

UNIT VI

Chapter 34:

GUYTON AND HALL TEXTBOOK OF **MEDICAL PHYSIOLOGY** THIRTEENTH EDITION



Resistance of the Body to Infection: I. Leukocytes, Granulocytes, the Monocyte- Macrophage System, and Inflammation

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Defense Against Infection

Leukocytes

- **Microorganisms coexist with us and within us , which can be beneficial or harmful.**
- **Phagocytes can ingest and destroy invading organisms and participate in tissue reactions that “wall off” infection.**
- **Other white cells (lymphocytes, chapter 35) mediate responses that destroy or neutralize *specific* microorganisms.**

White Blood Cells

- Circulate in blood and may enter the tissues
- Are of six types:
 - Polymorphonuclear neutrophils
 - “ eosinophils
 - “ basophils
 - Monocytes
 - Lymphocytes (plasma cells)
 - Platelets (from megakaryocytes)

White Blood Cell Counts

- **Total WBC ~ 7,000 / mm³**
(almost 1,000-fold fewer than RBCs)
- **Proportions:**
 - Neutrophils 62%
 - Eosinophils 2.3%
 - Basophils 0.4%
 - Monocytes 5.3%
 - Lymphocytes 30%
- **Platelets ~ 300,000 / mm³**

Leukopoiesis

Genesis of Myelocytes

Bone marrow only **1 myeloblast**

3 megakaryocyte

2 promyelocyte

**Monocyte
genesis**

**4 neutrophil
myelocyte**

**8 eosinophil
myelocyte**

**11 basophil
myelocyte**

**5 Young neutrophil
metamyelocyte**

**9 eosinophil
Meta-
myelocyte**

**Polymorph-
nuclear
basophil**

**6 band neutrophil
metamyelocyte**

**7 Polymorph-
nuclear
neutrophil**

**10 polymorphnuclear
eosinophil**

12

16

Genesis of Lymphocytes

Mainly in
lymphogenous
tissues, lymph
glands,
thymus.....

Genesis of White Blood Cells

- **Granulocytes and monocytes develop in the bone marrow, and most remain there until needed peripherally (number in marrow ~3x blood; 6-day supply)**
- **Lymphocytes develop mostly in the peripheral lymphoid organs (thymus, spleen, tonsils, lymph nodes, Peyer's patches), less found in blood**
- **Megakaryocytes develop and reside in the marrow, fragment to release platelets**

Life Span of White Blood Cells

- **Granulocytes:**

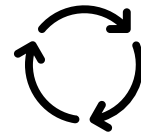
- Circulating, 4 – 8 hours
- In the tissues, 4 – 5 days
(shorter timelines with infection, inflammation)

- **Monocytes / Macrophages:**

- Circulating, 10 – 20 hours
- As tissue macrophages, months or longer

- **Lymphocytes:**

- Continuously re-circulate:
lymph...nodes...blood.. tissues (diapedesis)
- Long-lived... weeks, months, longer



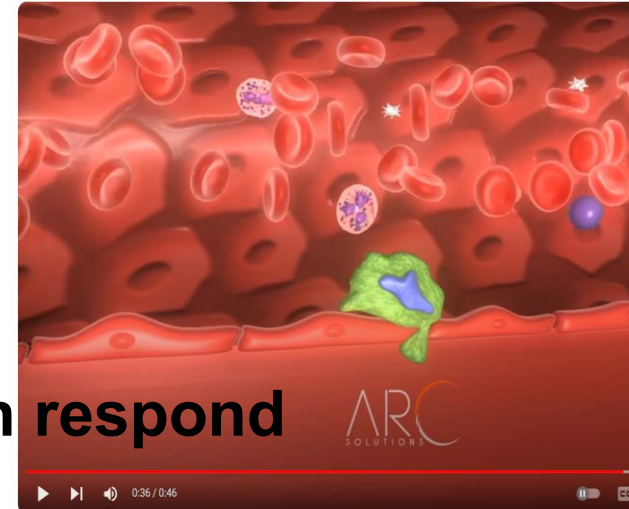
- **Platelets:** ~ Replaced every ten days~ 30K each day

Neutrophils and Macrophages

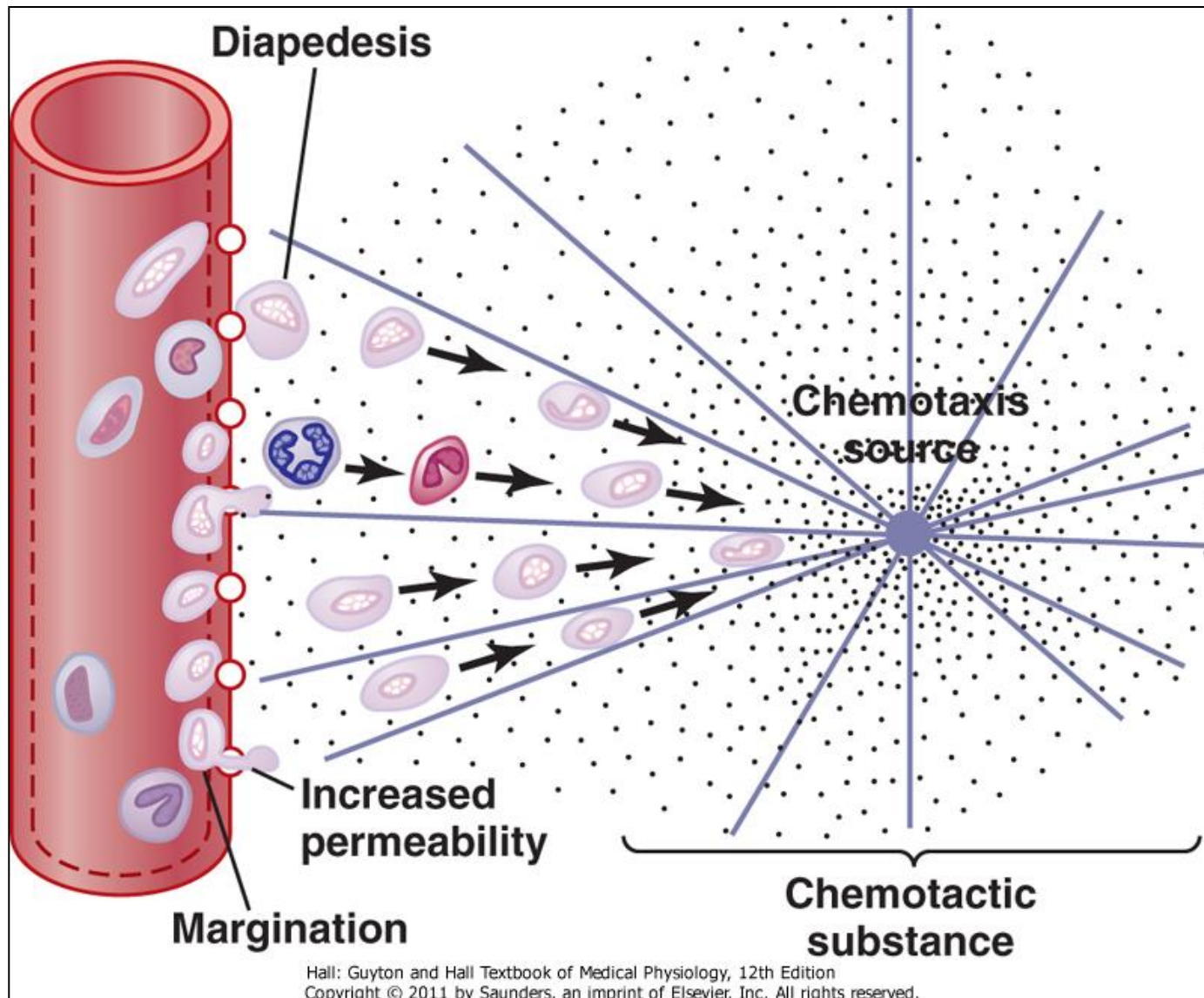
- Neutrophils are mature cells that can respond immediately to infection
- Monocytes mature in the tissues to become macrophages (monocytes in blood little ability)
- Both exhibit motility:
 - Diapedesis

[Diapedesis - Medical Animation by Arc Solutions - YouTube](#)

- Ameboid motion
- Chemotaxis (Chemoattractants: bacterial or tissue degradation products, complement fragments, other chemical mediators)



Neutrophil Margination & Migration



Phagocytosis

- “Phagocytosis” is the ingestion of particles
- Phagocytes must distinguish foreign particles from host tissues
- Appropriate phagocytic targets:
 - May have rough surfaces
 - Lack protective protein coats
 - May be immunologically marked for phagocytosis by *antibodies* or *complement components* that are recognized by receptors on the phagocytes
... this immunologic marking is called “opsonization”

Phagocytosis

- **Neutrophils**: can ingest 3-20 bacteria
- **Macrophages**: After being activated in the tissues, are extremely effective phagocytes (up to ~100 bacteria)
- **Macrophages can ingest larger particles...**
 - Damaged RBCs
 - Malarial parasites
- **Macrophages can extrude digestion products and survive and function for many months**

Digestion of Ingested Particles

- In both neutrophils and macrophages, *phagosomes* fuse with *lysosomes* and other granules to form *phagolysosomes* (*digestive vesicles*)
- These contain *proteolytic enzymes*, and in macrophages, *lipases* (important in killing tuberculosis bacillus and some other bacteria)

Bactericidal Agents

- Bacteria may be killed even if they are not digested
- Enzymes in the phagosome or in *peroxisomes* generate strongly bactericidal *reactive oxygen species...*
 - *Superoxide* (O_2^-)
 - *Hydrogen peroxide* (H_2O_2)
 - *Hydroxyl ions* (OH^-)
 - *Myeloperoxidase* catalyzes

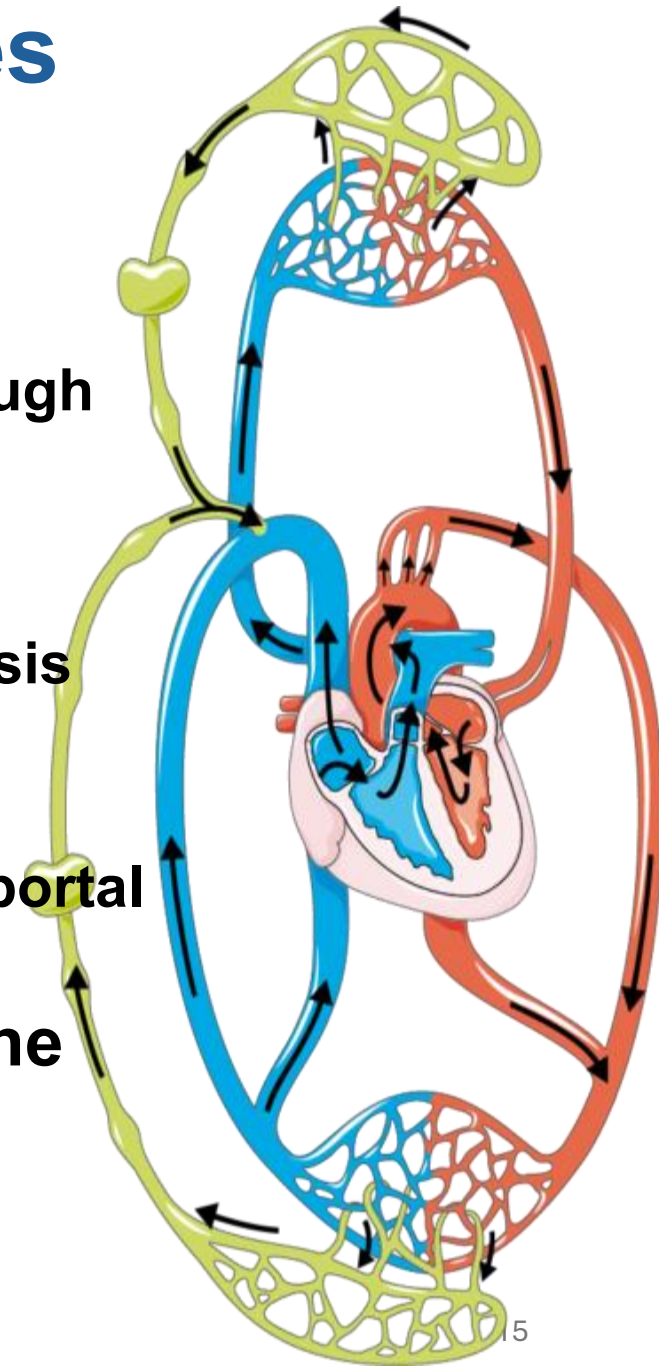


The Reticuloendothelial System

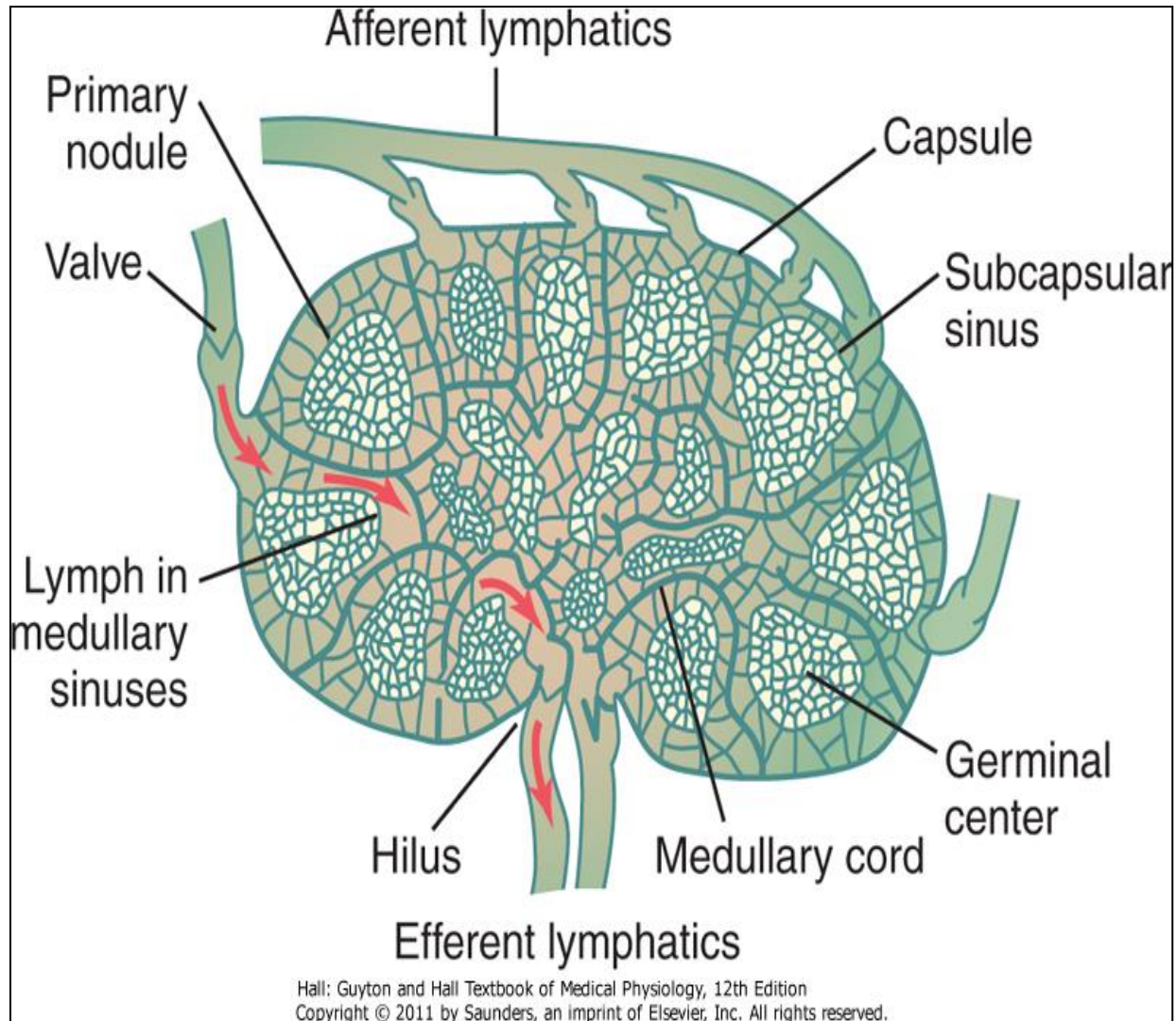
- **After entering the tissues, macrophages become fixed and may be resident for years**
- **When appropriately stimulated they can break away and move to sites of inflammation**
- **Circulating monocytes, mobile macrophages, fixed tissue macrophages, and some specialized endothelial cells form the *reticuloendothelial system*, almost all derived from monocytes, comprising a phagocytic system located in all tissues**

Specialized Macrophages

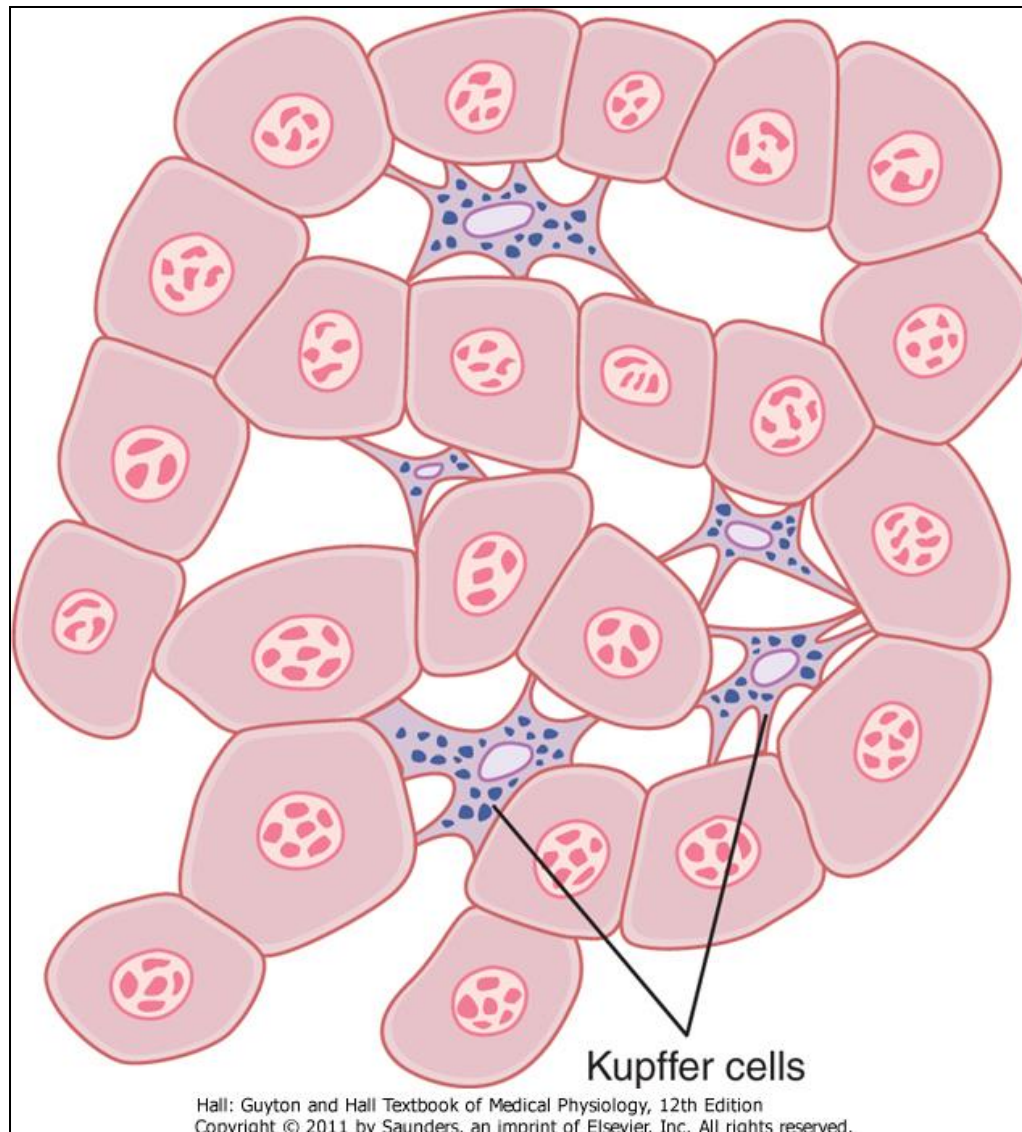
- **Skin, subcutaneous (histiocytes)**
- **Lymph nodes**
 - Ingest / sample particles arriving through the lymph
- **Alveolar macrophages**
 - Digest or entrap inhaled particles and microorganisms like silica, tuberculosis bacilli.
- **Kupffer cells**
 - Lining sinusoids, Surveillance of the portal circulation.
- **Macrophages in the spleen and bone marrow**
 - Surveillance of the general circulation



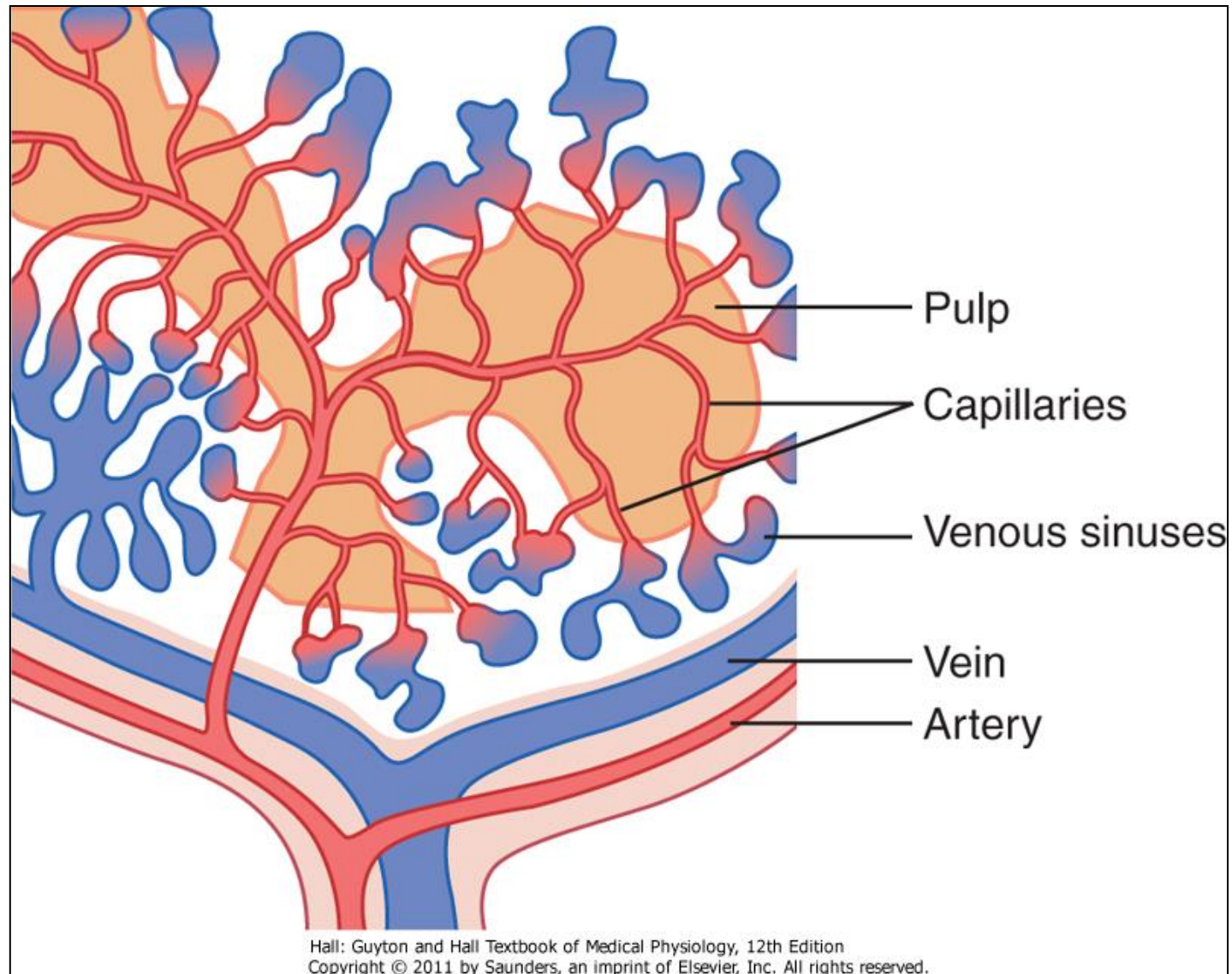
Structure of a Lymph Node



Kupffer Cells in the Liver Sinusoids



Structure of the Spleen



Neutrophils, Macrophages & Inflammation

- Inflammation is driven by chemical mediators and characterized by heat, redness, swelling, and pain
- Physiologically, it involves...
 - Vasodilatation and increased blood flow
 - Increased capillary permeability
 - Coagulation of interstitial fluids
 - Accumulation of granulocytes and monocytes
 - Swelling of tissue cells
- Mediators: *histamine, bradykinin, serotonin, prostaglandins, complement products, clotting components, lymphokines*

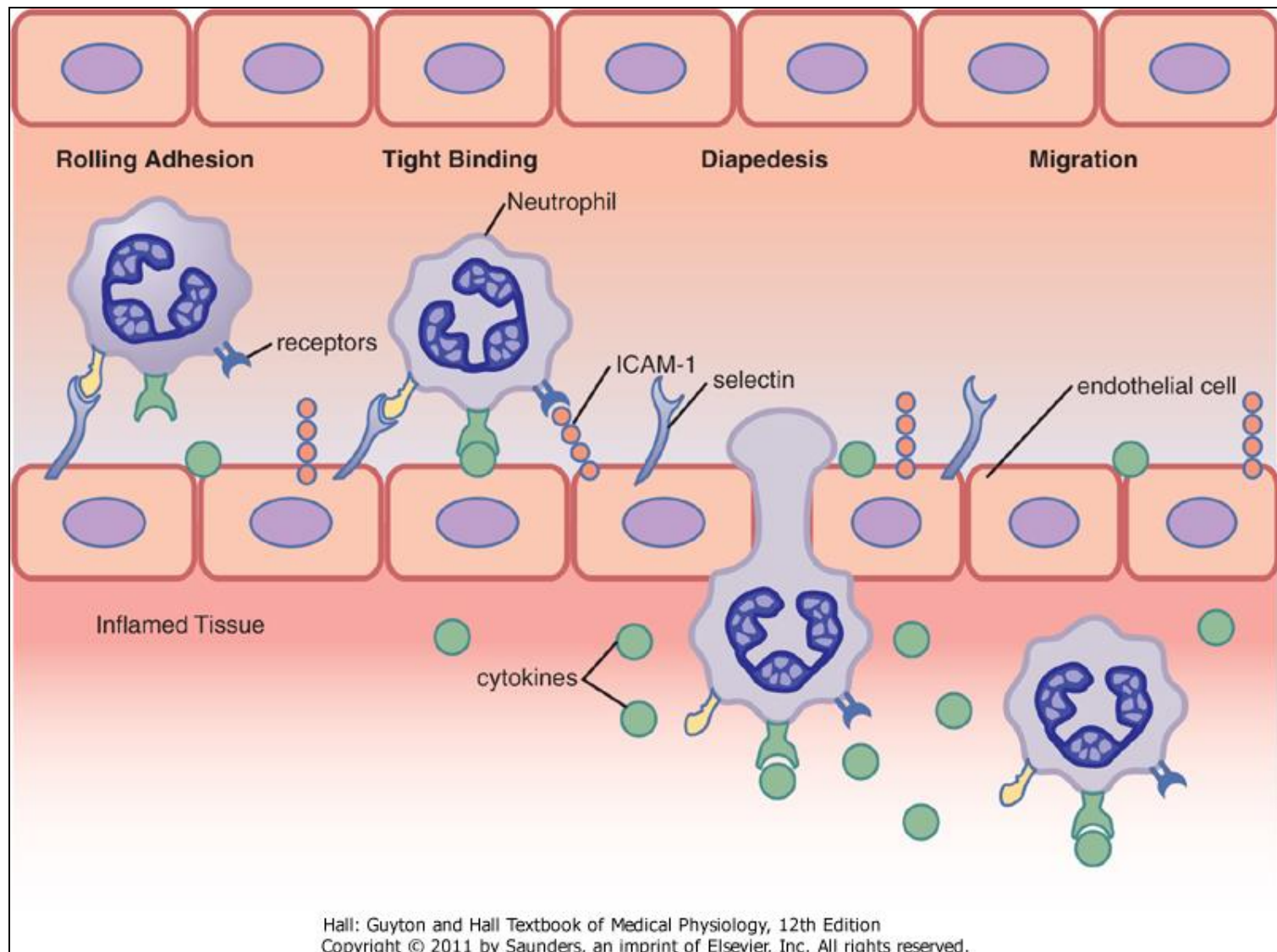
“Walling Off” Sites of Inflammation

- **Fibrinogen clots minimize fluid flow in and out of the inflamed area**
- ***Staphylococci* cause intense inflammation and are effectively “walled off”**
- ***Streptococci* induce less intense inflammation and may be more likely to spread than *staphylococci*, and cause death**

Neutrophils and Macrophages in Inflammation

- **Tissue macrophages** that encounter foreign particles enlarge and become mobile to provide **a first line of defense (min)**
- Within an hour neutrophils migrate to the area in response to inflammatory cytokines (TNF, IL-1)
2nd line of defense
- Upregulated ***selectins*** and ***ICAM-1*** on endothelial cells
- Bind to ***integrins*** on neutrophils, leading to ***margination***, followed by ***diapedesis***, and ***chemotaxis*** directing neutrophils into the inflamed tissues, to kill bacteria and scavenge

Neutrophil Migration to an Inflamed Site



Neutrophilia



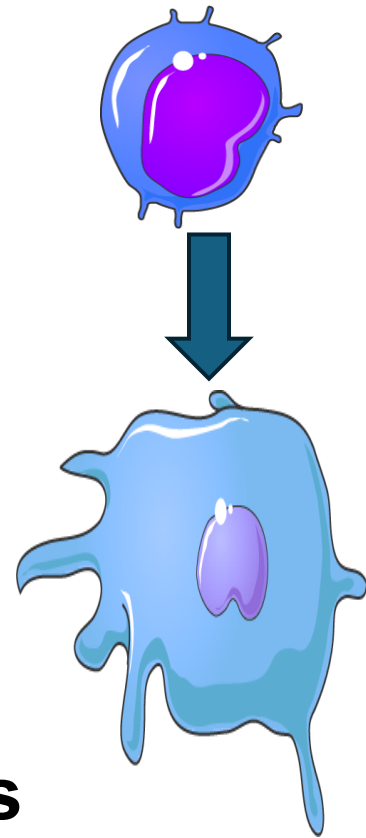
- **With intense inflammation neutrophil count can increase dramatically...**

4,000-5,000 **→** **15,000-25,000**

- **Results from mobilization of mature neutrophils from the bone marrow by inflammatory mediators**

Secondary Macrophage Invasion

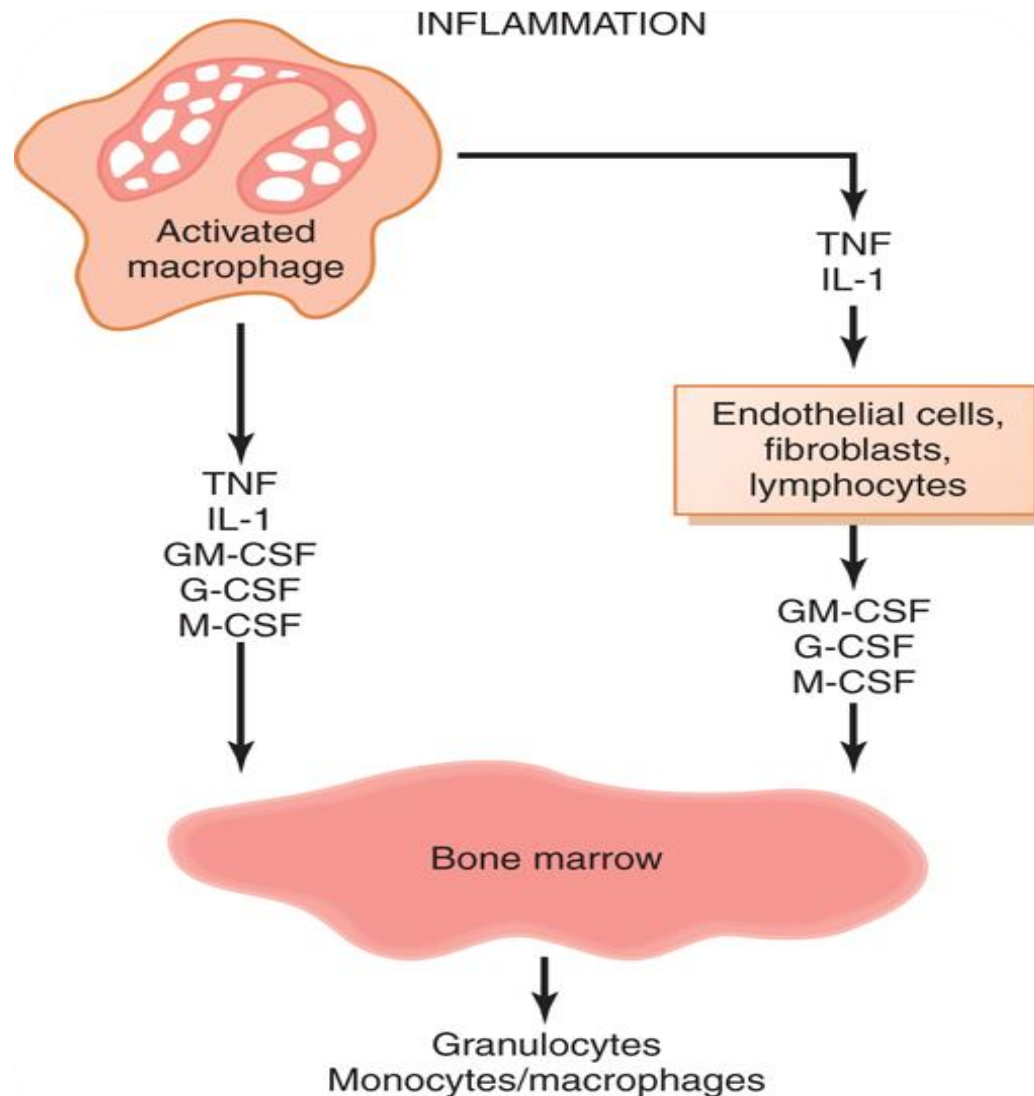
- In response to chemoattractants, monocytes gradually accumulate (slowly) and become macrophages (after ~ 8 hours mature)
- In part due to increased bone marrow production (store is low), macrophages become the dominant inflammatory cell over several weeks, cleaning up remaining bacteria, necrotic tissue, and directing tissue remodeling. **Third line of defense**



Bone Marrow Responses

- **Growth factors produced in response to infection and inflammation drive proliferation and differentiation of leukocyte precursors in the marrow**
- **First mature cells released after 3 – 4 days**
- **The bone marrow can increase production of granulocytes and monocytes by 20 – 50- fold and maintain this for months or years**
- **Fourth line of defense**

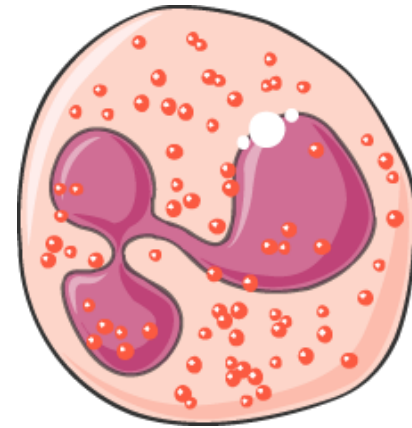
Bone Marrow Response to Inflammation



Formation of Pus

- **Pus is composed of dead bacteria and neutrophils, many dead macrophages, necrotic tissue that has been degraded by proteases, and tissue fluid, often in a cavity formed at the inflammatory site**
- **Over days and weeks it is absorbed into the surrounding tissue and lymph and disappears**

Eosinophils



- **Eosinophils are weak phagocytes and exhibit chemotaxis**
- **Particularly important in defense against parasites, Ex: schistosomiasis and trichinosis**
- **Can adhere to parasites and release substances that kill them (hydrolases, reactive oxygen species, major basic protein(larvacidal)).**
- **Also accumulate in tissues affected by allergies, perhaps in response to eosinophil chemotactic factor from basophils (eosinophils may detoxify some products of basophils)**

Basophils



- Similar to *mast cells* adjacent to **Capillaries**, both cell types release heparin
- Basophils and mast cells both release histamine, bradykinin, and serotonin
- When IgE bound to receptors on their surfaces is cross-linked by its specific antigen, mast cells and basophils *degranulate*, releasing...
 - *histamine, bradykinin, serotonin, heparin, leukotrienes, and several lysosomal enzymes*