UC San Diego UC San Diego Previously Published Works

Title

Pangenome comparison of Bacteroides fragilis genomospecies unveils genetic diversity and ecological insights

Permalink https://escholarship.org/uc/item/2xf4449c

Journal mSystems, 9(7)

ISSN 2379-5077

Authors

Oles, Renee E Terrazas, Marvic Carrillo Loomis, Luke R <u>et al.</u>

Publication Date

2024-07-23

DOI

10.1128/msystems.00516-24

Peer reviewed



8 Host-Microbial Interactions Observation



Pangenome comparison of *Bacteroides fragilis* genomospecies unveils genetic diversity and ecological insights

Renee E. Oles,^{1,2} Marvic Carrillo Terrazas,¹ Luke R. Loomis,¹ Chia-Yun Hsu,¹ Caitlin Tribelhorn,² Pedro Belda-Ferre,² Allison C. Ea,¹ MacKenzie Bryant,² Jocelyn A. Young,^{2,3} Hannah C. Carrow,¹ William J. Sandborn,^{4,5} Parambir S. Dulai,^{4,6} Mamata Sivagnanam,^{2,3} David Pride,^{1,5,7,8} Rob Knight,^{2,5,9,10,11} Hiutung Chu^{1,5,12}

AUTHOR AFFILIATIONS See affiliation list on p. 5.

ABSTRACT *Bacteroides fragilis* is a Gram-negative commensal bacterium commonly found in the human colon, which differentiates into two genomospecies termed divisions I and II. Through a comprehensive collection of 694 *B. fragilis* whole genome sequences, we identify novel features distinguishing these divisions. Our study reveals a distinct geographic distribution with division I strains predominantly found in North America and division II strains in Asia. Additionally, division II strains are more frequently associated with bloodstream infections, suggesting a distinct pathogenic potential. We report differences between the two divisions in gene abundance related to metabolism, virulence, stress response, and colonization strategies. Notably, division II strains harbor more antimicrobial resistance (AMR) genes than division I strains. These findings offer new insights into the functional roles of division I and II strains, indicating specialized niches within the intestine and potential pathogenic roles in extraintestinal sites.

IMPORTANCE Understanding the distinct functions of microbial species in the gut microbiome is crucial for deciphering their impact on human health. Classifying division II strains as *Bacteroides fragilis* can lead to erroneous associations, as researchers may mistakenly attribute characteristics observed in division II strains to the more extensively studied division I *B. fragilis*. Our findings underscore the necessity of recognizing these divisions as separate species with distinct functions. We unveil new findings of differential gene prevalence between division I and II strains in genes associated with intestinal colonization and survival strategies, potentially influencing their role as gut commensals and their pathogenicity in extraintestinal sites. Despite the significant niche overlap and colonization patterns between these groups, our study highlights the complex dynamics that govern strain distribution and behavior, emphasizing the need for a nuanced understanding of these microorganisms.

KEYWORDS pangenome, commensal bacteria, genomic diversity, niche adaptation, *Bacteroides*

B acteroides fragilis is a persistent colonizer of the human gut linked to both health and disease (1) and is composed of two genomospecies termed divisions I and II. They have primarily been differentiated through the presence of *cepA*, a beta-lactamase, which is unique to division I (2), and the chromosomally encoded carbapenemase gene (*cfiA* or *ccrA*), which is unique to division II and provides resistance to beta-lactamase inhibitors (3, 4). Due to their genetic similarity, traditional methods such as 16S rRNA gene analysis cannot distinguish between these divisions, yet they share an average nucleotide identity (ANI) of 87%, below the typical species cutoff of 96% (3, 5–10). Here, we conduct a comprehensive genomic comparison and identified genes conserved **Editor** Sean M. Gibbons, Institute for Systems Biology, Seattle, Washington, USA

Address correspondence to Hiutung Chu, hiuchu@health.ucsd.edu.

The authors declare a conflict of interest.

See the funding table on p. 6.

Received 15 April 2024 **Accepted** 31 May 2024 **Published** 27 June 2024

Copyright © 2024 Oles et al. This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license.

mSystems

within each *B. fragilis* division, but not shared between them, shedding light on the unique biological roles and functions of these divisions within their ecological niches.

We analyze 694 *B. fragilis* whole genome sequences, including 139 from our own collection, which we isolated and sequenced for the first time, and the remaining from public sources (Tables S1 and S2). To compare the genetic relatedness between divisions, we employed Mash, a whole genome k-mer-based approach (11) to determine the genetic distance between each strain (Fig. 1A). Metric multidimensional scaling (mMDS) reveals a clear separation of strains into two distinct divisions (Fig. 1A). To further support this distinction, we discovered a significant difference in GC content between the divisions (Welch's t-test, P = 8.1e-5; Cohen's effect size, d = 0.35) (Fig. 1B), although no differences were found in genome size (Welch's t-test, P = 0.22) (Fig. 1C). We also observe a difference in the average GC content in the core genes (present in >99% of



FIG 1 B. fragilis is composed of two monophyletic divisions. (A) Metric multidimensional scaling (mMDS) of the k-mer based Mash distances of 694 strains, colored by divisions I (green, n = 554) and II (purple, n = 140). (B) GC content (%) of isolate assemblies in division I and II isolates. Average for division I = 43.35% ± 0.19 and division II = 43.42% ± 0.16 (P = 8.1e-5, Welch's t-test with unequal variance; n = 694). (C) Genome size (bp) of isolate assemblies in division I and II isolates. Average for division I = 5.26×10^6 bp and division II = 5.22×10^6 bp (P = 0.22, Welch's t-test with unequal variance; n = 694). (D) The proportion of isolates originating from abscess (P = 0.18), blood (P = 0.0049), and fecal (P = 0.0011) samples in division I compared with division II, P-values from Fisher's exact test. Division I: total = 554, fecal = 228, blood = 30, abscess = 51; division II: total = 140, fecal = 56, blood = 21, abscess = 23. The proportion of isolates originating from Africa (P = 0.18), Asia (P = 2.2e-16), Europe (P = 0.00019), or North America (P = 2.2e-16) in division I compared with division II, P-values from Fisher's Exact Test. Division I: total = 554, Africa = 33, Asia = 46, Europe = 11, North America = 459; division II: total = 140, Africa = 13, Asia = 58, Europe = 13, North America = 56. (E) Distribution of isolates in each continent per division in the Pasoli et al., 2019 data set. The proportion of isolates originating from Africa (P = 1), Asia (P = 3.9e-08), Europe (P = 9.4e-07), North America (P = 0.029), Oceania (P = 0.00017), or South America (P = 1) in division I (green) compared with division II (purple), P-values from Fisher's exact test. Division I: n = 437, Africa = 1, Asia = 107, Europe = 263, North America = 63, Oceania = 1, South America = 2; Division II: n = 65, Africa = 0, Asia = 39, Europe = 18, North America = 3, Oceania = 5, South America = 0. (F) Phylogenetic tree of the core genome alignment of 694 strains through maximum likelihood, midpoint rooted, colored by divisions I (green) and II (purple). (G) The phylogenetic tree of the core genome alignment of division I strains through maximum likelihood, midpoint rooted, annotated with the inner ring, Group: healthy, infection, IBD, non-human animal, unknown; and outer ring, Continent: Asia, Africa, Europe, Oceania, North America, and South America (n = 554).(H) The phylogenetic tree of the core genome alignment of division Il strains through maximum likelihood, midpoint rooted, annotated with the inner ring, Group: healthy, infection, IBD, non-human animal, unknown; and outer ring, Continent: Asia, Africa, Europe, Oceania, North America, and South America (n = 140).

isolates) of divisions I (44.6% ± 4.1) and II (45.0% ± 4.0), demonstrating the same trend where division II strains have a moderately higher average GC content than division I (Welch's *t*-test, P = 8.3e-21). Of the shared core genes in division I versus II, the average GC content per gene in divisions I and II is 44.9% ± 3.7 and 45.0% ± 3.8, respectively (Welch's *t*-test, P = 0.0081). However, core genes exclusive to division I have an average GC content of 43.0% ± 5.5, whereas those unique to division II are 44.2% ± 5.7 (Welch's *t*-test, P = 4.2e-19), suggesting the differences in GC content stem from recent evolution between divisions. Although significant, the GC content difference is subtle and may not accurately categorize any given isolate as either division I or II. Finally, based on the maximum likelihood, midpoint-rooted phylogeny of the core genome alignment, divisions I and II separate into discrete clades (Fig. 1F).

We next investigated whether divisions I and II are associated with disease states, isolation sites, or other metadata categories. Division I strains are more prevalent (80% of the total; 554 of 694) than division II. Among the 409 isolates from abscesses, fecal samples, or blood, division I strains are more commonly isolated from fecal samples (74%) compared with division II (56%, Fisher's exact test, P = 0.0011) (Fig. 1D). Conversely, division II strains are more frequently associated with abscesses (23%) or blood (21%) compared with division I strains (16% from abscess, Fisher's exact test, P = 0.18, and 10% from blood, Fisher's exact test, P = 0.0049) (Fig. 1D). Notably, division I and II strains exhibit variations in the continent of isolation. Moreover, 84% (n = 459) of division I strains originate from North America, compared with only 40% (n = 56) of division II strains (Fisher's exact test, P = 2.2e-16) (Fig. 1D and H). In contrast, only 8% of division I strains originate from Asia (n = 46), compared with 41% (n = 58) of division II strains (Fisher's exact test, P = 2.2e-16) (Fig. 1D and G). To further explore the geographical distribution of these divisions, we examined 502 species-genome bins (SGBs) classified as B. fragilis, which were reconstructed from 9,428 human gut metagenomic samples worldwide (12). 87% and 13% of strains belong to divisions I and II, respectively. No host harbor both divisions, in line with reports from other studies (13-16). Most of the division I strains (75%) originate from Europe or North America, whereas most division II strains (60%) are from Asia (Fig. 1E). This aligns with previous reports indicating a higher rate of cfiA+isolates (division II) in Japan, Hong Kong, and India (17). This geographic disparity suggests the under-representation of division II strains in public databases may be due to the limited sampling of specific populations (18).

Using Panpiper (19), we compared the pangenomes of B. fragilis division I and II, and identified 794 genes with differential prevalence, including the exclusive presence of cfiA in division II and cepA in division I (Fig. 2A, B and E; Table S3) (2, 4). We next assessed the differential abundance of carbohydrate-active enzymes, along with reference metabolic (EC) and reference KEGG orthology pathways (KEGG KO) (Fig. 2C through E). Our analysis reveals division-specific metabolic capabilities and potential ecological niches. Division II strains have genes favoring the degradation of plant cell walls, including glycosyl hydrolases (GH5, GH9, GH51, and GH95) (Fig. 2C), suggesting adaptation to dietary variations. Specifically, BFAG_03498 (ko:K01179, GH9) is predicted to mediate the breakdown of cellulose, BFAG_02344 (GH51) is involved in the breakdown of arabinose-containing polysaccharides, and BFAG_0465 (GH95), an alpha-L-fucosidase, is involved in the cleavage of internal beta-1,4-glycosidic bonds present in plant cell walls (20) (Table S3). One possible explanation for the higher prevalence of plant cell wall degradation genes in division II strains may be dietary differences among hosts of divisions I and II, potentially linked to their distinct geographical distributions (Fig. 1D and E) (21). Division I strains harbor genes indicative of complex carbohydrate degradation, a hallmark feature of gut-resident commensal Bacteroides (1, 22). This includes two predicted alpha-L-rhamnosidases (BF9343_0522, BF9343_0310; GH78), which are core genes exclusive to division I (Fig. 2C; Table S3). Division I strains also exhibit an enrichment of GH33 sialidases (Fig. 2C), which catalyze the cleavage of terminal sialic acid residue. Although sialidases have been linked to virulence (23), the B. fragilis GH33 sialidase mediates intestinal colonization and persistence during early life (24).

Observation



FIG 2 *B. fragilis* divisions I and II segregated by multiple differentially abundant genes and gene categories. (A) Relative \log_2 gene abundance heatmap by division, where genes are clustered by R pheatmap complete method, annotated by regions of gene clusters core to both divisions, core only to division I, or core only to division II. (B) Histogram of \log_2 -fold change of prevalence between all genes in division I versus II. (C-E) \log_2 average number of genes per isolate in categories, (C) carbohydrate-active enzymes (CAZy) (\log_2 -fold change ≥ 0.5), (D) EC category (\log_2 -fold change ≥ 1), and (E) KEGG KO (\log_2 -fold change ≥ 0.5) between divisions I and II, displaying categories significant by Kruskal–Wallis test (corrected $P \leq 0.01$). Legend is \log_2 average number of genes per isolate in each category. (F) Total number of antimicrobial resistance (AMR) genes per isolate for each division; P = 0.004, Welch's *t*-test. (G) The percentage of isolates per division with each antimicrobial resistance gene. *cfr*, chloramphenicol–florfenicol resistance gene; *cfx*, cefuroxime resistance gene, *tet*, tetracycline resistance gene; *mef*, macrolide efflux gene; *msr*, macrolide efflux gene, *nim*, nitroimidazole resistance gene; *sul*, sulfonamide resistance gene; *tet*, tetracycline resistance gene.

Because sialic acid is identified in capsular polysaccharides and lipooligosaccharides (25), its presence may influence colonization and interactions within the host. Division I strains are also enriched in the type VI secretion system GA3, with 84.4% of division I strains having all T6SSiii GA3 structural genes (BF9343_1919–1925, 1931, 1940–1943) (26) compared with 48.1% in division II. This system, exclusive to *B. fragilis*, is recognized for mediating intra-strain competition and colonization dynamics (27–29). The differential abundance of glycosyl hydrolases and T6SSiii GA3 suggests distinct colonization strategies between division I and II strains within the gut.

Division I and II strains may occupy distinct ecological niches, distinguished by genes associated with metabolism and pathogenicity. Division II strains exhibit an increased abundance in genes related to proline degradation and glutamate synthesis pathways (EC 3.4.21.26, BFAG_03703; EC 1.5.5.2, BFAG_03859) (Fig. 2D; Table S3). Additionally, these strains have an increased abundance of the gene encoding DNA-formamidopyrimidine glycosylase (EC 3.2.2.23, BFAG_03121), crucial for DNA repair mechanisms against mutagenesis and cell death induced by alkylating agents (Fig. 2D; Table S3). We also observed differential prevalence in genes and pathways related to multidrug resistance. Division I strains have an increased prevalence of gamma-carboxymuconolactone

decarboxylase (EC 4.1.1.44) (Fig. 2D) associated with the breakdown of aromatic compounds and antimicrobial resistance (AMR) (30). We indentify a putative erythromycin esterase that detoxifies macrolides also more abundant in division I (31). Conversely, division II strains have a higher abundance of efflux proteins (K09771, K11741) (Fig. 2E; Table S3) and virginiamycin A acetyltransferase (*vat*, K18234), providing resistance to streptogramins (Fig. 2E; Table S3). Indeed, division II strains harbor a higher number of known AMR genes per isolate compared with division I (P = 0.004) (Fig. 2F and G), indicating a potential for increased virulence. Further experimental studies are necessary to determine the functional impact of division-specific genes to understand their roles and interactions within the intestinal ecosystem and host. Collectively, our comparative genomics study unveils distinct geographical distribution and genetic signatures within *B. fragilis* divisions, offering insights into their intricate interactions with the host and respective ecological niches.

ACKNOWLEDGMENTS

We thank members of the Chu lab for technical support and helpful discussions.

This work was supported by grants from the National Institute of Health (NIH) R01 Al167860 and P30 DK120515. Additional support was provided to H.C. by the Chiba University-UC San Diego Center for Mucosal Immunology, Allergy and Vaccines (cMAV), CIFAR Humans and the Microbiome Program, and The Hartwell Foundation.

Support to R.E.O. was provided by T32 AR064194 (NIAMS). Support to M.C.T. was provided by T32 DK007202 (NIDDK), the National Academies of Sciences, Engineering and Medicine through the Predoctoral Fellowship of the Ford Foundation, and the Howard Hughes Medical Institute (HHMI) Graduate Fellowships grant (GT15123). Support to J.A.Y. was provided by T32 DK007202 (NIDDK). Support to M.S was provided by NIH DK120515 and T32 DK007202. This publication includes data generated at the UC San Diego IGM Genomics Center utilizing an Illumina NovaSeq 6000 that was purchased with funding from a National Institutes of Health SIG grant (S10 OD026929).

W.J.S's current conflicts of interest are: Mirador Therapeutics (stock, employee, company officer, Ventyx Biosciences (stock, former employee), Prometheus Laboratory (board of directors), Shoreline Biosciences (stock, scientific advisory board), Forbion (consultant), Alimentiv (consultant). R.K.'s current conflicts of interest are: Gencirq (stock and SAB member), DayTwo (consultant and SAB member), Cybele (stock and consultant), Biomesense (stock, consultant, SAB member), Micronoma (stock, SAB member, co-founder), and Biota (stock, co-founder).

AUTHOR AFFILIATIONS

¹Department of Pathology, University of California, San Diego, California, USA

²Department of Pediatrics, School of Medicine, University of California, San Diego, California, USA

³Rady Children's Hospital, San Diego, California, USA

⁴Division of Gastroenterology, University of California, San Diego, California, USA

⁵Center for Microbiome Innovation, University of California, San Diego, California, USA

⁶Division of Gastroenterology, Northwestern University, Chicago, Illinois, USA

⁷Center for Innovative Phage Applications and Therapeutics (IPATH), University of California, San Diego, California, USA

⁸Center of Advanced Laboratory Medicine (CALM), University of California, San Diego, California, USA

⁹Shu Chien-Gene Lay Department of Bioengineering, University of California, San Diego, California, USA

¹⁰Department of Computer Science & Engineering, University of California, San Diego, California, USA

¹¹Halıcıoğlu Data Science Institute, University of California, San Diego, California, USA

¹²Chiba University-UC San Diego Center for Mucosal Immunology, Allergy and Vaccines (cMAV), University of California, San Diego, California, USA

AUTHOR ORCIDs

Renee E. Oles ^(b) http://orcid.org/0000-0001-5945-0215 Marvic Carrillo Terrazas ^(b) http://orcid.org/0000-0002-8830-5161 Chia-Yun Hsu ^(b) http://orcid.org/0000-0002-5283-1020 Pedro Belda-Ferre ^(b) http://orcid.org/0000-0001-6532-1161 MacKenzie Bryant ^(b) http://orcid.org/0000-0003-0749-2995 Jocelyn A. Young ^(b) http://orcid.org/0000-0002-9364-1928 Hannah C. Carrow ^(b) http://orcid.org/0000-0003-0463-0811 Mamata Sivagnanam ^(b) http://orcid.org/0000-0002-3643-1326 Rob Knight ^(b) http://orcid.org/0000-0002-975-9019 Hiutung Chu ^(b) http://orcid.org/0000-0001-7489-0446

FUNDING

Funder	Grant(s)	Author(s)
HHS NIH National Institute of Allergy and Infectious Diseases (NIAID)	R01 AI167860	Hiutung Chu
HHS NIH National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK)	P30 DK120515	Hiutung Chu
HHS NIH National Institute of Arthritis and Musculoske- letal and Skin Diseases (NIAMS)	T32AR064194	Renee E. Oles

ADDITIONAL FILES

The following material is available online.

Supplemental Material

Supplemental methods (mSystems00516-24-s0001.docx). Additional experimental details and methods.

Table S1 (mSystems00516-24-s0002.docx). Isolation data for newly isolated and/or sequenced strains.

Table S2 (mSystems00516-24-s0003.docx). Accession numbers and source for *B. fragilis* strains from public repositories.

 Table S3 (mSystems00516-24-s0004.docx). Genes differentially prevalent between division I and II strains.

REFERENCES

- 1. Wexler HM. 2007. Bacteroides: the good, the bad, and the nitty-gritty. Clin Microbiol Rev 20:593–621. https://doi.org/10.1128/CMR.00008-07
- Parker AC, Smith CJ. 1993. Genetic and biochemical analysis of a novel Ambler class A beta-lactamase responsible for cefoxitin resistance in Bacteroides species. Antimicrob Agents Chemother 37:1028–1036. https://doi.org/10.1128/AAC.37.5.1028
- Gutacker M, Valsangiacomo C, Piffaretti J-C. 2000. Identification of two genetic groups in *Bacteroides fragilis* by multilocus enzyme electrophoresis: distribution of antibiotic resistance (cfiA, cepA) and enterotoxin (bft) encoding genes. Microbiology (Reading) 146:1241–1254. https:// doi.org/10.1099/00221287-146-5-1241
- Rasmussen BA, Gluzman Y, Tally FP. 1990. Cloning and sequencing of the class B beta-lactamase gene (ccrA) from *Bacteroides fragilis* TAL3636. Antimicrob Agents Chemother 34:1590–1592. https://doi.org/10.1128/ AAC.34.8.1590
- English J, Newberry F, Hoyles L, Patrick S, Stewart L. 2023. Genomic analyses of *Bacteroides fragilis*: subdivisions I and II represent distinct species. J Med Microbiol 72. https://doi.org/10.1099/jmm.0.001768

- Johnson JL. 1978. Taxonomy of the *Bacteroides*. Int J Syst Evol Microbiol 28:245–256. https://doi.org/10.1099/00207713-28-2-245
- Podglajen I, Breuil J, Casin I, Collatz E. 1995. Genotypic identification of two groups within the species *Bacteroides fragilis* by ribotyping and by analysis of PCR-generated fragment patterns and insertion sequence content. J Bacteriol 177:5270–5275. https://doi.org/10.1128/jb.177.18. 5270-5275.1995
- Ruimy R, Podglajen I, Breuil J, Christen R, Collatz E. 1996. A recent fixation of cfiA genes in a monophyletic cluster of *Bacteroides fragilis* is correlated with the presence of multiple insertion elements. J Bacteriol 178:1914–1918. https://doi.org/10.1128/jb.178.7.1914-1918.1996
- Nagy E, Becker S, Sóki J, Urbán E, Kostrzewa M. 2011. Differentiation of division I (cfiA-negative) and division II (cfiA-positive) *Bacteroides fragilis* strains by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry. J Med Microbiol 60:1584–1590. https://doi.org/10.1099/ jmm.0.031336-0
- Wallace MJ, Jean S, Wallace MA, Burnham C-AD, Dantas G. 2022. Comparative genomics of *Bacteroides fragilis* group isolates reveals

species-dependent resistance mechanisms and validates clinical tools for resistance prediction. mBio 13:e0360321. https://doi.org/10.1128/mbio.03603-21

- Ondov BD, Treangen TJ, Melsted P, Mallonee AB, Bergman NH, Koren S, Phillippy AM. 2016. Mash: fast genome and metagenome distance estimation using MinHash. Genome Biol 17:132. https://doi.org/10.1186/ s13059-016-0997-x
- Pasolli E, Asnicar F, Manara S, Zolfo M, Karcher N, Armanini F, Beghini F, Manghi P, Tett A, Ghensi P, Collado MC, Rice BL, DuLong C, Morgan XC, Golden CD, Quince C, Huttenhower C, Segata N. 2019. Extensive unexplored human microbiome diversity revealed by over 150,000 genomes from metagenomes spanning age, geography, and lifestyle. Cell 176:649–662. https://doi.org/10.1016/j.cell.2019.01.001
- Zhao S, Lieberman TD, Poyet M, Kauffman KM, Gibbons SM, Groussin M, Xavier RJ, Alm EJ. 2019. Adaptive evolution within gut microbiomes of healthy people. Cell Host Microbe 25:656–667. https://doi.org/10.1016/j. chom.2019.03.007
- 14. Yassour M, Vatanen T, Siljander H, Hämäläinen A-M, Härkönen T, Ryhänen SJ, Franzosa EA, Vlamakis H, Huttenhower C, Gevers D, Lander ES, Knip M, Xavier RJ, DIABIMMUNE Study Group. 2016. Natural history of the infant gut microbiome and impact of antibiotic treatment on bacterial strain diversity and stability. Sci Transl Med 8:343ra81. https:// doi.org/10.1126/scitranslmed.aad0917
- Verster AJ, Ross BD, Radey MC, Bao Y, Goodman AL, Mougous JD, Borenstein E. 2017. The landscape of type VI secretion across human gut microbiomes reveals its role in community composition. Cell Host Microbe 22:411–419. https://doi.org/10.1016/j.chom.2017.08.010
- Rashidan M, Azimirad M, Alebouyeh M, Ghobakhlou M, Asadzadeh Aghdaei H, Zali MR. 2018. Detection of *B. fragilis* group and diversity of bft enterotoxin and antibiotic resistance markers cepA, cfiA and nim among intestinal Bacteroides fragilis strains in patients with inflammatory bowel diseas. Anaerobe 50:93–100. https://doi.org/10.1016/j. anaerobe.2018.02.005
- Cao H, Liu MC-J, Tong M-K, Jiang S, Lau A, Chow K-H, Tse CW-S, Ho P-L. 2022. Diversity of genomic clusters and CfiA/cfiA alleles in *Bacteroides fragilis* isolates from human and animals. Anaerobe 75:102567. https:// doi.org/10.1016/j.anaerobe.2022.102567
- Abdill RJ, Adamowicz EM, Blekhman R. 2022. Public human microbiome data are dominated by highly developed countries. PLoS Biol 20:e3001536. https://doi.org/10.1371/journal.pbio.3001536
- Oles R, et al. 2023. rolesucsd/Panpiper. GitHub repository. https://doi. org/10.5281/zenodo.11186447
- Wu H, Owen CD, Juge N. 2023. Structure and function of microbial α-lfucosidases: a mini review. Essays Biochem 67:399–414. https://doi.org/ 10.1042/EBC20220158
- 21. De Angelis M, Ferrocino I, Calabrese FM, De Filippis F, Cavallo N, Siragusa S, Rampelli S, Di Cagno R, Rantsiou K, Vannini L, Pellegrini N, Lazzi C,

Turroni S, Lorusso N, Ventura M, Chieppa M, Neviani E, Brigidi P, O'Toole PW, Ercolini D, Gobbetti M, Cocolin L. 2020. Diet influences the functions of the human intestinal microbiome. Sci Rep 10:4247. https://doi.org/10. 1038/s41598-020-61192-y

- Pudlo NA, Urs K, Crawford R, Pirani A, Atherly T, Jimenez R, Terrapon N, Henrissat B, Peterson D, Ziemer C, Snitkin E, Martens EC. 2022. Phenotypic and genomic diversification in complex carbohydratedegrading human gut bacteria. mSystems 7:e0094721. https://doi.org/ 10.1128/msystems.00947-21
- Godoy VG, Dallas MM, Russo TA, Malamy MH. 1993. A role for *Bacteroides fragilis* neuraminidase in bacterial growth in two model systems. Infect Immun 61:4415–4426. https://doi.org/10.1128/iai.61.10.4415-4426.1993
- Buzun E, Hsu C-Y, Sejane K, Oles RE, Ayala AV, Loomis LR, Zhao J, Rossitto L-A, McGrosso D, Gonzalez DJ, Bode L, Chu H. 2024. A bacterial sialidase mediates early life colonization by a pioneering gut commensal. Cell Host Microbe:2023.08.08.552477. https://doi.org/10.1016/j.chom.2023. 12.014
- Ghosh S. 2020. Sialic acid and biology of life: an introduction. Sialic Acids Sialoglycoconjugates Biol Life Health Dis:1–61. https://doi.org/10.1016/ B978-0-12-816126-5.00001-9
- Coyne MJ, Roelofs KG, Comstock LE. 2016. Type VI secretion systems of human gut *Bacteroidales segregate* into three genetic architectures, two of which are contained on mobile genetic elements. BMC Genomics 17:58. https://doi.org/10.1186/s12864-016-2377-z
- Sheahan ML, Coyne MJ, Flores K, Garcia-Bayona L, Chatzidaki-Livanis M, Sundararajan A, Holst AQ, Barquera B, Comstock LE. 2023. A ubiquitous mobile genetic element disarms a bacterial antagonist of the gut microbiota. bioRxiv. https://doi.org/10.1101/2023.08.25.553775
- Chatzidaki-Livanis M, Geva-Zatorsky N, Comstock LE. 2016. Bacteroides fragilis type VI secretion systems use novel effector and immunity proteins to antagonize human gut Bacteroidales species. Proc Natl Acad Sci U S A 113:3627–3632. https://doi.org/10.1073/pnas.1522510113
- Hecht AL, Casterline BW, Earley ZM, Goo YA, Goodlett DR, Bubeck Wardenburg J. 2016. Strain competition restricts colonization of an enteric pathogen and prevents colitis. EMBO Rep 17:1281–1291. https:// doi.org/10.15252/embr.201642282
- Rana S, Skariyachan S, Uttarkar A, Niranjan V. 2023. Carboxymuconolactone decarboxylase is a prospective molecular target for multi-drug resistant *Acinetobacter baumannii*-computational modeling, molecular docking and dynamic simulation studies. Comput Biol Med 157:106793. https://doi.org/10.1016/j.compbiomed.2023.106793
- Zieliński M, Park J, Sleno B, Berghuis AM. 2021. Structural and functional insights into esterase-mediated macrolide resistance. Nat Commun 12:1732. https://doi.org/10.1038/s41467-021-22016-3