## Parse and Converting workshop

## **BLAST** report

- Ecoli proteins
- S\_enterica proteins
- Ecoli-vs-Senterica.BLASTP.tab.gz

Write script to read in the BLAST report. - Calculate for each alignment what the % aligned of the Ecoli query protein is? - Calculate the % of proteome that was aligned (out of the total number of Ecoli proteins)

## Orthofinder parsing

Here is a data file - OrthoFinder result

• Let's write a script which will summarize the data by Species.

The data look like <code>ORTHOGROUP GENENAME\_SP1</code> , <code>GENENAME2\_SP1 GENENAME\_SP2</code>