



Intermediate Linux Course

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More Commandline Tools

Here is a quick list of useful commandline tools which will be used throughout the rest of the document. Many of these tools have quite extensive functionality and only a very limited part can be discussed here, so the reader is encouraged to read more about these using the links given in the *Links* section...

1.1 Command-line Tools

1.1.1 GZIP

gzip is a compression/decompression tool. When used on a file (without any parameters) it will compress it and replace the file by a compressed version with the extension '.gz' attached:

```
# ls textfile*
textfile
# gzip textfile
# ls textfile*
textfile.gz
```

To revert this / to uncompress, use the parameter -d:

```
# ls textfile*
textfile.gz
# gzip -d textfile
# ls textfile*
textfile
```

Note: As a convenience, on most Linux systems, a shellscript named gunzip exists which simply calls gzip -d

1.1.2 TAR

tar (tape archive) is a tool to handle archives. Initially it was created to combine multiple files/directories to be written onto tape, it is now the standard tool to collect files for distribution or archiving.

tar stores the permissions of the files within an archive and also copies special files (such as symlinks etc.), which makes it an ideal tool for archiving... Usually tar is used in conjunction with a compression tool such as gzip to create a compressed archive:

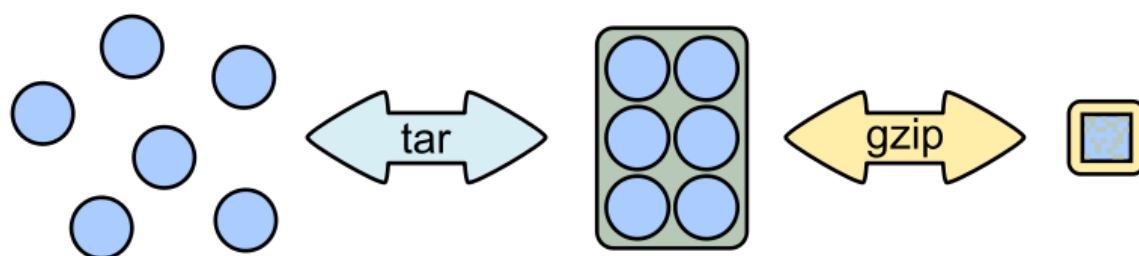


Figure 1.1: source: Th0msn80 (Wikipedia)

The most common command-line switches are:

Option:	Effect:
-c	create an archive
-t	test an archive
-x	extract an archive
-z	use gzip compression
-f	filename filename of the archive

Note: Don't forget to specify the target filename. It needs to follow the -f parameter. Although you can combine options like such: `tar -czf archive.tar` the order matters, so `tar -cfz archive.tar` will *not* do what you want...

Creating an archive containing two files:

```
# tar -cf archive.tar textfile1 textfile2
```

Listing the contents of an archive:

```
# tar -tf archive.tar
textfile1
textfile2
```

Extracting an archive:

```
# tar -xf archive.tar
```

Creating and extracting a compressed archive containing two files:

```
# tar -czf archive.tar.gz textfile1 textfile2
# tar -xzf archive.tar.gz
```

Creating a backup (eg. before doing something dangerous?):

```
# tar -czf /folder/containing/the/BACKUP.tgz /folder/you/want/to/backup
```

1.1.3 GREP

grep finds lines matching a pattern in textfiles.

Usage: grep [options] pattern file(s)

```
# grep -i ensembl P04637.txt

DR Ensembl; ENST000000269305; ENSP000000269305; ENSG000000141510.
DR Ensembl; ENST000000359597; ENSP000000352610; ENSG000000141510.
DR Ensembl; ENST000000419024; ENSP000000402130; ENSG000000141510.
DR Ensembl; ENST000000420246; ENSP000000391127; ENSG000000141510.
DR Ensembl; ENST000000445888; ENSP000000391478; ENSG000000141510.
DR Ensembl; ENST000000455263; ENSP000000398846; ENSG000000141510.
```

Useful options:

Option:	Effect:
-v	Print lines that do not match
-i	Search case-insensitive
-l	List files with matching lines, not the lines itself
-L	List files without matches
-c	Print count of matching lines for each file

Count the number of fasta sequences (they start with a ">") in a file:

```
# grep -c '>' twofiles.fasta
2
```

List all files containing the term "Ensembl":

```
# grep -l Ensembl *.txt
P04062.txt
P12931.txt
```

1.1.4 SED

sed is a Stream EDitor, it modifies text (text can be a file or a pipe) on the fly.

Usage: 'sed command file',

The most common usecases are:

Usecase	Command:
Substitute TEXT by REPLACEMENT:	's/TEXT/REPLACEMENT/'
Transliterate the characters x a, and y b:	'y/xy/ab/'
Print lines containing PATTERN:	'/PATTERN/p'
Delete lines containing PATTERN:	'/PATTERN/d'

```
# echo "This is text." | sed 's/text/replaced stuff/'
This is replaced stuff.
```

By default, text substitution are performed only once per line. You need to add a trailing 'g' option, to make the substitution 'global' ('s/TEXT/REPLACEMENT/g'), meaning all occurrences in a line are substituted (not just the first in each line). Note the difference:

```
# echo "ACCAAGCATTGGAGGAATATCGTAGGTAAA" | sed 's/A/_/'
_CCAAGCATTGGAGGAATATCGTAGGTAAA

# echo "ACCAAGCATTGGAGGAATATCGTAGGTAAA" | sed 's/A/_/g'
_CC__GC_TTGG_GG__T_TCGT_GGT____
```

When used on a file, sed prints the file to standard output, replacing text as it goes along:

```
# echo "This is text" > textfile
# echo "This is even more text" >> textfile
# sed 's/text/stuff/' textfile
This is stuff
This is even more stuff
```

sed can also be used to print certain lines (not replacing text) that match a pattern. For this you leave out the leading 's' and just provide a pattern: '/PATTERN/p'. The trailing letter determines, what sed should do with the text that matches the pattern ('p': print, 'd': delete)

```
# sed '/more/p' textfile
This is text
This is even more text
This is even more text
```

As sed by default prints each line, you see the line that matched the pattern, printed twice. Use option '-n' to suppress default printing of lines.

```
# sed -n '/more/p' textfile
This is even more text
```

Delete lines matching the pattern:

```
# sed '/more/d' textfile
This is text
```

Multiple sed statements can be applied to the same input stream by prepending each by option '-e' (edit):

```
# sed -e 's/text/good stuff/' -e 's/This/That/' textfile
That is good stuff
That is even more good stuff
```

Normally, sed prints the text from a file to standard output. But you can also edit files in place. Be careful - this will change the file! The '-i' (in-place editing) won't print the output. As a safety measure, this option will ask for an extension that will be used to rename the original file to. For instance, the following option '-i.bak' will edit the file and rename the original file to textfile.bak:


```
# sed -i.bak 's/text/stuff/' textfile
# cat textfile
This is stuff
This is even more stuff
# cat textfile.bak
This is text
This is even more text
```

1.1.5 AWK

awk is more than just a command, it is a complete text processing language (the name is an abbreviation of the author's names). Each line of the input (file or pipe) is treated as a record and is broken into fields. Generally, awk commands are of the form:

```
awk condition { action }
```

where:

- condition is typically an expression
- action is a series of commands

If no condition is given, the action is applied to each line, otherwise just to the lines that match the condition.

```
# awk '{print}' textfile
This is text
This is even more text

# awk '/more/ {print}' textfile
This is even more text
```

awk reads each line of input and automatically splits the line into columns. These columns can be addressed via \$1, \$2 and so on (\$0 represents the whole line). So an easy way to print or rearrange columns of text is:

```
# echo "Bob likes Sue" | awk '{print $3, $2, $1}'
Sue likes Bob

# echo "Master Obi-Wan has lost a planet" | awk '{print $4,$5,$6,$1,$2,$3}'
lost a planet Master Obi-Wan has
```

awk splits text by default on whitespace (spaces or tabs), which might not be ideal in all situations. To change the field separator (FS), use option '-F' (remember to quote the field separator):

```
# echo "field1,field2,field2" | awk -F',' '{print $2, $1}'
field2 field1
```

Note two things here: First, the field separator is not printed, and second, if you want to have space between the output fields, you actually need to separate them by a comma or they will be catenated together...

```
# echo "field1,field2,field2" | awk -F',' '{print $1 $2 $3}'
field1field2field3
```

You can also combine the pattern matching and the column selection techniques:

```
# awk '/more/ {print $3}' textfile
even
```

awk really is powerful in filtering out columns, you can for instance print only certain columns of certain lines. Here we print the third column of those lines where the fourth column is 'more':

```
# awk '$4=="more" {print $3}' textfile
even
```

Note the double equal signs "==" to check for equality and note the quotes around "more". If you want to match a field, but not exactly, you can use '~' instead of '==':

```
# awk '$4~"ore" {print $3}' textfile
even
```

1.1.6 Hints

Quoting

In Programming it is often necessary to “glue together” certain words. Usually, a program or the shell splits sentences by whitespace (space or tabulators) and treats each word individually. In order to tell the computer that certain words belong together, you need to “quote” them, using either single (') or double (") quotes. The difference between these two is generally that within double quotes, variables will be expanded, while everything within single quotes is treated as string literal. When setting a variable, it doesn't matter which quotes you use:

```
# MYVAR=This is set
-bash: is: command not found

# MYVAR='This is set'
# echo $MYVAR
This is set
# MYVAR="This is set"
# echo $MYVAR
This is set
```

However, it does matter, when using (expanding) the variable: Double quotes:

```
# export MYVAR=123
# echo "the variable is $MYVAR"
the variable is 123
# echo "the variable is set" | sed "s/set/$MYVAR/"
the variable is 123
```

Single quotes:

```
# export MYVAR=123
# echo 'the variable is $MYVAR'
the variable is $MYVAR
# echo "the variable is set" | sed 's/set/$MYVAR/'
the variable is $MYVAR
```

Weird things can happen when parsing data/text that contains quote characters:

```
# MYVAR='Don't worry. It's ok.'; echo $MYVAR
>
# you need to press Ctrl-C to abort
# MYVAR="Don't worry. It's ok."; echo $MYVAR
Don't worry. It's ok.
```

Expanding and Escaping

You already learned how to expand a variable such that its value is used instead of its name:

```
# export MYVAR=123
# echo "the variable is $MYVAR"
the variable is 123
```

“Escaping” a variable is the opposite, ensuring that the literal variable name is used instead of its value:

```
# export MYVAR=123

# echo "the \$MYVAR variable is $MYVAR"
the $MYVAR variable is 123
```

Note: The “escape character” is usually the backslash “\”.

1.2 Command Line Exercises

1.2.1 TAR & GZIP

1. Use gzip to compress the file P12931.txt
2. Decompress the resulting file P12931.txt.gz (revert previous command)
3. Use tar to create an archive containing all fasta files in the current directory into an archive called “fastafiles.tar”
4. Use gzip to compress the archive “fastafiles.tar”
5. How can you achieve the two previous steps “using tar to create archive” and “gzip the archive” in one command?
6. Test (list the contents of) the compressed archive “fastafiles.tar.gz”

7. Download the compressed PDB file for entry 1Y57 from rcsb.org (eg. `wget "http://www.rcsb.org/pdb/files/1Y57.pdb.gz"`) and decompress it.

1.2.2 GREP

1. Which of the DNA files ENST0* contains "TATATCTAA" as part of the sequence?
2. List only the names of the DNA files ENST0* that contain "CAACAAA" as part of the sequence.
3. Considering the previous example, would you consider grep a suitable tool to perform motif searches? Why not? Try to find the pattern "CAACAAA" by manual inspection of the first two lines of each sequence.
4. Count the number of ATOMs (lines starting with "ATOM") in the file 1Y57.pdb.
5. Does this number agree with the annotated number of atoms (Search the REMARKs for "protein atoms")

1.2.3 SED

1. Use sed to print only those lines that contain "version" in the files P05480.txt and P04062.txt
2. Use sed to change the text "sequence version 3" to "sequence version 4" in the files P05480.txt and P04062.txt (without actually changing the files, just printing)
3. Use sed to update the text "sequence version 3" to "sequence version 4" in the files P05480.txt and P04062.txt (this time, make the changes directly in the files)
4. Replace (transliterate) all occurrences of "r" by "l" and "l" by "r" (at the same time) in the file PROTEINS.txt (so that "structural" becomes "stluctular")

1.2.4 AWK

1. Use awk to print only those lines that contain "version" in the files P12931.txt and P05480.txt and think about how this procedure is different to sed.
2. For all FASTA files that begin with "P" ("P*.fasta") print only the second item of the header (split on "|") eg. for ">sp|P12931|SRC_HUMAN Proto-oncogene", print only "P12931"
3. The file "P12931.csv" contains phosphorylation sites in the protein P12931. (If the file "P12931.csv" does not exist, use `wget http://phospho.elm.eu.org/byAccession/P12931.csv` to download it).
 - (a) Column three of this file lists the amino acid position of the phosphorylation site. You are only interested in position 17 of the protein. Try to use "grep" to filter out all these lines containing "17".
 - (b) Now use awk to show all lines containing "17".
 - (c) Next try show only those lines where column three equals 17 (Hint: The file is semicolon-separated...).
 - (d) Finally print the PMIDs (column 6) of all lines that contain "17" in column 3.

1.2.5 Quoting and Escaping

1. Familiarize yourself with quoting and escaping.
1. Run the following commands to see the difference between single and double quotes when expanding variables:
2. Next, use ssh to login to a different machine to run the same command there, again using both quoting methods:
2. Closely inspect the results; is that what you were expecting? Discuss this with your neighbour.

I/O Redirection

Three IO “channels” are available by default:

- **Standard input (STDIN, Number: 0):** The input for your program, normally your keyboard but can be an other program (when using pipes or IO redirection)
- **Standard output (STDOUT, Number: 1):** Where your program writes its regular output to. Normally your terminal
- **Standard error (STDERR, Number: 2):** Where your programs normally write their error message to. Normally your terminal

Input, output and error messages can be redirected from their default “targets” to others. If using the file descriptor numbers (0, 1, 2) in redirections, then there must be no whitespace between the numbers and the redirection operators.

Hint: Redirect to `/dev/null` to discard the output of any command

Write the output of *cmd* into *afile*. This will **overwrite** *afile*:

```
$ cmd > afile
```

Write the output of *cmd* into *afile*. This will **append** to *afile*:

```
$ cmd >> afile
```

Discard the output of *cmd*

```
$ cmd > /dev/null
```

Write the output of *cmd* into *afile* (overwriting *afile*!) and write STDERR to the same place:

```
$ cmd > afile 2>&1
```

Append the output and error messages of *cmd* to *afile*:

```
$ cmd >> afile 2>&1
```

Same as above:

```
$ cmd > afile 2> afile
```

Append the output of *cmd* to *afile* and discard error messages:

```
$ cmd >> afile 2>/dev/null
```

Three times the same: Discard output and error messages completely:

```
$ cmd > /dev/null 2>&1  
$ cmd > /dev/null 2>/dev/null  
$ cmd >& /dev/null
```

Use output of *cmd2* as standard input for *cmd1*:

```
$ cmd1 < cmd2
```

See also

- [Bash One-Liners Explained, Part III: All about redirections](#) ¹
- [Bash Redirections Cheat Sheet](#) ²
- [Redirection Tutorial](#) ³

¹ <http://www.catonmat.net/blog/bash-one-liners-explained-part-three>

² <http://www.catonmat.net/blog/bash-redirections-cheat-sheet>

³ http://wiki.bash-hackers.org/howto/redirection_tutorial

The shell knows two types of variables: “Local” *shell* variables and “global” exported *environment* variables. By convention, environment variables are written in uppercase letters.

Shell variables are **only available to the current shell** and not inherited when you start an other shell or script from the commandline. Consequently, these variables will not be available for your shellscripts.

Environment variables are **passed on** to shells and scripts started from your current shell.

3.1 Setting, Exporting and Removing Variables

Variables are set (created) by simply assigning them a value

```
$ MYVAR=something
```

There must be no whitespace around the equal sign.

To create an environment variable, `export` is used. You can either export while assigning a value or in a separate step. Both of the following procedures are equivalent:

```
1. $ export MYGLOBALVAR="something else"
```

```
2. $ MYGLOBALVAR="something else"  
   $ export MYGLOBALVAR
```

Note: There is no `$` in front of the variable: To reference the variable itself (not its content) the name is used without `$`

Variables are removed with `unset`:

```
$ unset MYVAR
```

Note: Assigning a variable an empty value (i.e. `MYVAR=`) will *not* remove it but simply set its value to the empty string!

3.2 Listing Variables

You can list all your current environment variables with `env` and all shell variables with `set`. The list of shell variables will also contain all environment variables

3.3 Variable Inheritance

Only environment variables will be available in shells and scripts started from your current shell. However in shell commands run in subshells (i.e. commands run within round brackets) also local (shell) variables of your current shell are available.

3.4 Examples

Consider the following small shellscript *vartest.sh*:

```
#!/bin/sh
echo $MYLOCALVAR
echo $MYGLOBALVAR
echo -----
```

We will use it in the following examples to illustrate the various variable inheritances:

1. Set the variables and run the script i.e. in a new shell:

```
$ export MYGLOBALVAR="I am global"
$ MYLOCALVAR="I am local"
$ ./vartest.sh

I am global
-----
```

2. "source" the script, i.e. run it within your current shell:

```
$ ./vartest.sh
I am local
I am global
-----
```

3. Access the variables in a subshell:

```
$ (echo $MYGLOBALVAR; echo $MYLOCALVAR)
I am global
I am local
```

Basic Shell Scripting

4.1 What is a Script?

A script is nothing else than a number of shell command place together in a file. The simplest script is maybe just a complex oneliner that you don't want to type each time again. More complex scripts are seasoned with control elements (conditions and loops) which allow for a sophisticated command flow. scripts might allow for configuration and customization, thus allowing one script to be flexibly used in several different environments. Whatever you do in a script, you can also do on the commandline. This is also the first way to test your scripts step by step!

4.2 Script Naming and Organization

It is good practice - though not technically required - to give your scripts an extension which specifies their type. I.e. `".sh"` for Bourne Shell and Bourne Again Shell scripts, `".csh"` for C-Shell scripts. Sometimes `".bash"` for Bourne Again Shell scripts is used.

We recommend to either store all scripts in one location (e.g. `~/bin`) and add this location to your `$PATH` variable or to store the scripts together with the files that are processed by the script.

Hint: If you use scripts to process data, then the scripts should probably be archived together with the data files!

4.3 Running a Script

There are basically three ways to run a script:

1. the location to your script is not in your `$PATH` variable, then you have to specify the full path to the script:

```
$ /here/is/my/script.sh
[...]  
$
```

2. the location to the script is in the `$PATH` variable, then you can simply type its name:

```
$ script.sh
[...]  
$ man
```

In both situations, the script will need to have execute permissions to be run. If for some reason you can only read but not execute the script, then it can still be run in the following way:

3. specifying the interpreter (i.e. the program required to run the script). For shellscripts this is the appropriate shell). The full path (relative or absolute) to the script has to be provided in this case, no matter whether the script location is already contained in `$PATH` or not:

```
$ /bin/sh /here/is/my/script.sh
[...]  
$
```

4.3.1 Basic Structure of a Shellscript

Shellscripts have the following general structure:

- A line starting with “`#!`” which defines the interpreter. This line is called the *shebang line* and must be the first line in a script.
- A section where the configuration takes place, e.g. paths, options and commands are defined and it is made sure, that all prerequisites are met.
- A section where the actual processing is done. This includes error handling.
- A controlled exit sequence, which includes cleaning up all temporary files and returning a sensible exit status.

This is merely a recommendation to keep your scripts well structured. None of these sections are mandatory.

4.3.2 Readability and Documentation

Make your script easily readable. Use comments and whitespace and avoid super compact but hard to understand command lines. Always take into account that not only the shell, but also human beings will probably have to read and understand your script. (see [Breaking up long lines](#) (page 28)) Even if your script is very simple - document it! This helps others understand what you did, but - most important - it helps you remember what you did, when you have to reuse the script in the future.

Documentation is done either by writing comments into the script or by creating a special documentation file (`README.txt` or similar). Documenting in the script can be done in several ways:

- A preamble in the script, outlining the purpose, parameters and variables of the script as well as some information about authorship and and perhaps changes.
- Within the script as blocks of text or “End of line” comments.

To write a comments use the hash sign (" #"). Everything after a "#" is ignored when executing a script.

4.3.3 Anatomy of a Shellsript

Let's have a look at the following script, breaking it down into individual parts. First, the full script:

```
#!/bin/sh
```

Shebang line

```
#
# myscript.sh
#
# General purpose script for extracting Glycine
# occurrences in a datafile.
#
# Usage: myscript.sh datafile
#
# Exit values: 1: No datafile given or file
#               doesn't exist
#               2: No Glycine found
#
# Author: Me, myself and I
# Date:   Heidelberg, December 12., 2012
#
```

Preamble with a short description, usage information, authorship etc. etc.

```
# --- Configuration ---
GREPCMD=/bin/grep
DATAFILE=$1
```

Configuration

```
# --- Check prerequisites ---
# first check for $1
if [ -z $DATAFILE ]
then
    echo "No datafile given" 1>&2 # print on STDERR
    echo "USAGE: $0 datafile"
    exit 1
fi

# then check if the file exists
if [ ! -f $DATAFILE ]
then
    echo "Datafile $DATAFILE does not exist!" 1>&2
    exit 1
fi
```

Checking prerequisites and sane environment

```
# --- Now processing---
$GREPCMD -q Glycine $DATAFILE # Where is Glycine?
```

This is what you actually wanted to do

```
# --- Exit ---
if [ $? -eq 0 ]
then
    exit 0
else
    exit 2
fi
```

Ensure a valid and meaningful exit status

You can see from this example, that very often the actual computation is only a small part of the code. The rest of the scripts deal with prerequisites, error handling, user dialogue, exit status etc. etc.

4.3.4 Reporting Success or Failure - The Exit Status

Commands report their success or failure by their exit status. An exit status of 0 (zero) indicates success(!), while any exit status greater than 0 indicates an error. Some commands report more than one error status. Refer to the respective manpages to see the meanings of the different exit status. The exit status of a script is usually the exit status of the last executed command, which is reported by the environment variable `$?`:

`$?`: The exit status of the last run command

See *Ensuring a Sensible Exit Status* (page 27) about how to control the exit status of your script.

4.3.5 Command Grouping and Sequences

Commands can be concatenated to be executed one after the other unconditionally or based on the success of the respective previous command:

`cmd1; cmd2` – Execute commands in sequence

Example: Create a directory and change into it:

```
$ pwd
/home/fthommen
$ mkdir a; cd a
$ pwd
/home/fthommen/a
$
```

`cmd1 && cmd2` – Execute `cmd2` only if `cmd1` was successful:

Example: Confirm that `/etc` exists:

```
$ cd /etc && echo "/etc exists"
/etc/exists
$
```

`cmd1 || cmd2` – Execute `cmd2` only if `cmd1` was not successful:

Example: Warn if a directory doesn't exist:

```
$ cd /etc || echo "/etc is missing!"
$ cd /nowhere >&/dev/null || echo "/nowhere does not exist"
/nowhere does not exist
$
```

`(cmds)` – Group commands to create one single output stream: The commands are run in a subshell (i.e. a new shell is opened to run them)

Example: Change into `/etc` and list content. You are still in the same directory as you were before:

```
$ pwd
/home/fthommen
$ (cd /etc; ls)
[... directory listing here ...]
```

```
$ pwd
/home/fthommen
$
```

{ cmds; } – Group commands to create one single output stream: The commands are run in the current (!) shell.

Note: The opening “{” must be followed by a blank and the last command must be succeeded by a *semicolon* (“;”)

Example: Change into /etc and list its content. You are still in /etc after the bracketed expression (compare to the example above):

```
$ pwd
/home/fthommen
$ { cd /etc; ls; }
[... directory listing here ...]
$ pwd
/etc
$
```

4.4 Control Structures

The following syntax elements will be described for sh/bash *and* for csh/tcsh. However since this course is mainly about sh/bash, examples will only be given for sh/bash. Some notes about csh/tcsh specialities might be given in the text. This is only a selection of the most useful or most common elements. There are much more in the manpages. All shells offer myriads of possibilities which cannot possibly be demonstrated in this course. Some of the described features might be specific to bash and not be available in a classical Bourne Shell on other systems.

4.4.1 Conditional Statements

if - then - else

if - then - else is the most basic conditional statement: Do something depending on certain conditions. Its basic syntax is:

sh/bash	csh/tcsh
<pre>if condition1 then commands elif condition2 more commands [...] else even more commands fi</pre>	<pre>if (condition) then commands else if (condition2) then more commands [...] else even more commands endif</pre>

Conditions can be either the **exit status of a command** or the **evaluation of a logical or arithmetic expression**:

1. Evaluating the exit status of a command: Simply use the command as condition. For example:

```
if grep -q root /etc/passwd
then
    echo root user found
else
    echo No root user found
fi
```

Note: In *csh/tcsh*

1. To evaluate the exit status of a command in it must be placed within curly brackets with blanks separating the brackets from the command: `if ({ grep -q root /etc/passwd }) then [...]`
2. Redirection of commands in conditions does not work

Hint: Redirect the output of the command to be evaluated to `/dev/null` if you are only interested in the exit status and if the command doesn't have a "quiet" option.

2. Evaluating of conditions or comparisons:

Conditions and comparisons are evaluated using a special command `test` which is usually written as `[]` (no joke!). As `[]` is a command, it must be followed by a blank. As a speciality the `[]` command must be ended with `""]""` (note the preceding blank here)

Note: In *csh/tcsh* the `test` (or `[]`) command is not needed. Conditions and comparisons are directly placed within the round braces.

sh/bash	File condition	cshtcsh
-e <i>file</i>	<i>file</i> exists	-e <i>file</i>
-f <i>file</i>	<i>file</i> exists and is a regular <i>file</i>	-f <i>file</i>
-d <i>file</i>	<i>file</i> exists and is a directory	-d <i>file</i>
-r <i>file</i>	<i>file</i> exists and is readable	-r <i>file</i>
-w <i>file</i>	<i>file</i> exists and is writeable	-w <i>file</i>
-x <i>file</i>	<i>file</i> exists and is executable	-x <i>file</i>
-s <i>file</i>	<i>file</i> exists and has a size > 0	
	<i>file</i> exists and has zero size	-z <i>file</i>
	String Comparison	
-n <i>s1</i>	String <i>s1</i> has non-zero length	<i>s1</i> == <i>s2</i>
-z <i>s1</i>	String <i>s1</i> has zero length	<i>s1</i> != <i>s2</i>
<i>s1</i> = <i>s2</i>	Strings <i>s1</i> and <i>s2</i> are identical	
<i>s1</i> != <i>s2</i>	Strings <i>s1</i> and <i>s2</i> differ	
<i>string</i>	String <i>string</i> is not null	
	Integer Comparison	
<i>n1</i> -eq <i>n2</i>	<i>n1</i> equals <i>n2</i>	<i>n1</i> == <i>n2</i>
<i>n1</i> -ge <i>n2</i>	<i>n1</i> is greater than or equal to <i>n2</i>	<i>n1</i> >= <i>n2</i>
<i>n1</i> -gt <i>n2</i>	<i>n1</i> is greater than <i>n2</i>	<i>n1</i> > <i>n2</i>
<i>n1</i> -le <i>n2</i>	<i>n1</i> is less than or equal to <i>n2</i>	<i>n1</i> <= <i>n2</i>
<i>n1</i> -lt <i>n2</i>	<i>n1</i> is less than <i>n2</i>	<i>n1</i> < <i>n2</i>
<i>n1</i> -ne <i>n2</i>	<i>n1</i> is not equal to <i>n2</i>	<i>n1</i> != <i>n2</i>
	Combination of conditions	
! <i>cond</i>	True if condition <i>cond</i> is not true	! <i>cond</i>
<i>cond1</i> -a <i>cond2</i>	True if conditions <i>cond1</i> and <i>cond2</i> are both true	<i>cond1</i> && <i>cond2</i>
<i>cond1</i> -o <i>cond2</i>	True if conditions <i>cond1</i> or <i>cond2</i> is true	<i>cond1</i> <i>cond2</i>

Examples: Test for the existence of /etc/passwd:

```
if [ -e /etc/passwd ]
then
    echo /etc/passwd exists
else
    echo /etc/passwd does NOT exist
fi
```

or:

```
if test -e /etc/passwd
then
    echo /etc/passwd exists
else
    echo /etc/passwd does NOT exist
fi
```

Note: Bash supports an additional way of evaluating conditional expressions with `[[expression]]`. This syntax element allows for more readable expression combination and handles empty variables better. However it is not backwards compatible with the original Bourne Shell. See the bash manpage for more infor-

mation

case

The case statement implements a more compact and better readable form of if - elif - elif - elif etc. Use this if your variable (you can *only* check for variables with case) can have a distinct number of valid values. A typical usage of case will follow later.

The basic syntax is:

sh/bash	csh/tcsh
<pre>case variable in pattern1) commands ;; pattern2) commands ;; *) commands ;; esac</pre>	<pre>switch (variable) case pattern1: commands breaksw case pattern2: commands breaksw default: commands endsw</pre>

Note: for the patterns “*”, “?” and “[...]” can be used

Note: The “*)” (sh/bash) and “default:” (csh/tcsh) patterns are “catch-all” patterns which match everything not matched above. It is often used to detect invalid values of variable.

Note: Multiple patterns can be handled by separating them with “|” in sh/bash or by successive case statements in csh/tcsh.

Example: Check if /opt/ or /usr/ paths are contained in \$PATH::

```
case $PATH in
  */opt/* | */usr/* )
    echo /opt/ or /usr/ paths found in \ $PATH
    ;;
  *)
    echo '/opt and /usr are not contained in $PATH'
    ;;
esac
```

4.4.2 Loops

for / foreach

The for and foreach statements respectively will loop through a list of given values and run the given statements for each run:

sh/bash	csh/tcsh
<pre>for variable in list do commands done</pre>	<pre>foreach variable (list) commands end</pre>

list is a list of strings, separated by whitespaces

Examples: List all files in /tmp in a bulleted list:

```
for FILE in /tmp/*
do
    echo " * $FILE"
done
or
for FILE in `ls /tmp`
do
    echo " * $FILE"
done
```

while / until

The while and until loops execute your commands while (or until respectively) a certain condition is met:

sh/bash	csh/tcsh
<pre>while condition do commands done until condition do commands done</pre>	<pre>while (condition) commands end</pre>

The conditions are constructed the same way as those used in if statements.

Note: The until statement is not available in csh/tcsh.

“Manual” loop control

Instead of (or additionally to) the built-in loop control in for/foreach, while and until loops, you can control exiting and continuing them with `break` and `continue`: `break` “breaks out” of the innermost loop (loops can be nested!) and continues after the end of the loop. `continue` skips the rest of the current (innermost) loop and starts the next iteration

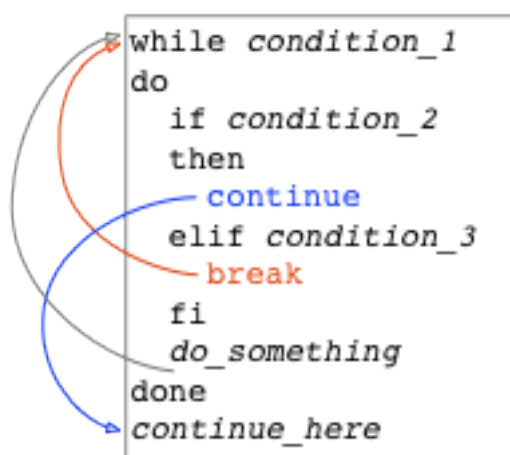





Figure 4.1: Loop control

Symbol	
	Regular loop cycle
	break due to <i>condition_2</i>
	continue due to <i>condition_3</i>

4.5 Making Scripts Flexible

Scripts are most useful, if they can be reused. Copying scripts and changing them to fit the new situation is time-consuming and error-prone. Additionally if you add an improvement to the current script, then all previous versions will stay without it. Having one script with the possibility to configure it, is usually the better way. Customization of scripts can be achieved by either using variables or by adding the possibility to use your own commandline options and arguments.

4.5.1 Configurable Scripts

Any value - be it paths, commands or options - that is specific to individual applications or your script, should not be “hardcoded” (i.e. used literally within the script) but assigned to variables:

Using Variables

Any value - be it paths, commands or options - that is specific to individual applications or your script, should not be hardcoded (i.e. used literally within the script). Instead you should use variables to refer to them:

Bad example: You have to change two instances of the path each time you want to list an other directory:

```
#!/bin/sh

echo "The directory /etc contains the following files:"
ls /etc
```

Good example: The path is now in a variable and only one instance has to be changed each time (less work, less errors):

```
#!/bin/sh

MYDIR=/etc

echo "The directory $MYDIR contains the following files:"
ls $MYDIR
```

Of course, you'll still have to modify the script each time you want to list the content of an other directory. A more flexible way of customization would be to use a settings file.

Using a Settings File

Instead of having your configurable section within the script, it can be "outsourced" to its own file. This file is basically a shellscript which is run within the primary script. To run commands from a file within the current environment, the commands `source` (bash, csh/tcsh) or `.` (dot) (sh/bash) are used:

The settings file, e.g. settings.ini:

```
MYDIR=/etc
```

The script:

```
#!/bin/sh

. ./settings.ini

echo "The directory $MYDIR contains the following files:"
ls $MYDIR
```

4.5.2 Defining your own Commandline Options and Arguments

The best way to configure a script is to allow for your own commandline options and arguments. Commandline arguments are available the script as so-called positional parameters `$1`, `$2`, `$3`: etc. `$0`: contains the name of the script. The variables important when dealing with commandline parameters are:

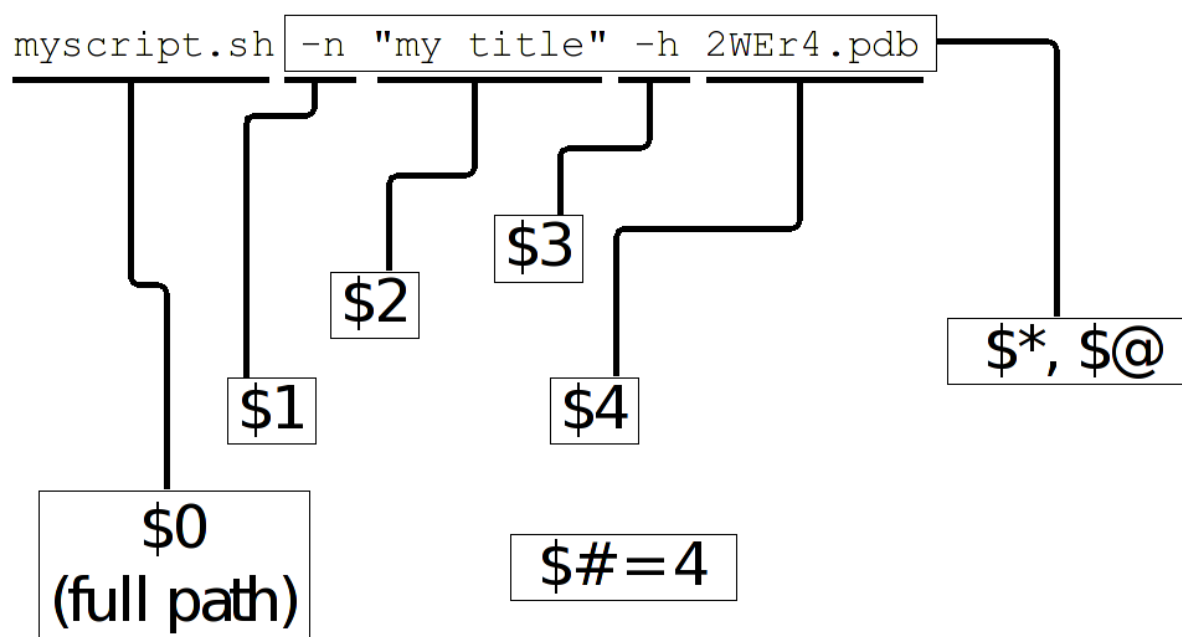
\$0: path to the script. Either the path as you specified it or the full path if the script was executed through `$PATH`

\$1, \$2, \$3, etc: Positional parameters (i.e. commandline arguments)

\$#: Current number of positional parameters

\$*: All positional parameters. If used within double quotes ("`$*`"), then it will expand to the list of all positional parameters, where the complete list is quoted

@: All positional parameters. If used within double quotes ("`@`"), then it will expand to the list of all positional parameters, where each parameter is individually quoted



```
"$@" = "-a" "my title" "-h" "2WEr4.pdb"
"$*" = "-a my title -h 2WEr4.pdb"
```

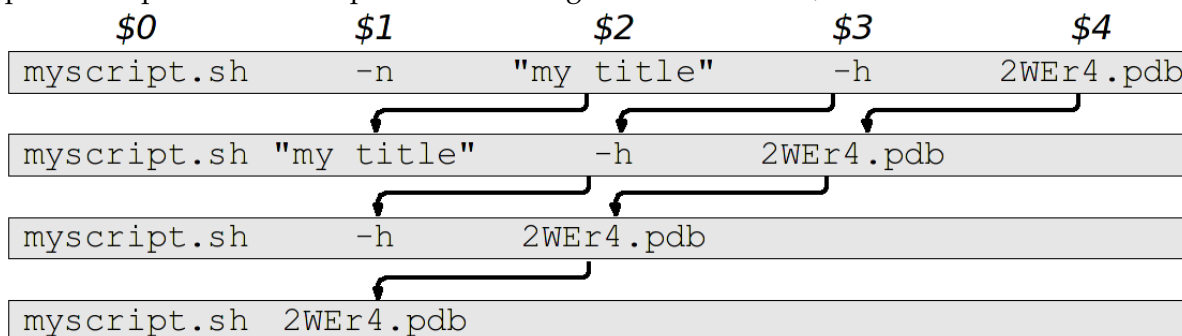
If you run the script

```
#!/bin/sh
echo The script is $0
echo The first cmdline option is $1
echo The second cmdline option is $2
```

with two arguments, you'll get the following output:

```
$ ./script.sh ABC DEF
The script is ./script.sh
The first cmdline option is ABC
The second cmdline option is DEF
$
```

In many cases you'll not know how many parameters are given on the commandline. In these cases you can use `shift` to loop through them. `shift` removes `$1` and moves all other positional parameters one position to the right: `$2` becomes `$1`, `$3` becomes `$2` etc.:



With the help of “\$#”, “shift”, “case” and the positional parameters we can now check all the commandline parameters:

```
while [ "$#" -gt 0 ]
do
  case $1 in
    -h) echo "Sorry, no help available!" # not very helpful, is it?
        exit 1                         # exit with error
        ;;

    -v) VERBOSE=1                      # we may use $VERBOSE later
        ;;

    -f) shift
        FILE=$1                       # Aha, -f requires an
                                     # additional argument
        ;;

    *)  echo "Wrong parameter!"
        exit 1                         # exit with error
  esac
  shift
done
```

4.6 Ensuring a Sensible Exit Status

If you don't provide your own exit status, then the script will return the exit status of the last executed command (See [Reporting Success or Failure - The Exit Status](#) (page 18)). In many cases this might be what you want, but very often it isn't. Consider the following script which is a real example from real life and happened to me personally:

```
#!/bin/sh

[... do something that fails ...]

echo "End of the script"
```

This script will *always* succeed, as the echo command hardly ever fails. You will - from the exit status of the script - never be able to detect that something went wrong. Instead in such cases you should manually handle the exit codes of the commands that are run within the script.

With it's help we can keep track of the exit status of all our important processing steps and finally return a sensible value:

```
#!/bin/sh
mystatus=0;

[... do something that might fail ...]
if [ $? -ne 0 ]
then
  mystatus=1
fi
```

```
[... do something else that might fail, too ...]
[ $? -ne 0 ] && mystatus=1          # same as above.  Do you understand
                                   # this?

echo "End of the script"
exit $mystatus
```

4.6.1 Why is the exit status important after all?

First when you use your script within other scripts, you'll probably need to be able to check, if it has succeeded. There might be other ways (e.g. checking outputfiles for certain strings, checking directly the textual output of the script etc.), but these ways are usually cumbersome and require lots of coding. Exit values are easy to check. Second: Other tools and systems might also use the exit status of your script. E.g. the cluster system uses your job's exit status to assess, if it has run successfully or not. Returning success even in case of failure will result in lots of complications in case a problem occurs. It took me several days to realize the bug above.

4.7 Tips and Tricks

4.7.1 Combining Variables with other Strings

When combining variables with other strings, then in some situations the variable name must be placed in curly brackets ("{}"):

```
$ A=Heidel
$ echo $Aberg

$ echo ${A}berg
Heidelberg
$
```

4.7.2 Filenames and Paths

If possible, try to avoid any special characters (blanks, semicolons (";"), colons (":"), backslashes ("") etc.) in file and directory names. All these special characters can lead to problems in scripted processing. Instead, stick to alphanumeric characters (a-z, 0-9), dots ("."), dashes ("-") and underscores ("_"). Additionally sticking to lowercase characters helps avoiding mistypes and makes the automatic filename expansion easier.

4.7.3 Breaking up Long Code Lines

Code lines can become pretty long and unreadable, wrapping onto the next line etc. You can use the escape character (backslash, "\") to break them up and enhance readability of your script. The escape character must immediately be followed by a newline (no intermediate blanks or other is allowed):


```
$ bsub -o output.log -e error.log -q clngnew -M 150000 -R "select[(mem > 15000)]" /g/so
```

becomes:

```
$ bsub -o output.log \
      -e error.log \
      -q clngnew \
      -M 150000 \
      -R "select[(mem > 15000)]" \
      /g/software/bin/pymol-1.4 -r -p < pymol.pml
```

Which is way better to read and to maintain

4.7.4 Script Debugging

sh/bash and csh/tcsh have both an option “-x” which helps debugging a script by echoing each command before executing it. This option can be set and unset during runtime with `set -x` / `set +x` (sh/bash) and `set echo` / `unset echo` (csh/tcsh).

4.7.5 Command Substitution

You can use the output of a command and assign it to a variable or use it right away as text string, by using the command substitution operator “`” (backticks, backquotes) or “\$(...)”. The backtick operator works in all shells, while “\$(...)” only works in bash.

Three variants for the same (print out who you are in English text):

```
$ ME='whoami'
$ echo I am $ME
I am fthommen
$

$ ME=$(whoami)
$ echo I am $ME
I am fthommen
$

$ echo I am `whoami`
I am fthommen
$
```

4.7.6 Create Temporary Files

You can create temporary files with `mktemp`. By default it will create a new file in `/tmp` and print its name:

```
$ mktemp
/tmp/tmp.Yaafh19370
$
```

4.7.7 Cleaning up Temporary Files

It is considered good practice and sometimes even important, to clean up temporary data before ending a script. A simple way - which will not cover all cases, though - could be to store all created temporary files in a variable and remove them all before exiting the script:

```
#!/bin/sh
ALL_TEMPFILES=""      # store a list of all temporary files here

TEMPFILE1=`mktemp`
ALL_TEMPFILES="$ALL_TEMPFILES $TEMPFILE1"

TEMPFILE2=`mktemp`
ALL_TEMPFILES="$ALL_TEMPFILES $TEMPFILE2"

[... process, process, process ...]

rm -f $ALL_TEMPFILES
exit
```

Solutions to the Exercises

5.1 TAR & GZIP

1. Use gzip to compress the file P12931.txt

```
$ gzip P12931.txt
```

2. Decompress the resulting file P12931.txt.gz (revert previous command)

```
$ gunzip P12931.txt.gz
```

or

```
$ gzip -d P12931.txt.gz
```

3. Use tar to create an archive containing all fasta files in the current directory into an archive called "fastafiles.tar"

```
$ tar -c -f fastafiles.tar *.fasta
```

4. Use gzip to compress the archive "fastafiles.tar"

```
$ gzip fastafiles.tar
```

5. How can you achieve the two previous steps "using tar to create archive" and "gzip the archive" in one command?

```
$ tar -c -z -f fastafiles.tar.gz *.fasta
```

Note: Note the -z

6. Test (list the contents of) the compressed archive "fastafiles.tar.gz"

```
$ tar -tf fastafiles.tar.gz
```

7. Download the compressed PDB file for entry 1Y57 from rcsb.org (eg. `wget "http://www.rcsb.org/pdb/files/1Y57.pdb.gz"`) and decompress it.

```
$ wget "http://www.rcsb.org/pdb/files/1Y57.pdb.gz"
$ gunzip 1Y57.pdb.gz
```

5.2 GREP

1. Which of the DNA files ENST0* contains "TATATCTAA" as part of the sequence?

```
$ grep "TATATCTAA" ENST0*

ENST00000380152.fasta:ACGGAAGAATGTGAGAAAAATAAGCAGGACACAATTACAATAAAAAATATATCTAA
ENST00000544455.fasta:ACGGAAGAATGTGAGAAAAATAAGCAGGACACAATTACAATAAAAAATATATCTAA
```

2. List only the names of the DNA files ENST0* that contain "CAACAAA" as part of the sequence.

```
$ grep "CAACAAA" ENST0*

ENST00000380152.fasta
ENST00000544455.fasta
```

3. Considering the previous example, would you consider grep a suitable tool to perform motif searches? Why not? Try to find the pattern "CAACAAA" by manual inspection of the first two lines of each sequence.

Note: Answer: When using grep as a motif searching tool, you need to keep in mind that grep (like sed and awk) is line-oriented, meaning that by default it only searches for a given motif in a single line. In the given example, upon manual inspection you will find the given motif also in the file ENST00000530893.fasta, which grep missed. You would need to think about how to do multi-line searches (eg. Removing line-breaks etc.)

4. Count the number of ATOMs (lines starting with "ATOM") in the file 1Y57.pdb.
5. Does this number agree with the annotated number of atoms (Search the REMARKs for "protein atoms")

```
$ grep -c "ATOM" 1Y57.pdb
3632
$ grep -i "protein atoms" 1Y57.pdb
REMARK    3      PROTEIN ATOMS                : 3600
```

This means there are 3600 atoms annotated in this PDB file, however we counted 3632. This is because grep also counted any occurrence of "ATOM" within REMARKS. We can avoid this by either filtering out the remarks:

```
$ grep -v REMARK 1Y57.pdb | grep -c ATOM
3600
```

...or by telling grep to only count those lines that start with "ATOM":

```
$ grep -c ATOM 1Y57.pdb
3600
```

5.3 SED

1. Use sed to print only those lines that contain “version” in the files P05480.txt and P04062.txt

```
$ sed '/version/p' P05480.txt P04062.txt
```

2. Use sed to change the text “sequence version 3” to “sequence version 4” in the files P05480.txt and P04062.txt (without actually changing the files, just printing)

```
$ sed 's/sequence version 3/sequence version 4/' P05480.txt P04062.txt
```

3. Use sed to update the text “sequence version 3” to “sequence version 4” in the files P05480.txt and P04062.txt (this time, make the changes directly in the files)

```
$ sed -i.bak 's/sequence version 3/sequence version 4/' P05480.txt P04062.txt
```

4. Replace (transliterate) all occurrences of “r” by “l” and “l” by “r” (at the same time) in the file PROTEINS.txt (so that “structural” becomes “stluctular”)

```
$ sed 'y/rRlL/lLrR/' PROTEINS.txt
```

5.4 AWK

1. Use awk to print only those lines that contain “version” in the files P12931.txt and P05480.txt and think about how this procedure is different to sed.

```
$ awk '/version/ {print}' P12931.txt P05480.txt
```

This is very similar to sed, you also have to use the slashes “/” to define the search pattern. However the sed notation is a little more concise...

2. For all FASTA files that begin with “P” (“P*.fasta”) print only the second item of the header (split on “|”) eg. for “>sp|P12931|SRC_HUMAN Proto-oncogene”, print only “P12931”

```
$ awk -F'|' ' />/ {print $2}' P*.fasta
```

3. The file “P12931.csv” contains phosphorylation sites in the protein P12931. (If the file “P12931.csv” does not exist, use wget <http://phospho.elm.eu.org/byAccession/P12931.csv> to download it).

- (a) Column three of this file lists the amino acid position of the phosphorylation site. You are only interested in position 17 of the protein. Try to use “grep” to filter out all these lines containing “17”.

```
$ grep 17 P12931.csv
```

- (b) Now use awk to show all lines containing “17”.

```
$ awk '/17/ {print}' P12931.csv
```

- (c) Next try show only those lines where column three equals 17 (Hint: The file is semicolon-separated...).

```
$ awk -F';' '$3==17 {print}' P12931.csv
```

- (d) Finally print the PMIDs (column 6) of all lines that contain “17” in column 3.

```
$ awk -F';' '$3==17 {print $6}' P12931.csv
```

5.5 Quoting and Escaping

1. Familiarize yourself with quoting and escaping.

1. Run the following commands to see the difference between single and double quotes when expanding variables:

```
$ echo "$HOSTNAME"
...
$ echo '$HOSTNAME'
```

2. Next, use ssh to login to a different machine to run the same command there, again using both quoting methods:

```
$ ssh pc-atcteach01 'echo $HOSTNAME'
...
$ ssh pc-atcteach01 "echo $HOSTNAME"
```

2. Closely inspect the results; is that what you were expecting? Discuss this with your neighbour.

6.1 Links and Further Information

6.1.1 Links

- A full 500 page book about the Linux commandline for free(!): [LinuxCommand.org](http://linuxcommand.org/) ¹
- Another nice introduction: “A beginner’s guide to UNIX/Linux” ²
- The “*commandline starter*” chapter of an O’Reilly book: [Learning Debian GNU/Linux - Issuing Linux Commands](http://oreilly.com/openbook/debian/book/ch04_01.html) ³
- A nice introduction to Linux/UNIX file permissions: “[chmod Tutorial](http://www.catcode.com/teachmod/)” ⁴
- [Linux Cheatsheets](http://www.cheat-sheets.org/#linux) ⁵
- For the technically interested: [Linux Filesystem Hierarchy Standard](http://www.pathname.com/fhs/) ⁶ and [Linux Standard Base](http://www.linuxfoundation.org/collaborate/workgroups/lsb) ⁷
- [Unix commands applied to bioinformatics](http://rous.mit.edu/index.php/Unix_commands_applied_to_bioinformatics) ⁸
- [BioPieces](http://code.google.com/p/biopieces) ⁹ are a collection of bioinformatics tools that can be pieced together in a very easy and flexible manner to perform both simple and complex tasks.
- [Google shell style guide](https://code.google.com/p/google-styleguide) ¹⁰
- [Useful bash one-liners for bioinformatics](https://github.com/stephenturner/oneliners) ¹¹
- Interactive explanation of your commandline: [Explain Shell](http://www.explainshell.com) ¹²

¹ <http://linuxcommand.org/>

² <http://www.mn.uio.no/astro/english/services/it/help/basic-services/linux/guide.html>

³ http://oreilly.com/openbook/debian/book/ch04_01.html

⁴ <http://www.catcode.com/teachmod/>

⁵ <http://www.cheat-sheets.org/#linux>

⁶ <http://www.pathname.com/fhs/>

⁷ <http://www.linuxfoundation.org/collaborate/workgroups/lsb>

⁸ http://rous.mit.edu/index.php/Unix_commands_applied_to_bioinformatics

⁹ <http://code.google.com/p/biopieces>

¹⁰ <https://code.google.com/p/google-styleguide>

¹¹ <https://github.com/stephenturner/oneliners>

¹² <http://www.explainshell.com>

6.1.2 Command Line Mystery Game

CLMystery¹³ is a game that you play on the commandline: There's been a murder in Terminal City, and TCPD needs your help to solve this crime *by using commandline tools only!*

To play the game, get the files from github and read the instructions:

```
wget https://github.com/veltman/clmystery/archive/master.zip
unzip master.zip
cd clmystery-master/
cat instructions
```

6.1.3 Real printed paper books

- Dietz, M., *"Praxiskurs Unix-Shell"*, O'Reilly (highly recommended!, german language only)
- Herold, H., *"awk & sed"*, Addison-Wesley
- Robbins, A., *"sed & awk Pocket Reference"*, O'Reilly
- Robbins, A. and Beebe, N., *"Classic Shell Scripting"*, O'Reilly
- Siever, E. et al., *"Linux in a Nutshell"*, O'Reilly

6.1.4 Live - CDs

A Live-CD is a complete bootable computer operating system which runs in the computer's memory, rather than loading from the hard disk drive. It allows users to experience and evaluate an operating system without installing it or making any changes to the existing operating system on the computer.

Just download an ISO-Image, burn it onto a CD/DVD and insert it into your DVD-Drive to boot your computer with Linux!

Fedora Live CD

This Live CD contains everything the **Fedora**¹⁴ Linux operating system has to offer and it's everything you need to try out Fedora - you don't have to erase anything on your current system to try it out, and it won't put your files at risk. Take Fedora for a test drive, and if you like it, you can install Fedora directly to your hard drive straight from the Live Media desktop.

Knoppix

Knoppix¹⁵ is an operating system based on Debian designed to be run directly from a CD / DVD or a USB flash drive, one of the first of its kind for any operating system. When starting a program, it is loaded from the removable medium and decompressed into a RAM

¹³ <https://github.com/veltman/clmystery>

¹⁴ <http://fedoraproject.org/wiki/FedoraLiveCD>

¹⁵ <http://knopper.net/knoppix>

drive. The decompression is transparent and on-the-fly. More than 1000 software packages are included on the CD edition and more than 2600 are included on the DVD edition. Up to 9 gigabytes can be stored on the DVD in compressed form.

BioKnoppix

Bioknoppix ¹⁶ is a customized distribution of Knoppix Linux Live CD. With this distribution you just boot from the CD and you have a fully functional Linux OS with open source applications targeted for the molecular biologist. Beside using RAM, Bioknoppix doesn't touch the host computer, being ideal for demonstrations, molecular biology students, workshops, etc.

Vigyaan

Vigyaan ¹⁷ is an electronic workbench for bioinformatics, computational biology and computational chemistry. It has been designed to meet the needs of both beginners and experts.

BioSlax

BioSLAX ¹⁸ is a live CD/DVD suite of bioinformatics tools that has been released by the resource team of the BioInformatics Center (BIC), National University of Singapore (NUS).

6.2 About Bio-IT

Bio-IT is a community project aiming to develop and strengthen the bioinformatics user community at EMBL Heidelberg. It is made up of members across the different EMBL Heidelberg units and core facilities. The project works to achieve these aims, firstly, by providing a forum for discussing and sharing information and ideas on computational biology and bioinformatics, focused on the **Bio-IT portal**. Secondly, we organise and participate in a range of different networking and social activities aiming to strengthen ties across the community.

6.2.1 Resources

A list of biocomputing-related resources associated with the project, in the top-left "Resources" menu, including, for example there is help provided for installing software on Linux computers at EMBL, instructions on using the Git versions control system server provided by EMBL, and various other kinds of information.

6.2.2 Training and Outreach

The "Training and Outreach" menu provides information on events (courses and conferences), both internal to EMBL and organised elsewhere by other organisations, that are related to biocomputing and bioinformatics

¹⁶ <http://bioknoppix.hpcf.upr.edu>

¹⁷ <http://www.vigyaan.cd.org>

¹⁸ <http://www.bioslax.com>

6.2.3 Networking

Several different kinds of networking events for the Bio-IT community are being organised, including beer sessions for the EMBL community, and within-Heidelberg events for the larger Heidelberg biocomputing community.

6.2.4 Biocomputing expertise at EMBL

You can use the Bio-IT portal to search for people working at EMBL who have experience working with data or tools you might be interested in.

If you've not yet got a page up on the portal describing your own expertise, please do so. If you need any help doing this, you can read about this in the portal's FAQ section, or get in touch with one of the site administrators.

6.2.5 Centers

EMBL Centres are 'horizontal', cross-departmental structures that promote innovative research projects across disciplines. All the EMBL Centres listed below have a strong computational component.



Biomolecular Network Analysis

The **CBNA** disseminates expertise, know-how and guidance in network integration and analysis throughout EMBL.

Statistical Data Analysis

The **CSDA** helps EMBL scientists to use adequate statistical methods for their specific technological or biological applications.

Molecular and Cellular Imaging

The **CMCI** makes your life in image processing/analysis easier and more fun.

Modeling

The **Centre for Biological Modeling (CBM)** aims to support people to adopt mathematical modeling techniques into their everyday research.

6.3 Acknowledgements

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