Software and Web-Security Assignment 1, Monday, February 2, 2015

Handing in your answers: Submission via Blackboard (http://blackboard.ru.nl)

Deadline: Monday, February 9, 24:00 (midnight)

Marks: You can score a total of 100 points.

- 1. Log into a Linux machine, either locally, or through ssh into lilo.science.ru.nl or stitch.science.ru.nl.
 - (a) Use the command line to create a directory called sws1-SNUMBER1-SNUMBER2 (replace SNUMBER1 and SNUMBER2 by your student numbers). In this directory, create 3 subdirectories called exercise1, exercise2, and exercise3.
 - (b) Write a text file called names.txt with your names and student S-numbers and place this file into the directory sws1-SNUMBER1-SNUMBER2.
 - (c) How large (in bytes) is the directory sws1-SNUMBER1-SNUMBER2? Write your answer to a file called 1b.txt and place this file in the exercise1 directory.
 - (d) Write all commands that you used for the other parts of this exercise to a text file called commands and place this file in the exercise1 directory.
- 2. Download the file http://cryptojedi.org/peter/teaching/genome.txt. You will see that it consists of 500 lines, each consisting of 100 characters, each of those characters being A,C,G, or T. See the content in this file as one long sequence of genome.
 - (a) Use Linux shell commands to find out how often the subsequence GATTACA is contained in this genome? Write your answer and how you found this answer into a file called 2a.txt. Place this file in the exercise2 directory.

Hint: Be careful, the sequence GATTACA may be spread over two lines in the file!

- (b) Write a shell script called genome.sh, which receives as first argument a filename and as second argument a string, and prints, how often the string appears in the file, also counting occurrences of the search string being spread over several lines. Place the script genome.sh into the exercise2 directory.
- (c) Write a shell (bash) script called gengenome.sh, which generates output that looks like the content of genome.txt, but with random choices of A,C,G, or T. Make sure that
 - the program generates new random output each time it is called;
 - the probability for each of the four letters at each position is 25%; and
 - the program prints exactly 500 lines of 100 characters each.

Place the script gengenome.sh into the exercise2 directory.

Note: It may be helpful to take a look at bash programming tutorials online.

- (d) Write a C program called parsegenome.c, which checks whether a file given as first command-line argument is of the format of the genome.txt file, i.e., whether it has exactly 500 lines with 100 characters (+ newline) each, where each of the 100 characters of each line is one of either A, C, G, or T. Let the program return -1 if the file does *not* have the correct format and 0 if the file has the correct format. Furthermore, if the file has the correct format, make sure that the program counts how often each of the 4 characters occurs and print these 4 counts to standard output. Place the file parsegenome.c into the exercise2 directory.
- (e) Write a Makefile that compiles parsegenome.c and place the Makefile into the exercise2 directory.
- 3. Generate a tar.gz archive of the whole sws1-SNUMBER1-SNUMBER2 directory. Submit this archive in Blackboard.