# Non-Specific Immunity

# Innate (natural, native, non-specific) immunity

- Always present, ready to recognise and eliminate microbes. May be stimulat also by non-microbial agents.
- Frequently eliminates microbes before the specific immunity becomes active.
- Receptors are encoded in the germline, are not a product of recombination of genes.

# Differences between the Innate and Acquired Immunity

- Innate Immunity
  - Universal
  - Rapid
  - Lacks memory

- Acquired Immunity
  - Not universal
  - 'Slow' to develop
  - Memory
  - Specific but in some situations reacts to autoantigens
  - 'Plays to the tune of the innate immune system'

# Signals of danger

EXOGENEOUS (PAMPs)

**ENDOGENEOU** (e.g. STRESS PROTEINS RELEASD FROM NECROTIC CELL)

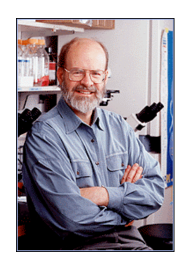
# Inborn immunity – activating signals (alarmins)

#### Pathogen-Associated Molecular Patterns (PAMPs)

(C. A. Jeneway, Jr, 1989)

microbial structures, motifs, present in large groups of microorganisms, necessary for their life

e.g. lipopolysachrides, lipopeptides, peptidoglycans, manose nonmetylated CpG present in bacterial DNA, dsRNA of RNA-vises



#### **Danger, Damage-Associated Molecular atterns (DAMPs)**

(Polly Celine Eveline Matzinger, 1994)

Molecular structures of the host macroorganism

e.g. HSP60, HSP70, fragments of fibrinogen, fibronectin, hyaluronan..



#### Differences between innate and specific immunity

	Innate immunity	Adaptive immunity
Specificity	For structures shared by classes of microbes ("molecular patterns")	For structural detail of microbial molecules (antigens); may recognize nonmicrobial antigens
	Different microbes Identical mannose receptors	Different microbes - Distinct antibody molecules
Receptors	Encoded in germline; limited diversity	Encoded by genes produced by somatic recombination of gene segments; greater diversity
	Toll-like receptor Mannose receptor	TCR YET YOU
Distribution of receptors	Nonclonal: identical receptors on all cells of the same lineage	Clonal: clones of lymphocytes with distinct specificities express different receptors
Discrimination of self and nonself	Yes; host cells are not recognized or they may express molecules that prevent innate immune reactions	Yes; based on selection against self-reactive lymphocytes; may be imperfect (giving rise to autoimmunity) bownloaded from: StudentCons

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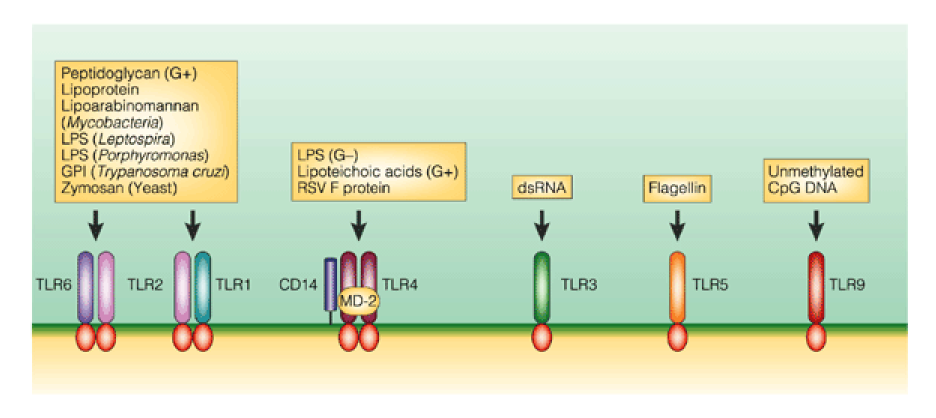
PAMPs – pathogen-associated molecular patterns exogenou substances activating non-specific immunity (Endotoxin, mannose, double-stranded RNA, unmelylated CpG nucleotides)

DAMPs – danger associated molecular patterns – endogenous substances stimulating non specific immunity – heat shock proteins, uric acid

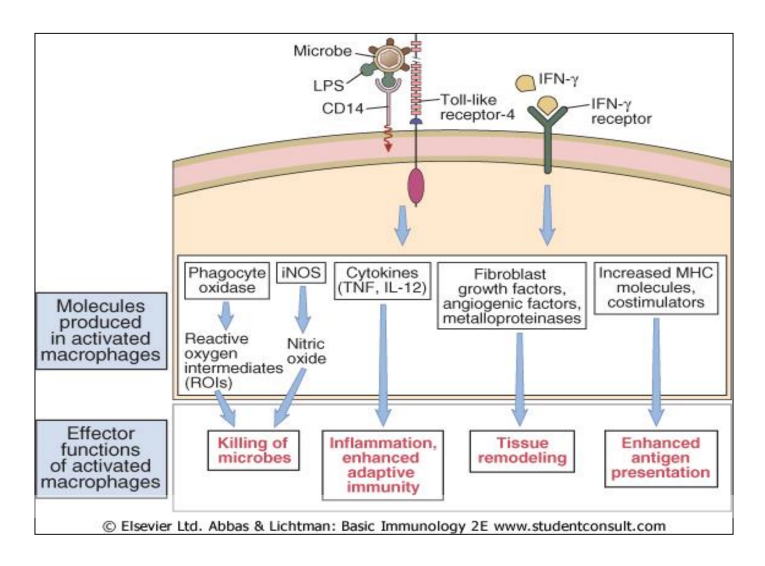
PRR- Pattern recognition receptors - recognize PAMPs, DAMPs.

TOLL-like receptors –surface or intracellular receptors recognizing various PAMPs. Expressed on dendritic cells, macrophages, granulocytes, epitelial cells.... They induce activation of these cells.

### **TOLL-LIKE RECEPTORS**



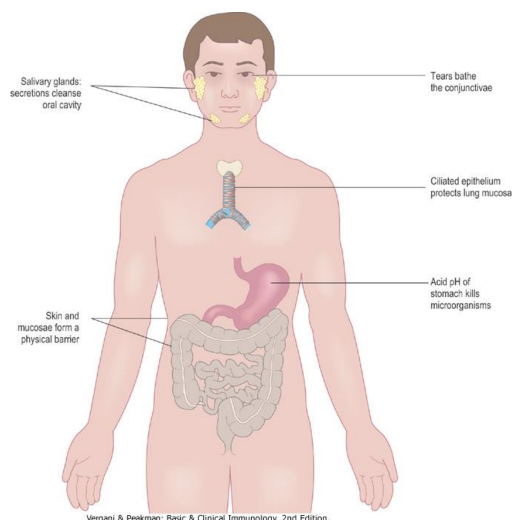
#### Activation by Toll-like Receptors and by Cytokine Receptors



# Basic components of non-specific defence

- Non Specific barriers
  - Anatomical/Physiological
- Acute phase reactants and Inflammation
  - Complement/Interferons/CRP
- Innate cells
  - PMN/Macrophages/NK cells

## Non-specific barriers of human body



Vergani & Peakman: Basic & Clinical Immunology, 2nd Edition.
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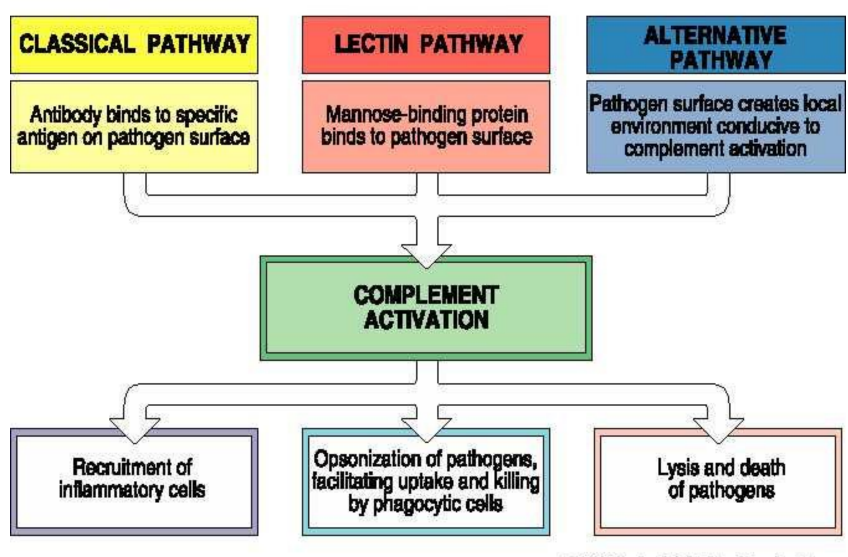
# The Complement System

# General features of the Complement System Activation

- Inactive, preformed protein is activated by the proteolytic cleavage.
- It is cleft into the smaller part (called a) and a bigger part (called b).
- Usually the bigger part has also proteolytic activity, while the smaller part has various other biological activities (chemotactic, anaphylatoxic).
- Component C6-C9 are activated without cleavage, they just "attach" to the complex of the other complement components.

#### **Activation of the complement sytsem**

Figure 7.27



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Complement system

## Alternative pathway spontaneous activation

C3  $\longrightarrow$  C3b Bf Factor D Bb

C3b/Bb

#### **Lectin pathway**

mannose binding

MBL

MASP-1

MASP-2

C4/C2 ----> C4b/C2b

C4b/C2b

## Classical pathway antibody binding

 $C1q/r_2/s_2$ 

C4/C2 ---- C4b/C2b

C4b/C2b

#### C3 convertase

C3 ——— C3a + C3b

### C5 convertase $C5 \longrightarrow C5b$

# lytic pathway formation of membrane-attack comlex, lysis of pathogens

C5b, C6, C7, C8, C9

#### opsonization phagocytosis

C3b

#### anaphylatoxin

inflammation, phagocyte recruitment

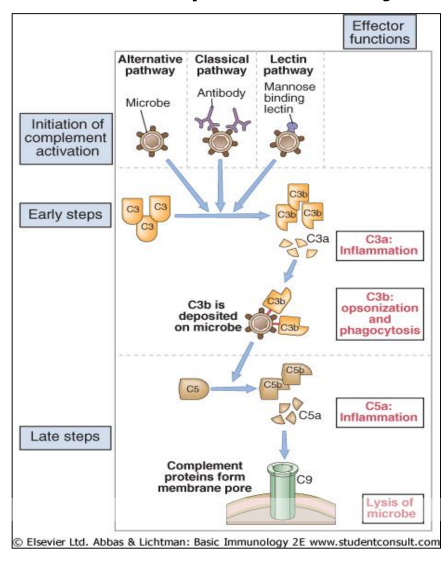
C4a, C3a, C5a

## Complement system activation

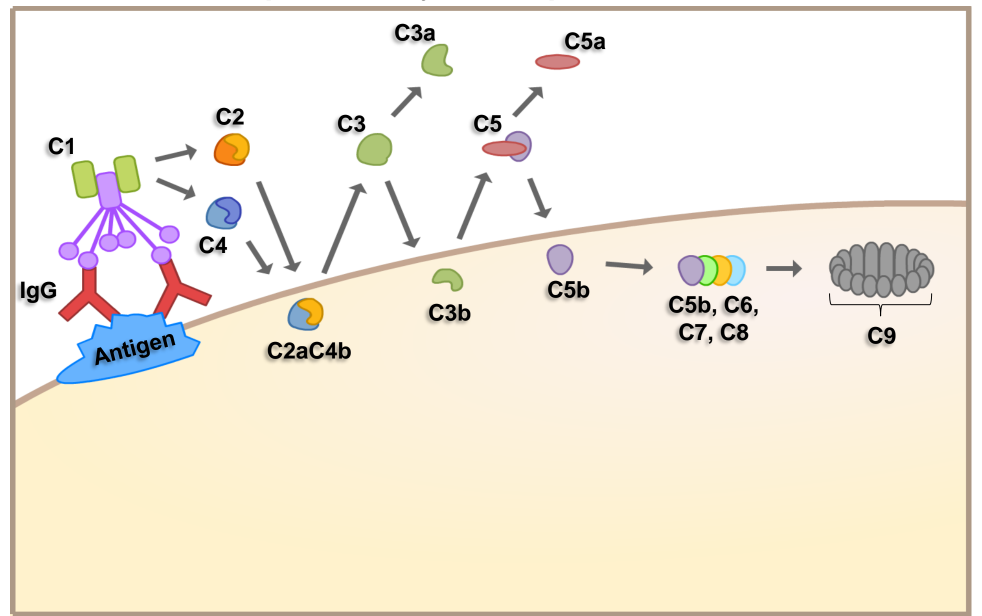
- Classical pathway:
  - Complexes IgG-antigen, IgM-antigen,
  - C-reactive protein
- Alternative pathwas
  - Lipopolysaccharide of G- bacteria
  - Cell wall of some bacteria
  - Cell wall of the yeasts (zymozan)
  - Aggregated IgA
- Lectin pathway:
  - Mannose and other sacharides



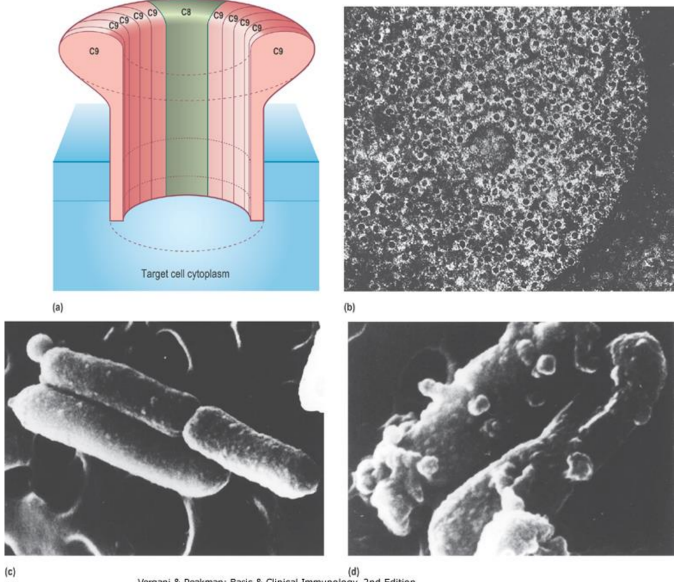
#### The Complement System



## Classical pathway complement activation

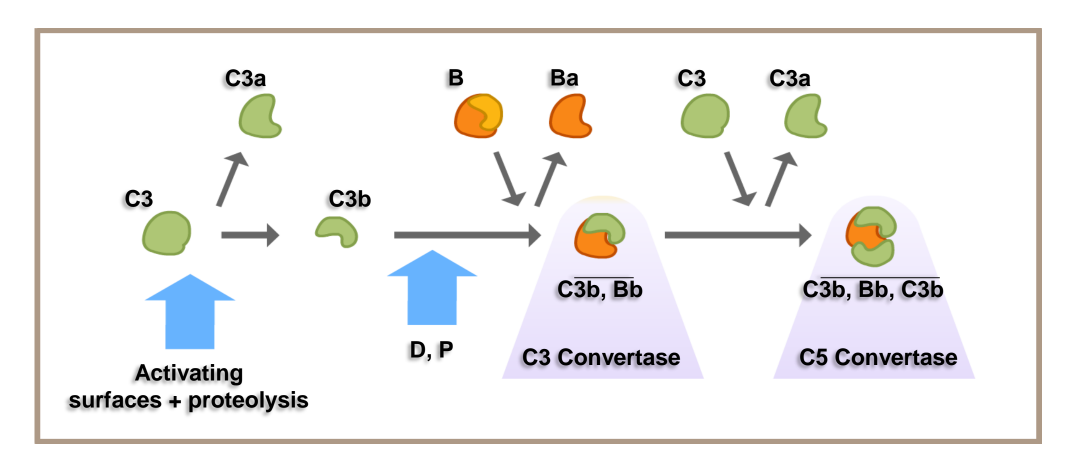


#### Effect of C9



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# Actiation of Alternative Pathway of the Complement system



# Regulation of the Complement System

- Factors present in plasma
  - Classical pathway mainly C1-inhibitor C1-INH (deficiency leads to hereditary angioedema).
  - Alternative pathway factor H, factor I
- Faktors on cell membranes:
  - CD59, CD55 (DAF decay accelerating factor) CD46

# Patothogenetic Significance of the Complement System

- Deficit of components of the classical and alternative pathways – proneness to bacterial infections and to systemic autoimmune diseases.
- C1-INH deficiency: hereditary angioedema.
- Mutations of factor H atypical hemolytic-uremic syndrome.
- Several polymorphisms of factor H are linked to senile macullar degeneration.
- Paroxysmal night hemoglobinuria caused by mutations of the gene PIG-A (product if this gene binds CD 55 and CD 59 into cytoplasmatic membrane.

# Biological effects of activated complement system

- C9 cytolytic effect
- C3b opsonisation
- C3a, C5a anaphylatoxins, liberation of histamine
- C5a chemotaxin

# Phagocytosis

## Phagocytic cells

- Polymorphonuclear granulocytes
- Monocytes + macrophages
- Dendritic cells mainly non-activated cells. After activation they loose most of their phagocytic activity.



### Polymorphonuclear granulocyte



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## Normal blood count (in adults)

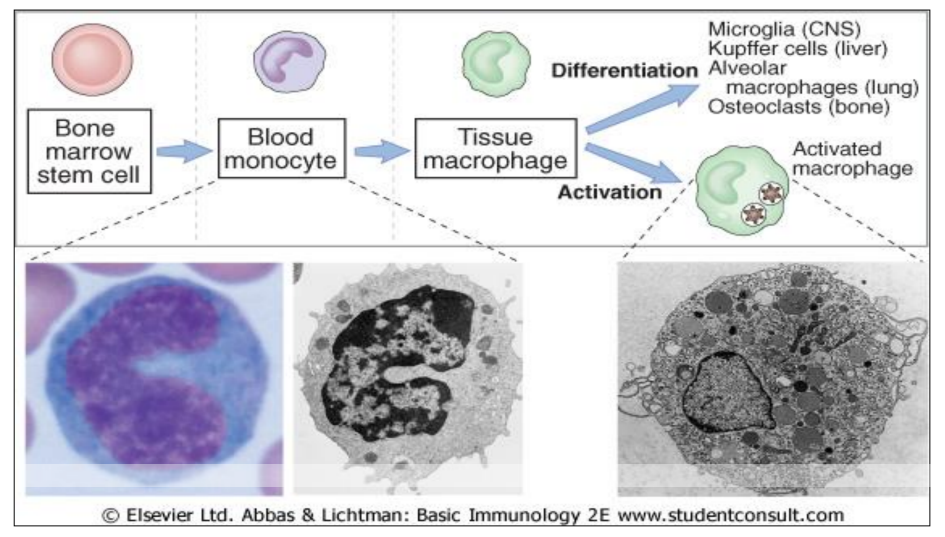
- Erythrocytes: 4-5 x 10<sup>12</sup>/I
- Thrombocytes: 150-300 x 10<sup>9</sup>/l
- Leukocytes: 4-9 x 10<sup>9</sup>/l
  - Granulocytes: 55-70%
  - Eosinophils: 1-4%
  - Basophils: 0-1%
  - Lymphocytes: 24-40%
  - Monocytes: 3-8%

## Macrophages

- Derived from blood monocytes.
- Connective tissue macrophages
  - Kupffer cells (liver)
  - Alveolar macrophages (lungs)
  - Microglia (CNS)
  - Osteoclasts (bone)
  - Peritoneal macrophages

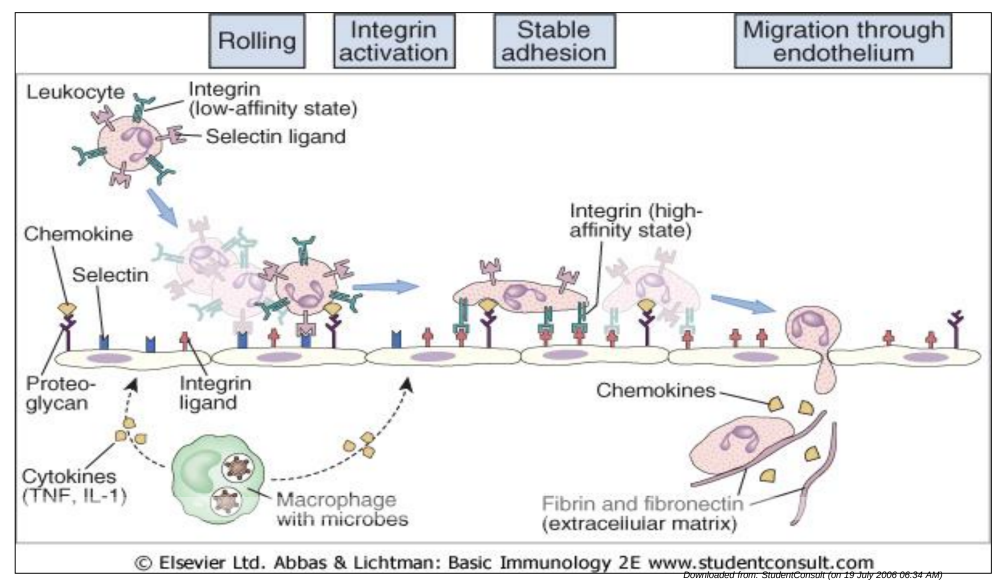


### Development of macrophages





### Extravasation of leukocytes



### Chemotaxins

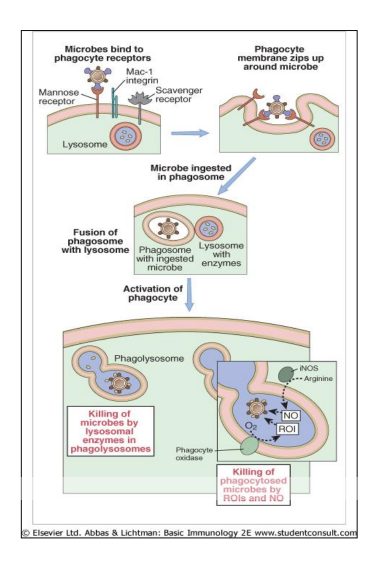
- Attract phagocytic cells
- Products of destroyed cells
- C5a
- IL-7, IL-1
- Leukotriens

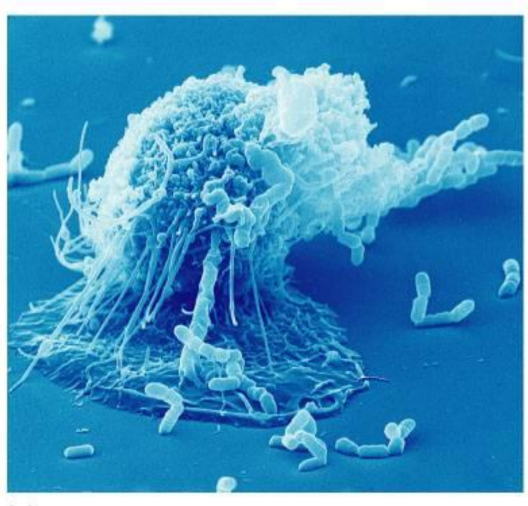
## **Opsonins**

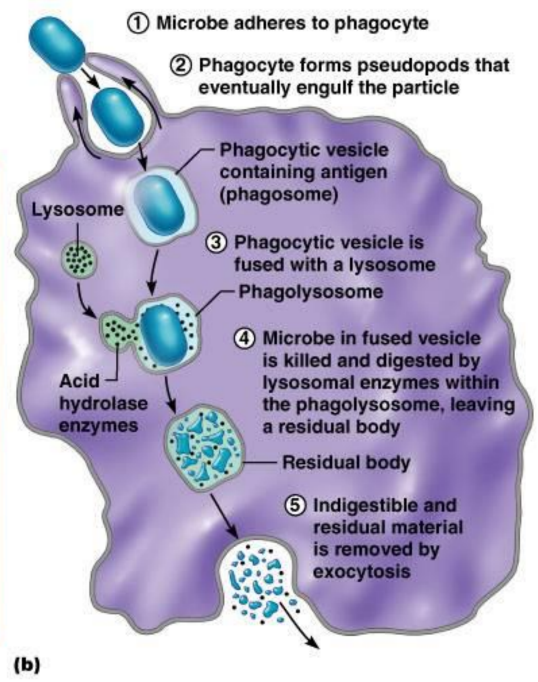
- Substances enhancing phagocytic process by improving attachment of the particle to the phagocytic cell.
- Specific: IgG, (IgM only indirectly by activation of the complement system)
- Non- specific: C3b, fibronectin....



### Steps of phagocytosis







(a)

# Killing mechanisms of phagocytic cells

- Reactive metabolites of oxygen (H<sub>2</sub>O<sub>2</sub>, hydroxyl radical (.OH), superoxide aniont (O<sub>2</sub>-), singletted oxygen (.O<sub>2</sub>)
- Reactive nitrogem intermediates (NO, NO<sub>2</sub>)
- Hydrolases: protease, lipases, DNAses
- Low pH
- Lysozyme
- Lactoferin
- Defensins antimicrobial polypeptides

Class of mechanism	Specific products	
Acidification	pH=~3.5 -4.0, bacteriostatic or bacteriocidal	
Toxic oxygen-derived products	Superoxide O <sub>2</sub> <sup>-</sup> , hydrogen peroxide H <sub>2</sub> O <sub>2</sub> , singlet oxygen <sup>1</sup> O <sub>2</sub> , hydroxyl radical OH; hypohalite OCI <sup>-</sup>	
Toxic nitrogen oxides	Nitric oxide NO	
Antimicrobial peptides	Defensins, cationic proteins	
Enzymes	Lysozyme — dissolves cell walls of some Gram-positive bacteria.  Acid hydrolases — further digest bacteria	
Competitors	Lactoferrin — binds Fe, vitamin B <sub>12</sub> binding protein	

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## Lysozyme

- Cleaves cell walls of G+ bacteria
- Present in granules of neutrophil granulocytes, in plasma, secretions.

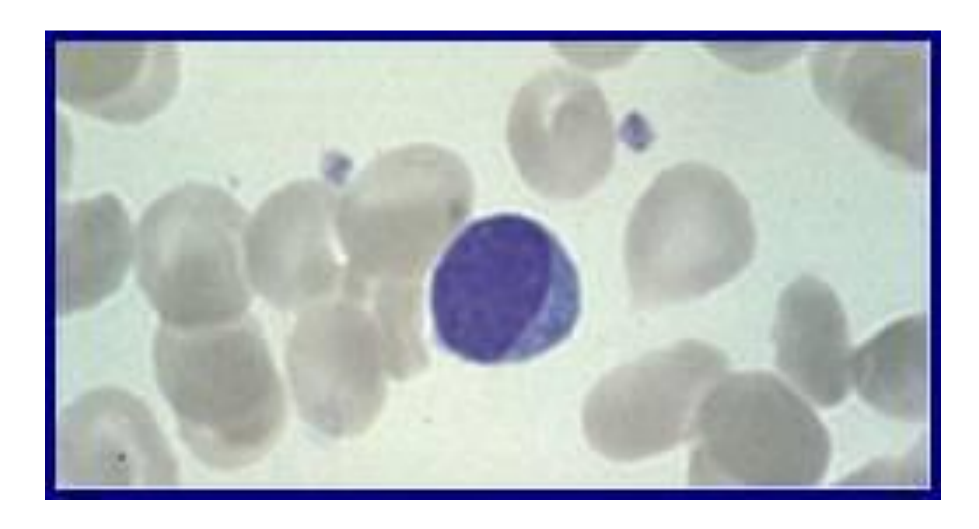
#### **Defensins**

- Polypeptides with antibiotic-like effect against bacteria and fungi.
- Produced mainly by granulocytes and by epithelial cells.
- Lead to desintegration of cell membranes, formation of pores in membranes.
- Main groups are  $\alpha$  and  $\beta$  defensins

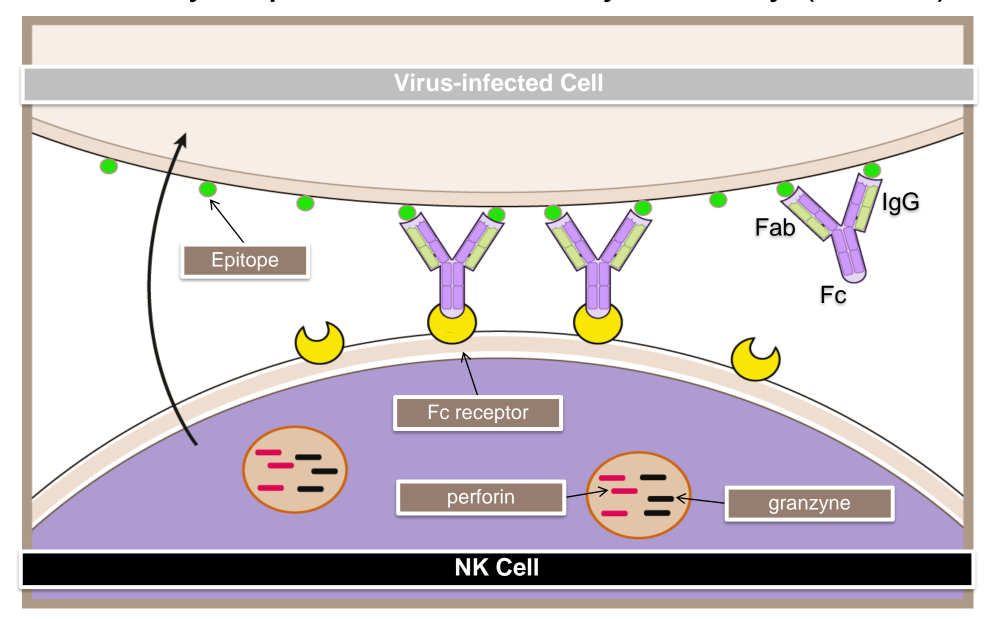
# Natural killers (NK cells)

- Originate in non-T non-B lymphocyte lineage.
- Morphologically: large granulated lymphocytes (LGL).
- Recognition of target cells in antigen non-specific.
- Virus infected and tumor cells are killed.
- Target cells are recognised mainly by decreased HLA-I expression.
- Cytotoxic mechanisms are similar to Tc cells: perforin and induction of apoptosis.
- Produce various cytokines, e.g. IFN-γ, IL-12

### Large granulated lymphocyte



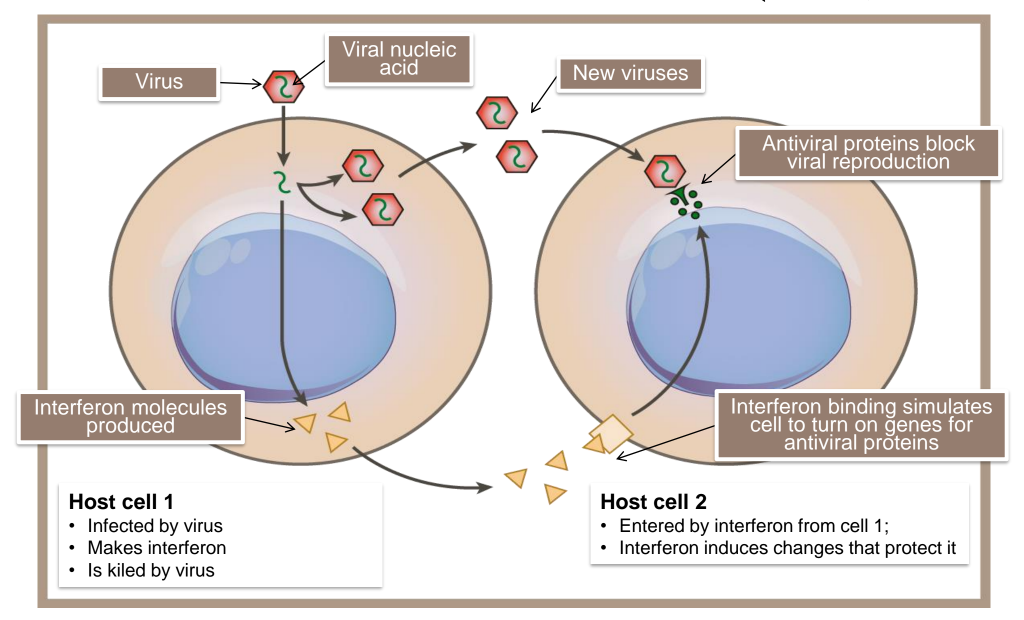
#### Antibody dependent cellular cytotoxicity (ADCC)



# Interferons (IFN)

- Type 1: IFN  $\alpha$ , IFN  $\beta$  produced mainly by virus-infected cells (fibroblasts, granulocytes. Lead to inhibition of the virus replication in target cells.
- Type 2 "Immune interferon": IFN  $\gamma$ : is produced by activated TH1+ cells, NK cells . Induces activation of macrophages.
- Interferon type 3 IFN  $\lambda$  IL28A,B, IL-29 similar to IFN-I

## The action of interferon (IFN)



#### Inflammation

- A rapid response to wounding and infection
- An important consequence of innate immunity
- Cardinal features
  - rubor (redness), calor (heat), tumor (swelling), dolor (pain)
- Local consequences of inflammation
  - Increased blood flow to affected area
  - Recruitment of phagocytes to affected area, particularly neutrophils and macrophages
  - Alteration of vascular permeability leading to entry of soluble molecules from the plasma

#### Local mediators of inflammation

- Products of activation of the kinin, complement, and coagulation systems.
   Usually C3a and C5a plays a significant role.
- Vasoactive amines histamin, serotonin reased from the damaged cells or stimulated macrophages.
- Metabolites of arachidonic acid
- Platelet activating factor
- Produkty of monocytes and granulocytes: IL-1, TNF-α, IL-6, IL-18, chemokines, NO,
- Produkts of activated lymphocytes: TNF-α, IL-6, IFN-γ, chemokines

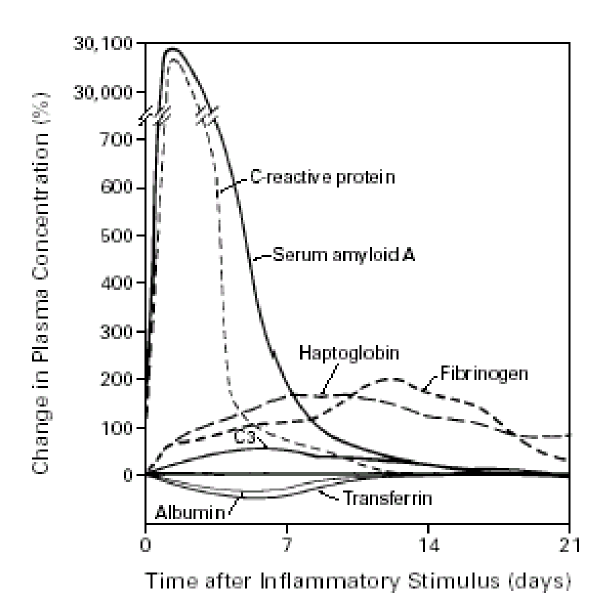
# General symptoms and signs of inflammation

- Orchestrated mainly by IL-1, IL-6, TNF-α
- Fever
- Fatigue, somnolence
- Loss of appetite
- Laboratory signs: leukocytosis, increased ESR, increase in accute phase proteins, decreased levels of iron and zinc in serum.

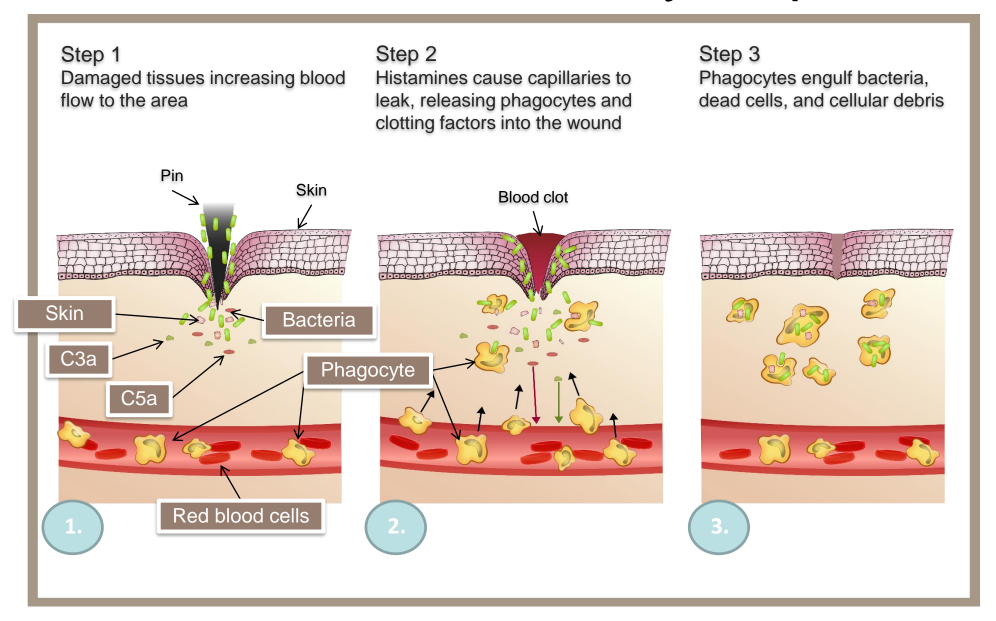
## Accute-phase proteins

- Serum levels are increased during inflammation
- Produced by the liver after stimulation by IL-1, IL-6, TNF- $\alpha$
- Best known: C-reactive protein
- Others: Complement components, A1-AT, fibronectin..

## Accute phase response



# Initiation of inflammatory response



## Drugs modulating inflammatory process

- Glucocorticoids
- Non-steroidal anti-rheumatic (anti-phlogistic) drugs (acidosalicylic acid, paracetamole,...)
- Antimalarics
- Gold
- Monoclonal antibodies against inflammatory cytokines and adhesion molecules