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***Clinical Trials Study***

**Hypoallergenic formula with *Lactobacillus rhamnosus GG* for babies with colic: A pilot study of recruitment, retention, and fecal biomarkers**

Fatheree NY *et al*. Probiotic formula for infants with colic

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**Abstract**

**AIM:** To investigate recruitment, retention, and estimates for effects of formula supplementation with *Lactobacillus rhamnosus GG* (LGG) on inflammatory biomarkers and fecal microbial community in infants with colic.

**METHODS:** A prospective, double-blind, placebo-controlled trial was conducted in otherwise healthy infants with colic. We screened 74 infants and randomized and analyzed results in 20 infants [9 receiving LGG (LGG+) and 11 not receiving LGG (LGG-)]. LGG was incorporated in the formula (Nutramigen®) (minimum of 3 × 107 CFU’s daily) in the LGG+ group. Fecal microbiota and inflammatory biomarkers, including fecal calprotectin, plasma cytokines, circulating regulatory T cells (Tregs), and crying + fussing time were analyzed to determine optimal time points and effect sizes for a larger trial.

**RESULTS:** Recruitment in this population was slow, with about 66% of eligible infants willing to enroll; subject retention was better (75%). These rates were influenced by parents’ reluctance to volunteer their infant for a clinical trial and by their tendency to change formulas. The maximal difference of crying + fussing time was observed at day 14, comparing the 2 groups, with a mean difference of -91 (95%CI: -76, 259) min (*P* = NS). Fecal calprotectin showed no significant difference, but the optimal time to determine a potential effect was at day 90 (with a mean difference of 121 (95%CI: -48, 291) g/g stool), observing a lower level of fecal calprotectin in the LGG+ group. The fecal microbial communities were chaotic, as determined by Shannon’s diversity index and not apparently influenced by the probiotic. No significant change was observed in plasma inflammatory cytokines or Tregs, comparing LGG+ to LGG- groups.

**CONCLUSION:** Designing future colic trials involving a probiotic-supplemented formula for infants in the United States will require consideration for difficult enrollment. Infants with colic have major variations in feal microbiota and calprotectin, both of which improve with time, with optimal time points for measurement at days 14 and 90 after treatment.

**Key words:** Barr diary; Crying; Fussing;Probiotic; Regulatory T cells; Cytokines; Newborn; Intestine; Inflammation; Biomarker

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**Core tip:** The “dysbiosis” theory proposes that newborns with abnormal colonization are predisposed to having gut inflammation and colic. Probiotics may reduce crying and diversify the fecal microbiota. A prospective, double-blind, placebo-controlled trial was conducted in healthy infants with colic. After 75% screen failure or dropouts, 20 infants were analyzed (9 receiving formula with *Lactobacillus* GG and 11 not receiving *Lactobacillus rhamnosus GG* in their formula). We found that: (1) recruitment/retention indicate future randomized controlled trials should enroll 80 patients with an optimal timepoint for observing a potential difference in crying at 14 d; (2) microbial communities were chaotic in infants with colic, even more so than reported in Dutch infants; and (3) our study was the first to analyze cytokine levels and circulating Tregs in infants with colic.

Fatheree NY, Liu Y, Ferris M, Van Arsdall M, McMurtry V, Zozaya M, Cai C, Rahbar MH, Hessabi M, Vu T, Wong C, Min J, Tran DQ, Navarro F, Gleason W, Gonzalez S, Rhoads JM. Hypoallergenic formula with *Lactobacillus rhamnosus GG* for babies with colic: A pilot study of recruitment, retention, and fecal biomarkers. *World J Gastrointest Pathophysiol* 2015; In press

**INTRODUCTION**

Colic has been defined as inconsolable crying and fussing, for greater than 3 h daily for more than 3 days per week in infants from 3 weeks to 3 months of age[1]. There have been many theories to explain the occurrence of colic, including bacterial overgrowth[1], the “fourth trimester” theory in which the baby is wishes to remain *in utero*[2], parental depression[3], excessive intestinal gas[1], and milk protein allergy[4]. Savino was the first to propose that an abnormal microbiota (“dysbiosis”) might be an important pathophysiological mechanism, by demonstrating a reduced abundance of *Lactobacilli* and increased abundance of *E. coli* in the stools of infants with colic[5].

Our previous studies showed that intestinal inflammation was a feature of colic[6]. This led to the hypothesis that there may be an association of dysbiosis and intestinal inflammation, both of which may improve with probiotic treatment. The probiotic *Lactobacillus rhamnosus GG* (LGG) has been shown to reduce diarrhea in children with acute infectious enteritis[7] and to facilitate the development of a more diverse fecal microbiota[8]. Others have hypothesized that early colonization of the immature small intestine with lactobacillus would reduce gut inflammation and symptoms in infants with colic. However, previous studies have focused on breast-fed babies.

In the current studies, our two major aims were (1) to investigate the feasibility of recruitment and retention of babies with colic randomized to receive a probiotic-containing formula; and (2) to determine effect size of LGG-supplemented formula on crying + fussing time, the intestinal microbiota, and the inflammatory biomarker calprotectin in infants with colic.

**MATERIALS AND METHODS**

***Ethical review and approval***

This protocol was approved by the Institutional Review Board of the University of Texas Health Science Center at Houston. Every 10 patients, a Data Safety Monitoring Board convened to review safety. A consort checklist is available as supporting information. The trial protocol was registered in <http://www.clinicaltrials.gov> NCT01279265.

***Study design, population and randomization***

This study was a prospective, double-blind, placebo-controlled trial in otherwise healthy infants with colic, designed as a pilot study for determining potential effects of treatment with casein-hydrolysate formula with LGG versus no LGG on selected biomarkers in infants with established colic. This formula (Nutramigen) was chosen because at the time it was the only formula that could be obtained in liquid form with or without probiotic, because it has been suggested to be beneficial in infants with colic[4], and because FDA-monitored safety and biomarker trials of direct probiotic supplementation in infants with colic had not been completed at this point. Partially of fully formula-fed infants age 3-13 wk old born full-term (> 37 wk gestation) were included if they fulfilled the colic definition of crying and fussing more than 3 h per day for at least 3 d weekly, documented at enrollment by at least 2 abnormal Barr diaries over a 3-d period[9]. Patients were actively recruited through university community clinics, 4 pediatric practices affiliated with our university, the pediatric gastroenterology clinic, and other local pediatricians *via* mailings, television coverage, and a website. Infants were excluded if they had failure to thrive, chronic lung disease, diarrhea, fever, and if they took a probiotic prior to enrollment.

Infants were randomly assigned by block randomization (groups of 4) to one of two formulas, either casein hydrolysate (Nutramigen®) with LGG (EnfloraTM) or casein hydrolysate without LGG (Nutramigen A+®). Initially, the protocol was to enroll children with or without colic to receive the above 2 formulas in white, unlabeled containers and to measure crying + fussing time and the biomarkers, but the protocol had to be changed because they were uncomfortable with this type of label. We changed the protocol so that containers were partially covered with a sticky label to ensure blinding. The randomization schedule was computer-generated, prepared by the study biostatistician and implemented by pharmacists in Department of Investigational Drugs Services (IDS) at Memorial Hermann Hospital. Infants with partial breast-feeding were required to take at least 240 mL of formula per day to ensure at least 3.6 x 107 CFU’s of LGG (if they were randomized to the LGG+ group). However, most infants were completely formula-fed (as shown in Table 1). The infants were required to take study formula for the entire 90 days of observation, with research visits on days 1, 14, 42, and 90. Patients were followed by telephone on a weekly basis. During each clinical visit, the medical history and clinical condition of each infant was evaluated by a pediatric gastroenterologist. Stool and blood were collected at baseline and follow up visits.

***Barr diary***

Infant crying and fussing time was quantified using the Barr Diary, a well-validated instrument, as previously described[9] at each study visit.

***Breath tests***

Parents/guardians were asked to have the infant fast for a minimum of 3 h before the baseline visit and before visit 2. After two baseline samples were collected (separated by 15 min), the infant was fed 60 mL of glucose water, and at time = 45 min, exhaled air was collected and breath hydrogen and methane were measured using the Quintron Model SC MicrolyzerTM (Quintron Instrument Co., Inc., Milwaukee, Wisconsin). In all infants, breath methane was negligible. A breath test was considered positive if the baseline hydrogen level was ≥ 20 parts per million (ppm) or if there was an increase from baseline of ≥ 12 ppm[6].

***Research lab protocols***

*Fecal calprotectin.* Stool samples was prepared and analyzed by using a quantitative calprotectin ELISA kit according to manufacturer’s instructions, as previously described[10]. The level of calprotectin was expressed as μg/g of stool weight.

**Plasma cytokines and percentage of Tregs:** Plasma cytokines were detected by using MSD Human ProInflammatory 7-Plex Ultra-Sensitive Kit (Meso Scale Discovery®, Gaithersburg, MD) which measures human IFN-γ, IL-1β, IL-6, IL-8, IL-10, IL-12p70, and TNF-α[10]. This inflammatory panel was chosen because LGG has been shown to prevent enterocyte apoptosis induced by IFN-γ, IL-1β, IL-6, IL-8, IL-10, IL-12p70, and TNF-α[11] Isolated peripheral blood mononuclear cells (PBMCs) were stained with surface CD4 and intracellular FOXP3 antibodies and analyzed by using flow cytometry[10].

**Fecal pyrosequencing analysis:** Parents were instructed to collect a stool sample within 48 h of the visit and to store stools frozen. In the lab, stool samples were subdivided and stored at -80 °C until analyzed. DNA extraction, PCR-amplification, pyrosequencing and taxonomic identification of 16S rRNA gene sequences in stool specimens were performed as previously described[12], using QIIME[13] to analyze microbial communities. A total of 185373 reads with an average length of 460 ± 62 bases were included in this study. The average number of reads per sample was 3783.

***Statistical analysis***

**Sample size and power:** This pilot study aimed to determine recruitment, retention, adverse events, and biomarkers but was not powered to detect differences in crying time between the two study arms. (Based on our previous study[6] showing 297 ± 142 min/d as a mean crying+fussing time in colicky infants, we determined that a sample size of 60 colicky infants (30 infants per study arm) would have been required to detect a mean difference of 100 min/d of crying + fussing time, with a power of 0.80 at the 5% level).

Baseline characteristics were compared between two groups using the two sample t-test or Wilcoxon rank sum test for continuous variables and the Fisher’s exact test for categorical variables. For estimating the effect size of Barr diary crying time and fecal calprotectin, we used generalized Estimating Equation (GEE) method with autoregressive covariance structure to account for potential correlation between measures at multiple visits during the follow up period. Time variable (*i.e.,* visits), treatment group indicator, as well as their interactions are included as covariates to estimate the differences in the outcomes between the two study arms (*i.e.,* LGG- values minus LGG+ values) over time. The adjusted means and the mean differences between LGG- *vs* LGG+ groups as well as their 95%CI were calculated. The Wilcoxon rank sum test was applied to compare the cytokines at baseline and at each of the follow up visits. All the above analyses were conducted using statistical software SAS 9.3 (SAS Institute, Cary, NC). Microbiota data were analyzed using Mann-Whitney *U-*test and one-way ANOVA using GraphPad Prism v5.0 for windows (GraphPad Software, San Diego, CA). Shannon’s Diversity Index was calculated using a locally developed pyrosequencing pipeline[12].

**RESULTS**

***Clinical characteristics***

Enrollment took place from September 2011 to January 2013. Seventy-four infants with colic were screened based on the above inclusion and exclusion criteria. Forty-four (59%) infants were not included in the study because they did not come to the scheduled first clinic visit or because the parents/guardians chose not to participate. Thirty were enrolled, but 4 changed their mind after signing consent, and 6 were uncomfortable when they received cans of formula with a white label. Subsequently, we changed by taping over the label to hide only the probiotic part of the label (The study initially was designed to include 30 infants in each group, but it was closed because of lack of funds to continue.). Twenty infants randomized to either the LGG+ group (*n* = 9) or to the placebo (LGG- group) (*n* = 11) group were able to be analyzed (Figure 1, Consort Diagram). Baseline characteristics including mean gestational age, age at enrollment, weight, and length showed no differences between the two groups (Table 1). Feeding method was exclusive formula feeding in 70%, although 3 in each group were receiving limited supplementation with breast milk. A wide range of formulas were given prior to enrollment, none of which contained a probiotic. Almost 5-h daily of crying and fussing provided evidence of severe symptoms in these infants.

***Clinical course***

Longitudinal analysis of outcome variables indicated that total crying + fussing times at baseline were comparable, and at each of the three treatment visits crying + fussing time decreased (Table 2). The maximal difference of crying + fussing time was observed at visit 2 (day 14) comparing the 2 groups, with a mean difference of -91 (95% CI: -76, 259) min, trending toward a shorter crying+fussing time in the LGG+ group. This difference was entirely the result of improved fussing time. After the immediate dropout following written consent of 4 infants, we observed a loss of 5 of the 20 children during follow-up. These children dropped out because of mild diarrhea (*n* = 1) or their parents’ decision to change the formula (*n* = 4). No adverse events during the period of observation were deemed attributable to the study product (LGG). These findings indicate that in studies of babies with colic, dropout is a common problem, with formula-changing being the major reason.

***Fecal calprotectin***

We previously demonstrated an elevated fecal calprotectin in babies with colic, compared to age-matched babies without colic[6]. Longitudinal analysis of fecal calprotectin at baseline and at follow up visits showed that the values were similar at baseline (Table 2), while the maximal mean difference in fecal calprotectin between the LGG+ and LGG- groups was seen at visit 4 (90 d of probiotic formula treatment), with a difference of -121 (-48, 291) μg/g stool, observing a statistically nonsignificant lower level of fecal calprotectin in the LGG+ group.

***Fecal microbiota***

**Distribution of predominant bacterial taxa in colicky infants:** No differences in fecal diversity at any of the visits were observed between the infants with LGG+ and LGG- formulae. At baseline, the most abundant bacterial phylum was Firmicutes (72%); followed by Proteobacteria (24%). Enterobacteriales was the most abundant order and Enterobacteriaceae the most abundant family in these infants. The genus level analysis at enrollment showed the major genera to be *Blautia*, *Escherichia/Shigella*, *Enterococcus*, *Streptococcus*, and *Coprobacillus* (Figure 2). There was no significant difference in averaged genus level microbial composition distribution between (LGG+)-fed colicky babies compared to (LGG-)-fed babies with regard to the major taxa. Minor differences might have been missed because of small sample size.

**Abundance of *L. rhamnosus* in stool specimens:**Previously, investigators reported that a 2-wk trial of LGG at similar doses resulted in positive LGG cultures of the stool of 85% of healthy infants during its administration and in half when measured 28d later[14]. Here, *L. rhamnosus* abundance increased to about 5% of total bacteria after 14d of LGG+ formula treatment (*P* = 0.006), which was significantly higher than baseline, and also higher than on visit 3 or visit 4 (*P* < 0.05) (Figure 3A).

**Diversity of microbiota and changes over time:** We had the opportunity to analyze the evolution of microbial communities of a set of dizygotic twins at the genus level, as shown in Figure 3B. One infant was randomized to the LGG- group, while the other was randomized to LGG+. Note that we were unable to obtain baseline sequence data for one twin despite two attempts, but data were available for both at each follow-up visit. At each of the *treatment* visits, we were able to compare the microbiota of the two infants, and we found that *Lactobacillus* was present in the twin babies’ stools at all follow-up visits. It was most abundant in the LGG+ infant at the 2nd visit. However, lactobacillus abundance declined to low levels in both infants by visit 4. Comparing visits 2, 3 and 4 in these two infants, the stool composition fluctuated greatly. However, comparison of the microbiota of twin A and B revealed remarkable convergence of the community profile among the twins, most evident at visit 4.

**Shannon’s diversity index:** varied wildly in all infants over time. Figure 4 shows data for all infants whose parents brought stools at each of the 4 clinic visits. There were no major differences noted between LGG+ and LGG- infants, because in both groups diversity fluctuated greatly from visit to visit. Thus, fecal microbial community was “chaotic” and did not show stabilization by the end of the 3-month study, when the infants were 4-5 months old.

***Other clinical and basic lab testing***

There were no major differences between the LGG+ and LGG- groups with respect to breath hydrogen levels (data not shown), plasma cytokine levels, or percentage of circulating Tregs (Table 3). With respect to breath hydrogen, 5 of the 20 infants (25%) had breath hydrogen increases of ≥ 12 ppm above baseline at their initial sampling, while two infants (10%) had increased breath hydrogen at visit 2. However, there was no significant difference between the LGG+ and placebo groups with respect to breath hydrogen changes, and there was also no correlation between changes in breath hydrogen and crying time.

**DISCUSSION**

***Recruitment and retention***

Recently published in JAMA Pediatrics, Sung’s systematic review of probiotics for colic emphasized that there remains “insufficient evidence to support probiotics to manage colic, especially in formula-fed infants”[15]. Our study which focused on this group, was not powered to determine efficacy of LGG+ formula, but we aimed to identify key information for future prospective randomized, controlled trials of treatments for infants with colic in the United States. Although not focused on colic, a recent double blind placebo-controlled trial focused on preterm infants (gestational age 32-36 wk.) who were randomized to receive LGG (> 109 cfu’s daily), a combination prebiotic or placebo. Authors reported that these infants were more likely to be contented (as opposed to “excessive criers”) during the first 2 months of life if they received prebiotic or LGG as compared to placebo[16]. Barr diaries were not used in this study. In addition to Sung’s trial, our results may be included in meta-analyses of probiotic supplementation for formula-fed infants with colic.

One key finding was that recruitment of babies with colic in this country is difficult. During the consent process, two-thirds of parents we interviewed declined to particiate in this trial. Possible reasons included the FDA- and IRB-mandated consent form (which contained the statement “very rare cases of blood infection, acidosis (acid in the blood), endocarditis (heart valve infection), and meningitis (swelling and irritation of the covering tissue around the brain) in patients who are already ill have been reported*.”* Parents who declined to participate said that they preferred to try formulas, herbal remedies, and probiotics that are advertised to be beneficial for babies with colic or “sensitive” intestines. To the authors, this indicated their uncertainty associated with clinical research and the informed consent process.

After recruitment, retention was about 75%, with dropouts related to infants not taking the formula or to the observation of loose stools. Based on our previous study of crying+fussing time in colicky infants[6], we determined that a sample size of 60 colicky infants (30 infants per study arm) would detect a mean difference of 100 min/d of crying + fussing time between the 2 study groups. We attempted to recruit this number over two years, but were unable to do so. Although we cannot be certain that enrollment and dropout rates would be the same for similar trials, we can estimate that a future probiotic formula trial conducted in the United States would require recruitment of > 80 enrollees for 60 infants to complete the trial. With our recruitment numbers in the 4th largest United States city, with 5300 pediatric gastroenterology visits annually, we suggest that a 3-center, 2-year trial would be optimal. Note that smaller differences in crying + fussing time were found to be significant in the Savino *et al*[5] and Szajewska *et al*[17] studies, but they required numbers of infants similar to the 30 in each group that we herein recommend.

**Safety:**We found no major side effects or safety problems contributing to product concerns or patient dropouts.

***Possible gut inflammation in colic***

We reported previously in infants with colic elevated fecal calprotectin (FC), suggesting a contribution of gut inflammation to this condition[18]. In the current study, we found that FC levels were high at virtually all time-points in the infants with colic, in both LGG+ and LGG- infants. The values of FC were consistently above the normal clinical range reported in adults (0-162 μg/g). An elevated FC in infants with colic and in normal infants at this age may reflect low-grade intestinal inflammation during this period of aggressive microbial colonization[19]. In addition, some of the infants were partially breast-fed, a condition arguably associated with a higher FC level[20], although there were only 3 such infants in each group. The difference in FC between groups was not significant. There could be a downward trend as a consequence of normal colonization, because FC levels in older children are substantially lower than those in infants[21]. Savino *et al*[22] recently published in abstract form a similar increase in fecal calprotectin in infants with colic.

While fecal calprotectin may be a helpful biomarker during the evolution of colic, plasma cytokines appear not to be informative. All of the cytokine levels in the infant plasma were detectable above the minimal detection level of the assay, and values were similar to those previously reported in other studies[23-25].

**Gas production in colic:**Our study provides evidence that excessive intestinal H2 gas production is less prevalent in our population than was reported previously[1] . Glucose breath testing and/or elevated fasting breath H2 in our study was abnormal in only 25% of our infants with colic. A previous study, in which infants with colic received the nonabsorbable sugar lactulose which raised breath H2 values, showed that most babies did not have any symptomatic response to lactulose[26]. It is possible that excessive gas could be a contributing factor to colic in a subset of infants.

***Microbial characterization of infant colic***

Pyrosequencing of 16S rRNA in the stool showed several interesting findings. First, longitudinally hyper-variable microbiota profiles during the 3-mo study (very evident in the twin pair shown in Figure 1B) support the concept of a “chaotic” pattern of colonization described by Palmer *et al*[27] early in life. This chaotic pattern has been challenged, with Koenig *et al*[28] suggesting a revised concept of population shifts attributable to major changes in life events. However in our study, patients were followed by telephone on a weekly basis for 3 mo. There were no major changes in life events, such as diet, diarrhea, antibiotic administration, or probiotic ingestion that were reported (Figure 1B).

One unexpected observation of our study was that inclusion of LGG in the formula had little impact on the overall composition or microbial diversity of the stool. This finding was particularly evident comparing the microbiota of the twin infants. The twins’ fecal microbial composition at 3 time-points fluctuated greatly, and there were Lactobacilli in both infants’ stools at visits 2 (day 14) to 4 (day 90), even though only one received LGG. Previous studies have shown that among adult monozyotic twins, the average microbiota similarity between twins is significantly higher than between unrelated subjects[29]. We suggest that factors other than *L. rhamnosus* were responsible for shifts in the fecal microbiota pattern of the infants. Such factors may include exposure to other people, animals, or environments.

In summary, our study showed that future trials of probiotics in formula for infants with colic at concentrations similar to those of Nutramigen with LGG should aim to recruit around 80 infants and should focus on determining efficacy on crying time at 14 d and on fecal calprotectin at 90 d. Changes in the microbiota (and/or their metabolic products) might be optimally observed at 2-4 wk of treatment.

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**COMMENTS**

***Background***

Colic has been defined as inconsolable crying plus fussing time equaling greater than 3 h daily for more than 3 d per week in infants from 3 wk to 3 mo of age. There have been many theories to explain the occurrence of colic, including bacterial overgrowth, the “fourth trimester” theory in which the baby would prefer to stay in utero, parental depression, excessive intestinal gas, and milk protein allergy. The probiotic Lactobacillus rhamnosus GG has been shown to reduce diarrheal volume and duration in children with acute infectious enteritis; it has also been shown to facilitate the development of more diverse fecal microbiota. Savino was the first to propose that an abnormal microbiota (“dysbiosis”) might be an important pathophysiological mechanism.

***Research frontiers***

Colic has begun to be studied as an example of possible dysbiosis with inflammation of the gut. Recent studies have centered on possible gut developmental issues, including the transition of environment from the intrauterine “bubble” to the external world and exposure to many new bacterial communities. Manipulation of the gut microbiota could be a major advance toward reducing the crying and fussing times of these infants.

***Innovations and breakthroughs***

This study is one of the first to investigate the impact on a probiotic on the microbiota of colicky babies and fecal calprotectin, as a marker of inflammation, during early life.

***Applications***

To summarize the practical applications of your research findings, so that readers may understand the perspectives by which this study will affect the field and future research. Future trials of probiotic-supplemented formulas may benefit from our pilot trial. The authors demonstrated robust numbers for crying+fussing time, fecal calprotectin, and microbial diversity in this population.

***Terminology***

*Lactobacillus rhamnosus* GG (LGG) is a health-promoting bacterial species which may be capable of reducing inflammation and regulating gut function. Microbiota is the term for bacterial community found in various locations such as skin and gut. Our focus is on the intestine, which has been shown to regulate diverse functions in the human body.

***Peer-review***

The topic is interesting and the manuscript well presented.

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**L-Editor: E-Editor:**

**Table 1 Comparison of baseline characteristics of randomized patients in the two study groups**

|  |  |  |  |
| --- | --- | --- | --- |
|  | **LGG+** **group** | **LGG-** **group** |  |
| Continuous variables | *n* | mean ±SD | *n* | mean ±SD | *P* |
| Age at the time randomized (d) | 9 | 57 ± 30 | 11 | 68 ± 28 | 0.341 |
| Gestational Age (wk) | 9 | 38 ± 2 | 11 | 37 ± 2 | 0.201 |
| Birth Weight (kg) | 9 | 3.1 ± 0.8  | 11 | 2.9 ± 1.1 | 0.471 |
| Birth Height (cm) | 7 | 51.6 ± 1.8 | 9 | 47.2 ± 5.3 | 0.041 |
| Discrete variables | *n* | mean ±SD | *n* | mean ±SD |  |
| Gender | 9 |  | 11 |  |  |
|  Female  |  | 4 ± 44.4 |  | 4 ± 36.4 | 1.0 |
|  Male |  | 5 ± 55.6 |  | 7 ± 63.6 |  |
| Race | 9 |  | 11 |  | 1.0 |
|  African-American |  | 3 ± 33.3  |  | 4 ±36.4 |  |
|  Caucasian |  | 6 ± 66.7 |  | 7 ± 63.6 |  |
| Ethnicity | 9 |  | 11 |  | 1.0 |
|  Hispanic or Latino |  | 2 ± 22.2 |  | 2 ± 18.2 |  |
|  Not Hispanic or Latino |  | 7 ± 77.8 |  | 9 ± 81.8 |  |
| Partial Breast Feed | 9 |  | 11 |  | 1.0 |
|  Yes |  | 3 ± 33.3 |  | 3 ± 27.3 |  |
|  No |  | 6 ± 66.7 |  | 8 ± 72.7 |  |
| Formula Type | 9 |  | 11 |  |  |
|  Earth’s Best |  | 1 ± 11.1 |  | 0 ± 0 |  |
|  Enfamil |  | 1 ± 11.1 |  | 0 ± 0 |  |
|  Enfamil Gentleease |  | 1 ± 11.1  |  | 0 ± 0 |  |
|  Gerber Good Start |  | 0 ± 0 |  | 1 ± 9.1 |  |
|  Isomil |  | 1 ± 11.1 |  | 1 ± 9.1 |  |
|  Neocate |  | 0 ± 0 |  | 1 ± 9.1 |  |
|  Nutramigen Ready Mix |  | 0 ± 0 |  | 1 ± 9.1 |  |
|  Similac |  | 2 ± 22.2  |  | 0 ± 0 |  |
|  Similac Advance |  | 1 ±11.1 |  | 0 ± 0 |  |
|  Similac Senstive |  | 0 ± 0 |  | 5 ± 45.5 |  |
|  Breast Milk |  | 2 ±22.2 |  | 2 ± 18.2  |  |

1Denotes p values obtained by non-parametric Wilcoxon rank sum test. All other *P* values are obtained by Fisher’s exact test. Q1: 1st quartile; Q3: 3rd quartile.

**Table 2 Longitudinal analysis of clinical variables by study group** (**LGG+** *vs* **LGG-) at baseline and follow up visits**

|  |  |  |
| --- | --- | --- |
|  | **Adjusted means (95%CI)** | ***P* value** |
| LGG+ group | LGG- group | Mean differences(95%CI) |
| Crying+fussing time (min)  |  |
| Visit 1 (Baseline) | 296 (210, 381) | 337 (251, 422) | 41 (-80, 161) | 0.51 |
| Visit 2 | 197 (117, 278) | 289 (142, 436) | 91 (-76, 259) | 0.29 |
| Visit 3 | 144 (54, 234) | 199 (69, 328) | 55 (-104, 213) | 0.50 |
| Visit 4 | 111 (65, 157) | 133 (60, 205) | 22 (-64, 107) | 0.62 |
|  |  |  |  |  |
| Fecal calprotectin (μg/g)  |  |
| Visit 1 (Baseline) | 285 (199, 371) | 294 (184, 404) | 9 (-131, 149) | 0.90 |
| Visit 2 | 226 (182, 270) | 305 (186, 423) | 79 (-48, 205) | 0.22 |
| Visit 3 | 229 (113, 345) | 250 (154, 347) | 21 (-130, 172) | 0.78 |
| Visit 4 | 211 (80, 342) | 332 (225, 440) | 121 (-48, 291) | 0.16 |

Longitudinal model: barr diary data = β0 + β1 × visit2 + β2 × visit3 + β3 × visit4 + β4  group + β5 × visit2 group + β6 × visit3 group + β7 × visit4 group. Here, visit2, visit3, visit4 are dummy variables; visit2 = 1 if at visit 2, 0 otherwise; visit3 = 1 if at visit 3, 0 otherwise; visit4 = 1 if at visit 4, 0 otherwise; group = 1 if in LGG group, 0 otherwise.**Table 3 Longitudinal analysis of cytokins by study group** (**LGG+** *vs* **LGG-) at baseline and follow up visits**

|  |  |  |
| --- | --- | --- |
|  | **Adjusted geometric means****(95%CI)** | ***P*** |
| **LGG+ group** | **LGG- group** |
| Cytokines |
| IFN-γ (pg/mL) |
|  Visit 1 (Baseline) | 0.7 (0.4, 1.3) | 0.6 (0.3, 1.2) | 0.84 |
|  Visit 2 | 1.2 (0.7, 2.0) | 0.5 (0.3, 0.9) | 0.04 |
|  Visit 4 | 0.4 (0.3, 0.6) | 0.2 (0.1, 0.4) | 0.07 |
|  |  |  |  |
| IL-10 (pg/mL) |
| Visit 1 (Baseline) | 3.7 (2.3, 6.0) | 4.2 (2.2, 8.1) | 0.78 |
| Visit 2 | 4.3 (3.2, 5.7) | 3.8 (2.3, 6.4) | 0.72 |
| Visit 4 | 2.7 (1.7, 4.2) | 2.0 (1.1, 3.7) | 0.47 |
|  |
| IL-12p70 (pg/mL) |
| Visit 1 (Baseline) | 0.2 (0.1, 0.4) | 0.2 (0.1, 0.4) | 0.85 |
| Visit 2 | 0.3 (0.2, 0.4) | 0.4 (0.3, 0.6) | 0.29 |
| Visit 4 | 0.2 (0.1, 0.5) | 0.3 (0.2, 0.4) | 0.57 |
|  |  |  |  |
| IL-1β (pg/mL) |  |  |  |
| Visit 1 (Baseline) | 0.08 (0.05, 0.13) | 0.19 (0.08, 0.43) | 0.09 |
| Visit 2 | 0.27 (0.08, 0.88) | 0.15 (0.06, 0.40) | 0.45 |
| Visit 4 | 0.11 (0.05, 0.21) | 0.16 (0.05, 0.49) | 0.52 |
|  |  |  |  |
| IL-6 (pg/mL) |  |  |  |
| Visit 1 (Baseline) | 0.3 (0.2, 0.5) | 0.4 (0.3, 0.6) | 0.36 |
| Visit 2 | 0.3 (0.2, 0.6) | 0.3 (0.2, 0.4) | 0.66 |
| Visit 4 | 0.2 (0.1, 0.4) | 0.2 (0.1, 0.3) | 0.63 |
|  |  |  |  |
| IL-8 (pg/mL) |  |  |  |
| Visit 1 (Baseline) | 6.5 (4.8, 8.7) | 7.0 (5.3, 9.4) | 0.69 |
| Visit 2 | 5.4 (4.1, 7.3) | 9.5 (6.6, 13.7) | 0.02 |
| Visit 4 | 4.9 (3.2, 7.5) | 4.6 (2.8, 7.8) | 0.88 |
|  |  |  |  |
| TNF-α (pg/mL) |  |  |  |
| Visit 1 (Baseline) | 4.5 (3.8, 5.3) | 4.8 (3.7, 6.2) | 0.69 |
| Visit 2 | 5.4 (4.4, 6.5) | 4.2 (3.2, 5.5) | 0.15 |
| Visit 4 | 4.4 (3.3, 5.9) | 5.7 (4.7, 6.9) | 0.14 |
|  |  |  |  |
| Treg (%) |  |  |  |
| Visit 1 (Baseline) | 7.2 (6.5, 8.0) | 6.7 (5.9, 7.5) | 0.37 |
| Visit 2 | 7.5 (6.8, 8.2) | 8.5 (7.0, 10.3) | 0.23 |
| Visit 4 | 7.5 (6.1, 9.3) | 9.2 (8.1, 10.5) | 0.10 |

Longitudinal model: ln(cytokines) = β0 + β1 × visit2 + β2 × visit4 + β3 × group + β4 × visit2 group + β5 × visit4 group; Here, visit2, visit4 are dummy variables; visit2 = 1 if at visit 2, 0 otherwise; visit4 = 1 if at visit 4, 0 otherwise; group = 1 if in LGG group, 0 otherwise. The geometric mean is the anti-log of arithmetic mean of log-transformed value. *P* values are obtained using wald test to test whether the differences of cytokines between study groups are significant at each visit.

**Figure 1 Participant flow diagram.**

Screened for eligibility (*n*=74)

**.**

**Follow-Up**

**Analysis**

**Follow-Up**

**Follow-Up**

**Allocation**

**Allocated to placebo (*n*=11)**

* Received allocated intervention (*n*=11)

**Measures:**

Barr Diary: 11 Stool: 11 Blood: 9

**Analyzed (*n*= 11)**

One patient was excluded from analysis due to missing baseline data: Barr diary.

**Visit 2 (*n*= 11)**

* Withdrew (*n*=2 ): Infant not taking formula well

**Measures:**

Barr Diary: 9 Stool: 8 Blood: 3

**Visit 4 (*n*= 7)**

* Withdrew from Study (*n*=0 )

**Measures:**

Barr Diary: 5 Stool: 6 Blood: 3

**Visit 3 (*n*=9)**

* Withdrew(*n*=2): Adverse Event Mild Diarrhea & Infant not taking formula

**Measures:**

Barr Diary: 5 Stool: 7

**Visit 3 (*n*= 9)**

* Withdrew: Infant not taking formula (*n*=1 )

**Measures:**

Barr Diary: 7 Stool: 8

**Visit 4 (*n*= 8)**

* Withdrew from Study (*n* =0 )

**Measures:**

Barr Diary: 8 Stool: 6 Blood: 5

**Visit 2 (*n*= 9)**

* Withdrew (*n* =0 )

**Measures:**

Barr Diary: 8 Stool: 8 Blood: 3

**Allocated to Formula with LGG (*n*= 9)**

* Received allocated intervention (*n*=9 )

**Measures:**

Barr Diary: 9 Stool: 6 Blood: 8

**Analyzed (*n*= 9)**

Two patients were excluded from analysis due to missing baseline data: Barr diary.

Excluded (*n*= 10)

* Protocol Change (*n*= 6 )
* Decided not to participate (n=4)

Excluded (*n*= 44)

* Screening “no show” (*n*= 10 )
* Refused to participate (*n*= 34 )

**Enrollment (*n*=30)**

Randomized (n=20)

**Consented (*n*=30)**

**Figure 2 Fecal microbiota of infants with colic before and after treatment with LGG *vs* placebo.** Average percent abundance of major bacterial groups at the genus level in colicky infants treated with LGG [LGG (+), *n* = 3] or placebo [LGG(-), *n* = 6]. Note that this was the subset of infants that had stools available for analysis at each of the 4 clinic visits. Visit 1 (day 1); visit 2 (day 14); visit 3 (day 42); and visit 4 (day 90).

**Figure 3 Abundance of *L. rhamnosus* and shifts in composition of stool specimens.** A: *L. rhamnosus* 16S rDNA sequence abundance in colicky infants at baseline and 14, 42 and 90 d after treatment with LGG (visits 2, 3 and 4). The median, 25th-75th percentiles (boxes) and 10th-90th percentiles (whiskers) are represented. Black dots represent outliers. a*P* < 0.05 compared to baseline and all other visits, respectively; B: Twins’ bacterial distribution at genus level. The abundance of major bacterial genera in stools of twin infants with colic: baseline (visit 1) and 14, 42 and 90 d after assignment to formula +/- LGG (visits 2, 3 and 4). Only one stool sample was evaluable at visit 1. The percent of abundance of Lactobacillus was indicated as orange box which was increased in LGG+ compared to LGG- at visit 2 (*P* < 0.05).

 **Figure 4 Shannon’s diversity of stool samples, measured over time.** LGG- infants’ stools are shown in blue and LGG+ infants’ stool results are shown in orange. Results are shown only for the patients for whom stools were available at each clinic visit. For two patients in the LGG+ group, one of the visits did not yield sequencing results. Note wide fluctuation at the various time points in infants in both groups.

