

Modified by: Nour Hussein



HYPERTENSIVE VASCULAR DISEASE

Arteriolosclerosis

→ Affects mainly
Arterioles.

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Hypertension(HTN)

- Cutoffs in diagnosing hypertension in clinical practice → sustained diastolic pressures >90 mm Hg, and/or sustained systolic pressures >140 mm Hg

- **Malignant hypertension**
 - 5% of HTN patients present with a rapidly rising blood pressure that, if untreated, leads to death within 1 to 2 years.
 - * usually due to patients not complying with their medications.
 - **systolic pressures \geq 200 mm Hg or diastolic pressures \geq 120 mm Hg**
 - either readings, or both
 - associated with renal failure and retinal hemorrhages
 - most commonly is superimposed on preexisting benign hypertension

Hypertension (HTN) has the following potential complications: "untreated hypertension."

- stroke (CVD)
- multi-infarct dementia
- atherosclerotic coronary heart disease
 - (*hypertensive heart disease*)
- cardiac hypertrophy and heart failure
- aortic dissection
- renal failure

Ischemic
Heart Disease

Types of hypertension

- 1- **essential (idiopathic) hypertension (95%)**
- 2- **secondary hypertension:** Most are due to renal disease, or renal artery narrowing (= renovascular hypertension), and to a lesser degree are due to many other conditions....

Essential Hypertension

Accounts for 90% to 95% of all cases

Secondary Hypertension

Renal

Acute glomerulonephritis

Chronic renal disease

Polycystic disease

Renal artery stenosis

Renal vasculitis

Renin-producing tumors

Endocrine

Adrenocortical hyperfunction (Cushing syndrome, primary aldosteronism, congenital adrenal hyperplasia, licorice ingestion)

Exogenous hormones (glucocorticoids, estrogen [including pregnancy-induced and oral contraceptives], sympathomimetics and tyramine-containing foods, monoamine oxidase inhibitors)

Pheochromocytoma

Acromegaly

Hypothyroidism (myxedema)

Hyperthyroidism (thyrotoxicosis)

Pregnancy-induced (pre-eclampsia)

Cardiovascular

Coarctation of the aorta

Polyarteritis nodosa

Increased intravascular volume

Increased cardiac output

Rigidity of the aorta

Neurologic

Psychogenic

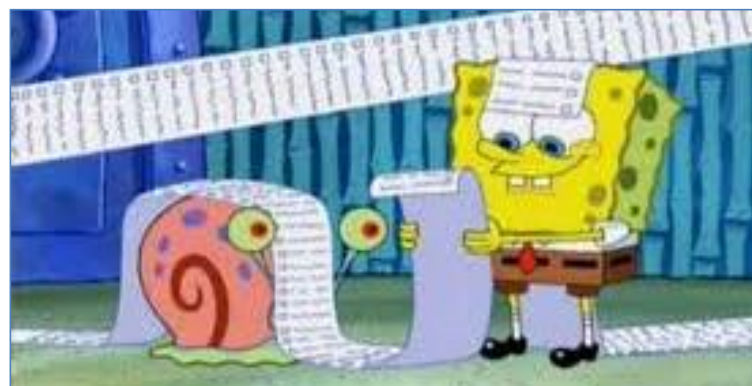
Increased intracranial pressure

Sleep apnea

Acute stress, including surgery

Most common of all

Most common of secondary causes



- *Pathogenesis of essential HTN*
- ? **Genetic factors**
- ? **familial clustering of hypertension**
- angiotensinogen polymorphisms and angiotensin II receptor variants; polymorphisms of the renin-angiotensin system.
- ? Susceptibility genes for essential hypertension: genes that control renal sodium absorption, etc...
** mutations in enzymes, proteins involved in fluid and salt balance in persons body.*

Pathogenesis of essential HTN

- **Environmental factors** → Aggravate hypertension.
- stress, obesity, smoking, physical inactivity, and high levels of salt consumption, modify the impact of genetic determinants.
- Evidence linking dietary sodium intake with the prevalence of hypertension in different population groups is particularly strong.

Morphology

- HTN is associated with **arteriolosclerosis** (small arterial disease)
 - Two forms of small blood vessel disease are hypertension-related:
 - 1- **hyaline arteriolosclerosis** → Benign.
 - 2- **hyperplastic arteriolosclerosis** → Malignant
- different forms of HT

→ Hardening of Arteriole wall
due to Hyalin material

1- Hyaline arteriolosclerosis

- Ass. with benign hypertension
- homogeneous, pink hyaline thickening of the arteriolar walls; luminal narrowing
- leakage of plasma components across injured endothelial cells into vessel walls and increased ECM production by smooth muscle cells in response to chronic hemodynamic stress.

- ① Arterioles
- ② Benign Hypertension
- ③ Over a long period of time.

→ Progression of
this disease is
slow

- **Hyaline arteriolosclerosis: Complications**

- **Most significant in kidneys → nephrosclerosis (glomerular scarring)**
+ Fibrosis.

- Other causes of hyaline arteriolosclerosis:

- 1- **elderly patients (normo-tensive)**

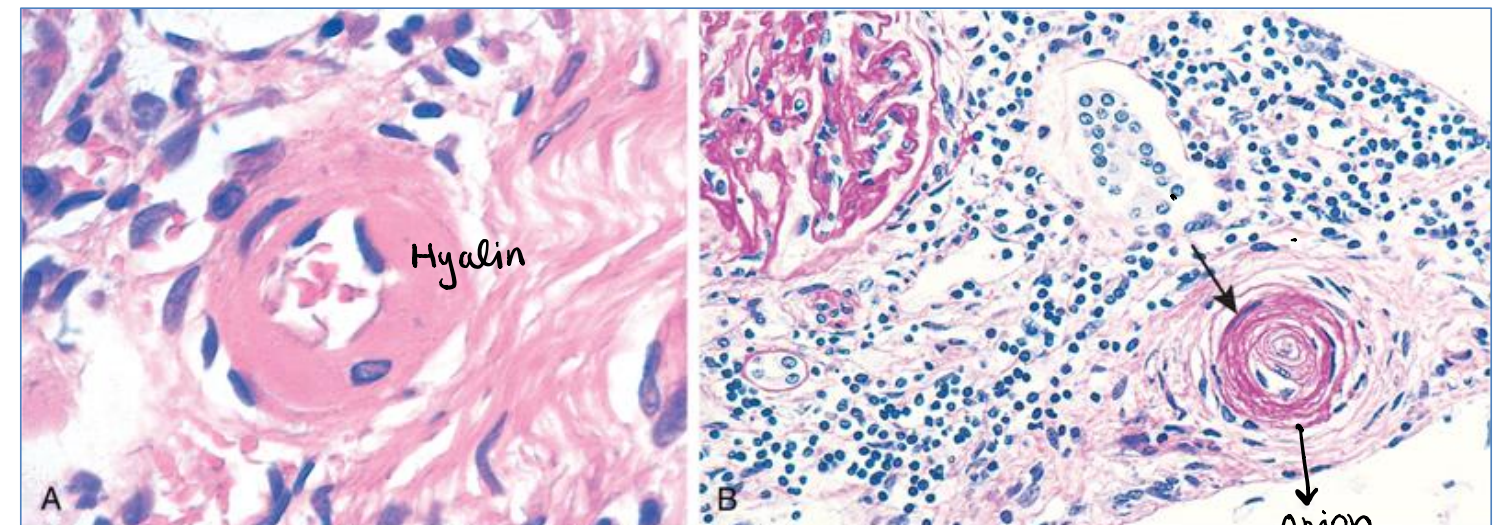
- 2- **diabetes mellitus**

+ Benign hypertension (as we said)

2- Hyperplastic arteriolosclerosis

- With severe (malignant) hypertension.
- "onionskin" concentric laminated thickening of arteriolar walls → luminal narrowing.
- = smooth muscle cells and thickened, reduplicated basement membrane. }^{Layering.}
- In malignant hypertension → fibrinoid vessel wall necrosis (necrotizing arteriolitis), which are particularly prominent in the kidney

A, Hyaline arteriolosclerosis. The arteriolar wall is thickened with the deposition of amorphous proteinaceous material, and the lumen is markedly narrowed. → ischemia of downstream organ



Kumar et al: Robbins Basic Pathology, 9e.
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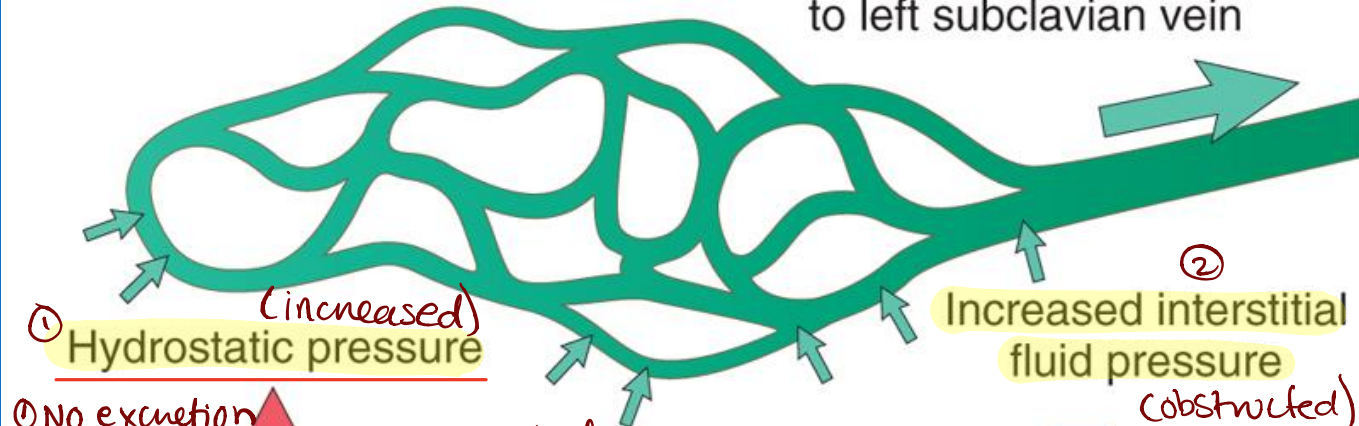
EDEMA

EDEMA

- 60% of lean body wt. = water
 - (2/3) intracellular.
 - (1/3) extracellular (interstitial fluid)
 - 5% blood plasma.
- ***edema* = accumulation of interstitial fluid within tissues.**
- ***Edema ≠ Extravascular fluid collection in body cavities:***
 - pleural cavity (*hydrothorax*)
 - the pericardial cavity (*hydropericardium*)
 - peritoneal cavity (*hydroperitoneum*, or *ascites*).

LYMPHATICS

To thoracic duct and eventually to left subclavian vein



Edema = Abnormality in one of these forces.

Nourishment.

Arterial end

Helps exit of fluid from Blood vessel to tissues.

- ① ↑ Synthesis (liver)
- ② ↑ Loss (kidney)
- ③ ↓ Synthesis (GI) ← malabsorption.

Helps trap fluid inside Blood vessel.

Plasma colloid osmotic pressure

(decreased)
Reduction in Plasma proteins.

Venous end

CAPILLARY BED

Mechanisms of edema

Increased Hydrostatic Pressure

Impaired Venous Return

Congestive heart failure; Constrictive pericarditis; Ascites (liver cirrhosis); Venous obstruction or compression; Thrombosis; External pressure (e.g., mass); Lower extremity inactivity with prolonged dependency

Arteriolar Dilation

Heat; Neurohumoral dysregulation

Reduced Plasma Osmotic Pressure (Hypoproteinemia)

Protein-losing glomerulopathies (nephrotic syndrome)
Liver cirrhosis (ascites); Malnutrition; Protein-losing gastroenteropathy

Lymphatic Obstruction

Inflammatory; Neoplastic; Postsurgical; Postirradiation

Sodium Retention

Excessive salt intake with renal insufficiency

Increased tubular reabsorption of sodium

Renal hypoperfusion

Increased renin-angiotensin-aldosterone secretion

Inflammation

Acute inflammation; Chronic inflammation; Angiogenesis

Local Causes.

Clinical Correlation of edema

- **Subcutaneous edema:**
 - the most common;
 - important to recognize as it signals potential underlying cardiac or renal disease
 - Can impair wound healing or the clearance of infections.

- **Brain edema:**
 - life-threatening → brain *herniation* (extrude) e.g. through the foramen magnum.

- **Pulmonary edema:** Accumulation of fluid in parenchyma of lung.
 - Common causes:
 - left ventricular failure - renal failure - ARDS
 - inflammatory and infectious disorders of the lung.
 - can cause death by interfering with normal ventilatory function & impeding oxygen diffusion
 - creates a favorable environment for infections