Age-Related Variation of Bacterial and Fungal Communities in Different Body Habitats across the Young, Elderly, and Centenarians in Sardinia

Lu Wu,a,b Tiansheng Zeng,a Massimo Deligios,b Luciano Milanesi,c Morgan G. I. Langille,d Angelo Zinellu,b Salvatore Rubino,b Ciriaco Carru,b David J. Kelvin,a,d

a Division of Immunology, International Institute of Infection and Immunity, Shantou University Medical College, Shantou, Guangdong, China
b Department of Biomedical Sciences, University of Sassari, Sassari, Italy
c Institute of Biomedical Technologies, National Research Council of Italy, Segrate, Italy
d Department of Microbiology and Immunology, Dalhousie University, Halifax, Nova Scotia, Canada

Ciriaco Carru and David J. Kelvin contributed equally to this work as corresponding authors.

ABSTRACT Human body microbes interact with the host, forming microbial communities that are in continual flux during the aging process. Previous studies have mostly focused on surveying a single body habitat to determine the age-related variation in the bacterial and fungal communities. A more comprehensive understanding of the variation in the human microbiota and mycobiota across multiple body habitats related to aging is still unclear. To obtain an integrated view of the spatial distribution of microbes in a specific Mediterranean population across a wide age range, we surveyed the bacterial and fungal communities in the skin, oral cavity, and gut in the young, elderly, and centenarians in Sardinia using 16S rRNA gene and internal transcribed spacer 1 (ITS1) sequencing. We found that the distribution and correlation of bacterial and fungal communities in Sardinians were largely determined by body site. In each age group, the bacterial and fungal communities found in the skin were significantly different in structure. In the oral cavity, age had a marginal impact on the structures of the bacterial and fungal communities. Furthermore, the gut bacterial communities in centenarians clustered separately from those of the young and elderly, while the fungal communities in the gut habitat could not be separated by host age.

IMPORTANCE Site-specific microbial communities are recognized as important factors in host health and disease. To better understand how the human microbiota potentially affects and is affected by its host during the aging process, the fundamental issue to address is the distribution of microbiota related to age. Here, we show an integrated view of the spatial distribution of microbes in a specific Mediterranean population (Sardinians) across a wide age range. Our study indicates that age plays a critical role in shaping the human microbiota in a habitat-dependent manner. The dynamic age-related microbiota changes we observed across multiple body sites may provide possibilities for modulating microbe communities to maintain or improve health during aging.

KEYWORDS 16S rRNA sequencing, ITS1 sequencing, Sardinian, aging, gut microbiota, human microbiome, oral microbiota, skin microbiota

The human body and microbes that inhabit it organize together into a superorganism. The distribution of microbes in the human body is largely and primarily determined by body habitats with different ecological niches (1, 2); moreover, it is also influenced by age, lifestyle, ethnicity, and geography (3–6). The resident microbes
include not only bacteria but also other microorganisms, such as fungi and archaea. Dysfunction of any members of this community may affect the health of the human body and may cause pathogenic diseases (7, 8). Moreover, the human microbiotas distributed across the body are not isolated from one another but are interconnected as a unit (9). Thus, multikingdom microbiota communities across the human body should not be considered in isolation but as part of an individual pool of interacting communities.

Aging is associated with a wide array of physiological, metabolic, and immunological function declines (10). As a complex and dynamic ecosystem, the human microbiota is subject to continual change in life. Site-specific physiological changes take place during the aging process and are associated with microbial communities also undergoing site-specific changes. For example, the aging-related decrease in sebum in the skin is associated with changes in the skin microbiota (11, 12). Physical and cognitive age-associated changes of a host lead to lifestyle alterations, such as dietary preference changes caused by reducing dental status and decreasing chewing ability (13). Dietary fiber deprivation can change the gut microbiota composition and promote intestinal barrier dysfunction (14). Interestingly, many aging-related clinical issues are closely correlated with the changed microbiota. For example, ulcers in the skin which commonly happen in the physically limited or bedridden elderly were associated with altered skin microbiota (15). A proinflammatory gut microbiota may contribute to the development of atherosclerosis and stroke (16, 17). Moreover, the microbiota also can modulate the aging process (18). The intestinal barrier dysfunction and systemic inflammation promoted by age-related microbial dysbiosis have been observed in mice (19). Therefore, investigating the age-related microbiota change in humans is critical to understanding the interplay between the human microbiota and host during aging.

Aging-related alterations in the gut bacterial community in different cohorts have been demonstrated in previous studies (20–28). The variation may be caused by diverse genetic, dietary, and environmental factors (29–32). Age-related skin bacterial community changes in a Japanese population and a Chinese population have been reported (33, 34). The oral microbiota is emerging as a promising indicator for heath (35, 36). However, the age-related variation in the oral microbiome at the population level has not been fully demonstrated (36, 37). Compared with extensive studies on site-specific bacterial communities in the human body, how aging affects the site-specific fungal communities is poorly understood.

In order to examine the variation of microbiota in different body habitats across different age groups, we recruited 65 subjects from Sardinia, Italy. This cohort formed part of the AKEntAnnos (AKEA) cohort study (38). The composition of the cohort was a healthy young group (age 19 to 33 years), a healthy elderly group (age 68 to 88 years), and a centenarian group (>100 years old). 16S rRNA gene and internal transcribed spacer 1 (ITS1) gene amplicon sequencing methods were carried out to survey the skin, oral, and gut microbiota communities at each site. Our study provides new insights into how age influences the diversity of bacterial and fungal communities in various body habitats in the Sardinian population.

RESULTS

16S rRNA gene and ITS1 gene amplicon sequencing of the microbiota from different body habitats. To compare community structures and the relationship between bacteria and fungi among body habitats in different age groups, a total of 65 Sardinians from three age groups were recruited (Table 1). For each subject, skin (four different sites, the left palm, right palm, forehead, and umbilicus), oral, and fecal samples were collected. The workflow is shown in Fig. S1 in the supplemental material. A total of 379 16S rRNA and 377 ITS1 quality control amplicon libraries were constructed and sequenced on an Illumina MiSeq platform. More than 9.6 million reads were generated for 16S rRNA sequencing, and 11.2 million reads were generated for ITS1 sequencing. On average, 25,341 clean reads for 16S rRNA sequencing and 29,813 clean reads for ITS1 sequencing were sequenced for each sample. Across all six body
sites, we detected members of 30 bacterial phyla assigned to 562 bacterial genera and 6 fungal phyla assigned to 691 fungal genera. The clinical characteristics and health measurements for the three age groups and the total distribution of the bacterial and fungal community taxonomy assignments for each sample are displayed in Data Set S1.

**Bacterial and fungal community α diversity in three age groups across the different body sites.** At a normalized sequence depth, the bacterial and fungal community diversity and richness evaluated by the Shannon diversity index and Chao1-estimated operational taxonomic units (OTUs) varied dramatically in different body sites (Fig. S2). Overall, both analyses revealed that the bacteria had a significantly higher α diversity than that of fungi in the same body sites (Fig. S2a; Kruskal-Wallis test, \( P < 0.001 \)). The left and right palms showed similar significantly higher bacterial and fungal community α diversity than did the forehead (Fig. S2b). The α diversity of the fungal communities in the oral cavity and gut was significantly lower than that in the palms (Fig. S2b). The Shannon diversity indexes of the bacterial communities were not significantly different in the gut and oral cavity, and neither were the α-diversity values of the fungal communities (Fig. S2b), while significantly higher Chao1 indexes of the bacterial communities and significantly lower Chao1 indexes of the fungal communities in the gut were observed than what was found with the oral cavity (Fig. S2b).

To further elucidate the age-related α diversity for bacterial and fungal communities, α diversity was compared between the three age groups (Fig. 1a to d). The significance of α diversity between age groups depended on the analyses. For bacterial communities in the gut, no significant difference in Chao1 index values was found among the three age groups, while significantly higher Shannon diversity index values were found in the elderly group than in the young and centenarian groups (Fig. 1a and b). For bacterial communities in the oral cavity, both analyses revealed that the centenarians had significantly lower α diversity than did the young and elderly, who shared similar α diversity (Fig. 1a and b). For bacterial communities in the umbilicus, significantly higher α diversity was found in the elderly than with the young and centenarians (Fig. 1a and b). For the fungal communities in the oral cavity and gut, the variation of α-diversity values between the three age groups was not significant (Fig. 1c and d). For fungal communities in the umbilicus and forehead, significantly higher α diversity was found in the elderly than with the young and centenarians; also, the fungal communities in the palms also displayed higher α diversity in the elderly than did those in the young and centenarians (Fig. 1c and d).

**Bacterial and fungal community β diversity in the different body sites.** Principal-coordinate analysis (PCoA) based on the Bray-Curtis distance of the microbiota compositional profiles at the genus level revealed that the primary clustering of the bacterial community in the body was driven by body site (Fig. 2a). Compared to the skin, where the communities were rather loosely clustered, the gut and oral communities were densely clustered. The Adonis test showed that the bacterial communities were significantly clustered by individual subjects (\( R^2 = 25\%, \ P = 0.001 \)). Meanwhile, the grouping of bacterial communities by body site was also statistically significant (\( R^2 = 21\%, \ P = 0.001 \)), with gut bacterial communities forming a cluster away from bacterial clusters of the skin and oral cavity. The clustering of fungal communities in the PCoA plots was less distinctly separated by body site than that of bacterial communities (Fig. 2b); however, the Adonis test showed that the grouping of fungal communities was statistically significant both for each body site (\( R^2 = 15\%, \ P = 0.001 \)) and in each

| TABLE 1 Clinical characteristics of the Sardinian cohort |
|-----------------------------------------------|
| **Characteristic** | **Data for population (n):** |
|                  | Centenarians (22) | Elderly (24) | Young (19) |
| Age (mean ± SD [range]) (yr)               | 102.0 ± 1.5 (99–107) | 77.2 ± 5.9 (68–88) | 24.9 ± 3.5 (19–33) |
| No. of males | 6 | 10 | 7 |
| No. of females | 16 | 14 | 12 |
individual ($R^2 = 30\%, P = 0.001$). Overall, although the human microbiota is grouped by body habitats, they are highly specific to an individual.

To assess the effect of different body habitats on community variation, the compositional differences of bacterial and fungal communities among individuals ($\beta$ diversity) for each body site were assessed (Fig. 2c and d). The distances within the defined body site revealed the different degrees of community composition dissimilarity. The Kruskal-Wallis test revealed that the variation in bacterial community compositions between individuals in the palms was significantly larger than that of the other body sites, while the variation was smallest in the oral cavity ($P < 0.05$). The fungal community compositional variations within the oral cavity and gut were significantly larger than with the skin (Kruskal-Wallis test, $P < 0.05$).

**Bacterial and fungal community $\beta$ diversity in the different age groups across body sites.** Analysis of similarity (ANOSIM) and a multiresponse permutation procedure (MRPP) test were applied to test statistically whether the microbiota communities were significantly different among the three age groups (Table 2). The tests revealed that the three age groups showed different degrees of subclustering in certain habitats. The skin and gut bacterial communities were statistically different for different age groups, but oral bacterial communities were not statistically different for different age groups. No significant difference among the three age groups was detected for the fungal com-
FIG 2 Bacterial and fungal community β diversity in different body sites. (a and b) PCoA of Bray-Curtis distances for the bacterial (a) and fungal (b) communities in each individual across the body sites at the genus level. Samples in each body site are clustered by 95% confidence ellipses. (c and d) Density distributions of bacterial and fungal Bray-Curtis dissimilarities. (e) Box plots of Bray-Curtis distances for bacterial and fungal communities across body sites for different age groups. (Continued on next page)
TABLE 2 Statistical evaluation of the differences in bacterial and fungal communities among age groups and body sites

| Community (sequencing) | Body site | ANOSIM | MRPP test |
|------------------------|-----------|---------|------------|
|                        |           | Test statistic | \(P\) value | Observed \(\Delta\) | Expected \(\Delta\) | \(C\) \(\Delta\) | \(E\) \(\Delta\) | \(Y\) \(\Delta\) | \(P\) value |
| Bacteria (16S rRNA)    | L         | 0.225   | 0.001     | 0.75     | 0.79     | 0.749    | 0.806   | 0.681    | 0.001     |
|                        | R         | 0.208   | 0.001     | 0.76     | 0.79     | 0.758    | 0.798   | 0.721    | 0.001     |
|                        | F         | 0.128   | 0.001     | 0.65     | 0.67     | 0.729    | 0.618   | 0.577    | 0.001     |
|                        | U         | 0.127   | 0.003     | 0.76     | 0.78     | 0.783    | 0.738   | 0.766    | 0.001     |
|                        | O         | 0.019   | 0.185     | 0.54     | 0.55     | 0.598    | 0.528   | 0.502    | 0.008     |
|                        | G         | 0.176   | 0.001     | 0.76     | 0.79     | 0.819    | 0.764   | 0.678    | 0.001     |
| Fungi (ITS1)           | L         | 0.091   | 0.002     | 0.66     | 0.68     | 0.635    | 0.762   | 0.546    | 0.001     |
|                        | R         | 0.083   | 0.009     | 0.67     | 0.7      | 0.612    | 0.773   | 0.616    | 0.001     |
|                        | F         | 0.088   | 0.003     | 0.54     | 0.57     | 0.563    | 0.609   | 0.428    | 0.001     |
|                        | U         | 0.08    | 0.015     | 0.65     | 0.67     | 0.648    | 0.707   | 0.572    | 0.001     |
|                        | O         | 0.029   | 0.125     | 0.84     | 0.84     | 0.829    | 0.848   | 0.801    | 0.071     |
|                        | G         | 0.04    | 0.087     | 0.85     | 0.85     | 0.850    | 0.860   | 0.790    | 0.054     |

**a** L, left palm; R, right palm; F, forehead; U, umbilicus; O, oral cavity; G, gut.

**b** ANOSIM and MRPP tests based on the Bray-Curtis distance at the genus level were used to statistically evaluate the differences in bacterial and fungal communities among the three age groups in each body site. The number of permutations is 999. \(P\) values of \(<0.05\) are highlighted in bold.

The fungal communities for the skin habitats showed a significant difference among age groups, but even so, the group dissimilarities were not sharp \(0 < R < 0.1\). To avoid biases related to the unequal distribution of sexes, permutational multivariate analysis of variance (PERMANOVA), ANOSIM, and MRPP test based on the Bray-Curtis dissimilarity at the genus level were used to look at the sex-related difference in microbiota distributions in each body site (Table S1a). Sex displayed a weak contribution to the variability in microbial distribution.

PCoA based on the Bray-Curtis distance of the microbiota compositional profiles at the genus level was used to visualize bacterial and fungal community structures for the different age groups (Fig. S3). We found that the age group clustering matched the ANOSIM and MRPP test results, as shown in Table 2. More specifically, ANOSIM based on the Bray-Curtis distance at the genus level was applied to statistically evaluate the differences between two age groups in six body sites for the bacterial and fungal communities (Table S1b). We observed that for the bacterial communities, the symmetric palms and forehead showed a similar pattern, where the young and centenarians had clearly separated clusters, while the elderly had clusters that overlapped with the young and centenarians (Fig. S3a to c). In the umbilicus sites, the clustering of the bacterial communities for age groups was slightly different, in that elderly and centenarian communities clustered closely but distinct from those of the young (Fig. S3d and Table S1b). For the fungal communities in the skin, the young and centenarians showed close clustering, whereas the elderly clustered separately (Fig. S3g to j and Table S1b).

Host health parameters may act as covariates and associate with the host gut microbiota composition. We next used envfit analysis to determine the significant
factors that can explain the gut microbiota variance (Table S1c). Several covariates were significantly associated with the gut bacterial community, with the functional independence measure (FIM) and the Mini-Mental State Examination (MMSE) indexes explaining the greatest amount of variance. Conversely, the factors we measured had no significant association with gut fungal community.

To investigate the age-related effects on community variation, the compositional differences of bacterial and fungal communities among individuals (β diversity) in the three age groups were assessed for each body site (Fig. 2e). We observed considerable variation in the bacterial and fungal community structures among individuals in the different age groups, with the exception of the umbilicus and oral cavity bacterial and gut fungal communities (Fig. 2e). For example, the variation in bacterial community compositions between centenarian individuals was significantly larger than those of the elderly and young, and the variation was smallest in the young (Kruskal-Wallis test, $P < 0.05$), while the fungal community compositional variation within each age group was not significantly different in the gut (Kruskal-Wallis test, $P > 0.05$).

**Bacterial and fungal community taxonomic compositions in the three age groups across the different body sites.** To obtain a taxonomic overview of the bacterial and fungal communities, a genus-level relative abundance comparison was performed across the six body sites in the three age groups (Fig. 3 and S4). The resulting profiles showed that each body habitat had a characteristic compositional pattern for both bacteria and fungi (Fig. 3). The dominant bacteria in the skin sites were *Propionibacterium*, *Staphylococcus*, and *Corynebacterium* spp. (Fig. 3a), while the dominant fungi were *Malassezia* spp. (Fig. 3b). The dominant bacteria in the oral cavity were *Streptococcus*, *Veillonella*, *Prevotella*, *Rothia*, and *Actinomyces* spp. (Fig. 3a), while the dominant fungi in the oral cavity were *Malassezia*, *Candida*, and *Saccharomyces* spp. (Fig. 3b). The dominant gut bacteria were *Bacteroides*, *Faecalibacterium*, *Blautia*, *Coprococcus*, and *Bifidobacterium* spp. (Fig. 3a), whereas *Penicillium* and *Saccharomyces* spp. were the dominant fungi in the gut (Fig. 3b).

Diversification of the bacterial and fungal community compositional signatures in the three different age groups was also observed (Fig. 3 and S4). For example, in the oral cavity, the dominant genus profiles were similar among the three age groups in the bacterial community (Fig. 3a and S4i). The dominant fungal genus profiles in the palms was similar for the young and centenarians but different from that for the elderly, which is in part due to the remarkably low average relative abundance of *Malassezia* spp. in the elderly (Fig. 3b and S4d and f). Several genera also had a wide range of average relative abundance in different age groups. For example, the average relative abundance of *Propionibacterium* spp. in the forehead for the young was 0.54, while it was 0.39 for the elderly and 0.21 for the centenarians (Fig. 3a). *Saccharomyces* spp. had an average relative abundance in the gut for the elderly and centenarians (0.10) similar to that for the young (0.27) (Fig. 3b).

To study which bacterial and fungal taxa had significant differences in abundance among the three age groups, the linear discriminant analysis (LDA) effect size (LEfSe) was determined. The results showed that 27 bacterial genera and 15 fungal genera had differential abundance in the three age groups (Fig. 4). For example, overrepresentation of *Propionibacterium* spp. was observed in the four skin sites (left palm, right palm, forehead, and umbilicus) for the young, while *Prevotella*, *Rothia*, and *Veillonella* spp. were enriched in the left palm, right palm, and forehead for the centenarians. In the gut, *Blautia*, *Roseburia*, and *Faecalibacterium* spp. were enriched in the young, *Prevotella* and *Ruminococcus* spp. were enriched in the elderly, and *Parabacteroides*, *Lactobacillus*, and *Clostridium* spp. were enriched in the centenarians. In the oral cavity, *Haemophilus* spp. were enriched in the young. Interestingly, we observed a significant enrichment of *Bifidobacterium* spp. in the left palms of the centenarians (Fig. 4a). The enrichment of *Bifidobacterium* spp. could be detected in other body sites of centenarians, although not all enrichment reached statistical significance (Fig. S5). Fungi showed accumulation in the skin in the elderly compared with the young and centenarian groups (Fig. 4b). For example, *Debaryomyces* and *Penicillium* spp. were enriched in...
Malassezia spp. were enriched in the left palm, forehead, and umbilicus for the young while in the right palm for the centenarians (Fig. 4b).

**Bacterial and fungal correlations at the community and genus levels.** To explore potential correlations between bacterial and fungal community diversity in the human body, community diversity was evaluated by the Shannon diversity index for each clinical sample within each body site (Fig. 5a). The Shannon diversity values for the bacterial and fungal communities in the gut and oral cavity were not statistically correlated, but a significant correlation was observed in the skin. The linear regression analysis indicated that for the skin, individuals with high diversity in the bacterial community were also associated with a high diversity in the fungal community. In the gut and oral sites, the diversity of bacterial and fungal communities appeared to be independent.

To determine the correlations between bacterial and fungal community dissimilarities between any two clinical samples within each body site, linear regression analysis and Mantel tests were performed between fungal and bacterial Bray-Curtis dissimilarity pairwise comparisons (Fig. 5b and Table S2). Interestingly, the greatest positive correlation was detected in the palms, indicating that two individuals with similar bacterial
communities were associated with a similar fungal community in palms. For other body sites, the cross-domain community correlations were weak or insignificant.

Cooccurrence analysis of each body habitat, based on Pearson’s correlation of fungal and bacterial taxonomic relative abundances, provided a preliminary evaluation of major fungus-bacterium genus associations (Fig. S6). It revealed that for different niches, the correlations of the bacterial and fungal taxa were diverse. In the skin, bacteria and fungi exhibit greater numbers of significant bacterium-bacterium, fungus-fungus, and cross-domain correlations, while in the oral cavity, most of those correlations belong to bacterium-bacterium correlations. For the gut, a small number of significant correlations were observed, but the number of cross-domain correlations was greater than those observed in the oral cavity. We also observed that the correlation between the same taxa in different niches can be distinct. For instance, Candida and Malassezia spp. had a significant negative correlation in the skin and oral cavity, but they did not show a significant correlation in the gut. In the gut, the strongest positive correlation occurred between Meyerozyma and Rhodotorula spp., Ruminococcus and Wallemia spp., and Dialister and Trichosporon spp. In the oral cavity, Rhodotorula spp. showed a significant positive correlation with Cryptococcus spp.; however, no significant correlation was observed with Meyerozyma spp., which was the case in the gut. A strong negative correlation was found between Prevotella and Streptococcus spp. in the oral cavity, but only a weak positive correlation was observed in the skin.

Comparative analysis of the age-related gut bacterial communities in different populations. To investigate the age-related features of gut bacterial community across populations, we compared our results with two previous studies from Italy and Japan which all targeted the 16S rRNA V3-V4 region (24, 26) (Fig. S7). In the PCoA plot, the distribution of extreme aging individuals in different populations shifts in the same direction away from the young and elderly and was conserved in the Italian and Japanese cohorts, similar to what we observed here (Fig. S7a). In comparing the Sardinian cohort with the cohort from Emilia-Romagna, Italy, we found that several age-related features were shared between the two populations at the genus level for the gut bacterial community (Fig. S7b). For example, lower abundances of Faecalibacterium spp. but higher abundances of Bifidobacterium spp. in centenarians were observed compared with the young and elderly in both populations.
DISCUSSION

Our study here systematically demonstrated age-related bacterial and fungal community variations in different body sites within a single population. Concurrent analysis of bacterial and fungal communities revealed that the body habitats and age differentially shape these two microbial communities. Our study provides a framework for future mechanistic investigation of age-related microbiota adaptation and interactions between fungal and bacterial communities in the human body.

We further demonstrated that the distribution of the microbiota in Sardinians was largely determined by body site and individual, consistent with the previous studies (2, 39). We also showed the various age-related bacterial and fungal community diversity measures in each body site. For example, the significantly low $\alpha$ and $\beta$ diversity of bacterial and fungal communities in the forehead of the young could be driven by the highly similar skin microbiome and mycobiome among the young with the presence of

FIG 5 Correlations between bacterial and fungal communities. (a) Correlation of bacterial and fungal community Shannon diversity indexes in each body site. The samples are differently shaped for each body site and colored by age groups labeled by C (centenarian), E (elderly), and Y (young). (b) Correlation of fungal and bacterial Bray-Curtis dissimilarity pairwise comparisons at the genus level within each body site. The adjusted (adj) $R^2$ of the linear regression analysis and Pearson correlation coefficient ($r$) are provided for each plot. All $P$ values in panel b are $<0.0001$. L, left palm; R, right palm; F, forehead; U, umbilicus; O, oral cavity; G, gut.
predominant species in the communities, while the remarkable high $\alpha$ and $\beta$ diversity of bacterial and fungal communities in the palms of the elderly indicated their highly personalized and complex community structures. Interestingly, in our cohorts, the centenarians have $\alpha$ diversity similar to that of the young, while the elderly had a higher Shannon diversity than that of the young and centenarians. A previous study in U.S., U.K., and Colombian cohorts had observed a positive association between $\alpha$ diversity and age but not in Chinese cohorts (40). It was also observed in a larger Irish cohort that the elderly displayed greater interindividual variation ($\beta$ diversity) than did younger adults for gut microbiota (20). The dynamic age-related $\alpha$- and $\beta$-diversity features across populations indicate that how aging shapes the gut microbiota may remain an open question.

Although we showed different degrees of variation in the taxonomic compositions of bacterial and fungal communities among the different age groups, bacterial and fungal communities in the skin sites all displayed an age-related clustering, consistent with findings from a previous study in Japan but differing from a study in China (33, 34). In the four skin sites we surveyed, bacterial communities between young and centenarians are significantly different, while fungal communities were more similar between the young and centenarians but distinct from those of the elderly. The skin bacterial and fungal community variations in different age groups were related not only to age; it is possible to assume additional factors, such as host physiological capabilities, environmental exposure, and different nutritional requirements for bacteria and fungi may contribute to the divergence.

The bacterial and fungal species with significantly different abundances among the three age groups were also identified in the skin. *Propionibacterium* spp. were enriched in the skin for the young compared with the elderly and centenarians, which could correlate with the decrease in the sebaceous gland activity in skin sites associated with aging (11), consistent with previous studies (33, 41). Several groups have explored the direct interactions between the skin microbiota and the host immune system (42, 43). High abundances of potentially pathogenic *Staphylococcus* and *Streptococcus* spp. were detected in the skin sites of centenarians, which suggests that further attention should be paid to the possible infection of these potentially pathogenic genera in centenarians. *Malassezia* spp., as dominant lipophilic fungi in human body, declined in the elderly compared with the young in our study, different from a previous study in a Toronto population where the elderly had the highest abundance of *Malassezia* spp. (44). Intriguingly, the centenarian group showed a higher abundance of *Malassezia* spp. than did the elderly, which suggests that the sebum in the skin was not the only determinant of *Malassezia* sp. colonization.

Evidence shows that specific microbes in the oral cavity are associated with oral health and disease and are even linked with other systemic diseases (45, 46). We found that both the bacterial and fungal communities in the oral cavity did not display distinct age-related clustering, although age-associated oral bacterial community structural changes were found in a Chinese cohort (37).

Among those microbial communities distributed across the body, the gut microbiota is the most comprehensively studied microbial community and has the strongest association with human disease and health (8). Numerous recent studies indicated that the gut microbiota had a major impact on the host metabolic status and can regulate host life span in animal models (47–50). With aging, the physiological decline of the host gradually causes dietary alterations and decreased immunity health, consequently leading to changes in the gut microbiota (51, 52).

Our data confirmed the results found in the NIH Human Microbiome Project (HMP) cohort which identified the basic fungal and bacterial community characteristics in the gut, such as the fungal diversity being lower than bacterial diversity, the independent relationship between the $\alpha$ diversity of the bacterial and fungal communities, and the dominant fungal genera, *Saccharomyces*, *Malassezia*, and *Candida* (53). In our study here, the Sardinian population showed certain specific features different from those of the HMP cohort. The dominant genera in the gut, for example, *Penicillium*, are higher
in mean relative abundance in the gut in the Sardinian cohort (>10%) than in the HMP cohort (<4%). On the other hand, the abundance of Cladosporium spp. was lower in the Sardinian cohort. The cause of these population differences in the human microbiota is unknown; however, environmental factors such as geography, ethnicity, diet, and lifestyle may be contributing factors (3–6). It must be pointed out that technical differences in collection and processing, as well as analysis of data, may also contribute to the observed differences.

We observed that the young and elderly had similar bacterial gut community compositions, consistent with previous studies (21, 27), but different from the bacterial gut community composition of most of the centenarians. Similar results were also obtained by our previous metagenomics sequencing (28), as well as the results from an Emilia-Romagna (Italy) cohort and a Japanese cohort (24, 26). The separated clustering of the gut bacterial communities in the centenarians across populations suggested that age shaped the gut bacterial community following the same trajectory with the aging process. For example, the low abundances of Faecalibacterium and Roseburia spp. (23, 24, 26, 54) in centenarians were detected in our study and other studies (23, 24). Parabacteroides spp. were enriched in the centenarian cohort in Sardinia and a Chinese centenarian cohort in Sichuan but decreased in another Chinese centenarian cohort in Guangxi (22, 23). Bifidobacterium and Lactobacillus spp. are believed to have health-promoting properties as probiotics (55, 56). However, previous studies reported the low abundances of Bifidobacterium and Lactobacillus spp. in the elderly (57, 58). However, high abundances of Bifidobacterium and Lactobacillus spp. were detected in our centenarians but not the elderly compared with the young, as well as the centenarians from Guangxi, China, and Emilia-Romagna, Italy (22, 24), suggesting an interesting gut microbiota feature of human longevity. Moreover, we also observed the enrichment of Bifidobacterium spp. in other body sites of the centenarians in Sardinia. We believe a larger cohort may be needed to confirm the importance and statistical significance of this enrichment in centenarians.

Compared to bacterial communities, the gut fungal communities in Sardinians displayed low α diversity and great individual variation, consistent with the HMP population from Houston, TX (53). However, diverse dominant species among populations were also observed. For example, Penicillium and Debaryomyces spp., which are found in fermented foods but cannot grow in the gut niche (59), had a higher prevalence in Sardinians than in the HMP population and may correspond to the high consumption of cheese in Sardinians (60). Besides, Debaryomyces spp. were significantly enriched in the young group compared with the elderly and centenarians in Sardinia. The variation in Debaryomyces spp. may partly be caused by the dietary preferences for different age groups. Previous studies have shown that diet can modulate the fungal communities in the murine gut (61). The subjects dominated by resident fungi may be associated with a diet enriched of fungi, further suggesting that caution should be made in the microbiome studies because separating the viable microbes from the nonliving microbes and resident from transient microbes can be difficult (62).

Our preliminary survey of the associations between clinical parameters and gut microbiota indicated that host health status-related factors such as FIM and MMSE could explain the greatest amount of variance among the three age groups of bacterial community, consistent with a previous study that also showed the correlation between gut microbiota and health (25). Moreover, the significant association between mini nutritional assessment (MNA) and gut bacterial community suggests that diet and nutrition could also act as important covariates contributing to the variance of the microbiome composition. Diet is considered one of the main determinants of the gut microbiota (63). Further longitudinal studies tracing how aging shapes the gut microbiota with the dietary and environmental factors will be necessary to determine the role those microbes play in the gut during aging.

With an integrated view of microbiota in different body habitats, our preliminary evaluation of the correlation between fungi and bacteria across body sites at both the community and genus levels revealed that the correlations were site and population
specific. At the community level, we detected positive correlations between the Shannon diversity of fungal and bacterial communities in each individual in all four skin sites but not in the oral cavity or the gut. This finding is consistent with previous studies of the skin microbiota in a Hong Kong cohort and the HMP gut microbiota study in a Houston, TX, cohort (53, 64). We detected the greatest significant positive cross-domain correlations in community composition dissimilarities in the palms. Interestingly, a skin microbiome study in a Hong Kong cohort also identified significant positive cross-domain correlations in the skin, especially in the palms (64). Although it seems that those cross-domain correlations are shared across populations, the underlying mechanisms are currently unknown. At the genus level, lipophilic Malassezia and Propioni bacterium showed a significant strong positive association in the skin, reflecting their similar nutrition requirements; in the mucosal habitats, however, the correlation was not statistically strong. Furthermore, in the skin, Actinomyces spp. were positively correlated with Staphylococcus and Streptococcus spp., but in the oral cavity, the correlation was negative. Also, for different populations, the cooccurrence pattern was different. In our study, a strong negative association was detected between Prevotella and Streptococcus spp. in the oral cavity, consistent with findings from an Italian study (65), while in a Japanese study, the correlation was positive (36). In the gut, the association relationship we observed, such as the positive association between Faecalibacterium and Saccharomyces spp., was also found in a U.S. study (61), although the correlation we detected based on the cooccurrence did not necessarily mean a true biological interaction between the species. Further cultivation assays are needed to gain insights into the importance and mechanisms of these correlations.

In summary, our study provides a picture of the diverse features of bacterial and fungal communities in different body sites and refines our understanding of the age-related human microbial community variations. However, to better understand the underlying mechanisms of age-related microbiota variations, we should focus more on the detailed functional descriptions of characteristics of diet, living conditions, community populations, as well as individual physiological characteristics and measurement of the variables for microbial ecological niches (e.g., pH range, dissolved oxygen range, temperature range, salinity range, etc.). Fundamental questions will be addressed about the age-related adaptation and diversification of the human microbiota under controlled environments with well-defined niches.

MATERIALS AND METHODS

Subject recruitment and clinical information collection. We recruited 65 subjects in Sardinia, Italy, as a part of the AKEA project, which studies the extreme longevity in Sardinia (38). Ethics approval was provided by the institutional local ethics committee of the Azienda Sanitaria Locale N.1 of Sassari, Italy.

The donors were volunteers recruited from the longevity AKEA project, and participants signed a written consent form. Subjects were divided into three age groups, young, elderly, and centenarians. Exclusion criteria for the young group and the elderly group included the following: (i) history of chronic medical conditions (diabetes and hypertension), (ii) the use of antimicrobial medication (antibiotic or antifungal treatments) 1 year before sampling, and (iii) with chronic dermatologic diseases (psoriasis, atopic dermatitis, vitiligo, or urticaria). Clinical and nutritional data were collected as described in the AKEA study (38).

Sample collection and DNA extraction. (i) Skin samples. To maximize microbial load, no prior cleaning of the skin was needed before sample collection. The skin samples were collected by professional staff with sterilized swabs (Catch-All sample collection swabs) at four different sites, including the forehead, two palms, and the umbilicus area. A 5- by 5-cm² area of skin was gently rubbed 10 times using a swab premoistened with sterilized enzyme lysis buffer. Each swab tip was placed into an Eppendorf tube containing 200 µl enzyme lysis buffer. The samples were kept on ice while shipping to the lab and were stored at −80°C. Negative-control specimens were collected by exposing swabs to room air and then were processed with the samples. Extracted DNA from the tip of the swabs was performed according to the manual’s instructions for the DNeasy blood and tissue extraction kits, with some modification. In brief, the samples in the lysis buffer were incubated at 37°C for 2 h. Five-millimeter zirconia beads (0.4 g; Sigma) were added. Then, each sample was subjected to a bead-beating step using multivortex V-32 (Biosan) at a maximum of 3,000 rpm for 30 min. Twenty-five microliters of proteinase K was added and incubated at 56°C for 30 min. Then, samples were heated at 95°C for 5 min, followed by ice for 1 min, and performed according to the DNeasy blood and tissue extraction kit protocol. The DNA was eluted with 200 µl Tris-EDTA (TE) buffer. The final DNA concentration was determined using an ND-1000 spectrophotometer (NanoDrop Technologies).
(ii) Oral samples. Participants were asked to avoid eating or drinking for 1 h prior to oral cavity sampling. Subjects were asked to let saliva collect in the mouth for at least 1 min. The subject was then asked to drool into a sterilized labeled 50-ml collection tube. This process may be repeated multiple times in order to collect larger volumes of saliva (2 to 5 ml). For the centenarians, it was difficult to collect the saliva, so oral washing samples were collected instead. Centenarians were asked to swish vigorously with 10 ml sterilized water for 30 s and then to expectorate into the tube. The wash was repeated twice. After shipping the samples on ice to the lab, they were centrifuged at 7,500 rpm for 10 min, the supernatant was discarded, and the DNA was extracted from the sediment following the same protocol as with skin samples.

(iii) Stool samples. Fecal samples were collected by the participants at home. Participants were provided with a sterilized stool specimen collection tube. After passing stool, a spoon was used to collect about a 1-g stool sample by scraping off the outer layer of solid feces and collecting the central part into the tube. Samples were immediately frozen at home at ~20°C and collected by laboratory personnel within 6 weeks. Long-term storage of samples was in a ~80°C freezer located at the University of Sassari. DNA was extracted from stool samples according to the manual instructions for the QIAamp DNA stool minikit (Qiagen), with some modifications. In brief, 200 mg of stool was suspended in 1.4 ml of buffer ASL, and 0.4 g of 5-mm zirconia beads (Sigma) was added. Then, each sample was subjected to a bead-beating step using Biosan at a maximum of 3,000 rpm for 30 min. Samples were heated at 95°C for 5 min and then centrifuged for 5 min at 13,000 rpm to pellet the stool particles. Next, 1.2 ml of supernatant was collected, the InhibitEX tablet was added, which was followed by incubation at room temperature (RT) for 1 min and centrifugation at 13,000 rpm for 3 min; then, 15 µl of proteinase K and 200 µl of buffer AL were added. DNA was isolated and incubated at 70°C for 10 min. Two hundred microliters of absolute ethanol was then added to the mixture, vortexed, and loaded on QIAamp mini spin columns. The columns were washed with buffer AW1 and buffer AW2, as per the QIAamp DNA stool minikit instructions. The DNA was eluted with 200 µl TE buffer. Finally, the DNA concentration was determined by using an ND-1000 spectrophotometer (NanoDrop Technologies).

Library construction and sequencing. Procedures for 16S rRNA and ITS1 library generation were performed as previously described (66). Briefly, the V3-V4 regions of the 16S rRNA gene and ITS1 gene were amplified using an improved dual-indexing approach. The primers for 16S rRNA amplification were 5'-ACTCCTACGGGAGGCAGCAG-3' and 5'-GACTACHVGGGTWTCTAAT-3' (67), and for ITS1 amplification, the primers were 5'-GTAAAGTCGTAACAAGGTTTTC-3' and 5'-GTCGACTAAGYTCGATGATTCAC-3' (41). 16S rRNA and ITS1 libraries were, respectively, normalized and pooled using the SequalPrep normalization plate kit (Invitrogen) prior to sequencing on an Illumina MiSeq platform.

Bioinformatics for 16S and ITS1 sequencing. Raw sequence reads were first trimmed by removal of the barcodes and linker sequences. Then, USEARCH was used for truncation of the reads not having an average quality of 15 based on the phred algorithm. Reads with less than 75% of their original length were removed (68). Further read processing was performed using QIIME (version 1.9.1) (69). Reads were clustered into operational taxonomic units (OTUs) at 97% similarity using the open-reference strategy by uclust based on the Greengenes database (2013_8 version) for 16S rRNA and UNITE fungal ITS database (version 7.2) (70, 71). The first cluster seed was chosen as the representative sequences for the OTUs. uclust for 16S rRNA sequencing and BLAST for ITS1 sequencing were used for taxonomic assignment of the representative sequences. Alignment of representative sequences was performed using PyNAST for 16S rRNA sequencing and MUSCLE for ITS1 sequencing. The aligned sequences were filtered and phylogenetic trees constructed using FastTree. Following rarefaction, an OTU table was generated in QIIME to minimize the difference in sequencing depths across samples (5,000 reads per sample for 16S rRNA libraries, 10,000 reads per sample for ITS1 libraries, and 10,000 reads per sample for cross-population gut bacterial community comparison). Rarefaction OTU counts were binned into genus-level taxonomic groups according to the taxonomic assignments described earlier. α-diversity indexes, including Chao1 richness and the Shannon diversity index, were calculated using QIIME.

Statistical analysis. All statistical analyses were performed using the R software (version 3.4.2). Multivariate analysis of community diversity employing PCoA was performed using vegan and visualized using ggplot2. A Bray-Curtis distance matrix was used as the dissimilarity index. LEfSe was performed to detect differentially abundant taxa between age groups using the default parameters (72). Only those taxa that showed a P value of <0.05 and the threshold logarithmic LDA score of ≥3 were ultimately considered. Kruskal-Wallis tests followed by Dunn's multiple-comparison test were used to determine whether significant differences existed among multiple groups (73). Linear regression was used to test the correlation between two variables (Shannon diversity index and Bray-Curtis distances). Discrimination among groups was detected by MRPP, ANOSIM, and the Adonis method in vegan in R. All permutation tests (i.e., MRPP, ANOSIM, Adonis, and the Mantel test) were conducted using 999 permutations. The significance and explained variance of covariate (referenced to as clinical parameters in Table S1c) were determined using the envfit function in vegan.

Data availability. The data sets generated and analyzed during the current study are available in the European Nucleotide Archive (ENA) under study accession number PRJEB25916.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, EPS file, 0.7 MB.
FIG S2, EPS file, 1.1 MB.
FIG S3, EPS file, 1.9 MB.
ACKNOWLEDGMENTS

We are grateful to Nikki Kelvin for editorial comments and assistance.

Funding support was provided by LKSF (to D.J.K.), the International Institute of Infection and Immunity, Shantou University Medical College (to D.J.K.), the Canadian Foundation for Innovation, the Dalhousie Medical Research Foundation (to D.J.K.), Ministero dell’Istruzione, dell’Università e della Ricerca (MIUR, Italy) Progetti di Ricerca di Rilevante Interesse Nazionale (PRIN) 2015 (protocol 20157ATSLF_002), and Consiglio Nazionale delle Ricerche Flagship InterOmics (code PB05) (to C.C. and A.Z.), and an NSERC discovery grant (to M.G.I.L.). D.J.K. is a recipient of a Tier I Canada Research Chair in Vaccinology and Inflammation.

L.W., S.R., D.J.K., and C.C. developed the study concept and design; L.W. and C.C. collected samples and acquired the data; L.W. performed data analysis; M.D., T.Z., A.Z., and L.M. contributed reagents, materials, and analysis tools; and L.W., M.G.I.L., and D.J.K. wrote the manuscript. All authors read and approved the final manuscript.

We declare no competing financial, professional, or personal interests that might have influenced the performance or presentation of the described work.

REFERENCES

1. Caporaso JG, Lauber CL, Costello EK, Berg-Lyons D, Gonzalez A, Stombaugh J, Knights D, Ager J, Pena R, Fierer N, Gordon JI, Knight R. 2011. Moving pictures of the human microbiome. Genome Biol 12:R50. https://doi.org/10.1186/gb-2011-12-5-r50.

2. Costello EK, Lauber CL, Hamady M, Fierer N, Gordon JI, Knight R. 2009. Bacterial community variation in human body habitats across space and time. Science 326:1694–1697. https://doi.org/10.1126/science.1177486.

3. Gupta VK, Paul S, Dutta C. 2017. Geography, ethnicity or subsistence-ties. ISME J 9:1979–1990. https://doi.org/10.1038/ismej.2015.11.

4. Yatsunenko T, Rey FE, Manary MJ, Trehan I, Dominguez-Bello MG, Contreras M, Magris M, Hidalgo G, Baldassano RN, Anokhin AP, Heath AC, Warner B, Reeder J, Kuczynski J, Caporaso JG, Lozupone CA, Lauber C, Clemente JC, Knights D, Knight R, Gordon JI. 2012. Human gut microbiome viewed across age and geography. Nature 486:222–227. https://doi.org/10.1038/nature11053.

5. Zhang J, Guo Z, Xue Z, Sun Z, Zhang M, Wang L, Wang G, Wang F, Xu J, Cao H, Xu H, Lu Q, Zhong Z, Chen Y, Qimugue S, Menghe B, Zheng Y, Zhao L, Chen W, Zhang H. 2015. A phylo-functional core of gut microbiota in healthy young Chinese cohorts across lifestyles, geography and ethnicity. ISME J 9:1979–1990. https://doi.org/10.1038/ismej.2015.11.

6. Oh J, Byrd AL, Deming C, Conlan S, NISC Comparative Sequencing Program, Kong HH, Sege JA. 2014. Biogeography and individuality shape function in the human skin metagenome. Nature 514:59–64. https://doi.org/10.1038/nature13786.

7. Cadwell K. 2015. The virome in host health and disease. Immunity 42:805–813. https://doi.org/10.1016/j.immuni.2015.05.003.

8. Lynch SV, Pedersen O. 2016. The human intestinal microbiome in health and disease. N Engl J Med 375:2369–2379. https://doi.org/10.1056/NEJMra1600266.

9. Thomas-White K, Forster SC, Kumar N, Van Kuiken M, Putluri N, Ajami NJ, Putluri N, Graf J, Bryan RM, McCullough LD. 2018. Age-related changes in the gut microbiota influence systemic inflammation and stroke outcome. Ann Neurol 84:23–36. https://doi.org/10.1002/ana.25250.

10. Brandsma E, Kloosterhuis NJ, Koster M, Dekker DC, Gijbels MJ, van der Velden S, Rios-Morales M, van Faassen MJ, Loreti MG, de Bruijn A, Fu J, Kuipers F, Bakker BM, Westerterp M, de Winther MPJ, Hofker MH, van Sluis B, Kooien D. 2019. A proinflammatory gut microbiota increases systemic inflammation and accelerates atherosclerosis. Circ Res 124:94–100. https://doi.org/10.1161/CIRCRESAHA.118.313234.

11. Heintz C, Mair W. 2014. You are what you host: microbiome modulation of the aging process. Cell 156:408–411. https://doi.org/10.1016/j.cell.2014.01.025.

12. Grice EA, Segre JA. 2011. The skin microbiome. Nat Rev Microbiol 9:244–253. https://doi.org/10.1038/nrmicro2537.

13. An R, Wilms E, Masdiee AAM, Smidt H, Zoetendal EG, Jonkers D. 2018. Age-dependent changes in GI physiology and microbiota: time to reconsider? Gut 67:2213–2222. https://doi.org/10.1136/gutjnl-2017-315542.

14. Desai MS, Seekatz AM, Koropatkin NM, Kamada N, Hickey CA, Wolter M, Pudlo NA, Kitamoto S, Terrapon N, Muller A, Young VB, Henriissat B, Wilmes P, Stappenbeck TS, Nunez G, Martens EC. 2016. A dietary fiber-deprived gut microbiota degrades the colon mucos barrier and enhances pathogen susceptibility. Cell 167:1339–1353.e21. https://doi.org/10.1016/j.cell.2016.10.043.

15. Wolcott RD, Hanson JD, Rees EJ, Koenig LD, Phillips CD, Wolcott RA, Cox SB, White JS. 2016. Analysis of the chronic wound microbiota of 2,963 patients by 16S rDNA pyrosequencing. Wound Repair Regen 24:163–174. https://doi.org/10.1111/wrr.12370.

16. Spychala MS, Venna VR, Jandzinski M, Doran SJ, Durgan DJ, Ganesh BP, Ajami NJ, Putluri N, Graf J, Bryan RM, McCullough LD. 2018. Age-related changes in the gut microbiota influence systemic inflammation and stroke outcome. Ann Neurol 84:23–36. https://doi.org/10.1002/ana.25250.

17. Brandsma E, Kloosterhuis NJ, Koster M, Dekker DC, Gijbels MJ, van der Velden S, Rios-Morales M, van Faassen MJ, Loreti MG, de Bruijn A, Fu J, Kuipers F, Bakker BM, Westerterp M, de Winther MPJ, Hofker MH, van Sluis B, Koonen D. 2019. A proinflammatory gut microbiota increases systemic inflammation and accelerates atherosclerosis. Circ Res 124:94–100. https://doi.org/10.1161/CIRCRESAHA.118.313234.

18. Heintz C, Mair W. 2014. You are what you host: microbiome modulation of the aging process. Cell 156:408–411. https://doi.org/10.1016/j.cell.2014.01.025.

19. Thevaranjan N, Puchta A, Schulz C, Naidoo A, Szamosi JC, Verschoor CP, Loukov D, Schenck LP, Cahon HH, Segre JA. 2014. Biogeography and individuality shape function in the human skin metagenome. Nature 514:59–64. https://doi.org/10.1038/nature13786.
Deane J, O'Connor M, Harnedy N, O'Connor K, O'Mahony D, van Sinderen D, Wallace M, Brennan L, Stanton C, Marchesi JR, Fitzgerald AP, Shanahan F, Hill C, Ross RP, O'Toole PW. 2012. Gut microbiota composition correlates with diet and health in the elderly. Nature 488:178–184. https://doi.org/10.1038/nature11319.

19. O'Toole PW, Jeffrey IB. 2015. Gut microbiota and aging. Science 350: 1214–1215. https://doi.org/10.1126/science.aac8469.

20. Wang F, Yu T, Huang G, Cai D, Liang X, Su H, Zhu Z, Li D, Yang Y, Shen P, Mao R, Yu L, Zhao M, Li Q. 2015. Gut microbiota community and its assembly associated with age and diet in Chinese centenarians. J Microbiol Biotechnol 25:1195–1204. https://doi.org/10.4146/jm.1410.10014.

21. Kong F, Hua Y, Zeng B, Ning R, Li Y, Zhao J. 2016. Gut microbiota signatures of longevity. Curr Biol 26:R832–R833. https://doi.org/10.1016/j.cub.2016.08.015.

22. Biagi E, Franceschi C, Rampelli S, Severgnini M, Ostani T, Turroni S, Consolandi C, Quercia S, Scurti M, Monti D, Capri P, Brigidip P, Candela M. 2016. Gut microbiota and extreme longevity. Curr Biol 26:1480–1485. https://doi.org/10.1016/j.cub.2016.04.016.

23. Jeffrey IB, Lynch DB, O'Toole PW. 2016. Composition and temporal stability of the gut microbiota in older persons. ISME J 10: 170–182. https://doi.org/10.1038/ismej.2015.88.

24. O'Toole PW, Tiku M, Suprakara H, Hashikura N, Takahashi S, Xiao JZ, Abe F, Osawa R. 2016. Age-related changes in gut microbiota composition from newborn to centenarian: a cross-sectional study. BMC Microbiol 16:90. https://doi.org/10.1186/s12866-016-0708-5.

25. Bian G, Gloor GB, Gong A, Li J, Jia C, Zhang W, Hu J, Zhang H, Zhang Y, Zhou Z, Zhang J, Burton JP, Reid G, Xiao Y, Zeng Q, Yang K, Li J, Li J. 2017. The gut microbiota of healthy aged Chinese is similar to that of the healthy young. mSphere 2:e00327-17. https://doi.org/10.1128/mSphere.00327-17.

26. Wu L, Zeng T, Zinelu A, Rubino S, Kelvin DJ, Carru C, Wu L, Zeng T, Zinelu A, Rubino S, Kelvin DJ, Carru C. 2019. A cross-sectional study of compositional and functional profiles of gut microbiota in Sardinian centenarians. mSystems 4:e00325-19. https://doi.org/10.1128/mSystems.00325-19.

27. Falony G, Joossens M, Vieira-Silva S, Wang J, Darzi Y, Faust K, Kurilshikov S, David LA, Maurice CF, Carmody RN, Gootenberg DB, Button JE, Wolfe BE, McDonald D, Wang J, Sadowski JD, Kriaucionis S, Li J, Jia H, Spector TD, Jia H. 2016. Shotgun metagenomics of 250 adult twins reveals genetic and environmental signatures of longevity. Curr Biol 26:R832–R833. https://doi.org/10.1016/j.cub.2016.04.016.

28. Shaw MA, Magrini V, Cross TJ, Palumbo CA, Leppert M, Car R, Antinucci F, Mardis ER, Venter JC, Myers EW, Cawley S, Lander ES, Wilson RK, Nickerson DA. 2010. 18 populations sequence with dynamic microbial communities. Environ Microbiol 12:321–337. https://doi.org/10.1111/j.1462-2920.2012.02502.x.

29. Kong F, Cogen AL, Radek KA, Park HJ, Maedler JD, Leichtle A, Ryan AF, Di Nardo A, Gallo RL. 2010. Activation of TLR2 by a small molecule produced by Staphylococcus epidermidis increases antimicrobial defense against bacterial skin infections. J Invest Dermatol 130:2211–2221. https://doi.org/10.1016/j.jid.2010.12.038.

30. Gupta AK, Kohli Y. 2004. Prevalence of Malassezia species on various body sites in clinically healthy individuals representing different age groups. Med Mycol 42:35–42. https://doi.org/10.1080/1369378031000161056.

31. Santos RE, Griffith AL, Dardis SR, Lee AM, Olsen I, Dewhirst FE, Ley EJ, Paster BJ. 2008. Bacteria of dental caries in primary and permanent teeth in children and young adults. J Clin Microbiol 46:1407–1417. https://doi.org/10.1128/JCM.01410-07.

32. Biagi E, Remers RT, Mallat Z, Sacco RL, Jacobs DR, Benensiano J, Tobsch A, Rundeck T, Kayser M, Savitski MN, Desvarieux M. 2015. Periodontal microbiota and phospholipases: The Oral Infections and Vascular Disease Epidemiology Study (INVEST). Atherosclerosis 242:418–423. https://doi.org/10.1016/j.atherosclerosis.2015.07.039.

33. Sonnenburg JL, Backhed F. 2016. Diet-microbiota interactions as modulators of human metabolism. Nature 533:56–64. https://doi.org/10.1038/nature18846.

34. Biagi E, Backhed F. 2012. Functional interactions between the gut microbiota and host metabolism. Nature 489:242–249. https://doi.org/10.1038/nature11152.

35. Han B, Sivaramakrishnan P, Lin CJ, Neve IAA, He J, Tay LWR, Sowa JN, Sizov A, Du G, Wang J, Herman C, Wang MC. 2017. Microbial genetic composition tunes host longevity. Cell 169:1249–1262.e13. https://doi.org/10.1016/j.cell.2017.05.036.

36. Smith P, Willemens D, Popkes M, Metge F, Gandiwa E, Reichard M, Valenzano DR, Smith P, Willemens D, Popkes M, Metge F, Gandiwa E, Reichard M, Valenzano DR. 2017. Regulation of life span by the gut microbiota in the short-lived African turquoise killifish. Elife 6:e27014. https://doi.org/10.7554/eLife.27014.

37. Sovran B, Hugenholtz F, El德merdon A, Van Beyk AA, Graawersen K, Huijskes M, Boekshoten MV, Savekouli HFS, De Vos P, Decker J, Wells JM. 2019. Age-associated impairment of the mucus barrier function is associated with profound changes in gut microbiota and immunity. Sci Rep 9:1437. https://doi.org/10.1038/s41598-018-35228-3.

38. Nagpal R, Mainali R, Ahmad S, Wang S, Singh R, Kavanagh K, Kitzman DW, Kushugulova A, Marotta F, Yadav H. 2018. Gut microbiome and aging: physiological and mechanistic insights. Nutr Healthy Aging: 1–10. https://doi.org/10.3233/NHA-170030.

39. Smith P, Willemsen D, Popkes M, Metge F, Gandiwa E, Reichard M, Valenzano DR, Smith P, Willemsen D, Popkes M, Metge F, Gandiwa E, Reichard M, Valenzano DR. 2017. Regulation of life span by the gut microbiota in the short-lived African turquoise killifish. Elife 6:e27014. https://doi.org/10.7554/eLife.27014.

40. Sonvar B, Hugenholtz F, Eldemerdon A, Van Beek AA, Graawersen K, Huijskes M, Boekshoten MV, Savekouli HFS, De Vos P, Decker J, Wells JM. 2019. Age-associated impairment of the mucus barrier function is associated with profound changes in gut microbiota and immunity. Sci Rep 9:1437. https://doi.org/10.1038/s41598-018-35228-3.

41. Biagi E, Nylund L, Candela M, Ostani T, Bucci L, Pini E, Nikkila J, Monti D, Satalori R, Franceschi C, Bridigip P, De Vos W. 2010. Through ageing, and beyond: gut microbiota and inflammatory status in seniors and centenarians. PLoS One 5:e1001667. https://doi.org/10.1371/journal.pone.01001667.

42. Collins MD, Gibson GR. 1999. Probiotics, prebiotics, and synbiotics:
approaches for modulating the microbial ecology of the gut. Am J Clin Nutr 69:10525–10575. https://doi.org/10.1093/ajcn/69.5.1052s.

56. Pan F, Zhang L, Li M, Hu Y, Zeng B, Yuan H, Zhao L, Zhang C. 2018. Predominant gut Lactobacillus murinus strain mediates anti-inflammatory effects in calorie-restricted mice. Microbiome 6:54. https://doi.org/10.1186/s40168-018-0440-5.

57. Hopkins MJ, Sharp R, Macfarlane GT. 2001. Age and disease related changes in intestinal bacterial populations assessed by cell culture, 16S rRNA abundance, and community cellular fatty acid profiles. Gut 48:198–205. https://doi.org/10.1136/gut.48.2.198.

58. Claesson MJ, Cusack S, O’Sullivan O, Greene-Diniz R, de Weerd H, Fitzgerald AP, Shanahan F, Twomey C, Hill C, Ross RP, O’Toole PW. 2011. Composition, variability, and temporal stability of the intestinal microbiota of the elderly. Proc Natl Acad Sci U S A 108(Suppl 1):4586–4591. https://doi.org/10.1073/pnas.1000971017.

59. Tamang JP, Watanabe K, Holzapfel WH. 2016. Review: diversity of microorganisms in global fermented foods and beverages. Front Microbiol 7:377. https://doi.org/10.3389/fmicb.2016.00377.

60. Buffa R, Floris G, Lodde M, Cotza M, Marini E. 2010. Nutritional status in a high-fat diet. Cell Host Microbe 26:265–272.e4.

61. Hoffmann C, Dollive S, Grunberg S, Chen J, Li H, Wu GD, Lewis JD, Telleria MT, Weiss M, Larsson K-H. 2013. Towards a unified paradigm for sequence-based identification of fungi. Mol Ecol 22:5271–5277.

62. Bisanz JE, Musser DC, Gozzi G, Turroni S, Ferrocino I, Lazzi C, Di Cagno R, Ercolini D. 2014. The same microbiota and a potentially discriminant metabolome in the saliva of omnivore, ovo-lacto-vegetarian and vegan individuals. PLoS One 9:e112373. https://doi.org/10.1371/journal.pone.0112373.

63. Rothschild D, Weissbrod O, Barkan E, Kurilshikov A, Korem T, Zeevi D, Lotan-Pompan M, Weinberger A, Halpern Z, Carmi S, Fu J, Wijmenga A, Zmora N, Pevsner-Fischer M, Israeli D, Kosower N, Malka G, Wolf BC, Avnit-Sagi T, Eberhardt U, Dueñas M, Grebenc T, Griffith GW, Hartmann M, Kirk PM, Kohout P, Larsson E, Lindahl BD, Lücking R, Martin MP, Matheny PB, Nguyen NH, Niskanen T, Oja J, Peay KG, Peintner U, Peterson M, Põldmaa K, Saag L, Saar I, Schüller A, Scott JA, Senés C, Smith ME, Suja A, Taylor DL, Telleria MT, Weiss M, Larsson K-H. 2013. Towards a unified paradigm for sequence-based identification of fungi. Mol Ecol 22:5271–5277. https://doi.org/10.1111/mec.12481.

64. DeSantis TZ, Hugenholtz P, Larsen N, Rojas M, Brodie EL, Keller K, Huber T, Dalevi D, Hu P, Andersen GL. 2006. Greengenes, a chimera-checked 16S rRNA gene database and workbench compatible with ARB. Appl Environ Microbiol 72:5069–5072. https://doi.org/10.1128/AEM.03006-05.

65. De Filippis F, Vannini L, La Storia A, Laghi L, Piombino P, Stellato G, Serrazanetti D, Gozzi G, Turroni S, Ferrocino I, Lazzi C, Di Cagno R, Gobbetti M, Ercole E. 2014. The same microbiota and a potentially discriminant metabolome in the saliva of omnivore, ovo-lacto-vegetarian and vegan individuals. PLoS One 9:e112373. https://doi.org/10.1371/journal.pone.0112373.

66. Fadrosh DW, Ma B, Gajer P, Sengamalay N, Ott S, Brotman RM, Ravel J. 2014. An improved dual-indexing approach for multiplexed 16S rRNA gene sequencing on the Illumina MiSeq platform. Microbiome 2:6. https://doi.org/10.1186/2049-2618-2-6.

67. Human Microbiome Project Consortium. 2012. Structure, function and diversity of the healthy human microbiome. Nature 486:207–214. https://doi.org/10.1038/nature11234.

68. Rognes T, Flouri T, Nichols B, Quince C, Mahe F. 2016. VSEARCH: a versatile open source tool for metagenomics. PeerJ 4:e2584. https://doi.org/10.7717/peerj.2584.

69. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EC, Fierer N, Pena AG, Goodrich JK, Gordon JJ, Huttley GA, Kelley ST, Knights D, Koenig JE, Ley RE, Lozupone CA, McDonald D, Muegge BD, Pirrung M, Reeder J, Sevinsky JR, Turnbaugh PJ, Walters WA, Widmann J, Yatsunenko T, Zaneveld J, Knight R. 2010. QIIME allows analysis of high-throughput community sequencing data. Nat Methods 7:335–336. https://doi.org/10.1038/nmeth.f.303.

70. Kõljalg U, Nilsson RH, Abarenkov K, Tedersoo L, Taylor AFS, Bahram M, Bates ST, Bruns TD, Bengtsson-Palme J, Callaghan TM, Douglas B, Drenkhan T, Eberhardt U, Dueñas M, Grebenc T, Griffith GW, Hartmann M, Kirk PM, Kohout P, Larsson E, Lindahl BD, Lücking R, Martin MP, Matheny PB, Nguyen NH, Niskanen T, Oja J, Peay KG, Peintner U, Peterson M, Põldmaa K, Saag L, Saar I, Schüller A, Scott JA, Senés C, Smith ME, Suja A, Taylor DL, Telleria MT, Weiss M, Larsson K-H. 2013. Towards a unified paradigm for sequence-based identification of fungi. Mol Ecol 22:5271–5277. https://doi.org/10.1111/mec.12481.

71. DeSantis TZ, Hugenholtz P, Larsen N, Rojas M, Brodie EL, Keller K, Huber T, Dalevi D, Hu P, Andersen GL. 2006. Greengenes, a chimera-checked 16S rRNA gene database and workbench compatible with ARB. Appl Environ Microbiol 72:5069–5072. https://doi.org/10.1128/AEM.03006-05.

72. Segata N, Izard J, Waldron L, Gevers D, Miropolsky L, Garrett WS, Knight R. 2011. Metagenomic microbial community profiling using parallelized sequence alignment. Genome Biol 12:R60. https://doi.org/10.1186/gb-2011-12-6-r60.

73. Parks DH, Tyson GW, Hugenholtz P, Beiko RG. 2014. STAMP: statistical analysis of taxonomic and functional profiles. Bioinformatics 30:3123–3124. https://doi.org/10.1093/bioinformatics/btu494.