1	Host-microbiome protein-protein interactions reveal mechanisms
2	in human disease
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# 8 Abstract

- 9 Host-microbe interactions are crucial for normal physiological and immune system development and are
- 10 implicated in a wide variety of diseases, including inflammatory bowel disease (IBD), obesity, colorectal
- 11 cancer (CRC), and type 2 diabetes (T2D). Despite large-scale case-control studies aimed at identifying
- 12 microbial taxa or specific genes involved in pathogeneses, the mechanisms linking them to disease have
- 13 thus far remained elusive. To identify potential mechanisms through which human-associated bacteria
- 14 impact host health, we leveraged publicly-available interspecies protein-protein interaction (PPI) data to
- 15 find clusters of microbiome-derived proteins with high sequence identity to known human protein
- 16 interactors. We observe differential presence of putative human-interacting bacterial genes in
- 17 metagenomic case-control microbiome studies. In 8 independent case studies, we find evidence that the
- microbiome broadly targets human immune, oncogenic, apoptotic, and endocrine signaling pathways in relation to IBD, obesity, CRC and T2D diagnoses. This host-centric analysis strategy provides a
- relation to IBD, obesity, CiC and 12D diagnoses. This nost-centric analysis strategy provides a
   mechanistic hypothesis-generating platform for any metagenomics cohort study and extensively adds
- 20 Incontains in provide sis-generating platform for any metagenomics conort study and ext 21 human functional appointation to commensal bacterial proteins
- 21 human functional annotation to commensal bacterial proteins.

# 22 One-sentence summary

- 23 Microbiome-derived proteins are linked to disease-associated human pathways by metagenomic and
- 24 protein-interaction analyses.

## 25 Main Text

- 26 Metagenomic case-control studies of the human gut microbiome have implicated bacterial genes in a
- 27 myriad of diseases. Yet, the sheer diversity of genes within the microbiome (1) and the lack of functional
- annotations (2) have thwarted efforts to identify the mechanisms by which bacterial genes impact host
- 29 health. In the cases where functional annotations exist, they tend to reflect the proteins' most granular
- 30 molecular functions (*e.g.* DNA binding, post-translational modification) rather than their role in
- 31 biological pathways (3) and few, if any, relate to host cell signaling and homeostasis. Associating any
- 32 commensal bacterial gene and a host pathway has thus far required experimental approaches catered to
- each gene or gene function (4, 5).
- 34 Protein-protein interactions (PPIs) have revealed the mechanisms by which pathogens interact with host
- tissue through in-depth structural studies of individual proteins (5–7), as well as large-scale whole-
- organism interaction screens (8, 9). We hypothesized that host-microbiome PPIs may underlie health
- 37 status and could serve to provide additional information, through annotation of human pathways, about
- the role of bacteria in modulating health. There are already canonical examples of protein-mediated
- 39 microbe-associated patterns (MAMPs) that directly trigger host-signaling pathways through pattern
- 40 recognition receptors present on epithelial and immune tissues (10), such as flagellin with Toll-like
- 41 receptor 5 (TLR5). Several recent observations have further underscored a role for commensal-host PPIs
- 42 in health: a protease secreted by *Enterococcus faecalis* binds incretin hormone glucagon-like peptide 1
- 43 (GLP-1), a therapeutic target for type 2 diabetes (T2D) (11); and a slew of ubiquitin mimics encoded by
- both pathogens (12) and gut commensals (13) play a role in modulating membrane trafficking.
- 45 Currently, few experimentally-verified PPIs exist between bacterial and human proteins (roughly 8,000 in
- the IntAct database (14)) and only a handful of these involve proteins pulled from the human gut
- 47 microbiome. Expanding the commensal-human interaction network through state-of-the-art structural
- 48 modeling (15) is untenable, as there are few sequences homologous to genes found in metagenomes
- represented in co-crystals from the Protein Data Bank (16) (PDB) (Fig. S1, Supplementary Note 1). In the
- absence of structure and experimental data, sequence identity methods have been used to great effect to
- 51 infer host-pathogen PPI networks for single pathogens (17–19), but such approaches have not yet been
- 52 applied at the community-level, as would be required for the human gut microbiome.
- 53 All pathogen-host interactions are initially implicated in virulence, whereas microbiome-associated
- 54 disorders tend not to follow Koch's postulates (20). To distinguish PPIs that may be associated with
- 55 health versus disease, we compared community-level PPI profiles in large case-control cohorts of well-
- 56 established microbiome-associated disorders—namely colorectal cancer (CRC) (21–24), T2D (25, 26),
- 57 inflammatory bowel disease (IBD) (27) and obesity (28) (Fig. 1A, Table S1). In order to build
- 58 community-level PPI profiles, we mapped quality-filtered metagenomic sequencing reads from eight
- 59 studies to a newly constructed database of bacterial human-protein interactors and the bacterial members
- 60 of their associated UniRef clusters (Fig. S2, Supplementary Methods), which represent homeomorphic
- 61 protein superfamilies through sequence identity (29). Using a normalized feature importance ranking
- from random forest classifiers trained on each disease cohort (Fig. S3, Supplementary Methods), we find
- 63 46,734 commensal bacterial proteins (comprising 579 clusters) associated with disease, by virtue of their
- 64 putative interactions with 1,145 human proteins.





80 Interaction does not need to be conserved across homologous proteins in different bacterial species. A key

- 81 concern is the disproportionate number of bacteria-human PPIs in IntAct derived from high-throughput
- 82 screens performed on three intracellular pathogens: *Yersinia pestis*, *Francisella tularensis* and *Bacillus*
- 83 *anthracis* (8). However, we find that patient-detected bacterial clusters are not biased towards the
- 84 originating classes of these three pathogens—Bacilli or Gammaproteobacteria—and rather, reflect the
- 85 breadth of taxa typically associated with human gut microbiomes (Fig. 1C and S4). We verified that
- 86 human microbiome proteins have high amino-acid similarity to experimentally-verified human interactors
- 87 in the same UniRef cluster (Fig. 1C and S5). Additionally, interspecies bacterial-human protein interface
- residues, in general, are highly similar, or even identical, between members of the same UniRef cluster
  (Fig. S6, Supplementary Note 2). Although we appreciate that there will be commensal-human PPIs that
- 90 are not captured by this approach due to the limited scope of experimental data available, this is the
- 91 largest and only dataset of microbiome-host associated PPIs.
- 92 Surprisingly, the 816 human proteins we associate with CRC via the microbiome contain a number of
- 93 previously identified CRC-associated genetic loci, including well-known cancer genes: tumor protein
- 94 p53, epidermal growth factor receptor (EGFR), matrix metalloprotease 2 (MMP2), and insulin-like
- 95 growth factor-binding protein 3 (IGFBP3), among others (Fig. 1D). This represents a larger trend: the
- 96 1,145 human interactors are overall enriched for proteins with previously-reported gene-disease
- 97 associations (GDA) in CRC, T2D, and IBD (Fig. 1E, Table S2), with the exception of obesity, where
- 98 annotation is generally scarce. In line with mixed etiologies of diseases, we see that GDAs are not
- disease-cohort specific (Fig. S7). In fact, 36 percent of our genes have more than one GDA for our
- 100 diseases of interest. We suspected this may extend to autoimmune diseases, which are increasingly
- studied in the context of the gut microbiome (30), and we confirm enrichment of GDAs for autoimmune
- 102 disorders in the human proteins implicated by our method. This concordance between known disease
- annotation and disease association through our method increases our confidence that we are capturing
- 104 relevant molecular heterogeneity underlying microbiome-related disease.
- 105 If these bacterial proteins are indeed modulating human health through PPIs, we should expect them to
- 106 contain signatures of surface localization or secretion. We find that a majority of disease-associated
- 107 bacterial protein clusters (90.2%) contain proteins that are transmembrane, are secreted by type 3 or type
- 4 secretion systems, and/or contain eukaryotic-like domains, another marker for secretion (Fig. 1F). The
- remaining 9.8% may also be adequately localized, but our annotations do not cover all bacterial secretion
- systems, and it is unclear whether bacterial lysis may result in protein delivery to the host.
- 111 One of the major advantages of our work is that through this new interaction network, we vastly improve
- 112 our ability to annotate host-relevant microbiome functions. 35.8% of our disease-associated bacterial
- 113 clusters contain no members with annotated microbial pathways in KEGG (Kyoto Encyclopedia of Genes
- and Genomes) (31) (Fig. 2A). Yet, most of these genes can now be annotated according to the pathways
- of their human targets, obtaining a putative disease-relevant molecular mechanism (Fig. 2B).
- 116 Interestingly, most of the bacterial clusters with KEGG pathway annotations also gain a secondary human
- pathway annotation. This dual function is not entirely surprising, as a number of these have orthologs that
- have been previously identified as bacterial 'moonlighting' proteins, which perform secondary functions
- in addition to their primary role in the cell (32). Mycoplasma pneumoniae GroEL and Streptococcus suis
- 120 enolase, a protein involved in glycolysis, bind to both human plasminogen and extra-cellular matrix
- 121 components (33, 34). Mycobacterium tuberculosis DnaK signals to leukocytes causing the release of the
- 122 chemokines CCL3-5 (35). Streptococcus pyogenes glyceraldehyde-3-phosphate dehydrogenase
- 123 (GAPDH), canonically involved in glycolysis, can be shuffled to the cell surface where it plays a role as
- an adhesin, and can also contribute to human cellular apoptosis (36). These examples distinctly illustrate
- how bacterial housekeeping proteins are used by pathogens to modulate human health. In this study, we
- 126 uncover commensal proteins that similarly may have 'interspecies moonlighting' functions and appear to
- 127 be pervasive throughout our indigenous microbiota.





- 129 (A) Paired stacked bar plots showing the disease-associated bacterial cluster pathways annotated by KEGG (left) and their
- inferred pathways according to the human proteins they target (right), as annotated by WikiPathways (59). (B) Human pathways
- 131 (annotated using WikiPathways) targeted by bacterial gene clusters detected in human microbiomes from these eight studies. The 132 top 75 human pathways that contribute the most annotations to bacterial clusters detected in the eight metagenomic cohorts that
- 133 previously lacked KEGG-based annotations are shown. (C) Human cellular pathways, enriched in our disease-associated human
- providesly lacked REGO-based annotations are shown. (c) runnan central pathways, enriched in our disease-associated numan proteins (with a Benjamini-Hochberg false discovery rate (BHFDR)  $\leq 0.05$ . –log(BHFDR), displayed on the barplot to the left),
- 135 are colored according to the percent of pathway members differentially targeted in each case-control cohort.

136 In evaluating the statistical significance of recurrent human functional annotations, we performed

137 pathway enrichment analysis on the implicated human proteins and find proteins with established roles in

cellular pathways coherent with the pathophysiology of CRC, IBD, obesity and T2D (Fig. 2C), namely 138

139 those involving immune system, apoptosis, oncogenesis, and endocrine signaling pathways. Though most 140 enriched pathways include human proteins associated with all four diseases, reflecting their associated

relative risks (37–41), there is heterogeneity in the identity and number of members associated with each 141

142 study. Far more human proteins from the antigen presentation pathway are associated with T2D, obesity 143 and IBD cohorts' microbiomes than with CRC, perhaps indicating a disease-specific association with this

144 process. We see this again with CRC, in the death receptor signaling pathway and remodeling of

epithelial adherens junctions. 145

We see specific examples of known molecular mechanisms for these diseases now implicated with 146

147 microbiome-host PPIs: We find that DNA fragmentation factor subunit alpha (DFFA) is associated with

148 T2D (in the Qin et al. cohort), and is involved in death receptor signaling, an important pathway for the

149 destruction of insulin-producing  $\beta$ -cells (42). Collagen alpha-1(I) chain (COL1A1) is also a significant 150 target associated with T2D (in the Karlsson et al. cohort), and plays a role in dendritic cell maturation and

hepatic fibrosis/hepatic stellate cell activation pathways, capturing known comorbidities between T2D

151

152 and hepatic steatosis and nonalcoholic steatohepatitis (NASH) (43). Proteins associated with CRC

153 spanned expected bacteria-associated pathways, such as the direct sensing of enterotoxins, e.g. heat-stable

enterotoxin receptor GUCY2C (in the Feng et al. and Zeller et al. cohorts); but also classical cancer-154

155 associated pathways, such as the maintenance of DNA integrity, e.g. protection of telomeres protein 1 (POT1) (in the Feng et al., Qin et al. and Schirmer et al. cohorts) and X-ray repair cross-complementing 156

protein 6 (XRCC6) (in the Feng et al. and Yu et al. cohorts), the latter of which is required for double-157

158 strand DNA break repair. Interestingly, actin-related protein 2/3 complex subunit 2 (ARPc2) (associated

159 in the Yu et al. and Karlsson et al. cohorts) regulates the remodeling of epithelial adherens junctions, a

160 common pathway disrupted in IBD (44), CRC (45) and, most recently, T2D (37). This host-centric

annotation is useful beyond large-scale analysis of metagenomic data, as it broadly enables hypothesis-161

driven research into the protein-mediated mechanisms underlying microbiome impacts on host health. 162

These data suggest a set of discrete protein interactions that induce physiological effects when delivered 163 to the host. Consistent with this idea, we find that indeed many associated human proteins are known drug 164 165 targets (Table S3). For example, both T2D cohorts' and the obesity cohort's microbiomes independently implicate human protein Rev-ErbA alpha (NR1D1), the target of the drugs GSK4112, SR9009 and 166 SR9011, which inhibit the binding of Rev-ErbA alpha with its natural ligand, heme (Fig. 3A). These 167 drugs have been shown to affect cellular metabolism *in vitro* and affect hyperglycaemia when given to 168 mouse models of metabolic disorder (46, 47). We also find instances where the off-label effects or side 169 effects associated with the drug match our microbiome-driven human protein association. For instance, 170 171 imatinib mesylate (brand name: Gleevec) has several human binding partners, including macrophage 172 colony-stimulating factor 1 receptor (M-CSF1R) (Fig. 3B), a human protein we associate with CRC (in 173 the Feng *et al.* and Zeller *et al.* cohorts), and platelet-derived growth factor receptor- $\beta$  (PDGFR-B), associated with obesity and T2D (in the Le Chatelier et al. and Qin et al. cohorts, respectively). Literature 174 175 on imatinib supports these findings: although imatinib is best known as a treatment for leukemia, it has 176 been shown to affect glycemic control in patients with T2D (48). Furthermore, imatinib can also halt the 177 proliferation of colonic tumor cells and is involved generally in inflammatory pathways, through its 178 inhibition of TNF-alpha production (49). Whereas the notion of microbiome-derived metabolites acting as

179 drugs is well-appreciated (50, 51), this work broadens the scope of microbiome-derived drugs to include

protein products acting through PPI. 180



## 181 Figure 3. Human proteins targeted by gut commensal proteins include known therapeutic drug targets. (A)RevErbA alpha

(NR1D1) binds several human proteins (not shown), DNA (not shown) and heme. GSK4112 competitively binds Rev-ErbA

alpha, inhibiting binding with heme. ParE is a microbiome protein present in a diverse range of organisms and has a high relative

184 risk associated with T2D. (B) Macrophage colony stimulating factor 1 receptor (CSF1R) is targeted by imatinib, among other

drugs, as well as the uncharacterized bacterial protein YqeH, a protein that has a low relative risk associated with CRC.

- 186 Here, we reveal for the first time an extensive host-microbiome PPI landscape. To achieve this, we
- 187 benefit from existing methods in pathogen-host PPI discovery, further informed by community-level PPI
- 188 profiles of genes differentially detected in human metagenomes. This work highlights the myriad host
- 189 mechanisms targeted by the gut microbiome and the extent to which these mechanisms are targeted across
- 190 microbiome-related disorders. However, this network is far from complete. Few of the interaction studies
- 191 on which this interaction network is based were performed on commensal bacteria and therefore, we may
- be missing interactions specific to our intimately associated bacteria. In addition to large-scale PPI studies
- 193 involving commensal bacteria and their hosts, further in-depth studies will be needed to fully characterize
- 194 these mechanisms, such as whether these bacterial proteins activate or inhibit their human protein
- interactors' pathways.
- 196 This platform enables a high-throughput glimpse into the mechanisms by which microbes impact host
- 197 tissue, allowing for mechanistic inference and hypothesis generation from any metagenomic dataset.
- 198 Pinpointing those microbe-derived proteins that interact directly with human proteins will enable the
- 199 discovery of novel diagnostics and therapeutics for microbiome-driven diseases, more nuanced definitions
- 200 of the host-relevant functional differences between bacterial strains, and a deeper understanding of the co-
- 201 evolution of humans and other organisms with their commensal microbiota.

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from random forest are available as supplementary material.

#### 416 **Methods**

### 417

#### 418 Building a putative bacteria-human protein-protein interaction (PPI) network

419 Interactions were downloaded from the IntAct database (14) [August 2018]. Only interactions with evidence codes that indicated binary, experimental determination of the interaction between UniProt 420 identifiers with non-matching taxa were preserved, thereby excluding co-complex associations, small 421 422 molecule interactions, and predicted interactions. This resulted in a set of 296,103 interspecies PPIs. 423 Interspecies protein interactors were mapped to their UniRef50 sequence clusters (52). UniRef50 Clusters 424 are calculated every week and are publicly available through the UniProt web service. UniRef50 clusters 425 are named after their seed sequence, which has at least 50% sequence identity to all other members in the 426 cluster. Additionally, all members in the cluster have at least 80% sequence identity to the seed sequence. 427 Given a UniRef cluster with an experimentally determined PPI with a human protein, all bacterial 428 members of the cluster are labeled as putative interactors. Human proteins that have not been verified by 429 the SwissProt curating platform are filtered out of the final interaction network. The latter step avoids the 430 over-annotation of human isoforms or homologs, or non-verified human proteins. Overall, we generate 8,808,328 bacteria-human PPIs involving 1,613,641 bacterial proteins and 4,186 reviewed human 431

- 432 proteins. This corresponds to 18,097 interactions between 33,123 UniRef clusters containing bacterial
- 433 proteins and the aforementioned 4,186 reviewed human proteins.
- 434

457

#### 435 Detection of human-targeting proteins in metagenomic shotgun sequencing data

436 Reads from eight metagenomic studies (Table S1) were downloaded from the Sequence Read Archive 437 (SRA) using fasterq-dump. Reads belonging to more than one replicate from the same patient were 438 concatenated and treated as a single run. Reads were then dereplicated using prinseq (v0.20.2) and trimmed using trimmomatic (v0.36) with the following parameters: 439

```
440
441
           Dereplication
442
           perl prinseq-lite.pl -fastq {1} -fastq2 {2} \
443
                    -derep 12345 -out format 3 -no gual header \
444
                    -out good \{3\} -out bad \{4\};
445
446
           {1,2} Refer to paired read input files
447
           {3,4} Refer to output filepaths
448
449
           Trimming
450
           java -Xmx8g -jar trimmomatic-0.36.jar \
451
                    PE -threads 5 {1}
452
                    ILLUMINACLIP:{2}:2:30:10:8:true \
                   SLIDINGWINDOW:4:15 LEADING:3 TRAILING:3 MINLEN:50
453
454
455
           {1} Refer to input files
456
           {2} Is the path to a fasta file of Nextera TruSeq adapters
```

458 Paired reads were combined into a single file and aligned to a protein library of all 1,613,641 human-459 interacting bacterial proteins generated above. This read-to-protein alignment was performed using 460 BLASTx through the DIAMOND (53) command line tool (v0.9.24.125). Read alignments were filtered to only consider results with an identity of at least 90% and no gaps. Bacterial proteins were considered 461 462 detected with sufficient depth and coverage: more than 10 reads across 95% of the protein sequence, 463 excluding 10 amino acids at each terminus. We assign any bacterial protein detection to its corresponding 464 UniRef homology cluster. Human-interacting bacterial clusters are marked as either 'detected' or 'not 465 detected' for each patient in each study. For each patient, we also generate a file of human proteins that 466 are targeted by their detected bacterial proteins based on our bacteria-human PPI network.

## 467 Identity, similarity, and conservation measurements

468 The sequence identity constraints imposed from a UniRef cluster's seed on all other member sequences

don't explicitly provide any information about the sequence identity, or similarity, between other pairs of

- 470 sequences in the cluster. We compute pairwise alignments in order to understand how appropriate our 471 annotation mapping is between proteins experimentally-verified to interact with human proteins and
- 472 bacterial members of the same UniRef cluster that were detected in metagenomic samples.
- 473 Experimentally-verified interactors are aligned to their metagenome-detected UniRef cluster members
- 474 using the Smith-Waterman local alignment algorithm with a BLOSUM62 matrix via python's parasail
- 475 (54) library (v.1.1.17). Amino acid identity is calculated as the number of identical matches in the
- 476 pairwise alignment, divided by the length of the experimentally-verified interactor. Amino acid similarity
- is likewise calculated as the number of matches in the pairwise alignment that represent frequent

478 substitutions (non-negative BLOSUM62 scores), divided by the length of the experimentally-verified479 interactor.

480

481 Each cluster has a different number of bacterial members, and thus, comparisons, so we need to

summarize the bacterial identity and similarity metrics per cluster. We represent the identity between

483 experimentally-verified and metagenomic-detected bacterial protein sequences for each cluster as mean,

- 484 median, or a weighted average (Fig. S5). Specifically, we calculate the weighted average of
- 485 identity/similarity as:
- 486
- 487

$$weighted\%match(cluster) = \sum_{member \in cluster} \frac{Prevalence_{member} \times \%match}{Prevalence_{cluster}}$$

488

489 Where Prevalence<sub>member</sub> is the percent of patients where the bacterial sequence was detected,

490 Prevalence<sub>cluster</sub> is the percent of patients where any bacterial sequence from the cluster was detected, and
491 %match is either the identity or similarity between the member and the experimentally-verified protein
492 interactor.

493

When necessary, we constructed multiple sequence alignments using only the experimentally-verified
interactor sequence and all the metagenome-detected members of its homology cluster in order to
quantify amino acid conservation at each site. We calculated the Jensen-Shannon divergence using the
code provided by Capra *et al.* (55) with a window size of 3.

498

## 499 Prioritization of disease-associated bacterial protein clusters and human targets

500 In order to identify heterogeneity in the prevalence of bacteria-human PPIs, we preprocessed the data into

501 two detection matrices. Each patient from each study is represented in two feature spaces: (a) a binary

502 vector of detected bacterial gene clusters or (b) a binary vector of putatively targeted human proteins.

503 Human proteins were considered redundant if they shared all the same bacterial protein partners in our

database, as their "detection" is, by definition, perfectly correlated in this design, and were treated as a

505 single feature. Additionally, we build a contingency table based on the case/control balance of the dataset 506 and the prevalence of each bacterial gene cluster or human protein. Features with an expected count of

507 less than 5 in any cell of the contingency table were also filtered out as being under- or over-detected.

508 We use these processed matrices to train a random forest machine learning classifier on the task of

separating case and control patients and, after verifying that they achieve reasonable performance on the

510 task using leave-one-out cross-validation (Fig. S3), we extract the feature importance from the classifiers.

511 Having used the scikit-learn (56) implementation of the random forest algorithm, feature importance

512 corresponds to the average Gini impurity of the feature in all splits that it was involved in. Gini feature

513 importance is a powerful prioritization tool, as it can capture the multivariate feature importance (whereas

simple metrics like log-odds ratio and corrected chi-squared statistics only capture univariate feature

importance). However, it has been noted that in sparse, binary decision tasks like our own (57, 58), this
feature importance can be can overestimate the importance of features based on their prevalence alone.

517 To obtain a normalized Gini feature importance, we perform a Monte Carlo estimate of the expected Gini

518 importance for each feature given the prevalence of all features in that dataset. On each iteration of the

simulation, we generate a random null feature matrix using a Bernoulli binary generator where:

520 
$$P(null[patient][feature] = 1) = Prevalence(feature)$$

521 We train two random forests on the disease labels for each patient, using either the real matrix or the null

522 matrix. Both real and null Gini feature importances are extracted for each feature and aggregated across

523 iterations of the simulation. The normalized Gini importance for each feature is expressed as a z-score:

524 
$$normGini_{f} = \frac{\overline{Gini_{f}^{real}} - \overline{Gini_{f}^{null}}}{\max(\sigma(Gini_{f}^{real}), \sigma(Gini_{f}^{null}))}$$

525 The simulation is repeated until the distance between the maximum and minimum normalized Gini

importance converges (at least 200 iterations of holding equal value). Code and preprocessed detection

527 matrices for each of the studies are provided in the Auxiliary Supplementary Materials.

528 This iterative procedure is a convenient way to generate a null feature importance for comparison, but

also provides a very robust measurement of Gini<sup>real</sup>. Random trees from a random forest act as

530 independent estimators: Given the same data, calculating the average importance of N forests with E

estimators is equivalent to the importance on a single forest with  $N \times E$  estimators. For the real Gini

feature importance calculation, our final estimates are equivalent to a forest with  $I \times E$  trees, where I is the

number of iterations at convergence. The entire procedure is analogous to iteratively increasing the

number of trained trees in a random forest (and its paired null model) with a step-size E (in our case,
 E=100) until normalized feature importance converges. An example is provided in the Auxiliary

535 E=100) until normalized feature importance converges. An example is provided in the Auxilia 536 Supplementary Materials.

536 Supplementary Materials.

537 Most of the normalized feature importances across studies fall at or below zero, indicating that their Gini

feature importance is not higher than would be expected in the null model (Fig. S8). This provides a

convenient cutoff (normalized Gini > 0) to prioritize a set of proteins, as human proteins with positive

540 normalized Gini importance capture proteins with large log-odds ratio magnitudes and rescues candidates

- that would've been missed through univariate analysis.
- 542

# 543 Human pathway annotation and enrichment analysis

544 Human pathway annotation was performed using the mygene python library. Specifically, we queried

- pathway annotations from WikiPathways (59), filtering out pathways from TarBase, as they specifically
   only include miRNA interaction annotation.
- 546 547
- 548 We performed pathway enrichment analysis using QIAGEN's Ingenuity Pathway Analysis (IPA) (60)

tool. All human proteins with a normalized feature importance greater than zero were uploaded as

550 UniProt identifiers into the desktop interface and submitted to their webserver for Core Enrichment

551 Analysis was conducted only on human tissue and cell lines and IPA's stringent evidence filter. Pathways

552 were considered enriched if they had both a  $-\log(p-value) \ge 1.3$  and a Benjamini-Hochberg False

- 553 Discovery Rate less or equal to 5%.
- 554

555 We additionally annotated all human proteins with any known drug targets from the probes-and-drugs

database (61) (04.2019 database dump), which aggregates drug-target interactions from the largest drug target databases.

## 558 Human gene-disease association

Disease annotations were extracted from all of GDAs from DisGeNET (62) (v.6.0). Lacking a simple
hierarchy of disease, we binned similar disease terms into the 5 larger categories relevant to our study.
Human protein identifiers were mapped to their Entrez gene ID's using the UniProt batch mapping
resource and then annotated with these 5 labels:

563

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605

564CRC: Adenocarcinoma of large intestine, Hereditary non-polyposis colorectal cancer syndrome,565Hereditary nonpolyposis colorectal carcinoma, Malignant neoplasm of colon stage IV, Malignant566neoplasm of sigmoid colon, Malignant tumor of colon, Microsatellite instability-high colorectal567cancer,

569 Diabetes: Brittle diabetes, Familial central diabetes insipidus, Fibrocalculous pancreatic diabetes, 570 Gastroparesis due to diabetes mellitus, Insulin resistance in diabetes, Insulin-dependent but 571 ketosis-resistant diabetes, Insulin-dependent diabetes mellitus secretory diarrhea syndrome, Insulin-resistant diabetes mellitus, Insulin-resistant diabetes mellitus at puberty, Latent 572 autoimmune diabetes mellitus in adult, Macroalbuminuric diabetic nephropathy, Maturity onset 573 diabetes mellitus in young, Maturity-onset diabetes of the young, type 10, Maturity-onset diabetes 574 575 of the young, type 11, Microalbuminuric diabetic nephropathy, Moderate nonproliferative diabetic retinopathy, Monogenic diabetes, Neonatal diabetes mellitus, Neonatal insulin-dependent 576 577 diabetes mellitus, Non-insulin-dependent diabetes mellitus with unspecified complications, 578 Nonproliferative diabetic retinopathy, Other specified diabetes mellitus, Other specified diabetes 579 mellitus with unspecified complications, Pancreatic disorders (not diabetes), Partial nephrogenic 580 diabetes insipidus, Prediabetes syndrome, Proliferative diabetic retinopathy, Renal cysts and diabetes syndrome, Severe nonproliferative diabetic retinopathy, Transient neonatal diabetes 581 mellitus, Type 2 diabetes mellitus in nonobese, Type 2 diabetes mellitus in obese, Type 2 582 583 diabetes mellitus with acanthosis nigricans, Visually threatening diabetic retinopathy, diabetes (mellitus) due to autoimmune process, diabetes (mellitus) due to immune mediated pancreatic 584 585 islet beta-cell destruction, diabetes mellitus risk, idiopathic diabetes (mellitus), postprocedural diabetes mellitus, secondary diabetes mellitus NEC 586

588 Autoimmune: Addison's disease due to autoimmunity, Adult form of celiac disease, Aneurysm of 589 celiac artery, Ankylosing spondylitis, Ankylosing spondylitis and other inflammatory 590 spondylopathies, Arteriovenous fistulas of celiac and mesenteric vessels, Blood autoimmune 591 disorders, Bullous systemic lupus erythematosus, Chilblain lupus 1, Dianzani autoimmune 592 lymphoproliferative syndrome, Dilatation of celiac artery, Hyperthyroidism, Nonautoimmune, 593 Latent autoimmune diabetes mellitus in adult, Maternal autoimmune disease, Multiple sclerosis in children, Neonatal Systemic lupus erythematosus, Subacute cutaneous lupus, Systemic lupus 594 erythematosus encephalitis, Venous varicosities of celiac and mesenteric vessels, Warm 595 596 autoimmune hemolytic anemia, diabetes (mellitus) due to autoimmune process, lupus cutaneous, 597 lupus erythematodes

599Obesity: Abdominal obesity metabolic syndrome, Adult-onset obesity, Aplasia/Hypoplasia of the600earlobes, Childhood-onset truncal obesity, Constitutional obesity, Familial obesity, Generalized601obesity, Gross obesity, Hyperplastic obesity, Hypertrophic obesity, Hypoplastic olfactory lobes,602Hypothalamic obesity, Moderate obesity, Overweight and obesity, Overweight or obesity,603Prominent globes, Simple obesity, Type 2 diabetes mellitus in nonobese, Type 2 diabetes mellitus604in obese

606 <u>IBD</u>: Acute and chronic colitis, Acute colitis, Allergic colitis, Amebic colitis, Chronic colitis,
 607 Chronic ulcerative colitis, Crohn Disease, Crohn's disease of large bowel, Crohn's disease of the
 608 ileum, Cytomegaloviral colitis, Distal colitis, Enterocolitis, Enterocolitis infectious, Eosinophilic

colitis, Food-protein induced enterocolitis syndrome, Hemorrhagic colitis, Ileocolitis, Infectious
colitis, Left sided colitis, Necrotizing Enterocolitis, Necrotizing enterocolitis in fetus OR
newborn, Neonatal necrotizing enterocolitis, Non-specific colitis, Pancolitis, Pediatric Crohn's
disease, Pediatric ulcerative colitis, Perianal Crohn's disease, Typhlocolitis, Ulcerative colitis in
remission, Ulcerative colitis quiescent

614

We additionally downloaded all human proteins involved in protein-protein interactions from the IntAct database and annotated them in the same manner in order to compare label frequencies.

617

# 618 Bacterial pathway, secretion, and taxonomy annotation

- 619 We submitted all bacterial protein sequences that were detected in human metagenomes to the
- 620 KofamKOALA (63) KEGG orthology search resource. We additionally submitted our bacterial sequences
- to EffectiveDB (64) in order to obtain predictions for EffectiveT3 (type 3 secretion based on signal
- 622 peptide), T4SEpre (type 4 secretion based on composition in C-terminus), EffectiveCCBD (type 3
- 623 secretion based on chaperone binding sites), and EffectiveELD (predicts secretion based on eukaryotic-
- 624 like domains). We used the single default cutoffs for T4SEpre, EffectiveCCBD, and EffectiveELD, and
- 625 chose the 'sensitive' cutoff (0.95) rather than the 'selective' (0.9999) cutoff for EffectiveT3.
- Transmembrane proteins or signal peptides were predicted using TMHMM (65) (v.2.0c), with a threshold
- 627 of 19 or more expected number of amino acids in transmembrane helices.
- 628
- 629 Bacterial taxonomy information was extracted from NCBI. UniProt identifiers and annotations were
- 630 downloaded using UniProt SPARQL endpoint.
- 631

# 632 Statistics

- 633 For Fig. 1D, we quantify the difference between the human proteins implicated in disease by our method
- 634 (StudySet) and all human proteins that have available protein interaction information (NullSet) by
- 635 comparing the proportion of these sets that have certain gene-disease associations. To do so we perform a
- 636 chi squared test (dof=1): The total number of proteins in these sets is 13,698 (NullSet) and 767
- 637 (StudySet). The breakdown of chi squared statistics and p-values can be found in Supplementary Table 6.

## 638 Supplementary Notes

## 639 Supplementary Note 1. Structural data available for these microbiome-human PPIs

640 Interaction network studies have increasingly moved towards structural interaction networks (66). These

641 networks represent not only the group of binary PPIs that have been detected, but also the partner-specific

642 interfaces on which these interactions occur. In the absence of resolved structural data for a given

bacterial-human PPI, structural PPI data of homologous proteins can be used to identify potential protein

644 interfaces.

645 We measured the extent to which structural interfaces could be used to infer gut commensal-human

- 646 protein-protein interaction by using DIAMOND (64) to query all amino acid sequences submitted to the
- 647 PDB for any templates that might match bacterial or human proteins in our putative interactor library. Out
- 648 of the 732 bacterial gene clusters that contain both members with experimentally-verified PPIs and were
- detected in human gut metagenomic sequences, 596 have BLASTP matches to a sequence in the PDB. A
- low-quality filter for at least 50% identity and 50% query coverage further lowers this set to 478 bacterial
- gene clusters. The same process and cutoffs detect PDB matches for 837 of the 2,140 human proteins in
- 652 our interaction network. The overlap of these two sets reveals 20 cocrystal structures that can provide
- 653 interface information for only 18 protein pairs including 15 bacterial gene-cluster proteins and 8 human
- 654 proteins (Fig. S1).

In order to identify interface residues between each pair of chains in the 20 cocrystal structures, we first

use NACCESS (67) to calculate the solvent accessibility of each residue in each chain. Chains with an

- accessible surface area of 15 Å or more are considered surface residues. We then calculate the change in
- accessible surface area for each residue when other chains in the same crystal structures are introduced.
- Residues which have a change in solvent accessible surface area above 1 Å are determined to be interfaceresidues (68).

661 While we identify interface residues in all 18 protein pairs (Table S4), 12 of these cases involve large

662 complexes where the human protein and bacterial protein match domains on more than one chain, and

sometimes the same chain (Table S5). Determining interface residues for two proteins with multiple

664 matches can complicate analysis, as they can result in multiple interfaces for the same protein partner. For

example, in PDB 2b3y, both the human protein IREB2 and the bacterial proteins from the Aconitate

666 hydratase cluster match domains in chains A and B. This would cause IREB2's interface residues to 667 contain interface residues from two sources in the same crystal structure. There are, however, 6 cases in

- 668 which the human protein and bacterial proteins match their respective chains exclusively. We highlight
- one example in which there are uniquely mapped chains, where 1p0s chains H and E match human
- 670 coagulation factor X and bacterial Ecotin, respectively (Fig. S10). Through this analysis, we demonstrate
- 671 the power of sequence homology searches in structural databases to confirm bacteria-human PPIs and
- 672 characterize their interfaces, but find that there are currently not enough representative sequences to do
- 673 structural prediction at a large scale for the commensal human microbiome.

# Supplementary Note 2. Conservation of interface residues in bacterial members of UniRef50 Clusters with human interactors in the PDB

- Functional annotations are commonly propagated between members of the same UniRef50 cluster (52,
- 677 69), yet it is not clear whether this intra-cluster conservation of function applies to exogenous interaction.
- To validate whether this is generally the case, we analyzed the conservation of interface residues across
- bacterial members of the same UniRef cluster for all bacteria-human protein-protein interactions
- 680 submitted to the Protein Data Bank (PDB).
- 681 Using UniProt's SPARQL API, we compiled a list of all PDB structures which contain both human
- proteins and bacterial proteins (751 structures as of January 2020), the UniRef50 cluster identifier for the
- bacterial protein, and all protein sequences in the corresponding cluster that also originate from bacterial

proteomes (68,434 unique bacterial protein sequences as of January 2020). Using Clustal Omega, we then

685 generated multiple sequence alignments for all the members of each UniRef50 clusters, excluding any

686 duplicated sequences. We calculated interface residues on all pairs of chains in each of the 751 structures 687 and measured the BLOSUM62 similarity between bacterial interface residues and their corresponding

amino acids in their respective UniRef50 cluster MSA.

689

690 Despite the small number of PPIs in our dataset that have representatives in the PDB (Fig. S1), examining

bacteria-human PPI co-crystal structures supports transfer of interaction among UniRef50 cluster

692 members. We find that there is high amino acid sequence identity and similarity between interface

residues in bacteria-human cocrystal structures and other bacterial members of the same cluster (Fig.S6).

We additionally calculate the Jensen-Shannon divergence on the columns of the MSA containinginterface residues and find that they are well-conserved (Fig.S6). Overall, we find evidence that interface

residues with a human protein interactor tend to be maintained between bacterial members of the same

697 UniRef50 cluster.

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#### **Supplementary Figures** 749

Figure S1. Few bacterial-human interaction sequences populate the Protein Data Bank.

750 751 752 A Venn diagram describing the number of detected bacterial clusters and human proteins in the eight metagenomic cohorts that 753 have any matching structure (using BLASTp) in the PDB and whether their structures appear on the same PDB cocrystal 754 structure.



- 755 Figure S2. An outline of our homology mapping procedure and alignment.
- 756 Depiction of the interaction network inference and protein detection pipeline. Note that only bacterial proteins found to be 757
  - human-interactors through the mapping procedure are used as candidates for detection in metagenomic studies.



### 758 Figure S3. Performance metrics.

A heatmap of precision, recall, and F1-scores for random forests with 5000 estimators, evaluated using leave-one-out cross-validation on each of the eight studies. Performances are listed for both the bacterial and human representations of the metagenomic sample. The bacterial representation lists all the bacterial genes detected in a patient that share a UniRef cluster with an experimentally-verified human-protein interactor. The vector of human proteins represents all the human proteins which might be targeted by the bacterial genes found in each metagenome.

100% 82.8 62.6 71.3 Feng2015-bacClust 62.7 84.9 72.1 Feng2015-humprot 96.1 98.0 97.0 Hannigan2017-bacClust 80% Hannigan2017-humprot 78.0 92.0 84.4 Karlsson2013-bacClust 71.1 99.0 82.8 72.3 92.2 81.0 Karlsson2013-humprot 60% 61.6 83.3 70.8 Lechatelier2013-bacClust Lechatelier2013-humprot 61.4 79.5 69.3 83.5 83.0 83.3 Qin2017-bacClust 72.6 66.7 69.5 Qin2017-humprot - 40% 80.2 97.7 88.1 Schirmer2018-bacClust 82.2 95.4 88.3 Schirmer2018-humprot 70.0 93.3 80.0 Yu2015-bacClust - 20% 67.4 75.3 85.3 Yu2015-humprot Zeller2014-bacClust 76.6 89.4 82.5 75.3 87.9 81.1 Zeller2014-humprot - 0% precision f1-score recall

26

## Figure S4. Taxonomic diversity in bacterial clusters detected in patients.

764 765 766 Histogram showing the number of species, genera, families, orders, classes and phyla for bacterial clusters with members detected in human microbiomes.



## Figure S5. Pairwise identity and similarity between proteins found in the human microbiome and those with

# respective text of the second second

Histogram showing the percent identity and similarity between bacterial proteins with experimental verification and their

corresponding detected proteins in human microbiomes in the same UniRef cluster. Three aggregation methods are used to

estimate each metric at a cluster level: median, average, and an average weighted by prevalence.



## 772 Figure S6. Interface similarity between bacterial proteins within a UniRef cluster.

Similarity, identity, and Jensen-Shannon divergence of interface residues across all bacterial members of the same UniRef cluster
 sourced from all cocrystal structures with human and bacterial interactors and no filtering based on our datasets.



## 775 Figure S7. Previous gene-disease associations for human interactors in our dataset.

The number of human interactors (with normalized feature importance greater than 0) according to their GDAs for CRC, T2D,
 obesity, IBD and autoimmunity.



Human proteins with gene-disease annotations

# Figure S8. Human protein interactors according to their normalized feature importance and log odds ratio. Volcano plots of the human protein interactors according to their normalized feature importance and log odds ratio

Volcano plots of the human protein interactors according to their normalized feature importance and log odds ratios in each case control cohort study.



### 781 Figure S9. Clustering of cases and controls is not due to disease status or study.

- 782 (A) Principal components analysis of patients by their detected human protein interactors, colored by study and label. (B)
- 783 Principal components analysis of detected human protein interactors for all samples in eight metagenomic studies colored by
- 784 disease status according to study. Controls are all colored together in blue.





## 786 Figure S10. Cocrystal structure of blood coagulation factor Xa in complex with Ecotin M84R.

787 Cluster Uniref50 Q1R9K8 contains several bacterial ecotins detected in human metagenomes. Using BLAST, we found highquality matches between members of this cluster and the structure 1p0s:E (Ecotin precursor M84R) in the PDB (identity of

789 97.2%, eval=1e-75). Our putative interactor to this cluster, coagulation factor X (P00742) likewise matched structure 1p0s:H

790 (coagulation factor X precursor) (identity of 100%, eval=3.8 e-150). Chain E is shown in blue, and chain H in orange, with their

791 interface residues highlighted as spheres. The linear model of both proteins is shown underneath. The linear model's colored

areas indicate the part of the proteins that were crystallized in this PDB, while the greyed-out areas indicate non-crystallized
 spans. The squares indicate the range of the BLAST match between our query proteins and the PDB reference sequences. Finally,

results in the squares indicate the location of interface residues as detected in this model. There are currently not enough

published structures to perform this analysis on all interactions involving detected bacterial genes (Fig. S1, Tables S4 and S5).



# 796 Supplementary Tables

# 797 Table S1. Metagenomic studies used in this research.

For each study, we list its disease focus, the labels in the cohort study, the patient count for each of the

799labels, how we grouped cases and controls, the number of detected bacterial clusters and inferred human

interactors, and the number of important bacterial and human proteins with normalized feature importancegreater than 0.

801 gl

# 803 Table S2. Human interactors that have known gene-disease associations.

Listed are the disease-associated human proteins (with normalized feature importance greater than 0) with GDAs in DisGeNET, along with the study in which they are found to be important.

806

# 807 Table S3. Human interactors that are known drug targets.

- For each disease-associated human protein (with normalized feature importance greater than 0), we list the drug interactor and the study in which it was found to be important.
- 810

# 811 Table S4. Interface residues from PDB chain pairs matching human and bacterial interactors in

- 812 our dataset.
- 813 All pairs of detected bacterial proteins and human proteins in the eight metagenomic datasets that have
- 814 BLASTp matches to two different chains within the same PDB cocrystal structure (totaling 15 bacterial
- protein clusters and 8 human proteins). Listed are the BLAST readouts for both matches, as well as the
- 816 interface residues for each chain at the PDB index, PDB sequence, and UniProt sequence mappings.
- 817

# 818 Table S5. Cocrystal structures representing interactions in our set.

- A summary of the PDB chain-pairs (presented in Table S4) that can be used as representatives to identify
- 820 interface residues for interactions in our set. We annotate each interaction by whether the bacterial and
- 821 human proteins match non-overlapping pairs of chains.
- 822

# 823 Table S6. Gene-disease association comparison statistics.

824 The set sizes, fractions, chi-squared statistics and p-values used to generate Fig. 1E.