

? What is Immunity ?



- Immunity = the protection against infectious disease conferred either by the immune response generated by immunization or previous infection or by other nonimmunologic factors...a.k.a. body's ability to resist infection

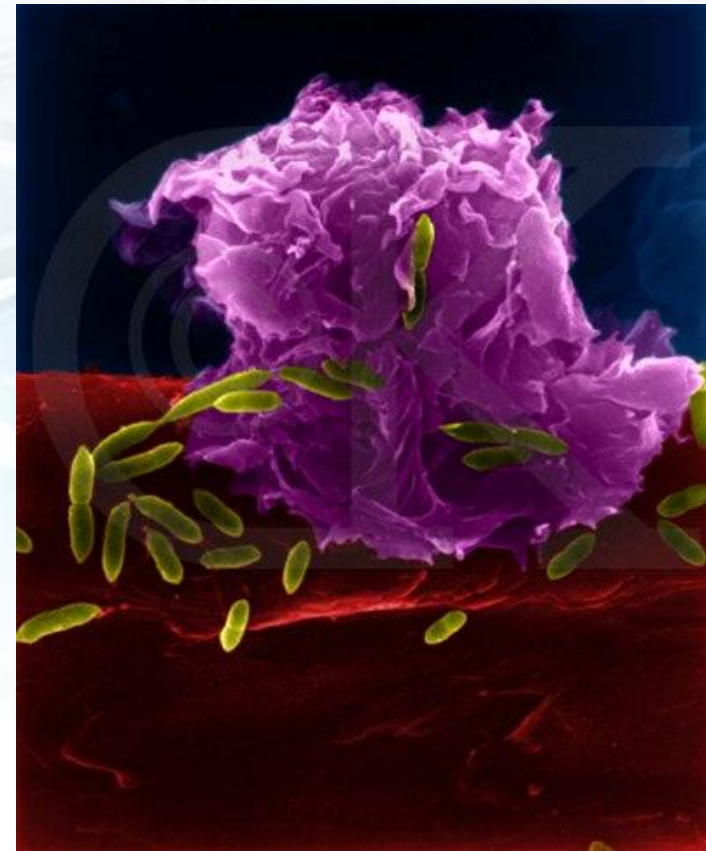
- 2 types of immunity
 - Non-specific (innate)
 - Specific (adaptive, acquired)



Innate Immunity



- Skin
 - What characteristics of the skin make it an effective mechanical barrier ?
- Mucous membranes (mechanical)
 - Cilia in respiratory tract
 - Lysozymes, pH
- Iron-binding proteins
 - Some bacteria require iron for growth
 - Transferrin, lactoferrin
- Phagocytosis
 - PMNs, monocytes and macrophages
- Complement



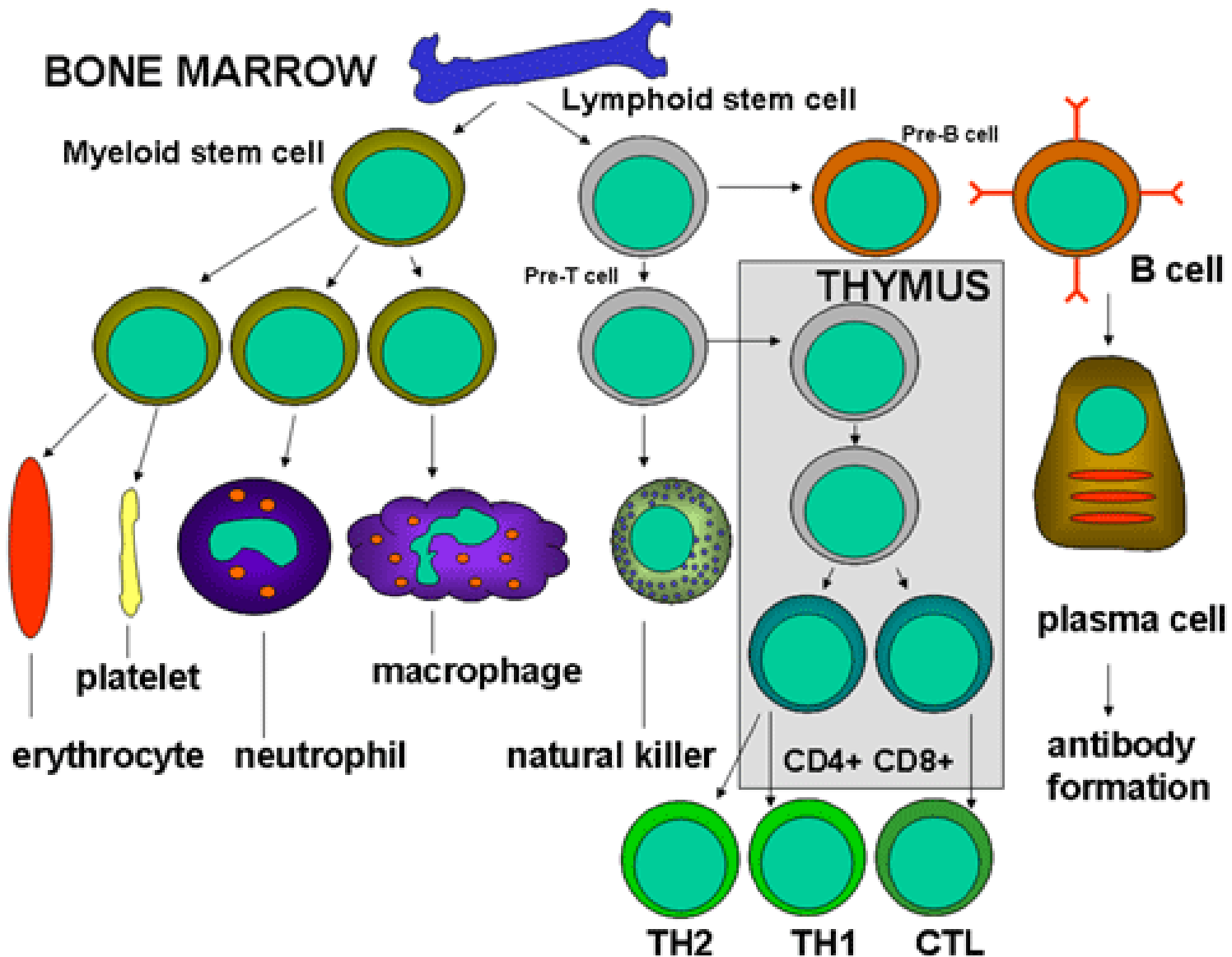
Specific Immunity



- Humoral and Cell-Mediated (CMI)
- What is the difference between innate immunity and adaptive immunity?
 - **Innate**: protects against ANY invader, does not discriminate
 - **Adaptive**: directed against one type of invader, dependant on past exposure



Where do immune cells come from?



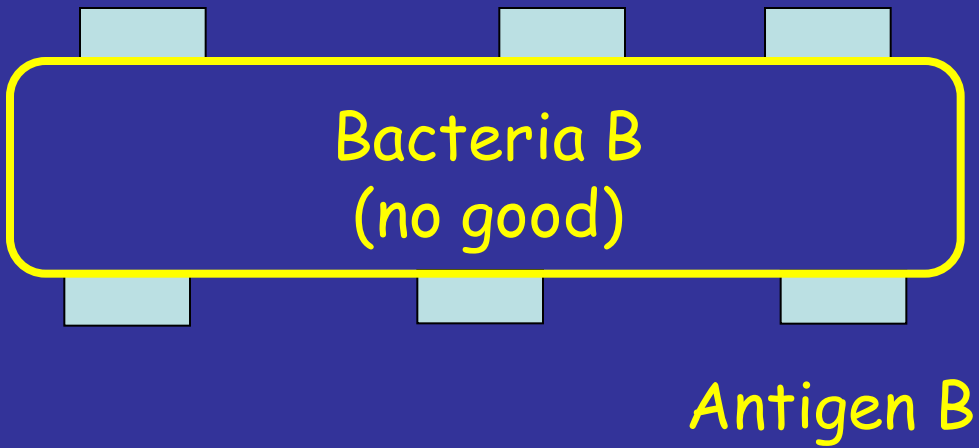
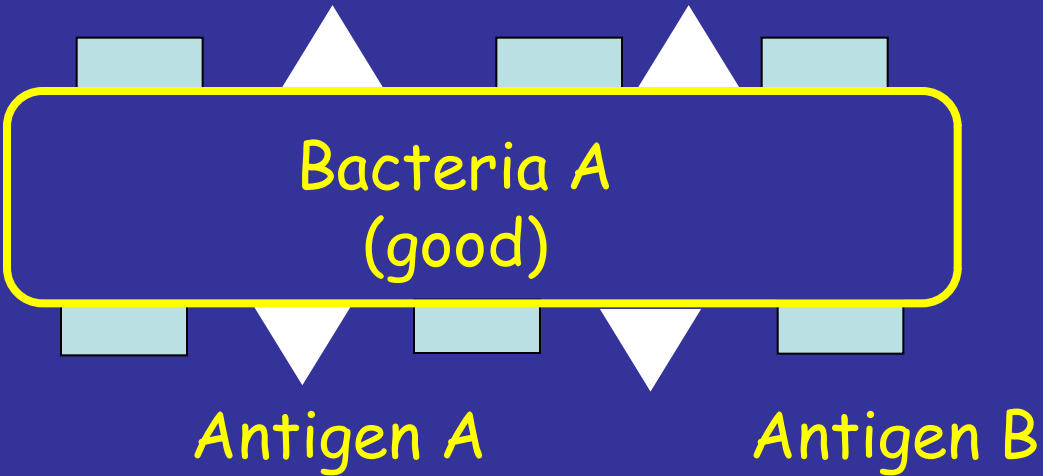
Humoral Immunity

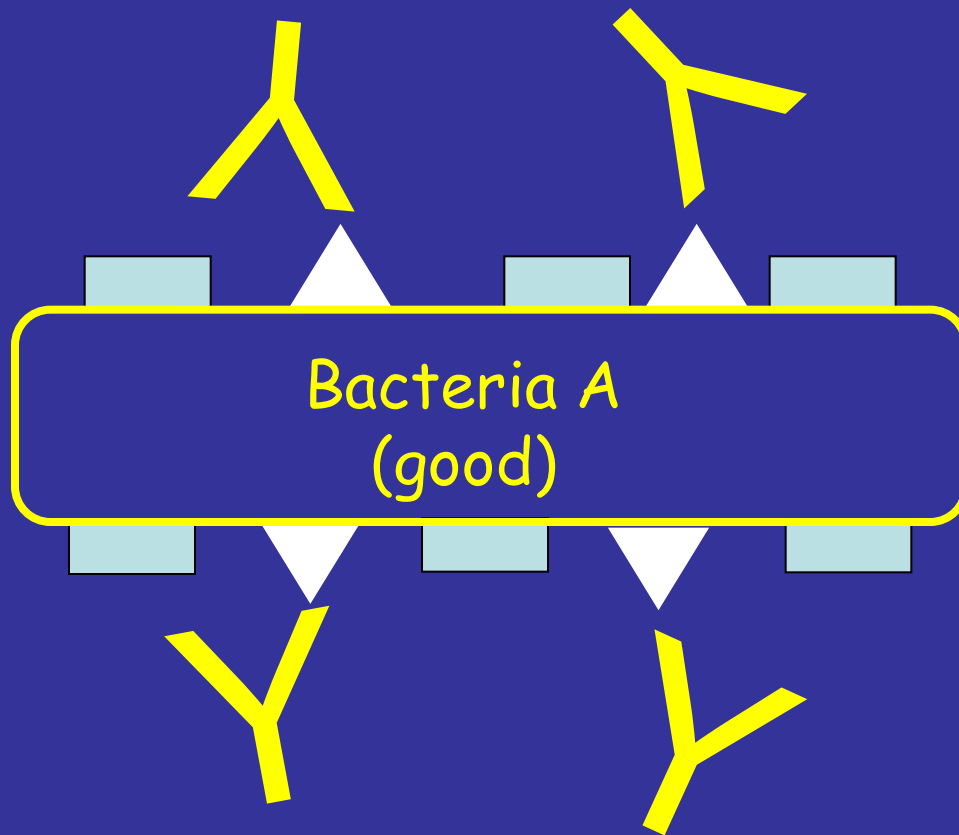


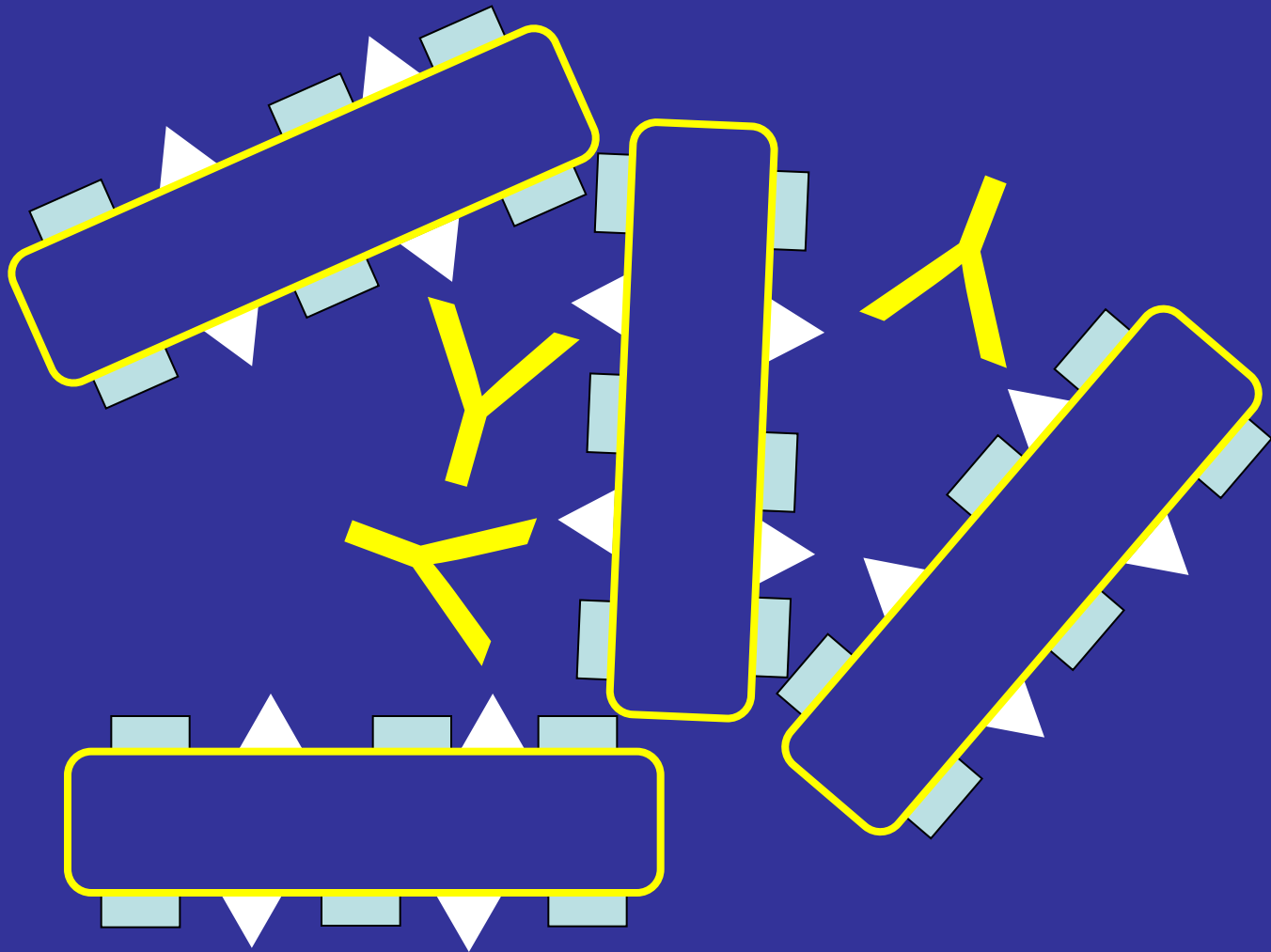
- Circulating antibodies
- **Antibody**: protein that binds specifically to a substance (its antigen)
 - Igs or immunoglobulins
 - Produced by B-lymphocytes upon stimulation from antigen presenting T-cells
 - Recognize toxins, capsules, some viral proteins
- **Antigen**
 - “non-self”
 - Protein, glycoprotein, lipoprotein, polysaccharide
 - What structures could be “antigenic” in a bacteria? Virus?



Antibody Binding: how does it occur?







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Immunoglobulins (Igs) a.k.a. Antibodies



- **Antibody:** Ig produced in response to stimulation by an antigen and reacting *specifically* with it.
- Distinguish “non-self” from “self”
- Constant and variable region
 - Variable region is responsible for antigen recognition



Figure 3-1 part 1 of 3 Immunobiology, 6/e. (© Garland Science 2005)

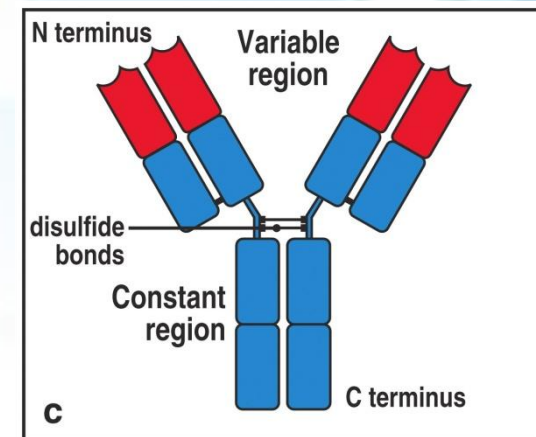


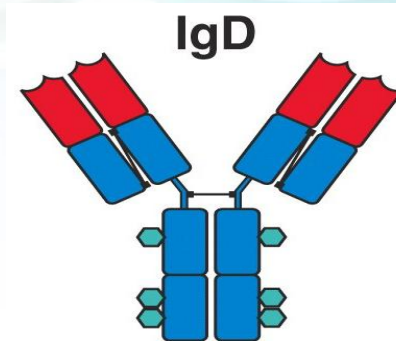
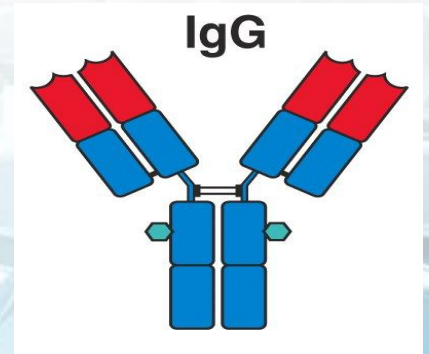
Figure 3-1 part 3 of 3 Immunobiology, 6/e. (© Garland Science 2005)



Classes of Igs



- 5 classes: IgG, IgA, IgM, IgE, IgD
- **IgG**
 - Host defense
 - Crosses placenta and protects newborn
- **IgD**
 - Role is unknown



Classes of Igs



- **IgA**
 - Host defense
 - Found in secretions
 - Tears, saliva, milk, respiratory, GI and genito-urinary tract
 - Dimer

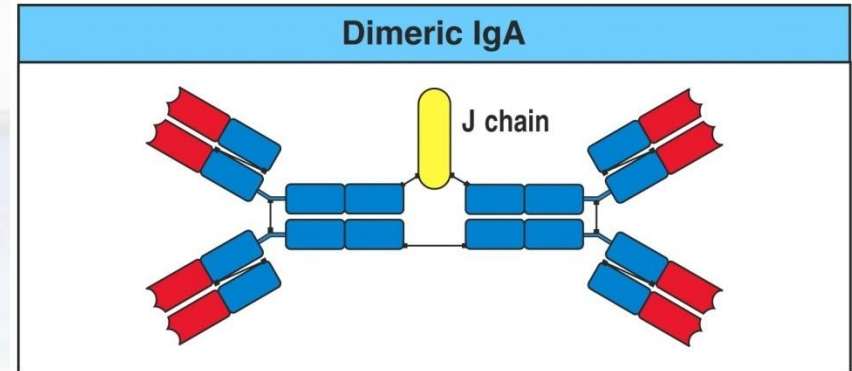


Figure 4-23 part 2 of 3 Immunobiology, 6/e. (© Garland Science 2005)

- **IgM**
 - Host defense
 - Early immune response
 - Pentamer
- **IgE**
 - Hypersensitivity (allergies)
 - Defends against parasites

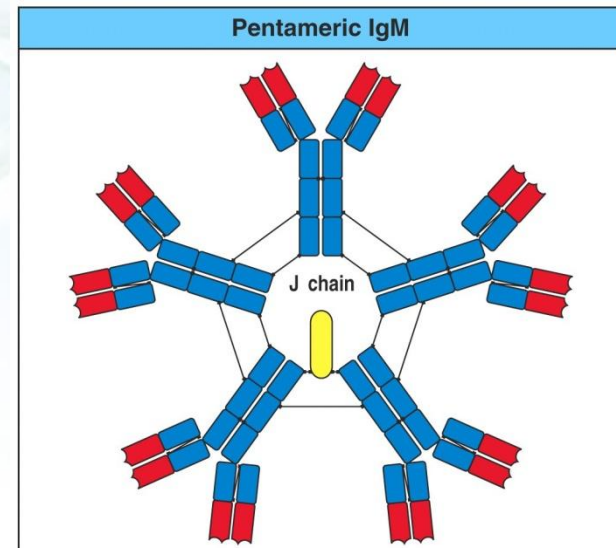


Figure 4-23 part 1 of 3 Immunobiology, 6/e. (© Garland Science 2005)

1^o and 2^o Immune Response



- **1^o Response**
 - Ab production triggered on first antigen introduction
 - Latent period of several days
 - Circulating antibody detectable after 5-10 days
 - Antibody in serum is maximum at ~21 days, then drops to low levels
- **2^o Response**
 - ★Basis for Immunizations★
 - Occurs when Ab is introduced 2nd, 3rd, 4th ...time
 - Lag, rapid Ab increase (2-3 days), slow decrease
 - Booster injections to maximize Ab levels



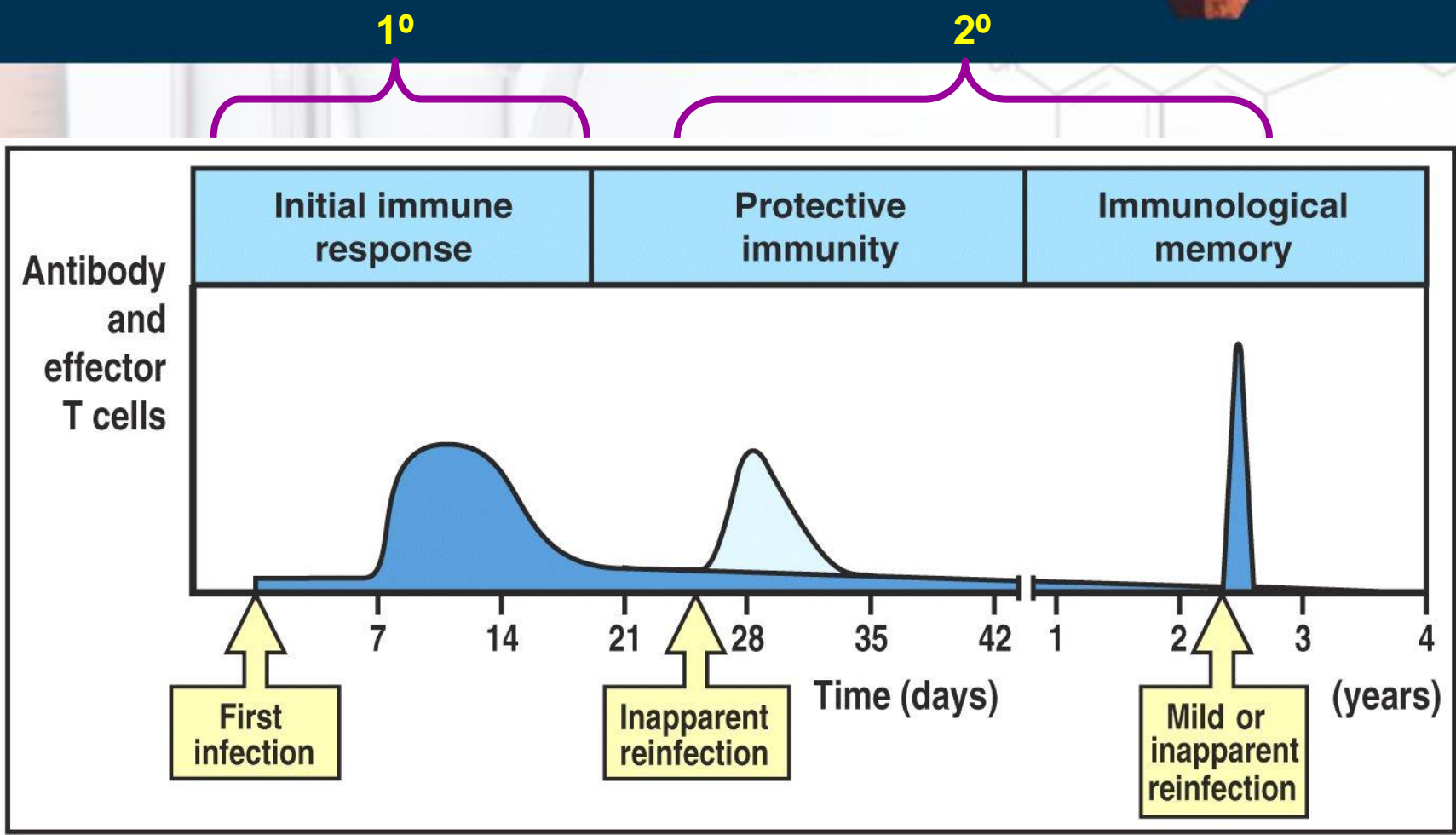


Figure 10-18 Immunobiology, 6/e. (© Garland Science 2005)

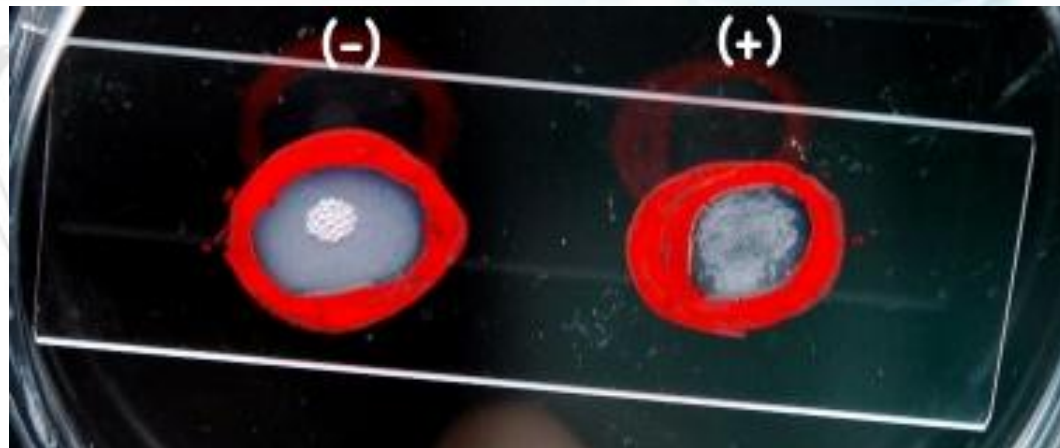


Antibody Detection



- **Serological Reaction**

- Detects presence of antibodies in serum sample
- Antigen and antibody interact; agglutination
- Antibody titration
- Detect unknown microorganisms using known antisera



So Far.....



- Innate Immunity ✓
- Adaptive Immunity
 - Humoral (Antibody) ✓
 - Cell-Mediated (Next)

Cell-Mediated Immunity (CMI)



- T-cells **NOT** antibodies!
 - Helper, suppressive, cytotoxic (killer) generated from memory T-cells
- Exposure to antigen induces response from trained T-cells
- Essential for defense against intracellular organisms, parasites, tumors and other foreign cells (i.e., transplants, grafts)
- Immune-suppressive medication for transplant recipients



Disorders of Immunity



1 Allergy and Hypersensitivity

- OVER-reaction to antigens in absence of true infection
- Can be fatal.....ANAPHYLAXIS

2 Auto-immune diseases

- Immune system reacts to its own “self” antigens
- “auto-antibodies”
- Type I diabetes, MS, rheumatoid arthritis, lupus

3 Immunodeficiency states

- Inability to produce antibodies and/or dysfunctional CMI
- Congenital, disease, AIDS

4 Graft rejection

- NORMAL immune reaction to “non-self”
- Control by immune-suppressive medication

Immunization



- **Passive Immunization**

- administration of pre-formed antibody against a *specific* microbial agent
- IgG animal origin: short lived, risk of hypersensitivity reaction
- IgG human origin: short lived, no risk of reaction
- **Gamma globulin (IgG)**: pooled from large grouped of blood donors and has antibodies to many common infections
- **Hyperimmune globulins (IgG)**: specific for a particular microbe



Active Immunization



- Stimulates immune system by administration of antigen
- LONGER LASTING
- **Live-attenuated vaccine**
 - Sub-clinical or mild illness mimicking the disease
 - Local (IgA) and humoral (IgG) immunity
 - Rapid immunity development
 - Serious illness in immuno-compromised individuals



Active Immunization (con't)



- **Killed vaccines, sub-unit vaccines and toxoids**
 - Antigen without infectivity
 - May require boosters
 - Adjuvant with toxoids
 - Polysaccharide vaccines can be conjugated to protein (see conjugate vaccines)
- **Recombinant vaccines**
 - DNA recombinant technology
 - Attenuates microorganism
 - Hep B vaccine
- **Adsorbed vaccines**
 - Vaccine mixed with inorganic salt for slower adsorption and longer-lasting immunity
 - Tetanus, diphtheria

Active Immunization (con't)



- **Conjugate vaccines**
 - Designed for poorly antigenic microorganisms
 - Conjugate antigen of interest to immunogenic, non-toxic protein
 - *Haemophilus influenzae* type b
- **Combined vaccines**
 - For ease of administration
- **Combined Active-Passive Immunization**
 - Immediate protection after possible exposure to microbe
 - Hyperimmune Igs and vaccine injected at DIFFERENT sites
 - Tetanus, Rabies, Hep B



Canada's Immunization Guide:

<http://www.phac-aspc.gc.ca/publicat/cig-gci/index.html>

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Immunization
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Bureau of Microbial Hazards

BMH  **BDM**

Bureau des dangers microbiens



Antibiotic Resistance



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Introduction



- The first antibiotic (?)
 - discovered in 1929 by Sir Alexander Fleming
- World War II
 - penicillin used to treat staphylococci and streptococci (1946)
- How effective was penicillin?

Introduction



- Resistance to penicillin recognized almost immediately
 - 80% of all strains of *Staphylococcus aureus*
 - *Streptococcus pyogenes* (Group A strep) still treated with penicillin
 - Interestingly, penicillin has never been effective against Gm-negatives (*Salmonella*, *Shigella*, *Bordetella pertussis*, *Yersinia pestis*, *Pseudomonas*) – why? 😊
- Late 1940s and early 1950s?

Antibiotic therapy



- Effective chemotherapy depends on selective toxicity
 - good against pathogen, does not affect host... 😊
- Exploit pathogen processes not seen in humans
 - cell wall, metabolism, etc.
- Knowledge of likely microorganism is crucial...
 - site
 - organism
 - allergy to host?

Antibiotic therapy



- Other considerations...
 - route of administration
- Monitoring therapy
- Adverse effects
 - GI-tract, skin, haemopoietic system, renal system, liver

Acquired resistance



- Three major mechanisms of resistance
 - Alteration in drug target
 - Production of inactivating enzymes
 - Decreased uptake of antibiotic

Antibiotic resistance



- Resistance occurs when a susceptible microorganism is no longer inhibited by an antibiotic agent
- Many reasons why this can happen
 - **intrinsic** - characteristics of microorganism vis-à-vis antibiotic's mechanism of action (inherent or “natural”)
 - **acquired** - new or added (driven by two genetic processes in bacteria...mutation and selection (vertical evolution); and exchange of genetic material (horizontal evolution))

The chromosome: role in antibiotic resistance...



- Mutations lead to
 - Change it site of antibiotic target (but protein for bacterial still works fine!)
 - Regulatory genes
 - turn on alternative path
 - turn on efflux mechanisms
 - Change cell permeability



Post-antibiotic era: is it possible?



- With current overuse of antibiotics, we are *forcing* bacteria to change (evolve) in order to survive
- How is this achieved/helped by us?

Decreasing antimicrobial resistance?



- Withhold antibiotics
 - self-limited viral infections (i.e., the “common cold”)
- Use narrowest spectrum antimicrobial agents
- Base decision about broadness of empiric antibiotic coverage on severity of illness
 - clinically stable and not at risk for significant morbidity... may be appropriate to wait culture results and MIC testing

Decreasing antimicrobial resistance?



- Prevention of infection
 - hygiene, **handwashing**
- Education
 - helps to achieve therapeutic and preventative goals
 - when are antibiotics needed?
 - how to take them?
 - **proper duration!!**
- Earlier detection of therapeutic failure
 - good for patients with antibiotic-resistant pathogens