferation) Fig. 2.2
Mitosis is a complex of processes leading to the division of a cell such that the two daughter nuclei receive identical complements of the number of chromosomes characteristic of the cells of the species. All cells arise from the division of pre-existing cells. The process of mitosis is divided into four phases: a) prophases-formation of paired chromosomes; disappearance of nuclear membrane, b) metaphase-chromosomes separate into exactly similar halves, c) anaphase-the two groups of daughter chromosomes separate and move along the fibers of a central "spindle", and d) telophase-the daughter chromosomes resolve themselves into a "reticulum" and the daughter nuclei are formed; the cytoplasm divides, forming two complete daughter cells.

The term mitosis is used interchangeably with cell division, but strictly speaking it refers to nuclear division whereas cytokinesis refers to division of the cytoplasm.

2.4.1.2 Synthesis (e.g., ECM structural proteins, enzymes)
Synthesis refers to the putting together of a chemical compound by the cell through the union of the elements comprising the compound or from other suitable starting materials. The chemical compounds can be proteins that serve as structural elements (e.g., collagen), enzymes (e.g., collagenase), regulators of other cells (e.g., hormones), immunoglobulins, etc. The compounds can be carbohydrates, lipids, or other macromolecules required in cell, tissue, or organ function. The products of cell synthesis can be stored in the cytoplasm or excreted by the cell.

2.4.1.3 Exocytosis (including degranulation)
Exocytosis refers to the process by which a cell excretes particles that are too large to diffuse through the cell membrane. It is considered to be the opposite of endocytosis.

2.4.1.4 Endocytosis
Endocytosis refers to the uptake by a cell of material too large to diffuse through its membrane. In the process of ingesting the material the cell invaginates its membrane, collecting the material in the fold produced by the invagination. Particles of the material larger than approximately one micrometer are "phagocytosed." In this process the cell membrane binds to the particle through a membrane receptor. Particles less than one micrometer are collected in the folds of the invaginated membrane with or without receptor binding, referred to as receptor-mediated endocytosis and pinocytosis, respectively. Fluid is taken up by the cell through the process of pinocytosis.

2.4.1.5 Migration
Migration refers to the movement of cells as a result of the action of intracellular cytoskeletal and contractile proteins and the receptor-mediated adhesion of the cell to extracellular matrix components (by integrins).
2.4.1.6 Contraction
At the cell level, contraction refers to the shortening of a cell and the concomitant development of tension in the matrix, as a result of the action of intracellular contractile proteins. The contractile proteins facilitating cell contraction are of the same family as facilitate cell migration.

2.4.2 Cell Protagonists
Requires a specific differentiated cell protagonist (Table 2.1). The process begins when the cell becomes the protagonist when expresses a specific phenotype or engages in a specific activity.

2.4.3 Chronology
Can appear at different sites at different times in different physiological and pathological sequences (i.e., modular; Table 2.2).

2.4.4 Matrix
Requires an insoluble matrix component/substrate (Table 2.3) for the cell activity. Cell acts on a substrate (e.g., extracellular matrix) through a membrane receptor (e.g., integrin) and a series of stereospecific biochemical reactions initiated by a signal (i.e., soluble regulator; see sec. 2.4.8). The matrix can serve as an insoluble regulator of cell function/activity.

2.4.5 Products
Results in a product that is a stable insoluble structure, soluble fragment, and/or mechanical force (Table 2.4). The quality of the product is invariant but the quantity is not; the product is quantifiable with respect to amount and direction/orientation, thus making the product a vector quantity.

2.4.5.1 Macromolecular Aggregates (Macromolecule) or Soluble Fragments
An example is collagen. The structure of a macromolecule is its conformation. In this case only one cell type is required (e.g., fibroblast) in the unit cell processes of synthesis and degradation.

2.4.5.2 Acellular Multicomponent Structure (Matrix)
Examples are basement membrane and clot. The structure comprises two or more types of macromolecules and may include nonviable cells. The structure of a matrix is its architecture.

2.4.5.3 Tissue (Cell + Matrix)
A system of multicomponent structures with viable cells. The structure of a tissue is its morphology.

2.4.6 Examples
Examples of unit cell processes are given in Tables 2.5 and 2.6.

2.4.7 Composite Processes
Two or more unit processes can combine to form a composite process (Table 2.7).
2.4.8 Regulators

Unit cell processes are regulated by diffusible soluble substances (Tables 2.8 - 2.11) acting on the cell protagonist directly or by mechanical forces acting on the cell indirectly through the matrix (e.g., by deforming the matrix). The regulators signal the start and the termination of the process. They also serve as connecting links between processes that comprise a composite process (e.g., the "coupling factor" between bone resorption and bone formation). In addition, regulators can control the rate of the process by acting on the cell and the substrate.

2.4.8 Rates of Processes

If the time period over which the unit cell process acts is t, then the rate of the process could be considered to be the quantity of product divided by t.
2.4 UNIT CELL PROCESSES

TABLE 2.1

Cell Protagonists

Cells of the same kind associate to form tissues. Tissues are divided into four types: connective tissue, epithelia, muscle, and nerve. Organs are formed by the combination of two or more tissues.

A. Connective Tissue (Matrix-Continuous)
   1. Blood (Cells floating freely in a fluid matrix until clotting; then a fibrillar matrix)
      a. Neutrophil
      b. Eosinophil
      c. Basophil
      d. Monocyte
      e. Lymphocyte
      f. Plasma cell
      g. Platelet
      h. Red blood cell

   2. Reticular Tissue (Cells in semi-solid matrix comprising reticular fibers, i.e., small diameter Type III collagen fibers. Examples include the framework of spleen, lymph nodes, bone marrow.)
      Fibroblast (synthesis of the Type III collagen)
      Reticular cell (macrophage-like)

   3. Loose Fibrous Tissue (Cells in semi-solid matrix comprising reticular and thicker collagen fibers. An example is stroma, the supporting tissue or matrix of an organ, as distinguished from its functional element, or parenchyma.)
      Fibroblast

   4. Dense Fibrous Tissue (e.g., dermis, ligament, tendon)
      Fibroblast

   5. Adipose (Fat) Cell

   6. Hyaline Cartilage and Fibrocartilage
      Chondrocyte

   7. Bone
      a. Osteoblast
      b. Osteocyte
      c. Osteoclast

   8. Tooth
      a. Odontoblast (synthesis of dentin)
b. Cementocyte (synthesis of cementum)

9. Other
   a. Macrophage/Histiocyte
   b. Mast cell
   c. Pericyte

B. Epithelia (Cell-Continuous)
   1. Simple
      a. Squamous cell (e.g., lining of blood vessels)
      b. Cuboidal cell (non-secretory and secretory)
      c. Columnar cell (e.g., lining of organs; including ameloblasts that synthesize enamel)
   2. Pseudostratified Columnar Ciliated Cell (respiratory passages)
   3. Compound (Stratified)
      a. Transitional cell (e.g., lining of urinary passages)
      b. Columnar cell (relatively uncommon)
      c. Squamous cell (uncornified and cornified, e.g., skin)

C. Muscle (Contractile Cells)
   1. Smooth muscle cell
   2. Cardiac muscle cell
   3. Skeletal (striated) muscle cell
   4. Myofibroblast

D. Nerve Tissue
   Nerve Cell
### TABLE 2.2

**Sites and Times at Which Unit Cell Processes Occur in Certain Clinical Sequences**

<table>
<thead>
<tr>
<th>Process/ (Protagonist Cell)</th>
<th>Site</th>
<th>Time</th>
<th>Clinical Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clotting (Platelet)</td>
<td>Vascular Tissue</td>
<td>Acute and Chronic</td>
<td>Wound Healing Stroke (thrombosis)</td>
</tr>
<tr>
<td>Collagen Degradation (Fibroblast)</td>
<td>Connective Tissue</td>
<td>Acute</td>
<td>Childbirth Wound healing</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Chronic</td>
<td>Collagen turnover Development Tumor resolution</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Acute and Chronic</td>
<td>Neoplasia Metastasis Bacterial infection (e.g. pneumonia, periodontal disease)</td>
</tr>
<tr>
<td>Collagen Synthesis (Fibroblast)</td>
<td>Connective Tissue</td>
<td>Acute and Chronic</td>
<td>Tumor growth Scar formation Regeneration</td>
</tr>
<tr>
<td>Epithelialization (Epithelial Cell)</td>
<td>Skin and oral mucosa</td>
<td>Acute</td>
<td>Wound healing</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Chronic</td>
<td>Epithelioma</td>
</tr>
</tbody>
</table>
### TABLE 2.3

**Substrates Involved in Unit Cell Processes:**

**Components of Extracellular Matrix**

- Collagen
- Elastin
- Adhesion Proteins (e.g., fibronectin, laminin)
- Glycosaminoglycans
- Proteoglycans
- Apatite (mineral)
### TABLE 2.4

Products of Unit Cell Processes

<table>
<thead>
<tr>
<th>PROCESS</th>
<th>PRODUCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mitosis</td>
<td>More cells/Cell proliferation</td>
</tr>
<tr>
<td>Synthesis</td>
<td></td>
</tr>
<tr>
<td>Insoluble matrix proteins</td>
<td>Matrix</td>
</tr>
<tr>
<td>Enzymes</td>
<td>Soluble matrix fragments</td>
</tr>
<tr>
<td>Cytokines</td>
<td>Regulators</td>
</tr>
<tr>
<td>Exocytosis (of stored granules/packets of regulators)</td>
<td>Regulators</td>
</tr>
<tr>
<td>Endocytosis</td>
<td>Solubilized fragments</td>
</tr>
<tr>
<td>Migration</td>
<td>Translocation</td>
</tr>
<tr>
<td>Contraction</td>
<td>Stress/Strain</td>
</tr>
</tbody>
</table>
2.4 UNIT CELL PROCESSES

TABLE 2.5

Examples of Unit Cell Processes

1. Clotting
   Platelets interacting with ("activated" by) collagen fibers (serving as an insoluble regulator) or reacting to injury/implants act on a collagen fiber to \textit{exocytose} granules of pre-packaged regulators (process of degranulation) and produce a clot.

2. Endocytosis
   Macrophages \textit{endocytose} fragments of the substrate. The substrate can be ECM, bacteria, or synthetic or natural materials related to implants. The products are small molecular weight metabolites.

3. Collagen Degradation
   Fibroblasts \textit{synthesize} the enzyme, collagenase, to depolymerize a collagen fiber to produce soluble peptide fragments.

4. Collagen Synthesis
   Fibroblasts attach to ECM components and \textit{synthesize} collagen molecules.

5. Contraction
   Myofibroblasts exert \textit{contractile} forces on collagen fibers.

6. Epithelialization
   Epithelial cells act on a basement membrane and undergo \textit{mitosis} in order to produce a confluent layer.

7. Bone Formation
   Osteoblasts attach to ECM components and \textit{synthesize} bone matrix.

8. Bone Resorption
   Osteoclasts attach to bone matrix and \textit{synthesize} protons and collagenase in order to solubilize the matrix, thereby producing peptide fragments and components of the apatite mineral.
### 2.4 UNIT CELL PROCESSES

**TABLE 2.6**

Examples of Unit Cell Processes

<table>
<thead>
<tr>
<th>Unit Cell Process</th>
<th>Cell Protagonist</th>
<th>Substrate</th>
<th>Activity/Function</th>
<th>Product</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clotting</td>
<td>Platelet</td>
<td>Collagen</td>
<td>Exocytosis (Degranulation)</td>
<td>Platelet Aggregation</td>
</tr>
<tr>
<td>Collagen Degradation</td>
<td>Fibroblast</td>
<td>Collagen</td>
<td>Synthesis: Collagenase</td>
<td>Soluble Collagen Fragments</td>
</tr>
<tr>
<td>Collagen Synthesis</td>
<td>Fibroblast</td>
<td>ECM Component</td>
<td>Synthesis: Collagen</td>
<td>Collagen</td>
</tr>
</tbody>
</table>
2.4 UNIT CELL PROCESSES

TABLE 2.7

Composite Unit Cell Processes

1. Collagen Remodeling
   Collagen degradation + collagen synthesis

2. Neovascularization
   Collagen synthesis + endothelialization

3. Scar Formation
   Collagen synthesis + contraction

Supercomposite Unit Cell Processes

1. Tissue Remodeling Associated with Wound Healing
   Collagen remodeling + neovascularization + scar formation

2. Bone Remodeling
   Bone resorption + bone formation

3. Acute Inflammation
   De-endothelialization + clotting + endocytosis + neovascularization

3. Granulation Tissue Formation
   Neovascularization + collagen synthesis + endocytosis

4. Chronic Inflammation
   Endocytosis + neovascularization + collagen degradation + collagen synthesis.
2.4 UNIT CELL PROCESSES

TABLE 2.8

Regulators of Unit Cell Processes

Cytokines*

Interleukins
  IL-1
  IL-6
Tumor Necrosis Factor (TNF)
Platelet Derived Growth Factor (PDGF)
Insulin-like Growth Factor (IGF)
  IGF-1
  IGF-2
Fibroblast Growth Factor (FGF)
  basic FGF
Transforming Growth Factor (TGF)
  TGF-β

Eicosanoids**

Prostaglandins
  PGE2
Leukotrienes
  LTB4

Differentiation Factors

Bone Morphogenetic Protein (BMP)

* Cytokines are polypeptides (proteins) that regulate many cell functions. They act on a target cell by binding to specific high-affinity receptors. Cytokines that act on the same cell that produced them are called autocrine factors; those that act on other cells are called paracrine factors; those that act systemically (through the vascular system) are referred to as endocrine factors. Molecules that switch on (i.e., regulate) mitosis are referred to as growth factors.

** Eicosanoids are chemically related signaling lipid molecules made primarily from arachidonic acid (fatty acid). Eicosanoids include prostaglandins, leukotrienes, thromboxanes, and lipoxins. Prostaglandins are continuously synthesized in membranes from precursors (20-carbon fatty acid chains that contain at least 3 double bonds, e.g., arachidonic acid) cleaved from membrane phospholipids by phospholipases, membrane-bound enzymes. They are continuously released by the cell, and are degraded by enzymes in the extracellular fluids. The subscript of PGE2 refers to the 2 double bonds outside the ring structure.
## 2.4 UNIT CELL PROCESSES

### TABLE 2.9

Certain Cytokines (Growth Factors) as Regulators of Cell Activities*

<table>
<thead>
<tr>
<th>Cytokine</th>
<th>Mitosis FB** EN</th>
<th>Migration (Chemotaxis) FB EN MC</th>
<th>Synthesis, FB Collagen</th>
</tr>
</thead>
<tbody>
<tr>
<td>Collagenase</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Platelet Derived Growth Factor (GF) (PDGF)</td>
<td>+ 0</td>
<td>+ 0 +</td>
<td>+ +</td>
</tr>
<tr>
<td>Fibroblast GF (FGF)</td>
<td>+ +</td>
<td>+ + ?</td>
<td>? +</td>
</tr>
<tr>
<td>Transforming GF-β (TGF-β)</td>
<td>-/+ -</td>
<td>+ ? +</td>
<td>+ +</td>
</tr>
<tr>
<td>Transforming GF-α and Epidermal GF (EGF)</td>
<td>+ +</td>
<td>0 + ?</td>
<td>? +</td>
</tr>
<tr>
<td>Interleukin-1 (IL-1) and Tumor Necrosis Factor (TNF)</td>
<td>+ 0/-</td>
<td>? ? +</td>
<td>+ +</td>
</tr>
</tbody>
</table>

+ - Stimulates  
- - Inhibits  
0 - No effect

** FB- Fibroblast; EN-Endothelial Cell; MC - Monocyte
### 2.4 UNIT CELL PROCESSES

**TABLE 2.10**

Regulators in Acute Inflammation

<table>
<thead>
<tr>
<th>Cell Regulator</th>
<th>Source</th>
<th>Vascular Leakage/Endoth. Cell Contraction</th>
<th>Leukocyte Migration/Chemotaxis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Eisoanoid</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leukotriene B₄ (LTB₄)</td>
<td>Leukocyte</td>
<td>0</td>
<td>+</td>
</tr>
<tr>
<td><strong>Cytokines</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Platelet Act. Factor (PAF)</td>
<td>Leukocyte</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>IL-1</td>
<td>Macrophage</td>
<td>0</td>
<td>+</td>
</tr>
<tr>
<td>TNF</td>
<td>Macrophage</td>
<td>0</td>
<td>+</td>
</tr>
<tr>
<td><strong>Amino Acid Derivatives</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Histamine Platelet</td>
<td>Mast Cell</td>
<td>+</td>
<td>0</td>
</tr>
<tr>
<td>Serotonin Platelet</td>
<td>Mast Cell</td>
<td>+</td>
<td>0</td>
</tr>
</tbody>
</table>
### TABLE 2.11

Regulators for Bone Cells

<table>
<thead>
<tr>
<th>Regulator</th>
<th>Cell Source (Regulator)</th>
<th>Osteoblast, OB Mitosis</th>
<th>Osteoblast, OB Synthesis</th>
<th>Osteoclast, OC Mitosis</th>
<th>Osteoclast, OC Synthesis*</th>
</tr>
</thead>
<tbody>
<tr>
<td>TGF-β</td>
<td>Mφ, OB, OC</td>
<td>+</td>
<td>+</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>bFGF</td>
<td>Mφ, OB</td>
<td>+</td>
<td>-</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>PDGF</td>
<td>Mφ, OB</td>
<td>+</td>
<td>+</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>IGF</td>
<td>OB</td>
<td>+</td>
<td>+</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>BMP</td>
<td>OB</td>
<td>+</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>IL-1</td>
<td>Mφ</td>
<td>+</td>
<td>+?</td>
<td>+</td>
<td>0</td>
</tr>
<tr>
<td>PGE₂</td>
<td>Mφ, OB</td>
<td>+</td>
<td>+?</td>
<td>?</td>
<td>+</td>
</tr>
<tr>
<td>TNF α</td>
<td>Mφ</td>
<td>+</td>
<td>?PGE₂</td>
<td>?</td>
<td>+</td>
</tr>
<tr>
<td>CSF-GM</td>
<td>Mφ, OB</td>
<td>0</td>
<td>0</td>
<td>+</td>
<td>0</td>
</tr>
<tr>
<td>Calcitonin (S)**</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>PTH (S)</td>
<td>0</td>
<td>PGE₂, CSF-GM</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>EGF (S)</td>
<td>+</td>
<td>-</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

* Osteoclast activity relates to synthesis of enzymes and hydrogen ions.
** Systemic factors not yet found to be produced in bone.

TGF: Transforming Growth Factor  
FGF: Fibroblast Growth Factor  
PDGF: Platelet Derived Growth Factor  
IGF: Insulin-like Growth Factor  
BMP: Bone Morphogenetic Protein  
IL: Interleukin  
TNF: Tumor Necrosis Factor  
CSF-GM: Colony Stimulating Factor-Granulocyte/Monocyte  
PTH: Parathyroid Hormone  
EGF: Epidermal Growth Factor
2.4 UNIT CELL PROCESSES

TABLE 2.12
Summary of Matrices, Regulators, and Unit Cell Processes According to Tissue Type

<table>
<thead>
<tr>
<th>Tissue Type</th>
<th>Matrices</th>
<th>Regulators</th>
<th>Mit</th>
<th>Syn</th>
<th>Migr</th>
<th>Cntr</th>
<th>Endo</th>
<th>Exoc</th>
</tr>
</thead>
<tbody>
<tr>
<td>Connective Tissue</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Epithelia</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Muscle</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Skeletal</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cardiac</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Smooth</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nerve</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
2.5 EXAMPLES OF UNIT CELL PROCESSES
2.6 CONTRACTION AND MIGRATION: ENDOGENOUS FORCE GENERATION

2.6.1 Functions of Cytoplasmic Actin Filaments
1) Contribute to maintaining cell shape.
2) Enable cell division.
3) Cell migration (many cell types).
4) Endocytosis (viz., polymorphonuclear neutrophils and macrophages).
5) Provide for cell contraction that results in tensile or compressive forces in the extracellular matrix. These forces cause a) tissue contraction in wound healing, b) the normal "tensioning" of structures such as tendons and ligaments, and c) pathological contracture associated with certain disorders (e.g., contracture of scar and the fibrous capsule around breast implants).
6) Intracellular movement of molecules and organelles.

2.6.2 Actin Filament Formation
1) Single molecules (i.e., monomer units of G-actin) aggregate (polymerize) to form small groups (2-3 G-actin units, dimers and trimers), which serve as "nuclei" to which other G-actin molecules attach.
2) Polymerization leads to formation of a long stiff rod (filament) of actin (F-actin). Because of the shape of G-actin units and the geometry of aggregation one end of the F-actin filament has the shape of an arrowhead and the other end is "barbed."

2.6.3 Mechanisms of Generating Tensile and Contractile Forces within the Cell

Tensile Forces
1) Reversible polymerization of G-to F-actin
2) Bundling of F-actin filaments (polymers) into a stiff fiber (stress fiber)
3) Reversible gelation - solvation of 3-D actin gels
   a. "Gelation" occurs with the formation of 3-D networks of orthogonally oriented F-actin filaments produced by bonding of "actin binding protein" (ABP) to two F-actin filaments. ABP serves as a flexible hinge.
   b. ABP facilitates actin gelation
   c. A calcium bonding protein (gelsolin) can dissolve actin - ABP gels (Ca dependent)

Contractile Forces
4) Myosin - actin interaction: sliding filament model
   Globular head of hinged myosin molecule attaches to an actin molecule. Alteration (conformational change) of the myosin molecule causes actin filament to slide. Myosin then attaches to the adjacent actin molecule and the process is repeated.
5) Myosin - actin interaction: Contraction of orthogonal actin network.
2.7 SUMMARY OF THE EFFECTS OF EXOGENOUS MECHANICAL FORCES ON CELLS AND GENERATION OF CONTRACTILE FORCE BY CELLS

2.7.1 Effects of Exogenous Forces

A. Increase in Force Leads to Increase in Tissue Mass
   1) Hypertrophy (e.g., skeletal muscle)
      Increase in synthesis of contractile proteins.
   2) Hyperplasia (e.g., bone)
      a. Increased number of cells (growth) resulting from mitosis of pre-osteoblasts.
         The cell cycle is shortened and/or cells in G₀ are recruited into the cell cycle.
         Strain might serve as a competence factor to render cells in G₀ or G₁ competent
         to synthesize DNA and/or as a progression factor that stimulates DNA synthesis
         in competent cells.
      b. Increased synthesis of extracellular matrix components (viz., collagen and
         glycosaminoglycans) by osteoblasts

B. Decrease in Force Leads to Decrease in Tissue Mass (Atrophy)
   1) Imbalance between the degradation and formation phases of tissue remodeling.
      Degradation may proceed at the normal rate but formation decreased in the absence
      of normal mechanical loading (strains).
   2) Mechanical strains may act to downregulate cell synthesis of enzymes and other
      agents. A decrease in strain would allow for the increased synthesis of these tissue-
      degrading agents. This has not yet been demonstrated.

2.7.2 Effects of Endogenous Forces

A. Cell Contraction (Contractile Forces in the Cell)
   Synthesis of contractile proteins (viz., actin).
   Increase in contraction due to the increase in amount of contractile protein
   (intracellularly) and/or increase in number of cells (mitosis) capable of contracting.

B. Cell Migration (Tensile Forces in the Cell)
   Intracellular assembly of actin networks.