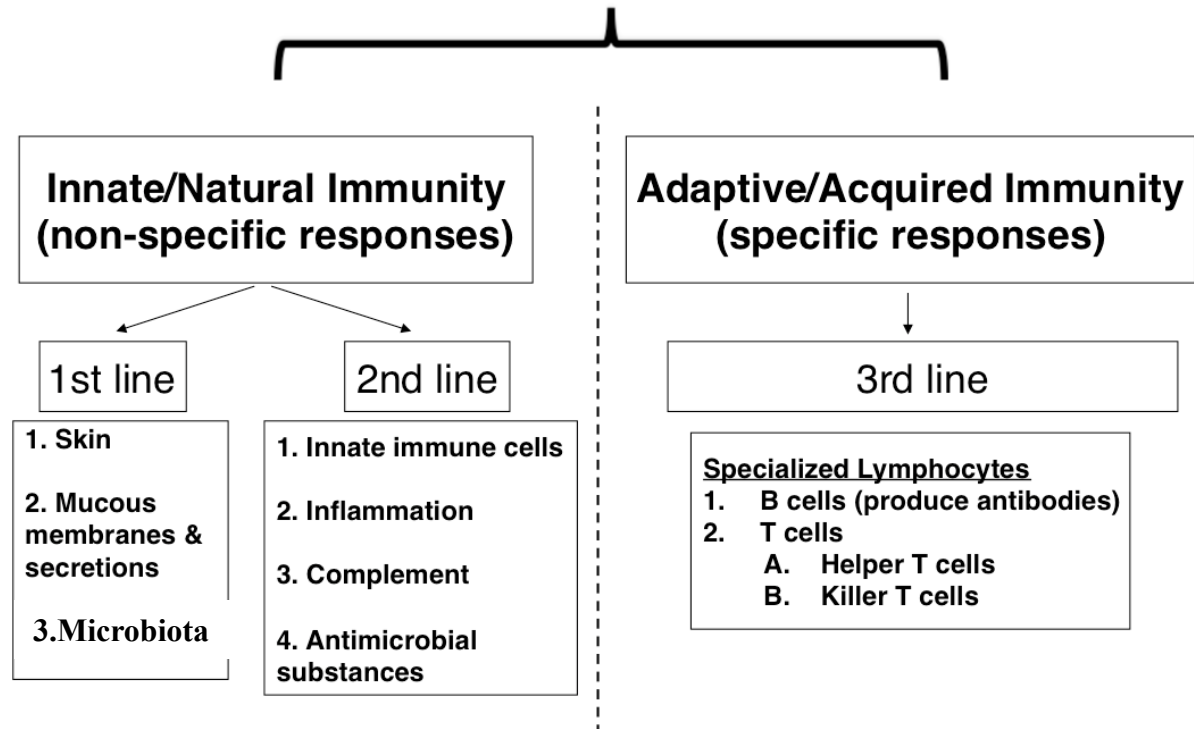


Chapter 9: Pathogen recognition in innate immunity

Properties of the Host : the immune system

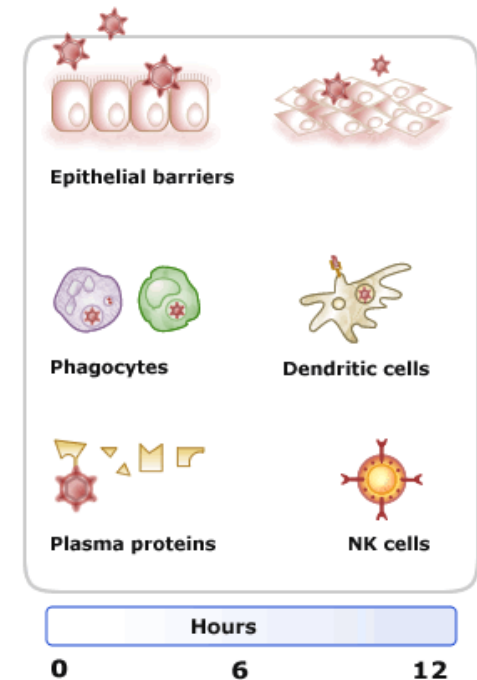
We are constantly being exposed to millions of potential pathogens daily, through contact, ingestion, and inhalation, but our immune system enables us to resist infections.

Infections triggers the immune response: **Immune system**



Innate immunity

- Innate immunity responses are the first defense against invading pathogens and spring into action immediately after an infection begins.
- During the first critical hours and days of exposure to a new pathogen, we rely on our **innate immune system** to protect us from infection.
- Innate immune responses are “**Inherent to the host**”. They do not depend on the host’s prior exposure to the pathogen and **are not specific to a particular pathogen**.
- Their responses rely on the body's ability to recognize **conserved features of pathogens** that are not present in the uninfected host. Basic mechanisms that regulate them are conserved.
- When activated, innate immunity triggers inducible mechanisms: complement cascade, phagocytosis, and produces inflammatory response.
- In vertebrates induction of innate immune response is required to initiate specific adaptive immune responses.



Adaptive immunity

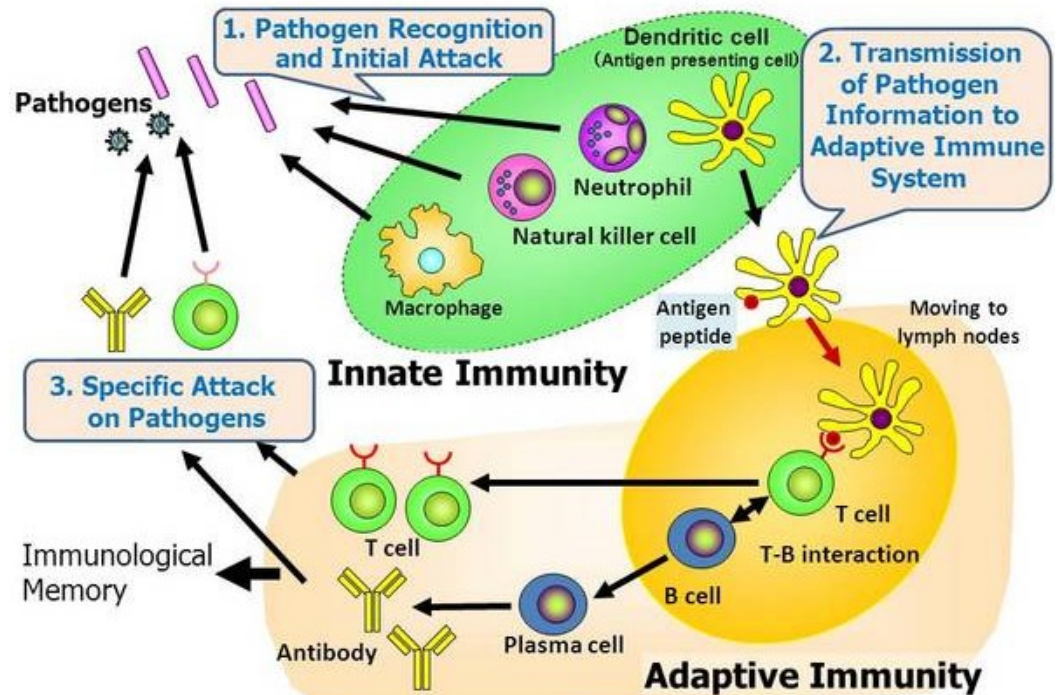
Adaptive mechanisms: not immediately ready to come into play. It must be turned on by host exposure to a pathogen (as during an infection).

Responses take at least some days to activate.

Highly specific: **directed against the invading pathogen** (immunological response based on antibody)

Diversity is extensive and resulting in a wide range of antigen receptors.

Memory is present: subsequent exposures to the same agent induce amplified responses



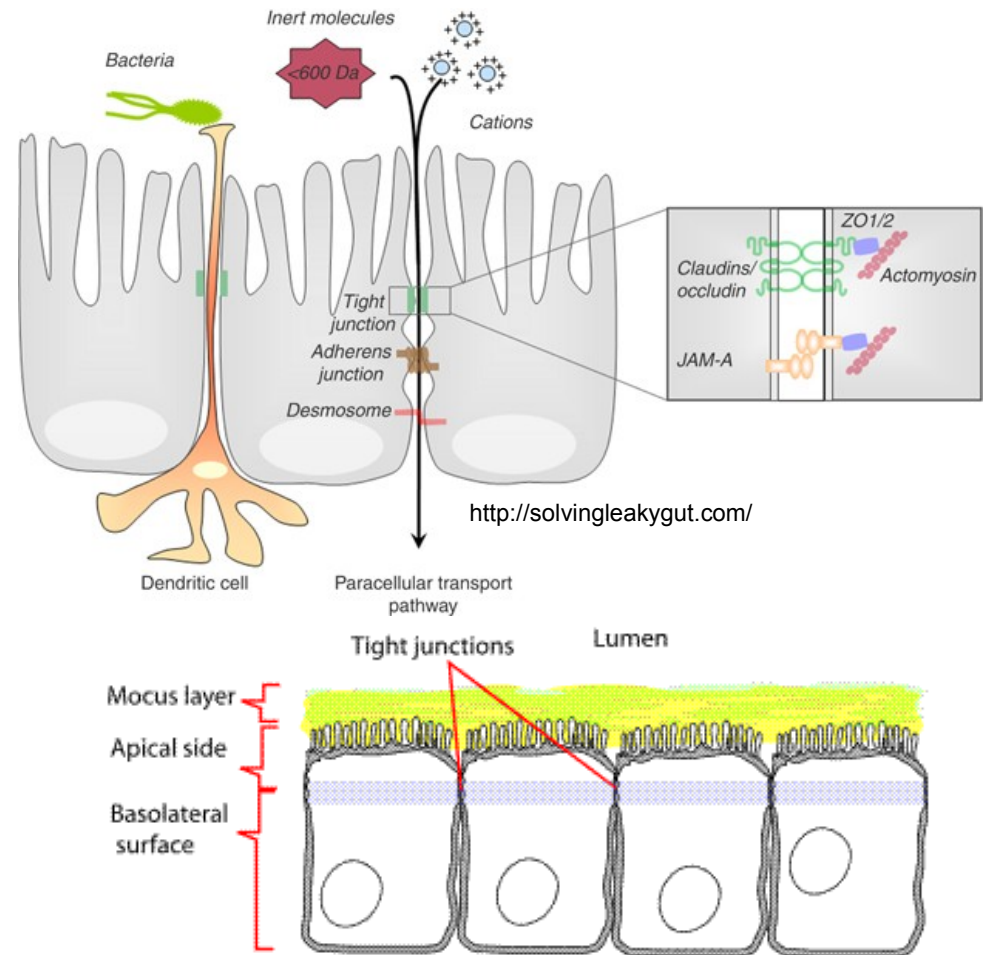
http://akira-pj.lserp.osaka-u.ac.jp/info_en/cafe_third.html

Epithelial Surfaces Help Prevent Infection

The skin and other epithelial surfaces, including those lining the lung and gut provide a **physical barrier** between the inside of the body and the outside world. No microbes can penetrate intact human skin, events have to breach the normal integrity of the skin.

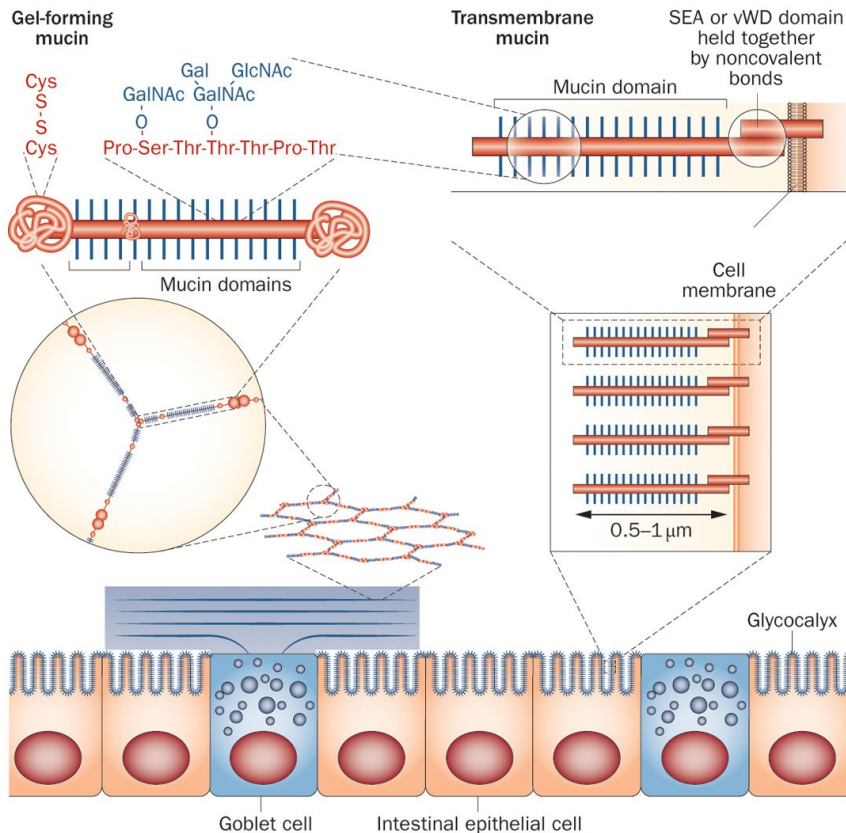
Epithelia surfaces: **tight junctions** between neighboring cells prevent easy entry by potential pathogens.

The interior epithelial surfaces are also covered with a **mucus layer** that protects these surfaces against microbial insults. It also facilitates their **clearance by beating cilia** on the epithelial cells.



Gel-forming mucin and transmembrane mucins attached to the apical membrane of cells

All the parts of mucosal surfaces are protected by mucin.



Mucins are a family of high MW glycoproteins. Mucin polymers are packed in the granules of the **goblet cells** found among the epithelia.

Two types: **gel-forming and transmembrane mucins.**

Mucin's key characteristic is to form **gels** for protection and lubrication. Secretion expands to form flat ring-like structures that stack under each other.

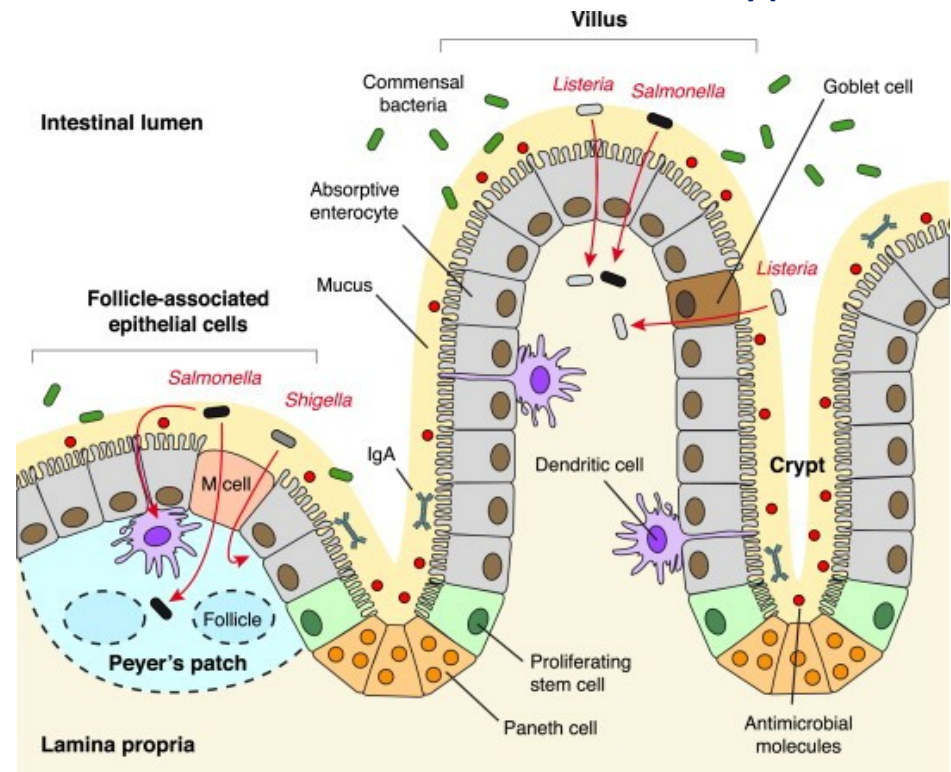
The mucus layer also contains substances such as **lysozyme, lactoferrin, and antimicrobial peptides** that kill pathogens or inhibit their growth.

The epithelium of the small intestine

Intestinal epithelial cells maintain a physical barrier against normal microbiota, The epithelium of the small intestine is composed of absorptive enterocytes, goblet cells, M cells, as well as proliferating stem cells and **Paneth cells** located in intestinal crypts.

GALT: gut-associated-lymphoid tissue: is a specialized lymphoid tissue allowing constant sampling of the luminal microbiota through **M cells**. Of the follicle-associated epithelium. Lymphocytes of GALT make **secretory IgA**.

Translocated bacteria (by transcytosis) are thus exposed to macrophages, dendritic cells (DCs), and B lymphocytes, they are usually captured, killed, processed, and presented to the immune system.



How bacterial pathogens colonize their hosts and invade deeper tissues
David Ribeta, b, c, , , Pascale Cossarta, b, c, ,

Article | Published: 01 April 2001

Dendritic cells express tight junction proteins and penetrate gut epithelial monolayers to sample bacteria

Maria Rescigno, Matteo Urbano, Barbara Valzasina, Maura Francolini, Gianluca Rotta, Roberto Bonasio, Francesca Granucci, Jean-Pierre Kraehenbuhl & Paola Ricciardi-Castagnoli ✉

Nature Immunology **2**, 361–367 (2001) | [Download Citation](#) ↓

Abstract

Penetration of the gut mucosa by pathogens expressing invasion genes is believed to occur mainly through specialized epithelial cells, called M cells, that are located in Peyer's patches. However, *Salmonella typhimurium* that are deficient in invasion genes encoded by *Salmonella* pathogenicity island 1 (SPI1) are still able to reach the spleen after oral administration. This suggests the existence of an alternative route for bacterial invasion, one that is independent of M cells. We report here a new mechanism for bacterial uptake in the mucosa tissues that is mediated by dendritic cells (DCs). DCs open the tight junctions between epithelial cells, send dendrites outside the epithelium and directly sample bacteria. In addition, because DCs express tight-junction proteins such as occludin, claudin 1 and zonula occludens 1, the integrity of the epithelial barrier is preserved.

Cells of the innate immune system

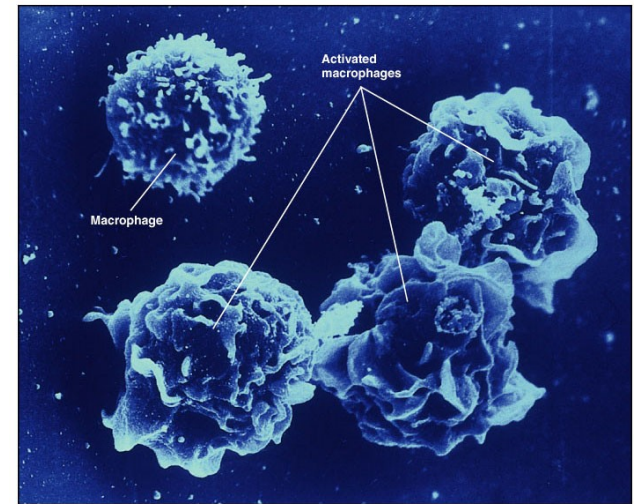
Skin and mucosal surfaces are highly effective in preventing pathogenic bacteria from entering the tissue and blood but from time to time, bacteria succeed in breaching these surfaces. Bacteria that get this far encounter a formidable defense force: the phagocytic cells and NK cells.

Macrophages reside in tissues throughout the body. These **long-lived cells** patrol the tissues of the body. Main function: to phagocytize (engulf) pathogens and particles. They also can alert and attract other immune cells. Specialized cells: In liver: Kupffer cells, in lungs: alveolar macrophages.

Monocytes: reside in the blood, ingest and kill bacteria, migrate to tissues. Precursors of macrophages.

Neutrophils, are differentiated, short-lived cells, which are abundant in blood but rapidly recruited to sites of infections. Neutrophils can phagocytize particles, release a respiratory burst and produce inflammatory cytokines.

Dendritic Cells are also phagocytic cells, but they have the special ability of initiating an adaptive immune response through a phenomenon called “antigen presentation” and through cytokine production (will be discussed later).

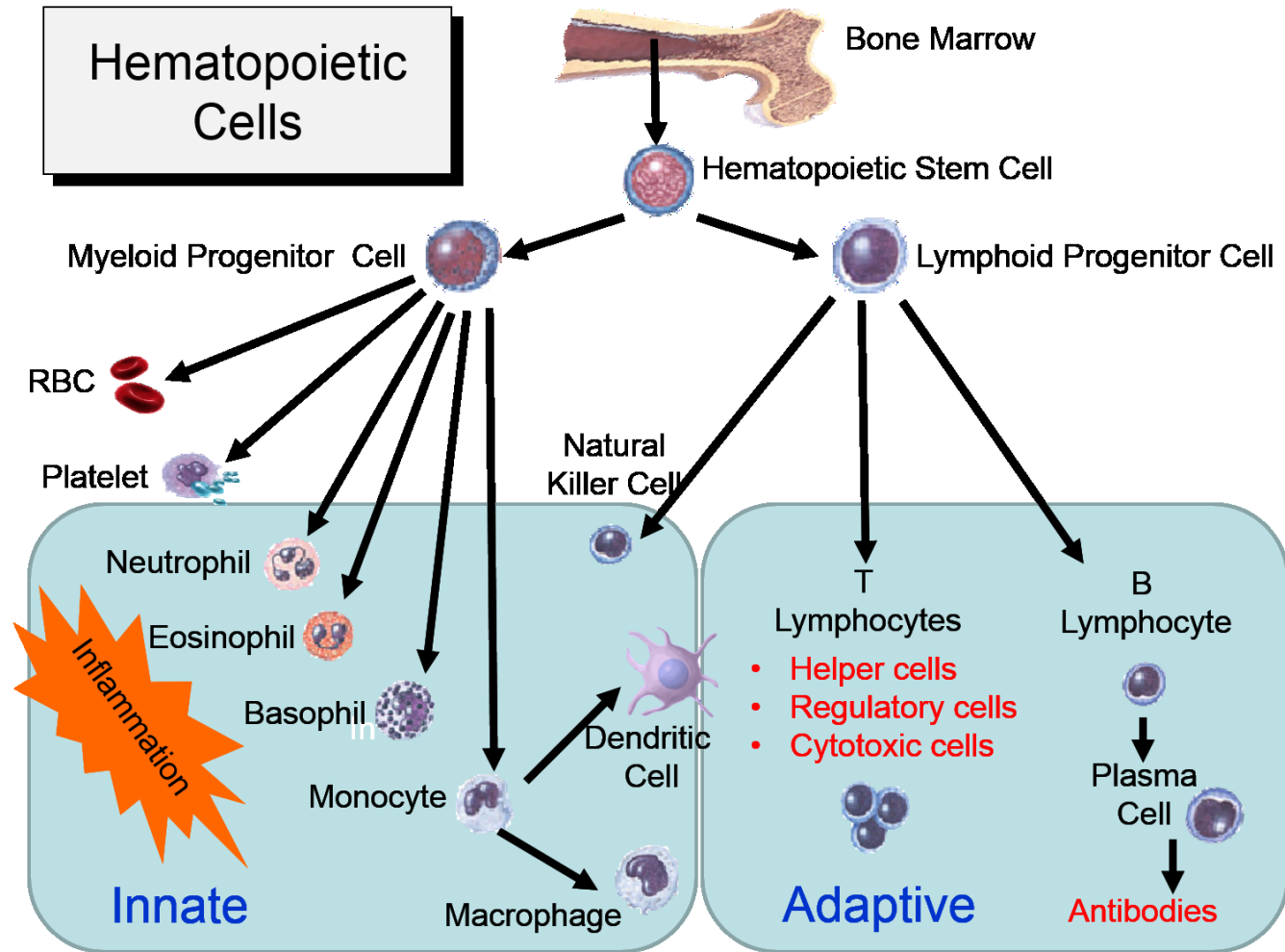


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macrophages



Cells of the Innate and Adaptive Immune Systems



Non-cellular Systems of the Innate Immune Response

Besides cells, there are also defenses in your body that are ready to react to pathogens as soon as they are encountered. These systems rely on small proteins that are found within the body fluids.

Complement system: The liver synthesizes the proteins of the complement system and they work in concert to aid in phagocytosis, bacteria lysing and immune cell attraction. It is a self-assembling machine that starts to assemble as soon as the first proteins are bound and in place.

Acute Phase Proteins: produced by the liver during inflammation when pro-inflammatory cytokines are produced. Many are designed to coat pathogens and have chemotactic properties. Some inhibit microbial growth by sequestering iron from the environment.

Anti-microbial Peptides: they function as natural antibiotics and are produced by cells that guard the external surfaces and internal surfaces such as the skin and the gastrointestinal system.



Human alpha defensin 1



Human beta defensin 1



Human cathelicidin LL-37



Human Cells Recognize Conserved Features of Pathogens

Innate and adaptive immune systems have to recognize and destroy infective microorganisms, without harming the host. Consequently, the immune systems must be able to distinguish **self from non-self**.

The innate immune system relies on the recognition of particular types of molecules or parts of them that are **common to many pathogens** but are absent in the host. Overall these molecules are called **pathogen-associated molecular pattern (PAMP)**. (The term Microbe-Associated Molecular pattern, MAMP, has also been proposed).

PAMPs are **essential structures** for the survival and therefore are **difficult to alter** for the microorganism. The various classes of pathogen-associated immunostimulants often occur on the pathogen surface in repeating **patterns**

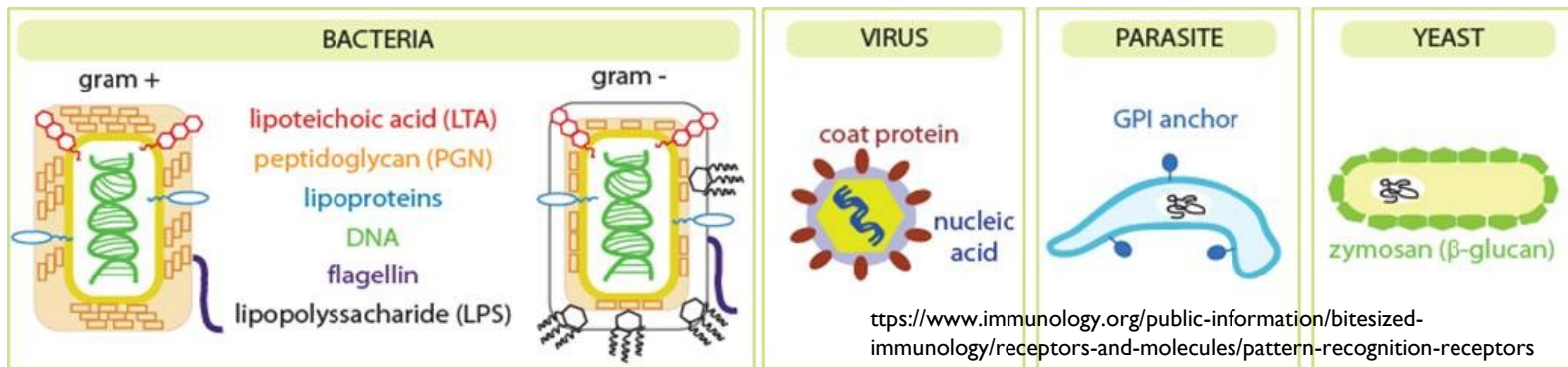
Various types of **PAMPs** are able to stimulate two types of innate immune responses: **inflammatory responses** (discussed below) and **phagocytosis** by cells such as neutrophils and macrophages.



Pathogen-associated molecular pattern

Microbial-associated immunostimulants are of various types:

- **Formylmethionine-containing peptides:** Procaryotic translation initiation differs from eucaryotic translation initiation in that formylated methionine. Any peptide containing formylmethionine at the N-terminus must be of bacterial origin.
- **The peptidoglycan cell wall** as well as **lipopolysaccharide (LPS)** on Gram-negative bacteria and **teichoic acids** on Gram-positive bacteria. **Flagellin** is also recognized.



- **Chitin** and other **β -glucans** (zymosan) which constitute cell walls of fungi
- **Short sequences in bacterial and viral DNA.** For example: “**CpG motif**” consists of the **unmethylated** dinucleotide CpG flanked by conserved motif. This sequence is highly less common in vertebrate DNA than in bacterial DNA. **Ss** and **Ds RNA** of viral origin.

PRRs: Pattern Recognition Receptors

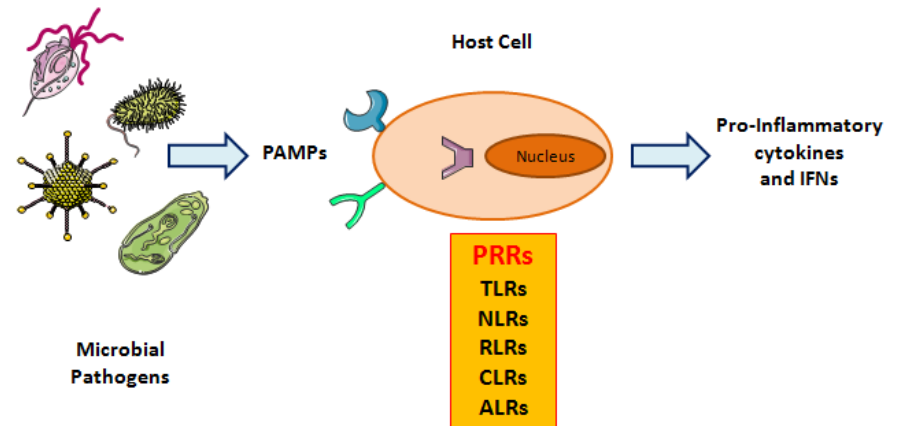
The innate immune system recognizes various classes of PAMPs/MAMPs via a limited number of well-conserved ancient receptors: **pattern-recognition receptors (PRRs)**.

PRRs possess common characteristics:

They are germline encoded, independent of immunologic memory and typically not clonally distributed.

PRRs are **expressed constitutively** in the host. They are mainly expressed by antigen presenting cells such as dendritic cells and macrophages, but they are also found in other immune (neutrophils, monocytes, lymphocytes) and non-immune cells.

The complex interactions between components allows a certain degree of pathogen-specific tailoring of the innate immune responses.



Pattern Recognition Receptors (PRRs)

<http://www.inbionet.eu/outreach-news/the-innate-immune-system/n>

Different class of PRRs act in concert

PRRs are **strategically localized** in the cell. There are present at the cell surface to recognize extracellular pathogens such as bacteria or fungi, in the endosomes where they sense intracellular invaders such as viruses and finally in the cytoplasm.

The PRR are divided in four families:

Toll-like receptors (**TLR**)

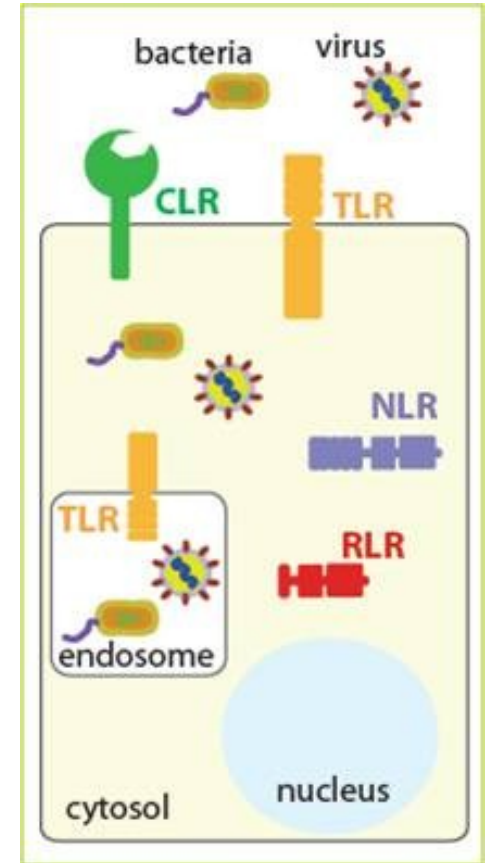
Nucleotide oligomerisation receptors (**NLR**)

RIG-I like receptors (**RLR**)

C-type lectin receptors (**CLR**)

This repertoire of molecules and cells **acts in concert** to form a network of host defense mechanisms. Number of different PRRs are engaged by a given pathogen via various PAMPs, hence securing a rapid and potent inflammatory response and also allowing for some specificity of the response.

The recognition of PAMPs by PRRs leads to production of pro-inflammatory cytokines, IFNs, activation of phagocytes which promotes activation of antigen-presenting cells that elicits adaptive immunity.



Toll-like Proteins Are an Ancient Family of Pattern Recognition Receptors

Many of the mammalian cell-surface pattern recognition receptors responsible for triggering host cell gene expression in response to pathogens are members of the **Toll-like receptor (TLR)** family.

Toll receptor has been originally discovered in *Drosophila*. It is involved in the adult fly's resistance to fungal infections.

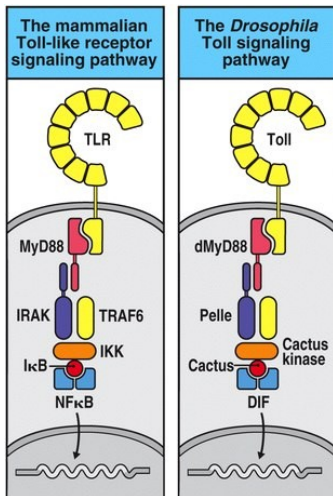
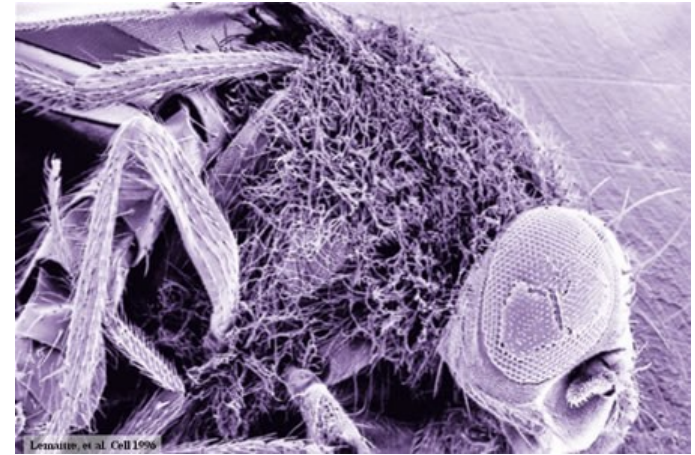


Figure 15-4 Immunobiology, 6/e. (© Garland Science 2005)

Intracellular signal transduction pathway activated downstream of Toll when a fly is exposed to a pathogenic fungus leads to activates the transcription of various genes, including those encoding the antifungal peptides: defensins.

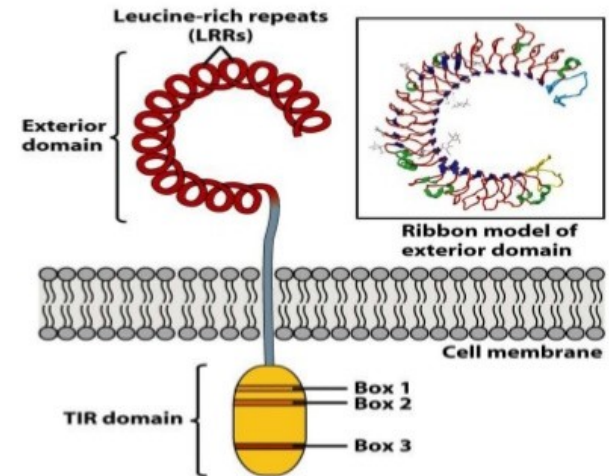
Toll was discovered in 1996 when in *Drosophila* a developmental gene *toll*, was knocked out. When this gene was knocked out it was apparent that the animals or the fruit flies succumbed to massive fungal infection

TLRs are evolutionarily conserved from the worm *Caenorhabditis elegans* to mammals. The human TLR family comprise 10 members, 12 in mouse.

The structure of a toll-like receptor

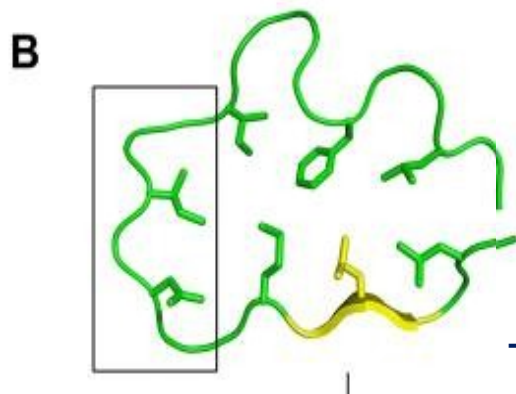
Toll-like receptors (TLRs) are type I integral membrane receptors, non-catalytic receptors that recognize structurally conserved molecules derived from microbes.

Three domains: 1) the extracellular **Leucine-Rich Repeat (LRR)** domain 2) short cysteine rich patches with short transmembrane portion, and 3) a conserved cytoplasmic signaling domain Toll/IL-1R homology (TIR domain).



LRR consensus sequences for TLR3

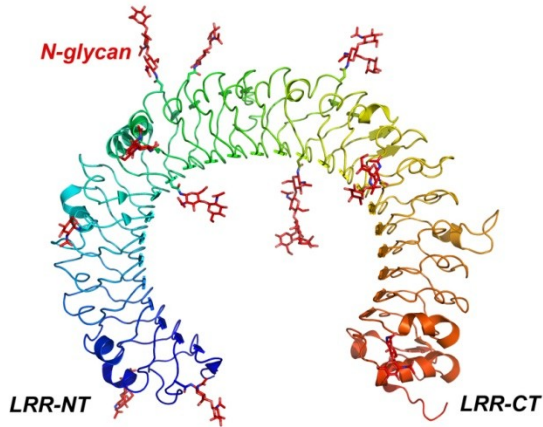
Leucine-Rich Repeat domain is composed of 19–25 tandem repeats of 22–30 residues in length and contains a conserved hydrophobic motif **XLXXLXLXX** spaced at distinctive intervals.



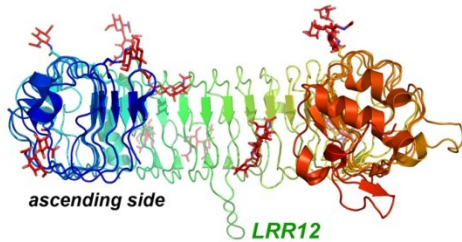
All LRRs adopt a **loop structure** (in figure TLR3), beginning with an extended stretch that contains three residues in the β -strand configuration (yellow), followed by an alpha-helical.

The boxed regions form the surfaces involved in ligand binding.

Structure of the TLR3 extracellular domain



When assembled into a protein, multiple consecutive LRRs form a **solenoid structure**, in which the consensus hydrophobic residues point to the interior to form a stable core and the β strands align to form a hydrogen-bonded parallel β sheet

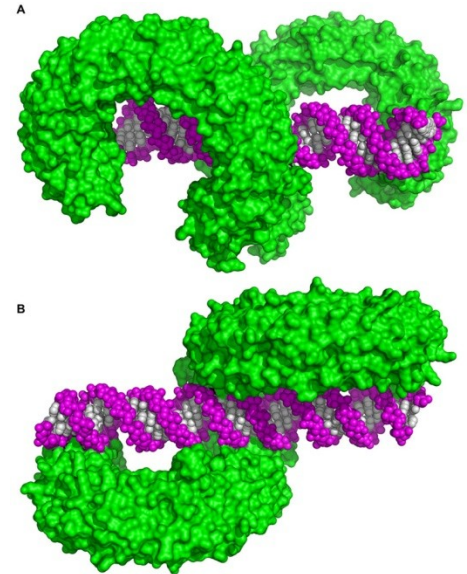


The structure of a TLR-ECD (hTLR3)

Structure. 2011 Apr 13; 19(4): 447–459.

Upon ligand binding, typically two extracellular domains form an “m”-shaped dimer sandwiching the ligand molecule bringing the transmembrane and cytoplasmic domains in close proximity and triggering a downstream signalling cascade.

Although the ligand-induced dimerization of these receptors has many common features, the nature of the interactions of the TLR extracellular domains with their ligands varies markedly between TLR paralogs.



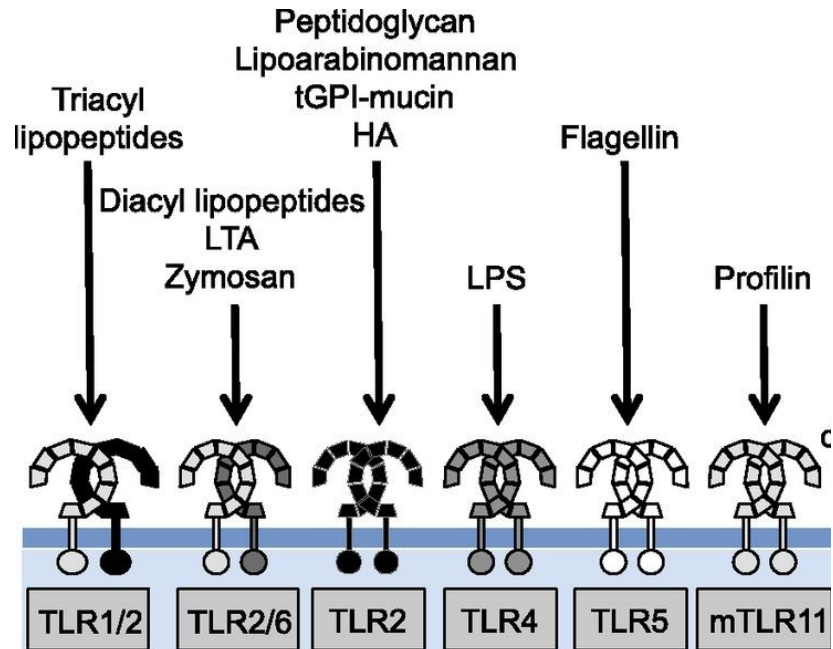
Structure of the TLR3/dsRNA complex

Structure. 2011 Apr 13; 19(4): 447–459.

Multiple TLRs are expressed in Humans

The human TLR family comprise 10 members (TLR1-TLR10). They are expressed on macrophages, neutrophils, DCs, B cells, specific types of T cells, and intestinal epithelial cells.

Expression of TLRs is not static but rather is modulated rapidly in response to pathogens, a variety of cytokines, and environmental stresses. Plasma membrane associated TLR: **TLR1, 2, 4, 5** and **6** recognise **microbial surface PAMPs**. They functions as homodimer or heterodimers.



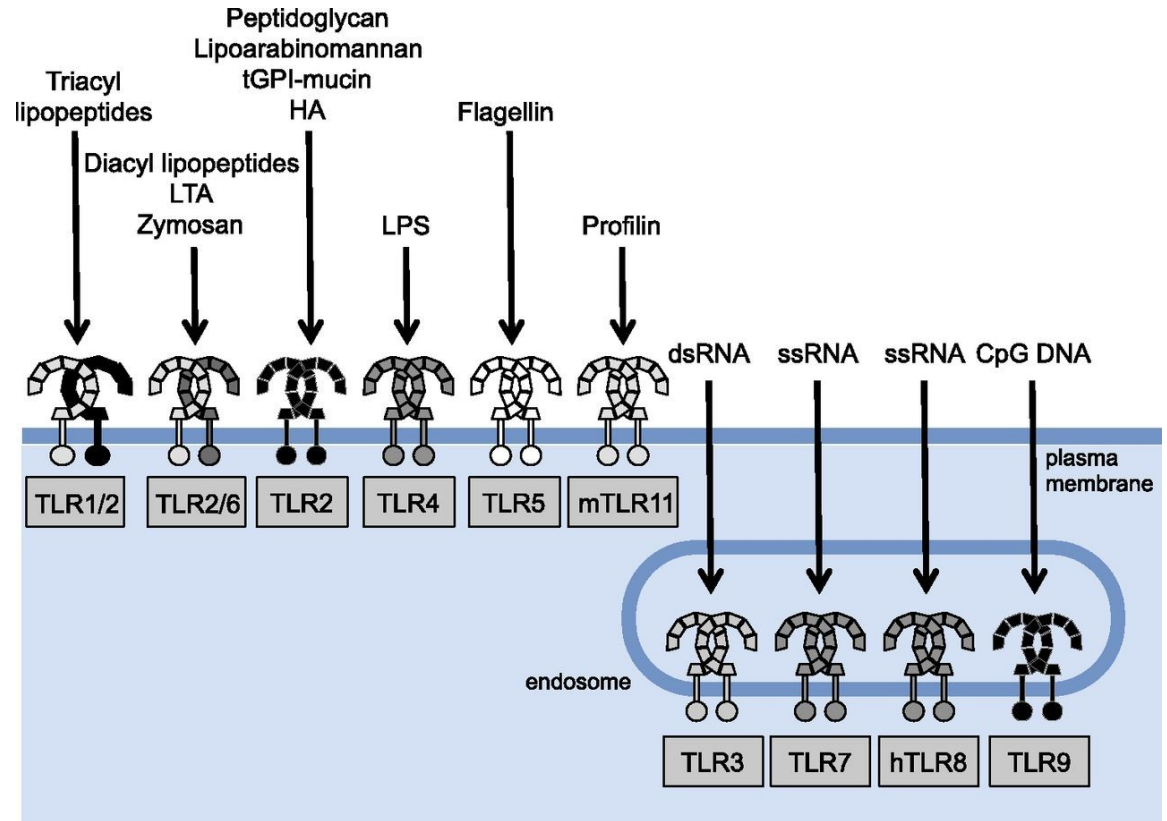
TLR2 recognizes its ligands by forming a heterodimer with either TLR1 or TLR6. The resulting complexes recognize distinct ligands: TLR1/2 triacyl lipopeptides and lipoproteins, TLR2/6 diacylipoproteins and zimosan

TLR4 recognizes lipopolysaccharide (LPS) on the cell surface. TLR5 recognizes flagellin. TLR-11 (mouse).

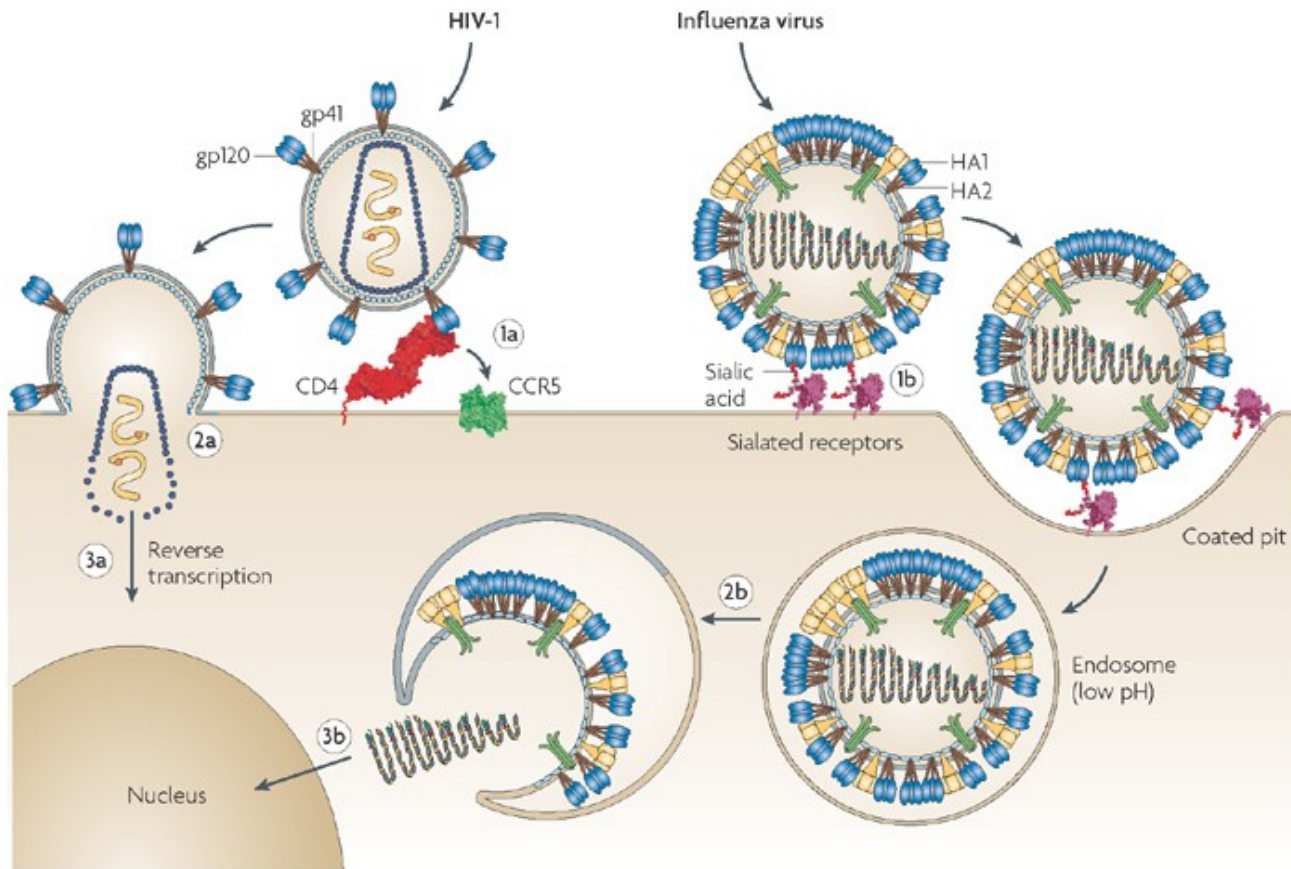
Endosome-associated TLRs

A set of TLRs, comprising TLR3, TLR7, TLR8, and TLR9, recognize nucleic acids derived from viruses and bacteria, as well as endogenous nucleic acids in pathogenic contexts.

Endosome-associated TLR:
TLR3 detects viral double-stranded (ds) RNA in the endosome;
Human heterodimeric **TLR7/8** complex recognize single-stranded (ss) RNAs from RNA viruses.
TLR9 senses unmethylated DNA with CpG motifs derived from bacteria and viruses.



Eukaryotic viruses entry pathways

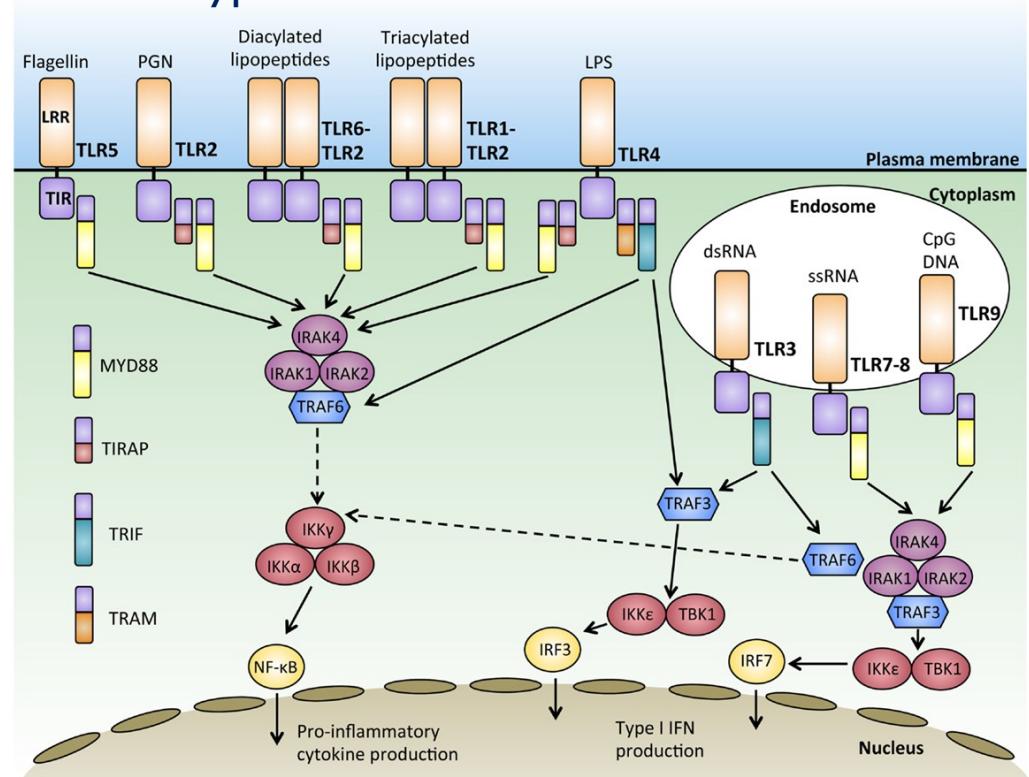


While some viruses can enter directly at the cell surface, (e.g. HIV-1) others exploit receptor-mediated endocytosis (e.g. influenza v) to bring them inside the cell, where viruses can eventually break free and begin replication

Signaling transduction pathway of TLRs

TLR-mediated microbial recognition is very important for host defense against pathogens. Recognition of PAMPs by TLRs leads to **transcriptional upregulation** of distinct genes, depending on the TLRs and cell types involved.

TLR signaling is roughly divided into two distinct pathways depending on the usage of the distinct adaptor molecules. Adaptor molecules **MyD88** and **TRIF** (see the figure) are responsible for the activation of distinct signaling pathways, culminating in the activation of transcription factors such as nuclear factor- κ B (NF κ B) and interferon-regulatory factors (IRFs), which regulate the production of pro-inflammatory cytokines and type I interferon (IFNs) respectively.



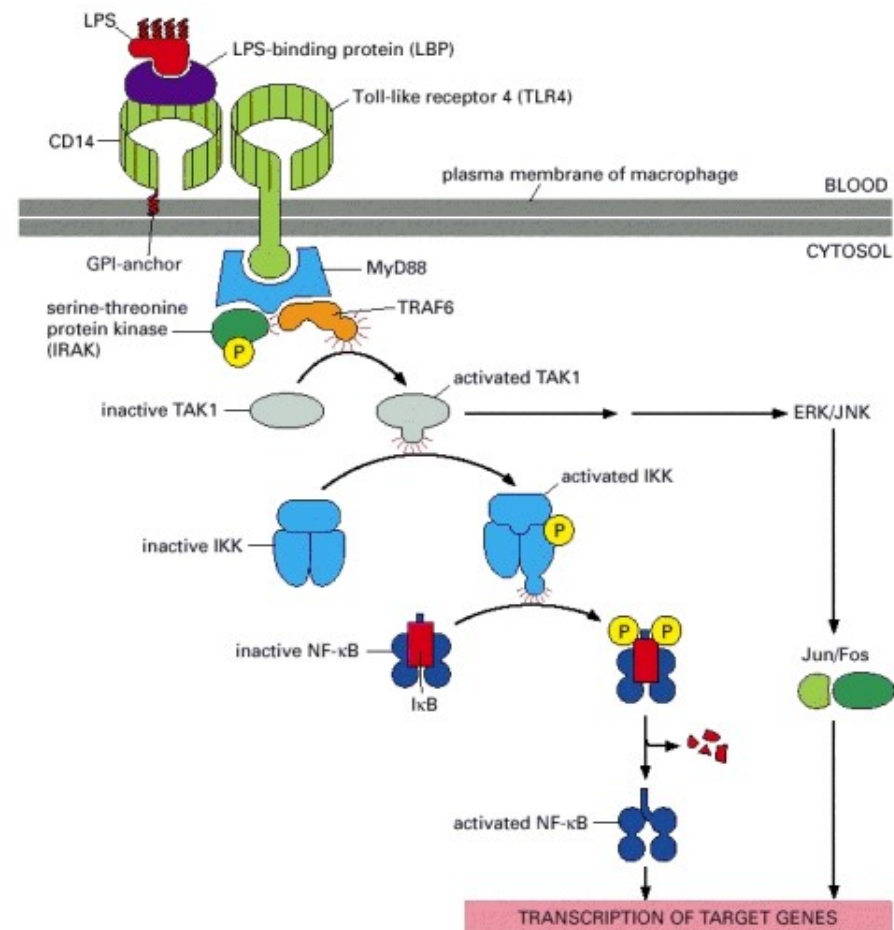
Stimulation with TLR3, TLR4, TLR7, and TLR9 ligands, but not the TLR2 ligand, induces type I IFN production, in addition to proinflammatory signals.

Activation of TLR4 by bacterial LPS.

LPS is recognized by the **TLR4** of the phagocytes in conjunction with the cell-surface co-receptor **CD14** and the soluble serum protein **LPS binding protein (LBP)**. LBP binds LPS in the blood and then it is bound by the receptor protein CD14 with leucine-rich repeat motifs.

TLR-4 binds to the CD14:LBP:LPS leading to recruitment of the adaptor proteins **MyD88** to the cytoplasmic domain of TLR4. The complex initiates a signaling cascade of phosphorylation events leading to activation of the kinase **IKK**. IKK phosphorylates **IκB**, an inhibitor bound to the transcription factor **NF-κB**.

P-IκB is degraded, releasing NF-κB, which migrates to the nucleus where it activates the transcription of proinflammatory genes.

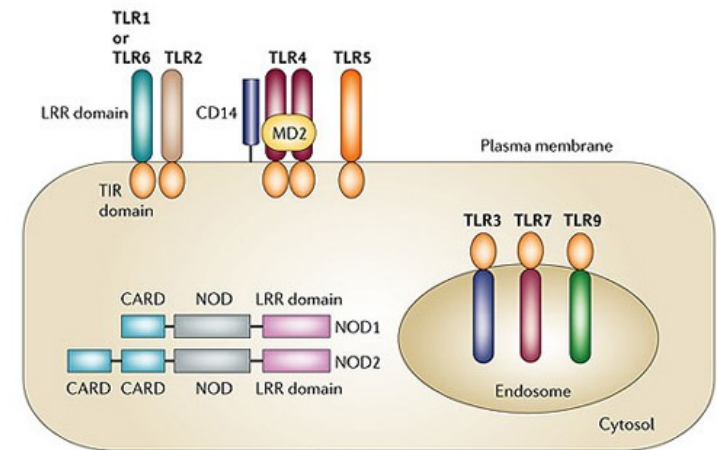


Cytoplasmic Pathogen Recognition System

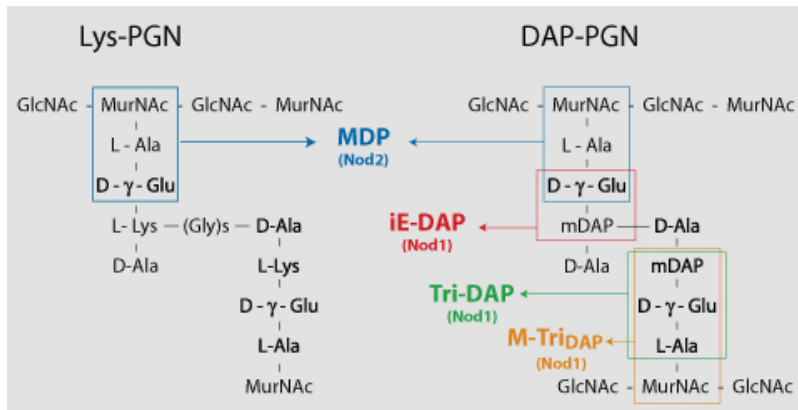
Pathogens that have invaded the cytosol are detected by a large family of cytoplasmic PRRs, which activate a number of signaling pathways.

NLR (NOD-like receptors) and **RLR (RIG-I-like receptor family)** are sensors for cytoplasmic pathogens.

NLRs: cytoplasmic proteins having a variety of functions in regulation of inflammatory and apoptotic responses. NLRs: C-terminal **leucine-rich repeat (LRR) region**, a central nucleotide-binding receptor (NOD) and a N-terminal protein-binding motifs (CARD). Signal transduction is mediated by CARD domain inducing transcriptional upregulation of proinflammatory cytokine genes.



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Nature Reviews | Immunology



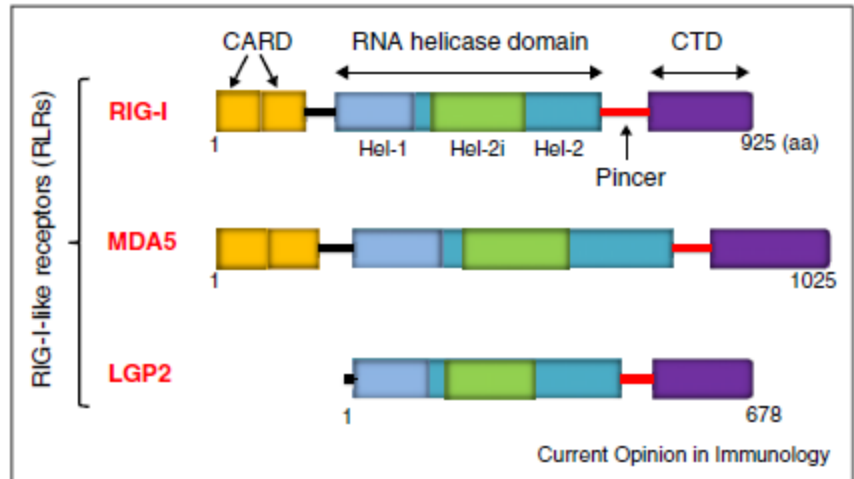
NOD-1 and **NOD-2** through LRR recognize distinct motifs derived from degradation of peptidoglycan (DAP-PGN) and **NOD-2** recognizes muramyl dipeptide (MDP) from bacteria peptidoglycan.

γ-D-glutamyl-meso-diaminopimelic acid (DAP)

RLRs and Virus Recognition

The CARD-helicase proteins/ **RIG-I-like** receptor family (RLRs) functions as an **intracellular PRR** serve to **recognize viruses** that have already entered into the cytoplasm of a cell, through the detection of viral replication by direct interaction with **ssRNA** and **dsRNA**.

RLR receptors (RIG-I) are RNA helicases that contain an N-terminal caspase recruitment domain (CARD) and a central helicase domain with ATPase activity required for RNA-activated signaling.



Current Opinion in Immunology 2015, 37:40–45

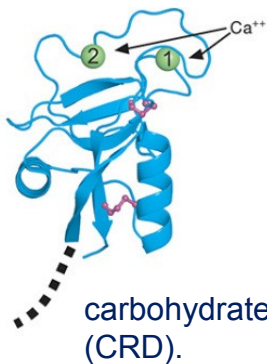
Binding of dsRNA or 5'-triphosphate RNA to the C-terminal domains of RLRs triggers signaling via CARD-CARD interactions, between the helicase and adaptor proteins, ultimately resulting in an antiviral response mediated by type I IFN production

C-type lectin receptors (CLR)

CLRs comprise large family of receptors that recognize **specific carbohydrate structures** (lectin activity) on microorganisms such as viruses, bacteria, and fungi. The lectin activity of these receptors is mediated by conserved **carbohydrate-recognition domain (CRD)**.

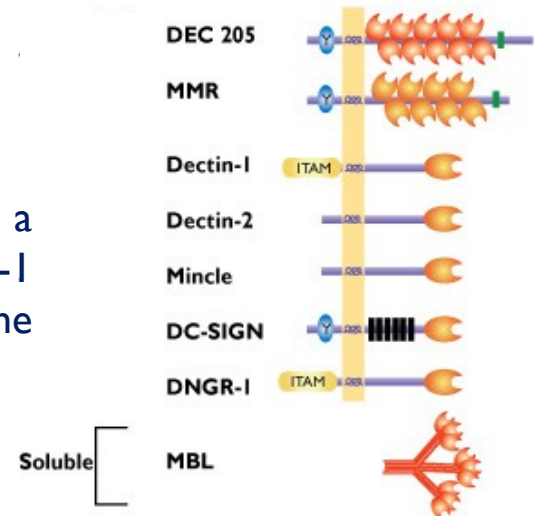
type I transmembrane proteins containing several CRDs or CRD-like domains. Macrophage mannose receptor (MMR) and DEC-205) are important in antigen uptake.

A



Type II transmembrane CLRs typically carry a single CRD domain and include Dectin-1 (specific receptor for β -glucans, Dectin-2 (the functional receptor for α -mannans).

CLR Types



<http://www.invivogen.com/review-clr>

Mannose-binding lectin (MBL) is a C-type lectin (collectins) with functions of soluble receptors for bacteria carbohydrates. The soluble CLR **Mannose binding lectin (MBL)** is an acute-phase serum protein that triggers one of the way to activate complement (**lectin pathway**).

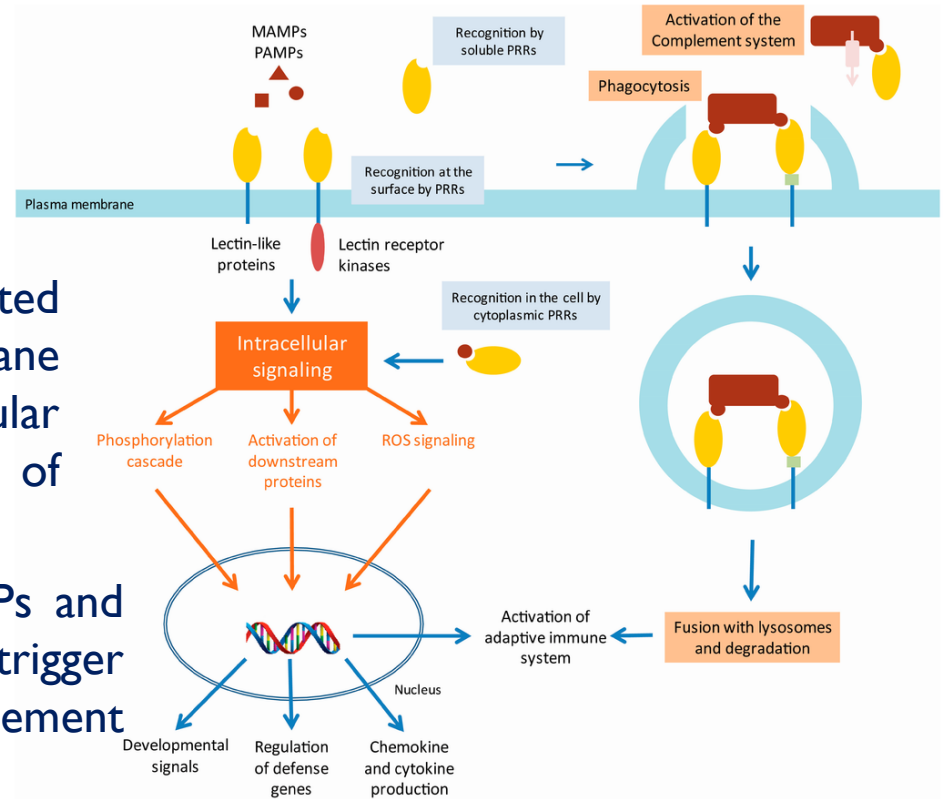


C-type lectin receptors (CLR)

CLRs signaling are involved in the **various steps for initiation of innate immune responses**

Detection of pathogen/microbe-associated molecular patterns (P/MAMPs) by membrane bound some CLR initiate both an intracellular signaling cascade and lead to phagocytosis of the pathogen

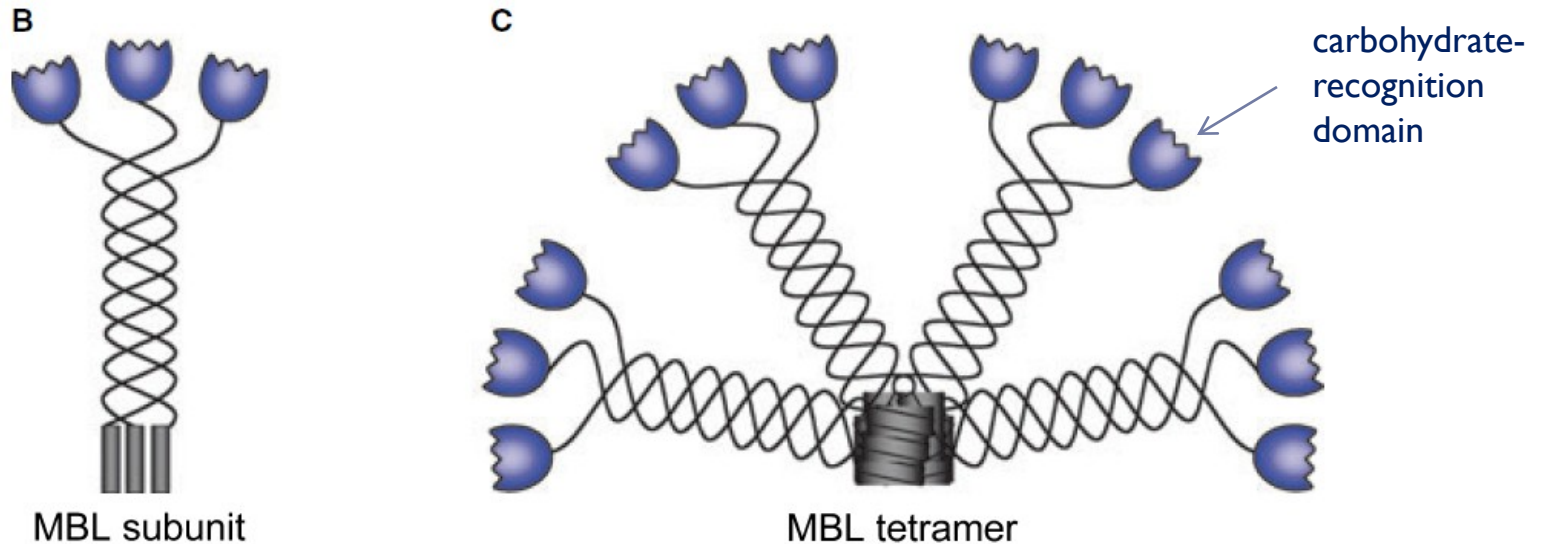
Soluble lectin CLR can recognize PAMPs and subsequently bind to receptors that will trigger phagocytosis or can activate the complement system.



Molecules 2015,
20(5), 9029-9053;



MBL structure and oligomerization



modified from 2009 John Wiley & Sons A/S
Immunological Reviews

MBL: as other collectins contain a collagen-like domain and usually assemble in large oligomeric complexes containing 9–27 subunits forms. MBL clusters of 2-9 carbohydrate-binding heads around a central collagen-like stalk. This assembly binds specifically to mannose and fucose residues in bacterial cell walls that have the **correct spacing and orientation** to match up perfectly with the number of carbohydrate-binding sites, providing a exemplary model of PRR.



Scheme of the complement activation systems

The complement system consists of about 20 interacting soluble proteins that are made mainly by the liver and circulate in the blood and extracellular fluid.

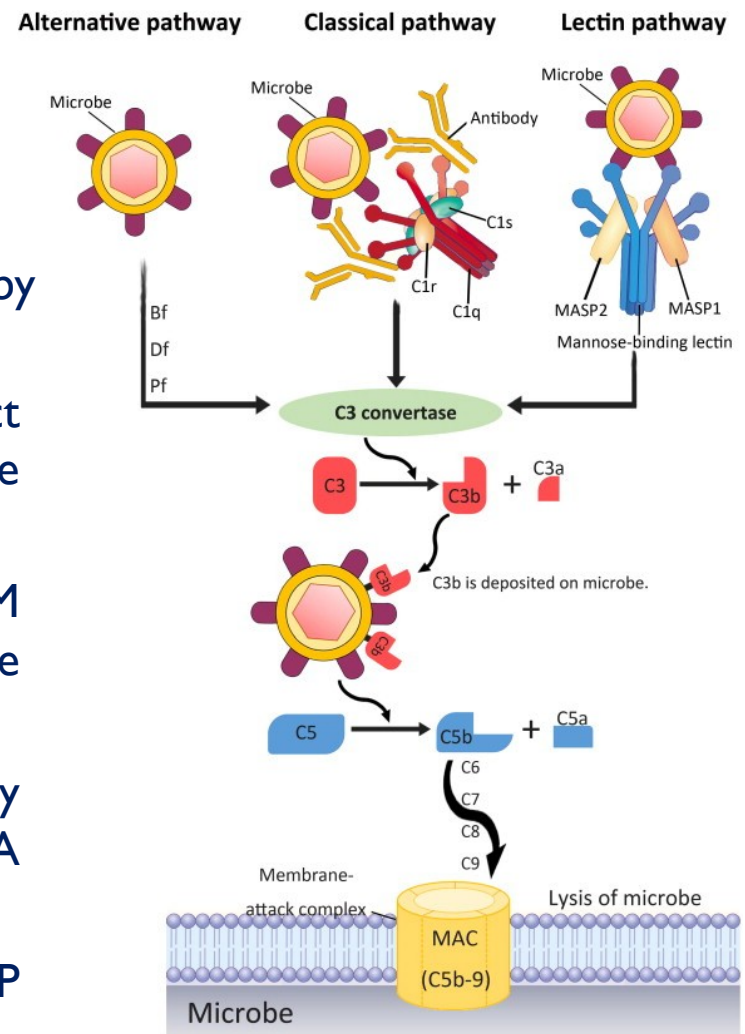
The principal stages in complement activation by the classical, lectin, and alternative pathways

Three distinct pathways of complement act locally to activate **C3** (C3a+C3b), which is the pivotal component of complement:

classical pathway: activated by IgG or IgM antibody molecules of the adaptive immune system bound to the surface of a microbe.

alternative pathway: C3 is spontaneously activated at low levels as C3b by LPS or TA found on the surfaces of bacteria.

Lectin pathway: activated by MBP/MASP complex (later)



The complement system

Three distinct pathways of complement act locally to activate **C3** (C3a+C3b), by the activity of the protease C3 convertase.

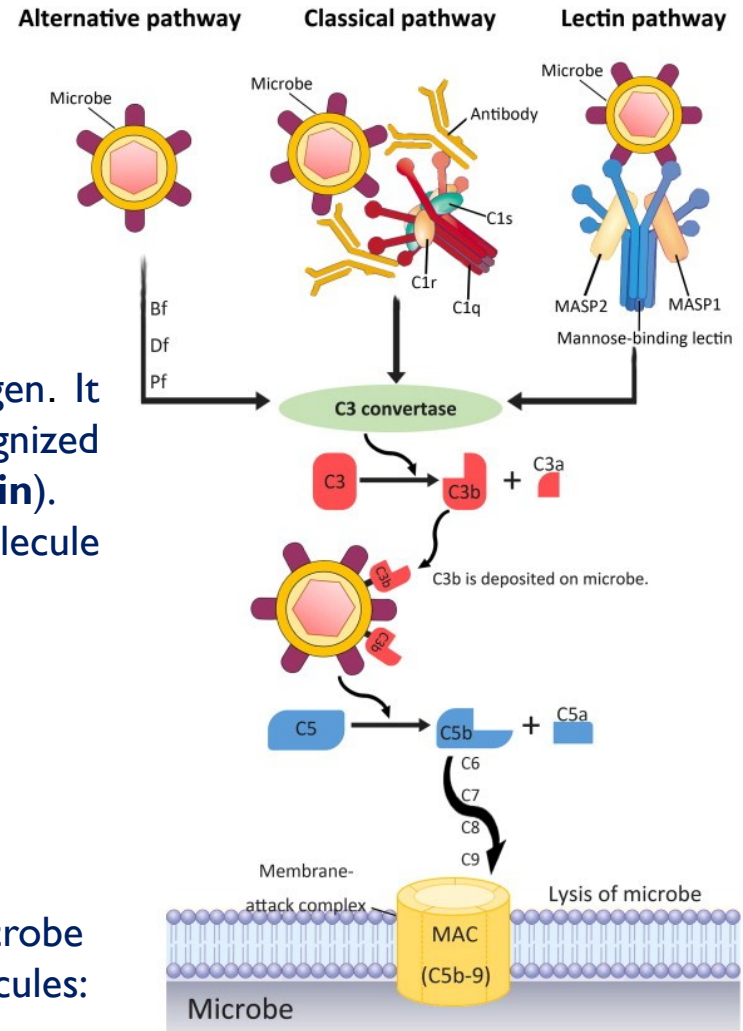
The early components and C3 are all proenzymes, which are activated sequentially by proteolytic cleavage (proteolytic cascade).

C3b, binds covalently to the surface of the pathogen. It acts as a protease (**C5 convertase**) but it is recognized by specific receptors (C3bR) on phagocytes (**opsonin**).

C3a is a major proinflammatory molecule (chemoattractant) and acts independently.

C5 convertase causes the cleavage of the **C5** to produce **C5a** (chemoattractant) and **C5b** which remains bound to C3 on the cell surface and assembles other late factors (C6, C7, C8)

A large transmembrane channel on microbe membrane is formed by the addition of C9 molecules: membrane attack complex, (**MAC**)

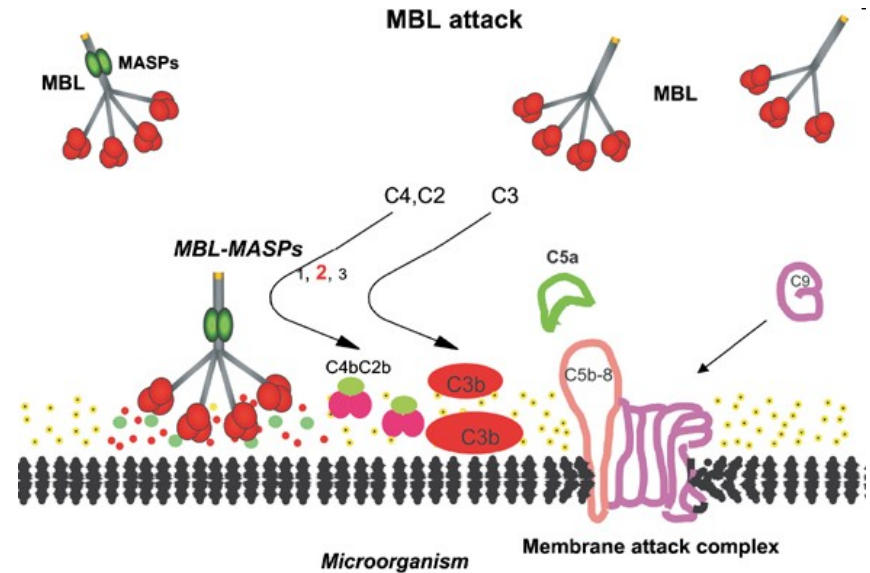


MBL activates lectin-pathway of complement

MBL when is activated by the binding of of mannose-containing glycans, complexed with the serine protease **MASPs** binds to sugar arrays on a microorganism.

Associated MASP is responsible for the release of C2a and C4b from the inactive complement factors C2 and C4.

The active complex C2aC4b form the **C3 convertase** that in turn cleaves C3 in **C3a** and **C3b**.



P Garred, F Larsen, J Seyfarth, R Fujita and H O Madsen. Genes and Immunity 2010

MBL-mediated complement attack complex. a complement attack through MASP. MASPs denote MBL-associated serine proteases.

