



Title	Two-week feeding of difructose anhydride III enhances calcium absorptive activity with epithelial cell proliferation in isolated rat cecal mucosa
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Citation	Nutrition, 22(3), 312-320 <a href="https://doi.org/10.1016/j.nut.2005.06.015">https://doi.org/10.1016/j.nut.2005.06.015</a>
Issue Date	2006-03
Doc URL	<a href="http://hdl.handle.net/2115/7404">http://hdl.handle.net/2115/7404</a>
Type	article (author version)
File Information	Nutrition22_3.pdf



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1 **ABSTRACT**

2

3 **OBJECTIVE:** Ingestion of Diffructose Anhydride (DFA) III enhances Ca absorption  
4 in rats. The aim of the present study was to determine the mechanism involved in  
5 increased Ca transport by DFA III ingestion. The short-term and long-term effects of  
6 DFAIII feeding on Ca transport were determined using isolated epithelium from the  
7 small and large intestine in rats.

8 **METHODS :** Male Sprague-Dawley rats were fed an 8% cellulose or 5% cellulose  
9 plus 3% DFAIII diet for 14 days. Net epithelial Ca transport in the small intestine,  
10 cecum and colon were compared between the two diet groups using an Ussing chamber.  
11 The contents and the epithelial tissues in the cecum were analyzed.

12 **RESULTS:** There were no differences in basal and luminal DFAIII-induced Ca  
13 transport in the isolated small intestinal and colonic mucosa between the two diet  
14 groups. Basal and lamina DFAIII-induced Ca transport in the cecum in the DFAIII fed  
15 group was higher than that in control group. A decrease in pH, and increases in Ca  
16 pools, short-chain fatty acids (SCFAs) or organic acids in the cecal contents and in the  
17 depth and number of cells in crypts in the cecal tissue were observed.

18 **CONCLUSIONS:** The increase in Ca transport involved two mechanisms, the  
19 presence of DFAIII in the small intestine directly affected the epithelial tissue and  
20 caused to increase Ca absorption as a short-term effect, and the degradation of DFAIII  
21 by microbial fermentation produced SCFAs and subsequently enhanced Ca absorption  
22 in the large intestine as a long-term effect, respectively.

23

24 **KEY WORDS:** Ca absorption, indigestible saccharide, Ussing chamber, intestine, rats

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## 1 INTRODUCTION

2

3 Difuctose anhydride (DFA) III (Fig. 1), a newly developed indigestible saccharide,  
4 consists of two fructose residues and is prepared from inulin with *Arthrobactor* sp.  
5 H65-7 fructotransferase (1). A procedure was established for the mass production of  
6 DFAIII using a bacterial enzyme (1) and the products of DFAIII have already been sold  
7 in Japan. Ingestion of DFAIII (2) and its isomer, DFAIV (3), increases Ca absorption  
8 in male rats as demonstrated in in vivo balance studies. The appearance Ca absorption  
9 was significantly higher in rats fed DFAIII than in rats wed well-known indigestible  
10 oligosaccharides such as raffinose or fructooligosaccharides (2). Feeding DFAIII has  
11 restored Ca absorption in female rats after ovariectomy, which induces ovarian hormone  
12 deficiency followed by a reduction in Ca absorption (4). DFAIII increases the Ca  
13 content, and the breaking force and mineral density of bone in exercised rats (5). Thus,  
14 the ingestion of DFAIII might play a beneficial role in the absorption and retention of  
15 Ca in the body.

16 In in vitro experiments, the application of DFA III or DFA IV in the mucosal  
17 medium was found to stimulate Ca absorption in everted sacs of the rat small intestine  
18 (2, 3) or colon (4). The application of DFAIII and DFAIV in the luminal medium was  
19 also found to increase net Ca absorption in the isolated epithelium of the small and large  
20 intestine in rats using the Ussing chamber technique (6). These findings suggest that  
21 DFAIII flowing into the lumen directly affects the epithelium of the small and large  
22 intestine and promptly enhances Ca transport. In our recent report, Ca transport in the  
23 isolated intestinal epithelium was increased by luminal DFAIII application and this  
24 increase in Ca absorption occurred via the paracellular route, which is regulated by tight  
25 junctions (TJs) (7).

26 In the above-mentioned in vivo balance studies using rats (2,4), DFAIII was  
27 repeatedly ingested for at least 2-4 weeks and the small and/or large intestinal mucosa

1 were continuously exposed to this sugar throughout the feeding period. The transport  
2 and absorption system of the intestinal mucosa might have adapted to the continuous  
3 stimulation of DFAIII in vivo, since the ingestion of DFAIII has been reported to  
4 increase the weight of the cecal wall in rats (2, 4). In addition, feeding DFAIII is  
5 reported to improve microbial fermentation and produce short-chain fatty acids  
6 (SCFAs) or other organic acids in the cecum in rats (2, 4).

7 In our latest report, ingestion of a 3%DFAIII diet for 14 d enhanced Ca absorption  
8 via intercellular passage in the isolated cecal epithelium in rats (8). This increase in  
9 Ca transport was not occurred in the small intestinal or colonic mucosa isolated from  
10 the same rats fed 3%DFAIII diet (8). In the cecum, which is thought to be a main  
11 portion of microbial fermentation, adaptive changes in its function should be induced by  
12 DFAIII ingestion for an extended period.

13 Thus, the aim of the present study was to clarify the mechanism for increase in Ca  
14 absorption through the small and large intestine in rats fed DFAIII for 2 weeks. As  
15 DFAIII itself directly stimulates Ca absorption from the intestine, Ca transport in the  
16 isolated epithelial tissue of the small and large intestine was compared with or without  
17 applying DFAIII into the mucosal medium using Ussing chamber system (9). In  
18 addition, since DFA III is known to be fermented in the cecum in rats, differences in  
19 morphological changes in the cecal epithelium, and pH, Ca pools and SCFA  
20 concentrations in the cecal contents were compared between in rats fed control diet and  
21 in rats fed DFAIII diet.

22

## 23 **MATERIALS AND METHODS**

24

25 This study was approved by the Hokkaido University Animal Committee and the  
26 animals were maintained in accordance with the Hokkaido University guidelines for the  
27 care and use of laboratory animals.

1

## 2 **Animals and diets**

3

4 Sixteen male Sprague-Dawley rats (4 wk old; Japan SLC, Shizuoka, Japan) were  
5 housed in individual stainless-steel metabolic cages with wire-mesh bottoms. The  
6 cages were placed in a room with controlled temperature (22-24°C), relative humidity  
7 (40-60%) and lighting (light 0800-2000 h). The animals were fed the stock diet shown  
8 in Table 1 for an acclimation period of 14 days and then assigned to two groups based  
9 on body weight. Compositions of the stock (control) and test diets are shown in Table  
10 1. DFA III was kindly provided by Nippon Beet Sugar MGF., Ltd. (Obihiro, Japan).  
11 One group (control group) was fed the control diet, while the other group (DFAIII  
12 group) was fed the test diet containing 30g of DFAIII per kilogram of diet. Rats were  
13 fed the assigned test diets for 14 days. All animals were allowed free access to  
14 deionized water throughout the test period. Body weight and food intake were  
15 measured every day.

16

## 17 **Sampling of the intestinal contents and tissue preparation**

18

19 On the last day of the experiment, the rats were anesthetized with pentobarbital sodium  
20 (30 mg/kg). The entire length of the small intestine from the ligament of Treitz to the  
21 ileocecal junction, the entire cecal sac, and the entire length of the colon were quickly  
22 removed. The contents of the small intestine, cecum and colon were collected,  
23 weighed and placed into test tubes. The intestinal contents were stored at -80 °C until  
24 future analysis. The outside and inside surface of the isolated intestine was washed  
25 with ice-cold (4 °C) saline (154 mmol/L NaCl), and the wet weight of each specimen  
26 was measured. Segments of the small intestine (a 6-cm section from the center),  
27 cecum (whole sac) and colon (a 6-cm section from the center) were quickly isolated,

1 and each specimen was then cut open along the mesenteric border to produce a flat  
2 sheet, and rinsed with ice-cold HEPES buffer solution (HBS). The HBS used in this  
3 study consisted of 125 mmol/L NaCl, 4 mmol/L KCl, 6 mmol/L L-glutamine, 30  
4 mmol/L 2-[4-(2-hydroxymethyl)-1-piperazyl] ethanesulfonic acid (HEPES), and 1.25  
5 or 10 mmol/L CaCl<sub>2</sub>·2H<sub>2</sub>O, and it was gassed with 100% O<sub>2</sub> to maintain a constant pH  
6 of 7.4. HBS containing 1.25 mmol/L Ca was used as the bathing solution for the  
7 serosal component during the experiment and as a stabilizing solution for both the  
8 mucosal and the serosal sides between experiments. The serosa and muscle layers  
9 were removed, and stripped preparations, consisting of the mucosa and the submucosa,  
10 were mounted onto Ussing chambers (Diffusion chamber system, Corning Costar Co.,  
11 Cambridge, UK) that exposed a circular area of the epithelium of 0.64 cm<sup>2</sup>. The  
12 serosal and mucosal sides of the segments were bathed in 1 mL of HBS containing 1.25  
13 mmol/L Ca and continuously exposed to 100% O<sub>2</sub> gas. After a 30-min stabilization  
14 period, the medium on both sides of the tissue was removed by aspiration and 1 mL  
15 portions of the appropriate solutions were added to the mucosal and serosal sides in the  
16 chamber system.

17

#### 18 **Saccharide and permeable marker**

19

20 DFAIII is soluble in water at 100 mmol/L. In our preliminary evaluation using the  
21 same chamber system, a maximal Ca transport response was obtained with 100mmol/L  
22 DFAIII on the luminal side (6,7). Lucifer Yellow CH dilithium salt (LY: FW 457.2,  
23 Sigma Chemical Co., St. Louis, MO) was used as the paracellular permeable marker  
24 (10) at a concentration of 21.8 μmol/L in the 10 mmol/L Ca-HBS mucosal medium.  
25 The pH of 10 mmol/L Ca plus LY-HBS containing any of these sugars at 0 (control) or  
26 100 mmol/L was found to remain constant at 7.4.

27

## 1 **Experimental procedure**

2

3 Fresh HBS containing 1.25 mmol/L Ca was applied to the serosal bath and 10 mmol/L  
4 Ca-HBS with or without 100 mmol/L DFAIII was applied to the mucosal bath in the  
5 experiments using the small intestine, cecum and colon preparations. After a 30-min  
6 incubation period, the serosal solution was transferred to a polyethylene test tube.

7

## 8 **Morphological index of cecal epithelium**

9

10 The cecal preparation for morphological observation was obtained by cutting a 6 mm  
11 wide strip of tissue along the circumference of the cross section of the cecum at 3/4 of  
12 the distance between the ileocecal valve to the end of the cecum. The strip was fixed  
13 in a 3.3 mmol/L buffered formaldehyde solution for future histological analysis. The  
14 fixed 6-mm cecal segments were cut into 7- $\mu$ m cross-sections on a glass slide using  
15 cryostat, and stained with hematoxylin and eosin. Only 10 well-oriented crypts, in  
16 which the base, lumen and the top of the crypt could all be seen, were selected per  
17 segment of the cecum. The consistency of the crypts was obtained by positioning the  
18 micrometer along the luminal surface and counting the number of complete crypts per 1  
19 mm. The depth of the crypts from the top to the base was measured in  $\mu$ m, and the  
20 number of cells in the crypt was counted (cells/crypt) using an optical microscope.  
21 The electrical portrait of the whole cross section of the cecal tissue was obtained and  
22 analyzed by computer system. The outside circumference along serosa and the inside  
23 circumference along the top of the crypts in the cecal section, which is expected to  
24 indicate the inner area of the lumen, was measured using soft ware program (Scino  
25 Image, Scino corptation, USA). The two parameters are expected to indicate the outer  
26 and inner area of the cecum, respectively.

27



## 1 **Bacterial enumeration in cecal content**

2

3 The population of anaerobic bacteria in the cecal contents was compared in rats fed the  
4 control and DFAIII diets. About 0.2 g (wet weight) of cecal content from one rat fed  
5 the control or DFAIII diet was immediately transferred into 1.8 ml of diluting solution  
6 containing (g/l):  $\text{KH}_2\text{PO}_4$ , 4.5;  $\text{Na}_2\text{HPO}_4$ , 6.0; L-cystein-HCl, 0.5; tween 80, 0.5; and  
7 Bacto Agar (DIFCO, USA), 1.0. The samples were kept under anaerobic conditions  
8 using Anaero Pack (Mitsubishi Gas Chem. Co. Inc. Japan) during transportation to our  
9 laboratory and were then spread on GAM agar plates containing (g/l): peptone, 10.0;  
10 soy peptone, 3.0; protease peptone, 10.0; hydrolyzed serum powder, 13.5; yeast extract,  
11 5.0; meat extract, 2.2; liver extract, 1.2; glucose, 3.0;  $\text{KH}_2\text{PO}_4$ , 2.5; NaCl, 3.0; soluble  
12 starch, 5.0; L-cystein-HCl, 0.3; thioglycolate-sodium salt, 0.3; and agar, 20.0. The pH  
13 of medium was adjusted to 7.1 before autoclaving. The bacterial population was  
14 counted after 2 days incubation at 37°C in an anaerobic chamber (Coy Laboratory  
15 Products, Inc. USA) and expressed as colony forming units (CFUs).

16

## 17 **Analyses**

18

19 The Ca concentrations in the serosal medium were measured by a colorimetric method  
20 using a commercial kit (Calcium C-Test, Wako Chemical Co., Osaka, Japan). Intra-  
21 and inter-assay coefficients of variation for Ca determination were below 2% (stock  
22 standard sample ) and 9% (same sample at 5 assays). The net  
23 transepithelial passage of Ca was expressed as nmol Ca transferred per min per square  
24 cm of surface area. The LY in the serosal solution was determined fluorometrically at  
25 430nm for excitation and 540nm for emission (FP-550, JAS.Co., Tokyo, Japan) after  
26 appropriate dilution of the solution with purified water. The contents of the small  
27 intestine, cecum or colon were diluted 4 vol of deionized water and homogenized with a

1 Teflon homogenizer. The pH of the intestinal homogenate was measured with a  
2 semi-conducting electrode (ISFET pH sensor 0010-15C, HORIBA, Ltd., Kyoto, Japan)  
3 as intestinal contents. Amount of total Ca in the homogenates were determined by  
4 atomic absorption spectrometry (Polarized Zeeman Atomic Absorption  
5 Spectrophotometer, A-5310, HITACHI, Tokyo, Japan) after wet-ashing with an acid  
6 mixture. Soluble Ca in the supernatant obtained after centrifugation (30000g for 20  
7 min at 4 °C) of the homogenate was determined by the above-mentioned atomic  
8 absorption spectrophotometry after deproteinization with 9 mol/L of perchloric acid.  
9 Concentrations of short-chain fatty acids (SCFAs) and organic acids (acetic, propionic,  
10 butyric, succinic and lactic acids) in the homogenates of the cecal and colonic contents  
11 were measured according to the method of Hoshi et al. (11) using high-performance  
12 liquid chromatography (LC-10 ADvp SHIMAZU, Kyoto, Japan) equipped with two  
13 Shima-packSCR-102H columns (8mm inner diameter and 30 cm long; SHIMAZU,  
14 Kyoto, Japan) and an electroconductibility detector (CDD-6A, SHMAZU).

15

## 16 **Statistical analyses**

17

18 All results are expressed as means  $\pm$  SEM (n=8). Statistical analyses were performed  
19 by unpaired t-test for comparison of values between the two diet groups. For  
20 comparison of the two diet groups (control and DFAIII diets, D), luminal application of  
21 DFAIII in the mucosal medium in vitro (with or without 100mmol/L DFAIII in the  
22 mucosal chamber, L), and interaction of (D) x (L), analyses were performed by  
23 two-way ANOVA following Duncan's multiple range test. A difference with  $P<0.05$   
24 was taken to be statistically significant. Regression analysis was used for analysis of  
25 the relationship between two factors.

26

## 27 **RESULTS**

1

2 There were no significant differences in initial and final body weights, body weight gain  
3 or food intake over the experimental period between groups fed the control and DFAIII  
4 diets (Table 2). Similarly, there were no differences in the wet weights of the small  
5 intestine or colon, or in their contents between the two diet groups (Table 3). The wet  
6 weights of the cecum ( $P < 0.005$ ) and cecal contents ( $P < 0.001$ ) in rats fed the DFAIII  
7 diet were, however, significantly greater than those in rats fed the control diet. The pH  
8 in the contents of the small intestine was similar between the two diet groups, while the  
9 pH in cecal and colonic contents were lower ( $P < 0.001$ ) in rats fed the DFAIII diets  
10 than in rats fed the control diets.

11 Regarding the morphological indices of the cecal epithelium (Table 4), the  
12 outside ( $P < 0.05$ ) and luminal circumference ( $P < 0.01$ ), and the ratio of luminal:  
13 outside circumference ( $P < 0.005$ ) of the cecal tissue preparations from rats fed the  
14 DFAIII diet were greater than those in rats fed the control diet. The depth of crypt ( $P$   
15  $< 0.001$ ) and the number of cells per crypt ( $P < 0.005$ ) in the DFAIII groups were  
16 significantly greater than those in the control diet groups. There were no significant  
17 differences in the width of epithelial cells in the crypts or the number of crypts per 1  
18 mm of cecal epithelial circumference between the two diet groups.

19 There were no significant differences in total and soluble Ca pools, soluble Ca  
20 concentration, or the ratio of soluble Ca pool in the contents of the small intestine and  
21 the colon between the two diet groups (Table 5). The total ( $P < 0.001$ ) and soluble Ca  
22 pools ( $P < 0.001$ ), and soluble Ca concentration ( $P < 0.05$ ) in the cecal contents of rats  
23 fed the DFAIII diets were higher than those in the rats fed the control diets. There  
24 were no significant differences in the percentage of the soluble Ca pool to total Ca pool  
25 in the cecal contents between the two diet groups.

26 There were no differences in basal (without luminal DFAIII application) net Ca  
27 absorption or LY permeability in the small intestinal mucosa between rats fed the

1 control diet and rats fed DFAIII diet (Fig.2AB). Application of 100 mmol/L DFAIII  
2 in the mucosal medium enhanced net Ca transport and LY passage in the small  
3 intestinal preparations in both diet groups ( $P < 0.05$ ). However, there were no  
4 differences in the increase in net Ca absorption or LY permeability induced by luminal  
5 application of DFAIII between the two diet groups.

6 For the large intestinal epithelial preparations, luminal application of DFAIII  
7 affected the net Ca absorption and LY permeability in the cecum (Fig. 3AB). Basal Ca  
8 transport in the cecal epithelium in rats fed the DFAIII diets was higher than that in rats  
9 fed the control diets. There were no differences in basal LY permeability between rats  
10 fed the control and the DFAIII diets. Application of DFAIII in the mucosal medium  
11 enhanced net Ca transport and LY passage in the cecum of both diet groups ( $P < 0.05$ ).  
12 The degree of the increase in net Ca absorption and LY passage were greater in the rats  
13 fed the DFAIII diet than in rats fed the control diet ( $P < 0.05$ ).

14 There were no differences in basal net Ca absorption or LY permeability in the  
15 colonic mucosa between rats fed the control diet and rats fed DFAIII diet (Fig. 4AB).  
16 Application of 100 mmol/L DFAIII in the mucosal medium enhanced net Ca transport  
17 and LY passage in the colonic preparations in both diet groups ( $P < 0.05$ ). However,  
18 there were no differences in the increase in net Ca absorption or LY permeability  
19 induced by luminal application of DFAIII between two the diet groups.

20 Regression analysis for the relation between the net Ca absorption and LY  
21 permeation was evaluated for the three portions of the intestine, respectively (Fig. 2-4C).  
22 Statistically significant ( $P < 0.001$ ) positive linear relations were observed between net  
23 Ca absorption and LY passage in the small intestine, cecum and colon isolated from the  
24 rats fed the control and the DFAIII diets.

25 The amount of acetic and propionic acid, total SCFA (sum of acetic, propionic,  
26 n-butyric, iso-butyric, n-valeric and iso-valeric acids), succinic and lactic acid in the  
27 cecal contents were higher ( $P < 0.001, 0.005$  or  $0.01$ ) in rats fed the DFAIII diets than in

1 rats fed the control diets (Table 6). However, there were no differences in the amount  
2 of main (acetic, propionic and butyric acid) and total SCFA, and lactic acid in the cecal  
3 contents between the control and test diet groups (Table 7). Anaerobic bacterial  
4 population in the cecal contents in one rat fed the control or DFAIII diet was  $2.8 \times 10^9$   
5 (control) and  $2.0 \times 10^{10}$  CFU (DFAIII), respectively.

6 The relation between net Ca absorption without luminal DFAIII application in  
7 chamber system (basal Ca transport) and some factors in the cecum or the cecal  
8 contents in rats fed control and DFAIII diets were evaluated by regression analysis  
9 (Table. 8). There was positive correlation between the basal Ca transport and the  
10 weight of the cecal wall, luminal circumference of cecal section, the ratio of lumen:  
11 outside length of section, the depth of crypt, or the number of cells per crypt height.  
12 There was a negative correlation between the Ca absorption rate in and the pH of the  
13 cecal contents. The Ca absorption in the cecal epithelium was correlated with the  
14 concentration of acetic, propionic or iso-butyric acids, total SCFA, succinic, and lactic  
15 acid in the cecal contents. There were no significant relations between the wet  
16 weight of cecal contents and Ca transport in cecal epithelium.

17

## 18 **DISCUSSION**

19

20 In the present study, ingestion of DFAIII for 14 days did not affect growth, food intake,  
21 or weight of the small intestine or colon in rats. The DFAIII feeding also have no  
22 affect on net absorption of Ca and permeability of LY in the isolated epithelium of the  
23 small intestine or colon. Repeated ingestion of DFAIII did increase in cecal weight,  
24 and net Ca transport and LY permeation in the isolated cecal epithelium. Positive  
25 linear regressions were obtained between net Ca absorption and LY permeation in the  
26 small intestine, cecum and colon ( $P < 0.001$ ). LY is reported to act as a marker of  
27 paracellular passage, in which is regulated by TJs (10). Thus, the increase in Ca

1 transport and LY passage with repeated feeding of DFVIII is thought to occur via the  
2 paracellular route in the cecal tissue. These changes induced by the feeding of DFVIII  
3 are similar to those in our latest report (8). The repeated ingestion of DFVIII affects  
4 the ability of the epithelial tissue in the cecum in rats.

5 Morphological indices of the cecal mucosa, such as the luminal and outer  
6 circumference of the cross section, depth of crypts, and the cell population of crypts,  
7 were also increased with DFVIII feeding. The increase in the weight of the cecal wall  
8 was due to an increase in the population of epithelial cells and their supporting tissue  
9 such as submucosal tissue and smooth muscle layer. At the same time, a decrease in  
10 pH and increases in acetic, propionic, total SCFA, succinic, and lactic acid  
11 concentrations were observed in the cecal contents. The feeding of DFVIII obviously  
12 affects the microbial environment and enhances the fermentation process in the cecum.

13 Although repeated feeding of DFVIII affects Ca transport activity and LY  
14 permeation in the cecum, this effect may be also induced by changes in the paracellular  
15 route via TJs in the epithelial tissue itself. An adaptation to enhance Ca transport  
16 ability might, therefore, occur in the epithelial tissue of the cecum after feeding of  
17 DFVIII for extended periods. Positive linear relations were obtained between the basal  
18 Ca transport and the depth of crypt, the number of cells per crypt and the intra-luminal  
19 circumference. Thus, the absorptive area in the cecal tissue was substantially  
20 increased and these morphological changes may be important factors in the increase in  
21 the efficiency of Ca transport in rats fed DFVIII diet for 14 days.

22 Ingestion of indigestible saccharides, such as sugar alcohols (12,13),  
23 oligosaccharides (14,15) and polysaccharides (16,17), has also been reported to increase  
24 Ca absorption in rats during balance studies in vivo. The effect of DFVIII ingestion on  
25 Ca absorption has also been reported in rats in vivo (2,3). To date, there are two  
26 aspects of Ca absorption known to be improved by indigestible saccharides; the direct  
27 effect of the sugar itself in the small and/or large intestine and the indirect effect of the

1 SCFAs or organic acids produced from indigestible sugars by microbial fermentation in  
2 the large intestine.

3 Intraluminal infusion of sorbitol increases Ca absorption in the small intestine of  
4 rats (18). Using a tracer technique with  $^{45}\text{Ca}$ , maltitol was found to stimulate Ca  
5 absorption in rats (19). In in vitro experiments using everted sacs or isolated  
6 preparations, maltitol (20), and polydextrose (21) were also seen to stimulate Ca  
7 absorption in the small intestine of rats. Various indigestible sugars were also found to  
8 increase net Ca absorption in the isolated epithelium of small and large intestine in rats  
9 using an Ussing chamber technique in vitro (6). Using everted sacs, DFA III in the  
10 mucosal medium has been found to stimulate Ca transport from the mucosal side to the  
11 serosal side in the rat jejunum, ileum (2) and colon (4). Our preliminary experiment  
12 using the same chamber system also revealed that luminal application of DFAIII  
13 increased Ca absorption in isolated epithelial tissue of the small and large intestine in  
14 rats (6,7). The present study also revealed that luminal application of DFAIII directly  
15 affected the epithelium itself, and increased Ca transport in the small and large intestine  
16 in rats.

17 Improved fermentation of resistant carbohydrates is also thought to be a factor in  
18 promoting transepithelial Ca transport in the large intestine (22,23). SCFAs and other  
19 organic acids (lactic acid, succinic acid, etc.) produced through microbial fermentation  
20 decrease the pH of the cecal contents and thereby convert insoluble Ca to a soluble ionic  
21 form (22,24). Indeed, in the present study, a decrease in the pH, and increases in the  
22 soluble Ca pools and the soluble Ca concentrations were observed in the cecal contents  
23 in rats fed the DFAIII diets.

24 In balance studies in vivo, a positive correlation exists between apparent Ca  
25 absorption and total SCFA, acetic, propionic and butyric acid concentrations in the  
26 cecum of rats fed oligosaccharides (25) or soluble soybean fiber (26). The infusion of  
27 SCFAs into the colon or rectum is known to stimulate Ca absorption in humans in vivo

1 (27,28). Luminal perfusion of SCFAs also enhances Ca absorption in the colon in rats  
2 (29). In our previous study using the same chamber technique, luminal application of  
3 SCFA enhanced net Ca transport from the mucosal side to the serosal side in isolated rat  
4 epithelial tissue in vitro (30). These in vivo and in vitro experiments indicate that  
5 SCFAs in the lumen promptly stimulate Ca transport in the epithelial tissue of the large  
6 intestine. However, in this experiment, the isolated cecal epithelial tissue was not  
7 exposed to SCFAs in the chamber system. Thus, changes in Ca transport activity,  
8 especially in the basal conditions, must have already occurred before isolation from the  
9 body.

10 SCFAs are an important factor in the stimulation of epithelial cells and the  
11 promotion of cell proliferation in the large intestine. Intraluminal infusion of SCFAs  
12 increased the relative weight of the mucosa and submucosa, crypt size, and mitotic  
13 index in the cecum in rats (31). Mucosal cell proliferation in the crypts was raised by  
14 the addition of SCFAs in vitro using isolated human cecal epithelium obtained through  
15 biopsy (32). In the present study, Ca transport activity in the cecal tissue was  
16 correlated to the weight of the cecal wall, depth of crypts, and outer circumference of  
17 the cecum, as well as with the concentration of some fermentation products (acetic acid,  
18 propionic acid, iso-butyric acid, total SCFA, succinic acid and lactic acid) in the cecal  
19 contents. Lactic acid stimulated mitosis of the cecal epithelial cell in rats (31). To  
20 date, there is no information about the trophic effect of succinic acid on cell  
21 proliferation in the gastrointestinal tract.

22 In balance studies with rats, cecectomy or ingestion of antibiotic drugs  
23 completely inhibited the increase in Ca absorption induced by the feeding of  
24 fructooligosaccharides or galactooligosaccharides, which are commonly used  
25 indigestible saccharides that affect Ca retention in the body (33,34). On the other  
26 hand, cecectomy only partially inhibited Ca absorption induced by DFAIII ingestion in  
27 rats in vivo (4). These results suggest that the main portion of the intestine for Ca



1 absorption may vary according to the indigestible saccharide.

2           In the present study, the increase in Ca absorption induced by repeated feeding  
3 of DFAIII may have occurred via two different mechanisms in the small and large  
4 intestine in vivo. Firstly, DFAIII flowing into the lumen of the intestine directly  
5 affected the epithelial tissue, and promoted Ca absorption. Prompt and direct action of  
6 DFAIII mainly occurs in the small intestine and to a minor extent in the large intestine  
7 in vivo because DFAIII is thought to be degraded by microbes. Secondly, chronic and  
8 indirect action of DFAIII occurred in the large intestine, particularly in the cecum in rats.  
9 DFAIII has a prebiotic effect generally induced by ingestion of indigestible saccharides  
10 via microbial fermentation. The increased population of epithelial cells and  
11 intra-luminal area induced by the increase in SCFA production are important factors in  
12 the increased Ca transport in the large intestine. The increase in total and soluble Ca  
13 pools, and the stimulatory effect of SCFAs on Ca transport also may contribute the  
14 increase in Ca absorption from the epithelial tissue of the large intestine. The  
15 paracellular route via TJs is involved in the promotion of diffusional Ca transport in  
16 both cases. Thus, in conclusion, DFAIII is thought to increase Ca transport by  
17 short-term and/or long-term mechanisms in the small and large intestine, and is thought  
18 to benefit Ca absorption and retention in the body.

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TABLE I

## COMPOSITION OF CONTROL AND TEST DIETS

Element	Control diet (g)	Test diet (g)
Casein <sup>1</sup>	200	200
Dextrin <sup>2</sup>	400	400
Soybean oil	70	70
Mineral mixture <sup>3</sup>	35	35
Vitamine mixtture <sup>3</sup>	10	10
Choline bitartate	2.5	2.5
L-cystine	3.0	3.0
Tert- butylhydroquinone	0.014	0.014
Crystallized cellulose <sup>4</sup>	80	50
DFAIII <sup>5</sup>		30
Sucrose	to make 1000 g	

1. Casein (ALACID; New Zealand Daily Bond, Wellington, New Zealand)

2. Dextrin (Pine-dex 4; Matsutani Chemical Industry, Tokyo, Japan)

3. Mineral and vitamin mixture were prepared according to the AIN-93G formulation

4. Crystallized cellulose (AVICEL; Asahi Kasei Ltd. Osaka, Japan)

5. DFAIII (Nippon Beet Sugar MGF., Ltd. Obihiro, Japan)

TABLE II

INITIAL AND FINAL BODY WEIGHT, BODY WEIGHT GAIN, AND FOOD  
INTAKE OF RATS FED THE CONTROL OR DFAIII DIET FOR 14 DAYS

	Control diet	DFAIII diet	<i>P</i> -value
Initial body weight (g)	213.1 ± 4.9	211.6 ± 6.1	0.848
Final body weight (g)	307.4 ± 7.4	307.4 ± 9.5	0.997
Body weight gain (g/d)	6.7 ± 0.3	6.7 ± 0.3	0.815
Food intake (g/d)	19.2 ± 0.5	18.9 ± 0.5	0.621

Values are mean ± SEM (n = 8). The values of initial and final body weight, body weight gain, and food intake were analyzed by unpaired *t*-test. Asterisk mean significant difference (*P* < 0.05).

TABLE III

WET WEIGHTS OF ISOLATED TISSUES AND THEIR CONTENTS, AND pH  
IN THE CONTENTS OF THE SMALL INTESTINE, CECUM AND COLON IN  
RATS FED THE CONTROL OR DFAIII DIET FOR 14 DAYS

	Control diet	DFAIII diet	<i>P</i> -value
Isolated tissue (g)			
Small intestine	6.40 ± 0.16	6.92 ± 0.30	0.154
Cecum	1.29 ± 0.08	1.66 ± 0.06*	0.003
Colon	1.03 ± 0.05	1.11 ± 0.05	0.305
Contents (g)			
Small intestine	1.05 ± 0.11	1.05 ± 0.09	0.785
Cecum	2.42 ± 0.20	4.64 ± 0.52*	0.001
Colon	1.10 ± 0.16	0.67 ± 0.15	0.075
pH in contents			
Small intestine	7.86 ± 0.16	7.99 ± 0.09	0.472
Cecum	7.91 ± 0.09	6.97 ± 0.13*	< 0.001
Colon	7.89 ± 0.07	7.47 ± 0.05*	< 0.001

Values are mean ± SEM (n = 8). The values of wet weight of isolated tissues and their contents, and the pH in the contents of the small intestine, cecum and colon were analyzed by unpaired *t*-test. Asterisks mean significant difference (*P* < 0.05).



TABLE IV

MORPHOLOGICAL INDICES OF ISOLATED EPITHELIAL TISSUE IN  
THE CECUM IN RATS FED THE CONTROL OR DFAIII DIET FOR 14 DAYS

	Control diet	DFAIII diet	<i>P</i> -value
Outer circumference (O)(mm)	13.7 ± 1.2	17.5 ± 1.2*	0.040
Inner circumference (I)(mm)	22.1 ± 2.1	36.6 ± 4.1*	0.007
Ratio of (I)/ (O)	1.61 ± 0.07	2.06 ± 0.12*	0.004
Crypt dept (µm)	195.2 ± 4.0	232.2 ± 4.0*	< 0.001
Number of cells/ crypt	32.8 ± 0.5	38.6 ± 1.0*	0.001
Width of one cell (µm)	6.0 ± 0.2	6.1 ± 0.2	0.776
Number of crypt/(mm)	19.2 ± 0.7	18.1 ± 0.4	0.201

Values are mean ± SEM (n = 8). The values of outer circumference (O), inner circumference (I), ratio of (I)/(O), crypt dept, number of cells / crypt, width of one cell, and number of crypt / mm in the cecal tissue were analyzed by unpaired *t*-test. Asterisks mean significant difference (*P* < 0.05).

TABLE V

TOTAL AND SOLUBLE CA POOL, SOLUBLE CA CONCENTRATION AND THE RATIO OF SOLUBLE CA POOL IN THE CONTENTS OF THE SMALL INTESTINE, CECUM AND COLON IN RATS FED THE CONTROL OR DFAIII DIET FOR 14 DAYS

	Control diet	DFAIII diet	<i>P</i> -value
Total Ca pool ( $\mu\text{mol}$ in contents)			
Small intestine	114.6 $\pm$ 13.1	148.2 $\pm$ 15.9	0.125
Cecum	421.2 $\pm$ 45.1	862.9 $\pm$ 77.6*	< 0.001
Colon	324.0 $\pm$ 44.6	272.6 $\pm$ 53.8	0.474
Soluble Ca pool ( $\mu\text{mol}$ in contents)			
Small intestine	34.9 $\pm$ 6.8	39.5 $\pm$ 3.9	0.565
Cecum	94.1 $\pm$ 8.8	193.6 $\pm$ 16.7*	< 0.001
Colon	72.1 $\pm$ 9.9	63.5 $\pm$ 14.5	0.631
Soluble Ca concentration ( $\mu\text{mol/g}$ contents)			
Small intestine	31.8 $\pm$ 4.5	36.8 $\pm$ 3.0	0.376
Cecum	37.5 $\pm$ 3.4	48.7 $\pm$ 3.3*	0.031
Colon	58.5 $\pm$ 5.1	72.9 $\pm$ 5.7	0.081
Soluble Ca pool (%)			
Small intestine	29.1 $\pm$ 2.2	27.4 $\pm$ 1.7	0.560
Cecum	22.7 $\pm$ 0.7	22.5 $\pm$ 0.4	0.854
Colon	22.2 $\pm$ 0.3	22.6 $\pm$ 1.0	0.701

Values are mean  $\pm$  SEM (n = 8). The values of total and soluble Ca pools, soluble Ca

1 concentration and the ratio of soluble Ca pool (100 x soluble Ca pool /total Ca pool) of  
2 the small intestine, cecum and colon were analyzed by unpaired *t*-test. Asterisks mean  
3 significant difference ( $P < 0.05$ ).

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TABLE VI

AMOUNT OF EACH AND TOTAL SCFA, AND OTHER ORGANIC ACIDS IN THE CECAL CONTENTS IN RATS FED THE CONTROL OR DFAIII DIET FOR 14 DAYS

	Control diet	DFAIII diet	<i>P</i> -value
	(μmol in cecal contents)		
Acetic acid	95.1 ± 12.5	243.8 ± 39.6*	0.003
Propionic acid	28.5 ± 3.6	83.8 ± 11.8*	< 0.001
n-Butyric acid	27.2 ± 3.6	29.1 ± 5.0	0.763
iso-Butyric acid	4.7 ± 2.4	6.4 ± 3.4	0.694
n-Valeric acid	8.7 ± 5.2	5.2 ± 1.1	0.391
iso-Valeric acid	8.8 ± 2.9	5.8 ± 1.6	0.581
Total SCFA	179.2 ± 20.4	481.4 ± 83.0*	0.003
Succinic acid	5.6 ± 1.3	102.1 ± 32.1*	0.009
Lactic acid	0.6 ± 0.3	5.2 ± 1.3*	0.004

Values are mean ± SEM (n = 8). The amount of each and total SCFAs (sum of acetic, propionic, n-butyric, iso-butyric n-valeric and iso-varelic acid) and other organic acids (succinic and lactic acid) in the contents of the cecum were analyzed by unpaired *t*-test. Asterisks mean significant difference (*P* < 0.05).

TABLE VII

AMOUNT OF EACH AND TOTAL SCFA AND OTHER ORGANIC ACIDS IN  
THE COLONIC CONTENTS IN RATS FED THE CONTROL OR DFAIII DIET  
FOR 14 DAYS

	Control diet	DFAIII diet	<i>P</i> -value
	(μmol in colonic contents)		
Acetic acid	1.49 ± 0.34	1.01 ± 0.24	0.266
Propionic acid	0.48 ± 0.10	0.56 ± 0.22	0.730
n-Butyric acid	0.45 ± 0.09	0.23 ± 0.07	0.091
iso-Butyric acid	0.17 ± 0.09	0.01 ± 0.01	0.082
n-Valeric acid	0.14 ± 0.07	0.15 ± 0.08	0.866
iso-Valeric acid	0.08 ± 0.03	0.05 ± 0.02	0.364
Total SCFA	2.98 ± 0.68	2.01 ± 0.49	0.266
Succinic acid	0.05 ± 0.04	0.80 ± 0.27*	0.017
Lactic acid	0.02 ± 0.01	0.03 ± 0.01	0.386

Values are mean ± SEM (n = 8). The amount of each and total SCFAs (sum of acetic, propionic, n-butyric, iso-butyric n- valeric and iso- varellic acid) and other organic acids (succinic and lactic acid) in the contents of the cecum were analyzed by unpaired *t*-test. Asterisks mean significant difference (*P* < 0.05).

TABLE VIII

CORRELATION BETWEEN NET CALCIUM ABSORPTION OF CECAL EPITHELIUM IN THE ABSENCE OF LUMINAL DFAIII APPLICATION AND CECAL WEIGHT, CECAL CONTENT WEIGHT, CECAL PH, LEVEL OF CECAL SOLUBLE CALCIUM, VARIOUS MORPHOLOGICAL INDICES IN CECAL TISSUE, CONCENTRATIONS OF VARIOUS SCFAS AND ORGANIC ACIDS IN THE CECAL CONTENTS IN RATS FED THE CONTROL AND DFAIII DIETS.

Correlation to net Ca absorption	<i>R</i> value	<i>P</i> value
Cecal weight	0.531*	0.034
Cecal content weight	0.291	0.273
Cecal content pH	- 0.594*	0.015
Inner circumference	0.482	0.058
Outer circumference	0.621*	0.010
Ratio of inner: outer length	0.544*	0.029
Crypt depth	0.604*	0.013
Number of cells· crypt <sup>-1</sup>	0.485	0.073
Cell width	0.174	0.517
Number of crypt· mm <sup>-1</sup>	0.023	0.931
Acetic acid	0.579*	0.019
Propionic acid	0.530*	0.035
n-Butyric acid	0.151	0.577
iso-Butyric acid	0.519*	0.039
n-Valeric acid	0.090	0.736
iso-Valeric acid	0.320	0.227

1	Total SCFA	0.594*	0.039
2	Succinic acid	0.681*	0.004
3	Lactic acid	0.579*	0.005

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5 Relationship between Ca absorption of the cecal epithelium, each cecal factor and the  
6 cecal contents were evaluated by regression analysis (n =16). Correlation between Ca  
7 absorption (nmol/min· cm) and cecal weight (g), cecal content weight (g), cecal content  
8 pH, morphological indices of cecal tissue cross-section, each and total SCFAs, and  
9 organic acid concentration ( $\mu\text{mol/ g}$  cecal contents) were expressed. Asterisks mean  
10 significant correlation ( $P < 0.05$ ).

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1 Legends

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3 FIG. 1. Structural Formula of Di-D-fructosefuranose 1,2' ; 2,3' Dianhydride  
4 (difructose anhydride III) Produced from Inulin with *Ar-throbacter* sp. H65-7 Inulase II  
5 (EC2.4.1.93).

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7 FIG. 2. Effects of luminal application of DFAIII on net Ca absorption and LY  
8 permeability, and their relationship in the isolated small intestinal epithelium of rats fed  
9 the control or DFAIII diet for 14 days. Values are mean  $\pm$  SEM (n = 8). The  
10 differences in net Ca transport and LY passage induced with (dotted column) or without  
11 (open column) the luminal application of 100mmol/L DFAIII in the small intestine  
12 isolated from the two diet groups was analyzed by two-way ANOVA (A and B). For  
13 net Ca absorption (A), *P* values estimated by two-way ANOVA were = 0.653 for diet  
14 (control and DFAIII diet, D), < 0.001 for DFAIII application (with or without DFAIII in  
15 luminal medium, L) and =0.391 for (D) x (L). For LY permeability (B panel), *P* values  
16 estimated by two-way ANOVA were = 0.889 for (D), < 0.002 for (L) and =0.6175 for  
17 (D) x (L). Values not sharing a superscript letter are significantly different according  
18 to Duncan's test (*P*< 0.05). Regression for net Ca transport and LY passage (n = 32)  
19 after 30-min incubation is shown for the ileum (C panel).

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21 FIG. 3. Effects of luminal application of DFAIII on net Ca absorption and LY  
22 permeability of the isolated cecal epithelium in rats fed the control or DFAIII diet for 14  
23 days. Values are mean  $\pm$  SEM (n = 8). The differences in net Ca transport and LY  
24 passage induced with (dotted column) or without (open column) the luminal application  
25 of 100mmol/L DFAIII in the cecum isolated from the two diet groups was analyzed by  
26 two-way ANOVA. For net Ca absorption (A), *P* values estimated by two-way  
27 ANOVA were < 0.002 for diet (control and DFAIII diet, D), < 0.001 for DFAIII



1 application (with or without DFAIII in luminal medium, L) and =0.894 for (D) x (L).  
2 For LY permeability (B panel), *P* values estimated by two-way ANOVA were = 0.021  
3 for (D), < 0.001 for (L) and =0.307 for (D) x (L). Values not sharing a superscript  
4 letter are significantly different according to Duncan's test ( $P < 0.05$ ). Regression for  
5 net Ca transport and LY passage ( $n = 32$ ) after 30-min incubation is shown for the ileum  
6 (C panel).

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8 FIG. 4. Effects of luminal application of DFAIII on net Ca absorption and LY  
9 permeability of the isolated colonic epithelium in rats fed the control or DFAIII diet for  
10 14 days. Values are mean  $\pm$  SEM ( $n = 8$ ). The differences in net Ca transport and LY  
11 passage induced with (dotted column) or without (open column) the luminal application  
12 of 100mmol/L DFAIII in the colon isolated from two diet groups was analyzed by  
13 two-way ANOVA. For net Ca absorption (A panel), *P* values estimated by two-way  
14 ANOVA were = 0.132 for diet (control and DFAIII diet, D), = 0.024 for DFAIII  
15 application (with or without DFAIII in luminal medium, L) and =0.682 for (D) x (L).  
16 For LY permeability (B panel), *P* values estimated by two-way ANOVA were = 0.239  
17 for (D), < 0.001 for (L) and =0.600 for (D) x (L). Values not sharing a superscript  
18 letter are significantly different according to Duncan's test ( $P < 0.05$ ). Regression for  
19 net Ca transport and LY passage ( $n = 32$ ) after 30-min incubation is shown for the ileum  
20 (C panel).

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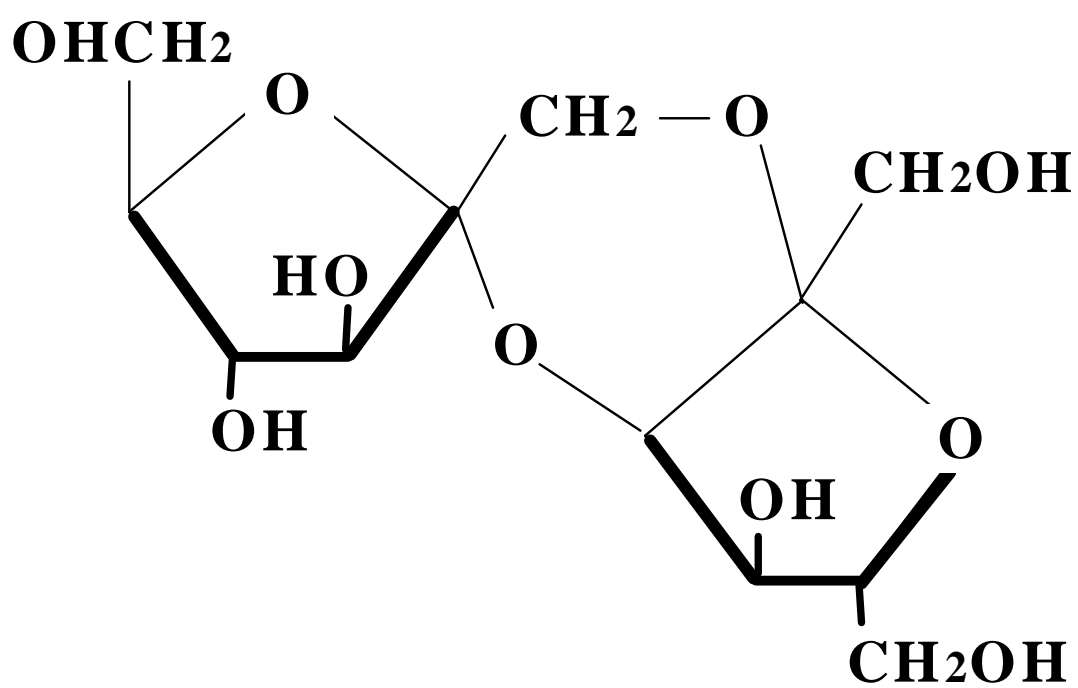
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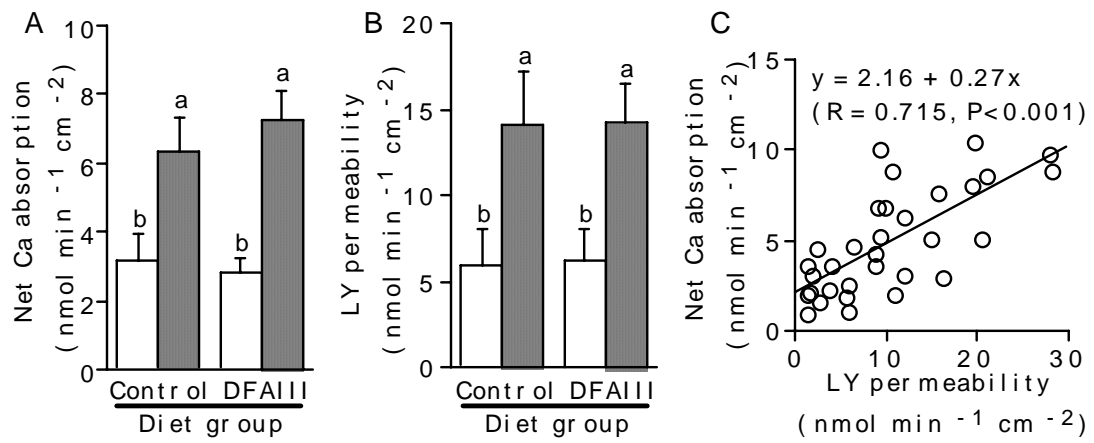
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Fig.1 MINEO H et al



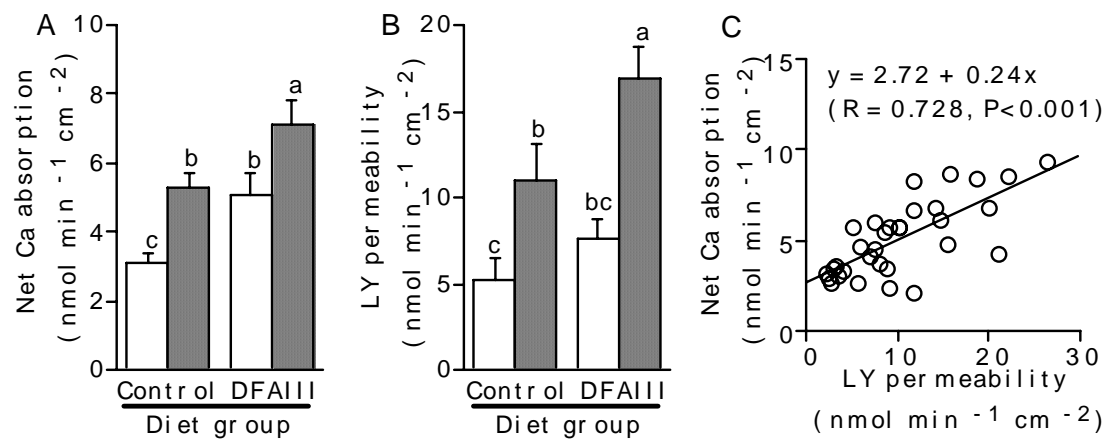
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Fig.2 MINEO H et al



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