Composition, variability, and temporal stability of the intestinal microbiota of the elderly

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Alterations in the human intestinal microbiota are linked to conditions including inflammatory bowel disease, irritable bowel syndrome, and obesity. The microbiota also undergoes substantial changes at the extremes of life, in infants and older people, the ramifications of which are still being explored. We applied pyrosequencing of over 40,000 16S rRNA gene V4 region amplicons per subject to characterize the fecal microbiota in 161 subjects aged 65 y and older and 9 younger control subjects. The microbiota of each individual subject constituted a unique profile that was separable from all others. In 68% of the individuals, the microbiota was dominated by phylum Bacteroides, with an average proportion of 57% across all 161 baseline samples. Phylum Firmicutes had an average proportion of 40%. The proportions of some phyla and genera associated with disease or health also varied dramatically, including Proteobacteria, Actinobacteria, and Faecalibacterium. The core microbiota of elderly subjects was distinct from that previously established for younger adults, with a greater proportion of Bacteroides spp. and distinct abundance patterns of Clostridium groups. Analyses of 26 fecal microbiota datasets from 3-month follow-up samples indicated that in 85% of the subjects, the microbiota composition was more like the corresponding time-zero sample than any other dataset. We conclude that the fecal microbiota of the elderly shows temporal stability over limited time in the majority of subjects but is characterized by unusual phylum proportions and extreme variability.

Recent advances in culture-independent techniques for microbial community analysis (1, 2) have led to major progress in understanding the complexity of the human intestinal ecosystem (reviewed in ref. 3). High-throughput amplicon pyrosequencing is emerging as the method of choice for determining the microbial phylotypes present in a sample (4). Total shotgun sequencing of the microbial metagenome is a robust approach to determine community composition with low amplification bias (5), but the cost is currently prohibitive for routine application to large numbers of samples.

The initial infant gut microbiota is a simple structure usually dominated by bifidobacteria, and through a series of successions and replacements, it migrates to a more complex adult pattern (2, 6). The adult intestinal microbiota has been shown to be relatively stable over time (7) and is sufficiently similar between individuals to allow identification of a core microbiome comprising 66 dominant operational taxonomic units (OTUs) that corresponds to 38% of sequence reads from 17 individuals (8). Estimates of the numbers of organisms present in the human gut range from 1,183 bacterial phylotypes based on a meta-analysis (9) to 3,180 OTUs (8). Changes in the adult gut microbiota have been linked to undesirable health conditions including obesity (10), inflammatory bowel disease (IBD) (11), and irritable bowel syndrome (IBS) (12). The link between obesity and the microbiota is likely to be more sophisticated than the simple phylum-level Bacteroidetes:Firmicutes ratio initially identified (13), and it is likely to involve microbiota–diet interaction (14–16). More subtle alterations in the levels of other bacteria in the gut may also impact on health. Some of the microbiota members most responsive to diet include the Clostridium clusters IV and XIVa; many of these species are producers of butyrate, which is an important energy source for intestinal cells (17, 18). Clostridium cluster IV also includes Faecalibacterium prausnitzii, which is depleted in IBD patients (11, 19, 20) and has anti-inflammatory activity in vitro and in mouse models (21). In contrast, increases in the numbers of adherent Escherichia coli cells have been associated with IBD (22, 23).

The proportion of people over the age of 65 y is increasing in many countries and will exceed the proportion of infants under 5 y of age on a global basis in 10 y (24). The aging intestine is subject to a number of physiological changes that impact on food digestion and absorption (25, 26) and immune function (27). Accompanied by a loss in the proportions of naïve CD4+ T cells, the inflammation process is characterized by persistent activation of innate immunity mediated by the NF-κB transcription factor (28). Given the potential for elements of the microbiota to impact on these dynamic processes, there is considerable interest in determining the composition of the gut microbiota of elderly subjects and characterizing its variation as a potential determinant of health (3). The gut microbiota composition of elderly subjects is expected to be in a state of flux, a theory supported by culture-dependent methods (29). More recent studies have suggested that the Bacteroides species diversity was marginally increased in the feces of elderly subjects and that of bifidobacteria

References

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reduced (30), whereas a further study suggested that both Bacteroides numbers and species diversity decline in the elderly (31, 32). The proportions of the phyla Firmicutes, Bacteroidetes, and Proteobacteria were 58%, 15%, and 27%, respectively (33) in three elderly subjects in Japan, using 16S rRNA gene-clonal sequence sequencing. In a study based in four European countries using labeled probe technology, significant variations in the proportions of major bacteria phyla were recorded between countries, emphasizing the danger of generalization of findings from one study to another (34) and illustrating the likelihood of diet-microbiota interactions being involved.

The ELDERMET consortium (http://eldermet.ucr.edu) was established in 2007 to investigate the role of the intestinal microbiota in elderly Irish subjects as an agent and indicator of health. The initial phase of the ELDERMET project is to establish the baseline microbiota composition in several hundred elderly subjects, for whom extensive clinical, health, and dietary intake information has been obtained. This data will allow identification of individuals and cohorts for full shotgun fecal microbiome sequencing to investigate functional linkages to health status. Here, we report the data for the microbiota composition of 161 elderly subjects at time 0, 26 of which were also sampled 3 mo later. The data are providing unprecedented insights into the composition and interindividual variability of the gut microbiota in the elderly and identifying several organisms whose high variability suggests they may be amenable to therapeutic modulation by dietary or other means.

Results

Study Design and Clinical Analysis of an Elderly Subject Cohort. Full ethical approval for this study was obtained from the Local Research Ethics Committee. One hundred sixty-one subjects entered the study; 82 female and 79 male (SI Materials and Methods). 161 baseline samples plus 26 3-mo follow-up samples, based on the same protocol, was more typical for published studies of younger adults (Firmicutes-dominant) (Fig. 1B). The Bacteroidetes-dominant character of the microbiota in elderly subjects was also apparent from the genus assignments, despite the fact that the RDP classifier does not efficiently classify as many amplicon sequences at genus level as it does at phylum level, on average, 78 ± 11% SEM of all reads are assigned at genus level compared with 99 ± 0.1% at phylum level. Noteworthy among the well-characterized bacterial genera were Faecalibacterium spp. at 6% (6.1 ± 0.3%) of the 15 most abundant phyla, Ruminococcus spp. at 4% (3.9 ± 0.3%), Roseburia spp. at 3% (3.2 ± 0.2%) (Fig. 1C), and Bifidobacterium spp. at 0.4% (0.4 ± 0.07%). Exclusion of the subjects who had been treated with antibiotics in the 4 wk before the time-0 sample (n = 43) or exclusion of the 26 3-mo follow-up samples had no impact on the taxon assignment proportions (Fig. S1).

The clostridia constitute a large class in phylum Firmicutes that has been divided into 19 groups or clusters (37). These clusters include organisms that are dominant or subdominant organisms in the human intestinal microbiota (9), and variations in some of them have been reported in stool of IBD patients (38); F. prausnitzii, a prominent member of Clostridium cluster IV, is also depleted in IBD (39). To assist in the assignment of the short V4 amplicon sequences to Clostridium groups, we developed an approach based on an association table of V4 region sequences from species previously assigned to Clostridium clusters (Materials and Methods). Thus, some 25% of 6.3 million amplicon reads were assigned to Clostridium clusters. Clostridium clusters IV and XIVa represented the dominant assignments, with 56% (56.6 ± 1%) and 35% (34.7 ± 1.1%) of the assigned sequences, respectively. Clusters XVI and XI comprised 3% (3.4 ± 0.4%) and 2% (1.7 ± 0.2%), respectively, of the total assignments. In the younger adults, Clostridium clusters IV and XIVa were also the dominant clusters with similar proportions, at 56% (56.0 ± 1.6%) and 36% (35.3 ± 1.9%), respectively.

Dramatic Interindividual Variability of the Elderly Gut Microbiota. Whereas the aggregate microbiota was dominated by the Bacteroidetes, the individual composition datasets showed extraordinary variation, with the proportion of the Bacteroidetes ranging from 3% to 92% and the proportion of the phylum Firmicutes ranging from 7% to 94% (Fig. 2A). Exclusion of the 43 antibiotic-treated subjects or 3-mo samples had no impact on the degree of variability (Fig. S2). The proportions of five other major phyla also varied substantially between individuals. For example, in several microbiota communities, the phylum Proteobacteria was unusually abundant, comprising between 11% and 25% of total reads, whereas the average was 2% (Fig. S2A). Although the average proportion of the phylum Actinobacteria was under 1% (0.44%) (Fig. 2), some samples harbored unusually high levels, at 8%, 5%,

Fig. 1. Aggregate microbiota composition at phylum level in (A) 187 (161 time-0 and 26 time 3-mo) fecal samples from elderly subjects, (B) 9 healthier young adult controls, and (C) at genus level in the 187 fecal samples from elderly subjects (the smaller pie chart to right is inserted for clarity). Only major taxonomic groups are shown; these cover 95% of all reads assigned to genus level.
and 4%. Similarly, considerable variation was detected for the proportions of the *Clostridium* clusters (Fig. 2B). Proportions of the total number of reads for clusters IV and XIVa ranged from 1.7% to 34% and 1.1% to 22%, respectively. None of the extreme values were from subjects treated with antibiotics or those returning after 3 mo (Fig. S3). *Faecalibacterium* and *Ruminococcus* varied most in their proportions, reaching 16.5% and 15.7% in some subjects, respectively (Fig. 2C). The two highest proportions were, perhaps surprisingly, detected in individuals who had been administered antibiotics (Fig. S4B), but unusually high proportions of ca. 10–12% of both *Faecalibacterium* and *Ruminococcus* were also recorded in subjects not taking antibiotics (Fig. S4A and C).

**Different Fecal Core Microbiota in Younger and Older Adults.** We identified and compared the core microbiota of younger adults [derived from previously published studies (8, 13, 40, 41) and ELDERMET subjects] (Table S2). Because procedures differ between individual studies, the bioinformatic analysis was harmonized by extracting V4 sequences from published full-length datasets from younger adults. Selection was based on full-length sequence availability (V4 extraction possible), being derived from healthy nonelderly adults and information on the subject from which they came. Although this comparison of high-coverage pyrosequencing data with lower coverage clone-library Sanger reads is complicated by the potential sampling bias of the latter, previous studies have shown similar distributions of the major taxa when measured by both full-length and partial pyrosequencing reads (4, 13, 41). We excluded from this comparison all subjects that had been treated with antibiotics, were subjected to gastric bypass (subjects gb1–3 in ref. 41), and were aged between 18 and 58 y among the healthy adults (subjects GG and BB in ref. 40). The extracted V4 region sequences were binned into unique sequences and grouped by subject. As had also been observed at the 97% OTU level (8), the subject specificity (proportion of unique sequences present in only one sample) was high in all datasets (Table S2). We subsequently defined the core microbiota as all of the unique sequences present in at least one-half of the subjects, following an approach applied in a recent study (8). Assignments to phylum, genus, and *Clostridium* cluster for each of these datasets revealed large differences between the elderly and younger adults (Fig. 3). More than one-half of the core microbiota (53%) in the elderly subjects comprised *Bacteroidetes* were made up by the genera *Bacteroides* (29%), *Alistipes* (17%), and *Parabacteroides* (7%) compared with 8–27% in the younger adults. Although less pronounced, assignments to the two major *Clostridium* clusters were also unequal, even more than when comparing aggregate microbiota, with cluster IV being the more predominant cluster among subjects older than 65 y, whereas cluster XIVa was more prevalent in the younger cohort.

Because the same variable region was used in this comparison of five studies, we were able to identify and classify core sequences that were unique to subjects either at least 65 y of age (400 sequences) or between 18 and 58 y (18 sequences) (Tables S3 and S4). The sequences unique to the elderly subjects correspond to a wide array of *Bacteroides, Alistipes*, and *Parabacteroides* species,
whereas younger subjects had one unique Bacteroides species, most similar to Bacteroides vulgatus. The elderly subjects also had a much higher abundance of Faecalibacterium spp. and consequently, Clostridium cluster IV. The composition of the (Bacteroides-dominated) core microbiota that was unique to elderly subjects was not markedly different when it was defined by either comparison with the datasets of younger subjects from this study combined with the four published datasets (Fig. 3B) or comparison exclusively with the data from the nine younger subjects generated in this study (Fig. 3C). The latter comparison resulted in 167 and 1,177 core sequences unique to the elderly and younger adults in this study, respectively. This shows that the low sampling size of the Sanger data in the former comparison did not compromise the outcome.

Intraindividual Fecal Microbiota Stability in a Majority of Elderly Subjects. To analyze changes in the microbiota over 3 mo in the same subject, we first analyzed the relatedness of the total pool of reads from all samples using UniFrac, as detailed in Materials and Methods. Briefly, a phylogenetic tree was built from the alignment of all ~1 million unique V4 sequence reads, from which UniFrac distances between all samples were calculated. These distances were used for hierarchical clustering, whereby relative abundances of unique sequences were factored (weighted UniFrac tree) (Fig. S5A) or only presence/absence data were factored (unweighted UniFrac tree) (Fig. S5B). In the unweighted tree for 161 time-0 and 26 time 3-mo samples, 85% (22/26) of the microbiota compositions for the time 3-mo samples localize most closely to their time-0 sample. This figure drops to 62% based on the weighted tree, presumably because of discrimination based on phylotypes being overwhelmed by weighting values of fewer taxa that each have more reads. However, by either approach, a given microbiota after 3 mo was more like to localize to its original composition than to any other sample analyzed.

Fig. S6 shows the fecal microbiota composition for the 26 pairs of time-0/time 3-mo fecal samples at phylum (Fig. S6A) and genus assignment levels (Fig. S6B). Four of six temporal pairs that matched only in the unweighted tree were still proximal in the weighted tree (Fig. S5B). It is evident that those temporal pairs that are separated in both weighted and unweighted trees also show larger variations of phyla and genus proportions. However, even for some temporal pairs that are each other’s closest neighbors in the unweighted tree only (e.g., EM047 and EM053), there are also relatively large phylum and genus variations.

Microbiota compositional differences were greater between individuals than within individuals (Fig. 4). The average unweighted UniFrac distances between 26 fecal microbiota communities were 0.87 ± 0.0016 SEM for the time-0 samples and 0.87 ± 0.0014 for the time 3-mo samples. The average unweighted UniFrac distance within the time-0 to time 3-mo samples was lower, at 0.76 ± 0.0104. The unweighted UniFrac distance within the 26 individuals from time 0 to time 3 mo was significantly lower than the between individual unweighted UniFrac distances at both time 0 (robust t test; \( P < 0.001 \)) and time 3 mo (robust t test; \( P < 0.001 \)). The weighted UniFrac analysis show a similar trend with average UniFrac distances between time-0 samples of 0.83 ± 0.0102 SEM, an average distance of 0.85 ± 0.0115 between all time 3-mo samples, and a much lower average UniFrac distance of 0.45 ± 0.0434 for between time-0 and time 3-mo samples. The weighted UniFrac distance within the 26 individuals from time 0 to time 3 mo was significantly lower than the between individual weighted UniFrac distances at both time 0 (robust t test; \( P < 0.001 \)) and time 3 mo (robust t test; \( P < 0.001 \)).

In some paired samples (e.g., EM059), the loss of phylogenetic proximity was not readily apparent at phylum level but was evident at genus level (Fig. S6B). Analysis of the genus assignments for these pair-wise comparisons showed that relative increases in the Firmicutes proportions, particularly Sporobacter spp. (order Clostridiales, family Ruminococcaceae), and the Lachnospiraceae family (order Clostridiales), were often present in microbiota pairs that diverged. Changes in the proportions of Alstipes spp. (order Bacteroidales, family Rikenellaceae), Parabacteroides (order Bac-

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teroidales), or Anaerophaga (order Bacteroidales) were also responsible for other divergent paired samples.

Effect of Antibiotics on the Microbiota. The impact of antibiotic usage was not readily evident from high-level comparisons of aggregate data (Fig. S1 B and C); a modest but significant effect of antibiotic use on the abundance of both phyla and genera was recorded. The average proportion of total reads assigned to the different phyla and genera were compared according to antibiotic use for the 161 baseline samples. Of the seven largest phyla in terms of composition proportion, a significantly increased relative abundance of Bacteroidetes (Mann–Whitney test; *P* = 0.0297) and a significant increase in the Bacteroidetes: Firmicutes ratio (Mann–Whitney test; *P* = 0.0363) were recorded in antibiotic users, with decreases in the relative abundance of both Firmicutes (Mann–Whitney test; *P* = 0.0445) and Proteobacteria [significant using the Mann–Whitney test (*P* = 0.0232) but not the *t* test (*P* > 0.05)]. At the level of genus, only the relative abundance of Parabacteroides was significantly increased (Mann–Whitney test; *P* = 0.0465) with antibiotic use.

Five of twenty-six subjects that provided both time-0 and time 3-mo fecal samples had been administered antibiotics between these visits, one belonging to the first group in Fig. S6 (EM059), three belonging to the second group (EM047, EM053, and EM077), and one belonging to the last group (EM090). There were significant differences in the time-0 and time 3-mo fecal microbiota profiles of these subjects according to antibiotic use. Antibiotic use significantly decreased the relative abundance of Actinobacteria (Mann–Whitney test; *P* = 0.0472). At genus level, Alistipes abundance was significantly increased in subjects who had taken antibiotics using the *t* test (*P* < 0.05) but not the Mann–Whitney test (*P* = 0.1263). Only two of these five subjects had been prescribed the same antibiotic, albeit at different dosages. A low, 250-mg dose of clarithromycin decreased the proportion of Bacteroidetes (Bacteroides and Parabacteroides) and equivalently increased the proportion of Firmicutes (Alistipes). In contrast, a higher, 500-mg dose of clarithromycin increased the proportion of Bacteroidetes (Parabacteroides) and nonequivalently decreased the proportion of Firmicutes (Alistipes), indicating changes in the proportions of other phyla.

Discussion

This study constitutes the largest and deepest sampling of the composition of the elderly gut microbiota reported to date. The fecal microbiota of younger adults was previously shown by temperature gradient gel electrophoresis (TGGE) and phylochip (HitChip) analysis to be stable and individual-specific (7, 42). In the present study, all 161 of the time-0 microbiota samples could be clearly separated by UniFrac analysis (Fig. S5), and fine-detail analysis of genus confirmed individual-specific microbiota configurations. The HitChip study also found that temporarily paired samples from elderly subjects were more similar than randomly compared samples from different subjects. However, the similarity of paired samples decreased from time 1 mo to time 2 mo. In the five elderly subjects that they investigated, the decrease in microbiota similarity was more pronounced in the Actinobacteria (42), whereas in younger adults, greatest divergence was recorded for Clostridium cluster IV. We observed a similar phenomenon in the 3-mo microbiota comparisons, suggesting that these levels of bacteria are commonly in flux in elderly subjects.

Subjects over 65 y of age often have morbidities ranging from mild to severe that increase in complexity and seriousness with advancing age. These conditions are treated with a complex repertoire of medications that are likely to have individual or combined effects on the microbiota. For the 4 of 26 (~15%) of the time-3 mo microbiota samples that were not stable over 3 mo in the unweighted UniFrac tree, no association was identified with the potentially influential factors of nonantibiotic prescription medication change or illness. Because antibiotic treatment is relatively common in the elderly, we did not exclude enrolling such subjects in the study but rather, treated them as a separate group for most analyses. Antibiotic treatment has been shown by culture-dependent methods (32), TGGE (43), and an elegant molecular method (4) to dramatically disturb the composition of the fecal microbiota. In three subjects treated with ciprofloxacin (4), the antibiotic caused a decrease in the taxonomic richness, diversely and evenness of the community. Interestingly, the magnitude of the changes and the taxa affected were different in the three subjects, and the overall community structure had been restored by 4 wk posttreatment. Stochastic or individual-specific effects on the microbiota were also described in another study (43). In the simplest comparison of antibiotic treatment that we could make, different doses of clarithromycin resulted in counterintuitive impacts on the microbiota, which could clearly be complicated and confounded by factors other than antibiotic use.

The gut microbiota of the elderly revealed here (by analysis of a fecal surrogate sample, similar to the majority of other studies) was substantially different from that of younger adults (13, 40, 44–46), particularly with regard to the lower proportion of phylum Firmicutes detected. Over 65% of the time-0 microbiota communities (Fig. 3) had atypical Bacteroidetes: Firmicutes ratios compared with those described for younger adults. In a previous pilot study using the same primers, we showed that V4-based phylogenetic assignments correlated well with microbiota composition data produced by pyrosequencing hybridizing full-length 16S rRNA gene amplicons to a phylochip (35). This indicates lack of a primer-based bias in favor of the Bacteroidetes. Furthermore, the aggregate composition of the nine young adults analyzed using our DNA extraction, amplification, sequencing, and phylogenetic methodologies was similar to that found in other studies of younger adults (45) and was not Bacteroidetes-dominant. The diversity and range of phylum proportions that we identified herein mean that comparison of aggregate figures must be interpreted judiciously. Nevertheless, some top-level comparisons are informative. The range of phylum Firmicutes in our study, across all 161 time-0 samples, varied from 8% to 80%, whereas phylum Bacteroidetes varied from 14% to 92%. The phylum Actinobacteria includes Bifidobacterium spp., which are among the best characterized of the commensal organisms considered beneficial or probiotic in nature (47). This phylum ranged from 0% to 8% in the total dataset from 161 fecal samples, comparing with values of 0.4% (41) to 11.4% (46) in other studies of younger adults. In our analysis of the core microbiota, we observed a clear shift to a more Clostridium cluster IV-dominated community in the elderly. These elderly-specific core sequences all belonged to the Ruminococcaceae family and essentially, comprised Faecalibacterium, Sporobacter, and Ruminococcus species. In a recent study, albeit using lower-resolution molecular methods (48), the proportion of Bacteroidetes in the fecal microbiota of 17 institutionalized elderly subjects was significantly higher than in younger adults. That study also noted relatively low Bifidobacterium proportions in elderly subjects, lower proportions of Clostridium cluster IV, and lower diversity overall (48).

This microbiota composition profiling provides the basis for a follow-up phase focusing on the function of the elderly gut microbiota. By first investigating the microbiota composition in a large panel of subjects of defined community setting, health status, and diet, we can rationally select samples for fecal shotgun metagenomics by reference to both host parameters and microbiota diversity. This approach is validated by the remarkable interindividual microbiota diversity that was evident in 161 time-0 fecal samples from phylum down to unique OTU level. Prior knowledge of microbiota composition will allow for more cost-effective application of shotgun microbiome sequencing to samples selected for ranges or types of phylogenetic diversity.

Materials and Methods

Subject Recruitment and Sample Collection. This study was approved by the Clinical Research Ethics Committee of the Cork Teaching Hospitals. Subjects, aged 65 y and older, were recruited and clinically investigated in two local hospitals. Inclusion criteria were identified by elderly referred here in an investigational drug evaluation within the last 30 days, or advanced organic disease. Informed consent was obtained from all subjects or in cases of cognitive impairment, next of kin in accordance with the local research ethics committee.
guidelines. Data collected included anthropometric measurements, clinical history and status, and medication history. Antibiotic use was also recorded for each subject. Nineteen adult subjects of ages ranging from 28 to 46 y that were not antibiotic-treated were also recruited by informed consent.

Molecular Methods and Bioinformatics. DNA was extracted from fecal samples, and the V4 region of the 16S rRNA gene was amplified, sequenced, and analyzed as described in SI Materials and Methods and ref. 35. In addition, a phylogenetic tree was built from an alignment of unique sequences (i.e., 100% identity OTUs) using fastTree (49). Based on this tree and abundances/presence/absence data of the unique sequences, Fast UniFrac (50) was used for hierarchical Unweighted Pair Group Method with Arithmetic Mean clustering and principal coordinate analysis. The resulting trees were visualized with Dendroscope (51). To assign reads to Clostridium clusters, 58,000 V4 sequences were extracted from full-length 16S rRNA genes with complete species assignments (RDP release 10.16). An association table was derived by comparing scores from an all-against-all BLAST search of the extracted sequences with a list of species previously assigned to Clostridium clusters. Subsequently, V4 sequence reads could be assigned to clusters if their BLAST scores against the custom database were above the threshold for the best-hit species in the association table. Sequences were deposited in the Short Read Archive.

Statistical Methods. Statistical analyses were carried out using Stata SE Release 11.0 (StataCorp 2009 Stata: Release 11. Statistical Software; StataCorp). Detailed statistical analysis is in SI Materials and Methods.

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