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 1-7 | sort -u | head -2 | wc -1
 - (b) cat casp13.txt | sort | cut -c 1-7 | head -2 | sort -u | wc -1
- 2. (2 points) What is the output of the following command?

17 RQFLDGIADGPVPFRYPAGDYHADLLPGS

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

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