# Classification by Nearest Shrunken Centroids and Support Vector Machines

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Practical DNA microarray analysis 2005

#### Two roads to classification

- 1. model class probabilities
  - → QDA, LDA, ...
- 2. model class boundaries directly
  - → Optimal Separating Hyperplanes, SVM



#### What's the problem?

In classification you have to trade off overfitting vs. underfitting and bias vs. variance.

In 12'000 dimensions even linear methods are very complex  $\rightarrow$  high variance!

### Simplify your models



## Discriminant analysis and gene selection



#### **Comparing Gaussian likelihoods**

Assumption: each group of patients is well described by a Normal density.

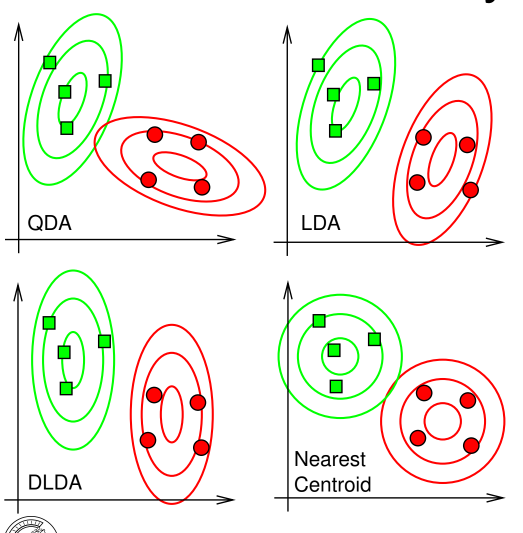
Training: estimate mean and covariance matrix for each group.

Prediction: assign new patient to group with higher likelihood.

Constraints on covariance structure lead to different forms of discriminant analysis.



#### Disriminant analysis in a nutshell



Characterize each class by mean and covariance structure.

- Quadratic D.A.
   different COVs
- Linear D.A. requires same COVs.
- Diagonal linear D.A. same diagonal COVs.
- Nearest centroids forces COVs to  $\sigma^2 \mathbf{I}$ .

#### **Feature selection**

#### **Next simplification:**

Base the classification only on a small number of genes.

Feature selection: Find the most discriminative genes.

This task is different from testing for differential expression. Genes can be significantly differential expressed, but still useless for classification.



#### **Feature selection**

#### 1. Filter:

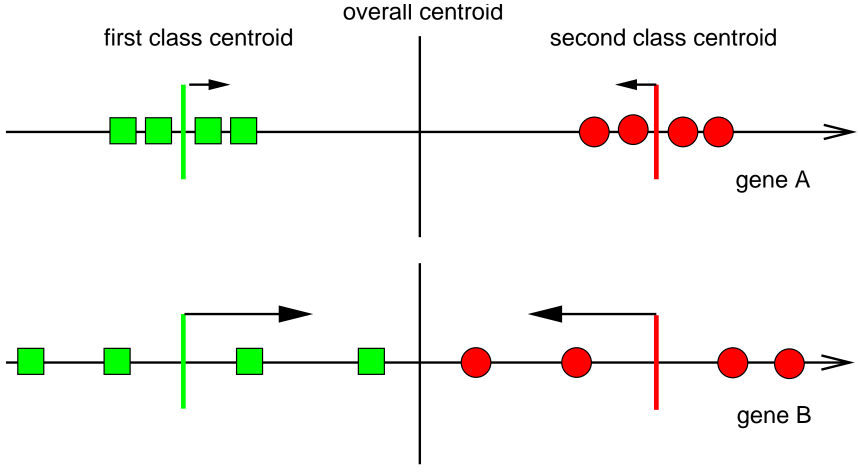
- Rank genes according to discriminative power by t-statistic, Wilcoxon, ...
- Use only the first k for classification.
- Discrete, hard thresholding.

#### 2. Shrinkage:

- Continously shrink genes until only a few have influence on classification.
- Example: Nearest Shrunken Centroids.



#### **Shrunken Centroids**





#### Nearest Shrunken Centroids cont'd

The group centroid  $\bar{x}_{gk}$  for gene g and class k is compared to the overall centroid  $\bar{x}_g$  by

$$\bar{x}_{gk} = \bar{x}_g + m_k(s_g + s_0) d_{gk}$$
,

where  $s_g$  is the pooled within-class standard deviation of gene g and  $s_0$  is an offset to guard against genes with low expression levels.

**Shrinkage:** Each  $d_{gk}$  is reduced by  $\Delta$  in absolute value, until it reaches zero. Genes with  $d_{gk}=0$  for all classes do not contribute to the classification.

(Tibshirani et al., 2002)



#### Shortcomings of filter and shrinkage methods

- 1. High correlated genes get similar score but offer no new information. But see (Jaeger *et al.*, 2003) for a cure.
- 2. Filter and Shrinkage work only on single genes.

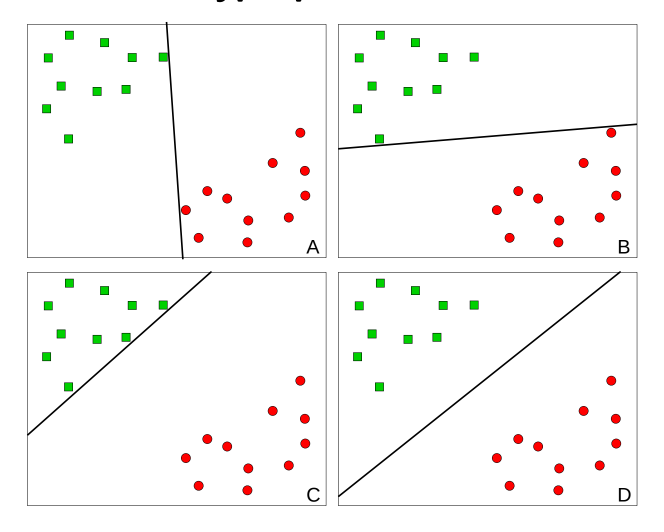
  They don't find interactions between groups of genes.
- 3. Filter and Shrinkage methods are only heuristics. Search for *best subset* is infeasible for more than 30 genes.



### Support Vector Machines — SVM —

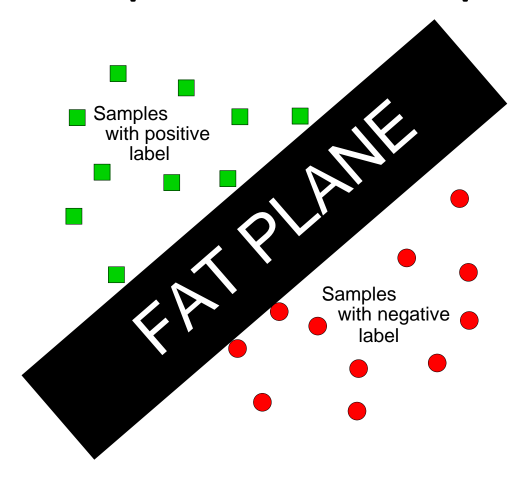


#### Which hyperplane is the best?



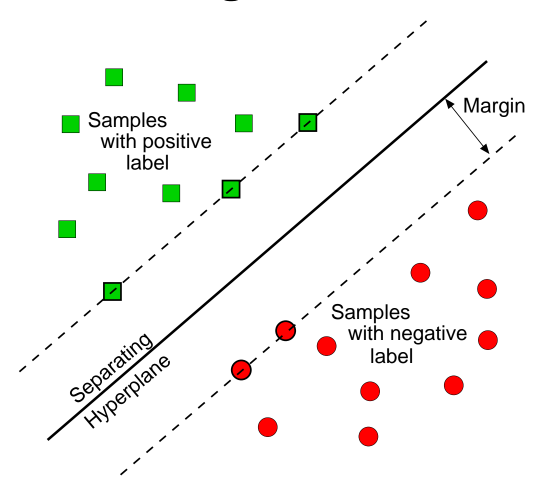


#### No sharp knive, but a fat plane





#### Separate the training set with maximal margin



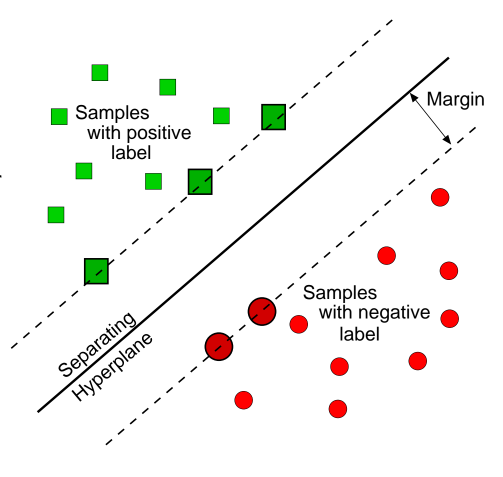


#### What are Support Vectors?

The points nearest to the separating hyperplane are called Support Vectors.

Only they determine the position of the hyperplane. All other points have no influence!

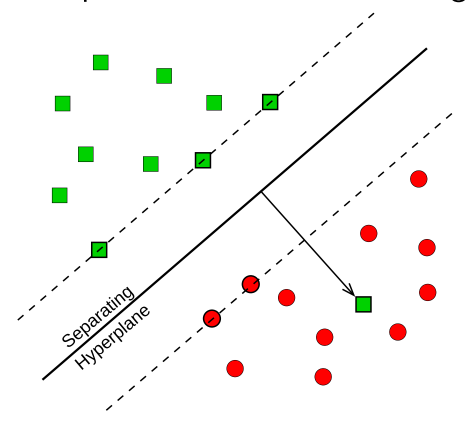
Mathematically: the weighted sum , of the Support Vectors is the normal vector of the hyperplane.





#### Non-separable training sets

Use linear separation, but admit training errors.



Penalty of error: distance to hyperplane multiplied by error cost C.



#### The end?

The story of how to simplify your models is finished.

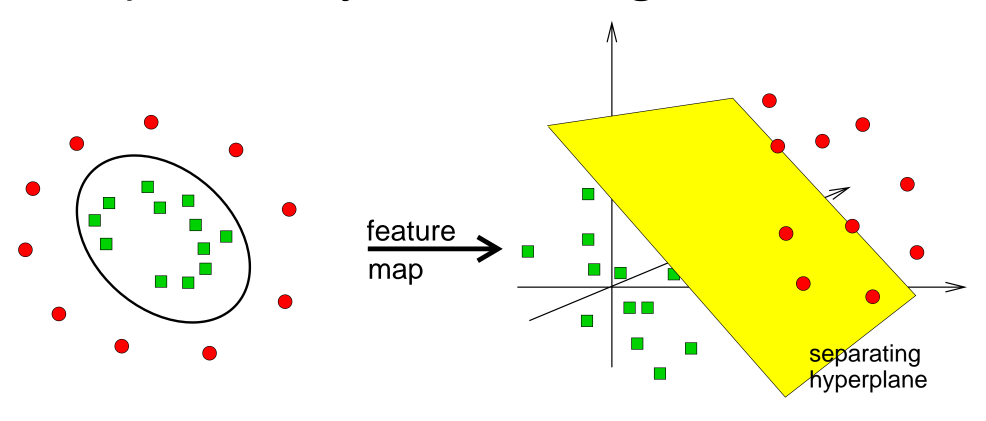
But for the sake of completeness:

How do we get from the simple linear Optimal Separating Hyperplane to a full-grown Support Vector Machine?

It's a trick, a kernel trick.



#### Separation may be easier in higher dimensions



complex in low dimensions

simple in higher dimensions



#### The kernel trick

#### Maximal margin hyperplanes in feature space

If classification is easier in a high-dimenisonal feature space, we would like to build a maximal margin hyperplane there.

The construction depends on inner products  $\Rightarrow$  we will have to evaluate inner products in the feature space.

This can be computationally intractable, if the dimensions become too large!

**Loophole** Use a kernel function that lives in low dimensions, but behaves like an inner product in high dimensions.



#### **Kernel functions**

Expression profiles 
$$p=(p_1,p_2,\ldots,p_g)\in\mathbb{R}^g$$
 and  $q=(q_1,q_2,\ldots,q_g)\in\mathbb{R}^g$ .

#### Similarity in gene space: INNER PRODUCT

$$\langle p, q \rangle = p_1 q_1 + p_2 q_2 + \ldots + p_g q_g$$

#### Similarity in feature space: KERNEL FUNCTION

$$\mathcal{K}(p,q) = \text{polynomial, radial basis, } \dots$$



#### **Examples of Kernels**

linear 
$$\mathcal{K}(p,q) = \langle p,q \rangle$$

polynomial 
$$\mathcal{K}(p,q) = (\gamma \langle p,q \rangle + c_0)^d$$

radial basis function 
$$\mathcal{K}(p,q) = \exp(-\gamma ||p-q||^2)$$



#### Why is it a trick?

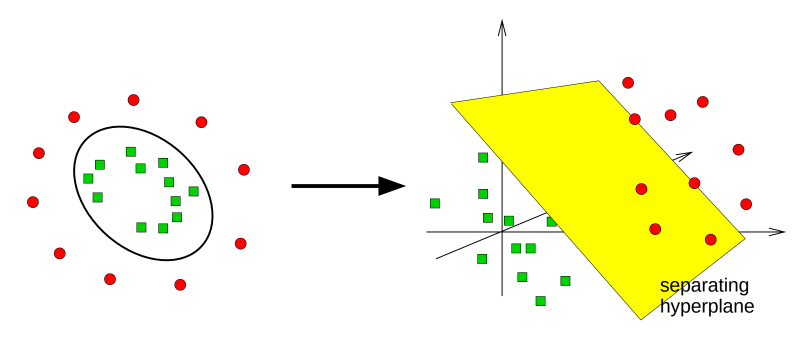
We do not need to know, how the feature space really looks like, we just need the kernel function as a measure of similarity.

This is kind of black magic: we do not know what happens inside the kernel, we just get the output.

Still, we have the geometric interpretation of the maximal margin hyperplane, so SVMs are more transparent than e. g. Artificial Neural Networks.



#### The kernel trick: summary



Non-linear separation between vectors in gene space using kernel functions

Linear separation between vectors in feature space using inner product



#### **Support Vector Machines**

A Support Vector Machine is

a maximal margin hyperplane in feature space

built by using a kernel function in gene space.



#### Parameters of SVM

Kernel Parameters  $\gamma$ : width of rbf

 $\gamma$ : width of rbf coeff. in polynomial ( =1 )

d: degree of polynomial

 $c_0$  additive constant in polynomial (=0)

Error weight C: influence of training errors



#### **SVM@work:** low complexity

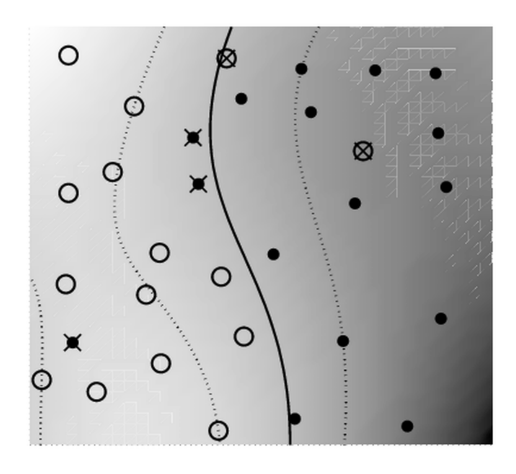


Figure taken from Schölkopf and Smola, Learning with Kernels, MIT Press 2002, p217



#### **SVM@work:** medium complexity

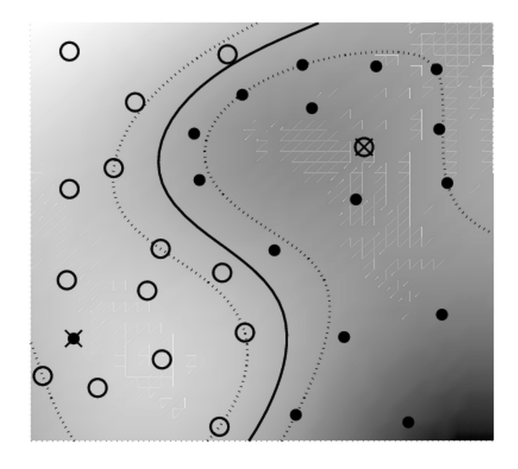


Figure taken from Schölkopf and Smola, Learning with Kernels, MIT Press 2002, p217



#### **SVM@work:** high complexity

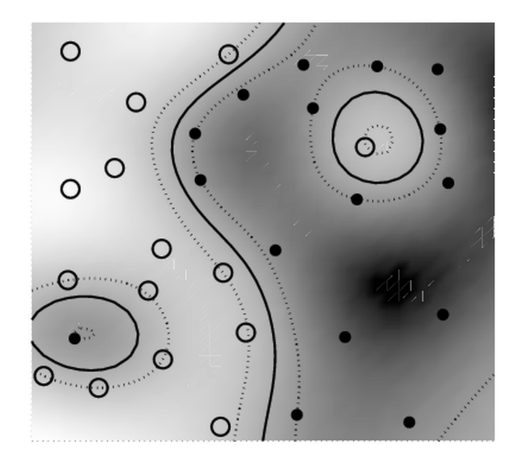


Figure taken from  $SCH\"{O}LKOPF$  and SMOLA, Learning with Kernels, MIT Press 2002, p217



#### References

- 1. Trevor Hastie, Robert Tibshirani, Jerome Friedman **The Elements of Statistical Learning**. Springer 2001.
- Bernhard Schölkopf and Alex Smola.
   Learning with Kernels. MIT Press, Cambridge, MA, 2002.
- 3. Robert Tibshirani, Trevor Hastie, Balasubramanian Narasimhan, Gilbert Chu **Diagnosis of multiple cancer types by shrunken centroids of gene expression**, PNAS, 99(10), 6567–6572, 2002.
- 4. Jochen Jäger, R. Sengupta and W.L. Ruzzo
  Improved Gene Selection for Classification of Microarrays, Proc. PSB 2003



### Intro into practical session



#### **Computational Diagnosis**

#### TASK:

For 3 new patients in your hospital, decide whether they have a chromosomal translocation resulting in a BCR/ABL fusion gene or not.

#### IDEA:

Learn the difference between the cancer types from an archive of 76 expression profiles, which were analyzed and classified by an expert.



#### Training ... tuning ... testing

#### TRAINING:

#### **TUNING:**

#### **TESTING:**

diagnosis <- predict(svm.doctor, new.patients)</pre>



#### Training ... tuning ... testing

#### TRAINING:

```
model <- pamr.train( data , labels )</pre>
```

#### **TUNING:**

```
pamr.cv( data, labels )
```

#### **TESTING:**

diagnosis <- pamr.predict(new.patients, best.treshold)</pre>