

رج تلاحظ انو في بعض الرسمات والمعلومات من مصدر
خارجي ولكن تم الاستعانتة فيهم للفهم
دعواتكم وبال توفيق



PHYSIOLOGY

FACULTY OF PHARMACEUTICAL SCIENCES

DR. AMJAAD ZUHIER ALROSAN

LECTURE 8, PART (2):ACTION POTENTIAL AND CONTRACTION
OF CARDIAC CONTRACTILE FIBERS

Objectives

1. Discuss **histology of cardiac muscle tissue**.
2. Discuss **action potential and contraction of contractile fibers**.
3. Describe **electrocardiogram as well as the cardiac cycle**.

(Pages 702-718, 720-726 of the reference).

THE CARDIOVASCULAR SYSTEM: THE HEART

Involuntary control

electrical synapse

skeletal \rightarrow chemical synapse

skeletal \rightarrow Ca^{+2} \rightarrow ATP \rightarrow Cardiac muscular

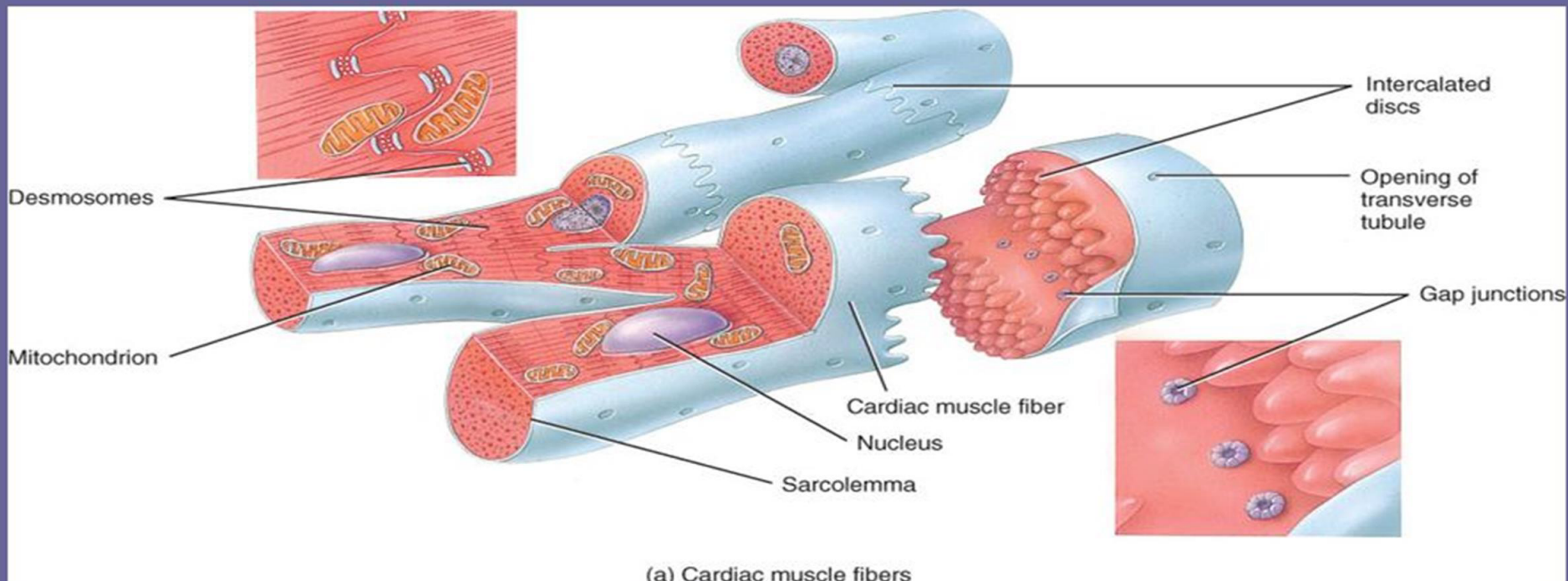
- The **heart contributes to homeostasis** by pumping blood through blood vessels to the tissues of the body to deliver oxygen and nutrients and remove wastes.
- The cardiovascular system consists of the blood, the heart, and blood vessels.

HISTOLOGY OF CARDIAC MUSCLE TISSUE

- Compared with skeletal muscle fibers, **cardiac muscle fibers are (shorter)** in length. They also **exhibit branching**, which gives individual cardiac muscle fibers a “stair-step” appearance.
→ Chemical signal
- Cardiac muscle fibers **connect** to neighboring fibers by **intercalated discs**, which contain **desmosomes**, which **hold the fibers together**, and **gap junctions**, which allow muscle action potentials to conduct from one muscle fiber to its neighbors.
- **Gap unit** junctions allow the entire myocardium of the **atria** or the **ventricles** to contract as a single, coordinated.
→ Unit 2

atrium + atrium

Cardiac Muscle Histology



- Branching, intercalated discs with gap junctions, involuntary, striated, single central nucleus per cell

AUTORHYTHMIC FIBERS:THE CONDUCTION SYSTEM

WLS action potential (easy ←)

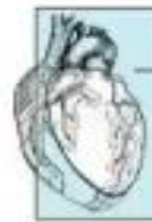
- ✓ An inherent and rhythmical electrical activity is the reason for the heart's lifelong beat.
- ✓ The **source of this electrical activity** is a network of specialized cardiac muscle fibers called **autorhythmic fibers** because they are **self-excitible**.
WLS lajwli 3,61
- ✓ Autorhythmic fibers **repeatedly generate action potentials that trigger heart contractions.**

Locations of autorhythmic cells

pacemaker

Sinoatrial node (SA node)

Specialized region in **right atrial** wall
near opening of **superior vena cava**
* العود الأسمون العلوي



Frontal plane

Atrioventricular node (AV node)

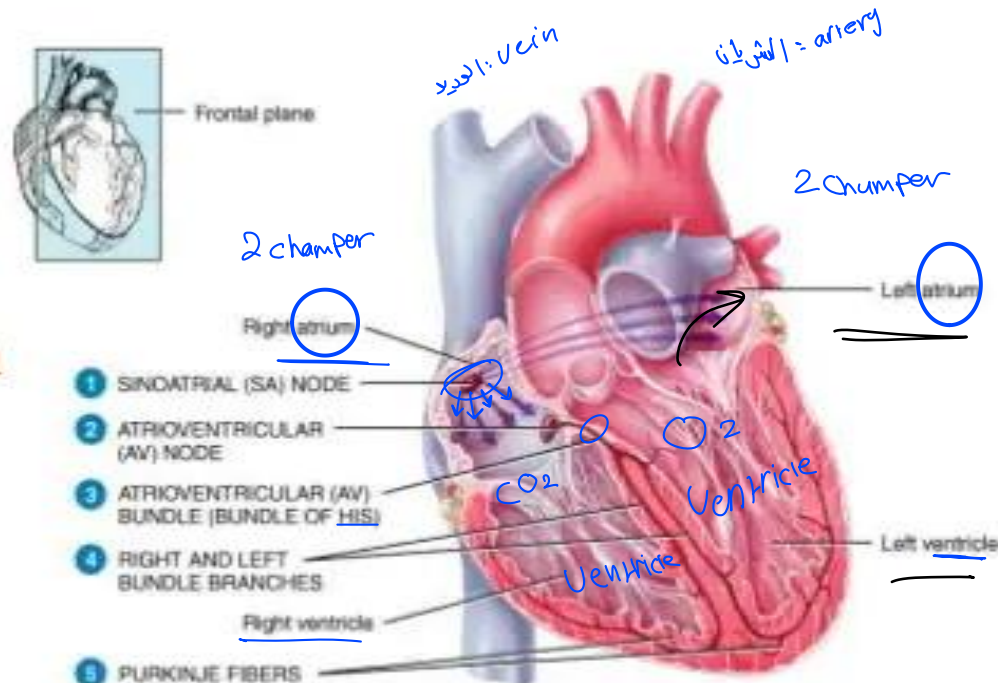
Small bundle of specialized cardiac cells located at **base** of right atrium near **septum**
ال حاجز

Bundle of His (atrioventricular bundle)

Cells originate at AV node and enters interventricular septum
Divides to form **right and left bundle branches** which travel down septum, curve around tip of **ventricular chambers**, travel back toward atria along outer walls

Purkinje fibers

Small terminal fibers that extend from **bundle of His** and spread throughout **ventricular myocardium**



الخلايا تنشأ من العقدة AV وتدخل

في الحاجز بين البطينين.

تنقسم إلى فرعين أيمن وأيسر يمتدان

في الحاجز ثم يلتفان حول قمة القلب

ويمتدان على جدران البطينين

(a) Anterior view of frontal section

20.10a



41 chapter

* ينحو الصلب من بطيئين وأذينين (عن فوق)

Filling ω_{max}

Right atrium

Right atrium

Right atrium

diastole

Vein ← ورجان - شريان

atrium: (n.) -

Ventricles: sub 11-2

artery وعاء دموي خارج من القلب \Rightarrow شريان :

Contraction \downarrow Right atrium \downarrow Right ventricle \downarrow Contraction is slow \times

وہ ایسی وسیعہ دھوئی دا احیل لئے آنکھاں تے وریڈ : Vein

پہلی جگہ سیارہ 1P: Unpaired 1st rib کا نام - Aorta

Pilling: dia stroke

Contraction:- is sole

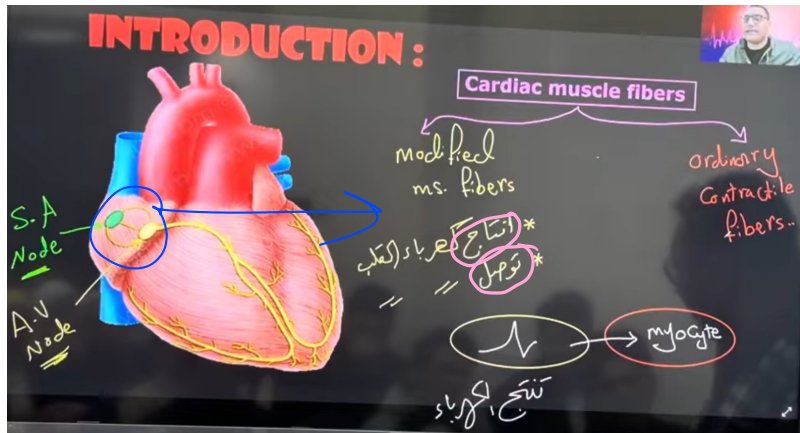
الحقوق المدنية والسياسية Right Human and Political Rights

palmonary artery \rightarrow CO₂

يوجد في جسم الإنسان دورةان دموييان هما: دورة كبيرة و دورة صغيرة (pulmonary circulation) و (systemic circulation) .

• contraction لاحظ انتقال العصبية من العصب المحيطي إلى العصب الشرياني (العصبيات المحيطية تصل إلى العصب الشرياني)

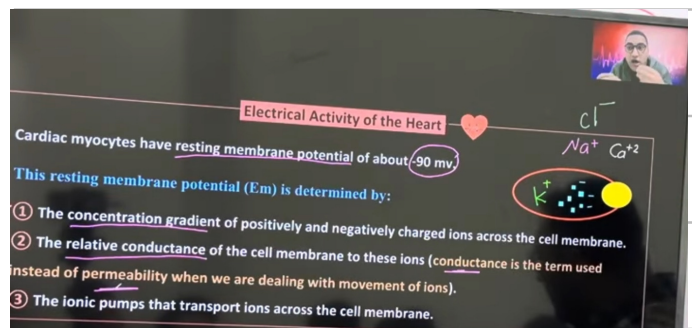
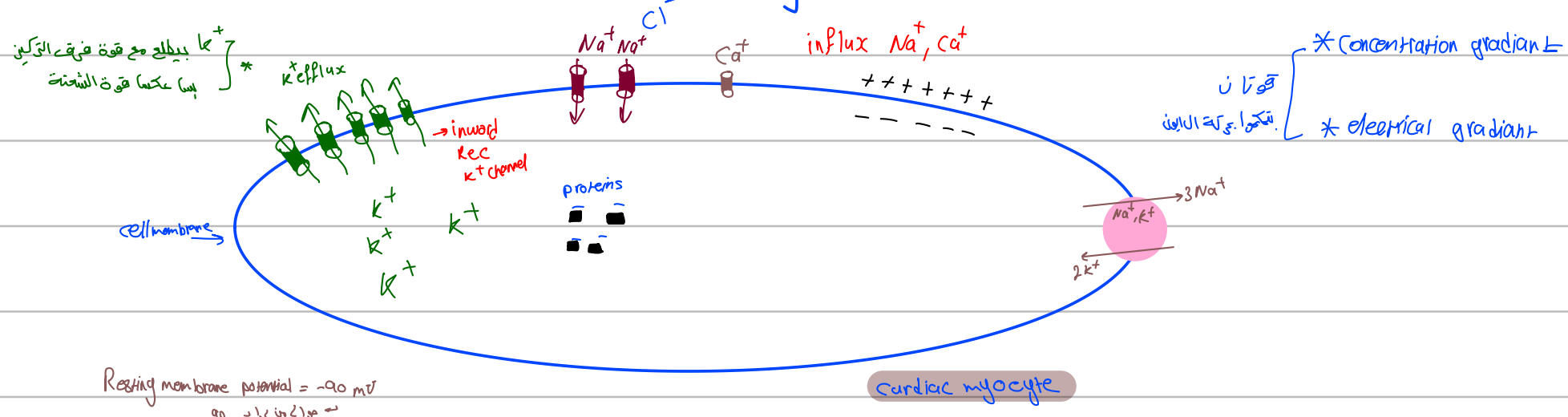
contraction لاحظ انتقال العصبية من العصب المحيطي إلى العصب الشرياني (العصبيات المحيطية تصل إلى العصب الشرياني) \therefore modified muscle fiber



SA node \longrightarrow AV node \longrightarrow bundle of His \longrightarrow Right & left bundle branches \longrightarrow Purkinje fibers

* مصدر خارجي دلائل ال نهاية الفيزيقيات الاتصال

electrical activity



$1 Na^+$ بدل $100 K^+$ يخرج *

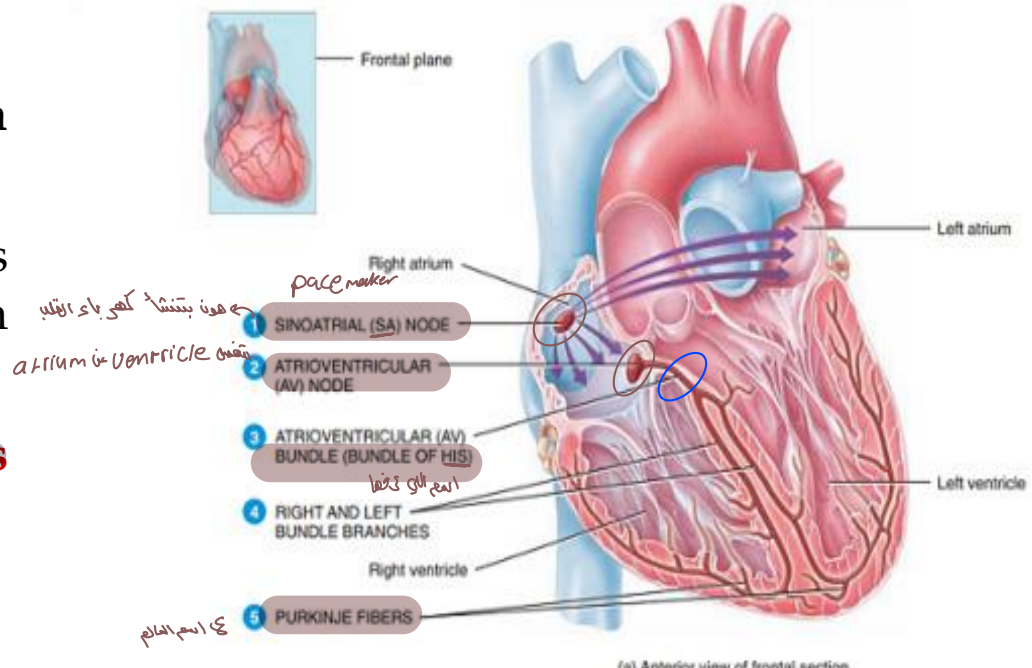
AUTORHYTHMIC FIBERS: THE CONDUCTION SYSTEM

1. They act as a pacemaker (electrical excitation that causes contraction of the heart).
2. They form the cardiac conduction system.
3. Cardiac action potentials propagate through the conduction system in the following sequence:
 - **Cardiac excitation normally begins in the (sinoatrial (SA) node)**

Pace maker

Figure 20.10 The conduction system of the heart. Autorhythmic fibers in the SA node (a), act as the heart's pacemaker, initiating cardiac action potentials (b) that cause contraction of the heart's chambers.

 The conduction system ensures that the chambers of the heart contract in a coordinated manner.



well this is action potential \downarrow so indep \rightarrow $*$

* Normal \rightarrow pacemaker \rightarrow SA node \rightarrow faster rate

ما بيعطيلووا كھو باد پس بديو جھلووا الکھر باد [AV node parkinje fibers]

- 1- spontaneous \Rightarrow Automaty
- 2- Regular \Rightarrow Rhythmicity

فقط Na^+ يدخلوا كم الين Na^+ funny

* لو اهذى خواسته من [SA node]

① Na^+ channels: Na^+ funny channels

دحول الـ N^+ بين O polarization اد دحول الـ O بـ N^+

Δ -type Ca^{2+} channel
↳ transient

voltage
potential

Piringlen

Depolarization

Living level

* Piring level

١- كل المفهومات التي قيل لها (40m) سُئل

transient Ca^{2+} channel - Ca^{2+} بفتح قنوات Ca^{2+} - 50 mV Ca^{2+} \rightarrow transient channel Ca^{2+} بفتح قنوات Ca^{2+}

لأنّ كُلّي عنصر لا يُجوا إلّا من SA node -

٤- دلائل التأثيرات المعاكسة (Depolarization)

AUTORHYTHMIC FIBERS: THE CONDUCTION SYSTEM

- SA node cells do **not** have a stable resting potential. Rather, they repeatedly depolarize to threshold spontaneously. The spontaneous depolarization is a pacemaker potential. *Well well well*
- When the pacemaker potential reaches threshold, it triggers an action potential. *Well*
Each action potential from the SA node propagates throughout both atria via gap junctions in the intercalated discs of atrial muscle fibers. Following the action potential, the two atria contract at the same time.

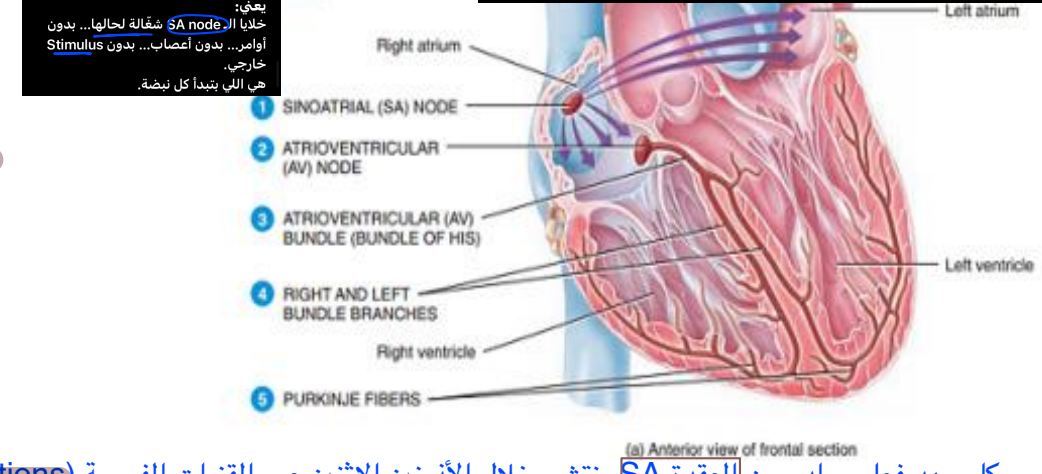
جود اطمئنهم للطلب

Figure 20.10 The conduction system of the heart.

⑥ The conduction system ensures that the



بمجرد ما يوصل للـ **Threshold** ينطلق **Action potential** في الخلية الجديدة.



رابعاً: النقاط المهمة للحفظ ◆

✓ خلايا SA node لا تملك resting potential ثابت.

✓ لديها **Pacemaker potential** يرتفع تلقائياً حتى

✓ عند الوصول للعتبة \rightarrow يخرج Action potential جديد.

✓ نتيجة ذلك: الأذينان ينقبضان في نفس الوقت.

كل جهد فعل صادر من العقدة SA يتنتشر خلال الأذنين عبر القنوات الفجوية (gap junctions) الموجودة في الأقراص البيضاء بين ألياف عضلة الأذنين.

AUTORHYTHMIC FIBERS: THE CONDUCTION SYSTEM

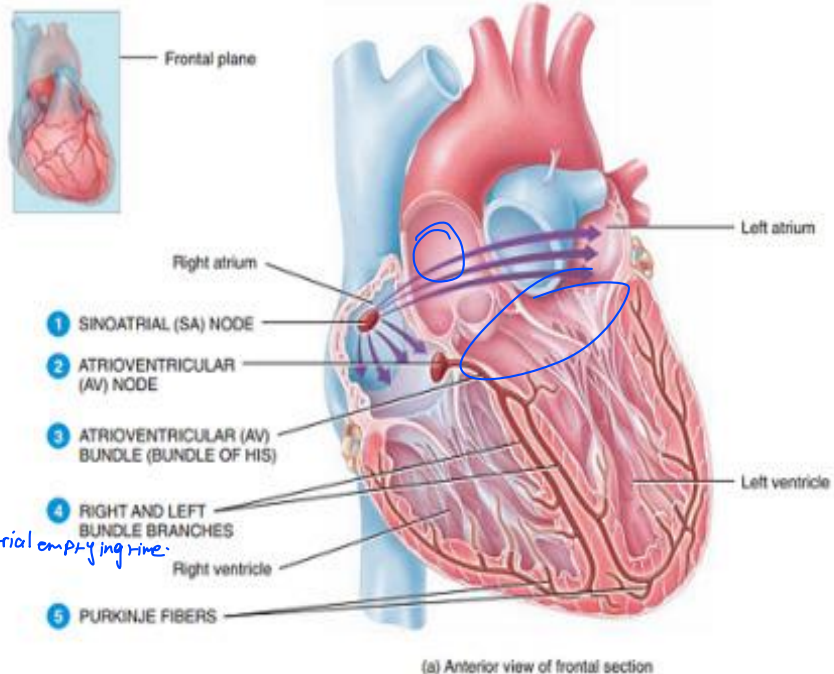
- By conducting along atrial muscle fibers, the **action potential reaches the atrioventricular (AV) node.**
- At the AV node, the **action potential slows** considerably as a result of various differences in cell structure in the AV node. This **delay provides time for the atria to empty their blood into the ventricles.**

عند العقدة AV يتباطأ جهد الفعل بشكل واضح نتيجة اختلافات في بنية خلايا هذه العقدة. هذا التأخير يمنح الأذينين الوقت ليفرغا دمهما في البطينين.

AV node = slow conduction = atrial emptying time.

Figure 20.10 The conduction system of the heart. Autorhythmic fibers in the SA node (a), act as the heart's pacemaker, initiating cardiac action potentials (b) that cause contraction of the heart's chambers.

 The conduction system ensures that the chambers of the heart contract in a coordinated manner.

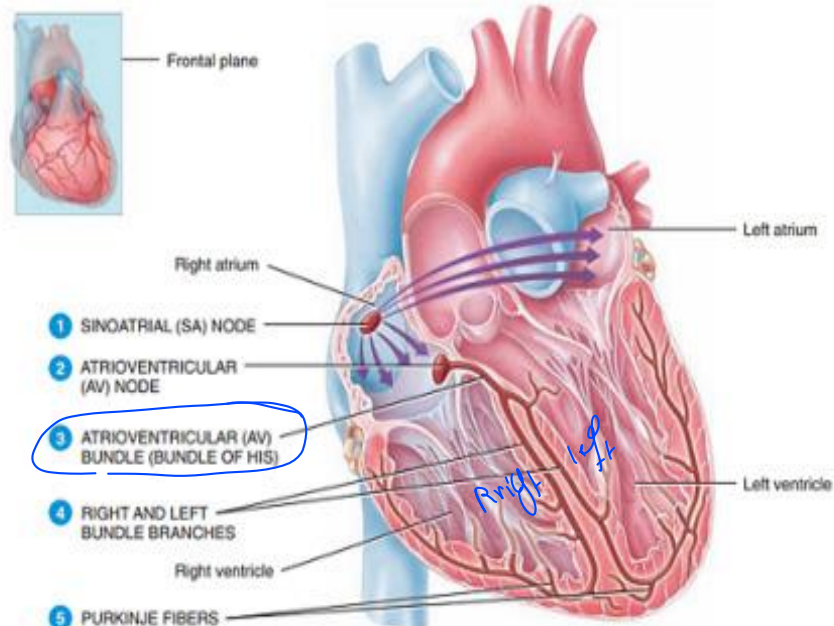


AUTORHYTHMIC FIBERS: THE CONDUCTION SYSTEM

- From the AV node, the action potential enters the atrioventricular (AV) bundle. This bundle is the only site where action potentials can conduct from the atria to the ventricles.
- After propagating through the AV bundle, the action potential enters both the right and left bundle branches.

Figure 20.10 The conduction system of the heart. Autorhythmic fibers in the SA node, located in the right atrial wall (a), act as the heart's pacemaker, initiating cardiac action potentials (b) that cause contraction of the heart's chambers.

The conduction system ensures that the chambers of the heart contract in a coordinated manner.



(a) Anterior view of frontal section

AUTORHYTHMIC FIBERS: THE CONDUCTION SYSTEM

- Finally, the **large-diameter Purkinje fibers** rapidly conduct the action potential beginning at the **apex of the heart** upward to the remainder of the ventricular myocardium. Then the **ventricles contract, pushing the blood upward toward the semilunar valves.**

homonogly, aorta Jicic - hole ← → intic - hole

- ① SA node → توليد الإشارة
- ② انتشار عبر الأذينين
- ③ AV node → تبطيء الإشارة
- ④ AV bundle (Bundle of His) → الطريق الوحيد للبطين
- ⑤ Right & Left bundle branches
- ⑥ Purkinje fibers → انقباض البطينين

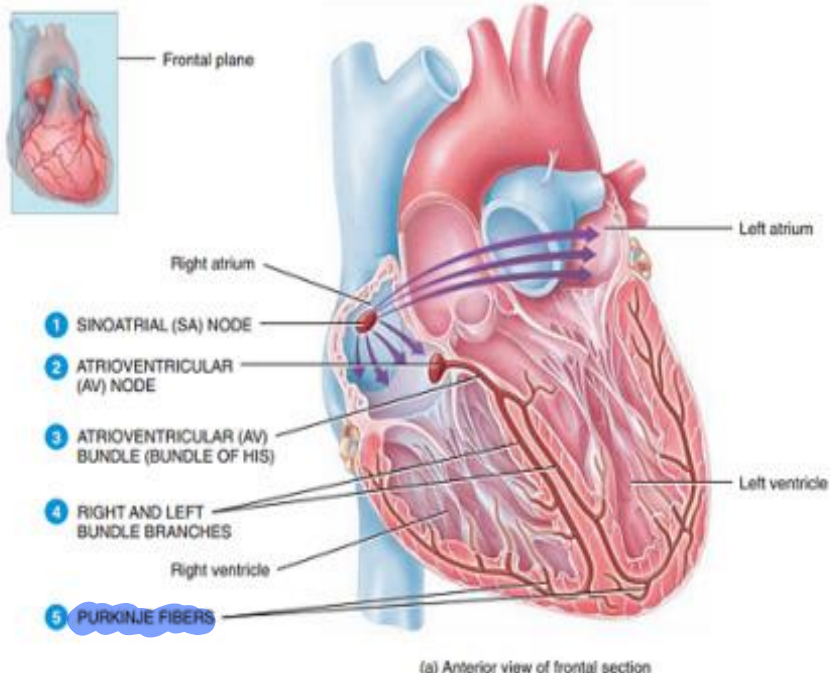
يعني الآن خلصنا كل نظام التوصيل الكهربائي للقلب

٤- **شو أحفظ؟ (المهم للامتحان)**

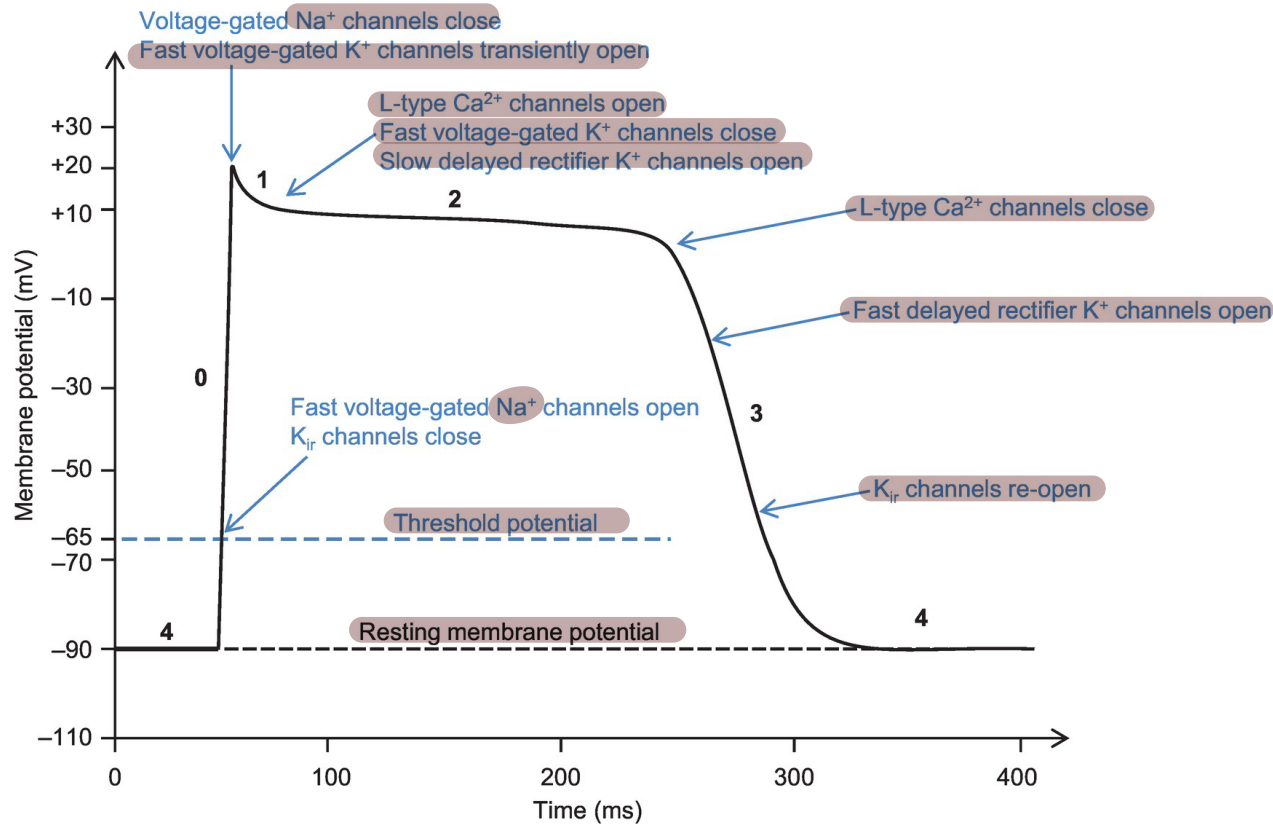
- أسرع اليف توصيل بالقلب سبب قطرها الكبيرة.
- ٢. ببدأ التوصيل 🔥
- ٣. السبب: دفع الدم بـ 🔥
- ٤. هي آخر محطة في 🔥
- ٥. مسؤولة عن **ventricular contraction** 🔥

Figure 20.10 The conduction system of the heart. Autorhythmic fibers in the SA node, located in the right atrial wall (a), act as the heart's pacemaker, initiating cardiac action potentials (b) that cause contraction of the heart's chambers.

6 The conduction system ensures that the chambers of the heart contract in a coordinated manner.



ACTION POTENTIAL AND CONTRACTION OF CONTRACTILE FIBERS



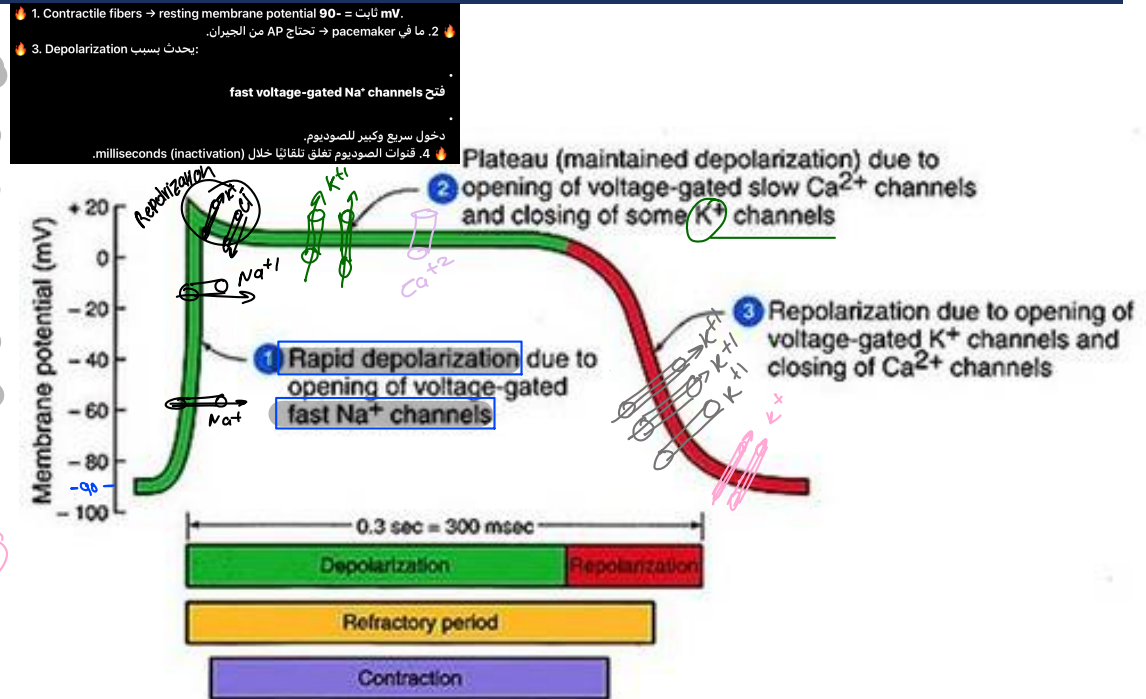
ACTION POTENTIAL AND CONTRACTION OF CONTRACTILE FIBERS

❖ Depolarization:

Unlike

autorhythmic fibers, **contractile fibers** have a stable resting membrane potential that is close to **-90 mV**.

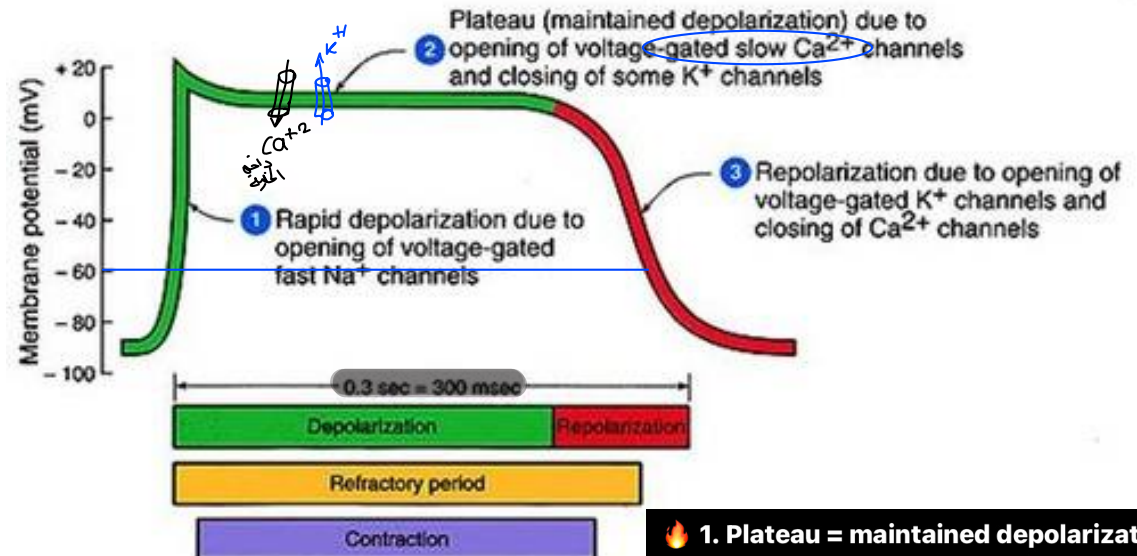
When a **contractile fiber** is brought to **threshold** by an action potential from neighboring fibers, its **voltage-gated fast Na ion channels open**. Inflow of **Na ions** down the electrochemical gradient produces a **rapid depolarization**. Within a few milliseconds, the **fast Na ion channels** automatically inactivate and **Na ions inflow decreases**.



rest membrane potential $\rightarrow \text{K}^+$
مسوّد عن عـ

ACTION POTENTIAL AND CONTRACTION OF CONTRACTILE FIBERS

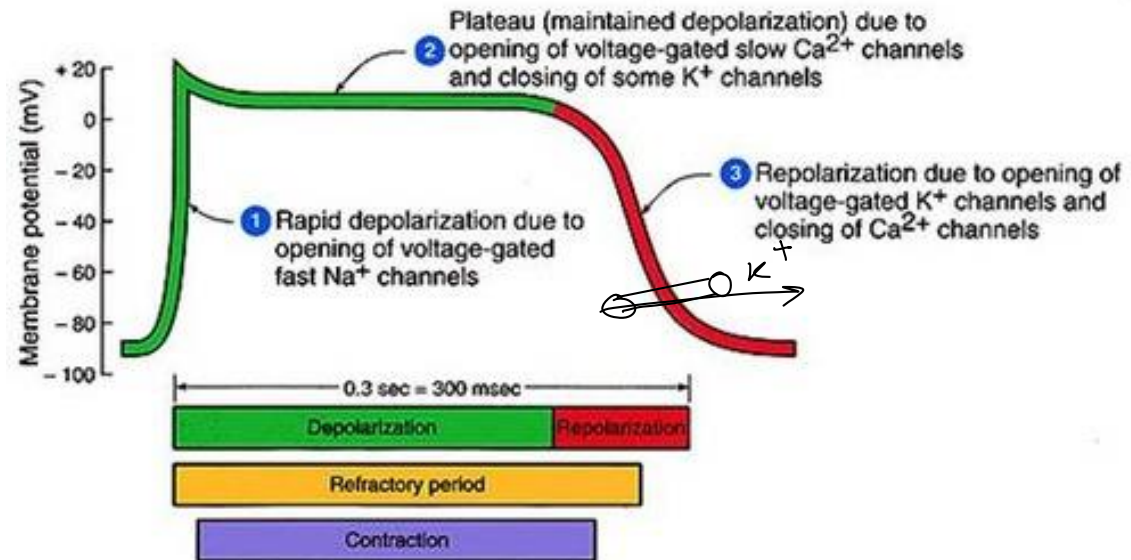
❖ **Plateau:** A period of maintained depolarization. It is due in part to opening of voltage-gated slow calcium ions channels in the sarcolemma. The increased calcium ions concentration in the cytosol ultimately triggers contraction. Several different types of voltage-gated potassium ions channels are also found in the sarcolemma of a contractile fiber (calcium ions inflow just balances potassium ions outflow).



1. Plateau = maintained depolarization slow Ca²⁺ channels
2. سبب الأساسي: فتح
3. Ca²⁺ inflow = trigger for contraction
4. بالتزامن يخرج K⁺ → يخلق توازن
5. أطول مرحلة في AP البطيني
6. تمنع حدوث tetanus (انقباض مستمر)

ACTION POTENTIAL AND CONTRACTION OF CONTRACTILE FIBERS

❖ **Repolarization:** After a delay (which is particularly prolonged in cardiac muscle), **additional voltage-gated potassium ions channels open**. Outflow of potassium ions restores the negative resting membrane potential (-90 mV). At the same time, the **calcium channels** in the **sarcolemma** and the **sarcoplasmic reticulum** are **closing**, which also contributes to repolarization.



ACTION POTENTIAL AND CONTRACTION OF CONTRACTILE FIBERS

- The **mechanism of contraction** is similar in cardiac and skeletal muscle:
 - ❖ The electrical activity (**action potential**) leads to the **mechanical response (contraction)** after a short delay.
 - ❖ As **calcium concentration rises inside a contractile fiber**, calcium ion binds to the regulatory protein troponin, which allows the **actin and myosin filaments to begin sliding past one another**, and tension starts to develop.
 - ❖ Substances that alter the movement of calcium ions through slow calcium ions channels influence the **strength of heart contractions**. **Epinephrine**, for example, increases contraction force by enhancing calcium ions flow into the cytosol.
- In muscle, the refractory period is the time interval during which a second contraction cannot be triggered. The refractory period of a cardiac muscle fiber lasts longer than the contraction itself . As a result, another contraction cannot begin until relaxation is well under way. Their **pumping function depends on alternating contraction (when they eject blood) and relaxation (when they refill)**.

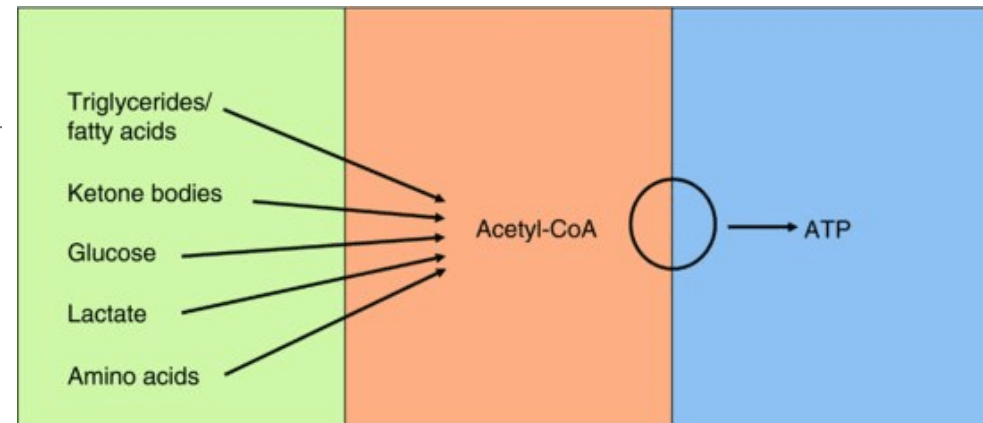
فترة المقاومة
لبعض صفات العضلات

الاسترخاء

ATP PRODUCTION IN CARDIAC MUSCLE

- In contrast to skeletal muscle, cardiac muscle produces little of the ATP it needs by anaerobic cellular respiration.
- Cardiac muscle fibers use several fuels to power mitochondrial ATP production. In a person at rest, the heart's ATP comes mainly from oxidation of fatty acids (60%) and glucose (35%), with smaller contributions from lactic acid, amino acids, and ketone bodies. During exercise, the heart's use of lactic acid, produced by actively contracting skeletal muscles, rises.

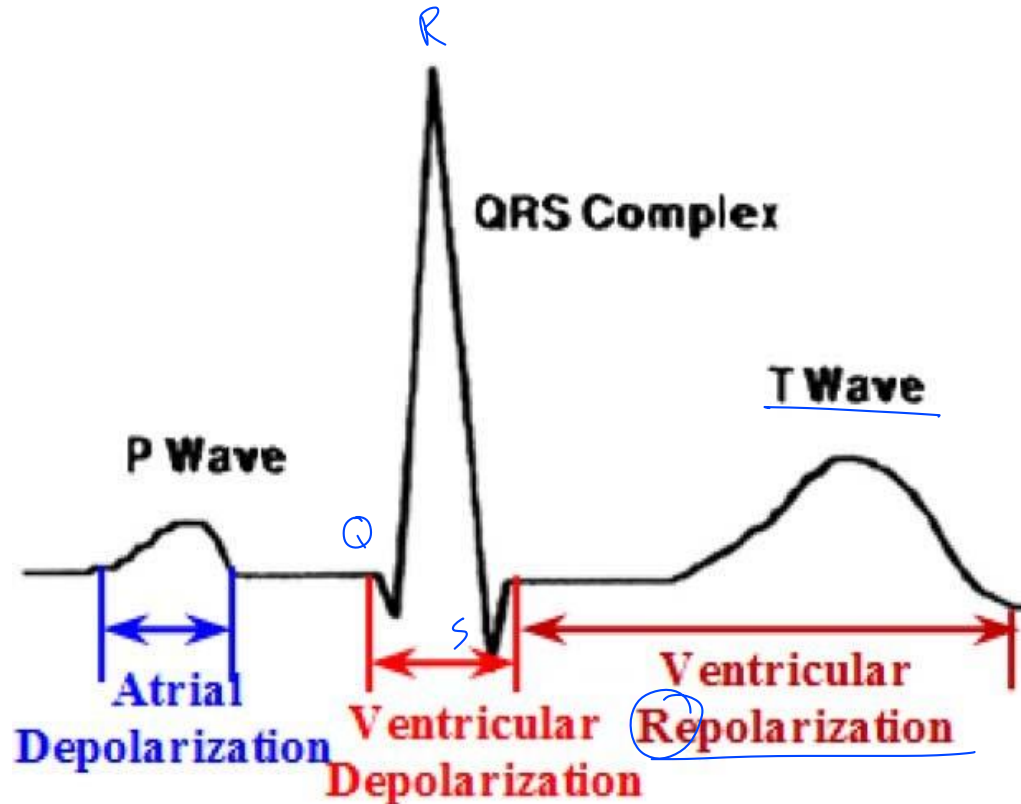
اللactic acid هو المُتجدد في العضلات





- As action potentials propagate through the heart, they generate electrical currents that can be detected at the surface of the body. An electrocardiogram, abbreviated either **ECG** or **EKG** (from the German word Elektrokardiogram), is a recording of these electrical signals.
- The instrument used to record the changes is an **electrocardiograph**.
- By comparing these records with one another and with normal records, it is possible to determine:
 - (1) if the conducting pathway is abnormal.
 - (2) if the heart is enlarged.
 - (3) if certain regions of the heart are damaged.
 - (4) the cause of chest pain.

ELECTROCARDIOGRAM



بيانات الـ ECG: دورة الكهربائية في القلب
Repolarization of Atrium

ELECTROCARDIOGRAM

- In reading an ECG, the size of the waves can provide clues to abnormalities.
وَالآن
- 1. Larger P waves indicate enlargement of an atrium.
- 2. An enlarged Q wave may indicate a myocardial infarction.
وَالآن
- 3. An enlarged R wave generally indicates enlarged ventricles.
- 4. The T wave is flatter than normal when the heart muscle is receiving insufficient oxygen—as, for example, in coronary artery disease. The T wave may be elevated in hyperkalaemia (high blood K ions level).
وَالآن

ELECTROCARDIOGRAM

فکات

- Analysis of an ECG also involves measuring the time **spans** between waves, which are called **intervals or segments**.
- **P–Q interval** is the time from the beginning of the P wave to the beginning of the QRS complex. It represents the conduction time from the beginning of atrial excitation to the beginning of ventricular excitation.
- The **S–T segment**, which begins at the end of the S wave and ends at the beginning of the T wave, represents the time when the ventricular contractile fibers are depolarized during the plateau phase of the action potential.

Segment: قطعة

interval: الفاصل \rightarrow wave + segment

ELECTROCARDIOGRAM

- The **Q-T interval** extends from the start of the QRS complex to the end of the T wave. It is the time from the beginning of ventricular depolarization to the end of ventricular repolarization. +

show the ECG clues the abnormality:-

? ECG ill to right ventricle

- 1 - conduction pathway abnorma.
- 2 - enlarged the heart muscle.
- 3 - cause of the chest pain.
- 4 - certain region of the heart damage

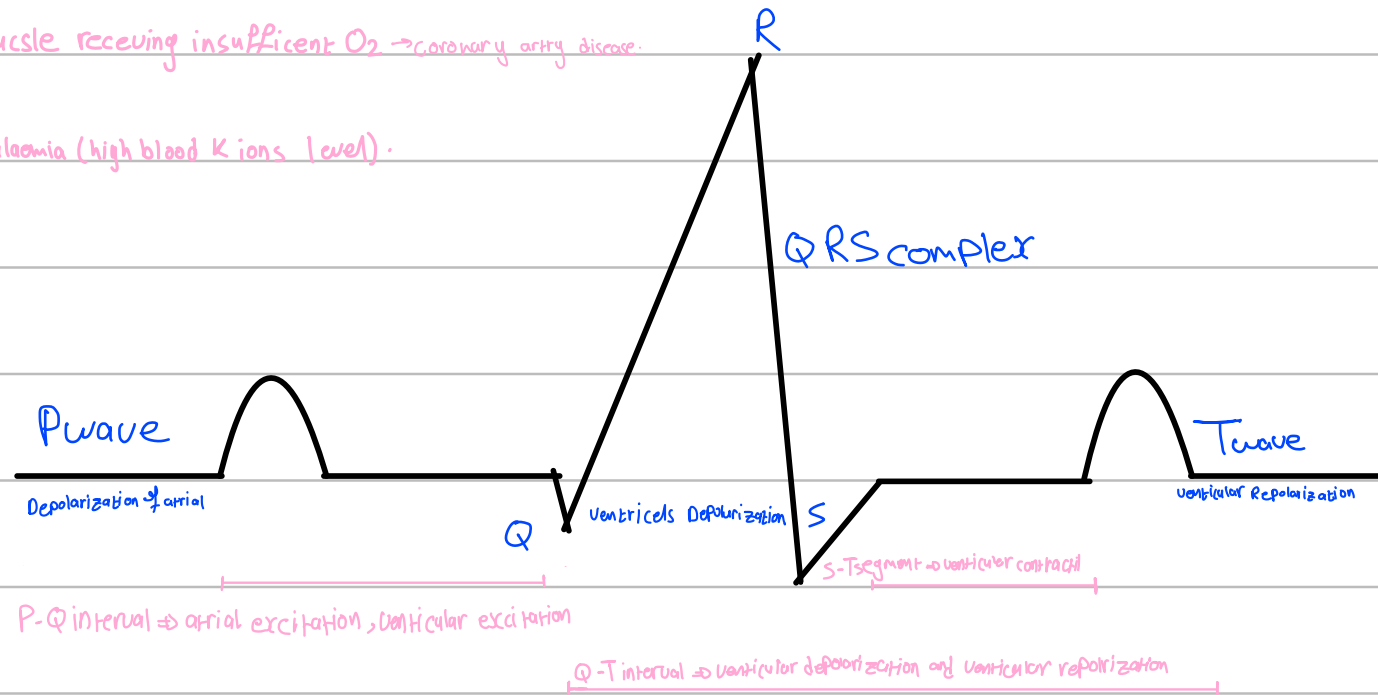
1 - large P wave → enlargement in atrium

2 - enlarged Q wave → myocardium infarction.

3 - enlarged R wave → enlargement in ventricles.

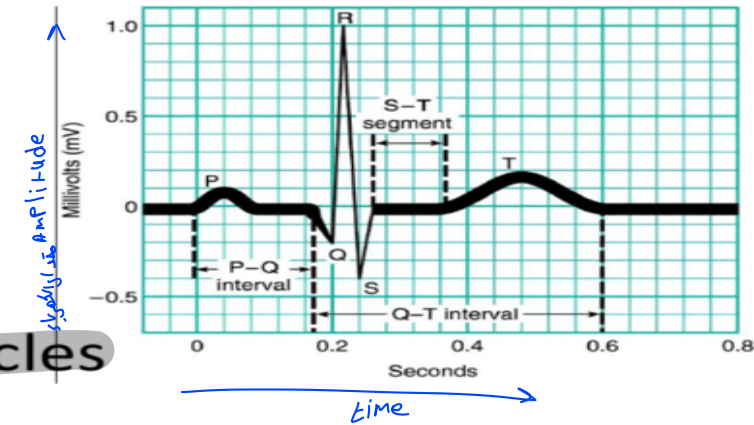
4 - T wave
more flatter → heart muscle receiving insufficient O₂ → coronary artery disease.

elevated → hyperkalaemia (high blood K⁺ level).



The Electrocardiogram

- The major deflections and intervals in a normal ECG include:
 - P wave** - atrial depolarization
 - P-Q interval** - time it takes for the atrial kick to fill the ventricles
 - QRS wave** - ventricular depolarization and atrial repolarization
 - S-T segment** - time it takes to empty the ventricles before they repolarize (the T wave)



CORRELATION OF ECG WAVES WITH ATRIAL AND VENTRICULAR SYSTOLE

- The term **systole** refers to the phase of contraction.
- The phase of relaxation is diastole.
- **The ECG waves predict the timing of atrial and ventricular systole and diastole.**
 - ❖ As the atrial contractile fibers depolarize, the P wave appears in the ECG.
 - ❖ After the P wave begins, the atria contract (atrial systole).
 - ❖ The action potential propagates rapidly again after entering the AV bundle. About 0.2 sec after onset of the P wave, it has propagated through the bundle branches, Purkinje fibers, and the entire ventricular myocardium.
 - ❖ Contraction of ventricular contractile fibers (ventricular systole) begins shortly after the QRS complex appears and continues during the S-T segment. جنب QRS جلطة
 - ❖ Repolarization of ventricular contractile fibers produces the T wave in the ECG about after the onset of the P wave.
 - ❖ Shortly after the T wave begins, the ventricles start to relax (ventricular diastole). Ventricular repolarization is complete and ventricular contractile fibers are relaxed.

1. Which statement best describes the onset of atrial systole?

- It occurs immediately after the QRS complex.
- It begins once the P wave has fully ended.
- It starts after the P wave begins.
- It starts 0.2 seconds after the onset of the P wave.

2. The P wave on the ECG directly represents:

- Repolarization of atrial fibers.
- Depolarization of atrial contractile fibers.
- Depolarization of ventricular fibers.
- Repolarization of ventricular contractile fibers.

3. Ventricular systole begins:

- At the onset of the T wave.
- Shortly after the QRS complex appears.
- Before the depolarization reaches the Purkinje fibers.
- When atrial systole ends.

4. The S-T segment corresponds to:

- The period before ventricular contraction begins.
- The duration of ventricular repolarization.
- The continuation of ventricular systole after it begins.
- Atrial depolarization spreading through the myocardium.

5. Ventricular repolarization is responsible for producing:

- The QRS complex.
- The P wave.
- The T wave.
- The PR segment.

6. The ventricles begin to relax:

- Immediately after the P wave ends.
- Shortly after the T wave begins.
- At the midpoint of the S-T segment.
- Simultaneously with atrial systole.

7. The passage of the action potential through the AV bundle and into the Purkinje fibers occurs:

- Just before the P wave begins.
- Approximately 0.2 seconds after the onset of the P wave.
- Immediately following ventricular systole.
- During ventricular diastole.

8. Which of the following correctly predicts the timing of systole and diastole?

- Mechanical events determine ECG waves.
- ECG waves occur after the mechanical events begin.
- ECG waves predict when atrial and ventricular systole and diastole will occur.
- Mechanical contraction produces the ECG waves.

9. The QRS complex occurs when:

- Ventricular systole ends.
- Atrial systole is fully completed.
- Ventricular contractile fibers are depolarized.
- Ventricular contractile fibers are repolarized.

10. Ventricular repolarization becomes complete during:

- Early ventricular systole.
- The end of the T wave.
- The onset of the S-T segment.
- The beginning of the QRS complex.

MCQ Answers

- C
- B
- B
- C
- C
- B
- B
- C
- C
- B

THE CARDIAC CYCLE: PRESSURE AND VOLUME CHANGES DURING THE CARDIAC CYCLE

Contraction, relaxation
= systole, diastole

○ Atrial Systole:

- Atrial depolarization causes atrial systole.
- The ventricles are relaxed (The end of atrial systole is also the end of ventricular diastole (relaxation)).

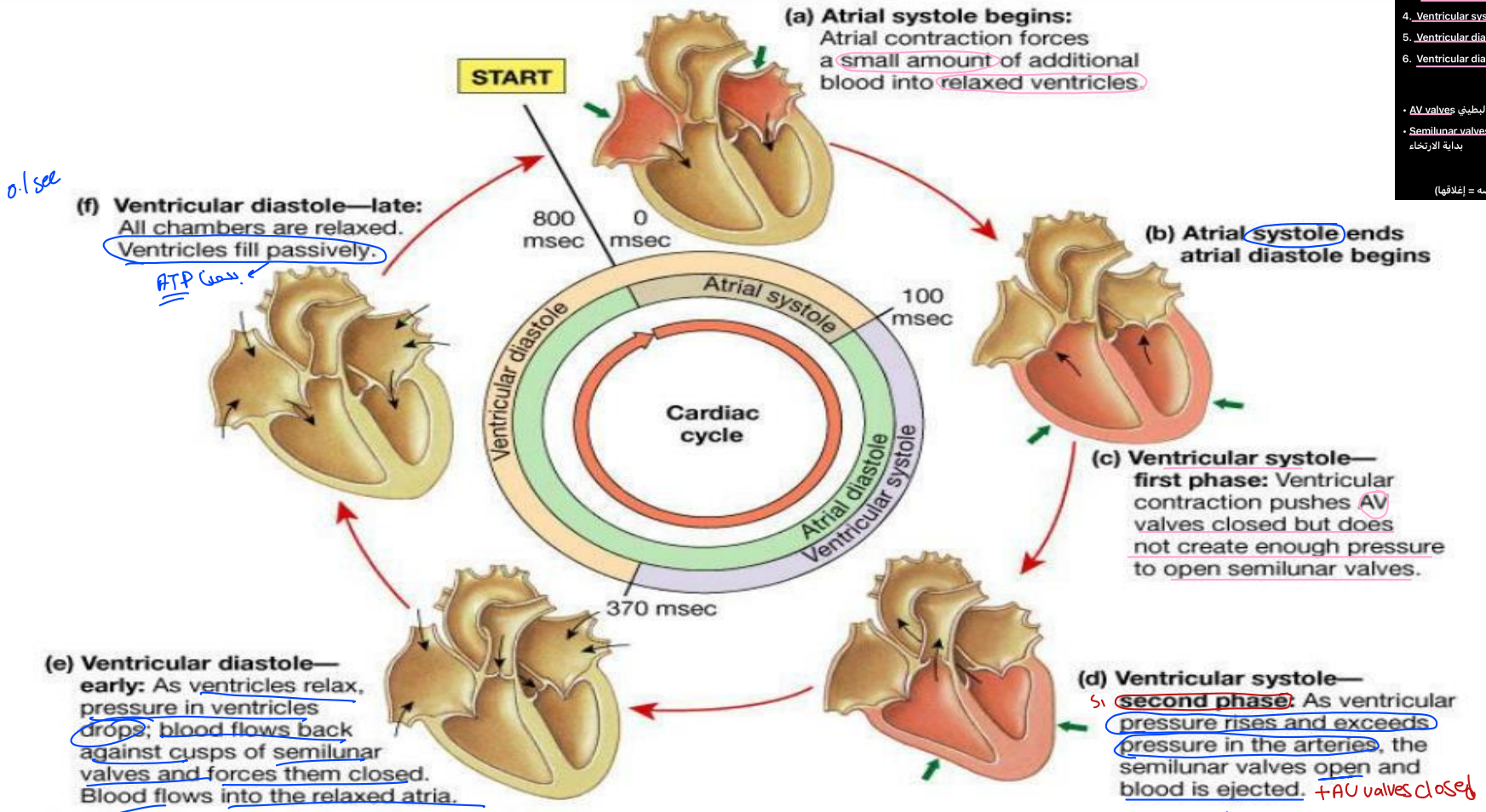
○ Ventricular Systole:

- The ventricles are contracting.
- At the same time, the atria are relaxed.

○ Relaxation Period:

- The atria and the ventricles are both relaxed.
- Ventricular repolarization causes **ventricular diastole**.

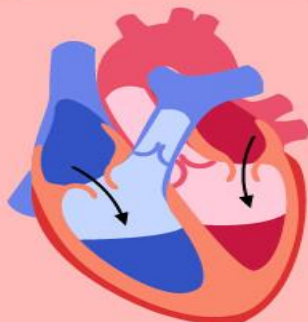
Figure 20.16 Phases of the Cardiac Cycle



PHASES OF THE CARDIAC CYCLE

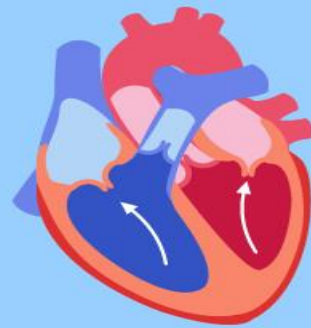
Atriole systole begins

Atrial contraction forces blood into ventricles



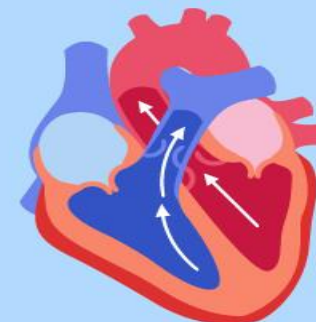
Ventricular systole (first phase)

Ventricular contraction pushes AV valves closed



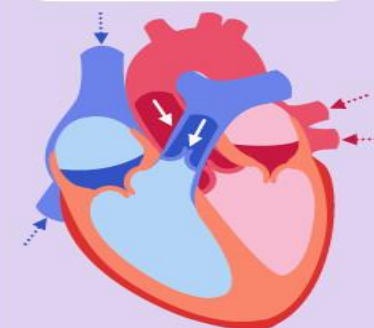
Ventricular systole (second phase)

Semilunar valves open and blood is ejected
AV valves closed \rightarrow S-e-lub-b



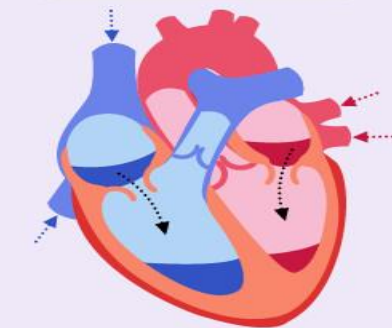
Ventricular diastole (early)

Semilunar valves close and blood flows into atria



Ventricular diastole (late)

Chambers relax and blood fills ventricles passively



P-Wave
Atria depolarization

R

Q S
QRS Complex
Ventricle depolarization

T

T - Wave
Ventricular repolarization

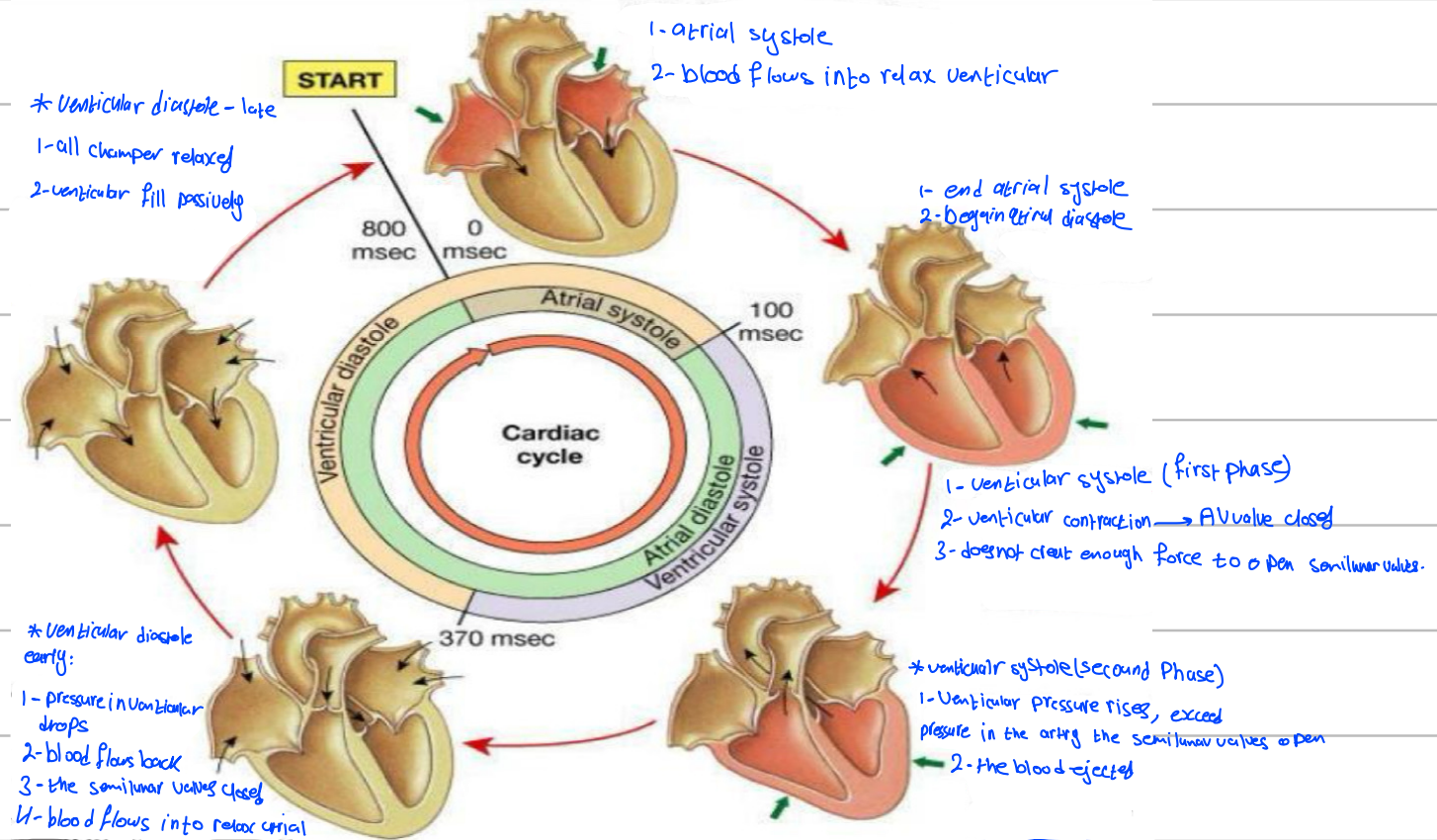
Atrial Diastole

Atrial Systole

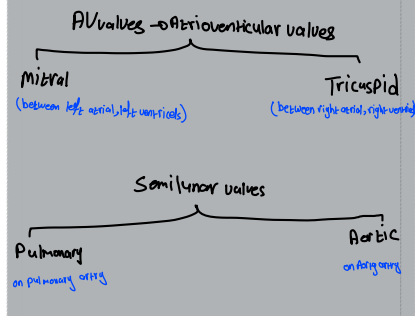
Ventricular Diastole

Atrial Diastole

Ventricular Diastole



Note:

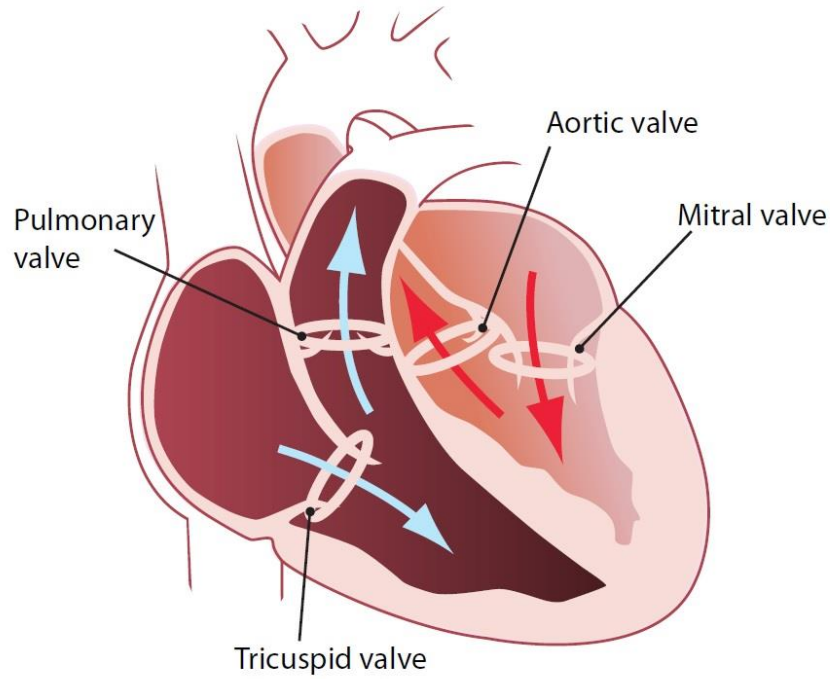


contraction $\xrightarrow{0.3 \text{ sec}}$ diastole $\xrightarrow{0.7 \text{ sec}}$ relaxation $\xrightarrow{0.3 \text{ sec}}$ diastole

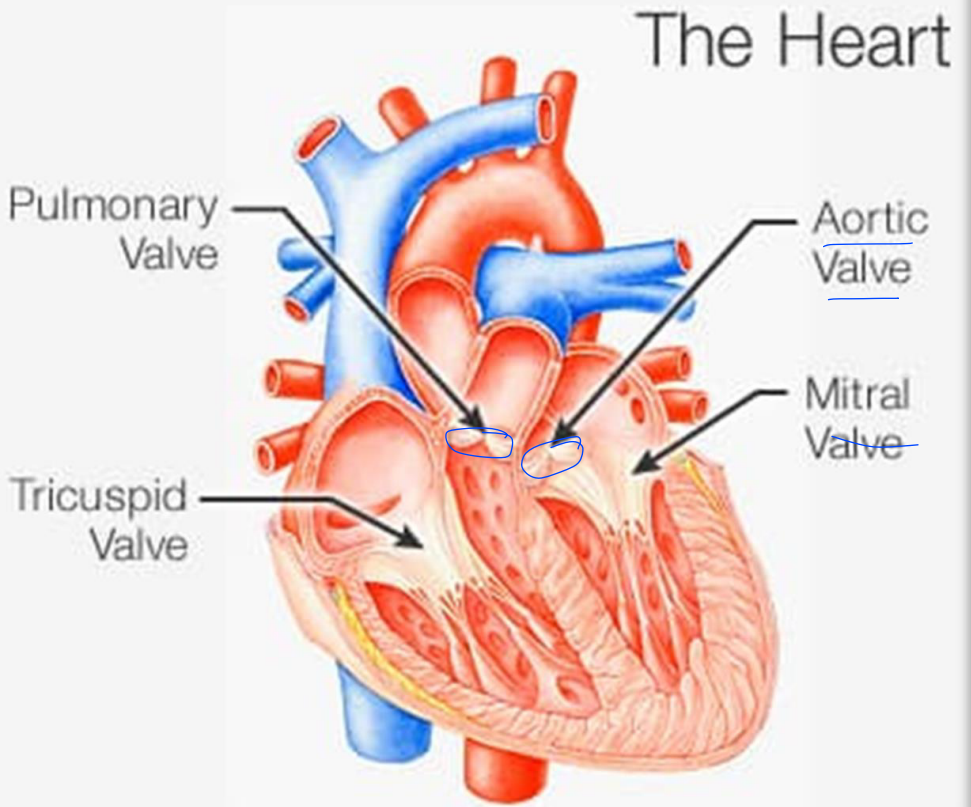
time

دعايات تسمى بركات الدم بال唼جه او اهداف خطط من المفهود الاعلى
على المفهود الاعلى

HEART VALVES



Semi-lunar valves Aortic, pulmonary valves



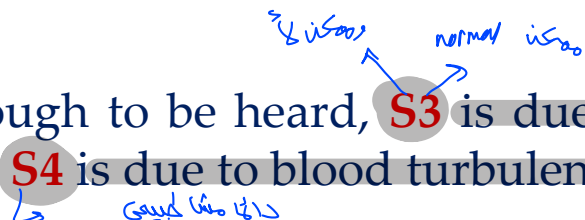
HEART SOUNDS

Auscultation: الاستماع إلى الأصوات الطبيعية والغير طبيعية
stethoscope: سمعة طبية

- **Auscultation**, the act of listening to sounds within the body, is usually done with a **stethoscope**.
- During each cardiac cycle, there are four heart sounds, but in a normal heart only the first and second heart sounds (**S1** and **S2**) are loud enough to be heard through a stethoscope.
- **The first sound (S1)**, which can be described as **a lubb sound**, is louder and a bit longer than the second sound. S1 is caused by **blood turbulence associated with closure of the AV valves soon after ventricular systole begins**.
- **The second sound (S2)**, which is shorter and not as loud as the first, can be described as a **dupp sound**. S2 is caused by **blood turbulence associated closure of the semilunar (aortic and pulmonary) valves at the beginning of ventricular diastole**.
- Normally not loud enough to be heard, **S3** is due to blood turbulence during **rapid ventricular filling**, and **S4** is due to blood turbulence during **atrial systole**

$S_1, S_2 \rightarrow$ **لوب** / **داب**

$S_3, S_4 \rightarrow$ **داب** / **لوب**



Heart sounds

- Auscultation – listening to heart sound via stethoscope
- Four heart sounds
 - S_1 – “lubb” caused by the closing of the AV valves
 - S_2 – “dupp” caused by the closing of the semilunar valves
 - S_3 – a faint sound associated with blood flowing into the ventricles \sim ywave
 - S_4 – another faint sound associated with atrial contraction



CARDIAC OUTPUT

خواص افعال

حجم الدم الى البطين
الAo من اورتیک
الAo

- **Cardiac output (CO)** is the volume of blood ejected from the left ventricle (or the right ventricle) into the aorta (or pulmonary trunk) each minute. Cardiac output equals **the stroke volume (SV)**, the volume of blood ejected by the ventricle during each contraction, multiplied by the **heart rate (HR)**, the number of heartbeats per minute:

$$\underline{CO} \text{ (mL/min)} = \underline{SV} \text{ (mL/beat)} \times \underline{HR} \text{ (beats/min)}$$

- **Cardiac reserve** is the difference between a person's **maximum cardiac output** and **cardiac output at rest**. The average person has a cardiac reserve of four or five times the resting value.

REGULATION OF STROKE VOLUME

حجم الدم الذي يُفخّج في خلية واحدة

- A healthy heart will pump out the blood that entered its chambers during the previous diastole.
- Three factors regulate stroke volume and ensure that the left and right ventricles pump equal volumes of blood: (1) preload the degree of stretch on the heart before it contracts; (2) contractility, the forcefulness of contraction of individual ventricular muscle fibers; and (3) afterload the pressure that must be exceeded before ejection of blood from the ventricles can occur.

PRELOAD: EFFECT OF STRETCHING

- Within limits, the more the heart fills with blood during diastole, the greater the force of contraction during systole. This relationship is known as the Frank–Starling law of the heart.
- The preload is proportional to the end-diastolic volume (EDV), (the volume of blood that fills the ventricles at the end of diastole). Normally, the greater the EDV, the more forceful the next contraction.
- Two key factors determine EDV: (1) the duration of ventricular diastole and (2) venous return, the volume of blood returning to the right ventricle.

CONTRACTILITY

Contractility علی‌نیز سرمهکایی را نیز می‌نامند

- ❑ **Myocardial contractility**, the strength of contraction at any given preload.
- ❑ Substances that increase contractility are **positive inotropic agents** (promote calcium ions inflow during cardiac action potentials), those that decrease contractility are **negative inotropic agents** (reducing calcium ions inflow).

AFTERLOAD

دیلی ایمیڈ میڈی

دیلی ایمیڈ میڈی

- Ejection of blood from the heart begins when pressure in the right ventricle exceeds the pressure in the pulmonary trunk, and when the pressure in the left ventricle exceeds the pressure in the aorta.
- At that point, the higher pressure in the ventricles causes blood to push the semilunar valves open. The pressure that must be overcome before a semilunar valve can open is termed the **afterload**.
- Conditions that can increase afterload include hypertension (elevated blood pressure) and narrowing of arteries by atherosclerosis.

REGULATION OF HEART RATE

■ Autonomic Regulation of Heart Rate:

النَّرْقَنُ الْقَلْبِيُّ الْمُعَنَّىُ

- ❖ Nervous system regulation of the heart originates in the cardiovascular center in the medulla oblongata. The cardiovascular center then directs appropriate output by increasing or decreasing the frequency of nerve impulses in both the **sympathetic and parasympathetic branches of the ANS.**
بِإِرْسَالِ الْأَعْصَارِ الْأَنْسَابِيِّةِ عَيْنَ دِلَاءَةٍ أَوْ تَعْلِيَةٍ لِرَدَدِ النَّفَّاعَاتِ الْعَلَيِّيَّةِ
- ❖ Proprioceptors that are monitoring the position of limbs and muscles send nerve impulses at an increased frequency to the cardiovascular center.
- ❖ Proprioceptor input is a major stimulus for the quick rise in heart rate that occurs at the onset of physical activity.
- ❖ Other sensory receptors that provide input to the cardiovascular center include chemoreceptors, which monitor chemical changes in the blood, and baroreceptors, which monitor the stretching of major arteries and veins caused by the pressure of the blood flowing through them. Important baroreceptors located in the arch of the aorta and in the carotid arteries.
 - Aortic arch → فوق القلب مباشرة
 - Carotid arteries → جانبي الرقبة

- ♦ **Proprioceptors** = مستقبلات حسية في العضلات والمفاصل تراقب وضع وحركة الجسم.

- ♦ أول ما تبدأي حركة → تبعث إشارات للميدولا → فيرفع نبض القلب مباشرة.
- ♦ هم أهم سبب للارتفاع السريع في **HR** عند بداية النشاط.

♦ شو تحفظي؟ (المختصر المفيد)

- 1. تنظيم الجهاز العصبي لنبض القلب يبدأ من الميدولا 
- 2. تتجمع المعلومات من المستقبلات في **Cardiovascular Center** 
- 3. المركز القلبي الوعائي يتحكم نبض القلب عن طريق:
 - Sympathetic ↑ HR 
 - Parasympathetic ↓ HR 

سم الغرفة: CVC أو Cardiovascular center

شو بتعمل هاي الغرفة؟

تراقب المعلومات اللي جاية من الجسم (مثل:
(proprioceptors, chemoreceptors, baroreceptors)

بعدين تقرّر:

- ٠ هل بدنا نسرّع نبض القلب؟
- ٠ ولا نبِطّئه؟

وبتبعث أوامرها عن طريق:

الأعصاب الوديّة (Sympathetic) → ترفع النبض

الأعصاب اللاؤدية (Parasympathetic) → تخفّض
لنبض

بعي... القرار النهائي لسرعة القلب يطلع من الميدو ولا.

* **Proprioceptors**: مستقبلات موجودة بالهيكلات بالذمار وبالظاهرات بباباواح كثة الحس

اول ما گفته شد که مسکن ملی ایران این سه میلیون هزار نفر است و بین سه و نه میلیون نفر باشد.

وهي في المولدة تحيط بالقلب **medulla oblongata** هي القسم الودي.

* Chemoreceptor: monitor chemical change

in the blood

* baroreceptors:- monitor stretching

of major arteries and veins caused by

the pressure blood inflowing

located in the arch of aorta and coronary arteries

REGULATION OF HEART RATE

- Autonomic Regulation of Heart Rate:
 - ❖ Through the sympathetic cardiac accelerator nerves: In SA (and AV) node fibers, norepinephrine speeds the rate of spontaneous depolarization so that these pacemakers fire impulses more rapidly and heart rate increases; in contractile fibers throughout the atria and ventricles, norepinephrine enhances calcium ions entry through the voltage-gated slow calcium ions channels, thereby increasing contractility.
Ca⁺⁺
 - ❖ Through Parasympathetic nerve impulses reach the heart via the right and left vagus (X) nerves: Vagal axons terminate in the SA node, AV node, and atrial myocardium. They release acetylcholine, which decreases heart rate by slowing the rate of spontaneous depolarization in autorhythmic fibers. As only a few vagal fibers innervate ventricular muscle, changes in parasympathetic activity have little effect on contractility of the ventricles.

* Sympathetic :-

1- cardiac accelerator nerve

2- SA, AV node \rightarrow NE \rightarrow speeds rate of spontaneous depolarization \rightarrow ↑ pacemaker fire impulses \rightarrow ↑ HR \rightarrow enhances Ca^{2+} entry
 \hookrightarrow voltage-gated slow

* Parasympathetic :-

1- Vagus axon terminates \rightarrow releases ACh \rightarrow ↓ slowing HR \rightarrow ↓ rate of spontaneous depolarization \rightarrow ↓ in autoregulatory fibers
SA node AV node atrial myocardium
few vagal fibers innervate to ventricular muscle \rightarrow ↓ chance of parasympathetic activation have a little effect on contracting of ventricles

① A utonomic activity

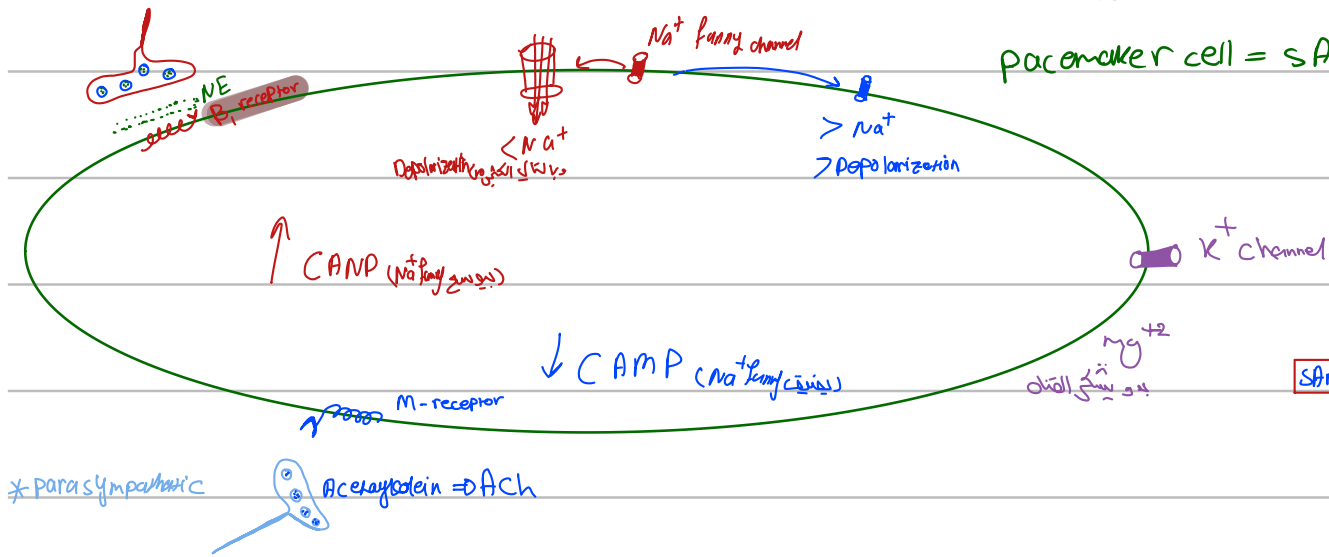
Diagram illustrating heart rate regulation:

Parasympathetic Pathway: Stimulus (e.g., *) → Parasympathetic → Ach (Acetylcholine) → M-receptor → ↓ cAMP → ↓ Inward current → ↓ heart rate

Sympathetic Pathway: Stimulus (e.g., *) → Sympathetic → NE (Norepinephrine) → ^{↓ cAMP} B receptor → ↑ cAMP → ↑ Inward current → ↑ heart rate

Annotations: "↓ cAMP" is written above the arrow from NE to B receptor. "↓ Inward current" and "↑ Inward current" are written above the arrows from cAMP to heart rate. "↓ heart rate" and "↑ heart rate" are circled in pink.

pacemaker cell = SA node cell \rightarrow en



شنايدر \rightarrow يزوج المبادرات المعاكسة التي ينطلق من الـ sympathetic

Tachycardia = heart rate 110 beats/min. Symptomatic with ≥ 100 beats/min.

• يوجد اصحاب خارجية تقبل لغتهم ولكن ليس المنشئ الكوادر بـ للتقطهم بها.

CHEMICAL REGULATION OF HEART RATE

hyperthyroidism:
sign of ↑ HR during sleep

NE, E \rightarrow ↑ heart rate effectiveness

↑ HR
+ Ca^{2+} + thyroid hormones

الدورة الكهربائية (فرقة القلب)

1. **Hormones:** Epinephrine and norepinephrine (from the adrenal medullae) enhance the heart's pumping effectiveness. These hormones affect cardiac muscle fibers in much the same way as does norepinephrine released by cardiac accelerator nerves—they increase both heart rate and contractility. One sign of hyperthyroidism (excessive thyroid hormone) is tachycardia, an elevated resting heart rate.

2. **Cations:** Given that differences between intracellular and extracellular concentrations of several cations (for example, sodium and potassium ions) are crucial for the production of action potentials in all nerve and muscle fibers. Elevated blood levels of potassium ions or sodium ions decrease heart rate and contractility. Excess sodium ions blocks calcium inflow during cardiac action potentials, thereby decreasing the force of contraction, whereas excess potassium ions blocks generation of action potentials. A moderate increase in interstitial (and thus intracellular) calcium ions level speeds heart rate and strengthens the heartbeat.

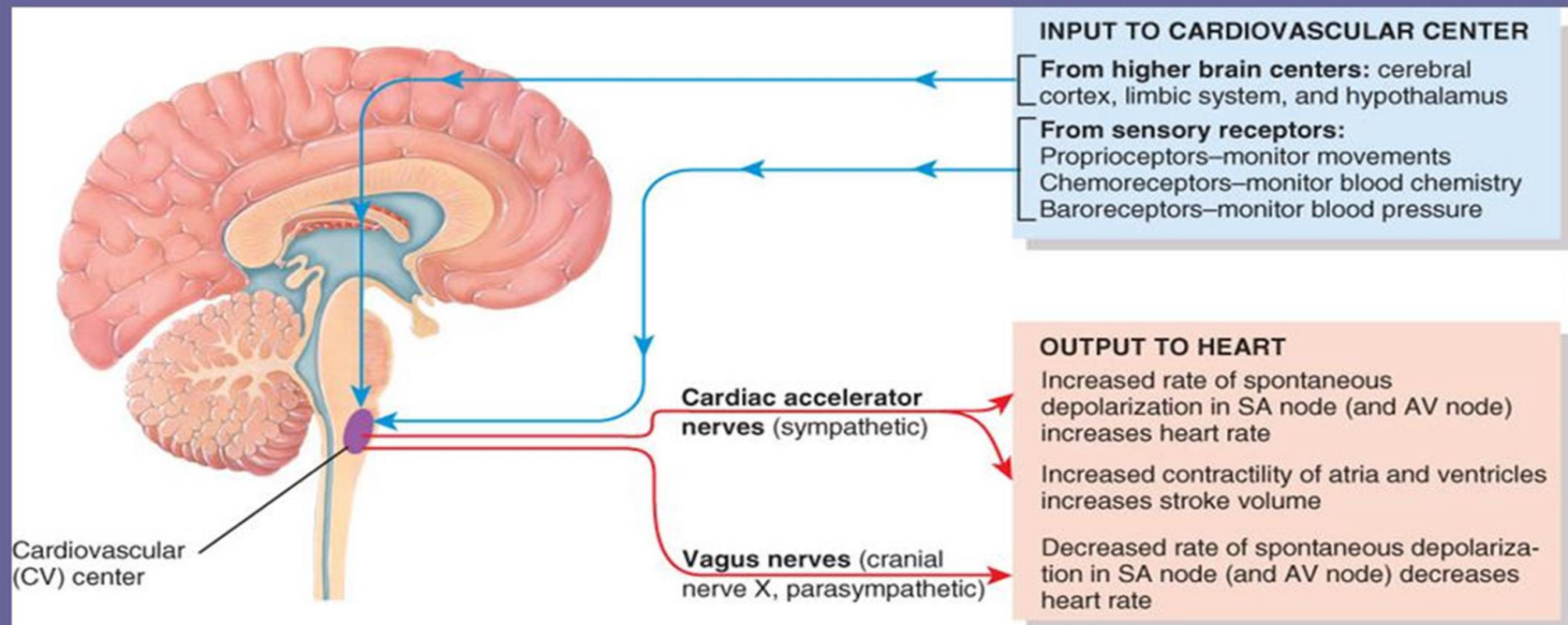
* E, NE \rightarrow enhances the heart's pumping effectiveness

$Ca^{2+} \rightarrow$ ↑ HR \rightarrow ↑ contractility
 $K^+ \rightarrow$ ↓ HR \rightarrow ↓ contractility
 $Na^+ \rightarrow$ ↓ HR \rightarrow ↓ contractility

الدورة الكهربائية

↑ HR \rightarrow ↑ contractility

Regulation of Heart Rate



OTHER FACTORS IN HEART RATE REGULATION

- Age, gender, physical fitness, and body temperature also influence resting heart rate.

كل درجة الحرارة تزيد على متوسطها
كل درجة اعلى من درجة متوسطها
- A physically fit person may even exhibit bradycardia, a resting heart rate under 50 beats/min.
- During surgical repair of certain heart abnormalities, it is helpful to slow a patient's heart rate by hypothermia, in which the person's body is deliberately cooled to a low core temperature.

أولاً: Neural Regulation (التنظيم العصبي)

1. Sympathetic (الجهاز الودي)

- ينطلق من: **cardiac accelerator nerves**
- يفرز: **norepinephrine**
- التأثير:
 - ↑ سرعة النبض (يزيد معدل depolarization لل SA node)
 - ↑ قوة الانقباض (يزيد دخول الكالسيوم لعضلة القلب)

2. Parasympathetic (الجهاز نظير الودي)

- ينطلق من: **vagus nerve** (العصب الحائر)
- يفرز: **acetylcholine**
- التأثير:
 - ↓ يقلل سرعة النبض
 - لا يؤثر على قوة الانقباض تقربياً

ثالثاً: Other Factors (العوامل الأخرى)

العامل	التأثير
العمر	معدل القلب يميل للانخفاض مع العمر
الجنس	النساء نبضهن أعلى قليلاً من الرجال
اللياقة البدنية	الرياضي → Bradycardia (أقل من 50 bpm)
في الجراحة Hypothermia	↑ حرارة → ↑ معدل القلب → حرارة → ↓ معدل القلب يُستخدم لتقليل معدل القلب وحماية عضلة القلب

ملخص الحفظ الإجباري (ثبتت سريع)

♦ Sympathetic → NE → ↑ HR + ↑ Contractility

♦ Parasympathetic → ACh → ↓ HR

♦ Epinephrine/Norepinephrine → (من الغدة الكظرية) → ↑ HR

♦ Hyperthyroidism → Tachycardia

♦ ↑ Na⁺ → ↓ القوة

♦ ↑ K⁺ يمنع → ↓ AP → ↓ HR

♦ ↑ Ca²⁺ → ↑ HR + ↑ القوة (معتدل)

♦ Athlete → Bradycardia

♦ Hypothermia → ↓ HR (مفيدة بالجراحة)

ثانياً: Chemical Regulation (التنظيم الكيميائي)

1. الهرمونات

• Epinephrine + Norepinephrine من الـ adrenal medulla

: تعمل مثل الأعصاب الودية تماماً

• ↑ تزيد معدل القلب

• ↑ تزيد قوة الانقباض

علامة مهمة سريرياً

• Hyperthyroidism يسبب → Tachycardia

(زيادة هرمون الغدة الدرقية ترفع النبض حق أثناء الراحة)

2. الأيونات

الصوديوم ⁺Na

إذا زاد كثيراً → ↓ يقلل قوة الانقباض

السبب: يمنع دخول الكالسيوم إلى الخلية

البوتاسيوم ⁺K

إذا زاد → ↓ يمنع توليد جهد الفعل → يقلل معدل القلب بشدة

الكالسيوم ²⁺Ca

زيادة معتدلة → ↑

↑ يزيد معدل القلب

↑ يزيد قوة الانقباض

HELP FOR FAILING HEARTS

وہاں option ہے

- Cardiac transplantation is the replacement of a severely damaged heart with a normal heart from a brain-dead or recently deceased donor.

وہاں ملکاں پر لے لیں

- Cardiac transplants are performed on patients with end-stage heart failure or severe coronary artery disease.

کوئی گلہ بیوی

کوئی

کوئی نہیں



THANK YOU

AMJADZ@HU.EDU.JO