Chapter 43

The Immune System

PowerPoint® Lecture Presentations for



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Overview: Reconnaissance, Recognition, and Response

- Barriers help an animal to defend itself from the many dangerous pathogens it may encounter
- The immune system recognizes foreign bodies and responds with the production of immune cells and proteins
- Two major kinds of defense have evolved: innate immunity and acquired immunity

Fig. 43-1



- Innate immunity is present before any exposure to pathogens and is effective from the time of birth
- It involves nonspecific responses to pathogens
- Innate immunity consists of external barriers plus internal cellular and chemical defenses

- Acquired immunity, or adaptive immunity, develops after exposure to agents such as microbes, toxins, or other foreign substances
- It involves a very specific response to pathogens



Concept 43.1: In innate immunity, recognition and response rely on shared traits of pathogens

- Both invertebrates and vertebrates depend on innate immunity to fight infection
- Vertebrates also develop acquired immune defenses

Innate Immunity of Invertebrates

- In insects, an exoskeleton made of chitin forms the first barrier to pathogens
- The digestive system is protected by low pH and lysozyme, an enzyme that digests microbial cell walls
- Hemocytes circulate within hemolymph and carry out phagocytosis, the ingestion and digestion of foreign substances including bacteria



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• Hemocytes also secrete antimicrobial peptides that disrupt the plasma membranes of bacteria

Fig. 43-4



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- The immune system recognizes bacteria and fungi by structures on their cell walls
- An immune response varies with the class of pathogen encountered

Fig. 43-5

RESULTS





Fig. 43-5a

RESULTS



Fig. 43-5b

RESULTS



Innate Immunity of Vertebrates

- The immune system of mammals is the best understood of the vertebrates
- Innate defenses include barrier defenses, phagocytosis, antimicrobial peptides
- Additional defenses are unique to vertebrates: the inflammatory response and natural killer cells

- Barrier defenses include the skin and mucous membranes of the respiratory, urinary, and reproductive tracts
- Mucus traps and allows for the removal of microbes
- Many body fluids including saliva, mucus, and tears are hostile to microbes
- The low pH of skin and the digestive system prevents growth of microbes

- White blood cells (leukocytes) engulf pathogens in the body
- Groups of pathogens are recognized by TLR, Toll-like receptors

Fig. 43-6



- A white blood cell engulfs a microbe, then fuses with a lysosome to destroy the microbe
- There are different types of phagocytic cells:
 - Neutrophils engulf and destroy microbes
 - Macrophages are part of the lymphatic system and are found throughout the body
 - **Eosinophils** discharge destructive enzymes
 - Dendritic cells stimulate development of acquired immunity



Antimicrobial Peptides and Proteins

- Peptides and proteins function in innate defense by attacking microbes directly or impeding their reproduction
- Interferon proteins provide innate defense against viruses and help activate macrophages
- About 30 proteins make up the complement system, which causes lysis of invading cells and helps trigger inflammation

- Following an injury, mast cells release histamine, which promotes changes in blood vessels; this is part of the inflammatory response
- These changes increase local blood supply and allow more phagocytes and antimicrobial proteins to enter tissues
- Pus, a fluid rich in white blood cells, dead microbes, and cell debris, accumulates at the site of inflammation



- Inflammation can be either local or systemic (throughout the body)
- Fever is a systemic inflammatory response triggered by pyrogens released by macrophages, and toxins from pathogens
- Septic shock is a life-threatening condition caused by an overwhelming inflammatory response

- All cells in the body (except red blood cells) have a class 1 MHC protein on their surface
- Cancerous or infected cells no longer express this protein; natural killer (NK) cells attack these damaged cells

Innate Immune System Evasion by Pathogens

- Some pathogens avoid destruction by modifying their surface to prevent recognition or by resisting breakdown following phagocytosis
- Tuberculosis (TB) is one such disease and kills more than a million people a year

Concept 43.2: In acquired immunity, lymphocyte receptors provide pathogen-specific recognition

- White blood cells called lymphocytes recognize and respond to antigens, foreign molecules
- Lymphocytes that mature in the thymus above the heart are called T cells, and those that mature in bone marrow are called B cells

- Lymphocytes contribute to immunological memory, an enhanced response to a foreign molecule encountered previously
- Cytokines are secreted by macrophages and dendritic cells to recruit and activate lymphocytes

Acquired Immunity: An Overview

- B cells and T cells have receptor proteins that can bind to foreign molecules
- Each individual lymphocyte is specialized to recognize a specific type of molecule

Antigen Recognition by Lymphocytes

- An antigen is any foreign molecule to which a lymphocyte responds
- A single B cell or T cell has about 100,000 identical antigen receptors



(a) B cell receptor



(a) B cell receptor



The Antigen Receptors of B Cells and T Cells

- B cell receptors bind to specific, intact antigens
- The B cell receptor consists of two identical heavy chains and two identical light chains
- The tips of the chains form a constant (C) region, and each chain contains a variable (V) region, so named because its amino acid sequence varies extensively from one B cell to another

 Secreted antibodies, or immunoglobulins, are structurally similar to B cell receptors but lack transmembrane regions that anchor receptors in the plasma membrane
- Each T cell receptor consists of two different polypeptide chains
- The tips of the chain form a variable (V) region; the rest is a constant (C) region
- T cells can bind to an antigen that is free or on the surface of a pathogen



- T cells bind to antigen fragments presented on a host cell
- These antigen fragments are bound to cellsurface proteins called MHC molecules
- MHC molecules are so named because they are encoded by a family of genes called the major histocompatibility complex

- In infected cells, MHC molecules bind and transport antigen fragments to the cell surface, a process called antigen presentation
- A nearby T cell can then detect the antigen fragment displayed on the cell's surface
- Depending on their source, peptide antigens are handled by different classes of MHC molecules

Fig. 43-11



- Class I MHC molecules are found on almost all nucleated cells of the body
- They display peptide antigens to cytotoxic T cells



- Class II MHC molecules are located mainly on dendritic cells, macrophages, and B cells
- Dendritic cells, macrophages, and B cells are antigen-presenting cells that display antigens to cytotoxic T cells and helper T cells

- The acquired immune system has three important properties:
 - Receptor diversity
 - A lack of reactivity against host cells
 - Immunological memory

Generation of Lymphocyte Diversity by Gene Rearrangement

- Differences in the variable region account for specificity of antigen receptors
- The *immunoglobulin* (*Ig*) gene encodes one chain of the B cell receptor
- Many different chains can be produced from the same Ig chain gene by rearrangement of the DNA
- Rearranged DNA is transcribed and translated and the antigen receptor formed

Fig. 43-13

DNA of undifferentiated B cell



- Antigen receptors are generated by random rearrangement of DNA
- As lymphocytes mature in bone marrow or the thymus, they are tested for self-reactivity
- Lymphocytes with receptors specific for the body's own molecules are destroyed by apoptosis, or rendered nonfunctional

Amplifying Lymphocytes by Clonal Selection

- In the body there are few lymphocytes with antigen receptors for any particular epitope
- The binding of a mature lymphocyte to an antigen induces the lymphocyte to divide rapidly
- This proliferation of lymphocytes is called clonal selection
- Two types of clones are produced: short-lived activated effector cells and long-lived memory cells



- The first exposure to a specific antigen represents the primary immune response
- During this time, effector B cells called plasma cells are generated, and T cells are activated to their effector forms
- In the **secondary immune response**, memory cells facilitate a faster, more efficient response





Primary immune response to antigen A produces antibodies to A. Secondary immune response to antigen A produces antibodies to A; primary immune response to antigen B produces antibodies to B.



Concept 43.3: Acquired immunity defends against infection of body cells and fluids

- Acquired immunity has two branches: the humoral immune response and the cellmediated immune response
- Humoral immune response involves activation and clonal selection of B cells, resulting in production of secreted antibodies
- Cell-mediated immune response involves activation and clonal selection of cytotoxic T cells
- Helper T cells aid both responses

Fig. 43-16







Cell-mediated immune response



Helper T Cells: A Response to Nearly All Antigens

- A surface protein called CD4 binds the class II MHC molecule
- This binding keeps the helper T cell joined to the antigen-presenting cell while activation occurs
- Activated helper T cells secrete cytokines that stimulate other lymphocytes





Cytotoxic T Cells: A Response to Infected Cells

- Cytotoxic T cells are the effector cells in cellmediated immune response
- Cytotoxic T cells make CD8, a surface protein that greatly enhances interaction between a target cell and a cytotoxic T cell
- Binding to a class I MHC complex on an infected cell activates a cytotoxic T cell and makes it an active killer
- The activated cytotoxic T cell secretes proteins that destroy the infected target cell



Animation: Cytotoxic T Cells



B Cells: A Response to Extracellular Pathogens

- The humoral response is characterized by secretion of antibodies by B cells
- Activation of B cells is aided by cytokines and antigen binding to helper T cells
- Clonal selection of B cells generates antibodysecreting plasma cells, the effector cells of humoral immunity





- The five major classes of antibodies, or immunoglobulins, differ in distribution and function
- Polyclonal antibodies are the products of many different clones of B cells following exposure to a microbial antigen
- Monoclonal antibodies are prepared from a single clone of B cells grown in culture

Class of Immuno- globulin (Antibody)	Distribution	Function
lgM (pentamer)	First Ig class produced after initial exposure to antigen; then its concentration in the blood declines	Promotes neutraliza- tion and cross- linking of antigens; very effective in complement system activation
lgG (monomer)	Most abundant Ig class in blood; also present in tissue fluids	Promotes opsoniza- tion, neutralization, and cross-linking of antigens; less effec- tive in activation of complement system than IgM Only Ig class that crosses placenta, thus conferring passive immunity on fetus
lgA (dimer) J chain Secretory component	Present in secretions such as tears, saliva, mucus, and breast milk	Provides localized defense of mucous membranes by cross-linking and neutralization of antigens Presence in breast milk confers passive immunity on nursing infant
lgE (monomer)	Present in blood at low concen- trations	Triggers release from mast cells and basophils of hista- mine and other chemicals that cause allergic reactions
IgD (monomer) Trans membrane region	Present primarily on surface of B cells that have not been exposed to antigens	Acts as antigen receptor in the antigen-stimulated proliferation and differentiation of B cells (clonal selection)

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IgM (pentamer)	First Ig class produced after initial exposure to antigen; then its concentration in the blood declines	Promotes neutraliza- tion and cross- linking of antigens; very effective in complement system activation

Class of Immuno- globulin (Antibody)	Distribution	Function
IgG (monomer)	Most abundant Ig class in blood; also present in tissue fluids	Promotes opsoniza- tion, neutralization, and cross-linking of antigens; less effec- tive in activation of complement system than IgM
Convright © 2008 Pearson Education. Inc., publishing as Pearson Benjamin Cumm	005.	Only Ig class that crosses placenta, thus conferring passive immunity on fetus



Class of Immuno- globulin (Antibody)	Distribution	Function
lgE (monomer)	Present in blood at low concen- trations	Triggers release from mast cells and basophils of hista- mine and other chemicals that cause allergic reactions

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The Role of Antibodies in Immunity

- Neutralization occurs when a pathogen can no longer infect a host because it is bound to an antibody
- Opsonization occurs when antibodies bound to antigens increase phagocytosis
- Antibodies together with proteins of the complement system generate a *membrane attack complex* and cell lysis





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Viral neutralization



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Fig. 43-21c



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- Active immunity develops naturally in response to an infection
- It can also develop following immunization, also called vaccination
- In immunization, a nonpathogenic form of a microbe or part of a microbe elicits an immune response to an immunological memory

- Passive immunity provides immediate, shortterm protection
- It is conferred naturally when IgG crosses the placenta from mother to fetus or when IgA passes from mother to infant in breast milk
- It can be conferred artificially by injecting antibodies into a nonimmune person

- Cells transferred from one person to another can be attacked by immune defenses
- This complicates blood transfusions or the transplant of tissues or organs

- Antigens on red blood cells determine whether a person has blood type A (A antigen), B (B antigen), AB (both A and B antigens), or O (neither antigen)
- Antibodies to nonself blood types exist in the body
- Transfusion with incompatible blood leads to destruction of the transfused cells
- Recipient-donor combinations can be fatal or safe

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- MHC molecules are different among genetically nonidentical individuals
- Differences in MHC molecules stimulate rejection of tissue grafts and organ transplants

- Chances of successful transplantation increase if donor and recipient MHC tissue types are well matched
- Immunosuppressive drugs facilitate transplantation
- Lymphocytes in bone marrow transplants may cause the donor tissue to reject the recipient

Concept 43.4: Disruption in immune system function can elicit or exacerbate disease

Some pathogens have evolved to diminish the effectiveness of host immune responses

Exaggerated, Self-Directed, and Diminished Immune Responses

 If the delicate balance of the immune system is disrupted, effects range from minor to often fatal

Allergies

- Allergies are exaggerated (hypersensitive) responses to antigens called allergens
- In localized allergies such as hay fever, IgE antibodies produced after first exposure to an allergen attach to receptors on mast cells



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- The next time the allergen enters the body, it binds to mast cell-associated IgE molecules
- Mast cells release histamine and other mediators that cause vascular changes leading to typical allergy symptoms
- An acute allergic response can lead to anaphylactic shock, a life-threatening reaction that can occur within seconds of allergen exposure

- In individuals with autoimmune diseases, the immune system loses tolerance for self and turns against certain molecules of the body
- Autoimmune diseases include systemic lupus erythematosus, rheumatoid arthritis, insulindependent diabetes mellitus, and multiple sclerosis

Exertion, Stress, and the Immune System

- Moderate exercise improves immune system function
- Psychological stress has been shown to disrupt hormonal, nervous, and immune systems

- Inborn immunodeficiency results from hereditary or developmental defects that prevent proper functioning of innate, humoral, and/or cell-mediated defenses
- Acquired immunodeficiency results from exposure to chemical and biological agents
- Acquired immunodeficiency syndrome (AIDS) is caused by a virus

Acquired Immune System Evasion by Pathogens

 Pathogens have evolved mechanisms to attack immune responses

- Through antigenic variation, some pathogens are able to change epitope expression and prevent recognition
- The human influenza virus mutates rapidly, and new flu vaccines must be made each year
- Human viruses occasionally exchange genes with the viruses of domesticated animals
- This poses a danger as human immune systems are unable to recognize the new viral strain



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- Some viruses may remain in a host in an inactive state called latency
- Herpes simplex viruses can be present in a human host without causing symptoms

- Human immunodeficiency virus (HIV) infects helper T cells
- The loss of helper T cells impairs both the humoral and cell-mediated immune responses and leads to AIDS
- HIV eludes the immune system because of antigenic variation and an ability to remain latent while integrated into host DNA



Animation: HIV Reproductive Cycle

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Fig. 43-26



- People with AIDS are highly susceptible to opportunistic infections and cancers that take advantage of an immune system in collapse
- The spread of HIV is a worldwide problem
- The best approach for slowing this spread is education about practices that transmit the virus

- The frequency of certain cancers increases when the immune response is impaired
- Two suggested explanations are
 - Immune system normally suppresses cancerous cells
 - Increased inflammation increases the risk of cancer

- 1. Distinguish between innate and acquired immunity
- 2. Name and describe four types of phagocytic cells
- 3. Describe the inflammation response

- 4. Distinguish between the following pairs of terms: antigens and antibodies; antigen and epitope; B lymphocytes and T lymphocytes; antibodies and B cell receptors; primary and secondary immune responses; humoral and cell-mediated response; active and passive immunity
- 5. Explain how B lymphocytes and T lymphocytes recognize specific antigens
- 6. Explain why the antigen receptors of lymphocytes are tested for self-reactivity

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- 7. Describe clonal selection and distinguish between effector cells and memory cells
- 8. Describe the cellular basis for immunological memory
- 9. Explain how a single antigen can provoke a robust humoral response
- 10. Compare the processes of neutralization and opsonization

- 11. Describe the role of MHC in the rejection of tissue transplants
- 12. Describe an allergic reaction, including the roles of IgE, mast cells, and histamine
- 13. Describe some of the mechanisms that pathogens have evolved to thwart the immune response of their hosts
- 14. List strategies that can reduce the risk of HIV transmission