



#### **Exchanging Materials**

- Every organism must exchange materials with its environment.
  - This exchange ultimately occurs at the cellular level.



# Circulatory Systems Reflect Phylogeny

• Transport systems functionally connect the organs of exchange with the body cells.

Solubility of Gases in Distilled Water							
°C	Oxygen	Carbon Dioxide	Nitrogen	Helium			
0	21.7	767.5					
10	16.9	531.2	_				
20	13.7	386.8	6.82				
30	11.6	294.9	_				
37	10.6	250.5	5.61	3 75			
40	10.2	234.8		5.75			

°C	Salinity	Office	10%	20%	1011	
1,25			THE REAL OWN		30 %	40%a
0		21.7	20.2	18.9	17.7	16.6
10		16.9	15.8	14.8	13.9	13.1
20		13.7	12.9	12.2	11.5	10.8
30		11.6	11.0	10.4	9.86	9.31
40		10.2	9.71	9.26	8.73	8 24

Altitude	P.	Ambient nO.	p.O.	p <sub>A</sub> CO <sub>2</sub>
	~ 0	rimorene poz	PA-2	
0	101	21.1	13.8	5.3
3100	70.6	14.6	8.9	4.8
4340	61.9	12.8	6.0	
6200	46	9.7	5.3	3.2
7100		normal "cei	ling'' V	
8848	33	6.9	4.0	1.5
9200	30	6.3	2.8	-
12300	19	3.9	1.1	-
14460	punning	"ceiling" with	pure O <sub>2</sub>	
15400	12	2.4	0.1	-
20000	6	1.3	0	0

liter <sup>-1</sup> kPa <sup>-1</sup> disso in equilibrium with as a function of de	blved $N_2$ ) for h the ambi epth of div	or a hum ent hydr ing.	an scuba ostatic p	diver
Partial pressures (	(kPa) for o	xygen an	and nitrogenerative colar card	en (dry,
$CO_2$ -free values for	or ambient	air), alvo	$I O_2 \text{ per } I$	bon
dioxide, and plasm	na-dissolve	ed $O_2$ (mi	$O_2 \text{ is } O_2$	iter
plasma; assuming	plasma so	lubility o	$N_2 \text{ per } I$	209 ml
liter <sup>-1</sup> kPa <sup>-1</sup> ) and	fat N <sub>2</sub> con	tent (ml	$N_2 \text{ per } I$	ter
body fat; assuming	g fat solub	ility of N	$N_2 \text{ is } 0.67$	ml

Ambient Pressure	101	202	1111	5151
pO <sub>2</sub>	21.1	42.4	233.5	1082.6
pN <sub>2</sub>	79.8	159.7	878.3	4072.5
Alveolar pCO <sub>2</sub>	5.32	5.32	5.32	5.32
Plasma O <sub>2</sub>	4.4	8.8	48.3	223.9
Fat N <sub>2</sub>	53	106	582	2700

#### Sea level

 $P_{O2} = 760 * 0.2094 = 159 \text{ mmHg}$ 

 $P_{O2} = (760-18)*0.2094 = 155 \text{ mmHg}$ 

#### The Respiratory System

- Major function-respiration
  - Supply body with O<sub>2</sub> for cellular respiration; dispose of CO<sub>2</sub>, a waste product of cellular respiration
  - Its four processes involve both respiratory and circulatory systems
- · Also functions in olfaction and speech

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#### **Processes of Respiration**

- Pulmonary ventilation (breathing)movement of air into and out of lungs
- External respiration-O<sub>2</sub> and CO<sub>2</sub> exchange between lungs and blood
- Transport-O<sub>2</sub> and CO<sub>2</sub> in blood
- Internal respiration-O<sub>2</sub> and CO<sub>2</sub> exchange between systemic blood vessels and tissues

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system

#### **Bronchi and Subdivisions**

- Air passages undergo 23 orders of branching → bronchial (respiratory) tree
- From tips of bronchial tree → conducting zone structures → respiratory zone structures

#### **Conducting Zone Structures**

- Trachea → right and left main (primary) bronchi
- Each main bronchus enters hilum of one lung
  - Right main bronchus wider, shorter, more vertical than left
- Each main bronchus branches into lobar (secondary) bronchi (three on right, two on left)
  - Each lobar bronchus supplies one lobe

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Figure 22.7 Conducting zone passages

#### **Conducting Zone Structures**

- Each lobar bronchus branches into segmental (tertiary) bronchi
  - Segmental bronchi divide repeatedly
- Branches become smaller and smaller →
  - Bronchioles-less than 1 mm in diameter
  - Terminal bronchioles-smallest-less than 0.5 mm diameter

Superior lobe of right lung birding lobe of right lung **Conducting Zone Structures** 

- From bronchi through bronchioles, structural changes occur
  - Cartilage rings become irregular plates; in bronchioles elastic fibers replace cartilage
  - Epithelium changes from pseudostratified columnar to cuboidal; cilia and goblet cells become sparse
  - Relative amount of smooth muscle increases
     Allows constriction

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Figure 22.8a Respiratory zone structures

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#### **Respiratory Zone**

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- Begins as terminal bronchioles → respiratory bronchioles → alveolar ducts → alveolar sacs
  - Alveolar sacs contain clusters of alveoli
    - ~300 million alveoli make up most of lung volume
    - · Sites of gas exchange

Aveolar duct reminate bronchides tronchides tronch

Figure 22.8b Respiratory zone structures.



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#### **Respiratory Membrane**

- Alveolar and capillary walls and their fused basement membranes
  - ~0.5-µm-thick; gas exchange across membrane by simple diffusion
- · Alveolar walls
  - Single layer of squamous epithelium (type I alveolar cells)
- Scattered cuboidal type II alveolar cells secrete surfactant and antimicrobial proteins

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(a) Diagrammatic view of capillary-alveoli relationships

Figure 22.9b Alveoli and the respiratory memb



(b) Scanning electron micrograph of pulmonary capillary casts (70x) 9 2013 Feason Educator, Inc.

#### Alveoli

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- Surrounded by fine elastic fibers and pulmonary capillaries
- Alveolar pores connect adjacent alveoli
   Equalize air pressure throughout lung
- Alveolar macrophages keep alveolar surfaces sterile
  - 2 million dead macrophages/hour carried by cilia  $\rightarrow$  throat  $\rightarrow$  swallowed

#### Figure 22.9c Alveoli and the respiratory membrane.



(c) Detailed anatomy of the respiratory membrane

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#### Lungs

- · Apex-superior tip; deep to clavicle
- · Base-inferior surface; rests on diaphragm
- **Hilum**-on mediastinal surface; site for entry/exit of blood vessels, bronchi, lymphatic vessels, and nerves
- Left lung smaller than right
  - Cardiac notch-concavity for heart
  - Separated into *superior* and *inferior* **lobes** by *oblique fissure*

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(a) Anterior view. The lungs flank mediastinal structures laterally.

#### **Mechanics of Breathing**

- Pulmonary ventilation consists of two phases
  - Inspiration-gases flow into lungs
  - Expiration-gases exit lungs

## Pressure Relationships in the Thoracic Cavity

- Atmospheric pressure (P<sub>atm</sub>)
  - Pressure exerted by air surrounding body
  - 760 mm Hg at sea level = 1 atmosphere
- Respiratory pressures described relative to  $\mathsf{P}_{\mathsf{atm}}$ 
  - Negative respiratory pressure-less than Patm
  - Positive respiratory pressure-greater than Patm
  - Zero respiratory pressure = P<sub>atm</sub>

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#### **Intrapulmonary Pressure**

- Intrapulmonary (intra-alveolar) pressure (P<sub>pul</sub>)
  - Pressure in alveoli
  - Fluctuates with breathing
  - Always eventually equalizes with Patm

#### **Intrapleural Pressure**

- Intrapleural pressure (P<sub>ip</sub>)
  - Pressure in pleural cavity
  - Fluctuates with breathing
  - Always a negative pressure (<P<sub>atm</sub> and <P<sub>pul</sub>)
  - Fluid level must be minimal
    - Pumped out by lymphatics
    - If accumulates  $\rightarrow$  positive  $\mathsf{P}_{\mathsf{ip}}$  pressure  $\rightarrow$  lung collapse

#### **Intrapleural Pressure**

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#### Negative P<sub>ip</sub> caused by opposing forces

- Two inward forces promote lung collapse
  - · Elastic recoil of lungs decreases lung size
  - · Surface tension of alveolar fluid reduces alveolar size
- One outward force tends to enlarge lungs
  - · Elasticity of chest wall pulls thorax outward

#### **Pressure Relationships**

- If  $P_{ip} = P_{pul}$  or  $P_{atm} \rightarrow$  lungs collapse
- (P<sub>pul</sub> P<sub>ip</sub>) = transpulmonary pressure - Keeps airways open
  - Greater transpulmonary pressure → larger lungs



#### Homeostatic Imbalance

- · Atelectasis (lung collapse) due to
  - Plugged bronchioles  $\rightarrow$  collapse of alveoli
  - Pneumothorax-air in pleural cavity
    - · From either wound in parietal or rupture of visceral pleura
    - · Treated by removing air with chest tubes; pleurae heal → lung reinflates

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#### **Pulmonary Ventilation**

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- Inspiration and expiration
- · Mechanical processes that depend on volume changes in thoracic cavity
  - Volume changes  $\rightarrow$  pressure changes
  - Pressure changes  $\rightarrow$  gases flow to equalize pressure

#### **Boyle's Law**

- · Relationship between pressure and volume of a gas
  - Gases fill container; if container size reduced → increased pressure
- Pressure (P) varies inversely with volume (V):

 $-P_1V_1 = P_2V_2$ 

#### Inspiration

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- · Active process
  - Inspiratory muscles (diaphragm and external intercostals) contract
  - Thoracic volume increases → intrapulmonary pressure drops (to –1 mm Hg)
  - Lungs stretched and intrapulmonary volume increases
  - Air flows into lungs, down its pressure gradient, until  $P_{pul} = P_{atm}$

#### **Forced Inspiration**

 Vigorous exercise, COPD → accessory muscles (scalenes, sternocleidomastoid, pectoralis minor) → further increase in thoracic cage size

Figure 22.13 Changes in thoracic volume and sequence of events during inspiration and expiration. (1 of 2) Slide



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#### Expiration

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- Quiet expiration normally passive process
   Inspiratory muscles relax

  - Thoracic cavity volume decreases
  - Elastic lungs recoil and intrapulmonary volume decreases → pressure increases (P<sub>pul</sub> rises to +1 mm Hg) →
  - Air flows out of lungs down its pressure gradient until  $P_{pul} = 0$
- Note: forced expiration-active process; uses abdominal (oblique and transverse) and internal intercostal muscles

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Figure 22.13 Changes in thoracic volume and sequence of events during inspiration and expiration. (2 of 2) Slide 1



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Figure 22.14 Changes in intrapulmonary and intrapleural pressures during inspiration and expiration

## Physical Factors Influencing Pulmonary Ventilation

- Three physical factors influence the ease of air passage and the amount of energy required for ventilation.
  - Airway resistance
  - Alveolar surface tension
  - Lung compliance

#### **Airway Resistance**

- Friction-major nonelastic source of resistance to gas flow; occurs in airways
- Relationship between flow (F), pressure (P), and resistance (R) is:

$$F = \frac{\Delta P}{R}$$

- $-\Delta P$  pressure gradient between atmosphere and alveoli (2 mm Hg or less during normal quiet breathing)
- Gas flow changes inversely with resistance

#### **Airway Resistance**

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- · Resistance usually insignificant
  - Large airway diameters in first part of conducting zone
  - Progressive branching of airways as get smaller, increasing total cross-sectional area
  - Resistance greatest in medium-sized bronchi
- Resistance disappears at terminal bronchioles where diffusion drives gas movement

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#### **Homeostatic Imbalance**

- As airway resistance rises, breathing movements become more strenuous
- Severe constriction or obstruction of bronchioles
  - Can prevent life-sustaining ventilation
  - Can occur during acute asthma attacks; stops ventilation
- Epinephrine dilates bronchioles, reduces air resistance

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#### **Alveolar Surface Tension**

#### Surface tension

- Attracts liquid molecules to one another at gas-liquid interface
- Resists any force that tends to increase surface area of liquid
- Water–high surface tension; coats alveolar walls → reduces them to smallest size

#### **Alveolar Surface Tension**

- Surfactant
  - Detergent-like lipid and protein complex produced by type II alveolar cells
  - Reduces surface tension of alveolar fluid and discourages alveolar collapse
  - Insufficient quantity in premature infants causes infant respiratory distress syndrome
    - $\boldsymbol{\cdot} \rightarrow \text{alveoli collapse after each breath}$

#### Lung Compliance

- Measure of change in lung volume that occurs with given change in transpulmonary pressure
- Higher lung compliance → easier to expand lungs
- Normally high due to
  - Distensibility of lung tissue
  - Surfactant, which decreases alveolar surface tension

#### Lung Compliance

• Diminished by

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- Nonelastic scar tissue replacing lung tissue (fibrosis)
- Reduced production of surfactant
- Decreased flexibility of thoracic cage

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#### **Total Respiratory Compliance**

- The total compliance of the respiratory system is also influenced by compliance (distensibility) of the thoracic wall, which is decreased by:
  - Deformities of thorax
  - Ossification of costal cartilage
  - Paralysis of intercostal muscles

#### **Respiratory Volumes**

- Used to assess respiratory status – Tidal volume (TV)

  - Inspiratory reserve volume (IRV)
  - Expiratory reserve volume (ERV)
  - Residual volume (RV)

Figure 22.16b Respiratory volumes and capacities.

	Measurement	Adult male average value	Adult female average value	Description
ſ	Tidal volume (TV)	500 ml	500 ml	Amount of air inhaled or exhaled with each breath under resting conditions
Baurinten	Inspiratory reserve volume (IRV)	3100 ml	1900 ml	Amount of air that can be forcefully inhaled after a normal tidal volume inspiration
volumes	Expiratory reserve volume (ERV)	1200 ml	700 ml	Amount of air that can be forcefully exhaled after a normal tidal volume expiration
	Residual volume (RV)	1200 ml	1100 ml	Amount of air remaining in the lungs after a forced expiration
	Total lung capacity (TLC	) 6000 ml	4200 ml	Maximum amount of air contained in lungs after a maximum inspiratory effort: TLC = TV + IRV + ERV + RV
Bausinstan	Vital capacity (VC)	4800 ml	3100 ml	Maximum amount of air that can be expired after a maximum inspiratory effort: VC = TV + IRV + ERV
capacities	Inspiratory capacity (IC)	3600 ml	2400 ml	Maximum amount of air that can be inspired after a normal tidal volume expiration: IC = TV + IRV
	Functional residual capacity (FRC)	2400 ml	1800 ml	Volume of air remaining in the lungs after a normal tidal volume expiration: FRC = ERV + RV

(b) Summary of respiratory volumes and capacities for males and females

Figure 22.16a Respiratory volumes and capacities.



<sup>(</sup>a) Spirographic record for a male

#### **Dead Space**

- Anatomical dead space

   No contribution to gas exchange
   Air remaining in passageways; ~150 ml
- Alveolar dead space-non-functional alveoli due to collapse or obstruction
- Total dead space-sum of anatomical and alveolar dead space

#### **Pulmonary Function Tests**

- **Spirometer**-instrument for measuring respiratory volumes and capacities
- · Spirometry can distinguish between
  - Obstructive pulmonary disease—increased airway resistance (e.g., bronchitis)
     TLC, FRC, RV may increase
  - Restrictive disorders—reduced TLC due to disease or fibrosis
    - VC, TLC, FRC, RV decline

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#### **Pulmonary Function Tests**

- To measure *rate* of gas movement
  - Forced vital capacity (FVC)—gas forcibly expelled after taking deep breath
  - Forced expiratory volume (FEV)—amount of gas expelled during specific time intervals of FVC

#### **Alveolar Ventilation**

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- Minute ventilation—total amount of gas flow into or out of respiratory tract in one minute
  - Normal at rest = ~ 6 L/min
  - Normal with exercise = up to 200 L/min
  - Only rough estimate of respiratory efficiency

Table 22.2 Effects of Breathing Rate and Depth on Alveolar ventilation of Three Hypothetical Patients

	Breathing Rate					
BREATHING PATTERN OF HYPOTHETICAL PATIENT	DEAD SPACE VOLUME (DSV)	TIDAL VOLUME (TV)	RESPIRATORY RATE*	MINUTE VENTILATION (MVR)	ALVEOLAR VENTILATION (AVR)	% EFFECTIVE VENTILATION (AVR/MVR)
I-Normal rate and depth	150 ml	500 ml	20/min	10,000 mVmin	7000 ml/min	70%
II-Slow, deep breathing	150 ml	1000 ml	10/min	10,000 ml/min	8500 ml/min	85%
III-Rapid, shallow breathing	150 ml	250 ml	40/min	10,000 ml/min	4000 ml/min	40%

\*Respiratory rate values are artificially adjusted to provide equivalent minute ventilation as a baseline for comparing alveolar ventilation

### Gas Exchanges Between Blood, Lungs, and Tissues

- External respiration-diffusion of gases in lungs
- Internal respiration-diffusion of gases at body tissues
- · Both involve

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- Physical properties of gases
- Composition of alveolar gas

### Basic Properties of Gases: Dalton's Law of Partial Pressures

• Total pressure exerted by mixture of gases = sum of pressures exerted by each gas

#### · Partial pressure

- Pressure exerted by each gas in mixture
- Directly proportional to its percentage in mixture

#### **Basic Properties of Gases: Henry's Law**

- · Gas mixtures in contact with liquid
  - Each gas dissolves in proportion to its partial pressure
  - At equilibrium, partial pressures in two phases will be equal
  - Amount of each gas that will dissolve depends on
    - Solubility–CO $_2$  20 times more soluble in water than O $_2;$  little N $_2$  dissolves in water
    - Temperature-as temperature rises, solubility decreases

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**Composition of Alveolar Gas** 

- Alveoli contain more CO<sub>2</sub> and water vapor than atmospheric air
  - Gas exchanges in lungs
  - Humidification of air
  - Mixing of alveolar gas with each breath

Table 22.4	and in the Alveoli	al Pressures and Approximate	Percentages in the Atmos		
	ATMOSPHE	RE (SEA LEVEL)	ALVEOLI		
GAS	APPROXIMATE PERCENTAGE	PARTIAL PRESSURE (mm Hg)	APPROXIMATE PERCENTAGE	PARTIAL PRESSURE (mm Hg)	
N2	78.6	597	74.9	569	
01	20.9	159	13.7	104	
CO2	0.04	0.3	5.2	40	
H <sub>2</sub> O	0.46	3.7	6.2	47	
	100.0%	760	100.0%	760	

Table 22.4 Comparison of Gas Partial Pressures and Approximate Percentages in the Atmosphere and in the Alveoli

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#### **External Respiration**

- Exchange of O<sub>2</sub> and CO<sub>2</sub> across respiratory membrane
- · Influenced by

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- Thickness and surface area of respiratory membrane
- Partial pressure gradients and gas solubilities
- Ventilation-perfusion coupling

### Thickness and Surface Area of the Respiratory Membrane

- Respiratory membranes
  - 0.5 to 1  $\mu$  m thick
  - Large total surface area (40 times that of skin) for gas exchange
- Thicken if lungs become waterlogged and edematous → gas exchange inadequate
- Reduced surface area in emphysema (walls of adjacent alveoli break down), tumors, inflammation, mucus

#### **Partial Pressure Gradients and Gas Solubilities**

- Steep partial pressure gradient for O<sub>2</sub> in lungs
  - Venous blood  $Po_2 = 40 \text{ mm Hg}$
  - Alveolar Po<sub>2</sub> = 104 mm Hg
    - · Drives oxygen flow to blood
    - · Equilibrium reached across respiratory membrane in ~0.25 seconds, about 1/3 time a red blood cell in pulmonary capillary →
      - Adequate oxygenation even if blood flow increases 3X

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**Partial Pressure Gradients and Gas Solubilities** 

- Partial pressure gradient for CO<sub>2</sub> in lungs less steep
  - Venous blood  $Pco_2 = 45 \text{ mm Hg}$
  - Alveolar  $Pco_2 = 40 \text{ mm Hg}$
- Though gradient not as steep, CO<sub>2</sub> diffuses in equal amounts with oxygen
  - CO<sub>2</sub> 20 times more soluble in plasma than oxygen



**Ventilation-Perfusion Coupling** 

- Perfusion-blood flow reaching alveoli
- · Ventilation-amount of gas reaching alveoli
- Ventilation and perfusion matched (coupled) for efficient gas exchange
  - Never balanced for all alveoli due to
    - · Regional variations due to effect of gravity on blood and air flow
    - · Some alveolar ducts plugged with mucus

#### **Ventilation-Perfusion Coupling**

Perfusion

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- Changes in Po2 in alveoli cause changes in diameters of arterioles
  - Where alveolar O22 is high, arterioles dilate
  - · Where alveolar O2 is low, arterioles constrict
  - · Directs most blood where alveolar oxygen high

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#### Ventilation-Perfusion Coupling

- Changes in Pco<sub>2</sub> in alveoli cause changes in diameters of bronchioles
  - Where alveolar CO2 is high, bronchioles dilate
  - Where alveolar  $\text{CO}_2$  is low, bronchioles constrict
  - Allows elimination of CO<sub>2</sub> more rapidly

Figure 22.19 Ventilation-perfusion coupling.



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#### Transport of Respiratory Gases by Blood

• Oxygen (O<sub>2</sub>) transport

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Carbon dioxide (CO<sub>2</sub>) transport

#### O<sub>2</sub> Transport

- Molecular O<sub>2</sub> carried in blood
  - 1.5% dissolved in plasma
  - 98.5% loosely bound to each Fe of hemoglobin (Hb) in RBCs
     4 O<sub>2</sub> per Hb

O<sub>2</sub> and Hemoglobin

- **Oxyhemoglobin** (HbO<sub>2</sub>)-hemoglobin-O<sub>2</sub> combination
- Reduced hemoglobin (deoxyhemoglobin) (HHb)-hemoglobin that has released O<sub>2</sub>

 $\begin{array}{c} \text{Lungs} \\ \text{HHb} + \text{O}_2 & \longleftrightarrow & \text{HbO}_2 + \text{H}^+ \\ & \text{Tissues} \end{array}$ 

### O<sub>2</sub> and Hemoglobin

- Loading and unloading of  $\mathrm{O}_{\mathrm{2}}$  facilitated by change in shape of Hb
  - As  $O_2$  binds, Hb affinity for  $O_2$  increases
  - As  $\mathrm{O}_2$  is released, Hb affinity for  $\mathrm{O}_2$  decreases
- Fully saturated (100%) if all four heme groups carry O<sub>2</sub>
- Partially saturated when one to three hemes carry O<sub>2</sub>

Figure 22.20 The amount of oxygen locally. (1 of 3) 89 88 22 88 40 60 PO<sub>2</sub> (mm Hg) © 2013 Pearson Education, In

in depends on the Po, (the amount of oxygen) available

#### Influence of Po<sub>2</sub> on Hemoglobin Saturation

- · In arterial blood
  - $-Po_2 = 100 \text{ mm Hg}$
  - Contains 20 ml oxygen per 100 ml blood (20 vol %)
  - Hb is 98% saturated
- Further increases in Po<sub>2</sub> (e.g., breathing deeply) produce minimal increases in O<sub>2</sub> binding

#### Influence of Po<sub>2</sub> on Hemoglobin Saturation

- · In venous blood
  - $-Po_2 = 40 \text{ mm Hg}$
  - Contains 15 vol % oxygen
  - Hb is 75% saturated
  - Venous reserve

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· Oxygen remaining in venous blood

Figure 22.20 The locally. (2 of 3)



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Figure 22.21 Effect of temperat

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#### **Other Factors Influencing Hemoglobin Saturation**

- Increases in temperature, H+, Pco<sub>2</sub>, and BPG
  - Modify structure of hemoglobin; decrease its affinity for O<sub>2</sub>
  - Occur in systemic capillaries
  - Enhance O<sub>2</sub> unloading from blood
  - Shift O2-hemoglobin dissociation curve to right
- · Decreases in these factors shift curve to left
  - Decreases oxygen unloading from blood

hemoglobin dissociation cu 40 Po 6

## Factors that Increase Release of $\rm O_2$ by Hemoglobin

- As cells metabolize glucose and use O<sub>2</sub> – Pco<sub>2</sub> and H<sup>+</sup> increase in capillary blood →
  - Declining blood pH and increasing  $Pco_2 \rightarrow$ 
    - Bohr effect Hb-O₂ bond weakens → oxygen unloading where needed most
  - Heat production increases → directly and indirectly decreases Hb affinity for O<sub>2</sub> → increased oxygen unloading to active tissues

#### **Homeostatic Imbalance**

• Hypoxia

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- Inadequate O<sub>2</sub> delivery to tissues → cyanosis
- Anemic hypoxia–too few RBCs; abnormal or too little Hb
- Ischemic hypoxia-impaired/blocked circulation
- Histotoxic hypoxia–cells unable to use O<sub>2</sub>, as in metabolic poisons
- Hypoxemic hypoxia–abnormal ventilation; pulmonary disease
- Carbon monoxide poisoning–especially from fire; 200X greater affinity for Hb than oxygen

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#### CO<sub>2</sub> Transport

- CO<sub>2</sub> transported in blood in three forms - 7 to 10% dissolved in plasma
  - 20% bound to *globin* of hemoglobin (carbaminohemoglobin)
  - 70% transported as bicarbonate ions (HCO<sub>3</sub><sup>-</sup>) in plasma

#### Transport and Exchange of CO<sub>2</sub>

• CO<sub>2</sub> combines with water to form carbonic acid (H<sub>2</sub>CO<sub>3</sub>), which quickly dissociates

$CO_2 +$	$-H_2O \equiv$	$\Longrightarrow$ H <sub>2</sub> CO <sub>3</sub> $\equiv$	$\rightarrow$ H <sup>+</sup>	+ HCO <sub>3</sub> <sup>-</sup>
carbon	water	carbonic	hydrogen	bicarbonate
dioxide		acid	ion	ion

 Occurs primarily in RBCs, where carbonic anhydrase reversibly and rapidly catalyzes reaction

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#### Transport and Exchange of CO<sub>2</sub>

- In systemic capillaries
  - HCO<sub>3</sub><sup>–</sup> quickly diffuses from RBCs into plasma
    - · Chloride shift occurs
      - Outrush of HCO<sub>3</sub><sup>−</sup> from RBCs balanced as CI<sup>−</sup> moves into RBCs from plasma

Figure 22.22a Transport and exchange of  $CO_2$  and  $O_2$ 



(a) Oxygen release and carbon dioxide pickup at the tissues

#### Transport and Exchange of CO<sub>2</sub>

- · In pulmonary capillaries
  - HCO<sub>3</sub><sup>-</sup> moves into RBCs (while Cl<sup>-</sup> move out); binds with H<sup>+</sup> to form H<sub>2</sub>CO<sub>3</sub>
  - $\rm H_2CO_3$  split by carbonic anhydrase into  $\rm CO_2$  and water
  - CO2 diffuses into alveoli

Figure 22.22b Transport and exchange of CO<sub>2</sub> and O<sub>2</sub>



(b) Oxygen pickup and carbon dioxide release in the lungs

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#### **Control of Respiration**

- Involves higher brain centers, chemoreceptors, and other reflexes
- · Neural controls

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- Neurons in reticular formation of medulla and pons
- Clustered neurons in medulla important
  - Ventral respiratory group
  - Dorsal respiratory group

#### **Medullary Respiratory Centers**

- Ventral respiratory group (VRG)
  - Rhythm-generating and integrative center
  - Sets eupnea (12–15 breaths/minute)
     Normal respiratory rate and rhythm
  - Its inspiratory neurons excite inspiratory muscles via phrenic (diaphragm) and intercostal nerves (external intercostals)
  - Expiratory neurons inhibit inspiratory neurons

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#### **Medullary Respiratory Centers**

- Dorsal respiratory group (DRG)
  - Near root of cranial nerve IX
  - Integrates input from peripheral stretch and chemoreceptors; sends information → VRG



#### **Generation of the Respiratory Rhythm**

· Not well understood

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- One hypothesis – Pacemaker neurons with intrinsic rhythmicity
- Most widely accepted hypothesis
  - Reciprocal inhibition of two sets of interconnected pacemaker neurons in medulla that generate rhythm



Figure 22.26 Location and innervation of the peripheral chemoreceptors in the carotid and aortic bodies.



#### **Inflation Reflex**

- Hering-Breuer Reflex (inflation reflex)
  - Stretch receptors in pleurae and airways stimulated by lung inflation
    - Inhibitory signals to medullary respiratory centers end inhalation and allow expiration
    - Acts as protective response more than normal regulatory mechanism

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#### **Respiratory Adjustments: Exercise**

- Adjustments geared to both intensity and duration of exercise
- Hyperpnea

- Increased ventilation (10 to 20 fold) in response to metabolic needs
- Pco<sub>2</sub>, Po<sub>2</sub>, and pH remain surprisingly constant during exercise

#### **Respiratory Adjustments: Exercise**

- Three neural factors cause increase in ventilation as exercise begins
  - Psychological stimuli-anticipation of exercise
  - Simultaneous cortical motor activation of skeletal muscles and respiratory centers
  - Excitatory impulses to respiratory centers from proprioceptors in moving muscles, tendons, joints

#### **Respiratory Adjustments: High Altitude**

- Quick travel to altitudes above 2400 meters (8000 feet) may → symptoms of acute mountain sickness (AMS)
  - Atmospheric pressure and Po<sub>2</sub> levels lower
  - Headaches, shortness of breath, nausea, and dizziness
  - In severe cases, lethal cerebral and pulmonary edema

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#### Acclimatization to High Altitude

- Acclimatization—respiratory and hematopoietic adjustments to long-term move to high altitude
  - Chemoreceptors become more responsive to  $Pco_2$  when  $Po_2$  declines
  - Substantial decline in Po<sub>2</sub> directly stimulates peripheral chemoreceptors
  - Result—minute ventilation increases and stabilizes in few days to 2–3 L/min higher than at sea level

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#### **Homeostatic Imbalances**

- Chronic obstructive pulmonary disease (COPD)
  - Exemplified by chronic bronchitis and emphysema
  - Irreversible decrease in ability to force air out of lungs
  - Other common features
    - History of smoking in 80% of patients
    - Dyspnea labored breathing ("air hunger")
    - · Coughing and frequent pulmonary infections
    - Most develop respiratory failure (hypoventilation)
    - accompanied by respiratory acidosis, hypoxemia

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#### **Homeostatic Imbalance**

- · Emphysema
  - Permanent enlargement of alveoli; destruction of alveolar walls; decreased lung elasticity →
    - Accessory muscles necessary for breathing
       → exhaustion from energy usage
    - Hyperinflation → flattened diaphragm → reduced ventilation efficiency
    - Damaged pulmonary capillaries → enlarged right ventricle

#### **Homeostatic Imbalance**

- · Chronic bronchitis
  - Inhaled irritants  $\rightarrow$  chronic excessive mucus  $\rightarrow$
  - Inflamed and fibrosed lower respiratory passageways  $\rightarrow$
  - Obstructed airways  $\rightarrow$
  - − Impaired lung ventilation and gas exchange →
  - Frequent pulmonary infections



#### **Homeostatic Imbalances**

- Asthma–reversible COPD
  - Characterized by coughing, dyspnea, wheezing, and chest tightness
  - Active inflammation of airways precedes bronchospasms
  - Airway inflammation is immune response caused by release of interleukins, production of IgE, and recruitment of inflammatory cells
  - Airways thickened with inflammatory exudate magnify effect of bronchospasms

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#### **Homeostatic Imbalances**

#### Tuberculosis (TB)

- Infectious disease caused by bacterium Mycobacterium tuberculosis
- Symptoms-fever, night sweats, weight loss, racking cough, coughing up blood
- Treatment- 12-month course of antibiotics
   Are antibiotic resistant strains

#### **Homeostatic Imbalance**

- Cystic fibrosis
  - Most common lethal genetic disease in North America
  - Abnormal, viscous mucus clogs passageways
     → bacterial infections
    - Affects lungs, pancreatic ducts, reproductive ducts
  - Cause–abnormal gene for Cl<sup>-</sup> membrane channel

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#### **Homeostatic Imbalances**

#### Lung cancer

- Leading cause of cancer deaths in North America
- 90% of all cases result of smoking
- Three most common types
  - Adenocarcinoma (~40% of cases) originates in peripheral lung areas - bronchial glands, alveolar cells
  - Squamous cell carcinoma (20–40% of cases) in bronchial epithelium
  - Small cell carcinoma (~20% of cases) contains lymphocytelike cells that originate in primary bronchi and subsequently metastasize

· Extra content from text

Figure 22.28 Embryonic development of the respiratory system.



#### **Developmental Aspects**

- By 28th week, premature baby can breathe on its own
- During fetal life, lungs filled with fluid and blood bypasses lungs
- · Gas exchange takes place via placenta

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#### **Homeostatic Imbalance**

- · Treatments for cystic fibrosis
  - Mucus-dissolving drugs; manipulation to loosen mucus; antibiotics
  - Research into
    - Introducing normal genes
    - Prodding different protein  $\rightarrow$  Cl<sup>-</sup> channel
    - Freeing patient's abnormal protein from ER to  $\rightarrow$  Cl  $^{\rm c}$  channels
    - · Inhaling hypertonic saline to thin mucus

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#### Haldane Effect

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- Amount of  $CO_2$  transported affected by  $Po_2$ 
  - Reduced hemoglobin (less oxygen saturation) forms carbaminohemoglobin and buffers H<sup>+</sup> more easily →
  - Lower Po<sub>2</sub> and hemoglobin saturation with O<sub>2</sub>; more CO<sub>2</sub> carried in blood
- Encourages CO<sub>2</sub> exchange in tissues and lungs

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#### Haldane Effect

- At tissues, as more CO<sub>2</sub> enters blood
  - More oxygen dissociates from hemoglobin (Bohr effect)
  - As HbO<sub>2</sub> releases O<sub>2</sub>, it more readily forms bonds with CO<sub>2</sub> to form carbaminohemoglobin

#### Influence of CO<sub>2</sub> on Blood pH

- Carbonic acid–bicarbonate buffer system–resists changes in blood pH
  - If H<sup>+</sup> concentration in blood rises, excess H<sup>+</sup> is removed by combining with  $HCO_3^- \rightarrow H_2CO_3$
  - If H<sup>+</sup> concentration begins to drop,  $H_2CO_3$  dissociates, releasing H<sup>+</sup>
  - HCO<sub>3</sub><sup>-</sup> is alkaline reserve of carbonic acidbicarbonate buffer system

#### **Chemical Factors**

- · Influence of arterial pH
  - Can modify respiratory rate and rhythm even if CO<sub>2</sub> and O<sub>2</sub> levels normal
  - Mediated by peripheral chemoreceptors
  - Decreased pH may reflect
    - CO<sub>2</sub> retention; accumulation of lactic acid; excess ketone bodies
  - Respiratory system controls attempt to raise pH by increasing respiratory rate and depth

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#### **Summary of Chemical Factors**

- Rising CO<sub>2</sub> levels most powerful respiratory stimulant
- Normally blood Po<sub>2</sub> affects breathing only indirectly by influencing peripheral chemoreceptor sensitivity to changes in Pco<sub>2</sub>

#### **Summary of Chemical Factors**

- When arterial Po<sub>2</sub> falls below 60 mm Hg, it becomes major stimulus for respiration (via peripheral chemoreceptors)
- Changes in arterial pH resulting from CO<sub>2</sub> retention or metabolic factors act indirectly through peripheral chemoreceptors

#### Acclimatization to High Altitude

- Always lower-than-normal Hb saturation levels
  - Less O<sub>2</sub> available
- Decline in blood O<sub>2</sub> stimulates kidneys to accelerate production of EPO
- RBC numbers increase slowly to provide long-term compensation

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