Biogeography and Environmental Dynamics Shape Ecological

Networks of the Human Microbiome

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21 Introduction

Viruses and bacteria are critical components to the human microbiome and play important roles in health and disease. Bacterial communities have been associated with diseases including a wide range of skin conditions¹, acute and chronic wound healing conditions^{2,3}, and gastrointestinal diseases including inflammatory bowel disease^{4,5}, Clostridium difficile infections⁶, and colorectal cancer^{7,8}. Altered virones (consisting primarily of bacteriophages) have also been associated with various diseases and environmental perturbations including inflammatory bowel disease^{5,9}, periodontal disease¹⁰, and others^{11–16}. Individual phages are capable of lysing their bacterial hosts and modulating their functionality through horizontal gene transfer (i.e. transduction) and altering bacterial host gene expression. At the community level, the human virome has begun to be implicated in promoting antibiotic resistance throughout human-associated microbial communities 12,17. These virus community shifts are not reflections of the bacterial communities, but rather act in concert with bacteria as a single overall community 5,18 . This previous work has relied on studing bacteria and phages in isolation (e.g. taxonomic classification, alpha diversity, and beta diversity), reducing them to two seperate communities. In fact, bacteria and phage communities are dynamic and complex, acting in concert and sharing information (e.g. horizontal gene transfer) to maintain balanced, efficient ecosystems and their removal can disrupt or even collapse the ecosystem ^{18–28}. Relationship-based network approaches allow us to capture this information, including community resilience to instability and the degree of information flow across microbes. The reductionist approach to understanding the human microbiome is unable to capture this abundant information of how these communities interact and influence each other. Here we bridge this knowledge gap by characterizing the human bacterial and phage communities through their relationships, using machine learning and graph theory techniques. We focus on characterizing the network dynamics of the healthy human microbiome (including the skin and gut), so as to provide a foundation on which we and other can build further studies into the network dynamics associated with disease states. To begin characterizing the networks of bacteria and phages of the human microbiome, we leveraged three published microbiome datasets with paired virus and bacterial metagenomic sequence sets^{12,13,29,30}. These datasets included gut and skin samples, thus allowing us to more gain broader insights into the human microbiome as a whole. We built off of previous work on large-scale phage-bacteria ecological network analysis by inferring interactions using metagenomic datasets, instead of previous culture-based techniques^{23,24}. Our metagenomic interaction inference model is powered beyond previous models by its inclusion of protein interaction data, inclusion of negative interactions as well as positive, and the use of a more sophisticated

machine learning algorithm instead of inappropriate linear models³¹.

Using this approach we go beyond contemporary, isolated approaches to uncover a basic understanding of
the relationship dynamics associated with healthy human phage and bacterial communities. We find that
different people, body sites, and anatomical locations not only support distinct mcirobiome membership
and diversity, but also support communities with distinct communication structures and propensities toward
ecological instability. The importance of these findings is twofold. First, we provide a basic picture of healthy
microbiome network structure across the body. By understanding the healthy state, we empower future
studies to begin investigating how these community structures change in disease or otherwise altered states.
Second, by providing a new level of understanding for how the microbiome interacts and communicates, we can
begin to inform microbiome-based therapeutic development. Probiotic drugs and fecal microbiota transplants
(FMTs) rely on understanding community stability and the influence of microbes on each other so that they
may engineer the community. Our model provides a new level of understanding what those communities look
like and what degree of communication we should strive for. Understanding horizontal gene transfer of agents
such as antibiotic resistance genes also benefits from a map of the community's communication geography.

66 Results

67 Modeling Phage-Bacteria Infectious Networks

68 We studied the impact of human system environment (e.g. skin, gut, defined diets) on microbiome phage-

bacteria interaction networks by leveraging previously published sequence sets containing purified virome

samples paired with bacterial metagenomes from whole metagenomic shotgun sequences. Our study contained

datasets for three human virome studies, including a study of the impact of diet on the gut virome¹³, the impact

of anatomical location on the skin virome¹², and the virome of monozygotic twins and their mothers^{29,30}. The

viromes associated with these datasets were subjected to virus-like particle (VLP) purification to eliminate

other organism DNA including bacteria, fungi, and humans.

₇₅ Bacterial and viral sequences were quality filtered and assembled into contigs from their respective sample

sets. These studies were performed over five years using different methods and technologies, so therefore

yielded different sequence abundances (Supplemental Figure S1 A-B). Because contig assembly rarely

reconstructs entire genomes, we clustered related bacteria and phage contigs by k-mer frequency and co-

abundance using the CONCOCT algorithm. These clusters represent operationally defined clusters of related

bacteria and phage genomes that we defined as operational genomic units (OGUs), which are conceptually

similar to the operational taxonomic unit (OTU) and operational protein family (OPF) definitions used for grouping highly similar 16S rRNA and open reading frame sequences, respectively³². Contigs and OGUs demonstrated high sequence coverage and length (Supplemental Figure S2 - S3).

We predicted which phage OGUs will infect which bacteria using a random forest model trained on experimentally validated infectious relationships from six previous publications^{31,33–37}. This training set contained
diverse bacteria and phages, with both broad and specific infectious capabilities (Supplemental Figure S4

A - B). Phages with linear and circular genomes, as well as ssDNA and dsDNA genomes, were included
in the analysis. Because this was a DNA sequencing study, RNA phages were excluded from the analysis
(Supplemental Figure S4 C-D). This training set included both positive relationships (a phage infects a
bacterium) and negative relationships (a phage does not infect a bacterium). This built on previous work,
which focused almost exclusively on positive relationships only, by allowing us to validate the false positive
and false negative rates associated with our candidate models.

Four phage and bacterial genomic markers were used to predict infectious relationships between bacteria and phages: 1) Genome nucleotide similarities, 2) gene amino acid sequence similarities, 3) CRISPR targeting of phages by bacterial CRISPR spacer sequences, and 4) similarity of protein families known to be associated with known protein-protein interactions. The resulting random forest model exhibited high performance with an AUC of 0.853, a sensitivity of 0.851, and a specificity of 0.774 (Figure 1 A). The most important predictor in the model was nucleotide similarity between genes, followed by nucleotide similarity of whole genomes (Figure 1 B). Protein family interactions were moderately important to the model, while CRISPRs were minimally important, likely because they were redundant with the blast information. Approximately one third of the relationships yielded no score and were automatically classified as being non-infectious (Figure 1 C).

Upon validation and completion, we used the random forest model to classify the relationships between 103 bacteria and phages in the experimental datasets. The relationships within the three studies were used to construct one master network, containing the three study sub-networks, which themselves each contain 105 sub-networks for each sample (Figure 1 D). Metadata including study, sample ID, disease, and OGU abundance within the community (based on sequence count) were also stored in the multi-study master 107 network to allow for effective parsing and downstream analysis (Supplemental Figure S5). The resulting master network was highly connected and contained 72,287 infectious relationships among 578 nodes, 298 of 109 which represented phages and 280 that represented bacteria. Although the network was highly connected, not 110 all relationships were present in all samples. Furthermore, relationships were weighted by relative abundance 111 of their associated bacteria and phage, meaning that lowly abundant relationships could be present but insignificant compared to those that were more highly abundant. Like the master network, the skin network exhibited a diameter of 4 (measure of graph size; the greatest number of traversed vertices required between two vertices) and included almost all of the master network nodes and edges (Figure 1 E - F). The gut diet and twin sample sets each contained less than 200 vertices, less than 20,000 relationships, and diameters of 3, suggesting more sparsely related phages and bacteria (Figure 1 E - F).

Interpersonal and Intrapersonal Diversity of Gut Microbial Networks

Previous work has show reduced intra-personal (within person) diversity in the viromes and bacterial communities of the human gut and skin compared to inter-personal (between people) diversity^{12,38,39}. Understanding this conservation of the microbiome has been important for establishing a basic understanding of microbiome individuality and aids in interpreting microbiome study results. We hypothesized that like other aspects of these microbial communities (e.g. diversity, community composition), there is a strong conservation of network structure in the skin and gut.

We tested this hypothesis by calculating the degree of dissimilarity between subject graphs while incorporating

125 both phage and bacteria abundance, as well as centrality within the node. This was accomplished by 126 first calculating the weighted Eigenvector centrality of all bacteria and phages within each sample graph. Conceptually, this metric defines central phages as those that are highly abundant and infect many bacteria 128 which themselves are abundant and infected by many other phages. Bacterial centrality is defined in the same way. This metric allowed us to identify those bacteria and phage hubs that were capable of broadly 130 disseminating genetic material and most significantly impact community dynamics through fluctuations in abundance. Having more microbial hubs is also associated with greater stability and resilience to perturbations 132 such as extinction events, which could disintegrate a poorly connected network. We compared the centrality profiles between graphs using the Bray-Curtis metric which accounts for shared bacteria and phages, as 134 well as their degree of weighted centrality. The final result was a robust measurement of graph structure 135 dissimilarity that accounted for abundance, centrality, and shared community membership.

We found that gut microbiome network structure was highly individual specific, with networks clustering tightly based on the human host (Figure 2 A). Intrapersonal (within person) network dissimilarity over the 8-10 day sampling period was significantly less than the average interpersonal dissimilarity (between people) associated with each individual (Figure 2 B). The probability of intrapersonal diversity of an individual being less than interpersonal diversity in our dataset was 98% (absolute error < 1.1e-4) (Figure 2 B).

42 Unlike the strong intrapersonality of the gut, skin network structure was weakly intrapersonally conserved.

The probability of a given skin site being more similar to itself after a month compared to a different person at the same time was 83% (absolute error < 7.5e-6) (Figure 2 C). This distribution was similar for each separate anatomical site, supporting this as an accurate representation of skin intrapersonal and interpersonal diversity (Supplemental Figure S6).

The individuality of gut network structures did not extend to families. The gut network structures were no more similar between twins and their mothers (intrafamily) compared to other families of twins and families (inter-family) (Figure 2 D). The probability of a family member being more similar to another family member compared to non-family members was 63% (absolute error < 1.1e-4).

Role of Diet & Obesity in Gut Network Structure

Diet is a major environmental factor that influences resource availability and gut microbiome composition and diversity, including bacteria and phages^{13,40,41}. Additional work in isolated culture-based systems has suggested that changes in nutrient availability are associated with altered phage - bacteria network structures, although this has yet to be applied to humans²¹. We therefore hypothesized that network structure would be altered by changes in diet. We additionally evaluated the potential association between gut network structure and obesity, a disease linked to diet and potentially the microbiome⁴².

We evaluated the differences in gut network structure by quantifying graph connectedness as the overall degree 158 of network centrality. We accomplished this by utilizing two common centrality metrics: degree centrality 159 and closeness centrality. Degree centrality, the simplest centrality metric, was defined as the number of connections each phage made to bacteria, or each bacterium made to phages. Said another way, degree 161 centrality was defined as the number of relationships associated with each bacterium or phage. Because this metric alone offers only minimal insight, we supplemented it with measurements of closeness centrality. 163 Closeness centrality is a measure of how close each phage or bacterium is to all of the other phages and bacteria in the network. A higher closeness centrality suggests genetic information or the effects of altered 165 abundance would be more more impactful to all other microbes in the system. Because these values are assigned to each phage and bacterium within each network, we calculated the average connectedness and 167 corrected for the maximum potential degree of connectedness to obtain a single value for the connectedness of each graph. 169

We found that gut microbiome network structures associated with high fat diets were less connected than
those of low fat diets (Figure 3 A-B). Tests for statistical differences were not performed, so as to prevent
misleading interpretations from the small sample size. High fat diets exhibited less degree centrality (Figure

3 A), meaning bacteria were overall targeted by less phages and phage tropism was more specific. High fat
diets also exhibited decreased closeness centrality (Figure 3 B), meaning the microbes were more distant
from other microbes in the community, making information transfer and the impact of altered abundance less
likely to impact other bacteria and phages within the network.

In addition to diet, there was also an association between obesity and network structure (Figure 3 C-D).

The obesity-associated network was associated with a higher degree centrality (Figure 3 C) but a less
closeness centrality, compared to the healthy controls (Figure 3 D). This means that the obesity network
was overall less connected, with more relationships per bacteria and phage, but microbes being further from
all other microbes within the community.

Variation of Network Structure Across the Human Skin Landscape

Extensive previous work has shown differences in microbial communities between anatomical sites, including bacteria, viruses, and fungi^{12,38,43}. These communities vary by degree of skin moisture, oil, and environmental exposure. We hypothesized that like microbial composition and diversity, microbial network structure differs between anatomical sites. We addressed this hypothesis by evaluating the changes in network structure between anatomical sites within our skin dataset.

We quantified the average centrality of each sample using the weighted Eigenvector centrality metric, which
provides a higher value for more abundant phages and bacteria that have more relationships to other bacteria
and phages that themselves have more relationships. We found that intermittently moist skin sites (dynamic
sites that fluctuate between being moist and dry) were significantly less connected than the more stable
moist and sebaceous environments (Figure 4 A). We also found that skin sites that were protected from
the environment (occluded) were much more highly connected than those that were constantly exposed to
the environment or intermittently occluded (Figure 4 B).

We supplemented this analysis by comparing the network signatures using the centrality dissimilarity approach described above for our intrapersonal diversity analysis. The dissimilarity between samples was a function of shared relationships, degree of centrality, and bacteria/phage abundance. When using this more sophisticated approach, we found that network structures significantly clustered by moisture, sebaceous, and intermittently moist status (Figure 4 C,E). We also found that occluded sites were significantly different from exposed and intermittently occluded sites, but there was no difference between exposed and intermittently occluded sites (Figure 4 D,F).

Discussion

We developed and implemented a network-based method to understanding the relationships between bacteria and bacteriophages across the human microbiome. This network approach was advantageous over previous studies because it did not rely on inappropriate linear correlations of abundance, it was trained on positive and negative relationships, and it involved implementation of network theory concepts. By utilizing links between bacteria and phage community members, we gained new insights into communication pathways throughout the communities, including potential routes of horizontal gene transfer, the influence of fluctuating abundance, and overall community stability and resilience to extinction events.

We used this approach to build off of previous human microbiome and virome work and show that the
structural properties of bacteria and phage community relationships differ based on environmental conditions.
We found that, just as gut virome composition and diversity are conserved in individuals, gut network
structure was also conserved within people over time. This conservation did not extend to families, as gut
network structures were just as similar to each other as they were to members of other families.

While the gut networks were highly conserved within individuals, the network structure of the skin was weakly 215 conserved. This weaker conservation in the skin compared to the gut could have been due to the greater 216 exposure of the skin to the external environment. Comparison of the networks by skin micro-environment, 217 including moisture levels and environmental exposure, revealed a significant difference in network structures 218 between anatomical micro-environments. The more dynamic regions, including the exposed, intermittently 219 moist, and intermittently occluded sites, exhibited decreased connectedness compared to the more stable 220 micro-environments. Thus the protected and stable environments harbored more stable microbiome networks 221 that were amenable to broad horizontal gene transfer and resilience to network disruption and disintegration. This link between network structure and micro-environment also suggests a role for nutrient availability in 223 defining network structure.

Because diet has been linked to alterations in gut bacterial and viral communities, and because nutrient
availability has been associated with altered network structures in vitro, we evaluated the association of high
and low fat diets with microbiome network structure. Although our sample size was small, we found evidence
that there is indeed a difference in network structure between high and low fat diets. We also observed a
difference in network structure between healthy and obese individuals, with obesity being a disease associated
with altered nutrient availability. Although the obese individual's gut microbiome network had nodes with
more relationships, it was associated with reduced connectedness similar to what was observed in high fat
diets. Thus the food we eat may not only impact what microbes colonize our guts, but may also impact their

233 ability to communicate and remain stable.

Together these findings suggest that human microbiome network structure and stability depends on the
environmental conditions. More specifically, exposed and fluctuating micro-environments seem to be associated
with microbial communities less resilient to perturbations and with less capacity for intra-community
communication. Nutrient availability also appears to play a role in these community traits.

This work represents an initial step toward understanding the microbiome through its relationships, in addition
to its membership. While these findings are informative, there are certainly caveats and future directions that
are beyond the scope of this initial study. First, while our infection classification model is advantageous over
existing models, we recognize that, like most classification models, there remains opportunity for improvement.
For example, such a model is only as good as its training set, and future large-scale endeavors into infectious
relationships (and the associated genomes) will provide more robust training and higher model accuracy. New
and creative scoring metrics can also be integrated into this model to further improve model performance.

Second, while informative, this work was done retrospectively and relied on published research from as much many years ago. These archived datasets were limited by the technology and associated costs of the time, meaning the datasets are poorly powered for the statistical analysis we strive for today. While we were able to present initial observations, follow-up studies will certainly be required to validate our observations.

Overall these findings and methodologies are exciting because they represent a new way of understanding
the human microbiome. We expect this work will contribute toward understanding the human microbiota
as complex interacting communities instead of the reductionist approach often utilized today. Our findings
confirm that, in addition to community membership and diversity, microbiome interactions differ between
human environments. Some skin sites, diets, and obesity are associated with less connected communities,
which could impact gene transfer and community stability. Other highly connected communities are expected
to be more capable of transferring genetic material, such as antibiotic resistance genes. Finally, the link
between network structure and nutrient availability could be important for more effective design of probiotics,
fecal microbiota transplants, and other microbiome-based therapeutics.

$_{258}$ Materials & Methods

Data Availability

All associated source code is available on GitHub at the following repository:

https://github.com/SchlossLab/Hannigan-2016-ConjunctisViribus.

Data Acquisition & Quality Control

Raw sequencing data and associated metadata was acquired from the NCBI sequence read archive (SRA).

Supplementary metadata was acquired from the same SRA repositories and their associated manuscripts.

The gut virome diet study (SRA: SRP002424), twin virome studies (SRA: SRP002523; SRP000319), and skin virome study (SRA: SRP049645) were downloaded as .sra files. Sequencing files were converted to fastq format using the fastq-dump tool within the NCBI SRA Toolkit (version). Sequences were quality trimmed using the Fastx toolkit to exclude bases with quality scores below 33 and shorter than 75 bp. Paired end reads were filtered to exclude and sequences missing their corresponding pair using the get_trimmed_pairs.py

271 Contig Assembly

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available in the source code.

Contigs were assembled using the Megahit assembly program (*version*). A minimum contig length of 1 kb was used. Iterative k-mer stepping began at a minimum length of 21 and progressed by 20 until 101. All other default parameters were used.

275 Contig Abundance Calculations

Contigs were concatenated into two master files prior to alignment, one for bacterial contigs and one for phage contigs. Sample sequences were aligned to phage or bacteria contigs using the Bowtie2 global aligner (version). We defined a mismatch threshold of 1 bp and seed length of 25 bp. Sequence abundance was calculated from the Bowtie2 output using the calculate_abundance_from_sam.pl script available in the source code.

Operational Genomic Unit Binning

Contigs often represent large fragments of genomes, due to insufficient sequencing depth. In order to reduce redundancy and artificially inflated genomic richness within our dataset, it was important to bin contigs based on their likelihood to be of the same phylogenetic groups. This approach is conceptually similar to the clustering of related 16S rRNA sequences into operational taxonomic units (OTUs), although here we are clustering contigs into operational genomic units (OGUs).

We clustered contigs using the CONCOCT algorithm (*version*). Because of our large dataset and barriers in computational efficiency, we randomly subsampled the dataset to include 25% of all samples, and used these to inform contig abundance within the CONCOCT algorithm. CONCOCT was used with a maximum of 500 clusters, a k-mer length of four, a length threshold of 1 kb, 25 iterations, and exclusion of the total coverage variable.

OGU abundance (Ao) was obtained as the sum of the abundance of each contig (Aj) associated with that
OGU. The abundance values were length corrected such that:

$$A_O = \frac{10^7 \sum_{j=1}^k A_j}{\sum_{j=1}^k L_j}$$

Where L is the length of each contig j within the OGU.

²⁹⁵ Open Reading Frame Prediction

Open reading frames (ORFs) were identified using the Prodigal program (version) with the meta mode parameter and default settings.

²⁹⁸ Classification Model Creation and Validation

The classification model for predicting interactions was built using experimentally validated bacteria-phage infections or validated lack of infections from six studies^{31,33–37}. Associated reference genomes were downloaded from the European Bioinformatics Institute. The model was created based on the four metrics listed below.

The four scores were used as parameters in a random forest model to classify bacteria and bacteriophage pairs as either having infectious interactions or not. The classification model was built using the Caret R package.

The model was trained using five-fold cross validation with ten repeats. Pairs without scores were classified as not interacting. The model was optimized using the ROC value. The resulting model performance was plotted using the plotROC R package.

307 Identify Bacterial CRISPRs Targeting Phages

CRISPRs were identified from bacterial genomes using the PilerCR program (version). Resulting spacer sequences were filtered to exclude spacers shorter than 20 bp and longer than 65 bp. Spacer sequences were

aligned to the phage genomes using the nucleotide Blast algorithm with default parameters. The mean percent identity for each matching pair was recorded for use in our classification model.

312 Detect Matching Prophages within Bacterial Genomes

Temperate bacteriophages infect and integrate into their bacterial host's genome. We detected integrated
phage elements within bacterial genomes by aligning phage genomes to bacterial genomes using the nucleotide
Blast algorithm and a minimum e-value of 1e-10. The resulting bitscore of each alignment was recorded for
use in our classification model.

317 Identify Shared Genes Between Bacteria and Phages

Phages may share genes with their bacterial hosts, providing us with evidence of phage-host infectious pairs.

We identified shared genes between bacterial and phage genomes by assessing amino acid similarity between
the genes using the Diamond protein alignment algorithm. The mean alignment bitscores for each genome
pair was recorded for use in our classification model.

322 Protein - Protein Interactions

The final method we used for predicting infectious interactions between bacteria and phages was by detecting
pairs of genes whose proteins are known to interact. We assigned bacterial and phage genes to protein
families by aligning them to the Pfam database using the Diamond protein alignment algorithm. We then
identified which pairs of proteins are predicted to interact using the Pfam interaction information within the
Interact database (version). The mean bitscores of the matches between each pair were recorded for use in
our classification model.

329 Virome Network Construction

The bacteria and phage operational genomic units (OGUs) were scored using the same approach as outlined above. The infectious pairings between bacteria and phage OGUs were classified using the random forest model described above. The predicted infectious pairings and all associated metadata was saved as a graph database using Neo4j software. This network was used for downstream community analysis.

334 Centrality Analysis

- Degree and closeness centrality were calculated using the associated functions within the igraph R package.
- Briefly, the closeness centrality of Vi was calculated taking the inverse of the average length of the shortest
- path (d) between nodes Vi and the k other nodes Vj, such that closeness centrality of node Vi is:

$$C_{C}\left(V_{i}\right) = \left(\sum_{j=1}^{k} d\left(V_{i}, V_{j}\right)\right)^{-1}$$

- The distance between nodes (d) was calculated as the shortest number of edges required to be traversed to move from one node to another.
- The simple metric of **degree centrality** of node Vi was defined as the sum of the number of k edges Ei associated with that node, such that:

$$C_D\left(V_i\right) = \sum_{i=1}^k E_i$$

- The Eigenvector centrality was calculated using the values of the first eigen vector of the associated adjacency matrix that are associated with each vertex Vi. Conceptually, this function results in a centrality value that reflects the connections of the vertex, as well as the centrality of the connected vertices.
- The **centralization** metric was used to assess the average centrality of each sample graph G. Centralization was calculated by taking the sum of each vertex Vi centrality from the graph maximum centrality Cw, such that:

$$C(G) = \frac{\sum_{i=1}^{k} \max_{w} c(w) - c(v_i)}{T}$$

The values were corrected for uneven graph sizes by dividing the centralization score by the maximum theoretical centralization (T) for a graph with the same number of vertices.

350 Network Relationship Dissimilarity

We assessed similarity between graphs by evaluating the shared centrality of their vertices, as has been done previously. More specifically, we calculated the dissimilarity between graphs Gi and Gj using the Bray-Curtis dissimilarity metric and Eigenvector centrality values such that:

$$B\left(G_{i},G_{j}\right)=1-\frac{2C_{ij}}{C_{i}+C_{j}}$$

- Where Cij is the sum of the lesser centrality values for those vertices shared between graphs, and Ci and Cj
- are are the total number of vertices found in each graph. This allows us calculate the dissimilarity between
- graphs based on the shared centrality values between the two graphs.

357 Acknowledgments

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Figures 560

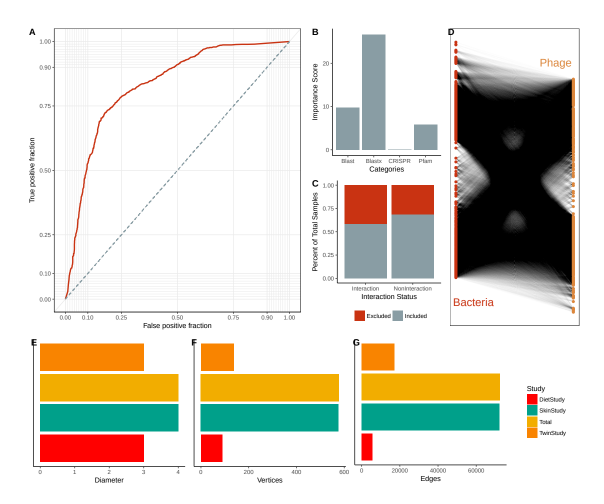


Figure 1: Summary of Multi-Study Network Model. (A) ROC curve resulting from the ten iterations used to create the phage - bacteria infection prediction model. (B) Importance scores associated with the metrics used in the random forest model to predict relationship between bacteria and phages. The importance score is the mean decrease in accuracy of the model when the model is rebuilt without that factor. (C) Proportions of samples excluded from model learning due to a lack of scores. The true interaction status of the sample is noted on the x-axis and bars are colored by the proportion of sample excluded (red) and included (grey) in model training. (D) Bipartite visualization of the resulting phage-bacteria network. This network includes information from all three published studies. (E) Network diameter (measure of graph size; the greatest number of traversed vertices required between two vertices), (F) number of vertices, and (G) number of edges (relationships) for the total network (yellow) and the individual study sub-networks (diet study = red, skin study = green, twin study (orange).

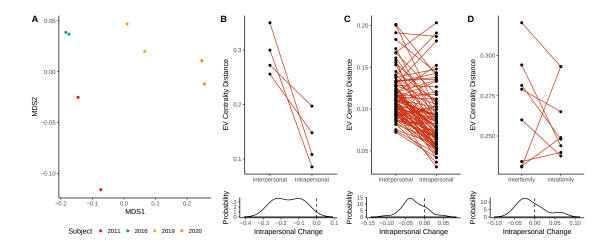


Figure 2: Intrapersonal vs Interpersonal Network Dissimilarity Across Different Human Systems. (A) NMDS ordination illustrating network dissimilarity between subjects over time. Each sample is colored by subject, with each sample pair collected 8-10 days apart. Dissimilarity calculated using the Bray-Curtis metric based on abundance weighted Eigenvector centrality signatures, with a greater distance representing greater dissimilarity in bacteria and phage centrality and abundance. (B) Quantification of gut network dissimilarity within the same subject over time (intrapersonal) and the mean dissimilarity between the subject of interest and all other subjects (interpersonal). Below is the probability distribution of the changes between intrapersonal and interpersonal diversity, representing the probability that intrapersonal dissimilarity will be lower than interpersonal dissimilarity (intrapersonal change less than zero). (C) Quantification of skin network dissimilarity within the same subject and anatomical location over time (intrapersonal) and the mean dissimilarity between the subject of interest and all other subjects at the same time and the same anatomical location (interpersonal). Probability distribution the same as for pannel B. (D) Quantification of gut network dissimilarity within subjects from the same family (intrafamily) and the mean dissimilarity between subjects within a familiy and those of other families (interfamily). Probability distribution the same as for pannel B.

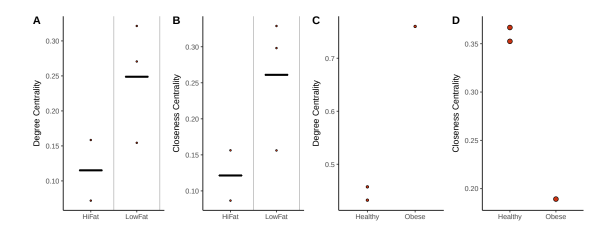


Figure 3: Impact of Diet and Obesity on Gut Network Structure. (A) Quantification of average degree centrality (number of edges per node) and (B) closeness centrality (average distance from each node to every other node) of gut microbiome networks of subjects limited to exclusively high fat or low fat diets. Lines represet the mean degree of centrality for each diet. (C) Quantification of average degree centrality and (D) closeness centrality between obese and healthy adult women. Tests for statistical significance were not performed as they would be misleadingly inappropriate for these small sample sizes.

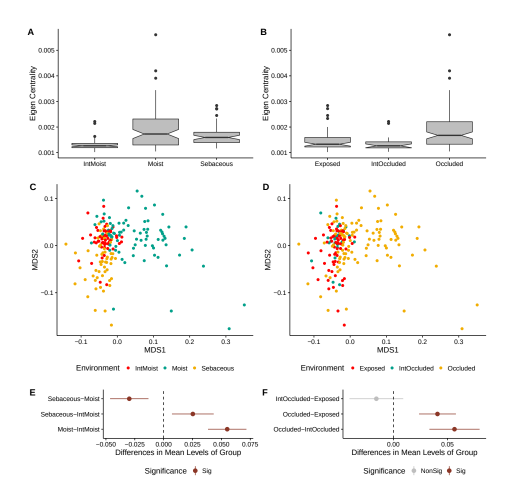


Figure 4: Impact of Skin Microenvironment on Microbiome Network Structure. (A) Notched boxplot depicting differences in average Eigenvector centrality between moist, intermittently moist, and sebaceous skin sites and (B) occluded, intermittently occluded, and exposed sites. Notched boxplots were created using ggplot2 and show the median (center line), the inter-quartile range (IQR; upper and lower boxes), the highest and lowest value within 1.5 IQR (whiskers), and the notch which provides an approximate 95% confidence interval as defined by 1.58 * IQR / sqrt(n). (C) NMDS ordination depicting the differences in skin microbiome network structure between skin moisture levels and (D) occlusion. Samples are colored by their environment and their dissimilarity to other samples was calculated as described in figure 2. (E) The statistical differences of networks between moisture and (F) occlusion status were quantified with an anova and post hoc Tukey test. Cluster centroids are represented by dots and the extended lines represent the assocaited 95% confidence intervals. Significant comparisons (p-value < 0.05) are colored in red, and non-significant comparisons are gray.*

361 Supplemental Figures

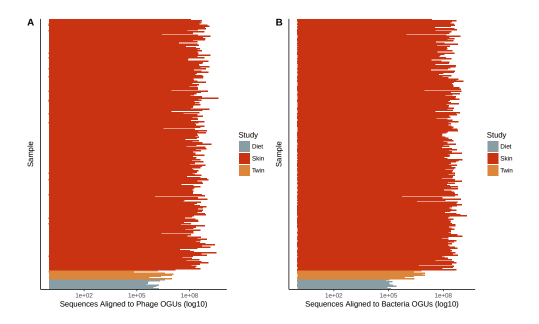


Figure S1: **Sequencing Depth Summary.** Number of sequences that aligned to (A) Phage and (B) Bacteria operational genomic units. Sequencing count aligned to OGUs was length corrected.

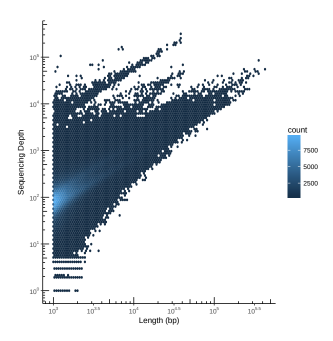


Figure S2: Contig Summary Statistics. Scatter plot heat map with each hexagon representing an abundance of contigs. Contigs are organized by length on the x-axis and the number of aligned sequences on the y-axis.

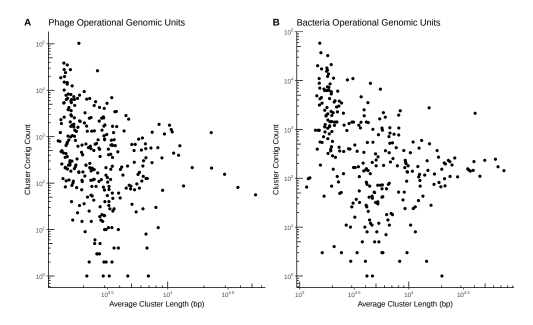


Figure S3: Operational Genomic Unit Summary Statistics. Scatter plot with operational genomic unit clusters organized by average contig length within the cluster on the x-axis and the number of contigs in the cluster on the y-axis. Operational genomic units of (A) bacteriophages and (B) bacteria are shown.

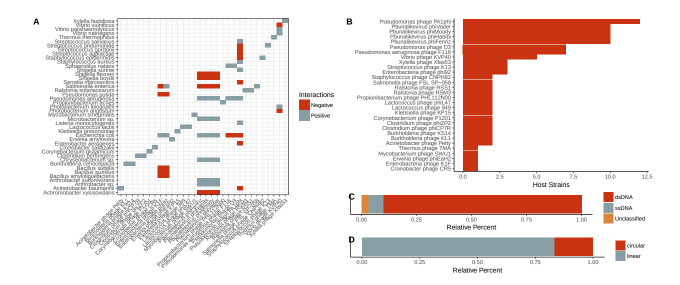


Figure S4: Summary information of validation dataset used in the interaction predictive model. A) Categorical heat-map highlighting the experimentally validated positive and negative interactions. Only bacteria species are shown, which represent multiple reference strains. Phages are labeled on the x-axis and bacteria are labeled on the y-axis. B) World map illustrating the sampling locations used in the study (red dots). C) Quantification of bacterial host strains known to exist for each phage. D) Genome strandedness and E) linearity of the phage reference genomes used for the dataset.

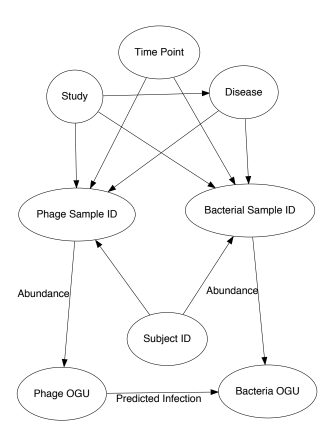


Figure S5: **Structure of the interactive network.** Metadata relationships to samples (Phage Sample ID and Bacteria Sample ID) included the associated time point, the study, the subject the sample was taken from, and the associated disease. Infectious interactions were recorded between phage and bacteria operational genomic units (OGUs). Sequence count abundance for each OGU within each sample was also recorded.

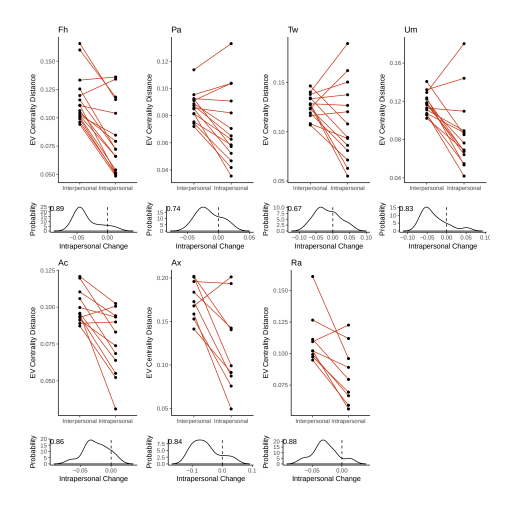


Figure S6: Intrapersonal vs Interpersonal Dissimilarity of the Skin. Quantification of skin network dissimilarity within the same subject and anatomical location over time (intrapersonal) and the mean dissimilarity between the subject of interest and all other subjects at the same time and the same anatomical location (interpersonal), separated by each anatomical site (forehead [Fh], palm [Pa], toe web [Tw], umbilicus [Um], antecubital fossa [Ac], axilla [Ax], and retroauricular crease [Ra]). Below is the probability distribution of the changes between intrapersonal and interpersonal diversity, representing the probability that intrapersonal dissimilarity will be lower than interpersonal dissimilarity (intrapersonal change less than zero). The probability that the slope will be less than zero (integral from negative infinity to zero) is provided in the top left corner.

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