## An Overview: Nonspecific vs. Specific Defense Mechanisms

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Body Defenses: An Overview

Nonspecific Defenses

• Hinder entry of most pathogens (disease causing organisms)
• Prevent spread of disease
• Strengthen immune response
Body Defenses: An Overview

Specific Defenses: The Immune System

» Destroys foreign cells that cause disease
» Recognize self from non-self
» Inactivate toxins and other foreign chemicals
Nonspecific Response
Surface Barriers: 1\textsuperscript{st} line of defense

1. **Intact Skin** (epidermis)
   - Forms mechanical barrier
   - Acid secretion inhibits bacterial growth
   - Keratin provides resistance against bacterial enzymes, acids, bases

2. **Mucus membranes**
   - Trap microorganisms in respiratory and digestive tracts
E. coli can cause urinary tract infections and traveler's diarrhea.
Flushing Away of Bacteria Lacking Pili

- Lacking any means of adherence, urine typically flushes bacteria from the urethra.

http://www.cat.cc.md.us/courses/bio141/lecguide/unit1/bacpath/pili/nopili.html
Bacterial fight back!

Bacteria with Pili Resisting Flushing

1. Bacteria such as *Neisseria gonorrhoeae* use pili to adhere to the mucous membranes of the urethra and thus resist the flushing action of the urine.

http://www.cat.cc.md.us/courses/bio141/lecguide/unit1/bacpath/pili/yespili.html
Bacterial fight back!
*Salmonella* Invading an Intestinal Mucosal Epithelial Cell

- *Salmonella* adheres to the microvilli of intestinal mucosal epithelial cells.
- Triggers pseudopods that engulf the *Salmonella* and place it in a vacuole.
- *Salmonella* replicate within the vacuole and are subsequently released from the cell.

http://www.cat.cc.md.us/courses/bio141/lecguide/unit1/bacpath/salinv.html
Nonspecific Response
1st line of defense (cont.)

1. **Gastric juice** → kills pathogens
2. **Acid lining of vagina** → kills pathogens
3. **Tears & saliva**
   » Contain lysozyme → kills pathogens
4. **Nasal hairs**
   » Filter and trap microorganisms
5. **Cilia** propel mucus away from lower respiratory tract
First-line respiratory defenses
2nd line of Nonspecific defense:
Nonspecific cellular defenses

1. Phagocytes: macrophages & neutrophils
   - Feed on pathogens that penetrate membrane barriers
   - Display antigen to enhance immune response (specific response)

2. Natural killer cells
   - Lyse and kill cancerous and virally infected cells
NK Cell destroying a target cell

Killer Cell → Target Cell

Target-oriented Granules

Surface Contact
Phagocytosis by a Macrophage
(a) Phases of phagocytosis

1. Chemotaxis and adherence of microbe to phagocyte.
2. Ingestion of microbe by phagocyte.
3. Formation of a phagosome.
4. Fusion of the phagosome with a lysosome to form a phagolysosome.
5. Digestion of ingested microbe by enzymes.
6. Formation of residual body containing indigestible material.
Phagocytosis Animation by Cells Alive: http://www.cellsalive.com

- How to get to the animation
  1. Click on “Immunology”
  2. Click on “Ouch—anatomy of a splinter”
  3. Go to bottom of page and Click on “Next”
  4. Click on “Phagocytosis—how white cells eat microbes”
2nd line of Nonspecific defense:
Nonspecific chemical defenses

Inflammatory response

1. Prevents spread of damaging agents
2. Removes cell debris and pathogens
3. Aids in tissue repair
4. Triggered when tissues are damaged or irritated
   • Physical trauma, heat, chemicals, pathogens
Inflammatory Response: a simplified view
Animation of Inflammatory Response
http://www.whfreeman.com/immunology/CH01/diapedesis.htm
Inflammatory Response (cont.)

- **Cardinal signs**: Redness, heat, swelling, pain
- **Sequence of events**
  1. Damaged tissue →
  2. Histamine & kinins →
  3. Vasodilatation + capillaries become leaky →
  4. Edema (swelling) →
  5. Pain receptors activated →
  6. WBC’s (B & T cells + phagocytes) attracted
Inflammatory Response: a detailed view

(a) Tissue damage

1. Chemicals such as histamine, kinins, prostaglandins, and leukotrienes (represented as blue dots) are released by damaged cells

2. Blood clot forms

3. Abscess starts to form (yellow area)

(b) Vasodilation and increased permeability of blood vessels

(c) Phagocyte migration and phagocytosis

4. Margination—phagocytes stick to endothelium

5. Emigration—phagocytes squeeze between endothelial cells

6. Phagocytosis of invading bacteria

(d) Tissue repair

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Nonspecific chemical defenses

Anti-microbial Chemicals

1. Complement protein system
   • More than 20 plasma proteins
   • Produced by the liver
   • Inactive until contact with pathogen

Roles
a. Enhance inflammatory response
   - Vasodilatation
   - Attract phagocytes
b. **Lyse foreign cells**
   - Poke holes in membrane → water enters → cells burst

c. **Opsonisation**
   - Cause foreign cells to stick together
   - Yummy for phagocytes → enhances phagocytosis
Animation of Phagocytosis Enhanced by Opsonization

http://www.catt.cc.md.us/courses/bio141/lecguide/unit1/bacpath/phagocytosis/phagsum.html
Bacterial fight back!

Bacterial Resistant to Phagocytosis

1. In some bacteria, a capsule covers the opsonin C3b on bacterial cell wall
2. Prevents binding to C3b receptors on the surface of phagocytes:
Bacterial fight back!

*Bacillus anthracis* in Action: Inhalation Anthrax

Source: *Scientific American* (March 2002)

1. Macrophage cells ingest *B. anthracis* spores and carry them to lymph nodes
   - In blood and inside macrophage: Spores transform into actively dividing cells
2. Proliferating *B. anthracis* cells erupt from macrophage cells
3. *How B. anthracis* cells evades destruction
   - Produce capsule that protects them from digestion by macrophages and antibodies from B-cells
   - Produce toxin enters and that impairs immune cell function
Bacterial fight back!

Blocking Phagosome Formation by Depolymerizing Actin

1. Some bacteria secrete proteins which depolymerize the phagocyte's actin microfilaments used for phagocytic engulfment.

http://www.cat.cc.md.us/courses/bio141/lecguide/unit1/bacpath/actinan.html
Anti-microbial Chemicals (Cont.)

2. Interferons
   - Released by virally infected cells
   - Stimulate neighboring cells to produce anti-viral proteins (AVP)
   - AVP’s inhibit viral reproduction in nearby cells
Interferons

1. Viral RNA stimulates host cell to synthesize interferon.
2. New viruses are produced by multiplication.
3. Meanwhile, alpha and beta interferons react with plasma membrane or nuclear membrane receptors on uninfected neighboring cell and induces synthesis of antiviral proteins (AVPs).
4. New viruses are released and infect neighboring cell.
5. AVPs block viral protein synthesis and thus interfere with viral multiplication.

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3. **Pyrogens**
   - Secreted by macrophages
   - Cause mild fever $\rightarrow$ Slows down bacterial reproduction
The 3rd Line of Defense: Specific Defense Mechanisms

1. The immune response
   a. Functional system that seeks to destroy or inactivate *specific* antigens (foreign molecules and cells)
   b. Antigen specific: responds to particular foreign substances
   c. Systemic response: body wide response
   d. Memory: quicker and stronger response to previously encountered pathogens
Branches Of The Immune System

1. Humoral immunity
   » Where B-cells circulating in the body’s fluids produce antibodies in response to a particular antigen

2. Cell-mediated immunity
   » Where killer T-cells (cytotoxic T-cells) destroy
     - cancer cells
     - virus and bacteria infected cells
     - cells of foreign grafts
Humoral and Cell-mediated Immunity

1. An antigen-presenting cell (APC) encounters and processes an antigen, forming MHC-antigen complexes on its surface.

2. A helper T ($T_H$) cell receptor binds to the complex, stimulating the APC to secrete interleukin-1.

3. This interleukin-1 stimulates the helper T cell to produce interleukin-2, which then stimulates that helper T cell to form a clone of helper T cells.

4. The cells of this clone in turn produce cytokines, stimulating cells of both immune systems.

(C) BENJAMIN/CUMMINGS
Immune Response: Roles of Cells involved

1. **Macrophages**
   - Antigen presenter (APC = Antigen Presenting Cell)
   - Activate Helper T Cells (by contact w/ displayed antigen)
   - Releases cytokines: Enhances Helper T Cell activation

2. **Helper T Cells**
   - Stimulate the humoral and cell-mediated response against specific antigens
   - Activated by Macrophages (Cytokines and contact w/ presented antigen)
   - Activate B-Cells and Killer T-cells by secreting cytokines
Humoral Immunity

1. **B lymphocytes (B-cells)**
   - Produced and mature in the bone marrow
   - Mature = become immunocompetent = ability to respond to an antigen by binding to it

2. B cells that bind to self-antigens are destroyed

3. Move to and concentrate in lymphatic tissue

4. Produce antibodies specific for specific antigens
   - Plasma cells vs. memory cells
Humoral Immunity

1. The microbial antigen is ingested by an APC and partially digested. Antigen fragments are presented on the APC surface and form complexes with self (MHC) molecules.

2. A helper T (T_H) cell specific for the presented antigen interacts with the complex.

3. The helper T cell then activates an appropriate B cell, probably one that itself has such complexes on its surface, as well as receptors for the microbial antigen. (There may also be direct stimulation by the microbial antigens.)

4. This interaction triggers the B cell to differentiate into a plasma cell, which secretes antibodies specific for the T-dependent antigen.
Clonal Selection Theory

Antigen receptor

Variety of B cells

Antigen molecules

Clone of memory cells

Clone of plasma cells

Antibody molecules
Antibody Structure:

Y-shaped molecules

(a) Antibody molecule

(b) Enlarged antigen-binding site bound to an antigenic determinant

(c) Antibody symbol
Each Antibody binds to a specific Epitope
Each Antibody binds to a specific Epitope
Major Antibody Types and their functions

- **IgM**
  - first to be made, good opsonizer and complement fixer

- **IgA**
  - Found in secretions: e.g. mucous and milk

- **IgG**
  - most abundant in blood, goes to fetus, good opsonizer and OK complement fixer, helps NK cells kill

- **IgE**
  - Causes allergies, hypersensitivities and anaphylactic shock; Defends against parasites
Mechanism of Antibody Action

Binding of antibodies to antigens inactivates antigens by:

- **Neutralization** (blocks viral binding sites; coats bacteria and/or opsonization)
  - Virus
  - Bacterium

- **Agglutination of antigen-bearing particles, such as microbes**
  - Bacteria

- **Precipitation of soluble antigens**
  - Soluble antigens

- **Complement fixation (activation of complement)**
  - Complement Lesion
  - Foreign cell

Enhances:

- **Phagocytosis**
  - Macrophage

Leads to:

- **Cell lysis**
Complement Protein-Ab interaction

Results in Cell Lysis

1. Pathogen’s membrane
2. Complement proteins
3. Membrane attack complex forming
4. Pore
1\textsuperscript{st} and 2\textsuperscript{nd} Exposures to a Specific Antigen

**Primary response**

- Initial exposure to antigen

**Secondary response**

- Second exposure to antigen
- IgG
- IgM

Antibody titer in serum (arbitrary units) vs. Time (days)
T Lymphocytes (T-Cells)

1. Produced in bone marrow
   » Multiply & mature in thymus gland (takes 2-3 days to become immunocompetent)
2. T-cells that bind to self-antigens are destroyed
3. Move to and concentrate in lymphatic tissue
4. Kinds of T-cells
   » Killer T-cells, helper T-cells, memory cells, suppressor T-cells
Development of Lymphocytes

Bone marrow or fetal liver

Pluripotent stem cell

Lymphocyte stem cell

Thymus

B cell

T cell

Lymphoid tissue (lymph nodes, spleen, blood, and lymph)
Lymphatic System

- Returns tissue fluid that has escaped the circulatory system back to the blood
- Lymph nodes filter out antigens from the lymph fluid
- Lymphocytes concentrate/reproduce in the lymph nodes, especially during infections
Lymph Node: Filters Lymph

- Afferent lymphatic vessel
- Capsule
- Cortex
- Medulla
- Sinuses
- Efferent lymphatic vessel
- Lymph node artery and vein
Killer T-Cell Activation requires:

1. Specific Pre-killer T-Cell binds to target cell
2. Cytokines secreted by Helper T-Cell
3. Helper T-Cell secretes cytokines after interacting with AP (macrophage)
Cytotoxic T-Cell inducing Apoptosis in Target Cell

- The binding of a Cytotoxic T-cell to the specific receptor molecules in a target cell may activate processes that lead to apoptosis: programmed cell suicide.
Cytotoxic T-Cell Lysing a Cell by Secreting Perforin

1. Cytotoxic T (T<sub>C</sub>) cell binds to infected cell.
2. T<sub>C</sub> cell releases perforin, which makes lesions in infected cell's membrane.
3. Infected cell lyses.
Animation of Cytotoxic T-Cell Lysing a Cell

http://www.cellsalive.com/ctl.htm
Active Immunity

- Naturally acquired during infections. How?
- Artificially acquired by vaccines
  - Vaccines contain dead or attenuated (weakened) organisms that stimulate the immune response
  - Booster Shots...Why needed?
Humoral and Cell-mediated Immunity

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Cell-mediated immunity (attack on infected cells)

Humoral immunity (secretion of antibodies by plasma cells)
Passive Immunity

• Antibodies obtained from outside source
• Effect is short term (2-3 weeks) — why?
• Examples
  – Via placenta and mother’s milk
  – Via injections:
    ➢ Gamma globulin
    ➢ Anti-venoms
    ➢ Anti-toxins (botulism, rabies, tetanus)
Disorders of the Immune System

1. **Allergies (Hypersensitivities)**
   - Immune Response against *harmless antigens* called **Allergens**
     - Results in damage to various body tissues

2. **Major Types of Allergies**
   - **Immediate or Acute Hypersensitivity**
     - Most common; Localized Reaction in Seconds
     - Plasma cells release IgE in response to allergen
     - IgE stimulates mast cells to release histamine
       - Blood vessels dilate and leak, causing...
         » Runny nose, watery eyes, itching, hives
Allergic Response

1. Allergen stimulates Ab production (humoral immunity)

2. Ab binds to and sensitizes mast cell

3. Binding of allergen to mast cell stimulates release of histamine

4. Histamine results in inflammatory response
Disorders of the Immune System (cont.):

**Anaphylactic Shock**

1. **Systemic Allergic Response to an allergen that enters directly into the blood**
   - Bee and ant stings, spider and snake bites
   - Injection of foreign substance: Penicillin, drugs, foreign proteins, mismatched blood types, etc.

2. **Allergen circulates quickly through entire body causing death in minutes**
   - Rapid Drop in Blood Pressure...Why?
   - Constriction of air passages (bronchioles) in lungs makes breathing difficult
   - Treat with epinephrine
Disorders of the Immune System (cont.):

Delayed Hypersensitivities

1. **Allergic contact Dermatitis**
   - Most common type
   - Direct skin contact with...
     - Poison oak/ivy, heavy metals (e.g. Pb, Hg, Ba, etc.), misc. chemicals in cosmetics, deodorants, foods, etc.
     - 1-3 days to respond

2. **Symptoms are caused by lymphokines released by Killer T Cells**
   - Treat with corticosteroid drugs (e.g. cortisone)
   - Why won’t antihistamines work?
Disorders of the Immune System (cont.):

Autoimmune Diseases

1. Killer T Cells or Antibodies attack the body’s own tissues as if they were foreign

2. Possible Causes..
   a. B and T Cells that attack self are not eliminated after they are produced
   b. Appearance of new self proteins/antigens
      - Possibly due to mutations, or
      - Bacterial or viral modification of self-proteins
   c. Cross-reaction of Antibodies or Killer T Cells produced during infection against self
Disorders of the Immune System (cont.):

AIDS: Acquired Immunodeficiency Syndrome

(a) HIV

- gp120
- gp41
- p32 integrase
- ssRNA
- p10 protease
- p17
- p24
- MHC proteins
- Reverse transcriptase

HIV
HIV from Cells Alive

- Go to the computer lab and study Human Immunodeficiency Virus (HIV) Infection at the Cells Alive website:
  
  http://www.cellsalive.com

Once at the website

- Click on Immunology from the menu on the left
- Then click HIV infection
HIV Infects and Kills Helper T-Cells

• What effect will this have on each of the following. Explain your reasoning.
  a. Macrophage activity
  b. Humoral Immunity
  c. Cell-mediated Immunity
  d. Complement protein activity
  e. NK Cells
  f. Inflammatory response
  g. Interferon activity
Case Study Questions

- Now is your chance to test your knowledge of the human immune system.
- Answer the following 11 case studies in as much depth and detail as your knowledge allows.
Case Study Question 1
Humoral and Cell-Mediated Immunity

• An individual failed to develop a thymus gland because of a genetic defect, what would happen to humoral and cell-mediated immunity?
Case Study Question 2
Nonspecific Defenses

a. What abnormalities would a person have that has a neutrophil deficiency?
b. A person with a macrophage deficiency?
Case Study Question 3
Nonspecific and Specific Defenses

• AIDS patients have a low helper T cell count. Would you expect patients with AIDS to develop fever in response to an infection?
Case Study Question 4
Specific Defenses

- A patient with symptoms of hyperthyroidism is found to have circulating antibodies that attack the receptors for TSH (thyroid stimulating hormone). What is the cause of hyperthyroidism in this patient?

  » **Background Information**: The thyroid gland produces thyroxin in response to a TSH, a hormone released by the pituitary gland. The binding of TSH to TSH receptors located on the cells of the thyroid stimulate it to release thyroxin, a hormone that helps to regulate the bodies metabolic rate. Hyperthyroidism involves elevated secretion of thyroxin and, hence, an elevated rate of metabolism.
Case Study Question 5
Specific Defenses

• If the thymus gland of an experimental animal is removed immediately following birth, the animal exhibits the following characteristics:
  » High susceptibility to infection
  » Decreased numbers of lymphocytes
  » Ability to reject graphs is reduced
• Explain these observations.
Case Study Question 6
Specific Defenses

• **Background Information:** Tetanus is caused by bacteria that enter the body through wounds in the skin. The bacteria produce and release a toxin that causes spastic muscle contractions. Death often results from the failure of the respiratory muscles.

• A patient comes to the emergency room after stepping on a nail. If the patient has been vaccinated against tetanus, the patient is given a tetanus booster shot, which consists of the toxin in an altered so that it is harmless. If the patient has never been vaccinated against tetanus, the patient is given an antiserum shot against tetanus toxin. **Explain the rationale behind these treatments.**
In reference to case study question 6: Sometimes both a booster shot and antiserum shot are given, but at different locations in the body. Explain why this is done, and why the shots are given in different locations.
In some cases, a booster shot is used as part of a vaccination procedure. A booster shot is another dose of the original vaccine given some time after the original dose was administered. Why are booster shots given?
Case Study Question 9

- Some people with a deficit of IgA, immunoglobulin A, have chronic sinus infections and respiratory tract infections. Explain these symptoms.
A few days after a walk in the hills you develop a severely itchy rash. What’s happening and what is the method of treatment?
Case Study Question 11

- Minutes after walking into a carpet showroom you begin suffering from symptoms of asthma (constrictions of the passages of the respiratory system which make breathing difficult). What’s happening and what is the method of treatment? Explain fully.