

1 **Lifelong temporal dynamics of the gut microbiome associated with longevity in mice**

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17

18 **Abstract**

19 The effect of lifelong dynamics on host longevity of the gut microbiome is largely unknown. Herein,
20 we analyzed the longitudinal fecal samples of seven sibling mice across their lifespan from birth to
21 natural death, spanning over 1,000 days of age, and maintained them under controlled environmental
22 and dietary conditions. Our 16S-rRNA sequencing analysis revealed 38 common “life-core” bacterial
23 species/OTUs (operational taxonomic units) detected in $\geq 80\%$ of all samples collected across the
24 lifespan of individual mice. Despite the shared genetic background and dietary habits, the gut
25 microbiome structure significantly diversified with age and among individuals. We found a strong
26 positive correlation between longevity and the alpha diversity in middle age (500-700 days) and
27 negative correlation in old age (>800 days). Furthermore, host longevity was significantly associated
28 with the abundance of 17 bacterial species/OTUs, most of which were “life-core” species. Our data
29 suggest that temporal dynamics of the gut microbiome are strongly linked to host longevity.

30

31

32 **Introduction**

33 It has long been hypothesized that the gut microbiome affects the longevity and aging of hosts. In
34 studies using germ-free animals show longer lifespans than conventional animals¹, and the
35 administration of antibiotics in mice and insects extends the lifespan of mice^{2,3}. In addition to
36 sterilization, fecal microbiota transplantation from wild-type mice into progeroid mice has been
37 reported to enhance both the health and lifespan of mice⁴. Moreover, the administration of the probiotic
38 bacterium *Bifidobacterium animalis* or *Akkermansia muciniphila* has been shown to extend the
39 lifespan of mice⁴⁻⁶. Low-calorie diets are also associated with a prolonged life span in many biological
40 species⁷⁻⁹, including worms, flies, mice, and rats, and the underlying mechanism of this effect was
41 recently reported to be associated with the gut microbiome¹⁰. These studies directly suggest that the
42 gut microbiome is involved in the prolongation of lifespan; however, it remains unclear how structural
43 variations in the gut microbiome play a role in the prolongation of lifespan.

44 In the last decade, culture-independent metagenomics with advanced sequencing
45 technologies have allowed the investigation of differences in the structure of the gut microbiome at
46 different life stages, from infancy to the elderly¹¹⁻¹⁸. Most of these metagenome-based human cohort
47 studies have shown covariance of age with the gut microbiome by comparing gut microbiomes
48 between different age groups. However, these approaches do not accurately determine whether the
49 age-associated differences in gut microbiomes are primarily associated with chronological age or
50 confounded by other factors such as lifestyles because individuals in different age groups might have
51 been exposed to different environmental and lifestyle events. In addition, patient history, including
52 medications and a history of gut microbial colonization, may also affect the gut microbiome of humans
53 throughout life. In contrast, mice have a lifetime of two-three years, enabling the monitoring of time

54 series variations in the gut microbiome over a lifetime. Therefore, the present study aimed to clarify
55 the intimate transition in the gut microbiome with age using mice instead of humans. We analyzed
56 fecal samples longitudinally collected from siblings of specific pathogen-free (SPF) mice throughout
57 their whole life from birth to death, spanning more than 1,000 days, to elucidate the association of gut
58 microbiomes with the lifespan of hosts and life events, including pregnancy, delivery, and cohousing,
59 under the same rearing environments.

60

61 **Results**

62 **Mice, collection of fecal samples, and 16S rRNA gene analysis.** We purchased and bred the parent
63 mice (Mo and Fa, SPF C57BL/6J strain), which thereafter had nonuplet mice (one male, seven females,
64 and one unknown) at the animal facility. Of these siblings, seven mice (M1, 2, 3, 4, 5, 7, and 8) died
65 of natural causes with a mean lifetime of 922.7 ± 71.6 days ranging from 827 to 1,044 days, one mouse
66 (M6) died with a shorter lifespan than the others at 524 days probably due to the development of a
67 tumor under the same controlled environmental, and one mouse (M9) died immediately after birth.

68 Throughout the experiment, the eight siblings were cohoused with their parents for 51 days
69 after birth, and thereafter cohoused with different combinations of individuals in different cages, and
70 three (M3, M7, and M8) and Mo experienced pregnancy and delivery (Table S1). We performed
71 immediate autopsies of the dead mice and observed tumors in five mice (M2, 6, 7, 8, and Mo) but not
72 in others (M1, 3, 4, 5, and Fa).

73 We longitudinally collected a total of 1,815 fecal samples from 10 mice (eight siblings and
74 parents) with an average sampling interval of 4.3 days and obtained a total of 18.0 million high-quality
75 reads of the 16S rRNA gene V1-2 region from fecal DNA samples using the Illumina MiSeq platform
76 (Table S1, Methods). Clustering of 16S reads from samples of the seven siblings with natural death
77 with 97% identity individually generated operational taxonomic units (OTUs) ranging from 4,094 to
78 6,984 OTUs (5,646 OTUs on average), of which 11 to 17% (13% on average) were OTUs with $\geq 0.1\%$
79 abundance in at least one sample of each mouse. In addition, clustering of 16S reads from all samples
80 of the 10 mice (eight siblings and parents) generated a total of 21,768 OTUs, including 6,531 OTUs
81 with $\geq 0.1\%$ abundance in at least one sample, which was used for analysis with common OTU IDs in
82 this study (Table S1).

83

84 **Temporal variations in α -diversity across a lifetime.** Analysis of the cumulative number of OTUs
85 with $\geq 0.1\%$ abundance in at least one sample over a lifetime of 200 days after birth revealed that over
86 50 and 90% of the total OTUs appeared within 28 and 167 days in the samples of the seven siblings,
87 respectively (Extended Data Fig. 1). The number of OTUs rapidly increased to 200–300 in the first
88 20 days, although 9–15 OTUs were detected in 0-day samples of four mice (M2, 3, 6, and 8), and were
89 relatively constant after 100 days in most mice (Extended Data Fig. 2). The number of OTUs in the

90 bins of four mice (M3, 5, 7, and 8) was relatively constant across the lifetime after 100 days compared
91 to that of the others (M1, 2, and 4), which tended to have a lower number of OTUs in the latter half of
92 the lifespan of mice. We observed an increase in the number of OTUs in the last two bins just before
93 death in six out of seven mice, and four of them were significantly increased (Student's t-test, $P < 0.05$).

94

95 **Temporal variations in β -diversity across a lifetime.** The principal coordinate analysis (PCoA) of
96 weighted and unweighted UniFrac distances of all samples across the lifetime of the seven siblings
97 revealed that the most prominent difference in β -diversity across samples was that in samples before
98 and after 20 days (Figs. 1A and B), consistent with the change in the number of OTUs in the early
99 lifetime shown in Extended Data Fig. 2. The data also showed significant differences in gut
100 microbiome composition between the individuals after 20 days (permutational multivariate analysis
101 of variance [PERMANOVA] $P < 0.05$; Figs. 1C and D), suggesting a high inter-individual diversity
102 in gut microbiomes between the seven siblings even under the same rearing environments.

103

104 **Variations in the relative abundance of OTUs across a lifetime.** We found nine distinct dynamic
105 patterns (clusters) in the relative abundance of OTUs ($>0.1\%$ abundance) across lifetimes in the seven
106 siblings, based on the Silhouette index calculation (Figs. 2 and Extended Data Fig. 3). Clusters_1 and
107 4 contained OTUs with a relatively high appearance frequency and constantly high abundance
108 throughout most of their lifetime. Cluster_2 contained OTUs with two transient peaks in both early
109 and late lifetimes. Cluster_3 contained OTUs detected immediately after birth, and almost disappeared
110 in the latter lifetime. Cluster_5 contained OTUs with the highest abundance in the first half of the
111 latter half of life, whereas Cluster_6 contained OTUs with higher abundance in the latter half of the
112 first half of life. Cluster_7 contained OTUs with a transient peak in early age and a slight increase in
113 late age. Cluster_8 contained OTUs with two transient peaks in the early (~ 50 days) and middle
114 lifetimes (~ 500 days), with a relatively lower abundance in the latter half of the first half of the lifetime.
115 Cluster_9 contained OTUs that transiently increased only at the early lifetime, which was limited to
116 one mouse (M3). For the transient OTUs in the early lifetime, OTUs in Clusters_2, 4, 5, 7, and 9
117 increased after the earliest OTUs in Cluster_3 appeared (Extended Data Fig. 4).

118

119 **Life-core and transient OTUs/species across a lifetime.** Histograms of the number of OTUs ($\geq 0.1\%$
120 abundance in at least one sample) along the appearance frequency showed a bimodal distribution in
121 all samples of the seven siblings (Fig. 3A). We then defined OTUs detected in $>80\%$ and $<20\%$ of all
122 samples from each mouse as “life-core” and “transient” OTUs, respectively. We found 74 non-
123 redundant life-core OTUs (varying from 48 to 60 in seven siblings), of which 38 (minimum life-core
124 OTUs, 51%) were common in all mice and belonged to Clusters_1 and/or 4, and seven (9.5%) were
125 specific to individuals (Table S2). In contrast, 989 non-redundant transient OTUs (varying from 317

126 to 448 in seven siblings) were found in the samples and included only six (0.6%) common OTUs
127 belonging to Clusters_2, 3, and 5, and 394 (39.8%) were specific to individuals (Table S3). Taxonomic
128 assignment revealed that all life-core OTUs belonged to the four major phyla of the gut microbiome,
129 of which 52 (70%) and 31 (82%) were assigned to Bacteroidetes in the core and minimum life-core
130 OTUs, respectively. In contrast, the transient OTUs belonged to more than nine phyla, of which 816
131 (83%) were assigned to Firmicutes (Fig. 3B). In addition, life-core OTUs had a relatively high
132 abundance compared to transient OTUs (Extended Data Fig. 5).

133

134 **Correlation of alpha diversity with the lifespan of the host.** As the seven siblings had different
135 lifespans ranging from 832 to 1,049 days under the same rearing environments (Table S1), we explored
136 the correlation between gut microbiome structure and their lifespans. To this end, we developed a
137 method to identify and assess microbial variables correlated with lifespan, based on Spearman's
138 correlation coefficient (Extended Data Fig. 6, see Methods). The analysis revealed strong correlations
139 between lifespan and α -diversity indices (OTU#, Shannon's index) in samples at certain ages
140 (Extended Data Fig. 7): significant positive correlations with the indices at ages from 500 to 700 days,
141 and significant negative correlations with the indices at ages after 800 days.

142

143 **Correlation of OTU abundance with the lifespan of the host.** Using the same method, we also
144 found that 17 OTUs in samples at certain ages were significantly correlated in abundance with lifespan
145 (Fig. 4). Seven OTUs (OTU00012, OTU00013, OTU00028, OTU00035, OTU00037, OTU00055, and
146 OTU00105) were negatively correlated in abundance with late age (late period of the lifetime, after
147 800 days); two OTUs (OTU00006 and OTU00076) were positively correlated in abundance with
148 middle age (middle period of the lifetime, from 100 to 600 days) and negatively correlated with late
149 age; two OTUs (OTU00041 and OTU01676) were positively correlated in abundance with late age;
150 four OTUs (OTU00023, OTU00045, OTU00063, and OTU00069) were positively correlated in
151 abundance with middle age; and two OTUs (OTU00019 and OTU00033) were negatively correlated
152 in abundance with middle age.

153

154 **OTUs that correlated with life events.** We also investigated OTUs associated with several life events,
155 such as pregnancy, body weight, tumorigenesis, and cohousing, and compared them with the 17
156 lifespan-associated OTUs.

157 Three mice (M3, M7, and Mo) underwent pregnancy/delivery during the experiment.
158 Comparison of UniFrac distances between samples 20 days before pregnancy, during pregnancy, and
159 after delivery in these mice revealed significant differences in unweighted UniFrac distance between
160 any pair of three samples and in weighted UiFrac distance between before and during pregnancy
161 samples and between before and after delivery samples (PERMANOVA, $P < 0.05$; Extended Data

162 Fig.8A). The number of OTUs was not significantly different between the samples; however,
163 Shannon's index was significantly increased in samples during pregnancy and after delivery compared
164 to samples before pregnancy (Extended Data Fig. 8B). The analysis also revealed 59 OTUs with
165 significant changes in abundance between samples (Table S4). These included four lifespan-associated
166 OTUs (OTU00023, OTU00033, OTU00063, and OTU00076), which were significantly different
167 before and after delivery.

168 We found that OTUs significantly correlated with body weight in samples collected from
169 later stages of life from day 555 to just before death, and identified 27 OTUs significantly correlated
170 in abundance with body weight ($P < 0.05$, average $\rho > 0.3$; Extended Data Fig. 9). Of these, two
171 OTUs (OTU00006 and OTU00045) were lifespan-associated OTUs; OTU00006 was negatively
172 correlated whereas OTU00045 was positively correlated with body weight.

173 We observed tumors in five mice (M2, 6, 7, 8, and Mo) by immediate autopsy of the dead
174 mice. UniFrac distance analysis of samples 30 days before death revealed that the gut microbiome
175 significantly differed between samples with and without tumors (Extended Data Fig. 10A). Both the
176 number of OTUs and Shannon's index were higher in the tumor samples than in the non-tumor samples
177 (Extended Data Fig. 10B). We also identified 32 and 33 OTUs that were significantly enriched and
178 depleted, respectively, in tumor samples compared to non-tumor samples (Table S5). Of these, five
179 (OTU00006, OTU00012, OTU00013, OTU00035, and OTU00105) and two OTUs (OTU00041 and
180 OTU00055) were lifespan-associated OTUs.

181 During the experiments, 10 mice (eight siblings and parents) were cohoused with different
182 combinations of individuals in different frequencies and periods after initially cohousing all mice
183 together for 51 days after birth (Extended Data Fig. 11). We divided the samples after 20 days into
184 those with and without cohousing and compared their UniFrac distances. The results revealed that
185 both weighted and unweighted UniFrac distances were significantly lower in housed samples than in
186 non-housed samples (Extended Data Fig. 12).

187

188 **Overlap between the lifespan- and life events-associated OTUs.** As mentioned above, we identified
189 17 lifespan-associated OTUs, 59 pregnancy-associated OTUs, 27 body weight-associated OTUs, and
190 65 tumor-associated OTUs. These 168 OTUs were clustered to generate 127 OTUs specific to either
191 of the four categories, and 29 and 6 OTUs overlapped with 2 and 3 host factors, respectively (Table.
192 S6). Of the 17 lifespan-associated OTUs, 7 (41%) overlapped with tumors, four with pregnancy, and
193 two with body weight.

194

195 **Discussion**

196 This is a comprehensive time-course study that elucidates the development of the gut microbiome over
197 the lifespan of cognate SPF mice. The relationship between the gut microbiome and host aging has

198 been well recognized; however, it is mostly based on the comparative analysis of gut microbiomes
199 between individuals of various ages with different dietary habits, lifestyles, and genetic backgrounds.
200 In this study, we used sibling mice and longitudinally monitored changes in the gut microbiome of
201 individuals from birth to death for approximately three years. Even though the mice were bred in a
202 controlled environment, we eventually observed phenotypic diversity between individuals,
203 accompanied by the development of the gut microbiome. As C57BL/6J mice were designed as an
204 inbred line and have common genetic backgrounds, these diverse individual differences are likely
205 caused by stochastic environmental factors.

206 In this study, we found, for the first time, a tendency for increased diversity of the gut
207 microbiome before death by observing the variation in gut bacteria over the course of a lifetime in
208 mice (Extended Data Fig. 2). Age-related gut microbiome transitions have been observed in several
209 cohort studies; however, changes in gut microbial diversity in adults have been controversial¹⁹. Many
210 studies in developed countries have suggested that the microbiota is relatively stable throughout
211 adulthood, and that aging induces significant shifts in gut microbiome composition and function that
212 are associated with a decline in diversity^{15,20}. In contrast, another study targeting hunter-gatherers in
213 Hadza reported an increase in microbial diversity in the older adult population²¹. The increase in
214 diversity found in this study is consistent with the Hadza study. The older adults in the industrialized
215 cohort might have been affected by a medical treatment such as antibiotics, whereas the increased
216 diversity seen in mice in our study and the Hadza study could be due to the absence of the effect of
217 medications. In the older adult population, a drop in immune fitness has been reported, accompanied
218 by impaired hematopoiesis, higher susceptibility to infections, and reduced responses to vaccination²²⁻
219 ²⁸. Therefore, the increase in gut microbial diversity found in this study and the Hadza study suggests
220 a relationship with immune weakening due to aging.

221 In the present study, the analysis of variation patterns of OTU abundance revealed nine
222 distinct dynamic patterns of OTU abundance across the lifetime of seven siblings (Fig. 2). At
223 approximately day 500–600, we also observed a significant change in the dynamics of OTUs in
224 clusters 5 and 6. OTUs in cluster 5 decreased after day 500–600 whereas the dynamics of the ones in
225 cluster 6 increased in the latter half of life. These results suggest that some mode changes related to
226 the dynamics of the gut microbiota may have occurred around day 500–600; however, it is unclear
227 whether this is due to changes in host homeostasis or the ecological balance of the gut microbiota.
228 Additionally, our results suggest that some microbes persist for almost a lifetime in high abundance
229 (Fig. 3); however, Caporaso et al. have suggested, based on two individual human time series analyses,
230 that no core temporal microbiome exists at high abundance²⁹. This discrepancy may be because the
231 mice in our study were raised in a stable environment and were not affected by medication. Moreover,
232 the bimodal distribution indicated that OTUs can be largely divided into “life-core” and “transient”
233 OTUs. “Life-core” OTUs were largely dominated by Bacteroidetes whereas transient OTUs were

234 mostly composed of Firmicutes. These results are consistent with a study by Faith et al. reporting that
235 Bacteroidetes and Actinobacteria are significantly more stable over time in humans³⁴. Bacteroidetes
236 and Firmicutes are both dominant phyla in gut microbiomes; however, our results indicated differences
237 in their persistence throughout life. The Bacteroidetes group tends to persist for a long time whereas
238 most of the Firmicutes group, although high in abundance, may be unable to persist. A previous report
239 based on fluorescent in situ hybridization imaging technology showed that Bacteroidetes are mostly
240 observed near the intestinal wall, whereas Firmicutes are separated from the gut wall in the mouse
241 intestinal tract³⁰. Therefore, the settling of Bacteroidetes in the gut wall may allow them to persist in
242 the host. In addition to their location in the gut, Firmicutes have been proposed to be more effective
243 in extracting energy from food than Bacteroidetes, thereby promoting more efficient absorption of
244 calories and subsequent weight gain^{31,32}. MacFarlane et al. reported that Bacteroidetes mainly produce
245 acetate and propionate whereas Firmicutes produces more butyrate³³. These differences in the
246 metabolic capacity of Firmicutes and Bacteroidetes might also affect their persistence.

247 In the present study, we developed an original method to determine the dynamic relationship
248 between the gut microbiome and lifespan at various times in life: a positive correlation in middle age
249 (day 500–700) and a negative correlation in old age (after day 800) (Extended Data Fig.7). As many
250 diseases have been reported to lower the diversity of gut bacteria^{35–37}, it is often discussed that higher
251 diversity is related to a healthier life. However, in this study, the average α -diversity throughout life
252 was not strongly correlated with the lifespan (Extended Data Fig. 7). Although higher diversity was
253 correlated with a longer lifespan only in the middle period of life (day500~600), an opposite
254 correlation was found in old age (after day 800). Moreover, the correlation between abundance and
255 lifespan of OTUs before day 400 tended to be weaker than that in later life. We further found 17 OTUs
256 that were strongly associated with lifespan, most of which were life-core OTUs (Fig. 4), although this
257 correlation was altered or even reversed for many OTUs around day 500–600. Among the 17 lifespan-
258 associated OTUs, *Lactobacillus* was negatively correlated with the middle period of life and showed
259 a significant positive correlation later in life (after day 700). The abundance of *Lactobacillus* has been
260 reported to decrease with age. Furthermore, an increase in *Lactobacillus* has been reported to be
261 associated with lifespan elongation under limited calorie conditions in mice¹⁰. In our study, a
262 consistent tendency was observed in aged mice but not in young mice. We also found that OTU33,
263 which was closest to *B. vulgatus* (93.3%), showed a positive correlation with lifespan in later life but
264 a negative correlation in early life (before day 500). *B. vulgatus* was reported to have a significantly
265 higher abundance in village communities in Korea, where longevity is common³⁸. This correlation
266 between a higher abundance of *B. vulgatus* and longevity is consistent with our results in later life but
267 not in early life. Additionally, immunosuppressive drug rapamycin and the anti-diabetic drug
268 metformin, have recently been reported to extend the lifespan of mice related with gut microbiome^{3,39}.
269 In our study, the lifespan-associated OTUs included the genera *Bacteroides* and

270 Lactobacillus/Alistipes, which are known to be affected by rapamycin and metformin in mice,
271 respectively. The change in correlation pattern with age suggest the possibility that the previously
272 reported relationship between microbes and lifespan can also be a correlation found only at certain
273 times in lifetimes.

274 In addition, some lifespan-associated OTUs were associated with pregnancy, body weight,
275 and tumor development (Table S6). OTU00033 (*B. vulgatus* 93.02%) decreased after pregnancy and
276 was negatively associated with longevity in young mice whereas OTU00023 (*C. fastidious* 83.44%),
277 OTU00063 (*P. capillosus* 90.77%), and OTU00076 (*E. coprostanoligenes* 90.29%) increased after
278 pregnancy, and were positively associated with longevity in young mice. Therefore, diversity
279 increased with pregnancy, and higher diversity in youth was associated with longevity. Despite the
280 limited number of mice in this study, these results indirectly suggest that pregnancy at a young age
281 might be associated with longevity. Moreover, OTU00006 (*B. caecigallinarum* 90.16%) was
282 negatively correlated with both body weight and longevity in old mice whereas OTU00045 (*P.*
283 *timonensis* 87.9%) was positively correlated with both body weight and longevity (Table S6). These
284 results indirectly suggest that weight gain in old age might be associated with longevity. Five OTUs
285 (OTU00006, OTU00012, OTU00013, OTU00035, and OTU00105) enriched in tumor-developed
286 mice were negatively associated with longevity in aged mice (Table S6). Additionally, diversity
287 increased with tumor development, and higher diversity in aged mice was negatively associated with
288 longevity. These results are consistent with the obvious conclusion that the absence of tumors in old
289 age is associated with longevity.

290 In this study, for the first time, we characterized the life-core microbiome and showed that
291 the temporal dynamics of some life-core OTUs are significantly correlated with lifespan. This study
292 strongly suggests a close relationship between bacterial dynamics and lifespan. Further investigation
293 of host immune changes or bacterial functional interactions will help clarify the mechanism underlying
294 the temporal dynamics of diversity and bacterial abundance observed in this study.

295

296 **Methods**

297 **Fecal collection in mouse experiments.** Two SPF C57BL/6J mice were maintained and bred under
298 specific pathogen-free conditions at RIKEN Center for Integrative Medical Sciences Animal Facility
299 or conventional facility at Yokohama City University. (Kanagawa, Japan). Male and female offspring
300 were separated after weaning for 50-51 days. The mouse cages were replaced primarily to control
301 reproduction. The starting point of this study was the birth of the second generation of SPF mice. Each
302 mouse in each group was recognized using ear punches. Nine offspring were born. All mice were kept
303 in a room at $23 \pm 2^\circ\text{C}$, $50 \pm 10\%$ relative humidity, on a 12-h light/dark cycle. The animals were
304 housed in cages (Extended Data Fig. 11) with wood shavings and fed (CE-2, CLEA Japan, Inc.) and

305 watered ad libitum. Male mouse (M1) was fed Oriental CMF (Oriental Yeast, Tokyo) from day 462-
306 559 for breeding. One of the nice offspring died at 6 days of age and was therefore excluded from the
307 analysis (M9). Another mouse (M6) was excluded from the analysis of the association of the gut
308 microbiome with lifespan because it died during retention during sample collection due to debilitation.
309 Mouse feces were collected, snap-frozen, and stored at -80°C . All animal experiments were approved
310 by the Institutional Animal Care and Use Committees of RIKEN Yokohama Branch and Yokohama
311 City University.

312

313 **Autopsies of dead mice.** Dead mice were immediately stored in a freezer (-20°C). After thawing,
314 they were dissected by a surgeon researcher, who visually checked the left and right kidneys, spleen,
315 lungs, heart, cervical lymph nodes, ovaries, prostate, and liver for the presence of tumors.

316

317 **DNA extraction.** Bacterial genomic DNA was extracted from mouse feces using enzymatic lysis
318 method. Frozen fecal pellets were thawed and suspended in 1 mL of TE10 (10 mM Tris-HCl, 10 mM
319 EDTA) buffer containing RNaseA (final concentration of 100 $\mu\text{g}/\text{mL}$; Nippon Gene, Tokyo, Japan).
320 Lysozyme (Sigma-Aldrich; St. Louis, MO, USA) was added at a final concentration of 15 mg/mL,
321 and the suspension was incubated for 1 h at 37°C with gentle mixing. Purified achromopeptidase
322 (Wako Pure Chemical, Osaka, Japan) was added at a final concentration of 2000 units/mL, and the
323 suspension was further incubated for 30 min at 37°C . Sodium dodecyl sulfate (final concentration, 1
324 mg/mL) and proteinase K (final concentration, 1 mg/mL; MERCK Group Japan, Tokyo, Japan) were
325 added to the sample, and the mixture was incubated for 1 h at 55°C . DNA was extracted with
326 phenol/chloroform/isoamyl alcohol (25:24:1), precipitated with isopropanol and 3 M sodium acetate,
327 washed with 75% ethanol, and resuspended in 200 μL of TE buffer. DNA was purified with a 20%
328 PEG solution (PEG6000 in 2.5 M NaCl), pelleted by centrifugation, rinsed with 75% ethanol, and
329 dissolved in TE buffer.

330

331 **16S rRNA gene amplicon sequencing.** In this study, we analyzed a total of 51.4 million (51,377,124
332 reads) high-quality reads of the 16S rRNA gene V1-2 region from eight siblings and the parents
333 (Supplementary Table 1). The isolated DNA (40 ng) was used for PCR amplification of the V1–V2
334 hypervariable regions of the 16S rRNA gene using the universal primers 27Fmod (5'-
335 AATGATACGGCGACCACCGAGATCTACACxxxxxxxxACACTCTTCCCTACACGACGCTC
336 TTCCGATCTagrgttgatymtgctcag-3') and 338R (5'-
337 CAAGCAGAAGACGGCATAACGATxxxxxxxxGTGACTGGAGTTCAGACGTGTGCTCTTCC
338 GATCTgtgctcctccgtaggagt-3'), containing the Illumina Nextera adapter sequence and a unique 8-bp
339 index sequence for each sample. Thermal cycling was performed on a 9700 PCR system (Life
340 Technologies, Carlsbad, CA, USA) using Ex Taq polymerase (Takara Bio, Tokyo, Japan) with the

341 following cycling conditions: initial denaturation at 96 °C for 2 min; 25 cycles of denaturation at 96 °C
342 for 30 s, annealing at 55° C for 45 s, and extension at 72 °C for 1 min; and final extension at 72 °C.
343 All amplicons were purified using AMPure XP magnetic purification beads (Beckman Coulter, Brea,
344 CA, USA) and quantified using the Quant-iT PicoGreen dsDNA Assay kit (Life Technologies). An
345 equal amount of each PCR amplicon was mixed and subjected to sequencing with MiSeq (Illumina)
346 using the MiSeq Reagent Kit v3 (600 cycles) according to the manufacturer's instructions.

347

348 **Data analysis.** After demultiplexing the 16S sequence reads based on a sample-specific index, paired-
349 end reads were joined using the fastq-join program. Reads with average quality values <25 and inexact
350 matches to the universal primer sequences were filtered; 3,000 reads that passed the quality filter were
351 randomly selected from each sample and subjected to downstream analysis. The selected reads of all
352 the samples were first sorted by the frequency of redundant sequences and grouped into operational
353 taxonomic units (OTUs) using UCLUST (<https://www.drive5.com/>) with a sequence identity
354 threshold of 97%. The taxonomic assignment of each operational taxonomic unit was determined by
355 similarity searching against the Ribosomal Database Project and NCBI genome database using the
356 GLSEARCH program.

357

358 **Temporal dynamics clusters analysis.**

359 Temporal dynamics clusters were calculated on the time-course data of OTU abundances. Bray-Curtis
360 distance between the OTU abundances were calculated and hierarchically clustered by 'hclust'
361 function R software program (v4.1.1) with 'ward.D2' method. The optimal number of clusters were
362 calculated by 'find_k' function from R library 'dendextend'.

363

364 **Statistical analysis.** All statistical analyses were conducted with the R software program (v4.1.1).
365 Wilcoxon rank sum test was used for comparison of alpha diversities and the proportion of gut
366 microbiome at the phylum, genus, and OTU levels. PERMANOVA was used to compare the gut
367 microbiome structure among groups based on both weighted and unweighted UniFrac distance. P
368 values were corrected for multiple testing using the Benjamin-Hochberg method as appropriate.

369 **Correlation analysis of the gut microbiome with lifespan.** In the correlation analysis of the gut
370 microbiome structure and lifespan of the seven siblings, the variables (abundance of the bacteria/OTUs
371 and diversity indices of communities) that were significantly correlated with lifespan were determined
372 using the following five steps (Extended Data Fig. 6): (1) we calculated Spearman's correlation
373 coefficients between lifespan and the variables of samples at each time point lifetime (days); (2) we
374 calculated the moving average of the correlation coefficients of samples as it shifted every 50 days
375 throughout the lifetime (see the trend of the entire period by taking the average of 50 days in shifts);
376 (3) we developed a time series in which the sampling time is completely randomly switched, and the

377 moving average of the correlation was calculated in the same way; (4) we defined the range of possible
378 values of the moving average of the correlation coefficients of the random time series (not strongly
379 correlated with life expectancy at a particular time) calculated in step 3 in terms of the average value
380 and the 3σ range; and (5) finally, if the actual time series showed a moving average value that clearly
381 deviates from the range defined in step 4, we considered it significant.

382

383 **Data availability**

384 The 16S rRNA V1–V2 region sequences analyzed in the current study were deposited in
385 DDBJ/GenBank/EMBL under accession XXXXXXXX.

386

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392

393 **Author contributions**

394 Experimental design, L.T. and W.S.; sampling and experiment, L. T., E.W., and T.U.; 16S rRNA
395 sequencing, R.K., Y.O, Y.K., H.M.; Statistical analysis, L.T.; Writing—Original Draft, L.T.;
396 Writing—Review & Editing, W.S., M.U., and M.H.; Supervision, W.S. and M.H.; Funding
397 Acquisition, L.T., W.S., M.H.

398

399 **Competing interests**

400 The authors declare no competing interests.

401

402 **Figure legends**

403 **Fig. 1. Principle coordinate analysis (PCoA) based on UniFrac distance.**

404 **A**, PCoA of seven mice from day 0 to day 1044 of life-based on the weighted UniFrac distance. The
405 color corresponds to the day after birth. **B**, PCoA of seven mice from day 0 to day 1044 based on the
406 unweighted UniFrac distance. The color corresponds to the day after birth. **C**, PCoA of seven mice
407 from day 20 to day 1044 based on the weighted UniFrac distance. Color corresponds to an individual
408 mouse. **D**, PCoA of seven mice from day 20 to day 1044 based on the unweighted UniFrac distance.
409 The color corresponds to an individual mouse.

410

411 **Fig. 2. The nine clusters of temporal operational taxonomic unit (OTU) dynamics.**

412 The average relative abundance of the OTUs included in each of the nine temporal dynamic clusters
413 is shown. Temporal dynamics clusters were optimally determined based on average silhouette width
414 (See Methods).

415

416 **Fig. 3. Life-core and transient microbiome.**

417 The upper figures are histograms of the observed frequency of the operational taxonomic units (OTUs)
418 (maximum relative abundance $\geq 0.1\%$). The observed frequency was calculated by dividing the
419 number of times when an abundance of more than 0 reads was observed by the total number of
420 observed times. The bar graphs below indicate the phylum-level composition of the OTUs in each bin.

421

422 **Fig. 4. Seventeen operational taxonomic units (OTUs) are significantly correlated with lifespan**
423 **in mice.**

424 Spearman's correlation coefficients between the abundance of 17 lifespan-associated OTU and
425 lifespans are shown. The orange dots indicate Spearman's correlation coefficient at each time point.
426 The blue dots indicate the simple moving average of the coefficients over 50 time points. The yellow
427 mesh indicates the confidence interval (3σ).

428

429 **References**

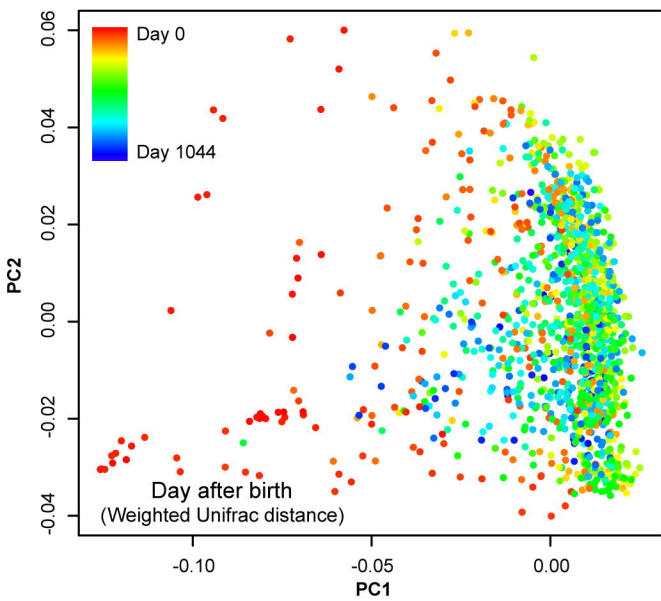
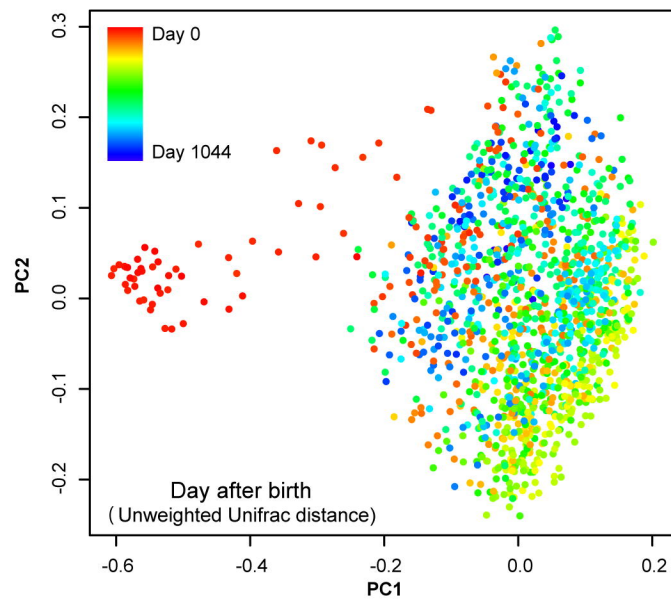
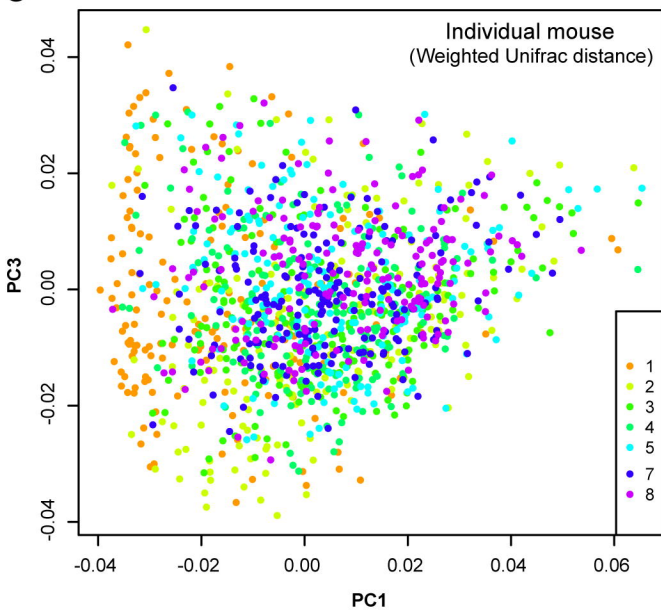
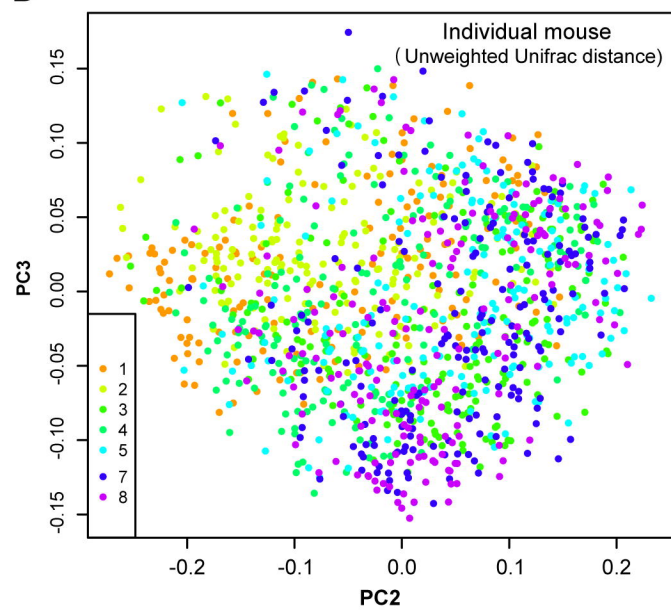
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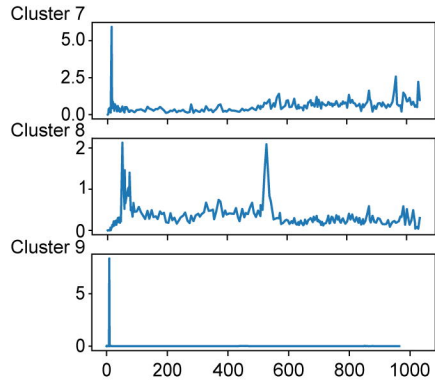
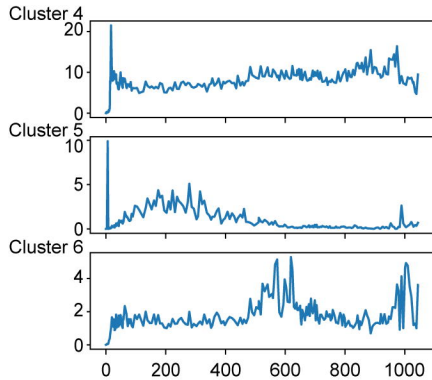
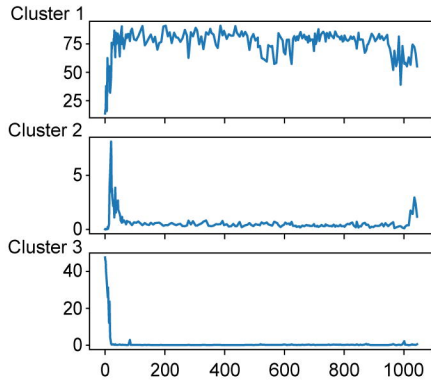
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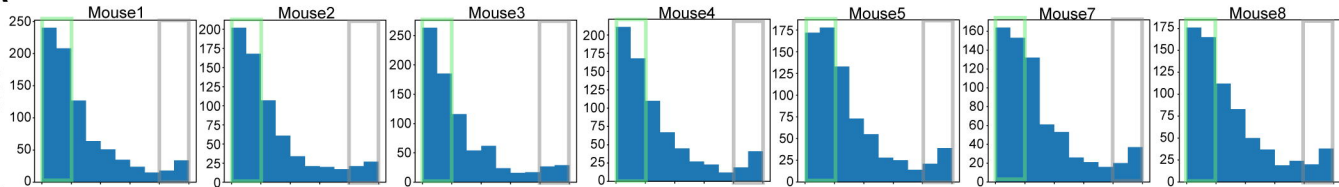
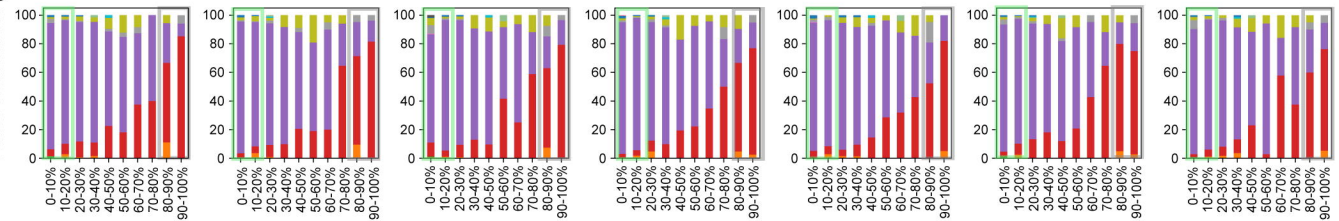
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A**B****C****D**

Relative abundance



Lifetime (days)

A**B**

Transient microbiome

Life-core microbiome

Acidobacteria TM7 Streptophyta Bacteroidetes Firmicutes Cyanobacteria Spirochaetes
 Proteobacteria Actinobacteria Deferribacteres Tenericutes Verrucomicrobia Fusobacteria UNDEFINED

Spearman's correlation coefficient

