

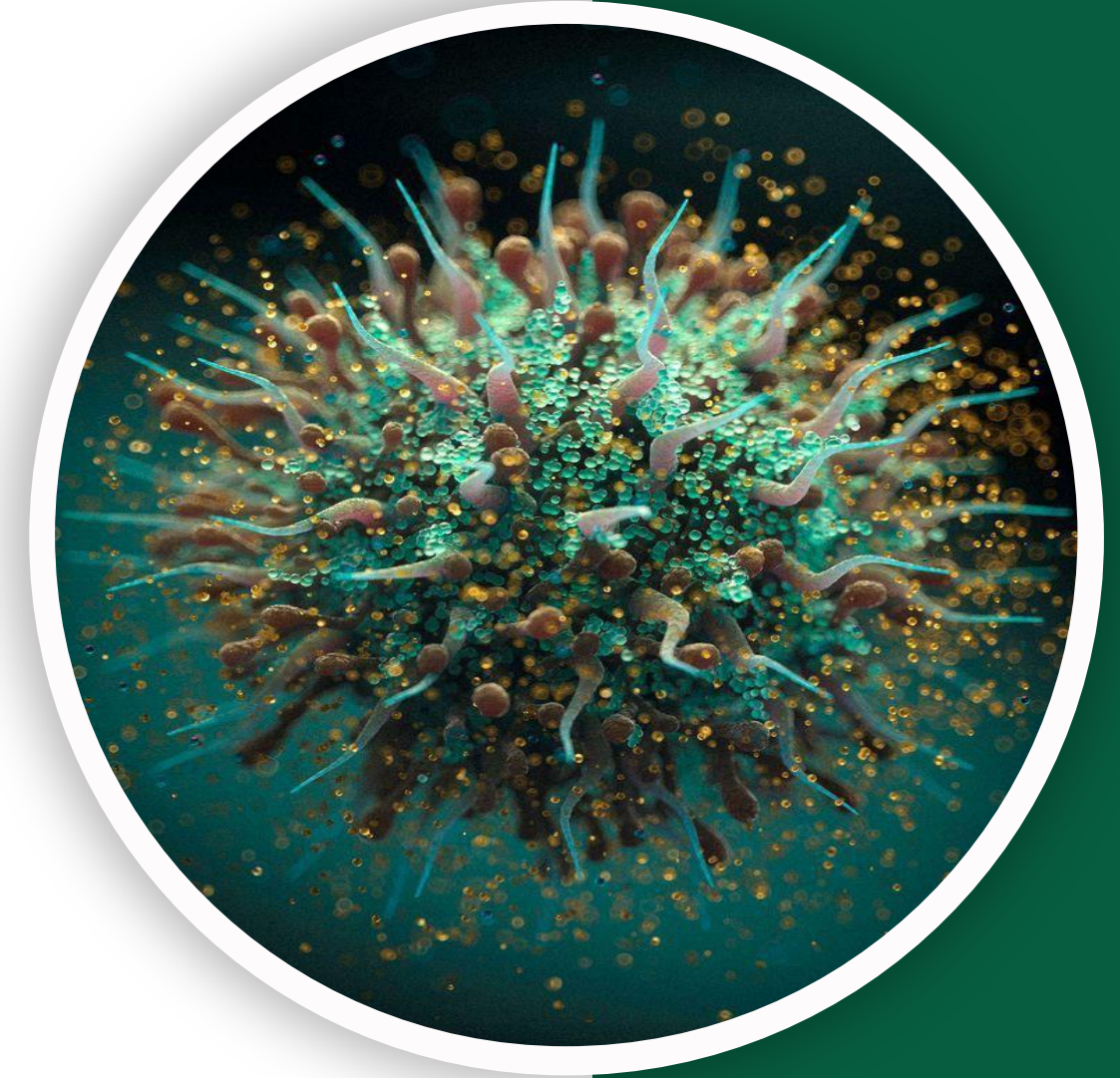
بِسْمِ اللّٰهِ الرَّحْمٰنِ الرَّحِیْمِ  
(وَفَوْقَ كُلِّ ذِي عِلْمٍ عَلِيمٌ)



جِلْدَانِ

GIS Pathology | MID 1

# Esophageal Diseases 1



Written by : DST

Reviewed by : Layan Bassam

Ruqaiya Moqbel

# Diseases of the esophagus-1

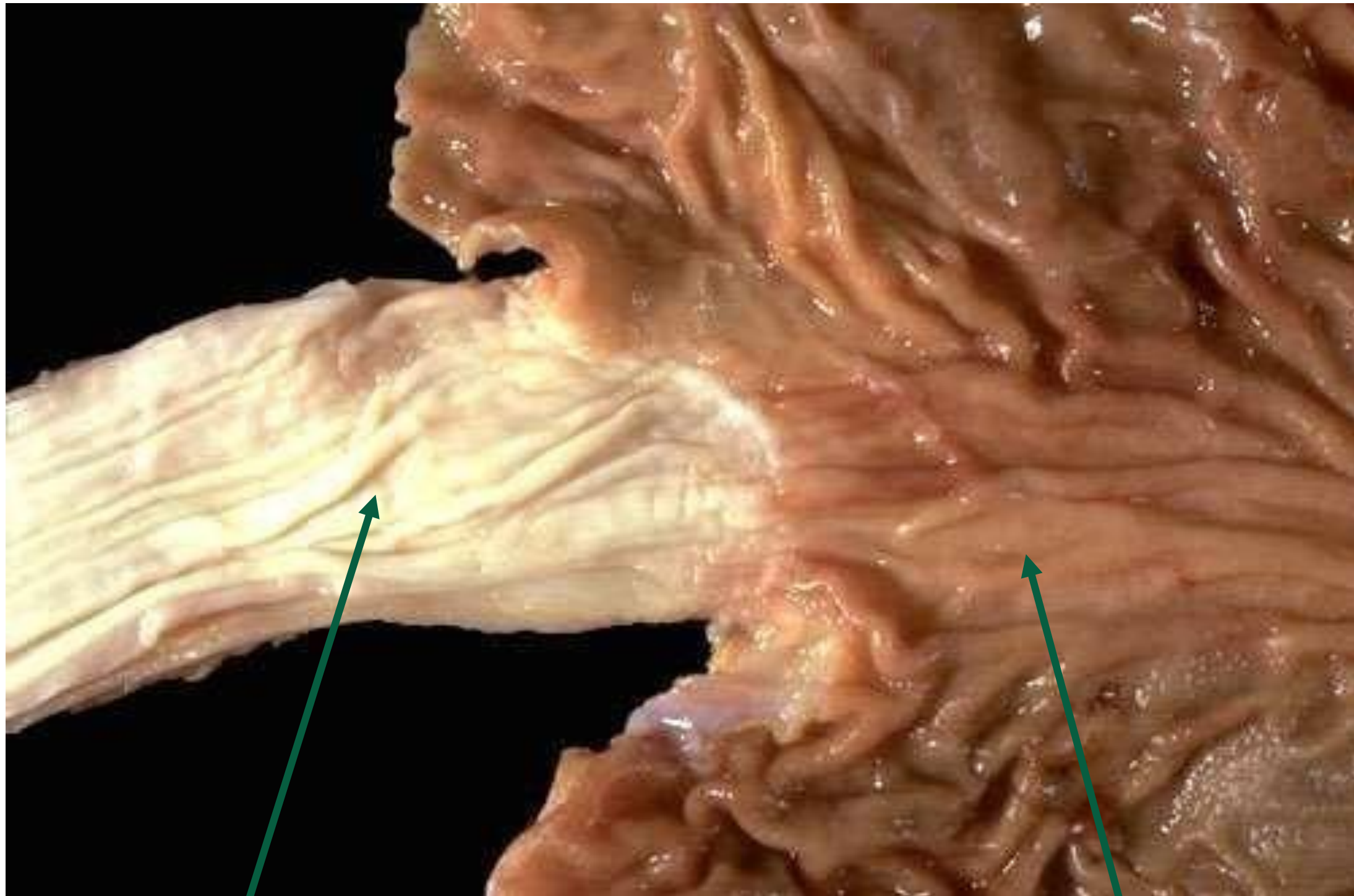
Manar Hajeer, MD, FRCPath University of  
Jordan, School of medicine

# Anatomy and histology:

Muscular tube  
extending from  
the epiglottis  
superiorly to the GEJ  
inferiorly.

Lined by stratified  
squamous  
epithelium in normal  
condition .

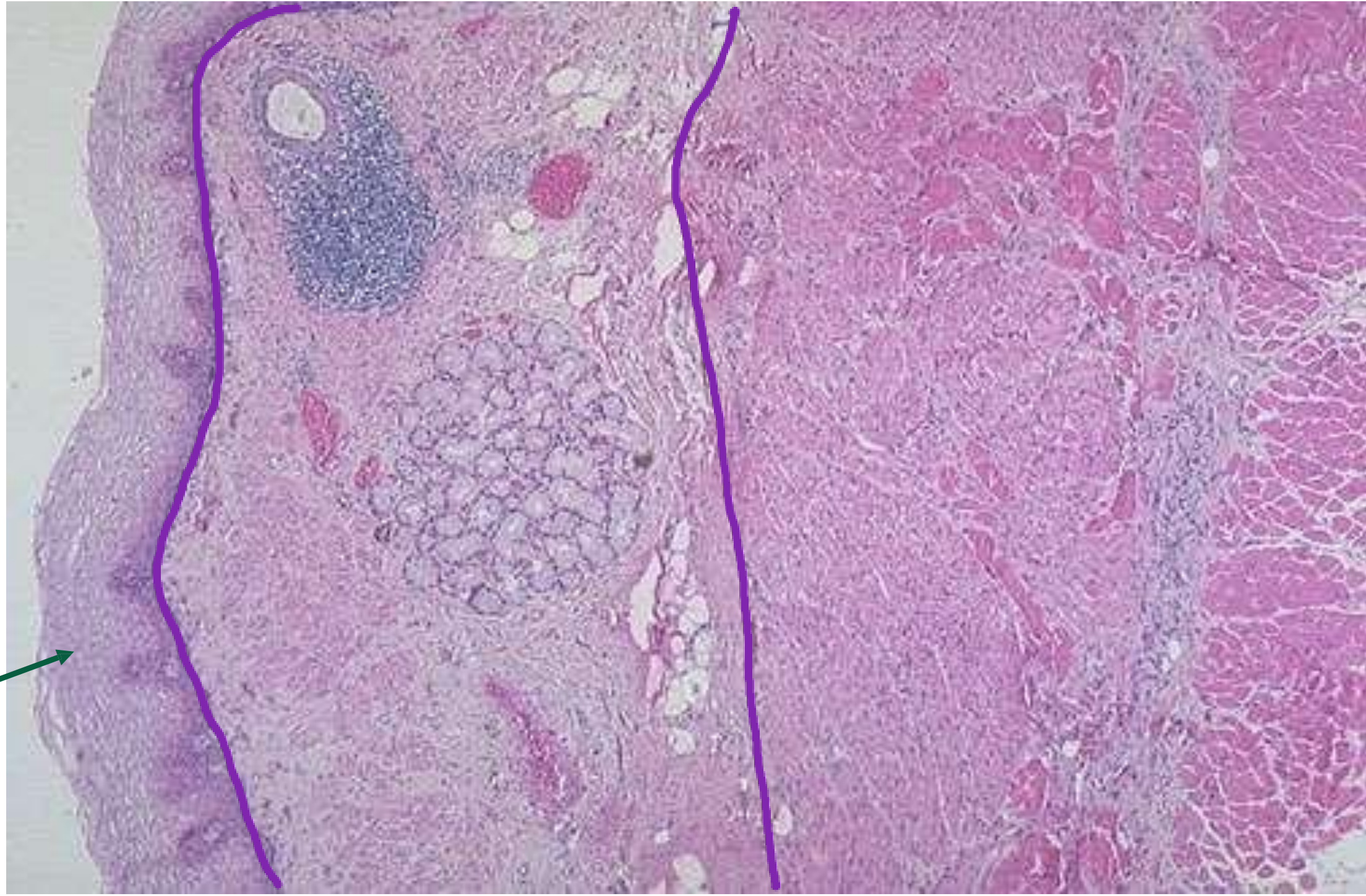
GEJ: gastroesophageal  
junction Junction = Sphincter



**Normal** esophageal mucosa which has Tan to Pale Pink color.

In contrast with the **normal** gastric mucosa which has a light brown color.

Stratified squamous epithelium lining



Submucosal layer

Muscularis propria

# Diseases that affect the esophagus

- 1. Obstruction: mechanical or functional.**
- 2. Vascular diseases: varices.**
- 3. Inflammation: esophagitis.**
- 4. Tumors.**

# -1 Mechanical Obstruction

- Congenital or acquired.
- Examples:

- **Atresia**
  - **Fistulas**
  - **Duplications**
  - **Aggenesis (very rare)**
  - Stenosis.
- usually congenital
- Atresia, Fistula, Duplication usually they present shortly after birth & they are non-compatible with eating and drinking or swallowing food.

↳ Most of the cases are acquired

Aggenesis means that the esophagus is not developed at all.

# Atresia

- Thin, non-canalized cord replaces a segment of esophagus.

This will interfere with the swallowing process and cause mechanical obstruction.

- Most common location: at or near the tracheal bifurcation.

- +- (sometimes associated with) fistula (upper or lower esophageal pouches to a bronchus or trachea).

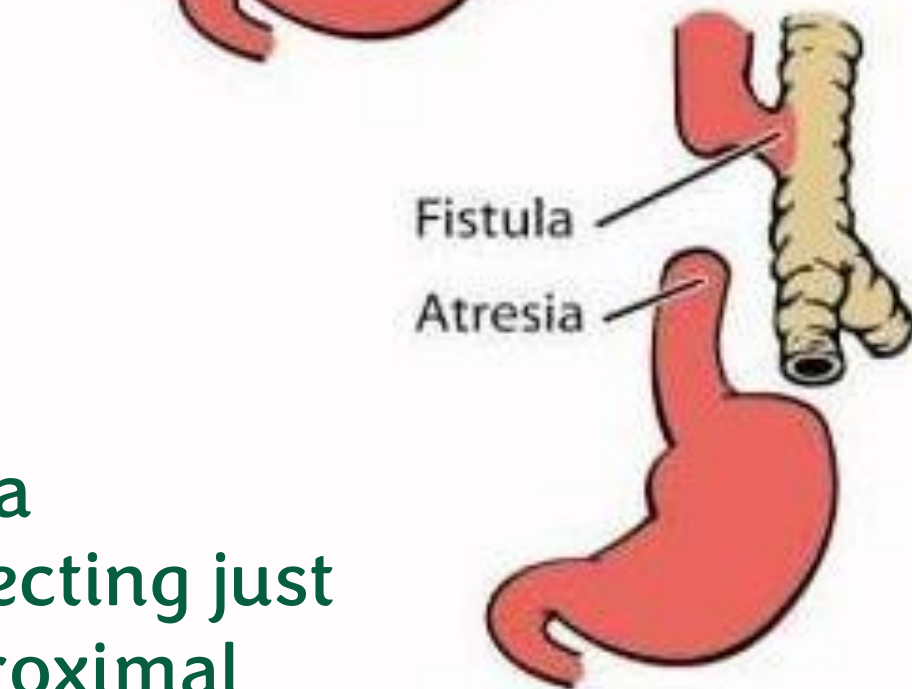
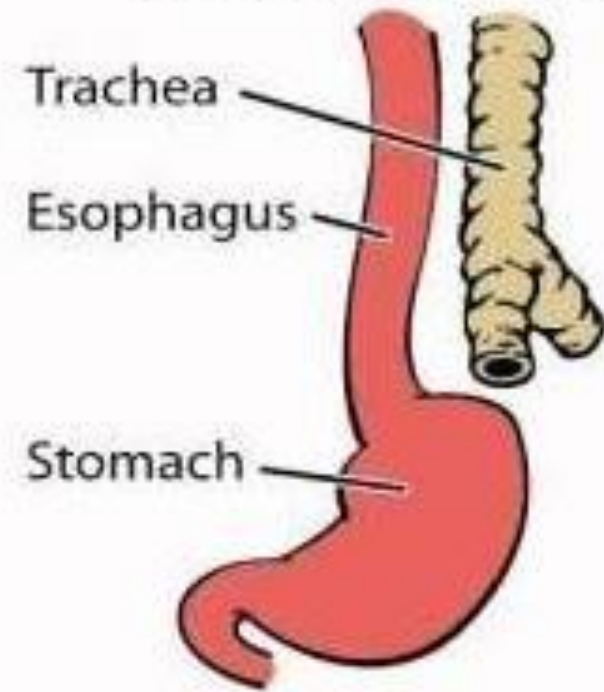
This association connects the upper part or the lower part of the esophagus with bronchus or trachea, with consequent risk of aspiration and aspiration pneumonia.

Aspiration is when something you swallow enters your airway instead of the esophagus.

- 1-Proximal pouch.
- 2-the white color here means this part non-canalized.
- 3-Distal pouch (connected by fistula to the trachea).

Full esophagus separated from the trachea and the bifurcation of the trachea into two main-bronchi

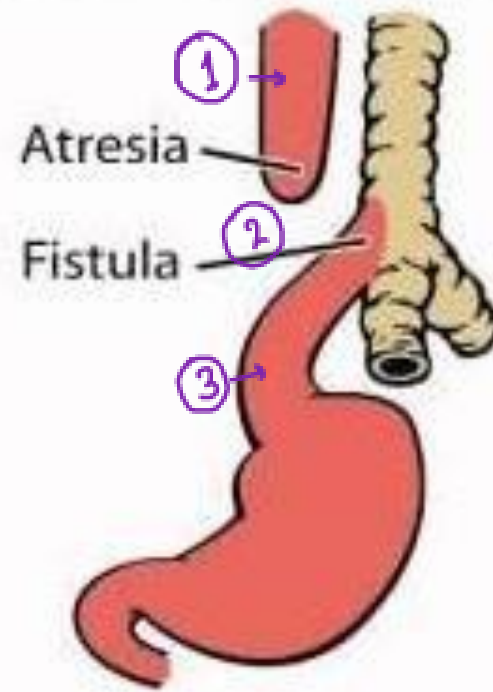
Normal Anatomy



Fistula connecting just the proximal pouch to the trachea

Atresia with proximal Fistula

Atresia with distal Fistula

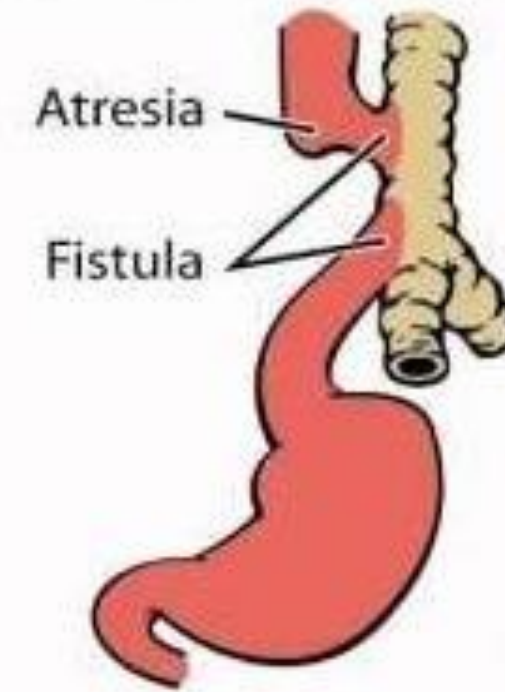


Atresia without fistula

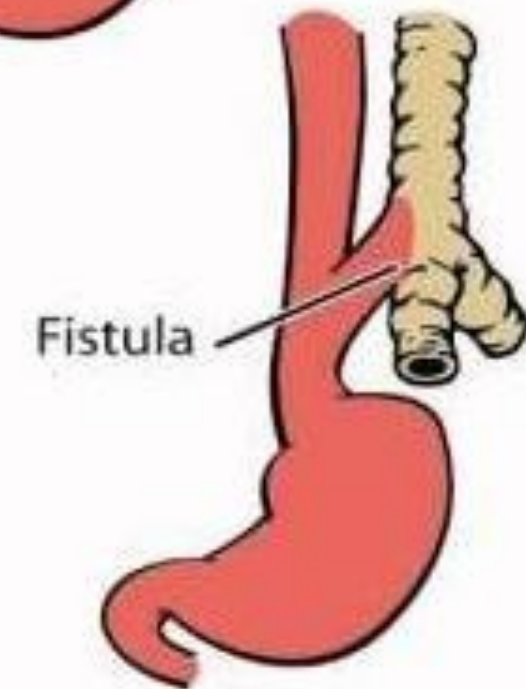


Atresia

Atresia with double Fistula



Both proximal and distal pouches are connected by fistulas to the trachea



Fistula without atresia

Fistula

# Clinical presentation:

- Shortly after birth: regurgitation **of food** during feeding  
**This is non-compatible with life.**
- Needs prompt surgical correction (rejoin).  
**in order for the baby to be able to eat and swallow.**  
**with**
- **Complications if w/ fistula:**
- Aspiration **of food contents**
- Suffocation **اختناق**
- Pneumonia
- Severe fluid and electrolyte imbalances.

**Because of the inability to eat, and nutritional problems.**

# Esophageal stenosis

## Most of the cases

- Acquired>>>Congenital.
- Characterized by: Fibrous thickening of the submucosa & atrophy of the muscularis propria. This will cause impedance of food flow through the esophagus.
- Due to inflammation and scarring upon a previous injury
- Causes:
  - Chronic GERD. GERD can be associated with ulcerations that are repaired by fibrosis leading to **stenosis** and narrowing of the esophagus.
  - Systemic sclerosis. Systemic sclerosis is due to fibrosis of the submucosa.
  - Irradiation therapy
  - Ingestion of caustic agents المواد الحارقة

Acids & Alkaline: they can cause chemical esophagitis which can be complicated later on by fibrosis and stenosis.

**GERD:**  
Gastroesophageal  
Reflux Disease

# Clinical presentation

- Progressive dysphagia.
- Difficulty eating solids that progresses to problems with liquids.

# 2-Functional Obstruction

: when you don't see something that interferes with the passage of food, but there is an abnormality in the innervation.

Efficient delivery of food and fluids to the stomach requires coordinated waves of peristaltic contractions.

Esophageal dysmotility: disorganized peristalsis or spasm of the muscularis.

Achalasia: the most important cause.

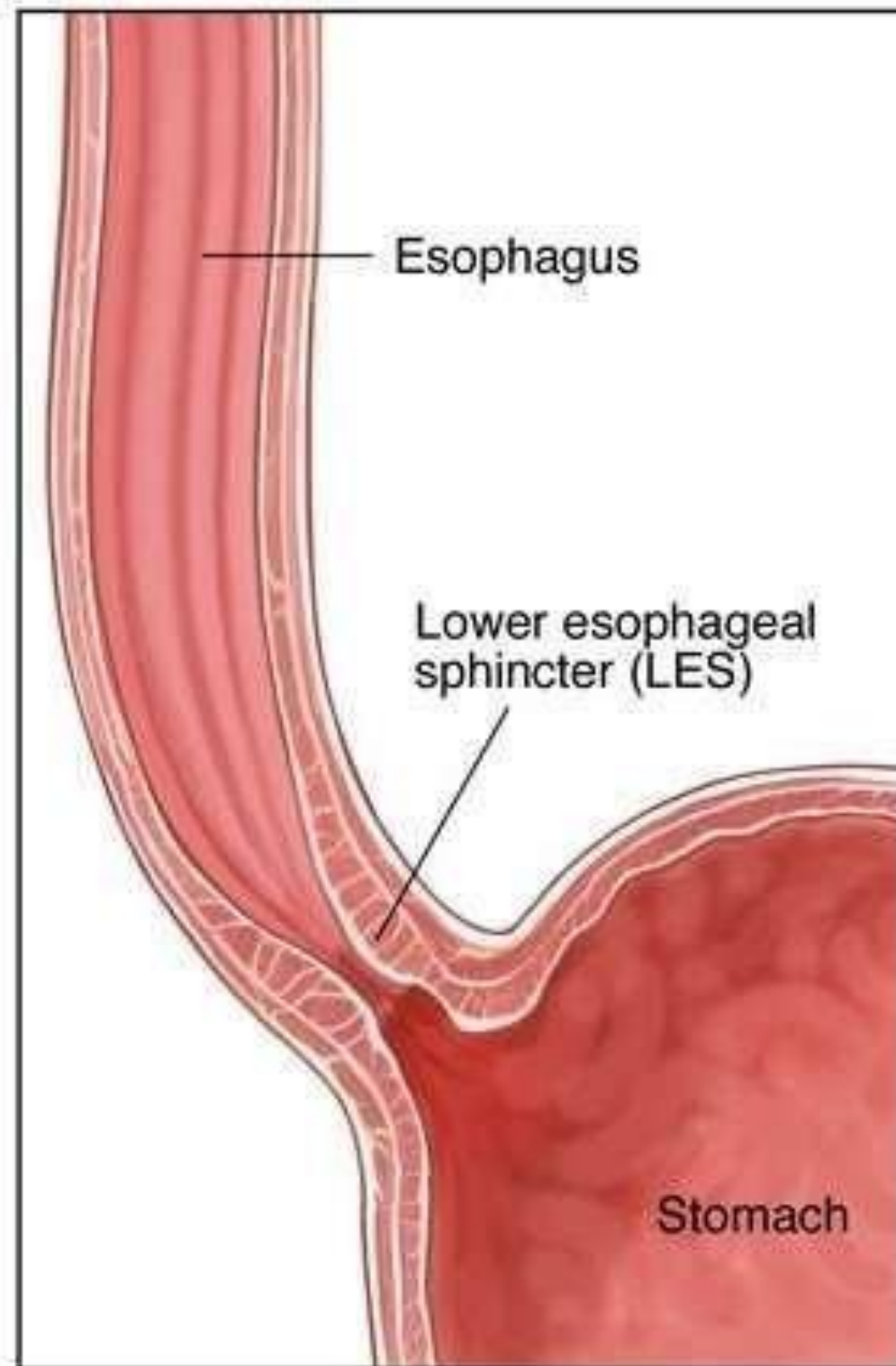
And this will lead to inefficient delivery of food through the esophagus to the stomach

# Achalasia

- **Triad:**
- Incomplete LES relaxation
- Increased LES tone  
→ no complete relaxation → the sphincter will be semi-closed.
- Esophageal aperistalsis. = No peristaltic movement.
- Primary >>>secondary.  
more common

LES: lower esophageal sphincter

## Typical features of Achalasia



Normal



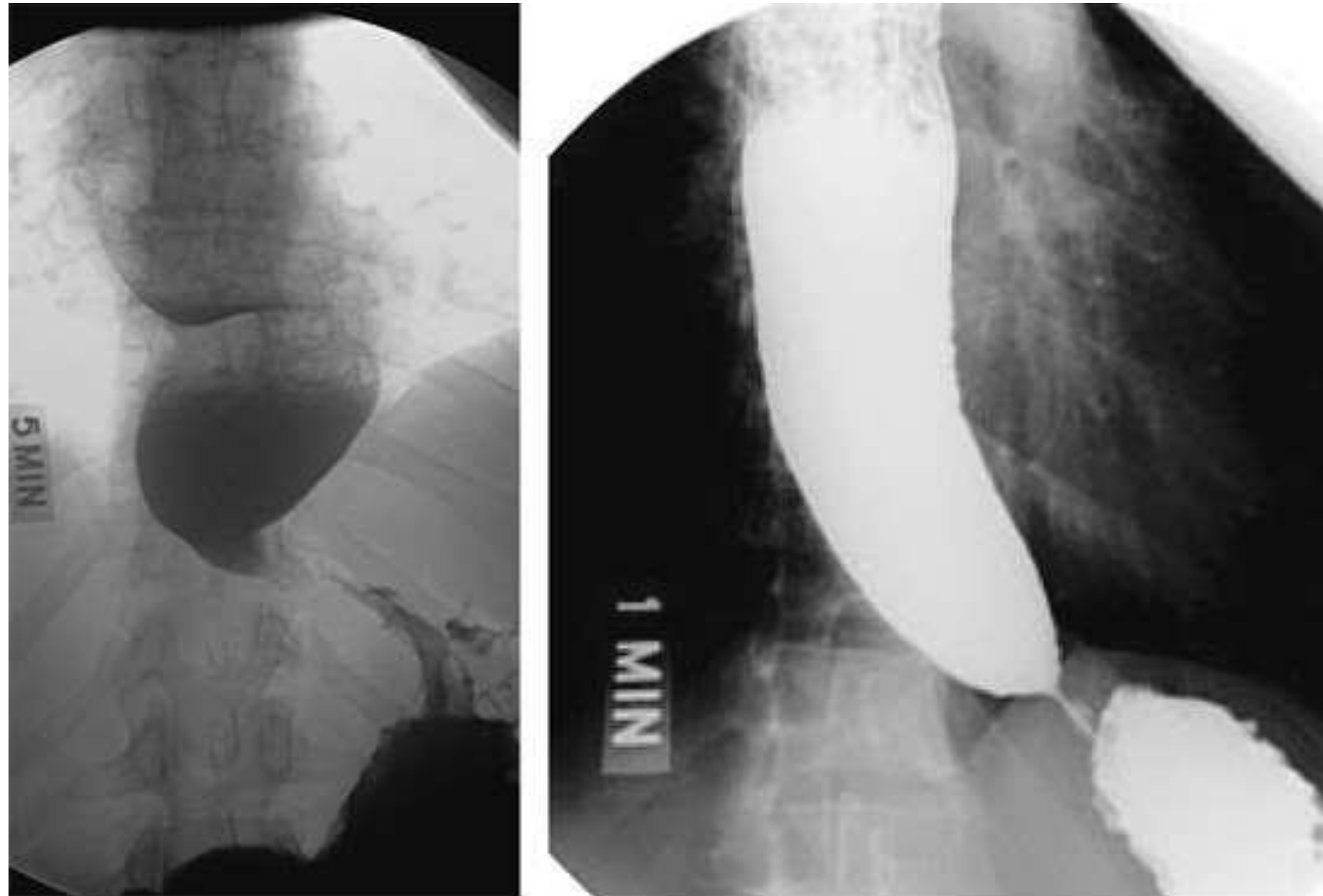
Achalasia

- Increased tone of LES and incomplete relaxation: Sphincter will be semi-closed (won't be open & won't be fully relaxed upon arrival of food) causing accumulation of food in the esophagus & the esophagus will dilate.

- The sphincter is semi-closed, while esophagus is dilated (due to accumulation of food).

## Barium swallow test:

We ask the patient to drink barium then we take X-ray images.



We can see that:

-the barium here will build up in the esophagus which is dilated.

-LES appears as a string as it is semi-closed, with a passage of very small amount of food to the stomach.

Source: Longo DL, Fauci AS, Kasper DL, Hauser SL, Jameson JL, Loscalzo J: *Harrison's Principles of Internal Medicine, 18th Edition*: [www.accessmedicine.com](http://www.accessmedicine.com)

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# Primary achalasia



-These neurons are responsible for the inhibition of the contraction leading to dilatation or relaxation of muscles.

-The degeneration of these neurons leads to increase tone of LES.

# Secondary achalasia

Less common

- Loss of neural innervation due to damage in:
- **Esophagus.**
- **Vagus nerve** (Which innervates the esophagus)
- **Dorsal motor nucleus of vagus**
- **One of the causes : Chagas disease, Trypanosoma cruzi infection**>>destruction of the myenteric plexus>> failure of LES relaxation>> esophageal dilatation.

Myenteric plexus is plexus of the gut which is responsible for the peristaltic movement.

# Clinical presentation

- Difficulty in swallowing
- Regurgitation **of food**
- Sometimes chest pain. **Due to aspiration.**

# 3-Vascular diseases: Esophageal Varices دوالي المريء

It is the most important vascular disease of the esophagus.

- Tortuous dilated veins within the submucosa of the distal esophagus and proximal stomach.
- Diagnosis by endoscopy or angiography.



These are dilated submucosal veins (the blackish vessels) engorged with blood in the distal esophagus.

**Pathogenesis:**  
Usually due to  
portal hypertension

- **Portal circulation:** blood from GIT >> portal vein >> liver (detoxification) >> (via hepatic vein) inferior vena cava.
- Diseases that impede portal blood flow >> portal hypertension >> esophageal varices.
- Distal esophagus : site of Porto-systemic anastomosis.
- **Portal hypertension** >> collateral channels in distal esophagus >> shunt of blood from portal to systemic circulation >> dilated collaterals in distal esophagus >> varices

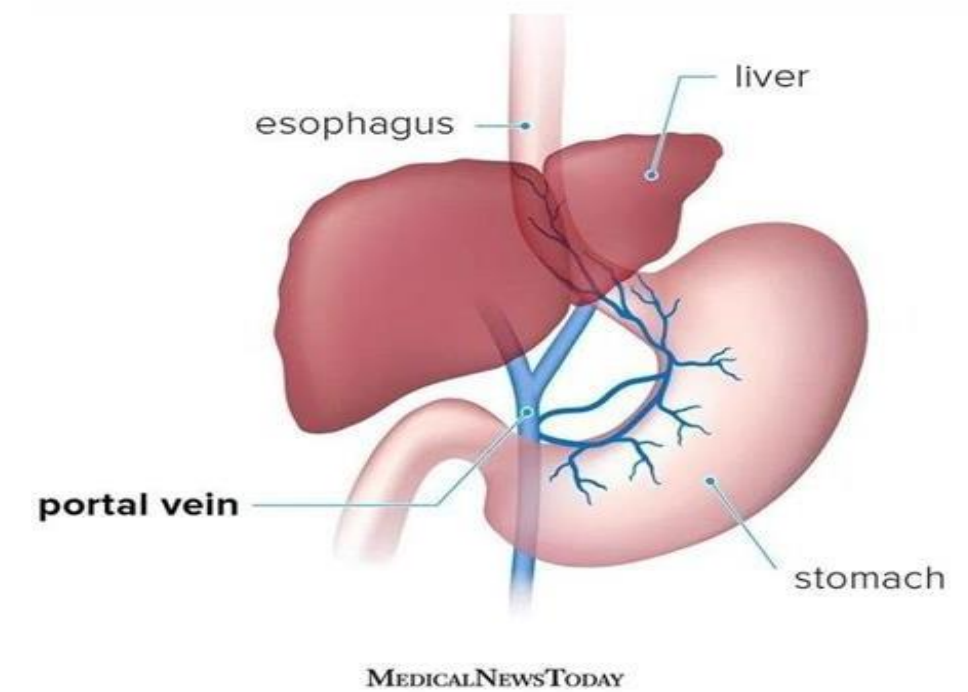
# Additional figure

-GI system is characterized by the presence of portal circulation.

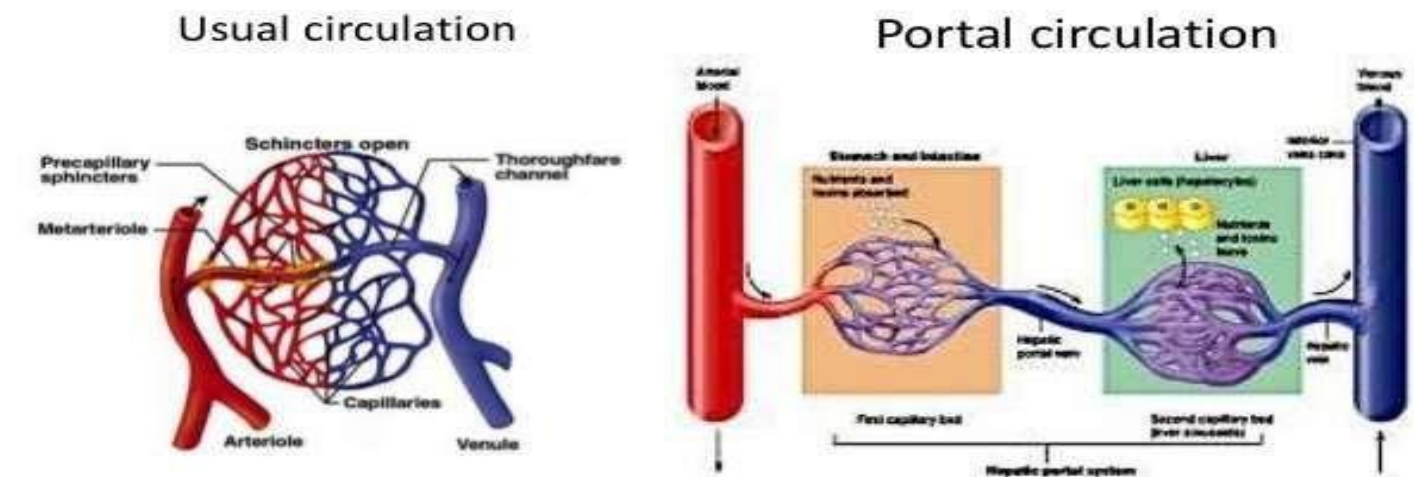
-What is a Portal Circulation?

It's a process in which the blood that is collected from GI tract will not go directly to the inferior vena cava, instead it will go through the portal vein to the liver (for detoxification), then through the hepatic vein into inferior vena cava.

-Any disease that impede this portal blood flow will lead to portal hypertension, leading to shunt of blood from the portal circulation to the systemic circulation through the area in which we have a collateral anastomosis between these two circulation, and one of these sites is **distal esophagus proximal stomach** leading to **Esophageal Varices**.

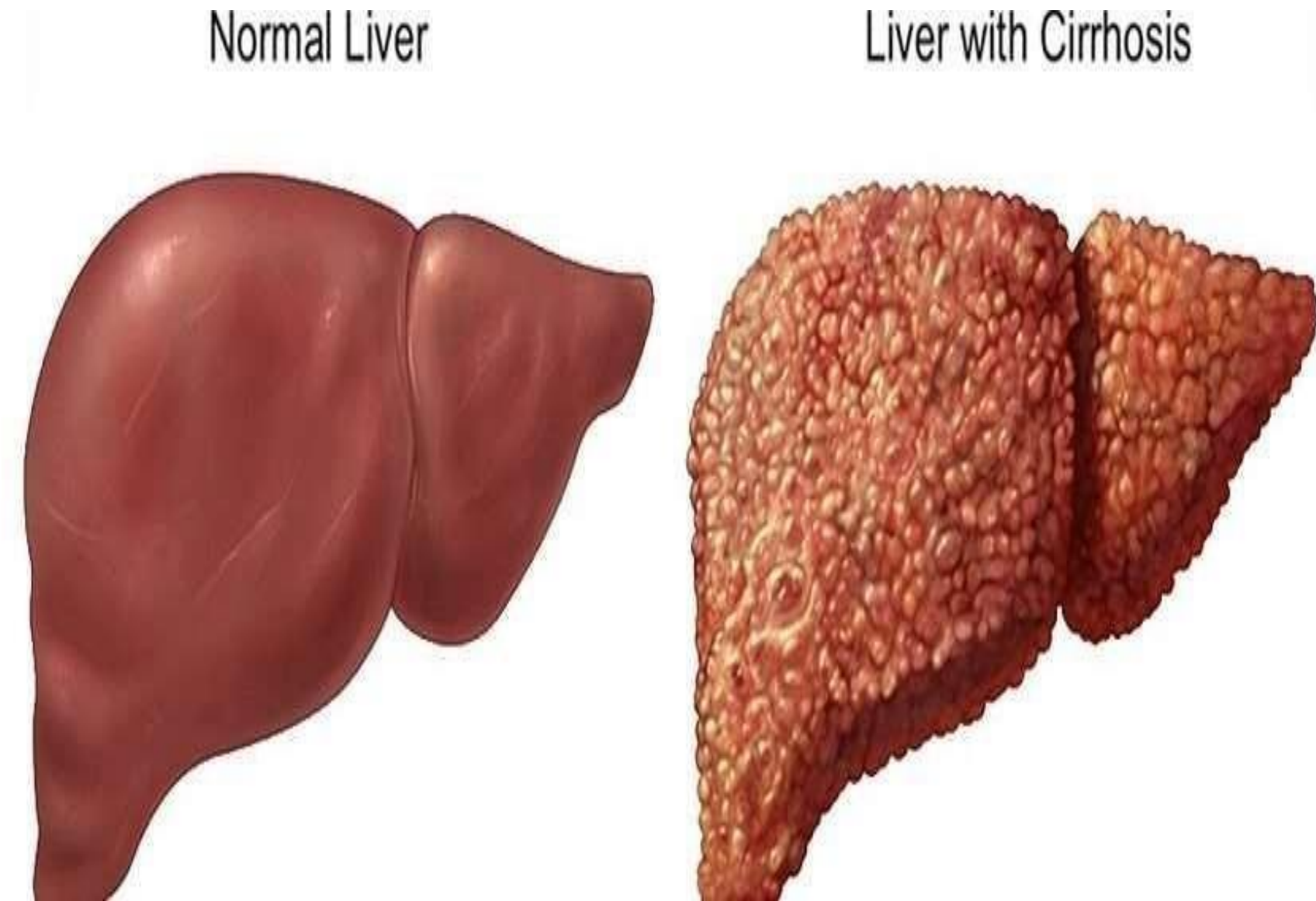


## Portal system



# Causes of portal hypertension

- ▶ Cirrhosis is most common  
Alcoholic liver disease.
- ▶ Hepatic schistosomiasis 2 **(second)** most common worldwide.



**Cirrhosis**



**liver will transform into nodular liver**



**portal Hypertension & Chronic liver disease**

# Clinical Features

- Often asymptomatic.

- Rupture leads to massive hematemesis and death.

- 20% of patients die from the first bleed despite interventions.

- Death due to hemorrhage, hepatic coma, and hypovolemic shock

- Rebleeding in 60%.

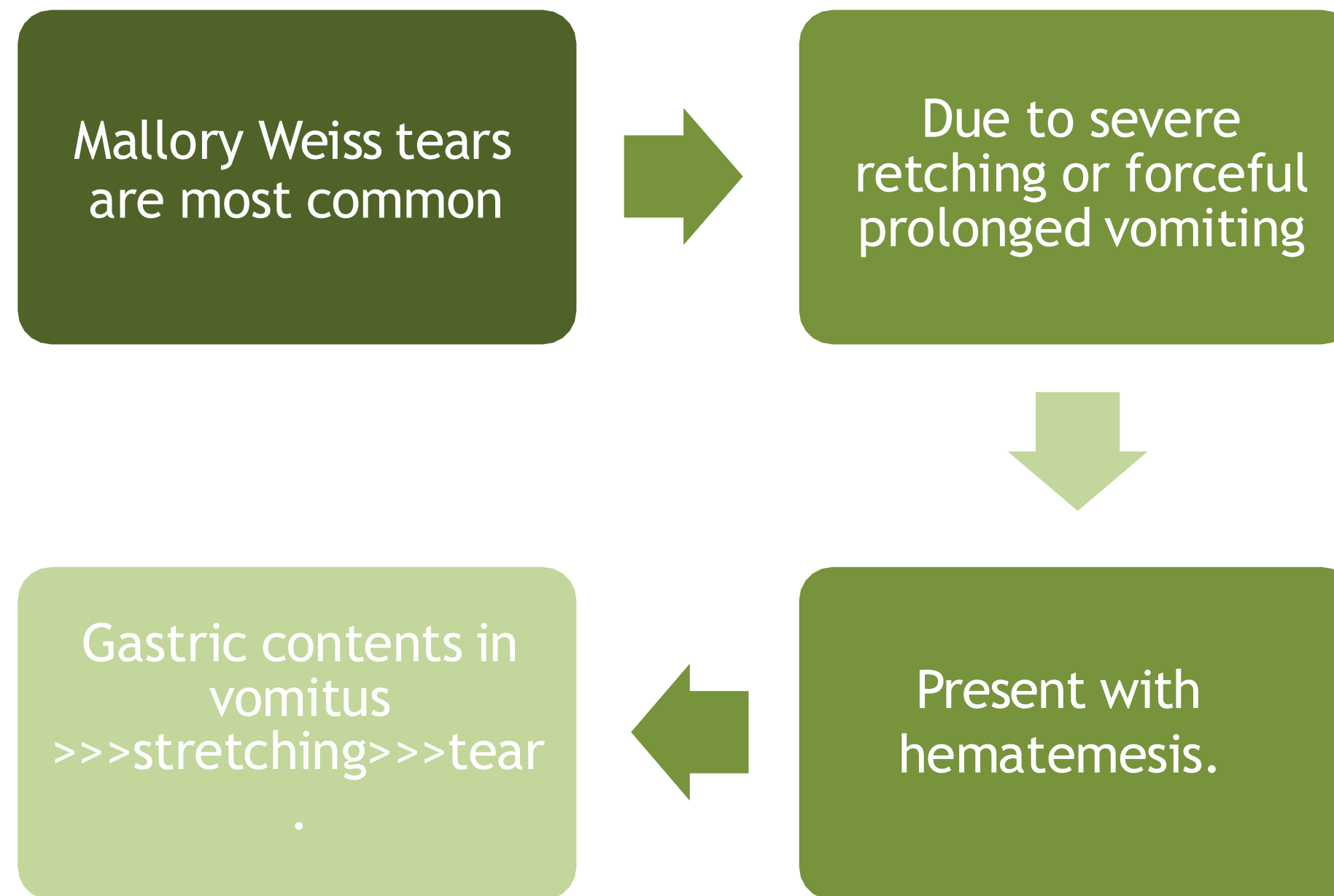
Discovered incidentally during endoscopy in patients with cirrhosis because they should undergo periodic surveillance for the development of varices .  
Hematemesis= Vomiting of blood

# 4-ESOPHAGITIS

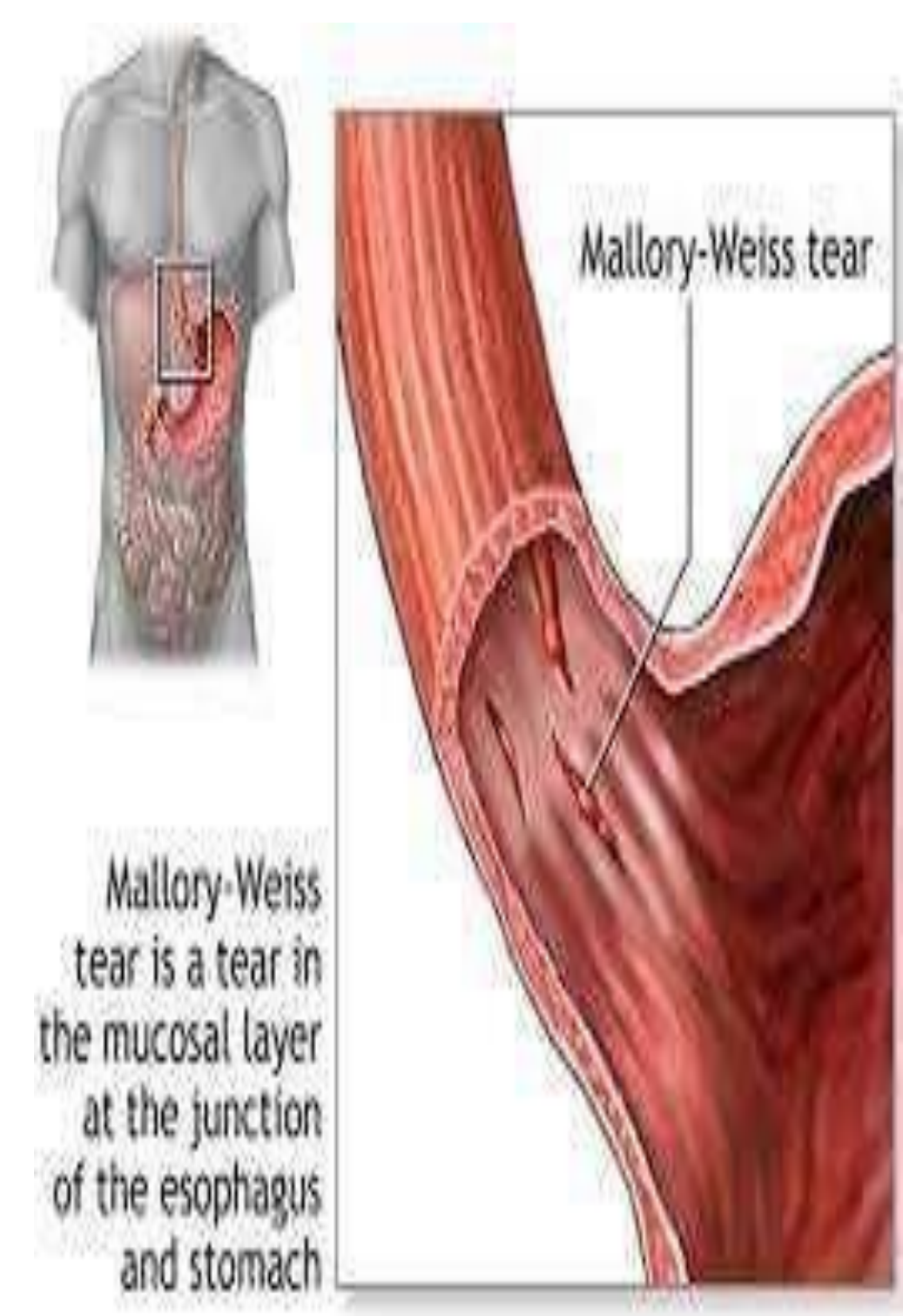
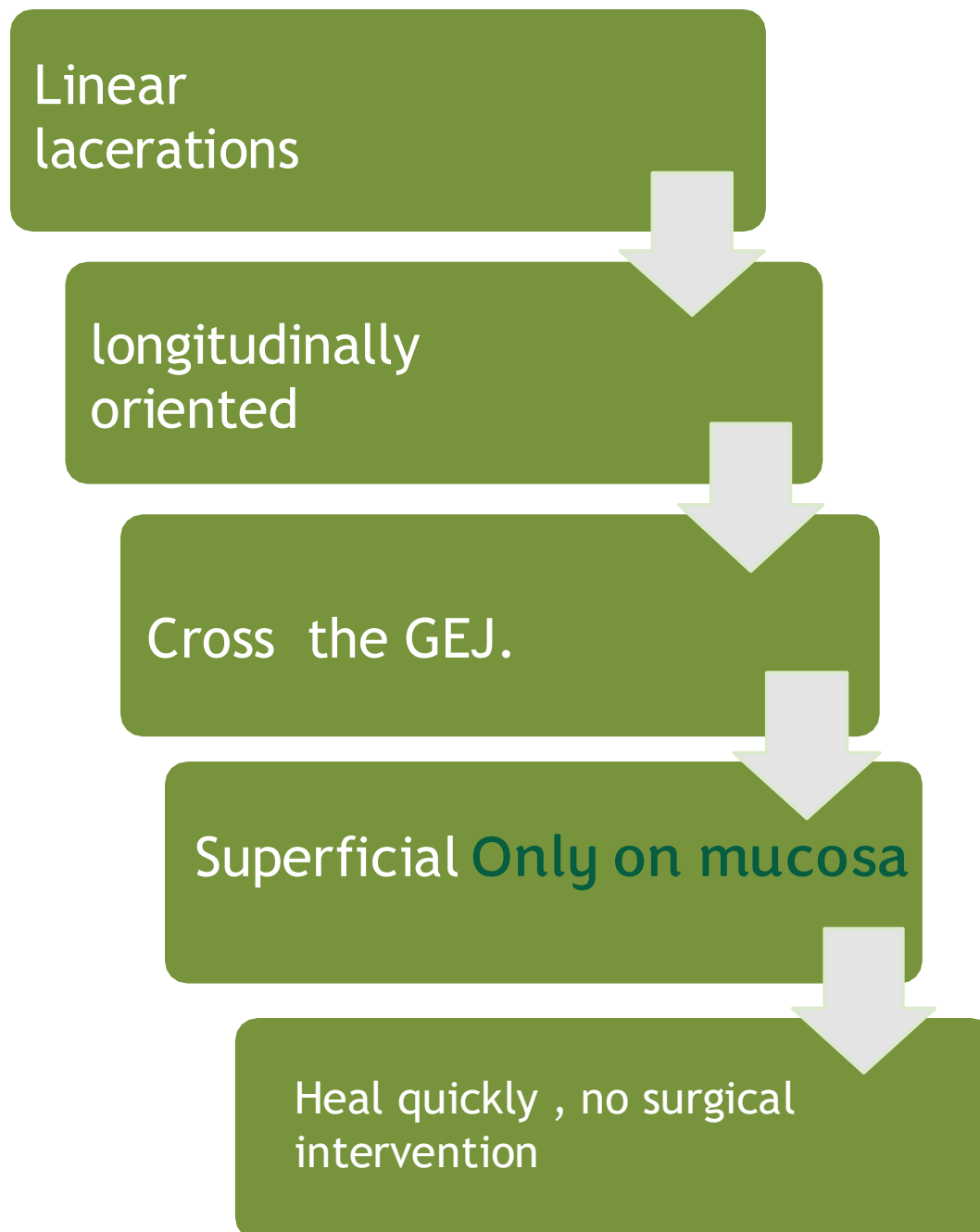
Inflammation of esophageal Caused  
by:

- ▶ Esophageal Lacerations.
- ▶ Mucosal Injury
- ▶ Infections
- ▶ Reflux Esophagitis
- ▶ Eosinophilic Esophagitis

# Esophageal Lacerations



-During vomiting which is severe & prolonged in this case, there will be no time for esophagus to relax. Consequently, a large amount of gastric content will pass through esophagus causing distension & stretching of esophagus. This leads to a tear presented with hematemesis (Fresh red colored blood)



The patient is presented with fresh red colored blood after forceful vomiting -> esophagitis  
The physician should reassure the patient that will not happen again and the bleeding or the tears will heal spontaneously

# Chemical Esophagitis

- ▶ **Damage to esophageal mucosa by irritants**
- ▶ Alcohol,
- ▶ Corrosive acids or alkalis
- ▶ Excessively hot fluids
- ▶ Heavy smoking
- ▶ Medicinal pills (doxycycline and bisphosphonates)
- ▶ Iatrogenic (chemotx, radiotx , GVHD)

**GVHD: Graft Versus Host Disease**

- **Biphosphonates are a major cause of medicine pill esophagitis due to large size of tablets that could be stuck in esophagus-  
Solution: We ask the patient to drink plenty of water & stay in an upright position for a while**

# Clinical symptoms & morphology

- ▶ Ulceration and acute inflammation.
- ▶ Only self-limited pain, odynophagia (pain with swallowing).
- ▶ Hemorrhage, stricture, or perforation in severe cases     **Stricture can lead to stenosis**

# Infectious esophagitis

- ▶ Can affect normal individuals, but is Mostly seen in debilitated or immunosuppressed patients.
- ▶ Viral (HSV, CMV)
- ▶ Fungal (candida >>> mucormycosis & aspergillosis)
- ▶ Bacterial: 10%. Bacteria is less common, and can be secondary to viral or fungal infection.

## Candidiasis :

- ▶ • Adherent.
  - Adherent to the esophageal mucosa as seen during endoscopy
- ▶ • Gray-white pseudo membranes
- ▶ • Composed of matted fungal hyphae and inflammatory cells
- ▶ • They can be seen microscopically upon biopsy examination-
- This infection can extend to oral mucosa causing oral thrush

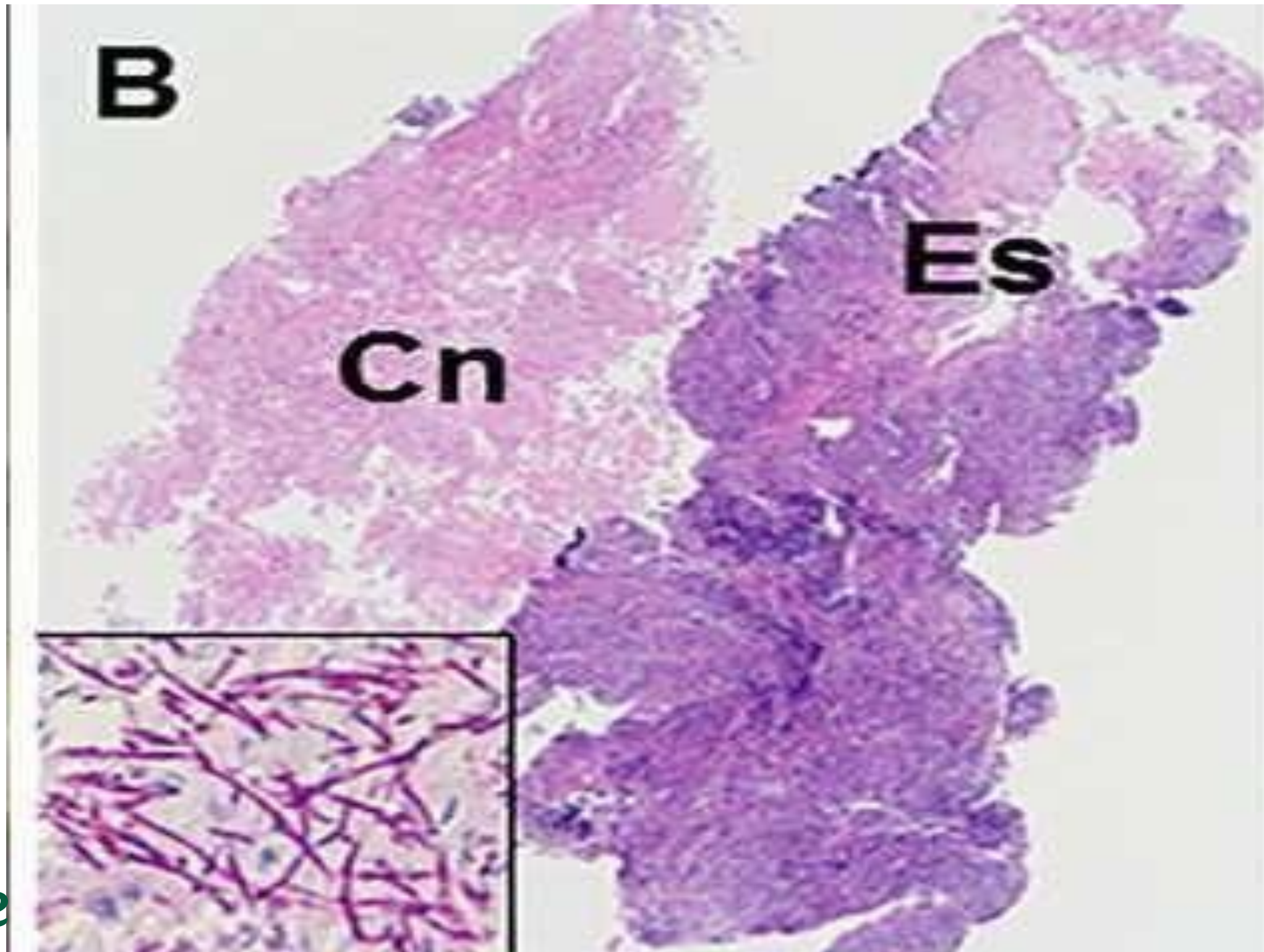
## Esophagus

-Oral mucosa with an oral thrush



<https://www.pinterest.com/pin/374291419013418659/>

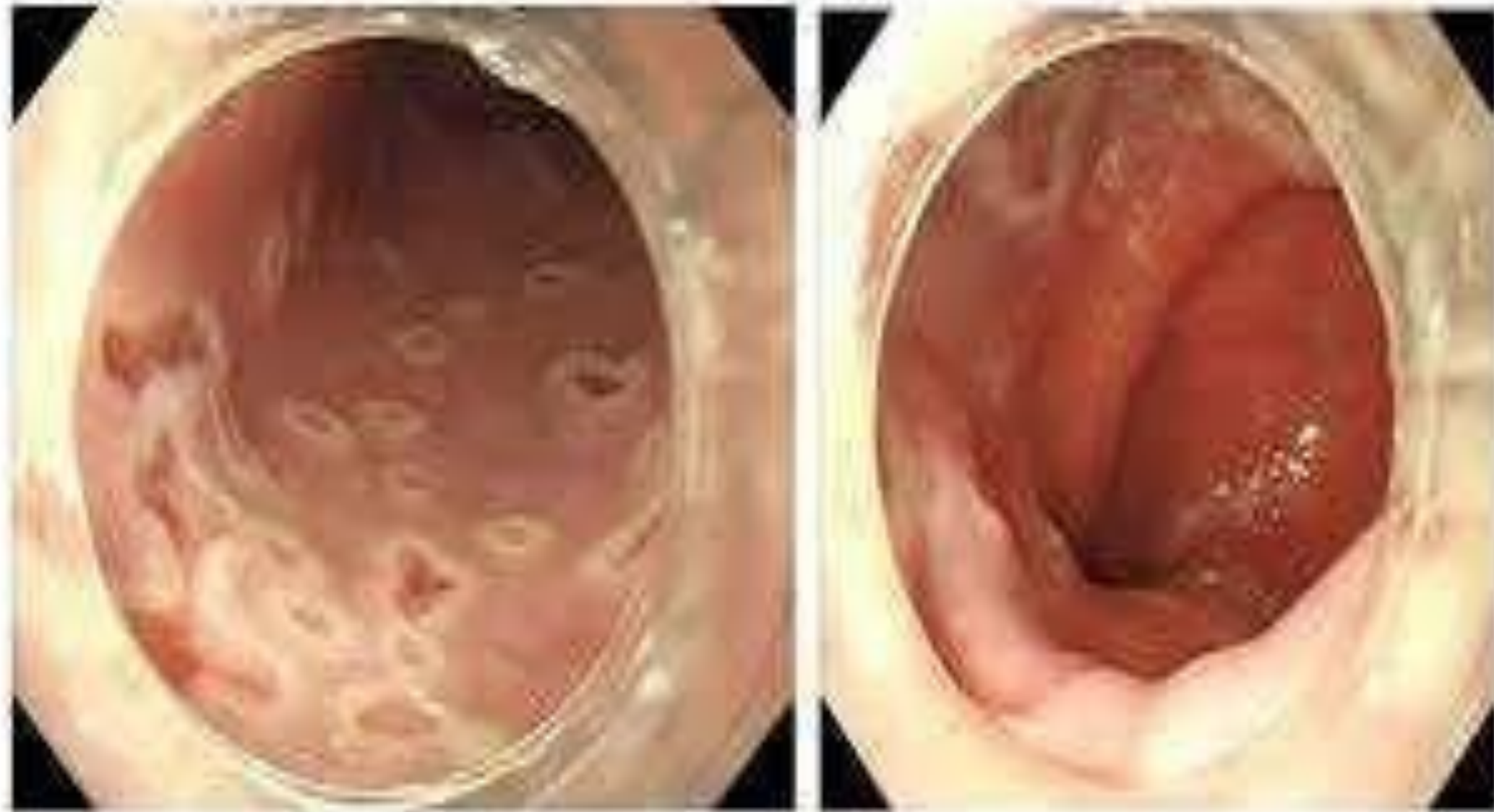
-Hyphae



We use periodic acid schiff stain  
to highlight fungal hyphae

[www.researchgate.net/publication/285369734\\_Esophageal\\_Candidiasis\\_as\\_the\\_Initial\\_Manifestation\\_of\\_Acute\\_Myeloid\\_Leukemia](http://www.researchgate.net/publication/285369734_Esophageal_Candidiasis_as_the_Initial_Manifestation_of_Acute_Myeloid_Leukemia)

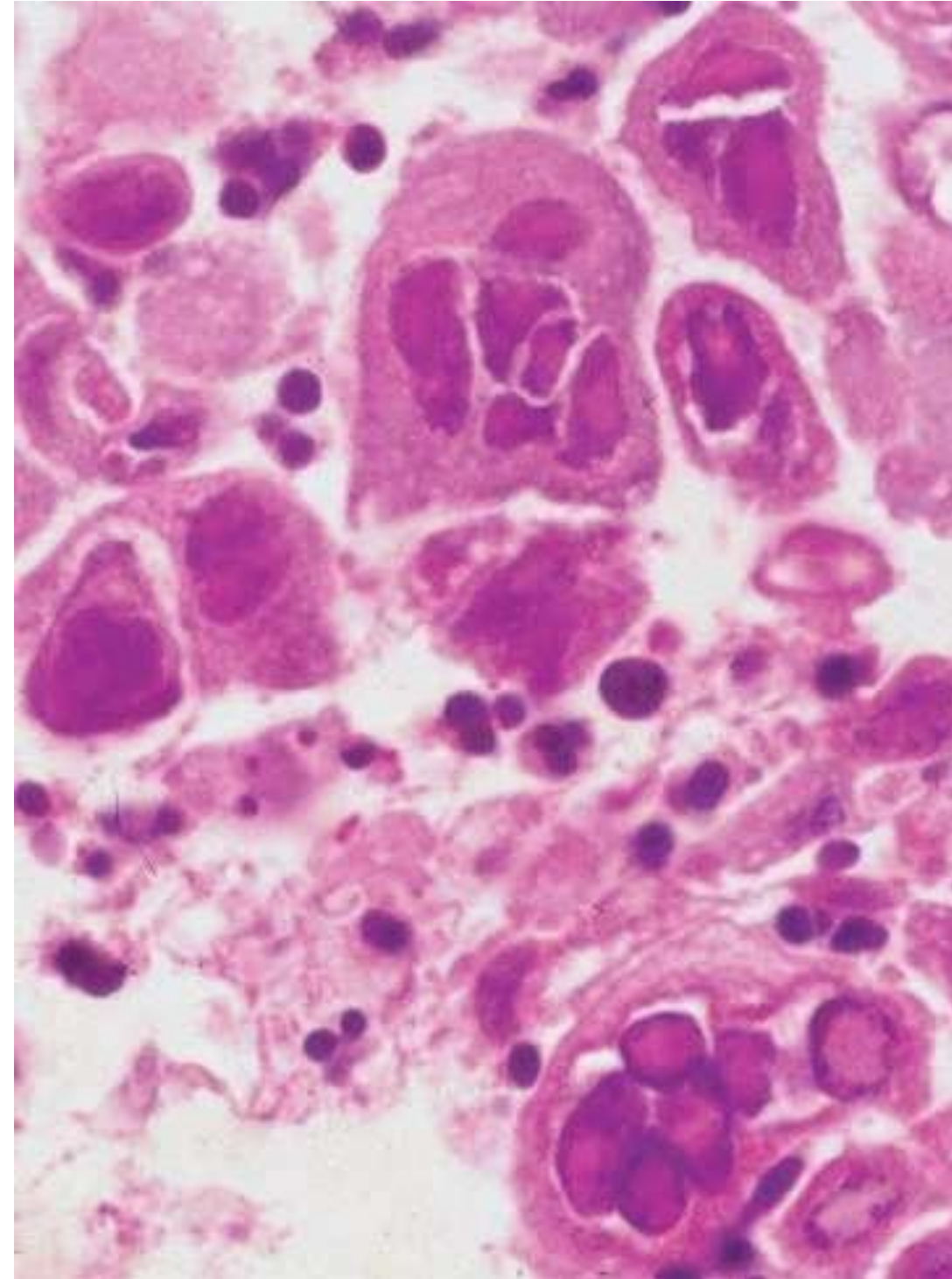
- ▶ **Herpes viruses**
- ▶ Punched-out ulcers **Can be seen by endoscopy**
- ▶ Histopathologic:
- ▶ Nuclear viral inclusions
- ▶ Degenerating epithelial cells ulcer edge
- ▶ Multinucleated epithelial cells (**fusion of epithelial cells to form one cell with multiple nuclei**).



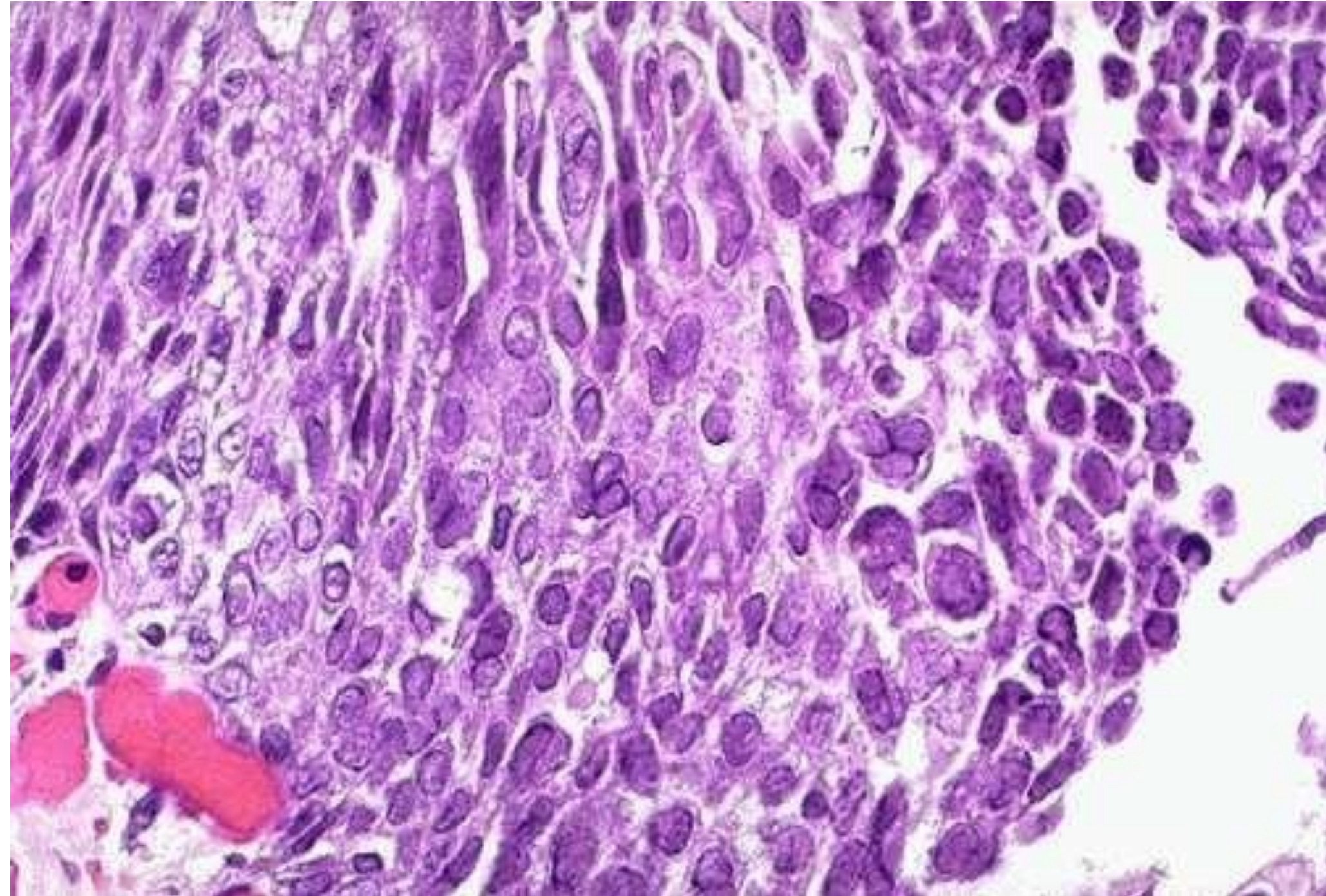
-Remember: The esophagus mucosa is normally pale-pink, but here we see the surrounding mucosa erythematous (red)

Here we can see punched-out ulcers

- HSV infection histology:  
Multinucleated cells, Viral  
nuclear inclusion,  
Degeneration of cells



- **Histology:-Multinucleated cells-The nuclei are characterized by intranuclear inclusion (A typical feature of HSV biopsies)**



- ▶ **CMV :**
- ▶ Shallower ulcerations.
- ▶ Biopsy: nuclear and cytoplasmic inclusions in capillary endothelium and stromal cells. (Mega cells) **Cytomegaly=Mega cells=Large cells**

CMV infects Endothelial & Stromal Cells in addition to epithelial cells, unlike HSV which only infects epithelial cells

-Large Stromal cells  
with nuclear inclusion



Large Endothelial cell

# رسالة من الفريق العلمي:

أنت عبد، فتأدب!

قال ابن القيم في كتابه الفوائد:  
الرَّبُّ سُبْحَانَهُ لَا يَمْنَعُ عَبْدَهُ الْمُؤْمِنُ شَيْئًا مِنَ الدُّنْيَا، إِلَّا وَيُؤْتِيهِ أَفْضَلَ مِنْهُ، وَأَنْفَعَ لَهُ!  
إِنَّ الطَّبَّيبَ لَا يُعْطِيكَ الدَّوَاءَ الَّذِي تَحِبُّهُ،  
وَلَكِنَّهُ يُعْطِيكَ الدَّوَاءَ الَّذِي يَشْفِيكَ (بِإِذْنِ اللَّهِ)  
وَلِلَّهِ الْمِثْلُ الْأَعْلَى، فَتَأَدَّبْ!  
أنت لا ترى المشهد الا من زاوية واحدة ، هي زاويتك أنت!  
والله يدبر لك الأمر من كل الزوايا!  
كان عمر بن الخطاب يقول: لو كُشِفَتْ حُجُبُ الْغَيْبِ،  
مَا اخْتَارَ الْعَبْدُ لِنَفْسِهِ غَيْرَ مَا اخْتَارَهُ اللَّهُ لَهُ!

[من كتاب رسائل من التابعين]

# For any feedback, scan the code or click on it.



Corrections from previous versions:

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