



PHYSIOLOGY

بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ



FINAL | Lecture 7

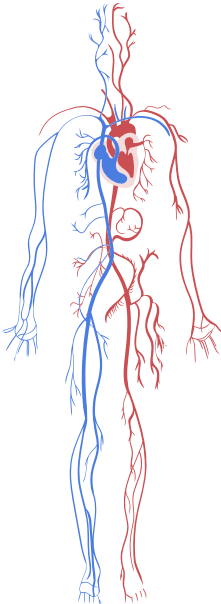
Special Circulations

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وَلَقَدْ خَلَقْنَا الْإِنْسَانَ وَنَعْلَمُ مَا تُوَسْوِسُ بِهِ نَفْسُهُ وَنَحْنُ أَقْرَبُ إِلَيْهِ مِنْ حَبْلِ الْوَرِيدِ
اللهم إنا نعوذ بك من شرور أنفسنا ومن سيئات أعمالنا

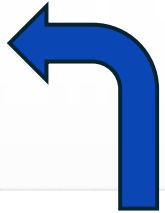


وَلِلّٰهِ الْأَسْمَاءُ الْحُسْنَىٰ فَادْعُوهُ بِهَا

المعنى: الذي سلم من العيوب والنقائص لكماله وكمال صفاته وأفعاله، وهو الذي يؤمن
الخلائق وحده ويسلمهم.

الورود: ورد مرة واحدة في القرآن.

الشاهد: ﴿الْمَلِكُ الْقُدُّوسُ السَّلَامُ﴾ [الحشر: ٢٣].



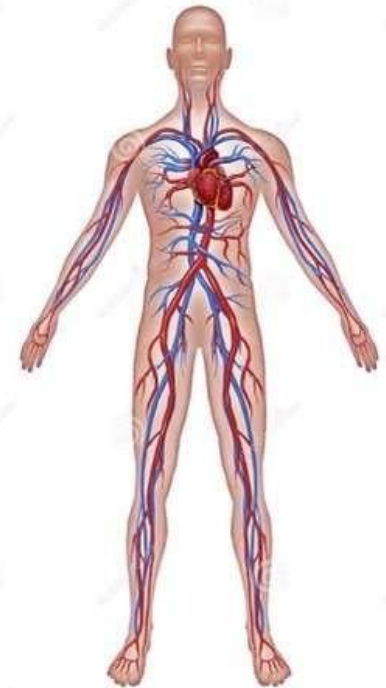
اضغط هنا لشرح أكثر تفصيلاً



Vascular Physiology

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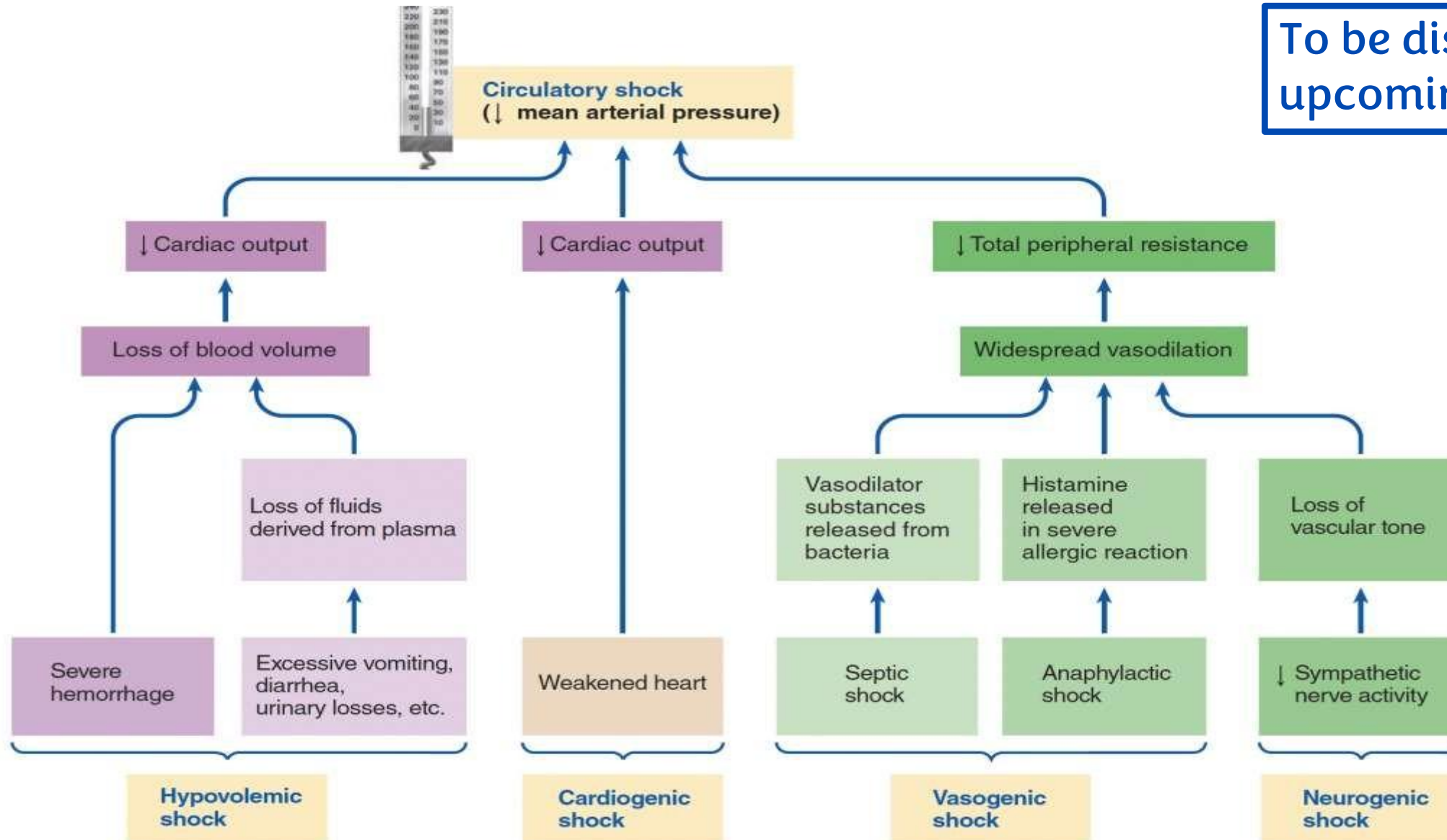


Pre-studying Notice

- It is highly recommended to tackle this lecture only after finishing all previous lectures because multiple previously studied concepts are required to fully understand this lecture.
- For this file not to reach a very high slide count, and to avoid repeating things, some details are thinned down if they have been extensively discussed in previous lectures.
- The focus of this file is more on this lecture's material; nevertheless, essential details will be pointed out even if they are not new.

Circulatory Shock

To be discussed in the upcoming slides...



Circulatory Shock

Circulatory shock is an inadequate tissue perfusion due to failure of the cardiovascular system, usually (not always) accompanied with Low BP.

➤ Causes of Circulatory Shocks include:

1. Hypovolemic Shock: Excessive Loss of intravascular volume, leading to:

↓ venous return → ↓ Cardiac Output → ↓ MAP → ↓ tissue perfusion.

➤ Types of Volume Loss include either:

A. Whole Blood Loss: Hemorrhage

B. Plasma Loss (Through excessive ECF loss), Ex: Vomiting, Diarrhea, Burns, Dehydration, Severe sweating

2. Cardiogenic shock: is a state in which the heart fails to pump blood adequately.

It can occur due to ischemia, severe heart failure, or arrhythmias, and carries an extremely low survival rate.

Circulatory Shock

3. Neurogenic Shock:

- Cause: Neurological impairment affecting the cardiovascular center, hypothalamus, or other parts of the central nervous system that leads to loss of tonic sympathetic stimulation to the blood vessels → widespread vasodilation
- Result: Loss of the vascular pressure required to drive blood through the vessels → Reduced blood flow to organs and tissues.

4. Vasogenic Shock

- Cause: release of vasodilator substances
- Result: widespread vasodilation → drop in TPR and pressure → reduced blood flow to organs.

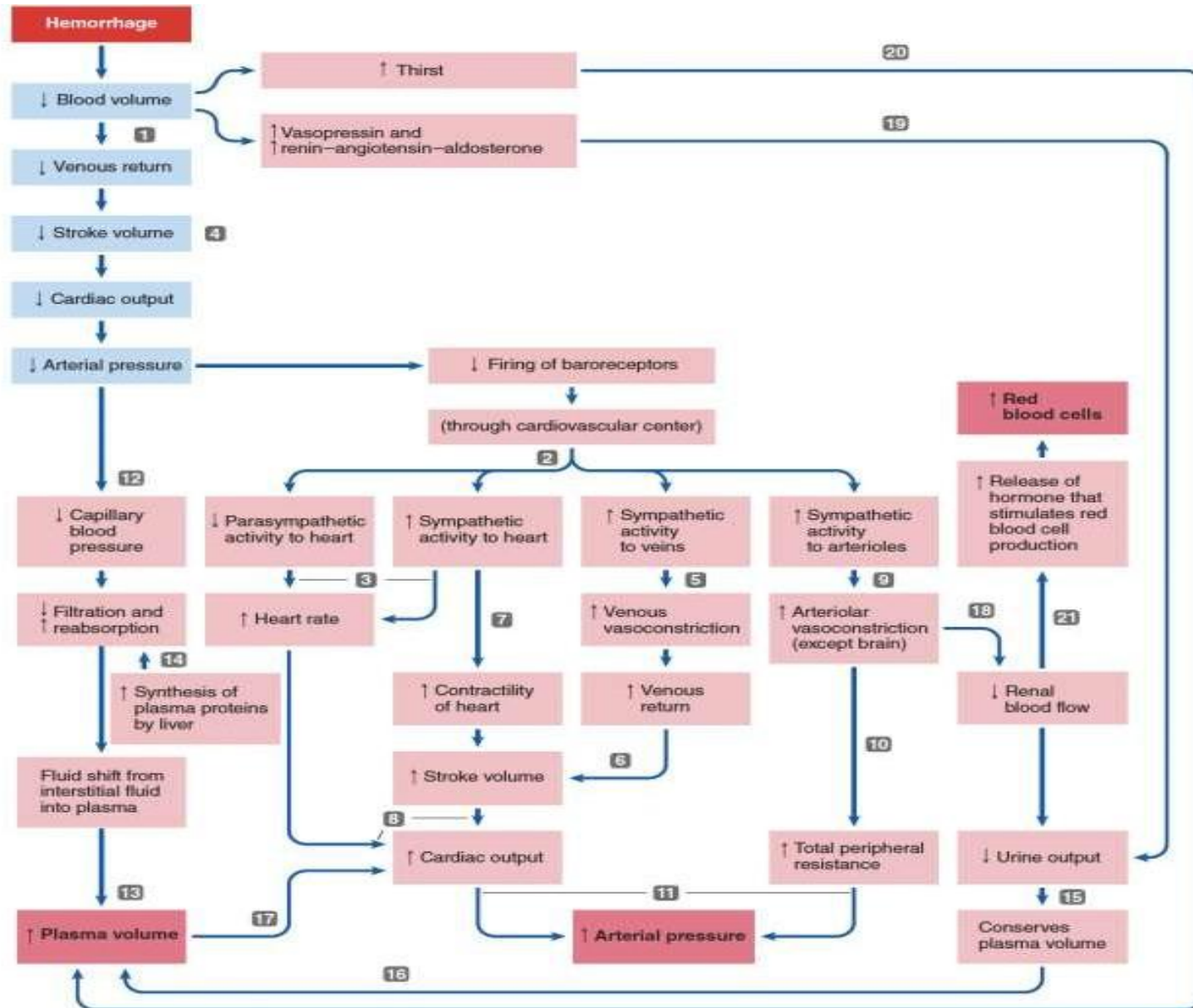
Occurs in two main pathways:

1. Allergic reaction: Allergen exposure → histamine release → marked vasodilation (Anaphylactic Shock)
2. Infection / bacterial toxins: Bacterial toxins → excessive vasodilator release → sepsis (Septic Shock)

Stages of Circulatory Shock

- Compensated
- Progressive
- Irreversible

Stages of circulatory Shock



To be discussed in the upcoming slides...

Stages of Circulatory Shock

1- Compensated Stage (As the Circulatory Shock starts): body activates **compensatory mechanisms** to restore blood pressure and maintain circulation, these mechanisms include:

➤ **Sympathetic Activation, leading to:**

→ **increased sympathetic tone**

→ **↑ heart excitability & contractility & ↑ heart rate**

→ **vasoconstriction** to raise systemic vascular resistance and venous return

➤ **Capillary Fluid Shift:**

Low capillary pressure leads to **↑ reabsorption of fluid from the interstitial space into the capillaries**

→ **helps restore intravascular volume**

➤ **Renal Response (RAAS Activation):**

Vasoconstriction leads to **↓ renal blood flow** → kidneys sense decreased perfusion → **RAAS activation:**

→ **↑ sodium reabsorption & ↑ water retention**

→ **↑ Further induces vasoconstriction**

➤ **Hormonal Mechanisms:**

Shock state leads to

→ **ADH release, which:**

→ **promotes water retention**

➤ **↑ Thirst (Polydipsia)**

Stages of Circulatory Shock

- If the compensatory mechanisms **restore blood pressure and perfusion**, the condition in this case is called "**Compensated Shock**"; however, if these mechanisms **fail** to restore adequate circulation the patient enters the "**Progressive Stage**".

2. **Progressive Stage** (Failure of compensatory mechanisms):

- Tissue perfusion continues to decline
- Medical intervention becomes **necessary**, **Ex: blood transfusion, fluids, vasopressors**
- If left without treatment the shock will continue to worsen, leading to the **Irreversible Stage**

3. **Irreversible Stage** (e.g., massive ischemic cardiogenic shock):

- **Irreversible tissue and organ damage (even if the circulation is restored).**
- **Medical intervention in this stage will not resolve the case; death is expected.**

Special Circulations

- **Coronary** circulation
- **Cerebral** circulation
- **Pulmonary** circulation
- **Renal** circulation
- **Skeletal muscle** circulation & Exercise
- **Skin** circulation & Temperature regulation

Generally, systemic control of blood flow to tissues is regulated by the sympathetic system; however, some tissues rely more on local control mechanisms to obtain either constant blood flow in cases of decreased perfusion, or increased blood flow in cases of increased demand.

Coronary circulation

- An unusual feature of the coronary circulation is the effect of mechanical compression of the blood vessels during systole in the cardiac cycle.
- Meaning that unlike all the vasculature, the coronary arteries gets supplied during the **diastolic heart phase** instead of the systolic phase. This mechanism of perfusion makes the coronary arteries completely dependent on **reactive hyperemia** in their supply (blood supply temporarily blocked during systole → accumulation of metabolites → induction of a strong (reactive) vasodilation once the blood supply is restored (during diastole)).
- This compression causes a brief period of occlusion and reduction of blood flow.
- When the period of occlusion (i.e., systole) is over, reactive hyperemia occurs to increase blood flow and O₂ delivery.

Coronary circulation

- The vital function of the heart makes it essential to maintain its blood supply regardless of the body's systemic regulatory responses. For this reason, the coronary arteries are specialized vessels that **rely primarily on local metabolic controls rather than systemic regulation**, ensuring that coronary blood flow is maintained—even when the rest of the body undergoes generalized vasoconstriction due to a systemic sympathetic response.
- Metabolic factors, especially myocardial oxygen consumption, are the major controllers of myocardial blood flow.
- Whenever the direct effects of nervous stimulation reduce coronary blood flow, the metabolic control of coronary flow usually overrides the direct coronary nervous effects within seconds.

Cerebral circulation

- The cerebral circulation is controlled almost entirely by local metabolites and exhibits autoregulation and active and reactive hyperemia.
- The brain's top priority is to maintain its blood flow, which is why sympathetic stimulation has no strong effect on cerebral circulation.
- Cerebral blood flow is regulated mainly by local factors.
- The most important local vasodilator in the cerebral circulation is CO₂ (or H⁺). An increase in cerebral PCO₂ (producing an increase in H⁺ concentration and a decrease in pH) causes vasodilation of the cerebral arterioles, which results in an increase in blood flow to assist in removal of the excess CO₂.
- It is interesting that many circulating vasoactive substances do not affect the cerebral circulation because their large molecular size prevents them from crossing the blood-brain barrier.

Pulmonary circulation

- The pulmonary circulation is controlled by O₂.
- Hypoxia induces vasodilation in the systemic circulation. When hypoxia occurs, metabolically active tissues are consuming more oxygen, so vasodilation occurs in the systemic circulation to increase blood flow and supply the tissues with their oxygen demand.
- The effect of O₂ on pulmonary arteriolar resistance is the exact opposite of its effect in other vascular beds.
- In the pulmonary circulation, hypoxia causes vasoconstriction.
- Regions of hypoxia in the lung cause local vasoconstriction, which effectively shunts blood away from poorly ventilated areas where the blood flow would be “wasted” and toward well-ventilated areas where gas exchange can occur.

Renal circulation

- Renal blood flow is tightly autoregulated so that flow remains constant even when renal perfusion pressure changes.
- Renal autoregulation is independent of sympathetic innervation, and it is retained even when the kidney is denervated (e.g., in a transplanted kidney).

Skeletal muscle circulation

- The degree of vasoconstriction of skeletal muscle arterioles is a major determinant of TPR because the mass of skeletal muscle is so large, compared with that of other organs.
In skeletal vasoconstriction, TPR increases, and vice versa.
- Blood flow to skeletal muscle is controlled both by local metabolites and by sympathetic innervation of its vascular smooth muscle.
This depends on the activity of the muscles (see next slide).

Skeletal muscle circulation

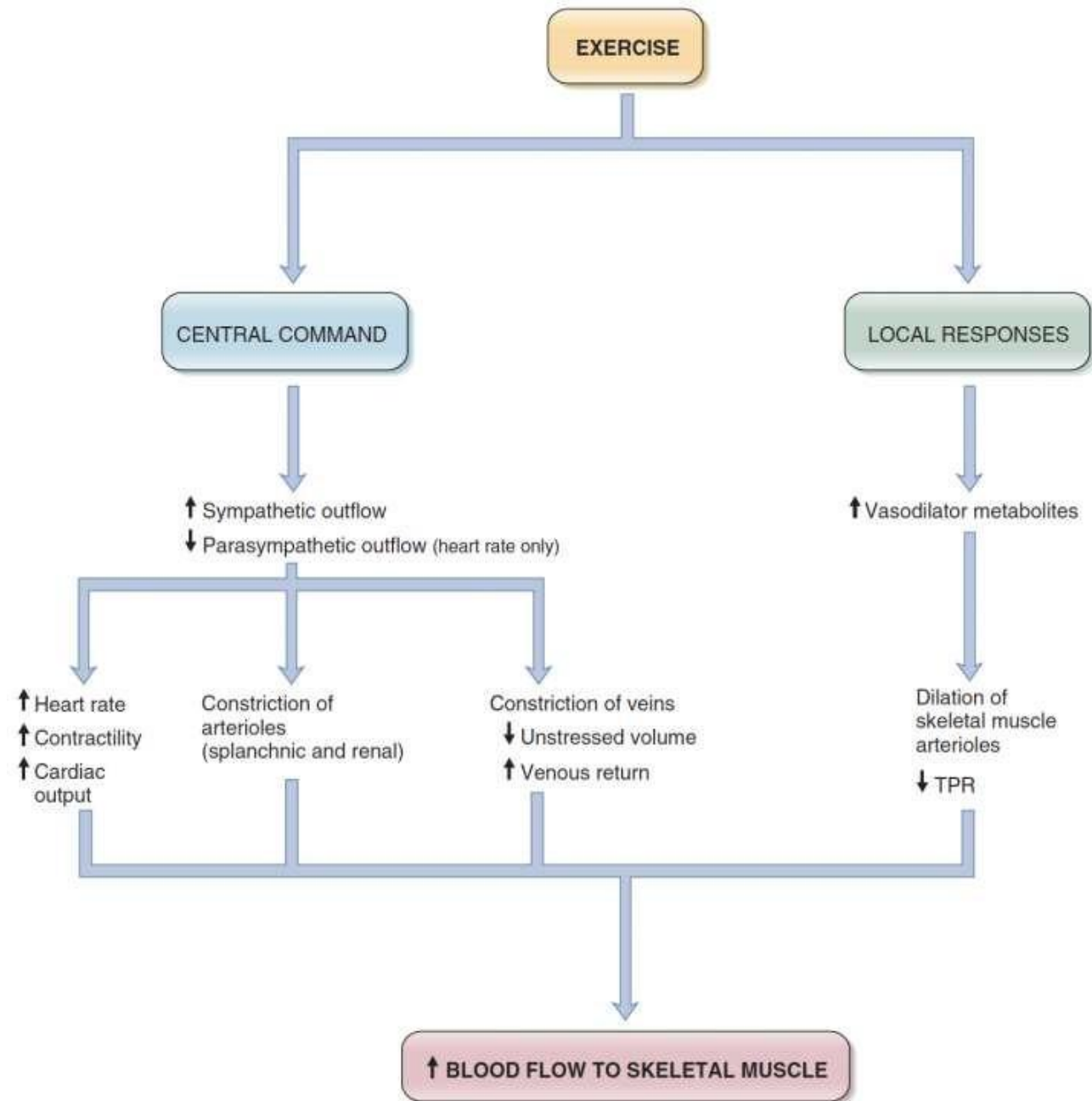
- At **rest**, blood flow to skeletal muscle is regulated primarily by its sympathetic innervation.
Vascular smooth muscle in the arterioles of skeletal muscle is densely innervated by sympathetic nerve fibers that are vasoconstricting ($\alpha 1$ receptors).
- There are also $\beta 2$ receptors on the vascular smooth muscle of skeletal muscle that are activated by epinephrine and cause vasodilation.
- Thus, activation of $\alpha 1$ receptors causes vasoconstriction, increased resistance, and decreased blood flow.
- Activation of $\beta 2$ receptors causes vasodilation, decreased resistance, and increased blood flow.
- Usually, vasoconstriction predominates

Skeletal muscle circulation

- During **exercise**, blood flow to skeletal muscle is controlled primarily by local metabolites.
- Each of the phenomena of local control is exhibited: **autoregulation and active and reactive hyperemia**.
- During exercise, the demand for O₂ in skeletal muscle varies with the activity level, and, accordingly, blood flow is increased or decreased to deliver sufficient O₂ to meet the demand.
- The local vasodilator substances in skeletal muscle are lactate, adenosine, and K⁺.
- Mechanical compression of the blood vessels in skeletal muscle can also occur during exercise and cause brief periods of occlusion.
- When the period of occlusion is over, a period of reactive hyperemia will occur, which increases blood flow and O₂ delivery.

Exercise

- The CNS responses include a central command from the cerebral motor cortex, which directs changes in the autonomic nervous system.
- The local responses include effects of metabolites to increase blood flow and O₂ delivery to the exercising skeletal muscle.



Exercise

- The central command refers to a series of responses, directed by the cerebral **primary** motor cortex, which are initiated by the anticipation of exercise **in the premotor cortex**.
- The efferent limb of the reflex produces increased sympathetic outflow to the heart and blood vessels and decreased parasympathetic outflow to the heart.

Exercise

- Cardiac output cannot increase without a concomitant increase in venous return (Frank-Starling relationship).
- In exercise, this concomitant increase in **venous return** is accomplished by two effects on the veins: The contraction of skeletal muscle around the veins has a **mechanical (squeezing) action**, and **activation of the sympathetic nervous system** produces venoconstriction.

Exercise

- Another consequence of the increased sympathetic outflow in the central command is **selective arteriolar vasoconstriction**.
- In the circulation of the skin, splanchnic regions, kidney, and inactive muscles, vasoconstriction occurs via α_1 receptors, which results in increased resistance and decreased blood flow to those organs.

Exercise

- In the exercising skeletal muscle, however, local metabolic effects override any sympathetic vasoconstricting effects, and arteriolar vasodilation occurs.
- Other locations where vasoconstriction does not occur are in the coronary circulation (where blood flow increases to meet the increased level of myocardial O₂ consumption) and the cerebral circulation.

Exercise

- In the **cutaneous circulation**, there is a **biphasic response**. Initially, vasoconstriction occurs (due to increased sympathetic outflow); later, however, as body temperature increases, there is selective inhibition of sympathetic cutaneous vasoconstriction resulting in vasodilation and dissipation of heat through the skin.
- Temperature regulation is the main systemic function of the skin circulation, so while the initial response is cutaneous vasoconstriction, after heat accumulates, vasodilation occurs to fulfill this function by dissipating the heat.

Exercise

- Local control of blood flow in the exercising skeletal muscle is orchestrated by active hyperemia.
- As the metabolic rate of the skeletal muscle increases, production of vasodilator metabolites such as lactate, K^+ , and adenosine also increases.
- These metabolites act directly on the arterioles of the exercising muscle to produce local vasodilation.
- Vasodilation of the arterioles results in increased blood flow to meet the increased metabolic demand of the muscle.
- This **vasodilation in the exercising muscle also produces an overall decrease in TPR.**

❖ Question from the doctor:

Which activity will cause a greater decrease in TPR, playing table tennis or swimming?

- Swimming involves a **larger mass** of skeletal muscles, leading to more vasodilation and a greater decrease in total peripheral resistance (TPR).

What about the effect of exercise on blood pressure?

- When the body exercises, sympathetic activity increases the heart rate, contractility and venous return. As a result, **systolic blood pressure rises** because its main controlling factor—stroke volume—also increases.
- In contrast, **diastolic blood pressure decreases**, since its main controlling factor is TPR, which is reduced during exercise.

Skin circulation

- The skin has blood vessels with dense sympathetic innervation, which controls its blood flow.
- The principal function of the sympathetic innervation is to alter blood flow to the skin for **regulation of body temperature.**
- **Local vasodilator metabolites have little effect on cutaneous blood flow.**

Temperature regulation

- Humans maintain a normal body temperature at a set point of 37°C (98.6°F).
- Because environmental temperatures vary greatly, the body has mechanisms, coordinated in the anterior hypothalamus, for both heat generation and heat loss to keep body temperature constant.
- When the environmental temperature decreases, the body generates and conserves heat.
- When the environmental temperature increases, the body reduces heat production and dissipates heat.

This slide was originally later in the file and was translocated to this position.

Regulation of body temperature

- The temperature-regulating center is located in the anterior hypothalamus.
- This center receives information about environmental temperature from thermoreceptors in the skin and about core temperature from thermoreceptors in the anterior hypothalamus itself.
- The anterior hypothalamus then orchestrates the appropriate responses, which may involve heat generating or heat-dissipating mechanisms.
- If core temperature is below the set-point temperature, then heat-generating and heat-retaining mechanisms are activated.
- If core temperature is above the set-point temperature, then heat-dissipating mechanisms are activated.
- These mechanisms include vasodilation of blood vessels of the skin (decreased sympathetic tone) and increased activity of sympathetic cholinergic fibers to sweat glands.

Mechanisms for generating heat

- When environmental temperature is less than body temperature, mechanisms are activated that increase heat production and reduce heat loss.
- These mechanisms include **stimulation of thyroid hormone** production, **activation of the sympathetic nervous system, and shivering.**
- **Behavioral** components also may contribute by reducing the exposure of skin to the cold (e.g., wrapping arms around oneself, adding more clothing).
- All changes in environmental temperature will be detected by higher cortical centers, making the individuals consciously aware that the environment is cold. As a result, heat-rising mechanisms may include behavioral changes, such as wearing more clothes, drinking something hot, etc

Mechanisms for generating heat

- Thyroid hormones are thermogenic: Their actions on target tissues result in heat production.
- Therefore, it is logical that exposure to cold temperatures activates thyroid hormones.

Mechanisms for generating heat

- Because **thyroid hormones are thermogenic**, it follows that an excess or deficit of thyroid hormones would cause disturbances in the regulation of body temperature.
- In hyperthyroidism, metabolic rate increases, O₂ consumption increases, and heat production increases, **and heat intolerance**.
- In hypothyroidism, there is a decreased metabolic rate, decreased O₂ consumption, decreased heat production, and extreme sensitivity to cold (**cold intolerance**).

Mechanisms for generating heat

- Cold environmental temperatures activate the **sympathetic nervous system**.
- One consequence of this activation is **stimulation of β receptors in brown fat**, which increases metabolic rate and heat production.
- This action of the sympathetic nervous system is synergistic with the actions of thyroid hormones:
- For thyroid hormones to produce maximal thermogenesis, the sympathetic nervous system must be simultaneously activated by cold temperatures.
- A second consequence of activation of the sympathetic nervous system is
- stimulation of α_1 receptors in vascular smooth muscle of skin blood vessels, producing vasoconstriction.
- Vasoconstriction **reduces blood flow to the surface of the skin and, consequently, reduces heat loss.**

Mechanisms for generating heat

- **Shivering**, which involves rhythmic contraction of skeletal muscle, is the most potent mechanism for increasing heat production in the body.
- Cold environmental temperatures activate centers in the posterior hypothalamus, which then activate the **α and γ motoneurons innervating skeletal muscle**.
- The skeletal muscle contracts rhythmically, generating heat and raising body temperature.

Mechanisms for dissipating heat

- When the environmental temperature increases, mechanisms are activated that result in increased heat loss from the body.
- Since heat is a normal byproduct of metabolism, the body must dissipate this heat just to maintain body temperature at the set point.
- Mechanisms for dissipating heat are coordinated in the anterior hypothalamus.
- Increased body temperature **decreases sympathetic activity** in skin blood vessels.

Mechanisms for dissipating heat

- This decrease in sympathetic tone results in increased blood flow through skin arterioles and greater arteriovenous shunting of blood to venous plexuses near the surface of skin.
- In effect, warm blood from the body core is shunted to the body surface, [increasing heat loss to the external environment](#).
- Shunting of blood to the surface is evidenced by redness and warmth of the skin.
- There also is increased activity of the **sympathetic cholinergic fibers innervating thermoregulatory sweat glands** to produce increased sweating (cooling).
- The **behavioral** components to dissipate heat include increasing the exposure of skin to the air (e.g., removing clothing, fanning).

Fever

- Fever is an abnormal elevation of body temperature.
- **Pyrogens** produce fever by **increasing the hypothalamic set-point temperature**. The result of such a change in set point is that a normal core temperature is “seen” by the hypothalamic center as too low relative to the new set point. The anterior hypothalamus then activates heat-generating mechanisms (e.g., shivering) to raise body temperature to the new set point.
- At the cellular level, the mechanism of pyrogen action is increased production of interleukin-1 (IL-1) in phagocytic cells. IL-1 then acts on the anterior hypothalamus to increase local production of **prostaglandins**, which increase the set-point temperature.

Fever

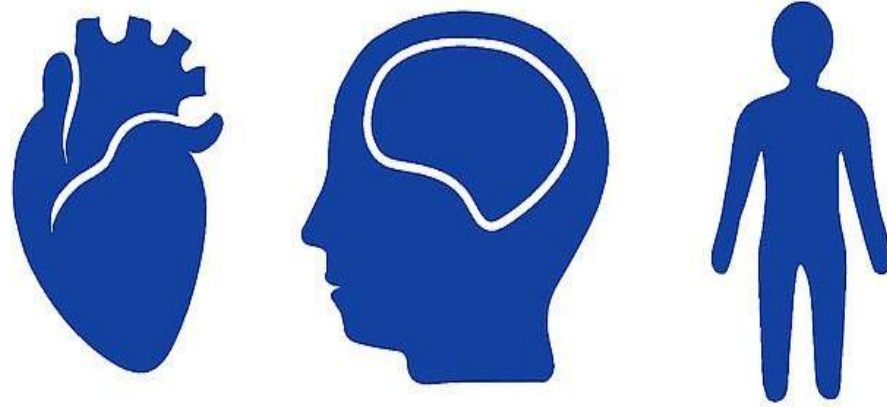
- When fever is treated, the temperature sensors in the anterior hypothalamus now “see” body temperature as too high relative to the set-point temperature and set in motion the mechanisms for dissipating heat including vasodilation and sweating.
- To treat fever and restore the normal set-point temperature, a drug that decreases prostaglandins, such as COX inhibitors, is given.

Disturbances of temperature regulation

- Heat exhaustion can occur as a consequence of the body's responses to elevated environmental temperature.
- Normally, the response to increased temperature includes vasodilation and sweating in order to dissipate heat.
- However, if the sweating is excessive, it can result in decreased ECF volume, decreased blood volume, decreased arterial pressure, and fainting.
- Heat stroke occurs when body temperature increases to the point of tissue damage.
- If the normal response to elevated environmental temperature is impaired (e.g., if sweating does not occur), then heat cannot be appropriately dissipated, and core temperature increases to dangerous levels.

Malignant hyperthermia

- Malignant hyperthermia (a disease with genetic predisposition) is characterized by a massive increase in metabolic rate, increased O₂ consumption, and increased heat production in skeletal muscle.
- The heat-dissipating mechanisms are unable to keep pace with the excessive heat production, and if the hyperthermia is not treated, body temperature may increase to dangerously high, or even fatal, levels.
- In susceptible individuals, malignant hyperthermia can be caused by inhalation anesthetics or muscle relaxants.
- The anesthesiologist should immediately act, and the patient should be labeled to indicate the condition for later procedures, and the family members should be alerted because they may have the same risk.



**PHYSIOLOGY
QUIZ
LECTURE 7**

رسالة من الفريق العلمي

قال الله تعالى :

إِنَّ الْمُتَّقِينَ فِي جَنَّاتٍ وَعُيُونٍ • ادْخُلُوهَا
بِسَلَامٍ آمِنِينَ • وَنَزَعْنَا مَا فِي صُدُورِهِمْ مِنْ
غِلٍّ إِخْوَانًا عَلَى سُرُرٍ مُتَقَابِلِينَ • لَا يَمَسُّهُمْ
فِيهَا نَصَبٌ وَمَا هُمْ مِنْهَا بِمُخْرَجِينَ

(الحجر : ٤٥ - ٤٨)

أي إن الذين اتقوا الله بامتنال ما أمر واجتناب ما نهى في بساتين وأنهار جارية يقال لهم:
ادخلوا هذه الجنات سالمين من كل سوء آمنين من كل عذاب. ونزعنا ما في قلوبهم من حقد
 وعداوة, يعيشون في الجنة إخواناً متحابين, يجلسون على أسرة عظيمة, تتقابل وجوههم
تواصلاً وتحابياً, لا يصيبهم فيها تعب ولا إعياء, وهم باقون فيها أبداً.

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Corrections from previous versions:

Versions	Slide # and Place of Error	Before Correction	After Correction
V0 → V1			
V1 → V2			