

Ch. 11: Respiratory Physiology

PowerPoint updated 3/2025

Objectives:

1. Review respiratory anatomy.
2. Understand mechanics of breathing, gas pressure vocabulary, and the principles of surface tension, compliance, and recoil.
3. Respiratory disorders and diagnosing them
4. How gas exchange occurs between the alveoli & pulmonary vessels, and between capillaries & tissue.
5. Regulation of breathing (voluntary vs involuntary) and blood pH
6. Hemoglobin & hemoglobin disorders

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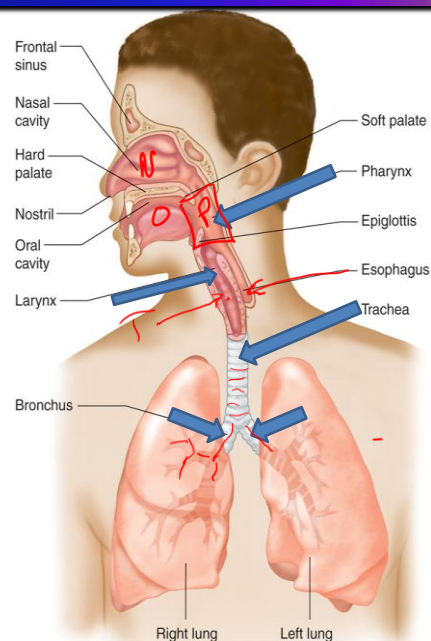
1. Respiratory Anatomy - REVIEW

2 Zones of Respiratory System:

Conduction and Respiratory zones

1) Conduction Zone = from oral/nasal cavities to:

Pharynx (throat)
 Larynx (voice box)
 Trachea
 Primary Bronchi
 Secondary Bronchi
 Tertiary Bronchi
 Terminal bronchioles

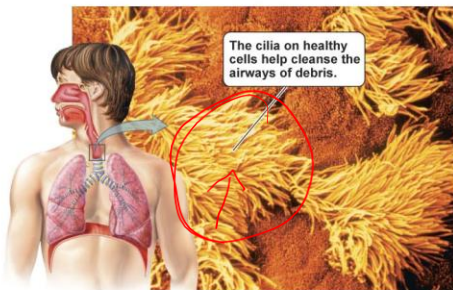


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Functions of Conducting Zone:

- Transports air to the lungs.
- Warms, humidifies, filters, and cleans the air.
 - **Mucus** traps small particles, and **cilia** move it away from the lungs.
 - **Expectoration** = coughing up the mucus that trapped debris and pathogens.

Do NOT Memorize



(a) The cilia are yellow in this color-enhanced electron micrograph. The cells without cilia secrete mucus.



(b) Cigarette smoke first paralyzes and then destroys the cilia. As a result, hazardous materials can accumulate on the surfaces of the air passageways.

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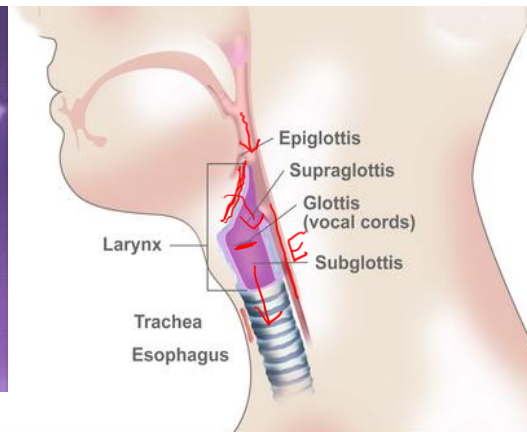
Structures in the larynx

glottis – opening between vocal cords

epiglottis – closes upon swallowing to prevent food from entering



Click [HERE](#) for GIF

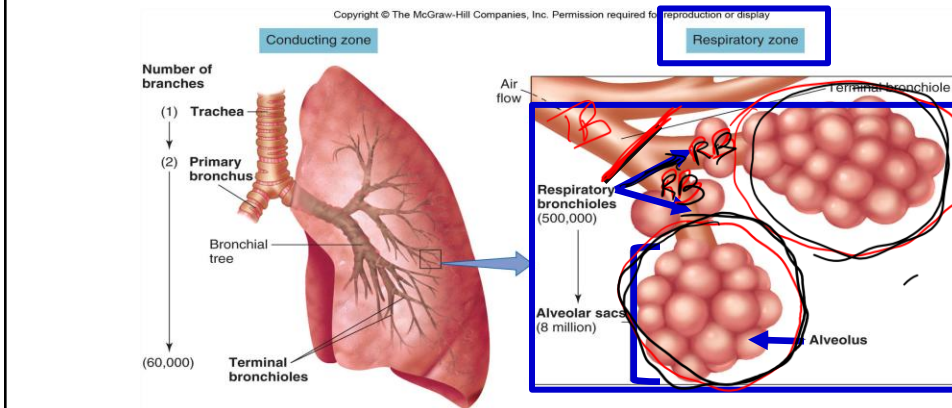


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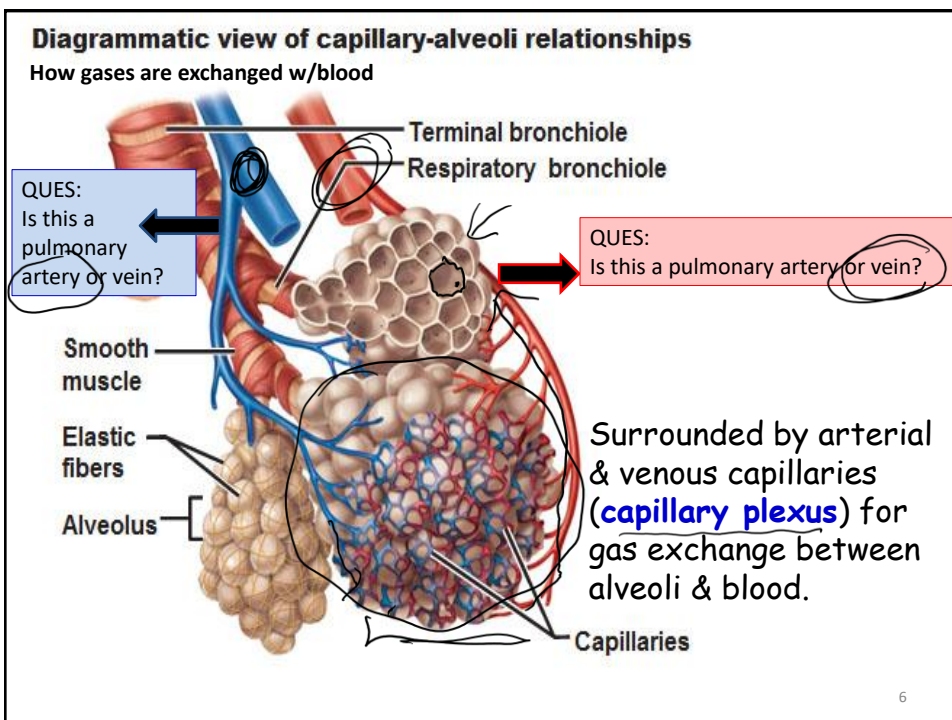
2) Respiratory zone

Respiratory bronchioles = smallest bronchioles, branch from tertiary bronchioles.

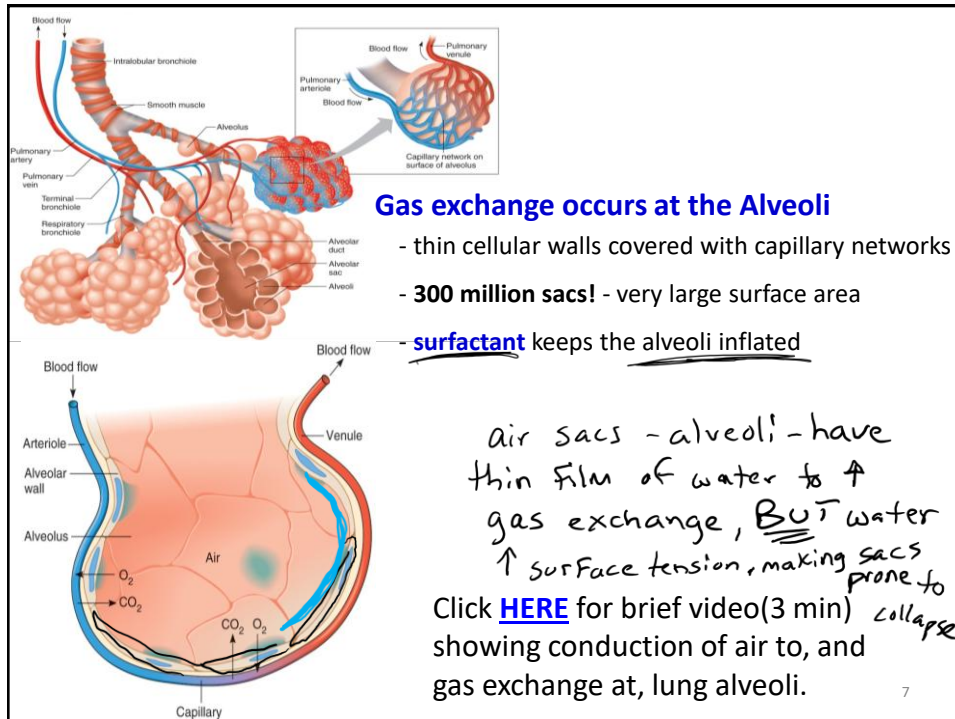
Alveolar sacs = honey-comb shaped, 1-cell thick sacs for gas exchange. [~600 mill in lungs!]



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2 Types Alveolar Cells:

Type 1 Alveolar Cells = make up alveolar walls

It's 97% of total lung surface area where most gas exchange occurs.

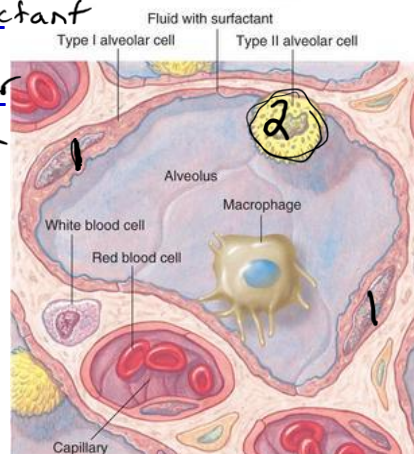
Type 2 Alveolar Cells = make surfactant

to ↓ surface tension from water

to prevent alveolar collapse

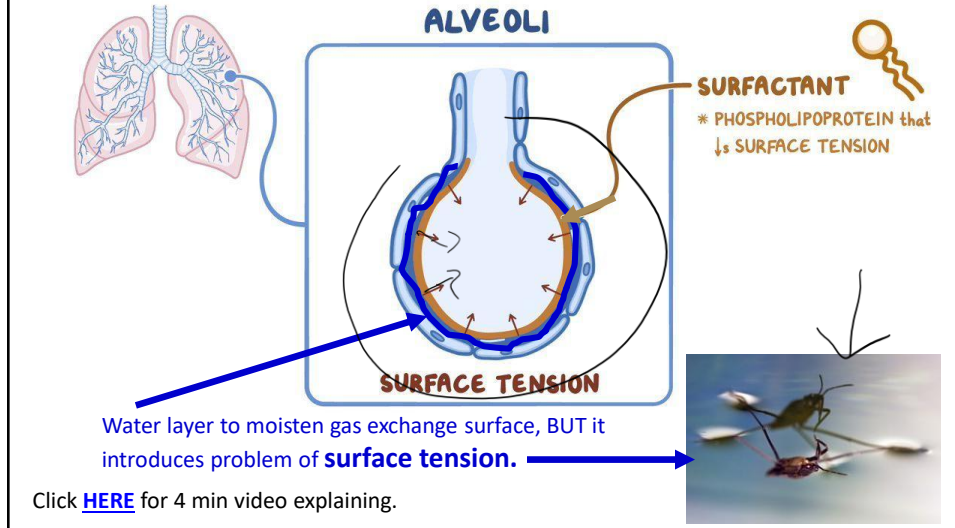
Surfactant

So when you exhale and alveolar sac shrinks, the walls don't stick together, causing collapsed alveoli (collapsed lungs!).



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Effect of surface tension and surfactant in alveolar sacs.



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Why is surfactant important?

no blockage
Non-obstructive Atelectasis =
collapsed lung (collapsed
alveolar sacs) *due to ↓ surfactant*

Leads to collapsed lungs.

LUNG COMPLICATIONS Respiratory distress syndrome

Normal alveoli Collapsed alveoli

Atelectasis

Not to be confused with

Obstructive Atelectasis =
lung collapse (alveolar
collapse) from obstruction.

Leads to collapsed lungs.

Normal Bronchiole

Blocked Bronchiole

Area of collapsed lung

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Surfactants ↓ intra-alveolar pressure & prevent collapse

Infant Respiratory Distress Syndrome (IRDS)



- Surfactant is produced > 28 weeks (7-8 months)
- Babies are born < 28 wks - not enough surfactant. High surface tension inside alveoli, results in collapsed alveoli, which collapses lung (non-obstructive atelectasis)
- Tx = synthetic surfactant delivered into baby's lungs & mechanical ventilator until Type 2 alveolar cells can make surfactant.

Acute Respiratory Distress Syndrome (ARDS)

- Due to inflammation from infection (septic shock) *lots of things*
- Results in protein (serum) secretion in lungs.
- Fluid dilutes surfactant, ↑ surface tension, alveoli collapse,
- could cause lung collapse (non-obstructive atelectasis)

COVID causes ARDS!
Click [HERE](#) to read more.

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Coronavirus (COVID) and ARDS

- Alveolar sac walls are very thin to allow for easy gas exchange.
- **Chronic inflammation of any kind**, which can occur with COVID-19, **leads to thickening of the alveolar walls**, making gas exchange difficult. (A patient with ARDS will have low oxygen levels – lower than the normal of 95% oxygen saturation of their arterial blood.)
- The **inflammation also causes serum buildup within the alveolar sacs**, which further decreases gas exchange.
lead to scar tissue. Permanent damage
- The increased fluid within the alveoli of the lungs is prone to **bacterial infection**. This is called **pneumonia**.

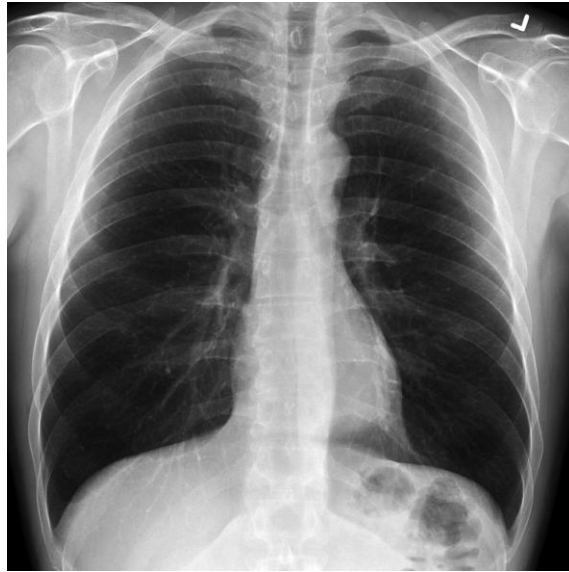
Click [HERE](#) for YouTube video explaining this, and respiratory treatment discoveries that decrease mortality of patients put on ventilators.

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Radiograph of healthy lungs

The black areas show air spaces. A normal x-ray of the lungs looks like this. There should be no white spots.



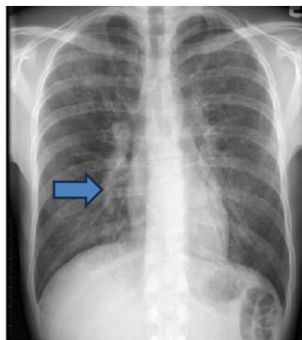
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Radiograph of lungs with **pneumonia & COVID-19**



(a) Normal



(b) Pneumonia

The arrow points to white spots, which are fluid pockets within the alveolar sacs that have become infected with bacteria.



(c) COVID-19

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Thoracic cavity: Anatomy REVIEW!

Membranes of the lungs:

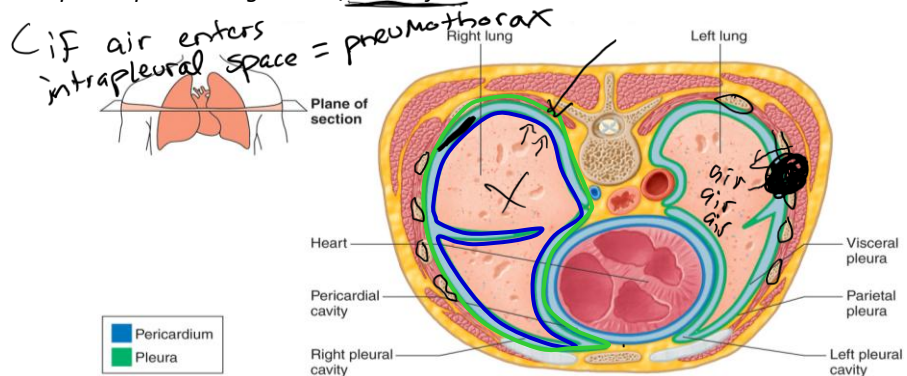
Visceral pleura = membrane covering the lungs. *Viscera = organ*

Parietal pleura = membrane lining the pleural cavity containing each lung.

- Parietal pleura held tight against thoracic wall by surface tension of water layer.
- As thoracic cage changes volume (w/ breathing) so do the lungs.

Intrapleural space = empty space between the 2 pleura.

- The 2 pleura pressed together w/serous fluid between them.

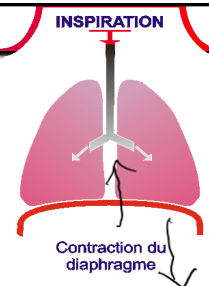


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2. Mechanics of Respiration

1) Air moves across membranes from high to low pressure

- Atmospheric air pressure = constant (760 mmHg)
- Lung air pressure depends on volume of thoracic cavity



2) Air pressure in lungs (closed chamber) changes with volume of chamber

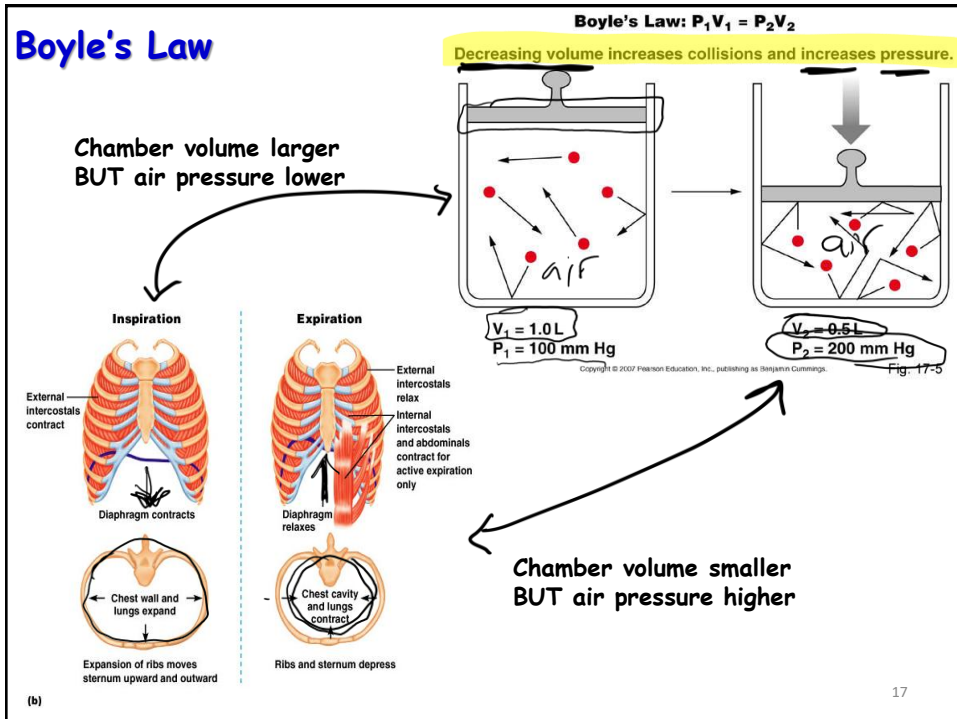
"Boyle's Law" = as volume of closed chamber \uparrow , air pressure within \downarrow
 as volume of closed chamber \downarrow , air pressure within \uparrow

within lungs
Translates to lung volume & air pressure within lungs ("intrapulmonary pressure")

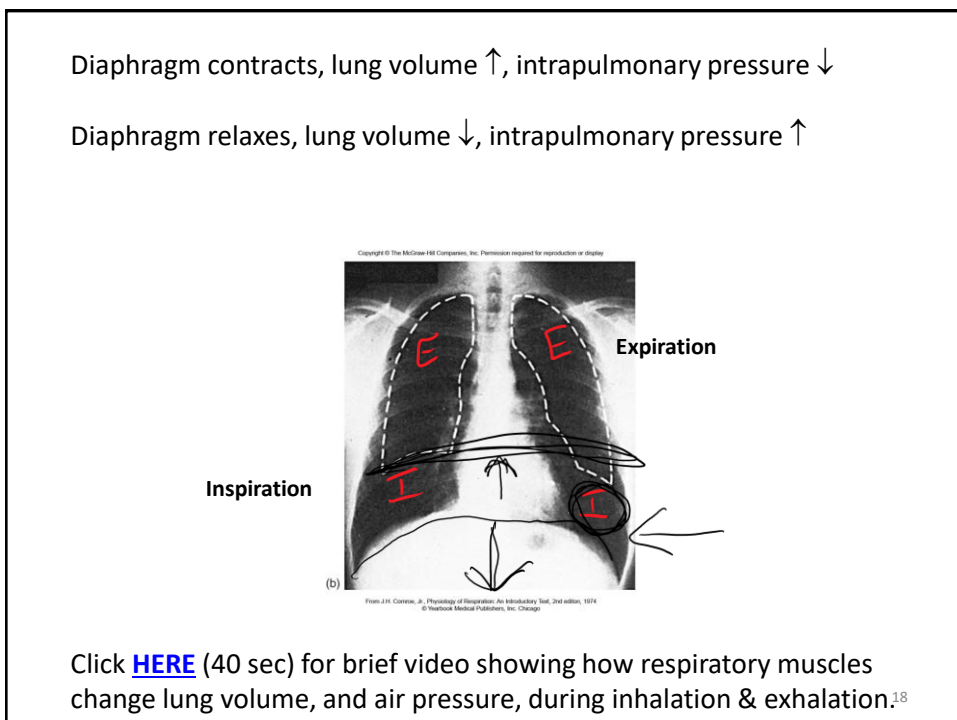
When diaphragm contracts, thoracic volume \uparrow , lung volume \uparrow , & intrapulmonary pressure \downarrow

When diaphragm relax, thoracic volume \downarrow , lung volume \downarrow , & intrapulmonary pressure \uparrow

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Gas Pressure Vocabulary:

Intrapulmonary pressure = pressure within lungs. Directly influenced by size of thoracic cage, depends on diaphragm.

- During inhalation – is lower than atmospheric pressure (-3 mmHg) below atmosphere
- During exhalation – is above atmospheric pressure ($+3 \text{ mmHg}$) above atmosphere

Intrapleural pressure = pressure between visceral & parietal pleura

- During inhalation – is lower than atmospheric (-6 mmHg)
- During exhalation – is still lower atmospheric (-3 mmHg)

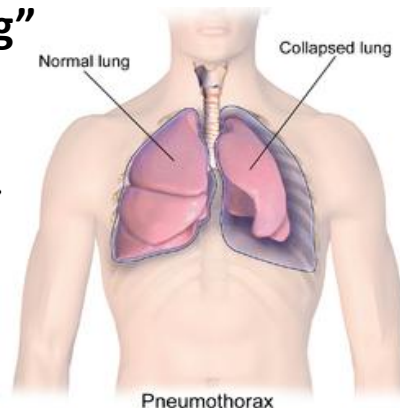
***** intrapleural pressure should ALWAYS be negative. If air enters this space, the lung can detach from thoracic wall, trapped air puts pressure on lung, & lung can collapse. ****

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Causes of a collapsed lung"

"Pneumothorax" = air enters the intrapleural space.



Result = can't expand lungs to get air to enter! Lung collapses.

Click [HERE](#) for 3 min YouTube video of pneumothorax.

Treatment = chest tube. Click [HERE](#)

2nd pneumothorax YouTube video where doctors

demonstrate with a set of lungs wrapped in a bag to simulate air invading the space between visceral parietal pleura. (As air escapes punctured lung, it fills the space making the bag expand and the lung collapse

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CLINICAL APPLICATIONS

If air enters the intrapleural space and thereby raises the intrapleural pressure, the difference in pressure between the inside of the lungs (intrapulmonary pressure) and the outside of the lungs (intrapleural pressure) is abolished. As a result, the lung is no longer stuck to the thoracic wall; this is like releasing a stretched rubber band, and the lung's elastic recoil causes it to collapse. The condition of air entering the intrapleural space and causing the collapse of a lung is known as a **pneumothorax**. Fortunately, a pneumothorax usually causes only one lung to collapse, because each lung is contained in a separate pleural compartment.

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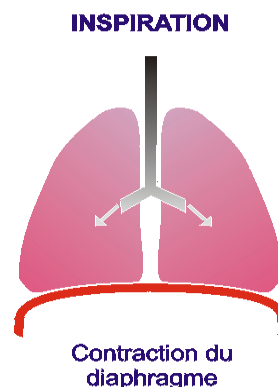
Important properties of the lungs:

A) Surface tension = pressure resulting from thin film of water lining alveoli that resists their expansion. Makes alveoli want to collapse with exhalation.

B) Compliance = lungs expand when stretched (when thoracic volume \uparrow).

- more lung compliance = greater capacity for "stretchiness"
- less lung compliance = less capacity for "stretchiness"

C) Elasticity/Recoil = tendency of lungs to return to normal shape after stretching. *(I use the word recoil, because it avoids confusion with the "stretch" of compliance.)* (When thoracic volume \downarrow , lungs volume also \downarrow parietal pleura keeps lungs "stuck" to thoracic wall).



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B) Lung compliance

Factors that increase compliance:

- pulmonary surfactants

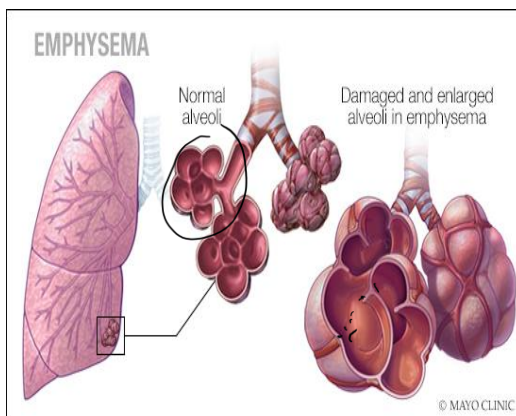
- **Emphysema** = air-trapping disorder from chronic inflammation (due to smoking) that destroys alveolar walls.

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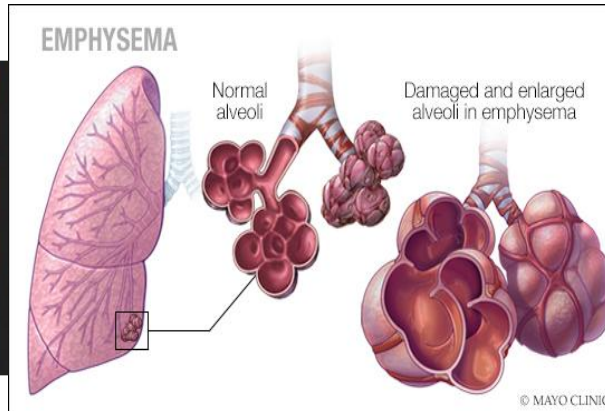
Smoking, Emphysema, and increased lung compliance

Smoking causes particles to settle into alveoli (where they never leave & cause chronic inflammation). The inflammation damages & destroys alveolar walls, leading to large air spaces within alveoli (thus **emphysema** is called an **air trapping disease**).



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In a [study](#) published in the *American Journal of Respiratory and Critical Care Medicine*, the UNC scientists found that the lungs of **vapes** – like the lungs of smokers – have elevated levels of protease enzymes, a condition known to cause emphysema in smokers. The researchers also found that the nicotine in vaping liquids is responsible for the increase in protease enzymes.



<https://808novape.org/scientists-show-how-vaping-induces-reactions-in-lungs-that-can-lead-to-disease/>

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For Ex. - Lung damage from smoking causes chronic bronchitis and formation of scar tissue ([pulmonary fibrosis](#)). *Click image for YouTube video demonstrating scarring of the lungs from smoking. (This will freak you out!)*



Comparison between normal lungs with smoker's lung

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B) Lung compliance

Factors that decrease compliance:

- many, many things!
- Anything that causes chronic inflammation can lead to ↓ compliance

Example: pulmonary fibrosis. = lung scar tissue

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Review

- The respiratory system
 - The conduction & respiration zones
- Airway, lung, and thoracic cavity anatomy
- Alveoli (gas exchange, surfactant, factors that affect intra-alveolar surface tension and pressure)
- Mechanics of breathing (Boyle's law and respiratory muscles), muscles of respiration.
- Gas pressure vocabulary, and pneumothorax,
- Important properties of the lungs
 - Surface tension, compliance, & recoil.
 - Factors that affect compliance: emphysema & pulmonary fibrosis

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3. Respiratory Disorders

Restrictive Disorder = Lung tissue is damaged. Lungs are stiff, or respiratory muscles are weak. Have difficulty filling lungs with air.

Example: **Pulmonary fibrosis** (also decreases compliance)

• Pulmonary sarcoidosis

~~Lung cancer~~

IRDS and ARDS

Obstructive Disorder = Lung tissue is normal, for time being, but resistance is increased (airways are narrowed) so air tends to remain in lungs even after exhaling.

Examples: **Asthma**, **COPD**, **Emphysema**, **Cystic fibrosis**,
Chronic bronchitis, and sleep apnea

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Restrictive Respiratory Disorders

- **Pulmonary fibrosis** = _____

HAS MANY CAUSES

> Breathing in small particles that accumulate in & irritates the lungs:

Ex: **Silicosis** = _____ ^{↳ of glass, sand, stone, dirt,}

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Restrictive Respiratory Disorders

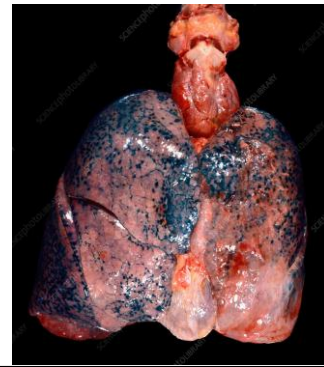
- Pulmonary fibrosis = _____

HAS MANY CAUSES

> Breathing in small particles that accumulate in & irritates the lungs:

Ex: **Anthracosis** (black lung disease) =

↳ coal dust
coal miner's disease



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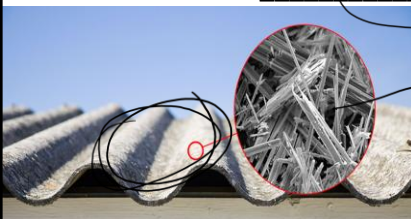
Restrictive Respiratory Disorders

- Pulmonary fibrosis = _____

HAS MANY CAUSES

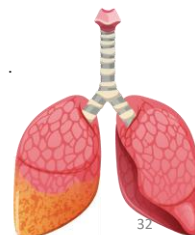
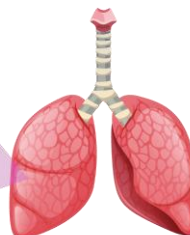
> Breathing in small particles that accumulate in & irritates the lungs:

Ex. **Mesothelioma** – asbestos and ↑ risk for lung cancer.



silica

Process of mesothelioma cause from asbestos



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Restrictive Respiratory Disorders

- **Pulmonary fibrosis** = _____

HAS MANY CAUSES

> Breathing in small particles that accumulate in & irritates the lungs:

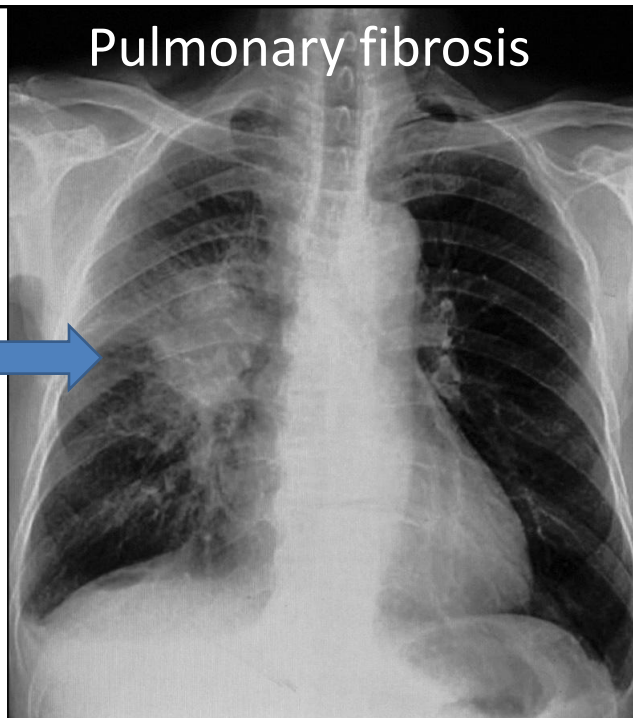
Ex. **Smoking** - _____



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Pulmonary fibrosis



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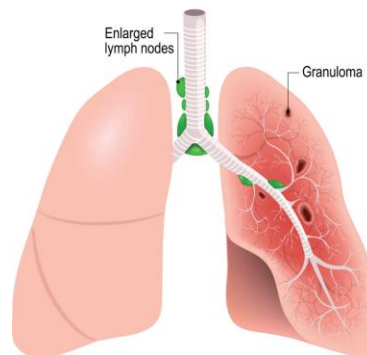
Restrictive Respiratory Disorders

- **Pulmonary sarcoidosis** = rare condition that causes formation of small patches of swollen tissue (granulomas). ~200,000 cases / year in US.

HAS MANY CAUSES

> Breathing in bacteria, viruses, or chemicals that irritates the lungs:

SARCOIDOSIS



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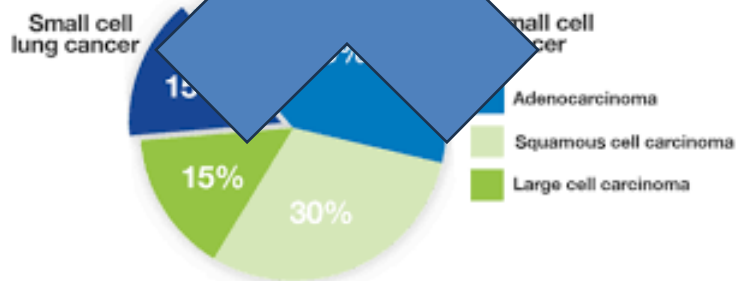
Restrictive Respiratory Disorders – Lung cancer

Main Types:

Non-small cell lung cancer (85% of all lung cancers in this group)

1. Adenocarcinoma (~40% of lung cancers)
2. Squamous cell carcinoma (~30% of lung cancers)
3. Large cell carcinoma (~10% of lung cancers)

Small cell lung cancer (~15% of all lung cancers in this group)



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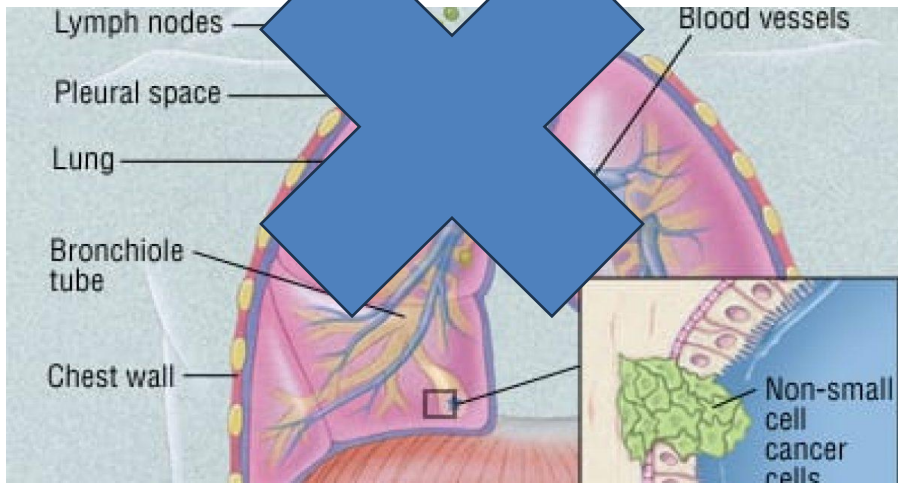
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Respiratory Illnesses – Lung cancer

Non-small cell lung cancer

1. Adenocarcinoma. This is the most common type (40% of all lung cancers), and starts in the mucus making gland cells in the lining of your airways. More likely to be localized to one area.

<https://www.2minutemedicine.com/patient-book/adenocarcinoma-of-the-lung/>



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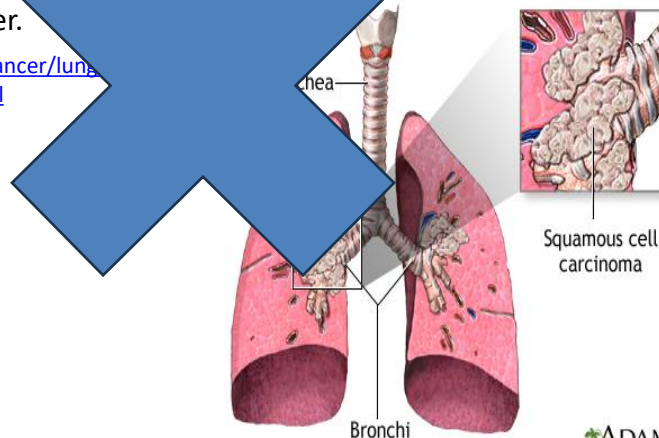
Respiratory Illnesses – Lung cancer

Non-small cell lung cancer

1. Adenocarcinoma. This is the most common type and starts in the mucus making gland cells in the lining of your airways.

2. Squamous cell cancer. 2nd most common lung cancer. Cancer of squamous cells lining the inside of the bronchi (bronchi) in the lungs. They are often found in the central part of the lung. Tends to be a slow growing cancer.

<https://www.cancer.org/cancer/lung-cancer/about/what-is.html>



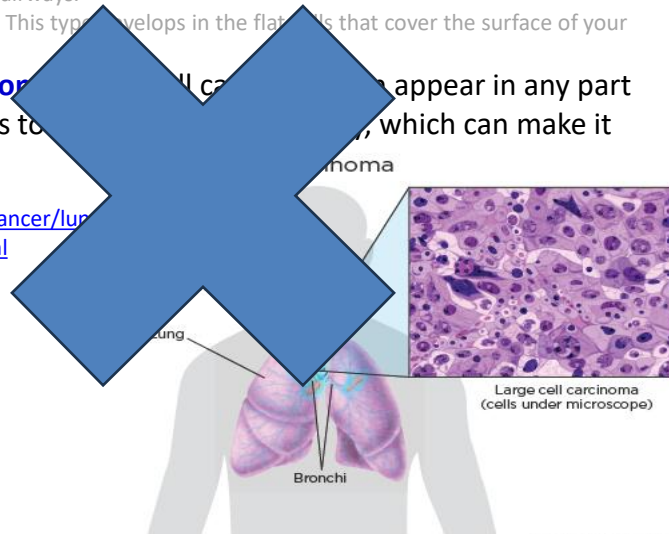
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Respiratory Illnesses – Lung cancer

Non-small cell lung cancer

1. **Adenocarcinoma.** This is the most common type and starts in the mucus making gland cells in the lining of your airways.
2. **Squamous cell cancer.** This type develops in the flat cells that cover the surface of your airways. ...
3. **Large cell carcinoma.** Large cell carcinoma can appear in any part of the lung. It tends to grow quickly, which can make it harder to treat.

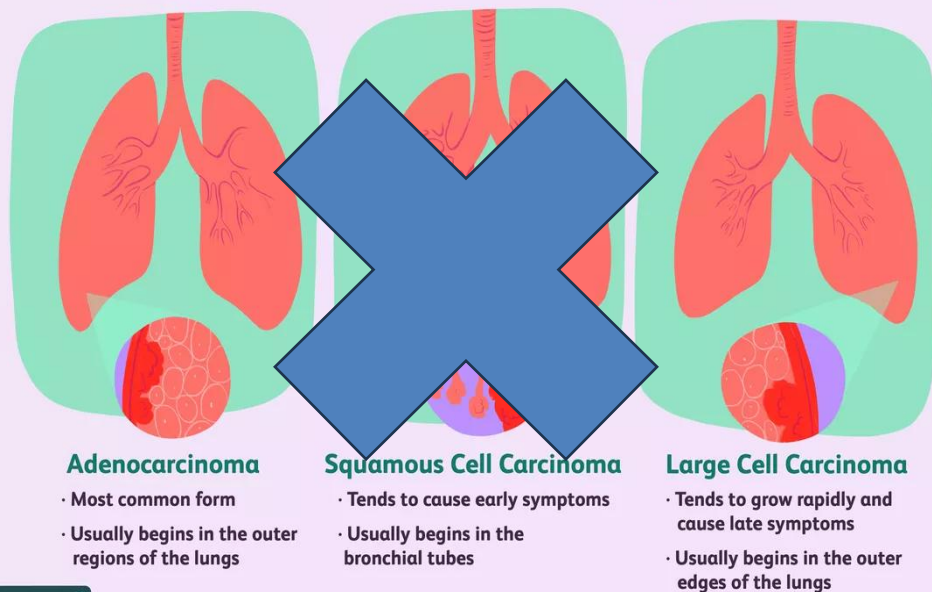
<https://www.cancer.org/cancer/lung-cancer/about/what-is.html>



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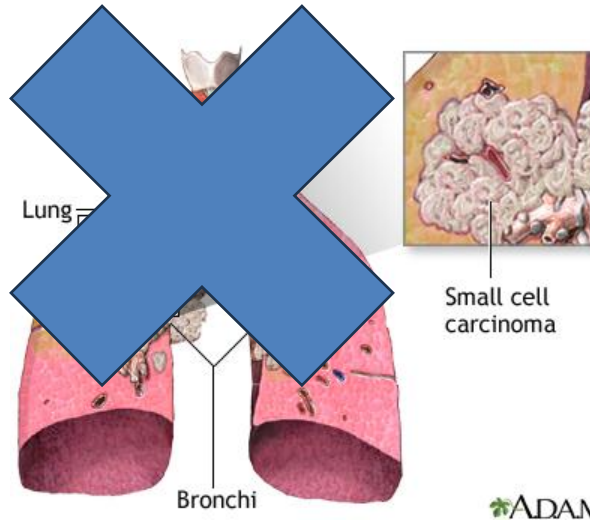
Types of Non-Small Cell Lung Cancer



verywell

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Small Cell Lung Cancer (SCLC): is the most aggressive form of lung cancer. It usually starts in the breathing tubes (bronchi) in the center of the chest. Although the cancer cells are small, they grow very quickly and create large tumors



ADAM 41

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5-year relative survival rates for non-small cell lung cancer

These numbers are based on people diagnosed with NSCLC between 2009 and 2015.

SEER stage	5-year relative survival rate
Localized	61% (has not spread from original spot)
Regional	35% (has spread from original spot, but not metastasized)
Distant	2% (has metastasized)
All stages combined	25%

5-year relative survival rates for small cell lung cancer

These numbers are based on people diagnosed with SCLC between 2009 and 2015.

SEER stage	5-year relative survival rate
Localized	27% (has not spread from original spot)
Regional	16% (has spread from original spot, but not metastasized)
Distant	3% (has metastasized)
All stages combined	6%

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Restrictive Disorder = Lung tissue is damaged. Lungs are stiff, or respiratory muscles are weak.

Example: **Pulmonary fibrosis**

Obstructive Disorders = Lung tissue is normal, but resistance is increased (airways are narrowed)

Examples: **Asthma**

COPD = Chronic obstructive ^{pulmonary} disorder
 - which includes **emphysema** &
 - **chronic bronchitis**
 • Cystic fibrosis
 • Chronic bronchitis ~ chronic bronchiole inflammation.
 • Sleep apnea

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Asthma

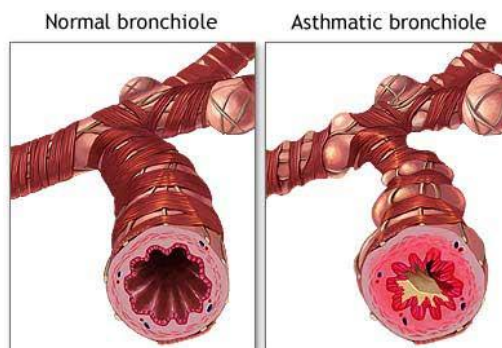
= inflamed airways from inhaling an an
 From inhaling antigens that person's immune system reacts to.

Muscles around the bronchioles are hyper-excitable

- Is an obstructive disorder due to bronchiole inflammation, mucous secretion, & narrowing of airways (**bronchoconstriction**).

Question:

What drug would you give as a treatment to dilate the bronchioles?
albuterol

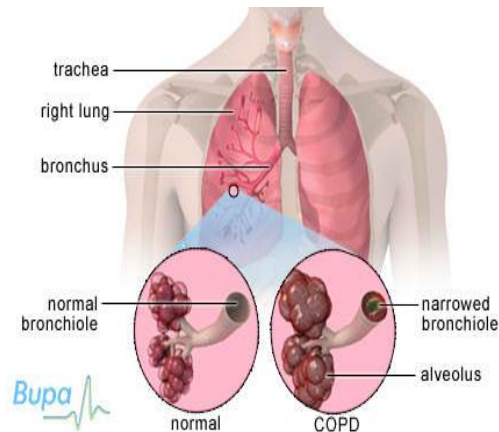


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Chronic Obstructive Pulmonary Disease (COPD)

Chronic inflammation of airways alveolar tissue

- narrows airways & destroys alveolar walls
- proliferation of mucus-secreting goblet cells
- development of scar (fibrous) tissue = **pulmonary fibrosis**
- **Obstructive disorder** – due to mucus buildup and narrowed airways.



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PHYSIOLOGY IN HEALTH AND DISEASE



People with pulmonary disorders frequently complain of **dyspnea**, which is a feeling of “shortness of breath.” The dyspnea, wheezing, and other symptoms of **asthma** are produced by increased resistance to airflow through the bronchioles (asthma is an obstructive pulmonary disorder, as discussed previously). The increased resistance to air flow is caused by bronchoconstriction and inflammation that may be provoked by allergic reactions (chapter 11).

Asthma may be treated on a sustained basis with glucocorticoid drugs (related to cortisol) that inhibit inflammation, thereby preventing or reducing the severity of “attacks.” New drugs (such as Singulair) that block the action of leukotrienes, a type of regulatory fatty acid (related to prostaglandins) that promote asthma, are now also available for this purpose. Acute asthma attacks are commonly treated with inhaled drugs (such as Albuterol) that stimulate the β_2 -adrenergic receptors (a type of receptor for epinephrine and norepinephrine; see chapter 6) that promote dilation of the bronchioles.

Alveolar tissue is destroyed in **emphysema**, resulting in fewer but larger alveoli (see fig. 12.8). The loss of alveoli reduces the ability of the bronchioles to remain open during expiration, causing *air trapping* during expiration when the bronchioles collapse. The most common cause of emphysema is cigarette smoking, which indirectly causes different protein-digesting enzymes to destroy the lung tissue. The loss of alveoli and air

trapping reduces gas exchange, so that people with emphysema have difficulty in both oxygenating the blood and eliminating carbon dioxide. Because of this, people with emphysema must often breathe from an oxygen tank.

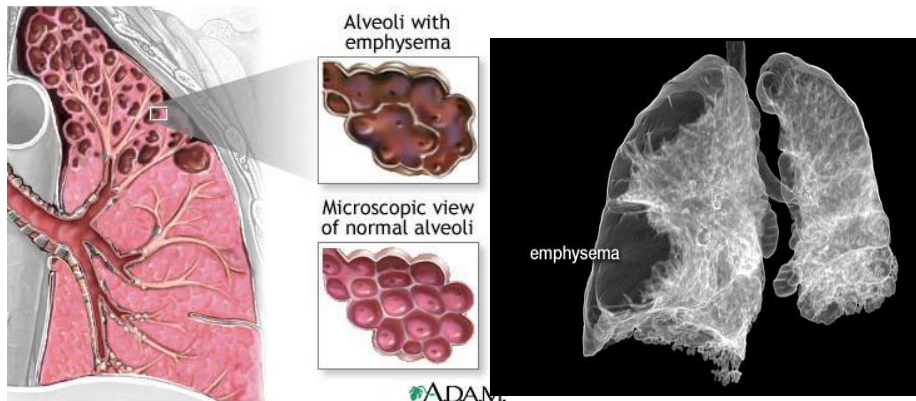
Chronic obstructive pulmonary disease (COPD) is characterized by chronic inflammation with narrowing of the airways and destruction of the alveolar walls. Included in the COPD category is *emphysema* and *chronic obstructive bronchiolitis*, which refers to fibrosis and obstruction of the bronchioles. The condition results in a faster age-related decline in the FEV_1 (discussed previously). COPD differs from asthma in that, unlike asthma, COPD is not reversible with the use of a bronchodilator such as Albuterol. Also unlike asthma, COPD is not helped much by inhaled glucocorticoids (drugs related to hydrocortisone). The vast majority of people with COPD are smokers, and stopping smoking once COPD has begun does not seem to stop its progression. In addition to the pulmonary problems directly caused by COPD, this condition increases the risk of pneumonia, pulmonary emboli (traveling blood clots), and heart failure. Patients with COPD may develop *cor pulmonale*—pulmonary hypertension with eventual failure of the right ventricle. COPD is now the fifth leading cause of death in the United States, and scientists have estimated that by 2020 it will become the third leading cause of death worldwide.

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Emphysema

Chronic destruction of alveolar tissue (walls between alveoli lost)

- reduces area for gas exchange
- alveoli expand easily, but can't empty easily (air-trapping disorder)
- **obstructive disorder**



<http://www.nlm.nih.gov/medlineplus/ency/images/ency/fullsize/17055.jpg>

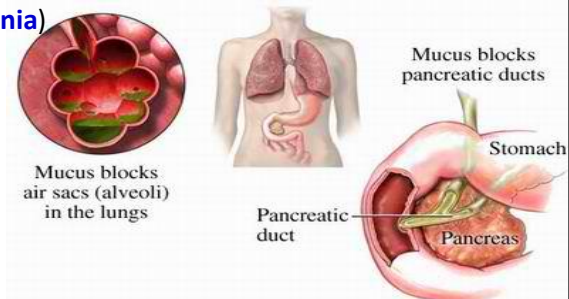
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Cystic Fibrosis

- Genetic disorder affecting Cl⁻ channels on alveoli membrane. **Obstructive disorder**

Results in buildup of mucus within alveoli causing:

- Dilutes surfactant
- ↓ decreased functional alveolar size
- ↑ surface tension & intra-alveolar pressure (harder for alveoli to expand)
- ↓ with gas exchange
- Warmth & moisture (mucus) aids bacterial growth. (Vulnerable to **pneumonia**)



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Sleep apnea *(added slide)*

Sleep apnea = muscles of throat relax during sleep (or nasal septum issues) allowing the tongue & soft palate to collapse blocking airway. Interrupted breathing during sleep. A-pnea = absence of breathing

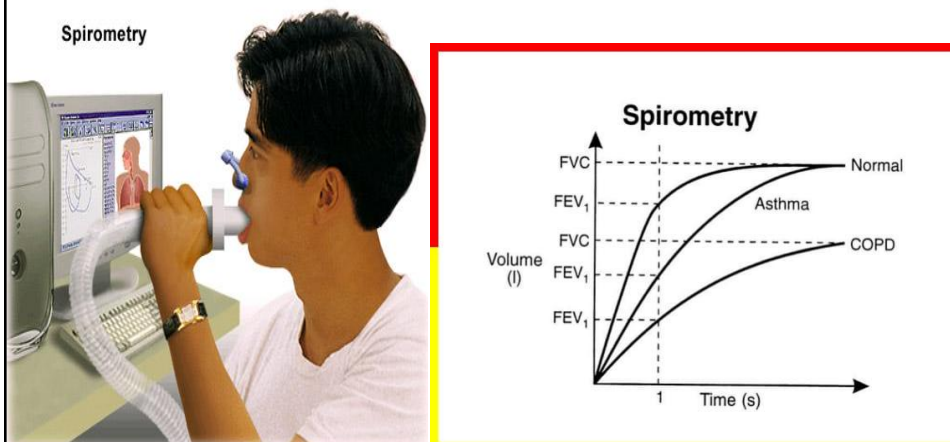
Treatment: Continuous positive airway pressure (CPAP) machines



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3. Respiratory Disorders – diagnosing them

- **Spirometry:** air movement during respiration recorded on a spirogram.
 - Measures lung volumes and capacities
 - Can diagnose restrictive and obstructive lung disorders



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Additional Respiratory Vocabulary (*I moved this slide up*):

Apnea = absence of breathing

Dyspnea = labored or difficult breathing

Eupnea = normal breathing at rest

Hyperventilation = excessively rapid ventilation (will decrease alveolar CO₂)

Hypoventilation = low ventilation (will increase alveolar CO₂)

Pneumothorax = presence of gas in intrapleural space causing lung collapse

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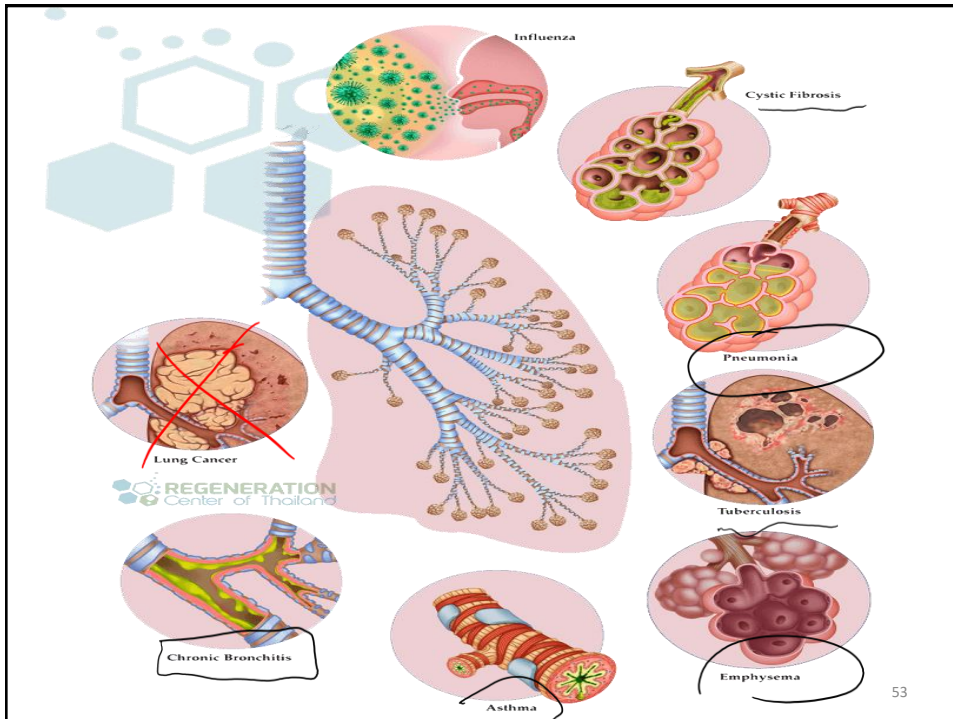
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3. Respiratory Disorders – diagnosing them

- Spirometry
- Lung X-ray
- Lung CT scan



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Review

Respiratory Disorders & diagnosing them

- Restrictive
 - pulmonary fibrosis
 - silicosis, mesothelioma
 - pulmonary sarcoidosis
 - ~~Lung cancer~~
 - Non-small cell lung cancer
 - » Adenocarcinoma, squamous cell carcinoma, large cell carcinoma
 - Small cell lung cancer
- Obstructive
 - Asthma
 - COPD
 - Emphysema
 - chronic bronchitis
 - cystic fibrosis
 - Sleep apnea

Spirometry

Additional respiratory vocabulary

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4. Basics of Gas Exchange at Lungs and at Body Tissues

Gas exchange between 2 structures is dependent on pressure gradient of dissolved O₂ & CO₂

* Gas moves from side with higher pressure (from dissolved gases) to side with lower pressure & visa versa

* Gas wants to move “downhill” from high to low pressure!



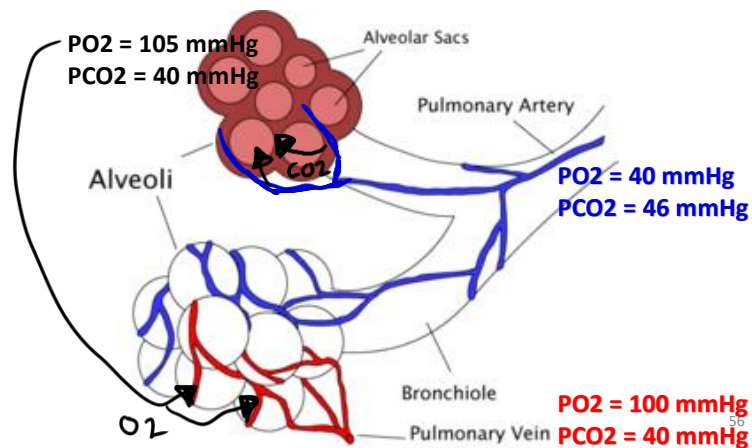
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Gas exchange between lung alveoli & pulmonary vessels:

> Alveolar PO₂ = 105 mmHg, higher than that in pulmonary arteries (40 mmHg)

> Alveolar PCO₂ = 40 mmHg, lower than that in pulmonary arteries (46 mmHg)

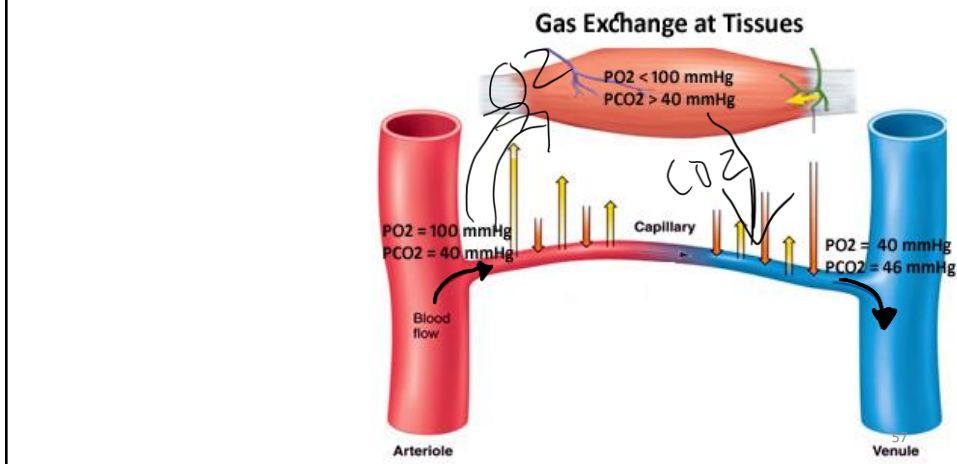


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Gas exchange between systemic capillaries & tissues:

> Tissue PO_2 (<100 mmHg) = lower than O_2 -rich arterial blood (100 mmHg)

> Tissue PCO_2 (>40 mmHg) = higher than that in arterial blood (40 mmHg)



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5. Regulation of Respiration – regulation of blood O_2 & CO_2

Autonomic motor control breathing involves:

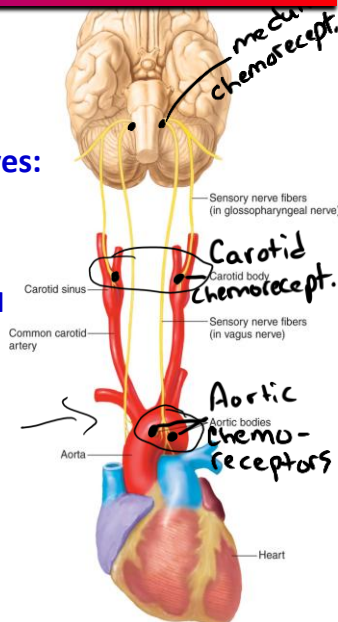
Sensors of pH = Chemoreceptors:

➤ **Aorta & carotid artery chemoreceptors (called peripheral chemoreceptors)**

- sense blood O_2 and CO_2 levels

➤ **Medulla chemoreceptors (called central chemoreceptors)**

- sense CSF O_2 and CO_2 levels



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5. Regulation of Respiration – regulation of blood O₂ & CO₂

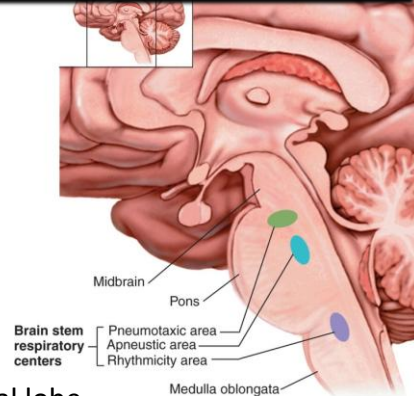
Motor neurons from 3 brain areas control breathing muscles:

1) Voluntary Breathing

= primary motor cortex of frontal cerebral lobe.

2) Involuntary Breathing (integrating center) =

- Medulla – respiratory center regulates respiratory rate.
- Pons – apneustic center (stimulate inhalation)
– pneumotaxic center (inhibit inhalation)



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5. Regulation of Respiration – regulation of blood pH

Normal Blood pH = 7.35 – 7.45

Blood pH maintained by buffering CO₂ with HCO₃⁻

Blood with high CO₂ or H⁺ content = acidic (acidosis)

Blood w/ lower CO₂ or high HCO₃⁻ content = alkaline (alkalosis)

Blood pH (Acid/Base balance) based primarily on blood CO₂ content and metabolic activities in body:

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Minute ventilation = the depth and rate of breathing.

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- **What happens to minute ventilation after hypoventilation?**

Holding breath increases CO₂ in blood (respiratory acidosis)

↓ pH

Stimulus = high blood CO₂

Sensors = chemoreceptors in aortic arch & carotid arteries

Integrating center = medulla oblongata (respiratory center), which stimulates increased minute ventilation.

Effectors = respiratory muscles ^{exhale more CO₂ (acid)} cause increased minute ventilation. More CO₂ exhaled

Effect = blood pH increases back to normal

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- What happens to minute ventilation **after hyperventilation?**

Rapid breathing decreases CO₂ in blood (causes respiratory alkalosis)

Stimulus = low blood CO₂

Sensors = chemoreceptors in aortic arch & carotid arteries

Integrating center = medulla oblongata (respiratory center)

Effectors = respiratory muscles cause decreased minute ventilation. Less CO₂ exhaled.

Effect = blood pH decreased back to normal

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Review

- Gas exchange at tissues & at alveoli of lungs
Depends on differences in partial pressures of O₂ and CO₂
- Regulation of breathing (voluntary vs involuntary)
 - Primary motor cortex (voluntary)
 - Medulla & Pons (involuntary)
- Acid / Base imbalance
 - Respiratory Acidosis & alkalosis
 - Changes to minute ventilation to fix blood pH

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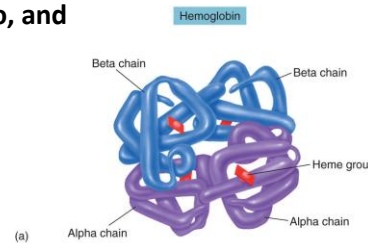
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6. Hemoglobin & Hemoglobin Disorders

Hemoglobin = respiratory pigment that binds to, and lets go of, oxygen.

- 4 protein chains w/4 iron-containing heme (pigments)
- Each heme group binds with 1 O₂ molecule
- Each RBC has ~280 million hemoglobin molecules (each RBC can carry ~billion O₂ molecules! (4 X 280 million))
- Hemoglobin bound to O₂ = “**oxyhemoglobin**” (Arterial blood 97% saturated w/oxyhemoglobin = bright red)
- Hemoglobin lacking O₂ = “**deoxyhemoglobin**” (venous blood dull red or maroon)

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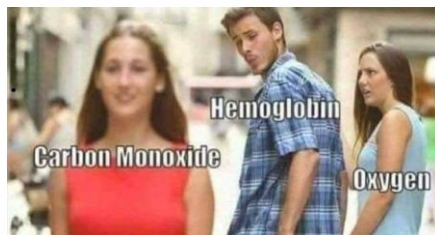


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Hemoglobin Disorders:

Carbon Monoxide = odorless, color-less gas that has greater binding to hemoglobin than to oxygen. Thus, carbon monoxide gas outcompetes O₂ for binding sites on hemoglobin, creating **carboxyhemoglobin** on RBCs. Known as “carboxyhemoglobinemia”



Result : Carboxyhemoglobin doesn't deliver O₂ to cells, and carbon monoxide can't be used in ETC to make ATP

> Hypoxia – tissues starved for O₂

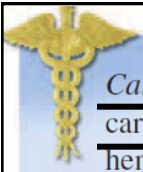
> Death (*CDC says 400 people die/year of this in US*)



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 **CLINICAL APPLICATIONS**

Carboxyhemoglobin is hemoglobin with its heme groups bound to carbon monoxide instead of oxygen. Because the bond that hemoglobin forms with carbon monoxide is 210 times stronger than the bond it forms with oxygen, carbon monoxide tends to displace oxygen and remain bound as it goes through the circulation, thereby reducing the oxygen carried by the blood. Carboxyhemoglobin has a cranberry juice color, compared to the tomato juice color of oxyhemoglobin. The body actually produces a tiny amount of carbon monoxide, and so there is always some carboxyhemoglobin; however, according to federal standards, active nonsmokers should not have more than 1.5% carboxyhemoglobin. Higher percentages of carboxyhemoglobin constitute **carbon monoxide poisoning**. Nonsmokers living in certain smoggy cities can have 3% carboxyhemoglobin levels, whereas smokers there may have 10% or more carboxyhemoglobin. Much higher percentages can be produced by smoke inhalation or suicide attempts, and may cause death. People with carbon monoxide poisoning are often given hyperbaric oxygen therapy (previously described).

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Hemoglobin Disorders contin...

Methemoglobinemia = disorder in which hemoglobin's iron (a component of heme) is "ferric" rather than "ferrous".

- > this hemoglobin called **methemoglobin** (pronounce as "met-hemoglobin")
- > Methemoglobin has ↓ ability to release (unload) O₂ at tissues.
- > Tissues chronically O₂-starved.
- > Patients are hypoxic & BLUE!

"Blue baby syndrome" = babies turn blue (hypoxia) from drinking milk made w/nitrate contaminated water. Nitrate causes formation of methemoglobin.

CDC says ~3,000 cases in babies/year in US



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Hemoglobin Disorders contin...

Neonatal jaundice At birth switch from **hemoglobin-F** (fetal) to **hemoglobin-A** (adult)

- Body removes RBCs with hemoglobin f.
- Liver removes biliruben from destroyed hemoglobin f.
- Liver sometimes not mature enough to remove biliruben.
- Bilirubin builds up (*hyperbilirubinemia*)
- Baby turns yellow. (*affecting about 60% of term and 80% of preterm babies*)

Treatment:

"blue light exposure" – breaks bilirubin down to water-soluble form excreted by kidneys.



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CLINICAL APPLICATIONS

The fetus has a different form of hemoglobin (called **hemoglobin F**) than its mother, who has adult hemoglobin (**hemoglobin A**).

The reason that the fetus has a different form of hemoglobin than its mother is because hemoglobin F has a stronger affinity for oxygen than does hemoglobin A, causing oxygen to move from the mother's to the fetus's blood. Hemoglobin F has a higher affinity for oxygen because it can't bind to 2,3-BPG, which works to reduce the bond strength for oxygen in the mother's red blood cells. The fetus stops producing hemoglobin F and begins producing hemoglobin A at about week 38 of pregnancy. When the fetus switches from hemoglobin F to hemoglobin A, it destroys its old red blood cells and converts the heme groups into bile pigment, or bilirubin (chapter 14). Too much bilirubin can cause jaundice (a yellowing of the skin and mucus membranes), and jaundice produced for this reason is called physiological neonatal jaundice. Putting the babies under blue light converts the bilirubin into a more water-soluble derivative that they can excrete in the urine.

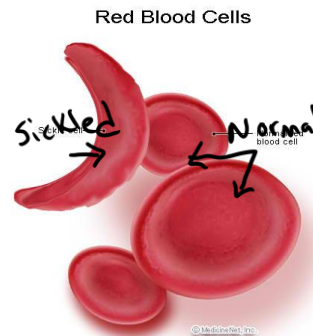
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Hemoglobin Disorders contin...

Sickle Cell Anemia = homozygous recessive condition in which body produces RBCs with **hemoglobin-S** rather than **hemoglobin-A**.

sickle cell is from 1 amino acid substitution (valine for glutamic acid) on hemoglobin.

- Hemoglobin-S turns RBCs into sickle-shape.
- Sickled RBCs carry less O₂ (cause hypoxia)
- Sickled RBCs tend to form clots (thrombus)
- Patients more prone to embolism.
- More prone to ischemic events.



CDC says 100,000 people in US, and globally millions of people, living with sickle cells dz.

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Review

- Hemoglobin O₂ transport:
 - Oxyhemoglobin & deoxyhemoglobin
 - Abnormal hemoglobin (carboxyhemoglobin, methemoglobin)
 - Neonatal jaundice
 - Sickle cell

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