III- ASTHMA



https://pixy.org/1266846/

III. ASTHMA

Chronic inflammatory disorder of the airways

 Causes recurrent episodes of wheezing, Dyspnea, chest tightness and cough particularly at night and/or early in the morning

its hallmarks are:

- a) Intermittent and reversible airway obstruction (bronchospasm)
- b) Chronic bronchial inflammation with eosinophils,
- Bronchial smooth muscle cell hypertrophy and hyperreactivity.
- d) increased mucus secretion.

MAJOR FACTORS:

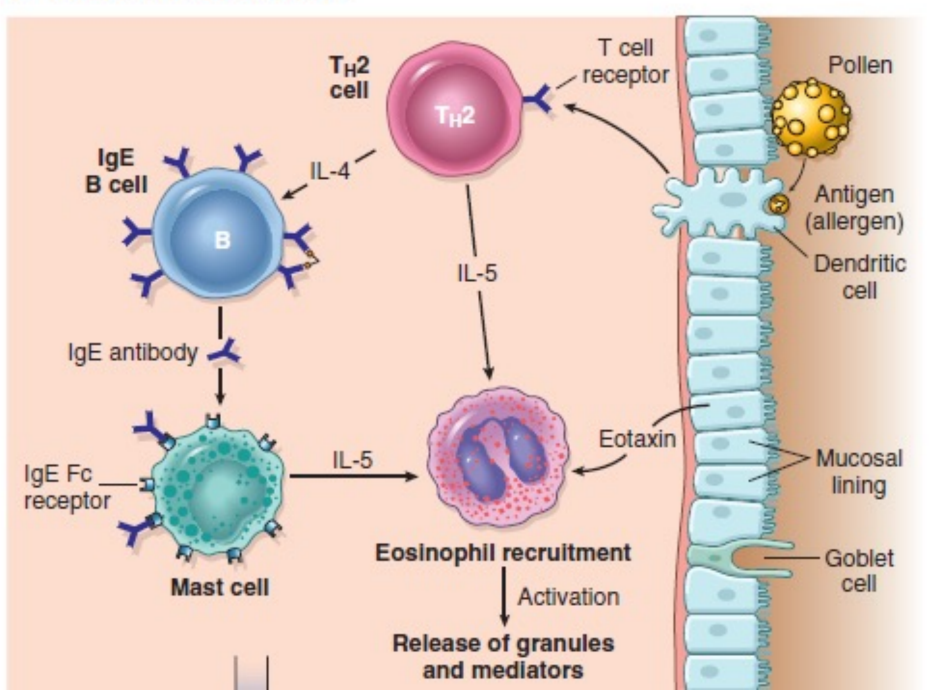
- ✓ Genetic predisposition to type I hypersensitivity (atopy)
- ✓ Acute and chronic airway inflammation
- ✓ Bronchial hyperresponsiveness to a variety of stimuli

CAN BE TRIGGERED BY:

- ✓ respiratory infections (especially viral)
- ✓ airborne irritants (smoke, fumes)
- ✓ cold air
- ✓ Stress
- ✓ exercise

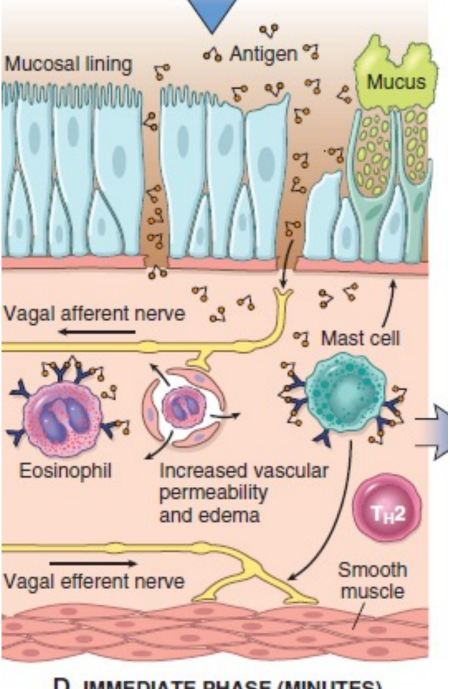
PATHOGENESIS

C TRIGGERING OF ASTHMA



The early-phase reaction is dominated by:

- ✓ bronchoconstriction
- ✓ increased mucus production
- ✓ vasodilation.



on re-exposure to antigen (ag)→ immediate reaction

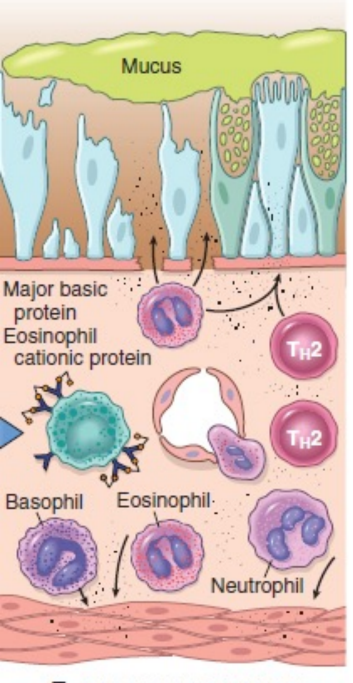
triggered by Ag-induced cross-linking of IgE bound to Fc receptors on mast cells.

mast cells release preformed mediators that directly and via neuronal reflexes induce: bronchospasm, increased vascular permeability, mucus production recruitment of leukocytes

IMMEDIATE PHASE (MINUTES)

The late-phase reaction is inflammatory:

Inflammatory mediators \rightarrow stimulate epithelial cells to produce chemokines (eotaxin) \rightarrow recruit TH2 cells, eosinophils, and other leukocytes \rightarrow amplifying the inflammatory reaction.



Leukocytes recruited to the site of reaction (neutrophils, eosinophils, and basophils; lymphocytes and monocytes)

→ release mediators → initiate the late phase of asthma.

eosinophils release major basic protein and eosinophil cationic protein that cause damage to the epithelium

E LATE PHASE (HOURS)

- Repeated bouts of inflammation lead to structural changes in the bronchial wall → called airway remodeling, including:
- ✓ deposition of subepithelial collagen
- ✓ hypertrophy of bronchial smooth muscle
- ✓ hypertrophy of Mucus glands
- ✓ increased vascularity

accumulation of mucus in the bronchial lumen

increased number of mucus-secreting goblet cells

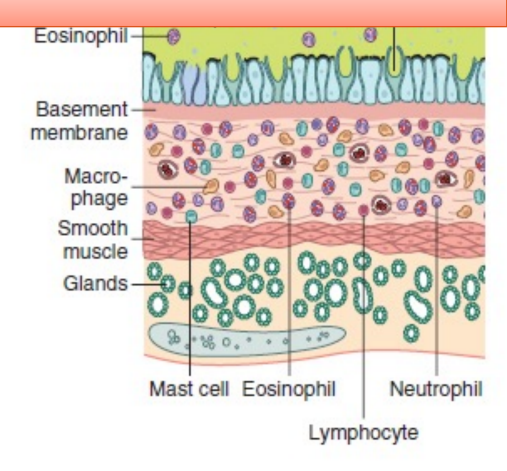
thickened basement membrane

intense chronic inflammation

hypertrophy and hyperplasia of smooth muscle cells

hypertrophy of submucosal glands





TYPES OF ASTHMA

ATOPIC ASTHMA:

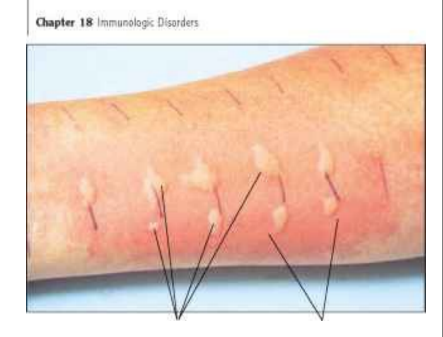
- The most common
- Classic example of type I IgE—mediated hypersensitivity reaction
- beginning in childhood
- Positive family history of atopy and/or asthma
- attacks are preceded by allergic rhinitis, urticaria, or eczema
- Attacks are triggered by allergens in dust, pollen, animal dander, or food, or by infections.

- Exposure to the antigen → excessive activation of type
- 2 helper cells → Cytokines production →
 - ✓ IL-4 and IL-13 stimulate IgE production
 - ✓ IL-5 activates eosinophils
 - ✓ IL-13 also stimulates mucus production
- IgE coats submucosal mast cells → upon re exposure → release of Mast cell-derived mediators → produce two waves of reaction:
 - early (immediate) phase of reaction
 - late phase of reaction

Diagnosis:

Skin test with the antigen: immediate wheal-and-flare reaction

(eg; skin prick test)



 serum radioallergosorbent tests (RASTs): blood test using radioimmunoassay to detect specific IgE antibodies

2- NON-ATOPIC ASTHMA:

No evidence of allergen sensitization

Negative skin test

A positive family history of asthma is less common.

- Triggered by:
 - viral respiratory infections (rhinovirus, parainfluenza virus)
 - inhaled air pollutants (sulfur dioxide, ozone, nitrogen dioxide).

3- DRUG-INDUCED ASTHMA:

- Eg: Aspirin induced asthma →
 - present with recurrent rhinitis, nasal polyps, urticaria, and bronchospasm.
- The precise pathogenesis is unknown → involve some abnormality in prostaglandin metabolism from inhibition of cyclooxygenase by aspirin

4- OCCUPATIONAL ASTHMA

Asthma attacks usually develop after
 repeated exposure to the antigen at the workplace

 triggered by fumes (plastics), organic and chemical dusts (wood, cotton, platinum), gases (toluene), and other chemicals.



MORPHOLOGY

MORPHOLOGY

- occlusion of bronchi and bronchioles by thick mucous plugs
- mucous plugs contain whorls of shed epithelium called Curschmann spirals.



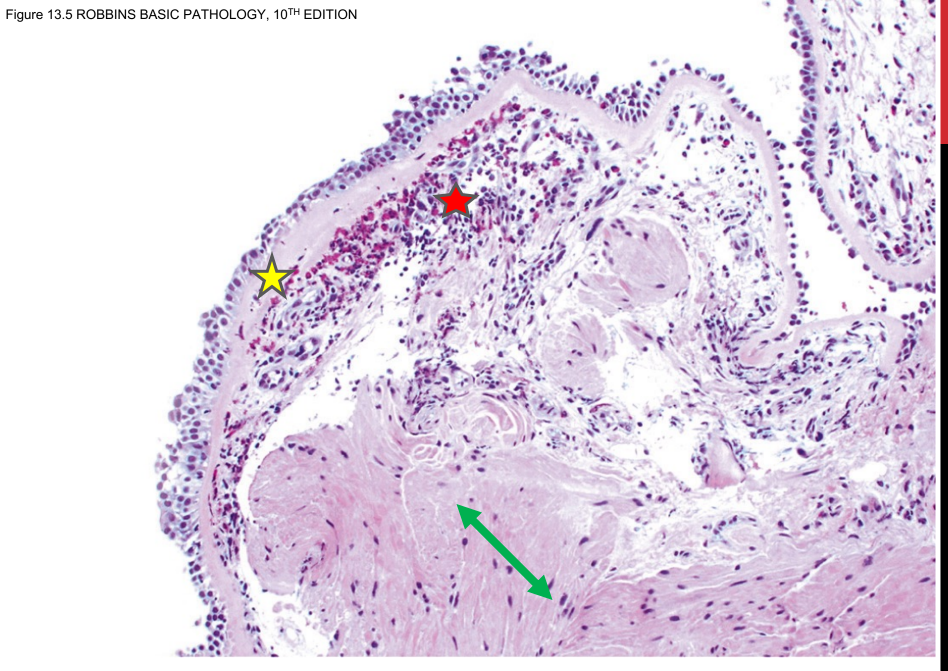
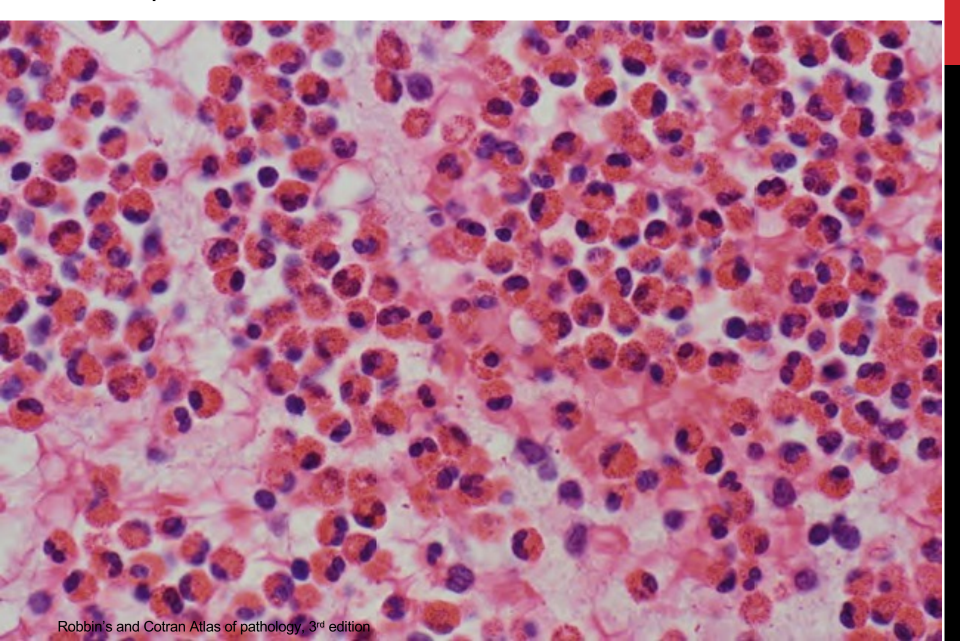
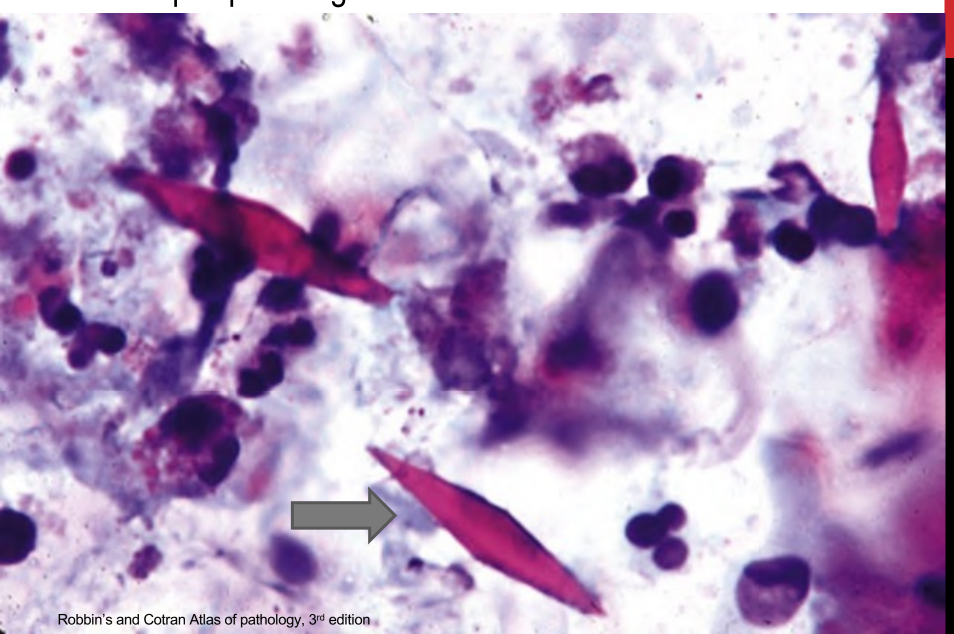


Fig. 13.11 Bronchial biopsy specimen from an asthmatic patient showing sub basement membrane fibrosis, eosinophilic inflammation, and smooth muscle hyperplasia

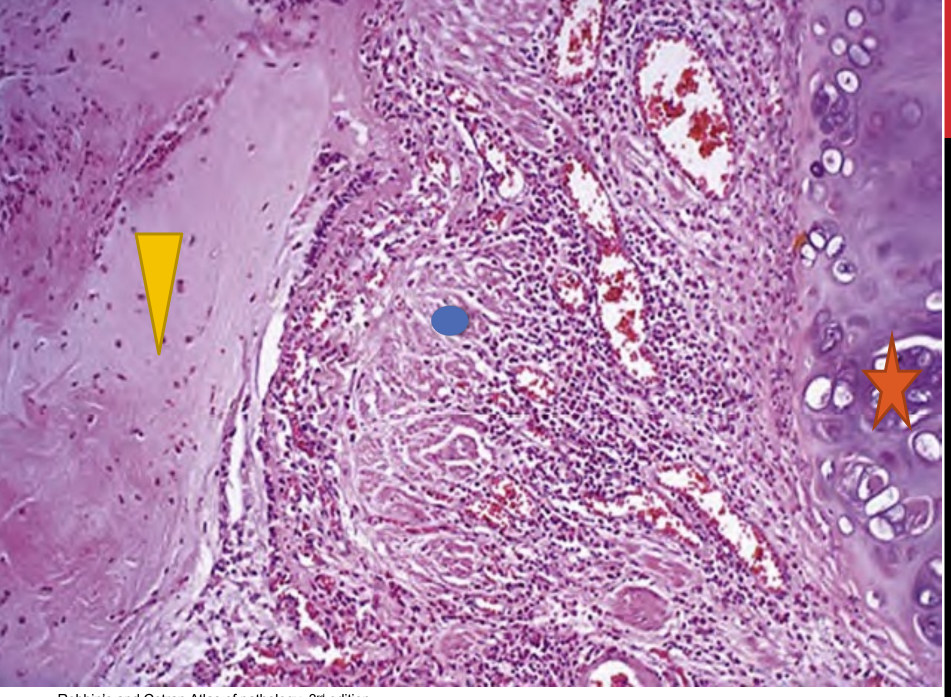
eosinophils



 Charcot-Leyden crystals: crystalloids made up of the eosinophil protein galectin-10

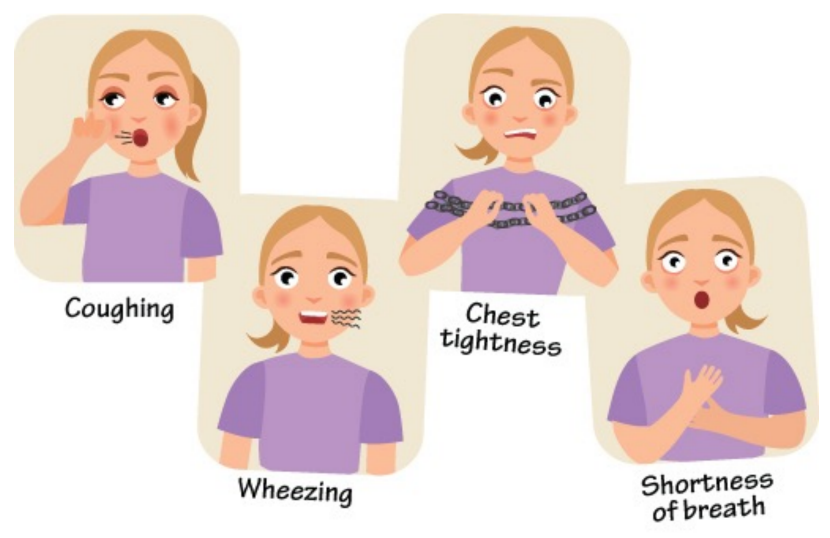


- airway remodeling, including:
 - Thickening of airway wall
 - Sub-basement membrane fibrosis
 - Increased submucosal vascularity
 - An increase in size of the submucosal glands and goblet cell metaplasia of the airway epithelium
 - Hypertrophy and/or hyperplasia of the bronchial muscle



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CLINICAL FEATURES



Status asthmaticus:

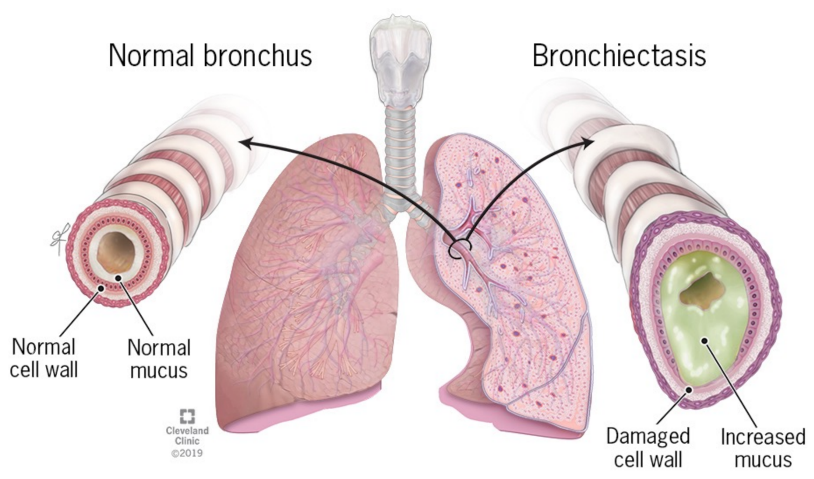
is a severe paroxysm that does not respond to therapy and persists for days or weeks. The associated hypercapnia, acidosis, and severe hypoxia may be fatal



MANAGEMENT:

- Standard therapies include:
 - Anti-inflammatory drugs(glucocorticoids)
 - Bronchodilators (beta-adrenergic drugs)
 - Leukotriene inhibitors

IV- BRONCHIECTASIS



IV- BRONCHIECTASIS

 Permanent dilation of bronchi and bronchioles caused by destruction of smooth muscle and the supporting elastic tissue

Typically results from or is associated with chronic necrotizing infections

 It is not a primary disorder, as it always occurs secondary to persistent infection or obstruction Clinically: cough and expectoration of copious amounts of purulent sputum

Diagnosis: appropriate history and radiographic demonstration of bronchial dilation.

PATHOGENESIS

- Two intertwined processes contribute to bronchiectasis:
 - **✓** obstruction
 - **✓** chronic infection

OBSTRUCTION → impairs clearance of secretions →
superimposed infection → inflammatory damage to the
bronchial wall + the accumulating exudate → airways
distention → irreversible dilation.

 PERSISTENT NECROTIZING INFECTION in the bronchi or bronchioles → poor clearance of secretions, obstruction, and inflammation with peribronchial fibrosis and bronchial walls damage → irreversible dilation

The conditions that most commonly predispose to bronchiectasis include:

Bronchial obstruction:

- By tumors, foreign bodies, and impaction of mucus OR as a complication of atopic asthma and chronic bronchitis
- bronchiectasis is localized

Congenital or hereditary conditions:

Cystic fibrosis:

- widespread severe bronchiectasis
- Due to obstruction caused by abnormally viscid mucus and secondary infections

Immunodeficiency states:

- Due to recurrent bacterial infections
- localized or diffuse

Primary ciliary dyskinesia (immotile cilia syndrome):

- rare autosomal recessive disorder → abnormalities of cilia
 → persistent infections.
- bronchiectasis + sterility in males

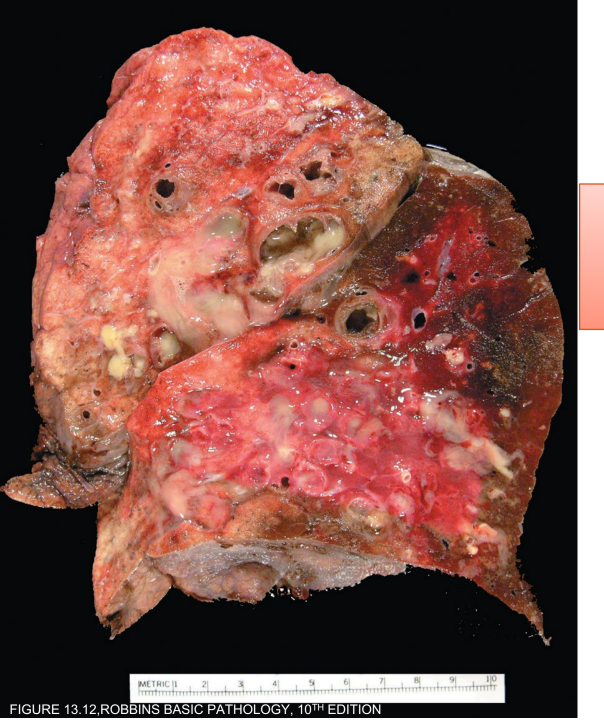
- Necrotizing, or suppurative, pneumonia:
 - particularly with virulent organisms such as Staphylococcus aureus or Klebsiella spp.

MORPHOLOGY, MACROSCOPIC:

Lower lobes bilaterally

most severe involvement in distal bronchi and bronchioles.

The airways may be dilated to as much as four times their usual diameter



markedly dilated bronchi filled with purulent mucus

MORPHOLOGY, MICROSCOPIC:

- In full-blown active cases:
 - intense acute and chronic inflammatory exudate within the walls of the bronchi and bronchioles → desquamation of lining epithelium and extensive ulceration
 - mixed flora are cultured from the sputum.

MORPHOLOGY, MICROSCOPIC:

- When healing occurs:
 - the lining epithelium may regenerate completely
 - abnormal dilation and scarring
 - fibrosis of bronchial and bronchiolar walls

- peribronchiolar fibrosis
- Abscess formation in some cases

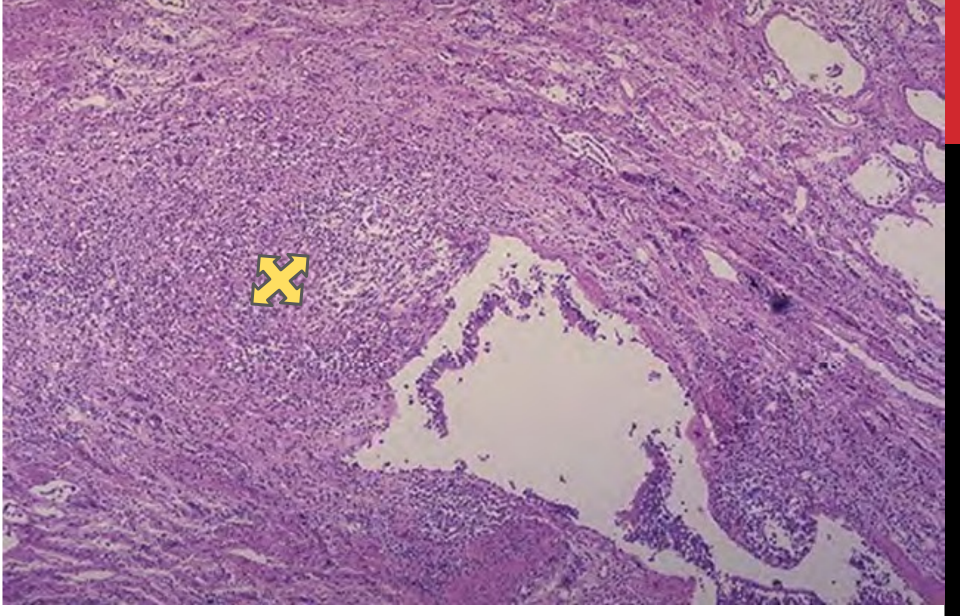


Figure 5-34 **Bronchiectasis**, **microscopic** dilated bronchus in which the mucosa and bronchial wall are not seen clearly because of the necrotizing inflammation with tissue destruction.

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CLINICAL FEATURES

- severe, persistent cough with mucopurulent sputum.
 - Other symptoms: dyspnea, rhinosinusitis, and hemoptysis.

• Symptoms are episodic

precipitated by URTI.

 Severe widespread bronchiectasis: significant obstructive ventilatory defects, hypoxemia, hypercapnia, pulmonary hypertension, and cor pulmonale.

IN SUMMARY:

Table 13.1 Disorders Associated With Airflow Obstruction: The Spectrum of Chronic Obstructive Pulmonary Disease

Clinical Entity	Anatomic Site	Major Pathologic Changes	Etiology	Signs/Symptoms
Chronic bronchitis	Bronchus	Mucous gland hypertrophy and hyperplasia, hypersecretion	Tobacco smoke, air pollutants	Cough, sputum production
Bronchiectasis	Bronchus	Airway dilation and scarring	Persistent or severe infections	Cough, purulent sputum, fever
Asthma	Bronchus	Smooth muscle hypertrophy and hyperplasia, excessive mucus, inflammation	Immunologic or undefined causes	Episodic wheezing, cough, dyspnea
Emphysema	Acinus	Air space enlargement, wall destruction	Tobacco smoke	Dyspnea
Small airway disease, bronchiolitis*	Bronchiole	Inflammatory scarring, partial obliteration of bronchioles	Tobacco smoke, air pollutants	Cough, dyspnea

^{*}Can be present in all forms of obstructive lung disease or by itself.

THANK YOU!