Failure of tropisetron to inhibit jejunal water and electrolyte secretion induced by 5-hydroxytryptamine in healthy volunteers

L K Munck, O Eskerod, M B Hansen, K Bukhave, J Rask-Madsen

Abstract
The effects of the 5-hydroxytryptamine (5-HT3) receptor antagonist, ICS 205-930 (tropisetron), on basal and 5-HT induced jejunal secretion of water and electrolytes were examined using a double blind, randomised crossover design. In seven healthy volunteers steady state perfusions of the proximal jejunum were performed twice with the Loc-I-Gut tube after 5+5 mg ICS 205-930 or placebo capsules were given. After equilibration for 60 minutes and completion of a 120 minute basal period 5-HT (10 μg/kg×min intravenously) was infused for 120 minutes. Net water absorption (mean (SEM)) in the basal period was 0.65 (0.04)-2.64 ml/cm×h and 0.67 (0.05) ml/cm×h after placebo and ICS 205-930, respectively (P<0.05). Infusion of 5-HT caused significant net secretion of water after placebo (2.05 (0.58) ml/cm×h; P<0.02) as well as ICS 205-930 (2.60 (0.89) ml/cm×h; P<0.05). As ICS 205-930 exerted no effects on either basal or 5-HT induced water and electrolyte transport in the intact human jejunum the compound is probably not efficacious as an anti-secretory drug in patients with 5-HT induced diarrhoea.

(Gut 1994; 35: 637–640)

Numerous in vivo studies have shown that exogenous 5-hydroxytryptamine (5-HT) elicits secretion of chloride and water in the small intestine of rodents,7,8 cats,9 dogs,10 and pigs (unpublished data) as well as in the human small intestine.11 Jejunal secretion elicited by bacterial toxins, such as cholera toxin12 and Escherichia coli heat stable enterotoxin,10 by Entamoeba histolytica,11 and by antigens in sensitised animals12,13 is mediated, in part least, by local release of 5-HT from enterochromaffine cells3 and immune cells, such as mast cells and phagocytes in the lamina propria.14 These cells play an important part in intestinal secretion through the release of prostaglandins15;16 by stimulation of the enterocyte12,17;18 or activation of neural reflex pathways.14,15

More specifically, 5-HT3 receptors have been implicated in the in vitro secretory response to both exogenous 5-HT16,17 and the 5-HT3 receptor antagonist, ICS 205-930 (tropisetron),17 which inhibits in a dose dependent manner part of the jejunal water secretion occurring in rodents in response to 5-HT19;20 choler toxin,19,20 and Escherichia coli heat stable enterotoxin.10 Furthermore, abnormally high plasma concentrations of 5-HT17 and luminal jejunal release of prostaglandin E219;20 are associated with profuse diarrhoea. Thus, case reports have provided evidence that 5-HT receptor antagonists, such as ICS 205-930,20 reduce the diarrhoeal volume in these patients, while high doses of ICS 205-930 alone acts as a secretagogue in the pig (unpublished data) and in combination with ketanserin increases choler toxin induced jejunal water secretion in humans (unpublished data).

The purpose of this study was to examine the effects of ICS 205-930 on basal and 5-HT induced intestinal water and electrolyte transport in the human small intestine. Steady state perfusions of jejunum were performed in healthy volunteers using a randomised, double blind, crossover design.

PARTICIPANTS AND SAFETY MEASUREMENTS
Seven healthy male volunteers (median age 22 years, range 20–26) not receiving any treatment consented to the study protocol, which was approved by the regional ethical committee and the Danish National Health Service. All subjects were seen before inclusion for clinical assessment, electrocardiography, and laboratory screening (haemoglobin and packed cell volume; red cell, platelet and, leucocyte counts; plasma concentrations of creatine, sodium, and potassium; and activities of aspartate aminotransferase and alkaline phosphatase.) Laboratory tests were repeated at the end of each perfusion.

EXPERIMENTAL DESIGN
The study was carried out in a randomised, double blind, crossover design. Each participant had two jejunal perfusions with an interval of at least one week. Either placebo capsules or 5 mg capsules of ICS 205-930 were given orally 12 hours and one hour before jejunal intubation.

SEGMENTAL JEJUNAL PERFUSION
After an overnight fast the Loc-I-Gut tube (Kabi Pharmacia, Uppsala, Sweden) was inserted orally.20 A guidewire was used for intubation and the position of the tube controlled by fluoroscopy. Proximal to a distal tungsten weight two inflatable latex balloons were placed 10 cm apart. The proximal balloon was placed at the ligament of Treitz and both balloons inflated with 35–40 ml of air. These volumes were adjusted during the perfusion and always at the start of 5-HT infusion to minimise leakage from the closed study segment. The volumes were, therefore, kept as high as possible, that is 1–2 ml less than the volume that elicited abdominal discomfort.

The closed jejunal segment was perfused at 3 ml/min (LKB 2115 Multiperpex Pump,
Bromma, Sweden). The composition in mmol/l of the isotonic perfusion solution was: Na\(^+\) 145, K\(^+\) 5, Cl\(^-\) 140, and HCO\(_3\)\(^-\) 10. \(^{51}\)Cr-EDTA (15 \(\mu\)Ci/l) was added as a non-absorbable marker.

Aspiration was by gravity drainage from ports next to the balloons. The aspirates were collected in 20 minute periods and immediately frozen to \(-20^\circ\)C until analysis. The volume of the study segment was about 80 ml.

A second double lumen tube was positioned in the antrum of the stomach for aspiration of gastric juice. Phenol red (50 mg/l; 154 mM NaCl) was infused at a rate of 1 ml/min into the stomach and analysed in the effluents from the closed jejunal segment to rule out leakage. After a 60 minute equilibration period, during which the closed jejunal segment was emptied for mucus, as judged from the appearance of clear aspirates, samples were collected in consecutive 20 minute periods. Both during the equilibration period and the control period (120 minutes) isotonic saline was infused intravenously at a rate of 50 ml/h, when the infusion solution was changed to 5-HT (5-hydroxytryptamine creatinine sulphate (Serva, Heidelberg, Germany) dissolved in isotonic saline (10 \(\mu\)g/kg\(\times\)min, 50 ml/h intravenously) for 120 minutes.

**ANALYTICAL PROCEDURES**

Sodium and potassium were determined by flame photometry (FLM 2, Radiometer, Copenhagen, Denmark) and chloride was measured in a chloride metre (CMT 10, Radiometer). \(^{51}\)Cr activity was measured by gammaspectrometry (model 1185, Searle Nuclear Chicago Division, Chicago, USA). Phenol red was measured spectrophotometrically at 560 nm after alkalisation (pH 11) with a Na\(_2\)PO\(_4\) buffer.

**DETERMINATION OF WATER AND ELECTROLYTE TRANSPORT**

Net water and electrolyte transport rates were determined as described by Knutson et al\(^a\) and expressed as ml/cm\(\times\)h and mmol/cm\(\times\)h, respectively. Net absorption was shown by a positive sign and net secretion by a negative sign.

**STATISTICAL ANALYSES**

All results are given as means with standard errors (SEM). To discover if analysis of variance could be applied the results were tested for carryover and period effects on treatment outcomes using Student’s \(t\) test. This was done on the assumption that there were no residual effects. Pairs of group means were tested by one way analysis of variance and Duncan’s multiple range test. Probability values less than 0·05 (two sided test) were considered significant.

**Results**

**DESIGN**

According to the protocol, active and placebo periods should be defined in each subject by the investigators from the change, if any, in net water transport rates calculated before breaking the code. Active or placebo periods were correctly defined only in four cases, suggesting that ICS 205-930 had no clinically relevant influence on water transport. Pairs of group means were tested by one way analysis of variance, because no statistically significant carryover effect (\(p>0\cdot05\)) or period effect (\(p>0\cdot01\)) was detected. The contamination by phenol red did not exceed 0·1% in any aspirate analysed. The recovery of \(^{51}\)Cr-EDTA was 99·7% (0·8%) (\(n=16\)), showing that the test segment was effectively isolated.

**NET WATER AND ELECTROLYTE TRANSPORT**

Net water transport rates (Figs 1 and 2) were insignificantly (\(p>0\cdot05\)) different from zero during basal periods, both after placebo (0·55 (0·84) ml/cm\(\times\)h) and ICS 205-930 (0·74 (0·72) ml/cm\(\times\)h) was given. Subsequently, pretreatment with ICS 205-930 caused no change in basal net water absorption compared with placebo (Figs 1 and 2). Net secretion of water in response to 5-HT infusion was seen in all subjects who had received placebo capsules (2·05 (0·58) ml/cm\(\times\)h; \(p<0\cdot02\)) and in all but one subject after ICS 205-930 capsules (2·60 (0·89) ml/cm\(\times\)h; \(p<0\cdot05\)) (Figs 1 and 2), but there was no difference (\(p>0\cdot05\)) in secretory response between active and inactive treatment.

Also net absorption of sodium, potassium, and chloride was insignificantly (\(p>0\cdot05\)) different from zero during basal periods after placebo (18·3 (2·46), 1·1 (0·7), and 5·5 (19·3) mmol/cm\(\times\)h) and ICS 205-930 (21·1 (16·2), 1·1 (0·5), and 11·7 (15·3) mmol/cm\(\times\)h), respectively. The 5-HT induced secretion of sodium, potassium,
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and chloride was not changed (p>0.05) by ICS 205-930 (Fig 3).

ADVERSE EVENTS

No adverse events from treatment with ICS 205-930 or placebo were reported and no abnormalities or significant differences in laboratory screening values were seen before or after the study, respectively. The intravenous infusion of 5-HT caused pain at the infusion site and vasodilation. Minor oedema of the hands and flushing were seen in two cases. Most subjects reported nausea and abdominal discomfort, which was relieved by withdrawing a few ml of air from the balloons. All symptoms disappeared immediately after the 5-HT infusion was stopped. No effects on blood pressure or heart rate were registered.

Discussion

Intestinal water and electrolyte transport is controlled by the enteric nervous system, comprising intrinsic and extrinsic neurons, in addition to neuroendocrine cells, hormones, bacterial enterotoxins, and the immune system, all of which are anatomically and functionally intermingled. The neurotransmitter, 5-HT, is present predominantly in the enterochromaffine cells along the entire small intestine, but a minor part is found in the myenteric plexus and in the mast cells situated in the lamina propria. The 5-HT induced, calcium dependent secretion is the result of inhibition of electroneutral sodium chloride absorption and stimulation of chloride secretion and is independent of changes in intestinal blood flow or permeability.

In this study we used a glucose free perfusion solution to permit comparison with previously published data on 5-HT induced jejunal secretion in humans. The results confirm that exogenous 5-HT induces jejunal secretion of water and electrolytes, as previously shown in vivo in various experimental animal models and in humans. The results also show that 10 mg of ICS 205-930 (tropisetron) given orally has no effect on either basal water and electrolyte transport or 5-HT induced secretion. In a model of human secretory diarrhoea, in which 6.25 μg cholera toxin given intrajejunally induced a secretory response of the same magnitude as seen in our study, ketanserin given parenterally and ICS 205-930 in combination enhanced the secretory effects of cholera toxin, while ICS 205-930 had no effect on basal water transport (unpublished data). In addition, ICS 205-930 failed to inhibit the secretory effects of cholera toxin in the pig small intestine in vivo and induced secretion at high rates (unpublished data).

Our results as well as those of others seen in pigs and in humans are not comparable with those obtained in vivo in rodents, because higher doses of the 5-HT receptor antagonists were given intravenously in the latter studies. In this study ICS 205-930 was given 12 hours and one hour before the experiment to ensure absorption and stable plasma concentrations of the drug. The dose is equivalent to that given intravenously by Eherer et al (unpublished data) who saw no effect on net water absorption during control periods and, contrary with what has been seen in rodent experiments, an enhancement rather than inhibition of cholera toxin induced secretion by ICS 705-930. Thus, the lack of effect of basal and 5-HT stimulated water and electrolyte transport is probably not explained by insufficient absorption of the drug. The optimum therapeutic dose for prevention of nausea and vomiting associated with chemotherapy is 5 mg. It should also be noted that the risk of overlooking a true effect of ICS 205-930...
on net water transport — that is, type II error, exceeding 1.5 SD was less than 20%. Thus, the species differences seen between rodents and humans are probably not explained by differences in dose or route of administration.

Net absorption of water and electrolytes is the most common finding in perfusion studies of the human small intestine using the triple lumen technique or the double lumen technique with a proximal occluding balloon (unpublished data). In this study net water absorption was minimal and insignificantly different from zero (p > 0.05).

Minimal net secretion of water occurred in several subjects during basal conditions, both after placebo and ICS 205-930 (Fig 1). This discrepancy between our previous study using the same perfusion solution, but the triple lumen technique, cannot be explained by the absence of glucose in the perfusate. It might be the result, however, of jejunal distension by occluding balloons, which has been shown to elicit net chloride secretion in rat colon. On the other hand, distension of the normal human jejunum with a balloon did not interfere with intrinsic myoelectrical activity. As the perfusions were performed during 'steady state' conditions any influence of the distended balloons on basal intestinal secretion probably will not have obscured an anti-secretory effect of ICS 205-930, although it might have decreased the response seen with 5-HT.

Presently, we cannot explain the conflicting results of ICS 205-930 on water and electrolyte transport in the small intestine of rodents and humans. The results of this study emphasise the risk of drawing conclusions concerning drug actions in humans from findings made in rodent studies, but suggest that the investigational drug, ICS 205-930, is probably not efficacious as an anti-secretory drug in patients with diarrhea.

This work was supported by a research grant from Sandoz Ltd., who also kindly provided ICS 205-930 and placebo capsules. Part of the work has been presented at XXXXIVnd International Congress of the International Union of Physiological Sciences in Glasgow, August 1991.

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Gut 1994 35: 637-640
doi: 10.1136/gut.35.5.637

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