

Genomics - Walkthrough

Notebook: Thesis Methods

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Author: Jess Friedersdorff

Genomics - Phylogenetic Tree Using Set of 40 Conserved Genes

Building a BLAST database:

- First, build a BLAST database using a file containing all of the template genes for each of the 40 cogs.
 - (There are over 1000 template genes, each from a different organism or species which these 40 genes have been found in).
- To do this, create a FASTA file and use DIAMOND (DIAMOND V0.7.9):

```
for f in *.fa; do sed -i "s/>/>$f\s/" "$f"; done

cat *.fa > ALLCOG.faa

diamond makedb --in ALLCOG.faa -d ALLCOG_database

for i in prokka_*; do cd $i; for j in *.faa; do sed "s/>/>$j./g" $j; done; cd
../; done > faafiles.faa
```

- The database was then queried using the .faa file of (all of the) Hungate species' genes using DIAMOND with the following script:

```
diamond blastp -d path/to/database.dmnd -q path/to/faafiles.faa -a path/to/output
diamond view -a path/to/input -o pat/to/output.tab -f tab
```

Nearly all of the 494 genomes have all 40 of The 40 Genes. The fewest that a bacteria had was 31:

```
more Hungatererun_vs_40genes_results.tab | awk 'BEGIN{prev=""};{if($1 != PREV){
print $0; PREV=$1;}}' > tophits_cog.txt
```

- Since it is always sorted by Bitscore, the top hit for each gene is the best. This does mean that a single cog might occur more than once in more than one gene per genome and doesn't account for this.

```
cat tophits_cog.txt | sed 's/sp\./sp/g' | awk -f ./awk_script | sort >
cogsforeachbacteria.txt
```

Where the awk script was:

```
#Script to give to awk.
#finds the highest bit score (value in the 12th column) for every unique
combination of bacteria and cog (column 1 and column 2)

{
sub("\.\.*", "", $1);
sub("\.\.*", "", $2);
if ($12 > vals[$1"."$2]) {
vals[$1"."$2] = $12;
}
}
```

```
END {
  for (key in vals) {
    print key "\t" vals[key];
  }
}
```

- The result was a file, with lines consisting of the name of each bacteria, the name of the cog and the bit score that cog best hit at. Then, the amount of cogs for each bacteria was counted:

```
cat cogforeachbacteria.txt | cut -f1 -d'.' | uniq -c >
count_cogforeachbacteria.txt
```

- Which lead to the findings that nearly all of the 494 have all 40 of The 40 Genes. The fewest that a bacteria had was 31.

Extracting Data to Use Further:

40 individual FASTA files, one for each cog, containing the sequences in the Hungate genomes was made.

- Ordered the query blast results first by cog name, then organism name, then bitscore:

```
sed 's/\.fa[as]/ /g' Hungatererun_vs_40genes_results.tab | sort -k3,3 -k1,1 -
k14,14rn > Hungatererun_vs_40genes_results.sorted.tab
```

- Then, using awk, find the first instance of the cog in the genome, keeping the gene name. Only those with bitscores above 60 are kept. (Some cogs only hit once per genome and some of these had low bitscores. Setting it to 60 removes those. Some missing data like this is not a problem when building trees)

```
cat Hungatererun_vs_40genes_results.sorted.tab | awk 'PREV{genome=""; cog=""};
{if($1 != genome || $3 != cog) { genome=$1; cog=$3; if($14 > 60) { print $0}}}' >
Hungatererun_vs_40genes_results.matches.tab
```

- So that the .faa file of Hungate genes can be searched, the file needs to be changed from a two line FASTA to a single line FASTA (name \t sequence \n name \t sequence etc).

```
$ cat faafiles.faa | cut -d' ' -f1 | sed '/>/s/$/\t/g' | sed 's/>/#>/g' | tr -d
'\n' | tr '#' '\n' | sed '1d' | tr -d '>' > editedfaafiles.faa
```

- Using the file with the best matches of 40 genes to Hungate genomes, search the Hungate file of genes, and extract the name and sequence, and output this to 40 different files, one for each Cog.

```
$ cat Hungatererun_vs_40genes_results.matches.tab | awk 'BEGIN{ while(getline <
"editedfaafiles.faa"){ split($0, a, "\t"); array[a[1]]=a[2]; }}{ printf
">"$1$2"\n"array[$1].faa"$2"\n" >> $3".faa" }'
```

Alignment of 40 COGs using MUSCLE:

MUSCLE v3.8.31 (Edgar, R.C. Nucleic Acids Res 32(5), 1792-97)

Example command:

```
$ muscle -in COG0012.faa.1 -out COG0012.faa.1.aligned
```

Concatenating Sequences Together using Catsequences

Now all of the sequences need concatenating together, to create one FASTA alignment file, where each entry in the file is one long sequences of the aligned 40 genes pasted together into one long sequence.

To do this, a programme called Catsequences was used, available at [Creevey GitHub](#).

- A list of files was created in the directory COG.faa.aligned_files:

```
ls *COG*.faa_aligned > listoffiles.txt
catsequences.exe listoffiles.txt
cat allseqs.fas | tr '?' '-' > allseqs_Hungate_Cog.fas
```

This file was then copied into the main directory (40genes) and given to RAxML.

Tree Building with RAxML:

RAxML version 8.2.9

Tree was built using this command:

```
raxmlHPC-PTHREADS-SSE3 -f a -N 100 -x 6543 -T 4 -m PROTGAMMALG -p 12345 -s
allseqs_Hungate_Cog.fas -n RAxML_COG2_bootstraps
```

Resulting tree could be seen here: <https://itol.embl.de/tree/14412410566444881508838827> (as of 15th September 2020)

Making Count Tables:

Requires a file containing a list of things to count, the file(s) where these things should be counted in, then querying whether the thing to be searched is present, so that if it is not, this can also be recorded, and if it is, then it can be counted. for each file, these are outputted to a new file that contains just the two columns (or even one column with counts) then are pasted together to make a new table.

For example, for EGN gene family counts (where HAP_NAP_SAP.list contains the species of interest which are the things to count and the .faa are gene family files containing the things that should be counted):

```
for j in *.faa; do
  for i in $(cat HAP_NAP_SAP.list); do
    count=$(if grep -q ${i} ${j};
      then grep -c ${i} ${j};
      else echo "x";
    fi);
    echo -e ${j}"\t"${count} >> ${i}.col;
  done;
done

for i in *.col; do
  echo ${i} > ${i}.edit;
  cat ${i} >> ${i}.edit;
done

paste *.col.edit > HAP_NAP_SAP_EGN_counts.table
```

bactCOGs and functional counts in genomes

Use the @2 (bacterial functions) as a high up hierarchy of bacterial groups:

First make a mapping file, for all the genes with the "@2" function:

```
for i in *.annotations; do awk 'BEGIN{FS=OFS="\t"}{split($19, a, ","); for (i in a) {if(a[i] ~ /@2$/ ) {print $1, a[i]} } } ' ${i}; done > gene_to_bactOG_map.list
```

output: (one line per COG, not per gene!)

```
ECADDMPC_03180  COG1690@2
ECADDMPC_03181  COG2304@2
ECADDMPC_03182  COG2819@2
ECADDMPC_03183  COG4870@2
ECADDMPC_03183  COG5263@2
ECADDMPC_03184  COG1578@2
ECADDMPC_03185  COG4633@2
```

Then make one per sample:

```
for i in *.annotations; do awk 'BEGIN{FS=OFS="\t"}{split($19, a, ","); for (i in a) {if(a[i] ~ /@2$/ ) {print $1, a[i]} } } ' ${i} > ${i}_bactOGonly; done
```

also make a list of all the bactOGs that have occurred in the annotations:

```
for i in *.annotations; do awk 'BEGIN{FS=OFS="\t"}{split($19, a, ","); for (i in a) {if(a[i] ~ /@2$/ ) {print $1, a[i]} } } ' ${i}; done | cut -f2 | sort | uniq > list_of_bactOGs.list
```

```
wc -l list_of_bactOGs.list
```

```
4916 list_of_bactOGs.list
```

There are **4916** bacterial OGs. It seems that an annotation can't have an X@2 and a COGX@2. but can have multiple COGX@2.

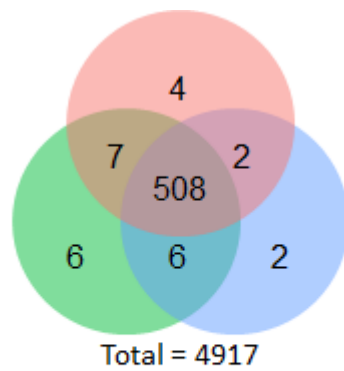
Then for each of those bactOGs that have occurred in any sample, count how many times they occur in each sample:

```
for i in *bactOGonly; do for j in $(cat list_of_bactOGs.list); do count=$( grep -wc ${j} ${i}); echo -e ${j} "\t" ${count} ; done > ${i}.col; done
```

```
paste *.col > bactOG_counts.tsv
```

Counts of bactCOG families shared between the ammonia phenotype groups, where all three species must have a representative.

shared/unique OGs	HAP	SAP	NAP	ALL
HAP	4	2	7	508 Total = 4917
SAP	2	2	6	
NAP	7	6	6	



Number of Bacterial orthologous genes
(OGs) shared between and unique to
HAPs, NAPs, SAPs

4 COGs for HAPs

COG0384@2 - isomerase, unknown function

COG1292@2 - Cell wall/membrane/envelope biogenesis; Belongs to the BCCT transporter (TC 2.A.15) family

COG2202@2 - Signal transduction mechanisms; Pas domain

COG5505@2

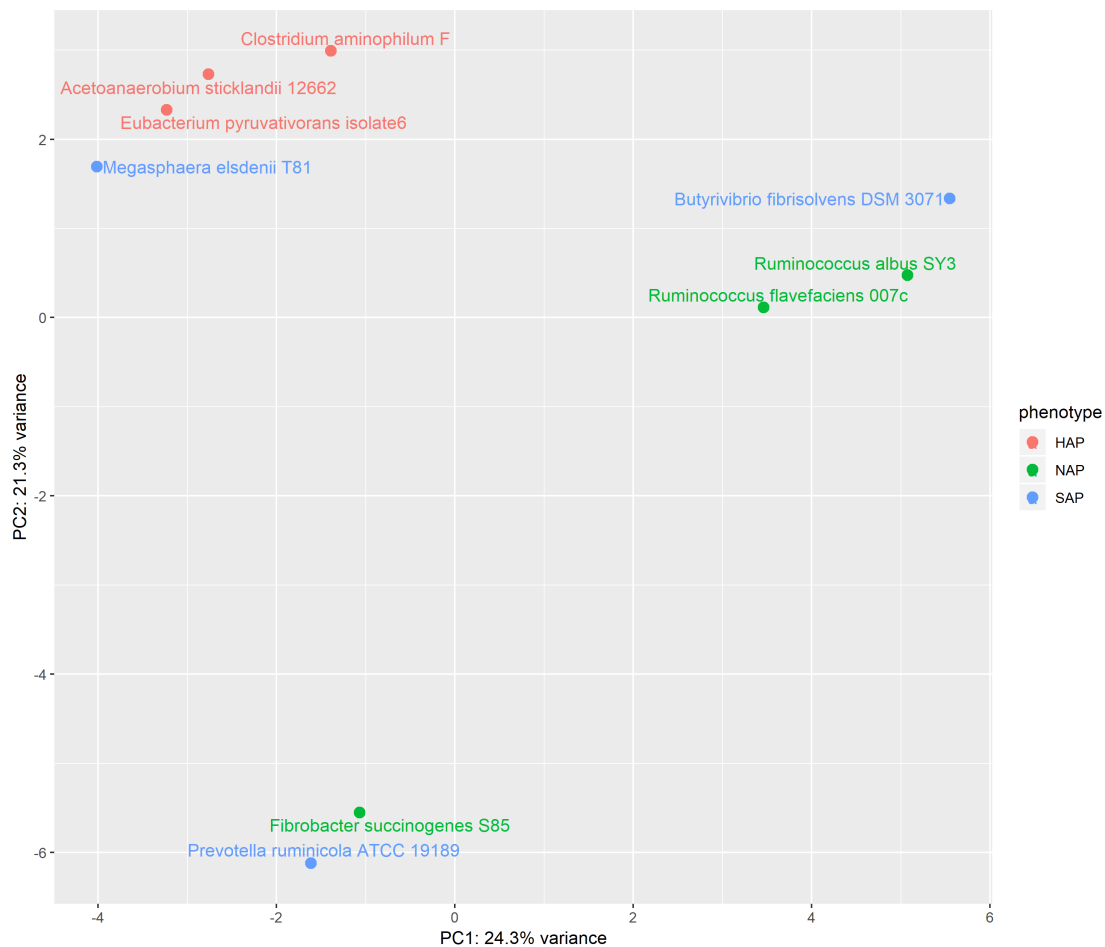
2 for HAPs SAPs

COG1063@2 - Amino acid transport and metabolism; alcohol dehydrogenase

COG4122@2 - Amino acid transport and metabolism; O-methyltransferase activity

R:

Use the R script genomes_bact_OGs_pcas.R



Loadings:

7 components needed to describe 94.285% of variance.

COG0395 influences PC1 most - inorganic ion transport and metabolism; glycerophosphodiester transmembrane transport

COG0823 influences PC2 most - Intracellular trafficking, secretion, and vesicular transport; Involved in the tonB-independent uptake of proteins

Making proportional stacked graphs for COG functional identifiers in genomes

First make a count table for the numbers of genes in each of the categories for each of the genomes: (COG Functional Category)

```

INFORMATION STORAGE AND PROCESSING
[J] Translation, ribosomal structure and biogenesis
[A] RNA processing and modification
[K] Transcription
[L] Replication, recombination and repair
[B] Chromatin structure and dynamics

CELLULAR PROCESSES AND SIGNALING
[D] Cell cycle control, cell division, chromosome partitioning
[Y] Nuclear structure
[V] Defense mechanisms
[T] Signal transduction mechanisms
[M] Cell wall/membrane/envelope biogenesis
[N] Cell motility
[Z] Cytoskeleton
[W] Extracellular structures
  
```

[U] Intracellular trafficking, secretion, and vesicular transport
 [O] Posttranslational modification, protein turnover, chaperones

METABOLISM

[C] Energy production and conversion
 [G] Carbohydrate transport and metabolism
 [E] Amino acid transport and metabolism
 [F] Nucleotide transport and metabolism
 [H] Coenzyme transport and metabolism
 [I] Lipid transport and metabolism
 [P] Inorganic ion transport and metabolism
 [Q] Secondary metabolites biosynthesis, transport and catabolism

POORLY CHARACTERIZED

[R] General function prediction only
 [S] Function unknown

obtained from: <ftp://ftp.ncbi.nih.gov/pub/wolf/COGs/COG0303/fun.txt> (accessed 06/01/2020)

```
for i in $(cat COG_functional_identifiers.list | cut -f1); do for j in
*.annotations; do name=$(echo ${j} | cut -d "." -f1); count=$(cut -f21 ${j} |
grep -c ${i}); echo -e "${i}"\t"${count}" >> ${name}_func.col; done; done
for i in *.col; do echo -e "\t" ${i} > ${i}.edit; cat ${i} >> ${i}.edit; done
paste *.edit > COG_functional_identifiers_counts.table
```

Use the insert -> graph -> percentage stacked graph -> switch rows/columns to get a stacked graph in Excel.

Transporters - Amino acids and peptides.

Table of transporter codes:

1.B.83	The Alpha-Aminoxy Acid Channel (AAAC) Family	AAAC
2.A.118	The Basic Amino Acid Antiporter (ArcD) Family	ArcD
2.A.3	The Amino Acid-Polyamine-Organocation (APC) Family	APC
2.A.95	The 6 TMS Neutral Amino Acid Transporter (NAAT) Family	NAAT
3.A.1.3	The Polar Amino Acid Uptake Transporter (PAAT) Family	PAAT
3.A.1.4	The Hydrophobic Amino Acid Uptake Transporter (HAAT) Family	HAAT
2.A.120	The Putative Amino Acid Permease (PAAP) Family	PAAP
2.A.42	The Hydroxy/Aromatic Amino Acid Permease (HAAAP) Family	HAAAP
2.A.18	The Amino Acid/Auxin Permease (AAP) Family	AAP
2.A.23	The Dicarboxylate/Amino Acid:Cation (Na ⁺ or H ⁺) Symporter (DAACS) Family	DAACS
2.A.26	The Branched Chain Amino Acid:Cation Symporter (LIVCS)	LIVCS

	Family	
2.A.21.2	The Solute:Sodium Symporter (SSS) Family	SSS
2.A.25	The Alanine or Glycine:Cation Symporter (AGCS) Family	AGCS
2.A.78	The Branched Chain Amino Acid Exporter (LIV-E) Family	LIV-E
2.A.114	The Putative Peptide Transporter Carbon Starvation CstA (CstA) Family	CstA
2.A.67	The Oligopeptide Transporter (OPT) Family	OPT
2.A.17	The Proton-dependent Oligopeptide Transporter (POT/PTR) Family	POT/PTR
1.A.11	Ammonium channel	Amt

Get all the tcdb fasta files:

<http://www.tcdb.org/public/tcdb> downloaded 14/01/2020,

- [TCDB FastA Sequences](#)  (Last modified: January 14 2020)

Made a blast database:

```
makeblastdb -in tcdb.fasta -out tcdb -dbtype prot
```

Make submission file and blast the .faa files of the HAPs, NAPs and SAPs against the database:

```
subheader.sh blast_tcdb 25G > blast_tcdb.sub
for i in *.faa; do echo "blastp -query "${i}" -db tcdb -out "${i}"_tcdb -outfmt
6"; done >> blast_tcdb.sub
```

Get tophits of results:

```
for i in *_tcdb; do awk 'BEGIN {prev=""; score=""} {if ($1 != prev && $12 >= 60)
{print $0; prev=$1}}' "${i}" > "${i}_tophits; done #Remove any hits that are under
60 bitscore
```

make a file of just the amino acid transporters codes.

```
for i in *_tophits; do grep -f aatransporters.list "${i}" >
"${i}_aatransporteronly; done
```

Count the occurrences of each of the transporters:

```
for i in *_tophits; do name=$(echo "${i}" | cut -d "." -f1); echo -e
"transportercode" "\t" "${name}" > "${name}_aatransporter.col; for j in $(cat
aatransporters.list); do count=$(grep -c "${j}" "${i}"); echo -e "${j}" "\t"
"${count}" >> "${name}_aatransporter.col; done ; done
```

Paste columns and make table:

```
paste *col > aa_transporter_counts.tsv
```

Then run the R script genomes_aatransporters.R and make the graph:

