Introduction to Unix shell - EXERCISES

- 2016/17 Part II BBS Bioinformatics
- 16 Jan 2017, 15:00-17:00
- Bioinformatics Training Room, Craik-Marshall Building, Downing Site
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and

- 2016/17 Part III Systems Biology SEB module
- 8 Feb 2017, 10:00-13:00
- Bioinformatics Training Room, Craik-Marshall Building, Downing Site
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- 1. Shakespeare
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Download and unpack the text files and the human reference genome annotation.

Exercises - Shakespeare

- 1. Check you have the shakespeare.txt file (all works of Shakespeare as text). Process it to output a list of words with frequency counts. Be careful not to count capitalised and non-capitalised words separately, and take care of the apostrophe!
- 2. Check how many times some country names are mentioned by Shakespeare. What are the most common words to co-occur in the same line with a country name? Filter your output for words shorter than four letters.
- 3. Find the most common bigrams Shakespeare uses. Trigrams?

Exercises - Human genome

- 1. How many genes are there in the reference genome? Don't forget to unpack the file.
- 2. How many transcripts does your favourite gene have, e.g. ENSG00000001461?
- 3. How many exons?
- 4. Produce a tab separated file with these columns: transcriptID, exon_number, exon_length.
- 5. Which exon is the longest?

Exercises - PDB

- 1. Extract the protein sequence from the PDB file 1A8Q.pdb (ATOM instances).
- 2. Chech if the sequence from ATOM instances matches the one in SEQRES.

Exercises - More sed

1. Write a script that would combine the split lines in split_lines.txt using sed.

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