Effects of an Enteral Formula Containing Fermented Dairy Products on Epithelial Ion Transport in Rat Intestines

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Summary Diarrhea is the most common complication of enteral nutrition (EN). Pro/prebiotics are typically used to prevent diarrhea during EN. This study aimed to demonstrate the effects of enteral formula containing fermented dairy products (FDPs) and galacto-oligosaccharides on intestinal mucosal functions in rats. After feeding rats with regular rodent chow (RRC), standard formula (STD-F), and FDP-containing formula (FDP-F) for 2 wk, the rats were sacrificed with their intestines removed. Then, the electrophysiological properties of intestinal epithelia were measured using the Ussing chamber. In addition, organic acids and microbiota in the cecal contents were analyzed. In FDP-F-fed rats, electrical nerve activation-evoked increase in short-circuit current (Isc) in the cecum and middle colon was reduced compared with STD-F-fed rats. Mucosal propionate-evoked changes in Isc in FDP-F-fed rats were also reduced in the terminal ileum. The total cecal organic acid concentration in STD-F-fed rats decreased compared with RRC-fed rats, and approximately half was recovered in FDP-F-fed rats, which contributed to the recovery of acetate and butyrate concentrations. In microbiota analysis, the density of total bacteria, particularly Bifidobacterium, in cecal contents increased in FDP-F-fed rats. In conclusion, the consumption of FDP-F changed the total amounts and components of gut microbiota and organic acids, and resulted in inhibitory changes in mucosal luminal stimulant- and nervous system-mediated fluid secretory function. These findings suggest that FDP-F might prevent the incidence of diarrhea during EN.

Key Words enteral nutrition, diarrhea, probiotics, prebiotics, short-chain fatty acids, Ussing chamber

Enteral nutrition (EN) provides nutrients directly into the gastrointestinal (GI) tract bypassing the oral cavity. It is required for patients with swallowing or chewing problems in the hospital or community setting. EN is a physiological feeding associated with fewer serious complications compared with parenteral nutrition (central venous feeding). However, complications, such as nausea, bloating, diarrhea, excessive ostomy output, constipation, abdominal discomfort or pain, and reflux, often occur during EN (1). In particular, diarrhea is the most common complication of EN; it affects 2% to 95% (this wide range resulted from the differences in the patient groups investigated and variations in the definition of diarrhea used) of patients (2).

Several causes of diarrhea during EN, including malabsorption, infection, bacterial contamination of diet, medical diagnosis of the patient, medication therapy, and formula-related causes, are considered (3). GI microbiota and short-chain fatty acids (SCFAs), which are fermentation products of GI microbiota, are implicated in the incidence of diarrhea during EN (4). Therefore, the roles of probiotics, defined as “live microorganisms which when administered in adequate amounts confer a health benefit on the host,” (5) and prebiotics, defined as “selectively fermented ingredients that allow specific changes, both in the composition and/or activity in the gastrointestinal microflora that confers benefits upon host well-being and health.” (6) in the reduction of the incidence of diarrhea during EN have been investigated (2, 7). Probiotics, such as Lactobacillus rhamnosus GG, have not shown promise as a treatment for diarrhea that is already occurring in critically ill patients (8). However, Saccharomyces boulardii as a probiotic has been reported to prevent diarrhea in critically ill patients (9). In a meta-analysis study of prebiotics, fiber-supplemented enteral formula (EF) has been found to have clinical benefit in reducing the incidence of diarrhea (10), but fructo-oligosaccharides (FOS), oligofructose, and inulin (as prebiotics) supplementations in EF did not minimize the incidence of diarrhea in EN (11, 12).

Fermented milk by lactic acid bacteria (LAB) is the first probiotic advocated by Metchnikoff in 1908 (13). Therefore, LAB-fermented milk has prospect of beneficial effects on diarrhea during EN. LAB-fermented milk-containing EF is at least clinically speculated to prevent the incidence of diarrhea; however, its mechanism is still unclear.

The fermented dairy product-containing formula (FDP-F) in the present study contained LAB (Lactobacil-
Enteral Formulas and Intestinal Ion Transport

N. bulgaricus and Streptococcus thermophilus)-fermented milk, propionic acid bacteria (PAB) (Propionibacterium freudenreichii ET-3)-fermented milk whey, and galactooligosaccharides (GOS). L. bulgaricus and S. thermophilus are the traditionally used combination for the production of yoghurt (14). PAB-fermented milk whey has a prebiotic function that enhances the growth of Bifidobacterium in normal adult humans (15), and 1, 4-dihydroxy-2-naphthoic acid is isolated and identified as a bifidogenic growth stimulator (16).

Infusion of SCFAs directly into the cecum has been reported to reverse the fluid secretion found in the ascending colon during EN (17). We therefore hypothesized that FDP-F-feeding may change the GI microbiota, increase SCFAs in the large intestinal contents, and induce an inhibitory effect on the secretory function of epithelia. Thus, in the present study, we aimed to investigate the effects of the consumption of sterilized FDP-F on the transepithelial ion transport inducing fluid secretion or absorption in the rat small and large intestines.

In the present study, fluid secretion in the intestinal epithelia was measured by Ussing chamber technique as an electrophysiological method (18). Short-circuit current (Isc) measured by the Ussing chamber technique is an index of net transepithelial ion transport. Driving force of intestinal fluid secretion is produced by the increase in luminal osmotic pressure. Active and electrogenic anion (Cl– and HCO3–) transport from serosal to luminal side of the epithelium produce a lumen negative potential difference, which induces passive cation (Na+) secretion and then increases luminal NaCl concentration and osmotic pressure. Therefore, measurement of the electrogenic anion secretion by Ussing chamber technique as a change in ΔIsc is relevant for evoking a massive fluid secretion and diarrhea. In addition, the epithelial sodium channel (ENaC) expressed in the apical membrane especially in the distal colonic and rectal epithelia induces an electrogenic Na+ absorption that increases Isc, and potently absorbs luminal water. Therefore, the inhibition of ENaC activity that decreases Isc induces an inhibition of water absorption in the distal colon and rectum, possibly inducing diarrhea. These indicate that both increase and decrease in Isc possibly induces diarrhea. Therefore, in the present study, measurements of the changes in Isc were performed to investigate the effects of the consumption of FDP-F.

MATERIALS AND METHODS

Experimental diets. Powder diets of two freeze-dried formulas and one regular rodent chow (RRC) were used. Meibalance (Meiji, Tokyo, Japan) as a standard formula (STD-F), YH Flore (Meiji) as an FDP-F, and CRF-1 (Oriental Yeast Co., Ltd., Tokyo, Japan) as an RRC were used. YH Flore, but not Meibalance, contained sterilized LAB (L. bulgaricus and S. thermophilus)-fermented milk, sterilized PAB (P. freudenreichii ET-3)-fermented milk whey (Profec®, Meiji), and GOS. The components of these experimental diets are shown in Table S1 (Supplemental Online Material). These enteral formulas were kindly gifted by Meiji Co., Ltd.

Animals. The animals were handled and killed in accordance with the Guidelines for the Care and Use of Laboratory Animals of the University of Shizuoka, and the study was approved by the University of Shizuoka Animal Usage Ethics Committee (approval No. 145069 and 1551103 approved on 11 August, 2014 and 9 September, 2015, respectively). Male Sprague-Dawley rats aged 6 wk (n = 18) were purchased from Japan SLC, Inc. (Hamamatsu, Japan). After a 1-wk acclimatization period, they were divided into three groups (six rats per group) by body weight-stratified randomization: RRC-, STD-F-, and FDP-F-fed groups. The animals were fed with each diet and given water ad libitum for 2 wk. In the present study, food intake could not be measured because the powder diet was spilled during the feeding and the metabolic cages were not used.

Sampling and tissue preparation for the Ussing chamber experiments measuring the electrogenic transepithelial ion transport. Two-week RRC-, STD-F-, and FDP-F-fed rats were anesthetized with isoflurane inhalation and decapitated with a guillotine. The GI tract en bloc from the stomach to the anus was removed and then immersed in ice-cold Krebs-Ringer solution containing (mm) 117 NaCl, 4.7 KCl, 1.2 MgCl2, 1.2 NaH2PO4, 25 NaHCO3, and 2.5 CaCl2. The solution was saturated with 95% O2–5% CO2. Mesenteric and fat tissues were removed from the specimens, and the isolated GI tracts were extended on a Ringer-soaked paper towel. Photo images were taken with a ruler using a digital camera to measure the length of the intestine. During the middle of experiments, I found that the size of the cecum was very different among the experimental groups. Therefore, I added the analysis of cecal tissue and the contents in residual 3 animals from that time. The cecum was removed, and gross weight (cecal tissue + contents) was measured. The cecum was cut open along the mesentery of the lesser curvature, and the cecal contents were taken to measure the organic acid concentrations. The contents were washed out, and cecal tissue weight was measured. The cecal content samples were frozen and stored at −80°C until analysis.

For the Ussing chamber experiments, the terminal ileum, cecum, proximal colon, middle colon, and distal colon were removed, cut open along the mesenteric border, and pinned flat to the silicon-rubber-lined petri dish with Krebs-Ringer solution. To investigate the electrophysiological properties of the epithelium, mucosa-submucosa preparations were made by removing the muscle layer under stereomicroscopy.

Ussing chamber experiments. Ussing chamber experiments were performed following a method described previously (19). Mucosa-submucosa preparations were mounted on the Ussing chambers (cross-sectional area: 0.64 cm²; CHM2, World Precision Instruments (WPI), Inc., Sarasota, FL, USA) with a pair of aluminum foil ribbon electrodes for electrical field stimulation (EFS). Both sides were perfused with 10 mL of Krebs-Ringer solution each by recirculation from the glass circulation reservoir (#5210, WPI) maintained at 37°C and gassed with 95% O2–5% CO2. D-Glucose (5 mM) was added
Table 1. Body weight, lengths of intestines, and cecal conditions in RRC-, STD-F- and FDP-F-fed rats.

|                      | RRC          | STD-F        | FDP-F        |
|----------------------|--------------|--------------|--------------|
| Body weight (g), n=6 | 319.9±8.1    | 308.2±8.1    | 305.7±7.8    |
| Length of small intestine (mm), n=6 | 693.4±40.4  | 592.3±42.1   | 613.5±45.3   |
| Length of large intestine (mm), n=6 | 152.7±6.5a  | 126.1±6.9b   | 135.9±4.6ab  |
| Cecal tissue weight (g), n=3 | 1.13±0.09a   | 1.12±0.10a   | 1.69±0.12ab  |
| Wet weight of cecal contents (g), n=3 | 5.50±0.033a | 3.66±0.43b   | 4.49±0.33ab  |

Data are expressed as mean±SE. Different superscript alphabets differ significantly (p<0.05) by Tukey test.

Table 2. Basal electrophysiological parameters of the mucosa-submucosa preparations in a variety of intestinal segments in RRC-, STD-F- and FDP-F-fed rats.

|                      | RRC          | STD-F        | FDP-F        |
|----------------------|--------------|--------------|--------------|
| Basal I_sc [µA/cm²]  |              |              |              |
| Terminal ileum       | 17.95±2.25   | 30.95±9.37   | 13.21±3.86   |
| Cecum                | 21.84±6.62   | 32.77±11.17  | 8.76±3.36    |
| Proximal colon       | 30.80±3.97   | 35.73±1.86   | 34.63±2.91   |
| Middle colon         | 16.06±0.58   | 17.28±1.29   | 21.22±3.66   |
| Distal colon         | 13.29±2.33   | 21.44±5.41   | 20.62±3.21   |
| Rectum               | 3.25±1.78    | 12.42±5.16   | 1.38±3.68    |
| Basal G_sc [mS/cm²]  |              |              |              |
| Terminal ileum       | 20.12±0.44   | 19.46±0.80   | 21.52±0.53   |
| Cecum                | 13.63±0.99a  | 15.35±0.78a  | 25.82±2.74b  |
| Proximal colon       | 20.27±1.07   | 15.22±1.51   | 22.03±3.38   |
| Middle colon         | 13.42±1.96a  | 15.50±1.41a  | 29.53±4.94b  |
| Distal colon         | 7.58±0.83a   | 12.37±2.05a  | 13.03±1.29b  |
| Rectum               | 7.82±1.22    | 17.15±3.45a  | 10.63±1.68   |

Data are expressed as mean±SE (n=4–6). Different superscript alphabets differ significantly (p<0.05), and a,b indicates tendency (p<0.10) of difference vs RRC by Tukey test (distal colon) or Games-Howells test (cecum, middle colon and rectum).

to the serosal solution only. The transepithelial potential difference (the serosal electrode served as the reference) was detected by paired Ag-AgCl electrodes (EKV, WPI) through Krebs-agar bridges, and was clamped at 0 mV by applying an I_sc with another pair of Ag-AgCl electrodes (EKV, WPI) connecting a voltage-clamp apparatus (CEZ-9100; Nihon-Kohden, Tokyo, Japan). The lumen negative electronic current indicates a value of positive I_sc per unit area of tissue (µA/cm²). To record tissue conductance (G_sc [mS/cm²]), voltage command pulses (10 mV, 3-s duration) were applied at 1-min intervals, and G_sc was calculated by Ohm’s law from necessary to current necessary to change the clamped voltage. The current output was continuously recorded on a data acquisition and analog-to-digital conversion system (PowerLab 4/26; ADInstruments, Cattle Hill, Australia). Immediately after mounting the tissues, I_sc and G_sc showed higher value, then gradually decreased, and achieved plateau within 1 h (Fig. 1). Thus, tissues were stabilized for 1 h before the experiments.

Protocol of the Ussing chamber experiments. After a 1-h stabilization period, basal I_sc and basal G_sc were determined, and the tissues were electrically stimulated at 25 V, 5 Hz, 0.5 ms-duration for 2 min via a pair of aluminum foil ribbon electrodes. Propionate (1 mM) was added to the mucosal bathing solution 1 h after the EFS, and changes in I_sc were measured. Further 1 h after the addition of propionate, carbachol (CCh) was added to the serosal bathing solution. Representative I_sc traces of the RRC-fed group from the time immediately after mounting tissues to the time after the final addition of CCh are shown in Fig. 1.

Chemicals. Sodium propionate was purchased from Wako Pure Chemical Industries, Ltd. (Osaka, Japan), and CCh was obtained from Sigma (St. Louis, MO, USA).

Organic acid concentrations in the cecal contents. Approximately 200 mg of cecal content was diluted with 2 volumes of Milli-Q water and homogenized, followed by centrifugation for 10,000 rpm at 4°C for 10 min. About 2.5 µL of Carrez Reagent I (53.5 g ZnSO₄·7H₂O/100 mL Milli-Q) and 2.5 µL of Carrez Reagent II (17.2 g K₃Fe(CN)₆·3H₂O/100 mL Milli-Q) were added to 200 µL of the supernatant to precipitate protein (20). The supernatant was passed through a 0.22-µm filter and used for the assay. Organic acids were measured with a post column pH buffering electric conductivity detection method using an electric conductivity detector (CDD-10A, Shimadzu Corporation) according to the modified method of Niwa et al. (21). A double-connected polymer column for organic acid
analysis (ICSep-ORH-801 6.5 mm×300 mm; Tokyo Chemical Industry Co., Ltd.) was used at a column temperature of 50°C, mobile phase of 5 mM p-trinitrosulfonic acid solution, reaction solution of 20 mM Bis-Tris aqueous solution containing 5 mM p-trinitrosulfonic acid solution and 100 μM EDTA (2Na), and flow rate of 0.5 mL/min. The total concentration was calculated as the sum of each organic acid.

**Analysis of microbiota.** Analysis of microbiota in the cecal contents was performed using the same method previously reported (22, 23). The microbiota DNA in the cecal content samples was extracted using QIAamp DNA stool mini kit (QIAGEN, Tokyo, Japan) and multi-beads shocker (Yasui Kikai, Osaka, Japan). Counts of total and individual bacterium [Bifidobacterium (24), Lactobacillus (25), Bacteroides (24), Clostridium cocoides (24), Clostridium leptum (24), Enterobacteriaceae (26), Enterococcus (27)] were determined with the ABI7300 real-time PCR system (Applied Biosystems, Tokyo, Japan) using QuantiTect SYBR Green RT-PCR (QIAGEN) and published primer base sequences and reaction conditions. The numbers of total bacteria and individual bacterium per gram of wet cecal contents were determined.

**Statistics.** Data are expressed as mean±SE after outlier values were removed by Smirnov-Grubbs test. Tukey-HSD test in homoscedasticity and Games-Howells test in heteroscedasticity were used to determine statistically significant difference (p<0.05) and tendency (0.05≤p<0.10) between each group using SPSS Statistics version 23 (IBM, Armonk, NY, USA).
RESULTS

Body weight, lengths of the small and large intestines, and weights of cecal tissue and its content

Data on body weight, lengths of intestines, and cecal conditions are shown in Table 1. The body weight and length of the small intestine (from the pyloric sphincter to the ileocecal junction) were not significantly different between each diet-fed group. However, the length of the large intestine (from the cecocolonic junction to the anus) in the STD-F-fed group was significantly shorter than that in the RRC-fed group. In the FDP-F-fed group, the length of the large intestine was not significantly different from the RRC- or STD-F-fed group.

The wet cecal tissue weight in the FDP-F-fed group was significantly heavier than that in the RRC- and STD-F-fed groups. Moreover, the wet weight of cecal contents in the STD-F-fed rats was significantly lighter than that in the RRC-fed group, but not than that in the FDP-F-fed group. The contents of the large intestine in the FDP-F-fed group seemed to be stickier in consistency than that in the other groups.

Basal electrophysiological parameters in the intestinal mucosa-submucosal preparations

Basal $I_{sc}$ and $G_t$ in mucosa-submucosal tissue preparations from each diet-fed rat are shown in Table 2. In basal $I_{sc}$, no significant difference and tendency were found between each other. In basal $G_t$, the values of the cecum, middle colon, and distal colon isolated from the FDP-F-fed group were significantly higher than those from the other diet groups. Basal $G_t$s in the distal colon and rectum from the STD-F-fed group tended to be higher ($p<0.10$) than that from the RRC-fed group.

EFS-evoked increase in $I_{sc}$ in the intestinal mucosa-submucosal preparations

During the EFS, $I_{sc}$ gradually increased and achieved maximum approximately until 2 min in all the segments, and returned to the basal level for several minutes after the EFS was completed as shown in Fig. 1 (EFS). In the cecum and middle colon, the EFS-evoked $I_{sc}$ response in the FDP-F-fed rat cecum was significantly reduced in comparison with the RRC- and STD-F-fed groups (Fig. 2, cecum and middle colon). In the proximal colon, the EFS-evoked $I_{sc}$ response in both the STD-F- and FDP-F-fed groups was significantly smaller than that in RRC-fed group (Fig. 2, proximal colon). In other segments, the EFS-evoked increases in $I_{sc}$ were not significantly different among the feeding groups.

Mucosal propionate-evoked changes in $I_{sc}$

The mucosal addition of one of the SCFAs, propionate (1 mM), which was reported to evoke an increase in $I_{sc}$ only at the mucosal side in the rat colon (28), evoked changes in $I_{sc}$ except in the proximal colon (Fig. 1, ▲ propionate). Propionate evoked a fast-phasic negative change in $I_{sc}$ within several seconds (P-1) in the terminal ileum, distal colon, and rectum, but not in the cecum and middle colon, and then a phasic positive change in $I_{sc}$ in a few minutes (P-2) in all segments except the...
In the terminal ileum of the STD-F-fed group, the mucosal propionate-evoked P-1 $I_{sc}$ response tended to be enhanced ($p<0.10$; Fig. 3B terminal ileum), and the P-2 response was significantly enhanced compared with the RRC-fed group (Fig. 3A terminal ileum). Although the P-2 response in the FDP-F-fed group was also significantly enhanced, the P-1 response in the FDP-F-fed group was not significantly different compared with each other.

In the cecum, although the propionate-evoked P-2 $I_{sc}$ response in the STD-F-fed group was not significantly reduced compared with the RRC-fed group, the response in the FDP-F-fed group was significantly reduced compared with the RRC-fed group (Fig. 3A terminal ileum). Although the P-2 response in the FDP-F-fed group was also significantly enhanced (Fig. 3A terminal ileum), the P-1 response was not significantly different compared with the RRC-fed group (Fig. 3B terminal ileum).

In the middle colon, the propionate-evoked P-1 negative $I_{sc}$ response in the STD-F-fed group tended to be reduced compared with the RRC-fed rats ($p<0.10$; Fig. 3B middle colon) although the responses were much smaller than the other segments. The P-2 $I_{sc}$ response in the FDP-F-fed group tended to be reduced compared with the STD-F-fed group ($p<0.10$; Fig. 3A). In the other segments, the propionate-evoked increases in $I_{sc}$ were not significantly different with each other.

**Table 3.** Organic acid concentrations [mM] in cecal contents isolated from RRC-, STD-F- and FDP-F-fed rats.

| SCFAs          | RRC       | STD-F     | FDP-F     |
|----------------|-----------|-----------|-----------|
| Acetate        | 91.0±4.6a | 53.0±5.1b | 78.1±5.5a |
| Propionate     | 19.3±0.9a | 33.0±2.4b | 19.4±4.0a |
| Butyrate       | 52.7±1.5a | 9.2±0.8b  | 21.3±1.5c |
| Valerate       | 3.2±0.2c  | 1.7±0.5b  | 0.0±0.0b  |
| Other organic acids | 0.2±0.0  | 0.3±0.0  | 1.7±0.7  |
| Formate        | 0.2±0.0   | 0.3±0.0   | 1.7±0.7   |
| Lactate        | 0.3±0.0   | 0.3±0.1   | 8.3±5.7   |
| Succinate      | 0.3±0.01  | 0.0±0.0   | 3.0±2.8   |
| Total SCFAs    | 166.2±3.4a| 96.8±5.1b | 118.8±10.8b|
| Total organic acids | 166.9±3.4a| 97.5±5.2b | 131.8±7.0c|

Data are expressed as mean±SE ($n=3$). Different superscript alphabets differ significantly ($p<0.05$) by Tukey test (acetate, propionate, butyrate, total SCFAs and total organic acids) or Games-Howells test (valerate).

**Table 4.** Bacterial densities in wet cecal contents isolated from RRC-, STD-F- and FDP-F-fed rats.

|                     | RRC       | STD-F     | FDP-F     |
|---------------------|-----------|-----------|-----------|
| Bifidobacterium     | 7.10±0.15a| 7.56±0.25a| 10.37±0.09b|
| Lactobacillus       | 9.39±0.14 | 9.51±0.06 | 9.62±0.28 |
| Bacteroides fragilis group | 7.83±0.11a | 10.28±0.10b | 10.02±0.23b |
| Clostridium coccoides group | 9.96±0.01a | 10.42±0.12b | 10.81±0.21b |
| Clostridium leptum subgroup | 8.83±0.08b | 10.25±0.07b | 9.28±0.60ab |
| Enterobacteriaceae  | 6.80±0.28a| 8.04±0.17ab| 8.48±0.56b |
| Enterococci         | 7.15±0.24a| 7.81±0.15a | 8.73±0.11b |
| Total bacteria      | 10.43±0.03a| 11.26±0.02a| 11.44±0.09b|

Data are expressed as mean±SE ($n=3$). Different superscript alphabets differ significantly ($p<0.05$) by Tukey test (Bifidobacterium, Bacteroides fragilis group, Clostridium coccoides group, Enterobacteriaceae, and Enterococci) or Games-Howells test (Clostridium leptum subgroup and total bacteria).

**Serosal CCh-evoked increase in $I_{sc}$**

CCh ($10^{-5} M$), which is an analog of acetylcholine (ACh), the predominant secretagogue in the intestinal glands, evoked an increase in $I_{sc}$ in all preparations (Fig. 1, A), and no significant difference was found among the feeding groups (Fig. S1 (Supplemental Online Material)).

**Concentrations of cecal organic acids**

Organic acid concentrations in cecal contents are shown in Table 3. The total organic acid concentration of cecal contents in the STD-F-fed group was significantly reduced compared with the RRC- and STD-F-fed groups. Although the total organic acid concentration in the FDP-F-fed group was significantly lower than that in the RRC-fed group, the concentration in the FDP-F-fed group was significantly higher than that in the STD-F-fed group. The total SCFA concentrations of cecal contents both in the STD-F- and FDP-F-fed groups were significantly reduced compared with the RRC-fed group. The butyrate and acetate concentrations in the STD-F-fed group were significantly reduced compared with those in the RRC-fed group, but these concentrations in the STD-F-fed group were significantly increased compared with the FDP-F-fed group. Although the butyrate concentration in the FDP-F-fed group was significantly
reduced compared with the RRC-fed group, it was significantly increased compared with the SDT-F-fed group. The acetate concentration in the SDT-F-fed group was also reduced compared with that in the RRC- and FDP-F-fed groups, but the concentration in the FDP-F-fed group was not significantly different from that in the RRC-fed group. On the other hand, the propionate concentration in the STD-F-fed group was significantly increased compared with the RRC- and FDP-F-fed groups, and no significant differences were found between the RRC- and FDP-F-fed groups.

**DISCUSSION**

The present study showed that the consumption of STD-F enhances mucosal stimulant-mediated transepithelial ion transport in compared with RRC in terminal ileum of rats. This reason is unclear, but it might be due that RRC was composed of crude materials like crude grain in compared that STD-F was composed of purified materials. Whereas, FDP-F attenuated the enhancement, moreover FDP-F further attenuated the nervous system-mediated ion transport in cecum. It has been suggested that the inhibitory effects of FDP-F on the intestinal ion transport is possibly due to the changes in the components of gut microbiota and organic acids. These findings suggest that fermented dairy product- and GOS-containing EF might possibly prevent the incidence of diarrhea during EN.

**Effects of diets on electrophysiological parameters**

The basal conditions of electrogenic transepithelial ion transport in all intestinal segments appear not to be changed by the diets, because basal $I_{sc}$ in all intestinal segments were not significantly different among treatments (Table 2). Therefore, the cause of diarrhea during EN was suggested not to be attributed to the change in basal transepithelial ion transport. On the other hand, basal $G_{s}$ in the cecum, middle colon, and distal colon were significantly higher than the others only in the FDP-F-fed group (Table 2). The reason is unknown, but it is indicated to be artificially due to the preparation process of removing contents, which were stickier in the FDP-F-fed group than in the other groups. The mucus layer of the FDP-F-fed rat might be removed more than that of the other groups.

Electrical field stimulation (EFS) to the intestinal mucosa-submucosal preparation activates submucosal and mucosal neurons of the enteric nervous system (ENS), inducing transepithelial anion secretion (29). The EFS-evoked secretory response has been reported to be mediated via ACh and other secretagogue neurotransmitters, predominantly vasoactive intestinal polypeptide (VIP) (30, 31). ACh and VIP directly activate epithelial cells, increase intracellular Ca$^{2+}$ and cAMP, respectively, and then induce anion secretion. The present results suggest that FDP-F-feeding affects the ENS to reduce the intestinal secretion in the cecum and middle colon (Fig. 2), but not reducing that of the epithelial cells, because the direct activation of the epithelium by CCh (Fig. S1) was not affected by the dietary treatment.

Luminal propionate is reported to activate intestinal epithelial cells, releasing ACh from the epithelial cells, and activating neighboring epithelial cells and sensory nerve terminals (32). This induces epithelial secretory response mediated partially through the neural and partially through the non-neural direct pathways (28). The propionate-evoked negative (P-1) and positive (P-2) changes in $I_{sc}$ are due to potassium (K$^{+}$) and anion (Cl$^{-}$ and HCO$_3^-$) secretions, respectively (33). The transepithelial K$^{+}$ and anion secretions contribute in generating the driving force of epithelial fluid secretion. In the present study, STD-F-feeding enhanced the propionate-evoked P-2 change in $I_{sc}$ (anion secretion) in the terminal ileum compared with RRC-feeding (Fig. 3, terminal ileum). The terminal ileum is a segment that absorbs the most water in the intestine, so the enhancement of water secretion in the terminal ileum might possibly induce diarrhea. The cause of diarrhea during EN is possibly due to the higher sensitivity to luminal propionate. On the other hand, the propionate-evoked P-1 change in $I_{sc}$ (K$^{+}$ secretion) in the FDP-F-fed group was significantly reduced compared with the STD-F-fed group (Fig. 3, terminal ileum). Moreover, in the FDP-F-fed group, the propionate-evoked P-1 change in $I_{sc}$ (anion secretion) in the cecum was significantly reduced compared with the RRC-fed group (Fig. 3, cecum), suggesting the prevention of massive fluid secretion in the FDP-F-fed group. It is therefore suggested that STD-F-feeding makes the epithelium more sensitive to luminal propionate. These results suggest that FDP-F-feeding may reduce propionate-evoked fluid secretion by decreasing the sensitivity to propionate in the ileal and cecal epithelia and by inhibitory regulation of ENS in the cecum.

**Analysis of cecal organic acids and microbiota**

It is unclear whether the changes in the electrophysiological activities of intestinal epithelia are regulated by the cecal organic acid concentrations and microbiota. However, the decrease in the luminal total organic acid concentration in both STD-F and FDP-F might induce an increase in the sensitivity of epithelial cells to propionate in the terminal ileum. In the FDP-F-fed group, the concentrations of acetate and butyrate significantly increased, whereas the propionate decreased compared with the STD-F-fed group (Table 3). These differences between the STD-F- and FDP-F-fed groups were possibly involved in the differences in the propionate-evoked responses, but this is unclear given the present results. Further studies are warranted in the future.

In the present analysis of cecal microbiota, the total bacterial densities in the cecal contents in the STD-F- and FDP-F-fed groups increased compared with the
RRC-fed group (Table 4), although the total organic acid concentration in the STD-F-fed and FDP-F-fed groups was decreased (Table 3). The density of *Bifidobacterium* in FDP-F was significantly increased (Table 4), suggesting that the increase in *Bifidobacterium* might contribute to the increase in acetate and butyrate.

**Length of the intestines**

Colonic mucosal atrophy is induced by liquid elemental diet in rats (34), and epithelial proliferation is decreased in the colon of rats fed a liquid diet (35). Although I did not perform histological analysis of mucosa, but measured the lengths of small and large intestines. The results showed that the length of large intestine was shorter in the STD-F-fed rats, but not in the FDP-F-fed rats compared to the RRC-fed rats (Table 1).

SCFAs, especially butyrate, are known as a trophic substance to the intestinal mucosa (36, 37). The trophic effect of FDP-F is suggested to be possibly due to the increase in SCFA, especially butyrate production in the large intestine. We previously have reported that the SCFA receptors, FFA2 (GPR43) and FFA3 (GPR41), are expressed in the intestinal epithelium, particularly peptide YY (PYY) and/or glucagon-like peptide 1 (GLP-1) containing L-type enteroendocrine cells in the ileum and colon of humans and rats (38–40). GLP-2, which is produced from the same precursor with GLP-1, is known to have a trophic effect that is highly specific for the GI tract (41). In addition, it has been reported that GLP-2 increases a length of small intestine in parenterally fed rats with short bowel syndrome (42). Therefore, it is suggested that the increase in luminal SCFA enhances GLP-2 endocrine, inducing the increase in the length of the large intestine.

**Conclusions**

The present study showed that STD-F consumption increased the sensitivity for luminal propionate. Moreover, FDP-F consumption decreased the secretory nerve activity in the cecum, proximal colon, and middle colon, and the sensitivity for propionate in the cecum. These might be due to the total amounts and components of gut microbiota and organic acids, and inhibitory changes in mucosal luminal stimulant- and nervous system-mediated fluid secretory function.

**Disclosure of state of COI**

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**Supporting information**

Supplemental online material is available on J-STAGE.

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