Effects of serotonin on rat ileocolonic transit and fluid transfer in vivo: possible mechanisms of action

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Abstract
The aim was to investigate the action of serotonin (5HT) on function of the ileocolonic junction (ICJ) in vivo. In anaesthetised rats, models were developed to study the effects of intraperitoneal (ip) serotonin on ileocolonic and colonic transit, and the effects on transit of a number of 5HT receptor antagonists. In the first series of experiments, a bolus of saline labelled with 99mTc DTPA was instilled 20 cm proximal to the ICJ and transit was assessed three hours later by the geometric centre of the spread of isotope. In the second series, similar techniques were used on the postcaecal colon and transit assessed two hours later. In the third series of experiments, the effects of ip 5HT on ileal net fluid flux was evaluated by standard perfusion experiments with 3H-labelled polyethylene glycol (PEG) 4000 as a non-absorbable marker in rat plasma-like electrolyte solution. Compared with ip saline, 5HT accelerated ICJ transit significantly (p<0.05). This acceleration was comparable with the effect of ip bethanecol. The effects of 5HT on ICJ transit were inhibited by the intraperitoneal (ip) infusion of atropine, the 5HT receptor antagonists, methysergide, ketanserin, zacopride, and the 5HT4 agonist, 5CS3116. Methysergide, zacopride, and 5CS3116 given with ip 5HT slowed ICJ transit to rates below those of ip saline alone. When these same agents were given together with ip saline, the ICJ transit was not significantly altered. Serotonin, at the dose that accelerated ICJ transit, did not significantly alter transit in post-ileal fluid transport. In conclusion, 5HT is a potent pharmacological stimulant of transit across the rat ICJ in vivo; the action of 5HT is mediated partly through muscarinic neurones and several 5HT receptor subtypes.

Serotonin (5 hydroxytryptamine, 5HT) is believed to be an important humoral modulator of carcinoid diarrhoea. Agents that release 5HT exacerbate diarrhoea in the carcinoid syndrome; agents that block the synthesis or the peripheral effects of 5HT sometimes control diarrhoea in patients with carcinoid syndrome. Serotonin stimulates contraction or myoelectric spike activity of small intestinal muscle in vivo, and gastrointestinal transit is altered in patients with the carcinoid syndrome. Serotonin also stimulates gastric emptying in rats in vivo; however, the specific effects of 5HT on propulsion and transit in vivo are less clear.

In an in vitro study of an isolated segment of guinea pig ileum, Bulbring and Lin found that 5HT stimulated propulsive motility when applied intraluminally. By contrast, serosal 5HT had no effect on transit; others have even reported inhibition of peristalsis by 5HT. Serotonin stimulates nerve muscle preparations from the guinea pig small intestine and colon in vitro. Thus there is evidence that 5HT can both inhibit and facilitate gastrointestinal motor function.

Intestinal smooth muscle cells and enteric nerves have 5HT1, 5HT2, and 5HT3 receptors. More recently, a new class of 5HT4 receptors has been found in several tissues, including the guinea pig ileum and ascending colon, the cardiovascular system of the pig, and the human heart. Acetylcholine is thought to play an important part in mediating 5HT induced contraction of intestinal smooth muscle in vivo and in vivo. The relative roles of the different subtypes of 5HT receptors inducing propulsion through different levels of the alimentary tract are unclear. Further understanding of serotonergic control of motility could lead to new ways to control diarrhoea in diseases such as the carcinoid syndrome.

The ileocolonic junction (ICJ) is an important area whereby gastrointestinal absorptive and motor functions are controlled, by virtue of its regulation of the passage of chyme from the small to the large bowel. In carcinoid diarrhoea, transit through the small bowel and proximal colon is considerably accelerated, and this acceleration is associated with high urinary concentrations of 5-hydroxyindole-acetic acid (5HIAA) and circulating 5HT. In view of the rapid ICJ transit in patients with carcinoid diarrhoea, our aim was to develop a model in which the effects and mechanisms of hypersero-toninaemia on transit and fluid transport in this region could be investigated.

Materials and methods
Experiments were performed on 100 adult male Sprague-Dawley rats weighing 300–400 mg; the study was approved by the institutional animal care and use committee and the radiation control committee of the Mayo Clinic. Animals were housed in an animal care facility that is approved by the American Association for Accreditation of Laboratory Animal Care and they were exposed to 12 hours of alternating light and dark cycles. Their health was monitored by fully trained veterinary staff.

EXPERIMENTAL PROCEDURES
Preparation of experimental models
Rats were anaesthetised with intraperitoneal (ip) chloral hydrate (300 mg/kg). In all animals, a
Effects on transit were counted in Tygon of means minutes; the throughput 99mTc DTPA Colonic transit calibrator by ligatures segment after transit Ileocolonic method of beyond dissemination. Preliminary segment. five volume was returned was set up, segments 20 cm. The was postcaecal placed in the caecum, the distal ileum, and the proximal colon that was accessible at the time of laparotomy. The average length of the colonic loop studied was 20 cm. In a third series of experiments (fluid transport), the distal 20 to 25 cm of ileum was isolated for perfusion studies by two Tygon cannulae. After one of these three experimental segments was set up, the intestines were returned to the abdominal cavity and the incision was closed.

Ileocolonic transit
One ml of 99mTc DTPA radio labelled saline was introduced into the ileum by slow infusion over five minutes; 0·3 ml saline was gently flushed through the cannula to ensure delivery of the entire volume of radioisotope into the ileum. Preliminary studies showed that, with this method of delivery, the radiolabel did not disperse beyond the first 5 cm of the ileal segment. Animals were kept under light anaesthesia thereafter with ip chloral hydrate. Three hours after the ileal infusion, the abdomen was opened and the mesenteric vessels were tied. The ileocolonic segment was identified, closed by ligatures at each end, and divided into five equal portions by separating 4 cm segments by means of silk ligatures. The caecum and colon were identified down to the pelvis and ligatures tied. The rats were then killed, and all these segments were counted in a Capintech radioisotope dose calibrator (CRC-5, Montvale, NJ).

Colonic transit
With the animals under light anaesthesia, 1 ml of 99mTc DTPA radio labelled saline was introduced into the colon by slow infusion over five minutes; 0·3 ml of saline was gently flushed through the cannula. Two hours after this infusion, the abdomen was opened, and the loop of colon was divided into four equal segments by means of silk ligatures. The effluent from the distal Tygon cannula was collected separately during the experiment, and constituted the fifth segment. Each segment of colon and the effluent were counted in a dose calibrator.

Ileal fluid transport
The ileum was perfused through the proximal cannula with an iso-osmolar electrolyte solution containing 14C PEG 4000. The electrolyte content (Na+, 151 mmol/l; K+, 5·6 mmol/l; Cl-, 128 mmol/l; Ca++, 2·09 mmol/l; HCO3-, 23·1 mmol/l) was selected to mimic rat plasma (osmolarity: 324 mOsm/kg; pH, 7·4). The ileal loop was perfused at a rate of 1·0 ml/min by a peristaltic pump (Watson-Marlow Inc, Marblehead, MA); experiments started with 30 minutes equilibration followed by three hours during which samples of ileal effluent were collected every 10 minutes. During the first and third hours, saline was infused into the aorta; during the second hour, 5HT was infused. Net fluid flux was assessed by changes in the concentration of the non-absorbable marker; 14C PEG 4000.

Choice of agonist and antagonist doses
The dose of 5HT (Sigma Chemical Co, St Louis, MO) used in this study (7·74 mmol/kg) was based on previous in vivo dose response studies that showed that 5HT stimulated gastric emptying.3 Antagonist doses were based on previous publications.8,9 In the first series of experiments on motor function of the ileocolonic junction, we studied the effects of ia saline, 5HT, and bethanechol chloride, at a dose of 3·05 mmol/kg (Merck, Sharp, and Dohme, West Point, PA). These agents were given in a total volume of 1·5 ml, a 0·5 ml bolus injection, followed by a continuous infusion of 1 ml over a one hour period.

In experiments to study the mechanism of 5HT induced alterations of ileocolonic function, atropine and several 5HT receptor subtype antagonists were given intraperitoneally 60 minutes before the laparotomy; these were atropine sulphate (0·14 mmol/kg, Elkins-Sinn, Cherry Hill, NJ); methysergide (2·83 mmol/kg, a 5HT1 and 5HT2 antagonist donated by Sandoz Research Institute, East Hanover, NJ, USA); ketanserin (1·83 mmol/kg, a 5HT2 antagonist donated by Janssen Pharmaceutica Inc, Beerse, Belgium); R-zacopride (2·5 mmol/kg, a selective 5HT3 antagonist donated by Searle, IL, USA), and SC53116 (2·4 mmol/kg, a 5HT4 agonist donated by Searle, IL, USA).

Experimental design
Five series of experiments were performed in rats in vivo to study: (1) the effects of ia saline, 5HT, and bethanechol on ileocolonic transit; (2) the effects of ip atropine, a 5HT4 agonist, and several 5HT antagonists on ileocolonic transit induced by ia 5HT; (3) the effect of the same ip 5HT agonist and antagonists on ileocolonic transit; (4) the effect of ia saline, 5HT, and bethanechol on colonic transit; and (5) a comparison between the effects of ia saline and ia 5HT on ileal net fluid flux.

DATA AND STATISTICAL ANALYSIS
The proportions of counts in the ileal or colonic segments at the end of the experiment were calculated. 1CJ transit was assessed by finding
the geometric centre (weighted average) of
counts over the five ileal segments, caecum, and
colon with an adaptation of a previously pub-
lished method used to measure intestinal transit
in rats.\(^\text{15}\) For studies of ICJ transit, with this
method, a value of 1 indicated that all the isotope
was in the first 4 cm of ileum; a value of 6
indicated that all isotope was in the caecum or
colon. Colonic transit was similarly assessed with
the four colonic segments and effluent as the fifth
segment.

Statistical analysis compared the effects on
transit of 5HT and its agonist and antagonists by
ANOVA followed by Dunnett’s multigroup
comparison, with the ia saline or the ia 5HT
alone as the control groups. Effects of ip agents
during ia saline were also compared by ANOVA
with Newman-Kuels multigroup comparison.
All results are expressed as mean (SEM).

Net fluid flux was summarised for the ia saline
and 5HT hours and expressed as \(\mu\)l/cm loop/
hour in each rat. Statistical analysis was by
paired \(t\) test.

Results

ILEOCOLONIC JUNCTION TRANSIT: EFFECTS OF 5HT
AND ITS ANTAGONISTS

Serotonin and bethanechol (Fig 1) significantly
accelerated ICJ transit compared with saline.

**Table 1.** Ileocolonic transit: effect of combined intra-aortic
serotonin (5HT) and intraperitoneal 5HT agonist (SC53116)
or 5HT antagonists

<table>
<thead>
<tr>
<th>Agents</th>
<th>(n)</th>
<th>Geometric centre (mean (SEM))</th>
<th>(p^*)</th>
<th>(p^{*})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saline (ia)</td>
<td>5</td>
<td>3.1 (0.2)</td>
<td>&lt;0.05</td>
<td></td>
</tr>
<tr>
<td>5HT alone (ia);</td>
<td>5</td>
<td>4.3 (0.1)</td>
<td>&lt;0.05</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>+ Atropine (ip)</td>
<td>5</td>
<td>2.4 (0.2)</td>
<td>&lt;0.05</td>
<td>NS</td>
</tr>
<tr>
<td>+ Methysergide (ip)</td>
<td>5</td>
<td>1.9 (0.2)</td>
<td>&lt;0.05</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>+ Ketanserin (ip)</td>
<td>5</td>
<td>2.4 (0.2)</td>
<td>&lt;0.05</td>
<td>NS</td>
</tr>
<tr>
<td>+ R-Zacopride (ip)</td>
<td>5</td>
<td>1.6 (0.1)</td>
<td>&lt;0.05</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>+ SC53116 (ip)</td>
<td>5</td>
<td>1.9 (0.1)</td>
<td>&lt;0.05</td>
<td>&lt;0.05</td>
</tr>
</tbody>
</table>

\(\ast\)Significance compared with 5HT ia without antagonists.
\(\dagger\)Significance compared with ia saline.
ia=Intra-aortic; ip=intraperitoneal.

**Table 2.** Ileocolonic transit: effects of serotonin (5HT)
antagonists and 5HT agonist (SC53116)

<table>
<thead>
<tr>
<th>ip Agents</th>
<th>(n)</th>
<th>Geometric centre (mean (SEM))</th>
<th>(p) v ip Saline</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saline</td>
<td>6</td>
<td>3.0 (0.2)</td>
<td>NS</td>
</tr>
<tr>
<td>Methysergide</td>
<td>6</td>
<td>2.4 (0.4)</td>
<td>NS</td>
</tr>
<tr>
<td>Ketanserin</td>
<td>6</td>
<td>2.8 (0.4)</td>
<td>NS</td>
</tr>
<tr>
<td>R-Zacopride</td>
<td>6</td>
<td>3.0 (0.3)</td>
<td>NS(^*)</td>
</tr>
<tr>
<td>SC53116</td>
<td>6</td>
<td>2.0 (0.3)</td>
<td>NS(^*)</td>
</tr>
</tbody>
</table>

\(\ast\)Significance for comparison of R-Zacopride v SC53116.
\(\dagger\)All animals received a background ia infusion of saline.
Abbreviations as for Table 1.

The effects of 5HT and bethanechol were not
statistically different.

Table 1 shows the geometric centres of isotope
in the ileocolonic segment after the concomitant
5HT and the series of ip drugs were given.
Methysergide, R-zacopride, and SC53116
significantly inhibited the increased transit
induced by 5HT; indeed, the geometric centres of
isotope with ia 5HT and these agents given ip
were less than the value observed with ia saline
alone (Table 1). Atropine and the 5HT2
antagonist, ketanserin, also significantly inhibited
5HT induced ICJ transit, but not to concentrations
below that of ia saline.

**EFFECT OF INTRAPERITONEAL SC53116 AND 5HT
ANTAGONISTS ON BASAL ILEOCOLONIC JUNCTION
TRANSIT**

None of the 5HT antagonists or the 5HT2
agonist, SC53116, significantly altered ICJ
transit when they were given ip during ia
infusion of saline (Table II). There were no
significant differences relative to ip saline, but
the difference between groups treated with
SC53116 and R-zacopride was significant (Newman-Kuels test).

**EFFECT OF 5HT AND BETHANECHOL ON COLONIC
TRANSIT**

Intra-aortic bethanechol significantly accelerated
colonic transit. The effect of ia serotonin was
variable; however, there was no overall acceleration
of colonic transit (Fig 2).

**EFFECT OF 5HT ON ILEAL FLUID FLUX**

Intra-aortic 5HT, given at the same dose and
route as had induced acceleration of ICJ transit,
failed to significantly alter baseline absorption
found during the ia infusion of saline (Fig 3).

Discussion

The questions we considered required that an in
vivo model be developed for measurement of
transit across the ileocolonic junction and colon.
We elected to use a static rather than a dynamic
method to evaluate transit because our previous
studies with dynamic scintigraphy\(^\text{16}\) showed that
emptying of the unprepared rat ileum into the
caecum occurred very infrequently. Infusion of
the liquid bolus at a slow rate ensured that the

![Figure 1: Geometric centre of isotope in the ileocolonic segment three hours after intra-aortic saline, 5HT, or bethanechol. Results are shown as mean (SEM).](Image)

![Figure 2: Geometric centre of isotope in the colon, two hours after intra-aortic saline, 5HT, or bethanechol. Note that bethanechol stimulates colonic transit, but 5HT does not.](Image)
Figure 3: Net fluid ideal flow during infusion of saline and intra-aortic SHT. Note the lack of effect of intra-aortic SHT (total dose 7-14 mmol/kg).

Absorption (µL/cm²/h)

Saline (before) 0 20
Saline (after)
Serotonin

isotope did not disperse beyond the first ileal segment; the present results confirmed the effect of a cholinergic agonist and antagonist on transit in the ICJ and colon, suggesting that the model is valid.

Serotonin accelerated transit of liquid across the ICJ, but not in the colon in vivo. Previous reports have shown in vivo effects of 5HT on intraluminal pressure and myoelectric activity of the ICJ in the cat, but they did not assess the propulsive effects of 5HT in vivo or analyse which subtypes of 5HT receptors were responsible. The effects of serotonin on gastrointestinal motor function are generally believed to be mediated through myenteric plexus neurons. Although the mucosal application of 5HT activated the peristaltic reflex, suggesting a possible role for mucosal enterochromaffin cells, Gershon and Tamir provided convincing data that an intramural barrier separates the myenteric plexus from the enterochromaffin cells in the epithelium. It is possible, however, that mucosal application of 5HT might stimulate submucosal 5HT neurons, which may in turn activate interneurons connecting the submucosal and enteric plexuses.

Effects of 5HT on colonic motor function were previously studied in vivo by Jacoby et al. Intracerebroventricular 5HT inhibited propulsive motility in the mouse colon; by contrast, ip 5HT stimulated propulsion of a glass bead through the colon. In vitro, 5HT stimulated rapid colonic circular muscle but inhibited neurons responsible for the off contraction. Others have also shown in the rat that high doses (10⁻⁵ M) of 5HT relaxed the caecum and internal anal sphincter. Our data suggest that 5HT had no significant physiological effect on transit of liquid through the rat colon and are, therefore, consistent with these two studies.

The availability of novel, relatively specific antagonists of the 5HT receptor subtypes allowed us to evaluate the mechanism of 5HT induced acceleration of ICJ transit. Generally, our results confirm some of the findings from in vitro studies. We showed that the effects of 5HT on ICJ transit are partially mediated by cholinergic mechanisms and that they involve several 5HT receptor subtypes. Thus 5HT stimulated an excitatory muscarinic cholinergic pathway that was inhibited by atropine. This is consistent with the previously described role of acetylcholine in 5HT induced contraction of intestinal smooth muscle in vitro and in vivo, which is mediated through neuronal 5HT₁ receptors. Our findings of inhibition of the 5HT effect by subtypes 1 and 3 antagonists and by a 5HT₄ agonist suggests more complex interactions of 5HT receptors on enteric muscles or nerves. 5HT may be directly stimulating smooth muscle or antagonising an inhibitory interneuron in the myenteric plexus. Several motor effects of 5HT agents have been reported. For example, 5HT₁I receptors occur on enteric smooth muscle, and their stimulation resulted in smooth muscle contraction. Receptors for 5HT₁I may alter pyloric and caecal function. Others have identified 5HT₄ receptors in neurons of guinea pig ileum and ascending colon; the inhibitory effects of the 5HT₄ agonist, 5CS11, 5HT induced ICJ transit in the present study suggest that there may be a 5HT₄ mediated stimulation of the ileocaecal sphincter itself. The precise sites of action of 5HT were not considered in our experiments, and further studies on isolated tissues will be necessary.

Our findings help elucidate the in vivo effects of 5HT, such as in patients with carcinoid diarrhoea who have rapid small bowel and colonic transit and decreased capacitance of the proximal colon. Thus correction of these motor dysfunctions may help control carcinoid diarrhoea. Previous studies in a few patients suggest that carcinoid diarrhoea is, at least temporarily, improved with methysergide, ICS205-930 (a 5HT₁ antagonist without 5HT₃ effects), and ketanserin. Intracolonic 5HT, given at the same dose that stimulated ICJ transit, had no significant effect on net fluid flux in the rat ileum. These findings contrast with those of Beubler et al who used close intra-arterial injections of 5HT. In the absence of dose-response studies, the only important point we can make is that transit was accelerated across the ICJ by a dose of ip 5HT that did not induce ileal secretion in vivo. It is possible that secretory and motor mechanisms contribute to carcinoid diarrhoea, and the effects of 5HT and its antagonists on both processes deserve further study.

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