CVS MODULE PHYSIOLOGY (LECTURE 1) Physiology of Cardiac Muscle I

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The **cardiovascular system** is a **closed system** of tubes (i.e. vessels) inside which blood circulates continuously by the **pumping action** of the **heart** in **one direction** only by the action of **valves** present in the **heart** and **veins**.

It is composed of :

1. <u>The Heart:</u> as a **pumping organ** of the system.

<u>Blood Vessels</u>: as containers, through which the circulation occurs.

- Cardiac muscle is a striated muscle like the skeletal muscle, but it is different from the skeletal muscle in being involuntary and syncytial.
- <u>Syncytium</u>: It means that cardiac muscle cells are able to excite and contract together as one unit due to the presence of gap junctions between adjacent cardiac cells.
- Both atria contract together as one unit (upper syncytium) and both ventricles contract together as one unit (i.e. lower syncytium), which are completely separated from each other by the fibrous A-V ring. So, the excitation waves cannot be directly transmitted from one syncytium to the other.

Myocardium of the heart is composed of two types of cardiac muscle cells (fibers):

A. Contractile Cells :

- Form about 98-99% of the cardiac muscle).
- Their action potential (AP) is called fast AP.

B. Non-contractile (auto-rhythmic) Cells:

- Form about 1-2 % of the cardiac muscles and are the cells that form excitatory- conductive system of the heart).
- Their AP is called slow or pacemaker AP.

Functions of the atria and ventricles

Atria:

- They are the entry-ways to the ventricles: they receive and store the venous return then pass it to the ventricles during ventricular diastole.
- The atrial walls contain stretch receptors the monitor changes in the intra-atrial pressure.
- Secret the atrial natriuretic peptide (ANP) which favors Na⁺ and water excretion by the kidneys and also causes VD.

Ventricles:

- Chambers whose contractions produce the pressures that drive blood through the pulmonary and systemic vascular systems and back to the heart.
- The pumping action of ventricles is the main force that propels blood to the peripheral circulatory system, thus loss of such function is fatal.

Physiological Properties of the Cardiac Muscle

Cardiac muscle has four properties, due to which the heart is able to fulfill its function as a pumping organ.

They include:

- 1. Automaticity & Rhythmicity (Chronotropism).
- 2. Excitability (Bathmotropism).
- 3. Conductivity (Dromotropism).
- 4. Contractility (Inotropism).

1. Automaticity and Rhythmicity (Auto-rhythmicity)

Automaticity:

it is the property of self-excitation; the ability of spontaneous generation of action potentials independent of any extrinsic stimuli.

Rhythmicity:

The regular generation of these action potentials (the heart can beat regularly).

Spontaneous automaticity and rhythmicity (auto-rhythmicity) of the cardiac muscle is due to the existence of a specialized excitatory-conductive system, which is composed of modified selfexciting, non-contractile cardiac muscle cells called pacemaker cells. Auto-rhythmicity is myogenic in origin (i.e. starts from the muscle itself independent from autonomic nerve supply that only controls the heart rate (either \uparrow or \downarrow) but don't initiate the beat.

Pacemaker(s) of the heart:

Pacemaker means the part of the heart that has the highest rhythmicity and the whole other parts of the heart follow its rhythm. <u>They include:</u>

1. <u>Sinoatrial (auricular) (SA) Node (1ry; normal Pacemaker):</u>

It has the highest rhythm (90-110; 100/minute).

So, it is called the **normal** or **1ry pacemaker** of the heart. Its rhythm is called **sinus rhythm**.

2. Atrioventricular (AV) Node (2ry Pacemaker):

Its rate is 45-60 /minute.

It acts only if SA node is damaged or blocked.

Its rhythm is called **nodal rhythm**.

3. Purkinje fibres (3ry Pacemaker):

Its rate is 25-40 /minute.

It takes over **only** if the conduction in **AV node is completely blocked**. Its rhythm is called **idioventricular rhythm**.





Mechanism of Autorhythmicity (Prepotential & Pacemaker AP)

- ✓ Pacemaker cells in the nodal tissue (SAN and AVN) have a resting membrane potential of about - 55 to -60 mV. However, it is not stable.
- ✓ After each impulse, gradual depolarization occurs spontaneously till a firing level is reached at which action potential (an impulse) is initiated.
- They can spontaneously and regularly initiate action potential and so they are responsible for automaticity and rhythmicity of the heart.
- ✓ This gradual depolarization is called prepotential or diastolic depolarization.

Pacemaker action potential is composed of: Phase 4; (Prepotential):

- ✓ Resting membrane potential is about -55 to -60 mV (unstable).
- ✓ At this potential there is activation of a special type of Na⁺ channels known as Na⁺ funny channels → funny current; F (inward Na⁺).
- ✓ Another type of channels called **transient** or <u>T-type</u> Ca⁺⁺ channels opens→ Ca⁺⁺ enters down its electrochemical gradient → depolarizes the cell to -40 mV.

Phase 0; (Depolarization):

- As the membrane is depolarized to the firing level (about 40 mV) another type of Ca⁺⁺ channels opens. These are known long-lasting or L-type Ca⁺⁺ channels → entrance of Ca⁺⁺.
- Because the movement of Ca⁺⁺ through these channels into the cell is not rapid, the rate of depolarization (slope of phase 0) is much slower than found in other types of cardiac cells. Therefore AP of pacemaker cells is called slow response action potential.

Phase 3; (Repolarization):

- It occurs due to opening of K⁺ channels→ outward directed K⁺ current along concentration and electric gradients. At the same time, L-type Ca⁺⁺ channels become inactivated and close → stop entrance of Ca⁺⁺.
- Repolarization continues until the membrane potential reaches -60 mV. At this potential the outward K⁺ current becomes gradually inactivated, while the inward Na⁺ current becomes activated again due to opening of funny current channels and a new phase 4 is initiated and the whole cycle is spontaneously repeated.

■ <u>N.B.:</u>

Phase 1 and 2 (which are present in fast AP) are absent.

Pacemaker Potential



Action potential of cardiac muscles

Grigoriy Ikonnikov and Eric Wong





Characters of pacemaker (slow) action potential

- The resting potential is 55 to 60 mV. It is unstable leading to prepotential (phase 4).
- Its upstroke (depolarization phase; phase 0) is slow, of small magnitude (up to +10 mV) and is mainly due to Ca⁺⁺ influx due to opening of L-type Ca⁺⁺ channels.
- The action potential duration is about 200- 250 ms.
- There is no plateau.
- The falling phase (repolarization; phase 3) is one phase.

<u>N.B.</u>

Although the rhythmicity of the SAN is ~ **100 /min**, the resting heart rate is **only about 75 beat/min**, **why**?

ANSWER:

This is due the **continuous inhibitory discharge** from the vagus nerve on SAN **decreasing** its inherited rhythm from **100 to 75 beat/min**. This called <u>VAGAL TONE</u>.

The vagus nerve **supplies** the whole cardiac muscle **except** the **ventricles** (i.e. called **VAGAL ESCAPE PHENOMENON**). This phenomenon **protects** the ventricles from **abnormally high** vagal stimulation (which can cause **cardiac arrest**).

Factors affecting rhythmicity (chronotropism)

- The effect of various factors on rhythmicity is called chronotropism.
- The factors that stimulate rhythmicity (accelerating the heart rate) are called + ve chronotropic factors, while factors that inhibit rhythmicity (slowing the heart rate) are called - ve chronotropic factors.

A. <u>Nervous Factors:</u>

1. <u>Parasympathetic (Vagal) stimulation via</u> acetylcholine (ACh) \rightarrow (-ve) chronotropic effect.

Mechanism:

Ach <u>stimulates M₂</u> cholinergic receptors \rightarrow decreased level of cyclic-AMP \rightarrow <u>inhibits F-type (Na⁺) channel</u> $\rightarrow \downarrow$ slope of prepotentials $\rightarrow \downarrow$ \downarrow rhythmicity. In addition, Parasympathetic stimulation also hyperpolarizes the plasma membranes of SAN cells by increasing their permeability to K⁺ \rightarrow K⁺ efflux. So, the rate of discharge is decreased (e.g. bradycardia). 2. Sympathetic stimulation via noradrenaline (NA) → (+ve) chronotropic effect.

Mechanism:

NA stimulates B_1 adrenergic receptors \rightarrow increased formation of cyclic-AMP \rightarrow increase inward Na⁺ current (F-type channel) & \uparrow Ca⁺² influx \rightarrow \uparrow slope of prepotentials \rightarrow \uparrow rhythmicity. So, the rate of discharge is increased (e.g. tachycardia).



B. Chemical Factors:

- **1. Catecholamines**: ↑ rhythmicity.
- **2. Thyroxine**: ↑ rhythmicity
- **3. Acetyl choline (ACh):** \downarrow rhythmicity.
- 4. Blood gases:
- Mild to moderate O_2 lack (hypoxia $\rightarrow \uparrow$ rhythmicity.
- Mild to moderate ↑ CO₂ (hypercapnia) or ↑ H+ (acidosis) → weakly inhibit SAN rhythmicity but they increase heart rate (HR).
- Severe O_2 lack ($\downarrow O_2$), \uparrow H+ or \uparrow $CO_2 \rightarrow \downarrow$ rhythmicity.
- 5. Effect of Drugs:
- Sympathomimetic drugs → ↑ rhythmicity.
- Parasympathomimetic drugs (i.e. cholinergic drugs) $\rightarrow \downarrow$ rhythmicity.
- Digitalis: Although it increases myocardial contractility, it inhibits SAN activity and decreases HR (i.e. vagal like effect)

C. Physical Factors:

A rise of body temperature (e.g. in muscular exercise or fever) $\rightarrow \uparrow$ rhythmicity (heart rate; HR) due to increased rate of discharge of SAN.

Hypothermia $\rightarrow \downarrow$ rhythmicity due to decreased rate of discharge of SAN.

D. Mechanical factors:

Rise of right atrial pressure个rhythmicity (Bainbridge reflex).

Right atrial distension may directly excite SAN leading to tachycardia (= Bainbridge effect).

