Effect of vasoactive intestinal polypeptide (VIP) antagonism on rat jejunal fluid and electrolyte secretion induced by cholera and *Escherichia coli* enterotoxins

F H Mourad, C F Nassar

Abstract

Background—The enteric nervous system is important in the pathophysiology of intestinal fluid secretion induced by cholera toxin (CT), *Escherichia coli* heat labile (LT), and heat stable (STa) toxins. The neurotransmitters involved are not fully elucidated. Vasoactive intestinal polypeptide (VIP), a potent intestinal secretagogue present in the enteric nervous system, is increased after exposure of the cat intestine to CT. Whether VIP is involved in the pathogenesis of cholera and other toxins is not known.

Aim—To study in vivo the effect of VIP antagonism on jejunal fluid secretion induced by CT, LT, and STa.

Methods—CT, LT (25 µg), or 0.9% NaCl was instilled in an isolated 25 cm segment of rat jejunum, and the VIP antagonist (VIPa) [4Cl-D-Phe6, Leu17]-VIP (0.2 or 2 µg/kg/min) or 0.9% NaCl was given intravenously. Two hours later, single pass in vivo jejunal perfusion was performed to assess fluid movement. In STa experiments, intravenous VIPa or 0.9% NaCl was given and 30 minutes later the jejunal segment was perfused with a solution containing STa 200 µg/l.

Results—VIPa had no effect on basal intestinal fluid absorption. CT induced net fluid secretion (median −68 µl/min/g dry intestinal weight (interquartile range −80 to −56)) which was dose dependently reversed by VIPa (6.2 (−16 to 34) and 29 (17 to 42); p<0.01). Similarly, LT induced secretion (−63 (−73 to −30)) was attenuated by VIPa (0.2 µg/kg/min) (−15 (−24 to −1); p<0.01) and totally reversed to normal levels by VIPa (2 µg/kg/min) (37 (28–56); p<0.01 compared with LT and not significantly different compared with normal controls). STa induced secretion (−17 (−19 to −2)) was also reversed by VIPa (12 (9–23) and 14 (0–26); p<0.01).

Conclusion—VIP plays an important role in CT, LT, and STa induced intestinal secretion and may be the final putative neurotransmitter in the pathophysiology of these toxins.

(Intestional secretion induced by cholera toxin (CT) and the structurally related enterotoxin, *Escherichia coli* heat labile toxin (LT), has been attributed solely to direct activation of adenylate cyclase with a corresponding increase in intracellular cAMP. However, evidence has been accumulating on the importance of other mediators such as 5-hydroxytryptamine (5-HT), prostaglandins, neurotensin, and of the enteric nervous system in the pathophysiology of CT induced secretion. In addition, vasoactive intestinal polypeptide (VIP) has been found to be released in response to administration of CT. VIP is present in high concentrations in the subepithelial neural plexuses and is a potent secretagogue which acts via specific adenylate cyclase linked receptors. For these reasons, it seems likely that VIP plays an important physiological and pathophysiological role in the intestine. The proposed mechanism for CT induced secretory diarrhoea includes a neural reflex with a common final pathway through VIP to the epithelial cells. However, this proposal remains to be confirmed in studies proving that VIP is important in the pathophysiology of CT induced secretion and is not just an innocent bystander.

The pathophysiology of diarrhoea caused by other enterotoxins has not been as extensively investigated. LT, which is structurally similar to CT, binds to the same receptor on enterocytes (GM1 ganglioside) and has been shown to stimulate adenylate cyclase with a corresponding increase in cAMP production. However, the two toxins are not identical in their amino acid composition and their binding affinity is different. In addition, we have previously demonstrated that 5-HT plays an important role in CT but not LT induced secretion suggesting that there are fundamental differences in the pathophysiology of fluid secretion caused by these toxins. Furthermore, Nzegwu and Levin have shown that intraluminal capsaicin inhibits LT induced secretion implying that LT acts through afferent nerve fibres. The neurotransmitters involved in the effenter loop of this neural reflex are not known. Whether VIP plays a role in LT...
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It has been shown that, stable toxin (STa) induced intestinal secretion with a plasma electrolyte solution (PES) after two hours the clamps were removed and abdomen closed. The intestine was returned to the abdominal cavity and the abdomen closed. Cannulae. The intestine was returned to the isolated segments followed by clamping of both the proximal and distal ends. The intestine was instilled in the isolated segments followed by clamping of both the proximal and distal cannulae. The intestine was returned to the abdominal cavity and the abdomen closed. After two hours the clamps were removed and the intestine perfused at a rate of 0.5 ml/min with a plasma electrolyte solution (PES) containing Na 140, K 4, Cl 104, and HCO3, 40 mmol/l to which 5 g of cold polyethylene glycol 4000 (PEG) and 4 µCi/l of [14C]-PEG were added. Thirty minutes were allowed to elapse to ensure establishment of a steady state following which consecutive 10 minute collections of the effluent were obtained from the distal cannula for one hour. All rats had femoral veins were cannulated. VIP antagonist (0.2 or 2 µg/kg/min) was started in the second femoral vein and 30 minutes later VIP infusion (0.02 or 0.2 µg/kg/min) was started in the second femoral vein. One hour later, intestinal perfusion was performed as mentioned above to calculate fluid and electrolyte movement.

In a different group of animals, both femoral veins were cannulated. VIP infusion (0.2 or 2 µg/kg/min) was started directly after instillation of STa EXPERIMENTS. The mean of net fluid and solute movement of six consecutive effluent samples were calculated and expressed, respectively, as µl/min/g and µmol/min/g of dry intestinal weight. Positive values denote net absorption and negative values net secretion.

Materials and methods
CT AND LT EXPERIMENTS
Adult Sprague-Dawley rats (180–220 g body weight) fasted for 18 hours were anesthetised with intraperitoneal injection of sodium pentobarbital (60 mg/kg) and maintained throughout the experiments by interval intraperitoneal injections (15–30 mg/kg) as necessary. The abdomen was opened through a midline incision and cannulae inserted into the small intestine proximally (5 cm distal to the duodenojejunal junction) and 25 cm distally in the jejunum fixed by ligation as previously described. CT, LT (25 µg in 3 ml of 0.9% NaCl), or 3 ml of 0.9% NaCl alone (controls) was instilled in the isolated segments followed by clamping of both the proximal and distal cannulae. The intestine was returned to the abdominal cavity and the abdomen closed. After two hours the clamps were removed and the intestine perfused at a rate of 0.5 ml/min with a plasma electrolyte solution (PES) containing Na 140, K 4, Cl 104, and HCO3, 40 mmol/l to which 5 g of cold polyethylene glycol 4000 (PEG) and 4 µCi/l of [14C]-PEG were added. Thirty minutes were allowed to elapse to ensure establishment of a steady state following which consecutive 10 minute collections of the effluent were obtained from the distal cannula for one hour. All rats had femoral veins were cannulated. VIP antagonist (0.2 or 2 µg/kg/min) was started in the second femoral vein and 30 minutes later VIP infusion (0.02 or 0.2 µg/kg/min) was started in the second femoral vein. One hour later, intestinal perfusion was performed as mentioned above to calculate fluid and electrolyte movement.

In a different group of animals, both femoral veins were cannulated. VIP infusion (0.2 µg/kg/min) on intestinal secretion induced by 1 mM carbachol added to the perfusate was tested. Steady state conditions in all experiments were demonstrated by less than 5% variation in fluid movement between consecutive 10 minute collections. Also, values were accepted only if recovery of radioactive PEG was 95–105%.

ANALYTICAL METHODS
[14C]-PEG concentrations in the effluent were measured in duplicate by liquid scintillation spectroscopy in an LKB Wallac 1409 scintillation counter. Sodium and chloride concentrations were determined by an Hitachi 911 analyser.

The mean of net fluid and solute movement between pairs were tested using the Wilcoxon rank sum test.

Materials
Cholera toxin, Escherichia coli heat stable and heat labile toxins, VIP, the VIP antagonist, and carbachol were obtained from Sigma Chemical Company (St Louis, Missouri, USA). Radio-labelled polyethylene glycol ([14C]-PEG 4000) was obtained from Amersham International (Buckinghamshire, UK) and all other chemicals were supplied by British Drug House (BDH Chemicals, Poole, UK).

Results
Basal fluid absorption (median 61 µl/min/g (interquartile range 49–81); n=11) was not affected by the VIP antagonist at 0.2 or 2 µg/kg/min (48 (28–79), n=7, and 55 (34–75), n=6, respectively). Chloride and sodium movement paralleled that of fluid (table 1).

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CT AND LT EXPERIMENTS
Net fluid secretion occurred in all animals who received CT (~68 µl/min/g (~80 to ~56), n=7)
Table 1: Effect of vasoactive intestinal polypeptide (VIP) antagonism on net Na⁺ and Cl⁻ flux in normal controls, in cholera toxin (CT), Escherichia coli heat labile toxin (LT), and Escherichia heat stable toxin (STa) induced secretion, and on VIP induced decrease in absorption

<table>
<thead>
<tr>
<th>Condition</th>
<th>n</th>
<th>Net Cl⁻ flux (µmol/min/g)</th>
<th>Net Na⁺ flux (µmol/min/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>11</td>
<td>2.8 (2.4 to 3.3)</td>
<td>6.7 (5.7 to 11.1)</td>
</tr>
<tr>
<td>+VIPa0.2</td>
<td>7</td>
<td>3.5 (2.0 to 4.3)</td>
<td>7.4 (4.5 to 12.7)</td>
</tr>
<tr>
<td>CT</td>
<td>7</td>
<td>−3.1 (−1.4 to −9.9)</td>
<td>−7.9 (−10.9 to −6.5)</td>
</tr>
<tr>
<td>+VIPa0.2</td>
<td>7</td>
<td>−3.3 (−1.3 to −1.35)</td>
<td>−1.5 (−3.23 to 2.23)</td>
</tr>
<tr>
<td>LT</td>
<td>6</td>
<td>−12 (−4 to 2)</td>
<td>−0.5 (−3.5 to 3.5)</td>
</tr>
<tr>
<td>+VIPa0.2</td>
<td>7</td>
<td>−7 (−11.2 to −5.7)</td>
<td>−2 (−6.5 to −1.6)</td>
</tr>
<tr>
<td>+VIPa2</td>
<td>6</td>
<td>−1 (−2.5 to 2)</td>
<td>2 (1.3 to 3)</td>
</tr>
<tr>
<td>STa</td>
<td>5</td>
<td>−10 (−15 to −5.5)</td>
<td>−7 (−9.2 to −4)</td>
</tr>
<tr>
<td>+VIPa0.2</td>
<td>5</td>
<td>−3 (−4.5 to 0.5)</td>
<td>−2 (−3.2 to 0)</td>
</tr>
<tr>
<td>+VIPa2</td>
<td>5</td>
<td>−2.2 (−3.0 to 0.2)</td>
<td>−1.5 (−3.5 to 2)</td>
</tr>
<tr>
<td>VIP0.02</td>
<td>5</td>
<td>1.2 (0.5 to 2.1)</td>
<td>3.4 (2.2 to 5.1)</td>
</tr>
<tr>
<td>+VIPa0.2</td>
<td>5</td>
<td>2.6 (2.2 to 4)</td>
<td>6 (5 to 10.2)</td>
</tr>
<tr>
<td>+VIPa2</td>
<td>5</td>
<td>2.9 (2.2 to 3.8)</td>
<td>7.1 (6.5 to 9.8)</td>
</tr>
<tr>
<td>VIP0.2</td>
<td>5</td>
<td>−1 (−1.9 to 1.1)</td>
<td>0 (1 to 2.1)</td>
</tr>
<tr>
<td>+VIPa0.2</td>
<td>5</td>
<td>1.9 (1.3 to 2.2)</td>
<td>3.1 (2.5 to 4.5)</td>
</tr>
<tr>
<td>+VIPa2</td>
<td>5</td>
<td>3.2 (2.5 to 4.1)</td>
<td>6.3 (5.2 to 10)</td>
</tr>
</tbody>
</table>

Data are median (interquartile range).

VIPa0.2, VIPa2, VIP antagonist [4Cl-D-Phe⁶, Leu¹⁷] VIP at a dose of 0.2 (VIPa0.2) or 2 µg/kg/min (VIPa2).

††p<0.01, ‡‡p<0.05, and †p<0.05 compared with the corresponding toxin or VIP alone; **p<0.01 compared with VIP alone.

Figure 1: Effect of the VIP antagonist [4Cl-D-Phe⁶, Leu¹⁷] VIP given intravenously at a dose of 0.2 and 2 µg/kg/min (VIPA0.2 and VIPa2, respectively) on cholera toxin (CT) induced secretion. Results are expressed as median (interquartile range); positive values denote absorption and negative values denote secretion. **p<0.01 compared with controls; †† ††p<0.01 compared with CT and controls.

Figure 2: Effect of the VIP antagonist [4Cl-D-Phe⁶, Leu¹⁷] VIP given intravenously at a dose of 0.2 and 2 µg/kg/min (VIPa0.2 and VIPa2, respectively) on Escherichia coli heat labile toxin (LT) induced secretion. Results are expressed as median (interquartile range); positive values denote absorption and negative values denote secretion. **p<0.01 compared with controls; †† ††p<0.01 compared with LT and controls; †‡† †‡†p<0.01 compared with LT and p<0.05 compared with controls.

Figure 3: Effect of the VIP antagonist [4Cl-D-Phe⁶, Leu¹⁷] VIP given intravenously at a dose of 0.2 and 2 µg/kg/min (VIPa0.2 and VIPa2, respectively) on Escherichia coli heat stable toxin (STa) induced secretion. Results are expressed as median (interquartile range); positive values denote absorption and negative values denote secretion. **p<0.01 compared with controls; †† ††p<0.01 compared with STa and controls; †‡† †‡†p<0.01 compared with STa and p<0.05 compared with controls.

Discussion

The present experiments demonstrate that VIP plays an important role in intestinal fluid and electrolyte secretion induced by CT and E coli enterotoxins. It has previously been shown that activation of the enteric nervous system is cru-
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**p<0.01 compared with controls; †p<0.05 compared with µg/kg/min given intravenously. Results are expressed as induced decrease in fluid absorption at 0.02 and 0.2 µg/kg/min (VIPa0.2 and VIPa2, respectively) on VIP induced water absorption in the small intestine in response to secretomotor reflexes.  

In cholera, the secretagogue 5-HT has been identified as an indispensable factor in the pathophysiology of small intestinal secretion.  

Thus 5-HT released from enterochromaffin cells activate, among other receptors, neuronal (5-HT₃) receptors leading to intestinal secretion.  

However, whether or not activation is direct or via release of other neurotransmitters or activation of other neurones is not clear. Neurotransmitters, such as substance P, have been found to be recruited by CT implying that 5-HT is not the only transmitter involved in CT. In the present experiments, VIP antagonism dose dependently decreased fluid secretion induced by CT and partially reversed it, implying that VIP plays an important role in CT induced secretion but some other mediators may also be involved.

LT is a toxin structurally similar to CT and was thought to act via the same pathways. However, recent evidence suggests that these two toxins, although both stimulate adenylate cyclase, have totally different pathophysiological mechanisms. It has been shown definitively that 5-HT is not involved in LT induced secretion. In addition, blocking substance P receptors does not affect LT induced secretion. Hence, how does LT stimulate secretion? LT activates intracellular CAM but this does not explain inhibition of secretion by neuronal blockade. By binding to some receptor sites on enterocytes, it may release an unknown transmitter which activates the neuronal reflex. Alternatively, it has been postulated that the toxin may bind directly to enteric neurones and thus activate secretion. Although the afferent capsaicin sensitive neurones are stimulated during LT exposure, it is not known if LT binds directly to these neurones. Cholera toxin B subunit was found to bind to specific receptors exclusively on VIP containing neurones; however, there are no such data for LT. In the present experiments, we demonstrated that VIP plays an important role in this secretory process as VIP antagonism completely prevented LT induced secretion.

STa is structurally different from the two other toxins and induces intestinal secretion through an increase in cGMP. In addition, it has been demonstrated that, in common with CT and LT, the enteric nervous system is important in STa induced secretion as shown by the inhibitory effect of neuronal blockade. The neuronal mediators involved have not been fully elucidated. 5-HT was not found to be implicated in STa induced competitive inhibitors of VIP, in vivo studies have demonstrated that they can antagonise or reverse the secretory effect of VIP in the intestine. In our study, the validity and specificity of [4Cl-D-Phe⁶,Leu¹⁷]VIP as a VIP antagonist was tested by its effect on reversing the VIP induced decrease in jejunal fluid absorption and by its failure to reverse carbachol induced secretion. Moreover, there are no reports that show that VIPs crosses the blood-brain barrier. Accordingly, we assume that the effect of VIPs is peripheral, although a central effect cannot be excluded.

Whether VIP has a direct effect on inducing secretion in CT and in other toxins needs to be confirmed by examining the effect of VIP inhibition on the secretory effect of these toxins. The recent advent of VIP antagonists has permitted a better understanding of the importance of VIP in many physiological and pathophysiological states. While some in vitro studies failed to show these antagonists as
secretion. However, Rolfe and Levin have recently shown that nitric oxide synthase inhibition as well as capsaicin inhibit STa secretion in the rat ileum and concluded that STa activates a nitric oxide dependent myenteric plexus secretory reflex mediated by capsaicin sensitive C fibres. In addition, it seems that tachykinin NK2 receptors are activated to induce secretion during STa exposure. In the present experiments, VIP antagonism partially prevented STa induced secretion demonstrating that VIPergic neurones are activated during STa induced secretion. VIPergic neurones and nitric oxide synthase containing neurones coexist in the enteric nervous system. In addition, nitric oxide may release VIP from nerve terminals, and VIP may cause release of nitric oxide. Thus we may speculate that nitric oxide and VIP interact to induce secretion in STa. This speculation needs to be confirmed in further studies.

In conclusion, our findings support the view that VIP participates in cholera toxin and E. coli enterotoxin induced secretion, presumably through activation of an effenter limb of the secretory neuronal reflex. These findings may have clinical applications in the future when VIP antagonists for human use become available.

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