#### Goals

- Get a chance to process dataset to explore genome evolutionary questions
- Practice creating graphical plots of data in R using existing templates of scripts
- Get a feel for some of the datatypes and datasets you might work with

### Get the Tutorial

```
$ git clone https://github.com/hyphaltip/Tutorial GenomeEvolution
$ cd data
$ bash download.sh # will download datasets
$ cd ..
$ Rscript scripts/install pkg.R
# run the 3 tutorial scripts to see plots
# generate plots/pheatmap example.pdf
$ Rscript scripts/plot heatmap family.R
# generate genome summary plots plots/chrom features.pdf
# and plots/summary stats.pdf
$ Rscript scripts/plot chroms 1.R
# summarize single-copy A.fumigatus orthologs only from
# the larger Orthologs table
$ ./scripts/extract orthologs single-copy Afum.py
```

## Install some R packages

- May need to install extra libraries, in R:
  - install.packages("ggplot2","gridExtra","dplyr","RColorBrew er","pheatmap")
  - if (!requireNamespace("BiocManager", quietly = TRUE))
     install.packages("BiocManager")
     BiocManager::install("AnnotationDbi", version = "3.8")
     BiocManager::install("tximport", version = "3.8")
  - see https://bioconductor.org/packages/release/bioc/html/
    AnnotationDbi.html
    http://bioconductor.org/packages/release/bioc/html/
    tximport.html
- open up the code in an editor or RStudio and take a look

## **Explore datasets**

- data folder has genome, protein, and GFF annotation in it
   or it will when you run the download.sh script
- How many annotated genes (gene features) are in the GFF file for each species? How many transcripts (mRNA features)
  - Can solve this many ways, unix commands grep, awk, wc will suffice
  - Can also do this in R

## **Explore datasets**

- analysis/ortho\_set1/Results
- Has OrthoFinder pre-computed results
- Explore these result files

# Tasks/Questions: OrthoFinder data

- Make a heat map of gene family sizes pick a cutoff like gene family total size > 25 but experiment with this
- scripts/plot\_heatmap\_family.R shows you how to work with this. Run it with Rscript scripts/plot\_heatmap\_family.R on UNIX
- I recommend Rstudio for interactive session (comment out the pdf() line if you do this so you can see the plots in your session
- You may need to install pheatmap in your R installation Do install.packages("pheatmap") in your R console.

#### Other challenges/questions

- How many singletons genes that have no homologs so aren't in a orthogroup - are there per species
  - Make a table with these numbers
  - How many single copy gene families are there across this data (eg 1:1:1 ... orthologs)

## Explore genome statistics

- Run the Rscript scripts/plot\_chroms\_1.R
- Examine some genome wide statistics like intron density and exon size and chromosome-wide plots of genes or introns/per gene etc.
- Explore and experiment with plotting different things like number of genes histograms across chromosomes

#### Some reference links

**TxDB** and Genomic Features

https://kasperdanielhansen.github.io/genbioconductor/html/GenomicFeatures.html

pheatmap

https://davetang.org/muse/2018/05/15/making-a-heatmap-in-r-with-the-pheatmap-package/