



# 12th practice: Immunological aspects of organ transplantation. HLA typing

**Basic Immunology** 

University of Pécs, Clinical Center

Department of Immunology and Biotechnology

Pécs, 2025.

#### **Basic terms**

autolog, allogeneic, xenogeneic graft

auto-, allo-, xeno-transplantation

#### Cornea

From cadaver Immunosuppression not required 40,000 transplants per year

#### Lung

From brain-dead donor Procedure recently developed; little data available 845 transplants in 1998 Often heart/lung transplant (45 in 1998)

#### Heart

From brain-dead donor HLA matching useful but often impossible Risk of coronary artery damage, perhaps mediated by host antibody 2,340 transplants in 1998

#### Liver

From cadaver Surgical implantation complex Resistant to hyperacute rejection Risk of GVHD 4,450 transplants in 1998

#### Skin

Mostly autologous (burn victims) Temporary grafts of nonviable tissue Allogeneic grafts rare, require immunosoppression

#### Blood

Transfused from living donor ABO and Rh matching required Complications extremely rare An estimated 14 million units used each year

#### **Pancreas**

From cadaver Islet cells from organ sufficient 253 transplants in 1998 Increasingly, panreas/kidney transplant for advanced diabetes (965 in 1998)

#### Kidney

From live donor or cadaver ABO and HLA matching useful Immunosuppression usually required Risk of GVHD very low 11,900 transplants in 1998

#### Bone marrow

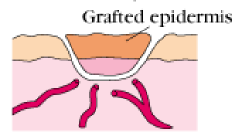
Needle aspiration from living donor Implanted by IV injection ABO and HLA matching required Rejection rare but GVHD a risk

### **Graft acceptance and rejection**

(a) Autograft acceptance Grafted epidermis

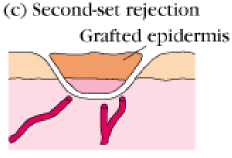


Days 3-7: Revascularization

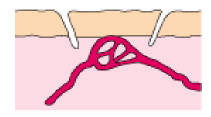


(b) First-set rejection

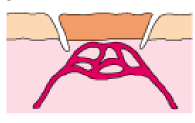
Days 3-7: Revascularization



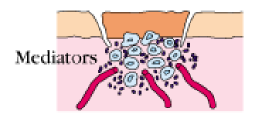
Days 3-4: Cellular infiltration



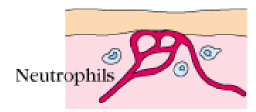
Days 7-10: Healing



Days 7-10: Cellular infiltration



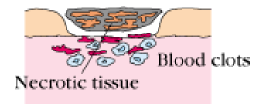
Days 5-6: Thrombosis and necrosis

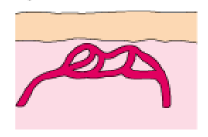


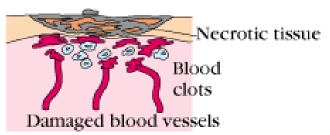
Days 12-14: Resolution



Days 10-14: Thrombosis and necrosis





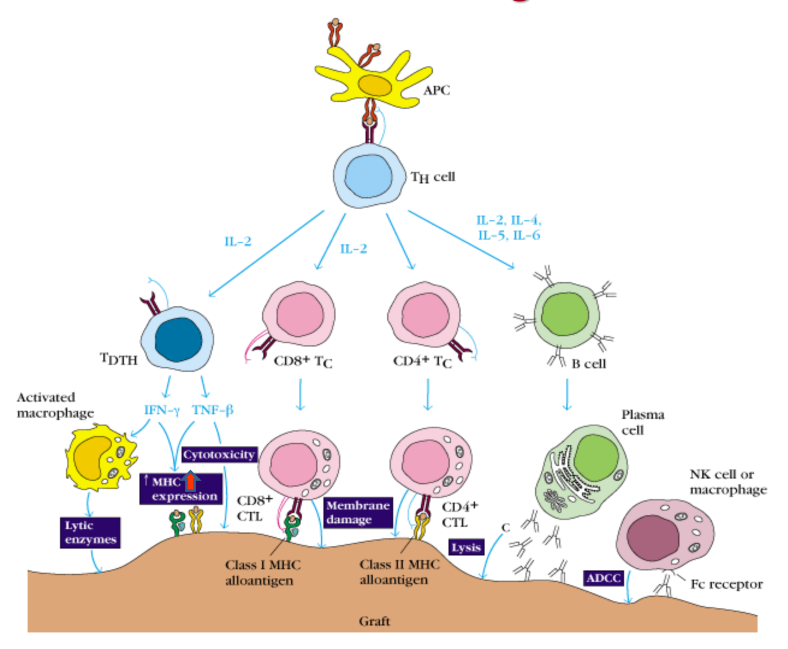


#### Host versus graft reaction

- <u>hyperacute</u> rejection caused by pre-existing antibodies
- <u>acute</u> rejection managed by T cells, ADCC and DTH

 <u>chronic</u> rejection induced by permanent endothelial injuries and complement activation

#### Mechanisms of host versus graft reactions



## Graft versus host reaction – bone marrow or hemopoietic stem cell transplantation

acute GVHD (acute tissue necrosis of the targeted organs)





chronic GVHD (autoimmune-like phenomenon)

## **Bone marrow transplantation**

Advantage	Disadvantage
Autologous	Allogeneic
no GVH	GVH
no rejection	rejection
no matching needed	need matching
	tumour in donor cells
Allogeneic	Autologous
no tumour transfer	grafting tumour cells
graft vs. tumour	(myelosuppression
myelosuppression avoided	possible)

## Discovery of HLA system

> George Snell performed experiments on mice.

Vaccined different individuals with mouse tumours.

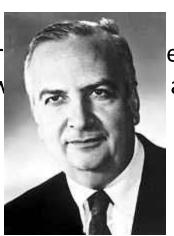
The rejection or engraftment was determined by the genetical difference or identity of the cells.

Genetic system: MHC (Major Histocompatibility Complex)

➤ **Jean Dausset** the same system in humans (1950s): HLA system: Human Leukocyte Antigen

Baruj Benacerraf: the str histoincompatibility betw

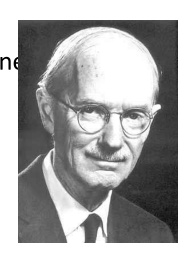
Nobel prize (1980)



Baruj Benacerraf (1920 - 2011)



Jean Dausset (1916 - 2009)



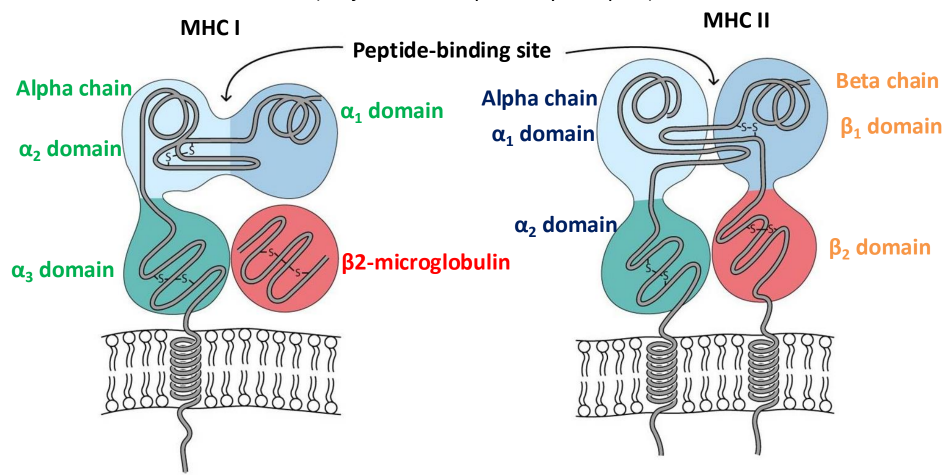
George D. Snell (1903 - 1996)

## Clinical significance of HLA typing

- Checking the immunological compatibility of the donor and the recipient before transplantations to prevent rejections.
- Further diagnostic confirmation of some autoimmune disorders as certain HLA types are more frequent in autoimmune conditions than others, e.g.:
  - HLA-B27: Becheterew's disease, Inflammatory bowel diseases (IBD), Psoriasis
  - HLA-DR1: Rheumatoid arthritis, Ulcerative colitis
  - HLA-DR3: Type I diabetes mellitus, Myasthenia gravis, Hashimoto's thyroiditis
  - HLA-DR4: Rheumatoid arthritis, SLE
  - HLA-DQ2: Celiac disease, Type I diabetes mellitus
  - HLA-DQ8: Celiac disease, Type I diabetes mellitus

## Basics of HLA typing I.

HLA (Human leukocyte antigen)MHC (Major histocompatibility complex)



On all nucleated cells and thrombocytes!

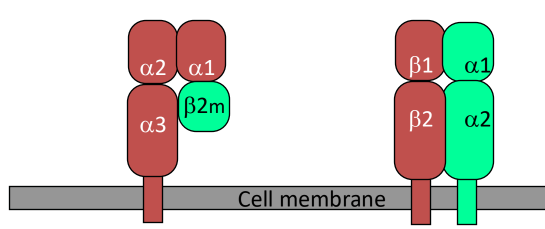
On antigen presenting cells! (e.g. macrophage, dendritic cell, B cell)

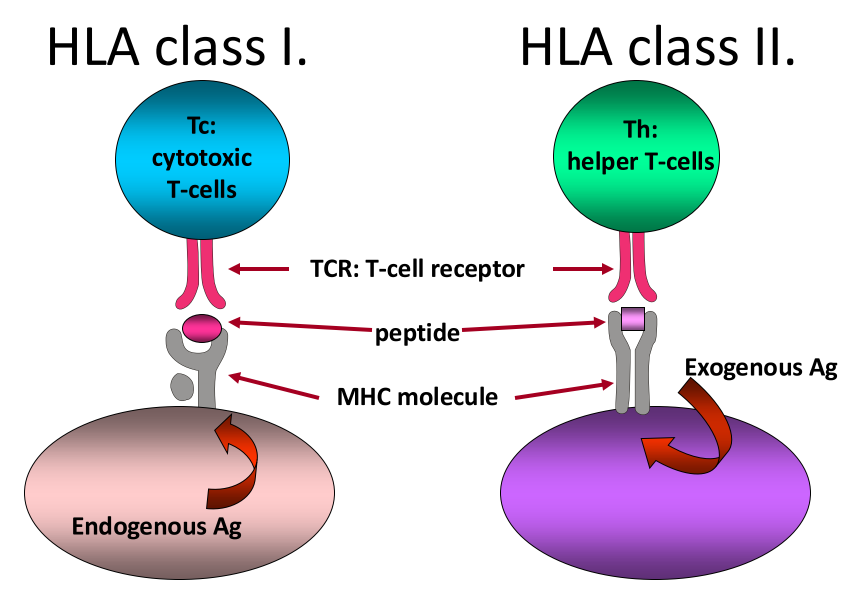
#### HLA class I.

- HLA-A, B, C genes
- Present in all nucleated cells and platelets.
- Different level of expression:
  - high on the cells of the immune system;
  - low eg. on nerve cells, on myocytes
- Consists of alpha chain and β2 microglobulin.

#### HLA class II.

- HLA-DR, DQ, DP genes
- Present on the surface of the cells of the immune system (professional antigen presenting cells):
  - B cells
  - dendritic cells
  - macrophages
- Can be induced on endothel.
- Consists of alpha and beta chains.



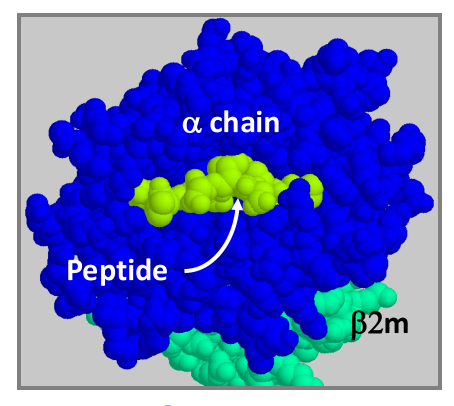


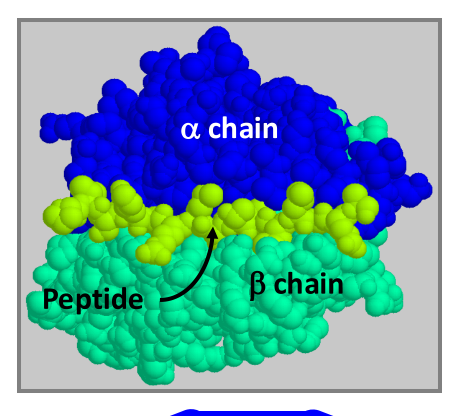
Peptides from endogenous proteins (own, virus, intracellular bacteria, tumour)

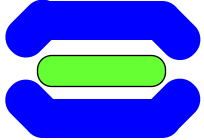
Peptides from exogenous proteins (extracellular pathogen, allergen)

#### HLA class I.

#### HLA class II.





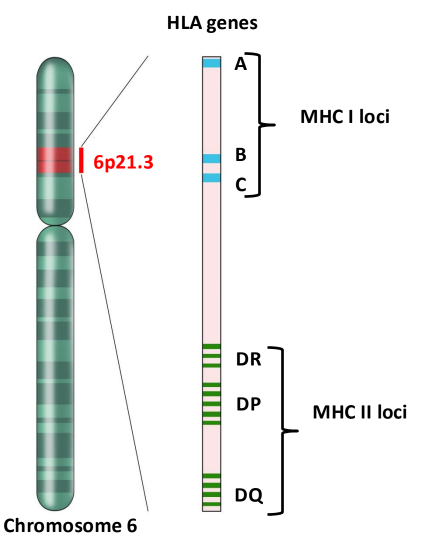




Binds peptides of 8-10 amino acids

Binds peptides of >13 amino acids

#### Inheritance of HLAs: MHC loci



In humans both HLA-A, B, C, DQ, DR, DP and DP are expressed simultaneously both from the maternal and paternal chromosomes.  $\rightarrow$  Many types of MHC are present on the cells.

#### Basics of HLA nomenclature

- MHC I has only 1 chain, e.g.: HLA-B\*27:01 → B type MHC I of the 27 serological group
- The MHC II is a heterodimer with two chains.

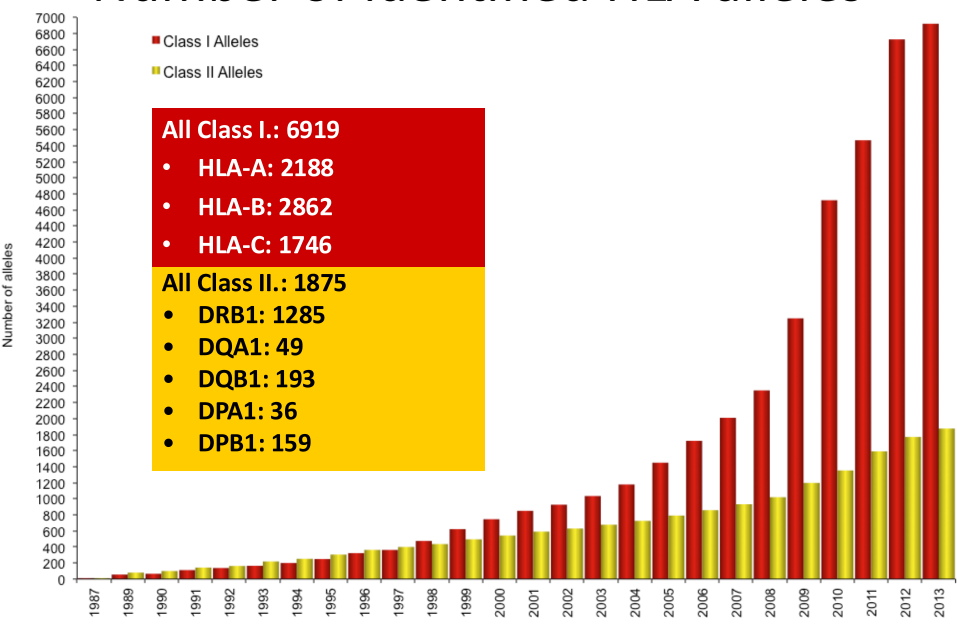
#### HLA-DQA1:05:01

- 1. What type of MHC does it encode? E.g.: A, B, C, DQ, DR, DP
- 2. Locus (A1 = encodes alpha chain, B1= encodes beta chain)

- 3. Into which serological group does the chain belong? (05 = alleles resulting a  $\alpha^5$  chain)
  - 4. The specific allele in the group

**Attention!** Slide is solely for illustration, we will not ask HLA nomenclature.

#### Number of identified HLA alleles



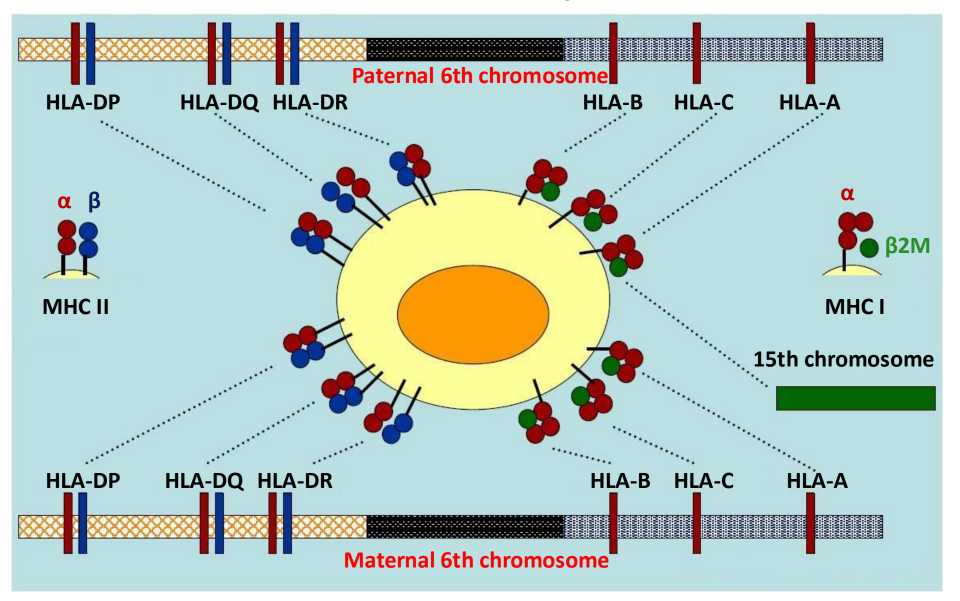
Year

#### Inheritance of HLAs

- Polygenic: Several genes encode MHC I and MHC II molecules. (e.g. HLA-A, B and C type I MHCs and DP, DQ and DR type II MHCs)
- Polymorphic: There are many different alleles of each gene in the population, therefore it is highly variable.
- Codominant: Both the maternal and the paternal alleles are expressed in an individual.

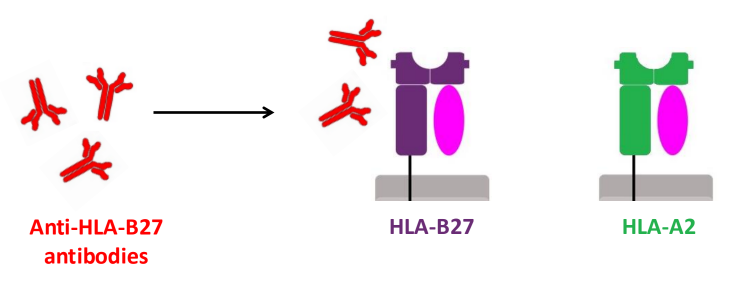
Each person has a characteristic **MHC pool** with different peptide-binding capabilities.

## Codominant expression



## HLA serotype vs genotype

HLA serotype: MHC molecules are distinguished by their different antigenicity. E.g.:



HLA genotype: MHC molecules are distinguished by identifying the encoding HLA alleles. A different genotype will not necessarily lead to a different serotype, there are more alleles than HLA serotypes. (6959 HLA alleles were known in 2010 but it is increasing each year. → Polymorphism)

## Methods of HLA typing

- Serological methods:
  - Microcytotoxicity assay (MCA)
  - Mixed lymphocyte culture(MLC)
- Molecular biological methods: (→ see from molecular cell biology)
  - Restriction fragment length polymorphism (RFLP)
  - Sequence-specific oligonucleotide probes (SSOP) → DNA hybridization
  - Sequence-specific primers → SSP-PCR
  - DNA sequencing

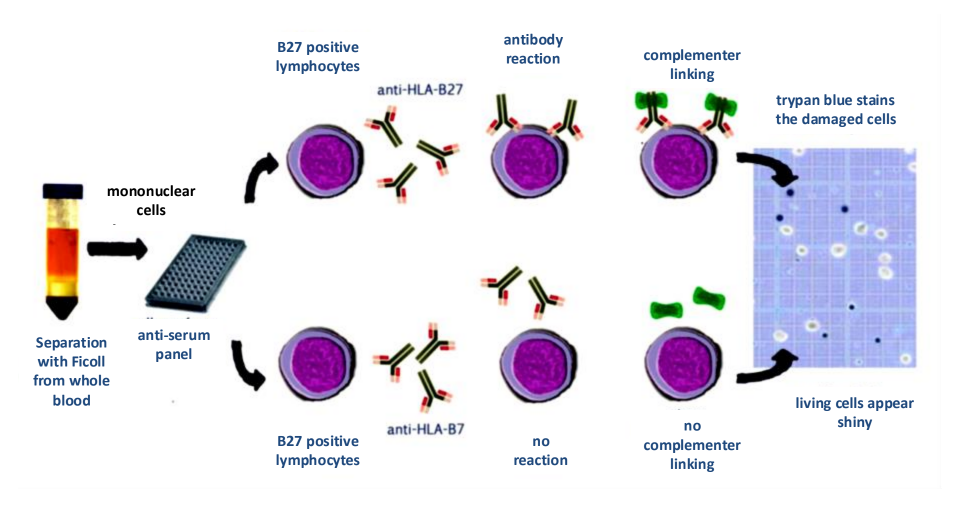
#### The molecular biological methods are the preferred ones because they are:

- More specific (well-defined probes and primers are used)
- More flexible (new oligonucleotide probes or primer can be designed as soon as a new allele described)
- More reliable (doesn't require a specific cell type and is less dependent on the condition of the patients)

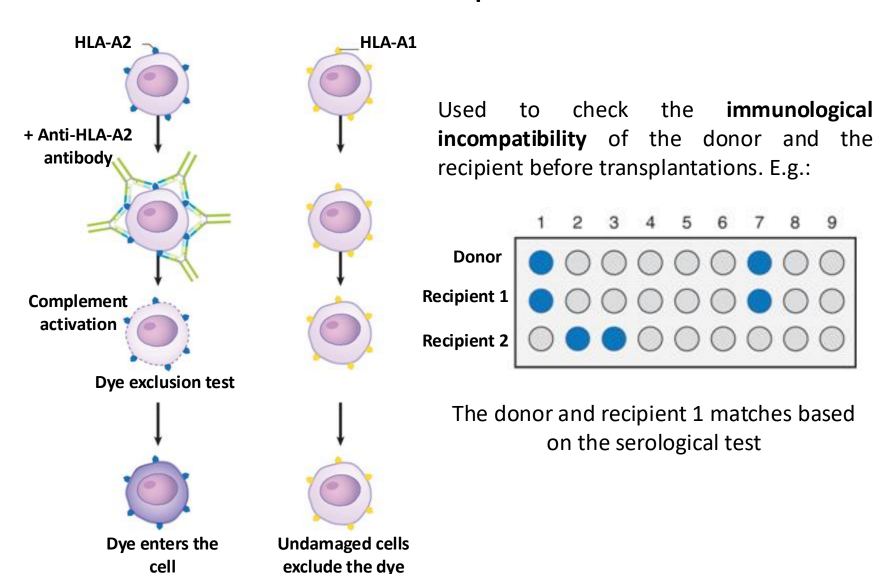
## **HLA** serology

- **1. Typing** investigation of polymorphism of HLA antigens: lymphocytes with unknown antigenicity added to known specificity antibodies.
- 2. Antibody screening PRA (panel reactive antibody) test detection of antibodies against HLA antigens: the sera that need to be tested is added to known antigenicity cell panel.
- 3. Cross match: in vitro model of antigen-antibody reaction that might occur during transplantation, might result in graft rejection in vivo.

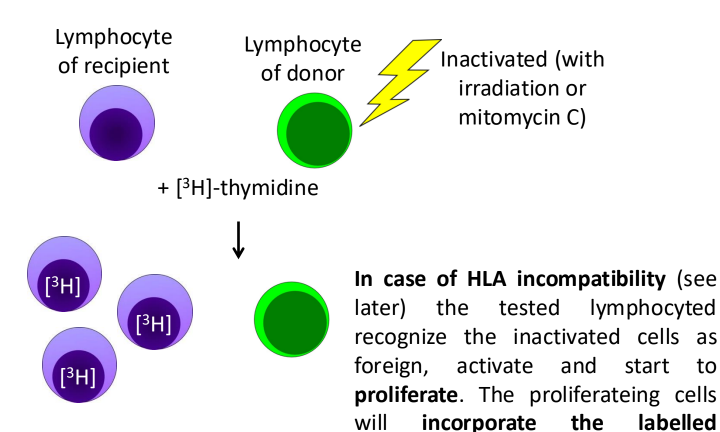
## Microcytotoxicity assay (Terasaki)



#### Microcytotoxicity assay (MCA) – Cross match Terasaki plate



## Mixed lymphocyte culture



lymphocyted

start

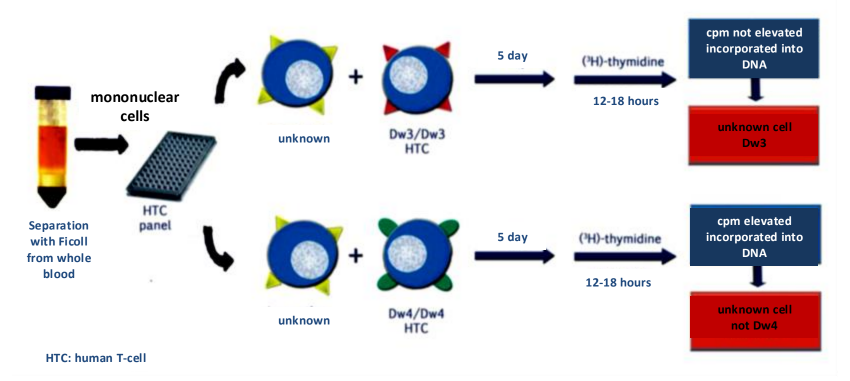
thymidine into their DNA.

labelled

#### Application:

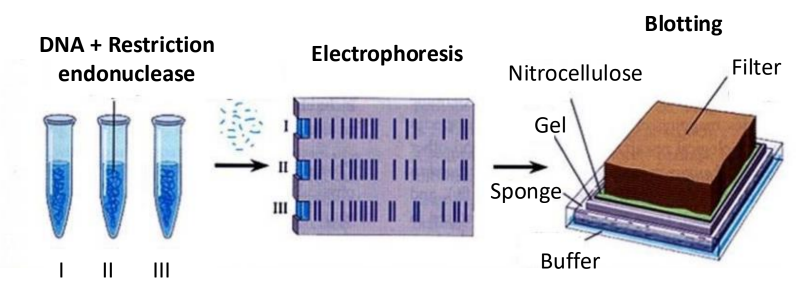
To check the **immunological incompatibility** of the donor and the recipient before transplantations.

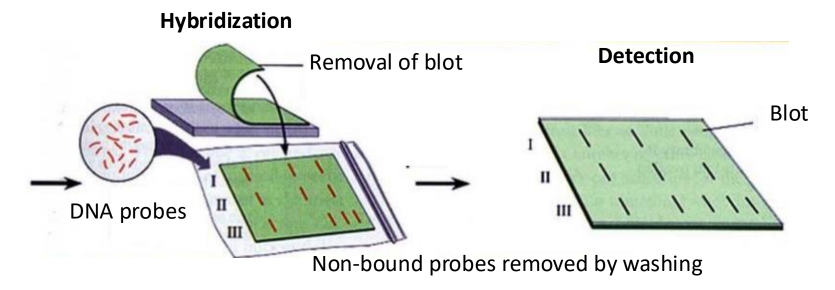
## Mixed lymphocyte culture (MLC)



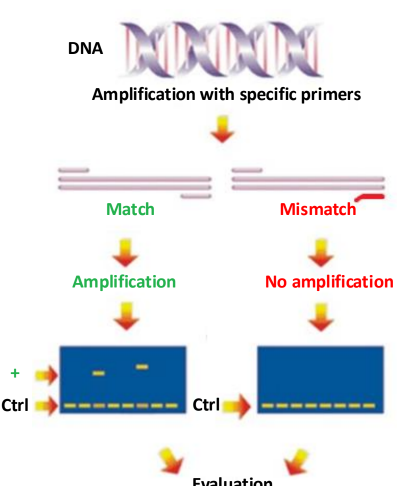
<u>Mixed Lymphocyte Culture (MLC)</u>: The lymphocytes of two individuals (eg. a patient's unknown lymphocyte and a laboratory sustained known lymphocyte expressing HLA-Dw) are mixed in a Petri dish and incubated in cell culture for days. If the unknown (patient's) lymphocyte doesn't carry HLA-Dw, which can be found on the test lymphocyte, then the lymphocyte becomes stimulated and proliferates, which can be measured by the thymidine incorporation. *Used solely to determine MHCII*, it has clinical significance in case of transplantation. Requires tissue culture laboratories that sustain homogeneous cell population. Isotope test.

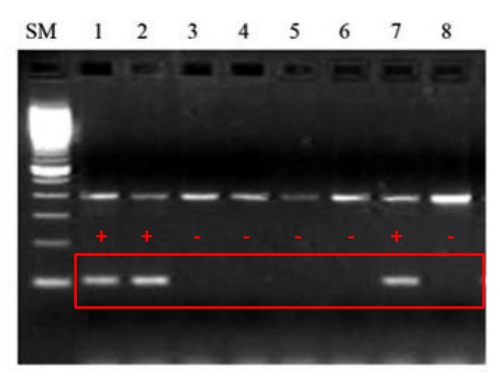
RFLP (Restriction fragment length polymorphism)





#### PCR with sequence-specific primers





HLA-A\*01 genotyping: The visible bands in samples 1, 2 and 7 mark HLA-A\*01 alleles.



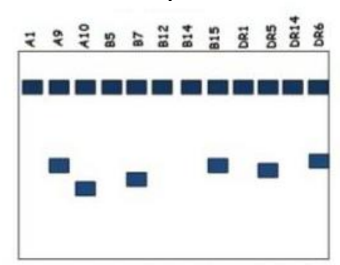
## Steps of organ transplantation alert

- The blood sample of the donor arrives
- Selection of possible recipients from the registry
- DNA isolation, SSP PCR for HLA-A, B and DRB1 genes, gel electrophoresis, evaluation
- Cross match → testing of donor's cells with the sera of the possible recipients in the presence of complement. Checking the recipient's sensitization

## HLA matching before kidney transplantations

The survival of the graft is mainly determined by the degree of matching of the **HLA-A**, **HLA-B** and **HLA-DR** alleles, these are the one investigated before transplantations.

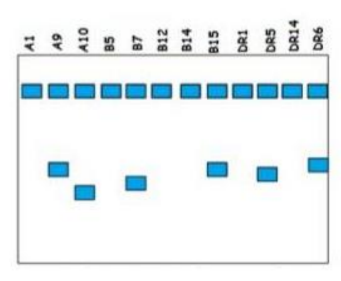
#### Recipient



#### **HLA** genotype:

- A9, A10
- B7, B15
- DR5, DR6

#### **Donor**



#### **HLA** genotype:

- A9, A10
- B7, B15
- DR5, DR6



## Graft versus host disease 1. (GVHD)

- May develop after allogeneic hematopoietic stem cell transplantations (HSCT).
- The donor-derived immune cells attack and damage the host tissues.
- Main risk factor: HLA mismatch between the donor and the recipient.
- Therapy: Steroids (immunosuppression), lethality is roughly 15%, but steroid resistant acute GVHD has a lethality of 90%.

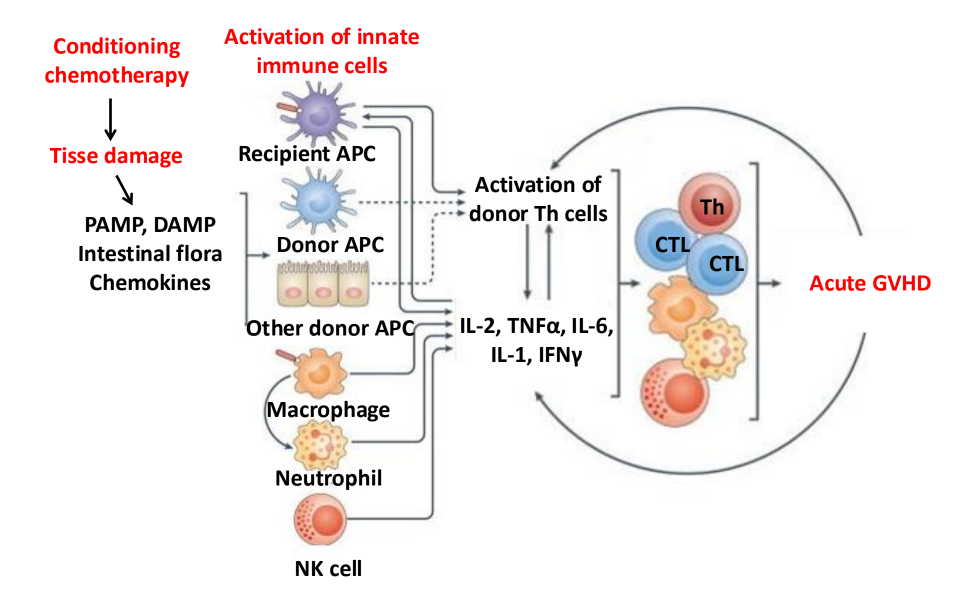


Severe skin GVHD



Acute intestinal GVHD (endoscopic image)

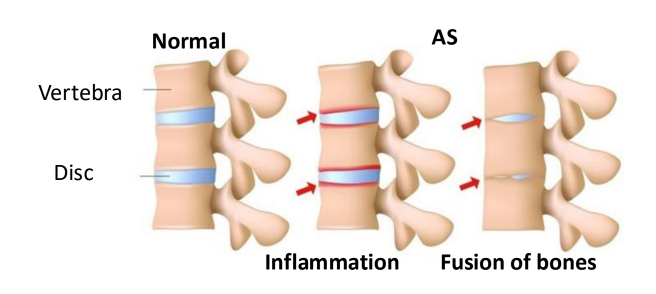
## Graft versus host disease 2. (GVHD)



#### HLA and disease associations 1.

- Bechterew's disease (Ankylosing spondylitis, AS): HLA-B27
- Approx. 90% of AS patient are HLA-B27 positives.
- Prevalence of HLA-B27 in the Caucasian race is 8%, in Scandinavia it reaches 24%.
- Approx. 1,8% of HLA-B27 positive individuals develop manifest AS.

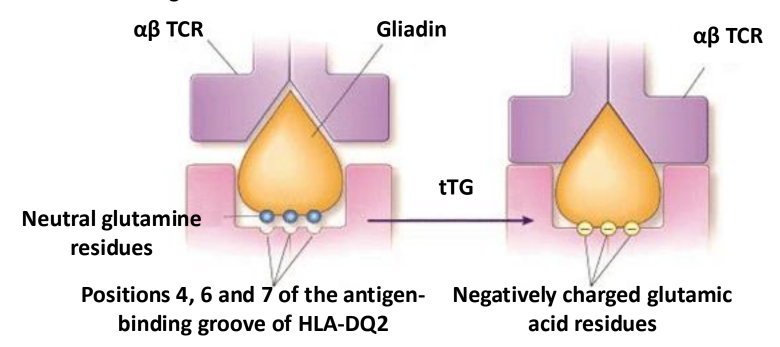
HLA-B27 positivity **only increases the risk of the disease**, it is **not enough on its own** for developing the disease! (This is also true for all the HLA associations.)





#### HLA and disease associations 2.

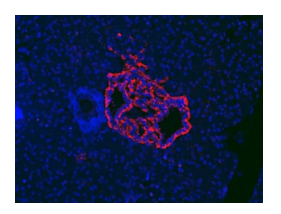
- Celiac disease (gluten-sensitive enteropathy): HLA-DQ2 and HLA-DQ8
- At least one of them is present in roughly **98%** of the patients. (Strongest known HLA association and the most well-understood role in the pathogenesis)
- Prevalence of HLA-DQ2 in the Caucasian race is 30%, but the prevalence of the disease is only 1%. → Positivity is not enough for developing the disease.
- These show a **higher affinity towards gliadin** than other MHC types, especially when binding the deamidated forms.



#### HLA and disease associations 3.

- Type I diabetes mellitus (IDDM): HLA-DR3, HLA-DR4
- HLA-DR3-DQ2  $\rightarrow$  3X risk
- HLA-DR4-DQ8  $\rightarrow$  10X risk
- HLA-DR3-DR4 heterozygotes → 25X risk
- HLA-DQ6.2 → 0,1X risk (protective role)





Direct IF: Human Langerhans islet (patient with IDDM)

**Green: Insulin (lacking)** 

Red: glucagon Blue: cell nuclei

#### Student Research

- Main fields of research at the department:
  - Differentiation of lymphoid tissues and recirculation of immune cells
  - Role of T cells in the murine model of rheumatoid arthritis
  - Signaling and role of regulatory T cells in autoimmune disease (mainly SSc)
  - Effects of glucocorticoids on T cells
  - Evolution of the immune system in invertebrate animal models
  - Efficacy monitoring of vaccines by immunoserological techniques

The exact topics can be found on the link below, for further info please contact the supervisors. Own topics are also welcomed if they are related to immunology.

http://aok.pte.hu/en/egyseg/tdk\_temak/120



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