#### Imunodeficiency states

- Primary
  - Caused by defined genetic defects
  - Usually rare, but severe (exception: IgA deficiency)
- Secondary
  - Consequence of some other disease, treatment, environmental factors...
  - Usually frequent, but usually clinically mild (exceptions: HIV disease, secondary aganulocytosis).

### Severe combined immunodeficiency (SCID)

- Early clinical manifestation (weeks-months)
- Severe and complicated infections affecting respiratory and gastrointestinal tract and the skin
- Failure to thrive
- Frequent diarrhea
- Usually lymphocytopenia
- T-cell deficiency, B cell present in some patients
- Decreased immunoglobulin levels



# SCID infections caused by atypical patogens

- Pneumocystis pneumonia
- Cytomegalovirus pneumonitis
- Disseminated BCG-itis
- Infections caused by atypical mycobacteria
- Candidiasis of oropharynx, skin



#### Immunoglobulin Deficiencies

Clinical manifestations begins at 6-12 months (or late).

Susceptibility to infection by encapsulated bacteria (Pneumococcus, Haemophilus).

Respiratory tract predominantly affected; patients suffer from recurrent otitis media, bronchitis, sinusitis, pneumonia.

Some patients also suffer from meningitis or chronic diarrhea.

#### X-linked agammaglobulinemia

- Only boys affected
- Clinical manifestation usually begins at 6-12 months
- Severe and complicated respiratory tract infections.
- Very low levels of all immunoglobulin isotypes.
- B-cell not detected

### Common variable immunodeficiency (CVID)

- Both sexes affected
- Clinical manifestation initiates at any age
- Frequent and severe respiratory tract infections
- Proneness to autoimmune diseases
- Variable decrease of immunoglobulin isotypes, usually markedly decreased IgA and IgG leves
- B-lymphocytes usually present

#### Selective IgA deficiency

- Frequency: 1:400
- Usually only mild manifestation
- Predominantly respiratory tract infections
- Patients are prone to autoimmune diseases
- Beware of anti-IgA antibodies that can cause a severe anaphylactic reaction after artificial IgA administration (by blood, immunoglobulin derivates)!

#### **T-cell Deficiences**

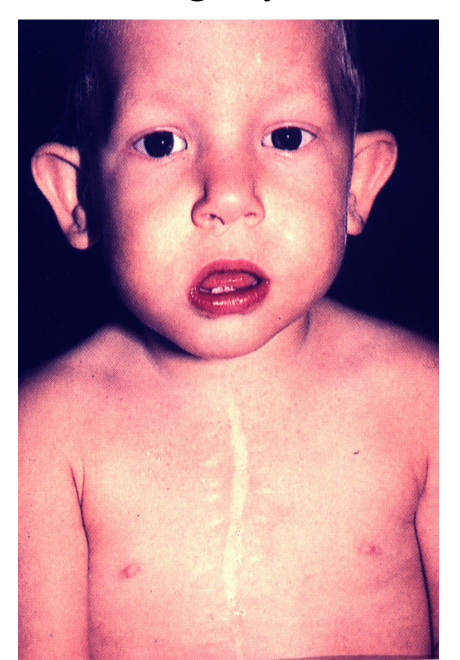
- -Early onset of clinical manifestation.
- -Extreme susceptibility to viral, fungal, mycobacterial, and protozoal infections.
- Respiratory system most frequently affected, but also other systems can be involved.

#### DiGeorge syndrome

- Defect in embryonic development of the 3rd and 4th pharyngeal pouches.
- Cardiovascular defects
- Hypoparathyroidism → hypocalcemia → seizures
- Thymic hypoplasia → T cell deficiency
- Typical facies: hypertelorism, micrognatia, low-set, posterior rotated ears.



#### DiGeorge syndrome

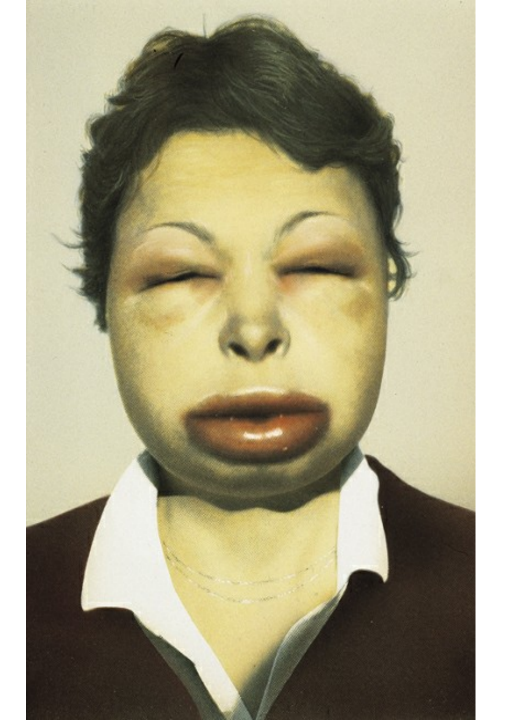


#### Complement deficiencies

- Deficiency of C1-C4: autoimmune systemic disorders, susceptibility to bacterial infections
- Deficiency C5-C9: susceptibility to bacterial infections, mainly to meningococcal meningitis
- Deficiency of C1 INH: hereditary angioedema

#### Hereditary angioedema

- Deficiency of C1 inhibitor (C1 INH)
- Uncontrolled activation of the complement system after trauma, infection, surgical operation....
- Vasoactive peptides (bradykinin, C3a,C5a) cause increased vascular permeability
- Oedema of the skin, respiratory tract (dyspnoe), gastrointestinal tract (cramps, vomiting)



#### HEREDITARY ANGIOEDEMA (HAE)

#### Phagocytic dysfunction

- Early onset of clinical manifestation
- Susceptibility to bacterial and fungal infections
- Abscess formation, mainly of the skin, periproctal area, liver, but any area may be affected.

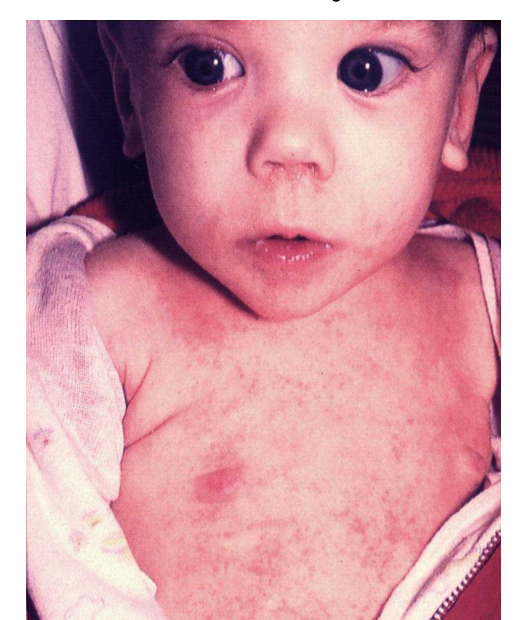
#### Chronic granulomatous disease

- Recurrent abscesses mainly of the liver, lungs, periproctal area, suppurative lymphadenitis, osteomyelitis
- Infections are caused mainly by catalasepositive organisms: St. aureus, Candida sp., Serratia marcescens
- Usually early onset of symptoms
- Production of reactive metabolites of oxygen is disturbed (defect of NADPH oxidase)

#### Wiskott-Aldrich syndrome

- X-linked disease
- Thrombocytopemia → bleeding tendency
- Severe eczema
- Immunodeficiency
- Severe allergic and autoimmune manifestations
- B-cell lymphomas

#### Wiskott-Aldrich syndrome



#### Ataxia telangiectasia

- Autosomal recessive
- Progressive cerebellar ataxia
- Telangiectasis especially on ear lobes and conjunctival sclera
- Immunodeficiency
- Frequent tumors
- Cause: mutation in ATM gene

#### Ataxia telangiectasia



### Treatment of primary immunodefciencies

- SCID and other severe immunodeficiencies: bone marrow transplantation, gene therapy in some cases.
- Antibody deficiencies: immunoglobulin replacement
- Antibiotic prohylaxis

### Clinical use of non-specific immunoglobulin derivates

- Types of derivates:
  - "Normal immunoglobulin" for intramuscular use. Used very rarely at present bcause only low dose can be given..
  - Intravenous immunoglobulins, subcutaneous immunoglobulins can be used in high doses
- Indications:
  - Replacement treatment in patients with antibody deficiencies
  - Prophylaxis of infections against which there is no specific immunoglobulin derivate (hepatitis A)
  - High doses of i.v. immunoglobulins are used in autoimmune diseases, systemic vasculitic diseases.

### Causes of secondary immunodeficiency

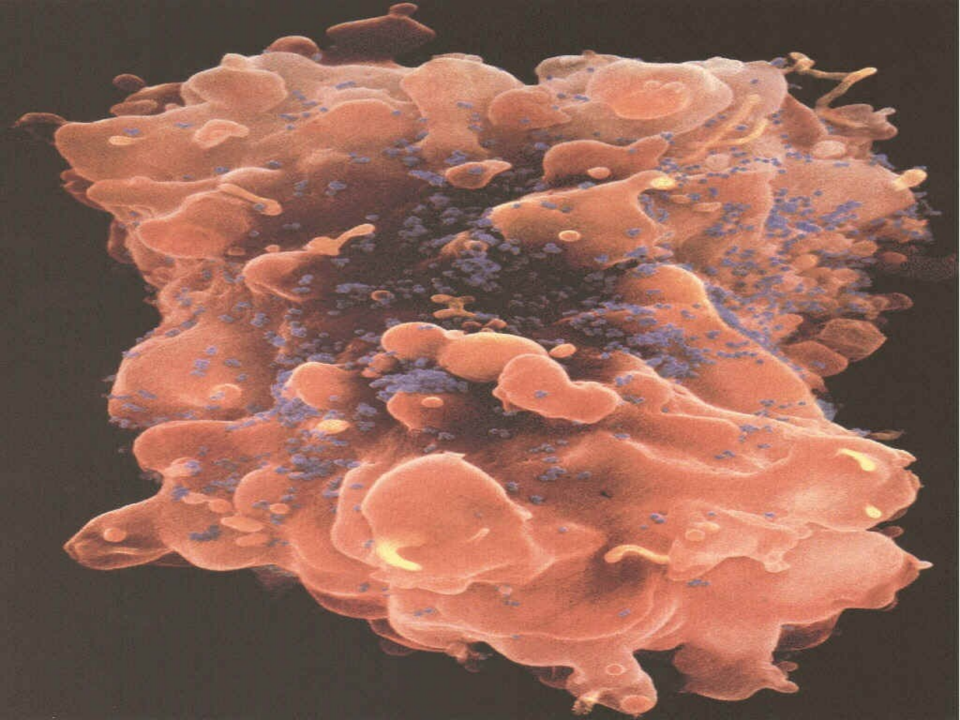
- Metabolic uremia, diabetes, malnutrition
- latrogenic cytostatics, immunosuppressants
- Malignat tumors
- Viral infections HIV, CMV, measles, infectious mononucleosis
- Splenectomy
- Stress
- Injuries, operations, general anestesia

### Imunodeficiency after splenectomy

- Disturbed phagocytosis, decreased production of antibodies.
- The most severe complication is hyperacute pneumococal sepsis.
- Prevention: vaccination against Pneumococcus, Haemophilus infl. B, Meningococcus. PNC prophylaxis.

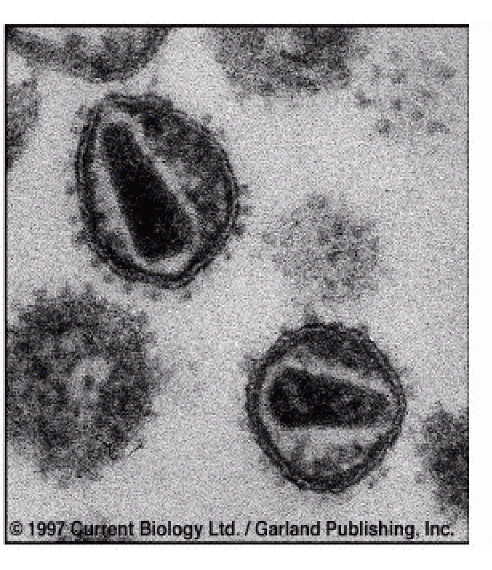
#### Secondary hypogammaglobulinemia

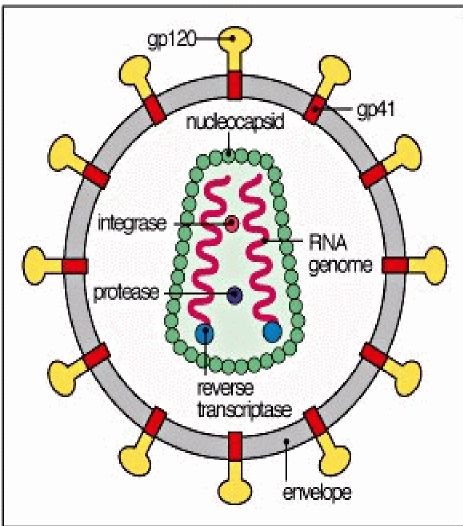
- Decreased production of immunoglobulins
  - Chronic lymphatic leukemia
  - Lymphoma
  - Myeloma
- Loss of immunoglobulins
  - Nephrotic syndrome
  - Exudative enteropathy



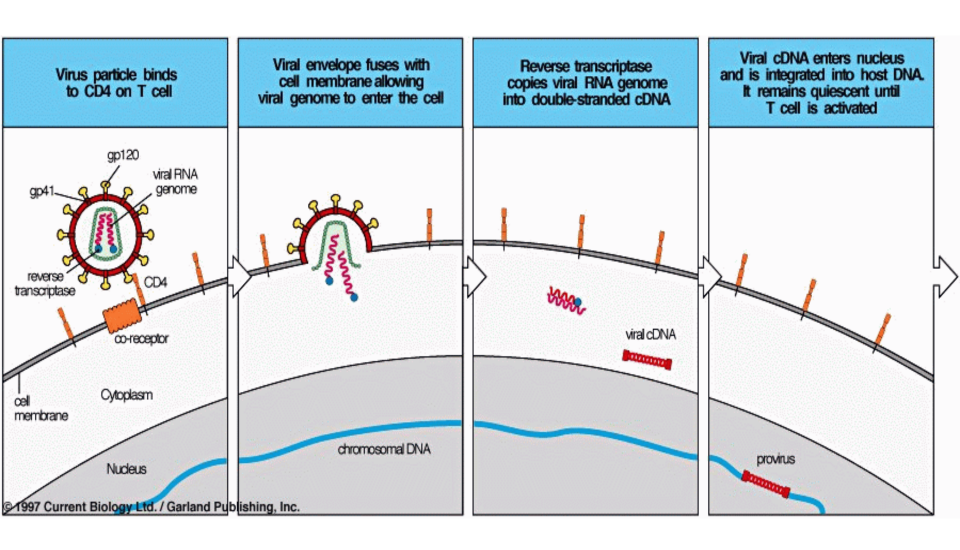
#### Ways of transmission

- 1. Sexual
- 2. <u>Parenteral</u> intravenous drug addicts previously blood products
- 3. <u>Vertical</u> mother to child transplacental, during delivery, by brestfeeding

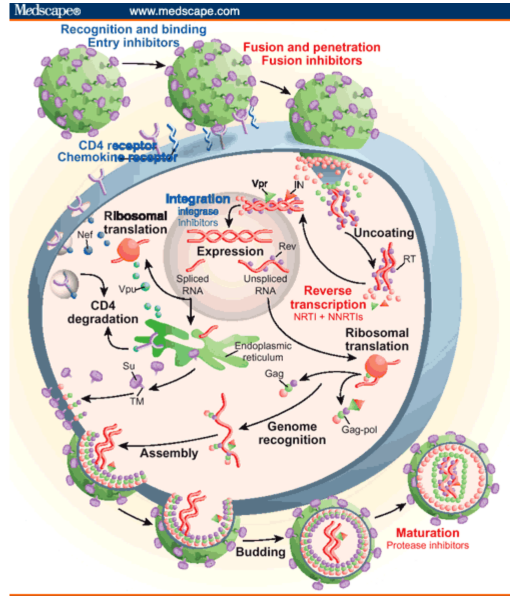


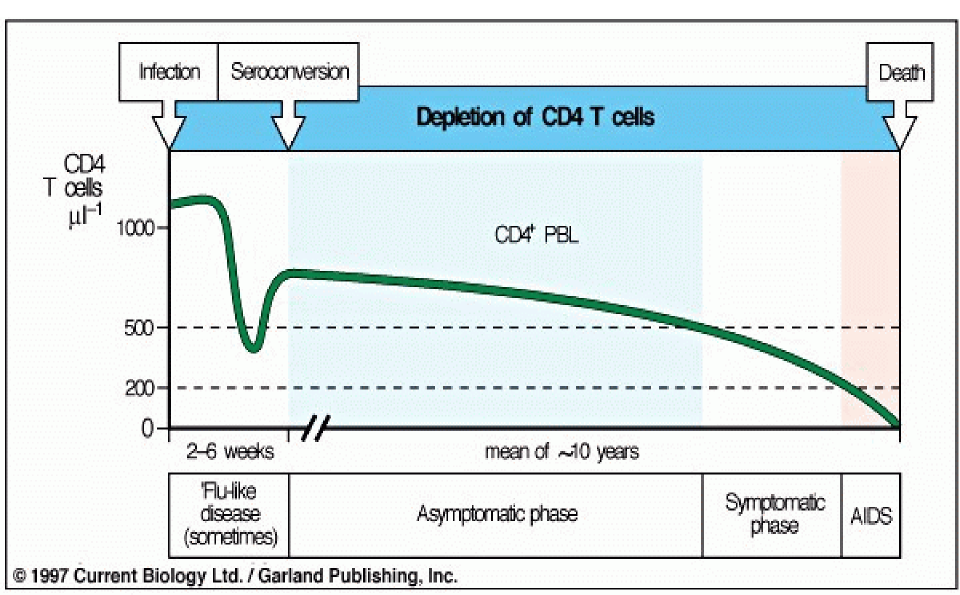


#### **Infection of CD4+ cell by HIV**



#### **HIV Cycle**





## Classification of HIV disease (CDC) 3 clinical categories

- A Asymptomatic disese
- B "small" opportunistic infections
- C "big" opportunistic infections and other states that define AIDS

#### Clinical category A

- Accute (primary) HIV infection
- Asymptomatic HIV infection
- Persistent generalised lymphadenopathy (PGL)

#### **HIV PRIMOINFECTION**

- Acute retroviral syndrome, ("mononucleosis-like syndrome")
- Present in 50-70% patients
- 2-6 weeks after infection

# Clinical presentaioon of HIV primoinfection

- Fever, lympadenopathy, pharyngitis
- Rash
- Myalgia, arthralgia, diarrhoea, cephalea
- nausea, vomiting
- thrush
- Neurologic symptoms
- Aphtous stomatitis

# Perzistent generalized lympadenopathy

- More than 3 months
- 1/3 HIV-infected persons
- Lymph nodes 0,5-2,0 cm, painless

### Clinical category B

- Fever >38,5 C more than 1 month
- Diarrhoea more than 1 month
- Oropharyngea candidiasis
- Vulvovaginal candidiasis (chronic or difficult to treat)
- Recurrent herpes zoster

## Clinical category C (AIDS)

- Pneumocystis pneumonia
- Brain abscess caused by toxoplasma
- Esofageal, tracheal, bronchial or lung candidiasis
- Chronic anal herpes, herpetic bronchitis, pneumonia
- CMV retinitis, generalized CMV infection
- Progressive multifocal leukoecephalopathy
- Mycobacterial infections

# Opportunistic Infections in AIDS Patients

- Pneumonia due to Pneumocystis jiroveci (carinii)
- Toxoplasma brain abscess
- Cytomegalovirus infection (retinitis, colitis)
- Mycobacterial infections
- Herpes virus and Varicella-Zoster infections

# Clinical category C (AIDS) - tumors

Kaposhi sarcoma

Brain lymphoma





Wasting syndrome



#### Treatment of HIV-disease

- Antiretroviral
  - Nucleoside inhibitors of reverse transcriptase: azidothymidin (syn. zidovudin), didanosin, zalcitabin, stavudin, lamivudin
  - Nonnucleoside inhibitors of reverse transcriptase: Nevirapin, delavirdin, efavirenz
  - HIV protease inhibitors: Saquinavir, ritonavir, indinavir
- Prophylaxis of *Pneumocystis carinii* pneumonia (co-trimoxazol), antiviral and antimycotic antibiotics

## Strategy of treatment

HAART - Highly

**Active** 

**Anti** 

Retroviral

**Therapy** 

Mega-HAART

## Diagnosis of HIV infection

- Detection of anti-viral antibodies
  - ELISA
  - -Western blott

Detection of antigen p 24