Bioinformatics and Genome Analyses

September 18 – December 15, 2017. Institut Pasteur Tunis https://webext.pasteur.fr/tekaia/BCGAIPT2017.html

Perl practical sessions

References:

Programming Perl (Book by Larry Wall) 3rd edition https://docstore.mik.ua/orelly/perl/prog3/index.htm

Perl Tutorial:

https://www.tutorialspoint.com/perl/index.htm

Common Practices (Programming Perl):

https://docstore.mik.ua/orelly/perl/prog3/ch24 01.htm

Sample Perl scripts (examples):

http://sifaka.cs.uiuc.edu/czhai/learnperl.html

Practical session:

- I. Consider simple sequences (dna or amino acids (aa)) Example: YAL068c.dna and YAL068c.prt
- Write a script to calculate:
- -base composition; GC% for nucleotide sequences
- Write a script to calculate:

Amino-acids composition for protein sequences

- -sequence size (base or amino-acids);
- -codon counts
- -extract a segment from a sequence corresponding to start and position;

Extractpartseq.pl (sequence identification, start position, end position)

- II. Use "goldData.xls" to compute different statistics about genome sequencing projects. goldData.xls was downloaded from: https://gold.jgi.doe.gov/downloadexceldata in august 2017. Most recent data can be downloaded from this URL.
- -Save the file goldData.xls into a text formatted file: goldData.txt with <tab> as column separator.
- -Write scripts to transform "\r" into "\n" in goldData.txt: trcr2nl.pl goldData.txt &

output file is: goldData.txt.nl mv goldData.txt.nl goldData.txt

-Replace white space "" in goldData.txt by "_" and Extract subsets of columns that will be used for next steps and keep in the output file solely the following columns:

GOLDSTAMP PROJECT_TYPE PROJECT_STATUS SEQUENCING_STATUS DOMAIN PHYLUM SPECIES (corresponding to columns: 0, 5, 6, 7, 12, 14, 19). extracttabgenomes.pl
The output file is goldDatafinal.txt

-Extract from goldDatafinal.txt, data tables each corresponding to a Domain (create a table per domain: Bacteria/Archaea/Eukaryotes/Viruses): output should be: Bateria.tab, Archaea.tab, Eukaryotes.tab, Viruses.tab. extractDomain.pl &

-Statistics: stats.pl &

The script *stats.pl* performs the following steps:

For each domain extract the following data table:

-WGS_Complete output file: Domain_WGS_Complete.tab
-WGS_Incomplete output file: Domain_WGS_Incomplete.tab
-WGS_Permanent_Draft output file: Domain_WGS_Permanent_Draft.tab
-Transcriptome_Complete output file: Domain_Transcriptome_Complete.tab
-Transcriptome incomplete output file: Domain_Transcriptome_incomplete.tab
and corresponding number of elements (size of the table).

Note "Domain" is Archaea/Bacteria/Eukaryote/Viruses.

The corresponding statistics output file is: statisticsgoldData.txt.

-Compute the PHYLUM distributions in each of the above output files related to a given domain.

Distributions of phylum per Project type:

WGS Complete

Bacteria: Bacteria_Complete.freq Archaea: Archaea_Complete.freq Eukaryotes: Eukaryotes_Complete.freq

WGS incomplete

Bacteria: Bacteria_Incomplete.freq Archaea: Archaea_Incomplete.freq Eukaryotes: Eukaryotes Incomplete.freq

WGS Permanent draft

Bacteria: Bacteria Permanent Draft.freq

Archaea: Archaea Incomplete.freq

Eukaryotes: Eukaryotes Permanent Draft.freq

Transcriptome Incomplete

Eukaryotes: Eukaryotes Transcriptome Incomplete.freq

For example: "PHYLUM" (column 6) distribution (in completely sequenced genomes). From each of these tables (Domain_WGS_Complete.tab) compute the Phylum distribution/Domain (number of occurrences/per phylum in decreasing order)

Phylum distribution in Bacteria: cut -f 6 Bacteria WGS Complete.tab | sort > ph

Write a script (freqsortednames.pl) that computes occurrences/identification. freqsortednames.pl ph & #(output file: ph.freq)

Note: eliminate "Phylum" from ph

sort -n -k 2 -r ph.freq > Bacteria_Complete.freq (insert a column header : « Phylum Nb » at the first line).

Phylum distribution in Archaea: cut -f 6 Archaea_WGS_Complete.tab | sort > ph freqsortednames.pl ph & Note: eliminate "Phylum" from ph

sort -n -k 2 -r ph.freq > Archaea_Complete.freq (insert a column header : « Phylum Nb » at the first line).

and so on for Eukaryotes and Viruses.

For example: Eukaryotes Complete.freq

Phylum Nb

Ascomycota 163
Streptophyta 50
Chordata 34
Apicomplexa 19
None 15
Nematoda 11
Arthropoda 10
Platyhelminthes 6
Basidiomycota 6

Microsporidia 5 Mucoromycota 2 Chlorophyta 2 Eustigmatophyceae 1

Bacillariophyta 1

III. Data manipulation:

Consider the GFF file (to be download from the ncbi server): GCF_000018785.1_ASM1878v1_genomic.gff

-What is the GFF format (search using google – look at the ncbi FAQ,..)?

od -c GCF_000018785.1_ASM1878v1_genomic.gff | more

can you guess the structure of this file (how many columns/line)?

Fredj Tekaia (tekaia@pasteur.fr)