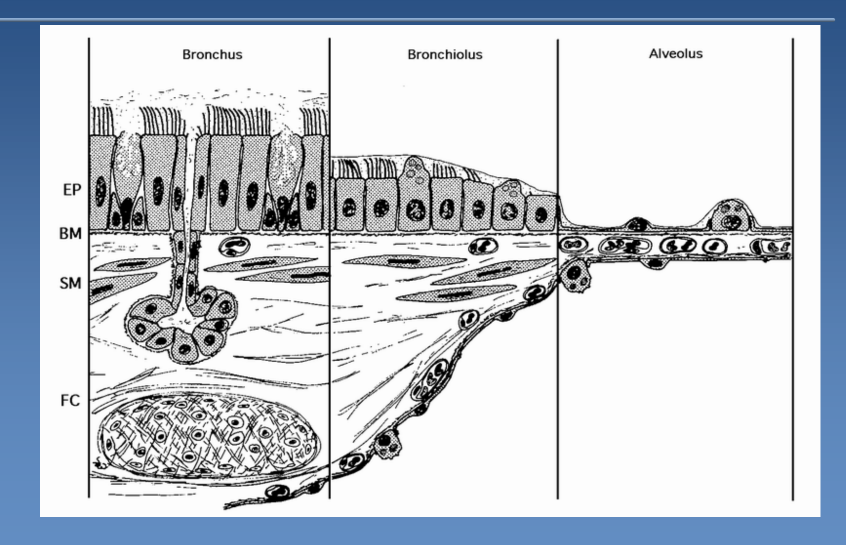
## Systemic pathology



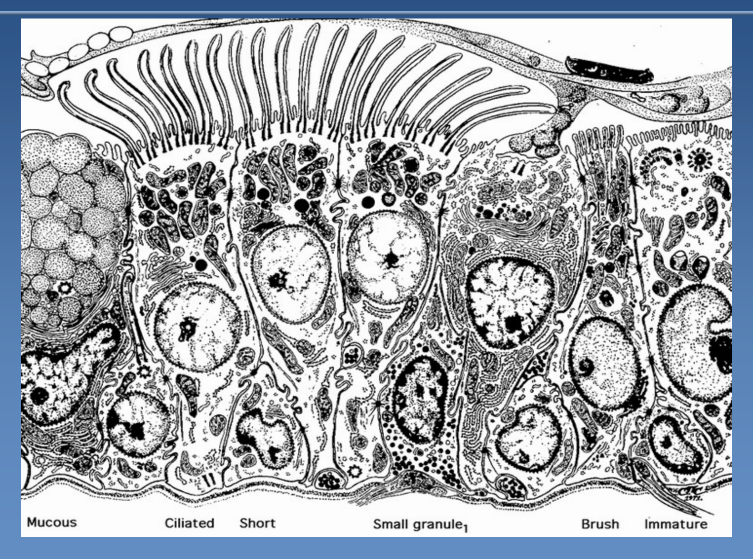
The respiratory tract

### Histology of respiratory tract

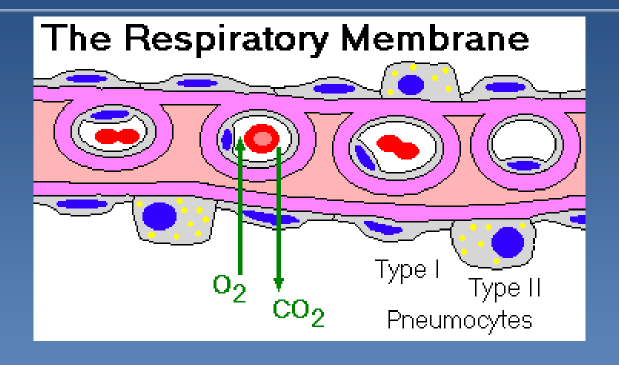


# Cellular components of bronchial mucosa





## The respiratory membrane



## Chronic polypous rhinitis



chronic proliferative inflammation

- etiology:
  - chronic irritation
  - allergy
  - repeated acute inflammations

## Chronic polypous rhinitis



#### Gross:

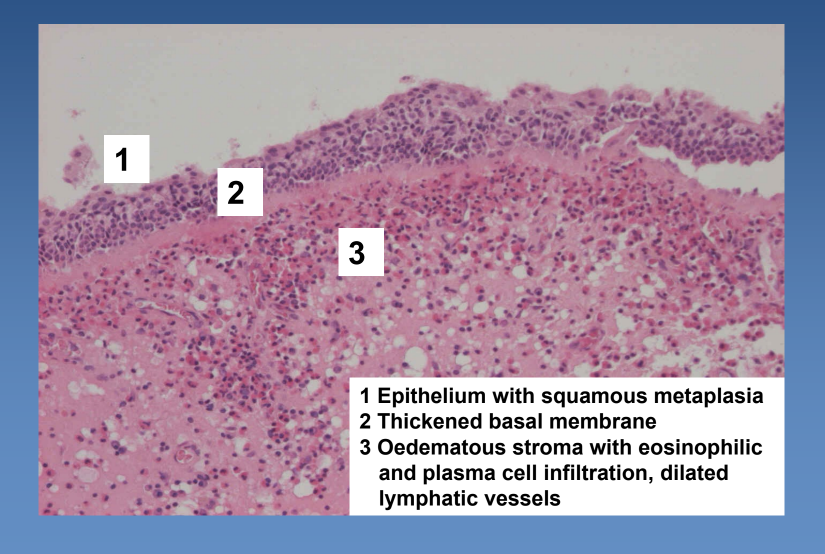
- mucosal polyps, often multiple
- variable size (mm 2 cm)

#### Micro:

- oedematous mucosal connective tissue
- Iymphoplasmocytic reactive infiltration, admixture of eosinophils, +/- neutrophils
- mucinous hyperplasia
- covered by hyperplastic respiratory epithelium, squamous metaplasia possible

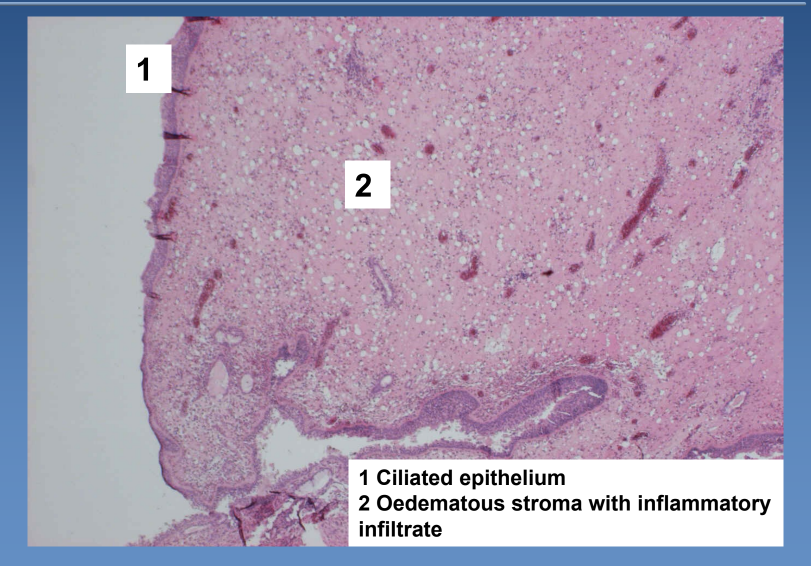
## Polypous chronic rhinitis





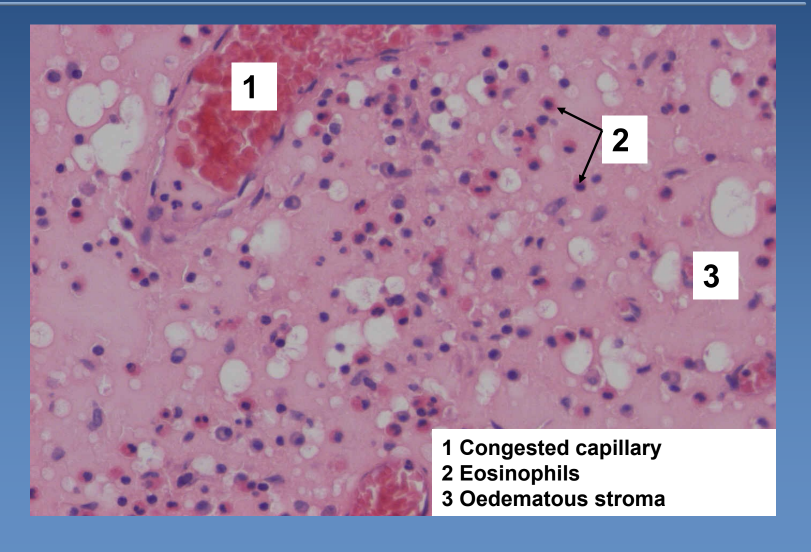
### Polypous chronic rhinitis





## Polypous chronic rhinitis





#### Laryngeal carcinoma



- Sequence: in squamous epithelium: hyperplasia atypical hyperplasia intraepithelial neoplasia (dysplasia carcinoma in situ) invasive carcinoma.
- In respiratory epithelium: squamous metaplasia intraepithelial neoplasia (dysplasia carcinoma in situ) invasive carcinoma.
- Commonly multiple dysplastic foci and/or sequential carcinomas in upper respiratory/GIT – same oncogenic factors, field theory

#### Laryngeal carcinoma



- Risk factors: smoking, alcohol, HPV, (asbestos, irradiation)
- **Papilloma:** HPV, solitary (adults) x multiple (papillomatosis in children). Benign, possible recurrence
- Carcinoma: mainly squamous cell ca, rare adenocarcinoma
- On vocal cords, supravocal, infravocal
- Clinical features: hoarseness, later pain, dysphagia, bleeding

#### Pseudomembranous tracheitis



- Diphteria, influenza, scarlet fever, mumps, etc.
- Iatrogenic intubation; uremia.
- Risk of ulceration chondromalatia cartilage breakdown perforation mediastinitis

### **Pulmonary infarction**



- aetiology:
  - thrombembolism of a. pulmonalis branches in the setting of compromised cardiovascular status (passive venous congestion)
- typically hemorrhagic
- often in lower lung lobes adjacent to pleura
- often multiple
- healing:
  - granulation tissue, later formation of fibrous scar

### Pulmonary infarction



- aetiology:
  - thrombembolism of a. pulmonalis branches in the setting of compromised cardiovascular status (passive venous congestion)
  - uncommonly local thrombosis/arterial closure (in carcinoma)
- typically hemorrhagic
- often in lower lung lobes adjacent to pleura
- often multiple
- healing:
  - granulation tissue, later formation of fibrous scar

### Hemorrhagic pulmonary infarction

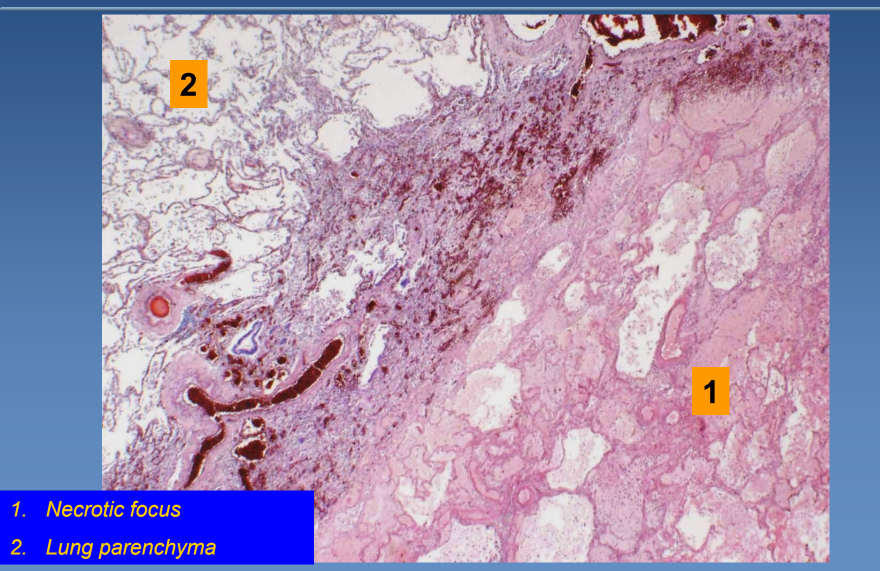
#### Gross:

- wedge-shaped sharply demarcated focus
- dark red-blue (new), yellowish-grey (older)
- > variable size
- firmer consistency

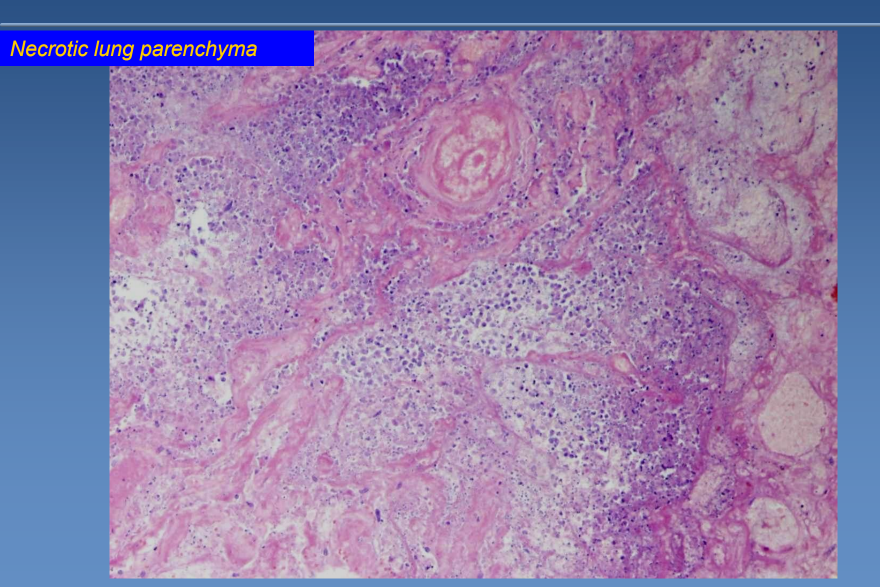
#### Micro:

- coagulative necrosis of lung parenchyma
- large extravasations of erythrocytes
- formation of abscess by secondary infection
- reactive acute fibrinous pleuritis
- healing scarring + emphysema (diff.dg. x tumor)

## Hemorrhagic pulmonary infarction



## Hemorrhagic pulmonary infarction



#### Alveolar oedema



- \* fluid accumulation in alveoli
- clinically:
  - expectoration of bubbly watery pinkish sputum
- pathogenesis:
  - → ↑ vascular permeability (injury to the alveolar-capillary wall)
  - → ↑ vascular hydrostatic pressure
  - → ↓ intravascular osmotic pressure
  - lymphatic drainage obstruction

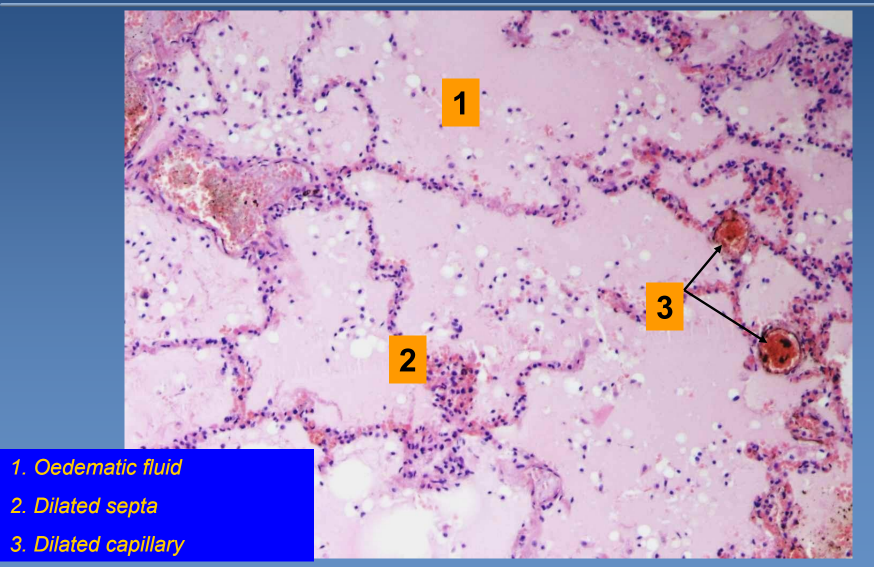
#### Alveolar oedema



- Complications: ↑ risk of infection
- Gross:
  - lungs enlarged, heavy, congested
  - bubbly fluid flowing out of the tissue +/- present in bronchi
- Micro:
  - alveoli filled with pink, homogenous fluid + air bubbles
  - dilatation and hyperemia of alveolar wall capillaries

#### Alveolar oedema





## Amniotic fluid aspiration



- minor aspiration usual during birth
  - clinically insignificant
- massive aspiration associated with fetus asphyxia
  - umbilical cord or placental disorders
- clinic:
  - changes in fetal heart rate immediate medical intervention necessary!

## Amniotic fluid aspiration

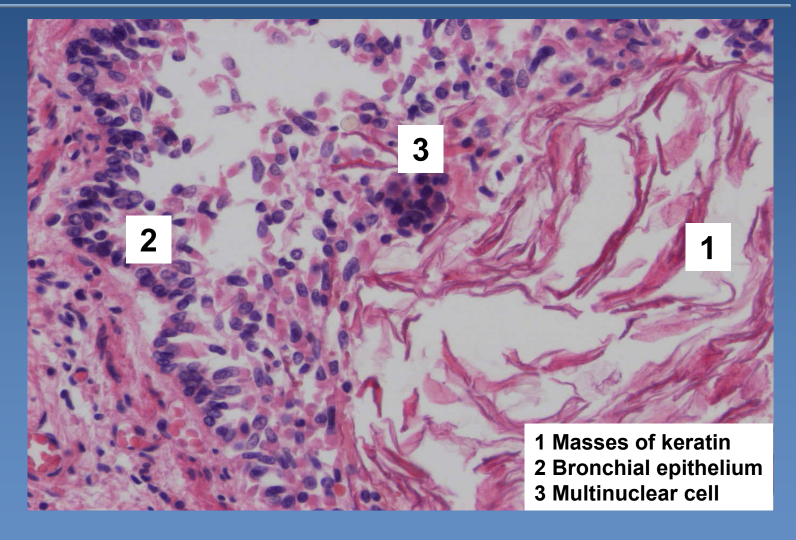


#### Micro:

- > keratin masses in bronchi and alveoli
- **⇒** amniotic cells
- lanugo (thin primary hairs)
- meconium bodies (from fetus intestinal content)
- infected amniotic fluid >> fetal death, adnate pneumonia

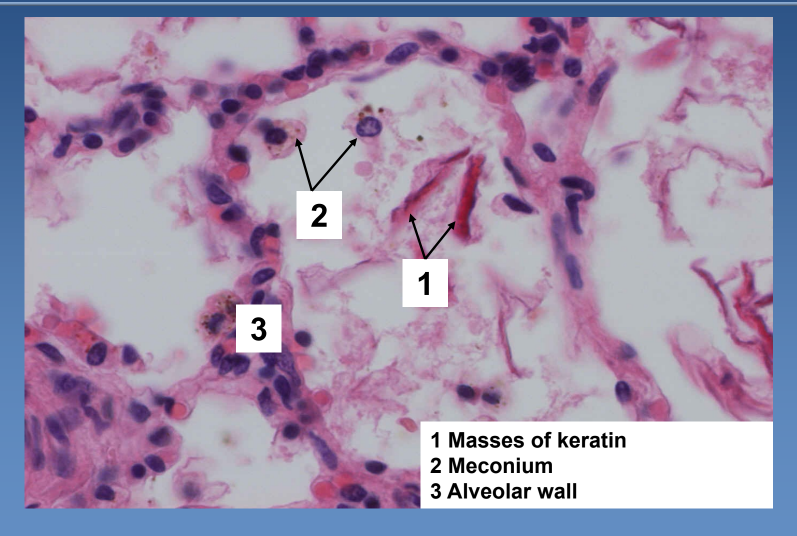
#### Amniotic fluid aspiration, keratin in bronchiole





#### Amniotic fluid aspiration, keratin in alveoli







- associated with chronic <u>left-sided</u> cardiac insufficiency
  - **>** etiology:
    - ischemic heart disease, systemic hypertension, valvular disorders, cardiomyopathy
- clinically ("asthma cardiale"):
  - cough
    - rusty sputum
  - shortness of breath (dyspnoea)
    - ortopnoea
    - paroxysmal nocturnal dyspnoea
      - relieved by sleeping with elevated head ("additional pillows needed")



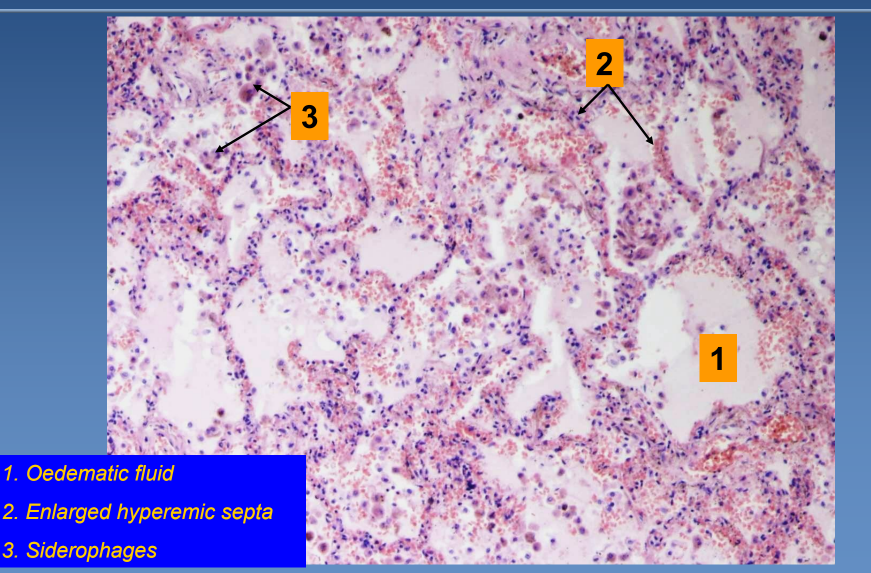
#### Gross:

- slightly enlarged lungs
- **solid** consistency
- rusty-brown color
  - rusty/cyanotic lung induration

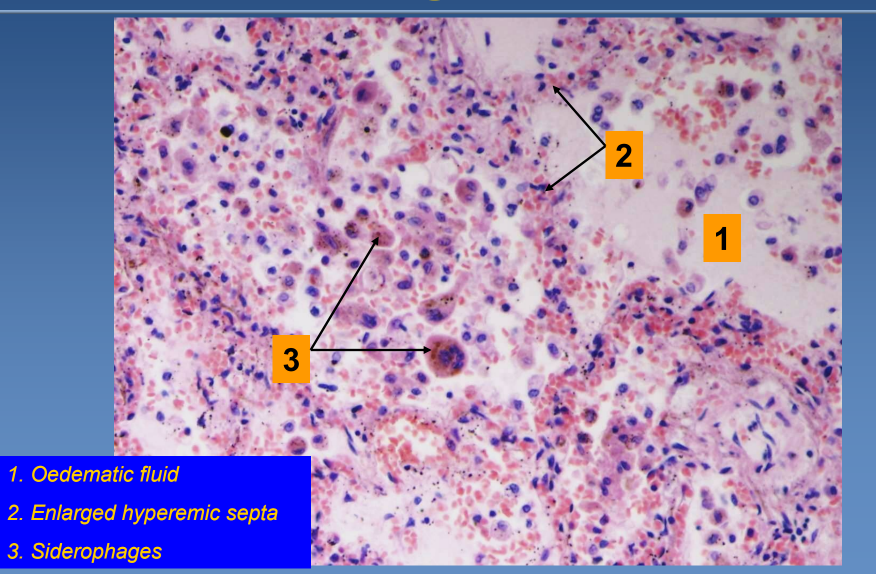
#### Micro:

- congestion of alveolar capillaries
- alveolar hemorrhage with siderophages:
  - histiocytes with cytoplasmic granules of hemosiderin
- fibrotization of alveolar walls

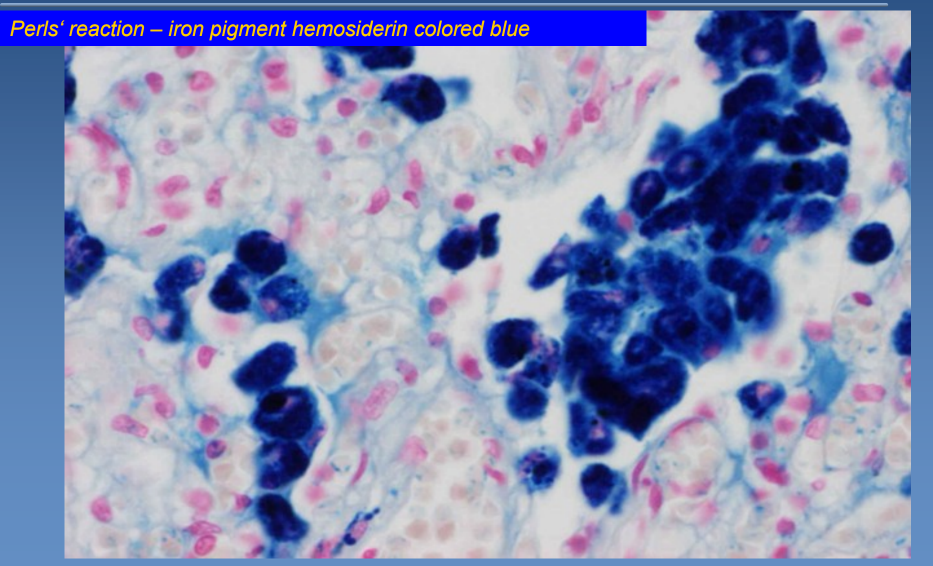












## Chronic pulmonary diseases

- Cobstructive airway d.-↑in resistance to airflow due to partial/complete obstruction at any level (trachea bronchi bronchiolog)
  - bronchioles)
    - **chronic** bronchitis
    - **⇒** bronchiectasis
    - **⇒** asthma
    - **bronchiolitis**
    - emphysema
- Restrictive reduced expansion +/- decreased total lung capacity.
  - chronic interstitial and infiltrative disorders
  - chest wall disorders

## Chronic obstructive pulmonary diasease



- Clinical syndrome productive cough, dyspnoea, end-stage – respiratory failure
- Pathology: chronic bronchitis +/- emphysema
- Major trigger cigarette smoking, air pollution
- Complications: recurrent bacterial/viral infections, cor pulmonale, pneumothorax, lung cancer, may progress to respirátory failure

#### Chronic bronchitis



- part of spectrum of ch. obstructive pulmonary disease, duration at least 3 months in 2 years
- Simple ch. b.
  - productive cough, no airflow obstruction
- Chronic asthmatic bronchitis
  - intermittent bronchospasm, hyperreactive bronchi
- **Solution** Obstructive ch. b.
  - chronic obstruction, usually + emphysema



- chronic inflammatory disease of bronchial tree, recurrent attacks of bronchospasm with exspiratory dyspnoea, cough, mucus hypersecretion
- increased irritability of the bronchial tree with paroxysmal narrowing of the airways.
- status asthmaticus:
  - increased frequency of attacks permanent bronchospasm
  - may be lethal
- variants:
  - atopic (extrinsic):
    - environmental factors, type I hypersensitivity reaction, IgE, mast cells degranulation, increased vascular permeability and mucus secretion + eosinophils activation
    - bronchioloconstriction, distal collapse or overinflation
  - non atopic (intrinsic):
    - triggered by infection (viral), subsequent hyperreactive state of vagal receptors, reaction after nonspecific irritation
  - drug-induced: i.e. aspirin, NSAID, cytokine dysbalance, commonly + urticaria, rhinitis
  - occupational: variable etiology (type I+III hypersensitivity) and stimulating agents



#### Gross:

- acute changes: bronchospasm + emphysema/collapse, mucus plugs in peripheral bronchi and bronchioles, bronchial inflammatory infiltrate
- chronic airway remodeling: hypertrophy/hyperplasia of smooth muscle and mucous glands

#### Micro:

- intraluminal:
  - mucus (Curschmann spirals), eosinophils, Charcot-Leyden crystals, cellular detritus

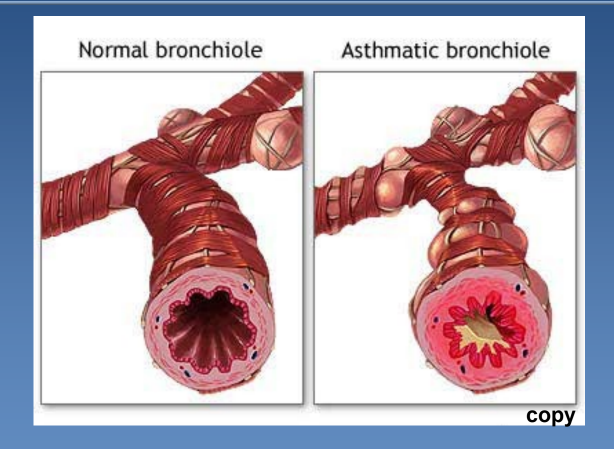
#### **bronchial** wall:

- oedema of the mucous membrane
- thickening (collagenisation) of the sub-basement membrane tissue
- mucous glands hypertrophy, eosinophil-rich inflammatory infiltrate, 

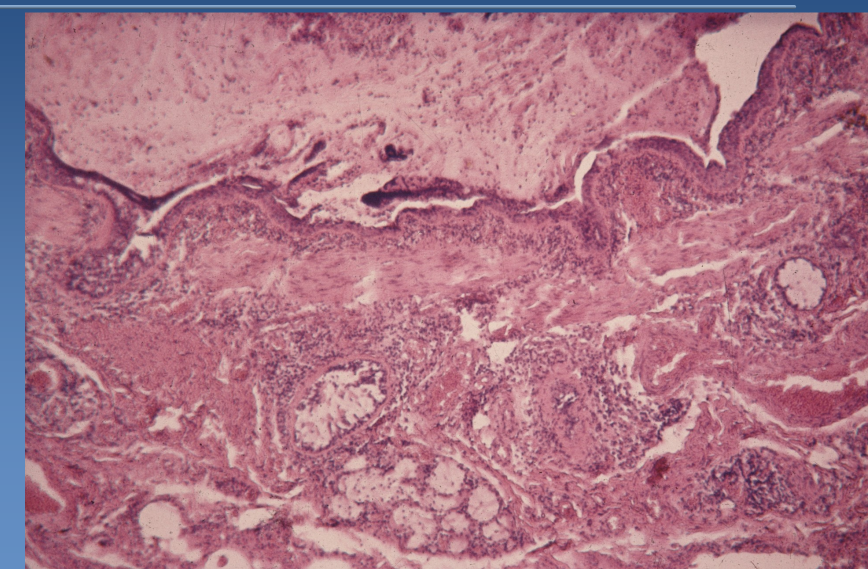
   vascularity,

   MALT











- permanent abnormal dilatation of bronchi
- arising from the weakening of the walls or changes in air pressure
- morphology:
  - cylindrical
  - ⇒ saccular
  - fusiform



### aetiology:

- congenital/hereditary conditions:
  - cystic fibrosis
  - Kartagener syndrome (structural abnormalities of the cilia, leading to persistent infections)

#### acquired:

chronic inflammations

Postinfectious (incl. necrotizing pneumonia)

Bronchial obstruction (tumor, foreign bodies, mucus)

Other (SLE, rheumatoid arthritis, etc.)

- radiotherapy
- changes of the pressure
  - chronic pulmonary collapse

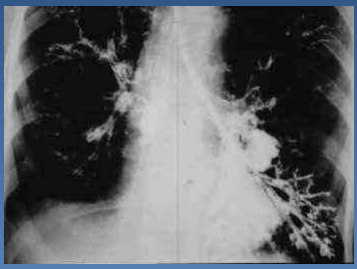


### complications:

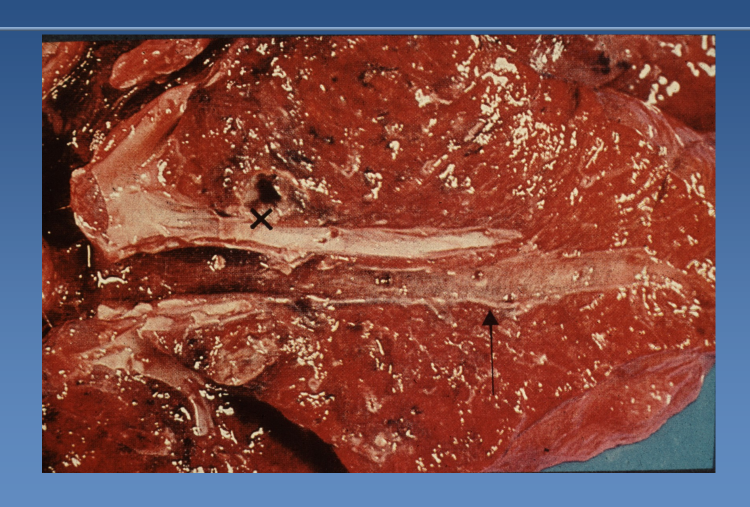
- inflammations:
  - chronic purulent bronchitis
  - bronchopneumonia including abscess formation
  - secondary infection incl. fungal (aspergilloma)
  - metastatic infection (brain abscess)
- emphysema
- fibrosis, pulmonary hypertension and cor pulmonale
- secondary AA amyloidosis











## Pulmonary emphysema



- regressive change
- abnormal permanent enlargement of the airspaces + alveolar wall destruction in the pulmonary tissue
- aetiology (combination of several factors):
  - smoking
  - $\Rightarrow$  deficiency of  $\alpha$ 1-antitrypsin
  - other
- types:
  - alveolar:
    - acute
    - chronic
  - interstitial airway rupture (trauma)

## Alveolar emphysema



### acute:

- alveolar septa are not destroyed
- rather pulmonary hyperinflation or distention

### chronic:

- permanent enlargement of airspaces distal to terminal bronchioles
- destruction of alveolar walls
- part of COPD (chronic obstructive pulmonary disease)
  - · combination of chronic bronchitis and chronic emphysema





pathogenesis and complications: proteaseantiprotease + oxidant-antioxidant imbalance in the setting of inflammatory response, bronchiolitis, later possible maladaptive immune response

```
thinning of alveolar walls and capillaries →
reduced blood supply →
complete destruction of alveolar walls →
difficult expiration + decreasing of lung capacity →
hypoxemia → endothelial cell dysfunction
medial hypertrophy, intimal fibrosis + vasoconstriction →
secondary pulmonary hypertension → →
cor pulmonale
```

## Alveolar emphysema



### types:

- ⇒ centrilobular (centriacinar):
  - upper lobes apex, more in males,
  - most commonly seen in smokers without congenital antitrypsin deficiency (but + chronic bronchitis), possible professional disease dust

### panacinar:

often lower lung zones; significant microscopic changes; antitrypsin deficiency, old age

### ⇒ distal acinar (paraseptal):

 adjacent to pleura, upper lobes foci of fibrosis, formation of cystlike structures – bullae (pneumothorax risk)

#### **⇒** irregular:

associated with scarring, usually postinflammatory

## Alveolar emphysema



### Gross:

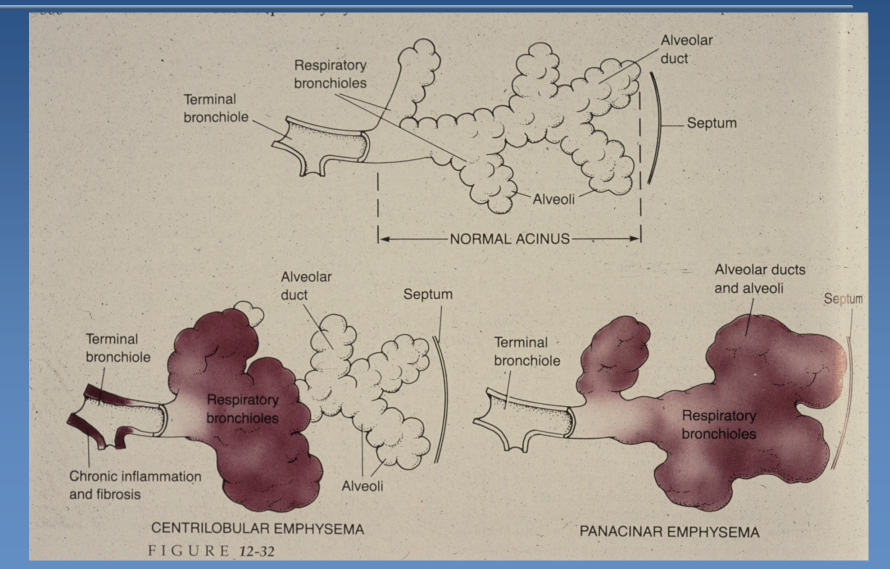
enlarged, voluminous lungs, light, pale, dry, emphysematous bullae

### Micro:

- thinning and destruction of alveolar walls
- deformation of bronchiolar walls
- chronic inflammatory changes

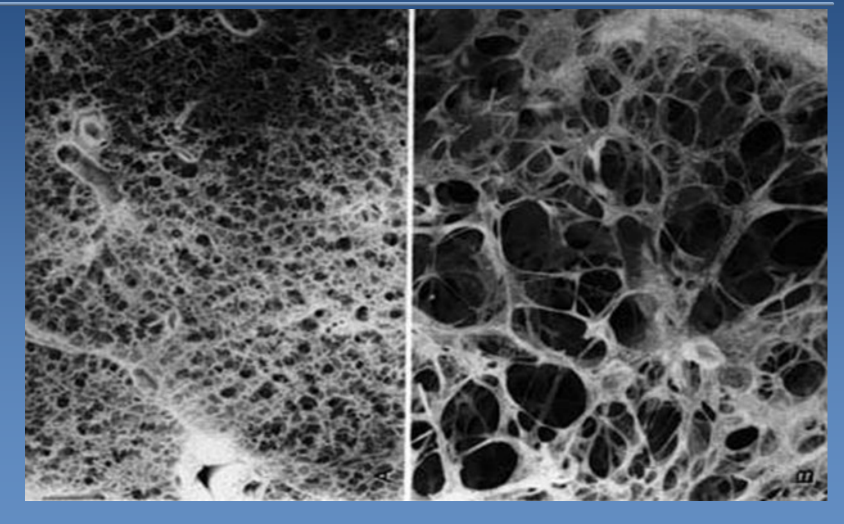
## Emphysema





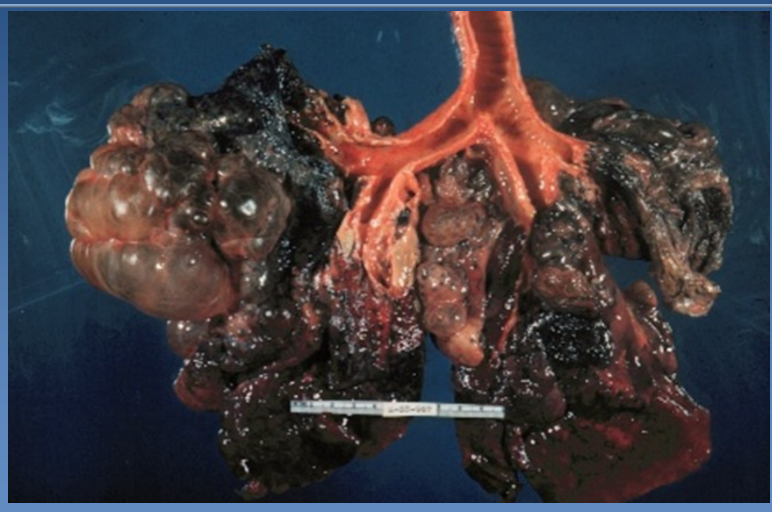
## Normal lung and pulmonary emphysema





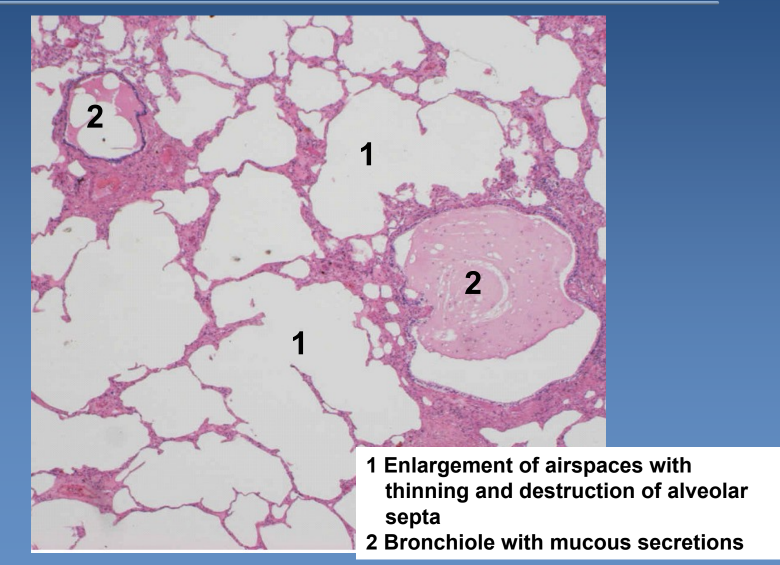
## Bullous emphysema





## Panacinar emphysema





## Pulmonary inflammations - classification



### Etiology

- Infections
- Non-infectious, commonly from the group of chronic interstitial lung disease (hypersensitivity pneumonitis, nonspecific interstitial pneumonia, etc.)

## Pulmonary inflammations - classification



- superficial:
  - > lobar pneumonia
  - bronchopneumonia
- interstitial
  - purulent (abscess, gangrene)
  - non-purulent
    - infectious (acute) atypical pneumonia
    - non-infectious (chronic)



- Community acquired acute pneumonia
  - ⇒ Str. pneumoniae
  - Haemophilus influenzae
  - Staph. aureus
  - Legionella pneumophila
  - Klebsiella pneumoniae
  - Pseudomonas
  - others (Moraxella, ...)



- Community acquired atypical pneumonia
  - Mycoplasma pneumoniae
  - Chlamydia ssp.
  - Coxiella burnetii (Q-fever)
  - ⇒ viruses influenza, parainfluenza, adenovirus, RS virus, etc.





- Hospital acquired pneumonias
  - ⇒ G- rods, Enterobacteriaceae (Klebsiella, E.coli, Pseudomonas)
  - Staph. aureus (methicillin resistant)

- Aspiration pneumonia
  - anaerobic oral flora + aerobic bacteria (incl. Str., Staph., Haemophilus etc.)



- Chronic pneumonia
  - > Nocardia
  - **⇒** Actinomyces
  - Granulomatous: mycobacteria (TBC, atypical), Histoplasma, other fungi
- Necrotizing pneumonia and lung abscess
  - Anaerobic bacteria (+/- mixed aerobic infection)
  - ⇒ Staph. aureus, Klebsiella, Str. pyogenes
  - some anthropozoonozes (plague, anthrax)



- P. in the immunocompromised host
  - ⇒ CMV
  - Pneumocystis jirovecii
  - Mycobacterium avium-intracellulare
  - ➡ Invasive aspergillosis
  - > Invasive candidiasis
  - "usual" infections



- \* superficial diffuse fibrinous inflammation
- affecting major part / entire lobe of a lung
  - similar histological features in the same time
  - ⇒ older/immunocompromised patients → lethal without antibiotic therapy
- untreated 4 stages:
  - congestion (+ oedema)
  - red hepatization (inflammatory infiltrate + congestion)
  - grey hepatization (fibrin)
  - resolution (resorption)



- healing:
  - ad integrum
  - **complications:** 
    - empyema
    - abscess
    - carnification
    - sepsis
    - metastatic purulent inflammation
      - e.g.leptomeningitis, pericarditis, endocarditis...

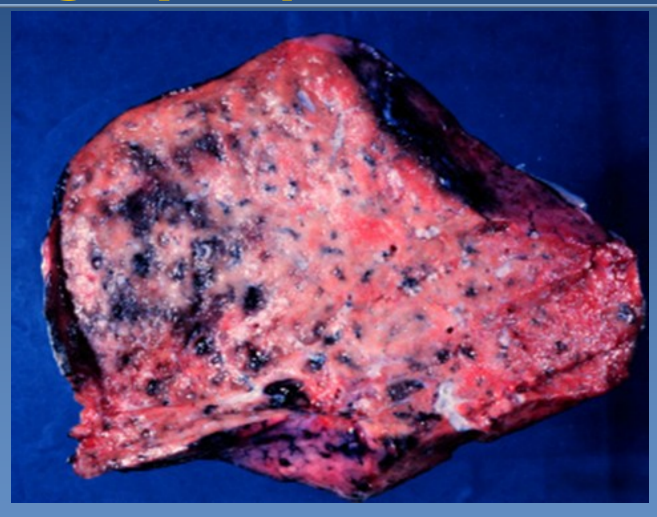
# Lobar pneumonia, red hepatization



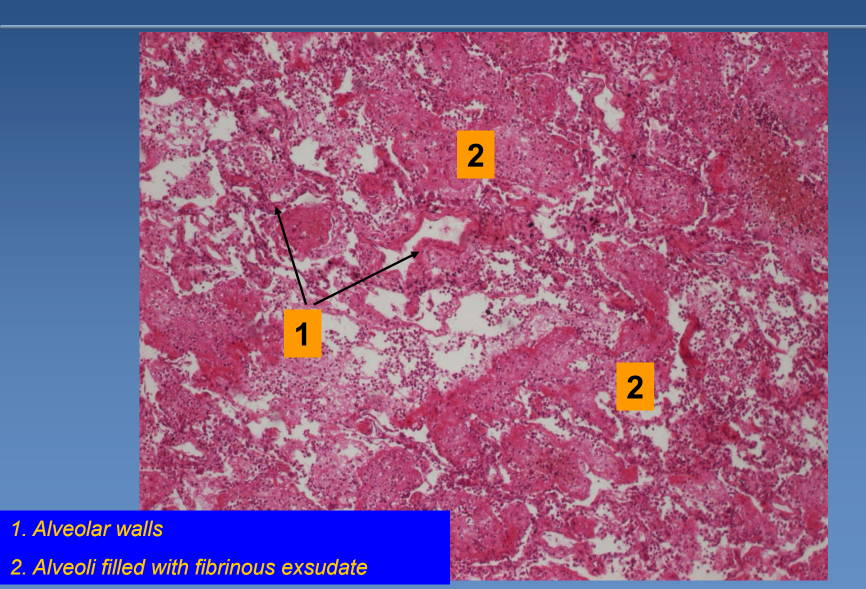


# Lobar pneumonia, grey hepatization

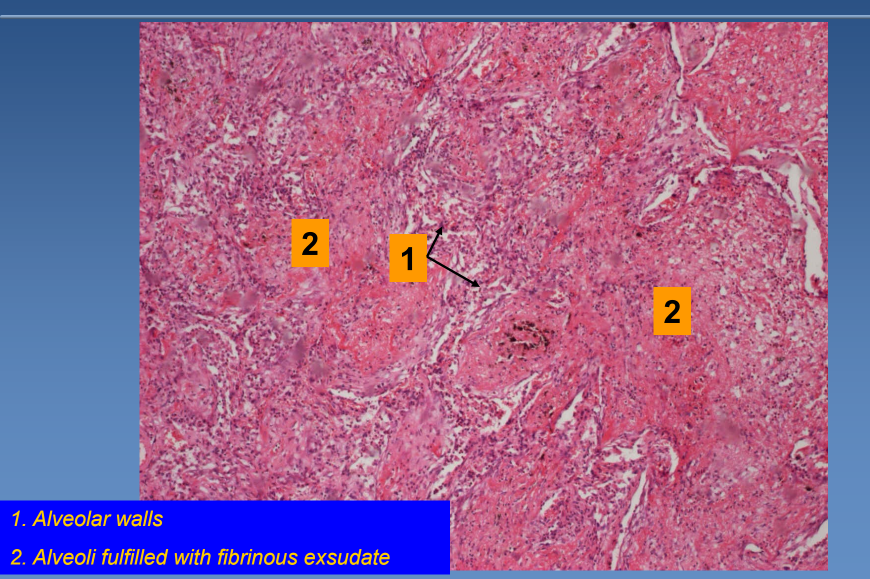




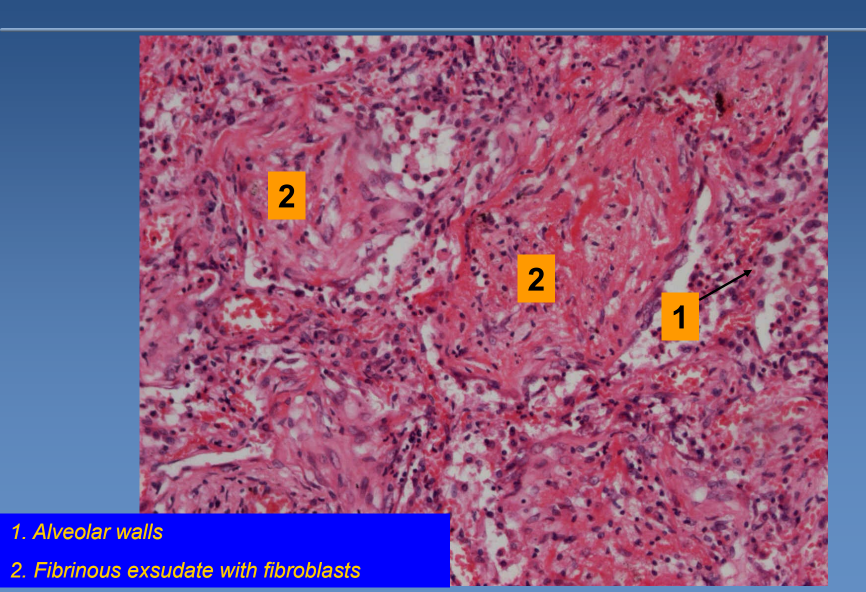












## Bronchopneumonia



- superficial type of pneumonia characterized by multiple foci of isolated, acute consolidation, affecting one or more pulmonary lobules
- inflammation spreads from bronchi
- aetiology:
  - streptococcus, staphylococcus, haemophilus, klebsiella
  - ⇒ legionella micro:
    - fibrinous purulent bronchopneumonia associated with fibrinous pleuritis
- possible secondary confluent inflammation, overlap patterns
- inflammatory complications:
  - pleuritis
  - ⇒ abscess
  - sepsis 🗢





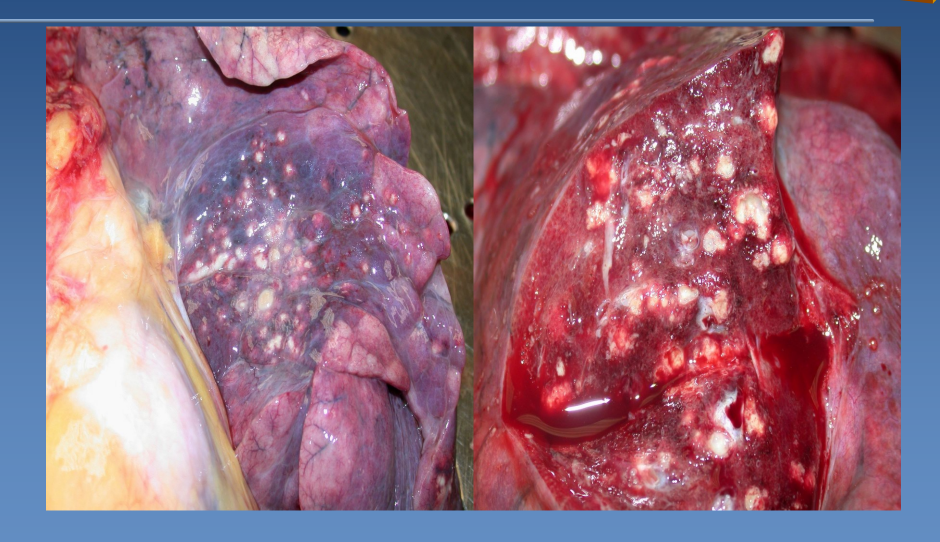
- Commonly secondary prior viral pneumonia, in chronic lung diseases, debilitating diseases, immunologic defect, aspiration, coma ...
  - various stages of inflammation in the same time
- **Gross:** 
  - oedema, hyperemic tissue with small grey-yellow foci
- Micro:
  - types of exsudate:
    - serous
    - suppurative (purulent) +/- fibrinous
  - abscessing form suppurative destruction of alveolar walls

## Bronchopneumonia



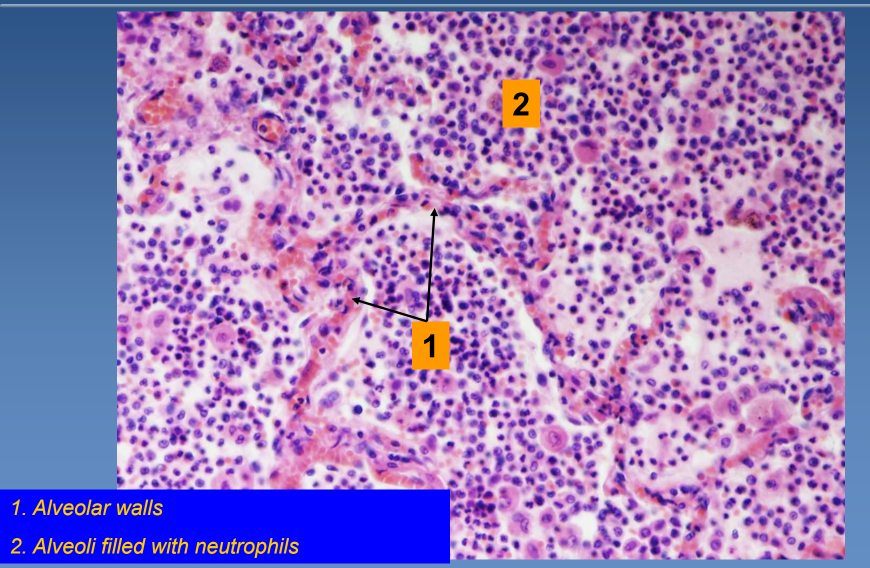


## Abscessing bronchopneumonia

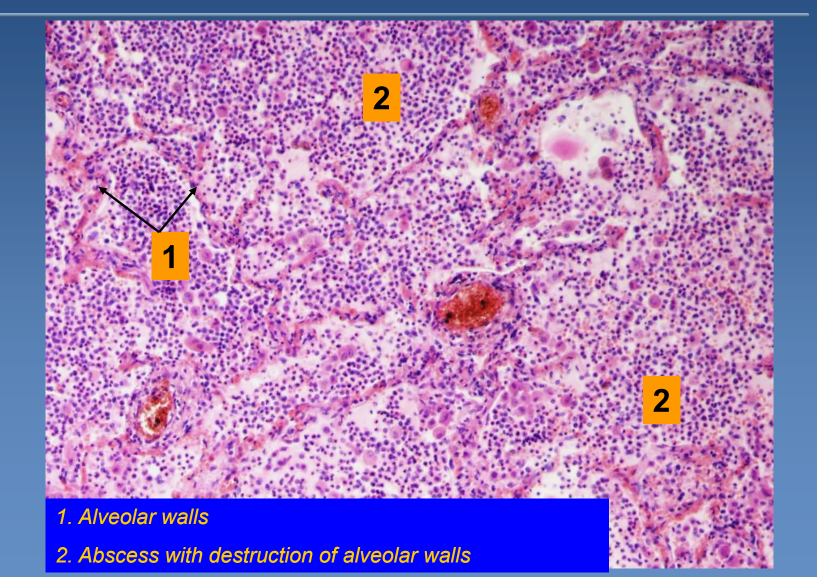


## Purulent bronchopneumonia





## Abscessing bronchopneumonia



# Infectious interstitial pneumonia



### Etiology:

- > viruses (incl. rubeola, varicella)
- mycoplasma, chlamydia, coxiella, etc.
- pneumocystis

### Symptoms:

fever, dyspnoea, dry cough, auscultation may be normal (empty alveoli), x massive changes on X-ray

### Healing:

- **⇒** ad integrum
- secondary bacterial pneumonia
- cryptogenic organizing pneumonia possible

# Infectious interstitial pneumonia



- Gross: focal / confluent, red-blue, congested, usually no pleuritis
- Micro:
  - 1) common histological features:
    - oedema and dilatation of alveolar walls
    - interstitium with mononuclear infiltrate (lymphocytes, macrophages, plasma cells)
    - possible ARDS "hyaline membranes" formation
      - necrotic pneumocytes and fibrin
      - eosinophilic material lining the lumen of alveoli

# Infectious interstitial pneumonia

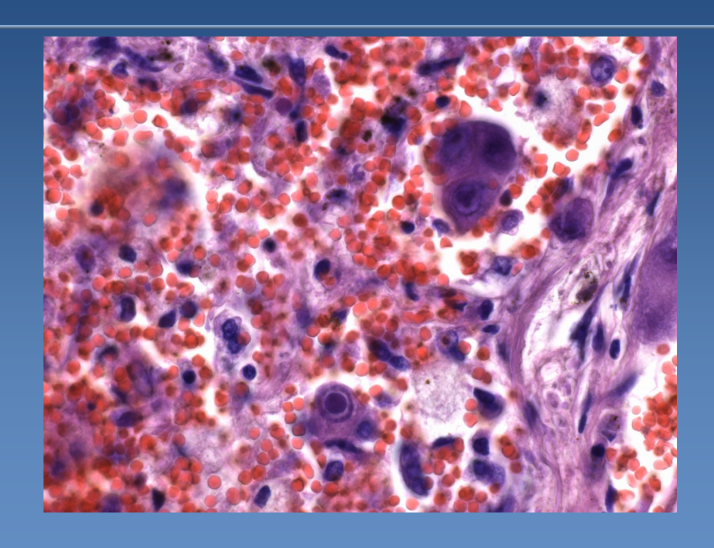


#### 2) inclusion pneumonia:

- typical inclusions and cytopatologic changes of pneumocytes
- CMV:
  - large pneumocytes with basophilic intranuclear inclusions
- Varicella, adenovirus:
  - intranuclear inclusions
- Measles:
  - giant cell pneumonia
  - multinucleated cells in alveoli and bronchioli (Warthin-Finkeldey cells)
- Pneumocystis pneumonia

## CMV pneumonia





# Pneumocystis pneumonia



#### etiology:

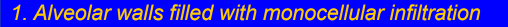
→ Pneumocystis jirovecii
(opportunistic fungal infection, immunocompromised patients)

#### Micro:

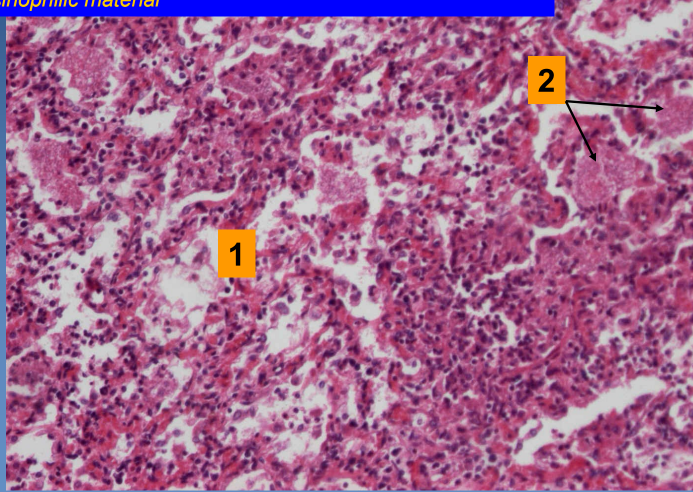
- widened alveolar septa, intraalveolar bubbly eosinophilic material:
  - pneumocystis capsules
- special histological stains:
  - Groccott silver impregnation (black)
  - Giemsa (blue)
  - PAS

# Pneumocystis pneumonia



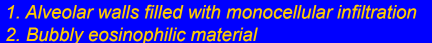


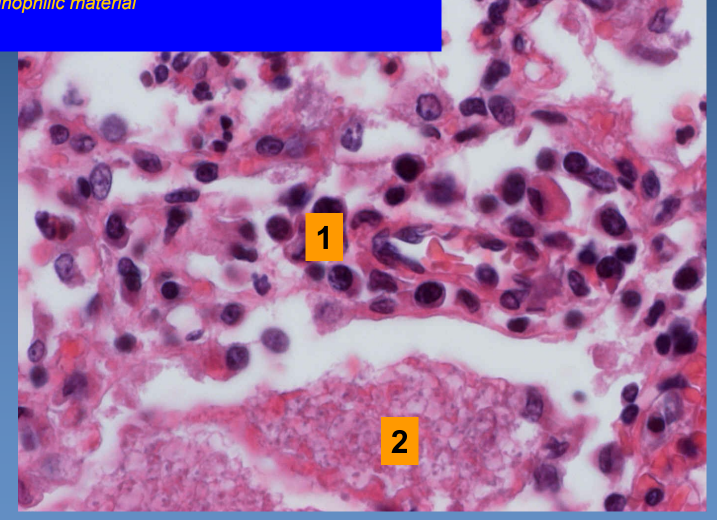
2. Bubbly eosinophilic material



## Pneumocystis pneumonia







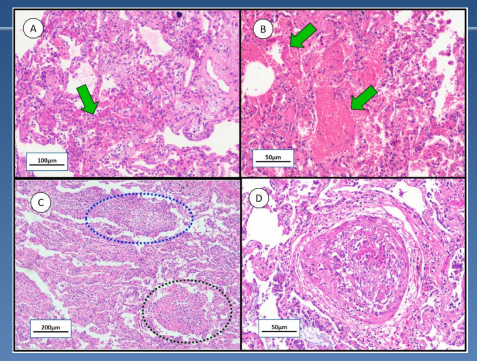
### Covid -19 in the lungs



- General appearance of interstitial viral pneumonia
- Pathogenesis
  - Cythopatic viral effect on mucosa + cillia damage
  - Surface S protein binding on ACE2 receptor
- Vasculopathy of the alveolocapillary membrane, microangiopathy + thrombosis
- Complications: ARDS, septic shock, secondary bacterial superinfection

### Covid -19 in the lungs





copy

★ Histopathological findings in COVID-19 lungs: the virus-induced lung injury with temporal heterogeneity: A - alveolar hyaline membrane (green arrow); B - alveolar-capillary barrier injury with hemorrhage (green arrows); C - acute fibrinous organizing pneumonia (dark blue circle) and organizing pneumonia (dark green circle); and D - pulmonary intravascular thrombotic events.

## Interstitial lung diseases



#### Form:

- ⇒ acute alveolar damage (ARDS, radiation pneumonitis, diffuse intrapulmonary haemorrhage Goodpasture'sy)
- chronic interstitial lung disease
  - Fibrosing
    - Idiopathic pulmonary fibrosis (Usual interstitial pneumonia)
    - Nonspecific interstitial pneumonia
    - Cryptogenic organizing pneumonia
    - Associated w. connective tissue diseases (rheumatoid arthritis)
    - Drug reaction
    - Pneumoconioses
  - Granulomatous (sarcoidosis, hypersensitivity pneumonitis extrinsic allergic alveolitis)
  - Eosinophilic
  - Smoking related (desquamative interstitial pneumonia etc.)
  - Other

- DAD (ARDS, RDS)
- clinical:
  - progressive respiratory insufficiency associated with shortness of breath and hypoxia, high mortality
- Etiology:
  - Primary ARDS:
    - lung inflammation/infection, aspiration of gastric content, mechanical trauma incl. chest contusion, fat embolism, near-drowning, ionizing radiation, inhaled irritants (smoke, chemicals),
  - **⇒** Secondary ARDS:
    - trauma (head) or sepsis
    - acute pancreatitis
    - renal insufficiency (uremia)
    - burns
    - hematologic conditions DIC, multiple transfusions
    - chemical injury (heroin overdose, acetylsalicylates, ...)

#### Gross:

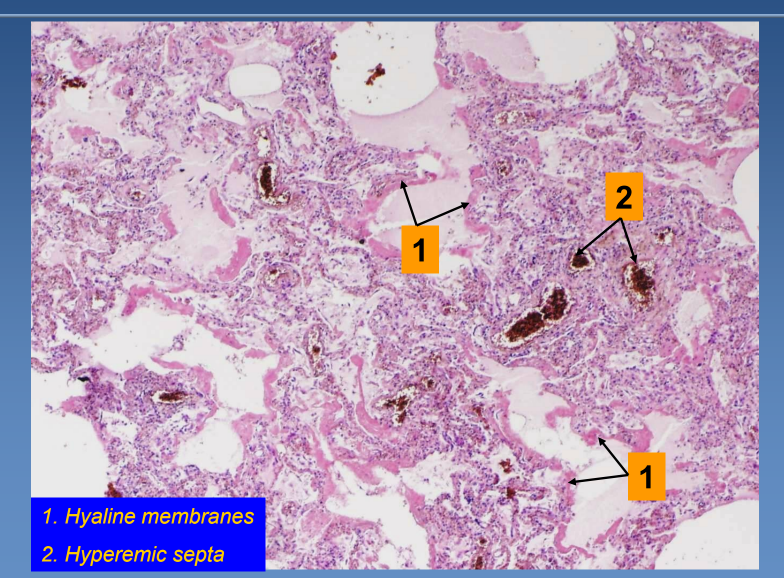
- > heavy lung
- dark red color
- **⇒** boggy

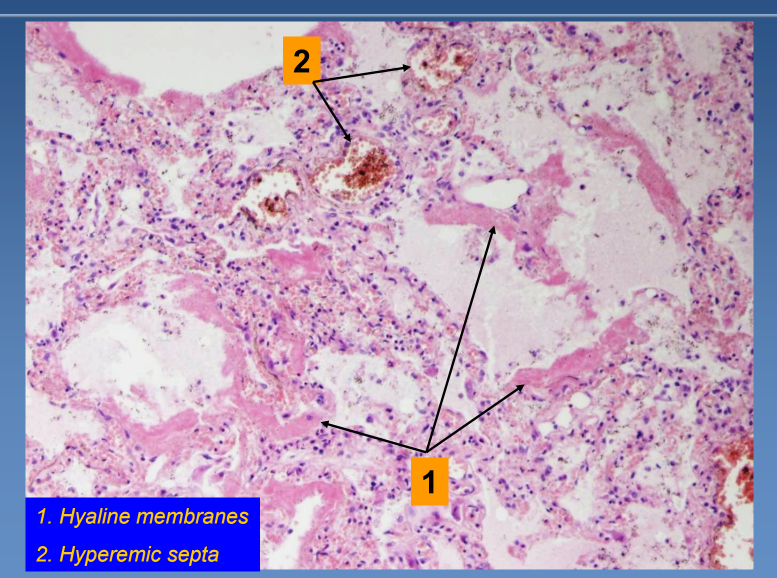
#### Micro:

- exsudative phase:
  - capillary congestion, oedema, hyaline membranes formation within 48 hours

#### proliferative phase:

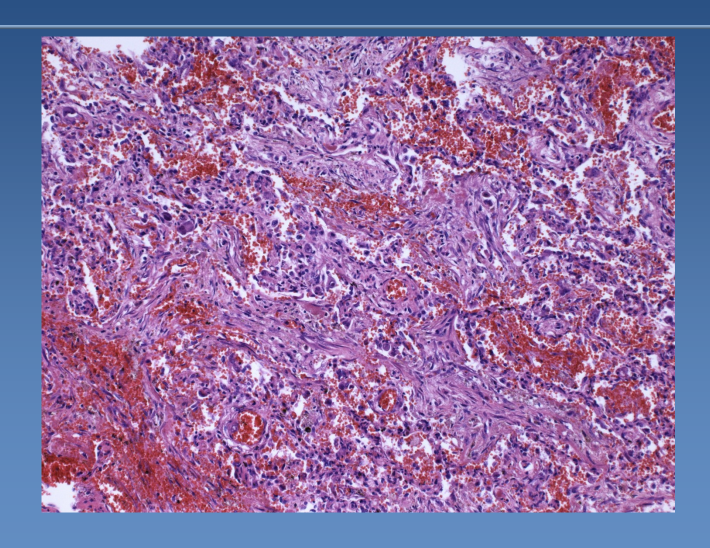
- epithelium regeneration (type II. pneumocytes)
- hyaline membranes ingested by macrophages
- proliferation of fibroblasts in alveolar walls -> pulmonary fibrosis possible





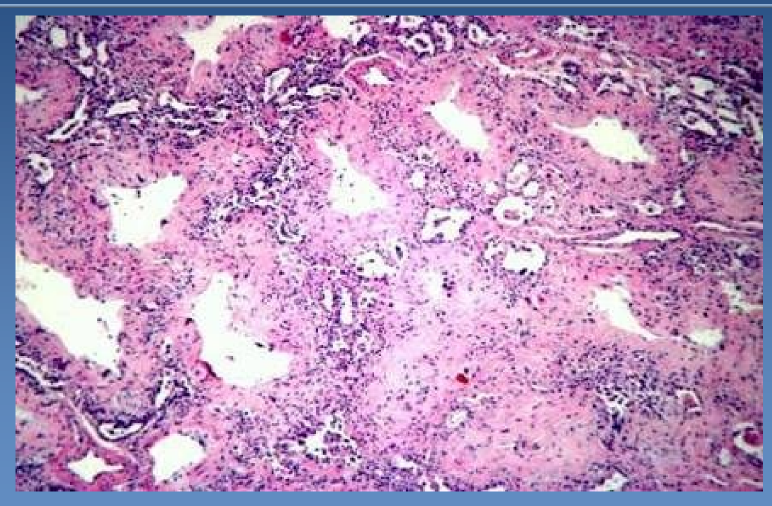
# CMV pneumonia - ARDS





### 6.3 DAD, proliferative phase - fibrotic stage - distinctly thickened interalveolar septa with a chronic inflammatory infiltrate.





# Idiopathic pulmonary fibrosis



- Clinical-radiologic-pathologic diagnosis
- synonymic cryptogenic fibrosing alveolitis
- histologic pattern of "usual interstitial pneumonia" (UIP):
- Etiology: abnormal epithelial repair myo/fibroblastic proliferation
  - intrinsic problem + exogenous factor (? occupational, smoking)
- Dismal prognosis: progressive dyspnoea, hypoxemia, lung failure in cca 3 yrs, therapy lung transplantation only

# Idiopathic pulmonary fibrosis



- usual interstitial pneumonia" (UIP):
  - >70% of all of idiopathic interstitial pneumonias
  - **d** etiology:
    - in some connective tissue diseases or in association with abnormalities of serum proteins
    - smoking, asbestosis
    - unclear

#### **→** Micro:

- •subpleural and a paraseptal foci of fibroblasts/fibrosis and chronic inflammatory infiltrate, cystic spaces honeycombing
- irregular distribution of histological features temporal heterogeneity

# Idiopathic pulmonary fibrosis



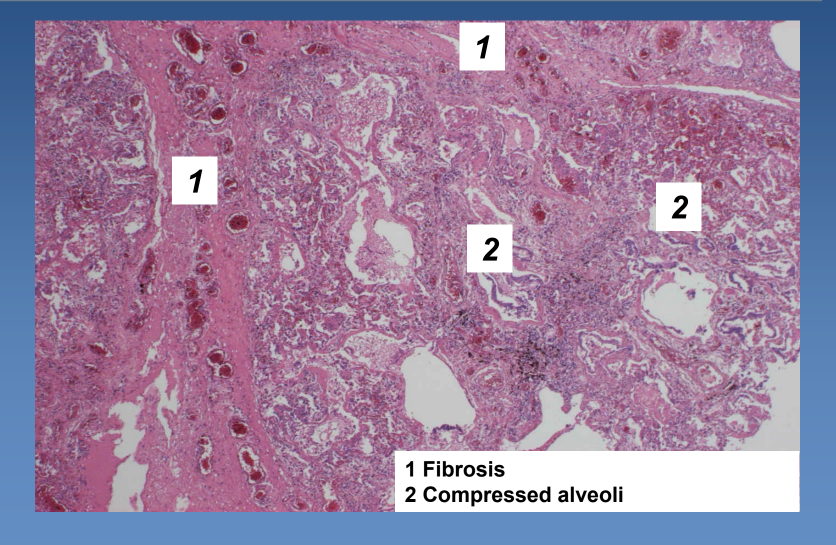
- non-specific interstitial pneumonia (NSIP):
  - different histologic/clinical pattern
  - commonly women, without smoking association
  - better prognosis
    - treated with corticosteroids

#### Micro:

- chronic interstitial inflammation +/- fibrosis
- no honeycombing
- regular distribution of changes

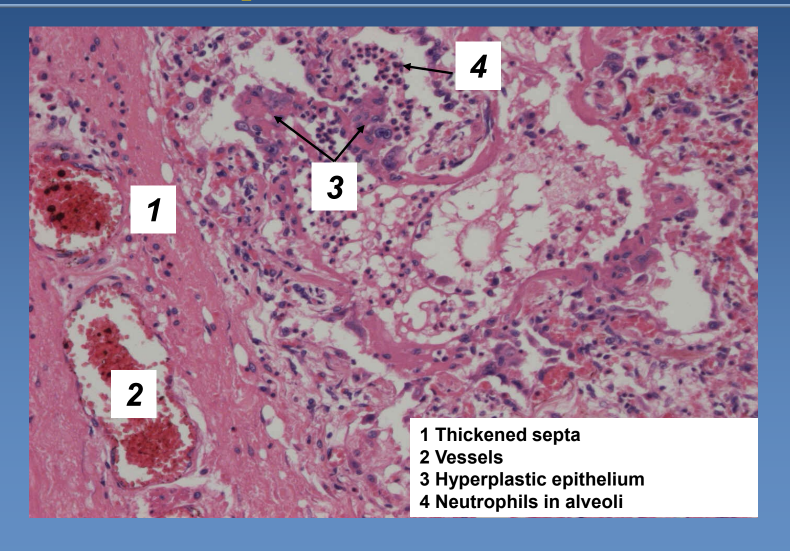
# Usual interstitial pneumonia





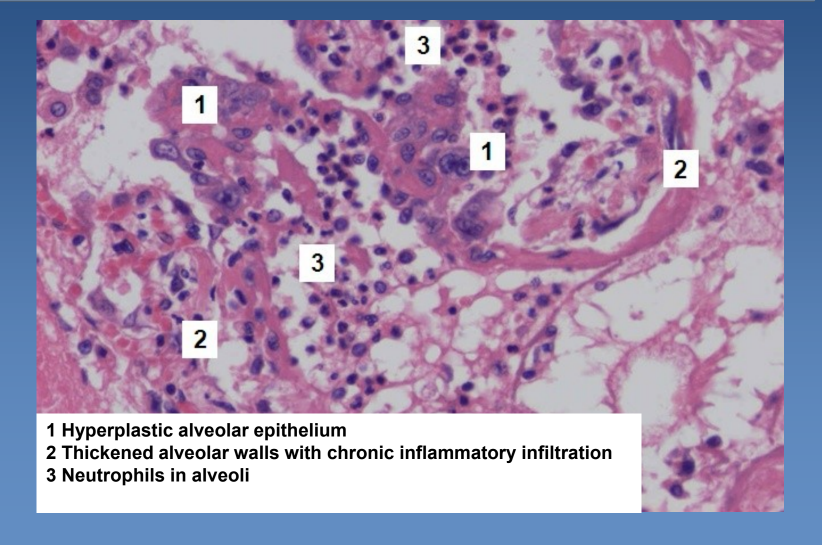
# Usual interstitial pneumonia





# Usual interstitial pneumonia





#### Pneumoconiosis



- an occupational and restrictive lung disease caused by the inhalation of specific dust
- sequels: inert (simple), fibrous, allergic, neoplastic
- high fibrogenicity of cristalline silica dust and asbestos
- 3 basic types:
  - coal-worker`s pneumoconiosis
  - **⇒** silicosis
  - ⇒ asbestosis

### Silicosis



- Chronic progressive pneumoconiosis
- Silicone dioxide particles (0,2-2µm) toxic to macrophages – focal necrosis + release of fibrogenic factors - fibrosis
- X-ray reticular fibrosis, nodules, diffuse fibrosis
- lung insufficiency
- cor pulmonale

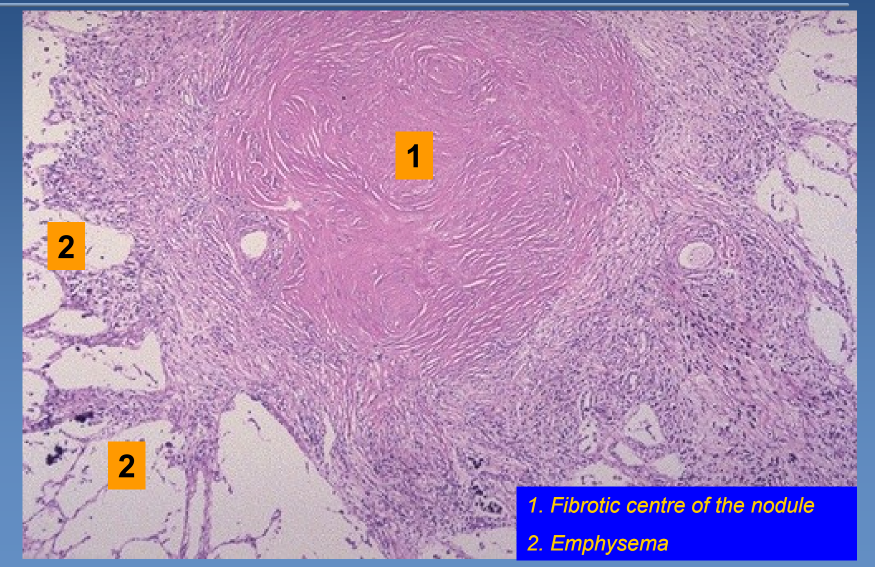
### Silicosis



- Gross (stages):
  - reticular fibrosis
  - **⇒** silicotic nodules
  - progressive massive fibrosis
- Micro:
  - nodules with concentric arrangement of hyalinized fibers and necrosis
  - anthracophages in the periphery of the nodule
  - emphysema in adjacent pulmonary tissue
  - particles seen under polarized light

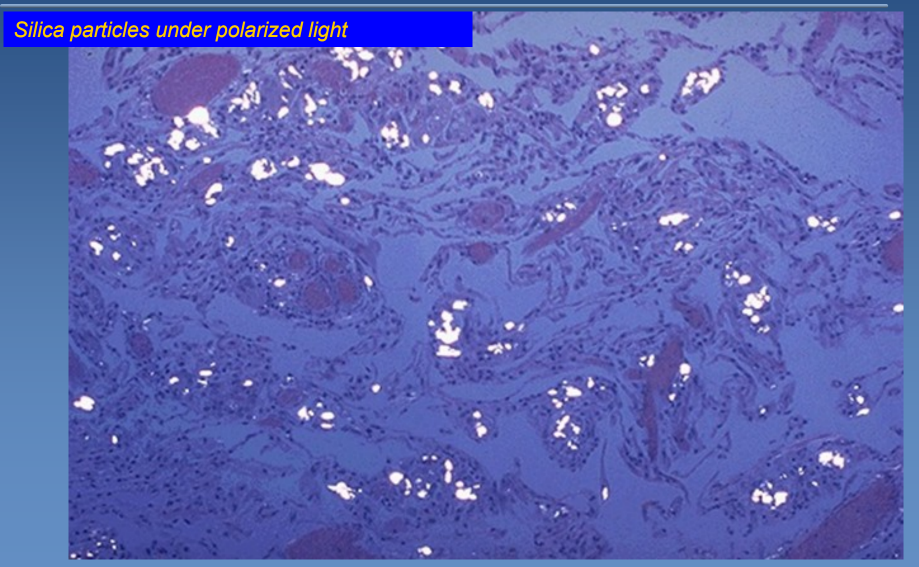






## Pulmonary silicosis





# Granulomatous inflammations - Tuberculosis

- aetiology
  - > Mycobacterium tuberculosis, M. bovis
  - special **Ziehl-Neelsen** stain
    - PCR more sensitive
- delayed-type hypersensitivity

(type IV. hypersensitivity)

→ T cells-mediated immune memory response to TBC antigens (granulomas)

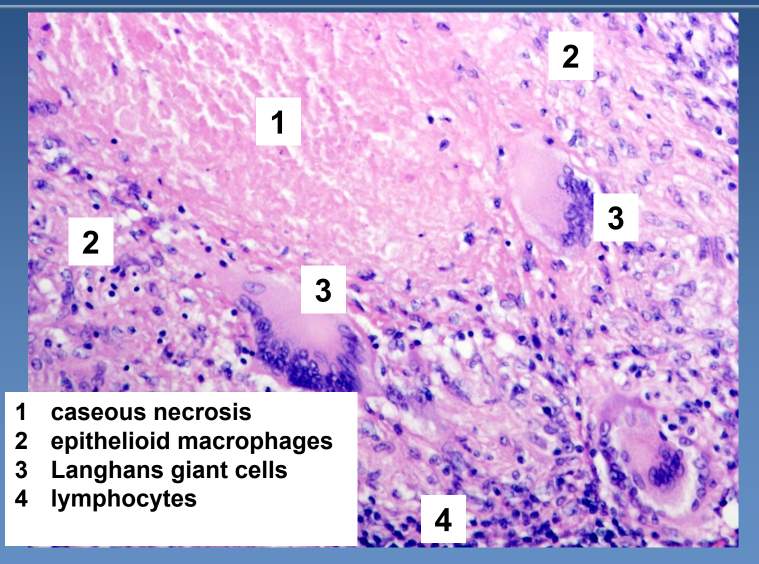
# Tuberculosis – morphological features



- tbc granuloma proliferative form
  - > host resistance
  - ⇒ specific granulation tissue: epithelioid macrophages + Langhans giant cells
- tbc exsudate exsudative form (meningitis)
  - ⇒ allergy
  - serofibrinous exsudate + Orth cells (macrophages)
- caseification
  - cheese-like, caseous necrosis sensibilization?
- + colliquation (liquefaction)
  - after release of proteolytic enzymes by neutrophils
- calcification

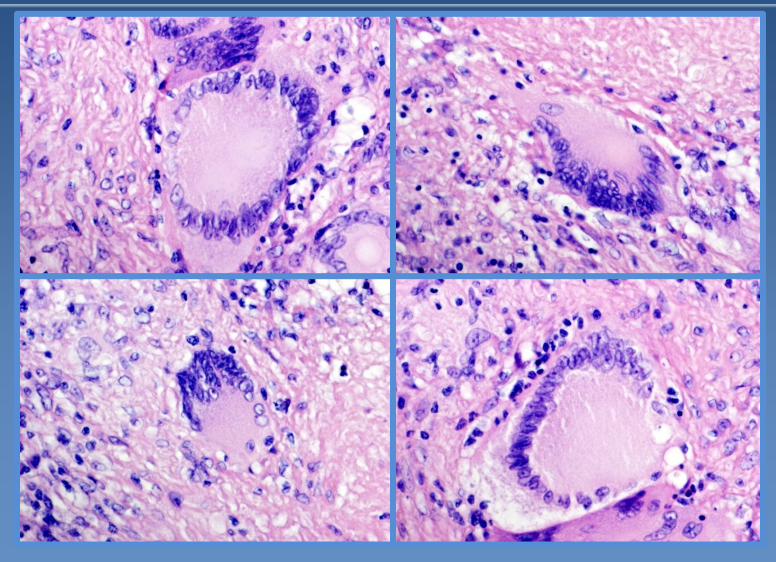
## Tbc granuloma





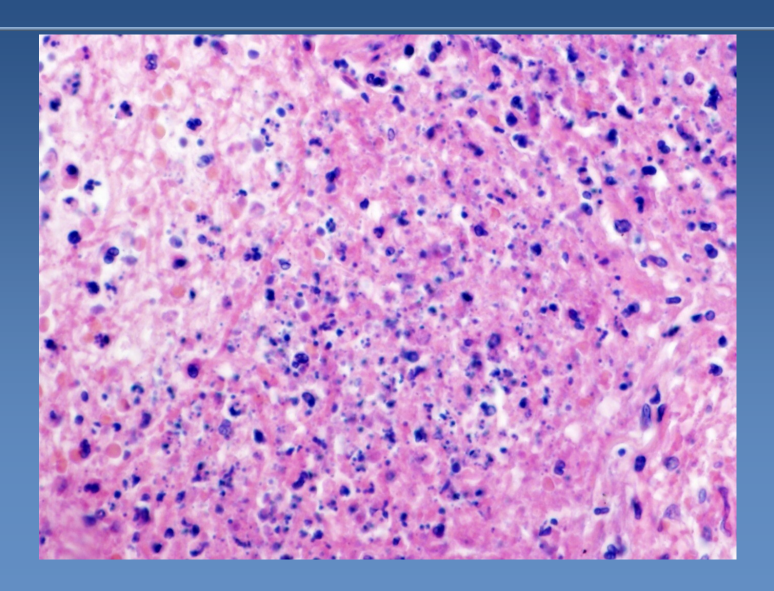
## Langhans giant cells











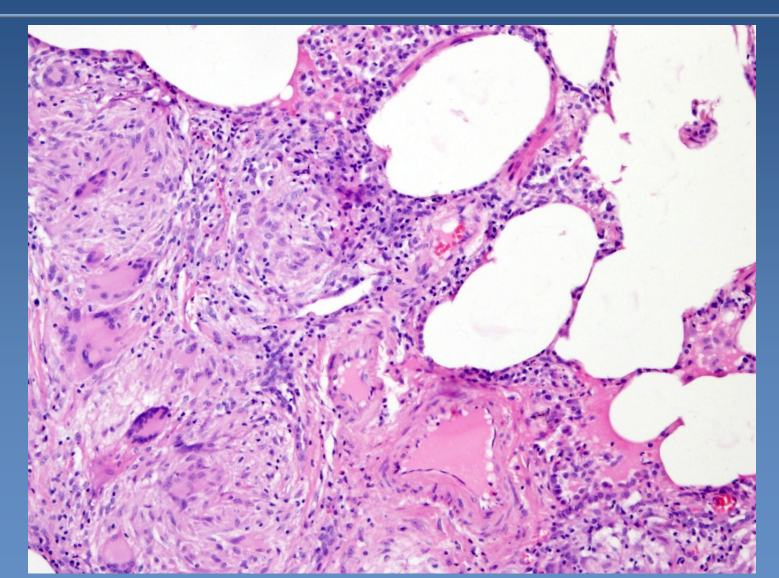
#### Sarcoidosis



- chronic granulomatous inflammatory disease of unknown aetiology
- \* affected tissue:
  - mediastinal lymph nodes, lungs, skin, eye
  - granulomas can affect any organ
- small regular granulomas similar to TBC granulomas, but without caseous necrosis, fibrosis usually more pronounced
- cytoplasmic bodies of Langhans giant cells, not specific:
  - asteroid inclusions
  - **⇒** Schaumann bodies
- dg. per exclusionem necessary elimination of TBC, fungal infection etc.

## Sarcoidosis



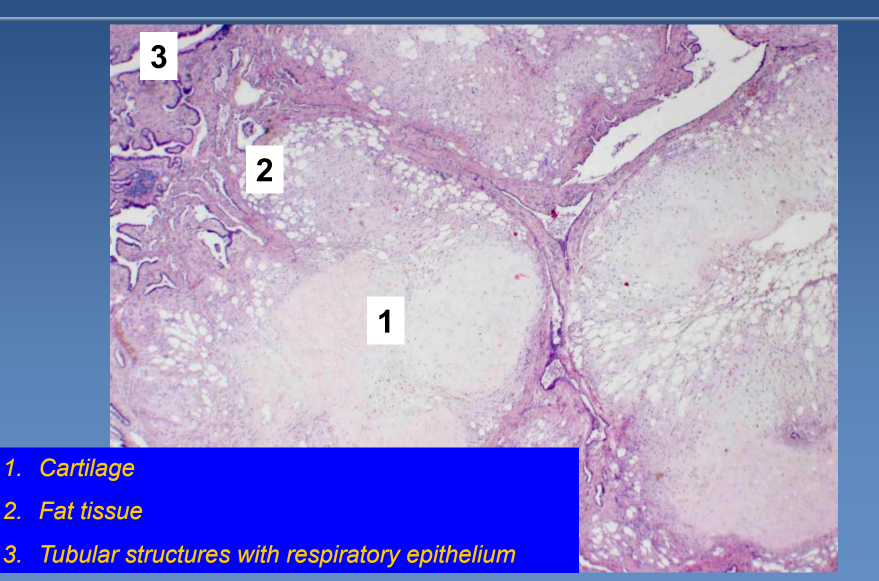


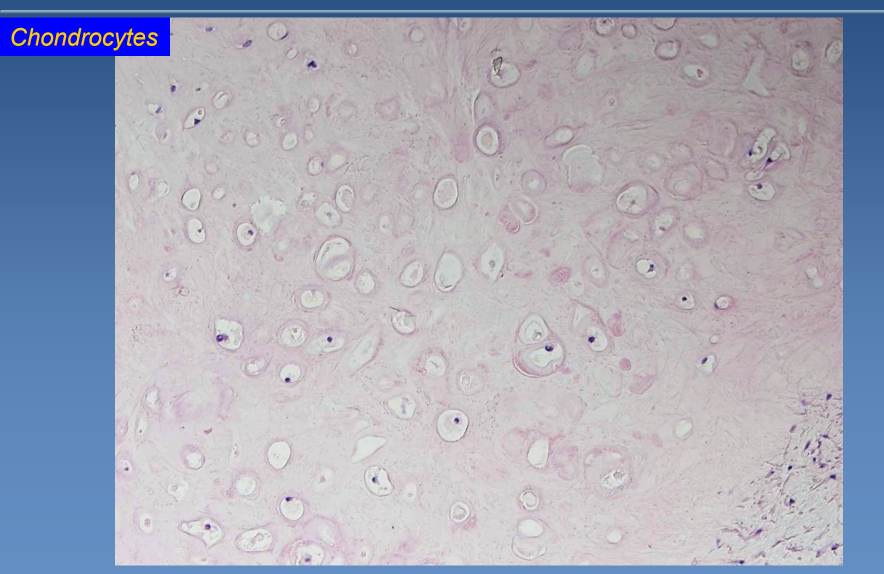
hamartoma? benign tumor?

- incidental X-ray finding
- differential diagnosis x malignant tumors important!

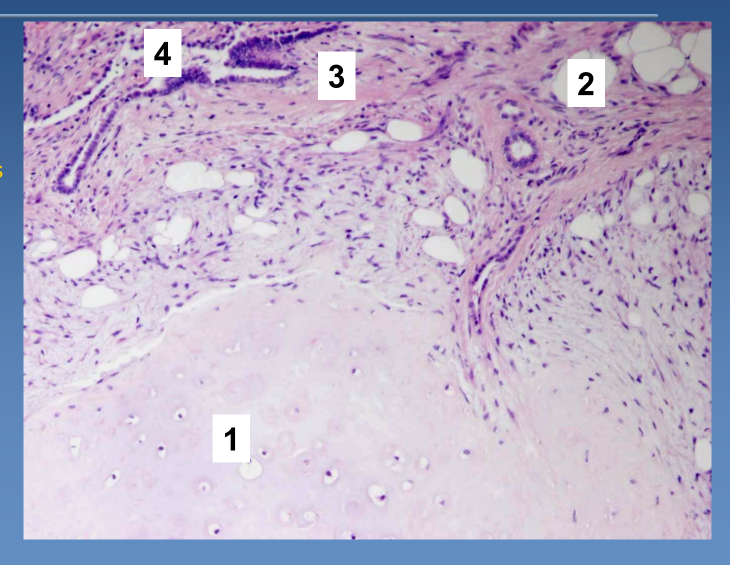
#### **×** Gross:

- whitish yellow
- well demarcated
- lobular structure
- Generally formed of mixture of homologous nonorganised afunctional tissues:
  - cartilage
  - connective tissue
  - ⇒ fat
  - tubular structures with epithelium





- Pulmonary chondrohamartoma
- 1. Cartilage
- 2. Fat tissue
- 3. Connestive tissue
- 4. Tubular structures



## Malignant lung tumors



- Primary epithelial tumors carcinomas
  - ⇒Squamous cell
  - Small cell (undifferentiated neuroendocrine ca)
  - **⇒** Adenocarcinoma
  - Large cell undifferentiated carcinoma
  - Other types (carcinoid tumors, adenosquamous ca, salivary gland tumors)
- Primary mesenchymal tumors sarcomas
- Lymphoproliferative neoplasias
- Metastatic tumors

## Bronchogenic carcinoma



- Very common primary malignancy
- ★ 5 year survival 5 7 %
- ★ 4 7 decenium, more commonly males
- Clinical symptoms: late
  - weight loss, chronic cough, haemoptysis, dyspnoea, chest pain, paraneoplastic syndromes (ACTH, ADH, PTH)

## Bronchogenic carcinoma



#### **\*** incidence:

- in CZE males 100/100 000 (the most common malignancy of men),
- ⇒ females 25/100 000 (the 3rd most common malignancy of women,

  ↑ tendency)

#### aetiology:

- **smoking** 
  - generally 20X higher risk in smokers
  - 20 cigarettes/day = 20 years, 40 cigarettes/day = 10 years...
- asbestos, Hg, Ni, As
- ionization
- radioactive radon
- dust particles
- familial predisposition

## Bronchogenic carcinoma



- local complications:
  - depends on the localization of the tumor:
    - lung collapse, bronchiectasis, bronchopneumonia, gangrene
    - widespread necrosis (more extensive in squamous cell ca)
      - destruction of vascular wall by tumor
      - fatal bleeding
- clinical types:
  - ⇒ small cell lung carcinoma (SCLC)
  - non-small cell lung carcinoma (NSCLC)

# Neuroendocrine carcinomas



- Neuroendocrine differentiation typical organoid growth pattern, neurosecretory granules, may be paraneoplastic syndromes – aberrant production of peptide hormones
- **Well-differentiated n. tumor, G1** carcinoid, i. e. in GIT, bronchi ...
- Moderately differentiated n. t. atypical carcinoid, G2
- ➤ Undifferentiated n. c. variable cell size, most common small cell carcinoma



- undifferentiated (high grade) neuroendocrine tumor
- 20 % of all bronchogenic carcinomas
- associated with smoking
- localized in lung hilus
- early metastatic spread, widespread dissemination
  - ⇒ lymphatic and hematogenous (LN, liver, brain, bones, kidney, adrenals, ...)



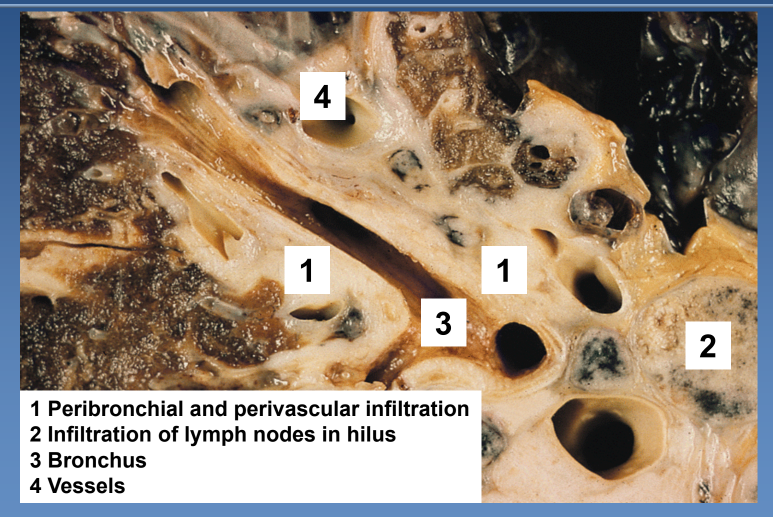
#### histologic types:

- ⇒small cell ("oat cell carcinoma")
- intermediate (now included into small cell type)
- **⇒** combined

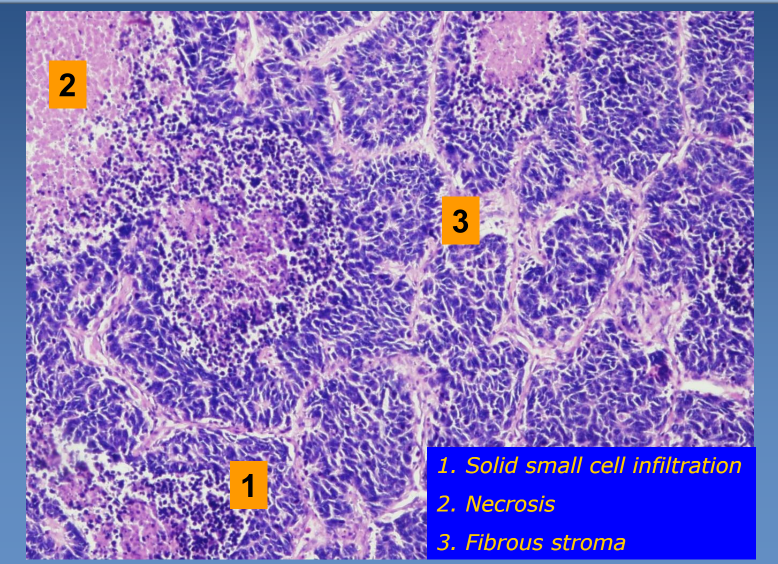
#### Micro:

- small cells with scant cytoplasm (size < 3 lymphocytes)</p>
- small round elongated dark blue nuclei without obvious nucleoli (oat cell carcinoma)
- solid growth
- neurosecretory granules in cytoplasm
  - chromogranin, synaptophysin

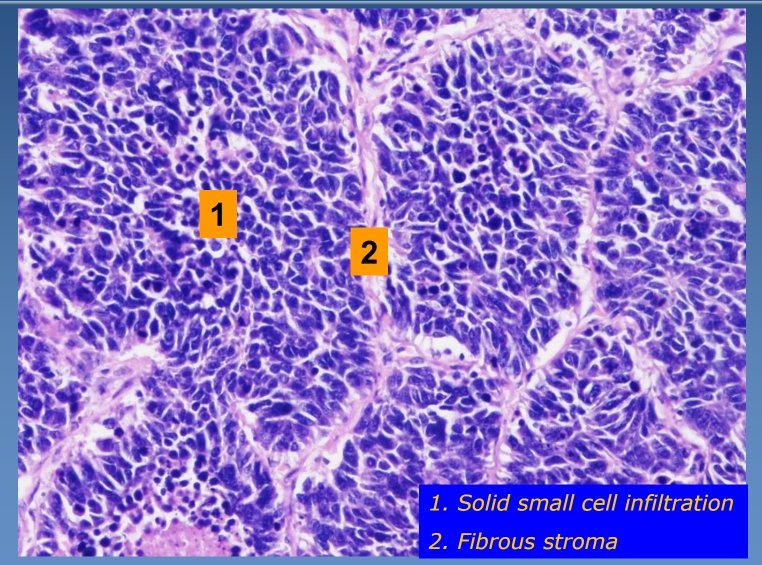












# Non-small cell lung carcinoma



- squamous cell carcinoma
- adenocarcinoma
  - adenocarcinoma in situ
  - minimally invasive:
    - non-mucinous
    - mucinous
    - mixed
  - invasive:
    - lepidic
    - acinar
    - papillary
    - micropapillary
    - solid
- ✗ large cell lung carcinoma
- other, incl. mixed

## Squamous cell carcinoma



- **\*** male 40%, female 20%
- strongly associated with smoking
- typical perihilar localisation (central>peripheral)
- commonly slow progression from squamous metaplasia – dysplasia – ca in situ
  - ⇒late metastases
- Micro:
  - squamous cell carcinoma of common type
    - polygonal shaped cells in solid nests, keratin pearls, cell junctions
  - variable differentiation

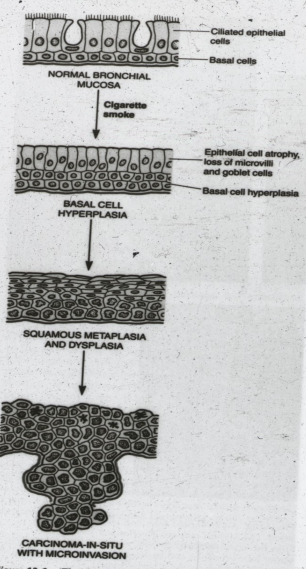
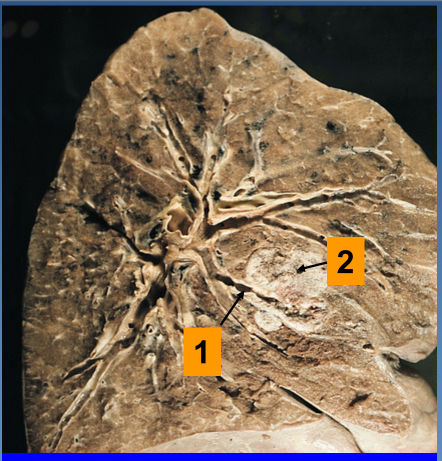


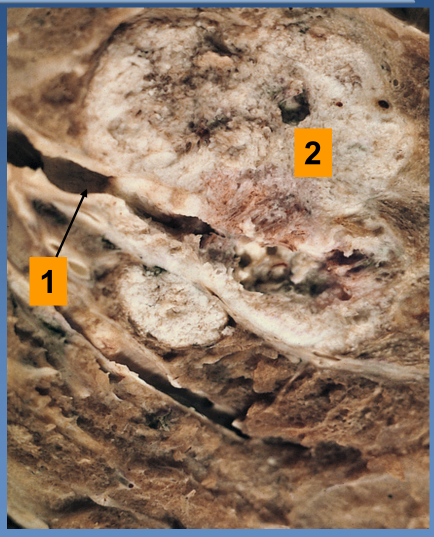
Figure 12-8. The development of squamous cell carcinoma. First, minor changes occur in the bronchial epithelium (basal cell hyperplasia), followed by squamous cell metaplasia with dysplasia. Then carcinoma in situ ensues, first localized to the epithelium and then extending to bronchial glands and penetrating the basement membrane. Extensive invasion and metastases follow.



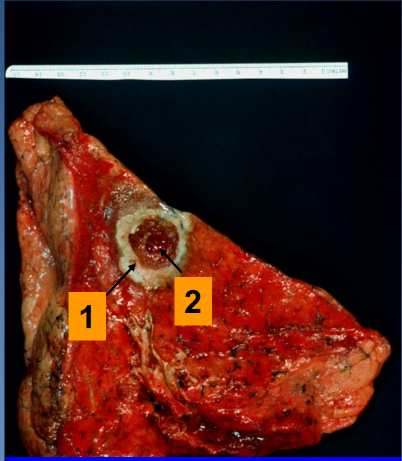
## Squamous-cell lung carcinoma



- 1. Segmental bronchus
- 2. Tumor



## Squamous cell lung carcinoma



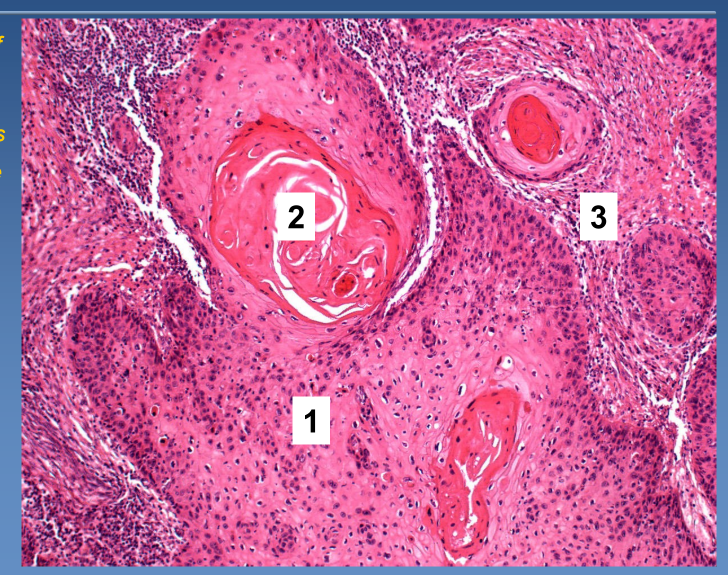
- 1. Tumor localized in the periphery
- 2. Central necrosis



- 1. Tumor in bronchus
- 2. Segmental bronchus

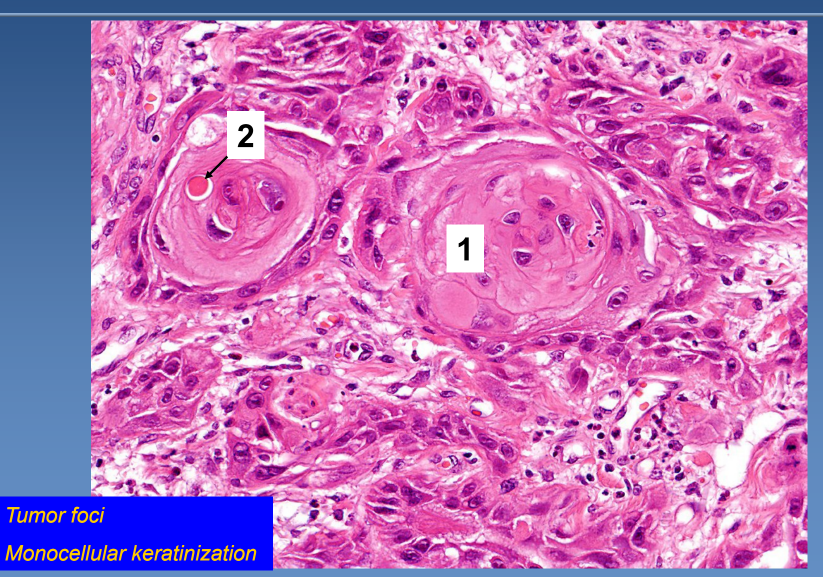
## Squamous cell carcinoma

- 1. Solid nests of malignant keratinocytes
- Keratin pearls
- Stroma of the tumor



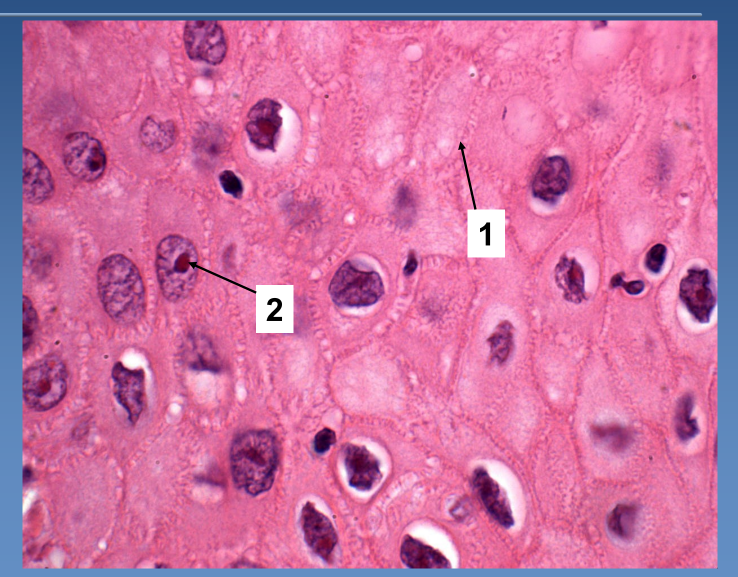
## Squamous cell carcinoma





## Squamous-cell carcinoma

- 1. Cell junctions
- 2. Nucleus with prominent nucleoli





- male 20%, female 40%;
- most cases in smokers, but the most common type in non-smokers
- typically localized in the periphery, subpleural
  - □ late symptoms !!! Commonly accidental finding on X-ray/CT
- formerly used term:
  - bronchioloalveolar adenocarcinoma (BAC) no more in use (but still present in WHO classification of lung tumors)



#### classification:

- Adenocarcinoma in situ AIS (size ≤3 cm):
  - non/mucinous (earlier BAC),
  - mucinous
  - mixed
  - no stromal/vascular/pleural invasion present

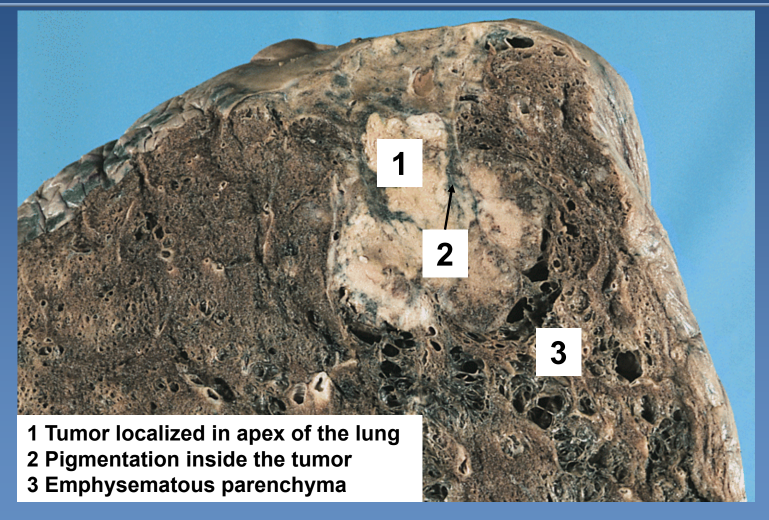
#### Minimally invasive ACA (size ≤3 cm and ≤ 5 mm invasion): idem

- apart of lepidic growth other types of spread (papillary, solid....) or stromal invasion present
- no vascular/pleural invasion present

#### > Invasive ACA:

- Lepidic
- Acinar
- Papillary
- Micropapillary
- Solid

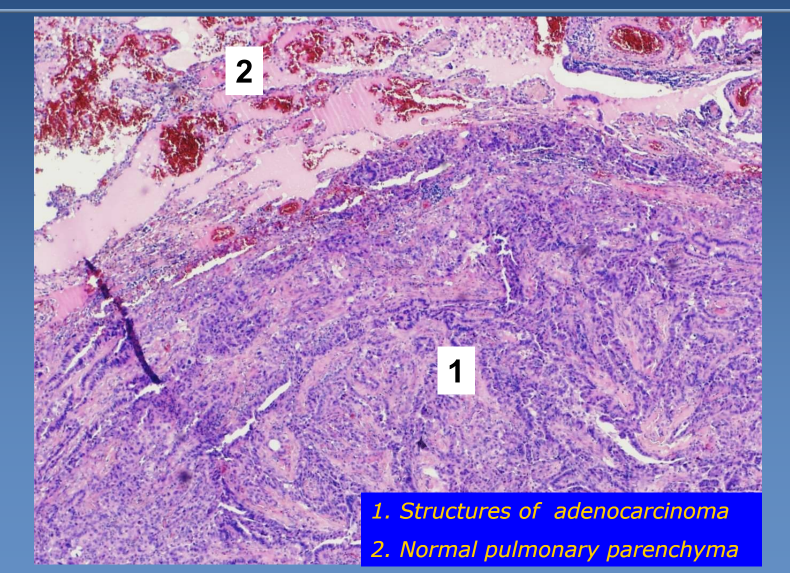






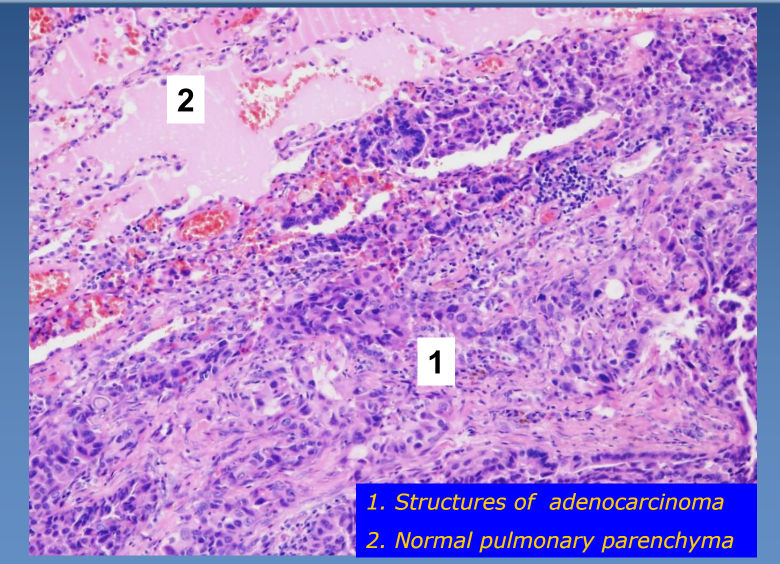




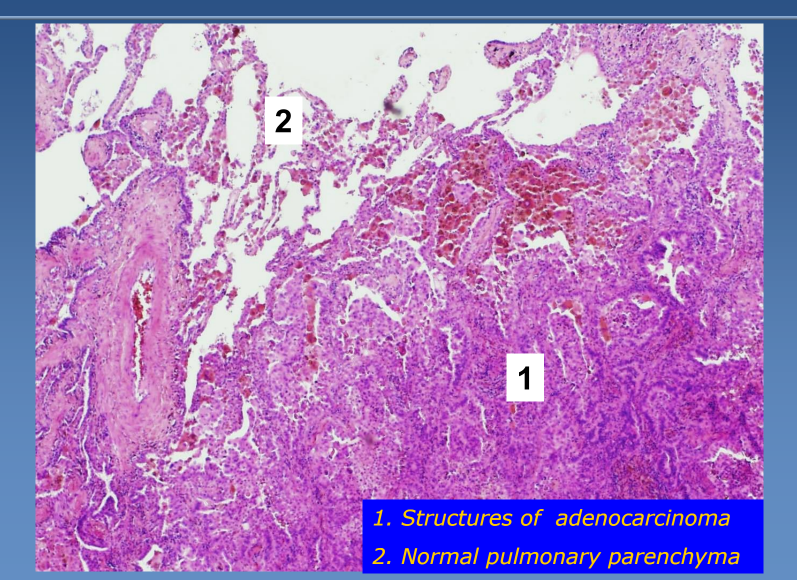






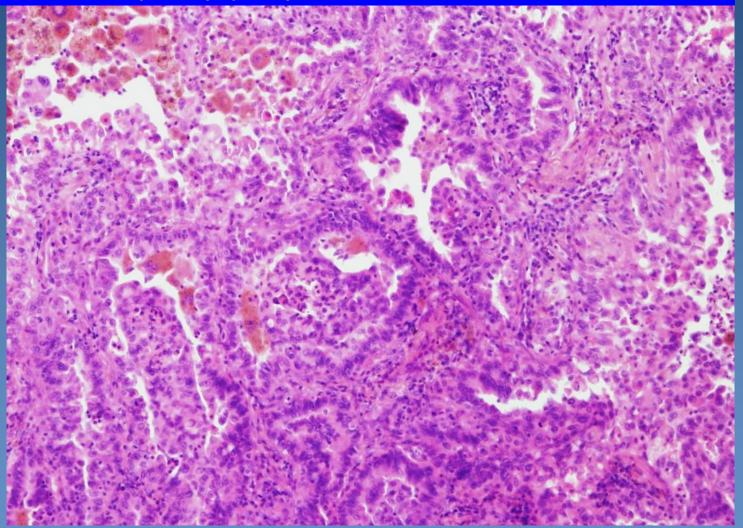






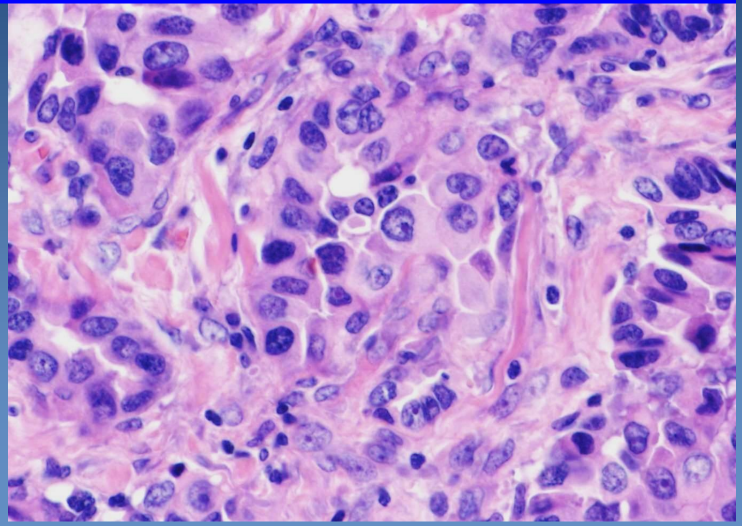


Structures of an acinary and papillary formed adenocarcinoma



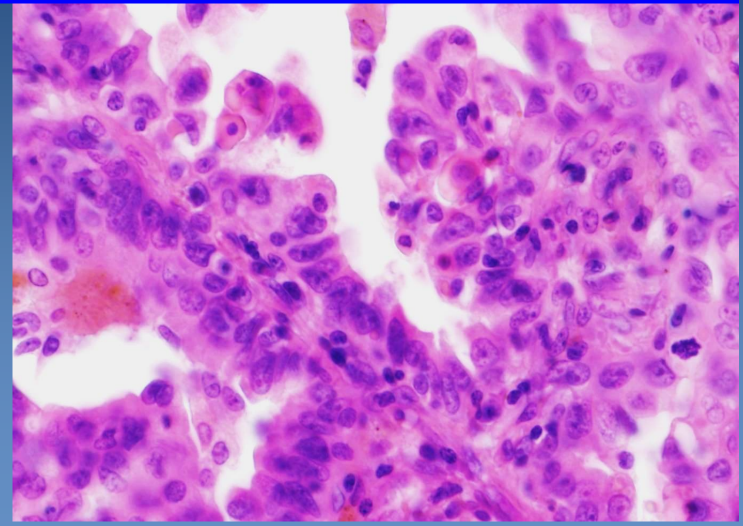


Cytology of malignant cells - anisocytosis and anisokaryosis

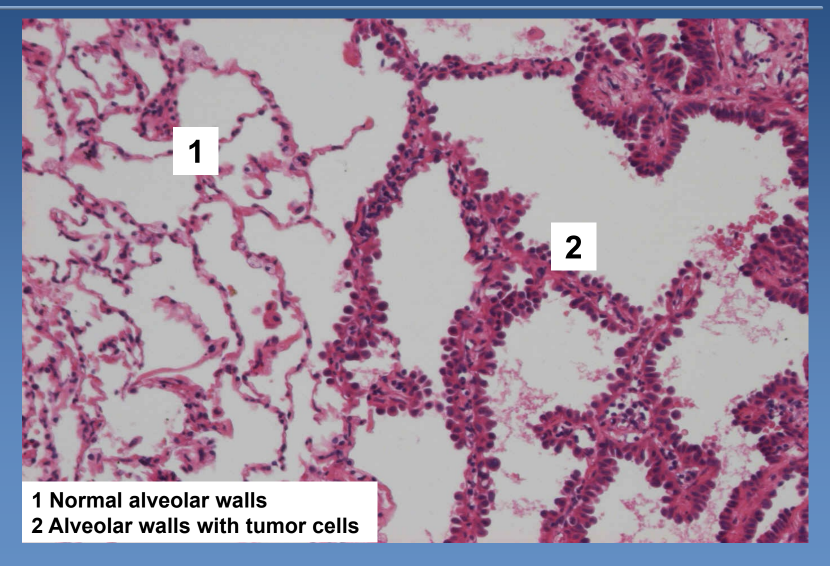




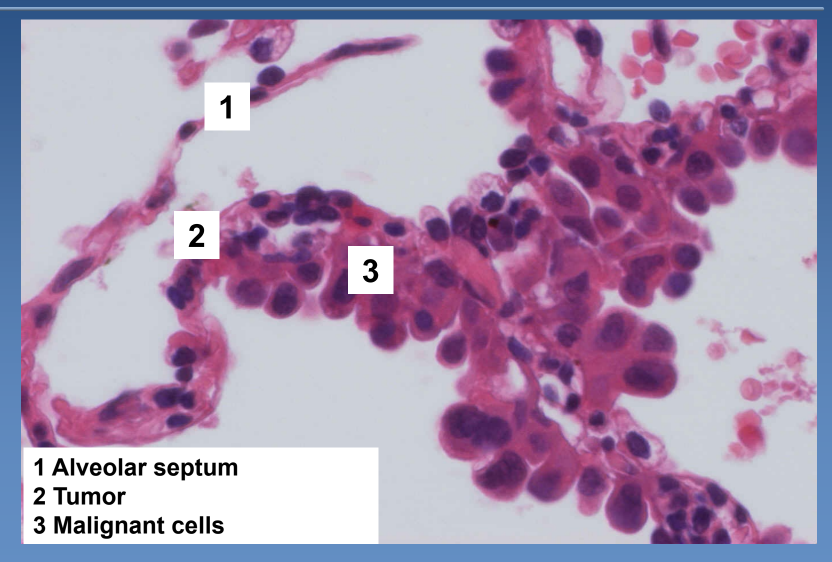
Cytology of malignant cells - anisocytosis and anisokaryosis



# AIS/minimally invasive ACA non/mucinous (earlier BAC)



# AIS/minimally invasive ACA non/mucinous (earlier BAC)



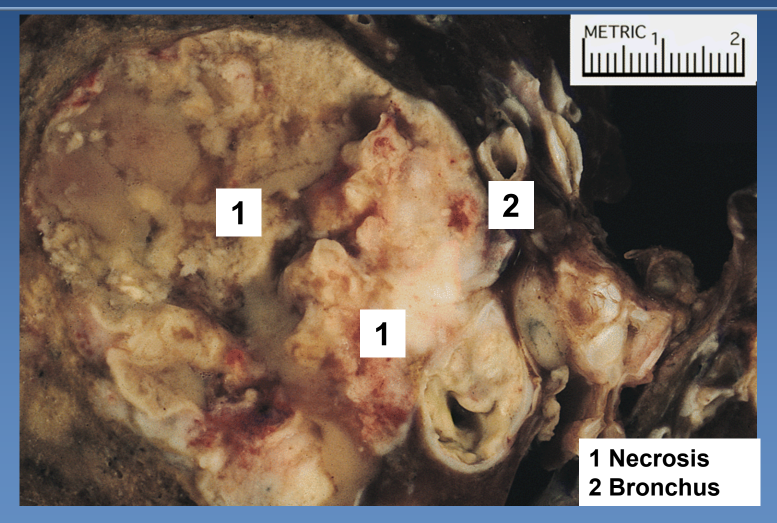


undifferentiated non-small cell carcinoma

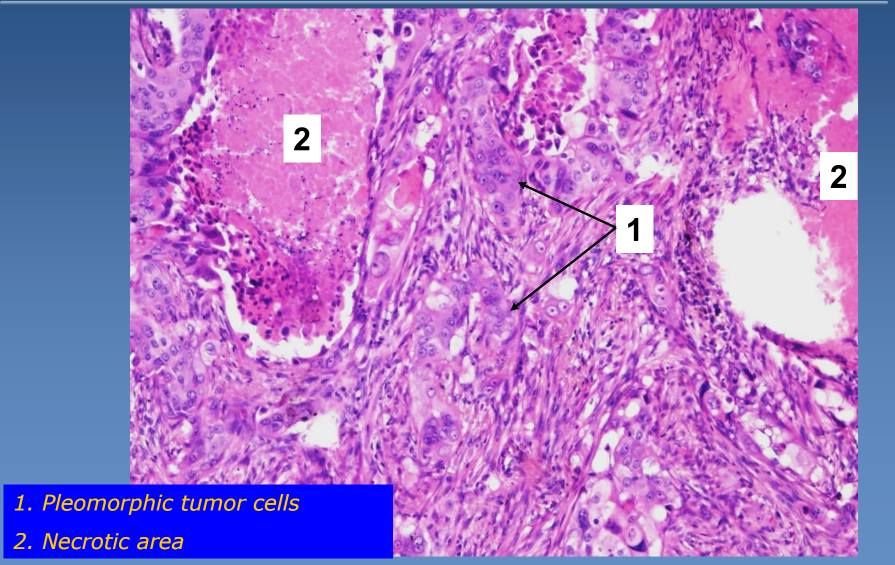
- Micro:
  - atypical pleomorphic cells

absent features of small cell carcinoma, adenocarcinoma or squamous cell carcinoma

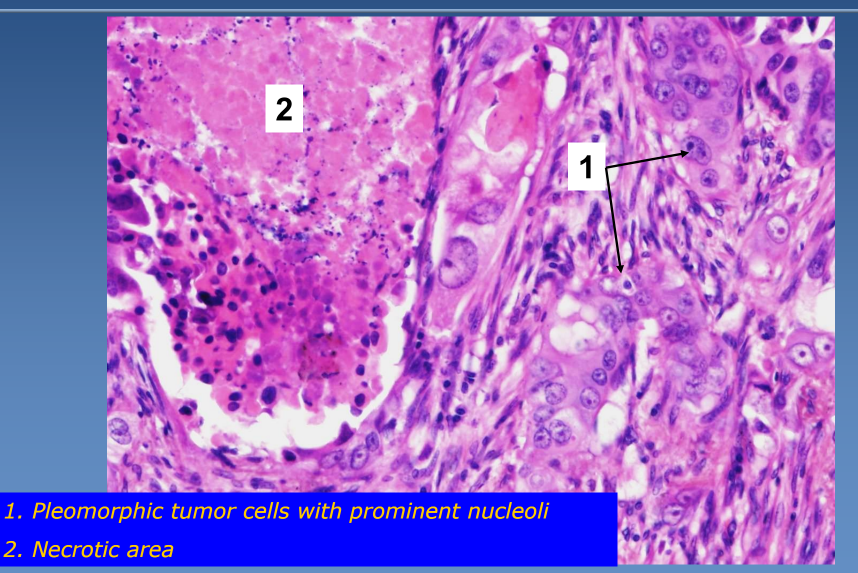












### Mesothelioma



- primary pleural tumor
- by far less common than secondary metastases of other tumors
- mostly malignant
- risk factor: chronic exposition to asbestos
- **\***Gross:
  - localized form
  - diffuse form
- **™**Micro:
  - pepithelioid, sarcomatoid, biphasic, desmoplastic