Basic Immunology

T-cell development in the thymus. Stages of maturation and the role of environmental factors.

Structure of the thymus



Structure of the thymus

Figure 5.3



The thymic stoma creates the microenvironment that is essential for T-cell development

T-cell source: thymus and beyond.

- Thymus: prime T-cell source
- Absence of thymus nude mouse (FoXN1)
 Ectodermal (skin) and endodermal developmental abnormalities. Profoundly reduced T-cell pool.
- <u>DiGeorge's syndrome</u>: Complex cardiac, facial, endocrine and immune defects

- Extrathymic T-cell source: intestinal "cryptopatch" (in LP 1000-5000 c-kit⁺, CD44⁺, IL7R⁺, CD25⁻ phenotype cells), maturation in MLN and PPs (based on RAG-expression).
- Second thymus





T cell repertoire

Total repertoire: TCR α , β : 10¹⁵ TCR γ , δ : 10¹⁶

T cell precursors are produced in the **<u>bone marrow</u>** from the common haemopoietic stem cell They migrate through the blood circulation to the thymus

<u>In the thymus</u>: T cell maturation, educational steps ,,double recognition" (MHC and peptide)

<u>Periphery</u>: mature, TCR expressing, CD4 or CD8 positive T cells

Self-MHC restricted

Self-tolerant T cells

The 'clonal selection theory' first proposed by Frank Macfarlane Burnet in 1956 predicts that each lymphocyte expresses a single, unique, antigen-specific receptor.





Antigen receptors: BCR és TCR



T cell receptor complex on mature T cells



αβ TcR – SP(CD4+ or CD8+) γδ TcR – DN (CD4-CD8-)

TCR protein chains and their CDR regions

CDR3



TCR β, α, δ γ loci



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V-D-J combinatorial and junctional diversity of TCR

	α	β	
Number of V Gene Segments	45	50	
Number of diversity (D) gene segments	0	2	
Number of joining (J) gene segments	~50	12	
Combinational Diversity Number of Possible V-(D)-J Combinations	Vn D2 J2 C 		
Junctional Diversity Total potential repertoire with junctional diversity	V1 D1 J1 C Addition of Nucleotides TCR: ~10 ¹⁶		

TCR diversity



NOTCH1 is required for the commitment of multipotent hematopoietic progenitors to the T-cell lineage



NOTCH1 support cell growth, proliferation and survival at multiple stages of thymocyte developmentis involved in the progression through the early DN1, DN2 and DN3 stages of thymocyte development and in the regulation of TCRB rearrangement

In: Transdifferentiation and regenerative medicine (Dr. Péter Balogh, Dr. Péter Engelmann (2011); University of Pécs)

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				DP-			
Stage of maturation	Stem cell	Pro-T	Pre-T	Double positive	Single positive (immature T cell)	Naive mature T cell	
Proliferation				1			
RAG expression							
TdT expression							
TCR DNA, RNA	Unrecombined (germline) DNA	Unrecombined (germline) DNA	Recombined β chain gene [V(D)J-C]; β chain mRNA	Recombined β, α chain genes [V(D)J-C]; β and α chain mRNA	Recombined β, α chain genes d [V(D)J-C]; β and α chain mRNA	Recombined β, α chain genes [V(D)J-C]; β and α chain mRNA	
TCR expression	None	None	Pre-T receptor (β chain/pre-T α)	Membrane αβ TCR	Membrane αβ TCR	Membrane αβ TCR	
Surface markers	c- <i>kit</i> + CD44+ CD25-	c- <i>kit</i> + CD44+ CD25+	c- <i>kit</i> + CD44+ CD25+	CD4+CD8+ TCR/CD3 ^{lo}	CD4+CD8 ⁻ or CD4-CD8+ TCR/CD3 ^{hi}	CD4+CD8 ⁻ or CD4-CD8+ TCR/CD3 ^{hi}	
Anatomic site	Bone marrow		Thy	mus		Periphery	
Response to antigen	None	None	None	Positive and negative selection	Negative selection	Activation (proliferation and differentiation)	
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Thymocyte populations based on their cell surface markers



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T-cell development in the thymus



Young mouse: 5x10⁷ T-cells daily During selection 98 % of thymocytes die by apoptosis 108

Daily 1-2 x 10⁶ mature T-cell migrate to the periphery

Thymocytes traffic through various regions of the thymus as they undergo positive and negative selection.



Thymic Microenvironment and T-cell Development

1. <u>Migration:</u> Chemokines

2. Proliferation IL-7

3. <u>Differentiation</u>
•TcRrearragement
•Phenotypes

4. <u>Selection</u> Apoptosis



The thymus selects for self-reactive T cells



Decision-making during the development of T cells



$\gamma\delta$ -and $\alpha\beta T$ cell maturation



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Structure and role of preTCR





Positive selection:

Epithelial cell - thymocyte interaction in the thymus cortex

Survival of DP cells whose TcR is appropriate for self MHC recognition

Negative selection:

APC (macropahge or DC) – thymocyte Interaction in thymus medulla

Death of DP cells with high affinity TcR for self MHC + self peptide recognition

Differentiation into SP stage

AIRE



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MHC recognition determines T cell lineage and function.



Role of mTEC in negative selection



mTEC = medullary thymic epithelial cell AIRE = autoimmune regulator transcription factor TRA = tissue related antigens

AIRE: autoimmune regulator

- AIRE is a transcription factor expressed in the medulla (inner part) of the thymus and controls a mechanism that prevents the immune system from attacking the body.
- In the thymus, the AIRE causes transcription of a wide selection of organ-specific genes. These self antigen reactive T cells that bind strongly to self-<u>antigen</u> are eliminated in the thymus in the negative selection.
- The AIRE gene is mutated in the rare autoimmune syndrome Autoimmune Polyendocrinopathy Syndrome type 1 (<u>APS-1</u>), also known as Autoimmune Polyendocrinopathy-Candidiasis-Ectodermal Dystrophy (<u>APECED</u>). Disruption of *AIRE* results in the development of a range of autoimmune diseases,

Clonal deletion and regulatory T cell selection are the major thymic tolerance





Affinity model of Tcell selection



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The affinity model of thymocyte selection centres on the strength of the interaction of the T cell receptor (TCR) with self-peptide–MHC complexes as a crucial determinant of cell fate. Weak interactions are required to protect thymocytes from death by neglect and to promote the positive selection of naive T cells. Strong interactions cause negative selection by apoptosis

Phases of T-cell maturation in the thymus.

- 1. Initiation of either TCR β or γ/δ chain gene rearrangement.
- 2. Formation of pT α /TCR β /CD3 (pTCR), allelic exclusion, IL-7-dependent proliferation β -selection.
- 3. Initiation of TCR α gene rearrangement.
- 4. Completion of TCR α/β gene rearrangement, co-expression of CD4/CD8 molecules.
- 5. Recognition of MHC/peptide complexes displayed by thymic cortical epithelium *positive selection*.
- 6. Binding to MHC/peptide complex displayed by thymic APC/medullary epithelial cells *negative selection*.
- Influence of stronger/more persistent signal: commitment towards CD4 or Treg (CD4/CD25+) subset.

Malignant haematopoietic diseases originated from immature cells of lymphoid cell lineage



Figure 7-43 Immunobiology, 7ed. (© Garland Science 2008)