Medical schools and residency training programs rarely talk about smoking. In this lecture, we will examine the habit, the puff and the respiratory consequences of smoking.
Objectives

What are the components of the puff
What are the four pathologic responses to the puff
How does physiology affect the distribution of smoking related diseases

At the end of this lecture, you should be able to answer the following objectives.
Smoke…Smoke…Smoke that cigarette

- Puff duration 2 seconds
- 10 puffs/cigarette
- Total duration 3-12 minutes/cigarette
- 1pk/day for 40 yrs = 2,920,000 puffs

The unit dose in cigarette smoking is the puff. A typical smoker accumulates nearly 3 million puffs.
The puff contains gases and particulate material called tar. Inhaled particles are heterogeneous in size. Smaller particles deposit primarily within respiratory bronchioles. Total particulate dose is large.

Puff

50 ml diluted in tidal volume 1000 ml
Mean aerodynamic diam 0.46 mm
Particulate concentration $10^{11}$
50% deposited (9:1 lobule to airways)
1 pk/day 20 mg “tar”/cig = 6000 gm/40 yrs
300,000,000,000,000,000,000 particles
Cigarettes contain a wide variety of materials besides tobacco leaf. From the Brown & Williamson files, foreign bodies included insect parts, animal hairs, and fungal debris.
Tobacco burns at approximately 900 degrees centigrade. This combustion produces a wide variety of organic compounds, not all of which have been studied. The puff is mostly gas, including carbon monoxide, and particulate material called tar.
Components

- Plutonium, arsenic, lead, cadmium
- Hydrogen cyanide
- Formaldehyde
- Ammonia
- Methane
- Toluene
- Methanol
- Acetaldehyde
- Acetone

These well known poisons are all known constituents of the puff.
What happens to particles?

Let’s now turn our attention to what happens to the puffs constituents when inhaled.
What happens to particles?

<table>
<thead>
<tr>
<th>Bronchus</th>
<th>Central</th>
<th>Peripheral</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bifurcations</td>
<td>Respiratory</td>
<td></td>
</tr>
<tr>
<td>Volume</td>
<td>150 ml</td>
<td>4500 ml</td>
</tr>
<tr>
<td>Flow</td>
<td>High</td>
<td>None</td>
</tr>
</tbody>
</table>

Particles will deposit somewhere along the airways. Large particles deposit on the ciliated airways from impaction. Small particles less than 5 microns in size will suspend in air currents until they reach the respiratory bronchioles. At this point airflow markedly decreases and particles are deposited.
Central Airways

High turbulent flow
Deposition studied
Airway casts
Ferric oxide microspheres
(1.7 - 12.2 mm)

Central airway anatomy is complex. Airflow through these tubes produces complex turbulence. Particle deposition is generally studied using airway casts with varying size particles.
Such studies show that most particles are deposited in the upper lobe airways at bifurcation points. Note the concordance between this experimental data and the historical frequency of the origin of central lung cancers.
Studying particulate deposition in the peripheral airways is more difficult. This typical smoking apparatus adds a nebulized radionuclide particle to the puff.
Small particles are deposited in the bases. This conforms to pulmonary physiology where the lower lobes are better ventilated. Compared to nonsmoking controls, smoking skews the deposition towards a more uniform deposition, however, more particles accumulate in the lower lobes.
Deposition is only one aspect of how the lung handles particles. Once deposited particles are gradually removed. The chronic retention of particulate material can be determined by autopsy examination of different elements. Even though the analyses involve different methods, the results are remarkably similar. Particles tend to accumulate over time in the upper lobes, approximately 1.25 times that of the lower lobes.
Note that the respiratory bronchiole and alveolar unit are devoid of lymphatics. The lung lymphatics are composed of two systems, one along the bronchovascular bundle and the second peripheral system along the pulmonary veins. What happens to particles that reach the respiratory bronchioles? Well they can be absorbed or engulfed by macrophages. Some are trapped forever. Those particles which can be engulfed by macrophages are removed through the lymphatics (or migrate to the terminal bronchiole and escape via the cilia). As we’ve seen, removal of particles is slow. Lymphatic flow is, in part, determined by arterial pressure and respiratory motion, which aids lymphatic flow.
Arterial pressure is nonuniform, the low pressure pulmonary artery system is linearly affected by gravity. Normal pulmonary pressure is just sufficient to perfuse the apex of the lung, thus the driving pressure for lymphatic flow will diminish towards the lung apex. In addition, passive respiratory motion is nonuniform, the lung bases and the anterolateral portion of the lung undergoes greater degree of respiratory expansion than the apices and dorsal aspect of the lung.
If gravity is important in the distribution of dusts and particles in the lung then the anatomical analysis used in these investigations is the incorrect method. For example, in a sagital section of the lung, the lower portion of the upper lobe and the upper portion of the lower lobe occupy the same gravitational plane, thus an analysis based on traditional anatomic boundaries will blur the true distribution of particulate material in the lung. The correct analysis will bread loaf the lung, much as we do with CT. Note that in this lung, pigmentation gradually diminish toward the base. Note the degree of pigmentation in the superior segment of the lower lobe is the same as the upper lobe.
Thus, even though the lung bases acutely accumulate the majority of inhaled particles, the lung bases are also cleared more effectively and particles accumulate in the upper lung zones.

Chronically, particles tend to accumulate in the upper lung zones. This is true of most of the dust diseases, from silica and coal, and as will we see, the smoking related diseases.
Harmful substances are also contained in the gas. Let’s quickly examine how the lung handles this gaseous component.
The concentration of gas in the alveolus depends on the linear relationship between blood flow and ventilation. Both are affected by gravity, higher at the base in the upright lung. Because the rate of change between blood flow and ventilation differs in the upright lung, the V/Q ratio is highest in the upper lung zones.
This diagram illustrates the extreme, no ventilation or no perfusion. As the V/Q ratio increases the concentration of inhaled gas also increases.
We will examine four diseases related to the harmful products in the puff.
Respiratory bronchiolitis was first described in young victims of motor vehicle accidents. This pathologic condition develops within 2 years of the first puff. The majority of patients are asymptomatic and have normal pulmonary function. This significance of this lesion is unknown. The condition is not specific to cigarette smoke but is seen with other dust pollutants.
Pathology

Pigmented macrophages
Centered on respiratory bronchioles

The hallmark is the pulmonary macrophage. This one is stuffed with particulate yellowish-brown material.
Clustered dirty macrophages associated with epithelial hyperplasia that extends into adjacent alveolar walls in a stellate fashion

In this example, macrophages are clustered in the wall of the respiratory bronchiole, the adjacent arteriole is on the left
Chest x-rays are typically normal. Even though all smokers have respiratory bronchiolitis, even HRCT is insensitive for this condition.
The nodules have a centriacinar distribution. A perilymphatic distribution would demonstrate beading of the vessels, involvement of the subplueral lung, fissures, and septa.
Smoking related diseases

- Respiratory bronchiolitis
- Centriacinar emphysema
- Bronchogenic carcinoma
- Eosinophilic granuloma

Let's now turn to centriacinar emphysema and compare and contrast this disease with respiratory bronchiolitis.
Centriacinar emphysema

Long term smokers
Dose and time dependent
100% ( > 40 pk/yrs)
Average age @ diagnosis 64
Usually asymptomatic < 30% destruction

This common disorder is dose and time dependent. A large portion of the lung must be destroyed before symptoms occur. Most patients with emphysema are asymptomatic.
Centriacinar emphysema is the dilatation and destruction of the respiratory bronchiole. In this pathologic example of a portion of the secondary pulmonary lobule, septa are visible along the top and left side of the illustration. The smaller holes are normal alveolar sacs.
Again this is a disease of the upper lung zones. Note the vertical distribution equally involving the upper and lower lobes at any given level.
Neutrophils

Proteolytic reaction to activated neutrophils which release elastases and oxygen radicals

Animal models use trypsin to produce similar lesions. Neutrophils are considered the culprit because these cells are rich in elastases and proteases.
Chest x-rays are insensitive for emphysema. HRCT is the most sensitive radiographic exam. Note the holes have no discernable walls.
Respiratory bronchiolitis and centriacinar emphysema were juxtaposed for this reason. Are they related? One occurs early and the other late in the natural history of smoking. The location and distribution of pathology are identical in each. Respiratory bronchiolitis is composed of inflammatory cells. Confirmation of this hypothesis would require a longitudinal study of smokers with HRCT.
Longitudinal study

Mean interval 5.5 yrs (4.5 - 7.5 yrs)
19 with initial micronodules
  7 no change
  7 increase profusion
  5 evolved into emphysema

There is now good evidence that indeed respiratory bronchiolitis is the precursor of emphysema. In a longitudinal study of smokers, micronodules (in the upper lung zones) evolved into emphysema.
Smoking related diseases

- Respiratory bronchiolitis
- Centriacinar emphysema
- Bronchogenic carcinoma
- Eosinophilic granuloma

Bronchogenic carcinoma is the leading cause of cancer death in both men and women. 90% are related to smoking.
Bronchogenic carcinoma
Peak incidence 50 - 70
Rare in nonsmokers
Risk: dose related
40 carcinogens in the puff

Over 40 carcinogens are identified in the puff. The latent period between the onset of smoking and the development of lung cancer is generally 30 - 50 years.
Through the 1980’s, squamous cell carcinoma was the most common histologic type. Primarily involving the central lobar and segmental airways. Since then, adenocarcinoma, usually a peripheral SPN has become the most common subtype. Why the change, In response to the health effects of smoking, tobacco companies introduced the filtered (‘safer’) cigarette in the early 1950’s. By the mid 50’s, the filtered cigarette had become the most popular form of tobacco consumption. The filter only removed the largest particles, those that because of their size impacted on the central airways, leaving the small particles to be deposited in the smaller airways in the lung periphery.
The radiographic manifestations are many. In general, they can be divided into peripheral tumors which present as SPN’s or central tumors which obstruct the airways.
Similar to the other smoking related diseases. The distribution primarily affects the upper lobes, particularly the right.
The predilection for the upper lobes, especially the right upper lobe is most pronounced in the young individual. Note that the gap between the upper and lower lobes gradually narrows with age.
Eosinophilic granuloma is an interesting disorder now thought to be related to smoking related.
Over 95% of patients with EG smoke. This disease usually occurs in the younger - heavier smokers. Curiously 25% of cases are asymptomatic, discovered on chest radiographs obtained for other reasons. This high percentage suggests that the disease is more common than appreciated. All smokers have increased numbers of Langerhan’s cells in the lung even though they don’t have EG.
The Langerhans cell normally resides in the lung, skin, and reticuloendothelial system. It is an antigen processing cell. In this granuloma the Langerhans cell stains black with S-100 protein.
The granulomas of EG have tentacles, similar to a starfish.
EG may be an allergic reaction to some constituent of the puff. Evidence includes, the role of the Langerhans cell in processing antigen and the granulomatous response similar to other allergic pathologies.
The allergen is unknown. One pathology report links the Birbeck granule, shown here, with a component of the soil that the tobacco plant grows in.
The disease primarily involves the mid and upper lung zones. The pattern is that of nodules or honeycombing. In contrast to other interstitial lung diseases, the lung volumes are normal or increased. Many patients present with spontaneous pneumothorax, as did this patient.
The nodules in EG are centriacinar in distribution. As they get larger they may have tentacles.
Centriacinar nodules are thought to evolve into thin-walled cysts. These examples are from different patients.
Cysts may aggregate and form shapes which are bizarre, this is nearly pathognomonic of EG. Note the chest tube anteriorly.
The puff produces a wide variety of responses in the lung. The distribution is similar and reflects how the lung handles gas and particulate material.
Thanks for stopping by.

Credits
Smoke! Smoke! Smoke! (That cigarette)
Commander Cody & His Lost Planet Airman, Warner Bros 1976
Smoke! Smoke! Smoke! (That cigarette)
Tex Williams & His Western Caravan Capital 1996