Phylogenomics and Population Genomics: Inference and Applications

ORTHOLOGY PREDICTION FOR PHYLOGENOMIC ANALYSES

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Before starting:

Log in to your session

ssh username@ec2-34-242-61-70.eu-west-1.compute.amazonaws.com

Copy the github session into your main folder:

svn export

https://github.com/ppgcourseUB/ppgcourse2022/trunk/Orthology_prediction_for_phylogenomic_analyses.MARINA_MARCET

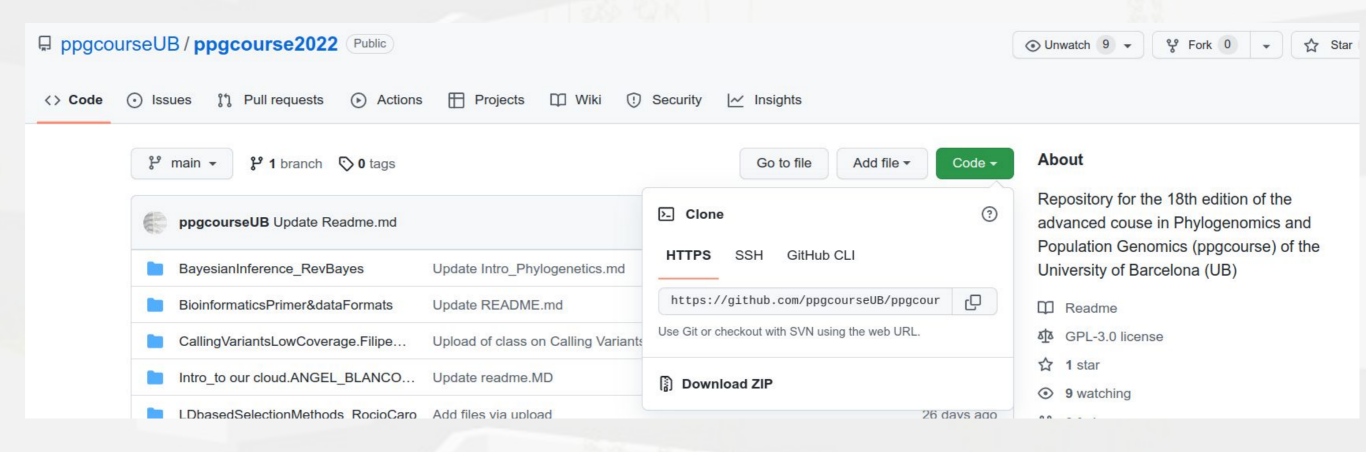
Move into the folder of the session:

cd Orthology_prediction_for_phylogenomic_analyses.MARINA_MARCET/



Just in case the svn export does not work:

Copy the main repository into your computer:



- Unzip the file
- Use scp -r folderName username@ec2-34-242-61-70.eu-west-1.compute.amazonaws.com:///home/username/

How to use VIM to edit files

(You can also use emacs if you prefer)

To open a file: vim fileName

```
##This is a script to run orthofinder

#SBATCH -p normal

#SBATCH -c 8

#SBATCH --mem=6GB

#SBATCH --job-name orthofinder-job01

#SBATCH -e %j.out

#SBATCH -e %j.err

#module loadding. Check available modules with `module avail`
module load orthofinder

#running orthofinder

orthofinder -f proteomes -t 8 -a 2
```



How to use VIM to edit files

(You can also use emacs if you prefer)

Before you start to write, press

i

```
#!/bin/bash

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-- INSERT -- 1,1 All
```

You now should have the word insert at the bottom



How to use VIM to edit files

(You can also use emacs if you prefer)

 Once you have edited what you wanted, press ESC (you will see that the --- insert --- will disappear)

```
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```

Now to save write



and press enter



Reminder: How to move through the terminal

To go to a folder:

cd folderName/folderName1

To move back to the previous folder:

cd ..

If you're completely lost:

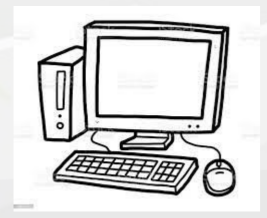
cd

This will just bring you to your home folder



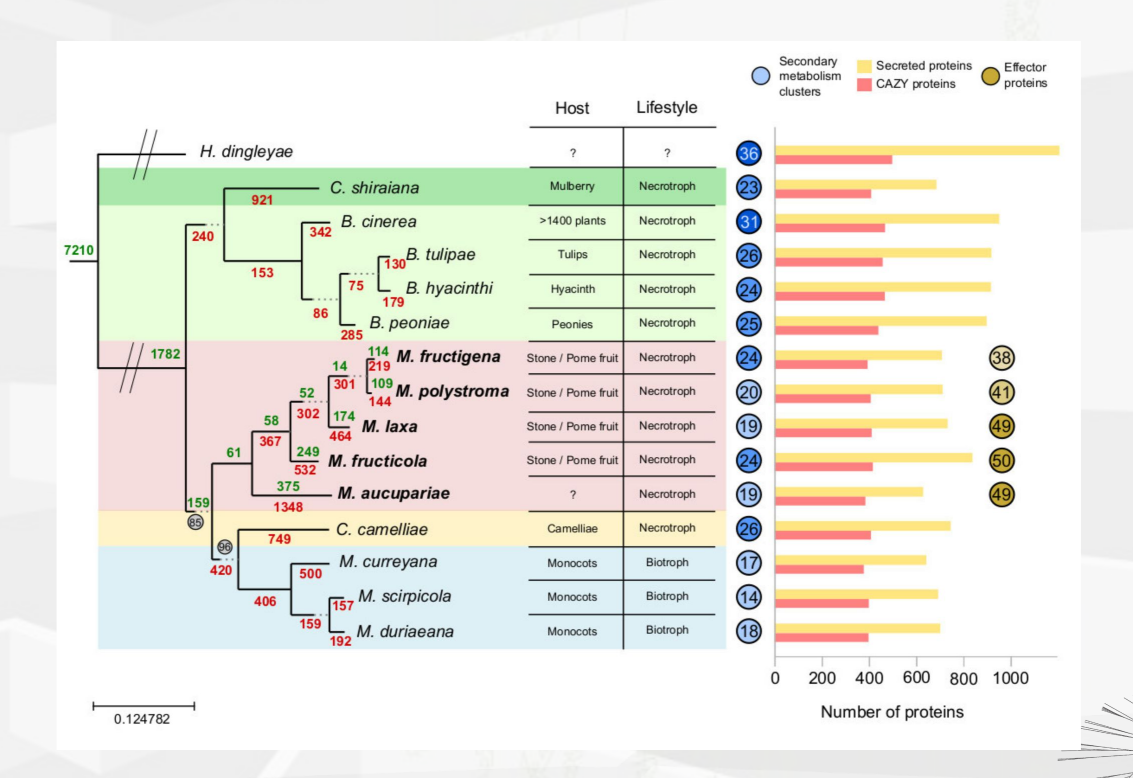
About me:

- Degree in Biochemistry URV
- PhD in Fungal evolution Centro de investigación Príncipe
 Felipe and Center for Genomic Regulation
- Center for Genomic Regulation
- Barcelona Supercomputing Center & Institut de Recerca
 Biomedica





About my research:



Outline

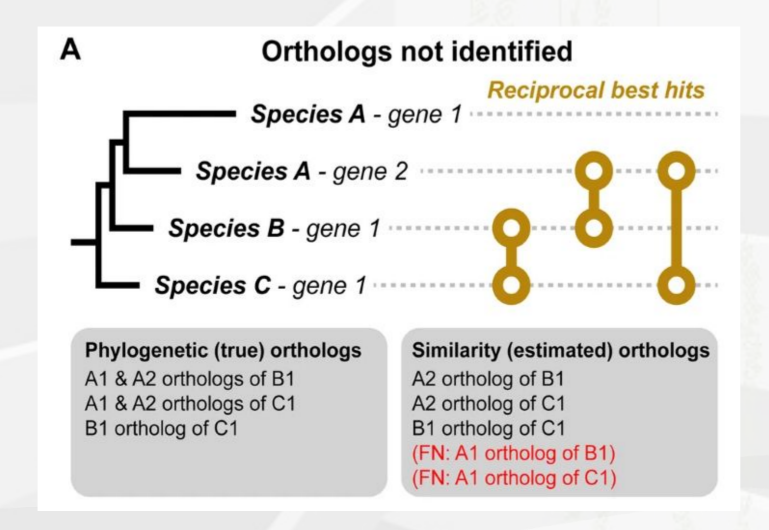
- Reminder
- Previous considerations
- OrthoFinder

Reminders

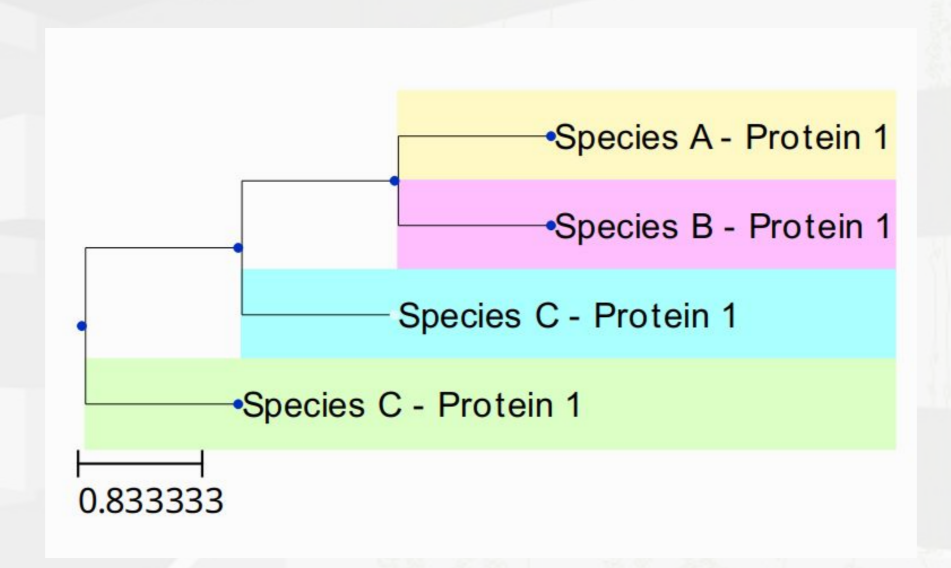
Homologs: Sequences that descend from a common ancestor.

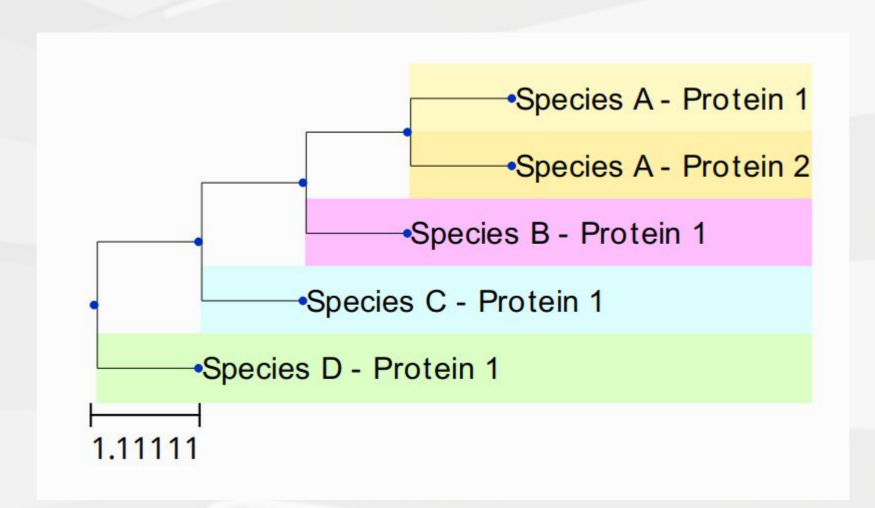
Orthologs: Sequences that come from a speciation event.

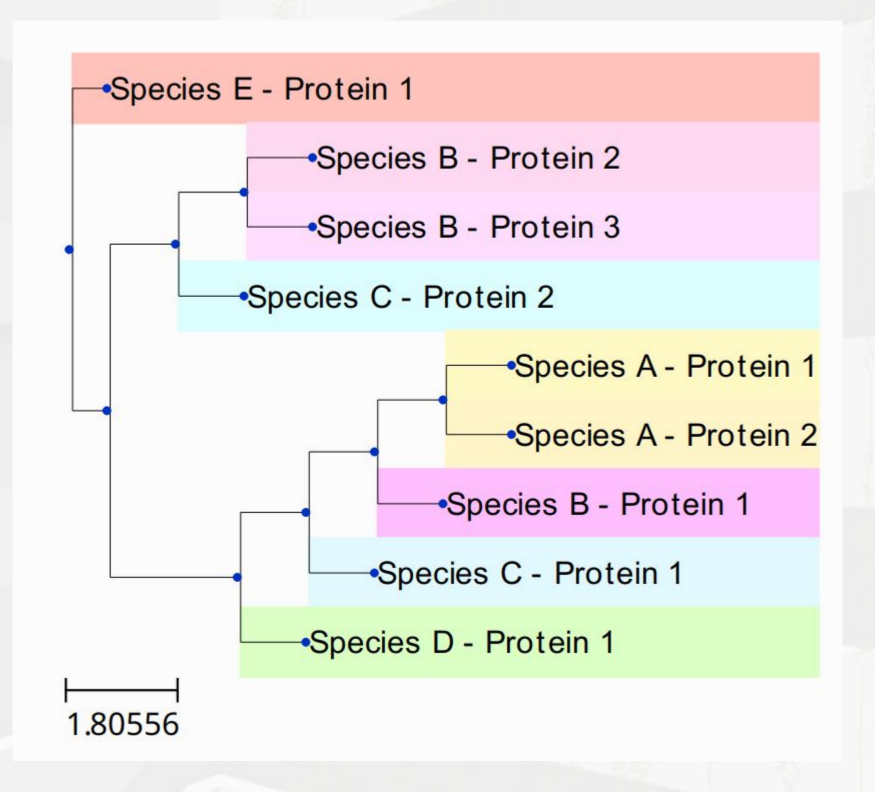
Paralogs: Sequences that come from a duplication event.



Orthogroups: Group of sequences that descend from a speciation event and can contain orthologs and in-paralogs.





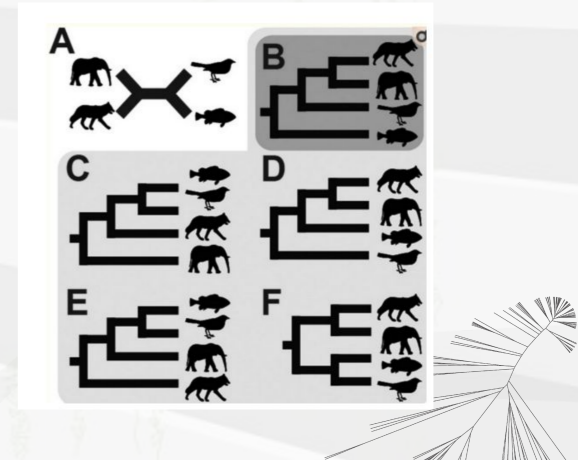


First considerations: What do you need to think about before starting.

- Species selection, specially outgroups
- Filtering of isoforms
- Fasta headers
- Computational resources

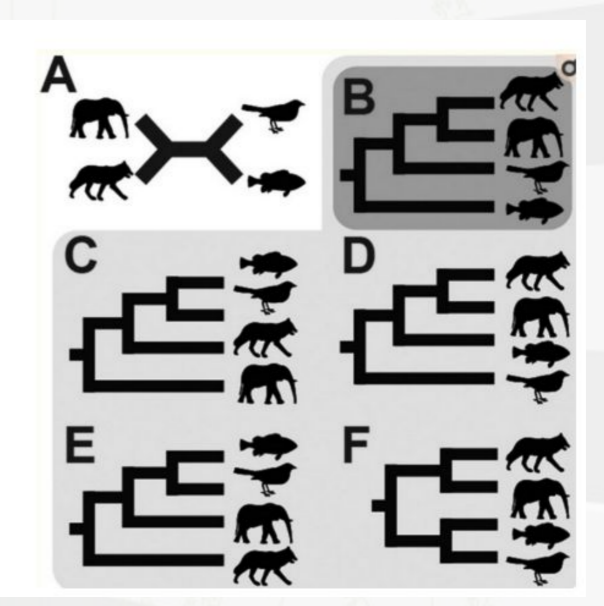
Species selection, specially outgroups

- How many species should we use?
- Genomes? Transcriptomes?
- Outgroups? How many?



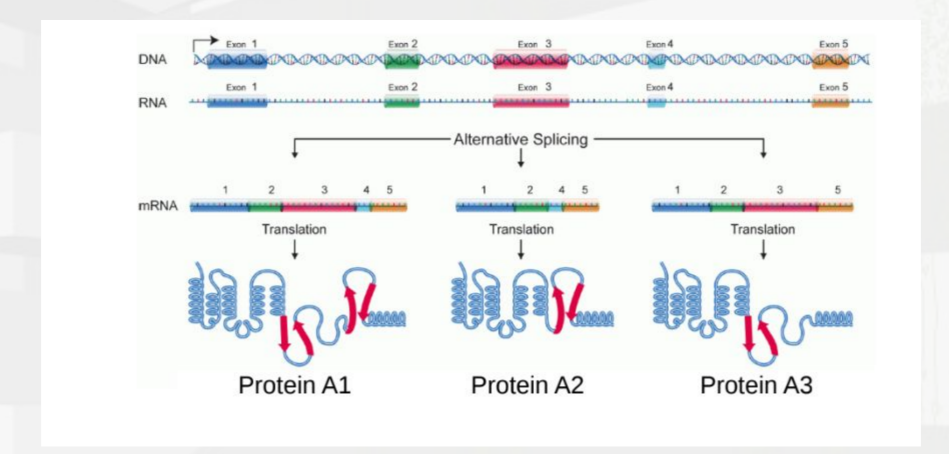
Outgroups

- When building a species tree, it is very important to use an outgroup in order to give directionality to the tree.
- Outgroups will also be necessary to root gene trees and perform orthology and paralogy predictions.
- If possible add at least two outgroups.



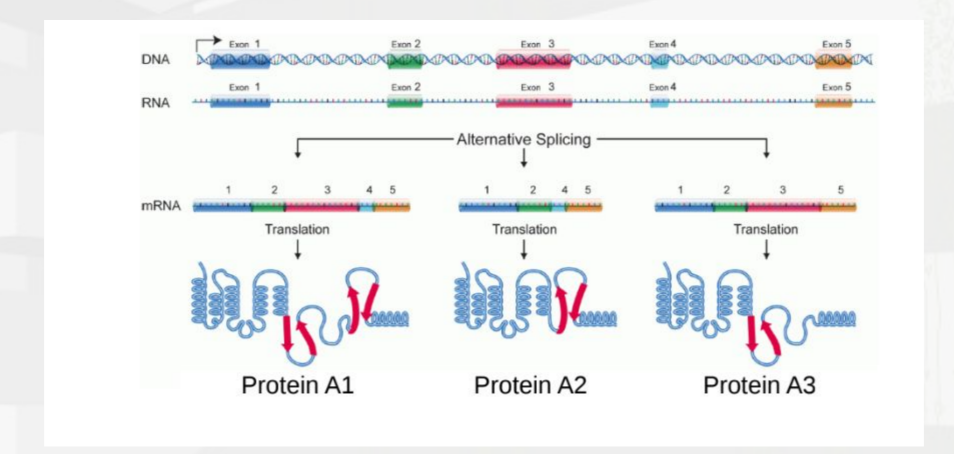


Isoforms

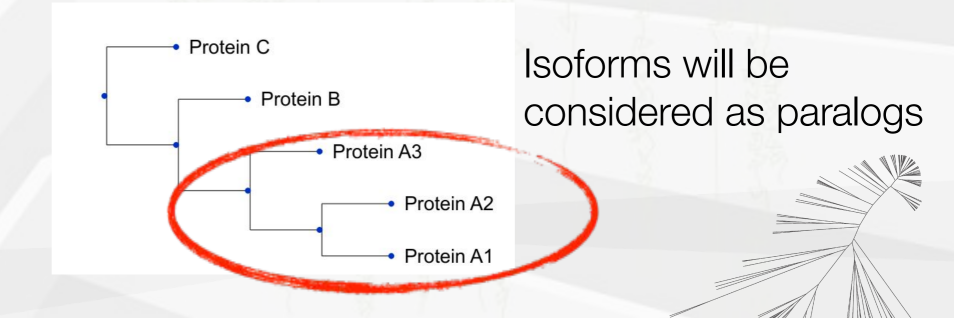


Should we add isoforms in our analysis?

Isoforms



Should we add isoforms in our analysis?



Headers

Fasta files contain headers that can be complicated. At first it will not bother you, but the downstream analysis can become much more complicated.

>sp|D2H788|RN182_AILME E3 ubiquitin-protein ligase RNF182 OS=Ailuropoda melanoleuca OX=9646 GN=RNF182 PE=3 SV=1

This is a typical Uniprot header.

Do you think it's a good idea to use it as such?



Computational resources

There are many ways to calculate orthology relationships, and some are more computationally expensive than others.

What would you use to do a homology search?

Computational resources

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Homology search: **Blast** is the tool by default, yet **Diamond** is much faster when the database is big.

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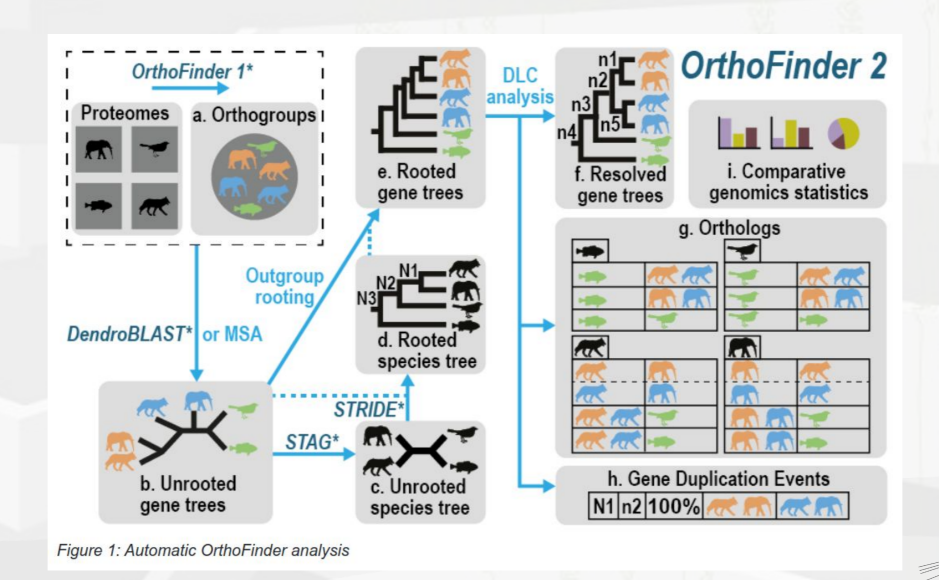
Orthology prediction: **Tree based** orthology prediction is more accurate, yet **similarity based** methods are faster.

Species selection: **More species** give more resolution, yet everything becomes more computationally expensive.

Before running an analysis always consider what you need and if you have the resources to get it.

OrthoFinder

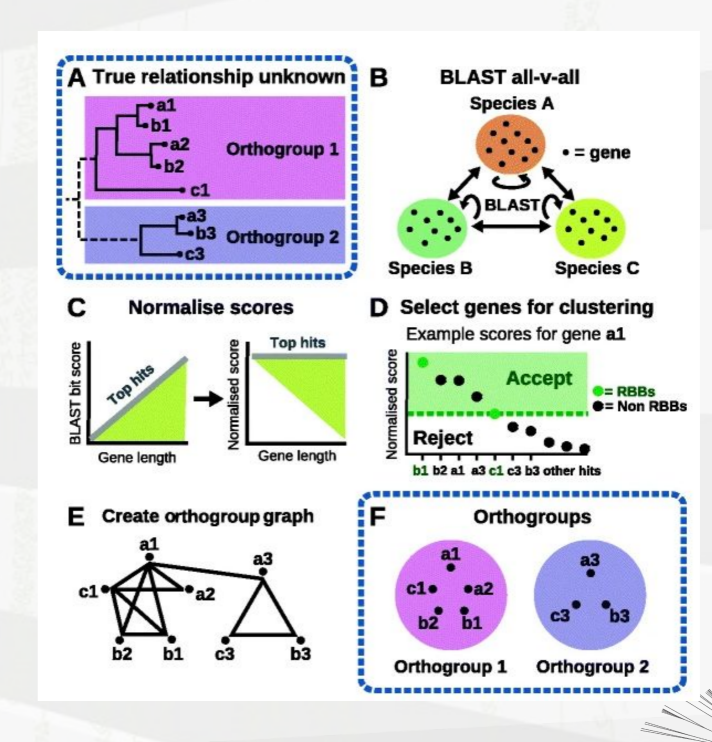
OrthoFinder is a fast, accurate and comprehensive pipeline for comparative genomics. It finds orthogroups and orthologs, infers rooted gene trees for all orthogroups and identifies all of the gene duplication events in those gene trees.



OrthoFinder

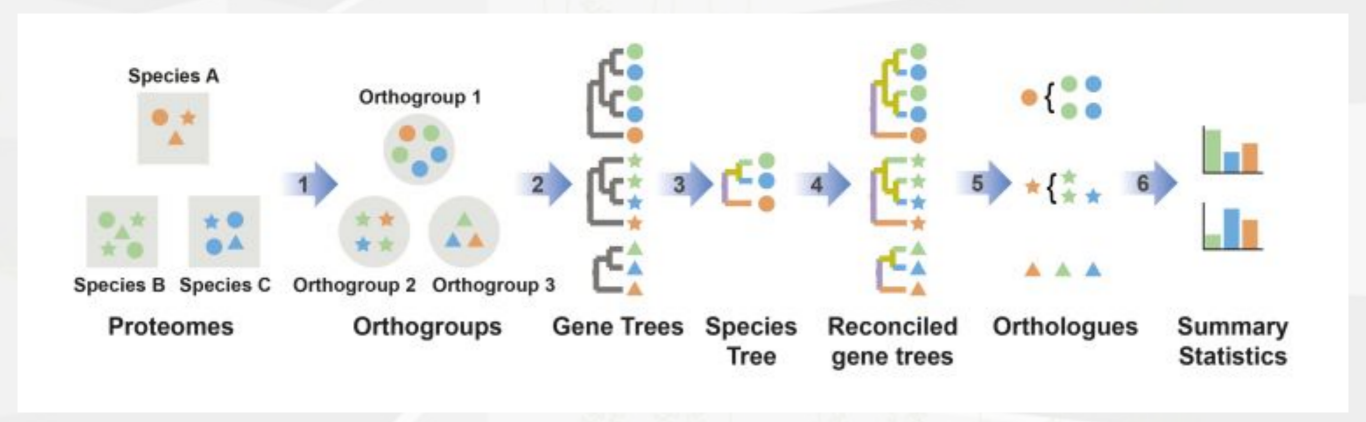
Things that Orthofinder solves compared to other algorithms:

- Bias towards gene length.
- Bias towards distantly related species.

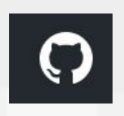


OrthoFinder

The pipeline goes from a set of proteomes to fully resolved gene trees and their orthologs and paralogs



Time for the practical!



https://github.com/ppgcourseUB/ppgcourse202 2/tree/main/Orthology