



Title	Effect of trehalose supplementation in milk replacer on the incidence of diarrhea and fecal microbiota in preweaned calves
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1 Running title: Fecal microbiota of calves fed trehalose

2

3 **Effect of trehalose supplementation in milk replacer on the incidence of diarrhea**
4 **and fecal microbiota in preweaned calves¹**

5

6 Hiroto Miura*, Kazuhisa Mukai†, Keigo Sudo‡, Satoshi Haga§, Yutaka Suzuki*,

7 Yasuo Kobayashi* and Satoshi Koike*²

8 *Graduate School of Agriculture, Hokkaido University, Hokkaido 060-8589, Japan

9 †Hayashibara Co., Ltd., Okayama 702-8006, Japan

10 ‡Top Farm Group, Hokkaido 093-0506, Japan

11 §Grazing Animal Unit, Division of Grassland Farming, Institute of Livestock and

12 Grassland Science, NARO, Tochigi, 329-2793, Japan

13

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17 ² Corresponding author: skoike7@anim.agr.hokudai.ac.jp

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ABSTRACT

20
21 Trehalose, a nonreducing disaccharide consisting of D-glucose with α,α -1,1
22 linkage, was evaluated as a functional material to improve the gut environment in
23 preweaned calves. In Experiment 1, 173 calves were divided into two groups; the
24 trehalose group was fed trehalose at 30 g/animal/day with milk replacer during the
25 suckling period, and the control group was fed nonsupplemented milk replacer.
26 Medication frequency was lower in the trehalose group ($P < 0.05$). In Experiment 2,
27 calves ($n = 20$) were divided into two groups (control group, $n = 10$, and trehalose group,
28 $n = 10$) based on their body weight and reared under the same feeding regimens as in
29 Experiment 1. Fresh feces were collected from individual animals at the beginning of the
30 trial (average age 11 days), 3 weeks after trehalose feeding (Experimental Day 22), and
31 one day before weaning, and the fecal score was recorded daily. Fecal samples were
32 analyzed for fermentation parameters and microbiota. Fecal score was significantly lower
33 in the trehalose group than in the control group in the early stage (at an age of 14–18 days;
34 $P < 0.05$) of the suckling period. Calves fed trehalose tended to have a higher proportion
35 of fecal butyrate on Day 22 than calves in the control group ($P = 0.08$). Population sizes
36 of *Clostridium* spp. were significantly lower ($P = 0.036$), whereas those of *Dialister* spp.
37 and *Eubacterium* spp. tended to be higher in the feces of calves in the trehalose group on
38 Day 22 ($P = 0.060$ and $P = 0.083$). These observations indicate that trehalose feeding
39 modulated the gut environment and partially contributed to the reduction in medication
40 frequency observed in Experiment 1.

41

42

43 ***Key words:*** calf, fecal microbiota, diarrhea, trehalose

44

45 **List of abbreviations**

46 VFA, volatile fatty acid

47 CP, crude protein

48 PCoA, principal coordinate analysis

49 OTU, operational taxonomic unit

50

INTRODUCTION

51

52 Calf health in the suckling period is an important aspect for future production
53 of cattle. Prevention of infectious diseases, particularly diarrhea (Uetake, 2013), is one of
54 the major concerns in calf management because innate and adaptive immune systems in
55 preweaned calves are not sufficiently established (Weaver et al., 2000) and develop
56 gradually by weaning (Jami et al., 2013; Steele et al., 2016).

57 Gut functions play a vital role not only in nutrient acquisition but also in defense
58 against infectious pathogens (Bischoff, 2011). Gut microbiota is a key factor involved in
59 the maintenance of the gut function (Yeoman and White, 2014). Oikonomou et al. (2013)
60 suggested that the gut microbiota of preweaned calves were related to susceptibility to
61 enteric infections. Therefore, appropriate management focusing on gut microbiota is
62 considered effective for the healthy growth of preweaned calves. In this regard, feeding
63 oligosaccharides is one of the useful approaches to maintain the stability of gut microbiota
64 in calves (Ghosh and Mehla, 2012; Uyeno et al., 2015).

65 In this study, we focused on trehalose, a nonreducing disaccharide consisting
66 of D-glucose with an α,α -1,1 bond. Studies have shown that trehalose feeding exhibited
67 beneficial effects in poultry and dairy cattle, increased the growth of juvenile chicks by
68 improving their intestinal innate immunity (Kikusato et al., 2016) and improved the
69 antioxidative activity in the rumen fluid, blood, and milk of dairy cattle (Aoki et al., 2010;
70 2013) through the alteration of rumen microbiota rather than a direct effect of trehalose
71 as an antioxidant. In addition to these beneficial effects on the poultry and dairy cattle,
72 trehalose can be used as a functional material for preweaned calves through modification

73 of the gut microbiota since trehalose has slower digestion and absorption rate in the
74 human small intestine (Oku and Nakamura, 2000). On the other hand, our preliminary in-
75 vitro experiment suggests that trehalose increases lactate production in calf small intestine
76 (unpublished data). We hypothesized that the excess lactate that flowed into the large
77 intestine is converted into butyrate by lactate-utilizing bacteria. Increased butyrate could
78 exhibit beneficial effects contributing to the reduction of diarrhea incidences, such as the
79 increase of mucin production (Canani et al., 2011) and attenuation of pathogenic bacteria
80 (Namkung et al., 2011) in the calf gut. Therefore, we conducted this study to investigate
81 the effects of trehalose feeding on the incidence of diarrhea in preweaned calves
82 (Experiment 1), followed by an investigation of the fecal microbiota and fermentation
83 parameters of preweaned calves to clarify the mode of action of trehalose as a functional
84 material (Experiment 2).

85

86 **MATERIALS AND METHODS**

87 The calves (crossbreed of Japanese Black and Holstein) used for the two
88 feeding trials in this study were raised in the same farm located in the northern part of
89 Japan (Hokkaido, 44°06 N, 143°79 E). The farm was selected based on two criteria: a
90 sufficient number of calves (c.a. 6,000 calves per year) were available for the experiment,
91 and the experienced professional farmers keep detailed records about feeding
92 management and health condition of individual calves. The animal protocol was
93 conducted in accordance with the Guidelines for Animal Experiments and the Act on
94 Welfare and Management of Animals, Hokkaido University.

95

96 ***Feeding trial 1 (Experiment 1)***

97 A total of 173 crossbred calves with similar age and body weight were chosen
98 and used in Experiment 1. The animals were blocked by birthdate and body weight and
99 randomly assigned to either the control (n = 83) or the trehalose (n = 90) group (Table 1).

100 The number of animals used in the present study was determined following the previous
101 reports in which the effect of feeding oligosaccharides on calf performance was evaluated
102 (Heinrich et al., 2003; Ghosh and Mehla., 2012). To maximize the animal number for the
103 feeding trial, all calves raised in the experimental period (January 2012 to April 2012) at
104 the same feeding area on the farm were subjected to the trial. Because of this technical
105 reason, there was a difference in the number of calves between the control and trehalose
106 groups, but as mentioned above, the animals were randomly allocated to each group.

107 Feeding regimens were similar in both calf groups, with the exception of trehalose feeding.
108 All calves were housed in individual calf hutches and fed 2.5 L of milk replacer (38 to
109 40°C) [27.0% crude protein (CP), 15.5% fat] by feeding bottle once a day at 0800 h.
110 Trehalose was dissolved in a milk replacer, and calves in the trehalose group received 30-
111 g trehalose per day via milk replacer throughout the suckling period. Based on the
112 preliminary results, where feeding 5- or 10-g trehalose per day showed no remarkable
113 beneficial or adverse effects on calf health and growth performance (unpublished data),
114 the feeding level of trehalose (30 g/animal/day) in the present study was determined. Calf
115 starter (20.0% CP, 10.0% crude fiber, 9.0% crude ash, and 2.0% crude fat), timothy hay
116 (10.1% CP, 64.8% neutral detergent fiber, and 39.7% acid detergent fiber), and water

117 were provided on an *ad libitum* basis. Calves were weaned when more than 1.5 kg of calf
118 starter consumption was observed for three consecutive days. The date of weaning was
119 recorded for each calf. As a part of health indices, the incidences of diarrhea and
120 medications were recorded daily for individual animals. Fecal consistency was monitored
121 daily by the experienced professional farmers as a part of routine work on the farm, and
122 the feces showing primarily liquid appearance was recorded as diarrhea. Regarding the
123 medications, when the animals displayed any symptoms of diseases such as diarrhea and
124 respiratory disorder, a combination of antibiotics and antipyretics was administered.
125 Similar to the monitoring of fecal consistency, the necessity of medication was carefully
126 judged by the experienced professional farmers; blinding was not conducted in this
127 experiment, though. The number of calves that experienced diarrhea (%), the average
128 number of days with diarrhea (days/animal), the average frequency of medications
129 (times/animal), and calf mortality (%) were calculated for the control and trehalose
130 groups. In the calculation of medication frequency and mortality, not only diarrhea-
131 associated causes but also other causes, including respiratory diseases, were counted.

132

133 ***Feeding trial 2 for elucidating the mode of action of trehalose (Experiment 2)***

134 ***Experimental design***

135 Male crossbred calves (n = 20) were used in Experiment 2. To perform precise
136 fecal scoring and sampling, Experiment 2 was scaled-down compared to Experiment 1.
137 The number of animals used in Experiment 2 was determined following the previous
138 reports in which the effect of feeding oligosaccharides on the fecal microbiota of calves

139 was evaluated (Uyeno et al., 2013). Calves were divided into two groups (control group,
140 n = 10, and trehalose group, n = 10) on the day before the beginning of the trial (at the
141 age of 10.6 ± 2.0 days) based on their body weight (Table S1). Calves were reared under
142 the same housing condition and same feeding regimens as in Experiment 1. Body weight
143 was measured at the beginning of the trial (Experimental Day 1) and one day before
144 weaning (Experimental Day 55) using a digital body weight scale. Fresh feces were
145 collected from individual animals on Day 1, three weeks after trehalose feeding
146 (Experimental Day 22) and Day 55. Fecal samples were obtained by rectal stimulation of
147 calves to defecate and collected directly into 50 mL tubes. Tubes containing the feces
148 were immediately placed on the ice and then stored at -30°C until use. In Experiment
149 2, the fecal score was recorded daily by the experienced professional farmers according
150 to the following criteria, as reported in the previous studies (Heinrichs et al., 2003; Ghosh
151 and Mehla, 2012): 1 = normal feces; 2 = slightly liquid feces; 3 = moderately liquid feces;
152 and 4 = primarily liquid feces. Blinding was not conducted in this experiment.

153

154 *Measurement of fermentation parameters*

155 For measuring pH and volatile fatty acid (VFA), 0.3-g feces was mixed with
156 800 μL of saline and centrifuged at $16,000 \times g$ for 5 min at 4°C . The supernatant was
157 subjected to pH measurement using a pH electrode (LAQUAtwin B-712; HORIBA,
158 Kyoto, Japan) and VFA measurement using a gas chromatograph (GC-14B; Shimadzu,
159 Kyoto, Japan), as described previously (Oh et al., 2017).

160 For measuring ammonia nitrogen ($\text{NH}_3\text{-N}$) and D-/L-lactate, 0.1-g feces was
161 mixed with 500 μL of saline and centrifuged at $16,000 \times g$ for 5 min at 4°C . $\text{NH}_3\text{-N}$ levels
162 were measured using the phenol–hypochlorite reaction method (Weatherburn, 1967). The
163 levels of D-/L-lactate were measured using a commercial assay kit (Megazyme, Wicklow,
164 Ireland). For measuring indole and skatole, 0.1-g feces was mixed with 1-mL 0.1 M
165 phosphate-buffered saline and centrifuged at $16,000 \times g$ for 5 min at 4°C . The assays for
166 indole and skatole were conducted using colorimetric methods (Walstra et al., 1999).

167

168 *Analysis of microbiota*

169 Total DNA was extracted and purified from 0.3-g fecal sample using the
170 repeated bead-beating plus column method (Yu and Morrison, 2004) using a commercial
171 kit (QIAmp DNA Stool mini kit; Qiagen, Hilden, Germany). The DNA concentration was
172 quantified by NanoDrop 2000 (Thermo Fisher Scientific, Waltham, USA).

173 Total DNA was diluted to a final concentration of 5 ng/ μL for all samples and
174 subjected to the amplification of the V3–V4 region of 16S rRNA gene using the primer
175 set of S-D-Bact-0341-b-S-17 (5'-CCTACGGGNGGCWGCAG-3') and S-D-Bact-0785-
176 a-A-21 (5'-GACTACHVGGGTATCTAATCC-3') (Herlemann et al., 2011). The PCR
177 mixture consisted of 12.5 μL of $2 \times$ KAPA HiFi HotStart Ready Mix (KAPA Biosystems,
178 Wilmington, USA), 0.1 μM of each primer, and 2.5 μL of DNA (5 ng/ μL). The PCR steps
179 were performed according to the following program: initial denaturation at 95°C for 3
180 min, 25 cycles at 95°C for 30 s, 55°C for 30 s, and 72°C for 30 s, and a final extension
181 step at 72°C for 5 min. Amplicons were purified using AMPure XP beads (Beckman-

182 Coulter, Brea, USA) and subjected to sequencing on the Illumina MiSeq platform
183 (Illumina, San Diego, USA). Raw sequences have been deposited in the NCBI Sequence
184 Read Archive under the accession no. DRA010638.

185 Data obtained from MiSeq sequencing were analyzed using the QIIME tool kit
186 version 1.9.1 (Caporaso et al., 2010). Paired-end reads were merged and filtered to
187 remove low-quality reads based on the quality scores (sequences with a quality score of
188 >20 were retained for further analysis). The high-quality reads were assigned to
189 operational taxonomic units (OTUs) at a 97% identity threshold using the UCLUST
190 algorithm (Edgar, 2010), and taxonomy was assigned using the latest Greengenes
191 database (v.13_8). The taxonomic identification was performed at the genus level. Alpha
192 diversity indices, including Chao1, Shannon, ACE, and Simpson, were calculated using
193 the QIIME. The relative abundance of each bacterial taxon was calculated by dividing
194 the number of reads assigned to each bacterial taxon by the total number of reads. Only
195 taxa with a relative abundance of >0.1% in at least 5 individuals in either control and
196 trehalose groups on Day 22 or Day 55 were considered as being observed and used for
197 the analysis. Principal coordinate analysis (PCoA) was used to determine the differences
198 in the microbial community structure based on the Bray–Curtis dissimilarity matrices at
199 the genus level using the vegan package version 2.5.6 (Oksanen et al., 2019) in R.

200 Quantitative real-time PCR was performed to determine the population sizes of
201 major gut bacterial groups, including Ruminococcaceae, *Bacteroides–Prevotella–*
202 *Porphyromonas* group, *Prevotella* spp., *Faecalibacterium prausnitzii*, *Lactobacillus* spp.,
203 *Clostridium coccoides–Eubacterium rectale* group, *Bifidobacterium* spp., *Escherichia*

204 *coli*, *Akkermansia muciniphila*, *Clostridium perfringens*, *Eubacterium hallii*,
205 *Streptococcus bovis*, and *Megasphaera elsdenii*, using a LightCycler system with a
206 KAPA SYBR Fast qPCR Kit (Kapa Biosystems, Charlestown, USA) with the respective
207 primer sets (Table S2). The PCR thermal conditions and the PCR mixture were the same
208 as those reported earlier (Myint et al., 2017). The population size of each bacterial target
209 was expressed as the proportion (%) of the abundance of 16S rRNA genes of each
210 bacterial target against that of the total bacteria.

211

212 ***Statistical analysis***

213 The ages of calves, body weight, number of days with diarrhea, frequency of
214 medications, fecal score, fecal fermentation parameters, alpha diversity indices in fecal
215 microbiota, and the bacterial abundance quantified by MiSeq and real-time PCR were
216 compared between the control and trehalose groups by Welch's t-test using R version
217 3.6.2 (R Core Team, 2019). Prior to performing Welch's t-test, the same data set was
218 analyzed with a repeated measures model with the fixed effects of treatment, day, and
219 treatment \times day interaction and the random effect of calf within the groups. Results
220 showed the greater effect of day in all data, while the effects of treatment and interaction
221 between treatment \times day were less apparent (Table S3). To clarify the effect of feeding
222 trehalose at respective sampling points (i.e., Day 22 and Day 55), the values were
223 compared between the control and trehalose groups using Welch's t-test. The number of
224 calves that experienced diarrhea and calf mortality were compared (control vs. trehalose
225 groups) by the Chi-squared test using R. *P*-values of <0.05 and <0.10 were considered to

226 be statistically significant and trend, respectively. The Pearson correlation between the
227 molar ratio of each VFA and the bacterial abundance quantified by MiSeq was analyzed
228 using the corrplot package version 0.84 (Wei and Viliam, 2017) in R.

229

230

RESULTS

231 *Analysis of the effects of trehalose feeding on the incidence of diarrhea in preweaned*
232 *calves (Experiment 1)*

233 Table 1 shows the results obtained from Experiment 1. There were no
234 differences in the weaning age, number of calves that experienced diarrhea, and mortality
235 between the feeding groups. However, medication frequency was significantly lower in
236 the trehalose group than in the control group ($P < 0.05$).

237

238 *Feeding trial 2 for elucidating the mode of action of trehalose (Experiment 2)*

239 *Growth and fecal fermentation parameters*

240 The growth of calves and the medication frequency observed in Experiment 2
241 were not different between the feeding groups (Table S1). The average fecal score during
242 the suckling period is depicted in Figure 1. In the trehalose group, the average fecal score
243 was significantly lower at Experimental Days 4 and 8 ($P < 0.05$) and tended to be lower
244 at Experimental Days 5 and 6 than those in the control group ($P < 0.10$). The repeated-
245 measures model analysis also supported this time-dependent effect; there was a
246 significant interaction between treatment and day ($P = 0.001$).

247 The fecal fermentation parameters are presented in Table 2. There were no
248 significant differences in fecal pH, total VFA, ammonia, total lactate, indole, and skatole
249 concentrations between the control and trehalose groups. The proportion of butyrate in
250 the trehalose group tended to be higher on Day 22 ($P = 0.080$), whereas that of valerate
251 was significantly lower on Day 55 ($P < 0.05$).

252

253 ***Community structure and composition of fecal microbiota***

254 Amplicon sequencing of the fecal samples yielded a total of 878,751 high-
255 quality reads, with an average of $21,968 \pm 5,532$ reads per sample. No differences were
256 observed in the alpha diversity indices (Chao1, ACE, Shannon, and Simpson) between
257 the two groups on Day 22 and Day 55 (Table S4). Although the PCoA revealed a
258 distinction between Day 22 and Day 55, there was no clear difference in the microbial
259 composition between the control and trehalose groups (Figure S1).

260 A total of 43 OTUs at the genus level were detected from the fecal samples
261 (Table S5). The bacterial taxa exhibiting statistical significance or trend in the relative
262 abundance are listed in Table 3. On Day 22, the relative abundance of *Clostridium* spp.
263 was lower, whereas those of *Dialister* spp. and putative [Eubacterium] were higher in the
264 trehalose group than in the control group. On Day 55, the proportions of unclassified S24-
265 7, putative [Prevotella], *Coprococcus* spp., *Odoribacter* spp., *Dialister* spp., and
266 *Lachnospira* spp. were higher in the trehalose group, whereas the proportion of
267 unclassified Coriobacteriaceae was lower in the trehalose group. A correlation analysis
268 was conducted between the molar ratio of each VFA and bacterial abundance quantified

269 by MiSeq to explore the bacterial taxa related to butyrate proportion (Figure 2). The
270 results showed that the butyrate proportion correlated negatively with the relative
271 abundance of *Clostridium* spp. but positively with *Bifidobacterium* spp..

272 The population sizes of major gut bacterial groups quantified by real-time PCR
273 are depicted in Figure 3. On Day 22, the population sizes of targeted bacterial groups
274 were not significantly different, whereas the prevalence of *Clostridium perfringens* was
275 lower in the trehalose group; two calves in the control group harbored this species,
276 whereas it was not detected in calves in the trehalose group (Figure 3). On Day 55, the
277 population size of Ruminococcaceae tended to be lower in the trehalose group than in the
278 control group ($P = 0.09$).

279

280

DISCUSSION

281 Existing literature reports the beneficial effects of trehalose feeding in poultry
282 (Kikusato et al., 2016) and dairy cattle (Aoki et al., 2010; 2013). In the present study, we
283 evaluated trehalose as a functional material for preweaned calves. Lower medication
284 frequency in the trehalose group was confirmed in Experiment 1. Although this result
285 suggests the improvement of calf health by trehalose feeding, the mode of action needs
286 to be further investigated. It has been suggested that the severe disease which requires
287 antibiotic treatment during the preweaning period of calves leads to less milk production
288 in their lifetime (Soberon et al., 2010). Therefore, reducing the medication frequency (i.e.,
289 less antibiotic administration) via trehalose feeding could be beneficial in terms of the
290 lifetime productivity of cattle. The calves in the present study were fed milk replacer once

291 a day, and this feeding management is not common. Although the farm used in the present
292 study has been adopting this feeding management without adverse effects (i.e.,
293 comparable growth and health conditions with multiple feedings), the optimum amount
294 of trehalose under different feeding management needs to be considered.

295 Although the average number of days with diarrhea was numerically higher in
296 the trehalose group in Experiment 1, we speculated that diarrhea in the trehalose group
297 was a mild symptom that did not require medications. In fact, improved fecal score was
298 confirmed in Experiment 2 of this study. Earlier studies have reported that feeding with
299 mannan oligosaccharides (MOS) improves the fecal score of calves (Heinrichs et al.,
300 2003; Ghosh and Mehla, 2012). MOS is known to inhibit the attachment of infectious
301 pathogens to the gut epithelium, thus leading to the improvement of animal health (Spring
302 et al., 2000). In contrast, Uyeno et al. (2013) suggested that cello-oligosaccharides
303 primarily composed of cellobiose modulate the gut environment with higher population
304 size of *Clostridium coccooides*–*Eubacterium rectale* group and butyrate concentration.
305 Similar to cellobiose, trehalose is a dimer of D-glucose, and the improvement in fecal
306 score by trehalose is probably due to the modulation of gut microbiota rather than due to
307 the direct inhibition of infectious pathogens.

308 To elucidate the mechanism of modulation of the gut environment by trehalose
309 feeding, a further feeding trial was conducted as Experiment 2 for fecal analysis. The
310 trehalose group showed an increase in butyrate levels at 3 weeks after trehalose feeding
311 (Day 22). Butyrate is the primary energy source of gut epithelial cells, which mediate the
312 absorption of water, mineral, and nutrients (Bedford and Gong, 2018). Moreover,

313 butyrate increases the expression of mucin protein in the gut epithelia (Canani et al., 2011).
314 The mucosal layer in the gut plays a key role in preventing the invasion of infectious
315 pathogens (Johansson et al., 2011). Therefore, the increase in butyrate levels by trehalose
316 feeding may contribute to the maintenance of the gut function, which reduces the severity
317 of diarrhea.

318 The alpha diversity indices and the PCoA plot of fecal microbiota indicated that
319 there were no remarkable alterations in the microbial community structures by trehalose
320 feeding. However, some bacterial groups changed their abundances, and a decrease in
321 *Clostridium* spp. abundance and an increase in *Dialister* spp. and putative [Eubacterium]
322 abundances were observed in the trehalose group on Day 22. It has been reported that
323 *Clostridium* spp. was detected at a higher abundance in the feces of diarrheic cattle
324 (Zeineldin et al., 2018). Some *Clostridium* species are considered as pathogenic (e.g., *C.*
325 *perfringens* and *C. difficile*) (Blanchard, 2012; Cho and Yoon, 2014). Therefore, the
326 decrease in *Clostridium* spp. abundance by trehalose feeding can contribute to the
327 prevention of infectious diseases and may partially explain the lowered medication
328 frequency in Experiment 1. A negative correlation between the abundance of *Clostridium*
329 spp. and the butyrate molar ratio was observed in Experiment 2, and the reduction in
330 *Clostridium* spp. abundance by trehalose feeding might be attributed to butyrate
331 enhancement. Namkung et al. (2011) reported that butyrate attenuated *C. perfringens* *in*
332 *vitro*. In Experiment 2 of the present study, none of the calves fed trehalose harbored *C.*
333 *perfringens*, whereas this bacterium was detected from two calves in the control group.

334 Altogether, an increase in butyrate level by trehalose feeding can be one of the factors
335 contributing to the beneficial effect of trehalose.

336 *Eubacterium* spp. are considered as beneficial bacteria producing butyrate from
337 lactate in the animal gut (Rivière et al., 2016). Therefore, an increase in *Eubacterium* spp.
338 abundance by trehalose feeding probably contributed to butyrate enhancement on Day 22.
339 Judging from the positive correlation between the relative abundance of *Bifidobacterium*
340 spp. and the butyrate proportion observed in Experiment 2, lactate produced by
341 *Bifidobacterium* spp. could act as a substrate for butyrate production by *Eubacterium* spp.
342 However, the elevation of *Bifidobacterium* spp. abundance in the trehalose group was not
343 observed in both the MiSeq sequencing and real-time PCR quantification results.
344 Therefore, we hypothesized that lactate was produced in the small intestine where lactic
345 acid bacteria are abundantly colonized (Malmuthuge et al., 2014). Recently, it is reported
346 that trehalose supplementation increased the abundance of *Lactobacillus* spp. in the
347 duodenum and jejunum of broilers (Wu et al., 2020). We conducted batch culture tests
348 using the jejunum content of Holstein calves and confirmed a remarkable increase in
349 lactate levels (unpublished data). This increase in lactate levels indicates that trehalose
350 was used by lactic acid bacteria in the small intestine of calves. These findings suggest
351 that lactate production in the small intestine was stimulated by trehalose, and the excess
352 lactate that flowed into the large intestine was metabolized by lactate-utilizing bacteria
353 such as *Eubacterium* spp.

354 Previous studies indicate the positive correlation between ruminal or fecal
355 abundance of *Dialister* spp. and the calf performance such as body weight, feed intake

356 and feed efficiency (Myer et al., 2015; Meale et al., 2017). Wang et al. (2016) reported
357 that *Dialister* spp. in the rumen might be involved in starch and fiber digestion and
358 contribute to VFA production. Although the role of *Dialister* spp. in the calf gut is not
359 completely understood, previous studies have reported a higher abundance of *Dialister*
360 spp. in healthy individuals than in patients with colorectal cancer or Crohn's disease
361 (Joossens et al., 2011; Weir et al., 2013). Therefore, *Dialister* spp. is considered to
362 contribute to host health, and an increased abundance of this bacterial genus by trehalose
363 feeding might be beneficial for the gut environment of calves.

364 In this study, on Day 55 (one day before weaning), there were increases in the
365 abundance of unclassified S24-7, putative [Prevotella], *Coprococcus* spp., *Odoribacter*
366 spp., *Dialister* spp., and *Lachnospira* spp. in the trehalose group. Studies have shown the
367 involvement of unclassified S24-7 in cellulose digestion in the gastrointestinal tract (Naas
368 et al., 2014; de Mulder et al., 2017). *Prevotella* and *Lachnospira* genera are also
369 considered to be involved in fiber digestion (Bryant and Small, 1956; Matsui et al., 2000).
370 The increase in the abundance of fiber-degrading bacteria by trehalose feeding on Day 55
371 potentially stimulates the utilization of dietary fiber in the large intestine after weaning.

372 There are some limitations in the present study; the feeding trials were
373 conducted in the commercial farm, and experienced professional farmers monitored fecal
374 consistency as a part of routine work on the farm without blinding. Since early detection
375 of diarrhea in the calves is one of the most important tasks of the farm manager to prevent
376 severe illness that directly impacts animal productivity, fecal scoring was performed with
377 stringent criteria. However, the beneficial effect of feeding trehalose to the calves

378 (reduction of medication frequency and fecal score improvement) needs to be further
379 evaluated by a blinding test with a trained person for the research purpose. Although the
380 effect of feeding trehalose was detected at respective sampling points, the treatment effect
381 was not clearly observed in the statistical analysis with the repeated measure model. This
382 was probably due to the individual differences in fermentation parameters and microbiota
383 of the calves. Improvement of the gut environment by feeding trehalose needs to be
384 further clarified by the studies with larger sample size.

385 In conclusion, trehalose feeding reduced the medication frequency of
386 preweaned calves, and the growth inhibition of pathogenic bacteria through butyrate
387 enhancement might be a possible mode of action causing these beneficial effects.
388 Although further investigation using pure culture of gut bacteria is necessary to elucidate
389 the details, trehalose can be used as a functional material for calf production.

390

391

CONFLICT OF INTEREST

392 The authors declare no conflicts of interest.

393

394

LITERATURE CITED

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591 **TABLES AND FIGURES**

592

593 **Table 1.** Effect of trehalose feeding on the incidence of diarrhea, medication frequency, and mortality in the calves (Experiment 1)

Item	Control	Trehalose	<i>P</i> -value ¹
Number of calves	83	90	NA
Age at the beginning of trial, days	19.19 ± 7.68	19.80 ± 8.69	0.628
Body weight at the beginning of trial, kg	47.51 ± 5.54	48.20 ± 5.88	0.427
Age at weaning, days ²	72.96 ± 14.30	71.51 ± 14.08	0.511
Number of calves that experienced diarrhea (%) ³	16/83 (19.2)	22/90 (24.4)	0.524
Average number of days with diarrhea, days/animal ^{3,4}	2.81 ± 1.38	4.59 ± 4.62	0.108
Average frequency of medications, times/animal ^{5,6}	3.68 ± 4.70	2.38 ± 3.54	0.049
Mortality (%) ⁶	3/83 (3.6)	2/90 (2.2)	0.926

594

595 Values are shown as mean ± SD.

596 NA: not applicable

597 ¹ *P*-values were calculated based on Welch's t-test.

598 ² Calves were weaned when more than 1.5 kg of calf starter consumption was observed for 3 consecutive days.

599 ³ Fecal consistency was monitored daily by experienced professional farmers as a part of routine work on the farm, and the feces
600 showing primarily liquid appearance was recorded as diarrhea.

601 ⁴ The sum of the number of days with diarrhea in the calves that experienced diarrhea was normalized by the number of calves that
602 experienced diarrhea.

603 ⁵ Total frequency of medications in each herd was normalized by total animals in each herd. The necessity of medication was carefully
604 judged by the experienced professional farmers.

605 ⁶Not only diarrhea-related causes but also other causes, including respiratory diseases, were counted.

606

607 **Table 2.** Effect of trehalose feeding on fecal fermentation parameters in the calves at Day 22 and Day 55 (Experiment 2)

Item	Day 22			Day 55		
	Control	Trehalose	<i>P</i> -value ¹	Control	Trehalose	<i>P</i> -value ¹
pH	7.15 ± 0.49	7.23 ± 0.48	0.734	7.66 ± 0.30	7.55 ± 0.43	0.553
Total VFA, µmol/g feces	48.9 ± 11.9	56.0 ± 17.9	0.334	76.3 ± 15.9	84.5 ± 34.2	0.525
Acetate, mol/100 mol	55.5 ± 5.0	52.9 ± 5.3	0.305	60.0 ± 5.4	58.4 ± 2.7	0.431
Propionate, mol/100 mol	27.6 ± 3.0	28.0 ± 3.5	0.814	25.9 ± 2.4	27.8 ± 3.5	0.207
Butyrate, mol/100 mol	11.9 ± 2.7	14.5 ± 3.3	0.080	8.8 ± 1.6	9.4 ± 1.8	0.516
Isobutyrate, mol/100 mol	1.2 ± 0.7	1.6 ± 1.2	0.425	1.4 ± 0.8	1.9 ± 1.0	0.219
Isovalerate, mol/100 mol	1.5 ± 0.9	1.9 ± 1.7	0.640	1.5 ± 1.0	1.4 ± 1.1	0.762
Valerate, mol/100 mol	2.3 ± 1.8	1.2 ± 1.2	0.131	2.3 ± 1.5	1.2 ± 0.6	0.047
Ammonia, µgN/g feces	0.27 ± 0.12	0.30 ± 0.34	0.831	0.21 ± 0.12	0.23 ± 0.14	0.782
Total lactate, µmol/g feces	5.51 ± 3.14	6.31 ± 7.71	0.779	14.59 ± 5.67	13.74 ± 8.94	0.815
Indole, µg/g feces	12.67 ± 8.52	14.02 ± 18.54	0.846	6.62 ± 3.33	7.77 ± 3.32	0.474
Skatole, µg/g feces	104.8 ± 46.4	119.5 ± 101.3	0.700	77.4 ± 30.5	87.9 ± 30.4	0.474

608

609 Values are wet weight basis and shown as mean ± SD.

610 ¹ Each parameter was compared between control and trehalose groups at respective sampling points. *P*-values were calculated by

611 Welch's t-test.

612

613 **Table 3.** Effect of trehalose feeding on fecal microbial abundances in the calves at Day 22 and Day 55 (Experiment 2)

Taxa ¹	Control	Trehalose	P-value ²
Day 22			
<i>Clostridium</i> spp.	1.575 ± 1.347	0.448 ± 0.390	0.036
<i>Dialister</i> spp.	0.379 ± 0.693	3.680 ± 4.577	0.060
[Eubacterium]	0.128 ± 0.119	0.352 ± 0.332	0.083
Day 55			
Unclassified S24-7	10.268 ± 3.309	15.511 ± 5.709	0.031
[Prevotella]	2.894 ± 2.758	5.930 ± 2.765	0.031
Unclassified Coriobacteriaceae	0.773 ± 0.349	0.435 ± 0.209	0.025
<i>Coprococcus</i> spp.	0.506 ± 0.305	0.850 ± 0.493	0.095
<i>Odoribacter</i> spp.	0.151 ± 0.072	0.314 ± 0.229	0.068
<i>Dialister</i> spp.	0.036 ± 0.038	0.182 ± 0.214	0.073
<i>Lachnospira</i> spp.	0.025 ± 0.027	0.110 ± 0.095	0.027

614

615 Abundance of bacterial taxa with a relative abundance of >0.1% in at least 5 individuals in either control and trehalose groups on Day
616 22 or Day 55 were compared between groups at respective sampling points.

617 Name of taxa with square brackets indicates putative assignment.

618 Values mean relative abundance (% of total reads) and are shown as mean ± SD.

619 ¹ Bacterial taxa showing statistical significance or trend in the relative abundance are listed.

620 ² P-values were calculated by Welch's t-test.

622 **Figure legends**

623

624 **Figure 1.** Effect of trehalose feeding on the average fecal score of calves during the suckling period in Experiment 2. Fecal score was
625 recorded daily by the experienced professional farmers as a part of routine work on the farm according to the following criteria: 1 =
626 normal feces, 2 = slightly liquid feces, 3 = moderately liquid feces, and 4 = primarily liquid feces. Average ages at Experimental Day 1
627 were 11.9 ± 2.0 days for the control group and 11.4 ± 2.0 days for the trehalose group (Table S1).

628 * $P < 0.05$ and † $P < 0.10$ (compared between control and trehalose group at respective days by Welch's t-test).

629

630 **Figure 2.** Pearson correlation between proportion of VFA and bacterial abundance in the feces of calves at Day 22 (Experiment 2). The
631 values for respective VFA represents proportion against total VFA (mol/100 mol). Bacterial abundance represents relative abundance
632 (% of total reads). Taxa with a relative abundance of $>0.1\%$ in at least 5 individuals in either control and trehalose groups on Day 22 or
633 Day 55 were included in the matrix. Name of taxa with square brackets indicates putative assignment. Only the plots showing
634 significance or trend are indicated in the panel. The scale colors indicate the correlation; positive (closer to 1, red circle) or negative
635 (closer to -1, blue circle).

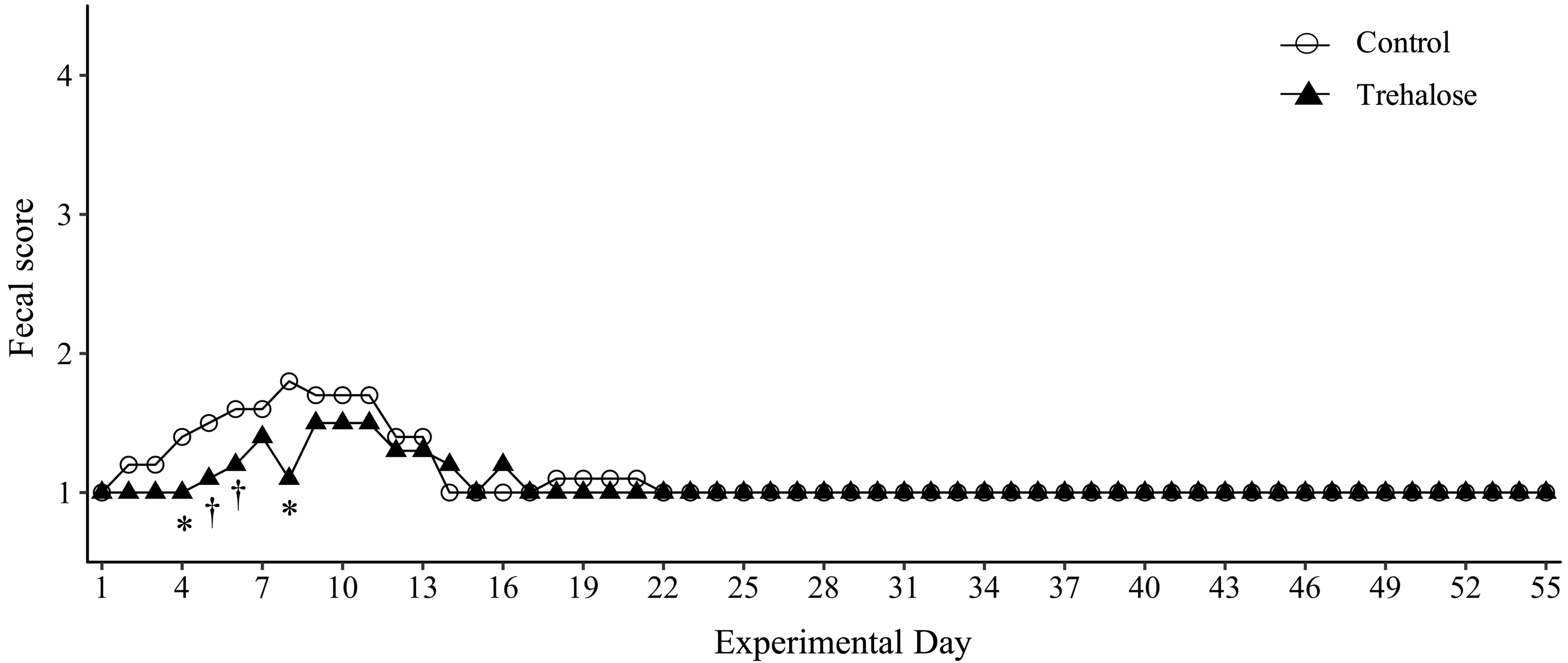
636

637 **Figure 3.** Effect of trehalose feeding on population sizes of major gut bacterial groups in the feces of calves at Day 22 and Day 55
638 determined by real-time PCR (Experiment 2). The population size of each bacterial target was expressed as the proportion (%) of the
639 abundance of 16S rRNA genes of each bacterial target against that of the total bacteria. Error bars indicate SD.

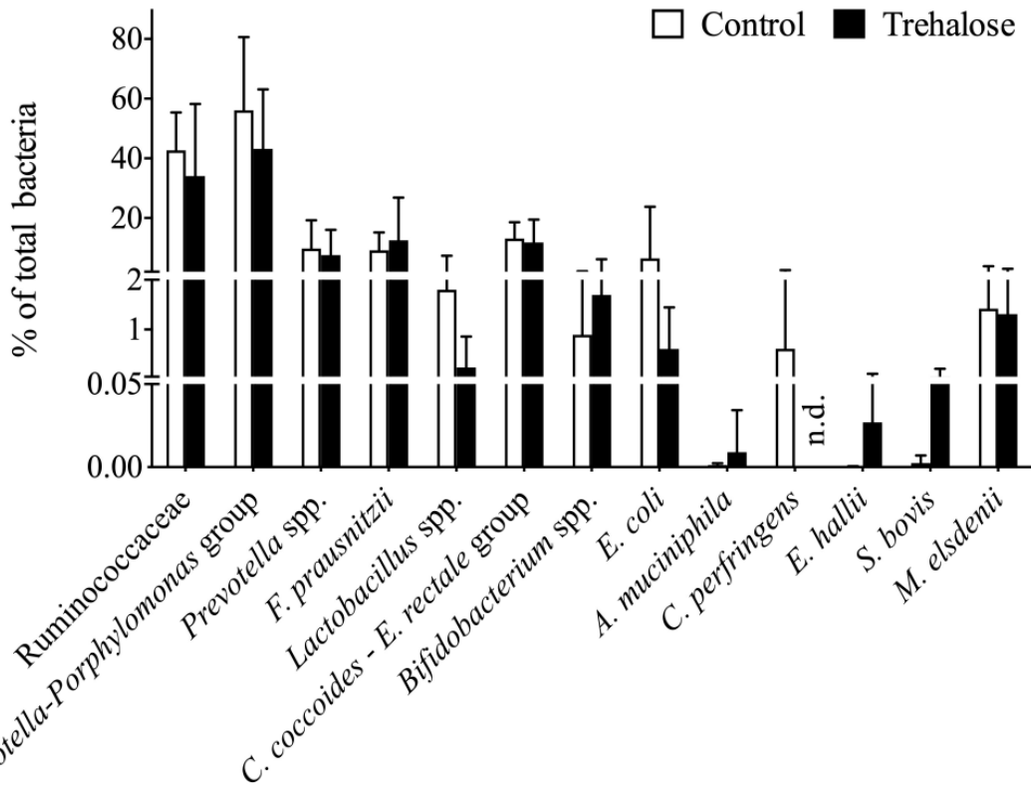
640 * $P < 0.05$ and † $P < 0.10$ (compared between control and trehalose group at respective days by Welch's t-test).

641 n.d.: not detected

642



Day 22



Day 55

