M1 - Immunology, Winter 2008

Fantone, J.; Pietropaolo, M. T.

<http://hdl.handle.net/2027.42/64939>
http://hdl.handle.net/2027.42/64939
Unless otherwise noted, the content of this course material is licensed under a Creative Commons Attribution - Non-Commercial - Share Alike 3.0 License.

Copyright 2008, Massimo T. Pietropaolo.

The following information is intended to inform and educate and is not a tool for self-diagnosis or a replacement for medical evaluation, advice, diagnosis or treatment by a healthcare professional. You should speak to your physician or make an appointment to be seen if you have questions or concerns about this information or your medical condition. You assume all responsibility for use and potential liability associated with any use of the material.

Material contains copyrighted content, used in accordance with U.S. law. Copyright holders of content included in this material should contact open.michigan@umich.edu with any questions, corrections, or clarifications regarding the use of content. The Regents of the University of Michigan do not license the use of third party content posted to this site unless such a license is specifically granted in connection with particular content objects. Users of content are responsible for their compliance with applicable law. Mention of specific products in this recording solely represents the opinion of the speaker and does not represent an endorsement by the University of Michigan.

Viewer discretion advised: Material may contain medical images that may be disturbing to some viewers.
The objectives of this lecture are to understand:

1. The Pathogenesis of Autoimmune Diabetes (Type 1A diabetes)
2. The role of T cells in Disease Pathogenesis
3. The role of Cytokines in Disease Pathogenesis
4. The role of Islet Autoantibodies
Diabetes Mellitus

A systemic disease with multiple metabolic abnormalities, chief among which is an elevation in plasma glucose.

In addition to the primary defect in carbohydrate metabolism defects in lipid metabolism are widespread, with elevations in plasma FFA and TG, and, in some circumstances, of ketones.
The Expert Committee on the Diagnosis and Classification of Diabetes Mellitus

I. Type 1 diabetes
   A. Immune mediated
   B. Idiopathic

II. Type 2 diabetes

III. Other specific types
Regulation of Plasma Glucose

Glucose Disposal

Peripheral Tissues

Liver

Steady State Plasma Glucose

Glucose Production

β Cells
Regulation of Plasma Glucose

Glucose Production

Liver

Glucose Disposal

Peripheral Tissues

Steady State Plasma Glucose

Insulin
Diabetes Mellitus- Type 1

Increased thirst (polydypsia)
Increased urination (polyuria)
Increased appetite (polyphagia)
Weight loss
Fatigue
Rapid, early onset (before age 15)
<table>
<thead>
<tr>
<th>Difference</th>
<th>Type 1</th>
<th>Type 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age of onset</td>
<td>Young</td>
<td>Older</td>
</tr>
<tr>
<td>Type of onset</td>
<td>Acute</td>
<td>Insidious</td>
</tr>
<tr>
<td>Genetic background</td>
<td>HLA related</td>
<td>Not HLA related</td>
</tr>
<tr>
<td>Islet cell antibodies</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Insulin secretion</td>
<td>Absent</td>
<td>Present</td>
</tr>
<tr>
<td>Nutritional status</td>
<td>Thin</td>
<td>Obese</td>
</tr>
<tr>
<td>Insulin dependence</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Insulin resistance</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Responsiveness to Orals</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Ketosis proneness</td>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>
Relative Proportions of Types 1 & 2 DM

95% Type 2
5% Type 1
Type 1 diabetes (IDDM)

Chronic autoimmune disease with juvenile onset, but may develop in adults as well as elderly (LADA).

**Polygenic disease**
- Strong MHC linkage
- Non-MHC genes

**Autoimmune etiology**
- Antibodies to islet autoantigens
- Autoreactive T cells

**Immune-modulation alters the course of disease**
- Antigen vaccination
- General immunosuppression
Stages in Development of Type 1 Diabetes

1. GENETICALLY AT RISK
2. MULTIPLE ANTIBODY POSITIVE
3. LOSS OF FIRST PHASE INSULIN RESPONSE
4. "PRE"-DIABETES
5. DIABETES

Genetic Predisposition
Insulitis
Beta Cell Injury
"Pre"-Diabetes
Diabetes

G. Eisenbarth, NEJM, 1986
Type 1 diabetes: a chronic inflammatory disease of the islets

Genetic Susceptibility
Empiric risk of developing Type 1 diabetes

<table>
<thead>
<tr>
<th>Empiric Risk</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>First degree relatives of T1DM</td>
<td>5-7%</td>
</tr>
<tr>
<td>probands*</td>
<td></td>
</tr>
<tr>
<td>Individuals without relatives with T1DM*</td>
<td>&lt;1%</td>
</tr>
<tr>
<td>Children of affected father**</td>
<td>~6%</td>
</tr>
<tr>
<td>Children of affected mother**</td>
<td>~2%</td>
</tr>
</tbody>
</table>

These estimates are for North American Caucasian* and Scandinavian populations**
The Wellcome Trust Case Control Consortium (WTCCC) primary genome-wide association (GWA) scan in T1DM

HLA

Human Leukocyte Antigen

human MHC

cell-surface proteins

important in self vs. nonself distinction

present peptide antigens to T cells

CLASS I: A,B,C

CLASS II: DR,DQ,DP
The Human Leukocyte Antigen Complex (6p21.31)

Class II (1.1 Mb)  
DP  DQ  DR

Class III (0.7Mb)  
Complement and Cytokines

Class I (2.2Mb)  
B  C  A

Centromere

Frequent Recombination

Telomere

Recombination is Rare

Recombination is Rare

Class I-like genes and pseudogenes

Complement and Cytokines
MHC Haplotype Sharing Increases DR3/4 Sibling Risk

Haplotype Determination:

**Siblings Share Both Haplotypes**
- **Family A**

**Siblings Share One Haplotype**
- **Family B**

**Siblings Share No Haplotype**
- **Family C**
MHC haplotype sharing increases risk in DR3/4-DQ8 siblings

% Autoantibody Positive
- Share 2
- Share 0 or 1

% Diabetic
- Share 2
- Share 0 or 1

Source: Aly T et al. PNAS, 2006
Multiple Factors May Drive Progressive Decline of $\beta$-Cell Function

- $\uparrow$ Apoptosis/Necrosis
- $\downarrow$ Islet Neogenesis
- Islet Autoantibodies
- Hyperglycemia (glucose toxicity)
- Environmental Factors

Autoreactive T cells

(Elevated cytokines IFN$\gamma$, IL-1$\beta$, TNF$\alpha$, etc.)
Environmental Factors
Congenital Rubella Syndrome

- 30% diabetic usually early T1DM, some T2DM
- incubation period 5-20 yrs
- HLA-DR3 or 3/4 in those with diabetes
- other autoimmune diseases (thyroid, AD)
- molecular mimicry with a 52kD autoantigen
- animal model - Syrian hamsters
- No diabetes after postnatal infection or MMR vaccination
Other Environmental factors involved in Type 1 diabetes pathogenesis

- Cocksakie B Virus? Molecular mimicry with the islet autoantigen glutamic acid decarboxylase (GAD)
- Enterovirus?
- Streptozotocin (low doses)?
Loss of self tolerance to self-antigens
Autoantigens in Diabetes

- Insulin
- Glutamic acid decarboxylase (GAD65)
- Islet autoantigen 512aa (ICA512/IA-2)
- Zinc Transporter Znt8
Is there a primary antigen or immune response to multiple antigens required for autoimmunity?

T cells specific for one antigen (insulin)

Insulitis

Epitope and antigen spreading, expansion

Diabetes

OR

T cells specific for multiple antigens

Insulitis

Expansion of T cells

Diabetes

Krishnamurthy et al  JCI:116:3258, 2006
Role of T cells
Pathogenic Cells in Type 1 diabetes

Cell-mediated Immunity

- CD4+ T cells-MHC class II molecules (APC) interaction
- CD8+ T cells-MHC class I molecules (APC) interaction
- NK cells
- Macrophages
- Dendritic cells
INSULITUS. PATIENT DIED FROM DKA
Type 1 diabetes pathogenesis: alteration between pathogenicity (T effector cells) and regulation (regulatory T cells)
Example of regulatory T cell defect: X-linked autoimmunity-immunodeficiency syndrome (XLAAD)

Gene defect: **FOXP3**

*This genetic defect can lead to Type 1 diabetes in the presence of other autoimmune disorders for abnormalities in regulatory T cell maturation.*
Regulatory T cells (Tregs)

Thymus

- Naïve CD4⁺
- CD4⁺CD25⁺ FoxP3⁺

Periphery

- DC
- TGF-β1
- IL-4
- IL-12
- IL-6
- IL-23
- TGF-β1

- Th2 → IL-4
- Th1 → IFN-γ
- Th17 → IL-17
- Treg → TGF-β1 others
- FoxP3
Role of cytokines
Differentiation of CD4+ T-cell Subsets

IL-12

Tn

IL-10

Tr1

IFN-α

Th1

IL-4

CD4+ T-cell

IL-10

IL-10

TGF-β

IFN-γ

IL-2

Cell-Mediated Immunity
Autoimmunity
Pro-Inflammation
Allograft Rejection

Suppressor/Regulatory

Humoral Immunity
Anti-Inflammatory
Oxidative Stress

IL-β, IFN-γ, TNF-α → Oxidative Stress

Nitric Oxide (NO) production

β-cell death
Role of autoantibodies
Cytoplasmic islet-cell-antibody staining

Positive reaction

Negative reaction


**Islet Cell Autoantibody Assays**

**GAD65 Autoantibodies**

**IA-2 Autoantibodies**

**Insulin Autoantibodies (IAA)**
New Radioimmunoassay [CV: inter-assay: 19.4%; intra-assay: 8%]

**Islet Cell Antibodies (ICA)**
Immunoperoxidase staining in rat and human pancreas
Prospective Studies in First Degree Relatives of T1DM Probands

Sibling/offspring cohort
Cumulative risk of developing clinical Type 1 diabetes in relatives of T1DM patients using islet autoantibodies (IAA, GAD65, IA-2, ICA)

Log Rank

P < 0.00001
Objective: To determine whether any immunomodulatory therapy can ameliorate insulin secretion in newly diagnosed T1DM (17-40 yr of age) and to ultimately prevent T1DM onset in first-degree relatives of T1DM probands. First trials in relatives started in 2003.

Criteria for enrolling T1DM patients in TrialNet: ≥ 2Ab to islet antigens.
Conclusions

• Type 1 diabetes mellitus is a polygenic disease. Although at least 19 T1DM-related candidate genes have been identified, polymorphic regions within the HLA complex confers the strongest diabetogenic effect.

• CD4+ and CD8+ T cell responses to islet autoantigens (insulin, GAD65 and IA-2) are pathogenic.

• A defect of Regulatory T cells in suppressing pathogenic autoimmune responses is associated with Type 1 diabetes.

• The proinflammatory cytokines IL-1\(\beta\), IFN-\(\gamma\) and TNF-\(\alpha\) can cause \(\beta\) cell death (increased NO production).

• Gene defects in FOXP3 and AIRE cause multiple autoimmune disease (APECED, APS-I respectively) including Type 1 diabetes

• The presence of multiple autoantibodies to insulin, GAD65, IA-2 are high risk markers of Type 1 diabetes progression.