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9.1 Evolution of Circulation



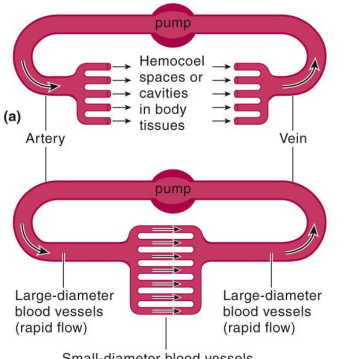
- Circulatory systems in multicellular organisms supplement diffusion with bulk transport
- Components of a circulatory system
 - Fluid that carries transported molecules and cells
 - A **pump** to move the fluids
 - Vessels to carry fluid between the pump and body tissues

9.1 Evolution of Circulation

- Open vs. closed circulatory systems
 - Open systems
 - Hemolymph moves through vessels that open into extracellular spaces
 - Closed systems
 - **Blood** is pumped from a heart through vessels that return blood to the heart
 - **Capillaries** are the primary structure distinguishing a closed from an open system

9.1 Evolution of Circulation

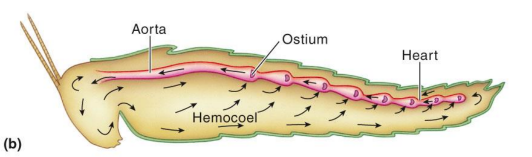




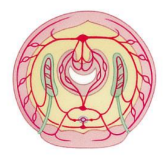
Small-diameter blood vessels (leisurely flow in diffusion zone) capillaries

(c)

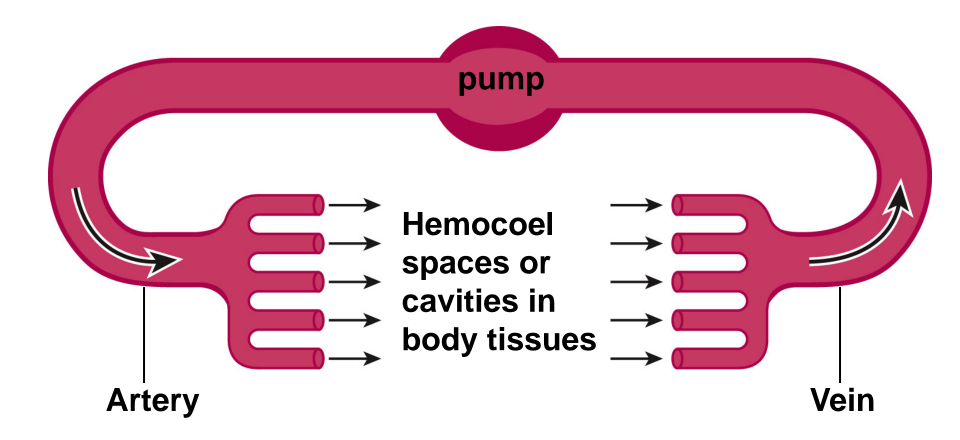
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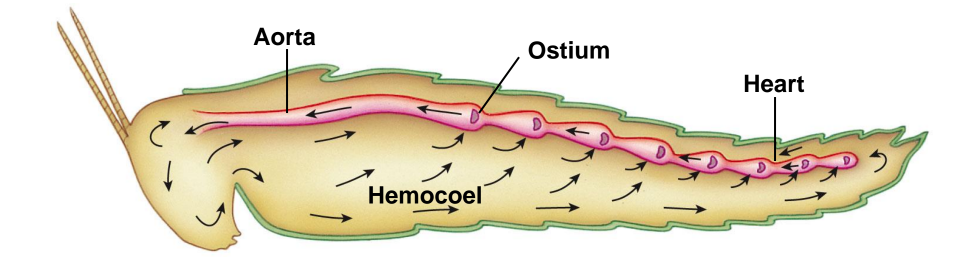


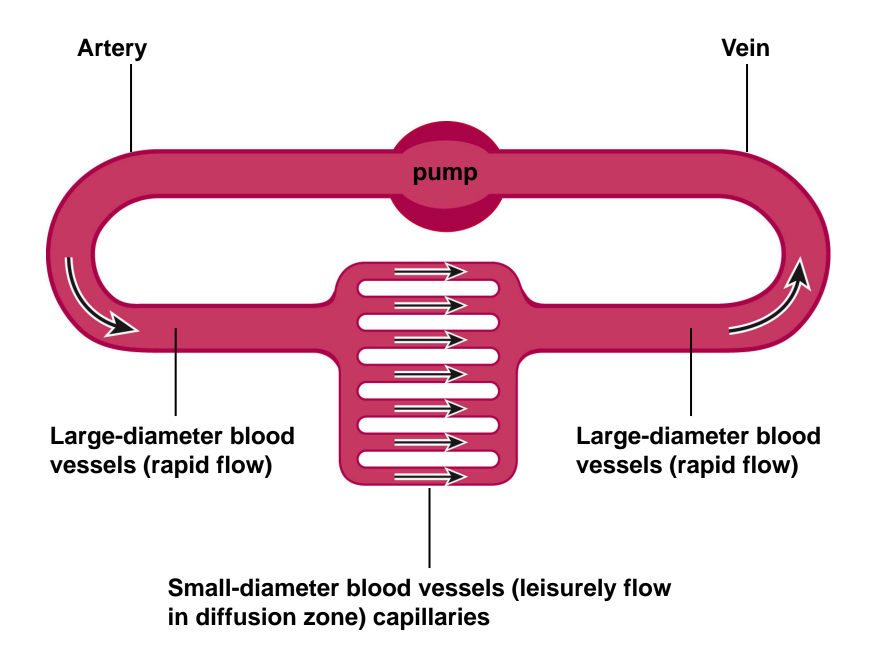
Dorsal blood vessel Two of five Ventral blood vessels Gut cavity

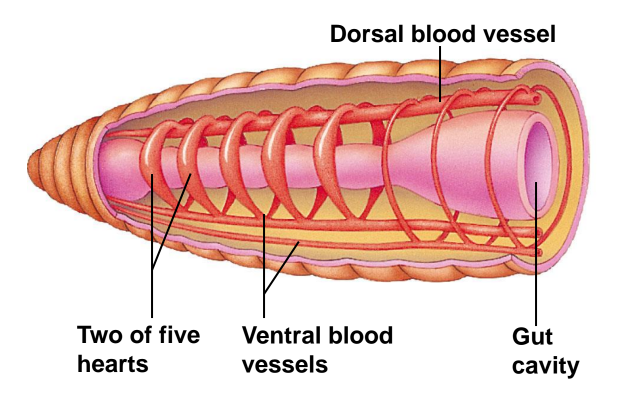


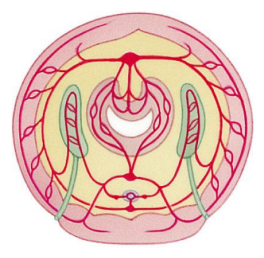
Segmented blood vessels (red) service muscles, nephridia, and other organs of each segment







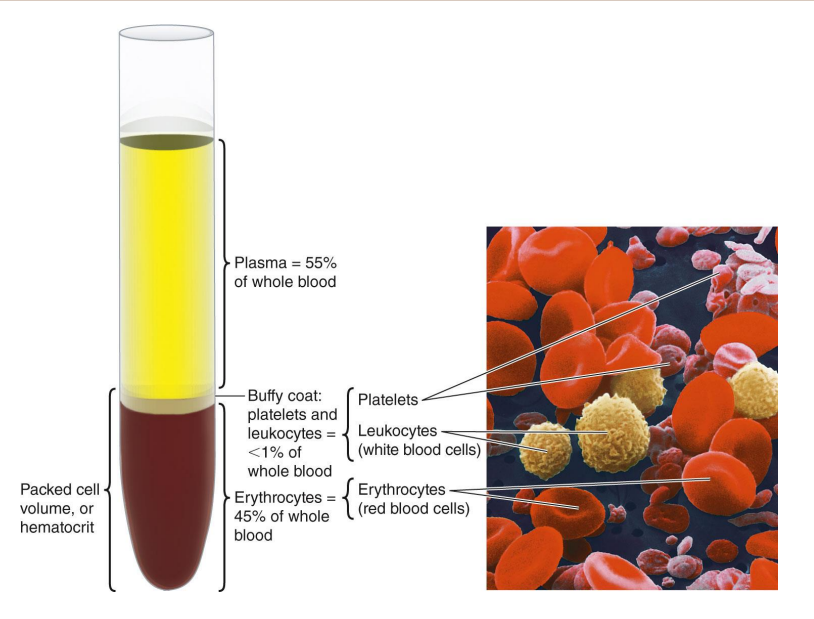




Segmented blood vessels (*red*) service muscles, nephridia, and other organs of each segment



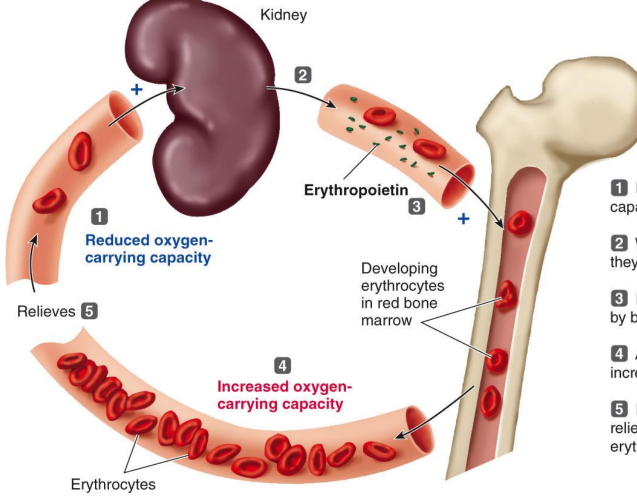
- Circulatory fluids are divided into plasma and cellular elements
 - Hematocrit is the percentage of blood volume occupied by cells
 - 45% in human males, 42% in human females
 - **Plasma** is an aqueous medium for transport of inorganic ions, gases and organic solutes
 - 90% water
 - Plasma proteins are the most plentiful organic solutes (6 8% of plasma)
 - Lipoproteins carry energy lipids (triglycerides) and structural lipids (phospholipids and cholesterol)





- Erythrocytes (red blood cells) transport oxygen from lungs or gills to tissues
 - Contain hemoglobin
 - Oval shaped in most vertebrates
 - Biconcave discs without nuclei or organelles in mammals
 - **Spleen** removes old erythrocytes and stores healthy erythrocytes, platelets and lymphocytes
 - Hemopoietic tissues (e.g. red bone marrow in birds and mammals) generate new erythrocytes (erythropoiesis)
 - **Erythropoietin** (EPO) is secreted by the kidneys to stimulate erythropoiesis . It is also **produced** in perisinusoidal cells in the **liver**.





1 Kidneys detect reduced O₂-carrying capacity of blood.

2 When less O_2 is delivered to the kidneys, they secrete erythropoietin into blood.

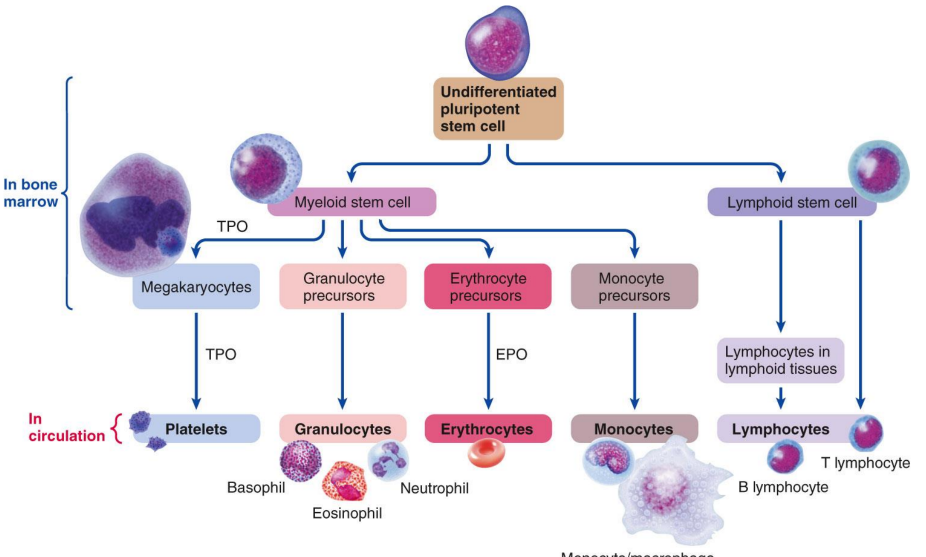
3 Erythropoietin stimulates erythropoiesis by bone marrow.

Additional circulating erythrocytes increase O₂-carrying capacity of blood.

5 Increased O₂-carrying capacity relieves initial stimulus that triggered erythropoietin secretion.



- Leukocytes (white blood cells) are key components of vertebrate immune systems
- Thrombocytes and platelets are involved in blood clotting
 - Thrombocytes are living cells found in all vertebrates except mammals
 - Break up into platelet-like fragments when activated by injury
 - Platelets are cell fragments circulating in mammalian blood
 - Shed from **megakaryocytes** in bone marrow



Monocyte/macrophage

9.3 Circulatory Fluids: Hemostasis

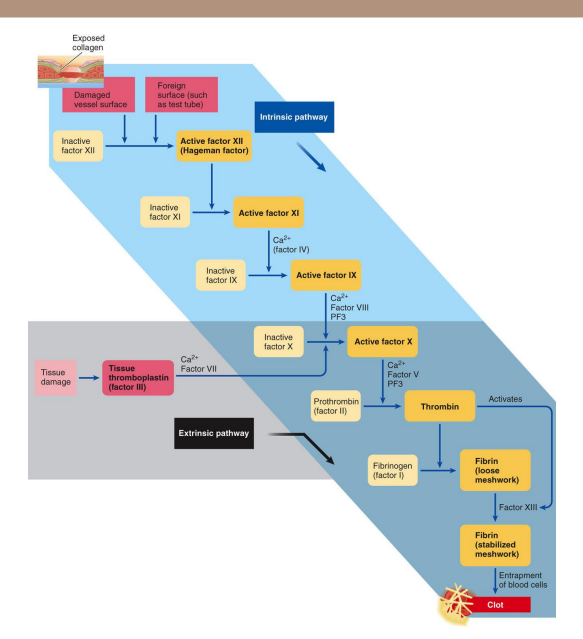


- Blood coagulation is the transformation of blood from a liquid into a solid gel.
 - Clotting cascade involves 12 clotting factors
 - Most are plasma proteins synthesized by the liver
 - Present in plasma in inactive forms
 - Intrinsic pathway involves 7 steps
 - Activation of the first factor (factor XII) triggers the clotting cascade
 - Extrinsic pathway involves 4 steps and requires contact with tissue factors outside of the blood

Final steps

- Thrombin converts fibrinogen into fibrin
- Fibrin, an insoluble thread-like molecule, forms a stabilized meshwork at the site of a platelet plug

9.3 Circulatory Fluids: Hemostasis



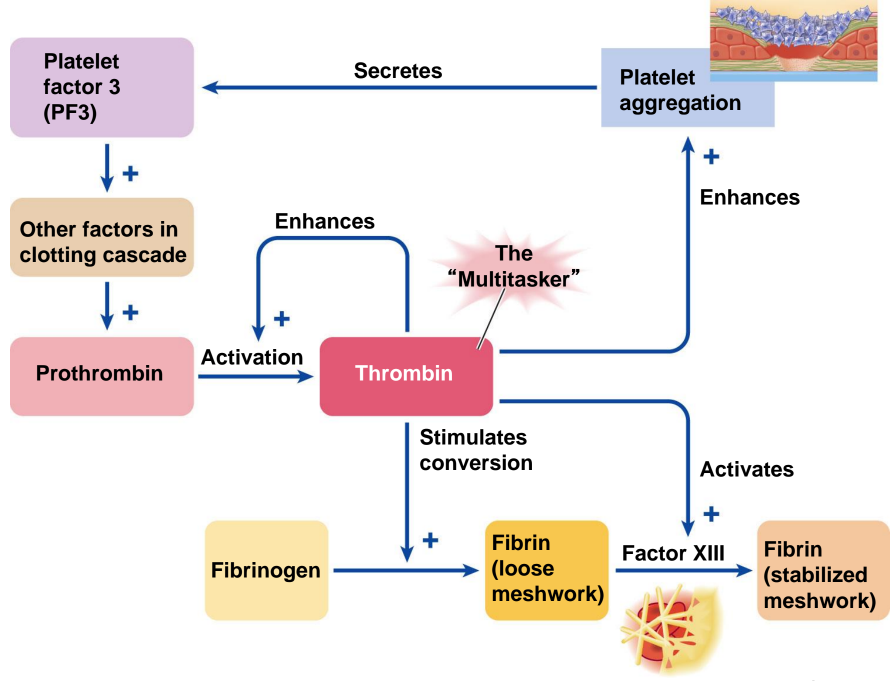
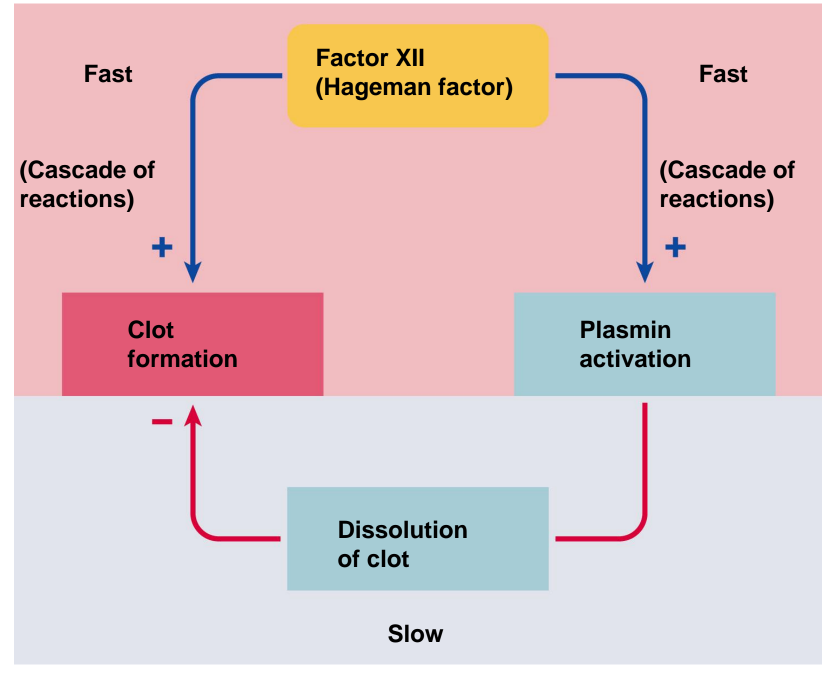


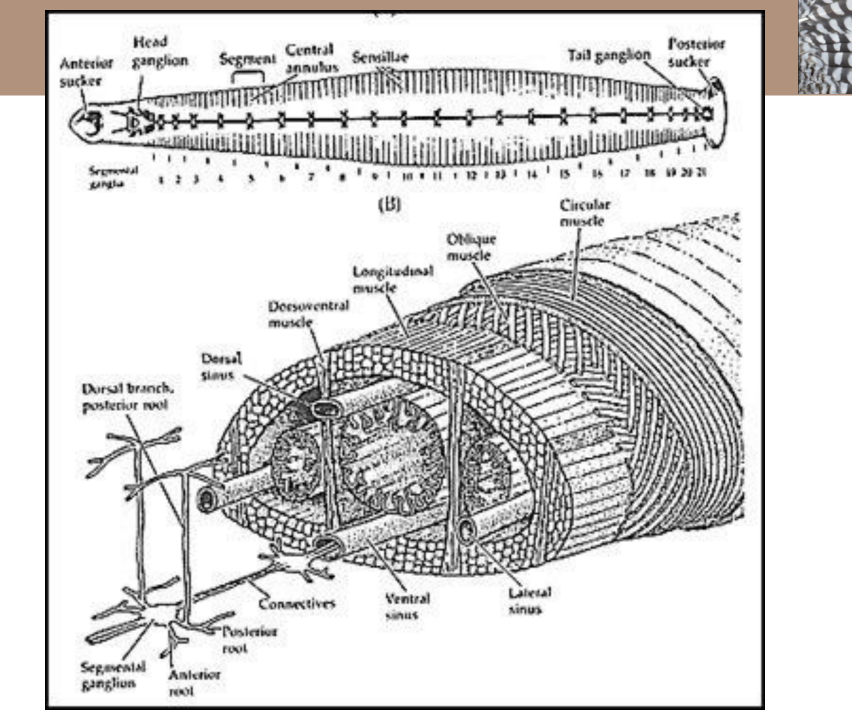
Figure 9-11 p397

9.3 Circulatory Fluids: Hemostasis

- Plasmin dissolves clots
 - **Plasminogen** is produced by the liver
 - Factor XII (which began the clotting cascade) triggers a cascade of reactions leading to activation of plasminogen to form **plasmin**
 - Tissue plasminogen activator (tPA) produces a low level of plasmin activation in the absence of clotting

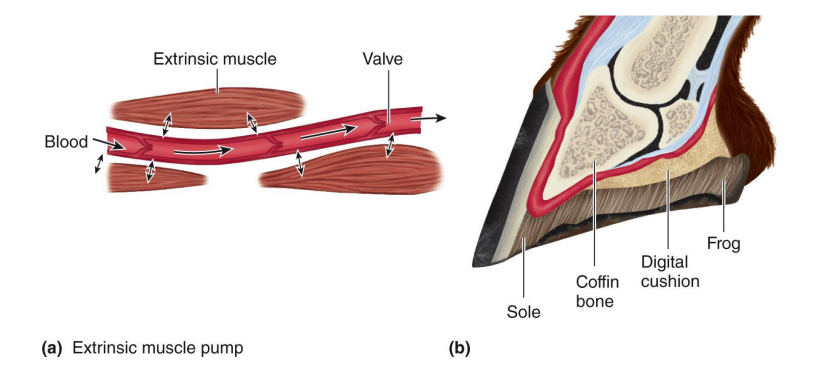


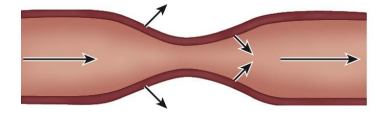




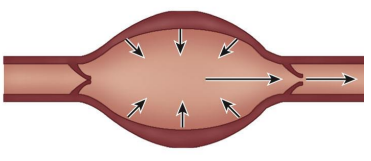
- Pumping mechanisms
 - Flagella
 - Flagella of epithelial cells create slow currents
 - Extrinsic muscle or skeletal pumps
 - Muscle contractions associated with locomotion
 - Peristaltic (tubular) muscle pumps
 - Walls of vessels contract in a moving wave
 - Chamber muscle pumps
 - One-way valves create flow in one direction
 - Auxiliary hearts
 - Boost flow to certain parts (e.g. gills) in animals with primary hearts





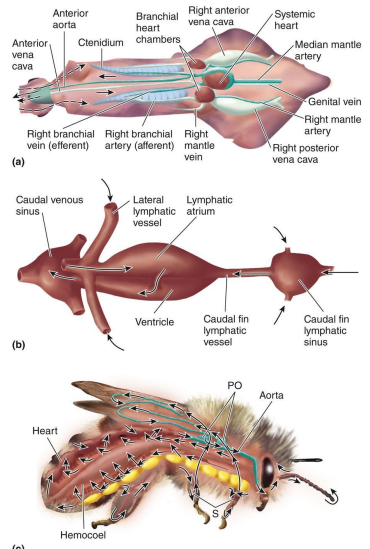


(c) Peristaltic heart



(d) Chambered heart





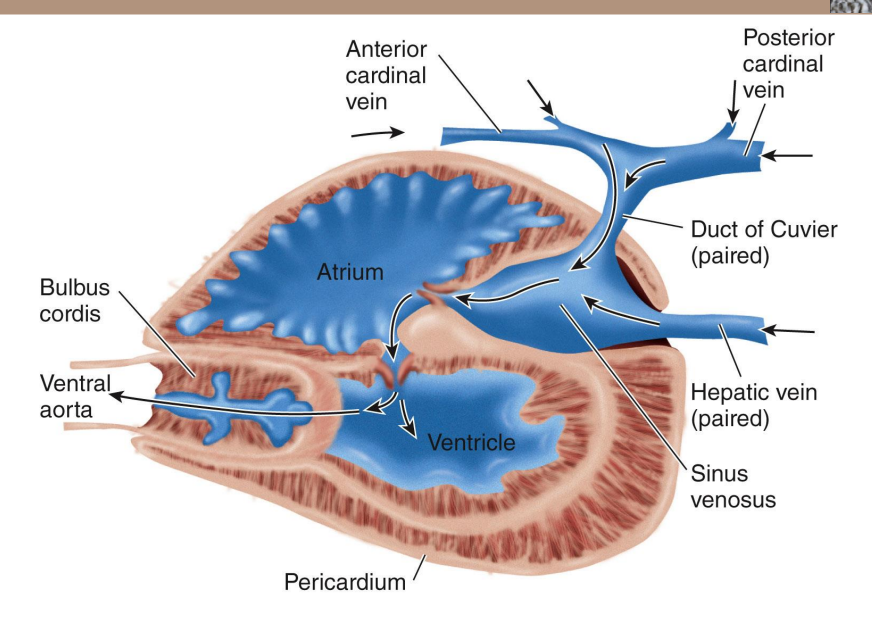
(c)

Arthropod hearts

- Dorsally located
- Single-chamber pumps (crustaceans)
- Tubular pumps (insects and arachnids)
- **Ostia** are pore-like openings that allow hemolymph to reenter the heart

Fish hearts have two primary chambers

- Atrium collects returning blood
- Ventricle pumps to body
- Auxiliary chambers
 - Sinus venosus collects blood from veins before entering atrium
 - Conus arteriosus (in cartilaginous fish) or bulbus arteriosus (in bony fish) dampen pulsatile pressure output of the ventricle

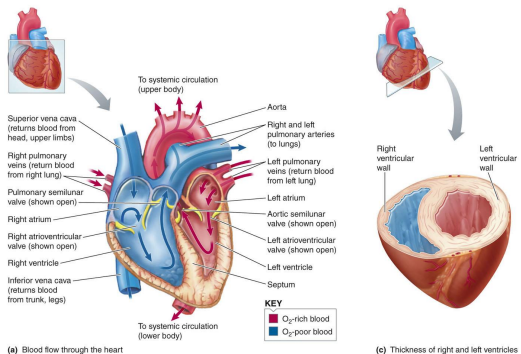


- Avian and mammalian hearts have dual pumps
 - Two atria receive blood
 - Two ventricles pump blood
 - Two pumps are separated by the **septum**
 - Right half pumps oxygen-depleted blood into the pulmonary circulation

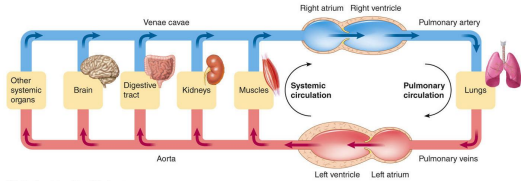
• Left half pumps oxygen-rich blood into the systemic circulation

• Both sides of the heart pump equal amounts of blood

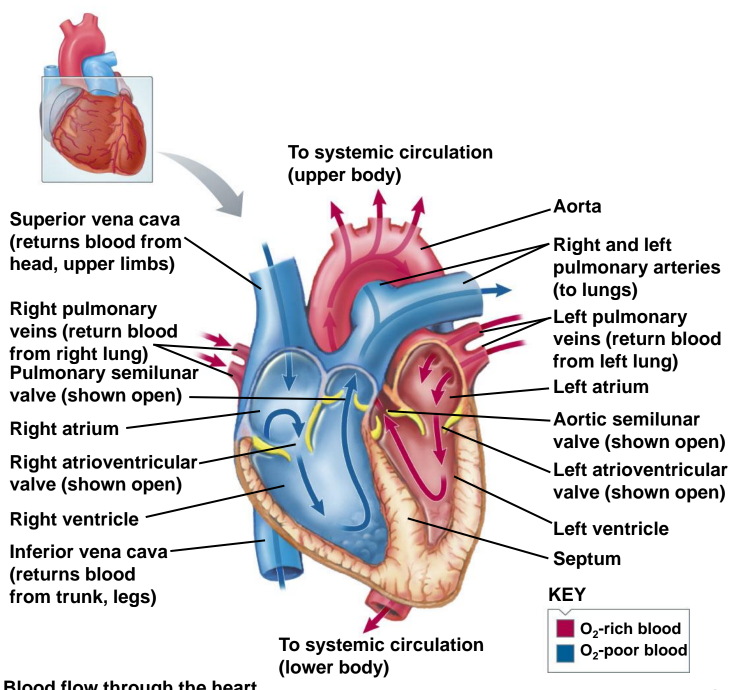




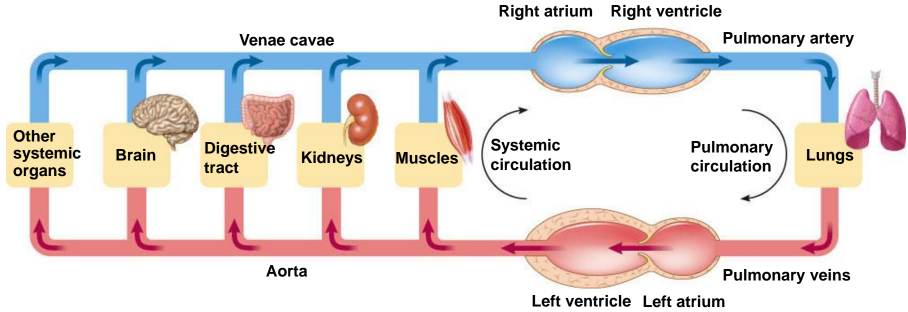




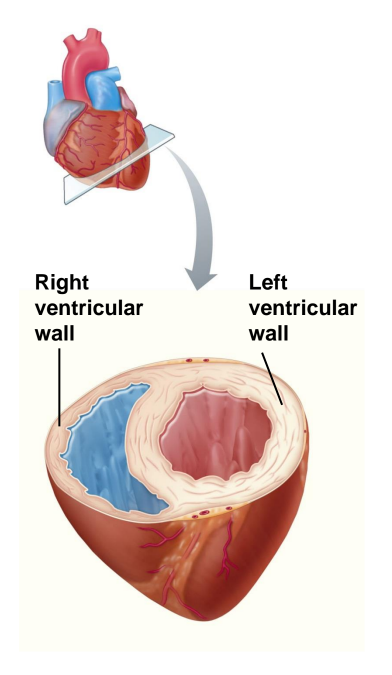
(b) Dual pump action of the heart



(a) Blood flow through the heart



(b) Dual pump action of the heart

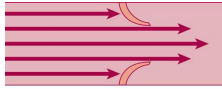


(c) Thickness of right and left ventricles

Figure 9-18c p404

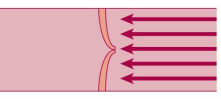


- Heart valves ensure unidirectional flow of blood
 - **Right** and **left atrioventricular** (AV) **valves** allow flow from atria into ventricles during ventricular filling
 - Aortic and pulmonary (semilunar) valves allow flow from ventricles into arteries during ventricular contraction
 - Backflow from atria into veins does not occur because atrial pressures are not much higher than venous pressures



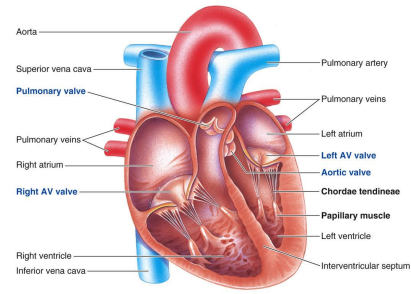
Valve opened

When pressure is greater behind the valve, it opens.



Valve closed; does not open in opposite direction

When pressure is greater in front of the valve, it closes. Note that when pressure is greater in front of the valve, it does not open in the opposite direction; that is, it is a one-way valve.



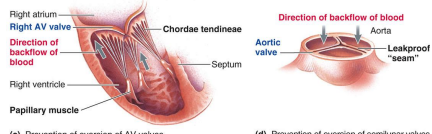
(a) Location of the heart valves in a longitudinal section of the heart



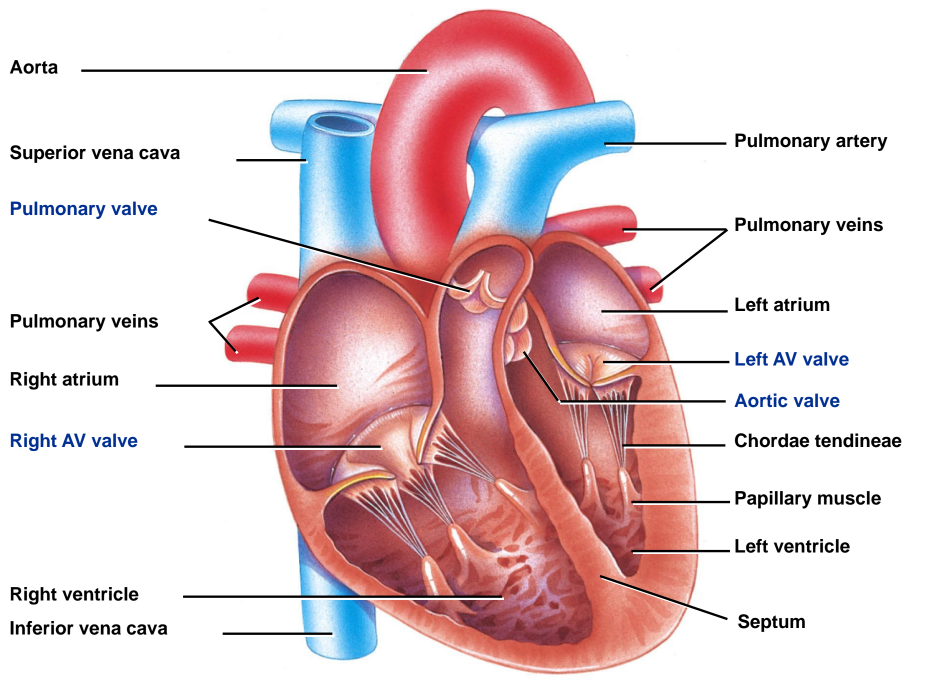




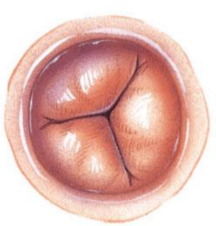
Aortic or pulmonary valve



(c) Prevention of eversion of AV valves



(a) Location of the heart valves in a longitudinal section of the heart





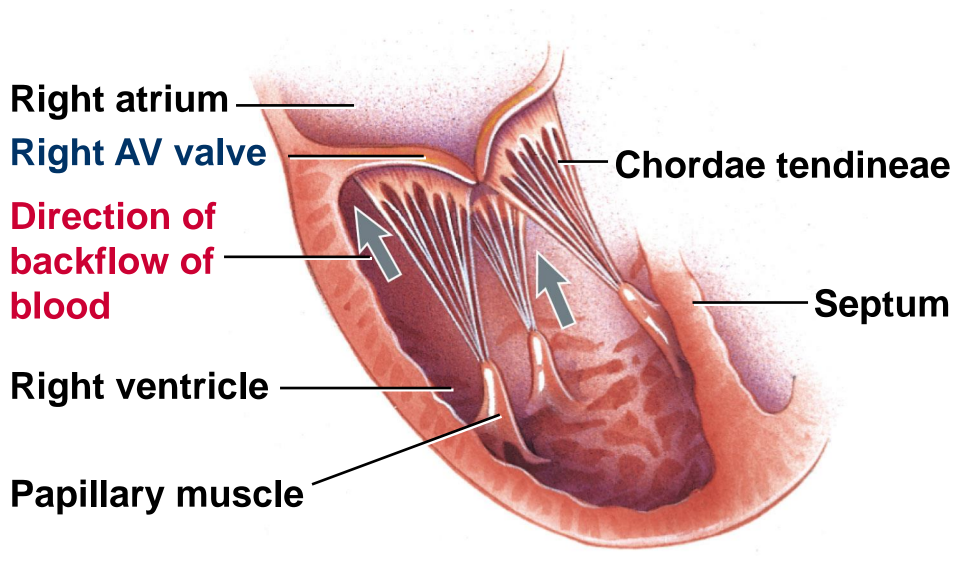
Right AV valve

Left AV valve

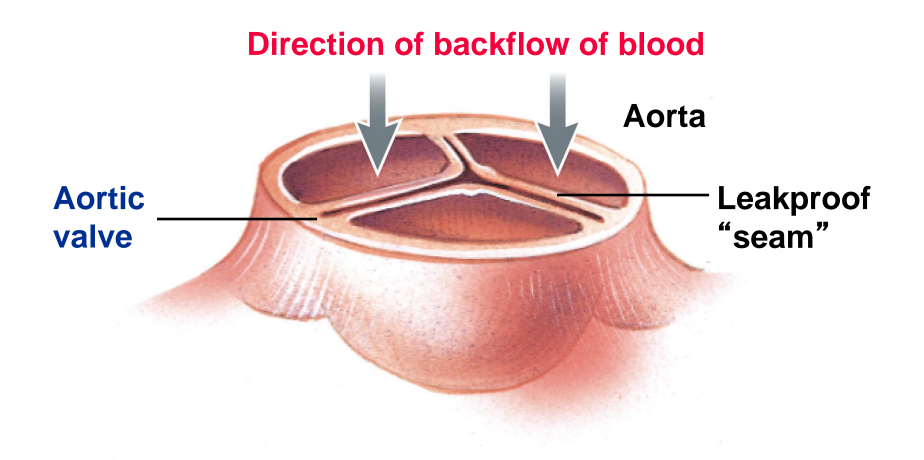
(b) Heart valves in closed position, viewed from above



Aortic or pulmonary valve



(c) Prevention of eversion of AV valves



(d) Prevention of eversion of semilunar valves

Figure 9-20d p405

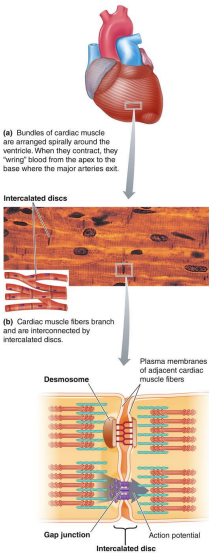
9.4 Circulatory Pumps: Evolution



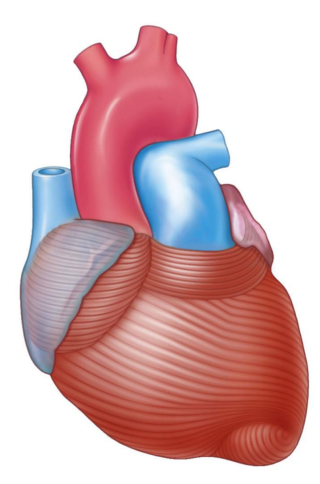
- Vertebrate heart walls
 - Thick, muscular myocardium is sandwiched between endocardium and epicardium
 - Myocardium consists of interlacing bundles of cardiac muscle fibers arranged spirally
 - Cardiac muscle cells form branching fibers with adjacent cells joined end-to-end at intercalated discs
 - Intercalated discs contain desmosomes and gap junctions
 - Impulses spread to all cells joined by gap junctions to form a functional syncytium

9.4 Circulatory Pumps: Evolution



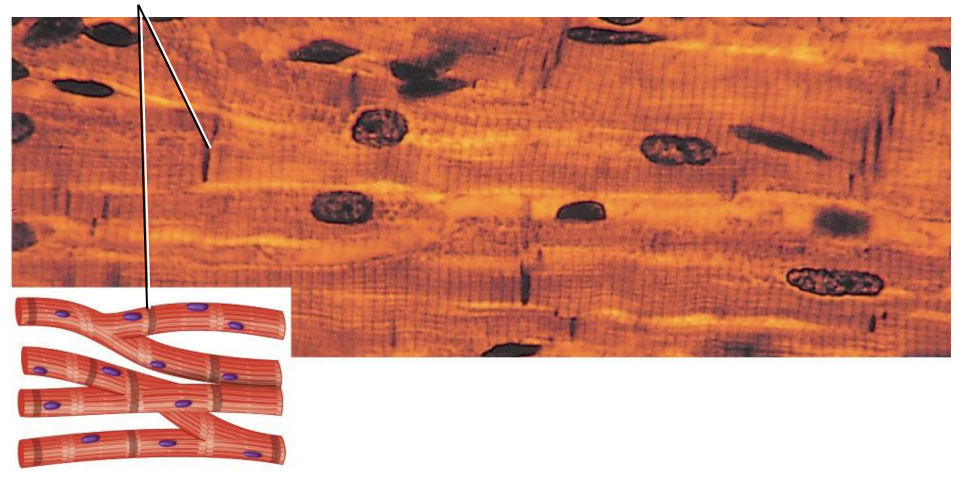


(c) Intercalated discs contain two types of membrane junctions: mechanically important desmosomes and electrically important gap junctions.

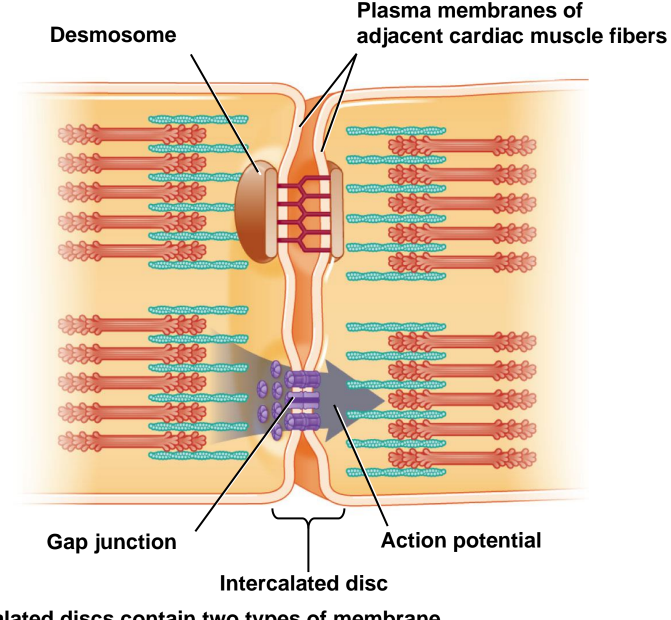


(a) Bundles of cardiac muscle are arranged spirally around the ventricle. When they contract, they "wring" blood from the apex to the base where the major arteries exit.

Intercalated discs



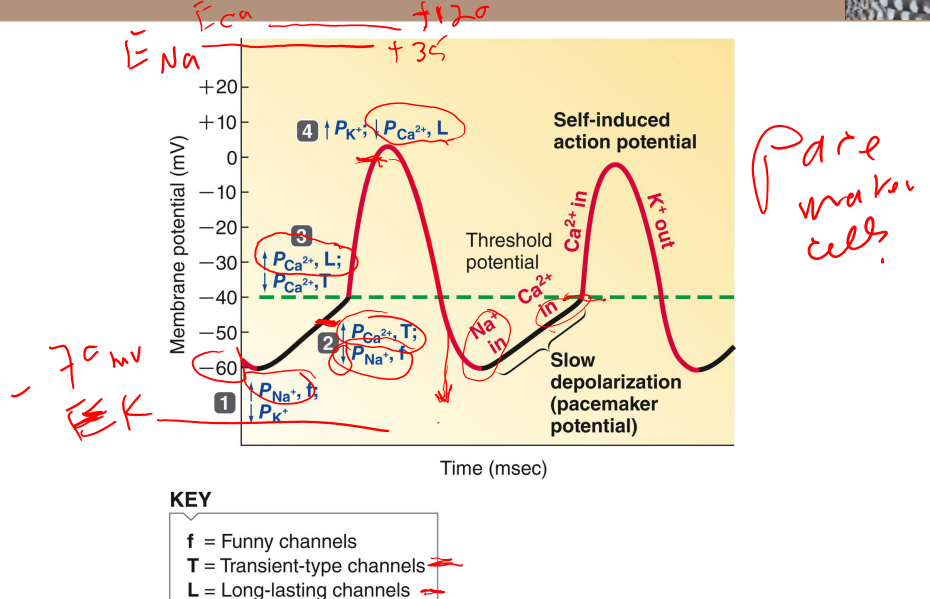
(a) Cardiac muscle fibers branch and are interconnected by intercalated discs.



(c) Intercalated discs contain two types of membrane junctions: mechanically important desmosomes and electrically important gap junctions.

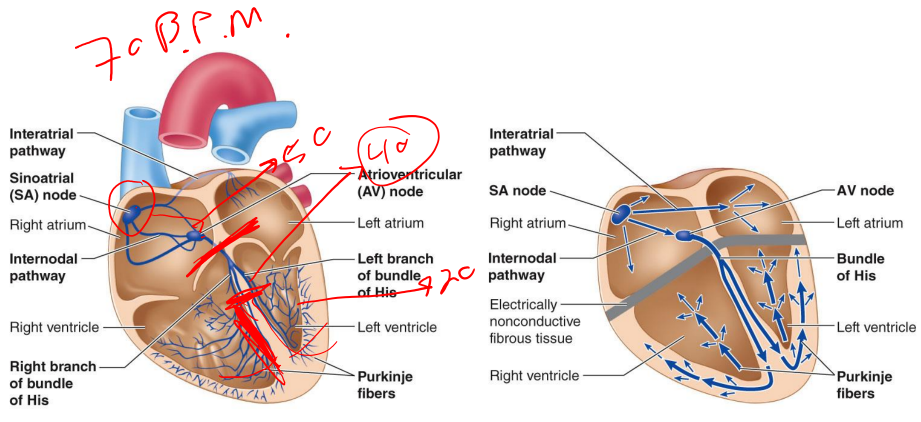


- Neurogenic hearts require external neural stimulus to beat (e.g. decapod crustaceans)
- Myogenic hearts have pacemaker cells
 - Membrane potential of pacemaker cells slowly depolarizes due to:
 - Increased inward Na⁺ current
 - Decreased outward K⁺ current
 - Increased inward Ca²⁺ current
 - Action potential is produced when L-type Ca²⁺ channels open at threshold
 - Large influx of Ca²⁺ causes rapid depolarization



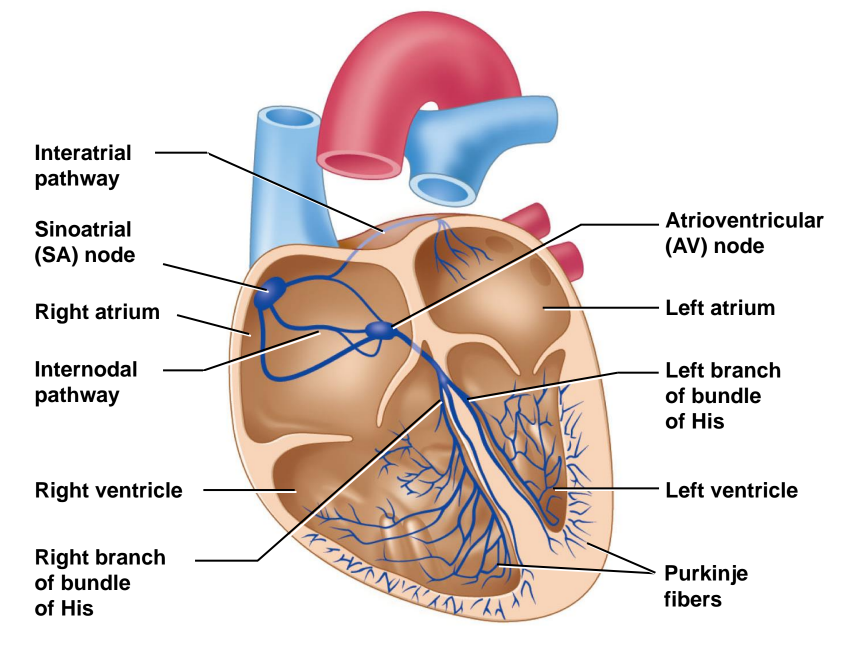
- Conduction pathway
 - 1. Sinoatrial (SA) node is the normal pacemaker
 - Fastest rate of autorhythmicity
 - 2. Atrial excitation
 - 3. Atrioventricular (AV) node transmits impulses between the atria and the ventricles
 - Slower rate of autorhythmicity
 - Impulse is delayed about 0.1 sec
 - 4. Ventricular excitation
 - Impulse passes down the bundle of His in the interventricular septum
 - Purkinje fibers extend through the ventricular myocardium



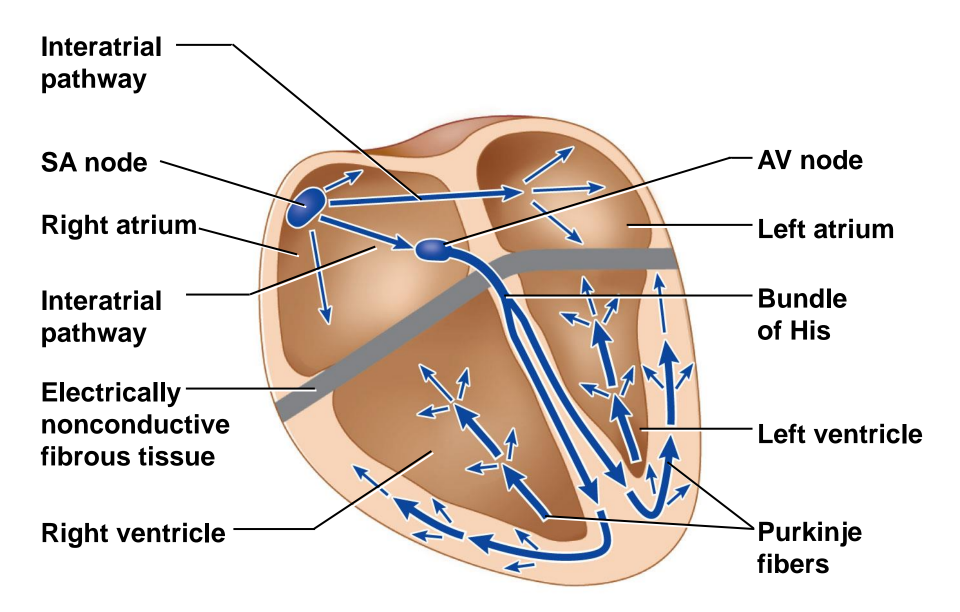


(a) Specialized conduction system of the heart

(b) Spread of cardiac excitation



(a) Specialized conduction system of the heart

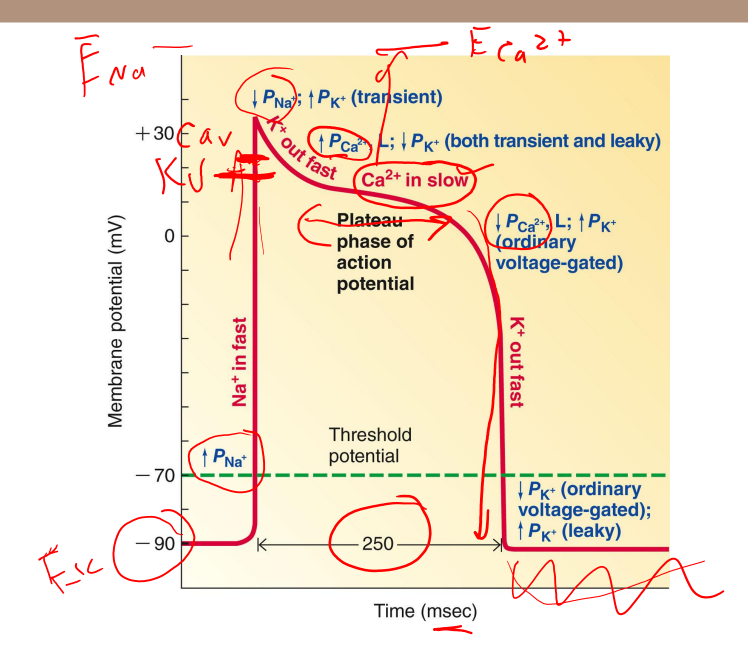


(b) Spread of cardiac excitation



- The action potential of contractile cardiac muscle cells shows a plateau phase.
 - Contractile cells remain relaxed until excited by adjacent cells through gap junctions
 - Rising phase
 - Opening of voltage-gated Na⁺ channels
 - K⁺ channels transiently open at the peak of action potential, producing slight repolarization
 - Plateau phase
 - Opening of "slow" L-type Ca²⁺ channels
 - Closing of K⁺ channels
 - Prolongs contraction to aid pumping
 - Falling phase
 - Opening of voltage-gated K⁺ channels

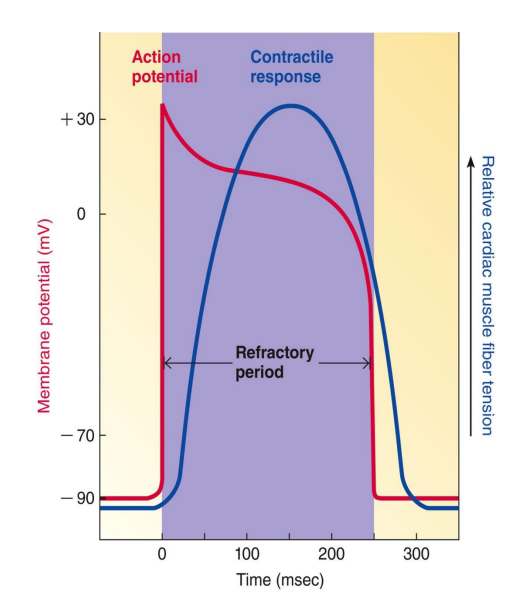






- Excitation-contraction coupling in the heart
 - Ca²⁺ enters the cytosol through L-type channels in T tubules
 - Triggers release of Ca²⁺ from sarcoplasmic reticulum
 - Ca²⁺ binds to troponin-tropomyosin complex to allow cross-bridge cycling
 - Extent and duration of cross-bridge activity varies with the amount of cytosolic Ca²⁺
 - Cardiac muscle has a long refractory period (250 msec)
 - Na⁺ channels remain inactivated during the plateau phase
 - Prevents summation and tetanus





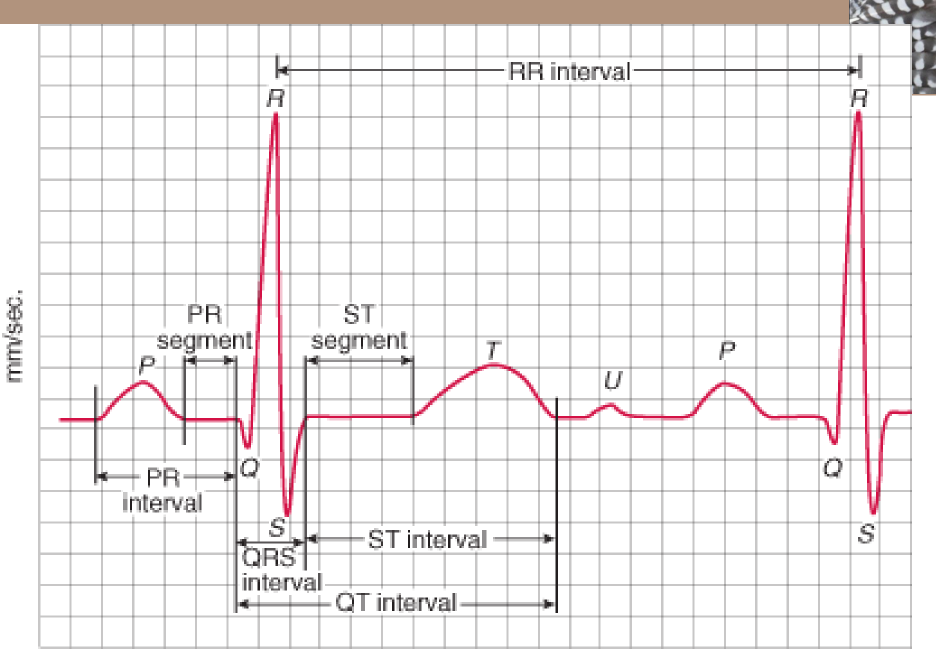


- Electrocardiogram (ECG)
 - Electrical currents generated by cardiac muscle can be detected using **recording electrodes** on the skin.
 - Interpretation of the ECG
 - P wave represents atrial depolarization
 - QRS complex represents ventricular depolarization
 - T wave represents ventricular repolarization
 - Periods of no current flow
 - PR segment represents AV nodal delay
 - ST segment represents plateau phase
 - **TP interval** represents **passive ventricular filling** while all chambers are at rest



(a) Electrocardiogram

```
P = Atrial
                 depolarization
        Q,R,S = Ventricular
                 depolarization
      R
             T = Ventricular
                 repolarization
Р
```

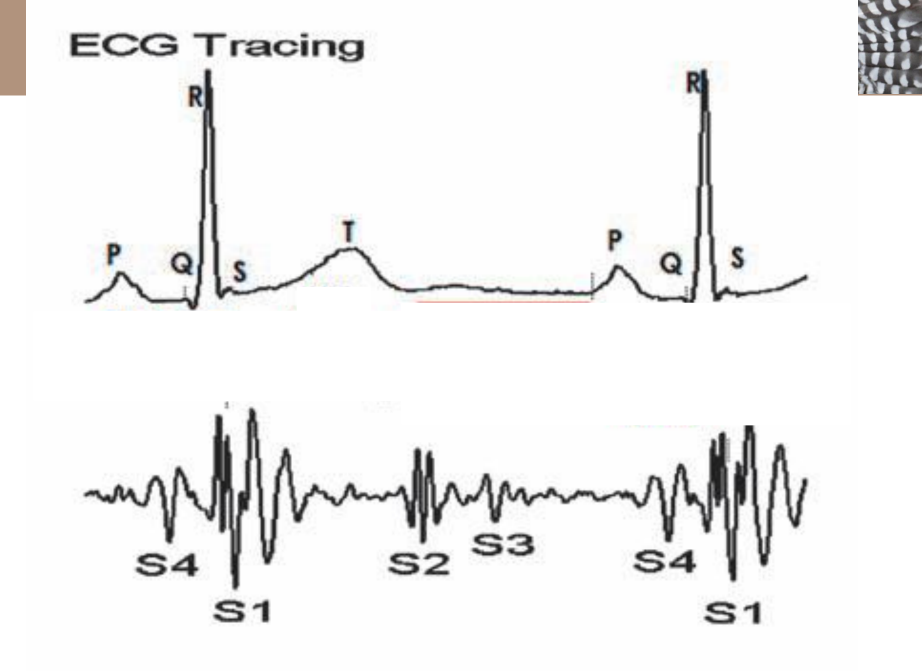


mm/mV 1 square = 0.04 sec/0.1mV



Table 5.1 Components of the ECG & Typical Lead II Values*

ECG COMPONENT		Measurement area	Represent	Duration (seconds)	Amplitude (millivolts)
Waves	P	begin and end on isoelectric line (baseline); normally upright in standard limb leads	depolarization of the right and left atria.	0.07 – 0.18	< 0.25
	QRS complex	begin and end on isoelectric line (baseline) from start of Q wave to end of S wave	depolarization of the right and left ventricles. Atrial repolarization is also part of this segment, but the electrical signal for atrial repolarization is masked by the larger QRS complex (see Fig 5.2)	0.06 – 0.12	0.10 - 1.50
	Т	begin and end on isoelectric line (baseline)	repolarization of the right and left ventricles.	0.10 - 0.25	< 0.5
	P-R	from start of P wave to start of QRS complex	time from the onset of atrial depolarization to the onset of ventricular depolarization.	0.12-0.20	
Intervals	Q-T	from start of QRS complex to end of T wave	time from onset of ventricular depolarization to the end of ventricular repolarization. It represents the refractory period of the ventricles.	0.32-0.36	
	R-R	from peak of R wave to peak of succeeding R wave	time between two successive ventricular depolarizations.	0.80	
υ.	P-R	from end of P wave to start of QRS complex	time of impulse conduction from the AV node to the ventricular myocardium.	0.02 - 0.10	
Segments	S-T	between end of S wave and start of T wave	period of time representing the early part of ventricular repolarization during which ventricles are more or less uniformly excited.	< 0.20	
	T-P	from end of T wave to start of successive P wave	time from the end of ventricular repolarization to the onset of atrial depolarization.	0.0 - 0.40	





Normal heart sounds are associated with heart valves closing, causing changes in blood flow

Common:

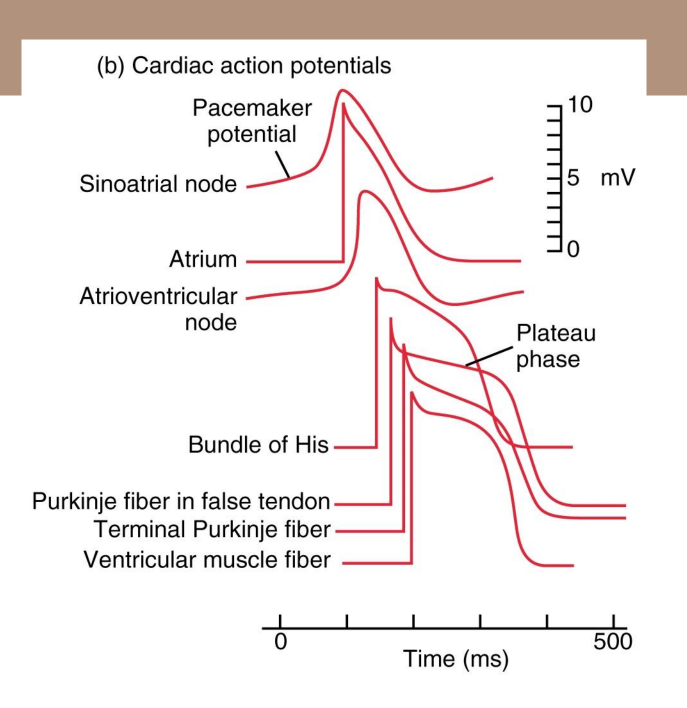
S₁: The first heart tone, or **S**₁, forms the "lubb" of "lubb-dub" Closure of the atrioventricular valves

S₂: The second heart tone, or **S₂**, forms the "dub" of "lubb-dub" Closure of the aortic valve and pulmonary valve at the end of ventricular systole

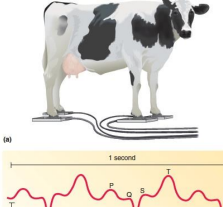
Not so common: In young children can hear. IN NEONATES ONE SHOULD NOT HEAR S3 AND S4.

S₃: Also called a **protodiastolic gallop**, **ventricular gallop**, or informally the "Kentucky" gallop (S1=ken; S2=tuc; S3=ky). It is not of valvular origin.

S3 is thought to be caused by the oscillation inrushing blood from the atria.



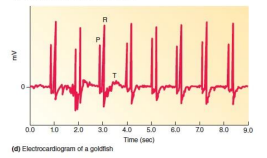


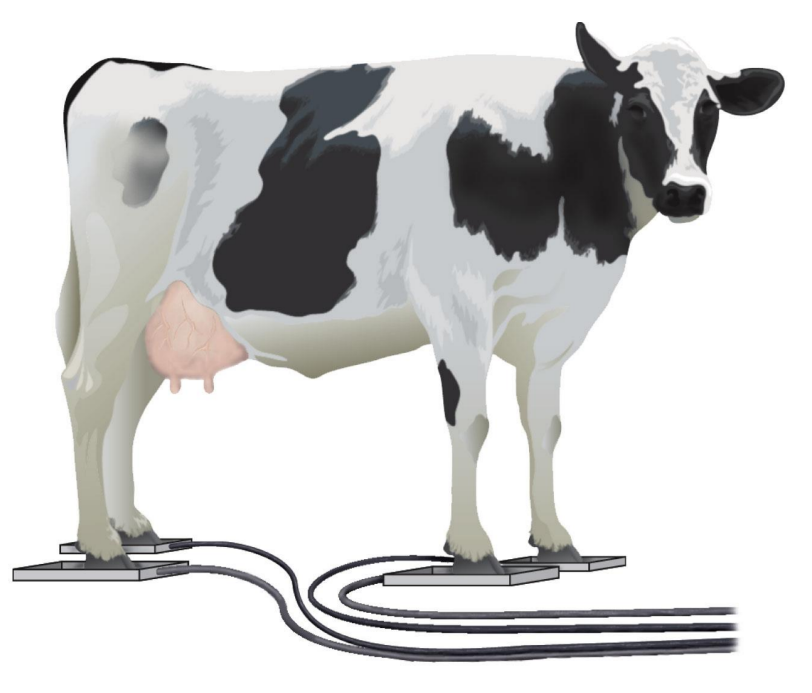


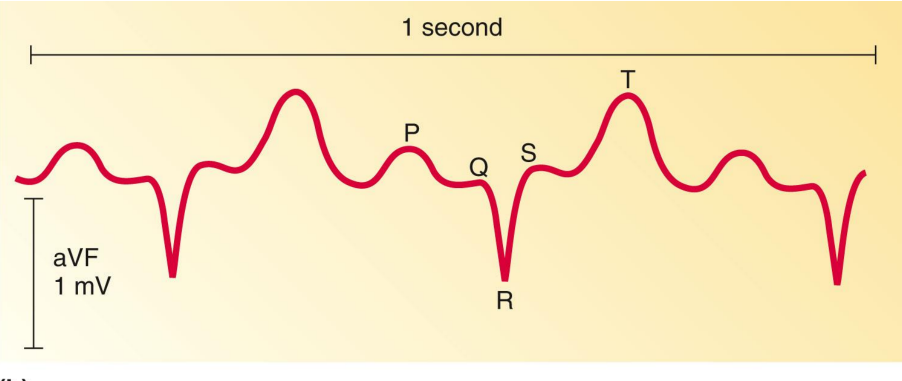




(c) Goldfish with implanted sensor for ECG

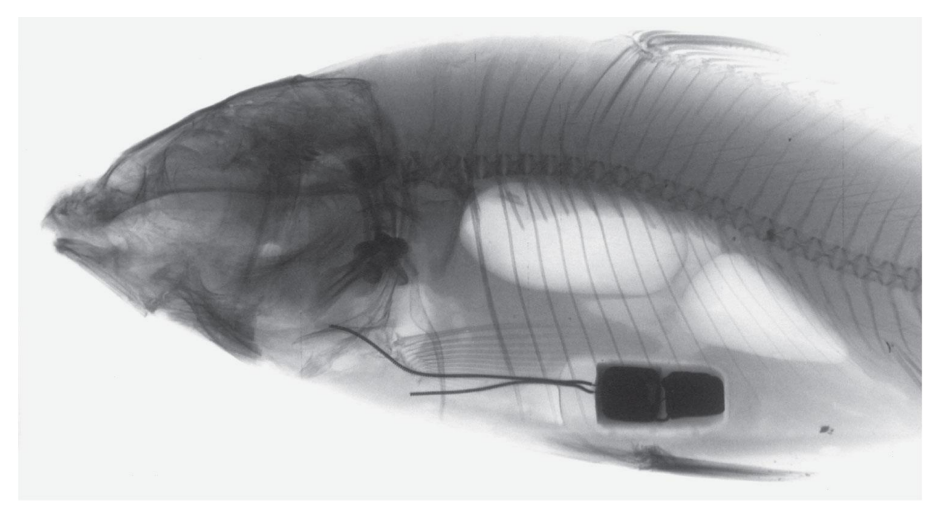




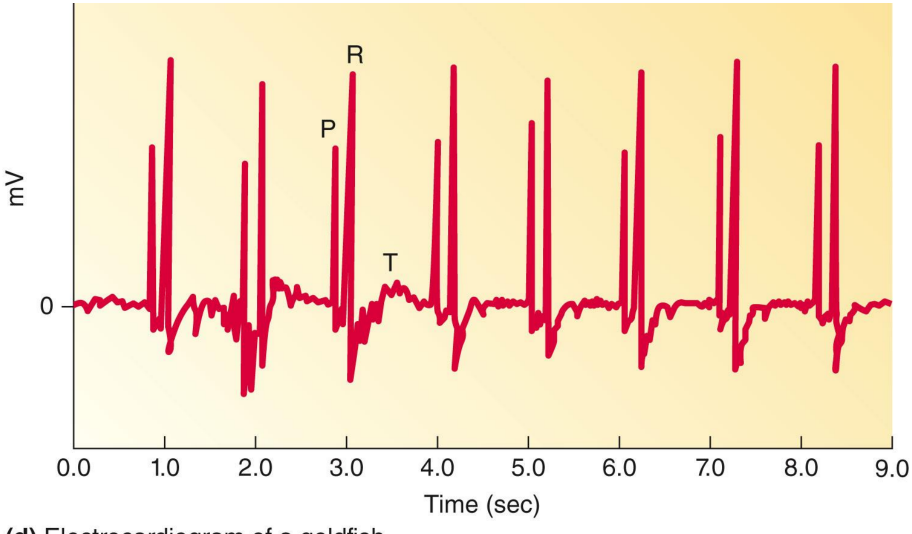


(b)

Figure 9-26b p411



(c) Goldfish with implanted sensor for ECG



(d) Electrocardiogram of a goldfish



- The cardiac cycle consists of alternating periods of systole and diastole
 - Systole is the period of contraction and emptying
 - **Diastole** is the period of relaxation and filling
 - Events are the same on the left and right sides of the heart
 - Pressures are lower on the right

Events of the cardiac cycle

1. Early ventricular diastole

- AV valves are open; aortic and pulmonary valves are closed
- Blood flows from veins into atria into ventricles

2. Late ventricular diastole

- SA node fires and atria depolarize (P wave)
- Atria contract, increasing atrial pressure

3. End of ventricular diastole

• Ventricular filling is completed (end-diastolic volume)

4. Ventricular excitation and onset of ventricular systole

- Impulse passes through AV node and bundle of His and depolarizes ventricles (QRS complex)
- Ventricles contract, increasing ventricular pressure
- AV valves close



- Events of the cardiac cycle
 - 5. Isovolumetric ventricular contraction
 - Ventricles are closed chambers while ventricular pressure rapidly rises
 - 6. Ventricular ejection
 - When ventricular pressure exceeds arterial blood pressure, aortic and pulmonary valves open
 - Blood is forced into arteries and ventricular volume decreases substantially

7. End of ventricular systole

- Ventricular emptying is completed (end-systolic volume)
- Amount of blood pumped out is the stroke volume
- 8. Ventricular repolarization and onset of ventricular diastole
 - Ventricles repolarize (T wave)
 - Ventricular pressure falls below arterial blood pressure and aortic and pulmonary valves close (dichrotic notch)



Events of the cardiac cycle

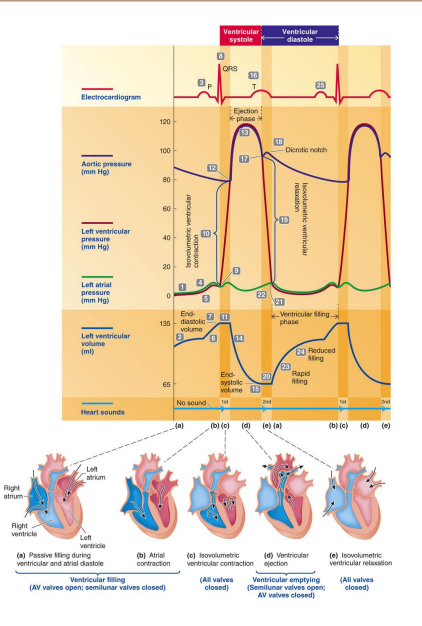
9. Isovolumetric ventricular relaxation

Ventricles are closed chambers while ventricular pressure rapidly falls

10. Ventricular filling

- When ventricular pressure falls below atrial pressure, AV valves open
- Ventricles begin to fill with blood
- Cardiac cycle repeats beginning with firing of SA node





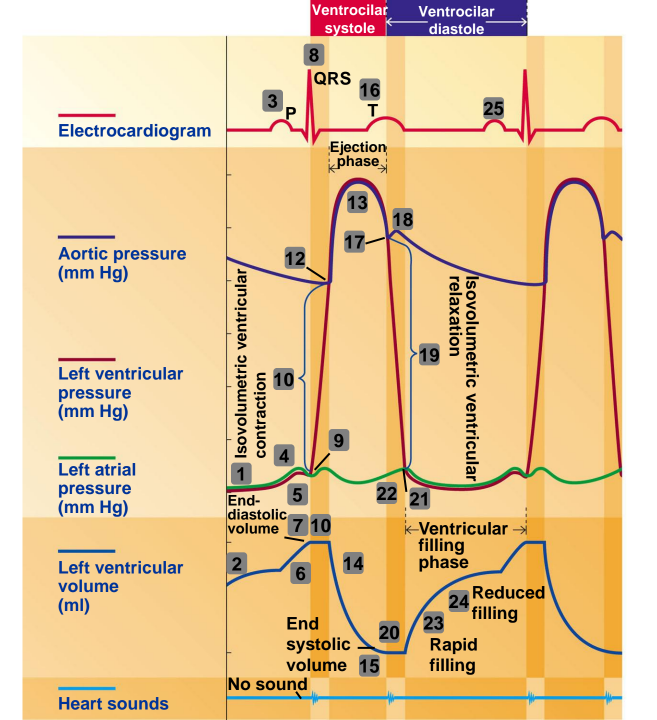
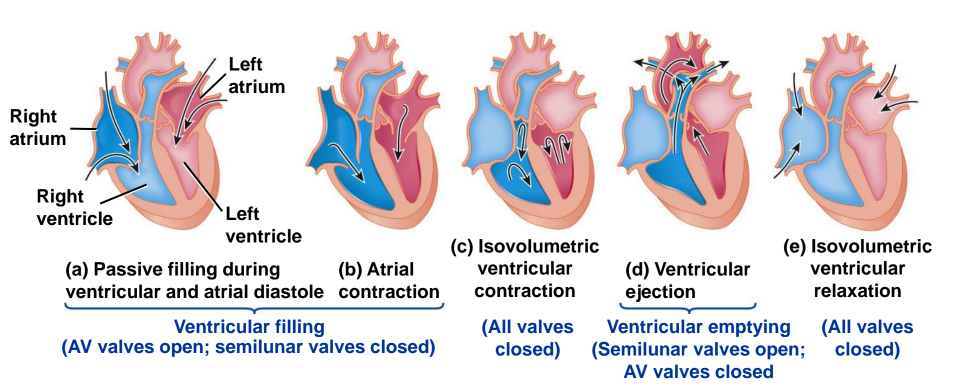


Figure 9-27 p413



9.7 Circulatory Pumps: Cardiac Output and Its Control



 Cardiac output (C.O.) is the volume of blood pumped per minute by a heart to the body.

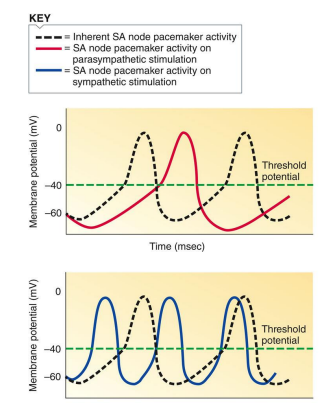
Cardiac output = heart rate x stroke volume

- Larger animals have slower heart rates, but larger stroke volumes
- Cardiac output increases with warmer body temperature, age during development, and increased activity level.

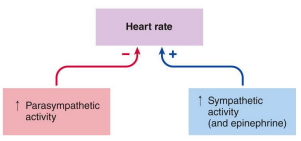


- Heart rate is determined by antagonistic regulation by the autonomic nervous system
 - Coordinated by the cardiovascular control center in the brain stem
 - ACh from vagus nerve binds to muscarinic receptors
 - Decreases heart rate (SA node)
 - Decreases excitability of the AV node
 - Shortens the plateau phase of atrial contractile cells
 - NE from sympathetic neurons and epinephrine from the adrenal medulla bind to β₁-adrenergic receptors
 - Increases heart rate
 - Reduces AV nodal delay
 - Speeds the spread of action potentials through the conduction pathway
 - Increases contractile strength of atrial and ventricular cells





Time (msec) (a) Autonomic influence on SA node potential



(b) Control of heart rate by autonomic nervous system

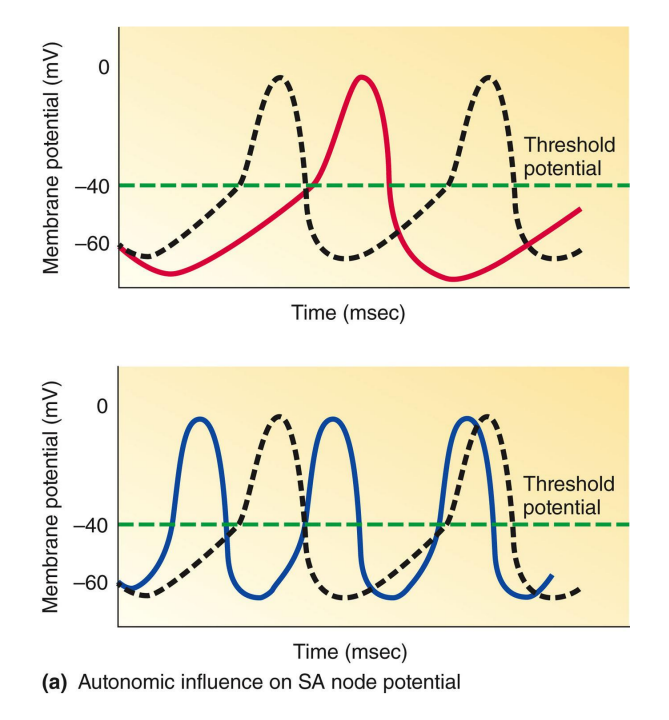
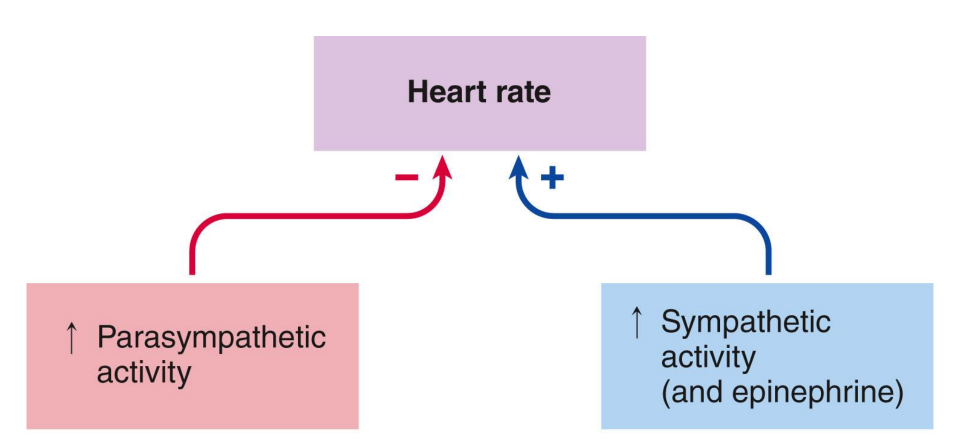


Figure 9-28a p416

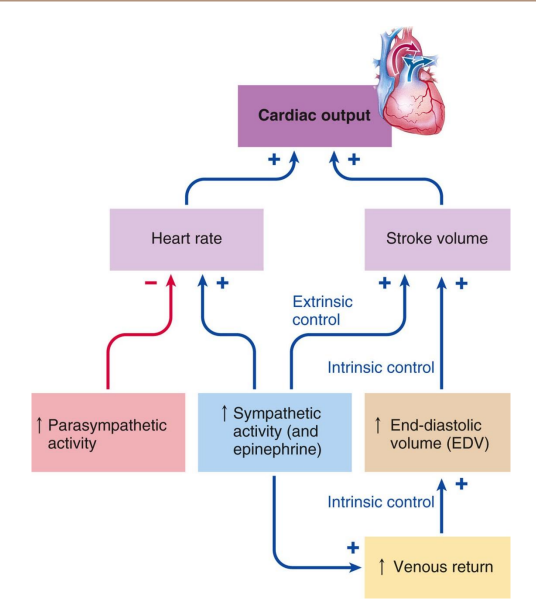


(b) Control of heart rate by autonomic nervous system



- Control of stroke volume
 - Intrinsic control
 - Direct correlation between end-diastolic volume (EDV) and stroke volume (SV)
 - Depends on the length-tension relationship of cardiac muscle
 - The greater the volume of blood entering the heart, the greater the volume ejected (Frank-Starling law of the heart)
 - Extrinsic control
 - Sympathetic stimulation enhances contractility of the heart
 - Sympathetic stimulation constricts veins, enhancing venous return and increasing stroke volume



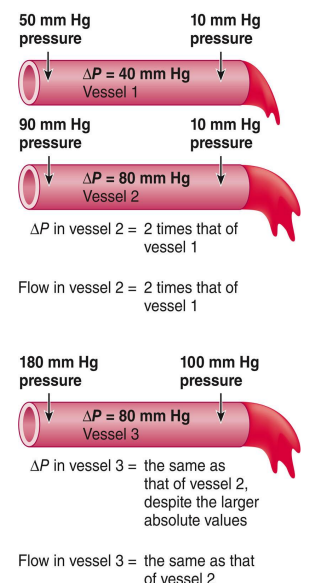




- The heart receives its blood supply through the coronary circulation
 - Heart muscle cannot extract oxygen or nutrients from blood
 within its chambers
 - Coronary arteries first evolved in active fishes
 - Branch off of brachial arteries leaving the gills
 - Coronary arteries branch off of the aorta in mammals
 - Coronary **blood flow increases** during activity
 - Dilation of coronary vessels is induced by adenosine
 - Adenosine is formed from ATP when oxygen supplies are low or cardiac activity is increased
 - Obstruction of coronary arteries is a leading cause of death in humans

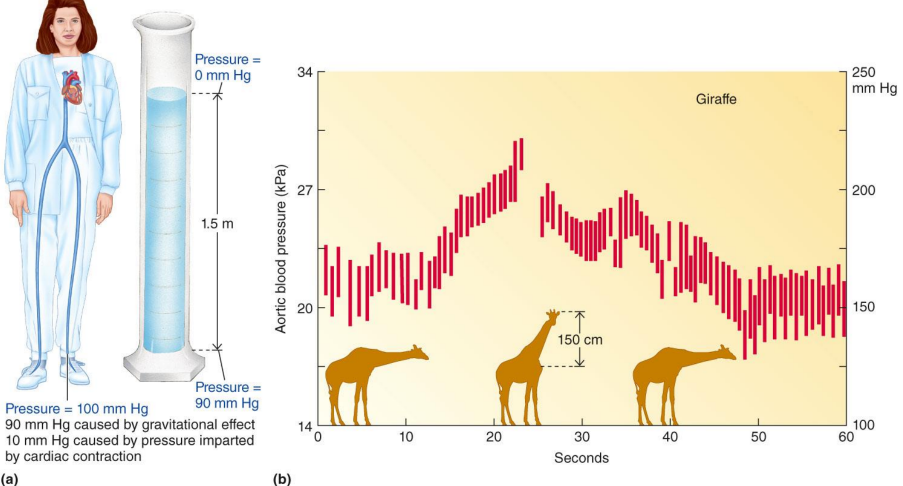


- Fluid flow obeys the hemodynamic flow law
 - $Q = \Delta P/R$
 - Q = flow rate of fluid through a vessel
 - ΔP = pressure gradient
 - R = resistance





- Fluid flow obeys the hemodynamic flow law
 Q = ΔP/R
 - Pressure gradient is the main driving force for flow through a vessel
 - Blood flows from an area of higher pressure to an area of lower pressure
 - Contraction of the heart generates pressure
 - Gravity also contributes to the pressure gradient
 - 70 mmHg pressure is required to push a column of water up 1 meter
 - A human needs an average driving pressure of 100 mmHg to overcome gravity
 - Fishes require a much lower driving pressure (~40 mmHg)

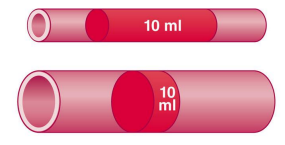


(a)

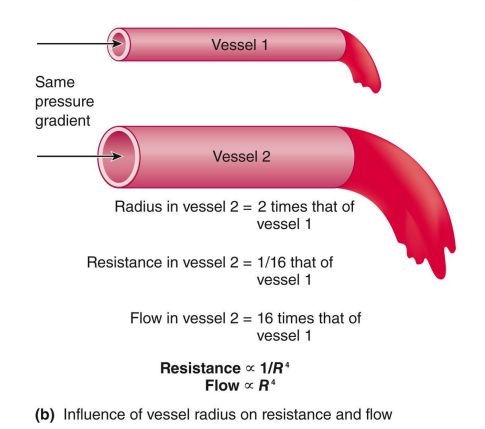


- Fluid flow obeys the **hemodynamic flow law** $Q = \Delta P/R$
 - Resistance is the hindrance to blood flow through a vessel caused by friction
 - $R = 8\eta L/\pi r^4$
 - R = resistance
 - η = viscosity of the fluid
 - L = length of the vessel
 - r = radius of the vessel
 - The major determinant of resistance to flow is the radius of the vessel
 - Resistance is inversely proportional to the 4th power of the radius

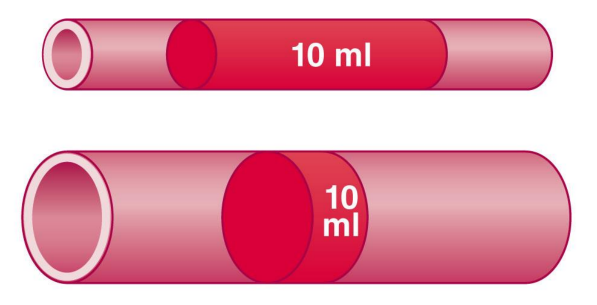




(a) Comparison of contact of a given volume of blood with the surface area of a small-radius vessel and a large-radius vessel







(a) Comparison of contact of a given volume of blood with the surface area of a small-radius vessel and a large-radius vessel

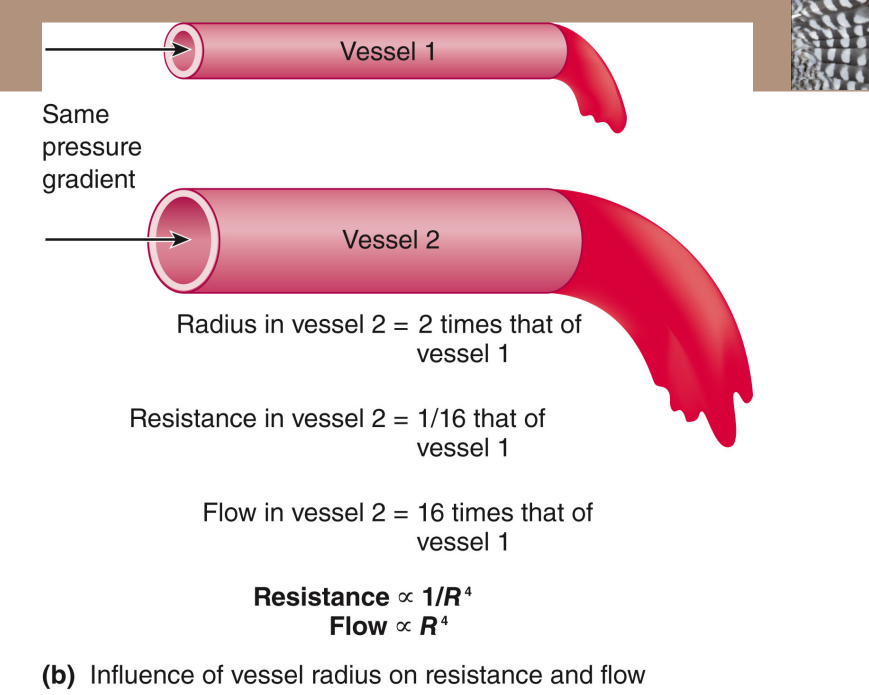


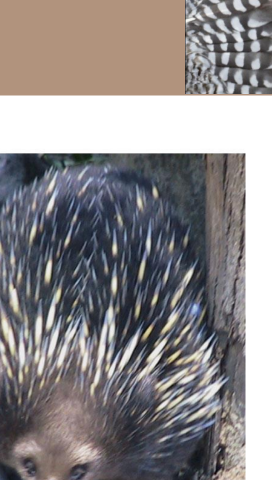
Figure 9-34b p421

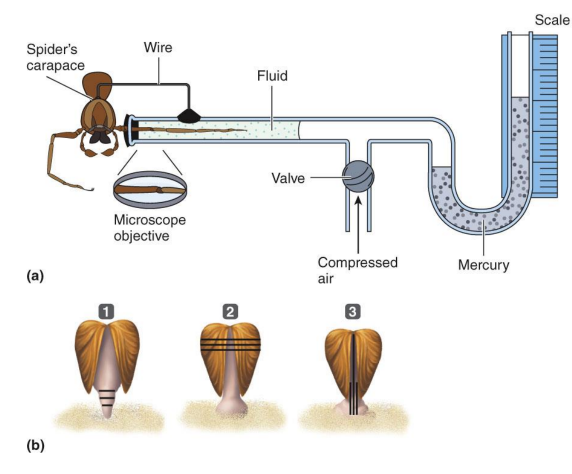


- Pressure can be used to exert force for noncirculatory functions
 - Movement
 - Extending legs of arachnids
 - Extending foot in bivalves
 - Ultrafiltration
 - Interaction between capillary blood and ECF
 - Initial process of urine formation

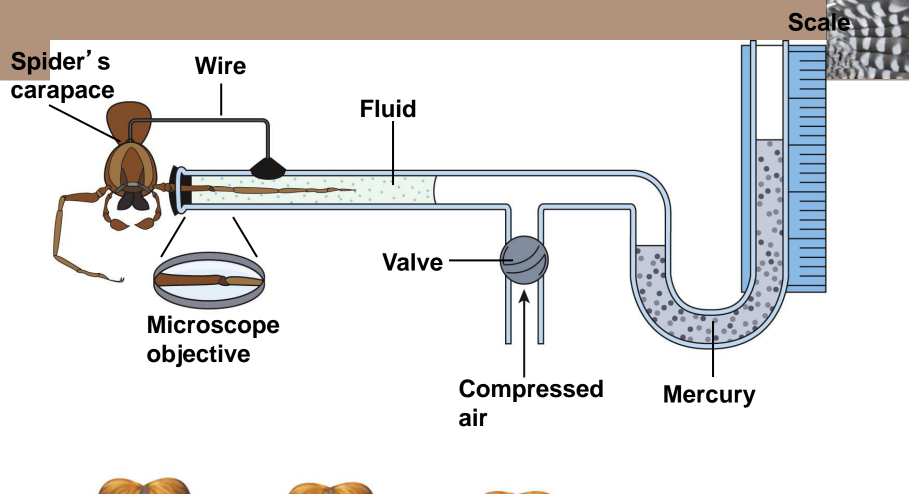
Erection

- Arousal of penis and clitoris
- Snout of echidna





(c)









9.9 Circulatory Pathways and Vessels: Open Circulation



- Some nonvertebrates have two or more separate ECF compartments
 - Example: Echinoderms have four different ECF compartments
- Non-cephalopod mollusks
 - Open circulation with myogenic chamber heart
 - Despite having no capillaries, there is directionality to hemolymph flow (e.g. clams control hemolymph for burrowing)

9.9 Circulatory Pathways and Vessels: Open Circulation



Crustacea

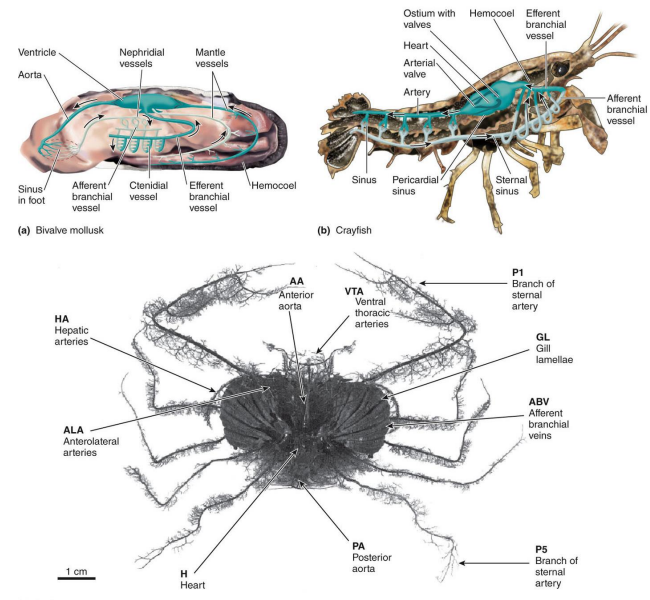
- Decapods have a well-developed circulatory system with a neurogenic chamber heart and numerous parallel structures
- Some decapods have capillary-like arteries opening into local lacunae and larger sinuses within target tissues
- All hemolymph flows to the gills before returning to the heart

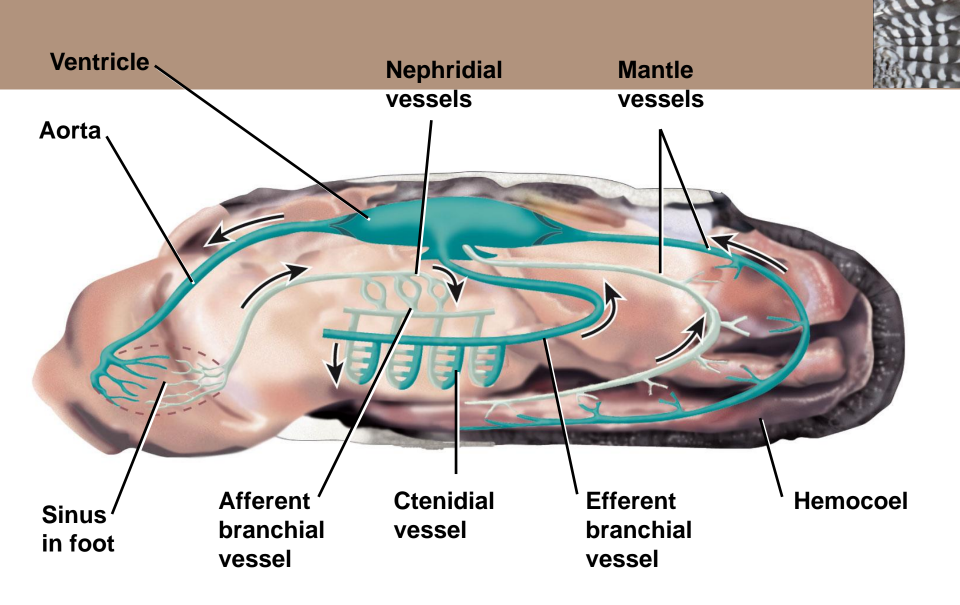
Insects

- Tubular heart and much less branching than crustaceans
- Insects do not rely on circulation for oxygen delivery

9.9 Circulatory Pathways and Vessels: Open Circulation

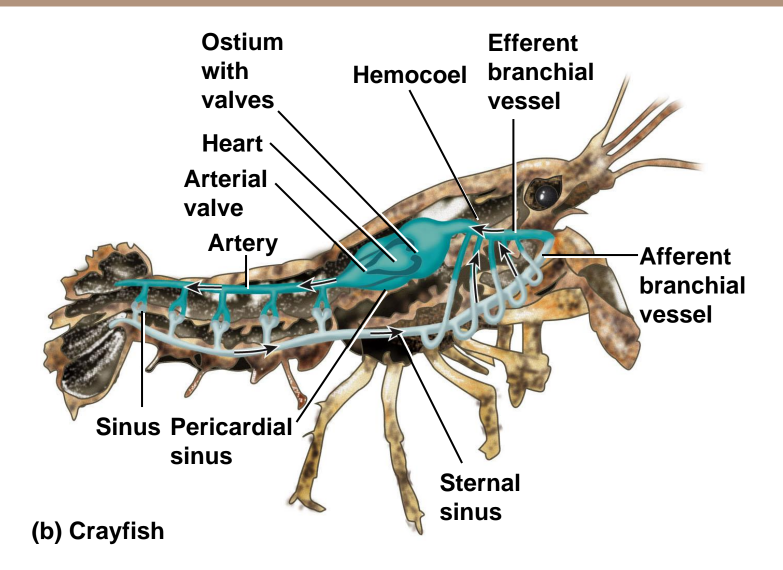


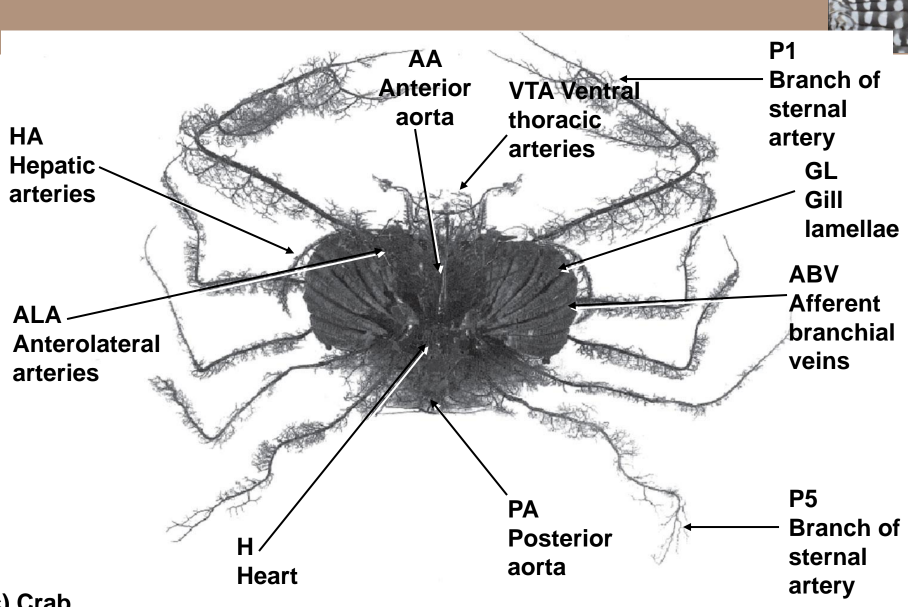




(a) Bivalve mollusk







(c) Crab

9.10 Circulatory Pathways and Vessels: Closed Circulation



Vertebrate circulatory system began as a single loop

Fishes

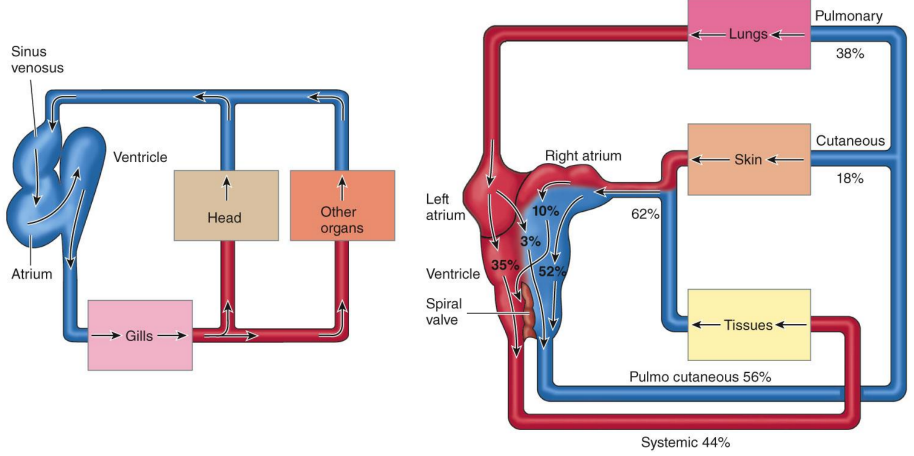
- Two-chambered heart (atrium and ventricle)
- Flow to the gills is in series with the rest of the circulation
- Parallel system distributes blood flow amongst the other organs

Amphibians

- Separate pulmonary circuit goes to lungs and skin
- Three-chambered heart with two atria
- Oxygenated and deoxygenated blood mix in single ventricle

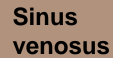
9.10 Circulatory Pathways and Vessels: Closed Circulation



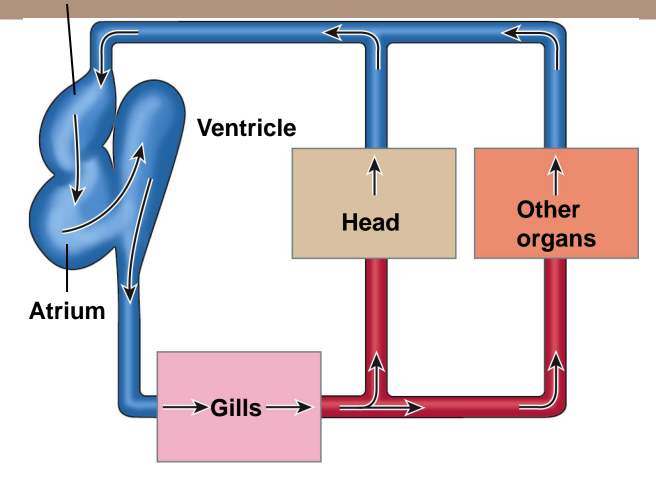


(a) Flow in a bony fish

(b) Flow in a bullfrog







(a) Flow in a bony fish

Figure 9-38a p426

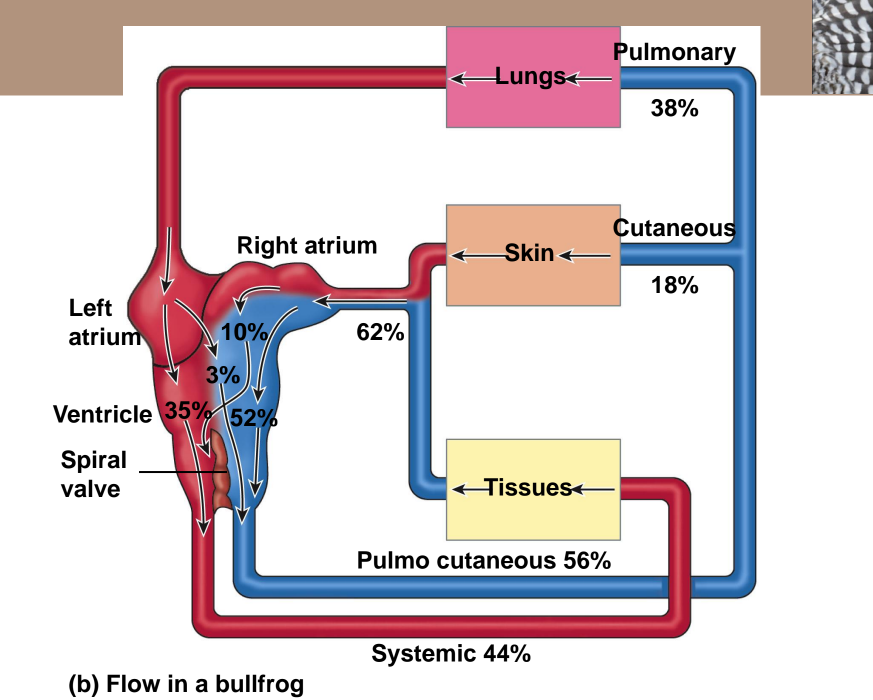
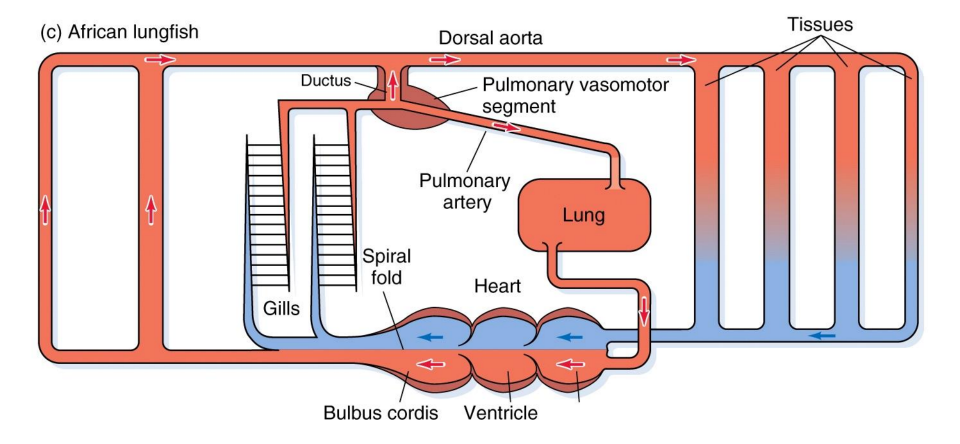


Figure 9-38b p426





9.10 Circulatory Pathways and Vessels: Closed Circulation



Reptiles

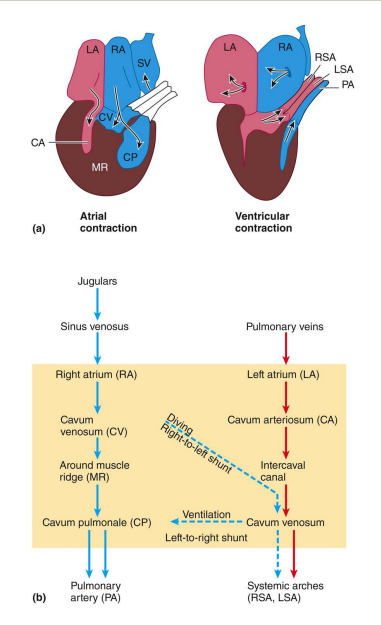
- Three-chambered heart
- Ventricle is divided into two large subchambers (cavum arteriosum and cavum pulmonale) by a thick muscle

Crocodiles

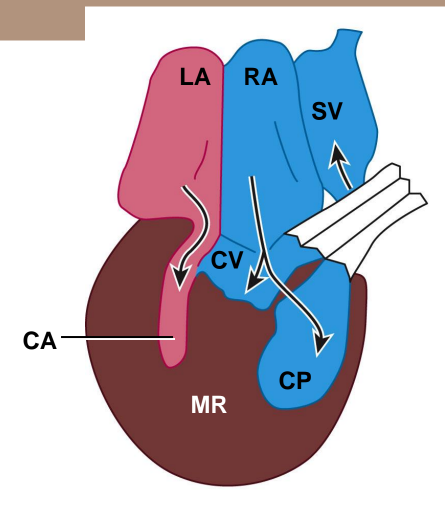
- Four-chambered heart with two atria and two ventricles
- Has two aortas, one exiting the right ventricle
- Foramen of Panizza connects the two aorta
 - Coglike valves between right ventricle and pulmonary arteries control diversion of blood flow
 - Adaptation for prolonged breath-holding during a dive

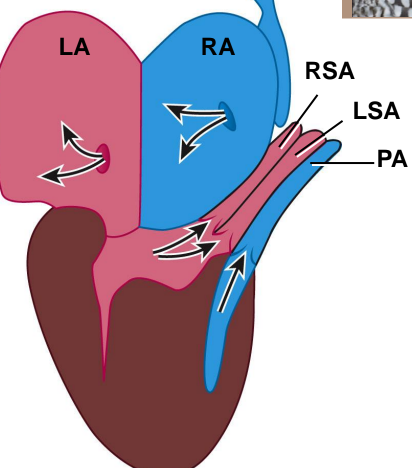
9.10 Circulatory Pathways and Vessels: Closed Circulation











Atrial contraction

Ventricular contraction

Figure 9-39a p427

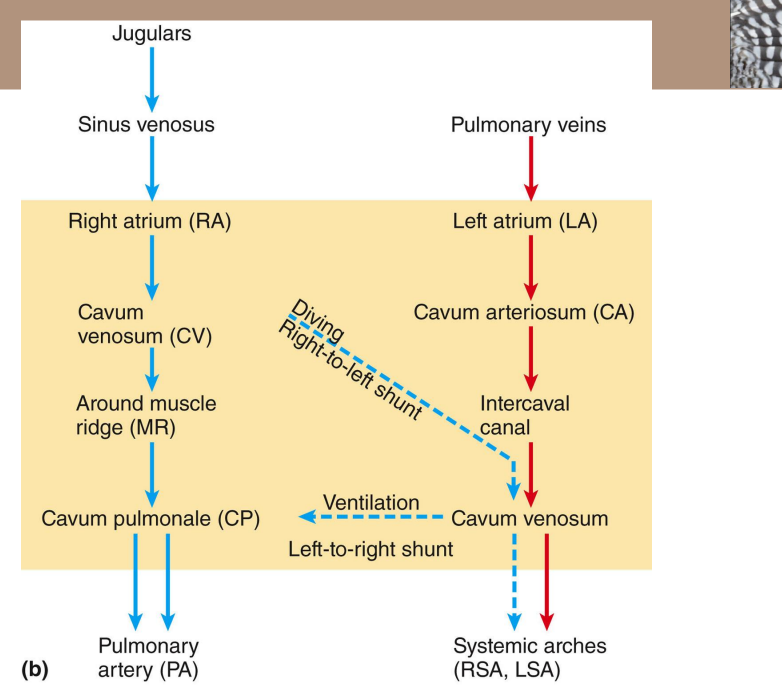
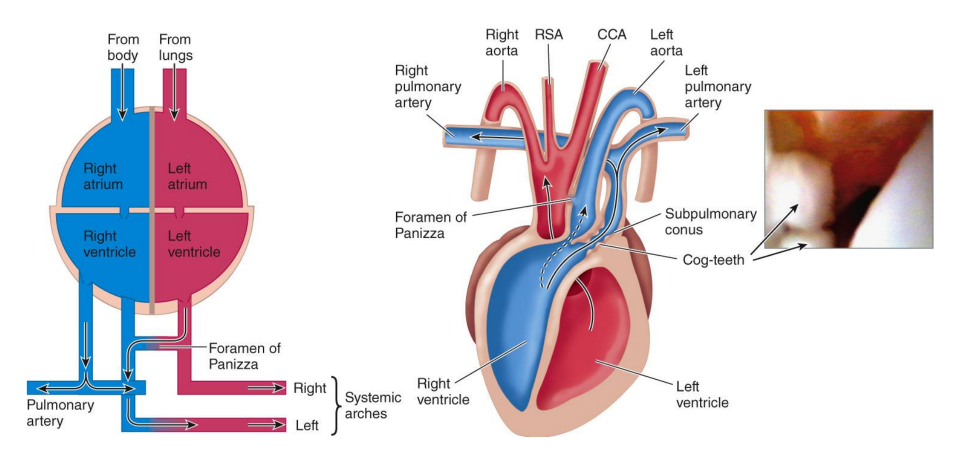


Figure 9-39b p427

9.10 Circulatory Pathways and Vessels: Closed Circulation





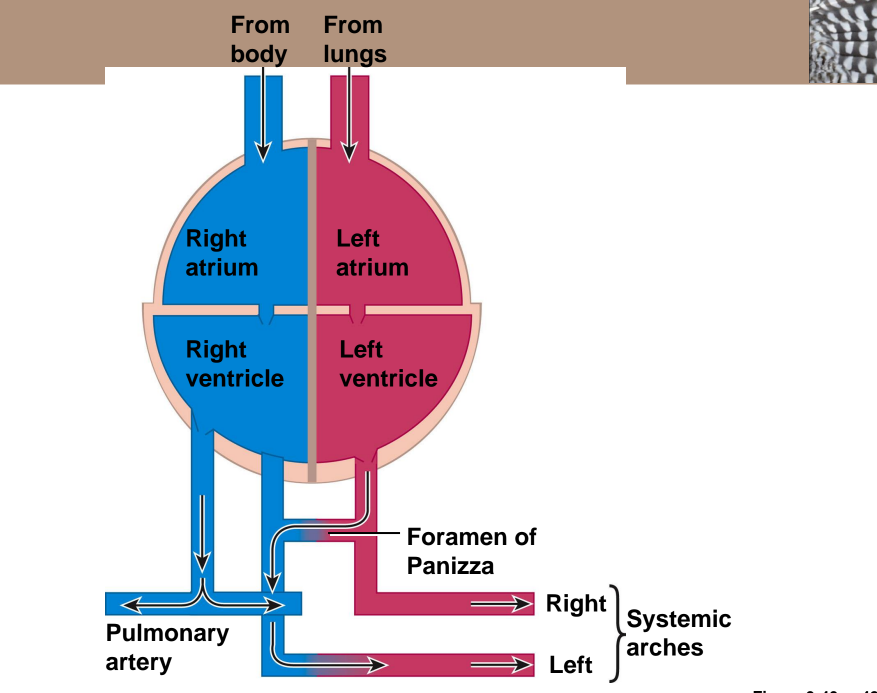
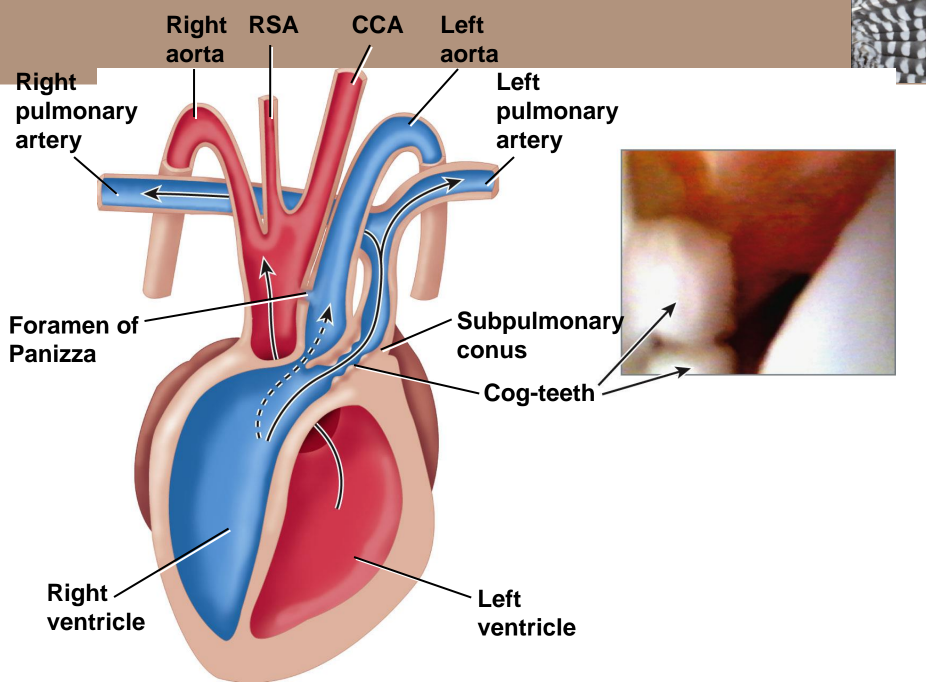


Figure 9-40a p428



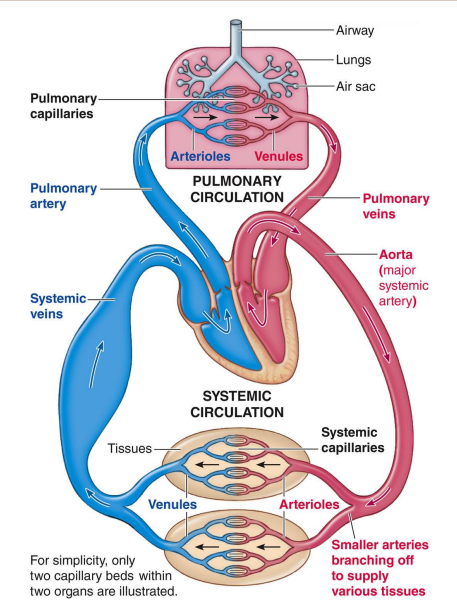
9.10 Circulatory Pathways and Vessels: Closed Circulation



- Birds and mammals
 - Four-chambered heart with no shunts
 - Complete separation of pulmonary and systemic flow evolved independently in birds and mammals to support their high endothermic metabolisms
 - Fetus of placental mammals has two bypasses since lungs are not functional
 - Foramen ovale -- opening in septum between right and left atrium
 - Ductus arteriosus -- connects pulmonary artery and aorta

9.10 Circulatory Pathways and Vessels: Closed Circulation

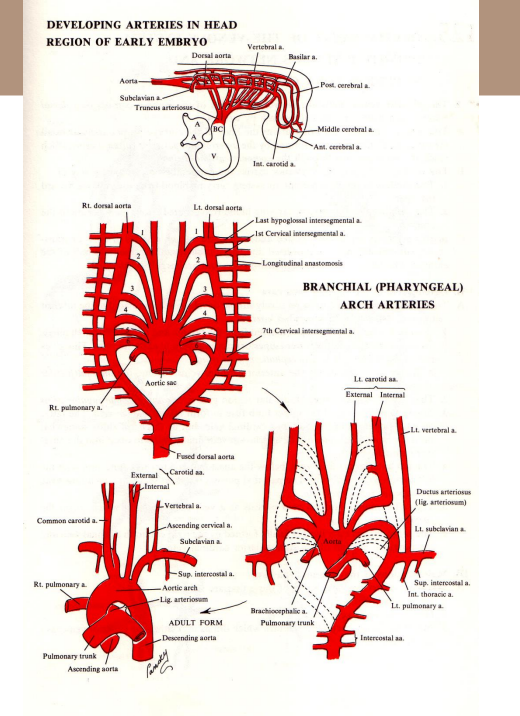




4. THE AORTIC ARCHES

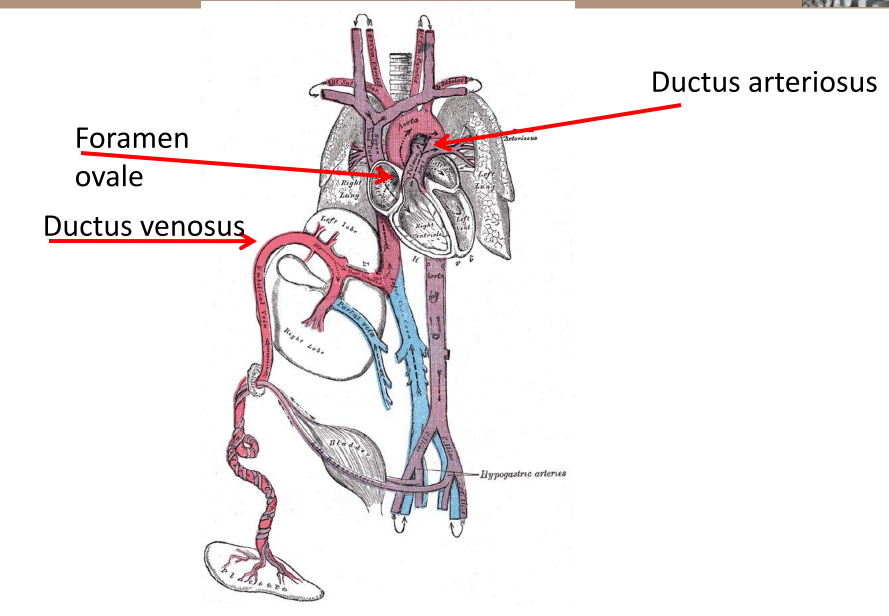
The first pair of aortic arches is formed by the curving of the ventral aorta into the primitive dorsal aorta. This arch is hidden in the mandibular arch and participates in formation of the *maxillary artery*, and contribute to the *external carotid artery*

- II. The second pair of aortic arches make their appearance in the middle of week 4. They cross the second branchial arches and give rise to the *stapedial* and *hyoid arteries*. (It should be noted that arches I and II regress rapidly and are not seen after day 31)
- III. The third pair of aortic arches make their appearance at the end of week 4. They give rise to the *common carotids* and *proximal portions of the internal carotid arteries*. The latter are the short cephalic prolongations of the primitive dorsal aortas and are associated with development and supply of the brain
- A. THE INTERNAL CAROTID ARTERIES are secondarily attached to the cranial portions of the dorsal aortas, which form the remainder of the carotid artery
- B. THE ORIGIN OF THE EXTERNAL CAROTID ARTERIES is controversial, but in later stages of development, they are found to sprout from aortic arch III. (Arch I, however, has been implicated in its developmental contribution)
- **IV.** The fourth pair of aortic arches make their appearance shortly after the third arches, at the end of week 4. Their development is different for the right and left sides
- A. ON THE RIGHT SIDE arch IV forms the proximal portion of the right subclavian artery and is continuous with the seventh segmental artery
 - 1. The caudal portion of the right primitive dorsal aorta disappears
 - 2. The distal portion of the subclavian artery forms from the right dorsal aorta and the right seventh intersegmental artery
- B. ON THE LEFT SIDE arch IV persists as the arch of the aorta, which grows significantly and is continuous with the primitive left dorsal aorta.
 - 1. The left subclavian artery (or seventh segmental) arises directly from the aorta
- C. THE SHORT PORTION of the right primitive ventral aorta, which persists between arches IV and VI, forms the *brachiocephalic arterial trunk* and the *first portion of the aortic arch*
- V. The fifth pair of aortic arches: in 50% of embryos, these arches are rudimentary vessels that degenerate with no derivatives. In fact, they may never even develop
- VI. The sixth pair of aortic arches make their appearance in the middle of week 5 and give rise to the *right* and *left pulmonary arteries*. After pulmonary vascularization is established, the communication with the corresponding primitive dorsal aorta regresses
- A. REGRESSION is total and complete on the right side. The proximal portion of the right arch forms the proximal part of right pulmonary artery; its distal portion degenerates
- B. THE PROXIMAL PORTION OF THE LEFT ARCH persists as the proximal part of the left pulmonary artery
 - The distal portion of the left arch, in which communication persists with the dorsal aorta until birth, forms the *ductus arteriosus* and diverts blood from the pulmonary artery to the aorta. Closure of the ductus arteriosus takes place in the neonatal period, and the functional duct becomes the anatomic *ligamentum arteriosum*
- C. THE DISTAL PORTIONS OF THE PULMONARY ARTERIES are derived from buds of the sixth aortic arches that grow into the developing lungs. After partitioning of the truncus arteriosus, the pulmonary arteries arise from the pulmonary trunk
- VII. Summary of the aortic arches: arch I regresses; arch II regresses; arch III forms the carotid system; arch IV forms the aortic arch (on the left) and the subclavian (on the right); arch V disappears; and arch VI forms the pulmonary arteries and the ductus arteriosus (on the left)





Fetus Heart: Before & After Birth



Sounds





http://www.easyauscultation.com/ cases-listing-details.aspx?caseID=7

Ventricular septal defect

Indomethacin



In humans, postnatal indomethacin can cause closure of the ductus arteriosus, and is used therapeutically when this structure remains patent in preterm neonates (Heymann et al., '76).

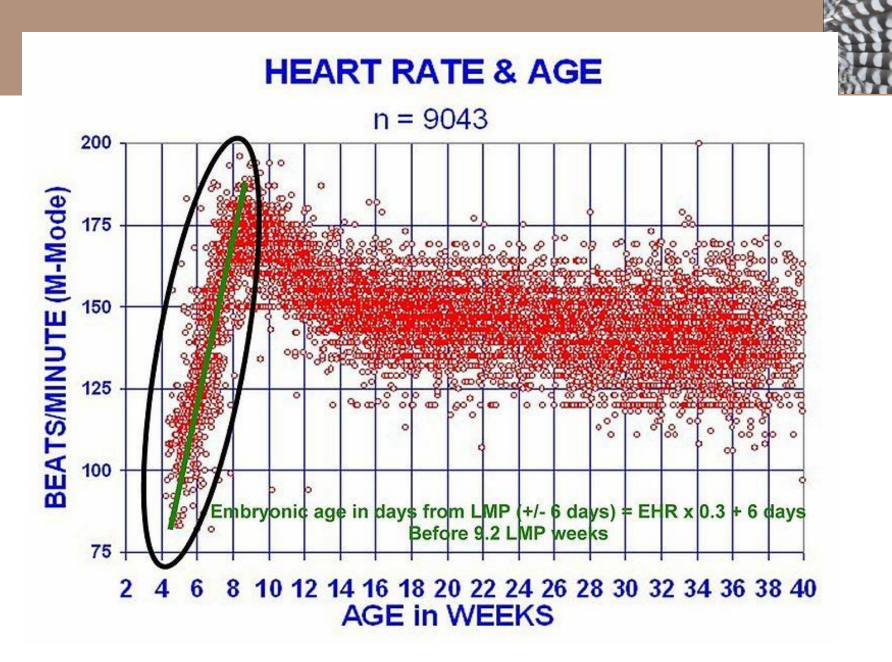
Ductal constriction can also occur in utero after maternal indomethacin administration (Moise et al., '88).

In addition, infants exposed to prenatal indomethacin were more likely to require surgical ligation of their PDA due to either a lack of response to postnatal indomethacin or a reopening of the duct after initial closure.

Pediatrics in Review Vol.28 No.4 April



- The human embryonic heart begins beating approximately 21 days after conception.
- The human heart begins beating at a rate near the mother's, about 75-80 BPM.
- The embryonic heart rate (EHR) then accelerates linearly for the first month of beating, peaking at 165-185 BPM during the early 7th week. This acceleration is approximately 3.3 BPM per day, or about 10 BPM every three days (increase of 100 BPM in the first month).
- Age in days = EHR(0.3) + 6After peaking at about 9.2 weeks after the normal menstrual period (LMP), it decelerates to about 150 BPM (+/-25 BPM) during the 15th week after the LMP.



Terry J. DuBose, M.S., RDMS; Director Diagnostic Medical Sonography Program



Congenital Heart Diseases: Neonate & Young Infant

- Significant congenital heart disease (CHD) may be diagnosed at virtually any age.
- Some conditions always are discovered in neonates; others rarely are identified during infancy.

Table 1. Mendelian Gene Syndromes Associated with Congenital Heart Anomalies*

	Frequency of Cardiac Anomaliest		
Etiologic Syndrome	All (%)	Distinctive or Most Common	Distinguishing Features
Autosomal Dominant			
Adams-Oliver syndrome	20	Left-sided obstruction (eg, COA, parachute MVP), TOF	Scalp cutis aplasia, terminal transverse limb defects
Alagille syndrome	95	(P)PS, TOF/TOF with PA, ASD, VSD	Bile duct paucity, chronic cholestasis, butterfly vertebrae, posterior embryotoxon
Char syndrome	60	PDA	Anomalies on fifth finger, supernumerary nipple
Cornelia de Lange syndrome	25	VSD, ASD, PS, TOF	Upper limb deficiency, GI anomalies
Holt-Oram syndrome	80	ASD ± other CVM, VSD, TA, TOF, PAPVC, conduction defect	Upper limb malformations
Neurofibromatosis	2	PSV, ASV, COA, HCM	Café au lait macules, optic glioma, scoliosis, pseudarthrosis, neurofibromas
Noonan syndrome	85	PSV, ASD, AVSD partial, COA, HCM	Short, webbed neck; pectus deformity; cryptorchidism
Rubinstein-Taybi syndrome	35	PDA, ASD, VSD, left-sided obstruction (eg, COA, HLHS)	Broad thumbs and great toes
Williams syndrome	60	SVAS, PS, other left-sided obstructions (eg, ASV, MS, COA)	Hypercalcemia, hypodontia, hypoplastic nails
Autosomal Recessive			
Ellis-van Creveld syndrome	60	AVSD, common atrium, ASD primum	Short limbs, polydactyly, hypoplastic nails, dental anomalies
Fryns syndrome	50	ASD, VSD, conotruncal	Diaphragmatic hernia, distal digital hypoplasia
Keutel syndrome	70	(P)PS	Short digits, mixed hearing loss, cartilage calcification
Smith-Lemli-Opitz syndrome	45	ASD, VSD, complete AVSD, TAPVC	Two- to three-toe syndactyly, cleft palate, lung anomalies, genital anomalies
X-linked Recessive			
Simpson-Golabi-Behmel	25	ASD; VSD; rare, variable	Macrosomia, cleft palate, supernumerary nipples,
syndrome		cardiomyopathy	hernias, hypospadias, poly/syndactyly
Suspected Gene Etiology Cardio-facio-cutaneous	75	ASD, HCM	Sparse, curly hair; low, rotated ears; hyperkeratosis
syndrome Hall-Hittner syndrome (CHARGE association)	80	Conotruncal/arch, assorted CVMs	Coloboma, choanal atresia, genital anomalies, ear
Costello syndrome	60	MVP, AV, thickening HCM, arrhythmia (atrial tachycardia)	Skin/joint laxity, fine/curly hair, deep palm creases, ulnar deviation, papillomata
PHACES syndrome	100	COA; IAA, A right; double, cervical aortic arch	Posterior fossa malformations, hemangiomas, eye anomalies
Ritscher-Schinzel syndrome (3C)	100	TOF, DORV, AVSD	Posterior fossa malformations, cleft palate, coloboma

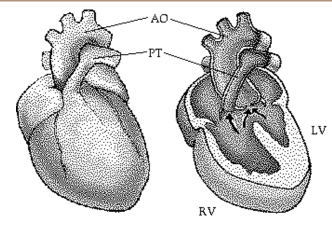
Table 2. Incidence of Most Common Cardiac Malformations*

	Prevalence (per 10,000 Births)			
Malformation	Metropolitan Atlanta Congenital Defect Program, 1995–1997	Baltimore-Washington Infant Study, 1981–1989		
Heterotaxy, L-TGA	1.6	1.4		
Outflow tract defects, total				
Tetralogy of Fallot	(4.7)	3.3		
D-TGA	2.4	2.3		
Double-outlet right ventricle	2.2	0.7		
Truncus arteriosus	0.6	0.5		
Atrioventricular septal defect				
With Down syndrome	2.4	2.3		
Without Down syndrome	1	1		
Ebstein anomaly	0.6	0.6		
Total APVC	0.6	0.7		
Right-sided obstruction		Not available		
Peripheral pulmonic stenosis	\sim	5.4		
Pulmonic stenosis, atresia	5.9	0.6		
Pulmonic atresia/intact septum	0.6	0.4		
Tricuspid atresia	0.3	0.4		
Left-sided obstruction		1.4		
Coarctation of the aorta	3.5	1.4		
Hypoplastic left heart	2.1	0.8		
Aortic valve stenosis	0.8	Not available		
Aortic arch atresia or hypoplasia	0.6	Not available		
Septal defects	210	11.2		
Ventricular septal defect	24.9	3.2		
Atrial septal defect	10	0.9		
Patent ductus arteriosus	8.1 9.7			
Other major heart defects	90.2	48.4		
Total	30.2			

APVC=anomalous pulmonary venous connection, TGA=transposition of the great arteries. *Reprinted from Lin AE, Holly HA. Genetic epidemiology of cardiovascular malformations. *Progr Pediatr Cardiol*. 2005;20:113–126 with permission from Elsevier.

Tetralogy of Fallot





This condition results from a single error: the conus septum develops too far anteriorly giving rise to two unequally proportioned vessels- - a large aorta and a smaller stenotic pulmonary trunk.

The four main characteristics of Tetralogy of Fallot are:

(1) pulmonary stenosis

(2) ventricular septal defect (VSD) of the membranous portion (the septum is displaced too far anteriorly to contribute to the septum)

(2) overriding parts (the parts straddles the VCD)



Tricuspid Atresia:

Total Correction: mortality less than 3%

Transposition of the great arteries Total Correction: mortality less than 2%

Pulmonic stenosis Total Correction: mortality less than 1%

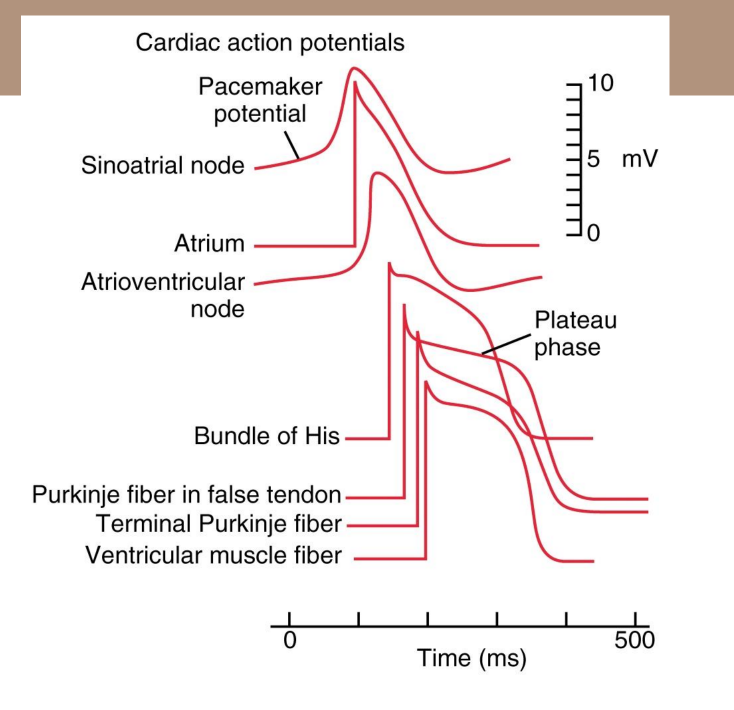
Truncus Arteriosus (various types) Mortality is > 10%

Hypoplastic left heart syndrome Mortality ~10%

ECG: Neonate

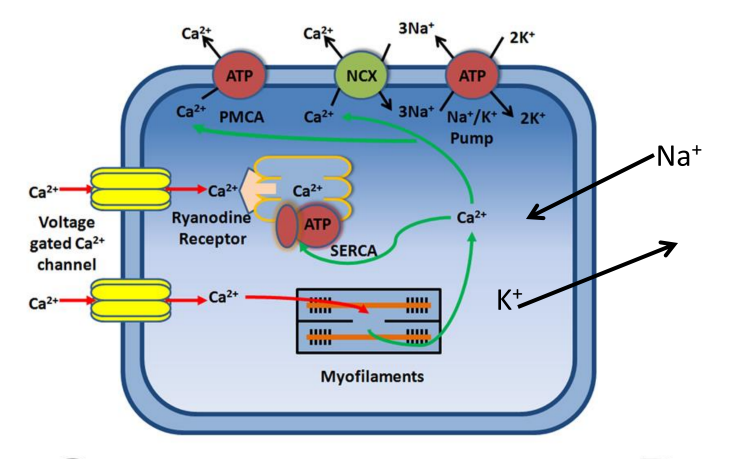


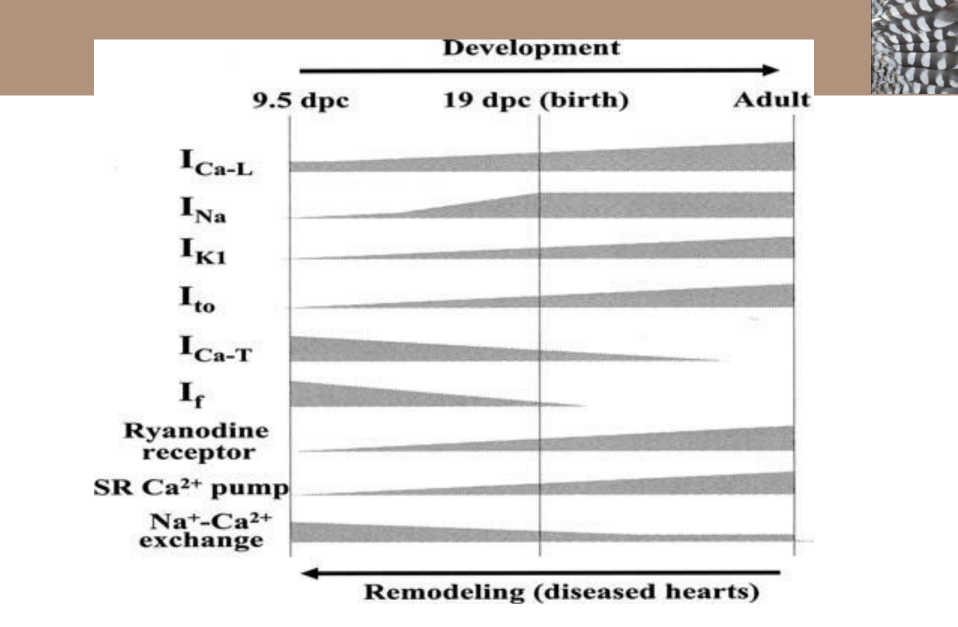
- Arrhythmias in fetuses and newborns are relatively common, occurring in up to 90% of newborns and in 1% to 3% of pregnancies (NeoReviews Vol.9 No.6 2008 e242,2008 American Academy of Pediatrics, Fetal and Neonatal Arrhythmias)
- Weak arterial pulses and right heart overload in the electrocardigram suggested the diagnosis of hypoplasia of the left heart. Impaired coronary perfusion to portions of the right and left ventricular myocardium. Pulmonary vasoconstriction from hypoxia. Myocardial ischemia on the electrocardiogram (The Journal of Pediatrics Volume 81 (2): 243-250)
- SIDS: A prolonged QT interval may be an important cause for SIDS. (Schwartz et al., The New England Journal of Medicine, 1998 338(24):1709-1714.





MUSCLE CONTRACTION





Brugada syndrome : Genetic disease, abnormal ECG sudden cardiac death (Sudden Unexpected Death Syndrome -SUDS). First described in 1992, ventricular fibrillation mutation in Na+ ion channel



• Recent evidence indicates that between 5 and 15% of SIDS cases carry potentially lethal **loss-of-function mutations in cardiac channelopathy genes**.

(Future Cardiol. 2009 Mar;5(2):201-7. Sudden infant death syndrome and cardiac arrhythmias. Morris JA, Harrison L, Brodison A, Lauder R. Department of Pathology, Royal Lancaster Infirmary, Lancaster LA1 4RP, UK.)

• Morphological changes in the mitochondrial network likely accompany the uncoupling with mitochondrial fission dampening the signals leading to **cardiomyocyte death**.

(J Bioenerg Biomembr. 2009 Apr;41(2):133-6. Uncouple my heart: the benefits of inefficiency.

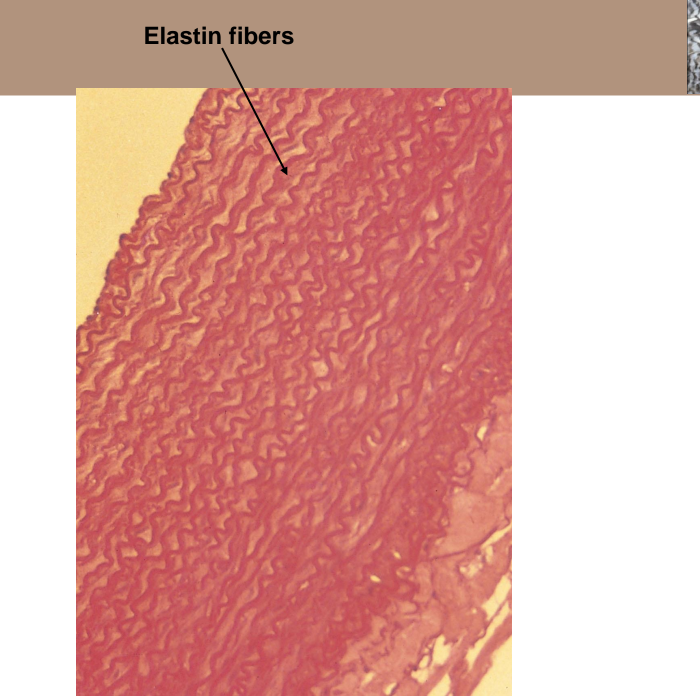
Sonographer: Pediatric echocardiography

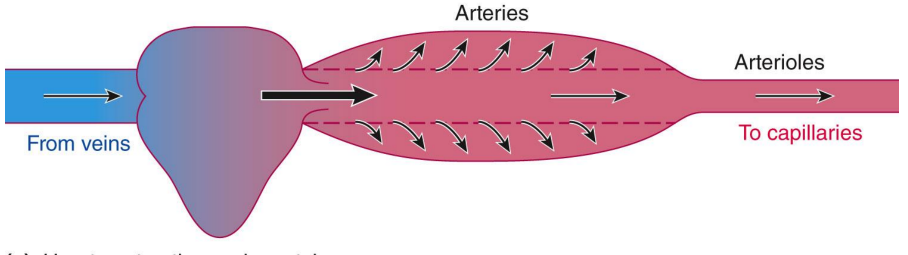




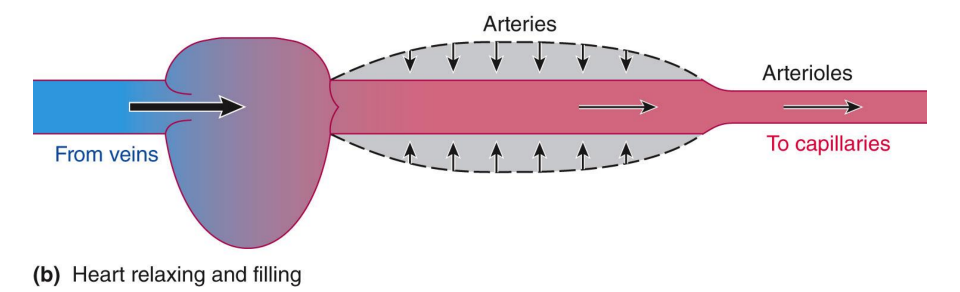
- Arteries provide rapid passage of blood from the heart to the tissues
 - Large radii offer little resistance to flow
 - Blood flows at high velocity
- Arteries serve as pressure reservoirs
 - Arteries' elasticity enables them to expand during ventricular systole
 - Elastic recoil is the driving force for continued flow of blood during diastole

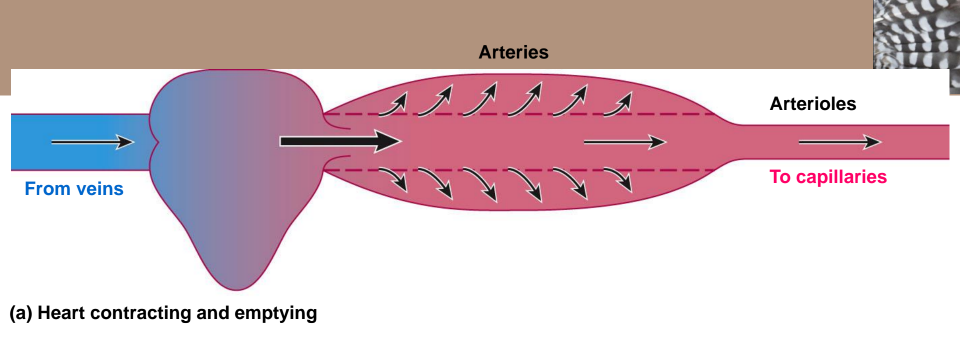






(a) Heart contracting and emptying





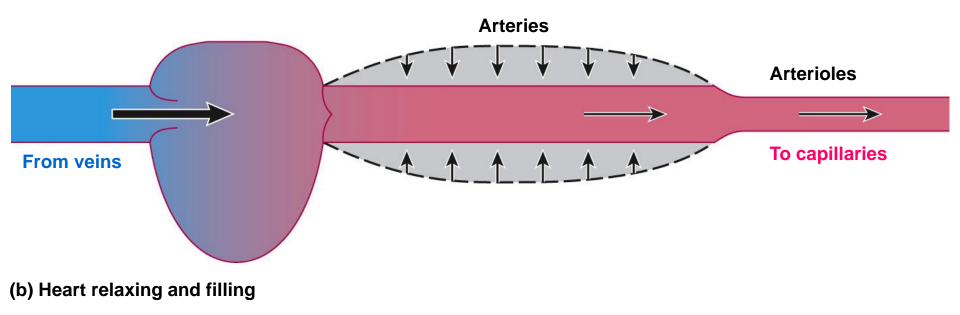
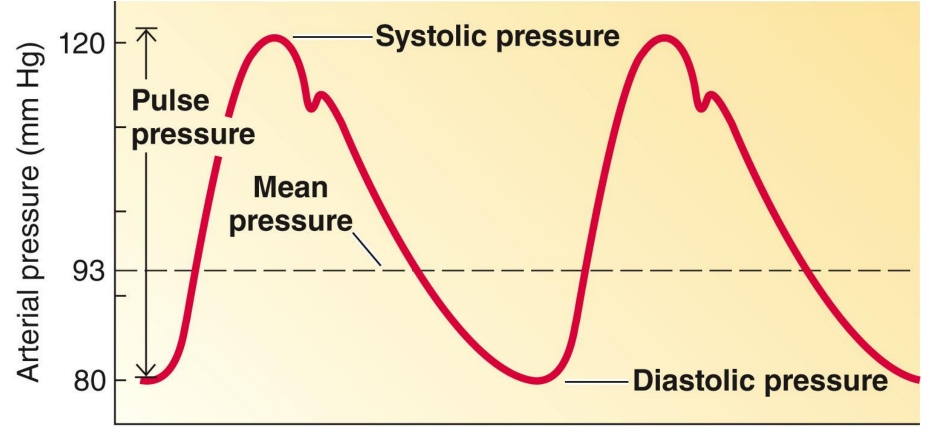


Figure 9-43 p430



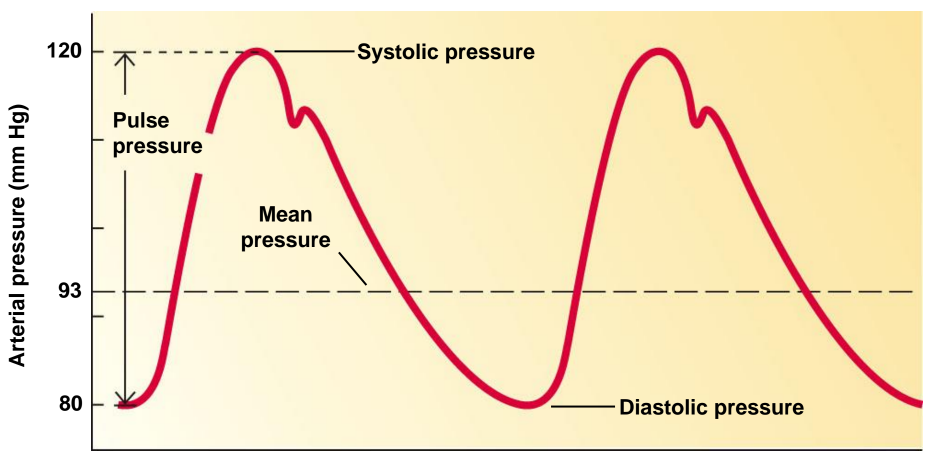
- Arterial blood pressure
 - Maximum pressure exerted on the arteries (systolic blood pressure) averages 120 mmHg in humans
 - Minimum pressure (diastolic blood pressure) averages 80 mmHg
 - Arterial blood pressure is expressed as a fraction (e.g. 120/80 mmHg)
 - Mean arterial pressure is the main driving force of blood flow

Mean arterial pressure = diastolic pressure + 1/3 (systolic pressure - diastolic pressure)



Time (msec)



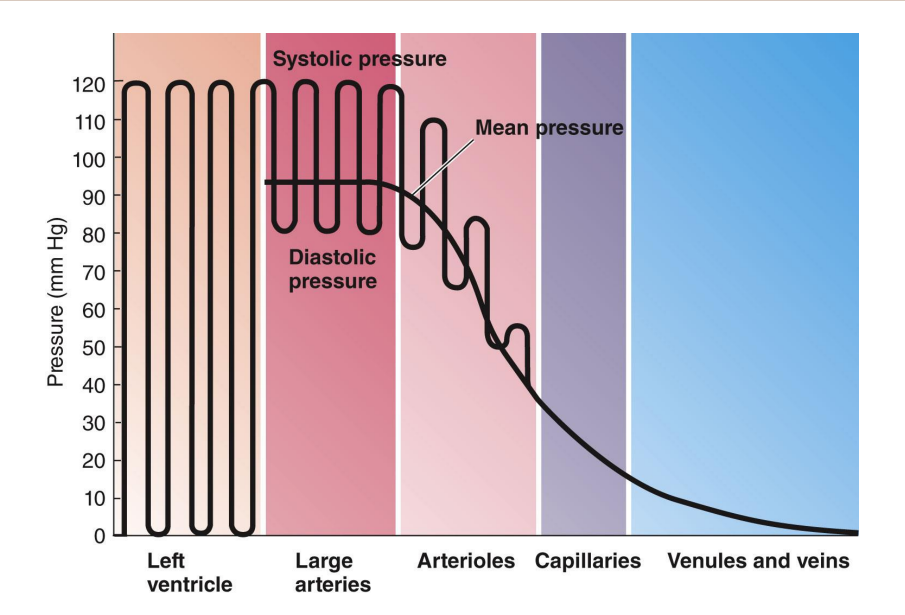


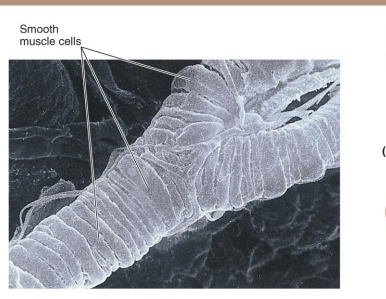
Time (msec)

Figure 9-44 p432

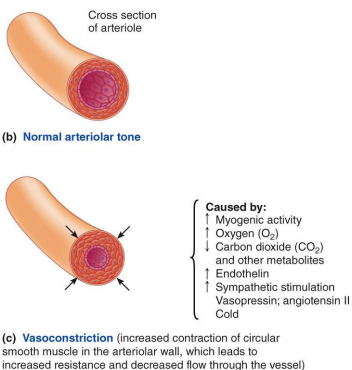


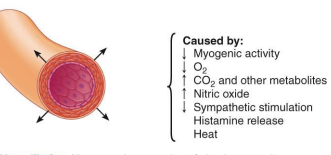
- Arterioles are the major resistance vessels
 - Radii of arterioles are small enough to offer considerable resistance to flow
 - Large drop in blood pressure through the arterioles
 - Mean arterial pressure of 93 mmHg drops to 37 mmHg where blood enters the capillaries
 - Eliminates pulsatile pressure swings
 - Thick layer of smooth muscle is innervated by sympathetic nerve fibers
 - Vasoconstriction results from smooth muscle contraction —-> decreased radius, increased resistance
 - Vasodilation results from smooth muscle relaxation —-> increased radius, decreased resistance





(a) Scanning electron micrograph of an arteriole showing how the smooth muscle cells run circularly around the vessel wall



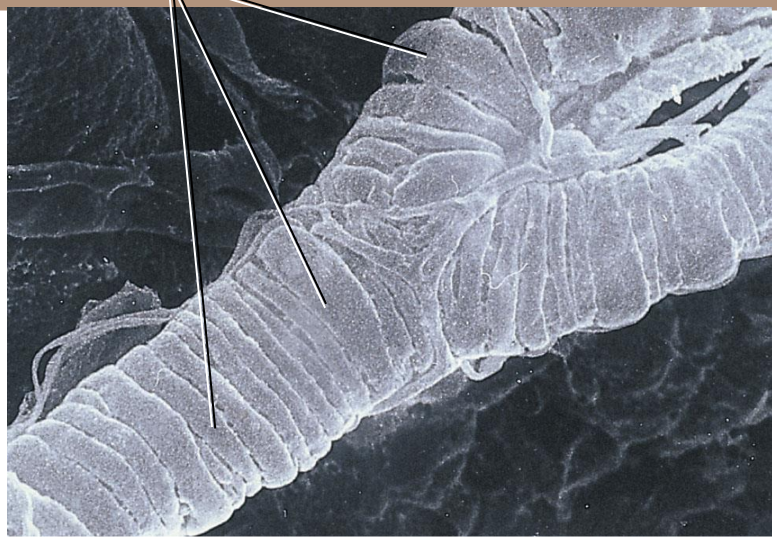


(d) Vasodilation (decreased contraction of circular smooth muscle in the arteriolar wall, which leads to decreased resistance and increased flow through the vessel)



Smooth muscle cells





(a) Scanning electron micrograph of an arteriole showing how the smooth muscle cells run circularly around the vessel wall

Figure 9-46a p433

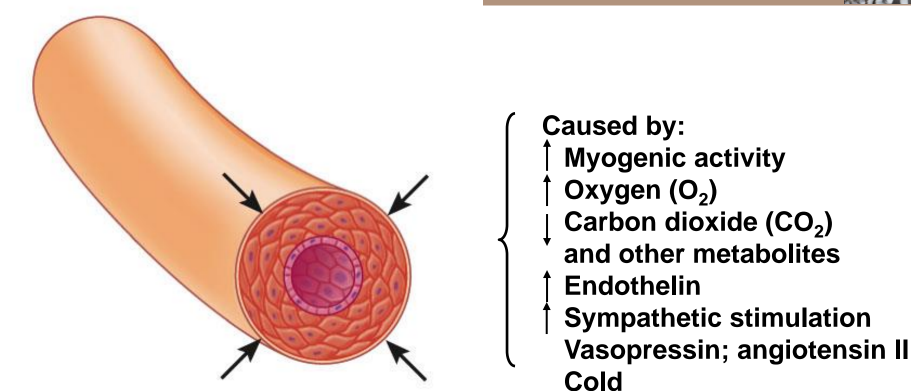


Cross section of arteriole

(b) Normal arteriolar tone

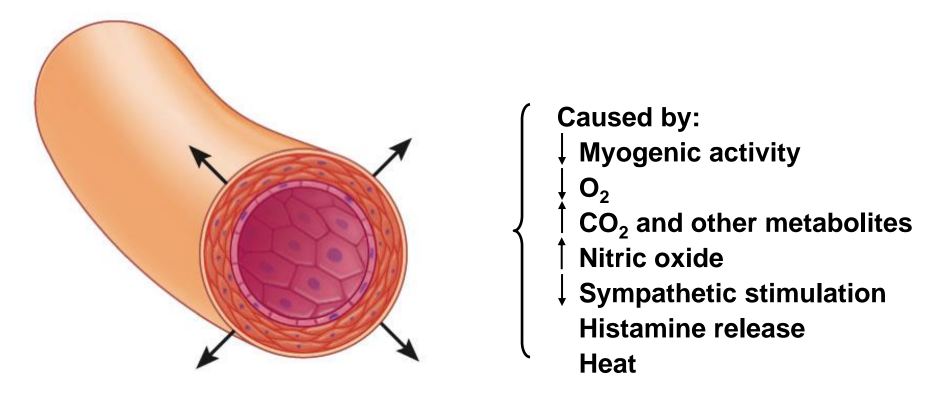
Figure 9-46b p433





(c) Vasoconstriction (increased contraction of circular smooth muscle in the arteriolar wall, which leads to increased resistance and decreased flow through the vessel)





(d) Vasodilation (decreased contraction of circular smooth muscle in the arteriolar wall, which leads to decreased resistance and increased flow through the vessel)

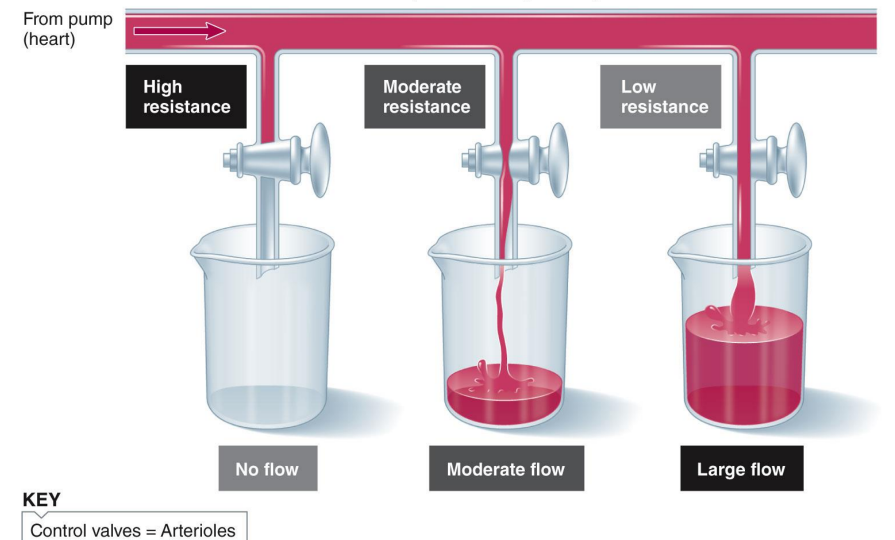


- Distribution of blood flow
 - Amount of blood flow received by each organ is determined by the number and diameter of its arterioles
 - Local (intrinsic) controls are changes within a tissue that alter the radii of arterioles
 - Local metabolic changes produce relaxation of arteriolar smooth muscle to increase blood flow to the organ (active hyperemia)
 - Histamine release causes vasodilation as an inflammatory response
 - Exposure to heat or cold
 - Stretch produces myogenic vasoconstriction

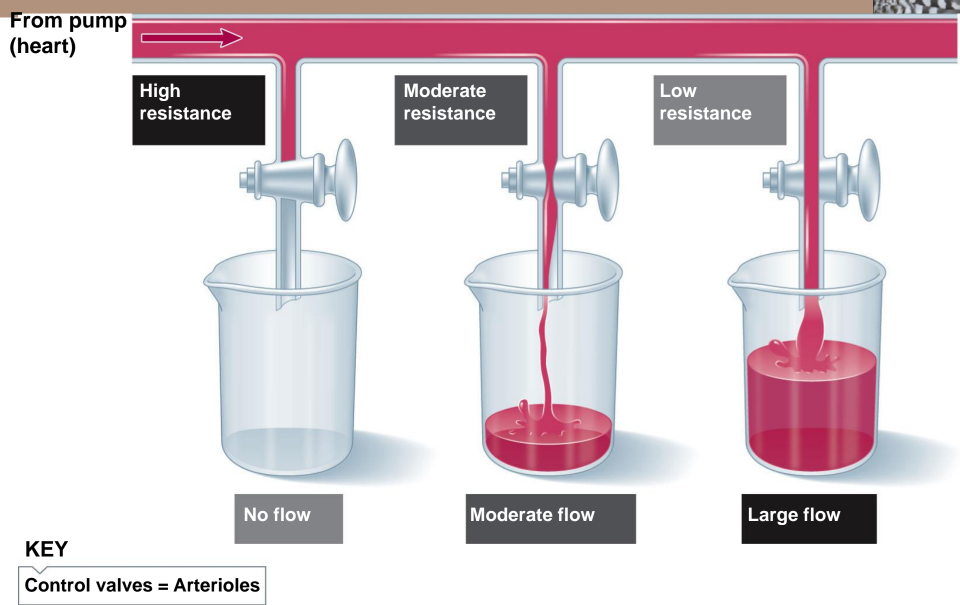
9.12 Circulatory Vessels: Arterioles



Constant pressure in pipe (mean arterial pressure)



Constant pressure in pipe (mean arterial pressure)

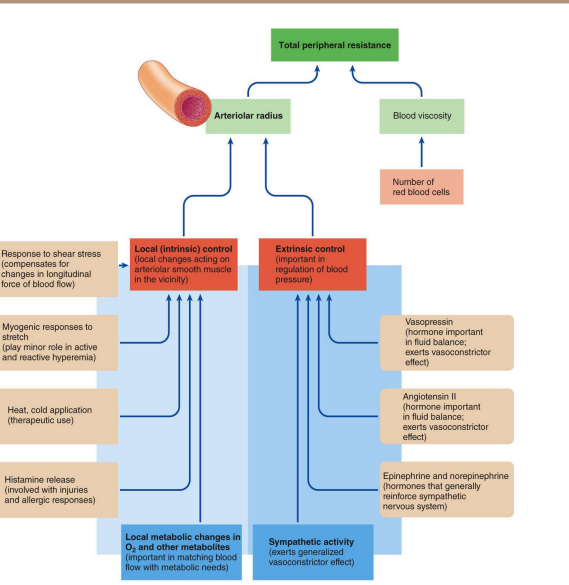


9.12 Circulatory Vessels: Arterioles

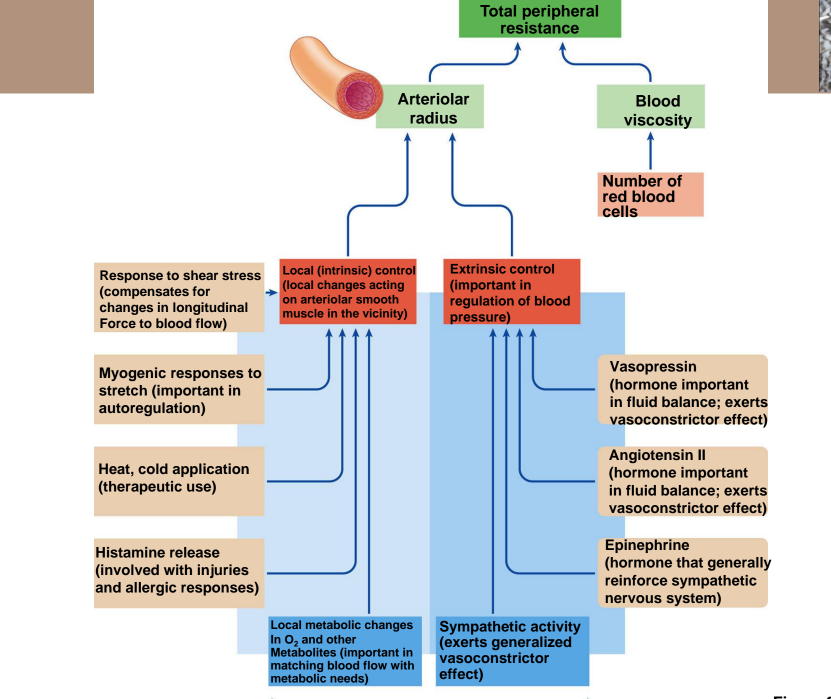


- Extrinsic control includes neural and hormonal influences with the sympathetic nervous system dominating
 - Sympathetic stimulation redistributes blood flow to the heart and skeletal muscle at the expense of other organs
 - Vasoconstriction of most arterioles due to activation of α₁-adrenergic receptors by NE or epinephrine
 - Lung and brain arterioles do not have α₁-receptors and do not constrict
 - Epinephrine activates β₂-adrenergic receptors in arterioles of skeletal muscle and heart to cause vasodilation
 - Mean arterial pressure is the product of cardiac output and total peripheral resistance

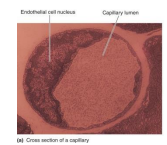
9.12 Circulatory Vessels: Arterioles



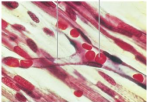
Major factors affecting arteriolar radius



- Capillaries maximize diffusion rates for exchange of materials
 - Diffusion distance is short
 - Capillary walls are thin
 - Capillaries are narrow
 - Capillaries branch extensively





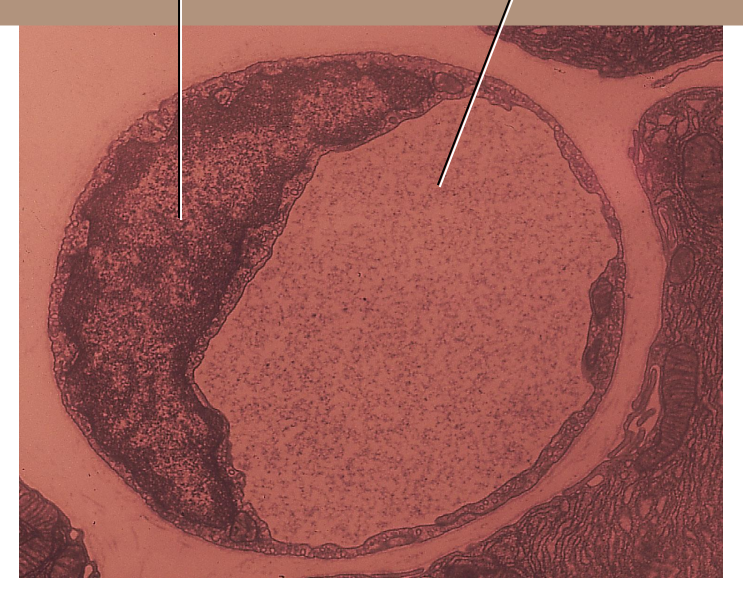


- Surface area is large
- Permeability is high
 - Pores between capillary endothelial cells
 - Tight junctions in brain capillaries form protective blood-brain barrier
 - Large pores in liver capillaries allow passage of proteins

Endothelial cell nucleus

Capillary lumen



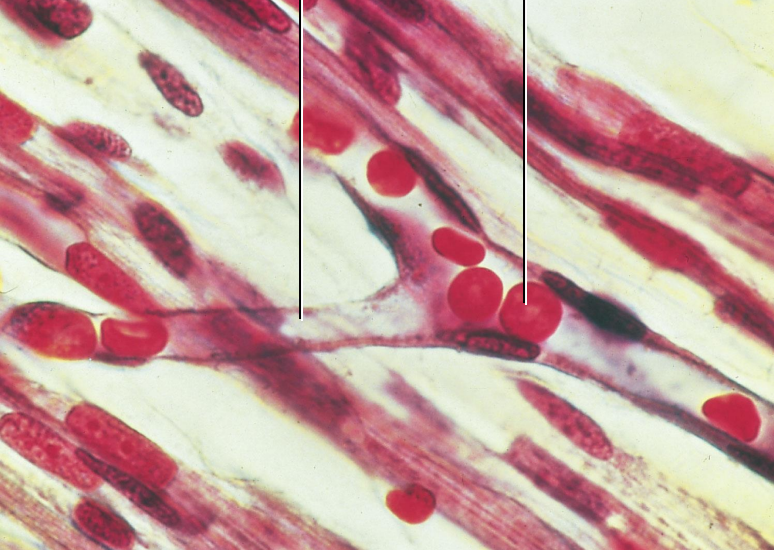


(a) Cross section of a capillary

Capillary

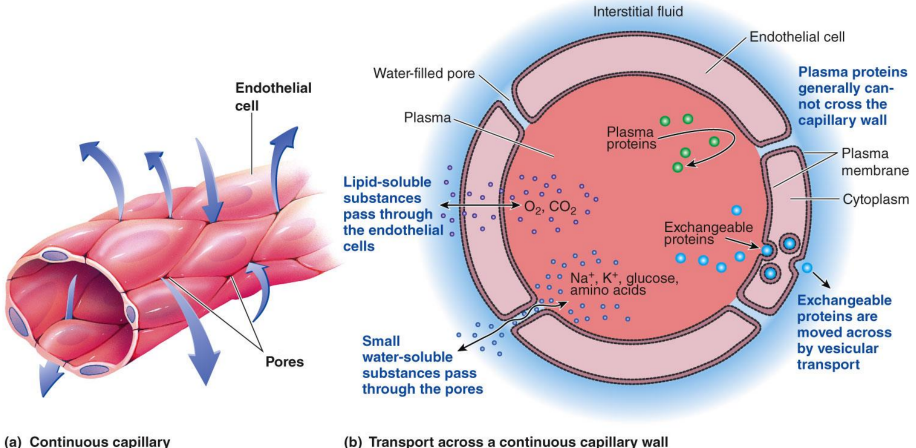
Red blood cell



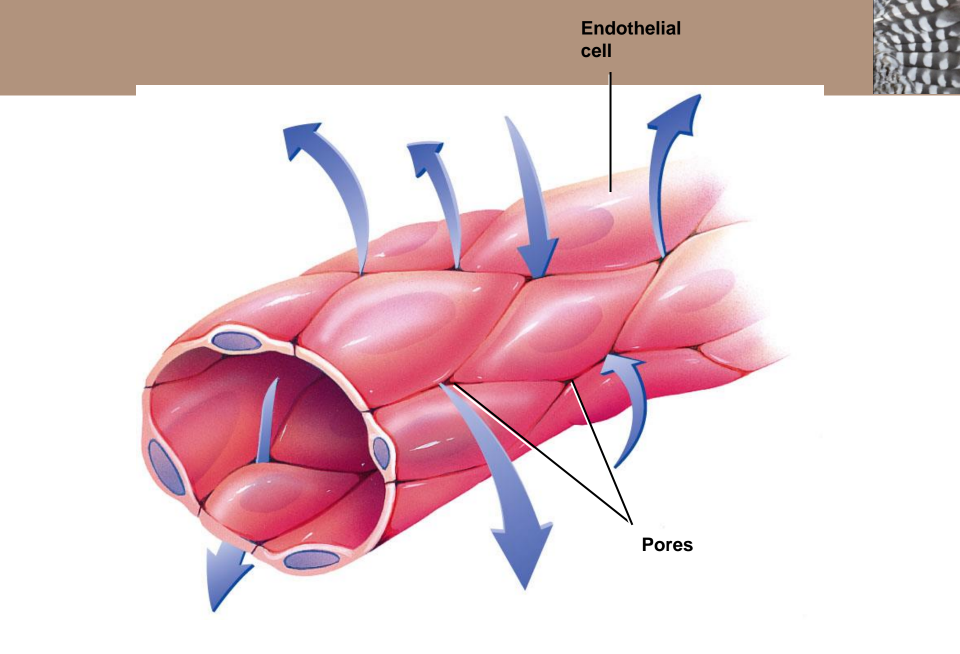


(b) Capillary bed

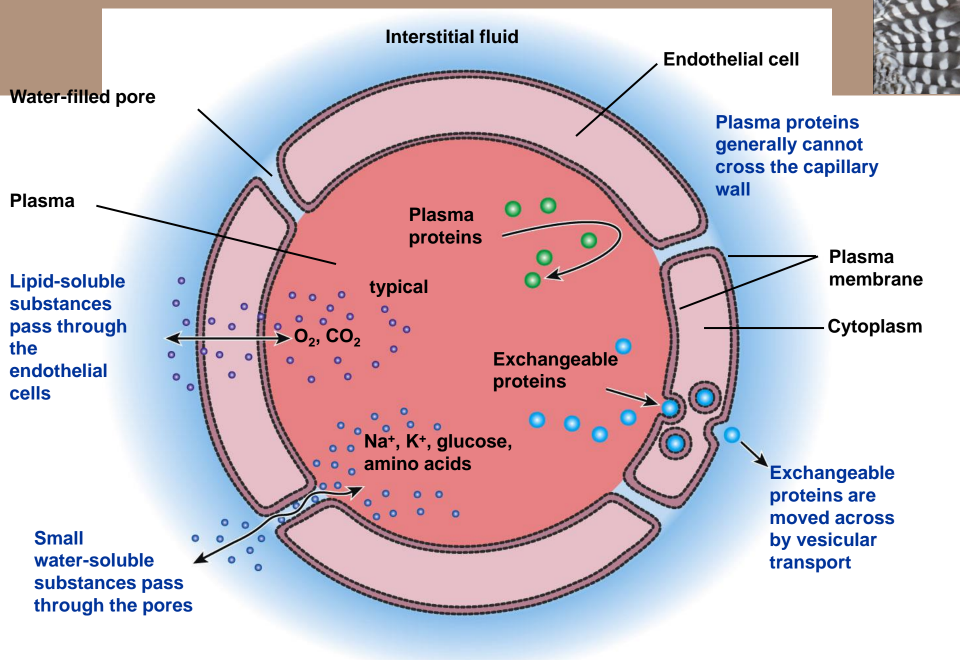




(b) Transport across a continuous capillary wall



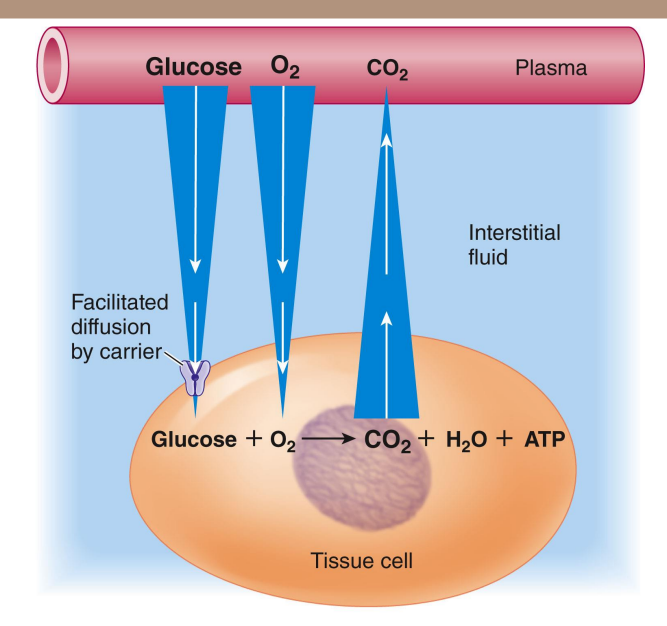
(a) Typical capillary





- Diffusion follows concentration gradients between blood and the surrounding cells
 - Cells consume glucose and oxygen
 - Cells produce CO₂ and other metabolic wastes
 - Precapillary sphincters controlling flow through individual capillaries are sensitive to local metabolic changes
 - Blood velocity is slowest due to the very large cross-sectional area of the capillaries





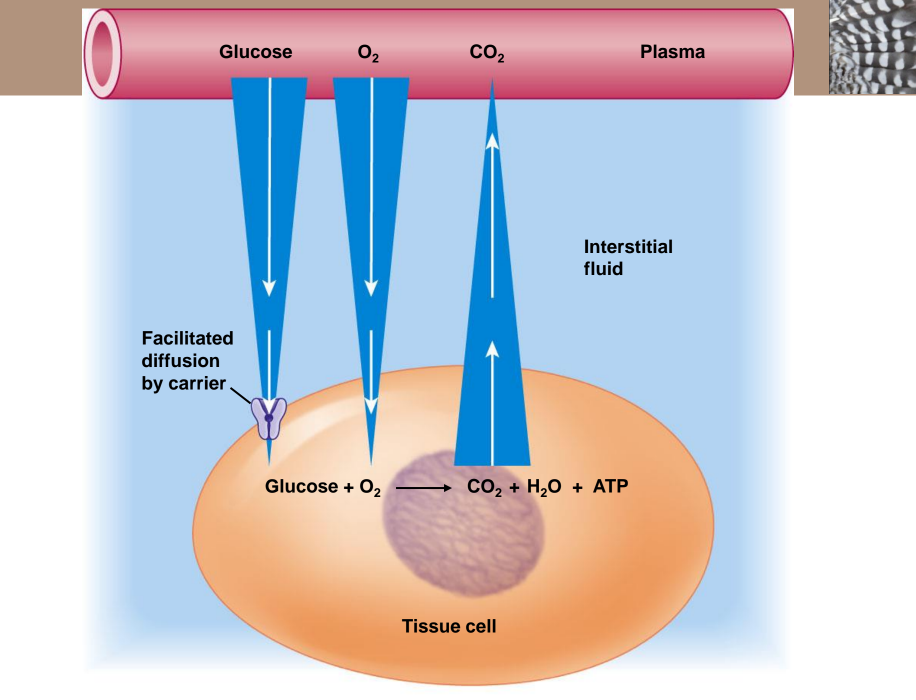
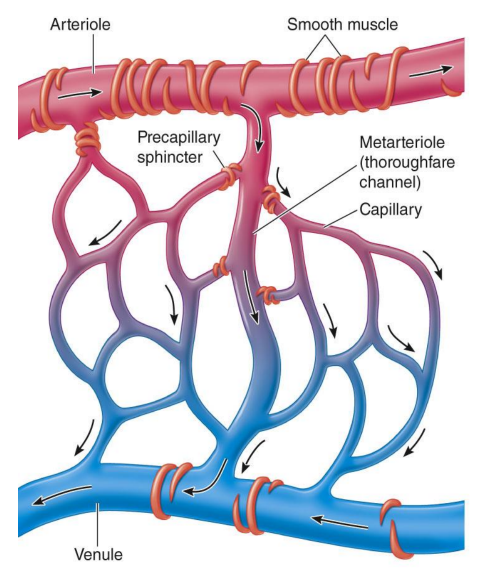


Figure 9-51 p440





⁽a) Sphincters relaxed

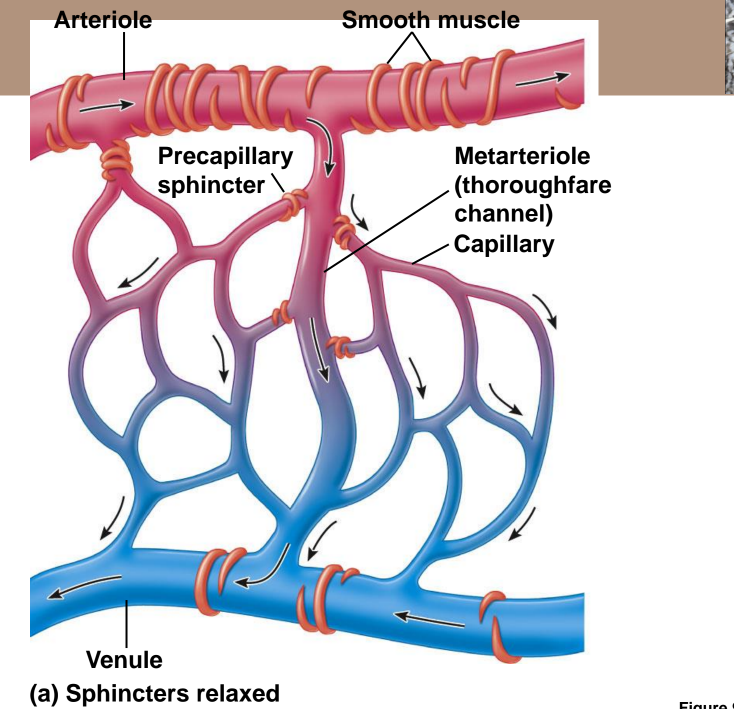
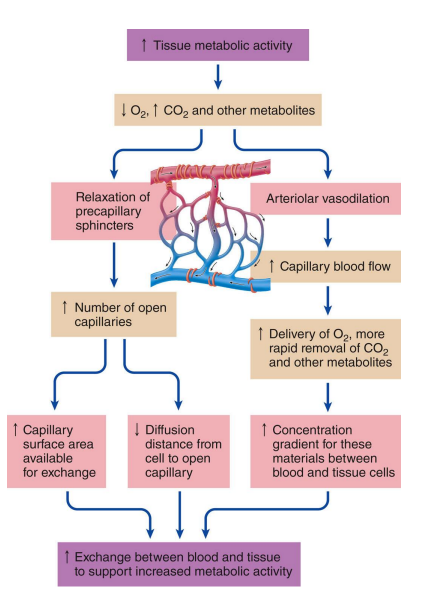
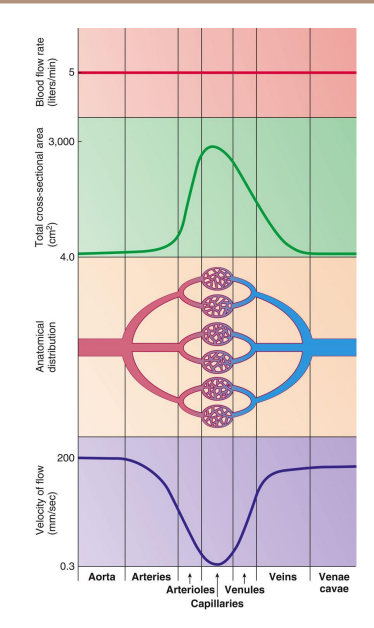


Figure 9-52 p440



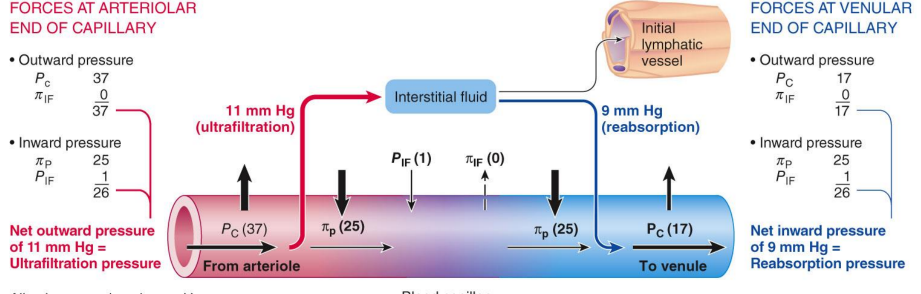






- Bulk flow across the capillary wall
 - Bulk flow occurs because of differences in the hydrostatic and colloid osmotic pressures between plasma and interstitial fluid
 - Capillary blood pressure (P_c) is the hydrostatic pressure exerted on the capillary walls
 - Plasma colloid osmotic pressure (π_p) is caused by plasma proteins
 - Interstitial fluid hydrostatic pressure (P_{IF}) is low
 - Interstitial fluid colloid osmotic pressure (π_{IF}) is very low
 - Protein-free plasma flows out of the capillary (ultrafiltration), mixes with interstitial fluid and then reenters the capillary (reabsorption)

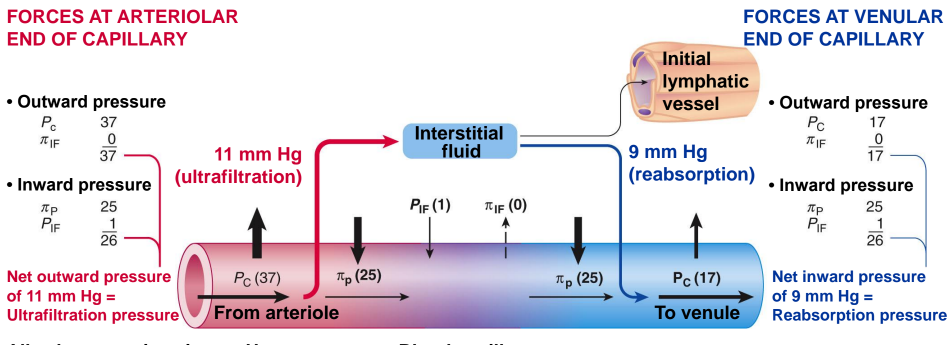




All values are given in mm Hg.

Blood capillary





All values are given in mm Hg.

Blood capillary

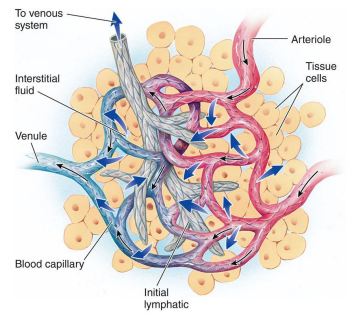
Figure 9-57 p443

9.14 Circulatory Vessels: Lymphatic System

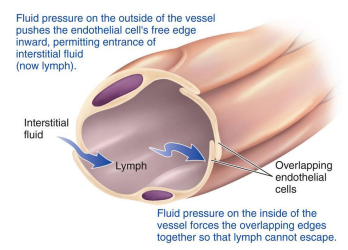


- The lymphatic system returns excess filtered fluid to the blood.
 - Network of one-way vessels
 - Interstitial fluid enters small blind-ended terminal vessels (initial lymphatics) to form lymph
 - Lymph vessels flow into the venous system near the heart
 - Passage of lymph through the lymph nodes aids in immune defense against disease
 - Lymph vessels **absorb fat** form the digestive tract

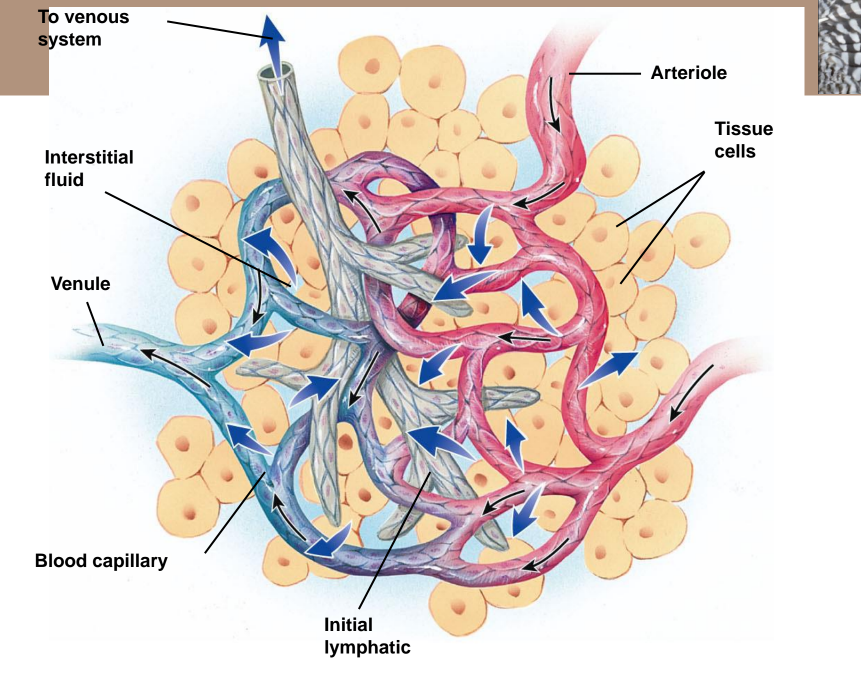
9.14 Circulatory Vessels: Lymphatic System



(a) Relationship between initial lymphatics and blood capillaries



(b) Arrangement of endothelial cells in an initial lymphatic



(a) Relationship between initial lymphatics and blood capillaries

Fluid pressure on the outside of the vessel pushes the endothelial cell's free edge inward, permitting entrance of interstitial fluid (now lymph).



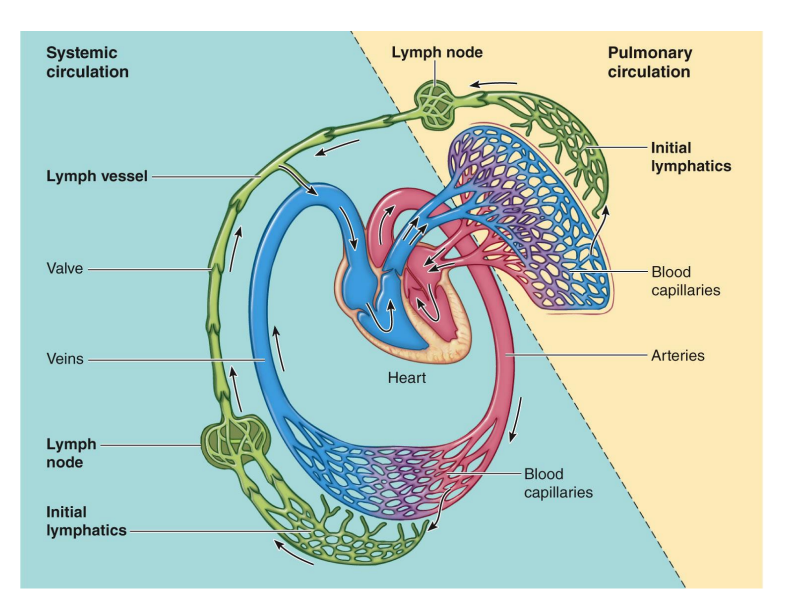
Interstitial fluid Lymph Overlapping endothelial cells

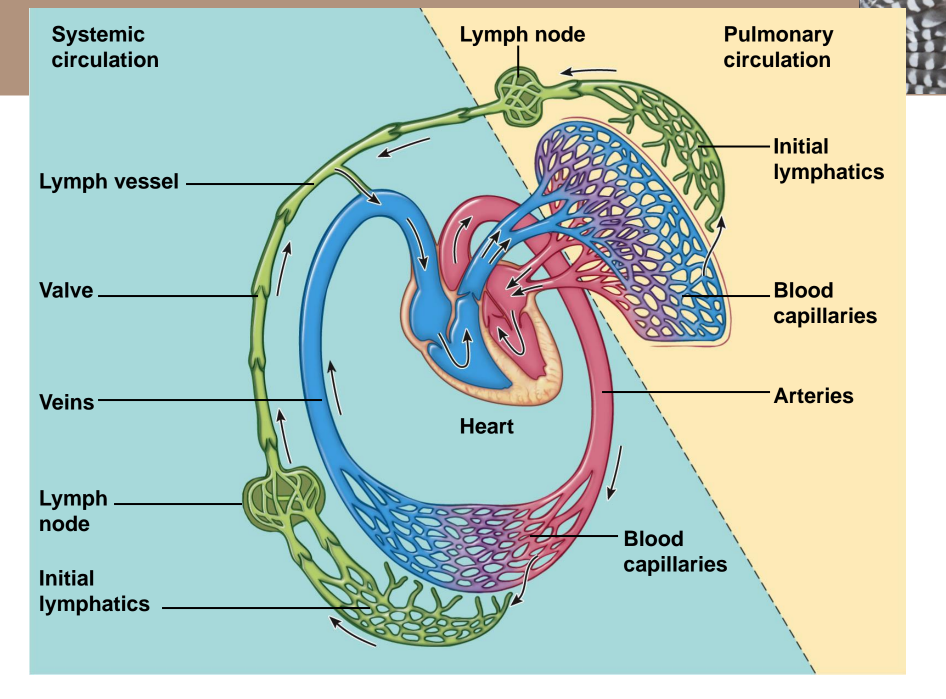
Fluid pressure on the inside of the vessel forces the overlapping edges together so that lymph cannot escape.

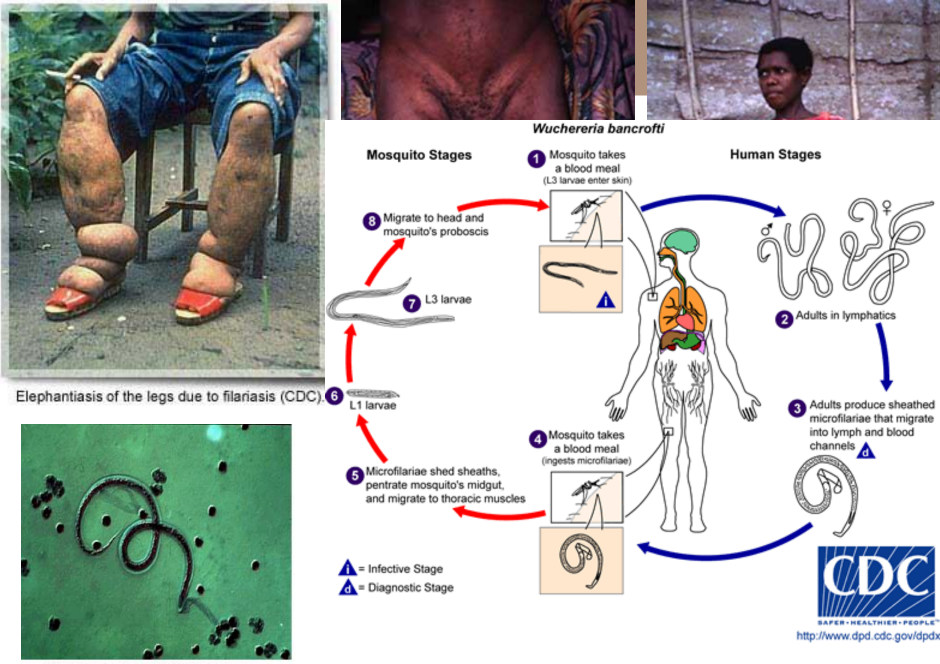
(b) Arrangement of endothelial cells in an initial lymphatic

Figure 9-58b p445

9.14 Circulatory Vessels: Lymphatic System



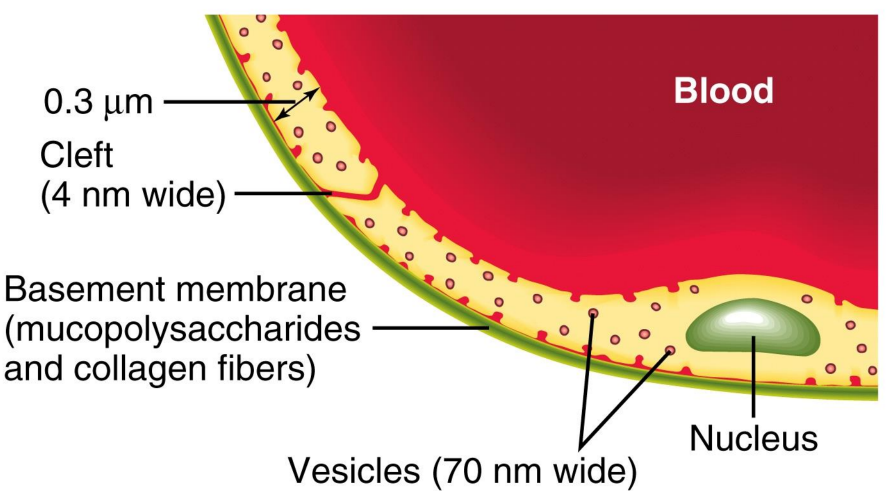




Wucheria bancrofti in blood WHO/TDR/Stammers

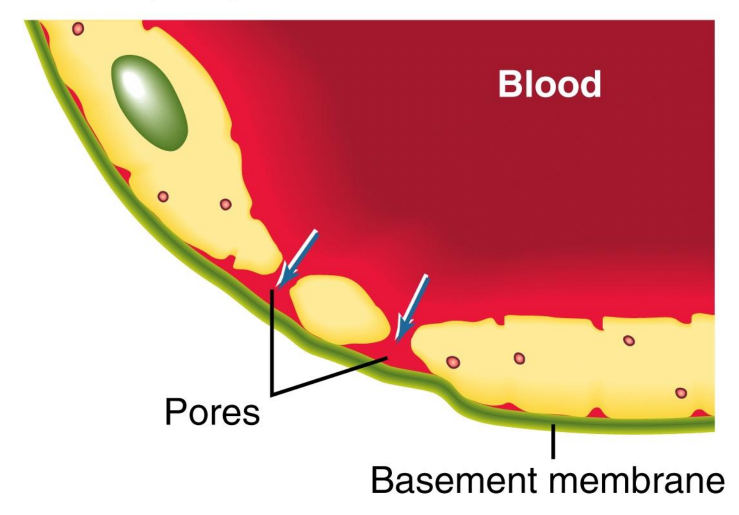


(a) Continuous capillary



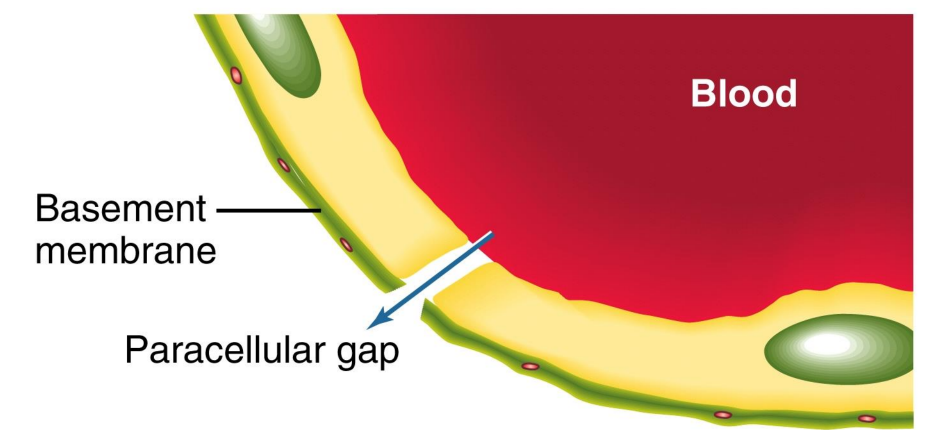


(b) Fenestrated capillary





(c) Sinusoidal capillary





- Veins serve as a blood reservoir and return blood to the heart.
 - Veins are **large** in diameter with little elasticity and low myogenic tone.
 - Easily **distend** to accommodate additional volumes of blood with no recoil (**capacitance vessels**)
 - Mammalian veins contain more than 60% of the total blood volume
 - Blood velocity accelerates on return from capillaries



TABLE 9-2 Features of Blood Vessels (with Human Values)

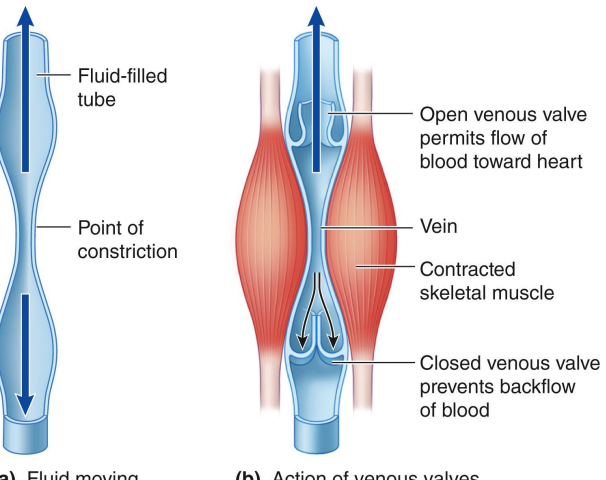
Feature	Arteries	Arterioles	Capillaries	Veins
Number	Several hundred*	Half a million	Ten billion	Several hundred*
Special Features Functions	Thick, highly elastic, walls; large radii* Passageway from heart to organs; serve as pres- sure reservoir	Highly muscular, well- innervated walls; small radii Primary resistance vessels; determine distribution of cardiac output	Very thin walled; large to- tal cross-sectional area Site of exchange; deter- mine distribution of extra- cellular fluid between plasma and interstitial fluid	Thin walled compared to arteries; highly distensible; large radii* Passageway to heart from or- gans; serve as blood reservoir

VESSEL TYPE

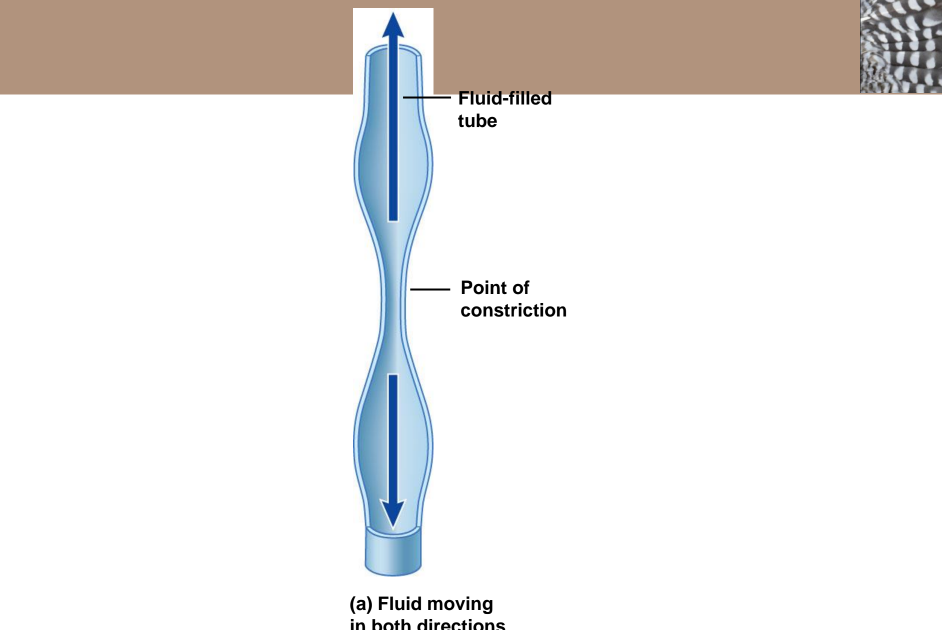


- Factors enhancing venous return to the heart
 - Driving pressure of cardiac contraction
 - Sympathetic activity produces venous vasoconstriction
 - Repeated contraction of skeletal muscles compresses veins (skeletal muscle pump)
 - Venous valves prevent blood from flowing backward (away from the heart)
 - Respiratory activity reduces pressure near the heart (respiratory pump)
 - Cardiac suction due to low ventricular pressures during diastole

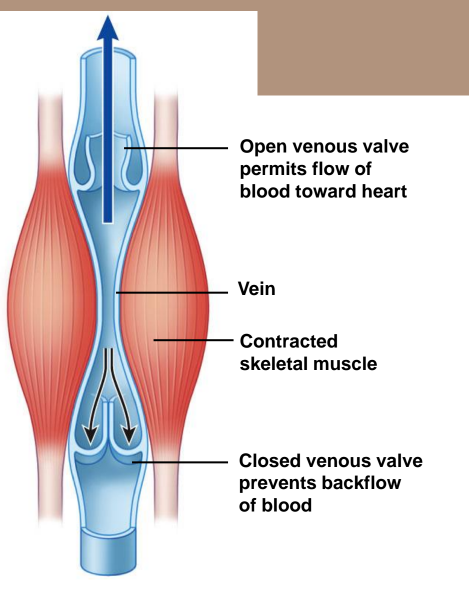




(a) Fluid moving in both directions on squeezing a fluid-filled tube (b) Action of venous valves, permitting flow of blood toward heart and preventing backflow of blood

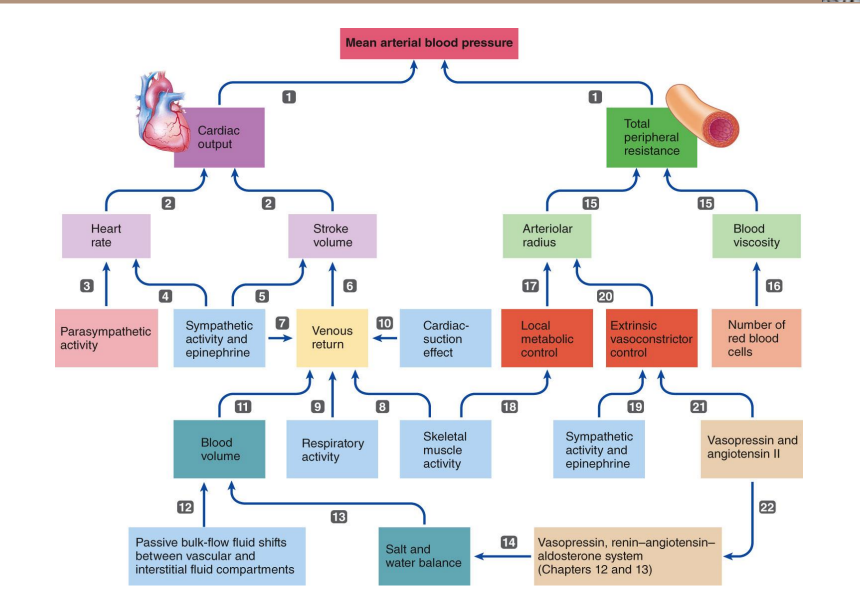


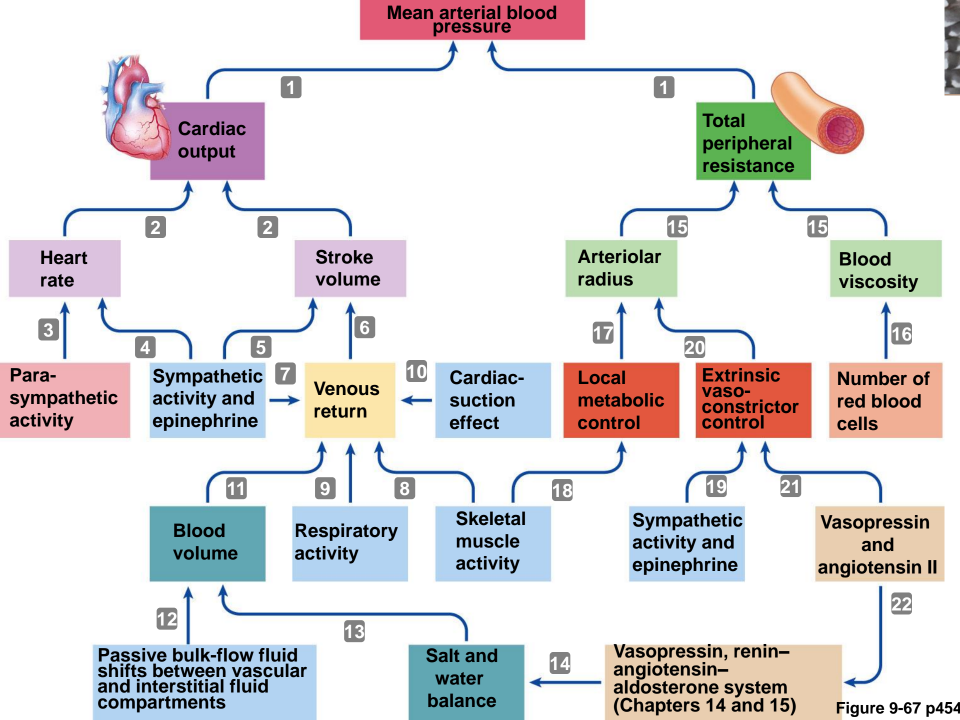
in both directions on squeezing a fluid-filled tube



(b) Action of venous valves, permitting flow of blood toward heart and preventing backflow of blood

9.15 Circulatory Vessels: Venules and Veins







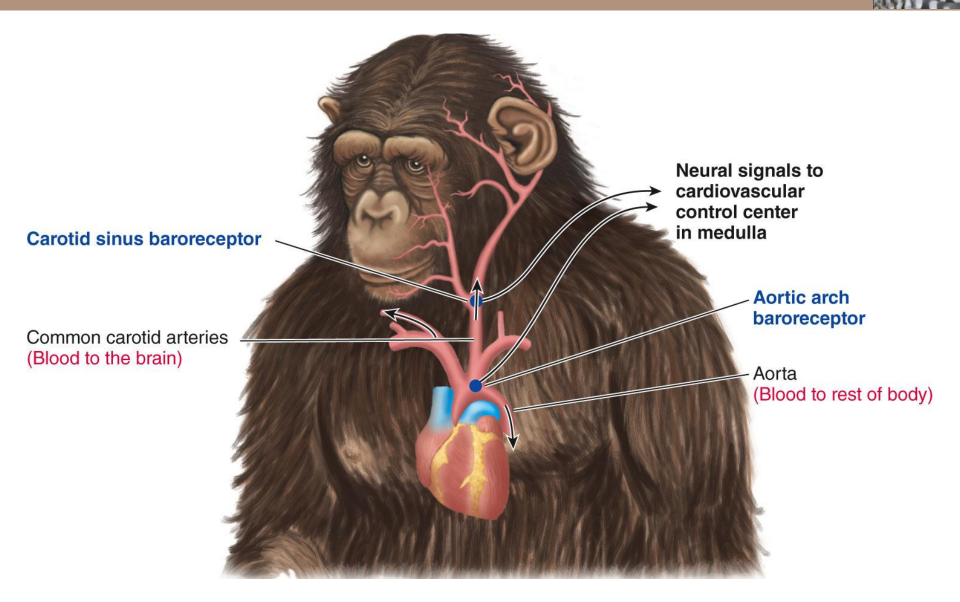
- Cardiovascular function is regulated in an integrated fashion
 - Two major goals:
 - Proper gas and heat transport
 - Maintaining arterial blood pressure
 - Cardiac output is homeostatically regulated when an animal is at rest to maintain consistent delivery of oxygen to key organs.
 - During activity, cardiac output is reset higher to boost blood flow to skeletal muscles and skin.



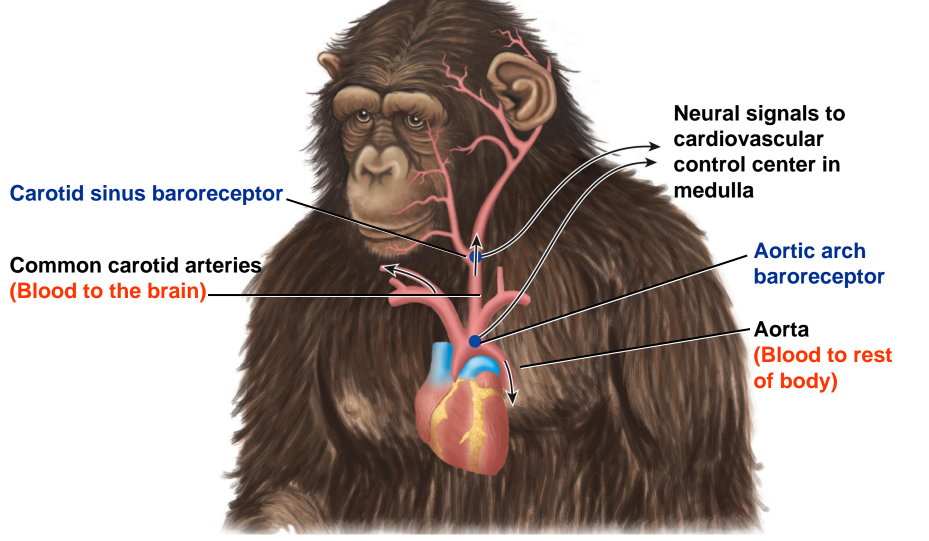
- Regulation of arterial blood pressure
 - Pressure must be high enough to overcome gravity, friction and other resistance factors
 - Pressure must be high enough for ultrafiltration in the kidneys
 - Pressure must not be so high that it creates extra work for the heart and increases the risk of vascular damage.
 - Chronic high blood pressure (hypertension) is due to excess resistance

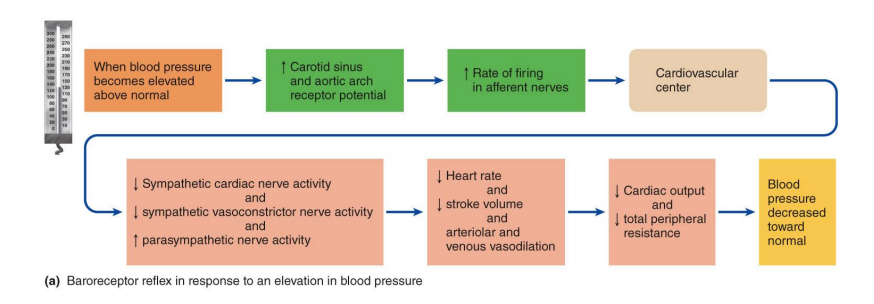


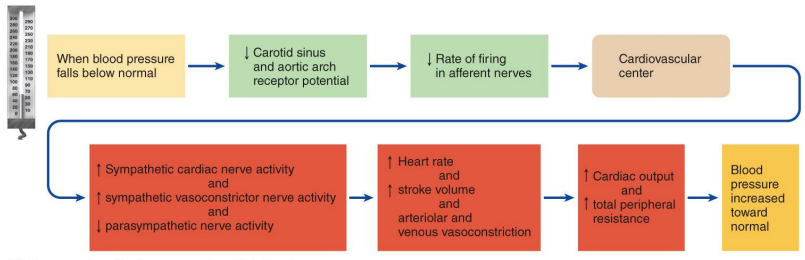
- Baroreflex regulation of arterial blood pressure
 - Arterial baroreceptors monitor blood pressure
 - The medullary cardiovascular control center alters sympathetic and parasympathetic activity
 - The heart and blood vessels respond to changing autonomic output
 - Example: Cardiovascular control center restores falling arterial blood pressure after standing by increasing sympathetic output, thus increasing heart rate and stroke volume and constricting blood vessels





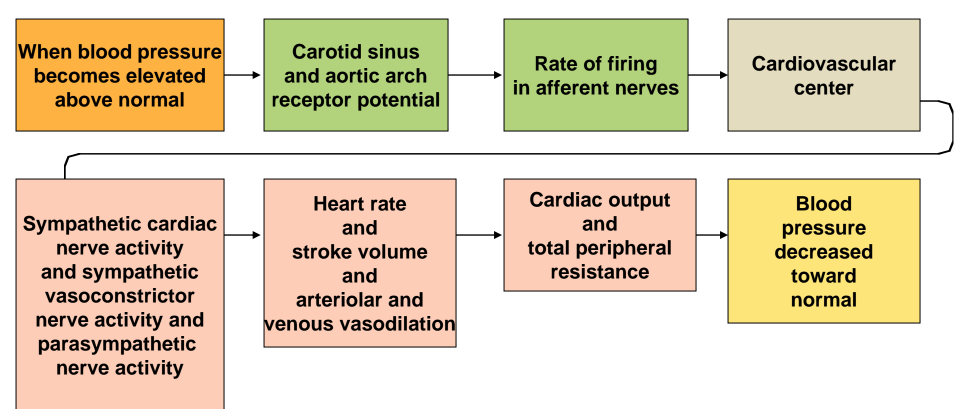






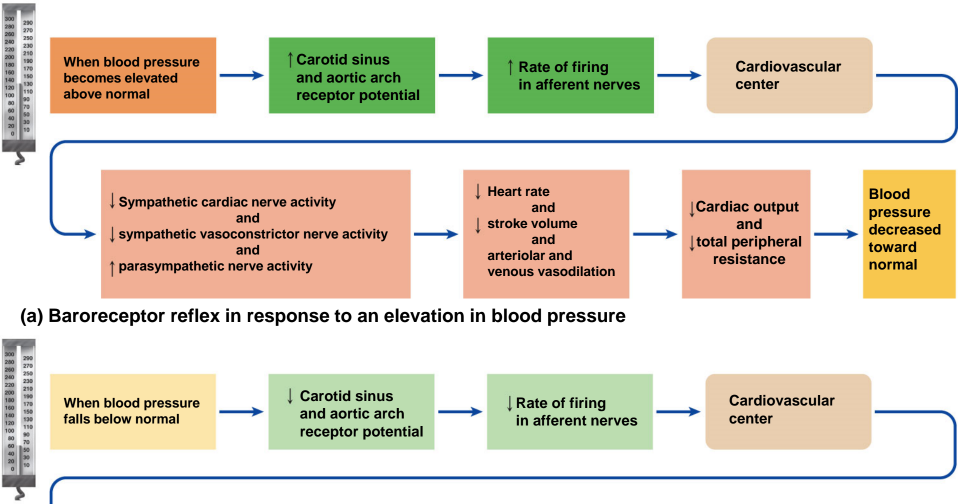
(b) Baroreceptor reflex in response to a fall in blood pressure

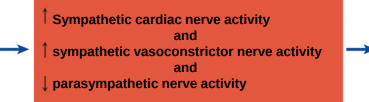


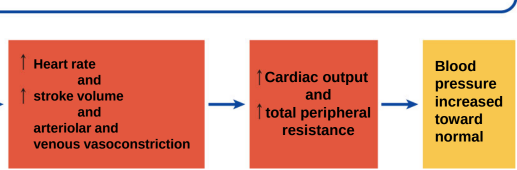


Stepped Art

Fig. 9-65a, p.451



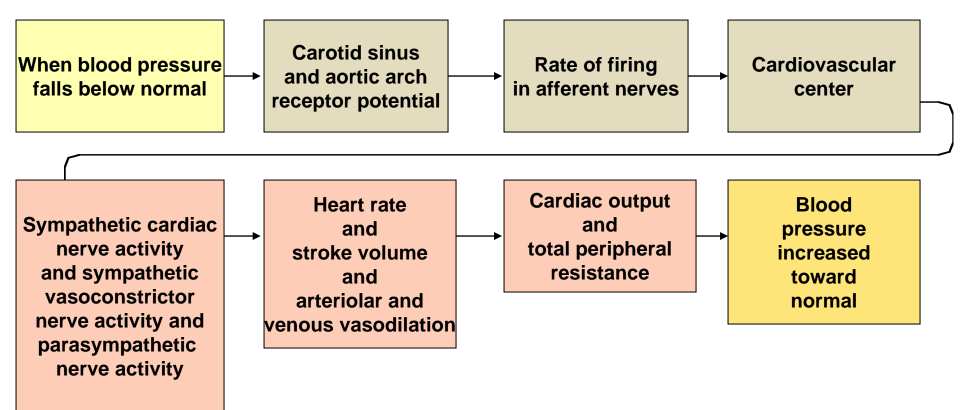




(b) Baroreceptor reflex in response to a fall in blood pressure

Figure 9-65 p451





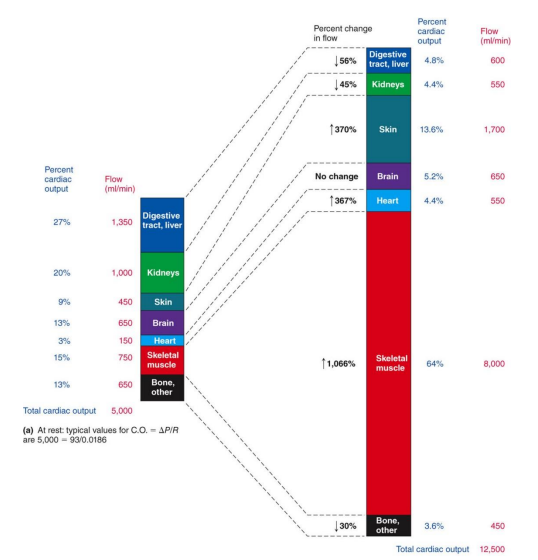
Stepped Art

Fig. 9-65b, p.451



- Response to increased locomotor activity
 - Increased heart rate and stroke volume result in increased cardiac output
 - Blood flow to skeletal muscle, heart and skin increases
 - Anticipatory responses
 - Increased locomotor activity induces cardiac and vascular changes before that activity leads to disturbances in blood gases.
 - Perception of stress can trigger the fight-or-flight response





(b) Activity: example values for C.O. = $\Delta P/R$ are 12,500 = 110/0.0088

