# ONE ARM Metagenomics Pipeline

Prepared for and by: Project 4

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# Part I Project overview

Project Overview 7

Metagenomic analysis of Antibiotic/Antimicrobial Resistance Genes (ARGs) in NCR (Metro Manila) hospitals, wastewaters, and surface waters.

# ✓ Done □ Pending ○ Optional ○ Needs refinement ○ Robust △ Unexpected issues → Drafted ◆ TBA - needs further investigation ◆ Moved

# Chapter 1

# Bioinformatics Progress

#### 1.1 Kanban Table

Table 1.1: Detailed Kanban Table for HPC and File Management Tasks  $\,$ 

Section	Task			Ö	Δ	<b>Ð</b>	<b>√</b>			
	Confirmation of Required Robustness						<b>✓</b>			
	Agree upon the type of analyses					Ð				
	SLURM request managen	ent			,					
HPC Preparation	Calculate using quotation from PGC						✓			
III C I reparation	Setup Docker containers for SLURM									
	SLURM container for simulations									
	Automation of SLURM requests									
	Send email for HPC usage requests			Ö						
File Management	BAM file parsing									
r ne management	Interconversion between SAM and BAM									
	Create the script									
	Raw reads QC						✓			
	raw.bash									
	Raw reads trimming						✓			
	trimmomatic.bash									
	fastp.bash									
	Looping mechanism for QC and trimming						✓			
	trimming-cleaning-checking.py									
	Data visualization									
	Determination of optimal tool/s									
	Parametric randomization						✓			
	trim_randomizer.smk									
	Parse QC of all metrics in parametric randomiz	er					✓			
	parseFastQC.py									

Raw Read Processing

Section	Task			Ö	$\triangle$	<b>(2)</b>	<b>√</b>			
	Determination of optimal parameters									
	Tool combination randomization script									
	Test on Datasets									
	Raw reads QC						<b>√</b>			
	Raw reads trimming						<b>√</b>			
	trimmomatic.bash	'		'						
	fastp.bash									
	trimming-cleaning-checking.py									
	Aggregate quality metrics						<b>√</b>			
	summary_stat.bash									
	Parametric randomization						$ $ $\checkmark$			
	Parse QC of all metrics in parametric randomizer									
	Integrate to pipelines									
	Pipeline integration of raw.bash									
	Raw reads trimming						<b>√</b>			
	Create the script									
	Kraken2						✓			
	krakenpipeline.bash									
	Bracken						$ $ $\checkmark$ $ $			
	krakenpipeline.bash									
	Diversity indices						$ $ $\checkmark$			
	calculate_diversity.py									
	MetaPhlan4									
	Test on Datasets									
Taxonomy	Kraken2						<b>√</b>			
Taxonomy	Bracken						<b>√</b>			
	Diversity indices						✓			
	Integrate to pipelines									
	Kraken2						✓			
	Bracken						<b>√</b>			
	Diversity indices						<b>√</b>			
	Test whole pipeline						<b>√</b>			
	metagenomics_general.smk									
	Bootstrapping script						$ $ $\checkmark$			
	bootstrapping_rawreads.smk									
	Create the script									
	metaSPAdes									
	MEGAHIT						✓			
	MASURCA									
	PLASS									
	AbySS									

Task			Ö	Δ	<b>(2)</b>	<b>√</b>	
Test on Datasets							
metaSPAdes							
MEGAHIT						✓	
MASURCA							
PLASS							
AbySS							
KMA-iterative							
Benchmarking between all of them							
Contig quality checking							
Integrate to pipelines							
MetaWrap binning				Δ			
Kraken (MEGAHIT)						✓	
Testing on datasets				Δ			
Testing on higher depth datasets							
Refinement of bins							
Testing binning pipeline							
RGI			Ŏ				
ShortBRED						<b>√</b>	
AMRFinder			Ö				
Test on datasets	ı	ı	ı	l		ı	
RGI							
ShortBRED							
AMRFinder							
	metaSPAdes MEGAHIT MASURCA PLASS AbySS KMA-iterative Benchmarking between all of them Contig quality checking Integrate to pipelines MetaWrap binning Kraken (MEGAHIT) Testing on datasets Testing on higher depth datasets Refinement of bins Testing binning pipeline RGI ShortBRED AMRFinder Test on datasets RGI ShortBRED	metaSPAdes MEGAHIT MASURCA PLASS AbySS KMA-iterative Benchmarking between all of them Contig quality checking Integrate to pipelines MetaWrap binning Kraken (MEGAHIT)  Testing on datasets Testing on higher depth datasets Refinement of bins Testing binning pipeline  RGI ShortBRED AMRFinder Test on datasets  RGI ShortBRED	Test on Datasets  metaSPAdes  MEGAHIT  MASURCA  PLASS  AbySS  KMA-iterative  Benchmarking between all of them  Contig quality checking  Integrate to pipelines  MetaWrap binning  Kraken (MEGAHIT)  Testing on datasets  Testing on higher depth datasets  Refinement of bins  Testing binning pipeline  RGI  ShortBRED  AMRFinder  Test on datasets  RGI  ShortBRED	Test on Datasets  metaSPAdes  MEGAHIT  MASURCA  PLASS  AbySS  KMA-iterative  Benchmarking between all of them  Contig quality checking  Integrate to pipelines  MetaWrap binning  Kraken (MEGAHIT)  Testing on datasets  Testing on higher depth datasets  Refinement of bins  Testing binning pipeline  RGI  ShortBRED  AMRFinder  Test on datasets  RGI  ShortBRED  ShortBRED	Test on Datasets  metaSPAdes  MEGAHIT  MASURCA  PLASS  AbySS  KMA-iterative  Benchmarking between all of them  Contig quality checking  Integrate to pipelines  MetaWrap binning  Kraken (MEGAHIT)  Testing on datasets  Testing on higher depth datasets  Refinement of bins  Testing binning pipeline  RGI  ShortBRED  AMRFinder  Test on datasets  RGI  ShortBRED  ShortBRED	Test on Datasets  metaSPAdes  MEGAHIT  MASURCA PLASS AbySS  KMA-iterative  Benchmarking between all of them  Contig quality checking  Integrate to pipelines  MetaWrap binning  Kraken (MEGAHIT)  Testing on datasets  Testing on higher depth datasets  Refinement of bins  Testing binning pipeline  RGI  ShortBRED  AMRFinder  Test on datasets  RGI  ShortBRED	

## Script Descriptions

This section covers all the scripts that were created during the Project.

Listen, I'm eternally curious and I don't just want to settle for any random journal. No offense, but I'm aiming for something high-impact!

These were created during off hours or during work hours, so some scripts might seem irrelevant at first, but trust me, there's a conscientiousness or meticulousness to this madness.

I have separated them into folders/repositories (shameless plug here: https://github.com/-GABallena) based on their relevance to the project.

- Project4 Essential scripts directly related to the core analyses of the project.
- Side Scripts that can potentially be used to increase the robustness of the paper.
- Main Scripts related to file organization and data management, crucial for handling large datasets.

Bioinformatics work

### Chapter 2

# Project 4 Scripts

#### 1.2.1 ARG-MGE.smk

- <sup>2</sup> Stage: Draft General Purpose
- This pipeline is designed for comprehensive metagenomic analysis of ARGs, it also includes
- 4 placeholders for analyses of mobile genetic elements (MGEs), and plasmid detection. It integrates
- 5 tools for read quality control, assembly, annotation, taxonomic profiling, and structural variant
- 6 (SV) detection to provide a high-resolution view of the genetic components in metagenomic
- 7 samples.

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9 Preprocessing

#### 2.1.1 Pair merging

- 12 Technical Notes: PEAR (Paired-End reAd mergeR) compares paired-reads to correct infer the
- 13 likely bases on its associated pair
- Rationale: The main goal here is to ensure good read quality for downstream analyses and to
- maximize the amount of data by reducing gaps in the sequence and improving confidence of
- base calls within that region. seqtk then converts compressed FASTQ files to FASTA.
- 17 Rationale
- Note: Another tool, PandaSeq which does the same thing is used later, this usage of PEAR here
- is because it is more optimized for larger datasets which in this case are trimmed reads. Note:
- 20 Conversion is necessary here as some tools cannot read FASTQ files, and instead rely on FASTA
- 21 formats.

#### 2 2.1.2 Translation and Reverse Translation

- 23 Technical Notes: transeq from EMBOSS translates nucleotide sequences to protein sequences
- based on standard genetic code. backtranseq reverses this to allow iterative alignments with
- 25 nucleotide sequences.
- 26 Rationale: This leverages conserved protein-level information, which is lost at the nucleotide
- 27 level due to synonymous mutations while also increasing the sensitivity to potential ARG

proteins from k-mer alignment. See Nature Methods, doi: doi.org/10.1038/s41592-019-0437-4 (2019) for more details.

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31 Metagenomic Assembly

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#### 2.1.3 Iterative alignment of ARG-contigs

Technical Notes: KMA (K-mer Alignment) is used find the reverse-translated ARGs with the

proximity filtering option to determine the surrounding regions of ARGs. This is done iteratively

for each gene increasing the ARG-associated database.

Rationale: Firstly, KMA is used because, unlike Bowtie2 and BWA-MEM, which were created

specifically for Human metagenomics, KMA does not suffer (or suffers less) from multi-allelic

<sup>39</sup> databases. Secondly, iterative alignment using this process allows us to contextualize the region

40 wherein ARGs reside - thereby narrowing our focus onto these local regions instead of looking

at the global genomic context. Notes: The script is designed to have a cap on the number of

42 iterations KMA creates, increasing the database size.

#### Sep 22 2024 Update

This is now deprecated as there is apparently another wrapper tool called ARG Profiler that has a module called ARG Extender that literally does the same thing but with an extra filtering step that applies filters on:

- 1. % query ID
- 2. % global consensus ID
- 3. Mean read depth

and is thus more robust. So I will instead be using their module and cite them as such

```
"This study utilized the ARGextender module from ARGProfiler (Martiny, et al., 2024) to extend genomic flanking regions around ARGs."
```

Personal Note: Feels bad a bit on my part that someone already published the idea before I did but hey, at least I know this sort of idea is publishable haha. Moreover, it does save us time because I only have to copy-paste their module (and associated scripts) into the existing pipeline.

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#### 2.1.4 Merging of paired reads

- 45 Technical Notes: PANDASeq is then used to further refine the paired-reads collated in the ARG-
- 46 related genes database.
- 47 Rationale: PANDASeq was chosen as, while being slower than PEAR, it is more accurate. This mer-

- 48 gins step is included to ensure that only high-confidence reads are assembled. Note: We leverage
- the fact that the ARG-related genes database is smaller compared to the raw metagenomic reads
- 50 database.

#### 51 2.1.5 Guided Metagenomic Assembly

- 52 Technical Notes: metaSPAdes is then used to create contiguous sequences from these local regions
- 53 by extending them using reads from the whole metagenomic pool. Additionally, Contigs are
- 54 filtered by length here to remove possible artefacts.
- Rationale: metaSPAdes was chosen as, while being slower than MEGAHIT, it is optimized in
- 56 handling highly diverse and mixed microbial populations. CARD is used here because it is
- 57 manually curated and updated regularly in order to be included in the database, there must
- be clinical data (e.g. ASTs(Antimicrobial susceptibility testing)) involved in the study.
- 59 Notably, this would decrease the sensitivity of our ARGs and would mostly be biased towards
- those reported in the clinical setting. To counter this, we could also incorporate other tools such
- as ResFinder, the NCBI AMR Database, and ARG-ANNOT
- Notes: Filtering of contig length is handled by a python script called minimum\_length\_CARD.py.
- Notes: Contigs are further extended using contigextender to form scaffolds
- Notes: Might add other contig extendending programs like GapFiller which leverages mate-
- 65 pair information

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67 Contig Quality Checks

#### 69 2.1.6 Confirmation of contigs with ARGs

- 70 Technical Notes: RGI scan the contigs and check whether which contigs created by metaSPAdes
- 71 have ARGs in them.
- Rationale: metaSPAdes may have created contigs that DO NOT contain ARGs, and have instead
- 73 assembled them into a more matching contig (a false-positive misassembly) this can happen
- 74 because of the different databases being used; also parallelization of methods like this increases
- 75 robustness because it has been confirmed independently from different starting points (bottom-up
- vs top-down approach). This allows us to filter ARG-containing contigs.
- 77 Notes: RGI is the official scanner of CARD.

#### 78 2.1.7 Standard Contig Quality Metrics

- 79 Technical Notes: A custom Python script calculate\_contig\_quality.py is created to do another
- 80 round of checking contig quality for downstream analysis, R scripts (TBA) are used to visualize
- 81 the data.
- 82 Notes: The Python script will measure standard contig quality metrics: N50, L50, GC-content,
- and coverage, as well as more robust metrics: N90 and L90.

#### $^{84}$ 2.1.8 Read-mapping

- Technical Notes: Samtools is used here to map the raw reads from the larger database back to
- $^{86}$  the assembled contigs and then calculates the coverage over the entire contig.
- 87 Rationale: Read-mapping is a quality control protocol used in metagenomics, to determine the
- 98 quality of the assembly. High-coverage means that many of the k-mers align well with that
- region of the contig, while low coverage is evidence of inconsistent mapping and that the contigs
- should be refined, split, or discarded.
- Note If there are persistent (after further refinement and reassemblies) sudden differences in
- 92 coverage across a contig, that contig could be chimeric, meaning, it could be from two different
- 93 populations.

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- Extra note A Python script plot\_and\_detect\_intermediate\_coverage.py is included in the
- 95 pipeline that is determine visualize and check how the coverage changes over contig regions. In
- general, they could be interpreted as the following:
- 1. Smooth, Uniform Coverage: Typically shown by well-assembled contigs.
- 2. Sharp Coverage Drop: May need to be split or flagged for reassembly. May also be a misassembly point (chimeric contig) or caused by a structural variant.
  - 3. Coverage Gaps: Regions with little to no read support; a strong indicator of misassembly.
- 4. Gradual Drops: Overlapping reads, repetitive or duplicate regions, partial HGT, sequence heterogeneity, or coverage differences due to a mixed population. Repeats and duplications can be filtered out using tools like RepeatMasker or BLAST (TBA).
  - 5. Sharp Increase: May be due to repetitive or duplicated regions, amplification bias from PCR, HGT, SV, chimeras.

#### 106 Read mapping parameters

Technical Notes: Four (4) Tools will be used in parallel to do the read mapping process BWA Bowtie KMA, and minimap2, their parameters have been adjusted to map reads at 95 % identity to the contigs.

Mapper	K-mer Length	Mismatch Penalty	Gap Opening	Gap Extension
			Penalty	Penalty
BWA	21	5	7	2
BWA	31	4	6	2
BWA	51	3	5	1
Bowtie2	-	4,2	5,2	5,2
KMA	Default	95% identity	Automatic	Automatic
Minimap2	-	5	7,2	4,1

Table 2.1: Example table of mapper configurations without command example

Rationale: 95 % identity is used to increase sensitivity - as is standard for determining homologous sequences. This adjustment was made because k-mers are either attach or don't. Parallelization is used to increase robusness.

Note K-mer extension is used to increase the accuracy of mapping. BWA k-mer lengths can be adjusted, while KMA does it by default. The others, cannot be adjusted.

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#### Note

I chose to change the script to not allow gaps during this phase as we already used reverse translation earlier to correct for synonymous codons, and protein sequences are more important when it comes to ARG function, the new values are below. I also added protein-based read mapping.

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Tool	K-mer Length	Mismatch	Gap Opening	Gap Extension
		Penalty	Penalty	Penalty
BWA	21, 31, 51	5, 4, 3	1000	1000
Bowtie2	-	4,2	1000,1000	1000,1000
KMA	-	95% identity	Automatic	Automatic
Minimap2	-	5	1000,1000	1000,1000

Table 2.2: Alignment parameters for ungapped alignments across BWA, Bowtie2, KMA, and Minimap2

Rationale: The main rationale for adding a protein-based read mapping protocol is because
ARGs are primarily about their protein-protein interactions (biological relevance). This method
also accounts for frameshifts with higher specificity to homologous regions.

On a personal note: This approach may also require further exploration into whether proteinprotein interactions are altered—perhaps by investigating changes in binding sites. Which is a story for another day (why do I do this to myself?)

#### 123 2.1.9 Taxonomic profiling of reads and contigs

Technical Notes: Kraken2 uses k-mer-based classification to assign taxonomy based on raw-reads.

While SprayNPray complements this by assigning taxonomy at the contig level.

Rationale: By comparing their respective databases with our ARG-related databases, we will be

able to connect our reads and/or contigs to their corresponding taxa, uncovering the microbial

hosts responsible for carrying and potentially spreading ARGs in the environment.

#### 2.1.10 Detect structural variants (SVs) in contigs

Technical Notes: Manta identifies large genomic rearrangements such as insertions, deletions, and duplications.

Rationale: Chimeric contigs may be due to systematic error or real biological signals. These chimeric contigs can be detected by Kraken2 and SprayNPray (i.e., when a portion of a contig is being assigned to different taxa). The rationale behind this step is to investigate whether structural variations are present — which may be evidence of horizontal gene transfer (HGT) events.

Tool	Input Files	Key Parameters
tblastn	<ul> <li>Protein sequences:     Translated     protein     sequences     contigs</li> <li>Nucleotide     database:     Cleaned     sequence     database</li> </ul>	<ul> <li>-outfmt 6</li> <li>-evalue 1e-5</li> <li>-gapopen 5</li> <li>-gapextend 2</li> <li>-matrix BLOSUM62</li> </ul>
blastp	<ul> <li>Protein sequences:         Translated         protein             sequences             contigs</li> </ul> <li>Protein database:             Translated</li> <li>cleaned</li> <li>sequence</li> <li>database</li>	<ul> <li>-outfmt 6</li> <li>-evalue 1e-5</li> <li>-gapopen 5</li> <li>-gapextend 2</li> <li>-matrix BLOSUM62</li> </ul>

Table 2.3: Protein read-mapping parameters for tblastn and blastp

#### 2.1.11 Sketching contigs followed by calculating Bray-Curtis diversity

Technical Notes: Mash Sketch uses a MinHash approach to generate a presence/absence profile of ARGs across contigs. This gives us a quick snapshot of what the contigs "look like" in terms of ARG content. For Bray-Curtis diversity, we calculate a dissimilarity matrix from the abundance data of ARGs, followed by a PCoA plot to visualize similarities between contigs based on their ARG profiles.

Rationale: The Mash Sketch helps rapidly identify the genetic makeup of contigs in terms of ARGs, which provides a foundation for further investigation. By applying Bray-Curtis diversity and using PCoA, we can group contigs based on their ARG similarity. If contigs with the same sketch group together, we can trace them back to their taxonomic IDs to identify the microbial hosts. However, if contigs have similar ARG profiles but belong to different taxa, this could serve as evidence for Horizontal Gene Transfer (HGT). This dual approach allows us to trace ARG spread and potential HGT events in a metagenomic context.

Note: Bray-Curtis (dis-)similarity is most often used as a presence or absence diversity metric.

152 Transposition

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2.1.12 Determination of Transposons (TBA)

155 Technical Notes: Tools such as the following can be used:

- HMMER3 suite
- Tnppred a transposon predictor tool

158 Rationale:

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160 Plasmids

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#### 162 2.1.13 Determination of putative plasmids

Technical Notes: The tools listed below will be used in parallel. Plasmids are considered valid when all 4 tools predict plasmid signatures in the contig.

- PlasPredict pipeline
- Recycler
  - PlasmidFinder
  - MOBSuite plasmid marker annotator

If plasmid signatures are present, plasmidSPAdes along with GapFiller will be used to check if the contig can circularize. oriTfinder will then be applied to contigs with fewer than 4 fragments. A Python script (TBA) will calculate GC skews of the chimeric contig and compare it to its taxonomic counterparts. Another Python script (TBA) will normalize the data according to 16S rRNA from trimmed reads. Lastly, plasmid percentage will be calculated based on reads mapped to putative plasmids over the total reads. A code snippet is present called calculate\_plasmid\_percentage.py.

Equation:

$$\label{eq:Plasmid Reads} \begin{aligned} \text{Plasmid Reads} &= \left(\frac{\text{Plasmid Reads}}{\text{Total Reads}}\right) \times 100 \end{aligned}$$

Rationale: oriTfinder looks for origin of transfer sites (oriT), which are characteristics of conjugative plasmid . This whole sub-pipeline is to look for evidence of conjugative plasmid transfer as the cause of these chimeric contigs. Normalization and percentage counts are used here to further check whether these "plasmids" align with our understanding of the average plasmid copy number.

Note: Will also be drafting a script (TBA) to do sliding window analysis of GC-skews - as different characteristics of this curve can be interpreted in different ways.

185 Phages

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#### 2.1.14 Phage influence signatures

- 188 Technical Notes: They will be determined using a variety of tools:
- After prediction with Prodigal, a sequence was considered a phage if it was identified by two-thirds (2/3) of the program stated below:
- VirSorter: Identifies viral signatures within microbial genomes and separates prophages from bacterial sequences.
- PHASTER: A web-based tool for phage search and annotation, identifying integrated prophages.
- VIBRANT: A tool that combines several approaches to identify and annotate phage elements in metagenomic sequences.

Rationale:This analysis aims to detect potential phage signatures in the chimeric contigs.
Since phages are mobile genetic elements, their involvement in transferring ARGs through transduction is highly relevant. Phages, especially temperate phages, can integrate into bacterial genomes and excise themselves, sometimes carrying host genetic material, such as ARGs, with them. The integration and excision signatures detected in contigs will provide evidence of possible transduction events in our datasets, supporting the hypothesis of ARG dissemination via phages.

#### 203 2.1.15 Annotion of Phage genes (TBA)

<sup>204</sup> Technical Notes All these proteins will be queried against the following databases.

- 205 1. PFAM
- 206 2. VOGdb
- 3. eggNOG
- Using a combination of eggNOG-mapper(mapper.py) and HMMER with the following thresholds: 208 E-value  $< 10^{-5}$ , score  $\ge 50$ . Active prophages were then separated from Inactive ones using 209 the tool Prophage Hunter - default scoring parameters. Results were considered false-positives 210 if they were considered active by Prophage Hunter but were 'not phage' for VirFinder and MetaPhinder as previously done by the authors of the tool. HMM profiles were downloaded 212 from proMGE database – which are calibrated to different recombinases: (huh y1, huh y2, 213 ser tn, ser ce, ser lsr, cas1) and used against the putative recombinases to further divide them 214 into distinct categories. To determine whether these genes are of viral or bacterial origin, CheckV 215 was used. 216
- 217 Rationale
- 218 Notes

#### 219 2.1.16 Phage Signature Extraction and Phylogenetic Analysis

Technical Notes: Phage-associated genes will be extracted from the chimeric contigs, followed by phylogenetic analysis to uncover evolutionary relationships. FastTree will be used to build

<sup>222</sup> a phylogenetic tree based on the extracted phage genes. For visualization, tools like iTOL or <sup>223</sup> FigTree can be used to generate an interpretable phylogenetic tree.

224 Rationale: Phage genes embedded in chimeric contigs (their taxonomy) may serve as strong

evidence of horizontal gene transfer (HGT) events. The aim here is to check the evolutionary ori-

226 gins of the phage genes found in our dataset and their potential involvement in the dissemination

of ARGs.

#### 228 2.1.17 Directory tree

Notes: The final directory tree should look like this to help you visualize. Notice the multiple path/to/ here as this is still (WIP):

/	project_root/
į	path/to/cleaned_reads/Prerequisite
	path/to/merged_reads/
	path/to/CARD_db/Prerequisite - Database
	path/to/output/
	final_annotated_plasmids.fasta
	fpkm_normalized_plasmids.txt
	categorized_MGEs.txt
	final_filtered_contigs.fasta
ı	path/to/logs/
	predict_plasmids.log
	run_mob_suite.log
	check_plasmidfinder.log
	calculate_gc_skew.log
,	path/to/plaspredict_output/
,	path/to/PFAM_db/Prerequisite - Database
,	path/to/TnpPred_db/Prerequisite - Database
,	path/to/plasmid_prediction/
	predicted_plasmids.fasta
	plasmidfinder_report.txt
	oritfinder_report.txt
	tnpred_report.txt
,	path/to/plasmidspades_output/
	plasmid_assembly.fasta
,	path/to/mob_suite_output/
	mob_suite_summary.txt
ı	path/to/gc_skew_analysis/
	gc_skew_plot.png
,	path/to/read_mapping/
	reads_mapped_to_plasmids.bam
	plasmid_read_count.txt
	total_read_count.txt

232 Homework?

231

#### 233 2.1.18 Future Considerations

Will continue improving this section (everything regarding HGT) by evaluating the results from these tools, aligning them with ARG presence, and refining the approach for identifying conjugation, transposon, and transduction events within chimeric contigs. This may also involve validating phage activity—again, why do I do this to myself?

#### 1 2.2 metagenomics\_general.smk

- 2 Stage: Done
- 3 General Purpose taxo-metagenomics
- <sup>4</sup> Specifics: This pipeline is designed for The essentials in metagenomics, which includes quality
- 5 checking, filtering, and trimming of raw reads to clean reads. As well as the usual taxo-
- 6 metagenomic analysis.

#### <sup>7</sup> 2.2.1 Quality control (Pre-processing)

- 8 Raw trimming of raw metagenomic data
- 9 Technical Notes: FastQC is used on raw reads
- Rationale: This is mainly used as a point of comparison determine whether the next step (trim-
- ming) was effective. This pre-processing step is the starting point in any and all metagenomics
- 12 pipelines.
- Notes: This whole quality control steps are interconnected with each other.

#### 14 2.2.2 Trimming

- 15 Technical Notes: Trimmomatic is used on raw reads
- 16 Rationale: Trimming involves removing low quality bases (often depending on something called
- 17 the Phred Score which is just a measure of how "confident" we are that the base on that site
- was accurate), adapters, and filtering reads that go below a specific length threshold.
- Notes: Journals often report the parameters on Trimmomatic (or any trimmer they decide to
- 20 use); this is often so that the study is reproducible, should one decide to actually reproduce the
- 21 study starting from scratch (raw reads).
- Notes: There are many different trimmers each with their own strengths and weaknesses,
- 23 Trimmomatic is just the most popular trimmer and is thus used here, though studies differ in
- 24 the parameters they used for trimming which often dictates how strict they are with what they
- define as "good enough".

26

37

- 27 Perspective: Why different people choose different trimmers depend on the strengths and
- weaknesses of the trimmer e.g. trimmers like "fastp" is used because it's fast making it suitable
- 29 for very large datasets like deep sequencing. While some trimmers like Sickle has automatic
- 30 adjustment over the entire sequence which makes it useful for very ancient datasets where DNA
- 31 is often highly-degraded. Other times, it's for convenience like Trim-Galore which combines
- 32 FastQC and Cutadapt trimmer in a single command. Another good example is BBDuk which is
- part of a larger package called BBtools, BBDuk also has an built-in contamination detection -
- 34 so it's particularly good at filtering out usual contaminants like sequences known to be from
- the human genome. so you can simply just use all the modules in that package for all-in-one
- processing. Other times, it's just familiarity.
- 38 Quality checks of post-trimmed data
- Technical Notes: FastQC is used on trimmed reads to determine how effective the trimming

- o process was.
- Rationale: If the trimming process was effective, we should notice a better quality reads here,
- otherwise, we might have to adjust the trimming parameters.
- Notes: Determination of whether the trimmed reads are "clean enough" is more of an art rather
- than actual science, though thresholds exist like Phred > 20 or Phred > 30 depending on how
- strict you are as a researcher.

46

48 Processing cleaned reads

49

56

#### 50 2.2.3 Metagenomic Taxonomy

- 51 Taxonomy from Cleaned Reads
- 52 Technical Notes: Kraken2 is used on raw reads to determine which species they came from.
- Rationale: Taxonomy based on reads is standard on metagenomics instead of using assembled
- 54 contigs because information is lost during the assembly process. By using cleaned reads we make
- 55 sure that:
  - 1. We are maximizing the amount of data
- 57 2. We are not being biased by low quality reads
- Notes: There are many ways in which taxonomy is assigned to raw reads the Kraken-based
- 59 packages use a curated database that links k-mers from your database to k-mers generated from
- their database (usually manually curated).
- 61 Notes: Others like MetaPhlan first create "markers" that are based off of known sequences, and
- 62 then scan your raw reads for these markers, which it then checks for under what taxon/taxa
- 63 that marker fell under.

64

#### 55 2.2.4 Diversity analysis

- 66 Diversity analysis per site or sample
- 67 Technical Notes: Here Bracken an extension of the Kraken packages or Qiime2 are often used
- 68 to calculate diversity. Instead I opted to create my own Python script calculate\_diversity.py
- 69 because:
- 1. I find that the diversity indices in either tools are limited to only the most often used i.e.
  the most popular indices so it is not comprehensive
- 2. I've had problems integrating them into the pipeline because some dependency limitations, un-updated scripts, and a pre-processing step that requires converting all of Kraken2,
- Bracken files, then importing them to Qimme2 which is too time-consuming and inefficient
- my script just automatically calulates from Bracken outputs and just puts out all the
- possible (non-phylogenetic-based-which you would need a phylogenetic tree to build first)
- indices out there.

- $\,$  Rationale: Taxonomy based on reads is standard on metagenomics instead of using assembled
- 79 contigs because information is lost during the assembly process. By using cleaned reads we make
- 80 sure that:
- 1. We are maximizing the amount of data
- 2. We are not being biased by low quality reads
- 83 Notes: There are many ways in which taxonomy is assigned to raw reads the Kraken-based
- packages use a curated database that links k-mers from your database to k-mers generated from
- 85 their database (usually manually curated).
- Notes: Others like MetaPhlan4 first create "markers" that are based off of known sequences,
- and then scan your raw reads for these markers, which it then checks for under what taxon/taxa
- that marker fell under.

#### $_{89}$ 2.2.5 Directory tree

```
/project_root/
  configs/
    config.yaml
  sample1_R1.fastq.gz
    sample1_R2.fastq.gz
    sample2_R1.fastq.gz
    sample2_R2.fastq.gz
  path/to/trimmed_reads_dir/ ...... Prerequisite - Trimmed Data
    sample1_R1_paired.fastq.gz
    sample1_R2_paired.fastq.gz
    sample2_R1_paired.fastq.gz
    sample2_R2_paired.fastq.gz
  path/to/fastqc_output_dir/
  path/to/kraken_output_dir/
    sample1.k2report
    sample1.kraken2
    sample2.k2report
    sample2.kraken2
  path/to/bracken_output_dir/
    sample1.bracken
    sample1.breport
    sample2.bracken
    sample2.breport
  path/to/cleaning_results_dir/
    summary_report.txt
  logs/
    calculate_diversity.log
```

\_\_path/to/output/ \_\_diversity\_matrices.tsv

#### <sub>1</sub> 2.3 trim randomizer.smk

- <sup>2</sup> Stage: Done
- 3 Purpose: Randomization of trimming parameters
- This is a module that will be part of a bigger pipeline. The idea here to is randomize
- parameters in a variety of trimmers in this case Trimmomatic, fastp, CutAdapt, BBDuk, and
- 6 Sickle famous bioinformatics trimming tools. Trimmomatic and Sickle in particular are
- 7 widely used in Illumina-based data.

8

#### 9 2.3.1 Random Parameter Generation

- 10 Technical Notes: Random parameters are generated for each trimming tool (Trimmomatic, Fastp,
- 11 Cutadapt, BBDuk, Sickle). This is done using the random module, which creates random values
- 12 for parameters such as quality scores, read length, adapter sequences, and error rates. These
- parameters are stored in the generated\_parameters dictionary to ensure consistency across
- 14 iterations for each sample.
- 15 Rationale:By randomizing parameters, the workflow allows for testing different parameter sets
- 16 across multiple iterations to find optimal settings for trimming and quality control.
- 17 Notes: This approach helps with parameter exploration, particularly when you are unsure which
- trimming settings will give the best results. The randomness provides variability, which can
- 19 highlight which parameters consistently lead to good results.

#### 20 2.3.2 Log Parameters to a TSV File

- 21 Technical Notes: Each set of generated parameters is logged into a separate TSV file (e.g.,
- trimmomatic\_params.tsv, fastp\_params.tsv). The file headers are written only once, and
- parameters are appended as the trimming steps proceed. This is done in a structured way so
- 24 that you can track the exact parameters used for each sample and iteration.
- 25 Rationale:Logging ensures reproducibility and transparency in bioinformatics workflows. Having
- 26 a record of all the parameter values used in each iteration is crucial for comparing results and
- 27 for future reference
- 28 Notes: This practice is a standard in scientific workflows where random parameter generation is
- 29 involved. It helps maintain a clear audit trail of the steps performed and aids in troubleshooting
- or refining workflows later.

#### 2.3.3 Define **Rules** for Each Tool

- Technical Notes: The script uses Snakemake rules to define separate rules for each tool which include
- 1. Input folder (as the script is designed to go through all the FASTQ samples within the folder)
- 2. Output folder, where the processed files will be saved (e.g., trimmed paired and unpaired reads). The script is designed to keep all the trimmed reads in separate files.

- 3. Params: Fetches the parameters to be randomized.
- 39 Rationale: Using Snakemake here allows for parallel execution of the workflow. This parallelization
- 40 if very important as the generation of random paramaters is created using a random seed. Using
- a random seed like this allows us to replicate what the parameters that had the optimal results
- were by tracking down what seed was assigned as dictated in the TSV file.
- Note keep in mind that this will create a large number of folders if you decide to iterate many
- times as each iteration, per tool, will have its own folder full of trimmed reads, per sample/site.
- <sup>45</sup> 2.3.4 Interpretation and Analysis of Results (TBA)
- 46 Technical Notes: Once all iterations have completed, the trimmed files can be analyzed to
- 47 determine which parameters led to the best results in terms of quality and length distribution of
- reads. There are many different metrics that can be used to interpret the results, including (but
- 49 not limited to)
- 1. Quality Scores significant differences in quality among sites and across entire sequences
  (Phred score, Contamination, Adapter Removal, N Content, Length Distribution, etc.).
  Can be done using tools like FastQC
- 2. Visualization can be done using Rstudio or Python's matplotlib to visually look for differences in abnormalities.
- Rationale:Evaluating read quality and assessing key metrics post-trimming helps to ensure that the data is suitable for downstream analyses. Optimal trimming should maximize the number of
- 57 high-quality, usable reads while eliminating low-quality bases and adapter contamination.
- Notes See Box on Biological Information for more possible details on this.
- <sub>59</sub> 2.3.5 Annotated Directory Tree with File Temporary files and Prerequisites

```
/project_root/
  raw_reads/ ......(permanent) (prerequisite)
    sample1 R1.fastq.gz .....(permanent) (prerequisite)
    sample1_R2.fastq.gz .....(permanent) (prerequisite)
    sample2 R1.fastq.gz .....(permanent) (prerequisite)
    sample2_R2.fastq.gz .....(permanent) (prerequisite)
  output_dir/
    fastp_output/ .....(temporary) (to be deleted)
      iteration_1/
        sample1_R1_fastp_trimmed.fastq.gz .....(temporary)
        sample1_R2_fastp_trimmed.fastq.gz ......(temporary)
        sample2_R1_fastp_trimmed.fastq.gz ......(temporary)
        sample2_R2_fastp_trimmed.fastq.gz .....(temporary)
      iteration 2/
        (same as iteration_1 but with iteration_2 files) ......(temporary)
      iteration 3/
```

☐ (same as iteration_1 but with iteration_3 files)(temporary)
trimmomatic_output/(temporary) (to be deleted)
iteration_1/
sample1_R1_paired.fastq.gz(temporary)
sample1_R1_unpaired.fastq.gz(temporary)
sample1_R2_paired.fastq.gz(temporary)
sample1_R2_unpaired.fastq.gz(temporary)
(same for sample2)(temporary)
iteration_2/
(same structure as iteration_1)(temporary)
iteration_3/
(same structure as iteration_1)(temporary)
cutadapt_output/(temporary) (to be deleted)
iteration_1/
sample1_R1_cutadapt_trimmed.fastq.gz(temporary)
sample1_R2_cutadapt_trimmed.fastq.gz (temporary)
iteration_2/
iteration_3/
(same structure as iteration_1)(temporary)
bbduk_output/(temporary) (to be deleted)
iteration_1/
sample1_R1_bbduk_trimmed.fastq.gz(temporary)
sample1_R2_bbduk_trimmed.fastq.gz(temporary)
iteration_2/
(same structure as iteration_1)(temporary)
iteration_3/
(same structure as iteration_1)(temporary)
sickle_output/ (temporary) (to be deleted)
iteration_1/
sample1_R1_sickle_trimmed.fastq.gz(temporary)
sample1_R2_sickle_trimmed.fastq.gz(temporary)
sample1_singles_sickle_trimmed.fastq.gz(temporary)
iteration_2/
(same structure as iteration_1)(temporary)
iteration_3/
(same structure as iteration_1)(temporary)
_ logs/(permanent) (prerequisite)
fastp_params.tsv(permanent) (prerequisite)
, , , , , , , , , , , , , , , , , , , ,
trimmomatic_params.tsv(permanent) (prerequisite)
cutadapt_params.tsv(permanent) (prerequisite)
bbduk_params.tsv(permanent) (prerequisite)

\_\_sickle\_params.tsv ......(permanent) (prerequisite)

#### <sup>1</sup> 2.4 bootstrapping\_rawreads.smk

- <sup>2</sup> Stage: Further refinement
- Purpose: Bootstrapping the taxo\_metagenomic pipeline itself
- Bootstrapping is the process of randomly selecting from a pool of samples (with replacement)
- 5 and using that in a specific process you want to bootstrap. This is a standard method used
- in molecular phylogenetics to determine the robustness of trees where a > 70 support from
- <sup>7</sup> bootstrapped data is considered robust enough.

#### The principle of bootstrapping in phylogenetics

Phylogenetics uses this statistical technique because (in principle) it effectively means that removing parts of the entire sequence does not alter the topology of the tree.

Here I'm adapting this method with Kraken2's taxonomic profiling to see whether the taxonomic support of its k-mer assignment is also consistent even with changes in the sampling sites.

This is still WIP because I plan to go step further and start bootstrapping the raw reads themselves to see if changes in reads changes the topology of taxonomic assignment.

#### 2.4.1 Directory setup of temporary files

- Technical Notes:Snakemake starts by ensuring the existence of necessary directories. Most
- 11 notably, the temporary bootstrap directories.
- 12 Rationale: Bootstrapping is sometimes done iteratively thousands of times, so making this a
- 13 temporary directory helps manage space.
- 14 Notes:

#### 15 2.4.2 Sample Identification

- 16 Technical Notes: The workflow identifies sample names by parsing filenames in the raw reads
- 17 directory.
- 18 Rationale: Inclusion of these in the script allows the user to flexibly configure the naming
- convention and the directory in which they want to bootstrap.
- Notes: Presently it looks for \_R1.fastq.gz and it's associated pair, \_R2.fastq.gz in the raw\_reads
- 21 directory.

#### 22 2.4.3 Bootstrapping

- 23 Technical Notes: This executes bootstrap\_reads.py in the scripts directory to start bootstrap-
- 24 ping the paired-end reads and outputs them in the temporary folders I mentioned earlier.
- 25 Rationale: Moving bootstrapping logic into a Python script leverages its ability to create ran-
- domizations from its libraries. Also it allows us to define a seed so it is reproducible (if you
- 27 want to reproduce) the results anyway.
- Note: The number of bootstraps and the fraction of samples you want to retain can be controlled

in the config.yaml file in the configuration folder.

30

#### Update Sep 19 2024

On a personal note: Before writing this part of the document, I decided to change the bootstrapping rule. It used to rely on a simplified approximation. The probability of not being selected after N independent draws from a sample of size N is given by:

$$P = \left(1 - \frac{1}{N}\right)^N$$

This is a well-known mathematical equation describing the probability of not selecting a sample at least once in N draws. As  $N \to \infty$ , this probability approaches:

1

Previously, I used the probability of a sample being selected, which is:

$$1-\frac{1}{e}$$

This was used as an approximation for bootstrapping by shuffling and adjusting the sample size. However, the updated script now performs actual bootstrapping, sampling with replacement, which is a more accurate statistical method for resampling.

31

#### 2 2.4.4 Annotated Directory Tree with File Movement and Prerequisites

```
/project_root/
 configs/
 config.yaml (prerequisite)
 path/to/raw_reads_dir/ ...... (permanent) (prerequisite)
   sample1_R1.fastq.gz .....(permanent) (prerequisite)
   sample1_R2.fastq.gz .....(permanent) (prerequisite)
   sample2_R1.fastq.gz .....(permanent) (prerequisite)
   sample2_R2.fastq.gz .....(permanent) (prerequisite)
 path/to/temp_bootstrap_dir/ .... (temporary) (to be deleted after pipeline resolves)
   sample1/ ..... (temporary)
    rep_1_R1.fastq.gz .....(temporary)
    rep_1_R2.fastq.gz .....(temporary)
    rep_2_R1.fastq.gz ......(temporary)
    rep_2_R2.fastq.gz ......(temporary)
    total reads.txt ......(temporary)
   sample2/ ..... (temporary)
    rep 1 R1.fastq.gz .....(temporary)
    rep_1_R2.fastq.gz .....(temporary)
    _rep_2_R1.fastq.gz .....(temporary)
```

```
rep_2_R2.fastq.gz ......(temporary)
    total_reads.txt .....(temporary)
  rep_1/ ..... (temporary)
    diversity_matrices_sample1_rep_1.tsv ......(temporary) (moved to
    diversity_results/)
    diversity_matrices_sample2_rep_1.tsv .....(temporary) (moved to
    diversity_results/)
  rep_2/ ..... (temporary)
    diversity_matrices_sample1_rep_2.tsv .....(temporary) (moved to
    diversity_results/)
    diversity matrices sample2 rep 2.tsv .....(temporary) (moved to
    diversity results/)
logs/ ...............(permanent) (prerequisite)
  metagenomics_pipeline_sample1_rep_1.log ......(permanent) (prerequisite)
  metagenomics_pipeline_sample1_rep_2.log ......(permanent) (prerequisite)
  metagenomics_pipeline_sample2_rep_1.log ......(permanent) (prerequisite)
  metagenomics pipeline sample2 rep 2.log ......(permanent) (prerequisite)
diversity_results/ .....(permanent) (contains moved files) (prerequisite)
  diversity_rep_1.tsv ......(moved from temp_bootstrap_dir)
  diversity_rep_2.tsv ......(moved from temp_bootstrap_dir)
aggregated_results/ .....(permanent) (prerequisite)
  diversity_aggregate.tsv .....(permanent)
```

#### $_{ exttt{3}}$ 2.4.5 Run kraken\_pipeline.bash

- Technical Notes: This Shellscript (or bash file) is used to automate the processing of all files from the bootstrapping. It runs them under Kraken2 then Bracken to generate taxonomy profiles for all of them. It also creates a log file for each replicate to provide traceability and error checking,
- 37 helping diagnose any issues with specific replicates or samples.
- Rationale: Read why I'm bootstrapping from the textbox above. The reason why I also included
- 39 Bracken and measurements diversity metrics in the analysis per sampling replicate is so we
- can analyze how the topology of diversity also changes similiar to how we look at topology of
- 41 phylogenetic trees. The final process consolidates all the diversity metrics (alpha diversity) and
- 42 matrices (beta diversity) into a TSV file.
- Note: The decision to use Snakemake and Shellscripts here is so that the bootstrapping comes
- 44 first before the pipeline is introduced. Otherwise Snakemake will run the entire thing in parallel,
- taking up so much memory because it runs Kraken2 for every single sample instead of doing it
- in batches which takes up so much unnecessary time.

#### 2.5 Binning.smk

- <sup>2</sup> Stage: To test on higher coverage data
- 3 Binning pipeline to create high quality (Metagenome Assembled Genomes) MAGs
- <sup>4</sup> Specifics: This pipeline passes through multiple quality checks during binning of using a variety
- of tools (both wrappers and modules) including MetaWrap, DasTool, CheckM2, MagPurify etc.

#### 6 2.5.1 Universal Configurations

- 7 Technical Notes: The pipeline allows the user to configure settings they want for the binning
- process. By default, the settings are Minimum Contig Length (2500bp), Completeness (50%),
- 9 Contamination (10%).
- Rationale: The default settings were curated by me and the reason I chose them is because of
- 11 the following
- 1. Longer contigs tend to represent more complete genomic fragments. Setting a threshold of 2500bp ensures better assembly quality. Others prefer a lower threshold for more sensitivity like 2000 bp.
- 2. Completeness 50% and Contamination 10% are actually based from a standard called the MIMAG standards.
- Notes: Making configurations universal this way creates consistency across the script, i.e. when specifically asked by a specific tool, this returns a universal parameter.
- Notes: Additionally, the user can specify the memory usage and number of threads they want
- to allocate per tool as well as other tool-specific parameters at the top of the script for ease of
- use. I plan to add this to the configuration file soon once I have tested the file to be working
- 22 at higher coverage since you can't make high quality bins with low read counts and my PC
- 23 can't practically handle that sorry.

#### 24 2.5.2 **FastUniq** (Deduplication)

- <sup>25</sup> Technical Notes: FastUniq used to remove duplicate reads.
- 26 Rationale: Since we are focused on creating MAGs or genomes based on populations of genomes,
- 27 removing duplicates is less risky during binning and is thus included. It also allows us to
- 28 completely remove amplification bias from PCR reactions. Moreoever, deduplication reduces
- <sup>29</sup> redundancy and thereby memory usage downstream.

#### 31 2.5.3 Seqtk (FASTA Conversion)

- 32 Technical Notes: Seqtk converts FASTQ to FASTA.
- Rationale: This conversion prepares sequences for tools that require FASTA inputs, such as
- 34 CD-HIT-EST.

30

35 Notes: I used to include a decompress-then-compress mechanism in the script to minimize

- memory storage but according to my calculations from the sequencing facility quotations, re-
- compressing may actually be more costly when done throughout the pipeline. Hence, it should
- be more cost-efficient to start compressing files ones the bins are done.

#### $_{39}$ 2.5.4 **CD-HIT-EST** (Clustering) at Identity: 90%

- Technical Notes: CD-HIT-EST clusters similar sequences at 90% identity.
- 41 Rationale: Clustering reduces redundancy in the contig data while maintaining closely related
- 42 sequences.

47 48

- Notes: I chose 90% ID because that's what I often see in published journals that is all Perhaps,
- 44 the 90% identity threshold balances removing duplicates while preserving diversity and that
- higher thresholds would result in fewer clusters but might oversimplify the data.
- 46 Fine, I'll make it my homework assignment why this specific threshold is used (TBA).

49 Bin Refinement

#### 51 2.5.5 MetaWRAP Binning and Reassembly

- Technical Notes: MetaWRAP is what is known as a wrapper program meaning it makes use of
- other tools as its modules. For the binning process in particular it uses 3:MetaBAT2, MaxBin2,
- and CONCOCT. Each binning process goes through internal quality control checks and the one
- 55 with the best bin-qualities are selected. It also has a reassembly feature wherein it reassembles
- 56 the contigs again to try and further refine the bins.
- 57 Rationale: Using 3 binners in parallel and choosing the best bins, quality checking, and then
- reassembling (not-so-good) bins make the binning process very robust, creating very refined bins
- not sponsored by the way, talking as a fellow researcher.
- Notes: No moving forward, the process of further refinement of bins seems redundant. But do
- 61 note that I have checked and validated that the process used in refining and checking by the
- 62 tools used here cover different metrics and therefore can be seen as parallel processes.

#### 63 2.5.6 DAS Tool (Bin Refinement)

- 64 Technical Notes: DAS\_Tool is very similar to MetaWrap in that they both choose the best bins
- 65 from a pool of bins from different binners (in this pipeline DAS Tool is designed to ALSO use
- 66 information from MetaBAT2, MaxBin2, and CONCOCT outputs to improve binning)
- 67 Rationale: However, as rationale for including it, is that it focuses more on single-copy gene (SCG)
- 68 analysis. In contrast, in MetaWrap, bins are evaluated using completeness and contamination
- 69 thresholds.
- 70 Notes: Notably, in the checking phase of this pipeline we will not be using DAS Tool for SCG
- <sub>71</sub> analysis. It is optimized for refining bins not quality checking bins. Instead we will use a more
- updated and optimized software for the latter called BUSCO.

- <sup>73</sup> 2.5.7 MAGpurify (Contamination Removal)
- Technical Notes: MAGpurify is also a bin refiner, in a sense that it uses several modules to
- detect and prune contamination in genome bins. It also uses other metrics to define bin quality
- <sub>76</sub> specifically it looks for differences in
- 1. Phylo-markers,
- 78 2. Clade-markers,
- <sup>79</sup> 3. Tetranucleotide-frequencies,
- 4. GC-content,
- 5. and then removes known-contaminants from it's manually curated database (created back in 2013)
- Rationale: This step improves genome quality by removing low-confidence contigs or contamina-
- tion from other taxa. Each module targets different contamination types (phylogenetic, clade,
- 85 etc.).

89 90

- Notes: Similar to DAS\_tool, MAGPurify is relatively old (in the bioinformatics world where new
- tools are being published every day). So in checking our bins we will be using more recently
- 88 updated tools.
- 91 Quality checking
- 93 2.5.8 MetaQUAST (Assembly Quality Assessment)
- Technical Notes: MetaQUAST assesses the quality of genome assemblies via the following:
- 1. N50 and L50 to determine contiguity
- 2. Number of contigs to determine fragmentation
- 3. GC content since a single MAG should have a constant GC content across its entirety (usually)
- 4. Alignment to a reference sequence
- Additionally, it also detects other metrics using modular tools
- 5. structural variations (requires GRIDSS)
- 6. presence or rRNA (requires SILVA)
- 7. Conserved gene sets (requires BUSCO)

Rationale: This step quantifies the completeness and accuracy of the assembled genomes, and is updated frequently.

Notes: Using reference genomes improves the accuracy of the assessment, but it's optional if references are unavailable.

108

#### A Personal Note

Personal Note: As of this writing, BUSCO has updated beyond the version required by QUAST (BUSCO od9). Unfortunately, this version is not available in the archives (you'll encounter a 404 error). Likewise, SILVA and GRIDSS frequently update. I recommend downloading each separately and manually linking their databases to avoid potential issues with QUAST's download management. Good luck and have fun!

109

#### 2.5.9 dRep (Dereplication)

111 Technical Notes: dRep dereplicates genomes by clustering them based on similarity.

Rationale: Dereplication reduces redundancy in the assembled genome data, ensuring unique genome representations. Basically CD-HIT but for whole genomes.

Notes: FastANI is utilized for quick, precise clustering, and is part of the dRep package. It defaults to a 95% ANI (Average Nucleotide Identity) threshold, a common yet somewhat subjective metric used to determine microbial species boundaries. This is often sufficient for microbial genomes due to their high gene density, frequently organized in operons. However, it may not be as suitable for eukaryotic genomes, which are laden with repetitive elements.

A Personal Note

Personal Note: I cannot find a newer version of either dRep or FastANI (both dating back to 2013, basically ancient in bioinformatics terms). There's a newer version called pyani, which is available in Bioconda, that might be a good replacement. However, I still need to reverse-engineer its source code to fully understand how it operates. Might be worth trying!

119

120

#### 2.5.10 CheckM2

Technical Notes: CheckM2 is used to predict genome completeness and contamination using low-memory mode (essential for resource-limited systems like mine; remember to adjust this on HPC systems). Rationale: CheckM2 is an updated version of CheckM, but many tools in this pipeline haven't been updated to recognize it. I haven't tested whether aliasing CheckM2 as CheckM would work, so it's added here as a final step to ensure the results meet MIMAG standards for completeness and contamination.

2.5.11 Annotated Directory Tree with File Movements and Temporary Files

```
/project_root/
__trimmed_reads/ ......(prerequisite)
```

	site1_R1_paired.fastq.gz(permanent)
	site1_R2_paired.fastq.gz(permanent)
	site2_R1_paired.fastq.gz(permanent)
	site2_R2_paired.fastq.gz(permanent)
ļ	fastuniq/(temporary) (to be deleted)
	site1_R1_uniq.fastq(temporary)
	site1_R2_uniq.fastq(temporary)
	site2_R1_uniq.fastq(temporary)
	site2_R2_uniq.fastq(temporary)
ļ	fasta/ (permanent)
	site1_R1_clean.fasta(permanent)
	site1_R2_clean.fasta(permanent)
	site2_R1_clean.fasta(permanent)
	site2_R2_clean.fasta(permanent)
ļ	megahit_output/(permanent)
	site1_assembly/
	site1_filtered_contigs.fa(permanent)
	site2_assembly/
	site2_filtered_contigs.fa(permanent)
	cdhit_contigs/(permanent)
	site1_cdhit_contigs.fasta(permanent)
	site2_cdhit_contigs.fasta(permanent)
ļ	CLEAN_READS/ (permanent) (moved)
	site1/
	site1_1.fastq(moved from fastuniq)
	site1_2.fastq(moved from fastuniq)
	site2/
	site2_1.fastq(moved from fastuniq)
	site2_2.fastq(moved from fastuniq)
ļ	binning_output/(permanent)
	site1_binning_output/
	concoct_bins.scaffolds2bin.tsv(permanent)
	metabat2_bins.scaffolds2bin.tsv(permanent)
	maxbin2_bins.scaffolds2bin.tsv(permanent)
	site2_binning_output/
	(same as site1)(permanent)
ļ	dastool_output/(permanent)
	DAS_Tool_bins_site1(permanent)
	DAS_Tool_bins_site2(permanent)
	reassembly_output/(permanent)
	site1_reassembly_output/
	assembly.fasta(permanent)

```
site2_reassembly_output/
  \_assembly.fasta ......(permanent)
magpurify_output/ ......(permanent)
 site1_cleaned.fasta .....(permanent)
 site2_cleaned.fasta .....(permanent)
metaguast output/ ..... (permanent)
 site1_metaquast_output/
   (MetaQUAST results) ......(permanent)
 site2_metaquast_output/
   (MetaQUAST results) ......(permanent)
dereplication_output/ ......(permanent)
 \_\mathtt{dereplicated\_genomes}/
  _dereplicated_genomes.fasta ......(permanent)
  _ checkm2_output/ ..... (permanent)
   __quality_report.tsv .....(permanent)
logs/ .....(permanent)
 run_megahit_site1.log ......(permanent)
 run_cdhit_site1.log ......(permanent)
 (logs for each step) ......(permanent)
```

#### 128 2.5.12 Alternatives

Vamb employs a multisplit approach wherein individual replicates (of assembled contigs) are first concatenated before performing binning. This approach can potentially enhance MIMAG standards by leveraging shared information across replicates, improving the completeness and quality of bins. By pooling data from replicates, the method also helps cancel out random noise.

This concept is analogous to image stacking in signal processing, where shared signals become more pronounced and differences or inconsistencies are easier to identify (see box on signal averaging)

One can visualize these improvements using Manhattan plots across the contigs, where lower variability leads to higher E-values. Here, E-values are used instead of P-values because they incorporate the statistical significance with respect to the reference database. The same principle applies when comparing coverage depths during read mapping on MAGs (Metagenome-Assembled Genomes). A higher read depth (more reads mapped back to the same site) increases confidence in the result, reinforcing the quality of the assembly and binning.

129

### 2.6 krakenpipeline.bash

- <sup>2</sup> Stage: Done
- The entire basic taxo-metagenomic pipeline with the usual tools

- 4 Specifics: This pipeline passes through a loop between FastQC and Trimmomatic trimming\_cleaning\_checking
- before progressing to Kraken2 then Bracken then calculate\_diversity.py.
- Notes: This does not include FastQC-ing of raw reads (yet), you have to run raw.bash for that.
- Additionally, it also currently does not automate the production of aggregated summaries of the
- 8 FastQC files for you (yet), you have to run summary\_stat.bash for that as well.

#### 9 2.6.1 Directory tree

```
/project_root/
  sample1_R1_paired.fastq.gz
    sample1_R2_paired.fastq.gz
    sample2_R1_paired.fastq.gz
    sample2_R2_paired.fastq.gz
  kraken_db/ ...... Prerequisite - Kraken2 database directory
    database files (.k2d, etc.)
  kraken_output/ ...... Output directory for Kraken2 reports
    sample1.k2report
    sample1.kraken2
    sample2.k2report
    sample2.kraken2
  bracken_output/ ...... Output directory for Bracken reports
    sample1.bracken
    sample1.breport
    sample2.bracken
    sample2.breport
```

### <sup>1</sup> 2.7 raw.bash; completely optional

- <sup>2</sup> Stage: Done
- 3 FastQC Automation script to run FastQC on raw reads
- 4 Specifics: This script takes raw reads and creates a summary report for each of them in
- 5 FasQC\_raw/
- 6 Notes: summary\_stat.bash can already do that for you, so it's a bit redundant. But if you're
- only interested in seeing QC from the raw reads, then have fun!
- 8 Personal notes: The only reason I am keeping this alive is because I am not quite sure yet if
- some pipelines rely on this and it's corresponding output files; I have to check.

### <sub>1</sub> 2.8 summary\_stat.bash

- <sup>2</sup> Stage: Done
- 3 FastQC Automation script
- 4 Specifics: This script takes reads from the raw reads directory and the trimmed reads directory,

- then creates a summary directory (if not already existent) (you can manually set this as user).
- 6 The summary directory will contain summary statistics extracted from FastQC sub-directories,
- 7 it also creates. It has two notable functions:
- 1. Counts the number of reads in raw and trimmed directories.
- 2. Extract quality metrics from FastQC subdirectories specifically: per base sequence quality, and per sequence quality scores.
- 11 It then saves them into a file the summary subfolder.
- Notes: The way it counts reads from FASTQ file is by reading the number of lines and dividing it
- by 4 (since each read in a FASTQ file consists of 4 lines: sequence identifier, sequence, plus sign,
- 14 and quality score).
- Notes: This does not include config file integration yet (TBA).
- Notes: Not integrated into any pipeline yet (TBA).

#### 17 2.8.1 Directory tree

```
/project_root/
  raw_reads/ ......(Prerequisite - Raw Data)
     sample1.fastq.gz
     sample2.fastq.gz
  trimmed_reads/ ......(Prerequisite - Trimmed Data)
     sample1_trimmed.fastq.gz
     sample2_trimmed.fastq.gz
  summary_statistics/
     fastqc_raw/
       \_\mathtt{sample1\_fastqc.zip}
       sample2_fastqc.zip
     fastqc_trimmed/
       sample1_trimmed_fastqc.zip
       sample2_trimmed_fastqc.zip
     extracted_metrics/
       sample1_base_quality.txt
       sample1_sequence_quality.txt
     raw_read_counts.txt
    _trimmed_read_counts.txt
```

### 2.9 plot\_and\_detect\_intermediate\_coverage.py

- 2 Stage: Done
- 3 Processing of coverage data
- $^4$  Specifics: This script takes the coverage data from a coverage file, which is expected to have 3
- 5 columns: chrom(chromosome/contig name), pos(position along the contig), cov(coverage value
- at that specific position). It then takes the latter two columns and uses matplotlib to create a
- 7 line plot of the coverage values and saves it as an image.
- 8 Notes: Presently, it determines, intermediate coverage by looking first looking for the minimum
- 9 and maximum value of coverage data.

10

#### Sept 20 2024 Update

I updated this one here to be a bit more robust in detection of chimeric contigs. Instead of visual inspection (which is a bit too subjective for my taste), I added 4 new features to this

- 1. Sliding window approach to detect changes coverage (rather than visualization alone) other than that we apply sliding window to detect changes in:
  - (a) Codon usage bias
  - (b) GC content
  - (c) Tetranucleotide frequencies
- 2. Each of these "windows" are then subjected to ANOVA, to determine statistical significance and
- 3. PCA, that combines all the (3) features to generate a 2D plot which to see whether different sections of the contig cluster differently which would indicate likely chimerism.

11

#### <sub>1</sub> 2.10 Shortbred ARG.bash

- <sup>2</sup> Stage: Back to drawing board and to test
- 3 Specifics: Marks trimmed reads with ARGs Technical Details This script takes cleaned reads
- 4 (dicated by the user), unzips them (to FASTQ), then starts finding ARG-markers on them.
- 5 Markers are determined by the function shortbred\_identify.py (instrinsic to the tool), taking
- 6 first the CARD database and aligning it with the RefSeq. After that, it clusters the RefSeq
- database at 95% similarity with those on the CARD database, to create markers. After that, it
- 8 uses another function shortbred\_quantify.py
- 9 Notes: I included an additional step here that filters out specific keywords from the FASTQ files
- 10 from a reference presently. It called filter\_keywords.txt which you can make yourself should
- 11 you want to filter out specific ARGs that you consider "low-confidence", or just create the file
- and leave it blank (up to you).

#### Sept 20 2024 Update

I am revisiting this process because:

- It turns out that ShortBRED is typically used on contigs, not raw reads. This makes sense because marker-based analyses are more accurate on contigs, as they reduce false positives and provide a clearer view of the genetic background.

  Note: ShortBRED is used to quantify genes, and raw reads are fragmented sequences.
  - Note: ShortBRED is used to quantify genes, and raw reads are fragmented sequences it is the CONTIGS THAT CONTAIN GENES.
- The current method of filtering low-confidence ARGs is inefficient: it creates a temporary file full of ARGs with the keywords, and then filters the marker database using those. There must be a more efficient way to skip this unnecessary step of generating temporary FASTQ files.

I have also streamlined the script to:

- Unzip FASTQ files in parallel.
- Remove the temporary file afterwards.
- Focus only on high-confidence ARGs. Previously, it also produced markers on the RefSeq database, but checking marker percentages turned out to be inefficient and pointless.
- Add checks when directory creation or unzipping fails.
- Use a consistent naming scheme.
- and finally, it now includes MEGAHIT assembly and a filtering step that removes contigs below 200 bp before ShortBRED processing.

The most important addition is the conversion from FASTQ to FASTA using seqtk. I had overlooked this step, as some tools cannot process FASTQ format.

Personal Note: Dear Reader, please be aware that ShortBRED is quite particular about FASTA headers. It requires the headers to be numeric identifiers, which adds an extra step of renaming all sequences to numbers. You'll also need to create an index to trace the original sequence headers back to their numeric counterparts.

While troubleshooting, I encountered persistent naming errors that were confusing at first. After diving into the source code, I eventually discovered that ShortBRED requires headers to be strictly numeric. This unexpected requirement added to the complexity of the process and was not immediately obvious.

### <sup>1</sup> 2.11 refseq\_bacteria.bash

- <sup>2</sup> Stage: Done
- 3 FastQC Automation script to run download from the bacteria RefSeq
- 4 Specifics: This only downloads the ones that end in .faa as opposed to .fna the latter are
- 5 genomes, the former are just all the annotated proteins themselves, with a genome or not.
- 6 Notes: This script is kept here because it was difficult to find the link to the FTP site, because
- 7 NCBI seemed to have had a recent restructuring.

#### Sept 20 2024 Update

Removed the hard-cap of 1-511.faa files, to add flexibility; now downloads without the need for the user to check how many to download. Also now uses aria2c, which has and intrinsic redownload unfinished files instead of wget function.

1 2.12 trimmomatic.bash; pipeline module

- 2 Stage: Done
- 3 Trimmomatic automation script to run FastQC on reads
- 4 Specifics: This script takes input reads (dictated by the pipeline) and trims them.
- 5 Note: This also contains the Trimmomatic parameters that I decided to use as placeholder at
- 6 the beginning of this whole project.

Note: Will likely become deprecated soon (or simply altered) depending on the results of my optimization tests on trimmers (TBA). Code snippet is below.

- 9 TRIMMOMATIC\_ADAPTERS="NexteraPE-PE.fa"
- TRIMMOMATIC\_SETTINGS="LEADING:10 TRAILING:10 SLIDINGWINDOW:4:20 MINLEN:60"
- 11
- 12 .
- 13 trimmomatic PE -phred33

#### <sup>1</sup> 2.13 renamingSIMS.bash

- <sup>2</sup> Stage: Done
- 3 renaming output files from raw-reads simulators i.e. CAMISIM
- 4 Specifics: This script takes the output folder of CAMISIM called the out directory and automates
- 5 renaming them, and moving them.
- 6 Note: Directory structure is below to show you why automation is necessary and manually
- 7 renaming is too time-consuming.

#### 9 2.13.1 Directory Tree with Notations

```
CAMISIM/
 out/
   sample1/ ...........Renamed from "2024.09.05 11.25.28 sample 0" to "sample1"
    _contigs_1/ ......Renamed from "contigs"
    reads_1/ ...... Renamed from "reads"
      anonymous_reads.fq.gz ......Original interleaved FASTQ file
      sample2/ ...........Renamed from "2024.09.05_11.25.28_sample_1" to "sample2"
    _contigs_2/ ......Renamed from "contigs"
    anonymous_reads.fq.gz ......Original interleaved FASTQ file
      raw_reads/
   sample1_R1.fastq.gz Copied from "CAMISIM/out/sample1/reads_1/R1.fastq.gz"
   sample1_R2.fastq.gz Copied from "CAMISIM/out/sample1/reads 1/R2.fastq.gz"
   sample2_R1.fastq.gz Copied from "CAMISIM/out/sample2/reads_2/R1.fastq.gz"
   sample2_R2.fastq.gz Copied from "CAMISIM/out/sample2/reads_2/R2.fastq.gz"
```

### 1 2.14 prokka ARG.bash

- <sup>2</sup> Stage: Done
- 3 Uses the wrapper tool prokka to predict genes from contigs
- 4 Specifics: This script first concatenates the CARD and NCBI-AMR databases then using the
- 5 --protein option, which replaces its database with the one the user specifies, uses that concate-
- 6 nated database to look for possible ARGs in contigs.
- 7 Note:

### Sept 21 2024 Update

I removed it's companion script prokka\_Uniprot.bash from the repo because it is redundant. Prokka already uses UniProt. I also updated it to include an EMBOSS transeq function to translate the fasta files to protein sequences, something that I overlooked and thought that prokka was able to translate by itself.

#### 2.14.1 Directory tree

```
project_root/
    databases/
```

```
CARD_sequences/
    extracted/
     \_ protein_fasta_protein_homolog_model.fasta CARD nucleotide sequences
      (Prerequisite)
  NCBI_AMR_sequences/
    AMRProt.fasta ......NCBI AMR nucleotide sequences (Prerequisite)
  arg_proteins.fasta Translated protein sequences (created by transeq) (Generated)
merged_contigs/
  SSRN_filtered_contigs.fa ............Contig files for other samples (Prerequisite)
prokka_output/
 _prokka_ARGs/
    SSR1/
     _ARG_SSR1.faa ....... Prokka protein FASTA output for SSR1 (Generated)
      ARG_SSR1.gbk ......Prokka GenBank file for SSR1 (Generated)
    SSR2/
      ARG_SSR2.faa ........ Prokka protein FASTA output for SSR2 (Generated)
      ARG_SSR2.gff ......Prokka GFF annotation for SSR2 (Generated)
     all_ARG_nucleotides.fasta .. Concatenated nucleotide sequences (created by script)
(Generated)
your_script.sh ......Bash script for the workflow (Script)
```

#### <sub>1</sub> 2.15 RefSeq.bash

- 2 Stage: Done
- 3 Downloads Bacterial RefSeq database
- Specifics: It uses aria2 a multi-connection download tool with -x 16 s -16 parameters meaning
- 5 16 connections. It then
- Verifies the downloads (confirming if the downloads are successful)
- Redownloads the unsuccessful downloads
- Extracts each individual file
- Concatenate or combine them into a single file called refseq bacteria.fasta
- Note: Used to be called nr\_download.bash, renamed it at Sep 22, 2024.

### <sup>1</sup> 2.16 Pavian\_analysis.R

- 2 Stage: Done
- <sup>3</sup> Visualization of Kraken2 via Pavian

- 4 Specifics: Uses Pavian to visualize specifically the \_kraken2\_summary.txt files. Additionally,
- it calls on ggplot2 to create a bar plot taxonomic classification counts (at different taxonomic
- 6 levels). It then redirects all output (tables and plots) into a pavian\_output.pdf.
- Note: You have to set the change the path to the correct Kraken2 output directory where the
- 8 .kraken files are.

### <sub>1</sub> 2.17 parseFastQC.py

- 2 Stage: Done
- 3 Interprets all the FastQC summaries
- 4 Specifics: It does a few things in sequence:
- Uses the summary.txt files from FastQC.
- 2. Creates a Legend that lists the abbreviations used in the file and puts them in a separate legend\_tsv file.
- 3. Extracts each FASTQC ZIP file into a temporary directory and looks for the summary.txt
- 4. Generates a summary report called output\_tsv for each sample (per direction R1 and R2).
- 12 Note: Make sure to set the correct path to the FastQC output directory where the .zip files are.

#### 13 2.17.1 Directory tree

```
project_root/

___fastqc_output/
___sample1_R1_fastqc.zip ......FastQC zip file for Sample 1 (Read 1) (Prerequisite)
___sample1_R2_fastqc.zip .....FastQC zip file for Sample 1 (Read 2) (Prerequisite)
___sample2_R1_fastqc.zip .....FastQC zip file for Sample 2 (Read 1) (Prerequisite)
___sample2_R2_fastqc.zip .....FastQC zip file for Sample 2 (Read 2) (Prerequisite)
___output/
___fastqc_report.tsv ......Generated FastQC summary report (Generated)
__legend.tsv ......Abbreviation legend for FastQC metrics (Generated)
__scripts/
___check_fastqc.py .......Python script for generating FastQC report (Script)
```

### <sub>1</sub> 2.18 minimum\_length\_CARD.py

- 2 Stage: Done
- 3 Description: Parses the CARD protein homologue FASTA file and identifies the shortest sequence
- 4 length.
- 5 Specifics: This script uses the Biopython library to read each sequence in the FASTA file,
- 6 comparing lengths and returning the smallest sequence length in the dataset.

- 7 Note: The input FASTA file should contain the CARD protein homologue sequences—so the
- 8 user has to set it to where it is.

### 1 2.19 template modulars: megahit\_binning.sh

- <sup>2</sup> Stage: Done
- Description: Template modular script that can be integrated into Snakefiles.
- 4 Specifics: This script loops through deduplicated paired reads (\*\_R1\_uniq.fastq) and runs
- <sup>5</sup> MEGAHIT, then filters out contigs > 200 bp. It is written specifically for integration into
- 6 Snakefiles, so its input and output directories are dictated by the Snakefile shell.
- 7 Note: MEGAHIT can read FASTQ files—I double-checked, haha.

The modified companion script, megahit\_binning.sh, is specifically designed for the Binning.smk pipeline, but they both perform the same task.

```
MEGAHIT_SETTINGS="--k-min 35 --k-max 141 --k-step 28"
```

Note: FastQC.bash is another template modular script that needs to be integrated into Snakefiles, but it instead uses FastQC.

### <sub>1</sub> 2.20 diversity bootstrap.R

2 Stage: Draft

10

- 3 Description: Reads and plots diversity indices.
- 4 Specifics: Parses diversity files for individual metrics and generates the following plots:
- Alpha-diversity plotting:
- Ridgeline plot: Shows the distribution of diversity metrics across samples, per bootstrap replicate.
- Violin plots: Displays diversity index distributions with individual points plotted for each bootstrap.
  - Beta-diversity plotting:
    - Heatmap: Visualizes the consistency of beta-diversity across bootstraps.
- NMDS (Non-Metric Multidimensional Scaling): Plots ordination based on Bray-Curtis
   distances.
- Note: This script needs updates to include: (TBA)
- Specific paths to diversity files.
- More customization options for the heatmap, such as clustering methods or color scales.

• Optional: Optimization for memory usage, as large datasets with many bootstraps can consume significant resources.

### 1 2.21 calculate\_plasmid\_percentage.py

2 Stage: Done

17

18

- 3 Description: This script calculates the percentage of reads that map to plasmids based on the
- 4 total and plasmid read counts.
- 5 Specifics: The script performs the following steps:
- 1. Reads the plasmid and total read counts from input files provided by Snakemake.
- 2. Calculates the percentage of plasmid reads relative to the total number of reads using the formula:

$$plasmid\_percentage = \left(\frac{plasmid\_reads}{total\_reads}\right) \times 100$$

- 3. Writes the percentage of plasmid reads (formatted to two decimal places) to the specified output file.
- Note: The input and output file paths are provided by Snakemake, ensuring integration into a larger pipeline.

### $_{\scriptscriptstyle 1}$ 2.22 calculate\_diversity.py

<sup>2</sup> Stage: Draft

13

- 3 Description: This script calculates both alpha and beta diversity metrics.
- 4 Specifics: The script reads species abundance data from .bracken files, calculates various
- 5 diversity metrics, and outputs the alpha and beta diversity results, both to the console and to a
- 6 file called diversity\_matrices.tsv.
- 7 Alpha Diversity Metrics: 13 Fisher's Alpha
- Using scikit-bio 14 Chao1
- Shannon Shannon
- Beta Diversity Metrics (via scikit-bio):
- Pielou's Evenness 17 Bray-Curtis
  - Custom implementations 18 Jaccard
- Note: There are two other beta diversity metrics not included here because they require phylogenetic trees (in Newick file format) beforehand, i.e., UniFrac and unweighted UniFrac (TBA). The Newick file format looks like so:
- 22 (A:0.1,B:0.2,(C:0.3,D:0.4):0.5);

 $^{23}$  where A, B, C, and D correspond to different taxa, the numbers represent branch lengths

24 (distances), and the bracketing specifies the clades.

# <sup>1</sup> Chapter 3

# <sup>2</sup> Project Side Scripts

### ₃ 3.1 kmer contam.smk

- <sup>4</sup> Stage: To test General Purpose
- 5 This pipeline is designed for the detection and removal of contaminant sequences using
- 6 k-mer-based filtering. It includes steps for contaminant k-mer generation, mapping k-mers to
- 7 metagenomic reads, and statistical testing for ambiguous sequences. The pipeline incorporates
- 8 tools for read quality control, contamination filtering, and re-validation using k-mers and BUSCO
- 9 analysis.

10

11 Preprocessing

### 3.1.1 Contaminant Databases Download

- 14 Technical Notes: The contaminant databases UniVec from NCBI, PhiX, and the 1000 Genomes
- 15 Project are downloaded using aria2c to ensure a fast and reliable download process.
- Rationale: These databases contain many known contaminants that may appear in metagenomic
- samples, which must be filtered out before further downstream analyses.

#### Personal Notes

Usually, in standard bioinformatics analyses:

- 1. The largely annotated human genome, GRCh38 (often referred to as HG38), is an upgraded version of the previous build, GRCh37 (sometimes called HG19), and is commonly used to filter out contaminants.
- 2. Tools like minimap2 or BBDuk are typically used to map reads to the human genome (or other contaminant references) and then remove contaminant sequences via k-mer matching or alignment.

So why did I create this more elaborate workflow?

- Tools like BBDuk and other k-mer-based mappers, such as Kraken2, often rely on exact matches to assign taxonomic profiles of contaminants or other sequences.
- While GRCh37 (or GRCh38) is a robust and well-annotated reference genome, it
  doesn't fully capture the breadth of human genetic diversity. It represents an
  amalgamation of human populations, but strict k-mer matching means it only
  identifies specific k-mers from that reference genome. The 1000 Genomes Project
  covers a broader range of human genetic diversity and is therefore included in this
  workflow to better account for this variability.

#### 19 3.1.2 BUSCO Validation

- 20 Technical Notes: BUSCO is used to validate the presence of Single Copy Genes (SCGs) across the
- 21 metagenomic data. This step ensures the retention of high-quality sequences that are biologically
- 22 meaningful.

18

- 23 Rationale: SCG validation confirms the biological integrity of the metagenomic sample after
- 24 contamination filtering. It uses all available BUSCO lineages for a comprehensive assessment.

#### 25 3.1.3 Marker K-mer Generation

- Technical Notes: K-mer counting is performed using Jellyfish for both SCGs and contaminants.
- 27 This process generates k-mers that represent sequences of interest (SCGs) and contaminants
- <sup>28</sup> (UniVec, PhiX, 1000 Genomes).
- 29 Rationale: K-mers allow rapid matching between known contaminants and sample reads, en-
- 30 abling efficient filtering and high-fidelity read retention.
  - Contamination Filtering and Statistical Testing

#### 34 3.1.4 Mapping K-mers

- 35 Technical Notes: KMA is used to map k-mers from SCGs and contaminants to raw metagenomic
- 36 reads.

31

- 37 Rationale: Mapping k-mers identifies sequences that match known contaminants, allowing for
- $^{38}$  removal of ambiguous or low-quality sequences from the data.

#### Personal Notes

The use of KMA here specifically, instead of other mappers like Bowtie2 and BWA, is because:

- While Bowtie2, BWA, and KMA are not restricted to exact matching (i.e., they can tolerate mismatches, increasing sensitivity),
- KMA performs better with highly redundant databases. Although Bowtie2 and BWA were originally optimized for mapping reads to human genomes or other low-redundancy datasets, they are not restricted to such uses. Both are excellent mappers, but KMA is specifically optimized for working with highly redundant databases, such as those commonly encountered in metagenomic or microbial datasets. For more information, see the RGI documentation.

40 3.1.5 Normalization and Testing

- 41 Technical Notes: Contaminant and SCG k-mers are normalized using wavelet transformation,
- and a statistical test (e.g., t-test or Z-test) is performed to detect ambiguous regions.
- Rationale: Normalization ensures that the k-mer counts are comparable across samples. Statisti-
- cal testing allows detection of ambiguous areas that may require further analysis or removal.
- Notes (For more details on wavelets, see Wavelets)

#### Personal Notes

The usage of wavelets here increases sensitivity and provides statistical robustness. Since we are no longer relying on exact matching but allowing for mismatches, we expect a distribution of k-mer bindings embedded within the coverage data. To analyze this distribution, we apply a wavelet transformation (similar to a Fourier transformation, but localized) to decompose the data into smaller components (frequencies) that contribute to the overall pattern.

Wavelet-based statistical testing is implemented in this step through a series of Python scripts:

- normalize\_scgs\_wavelet.py and normalize\_contam\_wavelet.py load the k-mer counts from SCGs (identified by BUSCO odb10) and from contaminant databases, respectively. These scripts apply continuous wavelet transformations (CWT) to create wavelet distributions and then calculate Z-scores for the wavelet coefficients to assess their statistical significance. Wavelet coefficients with Z-scores corresponding to a p-value of less than 0.05 (Z-score ≥ 1.96) are retained, while the others are filtered out.
- compare\_wavelets\_stats.py compares the wavelet distributions of SCGs and contaminants, testing whether they are statistically different (with the null hypothesis, H0, being that they are not different). If the null hypothesis is accepted (i.e., the

distributions are similar), the sequences are flagged as ambiguous and filtered out.

```
significant = np.where(p_value < 0.05, 'significant', 'ambiguous')</pre>
```

Update Sep 27, 5:33 am compare\_wavelets\_stats.py has also been now updated to handle:

- Decision making of using parametric or non-parametric wavelet tests
- Handle a battery of different tests
- Apply Bonferroni and FDR corrections for such tests

#### 48 3.1.6 Filtering Ambiguous Sequences

- <sup>49</sup> Technical Notes: Sequences identified as ambiguous are filtered out to retain only high-fidelity
- 50 reads in the final metagenomic assembly.
- 51 Rationale: This step ensures that the final assembly is free of contamination and contains only
- bigh-quality sequences for downstream analyses.

#### 53 3.1.7 Re-validation of SCGs

- Technical Notes: BUSCO is rerun on the final cleaned assembly to confirm that SCGs have been
- 55 retained after contamination filtering.
- Rationale: Re-validation with BUSCO ensures that the final dataset remains biologically
- meaningful after the filtering process.

#### 58 3.1.8 Directory tree

busco_outputs/	
dataset1/	
dataset2/	
busco_validation_outputs/	
short_summary.txt	
scripts/	(Prerequisite)
normalize_contam_wavelet.py	(Prerequisite)
normalize_scgs_wavelet.py	(Prerequisite)
compare_wavelets_stats.py	(Prerequisite)
env/	
busco_env.yaml	(Prerequisite)
jellyfish_env.yaml	(Prerequisite
raw_reads.fastq	` - ,

### Update Sep 27 3:57 am

Provided the limitations of the original Snakefile, which assumes normally distributed k-mer binding, I created another workflow that tests "normality" of the distribution. This determines whether a Z-test or t-test is valid. Here's a brief explanation of the scripts:

- normality\_test.py uses the Shapiro-Wilk and Kolmogorov-Smirnov tests to determine normality. If normality fails, we proceed with:
- fit\_distributions.py to check whether the data follow log-normal, exponential, or gamma distributions, which can be transformed using:
- transform\_data.py, which attempts to normalize the data. If normalization succeeds, we proceed with parametric tests.
- If normalization fails, we use check\_nonparametric.py to determine the type of non-parametric distribution the data likely follows.
- To increase confidence in the distribution type, I included two scripts that test goodness of fit:
  - goodness\_of\_fit\_parametric.py, and
  - goodness\_of\_fit\_nonparametric.py.

Using statistical tests rather than visual inspection removes human judgement bias and focuses purely on mathematical/statistical validation.

In line with this, researchers often settle on subjective p-values (e.g., < 0.05 or < 0.01) as thresholds, known as alpha. I included scripts that determine the best alpha for whichever distribution has the best goodness of fit:

- best\_alpha\_parametric.py and best\_alpha\_nonparametric.py, either of which feed into:
- bestfit\_and\_alpha.py, which determines the best alpha for downstream analyses.

- 60 Chapter 4
- <sup>61</sup> Project Main Scripts
- 1 4.1

Further investigations

- This section comprises of concepts or ideas that require further investigation. They are
- 5 written in boxes to help categorize them cleanly. I will refer to this section often as no script is
- perfect and can be further improved.

#### 7 1 Information

#### Biological Information

I have long pondered upon the idea of creating a graph with biological information on the y-axis and read length on the x-axis. I hypothesize that we will find a "sweet spot" wherein we can optimize the amount of information/read using such trimmers. Unfortunately, it is extremely difficult to define what a biological sequence really is, because technically you can generate any random sequence - whether protein or nucleotide. That's the main reason I have been stuck on this problem for quite a while, I've been trying to find the answer first.

The core difficulty here lies in defining what constitutes biological information in a meaningful, quantifiable way. Since any random sequence of nucleotides or proteins could be technically "valid" (false-positives).

When people are asked this question they often give DESCRIPTIONS of what a biological sequence is but not what DEFINES a biological sequence. I understand that there are many characteristics that can help in determining the signal from the noise, but I am just not satisfied with whether this "thing" checks most (if not all) the boxes - I need a non-subjective answer to this problem.

#### 9 2 Wavelets

#### Wavelets

What is a Wavelet?

A wavelet is a small, localized wave that is used to represent signals at different scales. Unlike sinusoids in the Fourier transform, which extend infinitely, wavelets are confined to a limited duration. This makes them highly useful for capturing both frequency and time (or space) information simultaneously.

Wavelets are particularly well-suited for analyzing signals with transient or localized features, such as sharp peaks, edges, or changes in behavior. Because of their ability to zoom in on fine details while also capturing broad trends, wavelets are widely used in signal processing, image compression, and data analysis.

How Do Wavelets Work?

Wavelets work by breaking down a signal into smaller, simpler components—each representing the signal at different scales. The signal is convolved with a family of wavelets, each scaled and shifted to analyze the data at various resolutions. The result is a multi-scale representation that provides insights into both high-frequency details (like sudden spikes)

and low-frequency trends (like global patterns).

Continuous Wavelet Transform (CWT)

The Continuous Wavelet Transform (CWT) is a specific type of wavelet transform that decomposes a signal continuously over a range of scales. CWT produces a 2D representation of the signal in both time and frequency domains, making it ideal for detecting localized features that vary across scales.

Why CWT for k-mer Normalization?

In bioinformatics, CWT is employed to normalize k-mer counts by analyzing the data across multiple scales. This multi-scale analysis captures localized features in the k-mer distributions, such as regions affected by contaminants or biologically significant regions (e.g., single-copy genes, SCGs). By applying CWT, we can detect and highlight these variations while preserving the overall structure of the data.

Signal vs. Noise Differentiation with CWT

One of the key strengths of wavelet transforms is their ability to differentiate signal from noise. By breaking down a signal across multiple scales, wavelets can identify high-frequency components often associated with noise, while preserving lower-frequency, biologically meaningful signals. This is especially important in this workflow, where tools that allow k-mer mismatches are used. Unlike tools like minimap2 or BBDuk, which prioritize specificity (via exact kmer matching), the wavelet-based approach prioritizes sensitivity, helping to detect real signals despite mismatches or noise in the data.

Key Advantages of CWT:

- Multi-scale analysis: CWT allows us to view the data at different scales, capturing both small localized features and broader trends.
- Localized feature detection: Unlike global methods (e.g., Fourier transform), CWT can detect localized anomalies in the data, such as contamination spikes that only affect certain regions.
- Non-stationary signal analysis: Many biological signals, including k-mer counts, are non-stationary (their statistical properties vary over time or space). CWT is well-suited for handling such data.
- Noise filtering: CWT can differentiate high-frequency noise from low-frequency signal, making it an excellent tool for extracting meaningful data even in noisy datasets.

#### Application in This Workflow:

In this workflow, CWT is applied to normalize k-mer counts for both contaminants and SCGs. After normalization, statistical tests like t-tests or Z-tests are used to detect ambiguous regions that may require further analysis or removal. This process ensures that k-mer counts are comparable across samples and that biologically meaningful patterns are preserved.

### <sup>12</sup> 3 ModelTesting

#### Model Testing in Phylogenetics

What is a Model Testing in Phylogenetics?

Note that especially in exploration of the tree space using Maximum Likelihood or Bayesian Inferences, you are often first required to test for the model of evolution - this serves two related purposes:

- To find the most fitting model given the dataset
- That is also the simplest one (avoids overfitting and thereby computational resources)

So what is a model in phylogenetics? A model is simply a possible explanation of how a specific gene or partition (discussed later) likely evolved.

Note Different models take into account different evolutionary processes (and parameters) into account (some account for more than others). Factors include:

- Transition vs Transversion rates
- Base frequencies
- Among-site rate variation
- The behavior or rate distribution Note Assuming differences in site variation is often known as  $\Gamma$  while I means proportion of sites is invariant.

To determine which is best tools such as ModelTest or jModeltest (if you prefer GUIs) use certain statistical criteria i.e.

- Akaike Information Criterion (AIC),
- Bayesian Information Criterion (BIC), or
- likelihood-ratio tests (LRT),

to rank models based on how well they explain the data while penalizing for model complexity.

Selection of a model affects phylogenetic trees in terms of

- Branch lengths
- Overall tree topology

Why bother penalize overfitting? First and foremost, overfitting may cause the model to assume random noise in the data as genuine evolutionary signals. Overfitting may also cause it to become too specific at explaining the data at the cost of generalizability.

Finally, it leads to unnecessary computational resources and time.

Personal Note TLDR, to balance between accuracy vs speed, and data specificity-generalized conclusions.

What to do when using concatenated sequences? When using concatenated sequences - likely from phylogenomics or using multiple marker genes. It is important to note that (more likely than not) evolutionary rates differ between different genomic regions. Case in point: coding vs non-coding regions or the mere fact that concept of conserved regions exist. In such cases as partitioned analysis is recommended - where you apply different models for each stretch of DNA or partition.

Note Doing so also helps you avoid Simpson's Paradox - a statistical bias - where trends from several groups disappear or reverse depending on how you partition your data based on metadata. Personal Note Other methods to avoid this bias include:

- Using Bayesian methods
- Coalescent models see Box on Coalescent Phylogenetics
- Phylogenetic Networks Box on Network Phylogenetics

### <sup>15</sup> 4 Phylogenetics

#### Coalescent

Networks in Phylogenetics

### s 5 Signal vs Noise

#### Signal Averaging

One can then check for variabilities in the averaged data and perform statistical analysis as to whether they cluster neatly.

#### 20 6 Robustness

#### Benchmarking

Why test on your own dataset?

Specific conditions, organisms, or complexities applicable to your metagenomic data
 which Gold standards might not be able to capture e.g. noise, biases, biological variation

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- Gold standards often reflect idealized situations, disallowing you to fine-tune or feature engineer based on your specific data.
  - Personal Note Every metagenomic dataset has unique challenges and represent a sample that no longer exists in the real world (because you took it). Testing the data allows you to specifically optimize tools or methods for those issues rather than for general use cases.
- Tools trained on gold standards may perform differently when exposed to new (real world) data, especially when your dataset has (unforeseen )biases or noise not accounted for in the development of the gold standard.
- Lastly, your research goals may differ from those who developed the simulations or gold standards. Testing directly on your dataset makes sure that any optimizations are specifically meaningful to your data.

Information dump

This is part is mainly created as an information dump outside of just bioinformatics, that perhaps we can one day apply to bioinformatic data analysis (mostly). This part could also serve as an index of papers that I've read regarding various topics that seem interesting to me and that I think can be applied in the wider field of computational or mathematical biology.

### <sup>29</sup> 2 Biological OFF Decay

In statistical mechanics, microstates (positions, velocities, energies, etc.) are often unpredictable—this is related to Schrödinger's thought experiment and further elaborated by Werner Heisenberg's Uncertainty Principle. However, macrostates (large-scale properties) that emerge from the ensemble of microstates are model-able. For \*\*radioactive decay\*\*, while we can't predict when individual atoms will decay, we can model the \*\*half-life\*\* of the entire ensemble. The \*\*exponential decay equation\*\* models this process:

$$N(t) = N_0 e^{-kt}$$

- Where:
- N(t) is the amount of substance remaining after time t,
- $N_0$  is the initial amount of the substance (at t = 0),
- e is the base of the natural logarithm ( $e \approx 2.718$ ),
- k is the decay constant, determining the rate of decay,
- t is the time elapsed.
- This equation describes a process where the quantity decreases over time at a rate proportional to its current value, leading to exponential decay. The negative exponent indicates that the amount is decreasing over time.
- How does this connect to gene expression?
- Individual cells in a tissue either have certain genes ON or turned OFF—this binary on/off behavior is analogous to the decay of atoms (binary: decayed or not). Thus, we can use the same exponential decay model to predict the number of genes that turn OFF in a population of cells (or tissue or organism) over time.
- In this model, we simply change the variables to
- N(t) is the number of cells with genes still ON at time t,
- $N_0$  is the initial number of ON genes (at t = 0),
- k is the decay constant, determining the rate at which genes are turned OFF (analogous to radioactive decay),
- t represents biological time (e.g., the time it takes for environmental conditions to cause changes in gene expression).

How can we determine the decay constant k?

In gene expression studies, k could represent the rate at which environmental or physiological conditions (such as cold temperatures or seasonal changes) cause genes to turn off. This decay constant could be derived from empirical data (e.g., using transcriptomics data like DeSeq2) by calculating how quickly gene repression occurs over a certain period of time.

The model can help us estimate when a certain percentage of genes will be repressed, akin to the half-life concept in radioactive decay. This framework allows us to mathematically predict e.g. when a plant or organism is preparing for overwintering, or to determine the rate of gene repression under specific environmental conditions.

### 66 3 Best Data science practices

- Here as some of the best practices and principles in data science which can be adopted to any scientific discipline dealing with large amount of data.
- 1. Defining the problem clearly which includes
- Objective/s
- Scope of the Project
- Key metrics for success
- 2. Knowing data data which including
- Data structure
- Data type

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- Data distribution
- Anomalies e.g. missing values
- Imbalances in data e.g. oversampling, undersampling, synthetic data generation etc.
- 3. Ensure reproducibility
  - Documentation: cleaning, analysis, modeling
  - Scripts and Notebooks
    - Makes your code easier to understand and facilitates collabs and updates.
- <sup>83</sup> 4. Validation
  - Split data into training and testing sets to ensure models generalize well to unseen data.
    - Prototype first before implementation
    - Feature Engineering
    - Optimize trade-off between complexity and interpret-ability
  - Implement Data Augmentation techniques to increase diversity and quantity of training data, improving robustness
  - 5. Consider ethical practices and biases; ensure fairness on treatment of data
  - Data privacy and Security is top priority
    - Encryption
- Access controls
- Anonymyzation techniques
- 6. Automate where possible to reduce the likelihood of human error this includes:
  - Pipelines

- Feature engineering updates 98 Feedback gq 7. Stay updated with the latest tools 100 8. Scalability: ensure models 101 • Can handle increasing amounts of data (data storage scalability) without 102 Significant performance degradation 103 Significant increases in computational resources 104 9. Impact: avoid "Analysis Paralysis" by focusing on insights that are actionable 105 10. Implement feedback loops based on 106 New data 107 Performance metrics 108 Stakeholder and end-user feedback 109 Alignment with broader objectives (be relevant and actionable) 110 Explore other models • Data leakage checks 112 Note that bias can arise from data 113 • Appropriate metrics are used 114 Always be skeptical of results 115 Feedback from domain expertise Auditing 117 11. Mind the end users; models must be 118 Accessible Interpretable 120 Usable 121 • Continuous Integration and Deployment (CI/CD) 122 12. Communication: 123 Uncertainties 124 Assumptions 125 Model explainability techniques: how complex models make decisions 126 13. Understand the data lifecycle 127
- (a) Collection to preprocessing
  (b) Analysis
  (c) Modeling
  (d) Eventual archival and deletion

### <sup>132</sup> 4 Staying updated

In line with best practices, here is a list of where to find the latest tools and the journals I personally consider to be of high-impact.

- 135 4.1 Where to find some protocols
- Nature Protocols, Nature Methods
- Journal of Visualized Experiments
- Current Protocols
- Cold Spring Harbor Protocols
- Protocol Exchange ......open repo hosted by Nature
- Bio-Protocol
- PLOS ONE Protocols
- STAR Protocols ...... more like a part of Cell
- 4.2 Bioinformatics Tools and Journals
- BMC Bioinformatics
- Briefings in Bioinformatics
- Nucleic Acids Research (NAR)
- Journal of Computational Biology
- PLoS Computational Biology
- Bioinformatics and Biology Insights ...... Bit on the low side of IF, but Open Access
- GigaScience ....... Also a DB repository for some custom datasets
- Genome Biology
- Scientific Data ...... hosted by Nature, aptly named
- Bioinformatics Advances

157	4.3 High quality (imo) Life-Science Journals
158	• General
159	- Nature
160	- Science
161	- Cell
162	– Annual Reviews
163	- Trends
164	– EMBO Mostly molecular biology
165	- PLoS (especially the specialized ones)
166	– Current Biology
167	• Mostly medical literature
168	- The Lancet
169	- NEJM
170	$-\mathrm{BMJ}$
171	- JAMA
172	– Cochrane Best place for Evidence-based Literature
173	4.4 Notable Labs and Groups for AMR research
174	– Knights Lab
175	– Dantas Lab Washington University in St. Louis
176	– Beiko Lab
177	- Fraser Lab
178	– Moran Lab
179	– Bhatt Lab Stanford University
180	– ph4ge group
181	– StaPH-B group
182	- phytools blog

## $_{183}$ 5 Beyond the scope of this study

 $^{184}$  Below are some of the analyses that are overlooked or not typically addressed - but not by high  $^{185}$  IF papers regarding antimicrobial resistance.

#### 186 5.1 In General

- Functional characterization of ARGs
- Experimental validation
  - Mechanisms of resistance
- Linking ARGs to specific pathways or cellular processes
- Ecological and Evolutionary Context
- How ARGs emerge, persist, and spread
- HGT, selective pressures, impact of environmental factors
- Innovative methodologies

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- Focus on mobile genetic elements
  - Plasmids, Transposons, Integrons
- Co-localization and dynamics of transfer
- Evolutionary dynamics of the resistome
- Public Health and Clinical Relevance Linking
- Clinical outcomes
- 201 Potential therapeutic strategies
- Policy recommendations
- Burden of ARGs in the environment
- Strategies for mitigating spread
- Quantitative Analysis and Modeling
- Prediction of spread
- Assessment of risk factors
- 208 Evaluation of mitigation strategies
- Evolutionary trajectory
- Policy and Societal Implications
  - Evidence-based recommendations for antibiotic use, environmental management, and global health strat
  - Ethical considerations and regulatory challenges
    - \* Responsible usage of antibiotics
      - \* Implications to public health policies
- \* Need for global cooperation

217	5.2	Bioinformatics
218	•	Comprehensive data integration multi-omics
219	•	Advanced assembly and binning techniques
220		- Hybrid assemblies
221		- Magnetic isolation or targeted metagenomics
222		- Deep sequencing
223	•	Usage of multiple specialized databases such as
224		- CARD
225		- ResFinder
226		- ARG-ANNOT
227		- Deep-ARG
228		- then cross-validation between them
229	•	HGT analysis often by specialized tools e.g.
230		- ICEberg
231		- oriTfinder
232		– plasmidSPAdes
233		
234		– Co-occurrence analysis with MGEs using tools such as
235		* MOB-suite
236		* PlasFlow
237		* cBar
238		* Cytoscape
239		* SpiecEasi
240		- Resistance Gene Quantification
241		* RPKM or TPM normalization
242		* Accounting for copy number variations
243		* Differential expression
244		- Phylogenetic and Evolutionary Analysis
245		* Evolutionary origins, dissemination pathways
246		* WGS to build phylogenies rather than markers
247		* MLST integration
248		* Phylogeography and Spatial Phylogenetics $\ldots\ldots{\rm SPREAD}$ or PhyloGeoBEARS
249		$\ast$ Selection pressure of genes or genomes and how they relate to antibiotic usage
250		* Phylogenetic networks and reticulate evolution

251	* Co-phylogeny and Host-Microbe
252	* Gene Tree-Species Tree Reconciliation or discrepancies
253	* Ancestral State Reconstruction
254	* Phylogenetic Comparative Methods
255	* Bayesian Phylogenetics and Model Selection
256	st Simulation studies for phylogenetic validation under different conditions
257	· Different rates of evolution
258	· Incomplete lineage sorting
259	· Varying levels of HGT
260	* Phylogenetic placement of uncultured or unknown
261	*
262	- Comparative genomics
263	* Syntenic regions
264	* Conserved domains
265	* Shared genomic islands
266	* Core vs accessory genomes
267	* Core vs shell vs cloud
268	<ul> <li>Deep Learning or Machine Learning Approaches</li> </ul>
269	* Neural networks
270	* Random forests
271	* Support vector machines
272	<ul> <li>Novel Pipelines of Workflows</li> </ul>
273	• Binning validation
274	• Strain-level resolution
275	• Detailed Eco-evolutionary context reservoirs and how they impact human health
276	• Advanced Contamination Control low-biomass samples
277	• ID of novel ARGs followed by experimental validation
278	• Data sharing and reproducibility
279	• Benchmarking and Validation against established standards or reference databases

### When to eliminate duplicates

FastUniq and other tools filter out duplicate reads thereby sacrificing depth - in exchange for a decrease in false-positives during read mapping. It also decreases the amount of data that needs to be stored or processed for downstream. This is similar to how dRep clusters OTUs and how marker-based annotators e.g. MetaPhlan4 and ShortBred align via a "representative" sequence.

The risk you run, removing duplicate reads is an increase in false-negatives (true signals otherwise assumed to be sequencing artefacts). On the flipside, duplicate removal prevents skewing by technical replication errors - by forcing normalization of the data.

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### 82 5.3 When to Split Papers

- Splitting papers into multiple publications can be a strategic and necessary move when the scope is too broad. Here are some key considerations:
- Distinct research questions and hypotheses that can be fully explored on their own
- When methodologies are distinct enough to require separate justifications
- Significant results in different areas
- Coherence in narrative flow
- Major differences in teams and co-authors
- Data and results need independent discussions in themselves
- It is a follow up study

### <sup>293</sup> 6 More on Determining Biological Data

- One of the most distinguising aspect of biological entities is the ability or potential (in the case of viruses) to self replicate under the appropriate conditions which could be
- Under specific environmental or metabolic processes

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#### 298 6.1 How genes evolve

Historically, a gene is a stretch of DNA that encodes information regarding a protein.

300 Motifs

#### 301 6.2 Information

- 302 Shannon Entropy
- 303 GC content nr database testing or calibration
- 304 Conserved sequences and database bias

#### Hierarchical information

Sites and parititions

#### Non-coding "genes"

content...

#### Signal and noise

dN/dS signals epigenetic markers - chance of protein interactions

### Viruses and the vines of life

Infection of the protocell.

#### Origin of Life

Infection of the protocell.

RNA World Hypothesis - self replication

Self replication - not limited to RNA e.g. transposition (though it does involve transcription or reverse transcription)

#### More data or better tools?

With the advancement of faster and more efficient sequencing technologies we really have to question whether we need more data.

A parallel discussion about this was in the field of physics - specifically the Einstein's Determinism and Bohr's probabilistic view of quantum objects. Let me explain. The exact tension between these two great scientific thinkers came about because Einstein was not satisfied that the randomness of quantum systems (only being collapsed after observation) is inherent to universal laws - and that there may be hidden variables we have either failed to take into account or our tools are not precise enough to detect. In contrast, Bohr's view (the Copenhagen interpretation) suggests inherent indeterminacy in quantum systems.

Ingenious thought experiments were made (and directly observed) as a result of these debates in the Solvay conferences which included well-known physicists such as Born, Heisenberg, and Schrodinger.

On the same vein, biological sequences seem to have an apparent randomness to them that either have variables or models that are too complex we fail to take into account or that we simply require more data to determine these biological hidden variables.

As at its heart, similar to Schrodinger's thought experiment or radioactive decay see section on Biological OFF Decay, where exactly mutations occur on a stretch of DNA is inherently random though we can model the rate at which it occurs (about 10e-10) depending on the organism.

Yes, be skeptical about noise, but also accept that some uncertainty is inherent. It's like a random walk, but constrained ultimately by developmental and evolutionary forces some of which are random e.g. genetic drift, transposition, and even horizontal transfer events. We might fall into the "we need more data" trap wherein eventually the signal will drown

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out the noise - but the reality is the noise will likely also increase with more data - and sometimes more complexity. Resolutions from the debates I mentioned earlier came from ingenious experiments carried out by better tools.

Karl Popper once expressed that to be able to test scientifically, it must be falsifiable - and thought experiments are just that "what ifs". Perhaps later on we develop better tools or algorithms that can confirm or deny our outstanding hypotheses later on.