Finding Orthologs and Paralogs

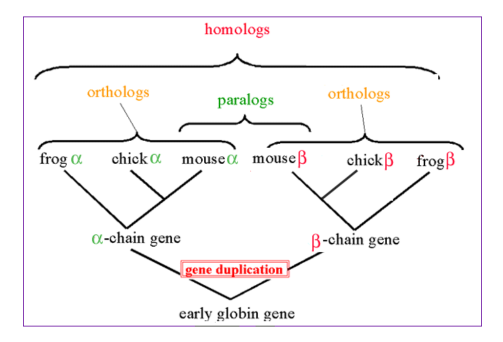


Figure 1: Orthologs

Gene families and Orthology

Problem: How to find "same" genes across multiple species.

Genes can duplicate (Paralogs) and can be identical due to descent (Ortholog)

Methods

- BLAST: reciprocal BLAST

Trees can help resolve relationships

Best hits can sometimes be wrong (B) though it can be resolved with phylogenetics.

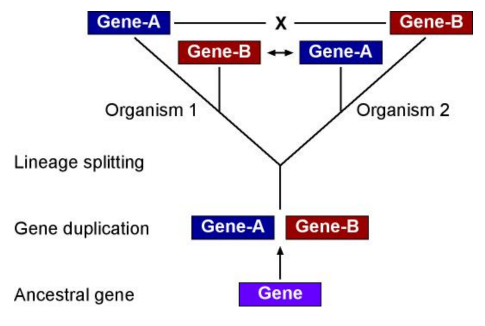


Figure 2: orthologs

Reciprocal Searches

• Bi-directional or Reciprocal BLAST

Implement Bidirectional

Method to find best top hit in one direction and the reverse.

Let's walk through the code

 $Will\ write\ this\ in\ Python\ in\ Class$

Clustering

- Lumping genes together based on similarity linkage
- Single-linkage means if there is a link between A-B then they are in a cluster

Code up single-linkage

Let's look at some code.

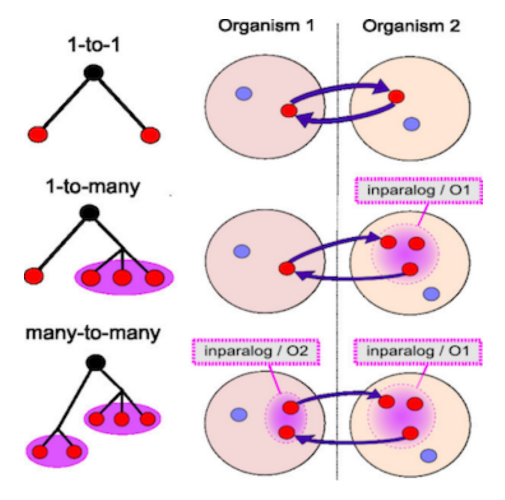


Figure 3: diagramorth

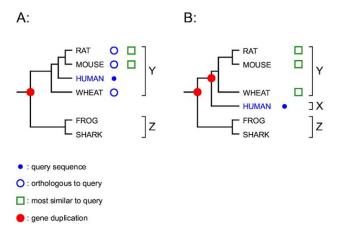


Figure 4: RIO

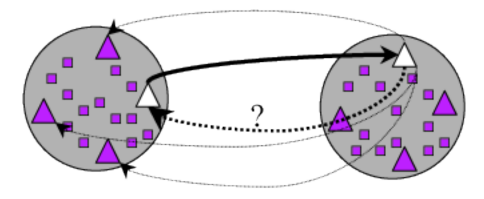


Figure 5: BRH

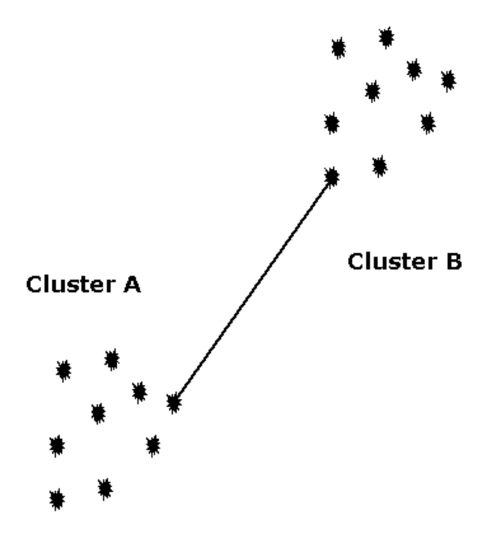


Figure 6: SingleLinkage

Issues

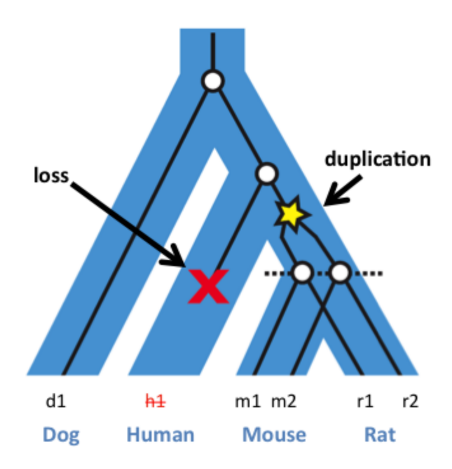


Figure 7: orthologsloss

Tools to go after Orthologous and Paralogous equences

• OrthoFinder

Steps to build orthologs on cluster

#!/usr/bin/bash

We will take 3 datasets of annotated Cyanobacteria, download and run analysis to generate Ortholog table.

```
#SBATCH --ntasks 16 --mem 8G -p short
module load ncbi-blast
module load orthofinder
module load miniconda2
CPU=8
mkdir -p cyanobacteria
cd cyanobacteria
curl -L -O ftp://ftp.ensemblgenomes.org/pub/bacteria/release-45/fasta/bacteria_10_collection
curl -L -O ftp://ftp.ensemblgenomes.org/pub/bacteria/release-45/fasta/bacteria_O_collection,
curl -L -O ftp://ftp.ensemblgenomes.org/pub/bacteria/release-45/fasta/bacteria_4_collection,
# uncompress files and name them all *.fasta
for file in *.fa.gz
do
m=$(basename $file .pep.all.fa.gz)
pigz -dc $file > $m.fasta
done
cd ..
```

Ortholog results

orthofinder.py -a \$CPU -f cyanobacteria

The output file by default will be the date of the anlaysis. Opening the file cyanobacteria/Results_XXX/Orthogroups.txt but I made a folder in the examples you look over. Here's one table

Format

```
GroupName\tSp1_Gene1, Sp1_Gene2\tSp2_Gene1, Sp2_Gene2\tSp3_Gene1, Sp3_Gene2
```

Cyanobacterium_aponinum_pcc_10605.ASM31767v1 Nostoc_punctiforme_pcc_73102.ASM2002v1 0G0000000 EKQ66605, EKQ66611, EKQ66662, EKQ66782, EKQ66954, EKQ66984, EKQ67084, EI EKQ67433, EKQ67590, EKQ67680, EKQ67799, EKQ67807, EKQ67983, EKQ68026, EKQ68032, EKQ68054, I EKQ69279, EKQ69300, EKQ69345, EKQ69368, EKQ69506, EKQ69549, EKQ69629, EKQ69630, EKQ69655, EI KQ70786, EKQ70840, EKQ70870, EKQ70894, EKQ71088, EKQ71090, EKQ71265, EKQ71335

OGO000001 AFZ52442, AFZ54265, AFZ54640 ACC78968, ACC78978, ACC79054, ACC79090, ACC7913881797, ACC82091, ACC82628, ACC82978, ACC83035, ACC83215, ACC83711, ACC84528, ACC84844, ACC869971, EKQ69995, EKQ70003, EKQ70556, EKQ70833, EKQ71286

ACC79344, ACC80485, ACC80595, ACC82143, ACC82836, ACC82962, ACC83849 DG0000002 AFZ55137 ACC84972, ACC84974, ACC84981, ACC84982, ACC84983, ACC85032 EKQ66950, EKQ67597, EKQ67615, El DG0000003 AFZ53198 ACC78875, ACC78976, ACC79256, ACC79524, ACC79759, ACC80145, ACC80528 ACC82769, ACC83025, ACC83081, ACC83457, ACC83602, ACC83721, ACC83749, ACC84422, ACC85331 ACC80422, ACC80525, ACC80662, ACC80851, ACC80857, ACC80914, ACC81440, ACC81 DG0000004 6, ACC83981, ACC84622, ACC84732, ACC85457 EKQ66830, EKQ66911, EKQ67039, EKQ67311, EKQ6999 AFZ52318, AFZ52611, AFZ52613, AFZ52925, AFZ52973, AFZ53626, AFZ53840, AFZ53841, DG0000005 CC82559, ACC83603, ACC83674, ACC85005, ACC85009 EKQ67574, EKQ67809, EKQ69976 AFZ52319, AFZ53394, AFZ54017, AFZ54472 ACC79360, ACC79745, ACC79853, ACC80832, 478, EKQ67551, EKQ67724, EKQ67810, EKQ68266

OGO000007 AFZ53704, AFZ54461, AFZ54462 ACC79786, ACC80242, ACC80282, ACC80538, ACC807682, EKQ68369, EKQ70142, EKQ70145, EKQ71300

The tool also generates summary statistics we can look through.

Could write a script to turn this into a table or use the summary count table provided.

ORTHOLOG_GRP SP1 SP2 SP3
ORTHO_0001 10 5
ORTHO_0002 1 1