**Pulmonary Ventilation**

The main functions of respiration are to provide oxygen to the tissues and remove carbon dioxide. The four major components of respiration are (1) *pulmonary ventilation*, which means the inflow and outflow of air between the atmosphere and the lung alveoli; (2) *diffusion of oxygen (O2) and carbon dioxide (CO2) between the alveoli and the* *blood*; (3) *transport of oxygen and carbon dioxide in the* *blood and body fluids* to and from the body’s tissue cells; and (4) *regulation of ventilation* and other facets of respiration.

**MECHANICS OF PULMONARY VENTILATION**

**MUSCLES THAT CAUSE LUNG EXPANSION AND CONTRACTION**

The lungs can be expanded and contracted in two ways: (1) by downward and upward movement of the diaphragm to lengthen or shorten the chest cavity, and (2)

by elevation and depression of the ribs to increase and decrease the anteroposterior diameter of the chest cavity. Normal quiet breathing is accomplished almost entirely

by the first method, that is, by movement of the diaphragm. During inspiration, contraction of the diaphragm pulls the lower surfaces of the lungs downward. Then,

during expiration, the diaphragm simply relaxes, and the *elastic recoil* of the lungs, chest wall, and abdominal structures compresses the lungs and expels the air. During

heavy breathing, however, the elastic forces are not powerful enough to cause the necessary rapid expiration, so extra force is achieved mainly by contraction of the

*abdominal muscles*, which pushes the abdominal contents upward against the bottom of the diaphragm, thereby compressing the lungs. The second method for expanding the lungs is to raise the rib cage. Raising the rib cage expands the lungs because, in the natural resting position, the ribs slant downward, thus allowing the sternum to fall backward toward the vertebral column. When the rib cage is elevated, however,

the ribs project almost directly forward, so the sternum also moves forward, away from the spine, making the anteroposterior thickness of the chest about 20 percent

greater during maximum inspiration than during expiration. *Therefore, all the muscles that elevate the chest cage* *are classified as muscles of inspiration, and the muscles*

*that depress the chest cage are classified as muscles of expiration*.The most important muscles that raise the rib cageare the *external intercostals*, but others that help are the

(1) *sternocleidomastoid* muscles, which lift upward on the sternum; (2) *anterior serrati*, which lift many of the ribs; and (3) *scaleni*, which lift the first two ribs.

The muscles that pull the rib cage downward during expiration are mainly (1) the *abdominal recti*, which have the powerful effect of pulling downward on the lower ribs at the same time that they and other abdominal muscles also compress the abdominal contents upward against the diaphragm, and (2) the *internal intercostals*.

The ribs during expiration are angled downward, and the external intercostals are elongated forward and downward. As they contract, they pull the upper ribs forward in relation to the lower ribs, which causes leverage on the ribs to raise them upward, thereby causing inspiration. The internal intercostals function exactly in the opposite manner, functioning as expiratory muscles because they angle between the ribs in the opposite direction and cause opposite leverage.

**PRESSURES THAT CAUSE THE MOVEMENT OF AIR IN AND OUT OF THE LUNGS**

The lung is an elastic structure that collapses like a balloon and expels all its air through the trachea whenever there is no force to keep it inflated. Also, there are no attachments between the lung and the walls of the chest cage, except where it is suspended at its hilum from the *mediastinum*, the middle section of the chest cavity. Instead, the lung “floats” in the thoracic cavity, surrounded by a thin layer of *pleural fluid* that lubricates movement of the lungs within the cavity. Further, continual suction of excess fluid into lymphatic channels maintains a slight suction between the visceral surface of the lung pleura and the parietal pleural surface of the thoracic cavity. Therefore, the lungs are held to the thoracic wall as if glued there, except that they are well lubricated and can slide freely as the chest expands and contracts.

**Pleural Pressure and Its Changes during Respiration.**

*Pleural pressure* is the pressure of the fluid in the thin space between the lung pleura and the chest wall pleura. As noted earlier, this pressure is normally a slight suction,

which means a slightly *negative* pressure. The normal pleural pressure at the beginning of inspiration is about−5 centimeters of water, which is the amount of suction

required to hold the lungs open to their resting level. During normal inspiration, expansion of the chest cage pulls outward on the lungs with greater force and creates

more negative pressure, to an average of about −7.5 centimeters.

These relationships between pleural pressure and changing lung volume. The lower shows the increasing negativity of the upper panel an increase in lung volume of 0.5

liter. Then, during expiration, the events are essentially reversed.

**Alveolar Pressure—The Air Pressure Inside the Lung Alveoli.** When the glottis is open and no air is flowinginto or out of the lungs, the pressures in all parts of therespiratory tree, all the way to the alveoli, are equal toatmospheric pressure, which is considered to be zero referencepressure in the airways—that is, 0 centimetersof water pressure. To cause inward flow of air into the alveoli during inspiration, the pressure in the alveoli mustfall to a value slightly below atmospheric pressure (below0). During normal inspiration, alveolar pressure decreases to about −1 centimeters of water. This slight negative pressure is enough to pull 0.5 liter of air into the lungs in the 2 seconds required for normal quiet inspiration. During expiration, alveolar pressure rises to about +1 centimeter of water, which forces the 0.5 liter of inspired air out of the lungs during the 2 to 3 seconds of expiration.

**Transpulmonary Pressure—The Difference between Alveolar and Pleural Pressures**

The *transpulmonary pressure* is the pressure difference between that in the alveoli and that on the outer surfaces of the lungs (pleural pressure), and it is a measure

of the elastic forces in the lungs that tend to collapse the lungs at each instant of respiration, called the *recoil* *pressure*.

**Compliance of the Lungs**

The extent to which the lungs will expand for each unit increase in transpulmonary pressure (if enough time is allowed to reach equilibrium) is called the *lung compliance*. The total compliance of both lungs together in the normal adult human averages about 200 milliliters of air per centimeter of water transpulmonary pressure. That is, every time the transpulmonary pressure increases 1 centimeter of water, the lung volume, after 10 to 20 seconds, will expand 200 milliliters.

**Compliance Diagram of the Lungs** is a diagram relating lung volume changes to changes in pleural pressure, which, in turn, alters transpulmonary pressure. Note that the relation is different for inspiration and expiration. Each curve is recorded by changing

the pleural pressure in small steps and allowing the lung volume to come to a steady level between successive steps. The two curves are called, respectively, the *inspiratory*

*compliance curve* and the *expiratory compliance curve*, and the entire diagram is called the *compliance diagram of the lungs*.The characteristics of the compliance diagram aredetermined by the elastic forces of the lungs. These forcescan be divided into two parts: (1) *elastic forces of the lung* *tissue* and (2) *elastic forces caused by surface tension of the fluid that lines the inside walls of the alveoli* and other lung

air spaces. The elastic forces of the lung tissue are determined mainly by *elastin* and *collagen* fibers interwoven among the lung parenchyma. In deflated lungs, these fibers are in an elastically contracted and kinked state; then, when the lungs expand, the fibers become stretched and unkinked, thereby elongating and exerting even more elastic force. The elastic forces caused by surface tension are much

more complex. The significance of surface tension which compares the compliance

diagram of the lungs when filled with saline solution and when filled with air. When the lungs are filled with air, there is an interface between the alveolar fluid and the air

in the alveoli. In lungs filled with saline solution, there is no air-fluid interface, and therefore, the surface tension effect is not present; only tissue elastic forces are operative in the lung filled with saline solution. Note that transpleural pressures required to expand air-filled lungs are about three times as great as those required to expand lungs filled with saline solution. Thus, one can conclude that *the tissue elastic forces tending* *to cause collapse of the air-filled lung represent only* *about one third of the total lung elasticity, whereas the* *fluid-air surface tension forces in the alveoli represent* *about two thirds*. The fluid-air surface tension elastic forces of the lungs

also increase tremendously when the substance called *surfactant* is *not* present in the alveolar fluid.

**Surfactant, Surface Tension, and Collapse of the Alveoli**

**Principle of Surface Tension.** When water forms a surface with air, the water molecules on the surface of the water have an especially strong attraction for one another. As a result, the water surface is always attempting to contract. This is what holds raindrops together—a tight contractile membrane of water molecules around the

entire surface of the raindrop. Now let us reverse these principles and see what happens on the inner surfaces of the alveoli. Here, the water surface is also attempting to contract. This tends to force air out of the alveoli through the bronchi and, in doing so, causes the alveoli to try to collapse. The net effect is to cause an elastic contractile

force of the entire lungs, which is called the *surface tension elastic force*.

**Surfactant and Its Effect on Surface Tension.**

Surfactant is a *surface active agent in water*, which means that it greatly reduces the surface tension of water. It is secreted by special surfactant-secreting epithelial cells

called *type II alveolar epithelial cells*, which constitute about 10 percent of the surface area of the alveoli. These cells are granular, containing lipid inclusions that are

secreted in the surfactant into the alveoli. Surfactant is a complex mixture of several phospholipids, proteins, and ions. The most important components are the phospholipid *dipalmitoyl phosphatidylcholine,* *surfactant apoproteins*, and *calcium ions*. The dipalmitoyl phosphatidylcholine and several less important phospholipids

are responsible for reducing the surface tension. They perform this function by not dissolving uniformly in the fluid lining the alveolar surface. Instead, part of the

molecule dissolves while the remainder spreads over the surface of the water in the alveoli. This surface has from one twelfth to one half the surface tension of a pure water surface. In quantitative terms, the surface tension of different water fluids is approximately the following: pure water, 72 dynes/cm; normal fluids lining the alveoli but without surfactant, 50 dynes/cm; normal fluids lining the alveoli and *with* normal amounts of surfactant included, between 5 and 30 dynes/cm.

**Pressure in Occluded Alveoli Caused by Surface Tension.**

If the air passages leading from the alveoli of thelungs are blocked, the surface tension in the alveoli tendsto collapse the alveoli. This collapse creates positive pressurein the alveoli, attempting to push the air out. Theamount of pressure generated in this way in an alveolus canbe calculated from the following formula:

Pressure =2.Surface tension/ Radius of alveolus

For the average-sized alveolus with a radius of about 100 micrometers and lined with *normal surfactant*, this calculates to be about 4 centimeters of water pressure

(3 mm Hg). If the alveoli were lined with pure water without any surfactant, the pressure would calculate to be about 18 centimeters of water pressure—4.5 times

as great. Thus, one sees the importance of surfactant in reducing alveolar surface tension and therefore also reducing the effort required by the respiratory muscles to expand the lungs.

**Effect of Alveolar Radius on the Pressure Caused by** **Surface Tension.**

Note from the preceding formula that the pressure generated as a result of surface tension in the alveoli is *inversely* affected by the radius of the alveolus, which means that the smaller the alveolus, the greater the alveolar pressure caused by the surface tension. Thus, when the alveoli have half the normal radius (50 instead of 100 micrometers), the pressures noted earlier are doubled. This phenomenon is especially significant in small premature babies, many of whom have alveoli with radii less than

one quarter that of an adult person. Further, surfactant does not normally begin to be secreted into the alveoli until between the sixth and seventh months of gestation, and in some cases, even later. Therefore, many premature babies have little or no surfactant in the alveoli when they are born, and their lungs have an extreme tendency to collapse, sometimes as great as six to eight times that in a normaladult person. This situation causes the condition called *respiratory distress syndrome of the newborn*. It is fatal if not treated with strong measures, especially properly applied continuous positive pressure breathing.

**EFFECT OF THE THORACIC CAGE ON LUNG EXPANSIBILITY**

Thus far, we have discussed the expansibility of the lungs alone, without considering the thoracic cage. The thoracic cage has its own elastic and viscous characteristics, similar to those of the lungs; even if the lungs were not present in the thorax, muscular effort would still be required to expand the thoracic cage.

**Compliance of the Thorax and the Lungs Together**

The compliance of the entire pulmonary system (the lungs and thoracic cage together) is measured while expanding the lungs of a totally relaxed or paralyzed subject. To measure compliance, air is forced into the lungs a little at a time while recording lung pressures and volumes. To inflate this total pulmonary system, almost twice as much pressure as is required to inflate the same lungs after removal from the chest cage is necessary. Therefore, the compliance of the combined lung-thorax system is almost exactly one half that of the lungs alone— 110 milliliters of volume per centimeter of water pressure for the combined system, compared with 200 ml/cm for the lungs alone. Furthermore, when the lungs are expanded to high volumes or compressed to low volumes, the limitations of the chest become extreme. When near these limits, the compliance of the combined lung-thorax system can be less than one fifth that of the lungs alone.

**“Work” of Breathing**

We have already pointed out that during normal quiet breathing, all respiratory muscle contraction occurs during inspiration; expiration is almost entirely a passive process caused by elastic recoil of the lungs and chest cage. Thus, under resting conditions, the respiratory muscles normally perform “work” to cause inspiration but not to cause

expiration. The work of inspiration can be divided into three fractions: (1) that required to expand the lungs against the lung and chest elastic forces, called *compliance work* or *elastic* *work*; (2) that required to overcome the viscosity of the lung and chest wall structures, called *tissue resistance work*; and (3) that required to overcome airway resistance to movement of air into the lungs, called *airway resistance work*.

**Energy Required for Respiration.**

During normal quiet respiration, only 3 to 5 percent of the total energy expended

by the body is required for pulmonary ventilation. However, during heavy exercise, the amount of energy required can increase as much as 50-fold, especially if the person has any degree of increased airway resistance or decreased pulmonary compliance. Therefore, one of the major limitations on the intensity of exercise that can be performed is the person’s ability to provide enough muscle energy for the respiratory process alone.

PULMONARY VOLUMES AND CAPACITIES

**RECORDING CHANGES IN PULMONARY VOLUME—SPIROMETRY**

Pulmonary ventilation can be studied by recording the volume movement of air into and out of the lungs, a method called *spirometry*. It consists of a drum inverted over

a chamber of water, with the drum counterbalanced by a weight. In the drum is a breathing gas, usually air or oxygen; a tube connects the mouth with the gas chamber.

When one breathes into and out of the chamber, the drum rises and falls, and an appropriate recording is made on a moving sheet of paper.

For ease in describing the events of pulmonary ventilation, the air in the lungs has been subdivided in this diagram into four *volumes* and four *capacities*, which are the average for a *young adult man*.

Four pulmonary lung volumes when added together, equal the maximum volume to which the lungs can be expanded. The significance of each of these volumes is the following:

1. The *tidal volume* is the volume of air inspired or expired with each normal breath; it

amounts to about 500 milliliters in the average adult male.

2. The *inspiratory reserve volume* is the extra volume of air that can be inspired over and above the normal tidal volume when the person inspires with full force; it is usually equal to about 3000 milliliters.

3. The *expiratory reserve volume* is the maximum extra volume of air that can be expired by forceful expiration after the end of a normal tidal expiration; this volume normally amounts to about 1100 milliliters.

4. The *residual volume* is the volume of air remaining in the lungs after the most forceful expiration; this volume averages about 1200 milliliters.

**Pulmonary Capacities**

In describing events in the pulmonary cycle, it is sometimes desirable to consider two or more of the volumes together. Such combinations are called *pulmonary capacities*.

The important pulmonary capacities, which can be described as follows:

1. The *inspiratory capacity* equals the *tidal volume* plus the *inspiratory reserve volume.* This capacity isthe amount of air (about 3500 milliliters) a personcan breathe in, beginning at the normal expiratorylevel and distending the lungs to the maximumamount.

 2. The *functional residual capacity* equals the *expiratory reserve volume* plus the *residual volume.* Thiscapacity is the amount of air that remains in thelungs at the end of normal expiration (about 2300milliliters).

3. The *vital capacity* equals the *inspiratory reserve volume* plus the *tidal volume* plus the *expiratory reserve volume.* This capacity is the maximumamount of air a person can expel from the lungsafter first filling the lungs to their maximum extentand then expiring to the maximum extent (about4600 milliliters).

4. The *total lung capacity* is the maximum volume to which the lungs can be expanded with the greatest possible effort (about 5800 milliliters); it is equal to the *vital capacity* plus the *residual volume.* All pulmonary volumes and capacities are usually about 20 to 25 percent less in women than in men, and they are greater in large and athletic people than in small and asthenic people.

**ABBREVIATIONS AND SYMBOLS USED IN PULMONARY FUNCTION STUDIES**

Spirometry is only one of many measurement procedures that the pulmonary physician uses daily. Many of these measurement procedures depend heavily on mathematical

computations. To simplify these calculations, as well as the presentation of pulmonary function data, several abbreviations and symbols have become standardized.

 more important of these . Using these symbols, we present here a few simple algebraic

exercises showing some of the interrelations among the pulmonary volumes and capacities; the student should think through and verify these interrelations.

VC = IRV + VT + ERV

VC = IC + ERV

TLC = VC + RV

TLC = IC + FRC

FRC = ERV + RV

 **DETERMINATION OF FUNCTIONAL RESIDUAL CAPACITY, RESIDUAL**

**VOLUME, AND TOTAL LUNG CAPACITY—HELIUM DILUTION METHOD**

The functional residual capacity (FRC), which is the volume of air that remains in the lungs at the end of each normal expiration, is important to lung function. Because its value changes markedly in some types of pulmonary disease, it is often desirable to measure this capacity. The spirometer cannot be used in a direct way to measure the

FRC because the air in the residual volume of the lungs cannot be expired into the spirometer, and this volume constitutes about one half of the FRC. To measure FRC,

the spirometer must be used in an indirect manner, usually by means of a helium dilution method, as follows. A spirometer of known volume is filled with air mixed

with helium at a known concentration. Before breathing from the spirometer, the person expires normally. At the end of this expiration, the remaining volume in the lungs is equal to the FRC. At this point, the subject immediately begins to breathe from the spirometer, and the gases of the spirometer mix with the gases of the lungs. As a result, the helium becomes diluted by the FRC gases, and the volume of the FRC can be calculated from the degree of dilution of the helium, using the following formula:

FRC=((CiHg/CfHg)-1)Vispir

where *FRC* is functional residual capacity, *CiHe* is initial concentration of helium in the spirometer, *CfHe* is final concentration of helium in the spirometer, and *ViSpir* is initial volume of the spirometer. Once the FRC has been determined, the residual

volume (RV) can be determined by subtracting expiratory reserve volume (ERV), as measured by normal spirometry, from the FRC. Also, the total lung capacity (TLC) can be determined by adding the inspiratory capacity (IC) to the FRC. That is,

RV= FRC- ERV and TLC= FRC+ IC

**MINUTE RESPIRATORY VOLUMEEQUALS RESPIRATORY RATE TIMES TIDAL VOLUME**

The *minute respiratory volume* is the total amount of new air moved into the respiratory passages each minute and is equal to the *tidal volume* times the *respiratory rate per* *minute*. The normal tidal volume is about 500 milliliters, and the normal respiratory rate is about 12 breaths per minute. Therefore, the *minute respiratory volume averages* *about 6 L/min*. A person can live for a short period with a minute respiratory volume as low as 1.5 L/min and a respiratory rate of only 2 to 4 breaths per minute. The respiratory rate occasionally rises to 40 to 50 per minute, and the tidal volume can become as great as the vital capacity, about 4600 milliliters in a young adult man.

This can give a minute respiratory volume greater than 200 L/min, or more than 30 times normal. Most people cannot sustain more than one half to two thirds of these

values for longer than 1 minute.

 ALVEOLAR VENTILATION

The ultimate importance of pulmonary ventilation is to continually renew the air in the gas exchange areas of the lungs, where air is in proximity to the pulmonary blood.

These areas include the alveoli, alveolar sacs, alveolar ducts, and respiratory bronchioles. The rate at which new air reaches these areas is called *alveolar ventilation*.

**“DEAD SPACE” AND ITS EFFECT ON ALVEOLAR VENTILATION**

Some of the air a person breathes never reaches the gas exchange areas but simply fills respiratory passages where gas exchange does not occur, such as the nose, pharynx,

and trachea. This air is called *dead space air* because it is not useful for gas exchange.

On expiration, the air in the dead space is expired first, before any of the air from the alveoli reaches the atmosphere. Therefore, the dead space is very disadvantageous

for removing the expiratory gases from the lungs.

**Measurement of the Dead Space Volume.**

A simple method for measuring dead space volume **.** In making this measurement,

the subject suddenly takes a deep breath of 100 percent O2, which fills the entire dead space with pure O2. Some oxygen also mixes with the alveolar air but does not completely replace this air. Then the person expires through a rapidly recording nitrogen meter, which makes the record shown in the figure. The first portion of the expired air comes from the dead space regions of the respiratory passageways, where the air has been completely replaced by O2. Therefore, in the early part of the record, only O2 appears, and the nitrogen concentration is zero. Then, when alveolar air

begins to reach the nitrogen meter, the nitrogen concentration rises rapidly, because alveolar air containing large amounts of nitrogen begins to mix with the dead space air.

After still more air has been expired, all the dead space air has been washed from the passages and only alveolar air remains. Therefore, the recorded nitrogen concentration

reaches a plateau level equal to its concentration in the alveoli, as shown to the right in the figure. With a little thought, the student can see that the gray area represents

the air that has no nitrogen in it; this area is a measure of the volume of dead space air. For exact quantification, the following equation is used:

VD= Gray area. VE/ Pink area+ Gray area

where *VD* is dead space air and *VE* is the total volume of expired air.

Let us assume, for instance, that the gray area on the graph is 30 square centimeters, the pink area is 70 square centimeters, and the total volume expired is 500 milliliters.

The dead space would be

30/(30+70).500 = 150ml

**Normal Dead Space Volume.**

The normal dead space air in a young adult man is about 150 milliliters. Dead space

air increases slightly with age.

**Anatomical Versus Physiological Dead Space.**

The method just described for measuring the dead space measures the volume of all the space of the respiratory system other than the alveoli and their other closely related gas exchange areas; this space is called the *anatomic dead* *space*. On occasion, some of the alveoli are nonfunctional or only partially functional because of absent or poor blood flow through the adjacent pulmonary capillaries. Therefore, from a functional point of view, these alveoli must also be considered dead space. When the alveolar dead space is included in the total measurement of dead space, this is called the *physiological dead space*, in contradistinction to the anatomical dead space. In a normal person, the anatomical and physiological dead spaces are nearly equal

because all alveoli are functional in the normal lung, but in a person with partially functional or nonfunctional alveoli in some parts of the lungs, the physiological dead space may be as much as 10 times the volume of the anatomical dead space, or 1 to 2 liters. These problems are discussed further in Chapter 40 in relation to pulmonary gaseous exchange and in Chapter 43 in relation to certain pulmonary diseases.

**RATE OF ALVEOLAR VENTILATION**

Alveolar ventilation per minute is the total volume of new air entering the alveoli and adjacent gas exchange areas each minute. It is equal to the respiratory rate times the

amount of new air that enters these areas with each breath.

VA=Freq .(VT –VD)

where *V*\_*A* is the volume of alveolar ventilation per minute, *Freq* is the frequency of respiration per minute, *VT* is the tidal volume, and *VD* is the physiologic dead space volume. Thus, with a normal tidal volume of 500 milliliters, a normal dead space of 150 milliliters, and a respiratory rate of 12 breaths per minute, alveolar ventilation equals 12 × (500 − 150), or 4200 ml/min. Alveolar ventilation is one of the major factors determining the concentrations of oxygen and carbon dioxide in the alveoli.

**Functions of the Respiratory Passageways Trachea, Bronchi, and Bronchioles**

The air is distributed to the lungs by way of the trachea, bronchi, and bronchioles.

One of the most important challenges in the respiratory passageways is to keep them open and allow easy passage of air to and from the alveoli. To keep the trachea from

collapsing, multiple cartilage rings extend about five sixths of the way around the trachea. In the walls of the bronchi, less extensive curved cartilage plates also maintain a reasonable amount of rigidity yet allow sufficient motion for the lungs to expand and contract. These plates become progressively less extensive in the later generations of

bronchi and are gone in the bronchioles, which usually have diameters less than 1.5 millimeters. The bronchioles are not prevented from collapsing by the rigidity of their

walls. Instead, they are kept expanded mainly by the same transpulmonary pressures that expand the alveoli. That is, as the alveoli enlarge, the bronchioles also enlarge, but not as much.

**Muscular Wall of the Bronchi and Bronchioles and Its Control.**

In all areas of the *trachea* and *bronchi* not occupiedby cartilage plates, the walls are composed mainly ofsmooth muscle. Also, the walls of the *bronchioles* are

almost entirely smooth muscle, with the exception of the most terminal bronchiole, called the *respiratory bronchiole*, which is mainly pulmonary epithelium and underlying fibrous tissue plus a few smooth muscle fibers. Many obstructive diseases of the lung result from narrowing of the smaller bronchi and larger bronchioles, often because of excessive contraction of the smooth muscle.

**Resistance to Airflow in the Bronchial Tree.**

Under *normal respiratory conditions*, air flows through the respiratory passageways so easily that less than 1 centimeter of water pressure gradient from the alveoli to the atmosphere is sufficient to cause enough airflow for quiet breathing. The greatest amount of resistance to airflow occurs not in the minute air passages of the terminal bronchioles but in some of the larger bronchioles and bronchi near the trachea. The reason for this high resistance is that there are relatively few of these larger bronchi in comparison with the approximately 65,000 parallel terminal bronchioles, through each of which only a minute amount of air must pass. In some disease conditions, the smaller bronchioles play a far greater role in determining airflow resistance because

of their small size and because they are easily occluded by (1) muscle contraction in their walls, (2) edema occurring in the walls, or (3) mucus collecting in the lumens of the bronchioles.

**Nervous and Local Control of the Bronchiolar Musculature “Sympathetic” Dilation of the Bronchioles.**

Direct control of the bronchioles by sympathetic nerve fibers is relatively weak because few of these fibers penetrate to the central portions of the lung. However, the bronchial tree is very much exposed to *norepinephrine* and *epinephrine* released into the blood by sympathetic stimulation of the adrenal gland medullae. Both these hormones, especially epinephrine because of its greater stimulation of *beta-adrenergic receptors*, cause dilation of the bronchial tree.

**Parasympathetic Constriction of the Bronchioles.**

A few parasympathetic nerve fibers derived from the vagus nerves penetrate the lung parenchyma. These nerves secrete *acetylcholine* and, when activated, cause mild to

moderate constriction of the bronchioles. When a disease process such as asthma has already caused some bronchiolar constriction, superimposed parasympathetic nervous

stimulation often worsens the condition. When this situation occurs, administration of drugs that block the effects of acetylcholine, such as *atropine*, can sometimes relax the

respiratory passages enough to relieve the obstruction. Sometimes the parasympathetic nerves are also activated by reflexes that originate in the lungs. Most of these reflexes begin with irritation of the epithelial membrane of the respiratory passageways, initiated by noxious gases, dust, cigarette smoke, or bronchial infection. Also, a bronchiolar constrictor reflex often occurs when microemboli occlude small pulmonary arteries.

 **Local Secretory Factors May Cause Bronchiolar Constriction.**

Several substances formed in the lungs are often quite active in causing bronchiolar constriction. Two of the most important of these are *histamine* and *slow reactive*

*substance of anaphylaxis*. Both of these substances are released in the lung tissues by *mast cells* during allergic reactions, especially those caused by pollen in the air.

Therefore, they play key roles in causing the airway obstruction that occurs in allergic asthma; this is especially true of the slow reactive substance of anaphylaxis. The same irritants that cause parasympathetic constrictor reflexes of the airways—smoke, dust, sulfur dioxide, and some of the acidic elements in smog—may also act directly on the lung tissues to initiate local, non-nervous reactions that cause obstructive constriction of the airways.

**Mucus Lining the Respiratory Passageways, and Action of Cilia to Clear the Passageways**

All the respiratory passages, from the nose to the terminal bronchioles, are kept moist by a layer of mucus that coats the entire surface. The mucus is secreted partly by individual mucous goblet cells in the epithelial lining of the passages and partly by small submucosal glands. In addition to keeping the surfaces moist, the mucus traps small particles out of the inspired air and keeps most of these particles from ever reaching the alveoli. The mucus is removed from the passages in the following manner. The entire surface of the respiratory passages, both in the nose and in the lower passages down as far as the terminal bronchioles, is lined with ciliated epithelium, with about 200 cilia on each epithelial cell. These cilia beat continually

at a rate of 10 to 20 times per second by the mechanism explained in Chapter 2, and the direction of their “power stroke” is always toward the pharynx. That is, the

cilia in the lungs beat upward, whereas those in the nose beat downward. This continual beating causes the coat of mucus to flow slowly, at a velocity of a few millimeters per minute, toward the pharynx. Then the mucus and its entrapped particles are either swallowed or coughed to the exterior.

**Cough Reflex**

The bronchi and trachea are so sensitive to light touch that slight amounts of foreign matter or other causes of irritation initiate the cough reflex. The larynx and carina (i.e.,

the point where the trachea divides into the bronchi) are especially sensitive, and the terminal bronchioles and even the alveoli are sensitive to corrosive chemical stimuli

such as sulfur dioxide gas or chlorine gas. Afferent nerve impulses pass from the respiratory passages mainly through the vagus nerves to the medulla of the brain. There, an automatic sequence of events is triggered by the neuronal circuits of the medulla, causing the following effect. First, up to 2.5 liters of air are rapidly inspired. Second, the epiglottis closes, and the vocal cords shut tightly to entrap the air within the lungs. Third, the abdominal muscles contract forcefully, pushing against the diaphragm while other expiratory muscles, such as the internal intercostals,

also contract forcefully. Consequently, the pressure in the lungs rises rapidly to as much as 100 mm Hg or more. Fourth, the vocal cords and the epiglottis suddenly

open widely, so that air under this high pressure in the lungs *explodes* outward. Indeed, sometimes this air is expelled at velocities ranging from 75 to 100 miles per hour.

Importantly, the strong compression of the lungs collapses the bronchi and trachea by causing their noncartilaginous parts to invaginate inward, so the exploding air actually

passes through *bronchial* and *tracheal slits*. The rapidly moving air usually carries with it any foreign matter that is present in the bronchi or trachea.

**Sneeze Reflex**

The sneeze reflex is very much like the cough reflex, except that it applies to the nasal passageways instead of the lower respiratory passages. The initiating stimulus of the sneeze reflex is irritation in the nasal passageways; the afferent impulses pass in the fifth cranial nerve to the medulla, where the reflex is triggered. A series of reactions similar to those for the cough reflex takes place, but the uvula is depressed, so large amounts of air pass rapidly through the nose, thus helping to clear the nasal passages of foreign matter.

**Normal Respiratory Functions of the Nose**

As air passes through the nose, three distinct normal respiratory functions are performed by the nasal cavities: (1) the air is *warmed* by the extensive surfaces of the conchae and septum, a total area of about 160 square centimeters ; (2) the air is *almost completely humidified* even before it passes beyond the nose; and (3) the air is 506

*partially filtered*. These functions together are called the *air conditioning function* of the upper respiratory passageways.Ordinarily, the temperature of the inspired air rises towithin 1°F of body temperature and to within 2 to 3 percentof full saturation with water vapor before it reaches thetrachea. When a person breathes air through a tube directlyinto the trachea (as through a tracheostomy), the coolingand especially the drying effect in the lower lung can leadto serious lung crusting and infection.

**Filtration Function of the Nose**

The hairs at the entrance to the nostrils are important for filtering out large particles. Much more important, though, is the removal of particles by *turbulent precipitation*. That is, the air passing through the nasal passageways hits many obstructing vanes: the *conchae* (also called *turbinates*, because they cause turbulence of the air); the septum; and the pharyngeal wall. Each time air hits one of these obstructions, it must change its direction of movement. The particles suspended in the air, having far more mass and momentum than air, cannot change their direction of travel as rapidly as the air can. Therefore, they continue forward, striking the surfaces of the obstructions, and are entrapped in the mucous coating and transported by the cilia to the pharynx to be swallowed.

**Size of Particles Entrapped in the Respiratory Passages.**

The nasal turbulence mechanism for removing particles from air is so effective that almost no particles larger than 6 micrometers in diameter enter the lungs through the

nose. This size is smaller than the size of red blood cells. Of the remaining particles, many that are between 1 and 5 micrometers *settle* in the smaller bronchioles as a result

of *gravitational precipitation*. For instance, terminal bronchiolar disease is common in coal miners because of settled dust particles. Some of the still smaller particles (smaller than 1 micrometer in diameter) *diffuse* against the walls of the alveoli and adhere to the alveolar fluid. However, many particles smaller than 0.5 micrometer in diameter remain suspended in the alveolar air and are expelled by expiration. For instance, the particles of cigarette smoke are about 0.3 micrometer. Almost none of these particles are precipitated in the respiratory passageways before they reach the alveoli. Unfortunately, up to one third of them do precipitate in the alveoli by the diffusion process, with the balance remaining suspended and expelled in the expired air. Many of the particles that become entrapped in the alveoli are removed by *alveolar macrophages*, and others are carried away by the lung lymphatics. An excess of particles can cause growth of fibrous tissue in the alveolar septa, leading to permanent

debility.

**Vocalization**

Speech involves not only the respiratory system but also (1) specific speech nervous control centers in the cerebral cortex, which are discussed in Chapter 58; (2) respiratory control centers of the brain; and (3) the articulation and resonance structures of the mouth and nasal cavities. Speech is composed of two mechanical functions: (1) *phonation*, which is achieved by the larynx, and (2) *articulation*,

which is achieved by the structures of the mouth

 **Phonation.** The larynx, is especially adapted to act as a vibrator. The vibrating elements are the *vocal folds*, commonly called the *vocal cords*. The vocal cords protrude from the lateral walls of the larynx toward the center of the glottis; they are stretched and positioned by several specific muscles of the larynx itself.

The vocal cords are seen when looking into the glottis with a laryngoscope. During

normal breathing, the cords are wide open to allow easy passage of air. During phonation, the cords move together so that passage of air between them will cause vibration. The pitch of the vibration is determined mainly by the degree of stretch of the cords, but also by how tightly the cords are approximated to one another and by the mass of their edges.

Immediately inside each cord is a strong elastic ligament called the *vocal ligament*. This ligament is attached anteriorly to the large *thyroid cartilage*, which is the cartilage that projects forward from the anterior surface of the neck and is called

the “Adam’s apple.” Posteriorly, the vocal ligament is attached to the *vocal processes* of two *arytenoid cartilages*. The thyroid cartilage and the arytenoid cartilages articulate from below with another cartilage the *cricoid cartilage*. The vocal cords can be stretched by either forward rotation of the thyroid cartilage or posterior rotation of the arytenoid cartilages, activated by muscles stretching from the thyroid cartilage and arytenoid cartilages to the cricoid cartilage. Muscles located within the vocal cords lateral to the vocal ligaments, the thyroarytenoid muscles, can pull the arytenoid cartilages toward the thyroid cartilage and, therefore, loosen the vocal cords. Also, slips of these muscles *within* the vocal cords can change the *shapes and masses of*

*the vocal cord edges*, sharpening them to emit high-pitched sounds and blunting them for the more bass sounds. Several other sets of small laryngeal muscles lie between

the arytenoid cartilages and the cricoid cartilage and can rotate these cartilages inward or outward or pull their bases together or apart to give the various configurations of the

vocal cords **Articulation and Resonance.** The three major organs of articulation are the *lips, tongue*, and *soft palate*. They need not be discussed in detail because we are all familiar with their movements during speech and other vocalizations. The resonators include the *mouth*, the *nose* and *associated* *nasal sinuses*, the *pharynx*, and even the *chest cavity*.

Again, we are all familiar with the resonating qualities of these structures. For instance, the function of the nasal resonators is demonstrated by the change in voice quality when a person has a severe cold that blocks the air passages to these resonators.