Molecular Ecological Analysis of Fecal Bacterial Populations from Term Infants Fed Formula Supplemented with Selected Blends of Prebiotics

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Supplementation of infant formulas with prebiotic ingredients continues the effort to mimic functional properties of human milk. In this double-blind, controlled, 28-day study, healthy term infants received control formula (control group; n = 25) or control formula supplemented with polydextrose (PDX) and galactooligosaccharide (GOS) (4 g/liter) (PG4 group; n = 27) or with PDX, GOS, and lactulose (LOS) (either 4 g/liter [PGL4 group; n = 27] or 8 g/liter [PGL8 group; n = 25]). A parallel breast-fed group (BF group) (n = 30) was included. Stool characteristics, formula tolerance, and adverse events were monitored. Fecal bacterial subpopulations were evaluated by culture-based selective enumeration (Enterobacteriaceae), quantitative real-time PCR (Clostridium clusters I, XI, and XIV, Lactobacillus, and Bifidobacterium), and fluorescence in situ hybridization (FISH). Fecal bacterial community profiles were examined by using 16S rRNA gene PCR-denaturing gradient gel electrophoresis. The daily stool consistency was significantly softer or looser in the BF group than in all of the groups that received formula. The formulas were well tolerated, and the incidences of adverse events did not differ among feeding groups. Few significant changes in bacterial subpopulations were observed at any time point. The bacterial communities were stable; individual profiles tended to cluster by subject rather than by group. Post hoc analysis, however, demonstrated that the bacterial community profiles for subjects in the BF, PG4, PGL4, and PGL8 groups that first received formula at a younger age were less stable than the profiles for subjects in the same groups that received formula at an older age, but there was no difference for the control group. These data indicate that formulas containing PDX, GOS, and LOS blends are more likely to influence gut microbes when administration is begun in early infancy and justify further investigation of the age-related effects of these blends on fecal microbiota.

Nondigestible food ingredients called prebiotics pass into the lower gastrointestinal tract and, by definition, may be selectively metabolized by mutualistic microorganisms, such as Lactobacillus spp. and Bifidobacterium spp., which in turn contribute to improved host health (12, 34). After lactose and lipids, oligosaccharides, which have prebiotic activity, are the third largest component of human breast milk (5 to 10 g/liter), and there are as many as 200 distinct molecular structures (5, 26). Lactobacilli and bifidobacteria are the predominant bacteria in the intestinal microbiota of breast-fed infants, whereas infants who receive cow’s milk-based infant formulas, which naturally contain low levels of oligosaccharides, often have higher concentrations of potentially pathogenic bacteria, such as Enterobacteriaceae and clostridia, in their intestinal microbiota (4, 15, 17).

Clinical investigations of infant formulas supplemented with galactooligosaccharide (GOS) and fructooligosaccharide (FOS) at a range of concentrations and a range of blends have revealed increases in the numbers of fecal bifidobacteria in preterm infants (7, 18) and in the numbers of both bifidobacteria and lactobacilli in term infants (2, 3, 6, 7, 13, 14, 18, 19, 23, 24, 31) compared with infants receiving unsupplemented formulas. The Bifidobacterium levels (3, 6, 7, 14) and stool consistencies (7, 23, 24) have often been comparable for infants in both prebiotic-supplemented formula groups and human breast milk-fed reference groups. Nondigestible polydextrose (PDX) and lactulose (LOS) also stimulate bifidobacterial growth (8, 16, 29). In a recent safety and tolerance study, healthy term infants who received a formula supplemented with prebiotic blends of PDX, GOS, and LOS when they were 14 to 120 days old exhibited normal growth and stool characteristics more similar to those of breast-fed infants than to those of infants fed an unsupplemented control formula (35).

In the present study, the effects of the same prebiotic blends on the fecal bacterial populations of formula-fed infants were investigated during a 28-day feeding period using 16S rRNA gene-based real-time quantitative PCR (qPCR), 16S rRNA-based fluorescence in situ hybridization (FISH), and PCR-denaturing gradient gel electrophoresis (DGGE). These culture-independent molecular techniques are now commonly used to identify, quantify, and determine the profiles of bacteria in complex microbial communities that are difficult to assess using standard culture methods, such as the communities in the gastrointestinal tract (2, 6, 9, 13–15, 19, 22, 28, 31, 36).

In this study, the Clostridium botulinum group (cluster I), the Clostridium butyricum group (cluster IV), the Clostridium coccoides-Eubacterium rectale group (cluster XIV), and...
the genera *Lactobacillus* and *Bifidobacterium* in infant fecal samples were targeted with 16S rRNA gene group-specific primers, and changes in the subpopulations were evaluated by qPCR. Potential changes in the bifidobacterial subpopulation and total bacteria were also evaluated by using FISH, and *Enterobacteriaceae* were enumerated on selective media. Specific changes in the sum of the levels of *Clostridium* clusters I and XI, which include the pathogenic organisms *Clostridium perfringens* and *Clostridium difficile*, respectively, were chosen for the primary comparison in this study since consistently lower levels of *Clostridium* spp. have been found in breast-fed infants than in formula-fed infants (10). Differences in bacterial community composition between feeding groups and between individual participants were analyzed by using DGGE.

Infant growth, stool characteristics, and tolerance of formulas were also monitored. The results were compared with the results for a parallel breast-fed reference group.

**MATERIALS AND METHODS**

**Subjects and experimental design.** Healthy term infants who were 13 to 92 days old were recruited for a multicenter, double-blind, controlled, parallel-group feeding trial in the United States. Infants who received antibiotics within 14 days of the beginning of the study were excluded. At the time of enrollment, formula-fed infants (n = 105) started receiving a commercially marketed formula, Enfamil LIPIl with iron (Mead Johnson & Company, Evansville, IN) (control group). Following a 7-day formula run-in phase, all infants participated in a feeding study over a 28-day period. At baseline (on day 1 of the feeding study), formula-fed infants were randomized and placed in one of four groups: (i) the control group (n = 25); (ii) a group that received the control formula supplemented with 4 g/liter of a probiotic blend containing PDX (Litesse Two PDX; Danisco, Copenhagen, Denmark) and GOS (Vivinal GOS; Friesland Foods Domo, Zwolle, The Netherlands) at a ratio of 1:1 (PG4 group; n = 27); (iii) a group that received the control formula supplemented with 4 g/liter of a probiotic blend containing PDX, GOS, and LPS (lactulose anhydride; Morinaga Milk Industry Co., Ltd., Tokyo, Japan) at a ratio of 3:2:1 (PGLA group; n = 27); or (iv) a group that received the control formula supplemented with 8 g/liter of a probiotic blend containing PDX, GOS, and LPS at a ratio of 3:2:1 (PGLA group; n = 26). A parallel breed-fed group of infants (BF group) was included as a reference group (n = 30). All of the infants in the reference group were exclusively or predominantly (no more than 120 ml water or juice per day) breast fed prior to the trial, were exclusively breast fed for 7 to 8 days before collection of the baseline stool samples, and then were exclusively breast fed for the remainder of the study. Body weight (in grams), length (in centimeters), and head circumference (in centimeters) were recorded at enrollment and day 28. A parent or caregiver kept a daily record of the formula intake (in ounces), the amount of gas (using the following scale: 0, none, 1; small amount, 2; moderate amount; 3, large amount), the amount of liquid (using the following scale: 1, hard; 2, formed; 3, soft; 4, loose; 5, watery), the amount of stool (using the following scale: 0, none, 1, small amount, 2, moderate amount, 3, excessive amount), and fussiness (using the following scale: 0, not fussy; 1, slightly fussy; 2, moderately fussy; 3, very fussy; 4, extremely fussy). Adverse events were monitored throughout the study. Institutional review boards for each site reviewed and approved the protocol and procedures, and parents provided written informed consent. The study was conducted in compliance with good clinical practices.

**Fecal collection and sampling.** Study personnel collected infant stool specimens at baseline, on day 14, and on day 28 no more than 2 h following defecation by using chemical-free diapers (Tushies; TenderCare International, Eau Claire, WI). Samples were evenly divided and placed into two separate 15-ml transport containers, and the aliquots were treated as follows: (i) shipped under ambient conditions to a central clinical laboratory (North Coast Clinical Laboratory, Sandusky, OH) for enumeration of *Enterobacteriaceae* or (ii) immediately frozen at −20°C and shipped on cold packs to a second central laboratory (Department of Animal Sciences, University of Illinois, Urbana) for DNA isolation and microbial community analysis. Following storage for approximately 3 years at −20°C, the second set of fecal samples was shipped on dry ice to a third laboratory (Department of Medical Microbiology, University Medical Center, Groningen, The Netherlands) for relative quantification of bifidobacteria by FISH.

**TABLE 1.** qPCR standards and detection limits

<table>
<thead>
<tr>
<th>Bacterial group</th>
<th>Representative strain used for standard curve</th>
<th>Detection limit (log ng/μl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total bacteria</td>
<td><em>B. fragilis</em> lab strain</td>
<td>−2.0</td>
</tr>
<tr>
<td><em>Lactobacillus</em></td>
<td><em>L. delbrueckii</em> lab strain</td>
<td>−3.7</td>
</tr>
<tr>
<td><em>Bifidobacterium</em></td>
<td><em>B. adolescentis</em> lab strain</td>
<td>−4.0</td>
</tr>
<tr>
<td><em>Clostridium cluster I</em></td>
<td><em>C. paraputrefaciens</em> 16480</td>
<td>−3.0</td>
</tr>
<tr>
<td><em>Clostridium cluster XI</em></td>
<td><em>C. difficile</em> 17450</td>
<td>−4.0</td>
</tr>
<tr>
<td><em>Clostridium cluster XIV</em></td>
<td><em>C. coccoides</em> ATCC 29236</td>
<td>−6.0</td>
</tr>
</tbody>
</table>

**Bacterial strains and culture conditions.** Bacterial strains used to obtain the qPCR standard curves are listed in Table 1. All *Clostridium* strains and *Bacteroides fragilis* were cultured in reconstituted clostralid medium broth, *Lactobacillus delbrueckii* was cultured in de Man-Rogosa-Sharpe broth, and *Bifidobacterium adolescentis* was cultured in Beltsil Minimal broth supplemented with hemin and menadione. All strains were grown at 37°C and harvested during exponential growth. *Enterobacteriaceae* were cultured on MacConkey agar at 37°C for at least 48 h, and the results were expressed in CFU.

**DNA extraction.** Genomic DNA in fecal samples (200 mg) and bacterial cultures was isolated using a previously described method (33).

**Real-time qPCR.** Assays were performed using the ABI PRISM 7900HT sequence detection system (Applied Biosystems, Foster City, CA). The PCR amplification protocols were the protocols described previously for total bacteria (9) and the genera *Bifidobacterium* (22), *Lactobacillus* (19), and *Clostridium* (32). All reaction mixtures contained 1 ng template DNA, a specific primer set (Table 1), and bovine serum albumin (0.01 mg/ml). SYBR green PCR master mixture (2X; Applied Biosystems) was used for detection of total bacteria, *Bifidobacte- rium*, and *Lactobacillus*. For detection of *Clostridium*, TaqMan PCR master mixture was used, and the internal probe was labeled with 6-carboxylfluorescein as the 5′ reporter dye and 6-carboxytetramethylrhodamine as the 3′ quencher dye. Standard curves were generated using serial dilutions of genomic DNA extracted from pure cultures of representative bacteria (Table 1). The initial data were expressed as target DNA concentrations, which were normalized by expression relative to the total bacterial DNA concentration.

16S rRNA gene PCR-DGGE. Template DNA (10 ng) was amplified using universal bacterial 16S-V3 primers (Table 2) (a 40-bp GC clamp was added to the forward primer) as described previously (25). PCR amplicons were loaded in 35 to 60% linear DNA-denaturing gradient gels, and DGGE was performed using the Bio-Rad D-code system (Bio-Rad, Los Angeles, CA). The reference ladder contained PCR amplicons from pure cultures of *B. fragilis*, *E. rectale*, *L. delbrueckii*, and *B. adolescentis* from the top to the bottom (both the fourth and fifth bands were derived from *B. adolescentis* and consistently appeared on the gel). Following electrophoresis, gels were silver stained and scanned using a GS-710 calibrated imaging densitometer (Bio-Rad). Images were analyzed using Diversity Database software (Bio-Rad). Each band on the gel was identified regardless of the band intensity. Pairwise similarity was determined using the Dice coefficient (C,), where patterns with no common bands received a score of 0 and identical band patterns received a score of 1.0 (21). Dendrograms were generated by using Ward’s algorithm, in which samples were clustered by the degree of similarity to visualize community relatedness.

**FISH.** Quantification of bifidobacteria by FISH was carried out by first suspending 0.5 g of each fecal sample in 4.5 ml filtered phosphate-buffered saline (PBS) with three to five glass beads (diameter, 4 mm) and homogenizing the preparation with a vortex mixer for 3 min. Debris was removed by centrifugation at 1,000 rpm for 1 min, and the supernatant was fixed by diluting it 1:4 in freshly prepared 4% (wt/vol) paraformaldehyde in PBS and incubating it overnight at 4°C. Depending on the probable number, the fixed samples were diluted as necessary in PBS, and 10-μl aliquots were spread completely across the surfaces of 1-cm² wells in custom-made slides (CBN, The Netherlands) pretreated with a gelatin solution (0.1% gelatin (Oxoid), 0.01% KCl(SeO₄)₂·12H₂O (Sigma)), allowed to dry evenly for 30 to 60 min at room temperature, and fixed in 96% (vol/vol) ethanol for 10 min. Bifidobacteria were labeled by hybridization for 16 h at 50.0°C with the genus-specific Bfd164 probe (20) that was 5′ end labeled with fluorescein isothiocyanate (Pharmacia) and added at a concentration of 10 ng/μl in buffer (20 mM Tris-HCl, 0.9 M NaCl, 0.1% sodium dodecyl sulfate; pH 7.2). After hybridization, the slides were rinsed in buffer (20 mM Tris-HCl, 0.9 M NaCl) for 30 min at 50°C and then with Milli-Q water, dried rapidly using compressed air, and mounted using 6 μl Vectashield (Vector Laboratories, Burlingame, CA) in each well and a coverslip. Total bacteria were labeled in a
obtained at baseline, day 14, and day 28. CS and Ward’s algorithm calculations
measures analysis that included all feeding groups was performed using data
with the baseline level were performed using analysis of covariance. A repeated-
HR600 image analysis software (Leica).
to 25 fields with 30 to 100 positive objects) or automatically using the Quantimet
many), and fluorescent cells were counted either visually (average number in 10
captured using a Leica DMRA2 epifluorescence microscope (Wetzlar, Ger-
similar way by hybridization with the Eub338 probe (1). Digital images were
using ANOVA. Race, ethnicity, gender, and study discontinuance were analyzed
based on U.S. National Center for Health Statistics reference data (http:/www
entered into the analysis. Anthropometric measurements were converted to z-scores
for-age z-scores for each group either remained the same or
increased between the time of enrollment and day 28. There
The feeding groups were similar with respect to gender
distribution and age of the participants. There was a statisti-
cally significant difference in race only between the BF group
(83% white, 3% black, and 13% more than one race) and the
group (56% white and 44% black) (P = 0.002). The mean
formula intake values were similar during the 7-day
formula run-in period (195.0 to 214.8 ml/kg per day) and
during the 28-day feeding trial (206.3 to 235.0 ml/kg per day). The
means for weight-, length-, and head circumference-for-age
z-scores at enrollment and day 28 are shown in Table 3. Sta-
tistically significant differences in mean weight- and length-for-
age z-scores among feeding groups were detected at enrol-
ment and day 28 (Table 3 shows individual P values for all
pairwise comparisons of z-scores). Mean weight- and length-
for-age z-scores for each group either remained the same or
increased between the time of enrollment and day 28. There

TABLE 2. PCR primers and probes used for PCR-DGGE, qPCR, and FISH

<table>
<thead>
<tr>
<th>Target bacterial group</th>
<th>Primer or probe</th>
<th>Oligonucleotide sequence (5’→3’)</th>
<th>Annealing temp (°C)</th>
<th>Reference</th>
</tr>
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<tbody>
<tr>
<td>Total bacteria</td>
<td>341F</td>
<td>CTCAGGGGAGGCGAGCGAG</td>
<td>55–65</td>
<td>25</td>
</tr>
<tr>
<td></td>
<td>543R</td>
<td>ATTACCCGGTGCTGG</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Eub338</td>
<td>GCTGCTTCCCCGTAGGAGT</td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>Lactobacillus</td>
<td>Lab-0159</td>
<td>GGAAGACRGTRCTAATACCG</td>
<td>56</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>Univ-0515</td>
<td>ATCGTATTACCCGCGGTGCTGCCA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bifidobacterium</td>
<td>g-Bifid-F</td>
<td>TCCTCTGAAACGGGGTGG</td>
<td>55</td>
<td>22</td>
</tr>
<tr>
<td></td>
<td>g-Bifid-R</td>
<td>GGTTGTCCTCCCGATATCTACA</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Bif164</td>
<td>CATCAGGCGTACCTCTACCA</td>
<td>60.2</td>
<td>20</td>
</tr>
<tr>
<td>Clostridium cluster I</td>
<td>CI-F1</td>
<td>TACCHRAAGGGAAGGCCCAC</td>
<td>63</td>
<td>32</td>
</tr>
<tr>
<td></td>
<td>CI-R2</td>
<td>GTTCTTCTAATCTCCTACGAT</td>
<td></td>
<td></td>
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<tr>
<td>Clostridium cluster XI</td>
<td>CXI-F1</td>
<td>ACGCTACTTGGAGGAGGA</td>
<td>58</td>
<td>32</td>
</tr>
<tr>
<td></td>
<td>CXI-R2</td>
<td>GAGCCGTTAGCCTTTACT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clostridium cluster XIV</td>
<td>CXIV-F1</td>
<td>GAWGAAGATAAYCGTTATGT</td>
<td>52</td>
<td>32</td>
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<td></td>
<td>CXIV-R2</td>
<td>CTACGCWCCCTTTACAC</td>
<td></td>
<td></td>
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<tr>
<td>Clostridium</td>
<td>Probe-I</td>
<td>GTGCCAGCAGCGCAGGTAATACG</td>
<td>32</td>
<td></td>
</tr>
</tbody>
</table>

* The sequence contained an additional 40-nucleotide GC clamp for DGGE.

similar way by hybridization with the Eub338 probe (1). Digital images were
captured using a Leica DMRA2 epifluorescence microscope (Wetzlar, Ger-
day 14, and day 28 were performed using the Wilcoxon rank sum test. Comparisons of changes in the
of total bacteria were not normally
distributed; therefore, data were expressed as median values. Changes from the
baseline values among groups of formula-fed infants were analyzed using the
Wilcoxon rank sum test; the sign test was used for within-group comparisons.
Comparisons of values for all feeding groups at baseline, day 14, and day 28
were performed using the Wilcoxon rank sum test. Comparisons of changes in the
levels of Enterobacteriaceae (log_{10} CFU/g stool) for formula groups compared
with the baseline level were performed using analysis of covariance. A repeated-
measures analysis that included all feeding groups was performed using data
obtained at baseline, day 14, and day 28. C, and Ward’s algorithm calculations
were performed using Diversity Database software (Bio-Rad). C, were analyzed
by using analysis of variance (ANOVA). Feeding group and age were factored
into the analysis. Anthropometric measurements were converted to z-scores
based on U.S. National Center for Health Statistics reference data (http://www.cdc.gov/nccdphp/dnpa/growthcharts/resources/sas.htm) and were analyzed by
using ANOVA. Race, ethnicity, gender, and study discontinuance were analyzed
by using Fisher’s exact test. Analyses were performed by using SAS, version 8
(Cary, NC).

RESULTS

Infant growth, stool characteristics, and formula tolerance.
There were no statistically significant differences in study dis-
continuation rates among the feeding groups, and 117 infants
completed the study, including 29 infants in the BF group, 21
infants in the control group, 23 infants in the PG4 group, 23
infants in the PGL4 group, and 21 infants in the PGL8 group.
One infant (PGL8 group) did not consume the study formula
after randomization and was not included in any of the anal-
yses, and one infant (BF group) did not meet the gestational
age requirement and was not included in analyses of intestinal
bacterial composition, stool characteristics, and tolerance.

TABLE 3. Mean weight-, length-, and head circumference-for-age
z-scores at enrollment and day 28

<table>
<thead>
<tr>
<th>Time</th>
<th>Group (n)</th>
<th>Mean z-score for age&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Enrollment</td>
<td>BF (30)</td>
<td>0.5 A</td>
</tr>
<tr>
<td></td>
<td>Control (25)</td>
<td>0.0 B</td>
</tr>
<tr>
<td></td>
<td>PG4 (27)</td>
<td>0.4 AB</td>
</tr>
<tr>
<td></td>
<td>PGL4 (27)</td>
<td>0.0 B -0.1 B</td>
</tr>
<tr>
<td></td>
<td>PGL8 (25)</td>
<td>0.1 A</td>
</tr>
<tr>
<td>Day 28</td>
<td>BF (29)</td>
<td>0.6 AB</td>
</tr>
<tr>
<td></td>
<td>Control (22)</td>
<td>0.2 AB</td>
</tr>
<tr>
<td></td>
<td>PG4 (24)</td>
<td>0.7 BC</td>
</tr>
<tr>
<td></td>
<td>PGL4 (24)</td>
<td>0.4 AC</td>
</tr>
<tr>
<td></td>
<td>PGL8 (21)</td>
<td>1.0 B</td>
</tr>
</tbody>
</table>

<sup>a</sup> The standard error for all means was 0.2. In each column means for the same
time point (enrollment or day 28) followed by the same letter are not significantly
different (P ≥ 0.05).
were no significant differences in mean head circumference-for-age z-scores at enrollment or at study day 28.

There were no significant differences among the feeding groups in the numbers of infants who experienced constipation or diarrhea during the run-in period or feeding trial (data not shown). The stool frequency (mean ± standard error) was significantly higher for the BF group (3.6 ± 0.3 stools/day) than for any of the formula groups (1.6 to 2.3 ± 0.3 stools/day) (P ≤ 0.018) during the run-in period; there were no significant differences among feeding groups in average mean stool frequency from baseline to day 28 (1.9 to 2.8 ± 0.3 stools/day).

The daily stool consistency scores (mean ± standard error) were significantly higher for the BF group (3.8 ± 0.1) than for all of the formula groups (control group, 3.3 ± 0.1; PG4 group, 3.3 ± 0.1; PGL4 group, 3.2 ± 0.1; PGL8 group, 3.1 ± 0.1) (P ≤ 0.004) during the run-in period; the scores for the BF group (3.8 ± 0.1) and each formula group were also significantly different when the averages from baseline to day 28 were compared (control group, 3.2 ± 0.1; PG4, PGL4, or PGL8 group, 3.4 ± 0.1) (P < 0.05). The only significant difference in the amount of gas (mean ± standard error) was the difference between the BF group (0.9 ± 0.1) and the PGL8 group (1.4 ± 0.1) (P = 0.006) from day 15 to day 21.

There were no significant differences among the feeding groups in the amount of fussiness (mean ± standard error) during the run-in period or feeding trial (1.0 to 1.3 ± 0.1). There were no differences among the study groups in the number of participants who experienced at least one adverse event, and there were no differences in the number of participants who experienced adverse events when body systems or individual events were analyzed. There were no adverse events that had particular clinical significance in any feeding group (data not shown).

**Quantification of Enterobacteriaceae.** Selective plating of fecal samples on MacConkey agar was used to enumerate fecal Enterobacteriaceae populations in all feeding groups (log_{10} CFU/g stool). No significant differences in the mean changes in the Enterobacteriaceae populations from baseline to day 28 were detected among the formula groups (control, PG4, or PGL4 group, 0.2 ± 0.1 log_{10} CFU/g stool; PGL8 group, 0.3 ± 0.1 log_{10} CFU/g stool). Within the PGL8 group there was a slight but statistically significant increase in the mean level of Enterobacteriaceae (P = 0.027) from baseline to day 28, but there were no significant changes within the other feeding groups. The means for Enterobacteriaceae for the different feeding groups at baseline, day 14, and day 28 are shown in Table 4. The mean for the BF group was significantly higher than the means for all other groups at baseline (P < 0.001). Significantly higher levels of Enterobacteriaceae were also observed on day 14 for the BF group than for the prebiotic-supplemented groups (P ≤ 0.002). The Enterobacteriaceae levels for the feeding groups at day 28 were similar.

**Quantification of fecal bacterial groups using real-time qPCR.** To compare the effects of prebiotics in infant formulas fed over a 28-day period on fecal bacterial composition, qPCR was used to determine the levels of *Clostridium* clusters I, XI, and XIV, *Lactobacillus* spp., and *Bifidobacterium* spp. expressed as percentages of the total bacteria. Data for eight samples were not included in the statistical analyses due to an unreasonably low density (four samples) or no detection (four samples) of total bacteria by qPCR. No significant differences among the formula groups in changes in the bacterial subpopulations from baseline to day 28 were observed for *Clostridium* cluster I, cluster XI, or cluster XIV, the sum for *Clostridium* clusters I and XI, *Bifidobacterium* spp., or *Lactobacillus* spp. As determined by using the sign test, a significantly greater number of samples exhibited decreases in the levels of *Clostridium* cluster XI (P = 0.049) and *Lactobacillus* spp. (P = 0.013) in the PGL8 group from baseline to day 28 (data not shown).

Median values for *Clostridium* clusters, *Lactobacillus* spp., and *Bifidobacterium* spp. for each feeding group at baseline, day 14, and day 28 are shown in Table 5. No significant differences among feeding groups in *Clostridium* cluster I values were observed at any time point. The median values for *Clostridium* cluster XI were significantly lower for the BF group than for the PG4 group (P = 0.006), the PGL4 group (P < 0.001), and the PGL8 group (P = 0.016) at baseline, for all other groups at day 14 (control group, P = 0.011; PG4 group, P < 0.001; PGL4 group, P = 0.003; PGL8 group, P = 0.019), and for the PG4 group (P = 0.005) and the PGL4 group (P = 0.015) at day 28. No significant differences among feeding groups for *Clostridium* cluster XIV, the sum for *Clostridium* clusters I and XI, *Lactobacillus* spp., or *Bifidobacterium* spp. were observed at any time point.

**Repeat quantification of the fecal bifidobacterial subgroup by FISH.** Given the generally low level of bifidobacteria detected by qPCR, fecal samples were reanalyzed by performing FISH using a probe specific for this subgroup. Compared to qPCR, FISH detected considerably higher median levels of bifidobacteria, expressed as a percentage of the total bacteria. The only significant difference was detected for samples collected during the baseline phase of the study, in which bifidobacteria were found to account for 20.7 and 83.5% of the total bacteria in the PGL8 group and human milk reference group, respectively (Table 5). No other significant differences in levels of bifidobacteria were observed for the various feeding groups at any time.

**Profiling of bacterial communities using DGGE.** Bacterial community profiles were generated by 16S rRNA gene PCR-DGGE. Samples collected from the same subject at baseline, day 14, and day 28 were run on the same gel to maintain electrophoretic consistency (representative results are shown in Fig. 1). Cₛ were calculated to quantify the pairwise similarity of DNA banding patterns within (intragroup) and among (intergroup) feeding groups at baseline, day 14, and day 28. The levels of intergroup similarity at baseline (Cₛ range, 18.0 to 23.4) were higher than the mean levels for all other groups at baseline (P ≤ 0.001).}

<table>
<thead>
<tr>
<th>Time</th>
<th>Conc of Enterobacteriaceae (log_{10} CFU/g stool)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>BF group</td>
</tr>
<tr>
<td>Baseline</td>
<td>7.9 A</td>
</tr>
<tr>
<td>Day 14</td>
<td>7.7 A</td>
</tr>
<tr>
<td>Day 28</td>
<td>7.5 A</td>
</tr>
</tbody>
</table>

*a* The standard error for all means was 0.1. For each time means followed by the same letter are not significantly different (P ≥ 0.05).
19.2), day 14 (Cs range, 19.2 to 21.7), and day 28 (Cs range, 16.4 to 19.9) were low. The intragroup Cs values were within similar ranges at all time points (data not shown). The intragroup and intergroup Cs values were not significantly different at any time point. A cluster analysis was performed using Ward's algorithm, and dendrograms were constructed to visualize relationships among banding patterns for samples. Samples obtained from the same subject at all time points examined tended to cluster together. No specific clustering patterns were observed for feeding groups, ages, or races (data not shown).

DGGE profiles generated for individual subjects at baseline, day 14, and day 28 were compared across three intervals to index the bacterial stability in response to the feeding regimen during the study period. Individual DGGE profiles obtained at different time points were compared to calculate the Cs value, which was used as an index of bacterial community stability; a higher Cs indicated greater bacterial stability and less population fluctuation during the corresponding feeding period. An analysis of Cs values by using ANOVA with feeding group and age as independent variables indicated that there was a significant age effect but there was not a feeding effect (data not shown). Because of the lack of a feeding effect during the 28-day period, a post hoc analysis was conducted, in which individuals in each feeding group were classified by age to obtain data for younger infants (21 to 50 days old at the beginning of the study) and older infants (51 to 104 days old at the beginning of the study). From baseline to day 28, the mean Cs values were lower for the younger infants in the BF group (P = 0.054) and significantly lower for the younger infants in all three supplemented-formula feeding groups (PG4 group, P = 0.048; PGL4 group, P = 0.040; and PGL8 group, P = 0.006) (Table 6). The mean Cs values for younger and older infants in the control group did not differ from baseline to day 28 (P = 0.511).

DISCUSSION

This study investigated the effects of prebiotic blends containing PDX and GOS (4.0 g/liter) or PDX, GOS, and LOS (4.0 or 8.0 g/liter) on fecal bacterial populations over a 28-day period in healthy term infants. Numerous clinical investigations of preterm and term infants who received formulas supplemented with GOS alone (3) or with GOS and FOS at a 9:1 ratio (2, 6, 7, 14, 18, 19, 23, 24, 31) at a wide range of concentrations (2.0 to 10 g/liter) have shown that the levels of *Bifidobacterium* spp. in the fecal bacterial population were higher in these infants than in infants who received unsupplemented...
formulation. This is the first clinical evaluation of the effects of new prebiotic blends of PDX, GOS, and LOS on the infant fecal microbiota.

Small but statistically significant differences in mean weight- and length-for-age z-scores were noted among feeding groups at enrollment and study day 28. For all groups, however, the mean weight- and length-for-age z-scores either remained the same or increased between enrollment and day 28, indicating that each group was growing normally. Consequently, differences among feeding groups were not considered clinically significant. The current study was not designed to evaluate growth as a primary clinical outcome. Ziegler et al. (35), however, fed infants from days 14 to 120 old the formulas given to the control, PG4, and PGL8 groups in a study designed specifically to evaluate infant growth according to the American Academy of Pediatrics Task Force recommendations for clinical testing of infant formulas (11). These workers found no differences in the rates of increases in weight or length over the duration of the 120-day study and concluded that infants fed these prebiotic-containing formulas exhibited normal growth (35). Stools from infants in the BF group in the current study were found to be softer and looser than stools from infants in groups that received formula. However, the stool consistency scores changed for the groups of infants who received prebiotic-supplemented formula over the 28-day feeding trial and approached the stool consistency scores for the breast-fed infants. Ziegler et al. (35) also reported that infants fed formula supplemented with prebiotics had stool characteristics more similar to those of breast-fed infants than to those of infants fed unsupplemented formula. Overall, infant formulas containing blends of PDX, GOS, and LOS were generally well tolerated and safe.

Few significant differences in the overall bacterial composition were found from baseline to day 28 within feeding groups or among feeding groups at any time point. Compared to the traditional culture methods used in previous prebiotic feeding studies, the more sensitive molecular methods for quantifying bacteria in complex microbial communities may make broad differences between breast-fed and formula-fed infants less apparent. For example, a recent examination of fecal samples from breast-fed and formula-fed infants using qPCR demonstrated that there were no significant differences in the *Bifidobacterium* levels (28). Although the levels of bifidobacteria tended to be higher in the BF group throughout the current study, the levels of neither bifidobacteria nor lactobacilli were significantly different in the formula-fed infants and the infants fed breast milk at any time point, with one exception. The exception occurred in the baseline phase, when fecal samples from infants in the PGL8 group contained significantly less bifidobacteria (20.7%) than fecal samples from infants in the human milk reference group (83.5%), as determined by FISH. The general lack of significance could have been due to the great variation in the data, which could be attributed to the wider range of infant ages (21 to 104 days at baseline) and the shorter feeding period (28 days) compared with other similar prebiotic studies.

In addition, subtle differences between the microbiota of breast-fed infants and the microbiota of formula-fed infants might occur only at the species level, where species in certain subpopulations of the gut microbiota shift toward the pattern observed in breast-fed infants (28, 30). Rinne et al. (30) demonstrated that the total number of bifidobacteria was significantly lower in infants who received control formula than in infants who received breast milk or a prebiotic-supplemented formula (GOS and FOS at a ratio of 9:1); additionally, species-specific PCR revealed that the *Bifidobacterium* species composition obtained for the breast-fed infants was found in infants who received prebiotic-supplemented formula. Using a species-specific qPCR assay, Penders et al. (28) found a significantly lower prevalence and significantly lower numbers of *C. difficile*, a member of *Clostridium* cluster XI, in breast-fed infants than in formula-fed infants. In the current study, a significantly greater number of subjects in the PGL8 group exhibited a decrease of the number of *Clostridium* cluster XI organisms between baseline and day 28. However, no species-specific prebiotic effects, such as a decrease in the level of a pathogen or similarity in species diversity, could be attributed to supplementation of infant formula with GOS, PDX, and LOS with the group-specific primers used.

The level of bifidobacteria observed expressed as a percentage of the total bacterial DNA in this study appeared to be low compared with the common assumption that *Bifidobacterium* spp. dominate the infant fecal microbiota. However, the results obtained by using a DNA concentration-based standard curve in the qPCR analysis may not have accurately reflected the number of bacterial cells in the complete microbial community. To estimate the number of bifidobacterial cells in the fecal microbiota, we performed post hoc FISH using *Bifidobacterium*-specific and universal bacterial probes. The percentages of bifidobacterial cells in the original samples indicated by the FISH results were much higher than the percentages estimated by DNA concentration-based qPCR. Consistent with the qPCR results, the percentages of bifidobacteria were higher in BF group than in the other groups, but there were few statistically significant differences among groups due to the great variation among subjects within groups. The difference between the percentages of bifidobacteria estimated by FISH and qPCR can probably be partially explained by inherent differences in methodology. *Bifidobacterium* spp. have relatively low numbers of 16S rRNA gene copies compared with other major intestinal bacterial species, which leads to potential underestimation by 16S rRNA gene-targeted qPCR. Palmer et al. (27) also observed a relatively low frequency and low abundance of bifidobacteria in infant fecal samples based on the results of molecular methods, including qPCR (as in this study) and

**TABLE 6. Cs for individual DGGE profiles between baseline and day 28**

<table>
<thead>
<tr>
<th>Group</th>
<th>Cs for different agesa</th>
<th>P valueb</th>
</tr>
</thead>
<tbody>
<tr>
<td>BF</td>
<td>0.49 ± 0.26</td>
<td>0.68 ± 0.23</td>
</tr>
<tr>
<td>Control</td>
<td>0.56 ± 0.25</td>
<td>0.62 ± 0.17</td>
</tr>
<tr>
<td>PG4</td>
<td>0.40 ± 0.16</td>
<td>0.60 ± 0.27</td>
</tr>
<tr>
<td>PGL4</td>
<td>0.41 ± 0.22</td>
<td>0.60 ± 0.20</td>
</tr>
<tr>
<td>PGL8</td>
<td>0.38 ± 0.17</td>
<td>0.60 ± 0.11</td>
</tr>
</tbody>
</table>

a The values are means ± standard deviations.

b P values for comparisons of younger and older infants in the same groups. No significant differences among age-matched feeding groups were detected by ANOVA.
microarray analysis, but in the latter analysis the primers used for broad-range amplification of rRNA genes were suboptimal and underestimated bifidobacteria eightfold. Another explanation for the low level of bifidobacteria detected by qPCR in the current study may be the difference in DNA extraction efficiency between gram-positive and gram-negative bacteria, which may result in a bacterial DNA composition bias. The differences in the percentages of bifidobacteria reported in different studies emphasize the importance of reevaluating PCR-related biases, including DNA extraction bias, in order to obtain more accurate estimates of the contribution of this bacterial group to the intestinal microbiota of infants.

An unexpected finding was that the BF group had significantly higher levels of Enterobacteriaceae than the formula-fed groups at the beginning of the study, since the levels of Enterobacteriaceae are reportedly lower in breast-fed infants (28). However, by the end of the study period, the levels of Enterobacteriaceae were comparable for all feeding groups, suggesting that neither prebiotic carbohydrates nor human milk oligosaccharides influenced the growth of these bacteria.

Analysis of the 16S rRNA gene PCR-DGGE results for infant fecal samples revealed little similarity in the bacterial community profiles among subjects regardless of the feeding practice. The calculated C, for DGGE banding patterns within or among feeding groups were low, and significant differences were not observed. Cluster analysis of DGGE profiles demonstrated that over time the individual bacterial community profiles clustered by subject rather than by feeding group. Significant differences in bacterial community stability were not observed among feeding groups during the 28-day feeding period, whereas a significant age effect was observed. A post hoc analysis of younger and older infants within each group revealed that the shifting microbial populations in younger infants in the PG4, PGL4, and PGL8 groups, as well as the BF reference group, were less stable from baseline to day 28. In contrast, the C, values for the control group appeared to be relatively high for both younger and older infants, indicating that the overall microbial profile was reasonably stable over the 28-day feeding period regardless of age. These age-related differences might have been due to the presence of prebiotics in the formulas and the presence of human milk oligosaccharides in breast milk, as corresponding age-related differences were not observed for the control group. Although further age-based statistical analyses were not performed, the results suggest that the prebiotic blends had a greater effect on the fecal bacterial communities of younger infants than on the fecal bacterial communities of older infants and imply that infant age is a variable that is more important than originally expected.

With the current study design, few statistically significant differences in the composition of fecal bacterial populations were observed in the infants who received formula supplemented with GOS, PDX, or LOS prebiotic blends compared with the infants who received a control formula or the reference BF group. However, the age-related differences in fecal bacterial community stability observed within the PG4, PGL4, and PGL8 feeding groups indicate that these prebiotic blends may have a greater impact on infant fecal bacterial populations in younger infants than in older infants. Further analysis is necessary before definitive conclusions concerning this possibility can be reached.

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REFERENCES