Mechanisms of histamine stimulated secretion in rabbit ileal mucosa*

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SUMMARY Histamine is present in high concentrations in the intestine and we investigated the possibility that it might have a role here in intestinal transport. When added to the basal side of rabbit ileal mucosa in vitro histamine (10⁻⁴M) induced a short-lived increase in electrical potential difference and short circuit current. It inhibited net chloride absorption but did not influence sodium transport. Alkali secretion, measured by a pH stat technique, was inhibited, suggesting that bicarbonate secretion was reduced. Both the electrical and ion flux responses to histamine were blocked by the H₁ receptor blocker diphenhydramine, but not by the H₂ receptor blocker cimetidine. The presence of specific H₁ histamine receptors was further supported by shifts in the dose-response curve to histamine by four different concentrations of diphenhydramine. Calculation of a pA₂ value from these 'Schild' plots provided a figure of 7·85, which is similar to that for H₁ receptors in other tissues. Aminoguanidine, a histaminase blocker, had no electrical effects alone but shifted the histamine dose response curve to the left. These studies indicate that histamine inhibits chloride absorption and alkali secretion, possibly by influencing a chloride/bicarbonate exchange process, through specific mucosal H₁ receptors. Enhancement of histamine effects by a histaminase inhibitor suggests that histaminases are present in the intestinal mucosa and supports the possibility of a role for endogenous histamine in influencing ion transport. The observations indicate a mechanism by which absorption might be impaired in diseases in which histamine is liberated locally in the intestine.

The intestinal mucosa contains a rapidly turning over pool of histamine which appears to be derived largely from non-mast cell sources.¹⁻³ Its physiological role here, however, is uncertain, although in systemic mastocytosis, in which histamine is liberated locally in high concentration, diarrhoea is a feature.³¹ It is possible therefore that histamine may act as an intestinal secretogogue and this possibility was examined in rabbit ileal mucosa in vitro.

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Methods

Using an in vitro technique previously described in detail,⁶ segments of stripped ileal mucosa were mounted in Perspex flux chambers. Tissues were bathed on both sides in buffer maintained at 37°C and of composition Na 146, K 4·2, Cl 125·8, HCO₃ 26·6, H₂PO₄ 0·2 HPO₄ 1·2, Ca 1·2, Mg 1·2, glucose 10 mMol/l, and pH 7·4. The buffer was stirred and oxygenated by a 95% oxygen 5% CO₂ gas lift system. In the experiments using a bicarbonate-free buffer, chloride was substituted for bicarbonate and 100% O₂ was bubbled through the buffer. Transmucosal potential difference was measured via saturated KC1 in agar bridges and calomel electrodes using a high impedance digital voltmeter. Sodium chloride in agar bridges to silver/silver chloride electrodes were used to transmit the short-circuit current. This was adjusted initially each minute and later every five minutes. Experiments conducted later were performed using
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an automatic voltage clamp apparatus, which continuously adjusted the short-circuit current. Corrections were made for the fluid gap resistance as described by Field et al.4 Tissue resistance (R) was calculated from the short-circuit current and potential difference and expressed in Ohms per cm².

Short-circuit current was converted to net ion flux in μmol/cm²/h by multiplication by a factor of 3.6 x 10⁸.

0.033 derived from ——— where A equals area of exposed tissue and F is Faraday’s constant.

Ion fluxes were determined in tissue pairs obtained from the same animal and with resistances varying by less than 25%. Usually eight tissues were set up at the same time. Steady state fluxes were usually obtained 15 minutes after addition of isotope and thus flux measurements were started 30 minutes after isotope addition to ensure equilibration. Fluxes were measured during the 15 minute period before addition of drugs (control flux) and for 15 to 30 minutes starting 15 minutes after addition of drugs when equilibration conditions had been re-established. Control fluxes were also measured in some experiments during four consecutive 15 minute periods. Unidirectional fluxes from mucosa to serosa and from serosa to mucosa were measured in each tissue pair after the addition of 0.5μCi²⁶Na and 2.5μCi⁳⁶Cl to the mucosal side of one tissue and the serosal side of its paired tissue. One millilitre samples were taken from mucosal and serosal solutions and replaced by 1 ml warm unlabelled buffer. Calculation of fluxes was performed as described previously4 and fluxes were determined under short-circuit conditions with minimal interruption to read the spontaneous potential difference.

With the use of a pH stat technique7 net alkali secretion was determined on the mucosal side of the tissue bathed in the bicarbonate free buffer and bubbled with 100% oxygen. The serosal bathing solution was a bicarbonate buffer bubbled with 95% oxygen 5% CO₂. The mucosal solution was maintained at pH of 7.4 by continuous titration with 5μmol/l hydrochloric acid titrant. This was carried out with an Autotitrator 11 and Autoburette ABU 13 (Radiometer, Copenhagen) which very accurately deliver the small volumes of hydrochloric acid needed to maintain the pH constant as measured by combined electrodes GK2321C and GK2332C. The volume of titrant needed to maintain the pH constant during a 30 minute control period and one hour after addition of histamine was recorded at five minute intervals. From the amount of acid added it is possible to calculate the rate of alkali (presumably bicarbonate) secreted. Potential difference and short-circuit current were also measured during this experiment.

An attempt was made to identify specific histamine receptors by studying the effect, on a histamine dose-response curve, of four different concentrations of the specific H₁ receptor antagonist diphenhydramine and calculating a pA₂ value from these dose response curves¹⁰. pA₂ values, a measure of drug antagonism, are an accepted basis for receptor classification based on the hypothesis that agonists and antagonists compete for receptors according to mass law¹⁰. The same pA₂ values should be obtained with similar receptors in different tissues with the same agonist/antagonist combination⁹. pA₂ is defined as the negative logarithm of the molar concentration of the antagonist which will reduce the effect of a multiple dose of the agonist (x) to that of a single dose. More simply, pA₂ is the negative logarithm of the molar concentration of antagonist which halves the sensitivity of the preparation to the agonist¹¹. These values are dependent on contact time between antagonist and tissue. A fixed time of 14 minutes was taken in these experiments to allow comparison with published data on pA₂ in other tissues. For dose response and agonist/antagonist interaction studies only one dose of agonist or antagonist was used in any single piece of mucosa to avoid persistence of effects and the possibility of tachyphylaxis.

Aminoguanidine, a specific inhibitor of the enzyme responsible for histamine catabolism, was used to determine whether histaminase was present in intestinal mucosa. Tissues were pretreated with 10⁻⁴M aminoguanidine (a dose which completely inhibits histaminase¹³) and histamine dose response, using maximal percentage potential difference increases, were determined and compared with results from control tissues.

Statistical analysis was made using Student’s t test for paired and unpaired samples and variability is expressed as mean ± one standard error of the mean. Materials used were histamine acid phosphate (Sigma Chemical Corporation), diphenhydramine (Park Davies Ltd), cimetidine (Smith, Kline & French Ltd), and aminoguanidine (Sigma Chemical Corporation).

Results

Histamine added to the aerosol side of ileal mucosa significantly increased potential difference, short-circuit current, and tissue resistance, the changes in potential difference and short-circuit current being maximal at two minutes post histamine (Fig. 1). Potential difference returned to control values within 10 minutes but resistance remained raised for
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Fig. 1 Changes in potential difference, short-circuit current, and resistance of ileal mucosa after histamine 10^{-4}M added to rabbit ileal mucosa bathed in glucose buffer (mean ± SEM shown). *Significantly different from control response (p < 0.001).

Fig. 2 Dose response curves for histamine added to the serosal and mucosal side of ileal mucosa. Mucosal application was ineffective. Significantly different, ∆p < 0.01, *p < 0.001.

Fig. 3 Effects of diphenhydramine (10^{-4}M) and cimetidine (10^{-4}M), on short-circuit current changes induced by histamine 10^{-4}M (glucose-free buffer). Diphenhydramine completely prevented the response to histamine, while cimetidine was without effect.

at least 30 minutes after histamine was added. The electrical response was reproducible and dose-related. A dose response curve (Fig. 2) demonstrated that the maximal response was achieved at a concentration of 10^{-4}M and the smallest dose at which a significant response was noted was 10^{-5}M. Histamine in concentrations greater than 10^{-3}M failed to increase potential difference or short-circuit current. Histamine added to the mucosal side of tissues failed to produce an electrical response (Fig. 2).

The H2 receptor blocker cimetidine in concentrations between 10^{-2} and 10^{-4}M did not influence electrical values in control tissues, nor did it influence the electrical response to histamine. However, equimolar concentrations of diphenhydramine, an H1 receptor blocker, completely prevented the...
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Histamine-induced electrical response (Fig. 3). In addition, diphenhydramine lowered potential difference and short-circuit current in tissues bathed in glucose-free buffer but not in those in glucose-containing buffer.

The influence of four different concentrations of diphenhydramine on the dose response curve to histamine is shown in Fig. 4, the curve being shifted to the right by the increasing concentrations of H1 blocker. The shift to the right of the concentration of histamine required to produce a half maximal control electrical response was used to calculate a pA2 value as described in the legend to Figs. 4 and 5 and shown in Fig. 5. From this graph a pA2 value of 7.85 was calculated, which is similar to pA2 values for H1 receptors in other tissues.

![Graph](image)

**Fig. 4** Dose response curves for histamine-induced increases in potential difference with and without diphenhydramine (DIPH) in rabbit ileum. The maximum increase in potential difference obtainable with histamine (2-2 mV) was taken as 100% and all other results related to that value. The dose response curve to histamine is increasingly shifted to the right by 10^-1, 5 x 10^-2, 10^-2 M diphenhydramine. The extent to which the dose of histamine required for a half maximal control response is shifted by the different concentrations of H1 blocker indicated at A, B, C, D, and E was used to calculate the pA2 value as shown in Fig. 5.

In the histaminase inhibition studies, amino-guanidine 10^-1 M had no electrical effect when given alone. However, in tissues pretreated with amino-guanidine, histamine provoked an electrical response at lower concentrations than in control tissues. The curve was shifted to the left by aminoguanidine pretreatment (Fig. 6), suggesting that histaminases are present in the tissues.

<table>
<thead>
<tr>
<th>Dose ratio (from Fig. 4)</th>
<th>Log (dose ratio-1) Diphenhydramine concentration (mol)</th>
<th>-Log diphenhydramine concentration</th>
</tr>
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<tr>
<td>BA = 3.0</td>
<td>Log 2 00 = 0.30</td>
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</tr>
<tr>
<td>CA = 4.75</td>
<td>Log 3 75 = 0.57</td>
<td>5 x 10^-7</td>
</tr>
<tr>
<td>DA = 6.50</td>
<td>Log 5 50 = 0.74</td>
<td>10^-6</td>
</tr>
<tr>
<td>EA = 12.5</td>
<td>Log 11.50 = 1.06</td>
<td>10^-5</td>
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**Fig. 5** Plot of log (dose ratio-1) vs—log antagonist concentration to determine pA2 (Schild plots).

**ION FLUXES**

Preliminary studies with control tissues bathed in glucose buffer alone showed that after 30 minutes' equilibration following addition of isotopes there was no significant change in sodium, chloride, or residual ion fluxes during four consecutive 15 minute periods (n = 8).

Steady state fluxes were achieved by 15 minutes after addition of histamine. Fluxes in control tissues and during the periods 15–30 minutes after histamine are shown in the Table (A). Net chloride absorption was significantly reduced (P < 0.01) due to a reduction in mucosa to serosa flux (P < 0.01). It should be noted that these fluxes were measured at a time when the transient electrical responses had largely resolved. There was no obvious effect on sodium transport. The calculated residual ion flux, which represents the unmeasured ions making up the total net ionic movement inferred from the short circuit current, was significantly reduced after histamine (P < 0.01). This result is compatible with a reduced secretion of bicarbonate ions (or increased secretion of hydrogen ions).

Alkali secretion, measured by the pH stat technique, confirmed that this residual ion flux was likely to be due, at least in part, to movement of bicarbonate or hydrogen ions. As shown in Fig. 7, alkali secretion was reduced after histamine from
Fig. 6 Dose response curves for histamine in control and aminoguanidine (10⁻⁴M) pretreated tissues. The curve is shifted to the left by aminoguanidine. (Mean ± SEM shown; *Significantly different from control response (p<0·001).

Table Ion fluxes and short-circuit current in rabbit ileal mucosa: responses to histamine (A); histamine in diphenhydramine pretreated tissues (C); diphenhydramine alone (B)

<table>
<thead>
<tr>
<th></th>
<th>Jms</th>
<th>Jnet</th>
<th>Jms</th>
<th>Jnet</th>
<th>C1⁻</th>
<th>Jnet</th>
<th>SCC</th>
<th>Jˀnet</th>
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<tr>
<td><strong>A</strong> Histamine 10⁻⁴M (n=9)</td>
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<td></td>
<td></td>
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<tr>
<td>Control</td>
<td>16·3</td>
<td>10·2</td>
<td>6·1</td>
<td>12·2</td>
<td>8·1</td>
<td>4·1</td>
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</tr>
<tr>
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<td><strong>B</strong> Diphenhydramine (10⁻⁴M) alone (n=11)</td>
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<tr>
<td>Diphenhydramine</td>
<td>12·2</td>
<td>8·9</td>
<td>3·3</td>
<td>9·5</td>
<td>8·1</td>
<td>1·4</td>
<td>2·8</td>
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</tr>
<tr>
<td></td>
<td>±1·0</td>
<td>±0·8</td>
<td>±0·6</td>
<td>±0·7</td>
<td>±0·7</td>
<td>±0·4</td>
<td>±0·3</td>
<td>±1·1</td>
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<tr>
<td><strong>C</strong> Histamine 10⁻⁴ after diphenhydramine 10⁻⁴M (n=9)</td>
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<td></td>
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<td></td>
<td></td>
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<tr>
<td>Control (no drug)</td>
<td>12·9</td>
<td>8·3</td>
<td>4·6</td>
<td>9·2</td>
<td>8·2</td>
<td>0·9</td>
<td>4·8</td>
<td>1·2</td>
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<tr>
<td></td>
<td>±0·5</td>
<td>±0·6</td>
<td>±0·6</td>
<td>±0·3</td>
<td>±0·3</td>
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</tr>
<tr>
<td>Histamine</td>
<td>12·6</td>
<td>7·7</td>
<td>4·8</td>
<td>9·6</td>
<td>7·8</td>
<td>1·8</td>
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<td>±0·6</td>
</tr>
</tbody>
</table>

In each instance control fluxes for the 15 minutes before addition of drugs and responses to histamine or diphenhydramine were measured in the periods 15 to 30 minutes after drug addition. In study C histamine was added 15 minutes after diphenhydramine and fluxes measured 15 to 30 minutes after histamine. Fluxes and short-circuit current are in μmol/cm²/h. Jsm = flux from mucosa to serosa. Jnet = net flux. Jˀnet = calculated residual ion flux. Mean ± SEM; *p<0·02; †p<0·01; ‡p<0·001.

2·0±0·3 to 1·4±0·3 μmol/cm²/h (n=8, p<0·01).

Although the flux data for the 15 minute period immediately after histamine addition are not presented, net chloride absorption was reduced (4·1 to 1·5 μmol/cm²/h, p<0·02, n=9) because of a decrease in mucosa to serosa flux (12·2 to 9·8 μmol/cm²/h, p<0·01, n=9). The time courses for the effect of histamine on alkali secretion (Fig. 7) and on chloride flux are thus similar.

Ion fluxes were measured in response to histamine after pretreatment with diphenhydramine (10⁻⁴M). The changes in chloride fluxes induced by histamine
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were shown to be completely prevented by this H1 blocker, although a slight fall in short-circuit current persisted (Table, C). Diphenhydramine (10⁻⁴M) alone did not influence sodium or chloride fluxes and a fall in residual ion flux was not statistically significant (Table, B). The decrease in short-circuit current induced by diphenhydramine in tissues in glucose buffer was not significantly different from the slight fall in short-circuit current seen in control tissues over the same time period.

Discussion

The presence of histamine receptors in the musculature of the small intestine has been supported by several investigations but their presence in intestinal mucosa has not been well described. Infusions of histamine into the mesenteric arterial system of dogs provoked marked secretion into the lumen, but it was suggested that this was due to leakage of fluid through a capillary network whose permeability had become markedly increased. Recent studies with a number of histamine receptor antagonists, however, have supported the idea that histamine receptors may be involved in intestinal ion transport, at least in rabbit ileum. Interpretation of the latter studies is dependent on the specificity of the histamine receptor antagonists used (see below) but the current studies lend additional weight to the concept of specific H1 receptors in rabbit ileal mucosa.

The electrical responses to histamine were shown to be specifically blocked by diphenhydramine in a dose-related manner. The calculated value of 7.85 for pA₂ is very similar to the values calculated for histamine/diphenhydramine relationships in other tissues (8.1, 8.0, and 7.8 in guinea-pig ileal muscle, 7.8 in guinea-pig trachea, and 7.8 in guinea-pig lung) supporting the interpretation that specific H1 receptors are involved.

The residual ion flux changes were shown, by the pH stat technique, to be due, at least in part, to a reduction in bicarbonate secretion. The residual ion flux change was larger than the change in titratable alkali but this is hardly surprising because the former was a calculated figure derived from the sum of Na and Cl fluxes and short-circuit current and the latter a direct measure in a different group of tissues at a different time. It is thus not possible to judge whether all or only a part of the change in residual ion flux in these studies could be attributed to bicarbonate (of H⁺) movement changes. The results presented here thus support the possibility of a chloride-bicarbonate exchange which is inhibited by histamine through an H1 receptor but additional effects on other unmeasured ions cannot be excluded.

It is conceivable that the changes in potential difference and short-circuit current were due not to effects of histamine on epithelium but rather to effects on submucosal tissue such as the muscularis mucosae. The specific H1 blockade demonstrated here could then theoretically represent a demonstration of H1 receptors on non-epithelial tissues. However, this seems somewhat unlikely, particularly as steady ionic flux changes, reached 15 to 30 minutes after histamine, were shown to be specifically blocked as well.

The observation from Fromm and Halpern that H1 receptor blockade alone significantly reduced residual ion flux and bicarbonate secretion suggests the possibility that endogenous histamine may be acting on the mucosa under the basal conditions of their study. However, a comparison of their and our results reveals some discrepancies. Firstly, we could not demonstrate a flux change in response to H1 receptor blockade alone. This may be due to our having used diphenhydramine, a blocker reputed to have minimal local anaesthetic activity compared with the pyrilamine primarily used by Fromm and Halpern. It is conceivable that the response to the latter agent was related to its local anaesthetic activity rather than its H1 receptor blocking activity. They report a similar response with diphenhydramine but only in a concentration 10 times higher than that used in our study. Secondly,
their response to H1 receptor blockade is not in keeping with that expected simply from antagonism to the actions of histamine demonstrated here. Our results would suggest that blockade of any endogenous histamine activity which may have been present should have enhanced bicarbonate secretion (and Cl− absorption) rather than the reverse. While the electrical and flux changes induced by histamine have been shown to be prevented by the H1 blocker used in our experiments, this and other blockers may also have an action independent of H1 blocking activity, particularly in high concentrations.

Our observations that aminoguanidine, a histaminase inhibitor, enhances the effect of histamine suggests that histaminases are present in intestinal mucosal preparations and indeed they have been localised in intestinal villous cells. This observation supports the possibility that histamine may have a physiological role in this site. The lack of effect of aminoguanidine alone suggests that it is unlikely that endogenous production of histamine is occurring to an appreciable extent in this in vitro preparation in the basal state.

Preliminary studies of electrical responses to histamine in rabbit jejunal and colonic mucosa and in human colonic mucosa suggest that these regions of the intestine respond in a similar manner (unpublished observations).

It is of interest to contrast the effects of histamine on the stomach with those on the intestine. In the former its activity seems to be mediated through H2 receptors, while in the intestine it is clear that H1 receptor mediated activity is responsible for its effect on transport. These data do not, of course, demonstrate a physiological role for histamine in intestinal transport but they do provide a possible mechanism by which absorption may be impaired in circumstances where histamine is liberated locally in the intestine. Such may be the case, for example, in intestinal allergies and systemic mastocytosis.

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References