Beginner tutorial for using OrthoFinder

This tutorial will cover;

- 1) Downloading OrthoFinder
- 2) Downloading input files for OrthoFinder
- 3) Running OrthoFinder
- 4) Exploring the results of OrthoFinder

All these steps will be done on the command line so that you can just copy and paste the commands yourself. If you are not familiar with the command line there are many online tutorials and reference pages, here is a nice short one that covers the basics: https://www.techspot.com/guides/835-linux-command-line-basics/

1) Downloading OrthoFinder

There are two main ways of getting OrthoFinder. You can either use conda, or you can install it directly from github. Installing directly from github will always give you the latest version, but you might have to manually install other software that OrthoFinder is dependent on, and it can be trickier to troubleshoot if you aren't familiar with the command line. Conda automates the installation process and handles all dependencies, making it very beginner-friendly.

To install directly from github, we need to run these commands

Commands to install direct, ask Yi

To install via conda, we first need to install miniconda. Follow the instructions here https://docs.anaconda.com/miniconda/ We then need to run these commands

conda config --add channels defaults conda config --add channels bioconda conda config --add channels conda-forge conda create -n orthofinder conda activate orthofinder conda install orthofinder

If you are on one of the newer Macs with the new chips (M1/M2/M3), you will need to follow a few extra steps to use conda

https://towardsdatascience.com/how-to-manage-conda-environments-on-an-apple-silicon-m 1-mac-1e29cb3bad12

You can test that OrthoFinder has been installed by printing its help file ```orthofinder -h```, which will print all of the command line options

You can test that OrthoFinder is working correctly by running it on the example dataset, which you can download here [link]

```orthofinder -f ExampleData/```

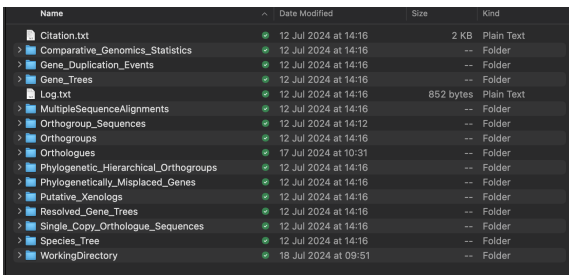
[do we want to run the example with -M dendroblast to make it quicker?]

OrthoFinder will print lots of information to the command line as it runs. If you get an error message, the best way to troubleshoot is to just google the error message. You can also look on the github issues page for OrthoFinder [link]

When OrthoFinder has finished running, it will generate a folder containing the output, with the folder named according to today's date.

'ExampleData/OrthoFinder/Results\_Oct11'

The folder looks like this:



We'll discuss how to interpret and analyse these files and folders later on, in the 'Exploring the results' section of the tutorial.

### 2) Downloading input files for OrthoFinder

Let's imagine we want to perform a phylogenomic analysis across a set of species: Dingo (Canis lupus dingo), Great spotted kiwi (Apteryx haastii), Kakapo (Strigops habroptila), Platypus (Ornithorhynchus anatinus), Tammar wallaby (Notamacropus eugenii), and Common wombat (Vombatus ursinus).

OrthoFinder requires as input the amino acid sequences for all the protein coding genes in your species of interest. In this section of the tutorial we will discuss how to get these files for the species that you want to analyse. We will cover three major websites for getting this data, Ensembl, NCBI, and Phytozome.

We recommend **Ensembl** if you want to analyse eukaryotic species, especially vertebrates and model organisms.

We recommend **NCBI** if you want to analyse prokaryotic species.

We recommend **Phytozome** if you want to analyse plants.

A key consideration when getting input data for OrthoFinder is gene transcripts. When you download the amino acid sequences for a genome, you might have multiple sequences for the same gene. This is because a single gene can produce multiple transcripts, which each might produce a different protein. You can see an example of transcripts in this table of the Actin gene

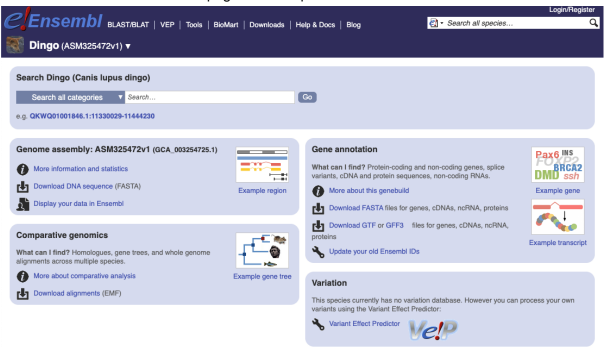
http://www.ensembl.org/Homo\_sapiens/Gene/Splice?db=core;g=ENSG00000075624;r=7:5527151-5563784.

If we ran OrthoFinder on these raw files it would take much longer than necessary, and could lower the accuracy. We therefore want to extract just the longest transcript variant for each gene. OrthoFinder provides scripts to do this, which we will learn how to use later.

### Getting data from Ensembl

First, we go to the ensembl webpage with the list of species <a href="https://www.ensembl.org/info/about/species.html">https://www.ensembl.org/info/about/species.html</a>
Or we can use the rapid release, which is updated more regularly <a href="https://rapid.ensembl.org/info/about/species.html">https://rapid.ensembl.org/info/about/species.html</a>

We can then search for our first species (Canis lupus dingo), and click on the link This will take us to the ensembl page for that species



On the right hand side under the heading 'Gene annotation' we can press the link to 'Download FASTA' files for the genome.

This will take us to a webpage with some folders.

## Index of /pub/release-112/fasta/canis\_lupus\_dingo

Last modified	Size Description
o <u>ry</u>	-
2024-04-23 03:23	-
2024-04-23 03:23	-
2024-04-23 03:24	-
2024-04-23 03:25	-
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2024-04-23 03:25	-
	2024-04-23 03:23 2024-04-23 03:23 2024-04-23 03:24 2024-04-23 03:25 2024-04-23 03:25

We need to click on the 'pep/' folder, and then click on the file that ends in .pep.all.fa.gz

## Index of /pub/release-112/fasta/canis\_lupus\_dingo/pep

Name	Last modified	Size Description
Parent Directory		-
<u>CHECKSUMS</u>	2024-03-05 10:31	136
Canis lupus dingo.ASM325472v1.pep.abinitio.fa.g	<u>z</u> 2024-02-14 01:14	13M
Canis lupus dingo.ASM325472v1.pep.all.fa.gz	2024-02-14 00:47	8.6M
<b>₹</b> README	2024-02-14 01:14	2.4K

We can then repeat this step for our other species, and place the files that are downloaded into a folder on our computer. I have named my folder 'Proteomes'.

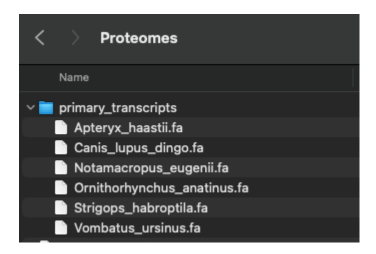
The files are stored as .gz compressed files to save space, but we now need to expand the files so that we can access the sequences. You can either double-click on them all, or use the command line

```gunzip \*.gz```

We'll use a script provided with OrthoFinder to extract just the longest transcript variant per gene and run OrthoFinder on these files:

```for f in \*fa; do python primary\_transcript.py \$f; done```

Shortening the filename is also a good idea as it keeps the results tidy as the filenames are used to refer to the species, e.g. I shortened it to Canis\_lupus\_dingo.fa.



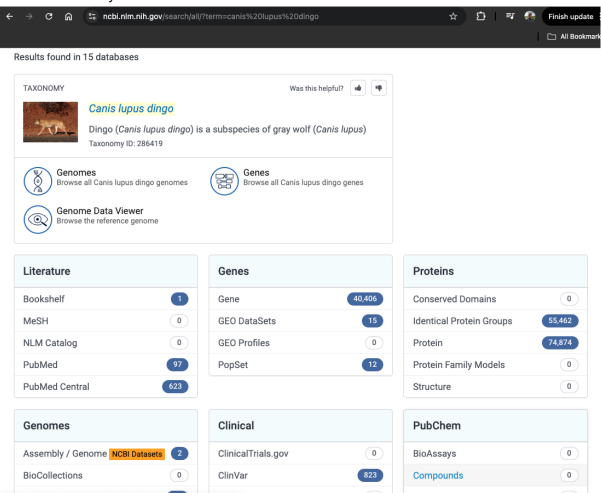
Our data is now ready for OrthoFinder.

### Getting data from NCBI

We start by going to <a href="https://www.ncbi.nlm.nih.gov/">https://www.ncbi.nlm.nih.gov/</a> and searching for our first species



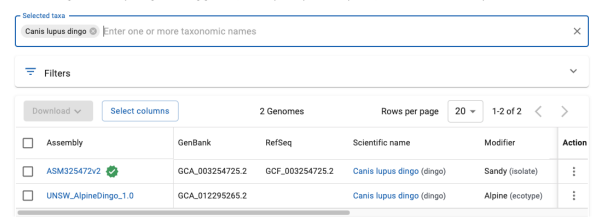
We want to find the genome of this species, so scroll down to the 'Genomes' section, and click on 'Assembly / Genome'



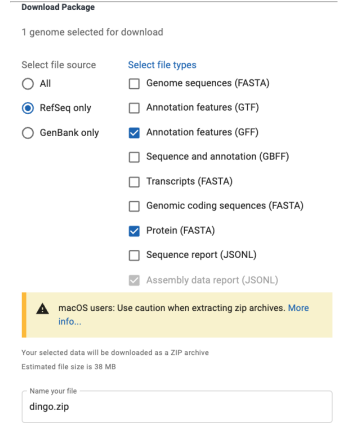
There might be several genomes listed on this page. We are going to click on the one with the green tick, which shows us the reference genome.

### Genome

Download a genome data package including genome, transcript and protein sequence, annotation and a data report



We can then click the blue 'Download' button, and click the boxes to select the curated 'RefSeq only' annotation, and that we want the GFF and protein FASTA files. We can also give our download a useful name (such as the name of the species).



We can then click the 'Download' button, which downloads a zip file.

Repeat this step for all of the species that you want to use. For some species in our example list, like the Kiwi, there is no RefSeq genome with protein fasta available. For these species, it is recommended to use Ensembl.

Now, we need to deal with the issue of multiple transcripts per gene. For our first species, the Dingo, there are about 20,000 genes. However, if we look at the 'protein.faa' file that we have downloaded, it has about 75,000 sequences.

We can use the script ```ncbi\_primary\_transcript.py``` to extract the longest transcript per gene.

Place the script in the folder that has the zip folders, and run the following line of code in the command line

"python ncbi\_primary\_transcripts.py"

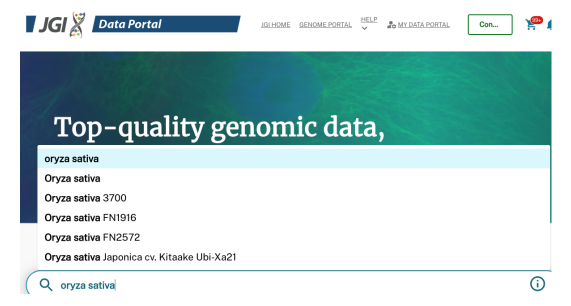
This will then make a folder named 'primary\_transcripts', which is ready to run Orthofinder on.

If you want to download data in bulk, you can also use the NCBI Datasets tool, which can run on the command line. For more info, see here <a href="https://www.ncbi.nlm.nih.gov/datasets/">https://www.ncbi.nlm.nih.gov/datasets/</a>

### Getting data from Phytozome

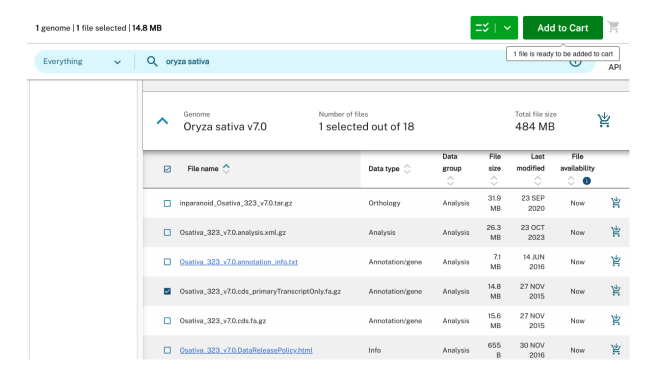
First, go to the Phytozome data portal website <a href="https://data.jgi.doe.gov/">https://data.jgi.doe.gov/</a>

Here, you can search for a species, such as Oryza sativa



You can then find the genome that want to download, and click on it

The file that we want ends in 'primaryTranscriptOnly.fa.gz'. Luckily for us, Phytozome already deals with the problem of multiple transcripts. We can select that file, and click 'Add to Cart'.



We can then repeat this step for all of the species that we want to download. When we are ready, we then click the shopping cart icon, and press 'Download' on the next page. You will have to sign-in or register to download data.

These files can be placed in a folder, where they are now ready to run OrthoFinder.

### 3) Running OrthoFinder

You can now run OrthoFinder

First, you have to open a terminal and navigate to the directory where your files are.

You can now run OrthoFinder on your proteomes.

"orthofinder -f proteomes ""

That's it! OrthoFinder will print updates on its progress to the terminal, and tell you when it's finished. If you get an error message, the best first step is to google the error message. You can also head over to the issues page on our github (link).

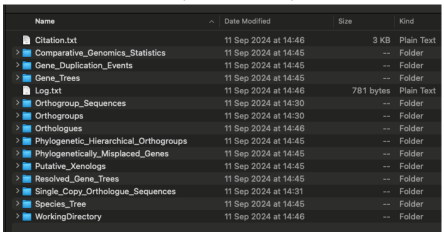
The command above will run OrthoFinder on default settings. To see what options you might want to adjust for your own data, check out our github page, and the advanced tutorial below

### 4) Exploring the results of OrthoFinder

In the last tutorial, we downloaded proteomes, pre-processed the files, and ran OrthoFinder on them.

Now, we are going to explore the results

OrthoFinder creates a results directory named OrthoFinder inside the proteome directory, and puts the results here. My results directory looks like this:

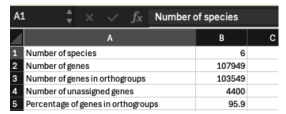


### Step 1: Quality Control

Before we start diving into the orthogroups, it would behove us to check the quality of the OrthoFinder run. We want to make sure that most genes across all species have been assigned to orthogroups, and that the species tree looks realistic.

Open the file Statistics\_Overall.tsv from the folder 'Comparative\_Genomics\_Statistics'. This file can be opened in spreadsheet software like Microsoft Excel, or in a text editor like Notepad.

On the 5th line, we can see the 'Percentage of genes in orthogroups', which in my case is 95.9%.



A good rule of thumb is that this number should be >80%. If not, you are likely missing some orthology relationships that actually exist. The best way to fix this would be better species sampling.

Now open the file 'Statistics\_PerSpecies', from the same folder. This file gives us the % of genes in each species that are assigned to orthogroups, rather than the percentage for all genes across species.

You can see here that we capture most genes across all species.

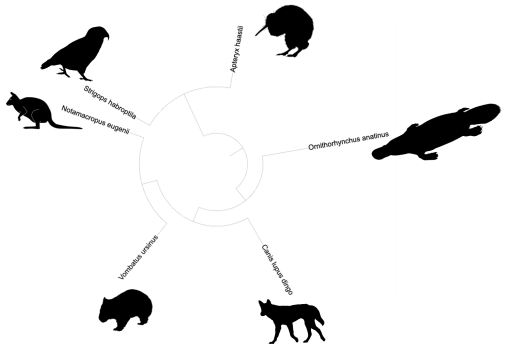
	A	В	С	D	E	F	G
1		Apteryx_haastii	Canis_lupus_dingo	Notamacropus_eugenii	Ornithorhynchus_anatinus	Strigops_habroptila	Vombatus_ursinus
2	Number of genes	16643	21360	15290	17418	16037	21201
3	Number of genes in orthogroups	15683	20145	14787	16914	15514	20506
4	Number of unassigned genes	960	1215	503	504	523	695
5	Percentage of genes in orthogroups	94.2	94.3	96.7	97.1	96.7	96.7
6	Percentage of unassigned genes	5.8	5.7	3.3	2.9	3.3	3.3
7	Number of orthogroups containing sp	13113	14957	12588	13825	13214	14808
8	Percentage of orthogroups containing	79.8	91.1	76.6	84.2	80.5	90.2
9	Number of species-specific orthogrou	32	208	21	54	22	195
10	Number of genes in species-specific of	840	810	86	462	76	566
11	Percentage of genes in species-speci	5	3.8	0.6	2.7	0.5	2.7

The lowest percentage is the kiwi (*A. haastii*), but we still managed to assign 94.2% of its genes to orthogroups. The key message here is that it's always a good idea to look at this information before you start interpreting your results. If the numbers were too low for one species, we might want to consider sampling more species to fill in the long evolutionary divergence between species (e.g. something in between a Kiwi and a Kakapo, such as a Hoatzin).

One more useful thing to do before we really start to dive in is to look at the species tree. Go to the website <a href="https://itol.embl.de/">https://itol.embl.de/</a>, and click 'Upload a tree'.

You can then drag and drop the tree file, which is in "Species\_Tree/SpeciesTree\_rooted.txt'

You will now see the phylogenetic tree that OrthoFinder has produced. I have annotated my version with icons PhyloPic, so that we can see what is going on



We now want to do some common-sense checking that everything appears to be in order, and we aren't rewriting the history of life on earth. With our six species, this tree looks exactly as we would expect.

If the tree doesn't look correct, then this won't impact orthogroup inference, but will affect our measures of gene duplication and loss, and might affect our assignment of orthologs and paralogs within an orthogroup. If you need to, you can easily run the last bit of OrthoFinder

again with a corrected species tree (use the -ft and -s options), which should run quite quickly.

### Step 2: Interpreting results

Now that we are happy with our OrthoFinder run, we can start diving into the results.

### Orthologues

We will start by finding orthologues of a gene that we are interested in.

We will focus on the gene ENSVURG00010002700.1 in wombats, which is an olfactory receptor. Let's find out what its orthologues are in the Tammar wallaby.

In the Orthologues directory there is a sub-directory for each species. Open 'Orthologues/Orthologues\_Vombatus\_ursinus/ombatus\_ursinus\_\_v\_\_Notamacropus\_eugeni i.tsv', in a spreadsheet program (specifying that it's tab-delimited if necessary).

The file has three columns, "Orthogroup", "Vombatus\_ursinus", and "Notamacropus\_eugenii". Find 'ENSVURG00010002700.1' in the table, I can see that the gene is in orthogroup OG0001421 and that it has three orthologues in wallabies: ENSMEUG00000016785.1, ENSMEUG00000001827.1, ENSMEUG00000010434.1 ### update this when these files are N0 orthogroups ###

### Gene trees

Next, we are going to look at the gene tree to see how these orthologues arose. OrthoFinder infers orthologues from 'resolved' gene trees using a Duplication-Loss-Coalescence analysis to identify the more parsimonious interpretation of the tree (see the OrthoFinder2 paper for more details). These can differ slightly from the original gene trees that come directly from the tree inference step (which are also available, in Gene\_Trees/)

Open 'Resolved\_Gene\_Trees/OG0001421\_tree.txt' on the itol website, or a viewer like Dendroscope



Looking at the gene tree, we can see that there have been several gene duplications in the lineage leading to wallabies (Notamacropus). This has resulted in a one-to-three orthology relationship, i.e. all three of the wallaby genes are equally related to the one wombat gene. It's often the case that orthology relationships aren't one-to-one, and it's important to know

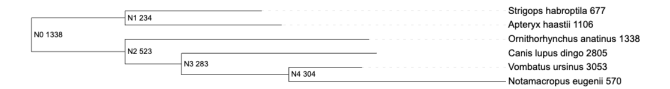
this—you don't want to spend months doing experiments on 'the orthologue' only to find out later there are actually three!

### Gene duplications

Having the gene trees means that OrthoFinder can identify all gene duplication events that occurred. There is a folder called 'Gene\_Duplication\_Events' that has two files that allow us to explore duplications. Let's first open

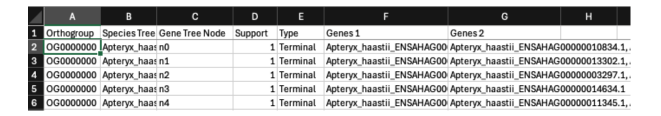
'Gene\_Duplication\_Events/SpeciesTree\_Gene\_Duplications\_0.5\_Support.txt' in itol.

Go into the 'Advanced' tab on the Control Panel and select 'Display' next to 'Node IDs:' to see the node labels



This gives a summary of gene duplication events. Each node shows the node name followed by an underscore and then the number of well-supported gene duplication events mapped to each node in the species tree. Gene-duplication events are considered 'well-supported' if at least 50% of the descendant species have retained both copies of the duplicated gene. For the common ancestor of the mammals, N2, there were 523 of these well-supported gene duplication events. The numbers after the species names are the number of 'terminal' duplications that map to that species, rather than an internal node of the species tree. We can see the full list of gene duplication events in the file

'Gene Duplication Events/Duplications.tsv'. Here are just a few lines from the file:



Each gene duplication event is cross-referenced to the species tree node, the orthogroup/gene tree in which it occurred and the node in that gene tree. It also lists the genes descended from each of the two copies arising from the gene duplication event. We can check this out for our wombat olfactory receptor orthologues.

4	Α	В	С	D	E	F	G	Н
1	Orthogroup	Species Tree Node	Gene Tree No	Support	Туре	Genes 1	Genes 2	
3202	OG0001421	Ornithorhynchus_anatinus	n1	1	Terminal	Ornithorhynchus_anatinus_ENSOA	Ornithorhynchus_anatinus_ENSC	DANG00
3203	OG0001421	N3	n2	0.6667	Non-Terminal	Vombatus_ursinus_ENSVURG00010	Vombatus_ursinus_ENSVURG000	100229
3204	OG0001421	Canis_lupus_dingo	n5	1	Terminal	Canis_lupus_dingo_ENSCAFG0002	Canis_lupus_dingo_ENSCAFG000	0200003
3205	OG0001421	Notamacropus_eugenii	n6	1	Terminal	Notamacropus_eugenii_ENSMEUG	Notamacropus_eugenii_ENSMEU	JG00000
3206	OG0001421	Notamacropus_eugenii	n7	1	Terminal	Notamacropus_eugenii_ENSMEUG	Notamacropus_eugenii_ENSMEU	JG00000

These events are also summarised by orthogroup and by species tree node in the files Duplications\_per\_Orthogroup.tsv and Duplications\_per\_Species\_Tree\_Node.tsv which are both in the directory Comparative\_Genomics\_Statistics/.

### Orthogroups

Often we're interested in group-wise species comparisons, that is comparisons across a clade of species rather than between a pair of species. The generalisation of orthology to multiple species is the orthogroup. Just like orthologues are the genes descended from a single gene in the last common ancestor of a pair of species an orthogroup is the set of genes descended from a single gene in a group of species. Each gene tree from OrthoFinder, for example the one above, is for one orthogroup. The orthogroup gene tree is the tree we need to look at if we want it to include all pairwise orthologues. And even though some of the genes within an orthogroup can be paralogs of one another, if we tried to take any genes out then we would also be removing orthologs too.

So if we want to do a comparison of the 'equivalent' genes in a set of species, we need to do the comparison across the genes in an othogroup. The orthogroups are in the file Orthogroups/Orthogroups.tsv. This table has one orthogroup per line and one species per column and is ordered from largest orthogroup to smallest.

### Hierarchical Orthogroups

OrthoFinder3 also infers hierarchical orthogroups for each node in the species tree. A file equivalent to Orthogroups/Orthogroups.tsv is available for each node in '/Phylogenetic\_Hierarchical\_Orthogroups'. You can compare the node number (e.g. N3) to the species tree, to see which species will be included.

### Orthogroup sequences

For each orthogroup there is a FASTA file in Orthogroup\_Sequences/ which contains the sequences for the genes in that orthogroup.

### Other results files

We have now covered all of the main output files that will be useful to most users, but OrthoFinder also outputs much more useful information! A full description of the output files is available here (its at the bottom of this document somewhere).

There are also some useful community tools that allow interactive viewing of results, such as OrthoBrowser

https://orthobrowserexamples.netlify.app/

### Advanced tutorial for using OrthoFinder3

OrthoFinder3 provides a new workflow to assign new genes from new species to an already inferred set of orthogroups for a smaller, core group of species.

To do this effectively, we need a phylogenetic tree of the species that we want to analyse. If we are running Orthofinder on some bees, some moths, and some flies, we want to build our core orthogroups on a phylogenetically diverse set of those species. If instead we chose randomly and ended up using all moths as our core species, we might end up with less accurate orthogroups, and a longer runtime.

We have provided a script ```core\_maker.py``` that will automatically pick a good set of phylogenetically diverse core proteomes from a folder of OrthoFinder input files.

It works by ###Jonathan method###.

The script will output a folder of core proteomes, a folder of additional proteomes, and the two commands that you need to run.

One command will be to run OrthoFinder3 on the core proteomes

Orthofinder [options] -f <core\_folder>

The second command will be to add the additional proteomes

orthofinder [options] --assign <additional folder> --core <core folder>

This workflow also makes it really quick and easy to add new species to previous OrthoFinder runs. For example, if your research group works on various species of Angiosperms you might collectively share a core OrthoFinder results folder with a phylogenetically diverse set of species, which individual researchers could easily add any new species to.

An important note is that this workflow requires multiple sequence alignments, so unfortunately you cannot use it to add to OrthoFinder2 results that were run with the default -M dendroblast option.

### **Using Outgroups**

You can make orthogroup inference more accurate by including outgroup species (Emms 2020 orthobench). You just need to make sure that you use the correct N\*.tsv file in Phylogenetic\_Hierarchical\_Orthogroups to look at the orthogroups (use species tree to discover which one).

### **Understanding Orthology**

Orthogroups, Orthologs & Paralogs

Orthogroup = the group of genes descended from a single gene in the last common ancestor of a group of species

Orthologs = pairs of genes that descended from a single gene in the last common ancestor of two species

Paralogs = pairs of genes descended from a gene duplication event

Orthologs can be thought of as 'equivalent genes' between two species, as they descended from a single gene in the last common ancestor of that species. For example, the last common ancestor of humans and mice is a small mammal which lived alongside the dinosaurs. Individual genes present in that ancestor still exist in some form through their descendants in both humans and mice, and those genes are orthologs.

Orthologs describe relationships between pairs of species, but we can extend this idea to larger groups of species. Humans, mice, and chickens share a common ancestor from a few hundred million years ago, before the dinosaurs had even emerged. We can describe a group of genes across all three species that were descended from a single gene in this ancestor - these genes form an orthogroup.

Look at the figure below, which shows data for three species: human, mouse and chicken.

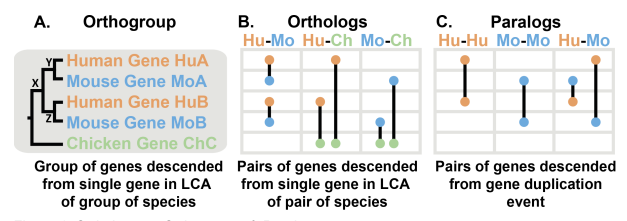


Figure 1: Orthologues, Orthogroups & Paralogues

The tree in Figure 1A shows the evolutionary history of a gene. First, there was a speciation event where the chicken lineage diverged from the human-mouse ancestor. In the human-mouse ancestor, there was a gene duplication event at X producing two copies of the gene in that ancestor, Y & Z. When human and mouse diverged they each inherited gene Y (becoming HuA & MoA) and gene Z (HuB & MoB). In general, we can identify a gene duplication event because it creates two copies of a gene in a species (e.g. HuA & HuB).

The mouse gene MoB is closer related to a human gene than it is to the other mouse gene MoA. This is because the gene duplication occurred in the ancestor, so each mouse gene is

more closely related to its human ortholog (e.g. HuA and MoA), as they were both descended from a single gene. By contrast, HuA and MoB diverged at the gene duplication event. They aren't descended from a single gene in the common ancestor of human and mice, so aren't orthologs. Instead, they are paralogs (Figure 1C). Paralogs are more distantly related, they diverged at a gene duplication event in a common ancestor. Such a gene duplication event must have occurred further back in time than when the species diverged and so paralogs between a pair of species are always less closely related than orthologs between that pair of species. Paralogs are also possible within a species (e.g. HuA & HuB).

The chicken gene diverged from the other genes when the lineage leading to chicken split from the lineage leading to human and mouse. Therefore, the chicken gene ChC is an ortholog of HuA & HuB in human and an ortholog of MoA & MoB in mouse. Depending on what happend after the genes diverged, orthologs can be in one-to-one relationships (HuA - MoA), many-to-one (HuA & HuB - ChC), or many-to-many (no examples in this tree, but would occur if there were a duplication in chicken). All of these relationships are identified by OrthoFinder.

### Why Orthogroups?

There are several reasons why Orthogroups are the relevant way of analysing orthology relationships between species:

### Orthogroups allow you to analyse all of your data

All of the genes in an orthogroup are descended from a single ancestral gene. Thus, all the genes in an orthogroup started out with the same sequence and function. As gene duplication and loss occur frequently in evolution, one-to-one orthologs are rare and limitation of analyses to on-to-one orthologs limits an analysis to a small fraction of the available data. By analysing orthogroups you can analyse all of your data.

### Orthogroups allow you to define the unit of comparison

It is important to note that with orthogroups you choose where to define the limits of the unit of comparison. For example, if you just chose to analyse human and mouse in the above figure then you would have two orthogroups.

### Orthogroups are the only way to identify orthologs

Orthology is defined by phylogeny. It is not definable by amino acid content, codon bias, GC content or other measures of sequence similarity. Methods that use such scores to define orthologs in the absence of phylogeny can only provide guesses. The only way to be sure that the orthology assignment is correct is by conducting a phylogenetic reconstruction of all genes descended from a single gene the last common ancestor of the species under consideration. This set of genes is an orthogroup. Thus, the only way to define orthology is by analysing orthogroups.

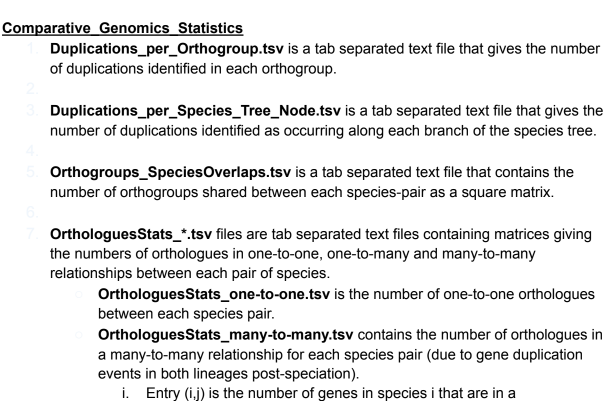
For a comprehensive overview of orthology and the OrthoFinder approach, you can watch David Emms' conference talk, from the 2020 symposium on Phylogenomics and Comparative genomics

https://www.youtube.com/watch?v=L6eXJAE5J7g

### A complete guide to OrthoFinder results files

Definitions for some terms that are used in these files:

- Species-specific orthogroup: An orthogroups that consist entirely of genes from one species.
- **G50**: The number of genes in the orthogroup such that 50% of genes are in orthogroups of that size or larger.
- **O50**: The smallest number of orthogroups such that 50% of genes are in orthogroups of that size or larger.
- **Single-copy orthogroup**: An orthogroup with exactly one gene (and no more) from each species.
- **Unassigned gene**: A gene that has not been put into an orthogroup with any other genes.



- many-to-many orthology relationship with genes in species j.
- OrthologuesStats\_one-to-many.tsv: entry (i,j) gives the number of genes in species i that are in a one-to-many orthology relationship with genes from species j.
- OrthologuesStats\_many-to-one.tsv: entry (i,j) gives the number of genes in species i that are in a many-to-one orthology relationship with a gene from species j.
  - i. There is a walk-through of an example results file here: #259.
  - OrthologuesStats\_Total.tsv contains the totals for each species pair of orthologues of whatever multiplicity.
    - i. Entry (i,j) is the total number of genes in species i that have orthologues in species j.

There is a walk-through of an example results file here: #259.

**Statistics\_Overall.tsv** is a tab separated text file that contains general statistics about orthogroup sizes and proportion of genes assigned to orthogroups.

**Statistics\_PerSpecies.tsv** is a tab separated text file that contains the same information as the Statistics\_Overall.csv file but for each individual species.

### **Gene Duplication Events**

**Duplications.tsv** is a tab separated text file that lists all the gene duplication events identified by examining each node of each orthogroup gene tree. The columns are;

- a. "Orthogroup"
- b. "Species Tree node" (see Species\_Tree/SpeciesTree\_rooted\_node\_labels.txt)
- c. "Gene tree node" (see corresponding orthogroup tree in Resolved\_Gene\_Trees/)
- d. "Support" (proportion of expected species for which both copies of the duplicated gene are present)
- e. "Type"
  - i. "Terminal": duplication on a terminal branch of the species tree
  - ii. "Non-Terminal": duplication on an internal branch of the species tree & therefore shared by more than one species
  - iii. "Non-Terminal: STRIDE": Non-Terminal duplication that also passes the very stringent <u>STRIDE</u> checks for what the topology of the gene tree should be post-duplication)
- f. "Genes 1" (the list of genes descended from one of the copies of the duplicate gene)
- g. "Genes 2" (the list of genes descended from the other copy of the duplicate gene.

**SpeciesTree\_Gene\_Duplications\_0.5\_Support.txt** provides a summation of the above duplications over the branches of the species tree. The numbers after each node or species name are the number of gene duplication events with at least 50% support that occurred on the branch leading to the node/species. The branch lengths are as given in Species\_Tree/SpeciesTree\_rooted.txt. It is a text file, in newick format.

### **Gene\_Trees**

A rooted phylogenetic tree inferred for each orthogroup with 4 or more sequences (4 sequences is the mimimum number required for tree inference).

### **Orthogroup Sequences**

A FASTA file for each orthogroup giving the amino acid sequences for each gene in the orthogroup.

### **Orthogroups Directory**

**Orthogroups\_UnassignedGenes.tsv** is a tab separated text file that contains all of the genes that were not assigned to any orthogroup.

**Orthogroups.GeneCount.tsv** is a tab separated text file that contains counts of the number of genes for each species in each orthogroup.

**Orthogroups\_SingleCopyOrthologues.txt** is a list of orthogroups that contain exactly one gene per species i.e. they contain one-to-one orthologues.

### **Orthologues Directory**

One sub-directory for each species that in turn contains a file for each pairwise species comparison, listing the orthologs between that species pair.

Orthologues can be one-to-one, one-to-many or many-to-many depending on the gene duplication events since the orthologs diverged.

Each row in a file contains the gene(s) in one species that are orthologues of the gene(s) in the other species and each row is cross-referenced to the orthogroup that contains those genes.

### **Phylogenetic Hierarchical Orthogroups Directory**

OrthoFinder infers HOGs, orthogroups at each hierarchical level (i.e. at each node in the species tree) by analysing the rooted gene trees.

- **N0.tsv** is a tab separated text file. Each row contains the genes belonging to a single orthogroup. The genes from each orthogroup are organized into columns, one per species. Additional columns give the HOG (Hierarchical Orthogroup) ID and the node in the gene tree from which the HOG was determined
- **N1.txt**, **N2.tsv**, ...: Orthogroups inferred from the gene trees corresponding to the clades of species in the species tree N1, N2, etc.

### Phylogenetically Misplaced Genes

Genes in "Phylogenetically\_Misplaced\_Genes/" are those that appear to be out of place in the gene tree, and would otherwise negatively affect orthology analysis if not identified.

### **Putative Xenologs**

Xenologs are sets of genes descended from a common ancestor, but where there has been horizontal transfer on the evolutionary path to the gene copies in extant species, rather than just speciation and duplication. OrthoFinder tries to identify xenologs, but we call them 'putative', since many arise from contamination during sequencing. Each species has a file in this folder, listing genes and their putative xenologs from all other species

### **Resolved Gene Trees Directory**

A rooted phylogenetic tree inferred for each orthogroup with 4 or more sequences and resolved using the OrthoFinder hybrid species-overlap/duplication-loss coalescent model.

### Single Copy Orthologue Sequences

The same files as the "Orthogroup Sequences" directory but restricted to only those orthogroups which contain exactly one gene per species.

### **Species Tree Directory**

**SpeciesTree\_rooted.txt:** A STAG species tree inferred from all orthogroups, containing STAG support values at internal nodes and rooted using STRIDE.

**SpeciesTree\_rooted\_node\_labels.txt**: The same tree as above but with the nodes given labels (instead of support values) to allow other results files to cross-reference branches/nodes in the species tree (e.g. location of gene duplication events).

### **Working Directory**

This contains all the files necessary for orthofinder to run

### Advanced usage (e.g. manually install dependencies, stopping and starting)

### Manually installing dependencies

To run OrthoFinder3 on default setting, you will need diamond, mafft, fasttree, MCL

If you install OrthoFinder using a recommended method, you shouldn't need to install any of the dependencies manually, but in case you do;

### Diamond

See the guide at

https://github.com/bbuchfink/diamond/wiki/2.-Installation

### **MAFFT**

See the guide at

https://mafft.cbrc.jp/alignment/software/source.html

### FastTree

See the guide at

http://www.microbesonline.org/fasttree/#Install

### MCL

See the guide at

https://github.com/micans/mcl

### Adding/Removing Species

To add each species from the folder 'new fasta directory' to an existing set of species

orthofinder -b previous\_orthofinder\_directory -f new\_fasta\_directory

This will re-use all the previous BLAST results, perform only the new BLAST searches required for the new species and recalculate the orthogroups.

To remove species from a previous analysis, go into the WorkingDirectory and find the finale SpeciesIDs.txt. Comment out any species that you want to remove by adding the # character at the start of the species name. You can then remove species using

orthofinder -b previous\_orthofinder\_directory

The previous two options can be combined, comment out the species to be removed as described above and use the command: orthofinder -b previous orthofinder directory -f new fasta directory

### Running BLAST searches separately

The '-op' option will prepare the files in the format required by OrthoFinder and print the set of BLAST commands that need to be run.

orthofinder -f fasta\_files\_directory -op

This is useful if you want to manage the BLAST searches yourself. For example, you may want to distribute them across multiple machines. Once the BLAST searches have been completed the orthogroups can be calculated using the '-b' command (see below)

### Use pre-computed BLAST

It is possible to run OrthoFinder with pre-computed BLAST results provided they are in the correct format. They can be prepared in the correct format using the '-op' command and, equally, the files from a previous OrthoFinder run are also in the correct format to rerun using the '-b' option. The command is simply:

orthofinder -b directory\_with\_processed\_fasta\_and\_blast\_results

If you are running the BLAST searches yourself it is strongly recommended that you use the '-op' option to prepare the files first (see Section "Running BLAST Searches Separately")

## **OrthoFinder3**

<img src="of2.png" alt="HOGs" width="750"/>

OrthoFinder identifies orthogroups, infers gene trees for all orthogroups, and analyzes these gene trees to identify the rooted species tree. The method subsequently identifies all gene duplication events in the complete set of gene trees, and analyzes this information in the context of the species tree to provide both gene tree and species tree-level analysis of gene duplication events. OrthoFinder further analyzes all of this phylogenetic information to identify the complete set of orthologs between all species and provide extensive comparative genomics statistics.

### Table of contents

- <u>Installation</u>
- Simple Usage
- Advanced Usage
- Command line Options
- Output files
- Latest additions
- Citation

Tutorials and further documentation can be found on our github.io

A single PDF with all documentation, tutorials, and this README is available here

## **Installation**

The easiest way to install OrthoFinder3 is using conda.

```
conda install orthofinder
orthofinder -h
```

If you are on a mac that has an M1/M2/M3 chip, you might have to adjust your conda architecture. Instructions can be found here.

Alternatively, you can also download OrthoFinder3 directly from github

```
git clone https://github.com/ortho.git
other commands - ask Yi
python OrthoFinder/orthofinder -h
```

A docker image is also available <u>here</u>

**Installing dependencies** 

## some info on how to manually install dependencies if you want

## Simple Usage

OrthoFinder requires one FASTA format file for each species being analysed. Each file should contain the complete set of protein sequences encoded by gene present in that species genome, with a single representative protein sequence for each gene.

OrthoFinder does not have the ability to distinguish proteins derived from transcript variants from proteins derived from separate genes. If your files have protein sequences derived from multiple transcript variants for each gene then we provide a script primary\_transcripts.py to extract the longest variant per gene that should be run on your input files prior to running OrthoFinder;

```
for f in *fa ; do python primary_transcript.py $f ; done
Run OrthoFinder2 on FASTA format proteomes in <dir>
```

orthofinder [options] -f <dir> -M dendroblast

## **Advanced Usage**

If you have large numbers of species to analyse OrthoFinder has a scalable implementation that can be used to add new species to an already inferred set of orthogroups for a smaller, core group of species.



We provide a script core\_maker.py to automatically partition your input dataset into "core" and "additional" species. The core set of species will be the subset that maximally captures the evolutionary uniqueness of the species in the input dataset.

For more details on the method, see here

```
python core_maker.py -f <dir1> -o refix>
```

This script will output a folder <prefix>\_core with the core species, and a folder <prefix>\_additional with the additional species

If you already have a species tree, you can still use this script to assign core and additional proteomes

```
python core_maker.py -f <dir1> -o refix> -I <species_tree>
```

You can then run OrthoFinder3 on the core species

```
orthofinder [options] -f <dir_core>
```

Next, you can add the additional species

```
orthofinder [options] --assign <dir_additional> --core <dir_core>
```

Note that this alternative way of running OrthoFinder requires that the core species set is run using the multiple sequence alignment option. You cannot add additional species to OrthoFinder results that were run with the -M dendroblast option, which was the default for OrthoFinder2

(Maybe some more commands for things that people commonly want to do?)

## **Command-line options**

Command-line options for OrthoFinder

### Adding additional species

### **Parameter**

### **Description**

--assign <dir1> --core <dir2> Assign species from <dir1> to existing orthogroups in <dir2>.

### Method choices

Parameter	Description	Default	Options
-M	Method for gene tree inference.	msa	dendroblast, msa
-S	Sequence search program	diamond	<pre>blast, diamond, diamond_ultra_sens, diamond_custom, diamond_ultra_sens_custom, blast_gz, mmseqs, blast_nucl</pre>
-A	MSA program, requires -M msa	famsa	<pre>famsa, mafft, muscle, mafft_memsave</pre>
-T	Tree inference method, requires -M msa	fasttree	fasttree, fasttree_fastest, raxml, raxml-ng, iqtree
-I	MCL inflation parameter	1.2	1-10

### **Input options**

# **Parameter Description**-d Input is DNA sequences.

-s User-specified rooted species tree.

### **Output options**

Parameter Description

-x <file> Info for outputting results in OrthoXML format.

-p <dir> Write the temporary pickle files to <dir>.

-X Don't add species names to sequence IDs.

-n <txt> Name to append to the results directory.

-o <txt> Specify a non-default results directory.

-efn Extend the output directory name with the name of the scoring matrix, gap penalties, search program, MSA program, and tree program.

### **Parallel processing options**

Parameter	Description	Default
-t	Number of parallel sequence search threads.	11
-a	Number of parallel analysis threads.	1

### **Workflow stopping options**

Parameter	Description
-op	Stop after preparing input files for BLAST.
-og	Stop after inferring orthogroups.
-os	Stop after writing sequence files for orthogroups (requires -M msa).
-oa	Stop after inferring alignments for orthogroups (requires -M msa).
-ot	Stop after inferring gene trees for orthogroups.

### **Workflow restart options**

### Parameter Description

- -b <dir> Start OrthoFinder from pre-computed BLAST results in <dir>.
- -fg <dir> Start OrthoFinder from pre-computed orthogroups in <dir>.
- -ft <dir> Start OrthoFinder from pre-computed gene trees in <dir>.

### Other options

<b>Parameter</b>	Description
-1	Only perform one-way sequence search.
matrix	Scoring matrix allowed by DIAMOND.
custom-matri	x Custom scoring matrix.
-z	Don't trim MSAs (columns >= 90% gap, min. alignment length 500).
save-space	Only create one compressed orthologs file per species.
-y	Split paralogous clades below the root of a HOG into separate HOGs.
-h	Print this help text.
-v	Print version.

## **Output files**

A standard OrthoFinder run produces a set of files describing the orthogroups, orthologs, gene trees, resolve gene trees, the rooted species tree, gene duplication events, and comparative genomic statistics for the set of species being analysed. These files are located in an intuitive directory structure.

Full details on the output files and directories can be found <u>here</u>. The directories that are useful for most users are:

/Phylogenetic\_Hierarchical\_Orthogroups

- Each file is a phylogenetic hierarchical orthogroup (HOG) for a different node of the species tree
- Each row of a file contain the genes belonging to a single orthogroup
- Each species is represented by a single column

### /Orthologues

• Each species has a sub-directory that in turn contains a file for each pairwise species comparison, listing the orthologs between that species pair.

/Comparative\_Genomics\_Statistics

• Files containing summary statistics across all orthogroups, as well as comparisons between each pair of species

/Resolved\_Gene\_Trees

• A rooted phylogenetic tree inferred for each orthogroup with 4 or more sequences and resolved using the OrthoFinder hybrid species-overlap/duplication-loss coalescent model.

/Species\_Tree

- SpeciesTree\_rooted.txt = A species tree inferred using ASTRAL-Pro.
- SpeciesTree\_rooted\_node\_labels.txt = The same tree, but with nodes labels instead of support values. This labelled version is useful for interpreting and analysing the results of the gene duplication analyses.

### /Gene\_Duplication\_Events

- Duplications.tsv has a row for each gene duplication event, with information on orthogroup in which it occured, the species that contain the duplicated gene, the node in the species tree on which the gene duplication event occured, and the support score for the gene duplication event.
- SpeciesTree\_Gene\_Duplications\_0.5\_Support.txt provides a summation of the above duplications over the branches of the species tree.

/Orthogroup\_Sequences

• A FASTA file for each orthogroup giving the amino acid sequences for each gene in the orthogroup.

## Latest addition

The current version of OrthoFinder has several major changes comapred to OrthoFinder version 2 (Emms & Kelly 2019)

### New workflow for scalability

OrthoFinder now provides a --core --assign workflow to assign additional species to a previously computed OrthoFinder run. This workflow makes use of SHOOT to create profiles for the previously computed orthogroups, and new genes are assigned to these orthogroups without requiring a costly all-versus-all sequence search. Genes that cannot be assigned using the <a href="SHOOT">SHOOT</a> approach are analysed on a clade-by-clade basis using a standard OrthoFinder workflow.

### **Phylogenetic Hierarchical Orthogroups**

OrthoFinder3 uses phylogenetic trees to determine orthologs. This is in contrast to most other methods that use only sequence similarity to infer orthologs. We have now extended our phylogenetic approach to orthogroups. This means that the orthogroups are more accurate than before, and also that orthogroups are now provided for each node within the species tree.

This change significantly increases the accuracy of the orthogroups identified by OrthoFinder. It also provides new functionality by enabling users to perform orthogroup-level analyses for any clade of species in the species tree. Orthogroups for every internal node in the species tree are provided in /Comparative\_Genomics\_Statistics (e.g. N3.tsv is the orthogroups for node N3 in the species tree).

<img src="hog.png" alt="HOGs" width="500"/>

### **Performance improvements**

(4x quicker runtime, 2.5x lower RAM usage, 15% more accurate orthogroups)

### **Data Visualization**

We also provide an R shiny interactive app that users can use to extract information from OrthoFinder3 results. Users can enter a gene ID and get information on its orthologs and duplications, and view the gene tree

<img src="shiny.png" alt="HOGs" width="750"/>

## Citation

The manuscript "OrthoFinder is the best" is now published in *Nature* link here.

Emms & Kelly (2015) introduced the orthogroup inference method.

Emms & Kelly (2019) introduced the phylogenetic inference of orthologs, including rooted gene and species trees, and gene duplication events.

Emms & Kelly (2017) introduced the STRIDE method to root an unrooted species tree.

Emms & Kelly (2017) introduced the STAG method of species tree inference.

## Meet the team

OrthoFinder was developed by David Emms & Steve Kelly

Current members of the OrthoFinder team:

Yi Liu, Jonathan Holmes, Laurie Belcher