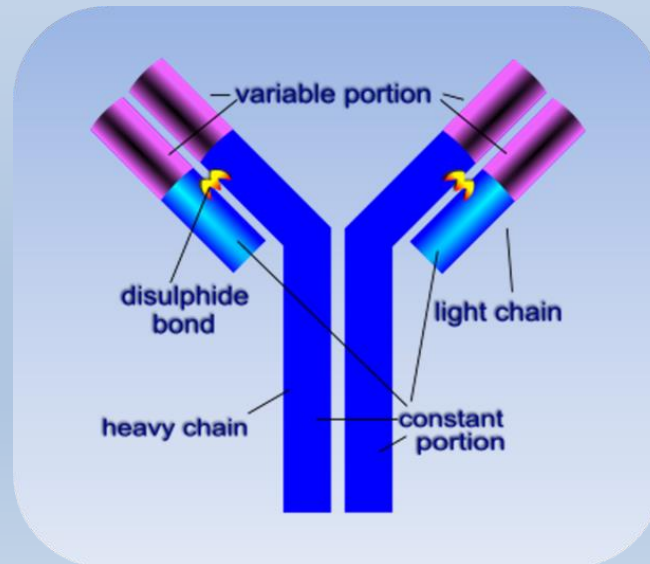


## Topics:

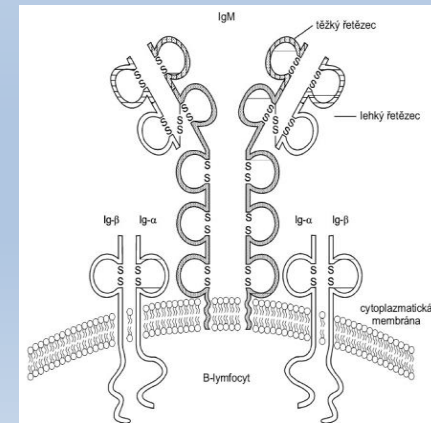
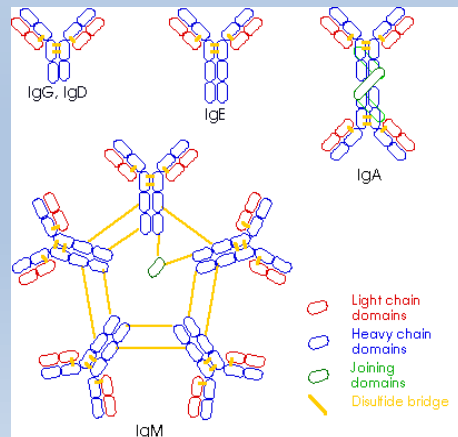
31. Immunoglobulins - structure.
32. Immunoglobulins - functions.
33. Genetic background of immunoglobulin production.
34. Biological and chemical characteristics of immunoglobulin classes IgG and IgA.
35. Biological and chemical characteristics of immunoglobulin classes IgM, IgD and IgE.
36. Isotype switching. Idiotypes and anti-idiotypes - their role. Immunological memory.
37. Ontogenesis of the immune response.
38. Primary immune response.
39. Secondary immune response. Effector functions of antibodies.

# Immunoglobulins (Antibodies)



# Immunoglobulins

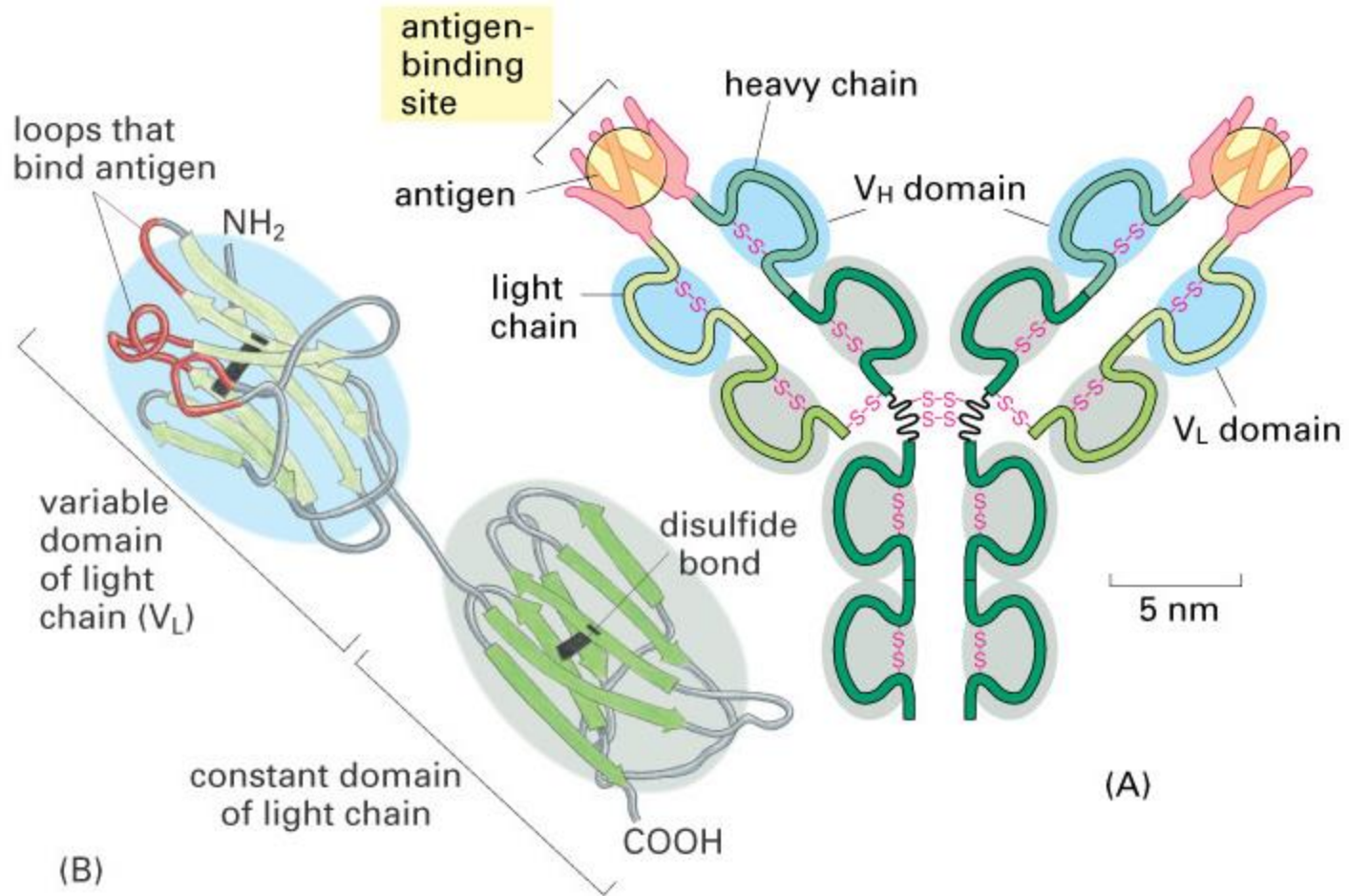
- Immunoglobulins or antibodies are glycoproteins, responsible for humoral part of **specific immune response**
- produced by B cells (plasma cells)
- secreted x membrane (BCR)



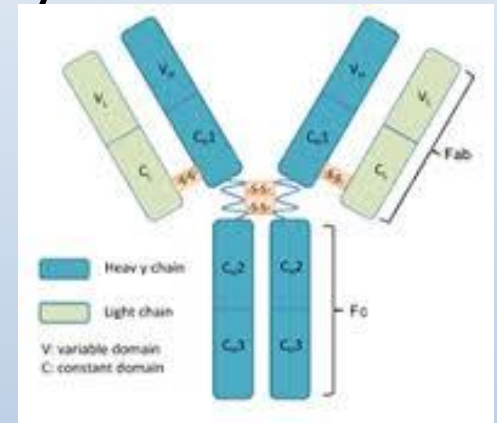
# Immunoglobulin structure

- 2 heavy (H) chains covalently linked by disulfide bonds, each H chain is connected to a light (L) chain by disulfide bonds
- H chain consists of 4 to 5 domains (1 variable, 3-4 constant)
- L chain consists of 2 immunoglobulin domains (1 variable, 1 constant)
  
- Types of L chains -  $\kappa$ ,  $\lambda$
- Types of H chains -  $\mu$ ,  $\delta$ ,  $\gamma$  ( $\gamma 1-4$ ),  $\alpha$  ( $\alpha 1$ ,  $\alpha 2$ ),  $\epsilon$

# Immunoglobulin structure



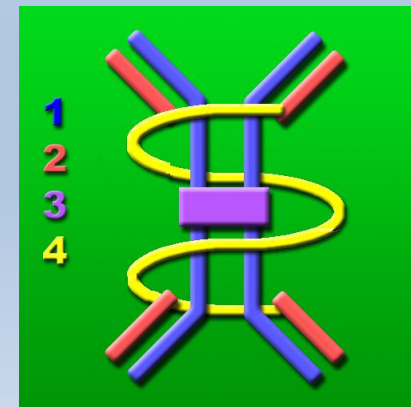
- Variable domains of L and H chain form **the binding site for Ag**
- **Hinge region** – place where are the heavy chains linked by disulfide bonds



- Immunoglobulins are glykoproteins (glycosylated **Fc part**)

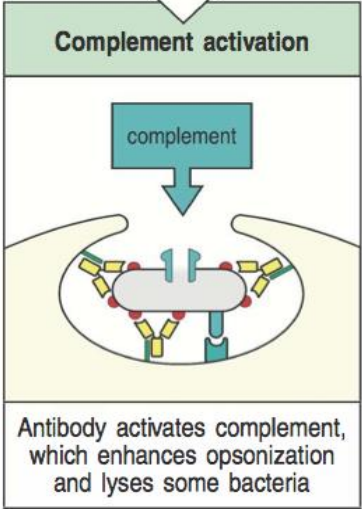
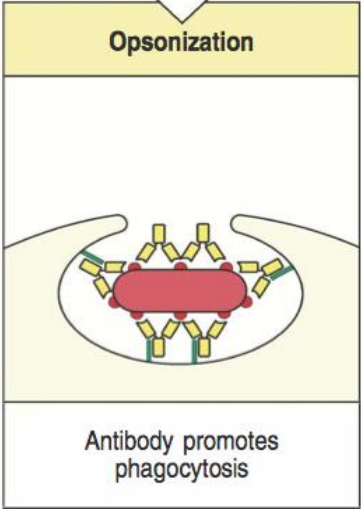
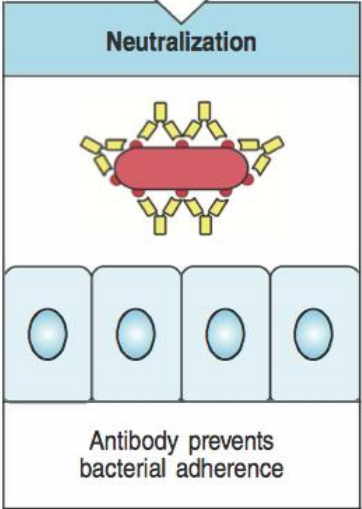
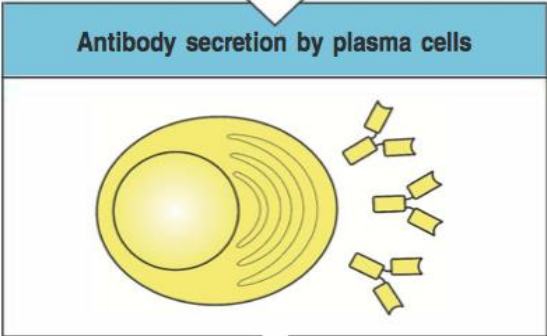
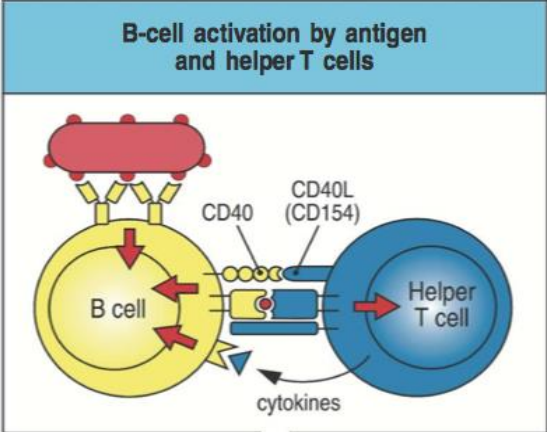
- **J chain** - molecules of immunoglobulin classes (IgM, IgA) consist of several monomer units – joined together by J chain

- **Secretory component** (IgA)



# Immunoglobulins functions

- **Antigen neutralization** Antibodies prevent bacterial adherence or inhibit activity of toxins, viruses and other microorganisms by binding to their important epitopes
- **Complement activation (IgM, IgG)** Antibody activates complement, which enhances opsonization and lyses some bacteria
- **Opsonization (IgA, IgG)** Antibodies promote phagocytosis by APC
- **Mast cell activation using IgE**
- **ADCC** (antibody-dependent cellular cytotoxicity)



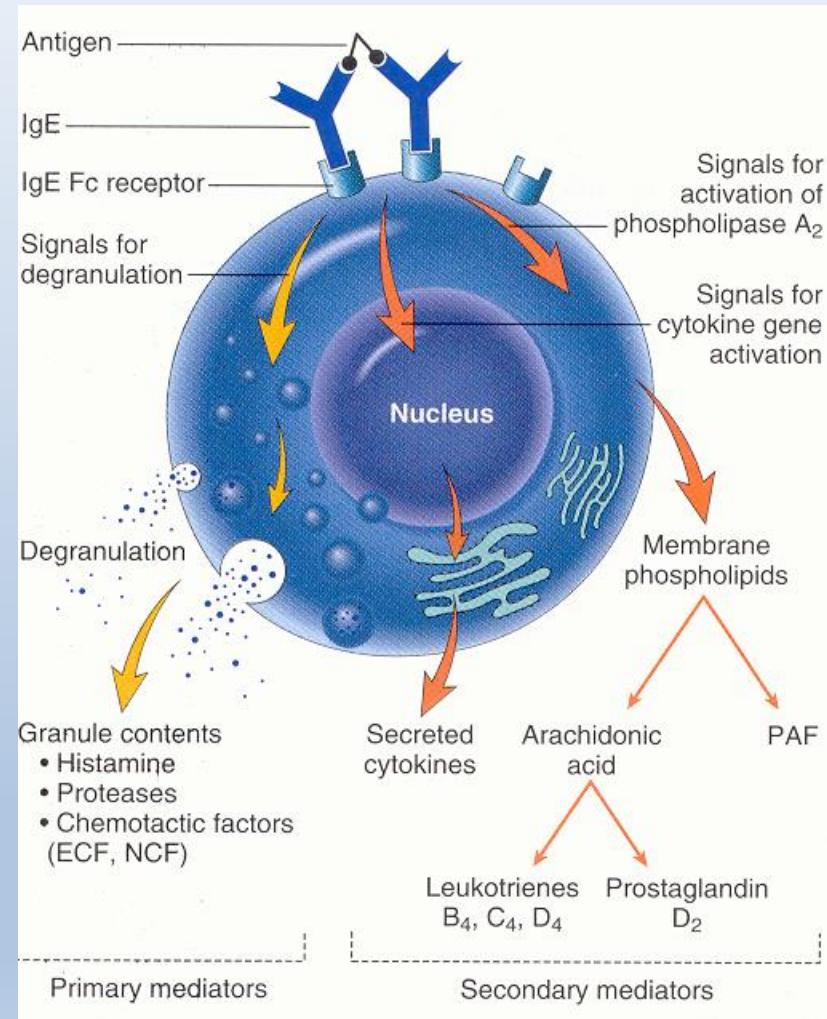


# Immunoglobulins functions

## ■ Mast cell activation using IgE

Mast cells can be stimulated by cross-linking of IgE receptors (Fce RI)

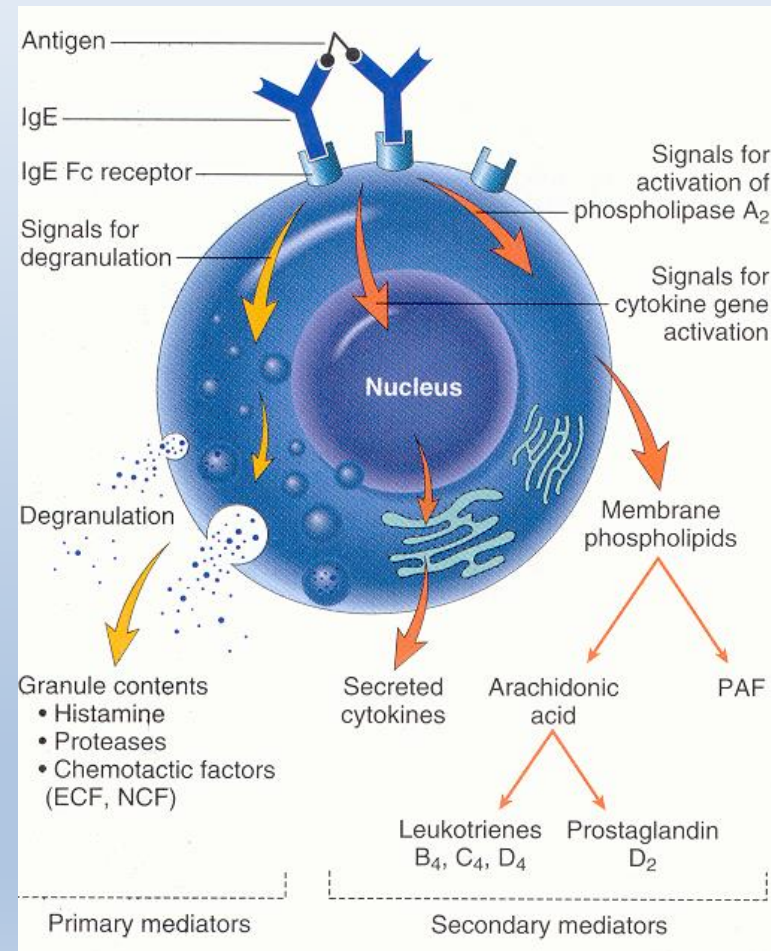
Basis of allergic reaction and of the defence against multicellular parasites



# Mast cell activation by cross-linking of IgE Fc receptors

Allergen or multicellular parasite binds to IgE on mast cell → cross-linking of several molecules Fc $\epsilon$ RI

- initiate mast cell **degranulation** (release of **histamin**, tryptase, serotonin...)
- **activation of arachidonic acid metabolism** (leukotriene C<sub>4</sub>, prostaglandin PGD<sub>2</sub>) - amplification of inflammatory responses
- **cytokine production** by mast cell (TNF, TGF $\beta$ , IL-4, 5, 6)

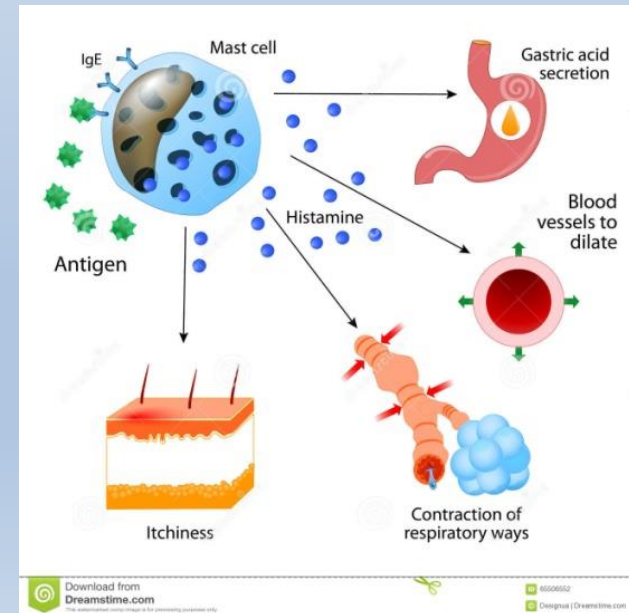


# Histamine

- vasodilatation, increase of vascular permeability (erythema, edema, itching)
- bronchoconstriction (cough, wheezing, dyspoe)
- increases intestinal peristalsis (diarrhea)
- increases mucus secretion (cough)

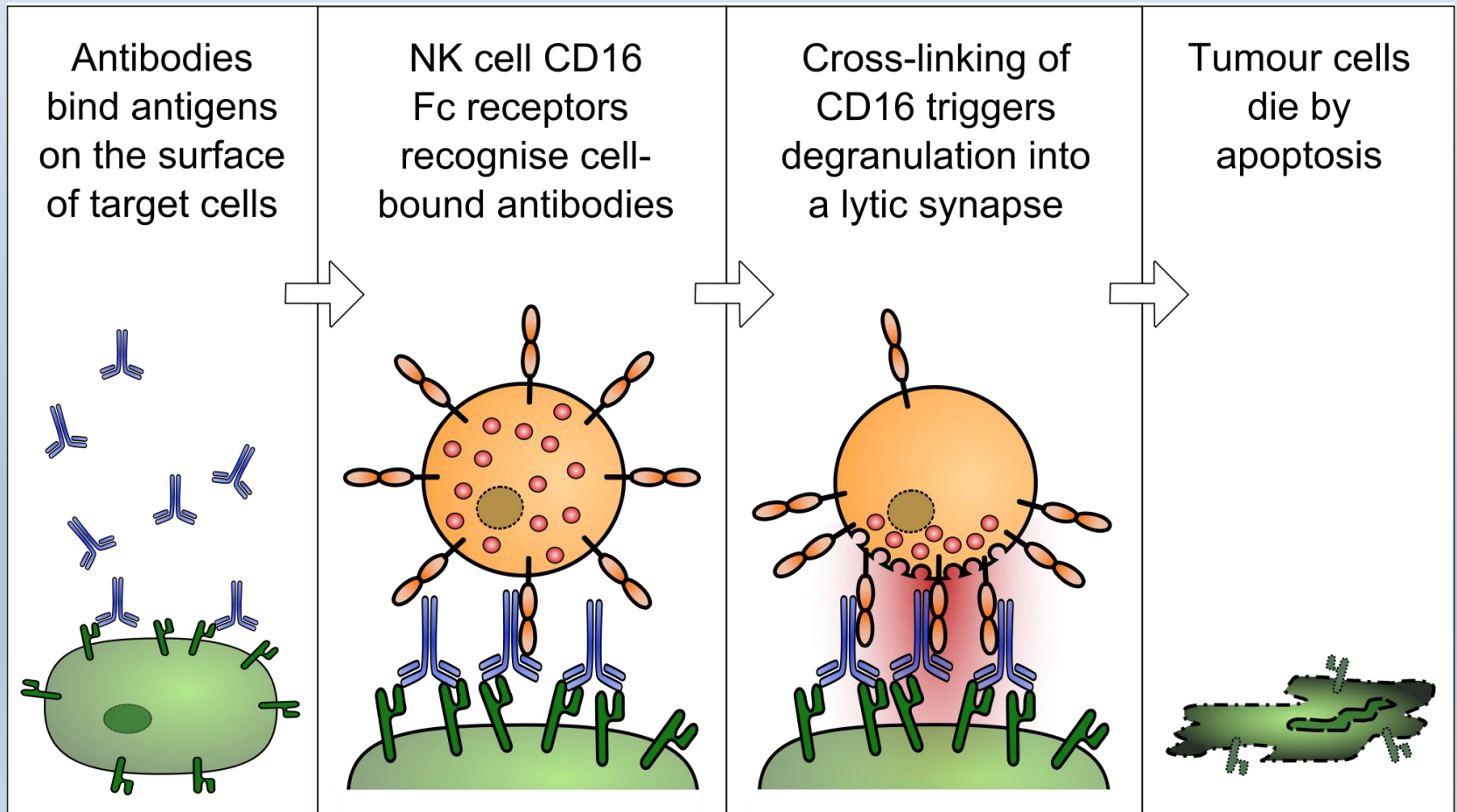
Responsible for the clinical signs of allergy.

Helps eliminate the parasite.



# Immunoglobulins functions

- **ADCC** (antibody-dependent cellular cytotoxicity) NK cells recognize cell opsonized with IgG antibodies by the Fc receptor CD16, this leads to the activation of cytotoxic mechanisms (NK degranulation)



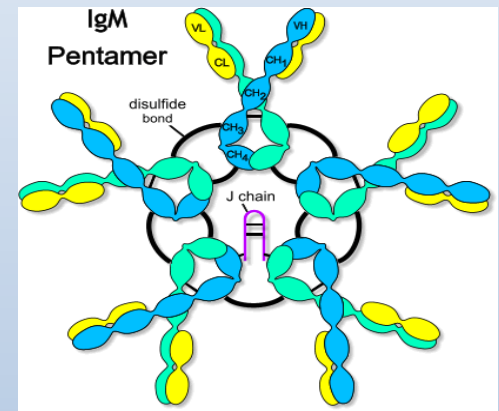
- <https://www.youtube.com/watch?v=N3L4kQqsGPO>

# Classes of immunoglobulins and their functions

- IgM, IgD, IgG (IgG1 - IgG4), IgA (IgA1, IgA2), IgE
- Distinguished by the constant part of H chain

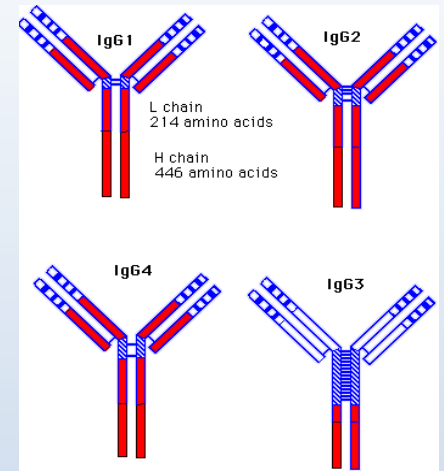
## ■ IgM

- first isotype that forms after the meeting with Ag
- as a monomer form BCR
- secreted as pentamer (10 binding sites)
- functions: Ag neutralization, complement activation, do not bind to Fc receptors on phagocytes
- (concentration of 0.9 to 2.5 g / l; biol. half-life 6 days)



## ■ IgG

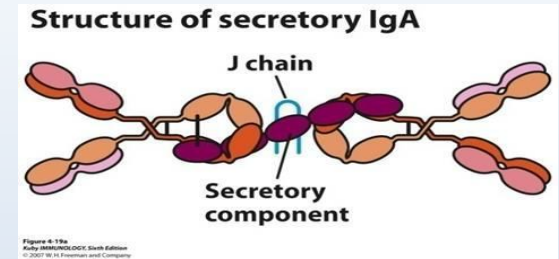
- predominantly formed in secondary immune response
- functions: Ag neutralization, opsonization, complement activation, ADCC
- isotypes IgG1-IgG4 with different ability of complement activation and binding to Fc receptors on phagocytes (opsonization)
- passes the placenta (protection of fetus in utero)
- (concentration of 8 to 18 g / l; biol. half-life of 21 days)



# IgA

## Secretory IgA

- the most significant mucosal immunoglobulin
- dimer with secretory component
  - transcytosis - IgA is transported across the epithelium using transport Fc receptor (polymeric-Ig receptor), on luminal side is IgA split off with the part of the receptor called secretory component, which protects Ig against intestinal proteases
- provides protection of mucous membranes
- functions: Ag neutralization, do not activate complement , opsonization
- saliva, tears, breast milk



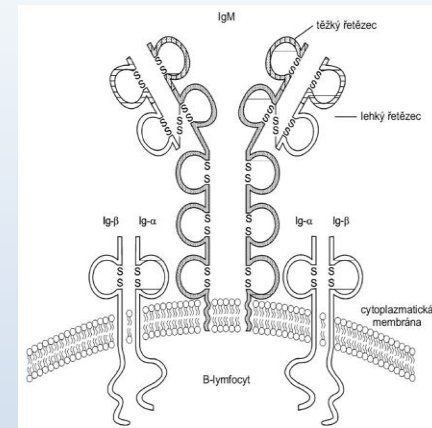
## Serum IgA

- monomer, dimer or trimer
- (0.9 to 3.5 g / l; biol. half-life of 6 days)



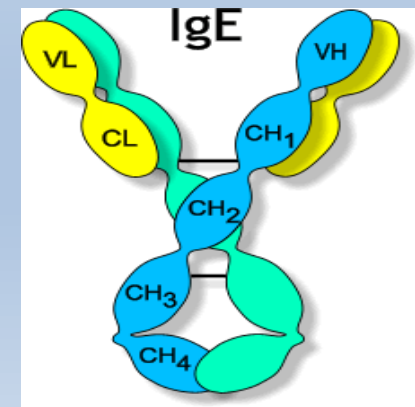
## ■ IgD

- form a BCR
- in serum is in a very low concentration
- (0.1 g / l; biol. half-life 3 days)



## ■ IgE

- applies in defense against multicellular parasites
- is the main cause of allergic reactions
- ( 0-100 kIU/l; biol. half-life 2 days)



# The genetic basis of the immunoglobulins development

- **Gene segments for H chains** – on chromosome 14

**V** (variable) segments

**D** (Diversity) segments

**J** (joining) segments

**C** segments



variable domain of H chain

constant domains of H chain

- **Gene segments for L chains** -  $\kappa$  on chromosome 2  
-  $\lambda$  on chromosome 22

**V** (variable)

**J** (joining)

**C**

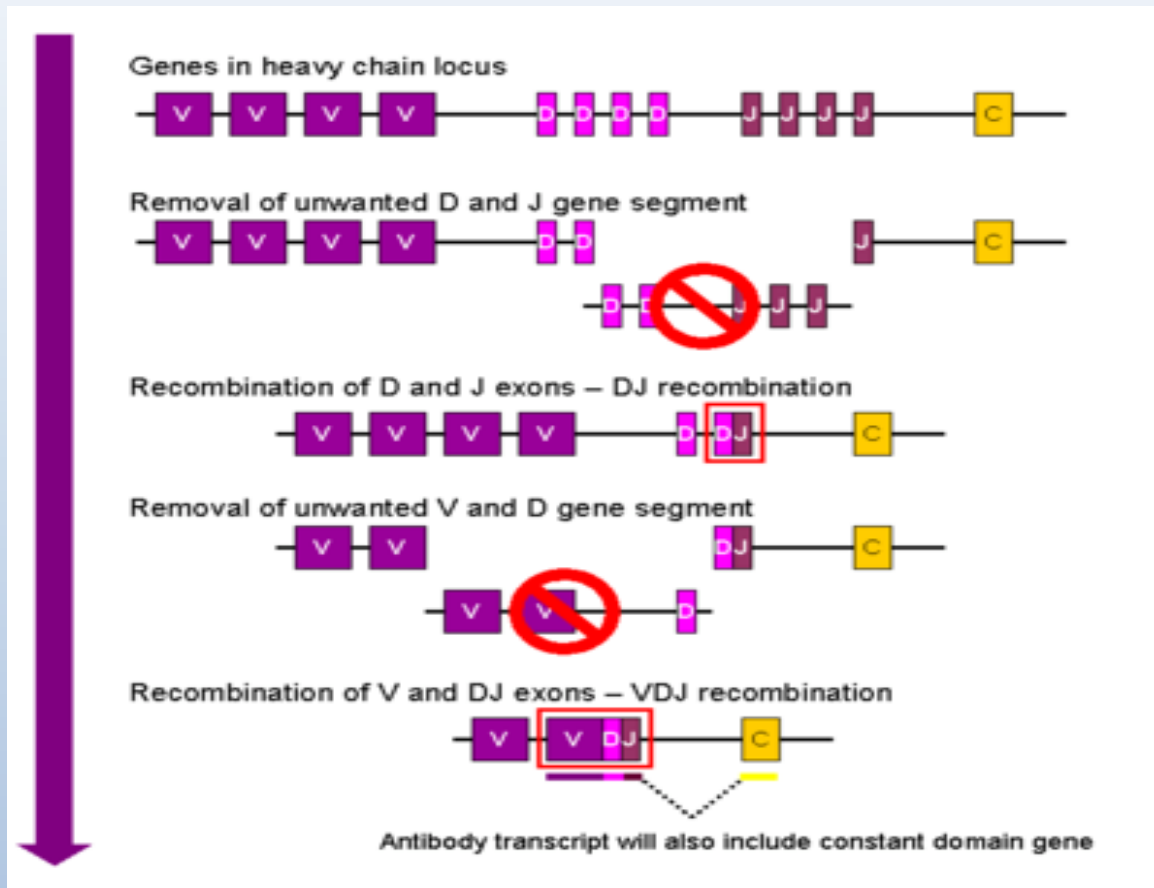


variable domain of L chain

constant domain of L chain

- At the ends of V, D, J segments there are signal sequences which are recognized by enzyme VDJ recombinase that carry out the rearrangement of these genes
- On the sides of C segments are so-called switch sequences, which are recognized by enzyme recombinase that carry out isotype switching

# The rearrangement of genes coding H chain

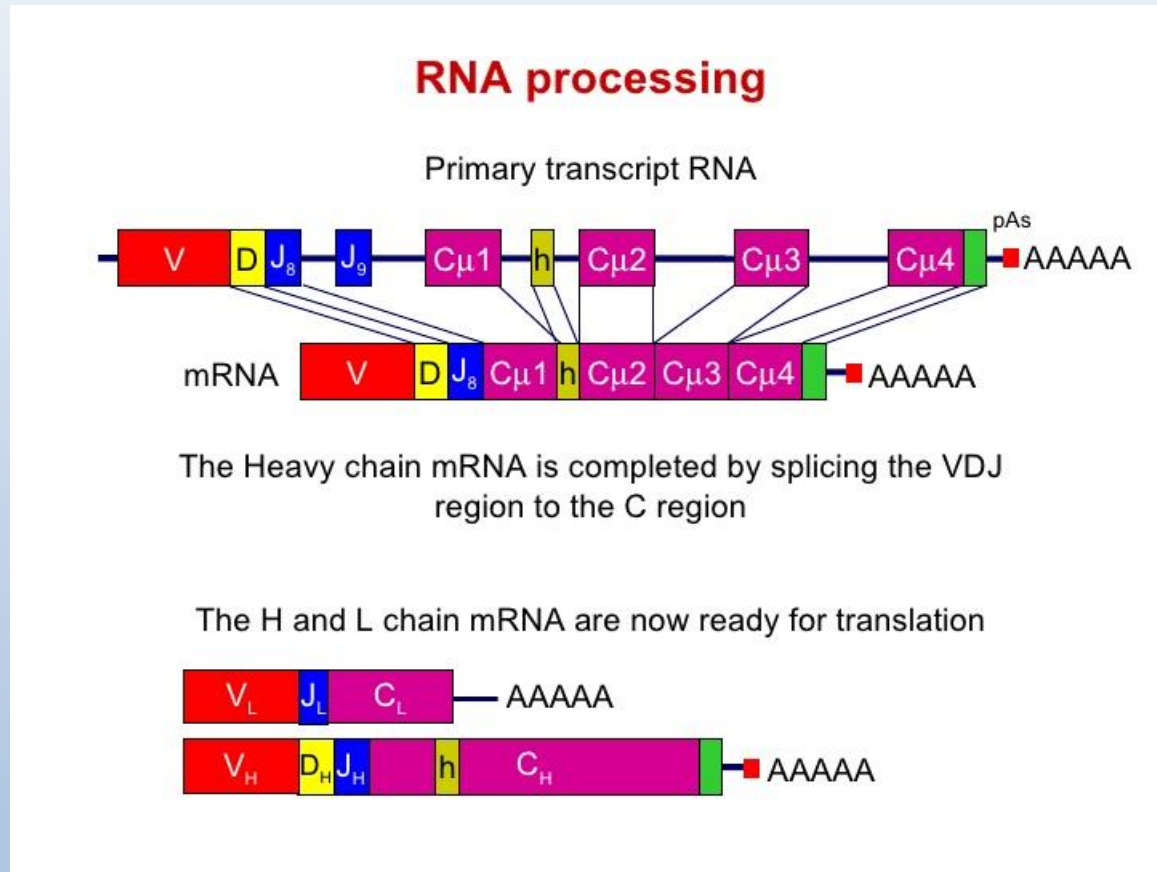


**1) DJ rearrangement** - excision a section of gene complex between some D and J segment

**2) VD rearrangement** - excision a section between some V segment and DJ

The rearranged IgH gene is transcribed into mRNA

# The rearrangement of genes coding H chain



The first formed H chain is  $\mu$ .

If rearrangement is unsuccessful, B lymphocyte die.

# The rearrangement of genes coding L chain

- 1) First, rearrange the genes encoding the L chain  $\kappa$ , there is excision of sections between a V and J segment
- 2) If regrouping of the  $\kappa$  genes is unsuccessful, start the regrouping genes  $\lambda$ .

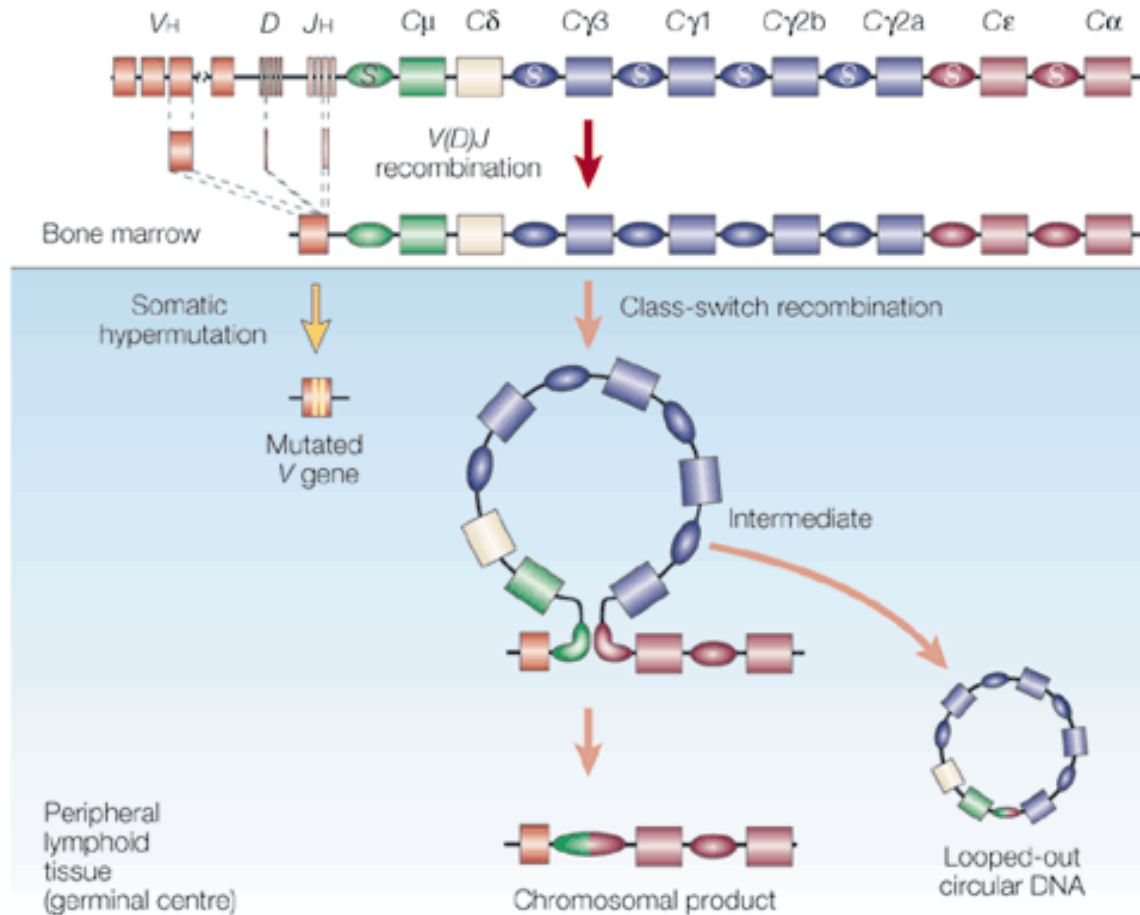
If regrouping is unsuccessful, B lymphocyte die.

- [https://www.youtube.com/watch?v=wm4qqkf\\_iY](https://www.youtube.com/watch?v=wm4qqkf_iY)

# Isotype (class) switching

- Occurs during the terminal differentiation of B lymphocyte after contact with Ag on the surface of FDC
- Enzymes recombinases recognize the switch sequences located on the sides of C segments and excise gene segments
- After elimination of some C segment, the closest segment to VDJ segment is transcribed into mRNA, and after splicing and translation arise corresponding isotype of the H chain

# Isotype switching





# Isotype switching

- Cytokines regulate which isotype will be produced:

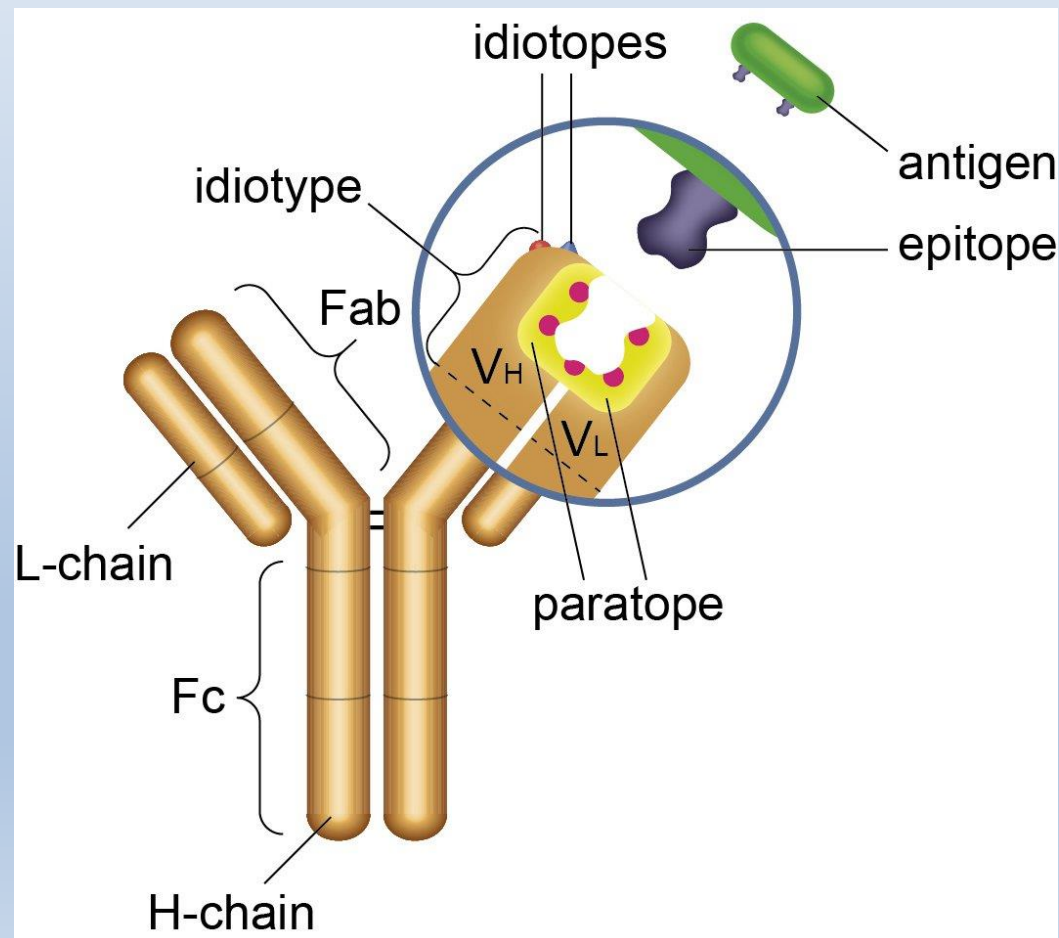
IL-4 stimulates switching to IgE and IgG4

TGF $\beta$  stimulates switching to IgG2 and IgA

<https://www.youtube.com/watch?v=gyTHXjVUPWw>

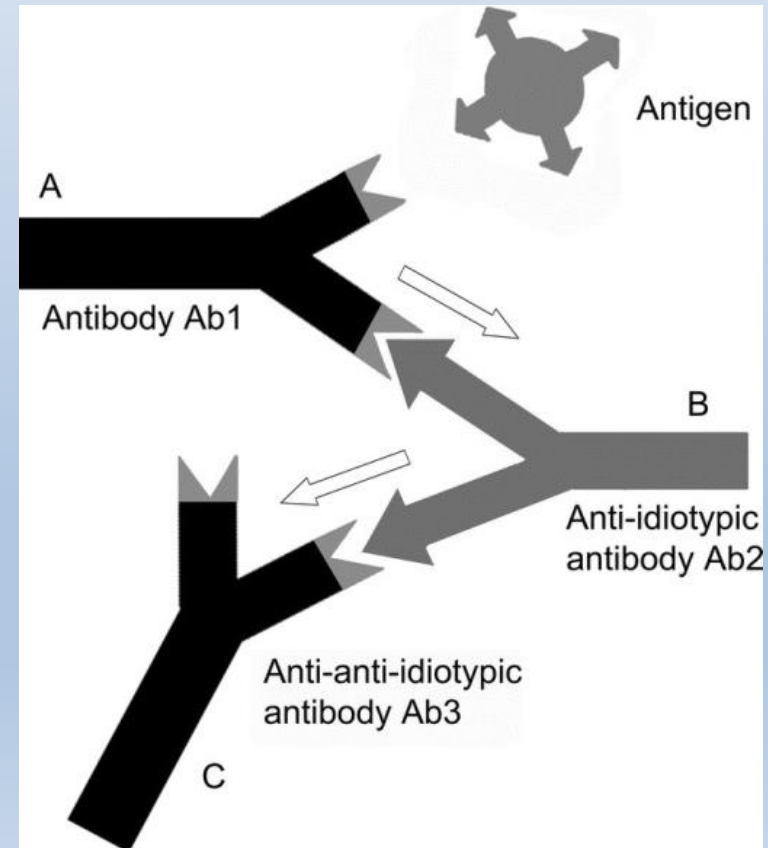
# Anti-idiotypic antibodies

- **IDIOTYP** = group of idiotops which are located on the variable part of antibody



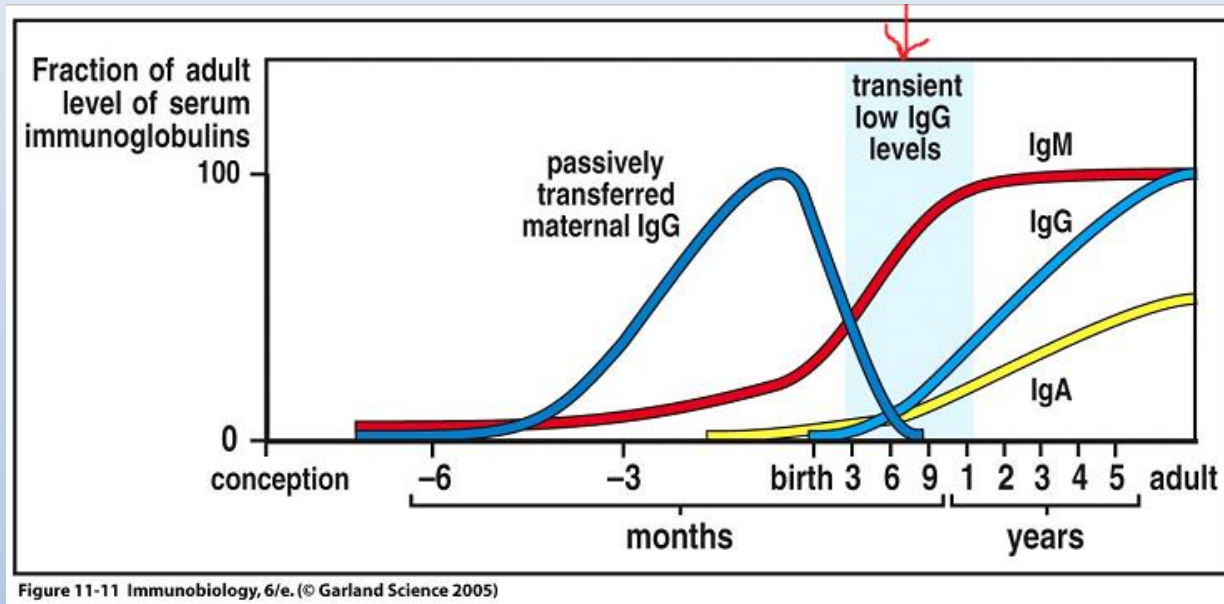
# Anti-idiotypic antibodies

- Idiotypic structures of 1st generation antibodies can be recognized by some B cells as antigens and can induce production of anti-idiotypic antibodies (2nd generation antibodies; some binding sites may remind Ag, which caused formation of 1st generation antibodies)
- Against the 2nd generation antibodies formate antibodies of 3rd generation (anti-antiidiotypic antibodies).
- The idiotypic network may play a role in regulation of antibody response



# Ontogenesis of antibodies

- Synthesis of specific antibodies begins around the 20.-24. week of gestation, the total concentration of IgA and IgM remains very low until birth, IgG begins to form after birth

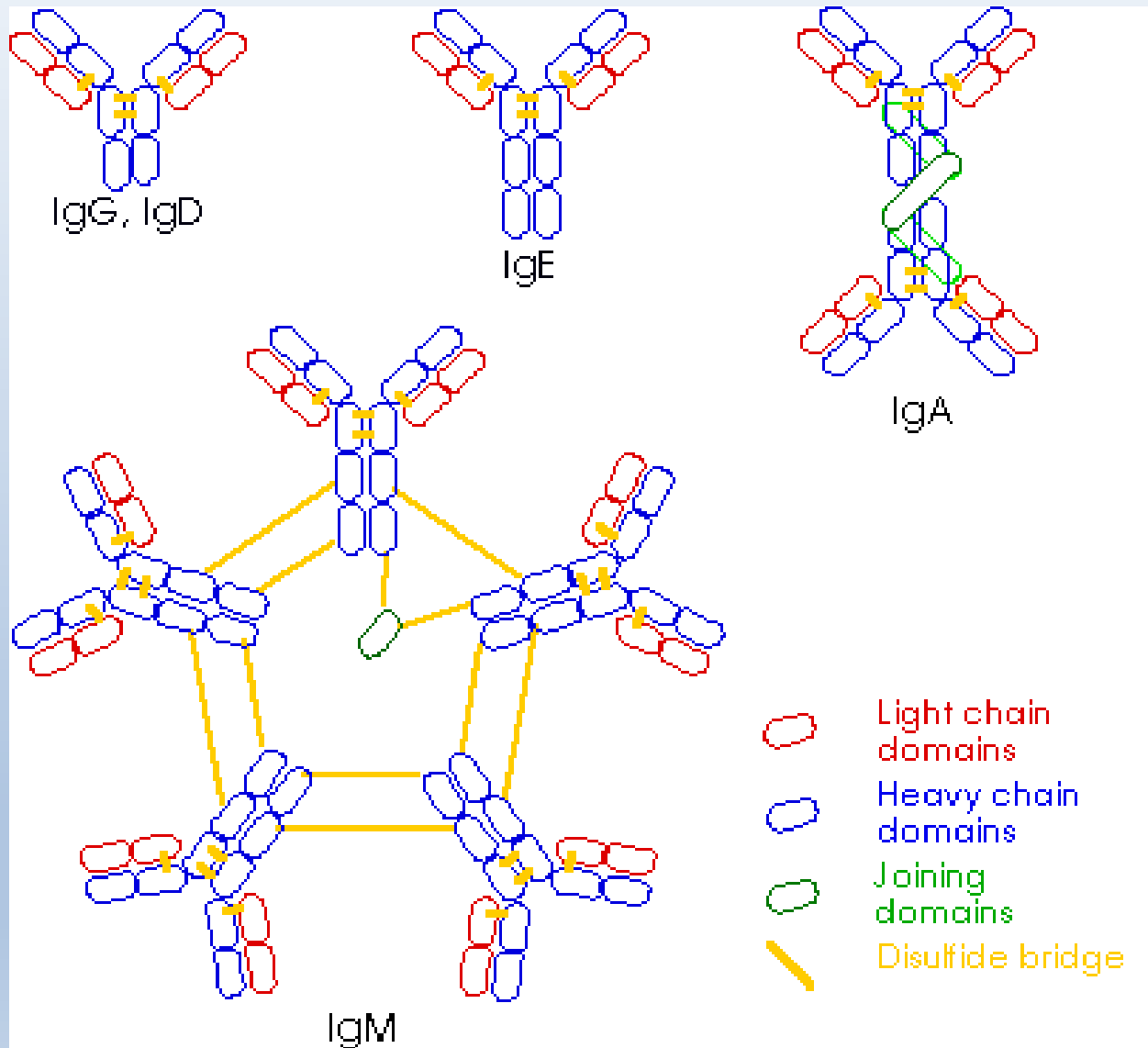


After birth begins slow growth of own IgG, which is accompanied by decline in maternal IgG (about 3. to 6.month)

- The IgM concentration reaches values comparable with adults in the 1- 3 year of life, IgG and IgA between 10.-15. year

- After birth B lymphocytes respond to immunization predominantly by IgM formation, switching to other isotype is slower
- Antibody response to polysaccharide antigens appears around 2. year of life
- In old age is a lower antibody response to new stimuli and increased autoantibodies production

# Immunoglobulins summary



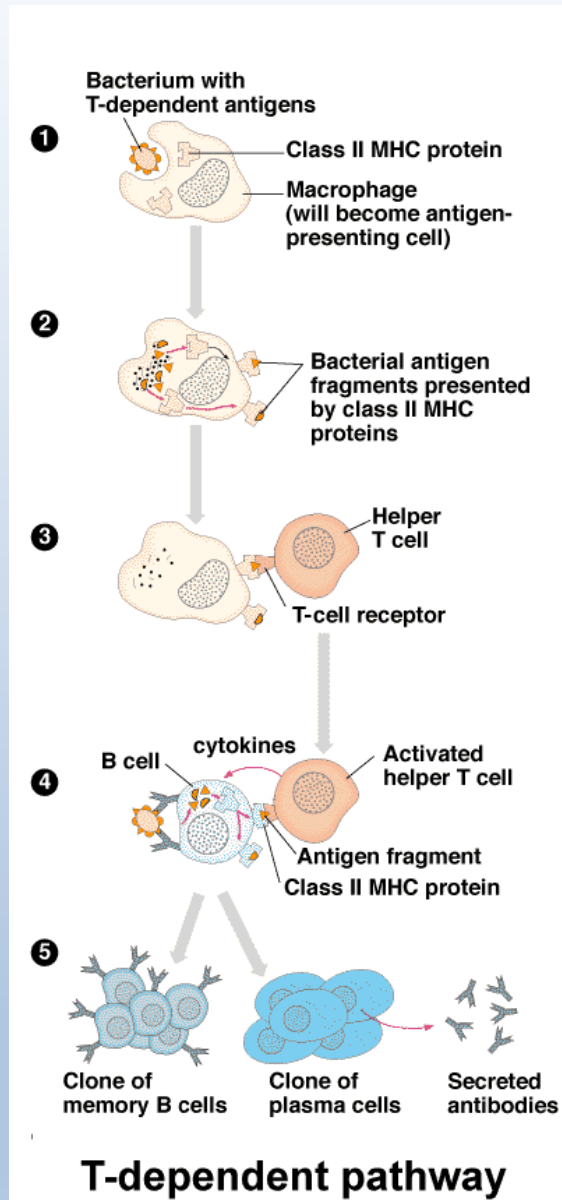
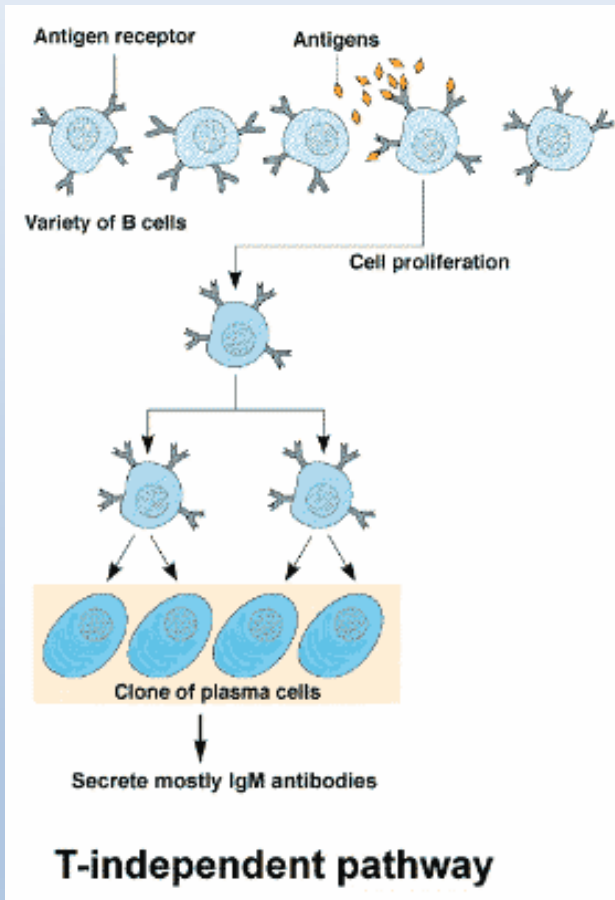
# **Humoral immune response**



# Humoral response induced by

- **T-independent antigens**
  - Cause predominantly IgM production
  - Bacterial polysaccharides, lipopolysaccharides
- **T-dependent antigens**
  - Reaction to these Ag occurs in two phases - primary and secondary
  - Initiate the development of memory cells and formation of high-affinity antibodies and different isotypes
  - Most of antigens (proteins)

# T-independent and T-dependent immune response



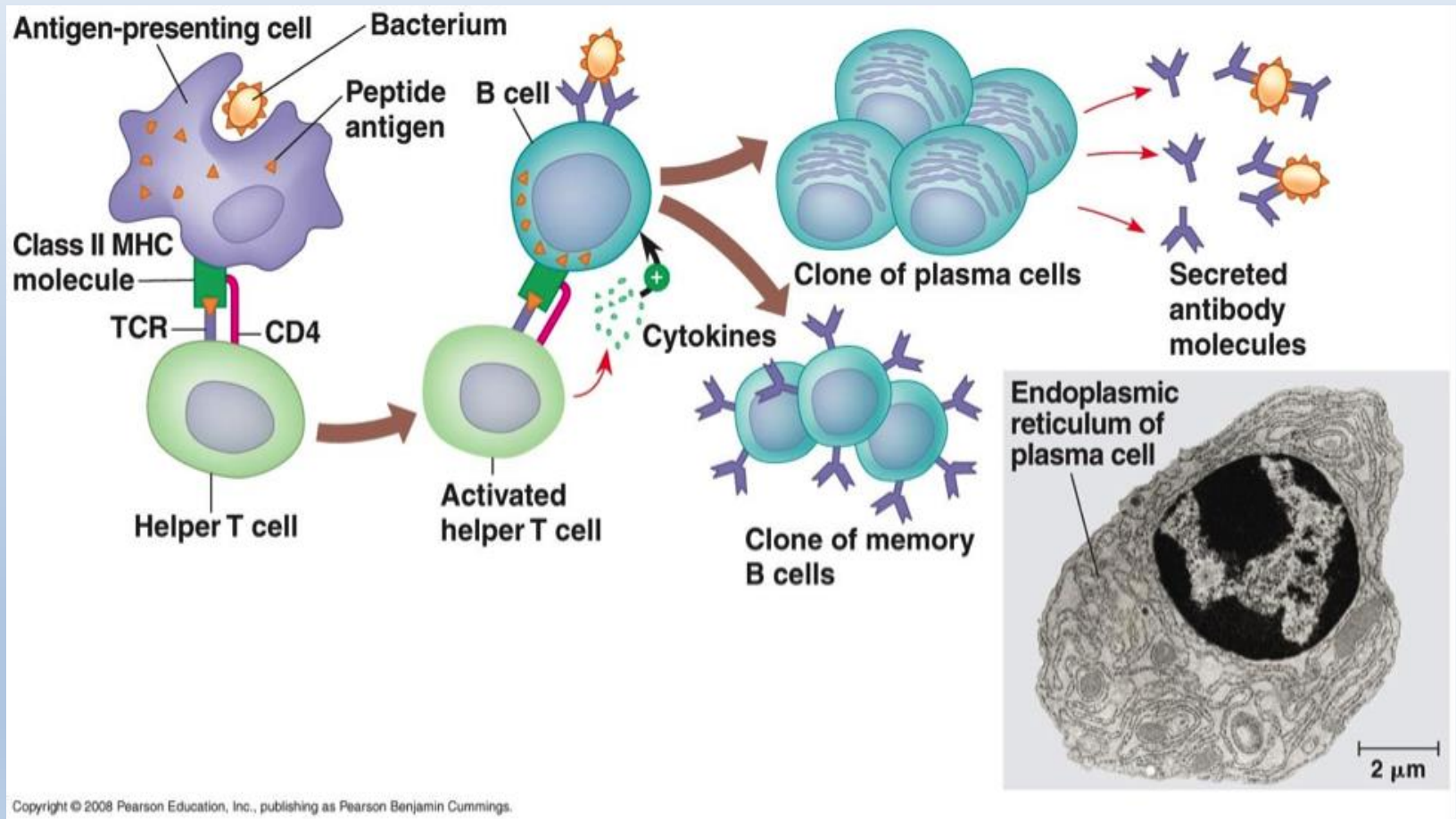
# Antibody response induced by T-dependent antigen

## Primary phase of antibody response

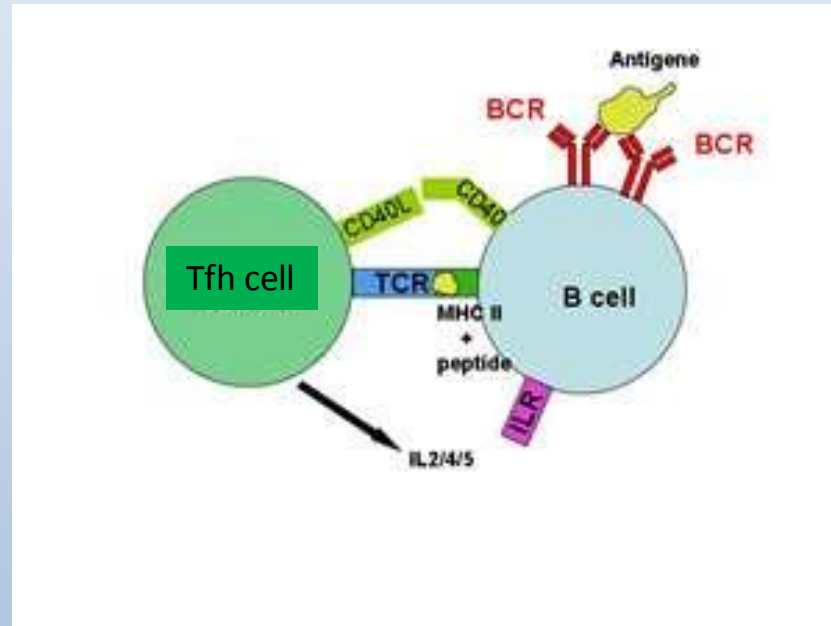
- Takes place in secondary lymphoid organs
- After the first contact with Ag 2 processes run simultaneously:
  - **stimulation of B** cells by Ag binding to BCR
  - Ag **absorption by APC** and its **presentation** via MHC gp class II to precursors of T<sub>H</sub> cell → formation of clone of antigen-specific Tfh cells

Tfh cells provide assistance to stimulated B cells - leading to their proliferation, differentiation into plasma (produce Ab) and memory cells

# T-dependent immune response



# Antigen presentation by B lymphocyte to Tfh lymphocyte



- Antibodies produced in the primary phase (3-4 days) are **IgM** and have a low affinity for Ag, create with Ag immune complexes
- Immune complexes are captured in the secondary lymphoid organs on the surface of **FDC** (follicular dendritic cells) - Ag presenting cells to B lymphocytes

## Secondary phase of antibody response

- When antigens in immune complexes on the surface of FDC are recognized by B cells, another cycle of proliferation and differentiation of B cells begins (with assistance of Tfh cells)
- This process is accompanied by somatic mutations of V segments of H and L chains → production of antibodies with higher affinity to Ag (4-6x higher) = **affinity maturation of antibodies**
- Takes place in germinal centres (contain B, Thf and FDC) of newly formed secondary lymphoid follicles = **Germinal center reaction**

# Secondary phase of antibody response

- Besides somatic mutations also **isotype switching** starts- instead of IgM other isotypes of immunoglobulins are produced, which isotypes (IgG, A, E) arise determines cytokine environment
- Contact between CD40 (B lymphocytes) and CD40L (Tfh lymphocytes) is essential for the initiation of somatic mutations, isotype switching and formation of memory cells



## Secondary phase of antibody response

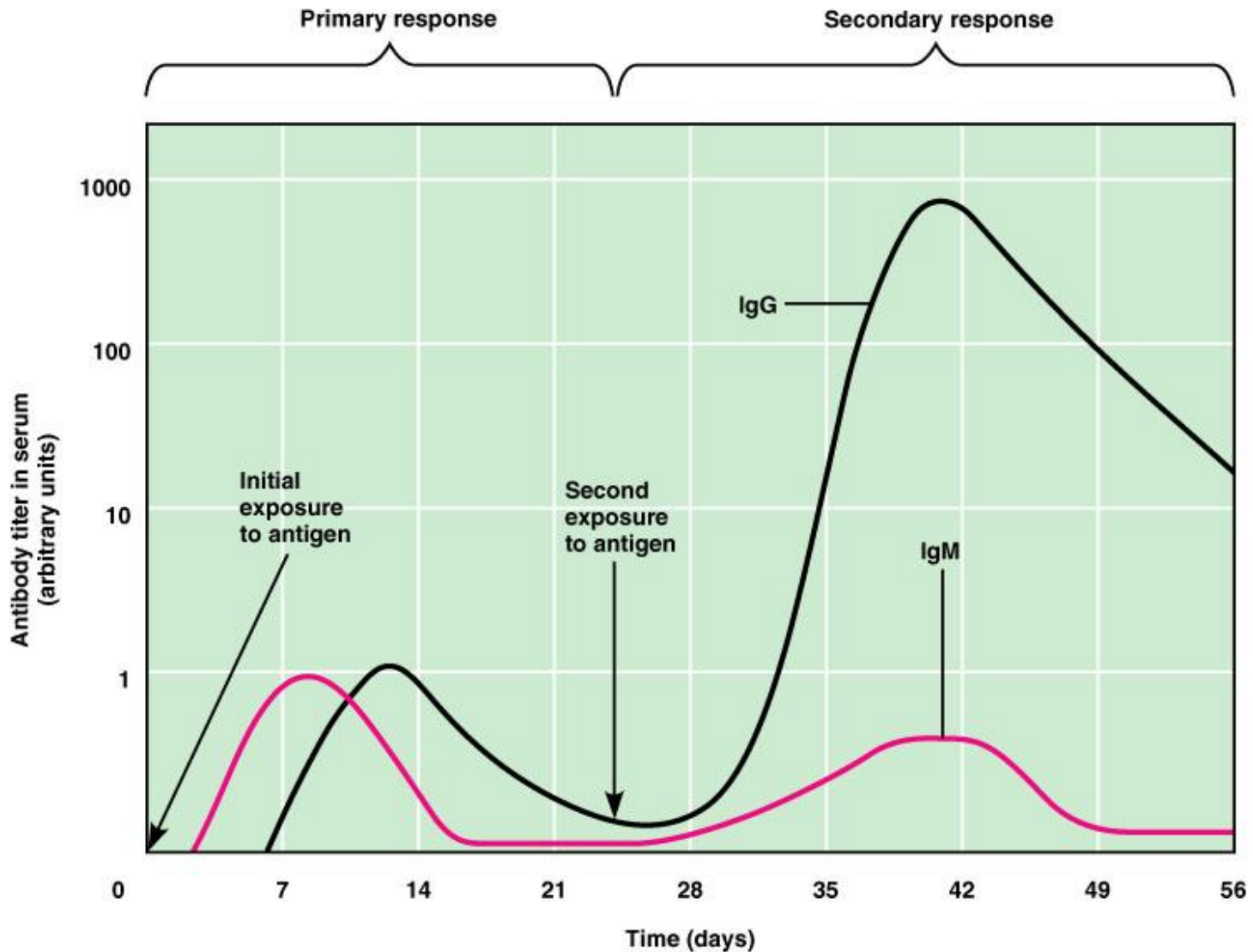
- In the secondary phase of the immune response there are generated antibodies with **higher affinity to Ag** and with **other effector characteristics** , which are dependent on isotype. During this phase also **memory cells** are formed, prepared for next meeting with the Ag
- Antibodies in the body persist for a long time after primary infection

# Primary and secondary immune response

- **Primary immune response** – occurs after the first exposure to antigen
- **Secondary immune response** – occurs after subsequent encounter with the same antigen and is more rapid due to the activation of previously generated memory cells

# Primary and Secondary Response

- Antibody response to initial antigenic stimulus is called **primary response**
  - differs both quantitatively and qualitatively
  - Slow, sluggish and short lived
  - Long lag phase and low titre of antibody
  - Predominantly IgM
- Subsequent to primary response is call **secondary response**
  - Prompt, powerful and prolonged
  - Short or negligible lag phase
  - much higher level of antibodies for longer period
  - Predominantly IgG



- [https://www.youtube.com/watch?v=LYZzvrg\\_qIM](https://www.youtube.com/watch?v=LYZzvrg_qIM)

**Thank you for your attention!**