Basic Immunology

Lecture 21.

Allergies and hypersensitive reactions

Cellular and molecular mechanism. T cell mediated macrophage activation = Type IV. hypersensitive reaction (DTH).

Hypersensitive reactions

- Pathological overreactions of the immune response with severe tissue damage (necrosis) in the effector phase.
- The immune system itself initiates these diseases.
- Different background mechanisms.
- Gell and Coombs divided 4 types of reactions.

Based on the immunological mechanisms we distinguish 4 types of hypersensitive reactions

Immunoglobulin-mediated

Type I.Atopy or Allergy(IgE-mediated immediate form)

- Type II.Humoral cytotoxic immune reactions(IgG against cellular antigens)
- Type III. Immuncomplex-diseases (soluble self or non-self antigens)

Cell-mediated

Type IV. T cell-mediated \rightarrow Th1- and Tc- cytokines (DTH=**D**elayed **T**ype **H**ypersensitivity)

	Туре І	Ту	pe II	Type III
Immune reactant	lgE	IgG		lgG
Antigen	Soluble antigen	Cell- or matrix- associated antigen	Cell-surface receptor	Soluble antigen
Effector mechanism	Mast-cell activation	Complement, FcR ⁺ cells (phagocytes, NK cells)	Antibody alters signaling	Complement, Phagocytes
	Ag	platelets complement		immune complex blood vessel complement
Example of hypersensitivity reaction	Allergic rhinitis, asthma, systemic anaphylaxis	Some drug allergies (eg, penicillin)	Chronic urticaria (antibody against FC∈R1α)	Serum sickness, Arthus reaction

Figure 12-2 part 1 of 2 Immunobiology, 6/e. (© Garland Science 2005)

		Туре IV	
Immune reactant	T _H 1 cells	T _H 2 cells	CTL
Antigen	Soluble antigen	Soluble antigen	Cell-associated antigen
Effector mechanism	Macrophage activation	IgE production, Eosinophil activation, Mastocytosis	Cytotoxicity
	IFN-γ T _H 1	IL-4 IL-5 ↓ eotaxin	CTL ₽
	chemokines, cytokines, cytotoxins	cytotoxins, inflammatory mediators	
Example of hypersensitivity reaction	Contact dermatitis, tuberculin reaction	Chronic asthma, chronic allergic rhinitis	Contact dermatitis

Figure 12-2 part 2 of 2 Immunobiology, 6/e. (© Garland Science 2005)



Type I., immediate hypersensitivity; Allergy, Atopy

Basic mechanism



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Allergens

Figure 10.1a



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Food antigens (milk, soy, gluten, nuts, additives etc.)

Most important characteristics of inhaled allergens which enhance IgE production through Th2 activation

Proteins	only proteins elicit T cell response
Enzime activity	often proteases
Low dose	enhance activation of IL-4-producing CD4- Th2 cells
Low molecular weight	the allergen can easily diffuse from the particle into the mucus.
Good solubility	the allergen can be released easily from the particle
Stabile	the allergen can be released even from exsiccated particles
Contain peptides that are able to bind to self MHCII	important at the first exposure for T cell activation



IgE-Receptors



Fcε-Receptor signaling



Gilfillan et al. Nature Reviews Immunology 6, 218-230 (March 2006) | doi:10.1038/nri1782



Mechanism of Type I. hypersensitivity



Degranulation of mast cells



Pharmacologic Mediators of Immediate Hypersensitivity

Preformed mediators in granules

histamine	bronchoconstriction, mucus secretion, vasodilatation, vascular permeability			
tryptase	proteolysis			
kininogenase	kinins and vasodilatation, vascular permeability, edema			
ECF-A (tetrapeptides)	attract eosinophil and neutrophils			
Newly formed mediators				
leukotriene B ₄	basophil attractant			
leukotriene C ₄ , D ₄	same as histamine but 1000x more potent			
prostaglandins D ₂	edema and pain			
PAF	platelet aggregation and heparin release:			

microthrombi

Antigen-IgE binding enhances IgE production



Figure 12-7 Immunobiology, 6/e. (© Garland Science 2005)

Late phase

Upon cytokine effect: recruitment of neutrophils and eosinophils, stimulation of B cells

IL-3, IL-5, GM-CSF \rightarrow local eosinophil proliferation \rightarrow Inflammation

Physiological role of the IgE response in the protection against parazites and fungi



Type I. diseases

- Systemic anaphylaxia anaphylactic sock
- Allergic rhinitis (=Hay fever)
- Allergic conjunctivitis
- Allergic asthma
- Urticaria
- Ekzema (atopic dermatitis)



Allergy – Environmental factors

Atopic allergy and asthma is the most frequent in the economically well-developed countries.

- changes in the infectious diseases in early childhood ("Hygiene-theory" / "Old Friends Hypothesis")
- Environmental pollution (air pollution in industrial regions, traffic)
- Altered allergen concentrations
- Changes in the diet (chemicals)
- Changes in the gut microbiota



Hygiene-theory



No data available

Not applicable

No preventive chemotherapy required

>2/3

<1/3

Hygiene-theory



Figure 1. Inverse Relation between the Incidence of Prototypical Infectious Diseases (Panel A) and the Incidence of Immune Disorders (Panel B) from 1950 to 2000.

Hygiene-theory

Environment

'Developing' countries Large family size Rural homes, livestock Intestinal microflora-variable, transient Low antibiotic use High helminth burden Poor sanitation, high orofaecal burden

Non-allergic

'Westernized' countries Small family size Affluent, urban homes Intestinal microflora-stable High antibiotic use Low or absent helminth burden Good sanitation, low orofaecal burden

> Allergic disorders (asthma, eczema and rhinitis)

Nature Reviews | Immunology

In: Marsha Wills-Karp, Joanna Santeliz & Christopher L. Karp: <u>The germless theory of allergic disease: revisiting the hygiene</u> <u>hypothesis.</u> *Nature Reviews Immunology* **1**, 69-75 (October 2001)doi:10.1038/35095579

Genes

Old Friends hypothesis

"Old Friends"=Organisms such as helminths and environmental saprophytes, that are part of mammalian evolutionary history.



In: Review series on helminths, immune modulation and the hygiene hypothesis: The broader implications of the hygiene hypothesis. Immunology, Volume 126, Issue 1, pages 3-11, 8 DEC 2008 DOI: 10.1111/j.1365-2567.2008.03007.x http://onlinelibrary.wiley.com/doi/10.1111/j.1365-2567.2008.03007.x/full#f2



In: Review series on helminths, immune modulation and the hygiene hypothesis: The broader implications of the hygiene hypothesis. Immunology, <u>Volume 126, Issue 1, pages 3-11, 8 DEC 2008 DOI: 10.1111/j.1365-2567.2008.03007.x</u> <u>http://onlinelibrary.wiley.com/doi/10.1111/j.1365-2567.2008.03007.x/full#f2</u>





Atopy

- increased susceptibility to allergic disease (eg. hay fever, asthma)
- strong IgE-answer to environmental antigens
- high <u>lgE</u> and <u>eosinophylia</u> in the blood
- Genetic background:
 - **Chromosome 11q** high affinity $Fc \in R \beta$ -chain polimorfism
 - Chromosome 5q IL-3, IL-4, IL-5, IL-9, IL-13 and GM-CSF genes IgE isotype switch, eosinophil granulocyte survival, mast cell proliferation
 - IL-4 promoter increased activity higher IgE cc.
 - > IL-4-receptor α -chain gain-of-function mutation increased signaling strength

Therapeutic possibilities

- Allergen free environment
- Antihistamines
- Desensitization
- Membrane-stabilizing drugs
- Non-specific immunosuppression
- CD23 (inhibiting IgE receptor) activation

Diagnosis:



1. Intradermal skintest

2. ELISA: allergen-specific IgE measurement

Type II. hypersensitivity antibody-mediated citotoxic form

Type II hypersensitivitycytotoxic reactions

- antibody and cell-mediated cytotoxicity
- complement-mediated lysis
- IgG and IgM
- K-cells, platelets, neutrophils, eosinophils and macrophages
- > Examples:
 - Rh antigen
 - transfusion reactions
 - autoimmune haemolytic anemia
 - hyperacute graft rejection
 - reactions to tissue antigens

Type II. diseases

- <u>Antigens</u> are usually endogenous, sometimes exogenous chemicals (haptens), which can bind to cell surface.
- Drug-induced-hemolitic anemia, granulocytopenia,
 trombocytopenia
- <u>Diagnosis</u>: circulating antibodies and immunfluorescence on biopsy from the lesion
- <u>Therapy</u>: anti-inflammatory- and immunsupressive drugs



Type II. hypersensitivity (1)





ADCC and complement-mediated lysis

Complement- and Fc receptor – mediated inflammation





Abnormal physiologic responses without cell/tissue injury



Type II. hypersensitivity -

Figure 10.26



B cells are activated by antigen and by help from activated T_H2 cells P Ce RBC Plasma cells secrete penicillinspecific IgG which binds to modified red blood cells olasma lgG 狑

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Drug-induced hemolytic anemia

Hemolysis
Rh incompatibility



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Prophylaxis: anti-RhD antibody prophylaxis after delivery



Antibody-mediated Glomerulonephritis (1)

Goodpasture-syndrome

Anti-basement membrane antibody-mediated glomerulonephritis



Light microscopy



Immunofluorescence

The pathologic lesion contains antibodies, complement and neutrophils.

Staining is smooth and linear.



Pemphigus vulgaris

<u>Target antigen:</u> skin intercellular proteins: cadherin, desmosome <u>Symptoms:</u> blisters in the skin





Type III. hypersensitivity

Immuncomplex disease

Type III. hypersensitivity

- Immuncomplex disease
- Antigens are exogenous (chronic bacterial, viral or parasitic infections) or endogenous tissue molecules (Autoimmun diseases)
- Antigens are soluble. The patologic lesion contains antibody and complement factors.
- Tissue damage caused by neutrophils (inflammation) and platelets (thrombosis).



Types of Antibody-Mediated Diseases (2)



Fig. 18-1B

Diseases

- Caused by dissolved immuncomplexes. The outcome of the disease is influenced by the size of the immuncomplexes.
- might be general (eg. serum sickness) or organspecific:

Skin (SLE, Arthus-reaction) Lung (Aspergillosis, Farmer's lung) Blood vessels (Polyarteritis) Limbs (RA) Kidneys (lupus Nephritis)

• **3-10 hours** needed for the development

For diagnosis immuncomplexes have to be verified in tissue biopsy.

Granular staining is characteristic.

Immuncomplexes and low complement concentration in the serum.

Arthus-reaction: immuncomplex-mediated vasculitis



Antibody-mediated Glomerulonephritis (2)

Immune complexmediated glomerulonephritis







Light microscopy

Immunofluorescence

Electron microscopy

Fig. 18-3B

Type III. hypersensitivity



Type III. hypersensitivity

Disease	Symptom	Therapy
Serum sickness (GN, Arthritis, Vasculitis)	fever, limb pain, dermatitis, lymphadenopathia, proteinuria, breathing insufficiency	Clearance of immuncomplexes, supportive treatment
Polyarteritis nodosa	Pain, high blood pressure	Immunosupression
SLE, RA	Polyarthralgia (limb pain), face redness (dermatitis), lung- and kidney failure	Immunosupression
allergic bronchopulmonary Aspergillosis	Asthma, recurrent fever, chest pain	Corticosteroids against inflammation
Some cancers	Similar to serum sickness	Tumor excision

Delayed type hypersensitivity (DTH)

Type IV. hypersensitivity

TABLE 14-3 INTRACELLULAR PATHOGENS AND CONTACT ANTIGENS THAT INDUCE DELAYED-TYPE HYPERSENSITIVITY

Intracellular bacteria Mycobacterium tuberculosis Mycobacterium leprae Listeria monocytogenes Brucella abortus Intracellular fungi Pneumocystis carinii Candida albicans Histoplasma capsulatum Cryptococcus neoformans Intracellular parasites Leishmania sp.

Intracellular viruses Herpes simplex virus Variola (smallpox) Measles virus Contact antigens Picrylchloride Hair dyes Nickel salts Poison ivy Poison oak

Self tissue antigens

Alloantigens (Transplantation)

Phase 1 and 2 of DTH



<u>1. Sensibilization</u>: 1-2 weeks after the first antigen contact. APCs (Langerhans-cells, endothel cells or macrophages) produce IL-12 and induce Th1-cell differentiation.

2. Activation: Th1-activation, proliferation, rarely CD8+ CTL-activation.

2. contact with the antigen



<u>Effector phase</u>: 2. antigen stimulus leads to Th1-cell activation, citokin secretion (24h), recruitment of macrophages and other non-specific inflammatory cells (48-72h). From the infiltrating cells only 5% is T cell, 95% is non-specific.

Type IV. hypersensitivity



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Stages of macrophage activation

Resting >IFNgamma	Activated Hyperactivated >LPS, Immuncomplex double stranded RNA	
Phagocytosis	Antigen presentation	Tumor cell and parasite killing
Chemotaxis	Tumor cell binding	
Proliferation No cytotoxicity	decreased prolif.	No proliferation. No APC
MHC II -, O2 low	MHC II+, O2 high	MHCII -, O2high TNF,cytotoxic Protease secretion

4. phase of DTH

- <u>Granulomatous-reaction</u>: if the intravesicular pathogen survives in the cells it induces a prolonged DTH response – <u>chronic infection</u>
- → continous macrophage activation leads to citokin- and growth factor production and granuloma formation.
- Giant cells, epitheloid cells, tissue damage, necrosis, fibrosis.

The structure of granulomas



Diseases

- Infections: intracellular bacteria eg. *Mycobacterium tuberculosis, M. leprae;* Viruses: *Herpes simplex*
- Contact dermatitis, atopic ekzema
- Autoimmun diseases: Type 1 Diabetes Mellitus, Rheumatoid arthritis, Inflammatory bowel disease (IBD), Multiple sclerosis, Peripheral neuritis, Autoimmune myocarditis
- Transplant rejection: allogen tissue transplantation

Type IV. hypersensitivity – Tuberculotic granulomas





Poison ivy (Toxicodendron) Contact dermatitis



Comparison of Different Types of hypersensitivity

	type-l (anaphylactic)	type-ll (cytotoxic)	type-III (immune complex)	type-IV (delayed type)
antibody	lgE	IgG, IgM	lgG, lgM	None
antigen	Exogenous	cell surface	soluble	tissues & organs
response time	15-30 minutes	minutes-hours	3-8 hours	48-72 hours
appearance	weal & flare	lysis and necrosis	erythema and edema, necrosis	erythema and induration
histology	basophils and eosinophil	antibody and complement	complement and neutrophils	monocytes and lymphocytes
transferred with	antibody	antibody	antibody	T-cells
examples	allergic asthma, hay fever	erythroblastosis fetalis, Goodpasture's nephritis	SLE, farmer's lung disease	tuberculin test, poison ivy, granuloma