

تفريغ علم وظائف الأعضاء المرضي



اسم الموضوع:

Inflammation

إعداد الصيدلاني/ة:

Shahed Abushosheh



لجان الرفعات

Shahed Abushosheh



اللَّهُمَّ أكرمنا بالفهم
وأفتح علينا بالعلم
وزدنا اللهم علماً وحلماً

@da44.0

Pathophysiology-Inflammation

Faculty of Pharmaceutical Sciences

Dr. Amjaad Zuhier Alrosan, Dr. Abdelrahim Alqudah

Inflammation is a protective response of vascularized tissues to infections and tissue damage that brings cells and molecules of host defense from the circulation to the sites where they are needed, to eliminate the offending agents.

الخلايا المصابة
الجهاز المناعي

شئ ضروري دائماً تكون النهاية انه يرجع الى الهربان
مرات يصير بالعدوى cell death

INFLAMMATION

STIMULI

e.g. PATHOGEN

بكتيريا
فيروسات



e.g. TOXINS & TRAUMA

مواد سامة
لوعنبرية



GOAL is RESPOND to STIMULI and RESTORE BALANCE

ELIMINATE CAUSE

CLEARING OUT NECROTIC CELLS

TISSUE REPAIR

انما حاربنا cell death
المسبب بتخلص
سببها و الخلايا بتصر
تعمل انقسامات حتى
تعودن الخلايا
ولكن في معظم الاميان
يصير زبي الذببة وصعب انه يعمل tissue (Fibrous scar)

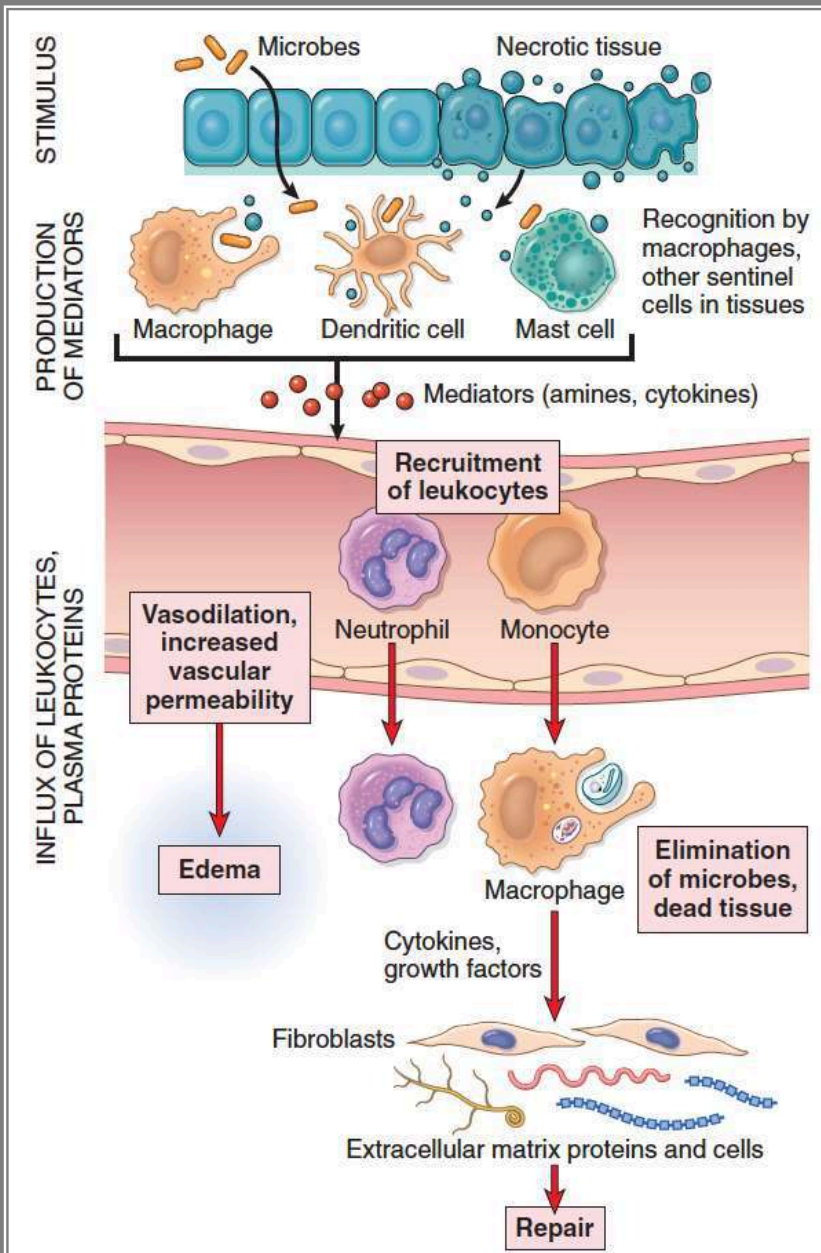


Fig. 3.1 Sequence of events in an inflammatory reaction. Macrophages and other cells in tissues recognize microbes and damaged cells and liberate mediators, which trigger the vascular and cellular reactions of inflammation. Recruitment of plasma proteins from the blood is not shown.

repair

PAMPS ← external كائنات *
(pathogen associated molecular patterns)

DAMPs ← internal كائنات *
Damage associated molecular patterns

DAMPs

released from damaged or dying cells and activate the innate immune

PAMPS

derived from microorganisms and recognized by pattern recognition receptor

Inflammation

② external stimuli factor
(pathogen/ toxins)

① internal stimuli factor
(cell damage) trauma

طريقة الاستجابة تختلف باختلاف stimuli

Causes of inflammation

Infections (bacterial, viral, fungal, parasitic) are the *most common* medically important causes of inflammation.

Tissue necrosis: from any cause, including ischemia

Trauma: blunt & penetrating

تكون
فجائية
وحادة

Physical agents: thermal injuries-burns/frostbite;
radiation

Chemicals: Agents as strong acids, alkalis

Foreign bodies: splinters, dirt, & sutures

Immune = hypersensitivity reactions:
types I, II, III, & IV,
against environmental substances or against self tissue

The typical inflammatory reaction develops through a series of sequential steps:

- The offending agent, which is located in extravascular tissues, is recognized by host cells and molecules.
- Leukocytes and plasma proteins are recruited from the circulation to the site where the offending agent is located.
- The leukocytes and proteins are activated and work together to destroy and eliminate the offending substance.
- The reaction is controlled and terminated.
- The damaged tissue is repaired.

LEUKOCYTES

GRANULOCYTES

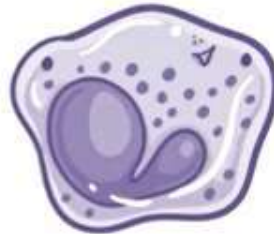
NEUTROPHILS



EOSINOPHILS



BASOPHILS



MAST CELLS

AGRANULOCYTES

LYMPHOCYTES



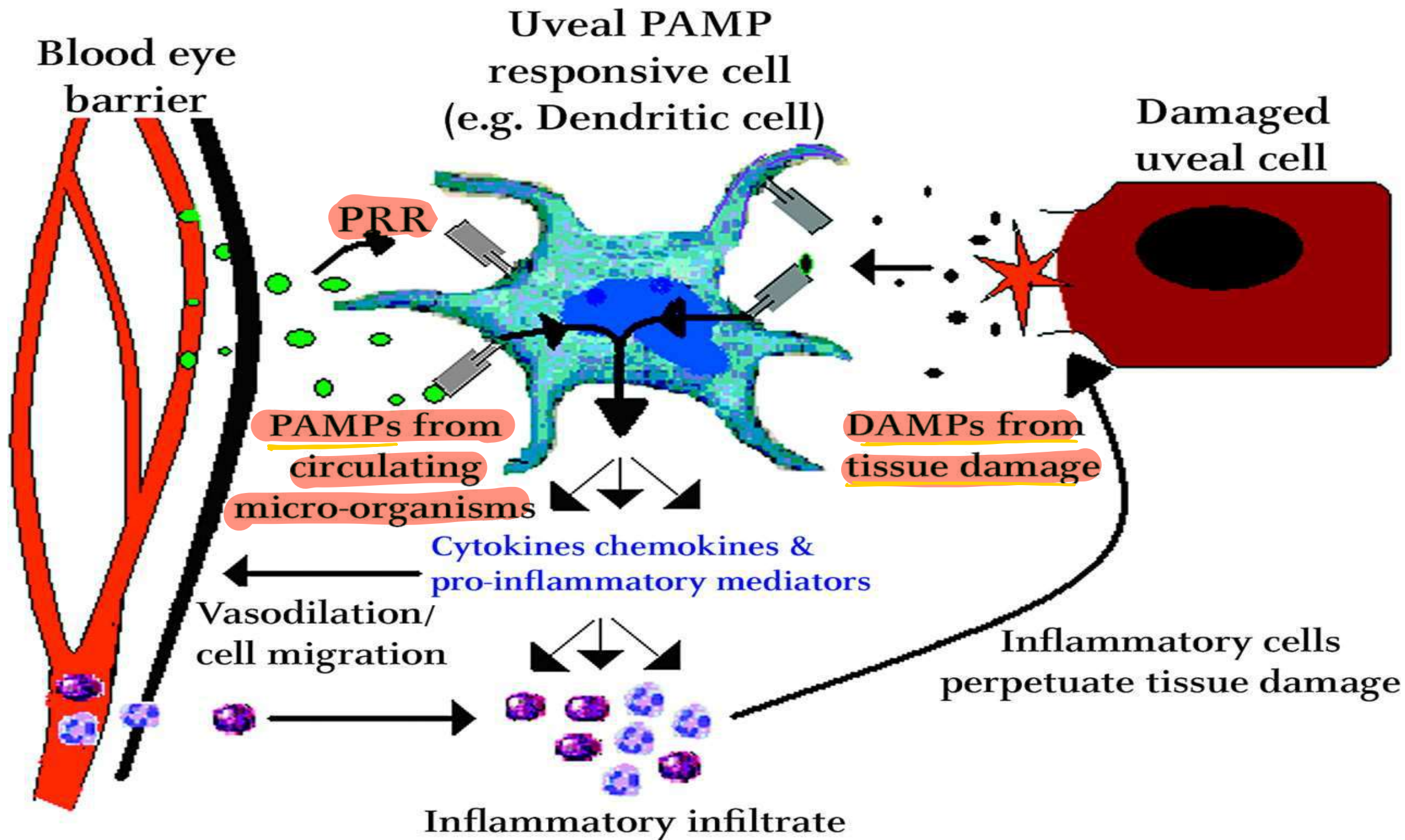
MONOCYTES



MACROPHAGES

DENDRITIC CELLS





histamine في فقط في عنا =
vasodilation ما يزيد فقط في عنا =

plasma protein factor
(clotting factor) / xIIa

↓
يحل كل تحفيز
kinin path wa

يحل كل إنتاج
brady kinin → vasodilation
↓
يغير بالهياج ما يكون
في histamine فقط
كان ينتج نتيجة تحفيز xIIa

(PAMPs)
↑
يقول عنا البكتيريا يكون عليها (antigen → endotoxin)
↓
mast cell يكون عليها receptors (PRR)
↓
mast cell (PRR) يعمل (recognition) لـ antigen الموجود
عنا البكتيريا

ال (endotoxin) يعمل تحفيز لـ receptors الموجودة
عنا mast cells

mast cell فيها → inflammatory mediators
↑
زيادة عن بهه
علية
inflammation
↓
له كل واحد سونول
عن سيمعنين (التاثيريكلن)
↑
نتيجة التحفيز بتصير تفرز
حل
histamine

عندما يحدث إصابة (injury) للخلية البطانية (Endothelial cells) في جدار

يتم إفراز بروتينات الالتصاق (adhesive substances) مثل Selectins على الخلايا البطانية
التي لها القدرة على الالتصاق بالخلايا المناعية
التي تهاجر نحو موقع الإصابة

تتجه الخلايا (Neutrophils) و (Monocytes) لتلتصق بالخلايا
التي تسمى action site

هنا يحدث تمايز (differentiation) للخلايا من monocytes إلى macrophages
التي تلتصق بـ selectins

macrophages و neutrophils تتعرفان على selectins
في جدار جدار جدار الخلية البطانية
التي تسمى rolling

عندما يحدث إصابة تنتج الخلية البطانية
مادة تسمى Leukin في الدم

في الدم يكون فيه microbes ويتجه macrophages
و neutrophils للالتصاق بـ selectins على الخلايا
البطانية

في الدم يحدث contact مع الخلايا البطانية

في الدم يحدث التصاق macrophages و neutrophils
مع microbes في الدم

التصاق الخلايا البطانية مع microbes
يحدث في البداية بـ rolling
التي تكون مرتبطة مع endothelial cells
في البداية وهذا الثلاثة يمكنها التصاق
الخلايا البطانية مع microbes

عندما يحدث إصابة
يتم إفراز بروتينات الالتصاق
التي تسمى rolling = margination

عندما يحدث إصابة

plasma membrane damage/injury

يتم تفعيل إنزيم

phospholipase A2

يحلل phospholipid إلى
arachidonic acid

يترجم إلى
inflammatory mediator
تنتجها خلايا mast cells

← بعد عملية Rolling

Leukocytes بغض النظر عن نوعها

يصير تطلع عن طريق gap بين وجوده
endothelial cells



بتصير تعمل (Squeezing)

بغض زكي بتختر حالي بين gap حتى
تطلع لا other side

كانه بالنهاية هاي Leukocytes بدعا تكرر
الجزئيات يلي جواتها وتخلص اي waste
ويصير لها excretion عن طريق kidneys
وتطلع عن طريق (urinary system)

هاي العملية ما بتكون محتايبة حتى ان تصير
لازم ان Leukocytes ترتبط مع substance
اسم (PECAM-1) وبعدين تحركها وتطلع

هاي العملية نسييرها **Diapedesis**

* positive chemotaxis / margination / diapedesis *



very fast

← ولكن بعد حاي الخطوات بتصير تاخذ وقتة اطول

leukocytosis - inducing factors

pro-inflammatory cytokines
انتاج كميات كبيرة من leukocytes بسبب اغزاز

عدوى تحفز hypothalamus الى انتاج

prostaglandin E2 ← هو السبب انه يصيرنا
حرارة (fever)

TNF-α و IL-1
de bone marrow
انتاج
white blood cells
red blood cells

يخرج (IL-1) و (TNF-α) ياثر على liver

C reactive peptide
ينتج

↓
يخفضه كتحص للتخفيف
by inflammation

positive chemotaxis



خلية حبيبي

neutrophils / Monocytes

الى منطقة

mast cells

نتيجة اغزاز inflammatory
mediators

خلال حاي العلية ال

macrophage و neutrophils

بتبلش تفرز

inflammatory mediators



pro-inflammatory
cytokines

Inflammatory Cascade: Pathogenesis and clinical findings

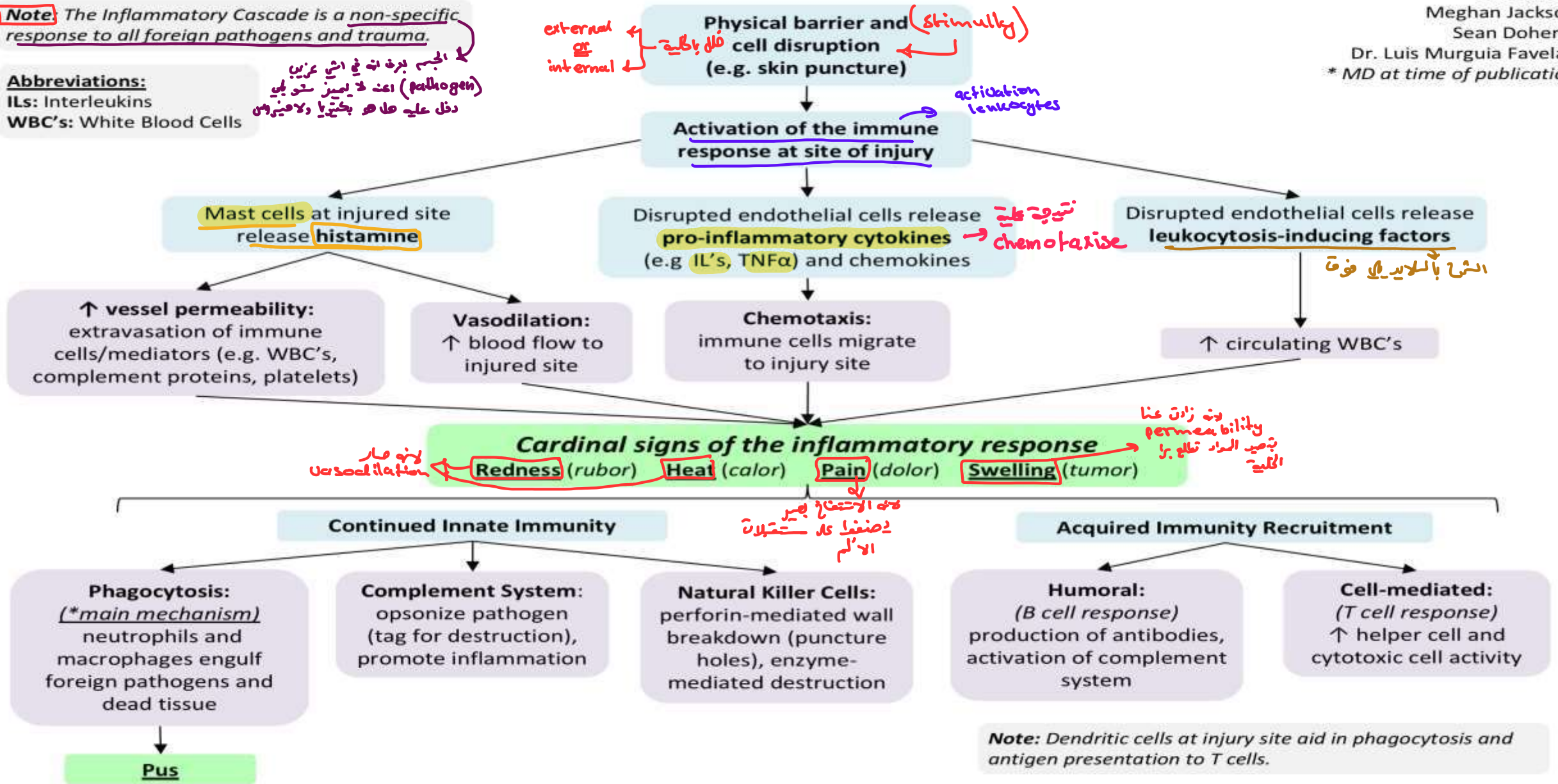
(الشكل رسم)

Authors:
Heather Yong
Reviewers:
Meghan Jackson
Sean Doherty
Dr. Luis Murguia Favela*
* MD at time of publication

Note The Inflammatory Cascade is a non-specific response to all foreign pathogens and trauma.

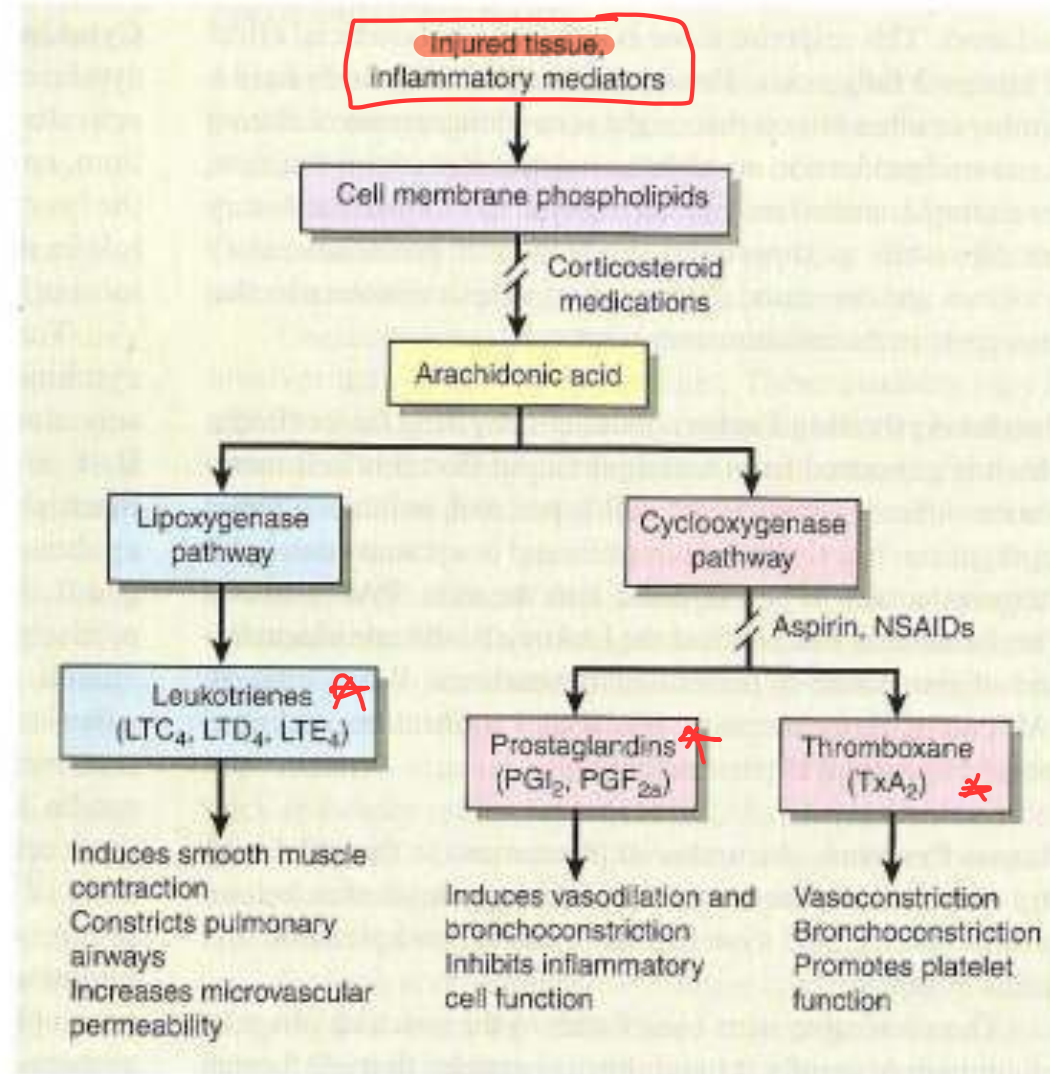
Abbreviations:
ILs: Interleukins
WBC's: White Blood Cells

الجسم يرد انه في اشي غريب (pathogen) اعنه لا يميز شو بين دقل عليه طاهو بختريا ولا فيروس

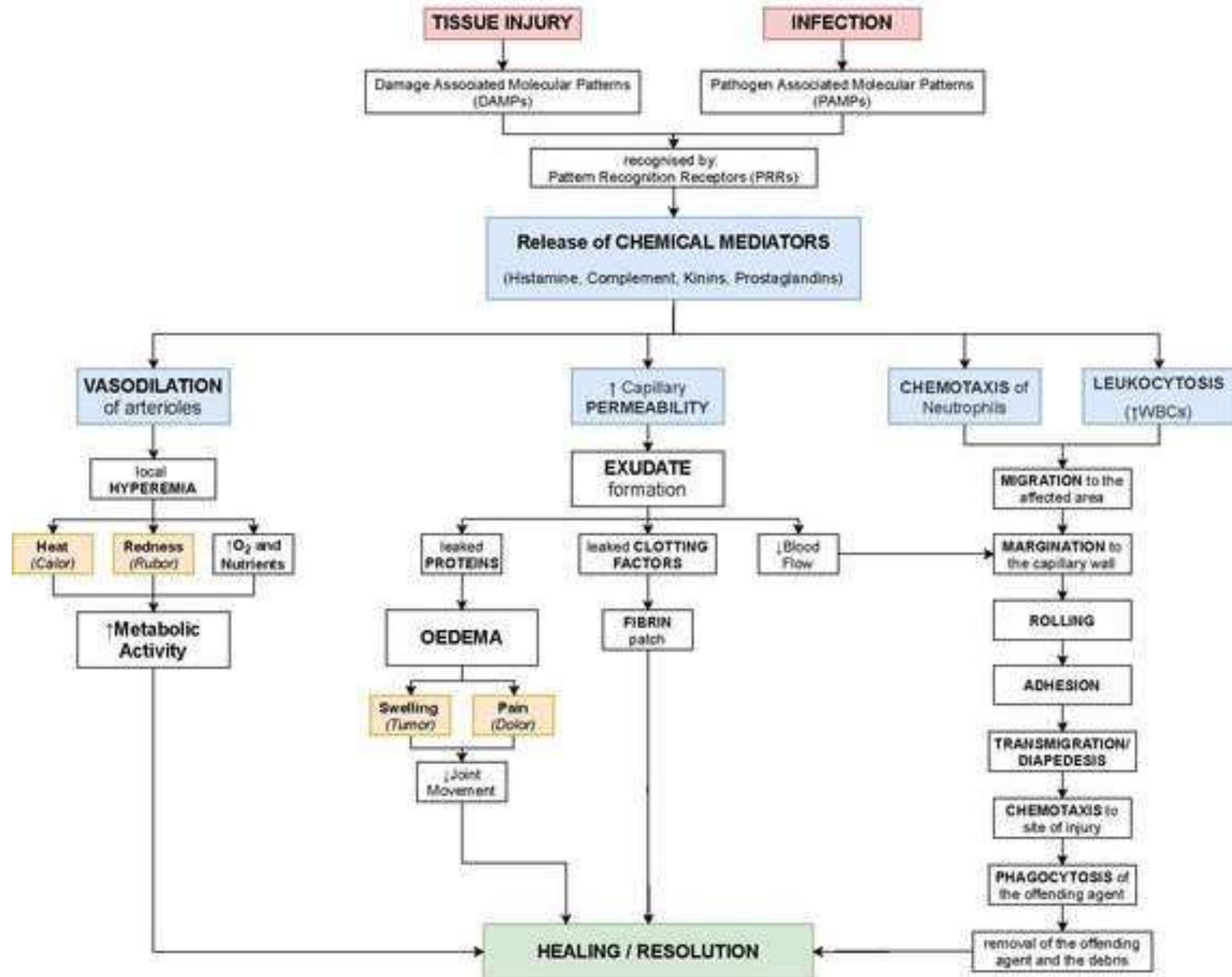


Note: Dendritic cells at injury site aid in phagocytosis and antigen presentation to T cells.

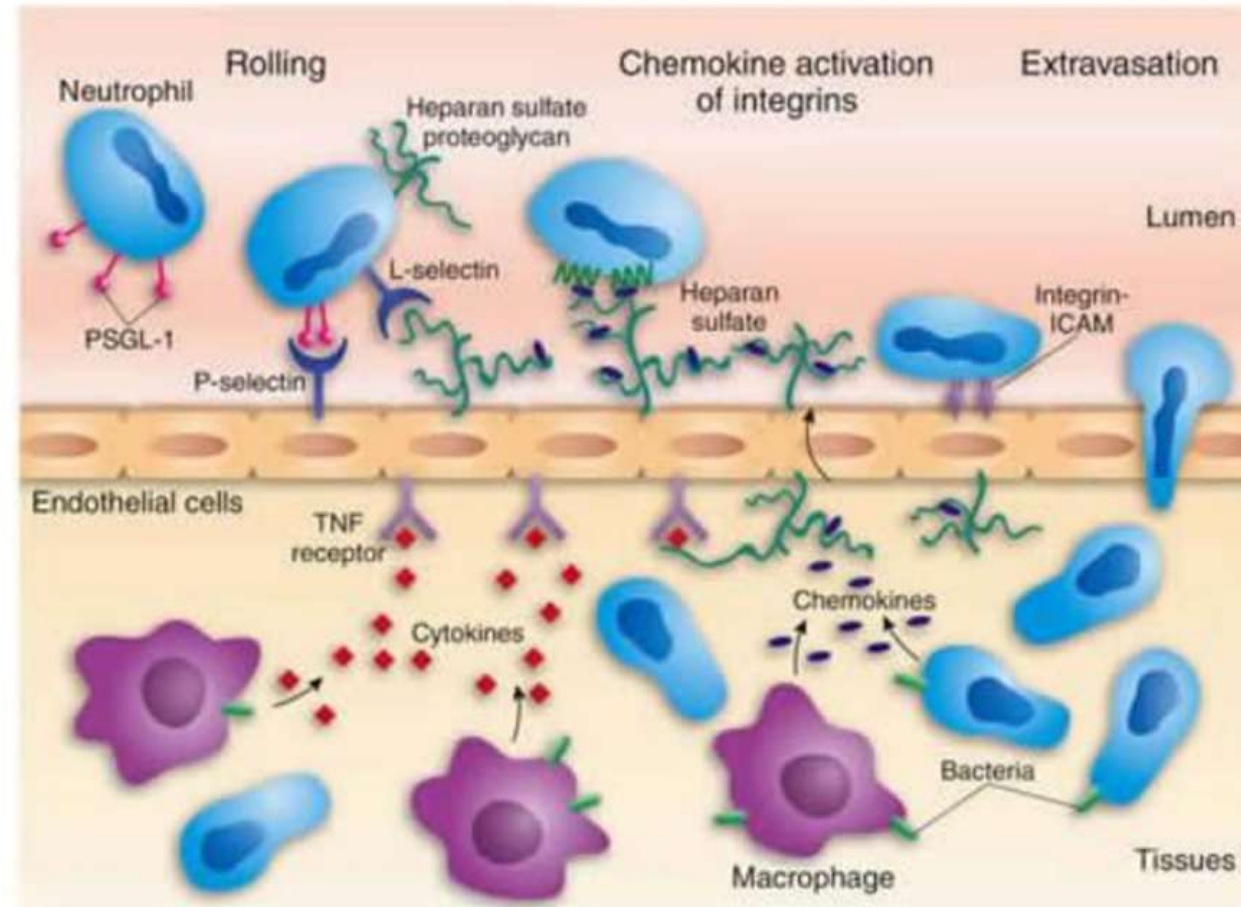
Arachidonic Acid Metabolites



* → Chem mediators



Neutrophils



The cardinal signs of inflammation

Heat (Calor)

Redness (Rubor)

Swelling (Tumor)

Pain (Dolor)

Loss of function (Functio laesa)

English	Latin	Cause
Heat	<i>Calor</i>	Vasodilation
Redness	<i>Rubor</i>	Vasodilation
Swelling	<i>Tumor</i>	Increased vascular permeability Increased granulation tissue
Pain	<i>Dolor</i>	Physical and chemical stimulation of nociceptors
Loss of function	<i>Functio laesa</i>	Pain Reflex muscle inhibition Disruption of tissue structure Fibroplasia and metaplasia

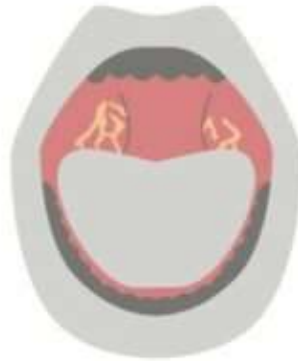
The cardinal signs of inflammation



Heat



Redness



Swelling



Pain



Loss of function

Due to increased
blood flow to site
of inflammation

Due to
accumulation
of fluid and cells

Due to stimulation of
sensory neurons by
inflammatory
mediators

Due to tissue
damage

Types of inflammation

تبدأ في وقت
مخف ح الوقت
لا لم تتم حاجت
Chronic يصير

Acute inflammation: Is the inflammatory response which is sudden in onset, of short duration, and is characterized by the classical *cardinal signs* of inflammation, in which the vascularized and exudative process predominates.

Chronic inflammation: Is an inflammation of a slow progress, marked chiefly by the formation of new connective tissue, it may be a continuation of an acute form or a prolonged low-grade form and usually causes permanent tissue damage.

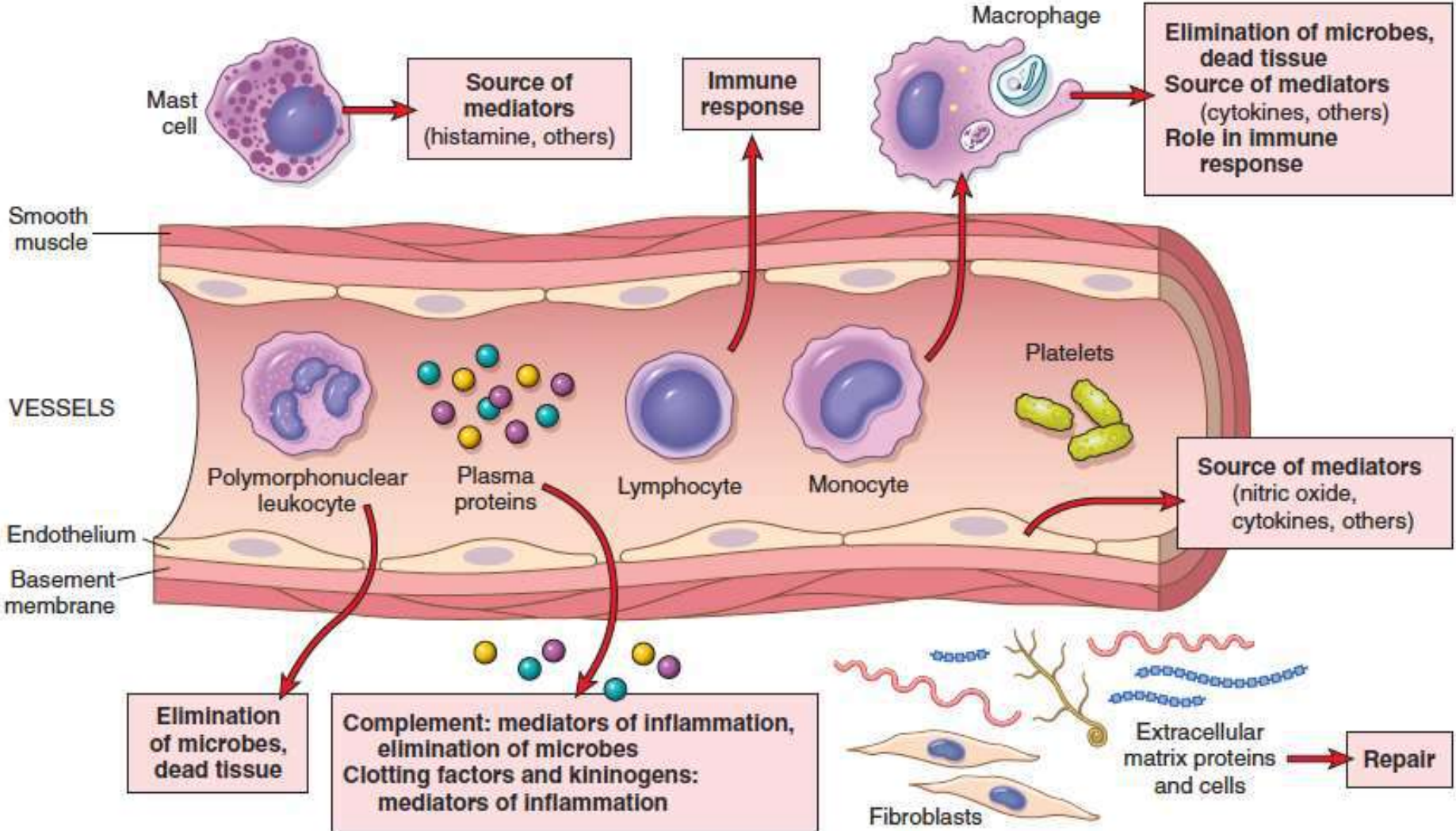
Table 3.1 Features of Acute and Chronic Inflammation

Feature	Acute	Chronic
Onset طريق	<u>Fast: minutes or hours</u>	<u>Slow: days</u>
Cellular infiltrate	Mainly neutrophils	Monocytes/macrophages and lymphocytes
Tissue injury, fibrosis	Usually mild and self-limited	May be severe and progressive
Local and systemic signs	Prominent	Less

Characteristics of inflammations

	Acute	Chronic
Cause	Single injury	Permanent present of the causing agent /bacteria, etc./
Duration	Hours, days	Weeks, months, years; depending on the causing agent
Presentative symptom	↑ permeability, exudation	Proliferative fibroblasts No exudation
Main components in the process	Liquid Proteins /proteases and antiproteases/ PMN leukocytes Macrophages	Macrophage Lymphocytes Eosinophyl granulocytes Connective tissue hiperplasy
Connecting reactions	Thrombosis	Immune response

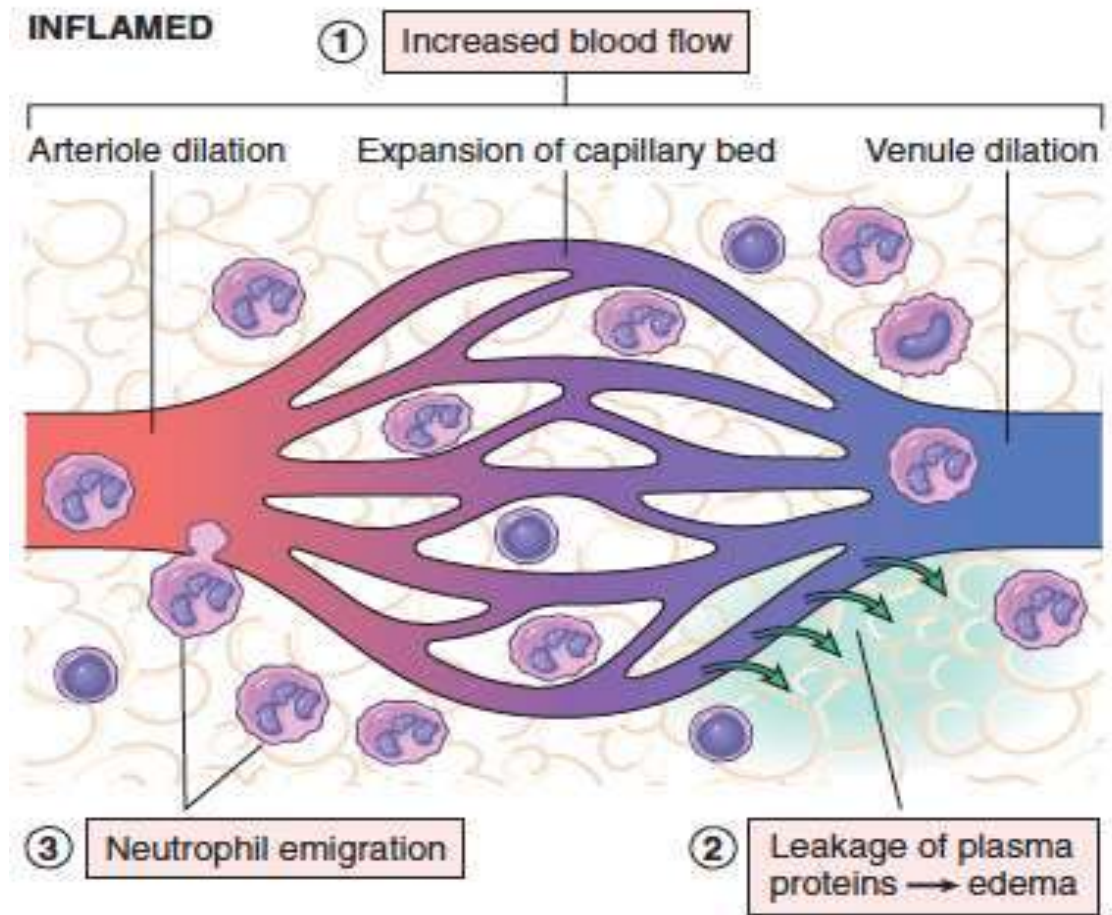
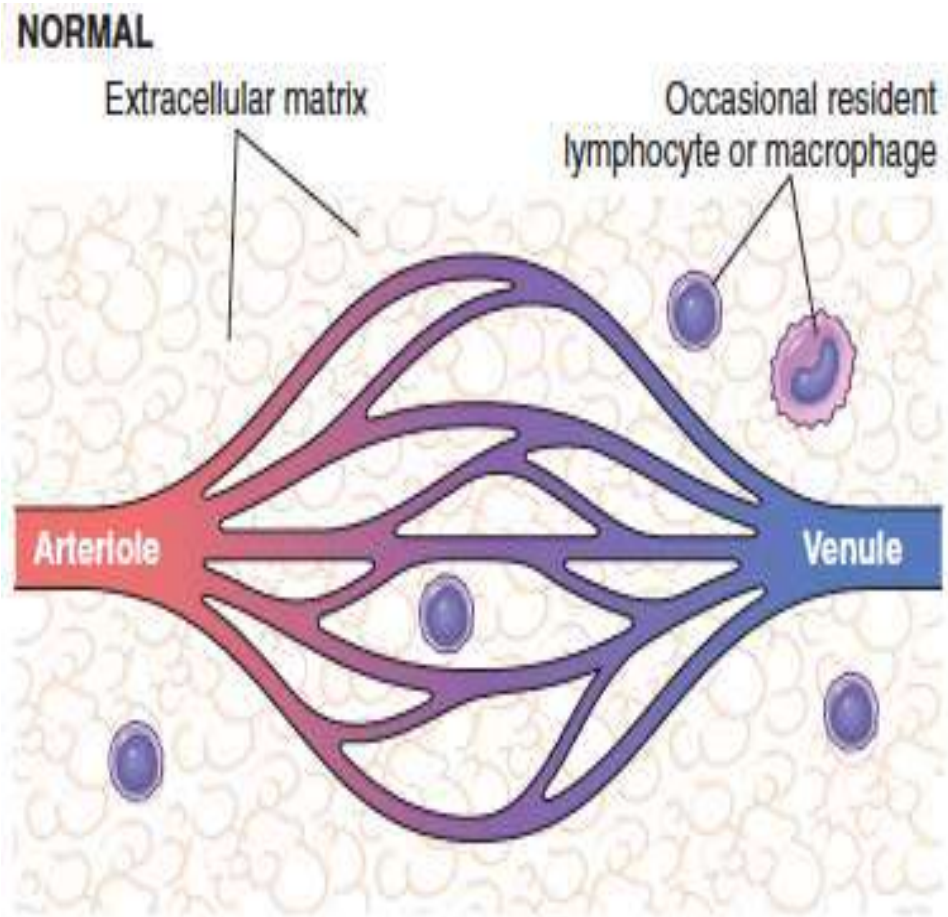
Components of Inflammatory Response



Components of Inflammatory Response

- 1. Circulating bone marrow-derived cells:** include leukocytes, neutrophils, eosinophils, basophils, lymphocytes, monocytes, & platelets.
- 2. Circulating proteins:** include clotting factors, kininogens, & complement components, all are synthesized by the liver.
- 3. Vascular wall cells:** include endothelial cells (EC), and the underlying smooth muscle cells (SMC).
- 4. Connective tissue cells:** include mast cells, macrophages & lymphocytes; to Phagocytose and the fibroblasts that synthesize the extracellular matrix (ECM).
تول tissue repair !عنه
- 5. The extracellular matrix (ECM):** consist of fibrous structural proteins (e.g., collagen & elastin).

Acute Inflammation



Major Events In Inflammation

1. Vasodilation causing increased blood flow (warmth and erythema)
2. Extravasation and leakage of plasma fluids & proteins (edema)
3. Emigration of the leukocytes from and their accumulation at the site of injury

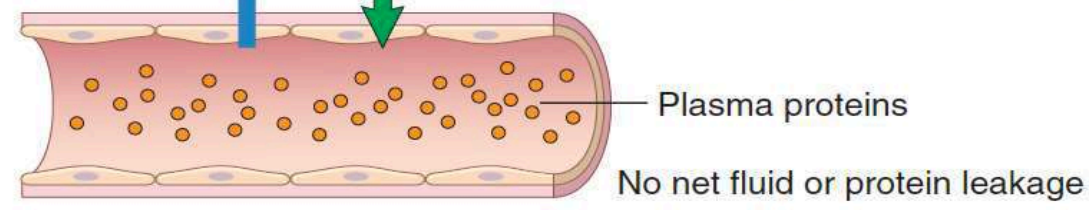
Vasodilation

→ blood flow
الى المنطقة بالي فيها التهاب

↓
يكون نتيجة
نزق الضغط

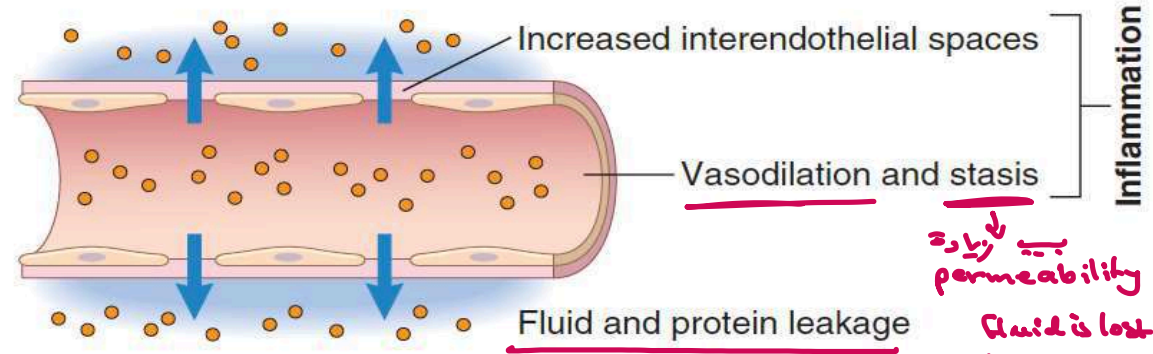
نتفه
Fluid pressure
Hydrostatic pressure = Colloid osmotic pressure

A. NORMAL



B. EXUDATE

high protein content, and
may contain some white
and red cells



C. TRANSUDATE

low protein content, few cells)

→ Increased hydrostatic pressure
(venous outflow obstruction,
[e.g., congestive heart failure])

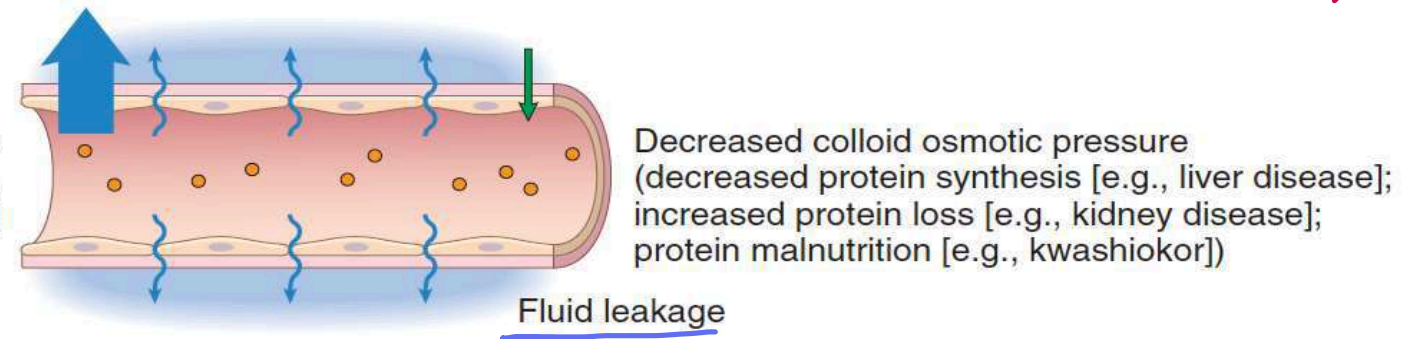


Fig. 3.2 Formation of exudates and transudates. (A) Normal hydrostatic pressure (blue arrow) is about 32 mm Hg at the arterial end of a capillary bed and 12 mm Hg at the venous end; the mean colloid osmotic pressure of tissues is approximately 25 mm Hg (green arrow), which is equal to the mean capillary pressure. Therefore, the net flow of fluid across the vascular bed is almost nil. (B) An exudate is formed in inflammation because vascular permeability increases as a result of retraction of endothelial cells, creating spaces through which fluid and proteins can pass. (C) A transudate is formed when fluid leaks out because of increased hydrostatic pressure or decreased osmotic pressure.

microbes يتصير ← بالناي بصير ← لكن مرات حكن انها ما تفل ← macrophages رولا كليه بكون catching ← micrope تامل catching لـ Neutrophils

lymphatic circulation

Responses of Lymphatic Vessels

- In addition to blood vessels, lymphatic vessels also participate in the inflammatory response.

- In inflammation, lymph flow is *increased and helps drain edema fluid*, leukocytes, and cell debris from the extravascular space.

- In severe inflammatory reactions, especially to microbes, the lymphatics may transport the offending agent, contributing to its dissemination. The lymphatics may become secondarily inflamed (*lymphangitis*), as may the draining lymph nodes (*lymphadenitis*).



↓
اذصار بار
lymphatic vessels
↓
اذصار بار
lymph nodes

Mechanisms of increased vascular permeability

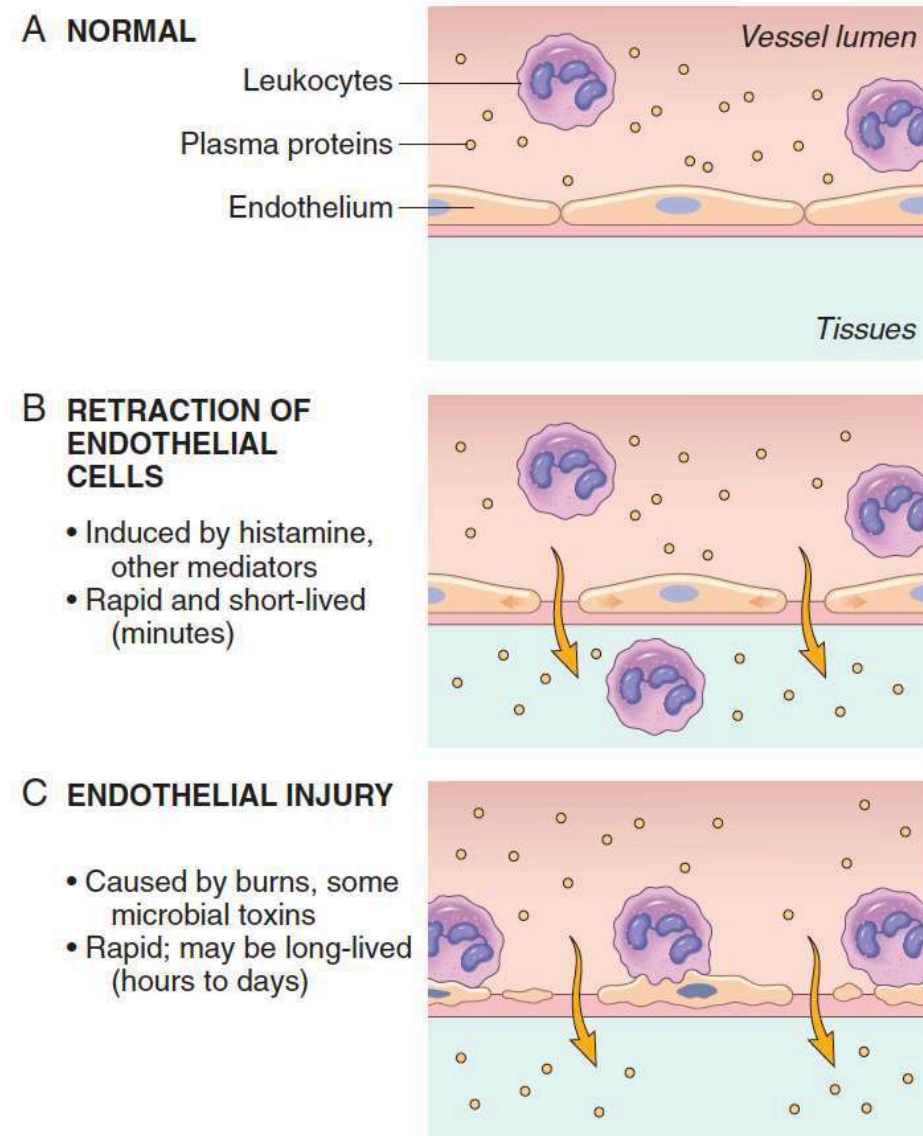


Fig. 3.3 Principal mechanisms of increased vascular permeability in inflammation and their features and underlying causes.

Increased Vascular Permeability

تسمى permeability بغير بنوعه المقدار

• There are three basic patterns of increased permeability:

- ① **Immediate transient response**: ^{قصيرة} Histamine-type.
- ② **Immediate-sustained response**: ^{اطول} which is seen in severe injury and is associated with damage to endothelial cells.
- ③ **Delayed prolonged response**: ^{اطول بزيادة} as in hypersensitivity type IV & exposure to ultraviolet light.

Leukocyte Recruitment to Sites of Inflammation

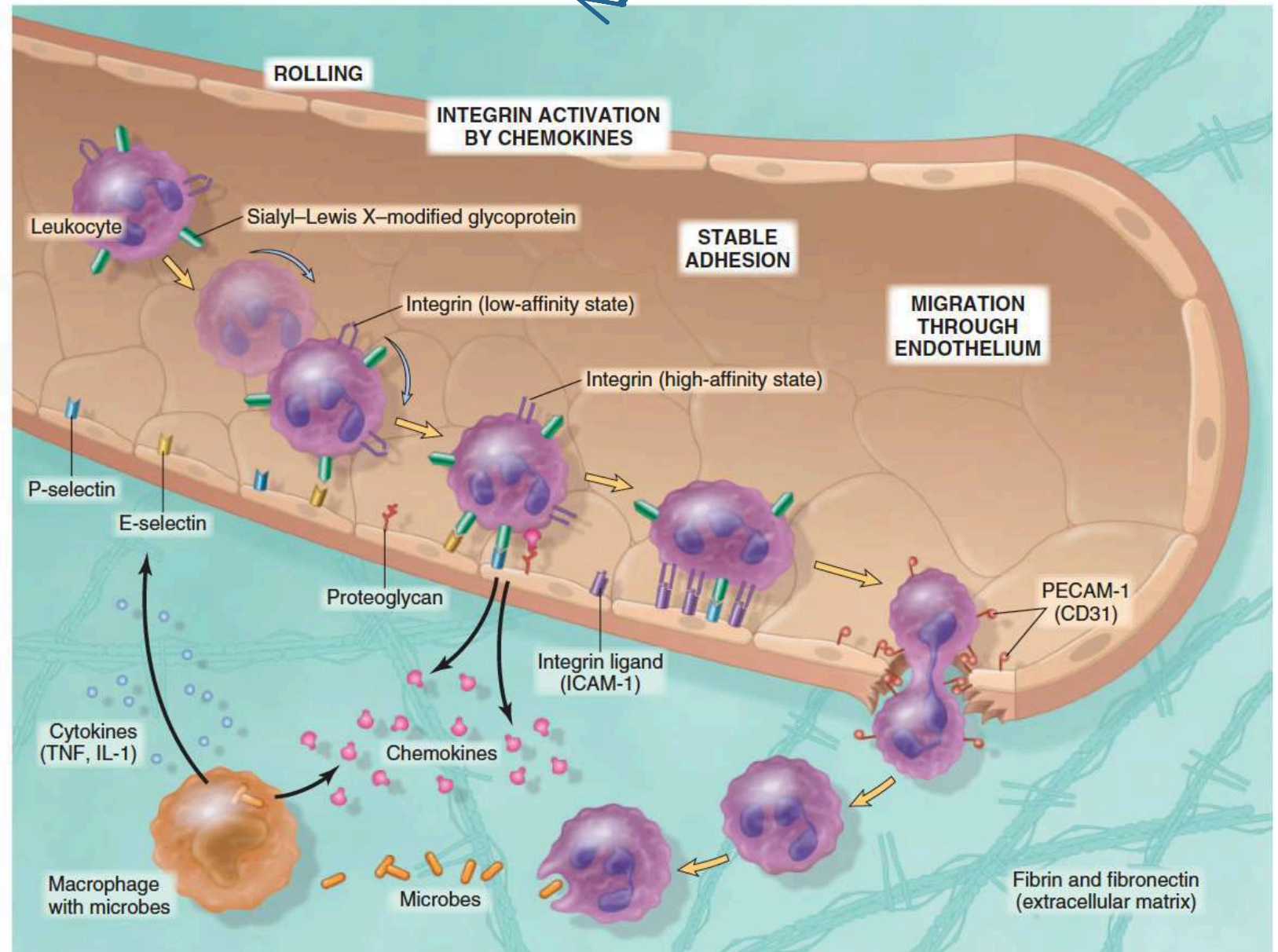
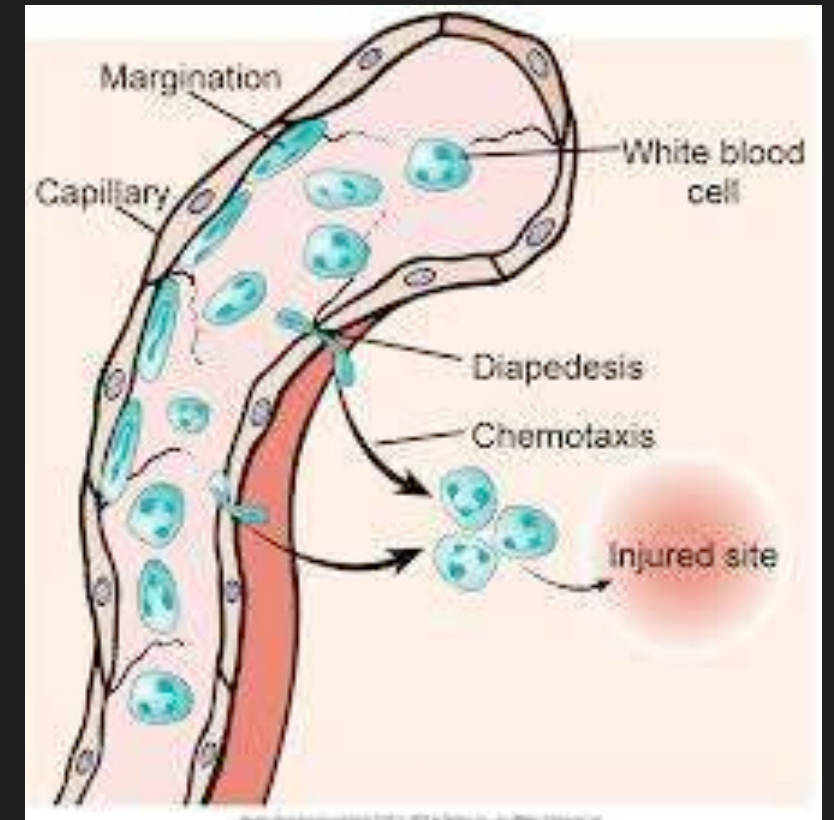


Fig. 3.4 The multistep process of leukocyte migration through blood vessels, shown here for neutrophils. The leukocytes first roll, then become activated and adhere to endothelium, then transmigrate across the endothelium, pierce the basement membrane, and move toward chemoattractants emanating from the source of injury. Different molecules play predominant roles at each step of this process: selectins in rolling; chemokines (usually displayed bound to proteoglycans) in activating the neutrophils to increase avidity of integrins; integrins in firm adhesion; and CD31 (PECAM-1) in transmigration. *ICAM-1*, Intercellular adhesion molecule-1; *PECAM-1 (CD31)*, platelet endothelial cell adhesion molecule-1; *TNF*, tumor necrosis factor.

• The sequence of events in the recruitment of leukocytes from the vascular lumen to the extravascular space consists of:

- margination and rolling along the vessel wall;
- firm adhesion to the endothelium;
- transmigration between endothelial cells; and
- migration in interstitial tissues toward a chemotactic stimulus

نقطة
diapedesis

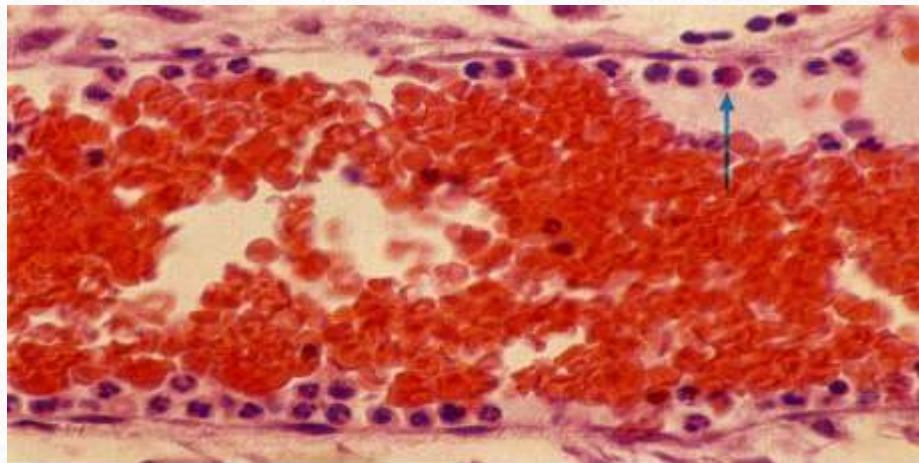
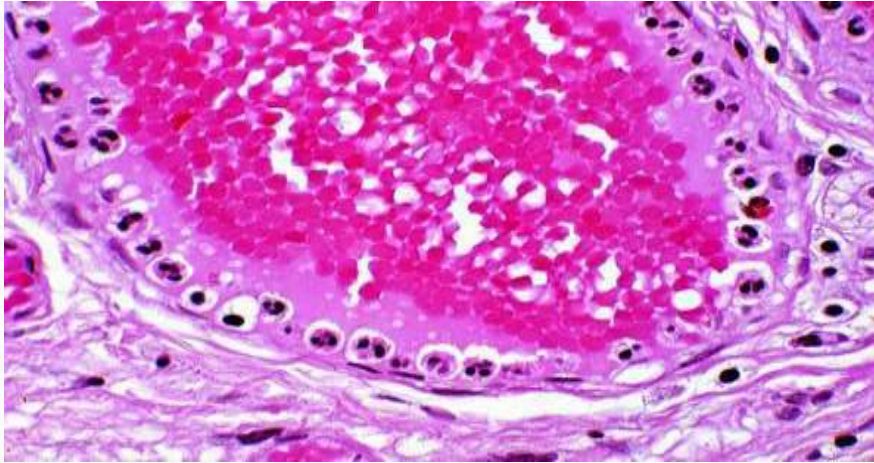


Mediators of cellular rolling and adhesion

- 1-The **selectins** which consist of
 - **E-selectin** is confined to the endothelial cells.
 - **P-selectin** is present on the platelets.
 - **L-selectin** is present on leukocytes, especially lymphocytes, it adheres lymphocytes to the endothelium.
- 2-The **immunoglobulin family molecules**: include two endothelial adhesion molecules:
 - An intercellular adhesion molecule 1 (**ICAM-1**) & Vascular cell adhesion molecule-1 (**VCAM-1**), both molecules interact with integrins found on leukocytes.
يعبر عن ارتباط الخلايا مع endothelium
- 3-The **integrins & mucin-like glycoprotein**: are transmembrane-adhesive glycoproteins that also function as receptors for the extracellular matrix.

(1) Leukocyte Margination and Rolling

**endothelium lined by WBC an appearance called *pavementing*

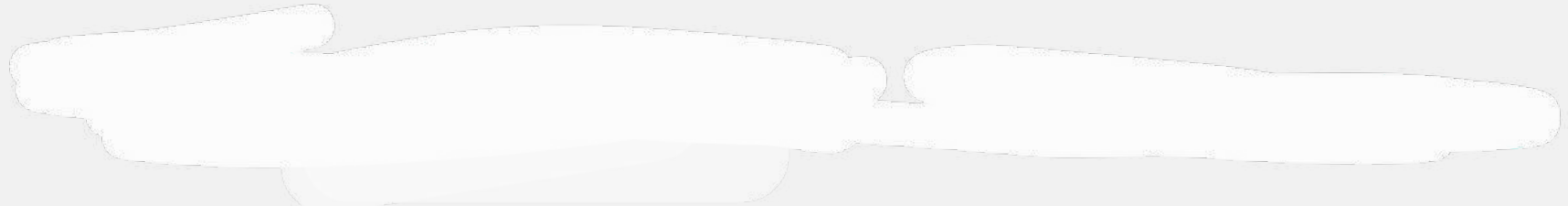


(2) Adhesion

- The rolling leukocytes are able to sense changes in the endothelium that allows them to adhere firmly to endothelial surfaces, this process is mediated by integrins expressed on leukocyte cell surfaces interacting with their ligands (selectins) on endothelial cells.

(3) Transmigration (Diapedesis)

• After being arrested on the endothelial surface, leukocytes migrate through the vessel wall primarily by squeezing between cells at intercellular junctions.



Phagocytosis

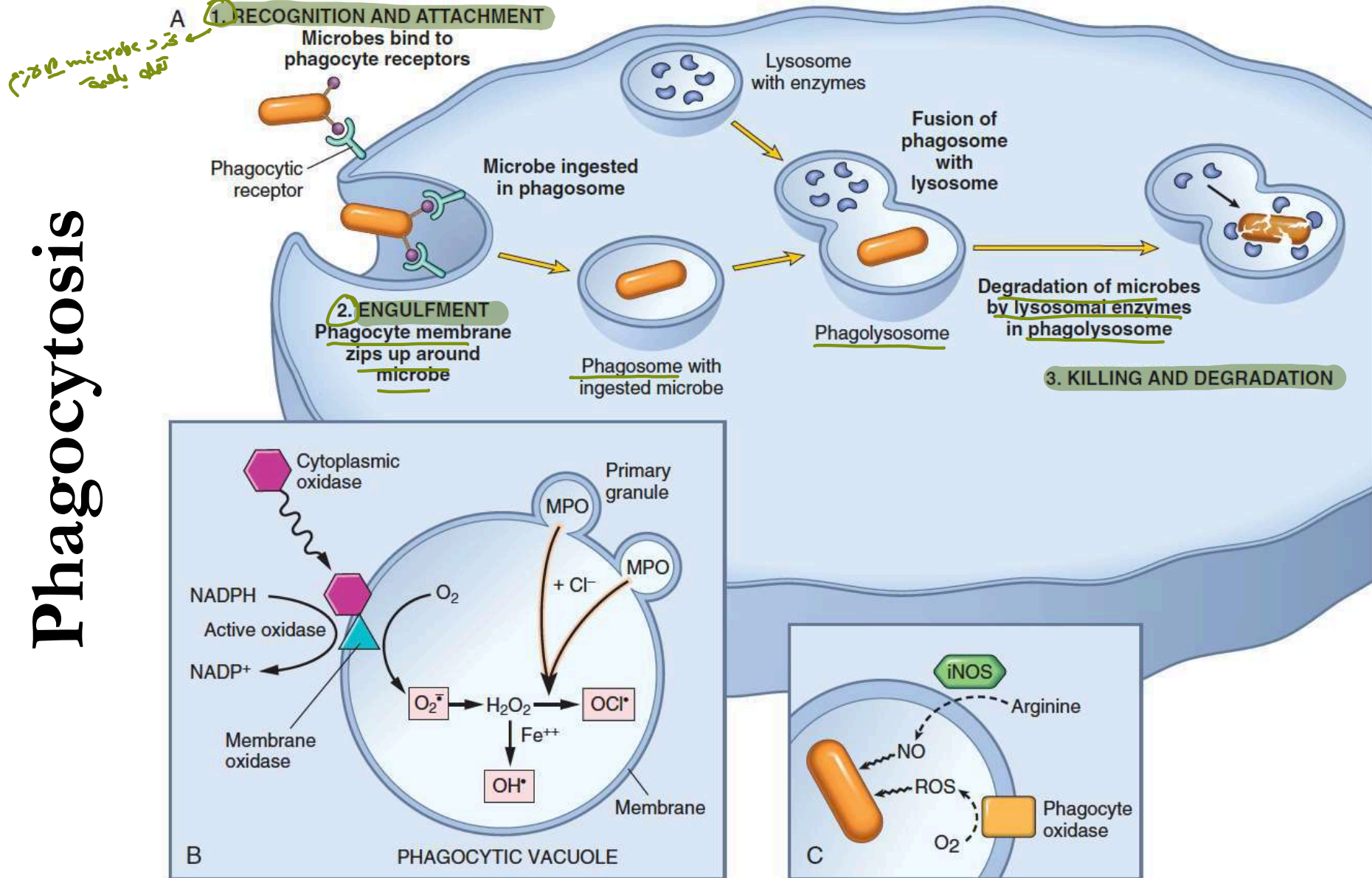


Fig. 3.7 Phagocytosis and intracellular destruction of microbes. (A) Phagocytosis of a particle (e.g., a bacterium) involves binding to receptors on the leukocyte membrane, engulfment, and fusion of the phagocytic vacuoles with lysosomes. This is followed by destruction of ingested particles within the phagolysosomes by lysosomal enzymes and by reactive oxygen and nitrogen species. (B) In activated phagocytes, cytoplasmic components of the phagocyte oxidase enzyme assemble in the membrane of the phagosome to form the active enzyme, which catalyzes the conversion of oxygen into superoxide (O_2^-) and H_2O_2 . Myeloperoxidase, present in the granules of neutrophils, converts H_2O_2 to hypochlorite. (C) Microbicidal reactive oxygen species (ROS) and nitric oxide (NO) kill ingested microbes. During phagocytosis, granule contents may be released into extracellular tissues (not shown). *iNOS*, Inducible NO synthase; *MPO*, myeloperoxidase; ROS, reactive oxygen species.

Chemical Mediators

Chemical substances are derived from either *plasma* or the *tissues*, they act as a link between the occurrence of injury and the onset of inflammation.

They are divided into two major categories :

• *Plasma Derived-Mediators*, these include:

• بتكون موجودة
في
plasma

1. The kinin system.
2. The complement system.
3. The coagulation & fibrinolytic system.
Coagulation

• بتكون موجودة
في
الخلايا

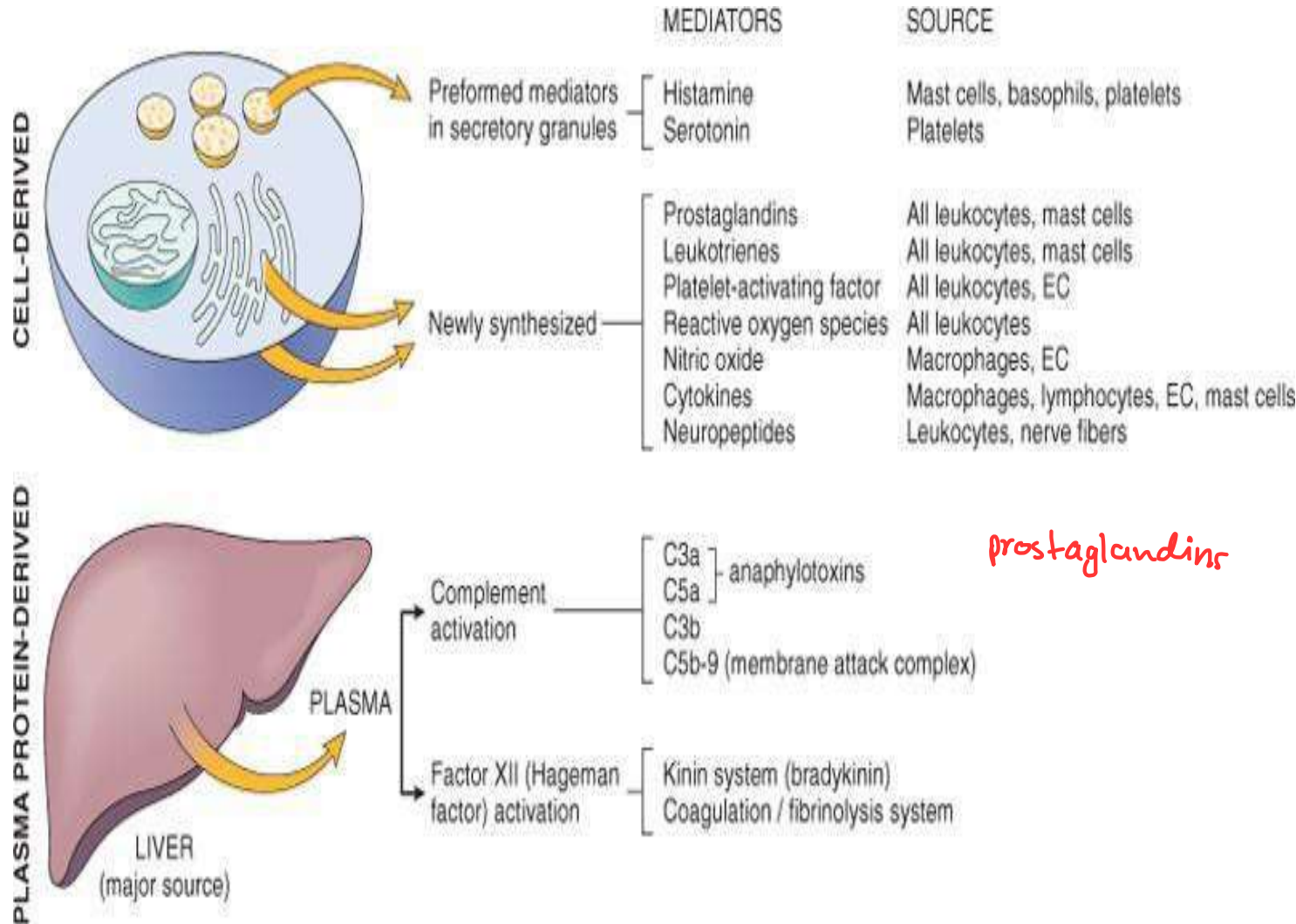
• *Cell derived mediators*; these include:

- circulating platelets, basophils, endothelial cells monocyte/macrophages, tissue mast cells and the injured tissue itself are all potential cellular sources of vasoactive mediators.

Chemical Mediators

30

Table 3.5 Principal Mediators of Inflammation



Mediator	Source	Action
Histamine	Mast cells, basophils, platelets	Vasodilation, increased vascular permeability, endothelial activation
Prostaglandins	Mast cells, leukocytes	Vasodilation, pain, fever
Leukotrienes	Mast cells, leukocytes	Increased vascular permeability, chemotaxis, leukocyte adhesion, and activation
Cytokines (TNF, IL-1, IL-6)	Macrophages, endothelial cells, mast cells	Local: endothelial activation (expression of adhesion molecules). Systemic: fever, metabolic abnormalities, hypotension (shock)
Chemokines	Leukocytes, activated macrophages	Chemotaxis, leukocyte activation
Platelet-activating factor	Leukocytes, mast cells	Vasodilation, increased vascular permeability, leukocyte adhesion, chemotaxis, degranulation, oxidative burst
Complement	Plasma (produced in liver)	Leukocyte chemotaxis and activation, direct target killing (membrane attack complex), vasodilation (mast cell stimulation)
Kinins	Plasma (produced in liver)	Increased vascular permeability, smooth muscle contraction, vasodilation, pain

Major roles of cytokines in acute inflammation

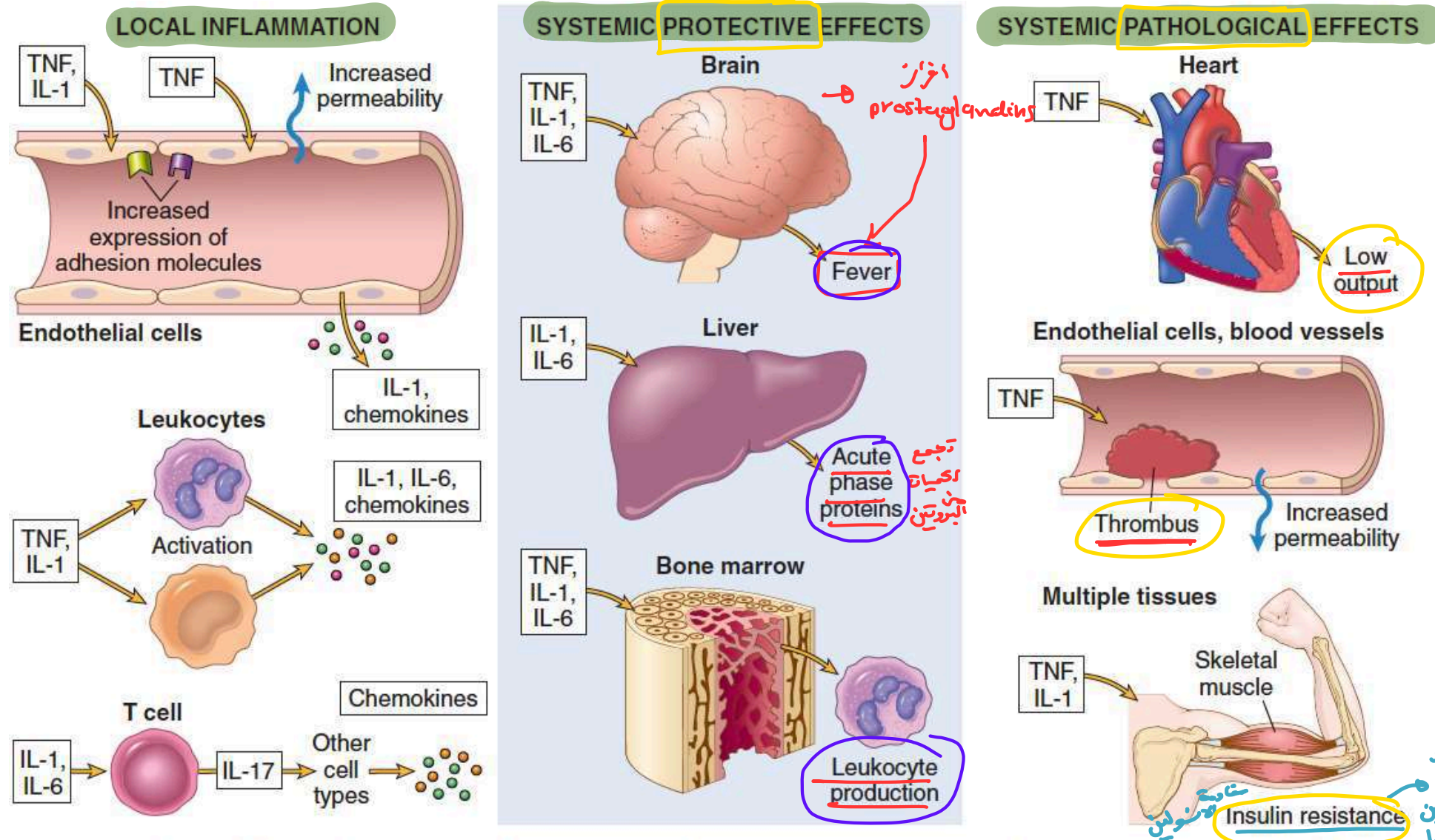


Fig. 3.10 Major roles of cytokines in acute inflammation. PDGF, Platelet-derived growth factor; PGE, prostaglandin E; PGI, prostaglandin I.

تصدير الجلوكوز
 قدرة على
 استهلاك الأنسولين
 وإدخاله إلى الخلايا
 بالتالي يمكن يصير
 hyperglycemia

The complement system

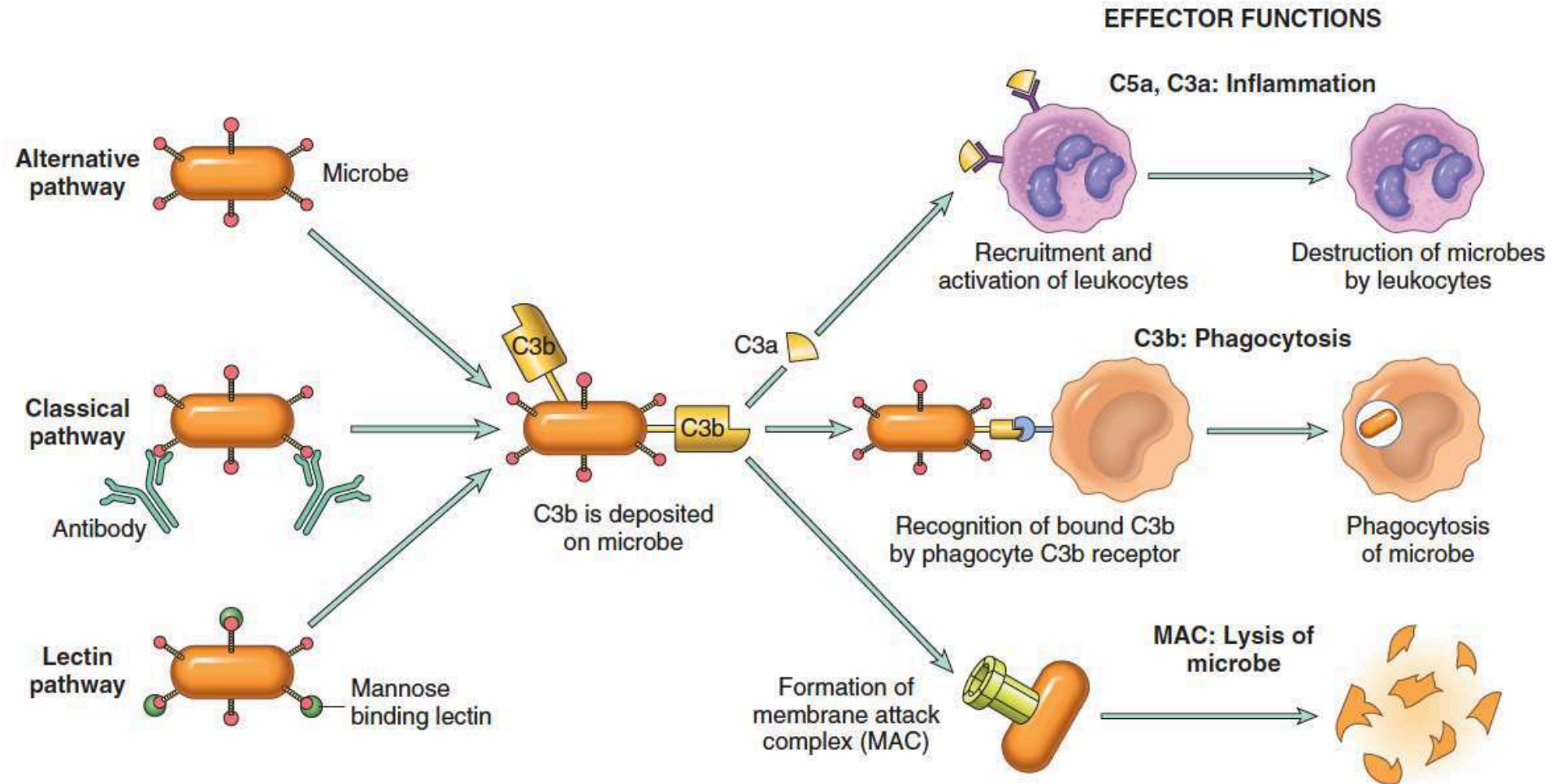
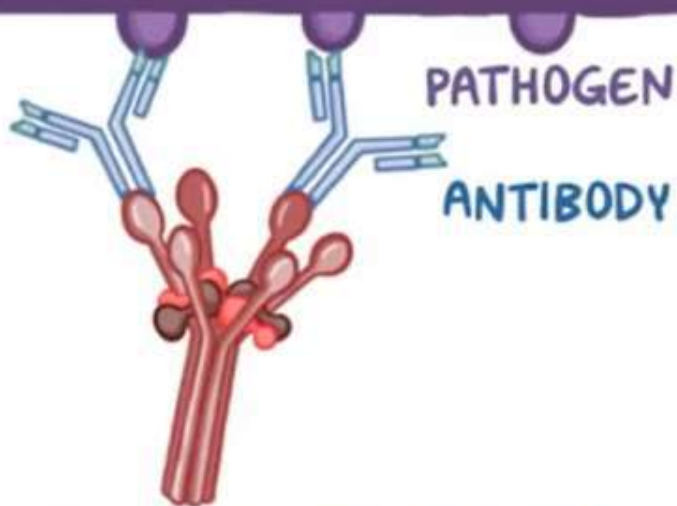


Fig. 3.11 The activation and functions of the complement system. Activation of complement by different pathways leads to cleavage of C3. The functions of the complement system are mediated by breakdown products of C3 and other complement proteins, and by the membrane attack complex (MAC).

CLASSICAL PATHWAY

→ antigen
یا وجود کی
factor is موجود کی
Antibody تـ ربطـ pathogen



LECTIN BINDING PATHWAY

→ mannose ربطـ
lectin protein is
Antibody یـ ربطـ



ALTERNATIVE PATHWAY

→ factor اول
کچھ تانی تانت رکھنا

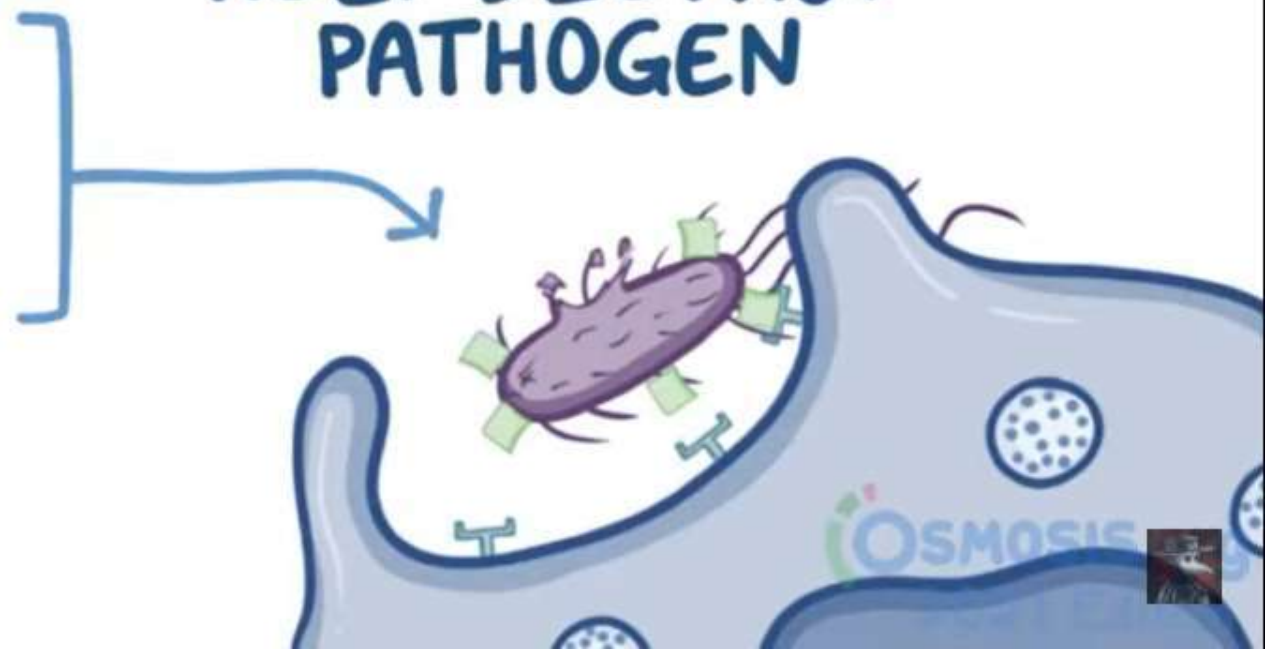


* ALONG THE WAY

- ↳ C5a + C3a → CHEMOTAXIN
- ↳ ANAPHYLATOXINS
- ↳ C3b → OPSONIN



HELP DESTROY PATHOGEN



↳ CONTRACTION OF SMOOTH MUSCLES

↳ BRONCHIAL CONSTRICTION

↳ ↑ VASCULAR PERMEABILITY

ANAPHYLATOXINS

MAST CELLS

BASOPHILS



↳ RELEASE PROINFLAMMATORY MOLECULES (HISTAMINE & HEPARIN)

C3b
OPSONIN



CHEMOTAXINS

NEUTROPHILS

EOSINOPHILS



MONOCYTES

MACROPHAGES



MAC

C9

C5

C6



C7

C8

Effects of the complement system

C3 → inactive

C3a
C3b → active

(1) **Vascular effects:** C3a & C5a (anaphylatoxins)
* increase vascular permeability & cause vasodilation.

(2) WBC activation, adhesion, & chemotaxis: C5a activates WBC, increases their adhesion to EC (integrins), & is a potent chemotactic agent for all WBC (except lymphocytes).

(3) Phagocytosis: When fixed to a microbial surface, C3b & its inactive proteolytic product C3b act as opsonins, augmenting phagocytosis by neutrophils & macrophages, which express receptors for these complement products.

(2) Kinin system activation

- In which factor XIIa convert plasma prekallikrein into kallikrein, which act on the circulating HMWK (*high molecular weight kininogen*) that leads finally to the formation of *bradykinin*.
- *Bradykinin*, like Histamine causes arteriolar dilatation, increases vascular permeability, & bronchial smooth muscle contraction.

(3) The clotting system:

The clotting system & inflammation are intimately connected processes.

The clotting system is divided into two pathways that play a role in the activation of thrombin and formation of fibrin.

I- The intrinsic clotting pathway

II- The extrinsic pathway



Clotting factors and inflammation

(1) Factor Xa increases vascular permeability & WBC emigration.

(2) Thrombin enhances WBC adhesion to EC.

(3) Fibrinogen cleavage results in the generation of fibrinopeptides that increases vascular permeability & are chemotactic for WBC.

Cell-derived mediators

1. Preformed mediators in secretory granules of cells: e.g.
Histamine is secreted by mast cells, basophils & platelets.
Serotonin is secreted by platelets.
Lysosomal enzymes are secreted by neutrophils & macrophages.

2. Newly-synthesized mediators: e.g.
Prostaglandins (PG) are secreted by all WBC, platelets, and endothelial cells.
Leukotriens (LT) are secreted by all leukocytes.
Platelets-activating factor (PAF) is secreted by leukocytes & endothelial cells.
Cytokines such as lymphokines are secreted by T-lymphocytes also by macrophages & mast cells.
Nitric oxide is secreted by macrophages.
Activated oxygen radicals (ROS) are released by all Leukocytes.

Other Cell- derived mediators

- **Reactive Oxygen Species (ROS):**

Released by neutrophils & macrophages during inflammation, have a role in microbial killing & tissue injury.

- **Nitric oxide (NO):**

A short-lived free radical gas produced by many cells, causing vasodilation & microbial killing.

- **Lysosomal enzymes:**

Granules in neutrophils & monocytes, cause microbial killing and tissue injury.

Table 3.8 Role of Mediators in Different Reactions of Inflammation

Reaction of Inflammation	Principal Mediators
Vasodilation	Histamine Prostaglandins
Increased vascular permeability	Histamine C3a and C5a (by liberating vasoactive amines from mast cells, other cells) Leukotrienes C ₄ , D ₄ , E ₄
Chemotaxis, leukocyte recruitment and activation	TNF, IL-1 Chemokines C3a, C5a Leukotriene B ₄
Fever	IL-1, TNF Prostaglandins
Pain	Prostaglandins Bradykinin
Tissue damage	Lysosomal enzymes of leukocytes Reactive oxygen species

الانحطاط
القائم

MORPHOLOGIC PATTERNS OF ACUTE INFLAMMATION:

The

- causative agents.
- severity of injury.
- the type of tissue involved,

Can all modify the basic morphologic patterns of acute inflammation, producing distinctive appearances.

لازم نعرف كل واحد
دوين بصير

جدران في -
pleura cavity
peritoneum cavity
pericardium cavity
synovial cavity

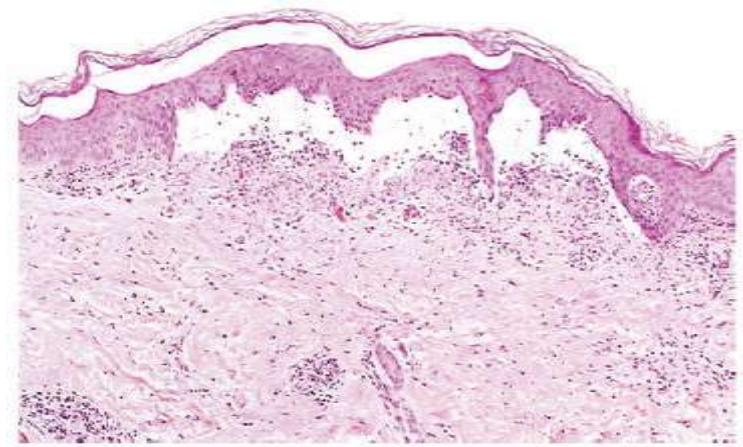
Serous inflammation

This is characterized by the outpouring of thin fluid that is either derived from the blood (serum) or the secretion of serous mesothelial cells of the pleura, peritoneum, pericardium, or the synovial cells lining the joint spaces.

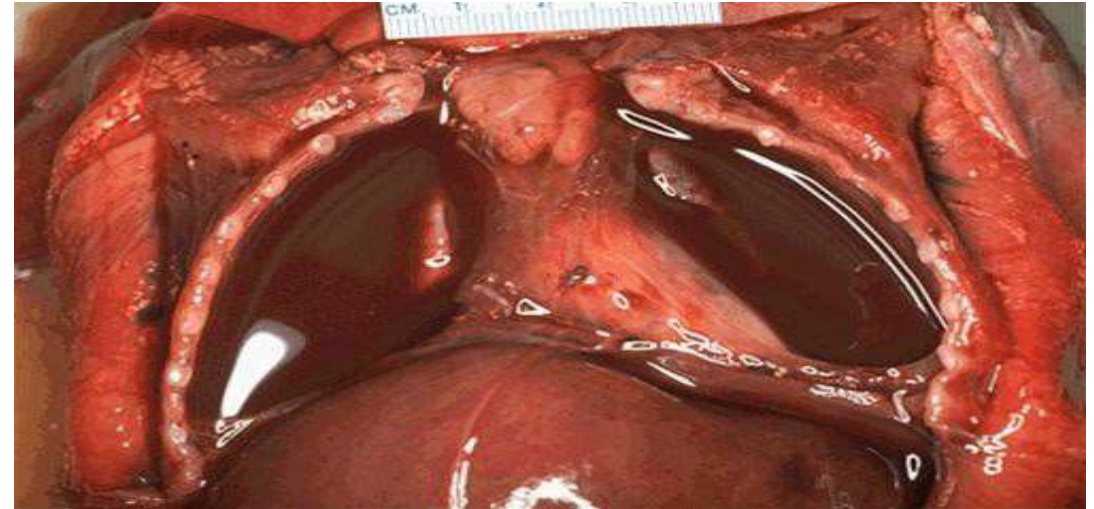
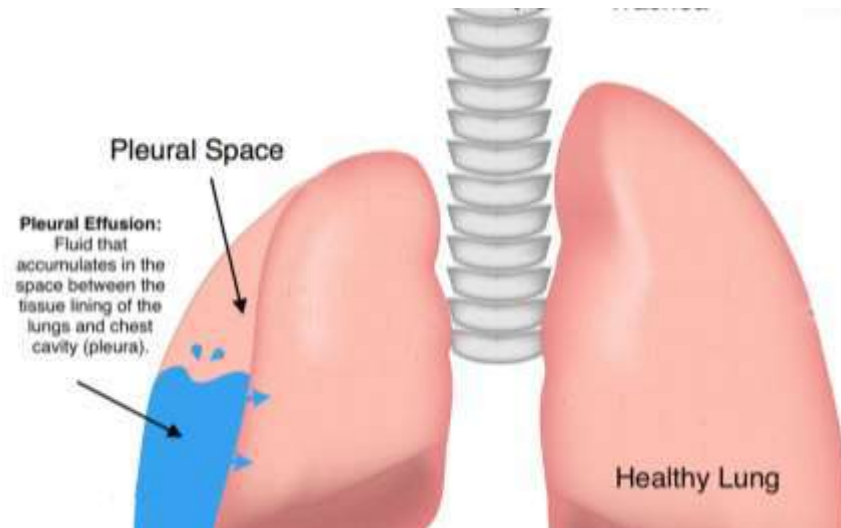
- Fluid accumulating in serous cavities is called effusions, as seen in Tuberculosis.

زني حرة المي ياي بتقله
الاشفا ياي بنحرقوا

□ Skin blister that results from a burn or viral infection is also an example of serous inflammation.



Serous inflammation



- Photographic appearance of serous inflammation, (TB) showing collection of a clear watery fluid in both pleural cavities compressing both lungs.

Fibrinous Inflammation

قسط في
pericardium

In this type of inflammation, there is the exudation of a large number of plasma proteins including fibrinogen with subsequent precipitation of masses of fibrin. This is characteristic of certain severe inflammatory responses.

Coagulation
تجلط

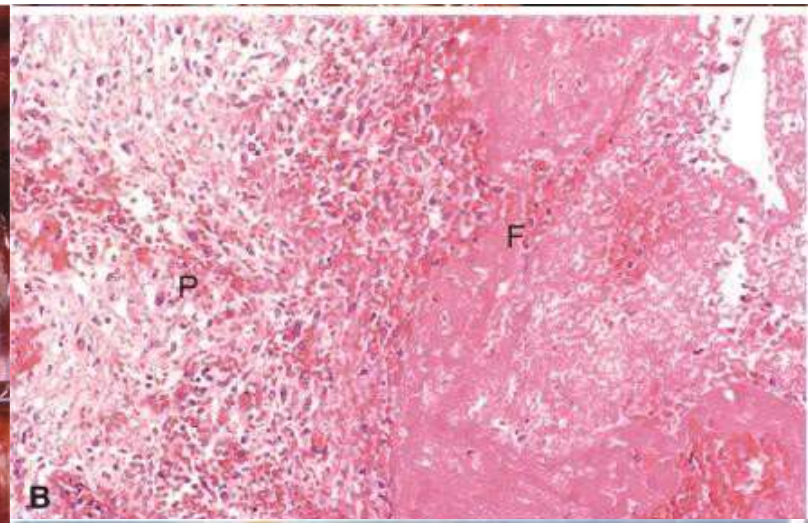
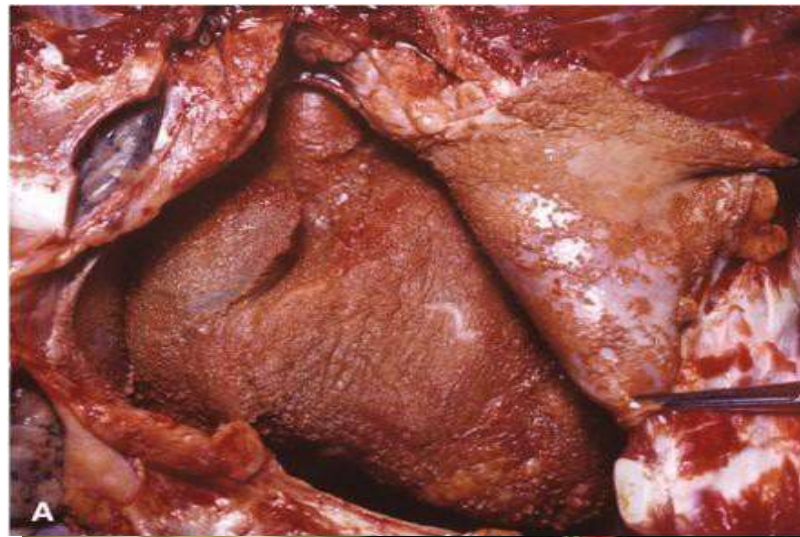
In rheumatic pericarditis, the pericardial space may become filled with large masses of fibrin, when the epicardium is stripped from the pericardium, the rubbery adherent fibrin coats both surfaces and simulating the appearance of bread and butter.

The organization of fibrinous exudates by the formation of new capillaries with fibroblasts obliterates the cavity.

Alternatively, there is fibrinolysis & resolution.

عمل الخثرة عكس
Coagulation

Fibrinous Inflammation



Top left- Gross appearance of fibrinous pericarditis, pericardium opened to show fine precipitates of fibrinous material.

Top right- microscopic view of same case showing homogenous pinkish-colored fibrinous material (F).

The bottom-Gross appearance of chronic fibrinous inflammation in rheumatic pericarditis showing thick bread & butter appearance of fibrinoid material.

كبد بال
appendix

Suppurative inflammation



This is characterized by the production of a large amount of pus (or purulent exudate).

السبب → Infection with Staphylococci ^{بكتيريا} produces localized suppuration as the skin pustule. ^{بال}

In suppurative appendicitis, ^{appendix} there is pus within the lumen and an intensive infiltration of polymorph neutrophils that are present in the mucosa, submucosa, muscularis & serosa of the appendix. ^{الزائدة الدودية}

Suppurative inflammation



- Gross appearance of the lower surface of the brain showing suppurative inflammation of meninges.

عنان صوتي
Colon / pharynx / larynx

Membranous or pseudomembranous inflammation

- This is a form of inflammatory reaction that is characterized by the formation of a membrane or more correctly a pseudomembrane because it is a non-viable structure.
انه اصلا
جاستفيد
به كليل
- It is usually made up of precipitated fibrin, necrotic epithelium & inflammatory leukocytes including polymorph neutrophils, red cells, bacteria & debris of dead tissue producing a false membrane over the inflamed surfaces looking like a bursting volcano.
waste
- This occurs when the inflammation is so severe as to cause epithelial necrosis and sloughing.

An example of this pattern is seen with Diphtheria affecting the larynx & pharynx. It may also affect the large bowel causing pseudomembranous colitis. The latter is caused by *Clostridium difficile* infection.

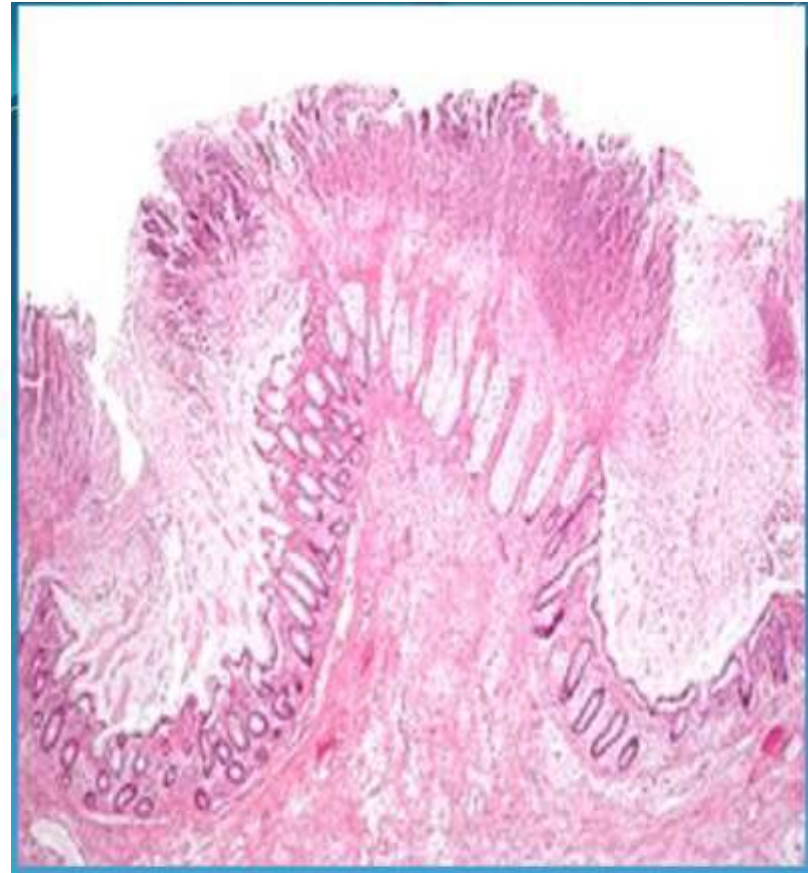
الاعمال
الفالطه



**Membranous or
pseudomembranous
inflammation**



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Left: Gross appearance of the colon showing pseudomembranous colitis showing numerous soft yellow pseudomembranes on the colonic mucosal surface.

Right: microscopic appearance of pseudomembranous colitis showing a bursting volcano-like precipitate of necrotic slough on colonic mucosa.

القرحة

ع

• An ulcer is a local defect, or excavation, of the surface of an organ or tissue that is produced by necrosis of cells and sloughing (shedding) of necrotic and inflammatory tissue.

• Ulcers are most encountered in

(1) the mucosa of the GIT and

(2) the subcutaneous tissues of the lower extremities in older persons who have circulatory disturbances

• Ulcerations are best exemplified by peptic ulcers of the stomach or duodenum, in which acute and chronic inflammation coexist.

• During the acute stage, there is intense polymorphonuclear infiltration and vascular dilation in the margins of the defect. As it develops into a chronic stage, the margins and base of the ulcer develop scarring with an accumulation of lymphocytes, macrophages, and plasma cells.

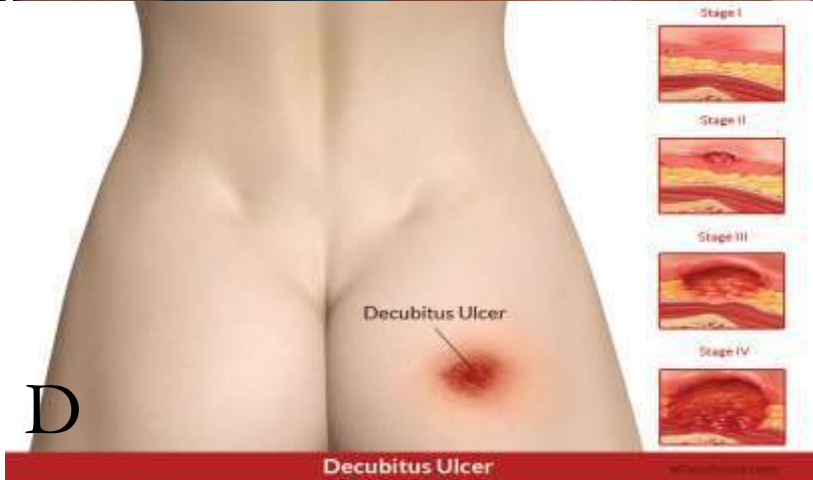
يمكن يكون
acute
و يمكن ليس
Chronic
بازالة تها
علاجها

مكان حدوثه
GI
lower extremities
mouth
back

Ulcerative inflammation

لعمل قرحة

**Ulcerative
inflammation**



A. Mouth Ulcer

B. Histological appearance of an ulcer

C. Diabetic foot ulcer

D. Decubitus Ulcer/Pressure Ulcer/Bedsore: occurs at sites with little fat and muscle over bony prominences. Ulceration occurs due to the breakdown and ulceration of tissue due to a combination of the weight of the body on the surface of the skin and the friction of a resistant surface such as a bed.

Outcomes of acute inflammation

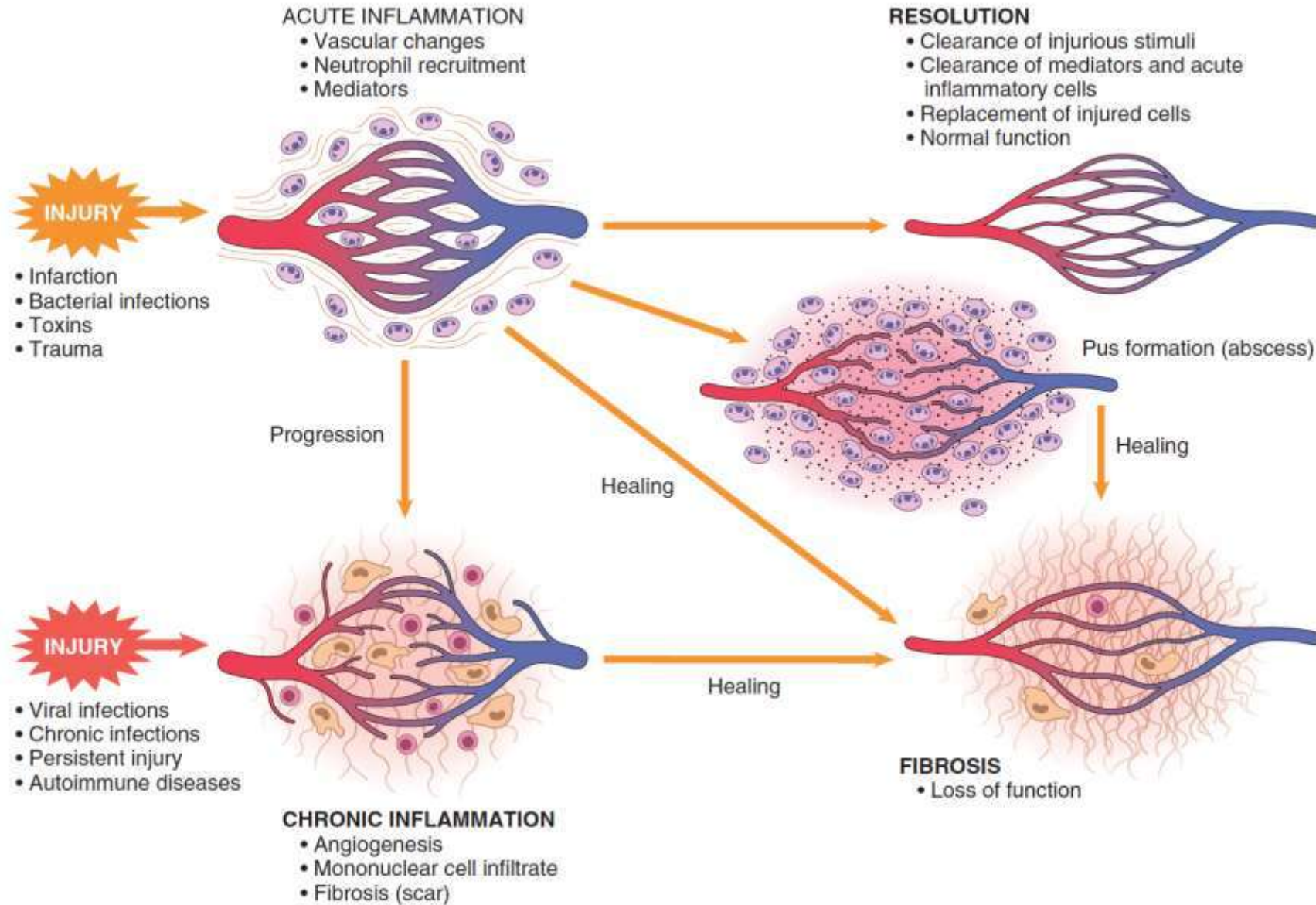


Fig. 3.16 Outcomes of acute inflammation: resolution, healing by fibrosis, or chronic inflammation. The components of the various reactions and their functional outcomes are listed.

Chronic Inflammation

Types of Chronic Inflammatory Cells
Effects of Inflammation

Chronic Inflammation

Inflammation of prolonged duration (weeks, months to years) in which active inflammation, tissue injury, & healing proceed simultaneously.

طی وقت واحد

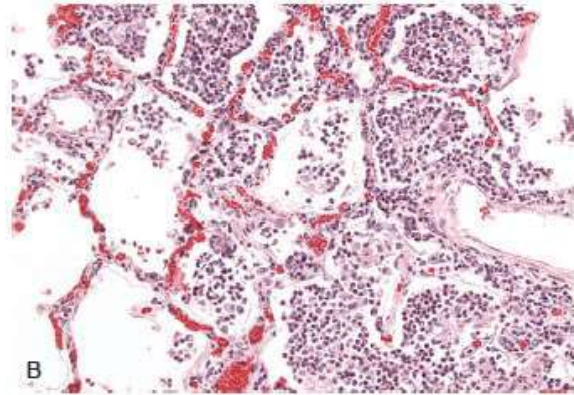
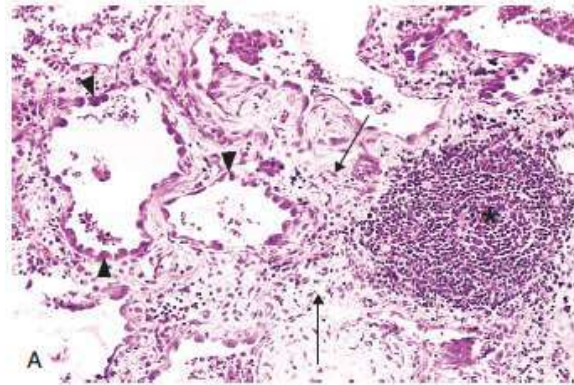
In contrast to acute inflammation, which is distinguished by vascular changes, edema, & neutrophilic infiltrate, chronic inflammation is characterized by chronic *inflammatory cells infiltration* together with *connective tissue deposition & scarring*.

Characteristics of Chronic Inflammation

1. Mononuclear chronic inflammatory cells infiltration, including macrophages, lymphocytes and plasma cells.
2. Tissue destruction, largely directed by the inflammatory cells.
3. Repair, involving new vessel proliferation (angiogenesis) & fibrosis.

لفني كحتموم بتحصير
كبيره

Chronic Vs Acute inflammation in the lungs



Chronic inflammation in the lung, showing the characteristic histologic features:

- * collection of chronic inflammatory cells;
- destruction of parenchyma (normal alveoli replaced by spaces lined by cuboidal epithelium); and
- replacement by connective tissue, resulting in fibrosis.

Acute inflammation of the lung (acute bronchopneumonia), neutrophils fill the alveolar spaces and blood vessels are congested.

Nature of leukocyte infiltrates in inflammatory reactions

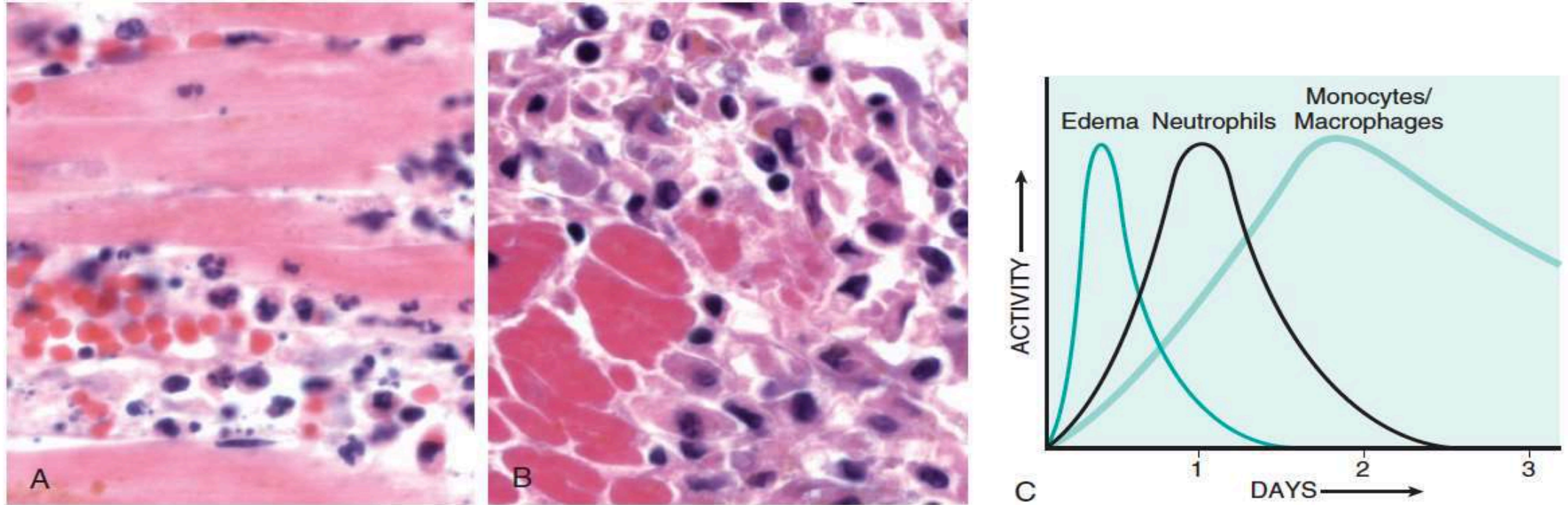


Fig. 3.5 Nature of leukocyte infiltrates in inflammatory reactions. The photomicrographs show an inflammatory reaction in the myocardium after ischemic necrosis (infarction). (A) Early (neutrophilic) infiltrates and congested blood vessels. (B) Later (mononuclear) cellular infiltrates. (C) The approximate kinetics of edema and cellular infiltration. For simplicity, edema is shown as an acute transient response, although secondary waves of delayed edema and neutrophil infiltration also can occur.

Causes of chronic inflammation

3. يمكن واصل لمرحلة
انه العدوى
ما يردج (ما يحتاج
بالمناد السوي)

1. Progression of acute to chronic inflammation
2. Viral infections = *Most viral infections elicit chronic inflammation*
3. Persistent infections = *microbes that are difficult to eradicate e.g., tubercle bacilli of T.B*
4. Immune-mediated inflammatory diseases, or hypersensitivity diseases = *Rheumatoid arthritis (RA) & inflammatory bowel syndrome (IBS) + bronchial asthma*
5. Prolonged exposure to potentially toxic agents =
 - Exogenous eg. inhaled silica & asbestos
 - Endogenous eg. plasma lipid components, which may contribute to atherosclerosis

Chronic Inflammatory cells and Mediators

- Macrophages

- Lymphocytes
- Plasma cells
- Eosinophils
- Mast cells
- Neutrophils

(1) Macrophages

The *most important* cell of chronic inflammation, are tissue cells that are derived from circulating blood monocytes.

Macrophages, scattered diffusely in most connective tissues normally, but found in increased numbers in certain organs (mononuclear phagocyte system):

- liver (Kupffer cells),
- CNS (microglial cells),
- Lungs (alveolar macrophages),
- Spleen & lymph nodes (sinus histiocytes).

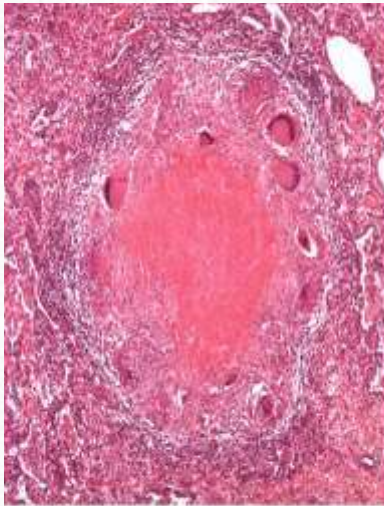
بعضها
تؤخذ واحدة

Activated macrophages

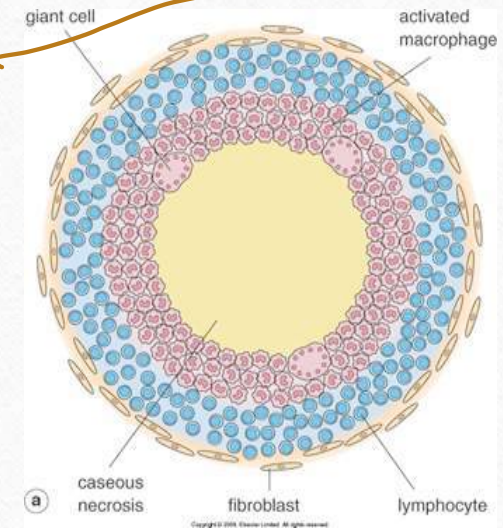
لا تكون
اصناما نستعين بها لكن
لما تصير
تعمل بالهمة
monocytes
macrophages

The activated macrophages appear large, flat pink, this appearance is similar to that of squamous cells & therefore, these cells are called *epithelioid macrophages* (epithelial-like). Focal aggregates of these cells are called a *granuloma*.

يكون
دائرا



اذا ضلت macrophage
نشطة، ضلت تتجمع بشكل
متر نتيجة وجود
inf]



Macrophage Products

1. Acid & neutral proteases, plasminogen activator.
2. ROS & NO.
3. (Arachidonic Acid) AA metabolites.
4. Cytokines, e.g., IL-1 & TNF.
5. GFs (PDGF, FGF, TGF- β) that influence the proliferation of fibroblasts, SMC, & the production of ECM.

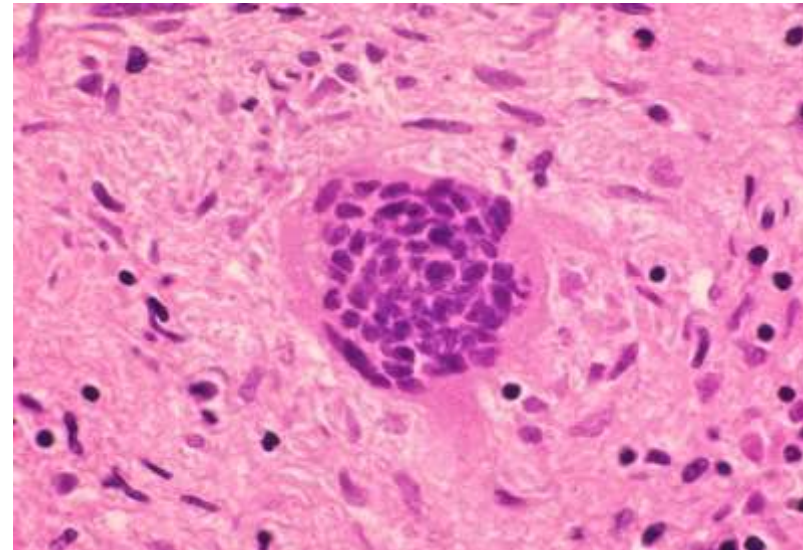
Fate of macrophages

- At sites of acute inflammation where the irritant is cleared & the process is resolved, macrophages die, or pass into lymphatics.
- However, in chronic inflammatory sites, macrophage accumulation persists, & proliferates. يحفز على
- Steady release of lymphocyte-derived chemokines & other cytokines (e.g., IFN- γ) من macrophages is an important mechanism by which macrophages are recruited to, or immobilized, in inflammatory sites.

نخبة كبيرة تحتوي على النسيج
Giant cell: من dead macrophages

IFN- γ can also induce macrophages to fuse into large, multinucleated cells called giant cells, a cell contains two or more (up to hundreds) nuclei.

كل
macrophages
في جواتها النسيج
من الخلايا



(2) Lymphocytes

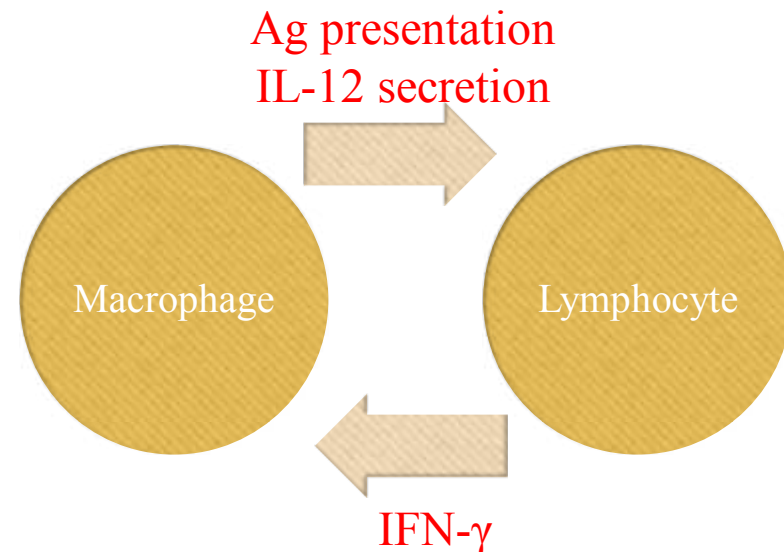
Both T & B lymphocytes migrate into inflammatory sites using some of the same adhesion molecule pairs & chemokines that recruit monocytes.

Lymphocytes are mobilized in the setting of: بتكون موجودة في ما بين

1. any specific immune stimulus (infections). محرزات خاصة محددة

2. in non-immune-mediated inflammation (e.g. due to infarction or tissue trauma). التهبات عن غير المناعي

T lymphocyte ← اذا صار عنا cell damage
B lymphocyte ← اذا كان في micropes



B cell
بالأخرى نطقنا
plasma cells

(3) Plasma cells

Are the terminally differentiated end-product of B-cell activation; they can
produce antibodies directed either against

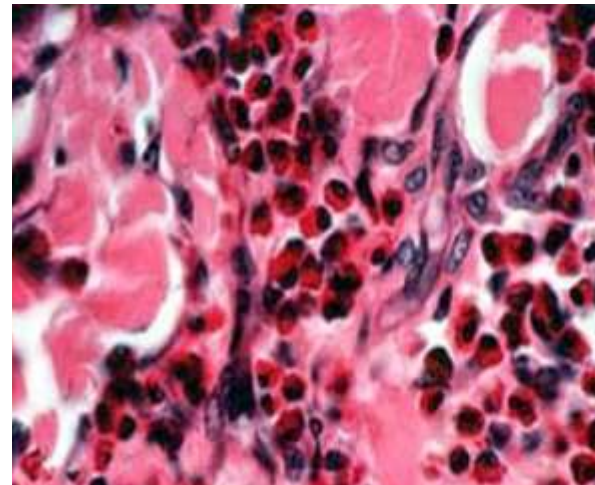
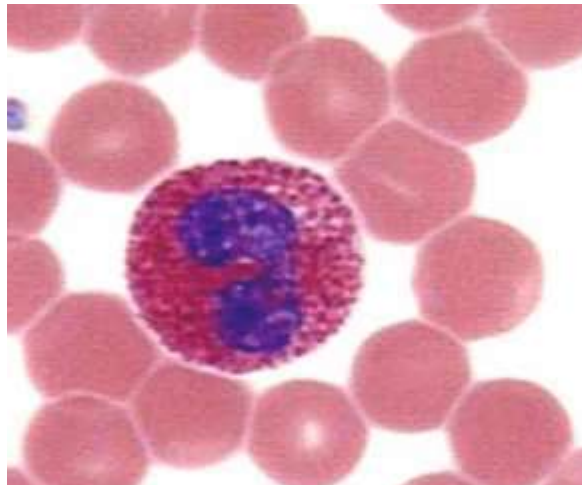
- (1) Persistent Ags in the inflammatory site, or
- (2) Altered tissue components.

(4) Eosinophils

Characteristically found in inflammatory sites around:

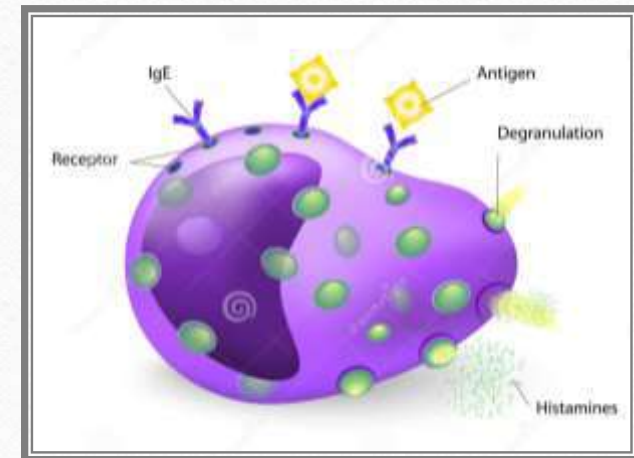
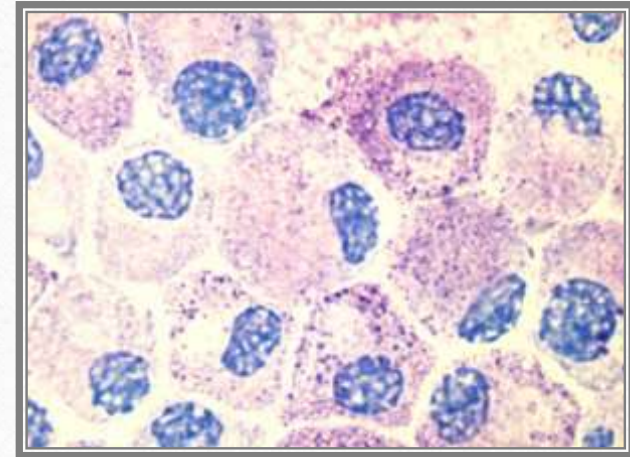
^{العدوى الطفيلية} ① Parasitic infection, eosinophil-specific granules contain a major basic protein, a highly charged cationic protein that is toxic to parasites.

^{ردود الفعل التحسسية المناعية} ② Part of Allergic immune reactions. ^{تكون عند تحسس متاسي معين} The emigration of eosinophils is driven by adhesion molecules similar to those used by neutrophils, & by specific chemokines derived from WBC or epithelial cells.



(5) Mast cells

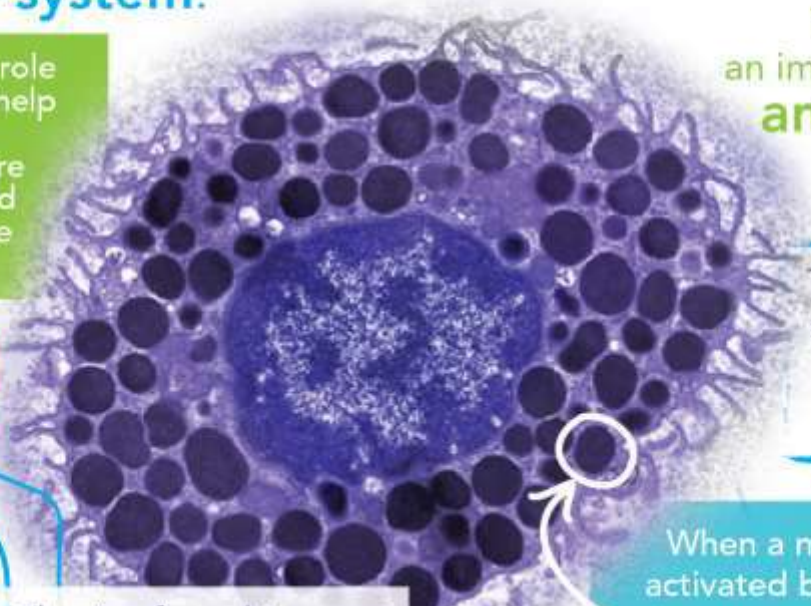
- Are sentinel (guard) cells, widely distributed in connective tissues throughout the body & can participate in both acute & chronic responses.
- In atopic individuals, mast cells are “armed” with IgE Antibodies (Ab) specific for certain Antigens (Ag).
- When these Ags are subsequently encountered, the prearmed mast cells are triggered to release histamine that elicits the vascular dilation & increased permeability of acute inflammation.
- IgE-armed mast cells are central players in allergic reactions including anaphylactic shock & can elaborate cytokines such as TNF & chemokines.



What is a MAST CELL?

Mast cells are a part of the immune system.

Mast cells play a role in inflammation, help defend against pathogens and are involved in wound healing and tissue repair.



Mast cells are well-known for releasing histamine during allergic reactions, such as in pollen or insect sting allergies.



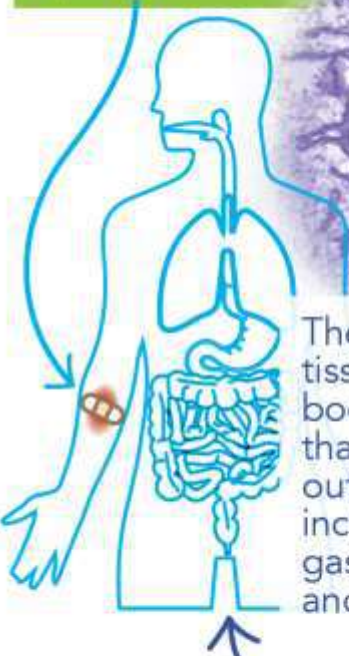
They play an important role in **anaphylaxis!**

They can detect and respond to foreign substances.

When a mast cell is activated by a trigger, these granules release many mediators (chemicals that mediate reactions leading to symptoms).

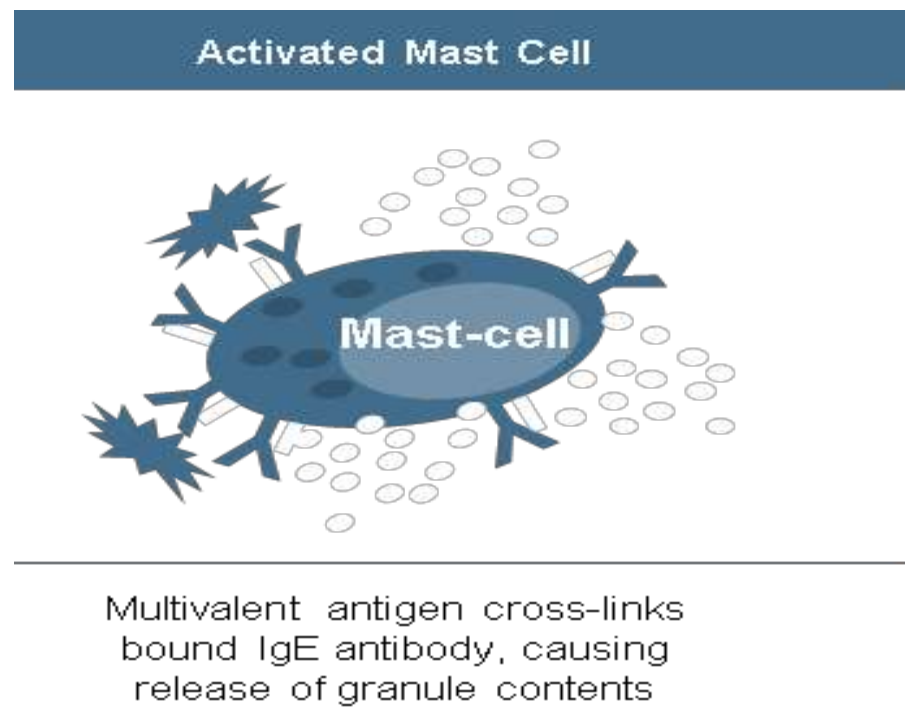
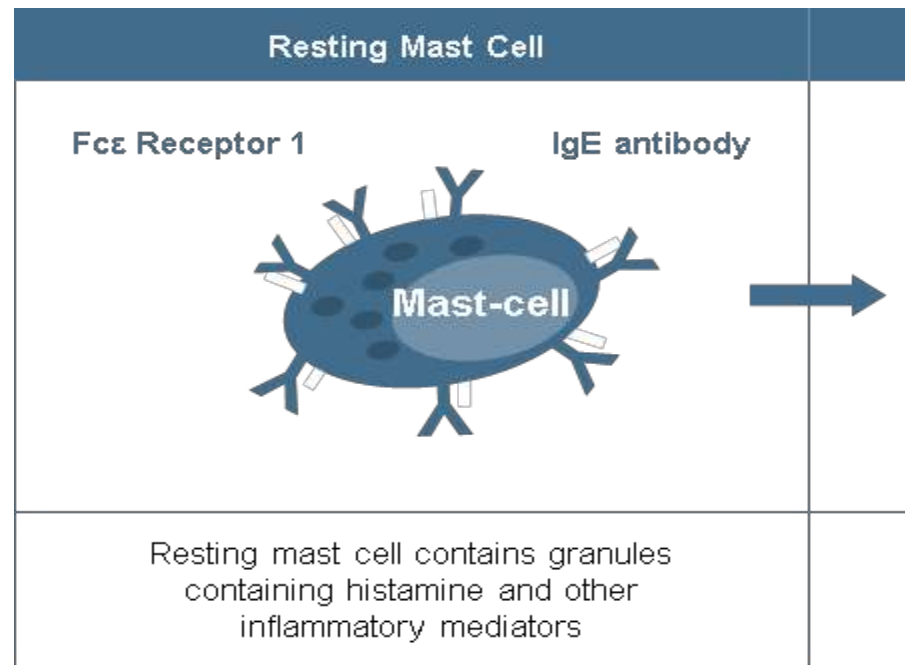
histamine
is a mediator

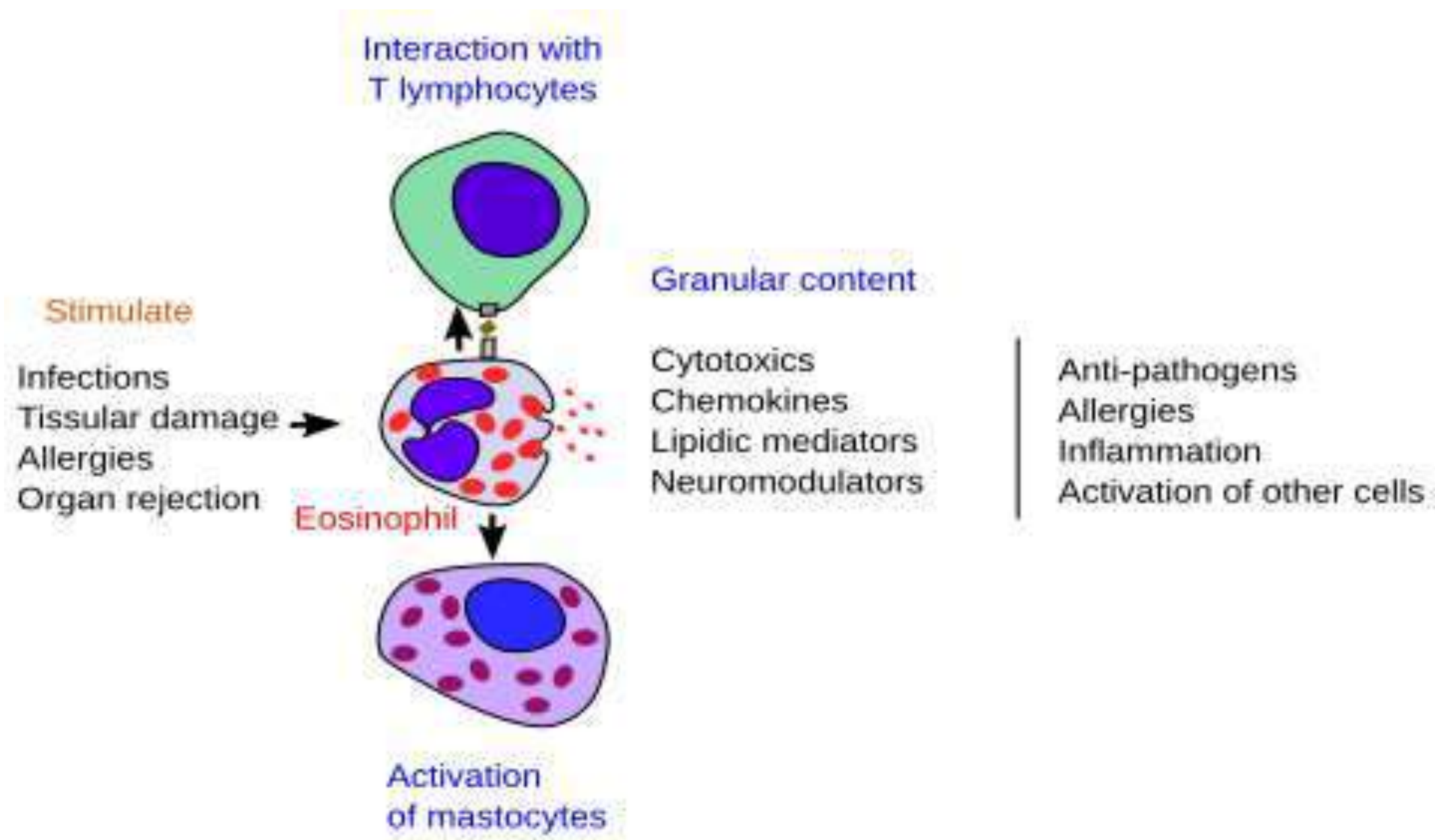
They're found in most tissues throughout the body, especially those that interact with the outside environment, including the lungs, gastrointestinal tract and skin.



MAST CELL DISEASE happens when these cells aren't behaving normally.

LEARN MORE AT tmsforacure.org





Interaction with
T lymphocytes

Stimulate

- Infections
- Tissue damage
- Allergies
- Organ rejection

Eosinophil

Granular content

- Cytotoxics
- Chemokines
- Lipidic mediators
- Neuromodulators

- Anti-pathogens
- Allergies
- Inflammation
- Activation of other cells

Activation
of mastocytes

تتضمن نتيجة إفراز
chemical
mediators
من macrophages

(6) Neutrophils

- Although neutrophils are the classic hallmarks of acute inflammation, many forms of chronic inflammation may continue to show extensive neutrophilic infiltrates, due either to persistent of
 - (1) microbes, or
 - (2) necrotic cells, or
 - (3) to mediators elaborated by macrophages.

Effects of inflammation

A- Beneficial effects:

- These act partly through the flow of exudates into the tissue & partly by the phagocytic & microbial effects of migrated WBC:
 - Dilution of toxins.
 - Protective antibodies.
 - Fibrin formation.
 - Promotion of immunity.

B-Harmful effects:

- Swelling: of acutely inflamed tissue may have serious mechanical effects e.g in acute laryngitis suffocation in children.
- Rise in tissue pressure

يصير عملية التنفس صعبة

Beneficial effects of inflammation

(1) Dilution of toxins:

Exudates dilute chemical and bacterial toxins & enhance their carriage by lymphatics.

الانزلاق كخفف
تنزير
الادوية اللقاحية

(2) Protective antibodies:

The proteins present in the exudates include antibodies, which have been already present in the plasma as a result of previous infection or immunization. These antibodies attack injurious agents in an attempt to destroy them immunologically.

المعامل الضارة
محاولة

Beneficial effects of inflammation

(3) Fibrin formation:

Fibrinogen of the blood is included in the exudates. In the extravascular space, it is converted into solid fibrin. A network of the deposited fibrin is seen in the inflamed tissue forming a mechanical barrier that precludes the movement & spread of bacteria, it may also aid in their phagocytosis.

شاهد

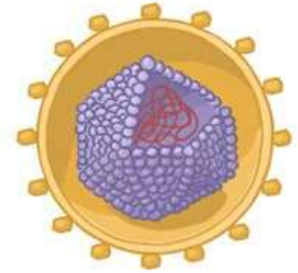
تعزيز المناعة

(4) Promotion of immunity:

Bacteria in the inflammatory exudates, whether free or phagocytosed, are carried to the lymph nodes by lymphatics. There they mount an immune response, which provides antibodies & cellular mechanisms that may appear after a few days and may remain for years. These immunological mechanisms help destroy microbial agents.

INFLAMMATION

COMPLEX RESPONSE to HARMFUL STIMULI



PATHOGEN

TRAUMA or TOXINS



BLOOD VESSELS DILATE

PERMEABLE

ATTRACT IMMUNE CELLS & FLUID

HEAT



REDNESS

PAIN

SWELLING

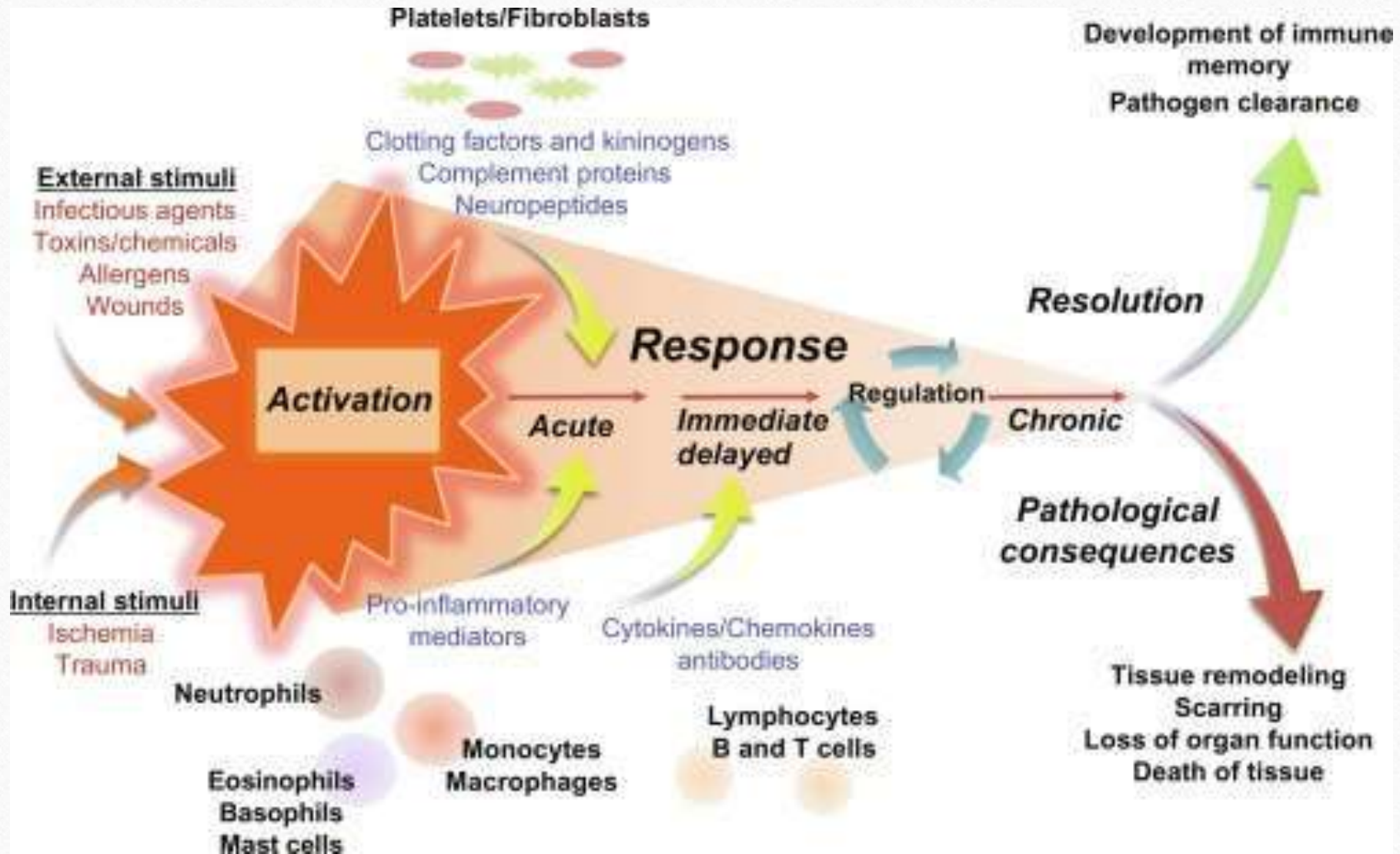
LOSS OF FUNCTION

WOUND REPAIR & RESOLUTION

INITIAL TISSUE INTEGRITY

FIBROUS SCAR





Thank You

اللهم إني استودعتك ما قرأت وما
حفظت وما تعلمت، فردّه عند حاجتي
إليه إنك على كل شيء قدير، بسم الله
الفتاح، اللهم لا سهل إلا ما جعلته
سهلاً، وأنت تجعل الحزن إذا شئت
سهلاً يارب العالمين

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