# Introduction to the Unix shell for biologists

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The source code can be found at:

https://github.com/konrad/Introduction to the Unix Shell for biologists/

## Motivation and background

In this course you will learn the basics of how to use the Unix shell. Unix is a class of operating systems with many different flavors including well-known ones like GNU/Linux and the BSDs. The development of Unix and its shell (also known as command line interface) dates back to the late 1960s. Still, their concepts lead to very powerful tools. In the command line you can easily combine different tools into pipelines, avoid repetitive work and make your workflow reproducible. Knowing how to use the shell will also enable you to run programs that are only developed for this environment which is the case for many bioinformatical tools.

### Work environment and test files

During this course all of you are working on Ubuntu (version 14.04) which is a widely used GNU/Linux distribution. The systems boots from a USB stick which offers you to run a live system or to install Ubuntu on your computer. We will run the live mode which will not change the system installed on your PC. After shutting the live system down and removing the stick everything on the computer will be as before.

To get test data click on the Dash button on the top left of your screen, type terminal and click on the Terminal icon. You will learn later what you are doing but for the moment just type the following commands into the command line interface. Do not write the dollar sign(\$). It just indicates the so called prompt:

```
$ wget http://data.imib-zinf.net/unix_course_files.tar.gz
$ tar xfz unix_course_files.tar.gz
$ rm unix_course_files.tar.gz
```

# The basic anatomy of a command line call

Running a tool in the command line interface follows a simple pattern. At first you have to write the name of the command (if it is not globally installed it's precise location needs to be given - we will get to this later). Some programs additionally require parameters and arguments. Parameters usually start with a dash (-). The common pattern looks like this (<> indicates obligatory items, [] indicates optional items):

```
cprogram name> [parameters] [arguments]
```

An example is calling the program 1s which lists the content of a directory. You can simply call it without any parameter

#### \$ ls

or with one or more parameters

```
$ ls -l
$ ls -lh
```

or with one or more arguments

```
$ ls test folder
```

or with one or more parameters and arguments

```
$ ls -l test_folder
```

The result of a command is written usually to the so called *standard output* of the shell which is the screen shown to you. We will later learn how to redirect this e.g. to the *standard input* of another program.

## How to get help and documentation

Especially in the beginning you will have a lot of questions what a command does and which arguments and parameters need to be given. One rule before using a command or before asking somebody about it is called RTFM (please check the meaning yourself). Maybe the most important command is man which stands for manual. Most commands offer a manual and with man you can read those. To get the documentation of 1s type

#### \$ man ls

To close the manual use q. Additionally or alternatively many tools offer some help via the parameter -h, -help or --help. For example 1s:

```
$ ls --help
```

Other tools present this help if they are called without any parameters or arguments.

# Bash keyboard shortcuts

There are different implementations of the Unix shell. You are currently working with Bash (**B**ourne-again shell). Bash has several keyboard shortcuts that improve the interaction. Here is a small selection:

- Ctrl-c Stop the command
- Ctlr-↑ Go backward in command history
- Ctlr-↓ Go forward in command history
- Ctrl-a Jump to the beginning of a line
- Ctrl-e Jump to the end of a line
- Ctrl-u Remove everything before the cursor position
- Ctrl-k Remove everything after the cursor position
- Ctrl-l Clean the screen
- Ctrl-r Search in command history
- Tab extend commands and file/folder names

### Files, folders, locations

Topics:

- ls
- pwd
- cd
- mkdir
- Relative vs. absolute path
- ~/

In this part you will learn how to navigate through the file system, explore the content of folders and create folders

At first we need to know where we are. If you open a new terminal you should be in your home directory (we will explain this below). To test this, call the program pwd which stands for print working directory.

# \$ pwd /home/ubuntu

The default user of the Ubuntu live system is called ubuntu. In general each user has a folder with its user name located inside the folder home. The next command we need and which has been already mentioned above is 1s. It simply lists the content of a folder. If you call it without any arguments it will output the content of the current folder. Using 1s we want to get a rough overview of what a common Unix file system tree looks like and learn how to address files and folders. The root folder of a systems starts with /. Call

#### \$ ls /

to see the content of the root folder. You should see something like

```
bin
      data
            etc lib
                        lost+found
                                   mnt
                                        proc
                                              run
                                                          tmp
                                                              var
                                                     srv
      dev
            home lib64
                       media
boot
                                opt
                                   root sbin sys
```

There are several subfolders in the so-called root folder (and yes, to make it a little bit confusing there is even a folder called root in the root folder). Those are more important if you are the administrator of the system. Normal users do not have the permission to make changes here. Currently your home directory is your little universe in which you can do whatever you want. In here we will learn how work with paths. A file or folder can be addressed either with its absolute or relative path. As you have downloaded and decompressed the test data you should have a folder unix\_course\_files located in your home folder. Assuming you are in this folder (/home/ubuntu/) the relative path to the folder is simply unix\_course\_files. You can get the content of the folder listed by calling 1s like this:

```
$ ls unix_course_files
```

This is the so called *relative path* as it is relative to the current work directory /home/ubuntu/. The *absolute path* would start with a / and is /home/ubuntu/unix\_course\_files. Call ls like this:

#### \$ ls /home/ubuntu/unix\_course\_files

There are some conventions regarding relative and  $absolute\ paths$ . One is that a dot (.) represents the current folder. The command

```
$ ls ./
```

should return the same as simply calling

#### \$ ls

Two dots (...) represent the parent folder. If you call

#### \$ ls ../

you should see the content of /home. If you call

#### \$ ls ../../

you should see the content of the parent folder of the parent folder which is the root folder (/) assuming you are in /home/ubuntu/. Another convention is that ~/ represents the home directory of the user. The command

#### \$ ls ~/

should list the content of your home directory independent of your current location in the file system.

Now as we know where we are and what is there we can start to change our location. For this we use the command cd (change directory). If you are in your home directory /home/ubuntu/ you can go into the folder unix\_course\_files by typing

#### \$ cd unix\_course\_files

After that call pwd to make sure that you are in the correct folder.

#### \$ pwd

/home/ubuntu/unix\_course\_files

To go back into your home directory you have different options. Use the absolute path

#### \$ cd /home/ubuntu/

or the above mentioned convention for the home directory ~/:

#### \$ cd ~/

or the *relative path*, in this case the parent directory of /home/ubuntu/unix\_course\_files:

#### \$ cd ../

As the home directory is such an important place cd uses this as default argument. This means if you call cd without argument you will go to the home directory. Test this behavior by calling

#### \$ cd

Try now to go to different locations in the file system and list the files and folders located there.

Now we will create our first folder using the command mkdir (make directory). Go into the home directory and type:

#### \$ mkdir my\_first\_folder

Here we can discuss the implementation of another Unix philosophy: "No news is good news." The command successfully created the folder my\_first\_folder. You can check this by calling ls, but mkdir did not tell you this. If you do not get a message this usually means everything went fine. If you call the above mkdir command again you should get an error message like this:

```
$ mkdir my_first_folder
mkdir: cannot create directory 'my_first_folder': File exists
```

So if a command does not complain you can usually assume there was no error.

## Manipulating files and folder

Topics:

- touch
- cp
- mv
- rm

Next we want to manipulate files and folders. We create some dummy files using touch which is usually used to change the time stamps of files. But you can also create empty files with it easily. Let's create a file called test\_file\_1.txt:

```
$ touch test_file_1.txt
```

Use 1s to check that it was created.

The command cp (copy) can be used to copy files. For this it requires at least two arguments: the source and the target file. In the following example we generate a copy of the file test\_file\_1.txt called a\_copy\_of\_test\_file.txt.

```
$ cp test_file_1.txt a_copy_of_test_file.txt
```

Use 1s to confirm that this worked. We can also copy the file in the folder my\_first\_folder which we have created above:

```
$ cp test_file_1.txt my_first_folder
```

Now there should be also a file test\_file\_1.txt in the folder my\_first\_folder. If you want to copy a folder and its content you have to use the parameter -r.

```
$ cp -r my_first_folder a_copy_of_my_first_folder
```

You can use the command mv (move) to rename or relocate files or folders. To rename the file a\_copy\_of\_test\_file.txt to test\_file\_with\_new\_name.txt call

```
$ mv a_copy_of_test_file.txt test_file_with_new_name.txt
```

With mv you can also move a file into a folder. For this the second argument has to be a folder. For example, to move the file now named test\_file\_with\_new\_name.txt into the folder my\_first\_folder use

```
$ mv test_file_with_new_name.txt my_first_folder
```

You are not limited to one file if you want to move them into a folder. Let's create and move two files file1 and file2 into the folder my\_first\_folder.

```
$ touch file1 file2
$ mv file1 file2 my_first_folder
```

At this point we can introduce another handy feature most shells offer which is called *globbing*. Let us assume you want to apply the same command to several files. Instead of explicitly writing all the file names you can use a *globbing pattern* to address them. There are different wildcards that can be used for these patterns. The most important one is the asterisk (\*). It can replace none, one or more characters. Let us explore this with a small example:

```
$ touch file1.txt file2.txt file3
$ ls *txt
$ mv *txt my_first_folder
```

The ls shows the two files matching the given pattern (i.e. file1.txt and file2.txt) while dismissing the one not matching (i.e. file3). Same for mv - it will only move the two files ending with txt.

We accumulated several test files that we do not need anymore. Time to clean up a little bit. With the command rm (remove) you can delete files and folders. Please be aware that there is no such thing as a trash bin if you remove items this way. They will be gone for good and without further notice.

To delete a file in my\_first\_folder call:

```
$ rm my_first_folder/file1.txt
```

To remove a folder use the parameter -r (recursive):

```
$ rm -r my_first_folder
```

# File content - part 1

Topics:

- less / more
- cat
- echo
- head
- tail
- cut

Until now we did not care about the content of the files. This will change now. Please go into the folder unix\_course\_files:

```
$ cd unix_course_files
```

There should be some files waiting for you. To read the content with the possibility to scroll around we need a so called pager program. Most Unix systems offer the programs more and less which have very similar functionalities ("more or less are more or less the same"). We will use the later one here. Let's open the file origin\_of\_species.txt

#### \$ less origin\_of\_species.txt

The file contains Charles Darwin's *Origin of species* in plain text. You can scroll up and down line-wise using the arrow keys or page-wise using the page-up/page-down keys. To quit use the key q. With pager programs you can read file content interactively, but sometimes you just want to have the content of a file given to you (i.e. on the *standard output*). The command cat (*concatenate*) does that for one or more files. Let us use it to see what is in the example file two\_lines.txt. Assuming you are in the folder unix\_course\_files you can call

#### \$ cat two\_lines.txt

The content of the file is shown to you. You can apply the command to two files and the content is concatenated and returned:

```
$ cat two_lines.txt three_lines.txt
```

This is a good time to introduce the *standard input* and *standard output* and what you can do with it. Above I wrote the output is given to you. This means it is written to the so called *standard output*. You can redirect the *standard output* into a file by using >. Let us use the call above to generate a new file that contains the combined content of both files:

```
$ cat two_lines.txt three_lines.txt > five_lines.txt
```

Please have a look at the content of this file:

```
$ cat five_lines.txt
```

The *standard output* can also be redirected to other tools as *standard input*. More about this below. With cat we can reuse the existing file content. To create something new we use the command echo which writes a given string to the standard output.

```
$ echo "Something very creative"
```

To redirect the output into a target file use >.

```
$ echo "Something very creative." > creative.txt
```

Be aware that this can be dangerous. You will overwrite the content of an existing file. For example if you call now

```
$ echo "Something very uncreative." > creative.txt
```

there will be only the latest string written to the file and the previous one will be overwritten. To append the output of a command to a file without overwriting the content use >>.

```
$ echo "Something very creative." > creative.txt
$ echo "Something very uncreative." >> creative.txt
```

Now creative.txt should contain two lines.

Sometimes you just want to get an excerpt of a file e.g. just the first or last lines of it. For this the commands head and tail can be used. Per default 10 lines are shown. You can use the parameter -n <NUMBER> (e.g. -n 20 or just -<NUMBER> (e.g. -20) to specify the number of lines to be displayed. Test the tools with the file origin\_of\_species.txt:

```
$ head origin_of_species.txt
$ tail origin_of_species.txt
```

You cannot only select vertically but also horizontally using the command cut. Let us extract only the first 10 characters of each line in the file origin\_of\_species.txt:

```
$ cut -c 1-10 origin_of_species.txt
```

The tool cut can be very useful to extract certain columns from CSV files (comma/character separated values). Have a look at the content of the file genes.csv. You see that it contains different columns that are tabular-separated. You can extract selected columns with cut:

```
$ cut -f 1,4 genes.csv
```

## File content - part 1

Topics:

- WC
- sort
- uniq
- grep
- cut
- tr

There are several tools that let you manipulate the content of a plain text file or return information about it. If you want for example some statistics about the number of character, words and lines use the command wc. Let us count the number of lines in the file origin\_of\_species.txt:

```
$ wc -l origin_of_species.txt
```

You can use the command sort to sort a file alpha-numerically. Test the following calls

```
$ sort unsorted_numbers.txt
$ sort -n unsorted_numbers.txt
$ sort -rn unsorted_numbers.txt
```

and try to understand the output.

The tool uniq takes a sorted list of lines and removes line-wise the redundancy. Please have a look at the content of the file redundant.txt. Then use uniq to generate a non-redundant list:

```
$ uniq redundant.txt
```

If you call uniq with -c you get the number of occurrence for each remaining entry:

```
$ uniq -c redundant.txt
```

With the tool grep you can extract lines that match a given pattern. For instance, if you want to find all lines in origin\_of\_species.txt that contain the word species call

```
$ grep species origin_of_species.txt
```

As you can see we only get the lines that contain **species** but not the ones that contain **Species**. To make the search case-insensitive use the parameter -i.

```
$ grep -i species origin_of_species.txt
```

If you are only interested in the number of lines that match the pattern use -c:

```
$ grep -ic species origin_of_species.txt
```

The program tr (translate) exchanges one character by another. It reads from the standard input and performs the replacement. To direct the content of a file as standard input into a program < is applied. Have a quick look at the content of the file DNA.txt.

```
$ cat DNA.txt
```

We now want to replace all Ts in the file by Us. For this we call:

```
$ tr T U < DNA.txt
```

## Connecting tools

Another piece of the Unix philosophy is to build small tools that do one thing optimally and use the standard input and standard output. The real power of Unix builds on the capability to easily connect tools. For this so-called *pipes* are used. To use the *standard output* of one tool as *standard input* of another tool the vertical bar | is used. For example, in order to extract the first 1000 lines from origin\_of\_species.txt, search for lines that contain species, then search in those lines the ones which contain wild and finally replace the ws by ms call (Please write this in one line in the shell and remove the \):

# Examples analysis

Equipped with a fine selection of useful programs and basic understanding of how to combine them, we will no apply them to analyze real biological data.

### Retrieving data

You have used the tool wget above to download the example files. It is very useful, especially, if you want to retrieve large data sets. We download the fasta file of the *Salmonella* Thyphimuirum SL1344 chromosome by calling (in this document the URL is split into three lines. Please write it in one line in the shell and remove the \).

```
wget ftp://ftp.ncbi.nih.gov/genomes/Bacteria/\
    Salmonella_enterica_serovar_Typhimurium_SL1344_uid86645/\
    NC_016810.fna
```

Additionally, we download the annotation in GFF format of the same replicon:

```
wget ftp://ftp.ncbi.nih.gov/genomes/Bacteria/\
    Salmonella_enterica_serovar_Typhimurium_SL1344_uid86645/\
    NC_016810.gff
```

### Counting the number of features

Use less to have a look at NC\_016810.gff. It is a tabular-separated file. The first 5 lines start with # and are called header. Then several lines with 9 columns follow. The third column contains the type of the entry (gene, CDS, tRNA, rRNA, etc). If we want to know the numbers of tRNA entries we could try to apply grep and use -c to count the number of matching lines.

```
$ grep -c tRNA NC_016810.gff
```

This leads to a suspiciously large number. The issue is that the string tRNA also occurs in the attribute column (the 9th column). We just want to select lines with a match in the third column. This can be achieved by combining cut and grep.

```
$ cut -f 3 NC_016810.gff | grep -c tRNA
```

To get the number of entries for all other features we could just replace the tRNA e.g. by rRNA. But we can also get the number for all of them at once using this constellation:

```
$ grep -v "#" NC_016810.gff | cut -f 3 | sort | uniq -c
```

Try to understand what we did here. You can use a similar call to count the number genes on the plus and minus strand:

```
$ cut -f 3,7 NC_016810.gff | grep gene | sort | uniq -c
```

### Calculate the GC content of a genome

Let us assume the GC content of the genome is not known to us. We can use a handful of commands to calculate this quickly. We can gain the number of nucleotides in the following manner.

```
grep -v ">" NC_016810.fna | grep -o "A" | wc -l
grep -v ">" NC_016810.fna | grep -o "C" | wc -l
grep -v ">" NC_016810.fna | grep -o "G" | wc -l
grep -v ">" NC_016810.fna | grep -o "T" | wc -l
```

As we only need to get the sum of As and Ts as well as Cs and Gs we can use an extended pattern for grep. The  $\mid$  means or:

```
grep -v ">" NC_016810.fna | grep -Eo "A|T" | wc -l
grep -v ">" NC_016810.fna | grep -Eo "C|G" | wc -l
```

Once we have the number we can calculate the GC content by piping a formula into the calculator bc.

```
echo "scale=5; 2332503/(2332503+2545509)*100" | bc
```

### Multiple sequence alignment with muscle

We cannot only work with the default tools of the Unix shell but additionally have now access to a plethora of command line tools. Let's assume we want to perform a multiple alignment of the members of the GlmZ family. We choose muscle for this purpose. Its web site offers compiled binaries which means we only have to download the containing archive via (again, please write it in one line in the shell and remove the \)

```
$ wget http://www.drive5.com/muscle/downloads3.8.31/\
    muscle3.8.31_i86linux64.tar.gz
```

and extract it:

```
$ tar xfz muscle3.8.31_i86linux64.tar.gz
```

As we might need this tool more often (this is purely hypothetical as once you shutdown the live system any data will be gone) we generate a folder bin in our home directory. This is by convention a place were those programs are stored.

#### \$ mkdir bin

Then we move the tool into the folder and rename it:

```
$ mv muscle3.8.31_i86linux64 ~/bin/muscle
```

and clean up a little bit:

```
$ rm muscle3.8.31_i86linux64.tar.gz
```

Now we download the sequences of the RNAs which we want to align (again, please write the URL in one line and remove the \).

```
$ wget -0 RF00083.fa "http://rfam.xfam.org/family/RF00083/\
alignment?acc=RF00083&format=fastau&download=1"
```

Have a look at the content of the file using less or cat.

If you call muscle without anything you will get a list of parameters.

#### \$ ~/bin/muscle

Please be aware that we have to give the path to muscle.

We want to specify an input file using (-in) and an output file (-out):

```
$ ~/bin/muscle -in RF00083.fa -out RF00083_aligned.fa
```

Now we have the alignments stored in RF00083\_aligned.fa.

# Very, very basic scripting

One huge advantage of the Unix shell is that you can script actions. For example you can write the command for the multiple alignment into a file e.g. using echo:

```
$ echo "~/bin/muscle -in RF00083.fa -out RF00083_aligned.fa" \
> run_me.sh
```

If you want to run the command in that script you can call the script in the following manner:

```
$ bash run_me.sh
```

Shell scripting offers very powerful options to program workflows. Due to time restriction we will not cover this here.

### What's next

Here we just covered a small selection of tools and possibilities and hope that you can extend your Unix skills based on this knowledge yourself. There are many basic tools we have not covered but which could be important, e.g., archiving and compression tools like tar, bzip2 and gzip. For more powerful text manipulation sed and awk are good choices. We also recommend to get familiar with text editors which can be used to interactively modify text files. Classic Unix environment editors are vi (and derivatives like vim) or Emacs. While they are very powerful they have a steep learning curve. For beginners gedit that offers a graphical user interface could be another option.