

# **Basic Immunology (Dentistry)**

## ***Lecture 7.-8.***

***Antigen recognition molecules: Immunoglobulins, T cell receptor. MHC and antigen presentation***

***Ferenc Boldizsar MD, PhD***

# Immune system

RECOGNITION

SELF

NON-SELF

normal immune-homeostasis

TOLERANCE

ELIMINATION



AUTOIMMUNITY

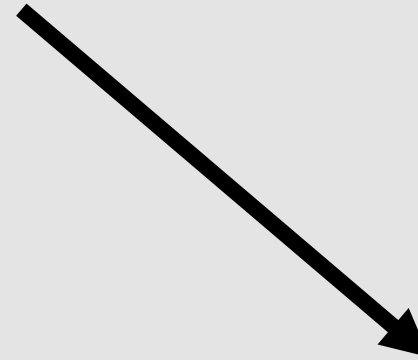
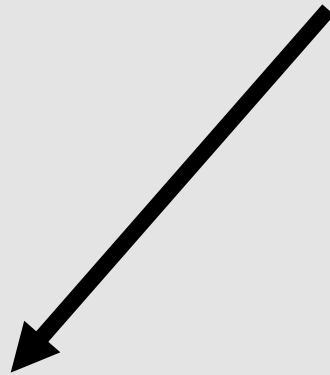
TUMORS

IMMUNO  
DEFICIENCIES

HYPERSENSITIVE  
REACTIONS

**ALTERED immune-homeostasis= IMMUNOPATHOLOGY**

# Immunological Recognition (**Receptors**)



## Innate immunity

general microbial  
Molecular PATTERNS

(„pattern recognition receptors”)

## Adaptive immunity

Antigen-specific (EPITOP)

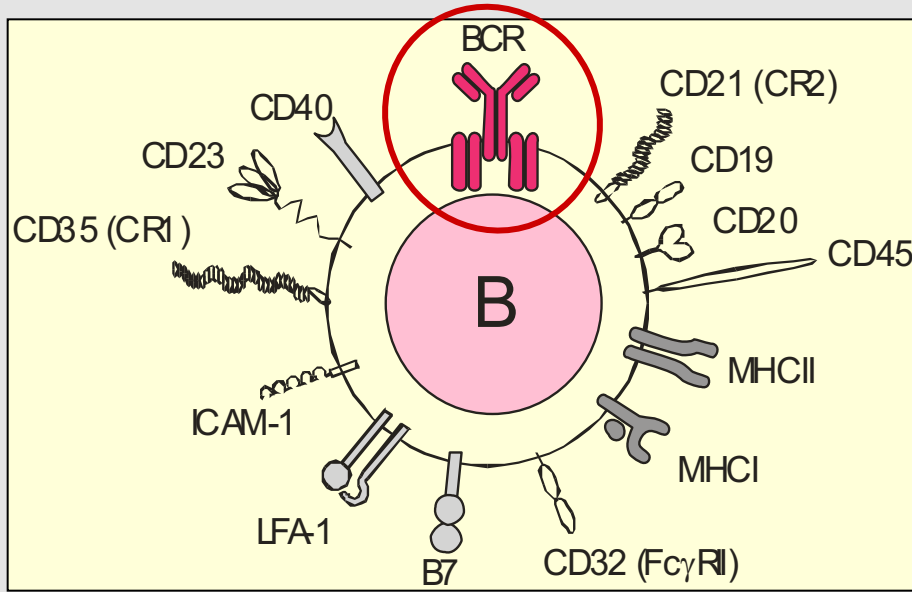


# Recognition molecules

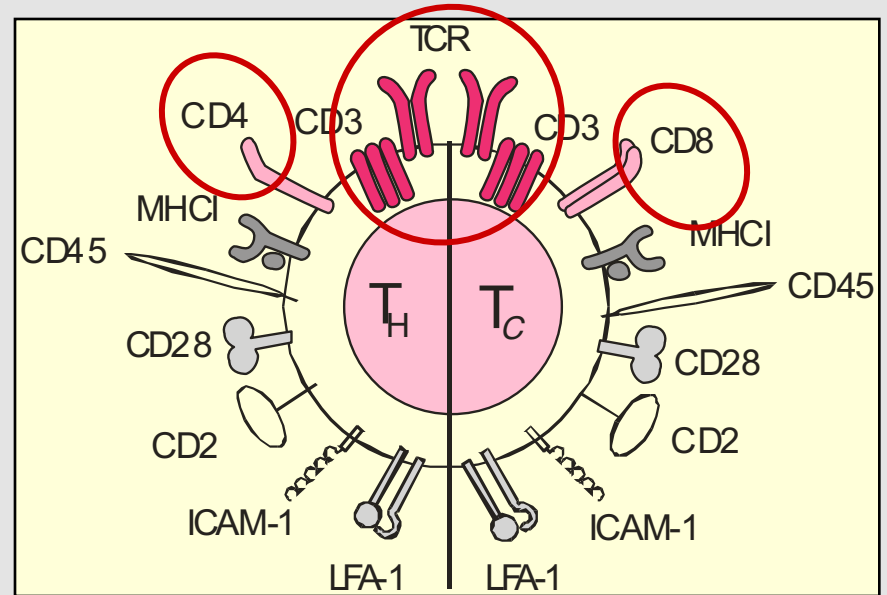
	Innate immunity	Adaptive immunity
Specificity	<p><b>For pathogen-associated molecular patterns (PAMPS)</b></p> <p>Different microbes</p> <p>Identical mannose receptors</p>	<p><b>For structural details of any molecules (antigens)</b></p> <p>Different microbes</p> <p>Distinct antibody molecules</p>
Receptors	<p><b>Encoded in germline (pattern recognition receptors)</b></p> <p>Toll-like receptor</p> <p>N-formyl methionyl receptor</p> <p>Mannose receptor</p>	<p><b>Encoded by lymphocyte genes produced by somatic recombination</b></p> <p>Ig</p> <p>TCR</p>
Distribution of receptors	<b>Non-clonal</b>	<b>Clonal</b>

Table 4-1

# Antigen-Receptors of Lymphocytes



**BcR= B-Zellrezeptor**

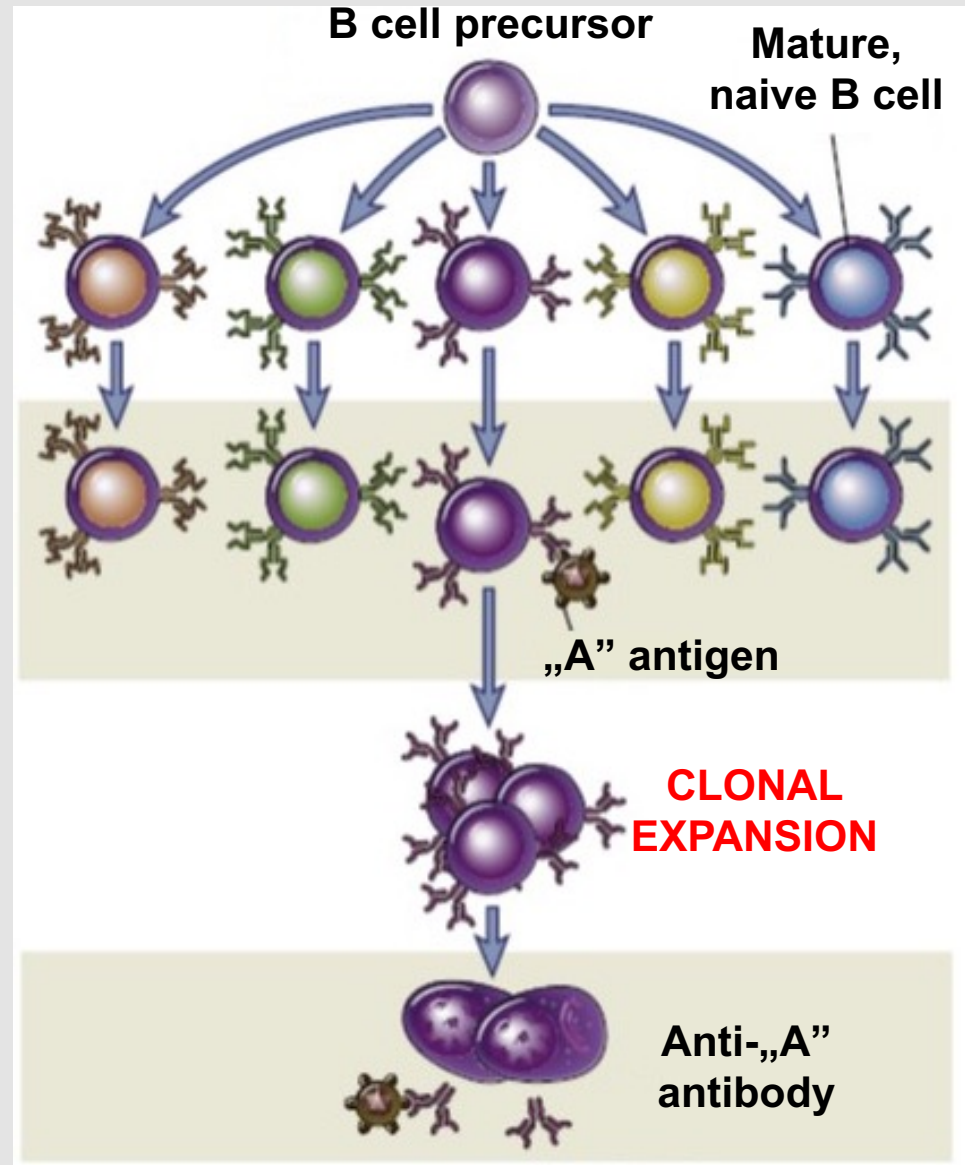


**TcR= T-Zellrezeptor**

BcR and TcR are **Antigen-Receptors**, which are different on each individual lymphocyte. Every single Antigenreceptor recognizes and binds only ONE specific Antigen (EPITOP)

# The Clonal Selection Hypothesis

1. Each newly produced lymphocyte expresses a **unique antigen-binding receptor**.
2. **Only those lymphocytes will become activated which recognize an antigen.** These selected cells will proliferate and produce **clones** of themselves with each sister cell having the same antigen-recognition receptor.
3. These clones will differentiate into **effector cells** which will participate in the immune response. (e.g. effector plasma cells produce antibodies)



# Recognition molecules in the adaptive immune system

Immunoglobulins

B cell receptors (BcR)

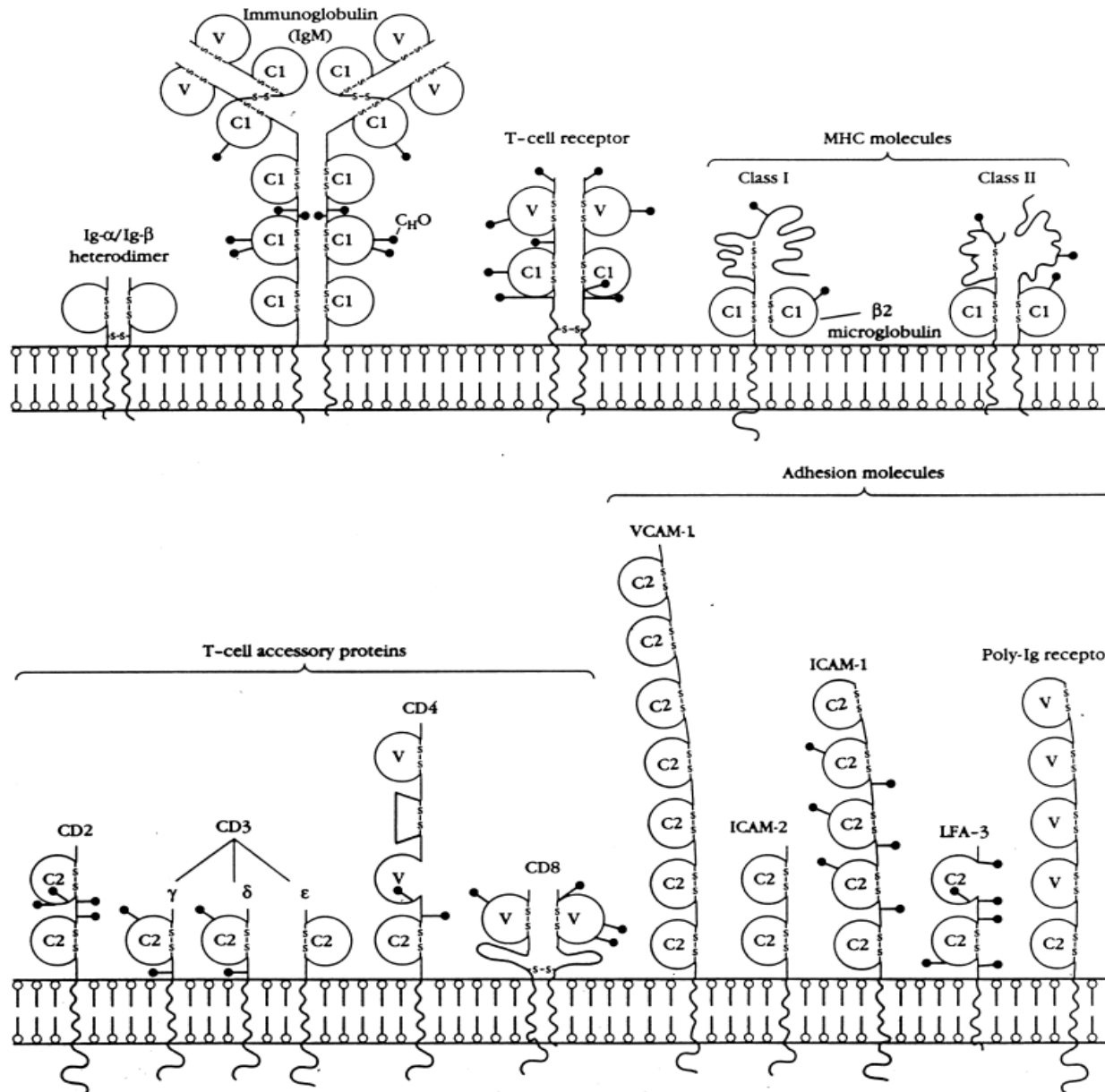
T cell receptors (TcR)

MHC class I and class II

Specialized molecules manage antigen recognition. The common structural features of these molecules are the well-conserved (constant) basic elements (designed by 110 amino acids domain units) containing variable, antigen specific parts (binding sites) for the recognition and ligand formation.

# Immune recognition molecules

**Antigen  
specific  
recognition  
molecules**



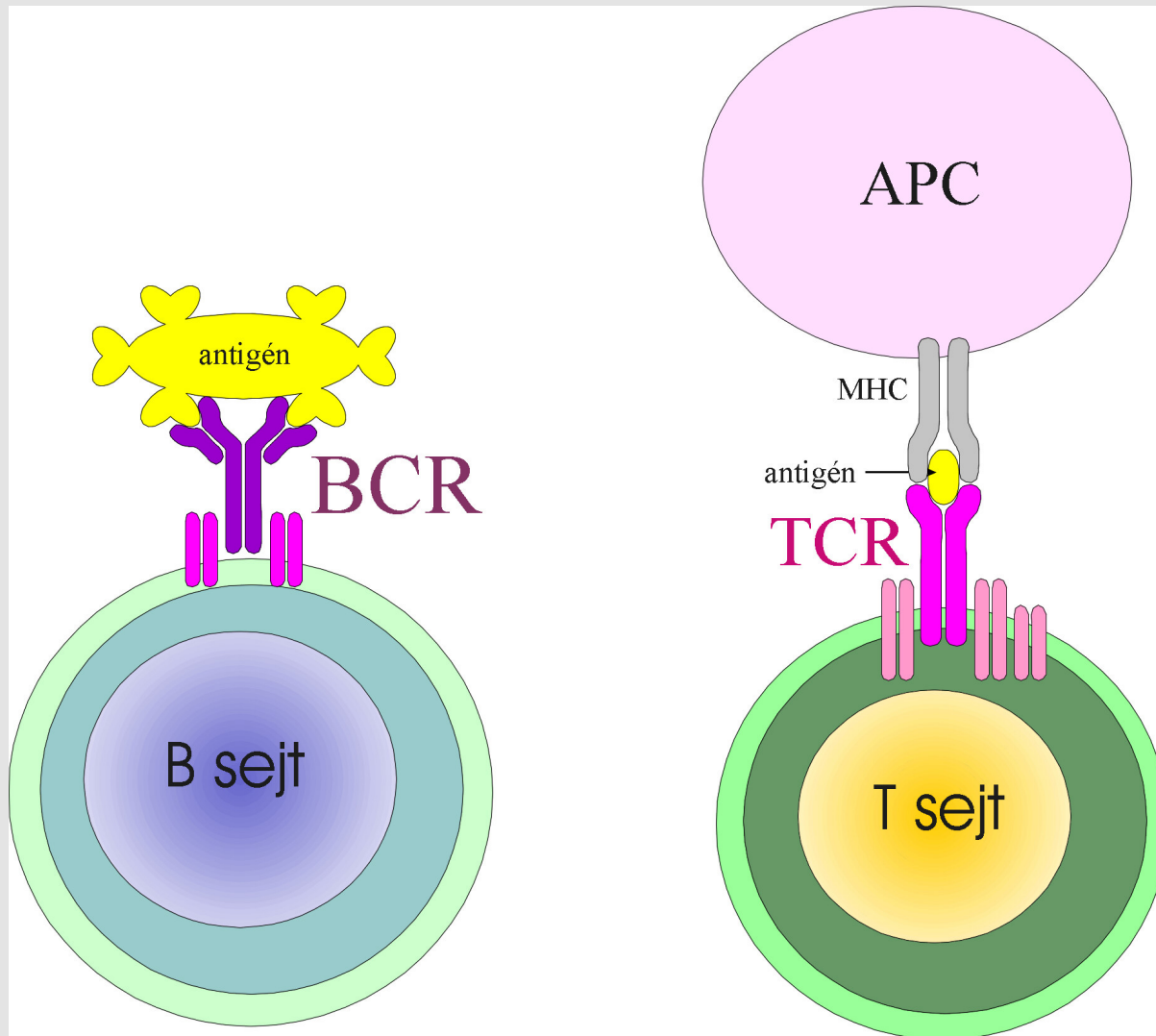
**Accessory  
molecules**



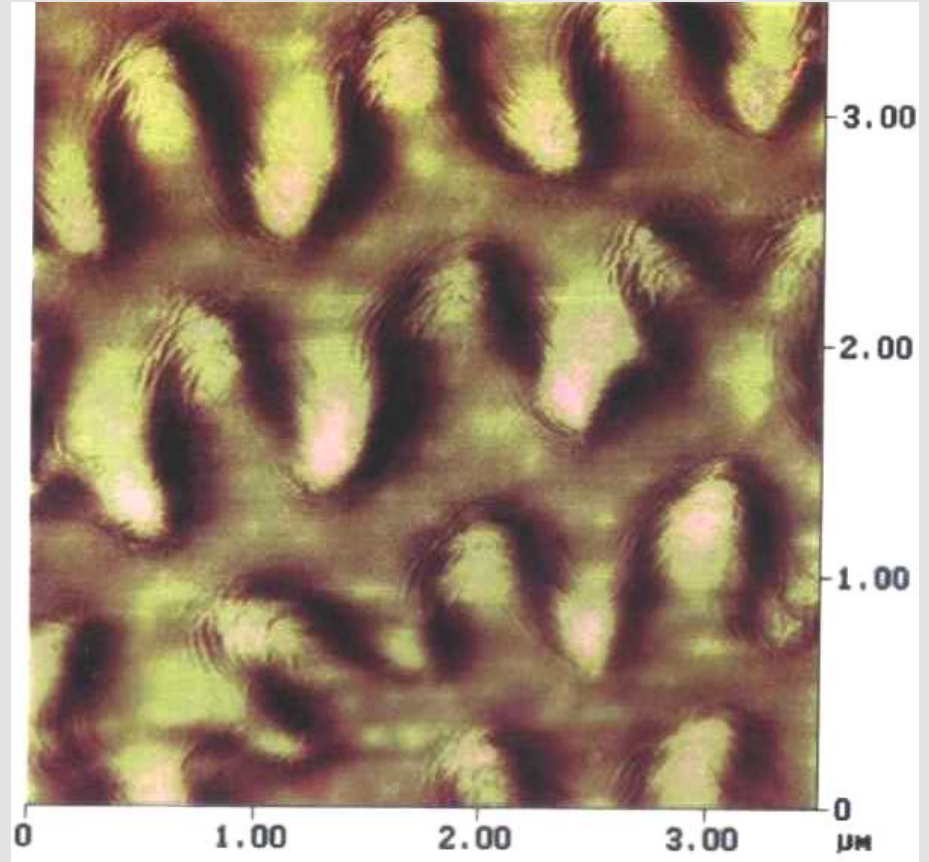
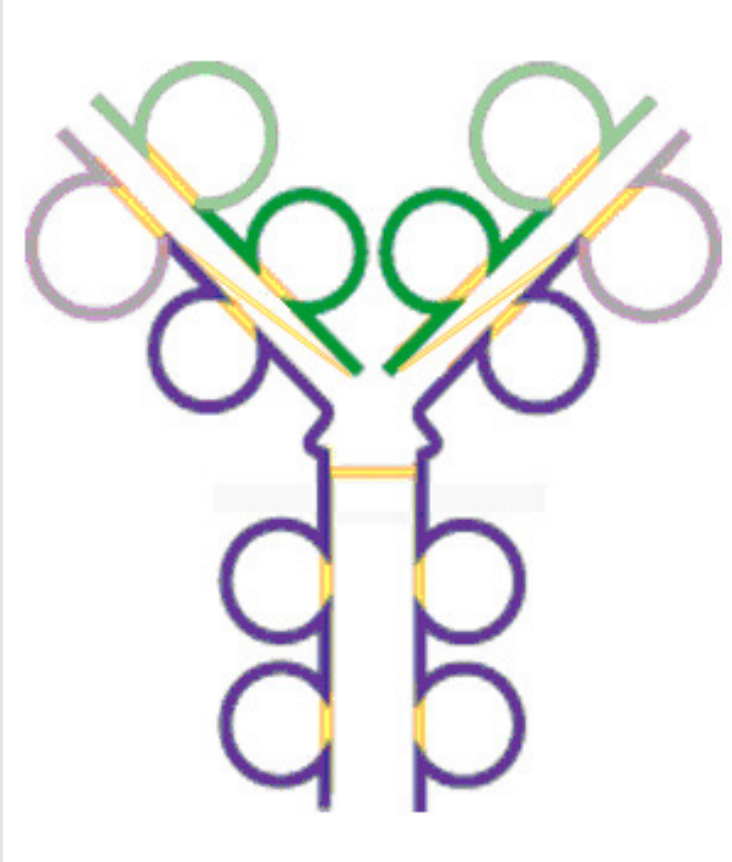
# Antigen recognition

	<b>B cells</b>	<b>T cells</b>
Receptor	BcR (Ig)	TcR
Antigen	native	denatured (presented)
APC	not needed	needed

# B cells and T cell antigen recognition

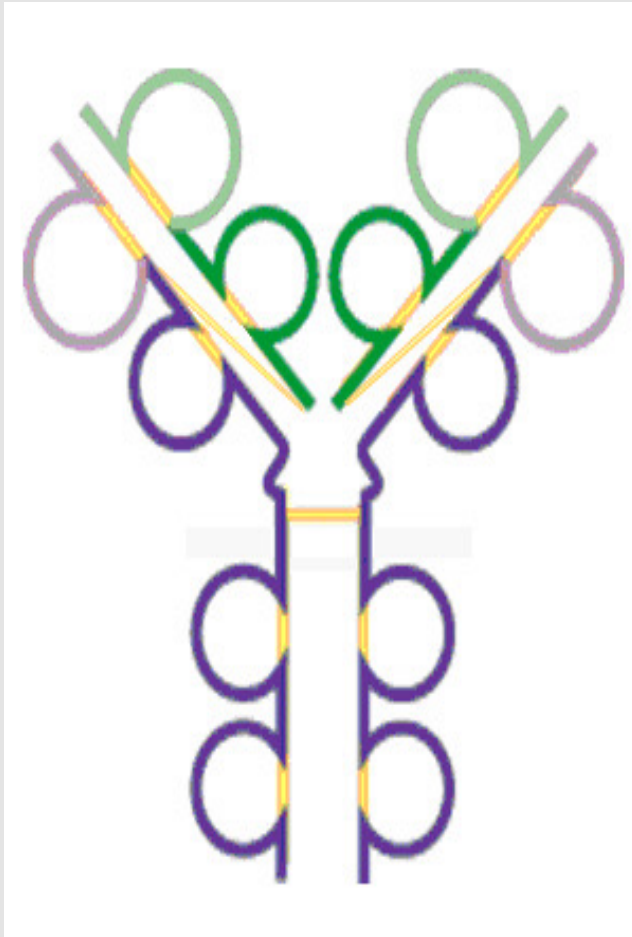


# Domain structure



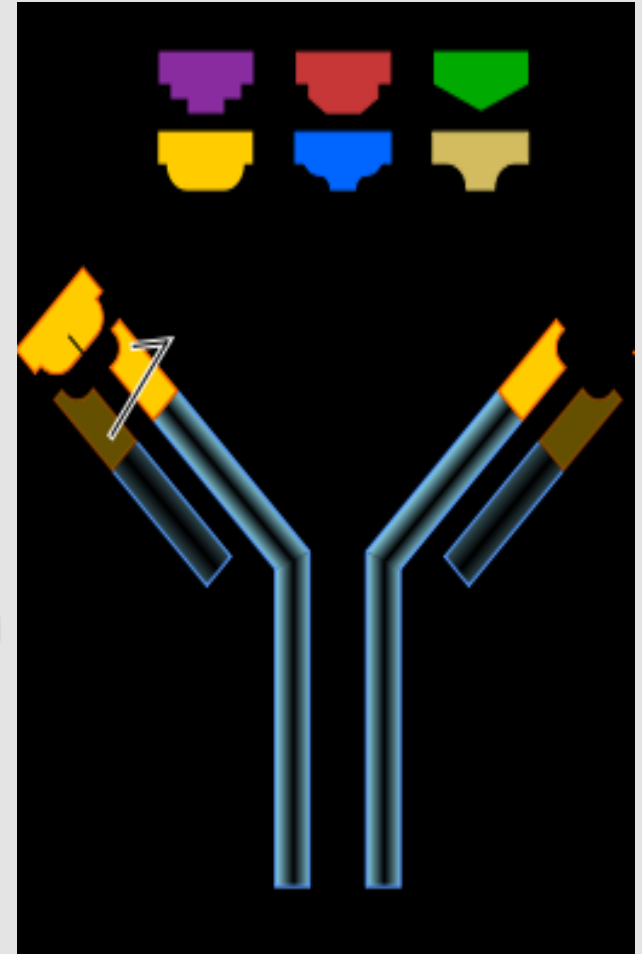
Well conserved amino acid sequence designed by 110 amino acids closed to a “ring shape” with disulphide bound.

# Immunoglobulin molecule



**CDR**  
**Variable region**  
**Idiotyp**  
**Fab fragment**

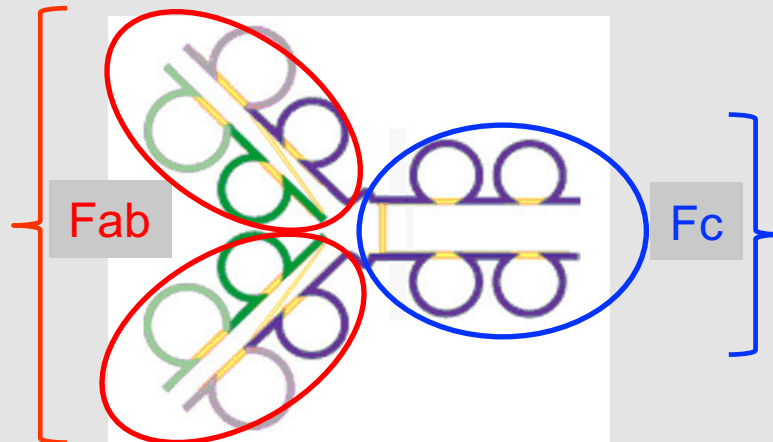
**Constant region**  
**Isotype**  
**Fc fragment**



# Immunoglobulin functions

## **Monofunctional** character:

Specific antigen recognition and - binding



## **Polyfunctional** character:

- Signaltransduction,
- Komplement fixation,
- Opsonisation,
- Immuncomplex formation
- FcR binding

# Immunoglobulin isotypes

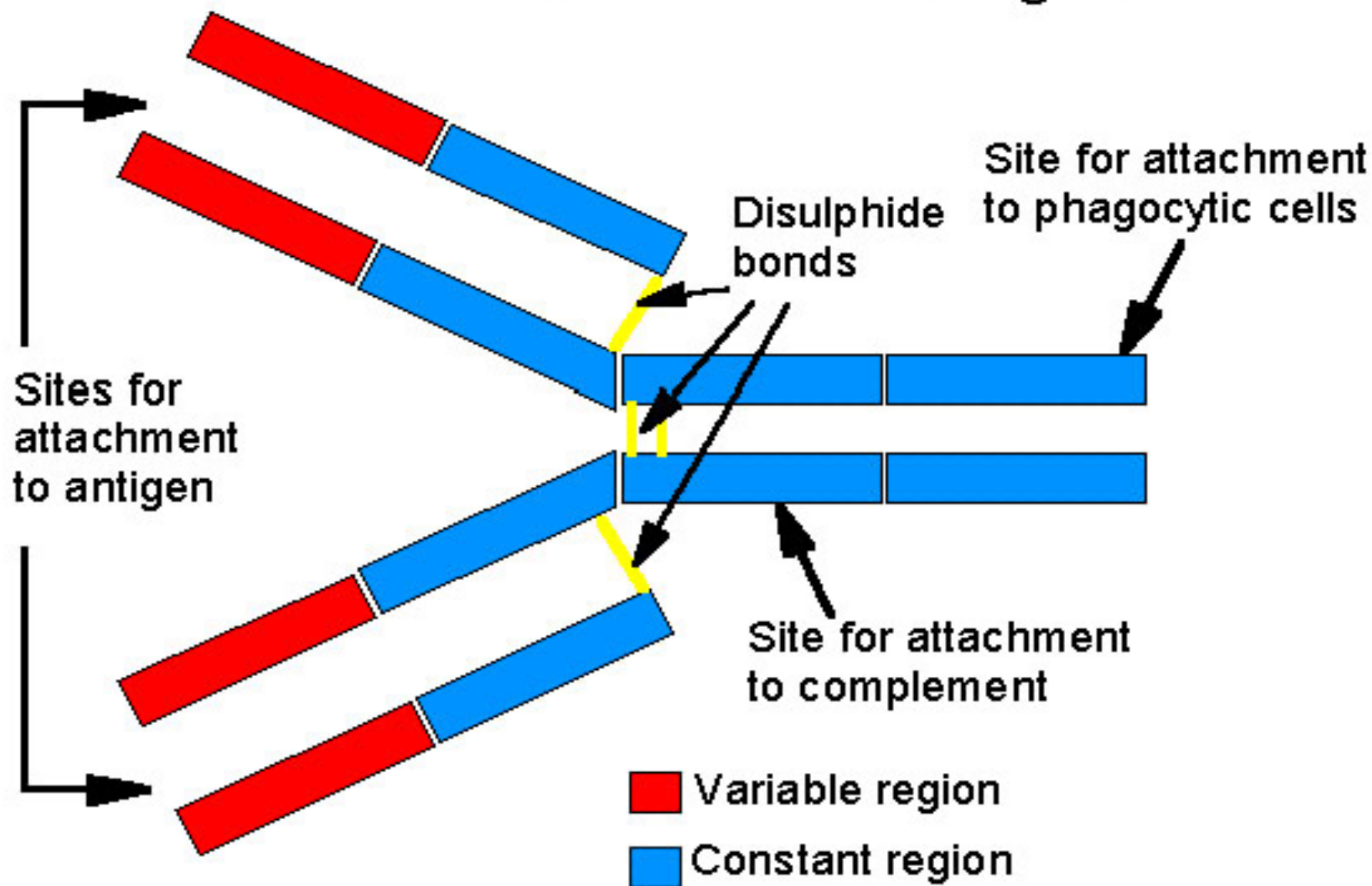
- Based upon the constant structures of heavy (**H**) and light (**L**) chains
- **CH isotypes:** called Ig classes and subclasses as **IgG, IgM, IgA, IgD** and **IgE**. All classes are represented in a normal serum (except the membrane bound IgD) as isotype variants.
- **CL** chain exists in two **isotypic forms:** kappa (**κ**) and lambda (**λ**), which can associate with all heavy chain isotypes.

<b>Heavy chain</b>	<b>Light chain</b>	<b>Immuno-globulin Class</b>	<b>Immuno-globulin Subclass</b>
$\gamma 1$	$\kappa$ or $\lambda$	IgG	IgG1
$\gamma 2$	$\kappa$ or $\lambda$		IgG2
$\gamma 3$	$\kappa$ or $\lambda$		IgG3
$\gamma 4$	$\kappa$ or $\lambda$		IgG4
$\alpha 1$	$\kappa$ or $\lambda$	IgA	IgA1
$\alpha 2$	$\kappa$ or $\lambda$		IgA2
$\mu$	$\kappa$ or $\lambda$	IgM	
$\delta$	$\kappa$ or $\lambda$	IgD	
$\epsilon$	$\kappa$ or $\lambda$	IgE	

Pronunciation of Greek letters:

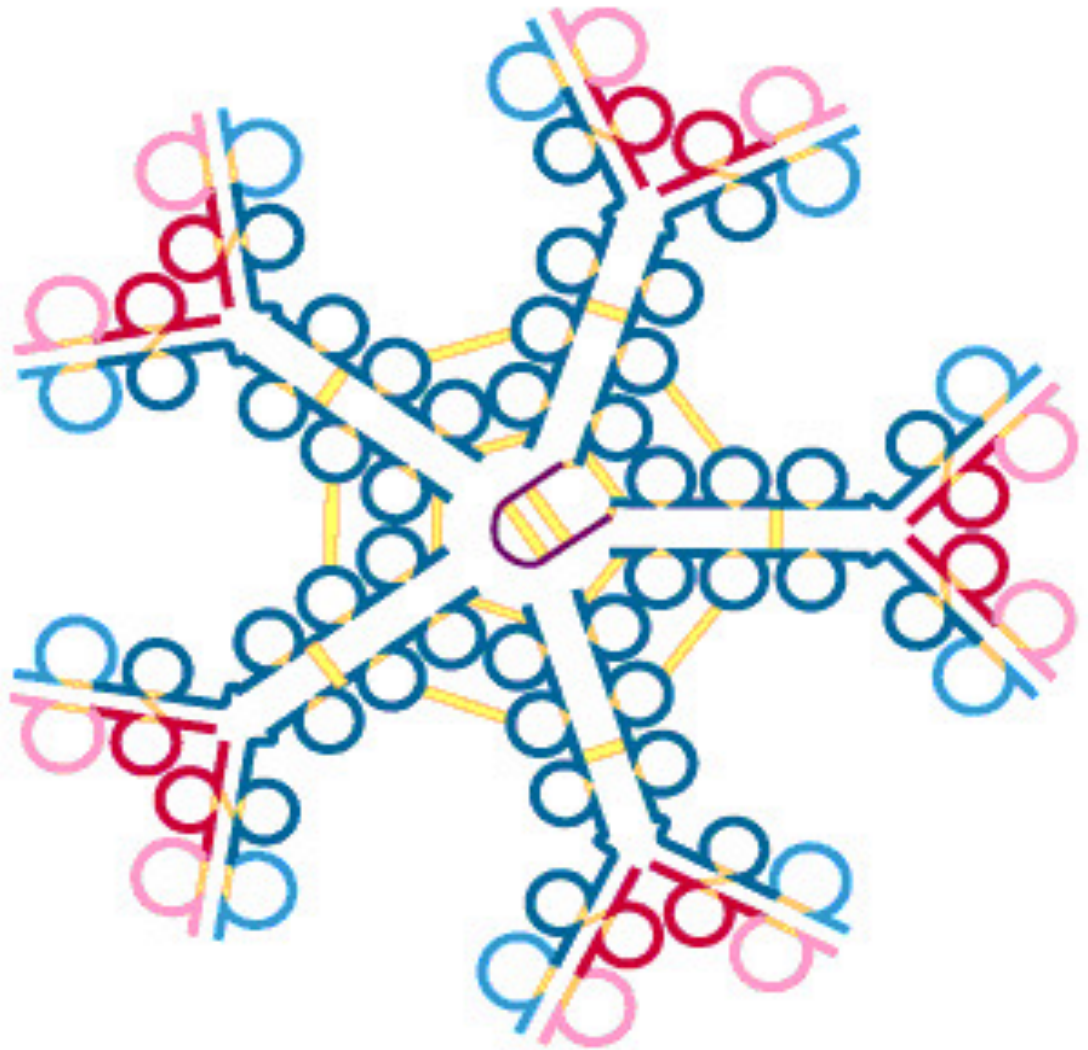
$\gamma$  gamma     $\alpha$  alpha     $\mu$  mu     $\delta$  delta  
 $\epsilon$  epsilon     $\kappa$  kappa     $\lambda$  lambda

# Structure of Immunoglobulin G1



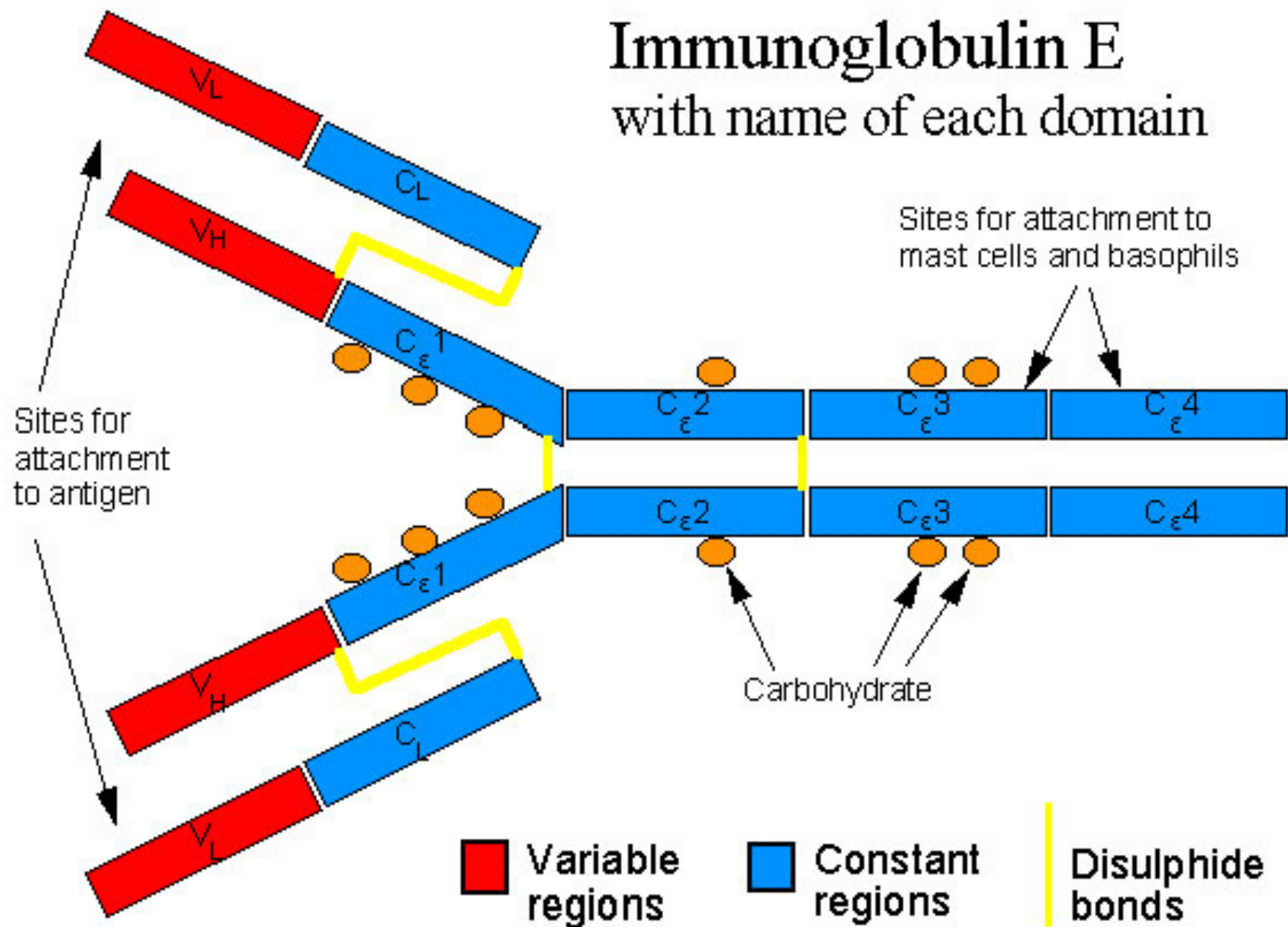


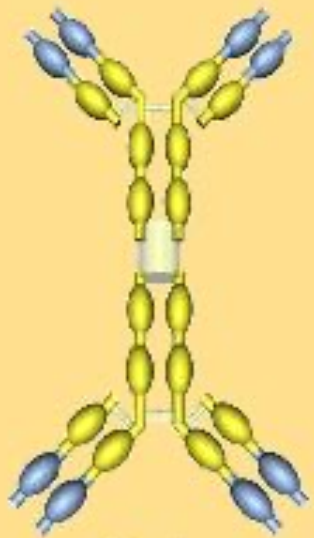
# IgA and IgM



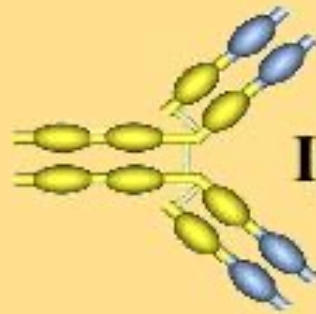
# Immunoglobulin E

with name of each domain

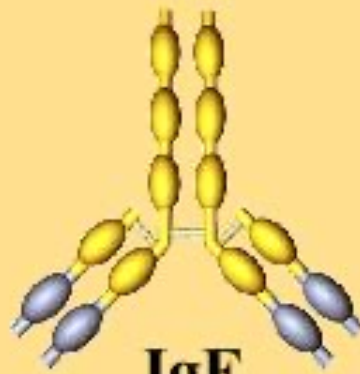




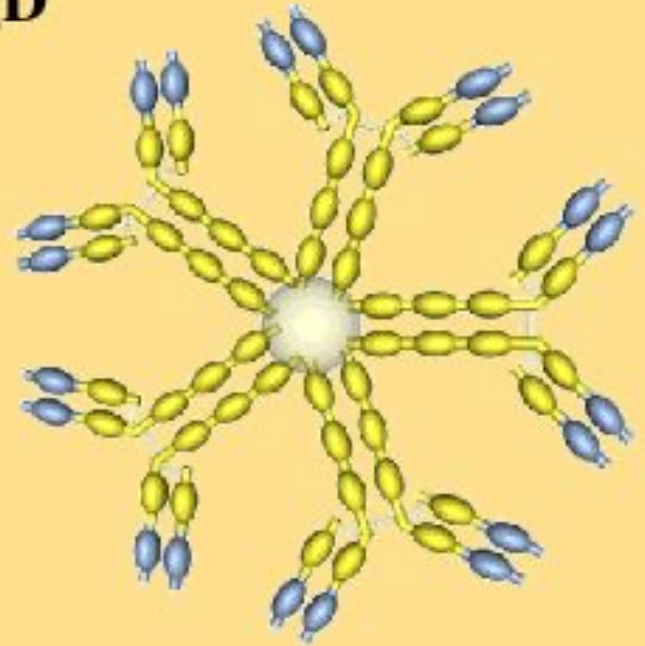
**IgA**



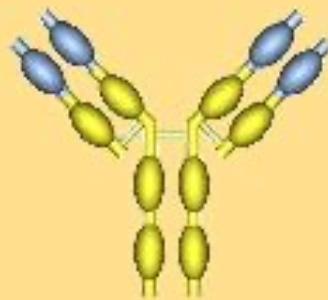
**IgD**



**IgE**



**IgM**

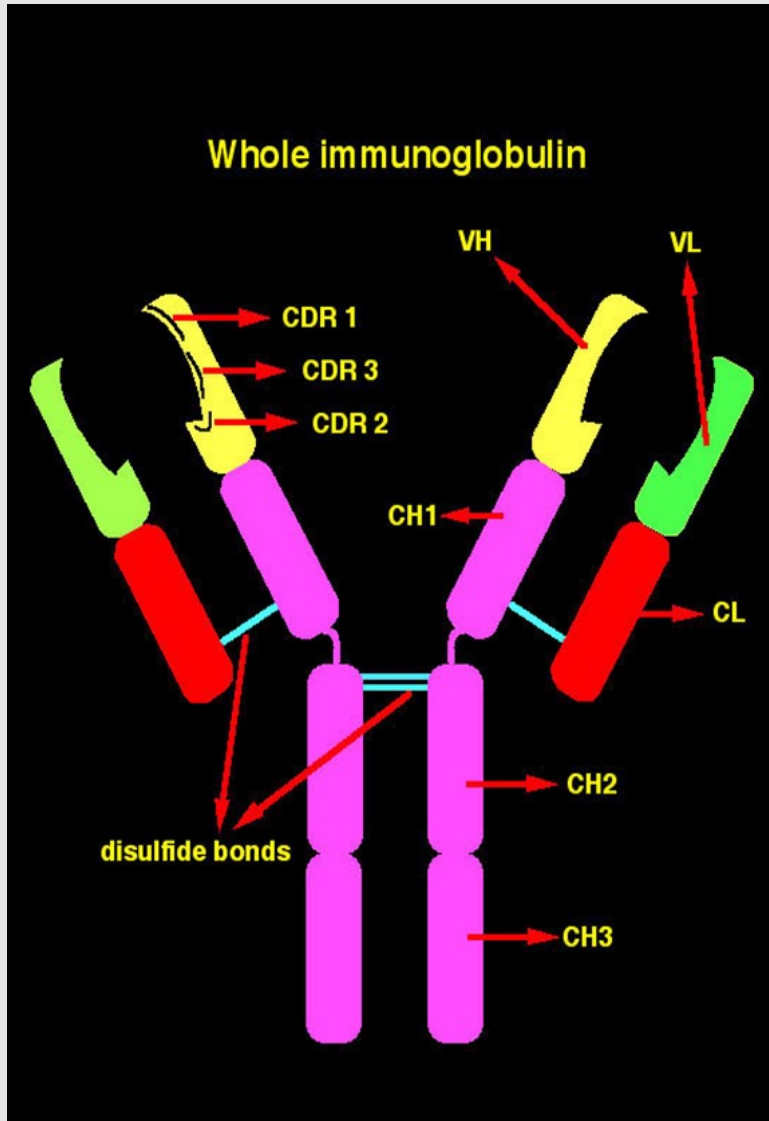


**IgG**

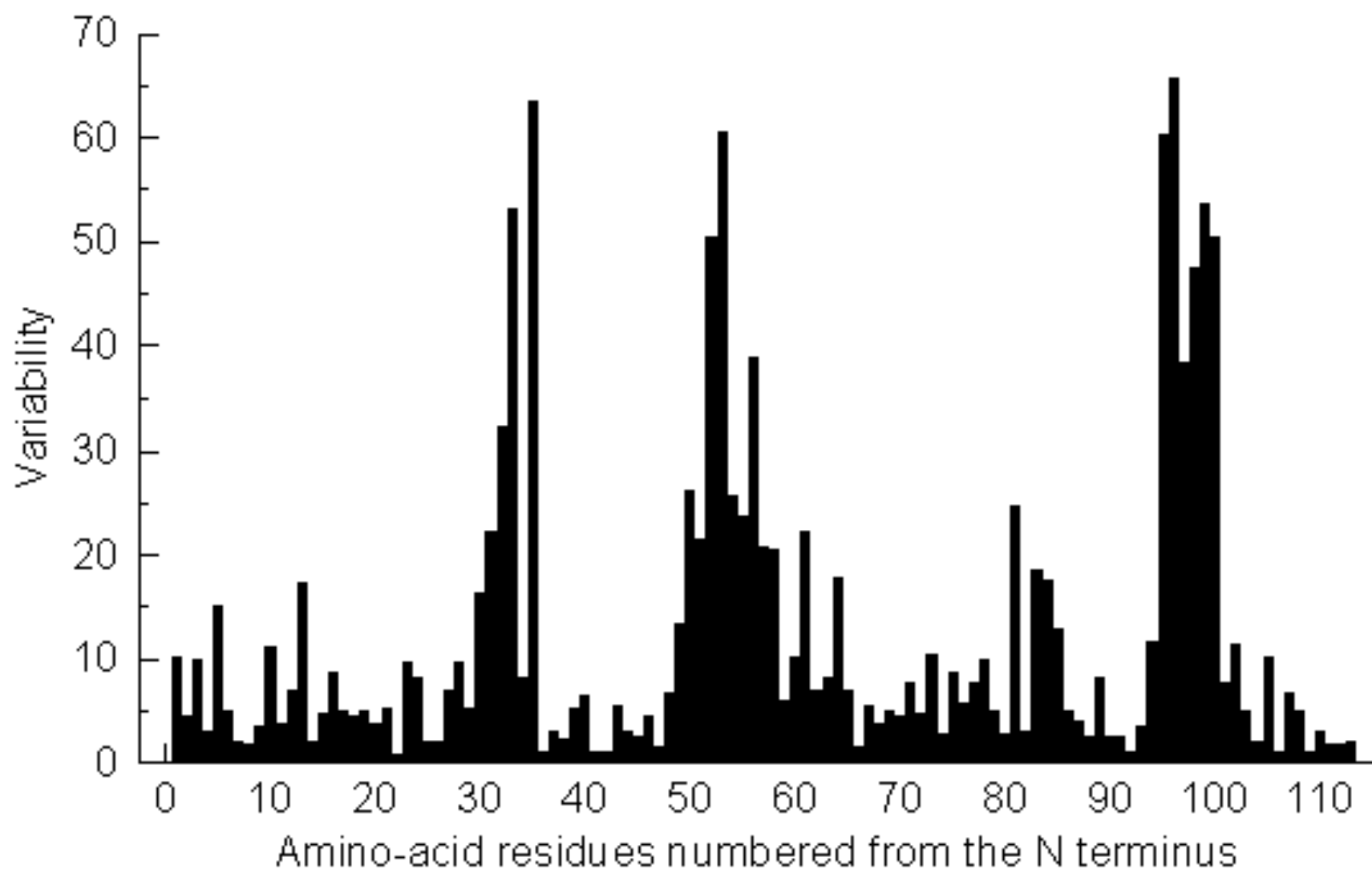
# Immunoglobulin idiotype

Individual determinants in **V regions**, specific for each antibody.

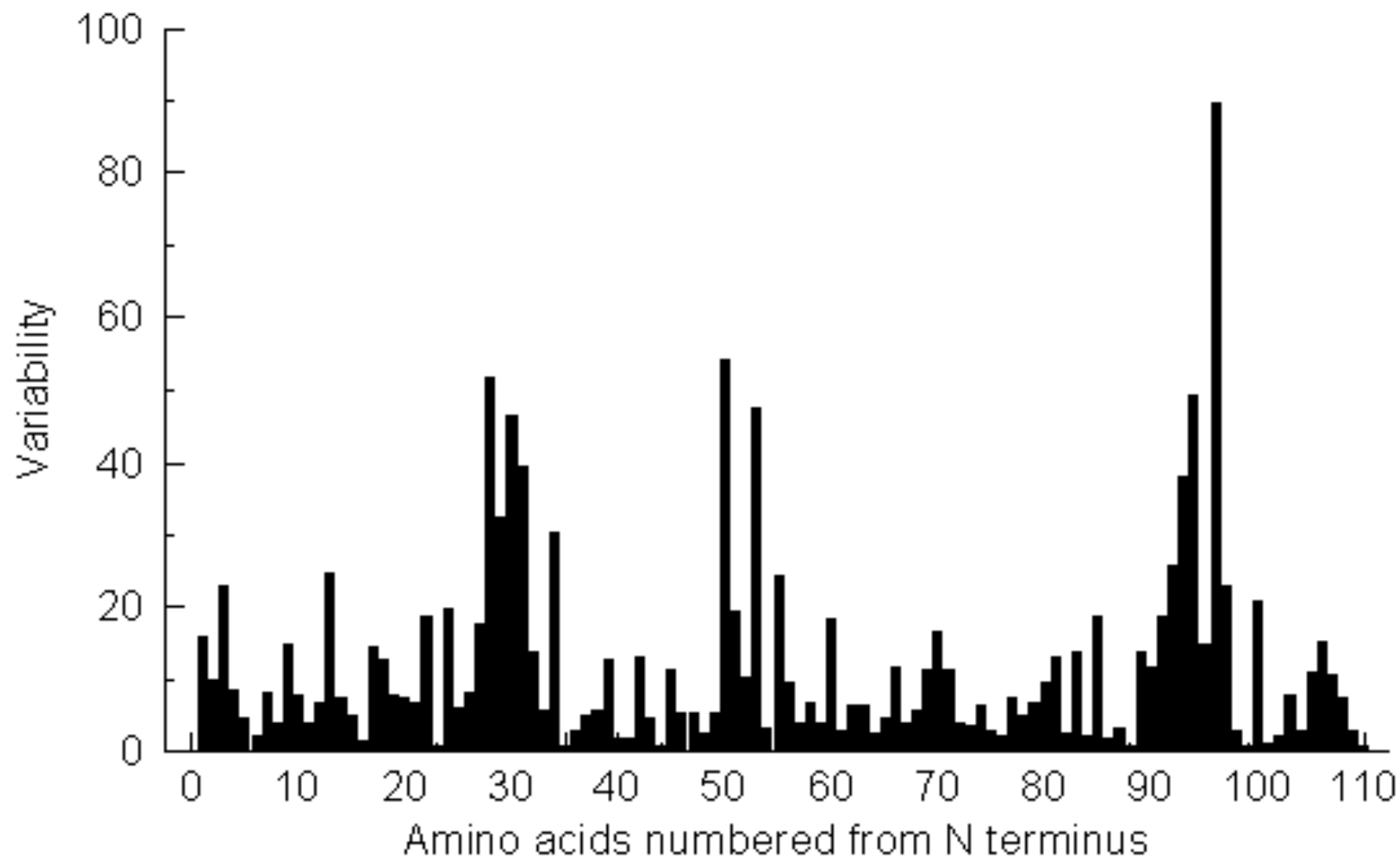
The N terminal Ig domain contains V region forming the antigen binding site: clustering the 3 hyper variable sequences close to each other on both chains - the variation of 3 x 3 results tremendous diversity.



## Variability of amino-acid residues in the variable region of immunoglobulin H chains



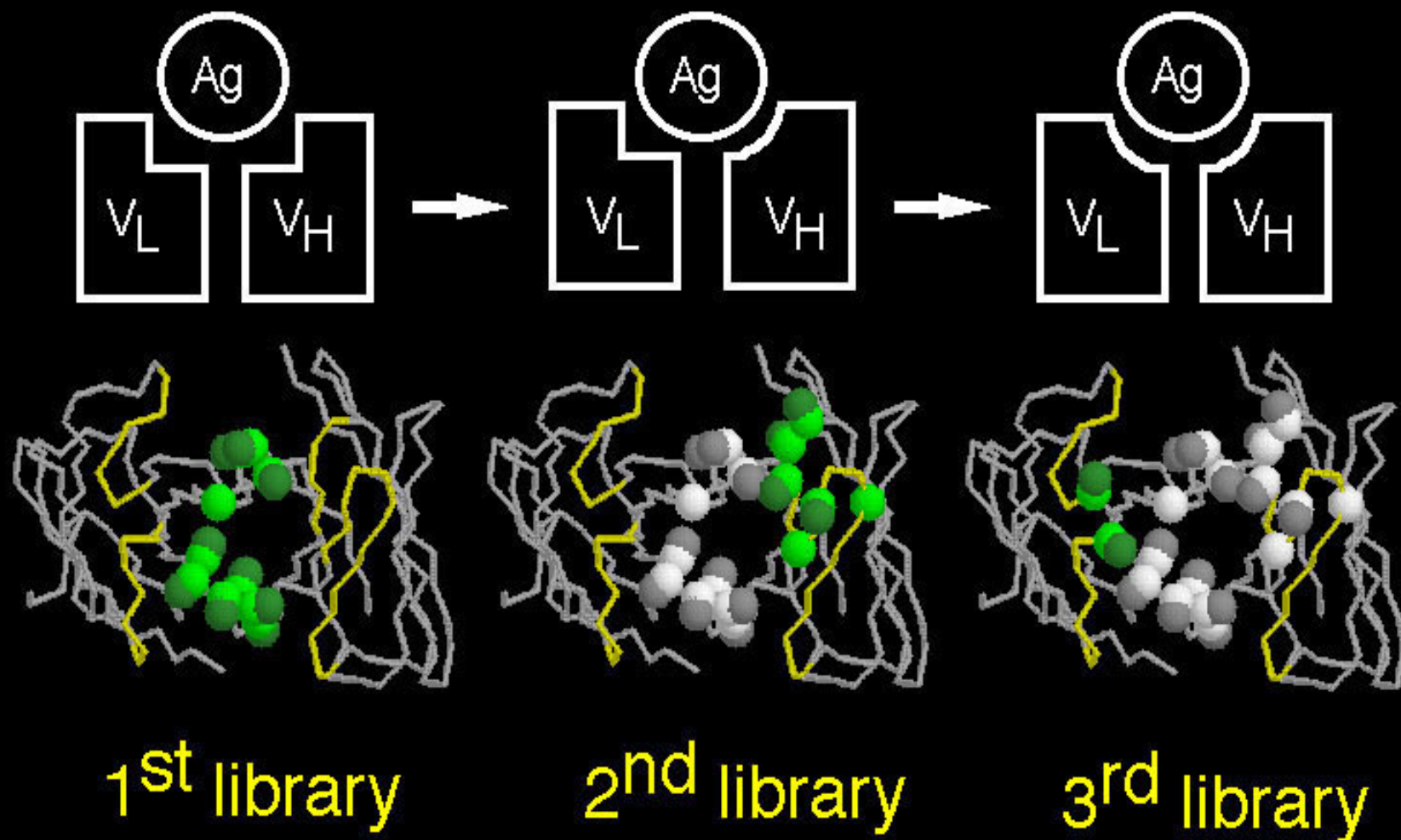
## Variability of amino-acid residues in the variable region of Immunoglobulin L chains





# Antibody affinity maturation

Pini et al. (1998) *J. Biol. Chem.* *273*, 21769-21776



# Antigen Recognition by T Cells

## “MHC-restriction”

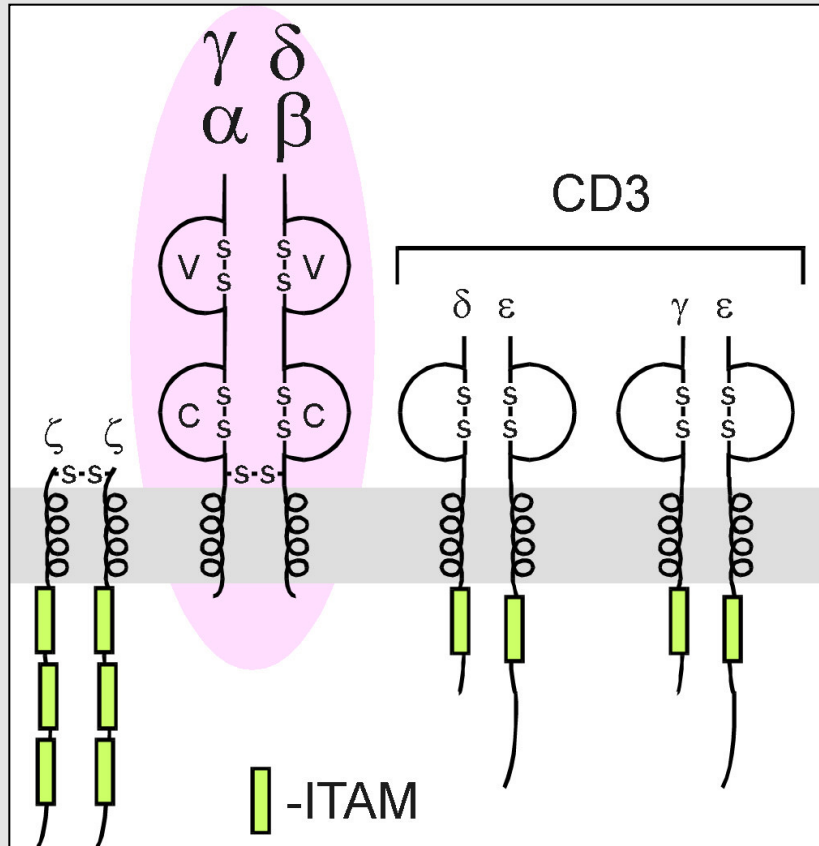
T cells recognize antigens only displayed on surfaces of the body's own cells as MHC-peptide complexes.

CD8+ (cytotoxic) T-cells	MHC I-peptide complex
CD4+ (helper) T-cells	MHC II-peptide complex

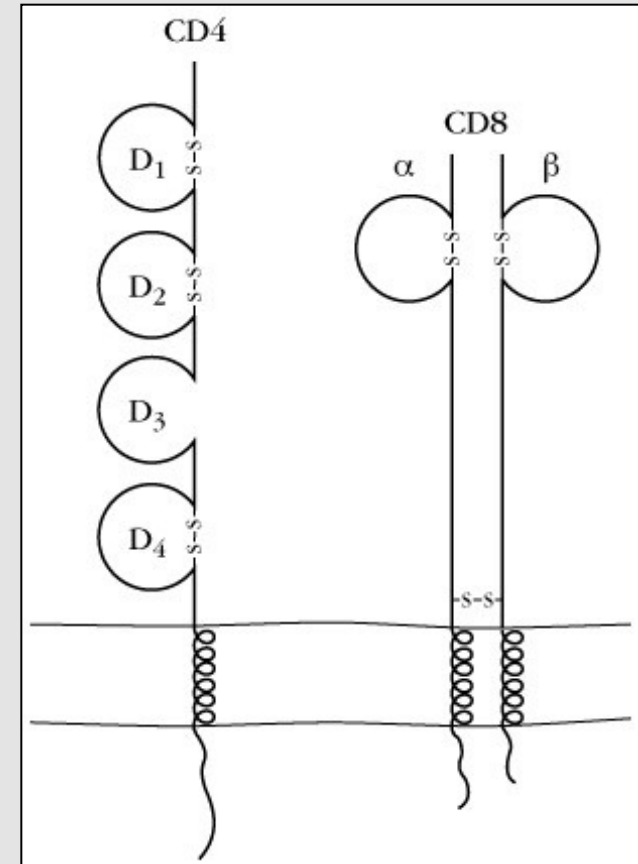
R. M. Zinkernagel & P. C. Doherty – Nobel Prize for Physiology or Medicine (1996.)



# T cell receptor complex on mature T cells



+

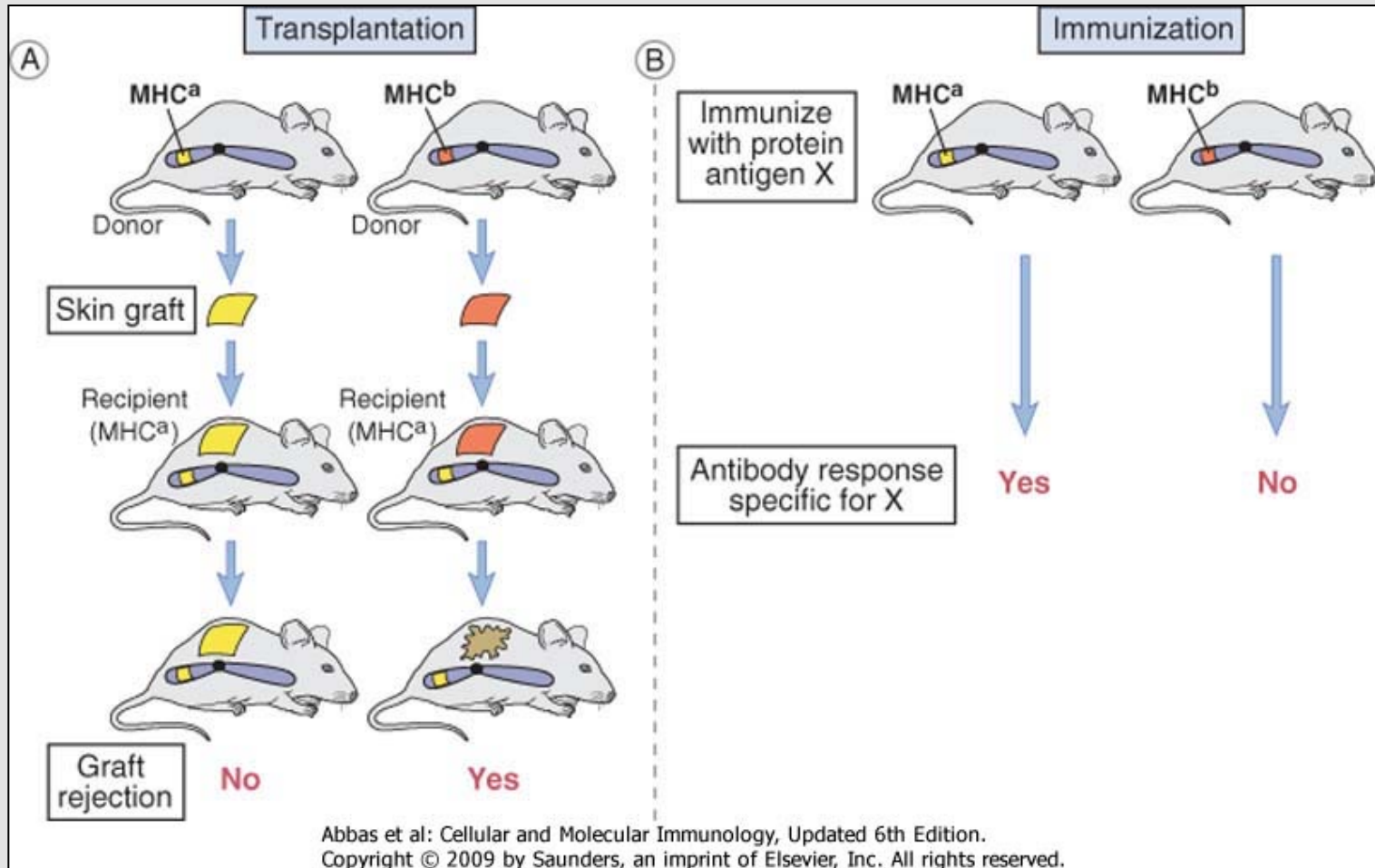


$\alpha\beta$  TcR – CD4+ vagy CD8+)  
 $\gamma\delta$  TcR – CD4-CD8-

# Definition

- **MHC=Major Histocompatibility Complex; HLA=Human Leukocyte Antigen**
- Discovery: transplantation experiments between inbred mouse strains expressing different MHC genes.
- **Inbred mouse strains:** mating of siblings for 20 generations → all mice are homozygous at every genetic locus (genetically identical = “*syngeneic*”)
- In case of polymorphic genes (eg. MHC) each inbred strain expresses a single allele from the original population
- Different inbred strains are “*allogeneic*” to each other = carry different alleles

# Discovery of the mouse MHC



Histocompatibility-2 (H-2) locus

K, D (MHC Class I) genes  
responsible for graft rejection

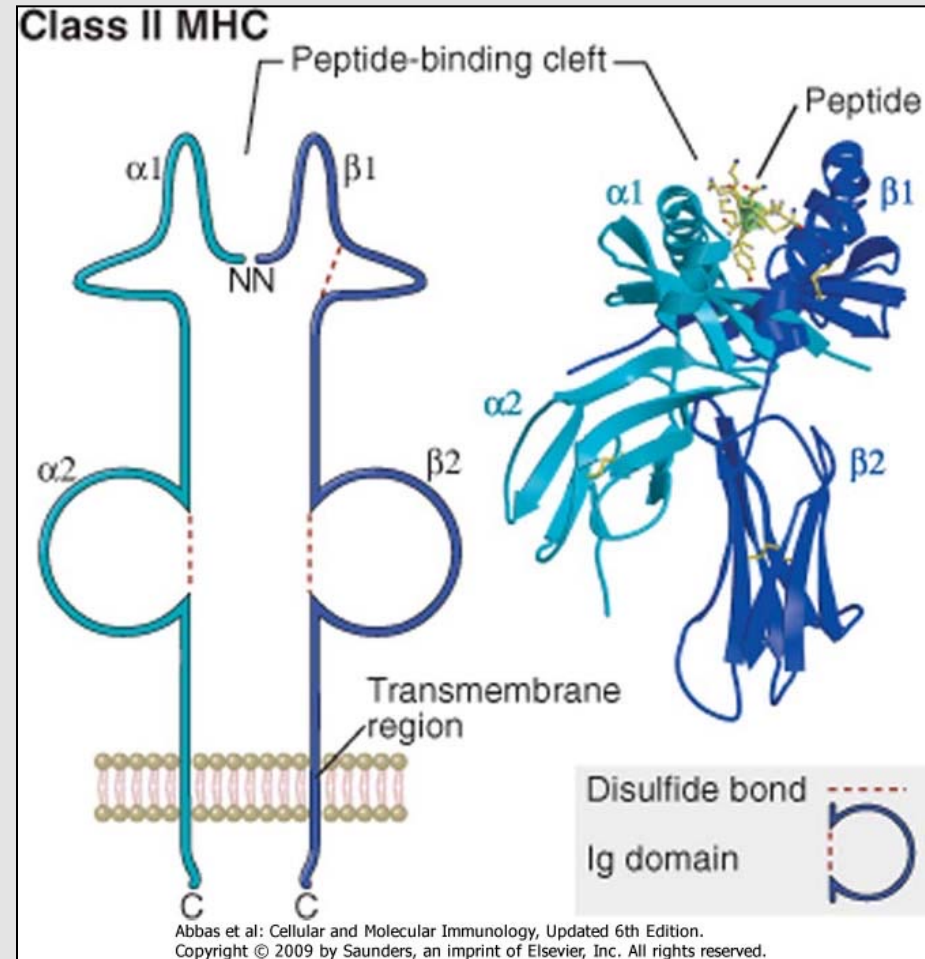
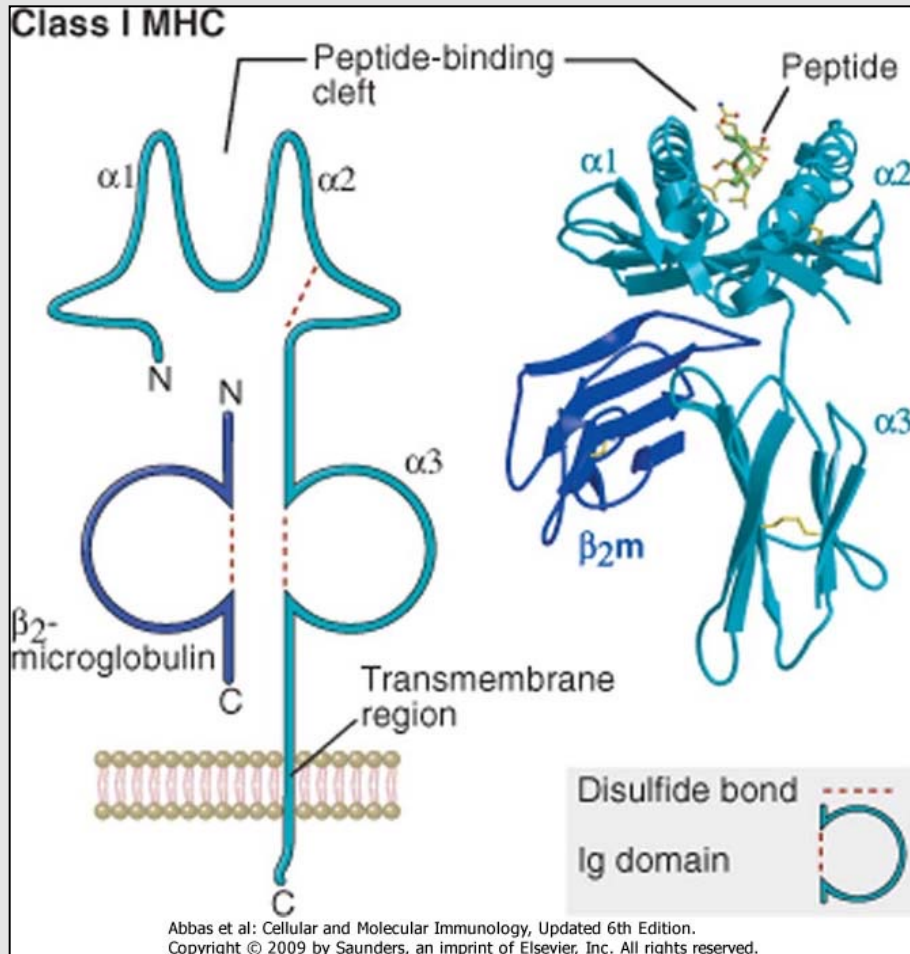
Immune response (I<sub>r</sub>) genes

A, E (MHC Class II) genes determine  
reactivity to different protein antigens

# Features of MHC-I and MHC-II molecules

Feature	Class I MHC	Class II MHC
Polypeptide chains	$\alpha$ (44-47 kD) $\beta_2$ -Microglobulin (12 kD)	$\alpha$ (32-34 kD) $\beta$ (29-32 kD)
Locations of polymorphic residues	$\alpha 1$ and $\alpha 2$ domains	$\alpha 1$ and $\beta 1$ domains
Binding site for T cell coreceptor	$\alpha 3$ region binds CD8	$\beta 2$ region binds CD4
Size of peptide-binding cleft	8-11 AA peptides	10-25 AA peptides
Nomenclature		
Human	HLA-A, -B, -C	HLA-DR, -DQ, -DP
Mouse	H-2K, H-2D, H-2L	I-A, I-E

# The structure of MHC-I and MHC-II





# Peptide binding of MHC-I and MHC-II

MHC-I

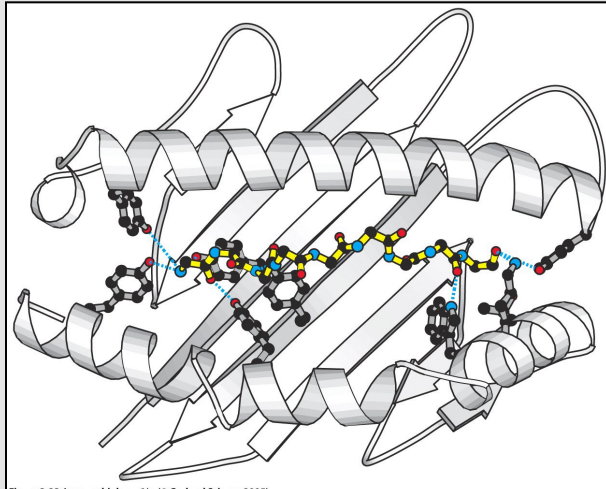


Figure 3-23 Immunobiology, 6/e. (© Garland Science 2005)

MHC-II

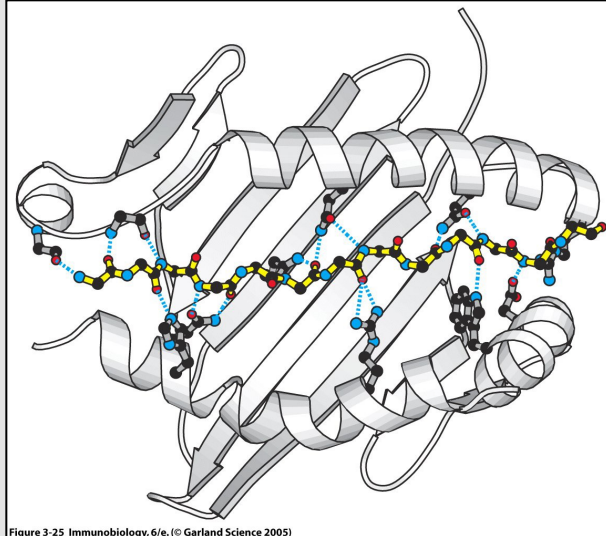
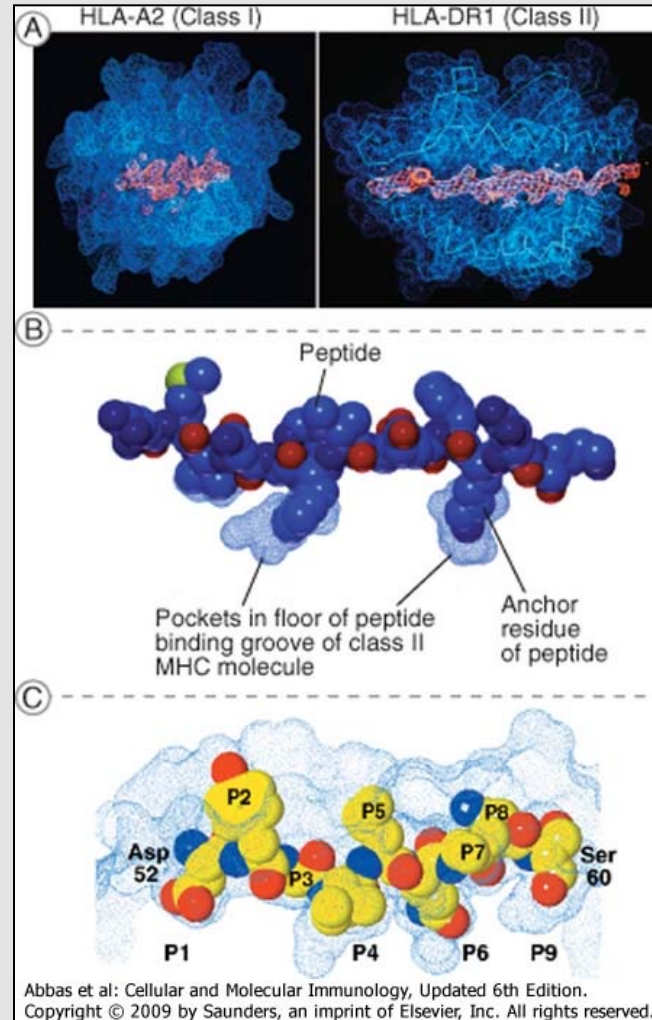
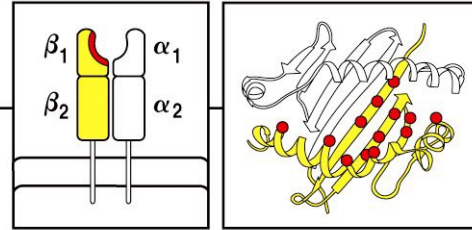
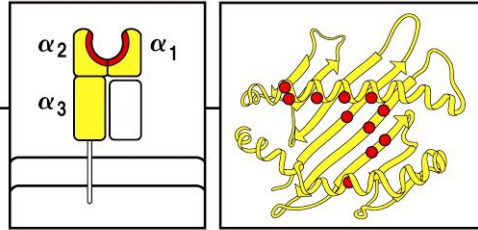


Figure 3-25 Immunobiology, 6/e. (© Garland Science 2005)



Non-covalent interaction between “**anchor**”-residues of the peptides and the small pockets in the  $\beta$ -sheet “**floor**” of the peptide-binding cleft.

# Peptide binding of MHC-I and MHC-II



1 MHC molecule can bind 3-500 different peptides which contain the appropriate “**anchor**”-residues at key positions.

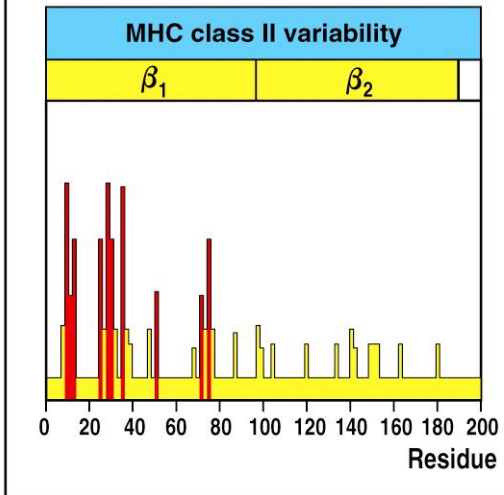
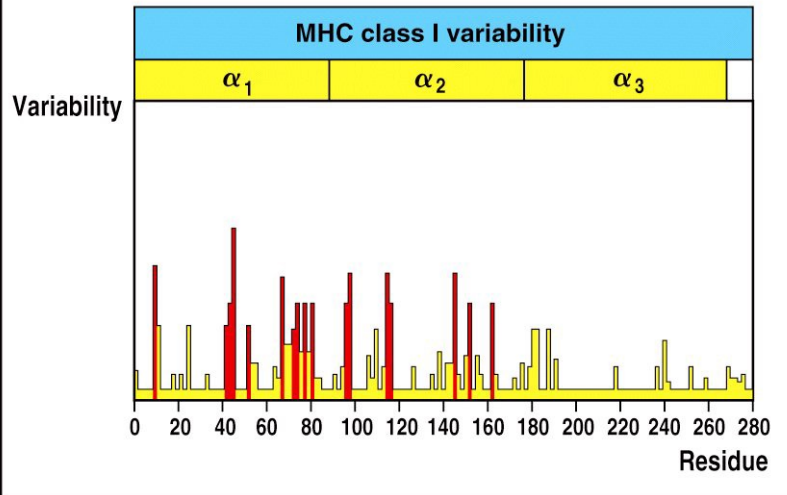
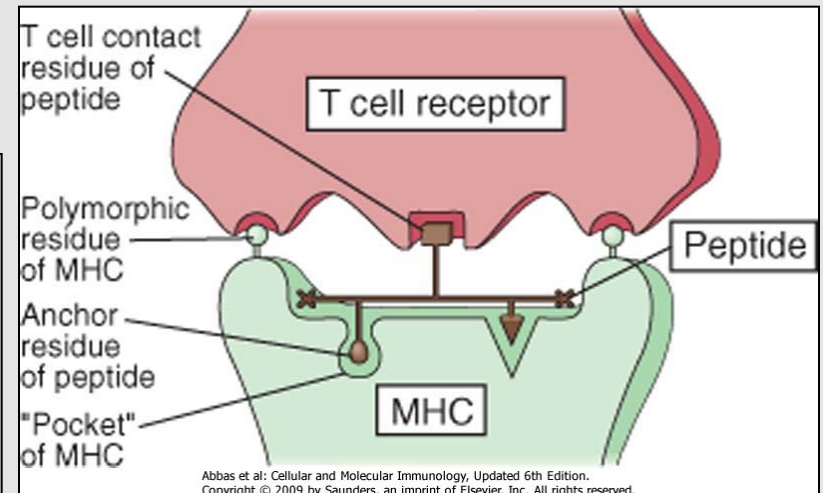


Figure 5-16 Immunobiology, 6/e. (© Garland Science 2005)

Polymorphic AA residues of the MHC molecules are located around the peptide-binding cleft – responsible for **peptide-specificity** and **TcR-binding**.



Abbas et al: Cellular and Molecular Immunology, Updated 6th Edition. Copyright © 2009 by Saunders, an imprint of Elsevier, Inc. All rights reserved.

# MHC-II peptide binding

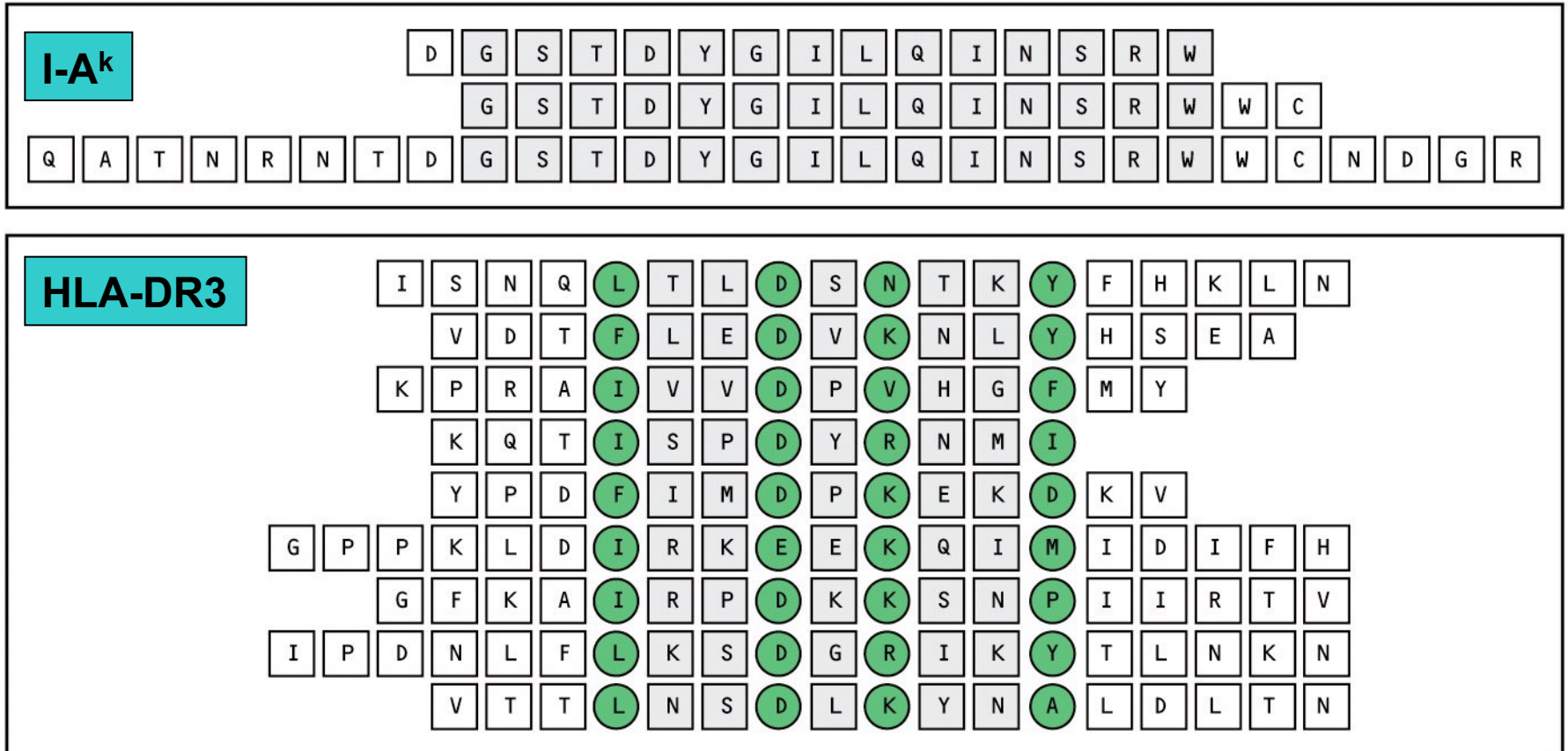
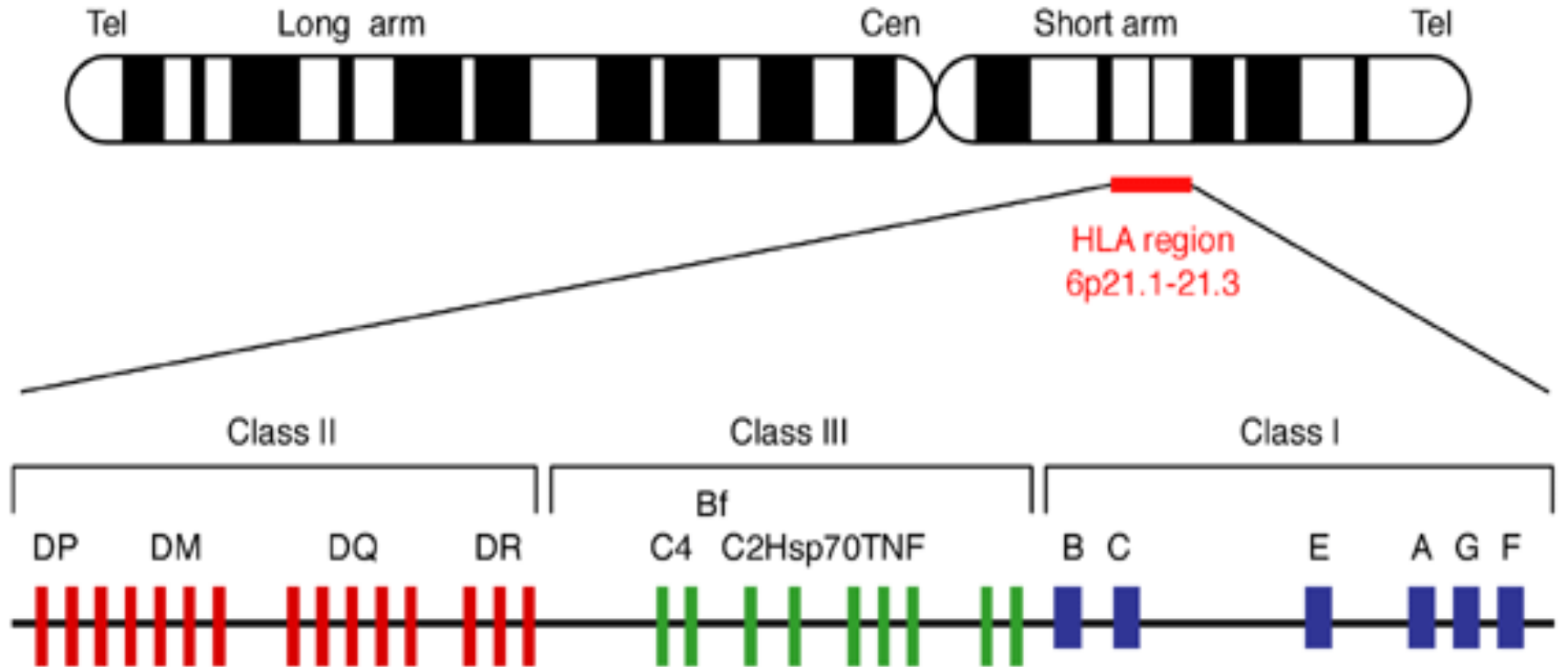


Figure 3-26 Immunobiology, 6/e. (© Garland Science 2005)



# HLA map

Chromosome 6



Gene map of the human leukocyte antigen (HLA) region

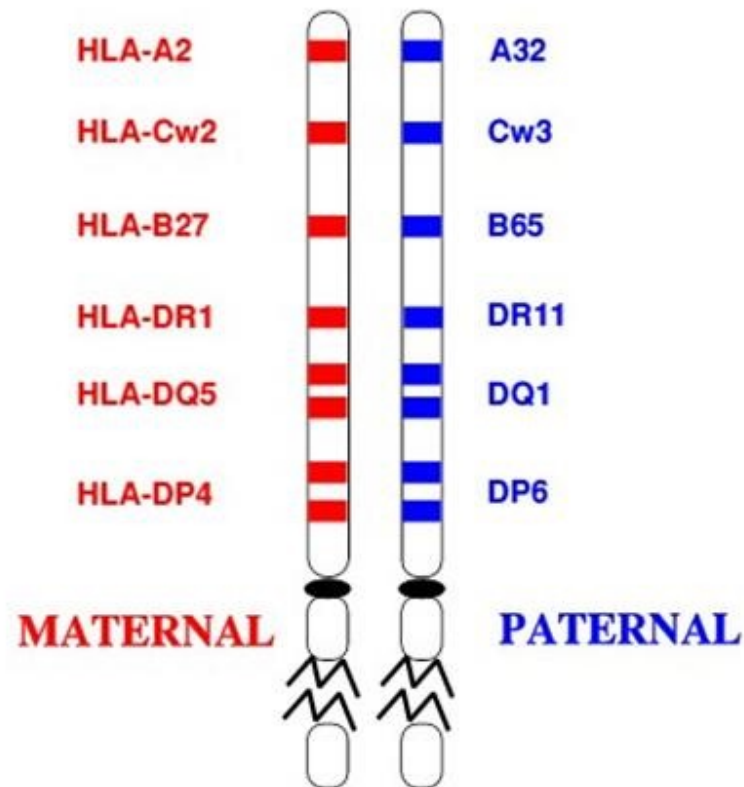
Expert Reviews in Molecular Medicine ©2003 Cambridge University Press

# Genetics of MHC (HLA)

1. **polygenic**: *several* different class I and class II **genes** encoding proteins with different specificities. In human there are 3 classical class I molecules (**HLA-A, B, C**) and 3 classical class II molecules (**HLA-DR, DP, DQ**).
2. highly **polymorphic**: *multiple alleles of each gene* (most individuals are likely to be heterozygous at each locus). The HLA-A has more than 20, B has more 50, and C more than 10 alleles. HLA-DR has 20, HLA-DQ has 9, and HLA-DP has 6 alleles.  
**Nomenclature**: eg. HLA-B\*2705= first 2 places – main alleles, last 2 places - suballeles. (w=workshop - not final)
3. **co-dominant**: Alleles are expressed from both MHC haplotypes in any one individual, and the products of all alleles are found on all expressing cells.

# Genetics of MHC (HLA)

Most Humans are heterozygous  
at the MHC



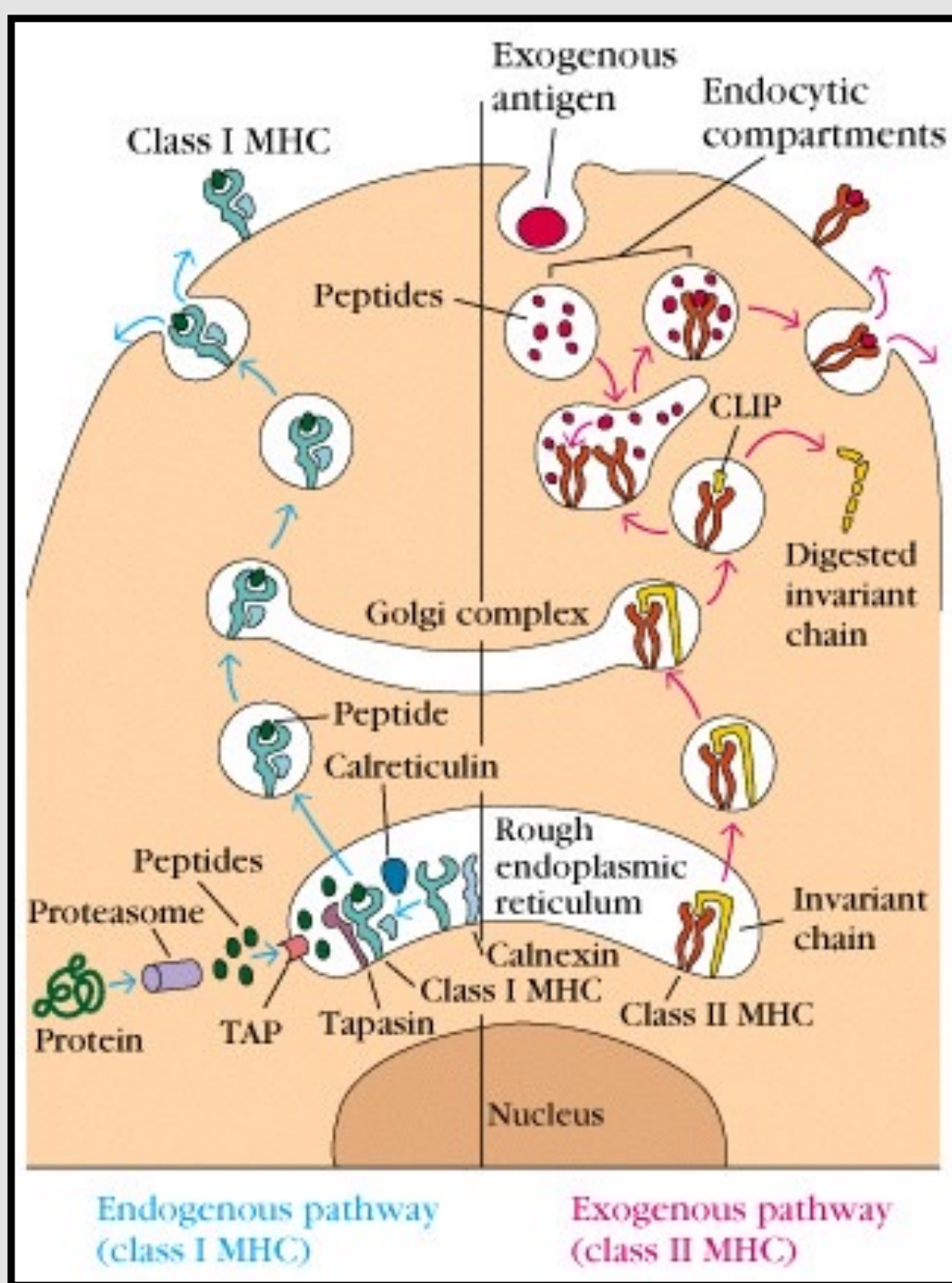
# Expression pattern of MHC I and MHC II

**MHC I**      All nucleated cells + platelets

**MHC II**      Professional antigen presenting cells

- Dendritic cells
- B cells
- Macrophages
- (Thymic epithelial cells)

Facultative antigen presenting cells  
eg. inflammatory epithel



Endogenous pathway  
(class I MHC)

Exogenous pathway  
(class II MHC)

# Antigen Presentation on MHC I

- Cytosolic, mainly normal or viral/modified proteins
  - Proteasomal degradation

Peptide transfer to the ER (TAP1&2)

MHC I chains produced into ER by ribosomes

Chaperons: calnexin, calreticulin, Erp57

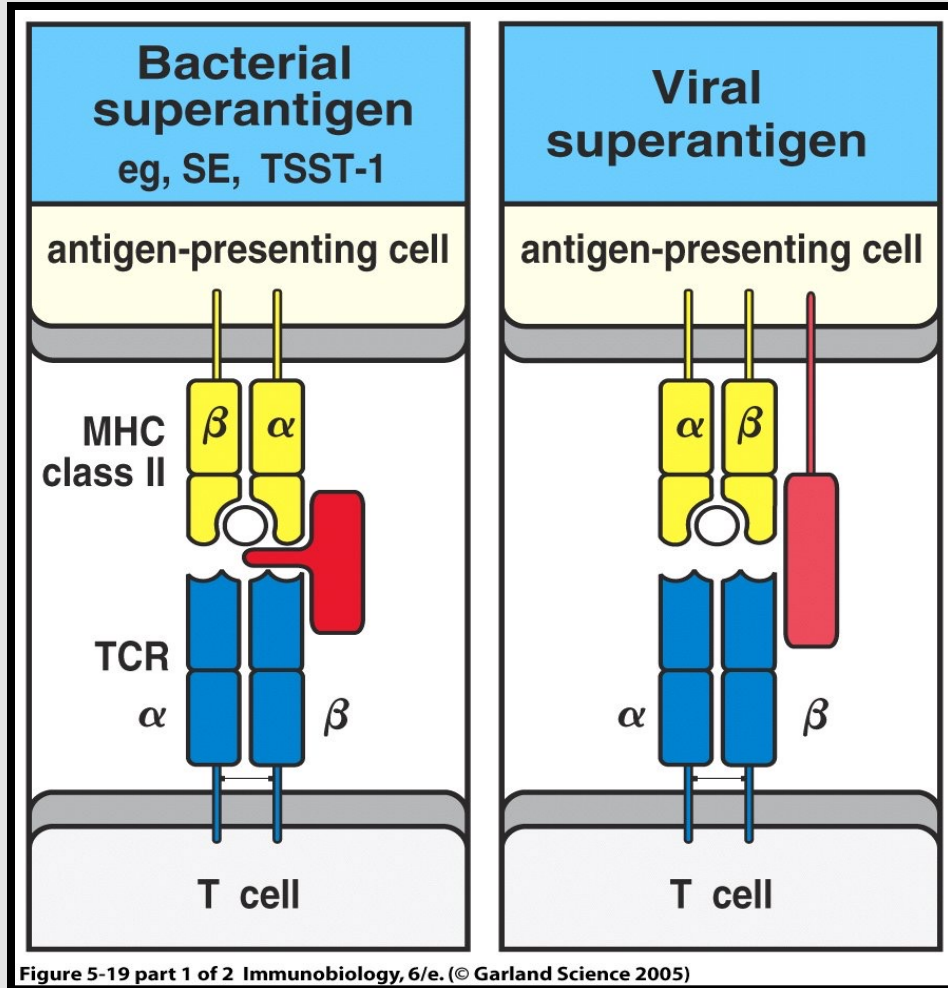
Tapasin and TAP1&2

MHCI & peptide binding within the ER

# Antigen Presentation on MHC II

- Endocytosed proteins: bacteria, bacterial product, internalised receptor bound peptide, parts of another cell
- Endosomal degradation
- MHCII chains produced into the ER by ribosomes
- invariant chain
- HLA-DM: MHC II specific chaperon
- CLIP=class II associated invariant chain peptide
- MHC II & peptide binding in endosomes outside the ER

# Superantigens



Compared to a normal antigen-induced T-cell response where 0.001-0.0001% of the body's T-cells are activated, SAGs (endotoxins) are capable of activating up to 20% of the body's T-cells. This causes a massive immune response (toxic shock syndrome) that is not specific to any particular epitope on the SAg.

T cells produce cytokines - systemic toxicity, suppression of adaptive immune response („Cytokine tsunami“)



# **Medical aspects of MHC**

- **Tissue/organ transplantation – donor and recipient must have matching HLA haplotype**
- **HLA-association of diseases (“disease susceptibility”)** – certain diseases appear more frequently in individuals with a specific HLA type

# HLA-association of diseases

Some HLA associated autoimmune diseases

Disease	HLA	Pts <sup>a</sup>	Ctrls <sup>a</sup>	RR <sup>b</sup>
Ankylosing spondylitis	B27	> 95	9	> 150
Subacute thyroiditis	B35	70	14	14
Psoriasis vulgaris	Cw6	87	33	7
Graves disease	DR3	65	27	4
Myasthenia gravis	DR3	50	27	2
Addisons disease	DR3	69	27	5
Rheumatoid arthritis	DR4(some)	81	33	9
Juvenile idiopathic arthiritis	DR8	38	7	8
Celiac disease	DQ2 (+DQ8)	92	28	30
Narcolepsy	DQ6(02)	> 95	33	> 40
Multiple sclerosis	DQ6(02)	86	33	12
Type 1 diabetes	DQ8(+)	81	23	14
Type 1 diabetes	DQ6(02)	< 0.01	33	0.02

<sup>a</sup> The figures show antigen frequencies in a Norwegian population.

<sup>b</sup> RR: relative risk; i.e. how many times more frequent the disease is in those having the corresponding HLA molecule compared to those lacking it.

In: E. Thorsby, B.A. Lie: HLA associated genetic predisposition to autoimmune diseases: Genes involved and possible mechanisms. *Transplant Immunology* 14 (2005) 175 – 182.

In: N. Singh, S. Agrawal, A.K. Rastogi Infectious Diseases and Immunity: Special Reference to Major Histocompatibility Complex. *Emerging Infectious Diseases* 3 (1997) 41-49.

Table 2. Association between human leukocyte antigen (HLA) and some infectious diseases

Disease	HLA Association
<i>Bacterial</i>	
Ankylosing spondylitis	B27
Reiter disease	B27
Acute anterior uveitis	B7
<i>Mycobacterial</i>	
Tuberculosis and leprosy (multibacillary forms)	DR2 (DRB1*1501, 1502)
lepromatous leprosy	DR2 and DQ1
paucibacillary tuberculoid	DR3
<i>Viral</i>	
Dengue fever virus	DR15
Human immunodeficiency virus 1	DR13 (DRB1*1301, 1302, 1303)
	DR2 (DRB1*1501)
	DRB1*03011
Hepatitis B virus	DR13
Hepatitis C virus	A2
	DR5
Epstein-Barr virus	B35.01
	A11
	B7
<i>Parasitic</i>	
Malaria	B53
Scabies	A11
Diffuse cutaneous leishmaniasis	A11, B5, B7
Localized cutaneous leishmaniasis	A28, Bw22, DQw8 Bw22, DR11, Qw7 Bw22, Dqw3
Schistosomiasis	B5, DR3
Visceral leishmaniasis	A26

Thank you for your attention!

