Lung diseases

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Restrictive lung disease
Granulomatous types
I. Sarcoidosis

- Although sarcoidosis is an example of a restrictive lung disease, it is a multisystem disease characterized by noncaseating granulomas in many tissues and organs.
Mycobacterial or fungal infections may also produce noncaseating granulomas; so the histologic diagnosis of sarcoidosis is one of exclusion.
Epidemiology

- Sarcoidosis occurs throughout the world, affecting both genders and all races and age groups.
- There is a predilection for adults younger than 40 years of age.
- Sarcoidosis is one of the few pulmonary diseases with a higher prevalence among nonsmokers.
ETIOLOGY AND PATHOGENESIS

- Although the etiology of sarcoidosis remains unknown, several lines of evidence suggest that it is a disease of disordered immune regulation in genetically predisposed persons exposed to certain environmental agents.
- Immunologic abnormalities in sarcoidosis suggest the development of a cell mediated response to an unidentified antigen and the process is driven by CD4+ helper T cells.

- These abnormalities include:
  1. Intra alveolar and interstitial accumulation of CD4+ TH1 cells
2. Increases in T cell derived TH1 cytokines such as IL2 and IFN-γ, resulting in T cell expansion and macrophage activation.

3. The bronchoalveolar lavage contains abundant CD4+ T cells

4. Anergy to common skin test antigens such as purified protein derivative (PPD)
The role of genetic factors is suggested by familial clustering and association with HLA classes such as HLA-A1 and HLA-B8.

After lung transplantation, sarcoidosis recurs in the new lungs in 75% of cases.
MORPHOLOGY
- The diagnostic histopathologic feature is noncaseating epithelioid granuloma, irrespective of the organ involved
Note: The giant cells may show

1. Schaumann bodies:
   - Are laminated concretions composed of calcium and proteins

2. Asteroid bodies
   - Are Stellate inclusions enclosed within giant cells

Note: Their presence (1&2) is not required for diagnosis of sarcoidosis. They also may occur in granulomas of other origins.
Schaumann bodies
Asteroid bodies
- Rarely, foci of central necrosis may be present in sarcoid granulomas.

- Caseation necrosis typical of tuberculosis is absent.
Involved organs:

1. The lungs are involved at some stage of the disease in 90% of patients:

- The granulomas predominantly involve the interstitium rather than air spaces, with some tendency to localize in the connective tissue around bronchioles and venules and in the pleura.
- In 5% to 15% of patients, the granulomas eventually are replaced by diffuse interstitial fibrosis, resulting in a so called honeycomb lung.

2. Intrathoracic hilar and paratracheal lymph nodes are enlarged in 75% to 90% of patients, while a third present with peripheral lymphadenopathy.
sarcoidosis
3. Skin lesions are encountered in approximately 25% of patients

A. Erythema nodosum:
- Is the hallmark of acute sarcoidosis consists of raised, red, tender nodules on the anterior aspects of the legs.
Sarcoidal granulomas are uncommon in these lesions.

b. Discrete painless subcutaneous nodules, and these usually reveal abundant noncaseating granulomas.
3. Involvement of the eye occurs in about one half of patients and the ocular involvement takes the form of iridocyclitis.

- These ocular lesions are frequently accompanied by inflammation in the lacrimal glands, with suppression of lacrimation.
4. Unilateral or bilateral parotitis with painful enlargement of the parotid glands occurs in less than 10% of patients with sarcoidosis; some develop xerostomia (dry mouth).
5- Other findings include hypercalcemia and is not related to bone destruction but is caused by increased calcium absorption secondary to production of active vitamin D by the mononuclear phagocytes in the granulomas.
Clinical Features

- In many affected persons the disease is asymptomatic, discovered on routine chest films as bilateral hilar adenopathy or as an incidental finding at autopsy.
In about two thirds of symptomatic cases.

a. gradual appearance of respiratory symptoms (shortness of breath, dry cough, or vague substernal discomfort) or

b. constitutional signs and symptoms (fever, fatigue, weight loss, anorexia, night sweats)
- Because of the variable and nondiagnostic clinical features, diagnosis is made by lung or lymph node biopsy.
- The presence of non caseating granulomas is suggestive of sarcoidosis, but other identifiable causes of granulomatous inflammation must be excluded.
Clinical course

- Sarcoidosis is characterized by either progressive chronicity or periods of activity interspersed with remissions

- 70% of affected persons recover with minimal or no residual manifestations
- 20% develop permanent lung dysfunction or visual impairment
- 10% develop pulmonary fibrosis and cor pulmonale
2. Hypersensitivity Pneumonitis

- Is an immunologically mediated inflammatory lung disease that primarily affects the alveoli and is often called allergic alveolitis.

- Most often it is an occupational disease that results from heightened sensitivity to inhaled antigens such as in moldy Hay.
- The damage occurs at the level of alveoli; so manifests as a predominantly restrictive lung disease.

- The occupational exposures are diverse, but the syndromes share common clinical and pathologic findings and probably have a very similar pathophysiologic basis.
Examples

Syndrome – Exposure– Antigens

a. Farmer's lung --------Moldy hay--Micropolyspora
b. Maple bark disease-Moldy maple bark - Cryptostroma

c. Pigeon breeder's lung-Pigeon droppings-Pigeon serum proteins in droppings
- Is an immunologically mediated disease

1. Bronchoalveolar lavage specimens consistently demonstrate increased numbers of T lymphocytes of both CD4+ and CD8+ phenotype.

2. Most patients have specific antibodies in their serum
3. Complement and immunoglobulins have been demonstrated within vessel walls by immunofluorescence, indicating type III hypersensitivity.

4. The presence of noncaseating granulomas in two thirds of patients with this disorder suggests a role for type IV hypersensitivity as well.
Morphology

- Patchy mononuclear cell infiltrates in the pulmonary interstitium mainly lymphocytes
- Interstitial non-caseating granulomas are present in about 2 thirds of th cases
- In advanced cases, diffuse interstitial fibrosis
Hypersensitivity Pneumonitis
Clinical Manifestations

- May manifest either as:

A. An acute reaction:

- Characterized by fever, cough, dyspnea, and constitutional signs and symptoms arising 4 to 8 hours after exposure.
With the acute form of this disease, the diagnosis is usually obvious because of the temporal relationship of symptom onset to exposure to the incriminating antigen.
b. or as a chronic disease
- characterized by insidious onset of cough, dyspnea, malaise, and weight loss.
If antigenic exposure is terminated after the acute attacks, complete resolution of pulmonary symptoms occurs within days.

Failure to remove the inciting agent eventually results in an irreversible chronic interstitial pulmonary disease.
Honeycomb lung
Diffuse alveolar hemorrhagic syndromes
1. Goodpasture Syndrome

- Is an uncommon condition characterized by a proliferative, usually rapidly progressive, glomerulonephritis (and) hemorrhagic interstitial pneumonitis.

- Both the renal and the pulmonary lesions are caused by antibodies targeted against the noncollagenous domain of the α3 chain of collagen IV.
These antibodies can be detected in the serum of more than 90% of persons with Goodpasture syndrome.
The characteristic linear pattern of immunoglobulin deposition (usually IgG, sometimes IgA or IgM) that is the hallmark diagnostic finding in renal biopsy specimens may be seen along the alveolar septa by immunofluorescence studies.
- Plasmapheresis and immunosuppressive therapy have markedly improved the once-dismal prognosis for this disease.
- Plasma exchange removes offending antibodies, and immunosuppressive drugs inhibit antibody production.
- With severe renal disease, renal transplantation is eventually required.
2. Idiopathic Pulmonary Hemosiderosis

- Is a rare disease of uncertain etiology that has pulmonary manifestations and histologic features similar to those of Goodpasture syndrome but

a. No associated renal disease

b. No circulating anti-basement membrane antibody.
- Most cases occur in children, although the disease is reported in adults as well, who have a better prognosis.
- With steroid and immunosuppressive therapy, survival has markedly improved from the historical 2.5 years;
- thus, an immune-mediated etiology is postulated.
Diffuse alveolar hemorrhage syndrome – perl’s stain
Lung tumors
- Primary lung cancer is also a common disease accounting for 95% of primary lung tumors.
- Carcinomas
- Carcinoma of the lung is the single most important cause of cancer-related deaths in industrialized countries.
- It accounts for about one third of cancer deaths in men, and has become the leading cause of cancer deaths in women.
- The peak incidence of lung cancer is in persons in their 50s and 60s.
- The prognosis with lung cancer is dismal:
  1. The 5-year survival rate for all stages of lung cancer combined is about 16%,
  2. disease localized to the lung, the 5-year survival rate is 45%
- The four major histologic types of carcinomas of the lung
  a. Adenocarcinoma
  b. Squamous cell carcinoma,
  c. Small cell carcinoma,
  d. and large cell carcinoma
Because of changes in smoking patterns in the U.S., adenocarcinoma has replaced squamous cell carcinoma as the most common primary lung tumor in recent years.
- Carcinomas of the lung were classified into two groups:
  
a. Small cell lung cancer (SCLC) and
  b. Non-small cell lung cancer (NSCLC), including adenocarcinomas and squamous cell carcinomas.
The reason for this historical distinction was that virtually all SCLCs have metastasized by the time of diagnosis and are not curable by surgery and are treated by chemotherapy, with or without radiation therapy.
By contrast, NSCLCs were more likely to be resectable and usually responded poorly to chemotherapy; however, now therapies are available that target specific mutated gene products present in the various subtypes of NSCLC, mainly in adenocarcinomas.

NSCLC must be classified into histologic and molecular subtypes.
ETIOLOGY

- There is strong evidence that cigarette smoking and, to a much lesser extent, other environmental insults are responsible for the genetic changes in lung cancers.

- About 90% of lung cancers occur in active smokers or those who stopped recently.
- The increased risk becomes 60 times greater among habitual heavy smokers (two packs a day for 20 years) than among nonsmokers.

- Since only 11% of heavy smokers develop lung cancer, however, other predisposing factors must be operative in the pathogenesis of this deadly disease.
- The mutagenic effect of carcinogens is conditioned by (genetic) factors.
- Many chemicals (procarcinogens) require metabolic activation via the P-450 monooxygenase enzyme system for conversion into ultimate carcinogens.
Persons with specific genetic polymorphisms involving the P-450 genes have an increased capacity to metabolize procarcinogens derived from cigarette smoke, and thus have the greatest risk for development of lung cancer.
- For reasons not entirely clear, women have a higher susceptibility to carcinogens in tobacco than men.

- Although cessation of smoking decreases the risk of developing lung cancer over time, it may never return to baseline levels.
Passive smoking increases the risk of developing lung cancer to approximately twice that of nonsmoker.

The smoking of pipes and cigars also increases the risk, but only modestly.
- There is increased incidence of lung carcinoma in asbestos workers; and workers exposed to dusts containing arsenic, chromium, uranium, nickel, and vinyl chloride
Note- Exposure to asbestos increases the risk of lung cancer fivefold in nonsmokers.

- Heavy smokers exposed to asbestos have an approximately 55 times greater risk for development of lung cancer than that for nonsmokers not exposed to asbestos.
PATHOGENESIS

- Smoking-related carcinomas of the lung arise by a stepwise accumulation of a multitude of genetic abnormalities that result in transformation of benign progenitor cells in the lung into neoplastic cells.
- The sequence of molecular changes is not random but follows a predictable sequence that parallels the histologic progression toward cancer.
Inactivation of tumor suppressor genes located on the short arm of chromosome 3 (3p) is a very early event, whereas TP53 mutations or activation of the KRAS
2. In Adenocarcinomas
   a. Activating mutations of the epidermal growth factor receptor (EGFR) and these tumors are sensitive to agents that inhibit EGFR signaling, but the response often is short-lived.
b. MET tyrosine kinase gene amplifications
c. In 4% of adenocarcinomas are EML4-ALK tyrosine kinase fusion genes and
4. ALK tyrosine kinase fusion genes and c-
- These abnormalities, while rare, are important because of their therapeutic implications, as they can be targeted with tyrosine kinase inhibitors.
The identification of genetic alterations producing overactive EGFR, ALK, and MET has opened up a new era of "personalized" lung cancer therapy, in which the genetics of the tumor guides the selection of drugs.
Among the major histologic subtypes of lung cancer, squamous and small-cell carcinomas show the strongest association with tobacco exposure.
MORPHOLOGY

1. Squamous cell carcinomas:
   a. are more common in men than in women
   b. are closely correlated with a smoking history;
c. They tend to arise centrally in major bronchi and eventually spread to local hilar nodes,

d. Disseminate outside the thorax later than do other histologic types
e. Large lesions may undergo central necrosis, giving rise to cavitation.

f. Are preceded by the development, over years, of squamous metaplasia or dysplasia in the bronchial epithelium,
h. Eventually, the small neoplasm reaches a symptomatic stage, when mass begins to obstruct the lumen of a major bronchus, often producing distal atelectasis and infection.
Squamous cell carcinoma of lung
2. Adenocarcinomas:
   a. May occur as central lesions but usually are more peripherally located, many with a central scar.
   b. Are the most common type of lung cancer in women and nonsmokers.
c. In general, adenocarcinomas grow slowly and form smaller masses than do the other subtypes.

d. They tend to metastasize widely at an early stage.

- On histologic examination, they may assume a variety of forms, including:
a. acinar(gland-forming),
b. and papillary,
b. Mucinous which often is multifocal and may manifest as pneumonia like consolidation)
d. and solid types. requires demonstration of intracellular mucin production by special stains to establish its adenocarcinomatous lineage
Note: Although foci of squamous dysplasia may be present in the epithelium proximal to resected adenocarcinomas, these are not the precursor lesions for this tumor.
- The putative precursor of peripheral adenocarcinomas is atypical adenomatous hyperplasia which progresses to:
  a. adenocarcinoma in situ (formerly bronchioloalveolar carcinoma),
  b. Minimally invasive adenocarcinoma (tumor less than 3 cm and invasive component measuring 5 mm or less),
  c. and invasive adenocarcinoma (tumor of any size that has invaded to depths greater than 5 mm).
Adenocarcinoma in situ (AIS), formerly called bronchioloalveolar carcinoma, often involves peripheral parts of the lung, as a single nodule.

- The key features of AIS are:
  a. Diameter of 3 cm or less,
  b. Growth along preexisting structures,
  c. and preservation of alveolar architecture
d. The tumor cells, which may be nonmucinous, mucinous or mixed, grow in a monolayer along the alveolar septa, which serve as a scaffold (this has been termed a “lepidic
growth pattern,
e. By definition, AIS does not demonstrate destruction of alveolar architecture or stromal invasion with desmoplasia, features that would merit the diagnosis of frank adenocarcinoma
3. Small cell lung carcinomas (SCLCs) are:
   a. Centrally located with extension into the lung parenchyma
   b. Early involvement of the hilar and mediastinal nodes.
   c. Are composed of tumor cells:
      1. with a round to fusiform shape, scant cytoplasm, and finely granular chromatin.
2. Mitotic figures frequently are seen
3. Necrosis is invariably present and may be extensive
5. Fragile cells that show fragmentation and "crush artifact".
6. Nuclear molding resulting from close apposition of tumor cells that have scant cytoplasm.
Small cell carcinoma of the lung
1. For all of these neoplasms, it is possible to trace involvement of successive chains of nodes in carina, in the mediastinum, and in the neck (scalene nodes) and clavicular regions and then distant metastases. 

- Involvement of the left supraclavicular node (Virchow node) is particularly characteristic and sometimes calls attention to an occult primary tumor.
3. They may infiltrate the superior vena cava to cause either venous congestion or the vena caval syndrome
3. Apical neoplasms (Pancoast tumors) causes Pancoast syndrome characterized by:
   a. Invasion of the brachial or cervical sympathetic plexus to cause severe pain in the distribution of the ulnar nerve resulting in Horner syndrome (ipsilateral enophthalmos, ptosis, miosis, and anhidrosis).
b. Is accompanied by destruction of the first and second ribs and sometimes thoracic vertebrae
Clinical Course

- Are silent, cancers that in many cases have spread so as to be unresectable before they produce symptoms.

- In some instances, chronic cough and expectoration call attention to still localized, resectable disease.
- By the time hoarseness, chest pain, superior vena cava syndrome, pleural effusion, or segmental atelectasis or pneumonitis makes its appearance, the prognosis is grim
- Too often, the tumor presents with symptoms resulting from metastatic spread to the brain (mental or neurologic changes), liver (hepatomegaly), or bones (pain).

- Although the adrenals may be nearly obliterated by metastatic disease, adrenal insufficiency (Addison disease) is uncommon,
- It is estimated that 3% to 10% of all patients with lung cancer develop clinically overt paraneoplastic syndromes.

1. Hypercalcemia: caused by secretion of a parathyroid hormone-related peptide by squamous cell carcinoma
- Osteolytic lesions may also cause hypercalcemia, but this would not be a paraneoplastic syndrome

2. Cushing syndrome (production of Adrenocorticotropic hormone); by small cell carcinoma
(3) syndrome of inappropriate secretion of antidiuretic hormone; by small cell carcinoma

(4) neuromuscular syndromes, including a myasthenic syndrome, peripheral neuropathy, and polymyositis

5) clubbing of the fingers and hypertrophic pulmonary osteoarthropathy by any type of carcinoma
6) coagulation abnormalities, including migratory thrombophlebitis, nonbacterial endocarditis, adenocarcinoma