### Hypersensitivity

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#### Origins of Hypersensitivity

### •"Hypersensitivity" first used clinically in 1893:

 During attempts to protect against diphtheria toxin, it was found that an animal would suffer enhanced responses and even death following its second exposure to toxin at a dose too small to injure normal untreated animals



Emil von Behring

- •The term "Allergy" is coined in 1906:
  - These hypersensitivity reactions were postulated to be the product of an "allergic" immune response, derived from the Greek allos ergos (altered reactivity)



Clemens von Pirque

Photos from Silverstein, AM. 1989. A History of Immunology. Academic Press, San Diego

#### **Definitions**

- Hypersensitivity:
  - Broadest (Abbas) Disorders caused by immune responses
    - Dysregulated response to foreign antigen
    - Failure of tolerance to self-antigen
  - Practical Used clinically to refer to aberrant or excessive immune responses generated against foreign antigens, although the same immune processes apply in many autoimmune diseases
- Allergy:
  - Symptoms elicited by encounter with foreign antigen in a previously sensitized individual

#### Manifestations of Hypersensitivity

•Symptoms frequently are localized to the anatomical site of antigen exposure:

Site of Exposure	Syndrome	Common Allergens	Symptoms
Respiratory	Allergic Rhinitis		Nasal Pruritis Rhinorrhea Congestion
Mucosa	Asthma	Who will be	Bronchospasm Chronic Airway Inflammation
G.I. Mucosa	Food Allergy		Cramping Vomit/Diarrhea Hives Anaphylaxis

### Manifestations of Hypersensitivity

Site of Exposure	Syndrome	Common Allergens	Symptoms
Claire	Contact Urticaria		Hives Pruritis
Skin	Contact Dermatitis		Rash Pruritis
Blood	Systemic Allergy		Hives/Edema Abd. Cramping Bronchospasm Hypotension

# Hypersensitivity: Gell & Coombs Classification

	Type I	Type II	Type III	Тур	e IV
Common Name	Immediate Hyper- sensitivity	Bystander Reaction	Immune Complex Disease	Delaye Hyperse	<i>J</i> 1
Example	Peanut Anaphylaxis	PCN-assoc. Hemolysis	Serum Sickness	Contact Dermatitis (Ni <sup>+</sup> ), PPD	Contact Dermatitis (poison ivy)
Mediator	IgE	IgG Monomer	IgG Multimers	CD4 T cell	CD8 T cell
Antigen	Soluble	Cell or Matrix Bound	Soluble	Soluble	Cell- associated
Effector Mechanism	Mast Cell Activation	Complement FcγR <sup>+</sup> Cells	Complement PMN, МФ	Macrophage Activation	Cytotoxicity (perforin/ granzyme)

#### Common to All Types

- Products of the adaptive immune system
  - Require at least one exposure for sensitization to occur
  - Sensitization can be long lived in the absence of re-exposure (>10 years) due to immunologic memory
  - Antigen is a protein or is capable of complexing with protein (e.g., nickel ion, penicillin)

#### Type I (Immediate) Hypersensitivity

- Antigens:
  - Exogenous, otherwise innocuous
  - Contact typically occurs via mucous membranes (respiratory, GI) and at low dose
- Immune Mechanism
  - Antigen contact first leads to IgE production: Sensitization
  - On re-exposure, pre-formed antigen-specific IgE triggers mast cell activation resulting in symptoms: hive, wheeze, itch, cramps
- Reactions:
  - Occur within seconds-minutes of exposure
  - Severity ranges from irritating to fatal

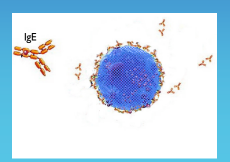
#### **IgE Production**



- Occurs as part of a secondary immune response (generally multiple or persistent exposures)
- Class switch to IgE is directed by IL-4 and IL-13 (Th2 cytokines), and requires T cell help via CD40L
- The propensity to make an IgE response to environmental antigens varies among individuals
- "Atopic" individuals are those with an inherited predisposition to form IgE responses

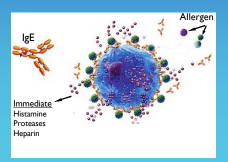
#### Type I Rxn: Sensitization Stage

- IgE produced by plasma cells has a short circulating half-life (serum  $T_{1/2}$ ~2 days; comp. to IgG~30 days)
- $\bullet$  Rapidly taken up by  $\mathsf{Fc}_\epsilon \mathsf{RI}$  on tissue mast cells and circulating basophils



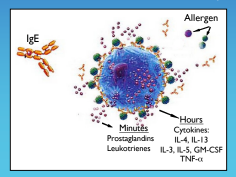
#### Type I Rxn: Effector Stage

- Early Phase Response: within seconds-minutes
  - IgE crosslinking by antigen → release of preformed mediators
  - histamine smooth muscle constriction, mucous secretion, vascular permeability, GI motility, sens. nerve stimulation



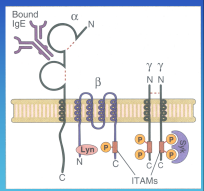
#### Type I Rxn: Effector Stage

- Late Phase Response: 6-24 hours after exposure
  - Mast cell production of newly synthesized mediators
    - Leukotrienes → smooth mm. contraction, vasodil., mucous prod.
    - Cytokines → recruitment of PMN and eosinophils

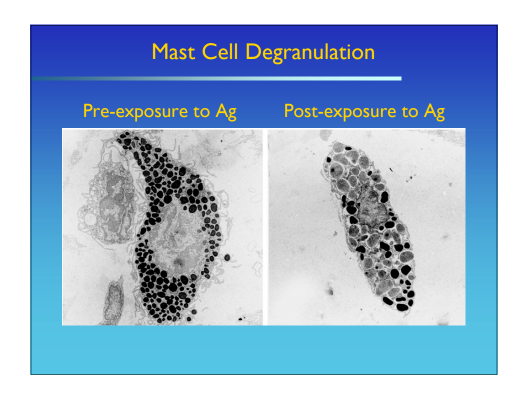


#### Fc<sub>ε</sub>RI Signaling

- Structure:  $\alpha\beta\gamma_2$ 
  - Alpha- binds IgE monomer
  - Gamma- shared by IgG FcR's I & III
- Receptors are aggregated
  - When pre-bound IgE binds multivalent Ag
  - Initiates ITAM phosphorylation
- ITAM's
  - Conserved tyrosine-containing sequence motifs within a variety of receptors (TCR, BCR, FcRs)
  - Serve as docking sites for downstream activating kinases, in this case, Syk



Immunoreceptor
Tyrosine-based
Activation
Motif



#### **Eosinophils**

- Innate responder cell in Type I hypersensitivity
- Production: Induced in the bone marrow by:
  - IL-5 Th2 cytokine, drives specifically eosinophil production
  - IL-3, GM-CSF drive granulocyte production in general
- Chemotaxis: Homing to tissue sites utilizes:
  - IL-5, Eotaxins-1, -2, & -3
- "Primed" for activation by IL-5, eotaxins, C3a & C5a
  - $\hat{U}Fc_{\gamma}R$  &  $Fc_{\alpha}R$  expression;  $\hat{U}C'$  receptor expression
  - induce  $Fc_{\epsilon}R$  expression
  - Uthreshold for degranulation

#### Eosinophils

- •Activation:
- Most potent trigger is Ig-crosslinking (IgA>IgG>IgE)
- Potentiated by IL-5, GM-CSF, granule proteins (MBP), C3a/C5a
- Results in exocytosis of pre-formed eosinophil toxic proteins
- Anti-microbial effect:
- major basic protein
- eosinophil cationic protein
- eosinophil-derived neurotoxin



All have pl's >10

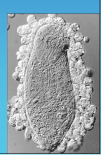
Directly toxic to helminths

Also cause tissue damage

- Mobilize more innate responders
- Secretion of IL-3, IL-5, GM-CSF (more eos), IL-8 (PMN)
- Elaboration of LT-C4, -D4

#### **Evolutionary Role of Type I Response**

- Mast cells line all subepithelial mucosa
  - Rapid recruitment of PMN, eosinophils, monocytes to sites of pathogen entry
  - 12 Lymph flow from peripheral sites to lymph node
  - ûG.l. motility → favors expulsion of G.l. pathogens
- Important role in parasite clearance
  - c-kit<sup>-/-</sup> mice have no mast cells → û susceptibility to *Trichinella*, *Strongyloides*
  - Eosinophil depletion (Ab-mediated) → ① severity of schistosomal infection



#### Allergy Epidemic

- •Type I Hypersensitivity diseases, including asthma and allergic rhinitis, have been increasing in prevalence in the economically "advantaged" parts of the world for 30 years
  - The "hygiene hypothesis" attributes increased allergic disease rates to generally decreasing microbial exposure in early life which would normally provide a Th1-promoting effect
    - Neonatal bias:  $\mbox{$\mbox{$\mbox{$$}$IL-12 (DC)$}}$  and  $\mbox{$\mbox{$\mbox{$$}$IFN-$$}$}\gamma$  (T cells)
    - Birth order: 

      ¬allergy rates among 3rd- and 4th-born children
    - Protective effect of day care
    - Hx of measles or HAV infection, or +PPD → \$\Pi\$ allergy rates
    - 1990 East/West Berlin immediately after the wall fell: East had
    - - 
       \$\Psi\$ vaccination rates, \$\hat{\Omega}\$ prev. childhood infection, but \$\Psi\$'ed asthmatically in the second content of the second content of

#### Allergy Epidemic

- Weighing against the Hygiene Hypothesis:
  - Despite this epidemiologic data, some evidence is hard to reconcile
    - Previous infection with helminths, which generates a strong Th2 response, is also associated with protection against allergy
    - Early life exposure to pathogens is also associated with decreased risk of autoimmune disease (e.g., type I diabetes), a classic Th I -mediated condition
  - Revised hygiene hypothesis early life exposure to microbial pathogens influences the balance of immune responsive vs. immune modulating influences, not simply Th1-Th2 balance

#### Type I Hypersensitivity in Allergy

- Manifestations of Type I Hypersensitivity:
- Allergic Rhinitis/conjunctivitis ("Hayfever")
- Asthma prevalence û 60% in the past 20 years
- Food/Medication reactions urticaria (hives)
- Contact urticaria
- Some forms of eczema
- Anaphylaxis systemic reaction induced by food, venom, medication, etc.

### Demonstrating Type I Hypersensitivity in the Patient

- Documenting allergic sensitivity: skin testing
- Allergen (airborne, food, venom, some medications) is introduced by prick or intradermal injection
- Sensitization is evident within 15-20 minutes as a wheal/flare at the allergen introduction site



#### **Anaphylaxis**

- Response to systemic circulation of allergen
  - Triggering of mast cells in peri-vascular tissue

  - High-output shock: ♥BP despite û'ed cardiac output
  - Other symptoms: flushing, urticaria, wheeze, laryngeal edema with airway compromise, G.l. cramping, diarrhea
- Rapid progression over seconds to minutes
- Treatment -
  - immediate administration epinephrine I.M., followed by antihistamines (HI and H2 blockade) → treat early phase
  - subsequent administration corticosteroids → prevent late phase

#### Type II Hypersensitivity

- Antibody-mediated "Bystander Reactions"
  - Immune effector is target-specific IgM and IgG
  - (Contrast with Type III Rxns in which the Ig is not specific for the tissue being damaged)

#### Clinical Manifestations:

- Classically manifests as a reaction to a foreign substance (most commonly a drug) acting as a hapten
- The same mechanisms, however, manifest with autoimmunity through the process of molecular mimickry

#### Type II Hypersensitivity

#### Drug Reactions

- Hapten a molecule too small to elicit an immune response itself, but capable of covalent conjugation to self proteins, creating a new (non-self) target or epitope
  - \_ example: penicillin is metabolized to yield the penicilloyl moiety which binds surface proteins on blood cells and platelets
  - \_ penicilloyl-proteins represent neoepitopes → break tolerance

#### Molecular Mimickry

- Pathogen elicits an appropriate Ab response
- Ab cross-reacts with self-tissue (very similar epitopes)
  - Group A Strep pharyngitis yields Ab's to the Strep M protein
     Ab's cross react with cardiac muscle and valves

### •Mechanisms of Type II Hypersensitivity: Exactly those of normal Ab function (plus some):

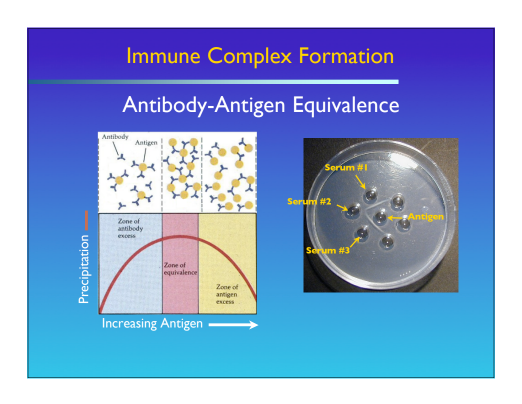
Ab Function	Target	Result	Syndrome	
Opsonization	Platelet surface proteins	Splenic clearance	Drug-induced thrombocytopenia	
Neutralization	Acetylcholine receptor	Receptor blocking	Myasthenia gravis	
ADCC	Glomerular basement membrane proteins	Glomerular destruction	Post-Streptococcal renal failure	
Complement- mediated lysis	Penicilloyl-RBC protein conjugates	RBC destruction	Drug-induced hemolytic anemia	
Non- Physiologic	TSH receptor	Receptor activation	Grave's disease	

# Type III Hypersensitivity: Immune Complex Disease

- First Description: Arthus Reaction
- Rabbit received an intravenous infusion of anti-toxin antibody

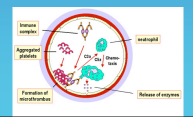


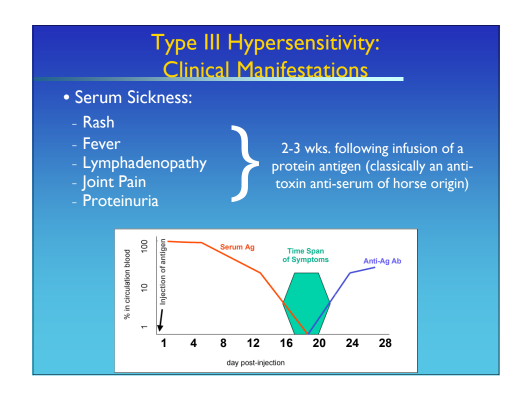
- Three days later, antigen (toxin) was injected subcutaneously
- Local erythema/tenderness with edema, necrosis, and hemorrhage developed within 8 hours = Arthus Reaction

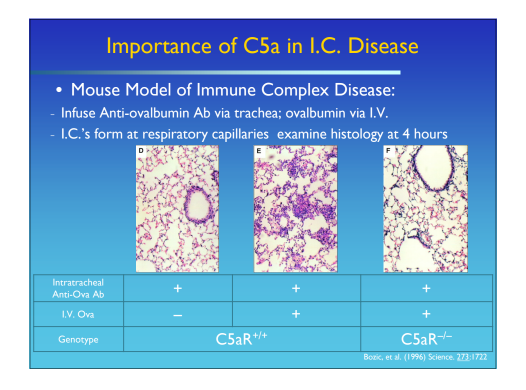


#### **Arthus Reaction**

- Immune Mechanism
  - Antibody-Antigen complexes form within blood vessel walls
  - Complement fixation generates C5a
    - Neutrophil chemoattractant → PMN infiltration
    - Anaphylatoxin local mast cell histamine release → tissue edema
  - Neutrophil activation by FcγR's → release of cytotoxic enzymes
  - Platelet aggregation by FcγR's → small vessel thrombosis, necrosis
  - Local macrophage release of IL-1, TNF- $\alpha$ , and IL-8  $\Longrightarrow$  propagation

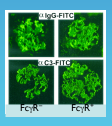


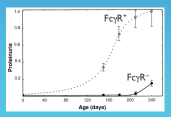


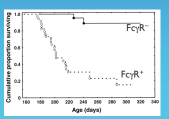


#### Importance of Fc<sub>7</sub>R's in I.C. Disease

- •B/W Mouse spontaneous accumulation of I.C.'s in the glomerulus leads to early death from renal failure
- •FcγRI and FcγRIII contain ITAM's; activating for phagocytes
- •Lack of FcγRI/FcγRIII protects against I.C.-mediated glomerular damage, despite accumulation of IgG/C3b-containing immune complexes







Clynes, et al. (1998) Science. <u>279</u>:1052

# Type IV (Delayed-Type) Hypersensitivity

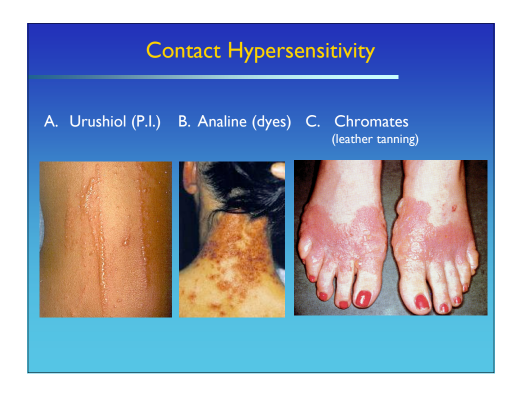
- •Group of related responses to antigen, all dependent on T cell-mediated immunity
- •Prior sensitization is required
- •Reactions occur over I-3 days following re-exposure
- •T cells: necessary and sufficient for DTH
- Athymic subjects (animal or human) do not get DTH rxns.
- T cell depletion (via anti-T cell Ab's) reverses sensitization
- Transfer of purified memory T cells confers sensitization

#### Manifestations of DTH Reactions

Туре	Site	Clinical Appearance	Antigen
Contact	Epidermis	Erythematous Papular Scaling Blistering	Poison ivy, latex, organic mols., metals (Ni <sup>++</sup> )
Tuberculin	Dermis	Local Induration	Mycobacteria, Candida, Mumps

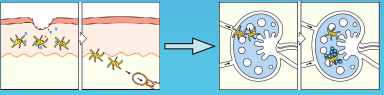
#### Common to all DTH Reactions

- Histology of the DTH reaction:
  - T Cells CD4 (Th1); some forms CD8
  - Macrophages/monocytes
  - Basophils
  - Tissue edema with fibrin extravasation
  - If persistent antigen: multinucleated giant cells; granulomata
- Cytokines found at the site of a DTH reaction:
  - IL-2
  - IFN-γ
  - TNF-o
  - Macrophage chemotactic protein (CCL-2)



#### Contact Sensitivity: Hapten DTH

- Phase One: Initial Exposure Sensitization
  - Antigen typically a small organic hapten, frequently lipophilic
  - Exposure crosses epidermal barrier by diffusion, associates with epidermal cell proteins ("haptenylation")
  - Processing haptenylated proteins are picked up by
    Langerhans cells → peptides → loaded onto MHC I and II
  - Presentation loaded LC's migrate to regional lymph nodes where they present haptenylated proteins to naive T cells



#### Contact Sensitivity: Hapten DTH

- Phase Two: Re-exposure Elicitation
  - Hapten-specific memory T cells bearing the cutaneous lymphocyte antigen (CLA-I) continuously migrate between lymphatics and skin
  - Re-encounter with haptenylated protein may occur on:
    - Langerhans cell (MHC II) → CD4<sup>+</sup> T cell activation → secretion of IFN-γ, MCP-1 → macrophage recruitment
    - Keratinocyte (MHC I) (lipophilic hapten) → CD8+ CTL activation release of perforins and granzyme → local tissue damage

## Hypersensitivity: Gell & Coombs Classification

	Type I	Type II	Type III	Тур	e IV
Common Name	Immediate Hyper- sensitivity	Bystander Reaction	Immune Complex Disease	Delaye Hyperse	
Example	Peanut Anaphylaxis	PCN-assoc. Hemolysis	Serum Sickness	Contact Dermatitis (Ni <sup>+</sup> ), PPD	Contact Dermatitis (poison ivy)
Mediator	IgE	IgG Monomer	IgG Multimers	CD4 T cell	CD8 T cell
Antigen	Soluble	Cell or Matrix Bound	Soluble	Soluble	Cell Associated
Effector Mechanism	Mast Cell Activation	Complement FcR <sup>+</sup> Cells	Complement PMN, МФ	Macrophage Activation	Cytotoxicity (perforin/ granzyme)

#### Hypersensitivity Progression

- Antigen-specific responses may progress from one type of hypersensitivity to another:
  - Latex allergy among healthcare workers
    - Initial reaction is typically a contact sensitivity (type IV reaction)
    - With recurrent latex contact, sensitivity progresses to latexspecific IgE, imparting risk of anaphylaxis
  - p-aminobenzoic acid (PABA), the active ingredient in many sunscreens, can act as a contact sensitizer
    - PABA DTH reactivity is associated with û'ed risk of immediate (type I) hypersensitivity to local anesthetics (e.g., benzocaine) due to cross-reactivity of the aromatic core

#### Penicillin Mediates All Types

•Immune-mediated adverse reactions occur at a rate of I per 100 administrations (!)

Type	Mechanism	Example
ı	IgE-mediated	Acute anaphylaxis, Urticaria
II	C'-mediated cytolysis Opsonization	Hemolytic anemia Thrombocytopenia
III	Immune Complex Damage	Serum sickness Drug fever, Vasculitis
IV	T Cell mediated	Contact sensitivity