## Shell scripting - 21/05/2020

## Programming for Bioinformatics - Module 3 Laurea Magistrale in Bioinformatics A.Y. 2019-2020

Consider the following data, saved into the space-separated file casp13.txt.

```
>H0974 048503/048504, , subunit 1, 72 residues;

MSYDYSSLLGKITEKCGTQYNFAIAMGLSERTVSLKLNDKVTWKDDEILKAVHVLELNPQ

DIPKYFFNAKVH

>H0974 048503/048504, , subunit 2, 95 residues;

MQTDTSNRLKQIMAERNLKQVDILNLSIPFQKKFGIKLSKSTLSQYVNSVQSPDQNRIYL

LAKTLGVSEAWLMGFDVPMVESSKIENDSHHHHHH

>H0980 Q3KP22-3 Q8NHR7, Human, subunit 1, 111 residues;

SLKPFTYPFPETRFLHAGPNVYKFKIRYGKSIRGEEIENKEVITQELEDSVRVVLGNLDN

LQPFATEHFIVFPYKSKWERVSHLKFKHGEIILIPYPFVFTLYVEMKWFHE

>H0980 Q3KP22-3 Q8NHR7, Human, subunit 2, 52 residues;

VNNMVTGYISIDAMKKFLGELHDFIPGTSGYLAYHVQNEINMSAIKNKLKRK

>H0993 MlaFA, E. coli, subunit 1, 269 residues;

MEQSVANLVDMRDVSFTRGNRCIFDNISLTVPRGKITAIMGPSGIGKTTLLRLIGGQIAP

DHGEILFDGENIPAMSRSRLYTVRKRMSMLFQSGALFTDMNVFDNVAYPLREHTQLPAPL

LHSTVMMKLEAVGLRGAAKLMPSELSGGMARRAALARAIALEPDLIMFDEPFVGQDPITM

GVLVKLISELNSALGVTCVVVSHDVPEVLSIADHAWILADKKIVAHGSAQALQANPDPRV
```

- 1. (2 points) What is the output of the following two commands?
  - (a) cat casp13.txt | sort | cut -c 1-4 | sort -u | head -2 | wc -1
  - (b) cat casp13.txt | sort | cut -c 1-4 | head -2 | sort -u | wc -l
- 2. (2 points) What is the output of the following command?

17 RQFLDGIADGPVPFRYPAGDYHADLLPGS

3. (3 points) What is the output of the following command?

```
cat casp13.txt | awk -F ',' '$0 ~ /;$/ {if($2 ~ /H/) print $2}' | sort -u | wc -l
```

4. (3 points) What is the output of the following command?

```
cat casp13.txt | sed '/;$/d' | sed '/^L/d' | wc -l
```