

Early life dynamics of the human gut virome and bacterial microbiome in infants

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The early years of life are important for immune development and influence health in adulthood. Although it has been established that the gut bacterial microbiome is rapidly acquired after birth, less is known about the viral microbiome (or 'virome'), consisting of bacteriophages and eukaryotic RNA and DNA viruses, during the first years of life. Here, we characterized the gut virome and bacterial microbiome in a longitudinal cohort of healthy infant twins. The virome and bacterial microbiome were more similar between co-twins than between unrelated infants. From birth to 2 years of age, the eukaryotic virome and the bacterial microbiome expanded, but this was accompanied by a contraction of and shift in the bacteriophage virome composition. The bacteriophage-bacteria relationship begins from birth with a high predator–low prey dynamic, consistent with the Lotka–Volterra prey model. Thus, in contrast to the stable microbiome observed in adults, the infant microbiome is highly dynamic and associated with early life changes in the composition of bacteria, viruses and bacteriophages with age.

The intestinal microbiome includes bacteria, eukaryotic viruses, bacterial viruses (bacteriophages), fungi and archaea. It has been established that some of these microorganisms interact with the immune system and influence their host's health^{1,2}. Alterations in the intestinal bacterial microbiome have been implicated in a wide range of human diseases including cirrhosis, diabetes and inflammatory bowel disease^{3–5}. Most therapeutic strategies targeting the microbiome, such as probiotics, prebiotics and fecal microbial transplantation, aim to modulate the bacterial microbial community^{6,7}. The bacterial microbiome is established soon after birth, and its composition changes over the next several years toward a stereotypical 'adult-like' bacterial community structure^{8–11}. This process can be influenced by multiple interacting factors such as nutrition, delivery route, antibiotic use and geographical setting^{8–15}. Studies of twins demonstrate that infants share a more similar bacterial microbiome with their co-twin than with unrelated individuals^{14,16–18}.

Much less is known about the viral microbiome (virome)¹⁹, a diverse community consisting of eukaryotic RNA and DNA viruses and bacteriophages. Emerging evidence indicates that the virome plays a role in human health. The burden of anellovirus (a eukaryotic DNA virus) is directly correlated with the degree of host immunosuppression and with organ transplant outcome and is an indicator of pediatric febrile illness and AIDS^{20–23}. Pathogenic simian immunodeficiency virus is associated with expansion of the enteric virome, which includes many eukaryotic RNA viruses²⁴. Additionally, chronic virus infection can confer increased resistance against pathogenic challenges²⁵, indicating that the virome may provide beneficial effects to the host¹.

The intestinal microbiota also contains diverse bacteriophages, which in healthy adults consist mostly of members of the order *Caudovirales* and family *Microviridae*. These bacteriophages typically maintain a stable community over time^{17,26–28}. Shifts in the enteric bacteriophage community composition have been associated with Crohn's disease and ulcerative colitis²⁹. However, unlike in environmental ecosystems, where changes in population dynamics of bacteriophage–bacteria interactions are known to follow a Lotka–Volterra "predator–prey" model^{30–32}, the predator–prey relationship between bacteriophages and bacteria has yet to be observed in the human intestinal microbiome¹⁷. Metagenomic studies of the healthy infant gut virome are limited to one study of a single infant in which the DNA virome was analyzed at a single time point using modest-depth Sanger sequencing³³. Targeted PCR and RT-PCR studies have determined that some eukaryotic viruses, such as picornaviruses and anelloviruses, can be frequently found in stools of healthy infants³⁴. Although metagenomic analyses of the gut virome of children with diseases such as diarrhea and acute flaccid paralysis have been described^{35–37}, to date, there has been no longitudinal analysis of the virome of a cohort of healthy infants.

Given that the bacterial microbiome is established during early infancy and is likely to affect long-term health^{8,9,14,16,38}, we examined the changes in the eukaryotic viruses and bacteriophages that accompany human development. To elucidate the degree of interindividual and intraindividual variability in the virome, we sequenced stools of a healthy monozygotic twin pair and three healthy dizygotic twin pairs. In this study, we defined 'healthy infants' as those having no apparent

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Virome of infants during early development

a

8 infants (4 pairs of twins): fecal specimens

0 3 6 12 18 24 months

Filtered through 0.4 μ M
Total nucleic acid extraction

Bead beating disruption
Total nucleic acid extraction

Multiple displacement
amplification
(MDA)

Sequence independent
DNA & RNA
amplification
(SIA)

16S
PCR
amplification

Next-generation sequencing

b

Method

A2-24 B1-6 B1-18 B2-6 B2-18 C1-6 C2-6

SIA MDA SIA MDA SIA MDA SIA MDA SIA MDA SIA MDA

Reads

1 100 >1,000

Alphaflexiviridae
Caliciviridae
Picornaviridae
Tombusviridae
Virgaviridae
Chrysovriidae

Anelloviridae
Circoviridae
Geminiviridae
Nanoviridae
Parvoviridae
Adenoviridae

Inoviridae
Microviridae
Corticoviridae
Podoviridae
Myoviridae
Siphoviridae
Tectiviridae

Unclass. Caudovirales
Unclass. dsDNA phages
Unclass. phages

Lipothirixviridae

Eukaryotic RNA viruses

Eukaryotic DNA viruses

Bacteriophages

Archaeal viruses

c

Age (0–24 months)

A1 A2 B1 B2 C1 C2 D1 D2

Alphaflexiviridae
Astroviridae
Caliciviridae
Picornaviridae
Tombusviridae
Virgaviridae
Chrysovriidae
Picobirnaviridae

Anelloviridae
Circoviridae
Geminiviridae
Nanoviridae
Parvoviridae
ssDNA satellites
Adenoviridae
Polyomaviridae

Inoviridae
Microviridae
Corticoviridae
Myoviridae
Podoviridae
Siphoviridae
Tectiviridae

Unclass. Caudovirales
Unclass. dsDNA phages
Unclass. phages

Lipothirixviridae

Unclass. viruses
Unclass. ssDNA viruses
Environmental samples

ssRNA

dsRNA

ssDNA

dsDNA

ssDNA

dsDNA

dsDNA

dsDNA

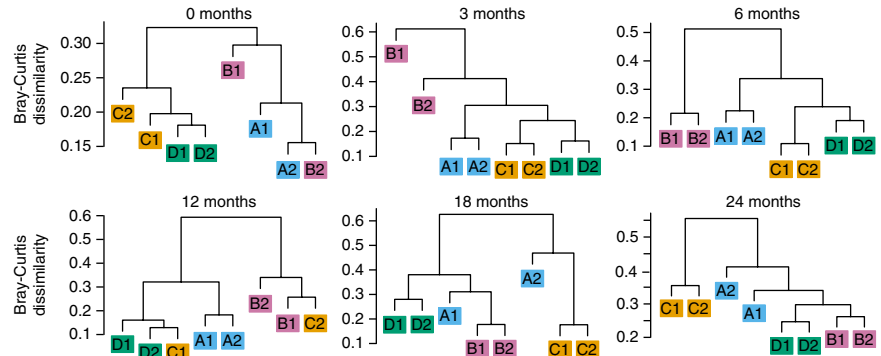
Archaeal viruses

Other viruses

Presence
Absence

1229

Figure 2 Analysis of virome beta diversity. Agglomerative hierarchical clustering of Bray-Curtis dissimilarity of virome communities (eukaryotic viruses and bacteriophages; genera) at the indicated ages.



sequence-independent DNA and RNA amplification (SIA) (Fig. 1a). MDA is commonly used in virome studies^{17,26,29} because of the high processivity of the phi29 polymerase used in this approach. However, MDA cannot detect RNA viruses, and its use also leads to preferential amplification of small circular DNA viruses⁴⁰. To complement MDA, therefore, we used the SIA approach, which is capable of detecting RNA as well as DNA viruses^{35,36}, although its sensitivity for DNA virus detection is generally lower than that of MDA (Fig. 1b). The libraries were pooled and sequenced on the Illumina MiSeq platform. On average, we obtained $549,301 \pm 207,521$ (mean \pm s.d.) reads per MDA sample library and $551,592 \pm 229,210$ reads per SIA sample library (Supplementary Fig. 1b). Sequencing reads were adaptor trimmed, quality filtered, taxonomically assigned and randomly subsampled to 200,000 reads per sample library for the analyses (details in Online Methods). To define the global virome composition, we merged the profiles of virus families identified by MDA and SIA into a presence-absence heatmap (Fig. 1c). Consistent with results from other PCR-based studies³⁴, we identified eukaryotic RNA viruses such as caliciviruses, astroviruses and picornaviruses in the infant fecal specimens. Additionally, bacteriophage families were more frequently detected than eukaryotic RNA and eukaryotic DNA viruses. Only one archaeal virus family (*Lipothrixviridae*) was identified.

To compare the virome biodiversity between individuals, we measured beta diversity at the virus genus level using the unweighted Bray-Curtis distance. Principal-coordinate analysis of the virome (eukaryotic viruses and bacteriophages) suggested that co-twin relatedness and age contributed to the variation in virome biodiversity of these infants (Supplementary Fig. 2a,b). Agglomerative hierarchical clustering supported the hypothesis that the virome community composition was typically more similar between co-twins than between unrelated infants after controlling for age (Fig. 2).

The eukaryotic virome is acquired after birth

We next focused on the assembly of the eukaryotic virome. Eukaryotic viral population richness was low in the earliest-in-life specimens and increased thereafter (Fig. 3a, Wilcoxon test $P < 0.05$), suggesting that the eukaryotic virome is primarily established through environmental exposures. Among the eukaryotic RNA viruses identified, the most commonly detected viral genera were enterovirus, parechovirus, tombamovirus and sapovirus (Fig. 3b). The relative sparsity of

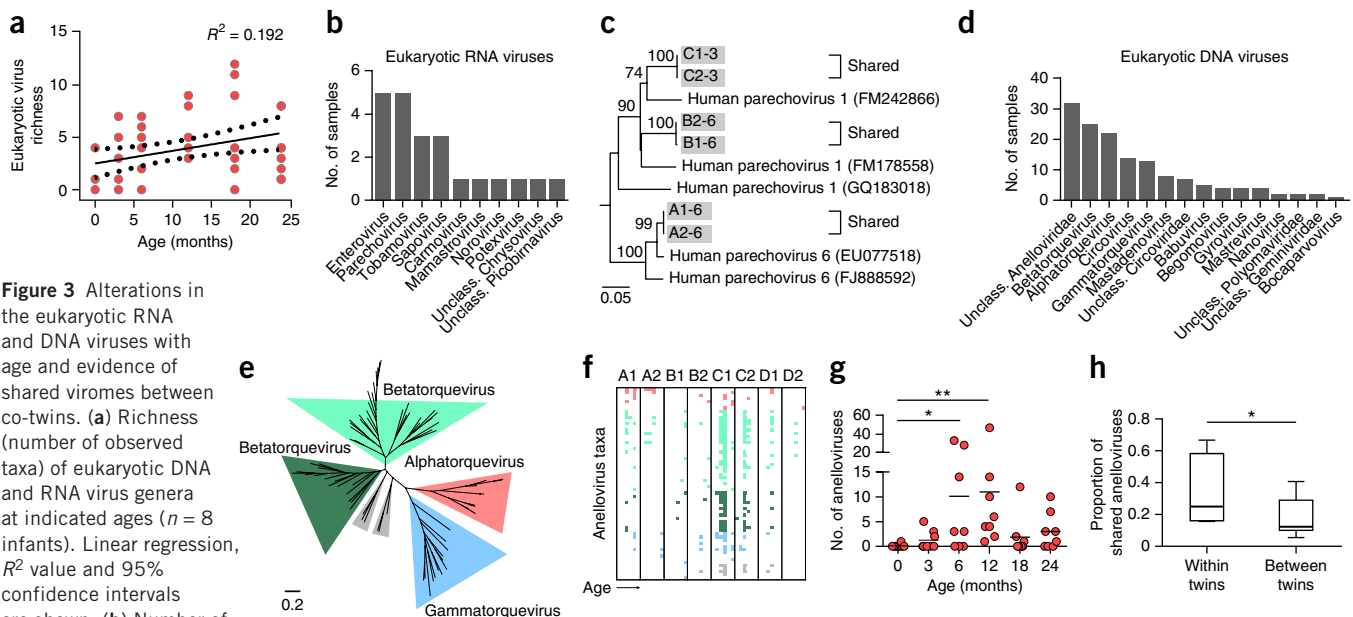
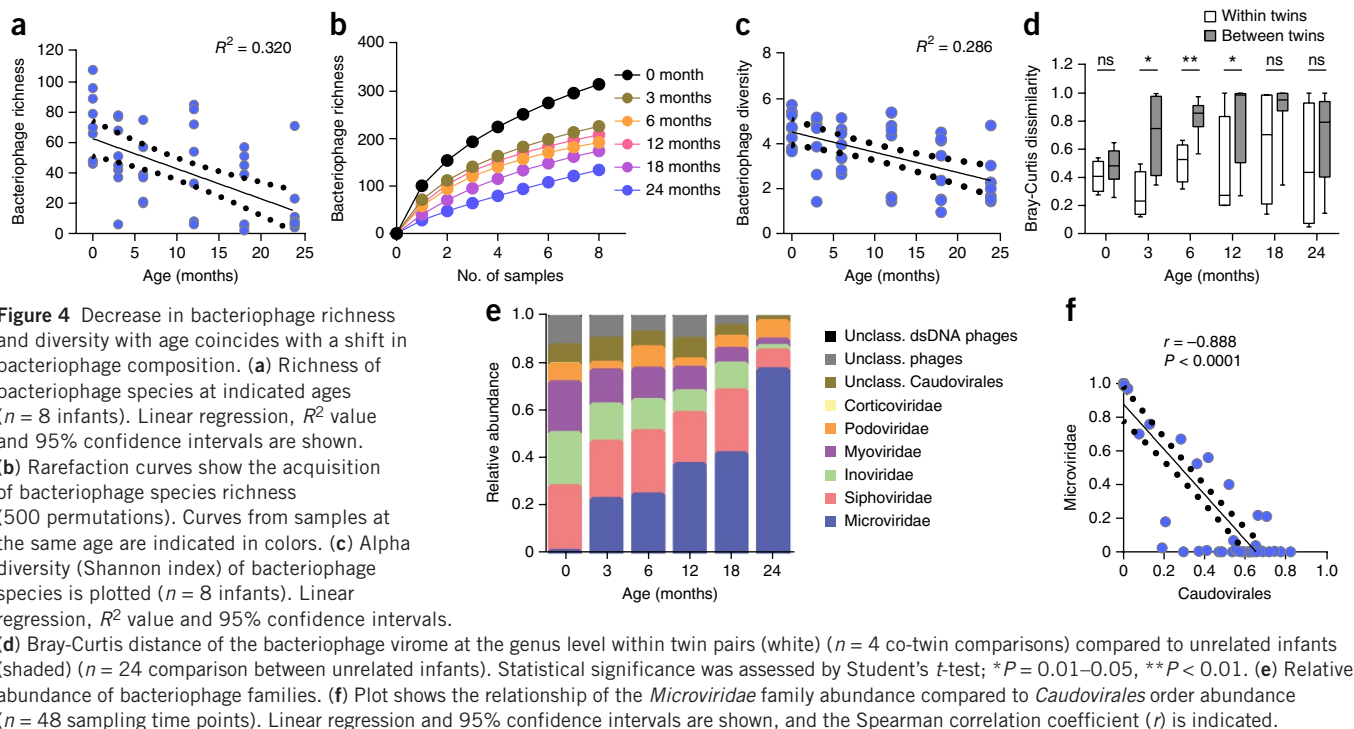


Figure 3 Alterations in the eukaryotic RNA and DNA viruses with age and evidence of shared viromes between co-twins. (a) Richness (number of observed taxa) of eukaryotic DNA and RNA virus genera at indicated ages ($n = 8$ infants). Linear regression, R^2 value and 95% confidence intervals are shown. (b) Number of specimens harboring indicated eukaryotic RNA virus genera. (c) Maximum-likelihood phylogeny of parechovirus sequences. Bootstrap values are indicated on the branches. (d) Number of specimens harboring indicated eukaryotic DNA virus genera. (e) Phylogenetic relationships of 61 anellovirus contigs and 12 reference strains inferred from the ORF1 amino acid alignment, generated by the maximum-likelihood method. Genera are highlighted in indicated colors. (f) Presence-absence heatmap sequencing reads mapped to the anellovirus contigs. Contigs are colored by their genus phylogenetic assignment from e. (g) Richness of anellovirus species at indicated ages ($n = 8$ infants). Statistical significance was assessed by Wilcoxon test (paired, nonparametric); * $P = 0.01$ – 0.05 , ** $P < 0.01$. (h) Comparison of the proportion of shared anellovirus taxa (genome contigs) acquired during the first 2 years of life between co-twins ($n = 4$ co-twin comparisons) and unrelated infants ($n = 24$ comparison between unrelated infants). Statistical significance was assessed by Student's t -test; * $P < 0.05$.



eukaryotic viruses precluded accurate assessment of common ecological parameters (including diversity measurements and rarefaction). To determine whether co-twins harbored the same virus strains at the same time points, we analyzed sequences of parechoviruses, one of the most prevalent RNA viruses detected in our study, by assembling the reads and mapping the resulting contigs to the human parechovirus genome (Supplementary Fig. 3a). Phylogenetic analyses demonstrated that infants within a twin pair shared nearly identical strains of parechovirus (>99.9% nucleotide identity), but different twin pairs harbored distinct strains (Fig. 3c). Consistent with this finding, strain-identical infection in co-twins was also observed for enterovirus (>99.6% nucleotide identity), the other highly prevalent eukaryotic RNA virus (Supplementary Fig. 3b). Although these findings indicate that infants within a twin pair are frequently infected with the same virus, we also observed instances in

which the virus was detected in only one twin but not in the other. For example, human parechovirus reads were detected in infant A1 (3 months) but not in co-twin A2. To determine whether the observed discordance might arise from sensitivity limitations of the sequencing or from differences in the viral load, we screened all the samples with a quantitative RT-PCR (qRT-PCR) assay and measured the number of human parechovirus viral copies. All five sequencing-positive samples were also positive by the qRT-PCR assay (Supplementary Fig. 3c). The three additional 'sequencing-negative' samples that were positive for human parechovirus by qRT-PCR had very low viral loads (18–730 viral copies/15 mg stool) (Supplementary Fig. 3c). Finally, RT-PCR and amplicon sequencing verified that 'discordant' infant A2 harbored the same human parechovirus isolate as its co-twin A1 (Fig. 3c). Thus, the qRT-PCR results independently validated the presence of the viruses

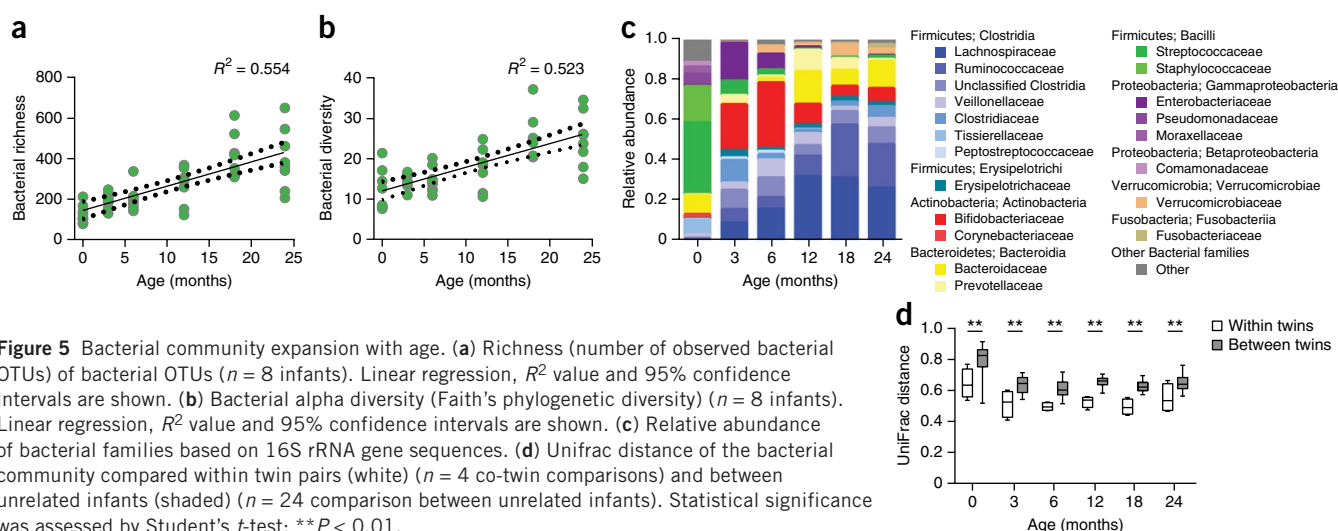
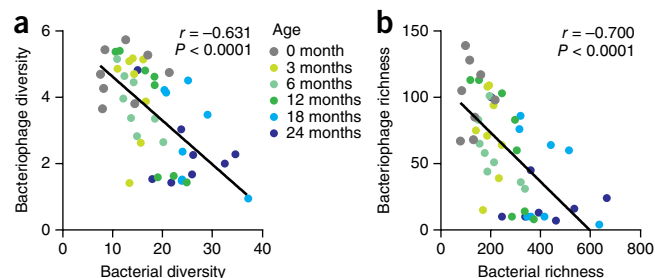


Figure 6 Inverse relationships between bacteriophages and bacteria. (a) Correlation between bacteriophage diversity and bacterial diversity ($n = 48$ sampling time points). Line indicates linear regression, and the Spearman correlation coefficient is shown. Color spectrum indicates age progression from 0–24 months. (b) Correlation plot between bacteriophage richness and bacterial richness ($n = 48$ sampling time points). Line indicates linear regression, and the Spearman correlation coefficient (r) is indicated. Color spectrum indicates age progression from 0 to 24 months.



identified by deep sequencing and further confirm that co-twins shared similar viromes.

Anelloviruses were the most prevalent eukaryotic DNA virus family detected (Fig. 3d). Almost all anelloviruses were previously unknown and highly divergent from previously described anelloviruses (Fig. 3e and Supplementary Fig. 3d). As anellovirus load has previously been associated with changes in host immune status^{20–22}, we hypothesized that changes in infant immunity (for instance, waning of maternal antibodies) may be reflected by changes in anellovirus prevalence or abundance. Because of the large number of novel anellovirus reads detected, we curated a set of unique anellovirus contigs (shared <95% nucleotide identity) to functionally serve as ‘reference genomes’. We then mapped sequencing reads from each specimen to the anellovirus reference contigs to determine the prevalence of each contig (Fig. 3f and Supplementary Fig. 3e). This approach yielded 98.6% concordance with PCR assays designed to detect three specific anellovirus contigs (Supplementary Fig. 3f). Anelloviruses were rarely detected earlier than 3 months of age, but soon increased significantly ($P < 0.05$), peaking at 6–12 months of age (Fig. 3g). Notably, one infant (C1) harbored at least 47 anellovirus species at 12 months of age. Moreover, co-twins shared a higher proportion of anelloviruses than did unrelated infants (Fig. 3h). Further, in some instances, the same anelloviruses could be detected from stools from the same infant that were collected up to 12 months apart (Fig. 3f), suggesting either persistence or a stable source of recurrent infection.

Bacteriophage community contracts and shifts in composition with age

Although DNA bacteriophages were detected in all samples, analysis of the SIA-generated data did not yield any RNA phages. In the absence of RNA phage, we chose to focus the subsequent bacteriophage analyses on the MDA-generated data so that our findings could be comparable to that of other bacteriophage virome studies that have used MDA^{17,26,29}. In contrast to the eukaryotic virome, bacteriophage richness was greatest in the earliest in life specimens at 0 months and decreased with age (Fig. 4a, Wilcoxon test $P < 0.01$). Richness rarefaction curves demonstrated that the rate of bacteriophage species accumulation indeed decreased with age, suggesting that the decrease in richness was unlikely to be attributable to sampling bias (Fig. 4b). Likewise, bacteriophage diversity decreased with age (Fig. 4c, Wilcoxon test $P < 0.01$). The interpersonal variation in the bacteriophage virome was lower (that is, the virome was more similar) between co-twins than between unrelated infants (Fig. 4d). The most abundant bacteriophages were from the *Caudovirales* order (*Siphoviridae*, *Inoviridae*, *Myoviridae* and *Podoviridae* families) and *Microviridae* family, consistent with other studies^{26,29,33}. However, there was a marked shift in the community composition toward an increased relative abundance of *Microviridae* bacteriophages by 24 months of age (Fig. 4e and Supplementary Fig. 4a). This shift in *Microviridae* abundance was also seen in the SIA data, indicating that it was not an artifact of method bias (Supplementary Fig. 4b).

An increase in *Microviridae* species richness was also observed, indicating that this expansion was not driven by a particular species (Supplementary Fig. 4c,d). The relative abundance of *Caudovirales* was inversely correlated with that of *Microviridae* (Fig. 4f). crAssphage, a globally ubiquitous bacteriophage⁴¹, was detected in only one specimen (infant A2, 24 month; 50,355 reads), suggesting that crAssphage is not acquired early in life. Thus, early infant development was marked by a contraction of bacteriophage community richness and diversity, accompanied by a shift toward a predominantly *Microviridae* composition.

Bacterial microbiome changes in infants

The ecological signatures of a healthy intestinal bacterial microbiota during early infancy have been characterized by increasing richness and diversity as bacterial populations mature into a more stable ‘adult-like’ population by 2–3 years of age^{8,9,11}. Given the dramatic changes we observed in the bacteriophage virome, we sought to understand whether these changes correlated with changes in the bacterial microbiome. Therefore, we performed bacterial 16S rRNA bacterial gene sequencing to generate an average of 67,569 reads per sample (s.d. $\pm 39,862$ reads). Quality-filtered sequence reads were clustered into operational taxonomic units (OTUs) at a 97% identity threshold. Principal-coordinate analyses of unweighted UniFrac distance matrices indicated that variation in the bacterial community was associated with age (Supplementary Fig. 5a,b). Consistent with other studies^{8–10}, bacterial richness (Fig. 5a, Wilcoxon test $P < 0.001$ and Supplementary Fig. 5c) and diversity (Fig. 5b, Wilcoxon test $P < 0.001$) also increased with age. Overall, we identified increasingly abundant *Clostridia* (*Firmicutes*) (Fig. 5c and Supplementary Fig. 5d). This was preceded by the predominance of *Bacilli* (*Firmicutes*) OTUs at 0 months, an increase in *Gammaproteobacteria* (*Proteobacteria*) and *Actinobacteria* (*Actinobacteria*) abundance at 3 and 6 months, and an increase in *Bacteroidia* (*Bacteroidetes*) abundance at 12, 18 and 24 months. The interindividual variation between co-twins was less than between unrelated infants (Fig. 5d). Hence, the bacterial microbiota of infants in this study was consistent with the expected trajectory of changes previously observed^{8–10}.

Predator-prey-like bacteriophage and bacteria relationships

Although bacteriophage-bacteria relationships in oceans display predator-prey dynamics^{30,31}, both bacteriophage and bacterial populations are relatively stable in the adult intestine^{17,26}. However, this relationship has yet to be defined in infants. In our cohort, bacteriophage diversity was inversely correlated with bacterial diversity (Fig. 6a). By examining temporal trends, we found that the microbiome shifted from a high bacteriophage–low bacterial diversity community at 0 months toward a low bacteriophage–high bacterial diversity community by 24 months of age. Consistent with this finding, bacteriophage and bacterial richness were inversely correlated in

an age-dependent manner (Fig. 6b). Further reflecting these ecological trends, correlations between specific genera of bacteriophages and bacteria, calculated using a linear mixed model, were dominated by negative correlations (Supplementary Fig. 6). Thus, the infant virome and bacterial microbiome evolves in a dynamic trajectory during the early years of life.

DISCUSSION

Interactions among the intestinal microbiota influence host physiology, development and immunity. Assembly of the infant intestinal bacterial microbiome likely ordains an adult microbiome that will have long-term implications for aspects of host phenotype such as obesity, inflammatory bowel disease and food allergies^{4,14}. However, much less is known about the way the virome develops in the infant, its impact on the bacterial microbiome or its role in human health. Here, we longitudinally defined the complete virome and bacterial communities of 8 infants (4 co-twin pairs) and uncover the microbial milestones of healthy infant virome development. The twin study design enabled us to determine that the infant microbiome (virome and bacterial community) is more similar between co-twins than between unrelated infants. This result contrasts with a study of adult co-twins in which the DNA virome was unique to each individual and twins were not more similar to each other than were unrelated individuals¹⁷. One possible interpretation is that although ‘twin-ness’ matters during infancy, this may reflect the fact that environmental exposures are the primary drivers of virome composition, as the infant twins generally still share a common environment. This was evidenced by the detection in co-twins of near-identical strains of eukaryotic viruses when sampled at the same time points.

Anelloviruses have been proposed to serve as biomarkers of functional immunocompetence because changes in anellovirus load in the serum have been associated with immunosuppression levels in transplant recipients^{20,21}. Additionally, anelloviruses have been frequently detected by PCR in the serum of infants⁴². We observed an expansion of anellovirus richness in the gut at 6 and 12 months of age; most of these anelloviruses were highly divergent from known anelloviruses, underscoring the value of using unbiased deep-sequencing approaches to systematically define the virome, as they may be difficult to detect using PCR assays. We speculate that this expansion could be the result of lowered immune state, as it coincides with the nadir of human IgG as maternal antibodies wane.

To date, no evidence indicates that “kill-the-winner”⁴³ dynamics occur in the human intestinal microbiota, whereby a peak in the bacterial (prey) population precedes the increase in bacteriophages (predator), which subsequently decreases bacterial (prey) populations. Although this is the most commonly recognized aspect of the classical Lotka-Volterra “predator-prey” model, the model also describes the reciprocal relationship whereby limited prey diversity controls predator abundance (that is, predator peaks precede prey peaks, also referred to as a reversed predator-prey cycle)^{32,44}. Our study (Fig. 6) suggests that bacteriophage-bacteria interactions in early infant development begin with the latter dynamics of the Lotka-Volterra model³². We posit that bacteriophage diversity is high at birth (Fig. 4, 0 month), but the bacteriophage population is unsustainable because of low bacterial colonization density (Fig. 5, 0 month). This leads to a contraction in the bacteriophage virome (Fig. 4), thereby relieving the predatory pressure on the bacterial community, allowing it to establish and colonize the gut (Fig. 5). In turn, this drives a shift in the bacteriophage composition (including increased *Microviridae* abundance) that has been selected for in the newly

established bacterial community. In a 2.5-year longitudinal study of a single adult, *Microviridae* were the predominant bacteriophage taxon present²⁶, raising a question about how and when the dynamic microbial state during infancy transitions into the stable community that has been reported in adults^{9,17,26}. Additionally, although our study is unable to address the source of the early bacteriophage diversity, our data raise the possibility of vertical or prepartum transmission (the median day of first sampling was day 2.6 of life). Nonetheless, our data will serve as a reference for healthy infant microbiota development. Although bacteriophage contraction associated with healthy development at birth might be a generalizable phenotype, it is possible that the specific founder bacteriophage composition (identities) might differ on the basis of external factors (such as geography and diet^{9,28,36}).

It has been well described that diet, antibiotics and mode of delivery influence the bacterial microbiome^{8,11–15,28}. Although our cohort included individuals who differed in terms of zygosity, breastfeeding status and mode of delivery, the small cohort size precluded assessment of the role of these factors on the microbiome. Regardless, our current study provides detailed insight into the dynamic interactions between viruses and bacteria in the gut during early development.

METHODS

Methods and any associated references are available in the [online version of the paper](#).

Accession codes. Sequence data have been deposited to the NCBI Sequence Read Archive under BioProject accession number [SRP058399](#).

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

E.S.L., L.R.H. and D.W. conceived and designed the experiments. L.D., I.K.B. and E.S.L. prepared samples for sequencing. E.S.L. and I.K.B. performed PCR experiments. E.S.L., G.Z. and Y.Z. processed and analyzed the sequencing data. P.I.T., B.B.W. and I.M.N. recruited the study participants and managed the metadata. E.S.L., L.R.H., D.W. and P.I.T. wrote and edited the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

Samples. This study was approved by the Human Research Protection Office of Washington University School of Medicine in St. Louis. We obtained consent from the mothers of twin infants to collect fecal specimens from their children monthly through age 3 years³⁹. Fecal specimens were couriered to the laboratory in insulated envelopes containing frozen packs and stored at -80°C until analysis, as described³⁹. Data collected included mode of delivery, medications given to infants, feeding content and episodes of illness (fever/vomiting/diarrhea) by regular interviews of parents or reviews of medical records from the physicians of the twins (**Supplementary Fig. 1a**). The 4 twin pairs (8 infants) in this study were chosen representative of healthy infants who varied in terms of zygosity, breastfeeding status, sex (6 males and 2 females) and mode of delivery. In this study, we defined these infants as 'healthy' as they had no apparent underlying genetic or chronic disorders. There were no other exclusion criteria. The infants had episodes of acute illness (**Supplementary Fig. 1a**). The nomenclature used to label specimens in the study begins with the infant twin pair designation followed by the age (for example, A2-24 refers to a specimen from the second infant in twin pair "A" collected at 24 months of age). For the purpose of clarity in the manuscript, the age of life was defined as 0 months (avg. 2.6 days, s.d. ± 1.1 days), 3 months (avg. 98.0 days, s.d. ± 2.7 days), 6 months (avg. 545.0 days, s.d. ± 17.4 days) and 24 months (avg. 718.5 days, s.d. ± 11.6 days).

Virome sequencing. Fecal specimens (approximately 200 mg) were diluted in phosphate-buffered saline (PBS) at a 1:6 ratio and filtered through a 0.45- μm -pore-size membrane. Total nucleic acid was extracted from the filtrate on the COBAS Ampliprep instrument (Roche) according to the manufacturer's recommendation. Samples were randomized using a random number generator (to define grouping of samples per sequencing run) and then subjected to the following amplification methods. The sequence independent DNA and RNA amplification (SIA) was performed on the total nucleic acid with primers consisting of a base-balanced 16 nucleotide (nt) specific sequence upstream of a random 15-mer (15 Ns) for random priming as previously described³⁵, and used for Nextera DNA library construction (Illumina). For multiple displacement amplification (MDA), total nucleic acid was amplified with phi29 polymerase (GenomiPhi V2 kit, GE Healthcare) according to the manufacturer's instructions and used for Nextera DNA library construction (Illumina). Libraries were purified and size-selected using Agencourt Ampure XP beads (Beckman-Coulter), followed by quantification using a 2100 Bioanalyzer (Agilent Technologies). Multiplexed SIA libraries were pooled and sequenced separately from multiplexed MDA libraries. One SIA sample (C2-0) failed library construction and was not sequenced. Additionally, to evaluate the level of specimen cross-contamination that might occur after library construction (for example, mixed clusters that lead to index misidentification⁴⁵, bioinformatic demultiplexing error, etc.), a uniquely indexed library of cDNA derived from the nematode Orsay virus RNA1 segment⁴⁶ was included in the pool for each sequencing run. Hence, 24 libraries and 1 Orsay virus control library were pooled at equimolar per sequencing run and sequenced on the Illumina MiSeq platform at the Center for Genome Sciences & Systems Biology at Washington University (total of 4 MiSeq sequencing runs, 2×250 paired-end reads, MiSeq v2 reagent kit).

Virome sequence processing. Investigators were blinded to the group allocation (i.e., age, twin pair, time point) during processing of virome sequences up to taxonomic assignment. Sequencing reads were demultiplexed and adaptor sequences were trimmed. Overlapping reads were joined using fastq-join in the ea-utils package⁴⁷. Low quality nucleotides were trimmed and discarded at a quality filter of Q30 Phred quality score. Candidate viral reads were identified by querying against a customized virus database. The customized virus database is comprised of all sequences with the "Viruses" superkingdom taxonomic classification from the publicly available NCBI NT and NR database (downloaded on 7 November 2013). CD-HIT⁴⁸ was used to minimize sequence redundancy (98% identity over 98% of the sequence length), resulting in a customized viral NT database of 449,469 sequences and a viral NR database of 621,095 sequences. The customized viral databases for this study can be downloaded at the following: (viral NT) http://pathology.wustl.edu/virusseeker/data/VirusDBNT_20131107_ID98.tgz; (viral NR) http://pathology.wustl.edu/virusseeker/data/VirusDBNR_20131107_ID98.tgz.

Sequencing reads were queried against the customized viral database sequentially using BLASTn (e-value cutoff 1E^{-10}), followed by BLASTx (e-value cutoff 1E^{-3}). False positive viral sequences were filtered by sequentially querying the candidate viral reads against the NCBI NT database using MegaBLAST (e-value cutoff 1E^{-10}), BLASTn (e-value cutoff 1E^{-10}), and the NCBI NR database using BLASTx (e-value cutoff 1E^{-3}) to remove sequences that have a top BLAST hit corresponding to a nonviral sequence (for example, human, fungal, etc.) as previously described⁴⁹. The taxonomic assignment for sequencing reads was determined by the taxonomy ID of the top BLAST result. Family and genus taxonomic assignments were parsed from the BLAST output using in-house perl and python scripts. Bacteriophage species taxonomic assignment was determined using the lowest-common-ancestor algorithm implemented in Megan (version 5.8.6) (ref. 50) with the following parameters: Min Support: 1, Min Score: 40.0, Max Expected: 0.01, Top Percent: 10.0, Min-Complexity filter: 0.44. Due to the presence of low complexity/repetitive regions in the reads, the following false-positive virus family taxonomic assignments were omitted from the analyses: Herpesviridae (3 reads), Mimiviridae (1 read) and Phycodnaviridae (2 reads). To assess for specimen cross-contamination, we evaluated each demultiplexed data set for the presence of reads that map to the control Orsay virus that was pooled into each MiSeq sequencing run as described above. The average number of Orsay virus sequencing reads obtained from the control library itself was 463,315 reads (s.d. $\pm 106,337$ reads). None of the 48 infant specimens yielded any Orsay virus sequencing reads.

Virome analysis. Sequencing reads generated by the SIA and MDA methods were rarefied (subsampling without replacement) to 200,000 reads per sample method (5 iterations) based on the sequencing depth (**Supplementary Fig. 1b**). The number of reads detected for a given viral taxon was plotted as a heatmap. Consistent results were obtained in the analyses across all iterations. Thus, results obtained from a representative iteration are shown. To define the global virome composition, we merged the rarefied SIA and MDA data and plotted it as an unweighted (presence/absence) heatmap. This merged data was used in analyses of the global virome (eukaryotic viruses and bacteriophages). The prevalence of eukaryotic viruses was inadequate to accurately assess most common ecological measurements (for example, diversity measurements, rarefaction) in a standalone analysis of the eukaryotic virome community. To analyze the bacteriophage virome community, we used the MDA data both because it had a better representation of DNA viruses (**Fig. 1b**) and so as to allow our findings to be comparable to other bacteriophage virome studies which historically have used MDA^{17,26,29}. Ecological analyses including richness and diversity measurements (Shannon index, Bray-Curtis dissimilarity), agglomerative hierarchical clustering and rarefaction curve analyses were performed using the *vegan* R package⁵¹. Rarefaction curves were performed using 500 permutations. Principal-coordinate analyses (PCoA) was performed using Emperor⁵².

To examine inter-individual virus isolates, sequencing reads were mapped to reference genomes (human parechovirus 1 (FM178558), human enterovirus B (NC_001472) and crAssphage (JQ995537)) using bowtie 2 and geneious^{53,54}. Maximum-likelihood (ML) phylogenetic trees were constructed with PhyML (version 3.00)⁵⁵ using appropriate evolution models as assessed by jModelTest2 (version 2.1.6) and ProtTest (version 2.4) accordingly^{56,57}. Support for ML trees was assessed by 1,000 nonparametric bootstraps, and analyses were performed at least twice.

Human parechovirus analysis. A previously described pan-parechovirus TaqMan RT-PCR assay targeting conserved sequences in the 5' UTR region of the genome was used to screen all samples for parechovirus⁵⁸. The following primers were used: AN345F (5'-GTAACASWGGCCTCTGGGSCCAAAAG-3') and AN344R (5'-GGCCCCWGRTCAGATCCAYAGT-3'), with probe AN257 (5'/6-FAM/CCTRYGGGTACCTYCWGGGCATCCTTC/TAMRA/-3'). The qRT-PCR was performed using the TaqMan Fast Virus 1-Step Master Mix (Applied Biosystems). The 20 μL reaction included 5 μL of extracted sample, 10 pmol of each primer, and 5 pmol of probe. The following cycling conditions were used: 50 $^{\circ}\text{C}$ for 5 min, 95 $^{\circ}\text{C}$ for 20 s, 40 cycles of 95 $^{\circ}\text{C}$ for 3 s and 58 $^{\circ}\text{C}$ for 30 s. To generate a standard curve for this assay, *in vitro* transcribed RNA was generated from a plasmid containing the region of interest using MEGAscript

(Ambion) per the manufacturer's protocol. Serial dilutions of the *in vitro* transcribed RNA from 5×10^6 to 5 copies were used to generate a standard curve and a limit of detection of 5 copies was defined. Samples were tested in a 96-well plate format with 5 water-only negative controls. All 5 negative controls were negative for parechovirus. We sought to obtain the orthologous 3D region from parechovirus-positive samples in order to perform a phylogenetic comparison. 3' RACE was performed with ThermoScript reverse transcriptase (Life Technologies) using an oligo (dT)₂₀ primer, and subsequently PCR amplified with a primer specific for human parechovirus (5'-CCAGGTTAACAATGAAGTATGGCAG-3'). Although human parechovirus was detected in specimens D1-12 and D2-12 by the Taqman assay (at low viral copies), we were unable to amplify the human parechovirus from these specimens by 3' RACE and RT-PCR.

Anellovirus analysis. Many of the anellovirus sequences shared limited identity to known anelloviruses, suggesting that they were highly divergent. Therefore, we first sought to curate the anellovirus genome sequences. Contigs were assembled *de novo* from all QC-filtered reads using Newbler (version 2.8) (ref. 59) and queried against the above customized viral database using BLASTx to identify anellovirus contigs. Anellovirus contigs greater than 500 nt were aligned to reference anellovirus genomes: alphatorquevirus TTV1 (AB008394), TTV-P1C1 (AF298585), TTV SIA109 (FJ426280), TTV8 genotype 22 (AB054647); betatorquevirus TTMV1 TLMV-CBD279 (AB026931), TTV-like TLMV-CLC062 (AB038625), TTV-like TTMV_LY2 (JX134045), TTMV5 TGP96 (AB041962), TTV-like LIL-γ1 (EF538880), TTV-like LIL-γ2 (EF538881), TTMV3 (NC_014088); gammatorquevirus TTMV1 MD1-073 (AB290918), TTMV MDJHem8-2 (AB303557), TTMV MDJN1 (AB303558). Contigs that shared >95% nucleotide identity were combined by taking the consensus. Additionally, to determine the phylogenetic relationships of the anelloviruses, only contigs that encoded ORF1 were used for further analyses. ML phylogenetic trees were constructed from the ORF1 amino acid alignment that included the above anellovirus reference genomes. Support for ML trees (LG + I + G + F) was assessed by 1,000 nonparametric bootstraps. Analyses were performed at least twice.

To determine the prevalence of the anelloviruses bioinformatically, sequencing reads from each sample were mapped to the curated anellovirus genome contigs using bowtie 2 and SAMtools^{53,60}. We evaluated the concordance of the *in silico* prevalence analysis with PCR assays for three curated anelloviruses: an alphatorquevirus anellovirus Contig2355 (forward primer 5'-GTAGCCAGAATAAGAACTATGCC-3', reverse primer 5'-TACTGTCTAAACCTGGAAGTTGC-3'); a betatorquevirus anellovirus Contig2737 (forward primer 5'-TCCAAGAGACTTTAAACAGGCC-3', reverse primer 5'-GGAAGCTCTGGATTGTCCATC-3'); a gammatorquevirus anellovirus Contig2393 (forward primer 5'-CTGATGTAGATGATGGACATGGC-3', reverse primer 5'-CATGAGCTTTGTTGCAGAAAGTC-3'). These three anellovirus contigs were chosen based on the following criteria: (1) a representative of each genus (alpha, beta, and gamma) was selected, and (2) based on sequence data, the contigs were either present in multiple time points from an infant or detected across multiple infants. PCR was performed with Taq DNA polymerase (Life Technologies) under the following cycling conditions: 95 °C for 5 min, 40 cycles of 95 °C for 30 s, 58 °C for 30 s, 72 °C for 23 s, followed by 72 °C for 10 min. Products were visualized by electrophoresis using 2% agarose gels.

Bacterial 16S rRNA gene sequencing. Nucleic acid was extracted from fecal specimens that were disrupted by bead beating as described previously²⁹. PCR was performed using Golay-barcoded primers specific for the V4 region (F515/R806) as previously described²⁹. Equimolar libraries were pooled and sequenced using an Illumina MiSeq sequencer (2 × 250 paired-end reads, MiSeq v2 reagent kit) at the Center for Genome Sciences & Systems Biology at Washington University. 4 specimens (A1-0, A2-0, D1-0, D2-0) yielded insufficient reads (<10,000 reads). Hence, we repeated the 16S PCR for these 4 specimens at 40 PCR cycles and re-sequenced their libraries in a subsequent MiSeq sequencing run.

Bacterial 16S rRNA gene analysis. 16S analysis was performed with QIIME (Quantitative Insights Into Microbial Ecology, version 1.8.0) (ref. 61). Sequences were quality filtered at Q20 Phred quality score and demultiplexed. Sequences were assigned to closed reference operational taxonomic units (OTUs) at a 97%

identity threshold using the Greengenes database (version 13.8) (ref. 62). Investigators were blinded to the group allocation (i.e., age, twin pair, time point) during processing of 16S rRNA gene sequences up to OTU taxonomic assignment. To account for inter-sample sequencing depth variability, all samples were rarefied to 10,000 reads per sample (10 iterations), which exceed the generally accepted minimum sequencing depth previously described⁶³. Consistent results were obtained in the 16S analyses across all iterations. Thus, results obtained from a representative iteration are shown. Alpha diversity (Faith's phylogenetic diversity), OTU richness and UniFrac distance were calculated using QIIME. Rarefaction curves were performed using the *vegan* R package using 500 permutations. PCoA plots were visualized using Emperor.

Correlation network. A linear mixed model was used to investigate the relative abundance changes of bacteria and bacteriophages over time. A linear mixed model takes into account repeated measurements from the same subjects at different time points. In the model, the relative abundance of bacteria and phage were log transformed, sample collection time was designated a fixed effect and subjects were designated as random effects. *P* values from multiple comparisons were corrected using false discovery rate (*q* value). A *q* value less than 0.05 was considered statistically significant. Bacteria and phage correlation was plotted using Cytoscape. The analyses were performed in R version 3.1.2.

Statistics. Student's *t*-test (two-sided) was performed with SAS. Normal distribution and equal variances between the groups were verified. Wilcoxon test (paired, nonparametric) was applied to compare the eukaryotic virus richness (Fig. 3a), bacteriophage richness (Fig. 4a), bacteriophage diversity (Fig. 4c), bacterial richness (Fig. 5a) and bacterial diversity (Fig. 5b) between matched samples at 0 months compared to 24 months. Wilcoxon test, linear regression and Spearman correlation was performed with GraphPad Prism. *P* values and 95% confidence intervals are shown accordingly.

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