ADDENDUM

Compositional analyses reveal correlations between taxon-level gut bacterial abundance and peripheral T cell marker expression in African infants

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ABSTRACT

Although exclusive breastfeeding has been linked to lower rates of postnatal HIV transmission compared to nonexclusive breastfeeding, mechanisms underlying this are unclear. Across a longitudinally sampled cohort of South African infants, we showed that exclusively breastfed (EBF) infants had altered gut bacterial communities when compared to nonexclusively breastfed (NEBF) infants, as well as reduced peripheral CD4+ T cell activation and lowered chemokine and chemokine receptor expression in the oral mucosa. We further demonstrated that the relative abundance of key taxa was correlated with peripheral CD4+ T cell activation. Here, we supplement those findings by using compositional data analyses to identify shifts in the abundance of several Bifidobacteria strains relative to select strains of Escherichia, Bacteroides, and others that are associated with the transition to NEBF. We illustrate that the abundance ratio of these taxa is tightly correlated with feeding modality and is a strong predictor of peripheral T cell activation. More broadly, we discuss our study in the context of novel developments and explore future directions for the field.

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Introduction

More so than by any other mechanism, gut bacterial communities are shaped by dietary practices.1,2 In infants, early life feeding practices are integral in shaping bacterial primary succession patterns in the gut and associated long-term health outcomes.3–5 For example, exclusive breastfeeding (EBF) during the first six months of life has been associated with protection against diarrhea, diabetes, and other morbidities, as compared to nonexclusive breastfeeding (NEBF), where infants are fed breastmilk with a combination of other liquids and solids, or formula feeding (FF).6–8 Recently, studies have begun to elucidate the link between infant feeding practices, microbial community composition, and infant health,9–11 revealing alterations in gut bacterial community composition and mucosal gene expression between feeding practices.5,10–13

Despite the benefits conferred by exclusive breastfeeding, it is not always a viable option due to maternal disease, dietary restrictions, or availability. In our recent study, we demonstrated that nonexclusive breastfeeding alters the community composition of gut bacteria, increases T cell activation and, possibly, mucosal recruitment of HIV target cells during at least the first 3 months of life. Here, we explore the results of our study in the current context of the field, apply novel compositional data analytical approaches to reveal models between bacterial abundance and measures of immune activation, and discuss next steps and remaining questions.

Feeding practices and HIV transmission

Despite the associated benefits, breastfeeding represents a conduit for HIV transmission in the absence of antiretroviral prophylaxis.14 Though the administration of antiretroviral therapy (ART) has greatly reduced the incidence of mother-to-child transmission, hundreds of thousands of infants still acquire HIV via this mechanism.15 However, previous
studies have found that EBF infants have a lower risk of HIV acquisition compared to NEBF infants, putatively due to reduced cell-free viral loads in the breastmilk of mothers that exclusively breastfeed. In our study conducted in Cape Town, South Africa, we showed that the abundance of specific bacterial taxa was correlated with decreased peripheral CD4+ T cell activation in EBF infants. Further, we demonstrated that gene expression of chemokines and associated receptors involved in HIV target cell recruitment was reduced in the oral mucosa of EBF infants relative to NEBF infants.

**Community composition of gut bacterial microbiota is altered between EBF and NEBF infants**

Many datasets exist about the effects of EBF vs FF on infant gut microbiota, though fewer have surveyed bacterial communities between EBF and NEBF infants. In our 2018 study, we demonstrated that EBF infants possessed altered community composition and diversity of gut bacteria through the first 14 weeks of life, even compared to infants fed breast milk, but along with other solids or liquids (NEBF). We reported consistently decreased phylogenetic diversity of EBF bacterial communities at both 6 and 14 weeks of life. Furthermore, we found that overall community profiles significantly differed between EBF and NEBF infants at both time points, using phylogenetically aware (weighted Unifrac) and traditional ecological (Jensen-Shannon) distance metrics. Specifically, we found reduced relative abundance of *Streptococcus luteciae* in EBF infants at 6 weeks of life compared to NEBF infants. At 14 weeks, we found that EBF infants displayed increased relative abundance of *Actinomyces* and *Atopobium* taxa and decreased relative abundance of two members within *Bacteroides* compared to NEBF infants. These results were recently corroborated by a meta-analysis of the effects of exclusive breastfeeding on gut microbiota across studies and populations, consistently finding reduced a-diversity and decreased relative abundance of *Bacteroides* in EBF infants.

Here, we supplement the findings in our original study by applying compositional analyses of bacterial abundance. With the application of compositional data analytical tools, we can further dissect differences in bacterial abundance between EBF and NEBF infants while reducing compositional effects (artifacts associated with relative abundance datasets). Microbial community datasets generated from marker gene surveys (e.g., via sequencing 16S/18S rRNA genes) yield abundance measurements that are influenced by the composition of the community, precluding the inference of true shifts in bacterial abundance. Here, we use a special application of the isometric log ratio transformation, a compositional data transformation, to generate ratios of highly informative bacterial taxa that enable more accurate inference of shifts in bacterial abundance. These ratios, termed balances, enable interpretable taxon-level inference of alterations that best discriminate EBF from NEBF infants (Figure 1). To identify the taxa used in Balance 1, we performed a centered log ratio (CLR) transformation on our bacterial abundance measurements. Once transformed, we performed a sparse principal component analysis to identify taxa positively and negatively associated with each sparse principal component and used these taxa to build a series of orthonormal compositional balances. Penalized (ℓ1) logistic regression was then used to identify compositional balances strongly associated with feeding practice. The ℓ1 penalty (lasso) is useful for selecting predictor variables strongly associated with specific responses. The ratio identified in Balance 1 yielded the greatest discrimination between feeding modalities.

Our results demonstrate that the log ratio abundance of three Bifidobacterial taxa (*B. breve*, *B. gallicum*, *B. bifidum*) to *Escherichia*, *Bacteroides dorei*, *Veillonella dispar*, *Bacteroides vulgatus*, and *Ruminococcus gnavus* is significantly reduced (*P* < .027) in NEBF infants, with non-Bifidobacterial taxa surpassing Bifidobacterial abundance by 6 weeks. Specifically, at birth, all infants in our dataset possessed a median value of 1.12 for Balance 1 and this value remained relatively stable for EBF infants during our study (0.7 at 6 weeks; 2.0 at 14 weeks). Conversely, the median value for NEBF infants fell to -0.16 at 6 weeks and -1.14 at 14 weeks, indicating greater abundance of taxa in the denominator (*Escherichia et al.*) than in the numerator (*B. breve*, *B. gallicum*, *B. bifidum*) to *Escherichia*, *Bacteroides dorei*, *Veillonella dispar*, *Bacteroides vulgatus*, and *Ruminococcus gnavus*. This decrease in Bifidobacteria in NEBF infants suggests that non-Bifidobacterial taxa are more abundant in this feeding modality.
B. bifidum. Bifidobacteria are canonically beneficial inhabitants of the breastfed infant gut microbiota, with select strains possessing the ability to metabolize human milk oligosaccharides. Conversely, B. dorei and B. vulgatus have been implicated in negative health outcomes such as the development of autoimmune disease and necrotizing enterocolitis in neonates and infants, and increases in R. gnavus have been associated with intestinal dysbiosis.

Alterations in the infant immune response associated with feeding modality

In our 2018 study, we assayed peripheral CD4+ T cell activation and buccal mucosal chemokine, chemokine receptor, and keratin gene expression levels longitudinally between EBF and NEBF infants, in order to gain insight into the mechanisms underlying the increased risk of HIV acquisition in NEBF infants. We found that peripheral CD4+ T cell activation was consistently decreased in EBF infants across all time points, with significant reductions in the frequency of CD4+ human leukocyte antigen-DR+ (HLA-DR+) and CD4+ HLA-DR+CD25+ T cells in EBF infants at 6 weeks of life. This relative immune quiescence in EBF infants may confer protection from HIV transmission, in line with previous findings reporting that systemic immune quiescence is associated with protection from sexual HIV transmission. Because HIV preferentially infects activated CD4+ T cells, our findings began to provide mechanistic insight into the underlying causes of reduced HIV transmission in EBF infants.

In the primary analysis, we further assayed chemokine, chemokine receptor, and keratin gene expression in the oral mucosa of EBF and NEBF infants, finding reduced levels of several of these in EBF infants at weeks 6 and 14. Specifically, we found significant reductions (measured as log_{10} fold changes in expression) in CCL5, CCL22, CXCR7, IL7R, and KRT5 in EBF infants, relative to NEBF counterparts. We hypothesized that these decreases in chemokine and chemokine receptor gene expression levels in EBF infants indicated lower immune activation in buccal mucosa, and a consequent...
decrease in HIV susceptibility. In line with previous work, our results of elevated cytoskeletal gene expression (KRT5) are suggestive of epithelial barrier compromise as a putative mechanism for increased mucosal HIV acquisition risk with NEBF. Together, these results suggest that activation of HIV target cells in conjunction with compromised mucosal barrier integrity may facilitate HIV virion access to target cells, consistent with observations from previous studies of genital mucosa.\textsuperscript{36–38}

The abundance of key bacterial taxa predicts mucosal immune activation and correlates with feeding practices

Our findings led us to propose that the increased bacterial diversity associated with NEBF may induce immune activation, thus explaining the higher incidence of HIV transmission in NEBF infants as compared to EBF infants. Indeed, it has been previously demonstrated that EBF infants have altered gene expression and peripheral T cell activation than formula fed infants.\textsuperscript{12,39} In our parent study, we identified correlations between ecological distance metrics (Jensen-Shannon and weighted Unifrac) and the expression of several T cell markers, suggesting that bacterial community composition shifted with T cell marker abundance. Using compositional data analytical approaches, we can move beyond correlations between community-wide distance metrics and immune activation markers and into more targeted models involving specific microbial taxa. Here, we use compositional analyses to enable modeling of interactions between bacteria in Balance 1 (Figure 1) and the expression of T cell activation markers (Figure 2). By reducing compositional effects associated with marker gene sequencing, we can more accurately infer interactions between bacterial abundance and external covariates (T cell marker expression).

Here, we demonstrate that the isometric log ratio of bacteria identified in Balance 1 as associated with feeding practice is a significant negative predictor of immune activation and co-receptor expression (CD25+: $P < .0049$; CCR5+: CD25+: $P < .007$; HLA-DR+: CD25+: $P < .002$). Further, this interaction is associated with feeding modality ($P < .027$) and is correlated with an increase in the ratio of several \textit{Bifidobacteria} taxa relative to \textit{Escherichia} and others (Figure 2). Broadly, these findings suggest that, in addition to discriminating feeding practices, Balance 1 also defines a spectrum of immune activation where samples with positive values (greater abundance of taxa in the numerator) display relative immune quiescence and samples with negative values (greater abundance of taxa in the denominator) display relative immune activation, with several plausible mechanisms in play. Concerning immune quiescence, \textit{Bifidobacteria} have been shown to reduce serum cytokine levels in a diverse range of disease conditions in adults and peripheral blood mononuclear cells,\textsuperscript{41} suggesting beneficial immunomodulatory effects. In adults, supplementation of \textit{B. infantis} has been shown to reduce cytokine levels in various inflammatory disorders.\textsuperscript{41} Conversely, several taxa in the denominator of Balance 1 have been associated with elevated immune activation. Increased fecal abundance of \textit{R. gnavus} has been associated with the development of allergies in infants, and work in murine models has linked \textit{R. gnavus} to elevated cytokine secretion and airway inflammation.\textsuperscript{34} Vatanen et al. demonstrated the immunostimulatory nature (upon initial exposure) of \textit{E. coli} lipopolysaccharide (LPS) compared to that of other gut commensals which are immunoinhibitory.\textsuperscript{32} Early exposure to immunostimulatory LPS produced by \textit{E. coli} has been shown to elicit significantly increased cytokine production in human cells in vitro as compared to LPS produced by other gut associates.\textsuperscript{32} In EBF infants, the reduced activation of these markers may be linked to reduced abundance of \textit{E. coli} LPS and a reduction in proinflammatory bacteria, and/or elevated abundance of immunomodulatory \textit{Bifidobacteria}, or a synthesis of both. With respect to HIV acquisition, the infant gut is known to host an abundance of CD4+ CCR5 + T cells which are highly susceptible to HIV-1.\textsuperscript{42} Our finding of a significant negative correlation between activated, CCR5+ expressing CD4 T cells and Balance 1 suggests that elevated abundance of \textit{Bifidobacteria} relative to \textit{Escherichia, Bacteroides}, and other inflammatory taxa may contribute to immune quiescence and reduced HIV target cell abundance, though this should be validated in additional cohorts and/or experimental systems. If these results are found to be reproducible, \textit{Bifidobacteria} probiotics may represent an attractive intervention to restore the health of NEBF infants and reduce HIV transmission.
Next steps and open questions

Despite the advances in our ability to quantitate and analyze bacterial and immunological dynamics in parallel, many questions remain regarding the effects of feeding practices on infant gut microbiota and immune development. That is, most of our insights into interactions between the microbiome and external variables are correlative in nature. As we progress toward causal studies of the microbiome using germ-free animal models, insights from large observational studies will translate into mechanistic insights. In our 2018 study, we demonstrated altered bacterial community compositions correlated with various measurements of immune activation. Here, we further interrogated those interactions by fitting models between specific bacterial taxa and feeding mode (Figure 1) and immune marker expression (Figure 2). However, whether altered bacterial communities elicited immune activation themselves, or if other variables between feeding practices induced immune activation, which in turn altered the associated bacterial communities, remains unknown. Our original study and further work here reveal that a link exists between these variables, but the directionality of these interactions remains an open question. Further efforts to interrogate these interactions in animal models may enable the fine-scale resolution needed to pinpoint mechanistic interactions and infer directionality between specific bacterial strains and immune activation. Validation of these models in additional cohorts will be imperative to determine...
the generality of these interactions between gut bacterial associates and immune responses.

As with any bacterial microbiome study, these data represent one piece of an incredibly elaborate network of interactions between bacteria, fungi, viruses, and the immune response. It is likely that these other microbial communities, as well as various environmental factors, act as determinants of gut bacterial succession and community dynamics. Our collective analyses on this cohort have begun to elucidate mechanistic links between the reduction in HIV susceptibility associated with exclusive breastfeeding versus nonexclusive breastfeeding in African infants. With this establishment, future studies will help to mitigate the risks associated with nonexclusive breastfeeding for all infants, regardless of HIV-exposure or geographic location. The bacteria included in models illustrated here (Balance 1) may guide targets for therapeutic interventions to mitigate the negative effects of NEBF. Our results, in conjunction with the growing body of literature analyzing the gut microbiota of EBF and NEBF infants, nominate *Bifidobacteria* as an attractive target for potential probiotics and therapeutics. Ultimately, insights gained from previous and future studies can be applied toward the improvement of health outcomes for all infants for which exclusive breastfeeding is not possible.

**Data availability**

The data and R scripts required to reproduce the analyses used in this study are available at [https://github.com/itsmis
terbrown/GutMicrobes_analyses](https://github.com/itsmis
terbrown/GutMicrobes_analyses)

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

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