# **Cardiovascular Physiology**

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- Functional components of the cardiovascular system:
  - Heart
  - Blood Vessels
  - Blood
- General functions these provide
  - Transportation
    - Everything transported by the blood
  - Regulation
    - Of the cardiovascular system
      - Intrinsic v extrinsic
  - Protection
    - Against blood loss
  - Production/Synthesis

## **Cardiovascular System Function**

- To create the "pump" we have to examine the Functional Anatomy
  - Cardiac muscle
  - Chambers
  - Valves
  - Intrinsic Conduction System

# **Cardiac Muscle**

- Characteristics
  - Striated
  - Short branched cells
  - Uninucleate
  - Intercalated discs
  - T-tubules larger and over z-discs

(b)



## **Chambers of the Heart**

- 4 chambers
  - 2 Atria
  - -2 Ventricles
- 2 systems
  - Pulmonary
  - Systemic



## Valves of the Heart

- Function is to prevent backflow
  - Atrioventricular Valves
    - Prevent backflow to the atria
    - Prolapse is prevented by the chordae tendinae
      - Tensioned by the papillary muscles
  - Semilunar Valves
    - Prevent backflow into ventricles





## **Intrinsic Conduction System**

- Consists of "pacemaker" cells and conduction pathways
  - Coordinate the contraction of the atria and ventricles



## Autorhythmic Cells (Pacemaker Cells)

- Characteristics of
  Pacemaker Cells
  - Smaller than contractile cells
  - Don't contain many myofibrils
  - No organized sarcomere structure
    - do not contribute to the contractile force of the heart



SA node cell

AV node cells

## Autorhythmic Cells (Pacemaker Cells)

- Characteristics of Pacemaker Cells
  - Unstable membrane potential
    - "bottoms out" at -60mV
    - "drifts upward" to -40mV, forming a pacemaker potential
  - Myogenic
    - The upward "drift" allows the membrane to reach threshold potential (-40mV) by itself
    - This is due to
      - 1. Slow leakage of K<sup>+</sup> out & faster leakage Na<sup>+</sup> in
        - » Causes slow depolarization
        - Occurs through I<sub>f</sub> channels (f=funny) that open at negative membrane potentials and start closing as membrane approaches threshold potential
      - 2. Ca<sup>2+</sup> channels opening as membrane approaches threshold
        - » At threshold additional Ca<sup>2+</sup> ion channels open causing more rapid depolarization
        - » These deactivate shortly after and
      - 3. Slow K<sup>+</sup> channels open as membrane depolarizes causing an efflux of K<sup>+</sup> and a repolarization of membrane

### **Characteristics of Pacemaker Cells**



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### **Altering Activity of Pacemaker Cells**

- Sympathetic activity
  - NE and E increase I<sub>f</sub> channel activity
    - Binds to  $\beta_1$  adrenergic receptors which activate cAMP and increase I<sub>f</sub> channel open time
    - Causes more rapid pacemaker potential and faster rate of action potentials

#### Sympathetic Activity Summary:

increased dromotropic effects ↑conduction of APs

increased inotropic effects ↑contractility

![](_page_10_Figure_9.jpeg)

## **Altering Activity of Pacemaker Cells**

- Parasympathetic activity
  - ACh binds to muscarinic receptors
    - Increases K<sup>+</sup> permeability and decreases Ca<sup>2+</sup> permeability
      - = hyperpolarizing the membrane
        - » Longer time to threshold = slower rate of action potentials

![](_page_11_Figure_6.jpeg)

![](_page_11_Figure_7.jpeg)

## **Contractile Cells**

- Special aspects
  - Intercalated discs
    - Highly convoluted and interdigitated junctions
      - Joint adjacent cells with
        - » Desmosomes & fascia adherens
      - Allow for synticial activity
        - » With gap junctions
  - More mitochondria than skeletal muscle
  - Less sarcoplasmic reticulum
    - Ca<sup>2+</sup> also influxes from ECF reducing storage need
  - Larger t-tubules
    - Internally branching
  - Myocardial contractions are graded!

- Special aspects
  - The action potential of a contractile cell
    - Ca<sup>2+</sup> plays a major role again
    - Action potential is longer in duration than a "normal" action potential due to Ca<sup>2+</sup> entry
    - Phases
      - 4 resting membrane potential @ -90mV
      - 0 depolarization
        - » Due to gap junctions or conduction fiber action
        - » Voltage gated Na<sup>+</sup> channels open... close at 20mV
      - 1 temporary repolarization
        - » Open K<sup>+</sup> channels allow some K<sup>+</sup> to leave the cell
      - 2 plateau phase
        - » Voltage gated Ca<sup>2+</sup> channels are fully open (started during initial depolarization)
      - 3 repolarization
        - » Ca2+ channels close and K+ permeability increases as slower activated K+ channels open, causing a quick repolarization
  - What is the significance of the plateau phase?

### **Skeletal Action Potential vs Contractile Myocardial Action Potential**

![](_page_14_Figure_1.jpeg)

(a) Skeletal muscle fast-twitch fiber: The refractory period (yellow) is very short compared with the amount of time required for the development of tension.

![](_page_14_Figure_3.jpeg)

- Plateau phase prevents summation due to the elongated refractory period
- No summation capacity = no tetanus

- Which would be fatal

![](_page_15_Figure_3.jpeg)

![](_page_15_Figure_4.jpeg)

#### TABLE 14-3

#### Comparison of Action Potentials in Cardiac and Skeletal Muscle

	SKELETAL MUSCLE	CONTRACTILE MYOCARDIUM	AUTORHYTHMIC MYOCARDIUM
Membrane potential	Stable at -70 mV	Stable at -90 mV	Unstable pacemaker potential; usually starts at –60 mV
Events leading to threshold potential	Net Na <sup>+</sup> entry through ACh- operated channels	Depolarization enters via gap junctions	Net Na <sup>+</sup> entry through I <sub>f</sub> chan- nels; reinforced by Ca <sup>2+</sup> entry
Rising phase of action potential	Na <sup>+</sup> entry	Na <sup>+</sup> entry	Ca <sup>2+</sup> entry
Repolarization phase	Rapid; caused by K <sup>+</sup> efflux	Extended plateau caused by Ca <sup>2+</sup> entry; rapid phase caused by K <sup>+</sup> efflux	Rapid; caused by K <sup>+</sup> efflux
Hyperpolarization	Due to excessive K <sup>+</sup> efflux at high K <sup>+</sup> permeability when K <sup>+</sup> channels close; leak of K <sup>+</sup> and Na <sup>+</sup> restores potential to resting state	None; resting potential is –90 mV, the equilibrium poten- tial for K <sup>+</sup>	Normally none; when repolariza- tion hits –60 mV, the I <sub>f</sub> channels open again. ACh can hyperpolar- ize the cell.
Duration of action potential	Short: 1–2 msec	Extended: 200+ msec	Variable; generally 150+ msec
Refractory period	Generally brief	Long because resetting of Na <sup>+</sup> channel gates delayed until end of action potential	None

- Initiation
  - Action potential via pacemaker cells to conduction fibers
- Excitation-Contraction Coupling
  - Starts with CICR (Ca<sup>2+</sup> induced Ca<sup>2+</sup> release)
    - AP spreads along sarcolemma
    - T-tubules contain voltage gated L-type Ca<sup>2-</sup> channels which open upon depolarization
    - Ca<sup>2+</sup> entrance into myocardial cell and opens RyR (ryanodine receptors) Ca<sup>2+</sup> release channels
    - Release of Ca<sup>2+</sup> from SR causes a Ca<sup>2+</sup> "spark"
    - Multiple sparks form a Ca<sup>2+</sup> signal

![](_page_17_Figure_9.jpeg)

![](_page_17_Picture_10.jpeg)

- Excitation-Contraction Coupling cont...
  - 2. Ca<sup>2+</sup> signal (Ca<sup>2+</sup> from SR and ECF) binds to troponin to initiate myosin head attachment to actin
- Contraction
  - Same as skeletal muscle, but...
  - Strength of contraction varies
    - Sarcomeres are not "all or none" as it is in skeletal muscle
      - The response is graded!
        - » Low levels of cytosolic Ca<sup>2+</sup> will not activate as many myosin/actin interactions and the opposite is true
    - Length tension relationships exist
      - Strongest contraction generated when stretched between 80 & 100% of maximum (physiological range)
      - What causes stretching?
        - » The filling of chambers with blood

![](_page_18_Figure_12.jpeg)

### Myocardial Physiology Contractile Cells

- Relaxation
  - Ca<sup>2+</sup> is transported back into the SR and
  - Ca<sup>2+</sup> is transported out of the cell by a facilitated Na<sup>+</sup>/Ca<sup>2+</sup> exchanger (NCX)
  - As ICF Ca<sup>2+</sup> levels drop, interactions between myosin/actin are stopped
  - Sarcomere lengthens

![](_page_19_Figure_6.jpeg)

## **Cardiac Cycle**

- Cardiac cycle is the sequence of events as blood enters the atria, leaves the ventricles and then starts over
- Synchronizing this is the Intrinsic Electrical Conduction System
- Influencing the rate (chronotropy & dromotropy) is done by the sympathetic and parasympathetic divisions of the ANS

- Electrical Conduction Pathway
  - Initiated by the Sino-Atrial node (SA node) which is myogenic at 70-80 action potentials/minute
  - Depolarization is spread through the atria via gap junctions and internodal pathways to the Atrio-Ventricular node (AV node)
    - The fibrous connective tissue matrix of the heart prevents further spread of APs to the ventricles
    - A slight delay at the AV node occurs
      - Due to slower formation of action potentials
      - Allows further emptying of the atria
  - Action potentials travel down the Atrioventricular bundle (Bundle of His) which splits into left and right atrioventricular bundles (bundle branches) and then into the conduction myofibers (Purkinje cells)
    - Purkinje cells are larger in diameter & conduct impulse very rapidly
      - Causes the cells at the apex to contract nearly simultaneously
        - » Good for ventricular ejection

![](_page_22_Figure_0.jpeg)

- The electrical system gives rise to electrical changes (depolarization/repolarization) that is transmitted through isotonic body fluids and is recordable
  - The ECG!
    - A recording of electrical activity
    - Can be mapped to the cardiac cycle

![](_page_24_Picture_0.jpeg)

![](_page_24_Picture_1.jpeg)

![](_page_24_Figure_2.jpeg)

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## Cardiac Cycle Phases

- Systole = period of contraction
- Diastole = period of relaxation
- Cardiac Cycle is alternating periods of systole and diastole
- Phases of the cardiac cycle
  - 1. Rest
    - Both atria and ventricles in diastole
    - Blood is filling both atria and ventricles due to low pressure conditions
  - 2. Atrial Systole
    - Completes ventricular filling
  - 3. Isovolumetric Ventricular Contraction
    - Increased pressure in the ventricles causes the AV valves to close...
      - Creates the first heart sound (lub)
    - Atria go back to diastole
    - No blood flow as semilunar valves are closed as well

- Phases of the cardiac cycle
  - 4. Ventricular Ejection
    - Intraventricular pressure overcomes aortic pressure
      - Semilunar valves open
      - Blood is ejected
  - 5. Isovolumetric Ventricular Relaxation
    - Intraventricular pressure drops below aortic pressure
      - Semilunar valves close = second heart sound (dup)
    - Pressure still hasn't dropped enough to open AV valves so volume remains same (isovolumetric)

Back to Atrial & Ventricular Diastole

![](_page_27_Figure_0.jpeg)

### Cardiac Cycle Blood Volumes & Pressure

![](_page_28_Figure_1.jpeg)

![](_page_29_Figure_0.jpeg)