

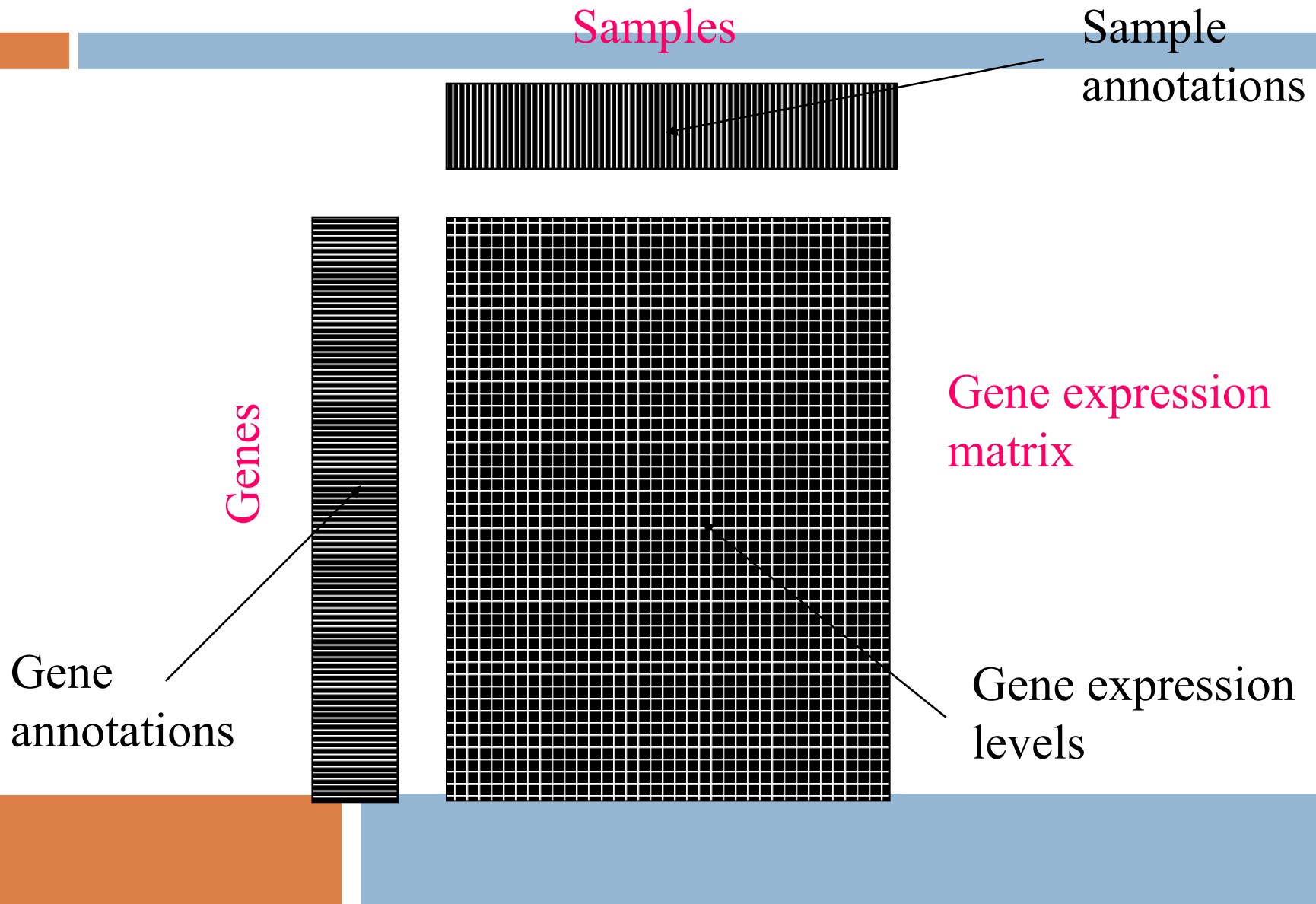
Bioinformatika

Gene expression data analysis

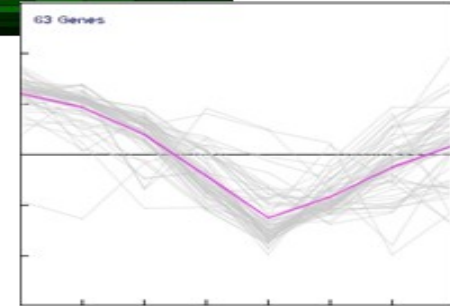
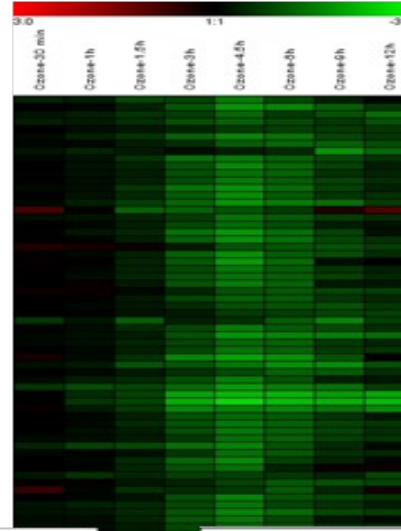
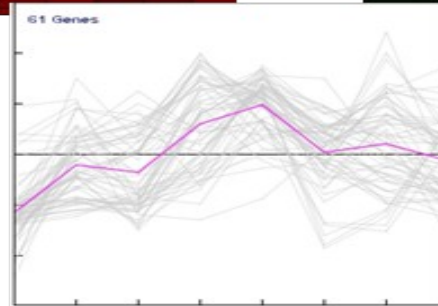
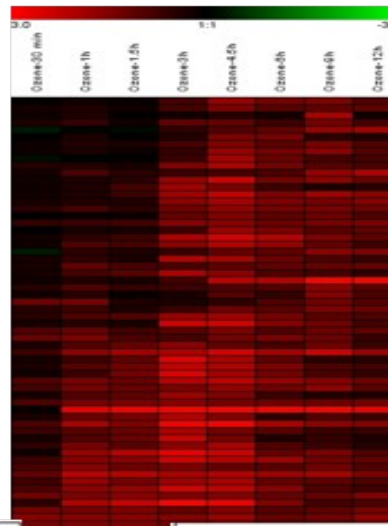
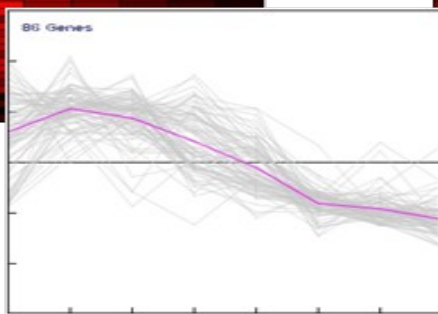
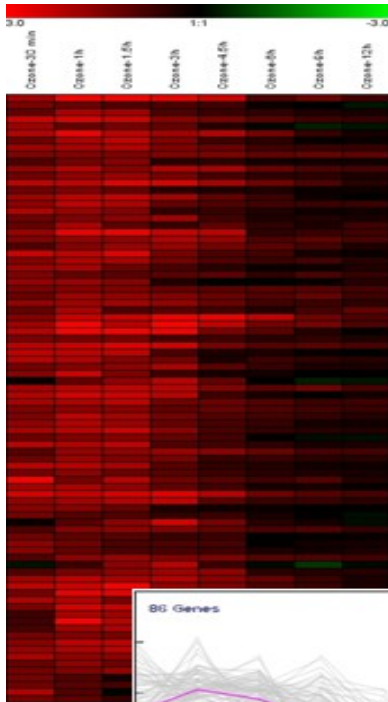
Michael Anděl

(some slides are courtesy of Mark Craven, U. of Wisconsin)

GE data – conceptual view



GE data – image view



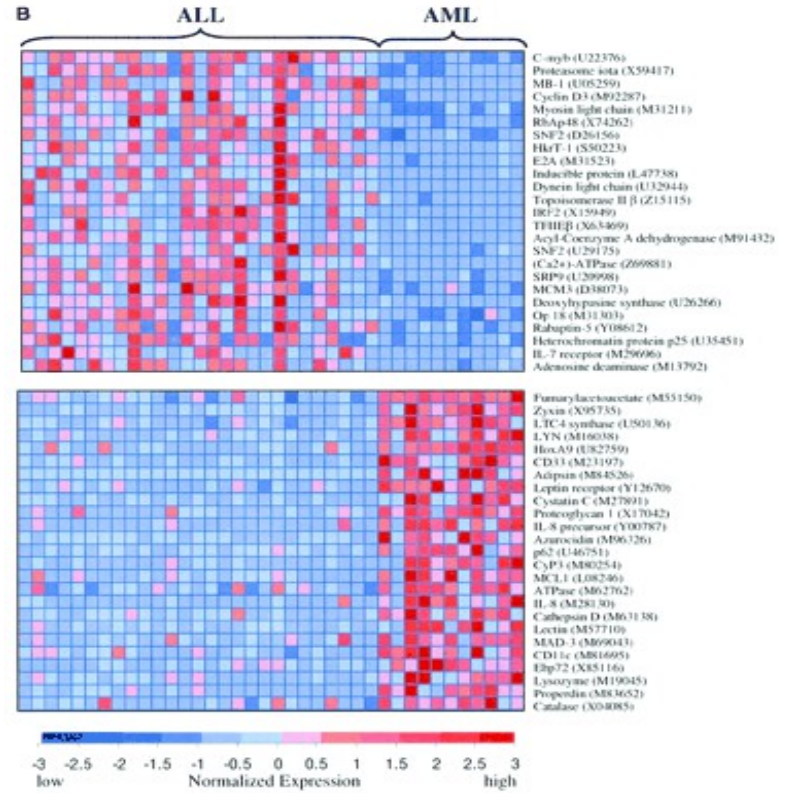
Classification task

Challenge:

- samples (10^1) x features (10^3)
- False hypotheses, overfitting
- Interpretability:
are the expressed genes the causal ones?

What to do?

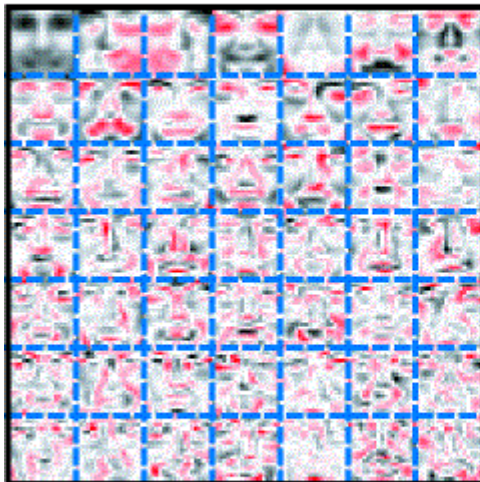
- Decrease number of hypotheses
- Analyze more abstract entities than genes, eg. principal components



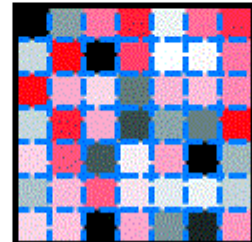
PCA – motivation

- $M \dots \# \text{ genes}$
- $N \dots \# \text{ samples}$
- $X (N \times M) \dots \text{GE data in the space of genes}$
- $V (M \times K) \dots \text{transformation basis, eigengenes}$
- $Z (N \times K) \dots \text{transformed GE data in the space of eigengenes}$
- $K \dots \# \text{ of eigengenes, i.e. the number of underlying concepts}$

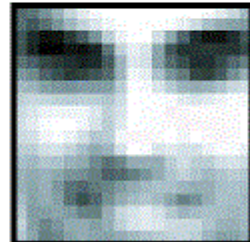
PCA



\times



$=$



Assignment

Data

- 7,129 GE profiles of 72 patients
- 25 samples ... acute myeloid leukemia (AML)
- 47 samples ... acute lymphoblastic leukemia (ALL)

Task

- Construct decision tree model to differentiate these types of tumours.

Assignment

Workflow

- 1) Learn a decision tree on subjected data. Use Matlab class **ClassificationTree**, for learning use method **fit()**.
- 2) Visualize resulting model (function **view()**) and enumerate the training error (function **predict()**)
- 3) Estimate the real error of the model by the means of crossvalidation. Use script **cv11.m**.
- 4) Compare the training error and real error estimation. Why are so different?

Assignment

Analyzing in terms of PCAs

Transform the data to the reduced space of eigengenes. Use attached function **pca.m**:

- 1) Do the transformation for the number of components K=5:5:64, i.e. $Z = X V(1:K,:)$.
- 2) For each of these transformed data **Z** learn the decision tree.
- 3) Choose an appropriate model according to its training error and model complexity.
- 4) Evaluate the chosen model by the crossvalidation
- 5) Compare the error estimation with the one of simple tree.

Assignment

Interpretation of results

- 1) Use web tool Phenopedia HuGE Navigator
- 2) Try to find some gene from the tree in the Phenopedia under the disease term ALL leukemia
- 3) Similarly find some genes from the **component tree**
- 4) To mine genes from the components use attached function **mineGenes.m** and **geneNames.mat**