Growth rate alterations of human colorectal cancer cells by 157 gut bacteria

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ABSTRACT
Several bacteria in the human gut microbiome have been associated with colorectal cancer (CRC) by high-throughput screens. In some cases, molecular mechanisms have been elucidated that drive tumorigenesis, including bacterial membrane proteins or secreted molecules that interact with the human cancer cells. For most gut bacteria, however, it remains unknown if they enhance or inhibit cancer cell growth. Here, we screened bacteria-free supernatants (secretomes) and inactivated cells of over 150 cultured bacterial strains for their effects on cell growth. We observed family-level and strain-level effects that often differed between bacterial cells and secretomes, suggesting that different molecular mechanisms are at play. Secretomes of Bacteroidaceae, Enterobacteriaceae, and Erysipelotrichaceae bacteria enhanced cell growth, while most Fusobacteriaceae cells and secretomes inhibited growth, contrasting prior findings. In some bacteria, the presence of specific functional genes was associated with cell growth rates, including the virulence genes TcdA, TcdB in Clostridiales and FadA in Fusobacteriaceae, which both inhibited growth. Bacteroidaceae cells that enhanced growth were enriched for genes of the cobalamin synthesis pathway, while Fusobacteriaceae cells that inhibit growth were enriched for genes of the ethanolamine utilization pathway. Together, our results reveal how different gut bacteria have wide-ranging effects on cell growth, contribute a better understanding of the effects of the gut microbiome on host cells, and provide a valuable resource for identifying candidate target genes for potential microbiome-based diagnostics and treatment strategies.

Introduction

Over the past few decades, thousands of microbial species have been identified in the human gut microbiome. Each person harbors an estimated 3.8 × 10^13 bacteria in their guts with a relatively stable species composition.1-3 Changes in the composition of the gut microbiome have been linked with various diseases including colorectal cancer (CRC), the third leading cancer worldwide with an incidence that is estimated to rise in the coming decennium.4,5 Besides important environmental and genetic factors, the initiation and development of CRC are affected by the human gut microbiome. During tumorigenesis, the bacterial composition on the affected intestinal mucosa changes in a process that is described by the bacterial driver-passenger model.6,7 Bacterial drivers may facilitate the acquisition of hallmarks of cancer8 in mucosal cells by generating DNA-damage, reducing the epithelial barrier function, and stimulating procarcinogenic immune responses. As the tumor microenvironment changes in the transition from adenoma to carcinoma, bacterial passengers compete with the drivers, further shifting the microbiome toward a CRC signature.9,10 The newly acquired bacteria may in turn affect tumor development, by potential inhibiting, enhancing, or neutral effects on tumor cell growth.6 These CRC-specific changes offer the opportunity to use the bacterial community for diagnostic, prognostic, preventive, and treatment measures.11-20

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In feces of CRC patients, the genera *Fusobacterium*, *Peptostreptococcus*, *Ruminococcus*, and *Escherichia/Shigella* are enriched, while *Bacteroides* and *Lachnospiraceae* (*Roseburia*, *Lachnospira*, *Anaerostipes*) have a lower relative abundance. Even though phyla such as *Bacteroidetes* are depleted, single species like enterotoxigenic *Bacteroides fragilis* may be enriched in some CRC patients. Comparing tumor tissue with adjacent normal tissue on the intestinal mucosa of CRC patients revealed that *Fusobacterium*, *Prevotella*, *Streptococcus*, and *Peptostreptococcus* were enriched in tumors while *Blautia*, *Escherichia*, *Pseudomonas*, and *Faecalibacterium* were enriched in the normal mucosa.

Many different bacteria have been associated with CRC tumors, but we are only beginning to understand the different mechanisms that are involved. The effect of bacteria on cell growth may be driven by direct bacterial-to-epithelial cell-cell contact or by secreted products present in the secretome. Membrane-bound bacterial proteins that require cell-cell contact can activate epithelial cell signaling. For example, the passenger bacterium *Fusobacterium nucleatum* encodes the membrane protein *Fusobacterium* adhesin A (FadA) that binds to E-cadherin, activating β-catenin signaling and resulting in increased tumor growth. Specific *E. coli* species with the *eae* gene express the adhesin protein intimin on their membrane surface which binds to and causes lesions to gut epithelial cells, allowing bacteria to breach the colonic barrier. Once the bacteria are bound to the epithelial cells, this allows them to inhibit DNA repair proteins, further contributing to lasting DNA damage. Several secreted bacterial toxins are known that can bind to receptors or pass through the cell membrane into the cytoplasm. The driver bacterium enterotoxigenic *B. fragilis* secretes the metalloprotease *B. fragilis* toxin (BFT) which leads to E-cadherin cleavage and increased wnt-signaling in colon epithelial cells and to tumor formation in mice and increased cell proliferation in vivo.

Bacteria such as *Escherichia coli* containing the *pks* island are able to produce a genotoxin called colibactin. Upon mucosal breach, colibactin reaches the epithelial cells and alkylates DNA, ultimately leading to tumorigenesis. These examples show that taxonomically diverse bacteria may lead to the acquisition of hallmark capabilities via diverse mechanisms.

The aim of this study was to examine the effects of bacterial cells and their secretomes on the growth rates of epithelial cells. We tested the effect of 157 different gut bacteria on the growth rates of five CRC cell lines and one immortalized kidney cell line. Our results revealed that different bacterial families specifically inhibit or enhance cell growth, although contrasting effects could be observed between some closely related strains. Both known virulence genes and novel microbial pathways were associated with the different growth rate changes. These results provide the first large-scale in vitro analysis of the effects of different microbial strains on epithelial cell growth.

Materials and methods

Bacterial strains

We selected 116 different gut bacteria, including species that are depleted or enriched in CRC patients whose genome sequences were available in the PATRIC database. Additionally, we selected specific bacteria without sequenced genomes (*n* = 39) that were strongly linked to CRC or were isolated from CRC tissue, including *Streptococcus bovis*, *Clostridium septicum*, *Clostridoides difficile*, *Bacteroides* sp. and the two potentially beneficial bacteria *Lactobacillus casei Shirotia* and *Akckermansia muciniphila* ATCC BAA-835™. Together, we analyzed 157 bacterial strains isolated from the human gut microbiome (Supplementary Table S1).

We purchased 96 bacterial strains from the reference catalog of the Human Microbiome Project (HMP, Prof. Dr. Emma Allen-Vercoe from the University of Guelph, Canada); 24 bacteria were kindly provided by Prof. Dr. Cynthia L. Sears from Johns Hopkins Medical Institutions, Baltimore, MD, USA; five strains were purchased from DSMZ (*Clostridium septicum* (Macé 1889) Ford 1927 DSMZ7534, *C. difficile* (Hall and O’Toole 1935) Lawson et al. 2016 DSM27543 (known as *Clostridium difficile* 630), *Fusobacterium nucleatum* Knorr 1922 DSM15643, *Fusobacterium nucleatum* subsp. *poly- morphum* (ex Knorr 1922) Dzink et al. 1990 DSM20482, and *Peptostreptococcus stomatis* Downes and Wade 2006 DSM17678); one strain
from ATCC (Streptococcus agalactiae ATCC13813); and 31 bacteria were in stock at the Radboud University Medical Center in Nijmegen, The Netherlands.\textsuperscript{58-60} Information about the bacterial strains, their origin, growth media, and their genome sequence is provided in Supplementary Table S1.

Bacteria were cultured in their respective media under anaerobic conditions at 37°C aired with nitrogen gas (N\textsubscript{2}, see Supplementary Table S1). Alternatively, Ralstonia sp. 5_2_56FAA, Ralstonia sp. 5_7_47FAA and Pseudomonas sp. 2_1_26 were grown aerobically at 37°C, and Bacillus smithii was grown anaerobically at its preferred temperature of 50°C. Eight media were prepared to culture bacteria (Supplementary Table S1): (i) brain heart infusion-supplemented (BHI-S, ATCC Medium 1293), (ii) BHI-S supplemented with 0.1% Tween 80 at pH 8.0 (BHI-T),\textsuperscript{61} (iii) NADC – 99X medium (ATCC Medium 1804), (iv) Desulfovibrio medium (ATCC Medium 2755), (v) Modified Reinforced Clostridial Medium (RCM, ATCC Medium 2107), (vi) Reinforced Clostridial medium with sodium lactate (60% solution) at a concentration of 1.5% (RCM+, from ATCC Medium 1252), (vii) differential RCM (DRCM), and (viii) nutrient broth (NB, ATCC Medium 3). Bacteria were grown until the medium was turbid and the optical density (OD) was at least 1.0. Absorbance was obtained by measuring OD at 600 nm.

Obtaining secretomes and bacterial cells

To investigate the effect of secreted molecules, we obtained secretomes from 154 bacterial cultures as follows. After culturing, bacteria were centrifuged at 4,700 rpm for 10 min and some bacteria subsequently at a higher speed to settle them down (see Supplementary Table S1). Next, supernatants were spun at 4,700 rpm for 10 min to pellet any remaining bacteria and filter-sterilized using 0.2 μm filters (Sigma-Aldrich, USA). Molecules larger than 10 kDa were concentrated using amicon ultra-15 centrifugal filters (Merck Millipore, Merck, USA). Concentrated supernatants were frozen at –80°C until further use and are further referred to as secretomes. To investigate the effect of bacterial surface-bound molecules, 145 bacteria were inactivated to allow cellular growth rates to be assessed by measuring metabolic activity (see the section “MTT assay” below). Cells of 12 spore-forming bacteria under our experimental conditions were excluded from this analysis (see Supplementary Table S1). Pelleted bacteria were resuspended in 70% ethanol and incubated for 5 min according to our inactivation protocol.\textsuperscript{62} After centrifugation at 20,000 rcf for 30 s, bacteria were washed in PBS twice and inactivated bacteria were frozen at –80°C until further use.

Culturing of cell lines and identification of their mutational landscapes

Five CRC cell lines (Caco-2, HCT15, HCT116, HT29, and SW480) and one immortalized embryonic kidney cell line (HEK293T) were selected based on their differences in mutational landscapes.\textsuperscript{63} Most commonly mutated oncogenes and tumor suppressor genes in CRC were confirmed with targeted single molecule molecular inversion probe (smMIP) mutation analysis as described,\textsuperscript{64} with additional smMIPs targeting CXCR4 (NM_001108540.1) codons 281–357, EZH2 (NM_0004456.4) codons 471–502, 618–645, 679–704, and SF3B1 (NM_012433.2) codons 603–471, 833–906 (smMIP sequences are available upon request), at the Radboud Diagnostics Department, Nijmegen, The Netherlands (Supplementary Table S2). Cells were split twice a week using Dulbecco’s Modified Eagle Medium (DMEM, Lonza, Switzerland) supplemented with 10% fetal bovine serum (FBS, Thermo Fisher Scientific, USA) and 1% Penicillin/Streptomycin (Pen/Strep, Thermo Fisher Scientific, USA). Cells used for this study were not split for more than 25 passages.

For MTT assays, optimal cell seeding in 96 well plates (Greiner Bio-One, Austria) was defined at 5,000 cells/well for Caco-2, 4,000 cells/well for HCT15, 1,500 cells/well for HCT116, 10,000 cells/well for HT29, 6,000 cells/well for SW480, and 20,000 cells/well for HEK293T. Cells were incubated overnight to attach before the addition of secretomes or bacterial cells.
**MTT assay**

Secretomes were diluted to their original concentrations in DMEM supplemented with 10% FBS and 1% Pen/Strep. Inactivated bacteria were diluted with DMEM supplemented with 10% FBS and 1% Pen/Strep to an OD of 0.1 and incubated on cell lines up to 72 h. For secretomes, filter-sterilized bacterial culture media used to generate secretomes served as a control for cell growth in the assay, while for bacterial cells, DMEM supplemented with 10% FBS and 1% Pen/Strep served as control for cell growth. MTT assays were performed to measure the cell metabolic activity every 24 h. These assays involve the conversion of the watersoluble yellow dye MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] to the insoluble purple formazan by the action of mitochondrial reductase. At 24, 48 and 72 h 10 μl of MTT (5 mg MTT dissolved per mL PBS) dye was added to the wells and cells were incubated for 3 h. Formazan was then solubilized with 150 μL MTT solvent (0.5 mL of 10% Nonidet (dissolved in water), 50 mL isopropanol (2-propanol) and 16.7 μL hydrochloric acid fuming 37% together) and the concentration determined by optical density at 570 nm (BioRad microplate reader model 680). Metabolic activity of cell lines was used as a measure for cell growth. All experiments were performed in quadruplicate.

**Cell growth analysis**

Absorbances at an optical density of 570 nm were measured by MTT assays at four time points (0, 24, 48, and 72 h). Growth rates per hour were calculated by dividing the difference in absorbance by the time interval between measurements (24 h). Results from four independent experimental replicates were averaged. Growth rates between 24 and 48 h were selected for comparative analysis because that time window captured the optimal growth for most cell lines (Supplementary Figure S1).

A growth rate score was defined by subtracting the growth of negative controls (cells cultured without bacterial products) from those cultured with bacterial products (secretomes or bacterial cells). Scores greater than 0 refer to enhanced growth, while scores lower than 0 refer to inhibited growth (see Supplementary Tables S3 and S4). To allow comparability between cell lines, normalized growth rate scores were calculated by dividing per cell line the scores (g) of each of the n bacteria by the norm (|g|) of the experimental condition $|g| = \sqrt{\sum_{i=1}^{n} g_i^2}$. The normalized growth rate scores were used to evaluate whether the enhancing or inhibiting effect of a bacteria is significantly stronger than the mean effect observed for all other bacteria. For this purpose, growth rate scores were transformed into z-scores and p-values were computed by separate one-tailed z-tests for enhancing or inhibiting effects. Based on these p-values, strains were identified as enhancers or inhibitors of growth by using a cutoff of $p < 0.05$. This cutoff represents the 5% extremes of the distribution of z-scores and were here used to identify the strongest effects. The FDR corrected p-values were also reported (Supplementary Tables S3 and S4).

The overall effects of bacterial secretomes and inactivated cells were visualized by projecting their normalized growth scores in a two-dimensional space with the t-SNE algorithm. To evaluate if bacteria of the same family have a similar overall effect on cell growth relative to bacteria from other families, we applied unsupervised nearest neighbor clustering and evaluated how often bacteria from the same family were nearest neighbors compared to a random sample without replacement. For this analysis, we only used bacteria from families with more than four representatives. Random groups were evaluated per family and were defined to contain the same number of strains as the family. Fisher’s exact test was used to evaluate if the nearest neighbor groups were enriched for bacteria of the same family compared to 1,000 random groups of the same size. The harmonic means of p-values from this comparison are reported.

**Association of bacterial virulence genes to growth rate changes**

Bacterial genomes were screened for 35 known virulence genes that may be associated with cancer or epithelial cell changes (Supplementary Table S5). The virulence factors Map (Mitochondrial associated protein), Type-3 secretion system (T3SS)
effect protein of *Citrobacter rodentium*, Shiga toxin 2A (Stx2A), Shigella enterotoxin 1 (ShET1), and colibactin (cyclomodulin toxin on polyketide synthase (pks) island of *Enterobacteriacea*) were identified in the *Enterobacteriacea* family, *B. fragilis* toxin (BFT) in *B. fragilis* species, and FadA in the *Fusobacteriacea* family (Supplementary Table S5). *Clostridium difficile* toxin A, and B (TcdA/TcdB) in *Clostridioides difficile* (Prévot 1938) Lawson et al. 2016, and Alpha-toxin of *Clostridium septicum* (Mace 1889). 1927 were identified as secreted factors in previous reports. Growth rate changes of strains containing virulence genes (if present in ≥2 strains) were compared to those of their non-virulent counterparts within the order (*Clostridiales*), family (*Fusobacteriacea* and *Enterobacteriacea*) or species (*B. fragilis*). Groups were compared using Mann-Whitney U-test of aggregate data of all cell lines. *P* < .05 was considered significant.

**Genes and subsystems identification**

To identify genes that are associated with the inhibiting/enhancing effects of bacterial strains within families, we sorted all strains based on their effect on growth rate scores and used a normalized Kolmogorov–Smirnov statistic to quantify the association of a gene with the observed effect. The significance of the rank-based statistic was estimated by comparing its value to values obtained from 10^3 random permutations of the same list. This provided all genes encoded by the genomes from a family with a *p*-value which quantified the association of the gene to the inhibiting/enhancing effect. We evaluated if genes that belong to the same functional subsystems were more often found to be significantly associated with the inhibiting/enhancing effect than to 10^3 random groups of genes. We defined this association by modeling the average count of genes belonging to a functional subsystem within the random groups as Poisson distributed variables. We then tested if the count of genes with significant *p*-values was statistically higher than expected from the Poisson model, with a significance level of 0.05 after Benjamini/Hochberg correction for multiple tests. We used the Poisson distribution as the null distribution since (i) the labels for functional subsystem within the random groups are independent; (ii) Counts are expressed as averages from many random samples; (iii) the mean and variance are equivalent, all features consistent with a Poisson distribution.

**Results**

**Reproducibility of growth rate alterations by bacterial cells and secretomes**

To identify alterations in the growth rates of CRC cell lines by bacterial cells or their secreted products, inactivated cells or secretomes were added to five CRC cell lines with different mutational landscapes and one embryonic kidney cell line HEK293T (see Methods and Supplementary Table S2). HEK293T cells are immortalized by DNA of human adenovirus type 5 resulting in senescence of the pRb and p53 pathway. While these cells are not normal, they have no mutations in the genes that are commonly mutated in CRC, as confirmed by our smMIP analysis (Supplemental Table 2). For each cell line independently, plain cell culture and bacterial culture media served as a control for cell growth for cells and secretomes, respectively. A total of 7,176 experiments were conducted where we measured the bacterial effect on cell growth. Each experiment was repeated four times. We evaluated the reproducibility of these replicates by performing linear regression over all pairwise combinations of replicates, i.e. measurements for all the possible combinations of two replicates for every strain appended into two separate lists. As shown in Supplementary Figure S2, we found high reproducibility between replicates (regression coefficient: 0.87 ± 0.05, intercept: 0.04 ± 0.02). Based on this reproducibility, we averaged the four experimental replicates for further analysis.

**Contrasting effects between bacterial cells and secretomes**

Figure 1(a) summarizes the effect of bacterial cells and secretomes on the growth of human cell lines. Overall, bacterial cells exhibit a stronger inhibiting effect than secretomes (*p* = 1.02e-75, Wilcoxon signed rank test, Figure 1(b)). There was a low correlation between the overall effects of
Figure 1. Growth rate scores of six human cell lines upon treatment with bacterial cells and secretomes. (a) Heatmap indicating low and high growth rate scores, respectively red and blue. Bacteria are sorted within bacterial families by the average growth rate score. Red numbered octagons highlight the strains discussed in the text: Bacteroides sp. 2_1_22 (1), B. fragilis K570 clinda R (ETBF) (2), B. sp. 4_1_36 (3), Clostridium septicum (Mace 1889) Ford 1927 (4), C. sp. D5 (5), Escherichia coli D9 (6), Klebsiella sp. 1_1_55 (7), E. coli 4_1_47FAA (8), F. nucleatum DSM 15643 (ATCC 25586) (9), F. nucleatum DSM 20482 (ATCC 10953) (10), F. necrophorum subsp. funduliforme 1_1_36S (11), F. nucleatum subsp. animalis 11_3_2 (12), Lachnospiraceae bacterium 8_1_57FAA (13), L. bacterium 3_1_46FAA (14), Streptococcus bovis 1212 (15), S. bovis 1459 (16), S. bovis 1417 (17), S. bovis 207 (18), Pediococcus acidilactici 7_4 (19), Pseudomonas sp. 2_1_26 (20), D. sp. 6_1_46AFAA (21), Ralstonia sp. 5_2_56FAA (22), Ruminococcaceae bacterium D16 (23),
Secretomes and inactivated bacteria on cell growth (Pearson r = 0.176). A low correlation was also observed when the measurements were stratified per bacterial family (Supplementary Table S6), indicating that specific factors within the two compartments (cell wall attached or secreted molecules) are of importance for the observed effects.

**Difference in growth rate alterations between CRC cell lines**

Growth rates were significantly enhanced or inhibited (p < .05) in one or more cell lines in 35 out of 145 (24.1%) inactivated bacterial strains (Figure 2(a), Supplementary Table S3) and 33 out of 154 (21.4%) secretomes (Figure 2(b), Supplementary...
Table S4). Growth rates of HEK293T cells were affected by the lowest number of bacteria (n = 21, n = 3 enhanced and n = 18 inhibited growth). On the other end of the spectrum were HCT116 cells that were affected by 28 different bacteria (n = 11 enhanced and n = 17 inhibited growth) and HCT15 cells that were affected by 27 different bacteria (n = 8 enhanced and n = 19 inhibited; Figure 2(c)). These two cell lines had microsatellite instability (MSI) and harbored the greatest number of pathogenic mutations in known tumor suppressor and oncogenes (n = 3 and n = 4, respectively, see Supplementary Table S2). In general, the number of pathogenic mutations of a cell line correlated with the number of bacteria that significantly affected its growth (Pearson r = 0.931, p < .01, see Figure 2(d)), suggesting that acquiring hallmark mutations might make cells more susceptible to growth rate changes by bacteria.

Some strains significantly enhanced or inhibited the growth rate of specific cell lines (Figure 1). Streptococcaceae cells generally had a neutral effect on cell growth, however, some Streptococcaceae secretomes specifically enhanced the growth of HT29 and SW480 cells (Figure 1). Three Streptococcus bovis secretomes (strains 207, 1212 and 1417) and one inactivated bacterium (strain 1459) selectively enhanced growth in HT29 cells, while no effects were observed in other cell lines (Figures 1 and 2(c)). The genomes of these enhancing Streptococcus bovis strains cluster together in a phylogenetic tree with Streptococcus galloplusus subsp. galloplusus (SGG) and subsp. pasteurianus (SGP) (Supplementary Figure 3). Other notable cell-line specific effects include Erysipelotrichaceae cells, and Tannerellaceae secretomes that significantly enhance the growth of HCT15 and HCT116 cells, respectively.

**Bacterial family-level and strain-level effects on cell growth rates**

To assess the overall effect of the different bacterial secretomes and cells on our six cell lines, we projected the growth rate scores into a two-dimensional t-SNE plot. Figure 3 shows this lower dimensional space overlayed with family-specific colors and a shade corresponding to the average growth effect. We observed a clear clustering of most, but not all of the bacterial families, i.e. cells or secretomes from the same families tend to have similar effects on the different cell lines (see also

![Figure 3. T-SNE plots summarizing the overall growth effects of bacterial cells (a) and secretomes (b) on six human cell lines. Colored circles correspond to strains belonging to particular bacterial families. Magenta and light blue shading around the circles indicate enhancing and inhibiting effects of strains, respectively. Families labeled as “Others” contains the following strains for bacterial cells (A): Lactobacillaceae (1), Peptostreptococccaece (2), Desulfovibrionaceae (3), Eubacteriaceae (4), Bifidobacteriaceae (5), Enterococcaceae (6), Veillonellaceae (7), Synergistaceae (8), Burkholderiaceae (9), Pseudomonadaceae (10), Propionibacteriaceae (11), Ruminococcaceae (12), Akkermansiaceae (13), Acidaminococcaceae (14). And the following strains for secretomes (B): Lactobacillaceae (1), Eubacteriaceae (2), Clostridiaceae (3), Peptostreptococcaceae (4), Bifidobacteriaceae (5), Enterococcaceae (6), Veillonellaceae (7), Synergistaceae (8), Burkholderiaceae (9), Desulfovibrionaceae (10), Propionibacteriaceae (11), Ruminococcaceae (12), Pseudomonadaceae (13), Akkermansiaceae (14), Acidaminococcaceae (15), Bacillaceae (16).]
Table 1. Secretomes and bacteria significantly enhancing or reducing cell growth rate (z-scores).

| Family                  | Bacteria                              | Caco-2 | HT29  | HCT15 | SW480 | HCT116 | HEK293T |
|-------------------------|---------------------------------------|--------|-------|-------|-------|--------|---------|
| **Bacteroidaceae**      | B. dorei 5_1_36/D4                    | 0.89   | 0.82  | 1.36  | 0.90  | 1.87   | 1.38    |
|                         | B. eggerthii 1_2_48FBA                | 1.82   | 0.59  | 1.51  | 1.16  | 1.21   | 1.48    |
|                         | B. fragilis KS70 clinda R (ETBF)      | 1.74   | 1.71  | 1.77  | 0.71  | 1.15   | 0.36    |
|                         | B. massiliensis 3972 T(B)2            | -0.44  | -1.63 | -0.83 | -0.51 | -1.45  | -2.01   |
|                         | B. ovatus 3_8_47FBA                   | 1.96   | 0.17  | 1.17  | 1.27  | 1.24   | 1.42    |
| **Bacteroides sp. 1_1_6** | 2.36   | -0.21 | 0.63  | 0.86  | 1.27  | 0.93    |
| **Bacteroides sp. 2_1_16** | 1.15   | 0.51  | 0.98  | 0.53  | 1.87  | 1.17    |
| **Bacteroides sp. 2_1_22** | -1.83  | -0.73 | -0.05 | -0.49 | 0.82  | -0.08   |
| **Bacteroides sp. 2_1_22** | -2.49  | -2.26 | -2.31 | -1.64 | -2.42 | -1.78   |
| **Bacteroides sp. 4_1_36** | 2.39   | 0.67  | 1.34  | 1.58  | 1.50  | 1.93    |
| **Bacteroides sp. 9_1_42FBA** | 1.71   | -0.06 | 0.71  | 1.09  | 1.46  | 0.87    |
| **Bacteroides sp. D2**   | -1.82  | -0.64 | -0.04 | -0.17 | 0.45  | 0.04    |
| **Clostridiaceae**      | C. septicum (Mace 1889) Ford 1927    | -4.64  | -5.02 | -4.11 | -2.63 | -1.84  | -4.49   |
| **Enterobacteriaceae**  | E. coli 4_1_47FBA                     | 1.63   | 1.05  | 1.26  | 1.17  | 2.12   | 1.65    |
|                         | E. coli D9                            | 0.48   | 0.64  | 1.12  | 0.88  | 1.75   | 1.28    |
|                         | Klebsiella sp. 1_1_55                 | 0.77   | 1.15  | 1.19  | 0.57  | 1.92   | 0.70    |
| **Erysipelotrichaceae** | Coprobacillus sp. 29_1                | -0.06  | 1.32  | 1.94  | 1.03  | 0.81   | 0.91    |
|                         | Coprobacillus sp. 29_1                | 0.40   | 1.84  | 1.64  | 1.15  | 0.72   | 0.21    |
|                         | Coprobacillus sp. 8_2_54FBA          | 0.68   | 1.78  | 1.59  | 1.15  | 0.82   | 0.46    |
|                         | Coprobacillus sp. D7                 | 1.02   | 1.64  | 1.74  | 1.18  | 0.80   | 0.58    |
|                         | Erysipelotrichaceae bacterium 2_2_44A | 1.07   | 1.26  | 1.70  | 0.95  | 1.82   | 0.68    |
|                         | Erysipelotrichaceae bacterium 21_3    | 1.19   | 1.51  | 1.85  | 1.13  | 1.47   | 1.10    |
| **Fusobacteriaceae**    | F. gordoniiformans 3-1-5R             | -0.43  | -1.52 | -2.62 | -3.32 | -2.16  | -1.16   |
|                         | F. necrophorum subsp. funditiforme 1_1_36S | 1.18  | 1.98  | 1.12  | 1.71  | 2.63   | 1.72    |
|                         | F. nucleatum subsp. animalis 11_3_2   | -2.50  | -2.86 | -3.56 | -3.09 | -2.70  | -0.77   |
|                         | F. nucleatum subsp. animalis 11_3_2   | 2.15   | 0.48  | 1.11  | 0.96  | 2.09   | 1.20    |
|                         | F. nucleatum subsp. animalis 21_1A    | -1.45  | -1.74 | -2.31 | -1.33 | -1.47  | -1.03   |
|                         | F. nucleatum subsp. animalis 3_1_33   | -0.66  | -1.74 | -2.07 | -1.24 | -1.60  | -1.25   |
|                         | F. nucleatum subsp. animalis 7_1      | 0.14   | -0.72 | -1.49 | -1.83 | -1.81  | 0.10    |
|                         | F. nucleatum subsp. animalis 7_1      | -1.86  | -1.40 | -2.48 | -1.73 | -1.30  | -1.75   |
|                         | F. nucleatum subsp. nucleatum ATCC 25586 | -1.64  | -1.67 | -1.67 | -1.89 | -1.67  | -2.04   |
|                         | F. nucleatum subsp. polymorphum ATCC 10953 | -1.51  | -1.39 | -1.33 | -1.94 | -1.60  | -1.88   |
|                         | F. nucleatum subsp. vincentii 3_1_27  | -0.48  | -1.88 | -2.13 | -1.65 | -1.56  | -1.20   |
|                         | F. nucleatum subsp. vincentii 3_1_36A2 | -0.95  | -1.62 | -2.24 | -1.72 | -1.17  | -1.26   |
|                         | F. nucleatum subsp. vincentii 4_1_13  | -0.97  | -1.92 | -2.06 | -1.52 | -1.76  | -1.45   |
|                         | F. periodonticum 1_1_41FBA            | -0.83  | -1.94 | -2.04 | -1.27 | -1.69  | -1.48   |
|                         | F. periodonticum 2_1_31               | -3.10  | -3.10 | -3.38 | -2.64 | -2.44  | -2.35   |
|                         | F. periodonticum 2_1_31               | -1.08  | -1.32 | -1.66 | -0.86 | -0.97  | -1.90   |
| **Lachnospiraceae**     | Anaerostipes sp. 3_2_56FBA            | -0.51  | -1.67 | -1.39 | -2.08 | -1.54  | -1.47   |
|                         | C. symbiosum WAL-14163                | -1.01  | -1.27 | -1.75 | -1.82 | -0.70  | -2.33   |

(Continued)
To confirm this observation, we tested if the growth effects of bacteria from the same family were more similar than one would expect by chance by permuting the data labels 1,000 times (see Methods). The effect of bacterial cells and secretomes was found to be significantly clustered for five and three bacterial families, respectively (Supplementary Table S7). For bacterial cells, the most striking effect was observed for *Fusobacteriaceae* that generally inhibited cell growth. Most *Bacteroidiaceae*, *Enterobacteriaceae*, and *Erysipelotrichiaceae* secretomes enhanced cell growth, although not always significantly (Figure 1). Contr astingly, *Clostridiaceae* secretomes generally inhibited growth rates.

In the *Bacteroidiaceae* family (n = 41 strains tested), secretomes and bacterial cells of seven and one strains significantly enhanced cell growth, respectively. The effects were most prominently observed in Caco-2 cells, while the growth of HEK293T cells was only significantly enhanced by the secretome of *Bacteroides* sp. 4_1_36 (Figure 2(c) and Table 1). Only three *Enterobacteriaceae* secretomes (n = 15) significantly enhanced the growth of HEK293T cells (*Escherichia coli* 4_1_47FAA) and HCT116 (*Escherichia coli* 4_1_163FAA).}

| Family                  | Species/Strain               | Effect on Caco-2 | Effect on HEK293T |
|-------------------------|-------------------------------|-----------------|------------------|
| *Lachnospiraceae*       | *bacterium 1_1_57FAA*        | -1.80 -1.20 -1.56 -2.32 -1.14 -1.14 |                |
| *Lachnospiraceae*       | *bacterium 3_1_46FAA*        | -2.41 -1.37 -2.59 -2.30 -1.84 -3.22 |                |
| *Lachnospiraceae*       | *bacterium 4_1_37FAA*        | 1.98 0.93 1.19 1.46 1.80 1.23 |                |
| *Lachnospiraceae*       | *bacterium 5_1_63FAA*        | 0.50 1.68 1.81 1.04 0.09 1.04 |                |
| *Lachnospiraceae*       | *bacterium 8_1_57FAA*        | -2.70 -1.52 -2.70 -2.30 -1.95 -3.24 |                |
| *Tannerellaceae*        | Parabacteroides sp. 2_1_7    | 0.06 0.17 0.56 0.88 -1.89 0.73 |                |
| *Tannerellaceae*        | Parabacteroides sp. 2_1_7    | 1.73 -0.64 0.92 -0.04 0.32 0.41 |                |
| *Tannerellaceae*        | Parabacteroides sp. D13      | 0.86 0.56 1.04 1.22 -2.21 1.10 |                |
| *Streptococcaceae*      | *S. bovis 1212*              | 0.23 1.64 0.40 0.62 -0.66 -0.41 |                |
| *Streptococcaceae*      | *S. bovis 1417*              | 0.49 1.65 0.39 0.67 -0.51 -0.02 |                |
| *Streptococcaceae*      | *S. bovis 207*               | 0.63 2.10 0.84 1.13 0.94 0.53 |                |
| *Streptococcaceae*      | *S. bovis 207*               | 0.35 2.29 0.45 0.81 -0.59 -0.42 |                |
| *Streptococcaceae*      | *S. salivarius NTB2*         | -0.98 -0.89 -1.73 -1.98 -1.52 -0.86 |                |
| *Streptococcaceae*      | *S. sp. 2_1_36FAA*           | -1.57 -1.84 -0.52 -0.51 -1.73 -1.54 |                |
| *Other*                 | *C. difficile* (Prévet 1938) | 0.13 -1.75 -1.26 -2.10 -1.22 -2.04 |                |
| *Others*                | *Desulfovibrio sp. 3_1_syn3* | 2.20 0.17 0.41 0.99 -0.37 -0.17 |                |
| *Others*                | *Desulfovibrio sp. 3_1_syn3* | -0.77 1.76 -0.40 -1.47 -1.77 -2.52 |                |
| *Others*                | *Desulfovibrio sp. 6_1_46FAA* | 0.90 0.06 -0.87 -0.55 -1.75 -1.35 |                |
| *Others*                | *Desulfovibrio sp. 6_1_46FAA* | -0.78 1.17 -0.69 -1.93 -2.06 -3.17 |                |
| *Others*                | *Eubacterium sp. 3_1_31*     | 1.25 1.02 1.24 1.85 0.96 1.43 |                |
| *Others*                | *P. acidilactici* 7_4        | -1.58 -0.82 0.71 0.33 -1.39 -1.96 |                |
| *Others*                | *Propionibacterium sp. 5_U_42FAA* | 0.20 0.20 2.02 0.96 -0.31 -1.33 |                |
| *Others*                | *Pseudomonas sp. 2_1_26*     | -4.60 -5.41 -4.35 -4.20 -2.87 -4.21 |                |
| *Others*                | *Pseudomonas sp. 2_1_26*     | -0.02 -1.26 -0.72 -1.91 -1.45 0.35 |                |
| *Others*                | *Ralestonia sp. 5_2_56FAA*   | -1.12 -0.28 0.99 -2.05 -0.36 -2.18 |                |
| *Others*                | *Ruminococcaceae bacterium D16* | -1.14 -1.53 -1.34 -1.23 -1.22 -2.50 |                |
| *Others*                | *Synergistes sp. 3_1_syn1*   | -1.40 -1.60 -1.57 -1.34 -1.68 -2.09 |                |

Secretomes are highlighted in gray, green represents significant enhancement ($p < 0.05$) in cell growth rate and red represents significant reduction ($p < 0.05$) in cell growth rate.

Figure 1 and Table 1. To confirm this observation, we tested if the growth effects of bacteria from the same family were more similar than one would expect by chance by permuting the data labels 1,000 times (see Methods). The effect of bacterial cells and secretomes was found to be significantly clustered for five and three bacterial families, respectively (Supplementary Table S7). For bacterial cells, the most striking effect was observed for *Fusobacteriaceae* that generally inhibited cell growth. Most *Bacteroidiaceae*, *Enterobacteriaceae*, and *Erysipelotrichiaceae* secretomes enhanced cell growth, although not always significantly (Figure 1). Contr astingly, *Clostridiaceae* secretomes generally inhibited growth rates.

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From the 12 out of 16 significant *Fusobacteriaceae* (81.3%) cells, 2 enhanced and 10 inhibited cellular growth. *Fusobacterium necrophorum* subsp. *funduliforme* 1_1_36S cells enhanced growth in HCT116, SW480, and HEK293T cells. Strikingly, while *Fusobacterium nucleatum* subsp. *animalis* 11_3_2 cells enhanced the growth of Caco-2 and HCT116, its secretome inhibited growth in those same cell lines. Especially the cell line HCT15 was sensitive to *Fusobacteriaceae* growth inhibition where 11 out of 16 strains significantly reduced growth rates. Alternatively, HEK293T and Caco-2 were less sensitive to *Fusobacteriaceae* with only four out of 16 strains inhibiting cell growth. Notably, the effect of *Fusobacteriaceae* secretomes on cell growth may be cell-specific depending on, for example, mutation status, since most secretomes that inhibited HCT116, SW480 and HCT15, did not inhibit the growth of Caco-2 and HEK293T cells.

**Families with fewer than 5 strains tested**

Among strains from families with less than five tested strains (n = 18), *Eubacterium* sp. 3_1_31 and *Propionibacterium* sp. 5_U_42FAA cells increased cell growth rates in SW480 and HCT15 cells, respectively (Table 1). Interestingly, *Pseudomonas* sp. 2_1_26 secretomes significantly inhibited the growth rates of all cell lines while the cells specifically inhibited SW480. *Pediococcus acidilactici* 7_4 and *Ruminococcaceae bacterium* D16 cells selectively inhibited HEK293T, while *Synergistes* sp. 3_1_syn1 also inhibited HCT116. Aerobically cultured *Ralstonia* sp. 5_2_56FAA secretomes inhibited 2 cell lines, SW480 and HEK293T. For *Desulfovibrio* sp. 3_1_syn3, the secretome increased Caco-2 growth while the bacterial cells increased HT29 growth.

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**Figure 4.** Influence of strains with and without encoded virulence factors on cellular growth rates. *B. fragilis* (a), *Fusobacteriaceae* (b), *Enterobacteriaceae* (c) and *Clostridiales* (d) encoding the virulence factors *bft, fadA, any toxin* (Map, pks, ShEt1, Stx2A), and any toxin (*TcdA, TcdB, alpha-toxin*), respectively, were compared to strains without these encoded virulence factors (Mann-Whitney U test). A trend toward cell growth enhancement was observed for *B. fragilis* bft+ secretomes (p = .07) (A), while significant inhibition of cell growth was observed for fadA+ *Fusobacteriaceae* cells (p < .0001), and secretomes (p < .05) (B), for *Enterobacteriaceae* secretomes encoding toxins (p < .0001) (C), and for *Clostridiales* secretomes encoding toxins (p < .0001) (D).

*4_1_47FAA and D9, and Klebsiella sp. 1_1_55* while no effects were observed for bacterial cells.
Contrastingly, these same bacterial cells decreased the growth of HCT116 and HEK293T. Similarly, *Desulfovibrio* sp. 6_1_46FAAA cells decreased the growth rates of HCT116, SW480 and HEK293T cells.

**Strain-specific effects that contrast their family**

While strains from the same bacterial families tend to have similar effects on the growth rates of cell lines (Supplementary Table S7), large variations were observed within certain families. For instance, within the *Clostridiaceae* family, the secretome of *Clostridium septicum* (Mace 1889) Ford 1927 strongly inhibited the growth of all tested cell lines, while the *Clostridium* sp. D5 secretome enhanced growth, although the significance threshold was not reached (Figure 1 and Supplementary Table S4). Similarly, *Bacteroides* sp. 2_1_22 cells strongly inhibited the growth rates of all cell lines while *B. fragilis* K570 cells enhanced growth in most cell lines (Figure 1 and Supplementary Table S3). A large growth rate variation was also observed in cell lines exposed to *Lachnospiraceae* cells (n = 11). For this family, the cells of *L. bacterium* 3_1_46FAAA and *L. bacterium* 8_1_57FAAA significantly inhibited the growth rates of all cell lines, while most of the other strains showed weak or low inhibition.

The observed variations in growth effect within bacterial families might either be explained by the phylogenetic distances between the strains or by characteristics with a non-phylogenetic distribution, such as accessory functions. To evaluate this, we measured the distances on a phylogenomic tree and compared them with the differences in growth rate effects over the cell lines. For most bacterial families, there is no association between the growth rate effects and phylogenetic distances (Supplementary Table S8, Supplementary Figure S4), suggesting that these effects may be associated with accessory functions that are independently gained or lost in different lineages. For *Erysipelotrichaceae* secretomes and cells, *Fusobacteriaceae* cells, and *Lachnospiraceae* secretomes, we found that the phylogenetic distance was significantly correlated with the growth rate effects, suggesting that these effects may be associated to variations in universally shared genes or to accessory functions with phylogenetically consistent dynamics.

**The presence of virulence genes is associated with growth rate inhibition**

To explain the complex, sometimes strain-dependent patterns of growth enhancement or inhibition, we analyzed the genome sequences of 144/157 tested gut bacteria. First, we checked whether 35 known bacterial family-specific virulence genes were present within the genomes of the tested strains (Supplementary Table S9), because some of the encoded virulence factors have been linked to cell proliferation changes. Nine toxin genes were found, including TcdA, TcdB,\(^{72}\) alpha-toxin\(^{68}\) of *Clostridiales*; FadA,\(^{31}\) colibactin present on the pks-island;\(^{73,74}\) Shiga toxin ShEt1; *Shigella* enterotoxin Stx2A; effector protein Map secreted by *Citrobacter rodentium* and *Escherichia coli;\(^{75,76}\) and BFT.\(^{38}\)

As shown in Figure 4, strains containing virulence genes tend to change growth rates, relative to related strains without those virulence genes. For example, the secretomes of *Clostridiales* strains significantly inhibited growth rate if their genomes contained TcdA, TcdB or alpha-toxin (Figure 4(a), \(p < .0001\)). TcdA and TcdB secretion by *Clostridioides difficile* (Prévot 1938) *Lawson et al. 2016* was confirmed by ELISA (data not shown). Similarly, *Fusobacteria* strains encoding the membrane protein FadA, which might be responsible for growth rate changes of epithelial cells,\(^{31}\) inhibited growth rates significantly (Figure 4(b), \(p < .0001\) and \(p < .05\) for bacterial cells and secretomes, respectively). Within the *Enterobacteriaceae* family, multiple secreted toxins were identified in the genomes, including colibactin, ShEt1, Stx2A, and Map. Secretomes of *Enterobacteriaceae* possessing any of these toxins inhibited cell growth relative to strains without these toxins (Figure 4(c), \(p < .0001\)). No differences were observed between bacterial cells of *Enterobacteriaceae* with or without the toxin genes. The secretomes of three *B. fragilis* strains encoding the secreted enterotoxin bft enhanced the growth rates of CRC cells, while the five *B. fragilis* strains without the toxin did not, although this trend was not significant (Figure 4(a), \(p = .075\)). The trend
was not observed for bacterial cells, where only B. fragilis K570 significantly enhanced growth rates (Table 1).

Functional categories associated with differential effects within families

Besides the toxin genes described above, other genes might also play a role in the observed effects. Thus, we next performed an unbiased comparative genome analysis where we associated all annotated bacterial genes to the effect of the strain on cell growth. We next grouped the genes by functional category and assessed whether genes within a category were significantly associated with cell growth alterations. Based on the PATRIC annotations we evaluated genes in two hierarchical levels of functions, namely ‘Subsystems’ and ‘Superclass’. All functional categories found to be significantly associated with cell growth alterations are listed in Supplementary Table S10.

The “Membrane transport” superclass was significantly associated with inhibiting Enterobacteriaceae cells and secretomes, and with enhancing Fusobacteriaceae secretomes. Other enriched superfamilies included “Cell envelope”, “Metabolism”, “Cellular processes”, “Protein processing”, “Energy”, and “Stress response, defense and virulence” (Supplementary Table S10). Subsystems including “Cobalamin synthesis”, “Ethanolamine utilization”, and the “Dpp dipeptide ABC transport system” were consistently found to be associated with growth enhancement or inhibition in multiple bacterial families and in multiple cell lines (Supplementary Table S10). The significant effects of other subsystems including “Vir-like type 4 secretion system”, “Glutamate fermentation”, “Flagellum”, “Branched-chain amino acid biosynthesis”, and others were associated with specific bacterial families. We expect that these statistical associations will provide valuable leads for future experiments.

Discussion

Over 150 human gut bacteria belonging to taxonomic groups that have previously been associated with CRC were tested for their effects on cell growth of five CRC cell lines (HT29, Caco-2, HCT116, SW480, and HCT15) with different mutation landscapes representing commonly mutated genes in CRC, and one immortalized kidney cell line (HEK293T). Our results show that cell growth enhancing and inhibiting effects of gut bacteria depend both on the mutational landscape of the cells and on the identity and functional content of the bacteria, including secreted or cell wall attached factors. CRC is a heterogeneous disease, as is reflected in the five different CRC cell lines with mutations in key CRC related genes (KRAS, APC, BRAF, TP53, CTNNB1, and PIK3CA). These mutations are absent from the HEK293T cells, an immortalized kidney cell line that was transformed by DNA of a human adenovirus, causing senescence in the pRb and p53 pathway. HEK293T cells were affected by the lowest number of gut bacterial strains, with only small differences.

Closely related bacteria may have distinct effects on cell growth

A striking observation was that the effects of bacterial cells and secretomes were strongly associated with bacterial families, although exceptions to this general pattern were observed. Previous studies have found that Fusobacteria, Streptococcus, Peptostreptococcus, Ruminococcus, and Escherichia/Shigella tend to be enriched in tumors, while Bacteroides, Pseudomonas, Lachnospira, and Anaerostipes are depleted. Regardless of the enrichment or depletion on CRC tumors in vivo, our in vitro results indicate that closely related bacteria may have distinct effects on cell growth. For example, we observed that different Fusobacteria strains, which were all cultured in the same media under the same conditions, may have opposing effects on cell growth, while these bacteria are consistently enriched in patients. This suggests that specific bacterial factors may play a role.

Fusobacteriaceae: enhancing or inhibiting growth?

Previous studies indicated that Fusobacteriaceae enhanced the growth of HCT116, SW480, and HT29 cells. F. nucleatum ATCC25586 was shown to enhance CRC cell growth via interaction with Toll-like receptors TLR4, TLR2, and myeloid differentiation primary response protein 88 (MYD88), and the
F. nucleatum strain Fn12230 was previously shown to enhance cell growth through the cell-wall attached adhesin FadA. Our results showed that most Fusobacteriaceae significantly inhibited cellular growth, including Fusobacterium nucleatum ATCC25586, contrasting previous observations, and that the presence of the FadA gene was significantly linked to growth inhibition by cells and secretomes (Table 1). Notably, the secretomes showed less pronounced inhibition than bacterial cells. While FadA is membrane-bound, secretomes likely include outer membrane vesicles (OMVs) that are 40–110 nm in size and express FadA. To rule out the possibility that these contrasting results were due to differences in methods, we compared our high-throughput MTT-based assay, used in this study, with the cell counting assay, that was employed in the referred studies, on six cell lines, using six different secretomes, and found a high correlation (Pearson’s r = 0.813, Supplementary Figure S5). These results confirm that the observed inhibitory effects by Fusobacteriaceae strains do not depend on the applied experimental techniques, but may partly be due to the use of different Fusobacteriaceae strains (Fn12230, ATCC25586, ATCC23726) isolated from extra-intestinal sites. Specifically, HCT15 cell growth was significantly inhibited by 11/16 Fusobacteriaceae strains (nine cells and three secretomes). HCT15 cells contain a missense mutation in MYD88 (p.Arg264Ter) resulting in an MYD88 deficiency, and may thus explain the insensitivity of this cell line to F. nucleatum subsp. animalis 11_3_2 and F. necrophorum subsp. funduliforme 1_1_36S, the two Fusobacteriaceae strains that promoted the growth of the other cell lines, but not of HCT15.

**Virulence factors are associated to cell growth inhibition**

The growth-inhibiting properties of Clostridiales secretomes may result from toxins encoded by this family. We observed that Clostridoides difficile (Prévot 1938) Lawson et al. 2016 and Clostridium septicum (Mace 1889) Ford 1927 secreting the virulence factors TcdA and TcdB, and lethal alpha toxin, respectively, significantly inhibited cell growth compared to the other Clostridiales secretomes. Clostridium septicum has been associated with CRC in the past.

Enterobacteriaceae strains encoding the toxins colibactin, Stx2A, ShEt1, and Map inhibited cell growth relative to non-toxigenic Enterobacteriaceae strains. Colibactin (present in four of our Escherichia strains) is a genotoxic compound that alkylates DNA and generates DNA double-strand breaks. The DNA-damaging effect of colibactin may not directly affect cell growth in our experimental setup, although colibactin may induce cell cycle arrest via SUMOylation of TP53 causing inactivation. Cell cycle arrest would stall cell growth, which can only happen in TP53 wild-type cell lines with active TP53 protein. Indeed, the only cell lines that were relatively inhibited by strains encoding colibactin were the wildtype TP53 cell lines HEK293T and HCT116.

An important virulence factor of B. fragilis is BFT. While in previous research, BFT was shown to enhance cell growth of HT29/c1 cells via β-catenin nuclear signaling, enterotoxigenic B. fragilis secretomes (VP113784 (bft1), 86–5443-2-2 (bft2), and K570 (bft3)) only marginally enhanced cell growth in our assay compared to the other B. fragilis secretomes in our screen. The mild effects as observed may depend on the level of toxin production and secretion under our experimental conditions. Interestingly, enterotoxigenic B. fragilis str. Korea 570 cells significantly enhanced growth, which might be independent of BFT and related to other cell-wall-attached factors.

**Streptococcus bovis strains specifically enhance growth of HT29 cells**

Streptococcus bovis infections have been linked to colorectal neoplasia. Our results demonstrate that secretomes of Streptococcus bovis strains that are closely related to known S. gallolyticus and S. pasteurianus strains (Supplementary Figure S3), selectively enhance the growth of HT29 cells. This is in line with a previous study showing cell growth enhancing effects in HT29 and HCT116 cells that were mediated through β-catenin, while no effect was observed in SW480 and HEK293T cells. An intact β-catenin pathway may, therefore, be required to observe the specific effect of S. gallolyticus on cell growth as is the case in HT29 cells, which may thus be sensitive to the effects of S. gallolyticus in our assay. HCT116 cells have a mutation in CTNNB1 (p.Ser45del) that interferes with β-catenin degradation, which may explain its insensitivity to S. gallolyticus.
Alternatively, mutations in the APC-gene in the other CRC-cells\textsuperscript{99} may interfere with β-catenin signaling depending on the specific mutation.

**Bacterial functions associated with cell growth alterations**

Several functional categories were significantly associated with cell growth. Most of these functions were identified in specific bacterial families and consistently inhibited or enhanced cell growth in different cell lines. These functions may reflect novel pathways of bacterial interference with cell growth which, to our knowledge, have not been previously identified. Most functions are related to cell metabolism, secretion, and transport systems. For example, secretomes of *Enterobacteriaceae* that encode the “Vir-like type 4 secretion system” inhibited the growth of all cell lines. Similarly, the gene superclass “Membrane transport” was mostly associated with secretomes that inhibited cell growth (Supplementary Table S10). Molecules that are secreted by these bacterial transport systems may be responsible for the inhibiting effect. For example, the Vir-like type 4 secretion system allows bacteria to secrete proteinaceous effectors that kill competitors.\textsuperscript{90} Here, we report an important first step in understanding the cell growth enhancing or inhibiting effects of human gut bacteria by identifying putative transport systems for effector molecules.

While the growth effects of secretomes were mostly associated with membrane transport functions, effects on cell growth by bacterial cells were associated with metabolic pathways (Supplementary Table S10). It is possible that these associations may be attributed to metabolic enzymes that may still be, although the bacterial cells were inactivated. In other ecosystems, metabolic enzymes of dead bacteria have been shown to remain active for up to 96 h.\textsuperscript{91} Our experiment was performed within 72 h. The synthesis of cobalam (vitamin B12), which is important for human cell proliferation \textit{in vitro} \textsuperscript{92} and a common addition to cell culturing media, was significantly associated with the cell growth enhancing effect of *Bacteroidaceae* and *Enterobacteriaceae* cells. Several possible scenarios can be envisioned to explain this observation. First, it remains to be tested whether vitamin B12 was retained in the bacterial fraction after washing, stimulating human cell growth on those cells. Second, different human gut-associated *Bacteroides* strains contain surface-exposed lipoproteins with high affinity for cobalam.\textsuperscript{93} Such molecules may remove cobalam directly from the media, making it unavailable to the human cells and impeding their growth.

The “Ethanolamine utilization” pathway was another metabolic process that we found to be associated with the inhibiting effect of *Fusobacteriaceae* cells on the growth rate of CRC cell lines. Ethanolamine (EA) is the basal component of phosphatidylethanolamine, a major phospholipid in animal cell membranes and is utilized by bacteria including *Fusobacteria* as a carbon source in the gut.\textsuperscript{94–96} EA sensing regulators have been proposed to regulate virulence in *Enterobacteriaceae*.\textsuperscript{96,97} It remains to be tested if the EA in human cell membranes triggers virulence factors in *Fusobacteria* that encode EA utilization genes, or perhaps if inactivated *Fusobacteria* cells retain the capacity to digest and kill human cells by the activity of EA utilization enzymes.

Overall, the statistical associations found in this exploratory analysis provide valuable clues about the possible functions and mechanisms that drive the interactions of bacteria with human cells in the gut. Future experiments are needed to confirm whether these factors are causal and to further clarify the molecular mechanisms involved.

**Outlook**

Our broad screen revealed many significant associations between gut bacteria, their encoded functions, and growth enhancement or inhibition of human cell lines. Many growth rate effects remain elusive and need to be further examined, such as their specificity for CRC. Moreover, it will be important to investigate the growth rate effects of living bacterial communities or mock communities, which are thought to synergize in affecting tumorigenesis,\textsuperscript{53} but fall outside the scope of our present study. Bacterial effects should be examined in more natural environments that mimic the conditions in the human gut, e.g. in co-culture with mucus, immune cells, and/or within organoid cultures.\textsuperscript{98} Our study forms an important baseline for such further research by identifying important traits in individual bacterial strains that are associated with enhancing or inhibiting cell growth, as well as important host factors that determine sensitivity to those bacterial traits. The influence of individual bacterial traits should be the focus of follow-up research and can be investigated
with add-back experiments, by introducing, e.g. virulence genes in non-virulent strains.

Conclusions

Above, we presented the results of the first large-scale analysis of the effects of gut bacteria on CRC cell growth. Our results with bacterial cells and secretomes reflect the complexity of the microbe–host interactions in the human gut. First, bacterial families tend to have consistent effects on cell growth, although these effects are not universal. This may partially be explained by the presence of virulence genes in some strains. Second, there are cell line dependent effects in cells due to their mutational landscape heterogeneity. Notably, these cell lines have acquired several different cancer hallmarks, yet their growth can still be influenced by specific bacteria. Our results suggest that the response of epithelial cells to gut bacteria may differ between patients due to differences in their gut bacteria. The relevance of our in vitro results for patients and CRC specificity remains to be studied. Cell growth enhancing and inhibiting bacterial traits could be important for determining risk for cancer or its progression, and these consequences could potentially be counteracted by microbiome modulation. Currently, microbiome medicine applications have already been adapted to some other diseases. For instance, Akkermansia muciniphila was demonstrated to improve diabetes type 2 and obesity in mice.17,18 Resistant life-threatening Clostridium difficile infections and ulcerative colitis were successfully treated with fecal microbiota transplants (FMT).19,20 Our study highlights the complex processes at play at the gut–microbiome interface and contributes to a better understanding of the role of different bacterial strains in altering colonic epithelial cell proliferation.

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