Medical Immunology



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Lecture 7

Antibodies and antigens

In this lecture we will discuss:

• Antibodies, their structure, types and function.

Antibodies/ overview

- The study of antibodies and their reactions with antigens is therefore classically called serology.
- Serum lacks coagulation factors but otherwise contains all the proteins found in plasma. Any serum sample that contains detectable antibody molecules that bind to a particular antigen is commonly called an antiserum.
- A healthy 70-kg adult human produces about 2 to 3 g of antibodies every day.



 Plasma proteomics holds great promise for the future of biomarker discovery, as well as in vitro diagnostics. Although plasma is readily accessible for analysis, the study of the plasma proteome is fundamentally limited by its vast dynamic range (10 orders of magnitude).



Antibodies and antigens

- Antibodies are circulating proteins that are produced in vertebrates in response to exposure to foreign structures known as antigens
- The strength of the binding between a single site of an antibody and an epitope of an antigen is called the affinity of the antibody.



Antibodies and antigens

Because the **hinge** region of antibodies gives them flexibility, a single antibody may attach to a single multivalent antigen **by more** than one binding site. the strength of attachment of the antibody to the antigen must take into account binding of all the sites to all the available epitopes. This overall strength of attachment is called the avidity.



FIGURE 5-13 Valency and avidity of antibody-antigen interactions. Monovalent antigens, or epitopes spaced far apart on cell surfaces, will interact with a single binding site of one antibody molecule. Although the affinity of this interaction may be high, the overall avidity may be relatively low. When repeated determinants on a cell surface are close enough, both the antigen-binding sites of a single IgG molecule can bind, leading to a higher avidity bivalent interaction. The hinge region of the IgG molecule accommodates the shape change needed for simultaneous engagement of both binding sites. IgM molecules have 10 identical antigen-binding sites that can theoretically bind simultaneously with 10 repeating determinants on a cell surface, resulting in a polyvalent, high-avidity interaction.

- Antibodies (immunoglobulins) are made of 2 heavy chains and 2 light chains, those chains combined give us an antibody binding region (Fab) and the fragment crystallizable region (Fc region) which is the tail region of an antibody that interacts with **cell surface receptors** called Fc receptors and some proteins of the **complement** system.
- Immunoglobulins are divided into different classes (isotypes).





- Antibodies can exist in two forms: membrane-bound antibodies on the surface of B lymphocytes function as receptors for antigen, and secreted antibodies that reside in the circulation, tissues, and mucosal sites
- Both heavy chains and light chains consist of amino terminal variable (V) regions that participate in antigen recognition and carboxyl-terminal constant (C) regions; the C regions of the heavy chains mediate effector functions.



- Question: How many polypeptide chains make up a typical antibody molecule?
- a) 1
- b) 2
- c) 3 • d) 4

 Most of the sequence differences and variability among different antibodies are confined to three short stretches in the V region of the heavy chain and to three stretches in the V region of the light chain. These diverse stretches are known as hypervariable segments, Because these sequences form a surface that is complementary to the three-dimensional structure of the bound antigen, the hypervariable regions are also called complementaritydetermining regions (CDRs).



Antibodies

- There are five immunoglobulin classes (isotypes) of antibody molecules found in serum: IgG, IgM, IgA, IgE and IgD. They are distinguished by the type of heavy chain they contain
- Antibodies of different classes differ in their location around the body, appear at different stages of an adaptive immune response.
- The heavy chain C regions of all antibody molecules of one isotype or subtype have essentially the same amino acid sequence. This sequence is different in antibodies of other isotypes or subtypes.



TABLE 5–2 Human Antibody Isotypes										
lsotope of Antibody	Subtypes (H Chain)	Serum Concentration (mg/mL)	Serum Half-life (days)	Secreted Form		Functions				
IgA	lgA1,2 (α1 or α2)	3.5	6	IgA (dimer) Monomer, dimer, trimer	Cα1 Cα2 Cα3 J chain	Mucosal immunity				
IgD	None (δ)	Trace	3	None		Naive B cell antigen receptor				
IgE	None (ε)	0.05	2	lgE Monomer	C_{1} C_{2} C_{2} C_{2} C_{2} C_{2} C_{2} C_{2} C_{2} C_{2}	Defense against helminthic parasites, immediate hypersensitivity				
IgG	lgG1-4 (γ1, γ2, γ3, or γ4)	13.5	23	lgG1 Monomer		Opsonization, complement activation, antibody- dependent cell-mediated cytotoxicity, neonatal immunity, feedback inhibition of B cells				
lgM	None (μ)	1.5	5	lgM Pentamer	Сµ1 Сµ3 Сµ3 Сµ4 Сµ4 Сµ4 Сµ4 Сµ4 Сµ4 Сµ4 Сµ4 Сµ4 Сµ4	Naive B cell antigen receptor, complement activation				

- Question: Which is the most abundant Ig class in serum?
- a) IgG
- b)lgA
- c) IgM
- d) IgE

- B lymphocytes are the cells responsible for antibody responses.
- During its development, each B-lymphocyte becomes genetically programmed through a series of gene-splicing reactions to produce an antibody molecule with a unique specificity
- It is estimated that the human body has the ability to recognize 10⁷ or more different epitopes, due to the wide range of possible combinations during gene splicing.





B Cell Activation

- Resting B cells become activated by antigen via binding to the BCR and internalization.
- Once internalized inside the B cell, the protein antigen is processed and presented with MHC II. The presented antigen is then recognized by helper T cells specific to the same antigen.
- Once activated by linked recognition, T-cells produce and secrete cytokines that activate the B cell and cause proliferation into clonal daughter cells.
- After several rounds of proliferation, additional cytokines provided by the T-cells stimulate the differentiation of activated B cell clones into memory B cells, which will quickly respond to subsequent exposures to the same antigen, and plasma cells that lose their membrane BCRs and initially secrete pentameric IgM





Antibodies/ Antibody Production

- Initial contact with a new antigen evokes the primary response, which is characterized by a lag phase of approximately 1 week between the challenge and the detection of circulating antibodies.
- Once antibody is detected in serum, the **levels rise exponentially** to attain a maximal steady state in approximately 3 weeks, **then decline gradually** with time.
- The **first antibodies** synthesized in the primary immune response **are IgM** and, **then IgG** antibodies arise and eventually predominate.

- After a subsequent exposure or booster injection of the same antigen, a different sequence called the **secondary response** ensues.
- In the secondary response;
- the lag time between the immunization and the appearance of antibody is shortened,
- the rate of exponential increase to the maximum steady-state level is more rapid,
- and the steady-state level itself is higher, representing a larger amount of antibody.
- Another key factor of the secondary response is that the antibodies formed are predominantly of the IgG class.



Antibodies/ Class switching

- After initial secretion of IgM, cytokines secreted by T- cells stimulate the plasma cells to switch from IgM production to production of IgG, IgA, or IgE.
- This process, called class switching or isotype switching, allows plasma cells cloned from the same activated B cell to produce a variety of antibody classes with the same antigen specificity.
- Class switching is accomplished by genetic rearrangement of gene segments encoding the constant region, which determines an antibody's class. The variable region is not changed, so the new class of antibody retains the original antigen specificity.

Class switch

- DNA rearrangement changing the heavy chain constant gene in memory cells.

	300 V Variable	10 D Diversity	4 J Joining	μ	δ	γ3	¥1	γ2b	γ2a	3	α	
C												\supset
	Genes coding	variable ro	egion	Ge Igi	nes 1 Ig	cod D	ing Ig	cor G	istar	nt re IgE	gions IgA	
	B Cell IgM											\supset
	Class switch to Ig6											
	Class switch to IgA									C		

Antibodies and antigens

- The ability of antibodies in any individual to specifically bind a large number of different antigens is a reflection of antibody diversity, and the total collection of antibodies with different specificities represents the antibody repertoire. The genetic mechanisms that generate such a large antibody repertoire occur exclusively in lymphocytes. This diversity is generated by random recombination of a limited set of inherited germline DNA sequences to form functional genes that encode the V regions of heavy and light chains as well as by the addition of nucleotide sequences during the recombination process.
- Somatic mutation in antigen-stimulated B lymphocytes that generates new V domain structures, some of which bind the antigen with greater affinity than did the original V domains. Those B cells producing higher affinity antibodies preferentially bind to the antigen and, as a result of selection, become the dominant B cells with each subsequent exposure to the antigen. This process is called affinity maturation.





Effector mechanisms of humoral immunity



Antibodies/ IgM

- Because of its many specific binding sites, IgM is particularly effective in agglutinating particles carrying antigens against which it is directed.
- IgM is particularly active in bringing about complement-mediated cytolytic damage to foreign antigen-bearing cells.



Antibodies/ IgG

- Immunoglobulin G (IgG) is the most abundant immunoglobulin in health and provides the most extensive and long-lived antibody response to the various microbial and other antigens.
- IgG antibody is characteristically formed in large amounts during the secondary response to an antigenic stimulus, and usually follows production of IgM in the course of a viral or bacterial infection.
- IgG is the only immunoglobulin class able to **cross the placental barrier** and, thus, provides passive immune protection to the newborn in the form of maternal antibody.



Antibodies/ IgA

- Immunoglobulin A (IgA) has a special role as a major determinant of so-called local immunity in protecting epithelial surfaces from colonization and infection.
- At the epithelia, two IgA molecules combine with another protein, termed the secretory piece, which is present on the surface of local epithelial cells. The complex, then termed secretory IgA (slgA),
- The major role of slgA is to prevent attachment of antigen-carrying particles to receptors on mucous membrane epithelia





Antibodies/ IgE

 IgE not only provides protective immunity against helminth parasites but can also mediate the type I hypersensitivity reactions that contribute to the pathogenesis of allergic diseases such as asthma, allergic rhinitis and atopic dermatitis.



Antibodies/ IgD

- IgD (IgD) is a monomeric antibody isotype that is expressed in the plasma membranes of immature Blymphocytes. **IgD** is also produced in a secreted form that is found in small amounts in blood serum.
- the function of IgD is to signal the B cells to be activated.



https://www.jacksonimmuno.com/secondary-antibody-resource/technicaltips/immunoglossary/#:~:text=Complementarity%20Determining%20Regions%20(CDRs)&text=An%20IgG%2 0antibody%20has%2012,each%20of%20the%20heavy%20chains.

Further reading:

• Cellular and Molecular Immunology. 7th Edition.. Chapter 5. Antibodies and antigens