One ARM Project 4 Metagenomics Pipeline

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4	File management	- NCBI AMR Database
	□ BAM file parsing	○ Script to identify ARGs from short-reads
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☼ ShortBRED Read quality control () AMRFinder Initial raw read QC ○ Script to quantify ARGs per sampling site Standard Trimming (Trimmomatic) ☼ ShortBRED Raw Read QC after trimming Script diversity calculations per site Parametric randomization script Parsing QC metrics of all iterations **Taxonomy** Bootstrapping script • Rapid taxonomy via short reads ☐ MetaPhlan4 Pipeline preparation for HPC Genomic context Calculate using quotation from PGC Set up config files Network analysis Taxo-metagenomics pipeline ARGs Associated pathogens ✓ Identification of relevant ARG databases - CARD NCBI AMR Database 8 Complete Script to identify ARGs from short-reads General taxo-metagenomics pipeline Detection of HGT Integration with config folder HGT mechanisms Testing with simulated datasets Pipeline integration FastQC Trimmomatic Kraken2 Bracken △ QHME2 Change QIIME2 to in-house diversity script Aggregate diversity metrics $[\Box]$ File management Compression and decompression scripts

Demultiplexing script

Script Descriptions

Pipelines

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.

Project4 - ARG-MGE.smk

Stage: Draft

General Purpose

This pipeline is designed for comprehensive metagenomic analysis of ARGs, it also includes placeholders for analyses of mobile genetic elements (MGEs), and plasmid detection. It integrates tools for read quality control, assembly, annotation, taxonomic profiling, and structural variant (SV) detection to provide a high-resolution view of the genetic components in metagenomic samples.

Specifics:

1.1 Preprocessing

• It first uses PEAR to merge paired-end reads.

Technical Notes: PEAR (Paired-End reAd mergeR) compares paired-reads to correct infer the 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.

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.

• Followed by conversion from FASTQ to FASTA using seqtk.

Technical Notes: seqtk converts compressed FASTQ files to FASTA.

Rationale: Some tools cannot work on FASTQ formats.

• Translation of the CARD-protein homologues database transeq, and then reverse-translates these proteins to nucleotides for alignment.

Technical Notes: transeq from EMBOSS translates nucleotide sequences to protein sequences based on **standard genetic code**. backtranseq reverses this to allow iterative alignments with nucleotide sequences.

Rationale: This leverages conserved protein-level information, which is lost at the nucleotide 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.

1.2 Metagenomic assembly of ARG-contigs

• Iterative alignment is performed using KMA to align reads against reversetranslated CARD sequences for high-sensitivity identification.

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 databases. Secondly, iterative alignment using this process allows us to contextualize the region 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 iterations KMA creates, increasing the database size.

• Merging ARG-related reads database with PANDASeq.

Technical Notes: PANDASeq is then used to further refine the paired-reads collated in the ARG-related genes database.

Rationale: PANDASeq was chosen as, while being slower than PEAR, it is more accurate. This mergins 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 database.

Merged reads are assembled into contigs using metaSPAdes.

Technical Notes: metaSPAdes is then used to create contiguous sequences from these local regions by extending them using reads from the whole metagenomic pool. Additionally, Contigs are filtered by length here to remove possible artefacts.

Rationale: metaSPAdes was chosen as, while being slower than MEGAHIT, it is optimized in handling highly diverse and mixed microbial populations. CARD is used here because it is manually curated and updated regularly - in order to be included in the database, there must be clinical data (ASTs) involved in the study.

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 that determines the minimum ARG sequence dictated by CARD.

Notes: Contigs are further extended using contigextender to form scaffolds

Notes: Might add other contig extendending programs like **GapFiller** - which leverages mate-pair information

• Python script is created to do another round of checking contig quality for downstream analysis, R scripts are used to visualize the data.

Technical Notes: The Python script will measure standard contig quality metrics e.g. N50, L50 etc.

Rationale: Should be obvious why

• Confirmation of contigs with ARGs using RGI.

Technical Notes: RGI scan the contigs and check whether which contigs created by metaSPAdes have ARGs in them.

Rationale: metaSPAdes may have created contigs that DO NOT contain ARGs, and have instead assembled them into a more matching contig (a false-positive misassembly) - this can happen because of the different databases being used; also parallelization of methods like this increases 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.

Notes: RGI is the official scanner of CARD.

1.3 Read-mapping

• Checking coverage statistics on contigs using raw reads

Technical Notes: Samtools is used here to map the raw reads from the larger database back to the assembled contigs and then calculates the coverage over the entire contig.

Rationale: Read-mapping is a quality control protocol used in metagenomics, to determine the 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 coverage across a contig, that contig could be chimeric, meaning, it could be from two different populations.

Extra note A Python script is included in the pipeline that is determine visualize and check how the coverage changes over contig regions. In general, they could be interpreted as the following:

Smooth, Uniform Coverage: Typically shown by well-assembled contigs.

Sharp Coverage Drop: May need to be split or flagged for reassembly. May also be a misassembly point (chimeric contitg) or caused by a structural variant

Coverage Gaps: Regions with little to no read support; a strong indicator of misassembly.

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

(to be added to the script)

Sharp increase: May be due to repetitive or duplicated regions, the assembler decided to collapse all the reads into that one contig, amplification bias from PCR, HGT, SV, chimeras.

1.3.1 Read mapping more details

Read mapping tools

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 percent identity to the contigs.

Mapper	K-mer Length	Mismatch	Gap Opening	Gap Extension
		Penalty	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 1: Example table of mapper configurations without command example

Rationale: 95 percent 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.

Note (September 18, 2024)

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.

Tool	K-mer	Mismatch	Gap Open-	Gap Exten-
	Length	Penalty	ing Penalty	sion 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: Alignment parameters for ungapped alignments across BWA, Bowtie2, KMA, and Minimap2

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

Table 3: Protein read-mapping parameters for tblastn and blastp

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 protein-protein interactions are altered—perhaps by investigating changes in binding sites. Which is a story for another day (why do I do this to myself?)

1.4 Contiguous chimeras

• On a parallel process: taxonomic profiling is done using Kraken2 and SprayNPray. and aligned with the ARG-raw reads and ARG-contigs databases

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

raw-reads. While SprayNPray complements this by assigning taxonomy at contig level. Rationale: By assigning 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.

• Detects structural variants (SVs) in contigs using Manta and identifies *chimeric* contigs based on SVs and taxonomic classification.

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.

• Or perhaps due to mobile genetic elements like transposons

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

- HMMER3 suite
- Tnppred a transposon predictor tool

Rationale:

• 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 similarity is most often used as a presence or absence diversity metric.

1.4.1 Plasmids

• 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 will calculate GC skews of the chimeric contig and compare it to its taxonomic counterparts. Another Python script 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.

Equation:

$$Plasmid\ Percentage = \left(\frac{Plasmid\ Reads}{Total\ Reads}\right) \times 100$$

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 to do sliding window analysis of GC-skews - as different characteristics of this curve can be interpreted in different ways.

1.5 Phages

• Phage influence signatures

Technical Notes: They will be determined using a variety of tools:

- 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.

• 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 a phylogenetic tree based on the extracted phage genes. For visualization, tools like iTOL or FigTree can be used to generate an interpretable phylogenetic tree.

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 origins of the phage genes found in our dataset and their potential involvement in the dissemination of ARGs.

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?

Project4 - metagenomics_general.smk

Stage: Draft

General Purpose

This pipeline is designed for **The essentials in metagenomics**, which includes **quality checking**, **filtering**, **and trimming of raw reads to clean reads**. As well as the usual **taxometagenomic analysis**.

Specifics:

Purpose: A general pipeline for preprocessing, assembly, and binning of metagenomic reads. **Inputs:**

- Raw sequencing data.
- Reference databases for assembly and binning.

Outputs:

- Assembled contigs.
- Binned genomes.

Key Steps:

- Preprocesses reads with quality control.
- Assembles reads into contigs using MEGAHIT.
- Bins the contigs into genomes using MetaWrap.

Modules

1. FastQC.bash

Purpose: Runs FastQC on sequencing reads for quality control checks. **Inputs:**

• Raw sequencing data in FASTQ format.

Outputs:

• FastQC reports (HTML and text).

Key Steps:

- Runs FastQC on input files.
- Generates quality control reports for each sample.

2. trimmomatic.bash

Purpose: Performs read trimming and filtering to remove low-quality bases and adapters. **Inputs:**

- Raw sequencing data in FASTQ format.
- Adapter sequences for trimming.

Outputs:

• Trimmed reads in FASTQ format.

Key Steps:

- Trims adapters from sequencing reads.
- Filters low-quality bases and short reads.

3. calculate_diversity.py

Purpose: Computes alpha and beta diversity metrics from metagenomic data. **Inputs:**

- Feature table in TSV format.
- Metadata for each sample.

Outputs:

- Diversity indices (e.g., Shannon, Simpson).
- Beta diversity distance matrices (e.g., Bray-Curtis).

Key Steps:

- Loads feature table and metadata.
- Computes diversity metrics for each sample.
- Outputs a summary of diversity indices.

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1.1 To-do List

1.2 Completed Tasks

- \checkmark raw API results now readable stored in TSV file
- \checkmark Created symlinks in private (repo) to .git folders in public repos to enable tracking of Git and Github activities
- ✓ Resolved residual time-tracking activities within the local computer

1.3 Scripts Worked On

✓ checkingAPI.sh (now updated to make the raw API results readable)

1.4 Issues Encountered

•

1.5 Need to Troubleshoot

•

September 17, 2024

1.6 To-do List

- Call for a group meeting regarding logistics
 - ✓ Offer to use landline to minimize on-site visits and reduce transportation costs; as point-persons have not been replying via email as quickly lately
 - ✓ Discuss PGC calculations and talk about allocatable budget
 - ✓ On-site orientation with PGC engineering department (OETS) regarding sampling sites taken care of by James and Roch
 - Raise concerns about data storage and management
 - Inquire if there are scripts I can develop to streamline the bureaucratic process

1.7 Completed Tasks

- Setup wakatime in VS Code and Ubuntu to track my coding productivity
- Helped in receiving and moving the autoclave and fridge needed by the Program

1.8 Scripts Worked On

- rawAPIformatter.sh
- ✓ checkingAPI.sh
- wakatime.sh

1.9 Issues Encountered

- ✓ Connection blockage from wakatime API
- raw API results is unreadable

1.10 Need to Troubleshoot

• conky display of wakatime API for desktop

September 16, 2024

1.11 To-do List

- Call for a group meeting regarding logistics
 - Offer to use landline to minimize on-site visits and reduce transportation costs; as point-persons have not been replying via email as quickly lately
 - Discuss PGC calculations and talk about allocatable budget
 - Raise concerns about data storage and management
 - Inquire if there are scripts I can develop to streamline the bureaucratic process

1.12 Completed Tasks

- ✓ Updated GitHub repositories; created new repositories: Documentation and Confidential
- ✓ Created a new bash script to track progress across all repositories
- ✓ Set up this research diary to document daily progress and tasks
- \checkmark Python script that calculates contig quality metrics L50, N50, but also L90, N90, GC skew etc.
- ✓ Python script that detects and calculates changes in coverage over contigs for read mapping
- ✓ Typesetting: explaining the process and rationale inside the ARG-MGE.smk pipeline
- ✓ Integration of the two scripts to the ARG-MGE pipeline

1.13 Scripts Worked On

- Project4/ARG_MGE.smk updated
- ✓ Project4/calculate_contig_quality.py
- ✓ Project4/plot_and_detect_intermediate_coverage.py

1.14 Issues Encountered

✓ Could not access GitHub, needed to reset the SSH keys and update my Git histories for version control

1.15 Need to Troubleshoot

• None identified at this time