## Commands/Workflow for Binning, Annotations, Anvio, Bin Curation of 2017 Lost City Metagenome

## **##Binning with ABAWACA**

srun prepare\_esom\_files.pl abawaca\_prep\_files H08.contigs.renamed.fa srun abawaca abawaca\_prep\_files/esom.names abawaca\_prep\_files/esom.lrn H08.contigs.renamed.fa abawaca\_bins & H08 bins: 13, 15, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 6, 8

## ##Prodigal Protein Prediction for each Bin

```
prodigal.sh
#! /bin/sh
#SBATCH --output prodigal.out
#SBATCH --error prodigal.err
#SBATCH --ntasks 19
#SBATCH --cpus-per-task 3
#SBATCH --nodes 1-2
#SBATCH --partition highmem
#SBATCH --mem-per-cpu 1G
srun prodigal -q -p meta -f gff -i 13.fasta -o 13.gff -a 13.faa &
srun prodigal -q -p meta -f gff -i 15.fasta -o 15.gff -a 15.faa &
srun prodigal -q -p meta -f gff -i 28.fasta -o 28.gff -a 28.faa &
srun prodigal -q -p meta -f gff -i 29.fasta -o 29.gff -a 29.faa &
srun prodigal -q -p meta -f gff -i 30.fasta -o 30.gff -a 30.faa &
srun prodigal -q -p meta -f gff -i 31.fasta -o 31.gff -a 31.faa &
srun prodigal -q -p meta -f gff -i 32.fasta -o 32.gff -a 32.faa &
srun prodigal -q -p meta -f gff -i 33.fasta -o 33.gff -a 33.faa &
srun prodigal -q -p meta -f gff -i 34.fasta -o 34.gff -a 34.faa &
srun prodigal -q -p meta -f gff -i 35.fasta -o 35.gff -a 35.faa &
srun prodigal -q -p meta -f gff -i 36.fasta -o 36.gff -a 36.faa &
srun prodigal -q -p meta -f gff -i 37.fasta -o 37.gff -a 37.faa &
srun prodigal -q -p meta -f gff -i 6.fasta -o 6.gff -a 6.faa &
srun prodigal -q -p meta -f gff -i 8.fasta -o 8.gff -a 8.faa &
```

# ##Annotations with Diamond for each protein prediction (done 2 times, with both T1000 and prokaryotes databases)

```
annotate.sh
#! /bin/sh
#SBATCH --output diamond.out
#SBATCH --error diamond.err
#SBATCH --ntasks 1
#SBATCH --cpus-per-task 5
#SBATCH -w node2
#SBATCH --partition batch
#SBATCH --mem-per-cpu 10G
diamond blastp --threads 5 --tmpdir /tmp --sensitive --evalue 0.000001 --outfmt 6 --out 13.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 13.faa
diamond blastp --threads 5 --tmpdir /tmp --sensitive --evalue 0.000001 --outfmt 6 --out 15.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 15.faa
```

```
diamond blastp --threads 5 --tmpdir/tmp --sensitive --evalue 0.000001 --outfmt 6 --out 28.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 28.faa
diamond blastp --threads 5 --tmpdir/tmp --sensitive --evalue 0.000001 --outfmt 6 --out 29.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 29.faa
diamond blastp --threads 5 --tmpdir/tmp --sensitive --evalue 0.000001 --outfmt 6 --out 30.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 30.faa
diamond blastp --threads 5 --tmpdir /tmp --sensitive --evalue 0.000001 --outfmt 6 --out 31.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 31.faa
diamond blastp --threads 5 --tmpdir /tmp --sensitive --evalue 0.000001 --outfmt 6 --out 32.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 32.faa
diamond blastp --threads 5 --tmpdir/tmp --sensitive --evalue 0.000001 --outfmt 6 --out 33.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 33.faa
diamond blastp --threads 5 --tmpdir /tmp --sensitive --evalue 0.000001 --outfmt 6 --out 34.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 34.faa
diamond blastp --threads 5 --tmpdir/tmp --sensitive --evalue 0.000001 --outfmt 6 --out 35.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 35.faa
diamond blastp --threads 5 --tmpdir /tmp --sensitive --evalue 0.000001 --outfmt 6 --out 36.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 36.faa
diamond blastp --threads 5 --tmpdir /tmp --sensitive --evalue 0.000001 --outfmt 6 --out 37.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 37.faa
diamond blastp --threads 5 --tmpdir/tmp --sensitive --evalue 0.000001 --outfmt 6 --out 6.KO.b6 -
-db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 6.faa
diamond blastp --threads 5 --tmpdir /tmp --sensitive --evalue 0.000001 --outfmt 6 --out 8.KO.b6 -
-db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 8.faa
diamond blastp --threads 5 --tmpdir/tmp --sensitive --evalue 0.000001 --outfmt 6 --out 14.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 14.faa
diamond blastp --threads 5 --tmpdir /tmp --sensitive --evalue 0.000001 --outfmt 6 --out 20.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 20.faa
diamond blastp --threads 5 --tmpdir/tmp --sensitive --evalue 0.000001 --outfmt 6 --out 23.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 23.faa
diamond blastp --threads 5 --tmpdir /tmp --sensitive --evalue 0.000001 --outfmt 6 --out 26.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 26.faa
diamond blastp --threads 5 --tmpdir/tmp --sensitive --evalue 0.000001 --outfmt 6 --out 27.KO.b6
--db /srv/databases/internal/diamond/KEGG-T10000.dmnd --query 27.faa
#! /bin/sh
#SBATCH --output diamond.proks.out
#SBATCH --error diamond.proks.err
#SBATCH --ntasks 1
#SBATCH --cpus-per-task 5
#SBATCH -w node2
#SBATCH --partition batch
#SBATCH --mem-per-cpu 10G
bins=("13" "15" "28" "29" "30" "31" "32" "33" "34" "35" "36" "37" "6" "8") for b in "${bins[@]}"; do
diamond blastp --threads 5 --tmpdir /tmp --sensitive --evalue 0.000001 --outfmt 6 --out
$b.KO.prok.b6 --db /srv/databases/internal/diamond/KEGG-prokaryotes.dmnd --query $b.faa
done
```

srun --ntasks 1 --cpus-per-task 4 --partition highmem --mem 200G --out checkm.faa.out --err checkm.faa.err checkm lineage\_wf --genes -f H08\_checkm\_table.tsv --tab\_table -t 4 --pplacer\_threads 4 -x faa prodigal\_proteins H08\_checkm\_faa.results &

#### ## I did the next steps to get geneCalls for Anvio

for sample in "13" "14" "15" "28" "29" "30" "31" "32" "33" "34" "35" "36" "37" "6" "8" do

srun -p single python get\_koID.py \${sample}.KO.prok.b6

/srv/databases/proteins/kegg/prokaryotes \${sample}.koID.proks.out &

done

for sample in "13" "14" "20" "23" "26" "27" "15" "28" "29" "30" "31" "32" "33" "34" "35" "36" "37" "6" "8"

do

srun python get\_koID.py \${sample}.KO.b6 /srv/databases/proteins/kegg/KO \${sample}.koID.KO.out &

done

## ##Annotate Features on metagenome (NOT BIN) from all .gffs from prodigal and b6 files from diamond

srun cat \*.gff > H08.gff

awk -F'\t' '/^#/{print \$0; next} {sub(/ID\=[^\_]\*\_/, "ID="\$1"\_", \$9); print

\$1"\t"\$2"\t"\$3"\t"\$4"\t"\$5"\t"\$6"\t"\$7"\t"\$8"\t"\$9}' H08.gff > H08.clean.gff

srun cat annotations/diamond\_output/\* > all.KO.b6

srun -p highmem --mem 70G annotate\_features --type CDS --mapping

/srv/databases/internal/json/KEGG-T10000.mapping.json,/srv/databases/internal/json/KEGG-prokaryotes.mapping.json --fields

gene,gene\_family,product,organism,database,md5,Ontology\_term,dummy --conflict quality --out H08.annotated.gff --b6 annotations/diamond\_output/all.KO.b6 binning/abawaca\_bins/final-clusters/H08.clean.gff > H08.annotate.log &

### ##Count features to calculate gene coverage for metagenome

srun -p highmem --mem 70G count\_features --attr gene --sorting name --units fpk,fpkm --out H08.gene.abundance data/H08.mapped.sorted.bam H08.annotated.gff 2> H08.fpk.gene.log &

#### ##Bin Refining

Pull out all contigs with same PhyloPythia id in bin, make new .fasta and .faa files just this list of contigs:

srun python ~/myScripts/makeFastaBin.py -c c d contigs in .txt file> -f <bin .fasta file from abawaca> -o <new .fasta file>

srun python ~/makeFaaBin.py -c d contigs in .txt file> -f <.faa file to be modified> -o <new .faa file>

#### ##CheckM on new bin with all same PhyloPythia id

srun --ntasks 1 --cpus-per-task 4 --partition highmem --mem 100G --out <outfile name> --err <errorfile name> checkm lineage\_wf --genes -f <sample. checkM table name> --tab\_table -t 4 --pplacer\_threads 4 -x <new .faa file> <directory of that file> <results\_name> &

#### ##Gene Call for Anvio

srun -p single python ~/myScripts/geneCall.py -f <new .fasta\_file> -k <get\_koID\_outfile> -a <new .faa file> -g <geneCall outfile> -o <function.outfile> -s <database name> &

#### ##Generate the Anvio DB Files

srun -p single --mem 10G --err <error.out> --out <out.out> anvi-gen-contigs-database -f <new .fasta file> -o anvio/\$sample.contigs.db --external-gene-calls <geneCall outfile> & srun -p single --mem 10G --out <out.out> --err <error.err> anvi-run-hmms -c anvio/\$sample.contigs.db &

## ##<function.outfile> from geneCall.py

srun -p single --mem 10G --err <error.err> --out <out.out> anvi-import-functions -c anvio/\$sample.contigs.db -i <function.outfile> & ##Map

srun -p batch --out <out.out> --err <error.err> --mem 20G bowtie2-build <new .fasta> <new.bin.name> &

sbatch bt.sh

-----

bt.sh

#! /bin/sh

#SBATCH --output bt2.out

#SBATCH --error bt2.err

**#SBATCH --ntasks 1** 

#SBATCH --cpus-per-task 4

**#SBATCH** --partition batch

#SBATCH --mem 20G

bowtie2 -p 4 --very-sensitive -I 130 -X 174 -x <> --interleaved

<sample>.interleaved.atrim.decontam.qtrim.derep.fq.gz 2> bowtie.mapping.log | samtools sort - @ 1 -m 10G -l 9 -O bam -T <new.bin.name> -o <new.bin.name>.mapped.sorted.bam

### ##Generate the PROFILE file for anvio

srun -p batch --mem 20G --error <error.err> --output <out.out> anvi-profile -i <mapped.sorted.filt.derep.bam for .fasta file anvio db made with> -c anvio/\$sample.contigs.db --min-contig-length 3000 --sample-name \$sample --output-dir anvio/\$sample --cluster-contigs & anvi-interactive -p \$sample.PROFILE.db -c \$sample.contigs.db

#### ##Visually split bin by basal branching, rerun checkM

#### ##Annotate features

#### ##Combine both T1000 and prokaryotes databases together to make one .b6 file

cat annotations/diamond\_output/<bin>.KO.\* > <bin>.all.b6

srun -p single --mem 50G annotate\_features --type CDS --mapping

/srv/databases/internal/json/KEGG-T10000.mapping.json,/srv/databases/internal/json/KEGG-prokaryotes.mapping.json --fields

gene,gene\_family,product,organism,database,md5,Ontology\_term,dummy --conflict quality --out <br/> <bin>.annotated.gff --b6 <bin>.all.b6 <bin>.clean.gff 2> <bin>.annotate.log

## ##Get annotated .gff for refined bin

srun python ~/myScripts/getgffBin.py -f <anvio split .fasta bin file> -g <bin>.annotated.gff -o <new bin>.annotated.gff

### ##Make mapping file for KEGG Mapper Online Tool

 $srun ~ \mbox{'myScripts/genesKO\_from\_gff.py < new bin>.annotated.gff < new bin>.forMapping.txt}$