Human Milk Oligosaccharides Reduce Murine Group B Streptococcus Vaginal Colonization with Minimal Impact on the Vaginal Microbiota

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ABSTRACT

Group B Streptococcus (GBS) colonizes the vaginal mucosa of a significant percentage of healthy women and is a leading cause of neonatal bacterial infections. Currently, pregnant women are screened in the last month of pregnancy, and GBS-positive women are given antibiotics during parturition to prevent bacterial transmission to the neonate. Recently, human milk oligosaccharides (HMOs) isolated from breastmilk were found to inhibit GBS growth and biofilm formation in vitro, and women that make certain HMOs are less likely to be vaginally colonized with GBS. Using in vitro human vaginal epithelial cells and a murine vaginal colonization model, we tested the impact of HMO treatment on GBS burdens and the composition of the endogenous microbiota by 16S rRNA amplicon sequencing. HMO treatment reduced GBS vaginal burdens in vivo with minimal alterations to the vaginal microbiota. HMOs displayed potent inhibitory activity against GBS in vitro, but HMO pretreatment did not alter adherence of GBS or the probiotic Lactobacillus rhamnosus to human vaginal epithelial cells. In addition, disruption of a putative GBS glycosyltransferase (Δsan_0913) rendered the bacterium largely resistant to HMO inhibition in vitro and in vivo but did not compromise its adherence, colonization, or biofilm formation in the absence of HMOs. We conclude that HMOs are a promising therapeutic bioactive to limit GBS vaginal colonization with minimal impacts on the vaginal microenvironment.

IMPORTANCE

During pregnancy, GBS ascension into the uterus can cause fetal infection or preterm birth. In addition, GBS exposure during labor creates a risk of serious disease in the vulnerable newborn and mother postpartum. Current recommended prophylaxis consists of administering broad-spectrum antibiotics to GBS-positive mothers during labor. Although antibiotics have significantly reduced GBS neonatal disease, there are several unintended consequences, including altered neonatal gut bacteria and increased risk for other types of infection. Innovative preventions displaying more targeted antimicrobial activity, while leaving the maternal microbiota intact, are thus appealing. Using a mouse model, we found that human milk oligosaccharides (HMOs) reduce GBS burdens without perturbing the vaginal microbiota. We conclude that HMOs are a promising alternative to antibiotics to reduce GBS neonatal disease.

KEYWORDS

antimicrobial activity, group B Streptococcus, human milk oligosaccharides, vaginal colonization, vaginal microbiota

Group B Streptococcus (GBS or Streptococcus agalactiae) is a Gram-positive bacterium that colonizes the gastrointestinal and vaginal tracts of ~18% of pregnant women globally (1), exposing ~20 million infants to GBS at, or prior to, delivery (2). The majority
of children born to GBS-positive women themselves become colonized without symptoms (3); however, a subset of these infants (∼300,000 annually) develop invasive GBS infections, accounting for upwards of 100,000 infant deaths each year worldwide (2). In addition, 57,000 annual stillbirths are attributed to GBS infections (2), though this may be an underestimate since GBS is also the most frequently cultured bacterium in midgestation spontaneous abortions (4). Because maternal colonization is a risk factor for neonatal infections, universal screening in late pregnancy and intrapartum antibiotic prophylaxis (IAP) to GBS-positive or at-risk mothers is the current standard of care in many countries. These preventative measures have decreased, but not eradicated, GBS early-onset disease (5). However, this early antibiotic exposure disrupts the infant microbiota, and the potential adverse consequences of this perturbation are not fully established (6–10).

Breastfeeding has long been associated with improved infant health, reduced risk of infectious disease, and accelerated immune and microbial maturation within the gut (11–13). Human milk oligosaccharides (HMOs), the third most abundant component of breastmilk, are a group of structurally complex, unconjugated glycans that are recalcitrant to host digestive enzymes. HMOs provide nutritional advantage for beneficial microbes in the infant gut and drive immune maturation at the gut epithelium (13–16). Moreover, HMOs may protect against neonatal pathogens by acting as soluble “decoy” receptors for enteric pathogens (17, 18), through neutralization of bacterial toxins (19, 20), or via direct antimicrobial activity, including against GBS (21–24). Although the mechanism of HMO-mediated GBS inhibition is not known, GBS expression of a putative glycosyltransferase (locus san_0913) is necessary for inhibitory activity (21), and HMO exposure lowers GBS sensitivity to antibiotics, including vancomycin, erythromycin, and trimethoprim (21, 25, 26). Additional support for HMO-mediated anti-GBS activity stems from clinical observations that mothers who produce a functional variant of the fucosyltransferase enzyme FUT3, which attaches fucose in an α1-3 or α1-4 linkage to form certain HMOs, are less likely to be vaginally colonized by GBS (27).

We hypothesized that HMOs may reduce GBS vaginal colonization in vivo either through direct antimicrobial activity, or through indirect activity on the vaginal epithelium and/or vaginal microbiota. Here, we test this hypothesis using a murine model of GBS vaginal colonization and pooled HMOs (pHMOs) isolated from human breastmilk. We further assess the impact of pHMOs on bacterial attachment to human vaginal epithelial cells and phenotypically characterize a GBS strain that is resistant to HMO inhibition (21). Combined, our findings support the continued exploration of HMOs as a therapeutic strategy for GBS in pregnancy and the neonatal period.

RESULTS

Topical pHMO treatment reduces GBS vaginal burdens in vivo. To determine the effect of HMOs on GBS vaginal colonization in vivo, wild-type C57BL/6J female mice were vaginally inoculated with GBS COH1, a serotype III ST17 neonatal sepsis clinical isolate (28). Mice were treated with pHMOs (1 mg/dose) 2 h after GBS inoculation, and dosing was repeated on the following two consecutive days. Lacto-N-tetraose (LNT), a commercially produced HMO that inhibits GBS growth in vitro (21) was included as a treatment condition to test the efficacy of a single HMO. Vaginal swabs were collected prior to pHMO treatment on days 0, 1, and 2, as well as on days 3 and 6 postinoculation (Fig. 1A). Treatment with pHMOs significantly reduced GBS vaginal burdens on day 1 (P = 0.023) and 2 (P = 0.009) during active treatment, but these differences were resolved at day 3 and 6 after pHMO treatment ceased (Fig. 1B). No differences between LNT and mock-treated groups were observed at any time point. In addition, endogenous vaginal Enterococcus spp. were distinguished on the Streptococcus selective media, but no differences between treatment groups were detected (see Fig. S1 in the supplemental material).

Vaginal epithelial HMO exposure does not impact adherence of GBS or probiotic Lactobacillus. Because HMOs can reduce pathogen adherence (29–31) and promote adherence of beneficial bacteria to the host epithelium (32), we tested the impact of epithelial HMO pretreatment on adherence of GBS or the probiotic Lactobacillus rhamnosus GG to human vaginal epithelial (VK2) cells. We observed no effect of pHMO or LNT
pretreatment on GBS adherence to VK2 cells at two different concentrations (Fig. 1C), nor did HMO pretreatment alter *L. rhamnosus* adherence to VK2 cells (Fig. 1D).

**HMO resistance conferred by disruption of san_0913 does not alter GBS biofilm formation, adherence, susceptibility to antibiotics, or *in vivo* colonization in the absence of HMOs.** Although the exact mechanism of HMO anti-GBS activity has yet to be established, increased GBS sensitivity to intracellular targeting antibiotics and enhanced cell membrane permeability occur following HMO exposure (21, 25, 26). In addition, HMO exposure perturbs multiple GBS metabolic pathways including those related to linoleic acid, sphingolipid, glycerophospholipid, and pyrimidine metabolism (26). A transposon mutant library screen identified the *gbs0738* gene (locus *san_0913* or GBSCOH1_RS04065 in COH1), a putative glycosyltransferase family 8 protein, as essential for GBS susceptibility to HMOs over a 7-h time course (21); however, the functional role of this glycosyltransferase in GBS-host interactions and resistance to antimicrobial compounds has not been characterized. Using a targeted insertional mutant of *san_0913* (COH1 Δ*san_0913*) (21), we assessed the growth of wild-type (WT) COH1 and Δ*san_0913* in the presence of 0 to 20 mg/mL pHMOs over 18 h. We found that growth of COH1 was significantly inhibited at all pHMO concentrations tested compared to the mock control (Fig. 2A and B). Concentrations of 20 and 10 mg/mL pHMO inhibited growth of Δ*san_0913* but to a lesser degree than seen with COH1 (Fig. 2A and B). To determine whether *san_0913* disruption altered GBS characteristics associated with colonization, we assessed the...
ability of $\Delta san_{0913}$ to form biofilms and attach to vaginal epithelial cells. We observed no differences between COH1 and $\Delta san_{0913}$ biofilm formation in either bacteriologic (Todd-Hewitt broth [THB]) or eukaryotic (RPMI 1640) media, as measured by crystal violet staining (Fig. 2C). In addition, we observed no differences in VK2 adherence between the COH1 and $\Delta san_{0913}$ strains (Fig. 2D). In our in vivo model, we found no differences in vaginal GBS burdens between COH1 and $\Delta san_{0913}$ (Fig. 2E). However, when mice were treated with pHMOs as in Fig. 1A, $\Delta san_{0913}$ displayed significantly higher GBS burdens at day 1 postinoculation ($P = 0.007$) during active pHMO treatment, but this difference resolved at later time points (Fig. 2F). Furthermore, we performed MIC assays of a variety of antibiotic classes, hydrogen peroxide, and dimethyl sulfoxide (DMSO) to determine the impact of $\Delta san_{0913}$ deficiency on GBS susceptibility. MICs were determined by a 90% reduction in control optical density at 600 nm (OD600) values. No differences in MICs between COH1 and $\Delta san_{0913}$ were observed with any compound tested (see Table S1 in the supplemental material).

pHMO treatment minimally impacts the endogenous murine vaginal microbiota in the presence or absence of GBS. We previously identified that GBS introduction to the murine vaginal tract causes community instability, particularly a decrease in Staphylococcus succinus, a dominant vaginal microbe in C57BL/6J mice (33). Because HMOs are metabolized by a variety of bacteria in the neonatal intestines (34–38), and since maternal serum HMO levels correlate with specific taxa in the maternal urinary and vaginal microbiota (39), we investigated whether pHMO treatment impacted the murine vaginal microbiota in the presence or absence of GBS perturbation. Using swabs from the murine experiments outlined in Fig. 1A, 16S rRNA amplicon sequencing was used to characterize shifts in the vaginal microbiota of Control (mock-treated, mock-infected), pHMO (treated, mock-infected), Control_GBS (mock-treated, GBS-infected), and pHMO_GBS (treated, GBS-infected) mice. Alpha diversity, as measured by Shannon’s diversity index, significantly increased in Control_GBS and pHMO_GBS groups compared to controls (Fig. 3A). However, in the absence of GBS, alpha
diversity was not impacted in the pHMO versus control groups at any time point (Fig. 3A). As observed previously (33), mice that received GBS showed heightened community instability compared to mock-infected controls as measured by Bray-Curtis distance between time points. This effect was seen both in the presence (pHMO_GBS, \(P = 0.0048\)) and absence (Control_GBS, \(P = 0.0073\)) of pHMO treatment for pairwise comparisons between days 2 and 3 (Fig. 3B). No impact on community stability was observed with pHMO treatment in the absence of GBS (pHMO, Fig. 3B).

Across all four conditions, no significant differences were observed in community richness over the 6-day time course, as measured by observed operational taxonomic units (OTUs; see Fig. S2A). Mice exposed to GBS (Control_GBS and pHMO_GBS), regardless of treatment, experienced a significant drop in the relative abundance of \(S.\ succinus\) compared to Control mice starting at day 1, and this effect continued throughout the sampling period (Fig. 3C). No differences in the relative abundance of \(Enterococcus\) spp. or \(Lactobacillus\) spp., the two next most abundant endogenous OTUs, were observed between groups (see Fig. S2B and C). ANCOM analysis (40) identified \(Bacteroides\) as the only significant differentially abundant taxa across the four groups, with increased abundance in pHMO_GBS mice compared to all other groups (Fig. 3D).

**Murine vaginal community state types display minimal differential stability upon pHMO treatment in the presence or absence of GBS.** The human vaginal microbiome, and more recently the murine vaginal microbiome, are classified into community state types (CSTs) (41) and murine community state types (mCSTs), respectively (33). In humans, four CSTs are each dominated by different \(Lactobacillus\) species, and the remaining CST had a

FIG 3 Alpha and beta diversity and differential taxa abundance, as measured by 16S rRNA amplicon sequencing. Mice were mock infected or GBS infected and then treated with pHMOs or mock treated: Control (mock-treated, mock-infected), pHMO (treated, mock-infected), Control_GBS (mock-treated, GBS-infected), and pHMO_GBS (treated, GBS-infected) as described in Materials and Methods. (A) Shannon’s diversity index of vaginal 16S amplicon sequencing from each condition over the time course. (B) Bray-Curtis pairwise distances between subsequent time points. (C and D) Relative abundances of \(S.\ succinus\) (C) and \(Bacteroides\) spp. (D) according to treatment group over time. Results are displayed as a Tukey’s box plot (A, B, and D) and min-to-max box-and-whisker plots (C) \(n = 11\) to 21/group per time point). Data were analyzed by two-way repeated measures ANOVA with Tukey’s multiple-comparison test. All comparisons shown are versus the Control group. ***, \(P < 0.001\); **, \(P < 0.01\); *, \(P < 0.05\). All other comparisons are not significant.
non-\textit{Lactobacillus} dominant taxa or diverse array of facultative and strictly anaerobic bacteria (41). In C57BL/6J mice from Jackson Laboratory, the vaginal microbiome is separated into 5 mCSTs dominated by either \textit{S. succinus}, \textit{Enterococcus}, a \textit{S. succinus-Enterococcus} mixture, \textit{Lactobacillus}, or a mixture of different taxa (33). In the present study, we detected all five of these mCSTs by hierarchical clustering with Ward’s linkage of Euclidean distances to generate mCSTs (top bar). The treatment (middle bar) and time point (bottom bar) per sample are displayed above the heatmap. Highest to lowest taxonomic abundances are indicated by heatmap intensity corresponding to the color bar (indicated in lower right corner), ranging from dark purple to white.

To assess whether mice differentially transitioned between mCSTs among treatment groups, we tracked mCSTs in individual mice over time. As our prior study (33), we found that mCSTs were relative unstable, with 43% of uninfected and 87% of GBS-infected mice categorized to two or more mCSTs over the time course (Fig. 5). Using Bray-Curtis first distances for microbial communities within individual mice, we compared the instability between the baseline composition and the subsequent time points. Although there were no differences
FIG 5 Vaginal microbiome stability over time with pHMO treatment and/or GBS infection. mCST designations for mouse cohort samples are displayed ordered by treatment group and time point (left panels). For each mouse, (Continued on next page)
in longitudinal stability between Control and pHMO groups ($P = 0.9615$) or Control and Control_GBS groups ($P = 0.9999$), Bray-Curtis first distances were higher in Control_GBS versus pHMO ($P = 0.042$) and pHMO_GBS versus pHMO mice ($P = 0.0003$; Fig. 5).

Although mCST I (S. succinus-dominant) was the most common mCST in Control and pHMO groups, mCST II appeared with significantly more frequency in the Control group ($P = 0.0404$) and mCST I appeared with more frequency in the pHMO group ($P = 0.0067$) (Fig. 6A). No significant differences in mCST frequencies were observed between Control_GBS and pHMO_groups with mCST II, mCST IV, and mCST VI representing the most abundant mCSTs in both GBS-infected groups (Fig. 6A). As seen previously (33), mCST I was the most stable community state: combining all conditions and samples with successfully sequenced consecutive time points, 84/109 (77%) of mCST I samples were assigned mCST I at the next time point (self-transitioning). mCST VI (GBS-dominant) was the next most stable, followed by mCST II, mCST III, mCST V, and mCST IV (Fig. 6B). When separated by treatment groups, we found that mCST I was more likely to self-transition in the pHMO group compared to the Control group ($P = 0.0401$), whereas mCST II was more likely to self-transition in the Control group compared to the pHMO group ($P = 0.0031$) (Fig. 6C). In GBS-infected animals, no significant differences in mCST self-transitions were observed between Control_GBS and pHMO_GBS (Fig. 6C).

**DISCUSSION**

GBS remains a pervasive pathogen in pregnancy and the neonatal period. Current IAP prevention strategies have not fully abolished GBS neonatal infections and IAP is ineffective in preventing GBS infection prior to parturition. Because of the adverse effects of antibiotic exposure on the endogenous microbiota and propagation of antibiotic resistance, discovery of more targeted antimicrobial therapies to control maternal GBS carriage is important for maternal and neonatal health. Here, we apply HMOs, natural products produced by the mammary gland during pregnancy and lactation to in vitro and murine models of GBS vaginal colonization. HMOs are known for simultaneous prebiotic benefits on commensal bacteria (14, 34, 38) and antimicrobial activity toward pathogens, including GBS (21–24). To our knowledge, this is the first application of HMOs as a vaginal therapy in vivo. We propose that HMOs possess promising anti-GBS activity in this environment with minimal impact on the vaginal microbiota.

Our animal model demonstrated that pHMO treatment reduced GBS vaginal carriage, but this effect was only seen during active treatment with no sustained impact observed after

**FIG 6** Frequency and transitions of mCSTs across treatment groups. mCST designations for mouse cohort samples were combined from all time points. (A) Frequency of mCST appearances within treatment groups. (B) Proportion of samples designated to each mCST grouped by the mCST from the previous time point. A self-transitioning mCST would be designated from a mCST to the same mCST at the next time point (e.g., from mCST I to mCST I). (C) Relative proportions of mCSTs that self-transitioned at the next time point separated by treatment group. Data were analyzed by chi-square test.
treatment ceased (Fig. 1). This finding aligns with other murine models showing protective effects of HMOs in reducing pathogen colonization (31, 42–44). We observed no changes in bacterial adherence when VK2 cells were pretreated with pHMOs. This observation is distinct from work showing HMO-mediated inhibition of pathogens (31, 45–47) or enhanced attachment of beneficial bacteria (32, 48–50) at the gastrointestinal mucosa. Other studies have observed no impact of pHMO treatment on certain pathogens (51) or on pathogen colonization of other epithelial surfaces such as the bladder (52). These results suggest that prior mechanisms seen with HMOs and the gut epithelium may absent in the vaginal epithelium or with the bacterial species we tested.

There are several limitations to this HMO treatment model. First, we did not optimize dosage, timing, or length of pHMO treatment, nor did we assess the impact of HMO treatment or san_0913 deficiency across multiple GBS strain or serotypes. Pilot studies failed to show an effect of a 0.3-mg HMO dose using the same treatment regimen applied in Fig. 1A (data not shown); however, it is possible that dosages greater than 1 mg used here may provide more significant reduction of GBS burden or alterations to the vaginal microbiota. Second, although LNT shows potent in vitro anti-GBS activity (21), this did not translate to an in vivo GBS reduction, and thus the specific HMOs responsible for GBS reduction in our animal model are currently unknown. A clinical study found that Lewis positive women, who generate certain fucosylated HMOs, display reduced GBS vaginal carriage and infant colonization at birth (27). Specifically, levels of lacto-N-difucohexaose I (LNDFHI) in breastmilk samples negatively correlated with maternal GBS colonization status and reduced GBS growth in vitro (27). Successful identification of the specific HMO(s) providing activity in vivo is a critical future step to translate our findings to the clinic. Third, HMOs and their fermentation products have multiple known gastrointestinal epithelial and immune modulatory activities (53–56). Likewise, it is possible that HMOs act indirectly through altering host vaginal responses to GBS; however, this was not evaluated in our study. Lastly, using murine models to test whether HMOs possess potential therapeutic activity in preventing GBS neonatal transmission and adverse birth outcomes (57, 58) will be an important application of our findings.

Although the exact mechanism of anti-GBS activity by HMOs is unknown, GBS susceptibility is linked to expression of a GBS-specific putative glycosyltransferase (locus san_0913) thought to catalyze the addition of glucose or galactose residues to the cell surface and thus may enable incorporation of HMOs into the GBS cell wall (21). In prior work, a glycosyltransferase-deficient Δsan_0913 strain showed resistance to HMO inhibition (5 mg/mL) over 7 h of culture (21). In our growth analysis, we confirmed this finding extended out to 18 h (Fig. 2). At higher concentrations (10 to 20 mg/mL) matching physiologic concentration of HMOs in human colostrum and breastmilk (59, 60), Δsan_0913 growth was inhibited, but not to the same extent as COH1, suggesting that this deficiency does not completely resolve anti-GBS activity of HMOs. Recent work has shown that HMOs induced multiple GBS stress responses related to cell membrane and cell wall components (26), but the role of san_0913 in this GBS response has not been established. While streptococcal glycosyltransferase activity has been implicated in biofilm formation and composition in S. mutans (61), our phenotypic analyses did not reveal any substantial deficits in the glycosyltransferase-deficient Δsan_0913 in biofilm formation, vaginal cell adherence, or in vivo vaginal colonization in the absence of HMO treatment. In the presence of HMOs, san_0913 deficiency conferred a colonization advantage early on during colonization, but not at later time points, suggesting that direct inhibition by HMOs at least partially contributes to HMO-mediated GBS reduction in vivo. These results may have important clinical implications for HMO therapies and emergence of spontaneous HMO-resistant GBS under selective pressure.

HMOs serve as prebiotics in the gut by promoting the establishment of Bifidobacteria and Bacteroides (37, 38, 62). Mammary HMO production begins early in pregnancy and is detected in maternal circulation in the first trimester (63). Moreover, maternal serum levels of two abundant HMOs (2′-FL and 3′-SL) positively correlate with vaginal Gardnerella spp. and L. crispatus, respectively (39), providing a basis for the hypothesis that HMOs might not only shape neonatal microbiota and immunity but also maternal vaginal microbiota. Whether HMOs have the potential to directly impact the vaginal microbiome in humans...
has not been determined, however, a common vaginal species, *L. gasseri*, lacks the ability to metabolize HMOs (34). Because of the well-known prebiotic effects of HMOs on the infant microbiota, we examined the impact of pHMOs on the murine vaginal microbiota in our colonization model. We found minimal pHMO-driven changes to the community composition in terms of alpha and beta diversity (Fig. 3). The most marked difference between groups in our model was the emergence of *Bacteroides* in mice dually inoculated with GBS and treated with pHMOs (Fig. 3D). While the relative abundance of *Bacteroides* remained below 5% of the entire microbial landscape in the majority of mice, 0.1 to 5% abundance is estimated to account for ca. $10^4$ to $10^6$ total CFU in the murine vaginal tract. In women, the vaginal microbiota postpartum shows community instability and increases in *Bifidobacterium* and *Bacteroides* (64, 65), but the mechanisms driving these changes are unknown. Whether HMOs can be detected in the human vagina during pregnancy and lactation, and whether human vaginal microbes can metabolize HMOs are important topics of future study.

There are several limitations to the interpretation of our murine microbiome data. First and foremost, the murine vaginal microbiome does not fully reflect the human vaginal microbiome in terms of species present; although there is an mCST dominated by a murine *Lactobacillus* (see Fig. S2), it is a rare community in C57BL/6J mice (33). As a future direction, we seek to use humanized microbiota mice to assess pHMO-mediated changes to the vaginal microbiota in the presence of human vaginal bacteria, such as that done in mouse models colonized with human gastrointestinal microbiota and treated with HMOs (42, 66). In women, GBS is present at a low relative abundance in the vagina (67), whereas in our mouse model GBS becomes a dominant member of vaginal community in some mice upon introduction (Fig. 4). This high relative abundance may alter dynamics of GBS and other vaginal taxa distinct from human vaginal communities. In addition, the length of HMO treatment may need to be extended to observe larger effects. Prior studies have described more pronounced HMO-mediated shifts to the gut microbiota of both conventional (44, 68) and humanized microbiota mice (66); however, the length of HMO treatment in these studies was longer than in our model (3 to 8 weeks versus 3 days, respectively).

By combining our prior (33) and current studies, we found that the vaginal microbiome of the C57BL/6J mice from one vendor is highly consistent across cohorts over several years. In both studies, we found that GBS introduction increases vaginal community instability and reduces the relative abundance of the most abundant taxa *S. succinus*. In addition, we confirmed our prior observation that mCST I (*S. succinus* dominant) is the most stable murine community over time. These consistencies highlight the utility of this murine model in comparing different experimental groups across cohorts and experimental variables.

In summary, we have demonstrated HMOs can reduce GBS vaginal colonization in an animal model with minimal impacts on the vaginal microbiota. There is mounting evidence that HMOs play an important role in shaping the infant gut microbiota and preventing pathogen colonization. HMO introduction to the vaginal tract may provide similar beneficial effects. These findings lay the framework to expand our knowledge of therapeutic applications of HMOs and support their continued development as a target for controlling GBS colonization in women.

**MATERIALS AND METHODS**

**Reagents, bacterial strains, and cell lines.** Pooled HMOs were isolated from human milk samples collected through a donation program at UC San Diego, lyophilized, and stored at $-20^\circ$C as described previously (69). Individual HMO lacto-N-tetraose (LNT) was purchased from Dextra Laboratories. Prior to use, HMOs were resuspended in molecular grade water to a final concentration of 100 mg/mL, and subsequent dilutions were made in cell culture media (in vitro) or molecular grade water (in vivo).

Group B Streptococcus (GBS) strains COH1 (ATCC BAA-1176) and isogenic Δssn, 0913 generated previously (21) were grown for at least 16 h at 37°C in Todd-Hewitt Broth (THB) prior to experiments with 5 μg/mL erythromycin added to Δssn, 0913 cultures. Prior to in vitro and in vivo experiments, overnight cultures were diluted 1:10 in fresh THB and incubated stationary at 37°C until mid-log phase (OD$_{600}$ = 0.4). *Lactobacillus rhamnosus* GG (ATCC 53103) was grown for 16 h at 37°C without shaking in de Man, Rogosa, and Sharpe (MRS) broth.

Immortalized human vaginal epithelial cells (VK2/E6E7, ATCC CRL-2616) were cultured in keratinocyte serum-free medium (KSF-M) (Gibco) with 0.5 ng/mL human recombinant epidermal growth factor and 0.05 mg/mL bovine pituitary extract. Cells were cultured in a 37°C incubator with 5% CO$_2$. Cells were split every 3 to 4 days at $\sim$80% confluence, and 0.25% trypsin–2.2 mM EDTA (Coming) was used to detach cells for passageing.
GBS growth kinetics. For growth curves, log phase cultures were diluted 1:10 in RPMI 1640 (Gibco) in 96-well microtiter plates with 20, 10, 5, or 2.5 mg/ml pHMOs or water control in a 200-μL total volume. Wells with pHMOs and media only were included to confirm absence of microbial contamination. Plates were incubated at 37°C, and the absorbance at OD₅₉₀ was read every 15 min for 18 h using a BioTek Cytation 5 multimode plate reader.

Biofilm assays. GBS biofilm assays were performed as described previously (70). Briefly, overnight cultures were diluted to an OD₅₉₀ of 0.1 in RPMI 1640 or THB, followed by incubation at 37°C for 24 h. The medium was removed, and the biofilms were washed twice with phosphate-buffered saline (PBS) before drying at 55°C for 30 min. Biofilms were stained with 0.2% crystal violet for 30 min, washed with PBS three times, and destained with 80:20 ethanol-acetone. Supernatant was transferred to a fresh 96-well plate, and the absorbance was measured at OD₅₉₀ using a BioTek Cytation 5 plate reader. Values were normalized to total bacterial growth prior to washing and staining, and data were expressed as a ratio of crystal violet staining to total bacterial growth (OD₅₉₀ to OD₆₀₀).

MIC assays. MICs were performed as described with minor adaptations (71). For within-mouse assessment of instability and mCST transitioning, samples with only one time point collected were excluded from Bray-Curtis analysis. The optimum number of clusters (5 mCSTs) was determined using wss and silhouette metrics generated through the R package stats (79) on the rarefied feature table with Ward’s linkage.

Adherence assays. GBS adherence assays were performed on confluent VK2 cells in 24-well plates as described previously (72, 73). For studies using HMOs, medium was replaced with KSFM containing 3 or 6 mg/mL pHMOs, LNT, or vehicle control for 18 h. Cells were infected with GBS COH1, Δsram_0093, or L. rhamnosus at MOI = 1 (assuming 1 × 10⁷ VK2 cells per well). Bacteria were brought into contact with the VK2 cells by centrifugation for 1 min at 300 g. After 30 min, the supernatant was removed, and the cells were washed six times with sterile PBS. Cell layers were incubated for 5 min with 100 μL of 0.25% trypsin-0.2 mM EDTA, and then 400 μL of 0.025% Triton-X in PBS was added. Wells were mixed 30 times to ensure detachment, and bacterial recovery was determined by plating on THB or MRS agar plates using serial dilution and counting CFU. Data are expressed as the percentage of adherent CFU compared to the original inoculum.

Animals. Animal experiments were approved by the UC San Diego Bay and approved by the College of Medicine Institutional Animal Care and Use Committees (IACUC) and conducted under accepted veterinary standards. Mice were allowed to eat and drink ad libitum. WT C57BL/6J female mice, originally purchased from Jackson Laboratories, aged 7 weeks, were allowed to acclimate for 1 week prior to experiments.

Murine GBS vaginal colonization model. Vaginal colonization studies were conducted as described previously (74). Briefly, mice were synchronized with 0.5 mg of β-estradiol administered intraperitoneally 24 h prior to inoculation with 1 × 10⁹ CFU total of GBS COH1 or PBS as a mock control into the vaginal tract. Where applicable, mice were administered 1 mg (10 μL of 0.025% Triton-X in PBS) pHMOs, LNT, or vehicle control into the vaginal lumen at 2 h postinoculation. Vaginal swabs were collected daily, about 24 h apart. In experiments testing HMOs, mouse received additional HMO or mock treatments on days 1 and 2 immediately after swab collection. Recovered GBS (identified as pink/mauve colonies) was quantified by plating on CHROMagar StrepB (DRG International, Inc.). Growth of blue colonies was considered endogenous Enterococcus spp. based on manufacturer protocols. Remaining swab samples were stored at −20°C until further use.

Sample processing and 16S rRNA amplicon sequencing. DNA was extracted from thawed bacterial suspensions using a Quick-DNA Fungal/Bacterial Microbe kit protocol (Zymo Research). The V4 regions of the 16S RNA gene were amplified using barcoded 315F-806R primers (75), and the resulting V4 amplicons were sequenced on an Illumina MiSeq. Raw sequencing data were transferred to Qiita (76). Sequences were demultiplexed, trimmed to 150-bp reads, and denoised using Deblur through QIIME2 v2020.8. QIIME2 was also used for rarefaction (1,900 sequences per sample) and calculation of the alpha diversity (Shannon and OTUs) and beta diversity (Bray-Curtis distance). For ANCOM (40) analysis for differentially abundant OTUs, the nonrarefied feature table was used. Taxonomic assignments used the naive Bayes sklearn classifier in QIME 2 trained on the 515F/806R region of Greengenes 13,8 99% OTUs. Since many of the samples were low biomass, DNA from sequencing reactions and kits had a substantial impact on the data set. Negative controls that went through the entire pipeline, from DNA extraction to sequencing, were used to catalog these contaminants (Pseudomonas veronii). Mitochondria and chloroplast 16S sequences were also removed. Output files generated through the Qiime2 pipeline were exported and analyzed with R version 3.6.1 (2019-07-05; “Action of the Toes”) using stats, factoextra, and phyloseq (78, 79). Data visualization was performed with ggplot2 (80) and Seaborn (81).

Community state type delineation. Feature tables and representative sequences generated from three independent studies were merged and used to generate a taxonomy file. Two more studies from our prior work (33) were downloaded from EBI accession number PRJEB25733 in addition to the present study (EBI accession PRJEB49304) for Fig. S2 depicting the Baseline CSTs. To assign mCSTs and create heatmaps, hierarchical clustering was performed using the R package stats (79) on the rarefied feature table with Ward’s linkage of Euclidean distances. The optimum number of clusters (5 mCSTs) was determined using wss and silhouette (kmeans) based on the dendrogram. For EBI accession number PRJEB49304 (this study) alone, including all experimental conditions and time points, we added an additional GBS-dominant mCST, as modeled previously (33). For within-mouse assessment of instability and mCST transitioning, samples with only one time point collected were excluded. Samples that did not successfully sequence at the baseline (day 0) time point were excluded from Bray-Curtis first distances analysis.

Statistics. All data were collected from at least three biological replicates performed in at least technical duplicate as part of at least two independent experiments. When biological replicates were not available (e.g., immortalized cell lines and bacteria only assays), experiments were performed independently at least three times. Mean values from technical replicates were used for statistical analyses, with independent experiment values or biological replicates represented in graphs with means, medians
with interquartile ranges, or box-and-whisker plots with Tukey’s, as indicated in the figure legends. All data sets were subjected to the D’Agostino and Pearson normality test to determine whether values displayed Gaussian distribution before selecting the appropriate parametric or nonparametric analyses. In instances where in vitro and in vivo experimental n values were too small to determine normality, the data were assumed to be nonparametric. GBS vaginal colonization burdens were assessed using Kruskal-Wallis with Dunn’s multiple-comparison test or two-stage Mann-Whitney test, as indicated in figure legends. GBS adherence to VK2 cells was assessed by or two-way analysis of variance (ANOVA) with Dunnett’s multiple-comparison test or the Wilcoxon matched-pairs signed rank test, as indicated in figure legends. GBS growth (area under curve) and biofilm formation were compared using two-way repeated measures ANOVA with Dunnett’s multiple-comparison test and two-way ANOVA with Sidak’s multiple-comparison test, respectively. Data from 16S rRNA amplicon sequencing were analyzed by a Bray-Curtis first distances were analyzed by using a Kruskal-Wallis test with a Dunn’s multiple-comparison posttest. mCST transition frequencies were compared by using a chi-square test. Statistical analyses were performed using GraphPad Prism, version 9.2.0 (GraphPad Software, Inc., La Jolla, CA). P values of <0.05 were considered statistically significant.

**Data availability.** Sequencing Data used in this study is available in EBI under accession number PRJEB49304; the code is accessible at GitHub under project “HMO_GBCategorization.”

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, TIF file, 0.2 MB.

**FIG S2**, TIF file, 0.2 MB.

**FIG S3**, TIF file, 1.4 MB.

**TABLE S1**, DOCX file, 0.01 MB.

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K.A.P., L.B., and V.N. conceived and designed experiments. K.A.P., M.E.M., S.O., A.V., P.B., J.J.Z., and D.M. performed experiments. K.A.P., M.E.M., and A.V. analyzed and interpreted results. M.E.M. and K.A.P. drafted the manuscript. All authors contributed to the discussion/manuscript edits.

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