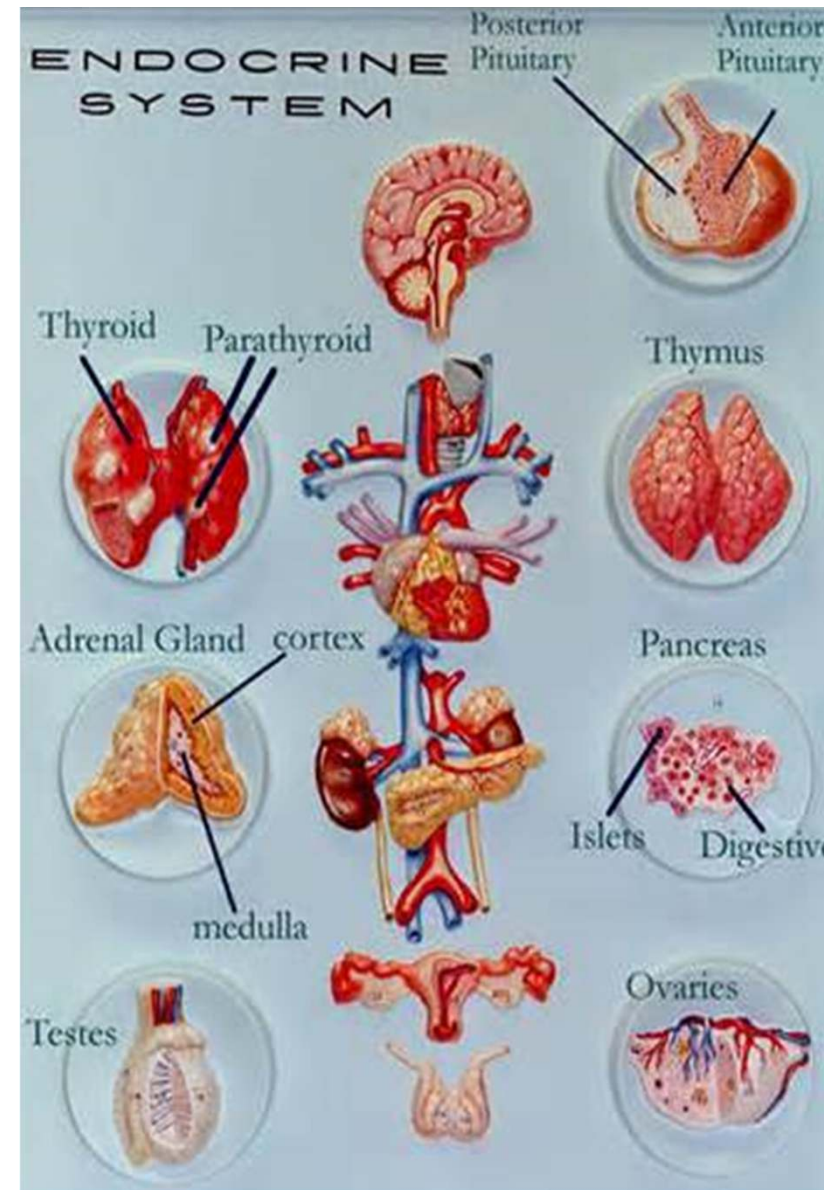
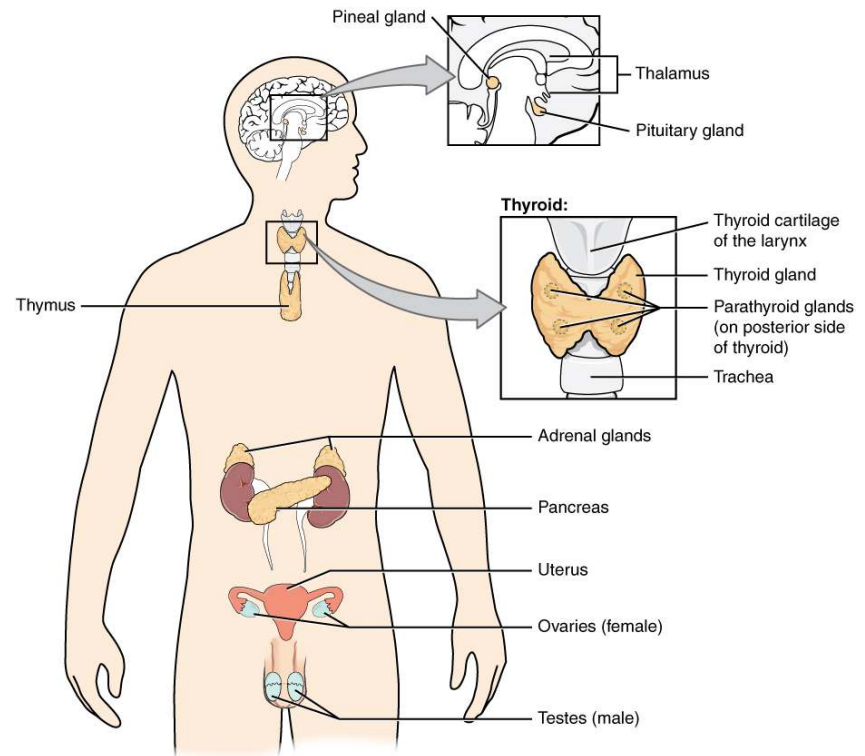
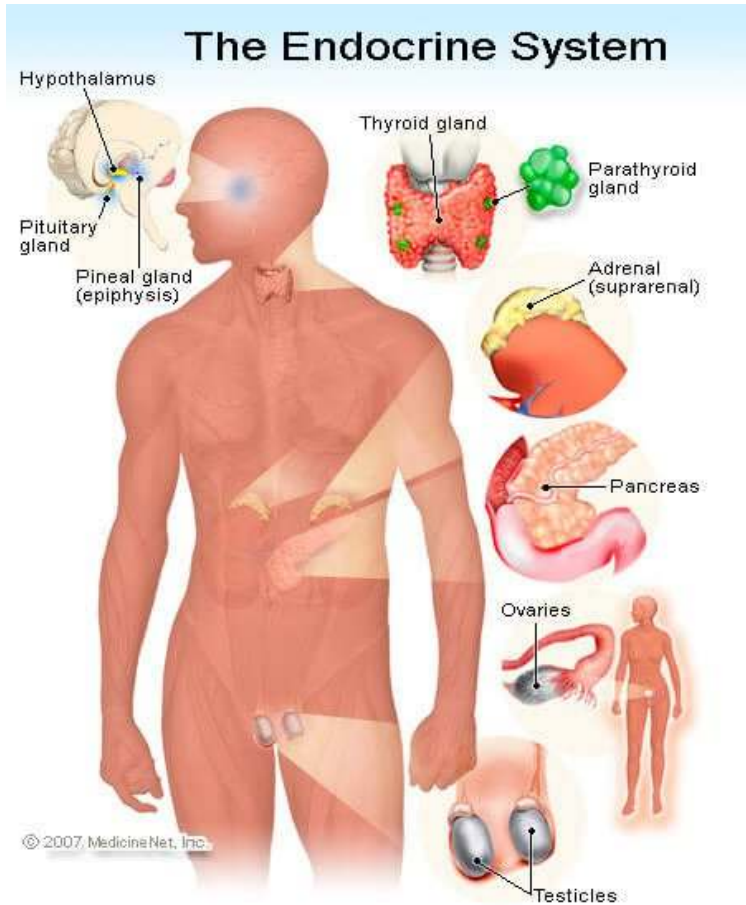


# The endocrine system

D.Hammoudi. MD



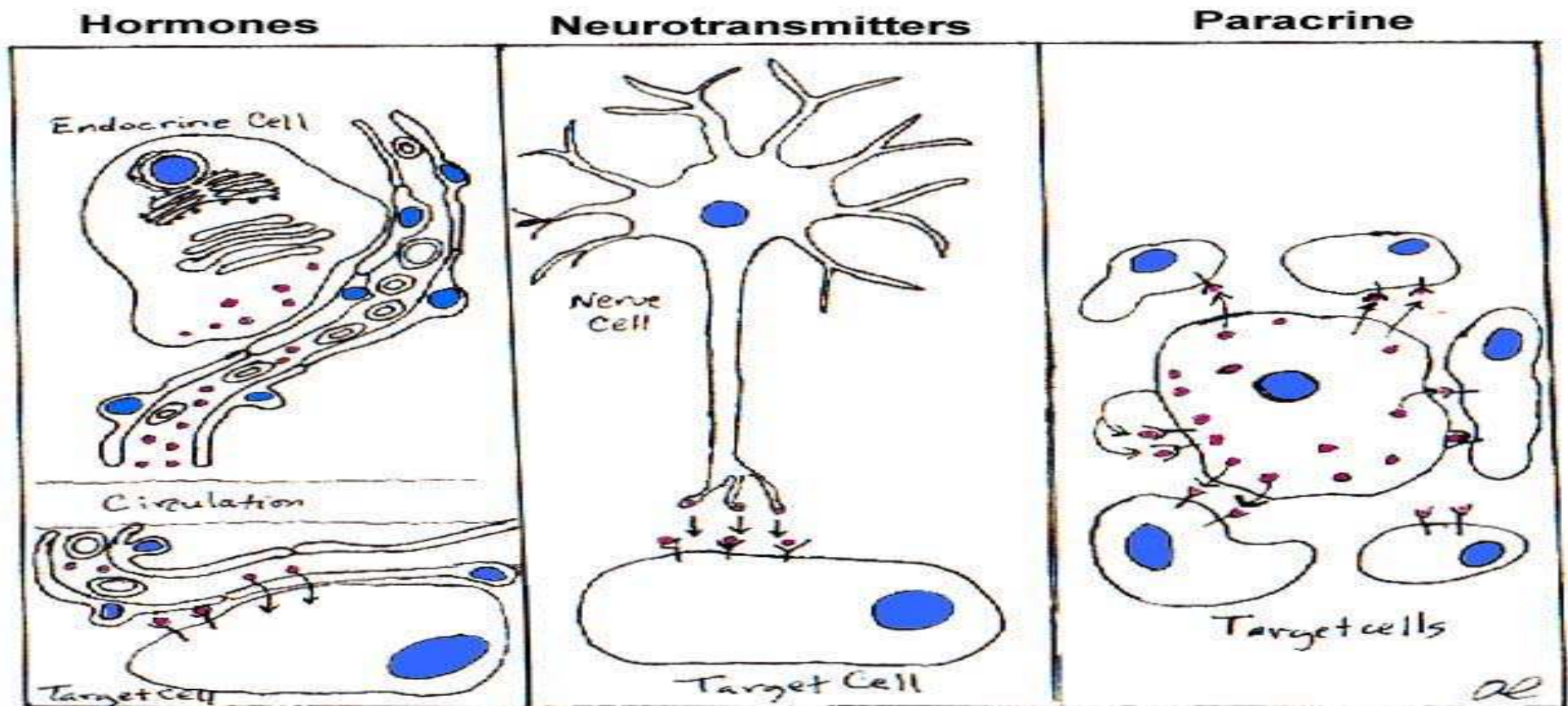
# GENERALITY



# Major Mechanisms for Signaling

- **Endocrine hormones** - small molecules released into the circulation to effect target cells at distant sites from the original release point.
- **Paracrine hormones** - small molecules released in a local area which has an effect only on cells within that local area of the body
- **Neurotransmission** - synaptic transmission

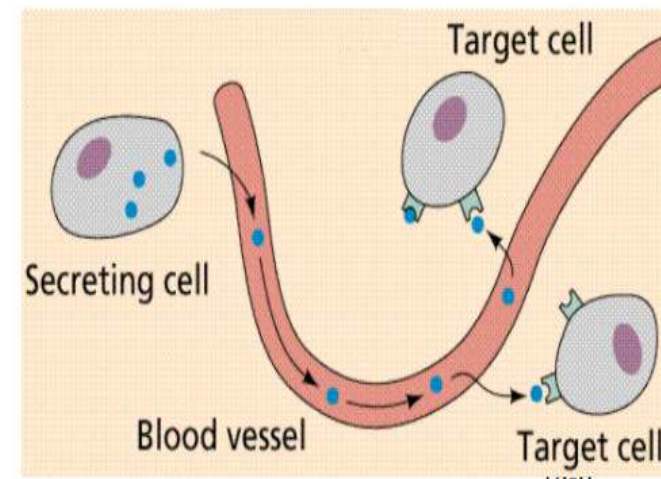
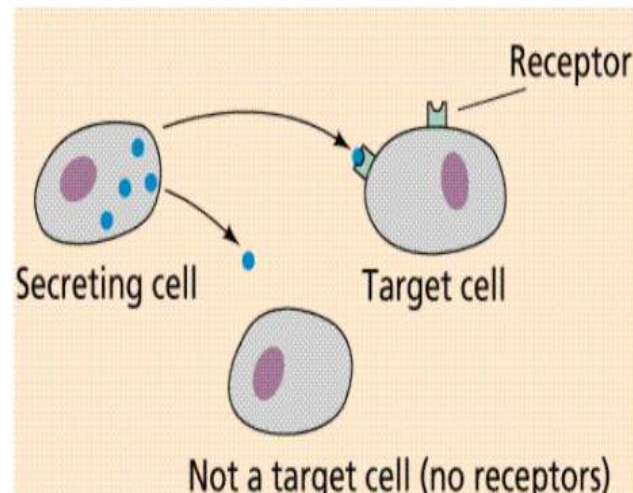
# Comparison of the three



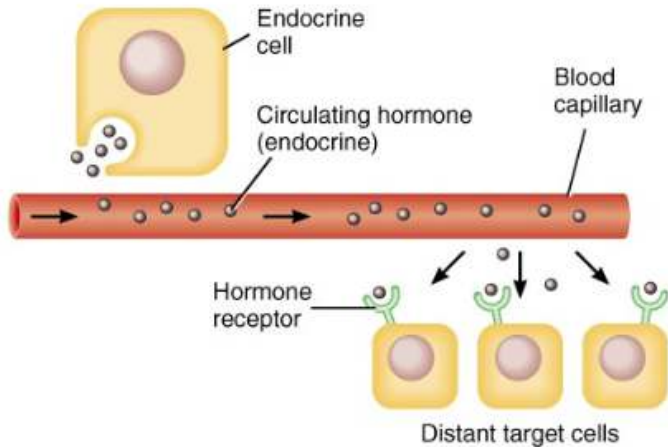


# Endocrine Glands Defined

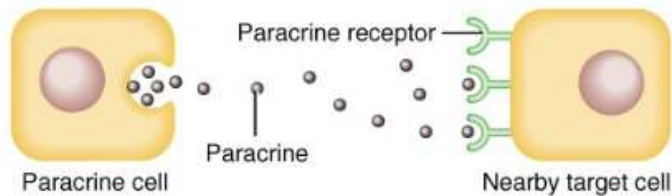
- Exocrine glands
  - secrete products into ducts which empty into body cavities or body surface
  - sweat, oil, mucous, & digestive glands
- Endocrine glands
  - secrete products (hormones) into bloodstream
  - pituitary, thyroid, parathyroid, adrenal, pineal
  - other organs secrete hormones as a 2nd function
- hypothalamus, thymus, pancreas, ovaries, testes, kidneys, stomach, liver, small intestine, skin, heart & placenta
- The endocrine system is a collection of glands that secrete chemical messages we call hormones.
- These signals are passed through the blood to arrive at a target organ, which has cells possessing the appropriate receptor.



## Circulating & Local Hormones



(a) Circulating hormones (endocrines)



(b) Local hormones (paracrines and autocrines)

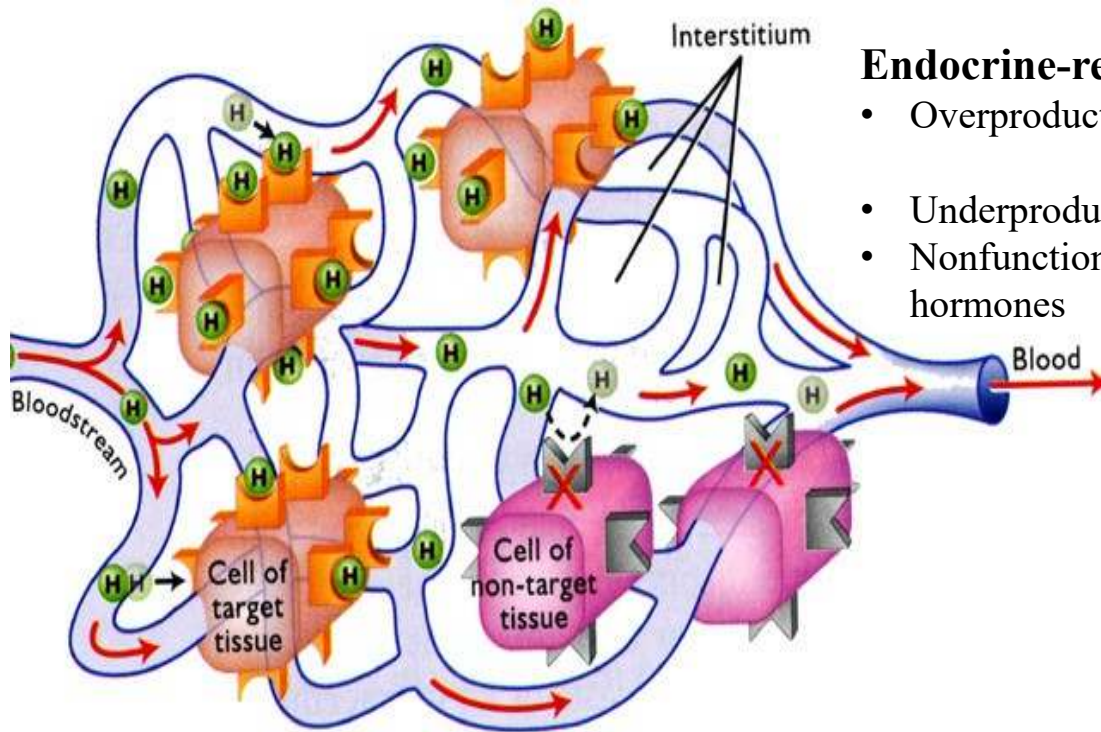
- **Circulating hormones**
  - act on distant targets
  - travel in blood
- **Local hormones**
  - paracrines act on neighboring cells
  - autocrines act on same cell that secreted them

## General Mechanisms of Hormone Action

- Hormone binds to cell surface or receptor inside target cell
- Cell may then
  - **synthesize new molecules**
  - **change permeability of membrane**
  - **alter rates of reactions**
- Each target cell responds to hormone differently
  - **liver cells---insulin stimulates glycogen synthesis**
  - **adipose---insulin stimulates triglyceride synthesis**

# Control of Hormone Secretion

- Regulated by signals from nervous system, chemical changes in the blood or by other hormones
- **Negative feedback control (most common)**
  - decrease/increase in blood level is reversed
- **Positive feedback control**
  - the change produced by the hormone causes more hormone to be released
- Disorders involve either hyposecretion or hypersecretion of a hormone

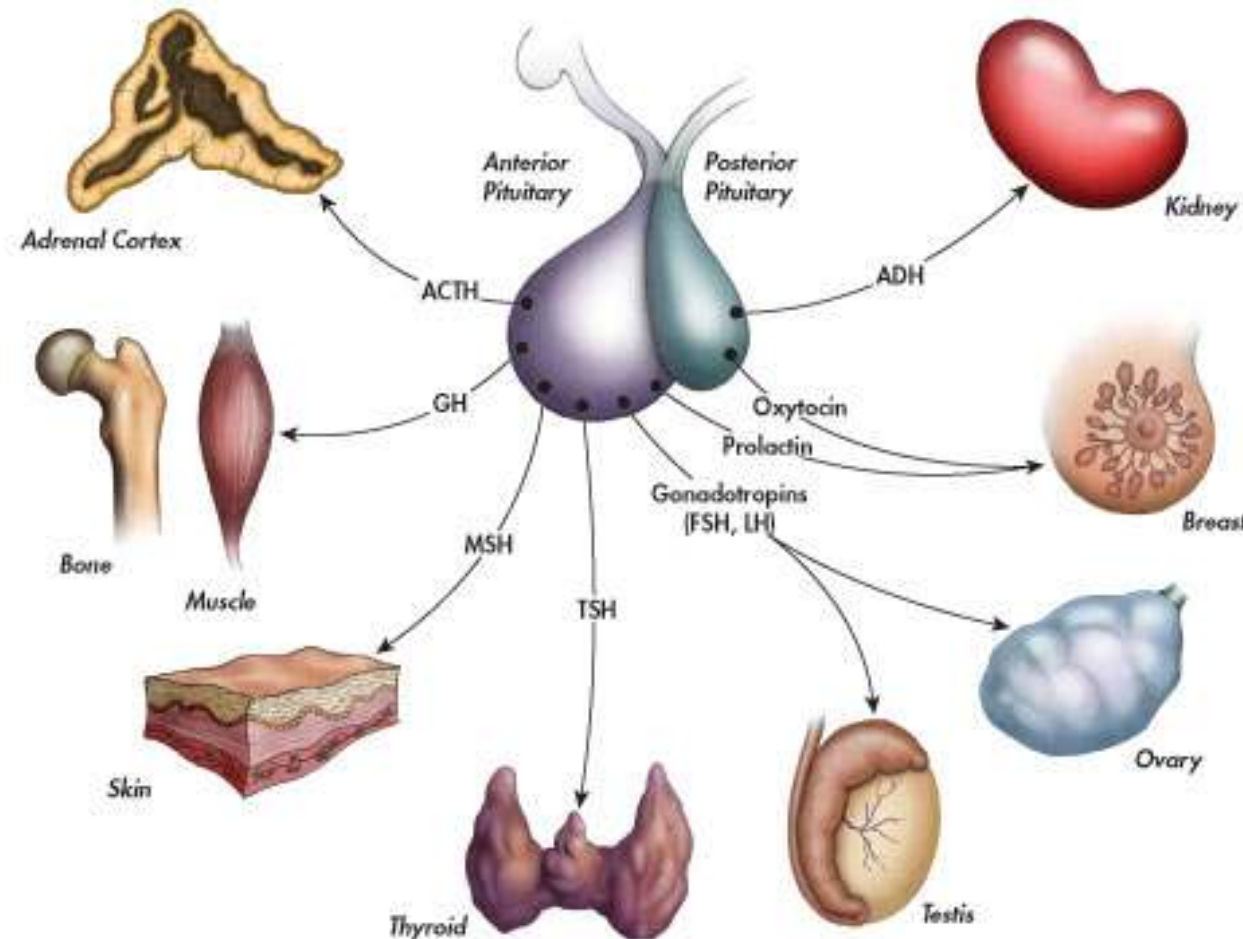


## Endocrine-related Problems

- Overproduction of a hormone
- Underproduction of a hormone
- Nonfunctional receptors that cause target cells to become insensitive to hormones

# FEEDBACK SYSTEMS

- CORTEX, SUBCORTEX? →
- HYPOTHALAMUS →
- ANTERIOR PITUITARY →
- ENDOCRINE GLAND →
- END ORGAN →
- HYPOTHALAMUS →





# Hormones

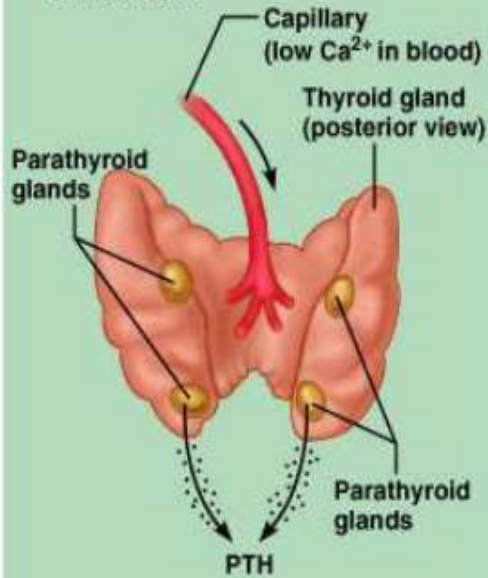
## 5 major classes

1. *Amino acid derivatives* → dopamine, catecholamine, and thyroid hormone;
2. *Small neuropeptides* → gonadotropin-releasing hormone (GnRH), thyrotropin-releasing hormone (TRH), somatostatin, and vasopressin;
3. *Large proteins* → insulin, luteinizing hormone (LH), and PTH produced by classic endocrine glands;
4. *Steroid hormones* such as cortisol and estrogen;
5. *Vitamin derivatives* such as retinoids (vitamin A) and vitamin D.

As a rule – protein based hormones act on the 'cell surface receptors' and steroid based hormones act on 'intracellular nuclear proteins'



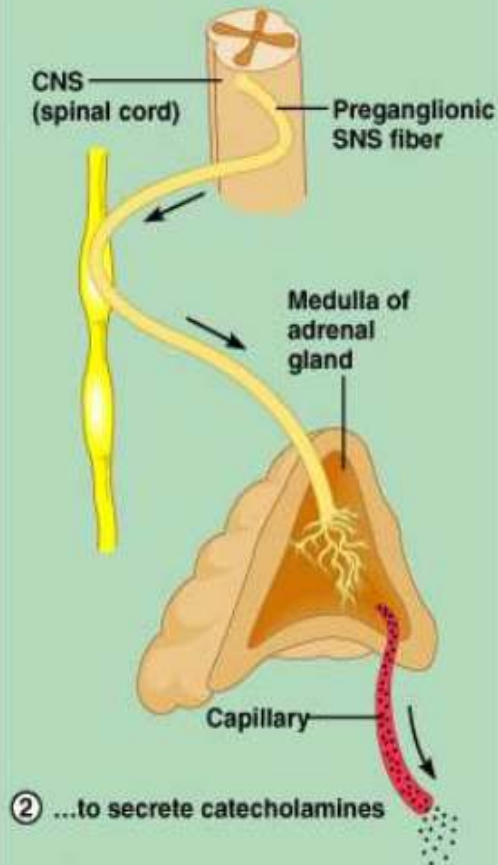
① Capillary blood contains low concentration of  $\text{Ca}^{2+}$ , which stimulates...



② ...secretion of parathyroid hormone (PTH)

(a) Humoral

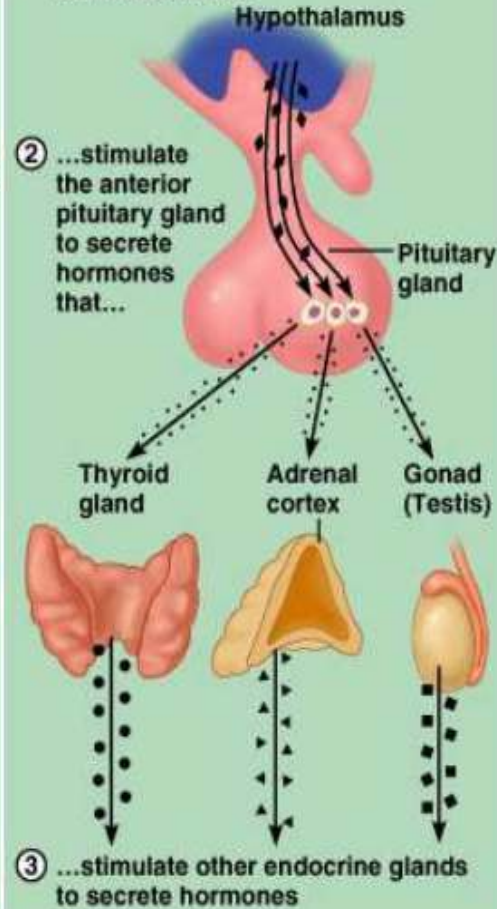
① Preganglionic SNS fiber stimulates adrenal medulla cells...



② ...to secrete catecholamines

(b) Neural

① The hypothalamus secretes hormones that...



③ ...stimulate other endocrine glands to secrete hormones

(c) Hormonal

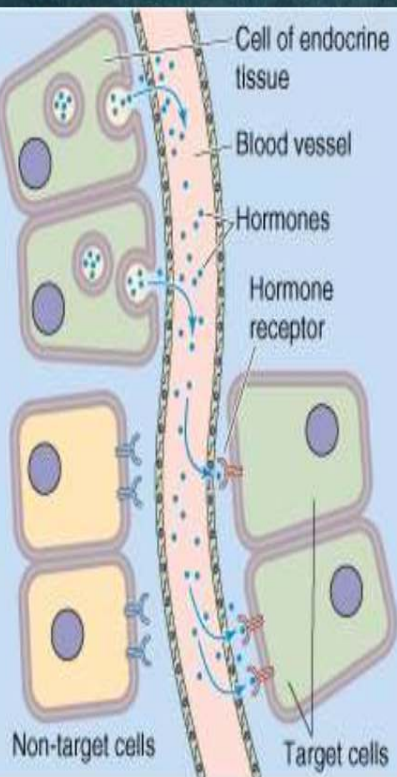
(a) **Humoral:** in response to changing levels of ions or nutrients in the blood

(b) **Neural:** stimulation by nerves

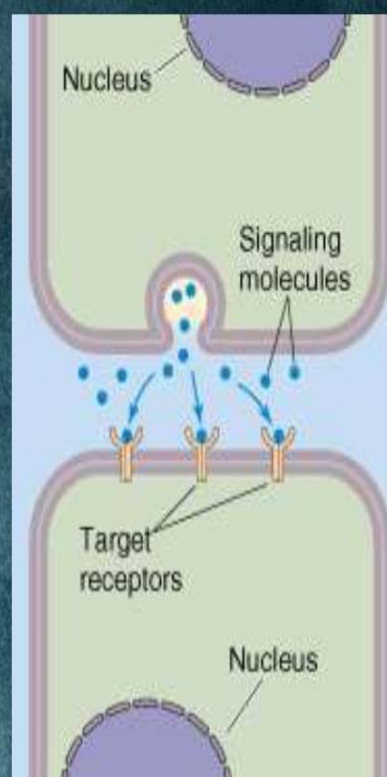
(c) **Hormonal:** stimulation received from other hormones

# Sites of hormone action

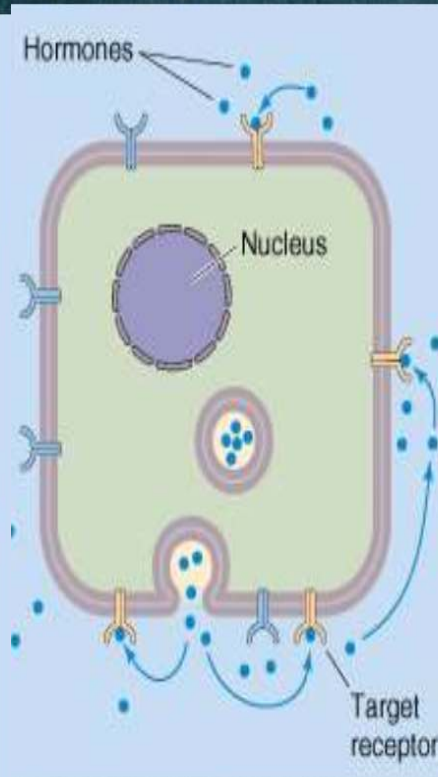
## Endocrine glands



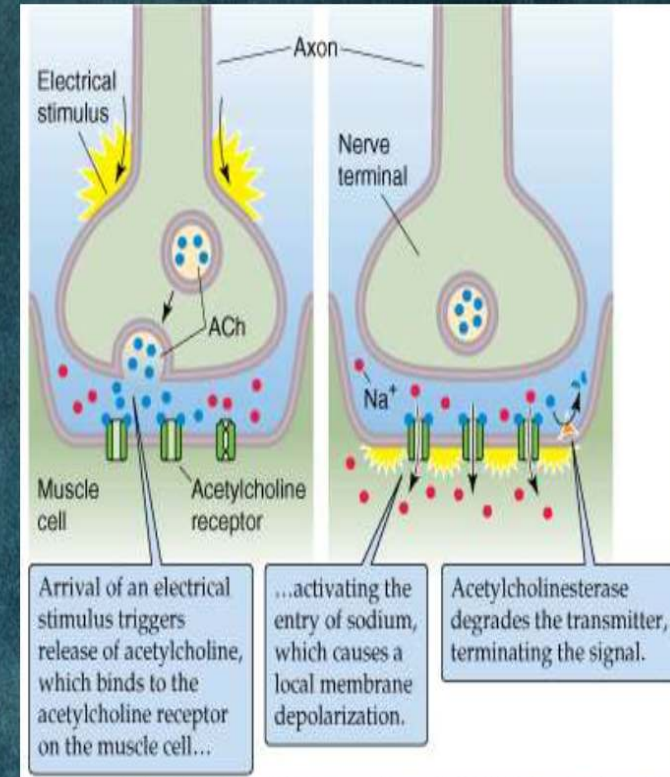
## Exocrine glands



## Autocrine glands



## Neurotransmission





**TABLE 50.1:** Mechanism of action of hormones (for expansions or abbreviations, see Appendix No.I)

Group	Mechanism of action	Examples of hormone
I A	Hormones bind with cell surface receptors with cAMP as the second messenger	ACTH, ADH, FSH HCG, LH, TSH MSH, PTH, CRH Glucagon, Calcitonin Catecholamines Retinoic acid
I B	Hormones having cell surface receptors; cGMP as second messenger	ANF (atrial natriuretic factor), NO (nitric oxide)
I C	Hormones having cell surface receptors; second messenger is calcium or phosphatidyl inositol (PIP2)	TRH, GnRH catecholamines Acetylcholine CCK, Gastrin Vasopressin Oxytocin, PDGF
I D	Hormones having cell surface receptors and mediated through tyrosine kinase	Insulin Somatomedin EGF, FGF PDGF, CGSF NGF, IGF
I E	Hormones having cell surface receptors, but intracellular messenger is a kinase or utilize phosphatase cascade	IL, GH, PRL, TNF, Adiponectin, Leptin, Resistin, Erythropoietin
II	Hormones that bind to intracellular receptors	Glucocorticoids Mineralocorticoids Estrogens, Progesterone Androgens Calcitriol, Thyroxine

There are two types of cells in signal transduction

- the **sender cell where the signal originates**
- the **target cell that receives the signal.**
- **The signal alters** or modulates the activity/function of the cell.
- **Autocrine signaling occurs when same cell acts as** sender and recipient, e.g. growth, differentiation, immune and inflammatory response.
- **Paracrine signaling is** effected by local mediators which have their effect near the site of secretion without entering the circulation.
- The effect is rapid and transient.
- **Juxtacrine signaling occurs when the two type of cells are adjacent to each other so that contact is established through gap junctions or through protein molecules on the surface of the two cells.**
- **Endocrine signaling is between cells which are located at a distance from each other and the signal may be hormones or chemical messengers secreted into circulation.**  
Once they reach the target cell, they bind to specific target cell receptors with high affinity.

# List three kinds of interaction of different hormones acting on the same target cell.

- **Permissiveness** – one hormone cannot exert its full effects without another hormone being present (ex. Reproductive system hormones regulate the development of the reproductive system. However thyroid hormone is also necessary for normal timely development of reproductive structures. Lack of thyroid hormone delays reproductive development.
- **Synergism** – occurs when more than one hormone produces the same effect at the target cell and their combined effects are amplified. (ex. both glucagon (pancreas) and epinephrine causes the liver to release glucose into the blood. When they act together, the amount of glucose released is about 150% of what is released when each hormone acts alone
- **Antagonism** – occurs when one hormones opposes the action of another hormone. (ex. insulin which lowers blood glucose levels, is antagonized by glucagon, which raises blood glucose levels.
- **Antagonists may:**
  - ☐ compete for the same receptor
  - ☐ Act through different metabolic pathways
  - ☐ Cause down-regulation of the receptors for the antagonistic hormone.

**Hormones stimulate adenyl cyclase:** ACTH, ADH, Calcitonin, CRH, FSH, Glucagon, epinephrine, hCG, LH, LPH, MSH, PTH and TSH.

**Hormones inhibit adenyl cyclase:** Acetylcholine, angiotensin II and somatostatin.

## Signaling pathways of endocrine hormones

<b>cAMP</b>	<b>FSH, LH, ACTH, TSH, CRH, hCG, ADH</b> (V <sub>2</sub> -receptor), <b>MSH, PTH</b> , calcitonin, GHRH, glucagon, histamine (H <sub>2</sub> -receptor)	<b>FLAT ChAMP</b>
<b>cGMP</b>	<b>BNP, ANP, EDRF (NO)</b>	<b>BAD GraMP<sub>a</sub></b> Think vasodilators
<b>IP<sub>3</sub></b>	<b>GnRH, Oxytocin, ADH (V<sub>1</sub>-receptor), TRH,</b> <b>Histamine (H<sub>1</sub>-receptor), Angiotensin II,</b> <b>Gastrin</b>	<b>GOAT HAG</b>
<b>Intracellular receptor</b>	<b>Progesterone, Estrogen, Testosterone, Cortisol,</b> <b>Aldosterone, T<sub>3</sub>/T<sub>4</sub>, Vitamin D</b>	<b>PET CAT on TV</b>
<b>Receptor tyrosine kinase</b>	Insulin, <b>IGF-1, FGF, PDGF, EGF</b>	MAP kinase pathway Think <b>Growth Factors</b>
<b>Nonreceptor tyrosine kinase</b>	<b>Prolactin, Immunomodulators</b> (eg, cytokines IL-2, IL-6, IFN), <b>GH, G-CSF, Erythropoietin,</b> <b>Thrombopoietin</b>	JAK/STAT pathway Think acidophils and cytokines <b>PIGGLET</b>



## Signaling Pathways of Endocrine Hormones

### General mechanism

- releasing hormone (IP<sub>3</sub>) → pituitary hormone (cAMP) → systemic hormone (steroid)
  - GnRH → FSH/LH → estrogen/testosterone/progesterone
  - TRH → TSH → T<sub>3</sub>/T<sub>4</sub>
- vasoactive hormones
  - cGMP
- growth factors
  - tyrosine kinase
- growth hormone, cytokines, hormones
  - receptor tyrosine kinase

IP <sub>3</sub>	cAMP	cGMP	Tyrosine kinase - intrinsic	Tyrosine kinase - receptor associated	Steroid
GnRH	FSH	ANP	Insulin	Prolactin	Glucocorticoid
Gastrin	LH	NO (EDRF)	IGF-1	Cytokines (IL-2,6,8)	Estrogen
Oxytocin	ACTH		FGF	GH	Progesterone
TRH	TSH		PDGF		Testosterone
ADH (V <sub>1</sub> )	CRH				Aldosterone
Histamine (H <sub>1</sub> )	hCG				Vitamin D
Angiotensin II	PTH				T <sub>3</sub> /T <sub>4</sub>
	Calcitonin				Cortisol
	Glucagon				
	GHRH (can act via IP <sub>3</sub> as well)				

# What are endocrine systems for?

## Endocrine Functions

- **Maintain Internal Homeostasis**
- **Support Cell Growth**
- **Coordinate Development**
- **Coordinate Reproduction**
- **Facilitate Responses to External Stimuli**

# What are the elements of an endocrine system?

- ***Sender*** = Sending Cell
- ***Signal*** = Hormone
- ***Nondestructive Medium*** = Serum & Hormone Binders
- ***Selective Receiver*** = Receptor Protein
- ***Transducer*** = Transducer Proteins & 2° Messengers
- ***Amplifier*** = Transducer/Effector Enzymes
- ***Effector*** = Effector Proteins
- ***Response*** = Cellular Response (2° Hormones)

# Functions

## Maintenance of growth & development

- – Growth hormone,
- Thyroxine,
- insulin,
- Glucocorticoid,
- Gonadal hormones

## Maintenance of internal environment

- ADH,
- Mineralocorticoids,
- PTH

## Regulation of energy balance and metabolism –

- Insulin,
- glucagon ,
- Leptin & Ghrelin

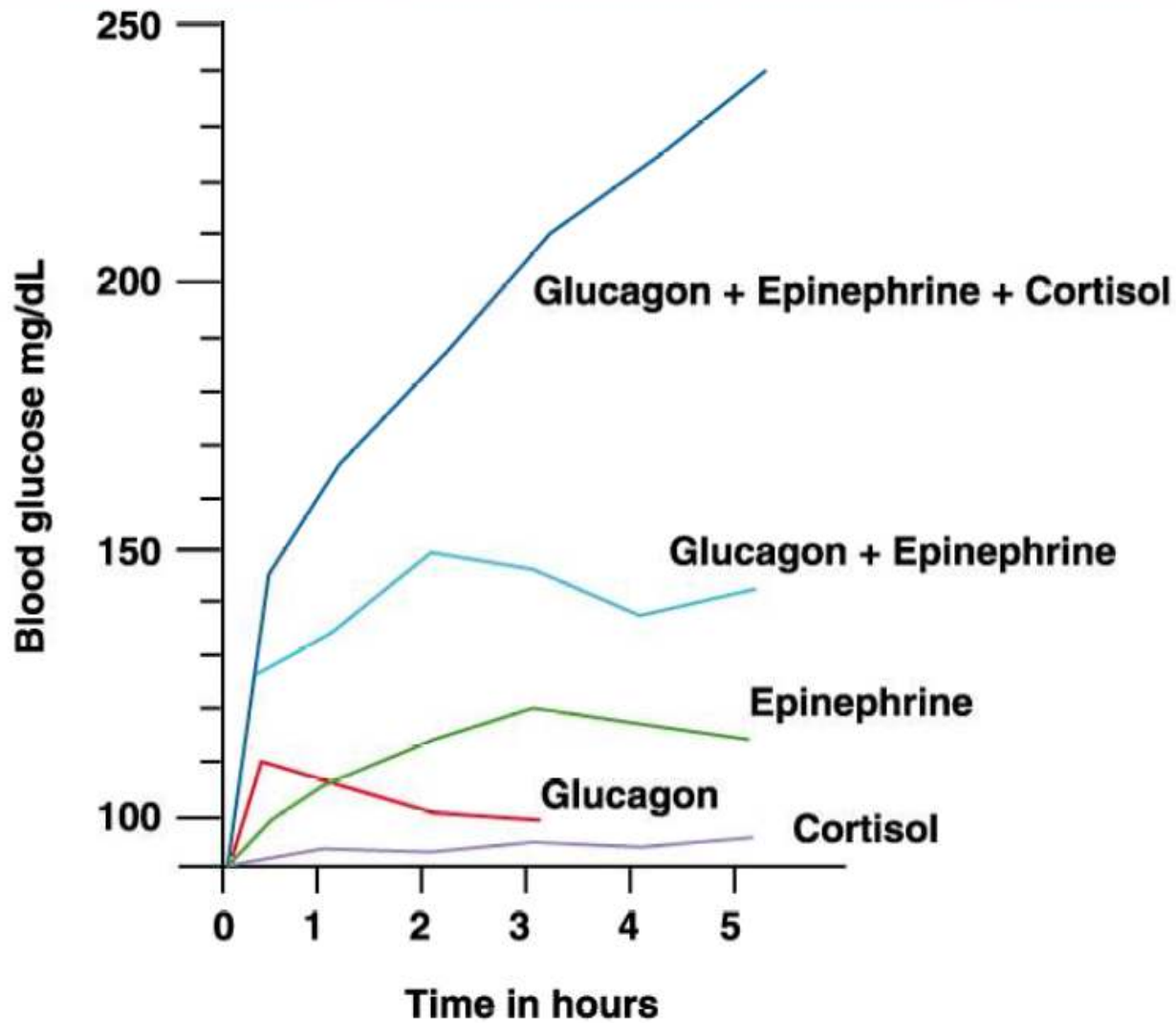
## Reproduction & species propagation – Gonadal & Pituitary hormones

When two or more hormones work together to produce particular result their effects are said to be **synergistic**.

- These effects may be additive or complementary.
- **Additive**: Same effect of the hormones on one target organ, for example, epinephrine and norepinephrine on heart rate
- **Complementary**: Work on different stages of a physiological procedure, for example, FSH (initiation) and testosterone (maintenance) on spermatogenesis



Synergistic effects



**HYPOTHALAMUS**  
 Production of ADH, oxytocin, and regulatory hormones

**PITUITARY GLAND**  
 Anterior lobe: ACTH, TSH, GH, PRL, FSH, LH, and MSH  
 Posterior lobe: Release of oxytocin and ADH

**THYROID GLAND**  
 Thyroxine (T<sub>4</sub>)  
 Triiodothyronine (T<sub>3</sub>)  
 Calcitonin (CT)

**THYMUS**  
 (Undergoes atrophy during adulthood)  
 Thymosins (Chapter 22)

**ADRENAL GLANDS**  
 Each adrenal gland is subdivided into:  
 Adrenal medulla:  
 Epinephrine (E)  
 Norepinephrine (NE)  
 Adrenal cortex:  
 Cortisol, corticosterone, aldosterone, androgens

**PINEAL GLAND**  
 Melatonin

**PARATHYROID GLANDS**  
 (on posterior surface of thyroid gland)  
 Parathyroid hormone (PTH)

**HEART**  
 Natriuretic peptides: ANP and BNP (Chapter 21)

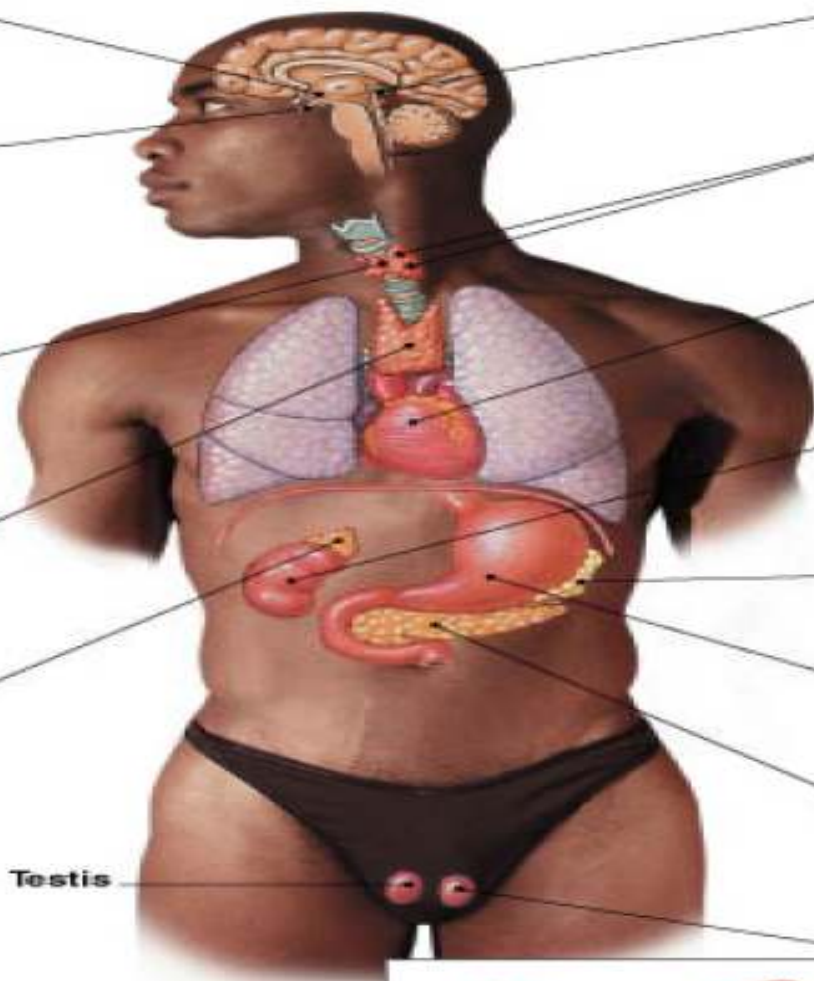
**KIDNEY**  
 Erythropoietin (EPO)  
 Calcitriol (Chapters 19 and 27)

**ADIPOSE TISSUE**  
 Leptin  
 Resistin

**DIGESTIVE TRACT**  
 Numerous hormones (detailed in Chapter 24)

**PANCREATIC ISLETS**  
 Insulin, glucagon

**GONADS**  
 Testes (male): Androgens (especially testosterone), inhibin  
 Ovaries (female): Estrogens, progestins, inhibin



**Table 37-1. Hormones and Their Sites of Production in Nonpregnant Adults**

Gland	Hormone
<b>Hormones Synthesized and Secreted by Dedicated Endocrine Glands</b>	
Pituitary gland	Growth hormone (GH) Prolactin Adrenocorticotrophic hormone (ACTH) Thyroid-stimulating hormone (TSH) Follicle-stimulating hormone (FSH) Luteinizing hormone (LH)
Thyroid gland	Tetraiodothyronine (T <sub>4</sub> ; thyroxine) Triiodothyronine (T <sub>3</sub> ) Calcitonin
Parathyroid glands	Parathyroid hormone (PTH)
Islets of Langerhans (endocrine pancreas)	Insulin Glucagon Somatostatin
Adrenal gland	Epinephrine Norepinephrine Cortisol Aldosterone Dehydroepiandrosterone sulfate (DHEAS)
Ovaries	Estradiol-17β Progesterone Inhibin
Testes	Testosterone Antimüllerian hormone (AMH) Inhibin

**Hormones Synthesized in Organs with a Primary Function Other Than Endocrine**

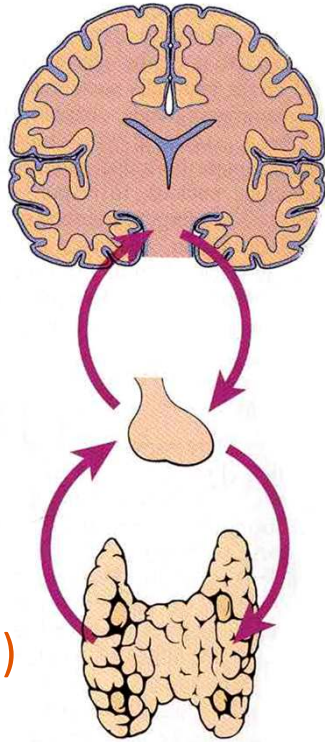
Brain (hypothalamus)	Antidiuretic hormone (ADH; vasopressin) Oxytocin Corticotropin-releasing hormone (CRH) Thyrotropin-releasing hormone (TRH) Gonadotropin-releasing hormone (GnRH) Growth hormone-releasing hormone (GHRH) Somatostatin Dopamine
Brain (pineal gland)	Melatonin
Heart	Atrial natriuretic peptide (ANP)
Kidney	Erythropoietin
Adipose tissue	Leptin Adiponectin
Stomach	Gastrin Somatostatin Ghrelin
Intestines	Secretin Cholecystokinin Glucagon-like peptide-1 (GLP-1) Glucagon-like peptide-2 (GLP-2) Glucose-dependent insulinotropic peptide (GIP; gastrin inhibitory peptide) Motilin
Liver	Insulin-like growth factor type I (IGF-I)

# Endocrine System in a Nutshell

Hypothalamus

Pituitary

Endocrine organ  
(for example, thyroid)



The hypothalamus tells the pituitary what to do

The pituitary tells the endocrine organ what to do

The endocrine organ releases hormone

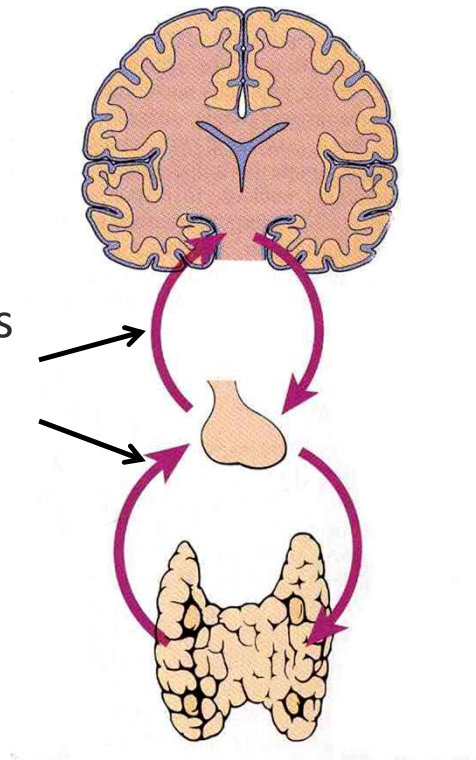
The hypothalamus is like a CEO but we don't talk about it much (not many diseases there)

The pituitary is like a COO. It basically tells everyone what to do.

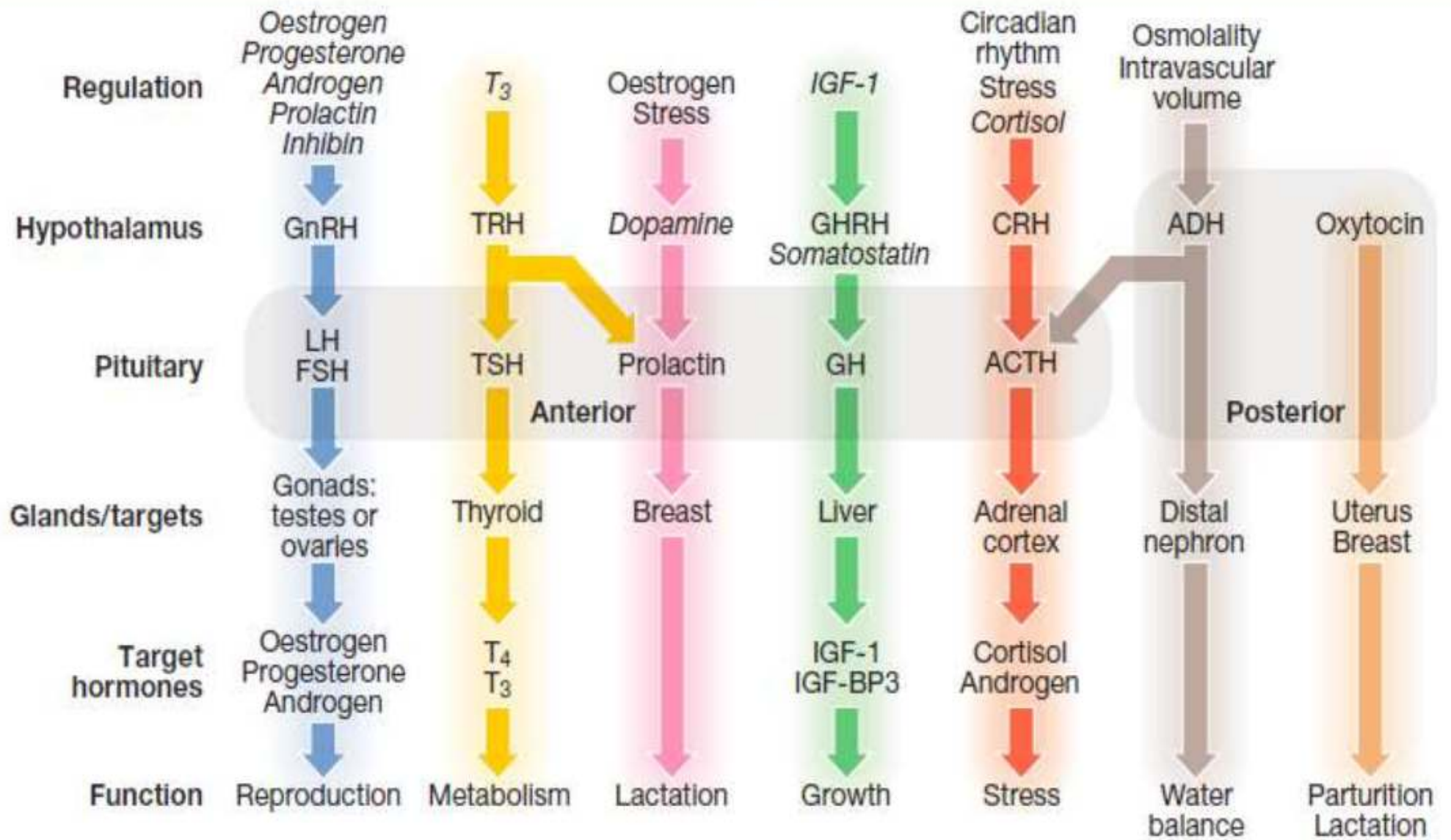
The endocrine organ is the worker drone. Poor guy.

## Endocrine System in a Nutshell

There are negative feedback loops that tell the system when to stop producing hormone.









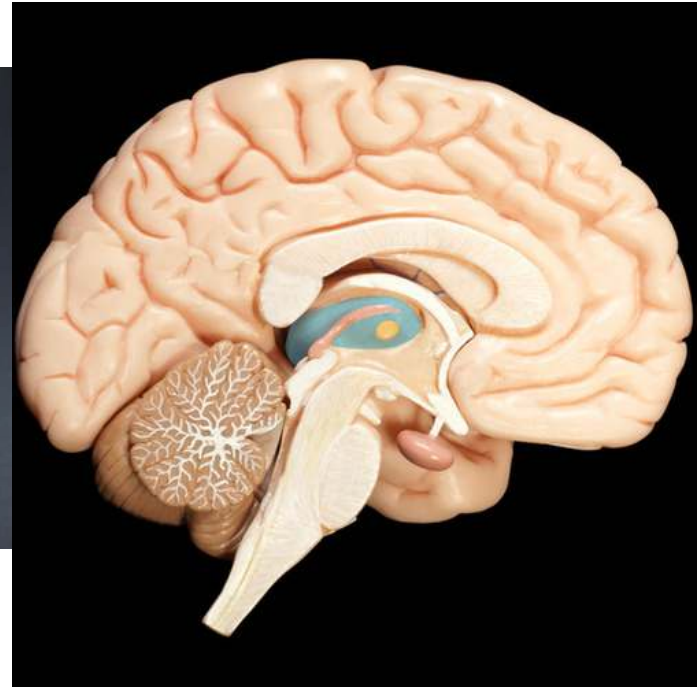
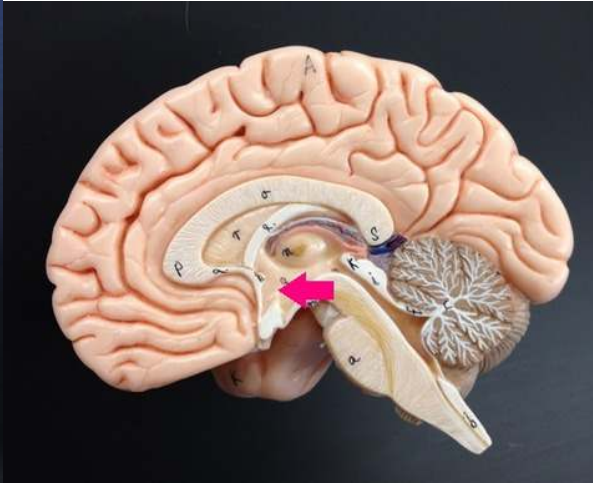
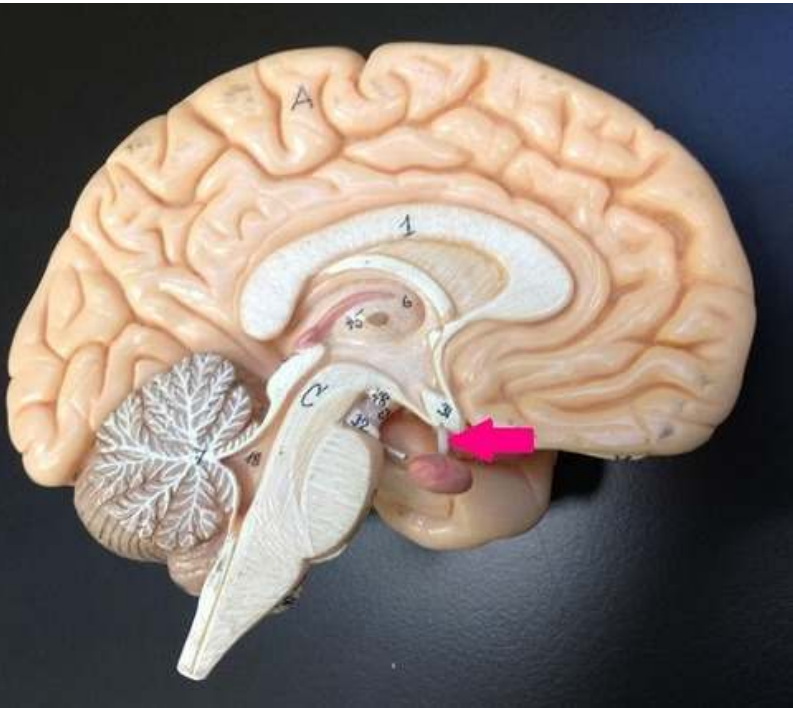
## Hormones Produced to a Significant Degree by Peripheral Conversion

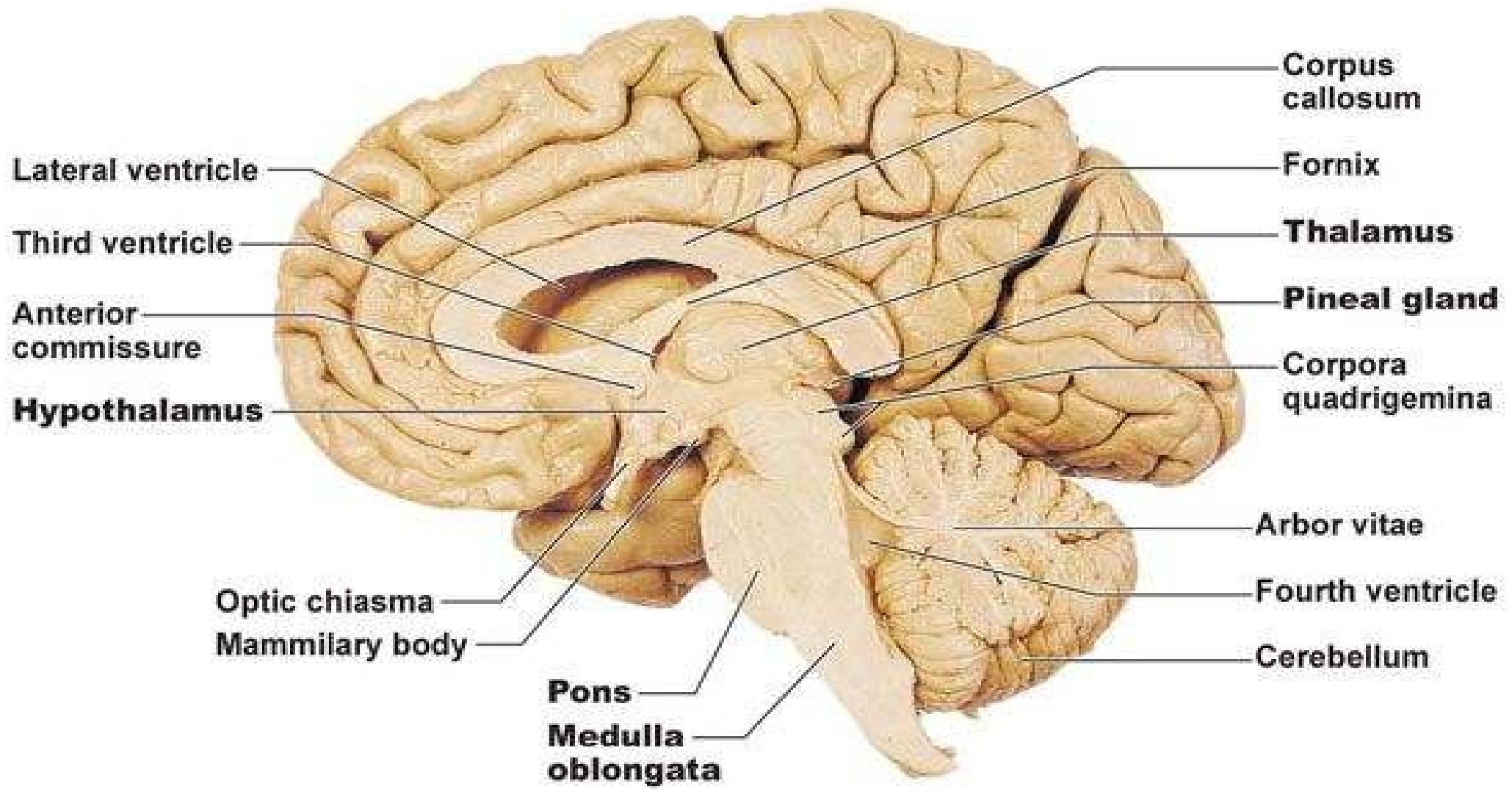
Lungs	Angiotensin II
Kidney	1,25-Dihydroxyvitamin D (vitamin D)
Adipose, mammary glands, other organs	Estradiol-17 $\beta$
Liver, sebaceous gland, other organs	Testosterone
Genital skin, prostate, other organs	5-Dihydrotestosterone (DHT)
Many organs	T <sub>3</sub>

**Table 37-2. Steroid Hormones**

Family	Number of Carbons	Specific Hormone	Primary Site of Synthesis	Primary Receptor
Progestin	21	Progesterone	Ovary Placenta	Progesterone receptor (PR)
Glucocorticoid	21	Cortisol Corticosterone	Adrenal cortex	Glucocorticoid receptor (GR)
Mineralocorticoid	21	Aldosterone 11-Deoxycorticosterone	Adrenal cortex	Mineralocorticoid receptor (MR)
Androgen	19	Testosterone Dihydrotestosterone	Testis	Androgen receptor (AR)
Estrogen	18	Estradiol-17 $\beta$ Estriol	Ovary Placenta	Estrogen receptor (ER)

# NEURO- ENDOCRINE

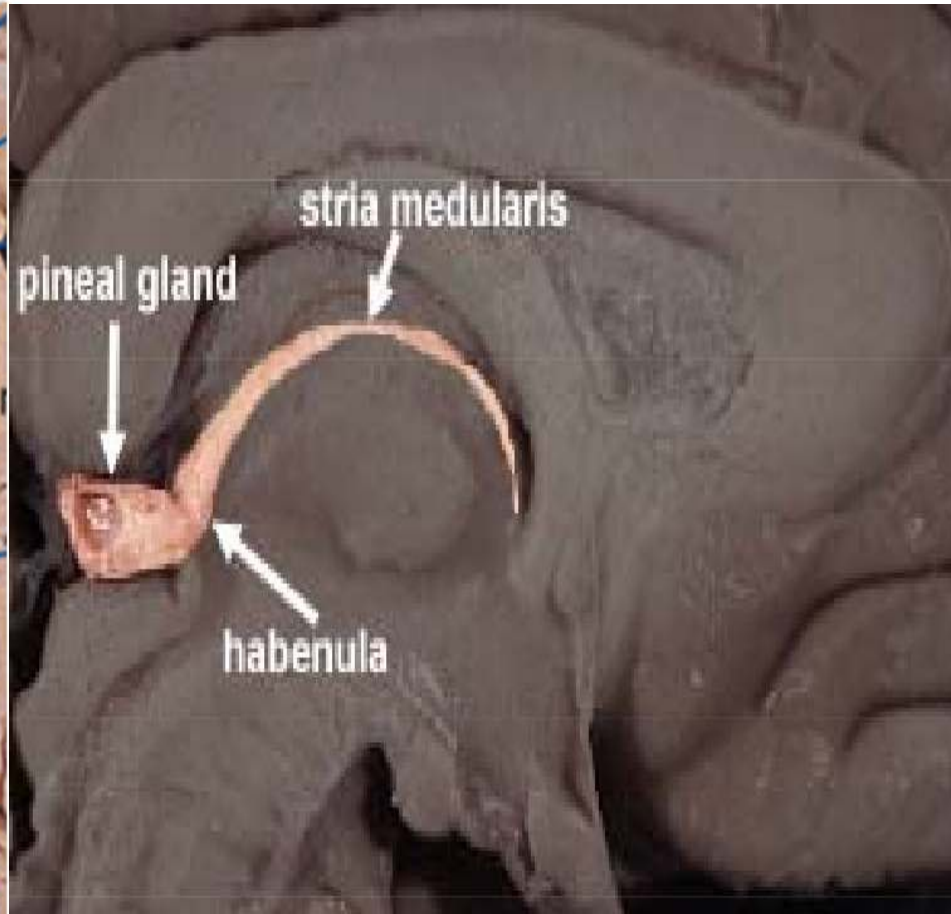
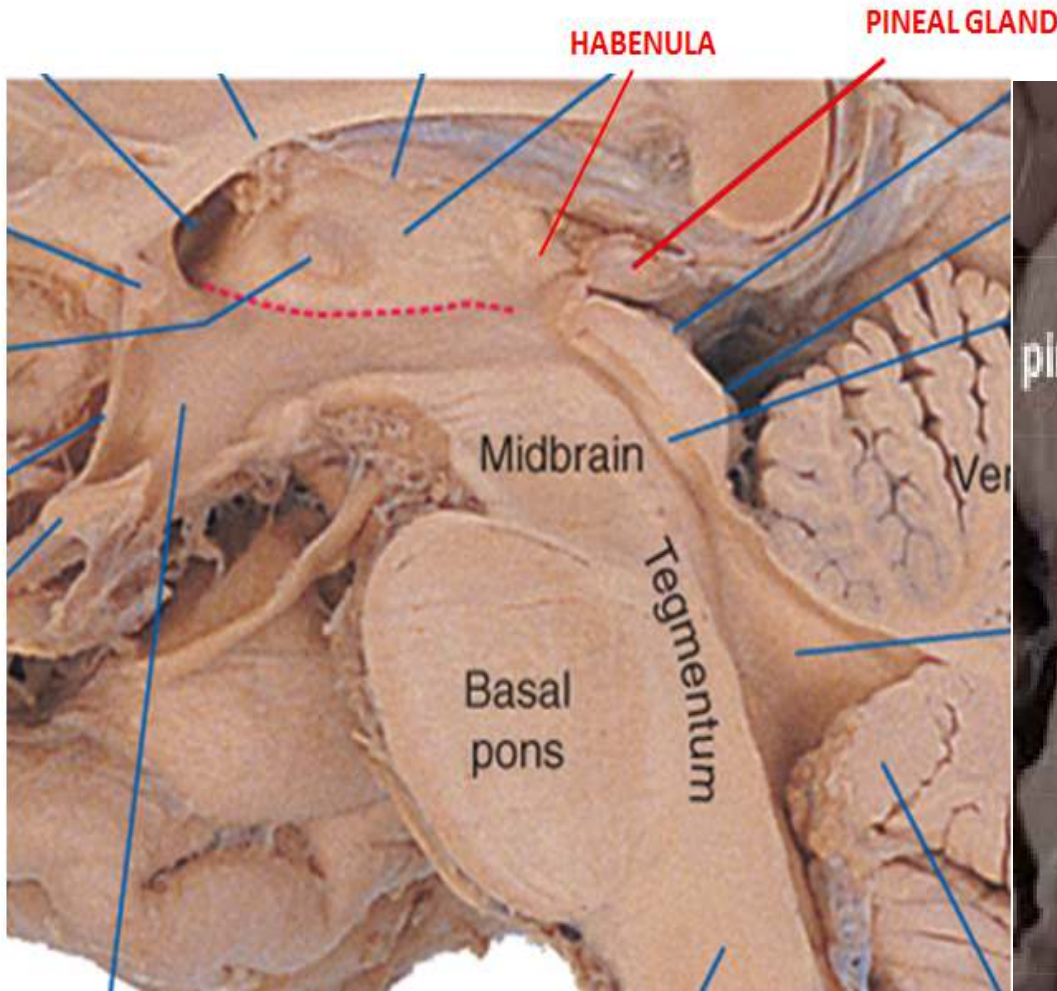




# EPITHALAMUS

MADE UP OF PINEAL GLAND AND HABENULA

- Habenula involved in food and water intake

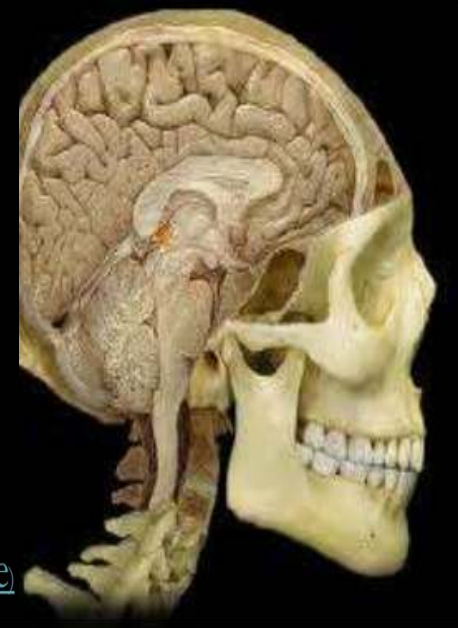
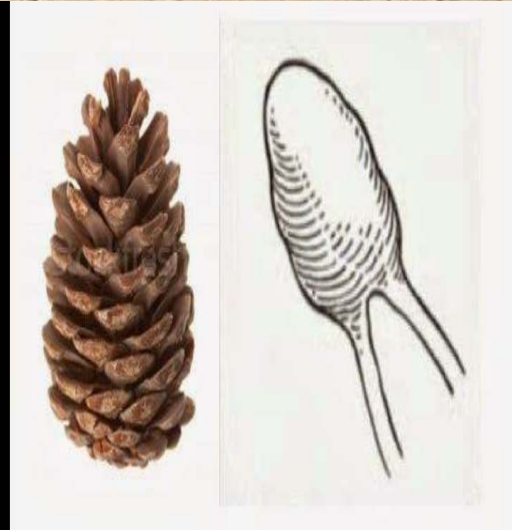




# Pineal Gland

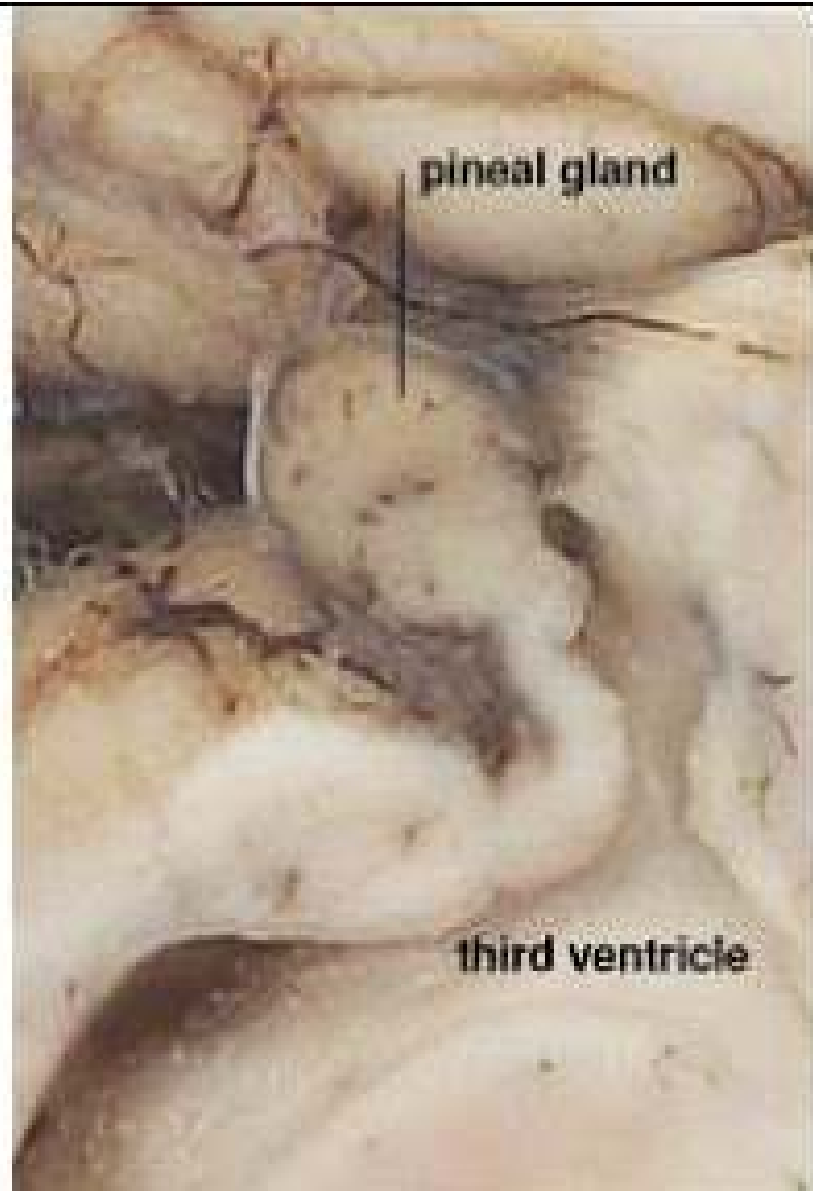
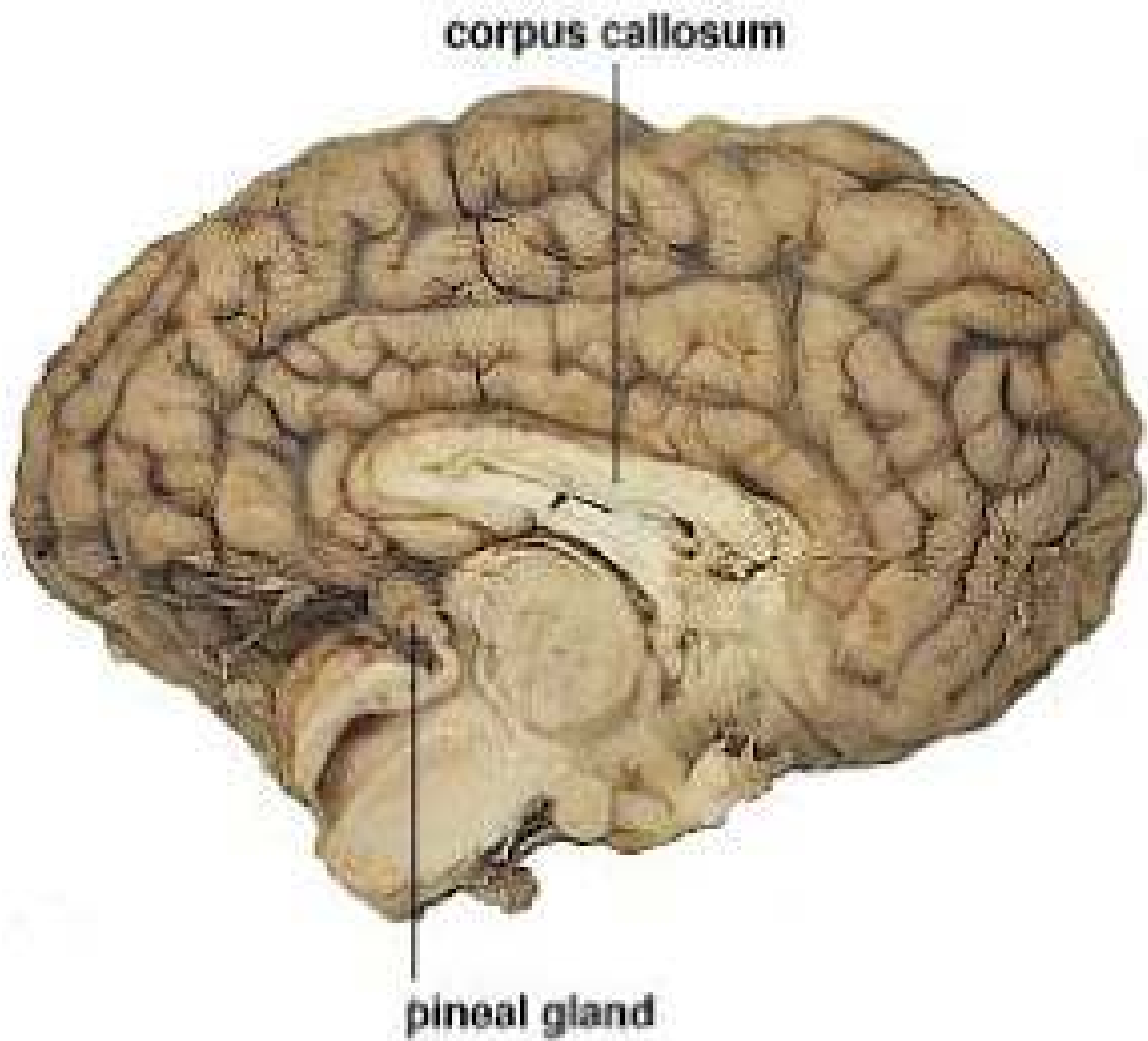
The 3<sup>rd</sup> eye of anubis

- AKA epiphysis cerebi
- **Pinealocytes secrete melatonin**
  - Involved in diurnal rhythms
  - Innervated by neurons of the ANS
- Brain Sand
  - Crystallized deposits of calcium carbonates and calcium phosphates



Its shape resembles a tiny pine cone (hence its name)





- The pineal body is **surrounded by pia mater**, which functions as its capsule and which sends connective tissue septa into the pineal body, subdividing it into lobules.
- **In the pineal we find two cell types:**
  - **pinealocytes** (about 95% of the cells; large, light and round nuclei)
  - **astrocytes** (glial cells; dark, elongated nuclei).
- Aside from the cells the pineal gland also contains ..... sand - well - **brain sand (or acervuli cerebri or - just for good measure - corpora arenacea)**. These are calcium-containing concretions in the pineal parenchyma, which increase in size and number with age.
- **The most prominent secretory product of the pineal body is melatonin.**
  - they may **"delay" puberty through anti-gonadotrophic effects.**
- blocks the secretion of gonadotropins (LH & FSH) from anterior pituitary gland. inhibit ovarian activity
- • These hormones aid in the proper development and functioning of the ovaries and testes
- Secretory activity in the pineal gland is stimulated by **darkness and inhibited by light.**
- Via the effects of pineal hormones on the adenohypophysis and sex hormones it is likely that the pineal body is involved in phenomena associated with **the circadian rhythm and seasonal phenomena** (e.g. seasonal affective disorder, SAD).
- The pineal body is innervated by **postganglionic sympathetic fibres derived from the superior cervical ganglion.**
- **serotonin** serotonin -neuro transmitter , vasoconstrictor
  - stimulates smooth muscles and inhibits gastric secretion

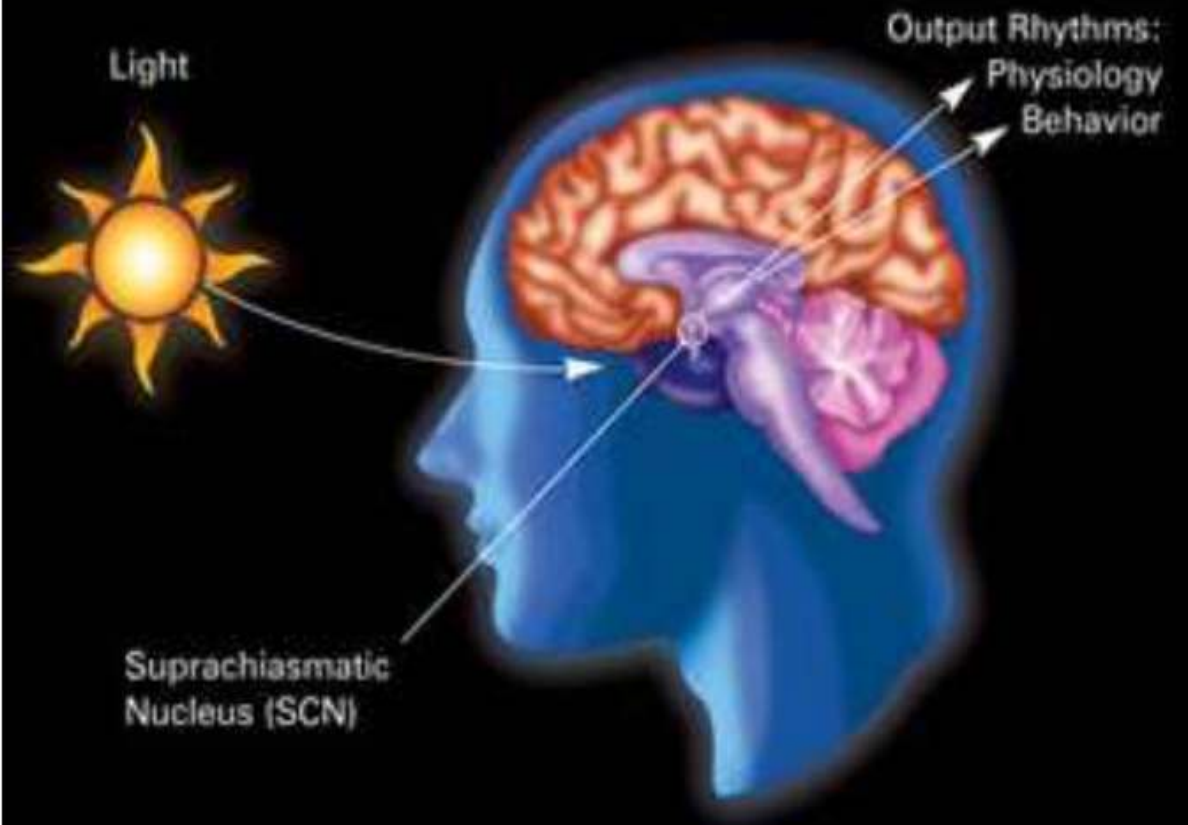
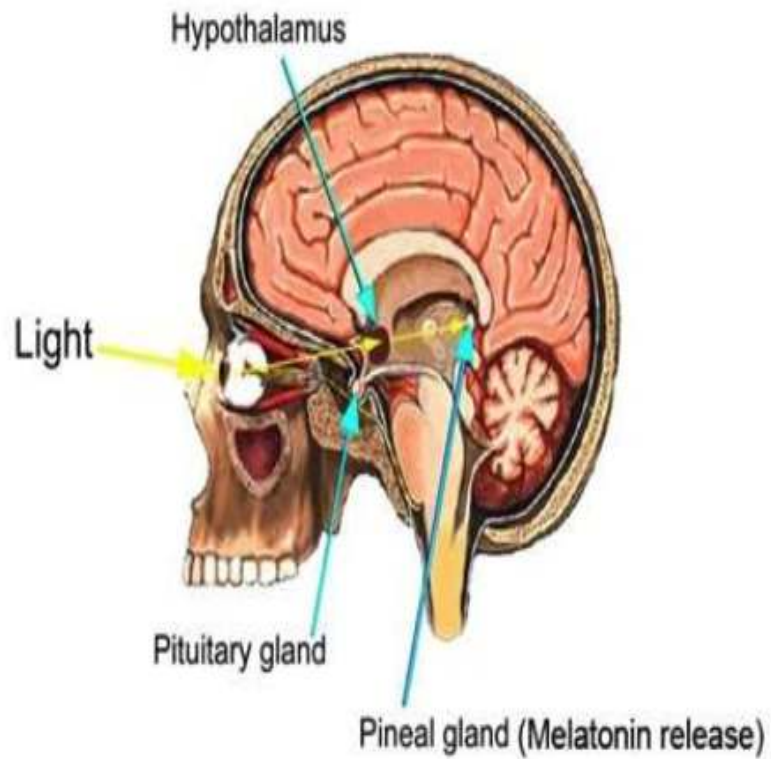
### **Melatonin effects :**

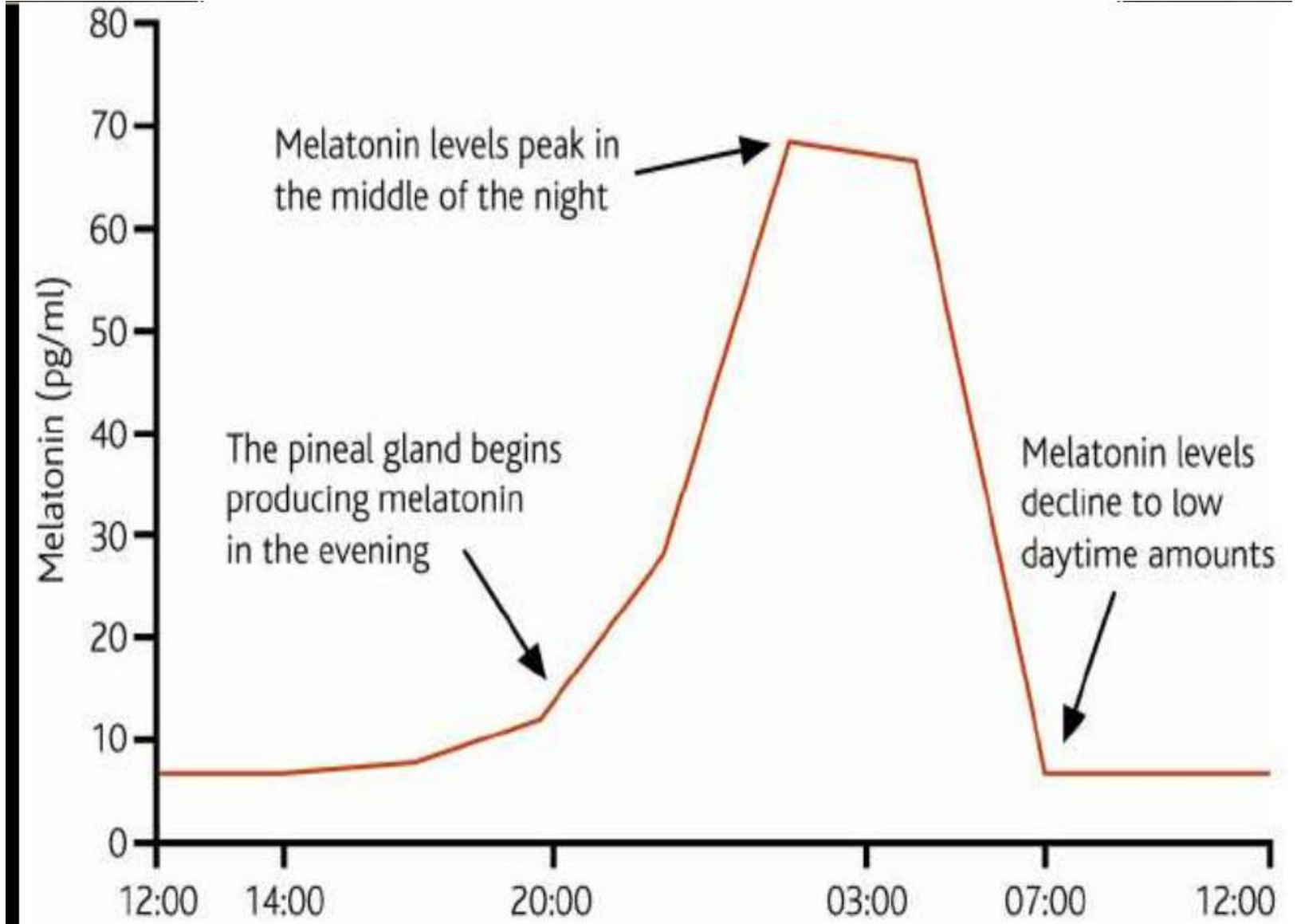
▣ **Dreaming:** Some supplemental melatonin users report an increase in vivid dreaming .

Extremely high doses of melatonin (50mg) dramatically increased REM sleep time and dream activity in both people with and people without narcolepsy .

▣ **Autism Individuals with autism spectrum disorders (ASD) may have lower than normal levels of melatonin**

- Natural Sleeping Tablet





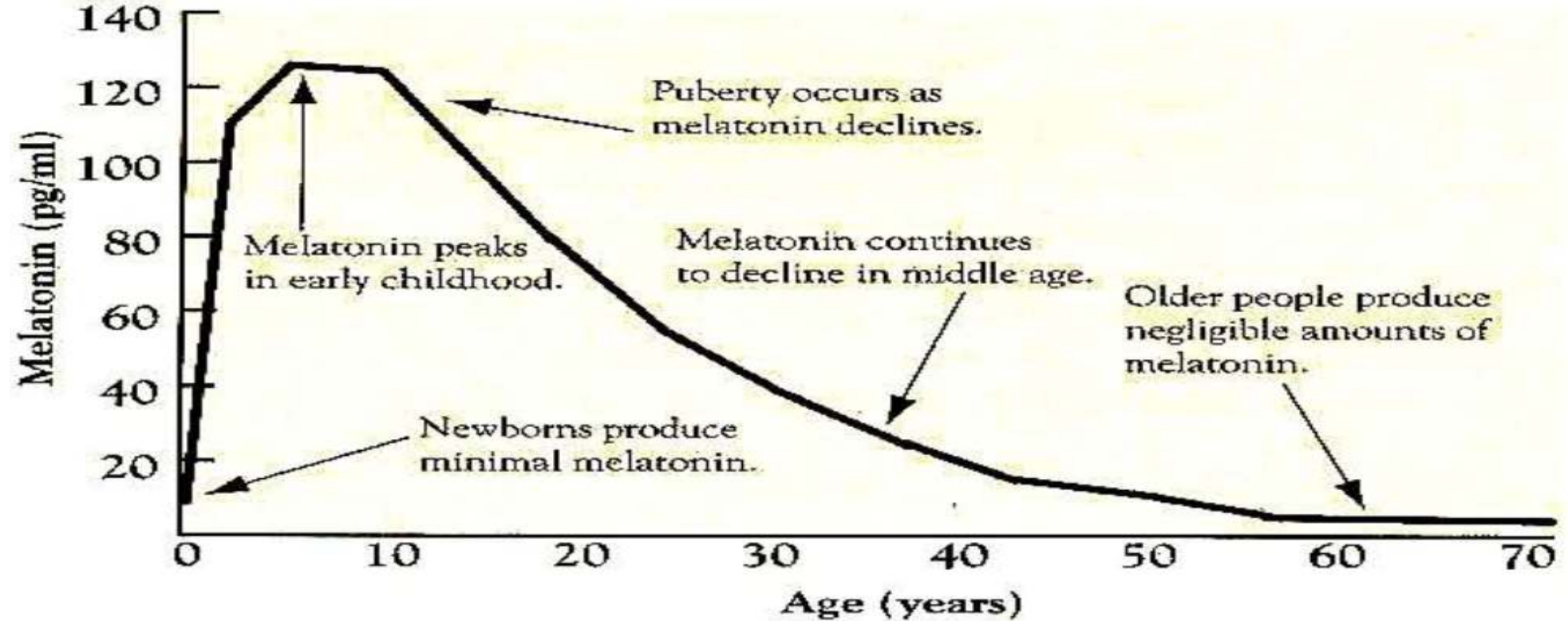
Melatonin levels peak in the middle of the night

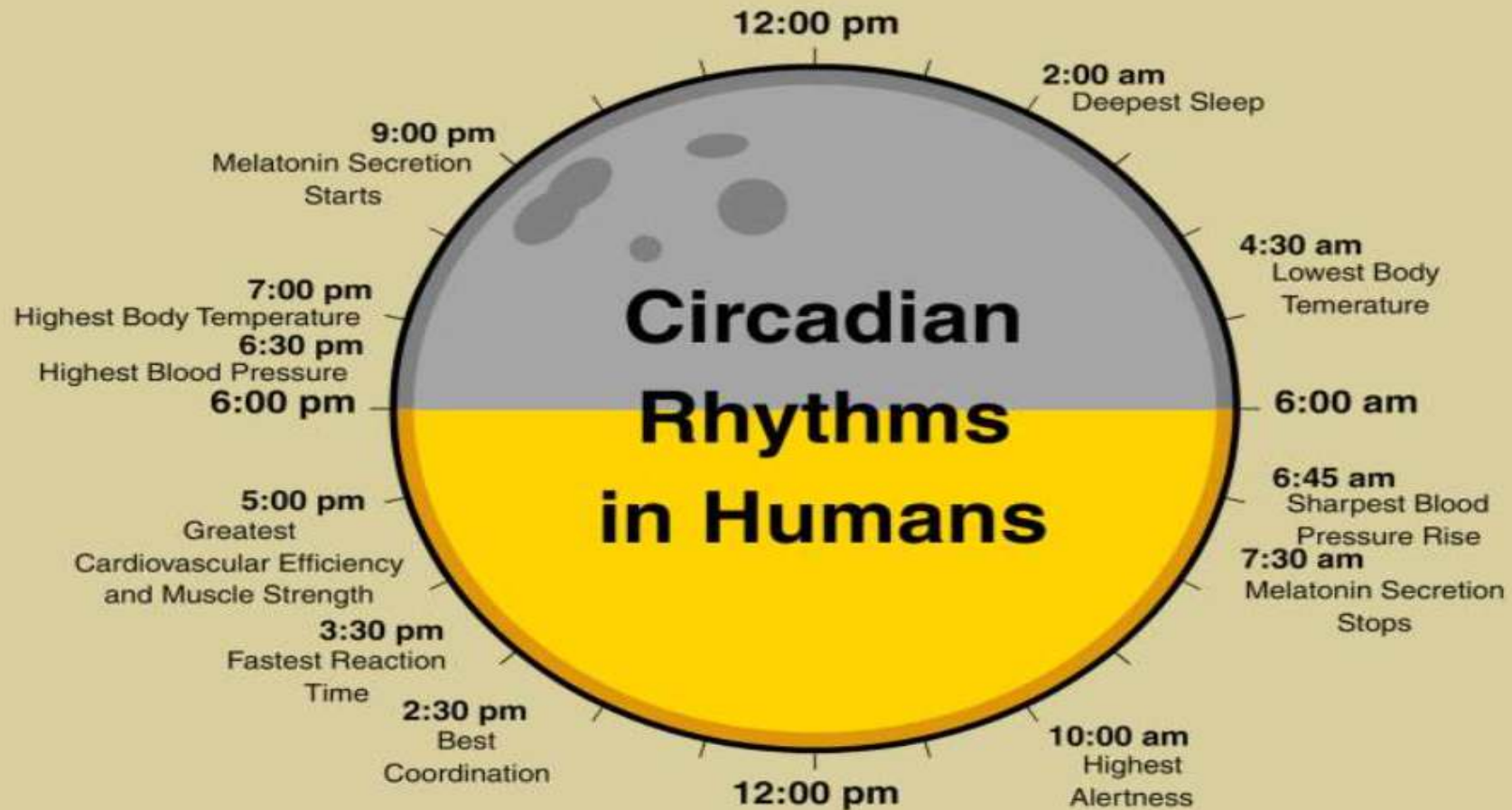
The pineal gland begins producing melatonin in the evening

Melatonin levels decline to low daytime amounts



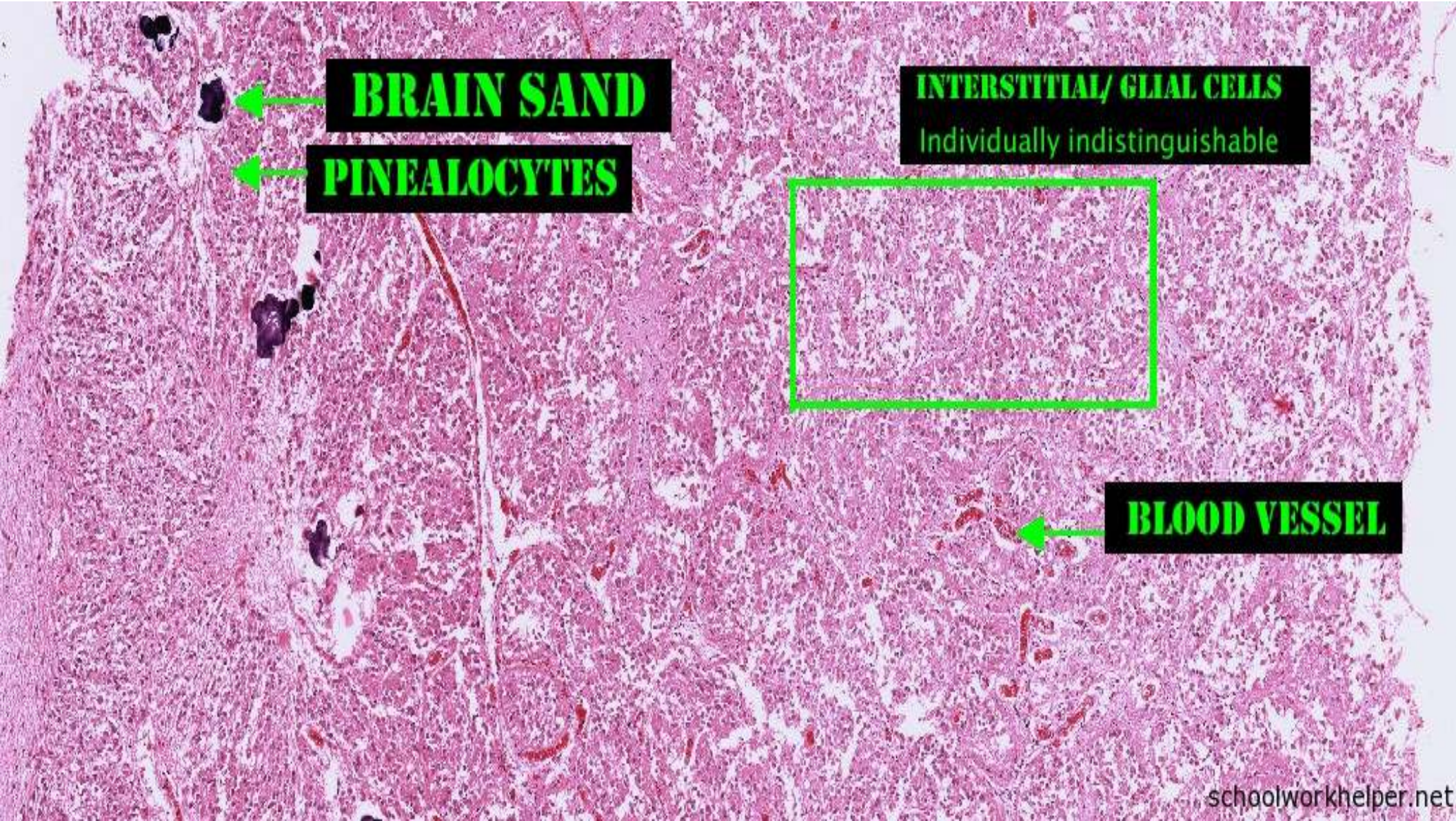
# MELATONIN VS AGE





Circadian rhythms are physical, mental and behavioral changes that follow a roughly 24-hour cycle, responding primarily to light and darkness in an organism's environment.





**BRAIN SAND**

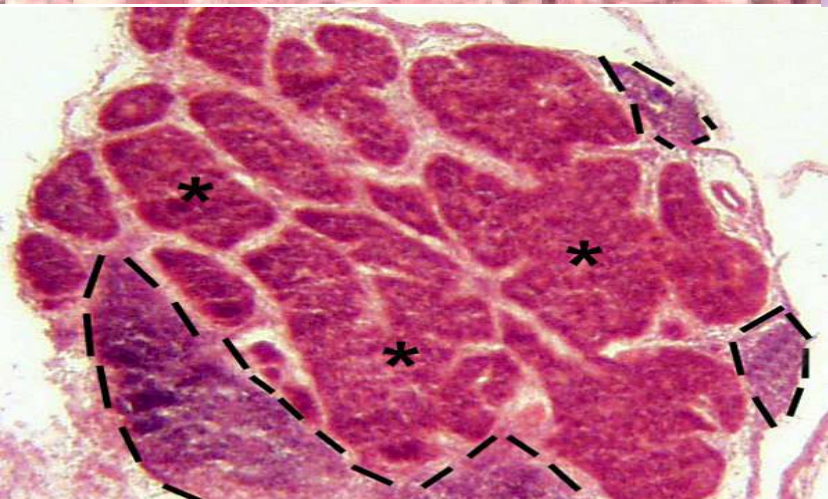
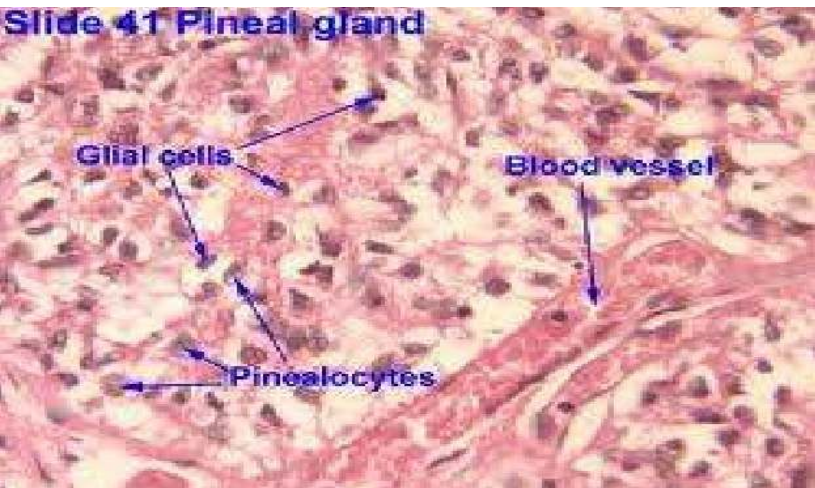
**PINEALOCYTES**

**INTERSTITIAL/ GLIAL CELLS**  
Individually indistinguishable

**BLOOD VESSEL**

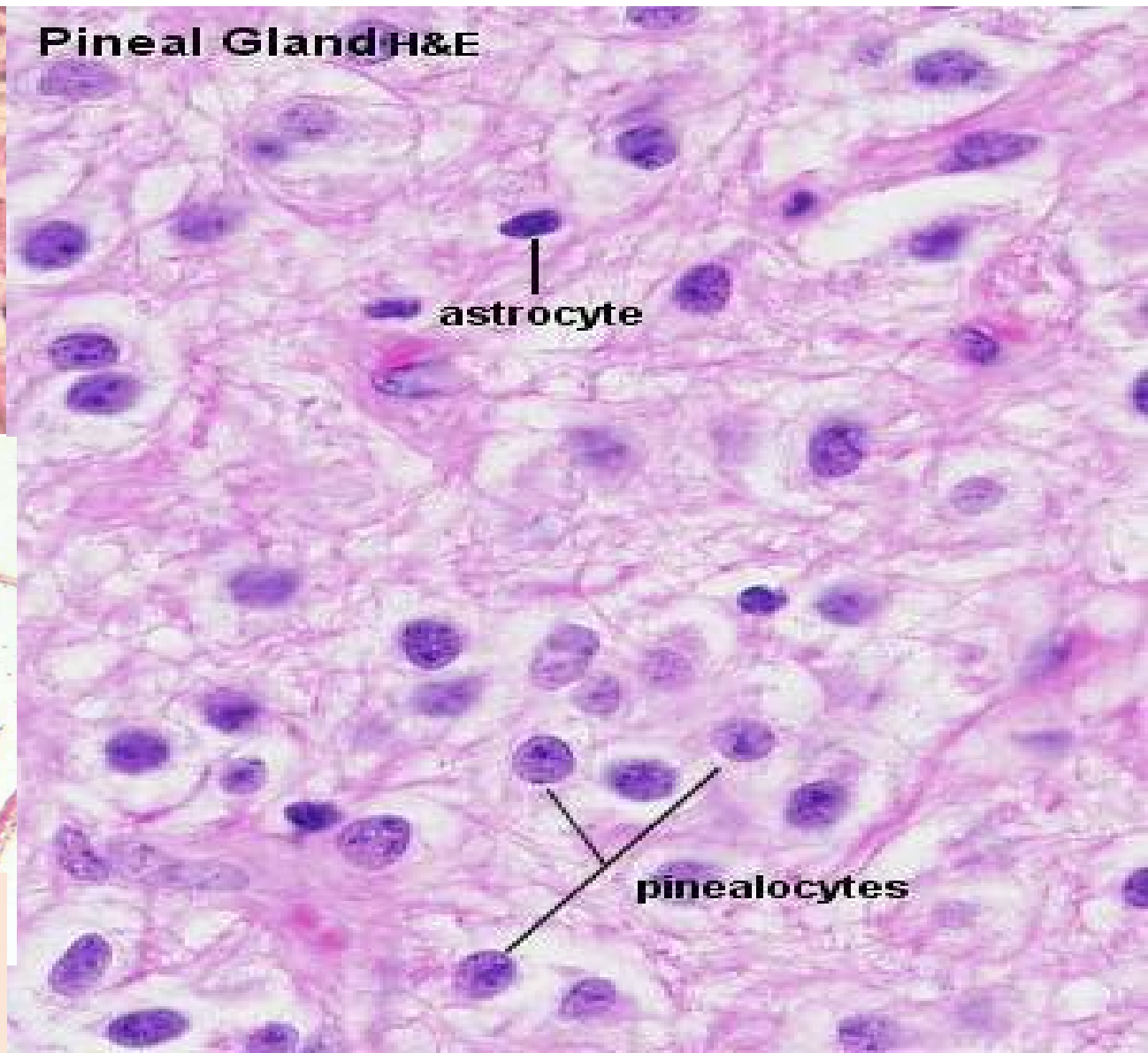


Slide 41 Pineal gland



The pineal follicles (\*) comprise pinealocytes and supportive cells arranged as epithelium. Prominent interstitial septa separate individual follicles.

Pineal Gland H&E



## **FUNCTIONS OF THE NEUROENDOCRINE SYSTEM**

Along with the nervous system, hormones provide the necessary communication between all the cells that constitute a multicellular animal

**Nervous system**-Is involved in rapid transfer of short-term events and coordination of short-term events. Electrochemical information involving neurons.

**Neurosecretory cells**-Neurons have electrical activity but involved in the production and release of neurosecretion that produces their effect as chemicals.

**Endocrine system**-Is involved in the integration and coordination of long-term events through chemicals called hormones.



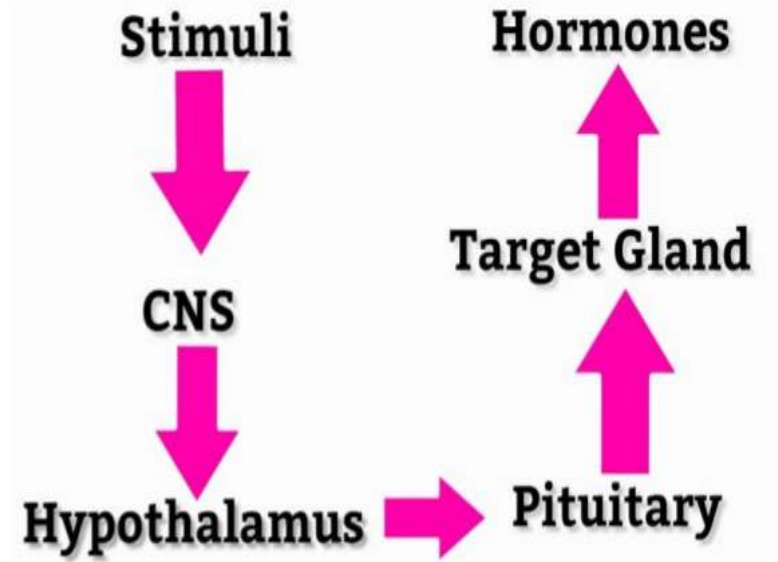
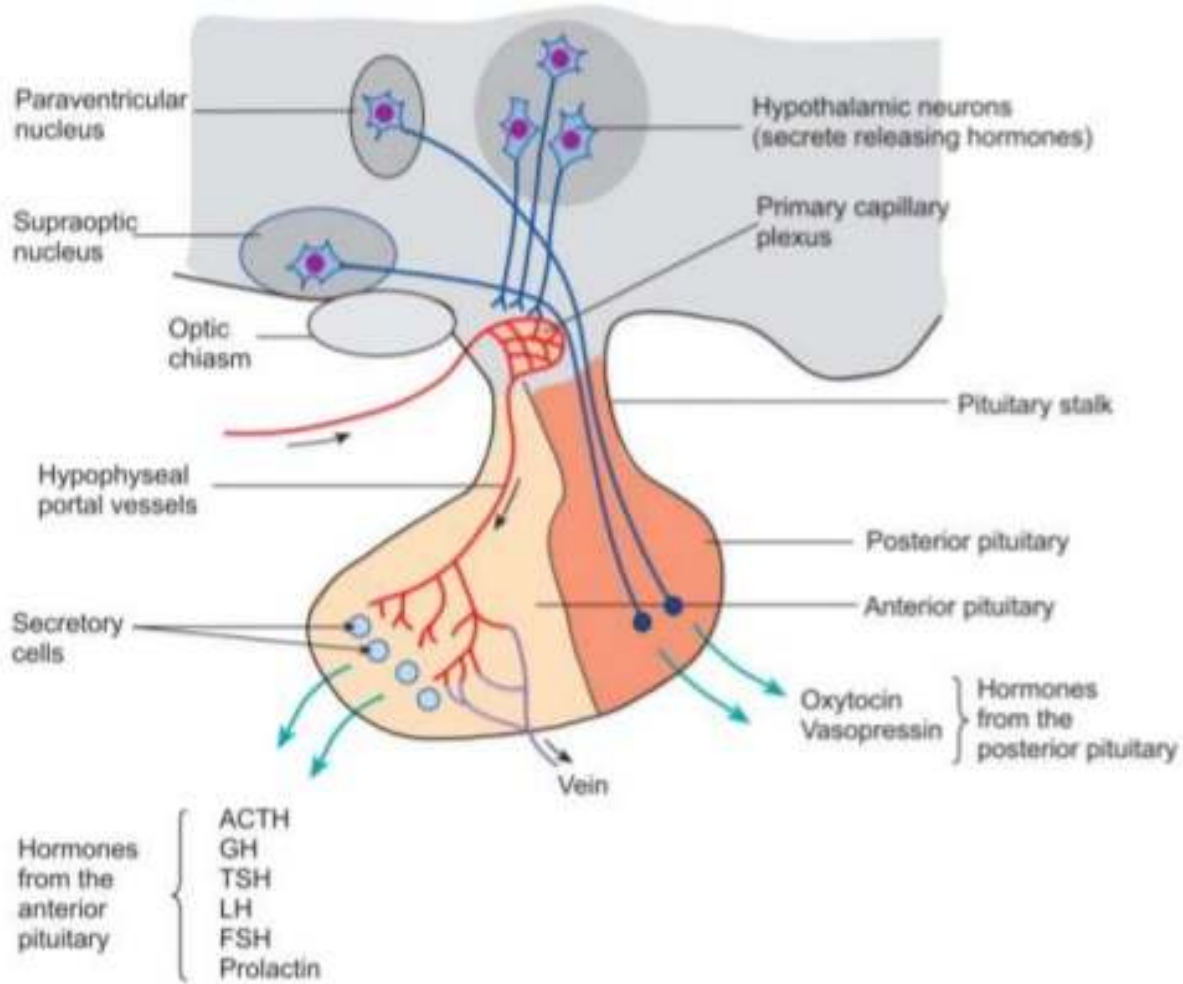
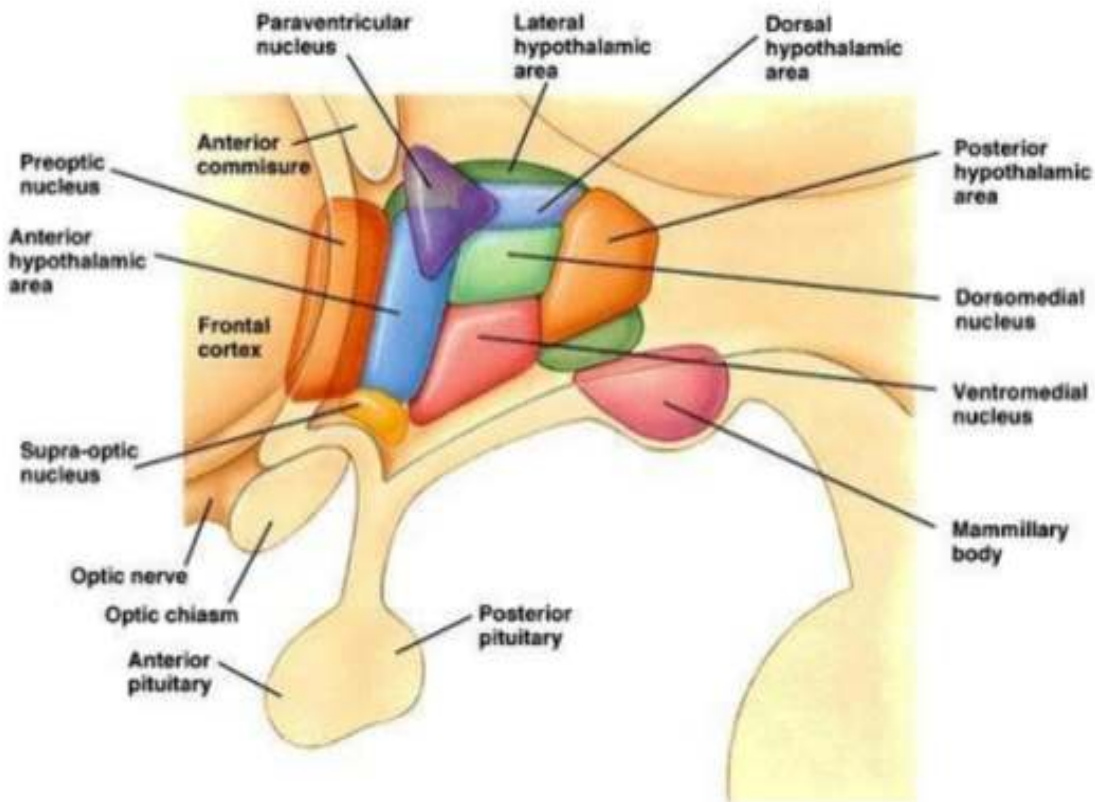


Fig. Showing ,The secretory activity of many endocrine glands is controlled by the nervous system.

# Hypothalamus

Nuclei of the Hypothalamus



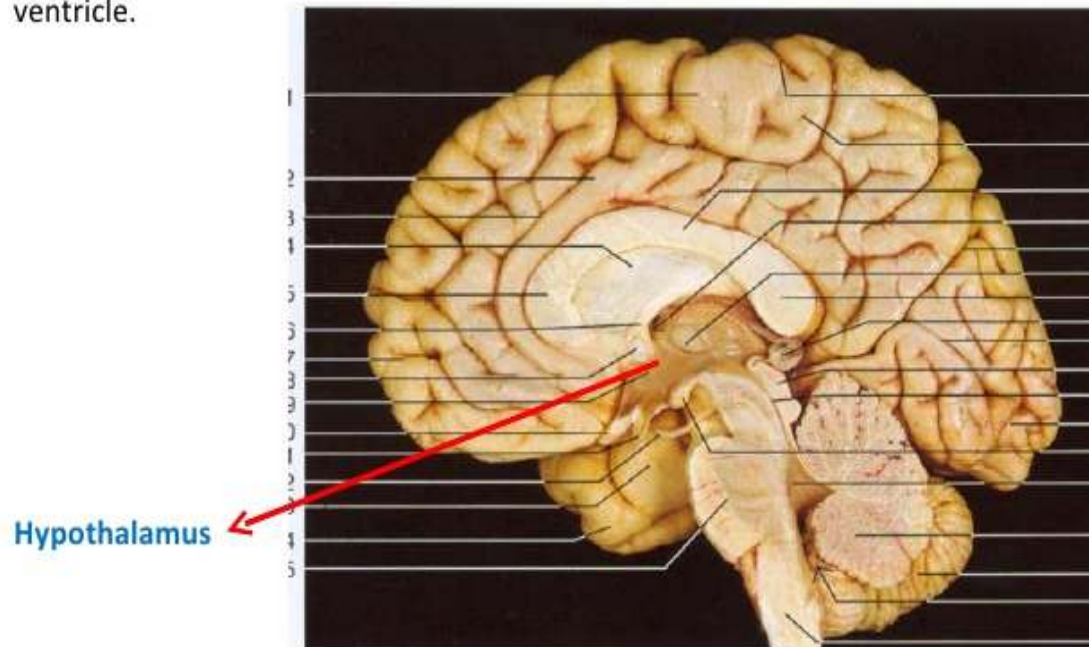
## Functions of Hypothalamus. (AS-RESPECT)

- Autonomic functions.
- Sleep –wake cycle.
- Reward & punishment centre.
- Endocrinal functions.
- Sexual behaviour & reproduction.
- Ph(F)ood intake regulation.
- Emotional & Instinctual behaviour.
- Circadian Rhythm control.
- Temperature regulation.

## Hypothalamus

The hypothalamus consists of only 4 cm<sup>3</sup> of neural tissue, or 0.3% of the total brain.

The hypothalamus extends from the lamina terminalis to a vertical plane posterior to the mammillary bodies, and from the hypothalamic sulcus to the base of the brain beneath the third ventricle.



- The hypothalamus contains neurons that control releases from the anterior pituitary.
- Seven hypothalamic hormones are released into a **portal system** connecting the hypothalamus and pituitary, and cause targets in the pituitary to release eight hormones.

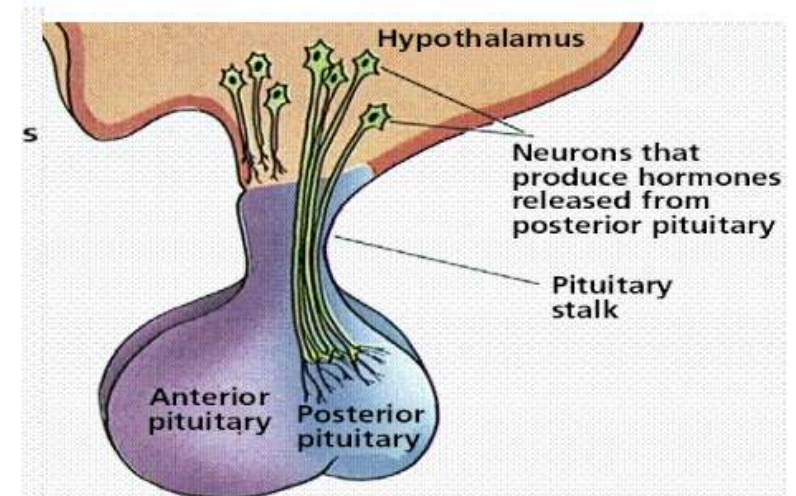
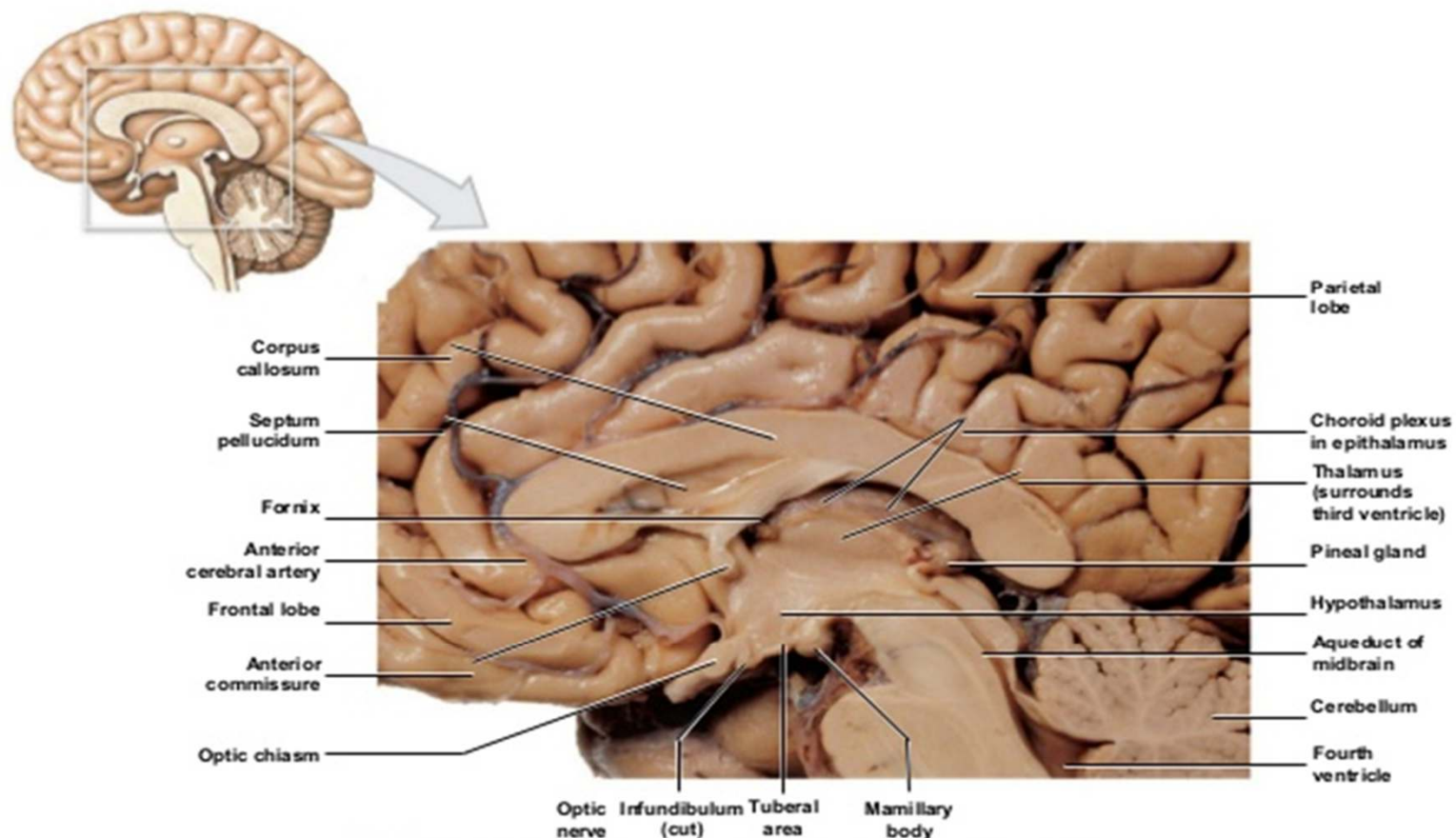




Figure 16.12a The Hypothalamus



■ Midsagittal section through the brain. This view shows the major features of the diencephalon and adjacent portions of the brain stem.

# PRINCIPAL HYPOTHALAMIC POLYPEPTIDE.

## Food intake increased by

- Neuropeptide Y.
- Orexin – A
- Orexin – B
- Melanin concentrating hormone. (MCH)
- Ghrelin.

## Food intake decreased by

- Cocaine & amphetamine regulated transcript. (CART)
- CRH.



## Endocrinal functions.

---

- Anterior Pituitary.(through Tubero-infundibular tract & hypophyseal portal system)
- Controls Thyroid G.
- Controls Metabolism through adrenal gland.
- Keep gonads inhibited.
- Control formation of milk by prolactin secretion.
- Posterior Pituitary regulate water balance through ADH.
- Regulation Of Uterine Contractility & regulation of Milk Ejection from breast through oxytocin.

# RAGE

**Rage** – violent & aggressive emotional state by strong stimulation of **Punishment Centre**.

Kept in check by counterbalancing activity of Ventromedial N of hypothalamus, hippocampus, amygdala & ant portion of limbic cortex.

- **Characterized by –**
  - Development of defense posture.
  - Extension of limbs
  - Lifting of tail.
  - Hissing & splitting
  - Piloerection.
  - Wide openings of eye.
  - Pupil dilation.

## Hypothalamic control of Anterior pituitary gland secretion

- **Hypothalamus controls the hormonal secretions of the anterior pituitary**, which in turn regulates other endocrine glands.
- Neurons in the hypothalamus **secrete releasing hormones and inhibiting hormones** into blood capillaries at the base of the hypothalamus.
- Releasing & inhibiting hormones released by **Paravocellular Neurosecretory cells of the hypothalamus**.

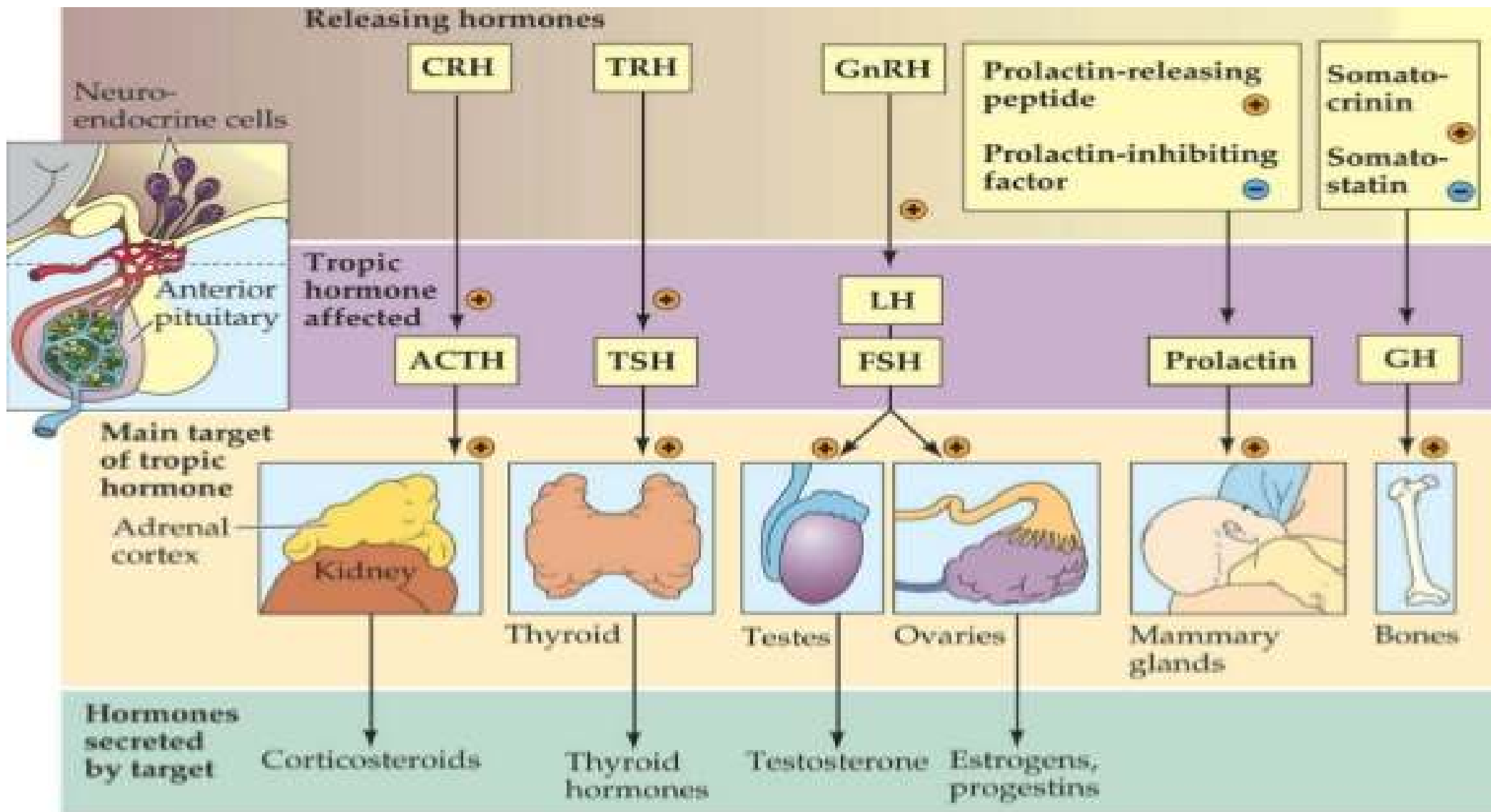
### Releasing and Inhibiting hormones

#### Hypothalamic -Releasing hormones :

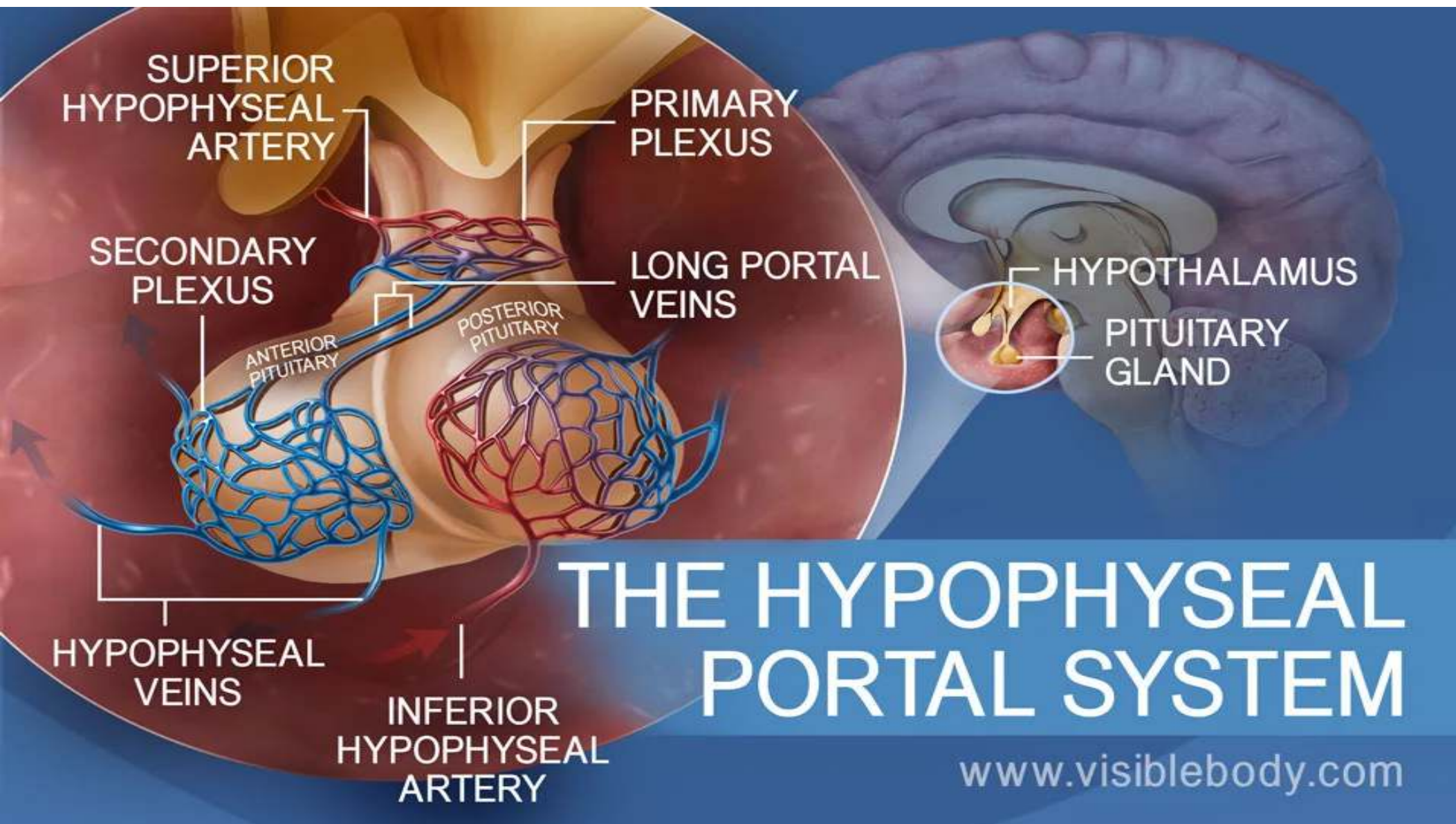
- CRH (Corticotropin releasing Hormone) ==> Stimulates the release of ACTH
- TRH (Thyrotropin-Releasing Hormone) ==> Stimulates the release of TSH
- GnRH (Gonadotropin-Releasing Hormone ) ==> stimulates the release of FSH& LH
- GHRH(Growth Hormone Releasing Hormone) ==> Stimulates the release of GH

# Hypothalamic releasing hormones

Hypothalamic releasing hormone	Effect on pituitary
Corticotropin releasing hormone (CRH)	Stimulates ACTH secretion
Thyrotropin releasing hormone (TRH)	Stimulates TSH and Prolactin secretion
Growth hormone releasing hormone (GHRH)	Stimulates GH secretion
Somatostatin	Inhibits GH (and other hormone) secretion
Gonadotropin releasing hormone (GnRH)	Stimulates LH and FSH secretion
Prolactin releasing hormone (PRH)	Stimulates PRL secretion
Prolactin inhibiting hormone (dopamine)	Inhibits PRL secretion



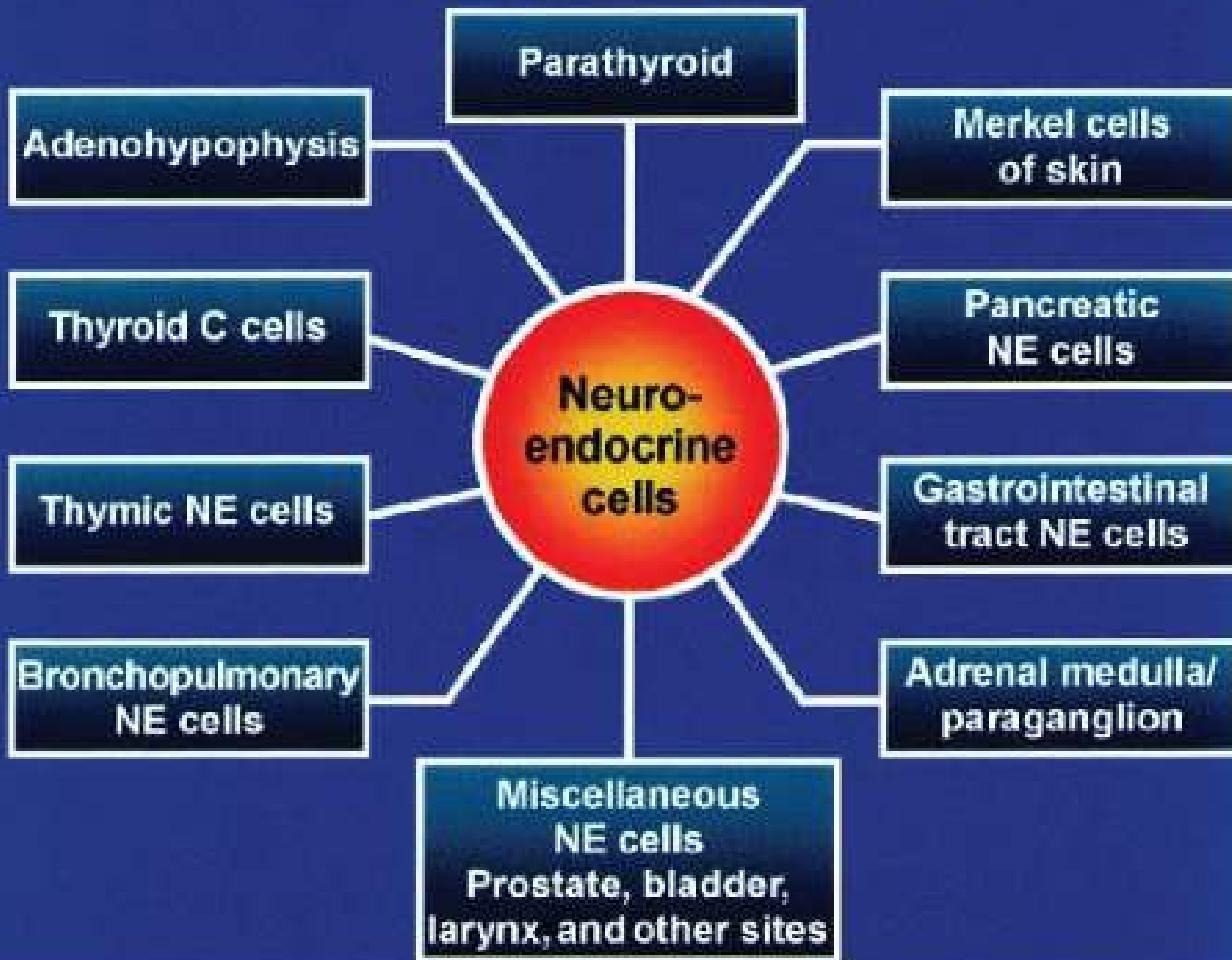




# THE HYPOPHYSEAL PORTAL SYSTEM

[www.visiblebody.com](http://www.visiblebody.com)

## Diffuse Neuroendocrine System



diffuse neuroendocrine system, which is composed of classic endocrine organs as well as scattered neuroendocrine cells in various organs and tissues.

- **Neurohypophysis**- It shows presence of axons of neuron , cell bodies of which are situated in hypothalamus. Interspersed among the nerve fibre few neuroglial cells called as pituicytes.

## Pituitary Gland

### MASTER GLAND

- Also known as Hypophysis Cerebri
- Situated in the hypophyseal fossa in middle cranial fossa
- Histologically it shows two parts-  
Adenohypophysis and Neurohypophysis  
**Adenohypophysis** shows presence of chromophobes & chromophils  
Chromophils are of two types- Acidophils & Basophils.

## Anterior Pituitary: Hormones

### •Anterior pituitary hormones

- FLAT PiG
  - FSH (follicle-stimulating hormone)
  - LH (luteinizing hormone)
  - ACTH (adrenocorticotrophic hormone)
  - TSH (thyroid-stimulating hormone)
  - Prolactin
  - Growth hormone (somatotropin)
- categories of hormones
  - corticolipotropins
    - ACTH and MSH (melanocyte-stimulating hormone)
  - glycoprotein hormones
    - FSH, LH, TSH
  - somatomammotropins
    - prolactin and growth hormone

### •Corticolipotropins

#### •synthesis

- corticolipotropins are derived from a single precursor, POMC

• POMC = pro-opiomelanocortin

- pathway details

#### •MSH

- corticolipotropin synthesis products (aka fragments) contain MSH
- increased MSH levels → skin pigmentation
- e.g., Addison's disease
- **↑ ACTH → ↑ MSH → skin pigmentation**

### •Glycoprotein hormones

- subunits of peptide hormones
  - glycoprotein hormones contain 2 subunits: **α and β subunit**
    - α subunits identical, β subunits non-identical
    - hormone specificity determined by β subunit
- human chorionic gonadotropin (hCG) structurally related to glycoprotein hormone
  - **hCG contains identical α subunit**

### ACIDOPHILS (growth)

- GROWTH HORMONE
- PROLACTIN

### BASOPHILS (trophy)

- TSH
- ACTH
- LH, FSH

- Somatotammotropins

- prolactin

- growth hormone

- secretion

- pulsatile secretory pattern

- secretory **bursts approximately every 2 hours**

- ↑ in secretory bursts during exercise and sleep

- functions

- ↑ linear growth and muscle mass

- growth mediated by production of somatomedins

- aka insulin-like growth factors (IGFs)

- diabetogenic effect

- insulin resistance

- decreases glucose uptake and utilization

- "diabetogenic"

- growth hormone produces increases in blood glucose



# Hormones secreted by anterior pituitary

1. **FSH** (follicle stimulating hormone)

1. **LH** (luteinizing hormone)

The above two are called **gonadotropins**

3. **TSH** (thyroid stimulating hormone, thyrotropin)

4. **ACTH** (adrenocorticotrophic hormone)

5. **GH** (growth hormone; somatotropin or somatotrophic hormone)

6. **PRL** (prolactin)

• **Tropic (trophic) hormones--** target other endocrine glands to release their own hormones.

**Hormones from basophils** :go to other endocrine glands, thyroid, adrenal cortex, ovary, testis. Cells from acidophils do NOT.

**Acidophils make GROWTH related hormones. Basophils make hormones which STIMULATE OTHER endocrine glands.**

**Chromophobes make NOTHING.**

- When stained with the PAS reaction all three types of basophils appear reddish

## **Chromophobe cells**

**anterior pituitary cells that lack granules and that do not react with acidophilic/basophilic stains**

**e.g., stromal cells and degranulated chromophils**

- **Chromophobe cells are unstained or weakly stained cells. appears relatively pale under the microscope**
- EM and immunocytochemistry are used.
- They are now thought to represent acidophil and basophilic cells in a dormant or recently degranulated stage (degranulation = release of most of the secretory vesicles), but may also include stem cells of the secretory cells.

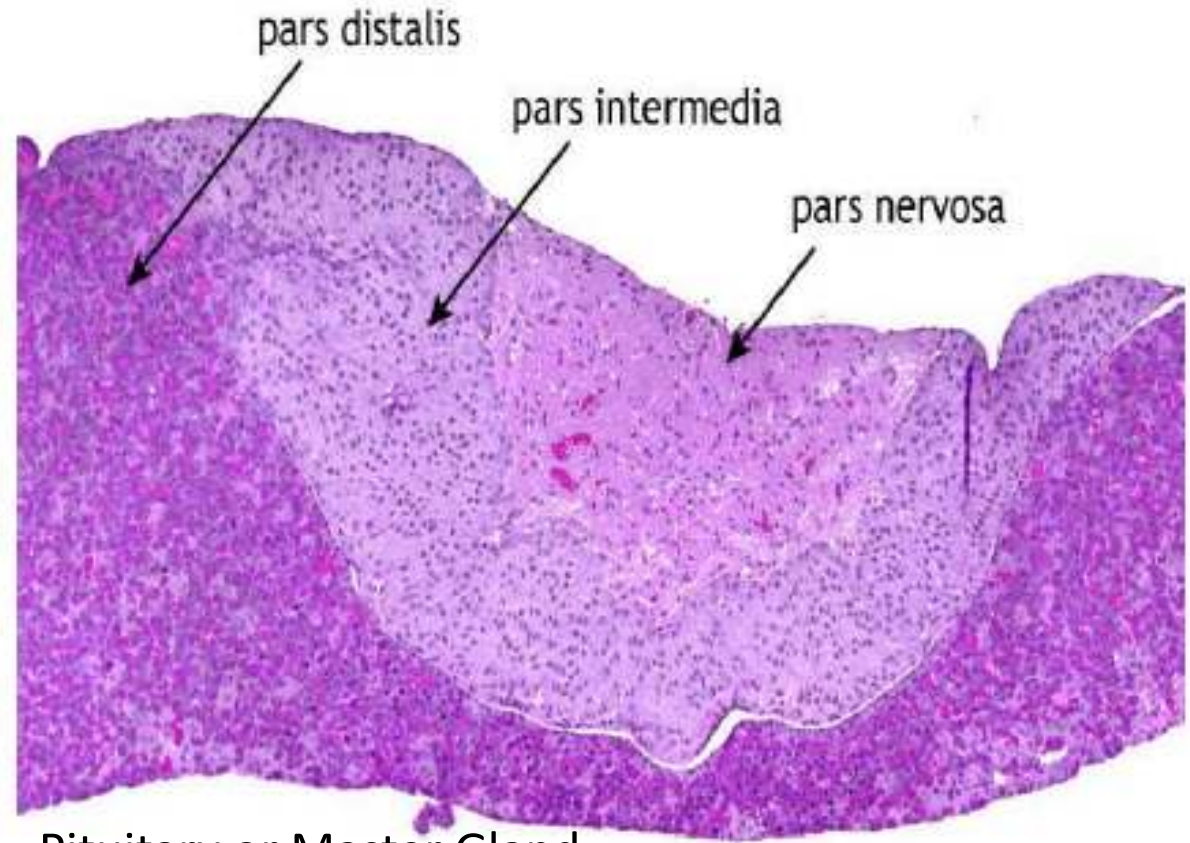
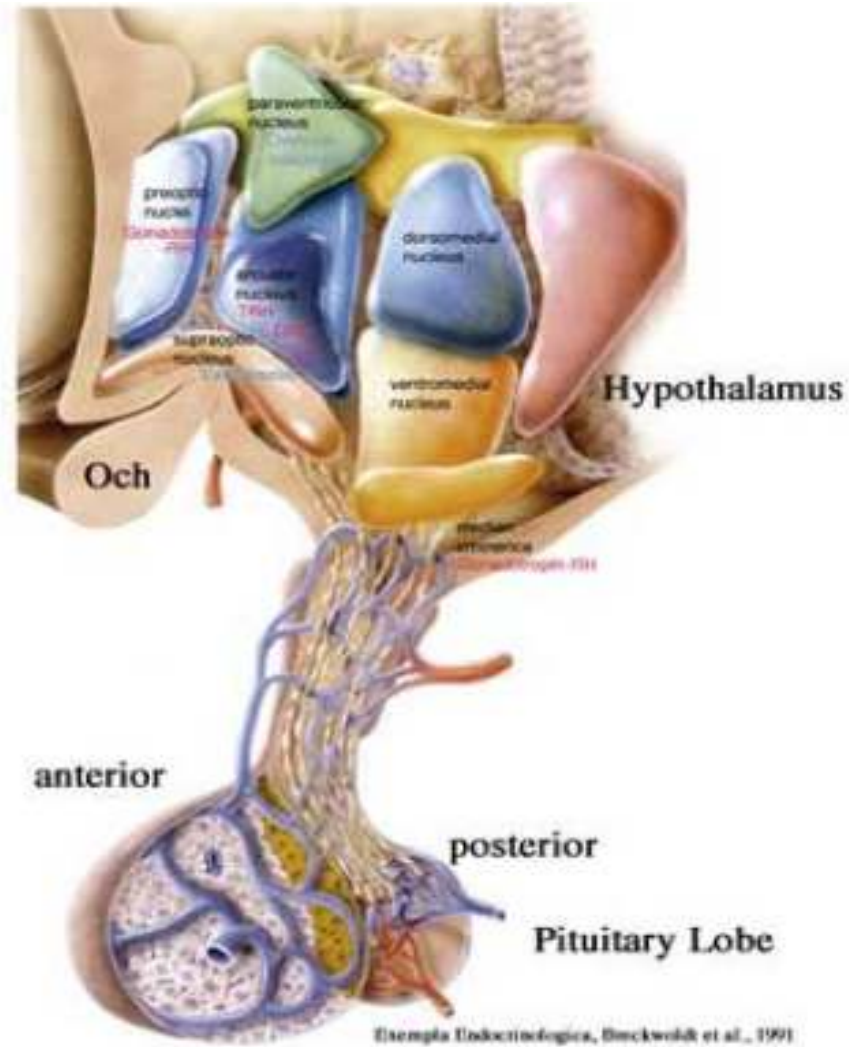
One type of chromophobe cell is known as amphophils.

- **Amphophils** are epithelial cells found in the anterior and intermediate lobes of the pituitary.
  - Together, these epithelial cells are responsible for producing the hormones of the anterior pituitary and releasing them into the bloodstream.
- **Melanotrophs** (also, Melanotropes) are another type of chromophobe which secrete melanocyte stimulating hormone (MSH).

Chromophobe" also refers to a type of renal cell carcinoma (distinct from "clear cell")  
30% of patients with Birt-Hogg-Dubé syndrome will also develop chromophobe renal cancer.

**Table 20.3****Pituitary Gland Hormones**

<b>Hormone</b>	<b>Target Cells</b>	<b>Effects of Hormone</b>
<b>HORMONES OF THE ANTERIOR PITUITARY</b>		
Adrenocorticotropic hormone (ACTH)	Adrenal cortex	Stimulates production of corticosteroid hormones
Follicle-stimulating hormone (FSH)	Female: Ovaries Male: Testes	Female: Stimulates growth of ovarian follicles Male: Stimulates sperm production
Luteinizing hormone (LH)	Female: Ovaries Male: Testes	Female: Stimulates ovulation, estrogen and progesterone synthesis in ovary Male: Stimulates androgen synthesis in testes
Thyroid-stimulating hormone (TSH)	Thyroid gland	Stimulates thyroid hormone synthesis and secretion
Prolactin (PRL)	Female: Mammary glands Male: Not known	Female: Stimulates milk production in mammary glands Male: May play a role in the sensitivity of the testes interstitial cells to LH
Growth hormone (GH)	Almost every cell in the body	Increased growth and metabolism in target cells; synthesis of somatomedin in the liver to stimulate growth at epiphyseal plate
Melanocyte-stimulating hormone (MSH)	Melanocytes	Stimulates synthesis of melanin and dispersion of melanin granules in epidermal cells
<b>HORMONES STORED IN THE POSTERIOR PITUITARY</b>		
Antidiuretic hormone (ADH) (also called vasopressin)	Kidney Smooth muscle in arteriole walls	Stimulates reabsorption of water from urine in kidneys Stimulates vasoconstriction in arterioles of body, thereby raising blood pressure
Oxytocin (OT)	Female: Uterus, mammary glands Male: Smooth muscle of male reproductive tract	Female: Stimulates smooth muscle contraction in uterine wall; stimulates milk ejection from mammary glands Male: Stimulates contraction of smooth muscle of male reproductive tract



Pituitary or Master Gland

posterior lobe

neurohypophysis

anterior lobe

adenohypophysis



### Acidophil cells (or acidophils)

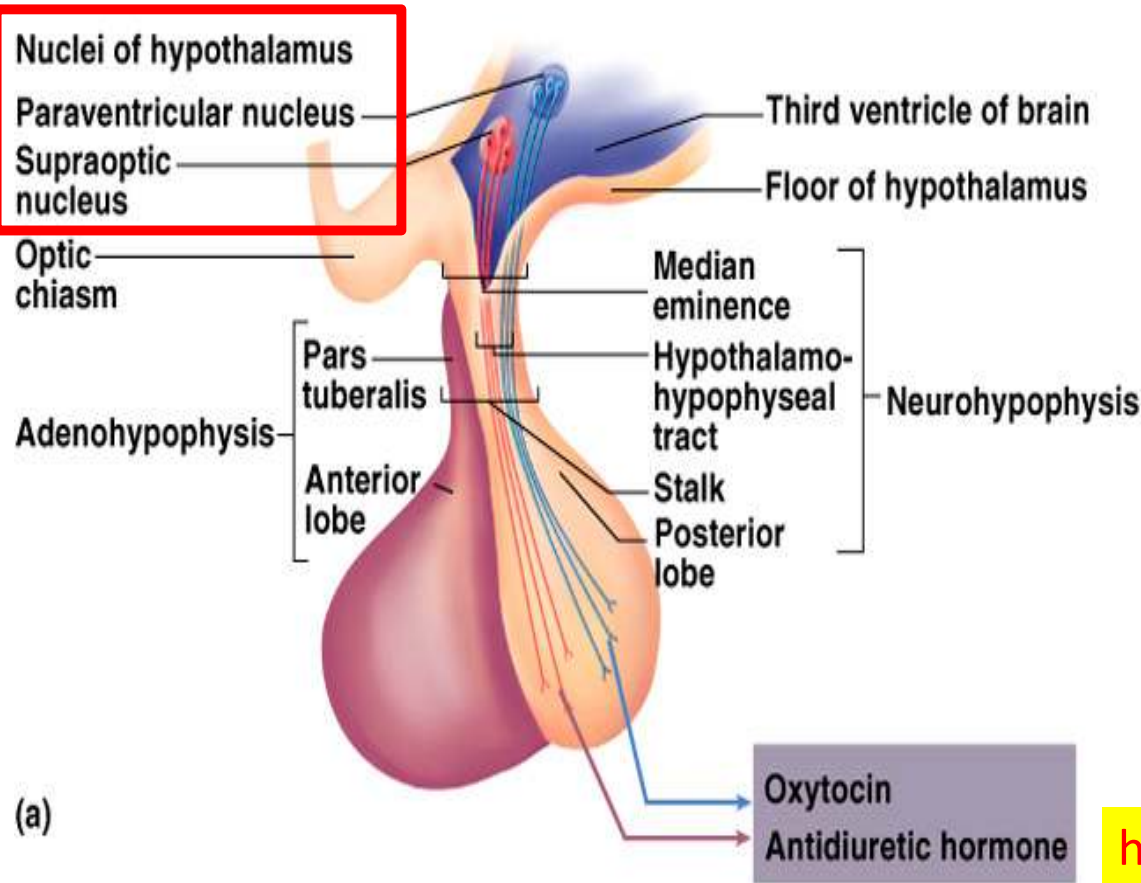
- Acidophils are rounded cells and typically smaller than basophil cells.
- Acidophils account for roughly 65% of the cells in the adenohypophysis.
- **The most frequent subtype of acidophils are the somatotrophs (which can be stained with the dye orange G).**
- **Somatotrophs produce growth hormone (GH or somatotropin)**, which e.g. stimulates liver cells to produce polypeptide growth factors which stimulate growth (e.g. somatomedin which stimulates epiphyseal cartilage - overproduction of this hormone may result in gigantism or acromegaly).
- **Mammotrophs (or lactotrophs)**, the second group of acidophils, secrete prolactin.
  - Their number increases significantly in late pregnancy and the early months of lactation.

### Basophil cells (or basophils)

Based on their hormone products basophils are divided into three subtypes.

- **Thyrotrophs produce thyroid stimulating hormone (TSH or thyrotropin).**
- **Gonadotrophs**
  - **produce follicle stimulating hormone (FSH), which stimulates the seminiferous epithelium in males in addition to early follicular growth in females.**
  - **Gonadotrophs also produce luteinizing hormone (LH), which stimulates production of testosterone by Leydig cells in males in addition to late follicular maturation, oestrogen secretion and formation of corpus luteum in females.**
- **Corticotrophs (or adrenocorticolipotrophs)**
  - **secrete adrenocorticotrophic hormone (ACTH or corticotropin) and lipotropin (LPH, no known function in humans).**
  - Corticotropes are the most frequent cell type in the pars intermedia.
  - In the pars intermedia, the precursor of ACTH and LPH undergoes further hydrolysis into melanocyte stimulating hormone (MSH, increased pigmentation in patients with Addison's disease) and a number of other peptides (among them endogenous opioids).

# The Posterior Pituitary



• **posterior pituitary, or neurohypophysis** = is the **neural portion of the pituitary**

- a **collection of unmyelinated axons**
  - axons **extend from cell bodies in hypothalamus**

• consists of:

- **pars nervosa,**
  - **infundibular stalk,**
  - **median eminence**
- **neurophysins** carry hormones made in the hypothalamus (ADH and oxytocin) from the hypothalamus to the posterior pituitary

• **embryological origin**

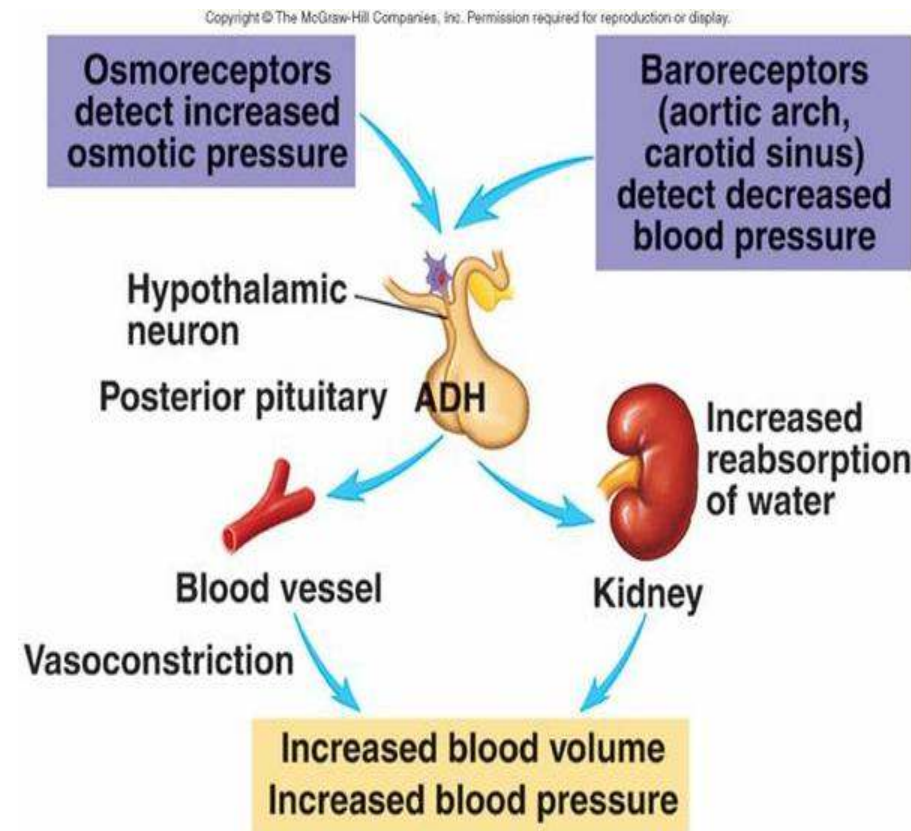
- **neural ectoderm**
  - **downgrowth of neural ectoderm (diencephalon)**

hormones are secreted by magnocellular neurons located in the supraoptic and paraventricular nucleus of hypothalamus

## Antidiuretic hormone (ADH; vasopressin)

- synthesis
  - hypothalamic supraoptic nucleus neuronal cell bodies synthesize ADH pro-hormone
    - **ADH pro-hormone contains ADH and neurophysin II**
  - ADH pro-hormones are packaged in secretory vesicles
    - secretory vesicles are transported via axonal transport to nerve terminals
      - **nerve terminals** in pars nervosa of posterior pituitary
  - ADH pro-hormone processing occurs in secretory vesicles during axonal transport
    - cleavage of neurophysin II and release of ADH hormone
- secretion
  - action potential depolarizes nerve terminals
    - neurosecretory vesicles fuse with plasma membrane
      - releases ADH and neurophysin II into perivascular space of highly fenestrated capillaries by which ADH enters systemic circulation

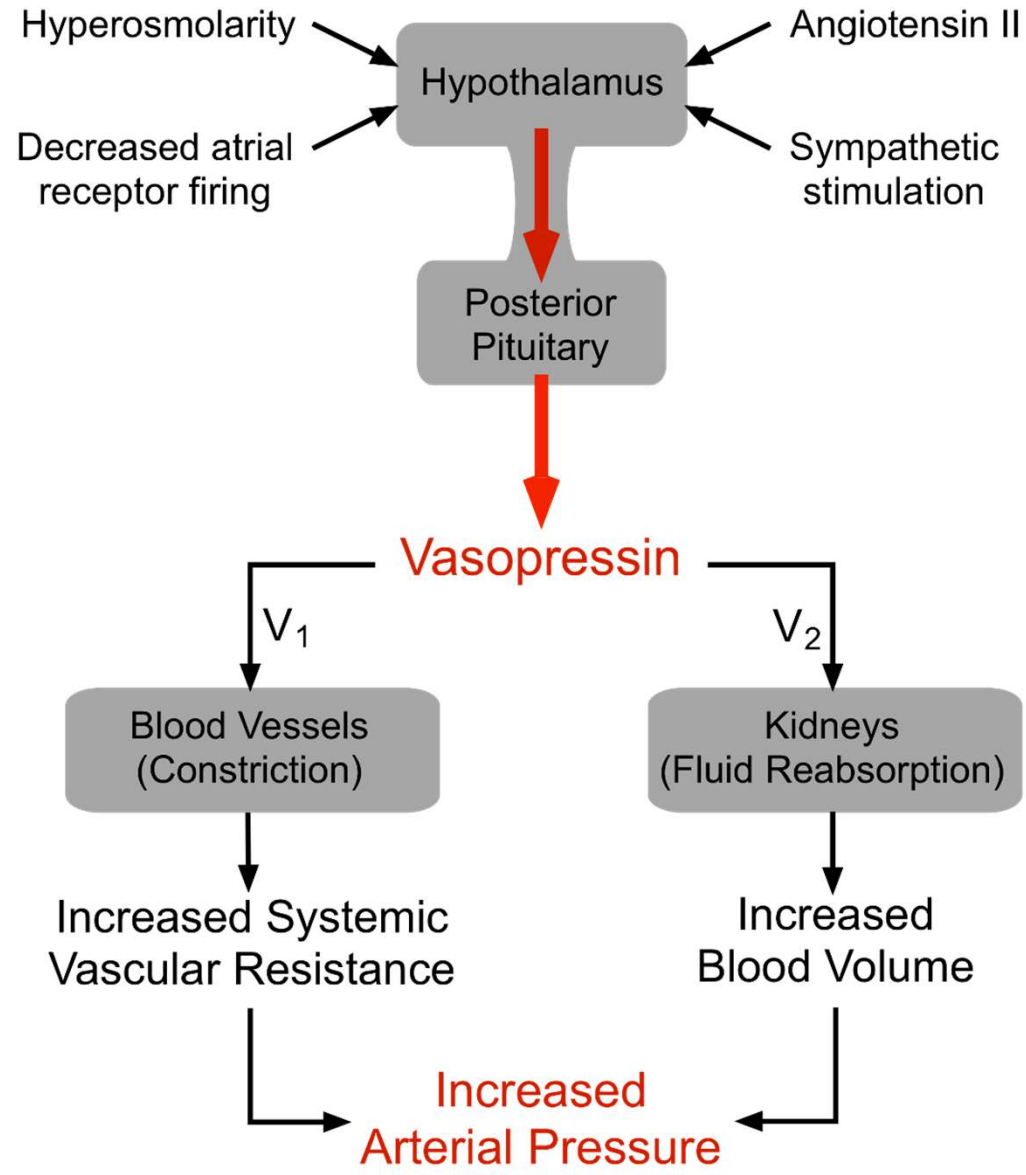
## The Posterior Pituitary

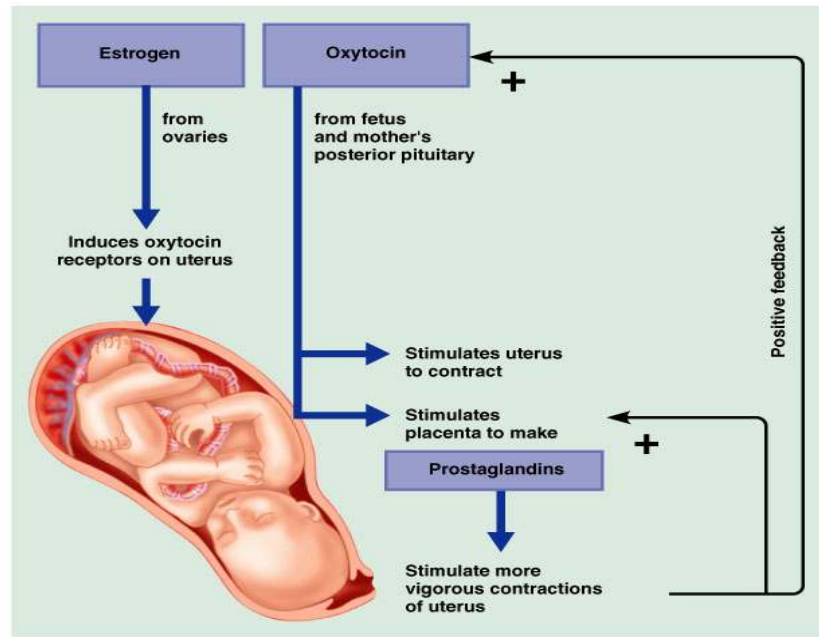
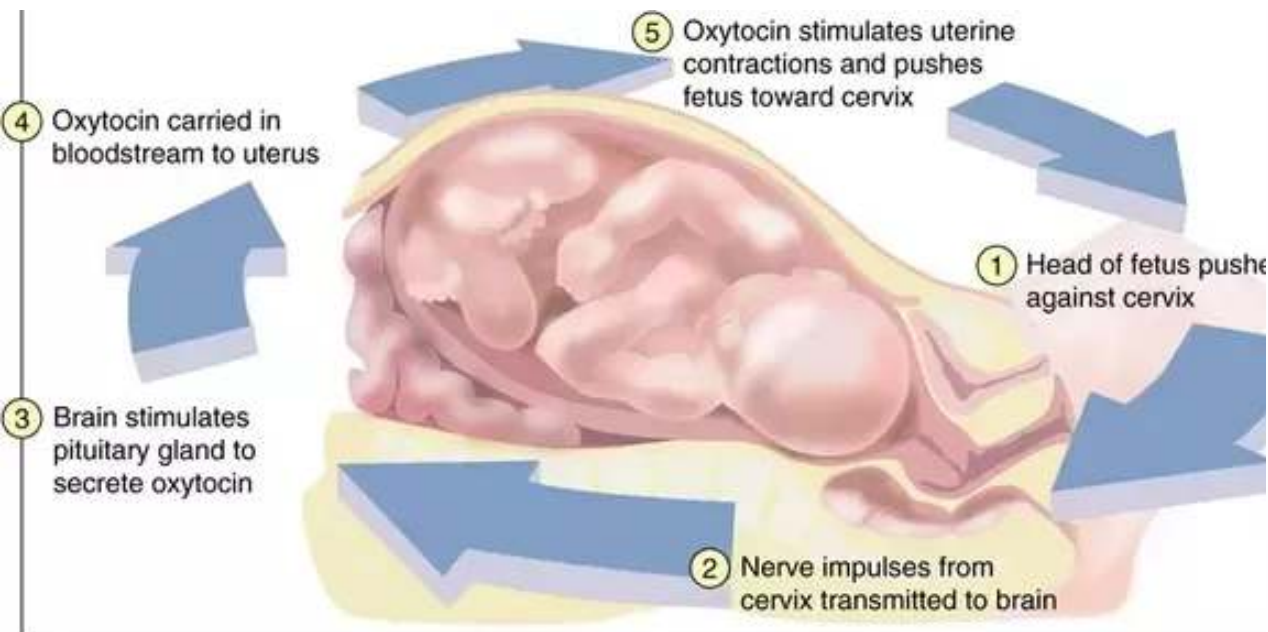


# POSTERIOR PITUITARY

- **OXYTOCIN (contracts uterine smooth muscle)**
- **VASOPRESSIN (ADH)**
  - **vasoconstriction,**
  - **gluconeogenesis,**
  - **platelet aggregation,**
  - **release of Factor-VIII and vWb factor,**
  - **concentrates urine, main effects on kidney and brain)**
- The posterior pituitary does not make these hormones, it just releases them.
- The hypothalamus actually makes the hormones and transfers it down the stalk to the neurohypophysis.







Copyright © 2004 Pearson Education, Inc., publishing as Benjamin Cummings.

## INCREASING OXYTOCIN

- LIGHT TOUCH
- SUCKING
- FOOD INTAKE
- WARMTH
- LIGHT PRESSURE
- MASSAGE-LIKE STROKING
- SEXUAL STIMULATION
- CALM & SUPPORTIVE ENVIRONMENT

Increases Oxytocin Release	Inhibits Oxytocin Release
<ul style="list-style-type: none"> <li>feeling secure</li> <li>privacy</li> <li>dim lighting</li> <li>people you know</li> <li>touch</li> <li>comfortable environment</li> <li>relaxation</li> </ul>	<ul style="list-style-type: none"> <li>feeling frightened</li> <li>feeling watched</li> <li>bright lights</li> <li>strangers</li> <li>questions</li> <li>uncomfortable environment</li> <li>thinking</li> </ul>

# BAHS\* of Posterior Pituitary Hormones

Hormone	Stimulates
Antidiuretic hormone	Water reabsorption in the kidney
Oxytocin	Contraction of uterine smooth muscle in labor. Contraction of breast cells to allow milk let down.

\* Boring as heck summary

# VIS\* of Oxytocin

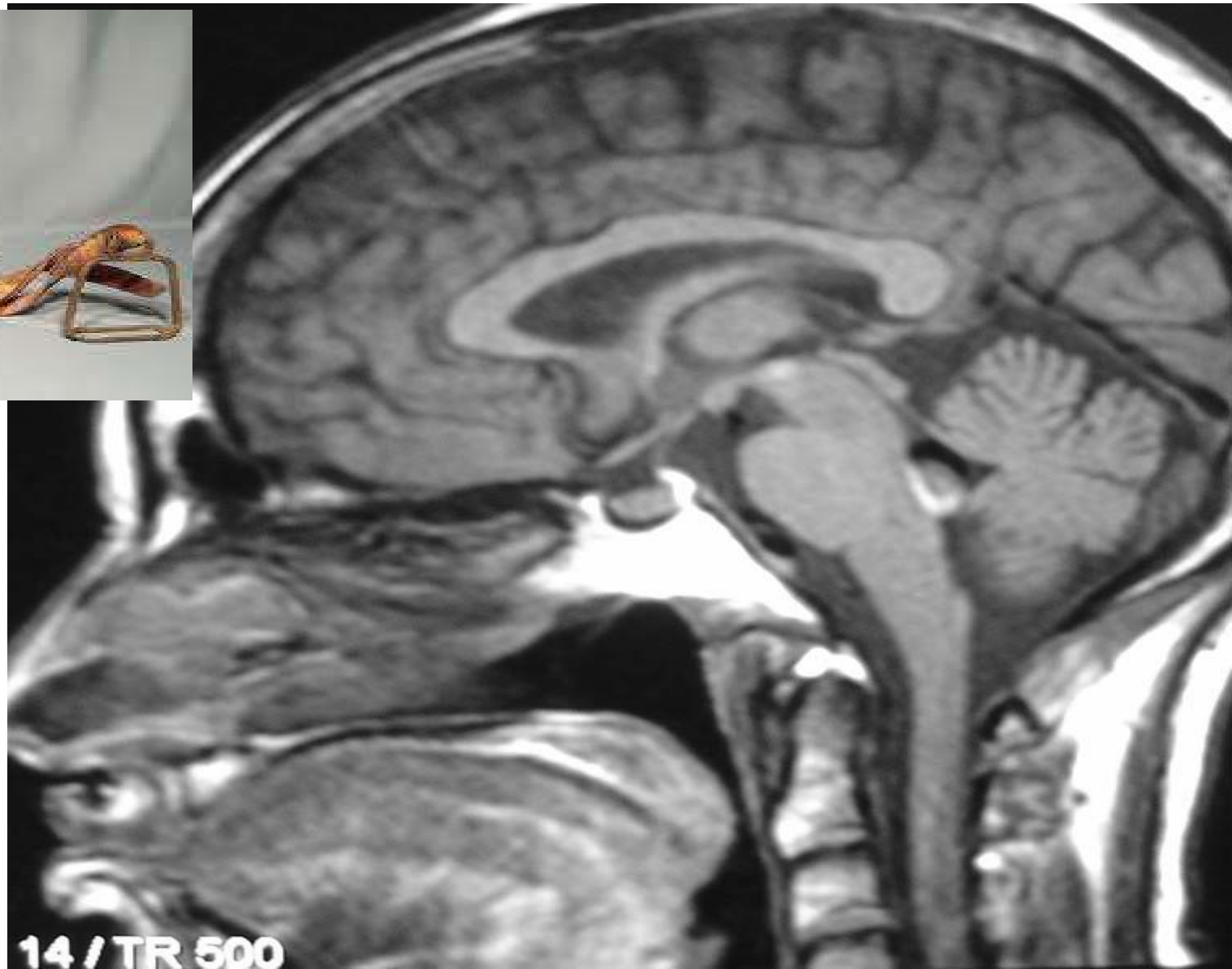
Situation	Stimulates
Interpersonal connection	Trust
Orgasm	Pleasure AND connection with that particular person
Intimate relationship	Monogamy
Sports teams	Better performance

\* Very interesting summary





Normal pituitary.  
With Turkish saddle,  
i.e., sella turcica.



# Pituitary Pathology

## □ Growth Hormone GH

- dwarfism -hyposecretion
- gigantism, acromegaly-hypersecretion

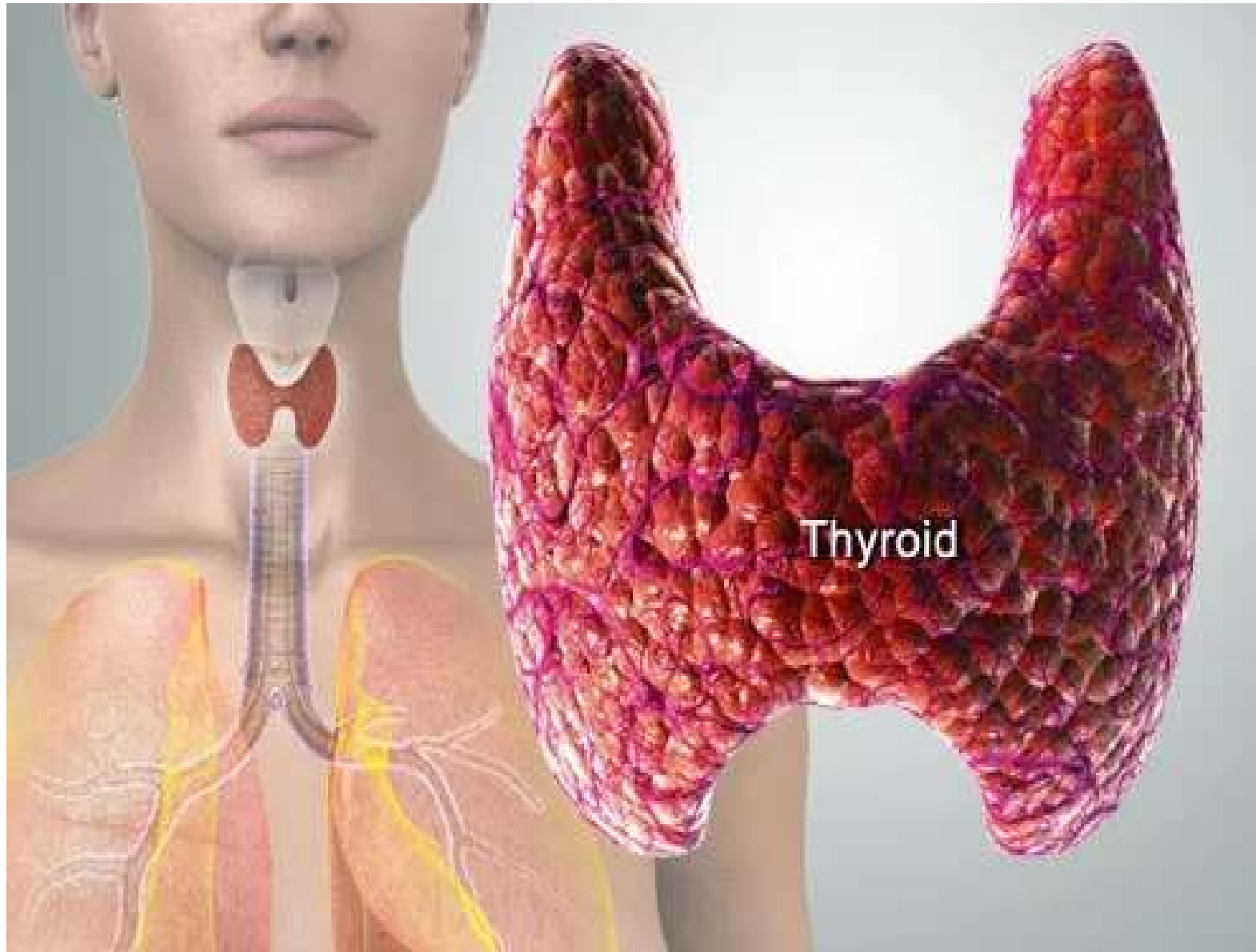
## □ Thyroid Stimulating Hormone TSH

- cretinism (infants) -hyposecretion
- myxedema( adults ) -hyposecretion
- Toxic goiter (adults -hypersecretion

## ■ exophthalmos



Thyroid



# What Does Thyroid Hormone Do?

- **Quick answer: increase growth and metabolism.**
- **More detailed answer:**
  - stimulate mitochondrial protein synthesis
  - increase absorption of carbohydrates
  - regulate fat metabolism
  - promote cell growth.
- **Bottom line: it increases basal metabolic rate and revs up most bodily functions (increases heart rate, raises body temperature, increases nervous reactivity, increases GI motility...the list goes on).**



## Iodine Metabolism

- i. Daily requirement of iodine is 150–200 mg/day.
  - Its sources are drinking water, fish, cereals, vegetables and iodinated salt.
- ii. Total body contains 25–30 mg of iodine.
  - All cells do contain iodine
  - but 80% of the total is stored in the thyroid gland.
  - Iodine level in blood is 5–10 µg/dL.
- iii. In most parts of the world, iodine is a scarce component of the soil.
  - Upper regions of mountains generally contain less iodine.
  - Such areas are called **goitrous belts**, e.g. **Himalayan region**.
- i v. Commercial source of iodine is seaweeds.

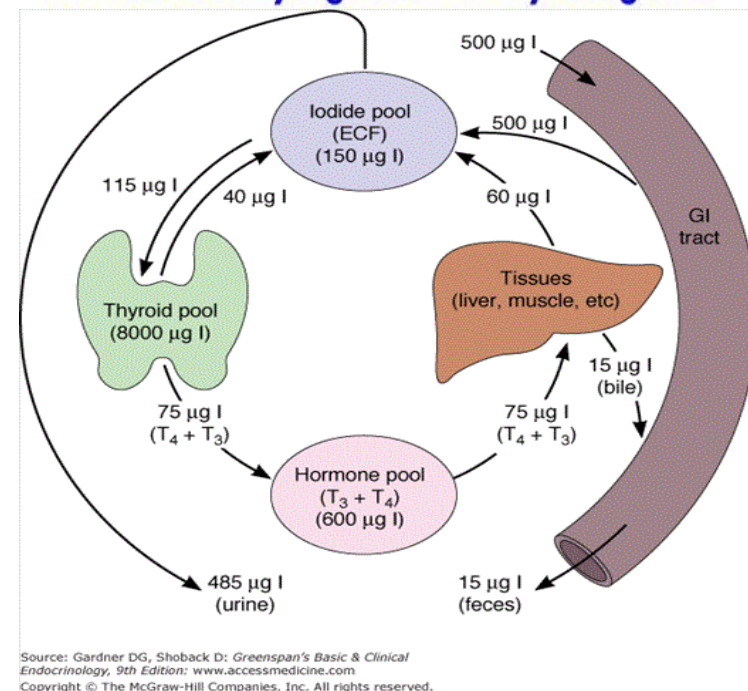
The program of iodination of common salt has resulted in increased availability of iodine.

v. Ingredients in foodstuffs, which prevent utilization of iodine are called **goitrogens**.

- Goitrogens are seen in cassava, maize, millet, bamboo shoots, sweet potatoes and beans.
- **Cabbage and tapioca contain thiocyanate, which inhibits iodine uptake by thyroid.**
- **Mustard seed contains thiourea, which inhibits iodination of thyroglobulin.**

vi. The only biological role of iodine is in formation of thyroid hormones, thyroxine (T<sub>4</sub>) and tri-iodo thyronine (T<sub>3</sub>).

- Iodine is absorbed from upper small intestine.
- Iodine is transported in plasma by loosely binding to plasma proteins.
- Iodine absorption also occurs through skin & lungs.
- 80% of body's iodine is stored in the organic form as **iodothyroglobulin** in thyroid gland.



# Thyroid Hormones

- **Thyroxine (T4) and Triiodothyronine (T3)-**
  - increases rate of energy release from carbohydrates
  - increases rate of protein synthesis
  - accelerates growth
  - stimulates activity in the nervous system
  - controlled by TSH
- **Calcitonin-**
  - **lowers blood calcium and phosphate ion concentrations by inhibiting release of calcium and phosphate from bones**
  - increases rate at which calcium and phosphate are deposited in bones

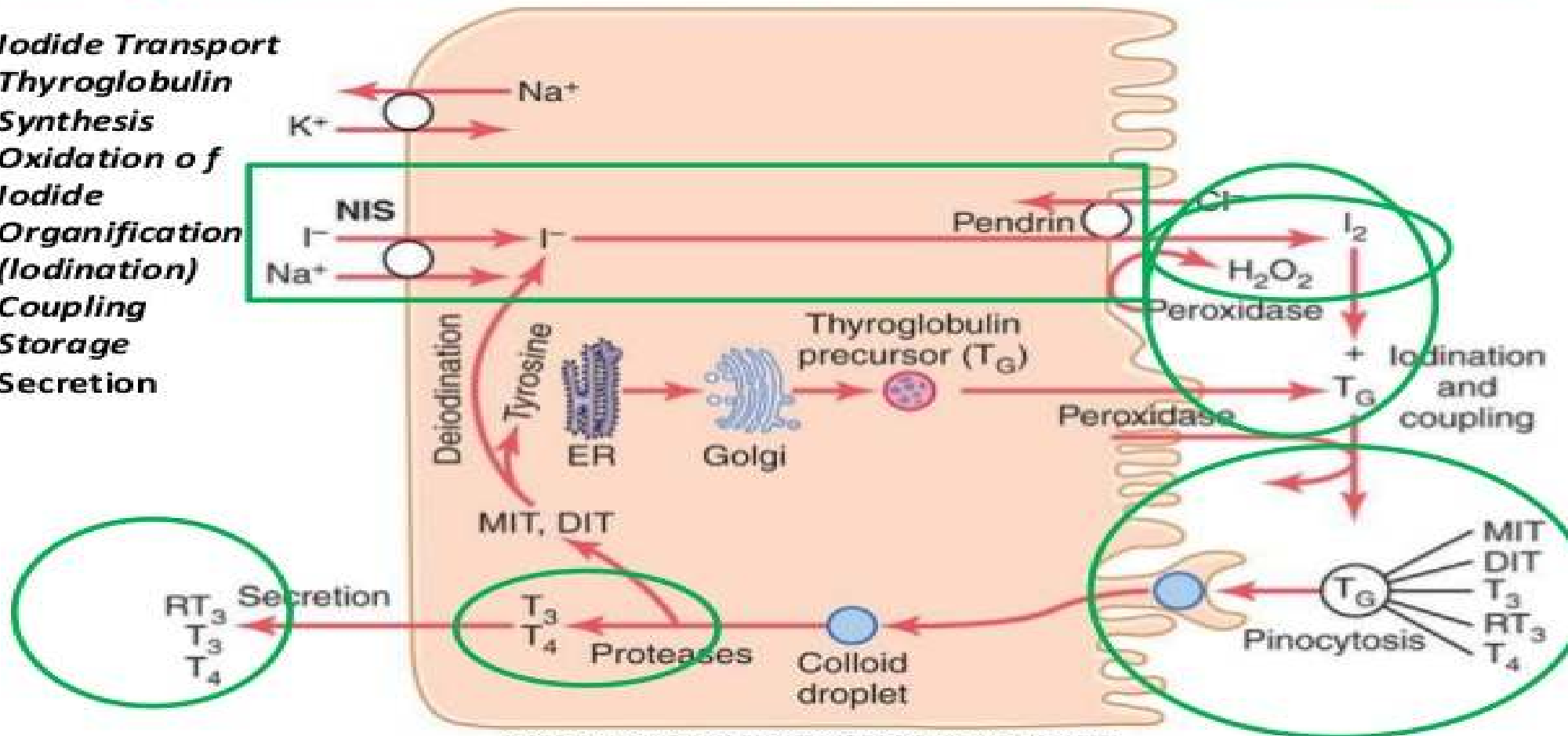
# Thyroid Gland

- Follicular cells synthesize **thyroglobulin** (a protein backbone) and secrete it into the colloid.
- Follicular cells take up iodide from the blood and attach it to tyrosine residues on thyroglobulin, forming T3 and T4 (thyroid hormones), which stay attached to thyroglobulin until needed.
- When stimulated by TSH, follicular cells eat a bit of colloid, digest it in a vesicle, cleave off the T3 and T4 and release it into the blood.

There are two groups of hormones derived from the amino acid tyrosine: Thyroid hormones are basically a "double" tyrosine with the critical incorporation of 3 or 4 iodine atoms. Catecholamines include epinephrine and norepinephrine, which are used as both hormones and neurotransmitters.

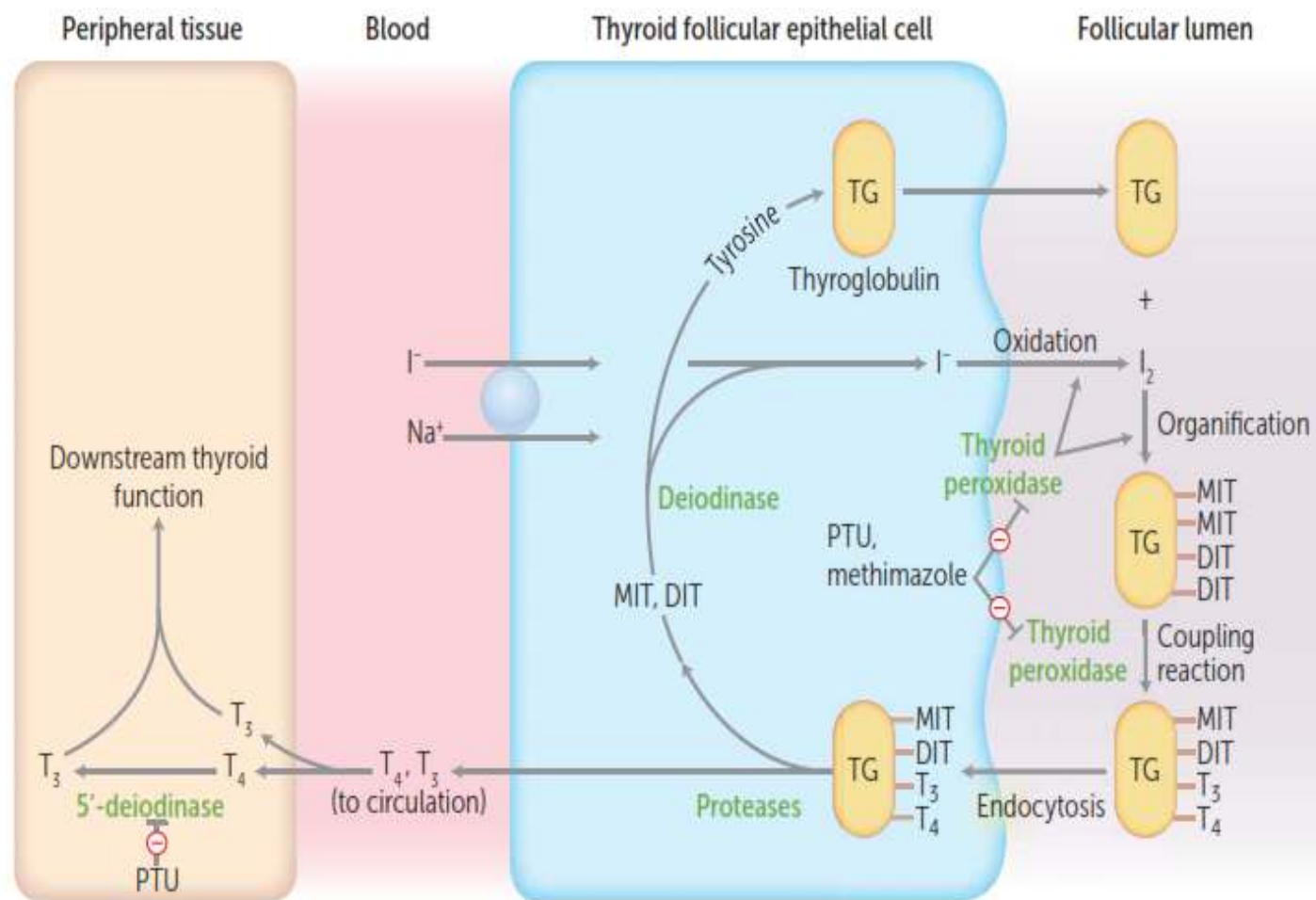
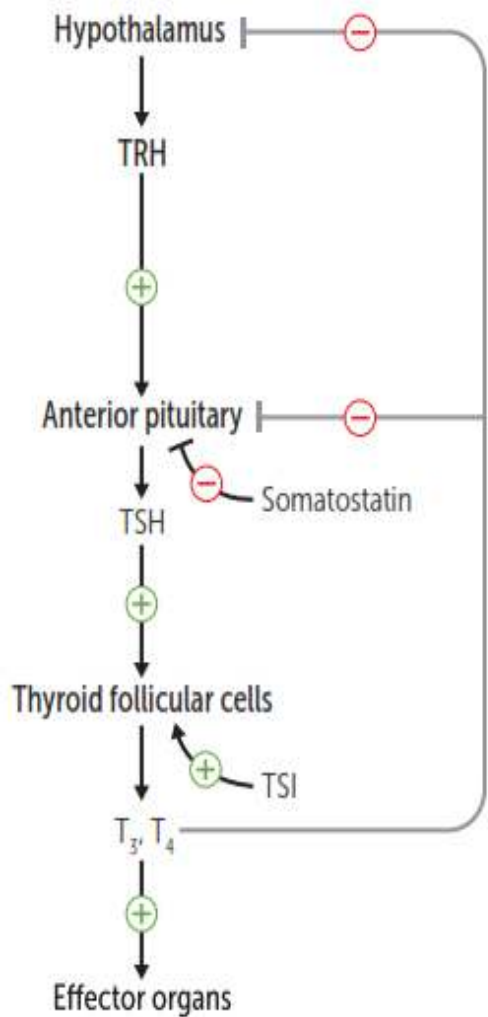
# Bio-synthesis and Secretion of Thyroid Hormone

1. Iodide Transport
2. Thyroglobulin Synthesis
3. Oxidation of Iodide
4. Organification (Iodination)
5. Coupling
6. Storage
7. Secretion



Hall: Guyton and Hall Textbook of Medical Physiology, 12th Edition  
 Copyright © 2011 by Saunders, an imprint of Elsevier, Inc. All rights reserved.

monoiodotyrosine (MIT) and diiodotyrosine (DIT)



monoiodotyrosine (MIT) and diiodotyrosine (DIT)



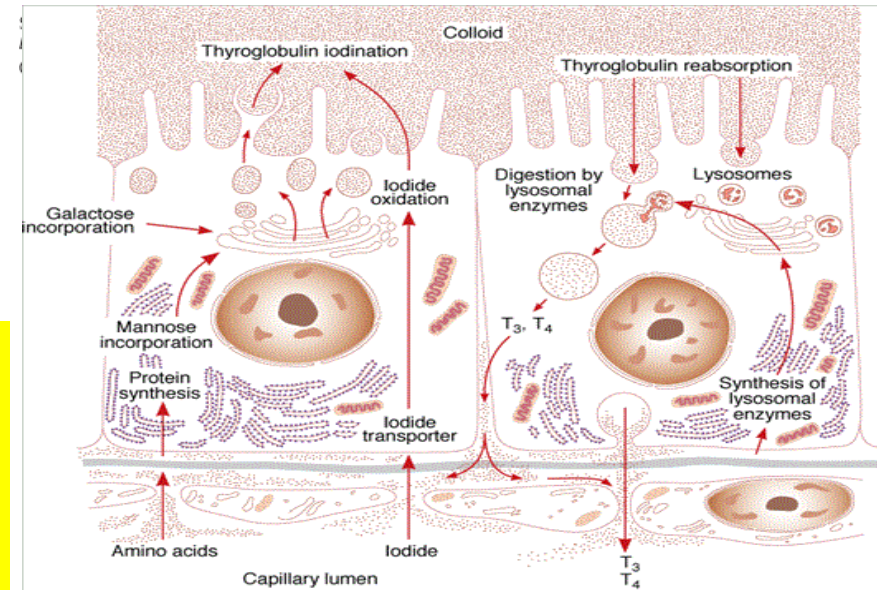
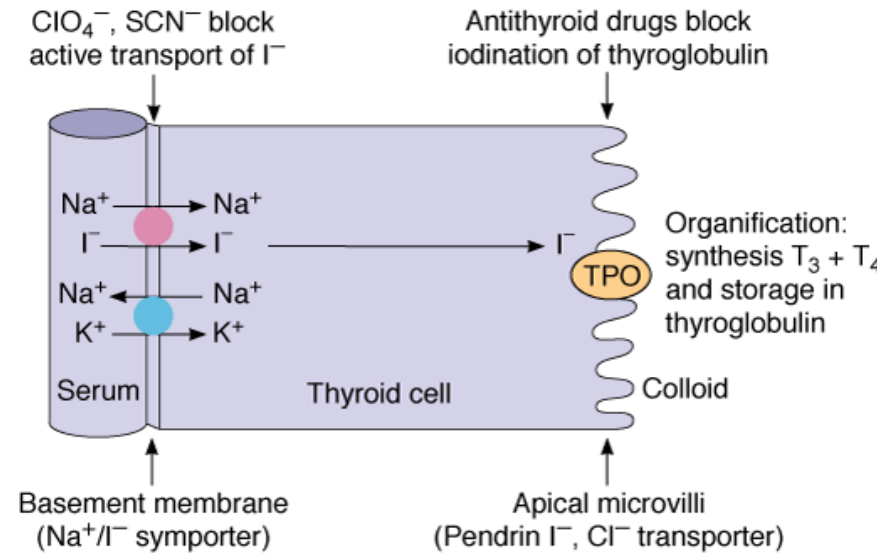
**Synthesis of T4 and T3 by the thyroid gland involves six major steps:**

- (1) active transport of iodide across the basement membrane into the thyroid cell (trapping)
- (2) oxidation of iodide and iodination of tyrosyl residues in thyroglobulin (organification)
- (3) linking pairs of iodotyrosine molecules within thyroglobulin to form the iodothyronines T3 and T4 (coupling)
- (4) pinocytosis and then proteolysis of thyroglobulin with release of free iodothyronines and iodotyrosines into the circulation
- (5) deiodination of iodotyrosines within the thyroid cell, with conservation and reuse of the liberated iodide
- (6) intrathyroidal 5'-deiodination of T4 to T3.

Thyroid hormone synthesis requires that NIS, thyroglobulin, and the enzyme thyroid peroxidase (TPO) all be present, functional, and uninhibited

The thiocarbamide drugs, including

- **methimazole,**
- **carbimazole,**
- **propylthiouracil (PTU)**
- **are competitive inhibitors of TPO. Their resulting ability to block thyroid hormone synthesis**



Source: Gardner DG, Shoback D: Greenspan's Basic & Clinical Endocrinology, 9th Edition: www.accessmedicine.com Copyright © The McGraw-Hill Companies, Inc. All rights reserved.

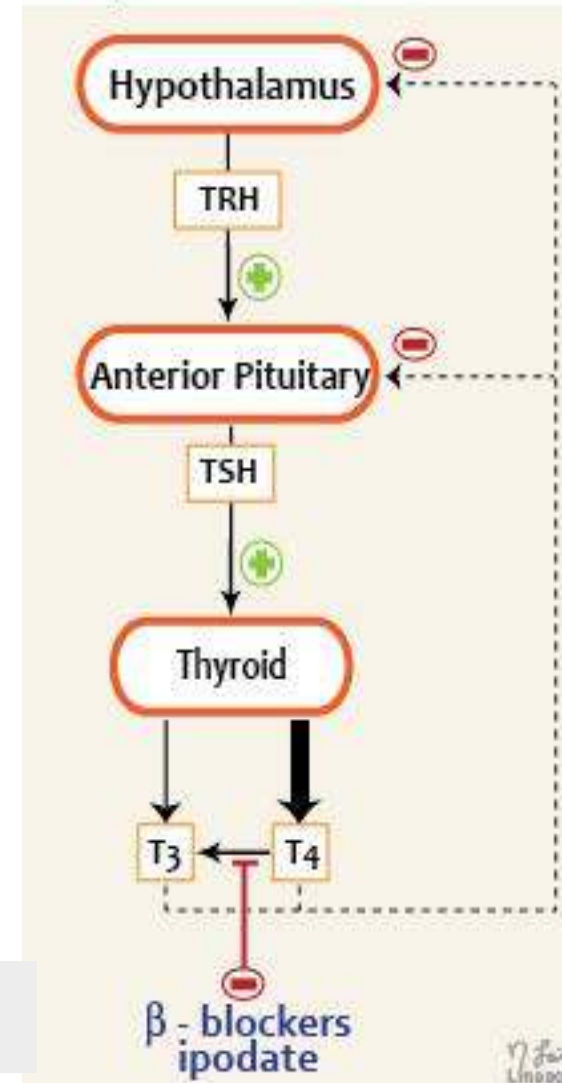
## •Synthesis

- created in the thyroid gland
- stored in thyroid follicles
- thyroid peroxidase** responsible for oxidation, organification, and coupling
  - forms  $I_2$  via oxidation of  $I^-$
  - forms thyroglobulin via organification of  $I_2$
- $T_4$  converted to  $T_3$  in peripheral tissues by outer ring deiodinase
- $T_4$  converted to  $rT_3$  by inner ring deiodinase

## •Regulation

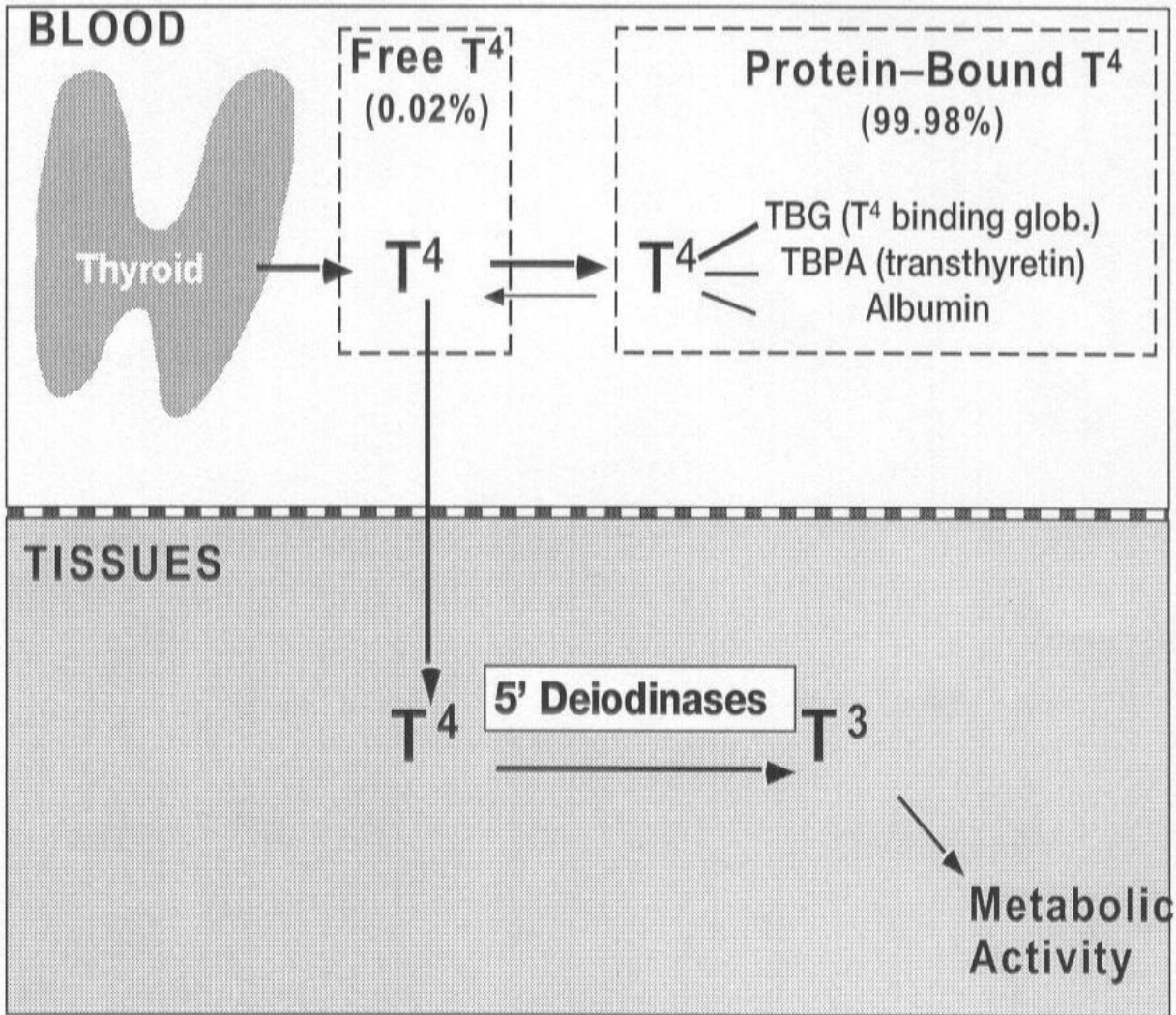
- TRH released from the hypothalamus to stimulates TSH release from the pituitary
  - TSH stimulates follicular cells to produce  $T_3$  and  $T_4$

# Thyroid Hormones



Abnormally low levels of  $T_4$  may indicate: dietary issues, such as fasting, malnutrition, or an iodine deficiency. medications that affect protein levels. hypothyroidism.

FIG 2. Thyroxine (T<sup>4</sup>) Distribution in the Circulation



**Function**

- bone growth
- CNS maturation
  - recall cretinism involves short stature and mental retardation
- increase the basal metabolic rate
  - via ↑ Na<sup>+</sup>/K<sup>+</sup>-ATPase activity
  - results in ↑ O<sub>2</sub> consumption, RR, and body temperature
- ↑ β<sub>1</sub> receptors in heart
  - results in ↑ CO, HR, SV, and contractility
  - recall the importance of treating hyperthyroidism with β-blockers
- ↑ glycogenolysis, gluconeogenesis, and lipolysis



**Thyroid hormones (T<sub>3</sub>/T<sub>4</sub>)**

Iodine-containing hormones that control the body's metabolic rate.

SOURCE	Follicles of thyroid. Most T <sub>3</sub> formed in target tissues.
FUNCTION	<p>Bone growth (synergism with GH)</p> <p>CNS maturation</p> <p>↑ β<sub>1</sub> receptors in heart = ↑ CO, HR, SV, contractility</p> <p>↑ basal metabolic rate via ↑ Na<sup>+</sup>/K<sup>+</sup>-ATPase activity → ↑ O<sub>2</sub> consumption, RR, body temperature</p> <p>↑ glycogenolysis, gluconeogenesis, lipolysis</p>
REGULATION	<p>TRH (hypothalamus) stimulates TSH (pituitary), which stimulates follicular cells. May also be stimulated by thyroid-stimulating immunoglobulin (TSI) in Graves disease. Negative feedback primarily by free T<sub>3</sub>/T<sub>4</sub> to anterior pituitary (↓ sensitivity to TRH) and hypothalamus (↓ TRH secretion).</p> <p>Wolff-Chaikoff effect—excess iodine temporarily inhibits thyroid peroxidase → ↓ iodine organification → ↓ T<sub>3</sub>/T<sub>4</sub> production.</p>

T<sub>3</sub> functions—**4 B's**:

- B**rain maturation
- B**one growth
- β**-adrenergic effects
- B**asal metabolic rate ↑

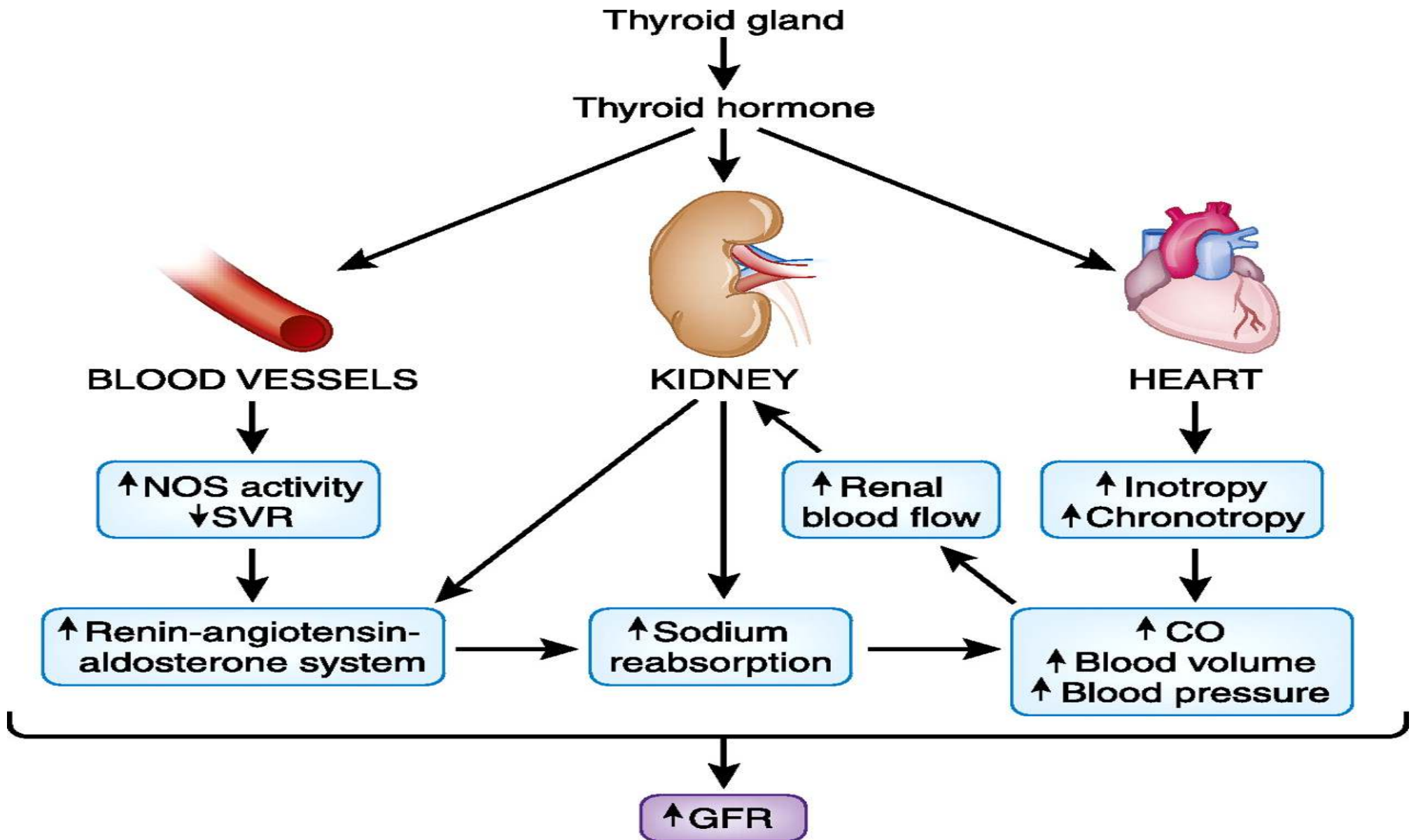
Thyroxine-binding globulin (TBG) binds most T<sub>3</sub>/T<sub>4</sub> in blood; only free hormone is active. ↓ TBG in hepatic failure, steroids; ↑ TBG in pregnancy or OCP use (estrogen ↑ TBG).

T<sub>4</sub> is major thyroid product; converted to T<sub>3</sub> in peripheral tissue by 5'-deiodinase.

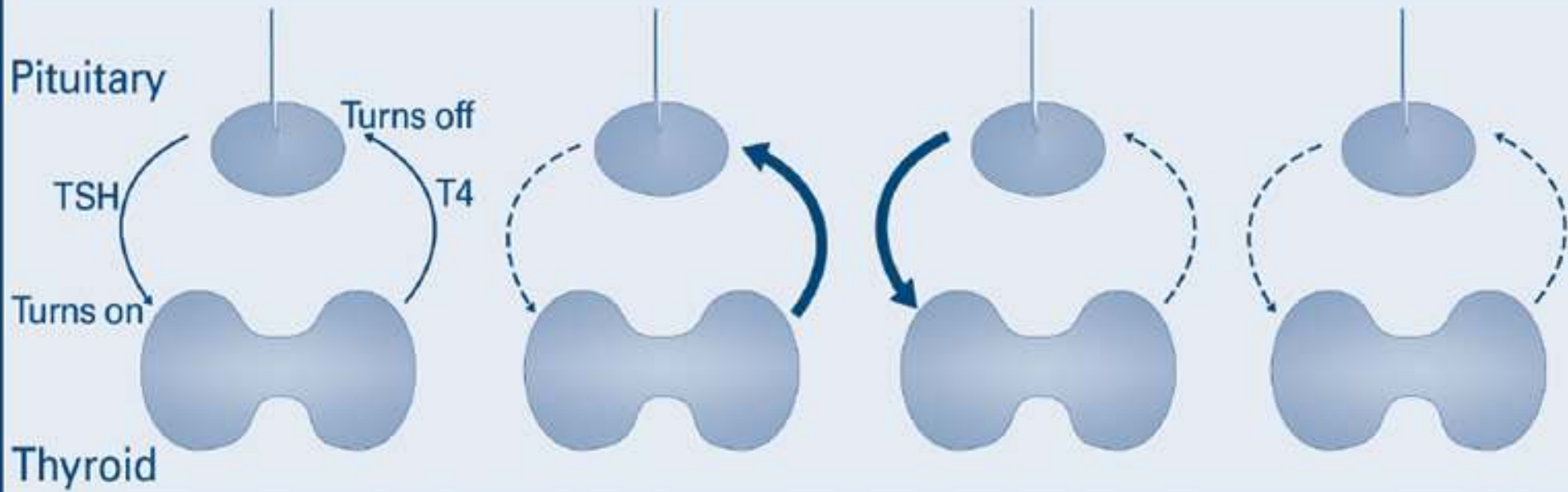
T<sub>3</sub> binds nuclear receptor with greater affinity than T<sub>4</sub>.

Thyroid peroxidase is the enzyme responsible for oxidation and organification of iodide as well as coupling of monoiodotyrosine (MIT) and di-iodotyrosine (DIT). DIT + DIT = T<sub>4</sub>. DIT + MIT = T<sub>3</sub>.

Propylthiouracil (PTU) inhibits both thyroid peroxidase and 5'-deiodinase. Methimazole inhibits thyroid peroxidase only. Glucocorticoids inhibit peripheral conversion of T<sub>4</sub> to T<sub>3</sub>.







**CONDITION: Normal**

**Hyperthyroidism**

**Hypothyroidism  
Primary**

**Hypothyroidism  
Secondary**

**TSH**      Normal

Low

High

Low

**T4**      Normal

High

Low

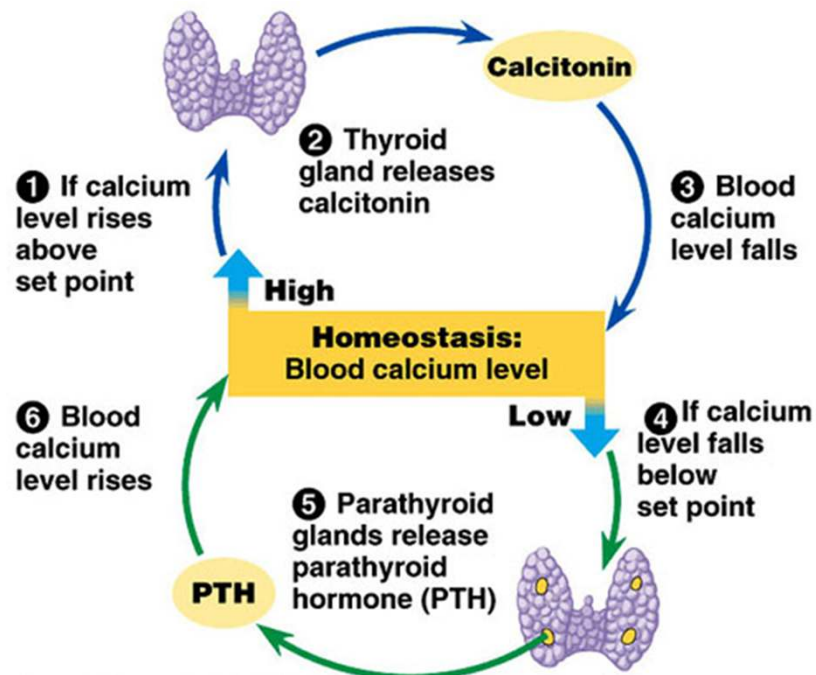
Low

# Parafollicular Cells (C Cells)

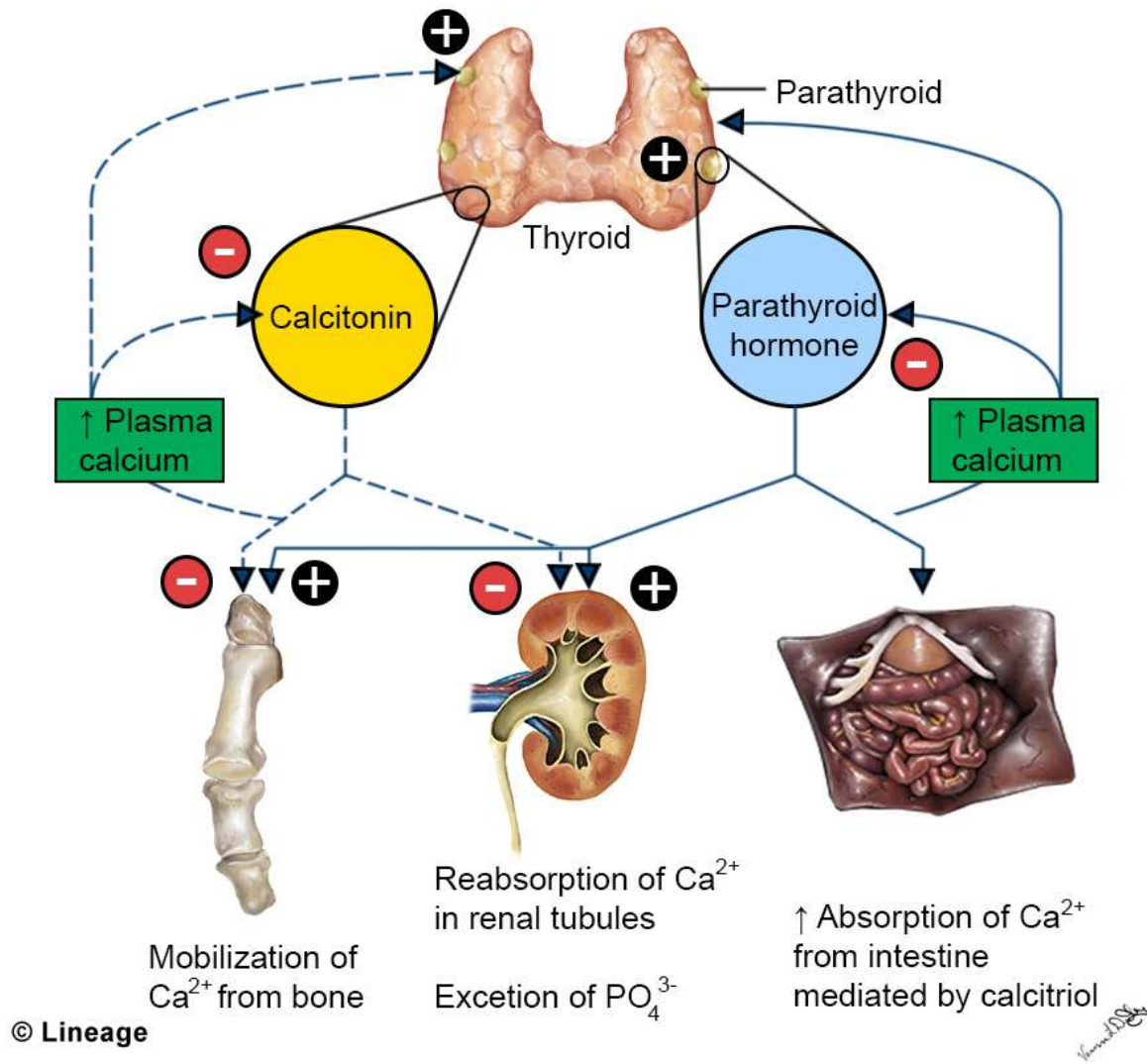
- Derived from neural crest ectoderm.
- Located between follicular cells and between follicles.
- Parafollicular cells are larger cells with clear cytoplasm and small secretory granules containing calcitonin.
- Calcitonin is made in response to high blood calcium (it's not affected by a pituitary hormone!).
- Calcitonin lowers blood calcium levels by inhibiting osteoclastic resorption.

## Calcitonin

SOURCE	Parafollicular cells (C cells) of thyroid.	Calcitonin opposes actions of PTH. Not important in normal $\text{Ca}^{2+}$ homeostasis. Calcitonin <b>tones</b> down serum $\text{Ca}^{2+}$ levels and keeps it in <b>bones</b> .
FUNCTION	$\downarrow$ bone resorption of $\text{Ca}^{2+}$ .	
REGULATION	$\uparrow$ serum $\text{Ca}^{2+} \rightarrow$ calcitonin secretion.	

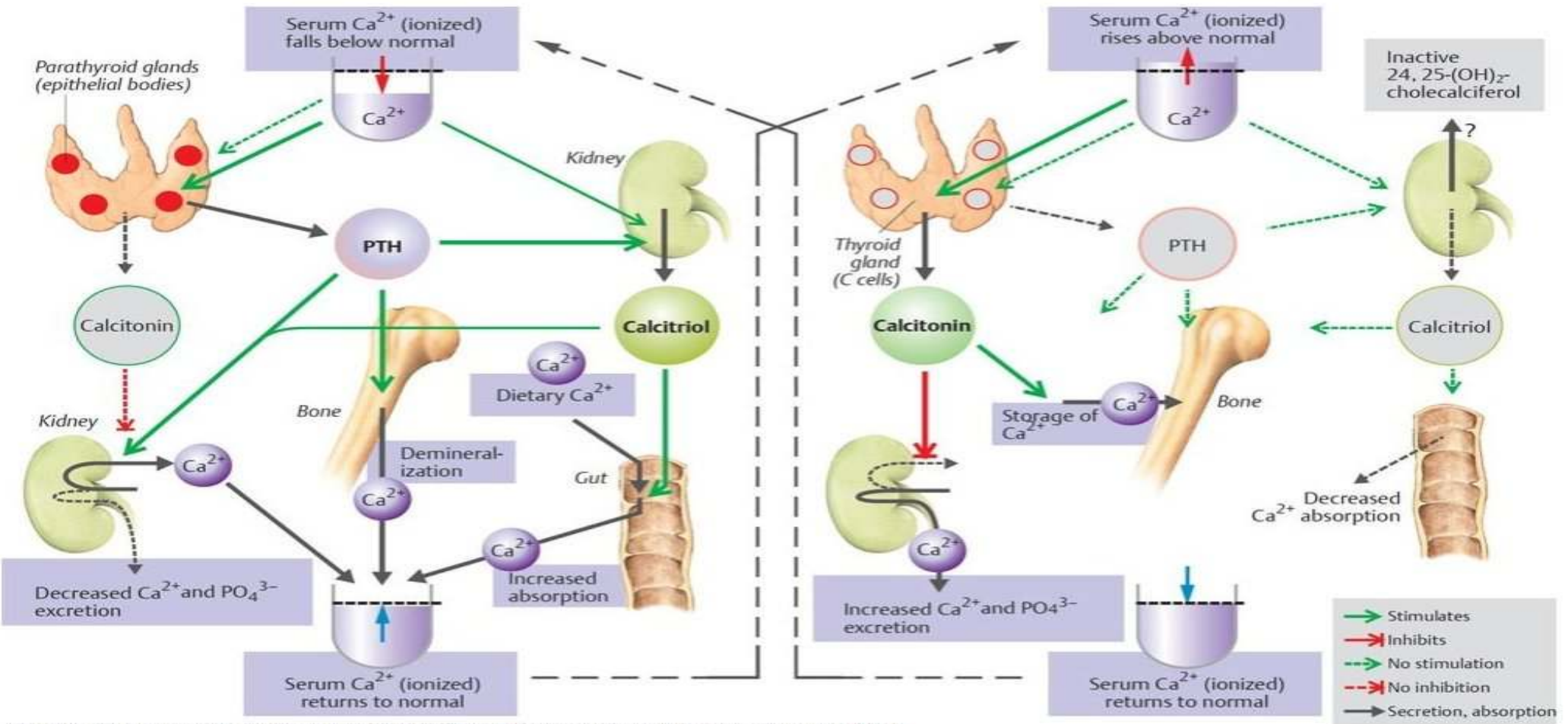


Copyright © Pearson Education, Inc., publishing as Benjamin Cummings.



### Hormonal regulation of the blood $\text{Ca}^{2+}$ concentration.

$\text{Ca}^{2+}$  homeostasis is achieved by three main hormones: parathyroid hormone (PTH, from parathyroid gland), calcitonin (from parafollicular cells of the thyroid gland), and calcitriol (mainly produced in the kidney). In low serum  $\text{Ca}^{2+}$  states, the actions of parathyroid hormone and calcitriol predominate, causing increased  $\text{Ca}^{2+}$  uptake from the gut and bone and decreased renal excretion. In high serum  $\text{Ca}^{2+}$  states, the action of calcitonin predominates, causing decreased  $\text{Ca}^{2+}$  uptake from the gut, increased renal excretion, and storage of excess  $\text{Ca}^{2+}$  in bone.

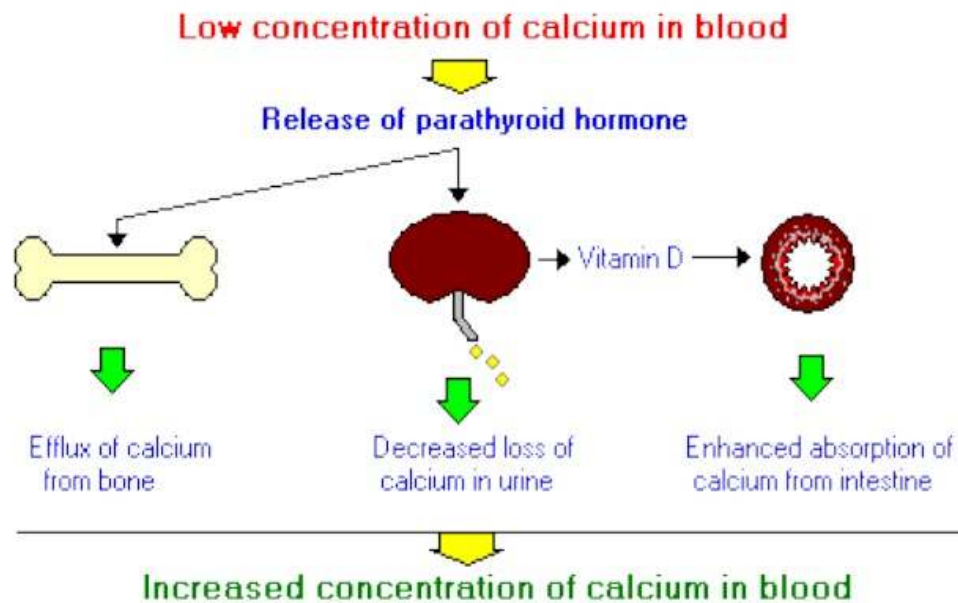


Source : Pharmacology - An Illustrated Review (Thieme Illustrated Review Series) - Simmons, Mark



	<b>PTH</b>	<b>Vitamin D</b>	<b>Calcitonin</b>
<b>Bone:</b>	↑ resorption.	↑ resorption & formation.	↓ resorption.
<b>Kidney:</b>	↑ tubular $\text{Ca}^{++}$ reabsorption. ↑ tubular $\text{PO}_4^-$ excretion.	↑ tubular $\text{Ca}^{++}$ & $\text{PO}_4^-$ reabsorptn	↓ tubular $\text{Ca}^{++}$ & $\text{PO}_4^-$ reabsorptn.
<b>G.I.T.</b>	Indirect through calcitriol (↑ $\text{Ca}^{++}$ & $\text{PO}_4^-$ reabsorption).	↑ $\text{Ca}^{++}$ & $\text{PO}_4^-$ reabsorption.	
<b>Serum <math>\text{Ca}^{++}</math></b>	↑	↑	↓
<b><math>\text{PO}_4^-</math></b>	↓	↑	↓

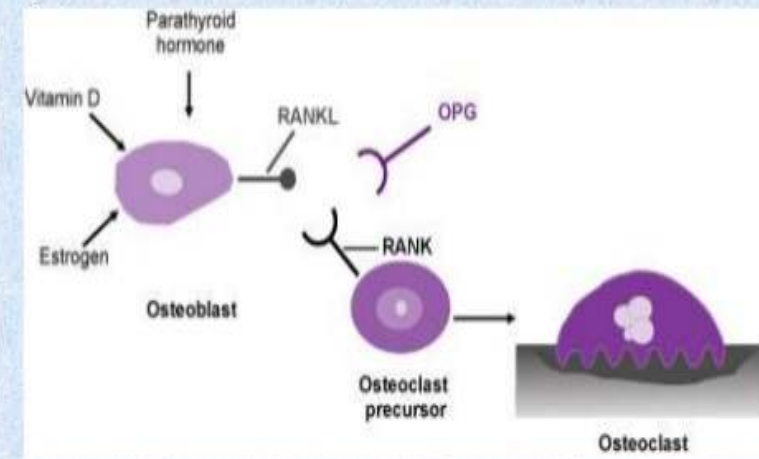
# Parathyroid



PTH binds to **osteoblasts**.

Osteoblasts **increase expression of RANK-L** and inhibits their expression of Osteoprotegerin (OPG). (*OPG blocks RANK-L*)

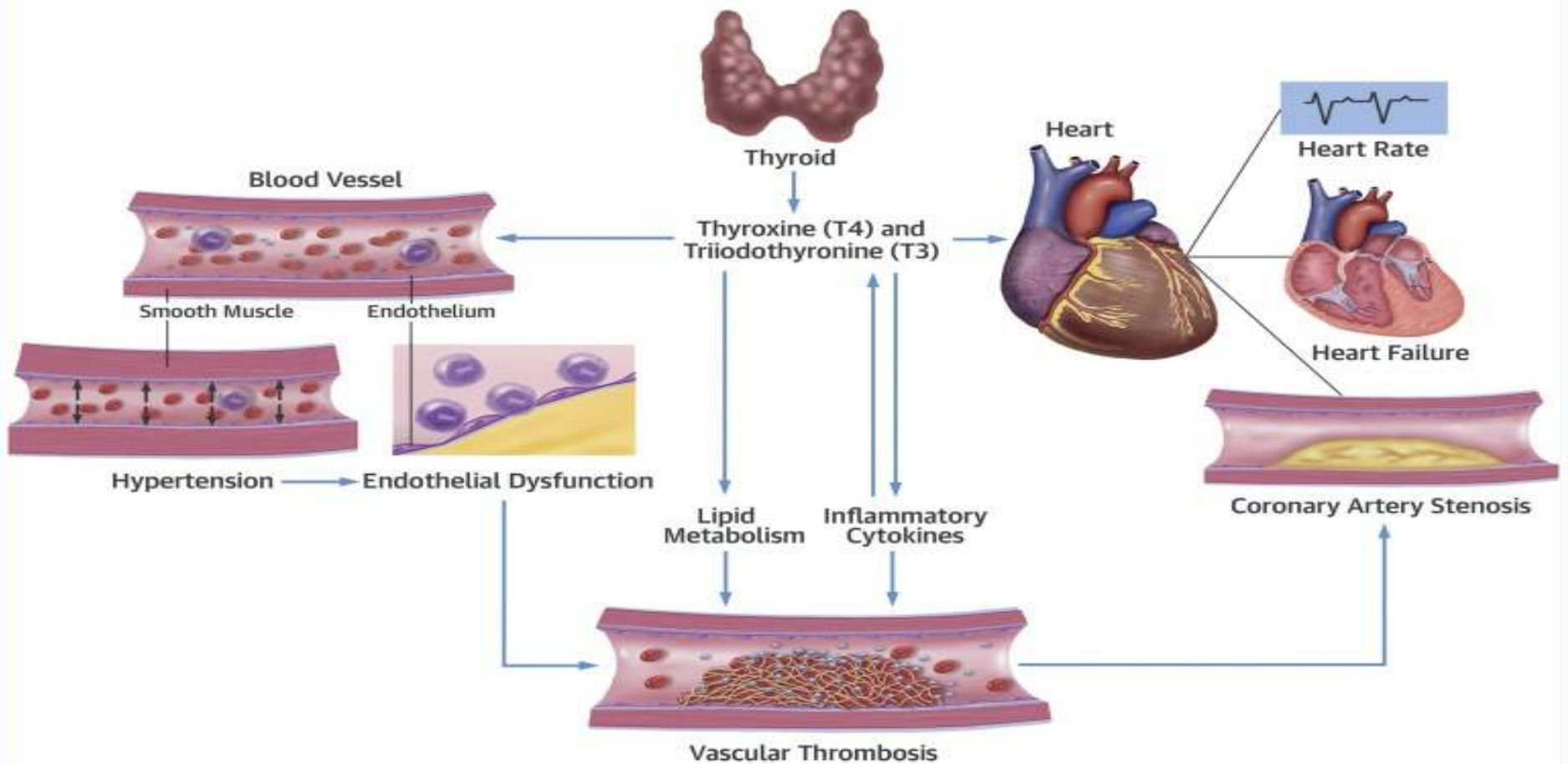
RANK-L **binds RANK** on **osteoclast precursors**, and they form new osteoclasts. And osteoclasts **enhance bone resorption** thus increasing Blood Calcium and Decreasing Bone Calcium



# Heart

- The natriuretic peptide family consists of three biologically active peptides: (will be discussing this in cardiovascular)
  - **atrial natriuretic peptide (ANP),**
  - **brain (or B-type) natriuretic peptide (BNP),**
  - **and C-type natriuretic peptide (CNP).**
- Among these, ANP and BNP are secreted by the heart and act as cardiac hormones.

## CENTRAL ILLUSTRATION: The Interactions Between Thyroid Hormones and the Cardiovascular System



Razvi, S. et al. J Am Coll Cardiol. 2018;71(16):1781-96.



# Pancreas

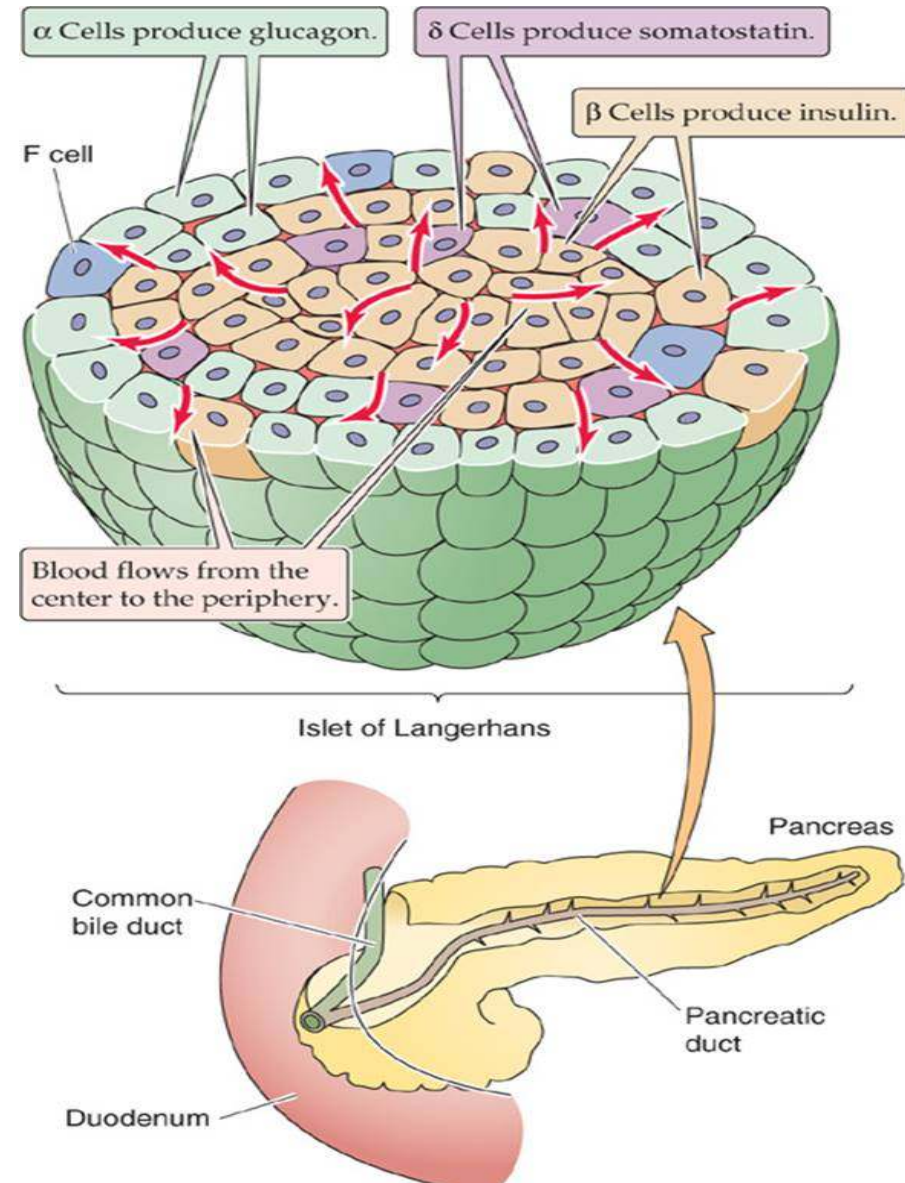
- Pancreas is having both Endocrine and Exocrine parts.
- The Endocrine part is made up of Islet of Langerhans which are aggregated in tail part of pancreas.
- Islet of Langerhans is an encapsulated structure bounded by thin capsule of reticular fibres.
- In the islet, following three different types of cells are mainly found. Alpha, Beta and D cells.

Beta ( $\beta$ ) cells produce INSULIN

• Alpha ( $\alpha$ ) cells produce GLUCAGON

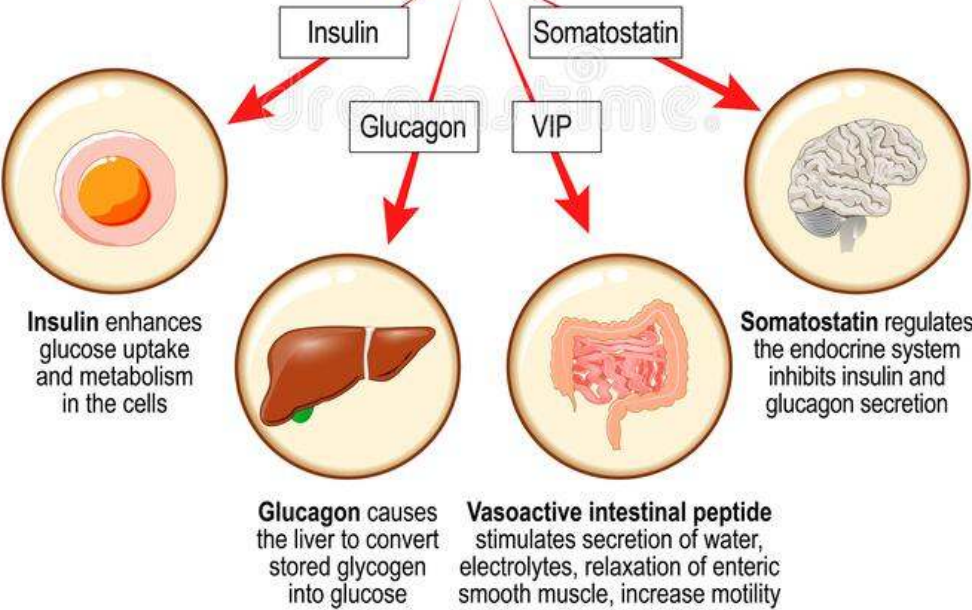
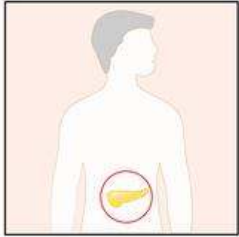
• Delta ( $\delta$ ) cells produce SOMATOSTATIN

• F cells produce PANCREATIC POLYPEPTIDE





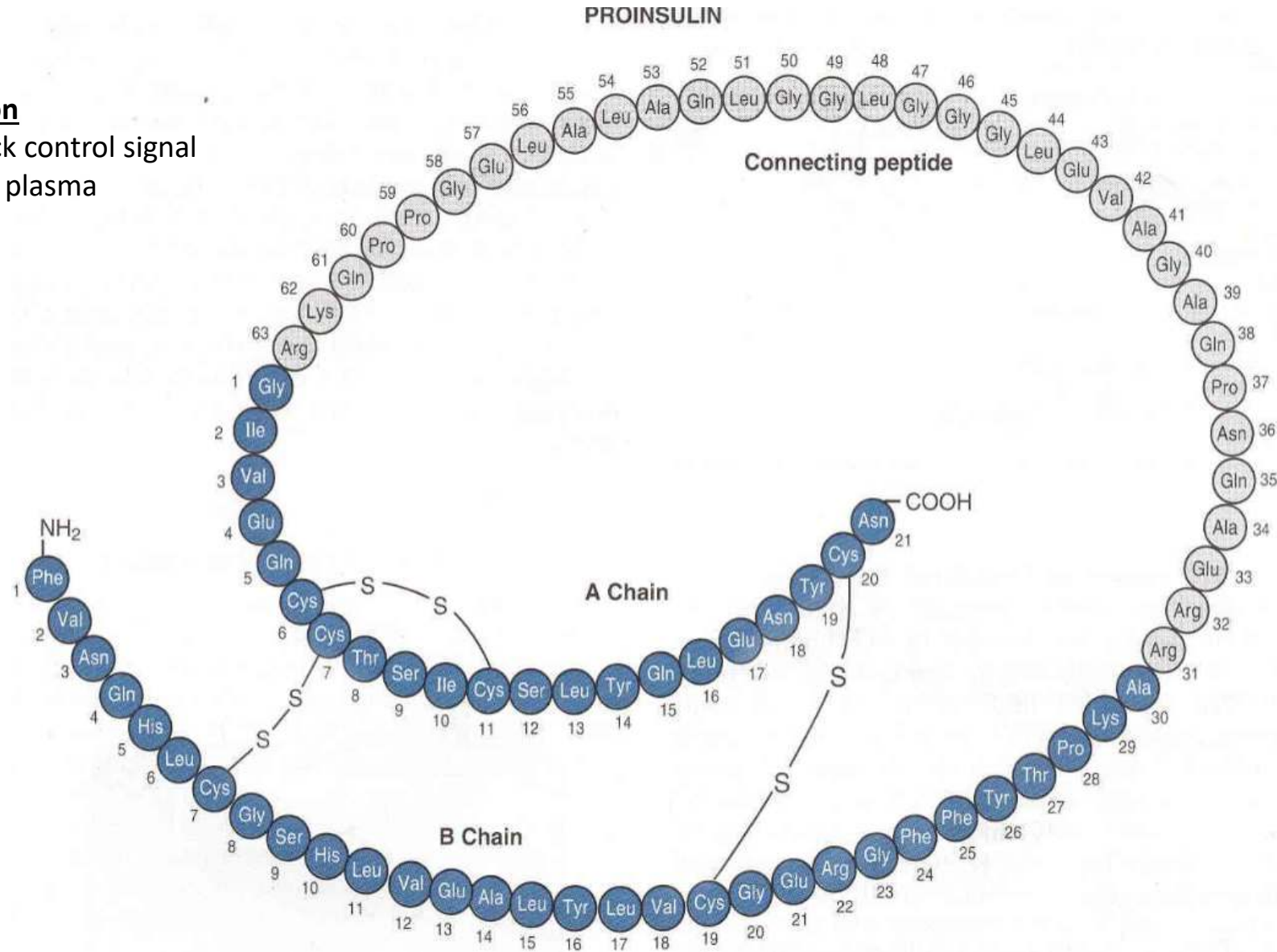
# Pancreas



# Insulin

## Regulation of insulin secretion

Mainly regulated by feed back control signal provided by nutrients level in plasma  
“Hormone of Abundance”



## Regulators of insulin secretion

### Stimulators of insulin secretion

↑ Serum glucose

↑ Serum amino acids

↑ Serum free fatty acids

↑ Serum ketone bodies

#### Hormones

Gastroinhibitory peptide (GIP)

Glucagon

Gastrin

Cholecystokinin (CCK)

Secretin

Vasoactive intestinal peptide (VIP)

Epinephrine ( $\beta$ -receptor)

Parasympathetic nervous system

### Inhibitors of insulin secretion

↓ Glucose

↓ Amino acids

↓ Free fatty acids

#### Hormones

Somatostatin

Epinephrine ( $\alpha$ -receptor)

Sympathetic nervous system stimulation

## INSULIN ACTION ON CARBOHYDRATE METABOLISM

### LIVER

- Stimulates glucose oxidation
- Promotes glucose storage as glycogen
- Inhibits glycogenolysis
- Inhibits gluconeogenesis

### MUSCLE

- Stimulates glucose uptake (GLUT4)
- Promotes glucose storage as glycogen

facilitates amino acids entry into muscle cells

- Facilitates protein synthesis in ribosomes by induction of gene transcription
- Inhibits proteolysis by decreasing lysosomal activity

**“ANABOLIC HORMONE”**

### ADIPOSE TISSUE

- Stimulates glucose transport into adipocytes
- Promotes the conversion of glucose into triglycerides and fatty acids

**“ANTI-DIABETOGENIC”**

## INSULIN ACTION ON FAT METABOLISM

### LIVER

- Anti ketogenic & Lipogenic
- Stimulates HMG-CoA reductase

### ADIPOSE TISSUE

- Promotes storage of fat
- Inhibits lipolysis by inhibiting Hormone sensitive lipase
- Promotes lipogenesis by stimulating lipoprotein lipase

“ANTI-KETOGENIC”

The absorptive state, or the fed state, occurs after a meal when your body is digesting the food and absorbing the nutrients (catabolism exceeds anabolism). Digestion begins the moment you put food into your mouth, as the food is broken down into its constituent parts to be absorbed through the intestine.

## INSULIN ACTION ON PLASMA K<sup>+</sup> CONCENTRATION

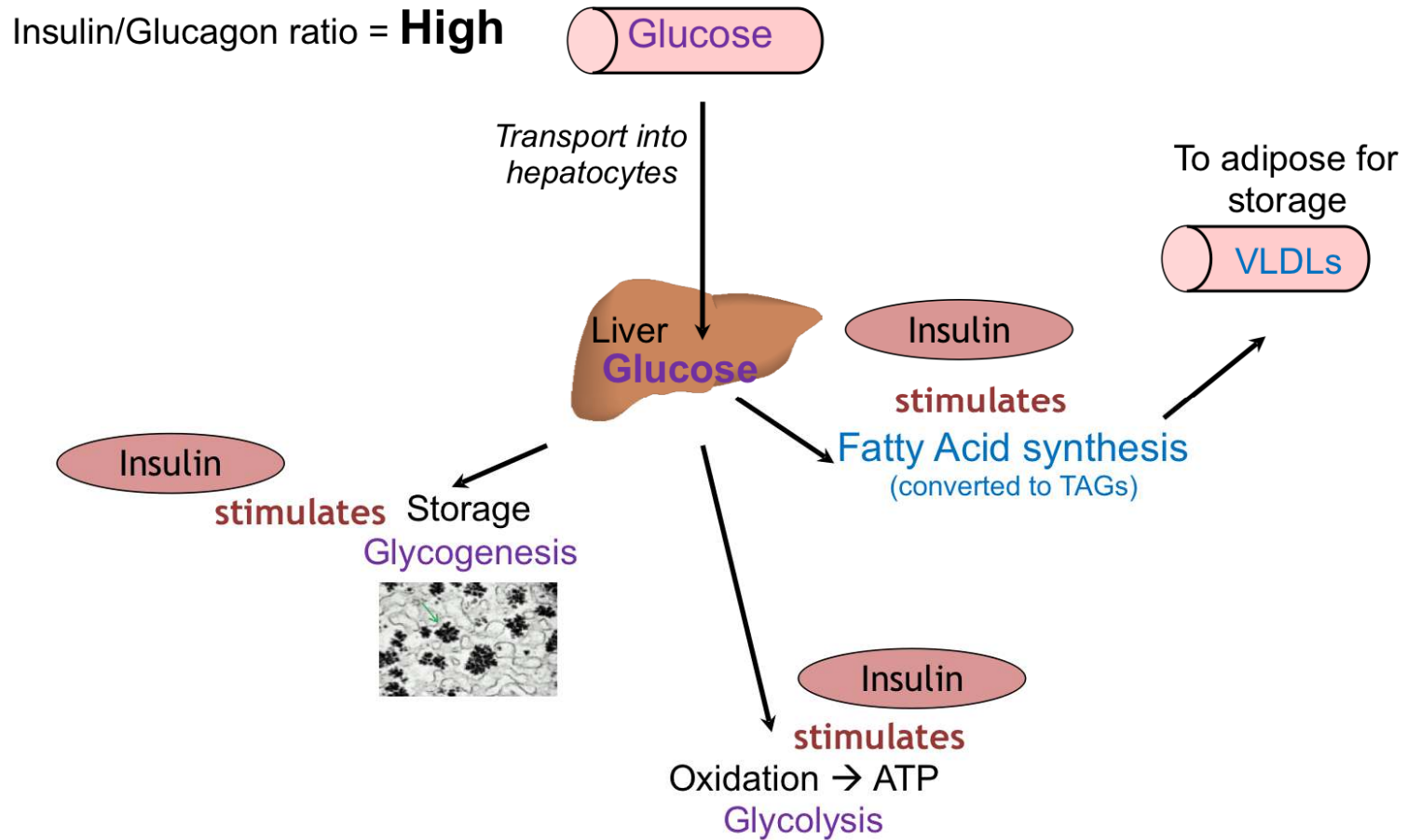
- Facilitates rapid entry of K<sup>+</sup> into cell by simulating Na-K ATPase activity
  - Thus decreases plasma concentration of K<sup>+</sup>
  - APPLIED: Insulin is given along with glucose in the treatment of Hyperkalemia that occurs in Acute Renal Failure
- “PHYSIOLOGICAL REGULATOR OF PLASMA K<sup>+</sup> CONCENTRATION”

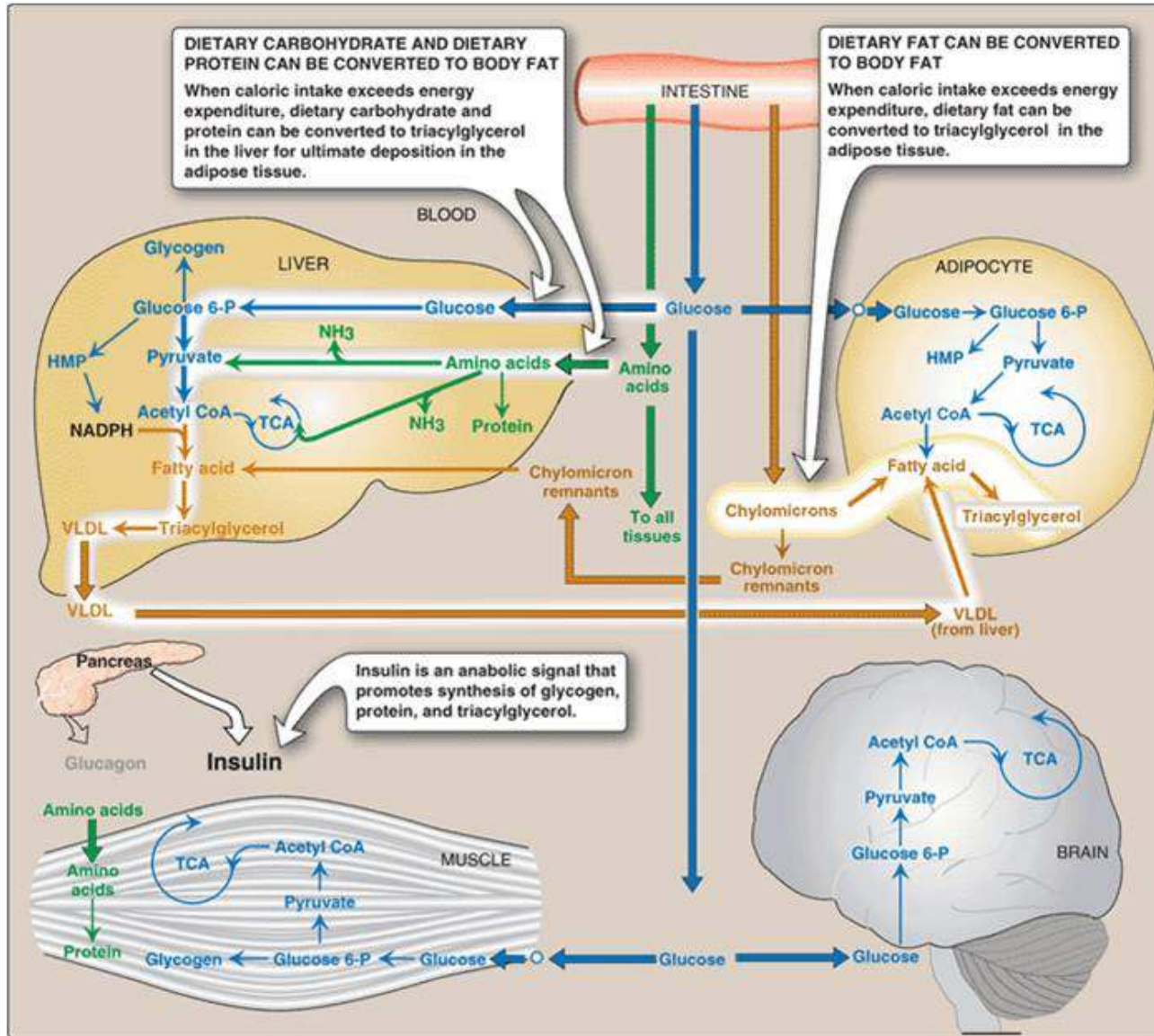
### Dominates in Fed State Metabolism

- INCREASE GLUCOSE UPTAKE IN MOST CELLS => Anti-Diabetogenic
- INCREASE GLUCOSE USE & STORAGE=> Anabolic
- INCREASE PROTEIN SYNTHESIS=> Anti-ketogenic
- INCREASE FAT SYNTHESIS=> Lipogenic



# Metabolism in Fed State: Liver





## GLUCAGON

Produced by alpha cells in the pancreas

- Its major target is the liver, where it promotes:
  - **Glycogenolysis** – the breakdown of glycogen to glucose
  - **Gluconeogenesis** – synthesis of glucose from lactic acid and non carbohydrates
  - **Release of glucose to the blood from liver cells**

Stimulates glycogenolysis, gluconeogenesis & inhibits glycogenesis

- Promotes lipolysis & ketogenesis
- Increases calorogenesis

“**Prodiabetogenic and Ketogenic**”

## INSULIN-GLUCAGON RATIO

- **Insulin** is hormone of **energy storage**
- **Glucagon** is hormone of **energy release**
- A balance should be maintained for normal metabolic functions
- After a normal balance diet is 3
- After overnight fasting decreases to 1, may decrease to as low as 0.4 after prolonged fasting
- Physiological significance – during neonatal period a low I/G ratio is critical for survival

## **Effects on Glucagon Secretion**

### **Stimuli for Glucagon Secretion**

↓ Blood glucose

↑ Serum amino acids (arginine, alanine)

Sympathetic nervous system stimulation

Stress

Exercise

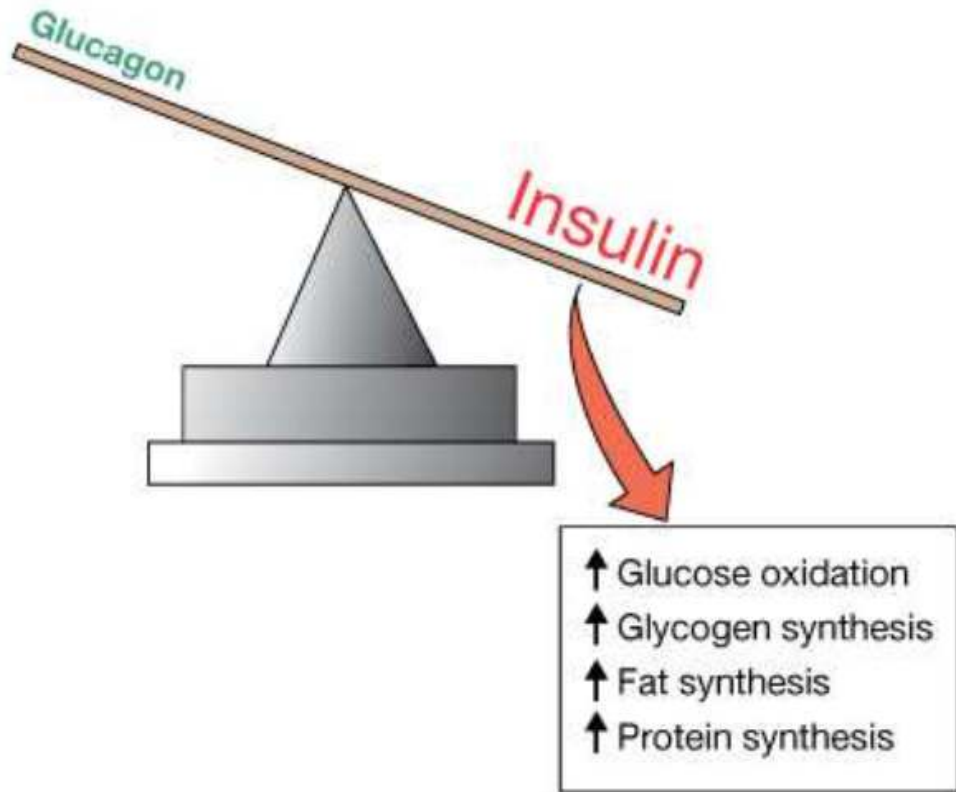
### **Inhibitors of Glucagon Secretion**

Somatostatin

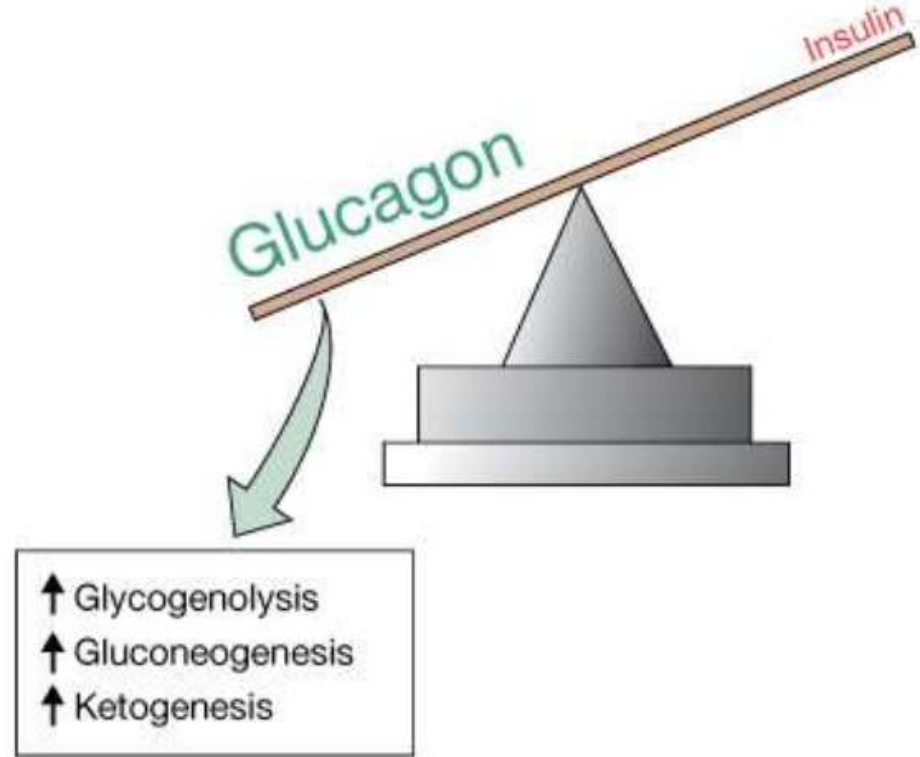
Insulin

↑ Blood glucose

(a) Fed state: insulin dominates



(b) Fasted state: glucagon dominates



## NORMAL PLASMA GLUCOSE LEVELS

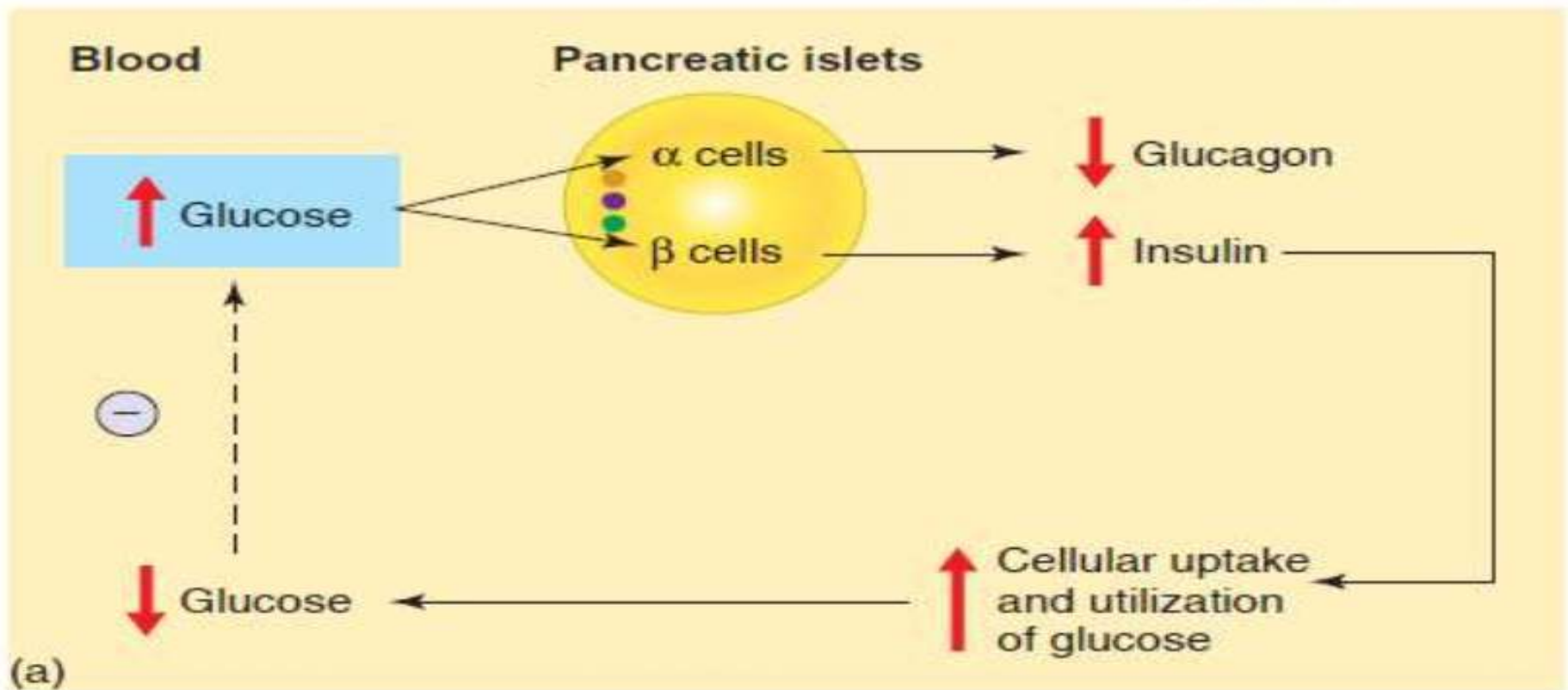
- Fasting : 70 – 100mg%
- Postprandial : 100 – 140mg%
- RBS : 80 – 120mg%

Rbs = random blood sugar

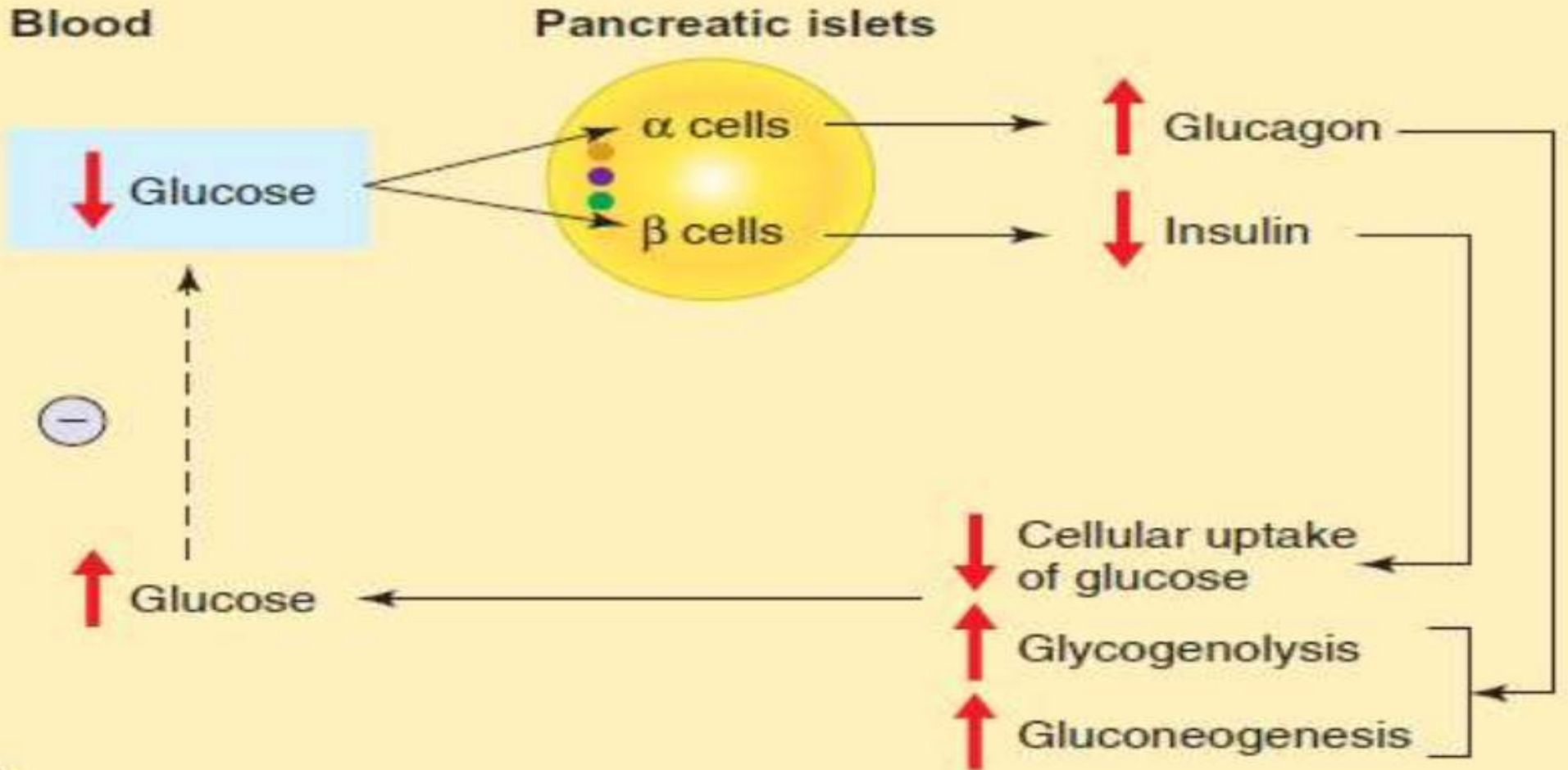


# GLUCOSE HOMEOSTASIS

- Sensor
- Integrating center
- Effector



# GLUCOSE HOMEOSTASIS



(b)

**Table: 49.4** Major differences between Type 1 and Type 2 diabetes mellitus.

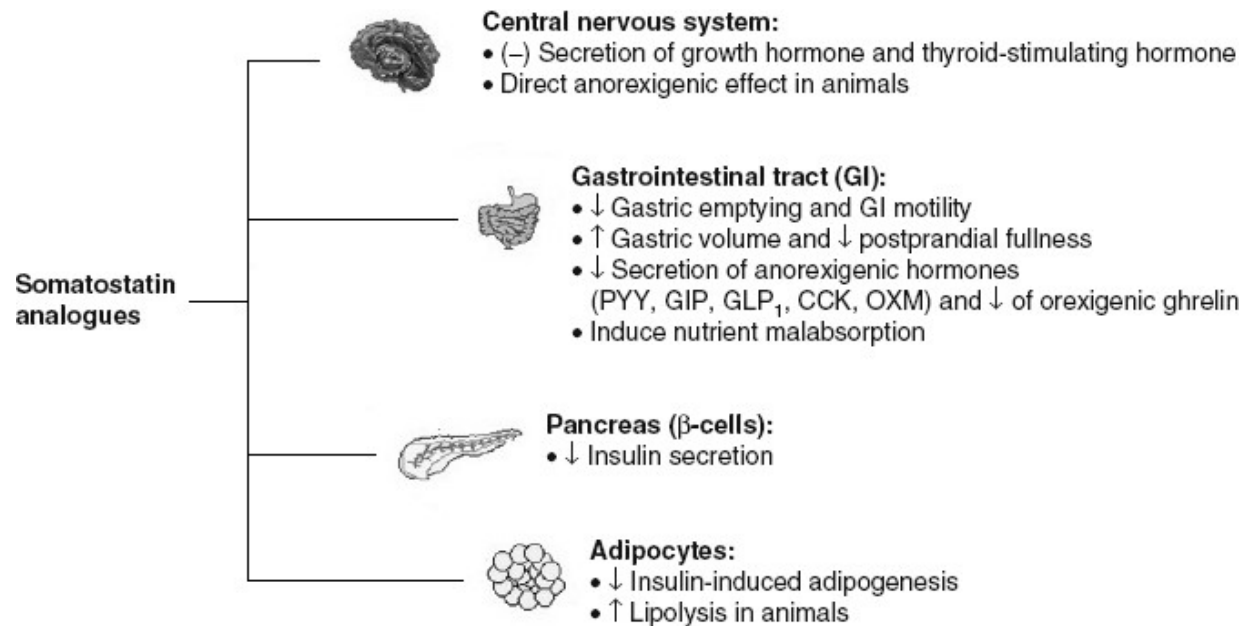
	Type I	Type II
1. Age of onset	Before the age of 40 (Juvenile onset diabetes)	After the age of 40 (Maturity onset diabetes)
2. Body fat mass	Not obese	Obese
3. Incidence	10% of the total diabetes	90% of the total diabetes
4. Genetic susceptibility	Concordance rate is < 50%	Concordance rate is > 50%
5. Incidence of ketoacidosis	High	Low
6. B cell mass of pancreas	B cells destroyed	B cells morphology is normal.
7. Nature of onset	Rapid	Gradual
8. Usual complication	Ketoacidotic coma	Hyperosmolar coma

## SOMATOSTATIN

Secreted from D cells of pancreas

- Also secreted in SOMATOSTATIN hypothalamus & GIT

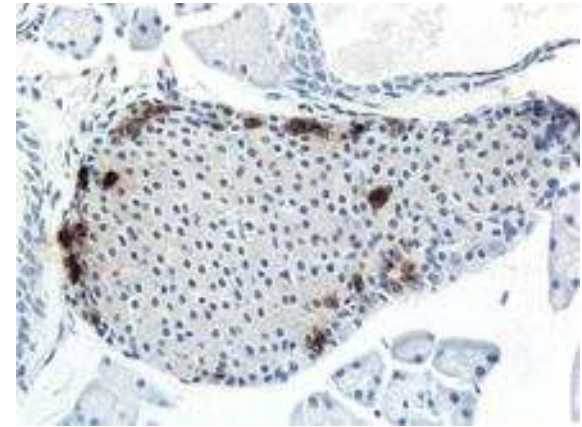
- **Inhibits secretion of insulin & glucagon**
- **Inhibits GI motility\* & GI secretions**
- **Regulates feedback control of gastric emptying**



## PANCREATIC POLYPEPTIDE

- Pancreatic polypeptide (PP) is a polypeptide secreted from F cells of pancreas or PP Cells = predominantly in the head of the pancreas.

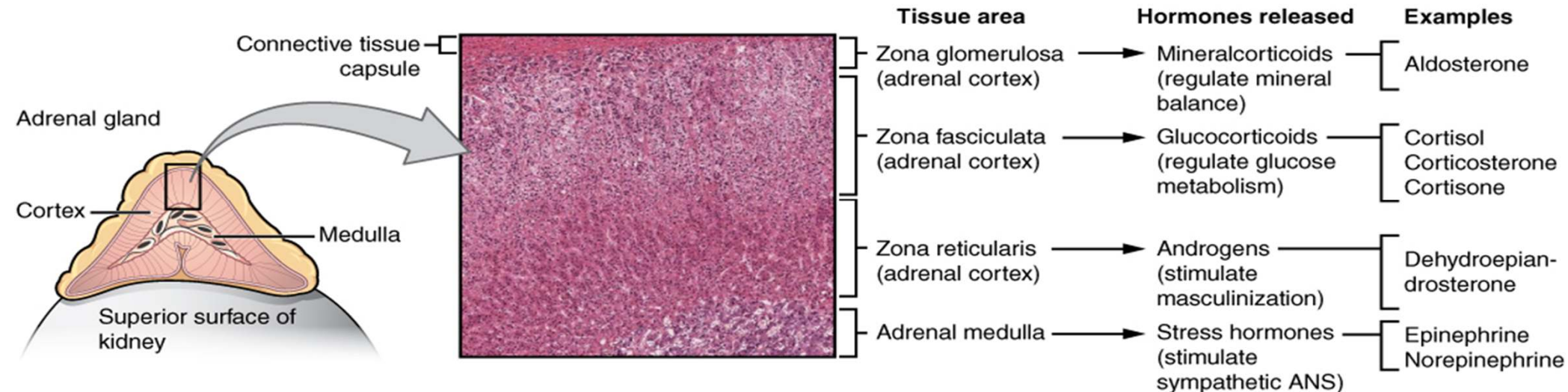
- Structurally similar to Neuropeptide Y secreted from hypothalamus
- Secreted in response to food intake
- Inhibits exocrine pancreatic secretion
- Slows the absorption of food from the GI tract



- The function of PP is to self-regulate pancreatic secretion activities (endocrine and exocrine).
- It also has effects on hepatic **glycogen** levels and gastrointestinal secretions.
- Its secretion in humans is increased after a protein meal, **fasting**, exercise, and acute **hypoglycemia**, and is decreased by **somatostatin** and intravenous **glucose**.
- Plasma PP has been shown to be reduced in conditions associated with increased food intake and elevated in **anorexia nervosa**. In addition, peripheral administration of PP has been shown to decrease food intake



# Adrenal glands

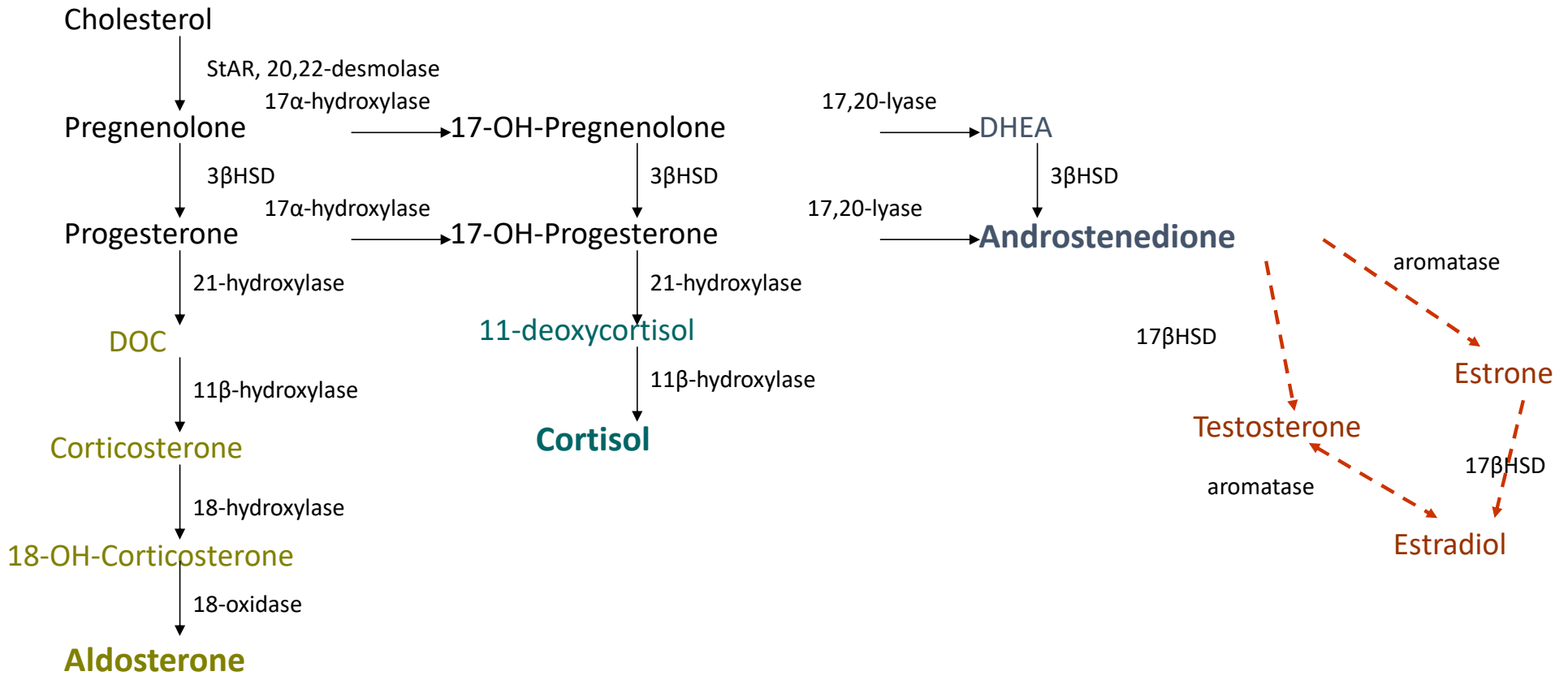


Glucocorticoids are chiefly produced in the zona fasciculata of the adrenal cortex  
Cortisol (or hydrocortisone) is the most important human glucocorticoid.

Glucocorticoids are corticosteroids that bind to the glucocorticoid receptor

ACTH

# Steroid Biosynthesis



- **GLUCOCORTICOIDS**

(regulate metabolism & are critical in stress response)

– CORTISOL responsible for control and metabolism of:

a. CHO (carbohydrates)

- increase glucose formed
- increase glucose released

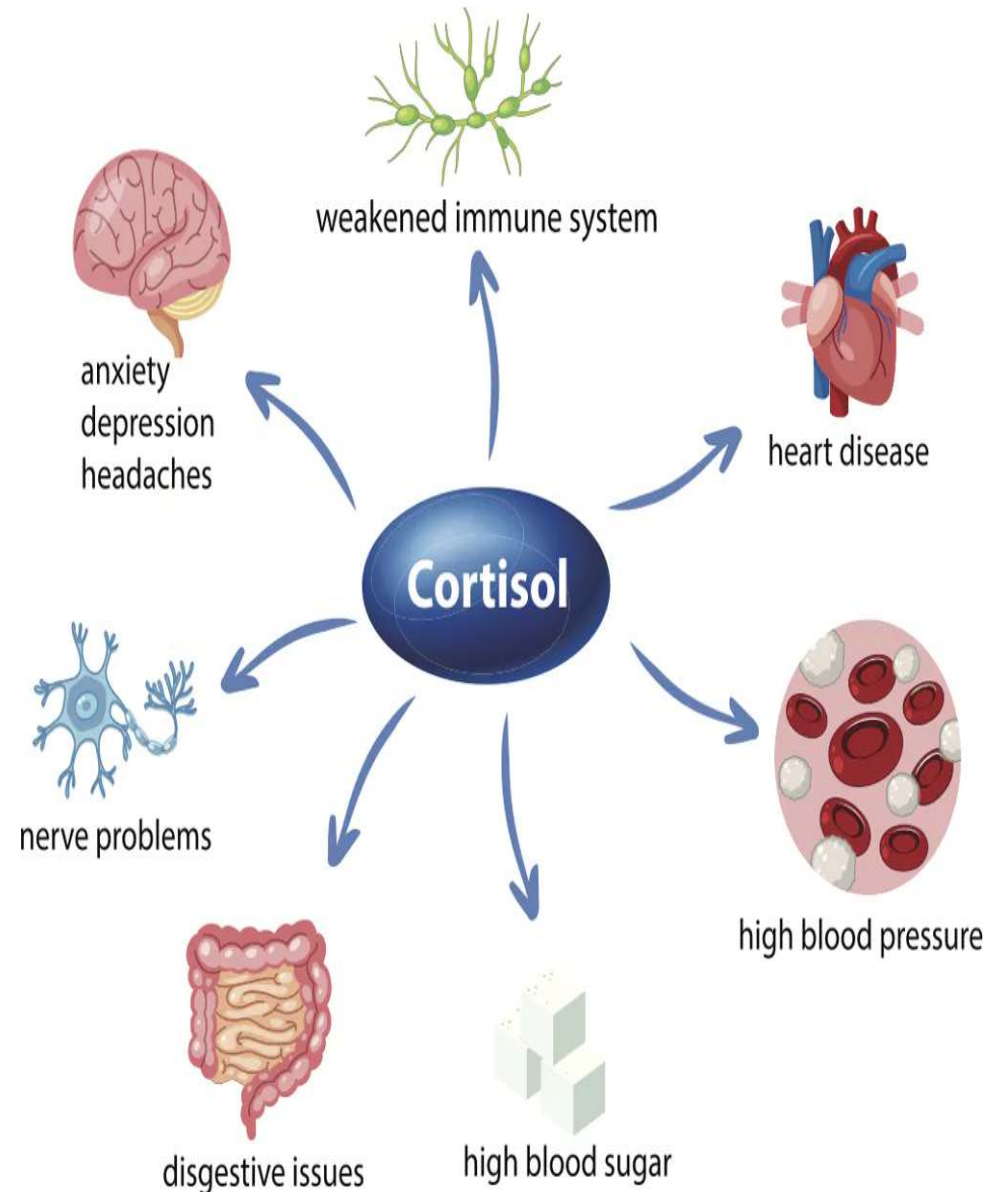
## **CORTISOL**

**FATS-control of fat metabolism**

- stimulates fatty acid mobilization from adipose tissue

**PROTEINS-control of protein metabolism**

- stimulates protein synthesis in liver
- protein breakdown in tissues
- decrease inflammatory and allergic response
- decrease immune system therefore prone to infection



**Glucocorticoid effects may be broadly classified into two major categories:**

**1/immunological**

**2/metabolic.**

**In addition, glucocorticoids play important roles in**

**1. fetal development and body fluid homeostasis.**

**2. Immune**

**3. Metabolic**

**4. Developmental**

**5. Arousal and cognition**

**6. Body fluid homeostasis**

## Cortisol and Immune

- up-regulate the expression of anti-inflammatory proteins.
- down-regulate the expression of proinflammatory proteins.
- Glucocorticoids are also shown to play a role in the development and homeostasis of T lymphocytes.
  - with either **increased or decreased sensitivity of T cell lineage to glucocorticoids.**



## Metabolic

Involved in glucose metabolism.

In the fasted state, cortisol stimulates several processes that collectively serve to increase and maintain normal concentrations of glucose in blood.

### Metabolic effects:

- **Stimulation of gluconeogenesis**, in particular, in the **liver**: This pathway results in the synthesis of glucose from non-hexose substrates, such as amino acids and glycerol from triglyceride breakdown.
- **Mobilization of amino acids** from extrahepatic tissues: These serve as substrates for gluconeogenesis.
- **Inhibition of glucose uptake** in muscle and adipose tissue: A mechanism to conserve glucose
- **Stimulation of fat breakdown** in adipose tissue: The fatty acids released by lipolysis are used for production of energy in tissues like muscle, and the released glycerol provide another substrate for gluconeogenesis.
- **Increase in sodium retention and potassium excretion** leads to hypernatremia and hypokalemia
- **Increase in hemoglobin concentration**, likely due to hindrance of the ingestion of red blood cell by macrophage or other phagocyte.
- **Increased urinary uric acid**
- **Increased urinary calcium and hypocalcemia**
- **Alkalosis**
- **Leukocytosis**

Excessive glucocorticoid levels resulting from administration as a drug or hyperadrenocorticism have effects on many systems.

Some examples include inhibition of bone formation, suppression of calcium absorption (both of which can lead to osteoporosis), delayed wound healing, muscle weakness, and increased risk of infection.

These observations suggest a multitude of less-dramatic physiologic roles for glucocorticoids.

## Developmental

- Glucocorticoids have multiple effects on fetal development.
- An important example is their role in **promoting maturation of the lung and production of the surfactant necessary for extrauterine lung function.**
- In addition, glucocorticoids are necessary for **normal brain development, by initiating terminal maturation, remodeling axons and dendrites, and affecting cell survival** and may also play a role in hippocampal development.
- Glucocorticoids **stimulate the maturation of the Na<sup>+</sup>/K<sup>+</sup>/ATPase, nutrient transporters, and digestion enzymes, promoting the development of a functioning gastro-intestinal system.**
- Glucocorticoids also support the development of the **neonate's renal system by increasing glomerular filtration.**

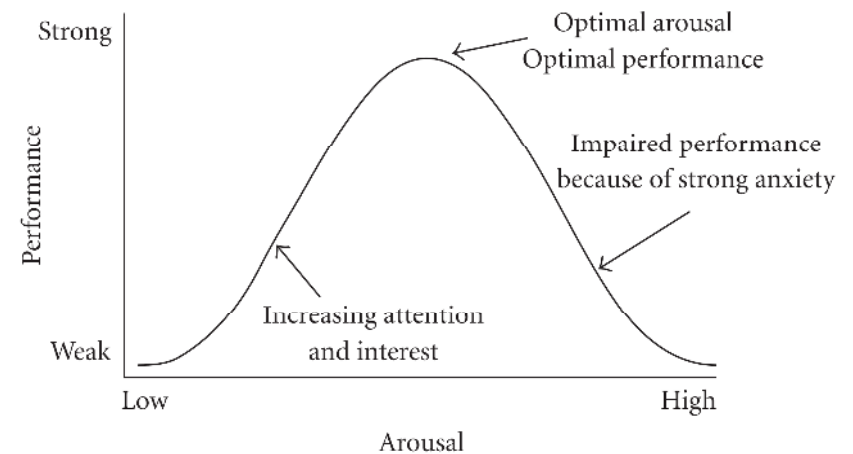
## **Body fluid homeostasis**

- Glucocorticoids could act centrally, as well as peripherally, to assist in the normalization of extracellular fluid volume by regulating body's action to atrial natriuretic peptide (ANP).
- Centrally, glucocorticoids could inhibit dehydration induced water intake
- Peripherally , glucocorticoids could induce a potent diuresis.

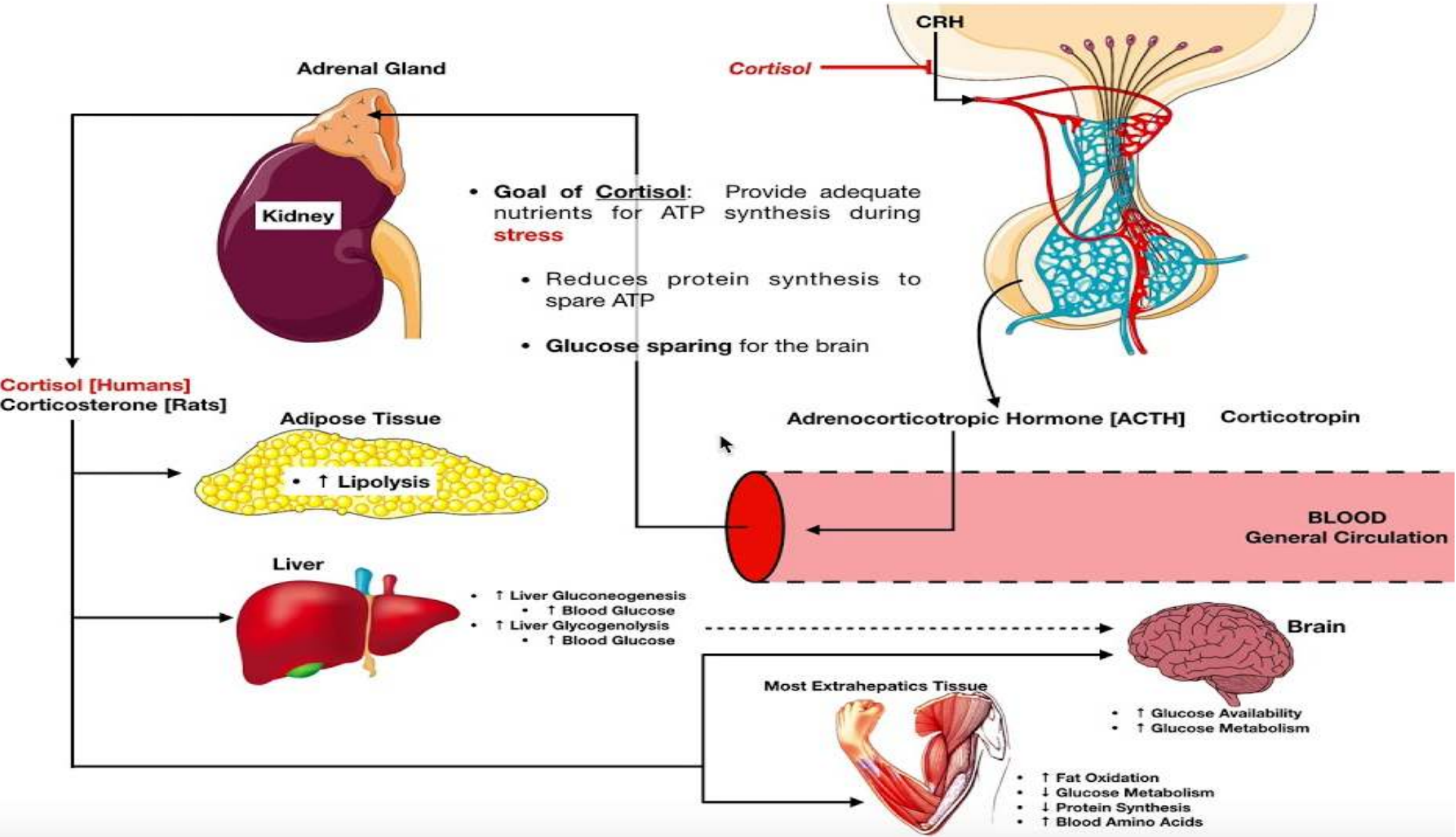
## Arousal and cognition

- A graphical representation of the Yerkes-Dodson curve
- Glucocorticoids act on the hippocampus, amygdala, and frontal lobes. Along with adrenaline, these enhance the formation of flashbulb memories of events associated with strong emotions, both positive and negative.
- Glucocorticoids have also been shown to have a significant impact on vigilance (attention deficit disorder) and cognition (memory).

Figure 1: The Yerkes-Dodson Human Performance and Stress Curve



The **Yerkes-Dodson law**, performance increases with physiological or mental arousal (stress) but only up to a point. When the level of stress becomes too high, performance decreases. There's more: The shape of the **curve** varies based on the complexity and familiarity of the task





## CORTISOL (THE STRESS HORMONE).



shuts down  
digestion



increases blood  
pressure



suppresses  
thyroid function



delays ovulation



raises blood sugar



impairs immune  
system

@NICOLEMJARDIM

# ACTH

## cortisol

- ↓ – levels cause stimulation of ACTH
- ↑ – levels cause dec. release of ACTH

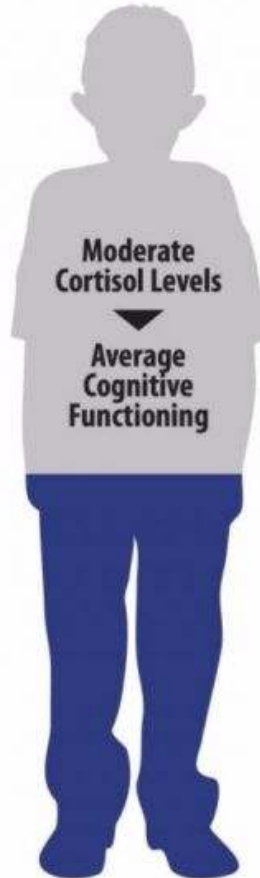
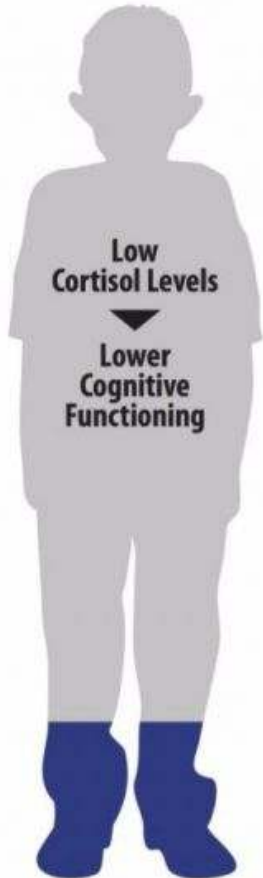
**At Risk Environment:**

Family Instability and Parental Emotional Unavailability

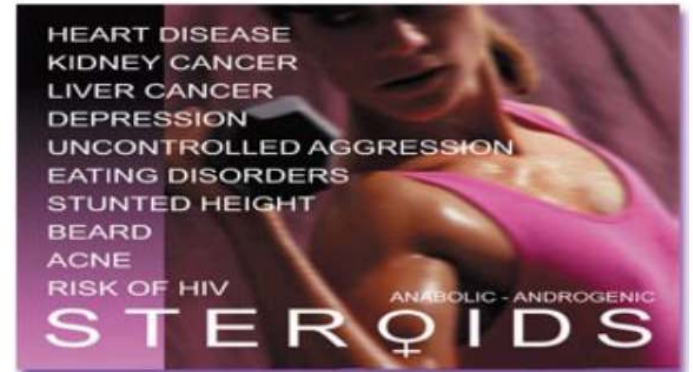


**At Risk Environment:**

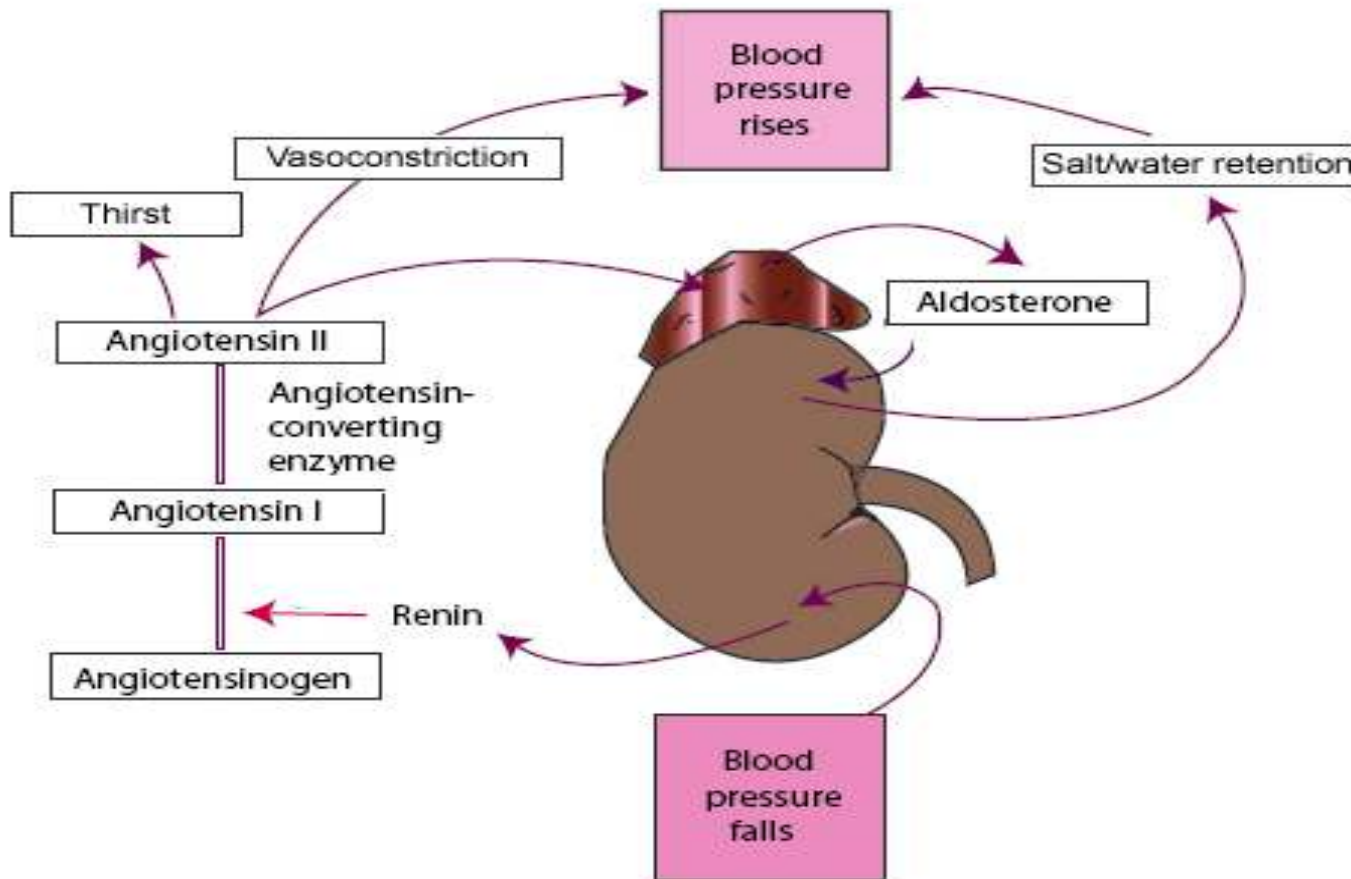
Parental Emotional Unavailability



# Too many steroids



# Adrenal physiology :Renin-angiotensin system



## **Mineralocorticoids** (F & E balance)

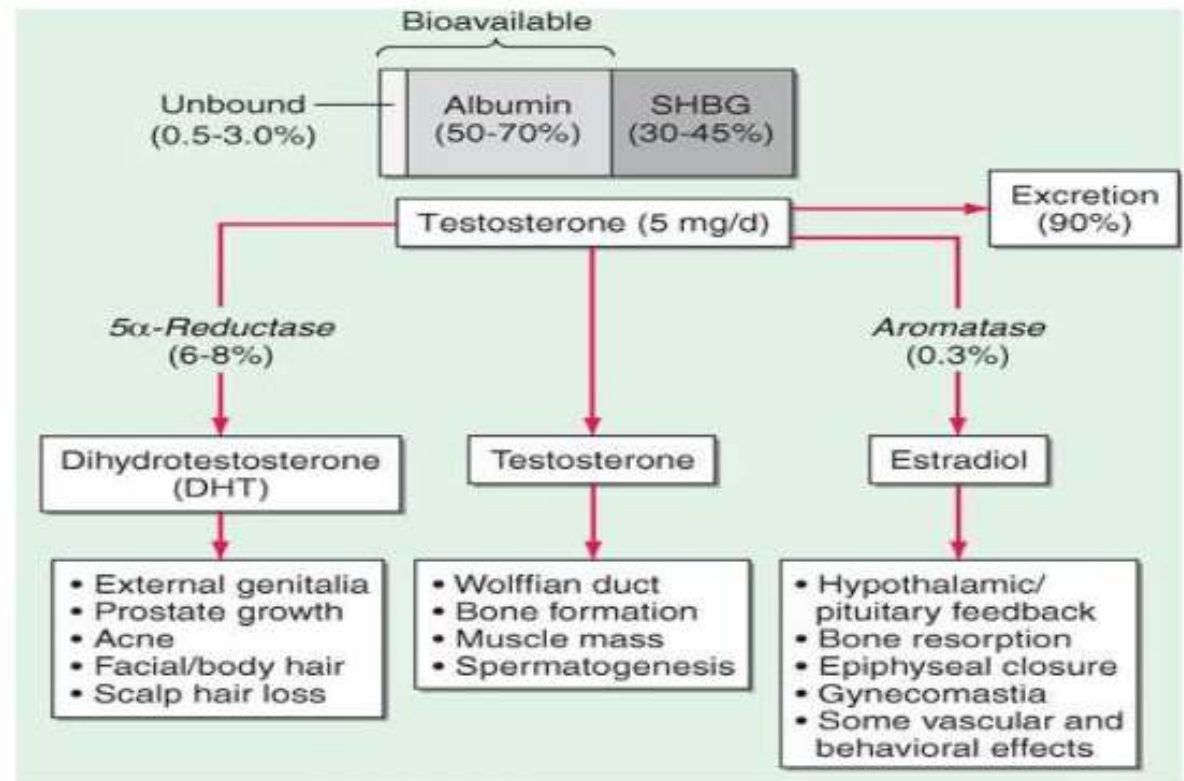
– Aldosterone (renin from kidneys controls adrenal cortex production of aldosterone)

- Na retention
- Water retention
- K excretion

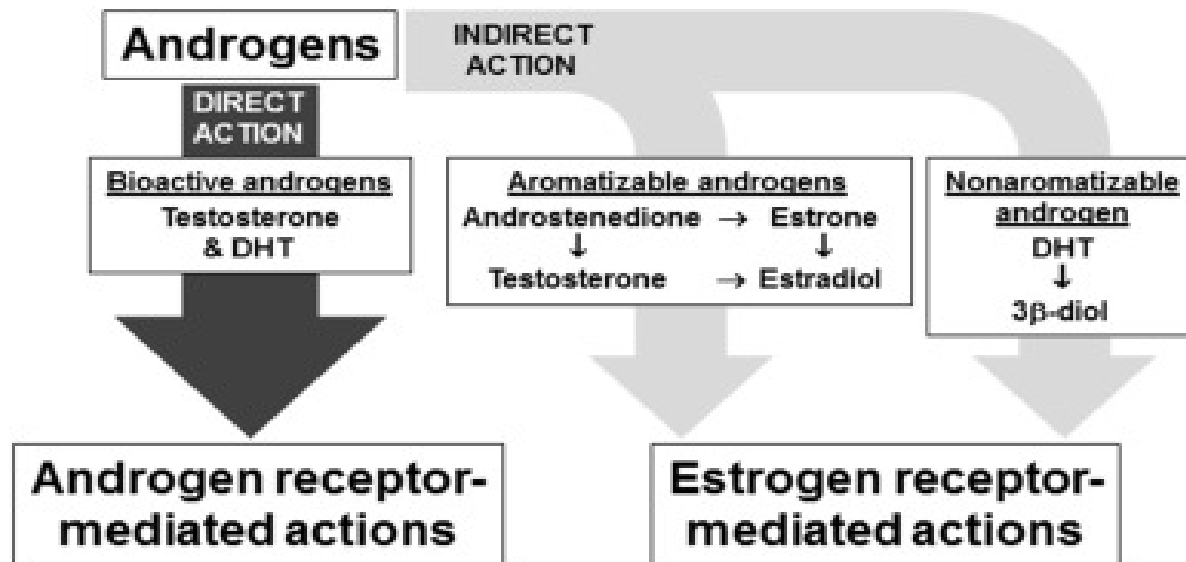
# ANDROGENS = SEX HORMONES

- – hormones which male characteristics
- • release of testosterone INCREASED
- Clear more in women than men

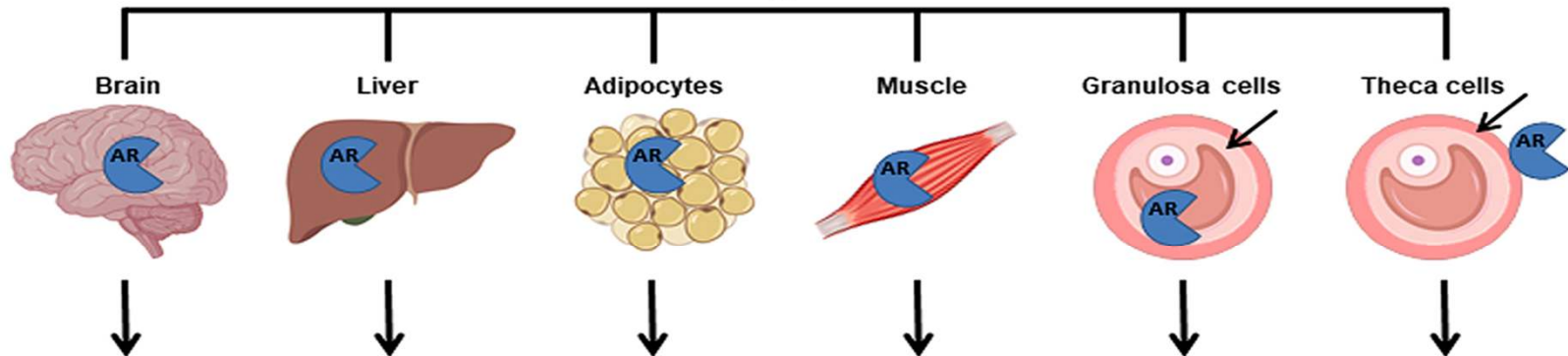
## ANDROGEN METABOLISM AND ACTIONS



Details will be discussed in male reproduction



### Androgen excess



Sites of AR actions hypothesised to be driving AR-mediated generation of PCOS-like traits

PCOS-like traits observed to be fully or partially ameliorated by a loss of site-specific AR actions

- Ovulatory dysfunction
- Polycystic ovaries
- Adiposity
- Adipocyte hypertrophy
- Dyslipidemia
- Hepatic steatosis

At present unknown.

At present unknown.

At present unknown.

- Irregular cycles
- Granulosa cell degeneration

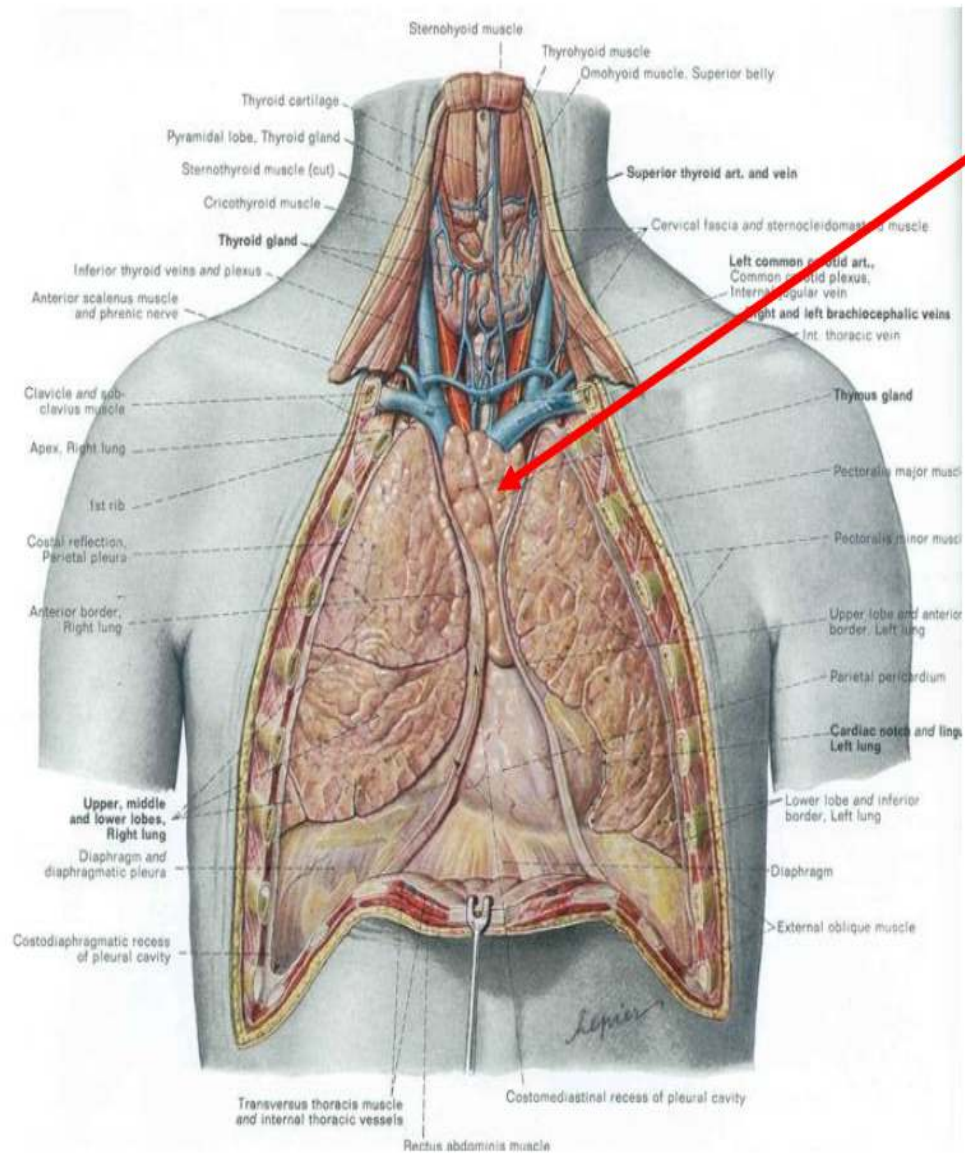
- Irregular cycles
- Ovulatory dysfunction
- Polycystic ovaries
- Granulosa cell degeneration

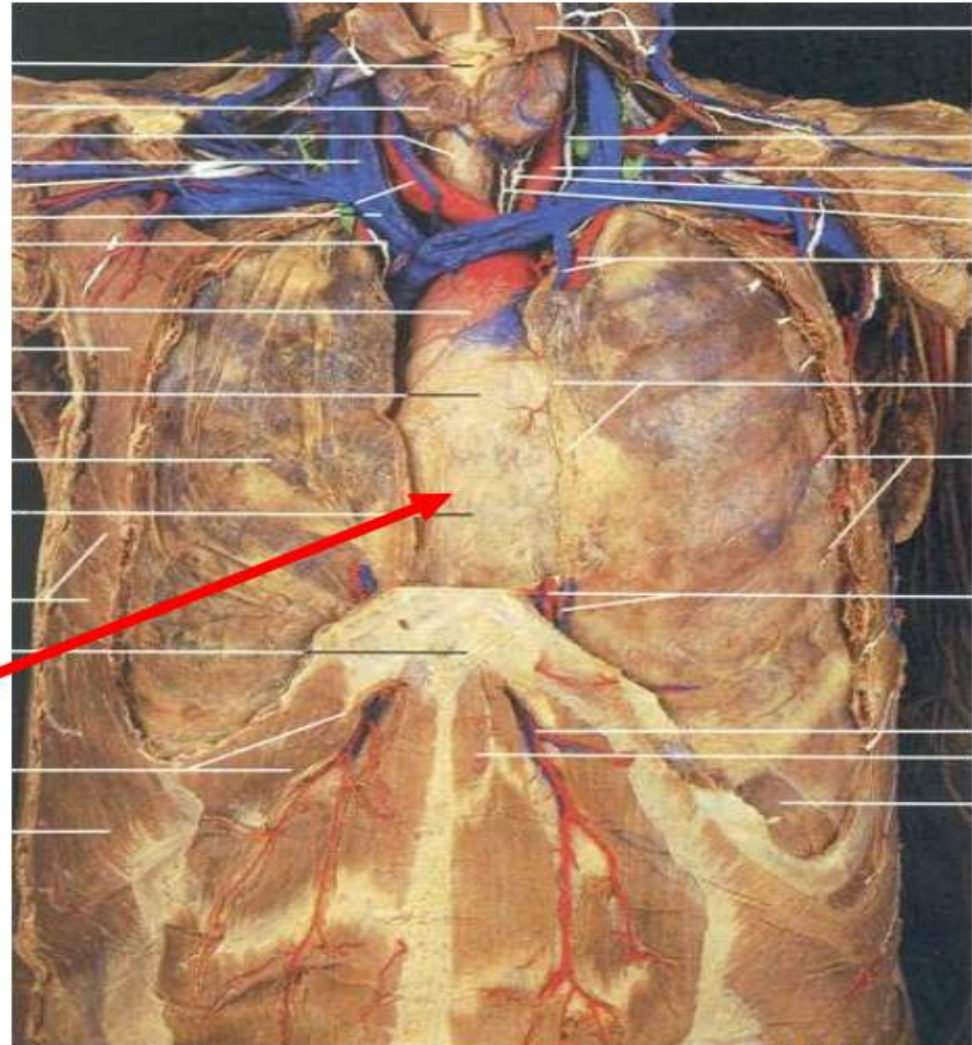
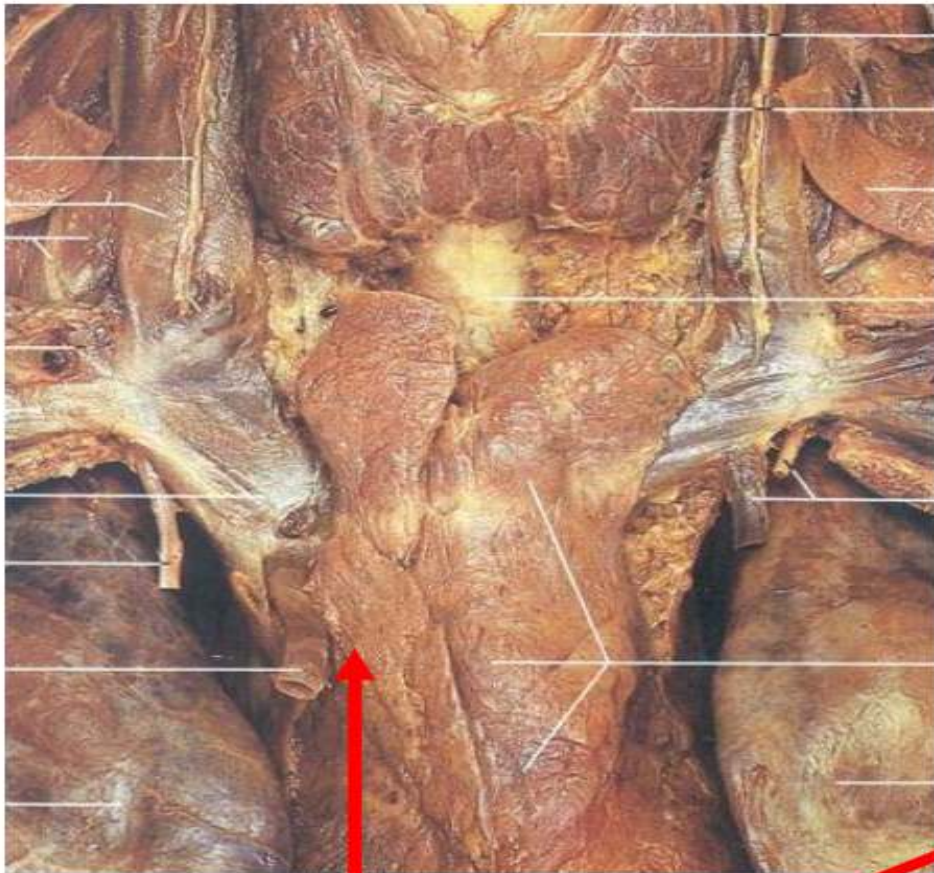


## THYMUS GLAND

## Adult THYMUS

- Located in the upper thorax region.
- Large in infants and children, it decreases in size throughout adulthood.
- By old age, it is composed mostly of fibrous connective tissue and fat.
- Thymus produces a hormone called **thymosin**.
- During childhood, it acts as an incubator for the maturation of a special group of whiteblood cells (T lymphocytes or T cells).
- T cells play a great role in immune response.





**Adult  
THYMUS**

- Many body organs not normally considered endocrine organs contain isolated cell clusters that secrete hormones.
- Examples include
  - the heart (atrial natriuretic peptide);
  - gastrointestinal tract organs (gastrin, secretin, and others);
  - the placenta (hormones of pregnancy—estrogen, progesterone, and others);
  - the kidneys (erythropoietin and renin);
  - the thymus; skin (cholecalciferol);
  - adipose tissue (leptin and resistin).
  - Bones

Hormonal tables

[https://en.wikipedia.org/wiki/List\\_of\\_human\\_hormones](https://en.wikipedia.org/wiki/List_of_human_hormones)



Name	Abbreviation	Tissue	Cells/Amino acid	Receptor	Target Tissue	Effect
Adrenaline, also known as epinephrine	EPI	adrenal gland	Adrenal medulla / Tyrosine	adrenergic receptor	nearly all tissues	blood pressure, glycogenolysis, lipolysis, etc.
Melatonin	MT	pineal gland	Pinealocyte / Tryptophan	melatonin receptor	CNS and peripheral tissue	circadian rhythm
Noradrenaline, also known as norepinephrine	NE	adrenal gland	Adrenal medulla / Tyrosine	noradrenergic receptor	nearly all tissues	blood pressure, glycogenolysis, lipolysis, etc.
Triiodothyronine	T <sub>3</sub>	peripheral tissue of thyroid gland	Thyroid follicular cell / Tyrosine	thyroid hormone receptor	nearly every cell in the body	increased metabolism
Thyroxine	T <sub>4</sub>	thyroid gland	Thyroid follicular cell / Tyrosine	thyroid hormone receptor	same as above	similar effect as T <sub>3</sub> but much weaker; converted to T <sub>3</sub> in target cells
Dopamine	DA	substantia nigra (mainly)	Phenylalanine / Tyrosine	D1 and D2	system-wide	regulation of cellular cAMP levels, prolactin antagonist



Eicosanoid for more information about this class of paracrine signalling chemicals and hormones.

Name	Abbreviation	Tissue	Cells	Receptor	Target Tissue	Effect
Prostaglandins	PG	seminal vesicle		prostaglandin receptor		vasodilation
Leukotrienes	LT	Blood	white blood cells	G protein-coupled receptors		increase vascular permeability
Prostacyclin	PGI <sub>2</sub>	endothelium		prostacyclin receptor		vasodilation, platelet activation inhibitor
Thromboxane	TXA <sub>2</sub>	Blood	platelets	thromboxane receptor		vasoconstriction, Platelet Aggregation

# Peptide

<a href="#">Vasoactive intestinal peptide</a>	VIP	<a href="#">gut, pancreas</a> , and <a href="#">suprachiasmatic nuclei</a> of the <a href="#">hypothalamus</a>			stimulates <a href="#">contractility</a> in the heart, causes <a href="#">vasodilation</a> , increases <a href="#">glycogenolysis</a> , lowers arterial <a href="#">blood pressure</a> and relaxes the smooth muscle of <a href="#">trachea</a> , stomach and <a href="#">gall bladder</a>
<a href="#">Uroguanylin</a>	UGN	renal tissues			regulates <a href="#">electrolyte</a> and <a href="#">water</a> transport in <a href="#">renal epithelia</a> .
<a href="#">Thyrotropin-releasing hormone</a>	TRH	<a href="#">hypothalamus</a>	<a href="#">Parvocellular neurosecretory neurons</a>	<a href="#">anterior pituitary</a>	Release <a href="#">thyroid-stimulating hormone</a> (primarily) Stimulate <a href="#">prolactin</a> release
<a href="#">Thyroid-stimulating hormone</a> (or thyrotropin)	TSH	<a href="#">anterior pituitary</a>	<a href="#">thyrotropes</a>	<a href="#">thyroid gland</a>	secrete <a href="#">thyroxine</a> (T <sub>4</sub> ) and <a href="#">triiodothyronine</a> (T <sub>3</sub> )
<a href="#">Thrombopoietin</a>	TPO	<a href="#">liver, kidney, striated muscle</a>	<a href="#">Myocytes</a>	<a href="#">megakaryocytes</a>	produce <a href="#">platelets</a> <sup>[6]</sup>
<a href="#">Somatostatin (or growth hormone–inhibiting hormone or growth hormone release–inhibiting hormone or somatotropin release–inhibiting factor or somatotropin release–inhibiting hormone)</a>	GHIH or GHRH or SRIF or SRIH	<a href="#">hypothalamus, islets of Langerhans, gastrointestinal system</a>	<a href="#">delta cells</a> in islets Neuroendocrine cells of the <a href="#">Periventricular nucleus</a> in hypothalamus		Inhibit release of <a href="#">GH</a> and <a href="#">TRH</a> from <a href="#">anterior pituitary</a> Suppress release of <a href="#">gastrin, cholecystokinin (CCK), secretin, motilin, vasoactive intestinal peptide (VIP), gastric inhibitory polypeptide (GIP), enteroglucagon</a> in <a href="#">gastrointestinal system</a> Lowers rate of gastric emptying Reduces <a href="#">smooth muscle</a> contractions and blood flow within the intestine <sup>[4]</sup> Inhibit release of <a href="#">insulin</a> from <a href="#">beta cells</a> <sup>[5]</sup> Inhibit release of <a href="#">glucagon</a> from <a href="#">alpha cells</a> <sup>[5]</sup> Suppress the exocrine secretory action of <a href="#">pancreas</a> .
<a href="#">Secretin</a>	SCT	<a href="#">duodenum</a>	<a href="#">S cell</a>		Secretion of <a href="#">bicarbonate</a> from <a href="#">liver, pancreas</a> and duodenal <a href="#">Brunner's glands</a> Enhances effects of <a href="#">cholecystokinin</a> Stops production of gastric juice
<a href="#">Renin</a>		<a href="#">Kidney</a>	<a href="#">Juxtaglomerular cells</a>		Activates the <a href="#">renin–angiotensin system</a> by producing <a href="#">angiotensin I</a> of <a href="#">angiotensinogen</a>
<a href="#">Relaxin</a>	RLN	<a href="#">Corpus luteum, Uterus, placenta, and Mammary gland</a>	<a href="#">Decidual cells</a>		Unclear in humans
<a href="#">Prolactin-releasing hormone</a>	PRLH	<a href="#">hypothalamus</a>			Release <a href="#">prolactin</a> from <a href="#">anterior pituitary</a>

<a href="#">Prolactin</a>	PRL	<a href="#">anterior pituitary, uterus</a>	<a href="#">lactotrophs</a> of anterior pituitary <a href="#">Decidual cells</a> of uterus	milk production in <a href="#">mammary glands</a> <a href="#">sexual gratification</a> after <a href="#">sexual acts</a>
<a href="#">Pituitary adenylate cyclase-activating peptide</a>	PACAP	multiple		Stimulates <a href="#">enterochromaffin-like cells</a>
<a href="#">Parathyroid hormone</a>	PTH	<a href="#">parathyroid gland</a>	<a href="#">parathyroid chief cell</a>	<ul style="list-style-type: none"> <li>•increase blood <math>Ca^{2+}</math>:indirectly stimulate <a href="#">osteoclasts</a></li> <li>•<math>Ca^{2+}</math> reabsorption in <a href="#">kidney</a></li> <li>•activate <a href="#">vitamin D</a></li> </ul> (Slightly) decrease blood <a href="#">phosphate</a> : <ul style="list-style-type: none"> <li>•(decreased reuptake in <a href="#">kidney</a> but increased uptake from bones</li> <li>•activate <a href="#">vitamin D</a>)</li> </ul>
<a href="#">Pancreatic polypeptide</a>		<a href="#">Pancreas</a>	<a href="#">PP cells</a>	Self-regulation of pancreatic secretions (endocrine and exocrine). It also affects hepatic glycogen levels and gastrointestinal secretions.
<a href="#">Oxytocin</a>	OXT	<a href="#">posterior pituitary</a>	<a href="#">Magnocellular neurosecretory cells</a>	release breast milkStimulates contraction of <a href="#">cervix</a> and <a href="#">vagina</a> . Involved in <a href="#">orgasm</a> , trust between people, <sup>[2]</sup> and <a href="#">circadian homeostasis</a> (body temperature, activity level, wakefulness). <sup>[3]</sup>
<a href="#">Osteocalcin</a>	OCN	<a href="#">Skeleton</a>	<a href="#">Osteoblasts</a>	Favors muscle function, memory formation, testosterone synthesis and energy expenditure <sup>[1]</sup>
<a href="#">Orexin</a>		<a href="#">hypothalamus</a>		wakefulness and increased energy expenditure, increased appetite
<a href="#">Motilin</a>	MLN	<a href="#">Small intestine</a>		stimulates gastric activity
<a href="#">Melanocyte stimulating hormone</a>	MSH or $\alpha$ -MSH	<a href="#">anterior pituitary/pars intermedia</a>	<a href="#">Melanotroph</a>	<a href="#">melanogenesis</a> by <a href="#">melanocytes</a> in <a href="#">skin</a> and <a href="#">hair</a>
<a href="#">Luteinizing hormone</a>	LH	<a href="#">anterior pituitary</a>	<a href="#">gonadotropes</a>	In female: <a href="#">ovulation</a> In male: stimulates <a href="#">Leydig cell</a> production of <a href="#">testosterone</a>
<a href="#">Lipotropin</a>	LPH	<a href="#">anterior pituitary</a>	<a href="#">Corticotropes</a>	<a href="#">lipolysis</a> and <a href="#">steroidogenesis</a> , stimulates <a href="#">melanocytes</a> to produce <a href="#">melanin</a>
<a href="#">Leptin</a>	LEP	<a href="#">adipose tissue</a>		decrease of <a href="#">appetite</a> and increase of <a href="#">metabolism</a> .
<a href="#">Insulin-like growth factor</a> (or somatomedin)	IGF	<a href="#">liver</a>	<a href="#">Hepatocytes</a>	insulin-like effectsregulate <a href="#">cell growth</a> and development

<a href="#">Insulin</a>	INS	<a href="#">pancreas</a>	<a href="#">beta cells</a>	Intake of <a href="#">glucose</a> , <a href="#">glycogenesis</a> and <a href="#">glycolysis</a> in <a href="#">liver</a> and <a href="#">muscle</a> from blood intake of <a href="#">lipids</a> and synthesis of <a href="#">triglycerides</a> in <a href="#">adipocytes</a> Other <a href="#">anabolic</a> effects
<a href="#">Inhibin</a>		<a href="#">testes</a> , <a href="#">ovary</a> , <a href="#">fetus</a>	<a href="#">Sertoli cells</a> of testes <a href="#">granulosa cells</a> of ovary <a href="#">trophoblasts</a> in fetus	Inhibit production of <a href="#">FSH</a>
<a href="#">Human placental lactogen</a>	HPL	<a href="#">placenta</a>		increase production of <a href="#">insulin</a> and <a href="#">IGF-1</a> increase <a href="#">insulin resistance</a> and <a href="#">carbohydrate</a> intolerance

<a href="#">Human chorionic gonadotropin</a>	hCG	<a href="#">placenta</a>	<a href="#">syncytiotrophoblast</a> cells	promote maintenance of <a href="#">corpus luteum</a> during beginning of <a href="#">pregnancy</a> Inhibit <a href="#">immune</a> response, towards the <a href="#">human embryo</a> .
<a href="#">Hepcidin</a>	HAMP	<a href="#">liver</a>		inhibits iron export from cells
<a href="#">Guanylin</a>	GN	<a href="#">gut</a>		regulates <a href="#">electrolyte</a> and <a href="#">water</a> transport in <a href="#">intestinal epithelia</a> .
<a href="#">Growth hormone-releasing hormone</a>	GHRH	<a href="#">hypothalamus</a>		Release <a href="#">GH</a> from <a href="#">anterior pituitary</a>
<a href="#">Growth hormone</a>	GH or hGH	<a href="#">anterior pituitary</a>	<a href="#">somatotropes</a>	stimulates <a href="#">growth</a> and <a href="#">cell</a> reproduction Release <a href="#">Insulin-like growth factor 1</a> from <a href="#">liver</a>
<a href="#">Gonadotropin-releasing hormone</a>	GnRH	<a href="#">hypothalamus</a>		Release of <a href="#">FSH</a> and <a href="#">LH</a> from <a href="#">anterior pituitary</a> .
<a href="#">Glucagon-like peptide-1</a>	GLP1	<a href="#">ileum</a>	<a href="#">L cells</a>	Stimulates the <a href="#">adenylyl cyclase</a> pathway, resulting in increased synthesis and release of <a href="#">insulin</a>
<a href="#">Glucagon</a>	GCG	<a href="#">pancreas</a>	<a href="#">alpha cells</a>	<a href="#">glycogenolysis</a> and <a href="#">gluconeogenesis</a> in <a href="#">liver</a> increases blood glucose level
<a href="#">Ghrelin</a>		<a href="#">stomach</a>	<a href="#">P/D1 cell</a>	Stimulate <a href="#">appetite</a> , secretion of <a href="#">growth hormone</a> from <a href="#">anterior pituitary gland</a>
<a href="#">Gastrin</a>	GAS	<a href="#">stomach</a> , <a href="#">duodenum</a>	<a href="#">G cell</a>	Secretion of <a href="#">gastric acid</a> by <a href="#">parietal cells</a>
<a href="#">Gastric inhibitory polypeptide</a>	GIP	mucosa of the <a href="#">duodenum</a> and the <a href="#">jejunum</a>	<a href="#">K cell</a>	Induces <a href="#">insulin</a> secretion

<a href="#">Galanin</a>	GAL	central nervous system and gastrointestinal tract			modulation and inhibition of <a href="#">action potentials</a> in <a href="#">neurons</a>
<a href="#">Gastric inhibitory polypeptide</a>	GIP	mucosa of the <a href="#">duodenum</a> and the <a href="#">jejunum</a>	<a href="#">K cell</a>		Induces <a href="#">insulin</a> secretion
<a href="#">Gastrin</a>	GAS	<a href="#">stomach, duodenum</a>	<a href="#">G cell</a>		Secretion of <a href="#">gastric acid</a> by <a href="#">parietal cells</a>
<a href="#">Ghrelin</a>		<a href="#">stomach</a>	<a href="#">P/D1 cell</a>		Stimulate <a href="#">appetite</a> , secretion of <a href="#">growth hormone</a> from <a href="#">anterior pituitary gland</a>
<a href="#">Glucagon</a>	GCG	<a href="#">pancreas</a>	<a href="#">alpha cells</a>		<a href="#">glycogenolysis</a> and <a href="#">gluconeogenesis</a> in <a href="#">liver</a> increases blood glucose level
<a href="#">Glucagon-like peptide-1</a>	GLP1	<a href="#">ileum</a>	<a href="#">L cells</a>	<a href="#">pancreatic beta cells</a>	Stimulates the <a href="#">adenyl cyclase</a> pathway, resulting in increased synthesis and release of <a href="#">insulin</a>
<a href="#">Gonadotropin-releasing hormone</a>	GnRH	<a href="#">hypothalamus</a>			Release of <a href="#">FSH</a> and <a href="#">LH</a> from <a href="#">anterior pituitary</a> .
<a href="#">Growth hormone-releasing hormone</a>	GHRH	<a href="#">hypothalamus</a>			Release <a href="#">GH</a> from <a href="#">anterior pituitary</a>
<a href="#">Hepcidin</a>	HAMP	<a href="#">liver</a>			inhibits iron export from cells
<a href="#">Human chorionic gonadotropin</a>	hCG	<a href="#">placenta</a>	<a href="#">syncytiotrophoblast cells</a>		promote maintenance of <a href="#">corpus luteum</a> during beginning of <a href="#">pregnancy</a> Inhibit <a href="#">immune</a> response, towards the <a href="#">human embryo</a> .
<a href="#">Human placental lactogen</a>	HPL	<a href="#">placenta</a>			increase production of <a href="#">insulin</a> and <a href="#">IGF-1</a> increase <a href="#">insulin resistance</a> and <a href="#">carbohydrate</a> intolerance
<a href="#">Growth hormone</a>	GH or hGH	<a href="#">anterior pituitary</a>	<a href="#">somatotropes</a>		stimulates <a href="#">growth</a> and <a href="#">cell</a> reproduction Release <a href="#">Insulin-like growth factor 1</a> from <a href="#">liver</a>
<a href="#">Inhibin</a>		<a href="#">testes, ovary, fetus</a>	<a href="#">Sertoli cells</a> of testes <a href="#">granulosa cells</a> of ovary <a href="#">trophoblasts</a> in fetus		Inhibit production of <a href="#">FSH</a>
<a href="#">Insulin</a>	INS	<a href="#">pancreas</a>	<a href="#">beta cells</a>		Intake of <a href="#">glucose</a> , <a href="#">glycogenesis</a> and <a href="#">glycolysis</a> in <a href="#">liver</a> and <a href="#">muscle</a> from blood intake of <a href="#">lipids</a> and synthesis of <a href="#">triglycerides</a> in <a href="#">adipocytes</a> Other <a href="#">anabolic</a> effects
<a href="#">Insulin-like growth factor</a> (or somatomedin)	IGF	<a href="#">liver</a>	<a href="#">Hepatocytes</a>		insulin-like effects regulate <a href="#">cell growth</a> and development



<a href="#">Leptin</a>	LEP	<a href="#">adipose tissue</a>			decrease of <a href="#">appetite</a> and increase of <a href="#">metabolism</a> .
<a href="#">Lipotropin</a>	LPH	<a href="#">anterior pituitary</a>	<a href="#">Corticotropes</a>		<a href="#">lipolysis</a> and <a href="#">steroidogenesis</a> , stimulates <a href="#">melanocytes</a> to produce <a href="#">melanin</a>
<a href="#">Luteinizing hormone</a>	LH	<a href="#">anterior pituitary</a>	<a href="#">gonadotropes</a>		In female: <a href="#">ovulation</a> In male: stimulates <a href="#">Leydig cell</a> production of <a href="#">testosterone</a>
<a href="#">Melanocyte stimulating hormone</a>	MSH or $\alpha$ -MSH	<a href="#">anterior pituitary/pars intermedia</a>	<a href="#">Melanotroph</a>		<a href="#">melanogenesis</a> by <a href="#">melanocytes</a> in <a href="#">skin</a> and <a href="#">hair</a>
<a href="#">Motilin</a>	MLN	<a href="#">Small intestine</a>			stimulates gastric activity
<a href="#">Orexin</a>		<a href="#">hypothalamus</a>			wakefulness and increased energy expenditure, increased appetite
<a href="#">Osteocalcin</a>	OCN	<a href="#">Skeleton</a>	<a href="#">Osteoblasts</a>	<a href="#">Muscle Brain Pancreas Testes</a>	Favors muscle function, memory formation, testosterone synthesis and energy expenditure
<a href="#">Oxytocin</a>	OXT	<a href="#">posterior pituitary</a>	<a href="#">Magnocellular neurosecretory cells</a>		release breast milkStimulates contraction of <a href="#">cervix</a> and <a href="#">vagina</a> . Involved in <a href="#">orgasm</a> , trust between people, and <a href="#">circadian homeostasis</a> (body temperature, activity level, wakefulness).
<a href="#">Pancreatic polypeptide</a>		<a href="#">Pancreas</a>	<a href="#">PP cells</a>		Self-regulation of pancreatic secretions (endocrine and exocrine). It also affects hepatic glycogen levels and gastrointestinal secretions.
<a href="#">Parathyroid hormone</a>	PTH	<a href="#">parathyroid gland</a>	<a href="#">parathyroid chief cell</a>		<ul style="list-style-type: none"> <li>•increase blood <a href="#">Ca<sup>2+</sup></a>; indirectly stimulate <a href="#">osteoclasts</a></li> <li>•Ca<sup>2+</sup> reabsorption in <a href="#">kidney</a></li> <li>•activate <a href="#">vitamin D</a> (Slightly) decrease blood <a href="#">phosphate</a>:</li> <li>•(decreased reuptake in <a href="#">kidney</a> but increased uptake from bones</li> <li>•activate <a href="#">vitamin D</a>)</li> </ul>
<a href="#">Pituitary adenylate cyclase-activating peptide</a>	PACAP	multiple			Stimulates <a href="#">enterochromaffin-like cells</a>

<a href="#">Prolactin</a>	PRL	<a href="#">anterior pituitary, uterus</a>	<a href="#">lactotrophs</a> of anterior pituitary <a href="#">Decidual cells</a> of uterus	milk production in <a href="#">mammary glands</a> <a href="#">sexual gratification</a> after <a href="#">sexual acts</a>
<a href="#">Prolactin-releasing hormone</a>	PRLH	<a href="#">hypothalamus</a>		Release <a href="#">prolactin</a> from <a href="#">anterior pituitary</a>
<a href="#">Relaxin</a>	RLN	<a href="#">Corpus luteum, Uterus, placenta, and Mammary gland</a>	<a href="#">Decidual cells</a>	Unclear in humans

<a href="#">Renin</a>		<a href="#">Kidney</a>	<a href="#">Juxtaglomerular cells</a>		Activates the <a href="#">renin-angiotensin system</a> by producing <a href="#">angiotensin I</a> of <a href="#">angiotensinogen</a>
<a href="#">Secretin</a>	SCT	<a href="#">duodenum</a>	<a href="#">S cell</a>		Secretion of <a href="#">bicarbonate</a> from <a href="#">liver</a> , <a href="#">pancreas</a> and duodenal <a href="#">Brunner's glands</a> Enhances effects of <a href="#">cholecystokinin</a> Stops production of gastric juice
<a href="#">Somatostatin (or growth hormone-inhibiting hormone or growth hormone release-inhibiting hormone or somatotropin release-inhibiting factor or somatotropin release-inhibiting hormone)</a>	GHIH or GHRIH or SRIF or SRIH	<a href="#">hypothalamus</a> , <a href="#">islets of Langerhans</a> , <a href="#">gastrointestinal system</a>	<a href="#">delta cells</a> in islets Neuroendocrine cells of the <a href="#">Periventricular nucleus</a> in hypothalamus		Inhibit release of <a href="#">GH</a> and <a href="#">TRH</a> from <a href="#">anterior pituitary</a> Suppress release of <a href="#">gastrin</a> , <a href="#">cholecystokinin</a> (CCK), <a href="#">secretin</a> , <a href="#">motilin</a> , <a href="#">vasoactive intestinal peptide</a> (VIP), <a href="#">gastric inhibitory polypeptide</a> (GIP), <a href="#">enteroglucagon</a> in <a href="#">gastrointestinal system</a> Lowers rate of gastric emptying Reduces <a href="#">smooth muscle</a> contractions and blood flow within the intestine <sup>[4]</sup> Inhibit release of <a href="#">insulin</a> from <a href="#">beta cells</a> Inhibit release of <a href="#">glucagon</a> from <a href="#">alpha cells</a> Suppress the exocrine secretory action of <a href="#">pancreas</a> .
<a href="#">Thrombopoietin</a>	TPO	<a href="#">liver</a> , <a href="#">kidney</a> , <a href="#">striated muscle</a>	<a href="#">Myocytes</a>	<a href="#">megakaryocytes</a>	produce <a href="#">platelets</a> <sup>[6]</sup>
<a href="#">Thyroid-stimulating hormone</a> (or thyrotropin)	TSH	<a href="#">anterior pituitary</a>	<a href="#">thyrotropes</a>	<a href="#">thyroid gland</a>	secrete <a href="#">thyroxine</a> (T <sub>4</sub> ) and <a href="#">triiodothyronine</a> (T <sub>3</sub> )
<a href="#">Thyrotropin-releasing hormone</a>	TRH	<a href="#">hypothalamus</a>	<a href="#">Parvocellular neurosecretory neurons</a>	<a href="#">anterior pituitary</a>	Release <a href="#">thyroid-stimulating hormone</a> (primarily) Stimulate <a href="#">prolactin</a> release
<a href="#">Vasoactive intestinal peptide</a>	VIP	<a href="#">gut</a> , <a href="#">pancreas</a> , and <a href="#">suprachiasmatic nuclei</a> of the <a href="#">hypothalamus</a>			stimulates <a href="#">contractility</a> in the heart, causes <a href="#">vasodilation</a> , increases <a href="#">glycogenolysis</a> , lowers arterial <a href="#">blood pressure</a> and relaxes the smooth muscle of <a href="#">trachea</a> , stomach and <a href="#">gall bladder</a>
<a href="#">Guanylin</a>	GN	gut			regulates <a href="#">electrolyte</a> and <a href="#">water</a> transport in <a href="#">intestinal epithelia</a> .
<a href="#">Uroguanylin</a>	UGN	renal tissues			regulates <a href="#">electrolyte</a> and <a href="#">water</a> transport in <a href="#">renal epithelia</a> .

# Steroid

Chemical class	Name	Abbreviation	Tissue	Cells	Target Tissue	Effect
<a href="#">androgen</a>	<a href="#">Testosterone</a>		<a href="#">testes, ovary</a>	<a href="#">Leydig cells</a>		<a href="#">libido</a> , <a href="#">Anabolic</a> : growth of <a href="#">muscle mass</a> and strength, increased <a href="#">bone density</a> , growth and strength, <a href="#">Virilizing</a> : <a href="#">maturation</a> of <a href="#">sex organs</a> , formation of <a href="#">scrotum</a> , deepening of voice, growth of <a href="#">beard</a> and <a href="#">axillary hair</a> .
<a href="#">androgen</a>	<a href="#">Dehydroepiandrosterone</a>	DHEA	<a href="#">testes, ovary, kidney</a>	<a href="#">Zona fasciculata</a> and <a href="#">Zona reticularis</a> cells of kidney <a href="#">theca cells</a> of ovary <a href="#">Leydig cells</a> of testes		<a href="#">Virilization</a> , <a href="#">anabolic</a>
<a href="#">androgen</a>	<a href="#">Androstenedione</a>		<a href="#">adrenal glands, gonads</a>			Substrate for <a href="#">estrogen</a>
<a href="#">androgen</a>	<a href="#">Dihydrotestosterone</a>	DHT	multiple			5-DHT or DHT is a male reproductive hormone that targets the prostate gland, bulbourethral gland, seminal vesicles, penis and scrotum and promotes growth/mitosis/cell maturation and differentiation. Testosterone is converted to 5-DHT by 5alpha-reductase, usually with in the target tissues of 5-DHT because of the need for high concentrations of 5-dht to produce the physiological effects.
<a href="#">mineralocorticoid</a>	<a href="#">Aldosterone</a>		<a href="#">adrenal cortex (zona glomerulosa)</a>			Increase <a href="#">blood volume</a> by reabsorption of <a href="#">sodium</a> in <a href="#">kidneys</a> (primarily) <a href="#">Potassium</a> and $H^+$ secretion in kidney.

<p><a href="#">estrogen</a></p>	<p><a href="#">Estradiol</a></p>	<p>E<sub>2</sub></p>	<p>females: <a href="#">ovary</a>, males <a href="#">testes</a></p>	<p>females: <a href="#">granulosa cells</a>, males: <a href="#">Sertoli cell</a></p>	<p><b>Females:</b>Structural:</p> <ul style="list-style-type: none"> <li>•promote formation of female <a href="#">secondary sex characteristics</a></li> <li>•stimulate <a href="#">endometrial</a> growth</li> <li>•increase <a href="#">uterine</a> growth</li> <li>•maintenance of <a href="#">blood vessels</a> and skin</li> <li>•reduce <a href="#">bone resorption</a></li> <li>•increase hepatic production of binding proteins</li> </ul> <p><u>Coagulation:</u></p> <ul style="list-style-type: none"> <li>•increase circulating level of <a href="#">factors 2, 7, 9, 10, antithrombin III, plasminogen</a></li> <li>•increase <a href="#">platelet</a> adhesiveness</li> </ul> <p>Fluid balance:</p> <ul style="list-style-type: none"> <li>•salt (<a href="#">sodium</a>) and water retention</li> <li>•increase <a href="#">growth hormone</a></li> <li>•increase <a href="#">cortisol, SHBG</a></li> </ul> <p>Gastrointestinal tract:</p> <ul style="list-style-type: none"> <li>•reduce bowel motility</li> <li>•increase cholesterol in <a href="#">bile</a></li> </ul> <p>Lung function:</p> <ul style="list-style-type: none"> <li>•promote lung function by supporting <a href="#">alveoli</a>.<sup>[7]</sup></li> </ul> <p><b>Males:</b> Prevent <a href="#">apoptosis</a> of germ cells<sup>[8]</sup></p>
---------------------------------	----------------------------------	----------------------	---	--	---




<a href="#">estrogen</a>	<a href="#">Estrone</a>		<a href="#">ovary</a>	<a href="#">granulosa cells</a> , <a href="#">Adipocytes</a>	
<a href="#">estrogen</a>	<a href="#">Estriol</a>	E <sub>3</sub>	<a href="#">placenta</a>	<a href="#">syncytiotrophoblast</a>	
<a href="#">glucocorticoid</a>	<a href="#">Cortisol</a>		<a href="#">adrenal cortex (zona fasciculata and zona reticularis cells)</a>		Stimulation of <a href="#">gluconeogenesis</a> Inhibition of glucose uptake in muscle and <a href="#">adipose</a> tissue Mobilization of <a href="#">amino acids</a> from <a href="#">extrahepatic</a> tissues Stimulation of <a href="#">fat breakdown</a> in adipose tissue <a href="#">anti-inflammatory</a> and <a href="#">immunosuppressive</a>
<a href="#">progesterone</a>	<a href="#">Progesterone</a>		<a href="#">ovary</a> , <a href="#">adrenal glands</a> , <a href="#">placenta</a> (when pregnant)	<a href="#">Granulosa cells</a> <a href="#">theca cells</a> of ovary	<ul style="list-style-type: none"> <li>•Support <a href="#">pregnancy</a>.<sup>[9]</sup>Convert <a href="#">endometrium</a> to secretory stage</li> <li>•Make <a href="#">cervical mucus</a> permeable to sperm</li> <li>•Inhibit <a href="#">immune</a> response, e.g. towards the <a href="#">human embryo</a>.</li> <li>•Decrease uterine <a href="#">smooth muscle</a> contractility<sup>[9]</sup></li> <li>•Inhibit <a href="#">lactation</a></li> <li>•Inhibit onset of <a href="#">labor</a></li> <li>•Support <a href="#">fetal</a> production of <a href="#">adrenal</a> mineralo- and glucosteroids</li> </ul> Other: <ul style="list-style-type: none"> <li>•Raise <a href="#">epidermal growth factor-1</a> levels</li> <li>•Increase core temperature during ovulation<sup>[10]</sup></li> <li>•Reduce <a href="#">spasm</a> and relax <a href="#">smooth muscle</a> (widen <a href="#">bronchi</a> and regulate <a href="#">mucus</a>)</li> <li>•<a href="#">Antiinflammatory</a>. Regulate <a href="#">immune response</a></li> <li>•Reduce <a href="#">gall-bladder</a> activity<sup>[11]</sup></li> <li>•Normalize <a href="#">blood</a> clotting and vascular tone, <a href="#">zinc</a> and <a href="#">copper</a> levels, <a href="#">cell oxygen</a> levels, and use of fat stores for energy</li> <li>•Assist in <a href="#">thyroid</a> function and <a href="#">bone</a> growth by <a href="#">osteoblasts</a></li> <li>•<a href="#">Resilience</a> in <a href="#">bone</a>, <a href="#">teeth</a>, <a href="#">gums</a>, <a href="#">joint</a>, <a href="#">tendon</a>, <a href="#">ligament</a> and <a href="#">skin</a> healing by regulating <a href="#">collagen</a></li> <li>•Nerve function and healing by regulating <a href="#">myelin</a></li> <li>•Prevent <a href="#">endometrial cancer</a> by regulating effects of estrogen</li> </ul>
<a href="#">secosteroid</a>	<a href="#">Calcitriol</a> (1,25-dihydroxyvitamin D <sub>3</sub> )		<a href="#">skin/proximal tubule</a> of <a href="#">kidneys</a>		Active form of <a href="#">vitamin D<sub>3</sub></a> Increase absorption of <a href="#">calcium</a> and <a href="#">phosphate</a> from <a href="#">gastrointestinal tract</a> and <a href="#">kidneys</a> inhibit release of <a href="#">PTH</a>
<a href="#">secosteroid</a>	<a href="#">Calcidiol</a> (25-hydroxyvitamin D <sub>3</sub> )		<a href="#">skin/proximal tubule</a> of <a href="#">kidneys</a>		Inactive form of <a href="#">vitamin D<sub>3</sub></a>

www.frankiejohn.com

www.frankiejohn.com

Grades don't measure  
intelligence and age doesn't  
define maturity...



 Share Y U NO GO? [www.frankiejohn.com](http://www.frankiejohn.com)

Monday, March 14, 2016