Effects of Ranitidine on Insulin and Lime - Induced Gastric Secretion in Albinowistar Rats

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Abstract

Purpose: To study the possible effect(s) of a relative H2-receptor blocker, ranitidine on lime and insulin-induced gastric secretion in male and female albino rats.

Methods: The rats were divided into 3 groups of lime juice, insulin and control in triplicates after 24hr starvation to empty the stomachand were cannulated (oesophageal, tracheal and gastric) using Gosh and Schild method. Using N saline, the acid content of the effluent was recorded. The 1st group of rats was perfused with lime solution (25% v/v, 50% v/v, 75% v/v and 95% v/v) and was used to modify the secretory rates of the parietal cells and the stomach effluent was collected 3 times in 30 minutes. In same manner, the 2nd group was perfused with 40 IU/kg insulin. The 3rd group (control) had no lime nor insulin. Ranitidine was administered (2.5ml ) intramuscularly and the results noted.

Results: The mean basal secretion significantly (P≤ 0.05) increased from 22.82 ±4.6mMol/L/hr to 52.94 ±10.23mMol/L/hr, while 2.5ml ranitidine (Zantac) injected intramuscularly decreased the basal stimulation from 52.94 ±10.23mMol/L/hr to 34.77 ± 5.09 mMol/L/hr. Insulin (40 IU/kg) was administered intravenously to the 2nd group of rats, and the mean basal secretion increased from 8.01±0.75 mMol/L/hr to 10.00±0.71 mMol/L/hr. Ranitidine was administered intramuscularly and that caused a significant decrease in the insulin stimulation from 10.00±0.71 mMol/L/hr to 8.01±0.75 mMol/L/hr.

Conclusions: Results obtained showed marked higher gastric secretion in females than in males, although steady increase was observed for both insulin and lime inducement. Generally, results obtained from this study indicated, a statistically significant decrease in both lime juice and insulin-induced gastric secretion by Ranitidine, and hence submits Ranitidine as a possible potent drug for peptic ulcer treatment.

Keywords: Agonists; Gastric acid secretion; Insulin; Lime; Inhibition

Introduction

One of the functions of the stomach is the secretion of gastric acid by oxyntic cell. Gastric juice secreted by the gastric gland of the stomach contains a number of gastric enzymes and acid, which were identified in 1824 to be hydrochloric acid. The acid plays a key role in digestion of potassium chloride and sodium chloride (Sembulingam and Sembulingam, 2010) and in digestion of proteins, by activating digestive enzymes, and making ingested proteins unravel so that digestive enzymes can break down the long chains of amino acids (Ward et al., 1963).

The ultimate source of H+ is water. The ultimate source of chloride ion is the sodium chloride found in blood. The reaction (overall) can best be summarized in the reaction below:

$$\text{CO}_2 + \text{H}_2\text{O} + 2\text{NaCl} \rightarrow 2\text{HCl} + \text{Na}_2\text{CO}_3$$

For each mole of HCl produced, a mole of Na2CO3 is formed. The carbon IV oxide used for the reaction is derived from the parietal cell metabolism itself and also from the cellular metabolism of other tissues in the body and for every H+ produced, a bicarbonate ion is released into the intestinal fluid and ultimately into the blood. This explains the so called alkaline tide seen in the urine during digestion (Soll and Read, 1991).

On stimulation, the morphological structure of the parietal cells have been found to change during secretion of gastric acid and...
The synthesis and secretion of hydrochloric acid by oxygenic cells have been studied extensively; H+ are secreted to obtain pH of 1.0 when intracellular pH is 7.1-7.2, achieving a 2 million fold concentration gradient. Carbonic anhydrase catalyze the formation of bicarbonate and H+ from carbon IV oxide and water (Davenport, 1939). An enzyme (H+/K+ ATPase), located on the surface of the oxygenic cell lumen, secretes H+ across the apical membrane in exchange for potassium ion, by a separate pathway. Potassium ion and chloride ion are secreted into the canalicular space, down their concentration gradient with water following the electrochemical and osmotic balance that is maintained across the apical membrane. Secretion of this large amount of acid into the lumen requires that an equivalent amount of base be secreted from the basolateral surface of the oxygenic cell to maintain the intracellular pH in the physiological range. Chloride ions are exchanged for hydroxyl ion (OH-) concentration. The Na+/K+ - ATPase located on the basolateral membrane maintains the higher intracellular potassium ion concentration (Vasudevan et al., 2007).

A variety of substances have been found to support acid secretion including glucose and fatty acids (Konturek et al., 1977; Roeckpe et al, 2006).

New classes of antisecretory drugs, the substituted benzimidazole appear to act by inhibition of this H+/K+/ATPase. Most recent data suggest that acid secretion at the canalicular membrane may be regulated by the potassium chloride permeability of the canalicular membrane. Potassium chloride permeability would increase at the transition to stimulated state to provide K+ a luminal face of the H+/K+/ATPase for exchange (Geibel, 2005).

Acid secretion is commonly divided into 3 phases. Viz: i) Cephalic Phase, ii) Gastric Phase and iii) Intestinal phase.

Cephalic Phase: Only a small proportion of gastrin is released during this phase, perhaps in part because, the fasting stomach is acidic and H+ inhibits gastrin release. There could be abundant secretion of gastric juice even though there is no food in the stomach. This is readily demonstrated by sham feeding (Sembulingam and Sembulingam, 2010).

Gastric Phase: As food and fluids enters the stomach, the buffering effect of the content raises the pH, removing the inhibitory effect of the low pH on gastric release; the effect is abolished by acidification of gastric content. Since secretion is also suppressed by atropine, and H2 antagonists, it is probably not due to direct effect on oxygenic cells. As food enters into the stomach, two factors are known to be operative; which are distension of the stomach and stimulation by chemical mediators (Yang, 2002).

Intestinal Phase: The secretion is mediated via release in the intestine of an unknown peptide, enterostin, the stimulus for its release appears to be the presence of peptides and some amino acids. The latent period of the intestinal phase is two to three hours, but the secretion when initiated may last an average of 6 hours. This phase is characterized by less activity than the other two phases. The following have been observed to execute gastric secretion when placed in the small intestine, water, extractive substances of meat, products of protein digestion, milk, alcohol, histamine, saponin, epinephrine, 0.1N HCl, 10% glycine solution and magnesium sulfate (Troidl et al., 1975).

The chemical mediators of gastric secretion are clearly acetylcholine (a secreting neuron which innervates the gastric gland, can directly activate parietal cells and also the antral G-cells which release the hormone gastrin terminals can directly activate parietal cells and also the antral G-cells which release the hormone gastrin), gastrin (A peptide hormone that caused a strong stimulation of gastric acid secretion, released from the G-cells of the pyloric antrum and the two principally active forms of gastrin are small gastrin (G-17) and big gastrin (G-34)) , and histamine (Uvans, 1969).

Insulin: Insulin was named by Demeyer in 1907 and obtained or isolated by Banting and Best in 1922 in Madeods Laboratory. It is a peptide hormone secreted by the beta cells of the islets of Langerhans, found in the pancreas. The hypoglycemia produced by insulin stimulates the sympathetic as well as parasympathetic centers. Stimulation of the latter causes the signals to be sent through the vagus nerves to cause gastric acid secretion and pepsin secretion (Schubert and Peura, 2008).

Histamine: The existence of histamine or -aminoethylimidazolide was demonstrated in 1910 by Sir Henry Deland Barge while working on ergot alkaloids. It was until 1927 that Best, Dale, Duddy and Thorpe isolated histamine from fresh sample of lungs and live, thereby establishing beyond doubt that histamine was a natural constituent of the body (Soll, 1978). Paton reported that histamine was present in all parts of the body, except bones and cartilages. All animal species have relatively high level of histamine in the glandular region of the stomach mucosa, where acid is secreted, and that there is a species difference on the distribution of histamine in the gastric mucosa, example in the rats, histamine is distributed in the enterochromaffin cells while in man, it is found in the mast cells. Histamine has two types of receptors located throughout the body, which when occupied by histamine or one of its analogues produces different effects. Some of these effects such as broncho-constriction and contraction of gut are mediated by the H1 receptors (Sandrik and Waldum, 1991) and are readily blocked by the classical antihistamine such as mepyramine and pyrilamine. Other effect such as gastric acid secretion is completely refractory of such antagonists and involves the activation of H2 receptor and is susceptible to the newly developed H2 antagonists.

Highly effective H2-antagonist, Buramamide retained the imidazole ring of histamine believed to be important for receptor recognition, but possessed a much bulkier side chain. Examples of such drugs are metiamide, mizatidine, and oxetidine, cimetidine and famotidine (Samuelson and Hinkle, 2003). It was found in Glaxo laboratory, that activity was retained in analogues of imidazole series when the imidazole moiety was replaced by an aminomethyl aromatic system. This gave rise to the compound ranitidine . Biochemical regulation of this secretion involves central signals conveyed by the vagus nerve and local mechanism mediated by cholinergic and peptidergic fibers of the gastric wall, as well as amine or peptides secreting cells located in the fundic and antral epithelia (Uvans, 1969). The blockade of this receptor by specific antagonist results in a surmountable inhibition of gastric acid secretion. The blockade of H+ K+ -ATPase is an alternative means of inhibition, this can be achieved by series of benzimidazole derivatives ion-dazopyrimidine which specifically accumulates in the
Parietal cell secretory canaliculus and covalently binds to ATPase inhibitory site (Roekpe et al., 2006; Code, 1949). It has longer than histamine antagonist. Therefore, H+ K+-ATPase inhibitors are of special interest in the treatment of acid-related diseases such as gastric and duodenal ulcer, as well as esophagitis reflux (Oliver, 1960; Feldman et al., 1998).

Peptic Ulcer: This is a breach in the mucosa of the digestive tract produced by digestion of the mucosa by pepsin and acid. This may occur when pepsin and acid are present in abnormally high conc. or when some other mechanism reduces the normal protective mechanisms of the mucosa. The results from Lee et al. (2001) correlated with peptic ulceration and evidence has shown that over secretion of gastric acid and gastric juice can be corrosive to the gastric mucosa and can consequently lead the development of peptic ulceration. Emotional and physical stress, hypermotility, vascular spasm and thrombosis, infection, allergy and reduced and altered mucosa production are some of the factors that could cause ulceration of the mucosa.

Ranitidine (Zantac): Accordingly, an improved H2-antagonist has been sought and results now described, lead us to believe that ranitidine, N-2, 5 (dimethyl amino), methyl-2-nitro-1, 1-ethenedia-mine is such a drug. Ranitidine was synthesized by members of the chemistry division, Glaxo Group research limited. Ranitidine is a new specific histamine antagonist which differ chemically from other histamine receptor blockers in having a furan rather than an imidazole or thiazole ring structure. Comparison with citrilidine has shown ranitidine to be four to ten times more potent antagonist. However, cimetidine has been shown to be a specific histamine H2-receptor antagonist despite the substitution of the imidazole by a furan ring. Ranitidine like cimetidine is effective clinically in the short term management of duodenal ulcer.

Ranitidine inhibits the acid and pepsin secretion which is induced by various stimuli (histamine, lime, pentagastrin, meal). The inhibitory effect on histamine is an expected of a competitive nature, that on pentagastrin is non-competitive. Depending on the test system used, ranitidine is up to ten times more potent than cimetidine on molar basis (Prichard et al. 1986). Ranitidine does not seem to affect cardiovascular function in a close range that can be useful. In duodenal ulcer patients, doses of ranitidine assumed to be therapeutically effective, does not change the concentration of follicle stimulating hormone, luteinizing hormone, prolactin, oestradiol and testosterone in the plasma, thus suggesting that the drug does not affect the endocrine system.

Lime (Citrus Aurantifolia): Lime is a term referring to citrus fruit which is typically round, green to yellow in color (Hulma, 1971), 3-6cm in diameter and containing sour and acidic pulp. Lime is a good source of vitamin and is often used to accent the flavors of foods and beverages. Lime juice may be squeezed from lime and is used as limeade, and as an ingredient in many cocktails. When the skin is exposed to ultraviolet light, after lime juice contact, a reaction known as phytophotodermatitis can occur, this can cause darkening of skin, swelling or blistering. The agent responsible for this is Psoralen. Lime juice and its oil are very beneficial for skin when consumed orally or applied externally. It rejuvenates the skin, keeps it shining protects it from infections and reduces body odor due to large amounts of Vitamin C and flavonoids both of which are antioxidants. When applied externally onto the skin, lime acids slough off dead cells (Kokwara, 2009). Lime also helps in areas where circulation is a problem; such is the case with allulite. The therapeutic properties of lime oil in skin are antiseptic, antiviral, disinfectant bactericidal, astrigent, restorative and tonic. Trivett and Meyer (1971) reported that citric acid is one of the most common acids in lime and is present in small quantity (8-9%). The quantity of the acid varies with season, ripeness and variety. The juice predominates in citric acid than the peel, while the peel of lime fruit has a higher content of oxalic acid than citric and malonic acid (Hulma, 1971). He also reported that chemical changes during maturation were broadly reverse of those of orange and grape juice. This study seeks to evaluate the following:

i) effects of insulin and lime juice on gastric acid secretion in the Wistar albino rats.

ii) effects of ranitidine on insulin and lime induced acid secretion in Wistar albino rat.

Materials And Methods (Gosh And Schild Method)

Albino Wistar rat were sourced from Animal house of the Biochemistry department of the University of Port Harcourt Nigeria. Each rat (200-250g) was starved for 24 hours/overnight in order to empty the stomach by the following day. Each rat was then weighed and divided into 3 groups of 5 each for males, females and control. Anesthesia (Urethane) was administered (0.6mg per 100gram body weight) intraperitionally along the linea alba. Each rat used, slept, within five minutes of the administration of the anesthesia. However, if the dose was not enough, an extra 0.1-0.2ml was given.

The rat was then pinned to the dissecting board and the dissecting light turned on (Fig 12). The skin around the neck of the rat was then removed, along the midline to expose the thyroid gland which was then separated by blunt dissection to expose the trachea (Fig 2.0). The trachea was then freed from the surrounding tissues. A skin - porter was passed under the trachea and a tracheostomy was performed by making a semi-transection of the upper rings of the trachea. A tracheal canula (dipped in Normal saline) was then inserted and it was then tied into position using needle and thread avoiding the vagus nerve. This allowed the animal to breathe freely. However, the canula was monitored to detect mucus accumulation due to irritation. If present, it was removed with a hypodermic needle. The wound was then covered with cotton wool damped with normal saline.

The skin over the linea Alba was then lifted up and out to expose the abdominal muscle. The abdominal muscle was then incised along the linea alba but the incision was then brought towards the left hypochondria. The stomach was then brought out by fingers along the linea alba but the incision was then brought towards the left hypochondria. The stomach was then brought out by fingers dipped in normal saline. After, a cut was then made on the anterior surface not too far from the pylorus. A gastric canula was introduced into the stomach and tied into position and switched back to keep temperature warm.

An esophageal tube connected to the perfusion bottle was then introduced into the stomach via the mouth and the esophagus. Normal saline kept at constant temperature of 370C was then introduced and allowed to flow through the tube from the perfusion bottle and the residual contents of the stomach were completely flushed out and the efficient discarded. When the stomach effluent was found to be free of food debris, the flow rate was regulated to give a volume of 10ml+ 4ml/minute at time interval of 10 minutes. More of the saline thus, had to be added to the bottle when needed.
After the flow rate had been adjusted, 3 basal collection of the stomach effluent were titrated against sodium hydroxide to find the concentration of the acid content of the effluent. The saline was removed from the perfusion bottle and quickly replaced with different contents of lime juice (25% v/v, 50% v/v, 75% v/v, and 95% v/v) and 3 basal collections were made at 10ml + 4mls/minutes at time interval of 10minutes. Each basal collection was titrated against sodium hydroxide and the results were noted (for n=5) and mean values taken accordingly.

Ranitidine (Zantal, 50mg) was administered (2.5ml) intramuscularly and 3 check 4 repetition basal collection was made at time interval of 10minutes. On each rat, lime of 25% v/v, 50% v/v, 75% v/v and 95% v/v were used to flush the stomach and 3 basal collections were titrated against sodium hydroxide and the results were noted.

The second group of rats was anaesthetized, dissected and canulated. Normal saline in the perfusion bottle was used to flush the stomach via the mouth to free the stomach from food and other debris. 3 basal collections were made and titrated against sodium hydroxide, for each rat. Insulin (0.5ml / body kg) was administered intramuscularly (1.m). The stomach effluent was then collected 3 times in 30 minutes. Ranitidine was then administered intramuscularly at a dose of 25ml/body weight, 3 basal collections were made and titrated against sodium hydroxide as described above. Statistical analysis was done using Student’s t – test to essentially compare the means of paired samples.

Results

Lime-Induced Gastric Secretion

The results obtained for lime-induced gastric secretion in albino wistar rats (3males and 2 females) with lime content of 25% v/v, 50% v/v, 75% v/v and 95% v/v within 30mins are as given in tables 3.1 and 3.2. Stimulation and inhibition are also presented.

Insulin-Induced Gastric Acid Secretion

The results obtained for insulin-induced gastric secretion in 5 wistar albino rats (male and female groups) is as given below. Stimulation and inhibition are also presented.

Discussion

Lime-induced gastric secretion in female Wistar albino rats.

Female albino Wistar rats were used to test the effect of relative new H2-receptor blocker, ranitidine. The mean basal secretion was 17.30±9.2mMol/L/hr and increased to 54.86±9.2mMol/L/hr after perfusion with lime of different concentration (25%, 50%, 75% and 95%) (Fig 2). This increase in acid secretion caused by lime juice was statistically significant on applying the student t-test and did not occur by chance. Administrations of the test drug, ranitidine (2.25mg/kg), after 30 minutes, caused a decrease in gastric secretion from 54.86±9.2mMol/L/hr to 29.78±7.67mMol/L/hr (Fig 3). Thus, the decrease in acid secretion caused by ranitidine was found to be significant and thus did not occur by chance.

Lime-induced gastric secretion in male Wistar albino rats.

Male albino Wistar rats gave a mean basal of 28.33±11.14 mMol/L/hr which increased to 51.71±10.20 (Table 3.2) with lime. Administration of the test drug decreased the stimulation by lime juice to 29.78±7.67 (graph 2). This significant increase by lime juice was inhibited by the test drug and did not occur by chance but was marked. Higher gastric secretion in females was observed (Figs 2 and 4) and secretion increased with weight (data not shown).

Insulin-induced gastric secretion in female Wistar albino rats.

Female albino wistar rats were induced with 0.5ml insulin (Actrapid) and the basal mean secretion increased from 8.70±0.62mMol/L/hr to 10.10±1.01mMol/L/hr. (Fig 4). Administration of the test drug inhibited secretion from 10.10±1.01...
mMol/L/hr to 8.70±0.62 mMol/L/hr. (Fig 5) Student’s t – test showed that these did not occur by chance but was significant.

**Insulin-induced gastric secretion in male Wistar albino rats.**

These rats were induced with insulin (0.5ml) which caused an increase from 7.89 1.10 mMol/L/hr mean basal secretion to 9.90±0.37 mMol/L/hr. (Fig 4). This decreased to 7.96±0.37 mMol/L/hr (Fig 5), on administration of the test drug. These increase and decrease did not occur by chance.

The great increase in acid secretion is believed to be strength dependent. The probable mechanism by which lime caused an increase in gastric secretion is through the gastric mechanism and by direct effect on the parental cells (Kay, 1953). Insulin on the other hand caused an average increase in acid secretion. Insulin has been found to stimulate acid secretion through its effect on the vagus nerve. Insulin-induced hypoglycemia stimulates the vagal nerve which in turn stimulates acid secretion (Forte and Zhu, 2010; Aihara et al., 2003).

Ranitidine which is a H2-receptor blocker, caused an inhibition due to its action on the H2-receptor, which mediates secretion, and its action is said to be specific. From this experiment, ranitidine exerted a greater effect on lime-induced gastric secretion than insulin. This explains that histamine is not a final common mediator for the action of other stimulation, but a potentiate (Bonfils et al., 1979; Pritchard et al., 1986). As previous studies have shown that, inhibition is not as a result of reduced mucosal blood flow (Prema and Smbulingam, 2010; Troidl et al., 1975), it was observed that the antisecretory activity of ranitidine stems from a selective action at H2-receptor. Since in addition to its blocking activity, ranitidine does not possess anti-androgenic activity function and also does not inhibit the mixed oxygenase metabolizing enzyme system in the liver (Feldman et al., 1998), it possesses a long duration of action (El – Serag ans Sonnenberg, 1998).

![Figure 2: Mean Basal (MBS) and mean lime-induced (LIS) acid secretion in albino wistar rats](image)

![Figure 3: Mean inhibition by ranitidine on lime-induced gastric secretion in albino wistar rats](image)
Conclusion

From the results obtained in this study, lime juice significantly enhanced gastric acid secretion and therefore may not be beneficial to peptic ulcer patients. Also, results obtained shows that there is significantly higher gastric secretion in females than in males. This means that peptic ulcer may be predominant in females than in male. Again, there was more secretion in rats with higher weight than in those with lower weight; this implies that peptic ulcer patients should be conscious of weight gain. Also, it was observed that insulin caused hyperacidity (hypoglycemia), therefore starvation should be avoided especially in peptic ulcer patients and since lime juiced caused a significant increase in gastric secretion, peptic ulcer patients should stay away from lime juice and derivatives. Finally, since Ranitidine produced a significant decrease in both lime juice and insulin-induced gastric secretion, it could pose a potent drug for peptic ulcer treatment.
References


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