Data and Text Processing for Health and Life Sciences Workbook

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http://labs.rd.ciencias.ulisboa.pt/book/

I - Multiple Choice¹:

- 1. A major problem of the nomenclature used in Health and Life Sciences:
 - (a) consistency
 - (b) inconsistency
 - (c) accuracy
 - (d) inaccuracy
- 2. Text files can contain data using a specific format, such as:
 - (a) TXT, CSV, and XLS
 - (b) TXT, XLS, and XML
 - (c) XLS, CSV, and XML
 - (d) TXT, CSV, and XML
- 3. One of the most common languages used to specify biomedical ontologies is:
 - (a) TXT
 - (b) OWL
 - (c) XLS
 - (d) CSV
- 4. A text format that can store multiple tables in a single file:
 - (a) CSV
 - (b) TSV
 - (c) XML
 - (d) XLS
- 5. To add a line to the end of file:
 - (a) echo 'line' >> file.txt
 - (b) echo 'line' > file.txt
 - (c) echo 'line' < file.txt</pre>
 - (d) echo 'line' | file.txt
- 6. To create a file containing the contents of a web page:
 - (a) curl 'https://www.ebi.ac.uk/...'
 - (b) curl 'https://www.ebi.ac.uk/...' > file.txt
 - (c) curl 'https://www.ebi.ac.uk/...' < file.txt</pre>
 - (d) curl 'https://www.ebi.ac.uk/...' | file.txt
- 7. To select only the lines of a XML file that specify the organism as Homo sapiens:
 - (a) cat '<name type="scientific">Homo sapiens</name>' file.xml
 - (b) sed '<name type="scientific">Homo sapiens</name>' file.xml
 - (c) grep '<name type="scientific">Homo sapiens</name>' file.xml
 - (d) xmllint --xpath '<name type="scientific">Homo sapiens</name>' file.xml
- 8. To obtain the species name of a protein in its XML file:
 - (a) grep '<name type="scientific">Homo sapiens</name>' P21817_entry.xml
 - (b) xmllint --xpath '//name/text()' P21817_entry.xml
 - (c) xmllint --xpath '//type/scientific/text()' P21817 entry.xml
 - (d) xmllint --xpath '//name[@type="scientific"]/text()' P21817_entry.xml
- 9. Which sequence matches the regular expression A?T*G+C:
 - (a) GGGC
 - (b) TTTG

¹ The answers are at the end of this workbook

- (c) AAAT
- (d) ATTC
- 10. Which sequence matches the regular expression ^A[^T]+G.C:
 - (a) TTGGC
 - (b) ATGGC
 - (c) AGGCC
 - (d) AGCCC

II - Output Identification:

1. Consider a file *test.txt* with the following three lines:

```
Caffeine is a compound.
I like caffeine drinks.
My favourite is the PowerCaffeine brand
```

(a) and the execution of the following two commands:

```
./reversemyfile.sh test.txt > test2.txt
tr -d 'a' < test2.txt > test3.txt
```

How many a's can we find in the first line of each file (test.txt, test2.txt and test3.txt)?

(b) and the execution of the following three commands:

```
grep 'is' test.txt > test1.txt
grep 'like' test.txt > test2.txt
grep -e 'is' -e 'like' test.txt > test3.txt
```

How many lines has each file (test1.txt, test2.txt and test3.txt)?

(c) and the execution of the following four commands:

```
echo $'is\nlike' > terms.txt
cat test.txt | xargs -I {} grep {} terms.txt > test1.txt
cat terms.txt | xargs -I {} grep {} terms.txt > test2.txt
cat terms.txt | xargs -I {} grep {} test.txt > test3.txt
How many lines has each file (test1.txt, test2.txt and test3.txt)?
```

2. Consider a file *test.txt* with the following two lines:

```
Caffeine is a compound. caffeine drinks are great.
```

Identify the output of the following command lines:

- (a) sed -E $'s/^c/n/'$ test.txt | wc -l
- (b) sed -E 's/^c/\n/i' test.txt | wc -l
- (c) sed -E 's/[c][ao]/ g ' test.txt | wc -l
- (d) sed -E $s/[^c][ao]/n/ig'$ test.txt | wc -l
- (e) sed -E 's/c[$^{\prime}$]+/\n/' test.txt | wc -l
- (f) sed -E $'s/c[^]+/n/i'$ test.txt | wc -l
- (g) sed -E $'s/c[^]+/n/ig'$ test.txt | wc -l
- (h) sed -E 's/ $[ca][^]+/n/ig'$ test.txt | wc -l
- 3. Consider a file *test.txt* with the following three lines:

```
Caffeine is a compound.
I like caffeine drinks.
My favourite is the PowerCaffeine brand.
```

Identify the output of the following command lines:

- (a) grep -c 'caffeine' test.txt
- (b) grep -c -e 'caffeine' -e 'Caffeine' test.txt
- (c) grep -c -i 'caffeine' test.txt
- (d) grep -c -i -w 'caffeine' test.txt

```
(e) grep -c -E 'caffeine' test.txt
(f) grep -c -E '(c|C)affeine' test.txt
(g) grep -c -E '.affeine' test.txt
(h) grep -c -E '[^c]affeine' test.txt

(i) sed -E 's/c/\n/' test.txt | wc -l
(j) sed -E 's/c/\n/i' test.txt | wc -l
(k) sed -E 's/c/\n/g' test.txt | wc -l
(l) sed -E 's/c/\n/ig' test.txt | wc -l
```

4. Consider a file *sequence.txt* with the following two lines:

```
ATAACGCTAG
AAACCCTTAG
```

Identify the output of the following command lines:

```
(a) grep -b -o -w -E 'AAA*' sequence.txt(b) grep -b -o -E '^AAA*' sequence.txt(c) grep -b -o -E 'AAA*$' sequence.txt
```

- (d) grep -b -o -E 'AAA*' sequence.txt
- 5. Consider a file *test.xml* with the following lines:

Identify the output of the following command lines:

```
(a) xmllint --xpath '/proteins/reference[1]/@id' test.xml
(b) xmllint --xpath '//proteins/reference[1]/@id' test.xml
(c) xmllint --xpath '//proteins/reference[2]/@date' test.xml
(d) xmllint --xpath '//proteins/reference[2]/text()' test.xml
```

6. Consider a file *test.owl* with the following lines:

```
<Class about="D3">
    <subClassOf resource="D1"/>
    <subClassOf resource="D2"/>
    <hasSynonym>MHN</hasSynonym>
    <hasSynonym>MHS</hasSynonym>
    <hasSynonym>MHE</hasSynonym>
    <label>MHX</label>
</Class>
```

Identify the output of the following command lines:

```
(a) xmllint --xpath '//Class[@about="D3"]' test.owl |
    grep -c 'MHN'
(b) xmllint --xpath '//subClassOf[@resource="D1"]/..' test.owl |
    grep -c 'MHN'
(c) xmllint --xpath '//subClassOf[@resource="D1"]/../hasSynonym' test.owl |
    grep -c 'MHN'
```

```
(d) xmllint --xpath '//subClassOf[@resource="D1"]/../label' test.owl |
    grep -c 'MHN'

(e) xmllint --xpath '//Class[@about="D3"]' test.owl |
    grep -o -E 'MH.' | wc -l

(f) xmllint --xpath '//subClassOf[@resource="D1"]/..' test.owl |
    grep -o -E 'MH.' | wc -l

(g) xmllint --xpath '//subClassOf[@resource="D1"]/../hasSynonym' test.owl |
    grep -o -E 'MH.' | wc -l

(h) xmllint --xpath '//subClassOf[@resource="D1"]/../label' test.owl |
    grep -o -E 'MH.' | wc -l
```

7. Consider a file *test.owl* with the following lines:

Identify the output of the following command lines:

```
(a) echo "D2" | ./getparents.sh test.owl | wc -l (b) echo "D2" | ./getancestors.sh test.owl | wc -l (c) echo "D3" | ./getparents.sh test.owl | wc -l (d) echo "D3" | ./getancestors.sh test.owl | wc -l
```

8. Consider a file *lexicon.txt* with the following two lines:

```
measles encephalitis
measles encephalitis, disorder
measles (encephalitis)
(Post) measles encephalitis, disorder
```

Identify the output of the following command lines:

```
(a) cat lexicon.txt |
    grep -c -E '^measles encephalitis$'
(b) tr -d '()' < lexicon.txt |
    grep -c -E '^measles encephalitis$'
(c) sed -E 's/, .*$//' < lexicon.txt |
    grep -c -E '^measles encephalitis$'
(d) sed -E 's/, .*$//' < lexicon.txt | tr -d '()' |
    grep -c -E '^measles encephalitis$'</pre>
```

III – Script Modification:

1. Consider this version of the *getpublications.sh* script:

```
ID=$1
rm -f chebi\_$ID\_*.rdf # Removes any previous files
grep -l '<name type="scientific">Homo sapiens</name>' chebi\_$ID\_*.xml | \
xargs -I {} grep '<dbReference type="PubMed"' {} | \
gawk -F\" '{ print $4 }' | sort -u | \
xargs -I {} curl 'https://www.uniprot.org/citations/{}.rdf' -o chebi\_$ID\_{}.rdf</pre>
```

Identify which line needs to be replaced and the new command lines to replace it, so the script:

- (a) saves all the publication links in a file named *links.txt*
- (b) uses a xpath guery to extract the PubMed identifiers
- 2. Consider this version of the *getpublications.sh* script:

Identify which line needs to be replaced and the new command lines to replace it, so the script:

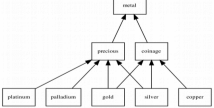
- (a) removes also all the numbers in the beginning of a label
- (b) removes the numbers that follow the words disease or syndrome
- 3. Consider this version of the *getproteins.sh* script:

Identify which lines needs to be replaced and the new command lines to replace it, so the script:

- (a) saves all the protein links in a file named *proteins.txt*
- (b) receives as argument a filename containing a list of ChEBI identifiers (one per line)

IV - Essay:

1. Which of the other four metals are more and less similar to platinum?



- 2. Both grep and awk were used as data filters, why we used these two commands instead of just one of them?
- 3. Both grep and xmllint were used as data filters, why we used these two commands instead of just one of them?
- 4. Both regular expressions and xpath queries were used for data selection, why we used these two techniques instead of just one of them?
- 5. Both regular expressions and exact match queries were used for text selection, why we used these two techniques instead of just one of them? Provide an example.

Answers

```
Ι
       1-b, 2-d, 3-b, 4-c, 5-a, 6-b, 7-c, 8-d, 9-a, 10-c
II
       1. (a) 2, 3, 0 (b) 2, 1, 3, (c), 0,2,3
       2. (a-h) 3, 4, 7, 6, 4, 4, 5, 7
       3. (a-l) 1, 3, 3, 2, 1, 3, 3, 2, 5, 6, 5, 7
       4. -, 11:AAA, -, 2:AA 11:AAA
       5. (a) XPath set is empty, (b) id="1", (c) date="2012", (d) third reference
       6. (a-h) 1, 1, 1, 0, 4, 4, 3, 1
       7. (a-d) 1, 1, 1, 2
       8. (a-d) 1, 2, 2, 3
III
       1(a) Line 6 needs to be replaced by:
      xargs -I {} echo 'https://www.uniprot.org/citations/{}.rdf' >> links.txt
       1(b) Line 4 needs to be replaced by:
      xargs -I {} xmllint --xpath '//dbReference[@type="PubMed"]/@id'
            chebi 27732 P21817 entry.xml | tr ' '\n' | tr 'id' '""' | \
      2(a) Line 6 needs to be replaced by
       sed -E 's/[,:;] .*$//; s/^ *//; s/ *$//'; s/^[0-9]+//' | \
      2(b) Line 6 needs to be replaced by
      sed -E 's/[,:;] .*$//; s/^ *//; s/ *$//';
            's/(disease|syndrome) [0-9]+/\1/' | \
      3(a) Line 6 needs to be replaced by
      xargs -I {} echo 'https://www.uniprot.org/uniprot/{}.xml' >> proteins.txt
      3(b) Line 3 needs to be replaced by
      cat $ID | xargs -I {} curl -s "https://www.ebi.ac.uk/chebi/
           viewDbAutoXrefs.do?d-1169080-e=1&6578706f7274=1&chebiId={}&
           dbName=UniProt" | \
```

IV

- 1. more similar is palladium, and less similar is copper
- 2. grep was used to filter lines (rows), while awk was used to filter data elements (columns) in the lines
- 3. grep was used to filter lines (rows), while xmllint for filtering XML data elements that may not be represented in a single line
- 4. Regular expressions were used in grep to match lines (rows) of text file, while xpath queries were used in xmllint to match data elements in a XML file.
- 5. Regular expressions enabled us to match text with some variations, while exact match only matches text that is completely equal to the given query. For example, the query A.C matches ABC and A.C when used as a regular expression, but only matches A.C when used as an exact match.