# Description of the project work for bioinformatic practicals

*3-6 pages long manuscript of 3 authors*

*Send:* 7 January 2019 as a **pdf** attachment of an email to [ELTEbioinfo@gmail.com](mailto:ELTEbioinfo@gmail.com). (If you send it earlier, we will correct it earlier.)

*Topic*: Description, comparison and summarization of certain properties of the 3 different proteins the authors analyzed during the semester.

*Description:* Write down the details of how you solved the below mentioned tasks and insert the results. Comment why you did certain steps and what is your opinion about the results. The sent document should be able to explain all above mentioned thing itself.

*Authors:* There should be three authors who have different proteins. (If necessary you can choose other proteins from this list: RASK, ERK1, JAK1, IGF1R, GSK3B, AXIN1, SMAD2, NOTCH1.) Write the name of the authors to the document.

***Tasks***:

1. Describe the proteins based on the basic descriptions you find in the Uniprot database. What are the unique and common properties of your proteins based on the Gene Ontology descriptions? (You can find the Gene Ontology categories of your proteins at Uniprot or at Ensembl Biomart.)
2. Draw a majority rule (not extended) consensus tree from the maximum-likelihood trees of the (enshortened) mRNA sequences of your proteins.[[1]](#footnote-1) (Species: human, chimp, dog, cow, mouse and opossum.) You can use the trees you calculated during the practicals or calculate new ones. Insert the trees to the document as figures. Use readable font size on the figures. Show the consensus values on the consensus tree (how many (or %) maximum-likelihood tree supported the branches). Discuss the the similarities and differences of trees.
3. Build 3 dot-plot figures from the genomic and the mRNA sequences of your proteins (human) with UGENE program. Paste the figures to your document. How many exons do you see on the dot-plots? Do these dot-plots have the real exon number of genes? Check the real exon numbers in NCBI. Are they equal with exons which you find in dot-plot figures?
4. Download the first neighbors of your proteins from two public databases using the Cytoscape software. Merge the networks. Discard all non-human proteins and all subgraphs that don’t connect to the greatest component (graph). Insert a figure in which the 3 proteins of yours are visually distinguishable from the others and from each other. Make their names readable (only for those three). Make the first neighbors and the common first neighbors of your proteins visually distinguishable too. Write a figure description for this.
5. Searching GSE21510 expression data on GEO2R webservice (<https://www.ncbi.nlm.nih.gov/geo/geo2r/>). Create two groups: stage1 és stage2. Assign each sample to the corresponding group based on the Stage és Tissue columns (Tissue column must contain "cancer"). Save the table and after that repeat the previous steps with Stage 3 and Stage 4. Download Affy HG U*133* PLUS *2* probeset table from Ensemble Biomart. Import all tables to Cytoscape (which has opened 3 proteins session from 4. task) and create a short animation using the CyAnimator, where you show the logFC values changes of the Stages1\_2 and Stage3\_4 with color (- values will be red, + values will be green. Make the animation at least 10 seconds long and create at least 3 key frames. Export the animation to video format and upload it to YouTube (make is publicly available). Insert the link of the video to the document.

\*for 2 task: Since you cannot use UGENE for this you can do the followings: Export all trees from UGENE to Newick format. Copy these trees to a single file. Upload the file to <http://bioinfo.nhri.org.tw/cgi-bin/emboss/fconsense>. Calculate the consensus, download the outtreefile. Read it into the Figtree software and (after formatting) export the tree as a figure.

03.12.2018

1. [↑](#footnote-ref-1)