Statistical Learning, Homework #3

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You should submit a single PDF file of the homework via Moodle, with the PDF rendered directly from a Quarto or RMarkdown source file and not converted from any other output format.

You should write your report like a mini scientific paper, following the guidelines and feedback provided at the previous homeworks. In particular, you should: introduce the analysis, discuss/justify the choices that you make, provide comments on the results that you obtain and draw some conclusions.

Please note that the maximum allowed number of pages is 10.

You will be working on a gene expression data set of 79 patients with leukemia belonging to two subgroups: patients with a chromosomal translocation ("1") and patients cytogenetically normal ("-1"). The data are provided in the attached gene_expr.tsv file, containing expression for 2,000 genes and an additional column with patient labels. You will perform a supervised analysis for prediction of the subgroups using support vector machines.

To this aim:

- Load the data and select a support vector machine for the task at hand. Evaluate different models and justify your final choice.
- A popular approach in gene expression analysis is to keep only the most variable genes for downstream analysis. Since most of the 2K genes have low expression or do not vary much across the experiments, this step usually minimizes the contribution of noise. Select then only genes whose standard deviation is among the top 5% and repeat the analyses performed in the previous task on the filtered data set.
- Draw some conclusions from all the analyses that you have conducted.