

Cold genes HMM

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Following anvio phylogenomics tutorial

<https://merenlab.org/2017/06/07/phylogenomics/>

Making own HMM collection

<https://merenlab.org/2016/05/21/archaeal-single-copy-genes/>

Get hmm-hits-matrix-txt

<https://anvio.org/help/main/programs/anvi-script-gen-hmm-hits-matrix-across-genomes/>

Making HMM file for anvio

HMM models downloaded from EggNOG website.

Custom HMM folder contains:

genes.hmm.gz - concatenated hmm profiles from EggNOG

genes.txt - list of genes + accession + source

kind.txt - gene

target.txt - AA:GENE

noise_cutoff_terms.txt - E 1e-30

Genomes used

Assembled using metaSPAdes, binned using MaxBin2 and scaffolded using SSPACE. Contamination and completeness estimated with CheckM.

D1: Joyce_1_Leptolyngbya

Scaffolds - 152

Largest Scaffold - 355994

N50 - 87779

Contamination - 1.57

Completeness - 97.2%

D2: Fryxell_1_Phormidesmis

Scaffolds - 322

Largest Scaffold - 272946

N50 - 31154

Contamination - 0.54

Completeness - 99.1%

D3: Fryxell_2_Leptolyngbya

Scaffolds - 232

Largest Scaffold - 237628

N50 - 49850

Contamination - 0.86

Completeness - 99.29%

D4: Fryxell_3_Anabaena

Scaffolds - 93

Largest Scaffold - 281882

N50 - 75233

Contamination - 0.88
Completeness - 100%

On anvio...

Using anvio 7.1

Pre analysis - Reformatted fasta file names:

```
anvi-script-reformat-fasta D1.sspace.final.scaffolds.fasta -o D1-contigs-fixed.fa -l 0 --simplify-names
```

1. Generate contigs database - each FASTA file should have a file with the same name that ends with '.db'.

```
for i in `ls *fa` | awk 'BEGIN{FS="."}{print $1}'
do
    anvi-gen-contigs-database -f $i.fa -o $i.db -T 4
    anvi-run-hmms -c $i.db -H Cold_HMM/
done
```

2. Use the program anvio-get-sequences-for-hmm-hits to get sequences out of genomes. 'external-genomes.txt' is list of genomes and their path.

```
anvi-get-sequences-for-hmm-hits --external-genomes external-genomes.txt \
                                --hmm-source Cold_HMM \
                                -o cold-genes-dna.fasta

anvi-get-sequences-for-hmm-hits --external-genomes external-genomes.txt \
                                --hmm-source Cold_HMM \
                                --get-aa-sequence \
                                -o cold-genes-aa.fasta
```

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3. Get table of hits

```
anvi-script-gen-hmm-hits-matrix-across-genomes --external-genomes external-genomes.txt \
                                                --hmm-source Cold_HMM \
                                                -o output.txt
```

Final files

output.txt - table of hmm hits for each genome
cold-genes-aa.fasta - amino acid sequences of hmm hits
cold-genes-dna.fasta - dna sequences of hmm hits