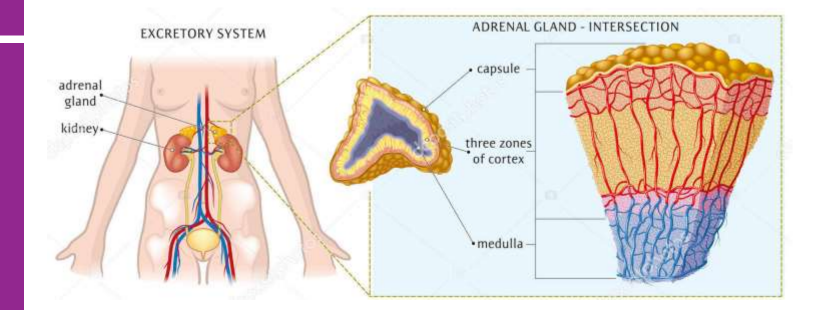
Endocrine system pathology ADRENAL GLAND

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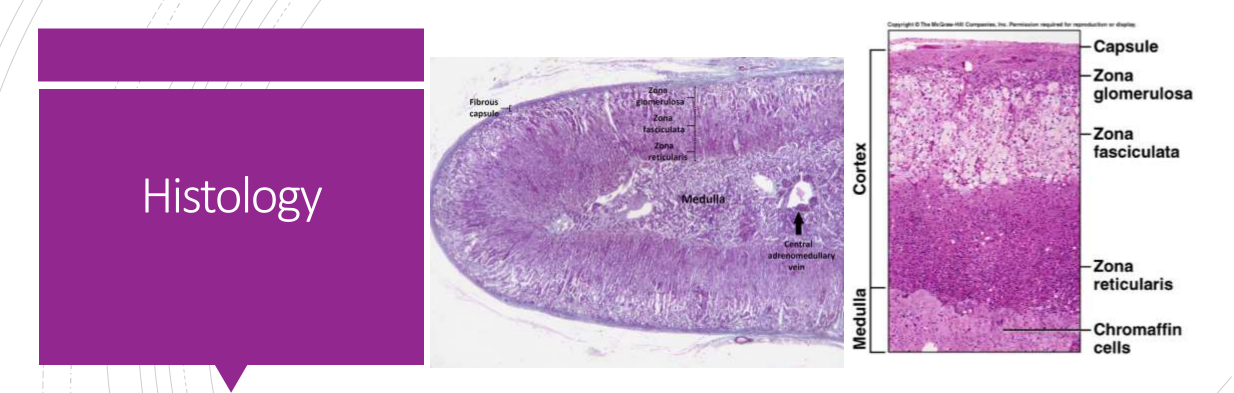
Anatomy



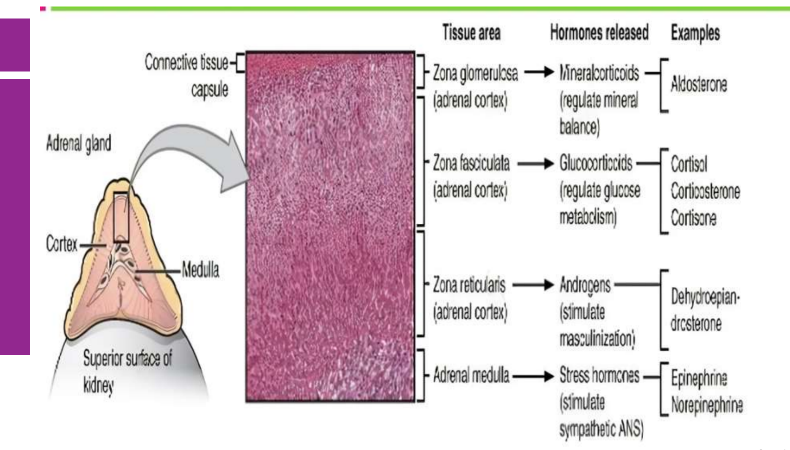








Hormones.



Adrenal disorders

Non-neoplastic:

- ✤Adrenal insufficiency:
- > Acute Adrenocortical Insufficiency.
- > Chronic Adrenocortical Insufficiency: Addison Disease.
- Adrenal hyperfunction:
- Cushing Syndrome.
- > Hyperaldosteronism.
- Adrenogenital Syndromes

Neoplastic:

- Adrenalcortical tumors.
- Adrenal medulla tumor

Cushing syndrome.

- Hypercortisolism (Cushing syndrome) is caused by elevated glucocorticoid levels.
- Endogenous causes :
- ✓ Hypothalamic/ pituitary hypersecretion ACTH.
- ✓ Paraneoplastic syndrome (lung CA).
- ✓ Adrenal tumor or hyperplasia .

- Exogenous cause :
- ✓ Steroid Therapy.

Signs and symptoms

- high blood pressure.
- abdominal obesity but with thin arms and legs.
- reddish stretch marks.
- round red face.
- fat lump between the shoulders.
- weak muscles and weak bones.
- acne and fragile skin







Morphology

 Morphologic changes in the adrenal glands depend on the cause of the hypercortisolism and include:

(1) cortical atrophy:

exogenous glucocorticoids, suppression of endogenous ACTH.

(2) diffuse hyperplasia:

ACTH dependent Cushing syndrome

(3) macronodular or micronodular hyperplasia:

primary cortical hyperplasia.

(4) an adenoma or carcinoma.

Hyperaldostero nism.

- Hyperaldosteronism is the generic term for a group of closely related conditions characterized by chronic excess aldosterone secretion.
- A-primary hyperaldosteronism are:
- Bilateral idiopathic hyperaldosteronism, characterized by bilateral nodular hyperplasia of the adrenal glands.
- Adrenocortical neoplasm, either an aldosterone-producing adenoma (the most common cause) or, rarely, an adrenocortical carcinoma.
- familial hyperaldosteronism may result from a genetic defect that leads to overactivity of the aldosterone synthase gene, CYP11B2.
- B- Secondary causes :
- Due to decreased renal perfusion (heart failure), activation of the renin angiotensin system.

Adrenogenital syndromes

- Adrenogenital syndromes refer to a group of disorders caused by androgen excess, which may stem from a number of etiologies, including primary gonadal disorders and several primary adrenal disorder.
 Could be caused by :
 - 1 Primary gonadal disorders(increase gonadal androgen).
 - 2 -Acquired :Adrenocortical Neoplasms. can occur at any age, frequently malignant .

CLINICAL FEATURES

- Virilization in female or precocious puberty in male.
- Patients have risk for acute adrenocortical insufficiency.
- Note :Adrenal androgen formation is regulated by ACTH , thus increase androgen can occur as apure syndrome or as a component of cushing syndrome.

ADRENOCORTIC AL INSUFFICIENCY

- May be primary adrenal(disease affecting the adrenal gland) :
- acute (called adrenal crisis).
- chronic (Addison disease).
- secondary to destruction of the pituitary as in SHEEHAN's syndrome or non functional pituitary adenoma

Fable 20.7 Causes of Adrenal Insufficiency

Acute

Waterhouse-Friderichsen syndrome	
Sudden withdrawal of long-term corticosteroid therapy	
Stress in patients with underlying chronic adrenal insuffic	iency
Chronic	
Autoimmune adrenalitis (60%–70% of cases in developed includes APSI (AIRE mutations) and APS2 (polygenic)	countries)—
nfections Tuberculosis Acquired immunodeficiency syndrome Fungal infections	
Hemochromatosis	
Sarcoidosis	
Systemic amyloidosis	
Metastatic disease	

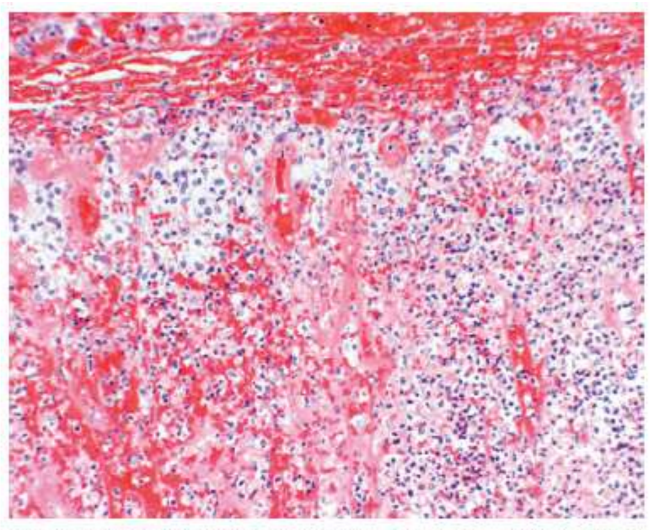


Figure 19–39 Waterhouse-Friderichsen syndrome. Bilateral adrenal hemorrhage in an infant with overwhelming sepsis, resulting in acute adrenal insufficiency. At autopsy, the adrenals were grossly hemorrhagic and shrunken; in this photomicrograph, little residual cortical architecture is discernible.

Chronic :(Addison's disease)

-Chronic adrenal cortical insufficiency , required immediate therapy .

-Progressive destruction of the adrenal. Causes include:

1-Autoimmune - 60-70 %, may be sporadic or

familial, linked to HLA-B8 or DR3

Often multisystem involvement.

2- Infections e.g. Tuberculosis, fungi.

3- Metastatic tumors destroying adrenal e.g. lung ,breast , ...others

4-AIDS.

Morphology & Clinical features in Chronic Adrenal Insufficiency :

Morphology depends on cause :

Autoimmune shows irregular small glands, with cortex heavily infiltrated by lymphocytes, medulla normal.

In T.B. \rightarrow Caseating Granuloma

In metastatic CA \rightarrow Type of primary tumor

In secondary to pituitary cause, the adrenal is shrunken

In general, clinical manifestations of adrenocortical insufficiency do not appear until at least 90% of the adrenal

Adrenal tumor:

✓ ADRENALCORTICAL TUMORS

✓ THE ADRENAL MEDULLA TUMOR

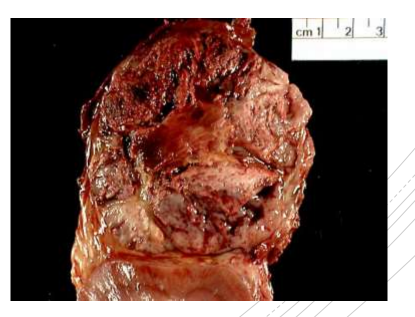
ADRENALCOR TICAL TUMORS

- Malignant epithelial tumor of adrenal cortical cells
- Adrenocortical carcinoma (ACC) is a rare endocrine tumor with high mortality
- More often involves left adrenal: left to right ratio = 1.2:1
- Functional adrenal cortical carcinomas have the following symptoms related to hormone production:
- 50% cortisol excess (Cushing syndrome, rapid onset)
- 20% sex hormone secretion (mainly androgens causing hirsutism, virilization and menstrual irregularities)
- *8% aldosterone (hypertension, hypokalemia)

Morphology

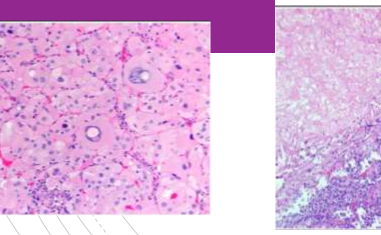
- Encapsulated , usually yellow color: single or multiple.
- Size variable 1-2 cm. Up to large tumors
- Malignant tumors may show necrosis, hemorrhage and

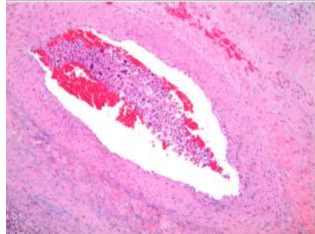
are usually larger.



Histology

- Encapsulated tumor composed of variably sized nests, large sheets and trabeculae
- Invasion of thick fibrous capsule
- Lymphovascular invasion (venous or sinusoidal)
- Areas of necrosis, hemorrhage, degeneration are common





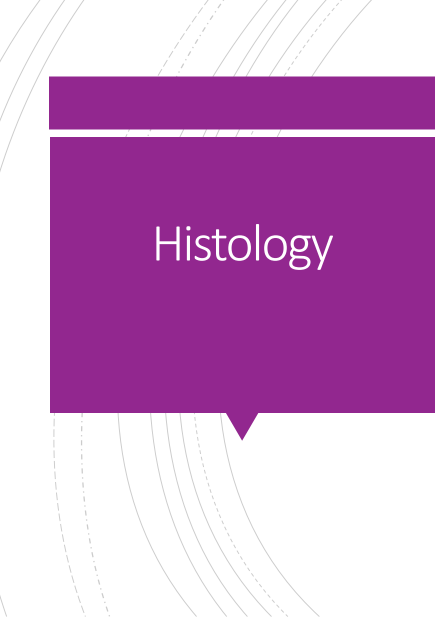
Tumor of the adrenal medulla *PHEOCHRO MOCYTOMA*

- Pheochromocytomas are neoplasms composed of chromaffin cells, which, like their nonneoplastic counterparts, synthesize and release catecholamin.
- Sometimes described as Ruleof 10% Tumor because :
 - * 10% bilateral.,
- 10 %multiple,
 - 10% non functional
 - * 10% familial, may be part of MEN syndrome.
 - * 10% Malignant.
 - * 10% extraadrenal site.
 - *25% associated with genetic mutation.

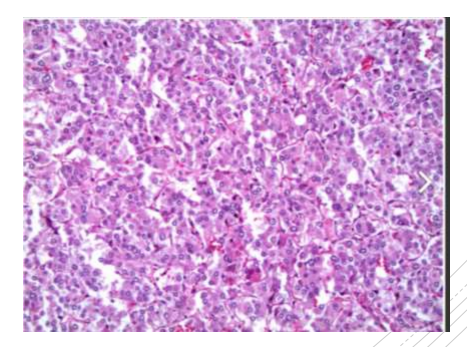
Morphology

well circumscribed, small to large in size





- Nested (zellballen), trabecular patterns.
- Nests of cells (Zellballen) with abundant cytoplasm filled with granules containing catecholamine.
- Malignancy confirmed by <u>METASTASES</u>



Thank you