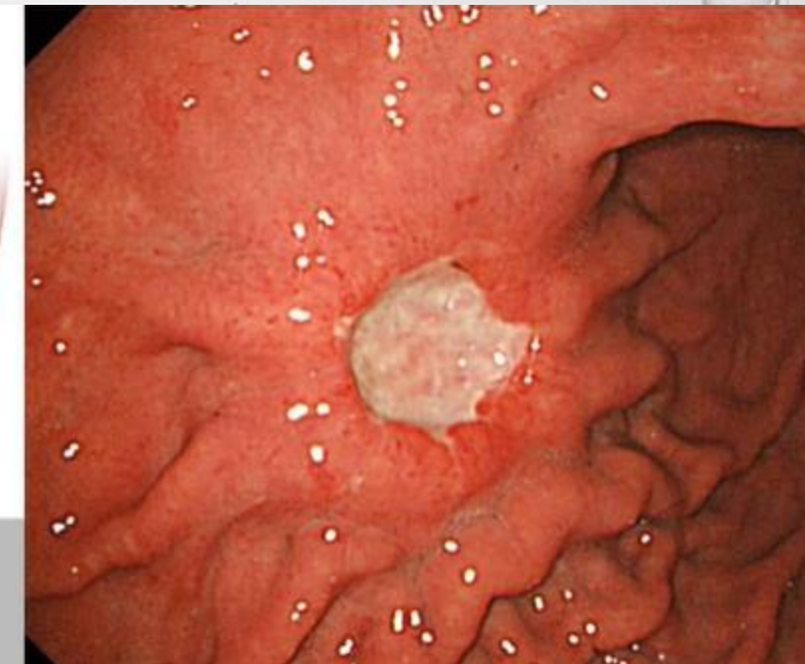
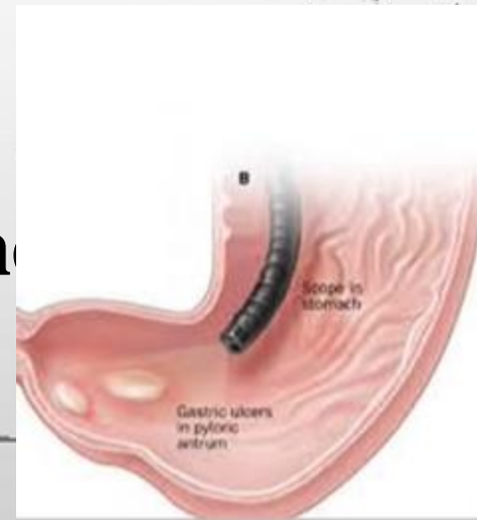
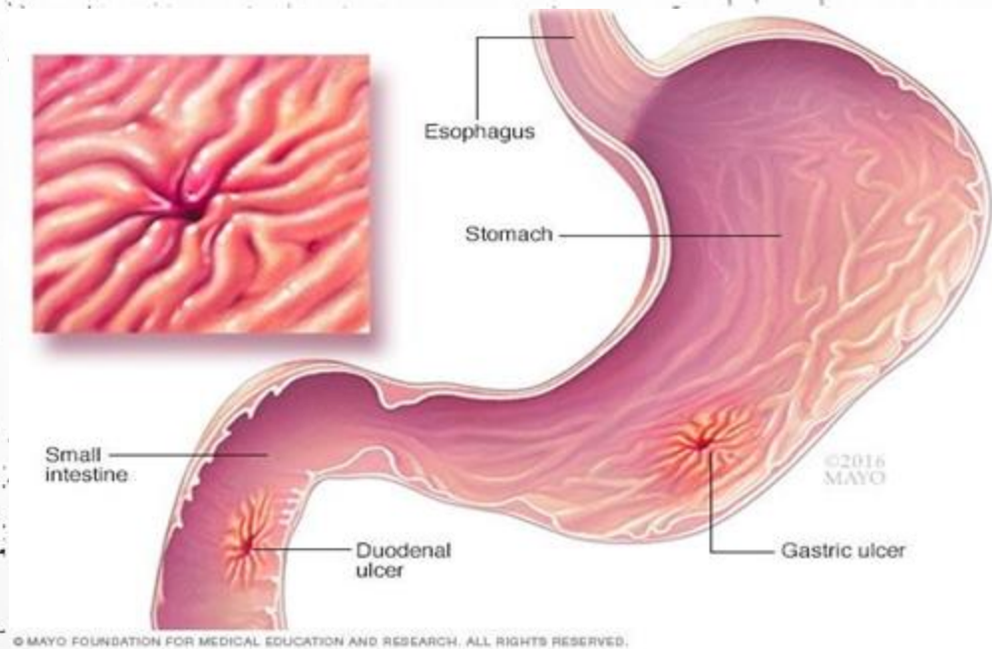


PEPTIC ULCER AND GERD TREATMENT

Dr/ Heba Ahmed Hassan

**Assistant professor of clinical
pharmacology, faculty of medicine
mutah university**



• PATHOGENESIS

- Unbalancing between aggressive factors & defensive factors.

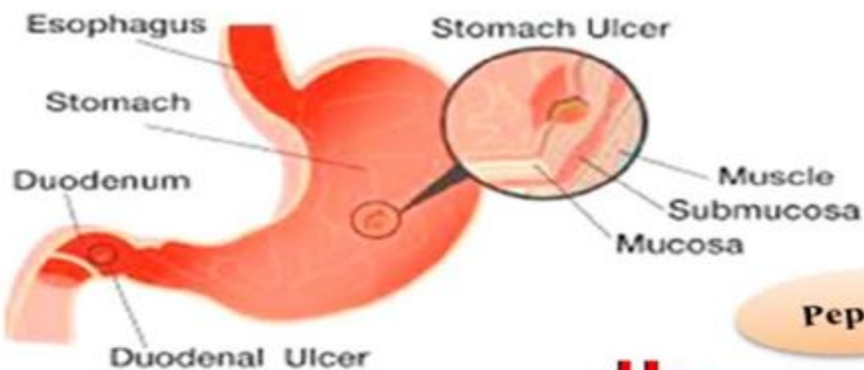
● A. Aggressive factors:

- Gastric acid secretion.
- Pepsin.
- Bile.
- Helicobacter pylori.

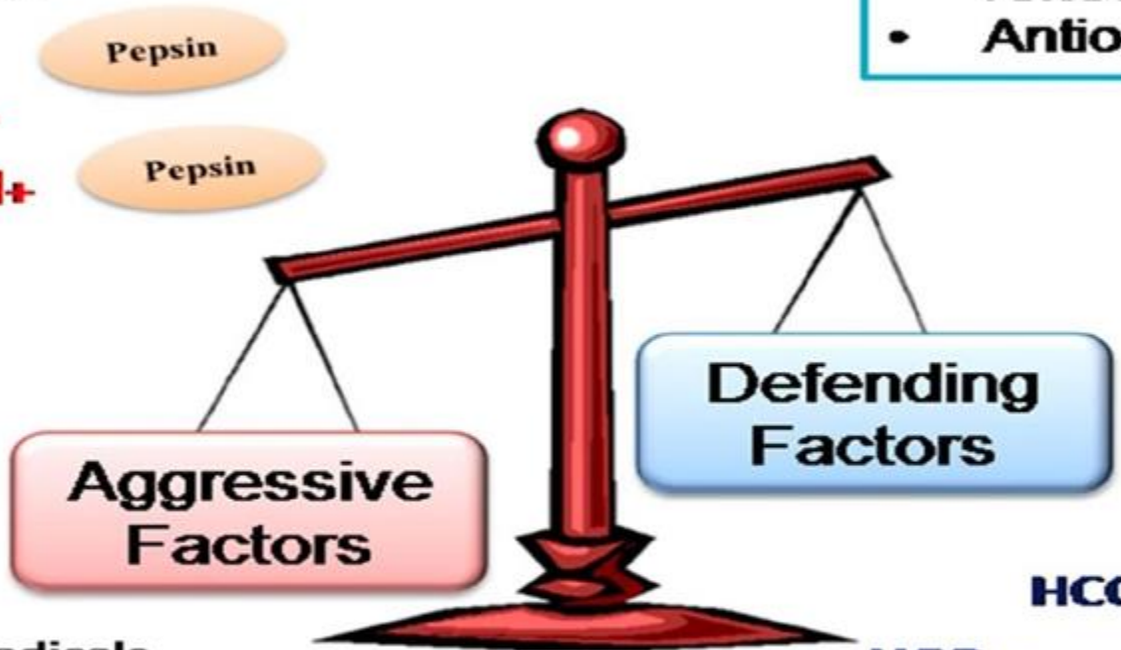
● B. Defensive factors:

1. Mucus & bicarbonate secretion
2. Thick lipoprotein coat.
3. Tight intercellular junctions.
4. Processes of restitution and regeneration after cellular injury.
5. Gastric mucosal blood flow.

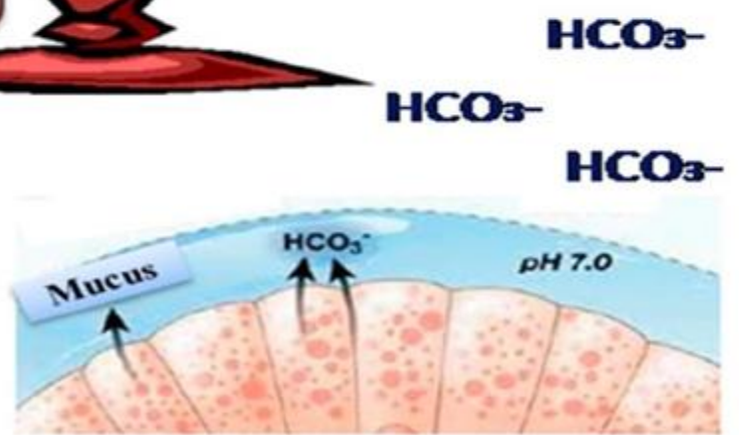
Peptic Ulcers



- Mucus and bicarbonate
- Blood flow
- NO, PGs
- Growth factor and cell renewal
- Antioxidant defense



- *Helicobacter pylori*
- Alcohol intake
- NSAIDs, stress and smoking
- Ischemic processes and RL
- Bile reflux
- Peptic acid secretion

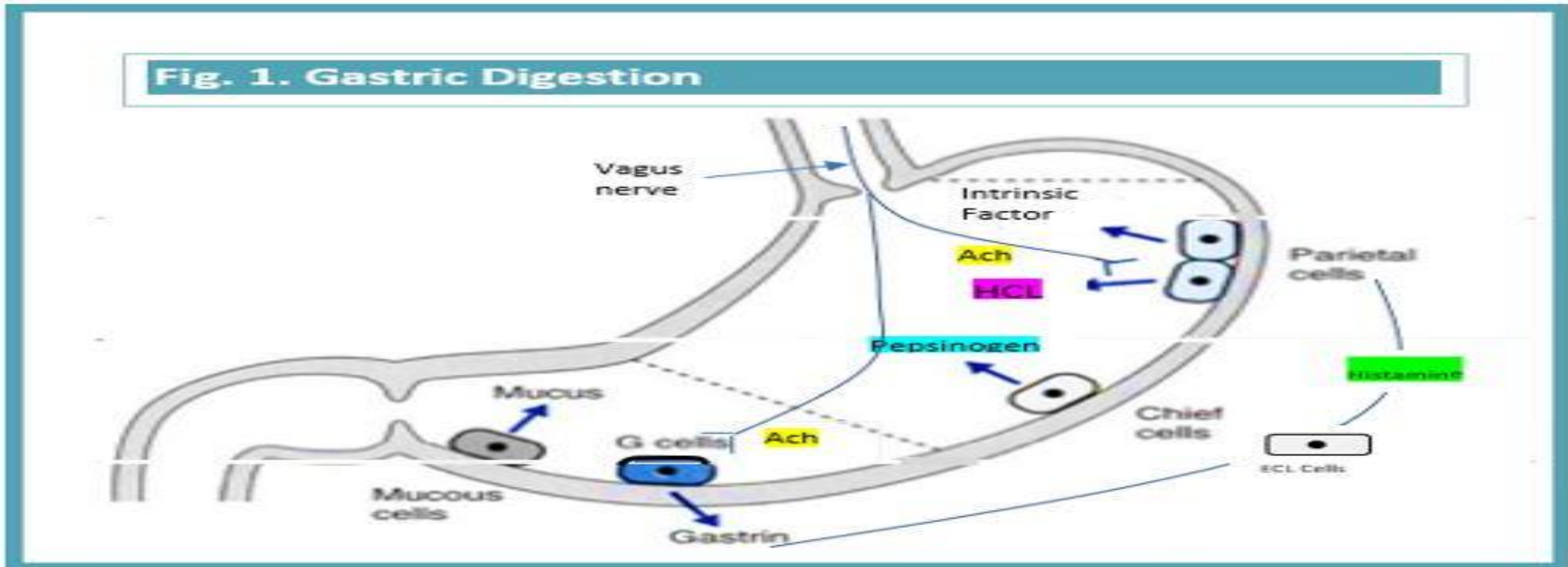


Mucus and bicarbonate



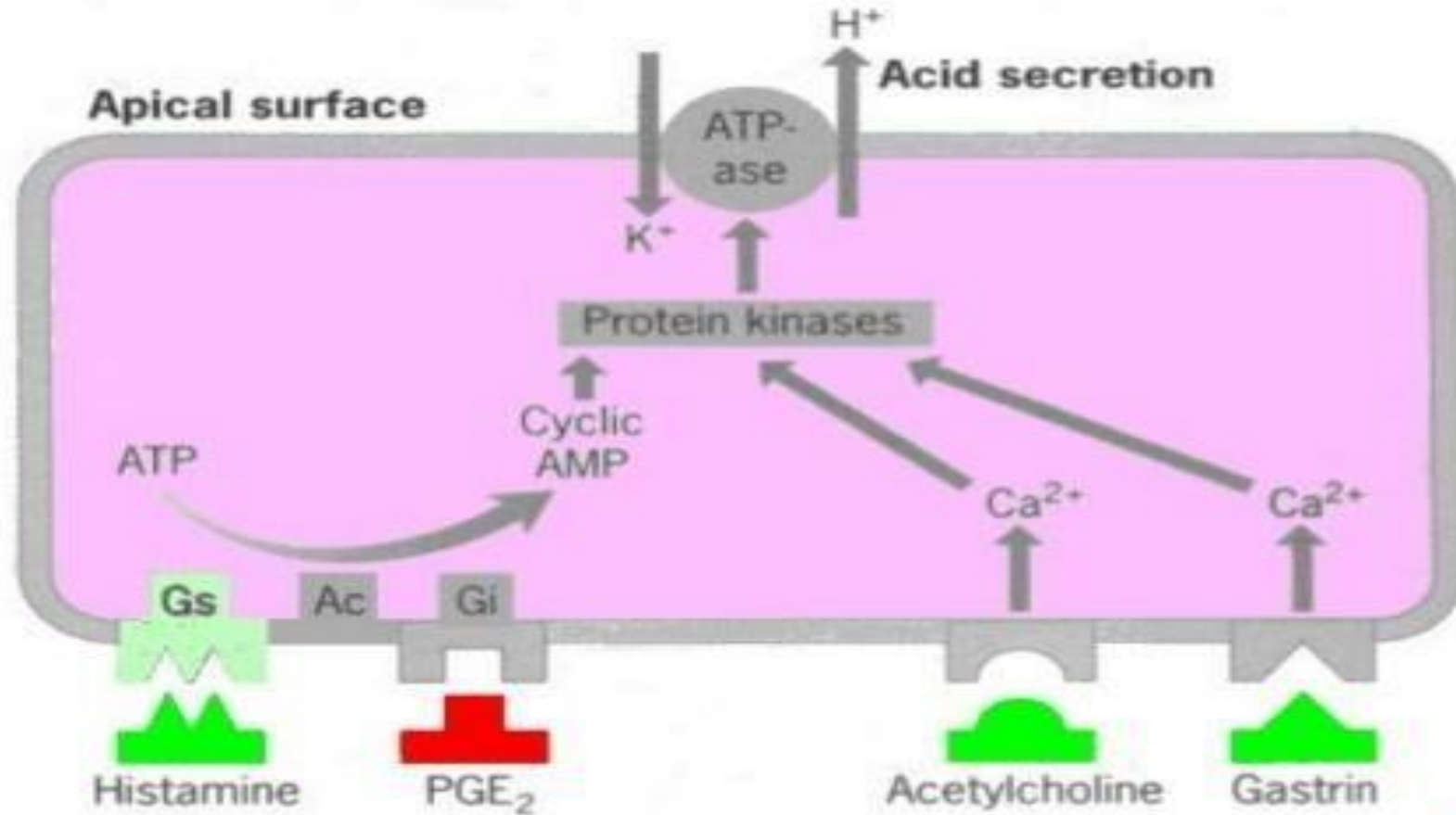
SECRETION OF HCL

- Nocturnal acid secretion (which depends largely on histamine)
- Meal-stimulated acid secretion (which is stimulated by gastrin, Ach and histamine).





Control Of Acid Secretion



HELICOBACTER PYLORI:

- H. pylori is a spiral shaped bacterium that is found in the gastric mucus layer or adherent to the epithelial lining of the stomach.
- 50% of world population is infected. It causes: duodenal/gastric ulcers and gastric cancer.
- H pylori causes more than 90% of duodenal ulcers and more than 60% of gastric ulcers

Clinical pictures:

Symptoms:

- **Pain (duodenal ulcer).**
- **Vomiting (gastric ulcer)**

Complications:

- Hemorrhage.**
- Perforation .**
- cancer (gastric ulcer).**

Goals of therapy:

1. Treatment of symptoms.
2. Promotion of healing (4-8 weeks for D.U. Or 8-16 weeks for G.U.).
3. Prevention of recurrence [maintenance dose (half the normal dose) for at least 6 months].

A -non pharmacological treatment

- ❖ sss (smoking, spices and stress)
- ❖ NSAIDS
- ❖ Drugs and alcohol

TREATMENT OF PEPTIC ULCER

1. drugs that reduce gastric acid secretion:

- a. proton pump inhibitors. PPIs
- b. H₂ histamine receptor antagonists.
- c. muscarinic antagonists .
- d. gastrin antagonists (proglumide).
- e. PG analogue.

2. Neutralization of gastric acidity:

Antacids.

3. Eradication of helicobacter pylori.

4. Cytoprotective agents

A- sucralfate.

B- colloidal bismuth

C- PG analogues (misoprostol).

D- carbenoxolon

DRUGS THAT REDUCE GASTRIC ACID SECRETION

(1) proton-pump inhibitors

- Omeprazole
- esomeprazole
- Lansoprazole
- Rabeprazole
- Pantoprazole

Proton Pump Inhibitor Drugs



PHARMACOKINETICS:

★ **Absorption:** Rapidly absorbed.

The bioavailability is decreased approximately 50% by food, hence drugs should be administered on an empty stomach.

➤ Acid inhibition lasts up to 24 hours owing to the irreversible inactivation of the proton pump.

★ **Distribution:** Bound to plasma protein (95%).

★ **Metabolism:** Hepatic metabolism [CYP3A4 & CYP2C19 (genotype)]. Rapid first-pass & systemic hepatic metabolism.

★ PPIs are administered as inactive prodrugs. To protect the acid-labile prodrug from rapid destruction within the gastric lumen.

Mechanism of action:

- Protonated within the canaliculus (depending on its Pka).
- **Irreversibly** inhibits H⁺-K⁺ ATPase (proton pump). At least **18hrs**. Are required for synthesis of new H⁺/K⁺ ATPase pump molecules.

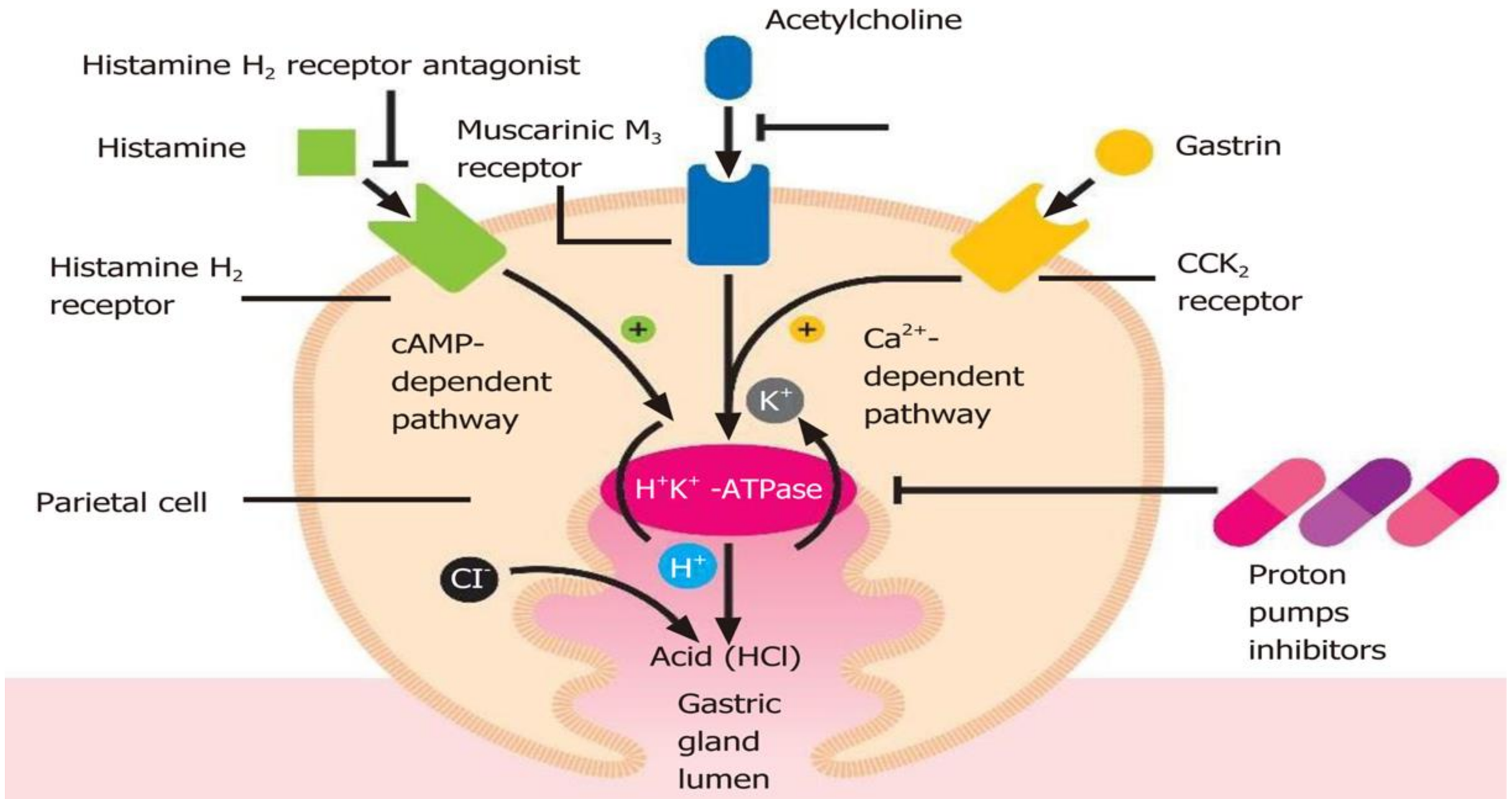
Pharmacological action:

1 inhibit both **fasting & meal-stimulated** gastric acid secretion (more than **95%**).

2 anti-H pylori:

A) direct.

B) ↑PH → ↓ minimal inhibitory concentrations of antibiotics against HP.



USES:

- 1- gastroesophageal reflux disease (GERD).
- 2- peptic ulcer
- 3- Zollinger-Ellison syndrome.
- 4- Prevention of stress-related mucosal bleeding (due to mucosal ischemia have normal or decreased acid secretion):

ADVERSE EFFECTS:

(RARE)

1. G.I.T. (Nausea, diarrhea, colic).

2. C.N.S. (Headache, drowsiness, dizziness).

3. Long-term elevation of gastric PH may cause:

A- hypergastrinemia → ECL hyperplasia which leads to:

□
Carcinoid tumors (rats).

□
Rebound hypersecretion upon discontinuation of the drug.

B-bacterial over growth in G.I.T. → ↑ Risk of respiratory and enteric infections.

4. Skin rash, subacute myopathy & arthralgias.

5. Chronic treatment decreases absorption of B12. (Acid is important in releasing vitamin B12 from food.)

6. Chronic treatment → ↑ risk of hip fracture. (Acid also promotes absorption of food-bound minerals (iron, calcium, zinc))

N.B. Point 5&6 called nutritional adverse effect

Drug interactions:

Because of the short half-lives of PPIs, clinically significant drug interactions are **rare**.

Enzyme **inhibition**: omeprazole may inhibit CYP2C19 (warfarin, phenytoin, and diazepam).

Enzyme **enhancer** Lansoprazole may enhance clearance of theophylline.

Rabeprazole and pantoprazole have no significant drug interactions.

↓ Gastric acidity may alter absorption of drugs for which intragastric acidity affects drug bioavailability, e.g. Ketoconazole, ampicillin ester, iron salts & digoxin.

(2) H2 HISTAMINE RECEPTOR ANTAGONISTS

Cimetidine Ranitidine Famotidine Nizatidine

Pharmacokinetics:

- **Absorption:** Rapidly absorbed.
- **Distribution:** Cross placenta. Therefore they should not be administered to pregnant women (CLASS B). Secreted in breast milk.
- **metabolism:** Cimetidine, ranitidine & famotidine undergo first-pass hepatic metabolism resulting in a bioavailability of approximately 50%
 - Nizatidine** has little first-pass metabolism and a bioavailability of almost 100%
- **Elimination:** H2 antagonists are cleared by a combination of hepatic metabolism, glomerular filtration, and renal tubular secretion (large part excreted by urine).

Pharmacodynamics:

- **Competitively** inhibit the interaction of **histamine with H2 receptors**.
- ↓ Gastric acid secretion.
- H2 antagonists are especially effective at inhibiting nocturnal acid secretion (which depends largely on histamine) but have a modest impact on meal-stimulated acid secretion (which is stimulated by gastrin and acetylcholine as well as histamine). Thus they block more than 90% of nocturnal acid but only 60-80% of day time acid secretion.

Uses:

1. Peptic ulcer.
2. Zollinger-ellison syndrome.
3. Gastro-esophageal reflux disease (GERD).
4. Other conditions (stress ulcer, Preanesthetic medication “emergency”).

Adverse effects

- Diarrhea, headache, fatigue, nausea, myalgia, constipation (common).
- Mental status changes (confusion, hallucination, agitation), commonly with cimetidine (I.V., Elderly, renal or hepatic dysfunction).
- Gynecomastia or impotence in men & galactorrhea in women (anti-androgen, ↑prolactin & estradiol).specific to cimetidine
- Cimetidine inhibits cytochrome P450 hepatic enzymes
- Rapid I.V. Infusion → bradycardia & hypotension through blockade of cardiac H₂ receptors.
- 4. thrombocytopenia
- 5. Reversible abnormalities in liver chemistry.

(3) SELECTIVE MUSCARINIC ANTAGONISTS (M1)

□ pirenzepine □ telenzepine

- ↓ Basal secretion (40- 50%).
- ↑ Gastric mucosal blood flow (M2 presynaptic on adrenergic fibers → ↓ Ne).
- ↑ Motility → ↑ LESF “lower esophageal sphincter pressure” (M1 receptors have a role in inhibitory motility pathway).

(4) PROSTAGLANDIN ANALOGUE, MISOPROSTOL (CYTOTEC)

- **A methyl analog of PGE1.**

Mechanism of action & pharmacodynamics:

1. Both acid inhibition & mucosal protection:

- Inhibits acid secretion (inhibits adenyl cyclase & gastrin release).
- Stimulates mucus and bicarbonate secretion.
- Increases blood flow.

• 2. Other actions:

- Stimulates intestinal electrolyte & fluid secretion.
- Stimulates intestinal motility.
- Stimulates uterine contraction.

Uses:

Prevention of NSAIDs-induced ulcers in high-risk patients.

Side effects:

1. Diarrhea & abdominal pain (10-20%).
2. Uterine contraction (abortion & vaginal bleeding).



2-neutralization of HCL

ANTACIDS

Chemical

Physical

Adsorb (HCL& pepsin) & Demulcent
1- Al^{+3} hydroxide gel. 2- Mg^{+2} trisilicate.

Systemic

❖ Na^{+} bicarbonate

Local (Non-systemic)

- 1 Mg^{+2} salts (Hydroxide & Trisilicate).
- 2 Al^{+3} salts (Hydroxide & Phosphate gel).
- 3 Ca^{+2} salts (Carbonate).

ANTACIDS

Pharmacological actions:

❑ Antipeptic effects:

❑ Reduction of gastric acidity will suppress activity of

pepsin: ❑ Activity decreases as PH increases above 2 and

❑ Irreversibly inactivated at PH 7

❑ Al⁺³ containing antacids → adsorb pepsin.



2. Effect on acid secretion: \uparrow PH (in gastric antrum) \rightarrow \uparrow gastrin \rightarrow rebound acid secretion.

3. Gastro- intestinal motor activity:

- A. \uparrow PH (of gastric content) \rightarrow \uparrow gastric motility (gastrin) \rightarrow \uparrow LESP.
- B. Al^{+3} \rightarrow relax smooth muscle of stomach (astringent) \rightarrow constipation.
- C. Mg^{+2} \rightarrow \uparrow cholecystokinin \rightarrow \uparrow motor activity.
- D. Mg^{+2} \rightarrow osmotic laxative effect.

MAGALDERATE [RIOPER]:

(AL HYDROXIDE + MAGNESIUM HYDROXIDE)

- Both magnesium and aluminum are absorbed and excreted by the kidney. Hence, patients with renal insufficiency should not take these agents for long-term therapy.

N.B. (milk-alkali syndrome)

Excessive doses of either sodium bicarbonate or calcium carbonate with calcium-containing dairy products can lead to hypercalcemia, renal insufficiency, and metabolic alkalosis .

3-ERADICATION OF HELICOBACTER PYLORI

B + M + A → FOR TWO WEEKS.

B

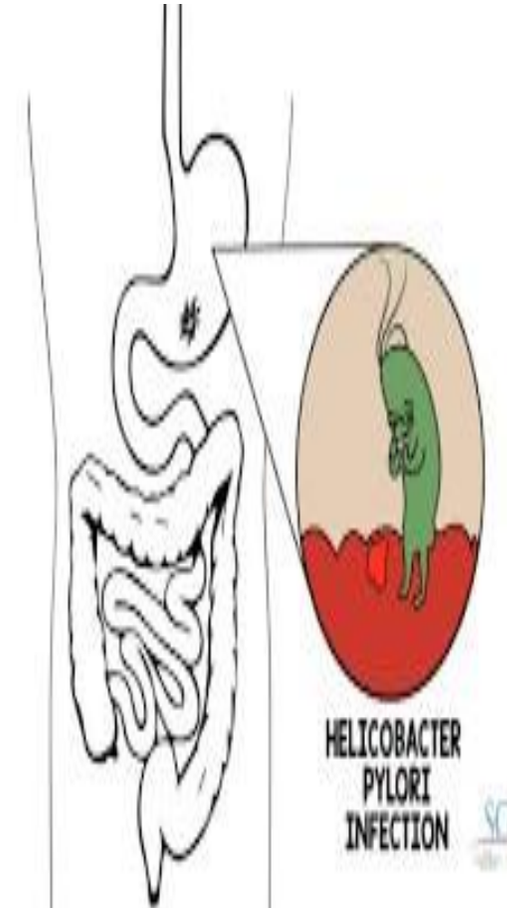
- Bismuth subcitrate (120mg four times daily).
- Bismuth subsalicylate (2 tablets; 262 mg each).

M

- Metronidazole (250 mg three times daily)
- Tinidazole (500mg bid)

A

- Amoxicillin (500mg three times daily).
- Tetracycline (500 mg four times daily).
- Clarithromycin (500mg three times daily).



Peptic ulcer & helicobacter pylori

Quadruple	<ul style="list-style-type: none">· Drugs that eradicate H Pylori + Anti-secretory drugs.
Triple	<ul style="list-style-type: none">· M + A + Antisecretory drugs. (Metronidazole+ Amoxicillin or Clarithromycin+ PPIs)
Dual	<ul style="list-style-type: none">· Amoxicillin + Omeprazole· Clarithromycin + Omeprazole

❖ These regimens are used for 10-14 days, then PPIs should be continued once daily for 4-6 weeks.

4-MUCOSAL PROTECTIVE AGENTS

A- Sucralfate: (sucrose octasulfate + Al^{+3} hydroxide)

Mechanism of action:

1. At acid PH (below 4) \rightarrow polymerization \rightarrow gel \rightarrow selective binding to necrotic ulcer tissues for up to 6 hrs. Sucrose sulfate (negatively charged) binds to proteins (positively charged) in the base of ulcers or erosion, forming a physical barrier.

2. Absorbs bile salts & pepsin.

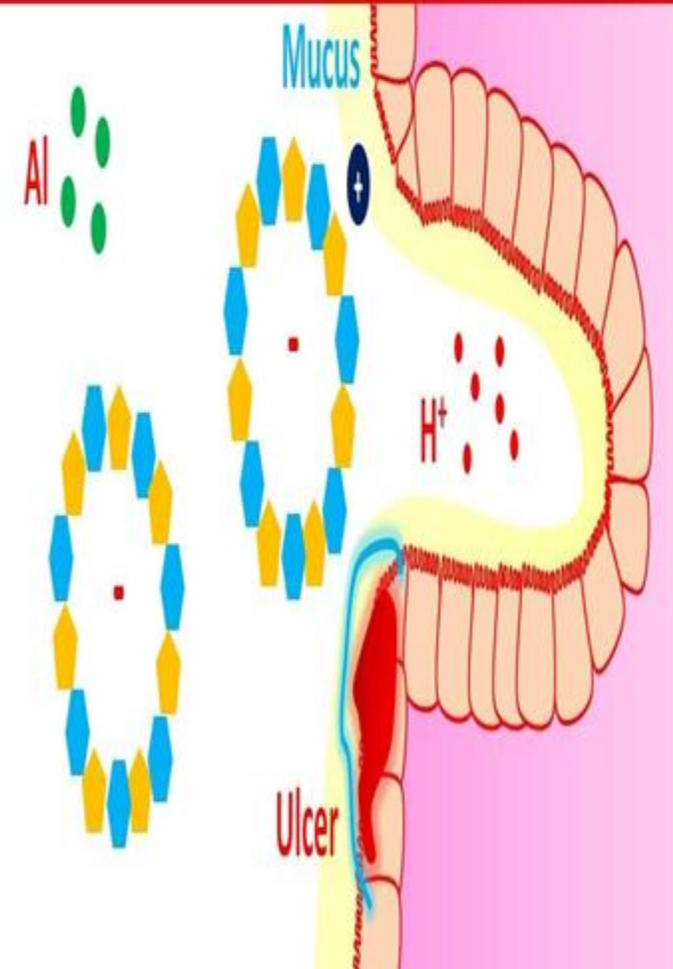
3. Stimulates PG & bicarbonate secretion

Side effects:

1- Constipation. 2- dry mouth.

3- 3% absorbed. Not be used for long period in patients with renal insufficiency. 4- adsorb [tetracycline, phenytoin, digoxin, cimetidine]

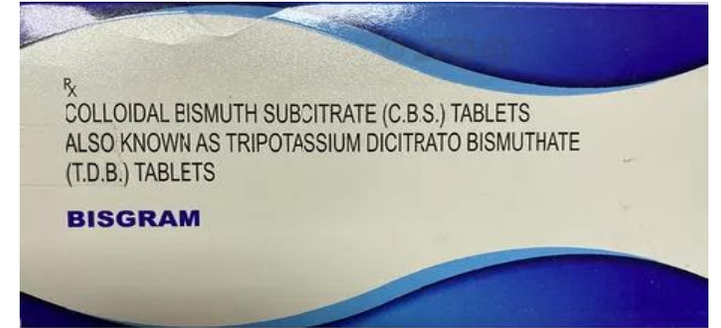
SUCRALFATE



B-BISMUTH COMPOUND: COLLOIDAL BISMUTH SUBCITRATE (DENOL):

□ **Mechanism of action:** (needs acid PH for activation).

- 1) Coats ulcer.
- 2) Stimulate the production of mucus and bicarbonates
- 3) Lysis of helicobacter pylori.
- 4) Decrease stool frequency and fluidity used in diarrhea of acute infections(travelers' diarrhea)



Side effects

-
- 1) Black color (oral cavity & stool). Blackening of stool, may be confused with G.I.T. Bleeding.
 - 2) Prolonged use → encephalopathy (ataxia, headaches, confusion, seizures). Thus, it should be used for short period only & avoid in renal impairment.

N.B.

Bismuth compound & sucralfate should not be administered simultaneously with antacids or H₂ antagonists.

C- Carbenoxolone (biogastrone):

- Synthetic derivative of liquorice.
- Mineralocorticoid activity → aldosterone-like side effect (salt and water retention).

Mechanism of action:

↑ Production, secretion & viscosity of intestinal mucus.

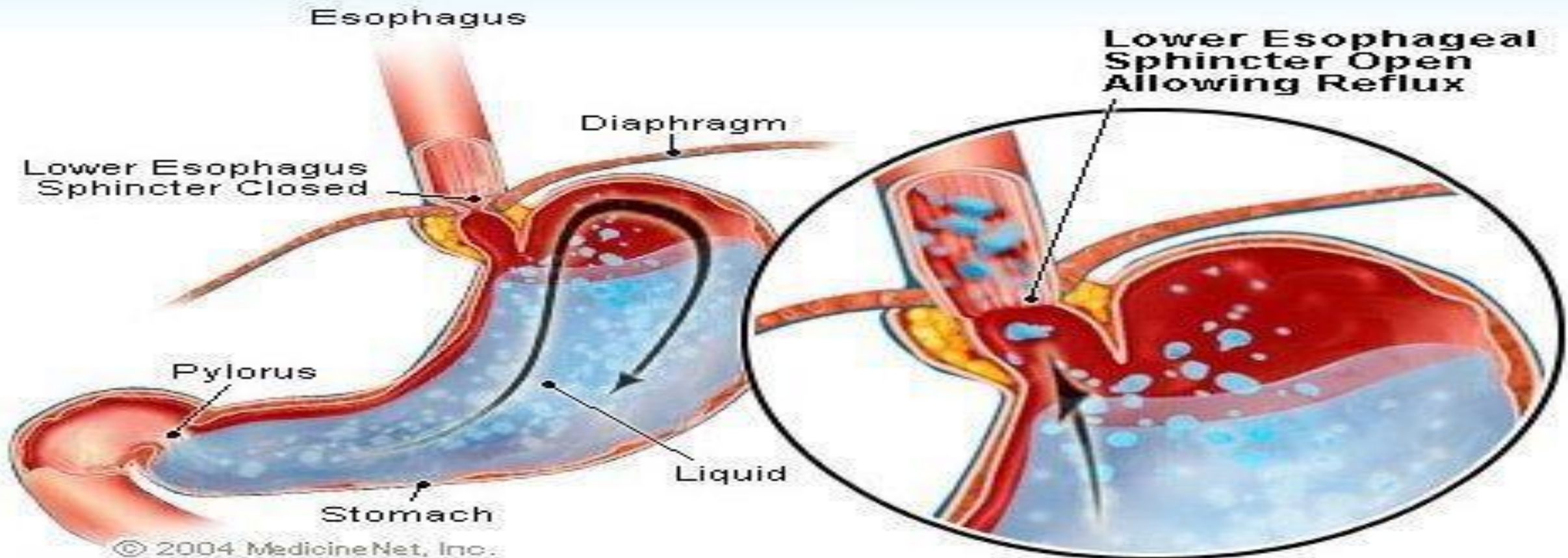
Side effects:

Na⁺ & water retention, hypokalemia & hypertension.



GASTRO-ESOPHAGEAL REFLUX DISEASE (GERD)

Gastroesophageal Reflux



MEDICAL MANAGEMENT ACCORDING TO SEVERITY OF GERD:

Stage I	<p>Sporadic uncomplicated heart burn, <u>less than 2-3 episodes/week</u>. Treated with:</p> <ul style="list-style-type: none">▪ <u>Life style</u> modification, including diet, weight loss, etc.▪ <u>Antacids and/or H₂-receptor</u> antagonists as needed.
Stage II	<p>Frequent symptoms more than <u>2-3 episodes/week</u> (with or without esophagitis).</p> <ul style="list-style-type: none">▪ Although <u>higher doses of H₂</u> antagonists increase healing rates, <u>PPIs</u> are preferred.
Stage III	<p>Chronic, unrelieved symptoms or immediate relapse after stopping therapy.</p> <ul style="list-style-type: none">▪ <u>PPIs either once or twice daily</u>.

GERD & pregnancy:

- Mild cases: conservatively, antacids or sucralfate.
- If symptoms persist: H2 receptor antagonists (ranitidine).
- Intractable symptoms or complicated reflux disease: lansoprazole.

GERD & children:

Omeprazole is safe and effective for treatment of erosive esophagitis & GERD.



- **Role of prokinetics in treatment of GERD:**
- Acid reflux is associated with transient LES relaxation that
 - occurs in absence of a swallow. The most effective therapy for
 - GERD still is suppression of acid production by the stomach.
- **Metoclopramide & domperidone:**
- used in treatment of symptomatic GERD but are not effective
in patients with erosive esophagitis.
- it is used mainly in combination with anti-secretory agents.



Thank You!