# PA196: Pattern Recognition

09. Feature selection and extraction

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Introduction
Filtering methods
Wrapper methods
Feature selection via regularization

## **Outline**

- Feature selection
  - Introduction
  - Filtering methods
  - Wrapper methods
  - Feature selection via regularization
- Peature extraction
  - Principal component analysis
  - Multidimensional scaling



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#### What?

- the vectors to be classified are elements of some d-dimensional space
- the problem is to identify those features that do not contribute to the classification task and eliminate them from classifier training
- thus, we seek the only d<sub>1</sub> < d features that contribute to classification



### Why?

- improve classification performance:
  - redundant features → unstable classifier, poor fit, etc.
  - sparser models have better generalization properties
- improve numerical stability
- reduce the required sample size
- reduce training time, classification time
- improve interpretability of the models



#### How?

- use some optimality criterion to select the optimal subset of features → search strategy
- optimality criterion:
  - single-variate or multi-variate
  - classifier-agnostic: filtering methods
  - use the classifier performance: wrapper methods
- can be implicit in some classifiers:
  - classification trees
  - AdaBoost with decision stumps
  - penalized methods (L<sub>1</sub> penalty)
  - etc
- hybrid: e.g. use a classification tree to select the features and another learner for final classifier



Feature selection

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#### WARNING

The feature selection should be included inside the cross-validation loop (or any other equivalent method) for performance estimation.



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## Filtering methods

Idea: replace using the classifier as criterion with some other criterion simpler (faster) to compute and find a subset of features satisfying this objective.

- single-feature/variable methods
- feature-set methods
- do not guarantee that selected features are relevant for a particular choice of classifier
- usually, introduces a new meta-parameter of the modeling process: the number of features to keep → this needs to be optimized!



## Single-feature methods

- evaluate each variable independently
- may or may not take into account the class label
- suitable approximation for high dimensional cases
- does not avoid selecting correlated features
- fast and easily integrated in modeling pipeline
- the simplest form: discard features that are constant or have low variability



### Using hypothesis testing for feature selection:

• use a statistical test (e.g. t-test) to test

 $H_0$ : there is no difference between classes  $H_1$ : there is a difference between classes

- if H<sub>0</sub> is rejected, we infer that the variable/feature is importance (it bears information about the difference between classes)
- the process involves computing a statistic and comparing it with the expected values under H<sub>0</sub>; if the value is too "unusual" then H<sub>0</sub> is rejected



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- in practice, this results in a "p-value" that is compared with a predefined cut-off  $\alpha$  (called  $\alpha$ -level, traditionally 0.05 or 0.01)
- p-value: the probability, under  $H_0$ , to obtain a statistic at least as extreme as the one from the data at hand
- α-level: probability of falsely rejecting H<sub>0</sub>



### Strategies for selecting the features:

- select top d<sub>1</sub> features (rank the features by statistic or p-values): d<sub>1</sub> needs to be chosen somehow (e.g. by cross-validation)
- select all features with p-value below a selected  $\alpha$ -level
  - this requires adjustment of p-values for multiple testing
  - the easiest is to control family-wise error rate e.g. divide each p-value by the number of tests (variables)
  - for indep. variables, the adjustment does not change the ordering of variables (except for the ties), only the number of variables with significant p-value
  - popular approach: control for false discovery rate (FDR)
  - more complex adjustments are possible also to take into account the correlation structure



## Mutual information-based feature selection

Let *X* and *Y* be two random variables, then their *mutual information* (*MI*) is defined as

$$I(X; Y) = \int_{Y} \int_{X} p(x, y) \log \frac{p(x, y)}{p(x)p(y)} dxdy$$

- provides a measure of dependency between 2 variables
- by considering X as a feature and Y the class to be predicted,
   I(X; Y) gives the relevance of X in predicting Y
- problem: estimation of MI? sample size?
- it can be extended to evaluate the relevance of a set of feature:  $I({X_1, ..., X_k}; Y)$
- can be extended to multi-class problems



#### Other similar measures:

Kullback-Leibler divergence,

$$KL(p||q) = \int p(x) \ln \frac{p(x)}{q(x)} dx$$

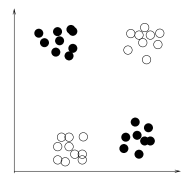
which is usually used in a symmetrized version

- entropy
- Gini index (as in classification trees)



### Issues with single-variable filtering:

- does not account for relationships between variables
- if two variables are not useful for classification when considered isolated, their combination might be useful





## Subset feature selection

- need a criterion to evaluate a set of features (e.g. MI)
- need a strategy to generate all or some of the possible sets of features



## Criteria

Let *J* be the criterion function, which is to be maximized by the best set of features.

- previous probabilistic measures can be extended to sets of features (MI, KL, etc)
- some allow immediate extension to multiclass (e.g. MI) others require a pair-wise approach:  $J(g_i, g_j)$  is computed for all pairs of classes  $g_i, g_j$
- popular choices measure are based on variance estimates.



### Example:

- $J = Tr(S_W^{-1}S_B)$  where  $Tr(\cdot)$  is the trace of a matrix, and  $S_B$  and  $S_W$  are the between-class and within-class scatter matrices, respectively
- $J = \frac{|\hat{\Sigma}|}{|S_W|}$  where  $\hat{\Sigma}$  is the estimated pooled variance matrix
- $J = \frac{Tr(S_B)}{Tr(S_W)}$
- Mahalanobis cr. (for 2 classes):

$$J = (\mu_1 - \mu_2)^T \left(\frac{\Sigma_1 + \Sigma_2}{2}\right)^{-1} (\mu_1 - \mu_2)$$

- in general, the criterion is chosen to allow a recursive decomposition (to compute *J* for *k* + 1 feature sets, one uses the result for *k* feature sets)
- also, to be monotone:

$$X \subseteq Y \Rightarrow J(X) \leq J(Y)$$



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## Search strategies

- Exhaustive search:
  - affordable only if the total number of features is small
  - involves enumerating all possible subsets
  - can be implemented in a breadth-first or depth-first fashion
  - can either start from full set of features (top-down) or from an empty set of features (bottom-up)
- if J is monotone, one can use a branch-and-bound procedure to avoid enumerating all sets; it will still produce a global optimum



## Suboptimal search strategies:

- sequential forward selection (SFS): add sequentially features to the current set such that at each step the added feature produces the maximum increase in J
- generalized SFS (GSFS): instead of adding a single feature, add k features
- sequential backward selection (SBS): start with the full set and remove at each step that feature that leads to minimum decrease in J
- generalized SBS (GSBS):...
- "plus I minus r": add best I and remove worse r features, using sequential selection
- generalized "plus I minus r"



### Floating search methods:

- sequential forward/backward floating selection
- a generalization of "plus I minus r" selection in which I and r can vary
- for example, (sketch of) SFFS:
  - start with empty set of features
  - use SFS to obtain a set of 2 features
  - add the feature that increases J the most and remove the one that decreases it the least; if it is the same feature, consider another feature not yet used
  - continue removing features until J decreases or only 2 features are left; then go to step 3



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## Wrapper methods

- the optimality criterion J is now the performance of the classifier to be trained
- any of the search strategies discussed before can be used in this case
- usually is it more computationally expensive than filtering methods
- produces features that are adapted to the intended classifier
- performance estimation is based on some re-sampling technique (e.g. cross-validation)



## Recursive feature elimination (RFE)

Idea: use the coefficients from the fitted model (classifier) to rank the features and eliminate those with low impact. Repeat the procedure until a convergence criterion is met.

- in LDA-like and (logistic) regression methods, the classifier has the form  $h(\mathbf{x}) = \sum_i w_i x_i$  and  $|w_i|$  can be used to judge the importance of i-th feature
- in the case of regression, one may compute p-values associated with the variables which, again, can be used for ranking them (smaller p-values correspond to more important variables)



## RFE for linear SVM

- use sensitivity analysis of a some cost function: what is the influence of adding/removing a feature?
- the decision function is

$$h(\mathbf{x}) = \sum_{i \in SV} y_i \alpha_i \langle \mathbf{x}, \mathbf{x}_i \rangle = \left\langle \mathbf{x}, \sum_{i \in SV} y_i \alpha_i \mathbf{x}_i \right\rangle$$

- then,  $\nabla_{\mathbf{x}} h(\mathbf{x}) = \sum_{i \in SV} y_i \alpha_i \mathbf{x}_i = \mathbf{w}$  and define the cost function as  $J(\mathbf{w}) = ||\mathbf{w}||$
- the smallest change is in the direction

$$w_j = \arg\min_j(|\mathbf{w}|)$$

i.e. the smallest amplitude coefficient corresponds to the least significant feature

 RFE proceeds by iteratively eliminating the least significant feature(s) and retraining the classifiers on the updated data



## RFE for non-linear SVM

- let  $\mathbf{H} = [y_i y_j K(\mathbf{x}_i, \mathbf{x}_j)]_{ij}$
- define the cost function  $J = \alpha^T H \alpha \langle \alpha, \mathbf{e} \rangle$ ,  $be = [1, ..., 1] \in \mathbb{R}^d$
- trick: to compute the change in J after removing feature k, assume no change in α when retraining on reduced-feature set (if the feature is not important it would change (much) α)
- with this,

$$H_{ij}^{(-k)} = y_i y_j K(\mathbf{x}_i^{(-k)}, \mathbf{x}_j^{(-k)})$$

the change in cost function is

$$\Delta J(k) = J - J^{(-k)} = \alpha^{\mathsf{T}} H \alpha - \alpha^{\mathsf{T}} H^{(-k)} \alpha$$



### Algorithm: to obtain a ranking of features

- let  $X^{\emptyset}$  be the original data set and let  $S = \emptyset$  be the initial ordered list of features (from least to most significant),  $X^{(S)}$  will denote the data set with features in S removed
- repeat 3-6 until no more features are left:
- $\odot$  train the SVM on  $X^{(S)}$  and let  $\alpha$  be the solution
- **o** compute  $\Delta J(i)$  for all features not yet removed  $(i \notin S)$



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## FS via regularization

Idea: introduce some constraints on the model such that the number of effective features is decreased.

- regularization has not necessarily as ultimate goal feature selection
- usually, reg. is imposed by penalizing some complexity measure of the model or the loss functions for classification:

Loss 
$$+ \lambda$$
Penalty

- if **w** is the vector of coefficients, the classical penalties are  $\|\mathbf{w}\|_2^2$  and  $\|\mathbf{w}\|_1$
- what about  $\lambda$ ?

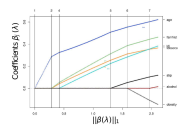


Example: penalized logistic regression.

The new loss function is

$$L(\theta = \{\mathbf{w}, w_0\}) = \sum_{i=1}^{n} [y_i(\langle \mathbf{w}, \mathbf{x}_i \rangle + w_0) - \log(1 + \exp(\langle \mathbf{w}, \mathbf{x}_i \rangle + w_0)] + \lambda \sum_{i=1}^{d} |w_i|$$

From ESL:





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- similar approaches exist for various classifiers
- there is a regularized version for classification trees
- ullet the best choice for  $\lambda$  is usually obtained from cross-validation



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## Feature extraction

Idea: transform the original feature space into a lower dimensional space where some important property (e.g. separability) of data is preserved or enhanced.

- the methods can be based on linear or non linear combinations of original variables
- some methods are variable-directed: they are primarily concerned with relations between variables
- while others are more individual-directed: the distances between data points are of main interest



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## Principal component analysis - PCA

- the goal is to find a linear transformation of the original variables such that the resulting variables are decorrelated
- the hope is to find a smaller number of variables that explain the data
- it is a basic data exploration technique

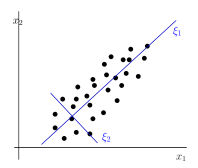


Find an orthogonal transformation **A** of the input data, such that the new variables

$$\boldsymbol{\xi} = \mathbf{A}\mathbf{x}$$

satisfy one of the following (equivalent) constraints:

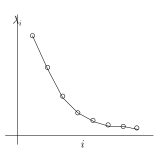
- the variance of  $\xi_i$  is stationary
- variables  $\xi_i$  are decorrelated
- the new axes are the best fit (least square) of the data and are orthogonal





The solution is given by the *eigenvectors* of the covariance matrix of data vectors  $\{\mathbf{x}_i\}$ .

- let λ<sub>i</sub> be the eigenvalues: their ranking gives the order of the principal axes
- $\sum_{i=1}^{d} \lambda_i = \sum_{i=1}^{d} \operatorname{Var}(\xi_i)$
- dimensionality reduction: keep only the first k principal components (e.g. to account for 80% of total variance)
- data approximation: reconstructing the original space from only k PCs





# PCA in pattern recognition

- lots of applications, either alone or in combination with some classifier
- common combinations: PCA + LDA, PCA + SVM
- kernel PCA
- classical application in computer vision: Turk and Pentland's eigenfaces (1991)



## Eigenfaces

Problem: face recognition. (Turk, Pentland: Eigenfaces for recognition. 1991)





#### Principal component analysis Multidimensional scaling







### Face recognition procedure (sketch):

- collect a number of images, say 4 images, per identity
- construct the eigenfaces, an keep the top k
- represent each face in terms of its coordinates in eigenface space  $\Omega = [\omega_1, \dots, \omega_k]$
- for each identity, average the ω-vectors to obtain a characteristic profile
- for a candidate face  $\Omega^*$  compute the Euclidean distance to each for the  $\Omega$ -vectors representing the identities
- if the distance is too large, the candidate face does not represent the claimed identity



- the algorithm can be used for face detection as well
- it generated a lot of follow-up methods
- the method is easy to implement and can be easily be adapted to low-bandwidth environments



#### Other methods:

- PCA is a special case of Karhunen-Loève transform
- generalizations of PCA: common principal components, non-linear PCA
- independent component analysis: no longer an orthogonal transform
- factor analysis, etc



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## Multidimensional scaling (MDS)

- tries to find a set of points that would explain the given dissimilarities (distances)
- it is used as an exploratory technique and for data visualization
- the representation follows the idea: similar objects are represent close together, dissimilar ones, farther apart
- can work both with metric and non-metric data
- in the classical approach, only one (dis)similarity matrix is required; other versions may use several



#### Classical MDS:

- for two objects i and j the similarity is represented in terms of Euclidean distance between points d<sub>ii</sub>
- d<sub>ij</sub> are related to the given similarity through a transformation
   f: δ<sub>ij</sub> = f(d<sub>ij</sub>)
- the problem is to find f such that an error measure is minimized



- MDS is just an instance of larger class of methods: "manifold learning"
- depending on the problem, other transformations could be more useful for visualization or classification
- isomap, local linearly embedding spaces, etc etc



# Example: Roweis and Saul: Nonlinear Dimensionality Reduction by Locally Linear Embedding, Science 2000.

