Antacids

Antacids are the substances which reduce gastric acidity resulting in an increase in the pH of stomach and duodenum. Gastric acidity occurs due to excessive secretion of HCl in stomach due to various reasons.

The pH of the stomach is 1.5-2.5 when empty and raises to 5-6 when food is ingested. Low pH is due to the presence of endogenous HCl, which is always present under physiological conditions. When hyperacidity occurs the result can range from:

1) gastritis (a general inflammation of gastric mucosa)
2) peptic ulcer or oesophageal ulcer (lower end of oesophagus)
3) gastric ulcer (stomach)
4) duodenum ulcers

Peptic ulcers occur due to defective oesophageal sphincter as in hiatal hernia. Gastric ulcers occur in lesser curvature and are found in first portion of duodenum. Symptoms include uncomfortable feeling from over eating, heart burn, growing hungry between meals. Complications involved are hemorrhage (being more common with duodenal ulcers), perforation. Depending upon the severity and location of an ulcer treatment will range from diet and antacids and/or anticholinergic therapy to complete bed rest to surgery. Small meals after short interval help in reducing acidity, stimulants of gastric acid must be avoided like coffee, alcohol, spicy food, oil or fried food.

Antacid therapy:

Antacids are alkaline bases used to neutralize the excess gastric HCl associated with gastritis or peptic ulcer. Since gastric HCl secretion is continuous, so is the administration of antacids.

Role of antacids:

1) Primarily in pain relief
2) Higher doses given continuously can promote ulcer healing
3) Superior to H$_2$ blockers in bleeding peptic ulcers

Criteria for antacids:

1) The antacid should not be absorber/or cause systemic alkalosis
2) It should not be constipative or laxative
3) It should exert effect rapidly and over a long period of time
4) The antacid should buffer in the range of pH 4-6
5) Reaction of antacid with HCl should not cause large evolution of gas.

Side effects of long term antacid therapy:

- If pH raises too high rebound acidity to neutralize the alkali occurs.
- Antacids which absorbed systemically exert alkaline effect on body’s buffer system.
- Some antacids cause constipation while others have laxative effect.
- Sodium containing antacids are problem for patients on sodium restricted diet.

Systemic antacids:

Systemic antacids are antacids which get systemically absorbed e.g. sodium carbonate is water soluble and potent neutralizer, but it is not suitable for the treatment of peptic ulcer because of risk of ulcer perforation due to production of carbon dioxide in the stomach.
Systemic absorption leads to alkalosis, may worsen edema and congestive heart failure because of sodium ion load.

**Non Systemic antacids**

They are insoluble and poorly absorbed systemically. In Magnesium salt, Magnesium carbonate is most water soluble and reacts with HCl at a slow rate, while Magnesium hydroxide has low solubility and has the power to absorb and inactivate pepsin and to protect the ulcer base. Aluminium hydroxide is a weak and slow reacting antacid. The aluminium ions relax smooth muscles and cause constipation. It absorbs pepsin at pH>3 and releases it at lower pH. It also prevents phosphate absorption. Calcium carbonate is a potent antacid with rapid acid neutralizing capacity, but on long term use, it can cause hypercalcemia, hypercalciuria and formation of calcium stone in kidney.

Every single compound among antacid have some side effect especially when used for longer period or used in elderly patients. To avoid certain side effects associated with antacids, **combinations of antacids** are used such as:

(i) Magnesium and aluminium containing preparation e.g. magnesium hydroxide a fast acting antacid with aluminium hydroxide which is a slow acting antacid.

(ii) Magnesium and calcium containing preparation where one is laxative and the later one is constipative in nature.

**Compounds used as Antacids**

(i)**Sodium Bicarbonate** (Baking soda)

\[ \text{NaHCO}_3 \]  

**M.W.** 84.01

**I.P. limit:** It contains not less than 99% and not more than 101% of NaHCO₃

**Properties:** White crystalline powder, odorless, with saline and slight alkaline taste, Stable in dry air, sparingly soluble in water, insoluble in alcohol.

**Preparation:**

1. By passing strong brine containing high concentrations of ammonia through a carbonating tower where it is saturated with carbon dioxide under pressure. The ammonia and carbon dioxide reacts to form ammonia bicarbonate which is allowed to react with NaCl to precipitate NaHCO₃ which is separated by filtration.

\[
\text{NH}_3 + \text{H}_2\text{O} + \text{CO}_2 \rightarrow \text{NH}_4\text{HCO}_3
\]

\[
\text{NH}_4\text{HCO}_3 + \text{NaCl} \rightarrow \text{NaHCO}_3 + \text{NH}_4\text{Cl}
\]

2. It can also be prepared by covering sodium carbonate crystals with water and passing carbon dioxide to saturation.

\[
\text{Na}_2\text{CO}_3 + \text{H}_2\text{O} + \text{CO}_2 \rightarrow \text{NaHCO}_3
\]

**Test for identification:** To 5ml of 5%w/v solution in carbon dioxide free water add 0.1ml phenolphthalein solution a pale pink color is obtained. On heating a gas is evolved and the solution turns red.
**For Sodium:**

1. To sample solution add 15% w/v potassium carbonate, heat, no precipitate is obtained add potassium antimonite solution heat to boiling, cool and if necessary scratch the inside of test tube with a glass rod, a dense white precipitate is produced.
2. Acidified the sample solution with 1M acetic acid and add excess of magnesium uranyl acetate solution yellow crystalline precipitate is obtained.

For bicarbonate: to sample add magnesium sulphate no precipitate is produced. On boiling a white colored precipitate is formed.

Assay: Weigh accurately 1gm and dissolve in 20ml of water, titrate the solution with 0.5N sulphuric acid using methyl orange as indicator. Each ml of 0.5N sulphuric acid = 0.0425gm of NaHCO₃

Use: It is used as antacid, and in electrolyte replacement.

(ii) Aluminium Hydroxide
Al(OH)₃ M. W. = 78.0

Aluminium hydroxide gel is an aqueous suspension of hydrated aluminium oxide with different amounts of basic aluminium carbonate and bicarbonate.

I.P. limit: It contains not less than 3.5% and not more than 4.4% of Al₂O₃

Properties: Aluminium hydroxide is a white, light odorless, tasteless amorphous powder. It is soluble in dilute mineral acids and in solution of alkali hydroxides but practically insoluble in water. It forms gel on prolonged contact with water at pH 5.5-8.0. It absorbs acids and carbon dioxide. The aluminium hydroxide gels are ideal buffers in the pH 3-5 range due to its amphoteric nature.

Preparation: It is prepared by dissolving sodium carbonate in hot water and the solution is filtered. To the filtrate add clear solution of alum (aluminium salt, chloride or sulphate) in water with constant stirring. Add more of water and remove all gas. The Aluminium Hydroxide precipitate out, collect the precipitate, wash and suspend in sufficient purified water flavoured with 0.01% peppermint oil and preserve with 0.1% sodium benzoate.

Test for identification: A solution in 2N HCl gives the characteristic reactions of aluminium salts. To sample add 5drops of freshly prepared 0.05%w/v solution of quinalizarin in 1% w/v solution of NaOH heated to boiling, cool, acidify with excess of acetic acid a reddish violet color is produced.

Assay: Accurately weigh 5gm and dissolve in 3ml HCl by warming on water bath, cool to below 20 °C and dilute to 100ml with water. To 20ml of this solution add 40ml of 0.05M disodium EDTA, 80ml water, 0.15ml methyl orange/red and neutralize by the dropwise addition of 1M sodium hydroxide. Again warm on water bath for 30 min, add 3gm hexamine and titrate with 0.05M lead nitrate using 0.5ml xylenol orange as indicator. Each ml of 0.05M disodium EDTA = 0.002549 gm of Al₂O₃

Uses: Aluminium hydroxide is used as antacid in the management of peptic ulcer, gastritis, gastric hyperacidity. It is also used as skin protectant and mild astringent.

(iii) Aluminium Phosphate
Aluminium phosphate consists of hydrated aluminium orthophosphate.
I.P. limit: It contains not less than 80.0% of Al PO₄

Properties: It occurs in nature as the minerals angelite, coeruleolactile, evansite, lucinite, sterrettite, wavellite etc. It is white infusible powder with some aggregates. It is insoluble in solutions of alkali hydroxide or acetic acid, water, alcohol etc; very slightly soluble in concentrated HCl and nitric acid. A 4% suspension in water has a pH of 5.5-6.5. Aluminium phosphate gel is a white, viscous suspension from which small amount of water may separate on standing. The gel has a pH in the range of 6.0-7.2.

Preparation:
1. It may be prepared by treating aluminium sodium oxide (NaAlO₂) with phosphoric acid
2. It can also be prepared by drying under suitable conditions the product of interaction in aqueous of an aluminium salt with an alkali phosphate such as sodium phosphate.

Test for Identification:

For aluminium: A solution in 2N HCl gives the reactions characteristic of aluminium salts. To sample add 5drops of freshly prepared 0.05%w/v solution of quinalizarin in 1% w/v solution of NaOH heated to boiling, cool, acidify with excess of acetic acid a reddish violet color is produced.

For phosphate: To neutral sample solution add silver nitrate solution, a light yellow precipitate forms, the color of which is not changed by boiling and is readily soluble in 10M ammonia and dilute HNO₃.

Assay: To acidify solution of weighed amount of the aluminium phosphate (0.8gm), 7.7gm disodium acetate is added, pH is adjusted to 4.5 with glacial acetic acid and dithizone in ethanol is added and sufficient quantity of ethanol is added. Titrate the solution with 0.05M zinc chloride until color turns red. A blank titration is also performed and the volume consumed by sample is calculated. Each ml of 0.05M disodium EDTA ≡ 6.098 mg of AlPO₄

Use: It is used as antacid.

(iv) Dihydroxy Aluminium Ammonium Acetate

I.P limit: It contains not less than 35.5% and not more than 38.5% of aluminum trioxide calculated on dried bases, contains small amounts of aluminum oxide and amino acetic acid.

Properties: It exists as very fine powder, bland taste, insoluble in water but forms suspension with water.

Preparation: It is prepared by adding solution of aluminum isopropoxide in isopropanolol to an aqueous solution of glycine.

Test for Identification:

For aluminium: A solution of sample in 2N HCl gives the characteristic reactions of aluminium salts. To the sample, add 5drops of freshly prepared 0.05%w/v solution of quinalizarin in 1% w/v solution of NaOH heated to boiling, cool, acidify with excess of acetic acid, a reddish violet color is produced.
For acetate: Heat the sample with an equal amount of oxalic acid, acidic vapours with characteristic smell of acetic acid are liberated
**Assay:** Transfer about 2.5gm of accurately weighed dihydroxy aluminium ammonium acetate and add 15ml of HCl and warm if necessary, to dissolve the sample completely. Transfer the solution with the aid of water to 500ml volumetric flask and dilute with water to volume and mix. Transfer 20ml and add 25ml 0.05M disodium EDTA and 20ml of acetic acid ammonium acetate buffer. Heat near boiling for 5min, cool and add 50ml of alcohol and 2ml of dithizone. Titrate with 0.05M zinc sulphate until green violet color turns rose pink. Perform blank and calculate the volume consumed by sample. Each ml of 0.05M disodium EDTA = 2.549 mg of Al$_2$O$_3$

**Use:** It is used as antacid.

**(v) Dihydroxy Aluminium Sodium Carbonate**

**I.P limit:** It contains not less than 90% and not more than 110.0% of Al$_2$O$_3$

**Preparation:** It is prepared by reaction between an alkoxide and sodium bicarbonate in water.

**Test for Identification:**

For aluminum: A solution in 2N HCl gives the reactions characteristic of aluminium salts. To sample add 5drops of freshly prepared 0.05%w/v solution of quinalizarin in 1% w/v solution of NaOH heated to boiling, cool, acidify with excess of acetic acid, a reddish violet color is produced.

For Carbonate: Suspend sample in 2ml of water in a test tube, add 2M acetic acid close the tube immediately with a stopper fitted with a glass tube bent at two right angles, heat gently and collect the gas in 5ml of 0.1M barium hydroxide a white precipitate is formed which is dissolves on addition of excess of dilute HCl.

For Sodium: To sample solution add 15% w/v potassium carbonate, heat, no precipitate is formed add potassium antimonite solution heat to boiling, cool and if necessary scratch the inside of test tube with a glass rod, a dense white precipitate is produced.

**Assay:** Weigh accurately powder not less than 200mg add 10ml of 2N HNO$_3$ cover and boil for 1 min, add 25ml of 0.1M disodium EDTA again boil for 1min cool, then add 10ml acetic acid ammonium acetate buffer and 50ml of acetone and 2ml of dithizone, adjust pH to 4.5 by the addition of ammonium hydroxide and titrate with 0.05M zinc sulphate maintaining pH at 4.5. Perform blank and calculate the volume consumed by sample. Each ml of 0.05M disodium EDTA = 5.098 mg of Al$_2$O$_3$

**Use:** It is used as antacid.

**(vi) Calcium Carbonate (precipitated chalk)**

Calcium carbonate is found in nature as limestone, marble, calcite, vaterite, aragonite and shell of sea animals.

**I.P. limit:** It contains not less than 98% and not more than 100.5% with reference to dried substance.

**Properties:** It occurs as a white, odorless tasteless microcrystalline powder which is stable in air. It exists in two crystal form and both are of commercial importance, one Aragonite and other is Calcite.
Precipitated chalk is prepared as a fine precipitate by adding a solution of ammonium carbonate and ammonia or sodium carbonate to a solution of calcium nitrate.

**Preparation:**
1) It can be prepared by mixing and boiling calcium and sodium carbonate solution and allowing the resulting precipitate to settle. The precipitate is collected, washed with boiling water until free from chloride and dried.
2) By passing carbon dioxide through lime water

**Test for Identification:**
**For Calcium:** Dissolve substance in 5M acetic acid and add 0.5ml of potassium ferrocyanide solution. The solution remains clear. Add ammonium chloride white crystalline precipitate is formed.

**For Carbonate:** Suspend sample in 2ml water in a test tube, add 2M acetic acid close the tube immediately with a stopper fitted with a glass tube bent at two right angles, heat gently and collect the gas in 5ml of 0.1M barium hydroxide a white precipitate is formed which is dissolves on addition of excess of dilute HCl.

**Assay:** Accurately weigh 0.1gm, dissolve in 3ml of dilute HCl, add 10ml of water. Boil the solution for 10min, cool, dilute with 50 ml with water. Titrate the solution with 0.05M disodium EDTA to with a few ml of the expected end point and add 8ml of NaOH solution and 0.1g of calcon mixture. Continue the titration until the color changes from pink to blue. Each ml of 0.05M disodium EDTA = 0.005004g of CaCO$_3$

**Uses:** It is used as fast acting antacid, in calcium deficiency, dentrifies and in combination with magnesium containing antacids due to its constipative properties.

**(vii) Tribasic Calcium Phosphate**

**I.P limit:** Tribasic calcium phosphate consist of variable mixture of calcium phosphate having approximate composition of 10CaO 3P$_2$O$_5$H$_2$O not less than 34% and not more than 40% of calcium and an amount of phosphate equivalent to not less than calcium phosphate calculated with reference to ignited substance.

**Properties:** It is a white odourless, tasteless, amorphous powder practically insoluble in water, alcohol or acetic acid but readily soluble in dilute HCl and HNO$_3$.

**Preparation:**
1. It is manufactured from bones which are calcined until white, powdered and digested with sulphuric acid. The insoluble tribasic calcium phosphate is converted to soluble phosphoric acid and insoluble calcium sulphate. The solution is filtered and the filtrate treated with calcium hydroxide to precipitate calcium phosphate.
2. Decomposition of calcium chloride and sodium phosphate in presence of aqueous ammonia at high temperature yield calcium phosphate. The white precipitate is filtered washed and freed from chloride and dried.
Test for Identification:
For Calcium: Dissolve substance in 5M acetic acid and add 0.5ml of potassium ferrocyanide solution. The solution remains clear. Add ammonium chloride white crystalline precipitate is formed.
For Phosphate: To neutral sample solution add silver nitrate solution, a light yellow precipitate forms, the color of which is not changed by boiling and is readily soluble in 10M ammonia and dilute HNO₃.

Assay:
For calcium: Weigh accurately 0.2gm and dissolve in HCl, triethanol amine and hydroxynaphthol blue indicator are added and titrate with 0.05M disodium EDTA until blue colour is obtained. Each ml of 0.05M disodium EDTA ≡ 0.002004gm calcium.
For phosphate: Acidify the aqueous solution of substance 0.2gm with dilute nitric acid, filter and add strong ammonium solution to produce slight precipitate. The precipitate is dissolved in dilute nitric acid, ammonium molybdate is added and precipitate is filtered, washed with potassium nitrate solution and redissovled in 1N sodium hydroxide add phenolphthalein and titrate the excess alkali with 1N sulphuric acid. Each ml of 1N NaOH ≡ 0.006743 gm of calcium phosphate.
Use: It is used as antacid, as non hygroscopic diluent, as an abrasive in tooth pastes.

(viii) Magnesium Carbonate
Magnesium carbonate is a hydrated basic magnesium carbonate containing 40-45% of magnesium oxide. It occurs in nature as the meniral magnate and lansfordite.
Heavy Magnesium Carbonate: 15 g occupy a volume of about 30ml
Light Magnesium Carbonate: 15 g occupy a volume of about 150ml
I.P. limit: It contains not less than 40% and not more than 45% of magnesium oxide
Properties: Both heavy and light magnesium carbonate are hydrated. Both are white, odorless powder practically insoluble in water and alcohol but solubilizes in dilute acids with strong effervescence.
Preparation: It is prepared by mixing hot solution of magnesium sulphate and sodium carbonate. The mixture is evaporated to dryness and the residue consisting of magnesium carbonate and sodium sulphate is digested for half an hour with boiling water. The precipitate of magnesium carbonate is collected on filter paper, washed with water until free from sulphate and then dry.

Test for Identification:
For Carbonate: Suspend sample in 2ml water in a test tube, add 2M acetic acid close the tube immediately with a stopper fitted with a glass tube bent at two right angles, heat gently and collect the gas in 5ml of 0.1M barium hydroxide a white precipitate is formed which is dissolves on addition of excess of dilute HCl.
For Magnesium: to solution of sample add dilute nitric acid solution a white precipitate is produced that is redissovled by adding 1ml of 2M ammonium chloride, add 0.25M disodium hydrogen phosphate a white crystalline precipitate is produced.
**Assay:** Accurately weigh 15g of magnesium carbonate and dissolve in a mixture of 20ml of water and 2ml 2M HCl. To this solution add 50ml of water and 10ml strong ammonia ammonium chloride solution titrate this with 0.05M disodium EDTA using mordant black II mixture as indicator until blue color is obtained. Each ml of 0.05M disodium EDTA = 0.002015g MgO

**Uses:** It is used as antacid and mild laxative. It is used as pharmaceutical aid (dispensing volatile oil for use in inhalants).

**(ix) Magnesium Oxide** (Magnesia)

**I.P. limit:** It contains not more than 98% of magnesium oxide

It occurs in nature as mineral periclase. It occurs in two varieties heavy magnesium oxide which is relatively dense white powder with 15 g occupying volume of about 30ml while light magnesium oxide is very bulky with 15 g occupying volume of about 150ml.

**Properties:** Both heavy and light magnesium oxide are odorless taste slightly alkaline, practically insoluble in water yield a solution which is alkaline. It readily dissolves in dilute acids with slight effervescence. In presence of acid, the oxide forms the magnesium hydroxide, therefore the chemistry and pharmacology are same as those of magnesium hydroxide.

**Preparation:**
1. It can be prepared by heating gently magnesium carbonate to redness.
2. It is also prepared by heating light magnesium carbonate to redness.

**Test for identification:**
**For Magnesium:** to solution of sample add dilute nitric acid solution a white precipitate is produced that is redissolved by adding 1ml of 2M ammonium chloride, add 0.25M disodium hydrogen phosphate a white crystalline precipitate is produced.

**Assay:** The assay of magnesium oxide is performed by complexometry. Accurately weigh 15g of magnesium oxide and dissolve in a mixture of 20ml of water and 2ml 2M HCl. To this solution add 50ml of water and 10ml strong ammonia ammonium chloride solution titrate this with 0.05M disodium EDTA using mordant black II mixture as indicator until blue color is obtained. Each ml of 0.05M disodium EDTA = 0.002015g MgO

**Uses:** It is used as antacid and laxative. It is ingredient of universal antidote along with tannic acid and charcoal. It is used for compounding and preserving fluid extract because of its absorptive power.

**(x)Magnesium Phosphate**

**I.P. limit:** It contains not less than 98% and not more than 101.5% substance ignited at 425°C to constant weight.

**Properties:** It is white, odourless, tasteless powder readily soluble in dilute mineral acids but practically insoluble in water. The chemistry is same as tribasic calcium phosphate.
Test for Identification:

For Magnesium: to solution of sample add dilute nitric acid solution a white precipitate is produced that is redissolved by adding 1ml of 2M ammonium chloride, add 0.25M disodium hydrogen phosphate a white crystalline precipitate is produced.

For phosphate: To neutral sample solution add silver nitrate solution, a light yellow precipitate forms, the color of which is not changed by boiling and is readily soluble in 10M ammonia and dilute HNO₃.

Assay: Weigh accurately 200mg previously ignited to 425°C to constant weight and dissolve in mixture of 25ml of water and 10ml of 2N HNO₃. Filter if necessary, wash any precipitate add sufficient 6N ammonium hydroxide to the filtrate to produce a slight precipitate and then add 1ml of 2N HNO₃. Adjust temperature to 50° add 7ml ammonium molybdate and maintain temperature for 30min with occasional stirring. Wash the precipitate once or twice with water by decantation. Transfer precipitate to the filtrate, wash with KNO₂ solution until the last washing is not acid to litmus. Transfer precipitate and filter to precipitate vessel add 50ml of water and 40ml of 1N NaOH, agitate until the precipitate is dissolved, add phenolphthalein and titrate excess of alkali with 1N H₂SO₄. Each ml of 1N NaOH ≡ 5.716mg of magnesium phosphate.

Uses: It is used as antacid and laxative.

(xi) Magnesium Trisilicate

I.P. limit: It contains not more than 29% of magnesium oxide and not more than 65% of silicon dioxide both calculated with reference to the ignited substance.

Properties: Fine, white, odorless, tasteless powder, free from grittiness. Magnesium silicate hydrate is a compound of magnesium oxide and silicon dioxide with varying amount of water.

Preparation: It is prepared by precipitation from solution of magnesium sulphate and sodium silicate.

Test for identification:

For Magnesium: to solution of sample add dilute nitric acid solution a white precipitate is produced that is redissolved by adding 1ml of 2M ammonium chloride, add 0.25M disodium hydrogen phosphate a white crystalline precipitate is produced.

For silicate: in a lead or platinum crucible mix by means of a copper wire to obtain a thin slurry the prescribed amount (0.25gm) of substance with 10mg of sodium fluoride and a few drops of sulphuric acid cover the crucible with thin transparent plate of plastic under which a drop of water is suspended and warm gently within a short time a white ring is formed around the drop of water

Assay:

For Magnesium Oxide: Weigh accurately 1gm substance and dissolve in 35ml of water allow to stand for 15min on water bath. Cool the contents to room temperature filter and wash the residue with water and dilute the combined filtrate and washing to 250ml with water. Neutralize 50ml of this solution with about 8ml of 10M NaOH and then add 10ml ammonia buffer pH 10, 50mg mordant black II mixture. Heat the contents to 40 °C and titrate with
0.05M disodium EDTA until color changes to deep blue. Each ml of 0.05M disodium EDTA = 0.002015g MgO

**For silicon dioxide:** weigh accurately 0.7gm of substance add 10 ml of 1M sulphuric acid, 10ml water and heat on water bath for 1.5hrs. shaking frequently and replacing the evaporated water. Allow to cool decant on to an ashless filter paper (7cm in diameter). Wash the precipitate by decantation with three quantities each of 5ml of hot water, transfer it to the filter paper and wash it with hot water, until 1ml of the filtrate remains clear on addition of 2ml of barium chloride solution and 0.5ml of 2N HCl. Ignite the filter paper and its contents in a tarred platinum crucible at 900 °C to constant weight. The residue is silicon dioxide.

**Uses:** It is used as non systemic antacid and adsorbent

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**PREPARED BY**

**B.KIRUTHIGA**
**LECTURER**
**DEPT OF PHARMACEUTICAL CHEMISTRY**