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## Metagenome analysis using the Kraken software suite

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### Abstract

In order to facilitate efficient and reproducible metagenomic analysis, we introduce Kraken Protocols, an end-to-end pipeline for the classification, quantification, and visualization of metagenomic datasets. Our protocol describes the execution of the Kraken programs, via a sequence of easy to use scripts, in two scenarios: (1) quantification of the species in a metagenomics sample, and (2) detection of a pathogenic agent from a clinical sample taken from a human patient. The protocols can be run by any users who are familiar with the Unix command-line environment.

### Introduction

Metagenomics sequencing has greatly improved our understanding of the microscopic world by revealing a vast range of microbial organisms that were previously unobserved, many of which cannot be grown in laboratory cultures<sup>1</sup>. Metagenomics experiments take many forms, two of which can be broadly categorized as either microbiome experiments or pathogen identification experiments. In microbiome experiments, researchers evaluate all of the microbial organisms identified in a given sample, often with the goal of describing what is present. In contrast, in a pathogen identification experiment, researchers focus on

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#### Author Contributions

J.L. and M.S. led the development of the protocol. N.R. executed and designed the Microbiome Analysis protocol and is the author of the KrakenTools alpha diversity tools. J.L. developed the Pathogen Identification protocol and is the author of Bracken and KrakenTools. M.S. authored the jupyter notebooks for the protocol. D.E.W. is the senior author of Kraken and Kraken 2. F.B. is the author of KrakenUniq. C.P. is an author for the KrakenTools beta diversity script. B.L. supervised the development of Kraken 2. S.L.S. supervised the development of Kraken, KrakenUniq, and Bracken. B.L. and S.L.S. supervised the development of this protocol. All authors contributed to the writing of the manuscript.

#### Competing interests

The authors declare no competing financial interest

identifying one or a few pathogenic microbes, with the goal of diagnosing an infection. The suite of tools in the Kraken package were developed to cover many of the bioinformatics needs of both microbiome and pathogen metagenomics experiments. These tools include Kraken<sup>2</sup>, KrakenUniq<sup>3</sup>, Kraken 2<sup>4</sup>, Kraken2Uniq (based on KrakenUniq), Bracken<sup>5</sup>, KrakenTools, and Pavian<sup>6</sup>.

Microbiome experiments begin with (1) removing host DNA and then (2) classifying a set of sequencing reads, with each read assigned to a taxonomic category (species, genus, or higher-level taxa), followed by (3) computing the relative abundance of different species in the sample. When computing relative abundance, researchers sometimes focus on a limited number of “marker” genes, classifying only the reads that align within these genes. The most widely used marker gene is ribosomal RNA, but protein-coding genes can also be used, particularly those that are expected to be present in exactly one copy per cell. However, many studies prefer to analyze all of the sequencing reads collected from a sample, which is the strategy we focus on in this protocol. Following characterization of the original data, downstream analyses may include (4) statistical methods for comparing the microbial compositions of different environments or (5) visualization methods for understanding microbial compositions. Kraken<sup>2</sup> and Kraken 2<sup>4</sup> (an improved version of Kraken) were previously developed by some of us for rapid, accurate classification of sequencing reads. Bracken<sup>5</sup> was developed to work in conjunction with Kraken to compute species abundance using Kraken classification results. Finally, KrakenTools and Pavian<sup>6</sup> provide a comprehensive set of tools for downstream statistical analysis and visualization of the classification and abundance estimation results.

In pathogen identification experiments, researchers and clinicians are interested in identifying pathogenic microbes that might be the cause of a harmful infection. Pathogen identification studies conceptually include the following steps: (1) removal of host DNA from the microbial reads; (2) classification of the remaining microbial reads; (3) comparison of sample reads against control samples; and (4) validation of pathogen classifications. The second and third steps are essentially the same as those done for microbiome experiments, while the first and last steps are needed to ensure the accuracy of any pathogen that is identified. For removal of host DNA, one can use Bowtie 2<sup>7</sup> (also developed by some of the authors of this protocol) as a fast, sensitive aligner that can compare sequencing reads to the human reference genome. Classification of the remaining reads is then accomplished by using KrakenUniq<sup>3</sup> and Kraken2Uniq. Finally, KrakenTools and Pavian<sup>6</sup> cover the remaining steps of pathogen identification experiments, assisting in the identification and verification of potential pathogens.

Here, we describe in detail how to use the Kraken suite to analyze metagenomic data for both microbiome and pathogen identification experiments. Figure 1 outlines the protocols for each of the two types of studies.

## Overview of the protocol

The protocols described here focus on two major categories of metagenomic experiments: microbiome analysis and pathogen identification. We detail the protocol steps using

DNA samples from the human gut and cornea. However, these protocols may also be applied to RNA data as well as other biomes and species. For microbiome analysis, we consider an experiment describing the change in gut microbiome for a single human patient who underwent a fecal microbiota transplant and antibiotic treatment. For pathogen identification, we consider 8 separate human corneal samples with bacterial, viral, or fungal infections and two non-infectious human corneal samples. For both samples, human reads were already removed prior to analysis in the protocol. However, we include the host removal step in the protocol to demonstrate how to remove host reads using Bowtie 2<sup>7</sup>.

For microbiome analysis, host-filtered microbial reads undergo classification against bacterial, viral, fungal, archaeal, and human genomes using Kraken 2 (Figure 1). The classification report is then used by Bracken for species abundance estimation, which provides estimated reads per species in the sample. Bracken output files are then passed to KrakenTools and Pavian for visualization. Pavian provides Sankey visualization of samples and read count comparisons between samples. For more in-depth downstream analysis, we recently developed KrakenTools (<https://github.com/jenniferlu717/KrakenTools>) as a set of individual programs for visualizing or transforming Kraken and Bracken output. For microbiome projects, KrakenTools provides functions to compute Krona plots, filter and combine reports, and calculate alpha and beta diversity metrics about each sample. The software is not previously published but is already packaged and available through github or bioconda (<https://bioconda.github.io/recipes/krakentools/README.html>).

For pathogen identification, we classify 8 infected patient samples and 2 control samples against the same Kraken 2 database used in microbiome analysis (Figure 1). However, the additional feature of unique k-mer counting from KrakenUniq<sup>3</sup> is enabled for accurate pathogen identification. We refer to this as Kraken2Uniq. Following classification by Kraken2Uniq, the Kraken report files are uploaded to the Pavian Shiny App<sup>6</sup>, which compiles all of the read counts per species and allows between sample visualization and comparison. The Pavian app then calculates z-scores between read counts, with higher z-scores resulting from samples with higher than usual read counts for a given species. We then sort the classified species by z-score, with the highest z-score species for each sample being the most likely pathogen. The filtered table can then be saved as a tab-delimited table that can undergo further visualization. Finally, pathogen identification can be validated using the extract kraken reads.py script from KrakenTools. The KrakenTools script allows users to extract classified reads and confirm potential pathogens using other tools such as BLAST<sup>8</sup> or Bowtie 2<sup>7</sup>.

The target audience for this protocol is biologists and clinicians working in microbiome or metagenomics analysis. This protocol does not require programming expertise, but it does assume familiarity with the Unix command-line interface. Users should be comfortable running programs from the command line in the Unix environment.

## Differences between Kraken methods

There are currently four major versions of the Kraken software: (1) Kraken<sup>2</sup>, (2) KrakenUniq<sup>3</sup>, (3) Kraken 2<sup>4</sup> and (4) Kraken2Uniq. All four versions are based on the same

classification algorithm which uses exact-matching of read k-mers against a Kraken database of k-mers from existing genomes. However, the methods differ in the ways they count, access and store the k-mer information, with each version improving upon the previous Kraken version.

Kraken and KrakenUniq apply the same k-mer retrieval strategy, and both tools can use the same databases. The reference database needs roughly 12 bytes per k-mer. Building and storing the Kraken index for a thousand distinct bacterial genomes of length 4Mb would result in an index of size  $\approx 45\text{ GB}$ .

However, the RefSeq<sup>9;10</sup> database already contains  $\sim 64K$  bacterial species, and trying to store all k-mers from this ever-growing database became the main bottleneck of Kraken and KrakenUniq. Therefore, Kraken 2 was developed as a much more efficient version of Kraken, reducing the memory usage by 85% over Kraken and KrakenUniq. In addition to the reduction in database size, Kraken 2 also runs about 5 times faster than Kraken/KrakenUniq.

Another distinguishing difference is in how the Kraken methods count reads. While Kraken and Kraken 2 provide cumulative statistics of the total read count per taxa, KrakenUniq and Kraken2Uniq additionally count and report the number of unique k-mers per taxon, using an efficient HyperLogLog implementation. In other words, for each species in the output, KrakenUniq will report how many distinct k-mers from that species were observed in the reads. Unique counts are especially useful for pathogen detection to distinguish real from spurious signals.

## Alternative analysis packages

The Kraken/KrakenUniq tools assign every read to a taxon of origin. Other software tools solve this problem, including CLARK<sup>11</sup>, Centrifuge<sup>12</sup>, and Kaiju<sup>13</sup>. Kraken 2 generally exhibits a more advantageous combination of speed, memory footprint, and accuracy compared to those tools<sup>4</sup>, but it is still possible to substitute those tools into earlier steps in this protocol. Recent benchmarking studies<sup>14, 15</sup> evaluated multiple metagenomic classifiers and found that Kraken, Kraken 2, and Bracken were among the best-performing methods.

Other tools seek to reduce the memory footprint of the index by including only the portions of the genome sequences that can distinguish between taxa, e.g. MetaPhlAn2<sup>16</sup>. This reduces the size of the index, but prevents the tool from classifying many of the reads. So, while these tools can be used in a microbiome analysis like the one described in this protocol, they are not appropriate for pathogen identification.

While approaches based on large-scale machine learning have been shown to have advantages in certain scenarios, e.g. for classifying reads from species that are absent from the database<sup>16</sup>, these approaches have tended to be computationally outperformed by classifiers that assign reads directly based on sequence content, without the need to first train a model<sup>17;18</sup>. For more details on competing paradigms for read classification, see Breitwieser et al.<sup>19</sup>.

The Bowtie 2 tool used to filter host reads can be replaced with similar read alignment tools such as BWA-MEM or minimap2.

## Limitations

### Removal of host reads.

For metagenomic studies, removal of host (usually human) DNA is critical: either as a pre-processing step, using a program such as Bowtie 2<sup>7</sup> to align all of the reads to the host, or by including the host genome in the Kraken database. The pre-processing strategy may be more effective because external alignment programs such as Bowtie 2 are more sensitive than Kraken, and because pre-processing yields a greatly reduced set of reads, making subsequent steps faster. Bowtie 2 and other alignment programs are suitable for the pre-processing step because they can identify and remove any reads belonging to a known host (e.g., human), while Kraken can then identify the wide variety of unknown microbial species in the remainder of the sample.

However, this pre-processing step may cause sequences of interest to be removed from the sample set, especially in the rare case of viral sequences found to be similar to a human endogenous retrovirus. In this protocol, we include the pre-processing with Bowtie 2 as an optional step that may or may not be included at the discretion of the users of the protocol.

### Reference Database.

Kraken 2's classification sensitivity and specificity highly depend on how (1) complete and (2) accurate the used reference database is. (1) The entire microbial biosphere is still far from being completely sequenced and thus missing or partial genomes may introduce biases and reduce classification performance. Reads without a reference in the database will be labeled as unknown or imprecisely assigned to the next closest taxon. (2) Additionally, genomes can also be contaminated with DNA from other organisms<sup>20;21</sup> introduced during sequencing. Contamination causes Kraken to assign wrong taxonomic labels to reads. While this kind of classification error affects only a small fraction of reads, it can make the detection of weak signals difficult.

For the purpose of this protocol, we use the complete genomes from RefSeq bacteria, archaea, viral, and plasmid libraries along with extensively screened eukaryotic pathogen genomes<sup>22</sup>. We use only complete genomes to avoid potential contamination from draft genomes.

For users that may have limited RAM, Kraken 2 provides the ability to generate a minikraken database. Minikraken databases compress the full Kraken 2 database by saving a subset, but representative set of the database k-mers. To allow all users to execute this protocol, we use an 8Gb minikraken database.

Alternatively, Kraken 2 can also classify reads against protein databases using six-frame-translation. Protein-level classification is more sensitive (but less specific) and therefore can help to reduce the missing reference bias. For even longer evolutionary distances, homology

detection alignment-based classification methods like DIAMOND<sup>23</sup> or MMseqs2<sup>24</sup> can be used.

Other issues can arise as reference databases grow and as the distribution of reference genomes per taxon becomes more lopsided. For example, Nasko et al.<sup>25</sup> showed that as the RefSeq database has grown over time, Kraken has tended to assign a greater proportion of reads to higher levels of the taxonomy. In particular, they showed that a greater proportion of reads tend to be assigned to the genus level, at the expense of assignments to the species level and below. As public databases of reference genomes continue to grow, it will be important to continue to evaluate how classification results are affected by large or lopsided numbers of genomes per taxon.

### Long read classification

This protocol is designed for Illumina sequencing reads, which are highly accurate (error rates under 0.5%) but short, usually 100–250bp. However, in some cases, users may want to evaluate longer, higher error-rate reads from Pacific Biosciences or Oxford Nanopore. For longer reads with a different error profile, users might remove host DNA using minimap2<sup>26</sup>, and they might also need to adjust the parameters of the Kraken software. Specifically, higher error rate reads, such as those from Pacific Biosciences or Oxford Nanopore, yield better classification with Kraken databases generated using smaller k-mer and minimizer lengths. Previous analyses have indicated better performance using k-mer and minimizer lengths of 26 (as opposed to the default k-mer length of 35 and minimizer length of 31).

Also the read count measurement might not be as useful for long reads, because these experiments typically generate fewer reads that differ significantly in length. For these experiments, it may make more sense to compare k-mer counts.

### Assembly-based approaches

When a species is present in sufficiently high amounts, one might wish to assemble the reads *de novo* to create large, contiguous sequences from that species. The protocols described here do not perform genome assembly, which requires other software methods.

## Materials

### Equipment

The Kraken 2 protocol website [http://ccb.jhu.edu/data/kraken2\\_protocol/](http://ccb.jhu.edu/data/kraken2_protocol/) summarizes all required software and data, along with details about how to download all required data for the protocol.

- Data (the example corneal reads, microbiome fecal reads, and Kraken 2 database for use in this protocol are available from NCBI SRA and Amazon AWS; see Equipment Setup for details)
- Kraken 2 software (<https://github.com/DerrickWood/kraken2/>, version 2.1.1 or later)

- Bracken software (<https://github.com/jenniferlu717/Bracken>, version 2.6.2 or later)
- KrakenTools software (<https://github.com/jenniferlu717/KrakenTools>, version 1.1. or later)
- Pavian software (<https://github.com/fbreitwieser/pavian>, version 1.0 or later)
- Bowtie 2 (<https://github.com/BenLangmead/bowtie2>, version 2.4.4)
- samtools (<http://www.htslib.org/download/>), version 0.1.20 or later)
- R (<https://www.r-project.org>)
- RStudio (<https://www.rstudio.com/>)
- Hardware (64-bit computer running either Linux or Mac OS X (10.7 Lion or later); 8GB of RAM; see Equipment Setup)

## EQUIPMENT SETUP

### Required data

- The microbiome analysis portion of this protocol is illustrated with a sample microbiome dataset of fecal samples from Xavier et al. (2018)<sup>27</sup>. Many samples from the paper are available at <https://www.ncbi.nlm.nih.gov/bioproject/PRJNA491657>. Here, we use 3 samples from a single patient, T11, who began antibiotic treatment at day 0, underwent stem cell engraftment on day 14, and received autologous fecal microbiota transplantation (auto-FMT) to return the patient's gut microbiota diversity at day 29.
- The infectious disease analysis portion of this protocol is illustrated with selected corneal samples from Li et al. (2018)<sup>28</sup>. All 20 samples from the paper are available at <https://www.ncbi.nlm.nih.gov/bioproject/PRJNA381365>. However, for clarity, we used a select group of 10 of the corneal samples (including 2 controls).
- Both protocols require a Kraken 2 database. For this protocol, we use the same pre-built Kraken 2 database, containing RefSeq complete genomes from December 2020 for bacterial, archaeal, and viral genomes, the Human reference genome GRCh38, and the cleaned eukaryotic pathogen genomes for EuPathDB48<sup>22</sup> (available at <http://ccb.jhu.edu/data/eupathDB/>). Additional pre-built Kraken 2 databases are available in the AWS cloud, detailed here: <https://benlangmead.github.io/aws-indexes/k2>. The following subsection details how to create a Kraken 2 Database if the desired database is not already available.

### Kraken 2 Databases

Several pre-built Kraken 2 databases are available at <https://benlangmead.github.io/aws-indexes/k2>. The most commonly used database is the standard Kraken 2 database (which includes RefSeq archaea, bacteria, viruses, plasmid complete genomes, UniVec Core, and the most recent human reference genome, GRCh38). In addition to the standard database,



we provide a database of the cleaned eukaryotic pathogens<sup>22</sup>, 16S rRNA databases, and expanded standard databases with RefSeq protozoa, fungi, and plant genomes. All databases are updated monthly to include the most recent genomes.

In this protocol, we use an 8GB minikraken database of the combined standard Kraken 2 database with the cleaned eukaryotic pathogen genomes.

If the desired database is not available, we describe here how to create a custom Kraken 2 database using the `kraken2-build` script options:

- First, download the NCBI taxonomy.

```
$ kraken2-build --db krakendb --download-taxonomy
```

- Second, download one or more reference libraries. (The full list of available libraries is at <https://github.com/DerrickWood/kraken2/wiki/Manual#custom-databases>.)

```
$ kraken2-build --db krakendb --download-library bacteria
$ kraken2-build --db krakendb --download-library archaea
$ kraken2-build --db krakendb --download-library viral
$ kraken2-build --db krakendb --download-library protozoa
$ kraken2-build --db krakendb --download-library UniVec_Core
```

- Third, download additional genomes by adding multi-FASTA or single-FASTA files. The FASTA sequence headers must include either 1) NCBI accession numbers or 2) the text `kraken:taxid` followed by the taxonomy ID for the genome (e.g. `>sequence100|kraken:taxid|9606|`). If this requirement is met, the following commands will add the sequences to the database:

```
$ kraken2-build --db krakendb --add-to-library chr1.fa
$ kraken2-build --db krakendb --add-to-library chr2.fa
```

- Finally, build the Kraken 2 database and generate the Bracken database files.

```
$ kraken2-build --db krakendb --build --threads 8
$ bracken-build -d krakendb -t 8 -k 35 -l 100
```

## Downloading and organizing required data

This section details how to organize the database and samples used in this protocol. You should first download the database from [https://genome-index.s3.amazonaws.com/kraken/k2\\_standard\\_eupath\\_20201202.tar.gz](https://genome-index.s3.amazonaws.com/kraken/k2_standard_eupath_20201202.tar.gz) and the samples from NCBI SRA. For details on how to download the SRA samples, see <https://www.ncbi.nlm.nih.gov/sra/docs/srdownload/>.



Table 1 lists the SRA accession IDs for each of the samples used in the Kraken Microbiome Analysis Protocol and the Kraken Pathogen Identification Protocol.

- Create four folders for required data: `k2protocol_db`, `m_samples`, `p_samples`, `b_index`
- Inside the `k2protocol_db` folder, unpack the database:

```
$ tar -xzvf k2_standard_eupath_20201202.tar.gz
```

- Inside the `b_index` folder, download the `k2protocol_bowtie2indices.tgz` file from
- [http://ccb.jhu.edu/data/kraken2\\_protocol/](http://ccb.jhu.edu/data/kraken2_protocol/) and unpack the files:

```
$ tar -xzvf k2protocol_bowtie2indices.tgz
```

- Download the fastq files for SRA samples for the microbiome analysis and for the pathogen identification analysis. Move the three microbiome analysis files into `m_samples` and move the 10 pathogen identification samples into `p_samples`.

### Downloading and installing software

There are two ways to install the required programs: (1) using conda and (2) downloading the binaries. Conda is an open-source package manager that helps to install software. All the software tools mentioned here are packaged in Bioconda<sup>29</sup>, which is a particular repository (“channel”) within conda. (1) To install the software using conda use the following command:

- Install all required using Conda

```
$ conda install -c conda-forge -c bioconda kraken 2 krakentools bracken \
  r bowtie2 samtools
```

- Pavian has no conda package. To run the protocol there are two ways:
- (a) Install it locally. Open the R console and use the following command:

```
$ R
if (!require(remotes)) { install.packages ("remotes")
remotes::install_github("fbreitwieser/pavian")
```

- (b) Use the pavian webserver <https://fbreitwieser.shinyapps.io/pavian/>.

(2) Alternatively all software can be also downloaded as binary using the following commands:

- Create a directory to store all of the executable programs used in this protocol (if none already exists):

```
$ mkdir $HOME/bin
```

- Add the above directory to your PATH environment variable: `$ export PATH=$HOME/bin:$PATH`
- To install Kraken 2, download the latest release from <https://github.com/DerrickWood/kraken2/releases>, unpack the Kraken 2 zip archive, `cd` to the unpacked directory, and run the install script.

```
$ unzip v2.1.1.zip
$ cd kraken2-2.1.1/
$ ./install_kraken 2.sh .
```

- To install Bracken, download Bracken version 2.6.2 from <https://github.com/JenniferLu717/Bracken>, unpack the Bracken zip archive, `cd` to the unpacked directory and run the install script.

```
$ unzip v2.6.2.zip
$ cd Bracken-2.6.2/
$ sh install_bracken.sh
```

- To install KrakenTools, download Kraken 2 version 2.1.1 from <https://github.com/JenniferLu717/KrakenTools> and unpack the KrakenTools zip archive.

```
$ unzip v1.2
```

- Copy the kraken executables to a directory in your PATH and create symlinks for the Bracken and KrakenTools executables.

```
$ cp kraken2-2.1.1/kraken2 $HOME/bin
$ cp kraken2-2.1.1/kraken2 -build $HOME/bin
$ cp kraken2-2.1.1/kraken2 -inspect $HOME/bin
$ ln -s Bracken-2.6.2/bracken $HOME/bin/bracken
$ ln -s Bracken-2.6.2/bracken -build $HOME/bin/bracken-build
$ ln -s KrakenTools $HOME/bin/KrakenTools
```

- Download Bowtie 2 version 2.4.4 (x86 64) from <https://github.com/BenLangmead/bowtie2/releases/>

```
$ unzip bowtie2 -2.4.4-linux-x86_64.zip
$ cp bowtie2-2.4.4-linux-x86_64/bowtie2* $HOME/bin
```

- To install Pavian, open RStudio or R and run the following:

```
if (!require(remotes)) { install.packages ("remotes") }
remotes::install_github("fbreitwieser/pavian")
```

## Procedure

There are two protocols included: Microbiome Analysis and Pathogen Identification. The Microbiome Analysis portion details how to use the Kraken suite to analyze the overall diversity of microbiome samples while also statistically differentiating between microbiome compositions. We illustrate this protocol using samples of a patient's fecal microbiome before and after antibiotic treatment, as studied in Taur et al.<sup>27</sup>. The Pathogen Identification protocol demonstrates how to identify pathogenic, atypical microbes within the overall microbiome using a set of 10 corneal samples from Li et al.<sup>28</sup>, 8 of which contain pathogenic microbes while 2 are control samples.

### Procedure 1: Microbiome Analysis

We will explore and visualize the depletion of patient T11's microbiota diversity due to antibiotic treatment, using our microbiome analysis pipeline. In the study by Taur et al.<sup>27</sup>, all of the patients underwent allogeneic hematopoietic stem cell transplantation (allo-HSCT) where antibiotic treatment is essential for optimal clinical outcomes. However, antibiotics deplete a patient's microbiome diversity. In the paper, the authors showed that auto-FMT treatment restores the microbial diversity to a patient's gut microbiome. In this example, we explicitly detail three samples from patient T11 collected before and during antibiotic treatment. Samples 1 and 2 were taken before T11 began antibiotic treatment and sample 3 was collected during treatment.

### Remove host DNA

1. Depending on the source of the metagenomic sample, users may remove host DNA by aligning reads to the host genome. The provided sample set has already undergone removal of host (human) DNA. We provide these command lines as a guide for how to remove human or host DNA from other samples using Bowtie 2<sup>7</sup>.

```
$ bowtie2 -x b_index/GRCh38 -p 8 -1 paired_reads_1.fastq -2
paired_reads_2.fastq \
--un-conc nonhuman_reads.fastq -S human_reads.sam
```

## Classify Microbiome Samples using Kraken

**Timing ~10 min**—2. Run Kraken 2<sup>4</sup> to classify all reads, providing each read with its taxonomy identification. Kraken 2 examines the *k*-mers within a query sequence and uses this information to query a database. In normal usage, the only required inputs are the database name, specified with `--db`, and the input FASTA file (or 2 files if the reads are paired). Below is the simplest example:

```
$ kraken2 --db $DBNAME seqs.fq
```

In our protocol, we specify the number of threads to improve the timing; with more threads the program will generally finish faster. With `--report` we specify the location and name of the `k2report` file and output the results of the run to a `kraken2` file with `> path/to/file`. The `--minimum-hit-groups` flag specifies the minimum number of "hit groups" needed to make a classification call. Hit groups are overlapping *k*-mers sharing the same minimizer. Kraken 2 uses minimizers to compress the input genomic sequences, thereby reducing storage memory needed and run time. In this example we increase the minimum number of hit groups from the default 2 groups to 3 groups for increased accuracy. Lastly, the `--report-minimizer-data` flag reports minimizer and distinct minimizer count information in addition to the normal Kraken 2 report.

```
$ kraken2 --db k2protocol_db --threads 8 --report kreports/
SRR14143424.k2report \
  --report-minimizer-data --minimum-hit-groups 3 samples/SRR14143424_1.fastq
\
  samples/SRR14143424_2.fastq > kraken_outputs/SRR14143424.kraken2
$ kraken2 --db k2protocol_db --threads 8 --report kreports/
SRR14092160.k2report \
  --report-minimizer-data --minimum-hit-groups 3 samples/SRR14092160_1.fastq
\
  samples/SRR14092160_2.fastq > kraken_outputs/SRR14092160.kraken2
$ kraken2 --db k2protocol_db --threads 8 --report kreports/
SRR14092310.k2report \
  --report-minimizer-data --minimum-hit-groups 3 samples/SRR14092310_1.fastq
\
  samples/SRR14092310_2.fastq > kraken_outputs/SRR14092310.kraken2
```

Kraken 2 has two output files, a standard Kraken 2 output, `.kraken2` file, and a Kraken 2 report, `.k2report` file. The standard Kraken 2 file outputs lines containing five tab-delimited fields; 1) "C"/"U" indicating classified or unclassified, 2) sequence ID, 3) taxonomy ID (0 if unclassified), 4) length of sequence in base pairs and 5) a space-delimited list indicating the lowest common ancestor mapping of each *k*-mer. The Kraken report file outputs lines containing 6 tab-delimited fields; 1) percentage of fragments covered by the clade rooted at this taxon, 2) number of fragments covered by the clade rooted at this taxon

3) number of fragments assigned directly to this taxon, 4) rank code such as S for species, 5) NCBI taxonomic ID and 6) indented scientific name. This information and more can be found in the Kraken 2 manual.

## Run Bracken for Abundance Estimation of Microbiome Samples

**Timing < 1 min**—3. Following classification, we use Bracken<sup>5</sup> to estimate species abundance in each sample. Here is a generic Bracken invocation:

```
$ bracken -d kraken_database -i sample.k2report -r read_length \
  -l taxonomic_level -t read_threshold -o sample.bracken -w sample.breport\
```

Below are the commands used to generate the abundance estimation in our example. Here we set the read length as the average read length in the dataset: 100bp. We set the level for abundance estimation to Species (with `-l S`), and with the `-t 10` we require 10 reads prior to abundance estimation to perform re-estimation. This effectively removes any species with fewer than 10 reads, thereby removing some noise from low abundance species.

```
$ bracken -d k2protocol_db -i kreports/SRR14143424.k2report -r 100 -l S -t
10 \
  -o bracken_outputs/SRR14143424.bracken -w breports/SRR14143424.breport
$ bracken -d k2protocol_db -i kreports/SRR14092160.k2report -r 100 -l S -t
10 \
  -o bracken_outputs/SRR14092160.bracken -w breports/SRR14092160.breport
$ bracken -d k2protocol_db -i kreports/SRR14092310.k2report -r 100 -l S -t
10 \
  -o bracken_outputs/SRR14092310.bracken -w breports/SRR14092310.breport
```

## Calculate Alpha Diversity

**Timing ~ 35 sec**—4. When trying to measure biodiversity, it is useful to quantify differences within and between ecosystems. Whittaker et al<sup>30</sup> defined several forms of alpha diversity as measures that capture the diversity within a particular ecosystem and can be expressed by the number of species in that ecosystem. Using the KrakenTools `alpha_diversity.py` script, you can calculate Berger Parker's<sup>31</sup> (BP), Fisher's<sup>32</sup> (F), Simpson's<sup>33</sup> (Si), Inverse Simpson's (ISi)<sup>33</sup>, and Shannon's<sup>34</sup> (Sh) alpha diversity for each sample after running Kraken 2 and Bracken. You can specify which alpha diversity metric you want to output to terminal with the `-a` flag, as shown below.

```
$ python KrakenTools/DiversityTools/alpha_diversity.py \
  -f bracken_outputs/SRR14143424.bracken -a BP
$ python KrakenTools/DiversityTools/alpha_diversity.py \
  -f bracken_outputs/SRR14143424.bracken -a Sh
$ python KrakenTools/DiversityTools/alpha_diversity.py \
```

```

-f bracken_outputs/SRR14143424.bracken -a F
$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14143424.bracken -a Si
$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14143424.bracken -a ISi

$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14092160.bracken -a BP
$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14092160.bracken -a Sh
$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14092160.bracken -a F
$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14092160.bracken -a Si
$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14092160.bracken -a ISi

$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14092310.bracken -a BP
$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14092310.bracken -a Sh
$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14092310.bracken -a F
$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14092310.bracken -a Si
$ python KrakenTools/DiversityTools/alpha_diversity.py \
-f bracken_outputs/SRR14092310.bracken -a ISi

```

### Calculate Beta Diversity

**Timing ~5 sec**—5. Beta diversity is useful when trying to examine the change in species diversity between two or more ecosystems. Next, we used the `beta_diversity.py` script to compute the Bray-Curtis<sup>35</sup> dissimilarity matrix, which will contain the pairwise dissimilarities among three microbiome samples. With this metric, an output of 0 means the two samples are exactly the same and 1 means they are maximally divergent.

```

$ python KrakenTools/DiversityTools/beta_diversity.py -i \
bracken_outputs/SRR14092160.bracken bracken_outputs/SRR14092310.bracken \
bracken_outputs/SRR14143424.bracken --type bracken

```

### Generate Krona Plots

**Timing < 1 min**—6. Krona plots are multi-layered pie charts frequently used in metagenomic visualization for viewing data in a phylogenetic hierarchy. Using these plots, we can view the hierarchical output from Bracken in a way to better visualize

the number of reads coming from different species, genera, etc. as percentages. We run `kreport2krona.py` to generate Krona plots<sup>36</sup> using the following commands.

```
$ python KrakenTools/kreport2krona.py -r breports/SRR14143424.breport \
  -o b_krona_txt/SRR14143424.b.krona.txt --no-intermediate-ranks
$ KronaScripts/ktImportText b_krona_txt/SRR14143424.b.krona.txt \
  -o krona_html/SRR14143424.krona.html
$ python KrakenTools/kreport2krona.py -r breports/SRR14092160.breport \
  -o b_krona_txt/SRR14092160.b.krona.txt --no-intermediate-ranks
$ KronaScripts/ktImportText b_krona_txt/SRR14092160.b.krona.txt \
  -o krona_html/SRR14092160.krona.html
$ python KrakenTools/kreport2krona.py -r breports/SRR14092310.breport \
  -o b_krona_txt/SRR14092310.b.krona.txt --no-intermediate-ranks
$ KronaScripts/ktImportText b_krona_txt/SRR14092310.b.krona.txt \
  -o krona_html/SRR14092310.krona.html
```

### Generate Pavian Plots using the Shiny App

**Timing < 1 min**—7. Using the .breport files from the Bracken run, you can use Pavian to create the plots shown in Figure 6A, B and C. Figure 2 shows the Pavian interface and the steps for creating classification networks. Open the Pavian shiny app via <https://fbreitwieser.shinyapps.io/pavian/>.

8. Click "Browse..." and Upload the 3 separate SRR\*.breport files.
9. Once files are uploaded, click "Sample" to see the hierarchical classification visualization results.
10. Click on the drop-down list to select the sample that you are viewing.
11. (Optional) Choose "Configure Sankey" to change what taxonomical ranks to display, number of taxa at each level, etc. there are 10 different settings that can be changed to customize the plots.
12. Save the network by clicking "Save Network"

### Procedure 2: Pathogen Identification

**Remove Human DNA using Bowtie 2**—1. (Optional) Remove human DNA from the sample set. The provided sample set has already undergone removal of host (human) DNA using the steps shown next, so they do not need to be run again. We provide these command lines as a guide for how to remove human or host DNA from other samples using Bowtie<sup>27</sup>.

```
$ bowtie2 -x bowtie_index/GRCh38 -p 8 -f -l paired_reads_1.fastq \
-2 paired_reads_2.fastq --un-conc nonhuman_reads.fa -S human_reads.sam
```



## Classify Reads with Kraken2-Uniq

**Timing ~6min, 42 seconds for 10 samples (8 Gb)**—2. Following removal of host DNA, the remaining reads undergo classification using Kraken2Uniq. The `--report-minimizer-data` flag forces Kraken 2 to provide unique k-mer counts per classification. Additionally, we use 8 threads for faster run times, and `--minimum-hit-groups 3` for increased classification precision (i.e., fewer false positives).

```
$ kraken2 --db k2protocol_db --threads 8 --minimum-hit-groups 3 \
--report-minimizer-data --report SRR12486971.k2report \
--paired SRR12486971_1.fastq SRR12486971_2.fastq > SRR12486971.kraken2
$ kraken2 --db k2protocol_db --threads 8 --minimum-hit-groups 3 \
--report-minimizer-data --report SRR12486972.k2report \
--paired SRR12486972_1.fastq SRR12486972_2.fastq > SRR12486972.kraken2
$ kraken2 --db k2protocol_db --threads 8 --minimum-hit-groups 3 \
--report-minimizer-data --report SRR12486974.k2report \
--paired SRR12486974_1.fastq SRR12486974_2.fastq > SRR12486974.kraken2
$ kraken2 --db k2protocol_db --threads 8 --minimum-hit-groups 3 \
--report-minimizer-data --report SRR12486978.k2report \
--paired SRR12486978_1.fastq SRR12486978_2.fastq > SRR12486978.kraken2
$ kraken2 --db k2protocol_db --threads 8 --minimum-hit-groups 3 \
--report-minimizer-data --report SRR12486979.k2report \
--paired SRR12486979_1.fastq SRR12486979_2.fastq > SRR12486979.kraken2
$ kraken2 --db k2protocol_db --threads 8 --minimum-hit-groups 3 \
--report-minimizer-data --report SRR12486981.k2report \
--paired SRR12486981_1.fastq SRR12486981_2.fastq > SRR12486981.kraken2
$ kraken2 --db k2protocol_db --threads 8 --minimum-hit-groups 3 \
--report-minimizer-data --report SRR12486983.k2report \
--paired SRR12486983_1.fastq SRR12486983_2.fastq > SRR12486983.kraken2
$ kraken2 --db k2protocol_db --threads 8 --minimum-hit-groups 3 \
--report-minimizer-data --report SRR12486988.k2report \
--paired SRR12486988_1.fastq SRR12486988_2.fastq > SRR12486988.kraken2
$ kraken2 --db k2protocol_db --threads 8 --minimum-hit-groups 3 \
--report-minimizer-data --report SRR12486989.k2report \
--paired SRR12486989_1.fastq SRR12486989_2.fastq > SRR12486989.kraken2
$ kraken2 --db k2protocol_db --threads 8 --minimum-hit-groups 3 \
--report-minimizer-data --report SRR12486990.k2report \
--paired SRR12486990_1.fastq SRR12486990_2.fastq > SRR12486990.kraken2
```

## Compare Samples to Controls using Pavian

**Timing ~5 min**—3. Following classification, we compare samples using Pavian <sup>6</sup> (Figure 3) Open the Pavian shiny app via <https://fbreitwieser.shinyapps.io/pavian/>.

4. Click "Browse..." and upload the 10 separate SRR\*.kreport2 files. Once files are uploaded, click "Comparison" to see how read counts vary across samples.f
5. Choose "Species" and "Z-score (reads)" to focus on species-level reads and calculate z-scores.
6. Sort by maximum z-scores.

The species with the max z-score for each sample is the most likely pathogen. Note that if two or more pathogens have a high z-score in a given sample, we should consider the possibility that the infection has multiple causes. Species with high read counts (or high z-scores) across all samples are considered background noise or contamination.

## Verify Classification

**Timing ~30 min**—7. For verifying the classification results, use the `extract_kraken_reads.py` script to extract reads for each of the top z-score species for each sample. This script extracts reads that matched a particular species, identified by the taxonomy ID that is provided with the `-t` parameter.

```
$ python extract_kraken_reads.py -k SRR12486971.kraken2 --include-children \
-s SRR12486971_1.fastq -s2 SRR12486971_2.fastq -t 723287 \
-r SRR12486971.k2report -o SRR12486971_A.algerae.tid723287.1.fa \
-o2 SRR12486971_A.algerae.tid723287.2.fa
$ python extract_kraken_reads.py -k SRR12486972.kraken2 --include-children \
-s SRR12486972_1.fastq -s2 SRR12486972_2.fastq -t 5059 \
-r SRR12486972.k2report -o SRR12486972_A.flavus.tid5059.1.fa \
-o2 SRR12486972_A.flavus.tid5059.2.fa
$ python extract_kraken_reads.py -k SRR12486974.kraken2 --include-children \
-s SRR12486974_1.fastq -s2 SRR12486974_2.fastq -t 5476 \
-r SRR12486974.k2report -o SRR12486974_C.albicans.tid5476.1.fa \
-o2 SRR12486974_C.albicans.tid5476.2.fa
$ python extract_kraken_reads.py -k SRR12486978.kraken2 --include-children \
-s SRR12486978_1.fastq -s2 SRR12486978_2.fastq -t 1774 \
-r SRR12486978.k2report -o SRR12486978_M.chelonae.tid1774.1.fa \
-o2 SRR12486978_M.chelonae.tid1774.2.fa
$ python extract_kraken_reads.py -k SRR12486983.kraken2 --include-children \
-s SRR12486983_1.fastq -s2 SRR12486983_2.fastq -t 10298 \
-r SRR12486983.k2report -o SRR12486983_HSV1.tid10298.1.fa \
-o2 SRR12486983_HSV1.tid10298.2.fa
$ python extract_kraken_reads.py -k SRR12486988.kraken2 --include-children \
-s SRR12486988_1.fastq -s2 SRR12486988_2.fastq -t 61605 \
-r SRR12486988.k2report -o SRR12486988_A.lugunensis.tid61605.1.fa \
-o2 SRR12486988_A.lugunensis.tid61605.2.fa
$ python extract_kraken_reads.py -k SRR12486989.kraken2 --include-children \
```

```

-s SRR12486989_1.fastq -s2 SRR12486989_2.fastq -t 1311 \
-r SRR12486989.k2report -o SRR12486989_S.agalactiae.tid1311.1.fa \
-o2 SRR12486989_S.agalactiae.tid1311.2.fa
$ python extract_kraken_reads.py -k SRR12486990.kraken --include-children \
-s SRR12486990_1.fastq -s2 SRR12486990_2.fastq -t 1280 \
-r SRR12486990.k2report -o SRR12486990_S.aureus.tid1280.1.fa \
-o2 SRR12486990_S.aureus.tid1280.2.fa

```

8. Align extracted reads to the species genomes using alignment programs such as Bowtie <sup>27</sup> or Minimap<sup>26</sup>. We use Bowtie 2 for alignment.

```

$ bowtie2 -x b_index/Aalgerae -f -p 8 \
-1 SRR12486971_A.algerae.tid723287.1.fa \
-2 SRR12486971_A.algerae.tid723287.2.fa \
-S SRR12486971_A.algerae_aligned.sam
$ bowtie2 -x b_index/Aflavus -f -p 8 \
-1 SRR12486972_A.flavus.tid5059.1.fa \
-2 SRR12486972_A.flavus.tid5059.2.fa \
-S SRR12486972_A.flavus_aligned.sam
$ bowtie2 -x b_index/Calbicans -f -p 8 \
-1 SRR12486974_C.albicans.tid5476.1.fa \
-2 SRR12486974_C.albicans.tid5476.2.fa \
-S SRR12486974_C.albicans_aligned.sam
$ bowtie2 -x b_index/Mchelonae -f -p 8 \
-1 SRR12486978_M.chelonae.tid1774.1.fa \
-2 SRR12486978_M.chelonae.tid1774.2.fa \
-S SRR12486978_M.chelonae_aligned.sam
$ bowtie2 -x b_index/HSV1 -f -p 8 \
-1 SRR12486983_HSV1.tid10298.1.fa \
-2 SRR12486983_HSV1.tid10298.2.fa \
-S SRR12486983_HSV1_aligned.sam
$ bowtie2 -x b_index/Alug -f -p 8 \
-1 SRR12486988_A.lugunensis.tid61605.1.fa \
-2 SRR12486988_A.lugdunensis.tid61605.2.fa \
-S SRR12486988_Alug_aligned.sam
$ bowtie2 -x b_index/Sagalactiae -f -p 8 \
-1 SRR12486989_S.agalactiae.tid1311.1.fa \
-2 SRR12486989_S.agalactiae.tid1311.2.fa \
-S SRR12486989_Sagalactiae_aligned.sam
$ bowtie2 -x b_index/Saureus -f -p 8 \
-1 SRR12486990_S.aureus.tid1280.1.fa \
-2 SRR12486990_S.aureus.tid1280.2.fa \
-S SRR12486990_Saureus_aligned.sam

```

9. Following alignment, we use SAMtools<sup>37</sup> to convert the SAM files to sorted BAM files.

```
$ samtools view -bS -F4 SRR12486971_A.algerae_aligned.sam \
> SRR12486971_A.algerae_aligned.bam
$ samtools view -bS -F4 SRR12486972_A.flavus_aligned.sam \
> SRR12486972_A.flavus_aligned.bam
$ samtools view -bS -F4 SRR12486974_C.albicans_aligned.sam \
> SRR12486974_C.albicans_aligned.bam
$ samtools view -bS -F4 SRR12486978_M.chelonae_aligned.sam \
> SRR12486978_M.chelonae_aligned.bam
$ samtools view -bS -F4 SRR12486983_HSV1_aligned.sam \
> SRR12486983_HSV1_aligned.bam
$ samtools view -bS -F4 SRR12486988_Alug_aligned.sam \
> SRR12486988_Alug_aligned.bam
$ samtools view -bS -F4 SRR12486989_Sagalactiae_aligned.sam \
> SRR12486989_Sagalactiae_aligned.bam
$ samtools view -bS -F4 SRR12486990_Saureus_aligned.sam \
> SRR12486990_Saureus_aligned.bam

$ samtools sort SRR12486971_A.algerae_aligned.bam \
-o SRR12486971_A.algerae_sorted.bam
$ samtools sort SRR12486972_A.flavus_aligned.bam \
-o SRR12486972_A.flavus_sorted.bam
$ samtools sort SRR12486974_C.albicans_aligned.bam \
-o SRR12486974_C.albicans_sorted.bam
$ samtools sort SRR12486978_M.chelonae_aligned.bam \
-o SRR12486978_M.chelonae_sorted.bam
$ samtools sort SRR12486983_HSV1_aligned.bam \
-o SRR12486983_HSV1_sorted.bam
$ samtools sort SRR12486988_Alug_aligned.bam \
-o SRR12486988_Alug_sorted.bam
$ samtools sort SRR12486989_Sagalactiae_aligned.bam \
-o SRR12486989_Sagalactiae_sorted.bam
$ samtools sort SRR12486990_Saureus_aligned.bam \
-o SRR12486990_Saureus_sorted.bam

$ samtools index SRR12486971_A.algerae_sorted.bam
$ samtools index SRR12486972_A.flavus_sorted.bam
$ samtools index SRR12486974_C.albicans_sorted.bam
$ samtools index SRR12486978_M.chelonae_sorted.bam
$ samtools index SRR12486983_HSV1_sorted.bam
$ samtools index SRR12486988_Alug_sorted.bam
$ samtools index SRR12486989_Sagalactiae_sorted.bam
$ samtools index SRR12486990_Saureus_sorted.bam
```

10. Finally, upload the sorted BAM file and the index of the sorted BAM file (.bam and .bai files) to Pavian's Alignment Viewer for the coverage plot and statistics. Figure 4 shows example coverage plots for the SRR12486978 reads aligned to the *Mycobacterium chelonae* genome and the SRR12486989 reads aligned to the *Streptococcus agalactiae* genome.

## Timing

### Troubleshooting

**Step 2. Run Kraken 2 results in large numbers of unclassified reads**—If your Kraken 2 output files have too many unclassified reads, it probably means there are organisms missing from your working Kraken 2 database. To classify more reads, you should download additional genomes and add them to the database.

**Steps 3–6. After Kraken 2 and Bracken calls**—If there are any errors during these steps, you should check that your `kraken2` and `bracken` files were successfully made. For example, if the output from `kraken2` looks like this, there was a problem in the `kraken2` call:

```
Loading database information ... done.
0 sequences (0.00 Mbp) processed in 0.001s (0.0 Kseq/m, 0.00 Mbp/m).
0 sequences classified (-nan %)
0 sequences unclassified (-nan %)
```

The command used to create the erroneous output shown above is as follows:

```
for sample in SRR14092160 SRR14092310; do
    kraken2 --db k2protocol_db --threads 8 --report kreports/$sample.k2report \
    --report-minimizer-data --minimum-hit-groups 3 samples/$sample_1.fasta \
    samples/$sample_2.fasta > kraken_outputs/$sample.kraken2
done
```

The problem in this example is the bash syntax. Here, the input filenames are not interpolated correctly because the “\$sample” variable is not protected against trailing underscores in the filenames. Therefore, bash is then attempting to reference a nonexistent variable. In this example, the commands that follow will throw an error because 0 sequences were classified and the `.kraken2` file is empty. To fix this, you should check that you are referencing the files correctly. The output of a successful Kraken 2 run for sample SRR14092160 is as follows (you can specify as many samples as you want, separated by spaces):

```
Loading database information ... done.
73021500 sequences (8907.00 Mbp) processed in 154.050 s (28440.6 Kseq/m,
```

```
3469 Mbp/m) .
11996432 sequences classified (16.43%)
61025068 sequences unclassified (83.57%)
```

There should be non-zero values for the number of sequences processed and for the number of reads classified. This can be done by protecting the code via some form of quoting. The following command, with double quotes around the filenames, does this:

```
for sample in SRR14092160 SRR14092310; do
kraken2 --db k2protocol_db --threads 8 --report " kreports/${
{sample}.k2report" \
--report-minimizer-data --minimum-hit-groups 3 "samples/${sample}_1.fasta" \
"samples/${sample}_2.fasta" > "kraken_outputs/${sample}.kraken2"
done
```

## Anticipated results

The accuracy of classification and abundance estimation is dependent on the quality and composition of the genomes used in the Kraken and Bracken databases. If the database is missing genomes present in the sequenced samples, there might be a high proportion of unclassified reads when using Kraken. If the database contains contaminated genomes (e.g. bacteria that have bits of human sequence in them<sup>20;21</sup>), this will lead to false positives, where sequencing reads will be identified as incorrect species. Unclassified reads and mis-classified reads will lead to additional problems downstream for Bracken, which will only report abundances for species identified by Kraken. To ensure a higher classification percentage, we recommend using a comprehensive Kraken database containing, at minimum, the human genome and bacterial, viral, archaeal, vector, and eukaryotic pathogen genomes. For a low rate of false positives, we recommend using only complete or quality-controlled genomes in the building of the Kraken database.

In this protocol, we use a "MiniKraken" version of this comprehensive database that includes the human genome, complete bacterial, viral, and archaeal genomes from RefSeq, and a filtered eukaryotic pathogen database. The MiniKraken version of the comprehensive database contains sequences from all of the aforementioned genomes, but it down-samples the k-mers from each genome sufficiently to allow Kraken to run with only 8GB of RAM. Kraken should still detect accurate bacterial, viral, archaeal, and eukaryotic microbial signatures, but as we limit the number of sequences, we also expect a significant fraction of unclassified reads.

## Microbiome Analysis

The three metagenomic sequencing samples that we use in this protocol illustrate the normal state of patient T11's microbiome prior to treatment and its depleted diversity state after antibiotic treatment. (The original study also measured diversity after a fecal microbial transplant restored much of the microbiome.) Supplemental Table 1 lists the read counts per species in the final Bracken analysis of the three samples. In Figure 5, we can

find the computed alpha diversity outputs for each sample, for each alpha diversity type currently available. We calculate these values using the abundance estimation calculation from Bracken using KrakenTools' alpha diversity script. In samples 1 and 2, both taken when patient T11 had normal microbiome diversity, we expect to see a higher value for alpha diversity than in sample 3, which was taken during antibiotic treatment. In Figure 5 A, we see exactly that across the board for all the alpha diversity types. In the original study, they published the alpha diversity values shown in the last row of table A. They specified that it is Inverse Simpson's alpha diversity however they did not include the exact tool or equations they used to find these values. The Inverse Simpson's alpha value on row 4 of table A is the value we found using Bracken's abundance estimation and using the equations shown. For sample 2, we found a significant discrepancy in the alpha diversity published in the original paper (10.55) and our calculation (21.5). However, this can be explained by the large percentage of unclassified reads for this sample (about 80%) and a difference in formulation. Since the Kraken 2 report has many unclassified reads, the Bracken abundance estimation is not being calculated for the entire sample but only for the 20% of reads that are classified. In order to have more classified reads, a larger database needs to be used.

Figure 5 also shows the output of the beta diversity calculation based on the Bray-Curtis dissimilarity matrix shown in section B. The heat map shows that when comparing the same two samples, for example Day -2 (sample 1) with itself, the beta diversity value is 0 because the two samples being compared are exactly the same. When comparing Day -2 with Day 12 (sample 3) the beta diversity is close to 1 because the diversity levels in each are very different from each other. Also, as expected, we see that samples 1 and 2 (day -2 and -9) are somewhat similar with a beta diversity value of 0.6.

In Figure 6, we have the Pavian and two of the Krona plots we created. First, focusing on the left-side, subplots A-C are the Pavian plots for samples 1–3 respectively. Here we see that plot C is dominated by a single species of bacteria, *Enterococcus faecium*, showing patient T11 only had one species of bacteria in his microbiome while undergoing antibiotic treatment. In plots A and B, we see several species of bacteria showing T11's normal variation in microbiome diversity. On the right-side of figure 6, we have two Krona plots made using samples 2 and 3, top to bottom. Here we see the top Krona plot is divided into several different sections which means represent the different species present in the sample. In the bottom plot, we see much fewer partitions showing again how *E. faecium* dominates the sample.

Using the MiniKraken database, there will be a large number of unclassified reads, as many as 42,899,387 in sample 1. For plotting purposes, we simply discarded all unclassified reads from the `kreport2krona.py` output when creating the plots shown in Figure 6. All of the visualizations methods used, Krona plot, Pavian plot, and Beta diversity heatmap, show the great difference in microbiome diversity between patient T11 before and during antibiotic treatment.

## Pathogen Detection

Because we are using the MiniKraken version of our comprehensive database, we expect a high proportion of unclassified reads across all infectious disease samples. Table 3 lists



the total read counts per sample and the number of unclassified and classified reads per sample. However, because the MiniKraken database was selected from a comprehensive set of genomes, we still expect to see classifications across all major taxonomic clades (e.g. bacteria, archaea, viruses, fungi, etc.). Table 4 lists the read counts for major clades, showing 80K–2.7M bacterial reads per sample, along with varying numbers of reads from archaea, fungi, and amoebae. Full species read counts for each sample are listed in Supplemental Table 2.

Despite the many clades detected in the corneal samples, the diagnostic challenge is determining the most likely pathogen(s) per sample. Samples SRR12486979 and SRR12486981 are control samples, providing a baseline for read counts from species are likely to be contaminants or possibly non-infectious components of the samples. For each of the remaining patient samples, we are interested in any species with higher than average read counts as compared to the control samples or other corneal samples. For example, sample SRR12486971 has 84,000+ *Anncaliia algerae* reads. All other samples have 12 *A. algerae* reads or fewer, making it appear that *A. algerae* is a likely pathogen in that sample (which indeed it is, as was confirmed in the original study).

The easiest method for finding the most likely pathogen is by uploading a set of similar samples to Pavian <sup>6</sup>, as described in Step 3 of our Pathogen Identification protocol. Using the comparison tab, Pavian can calculate and sort the species by z-scores, a metric used to determine whether a species (or any other taxonomic clade) has significantly higher read counts in one sample as compared to the remaining other samples. Table 5 lists the species with the highest z-score in each sample. As shown in the table, the highest z-score correctly identified the true pathogen for each of the non-control samples.

The highest z-score species can be further checked by using the k-mer-counting feature in KrakenUniq<sup>3</sup> (which is also available as a parameter in Kraken 2). As described in the KrakenUniq paper, for most species in a metagenomics sample, each read will be a random sample from the genome, which means that each read will tend to contribute approximately  $r-k$  distinct k-mers to the k-mer count, where  $r$  is the read length. However, if a bacterial genome is contaminated with a small amount of human sequence (for example), then large numbers of human reads will incorrectly match a small part of that genome. In these cases, the number of distinct k-mers that are observed from a genome will be very small compared to the number of reads. Thus, if the k-mer counts listed in the Kraken 2 report are relatively small, the species is probably not present but instead is a false positive caused by a contaminated genome in the database. Figure 7 summarizes the pathogen identification results for each individual sample, displaying a heat map of reads, k-mers, and z-scores across all samples.

Finally, we validate the classification results by extracting the classified reads using KrakenTools and aligning the reads against the suspected pathogen genomes using Bowtie 2<sup>7</sup>. We upload the alignment BAM files to Pavian <sup>6</sup> for a clear visualization of the read coverage across the genomes, confirming that the reads are derived from the full pathogen's genome.

As a result of this protocol, we have a set of potential infectious agents discovered in the samples. The infectious agents should be presented to pathologists and clinicians for further verification with other independent methods.

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

## Acknowledgments

Indexes for tools in the Kraken suite, including the indexes used in this protocol, are made freely available on Amazon Web Services thanks to the AWS Public Dataset Program.

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## Availability

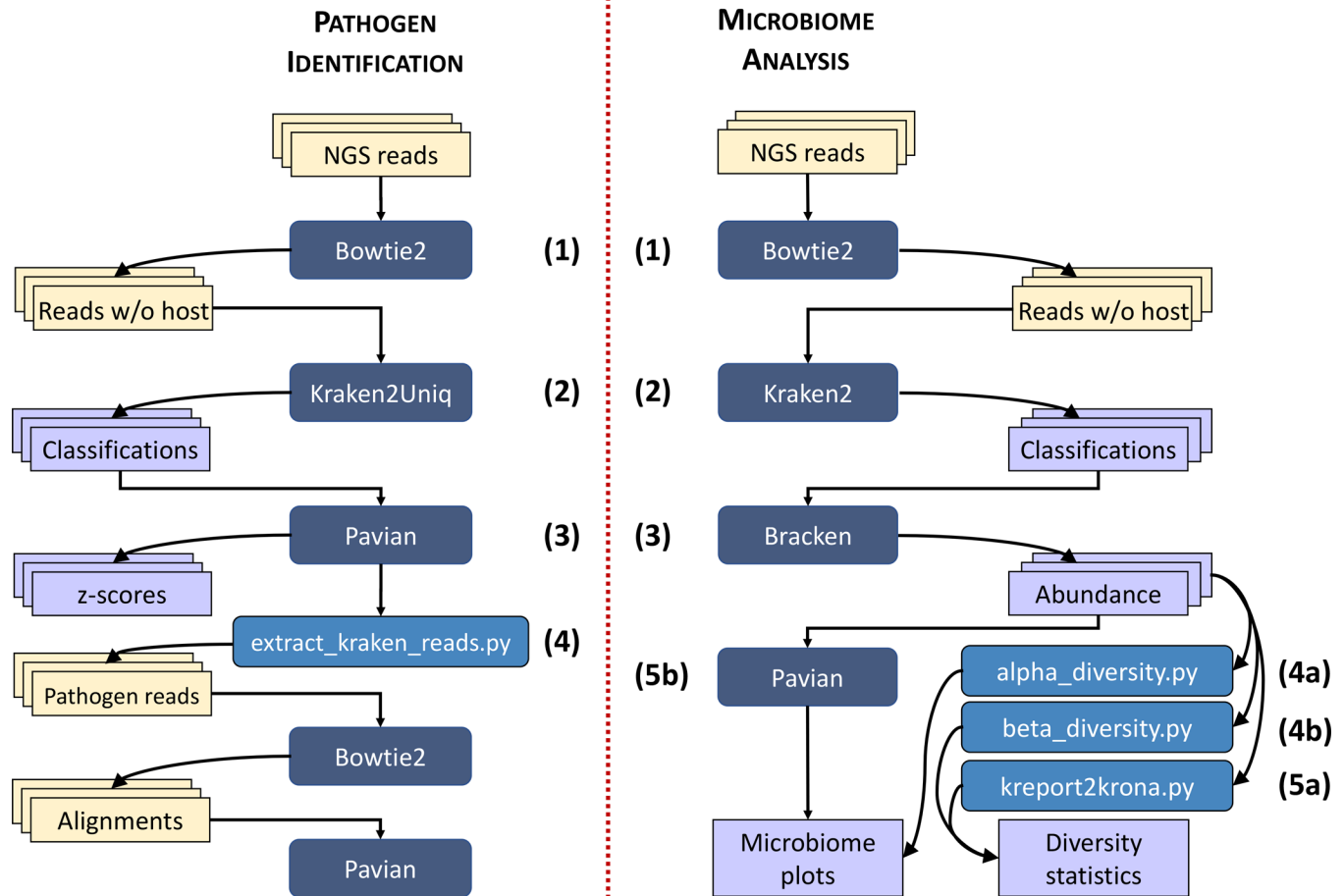
The following website details and links all software and databases used in this protocol: [http://ccb.jhu.edu/data/kraken2\\_protocol/](http://ccb.jhu.edu/data/kraken2_protocol/). We also provide easy-to-use jupyter notebooks for both workflows, which can be executed in the browser using Google Collab: <https://github.com/martin-steinegger/kraken-protocol/>

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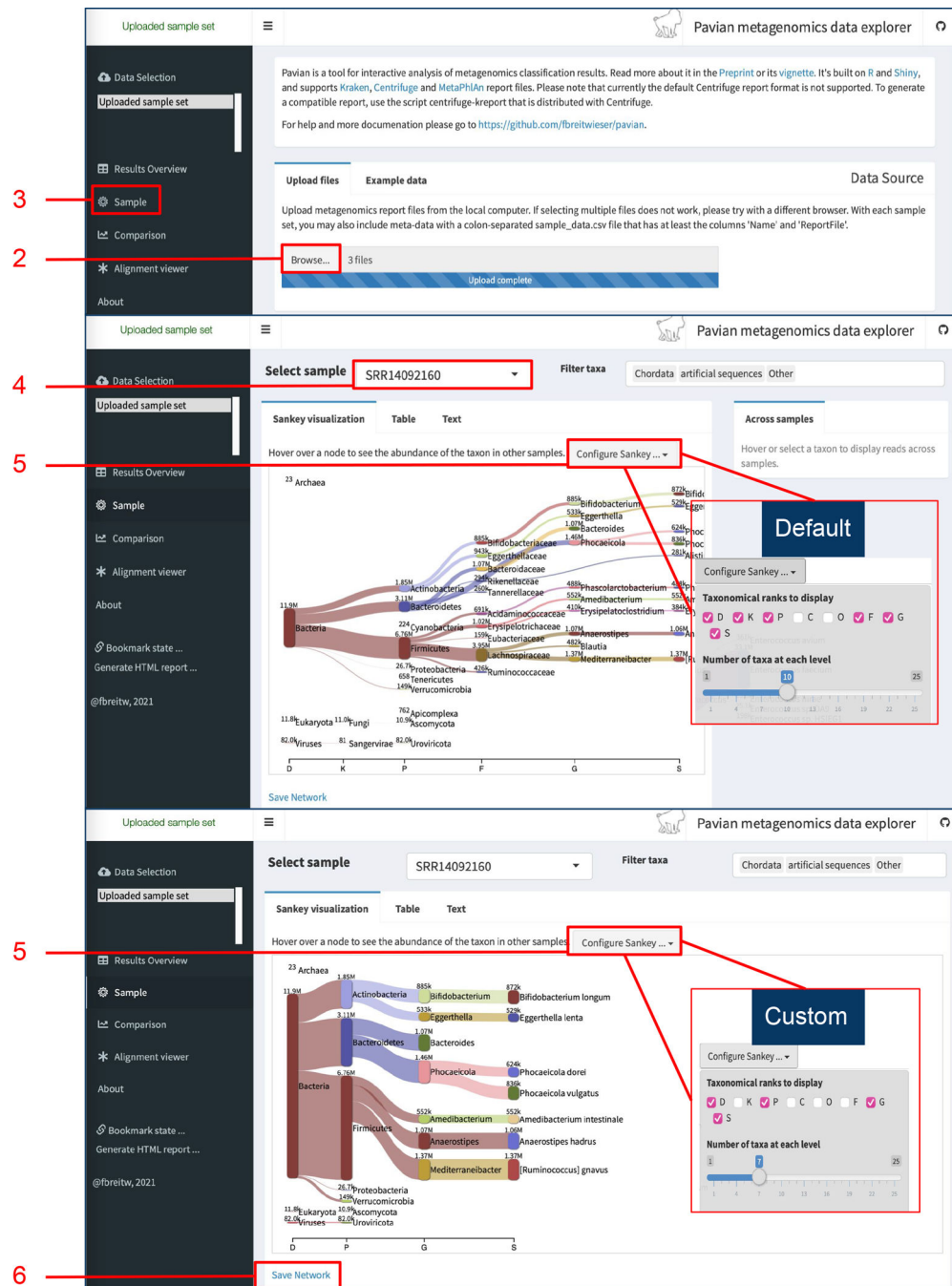
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**Fig 1. Protocol workflow.**

Overview of two workflows (1) pathogen identification and (2) microbiome analysis. (1) Here we try to detect an infectious agent using NGS reads. For this we start with a sample from the infection site and (ideally) a negative control. As a first step, host DNA is removed by excluding all reads aligning to the host genome using Bowtie 2. This step usually removes a large fraction of the reads. Remaining reads are then classified by Kraken2Uniq against a reference database, and the taxonomic reports are compared using Pavian. Pavian can distinguish large abundance changes between controls and infected samples using z-statistics. For all potential pathogen candidates, reads can be extracted using `extract_kraken_reads.py`. In workflow (2) we try to estimate the abundance of species in microbiome samples and compute the diversity changes between them. In the protocol, we start with multiple sets of reads from a microbiome before and after fecal transfer. All samples are classified using Kraken 2. Bracken takes the classified read counts and estimates the abundance of each taxon in the sample. Pavian can be used to explore and visualize this sample to spot the difference. Additionally, `alpha_diversity.py` can be used to quantify the diversity in a sample and `beta_diversity.py` can be used to compare diversity across samples.



**Fig 2. Pavian Output for Hierarchical Visualization**

Upon (1) opening the Pavian app, users should (2) upload the microbiome sample files. (3) Choose "Sample" to view classification visualization results. (4) Select sample from the drop-down menu. (5) Select plot settings to customize visualization. (6) Save image of network.

The screenshot displays the Pavian metagenomics data explorer interface. The left sidebar contains navigation options: Data Input, Results Overview, Sample, Comparison, Alignment viewer, and About. The main panel shows the 'Uploaded sample set' section with a 'Browse...' button and a progress bar indicating '10 files' uploaded. Below this, the 'Comparison' section is active, showing a table of read counts per sample per taxon. The table columns include Name, Rank, TID, Max, and various sample IDs (SRR12486971, SRR12486972, SRR12486974, SRR12486978, SRR12486979, SRR12486981, SRR12486983, SRR12486988). The table is sorted by 'Max' (z-score) in descending order. The bottom of the interface shows a 'Download full table in tab-separated value format' button.

2

3

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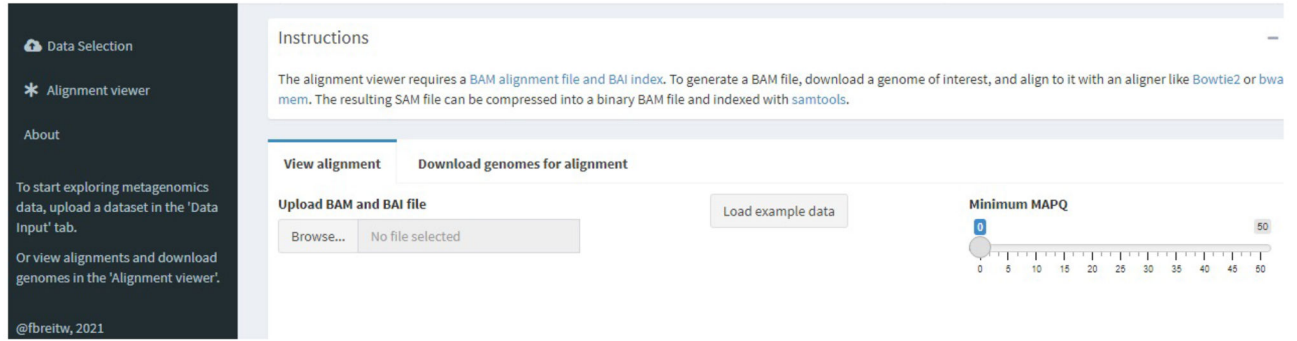
**Fig 3. Pavian Output for Pathogen Identification.**

Upon (1) opening the Pavian app, users should (2) upload the pathogen sample files. (3) Choose “Comparison” to view the table of read counts per sample per taxon. (4) Select ‘Species’ and ‘Z-score (reads)’ to filter the table and calculate z-scores. (5). Finally, sort by max z-scores to focus on species that are most likely pathogen candidates.



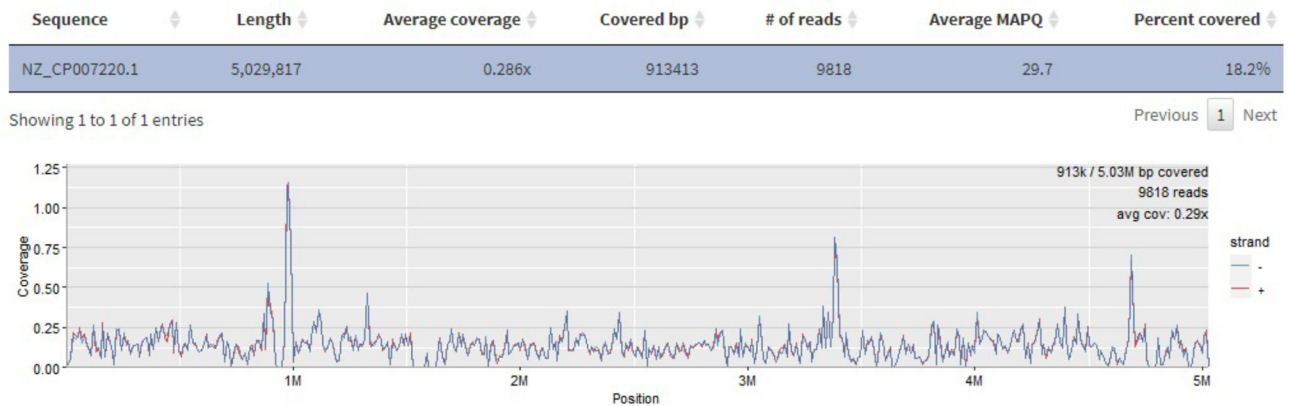
A

## Pavian alignment view



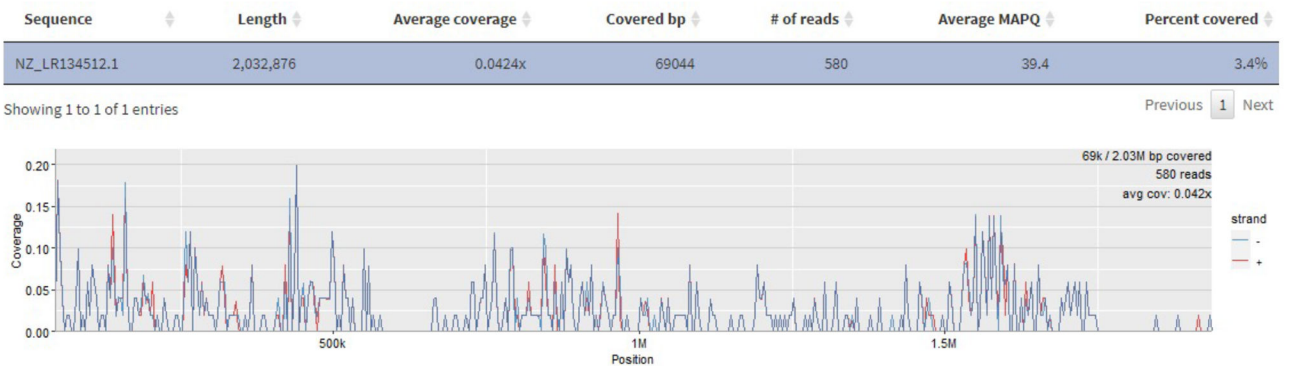
B

## SRR12486978 *Mycobacterium chelonae*



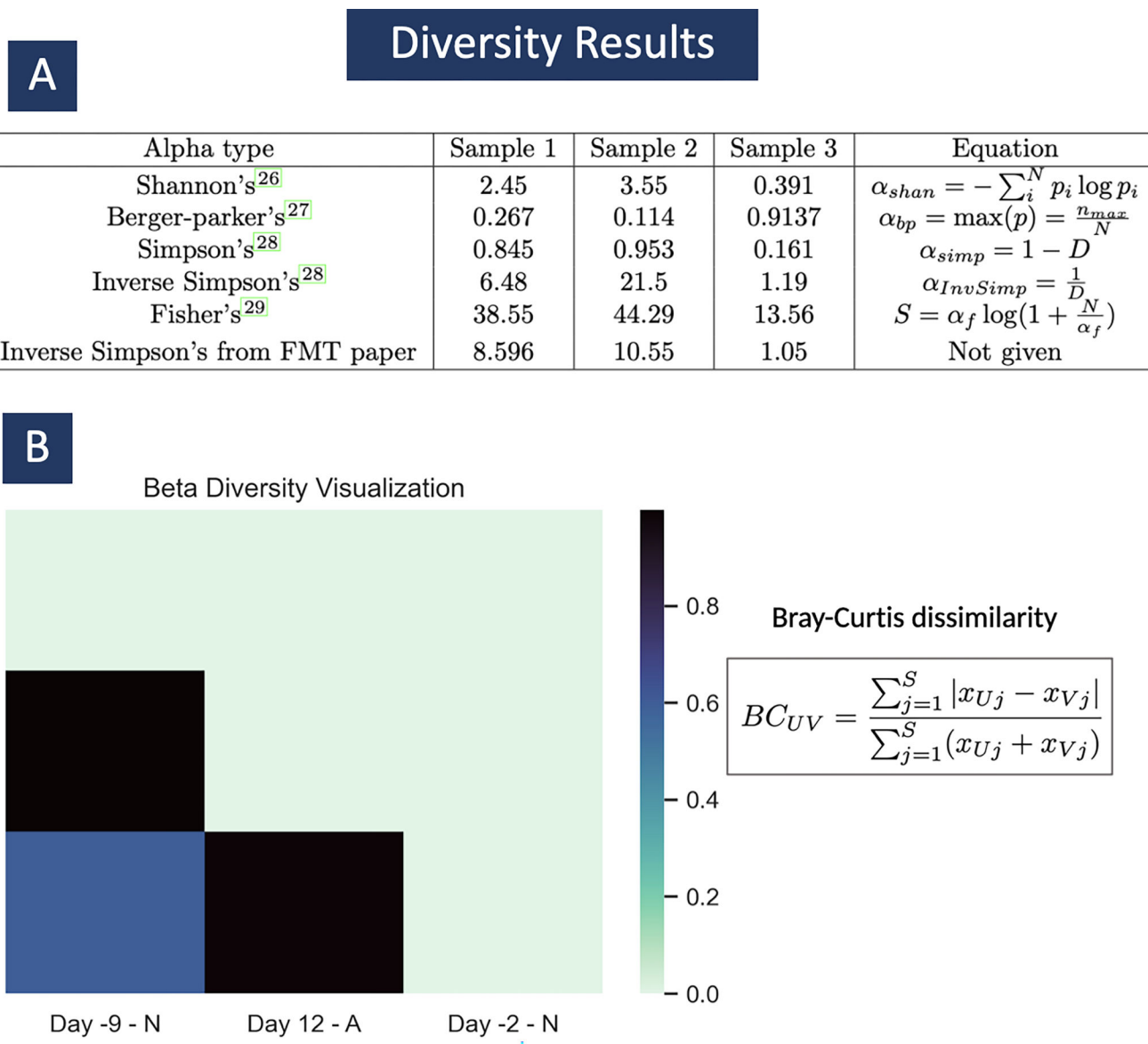
C

## SRR12486989 *Streptococcus agalactiae*



**Fig 4. Pavian Alignment Viewer.**

A) shows the graphical interface of Pavian's alignment viewer. Users should upload the .bam and .bai files to the alignment viewer. B) and C) show two example coverage plots for the pathogen identification samples. Pavian displays the coverage plot along with summarizing coverage statistics.



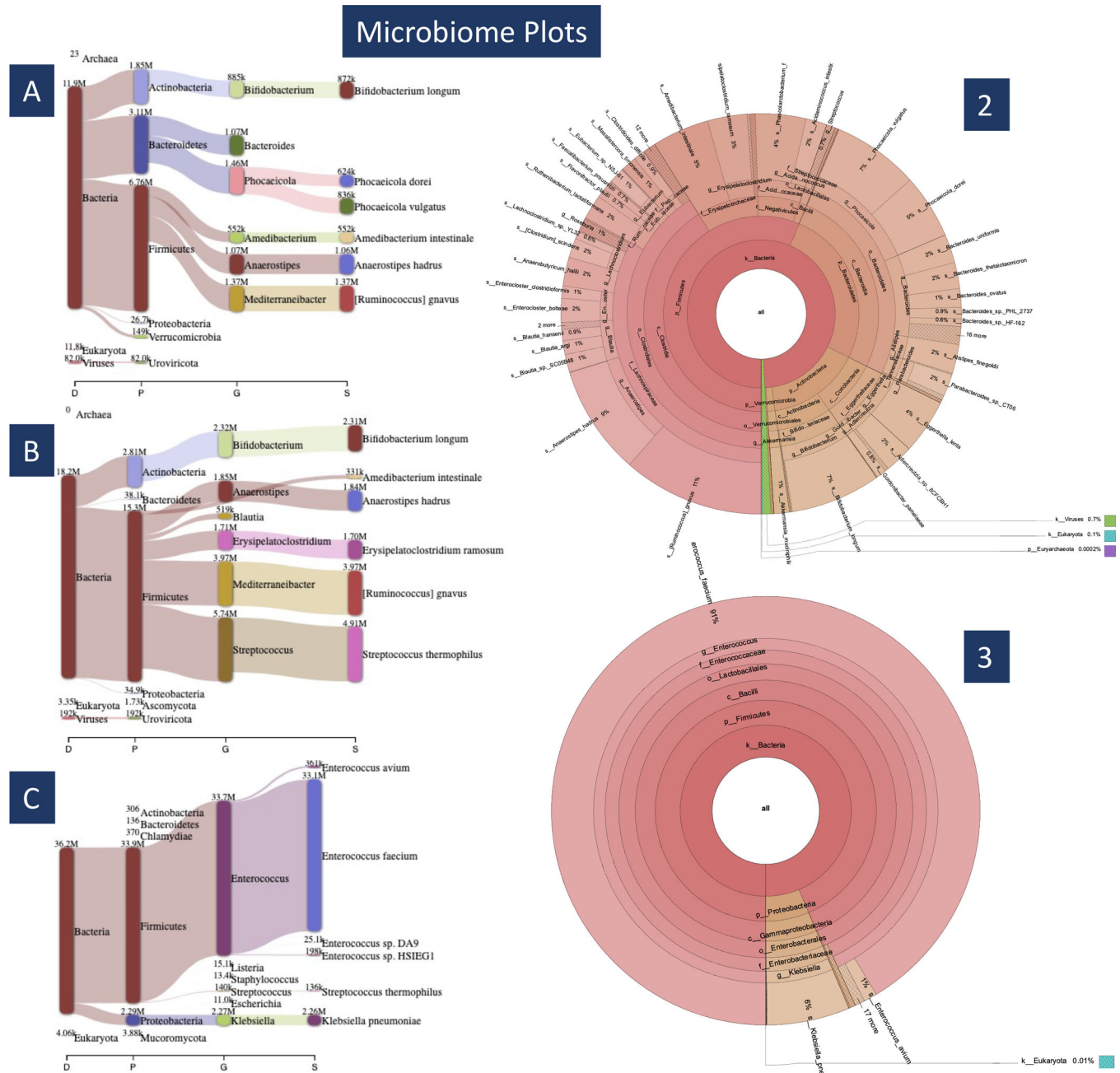
**Fig 5. Alpha and Beta diversity results.**

In subplot A, we can see the computed results for alpha diversity. In the equations  $p$  is

a vector of  $p_i$ s where  $p_i = \frac{(\text{number of individuals in } i^{th} \text{ species})}{\text{total number of individuals}}$  for all  $i$  species i.e.  $p_i = \frac{n_i}{N}$ . And

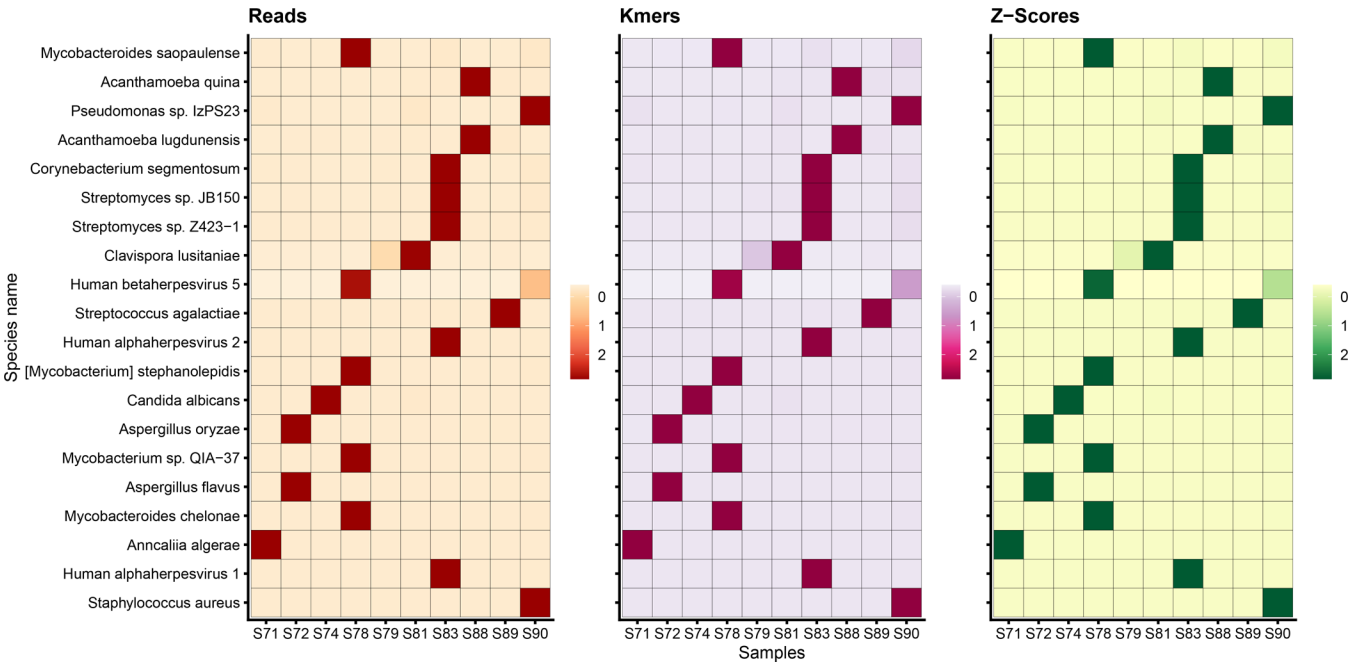
$D = \frac{\sum n_i(n_i - 1)}{N(N - 1)}$ . In subplot B, we can see a heatmap of the 3 samples from 3 different

time points in patient T11's treatment. The sample taken on day 12, was taken while T11 was taking antibiotics, marked with an 'A'. The samples taken from days -9 and -2, were taken before the commencement of antibiotic treatment, marked with an 'N'. Here we use `beta_diversity.py` to compare diversity across samples. This is the Bray-Curtis dissimilarity matrix. We see that the two samples taken before commencement of treatment are more similar to each other than either sample compared to sample A, from day 12.



**Fig 6. Microbiome Plots.**

In subplots A–C we see Pavian visualization for samples 1–3. Samples 1 and 2 have a similar taxonomic breakdown (A & B), corresponding to T11's normal microbiome diversity. Subplot C shows that sample 3 is dominated by a few bacteria when T11 is taking antibiotics. On the right are Krona plots generated from samples 2 and 3. The plot on top (2) shows the diversity of patient T11 before receiving any antibiotic treatment (sample 2) and the plot below it (3) shows how depleted his microbiome diversity while he is taking antibiotics (sample 3).



**Fig 7. Pathogen Identification Results.**  
The above plot summarizes the Kraken2Uniq results across the 10 corneal samples. The number of reads, number of k-mers, and z-scores reveals the most likely pathogen for each sample. For example, *Acanthamoeba quina* has high read and k-mer counts in S88 alone, *Staphylococcus aureus* is prevalent in S90, and *Human alphaherpesvirus 1* is likely to infect S83. For each sample, the true pathogen is the pathogen with the highest z-score for that particular sample.

**Table 1.**

## NCBI SRA Samples

Microbiome Analysis				
Sample	SRA ID	File size	Description	Expected Diversity
1	SRR14143424	5.9Gb	Paired, Day -2	Normal
2	SRR14092160	3.4Gb	Paired, Day -9	Normal
3	SRR14092310	2.3Gb	Paired, Day 12	Low
Pathogen Identification				
Sample	SRA ID	File size	Case ID	Expected Pathogen
1	SRR12486971	178.2Mb	Case 10	<i>Anncaliia algerae</i>
2	SRR12486972	318.5Mb	Case 9	<i>Aspergillus fumigatus</i>
3	SRR12486974	141Mb	Case 7	<i>Candida albicans</i>
4	SRR12486978	110.5Mb	Case 3	<i>Mycobacteroides chelonae</i>
5	SRR12486979	112.3Mb	Case 20	Control
6	SRR12486981	311.1Mb	Case 18	Control
7	SRR12486983	236.5Mb	Case 16	HSV 1
8	SRR12486988	351Mb	Case 11	<i>Acanthamoeba castellanii</i>
9	SRR12486989	228.2Mb	Case 2	<i>Streptococcus agalactiae</i>
10	SRR12486990	465.7Mb	Case 1	<i>Staphylococcus aureus</i>

**Table 2.**

Estimated time required for the microbiome analysis steps

Step	Description	Timing
1	Download and Setup	26 min
2	Classify/Estimate abundance	10 min
3	Alpha Diversity	15 sec
4	Beta Diversity	5 sec
5	Generate Krona	5 sec
6	Run Pavian	10 sec
	Total	37 min

**Table 3.**

Classification read counts from corneal samples using the MiniKraken database

Sample	Total Reads	Unclassified Reads	Unclassified %	Classified Reads	Classified %
SRR12486971	3,664,512	2,899,189	79.1%	765,323	20.9%
SRR12486972	7,594,644	7,285,624	95.9%	309,020	4.1%
SRR12486974	3,335,998	3,162,788	94.8%	173,210	5.2%
SRR12486978	2,625,249	2,496,107	95.1%	129,142	4.9%
SRR12486979	2,371,302	2,023,600	85.3%	347,702	14.7%
SRR12486981	6,730,160	6,093,775	90.5%	636,385	9.5%
SRR12486983	4,819,760	3,234,956	67.1%	1,584,804	32.9%
SRR12486988	8,369,736	8,122,882	97.1%	246,854	2.9%
SRR12486989	5,440,369	5,252,849	96.6%	187,520	3.4%
SRR12486990	9,402,750	6,619,669	70.4%	2,783,081	29.6%



**Table 4.**  
**Cornea samples per-clade read counts.**

This table lists the per-clade read counts for the major clades in these samples. Only select major clades are listed with read counts for clarity.

Sample	Bacteria	Archaea	Virus	Fungi	Amoeba
SRR12486971	649,685	34	228	85,396	45
SRR12486972	150,302	10	50	31,602	54
SRR12486974	85,361	7	14	36,833	18
SRR12486978	91,448	2	763	87	3
SRR12486979	317,804	21	64	885	36
SRR12486981	465,581	51	157	2,082	110
SRR12486983	888,749	44	646,017	1,496	54
SRR12486988	80,095	12	698	325	5587
SRR12486989	97,912	7	37	256	83
SRR12486990	2,674,036	152	1,109	4,449	166

**Table 5.**  
**Species with the highest Z-scores in cornea samples.**

This table lists the species with the highest z-scores for each of non-control samples.

Sample	True Infection	Z-score Species	Taxid	Reads	Z-score
SRR12486971	<i>Anncaliia algerae</i>	<i>Anncaliia algerae</i>	723287	84,409	56930
SRR12486972	<i>Aspergillus flavus</i>	<i>Aspergillus flavus</i>	5059	3,814	3814
SRR12486974	<i>Candida albicans</i>	<i>Candida albicans</i>	5476	36,609	1452
SRR12486978	<i>Mycobacterium chelonae</i>	<i>Mycobacterium chelonae</i>	1774	11,320	3817
SRR12486979	Control	-	-	-	-
SRR12486981	Control	-	-	-	-
SRR12486983	<i>HSV 1</i>	<i>HSV 1</i>	10298	635,691	65960
SRR12486988	<i>Acanthamoeba</i>	<i>Acanthamoeba lugdunensis</i>	61605	1,004	338
SRR12486989	<i>Streptococcus agalactiae</i>	<i>Streptococcus agalactiae</i>	1311	797	797
SRR12486990	<i>Staphylococcus aureus</i>	<i>Staphylococcus aureus</i>	1280	1,414,661	76330