

Unless otherwise noted, the content of this course material is licensed under a Creative Commons Attribution - Non-Commercial - Share Alike 3.0 License.

Copyright 2008, Arno Kumagai, Gary Hammer



The following information is intended to inform and educate and is not a tool for self-diagnosis or a replacement for medical evaluation, advice, diagnosis or treatment by a healthcare professional. You should speak to your physician or make an appointment to be seen if you have questions or concerns about this information or your medical condition. You assume all responsibility for use and potential liability associated with any use of the material.

Material contains copyrighted content, used in accordance with U.S. law. Copyright holders of content included in this material should contact open.michigan@umich.edu with any questions, corrections, or clarifications regarding the use of content. The Regents of the University of Michigan do not license the use of third party content posted to this site unless such a license is specifically granted in connection with particular content objects. Users of content are responsible for their compliance with applicable law.

Adrenal Physiology & Steroid Pharmacology



2008

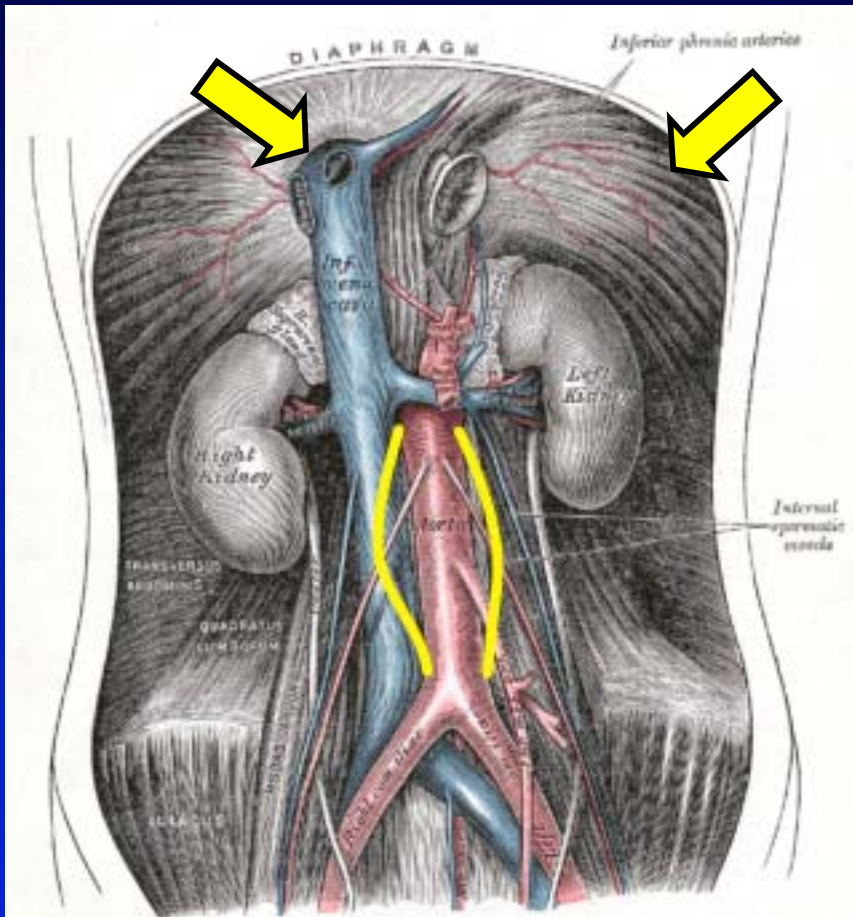
Gary D. Hammer, M.D., Ph.D.
University of Michigan
Ann Arbor, Michigan USA

Learning Objectives

After this lecture you should have an understanding of:

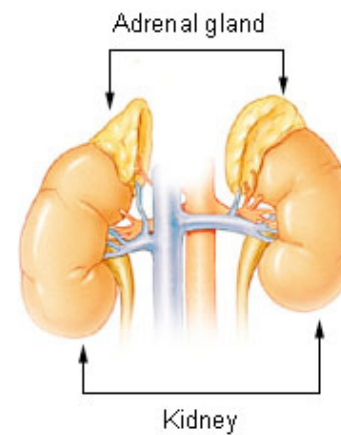
- The feedback loops regulating cortisol secretion.
- The physiologic actions of glucocorticoids (cortisol) + mineralocorticoids (aldosterone)
- The major pharmacologic uses of glucocorticoids.
- The major types of glucocorticoids.
- The major side effects of glucocorticoid therapy.

Anatomy of the adrenal glands

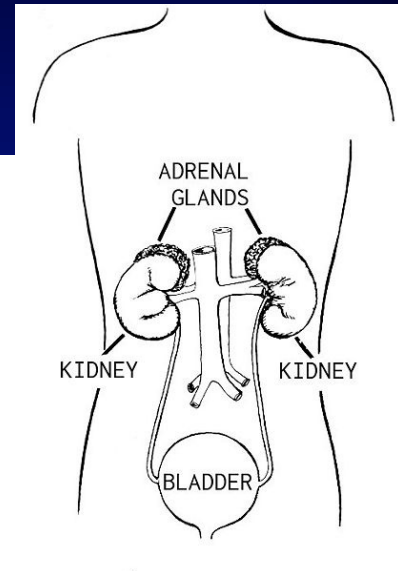


Public Domain
[Wikimedia Commons](#)

Adrenal Gland



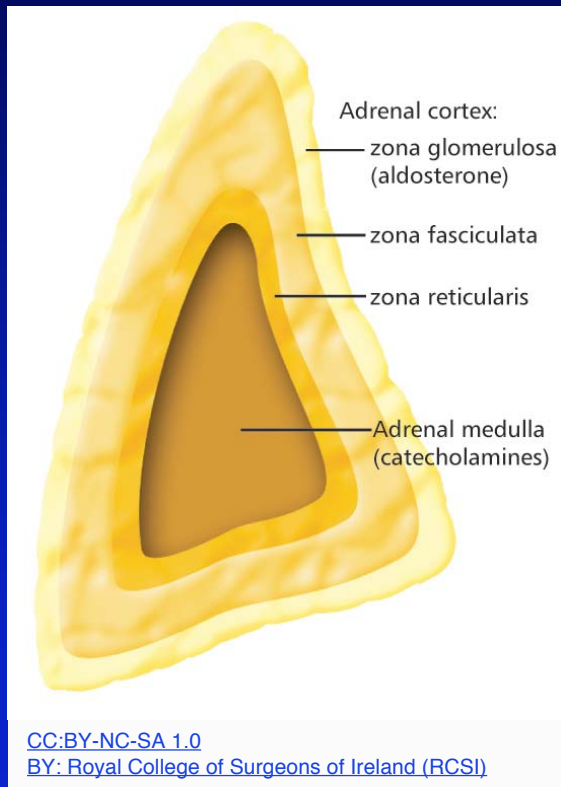
Public Domain
[Seer's.gov](#)



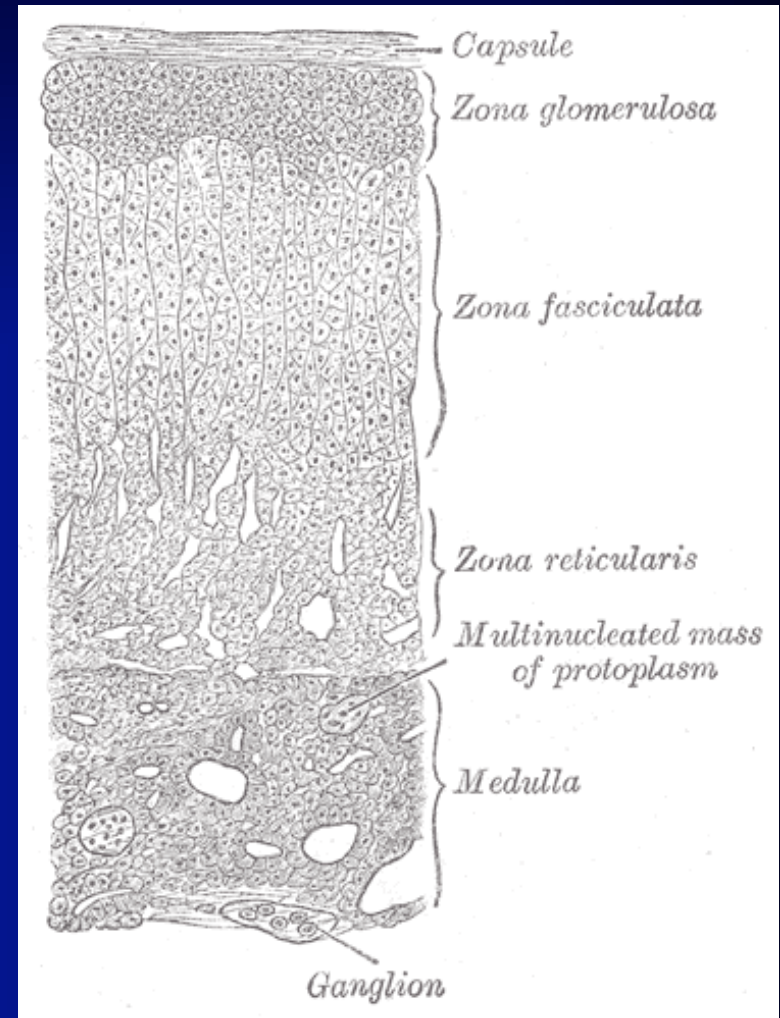
Public Domain
[Wikimedia Commons](#)

Histology of the Adrenal Gland

adrenal cortex



adrenal medulla



Public Domain
[Wikimedia Commons](#)

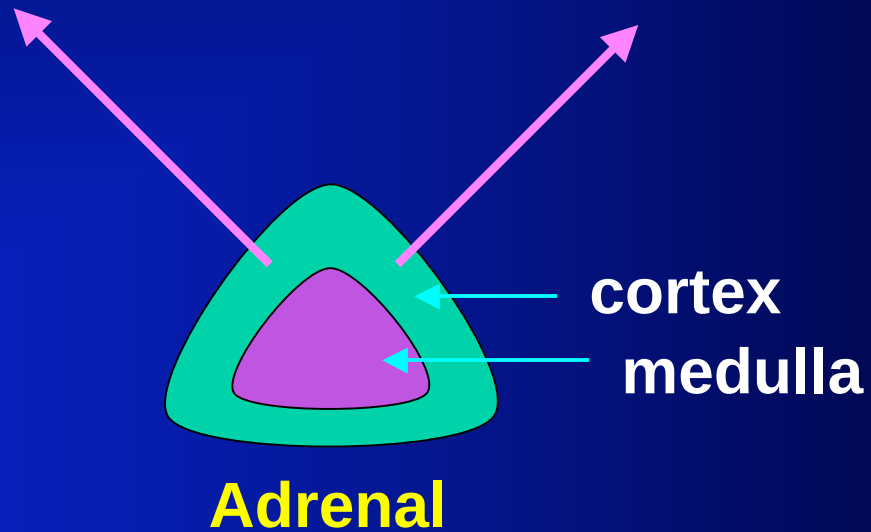
Adrenocortical Hormones = Steroids

GLUCOCORTICOID

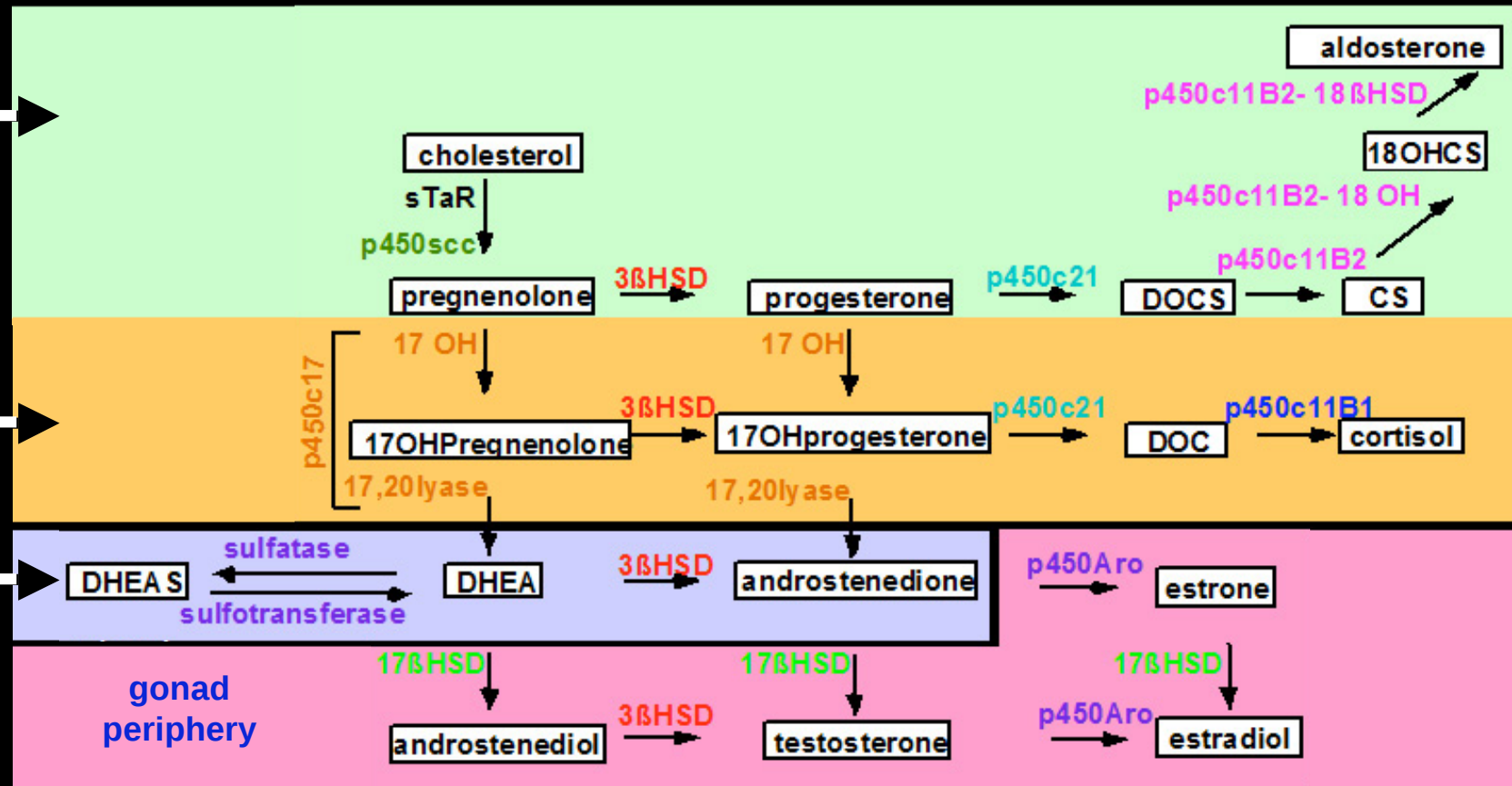
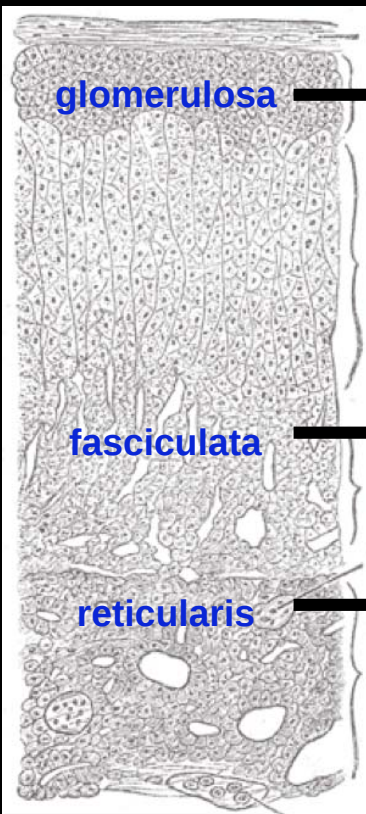
MINERALOCORTICOID

Cortisol

Aldosterone



Steroidogenesis



'Roids: The Bottom Line

**In the right amounts, steroids can be the body's
best friend....**

or

in the wrong amounts, the body's worst enemy....

Role of Glucocorticoids in Human Physiology

In the right amounts, glucocorticoids keep:

- Your blood pressure up (maintain cardiovascular stability).**
- Your blood sugar up (maintain metabolic homeostasis).**
- Your disposition sunny (maintain integrity of CNS function).**
- Your temperament cool (regulate response to stress).**

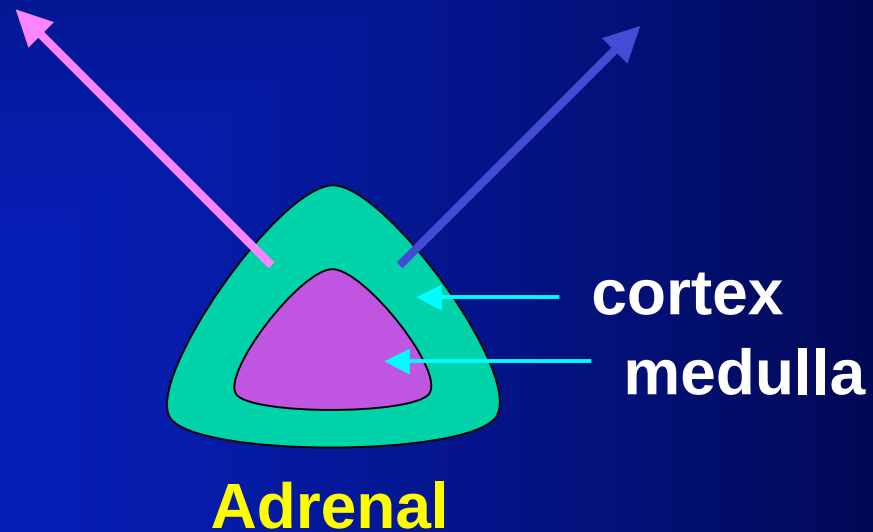
Adrenocortical Hormones = Steroids

GLUCOCORTICOID

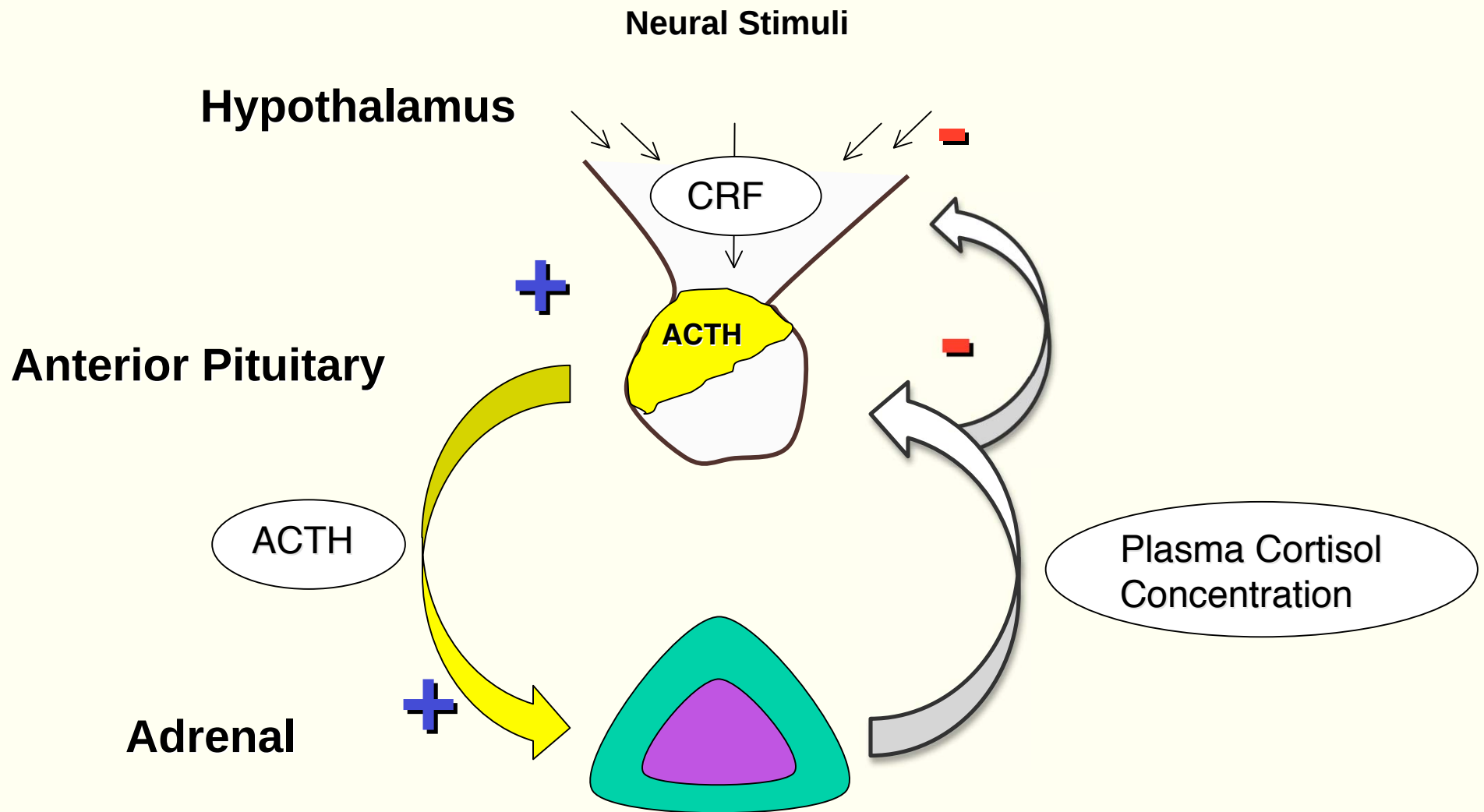
Cortisol

MINERALOCORTICOID

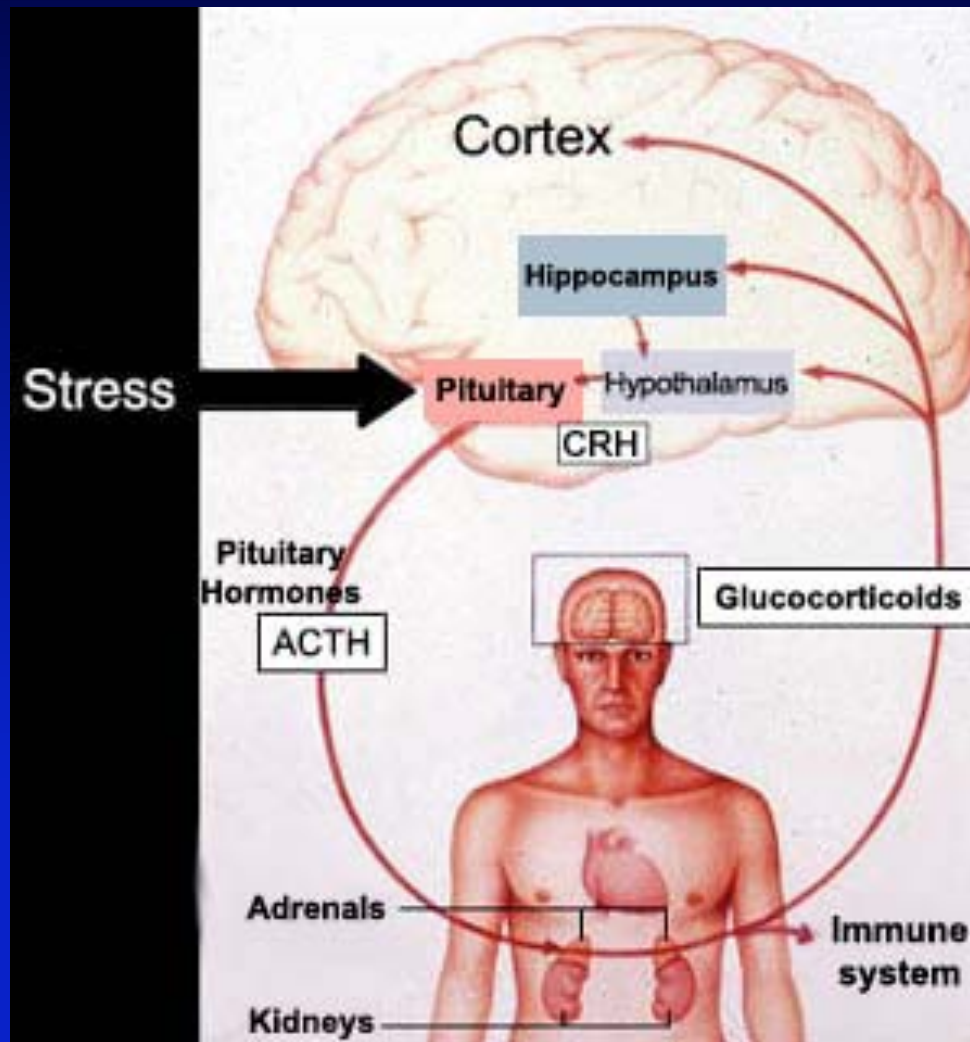
Aldosterone



The HPA axis

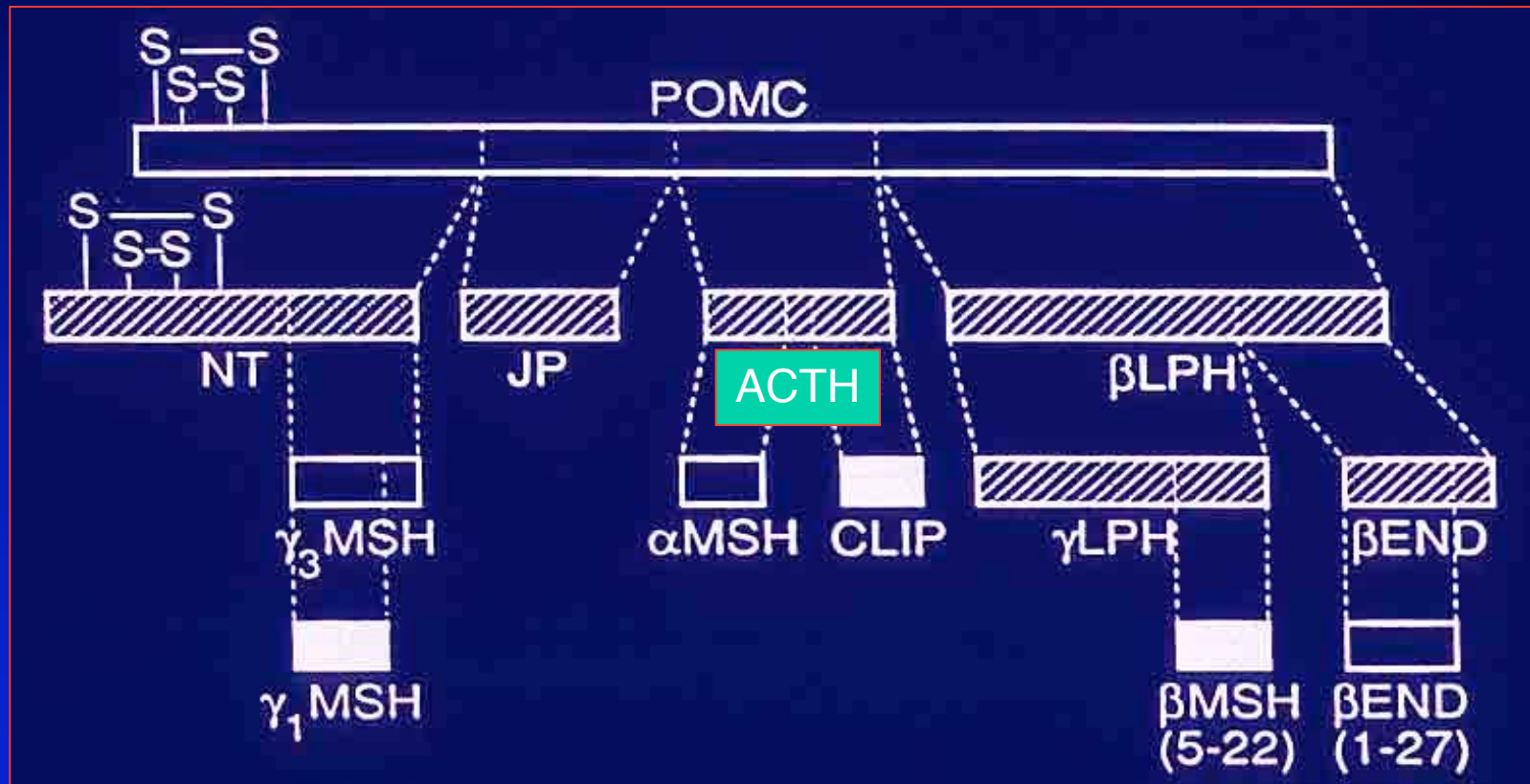


Regulation of ACTH Expression by CRH



Post-translational Processing of POMC in the Normal Pituitary

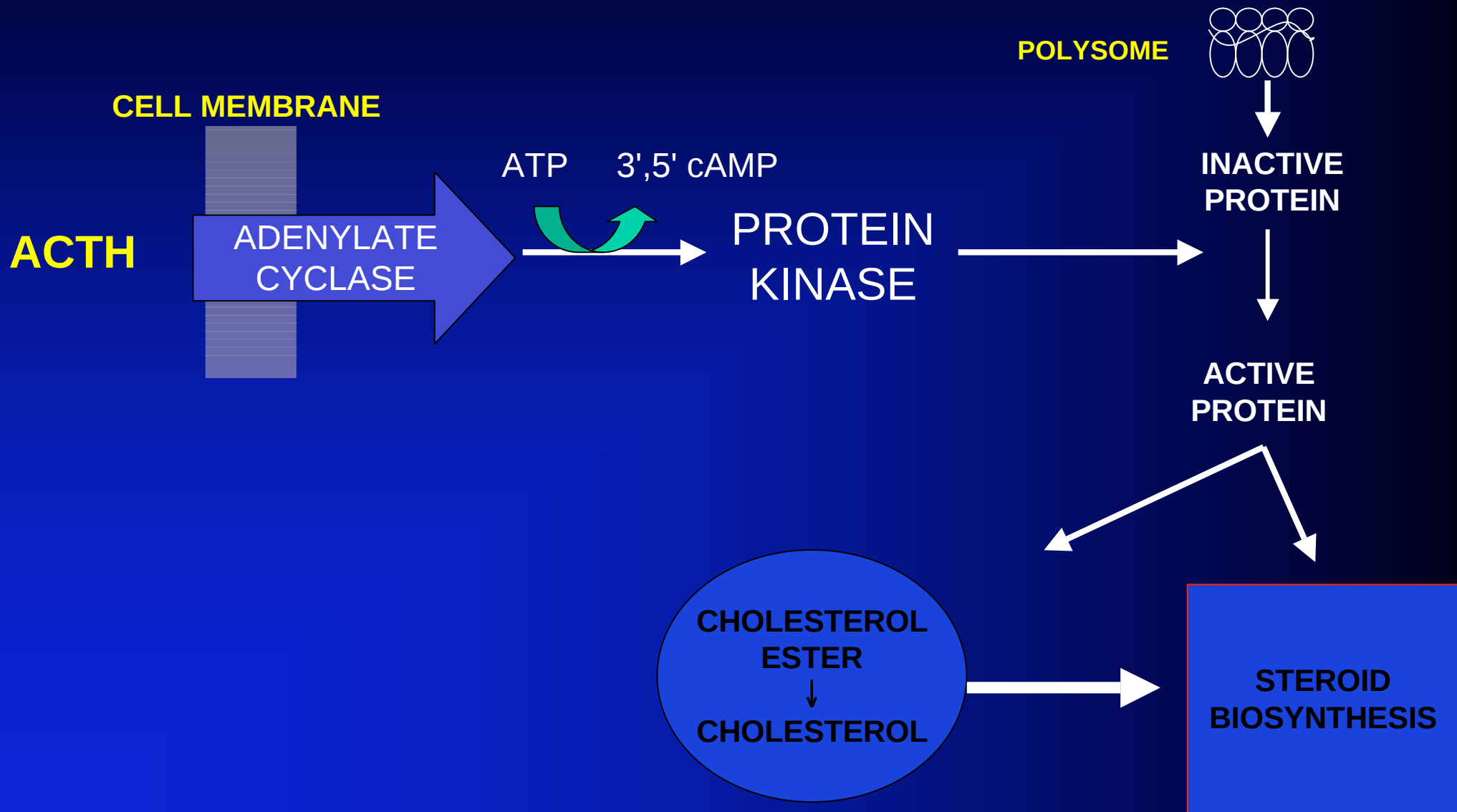
POMC = Pro-opiomelanocortin



Source: Undetermined

MSH = Melanocyte stimulating hormone

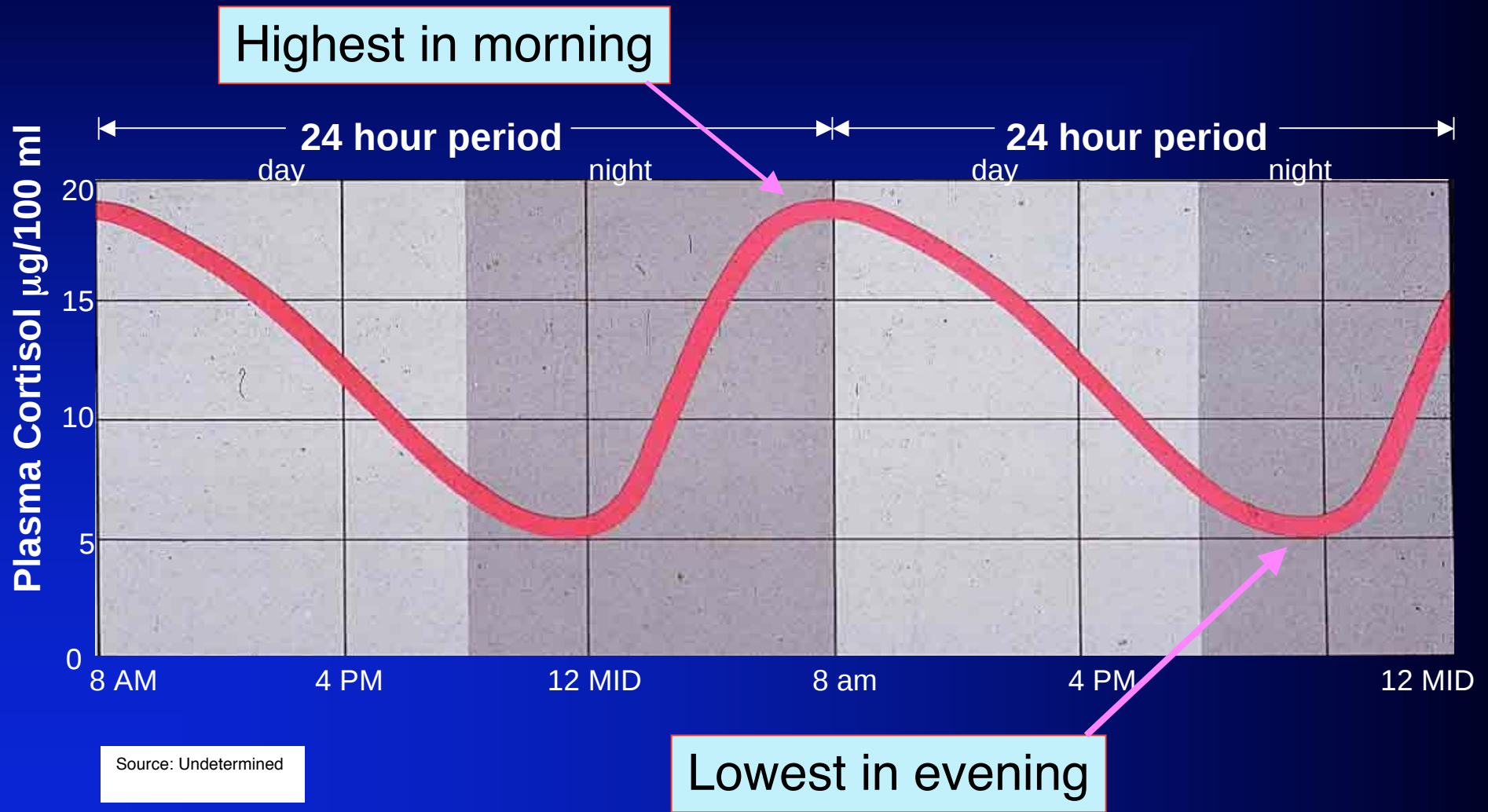
ACTH and Steroid Biosynthesis



Secretion, Transport and Metabolism of Cortisol



Circadian Rhythm of Cortisol Secretion



Corticosteroid Binding Globulin (CBG)



SHBG Grishkovskaya et al, 1999

- Acidic glycoprotein MW 52,000
- Produced in liver, lung, kidney, testes
- Regulates delivery of cortisol to tissues

Conditions that Affect Cortisol Metabolism

- **Increased Turnover:**
 - Thyroxine
 - Barbiturates
 - Phenytoin
- **Decreased Turnover:**
 - Liver disease
- **Increased Binding:**
 - Estrogens

Molecular Action of Glucocorticoids

Glucocorticoid receptors (GR) are transcriptional activators of a variety of gene products.



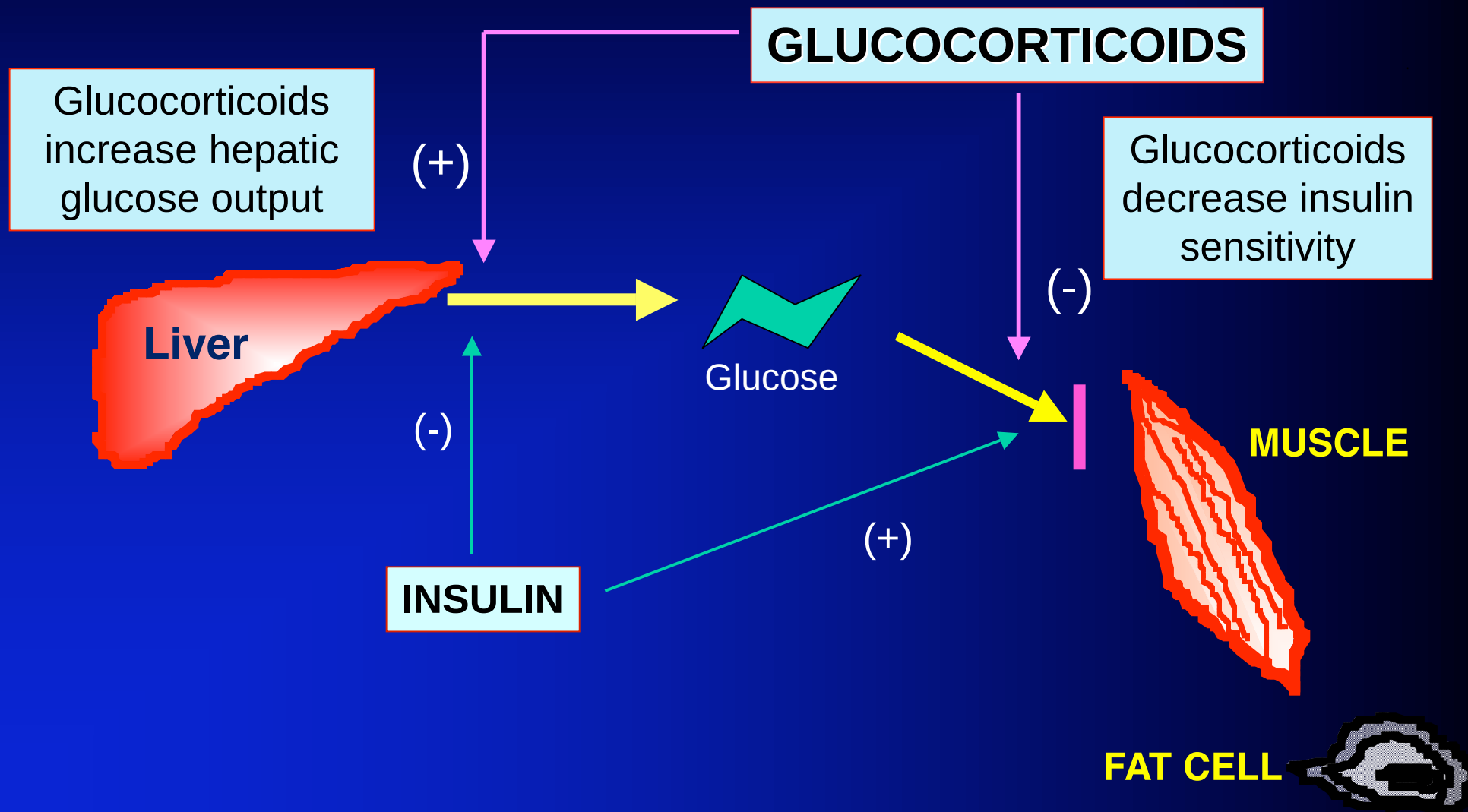
Metabolic Effects of Glucocorticoids

Prototypical Glucocorticoid = Cortisol

Glucocorticoids $\neq$ Insulin

**Glucocorticoids effects are generally
opposite those of insulin.**

Glucocorticoids & Carbohydrate Metabolism



Glucocorticoid Effects on Protein Metabolism

Insulin

- ↑ Anabolism (storage)
- ↑ Protein synthesis
- ↓ Protein breakdown
- ↓ Amino acid release



Glucocorticoids

- ↑ Catabolism
- ↓ Protein synthesis
- ↑ Protein breakdown
- ↑ Amino acid release

Glucocorticoid Effects on Lipid Metabolism

Insulin

- ↑ Anabolism (storage)
- ↑ Lipid synthesis
- ↓ Lipolysis
- ↓ Fatty acid release



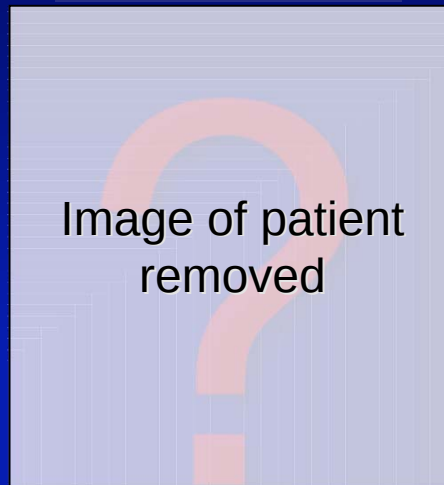
ADIPOCYTE

Glucocorticoids

- ↑ Catabolism
- ↓ Lipid synthesis
- ↑ Lipolysis
- ↑ Fatty acid release

Redistribution of fat

Redistribution of Fat in Glucocorticoid Excess



**Central obesity seen in Cushing's Syndrome
(Glucocorticoid Excess)**

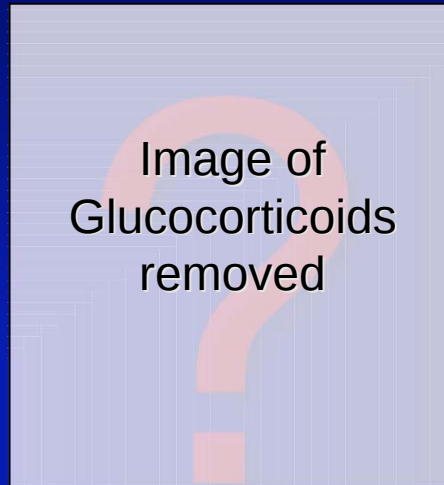
Glucocorticoid Effects on Inflammatory Mediators

Glucocorticoids INHIBIT inflammation.

Inhibit:

- 1) Arachidonic acid and its metabolites (prostaglandins; leukotrienes)
- 2) Platelet activating factor (PAF)
- 3) Tumor necrosis factor (TNF)
- 4) Interleukin-1 (IL-1)
- 5) Plasminogen activator

Sites of Action of Glucocorticoids in the Responses of Leukocytes During Antigenic Challenge/Inflammation



Glucocorticoids

Clinical Uses of Glucocorticoids

Steroid Therapy: Routes of Administration

- **Systemic**
 - Oral**
 - Parenteral**
- **Topical**
- **Inhalation**

Clinical Uses of Glucocorticoids

- **Replacement therapy**
- **Anti-inflammatory effect**
- **Immunosuppression**
- **Androgen suppression**

Glucocorticoids: Use as Anti-Inflammatory Agents



Source: Undetermined

Severe RA of hands

Emily Janz, a 36-year old woman presents with a 3-year history of rheumatoid arthritis. The disease has been progressive with involvement of PIP joints in both hands, wrists, elbows and TM joints. Treatment with non-steroidal anti-inflammatory drugs (NSAIDs) has not been successful.

Treatment with prednisone is begun using an alternate-day program.

Glucocorticoids: Use in Immunosuppression

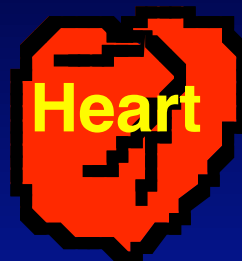


[CC-BY-ND 2.0](#)
[BY: wolfpix](#)

A 25-year old man was walking through a field when he was stung by an insect. He developed generalized edema, dyspnea, wheezing and dizziness. He was rushed by a friend to the emergency room, where a diagnosis of anaphylactoid reaction to insect bite was made.

He received a large dose of steroids parenterally and was subsequently advised on a program to taper the steroids over the next one week.

Glucocorticoids: Use in Immunosuppression



James Allen, a 55-year old man with a history of ischemic cardiomyopathy develops increasingly severe congestive heart failure. When he becomes totally incapacitated with a life-expectancy of less than 6 mo., he is placed on the cardiac transplantation list.

Two months later, he receives a heart and is subsequently placed on an immunosuppressive “cocktail” that includes prednisone, 5 mg daily.

Glucocorticoids: Use in Androgen Suppression



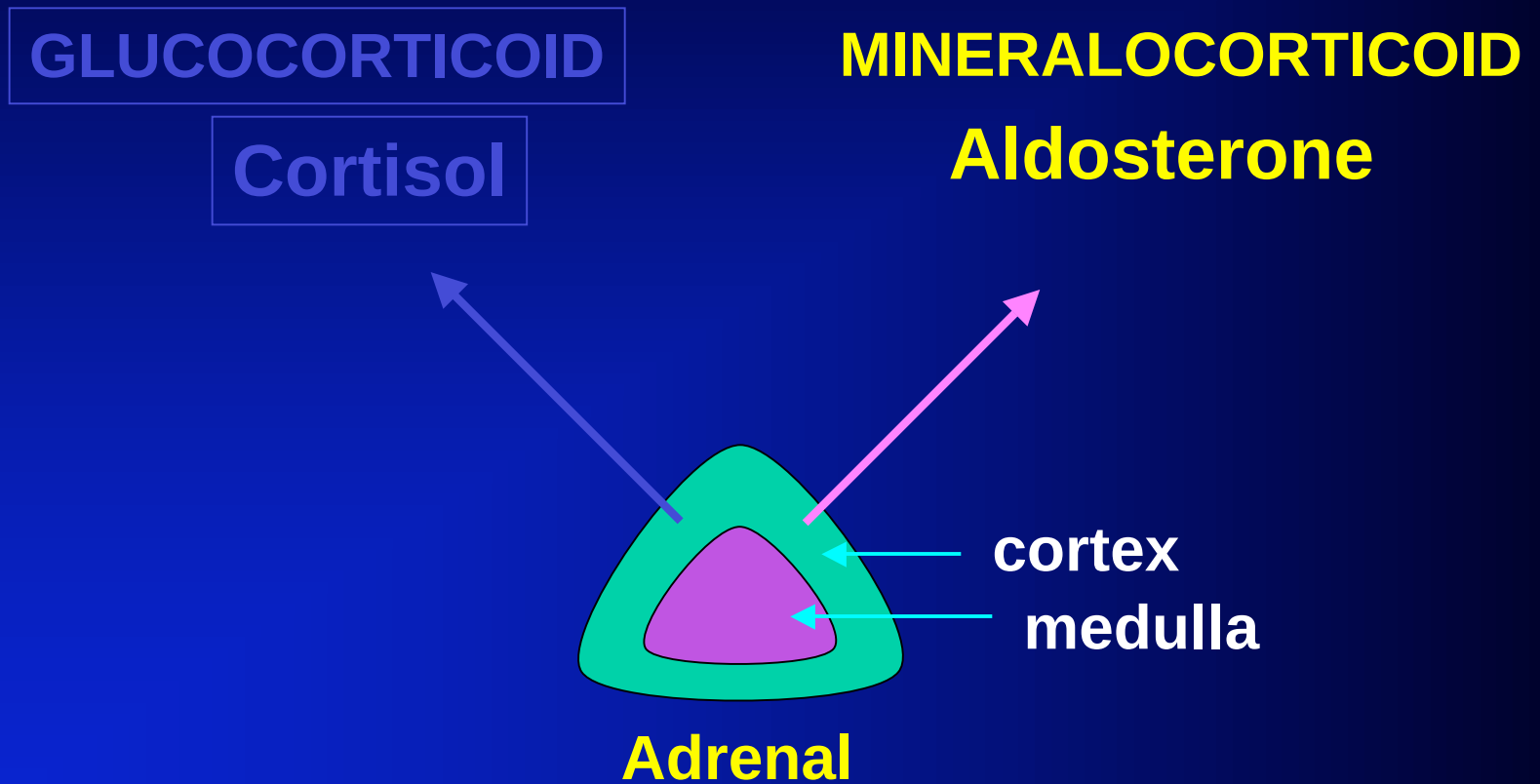
[CC:BY-NC-SA](#)
[BY: C. Matthew Peterson, MD](#)

Hirsutism in a young woman

A 25-year old woman comes in for evaluation of hirsutism present over the past 3 years. The hirsutism is of the androgen type and is associated with acne and irregular menses. Diagnostic studies reveal elevated serum dehydroepiandrosterone (DHEA) and testosterone levels.

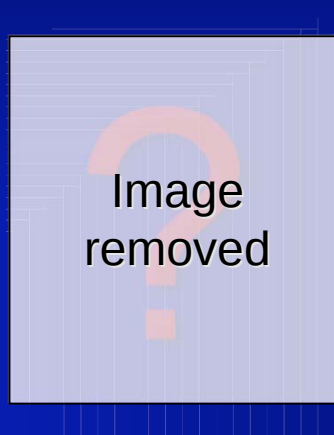
She receives Dexamethasone 2.0 mg daily, for seven days and serum DHEA and testosterone levels are measured the 8th day.

Adrenocortical Hormones = Steroids



Effects of Mineralocorticoid on Renal Tubule

Prototypical mineralocorticoid = Aldosterone



Aldosterone increases sodium resorption and potassium and hydrogen ion excretion.

Prototype of Steroid Compounds



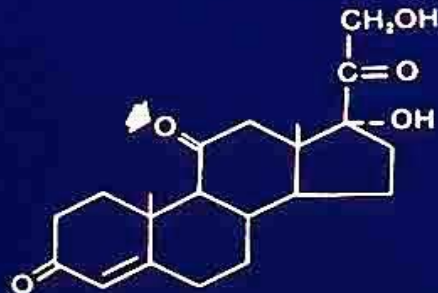
**CORTISOL
(HYDROCORTISONE)**



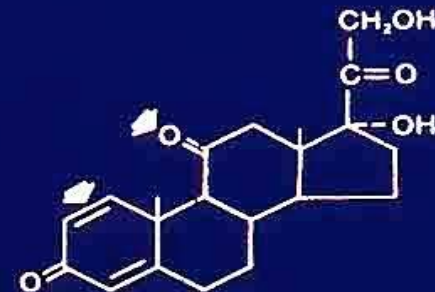
PREDNISOLONE



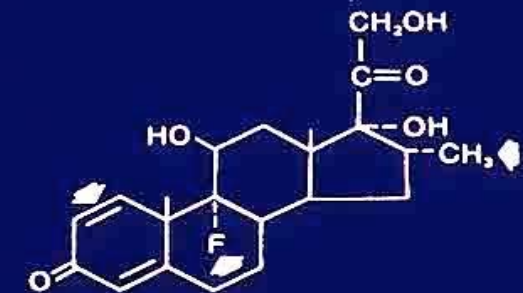
METHYL PREDNISOLONE



CORTISONE

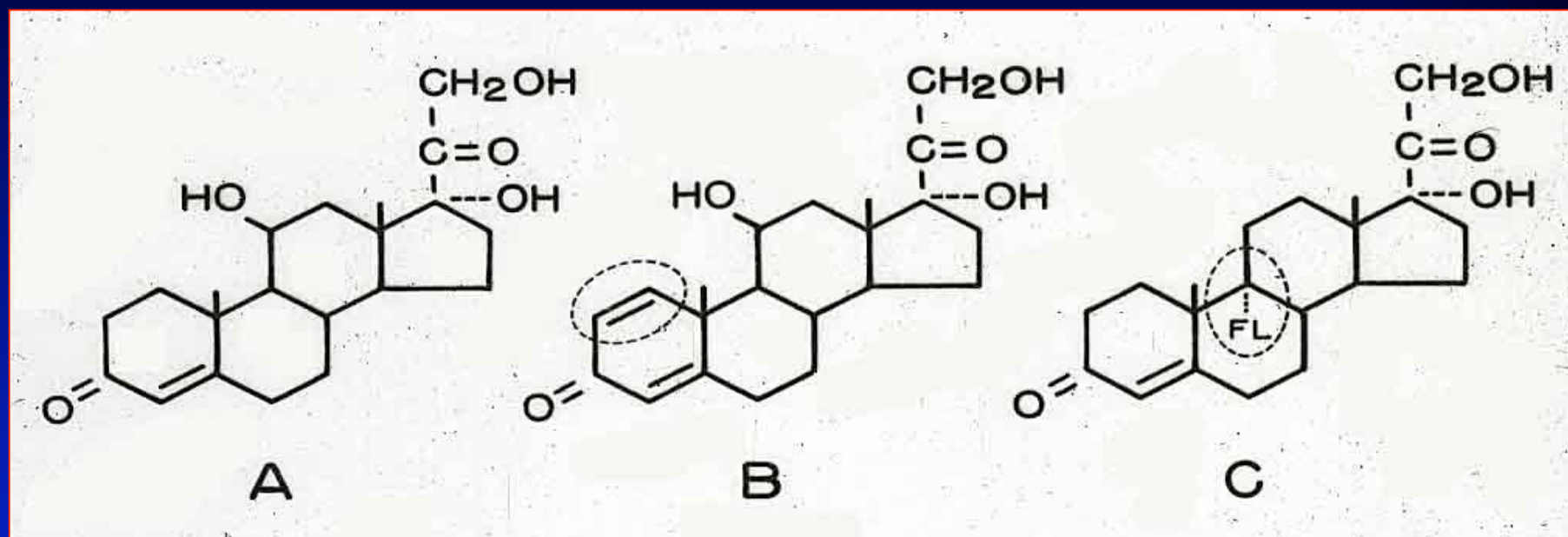


PREDNISONE



DEXAMETHASONE

Steroids: Structure-function Relationships



A. Hydrocortisone B. Prednisone C. 9- α -Fluorocortisol

Source: Undetermined

- Double-bond in 1,2 position increases glucocorticoid activity.
- Fluoro- group in 9- α position increases mineralocorticoid activity.

Steroids

Compound	Anti-Inflam. potency	Na-Retain. potency	Duration of action	Equivalent Dose
Cortisol	1	1	Short	20 mg
Prednisone	4	0.8	Intermediate	5 mg
9- α -fluoro- cortisone	10	125	Short	*
Dexa- methasone	25	0	Long	0.75 mg

Glucocorticoid effects: Dex > Prednisone > Cortisol

Mineralocorticoids: 9- α -fluorocortisone RULES

Glucocorticoid Therapy

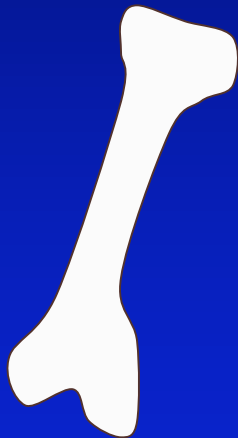
Side Effects

Or

“Yes, Virginia, there can be at times
‘Too Much of a Good Thing...’”

Glucocorticoid Effects on Calcium & Bone

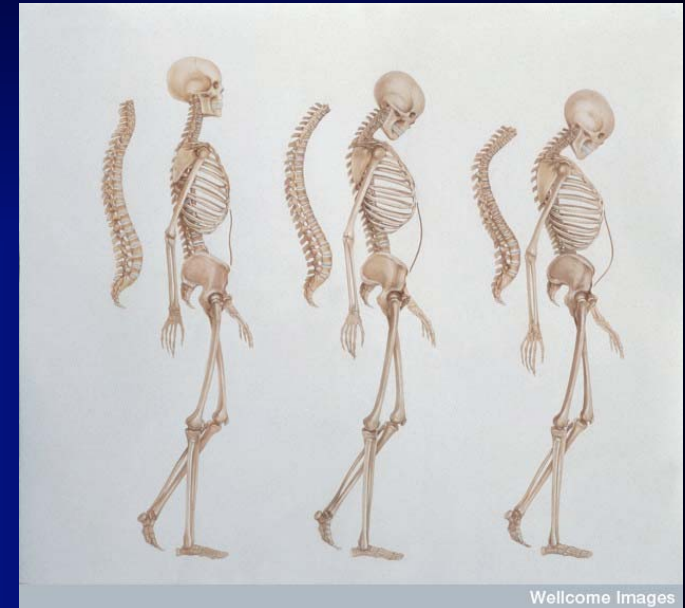
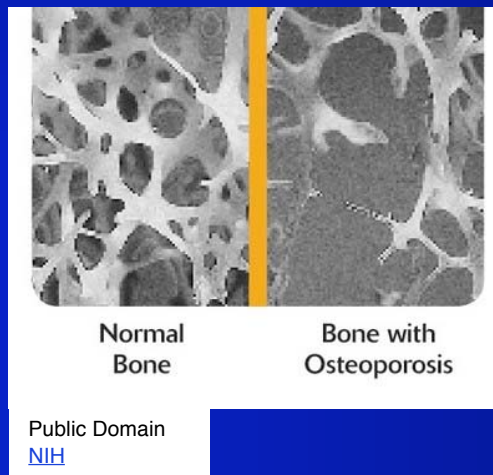
STERIODS → **“BRITTLE BONES”**



- ↓ Osteoblastic activity
- ↓ Calcium absorption from gut.
- ↑ PTH secretion
- ↑ Osteoclastic activity

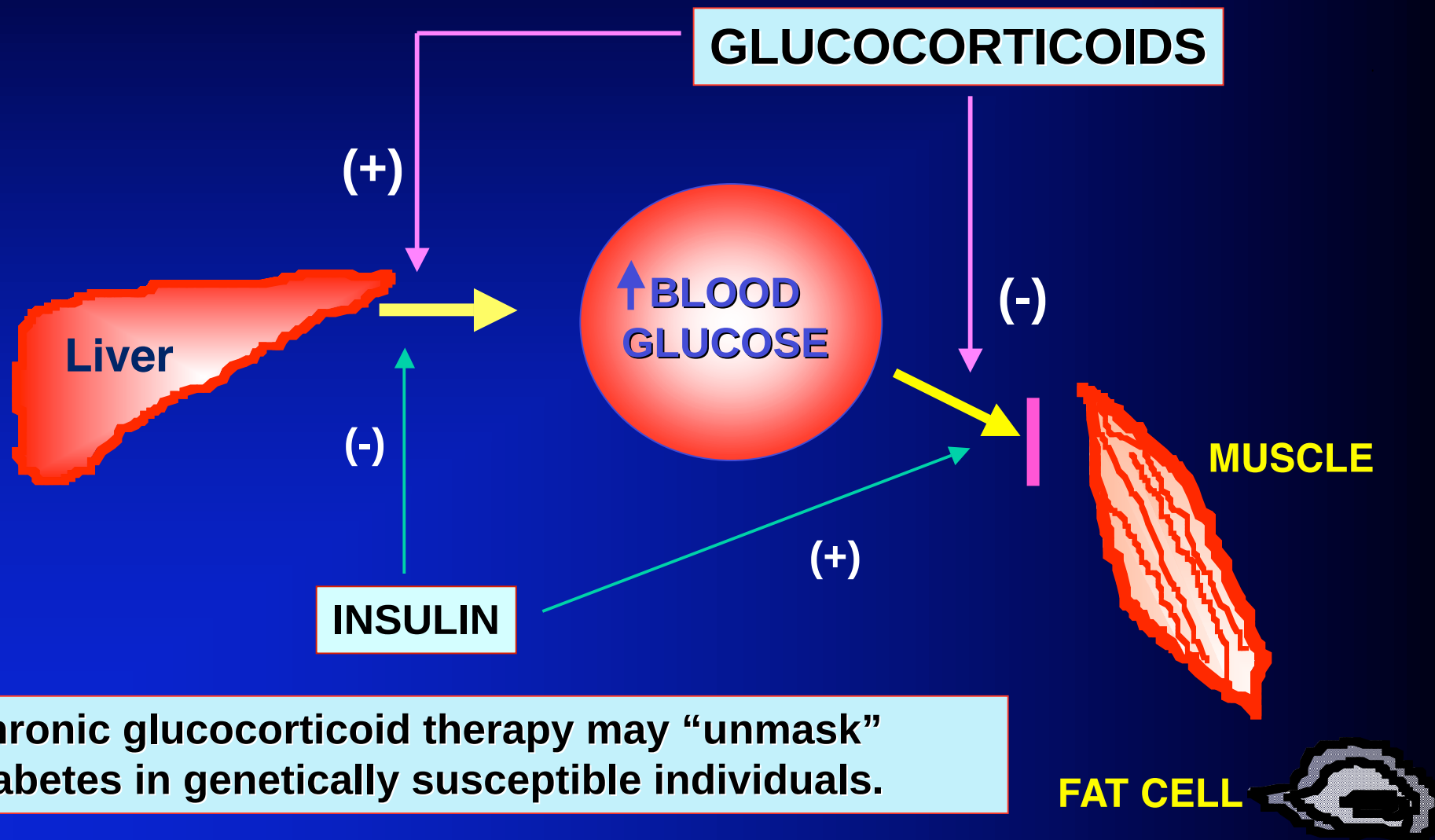
Steroids → “Brittle Bones”

A 60 yo postmenopausal woman was seen in clinic with acute onset of mid-thoracic back pain. She had complained of back pain for the past 2 years and a 2” loss of height. She had been on Prednisone, 10-15 mg daily, for the past 5 years for chronic polymyositis. Radiographic exam of the spine shows compression deformities in several vertebral bodies.

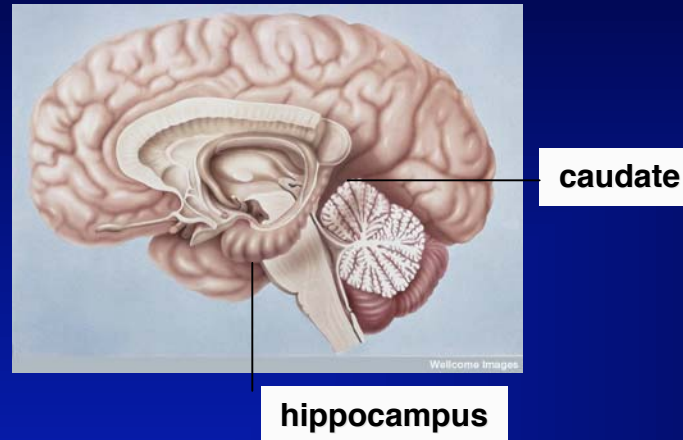


[CC:BY-NC-ND 2.0](#)
[BY: Wellcome Images](#)

Chronic Glucocorticoid Therapy & Carbohydrate Metabolism



Glucocorticoid Effects on the Central Nervous System



- Neuronal death or atrophy
- Structures affected: Hippocampus, caudate
- Neuropsychiatric symptoms:
 - Cognitive- memory, learning
 - Mood- irritability, depression
 - Sleep- insomnia

“Steroid Psychosis”

A 42-yo woman with an exacerbation of lupus nephritis was treated with high-dose prednisone for several days. Her nephritis improved markedly; however, she became increasingly euphoric and severely agitated with paranoid ideation and confusion.

Following tapering of the steroid, she returned to her “usual self.”

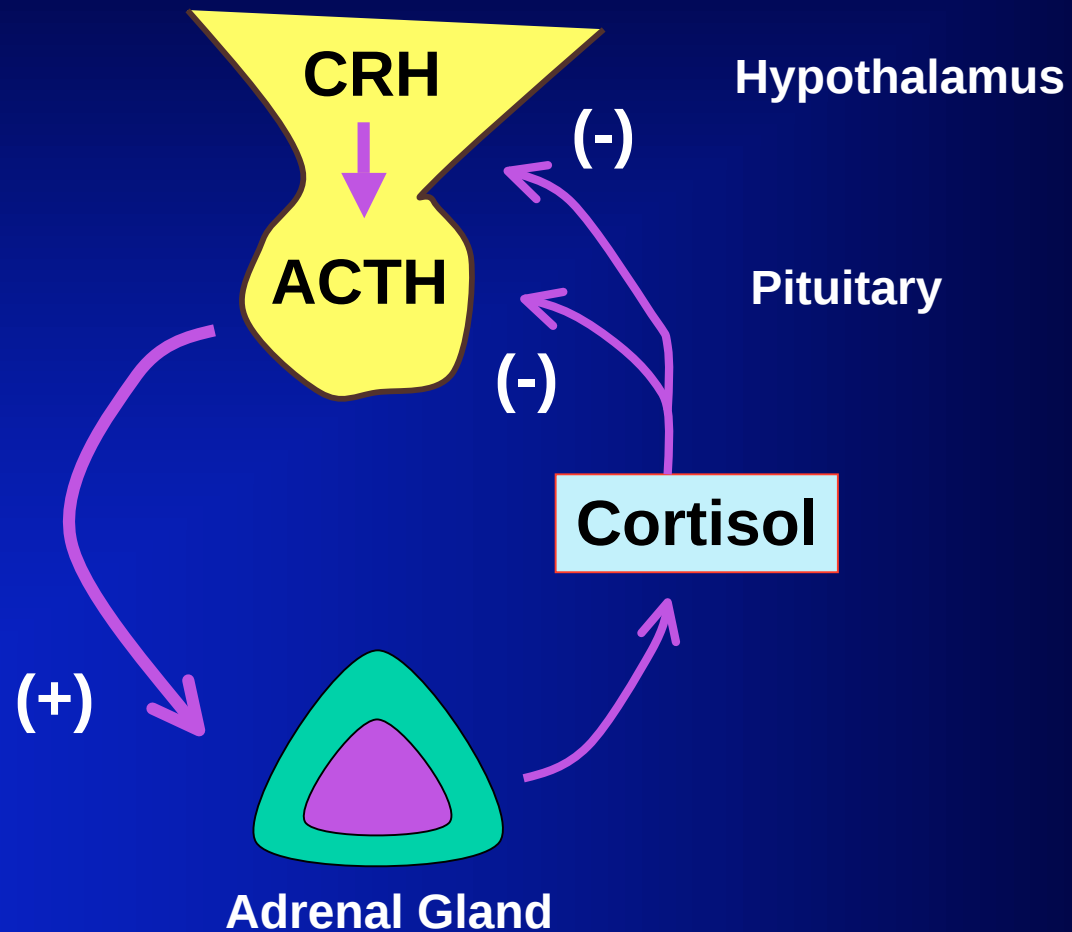
Glucocorticoid Effect on Gastric Function



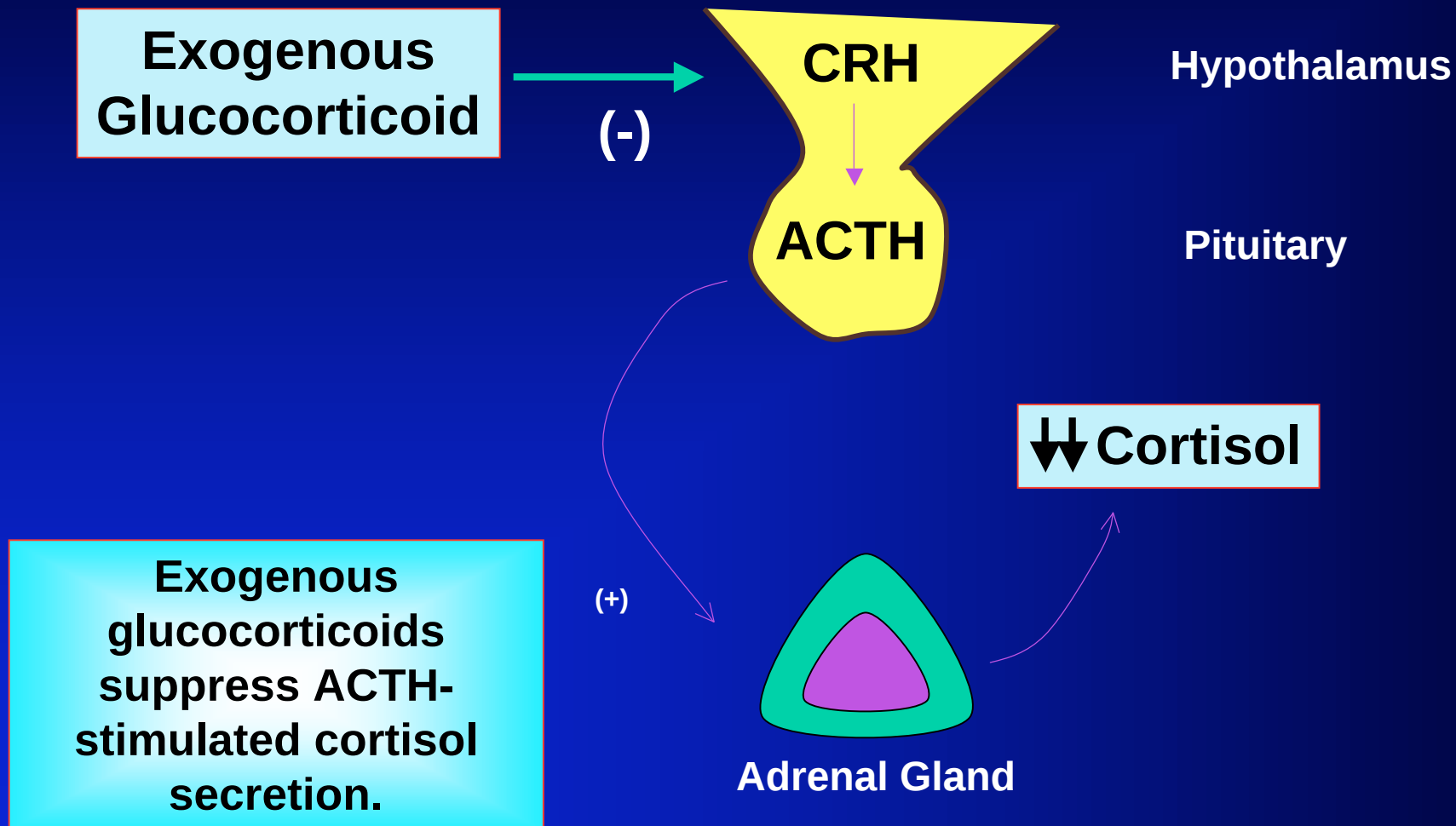
- ↑ Secretion of HCl and pepsin
- ↓ Protective barrier in the gastric mucosa

Glucocorticoid therapy may increase risk of ulcers.

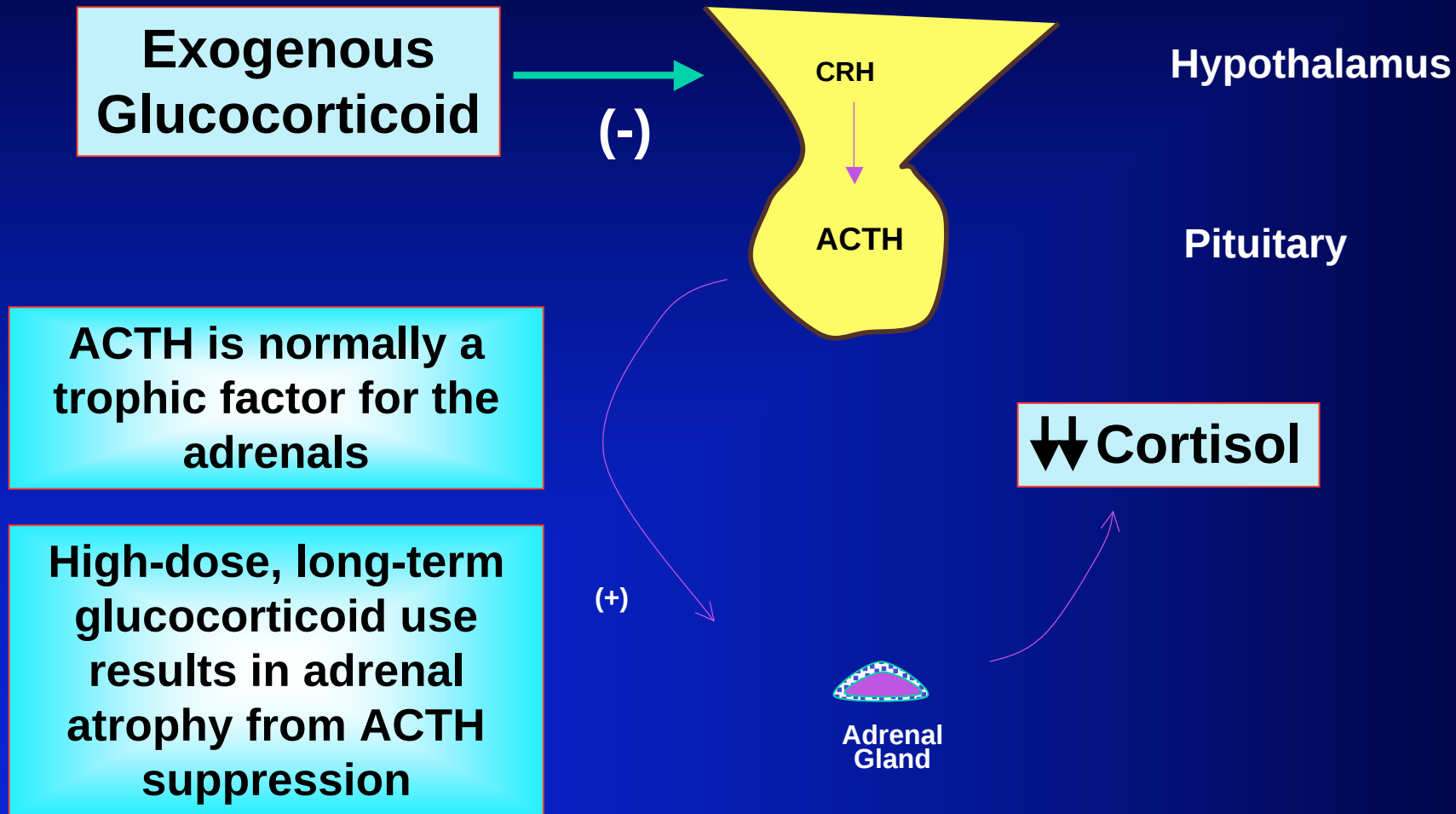
Complications of Chronic Exogenous Corticosteroid Use



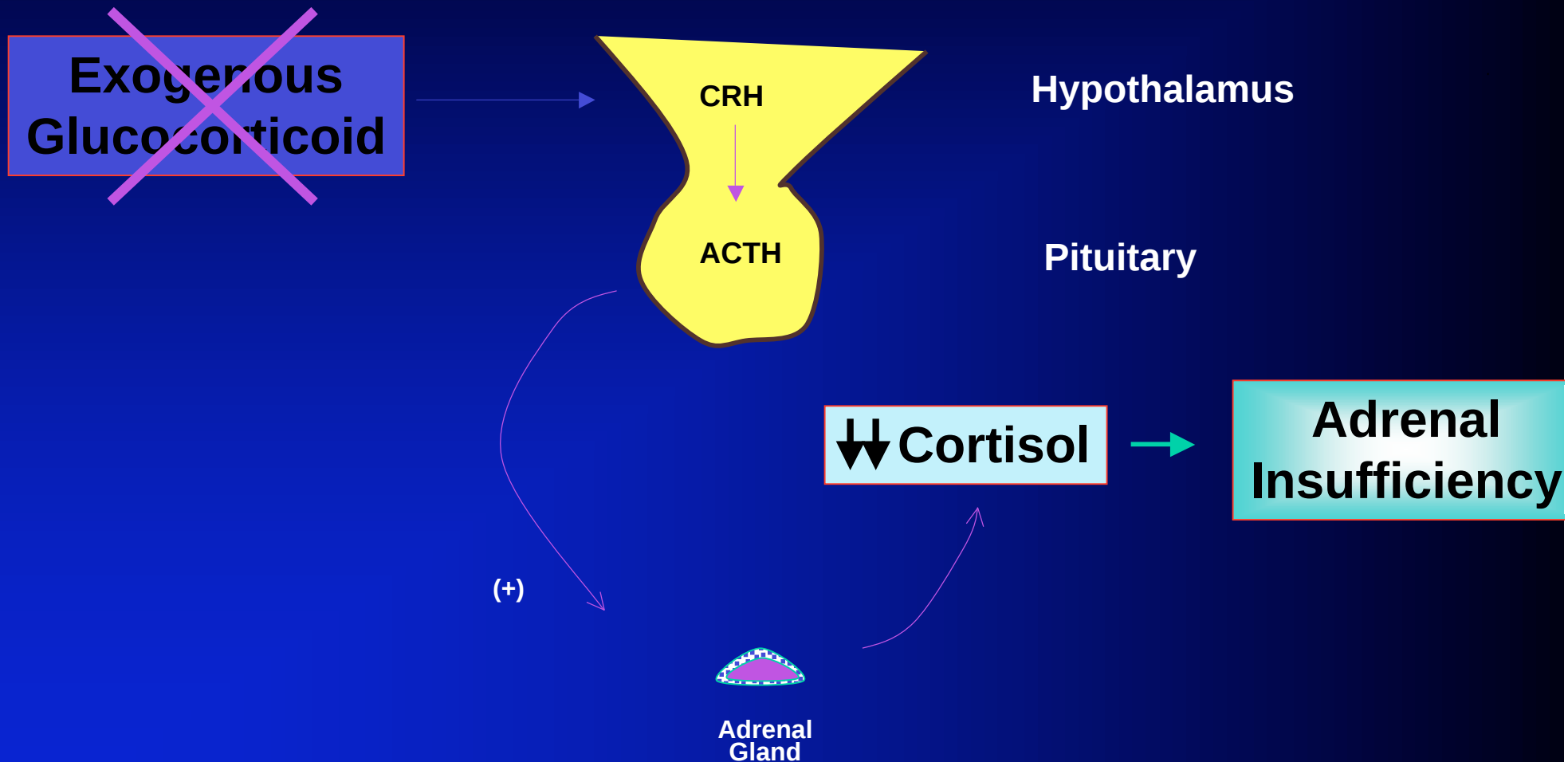
Complications of Chronic Exogenous Corticosteroid Use



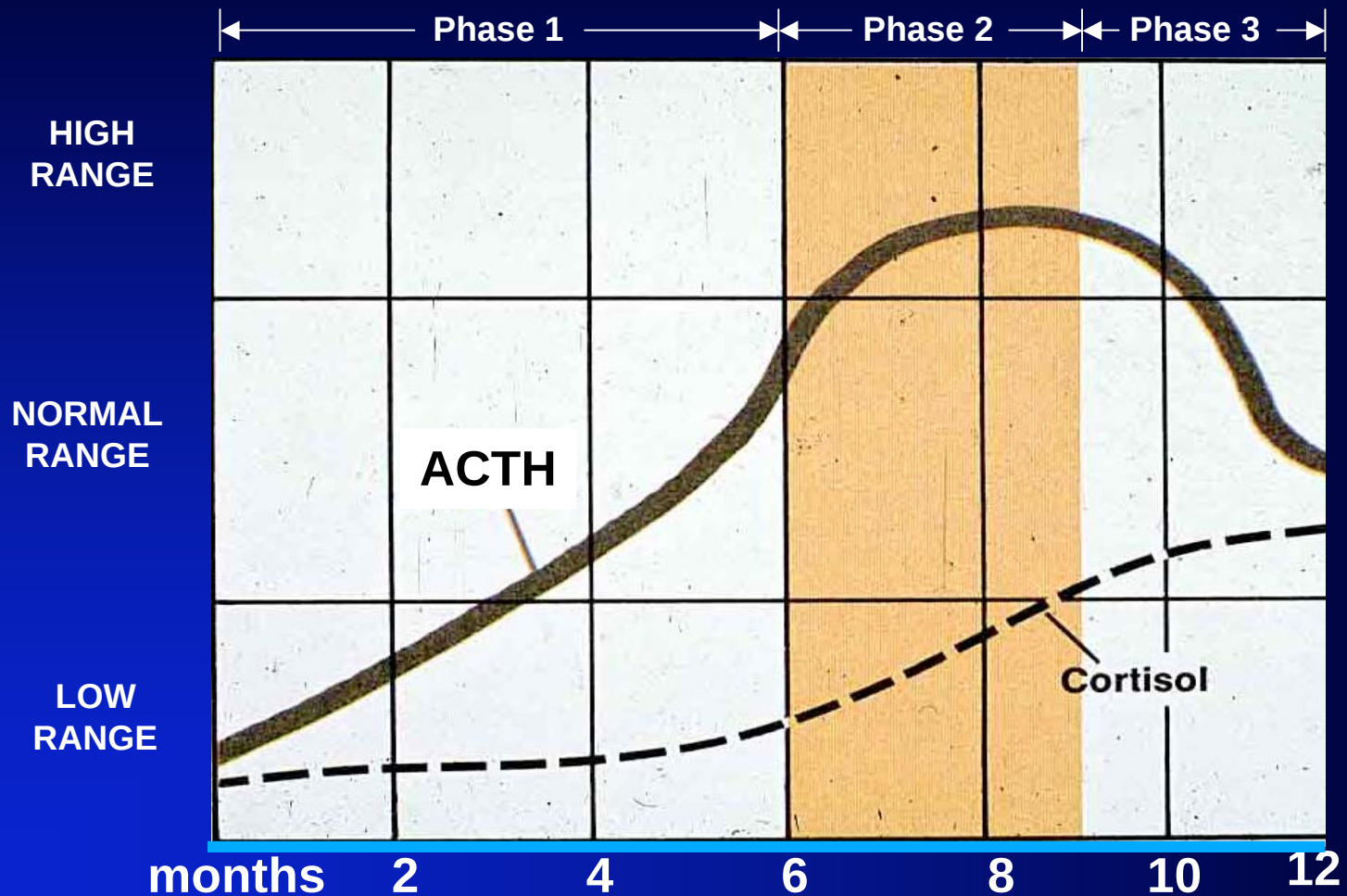
Complications of Chronic Exogenous Corticosteroid Use



Complications of Chronic Exogenous Corticosteroid Use



Recovery of Endogenous Cortisol Secretion Following Withdrawal of Exogenous Steroids



Source: Undetermined

Full recovery of endogenous cortisol secretion may require up to 18 months following steroid withdrawal.

Case #1

A 45-year old woman present with a two-month history of anorexia, nausea, fatigue, dizziness when assuming the upright posture, and increased pigmentation of the skin.

A diagnosis of Addison's disease (Cortisol and Aldosterone deficiency) is confirmed by appropriate testing.

Treatment is initiated with Cortisol 25 mg. (10/10/5) and 9- α fluorocortisol 0.05 mg QD.

Corticosteroid Therapy Considerations

- How serious is the underlying disorder?
- How long is therapy required?
- What is the anticipated effective dose range?
- Is patient predisposed to complications?
- Which preparation to use?
- Alternate day v. every day therapy.
- Program for withdrawal.

Complications with Prolonged Steroid Therapy

- Retarded longitudinal growth in children*
- GI Bleeding
- Osteoporosis*
- Diabetes*
- Cushing's Syndrome
- Steroid myopathy
- Hypertension
- Cataracts
- Psychiatric
- Adrenal suppression*

***Complications to remember**

Things to Remember if Dr. Lash put you to sleep and you're just waking up...:

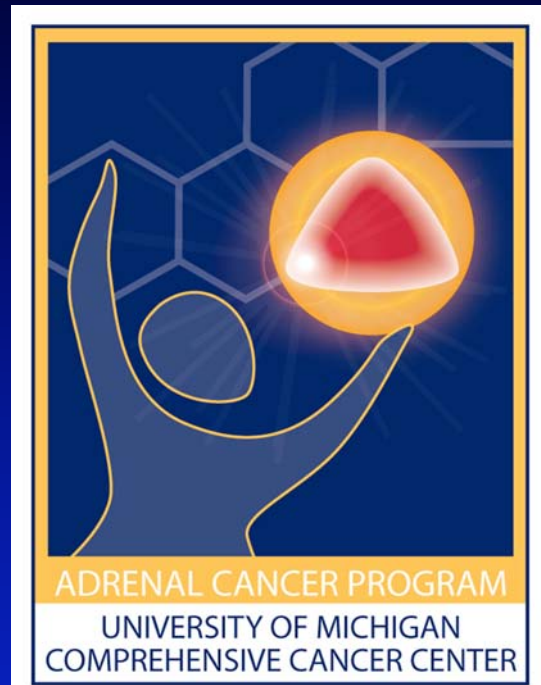
Understand:

- Feedback loops regulating cortisol secretion.
- The major physiologic actions of glucocorticoids (cortisol) and mineralocorticoids (aldosterone).
- The major pharmacologic uses of glucocorticoids.
- The major types of glucocorticoids--hydrocortisone, prednisone, dexamethasone, 9-a-fluorocortisol.
- The major side effects of glucocorticoid therapy.

Adrenal Steroid Physiology & Pharmacology

Questions?

Disorders of the Adrenal Cortex



Logo: All Rights Reserved
Regents of the University of Michigan.

2008

Gary D. Hammer, M.D., Ph.D.
University of Michigan
Ann Arbor, Michigan USA

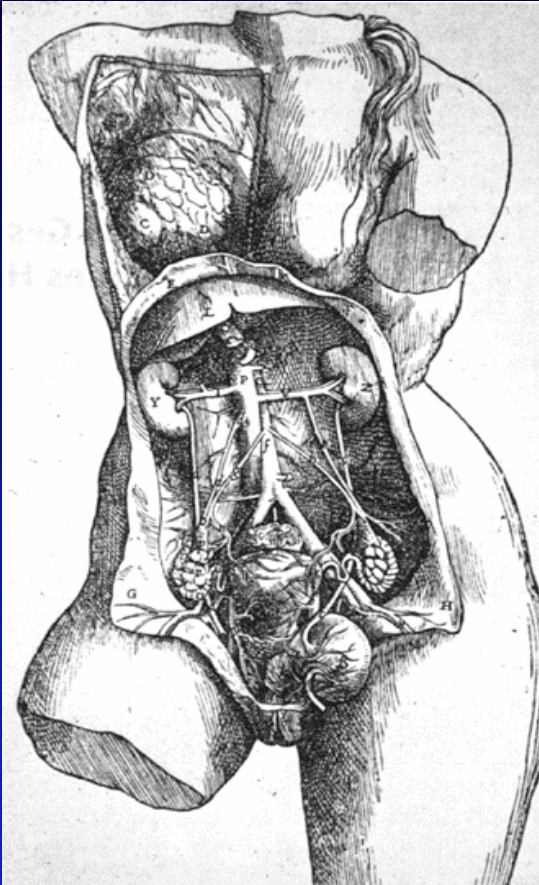
Goals/Objectives

- Remember the basic principles of the HPA axis: homeostatic control of plasma cortisol and aldosterone levels
- Remember the mechanism of action of glucocorticoids and mineralocorticoids
- Understand etiology, clinical features, differential diagnosis, evaluation and therapy of 3 classic adrenal disorders:
 - Adrenal Insufficiency
 - Cushing's Syndrome
 - Primary Hyperaldosteronism

Which Twin is Sick????

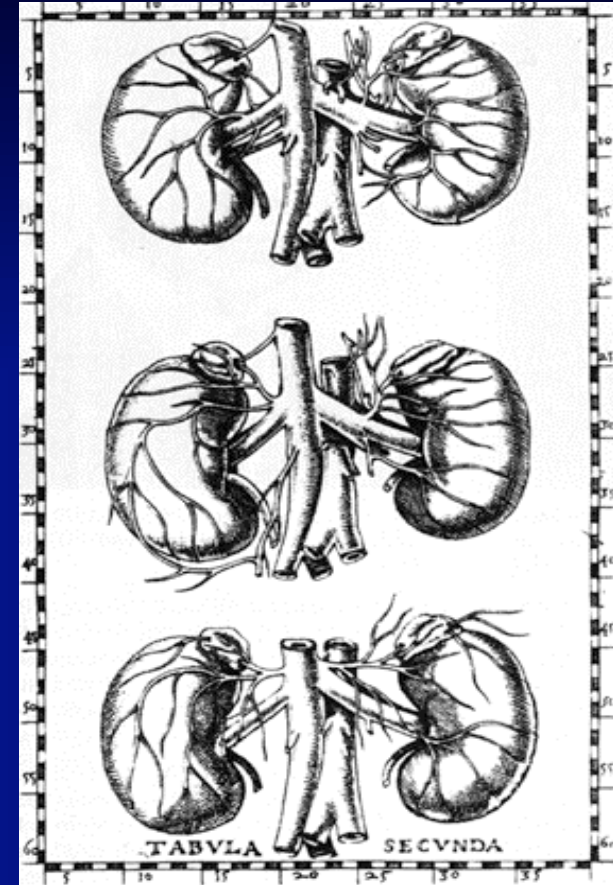


Adrenal Glands in Medical History



Public Domain

Andreas Vesalius (1543)
Book Five of De Corporis Humani Fabrica in 1543



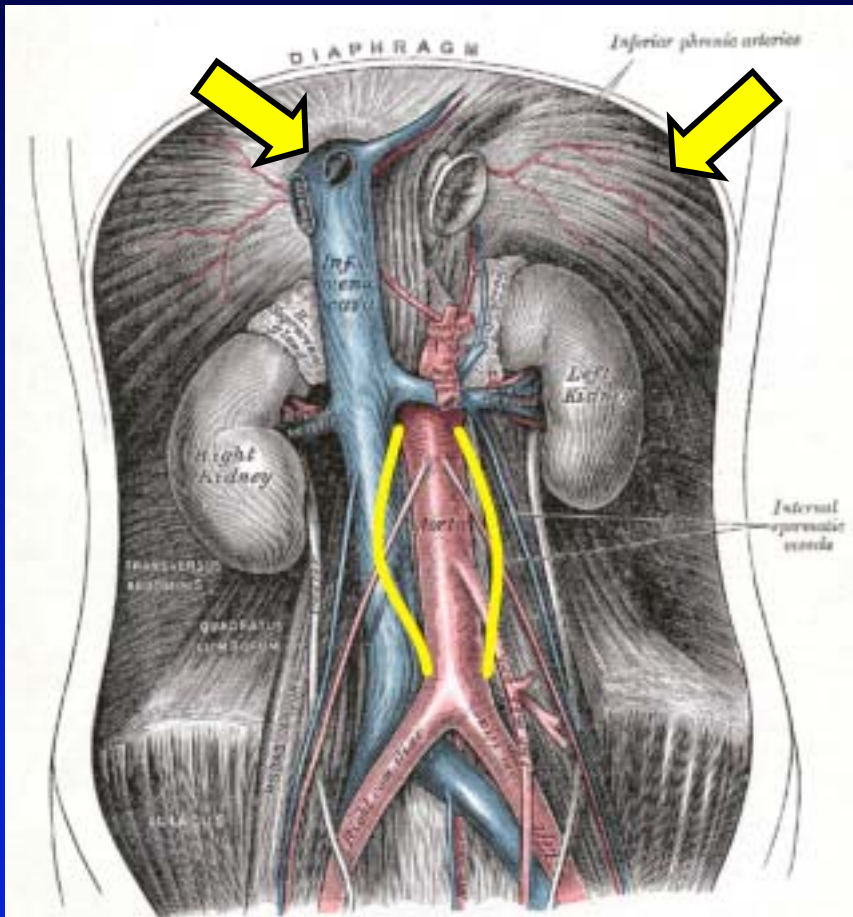
Public Domain

Bartholomäus Eustachius (1564)
"glandulae quae renibus incumbent" in 1564

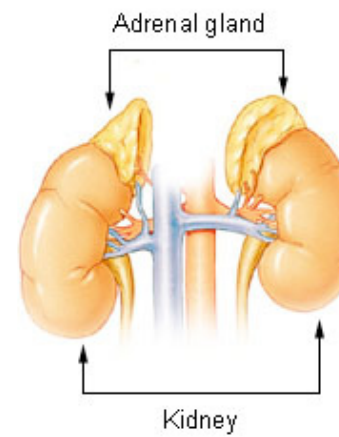
History of Adrenal

- 1716: Academie des Sciences of Bordeaux poses the question
"Quel est l'usage des glandes surrenales?"
- 1845: French thesis on organs of Undetermined function
"The adrenal cease(s) to be a secreting gland."
- 1855: Thomas Addison monograph
"On the constitutional and local effects of disease of the supra-renal capsules," described 10 cases marked by "anemia . . . feebleness of the heart action . . . a peculiar change of color in the skin occurring in connection with a diseased condition of the 'suprarenal capsules'."
- In 1945 Nobel Prize
Kendall, Pfiffner, and Reichenstein first tested adrenal extracts on a patient with Addison's disease, and the response was prompt and striking.

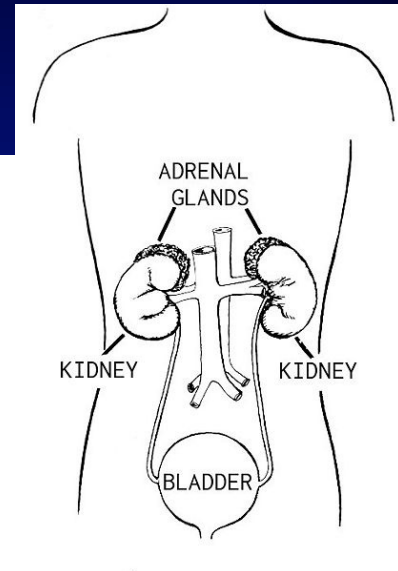
Anatomy of the adrenal glands



Public Domain
[Wikimedia Commons](#)



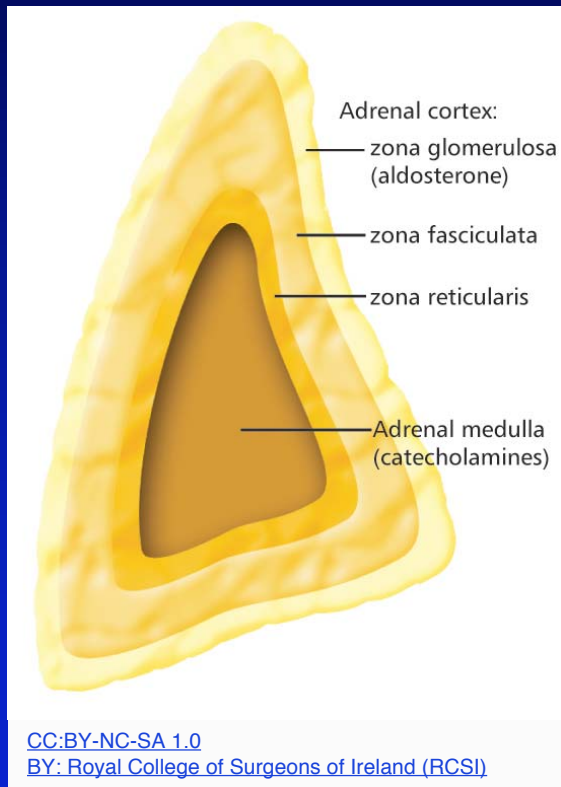
Public Domain
Seer's.gov



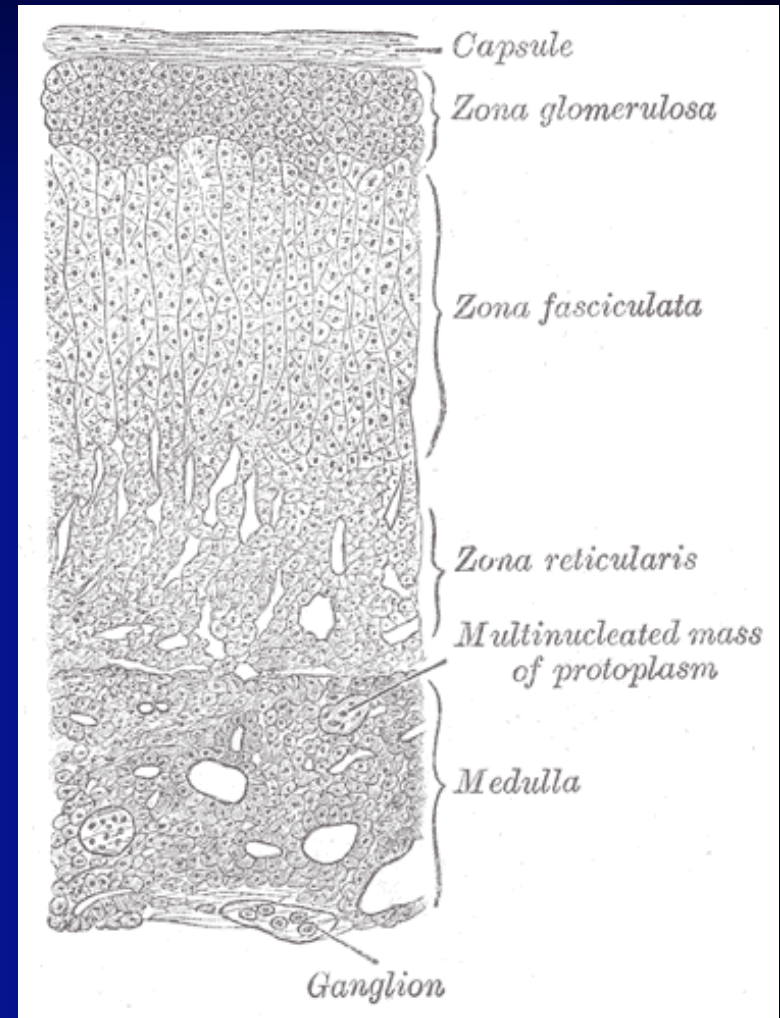
Public Domain
[Wikimedia Commons](#)

Histology of the Adrenal Gland

adrenal cortex



adrenal medulla



Public Domain
[Wikimedia Commons](#)

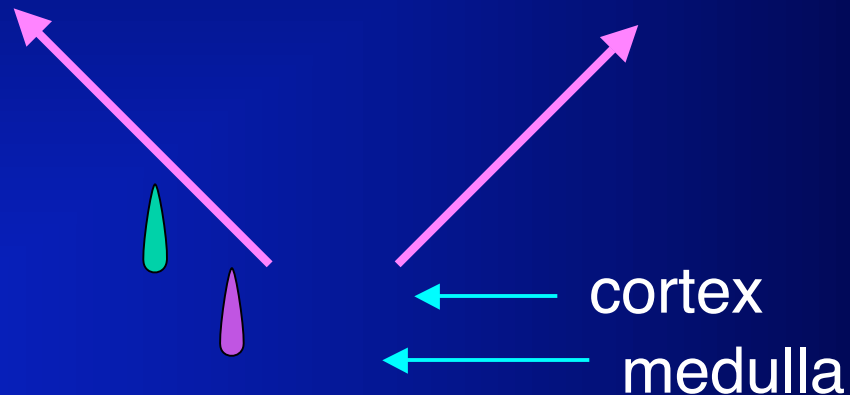
Adrenocortical Hormones = Steroids

GLUCOCORTICOID

Cortisol

MINERALOCORTICOID

Aldosterone



Adrenal

Definition of Adrenal Insufficiency

- “inappropriately low” adrenal steroid output
 - mineralocorticoids (aldosterone)
 - glucocorticoids (cortisol)
 - sex steroids (DHEAS)

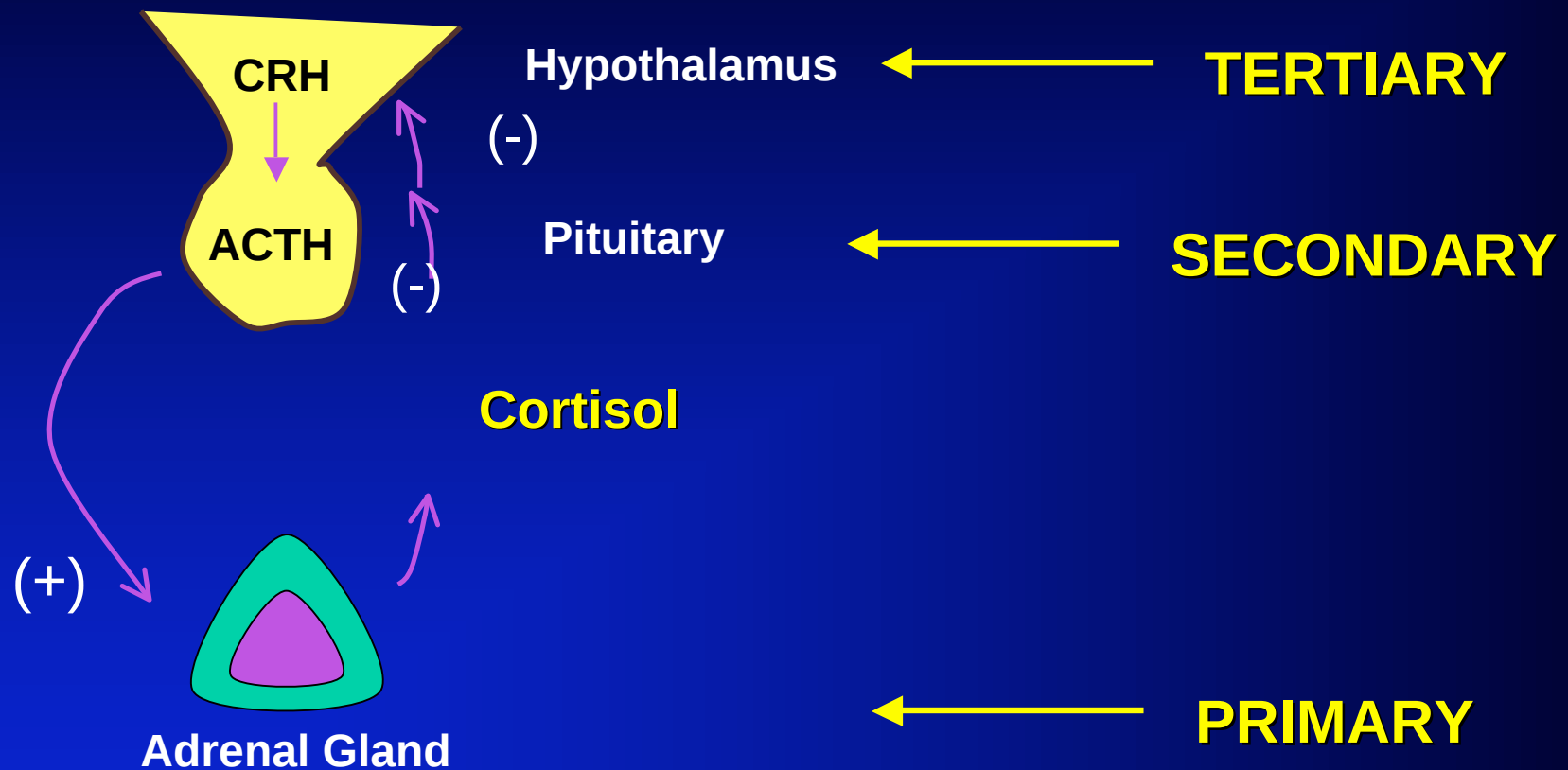
How Frequent Is Adrenal Insufficiency?

- In general, about 40-60 per million individuals have adrenal insufficiency
- 30,000-34,000 people in U.S.

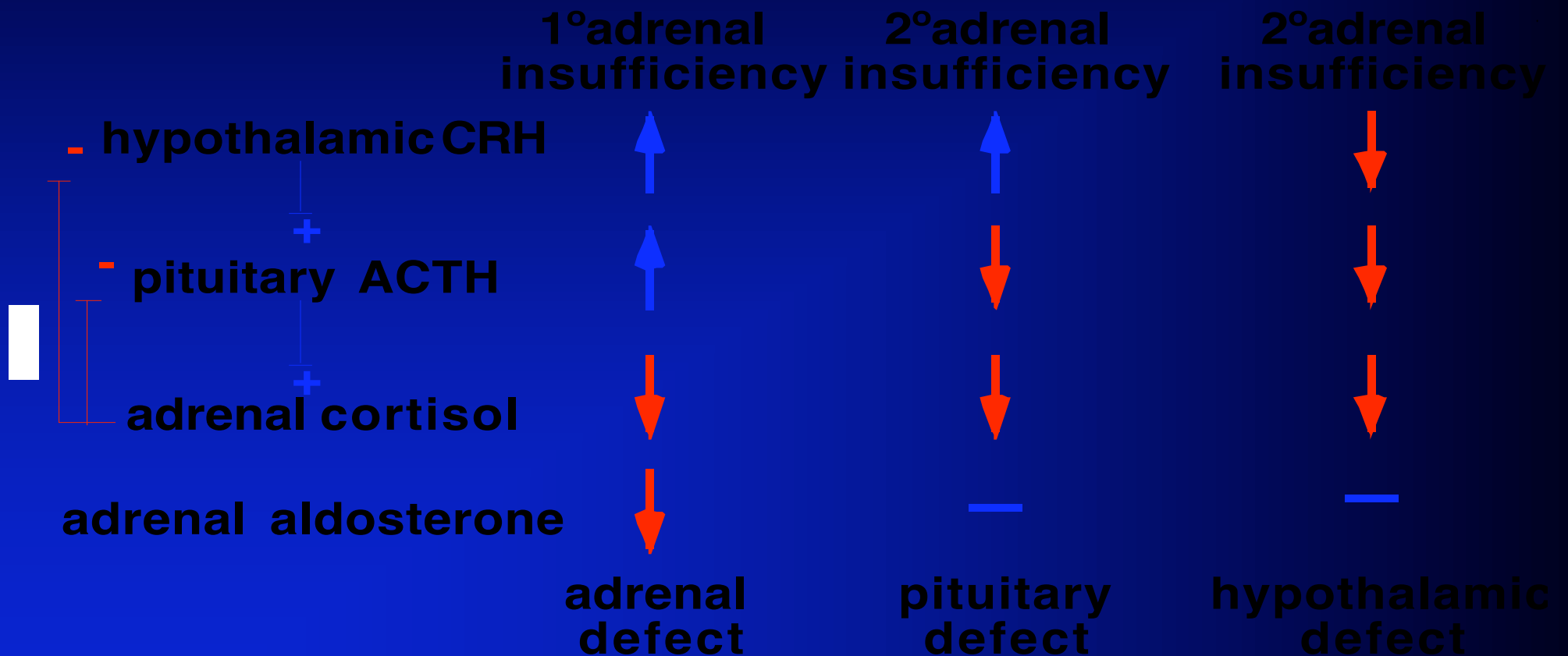


[CC:BY-NC-SA](#)
[BY: synnawang](#)

Types of Adrenal Insufficiency



Adrenal Insufficiency



Adrenal Insufficiency: Age Dependent Prevalence

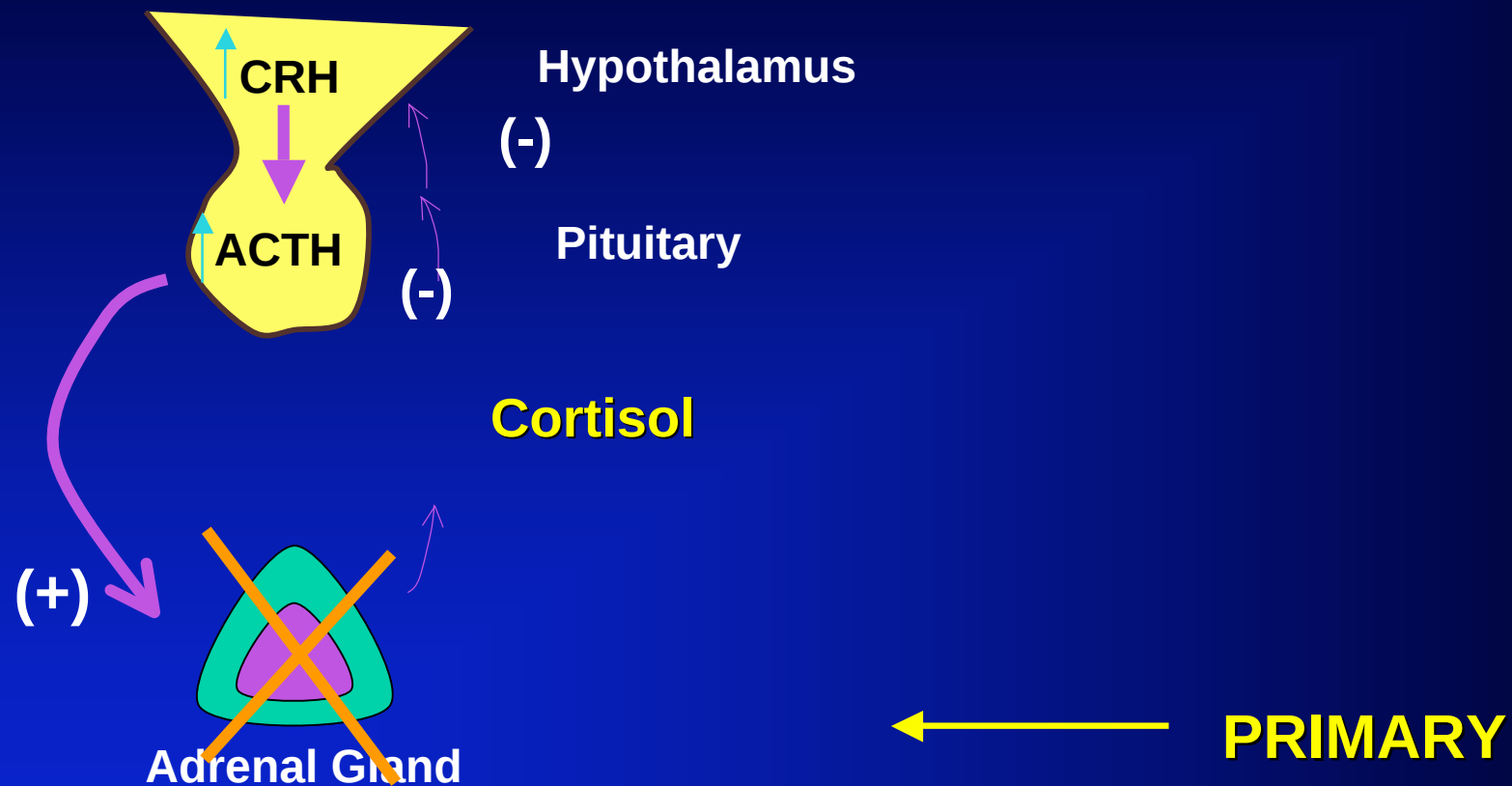
mean age 40 yo (range 17-72 yo)
autoimmune adrenalitis most common in all age groups

children: consider PGA or genetic defect

young men: adrenoleukodystrophy

adults and elderly: glucocorticoids for non -adrenal diseases

Types of Adrenal Insufficiency

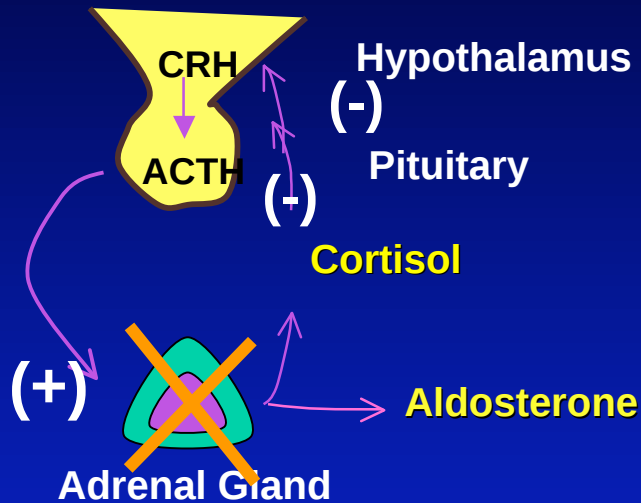


PRIMARY Adrenal Insufficiency



Public
Domain

Thomas Addison (1793-1860)



- Autoimmune adrenalitis (PGA I or II) 80%
- Infections: TB (20% - historically), CMV, fungal
- Vascular: hemorrhage, thrombosis, arteritis
- In cancer patients: metastatic cancer to adrenals
- In young men: adrenoleukodystrophy

IMPORTANT: In PRIMARY adrenal insufficiency, the adrenals are destroyed, and ALDOSTERONE is affected as well.

Adrenal Insufficiency

Autoimmune Adrenalitis

Image of
autoimmune
adrenalitis
removed

Adrenal Tuberculosis

Image of adrenal
tuberculosis
removed

Adrenal Hemorrhage

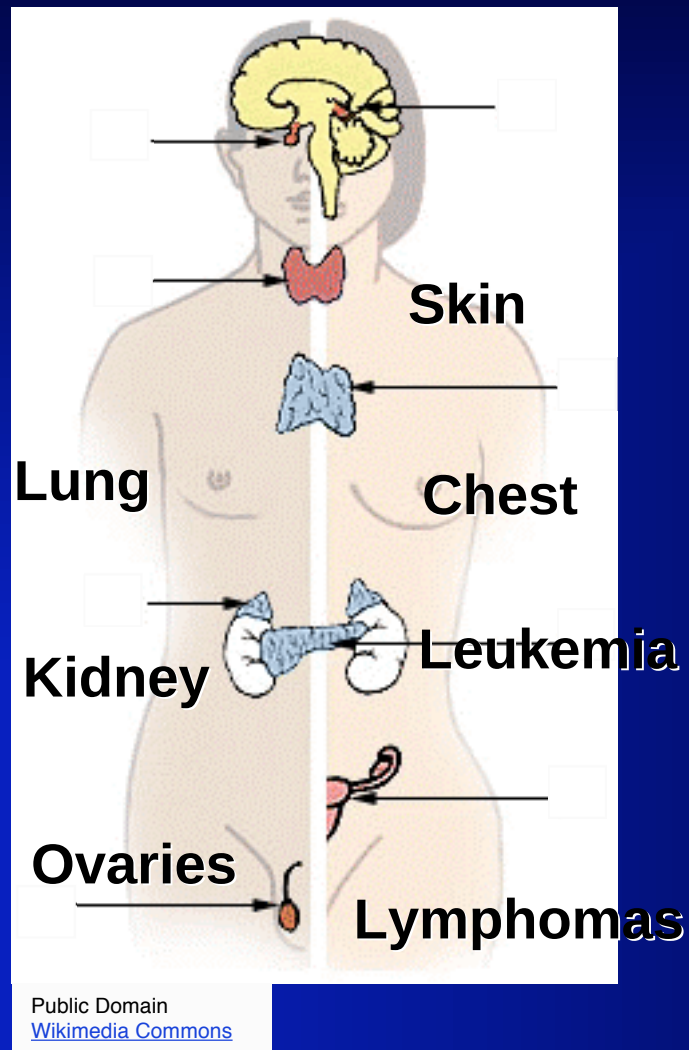


CC:BY-NC-SA
BY: University of Alabama at
Birmingham, Department of Radiology



CC:BY-NC-SA
BY: University of Alabama at
Birmingham, Department of Radiology

Metastases in the Adrenal Gland



Adrenoleukodystrophy/Adrenomyeloneuropathy

X-LINKED - ONLY IN MALES

PRESENTATION

- adrenal insufficiency (childhood)
- hypergonadotropic hypogonadism (puberty)
- spastic paraparesis/demyelination-**AMN**(20-30 yo) vs cerebral sclerosis-**ALD** (childhood)

PATHOPHYSIOLOGY: mutation in **Adrenoleukodystrophy protein(ALPD)**

ALPD function -peroxisomal transport protein anchors very long chain AcylCoA synthetase

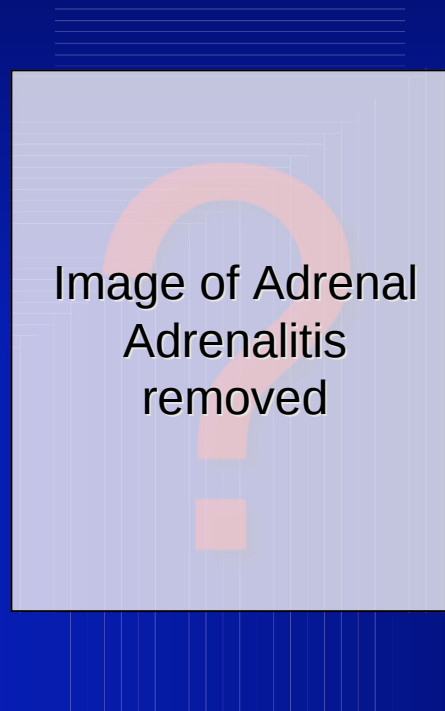
DISEASE - build up of chol. esters w unbranched saturated long chain FAs

TREATMENT: Cortisol replacement
Lorenzo's Oil helps serum level of VLCFA - but no clinical benefit in 3 yr F/U

MUST BE INCLUDED IN w/u of AI in young men and in w/u AI or hypoglycemia in infants

Primary Adrenal Insufficiency

Autoimmune adrenalitis results in **ADRENAL INSUFFICIENCY**



Autoimmune adrenalitis (and therefore its subsequent **ADRENAL INSUFFICIENCY**) can be found in specific genetic syndromes, **POLYGLANDULAR AUTOIMMUNE SYNDROMES**

Primary Adrenal Insufficiency

PGA I (Polyglandular Autoimmune Syndrome I)

autosomal recessive disease- Iranian Jewish heritage starting in childhood

APECED (Autoimmune Polyendocrinopathy-Candidiasis-Ectodermal Dystrophy)

autosomal recessive-Finnish heritage starting in childhood

2 of the following

- adrenal insufficiency (<15 yo)
- hypoparathyroidism (<10yo)
- chronic mucocutaneous candidiasis (<5 yo)

PLUS OFTEN

- dental enamel hypoplasia
- keratopathy/ectodermal dystrophy

occasionally

- chronic active HepB
- malabsorption
- cholelithiasis
- juvenile onset pernicious anemia
- alopecia/vitiligo
- primary hypogonadism
- hypothyroidism
- diabetes mellitus

Which Twin is Sick????



Famous Names in Endocrinology

Addison's Disease



Public Domain
[Wikimedia Commons](#)

John F. Kennedy



Public Domain

Jane Austin (1775-1817)

Addison's Disease & History



Public Domain
[Wikimedia Commons](#)

1960 Presidential Debate
John F. Kennedy vs. Richard M. Nixon
Chicago, Ill., September 21, 1960

Adrenal Insufficiency

Autoimmune adrenalitis

- **PGA II**
 - usually in middle age females
 - adrenal insufficiency
 - hyothyroidism or diabetes mellitus
 - *uncertain genetic component
 - autosomal dominant more likely
 - HAL-B8 chromosome 6
- **PGA III**
 - hypothyroidism
 - other autoimmune disorder (NOT adrenal insufficiency)

Primary Adrenal Insufficiency

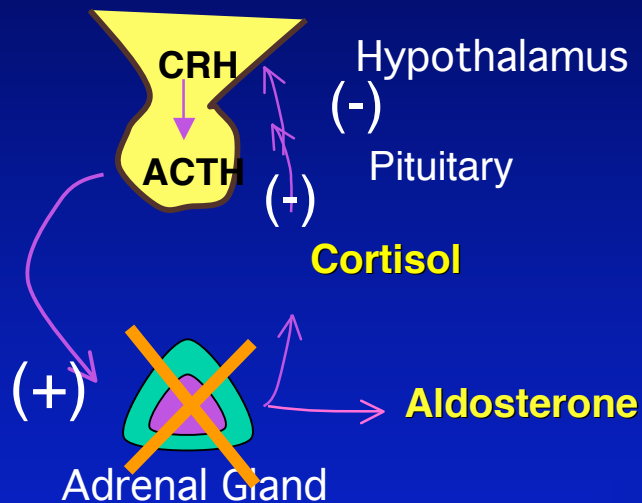
SYMPTOMS

Cortisol

- Fatigue
- Weakness & Malaise
- Anorexia
- Nausea and vomiting

Aldosterone

- Dizziness



SIGNS

- Proximal muscle weakness
- Orthostatic hypotension
- HYPERPIGMENTATION--Primary AI only
- HypoNa, HyperK—Primary AI only

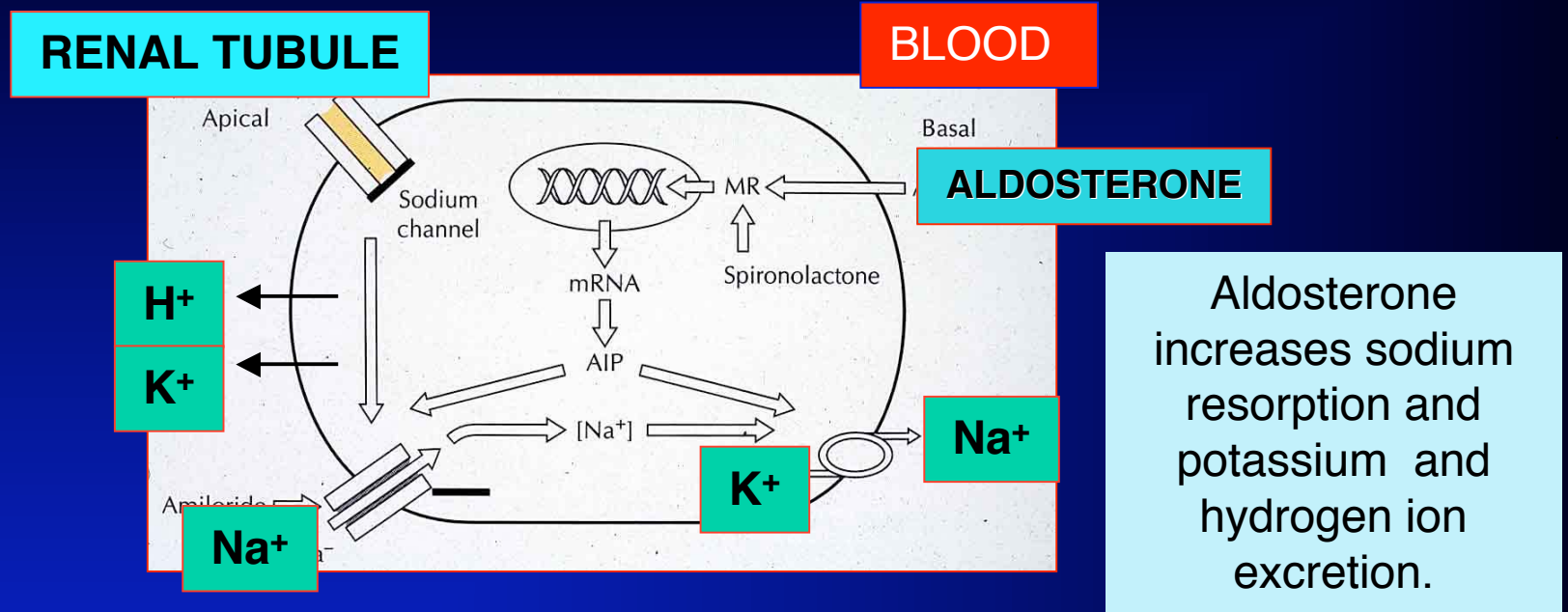
Hypoaldosteronism



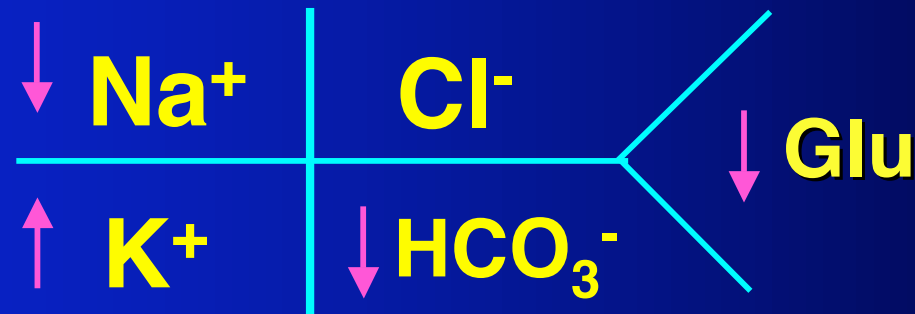
Hypotension
Hyperkalemia
Hyponatremia

Primary Adrenal Insufficiency

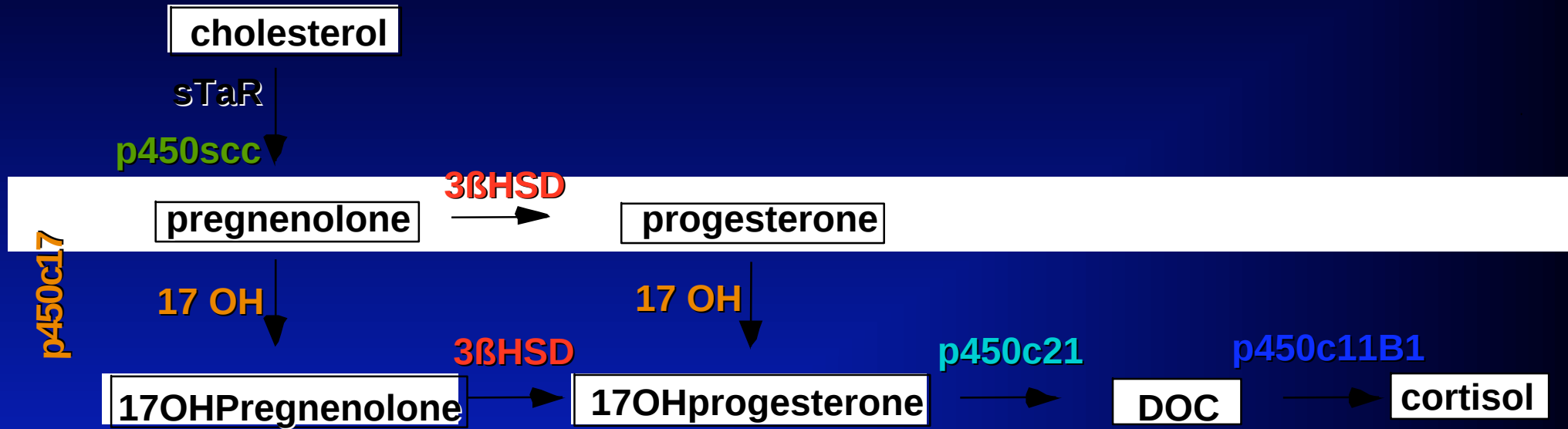
(ALDOSTERONE DEFECT ONLY SEEN IN PRIMARY AI not SECONDARY AI)



So, with aldosterone deficiency:



Glucocorticoid Deficiency

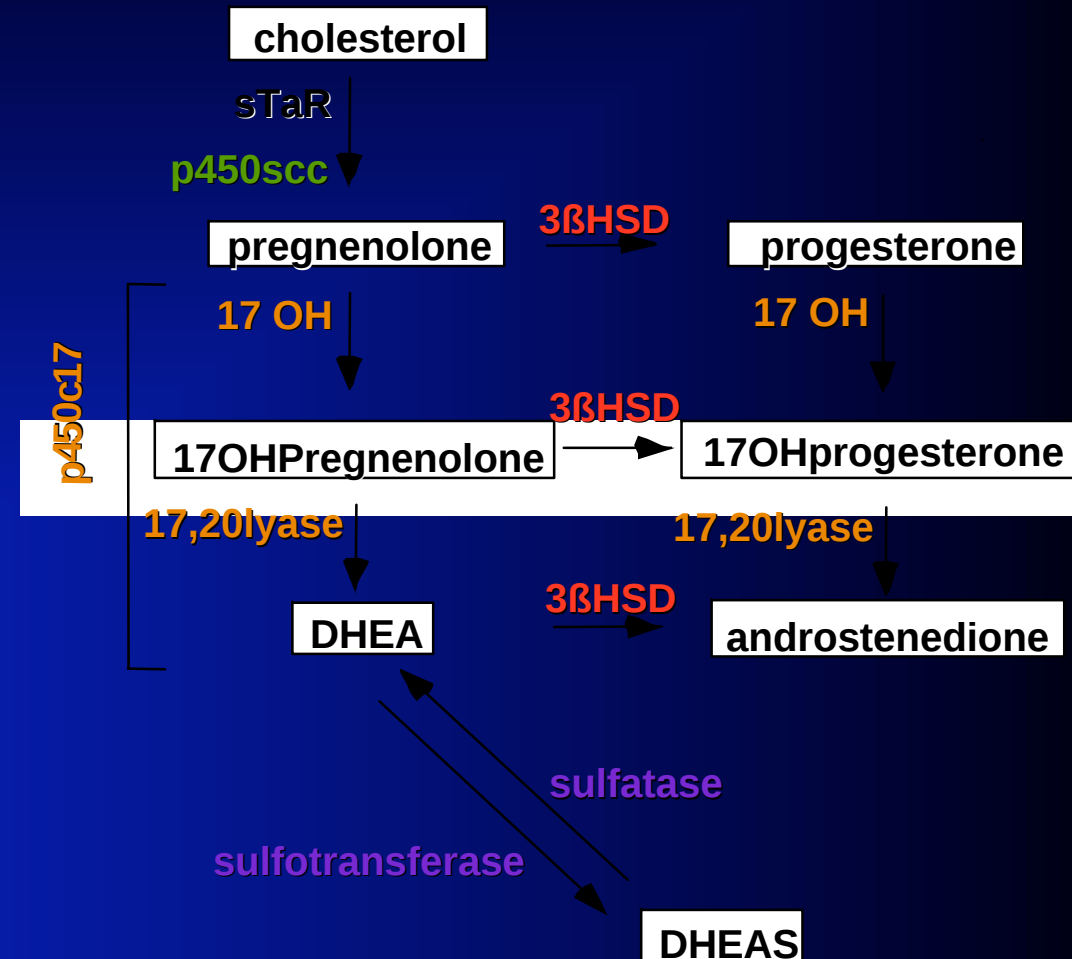


fatigue

hypotension

hypoglycemia

DHEAS Deficiency



Male: fatigue, Δ mood

Female: fatigue, Δ mood, libidinal dysfunction

Adrenal Insufficiency: Hyperpigmentation



Public Domain

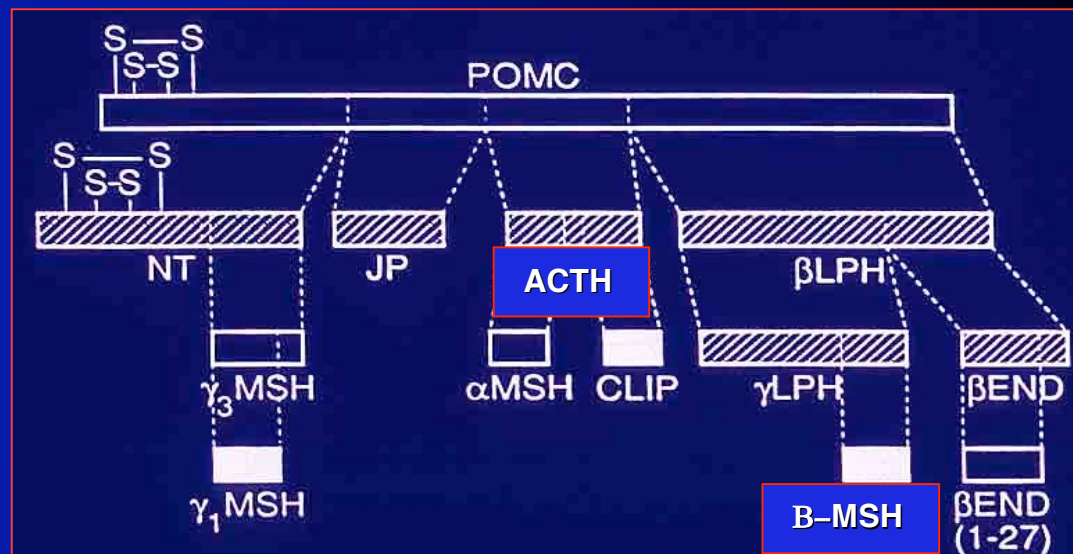
T. Addison
"On the constitutional and local effects of disease of the suprarenal capsules" 1855

Image of patient removed

Image of patient hands removed

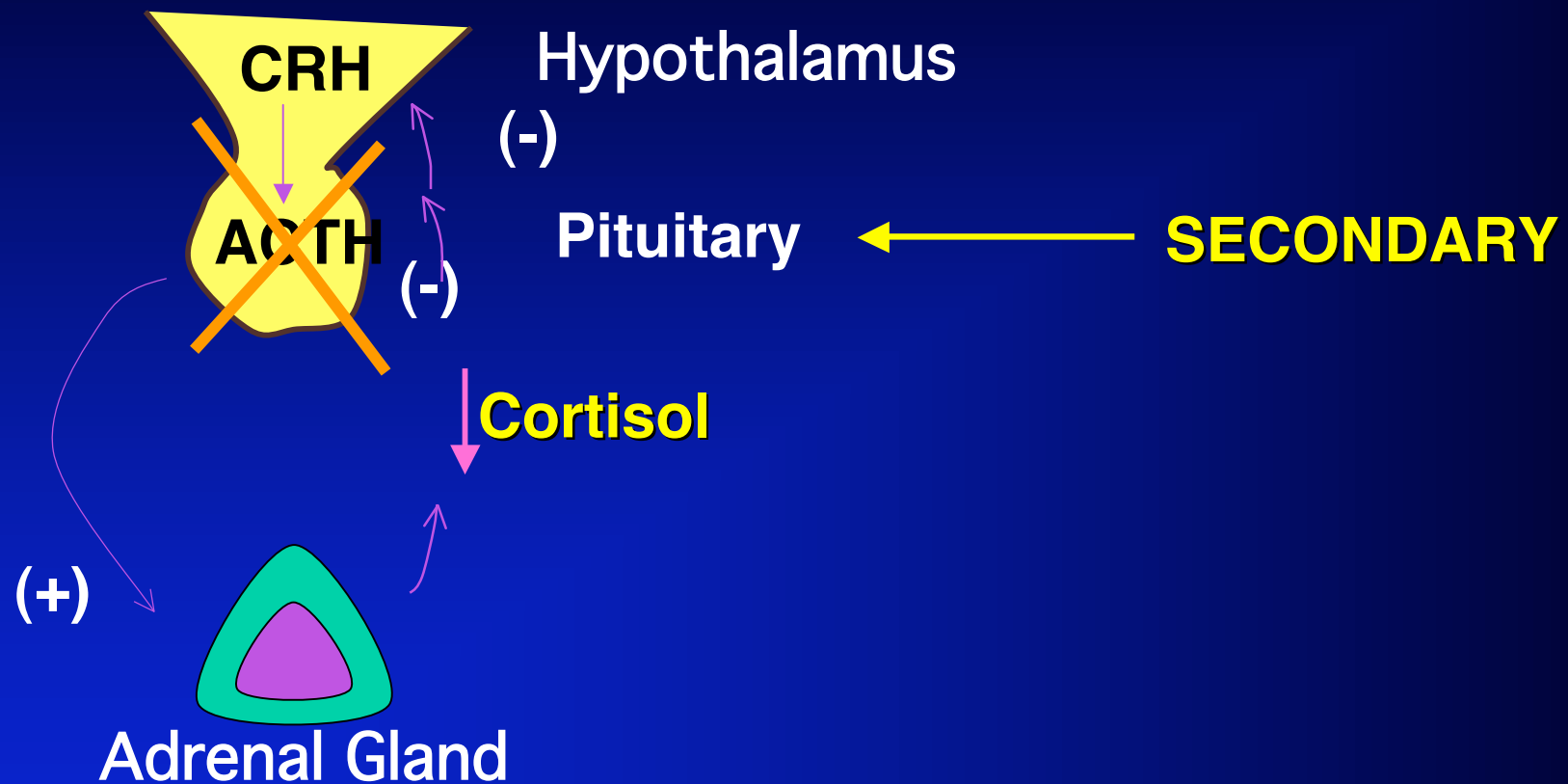
N Engl J Med
1997;337:1666.

Hyperpigmentation of palmar creases

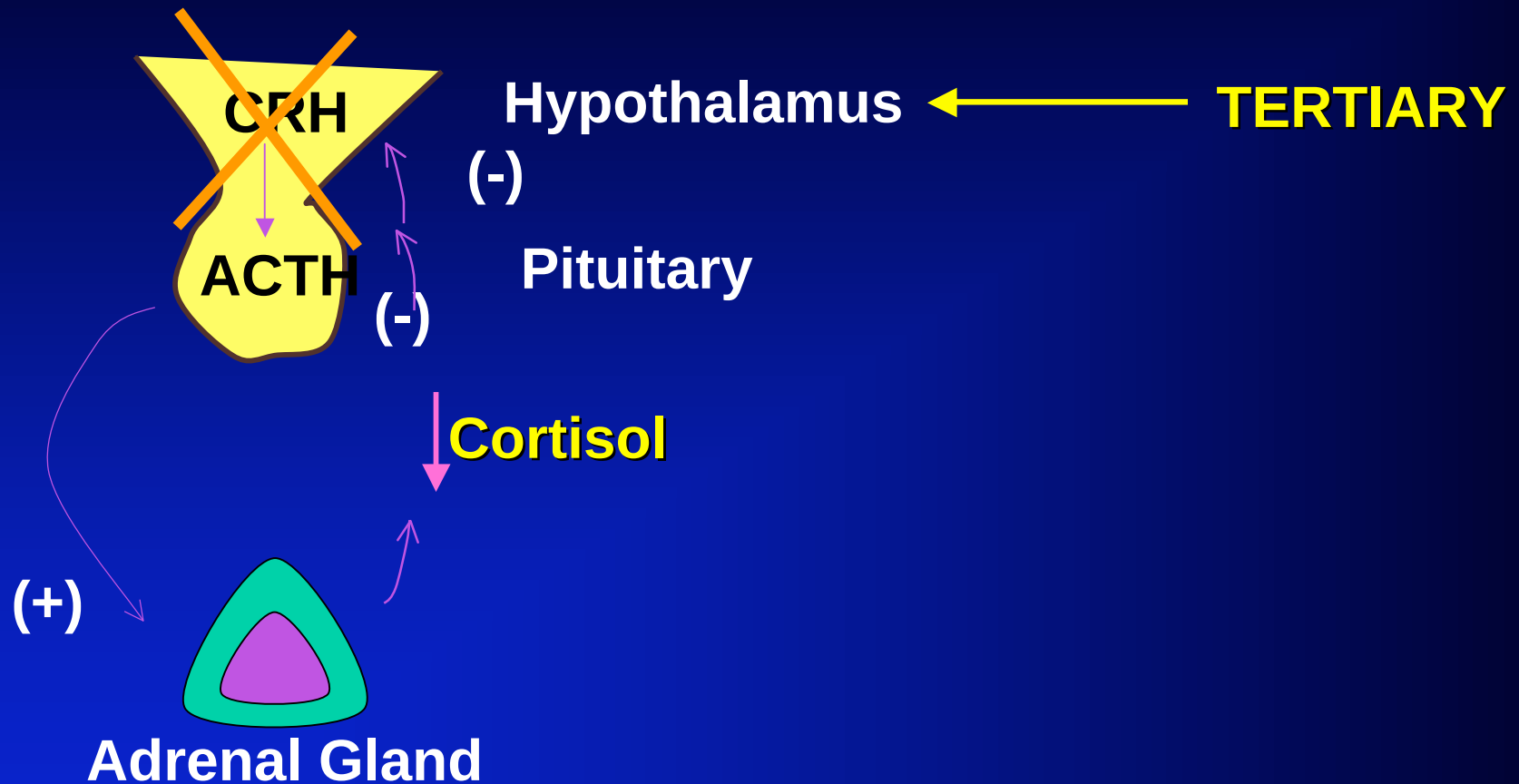


Source: Undetermined

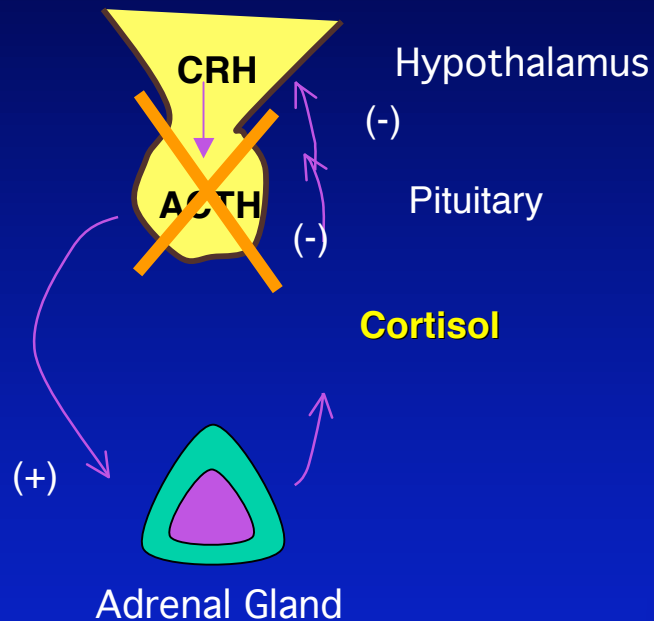
Types of Adrenal Insufficiency



Types of Adrenal Insufficiency



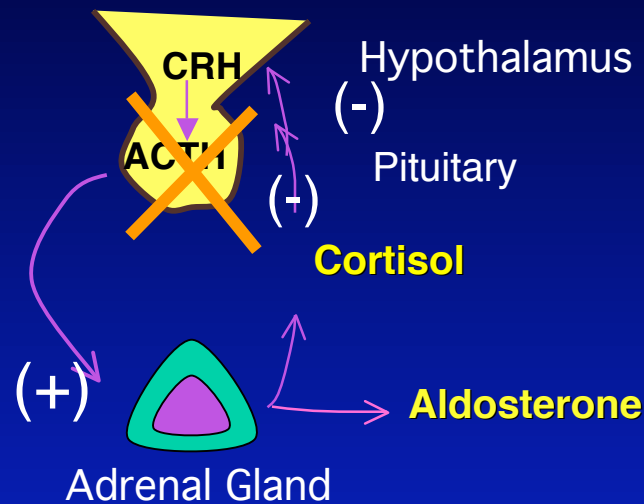
Secondary & Tertiary Adrenal Insufficiency



- **Vascular: Postpartum necrosis(Sheehan's)**
- **Lymphocytic hypophysitis**
- **Infiltrative diseases: Sarcoidosis, Histiocytosis X**
- **Tumor compression**
- **Following surgery or radiation**

- **Long term glucocorticoid treatment**
Pharmacologic Dose = more than physiologic replacement

Secondary Adrenal Insufficiency



SYMPTOMS

- Mild malaise, fatigue
- Proximal muscle weakness

SIGNS

- NO hyperpigmentation
- NO orthostatic hypotension

SIGNS & SYMPTOMS are generally milder than with primary adrenal insufficiency due to cortisol deficiency **ALONE** (ie: **NO ALDOSTERONE DEFICIENCY**)

Adrenal Insufficiency

REMEMBER TO DIFFERENCE BETWEEN PRIMARY AI AND SECONDARY AI

PRIMARY ONLY

- hyperpigmentation (92-96%)
- HYPERkalemia (52-64%)

associated features (ie can see if PGA)

- vitiligo (4%)
- hypothyroidism (primary)
- hypogonadism (primary)

SECONDARY ONLY

associated features (ie can see if entire pit. involved)

- growth delay
- HA
- DI (if stalk involved)
- hypothyroidism (secondary)
- hypogonadism (secondary)

Adrenal Crisis

*hemorrhage

- thromboembolic disease
- Coagulopathy
- anticoagulant therapy
- Waterhouse-Friderichsen Syndrome
 - Neisseria meningitidis septicemia
 - Streptococcus pneumoniae,
 - Pseudomonas aeruginosa
 - Staphylococcus aureus
 - Escherichia coli
 - Haemophilus influenzae
- *drugs - increase metabolism GC
 - phenytoin, phenobarbital, rifampin
- *drugs - decrease production GC
 - ketoconazole, AG, mitotane, metyrapone
- *withdrawal of exogenous glucocorticoids

Adrenal Crisis

- suspect in setting of:
 - catecholamine resistant hypotension
 - hypotension with abd pain
 - must r/o adrenal hemorrhage
- look for:
 - hyperpigmentation/decreased pubic hair
 - hyperkalemia
 - hyponatremia
 - hypoglycemia

If the diagnosis is missed, your patient will most likely die

Adrenal Insufficiency Diagnostic

SCREENING TEST:

AM CORTISOL: GOAL is to RULE OUT disease

Principle of test: Cortisol is highest in the AM allowing maximal chance of ruling out disease

- HI AM cortisol RULES OUT DISEASE

- BUT ONLY EXTREMELY LOW AM cortisol is DIAGNOSTIC

Most patients are neither EXTREMELY HI or EXTREMELY LOW and require DYNAMIC testing

Adrenal Insufficiency Diagnostic

DIAGNOSTIC TEST FOR PRIMARY ADRENAL INSUFFICIENCY:

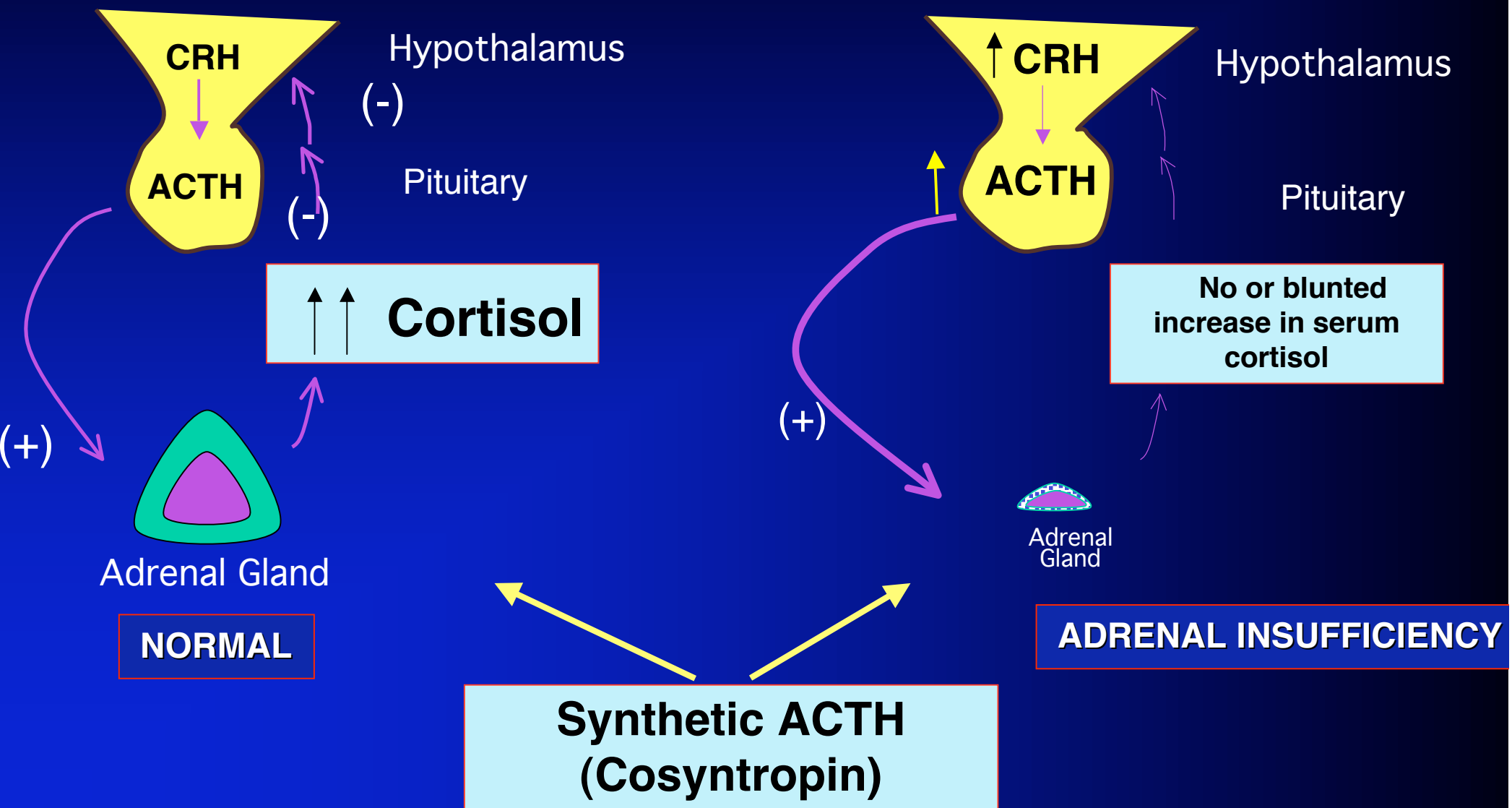
ACTH STIMULATION TEST: GOAL is to RULE IN disease

Principle of test: ACTH stimulates steroidogenesis and secretion of cortisol - normal levels well documented

- Cortisol level after ACTH that is SUBNORMAL is DIAGNOSTIC of AI

- ACTH level that is EXTREMELY HI is CONSISTENT with diagnosis of PRIMARY AI
but is NOT DIAGNOSTIC

The ACTH Stimulation Test



Adrenal Insufficiency Diagnostic

DIAGNOSTIC TEST FOR SECONDARY ADRENAL INSUFFICIENCY:

INSULIN HYPOGLYCEMIA TEST: GOAL is to RULE IN disease

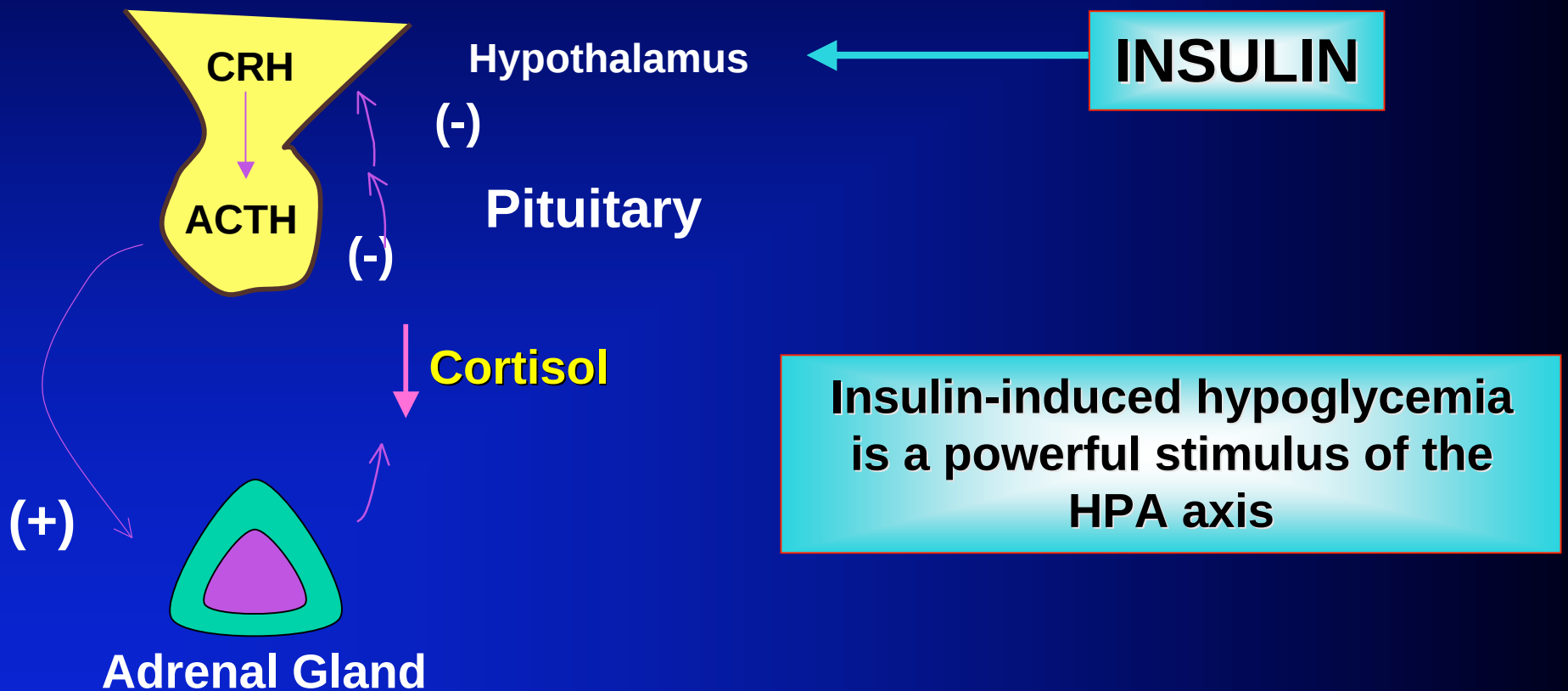
Principle of test: Insulin results in hypoglycemia that is the strongest stimulus for activation of HPA axis at the level of CRH

Cortisol level after IHT that is SUBNORMAL is DIAGNOSTIC of AI

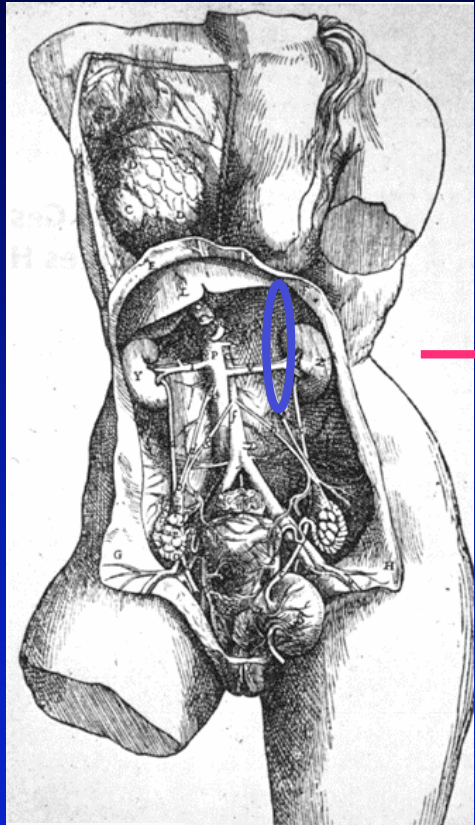
ACTH level after IHT that is SUBNORMAL is DIAGNOSTIC of SECONDARY AI

Diagnosis of Secondary/Tertiary Adrenal Insufficiency

The Insulin Tolerance Test



Therapy for Adrenal Insufficiency



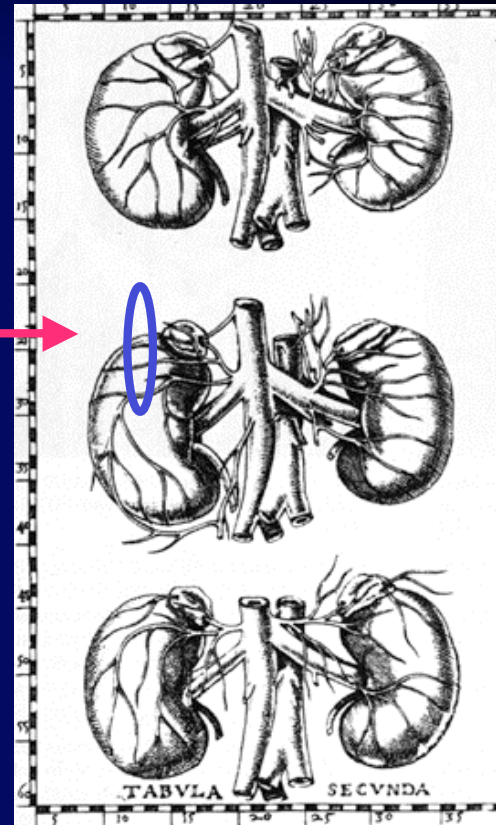
Public Domain

1543

Image of
adrenal
gland
removed

Therapy

Image of
adrenal
gland
removed



Public Domain

1564

Guidelines for Management

GUIDING PRINCIPLE: The more severe the stress the more cortisol patient needs

Acute Therapy (significant ill or Adrenal Crisis)

IV fluids

IV cortisol: HI DOSE

glucose

treat underlying precipitating events

Do not wait for labs!!!!

Maintenance Therapy

Glucocorticoids

hydrocortisone ~ 15-25 mg/d

titrate to a sense of well being and physical strength

avoid weight gain, hypertension, hyperglycemia and osteoporosis

Mineralocorticoids

fludrocortisone ~0.1 mg/d

titrate to salt craving and postural hypotension

together with serum K and upper range renin

DHEA -

Guidelines for Management

GUIDING PRINCIPLE: The more severe the stress the more cortisol patient needs!

Stress Dosing Glucocorticoids

Minimal

dental work
mild or non-febrile illness

no need for supplemental coverage

Minor

hernia repair

25 mg hydrocortisone - **day of procedure** (or onset of fever)

Moderate

hemicolectomy
significant febrile illness

50-75 mg hydrocortisone - day of procedure (or onset of fever)
rapid taper in **1-2 days**

Severe

cardiac surgery

100-150 mg hydrocortisone - day of procedure (or onset of fever)
rapid taper in **1-2 days**

Critically ill

sepsis

100 mg hydrocortisone i.v. bolus followed by -
50-100 mg hydrocortisone i.v. **q 6-8 hours** (or 0.18 mg/kg/hr)
0.05 mg/d fludrocortisone until shock resolves (**days to weeks**)

Discontinuing Glucocorticoids Following Long Term Suppression

GUIDING PRINCIPLE: The more glucocorticoid and the longer treated - the greater chance of long term suppression and atrophy of HPA axis

risk of suppression

Low risk: Low dose, short duration or short “bursts” of glucocorticoid
High dose and prolonged therapy ($\geq 1-4$ weeks) - risk is higher

time course for recovery

Larger doses for prolonged periods (months - years) - recovery can take from 9 MONTHS up to 1-2 years

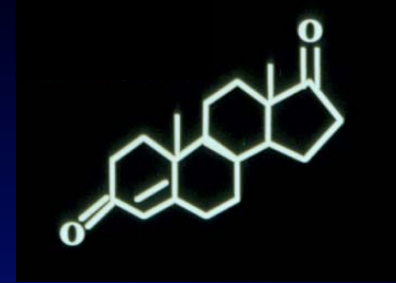
need for taper

taper from pharmacologic to physiologic (**determined by non-adrenal disease course**)
taper from physiologic to no treatment (**determined by adrenal suppression**)

DHEA: What is all the fuss?

- **Marker of aging**
 - ??pharmacologic reversal of aging process??
- **Predictor of morbidity/mortality**
- **Works wonders in rodents**
 - CNS, obesity, diabetes, immunity
- **Preliminary studies in humans**

DHEA what is it??



**19 carbon (androstane)
 $\Delta^5,3\beta$ -hydroxy,17-keto
SO₄, ester at 3 β**

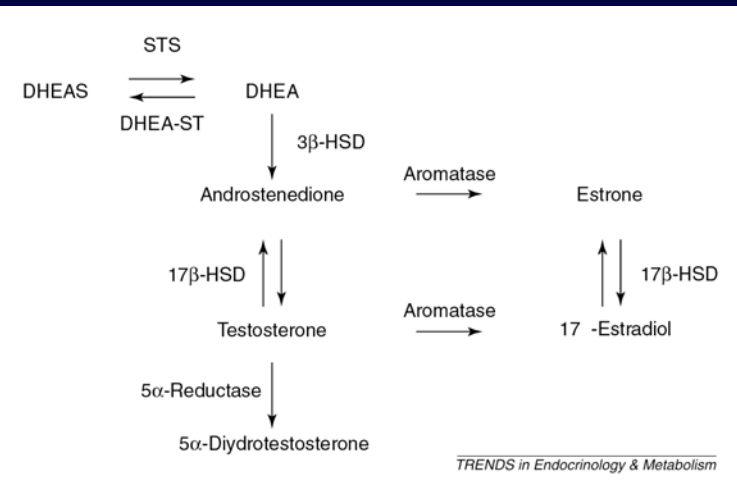
Source: Undetermined

- **Synthesized by adrenals only in humans and higher primates**
 - obligate precursor of all sex steroids in humans
- **More synthesized than all other steroids**
 - up to 25 mg/day in adults
 - major secretion of fetal adrenal
- **Most secreted as sulfate (DHEA-S)**
 - sulfation is ONLY in ADRENAL (NOT GONAD)
- **Inactive at androgen receptor**

DHEA: How Does it Work?

- **Conversion to androgens**
 - 50 mg/d raises testosterone in females
- **Intrinsic activity of DHEA-S in brain**
 - trophic effects on cultured neurons
 - GABA, NMDA, sigma receptor-channels
- **Actions of weird metabolites**
 - concept of NEUROSTEROIDS

Case for DHEAS



Source: TRENDS

DHEA + DHEAS

major secretory products of adrenal
peak in fetal life and adrenarche

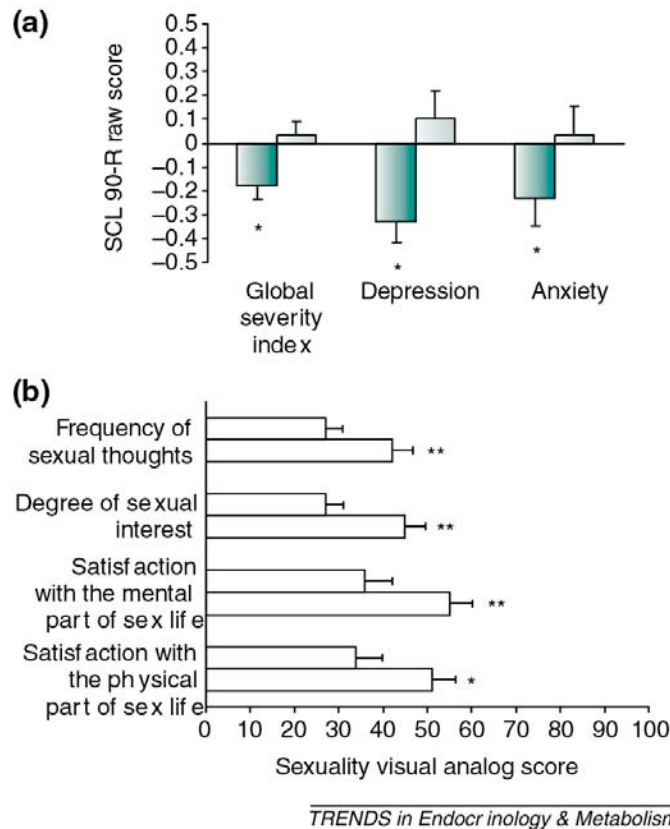
Decline throughout adult life to 20-20% by 70-80 yo

Advertisement as ANTI-AGING drug
In USA : FOOD SUPPLEMENT!!!!

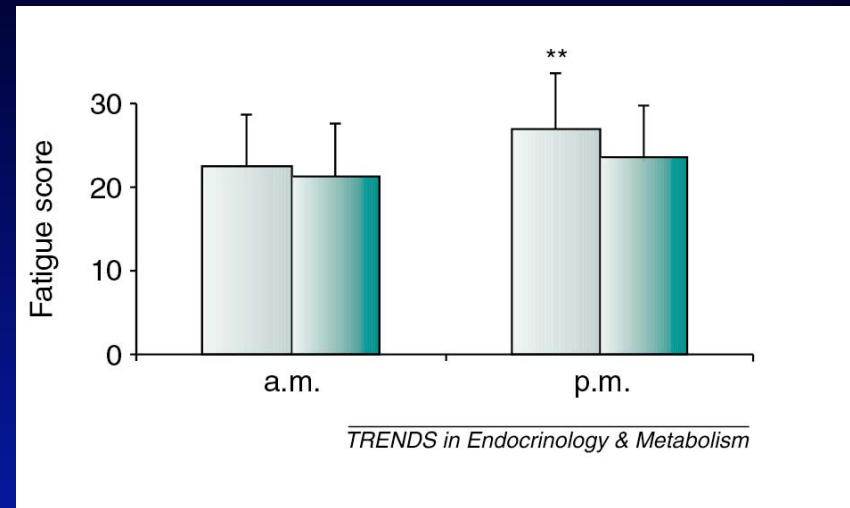
Images of DHEA
removed

classic steroid converted to testosterone peripherally
neurosteroid directly binding NMDA + GABA receptors

Case for DHEAS



Source: TRENDS



Source:
TRENDS

DHEA in **men and women** with primary adrenal insufficiency improves **mood** and **well-being**, irrespective of the patient's sex.

DHEA replacement in **women** with adrenal insufficiency improved overall well-being and mood, specifically **depression**, **anxiety** and both **sexual interest** and **sexual satisfaction**.

Guidelines for DHEA Treatment in Adrenal insufficiency

Adrenal Androgens

only in pts w AI who do NOT feel
“normal on replacement GC and MC”

DHEA: 25 mg po q a.m.

- may increase to 50 mg
- dictated by response and androgenic side effects
- monitor labs

DHEAS, androstendione and free test
LFTS and lipids at 4 + 12 w

Watch Out for Supplements

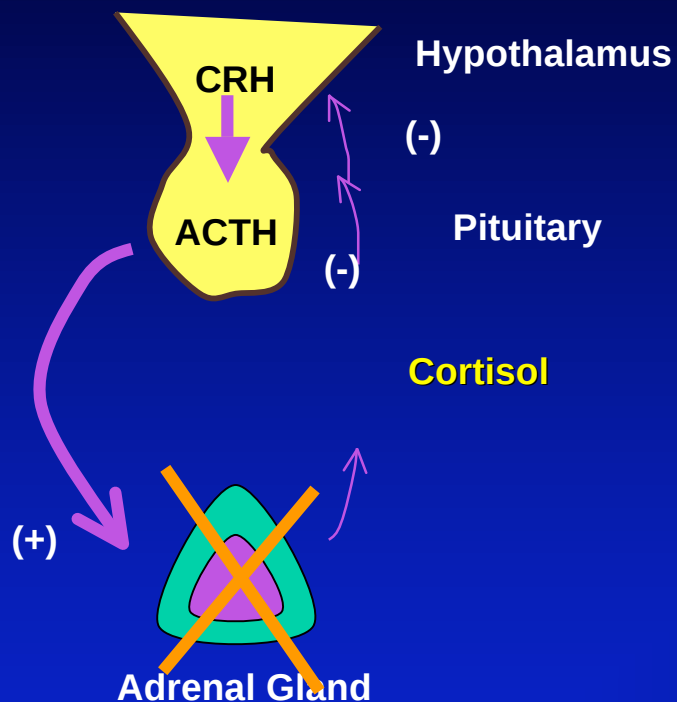
Steroids are lipophilic
Undetermined dosing
Undetermined purity

Image of Adrenal
Glandular Plus
Supplements
removed

Image of Adrenal
Cortex Complex
Supplements
removed

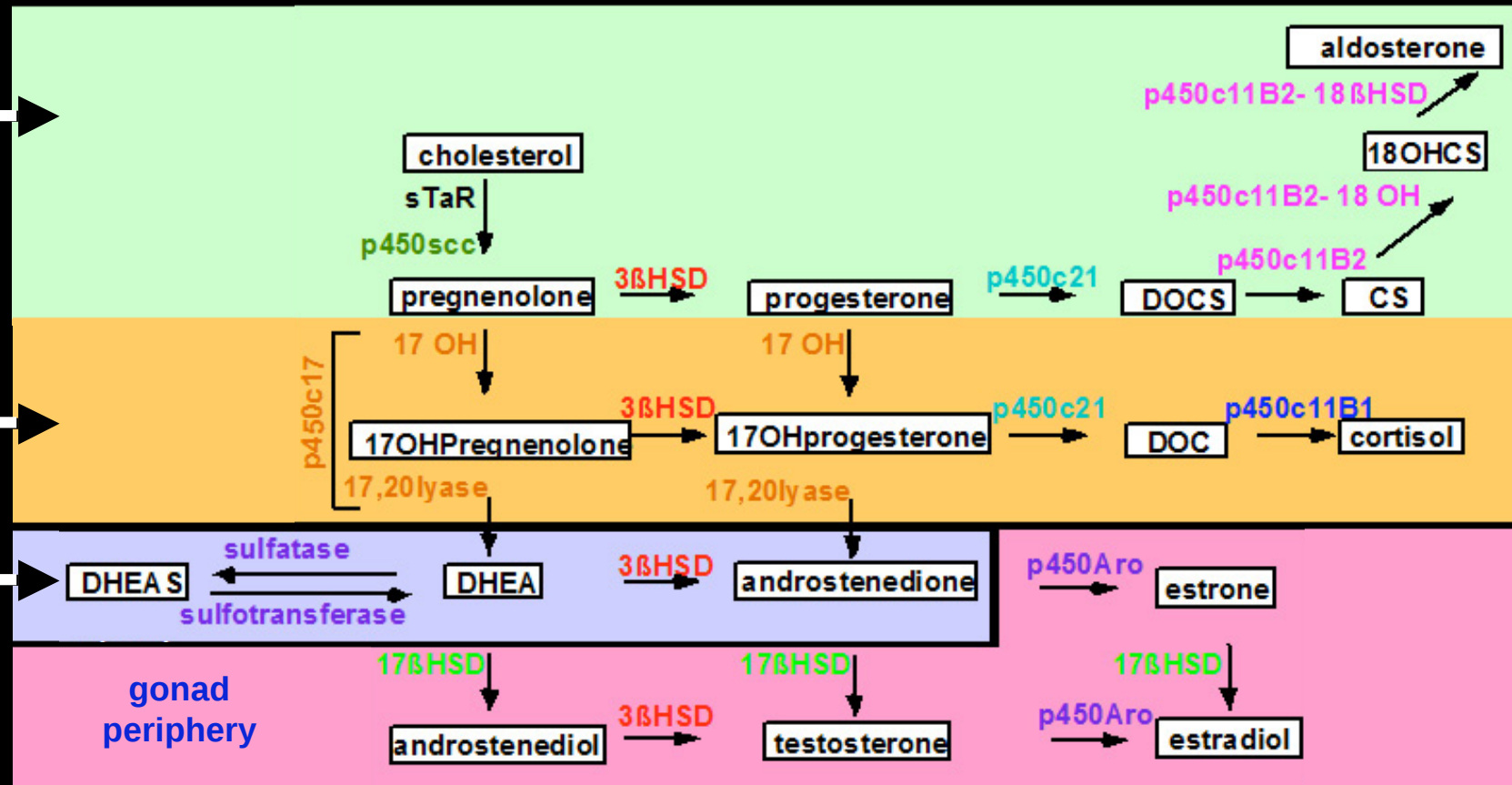
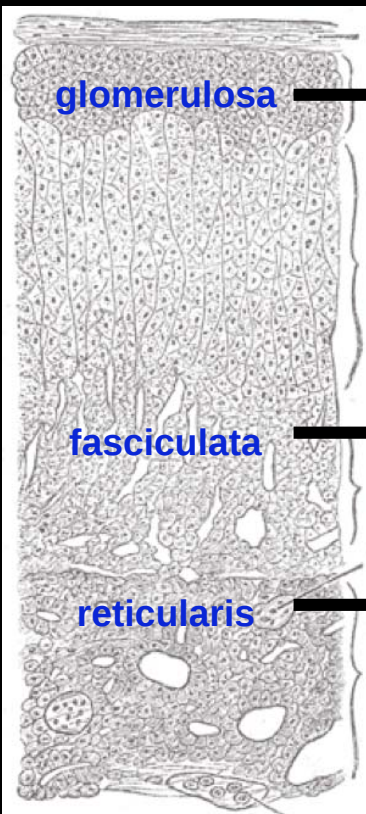
Image of Raw
Adrenal
Supplements
removed

Congenital Adrenal Hyperplasia

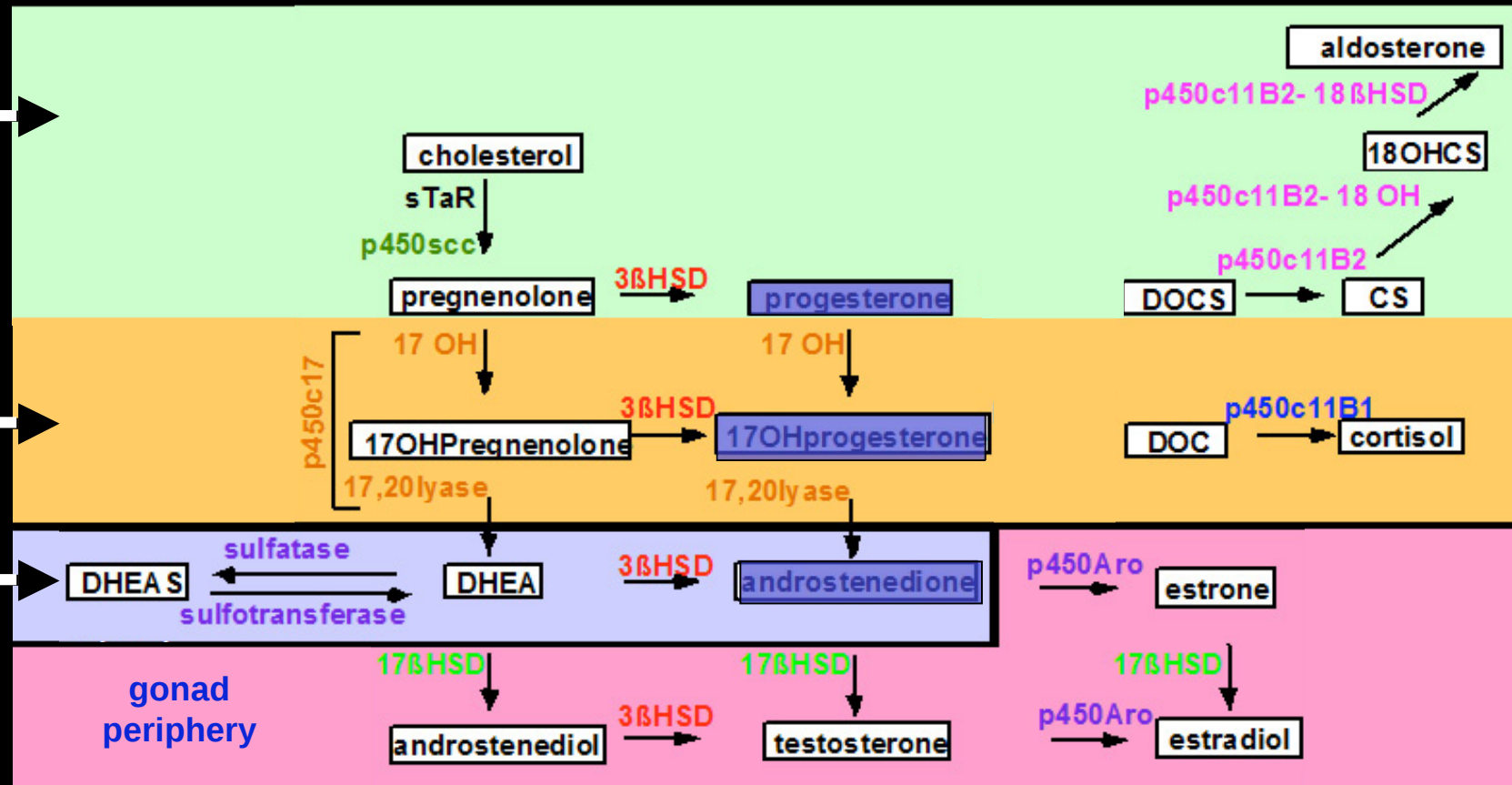
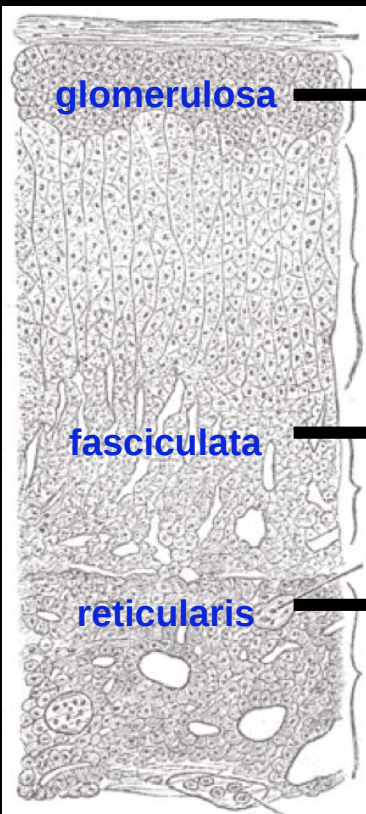


- Genetic block in biosynthetic pathway for cortisol and aldosterone result in primary adrenal insufficiency.
- Decreased feedback on hypothalamus and pituitary increase CRH and ACTH.
- Increased ACTH further stimulates adrenals and results in shunting and production of precursors.
- ACTH stimulates growth (HYPERPLASIA) of adrenals.

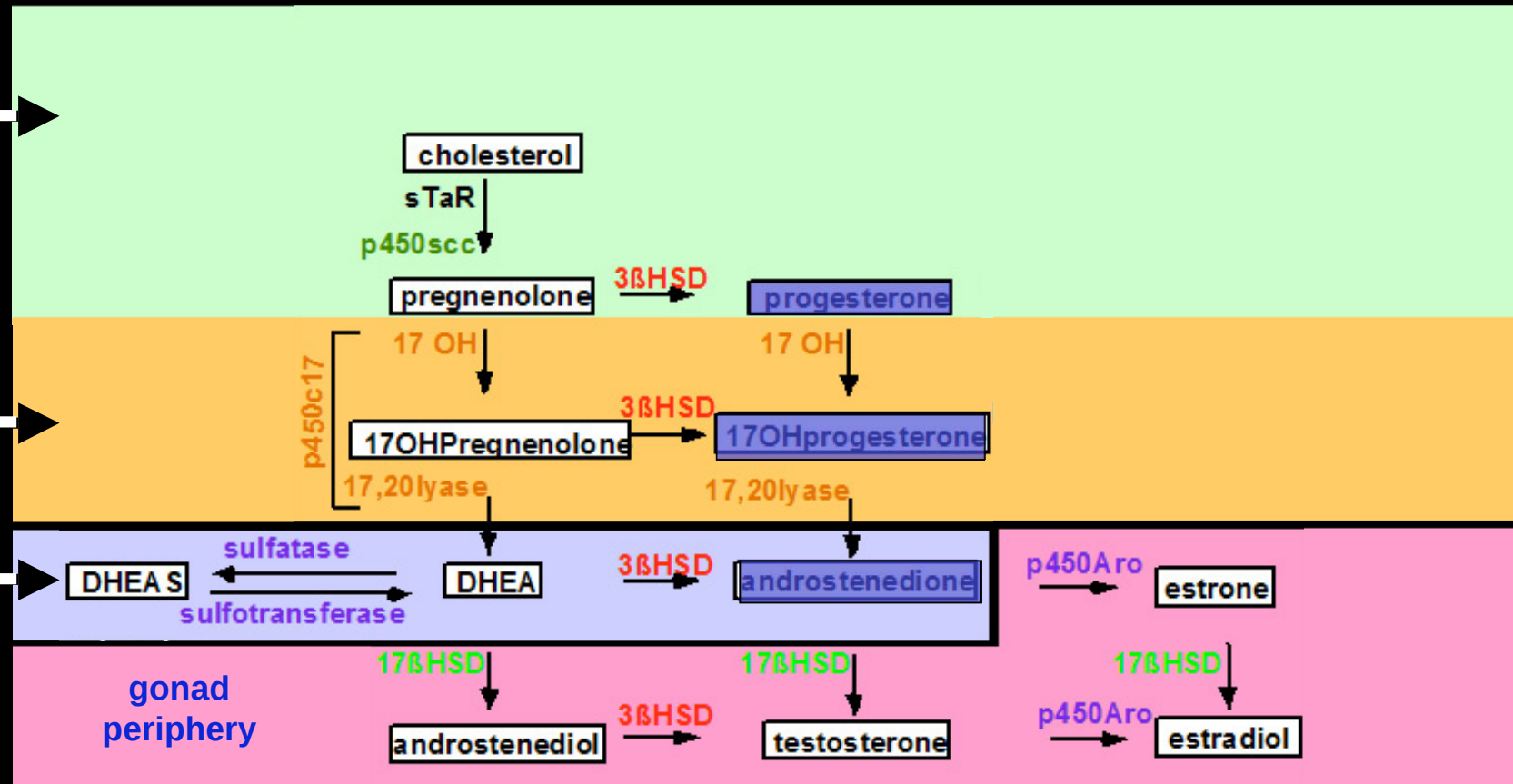
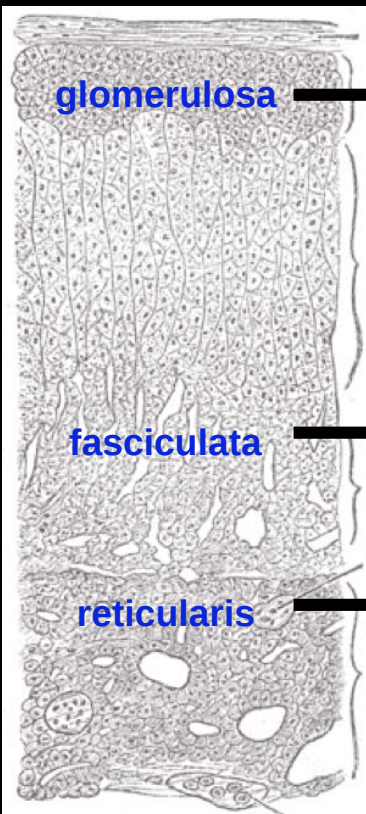
Steroidogenesis



Steroidogenesis



Steroidogenesis



SEVERE P45c21 Deficiency in FEMALE



	FEMALE PSEUDOHERMAPHRODITISM	
AGE -----	4 ⁴ / ₁₂ yrs.	23 yrs.
HT. AGE ----	5 yrs.	11 yrs.
BONE AGE --	9 yrs.	ADULT
17 K.S. ----	6.0mgm / 24 hrs.	50 mgm / 24 hrs
Pregnanetriol	13.6mgm / 24 hrs.	—

results in androgen excess in utero

MILD P45c21 Deficiency in FEMALE



CC:BY-NC-SA 1.0
BY: C. Matthew Peterson, MD

Cliteromegaly

Beard



CC:BY-NC-SA 1.0
BY: C. Matthew Peterson, MD

Hirsutism

results in androgen excess at puberty

Congenital Adrenal Hyperplasia

Important things to remember:

- Loss of function of enzyme in steroidogenesis pathway
- “Block” in pathway leads to shunting down alternate paths and abnormal build-up precursors before the block.
- Severe forms lead to virulization of females
- Milder forms (“non-classical”) may lead to hirsutism and menstrual abnormalities in women.
- Block in pathway may result in adrenal insufficiency during times of stress.

Adrenal Excess States

Causes of hypercortisolism

■ Physiological states

- Pregnancy
- Stress
- Chronic excessive exercise
- Malnutrition

■ Pathologic states

- Cushing's syndrome
- Diabetes mellitus
- Hyperthyroidism
- Severe chronic disease
- Glucocorticoid resistance
- Psychological states
- Anorexia nervosa
- Panic disorder
- Melancholic depression
- Obsessive-compulsive disorder

- PHARMACOLOGIC USE OF
GLUCOCORTICOIDS

Cushing's Syndrome



Public Domain

Harvey Cushing (far left) in 1895 during his House Pupilship (internship) at Massachusetts General Hospital.



Public Domain

DR. HARVEY WILLIAMS CUSHING
(1869-1939)

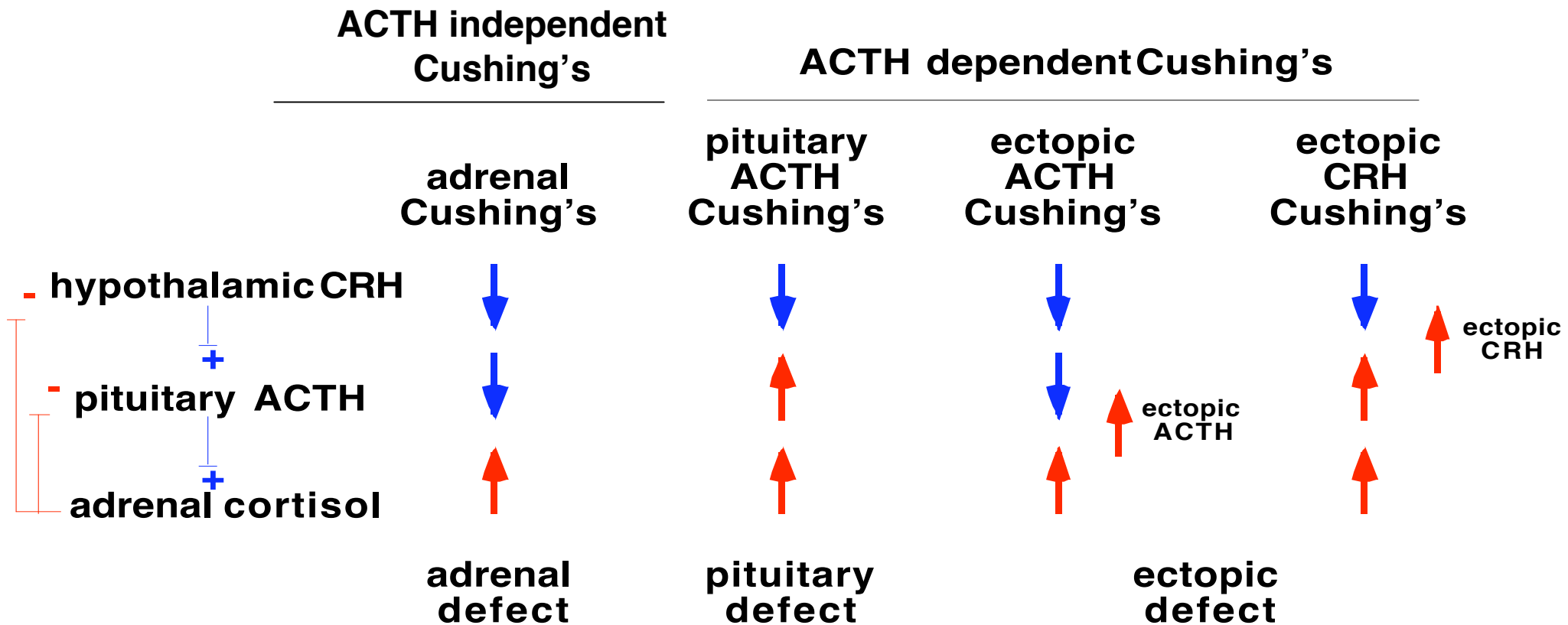
Cushing HW. The basophil adenomas of the pituitary body and their clinical manifestations (pituitary basophilism). Bulletin of the Johns Hopkins Hospital. 1932;50:137-95

His research on the pituitary body gained him an international reputation, and he was the first to ascribe to pituitary malfunction a type of obesity of the face and trunk now known as Cushing's disease, or Cushing's syndrome.

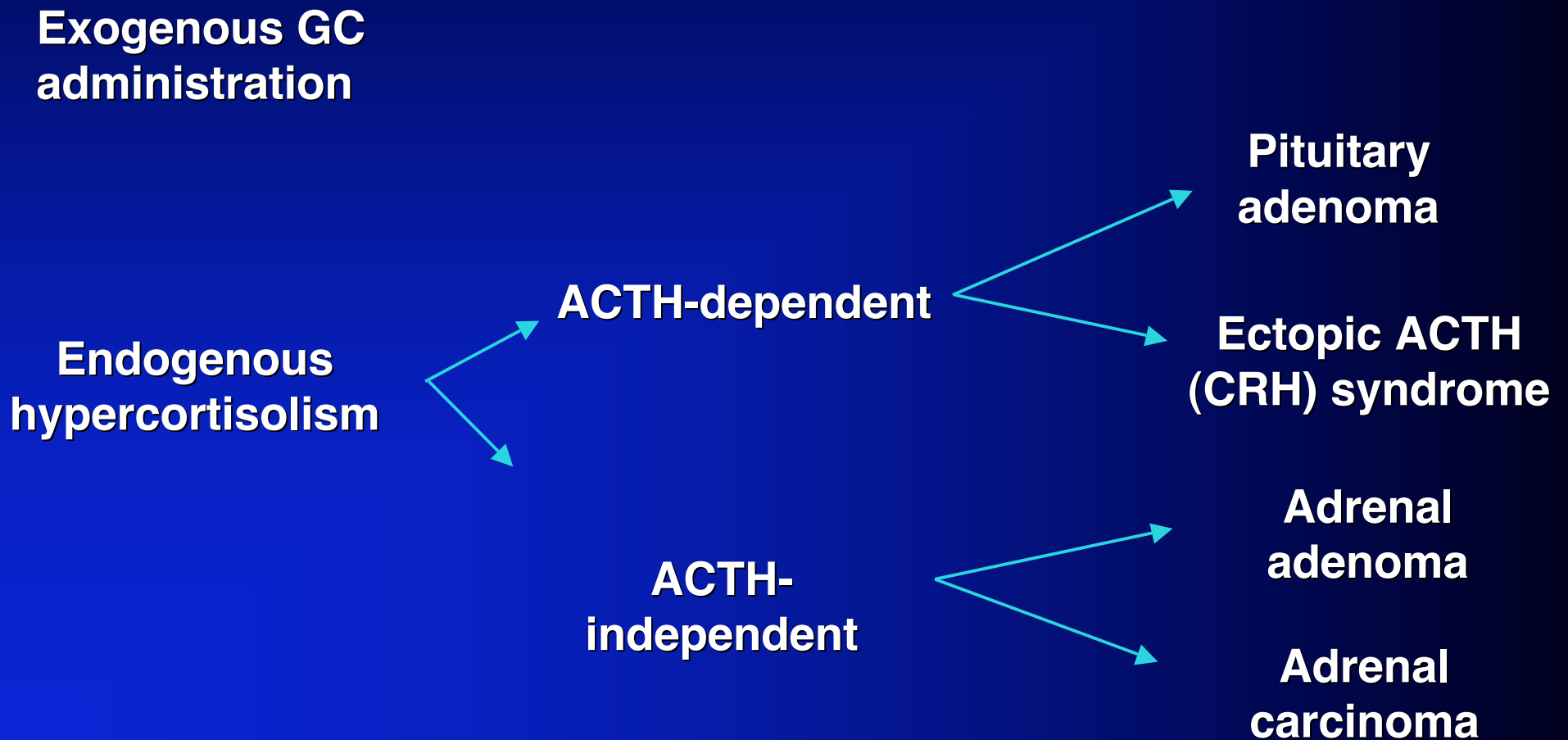
Cushing's Syndrome

- All types of Cushing's Syndrome
 - HI CORTISOL (urine and serum)
 - Absent circadian rhythm
- Adrenal Cushing's syndrome is autonomous and therefore has LOW ACTH
- Only ACTH-dependent Cushing's (by definition) has HI ACTH

Cushing's Syndrome



Cushing's Syndrome



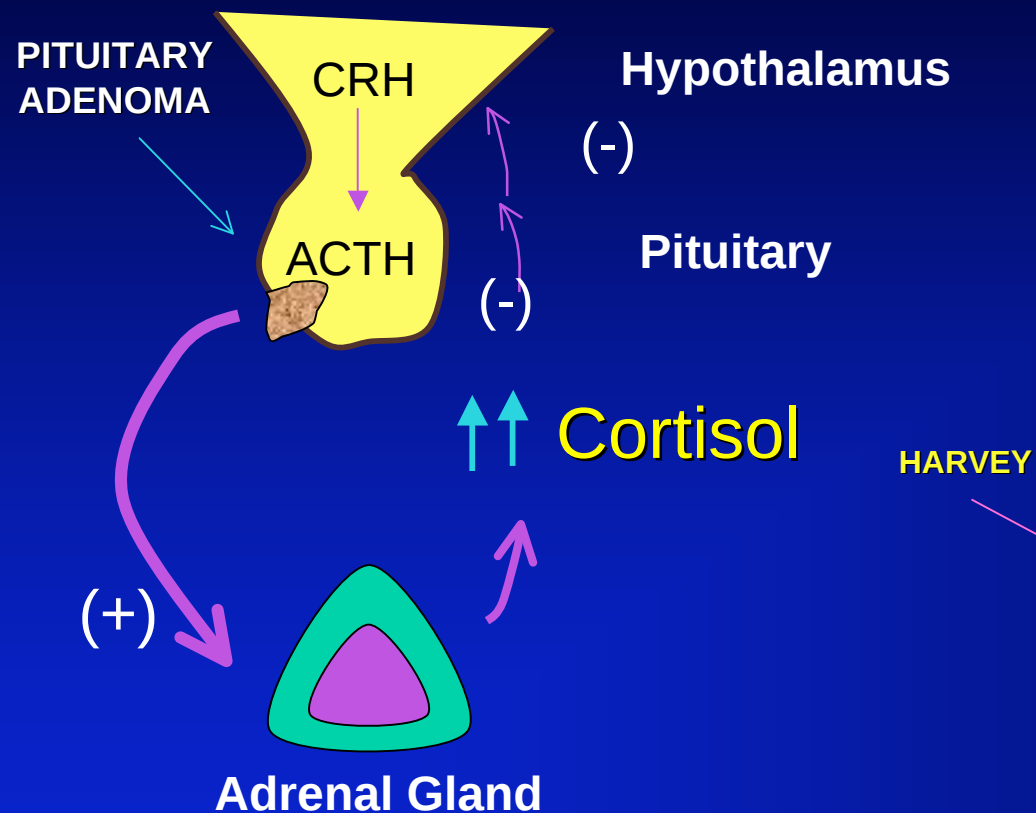
Cushing's Syndrome



- **ACTH-dependent Cushing's Syndrome**
 - **pituitary adenoma-ACTH (60%)**
 - **Ectopic hormone (10%)**
 - **ACTH**
 - **CRH**

all result in bilateral adrenal hyperplasia

Types of Cushing's Syndrome



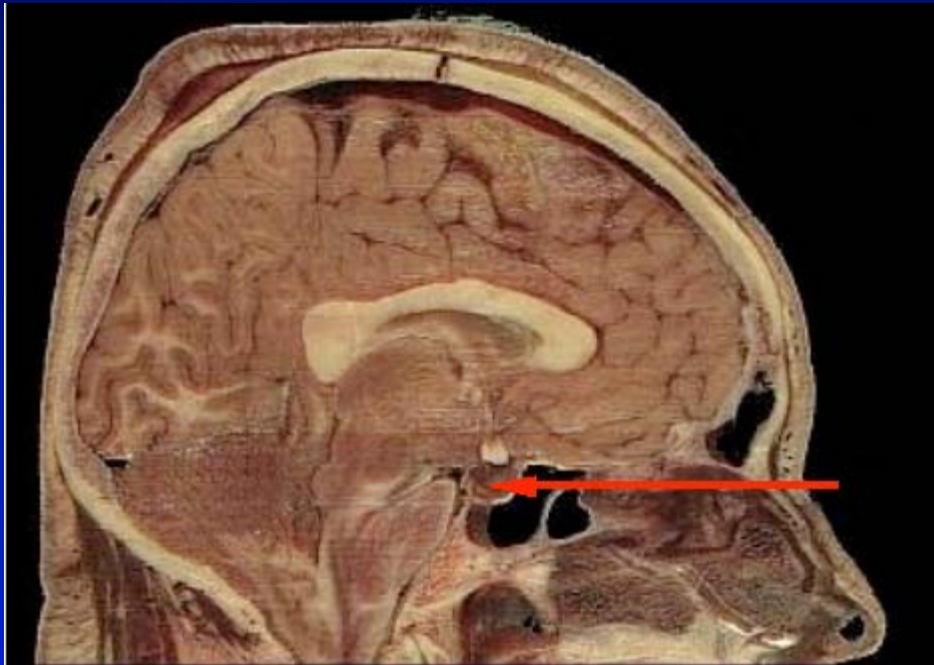
Cushing's Disease
("pituitary Cushings"):
hypercortisolism from a
pituitary adenoma

HARVEY

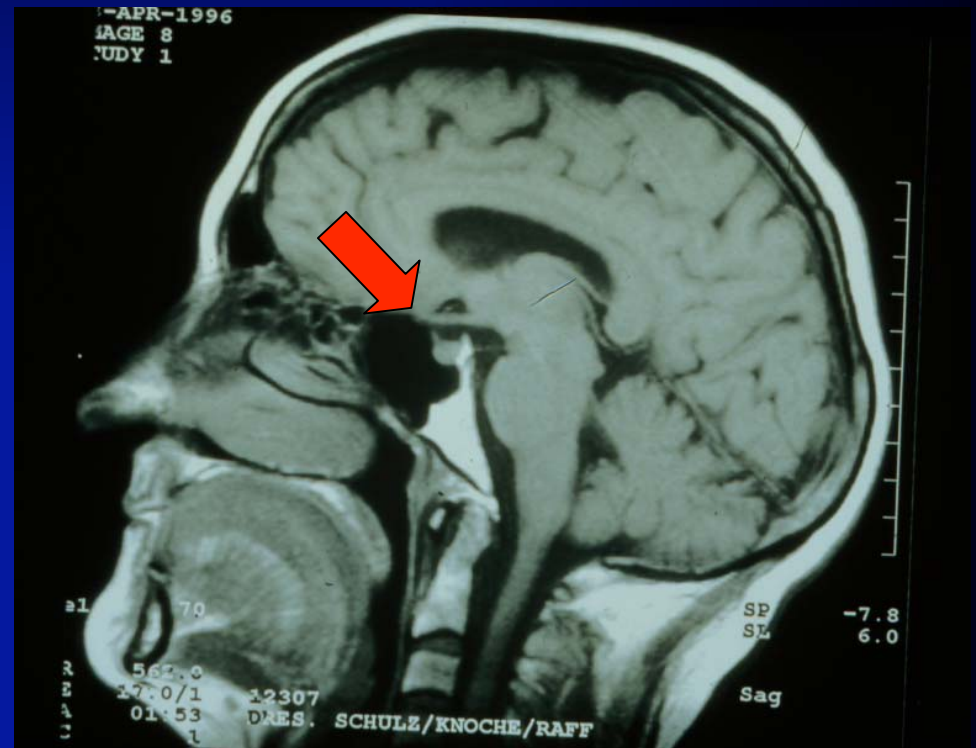


Public Domain

Normal Pituitary

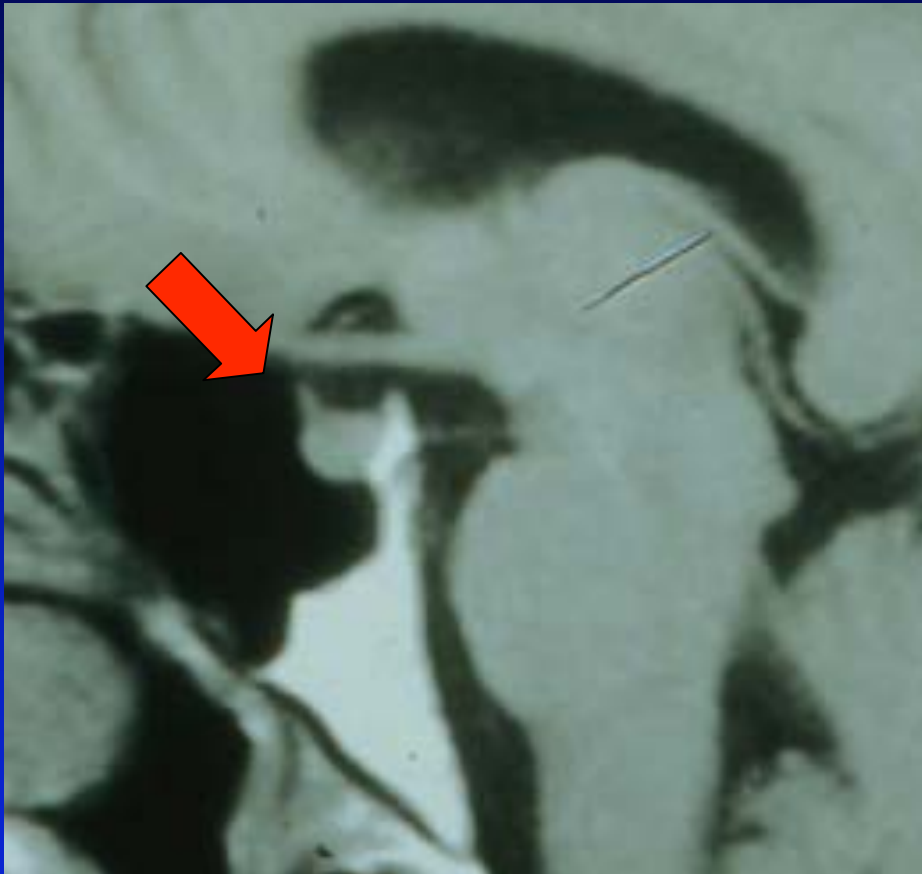


[GNU Free documentation license version 1.2](#)
[Wikimedia Commons](#)



Source: Undetermined

Pituitary Cushing's DISEASE



Source: Undetermined

normal

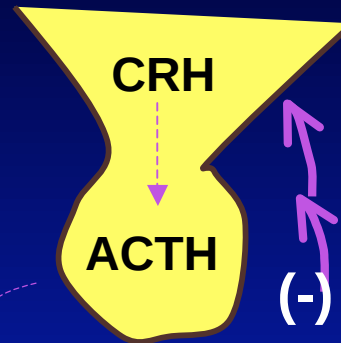


Source: Undetermined

Cushing's disease

Ectopic Cushing's

Hypercortisolism from
Ectopic production of
ACTH or CRH by tumor.



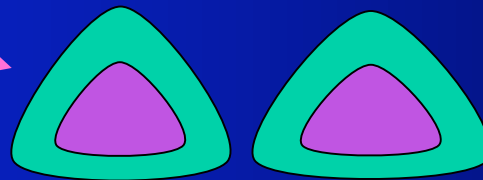
Hypothalamus

(-)

Pituitary

↑↑ **Cortisol**

ACTH



Images of oat cell
in lung removed

Small (Oat) cell ca of lung

Causes

- Bronchial carcinoid
- Oat cell carcinoma
- Thymic carcinoid
- Pheochromocytoma
- Medullary thyroid ca

Cushing's Syndrome

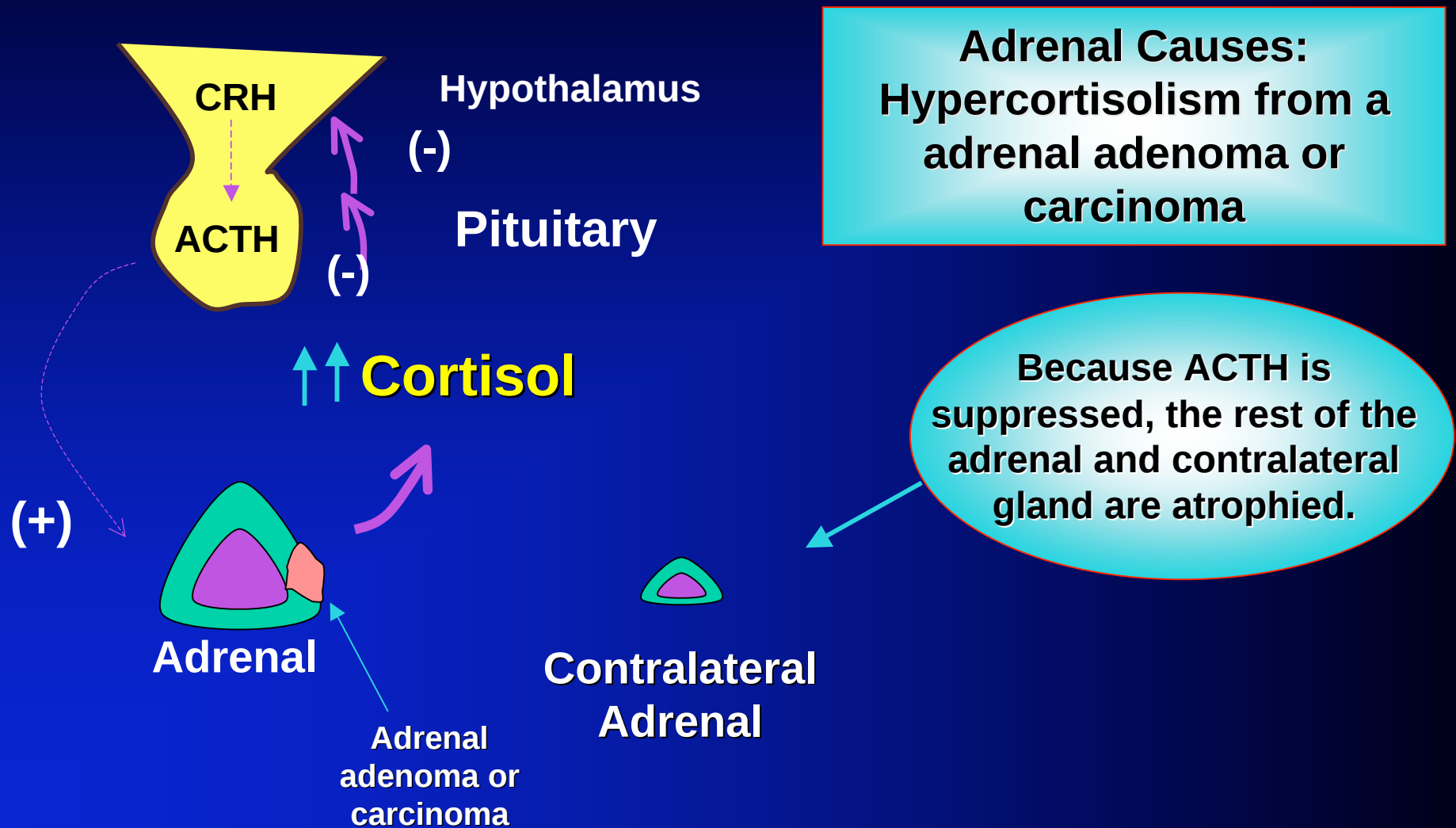


© Cornell University Medical College

[CC:BY-NC-SA 1.0](#)
[BY: Cornell University](#)

- **ACTH-independent Cushing's Syndrome**
 - **adrenal cortical neoplasm**
 - **adenoma**
 - **carcinoma**
 - **primary adrenal hyperplasia**

Adrenal Cushing's



Cushing's Syndrome

CLINICAL MANIFESTATIONS of CORTISOL EXCESS

- increased protein catabolism = striae, bruising, delayed wound healing, muscle wasting
- increased glucose production = DM
- redistribution of fat = truncal obesity
- bone breakdown = osteoporosis
- facilitation of catechol synthesis = hypertension
- anti-inflammatory = opportunistic infections
- Inhibition of HPG axis = amenorrhea, impotence
- CNS effects(limbic/hippocampus) = depression and memory difficulties

ACTH dependent ONLY

- Pigmentation (MSH)

ACTH dependent or Mixed Adrenal

- Androgen excess
 - Terminal hair hirsutism
 - Acne
 - Irregular menses
 - balding

Cushing's Syndrome

- **Physical examination:**
 - **adiposity**
 - **moon face, plethora**
 - **(pseudo-) gynecomastia**
 - **striae**



Image of patient
with striae on arms
removed



Image of patient
with moon face /
plethora removed

Cushing's Syndrome

- **Acanthosis nigricans**
- **Purple striae**



Image of patient
with acanthosis
nigricans on armpit
removed



Image of patient
with striae on
abdomen removed



Image of patient
with striae on
abdomen removed

Cushing's Syndrome

- **Myopathy**
 - Proximal muscle wasting
- Osteoporosis
- Oligo-Amenorrhea/Impotence
- Psychiatric Symptoms
 - depression, mania (Steroid psychoses)



Images of
myopathic
patient removed

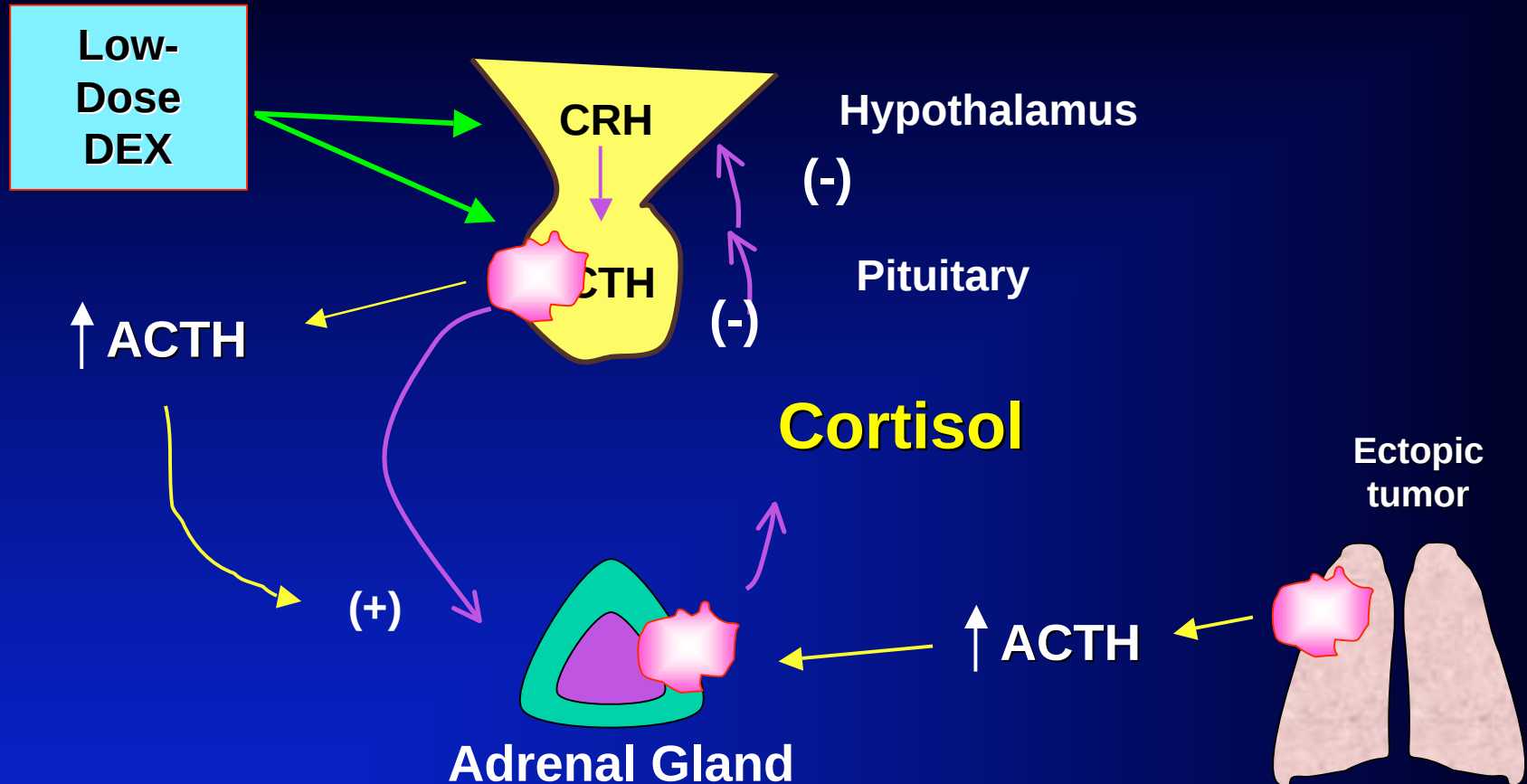
ACTH-Dependent Pituitary Cushing's Disease

- Symptoms due to pituitary mass
 - bitemporal hemianopsia
 - pituitary insufficiency
 - HA

Cushing's Syndrome: Diagnosis

- **Diagnosis**
 - **First diagnose CORTISOL EXCESS**
 - elevated 24 hr urine cortisol < 100 mg/24 hr
 - **Then diagnose PATHOLOGIC CORTISOL EXCESS**
 - r/o physiologic causes which suppress normally with low-dose DEX (Cort < 5 mg/dl)
- **ACTH dependent or NOT**
 - **Measure ACTH level**
 - if DETECTABLE > 9 pg/ml - must be ACTH dependent
 - (if NOT DETECTABLE < 9 pg/ml - must be ACTH independent)

Cushing's Syndrome: Low-dose DEX suppression



Low-dose Dex will suppress ACTH secretion in:

- normal patients
- physiologic hypercortisolism (stress)

Low-dose Dex will NOT suppress ACTH secretion in:

- ACTH dependent Cushing's syndrome (pituitary adenoma or ectopic ACTH producing tumors)
- ACTH independent Cushing's syndrome (adrenal tumors)

Cushing's Syndrome: Diagnosis

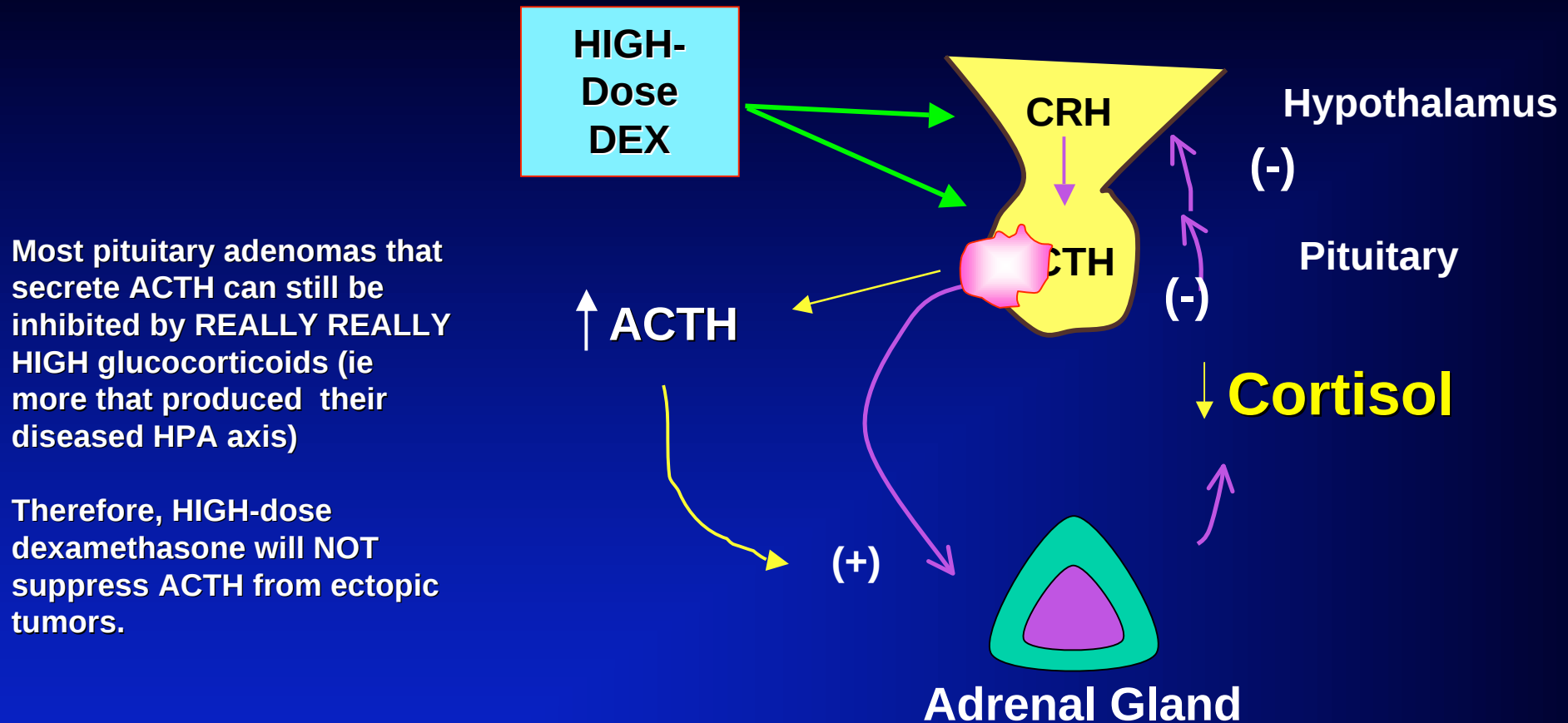
- **Diagnosis**
 - **First diagnose CORTISOL EXCESS**
 - elevated 24 hr urine cortisol < 100 mg/24 hr
 - **Then diagnose PATHOLOGIC CORTISOL EXCESS**
 - r/o physiologic causes which suppress normally with low-dose DEX (Cort < 5 mg/dl)
- **ACTH dependent or NOT**
 - **Measure ACTH level**
 - if DETECTABLE > 9 pg/ml - must be ACTH dependent
 - (if NOT DETECTABLE < 9 pg/ml - must be ACTH independent)

ACTH-DEPENDENT Cushing's Syndrome

Is it pituitary or ectopic????

- High dose DEX SUPPRESSION TEST
 - Pituitary Cushing's may suppress to high dose DEX
 - Ectopic NEVER suppresses to high dose DEX
- Inferior Petrosal sinus Sampling
 - Pituitary Cushing's - find HI ACTH near pituitary and low in the periphery
 - Ectopic Cushing's - find HI ACTH in the periphery and low near pituitary
- IMAGE the pituitary

Cushing's Syndrome: High-dose DEX suppression



Most pituitary adenomas that secrete ACTH can still be inhibited by REALLY REALLY HIGH glucocorticoids (ie more that produced their diseased HPA axis)

Therefore, HIGH-dose dexamethasone will NOT suppress ACTH from ectopic tumors.

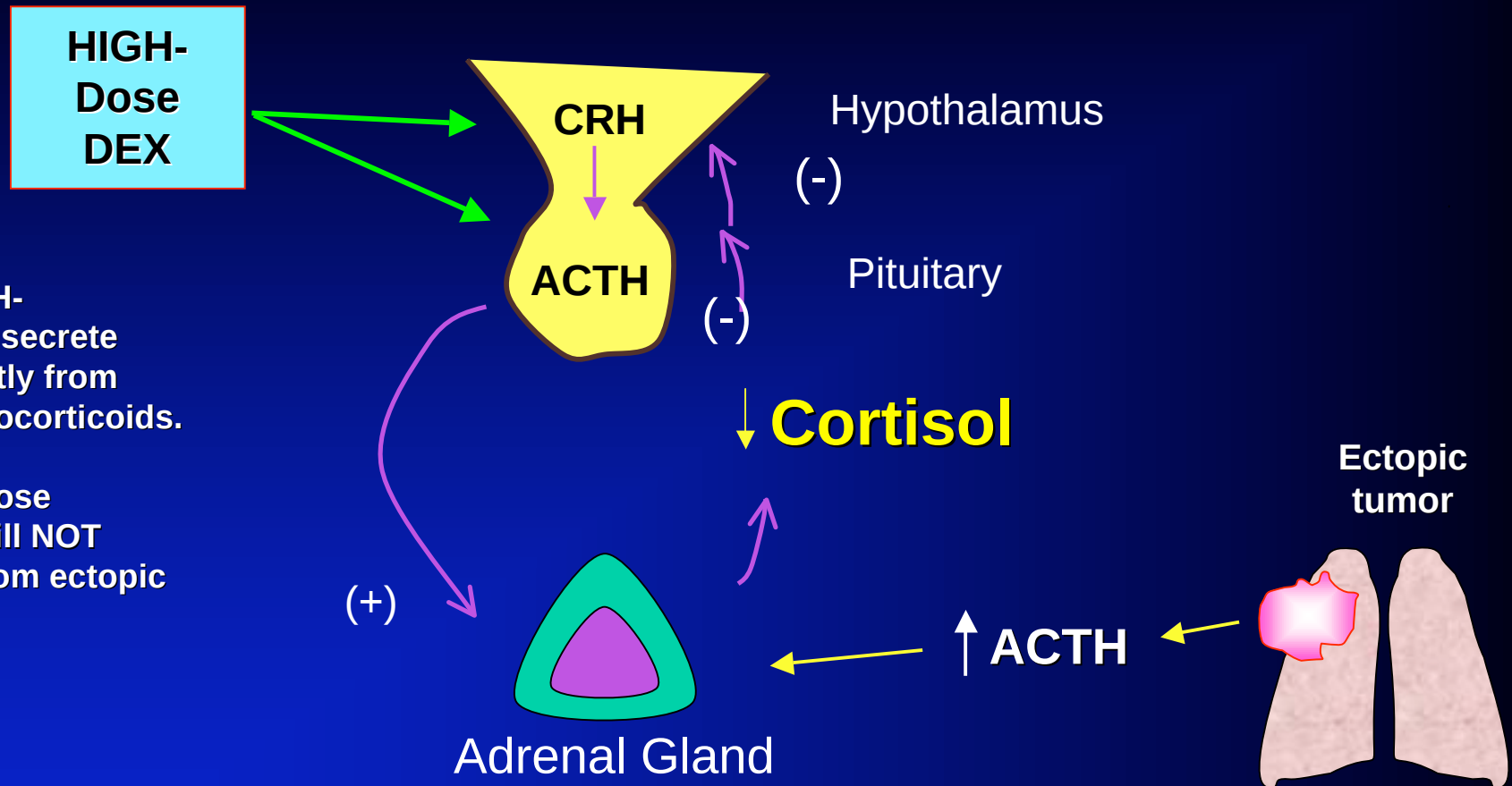
High-dose Dex will suppress ACTH secretion in:

-ACTH dependent Cushing's syndrome (pituitary adenoma)

High-dose Dex will NOT suppress ACTH secretion in:

-ACTH dependent Cushing's syndrome (ectopic tumors)

Cushing's Syndrome: Diagnosis



Most ectopic ACTH-producing tumors secrete ACTH independently from regulation by glucocorticoids.

Therefore, HIGH-dose dexamethasone will NOT suppress ACTH from ectopic tumors.

High-dose Dex will suppress ACTH secretion in:
-ACTH dependent Cushing's syndrome (pituitary adenoma)

High-dose Dex will NOT suppress ACTH secretion in:
-ACTH dependent Cushing's syndrome (ectopic tumors)

Imaging in Cushing Syndrome

- ADRENAL CT findings

- adrenals small = ?
- one adrenal large and 1 small =?
- Both adrenals large=?

- Pit MRI findings

- Mass or no mass
(some pituitary corticotrope tumors are too small to be seen on MRI)

- Search for ectopic ACTH or CRH producing tumor

- | | |
|---|-----|
| ▪ Lung: Bronchial Carcinoid and SCC | 50% |
| ▪ Thymic Carcinoid (epithelial thymoma) | 10% |
| ▪ Pancreatic Islet Cell Tumor | 10% |
| ▪ Pleochromocytoma | 10% |
| ▪ Abdominal Carcinoids | 5% |
| ▪ Medullary Thyroid Carcinoma | 5% |

Cushing's Syndrome Treatment

- adrenal adenoma

- resection
- cortisol replacement
- if not curative
 - XRT
 - bilateral adrenalectomy
 - adrenolytic therapy
 - mitotane
 - ketoconazole

- pituitary adenoma

- transphenoidal resection (TSR)
- cortisol replacement
- if not curative
 - XRT
 - bilateral adrenalectomy
 - adrenolytic therapy
 - mitotane
 - ketoconazole

- Ectopic ACTH or CRH

- Find the tumor!!!!!!!!!!!!
- if not curative
 - bilateral adrenalectomy
 - adrenolytic therapy
 - mitotane
 - ketoconazole

Cushing's Syndrome

before treatment

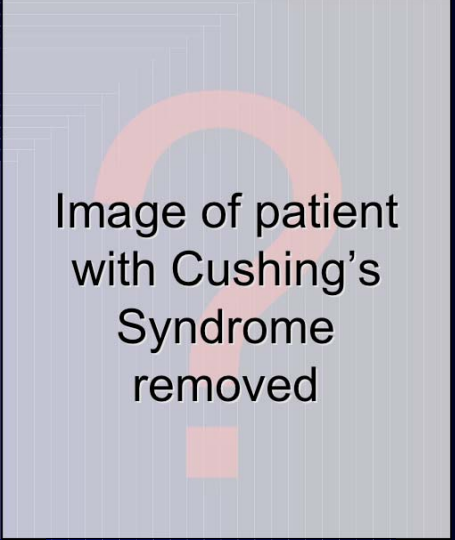


Image of patient
with Cushing's
Syndrome
removed

after treatment

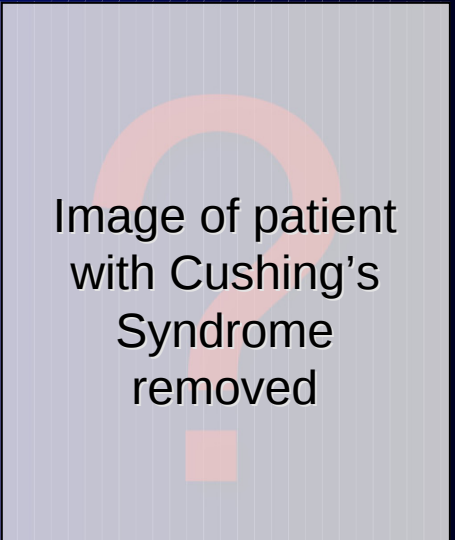


Image of patient
with Cushing's
Syndrome
removed

Cushing's Syndrome



[CC:BY 2.0](#)
[BY: tajai](#)



Public Domain
[Wikimedia Commons](#)



Public Domain
[Wikimedia Commons](#)

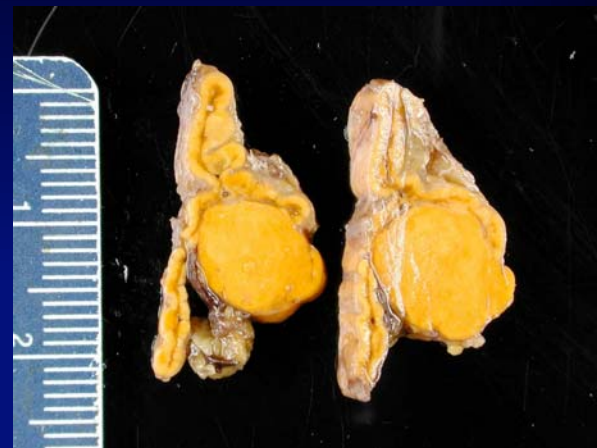
dogs
Ferrets
horses

UM Endocrinology in Adrenal History: Conn Syndrome

Image of patient
removed

Image of Jerome
Conn removed

Jerome Conn, M.D.



CC:BY 2.0
BY: Michael Feldman, MD

PRESIDENTIAL ADDRESS

PART I. PAINTING BACKGROUND

PART II. PRIMARY ALDOSTERONISM, A NEW CLINICAL SYNDROME*

JEROME W. CONN, M.D.

ANN ARBOR, MICH.

From the Metabolic Research Unit, Department of Internal Medicine, University of Michigan Medical School, Ann Arbor, Mich.

Presented at the Twenty-seventh Annual Meeting of the Central Society for Clinical Research, Chicago, Ill., Oct. 29, 1954.

Conn JW. **Primary aldosteronism, a new clinical syndrome.** J Lab Clin Med. 1955;45:3-17

Primary Aldosteronism

- Clinical Presentation

- Manifestations of HYPOKALEMIA and HTN

- LOW K

- neuromuscular

- paresthesias

- weakness

- tetany

- Renal

- Polyuria

- Carbohydrate

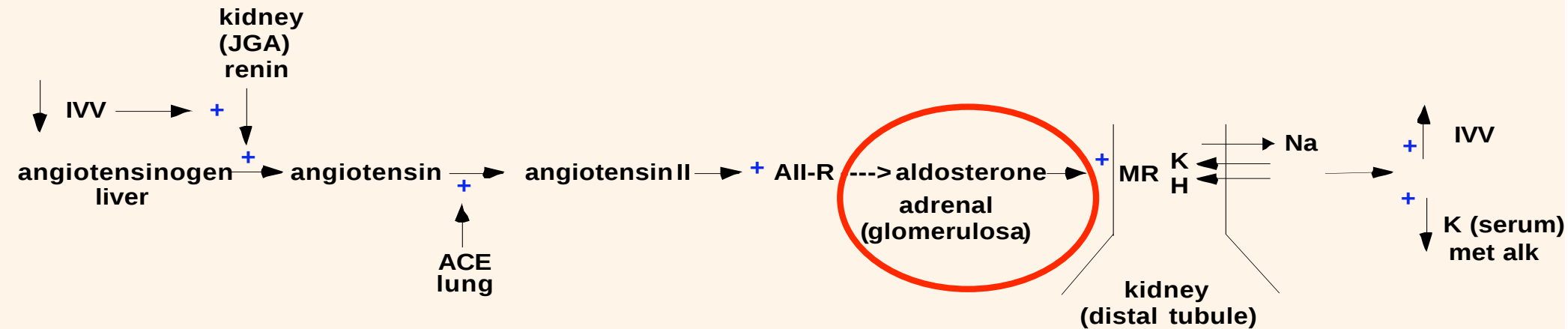
- abnormal GTT

- HTN usually not malignant

- early in disease may have HTN with NORMAL K

Causes of Hyperaldosteronism

Definition: syndrome of inappropriate excessive secretion of aldosterone by adrenal gland



Source: Undetermined

An increase in aldosterone ACTION can theoretically result from ANY defect in RAA pathway

- LOW IVV (real or perceived by kidney in renal artery stenosis)
- JGA renin tumor
- ACE polymorphisms
- overproduction of All by renal tumors
- ADRENAL overproduction of ALDO**
- constitutive MR or Na channel

Primary Aldosteronism

primary hyperaldosteronism (HI ALDO/LOW RENIN)

- ZG Aldo tumor 70%
- ZG Aldo hyperplasia 30%
- rare/rare/rare
 - Congenital adrenal hyperplasia <1%
(p450c11 β , p450c17)
 - ACE polymorphisms < 1%
 - All overproduction < 1%

secondary hyperaldosteronism (HI ALDO/HI RENIN)

- JGA renin tumor <1%
- renal artery stenosis <1%

apparent mineralocorticoid excess (LOW ALDO/LOW RENIN) (downstream of ALDO)

- constitutively active MR <1%
- Na/K/H channel <1%
- licorice <1%

Primary Aldosteronism

Consider in patients with:

- New HTN
- HTN with LOW K

EVEN THOUGH it only accounts for 0.5% of all HTN

**BECAUSE- IF YOU NEVER THINK OF THIS--
---YOU WILL NEVER FIND IT!!!**

Primary Aldosteronism

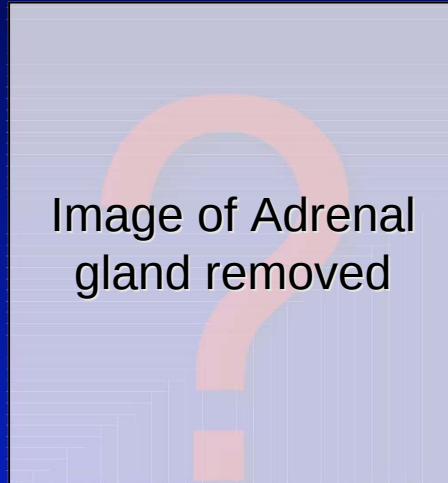
- Work-Up

- R/O other causes of LOW K
 - LOW intake (diet)
 - HI output
 - N/V/D
 - Diuretic use with loops + thiazides

- 24 h Urine ALDO

- If LOW- pt does not have PRIMARY ALDO
- IF HI (>10 ug/day)
 - check RENIN level (suppressed < 1 ng/ml/hr)
 - If RENIN HI ----JGA renin tumor or RAS
 - If RENIN LOW---- PRIMARY HYPERALDO
- IF NECESSARY (ie AMBIGUOUS) Volume expand to see if can suppress RAA
 - If can suppress --essential HTN

Adrenal Zona Glomerulosa Adenoma

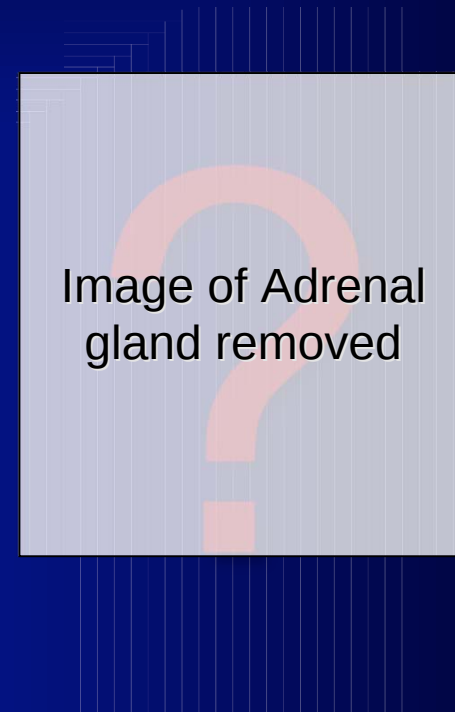


Primary Aldosteronism

- **IMAGING and TREATMENT**
 - CT scan
 - Adenoma
 - unilateral ADX
 - NO adenoma
 - selective venous cath to measure ALDO rt vs lt
 - If unilateral elevation-small adenoma
 - If no lateralization-bilateral hyperplasia
 - Medical trt with spironolactone or amiloride
 - bilateral ADX

Adrenocortical Carcinoma

- Larger adrenal mass
 - High probability NOT benign if >5 cm in diameter
 -
 - Development of Cushingoid features usually very rapid (several months rather than years)
- Often associated with elevated DHEA-sulfate and virulization



Remember:

Endocrine disorders are NOT diagnosed by means of imaging studies. Biochemical confirmation must come first before imaging is performed.



Public Domain
[Wikimedia Commons](#)

“Even our destiny is determined by our endocrine glands.”

Albert Einstein