Basic Immunology

Lecture 23-24th

Immunological tolerance

Physiological and pathological autoimmunity.

Autoimmune diseases.

Immune tolerance

The first study in dizygotic twin cattle with a common placenta and shared mixture of each other's red blood cells described the life time tolerance between each other. R.E. Billingham and Peter Medawar in 1953 injected foreign cells into fetal or neonatal mice, and they could become accepting of future grafts from the same foreign donor. Theories of immune tolerance formulated by Sir Frank McFarlane Burnet and Frank Fenner, who were the first to propose the deletion of self-reactive <u>lymphocytes to establish tolerance</u>. Burnet and Medawar were ultimately credited for "the discovery of acquired immune tolerance" and awarded with Nobel Prize in Physiology or Medicine in 1960. In their Nobel Lecture, Medawar and Burnet define immune tolerance as "a state of indifference or nonreactivity towards a substance that would normally be expected to excite an immunological response."

Cognitive paradigm of the immune tolerance

- In 1991, *Irun Cohen* proposed revising the clonal selection theory and replacing it with the cognitive paradigm.
- The natural autoimmune network is the *immunological homunculus*, which is the immune system's representation of the body.
- The particular self-reactivity comprising the homunculus could serve as a set of naturally occurring (auto)antibodies and cellular components of the natural immune system that act as biomarkers to help initiate and regulate tolerance and targeting-type immune response.

TOLERANCE & AUTOIMMUNITY

- Upon encountering an antigen, the immune system can either develop an immune response or enter a state of unresponsiveness called tolerance.
- Immunological tolerance is thus the lack of ability to mount an immune response to epitopes to which an individual has the potential to respond.
- Targeting type and tolerating type immune responses composed by the same cellular and molecular components, the differences are in the effector phase only.
- Targeting type immune response or tolerance needs to be <u>carefully regulated</u> since an inappropriate response – whether it be <u>autoimmune</u> reaction to self-antigens or tolerance to a potential pathogen – can have serious and possibly life-threatening immunodefficiencies.

TOLERANCE

- PASSIVE
- ACTIVE

AUTOIMMUNITY

- PHYSIOLOGIC REGULATION
- AUTOIMMUNE DISEASES

Mechanisms of the immunological tolerance

Central tolerance

- T cell mediated
- B cell mediated

Peripheral tolerance

- unresponsiveness
- local microenvironment mediated

Passive tolerance

Unresponsiveness: no MHC recognition or inhibited cellular differentiation.

- Tolerance induced by the nature of the antigen
- Tolerance induced by the body

Passive tolerance induced by the nature of the antigen

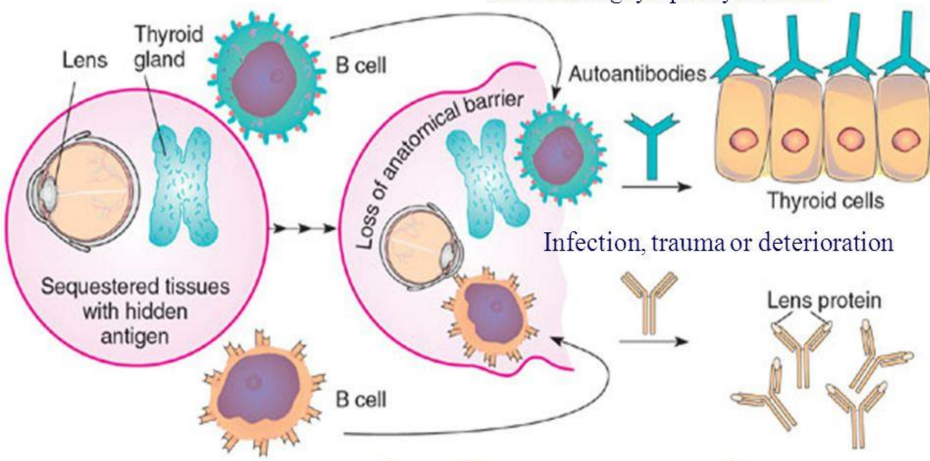
- chemical nature
- dose of the antigen
 - low dose tolerance
 (dominantly T cell mediated, long ranging)
 - high dose tolerance (B and T cell mediated, short ranging)
- mode of the administration

Tolerance induced by the body

- sequestered antigens
 no MHC recognition
 no antigen presentation
 no systemic response
- heredited or acquired immunodeficiency
- clonal anergies

Sequestered Antigen Theory Sequestered behind anatomical barriers

Copyright © The McGraw-Hill Companies, Inc. F Self reacting lymphocyte clones



(a) Sequestered Antigen Theory

Some tissues are not scanned by the immune system during embryonic growth. CNS, lens, thyroid & testes

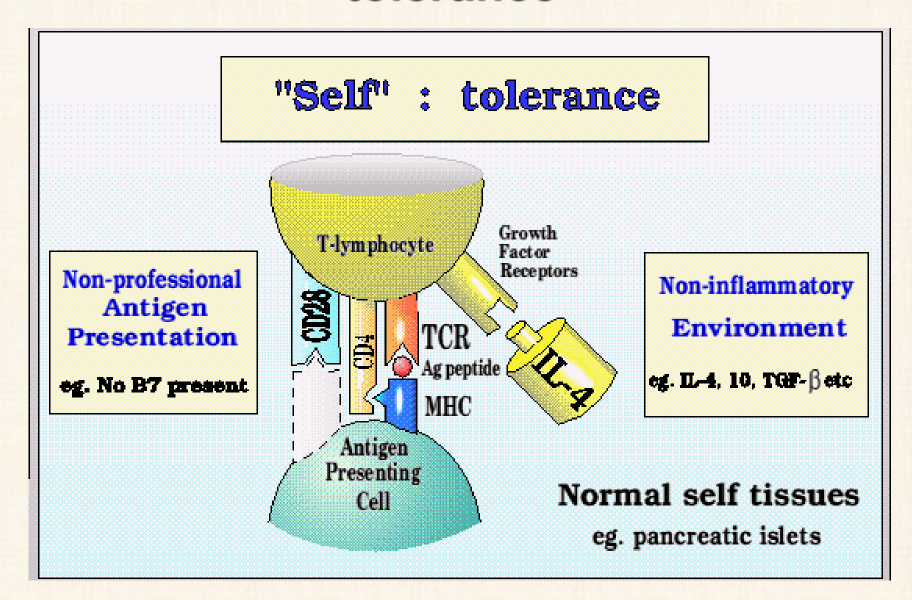
T-cell tolerance

Central Tolerance (selection in the Thymus)

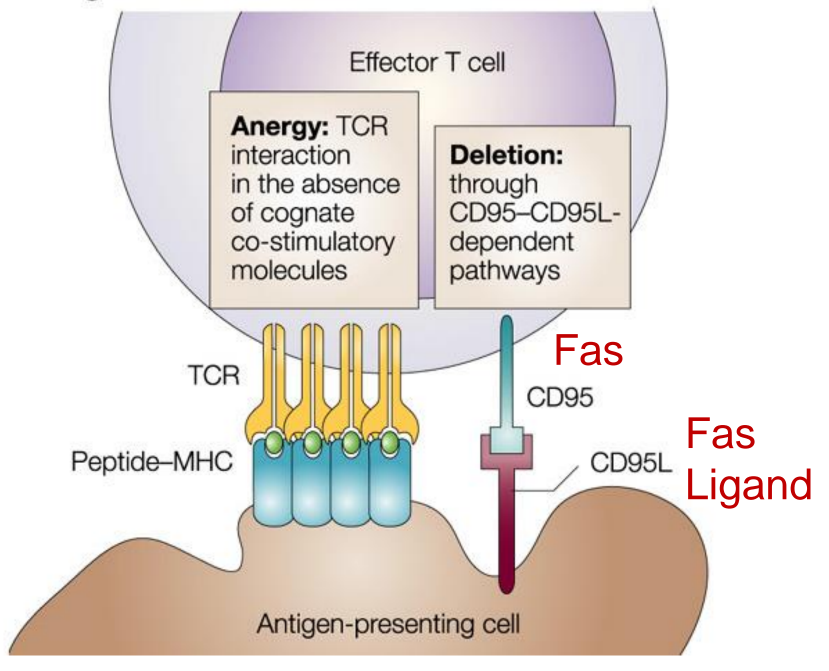
Peripheral Tolerance

- Lack of co-stimulation
- Receipt of death signal (high dose of antigen)
- Control by regulatory T cells

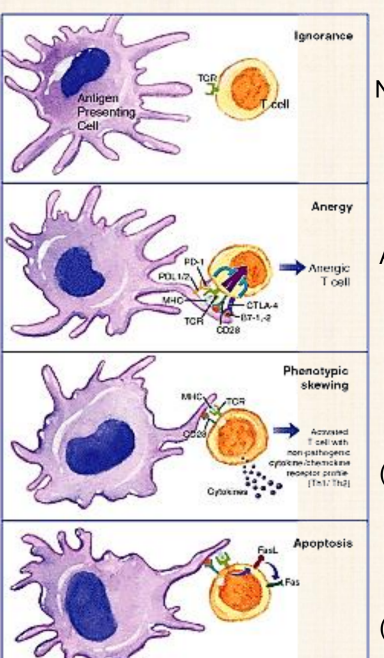
Failed co-stimulation results low dose tolerance



High-dose tolerance



Peripheral T cell tolerance

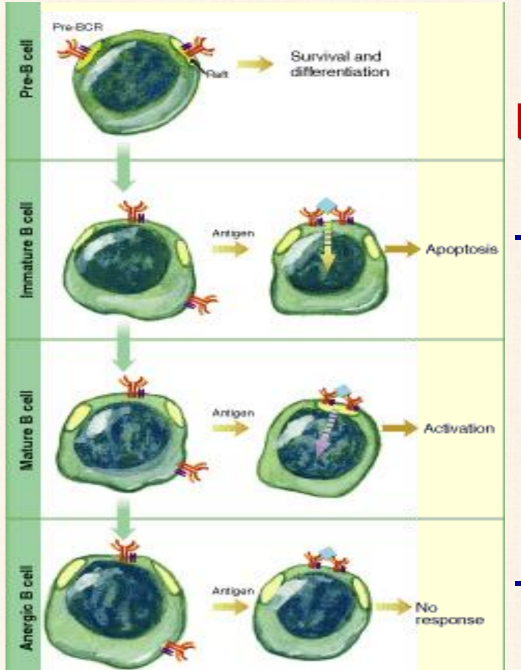


No response (Sequestred antigens)

Anergy
(Suppression by CTLA-4)

Anergy (Th1/Th2 cross regulation)

Deletion
(High dose of antigen)

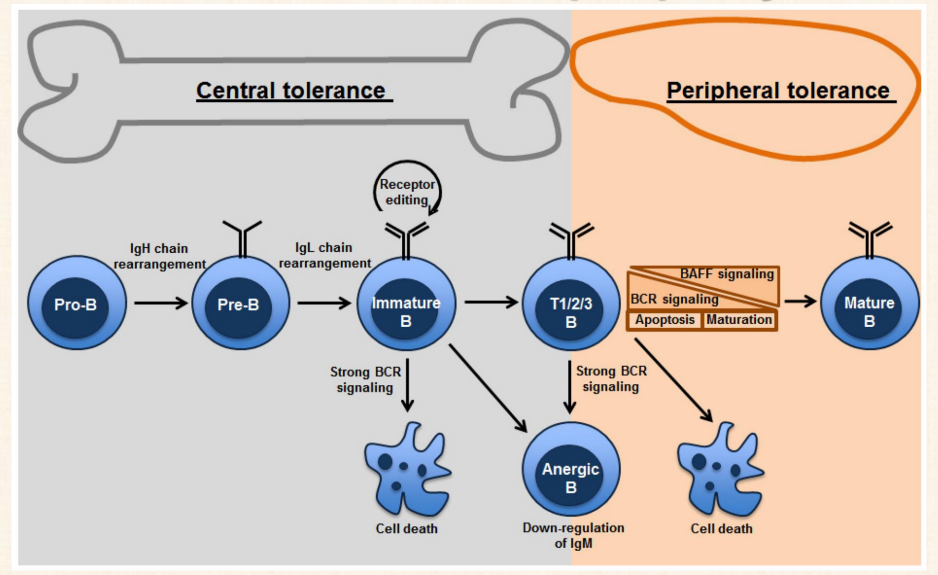


B-cell Tolerance

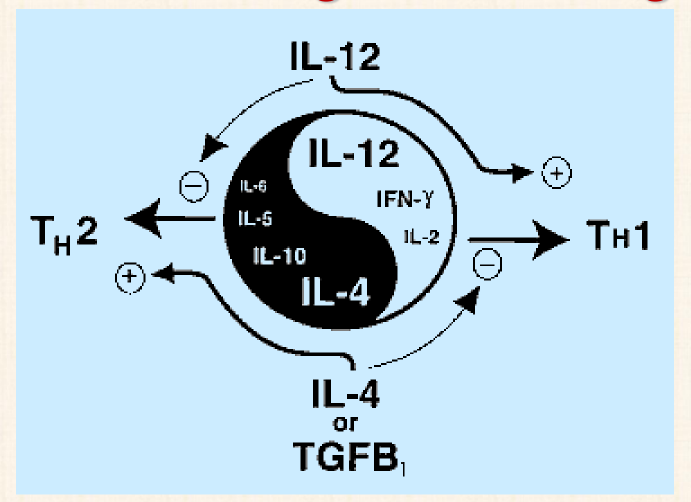
- Central tolerance

- Peripheral tolerance

Mechanisms of B-cell tolerance in bone marrow and periphery



Immunological Yin-Yang



The cytokines IL-12 and TGF beta 1 are predominant influences in "peripheral" and "mucosal" lymphatic tissues. Thus vectorial expression of these cytokines affect T cells and B cells in such a way that proliferating B cells become committed to secrete "peripheral" IgG or "mucosal" IgA, respectively.

ACTIVE TOLERANCE

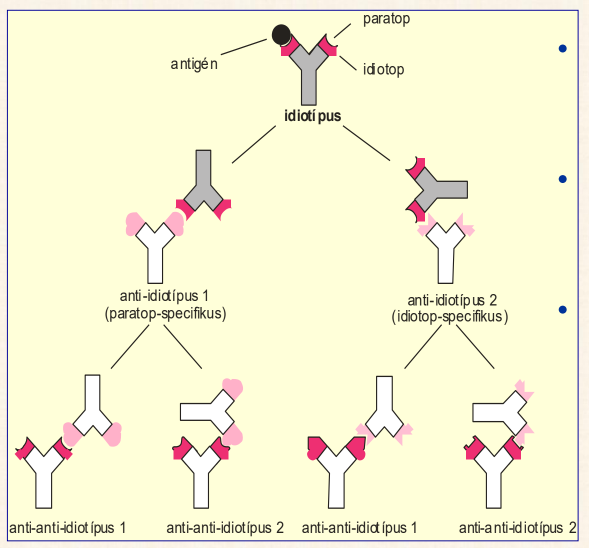
Anti-idiotype network

- Anti-idiotype antibodies against T cell and B cell receptors and immunoglobulins
- Antigen-specific inhibition and induction of memory

Natural immune system ("Immunological homunculus")

- Low affinity IgM, IgG or IgA natural autoantibodies produced by CD5+ B1B cells
- γ/δ T, iγ/δT, ILCs1,2,3, MAIT, IEL, iNKT cells

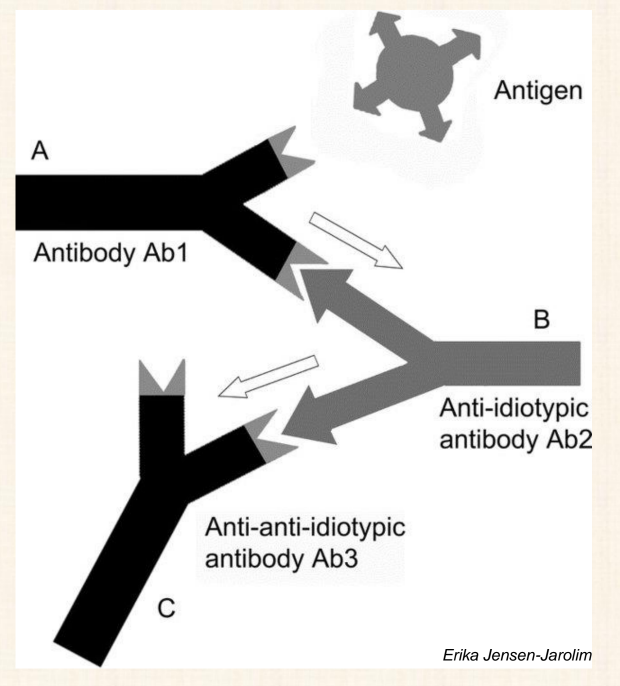
Anti-idiotype network (N. K. Jerne)



T- & B-cell suppression

Functional memory formation

Biological mimicri (insulin – anti-insulin – anti-anti-insulin ~ insulin)



The anti-idiotypic network amplifies antigenic signals. (A) An antibody Ab1 is produced in response to a specific antigen. (B) With a defined idiotype, Ab1 induces the production of an anti-idiotypic antibody Ab2. This Ab2 may resemble the original antigen as an internal image. (C) Ab2 can stimulate the synthesis of an anti(anti-idiotypic) antibody Ab3 which principally is of the same specificity as Ab1.

Naturally occurring (auto)antibodies

Autoantibodies of the IgM (mostly), or IgG and IgA classes, reactive with a variety of serum proteins, cell surface structures and intracellular structures, are 'naturally' found in all normal individuals. Present in human cord blood and in 'antigen-free' mice, their variable-region repertoire is selected by antigenic structures in the body and remains conserved throughout life. Encoded by germline genes with no, or few, mutations, natural autoantibodies are characteristically 'multireactive' and do not undergo affinity maturation in normal individuals. Natural autoantibodies may participate in a variety of physiological activities, from immune regulation, homeostasis and repertoire selection, to resistance to infections, transport and functional modulation of biologically active molecules.

Antigens recognized by natural autoantibodies

Heatshock proteins	hsp65, hsp70, hsp90, ubiquitin
Enzymes	aldolase, citockrom c, SOD, NAPDH, citrate synthase, DNA topoisomarase I.
Cell membrane components	β2-microglobulin, spectrin, acetylcholin receptor
Cytoplasmic components	actin, myosin, tubulin, myoglobin, myelin basic protein
Nuclear components	DNS, histones
Plasma proteins	albumin, IgG, transferrin
Cytokines, hormones	IL-1, TNF, IFN, insulin, thyreoglobin

Solid Organ Transplants

Bone Marrow Transplants



Autoimmune Diseases



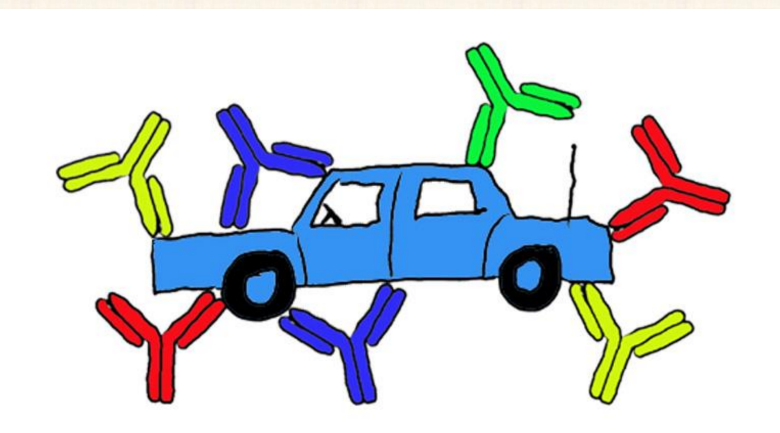
Immunologic Tolerance





Infectious Diseases/ Vaccine Development

Allergic Diseases



Autoimmunity

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Autoimmune diseases affect 5-8% of the population!

AUTOIMMUNITY

- Physiological autoimmunity: part of the normal immunological regulation
- Pathological autoimmunity: diseases caused by self reacting inflammatory immune responses with <u>permanent</u> <u>tissue/organ injury</u>

Pathomechanism of autoimmunity

- Chronic inflammation and tissue necrosis
 - Cellular components:

(T cells CD8 and Th1, NK, Mf, DC, Ne, Eo, Ba, Mc)

- Humoral components:

(Ig+complement, ADCC, cytokines, chemokines, tissue hormones and mediators)

Pathomechanism of autoimmunity

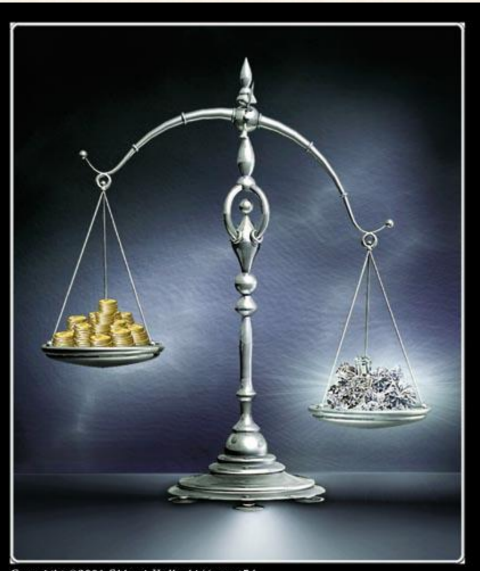
Multifactor mechanism

(general catastrophe of bio-regulation caused by external and internal factors)

- Autoimmune "steady state" (failure of dynamic balance on self tolerance and autoimmunity)
- Role of infections (molecular mimicry or inefficient natural antibody network)

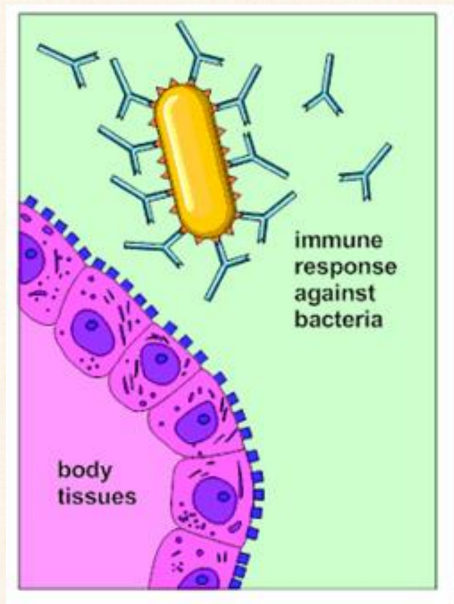
Autoimmune steady state

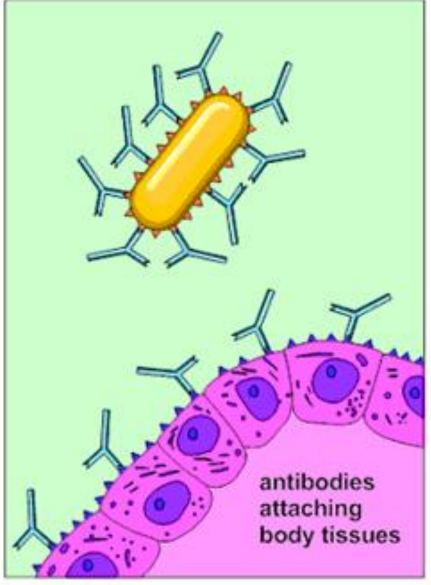
Self reacting immune response with tissues damages



Active tolerance and tissue repair

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Normal

Autoimmune Disorder

Pathogens and human antigens	Peptid residues	Overlaping sequences
Human cytomegalovirus IE2 HLA-DR molecule	79 60	PDP <u>LGRPD</u> ED VTE <u>LGRPD</u> AE
Poliovirus VP2 Acetylcholine receptor	70 176	STT <u>KESRGT</u> T TVI <u>KESRGT</u> K
Papilloma virus E2 Insulin receptor	76 66	SLH <u>LESLKD</u> S VYG <u>LESLKD</u> L
Klebsiella pneumoniae nitrogenase enzym HLA-B27 molecule	186 70	SRQTDREDE KAQTDREDL
Adenovirus 12 E1B Alfa-gliadin	384 206	LRRGMFRPSQCN LGQGSFRPSQQN
HIV p24 Human IgG	160 466	GVETTTPS GVETTTPS
Measles virus P3 Myelin basic protein	31 61	EISDNLGQE EISFKLGQE

Pathomechanisms of autoimmune diseases

- Autoimmunity by the antigen

Failed differentiation and selection of lymphocytes

- Genetic background

Autoimmunity by the antigen

Tissue injury or inflammation, leading to:

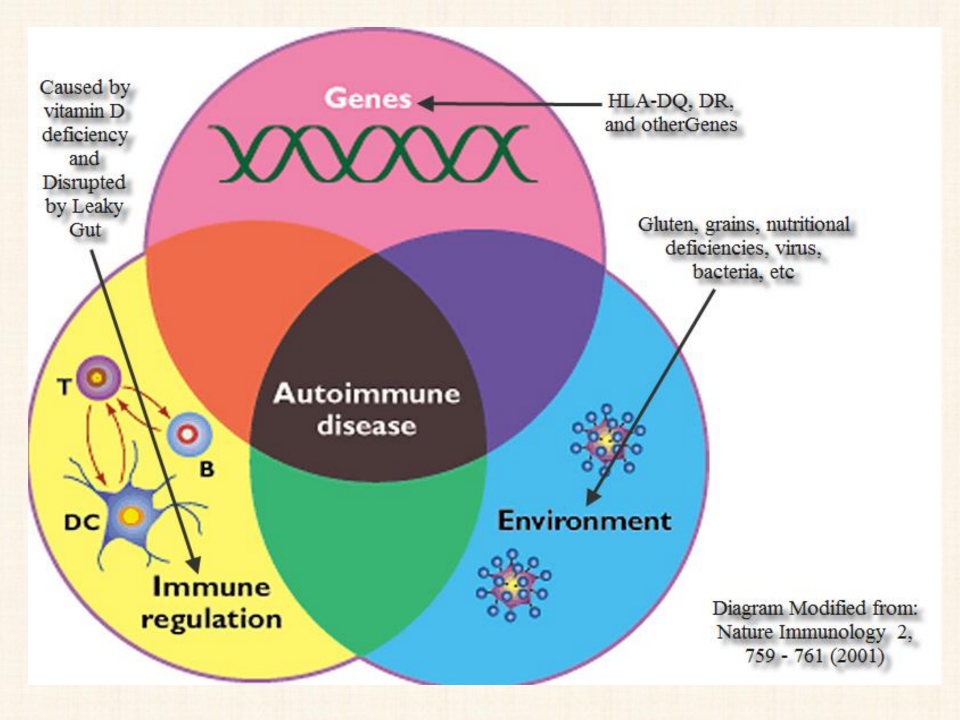
- Release of sequestered self antigens
- Structural alterations of self antigens
- Increased costimulation on tissue APCs

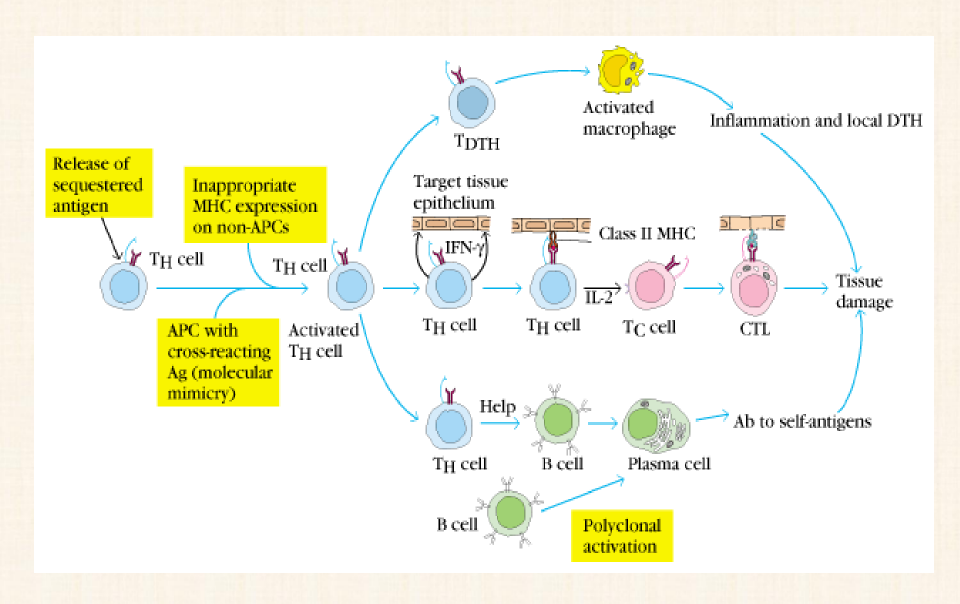
Autoimmunity by the failure of self tolerance

- Failed selection of lymphocyte repertoire
- Polyclonal activation of anergic selfreactive lymphocytes
- Stimulation by foreign antigens that cross-react with self

Thyroid gland Graves' disease Hashimoto thyreoiditis	DR3 DR5	3.7	TSH receptor ↑ Thyroid mikrosome peroxidase, thyroglobulin ↓
Pancreas IDDM	DR4/D R3 DQB 0302	20 100	B-inslet cellss ↓ GAD, HSP60, junB, insulin, pre/pro insulin
Neural system Sclerosis multiplex Myasthenia gravis	DR2 DR3	4.8 2.5	Brain white matter, MBP, PLP, MOG, MAG Peripheral neurons and striated muscle Acetycholin receptor
Heart: rheumatic fever Blood: AHA, thrombocytopenia	DR3,D R4		S. B-haemolyticus/Myosin Vvs gP Thrombocyte gP

SLE	DR3/		Kidney, serous layers
	DR2	5.8	ds/ssDNS, Sm-IC, SSA
Sjögren syndrome			Exocrine glands, salivary glands, liver, kidney, brain, thyreoid gland, heart, lung, gut
Rheumatoid arthritis (RA)	DR4 DR1	4.2	Joint connective tissues, collagen Type II, IgG RF
Spondyloarthritis	B27	<u>90</u>	Vertebra
(SPA) Reiter disease	B27	<u>33</u>	Clamydia, Yersinia
Salmonella/Shigella arthritis	B27	<u>20.7</u>	





Body Parts That Can Be Affected by Autoimmune Diseases

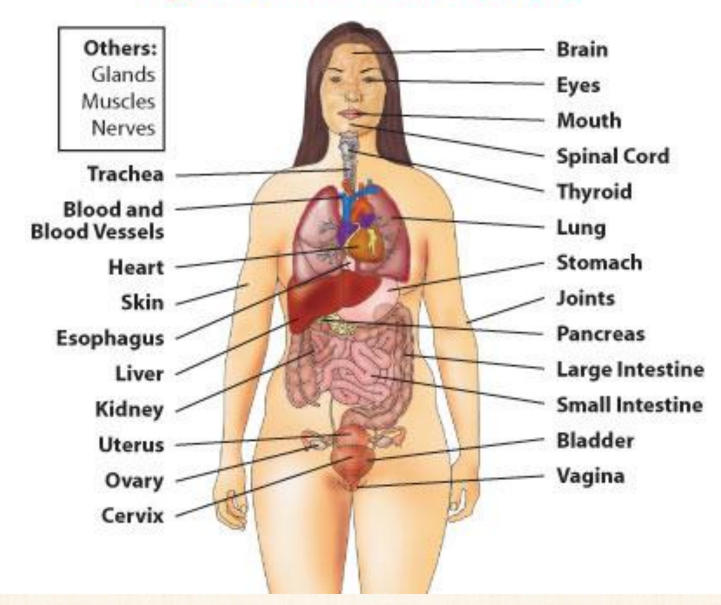
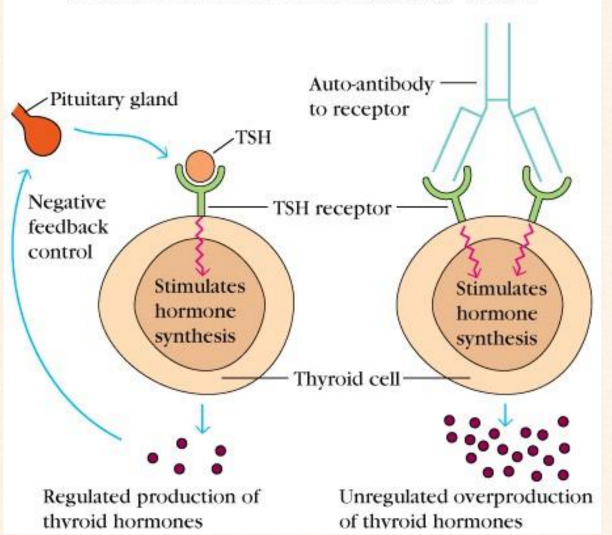


TABLE 20-1 SOME AUTOIMMUNE DISEASES IN HUMANS

Disease	Self-antigen	Immune response
	Organ-specific autoimmune diseases	
Addison's disease	Adrenal cells	Auto-antibodies
Autoimmune hemolytic anemia	RBC membrane proteins	Auto-antibodies
Goodpasture's syndrome	Renal and lung basement membranes	Auto-antibodies
Graves' disease	Thyroid-stimulating hormone receptor	Auto-antibody (stimulating)
Hashimoto's thyroiditis	Thyroid proteins and cells	T _{DTH} cells, auto-antibodies
Idiopathic thrombocyopenia purpura	Platelet membrane proteins	Auto-antibodies
Insulin-dependent diabetes mellitus	Pancreatic beta cells	T_{DTH} cells, auto-antibodies
Myasthenia gravis	Acetylcholine receptors	Auto-antibody (blocking)
Myocardial infarction	Heart	Auto-antibodies
Pernicious anemia	Gastric parietal cells; intrinsic factor	Auto-antibody
Poststreptococcal	Kidney	Antigen-antibody complexes
glomerulonephritis		
Spontaneous infertility	Sperm	Auto-antibodies
	Systemic autoimmune disease	
Ankylosing spondylitis	Vertebrae	Immune complexes
Multiple sclerosis	Brain or white matter	T_{DTH} and T_{C} cells, auto-antibodies
Rheumatoid arthritis	Connective tissue, IgG	Auto-antibodies, immune complexes
Scleroderma	Nuclei, heart, lungs, gastrointestinal tract, kidney	Auto-antibodies
Sjogren's syndrome	Salivary gland, liver, kidney, thryoid	Auto-antibodies
Systemic lupus erythematosus (SLE)	DNA, nuclear protein, RBC and platelet membranes	Auto-antobidies, immune complexes

Grave's disease

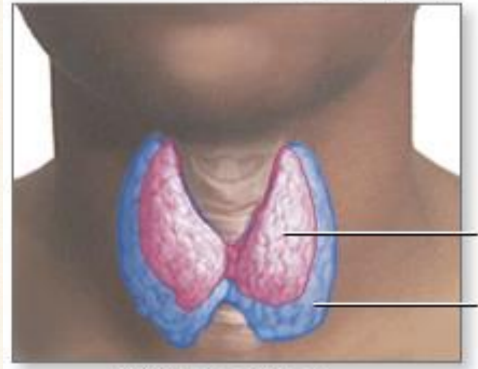
STIMULATING AUTO-ANTIBODIES (Graves' disease)



In Graves' Disease patient produces autoantibodies that bind to the receptors for thyroidstimulating hormone (TSH). TSH is produced by the pituitary gland and the receptors for TSH are present on thyroid cells. Binding of these autoantibodies mimics the normal action of TSH which is to stimulate the production of two thyroid hormones, thyroxine and triiodothyronine. However, the autoantibodies are not under a negative feedback control system and therefore lead to overproduction of the thyroid hormones. For this reason these autoantibodies have been termed long-acting thyroidstimulating (LATS) antibodies. Overproduction of thyroid hormones leads to many metabolic problems.



Exophthalmos (bulging eyes)



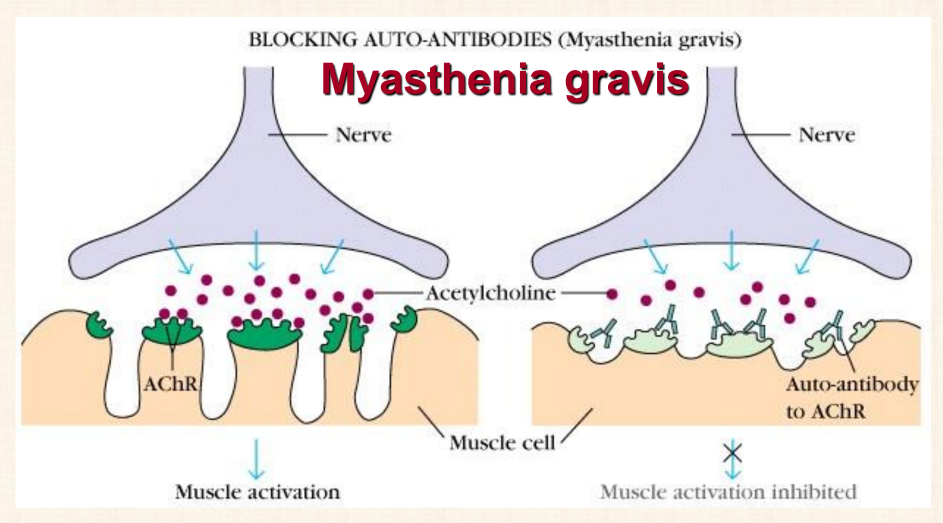
Diffuse goiter

Graves' disease is a common cause of hyperthyroidism, an over-production of thyroid hormone, which causes enlargement of the thyroid and other symptoms such as exophthalmos, heat intolerance and anxiety

Normal thyroid

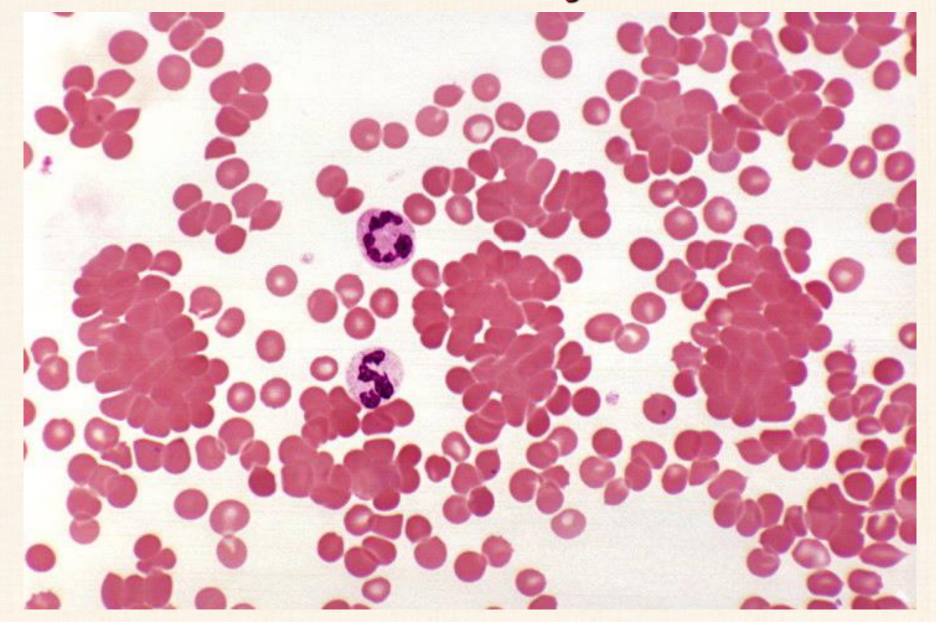
Enlarged thyroid



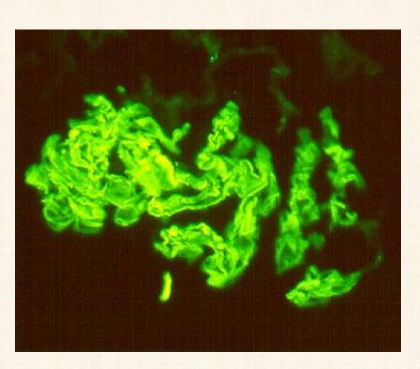


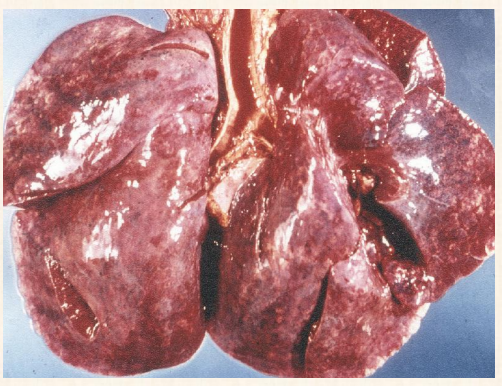
A patient with this disease produces autoantibodies to the acetylcholine receptors on the motor end-plates of muscles. Binding of acetylcholine in therefore blocked and muscle activation is inhibited. The autoantibodies also induce complement-mediated degradation of the acetylcholine receptors, resulting in progressive weakening of the skeletal muscles.

Autoimmune hemolytic anemia



Goodpasture syndrom



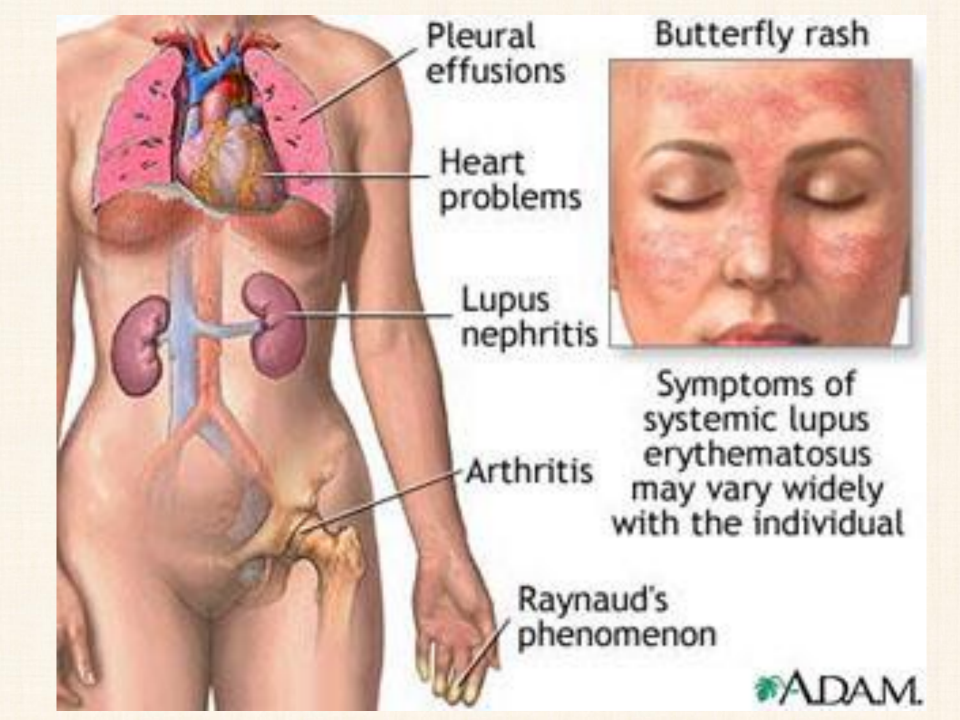


Autoantibodies are produced against alpha-3 subunit of type IV collagen in the basement membranes of glomeruli and lung causing bleeding necrosis.



Characteristic "butterfly" rash over the cheeks of a young girl with **SLE**.

Systemic Lupus Erythematosus (**SLE**) is characterized by fever, weakness, arthritis, skin rashes, pleurisy, and kidney dysfunction. Affected individuals may produce <u>autoantibodies</u> to a range of tissue antigens such as <u>DNA</u>, <u>histones</u>, <u>RBCs</u>, <u>platelets</u>, <u>leukocytes</u>, <u>and clotting factors</u>. SLE typically appears in women between 20 and 40 years of age with a <u>female:male ratio of 10:1</u>. An example of complications arising from SLE is when <u>immune complexes</u> are deposited along the walls of small blood vessels. This deposition <u>activates</u> <u>complement</u> system, resulting in glomerulonephritis and damage to the blood-vessel wall (<u>vasculitis</u>) causing widespread tissue damage.









Csökkent maximalis oralis apertura ill. teleangiectasia SSc-ben

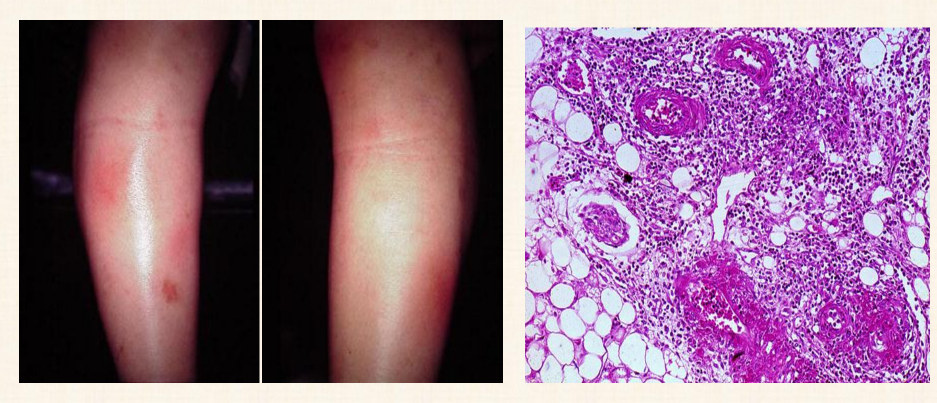
Diffuse cutaneous SSc (dcSSc): skin manifestation both on the extremities and on the trunk, severe internal organ involvement, poor prognosis

Limited cutaneous SSc (IcSSc): skin involvement only on the face and distal part of extremities, no internal organ involvement, good prognosis

The major autoantibody in SSc targets DNA topoisomerase I (Topo I or ScI-70)

Anti-Topo I autoantibodies are detected mainly, but not exclusively in dcSSc

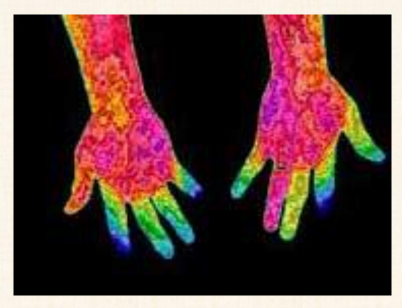
Periarteritis nodosa



The medium sized arteries in the fat tissue appear magenta red because their wall is impregnated with fibrin (fibrinoid necrosis). There is also marked inflammation in the wall of these blood vessels extending into the perivascular connective tissue (arteritis and periarteritis).

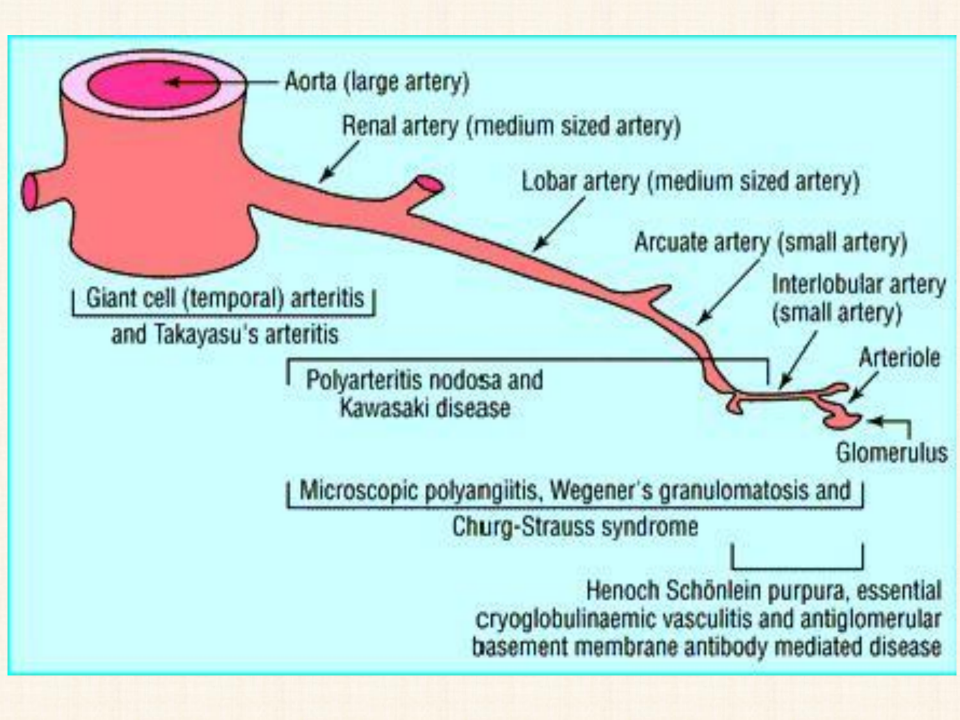
Periarteritis nodosa











Raynaud's Syndrome



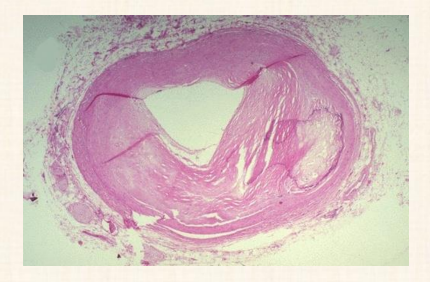




anti-Phopholipid syndrome

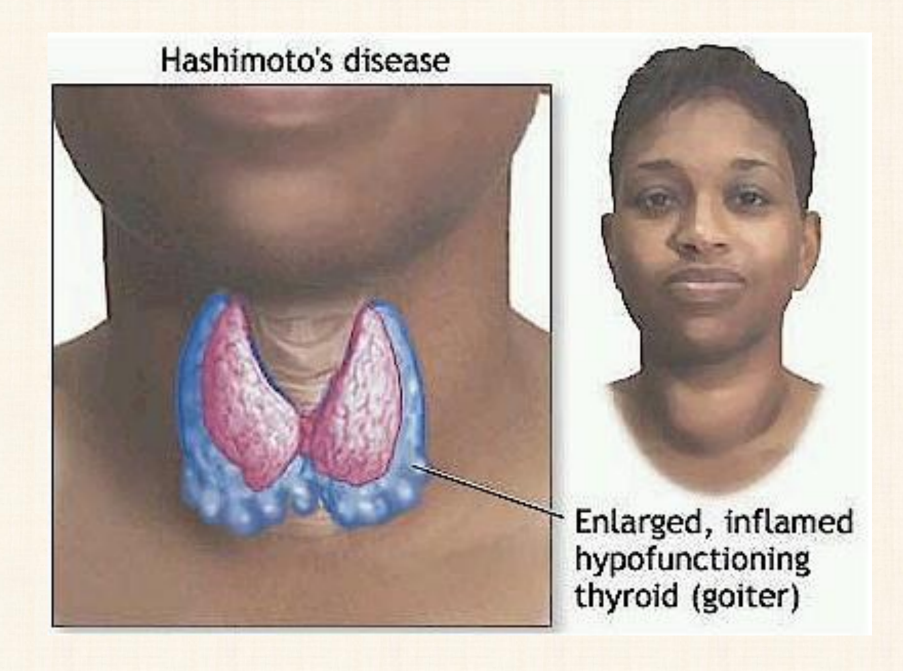




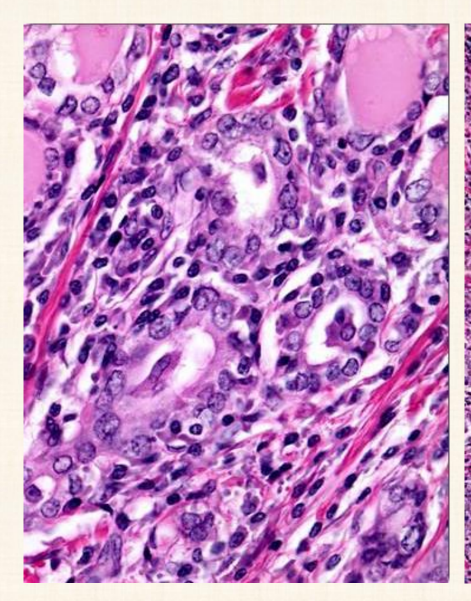


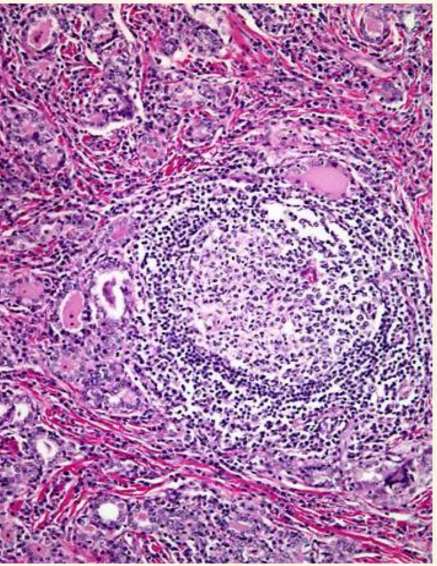
Livedo reticularis

Antiphospholipid Syndrome (APS, APLS, Hughes Syndrome, or Sticky Blood): abnormal antibodies linked to abnormal blood clots within veins and arteries.

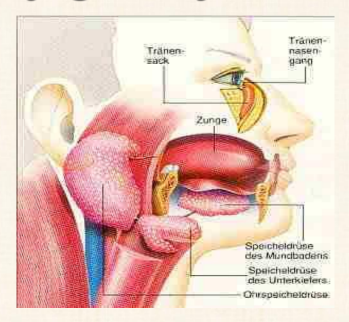


Hashimoto's disease

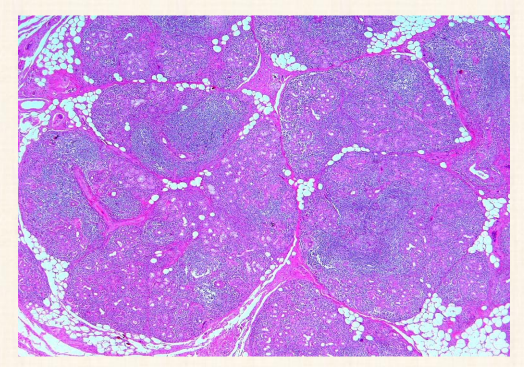


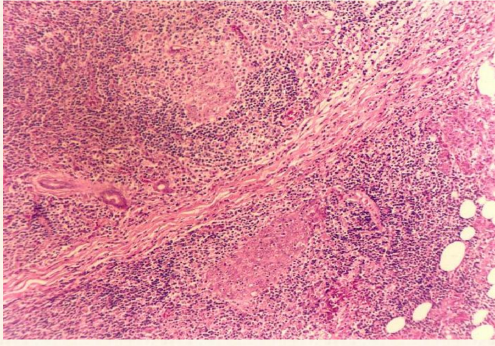


Sjögren syndrome

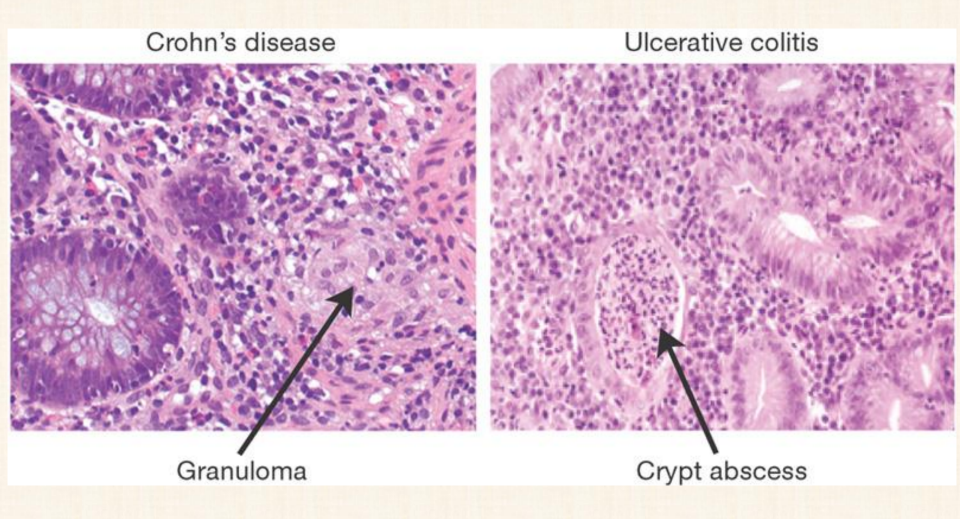




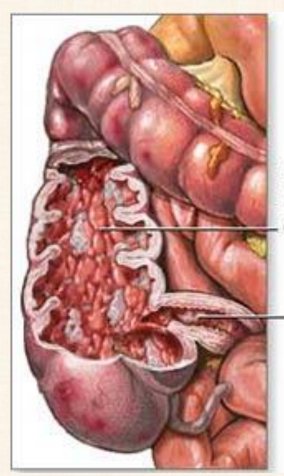




Inflammatory Bowel Diseases

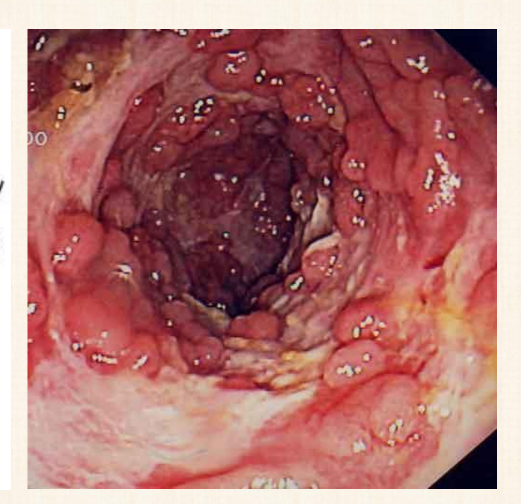


Crohn's disease



Inflammatory bowel disease (IBD)

Ileum portion of small intestine



Ulcerative colitis

