Comparative Metagenomics and Population Dynamics of the Gut Microbiota in Mother and Infant

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Accepted: 23 December 2009 Associate editor: Emmanuelle Lerat

Abstract

Colonization of the gastrointestinal tract (GIT) of human infants with a suitable microbial community is essential for numerous aspects of health, but the progression of events by which this microbiota becomes established is poorly understood. Here, we investigate two previously unexplored areas of microbiota development in infants: the deployment of functional capabilities at the community level and the population genetics of its most abundant genera. To assess the progression of the infant microbiota toward an adult-like state and to evaluate the contribution of maternal GIT bacteria to the infant gut, we compare the infant’s microbiota with that of the mother at 1 and 11 months after delivery. These comparisons reveal that the infant’s microbiota rapidly acquires and maintains the range of gene functions present in the mother, without replicating the phylogenetic composition of her microbiota. Microdiversity analyses for Bacteroides and Bifidobacterium, two of the main microbiota constituents, reveal that by 11 months, the phylotypes detected in the infant are distinct from those in the mother, although the maternal Bacteroides phylotypes were transiently present at 1 month of age. The configuration of genetic variants within these genera reveals populations far from equilibrium and likely to be undergoing rapid growth, consistent with recent population turnovers. Such compositional turnovers and the associated loss of maternal phylotypes should limit the potential for long-term coadaptation between specific bacterial and host genotypes.

Key words: Bacteroides, Bifidobacterium, gut microbiota, community genomics, bacterial population genetics.

Introduction

The gastrointestinal tract (GIT) is host to the most abundant and diverse bacterial community in humans, containing, in adults, on the order of $10^{14}$ bacterial cells compared with $10^{13}$ human cells for the entire body. This complex microbiota plays essential roles for gut maturation, food digestion, modulation of the immune system, and protection from pathogens (Savage 1977; Stark and Lee 1982; Cebra 1999; Mackie et al. 1999; Hooper et al. 2001; Favier et al. 2002; Fanaro et al. 2003; Schumann et al. 2005; Adlerberth et al. 2006; Ley, Peterson, and Gordon 2006; Ley, Turnbaugh, et al. 2006; O’Mahony et al. 2008). At birth, the GIT is germ free, but it rapidly enters an extensive and a complex process of colonization by a variety of microbes. Several studies have monitored this process at different stages by means of 16S ribosomal RNA (rRNA) analyses (Favier et al. 2002; Hopkins et al. 2005; Park et al. 2005; Penders et al. 2006; Palmer et al. 2007) and have shown that it is far
from a strict microbial succession. Rather, GIT colonization is highly variable among individuals and is influenced by numerous factors, including mode of birth, feeding habits, health status, and sanitary conditions (Fanaro et al. 2003; Adlerberth et al. 2006; Penders et al. 2006; Palmer et al. 2007). However, 16S rRNA analyses can only appraise the phylogenetic composition of a sample and provide no direct information about its functional capabilities. An additional issue of fundamental importance remains unexplored: no studies have addressed the population dynamics of individual bacterial species as the infant gut is colonized. However, an appreciation of the amount, pattern, origin, and persistence of population-level genetic variation within an infant’s GIT is crucial for understanding the evolutionary forces at work during the process of colonization. In particular, a specific aspect of population dynamics—the degree to which an infant’s microbiota is vertically inherited from the maternal GIT—will determine the potential for natural selection to foster long-term bacterial adaptations to specific host genotypes.

Here, we address these issues by focusing on the GIT microbiota in a single mother–infant pair, revealing not only the coding capabilities and phylogenetic composition of these bacterial communities but also the microdiversity, population biology, and degree of maternal inheritance within their main constituents. The infant in our analysis was a healthy male, vaginally delivered at full term in an urban hospital in Tucson, AZ. He was exclusively breast-fed for 5 months, after which solid food was introduced to his diet, although he continued to ingest breast milk until 2 years of age. We analyzed the GIT microbiota of this infant and that of his mother at two time points during the first year of life, at 1 month (I-1m and M-1m) and 11 months (I-11m and M-11m) after birth. These two stages represent well-differentiated phases of microbiota development in the infant (Cooperstock and Zead 1983). By 1 month, the initial phase of rapid acquisition of microbes after birth is thought to be over and, in exclusively breast-fed infants, to have produced a breast milk–adapted microbiota that will remain until the introduction of solid foods (Cooperstock and Zead 1983; Mackie et al. 1999). By 11 months, the baby has been exposed to a complex diet for a significant period of time and should be reaching an adult-like microbiota composition (Stark and Lee 1982; Mackie et al. 1999; Favier et al. 2002; Fanaro et al. 2003; Park et al. 2005; Penders et al. 2006; Palmer et al. 2007). Our analyses revealed that these two stages contained bacterial communities that were largely distinct in phylogenetic composition but remarkably similar and adult like in their global functional capabilities. Overall, this sampling strategy indicated that the GIT microbiota remains very dynamic throughout the first year of life, with highly diverse and actively growing bacterial populations, and that maternal GIT phylotypes can be transmitted to the infant but need not persist in the long term.

**Materials and Methods**

**Sample Collection** We obtained fecal samples from a healthy mother–infant pair at 1 and 11 months after delivery. Samples were collected at the University of Arizona, with informed written consent from the infant’s parents, using protocols approved by the institutional review boards of the Lawrence Berkeley National Laboratory and the University of Arizona.

**Preparation of High–Molecular Weight DNA and Fosmid Library Construction** In order to capture extensive genomic fragments from the bacterial species in the GIT microbiota, we generated fosmid libraries from high–molecular weight bacterial DNA isolated directly from the fecal samples. Fosmid libraries enable the elucidation of both the coding capabilities and the phylogenetic positions of the members of a given bacterial community while providing the possibility of recovering long contiguous sequences from interesting clones. To obtain high–molecular weight bacterial DNA, fresh fecal samples were harvested and immediately processed. In short, 5 g of sample was suspended in 50 ml phosphate buffer saline (PBS) and subjected to a series of filtration steps to eliminate particulate matter and large eukaryotic cells. Sequential filtrations were performed with a Micro Filtration System (Millipore) using 100-, 20-, and 11-μm nylon filters and were followed by centrifugation at 250 × g to remove eukaryotic nuclei. Bacterial cells were then pelleted, washed, and resuspended in 5 ml PBS. Visual inspection under the microscope revealed a large variety of bacterial cell types and no trace of eukaryotic cells. Bacterial cells were embedded in agarose plugs and lysed with a variety of enzymes (lysozyme, N-acetylmuramidase, and achr-romopeptidase) capable of digesting cell walls from phylogenetically diverse bacteria plus protease K and detergent (Hayashi et al. 2002). Agarose plugs containing bacterial DNA were washed exhaustively and stored at 4°C in 50 mM ethylenediaminetetraacetic acid. For fosmid library construction, DNA was extracted from the agarose plugs and mechanically fragmented by hydroshearing. Forty-kilobase DNA fragments were separated by pulsed-field gel electrophoresis and cloned into pCC1Fos (Epicentre Corp.).

**Sequencing and Analysis of Fosmid Ends** In order to characterize the phylogenetic composition and gene content of the obtained libraries, inserts from 1,536 randomly selected fosmids from each library were sequenced at both ends. All sequencing reactions were performed by the Sanger method using BigDye Terminators in ABI 3730 sequencers at the DOE Joint Genome Institute. Only high-quality sequences (≥Q20), as assessed by Phred (Ewing and Green 1998; Ewing et al. 1998), were retained for analysis after being subjected to LUCY (Chou and Holmes 2001) for quality trimming and vector contamination removal. Assembly was attempted with
Phrap (http://www.phrap.org), but due to the low percentage of reads assembling into contigs, all subsequent analyses were performed on unassembled collections of sequences for each sample. All sequences were deposited in GenBank (EF990944–EF998850) and IMG/M (Markowitz et al. 2008). Ab initio gene calling was performed in IMG/M using GeneMark (Besemer et al. 2001), and the identified protein-coding genes were further analyzed in IMG/M to provide estimates of the phylogenetic composition and the functional repertoire of each sample. Taxonomic affiliation was registered only for genes with best hits having >90% identity to IMG/M reference isolate genomes as determined by BlastP.

**Statistical Comparison of the Functional Profiles of GIT Samples** The abundances of different gene functions in each sample were compared by means of GIT Samples Statistical Comparison of the Functional Profiles of and annotated with NCBI’s open reading frame finder (Genome Biol. Evol. 2010). GeneMark (Besemer et al. 2001), and the identified protein-coding genes were further analyzed in IMG/M to provide estimates of the phylogenetic composition and the functional repertoire of each sample. Taxonomic affiliation was registered only for genes with best hits having >90% identity to IMG/M reference isolate genomes as determined by BlastP.

**Genus-Specific 16S rRNA Phylogenies** We investigated the extent of variation within *Bacteroides* and *Bifidobacterium*, two of the main constituent genera of the GIT microbiota in the infant and maternal microbial communities. 16S rRNA was polymerase chain reaction (PCR) amplified from the agarose-embedded DNA (above) using primers and reaction conditions determined for each genus. The primer pairs employed were forward Bac32F (5’-AACGCTAGCTACAGGCTT-3’) and reverse Bac708R (5’-CAATCGGATTCTCGTG-3’) for *Bacteroides* (Bernhard and Field 2000) and forward lm26 (5’-GATTCTGGCTCAAGGATGAACG-3’) and reverse lm3 (5’-CGGGTGCTCCATTCTCATG-3’) for *Bifidobacterium* (Kaufmann et al. 1997), yielding PCR products of 677 and 1,418 bp, respectively. PCR products were cloned into pCR4-TOPO (Invitrogen), and 50 randomly selected clones from each library were sequenced from both ends. Sequence reads were vector trimmed, assembled, quality checked, and chimera checked using the software package Genelib (Kirton E, unpublished data). The resulting sequence were aligned in ARB (Ludwig et al. 2004) to a backbone alignment from the Greengenes 16S rRNA database (http://greengenes.lbl.gov) (DeSantis et al. 2006). Alignments were refined manually taking into account rRNA secondary structures. Neighbor-joining trees were produced to identify those sequences in the Greengenes backbone alignment that were most similar to the reads from our samples. These sequences were exported for phylogenetic analysis along with our reads and sequences representing the extent of diversity within each genus analyzed. Prior to phylogenetic reconstruction, alignments were pruned of redundant sequences and poorly aligned regions in Gblocks (Castresana 2000), yielding 39- and 48-sequence alignments for *Bacteroides* and *Bifidobacterium*, respectively. Maximum likelihood phylogenetic analysis with 1,000 bootstrap replicates was performed using Garli (http://www.bio.utexas.edu/faculty/antisense/garli/Garli.html) (Zwickl 2006) with the best-fitting sequence evolution model (general time reversible with invariant sites and gamma distribution for site rate variation) and parameters determined in Modeltest v. 3.7 (Posada and Crandall 2001).

**Population Genetic Analyses** Genealogical relationships among the different haplotypes recovered within the main constituent genera of the GIT microbiota were estimated by statistical parsimony (Templeton et al. 1992), as implemented in the computer program TCS (Clement et al. 2000). Gene diversity (H) was computed as

$$H = n(1 - \sum x_i^2)/n - 1,$$

where $x_i^2$ is the estimated frequency of allele $i$ in a sample of $n$ alleles (Nei 1987). Expected numbers of total haplotypes $E(k)$ were computed according to Ewens (1972) as

$$E(k) = 1 + \theta/(\theta + 1) + \theta/(\theta + 2) + \ldots + \theta/(\theta + n - 1),$$
Table 1
Characterization of Mother and Infant Samples Based on Fosmid-End Sequencing

<table>
<thead>
<tr>
<th></th>
<th>M-1m</th>
<th>M-11m</th>
<th>I-1m</th>
<th>I-11m</th>
</tr>
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<tr>
<td>Analyzed reads</td>
<td>1,981</td>
<td>2,056</td>
<td>1,778</td>
<td>2,092</td>
</tr>
<tr>
<td>Analyzed bases</td>
<td>1,317,583</td>
<td>1,400,487</td>
<td>1,033,600</td>
<td>1,422,416</td>
</tr>
<tr>
<td>Total genes</td>
<td>1,916</td>
<td>2,041</td>
<td>1,522</td>
<td>2,077</td>
</tr>
<tr>
<td>Protein-coding genes</td>
<td>1,891</td>
<td>2,003</td>
<td>1,507</td>
<td>2,045</td>
</tr>
<tr>
<td>Functionally assigned genes(^a)</td>
<td>869</td>
<td>1,218</td>
<td>894</td>
<td>1,152</td>
</tr>
<tr>
<td>COG-assigned genes</td>
<td>719</td>
<td>1,067</td>
<td>767</td>
<td>1,030</td>
</tr>
<tr>
<td>Taxonomically assigned genes</td>
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<td>1,160</td>
<td>1,128</td>
<td>1,434</td>
</tr>
<tr>
<td>Identified genera</td>
<td>22</td>
<td>20</td>
<td>9</td>
<td>16</td>
</tr>
</tbody>
</table>

\(^a\) Functional assignment was performed within IMG/M (Markowitz et al. 2008) with basis on COG (Tatusov et al. 1997), Pfam (Bateman et al. 2004), TIGRfam (Selengut et al. 2007), and KEGG (Kanehisa et al. 2004) classifications.

where \( \theta = H/(1 - H) \). Expected numbers of haplotypes present singly (singletons) \( E(k_1) \) were computed according to Maruyama and Fuerst (1984) as

\[
E(k_1) = \frac{\theta n}{(n + \theta - 1)}.
\]

Data Deposition GenBank (EF990944–EF998850) and (GU362547–GU362642).

Results

Fosmid Libraries and End Sequencing  To characterize and compare the microbial communities present in the infant and the mother at 1 and 11 months after delivery, we generated four fosmid libraries from bacterial DNA isolated directly from fecal samples (50,000 clones each; 8 Gb total). Inserts from 1,536 randomly selected fosmids from each library were sequenced at both ends, yielding a total of 12,288 sequence reads and more than 8.4 Mb of high-quality sequence (Q20) as assessed by Phred (Ewing and Green 1998; Ewing et al. 1998). After further quality trimming and vector contamination removal, the sequences of each sample were independently assembled using phrap (http://www.phrap.org). The majority of sequences (62% on average) could not be incorporated into assemblies, and most of the assembled contigs contained only two reads. But because bacterial genes are compact and intronless, ab initio gene calling in IMG/M indicated that the four libraries contained only two reads. But because bacterial genes are compact and intronless, ab initio gene calling in IMG/M (Markowitz et al. 2008) was able to detect more than 7,500 genes in the unassembled reads of the four samples (table 1).

Similar Distribution of Gene Functions in the GIT Microbiota of Mother and Infant  The functional repertoire within the different GIT samples was appraised within IMG/M with basis on different classifications, including COG (Tatusov et al. 1997), Pfam (Bateman et al. 2004), TIGRfam (Selengut et al. 2007), and KEGG (Kanehisa et al. 2004). Combined, all classifications allowed for functional categorization of 45–60% of the genes detected within each sample, with 38–52% of total genes assigned to COG clusters (table 1). The overall distribution of COG functional classes was remarkably similar in all four libraries (fig. 1A). Carbohydrate and amino acid transport and metabolism were the most common COG categories overall—categories G and E—and together represented 21–27% of the COG assignments per sample. In contrast, lipid transport and metabolism (COG category I) accounted for only 2–3% of these assignments.

To evaluate any potential significance of function abundance differences among samples, we computed D scores for each individual COG as well as D ranks for COG pathways and categories as defined in IMG/M (Markowitz et al. 2008). Pairwise comparisons among samples did not identify significant differences in the distribution of individual COGs, pathways or categories (all \( P > 0.05 \)). In contrast, a PCA of the COG category profiles grouped closely the M-1m, M-11m, and I-11m samples to the exclusion of I-1m (fig. 1B). Inspection of the contribution of the different COG categories to the principal components indicates that it is the presence of the COG category W (Extracellular Structures) in I-1m that separates this sample on the second principal component (PC) axis, whereas the large distance between our samples and the larger adult human gut samples from Gill et al. (2006) (black and gray circles) on the first axis is due to the higher numbers of individual COGs represented within each category in the latter. The Extracellular Structures category is only represented in I-1m by four sequences; these sequences match COG5295, which codes for autotransporter adhesins, secreted bacterial proteins that may exhibit diverse virulence functions. Best hits to reference genomes in IMG indicated that the four I-1m COG5295 sequences were likely encoded by Escherichia coli.

To further evaluate potential differences in functionality between infant and maternal samples, we investigated the distribution of those COGs in the carbohydrate transport and metabolism category (G) that were previously found to be overrepresented in human gut microbiomes (Kurokawa et al. 2007). The human gut overrepresented G COGs include 32 COGs (group I) encoding proteins that could transport or metabolize the different sugars found in human milk, mostly lactose and other oligosaccharides containing glucose, galactose, N-acetylglucosamine, fucose, or sialic acid residues (table S1, Supplementary Material online). In addition, the human gut overrepresented G COGs also include 13 COGs (group II) encoding proteins specialized for the transport and metabolism of carbohydrates of plant origin (table S2, Supplementary Material online). To search for specific differences among our samples potentially related to diet, we specifically analyzed the distribution of human gut overrepresented G COGs belonging to these two groups.

We recovered in at least one of our samples 74% of the human gut overrepresented G COGs, including 13 COGs encoding glycosyl hydrolases and 5 for different components of sugar transporters. The two infant samples
(I-1m and I-11m) contained very similar numbers of group I COGs, 12 and 11, respectively, although only 5 of these were common to both (COG0676, COG1129, COG1263, COG1486, and COG2017). Notably, the two maternal samples (M-1m and M-11m) contained 18 group I COGs each, and 15 of these were in common. In terms of the overall number of reads assigned to group I COGs in each of the four samples (table S1, Supplementary Material online), there were no significant differences as evaluated by IMG/M D rank scores ($P > 0.05$), but some differences were apparent in terms of the presence/absence of specific group I COGs among samples. Six COGs were recovered exclusively from infants and not from the maternal samples, including COG0676, encoding uncharacterized enzymes related to aldose 1-epimerase, and COG1486, encoding $\alpha$-galactosidases/6-phospho-$\beta$-glucosidases. In addition to these two COGs common to both infants, COG1299 (fructose-specific IIC component of the phosphotransferase system) and COG4668 (galactose-1-phosphate uridylytransferase) were unique to I-1m, whereas COG3414 (galactitol-specific IIB component of the phosphotransferase system) and COG3669 ($\alpha$-L-fucosidase) were unique to I-11m. On the other hand, the maternal samples also contained several group I COGs not recovered in the infant samples, including COGs encoding enzymes for the metabolism of galactose (COG0153, COG1874, and COG3345), fructose (COG0205, COG1762, and COG2893), and fucose (COG0738 and COG2407) (table S1, Supplementary Material online).

Among group I COGs, we recovered in our samples three out of four glycosidase-encoding COGs found in a HMO utilization cluster recently discovered in *Bifidobacterium*.
**Different Phylogenetic Compositions across GIT**

**Microbiota Samples** Phylogenetic composition was established through BlastP queries of the identified protein-coding genes. Requiring a BlastP identity threshold to assign genes to a specific taxonomic group resulted in the identification and classification of 72% and 56% of genes in infant and mother samples, respectively. The distribution of these high-scoring BlastP hits revealed that, in spite of the functional similarity already noted, there were large differences in phylogenetic membership among libraries, indicating turnover in the GIT microbiota over time in both infant and mother. Although the total numbers of phyla represented were similarly low in all samples (four to five: Bacteroidetes, Actinobacteria, Proteobacteria, and Firmicutes, plus an occasional Fusobacteria), the presence and frequencies of different genera within these phyla varied substantially. In the 1-month-old infant, only 9 genera were recovered, but by 11 months, the number of genera increased and approached that found in the maternal samples (16 genera in I-11m vs. 22 in M-1m and 20 in M-11m). The microbiota in the 1-month-old infant consisted primarily of Escherichia (45%) and Bacteroides (21%), but by 11 months, these genera had been drastically reduced to 6% and 0.4%, whereas the previously absent Bifidobacterium (38%) and the related actinobacterium Collinsella (22%) dominated the community. During the same time period, the maternal microbiota also exhibited a substantial reduction in Bacteroides (from 48% at 1 month to 12% at 11 months) and the appearance of Bifidobacterium (10%), although in this case, these changes were accompanied by a large increase in enteric genera (3–29%) (fig. 1C).

**Microdiversity within Bacteroides and Bifidobacterium** In that Bacteroides and Bifidobacterium are main constituents of the gastrointestinal microbiota and play basic roles in the metabolism of carbohydrates in the gut, we investigated the extent of variation within these genera in the infant and maternal microbial communities. To this end, we employed 16S ribosomal DNA (rDNA) primers specific to Bacteroides (Bernhard and Field 2000) or Bifidobacterium (Kaufmann et al. 1997) and measured the species (97% sequence identity) and phylotype (99% sequence identity) diversity in each of these genera. The Bacteroides-specific primers amplified the 16S rDNA gene in all samples, but Bifidobacterium primers produced amplicons only in the samples recovered at 11 months after childbirth, in accordance with the results obtained by fosmid-end sequencing (fig. 1C). A maximum likelihood phylogeny of the 48 non-identical Bifidobacterium sequences (i.e., haplotypes) recovered from the 11-month samples resolved the main Bifidobacterium haplotypes into distinct clades according to their source of origin (fig. 2A). The 24 Bifidobacterium sequences from the infant at 11 months after delivery (I-11m) form a single phylotype, most closely related to an oral Bifidobacterium strain and to Bi. adolescentis, whereas the vast majority of haplotypes from the mother at this sampling point (M-11m) form a second phylotype related to Bi. angulatum (two other M-11m sequences reside within the Bi. animalis clade). The presence of distinct phylogenetic lineages in mother and infant suggests that each

**Fig. 2.**—Phylogeny of GIT haplotypes in relation to diversity within the corresponding genus. Phylogenies of the (A) Bifidobacterium and (B) Bacteroides 16S rRNA haplotypes with representative samples from each genus were derived by maximum likelihood with 1,000 bootstrap replicates under a general time-reversible model with invariant sites and gamma-distributed site rate variation. Scale bar represents 0.02 nucleotide substitutions per site. “Mixed” clades contain haplotypes recovered from both the mother and the infant.
independently acquired bifidobacteria from external sources during the period between 1 and 11 months after birth.

The phylogeny of the 39 *Bacteroides* 16S haplotypes obtained across the four libraries portrays a very different pattern of acquisition and transmission (fig. 2B). In contrast to the *Bifidobacterium* tree topology, two of the three *Bacteroides* clades (II and III) are intermixed with sequences from both maternal samples (M-1m and M-11m) and the 1-month infant sample (I-1m), whereas clade I consists only of sequences recovered from the 11-month infant (I-11m). The 19 sequences in clade I form a single phytype related to *Bacteroides vulgatus*. Clades II and III represent distinct but closely related phytypes (98.5% sequence identity), containing 5 and 15 haplotypes, respectively. Thus, the maternal GIT harbored the same two *Bacteroides* phytypes over a period of 10 months, and these phytypes were transmitted from the mother to the infant before 1 month of age. Nonetheless, 10 months later, a new phytype belonging to a different *Bacteroides* species had replaced the original maternal phytypes in the infant GIT.

**Fosmid Sequencing Confirms that Mother and 1-Month-Old Infant Contain the Same Bacteroides Strain** To test if the mother and the 1-month-old infant contained identical strains, we culled our collection of fosmid-end sequences and recovered two *Bacteroides* clones, one from the mother and one from the infant, with extended regions of 100% identity. From these, we generated draft shotgun sequences that provided large contigs with 26 kb of overlapping sequence. Alignment of the 26-kb regions from each host did not reveal a single nucleotide difference, confirming that mother and infant shared the same strain of *Bacteroides*.

**Bacteroides and Bifidobacterium Population Genetics** The population biology and transmission patterns in our mother–infant system were addressed by analyzing the distribution, frequency, and relationships of haplotypes within the *Bifidobacterium* and *Bacteroides* clades (fig. 2A and 8). Genealogical relationships among haplotypes were estimated by statistical parsimony (Templeton et al. 1992; Clement et al. 2000); this approach is required for population-level analyses because, unlike methods aimed at reconstructing lineage phylogenies, it integrates information about haplotype frequencies, allows for the ancestral haplotype to be represented in the sample, and does not force a bifurcating network topology.

In the case of *Bacteroides*, the maternal 1-month sample contained 12 haplotypes, the most common of which were represented by 28 (M-1m.h1, clade III) and 4 (M-1m.h12, clade II) identical sequences. All of the haplotypes in this sample were also recovered at nearly identical frequencies 10 months later in the maternal sample taken at 11 months after delivery (table 2). In addition, two novel haplotypes were represented by individual sequences in the 11-month maternal sample. In contrast, only three of the maternal haplotypes were recovered in the 1-month-old infant. These haplotypes included the two most prevalent and one of the singletons and were recovered in proportions similar to those observed in the mother (31, 8, and 1 sequences; table 2). Aside from harboring three haplotypes detected in the mother, the 1-month-old infant also contained seven unique singletons. Although the specific haplotypes differed between mother and infant, the total number of haplotypes, the proportion of singletons, and the overall gene diversity were similarly high in both individuals (table 2). Figure 3 presents the genealogy of all the *Bacteroides* haplotypes recovered from the mother and the 1-month-old infant (corresponding to clades II and III in fig. 2B). Irrespective of their sample of origin, all of the singletons and other low-frequency alleles were derived from the most common haplotypes by one or two mutational steps. These changes are unlikely to represent sequencing errors, as average Phred quality scores (Ewing and Green 1998; Ewing et al. 1998) for our quality-trimmed and assembled 16S contigs are approximately Q60, such that the average error is <10⁻⁶. Moreover, the fact that we recover very similar haplotype distributions in M-1m and M-11m, with nearly all the singletons being shared between the two, strongly indicates that the variants detected in the *Bacteroides* populations are real and not significantly affected by errors in sequencing.

In the three new populations present at 11 months (i.e., the mother and infant *Bifidobacterium* phylotypes and the new *Bacteroides* phylotype in the infant), we also detected very large numbers of haplotypes, many of which were singletons derived by one or a few mutations from the most common haplotype (table 2, fig. S1, Supplementary Material online), although in the case of the longer *Bifidobacterium* contigs, some of these changes might represent sequencing errors because most bases were covered by a single sequencing pass. The large numbers of singletons and total haplotypes present in all the GIT populations analyzed suggest that these populations are generally far from genetic equilibrium. Populations at equilibrium have characteristic haplotype frequency configurations that can be predicted based on the overall level of gene diversity (Ewens 1972; Nei and Li 1976; Maruyama and Fuerst 1984). For each of the *Bacteroides* and *Bifidobacterium* populations in our samples, we computed gene diversity indices based on the observed haplotype frequencies and derived the numbers of singletons and total haplotypes that would be expected in populations at equilibrium containing such level of gene diversity. In all cases, the predicted equilibrium frequencies of singletons and of total haplotypes were lower than those observed (table 2). Potential biases in the data introduced by undersampling are not likely to have caused the observed deviations from equilibrium frequencies; Good’s (1953) non-parametric coverage estimator indicates that we likely
recovered 70–80% and 40–50% of the haplotypes present in the *Bacteroides* and *Bifidobacterium* populations, respectively, and, as missed haplotypes would likely be those present at low frequencies, the numbers of haplotypes and singletons we detected are probably underestimated.

**Discussion**

The goal of our study was not to generate an exhaustive enumeration of the bacterial species in the human GIT but to compare the main compositional and functional features of the microbiota at different stages of development. Given the ease of obtaining human fecal samples, compared with the alternative invasive procedures to sample the contents of the GIT, we chose to analyze the bacterial composition of fecal material to characterize the gastrointestinal microbiota of mother and infant, as performed in most previous studies in humans (Zoetendal et al. 1998; Gewolb et al. 1999; Favier et al. 2002; Hayashi et al. 2002; Fanaro et al. 2003; Hopkins et al. 2005; Park et al. 2005; Adlerberth et al. 2006; Gill et al. 2006; Ley, Turnbaugh, et al. 2006; Park et al. 2005; Adlerberth et al. 2006; Gill et al. 2006; Ley, Turnbaugh, et al. 2006; Penders et al. 2006; Kurokawa et al. 2007; Palmer et al. 2007; Turnbaugh et al. 2009). Feces, which comprise 40% bacterial cells, are thought to be most representative of the microbiota in the colonic lumen, although, due to normal mucus excretion, epithelial turnover, and peristaltic movements, they are also likely to contain lumen- and mucosa-associated bacteria originating in other areas of the GIT (Moore and Holdeman 1975; Eckburg et al. 2005). Thus, although feces cannot fully represent the complexity of the GIT microbial ecosystem, they probably recover a substantial proportion of the bacterial species in this environment (Eckburg et al. 2005), and their analysis should

Table 2

<table>
<thead>
<tr>
<th>Persistence of Haplotypes of <em>Bacteroides</em> and <em>Bifidobacterium</em> in Mother and Infant Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>mother samples</td>
</tr>
<tr>
<td><strong>Bacteroides</strong> haplotypes(a)</td>
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<tr>
<td>-----------------</td>
</tr>
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<td><strong>M-1m</strong>.h1 (clade III)</td>
</tr>
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<tr>
<td><strong>M-1m</strong>.h30 (clade II)</td>
</tr>
<tr>
<td><strong>M-1m</strong>.h23 (clade II)</td>
</tr>
<tr>
<td><strong>M-1m</strong>.h5 (clade III)</td>
</tr>
<tr>
<td><strong>M-1m</strong>.h10 (clade III)</td>
</tr>
<tr>
<td><strong>M-1m</strong>.h13 (clade III)</td>
</tr>
<tr>
<td><strong>M-1m</strong>.h19 (clade III)</td>
</tr>
<tr>
<td><strong>M-1m</strong>.h20 (clade III)</td>
</tr>
<tr>
<td><strong>M-1m</strong>.h25 (clade III)</td>
</tr>
<tr>
<td><strong>I-1m</strong>.h441 (clade I)</td>
</tr>
<tr>
<td><strong>I-1m</strong>.h440 (clade I)</td>
</tr>
<tr>
<td><strong>I-1m</strong>.h445 (clade I)</td>
</tr>
<tr>
<td><strong>I-1m</strong>.h433 (clade I)</td>
</tr>
<tr>
<td><strong>I-11m</strong>.h4 (clade I)</td>
</tr>
<tr>
<td><strong>I-11m</strong>.h5 (clade I)</td>
</tr>
<tr>
<td><strong>I-11m</strong>.h658 (clade I)</td>
</tr>
<tr>
<td><strong>I-11m</strong>.h475 (clade I)</td>
</tr>
<tr>
<td><strong>I-11m</strong>.h576 (clade I)</td>
</tr>
<tr>
<td><strong>Gene diversity</strong>(H) and (h)(c)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Bifidobacterium</strong> haplotypes(a)</th>
<th><strong>M-1m</strong></th>
<th><strong>M-11m</strong></th>
<th><strong>I-1m</strong></th>
<th><strong>I-11m</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>M-1m</strong>.h450</td>
<td>0</td>
<td>8 (0.21)</td>
<td>0</td>
<td>0</td>
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<tr>
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<td>0</td>
<td>4 (0.11)</td>
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<td>0</td>
</tr>
<tr>
<td><strong>M-1m</strong>.h740</td>
<td>0</td>
<td>3 (0.08)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td><strong>M-1m</strong>.h549</td>
<td>0</td>
<td>2 (0.05)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td><strong>M-1m</strong>.h637</td>
<td>0</td>
<td>2 (0.05)</td>
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<td>0</td>
</tr>
<tr>
<td><strong>I-11m</strong>.h478</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>13 (0.29)</td>
</tr>
<tr>
<td><strong>I-11m</strong>.h658</td>
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<td>0</td>
<td>0</td>
<td>2 (0.04)</td>
</tr>
<tr>
<td><strong>I-11m</strong>.h475</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2 (0.04)</td>
</tr>
<tr>
<td><strong>I-11m</strong>.h576</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2 (0.04)</td>
</tr>
<tr>
<td><strong>Total singletons</strong>(b)</td>
<td>0</td>
<td>19 [12]</td>
<td>0</td>
<td>26 [9]</td>
</tr>
<tr>
<td><strong>Total haplotypes</strong>(b)</td>
<td>0</td>
<td>24 [21]</td>
<td>0</td>
<td>30 [18]</td>
</tr>
<tr>
<td><strong>Gene diversity</strong>(H) and (h)(c)</td>
<td>NA</td>
<td>0.95 [18.00]</td>
<td>NA</td>
<td>0.92 [11.22]</td>
</tr>
</tbody>
</table>

**Note.**—NA, not applicable.

\(a\) Only haplotypes recovered multiple times are shown individually. Clade designations are from figure 2B. Haplotype frequencies are shown in parentheses.

\(b\) Expected numbers of singletons or total haplotypes in a steady-state population are bracketed.

\(c\) \(h\) values are bracketed.
be adequate to investigate differences between individuals or sampling times (Turnbaugh et al. 2009).

Our community composition and microdiversity analysis based on fecal samples indicate that between 1 and 11 months, the infant GIT microbiota increased substantially in complexity while undergoing a significant turnover of taxa at several phylogenetic levels. In terms of taxonomic composition, we detected *Escherichia* and *Bacteroides* as the most abundant organisms in the 1-month-old infant, whereas *Bifidobacterium* only appeared and grew to dominate the infant GIT microbiota at some later stage between 1 and 11 months. Although *Bifidobacterium* is thought to dominate the early GIT microbiota in most breast-fed babies (Stark and Lee 1982; Mackie et al. 1999; Favier et al. 2002; Hopkins et al. 2005; Penders et al. 2006), a preponderance of *Escherichia* and *Bacteroides* has also been commonly observed, especially in cases where *Bifidobacterium* is not present (Stark and Lee 1982; Mackie et al. 1999). Remarkably, the mother followed a compositional shift at the genus level between 1 and 11 months that was similar to that of the infant: in both cases, *Bifidobacterium* appeared and rose to a substantial proportion of the GIT community, whereas *Bacteroides* decreased importantly (fig. 1C). However, it is important to note that the mother in our study followed a course of prescribed antibiotics a few days before delivery, and the perinatal administration of antibiotics, currently a frequent practice in industrialized countries, likely altered the composition of the maternal microbiota (Spaetgens et al. 2002). In particular, human GIT *Bifidobacterium* species are known to be highly sensitive to antibiotic treatment (Gewolb et al. 1999; Delgado et al. 2005), which might account for their absence in our study at 1 month after delivery.

Although our sequencing depth was not high and we did not repeat samplings for given time points, it is unlikely that undersampling could explain the large shifts in relative taxa abundances and the observed changes in dominant taxa between samples. Moreover, the general features of the pattern of changes in community complexity in the mother–infant pair concur with those reported in previous analyses based on 16S rDNA or on culture-dependent techniques, with a limited number of taxa in the 1-month-old infant, a large increase during the first year of life (Stark and Lee 1982; Mackie et al. 1999; Favier et al. 2002; Fanaro et al. 2003; Hopkins et al. 2005; Park et al. 2005; Penders et al. 2006; Palmer et al. 2007), and a stable number of taxa in the mother across the same time period (Zoetendal et al. 1998; Ley, Turnbaugh, et al. 2006).

Even though mother and infant incurred similar genus-level compositional shifts between 1 and 11 months (fig. 1C), microdiversity analyses revealed that the populations of *Bifidobacterium* that had been acquired by each individual by 11 months belonged to different species and that the *Bacteroides* populations of maternal origin present in the 1-month-old infant had been replaced at this point by a novel one. These results indicate that genera that are shared by mother and infant may have undergone species-level turnover or represent independent acquisitions from external sources, which might include food, other individuals, or other components of the environment rather than vertical inheritance. Two recent analyses compared the overall composition of the GIT microbiota between mothers and infants or adult daughters with varying conclusions regarding the potential role of vertical transmission (Palmer et al. 2007; Turnbaugh et al. 2009). However, these analyses did not report whether the most prevalent genera in their mother–child samples were represented by identical species.

At an even finer diversity scale, our analyses provide the first insights into the population biology of GIT bacteria. In the case of *Bacteroides*, identities between 16S and fosmid sequences recovered from mother and 1-month infant samples presented clear evidence for an early transmission of the maternal strain. In addition to the recognition of identical
sequences, the high similarity between the frequencies of the most abundant haplotypes in the maternal and the I-1m samples confirmed the likelihood of a transmission event (table 2), and further analysis of the frequency distribution of different 16S haplotypes across these samples provided additional information regarding the process by which Bacteroides was transmitted among hosts. The fact that the two most abundant maternal haplotypes, but only one of the low-frequency haplotypes, were identified in the infant suggests that colonization involved a small maternal inoculum, one in which most low-frequency alleles were not represented. Despite this likely bottleneck, the fact that I-1m contained seven new haplotypes, each derived from these inherited from the mother by a single nucleotide change (fig. 3), suggests that colonization of the infant gut by a small inoculum was followed by the quick appearance of novel genetic variants. This process is expected when a population expands rapidly after a brief bottleneck (Maruyama and Fuerst 1984), as would probably occur after the maternal Bacteroides strain entered the GIT of the young infant. Alternatively, the seven singletons found uniquely in I-1m might have been present in the mother at very low frequencies and be strongly selected for after transmission to the infant. Although this scenario is possible, it does not seem likely, as it would imply that the seven singletons were able to outcompete most of the haplotypes detected in both maternal samples, which would presumably also have been transmitted in the maternal inoculum, and without substantially altering the frequencies of the two major haplotypes. Similarly, it does not seem parsimonious that seven haplotypes, each differing from the most abundant maternal haplotype by a single nucleotide, would have been acquired from the environment through an independent inoculation.

In a growing population, new mutants accumulate and produce an excess of low-frequency haplotypes relative to the theoretical expectation for a population that has achieved a steady state; based on an observed level of gene diversity, it is possible to compute the total number of haplotypes and the number of singletons expected in a theoretical equilibrium population (Ewens 1972; Nei and Li 1976; Maruyama and Fuerst 1984). Such calculations confirmed that the Bacteroides population in the 1-month-old infant contained excesses of singletons and of total haplotypes (table 2), as is characteristic of a population expanding after a bottleneck, and supporting the hypothesis of colonization by a small inoculum followed by in situ diversification by mutation in a rapidly growing population. In fact, low-frequency haplotypes were also present in excess in each of the Bifidobacterium and Bacteroides populations analyzed from every sample (table 2). It is not surprising to detect indications of rapid growth in the populations that were novel at 11 months (i.e., Bifidobacterium in the mother, and both Bifidobacterium and the new Bacteroides strain in the infant) because these could have been recently founded. The lack of equilibrium in genetic diversity is more difficult to interpret for the older maternal Bacteroides population that was already present at the 1-month sampling, especially given the likely alterations in population dynamics caused by the administration of antibiotics before delivery. However, other 16S analyses of adult GIT samples have also detected the existence of very shallow fans of diversity for different genera (Eckburg et al. 2005; Ley, Peterson, and Gordon 2006), suggesting that bacterial populations in the gut may commonly exist far from genetic equilibrium and may be due to frequent oscillations in population size or to selection for rare variants. Experimental inoculation and subsequent monitoring of strains in model animals should enable a better understanding of bacterial population dynamics in the GIT.

In regard to coding capabilities, characterization of the COG functional classes present in the infant GIT at 1 and 11 months after birth identified no broad differences between these two stages or between the infant and the mother (D rank analysis; P > 0.05) in the face of remarkable change in phylogenetic composition (fig. 1A and C). Analysis of COG functional classes can distinguish bacterial communities inhabiting different environments (Tringe et al. 2005; Dinsdale et al. 2008). This suggests that, in spite of dietary and other differences, the infant and adult GIT offer largely similar environments for their bacterial inhabitants. In both, genes related to carbohydrate and amino acid transport and metabolism were the most abundant, whereas those involved in lipid transport and metabolism were rare, in accordance with previous metagenomic analyses of the human gut microbiota (Gill et al. 2006; Kurokawa et al. 2007; Turnbaugh et al. 2009). Carbohydrates and peptides, rather than fats, are therefore likely to represent the main sources of nutrients for GIT bacteria in the exclusively breast-fed 1-month-old infant as well as in the 11-month-old infant and the adult.

Furthermore, analyses targeted to functional groupings most likely to sense dietary differences between the two infant stages and the mother also failed to detect significant variation among samples. We hypothesized that 1) COGs encoding proteins able to transport or metabolize the different oligosaccharides that are abundant in human milk (group I) could be most frequent in the exclusively breast-feeding stage (I-1m) than in the adult and at the partially weaned stage when solid foods have already been introduced to the infant diet (I-11m), whereas 2) COGs encoding proteins specialized for the transport and metabolism of plant carbohydrates (group II) could be most frequent in the adult and partially weaned stages. However, the proportions of sequences belonging to these two COG groups were not different between the different samples by D rank analysis, and the number of group I COGs represented was actually higher in the maternal than in the infant samples. Previous analyses also indicated that similar families of glycosyl hydrolases were present in the infant and adult gut microbiota, including enzymes that
degrade carbohydrates of plant origin (Kurokawa et al. 2007). This suggests that the global makeup of the unweaned infant microbiota is not particularly adapted to a breast milk diet but rather that this bacterial community exploits the oligosaccharides of breast milk with a repertoire of enzymes similar to that present in the older infant and the mother.

It is necessary to note that although pairwise comparisons of the abundance of different functions and functional groupings between samples did not identify significant differences, a PCA of overall COG category profiles suggested that relative similarity was greatest for the two maternal microbiota samples and the infant sample taken at 11 months to the exclusion of I-1m (fig. 1). This result reflects the fact that one of the COG categories (W—Extracellular Structures) is uniquely represented in the 1-month infant sample. However, the few W COG category sequences recovered from I-1m are orthologs of E. coli secreted adhesins often associated with intestinal and nonintestinal infections (Restieri et al. 2007), and their presence is not likely to represent a functional adaptation specific to the gut environment of the 1-month-old infant.

In conclusion, our analyses present a dynamic GIT microbiota, capable of accommodating a significant turnover of taxa at different phylogenetic levels while maintaining a remarkable constancy in total functional capabilities. Furthermore, this study has generated the first insight into the bacterial population-level processes that shape the coevolution between the GIT microbiota and the human host. However, 16S rRNA analyses have repeatedly shown that there is a large degree of interindividual variability among infants in the development of the microbiota (Favier et al. 2002; Adlerberth et al. 2006; Penders et al. 2006; Palmer et al. 2007), and, therefore, numerous studies of this sort will be required to evaluate the generality of the trends detected here; large-scale studies of microbiota development in cohorts of infants are now enabled by the increasing facility of high-throughput sequencing. If typical, the dynamic patterns detected in this study would have significant evolutionary and medical implications. On one hand, our analyses suggest a lack of long-term maintenance of maternal phyotypes in the infant, although it is very difficult to rule out a low-level presence of a strain in a given environment, and further in-depth sequencing in other mother–infant pairs will be necessary to address this issue. If confirmed, the phyotype turnover suggested here would limit a tight coadaptation between specific strains and host genotypes and the adaptive diversification of bacterial lineages within host family lines (Ley, Peterson, and Gordon 2006); rather, an intermittent pattern of interactions between different bacterial strains and host genotypes would likely result in a diffuse process of coevolution with little local adaptation and moderate functional differentiation among strains (Futuyma and Slatkin 1983; Thompson 1994). On the other hand, the observed turnovers of genera, species, and strains offer heartening news for medical approaches aimed at modulating the composition of the gut microbiota because they suggest that the phylogenetic composition of this community is malleable and should be responsive to probiotic intervention. Finally, the observation that similar functional repertoires can be attained in the GIT microbiota in the face of substantial differences in phylogenetic composition supports the notion that diseased states of this community may be best identified by atypical distributions of functional gene categories, as recently shown for obesity (Turnbaugh et al. 2009).

Supplementary Material

Supplementary tables S1–S2 and figure S1 are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org/).

Acknowledgments

We thank Falk Warnecke for introducing us to the ARB package, Edward Kirton for processing sequencing data through the Genelib pipeline, Costas Mavrommatis and Ernest Szeto for assistance with IMG, Juan José Abellán for statistical advice, and Becky Nankivel and Shubhangi Kadhe for help in the preparation of the figures. This work was supported by the National Institutes of Health (grant number R01 DK66288 to M.P.F.). This work has been performed under the auspices of the US Department of Energy’s Office of Science, Biological and Environmental Research Program and by the University of California, Lawrence Berkeley National Laboratory under contract number DE-AC03-76SF00098.

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