The early life microbiota mediates maternal effects on offspring growth in a nonhuman primate

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Highlights
Infants of low parity females grow fast despite poor maternal milk production
These infants have reduced gut microbiota diversity, more *Bacteroides fragilis* at 2–5 days old
The infant gut microbiota shares more ASVs with milk than with the maternal gut
A milk-oriented infant gut microbiota mediates faster growth to 6 months of age

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The early life microbiota mediates maternal effects on offspring growth in a nonhuman primate

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SUMMARY
Maternal parity can impact offspring growth, but the mechanisms driving this effect are unclear. Here, we test the hypothesis that vertically transmitted microbiota may be one potential mechanism. We analyzed 118 fecal and milk samples from mother-offspring vervet monkey dyads across the first 6 months of life. Despite poorer milk production, offspring born to low parity females grew larger than their counterparts. These offspring exhibited reduced alpha diversity in the first days of life, stronger seeding of maternal milk microbiota, Bacteroides fragilis dominance, and a greater abundance of glycan utilization pathways. Moreover, the attainment of greater body mass by 6 months of age was mediated by reduced early life alpha diversity and B. fragilis dominance. This work demonstrates that the establishment of a specialized, milk-oriented gut microbiota promotes infant growth and suggests an evolutionarily conserved developmental role of B. fragilis in primates.

INTRODUCTION
In both humans and nonhuman mammals, impaired postnatal growth during early life can result in delayed maturation, dysregulated metabolic function, and increased disease risk during later life (Barker, 2004; Fest-Bianchet et al., 2000; Wauters et al., 1993). The maternal environment is a particularly strong determinant of the pace of early growth: for example, mammalian infants born to reproductively inexperienced (i.e., low parity) mothers typically grow slower, are smaller in body mass for their age, and exhibit higher rates of mortality than infants born to multiparous females (Altman and Alberts, 2005; Clutton-Brock et al., 1987; Clutton-Brock and Pemberton, 2004; Ibañez et al., 2013; Ruiz-López et al., 2010; Smuts and Nicolson, 1989). One potential explanation for these negative outcomes is that maternal differences in milk production result in poorer transfer of nutrients to fuel offspring growth in some mother-offspring pairs. Because milk volume increases with each successive reproduction, low parity mothers produce smaller quantities of milk than high parity mothers, resulting in fewer available nutrients for infants (Carnicella et al., 2008; Hinde et al., 2009; Lang et al., 2012; Roy et al., 2003; Sevi et al., 2000; Tanaka, 1997; Xiccato et al., 2004).

In captive nonhuman primates, however, infants of high and low parity mothers grow at similar rates despite poorer milk production in the latter group (Nuñez et al., 2015), indicating that milk volume alone may not fully explain parity effects on infant growth. Indeed, infant utilization of milk nutrients not only depends on milk quantity but also on the presence of milk-digesting host-associated microbiota in the infant gut (Marcobal and Sonnenburg, 2012; Zivkovic et al., 2013). In humans, gut microbes such as Bifidobacteria and Bacteroides encode genes responsible for breaking down milk glycans, otherwise indigestible compounds that regulate infant immunity and metabolism (Garrido et al., 2013; Marcobal et al., 2011). These microbes can affect infant growth directly by digesting milk oligosaccharides and glycosylated proteins (~70% of human milk proteins, Kirmiz et al., 2018). For instance, in rodent transplant models of the human microbiome, pups with a greater relative abundance of gut Bacteroides fragilis grew faster, but only in the presence of milk glycans (Charbonneau et al., 2016).

Microbes like B. fragilis may also indirectly impact infant growth by modifying the resource landscape in the gut and reshaping microbiome community composition to prioritize somatic growth. Although a less diverse gut microbiota can indicate dysbiosis in adults, microbial homogeneity during early life may instead
reflect a specialized microbiota canalized for nutrient assimilation. Studies in human infants have found mixed effects of alpha diversity on postnatal growth (Blanton et al., 2016; Gough et al., 2016), but studies in birds demonstrate a link between reduced early life diversity of the gut microbiota and faster later-life growth (Banerjee et al., 2018; Davidson et al., 2021). Alpha diversity may therefore approximate the micro-
biological phenotype directly associated with growth, reflecting taxonomic and functional homogeneity.

More broadly, the maternal environment may program the infant gut microbiota for enhanced milk utilization via vertical microbial transmission. Thus far, parity effects on maternal microbiota have been demonstrated in agricultural species (Berry et al., 2021; Bogado Pascottini et al., 2021), and more recently in humans (Kervinen et al., 2021; Lopez Leyva et al., 2021). However, in pigs, parity effects extend to the infant gut microbiota, suggesting that such effects may be vertically transmitted from maternal reservoirs (Berry et al., 2021). Large-scale investigations into maternal vertical transmission have identified the maternal gut as the largest source of colonizing bacteria to the infant gut (Backhed et al., 2015; Ferretti et al., 2018; Makino et al., 2013; Maqsood et al., 2019). Yet, milk remains notably absent from these studies, obscuring an understanding of its relative role in establishing the infant microbiota. Beyond nutrients and glycans, milk harbors an ephemeral yet diverse microbiota that colonizes the infant gut from birth through weaning (Fernández et al., 2013; Hunt et al., 2011; Meehan et al., 2018; Mulet-Wolz et al., 2019; Petruzzo et al., 2019; Fehr et al., 2020). Microbial strains exclusively shared between milk and infant gut communities provide evidence of a unique milk-infant gut transmission pathway independent of other maternal reservoirs (Martín et al., 2012; Pannaraj et al., 2017).

Here, we investigate the relationship between parity, maternal and infant microbiota, and early growth in a nonhuman primate — the vervet monkey (Chlorocebus aethiops sabaeus). Vervet monkeys are established biomedical models for humans, and are particularly appropriate for investigating the relationship between maternal and infant microbiota and early growth. Like humans, vervets possess a milk microbiota that is highly diverse and abundant in growth-associated taxa such as B. fragilis (Petruzzo et al., 2019). We characterize the milk and fecal microbiota of mother-infant vervet monkey dyads in a captive population with a high rate of growth-associated infant mortality (>30% in the first month of life, typically the smallest infants; Fairbanks and McGuire, 1984; Kavanagh et al., 2011). This high mortality rate offers a unique opportunity to investigate parity-dependent effects in a naturalistic system in which growth is under strong selective pressure and in which the environment is free of the confounds commonplace in human studies (e.g., socioeconomic status, birth mode).

We test the hypothesis that the infant microbiota mediates the effects of maternal parity on postnatal growth, and investigate if and how maternal microbiota contribute to these effects. We expect that maternal parity predicts the pace of infant growth, maternal and infant microbiota vary with parity, and the effects of maternal parity on postnatal infant growth occur via changes to the infant gut microbiota. We characterize infant gut microbiota in three ways, focusing on community diversity, temporal changes in the abundance of microbial taxa, and the sharing of microbiota with maternal reservoirs (i.e., vertical transmission). We further predict that parity-dependent variation in the infant gut microbiota originates in maternal microbial communities, and expect to find that these patterns are recapitulated in the maternal microbiota, particularly milk.

**RESULTS**

**Maternal parity predicts infant postnatal growth**

Mother-infant vervet monkey dyads were sampled at three time points across early postnatal life (T1 = 2–5 days old, T2 = 4 months, T3 = 6 months, Figure 1A). Despite poorer maternal milk production (Figure S1), infants born to low parity females were larger in body mass at T3 (B = --0.02 ± 0.004, t = --4.51 p < 0.01; Figure 1B). Each successive parity was associated with a 0.02 kg decrease in body size at both time points (mean infant body mass at T3 = 1.20 kg). There was no relationship between maternal parity and infant body mass in the first days of life (B = 0.02 ± 0.011, t = 1.42, p = 0.18), suggesting that parity-related differences in infant growth emerge primarily across the first 6 months of postnatal life and do not reflect variation in fetal growth or birth weight.

**Parity shapes the infant and maternal microbiota**

**Alpha and beta diversity**

Amplicon sequencing of the 16S rRNA gene identified 1,956 unique amplicon sequence variants (ASVs) across all infant gut samples (mean ± SD = 276 ± 144 ASVs per sample, range = 45-542) compared to 2,019 unique ASVs across maternal gut samples (mean ± SD = 399 ± 94 ASVs per sample, range = 176-661) and 2,714 unique ASVs across maternal milk samples (mean ± SD = 484 ± 176 ASVs per sample, range = 152-742). Infant gut alpha-diversity (measured as Shannon Index) was low compared to
maternal communities but increased sharply at T2 ($\beta = 57.00 \pm 7.63$, $t = 7.47$, $p < 0.0001$; Figure 2A). Compositional differences between the gut microbiota of infants and their own mothers (i.e., dyadic beta diversity) were greater at T1 compared to T2 and T3 ($\beta = -0.04 \pm 0.007$, $t = -6.38$, $p < 0.0001$; Figure S2), suggesting that infants achieve an adult-like gut microbiota by 4 months of age.

Infants born to low parity females exhibited lower alpha-diversity than high parity infants at T1 ($\beta = 6.13 \pm 2.02$, $t = 3.04$, $p < 0.01$; Figure 2B), but not at T2 or T3. Of the two maternal communities, only milk microbiota varied with maternal parity, and the pattern was opposite to the infant gut: low parity females exhibited more diverse milk microbiota than high parity females at T1 ($\beta = -12.391 \pm 3836$, $t = -3.23$, $p < 0.01$; Figure 2C), and there was no relationship between maternal alpha diversity and parity at any other time point. There was also no relationship between maternal parity and beta diversity in the maternal or infant microbiota.

**Differential abundance of taxa**

ASVs in the infant gut resolved to 78 bacterial families, 42 of which were highly abundant (>1% of total community composition; Figure S3 and Table S1). Of these 42 families, 21 exhibited statistically significant changes in relative abundance from T1 to T2, and 11 exhibited changes from T2 to T3 (Benjamini-Hochberg adjusted p values: $p_{adj} < 0.05$; Table S2). At T1, the infant gut was dominated by a single ASV from the Bacteroidaceae family (assigned to *B. fragilis*; mean relative abundance 28.5%). By T2, infants had almost entirely lost Bacteroidaceae from the gut community, which instead became dominated by Prevotellaceae (36.0% relative abundance).

To identify which specific ASVs within these families contributed the most to the maturation of the infant gut microbiota, we used a Compositional Data Analysis (CoDA) framework to account for the compositional nature inherent to microbiome data (Gloor et al., 2017). PC1 explained 18.7% of the total variance in composition among infant samples and showed a clear distinction between early life (T1) samples and older samples (T2 and T3) (Figure 3A). Clustering on PC1 was most strongly explained by two ASVs, each with the highest absolute loading scores: *B. fragilis*, which dominated early life neonatal samples (T1), and *Prevotella copri*, which dominated older (T2 and T3) samples (Figure 3B and Table S2).

Of these two taxa, only *B. fragilis* was associated with maternal parity. Infants born to low parity females exhibited stronger early life dominance of *B. fragilis* compared to high parity infants ($\beta = -0.40 \pm 0.14$, $t = -2.75$, $p < 0.01$; Figure 3C). Moreover, a greater abundance of infant gut *B. fragilis* predicted lower alpha diversity at T1 ($\beta = -1.15 \pm 0.16$, $t = -7.25$, $p < 0.001$), indicating that *B. fragilis* dominance drives homogeneity in the infant gut microbiota during early life.

**Vertical transmission from maternal reservoirs**

At T1, the vertical transmission of microbiota from milk to the infant gut was stronger than from the maternal gut ($\beta = 9.50 \pm 4.32$, $t = 2.20$, $p < 0.05$; Figure 4A): infants shared 60 ASVs on average with their
mother’s milk (51% transmission rate), compared to 41 ASVs with their mother’s gut (42% transmission rate). ASV sharing between the infant gut and maternal milk microbiota was greatest in low parity dyads ($b = 7.51 \pm 2.03, t = -3.69, p < 0.01$; Figure 4B). There was no effect of maternal parity on infant gut ASVs shared with the maternal gut. By T2, the average transmission rates from milk and the maternal gut to the infant gut were not significantly different, and there was no longer an effect of maternal parity on transmission rates from milk, indicating a critical period of vertical transmission in the first days of life.

To pinpoint which shared ASVs may be important for infant development, we identified ASVs that were both highly prevalent (i.e., present in all infant gut samples in our dataset) and shared at high frequency with the maternal milk and gut communities (i.e., present in matched mother-infant samples across ≥ 90% of dyads). Three ASVs, *B. fragilis*, *Collinsella aerofaciens*, and *Eubacteria biforme*, were prevalent and shared at high frequencies. *B. fragilis* and *E. biforme* were shared exclusively between maternal milk and infant gut microbiota, whereas *C. aerofaciens* was shared universally across all three communities. The relative abundance of *C. aerofaciens* and *E. biforme* were similar across maternal and infant communities, but the infant gut housed the greatest proportion of *B. fragilis* ($b = 0.09 \pm 0.04, t = -2.01, p < 0.05$) and *B. fragilis* in their milk compared to their gut (Figure 4C), providing evidence that infant gut *B. fragilis* may originate in milk.

There was no relationship between maternal parity and the relative abundance of *C. aerofaciens* or *E. biforme* in the infant gut, nor was there a relationship between maternal parity and these two microbial taxa in the maternal gut. However, low parity females housed more *E. biforme* ($b = -0.09 \pm 0.04, t = -2.47, p < 0.05$) and *B. fragilis* ($b = -0.09 \pm 0.04, t = -2.01, p < 0.05$) in their milk compared to high parity females.

**Predicted functional capacity for glycan utilization**

We generated data on predicted gene function for infant and maternal milk microbiota using the PICRUSt2 pipeline to determine whether the microbiota of low parity dyads was enriched in pathways for the utilization of milk glycans. Indeed, low parity infants exhibited a greater relative abundance of microbial gene functional pathways related to glycan biosynthesis and metabolism in their guts than high parity infants in the first days of life (T1) ($b = -0.14 \pm 0.05, t = -2.90, p = 0.006$; Figure 5A). The milk microbiota of low parity mothers also housed a greater relative abundance of glycan utilization pathways than high parity mothers at this same timepoint (T1) ($b = -0.01 \pm 0.05, t = -2.90, p = 0.04$; Figure 5B).

**Infant gut microbiota mediate parity effects on postnatal growth**

We constructed three path analyses corresponding to the three parity-dependent compositional measures of infant gut microbiota at T1 (alpha diversity, *B. fragilis* abundance, and ASVs shared with the maternal milk microbiota).
microbiota) and found that parity effects on infant body mass at T3 were mediated by the infant microbiota during early life. In the first model, the effect of maternal parity on neonatal infant gut alpha diversity was the strongest path present ($b = 0.709$, $p < 0.01$), and parity indirectly influenced infant mass via reduced infant gut alpha diversity ($b = -0.269$, $p < 0.05$) (Figures 6A and 6B). A reduction in each unit of early life Shannon Diversity was associated with a gain of approximately 0.3 kg in body mass. In the second model, maternal parity influenced infant mass via the relative abundance of $B. fragilis$ in the infant gut ($b = 0.359$, Figures 6C and 6D). With each unit increase in the relative abundance of $B. fragilis$ at T1, infants gained approximately 0.4 kg in body mass. The third model detected no mediation effect of the proportion of ASVs shared with milk on the rate of infant growth.

DISCUSSION
To date, parity effects on maternal and infant microbiota have emerged primarily in agricultural animals, and have focused on microbial community changes without establishing a direct link to infant development. Here, we show that parity not only impacts the infant and maternal microbiota in a nonhuman primate, but also influences the pace of early growth via modeling of the infant gut microbiota in the first days of life. Low parity females produced less milk than high parity females, but infants born to low parity females housed a specialized, milk-oriented microbiota dominated by $B. fragilis$ that promoted faster growth to 6 months of age. Patterns of $B. fragilis$ were recapitulated only in the maternal milk microbiota, revealing this maternal reservoir to be the likely source of parity-dependent variation in the infant gut. Consistent with previous studies in humans and rhesus macaques (Janiak et al., 2021; Thomson et al., 2018) but counter to a recent study in wild chimpanzees (Reese et al., 2020), alpha diversity of the vervet infant gut microbiota was lowest during early life (T1), increasing rapidly toward maternal levels by 4 months of age (T2). Changes in the infant gut microbiota with age were most strongly explained by $B. fragilis$ and $P. copri$. A trade-off between Bacteroides and Prevotella has been well-described in microbiome research for over a decade as a reflection of dietary differences between human populations consuming diets high in protein and fat (characterized by more Bacteroides) and those consuming a fiber-rich diet (characterized by more Prevotella) (Ley, 2016; Lozupone et al., 2012). In infant vervet monkeys, a trade-off between these taxa may reflect adaptive plasticity of the gut microbiome as infants transition from the consumption of milk to solid foods. Unlike $B. fragilis$’s role in milk glycan consumption (Coyne et al., 2008; Marcobal et al., 2011), $P. copri$ cannot digest glycans (Ley, 2016). Indeed, nursing piglets exhibit greater Bacteroidaceae, whereas weaned piglets exhibit greater Prevotellaceae (Frese et al., 2015).

We also found that low parity vervet infants exhibited less gut microbiota diversity than high parity infants, and low parity mothers harbored more diverse milk microbiota than high parity mothers. The opposing directionality of these relationships suggests that the reduced alpha diversity characteristic of the low diversity.
parity infant gut microbiota is not simply a result of vertical transmission from a less diverse maternal milk microbiota. Instead, this pattern provides evidence that infants raised by reproductively inexperienced mothers may selectively seed taxa from a highly diverse milk microbiota (Coyte et al., 2015).

All infants, regardless of parity, selectively seeded *B. fragilis* in the first days of life (T1), but *B. fragilis* was most abundant among infants born to low parity mothers and drove homogeneity of the infant gut microbiota in early life. A greater abundance of *B. fragilis* at T1 predicted the overall lower gut microbiota alpha diversity at both T1 and T2. *B. fragilis* can modulate the biochemical makeup of the gut environment (Chatzidaki-Livanis et al., 2017; Coyne et al., 2008), in turn altering subsequent colonization by other taxa. Though largely absent from the infant gut microbiota at T2, *B. fragilis* dominance at T1 may therefore explain the persistence of reduced community diversity via regulation of immigration order (Coyne et al., 2008; Marcobal et al., 2011).

Infants shared ~10% more ASVs with their mother’s milk microbiota than with her gut microbiota at T1 (i.e., a transmission rate from milk of ~51 vs. ~42% from the maternal gut), suggesting that vertical transmission from mothers to infants occurred predominantly via milk. These rates of transmission are higher than previously reported transmission rates in humans (milk: Pannaraj et al., 2017, maternal gut: Ferretti et al., 2018). Higher rates identified in our study may reflect stronger transmission via milk in nonhuman primates compared to humans, and contrast with previous studies in humans that suggest, despite the absence of milk samples, that the maternal gut is the largest source of taxa to the infant gut (Asnicar et al., 2017; Backhed et al., 2015; Ferretti et al., 2018). The incorporation of milk microbiota into the infant gut appears to be under strong evolutionary selection in primates, and may be otherwise obscured in human studies because of a historic focus on other maternal reservoirs.

Although there was no relationship between maternal parity and ASV sharing with the maternal gut microbiota at any age, infants of low parity females shared more ASVs with their mother’s milk microbiota than with higher parity infants at T1. There are a few potential explanations for this finding. First, the dispersal of milk microbiota may be enhanced in low parity females. The vervet monkey milk microbiota is highly individualized, with many ASVs unique to the host (Petrullo et al., 2019). Low parity females may possess more unique variants that are better adapted for dispersal to and survival within the infant gut (Duranti et al., 2017), an explanation also supported by our finding of greater taxonomic diversity in the milk of low parity females. Second, low parity females may exhibit variation in nonmicrobial components of milk (e.g., oligosaccharides, immunoglobulin A), which can modulate the infant gut environment in a manner that favors colonization by maternal milk ASVs over other taxa (Kirimiz et al., 2018; Rogier et al., 2014). Finally, low parity infants may selectively seed maternal milk microbiota more strongly than high parity infants via...
infant-specific mechanisms that promote colonization by maternal milk ASVs (e.g., phage activity, mucosal immunity, and pH) (Coyte et al., 2015).

High frequency sharing of particular ASVs in milk and infant gut communities provides evidence that the dyadic transmission and seeding of these taxa may be evolutionarily conserved (Korpela et al., 2018). At T1, there was a 15-fold higher abundance of \textit{B. fragilis} in the milk microbiota compared to the maternal gut microbiota. Moreover, the relative abundance of \textit{B. fragilis} was greater in the milk microbiota of low parity females compared to high parity females. Together, these results suggest that milk may be the primary reservoir of \textit{B. fragilis} to the infant gut. Given that neither the maternal vaginal nor gut microbiota explain variation in infant \textit{Bacteroides} colonization in prior studies in humans (Mitchell et al., 2020), our data suggest that milk may be the primary source of \textit{B. fragilis} in primates.

Stronger vertical transmission of \textit{B. fragilis} in low parity dyads may enhance host digestion of glycans from lower volumes of milk, resulting in a microbial community that is more efficient at assimilating milk components to fuel growth. Low parity females in this population can produce as much as 5x less milk than high parity females (Figure S1), necessitating a compensatory mechanism to mitigate the inequity in milk volume. We found that the maternal milk and infant gut microbiota of low parity dyads exhibited a greater abundance of gene pathways related to glycan metabolism compared to the maternal gut microbiota. Increased metabolism of milk glycans can provide the energetic resources to fuel host growth and development (Charbonneau et al., 2016). Thus, the \textit{B. fragilis} dominance and enhanced glycan utilization characteristic of the low parity infant gut microbiota may reflect a compensatory strategy toward a homogeneous gut microbiota suited to assimilate energy from lower volumes of milk.

Finally, infants born to low parity females were larger in body mass at 6 months of age, and this effect was mediated by reduced alpha diversity at T1, in line with previous studies on birds (Banerjee et al., 2018; Davidson et al., 2021). Mass at 6 months was also mediated by \textit{B. fragilis} dominance at T1, in line with experimental studies on the human microbiota (Blanton et al., 2016; Charbonneau et al., 2016). These findings support our hypothesis of a mediating role of microbiota on parity-dependent infant growth, but the faster, rather than slower, rates of growth among low parity infants are in contrast with prior work in wild mammals (e.g., Clutton-Brock and Pemberton, 2004; Ibáñez et al., 2013; Ruiz-López et al., 2010). One potential explanation for this pattern is that the nutrient abundance present in captivity may exploit the compensatory mechanisms that allow low parity mothers to successfully rear infants despite poor milk production in
the wild. Such exploitation, coupled with the “humanizing” effect of captivity on the nonhuman primate microbiota (Clayton et al., 2016; Houtz et al., 2021), may better approximate the microbial pathways that mediate maternal effects in modern human populations inhabiting calorie-rich environments. In addition, viability selection exerts substantial pressure on infant development in populations with high rates of early mortality (McAdam and Boutin, 2003; Mojica and Kelly, 2010). In our study population, infants face a growth-associated mortality rate of more than 30% (Fairbanks and McGuire, 1984) and individuals who die early in life are smaller in body size than their counterparts (Kavanagh et al., 2011), potentially amplifying growth-associated microbial effects. Our findings may thus extend to human populations experiencing stunted childhood growth and growth-associated early mortality by identifying 

\[ B. \text{fragilis} \]

as a key predictor of postnatal body mass.

**Limitations of the study**

This study was limited in a few ways. First, although maternal vaginal microbiota may contribute to or modify the above-described relationships, logistic constraints prevented vaginal microbiota sampling...
in this study. Despite this limitation, the vaginal microbiota is unlikely to be the primary contributor of maternal \textit{B. fragilis} to the infant gut as \textit{Bacteroides} are not common members of the human vaginal microbiota and even when present, do not appear to account for transmission to the infant gut (Mitchell et al., 2020). Given the importance of \textit{B. fragilis} in mediating parity effects on infant growth in this population, our interpretation of milk as the potential source of \textit{B. fragilis} is likely independent of vaginal microbiota’s effects. Second, the correlation between maternal parity and maternal age raises the question of whether the relationships described here may reflect age, rather than parity, effects. Because our hypotheses were centered on the effects of successive reproductions on maternal communities, and because prior work has shown that parity, not age, shapes the maternal microbiota (Berry et al., 2021), we did not include maternal age in our analyses to avoid issues of multicollinearity. Instead, we focused on parity because successive reproductions (independent of maternal age) shape the morphology of both the mammary gland (Rovai et al., 1999) and variation in milk production across mammalian species (e.g., Lang et al., 2012; Sevi et al., 2000), suggesting that similar effects may be found in maternal milk and infant gut microbiota.

**STAR METHODS**

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**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.isci.2022.103948.

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**AUTHOR CONTRIBUTIONS**

Concept, Design, and Interpretation of data, L.P. and A.L.; Data acquisition and Methodology, L.P., A.B., M.J.J., S.S., N.S.M., and A.L.; Writing – Original Draft, L.P. and A.L.; Review & Editing, A.B., M.J.J., S.S., and N.S.M.; Funding, L.P., M.J.J., N.S.M., and A.L.; Supervision, A.L. All authors have approved this version of the manuscript.
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**STAR METHODS**

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Biological samples** | | |
| Maternal (milk and fecal) and infant (fecal) samples used for microbiome analysis | Author’s collection (Wake Forest School of Medicine Vervet Research Colony) | https://doi.org/10.6084/m9.figshare.14561925 |
| **Deposited data** | | |
| 16S rRNA amplicon sequences from 118 fecal and milk samples | This paper | www.ncbi.nlm.nih.gov/sra/PRJNA728247 (NCBI Sequence Read Archive) |
| **Experimental models: Organisms/strains** | | |
| 18 adult female and 18 mixed-sex infant vervet monkeys (Chlorocebus aethiops sabaueus) | Vervet Research Colony, Wake Forest University | N/A |
| **Software and algorithms** | | |
| QIIME2 | Hall and Beiko, 2018 | https://qiime2.org/ |
| DADA2 | Callahan et al., 2016 | https://benjneb.github.io/dada2/ |
| MAFFT | Katoh et al., 2002 | https://mafft.cbrc.jp/alignment/software/ |
| PICRUSt2 | Douglas et al., 2020 | https://nephele.niaid.nih.gov/show_picrust2_details/ |

**RESOURCE AVAILABILITY**

**Lead contact**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Lauren Petrullo (petrullo@umich.edu).

**Materials availability**

This study did not generate new unique materials.

**Data and code availability**

- 16S rRNA gene sequences are available under BioProject ID PRJNA728247 at the NCBI Sequence Read Archive (SRA) (www.ncbi.nlm.nih.gov/sra/PRJNA728247) and associated metadata for this project are available at figshare: https://figshare.com/s/79f3cf6cf410f5a87d31.

- Code for this project is available at figshare: https://figshare.com/s/79f3cf6cf410f5a87d31.

- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

We studied 18 mother-infant captive vervet monkey (Chlorocebus aethiops sabaueus) dyads housed at the Vervet Research Colony (VRC) at the Wake Forest University School of Medicine in Winston-Salem, North Carolina, USA. Infant vervet monkeys <6 months of age of both sexes were included in this study. All animal use procedures and sample collection methods followed national and international guidelines and were approved by the Institutional Care and Use Committee (IACUC) at Wake Forest University School of Medicine. Vervets are highly social primates that breed once annually (Else et al., 1986). Mother-offspring dyads were housed at the VRC across 8 indoor-outdoor social groups organized by female kin. Thirteen of the dyads were provisioned daily with commercial monkey chow (Purina Monkey Chow, LabDiet 5038), while four dyads were fed a western-style lab diet (LabDiet SL3K) as part of a different study. In addition, some dyads were fed ad libitum, while feeding was time-restricted for others. However, it is important to note that infants in this population do not appear to incorporate solid foods into their diets until about 4-6 months (Fairbanks, 1988), thus the effect of diet on early life microbiota at T1 in this study is expected to...
be minimal. Animals were supplemented with fresh fruits and vegetables daily. Infants included in this study were delivered via unassisted vaginal births between June 18, 2017 and August 27, 2017 after typical ~5.5 month-long gestations.

**METHOD DETAILS**

We collected physiological data from dyads at three postnatal timepoints: T1 (2-5 days postpartum), T2 (4 months postpartum), and T3 (6 months postpartum). Two infants died within the first month of life as a probable consequence of poor maternal milk production: in one case, the mother was not producing any milk, and in the second case, the mother produced milk in only one mammary gland. Thus, both infants contributed fecal samples and somatometric data at T1 only. The T1 milk sample from the female with unilateral milk production was excluded from this study. Milk collection followed previously published protocols for the collection of milk from cercopithecine monkeys (Hinde et al., 2009; Petrullo et al., 2019). Both mammary glands were fully evacuated via manual expression into a single sample tube. Samples were placed immediately on ice, briefly vortexed, aliquoted into cryovials, and frozen at −80°C until shipment to the University of Washington. We collected maternal and infant fecal samples at all three sampling timepoints by briefly inserting standard (mothers) and pediatric (infants) flocked nylon swabs (FLOQSwabs, CO-PAN Diagnostics, Murrieta, CA) into the anal canal. Rectal swabs are reliable proxies for fecal samples when quantifying an animal’s gut microbiota, including among infants (Bassis et al., 2017; Reyman et al., 2019). Swabs were gently spun in the canal 2-3 times before removal and were snapped off into empty polypropylene tubes. Samples were immediately frozen at −80°C before being shipped to the University of Washington.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**DNA extraction, amplification, and library preparation**

We extracted microbial DNA from milk using the PowerFood Microbial kit (Qiagen) following the manufacturer’s kit protocols, with the addition of two front-end processing steps shown to increase overall yield from milk (Petrullo et al., 2019). For fecal swabs, microbial DNA was extracted using the Qiagen PowerLyzer PowerSoil kit. Samples were thawed to room temperature, swab tips were snapped into glass bead tubes (0.1 mm), and mechanically lysed (30 Hz for 2 min). Extractions then followed the manufacturer’s protocol.

We amplified the hypervariable V4 region of the 16S rRNA gene using PCR primer set 515F and 806R from The Human Microbiome Project (Gilbert et al., 2014; Yang et al., 2016), following previously published protocols (Petrullo et al., 2019). A fragment analyzer was used to confirm amplification of the V4 region prior to quantification via a qubit fluorometer.

**Sequencing and bioinformatics**

Amplicon libraries were balanced and pooled. Library complexity was increased by spiking libraries with PhiX prior to sequencing all libraries together on a single Illumina MiSeq flow cell using 301 bp paired-end sequences. This resulted in 1,456,972 reads for milk samples, 2,491,940 reads for adult fecal samples, and 2,147,458 reads for infant fecal samples. Sequences were analyzed using the QIIME2 platform (Caporaso et al., 2011; Hall and Beiko, 2018) and denoised with Divisive Amplicon Denoising Algorithm 2 (DADA2: Callahan et al., 2016) as a QIIME2 plug-in. In contrast to clustering sequencing reads based on a fixed dissimilarity threshold (e.g., the assignment of Operational Taxonomic Units [OTUs]), which can confound sequencing errors with biological variation, DADA2 infers sequences exactly (resulting in amplicon sequence variants, hereafter referred to as ASVs), providing higher taxonomic resolution than OTUs (Callahan et al., 2016, 2017). This approach is crucial for assessing changes in microbial composition (Tikhonov et al., 2015) as well as community diversity (Rosen et al., 2015), particularly in microbial communities presumed to be low in diversity based on prior OTU clustering methods (e.g., the human vaginal microbiota, Callahan et al., 2017), as well as determining vertical transmission between maternal and infant communities. Forward and reverse reads were trimmed to 240 bases long to remove the low-quality portion of the sequences. Next, the forward and reverse reads were merged and chimeric sequences removed. After filtering, trimming, merging, and chimera removal, we retained 811,225 16S rRNA gene sequences from 29 milk samples (28,972 ± 17,165 reads per sample), 1,547,442 sequences from 45 adult female fecal samples (48,357 ± 15,100 reads per sample), and 2,147,458 sequences from 45 infant fecal samples (47,721 ± 12,580 reads per sample). Only fecal samples with >15,000 reads were retained for analyses, resulting in N=44 infant fecal samples. No other samples were removed at this stage as sequencing depths...
across samples were highly balanced. ASVs were aligned using MAFFT (Katoh et al., 2002) and a phyloge-
netic tree was constructed using fasttree2 (Price et al., 2010). Taxonomic assignment of ASVs was per-
formed using the q2-feature-classifier in QIIME2 against the 13_8 version of the GreenGenes database
(McDonald et al., 2012) based on 100% similarity to the reference sequence. One milk sample was excluded
from the analysis as all of its 200,000+ reads were assigned to a single ASV.

Statistical analyses
All analyses for this study were performed in QIIME2 (Hall and Beiko, 2018) and R version 4.0.2 (R. Core
Team, 2015). Taxonomy and count tables were imported from QIIME2 into R using the qiime2R package
(Bisanz, 2018). As rarefaction of microbiota count data results in a loss of precision and variation (McMurdie
and Holmes, 2014), we instead used a compositional data, or “CoDa”, approach when necessary (Gloor
et al., 2017) and controlled for sequencing depth by including the log-transformed count of total reads
in a sample as an offset in all statistical models. Where microbial taxa were converted to relative abund-
dances, the taxonomy table was filtered to retain only those with > 1% relative abundance. Residuals of
all linear models described below were found to be normally distributed following visual evaluation by
Q-Q plots and Shapiro-Wilk tests. Because of the potential confounding effects of dietary composition
and feeding schedule, diet was controlled for as a fixed factor in all analyses. Figures were created in R us-
ing ggplot2 (Wickham, 2011) and visreg for partial residual plots (Breheny and Burchett, 2017). The graph-
ical abstract, Figures 1A, 6A, and 6C were created using biorender (www.biorender.com).

Milk yield
Linear models were used to test the effects of maternal parity on milk production (mL). Residuals for all
models were normally distributed upon visual inspection and evaluation with a Shapiro-Wilk test, and
therefore not transformed prior to analysis. We constructed models with milk yield (mL) as the dependent
variable, with maternal parity as a fixed factor while controlling for offspring sex (Hinde, 2009; Hinde et al.,
2009) and dietary composition.

Infant body mass
Linear models were used to test whether maternal parity predicted infant body mass at T3 (dependent var-
iable), controlling for infant sex, approximate birth weight (body mass at T1), and dietary category. T3 body
mass data were log-transformed prior to modelling to ensure normality of the model residuals.

Microbiome alpha & beta diversity
Microbiome samples were assessed at the community level using both alpha (within sample) and beta (be-
tween samples) diversity measures. Alpha diversity was calculated using the phyloseq package (McMurdie
and Holmes, 2013) and quantified using the Shannon Index of alpha-diversity (Shannon, 1948; Spellerberg
and Fedor, 2003), which takes into account both the richness and evenness of taxa. Shannon Indices were
transformed by Tukey transformation to achieve normality of residuals, and linear mixed models were con-
structed to test associations between infant age (categorical fixed effect: T1, T2, T3) and gut microbiome
diversity (Shannon Index, dependent variable), controlling for diet (fixed effect) and infant ID (random ef-
fact). Weighted UniFrac distance matrices were constructed in QIIME2 and were used to assess dissimilarity
between the gut microbiome of infants and their own mothers (dyadic weighted UniFrac distance) as a
measure of infant gut microbiome maturation. A linear mixed model was used to test whether infant age
(fixed effect) predicted dyadic weighted UniFrac distance (dependent variable), controlling for diet
(fixed effect) and infant ID (random effect).

Differential abundance testing
Differential abundance testing of taxa within the infant gut microbiome was performed in R using the
NBZIMM package (Zhang and Yi, 2020). Negative binomial models (glmer.nb() function) were used test
the effects of infant age (fixed effect) on the abundance of taxonomic families (dependent variable: raw
counts), controlling for diet (fixed effect), sequencing depth fit as an offset (log transformation of total num-
ber of reads in the sample), and infant ID (random effect). Models that did not converge due to a high pro-
portion of zeros were rerun using the same package as zero-inflated Gaussian mixed models (lme.zig() function).
P-values were adjusted using the Benjamini-Hochberg FDR multiple-test correction (Benjamini
and Hochberg, 1995) to account for multiple hypothesis testing and microbial taxa with adjusted p values <
0.05 were considered statistically significant.
Predicted functional profiles
To predict gene functional content from 16S rRNA data, we used the Phylogenetic Investigation of Communities by Reconstruction of Unobserved States 2 (PICRUSt2) v.2.1.3-b software (Douglas et al., 2020) pipeline with default options (picrust2_pipeline.py). PICRUSt2 predicts gene family abundance by using the table of ASVs generated by QIIME2 and their representative sequences. We then inferred Kyoto KEGG (Kyoto Encyclopaedia of Genes and Genomes) pathway abundances from the predicted KEGG ORTHOLOGY (KO) at Level 2 of the BRITE map (Kanehisa et al., 2014). Counts of functional pathways were transformed to relative abundance for each sample and a filtering threshold of 0.1% abundance was implemented to retain only the most abundant pathways. To estimate the accuracy of our PICRUSt2 predicted pathways, we calculated a weighted NSTI (Nearest Sequenced Taxon Index) score for each sample, which estimates the phylogenetic distance between OTUs and their sequenced reference genomes, to achieve an average NSTI score. The average NSTI score for all samples was 0.087 (SE ± 0.038, range: 0.011–0.28), reflecting high (~91%) distinguishability.

Linear mixed-effect models were used to test the effects of maternal parity on glycan utilization and metabolism pathways in the infant gut microbiome (dependent variable) as a function of sampling timepoint, controlling for diet (fixed factors) and infant ID (random effect).

Principal Components Analysis & CoDa framework
A Compositional Data (CoDa) framework was used to investigate how specific ASVs contributed to the compositional patterns of infant gut microbiome maturation (Gloor et al., 2017). Raw count data of ASV reads were normalized using a centered log-ratio (clr) transformation and a pseudocount of 0.65 in place of zeroes (Aitchison, 1982). We then ordinated the samples based on their compositional dissimilarity using a Principal Components Analysis (PCA; Hotelling, 1933) using pairwise Euclidean distances between samples (i.e., Aitchison distance). Unlike Principal Coordinates Analysis (PCoA), which is more commonly used in microbiome data analysis, PCA is not driven by presence/absence data, is more reproducible, and is more robust against sparsity (Gloor et al., 2017). The PCA was used to visualize sample clustering and generate loading scores for each ASV on the first PCA axis (which captured 18.7% of variability and was highly predicted by infant age). The loading scores were then sorted by absolute value to determine the microbial taxa with the highest influence (e.g., absolute value loading score) on the observed clustering of the first PCA axis (i.e., the most predictive of infant age) (Martino et al., 2019). We then calculated a log-ratio of the relative abundance of the two most influential taxa (Bacteroides fragilis and Prevotella copri) and used Pearson’s correlation tests to determine the correlation coefficient between the log-ratios and samples’ coordinates on the first PCA axis. Subsequently, we used a linear mixed model to test whether the log-ratio of these taxa (fixed effect) predicted dyadic weighted UniFrac distances between infant and maternal gut microbiomes (dependent variable), controlling for infant age (fixed factor), diet (fixed factor), and dyad ID (random).

Vertical transmission
The proportion of ASVs shared between maternal and offspring microbiota is a commonly-used index of vertical transmission using 16S rRNA gene data (Björk et al., 2019; Korpela et al., 2018; Maqsood et al., 2019; Renelies-Hamilton et al., 2021). We calculated the proportion of shared ASVs as being the proportion of all ASVs present within the infant gut microbiome (denominator) that were also found in maternal milk and gut microbiomes (numerator). To test whether these proportions differed significantly between infant gut/maternal gut versus infant gut/maternal milk comparisons, we fit linear mixed models with proportion of ASVs shared at T1 and T2 as the dependent variable and comparison (milk vs. infant gut or maternal gut vs. infant gut) as a predictor variable, controlling for individual ID. To test whether maternal parity affected vertical transmission, generalized linear models were used to test whether maternal parity predicted the raw count of shared ASVs (dependent variable), controlling for diet and the total number of ASVs present within the infant gut.

Path analysis
To test for a mediation effect of the microbiome on infant growth, we performed path analyses by constructing three separate path models (one for each parity-dependent measure of infant gut microbiome composition: alpha diversity, B. fragilis abundance, and vertical transmission via the milk microbiome) using the piecewiseSEM() in R (v. 2.0.1) (Lefcheck, 2016). Unlike simple structural equation modeling,
piecewiseSEM is robust to small sample sizes and can integrate mixed effects and generalized linear models (e.g., negative binomial models) (Lefcheck, 2016), making it more suitable for use with microbiome count data. Each path model was comprised of two recursive component models: 1) a model testing direct/indirect effects of both maternal parity and the relevant measure of infant gut microbiome composition at T1 on infant body mass at T3, controlling for confounding variables, and 2) a model testing the effect of maternal parity on the infant gut microbiome, controlling for confounding variables. Component models were integrated into a single causal network using the psem() function, which produced weighted coefficients and corresponding p-values to permit the assessment of the significance and relative strengths of each path within the overall network. The global goodness-of-fit was assessed for each model in piecewiseSEM() using Shipley’s test of d-separation (Shipley, 2013). All models had Fisher’s C statistics with $p > 0.05$, indicating that the models were suitable fits and no major paths were missing.