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Non-neuronal release of ACh plays a key role in secretory response to luminal

propionate in rat colon

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Non-technical summary

ACh is the best characterised neurotransmitter that is synthesized in cholinergic neurons in the brain and gut wall. In the gut, acetylcholine is released from the nerve endings in response to luminal stimuli and regulates the movement of gut contents via stimulating muscle contraction and epithelial ions secretion. We show that acetylcholine is synthesized in colonic epithelial cells and released into the serosal side by luminal chemical stimulation of short chain fatty acid, propionate, and causes chloride secretion. These results suggest that non-neuronal release of acetylcholine in response to luminal stimuli plays a role in colonic chloride secretion.

Abstract

Colonic chloride secretion is induced by chemical stimuli via the enteric nervous reflex. We have previously demonstrated that propionate stimulates chloride secretion via sensory and cholinergic systems of the mucosa in rat distal colon. In this study, we demonstrate non-neuronal release of ACh in the secretory response to propionate using an Ussing chamber. Mucosa preparations from the colon, not including the myenteric and submucosal plexuses, were used. Luminal addition of propionate and serosal addition of ACh caused biphasic changes in short-circuit current (I_{sc}). TTX (1 μ M) had no effects, while atropine (10 μ M) significantly inhibited the I_{sc} response to propionate and abolished that to ACh. In response to luminal propionate stimulation, ACh was released into serosal fluid. A linear relationship was observed between the maximal increase in I_{sc} and the amounts of ACh released 5 min after propionate stimulation. This ACh releases induced by propionate were not affected by atropine and bumetanide,

although both drugs significantly reduced the I_{sc} responses to propionate. Luminal addition of 3-Cl-propionate, an inactive analogue of propionate, abolished both ACh release and I_{sc} response produced by propionate. RT-PCR analysis indicated that isolated crypt cells from the distal colon expressed enzyme of ACh synthesis (ChAT) and transporters of organic cation (OCTs), but not neuronal CHT1 and VAChT. The isolated crypt cells contained comparable amounts of ACh to the residual muscle tissues including nerve plexuses. In conclusion, the non-neuronal release of ACh from colonocytes coupled with propionate stimulation plays a key role in chloride secretion, via the paracrine action of ACh on muscarinic receptors of colonocytes.

Abbreviations: ChAT, choline acetyltransferase; CHT1, high affinity choline transporter; OCT, organic cation transporter; SCFA, short chain fatty acid; VAChT, vesicular acetylcholine transporter

Introduction

Chemical stimulation of the intestinal mucosa can elicit secretory and contractile responses via sensory input to receptor cells and enteric nervous reflex (Hubel, 1985). Nutrients sensing regulates digestion, absorption and motility, while noxious stimuli evoke defence functions that flush out harmful microbes and toxins (Furness *et al.*, 2004).

Short-chain fatty acids (SCFAs), which are major products of microbial fermentation in the large intestine, not only act as energy sources (Livesey & Elia, 1995) but also as chemical stimulators for mesenteric vasodilatation (Knock *et al.*, 2002), colonic contraction (Yajima, 1985; Mitsui *et al.*, 2005) and peristalsis (Grider &

Piland, 2007), mucin secretion (Sakata & Setoyama, 1995), and chloride secretion (Yajima, 1988). The secretory response to SCFAs, in combination with the contractile response, seems to act as a lubricant for the movement of luminal content in the colon.

We previously demonstrated that luminal addition of propionate or other SCFAs transiently stimulated chloride secretion and resulted in an increase in short-circuit current (I_{sc}) and conductance in rat distal colon *in vitro* (Yajima, 1988). Hubel and Russ confirmed the I_{sc} responses to luminal propionate in rat distal colon and further showed that the propionate responses were not affected by tachyphylaxis to various transmitters localized in the intestinal wall, calcitonin gene-related peptide, 5-HT, histamine, neurotensin and substance-P (Hubel & Russ, 1993), whereas atropine (10 μ M) and local anesthetics (50–100 μ M) reduced the propionate responses by 81–90% and 76–82%, respectively (Yajima, 1988; Hubel & Russ, 1993). Furthermore, superficial mucosal damage with hypertonic Na sulphate (2 M) or xylose (4.5 M) reduced the propionate response by 90% and 86%, respectively (Hubel & Russ, 1993). Taken together, it could be speculated that ACh may be released from cholinergic secretomotor neurons or colonic epithelial cells; however, the exact site of ACh release was not fully revealed.

ACh is the best characterised neurotransmitter that is synthesized in cholinergic neurons and released via vesicular machinery in response to physiological and pharmacological stimulation. ACh acts through nicotinic and muscarinic receptors in nerves and peripheral tissues. The digestive tract is richly innervated by cholinergic neuron (Harrington *et al.*, 2010a). Epithelial cells and muscles express many subtypes of muscarinic receptors that are involved in the reflex motor and secretory responses to mechanical and chemical signals from luminal side of the digestive tract (Raybould *et al.*, 2004).

Besides neuronal ACh, the synthesis and storage of ACh in a broad variety of non-neuronal cells, particularly in surface epithelia of the airway, bladder, placenta and skin of rodents and humans has been recently discovered (Wessler & Kirkpatrick, 2008). The presence of non-neuronal ACh is also reported in the epithelial cells of the small and large intestines of rats and humans (Klapproth *et al.*, 1997). Non-neuronal ACh is physiologically expected to act as a local cellular signaling or trophic molecule (Klapproth *et al.*, 1997). However, the physiological significance of the synthesis and release of non-neuronal ACh in the intestine are still unclear, although there are a few reports regarding changes in the epithelial expression of ChAT in inflammatory bowel diseases, including ulcerative colitis (Gareau *et al.*, 2007; Jonsson *et al.*, 2007).

In the present study, we examined the ACh release from the mucosa preparation of the colon, not including the myenteric and submucosal plexuses, in response to the luminal addition of propionate, and non-neuronal ACh storage in colonic crypt cells of rat. We have demonstrated for the first time that luminal propionate stimulation released non-neuronal ACh from colonic epithelial cells into the serosal side, and then induced chloride secretion, probably via paracrine action on muscarinic receptors in colonocytes in rat.

Materials and Methods

Animals

This study was approved by the Hokkaido University Animal Committee, and all animals were maintained in accordance with the Hokkaido University guidelines for the care and use of laboratory animals. Male Sprague-Dawley rats (250–300 g) were used. They were fed a pelleted diet (Type MF, Oriental Yeast Co., Tokyo, Japan) *ad libitum*

with free access to water, but were starved overnight before use.

Tissue preparation

Rats were stunned with CO₂ gas and bled to death. The colonic segments were removed and opened along the mesenteric border, and the luminal contents were washed out by warm Krebs bicarbonate saline solution. For mucosa preparation, which histologically consisted of the mucosa and the muscularis mucosae (data not shown), the submucosa together with the tunica muscularis were removed with fine forceps under a stereomicroscope

Large and small epithelial sheets were made from the mucosa preparation. To measure the ACh release and electrical activity, two large sheets were prepared from each segment of the proximal or distal colon and were mounted in an Ussing chamber with a window size of 0.98 cm² and a volume of 5 ml. Four small sheets were prepared from the distal colon for the measurement of electrical activity and were mounted in an Ussing chamber with a window size of 0.20 cm² and a volume of 5 ml. To prepare four small sheets, the mucosa preparation of the distal colon was incised, first longitudinally and then transversely. Of the two small sheets from each transverse segment, one was treated with inhibitors of propionate response and the other, treated only with the solvent for the drug, served as control.

Short-circuit current measurement

The Ussing chambers were bathed on each side of the mucosa with a volume of 5 ml of bathing solution containing (in mM): NaCl, 119; CaCl₂, 1.25; MgCl₂ 1; K₂HPO₄, 2.2; KH₂PO₄, 0.2; NaHCO₃, 21 and Glucose, 10. The solution was bubbled with a gas mixture of 95% O₂ and 5% CO₂ (pH 7.4, 37 °C). The tissues were short-circuited by a

voltage clamp (Nihon Koden, Tokyo, Japan) at zero potential automatically with compensation for solution resistance. I_{sc} was continuously recorded and tissue conductance measured every min. The I_{sc} was referred to as positive when current flowed from the mucosal to the serosal side. The current was recorded by Power Lab system (ADInstruments, Bella Vista, Australia).

Propionate and ACh stimulation

The tissues were left for about 40 min to stabilize I_{sc} before the effects of propionate and drugs were studied. 25 or 50 μ l of sodium propionate (100 mM) and acetylcholine chloride (10 mM) were added to the mucosal and serosal sides in the Ussing chamber, respectively. Appropriate concentrations of other drugs in a volume of 5 to 10 μ l were added to the mucosal or serosal side 10 min prior to propionate or ACh additions.

ACh release

After I_{sc} of the mucosa preparation reached a stable baseline, ACh release experiments were initiated adding TTX (1 μ M) to the serosal side and eserine (0.1 mM) to both sides. Ten minutes later, propionate (1 mM) was added to the luminal side. A 200 μ l from both luminal and serosal fluids was taken at 5 min intervals to measure ACh concentration. After sampling, a 200 μ l of bathing solution was added to both fluids. The samples were stored at -80 °C until the ACh was analyzed.

Crypt isolation

The proximal and distal colons were removed according to the previously described anatomical divisions (Yajima, 1985). The colon was opened longitudinally and rinsed in

warm phosphate—buffered saline (PBS). Each colonic segment was incubated in a 10ml of Hank's balanced solution (Sigma Chemical Co, St. Louis, MO, USA) containing 5 mM dithiothreitol, 0.1% BSA, 1 mM glutamine and 30 mM EDTA (pH 7.4) for 15 min with shaking at 37 °C. After incubation, the colons were gently scraped with a rubber policeman to isolate the crypts, which were centrifuged at $100 \times g$ for 3 min at 4 °C. The crypt pellets were washed twice with cold PBS. The residual muscle tissues were vigorously scraped with a rubber policeman and washed with cold PBS. The crypt pellets and residual muscle tissues were weighed and stored at -80°C until analysis by RT-PCR and measurement of ACh.

RNA extraction and cDNA synthesis

Total RNA was extracted from the crypt pellets and residual muscle tissues with a QuickGene RNA tissue kit S II (Fujifilm Corp., Tokyo, Japan) and a RNase-free DNase set (Takara, Shiga, Japan) according to the manufacturer's instructions. cDNA was then synthesized from 150 ng of the total RNA with a ReverTra Ace qPCR RT kit (Toyobo, Osaka, Japan) according to the manufacturer's instructions.

RT-PCR analysis

RT-PCR was performed using a LightCycler 480 (Roche Applied Science, Tokyo, Japan). The expressions of six genes (ChAT, CHT1, OCT1, OCT2, OCT3 and VAChT) were assessed in all the samples. The expression of β-actin (housekeeping gene) was also assessed in this study. Specific primers and probe sets for each gene were designed according to the GenBank accession number with an online software provided by Roche Applied Science (https://www.roche-applied-science.com/). The sequences of the

primers and probes used in this study are listed in Table 1. PCR for all the genes was performed under the following conditions: an initial denaturation at 95 °C for 5 min, followed by 50 cycles of denaturation at 95 °C for 10 s and combined annealing-extension at 60 °C for 30 s. All reactions were performed using LightCycler 480 Probes Master and Universal ProbeLibrary Set (Roche) according to the manufacturer's instructions.

The relative expression level of each mRNA was calculated by the comparative Ct method (Nishimura *et al.*, 2003), wherein the Ct value of the target mRNA was subtracted from the Ct value of the β -actin mRNA. Data are presented as fold-differences to the expression level of the brain.

ACh measurement

The concentrations of ACh in the fluids obtained from the ACh release experiments, isolated crypts cells and residual muscle tissues were measured by high-performance liquid chromatography (HPLC) (Elite LaChrom, Hitachi Co. Ltd, Tokyo, Japan) combined with a post column enzyme reactor (AC-ENZYMEPAK, Eicom Co. Kyoto, Japan) and an electrochemical detector (ECD-700, Eicom Co. Kyoto, Japan). The mobile phase, applied at flow rate of 0.15 ml min⁻¹, comprised 50mM KHCO₃ containing 300mg 1⁻¹ sodium 1-decanesulfonate and 50mg 1⁻¹ EDTA-2Na. After separation by a styrene polymer column (AC-GEL, 2 × 150 mm, Eicom Co. Kyoto, Japan), ACh was converted to hydrogen peroxide by the post-column enzyme reactor with immobilized acetylcholine esterase and choline oxidase at 30 °C. The hydrogen peroxide was detected with the electrochemical detector which was equipped with a platinum electrode (+450 mV set potential). The ACh concentration was estimated by

the calibration curves of ACh standard using chromatography data software (D-2000 Elite, Hitachi Co. Ltd, Tokyo, Japan). Isopropyl homocholine was used as the internal standard.

The crypt pellets and residual muscle tissues were homogenized in 500 μ l of PBS containing eserine (0.1 mM) by Micro SmashTM (MS-100, Tomy Medical Ltd. Tokyo, Japan) at 3800 rpm for 3 min at room temperature. The lysates were centrifuged at $15000 \times g$ for 10 min at 4 °C and the supernatants were used for ACh measurement after appropriate dilution. Samples were loaded onto an autosampler (D-2000 Elite, Hitachi Co. Ltd, Tokyo, Japan) at 10 °C, and sample of 10 μ l was injected into the HPLC system.

Drugs

Sodium propionate and 3-Cl-propionic acid were obtained from Kanto Kagaku Co. Ltd (Tokyo, Japan). The free acid was neutralized with sodium hydroxide. Acetylcholine chloride, amiloride hydrochloride, atropine sulphate, bumetanide, dithiothreitol, EDTA, eserine hemisulphate, glutamine, tetrodotoxin and veratridine were obtained from Sigma Chemical Co. (St. Louis, MO, USA). Veratridine was dissolved in DMSO.

Statistics

Values are mean \pm S.E. The significance of the difference between data of the control and experimental groups was determined by Student's t test or one-way ANOVA followed by test of Dunnett. Linear regression was performed to determine the significance of differences in the relationship propionate-induced I_{sc} and ACh release. Differences were considered significant when P values were less than 0.05.

Results

Electrical activity of mucosa preparation

The basal I_{sc} of the mucosa preparation after stabilisation was $48.2 \pm 3.5 \,\mu\text{A cm}^{-2}$ (n = 16). Neuronal stimulations by bipolar rectangular electrical pulses (5 mA, 10 Hz) and serosal addition of veratridine (10 μ M) caused an increase in I_{sc} , 97.5 \pm 13.5 μ A cm⁻² (n = 4) and 149.4 \pm 4.0 μ A cm⁻² (n = 5), respectively. These neuronal responses were abolished by serosal addition of TTX (1 μ M), but were not affected by serosal addition of atropine (10 μ M).

Effects of TTX and atropine on I_{sc} responses to propionate and ACh

The serosal addition of TTX (1 μ M) decreased the basal I_{sc} by 28.4 \pm 4.8 μ A cm⁻² (n=8), but the serosal addition of atropine (10 μ M) had no effect on the basal I_{sc} . The residual I_{sc} after TTX treatment was almost abolished by the mucosal addition of amiloride (0.1 mM).

The luminal addition of propionate transiently increased I_{sc} not only across the mucosa-submucosa preparation but also the mucosa preparation as previously reported (Yajima, 1988). In this study, we confirmed that the mucosa preparation responded to luminal addition of propionate (0.5 mM) as well as serosal addition of ACh (0.05 mM), both of which caused a biphasic increase in I_{sc} . At first, a short downward spike followed by a higher transient increase in I_{sc} of similar duration was observed (Fig. 1).

The I_{sc} response to luminal propionate in the mucosa preparation was not significantly influenced by the serosal addition of TTX (1 μ M) (Fig. 2). On the other hand, the I_{sc} response to propionate was significantly inhibited by the serosal addition of

atropine (10 μ M) (Fig. 2). The serosal addition of TTX had no effect on ACh-induced I_{sc} responses, while the serosal addition of atropine abolished the I_{sc} responses to ACh (Fig. 2).

Relationship between propionate- and ACh-induced I_{sc} responses

Linear regression analysis of the maximal increases in I_{sc} between propionate- and ACh-induced I_{sc} responses showed a significant linear relationship (Fig. 3). This relationship, together with the results of the strong inhibition of I_{sc} response to propionate with atropine, suggests that ACh release mediates the luminal propionate-induced I_{sc} increase.

ACh release induced by luminal propionate

To address the involvement of ACh in the luminal propionate-induced I_{sc} increase, we directly measured ACh release from the mucosa preparation in response to luminal propionate in the presences of TTX (1 μ M) and eserine (0.1 mM). The mucosal and serosal fluids were sampled every 5 min. Before the propionate stimulation, ACh release was not observed in either fluid. After the propionate stimulation, ACh was detected in the serosal fluid but not in the mucosal fluid (Fig. 4). The amount of ACh release induced by propionate was 803 ± 181 pmol g⁻¹ tissue (n = 6) during the first 5 min period, 152 ± 61 pmol g⁻¹ tissue (n = 6) during the second 5 min period, and no release during the third 5 min period.

There was a significant linear relationship between the values of maximal I_{sc} increase and the amounts of ACh release after the propionate stimulation (Fig. 5).

Effects of atropine, bumetanide and 3-Cl-propionate on the increase in I_{sc} and ACh release induced by luminal propionate

Serosal additions of atropine (10 μ M) and burnetanide (50 μ M), a potent inhibitor of chloride secretion, in the presence of TTX significantly inhibited the I_{sc} increase induced by luminal propionate, but both drugs had no effects on ACh release into the serosal side (Fig. 6).

We previously reported that 3-Cl-propionate, an inactive analogue of propionate, had no effect on potential difference (PD) across the everted rat colon, but competitively inhibited the luminal propionate-induced increase in lumen-negative PD (Yajima, 1989). In this study, luminal addition of 3-Chloro-propionate had no effect on I_{sc} response, but completely blocked the propionate-induced I_{sc} increase as well as ACh release (Fig. 6). This result suggests that luminal propionate sensory input is received by a specific receptor site in the apical membrane of colonocytes and then causes ACh release.

Expression of genes related to ACh synthesis in isolated crypts

In light of the above data, it can be hypothesized that ACh release in response to luminal propionate in the mucosa preparation is of non-neuronal origin, and that epithelial cells are the most plausible candidates. To obtain colonic epithelium free of neuronal components, we isolated crypt cells from colonic sheets by means of Ca²⁺ free EDTA treatment (see details in Methods). The isolated crypt cells and residual muscle tissues were used for RT-PCR analysis of genes involved in ACh synthesis.

RT-PCR analysis showed that the crypt cells expressed higher mRNA levels of ChAT, an enzyme of ACh synthesis, compared to the residual muscle tissues (Fig. 7). On the other hand, mRNAs of neuron specific high affinity choline transporter (CHT1)

and vesicular acetylcholine transporter (VAChT) were scarcely detected in the crypt cells. This indicates that the isolated crypt cells were not contaminated by the enteric nerve components. The increased mRNA expression of organic cation transporters, OCT1, 2 and 3, suggest that these transporters probably play a role in choline uptake in epithelial cells compared to CHT1.

ACh content in isolated crypt cells and residual muscle tissues

The ACh content in the isolated crypt cells of the distal colon was 11.9 ± 2.0 nmol g⁻¹ wet wt, which was not significantly different than that in the residual muscle tissues of proximal and distal colons (Fig. 8). On the other hand, the ACh content in the crypt cells of the proximal colon was significantly lower than that of the distal colon.

Regional differences of propionate-induced ACh release and I_{sc} in the colon

Based on the difference of ACh contents in the crypt cells of the proximal and distal colons, we measured ACh release and I_{sc} response induced by luminal propionate stimulation in the proximal colon. As shown in Fig. 9, both ACh release and I_{sc} response were significantly lower in the proximal colon than in the distal colon. These results suggest that the regional difference of the secretory responses to luminal propionate depends on the difference of ACh content in the colonic crypt cells.

Discussion

In the present study, we have demonstrated for the first time that the chemical stimulation by luminal propionate induced non-neuronal ACh release from the colonic epithelium into the serosal side, which induced an increase in the I_{sc} , mainly due to

bumetanide sensitive chloride secretion, in rat distal colon. This result was substantiated by the direct measurements of ACh release after the luminal propionate stimulation and of ACh content in the isolated colonic crypt cells. Furthermore, these findings were supported by the expression of genes associated with ACh synthesis in the isolated colonic crypt cells.

The mucosa preparation of rat distal colon consists of the epithelium and the muscularis mucosae, not including the myenteric and submucosal nerve plexuses. Bridges et al. reported that the mucosa preparation of rat distal colon contains the neuronal network called the mucosal nerve plexus, which influences colonic ion transport (Bridges et al., 1986). It was shown that the Na⁺ channel activator, sea anemone toxin, and the electrical field stimulation (EFS) stimulate the neurons of the mucosal plexus, and induce the increases in I_{sc} and conductance across the mucosa preparation, which are abolished by TTX (Bridges et al., 1986; Diener et al., 1989). Furthermore, it was suggested that the mucosa plexus is involved in neuronally mediated cholinergic response according to the partial inhibition of the EFS-induced I_{sc} increase by atropine (50 µM) (Diener et al., 1989). In this study, we confirmed that nerve stimulations with the Na⁺ channel activator, veratridine, and EFS induced increases in I_{sc} in the mucosa preparation, which were completely abolished by TTX. It has been reported that veratridine causes ACh release from the submucosal neurons in rat colon (Gaginella et al., 1992). However, neither EFS nor veratridine could release ACh from the mucosal nerve plexus in the mucosa preparation since atropine had no effect on both-induced I_{sc} increase. On the other hand, the I_{sc} response to luminal propionate in the mucosa preparation was not affected by serosal addition of TTX (1 μ M) but was significantly inhibited by serosal addition of atropine (10 μ M). This

indicates that the I_{sc} response to propionate is mediated by ACh that is released by means other than excitation of the mucosal neurons.

It is well known that muscarinic ACh receptors are expressed on intestinal epithelial cells (Hirota & McKay, 2006). In rat colonic epithelial cells, M_1 and M_3 muscarinic receptors are responsible for ACh receptor-mediated chloride secretion (O'Malley *et al.*, 1995; Haberberger *et al.*, 2006). In this study, the ACh-induced increases in I_{sc} in the mucosa preparation were not affected by TTX but abolished by atropine, indicating a direct action of ACh on colonic epithelial muscarinic receptors. Hence, the TTX-resistant and atropine-sensitive I_{sc} responses to propionate (Fig. 2) suggest ACh release from non-neuronal components in the colonic mucosa.

As expected, the luminal addition of propionate induced the largest ACh release into the serosal side in the mucosa preparation 5 min after the stimulation, indicating that the I_{sc} increase observed during the same period must be elicited by the released ACh (Fig. 4). This speculation was supported by the observation of a highly significant linear relationship between the amount of ACh release and the maximal increase in I_{sc} (Fig. 5). The evidences that atropine and bumetanide inhibited the I_{sc} responses to luminal propionate but not the ACh release (Fig. 6), strongly suggests that the I_{sc} increase induced by the luminal propionate is mediated by the ACh that is released into the serosal fluid. In this study, we adopted the HPLC technique combined with enzyme reactors and electrochemical detector, a highly sensitive and specific method to measure ACh concentration in the experimental samples. In addition, sufficient concentrations of eserine to block acetylcholine esterase (AChE) activity in the mucosa preparation were added to both sides of Ussing chambers since AChE is richly involved in the nerve fibres in rat colonic mucosa (Mestres *et al.*, 1992).

It has been generally accepted that cells or tissues undergo a process of desensitization following sustained stimulation with a high concentration of receptor agonist. We have previously demonstrated that the secretory response to SCFAs is rapidly desensitized by the repeated stimulation of same or different acids. However, cross-desensitization between ACh and propionate does not occur in the mucosa-submucosa preparation of rat colon (Yajima, 1988). In the present study, the ACh release induced by propionate did not apparently affect the secretory response to ACh (Fig. 3), suggesting that the ACh release induced by luminal propionate is rapidly desensitized at the SCFA receptor and does not reach enough concentration to cause desensitization of muscarinic receptor.

Even though the ACh release was TTX-resistant, it may be not exclude the possibility that the ACh release in response to the luminal propionate in the mucosa preparation was of neural origin as it has been shown that the mucosa is richly innervated by the nerve fibres of postganglionic cholinergic neurons in the submucosal plexus (Harrington *et al.*, 2010a) Therefore, to evaluate the synthesis and storage of ACh in colonic epithelial cells, we prepared a colonic crypt free of mucosal nerve elements and measured the mRNA expressions of genes involved in ACh synthesis. The negligible expression of CHT1 and VAChT, specific markers of central and peripheral nerves (Harrington *et al.*, 2010b), in the crypt cells compared to the residual muscle tissues containing enteric nerves (Fig. 7) shows that the isolated crypt cells were free of the contamination of neuronal elements. Higher mRNA expression of ChAT gene in the crypt cells rather than in the residual muscle tissues indicates that the colonic epithelial cells have the ability to synthesize ACh. This was confirmed by direct measurement of the ACh contents in the crypt cells and the residual muscle tissues (Fig. 8).

Klapproth et al. previously reported that non-neuronal ACh content in the small and large intestines of rat were 1.3 ± 0.3 (n = 4), 1.4 ± 0.4 (n = 5) nmol g⁻¹ wet wt, respectively (Klapproth *et al.*, 1997). They sampled the tissues above the muscularis mucosae by scraping the mucosal surface with a cotton-tipped applicator, indicating that their samples contained submucosal nerve components. Their value of ACh content in the large intestine is close to that obtained in the crypts cells from the proximal colon (1.8 ± 0.2 nmol g⁻¹ wet wt, n = 4) and less than that from the distal colon (11.9 ± 2.0 nmol g⁻¹ wet wt, n = 4). The ACh content in the crypt cells from the rat distal colon is approximately three orders less than that in different regions of the rat brain (Vizi & Palkovits, 1978) and in the myenteric plexus in the guinea-pig distal colon (Giaroni *et al.*, 1999).

Choline is a key substrate for ACh synthesis. In the neurons, the uptake of choline is mediated by high-affinity choline transporter, CHT1 (Brandon *et al.*, 2004). In the crypt cells, the expression of CHT1 was not detected by RT-PCR, but higher expression of mRNAs for polyspecific organic cation transporters, OCT1, 2 and 3, was found in comparison with nerve-containing residual muscle tissues. Hence, the OCTs and not CHT 1, may play a role in choline uptake for ACh synthesis in the colonic epithelial cells as choline is one of the endogenous substrates of OCTs (Koepsell *et al.*, 2007).

The regional differences in the secretory responses to the luminal propionate in rat colon (Yajima, 1989) can be explained by the regional differences of the ACh content and release in the proximal and distal colon (Fig. 9). On the other hand, no significant difference in ACh content in the nerve containing muscle tissues of the proximal and distal colon strongly supports the hypothesis that the secretory response to the luminal

propionate depends on colonic epithelial ACh synthesis and release.

The ACh release from non-neuronal epithelial cells was reported for the first time in the human placenta (Wessler et al., 2001). By the superfusion of the villus isolated from the placenta, the ACh releases were observed at a rate of 0.13 nmol g⁻¹ wet weight min⁻¹. Interestingly, this value is comparable to that (0.16 nmol g⁻¹ wet wt min⁻¹) observed in the propionate-induced ACh release from rat colonic mucosa in this study. In the placenta villus, however, the ACh release spontaneously occurred without any stimulation, but not in the colonic mucosa. The stimulus-coupled release of non-neuronal ACh was recently reported in the human bladder strip with urothelium, in which the stretch tension enhanced the ACh release in the presence of TTX (Yoshida et al., 2008). To address physiological role of non-neuronal ACh release from the epithelium, it is important to know whether ACh is released into luminal or serosal side. However, the previous studies did not reveal the direction of ACh release from the epithelial cells of the placenta and bladder. In the present study, we clearly demonstrated that the ACh release coupled with the luminal propionate stimulation occurred into the serosal side, but not luminal side in the rat colon using an Ussing chamber.

In general, a stimulus-coupled release of transmitters is observed in excitable cells such as neurons and endocrine cells. The release of ACh in response to luminal propionate stimulation (Fig. 4) appears to be similar to the stimulus-coupled phenomenon, probably via ACh-containing sensory cells on colonic mucosa. In our previous study, we have speculated that the sensory stimulation of secretory response to luminal SCFAs is received at a specific receptor site in the mucosal surface of the rat colon (Yajima, 1988). The prior addition of luminal 3-Cl-propionate completely

blocked the I_{sc} response and also abolished the ACh release in response to the luminal propionate (Fig. 6). These results indicate that epithelial cells storing ACh have a receptor site for propionate, although a specific epithelial cell type storing ACh in the colonic epithelial cells has not yet identified. The SCFA receptor, free fatty acid receptor 2 and 3 (FFA2 and FFA3), that was immunohistochemically demonstrated in the colon of rats and humans (Karaki *et al.*, 2006; Tazoe *et al.*, 2008) is a possible receptor in the colonic secretory response to luminal SCFAs. Further studies are required to identify the specific cell type on colonic mucosa that respond to luminal propionate, and to determine the mechanism of ACh release.

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193-198.

Author contributions

Conception and design of the experiment: T.Y. Collection, analysis and interpretation of data: T.Y., R.I., M.M. and M.Y. Drafting the article or revising it critically: T.Y. and M.Y. All authors approved the final version of this manuscript.

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Figure legends

Fig. 1. A typical trace of short-circuit current (I_{sc}) measurement in the mucosa preparation.

After the basal Isc was stabilised, TTX (1 μ M) was added to the serosal fluid, 10 min later, propionate (0.5 mM) was added to the mucosal fluid. ACh (0.05 mM) was added to the serosal fluid after I_{sc} returned to the basal level.

Fig. 2. Effects of tetrodotoxin (TTX, 1 μ M) and atropine (10 μ M) on basal I_{sc} and luminal propionate-induced I_{sc} in mucosa preparation.

TTX (n = 8) and atropine (n = 8) were added to the serosal fluid 10min before the luminal addition of propionate (0.5 mM), respectively. Values are mean \pm S.E. of the maximal increase in I_{sc} and statistical significances were assessed with one-way ANOVA followed by test of Dunnett. *P<0.01 compared to the control.

Fig. 3. Relationship between propionate-induced Isc and ACh-induced I_{sc} .

Linear regression analysis was performed based on the difference between the maximal increases of I_{sc} induced by luminal propionate and that by serosal ACh in the same mucosa preparation.

Fig. 4. ACh release induced by luminal propionate in the mucosa preparation.

The concentration of ACh in the serosal fluid was measured every 5 min. Red line shows changes in I_{sc} plotted by the mean value every min (n = 6). Blue bars show ACh releases into the serosal fluid which were plotted as the mean value every 5 min (n = 6). The amount of ACh release is shown in pmoles per wet weight of the tissue covered on the window of the Ussing chamber. Propionate (1 mM) was added to the mucosal fluid 10 min after the start.

Fig. 5. Relationship between ACh release and I_{sc} increase induced by luminal propionate.

Linear regression analysis was performed based on the difference between the amount of ACh released for 5min and the maximal increases in I_{sc} induced by luminal propionate.

Fig. 6. Effects of atropine (10 μ M), bumetanide (50 μ M) and 3-Cl-propionate (1 mM) on I_{sc} response and ACh release induced by luminal propionate in the mucosa preparation.

Atropine and bumetanide were added to the serosal fluid and 3-Cl-propionate was added

to the mucosal fluid 10 min prior luminal propionate (1mM) stimulation. The concentration of ACh in the serosal fluid was measured at 5 min after luminal propionate stimulation. Values are mean \pm S.E. (n=6-16). Statistical significances of the maximum increases in I_{sc} and the ACh releases were assessed with one-way ANOVA followed by test of Dunnett. *P<0.001 compared to the control.

Fig. 7. Expression of genes related to ACh synthesis and storage.

RT-PCR analysis indicates that ChAT and OCTs mRNAs were expressed in the isolated crypt cells, while neuronal marker mRNAs of CHT1 and VAChT were scarcely expressed. A dot line indicates the level expressed in the brain. Values are mean \pm S.E. (n = 4).

Fig. 8. ACh content in the isolated crypt cells and the residual muscle tissues from the proximal and distal colon.

ACh concentrations in the tissues were measured after homogenizing with presence of eserine (0.1 mM) and centrifugation. The obtained supernatants were used in HPLC analysis. Values are mean \pm S.E. (n = 4). *P<0.05 compared between crypt cells and residual muscle tissues by Student's t-test.

Fig. 9. Regional difference in ACh content, ACh release and I_{sc} response between the proximal and distal colon.

A: ACh release induced by luminal propionate (1 mM). B: I_{sc} increase induced by luminal propionate (1 mM). Values are mean \pm S.E. (n = 4). *P<0.05 compared between the proximal and distal colon by Student's t-test.

Table 1. The sequences of primers and probes used in real-time PCR analysis

Table

Gene Name	Accession No.	P	Probe No.		
β-actin	NM_031144	Forward	cccgcgagtacaaccttct	#17	
	p-actiii	NWI_031144	Reverse	cgtcatccatggcgaact	#1/
ChAT	СЬАТ	NM 001170593	Forward	gcctcatctctggtgtgctt	#120
	11111_001170393	Reverse	cagtcagtgggaagggagtg	#120	
CHT1	NM_053521	Forward	cattggtttgttggttggtg	#83	
		Reverse	gctgtcccgttgatgtaacc	ποσ	
OCT1	NM_012697	Forward	gctagctgtgtccctgccta	#4	
		Reverse	gggattctgggacaaacca	π-	
OCT2	NM_031584	Forward	ctgtgactctgcccaacttct	#15	
		Reverse	ggagatcagccatcttggag	π13	
OCT3	NM_019230	Forward	gactggcgctatgtggagac	#22	
		Reverse	gccgcagacaaggtcaaa	$\pi \mathcal{L} \mathcal{L}$	
VAChT	NM_031663	Forward	tgcaagagcactgtccaact	#10	
		Reverse	cgaggggctagggtattcat	#10	

Listed probe numbers indicate the product number of the Universal ProbeLibrary Set and Human and Extension Set sold by Roche Applied Science.

Fig. 1

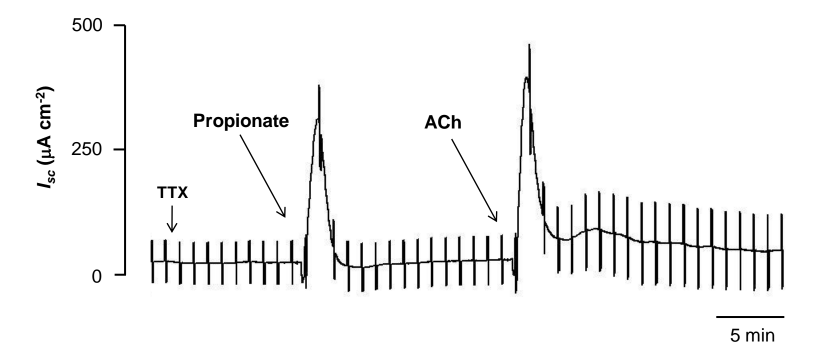


Fig. 2

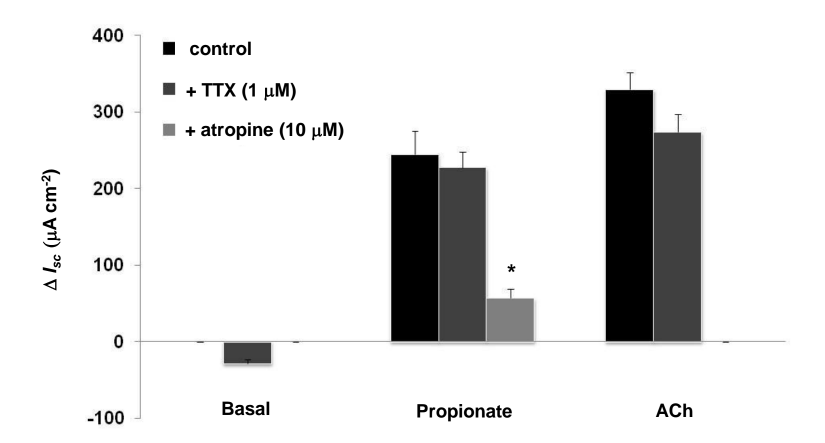


Fig. 3

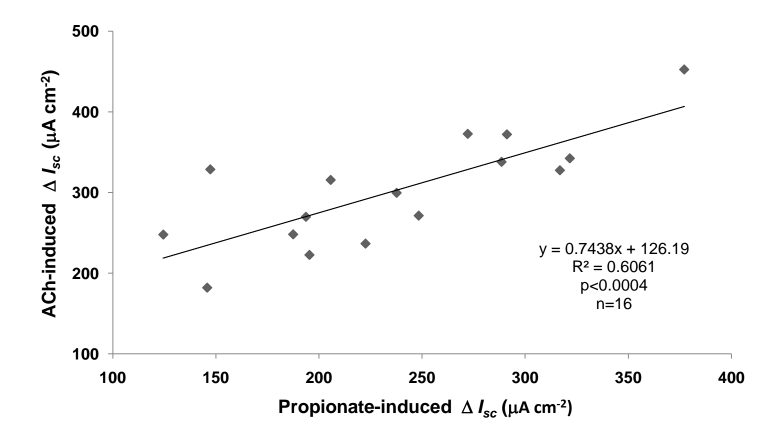


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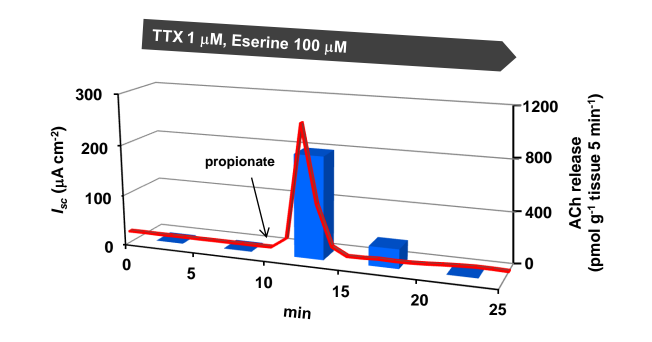


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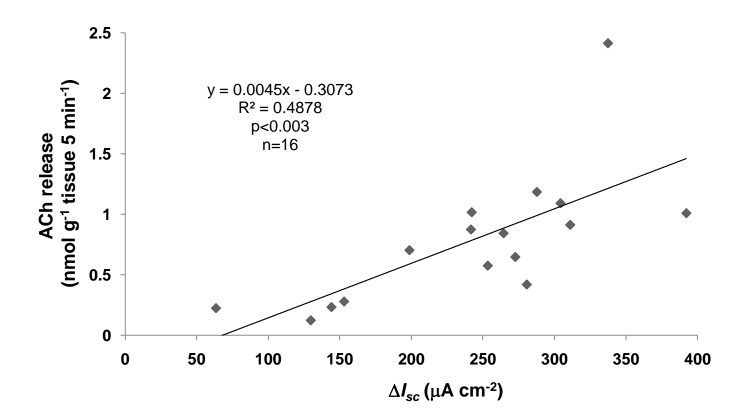


Fig. 6

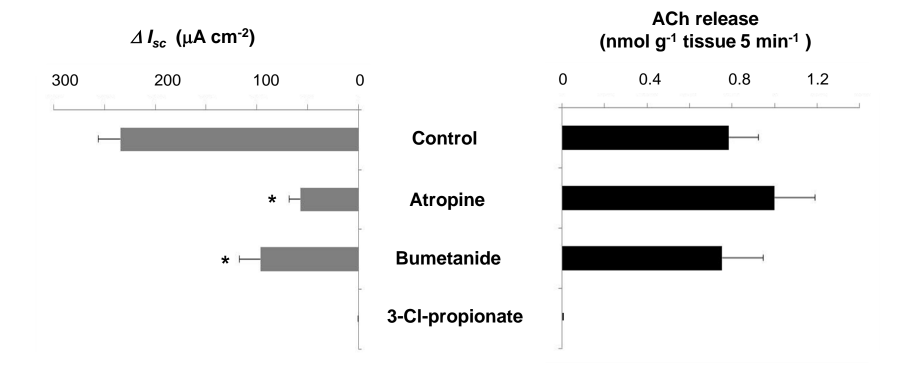


Fig. 7

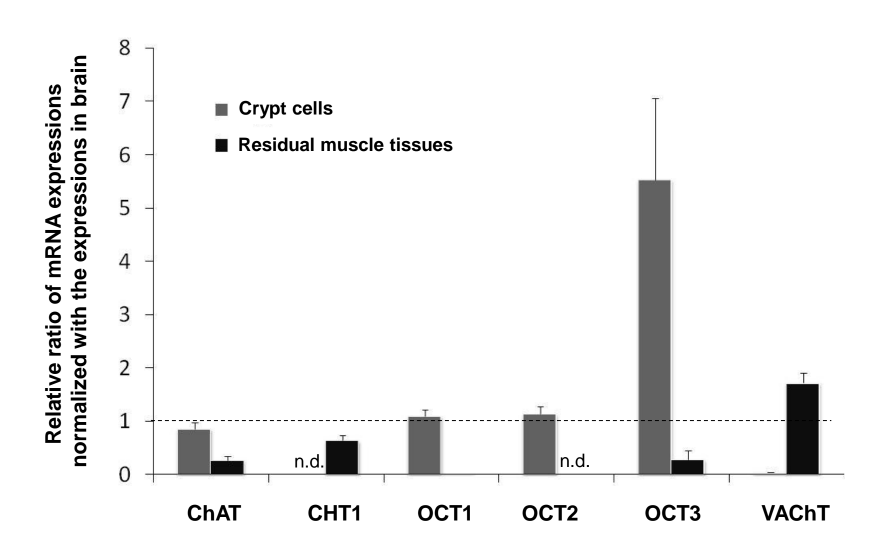


Fig. 8

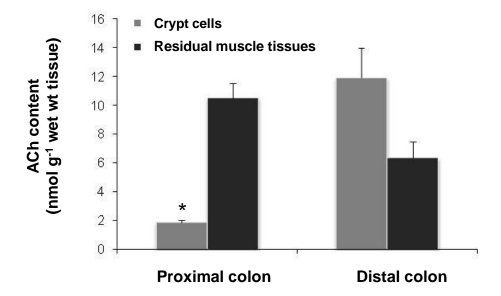


Fig. 9

