

Title	Gut microbiota composition correlates with diet and health in the elderly	
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Publication date	2012-07-13	
Original Citation	Claesson, M. J., Jeffery, I. B., Conde, S., Power, S. E., O'Connor, E. M., Cusack, S., Harris, H. M. B., Coakley, M., Lakshminarayanan, B., O'Sullivan, O., Fitzgerald, G. F., Deane, J., O'Connor, M., Harnedy, N., O'Connor, K., O'Mahony, D., van Sinderen, D., Wallace, M., Brennan, L., Stanton, C., Marchesi, J. R., Fitzgerald, A. P., Shanahan, F., Hill, C., Ross, R. P. and O'Toole, P. W. (2012) 'Gut microbiota composition correlates with diet and health in the elderly', Nature, 488, pp. 178–184. doi: 10.1038/nature11319	
Type of publication	Article (peer-reviewed)	
Link to publisher's version	10.1038/nature11319	
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Download date	2025-04-18 07:45:58	
Item downloaded from	https://hdl.handle.net/10468/10653	



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Figure 1 | Principal Co-ordinate Analysis based on 18 different non-UniFrac beta-diversity
 distances.

- 7 Subjects are colour coded according to community setting; Community (green), Day Hospital
- 8 (yellow), Rehabilitation (orange), Long-stay (red), and Young healthy control subjects (purple).



Figure 2 | Pearson correlations and linear regression of Healthy Food Diversity and three microbiota diversity metrics for the 168 subjects where dietary information was available. Subjects were colour coded as in Figure 1.











Figure 3 | Microbiota diversity and healthy food diversity correlations

A) Comparison of three microbiota and one healthy food diversity indices across the four dietary groups. B) Procrustes plots of unweighted and C) weighted UniFrac PCoA analysis of microbiota combined with FFQ PCA. The subjects in all panels are colour coded according to diet groups; DG1 (green closed circles), DG2 (yellow open circles), DG3 (orange open circles), and DG4 (red closed circles).



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Figure 4 | Duration in long-stay care affects microbiota

A) Unweighted and B) weighted UniFrac PCoA of faecal microbiota from 191 subjects. Panels C and D show Procrustes analysis combining unweighted (C) and weighted (D) UniFrac PCoAs (non-circle end of lines) with Food Type PCA (circle-end of lines). The subjects are colour coded according to duration in long-stay care: N/A (community, day hospital and young healthy controls; green closed circles), less than six weeks (rehabilitation; yellow open circles), from six weeks to one year (long-stay; orange open circles), and longer than one year (long-stay; red closed circles).



38 39

40 Figure 5 | Representative ¹H NMR spectrum of a faecal water extract.

41 Peaks are labelled to indicate a range of metabolites present in faecal water. The faecal sample was

42 from from a subject in the community residence group. Peak identiities: 1, cholate; 2, caprylate; 3,

43 valerate; 4 butyrate; 5, isovalerate; 6, valine; 7, leucine; 8, isoleucine; 9, propionate; 10, threonine; 11,

- lactate; 12, isocaproate; 13, alanine; 14, lysine; 15, acetate; 16, glutamate; 17, succinate; 18,
 glutamine; 19, methylamine; 20, aspartate; 21, trimethylamine; 22, malonate; 23, taurine; 24, glycine.
- 45 grutannne, 19, meuryrannne, 20, aspartate; 21, trimetnyramine; 22, maionate; 23, taurine; 24, gr



46

Figure 6 | Co-inertia analysis (CIA) of relationships between metabolome, microbiota composition and
 residence location.

51 The upper left panel shows the CIA of the metabolomics PCA and microbiota PCA, with arrows

52 indicating where samples position in the metabolite dataset relative to the microbiota dataset. The

53 upper right panel shows NMR loadings data; 95 % confidence intervals were calculated for individual

54 loadings using jack-knife analysis¹. Loadings that are significantly different from zero are presented in

55 the plot as black dots with those that failed to show significance presented in grey. Relevant

56 metabolites are labelled with dashed lines connecting NMR regions that represent the same metabolite.

57 The lower panel displays the associated microbiota at genus level. Only genera present in at least 20%

58 of the samples were used in the analysis.





Figure 7 | Predicted gene counts and assembly statistics for the faecal metagenome of 27 selected subjects.
 The graphs show total gene counts and sequencing assembly N50 values for shotgun sequence data of
 the faecal metagenome for 27 subjects of indicated residence location.







Figure 8 | Frequency of genes relayed to butyrate, acetate and propionate production in the faecal metagenome of 27 subjects.

91 Comparison of A) gene counts and B) average sequencing coverage, for enzymes involved in butyrate, 92 acetate and propionate production. Gene count values were normalised for 4.79 x 109 of sequenced 93 bases per subject, and coverage values were normalised for the average coverage in each metagenome. 94 BCoAt: Butyryl-CoA transferase/Acetyl-CoA hydrolase; ACS: Acetate-formyltetrahydrofolate 95 synthetase/Formate-tetrahydrofolate Propionyl-CoA:succinate-CoA ligase; PCoAt: 96 transferase/Propionate CoA-transferase.



101 Figure 9 | Cytokine and C Reactive Protein levels vary across subject residence location.

102 Box plots are color coded by residence location according to the scheme in Figure 1. Kruskal-Wallis P-values 103 refer to tests performed across all four community locations, and Mann-Whitney test was performed for each 104 pair-wise comparison. 105



106 107 **NEW FIG**



A. PPCCA score plots derived from ¹H NMR spectra of faecal water with FIM or Barthel index as 109 covariate. Dot size reflects covariate value. The influence of the covariate was quantified by examining 110 regression parameter estimates and the associated 95% CI. Barthel index has a significant effect on 111 112 PC1 with an intercept of 1.62 (2.96, 0.28) (95% CI). FIM has a significant effect on PCI with an intercept of 1.58 (0.19, 2.96). Spectra from community (n=10) are represented by green circles, 113 rehabilitation (n=9) are represented by orange circles and long-stay residents (n=10) are represented by 114 red circles. B. The top loadings for PC1 for each model. Acetate (1.925, 1.915 ppm), butyrate (0.905, 115 116 0.895, 0.915, 2.165 ppm) and propionate (2.175, 2.195 ppm) increase with increasing FIM and Barthel 117 values. Error bars represent the 95% confidence intervals.



147

148 Figure 11 | Enterotype definition is highly dependent on choice of clustering method.

A) Enterotype clustering when based on Jensen-Shannon distances according to approach by Arumagam et al. 2010. B) Enterotype clustering when based on unweighted, and C) weighted UniFrac distances according to the approach by Wu et al. 2011. Abundances of *Prevotella* and *Bacteroidales* in the

two clusters were displayed as boxplots and in the unweighted UniFrac Procrustes plots to the right of figure. Clusters generated from the three methods were

151 two clusters were displayed as boxplots and in the unweighted Omraa Floerastes plots to the right of right. Clusters generated from the three methods were 152 validated using both the silhouette technique (left bar chart) and the Calinski-Harabasz Index (right bar chart). D) In replacement of clustering approach, the

153 *Prevotella/Bacteroides* ratio was ordered by size and subject colour-coded by residence location (left), and displayed as colour gradient in the Procrustes plots.



156 Figure 12 | Definition of bacterial Co-abundance groups (CAGs)

157 CAGs were defined by A) heat plot showing Kendall correlations between genera clustered by the 158 Spearman correlation coefficient and Ward linkage hierarchical clustering; B) Network plot highlighting correlation relationships between five CAGs and a central group, for the whole cohort 159 studied. Circle sizes indicate genus abundances. The thickness of the lines is proportional to 160 161 correlation strength.



Figure 13 | Genera that are significantly differentially abundant between Community (green)

and Long-stay (red) subjects.

- Circle sizes indicate significant differential over-abundance in either of the two residence locations.



- 171

Figure 14 | Relationship between the main eight microbiota groupings and health

indicators.

- Box-plots showing health measures and indices, and food indices, as a function of main
- microbiota groupings 1a - 4b from UniFrac analysis in Fig.4.



Figure 15 | Aggregate reads classified at phylum level in faecal microbiota of young control and elderly subjects.



Figure 16 | Major microbiota differences by community location.

The box-plots show significant genus-level differences between microbiota of young control subjects,

the complete older person group studied here (all elderly), and cohorts defined by residence location.



Figure 17 | Hierarchical clustering of denoised and un-denoised OTUs, for both sequencing

platforms, based on the Pearson correlation similarity metric and average linkage.

The top horizontal line for each heat map shows community location and the bottom line shows the different pyrosequencing runs with one colour each.

207 Supplementary methods

209 Subject recruitment and sample collection

210 This study was approved by the Cork Clinical Research Ethics Committee. Subjects older than 211 64 years were recruited and clinically investigated in two local hospitals, which serve a population 212 base of ~481,000 in the Cork city and county region. They were defined as (a) community-dwelling 213 (Community); (b) attending an out-patient day hospital (Out-patient); (c) in short-term rehabilitation 214 hospital care (Rehabilitation; under six-weeks stay) or (d) in long term institutionalized care (long-215 stay; more than six weeks). The mean age of the subjects was 78 (\pm 8) years, with a range of 64 to 102 216 years. The subjects were all of Irish ethnicity. None of the faecal samples from elderly subjects from our previous study² were analyzed in the current analysis, because we did not have food frequency 217 218 data for all that cohort. Exclusion criteria were a history of alcohol abuse, participation in an 219 investigational drug evaluation or antibiotic treatment within prior 30 days, or advanced organic 220 disease. Informed consent was obtained from all subjects or, in cases of cognitive impairment, by 221 next-of-kin in accordance with the local research ethics committee guidelines. Data collected included 222 anthropometric measurements, clinical history and status and medication history. Antibiotic use prior 223 to the one-month exclusion period was also recorded for each subject. Thirteen younger adult subjects 224 of age ranging 28-46 years, which had not been treated with antibiotics within 30 days, were also recruited by informed consent. 225

226

227 Clinical and nutritional data collection

Habitual dietary intake was assessed using a validated, semi-quantitative, food frequency questionnaire (FFQ) based upon the SLAN study³. Food properties were determined using the UK Food Standards Agency Nutrient databank⁴. The Mini Nutritional Assessment (MNA) was used as a screening and assessment tool to identify subjects at risk of malnutrition.

Non-fasted blood samples were collected and analysed at Cork University Hospital clinical laboratories. Cytokines were measured using validated, commercial multi-spot microplates (Meso Scale Diagnostics). Anthropometric measures included height, weight, calf and mid-arm circumference. Charlson Comorbidity Index, Mini mental State Exam, Geriatric Depression Test, Barthel Score and Functional Independence Measures were carried out on all participants. For longterm care, day-hospital and rehabilitation subjects, a research nurse reviewed the medical records for
information on disease and current medication usage.

239

240 Molecular methods and bioinformatics

241 DNA was extracted from fecal samples, and the V4 region of the 16S rRNA gene was amplified, sequenced and analyzed, as described previously⁵. Briefly, V4 amplicons were sequenced 242 243 on a 454 Genome Sequencer FLX Titanium platform (Roche Diagnostics Ltd, West Sussex, UK and 244 Beckman Coulter Genomics SA, Grenoble, France). Raw sequencing reads were quality trimmed 245 using the QIIME pipeline⁶ according to the following criteria: i) exact matches to primer sequences 246 and barcode tags; ii) no ambiguous bases (Ns); iii) read-lengths not shorter than 150 bp or longer than 247 350 bp; iv) the average quality score in a sliding window of 50 bp not to fall below 25. For large-scale assignments into the new Bergey bacterial taxonomy⁷ we used the RDP-classifier version 2.2 with 248 249 50% as confidence value threshold. This was based on what was found suitable for V4 amplicons from 250 the human gut environment⁵. RDP classifications were imported into a MySQL database for efficient 251 storage and advanced querying.

The amplicon reads were truncated at 210 bp prior to OTU picking at 97% similarity level, and filtered for chimeric sequences using ChimeraSlayer. Representative sequences (the most abundant) for each OTU were aligned using PyNAST⁸ prior to tree building using FastTree⁹. These phylogenies were combined with absence/presence or abundance information for each OTU to calculate unweighted or weighted UniFrac distances, respectively¹⁰. Principal Co-ordinate Analysis and Procrustes superimposition were then performed from the UniFrac distances and Food Frequency data. The amplicon sequences were deposited in MG-RAST under the Project ID 154.

Metagenomes were sequenced from libraries with 91 bp paired-end 91 Illumina reads and 350 bp insert size and assembled using MetaVelvet¹¹. Samples EM039 and EM173 were sequenced from libraries of 101 bp paired-end Illumina reads with a 500 bp insert size, and subsequently assembled using MIRA¹² in hybrid with 551,726 and 665,164 454 Titanium reads, respectively. Protein sequences from enzymes were screened against the assembled metagenomes using TBLASTN with an amino acid identity cut-off of 30% and an alignment length cut-off of 200 bp. We screened the metagenome data for enzymes associated with production of butyrate (butyryl-CoA transferase/acetyl-CoA hydrolase), acetate (acetate-formyltetrahydrofolate synthetase/formate-tetrahydrofolate ligase), and propionate (propionyl-CoA:succinate-CoA transferase/propionate CoA-transferase). Genes were predicted using MetaGene¹³.

269

270 NMR analysis of the faecal water metabolome

271 Faecal water samples were prepared by the addition of 60 μ l D₂O and 10 μ l tri-methylsilyl-272 2,2,3,3-tetradeuteriopropionate to 540 µl faecal water. Spectra of samples were acquired by using a 273 Carr-Purcell-Meiboom-Gill (CPMG) pulse sequence with 32 K data points and 256 scans. Spectra 274 were referenced to TSP at 0.0 ppm, phase and baseline corrected with a line broadening of 0.3 Hz 275 using the processor on Chenomx NMR suite 7 (Chenomx Inc, Edmonton, Alberta, T5K 2J1). The 276 spectra were integrated at full resolution for data analysis (PCA, PLS-DA, CIA) with the water region (4-6 ppm) excluded and the data was normalized to the sum of the spectral integral. For PPCCA data 277 278 analysis, the spectra were integrated into spectral regions (0.01ppm).

279

280 Statistical methods and metabolome data analysis

Statistical analysis was carried out using R (version 2.13.2) or Stata (version 11) software 281 282 packages. Kruskal-Wallis and Mann-Whitney tests were used to find significant differences in 283 microbial taxa, clinical and biochemical measures, alpha diversity, and Healthy Food Diversity (HFD). 284 We used least square linear regression for comparing alpha diversity and HFD. Median regression¹⁴ 285 was used to compare clinical measures and microbiota, while adjusting for age, gender, medications, 286 and when appropriate residence location. For median regression, the median was modelled as a linear 287 function of independent variables. Model parameters are estimated such that they minimised the sum 288 of the absolute differences between observed and predicted values. P-values were generated using the wild bootstrap method¹⁵ to estimate variance. 289

290 A linear quantile (median) regression for two variables – a response variable (y) and a 291 predictor variable (x) – is the following: median (y) = $\beta 0 + \beta 1x$ where $\beta 0$ is the intercept (value when y = 0 and $\beta 1$ is the slope (change in median of y for a unit change in x). Together, these parameters describe the association between y and x, where x is a predictor of y. In the case of multiple predictor variables, each one is added to the regression equation and so the equation becomes median (y) = $\beta 0$ + $\beta 1x1 + \beta 2x2$ and now the slope $\beta 1$ is interpreted as the median change in x1 after adjusting for x2. This can be likened to a laboratory experiment where the specific effect of one variable on another is isolated by holding all other relevant variables constant.

Following statistical analysis of the taxonomic classifications, we estimated q-values to control for multiple testing using the qvalue and nFDR functions in the R package¹⁶. At the genus level we estimated the proportion of true null hypotheses with the qvalue function unless the estimated pi0 was less or equal to zero. In all other instances, we used the Benjamini & Hochberg method¹⁷.

302 Statistical analysis of the NMR data was performed using diverse software packages: PCA and 303 PLS-Da analysis was performed in SIMCA-P+ (Umetrics, Umeå, Sweden); Permutation testing was 304 performed in R and PPCCA was performed in R using the MetabolAnalyze package. The NMR data 305 was Pareto-scaled prior to data analysis. Assignment of the spectral peaks was performed using in-306 house libraries, statistical correlation analysis and 2D NMR spectra (TOCSY and COSY).

307 Supplementary Tables

308 Table 1. Taxa that are found in 50% of the subjects, significantly different across the four 309 community locations, according to a Kruskal-Wallis test, and that are more abundant in 310 Community subjects relative to Long-stay subjects.

311 C: Community subjects; DH: Day Hospital; R: Rehabilitation; LS: Long-stay. Community location

312 columns show medians of percentages of RDP-classified reads over the total number of reads for each

313 subject. The FDR value represents a Benjamini-Hochberg corrected P-value for the smaller data sets

314 Clostridium cluster and Family, and a Q-value for the larger data sets Genus and Species.

	С	DH	R	LS	FDR
Family					
Lachnospiraceae					
(Firmicutes/Clostridia/Clostridiales)	10.50	8.94	11.19	5.95	7.1E-05
Pasteurellaceae					
(Proteobacteria/Gammaproteobacteria/Pasteurellales)	0.03	0.05	0.01	0.00	0.0025
Alcaligenaceae					
(Proteobacteria/Betaproteobacteria/Burkholderiales)	0.28	0.53	0.33	0.09	0.0034
Genus					
Coprococcus					
(Firmicutes/Clostridia/Clostridiales/Lachnospiraceae)	1.52	0.67	1.23	0.40	8.40E-08
Roseburia					
(Firmicutes/Clostridia/Clostridiales/Lachnospiraceae)	1.79	2.51	2.26	0.28	8.40E-07
Butyricicoccus					
(Firmicutes/Clostridia/Clostridiales/Ruminococcaceae)	0.11	0.08	0.07	0.05	0.00002
Catenibacterium					
(Firmicutes/Erysipelotrichi/Erysipelotrichales/Erysipelotrichaceae)	0.06	0.02	0.00	0.00	0.00002
Oribacterium					
(Firmicutes/Clostridia/Clostridiales/Lachnospiraceae)	0.01	0.01	0.01	0.00	0.00006
Anaerosporobacter					
(Firmicutes/Clostridia/Clostridiales/Lachnospiraceae)	0.44	0.19	0.35	0.12	0.00018
Actinobacillus					
(Proteobacteria/Gammaproteobacteria/Pasteurellales/Pasteurellaceae)	0.03	0.05	0.01	0.00	0.00018
Lachnobacterium					
(Firmicutes/Clostridia/Clostridiales/Lachnospiraceae)	0.01	0.00	0.01	0.00	0.00027
Hydrogenoanaerobacterium					
(Firmicutes/Clostridia/Clostridiales/Ruminococcaceae)	0.47	0.07	0.43	0.12	0.00057
Sutterella					
(Proteobacteria/Betaproteobacteria/Burkholderiales/Alcaligenaceae)	0.09	0.11	0.06	0.00	0.00085
Parasutterella					
(Proteobacteria/Betaproteobacteria/Burkholderiales/Alcaligenaceae)	0.08	0.06	0.05	0.01	0.0011
Parasporobacterium					
(Firmicutes/Clostridia/Clostridiales/Lachnospiraceae)	0.08	0.04	0.06	0.02	0.0019
Barnesiella					
(Bacteroidetes/Bacteroidia/Bacteroidales/Porphyromonadaceae)	0.48	0.44	0.67	0.11	0.0048

315

317Table 2. Taxa identified as per criteria in SupplementaryTable 1 but showing taxa that are more318abundant in Long-stay subjects relative to Community-dwelling subjects.

	С	DH	R	LS	FDR
Family					
Desulfovibrionaceae					
(Proteobacteria/Deltaproteobacteria/Desulfovibrionales)	0.01	0.01	0.02	0.09	5.8E-05
Lactobacillaceae					
(Firmicutes/Bacilli/Lactobacillales)	0.00	0.01	0.15	0.02	7.1E-05
Eubacteriaceae					
(Firmicutes/Clostridia/Clostridiales)	0.18	0.87	0.19	0.89	7.1E-05
Porphyromonadaceae					
(Bacteroidetes/Bacteroidia/Bacteroidales)	4.41	7.20	5.57	8.53	0.00047
Enterobacteriaceae					
(Proteobacteria/Gammaproteobacteria/Enterobacteriales)	0.01	0.02	0.03	0.12	0.00059
Coriobacteriaceae					
(Actinobacteria/Actinobacteria/Coriobacteriales)	0.01	0.01	0.03	0.02	0.00087
Incertae Sedis XI					
(Firmicutes/Clostridia/Clostridiales)	0.00	0.00	0.00	0.01	0.0054
Streptococcaceae					
(Firmicutes/Bacilli/Lactobacillales)	0.05	0.09	0.11	0.10	0.042
~					
Genus					
Coprobacillus					
(Firmicutes/Erysipelotrichi/Erysipelotrichales/Erysipelotrichaceae)	0.01	0.02	0.05	0.15	5.7E-09
Lactonifactor	0.07	0.10	0.10		
(Firmicutes/Clostridia/Clostridiales/Ruminococcaceae)	0.06	0.10	0.18	0.21	3.7E-07
Anaerotruncus	0.00	0.00	0.10		1 25 05
(Firmicutes/Clostridia/Clostridiales/Ruminococcaceae)	0.09	0.08	0.12	0.26	4.3E-07
Acetanaerobacterium	0.01	0.02	0.00	0.04	5 (1) 07
(Firmicutes/Clostridia/Clostridiales/Ruminococcaceae)	0.01	0.02	0.02	0.04	5.6E-07
Lactobacillus	0.00	0.01	0.15	0.00	
(Firmicutes/Bacilli/Lactobacillales/Lactobacillaceae)	0.00	0.01	0.15	0.02	3.9E-06
Anaerofilum	0.00	0.00	0.00	0.01	7 0 1 0 <i>6</i>
(Firmicutes/Clostridia/Clostridiales/Ruminococcaceae)	0.00	0.00	0.00	0.01	7.8E-06
Eubacterium	0.16	0.97	0.10	0.90	1.05.05
(Firmicutes/Clostindia/Clostindiales/Eubacternaceae)	0.16	0.87	0.19	0.89	1.0E-05
Papillibacter (Firmi outor/Clostri dio/Clostri diolos/Dumino occosoco)	0.11	0.11	0.10	0.29	0.00002
(Filinicutes/Clostificia/Clostificiales/Rulliniccoccaceae)	0.11	0.11	0.10	0.38	0.00002
(Pastaroidatas/Pastaroidia/Pastaroidalas/Pornhyromonadasasa)	2.56	4 19	4 20	5.96	0.00000
(Bacteroideles/Bacteroidia/Bacteroidales/Porphyromonadaceae)	2.30	4.18	4.20	3.80	0.00009
(Firmicutes/Clostridia/Clostridiales/Puminococcaceae)	0.04	0.01	0.01	0.11	0.00018
Ethanoliganons	0.04	0.01	0.01	0.11	0.00018
(Firmicutes/Clostridia/Clostridiales/Puminococcaceae)	0.03	0.02	0.02	0.06	0.00038
Doreg	0.03	0.02	0.02	0.00	0.00038
(Firmicutes/Clostridia/Clostridiales/Lachnospiraceae)	0.30	0.44	0.47	0.66	0.00055
(Thinkutes/Elosthula/Elosthula/Es/Eachiosphaceae)	0.50	0.44	0.47	0.00	0.00033
(Protechacteria/Gammaprotechacteria/Enterchacteriales/Enterchacteriaceae)	0.00	0.02	0.02	0.06	0.0012
Subdoligranulum	0.00	0.02	0.02	0.00	0.0012
(Firmicutes/Clostridia/Clostridiales/Ruminococcaceae)	0.34	0.34	0.42	0.70	0.0026
Desulfovibrio	0.54	0.54	0.42	0.70	0.0020
(Proteobacteria/Deltaproteobacteria/Desulfovibrionales/Desulfovibrionaceae)	0.00	0.00	0.00	0.04	0.0032
Oxobacter	0.00	0.00	0.00	0.04	0.0052
(Firmicutes/Clostridia/Clostridiales/Clostridiaceae)	0.02	0.02	0.01	0.04	0.0052
Strentococcus	0.02	0.02	0.01	0.04	0.0032
(Firmicutes/Bacilli/Lactobacillales/Streptococcaceae)	0.04	0.08	0.11	0.09	0.0054

322 Table 3. Median values of food consumption for each of the four diet groups (DGs).

Food Names	Food Groups	DG1	DG2	DG3	DG4
White bread and rolls (including ciabatta & pannini)	cereals and potatoes	0.01	0.143	0.4285	0
Brown bread and rolls	cereals and potatoes	0.143	0.286	0.0215	0
Wholemeal bread and rolls	cereals and potatoes	0.714	0.033	0	0
Wheat-free; Rye bread; spelt bread (specify)	cereals and potatoes	0	0	0	0
Cream crackers, cheese biscuits	cereals and potatoes	0	0.01	0	0
Crisp bread	cereals and potatoes	0	0	0	0
Pancakes, muffins, oatcakes	cereals and potatoes	0	0	0	0
Scone (white)	cereals and potatoes	0.01	0.01	0.0215	0.005
Scone (brown)	cereals and potatoes	0	0	0	0
Non-ready to eat - Porridge	cereals and potatoes	1	0.286	1	1
High fibre (Bran/flakes, wheat biscuits, shredded wheat)	cereals and potatoes	0	0	0	0
Corn flakes, popped rice	cereals and potatoes	0	0	0	0
Muesli	cereals and potatoes	0	0	0	0
Sugar-coated cereals	cereals and potatoes	0	0	0	0
Boiled, instant or jacket potatoes	cereals and potatoes	1	0.714	0	0
Mashed potatoes	cereals and potatoes	0.033	0.286	1	1
Chips / Roast potatoes	cereals and potatoes	0.033	0.033	0.033	0.005
Potato Salad	cereals and potatoes	0.01	0.01	0	0
White Rice	cereals and potatoes	0.01	0	0	0
Brown Rice	cereals and potatoes	0.01	0	0	0
White/yellow or green pastas (e.g. spaghetti, noodles)	cereals and potatoes	0.01	0	0	0
Wholemeal pasta	cereals and potatoes	0	0	0	0
Cream (tablespoon)	dairy	0	0.01	0	0.2145
Creme fraiche/soured cream	dairy	0	0	0	0
Low-fat yoghurt	dairy	0	0	0	0
Full-fat yoghurt or Greek style yoghurt (125g carton)	dairy	0	0.01	0	0.286
Dairy desserts (125g carton)	dairy	0	0	0	0.005
Cheddar cheese (medium serving)	dairy	0.143	0.143	0	0
Low-fat cheddar cheese (medium serving)	dairy	0	0	0	0
Cottage cheese	dairy	0	0	0	0
Eggs as boiled, fried, scrambled, poached (one)	dairy	0.286	0.286	0.143	0
Butter	dairy	0	0	0	0.0715
Lite Butter	dairy	0	0	0	0
Low-fat margarine	dairy	0	0	0	0
Cholesterol Lowering margarine	dairy	0	0	0	0
Cream & Veg. Oil spread	dairy	0	0	0	0
Olive oil spread	dairy	0	0	0	0
Meat or cream soups: homemade / fresh (1 bowl)	dairy	0	0.01	0	0
Meat or cream soups: tinned / packet (1 bowl)	dairy	0	0	0	0
Whole milk (cup) - cow, goat, soya, rice milk -	dairy	0	0	0.143	1

specify					
Semi-skimmed milk (cup)	dairy	0	0	0	0
Probiotic Yoghurts / cheese / milk	dairy	0.286	0	0	0
Fish fried (batter/breadcrumbs), baked, grilled, fingers/cakes	fish	0	0.01	0.088	0.143
White fish, fresh or frozen (e.g. cod, haddock, plaice, sole)	fish	0.143	0.143	0	0
Oily fish, fresh or canned (e.g. mackerel, kippers, tuna, salmon, sardines, herring)	fish	0.143	0.033	0	0
Shellfish (e.g. crab, prawns, mussels)	fish	0	0	0	0
Apples	fruit	0.286	0.143	0	0.2145
Pears	fruit	0.143	0.033	0	0
Oranges, satsumas, mandarins	fruit	0.286	0.033	0	0
Grapefruit	fruit	0	0	0	0
Bananas	fruit	0.286	0.286	0.143	0.2145
Grapes	fruit	0.143	0.01	0	0
Melon	fruit	0.01	0	0	0
Peaches, plums	fruit	0.01	0	0	0
Apricots	fruit	0	0	0	0
Strawberries, raspberries, kiwi fruit	fruit	0.143	0.01	0	0
Blueberries	fruit	0	0	0	0
Tinned fruit (specify)	fruit	0	0.01	0	0
Dried fruit e.g. raisins	fruit	0.01	0	0	0
Frozen fruit (specify)	fruit	0	0	0	0
Other fruit	fruit	0	0	0	0
Pure fruit drinks e.g. orange juice (small glass)	fruit	0.033	0.033	0.286	1
Fruit squash (small glass)	fruit	0	0	0	0
Beef (roast / steak)	meat	0.033	0.143	0.143	0.286
Beef: stew	meat	0.033	0.143	0.143	0.143
Beef burger (1 burger)	meat	0	0	0	0
Pork (roast / chops / escalopes)	meat	0.01	0.033	0.143	0.143
Lamb (roast / chops / stew)	meat	0.033	0.143	0.143	0.143
Chicken or other poultry e.g. turkey: roast	meat	0.143	0.143	0.286	0.286
Breaded chicken, chicken nuggets, chicken burger	meat	0	0	0	0
Bacon / Ham	meat	0.033	0.143	0.143	0.143
Processed meat (Corned beef, Luncheon meats, sausages)	meat	0	0.033	0.143	0.143
Savoury pies (e.g. meat, pork, steak & kidney, sausage roll)	meat	0	0	0	0
Liver, heart, kidney, paté	meat	0	0	0	0
Chocolate coated sweet biscuits e.g. digestive (one)	sweets, cakes and alcohol	0.143	0.033	0	0
Plain biscuit e.g. digestives, rich tea (one)	sweets, cakes and alcohol	0.033	0.286	0.286	0.2145
Cakes e.g. fruit, sponge	sweets, cakes and alcohol	0.033	0.033	0.033	0
Buns, pastries e.g. croissants, doughnuts	sweets, cakes and alcohol	0	0	0	0
Fruit pies, tarts, crumbles	sweets, cakes and alcohol	0.01	0.033	0.005	0
Sponge puddings	sweets, cakes and alcohol	0	0	0	0.005
Milk puddings e.g. rice, custard, trifle	sweets, cakes and alcohol	0.01	0.033	0.286	0.286

Ice cream, choc ices, frozen desserts	sweets, cakes and alcohol	0.01	0.143	0.143	0.286
Chocolates, singles or squares	sweets, cakes and alcohol	0.033	0.033	0.005	0
Sweets, toffees, mints	sweets, cakes and alcohol	0.01	0.033	0	0
Sugar added to tea coffee, cereals (teaspoons)	sweets, cakes and alcohol	0	0	0	3
Sugar substitute e.g. canderel (teaspoon)	sweets, cakes and alcohol	0	0	0	0
Crisps or other packet snacks	sweets, cakes and alcohol	0	0	0	0
Jam, marmalade, honey, syrup (teaspoon)	sweets, cakes and alcohol	1	1	0.5	0
Wine (glass)	sweets, cakes and alcohol	0.143	0	0	0
Beer, Larger or Cider (half pint)	sweets, cakes and alcohol	0	0	0	0
Low alcohol / alcohol free beer / larger (half pint)	sweets, cakes and alcohol	0	0	0	0
Port, Sherry, Vermouth, liqueurs (glass)	sweets, cakes and alcohol	0	0	0	0
Spirits e.g. Gin, Whiskey (Single measure)	sweets, cakes and alcohol	0	0	0	0
Low calorie or diet soft fizzy drink (glass)	sweets, cakes and alcohol	0	0	0	0.005
Fizzy Soft drinks e.g.soda pop	sweets, cakes and alcohol	0	0	0	0.2145
Carrots	vegetables	0.286	0.286	0.286	0.714
Spinach	vegetables	0.01	0	0	0
Broccoli, spring greens, kale	vegetables	0.286	0.143	0	0
Brussels sprouts	vegetables	0.01	0.01	0.033	0.143
Cabbage	vegetables	0.033	0.143	0.143	0.143
Peas	vegetables	0.033	0.143	0.143	0.286
Green beans, broad beans, runner beans	vegetables	0.01	0	0	0.143
Courgettes	vegetables	0.01	0	0	0
Cauliflower	vegetables	0.033	0.143	0.143	0.143
Parsnips, turnips	vegetables	0.143	0.143	0.143	0.143
Leeks	vegetables	0.033	0	0	0
Onions	vegetables	0.286	0.143	0	0
Garlic	vegetables	0.143	0	0	0
Mushrooms	vegetables	0.143	0.033	0	0
Sweet peppers	vegetables	0.143	0	0	0
Green salad, lettuce	vegetables	0.286	0.143	0	0
Cucumber, celery	vegetables	0.143	0.033	0	0
Tomatoes	vegetables	0.286	0.286	0.088	0
Sweet-corn	vegetables	0	0	0	0
Beetroot	vegetables	0.033	0.01	0	0
Coleslaw	vegetables	0.01	0.01	0	0
Baked beans	vegetables	0.033	0.143	0	0
Dried lentils, beans, peas	vegetables	0	0	0	0
Tofu, soya meat, TVP, vege-burger	vegetables	0	0	0	0
Lasagne (meat based)	vitamins / minerals / tea etc.	0.01	0	0	0
Lasagne (vegetarian)	vitamins / minerals / tea etc.	0	0	0	0
Pizza	vitamins / minerals / tea etc.	0.01	0	0	0
Quiche (medium serving)	vitamins / minerals / tea etc.	0	0	0	0
Light salad cream or light mayonnaise (tablespoon)	vitamins / minerals / tea etc.	0.01	0	0	0
Salad cream, mayonnaise (tablespoon)	vitamins / minerals / tea etc.	0	0	0	0

French dressing (tablespoon)	vitamins / minerals / tea etc.	0.01	0	0	0
Other salad dressing (tablespoon)	vitamins / minerals / tea etc.	0	0	0	0
Sunflower margarine e.g. Flora	vitamins / minerals / tea etc.	0	0	0	0
Peanuts or other nuts	vitamins / minerals / tea etc.	0.01	0	0	0
Vegetable soups: homemade/fresh (1 bowl)	vitamins / minerals / tea etc.	0.033	0.143	0.286	0.286
Vegetable soups: tinned/packet (1 bowl)	vitamins / minerals / tea etc.	0	0	0	0
Sauces e.g. white, cheese, gravy (tablespoon)	vitamins / minerals / tea etc.	0.033	0.286	1	1
Tomato based sauces e.g. pasta sauces	vitamins / minerals / tea etc.	0.01	0	0	0
Curry-type sauces	vitamins / minerals / tea etc.	0.01	0	0	0
Pickles, chutney (tablespoon)	vitamins / minerals / tea etc.	0.01	0	0	0
Marmite, Bovril (tablespoon)	vitamins / minerals / tea etc.	0	0	0	0
Peanut butter (teaspoon)	vitamins / minerals / tea etc.	0	0	0	0
Tea (cup)	vitamins / minerals / tea etc.	2	4	4	4
Herbal tea (cup)	vitamins / minerals / tea etc.	0	0	0	0
Coffee instant (cup)	vitamins / minerals / tea etc.	0.143	0.033	0	0
Coffee ground (cup)	vitamins / minerals / tea etc.	0	0	0	0
Coffee, decaffeinated (cup)	vitamins / minerals / tea etc.	0	0	0	0
Coffee whitener e.g. Coffee-mate (teaspoon)	vitamins / minerals / tea etc.	0	0	0	0
Cocoa, Hot Chocolate (cup)	vitamins / minerals / tea etc.	0	0	0	0
Horlicks, Ovaltine (cup)	vitamins / minerals / tea etc.	0	0	0	0
Vitamin supplements (details)	vitamins / minerals / tea etc.	0	0	0	0
Mineral supplements (details)	vitamins / minerals / tea etc.	0	0	0	0
Other supplements	vitamins / minerals / tea etc.	1	0	0	0
Ready meal (specify)	vitamins / minerals / tea etc.	0	0	0	0
Takeaway (specify)	vitamins / minerals / tea etc.	0	0	0	0

329 Table 4 Assembly statistics for shotgun sequencing metagenomics.

Of the 27 assemblies, 25 were based on libraries with 91 bp paired-end 91 Illumina reads and 350 bp insert size. Samples EM039 and EM173 were assembled using 101 bp paired-end Illumina reads, from libraries with a 500 bp insert size, and in hybrid assembly with 551,726 and 665,164 454 Titanium reads, respectively. Only contigs larger than 500 bp were used for further analysis.

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- 335

Sample	Number	Total contig	Largest	N50	Total number of
	of contigs	length (bp)	contig (bp)	(bp)	bases $\frac{337}{228}$
EM039	57,873	102,436,267	495,772	2,873	3,317,557,3380
EM148	8,121	56,447,615	520,052	30,601	4,795,110,722
EM172	55,298	83,738,852	125,936	2,050	4,472,602,022
EM173	48,821	108,101,567	146,798	4,013	3,130,777,7 9 9 1
EM175	7,706	56,078,081	595,761	30,216	4,772,905,2 02 2
EM176	8,004	64,792,202	656,046	36,029	4,780,810,982
EM177	46,980	121,070,988	217,703	5,683	4,776,788,162
EM191	3,990	49,084,620	418,300	59,730	4,789,620,542
EM204	12,745	79,505,409	570,565	44,252	4,779,024,662
EM205	53,697	135,486,275	162,823	5,242	4,786,831,622
EM208	3,313	41,992,026	546,341	64,999	4,790,740,682
EM209	45,871	116,311,093	205,124	5,017	4,788,571,862
EM219	19,319	85,434,733	555,766	14,559	4,796,211,602
EM227	12,837	54,647,530	336,634	26,765	4,794,620,762
EM232	31,942	121,533,960	236,309	10,432	4,794,202,982
EM238	7,351	39,150,291	769,101	48,224	4,793,427,362
EM242	9,797	60,568,098	641,916	35,719	4,796,499,242
EM251	11,087	65,526,982	357,199	21,319	4,786,586,462
EM268	32,610	97,409,224	575,002	8,616	4,789,756,442
EM275	9,814	59,736,375	813,547	24,504	4,796,447,222
EM283	35,823	110,290,277	284,712	7,512	4,788,078,482
EM293	9,512	57,120,941	733,123	53,850	4,795,731,362
EM305	8,755	59,052,708	446,907	34,979	4,793,440,322
EM308	14,486	80,827,555	411,604	21,742	4,794,984,362
EM326	16,809	82,168,325	511,025	37,604	4,795,947,722
EM337	32,940	84,242,311	257,796	5,093	4,798,092,962
EM338	44,075	125,050,275	132,015	7,202	4,793,679,182
Sum	649,576	2,197,804,580	11,723,877	NA	125,889,048,067
Average	24,058	81,400,170	434,218	24,031	4,662,557,336

Table 5 Number of sequence matches for genes encoding three SCFA-producing enzymes
 against 27 assembled metagenomes.

345 BCoAt: Butyryl-CoA transferase / Acetyl-CoA hydrolase; ACS: Acetate-formyltetrahydrofolate 346 synthetase/Formate-tetrahydrofolate ligase; PCoAt: Propionyl-CoA:succinate-CoA transferase / 347 Propionate CoA-transferase. Columns titled Matching genes show number of TBLASTN hits of 348 enzymes (BCoAt: GI|71081820|; ACS: GI|218353245| and GI|7242549|; PCoAt: GI|29346147| and 349 GI|260588698|) with an amino acid identity greater than 30% and an alignment length longer than 200 350 bp.

- 351
- 352

Subject	Location	Predicted	Matching BCoAt gapos	Matching	Matching PCoAt genes
EM039	Community	136 611	9	13	9
EM148	Rehab	52,810	10	10	7
EM172	Community	150.449	14	18	11
EM173	Longstav	65.189	2	6	4
EM175	Community	56.117	4	11	7
EM176	Community	60,966	7	12	12
EM177	Community	156,739	12	18	17
EM191	Longstay	43,959	1	10	11
EM204	Community	78,779	8	11	10
EM205	Community	183,002	11	11	13
EM208	Longstay	37,155	0	9	6
EM209	Rehab	158,301	16	14	16
EM219	Rehab	94,356	9	13	15
EM227	Longstay	57,754	3	8	8
EM232	Rehab	141,771	18	14	13
EM238	Longstay	38,639	3	6	5
EM242	Longstay	58,756	3	9	9
EM251	Community	65,721	4	10	8
EM268	Community	123,398	13	15	7
EM275	Longstay	59,297	8	10	8
EM283	Community	137,351	13	16	9
EM293	Longstay	55,734	5	8	8
EM305	Rehab	57,727	6	10	8
EM308	Longstay	80,512	8	11	12
EM326	Rehab	85,907	8	12	8
EM337	Rehab	108,711	7	13	6
EM338	Rehab	161,152	13	13	15
Community Total		1,149,133	95	135	103
Long-stay Total		496,995	33	77	71
Community Average		114,913	9.5	13.5	10.3
Long-stay Average		55,222	3.7	8.6	7.9

355

359	Table 6 Descriptive statistics of clinical and health measurements in 178 elderly subjects across
360	the four residence locations.

	Community	Day Hospital	Rehabilitation	Long-stay		
	(n=83)	(<i>n</i> =20)	(<i>n</i> =15)	(n=60)		
	Mean (SD)					
Age (yrs)	73.6 (6.5)	80.9 (6.6)	77.3 (6.9)	83.6 (7.5)		
Male (%)	43	65	53	23		
Weight (kg)	75.4 (16.5)	79.2 (17.3)	82.6 (16.9)	59.8 (13.2)		
BMI^{1} (kg/m ²)	27.4 (5.0)	28.5 (6.1)	30.5 (4.9)	23.4 (4.6)		
CC ² (cm)	36.2 (4.1)	37.3 (5.7)	36.3 (5.2)	29.8 (4.4)		
MAC ³ (cm)	28.2 (3.2)	29.6 (3.8)	30.9 (4.1)	26.7 (3.3)		
SBP ⁴ (mmHg)	140.6 (17.1)	136.8 (17.0)	122.3 (15.0)	128.0 (18.8)		
DBP ⁵ (mmHg)	76.0 (10.2)	71.5 (11.4)	65.6 (9.6)	70.8 (13.0)		
		Median (interc	quartile range)			
CCI ⁶ (0, 9)	0 (0, 1)	3 (2, 4)	3 (1, 4)	3 (2, 4)		
Barthel ⁷ (0, 20)	20 (20, 20)	20 (18, 20)	12 (10, 16)	2 (1, 7)		
FIM ⁸ (0, 126)	126 (125, 126)	124 (119, 126)	94 (82, 114)	36 (20, 61)		
MMSE ⁹ (0, 30)	29 (27, 30)	27 (25, 29)	25 (21, 27)	14 (2, 20)		
$MNA^{10}(0, 30)$	27 (26, 28)	25 (24, 27)	25 (23, 25)	17 (14, 21)		

¹Body Mass Index, ²Calf Circumference, ³Mid-Arm Circumference, ⁴Systolic Blood Pressure, ⁵Diastolic Blood Pressure

⁶Charlson Index of Comorbidity: Score out of a total of 22. Higher scores indicate higher degree of comorbidity

⁷Barthel Index of Activities of Daily Living: Score out of a total of 20. Higher scores indicate higher degree of independence

⁸Functional Independence Measure: Score out of a total of 126. Higher scores indicate higher degree of independence.

⁹Mini-Mental State Exam: Score range 0 (worst performance) – 30 (best performance)

¹⁰Mini-Nutritional Assessment: Score out of a total of 30. >= 24 = well-nourished; 17-23.5 = at risk of malnourishment; <17 = undernourished

369 Table 7: Effect of medications on health measures tested for microbiota correlations

370 Medication types are coded as per foot-note. The Associated Difference column shows the median change in the health

371 parameter measurement in subjects receiving the medication compared to those who did not, and is derived from the slope of

the medication regression in the median regression model adjusted for age, gender, location (for all 178 subjects), unweighted

373 UniFrac PCoA axes 1, 2 and 3, and any other medications that were added to the model for a specific health value. Asterisks

 $374 \qquad \text{indicate significant p-values: } p <= 0.05 \ (*), p <= 0.01 \ (**), p <= 0.001 \ (***).$

	Unweighted Uni	Frac PCoA for all four reside	nce locations (n = 178)	
Health measurement	Medication type	Number of subjects	Associated difference	<i>p</i> -value
GDT	Med L	28	-0.68	0.2
	Med M	37	1.2	0.064
	Med P	10	1.15	0.13
Diastolic BP	Med C	103	3 39	0.053
	Med D	48	-1 19	0.61
116	Med O	13	3.06	0.01
	Med R	56	1 58	0.18
	Med S	10	1.54	0.32
	Med T	9	4 53	0.18
11.8	Med 0	13	6.61	0.19
IL8	Med R	56	-4.28	0.24
	Med S	10	-3.28	0.6
	Med T	9	0.87	0.87
ΤΝΕα	Med Q	13	-0.24	0.75
	Med R	56	0.51	0.40
	Med S	10	2.88	0.1
	Med T	9	0.28	0.71
	Unweighted U	niFrac PCoA for Community-	only subjects (n = 83)	
Health measurement	Medication	Number of subjects	Associated difference	p-value
Diastolic BP	Med C	44	2.3	0.44
	Med D	15	-3.51	0.48
GDT	Med L	5	-1.25	0.12
	Med M	7	2.89	0.012*
	Unweighted U	UniFrac PCoA for Longstav-o	nly subjects (n = 60)	
	Medication			
Health measurement	type	Number of subjects	Associated difference	p-value
Barthel	Med C	29	-1.03	0.31
	Med D	13	2.15	0.21
	Med I	9	-0.95	0.55
	Med L	23	-1.48	0.15
	Med N	15	-1.2	0.38
	Med O	7	1.54	0.46
	Med Q	10	-4.74	0.009**
	Med R	44	1.05	0.41
	Med S	4	6.41	0.001**
	Med T	6	-1.96	0.33
FIM	Med C	29	-7.45	0.17
	Med D	13	15.01	0.15
	Med I	9	-10.89	0.29
	Med L	23	-2.28	0.71
	Med N	15	-5.86	0.49
	Med O	7	15.79	0.11
	Med Q	10	-17.6	0.062
	Med R	44	-1.08	0.91
	Med S	4	47.9	0.0004***
	Med T	6	-14.98	0.21
Diastolic BP	Med C	29	0.52	0.84
	Med D	13	-5.18	0.17
Systolic BP	Med C	29	-4.6	0.33
	Med D	13	-7.42	0.34
IL8	Med Q	9	-2.56	0.71
	Med R	40	-3.46	0.48
	Med S	4	-1.66	0.84
	Med T	4	-1.78	0.79
CRP	Med T	4	1.05	0.78

376 377 377 Med C = Cardiovascular Medications (Chronic Cardiac Failure / Angina / Hypertension) Med D = Diuretics Med I = Bronchodilators / Inhalers medications

Med K = Arthritis (osteo and rheumatoid) medications Med L = Insomnia medications

Med M = Mood Disorders (antidepressants / psychoses / anxiety) medications

Med N = Epilepsy / Seizures / Neuropathic Analgesia medications

Med O = Parkinsons medications Med P = Alzheimers / dementia medications

Med Q = Analgesia - Opiod medications Med R = Analgesia (Non - opiod) - medications

Med S = NSAIDs (non-steroidal anti-inflammatory drug) Med T = Steroids

Table 6. Chemical shifts of metabolites identified by NMR metabolomics.

	Metabolite	Chemical shift (multiplicity) and assignment
1	Cholate	0.73 ppm (s) CH ₃
2	Caprylate	0.86 ppm (t) CH ₃ , 1.28 ppm (m) CH ₂
3	Valerate	0.88 ppm (t) CH ₃ , 1.27 ppm (m) CH ₂ , 1.51 ppm CH ₂
4	Butyrate ^a	0.90 ppm (t) CH ₃ , 1.56 ppm (m) CH ₂ , 2.16 ppm (t) CH ₂
5	Isovalerate	0.91 ppm (d) CH ₃ , 2.06 ppm (d) CH ₂
6	Valine	0.99 ppm (d) CH ₃ , 1.04 ppm (d) CH ₃
7	Leucine	0.95 ppm (t) CH ₃
8	Isoleucine	0.94 ppm (t) CH ₃ , 1.01 ppm (d) CH ₃
9	Propionate	1.06 ppm (t) CH ₃ , 2.18 ppm (q) CH ₂
10	Threonine	1.32 ppm (d) CH ₃ , 3.59 ppm (d) CH
11	Lactate	1.34 ppm (d) CH ₃
12	Isocaproate	0.88 ppm (d) CH ₃ , 1.44 ppm (m) CH
13	Alanine	1.48 ppm (d) CH ₃
14	Lysine	1.72 ppm (m) CH ₂ , 3.02 ppm (t) CH ₂
15	Acetate	1.92 ppm (s) CH ₃
16	Glutamate	2.34 ppm (m) CH ₂ , 2.05 ppm (m) CH ₂
17	Succinate	2.39 ppm (s) CH ₂
18	Glutamine	2.46 ppm (m) CH ₂ , 2.11 (m) CH ₂
19	Methylamine	2.60 ppm (s) CH ₃
20	Aspartate	2.66 ppm (m) CH ₂ , 2.82 ppm (m) CH ₂
21	Trimethylamine	2.88 ppm (s) CH ₃
22	Malonate	3.11 ppm (s) CH ₂
23	Taurine	3.23 ppm (t) CH ₂ , 3.36 ppm (t) CH ₂
24	Glycine ^b	3.55 ppm (s) CH ₂
25	Glucose ^b	3.25 ppm (dd) CH, $3.90 ppm$ (dd) CH ₂ , $4.63 ppm$ (d) CH
26	Glutarate ^a	1.79 ppm (m) CH ₂
27	Lipid ^b	1.17 ppm
28	Acetone	2.23 ppm (s) CH ₃
29	Phenylacetate	3.54 ppm (s) CH ₃ , $7.28 ppm$ (m) CH, $7.37 ppm$ (m) CH
30	Tyrosine	6.89 ppm (m) CH, 7.19 ppm (m) CH
31	Phenylalanine	CH, 7.33 ppm (m) CH, 7.37 ppm (m) CH, 7.43 ppm (m) CH
32	Formate	8.46 ppm (s) CH ₂

^ahigher in faecal water of community subjects relative to long stay subjects ^bhigher in faecal water of long stay subjects relative to community subjects 386

t, triplet; q; quartet; m, multiplet.

Letters in brackets indicate the multiplicity of the peak. s, singlet; d, doublet; dd, doublet of doublets;

392 <u>Supplementary Notes</u>393

394 Microbiota-metabolome co-inertia analysis395

To aid interpretation of the relationship between the genus-level microbiota and metabolomics datasets, we integrated them using a multivariate method known as co-inertia analysis (CIA¹⁸). Principal components analysis (PCA) can be applied to each of the individual datasets, and these PCAs are used as inputs for the CIA which then identifies ordinations of the two datasets that are maximally co-variant and so identifies the shared biological trends within the two datasets. The method is an unsupervised approach and is insensitive to a high variable-to-sample ratio¹⁹.

402 CIA was applied to the genus-level microbiota composition and metabolome data 403 (Supplementary Figure 6). As noted in Main text, this analysis revealed a significant 404 relationship between the two datasets with an RV coefficient of 0.474, indicating a strong co-405 structure between the two datasets. The first two components of the CIA account for 58.9% of 406 the variance in the datasets, with component 1 (horizontal) accounting for 48.4% of the 407 variance, and the second, component 2 (vertical) accounting for another 10.5%.

408 Community and long-stay samples are separated from each other along the primary axis in the 409 analysis. In addition to differentially abundant metabolites mentioned in the main text, there was an 410 increase in glycine, glucose and lipid levels in the long-stay subject faecal metabolomes. This could be 411 because the associated microbiota may have a reduced ability to metabolize these compounds, or that 412 there is a reduced uptake of these nutrients by the gut in long-stay subjects, that may or may not relate 413 their microbiota.

414 Using a cut-off of 1% abundance, the main genera associated with the community 415 metabolome along the first axis were *Bacteroides*, *Faecalibacterium*, *Ruminococcus*, *Oscillibacter*, 416 *Coprococcus* and *Sporobacter*. *Bacteroides* spp are known to produce acetate²⁰ but are found to be 417 associated with diets rich in animal produces and low in fruit and veg in this and other studies²¹. This 418 and their lack of cellulolytic ability²² may explain the strong association with acetate along the first 419 axis but the minor inverse association with the second axis. Instead *Sporobacter* is the genus most 420 closely related to the production of acetate according to this analysis. When the reads mapping to this 421 genus were investigated, we found that over 70% could be confidently assigned to the species 422 *Sporobacter termitidis. Sporobacter termitidis* is known to produce only acetate as an end 423 fermentation product²³.

A number of genera are associated with the higher faecal levels of butyrate -Faecalibacterium, Ruminococcus, Oscillibacter and Coprococcus. The identified species that have the highest correlation and abundance with these genera are Faecalibacterium prausnitzii, Ruminococcus bromii, Coprococcus eutactus and Oscillibacter valericigenes.

428

429 Properties of bacterial co-abundance groups (CAGs).

430 Associations between individual genera were determined using the Kendall correlation coefficient. 431 The full set of associations was visualized and clustered in R using the Made4 package²⁴ and the 432 function *Heatplot*, whereby Hierarchical clustering with the Pearson correlation distance metric and 433 Ward clustering was used to define co-abundant groups of genera (Supplementary Fig 12A). All 434 significant positive associations are shown in Supplementary Fig 12B. These associations were 435 controlled for multiple testing using the qvalue method¹⁶ and only those with false discovery rate < 436 0.05 were retained (Supplementary Table 8).

437 The Kendall correlations were converted to a correlation distance metric and used as input for 438 a Permutational MANOVA to determine if the CAGs were significantly different from each other. 439 Essentially this compared strength of the correlations between the groups to correlation strengths within the groups in a pairwise manner. Permutational MANOVA²⁵ was performed using the vegan 440 441 package in R. All five external CAGs displayed significantly different inter-relationships from each 442 other (p < 0.0001), except for the relationship between the *Bacteroides* CAG and the middle 443 Odoribacter CAG which was the least distinct, with a significance of 0.052. The Bacteroides CAG has 444 the smallest number of significant positive correlations between its members, giving it a weak 445 coherence, and is defined more by significant negative correlations with the other four external CAGs 446 but not against the middle *Odoribacter* group. Therefore the *Bacteroides* CAG was strongly negatively 447 associated with the other CAGs but not with the middle group.

448 The major genera that dominate the CAGs are characterized by their genus-associated abilities 449 to produce a wide range of short-chain fatty acids, including butyrate, acetate, lactate, propionate, 450 formate, and succinate as well as ethanol, hydrogen, and carbon dioxide. However, the Oscillibacter 451 CAG shows a large proportion of genera able to produce acetate, and only the acidaminococci are 452 reported to produce butyrate²⁶. The other CAGs also contain genera known to produce acetate, but 453 many of these fermentative genera are also able to synthesize a wider range of SCFAs when compared 454 to the genera in the Oscillibacter CAG. Genera able to use the fermentation products include 455 Desulfovibrio and Methanobrevibacter which are able to use lactate or ethanol as electron donors and 456 acetate as a carbon source respectively. The *Desulfovibrio* may also be using the sulphate liberated by 457 the Akkermansia species from mucin, as an electron acceptor.

458

459 **Description of Dietary Groups (DGs)**

DG1: The predominant features of this diet type include consumption of complex carbohydrates (including wholegrain breakfast cereals, breads, boiled potato), daily consumption of a wide range of fruit and vegetables and moderate (5 times/week) consumption of protein-rich white meat, fish and eggs. Red-meat was not consumed by this group while oily fish were consumed once/week with low intakes of dairy produce (approx. 3 times/week) and high-sugar/low-nutrient dense foods.

DG2: Both complex (wholegrain breakfast cereals and breads, boiled potatoes) and simple carbohydrates (white bread) were consumed frequently in this diet type. A lower variety of fruit and vegetables were consumed, less frequently (two-three times/daily) compared to DG1. Red-meat, fish or eggs were consumed daily with no consumption of oily fish. Intakes of dairy produce were lower among this group (approx. once weekly) with higher intakes of high-sugar/low-nutrient dense foods compared to diet type 1.

471 DG3: Overall, this diet group contained the least variety of all the DG's. Porridge and mashed potato 472 were the main daily staples while simple carbohydrate (white bread) was consumed frequently with no 473 inclusion of wholemeal varieties. Consumption and variety of fruit and vegetables were lowest overall 474 among this group (approx. 1-2 portions daily). Processed and unprocessed meat and eggs were 475 consumed once-twice/week with no fish consumed. Intakes of dairy produce are low (approx. 3 times
476 week) with the lowest consumption of high-sugar/low-nutrient dense foods.

477 DG4: Similar to diet type 3, mashed potato and porridge were the only staples in this diet type and 478 were consumed daily. Fruit and vegetable consumption was comparable with that of diet group 1 but 479 with much less variety. Processed and unprocessed meat and fried fish were consumed once-480 twice/week. Fish were not consumed by this group. Dairy products and high-sugar/low-nutrient dense 481 foods were consumed most frequently by this cohort (once-twice and 3-4 times daily, respectively), 482 predominantly full-fat dairy produce and puddings, and sweetened hot beverages which were 483 consumed 1-3 times daily.

484

485 Analysis of possible confounders of microbiota-health associations

486 Antibiotics. None of the subjects had been treated with antibiotics in the month prior to sampling; 487 extending this exclusion window would impact negatively on recruitment rates of older subjects. To 488 examine the effect of treatment prior to this exclusion window, we used quantile (median) regression 489 to examine the association between microbiota alpha diversity and time in days since last antibiotic 490 usage, extending before the one-month recruitment exclusion window. We also tested correlations 491 between the (microbiota) PCoA axes and time since last antibiotic usage. These models were adjusted 492 for age, gender and location, (as for the main microbiota-health analysis for Table 1). No relationship 493 between microbiota diversity and time since last antibiotic treatment was established (data not shown). 494 Other medications. We adjusted median regression models for the effect of medications on cognate 495 health read-outs. The actual effects of medication on relevant clinical scores in that model are 496 presented in Supplementary Table 8. The majority of medications had small non-significant effects on 497 the relevant health measure, which did not affect the significance of the relationship between the 498 health variables and the microbiota presented in Table 1.

Diet. Although diet clearly impacts on microbiota; the composition of the diet was expected have a direct health effect independent of a diet-microbiota-health effect. To explore this issue, we repeated the median regression analysis, controlling for diet, using the Health Food Diversity (HFD) index. Although diet explained a number of the associations, the majority of the frailty indicators-microbiota associations reported in Table 1 were still significant. Two variables, weight and BMI no longer had
 significant associations with microbiota after adjustment for HFD.

- 505
- 506 Aggregate microbiota composition in the elderly

507 We previously showed that there was a higher Firmicutes/Bacteroidetes ratio among nine 508 younger subjects compared to 118 elderly subjects². When we analysed 178 additional faecal samples 509 from elderly subjects, and four additional samples from younger healthy individuals, we did not 510 observe an aggregate microbiota dominated by phylum Bacteroidetes (Suppl. Fig 15). At genus level 511 there were significantly higher levels of Ruminococcus and Blautia and lower levels of 512 Escherichia/Shigella in young controls compared to the microbiota of the elderly subjects (Suppl. Fig 513 16). Clearly the balance in a study population of individuals from the Bacteroidetes-Prevotella-514 Ruminococcus-dominated groups (see main text) will affect aggregate estimations. To adequately infer 515 microbiota differences between young and old populations a larger number of samples from the 516 former cohort are needed, including information about food frequency consumption, which affects 517 phylum proportions (ref.²¹; this study).

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519 Denoised versus Un-denoised pyrosequencing analysis

Initially we denoised the pyrosequencing sequences using Denoiser²⁷ as part of the OTU 520 521 picking approach according to recommendations from the QIIME documentation. We detected a run-522 specific bias in the denoised dataset when comparing OTU clustering of OTUs generated with and 523 without a denoising step. Since OTUs were picked separately for each run at flow-gram level, prior to 524 an overall OTU picking at sequence level, an over-clustering within each run seemed to have taken 525 place. This happened in spite of using the recommended "exact" options for the UClust OTU picker. 526 Suppl. Fig. 17 shows the run-specific bias and also how the clustering according to community 527 location is retained for both denoised and un-denoised data sets.

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