Selected Topics of Pervasive Computing

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Overview and Structure

- 30.10.2013 Organisational
- 30.10.3013 Introduction
- 06.11.2013 Classification methods (Basic recognition, Bayesian, Non-parametric)
- 13.11.2013 Classification methods (Linear discriminant, Neural networks)
- 20.11.2013 -
- 27.11.2013 -
- 04.12.2013 -
- 11.12.2013 Classification methods (Sequential, Stochastic)
- 18.12.2013 Activity Recognition (Basics, Applications, Algorithms, Metrics)
- 08.01.2014 Security from noisy data (Basics, Entity, F. Commitment, F. Extractors)
- 15.01.2014 Security from noisy data (Error correcting codes, PUFs, Applications)
- 22.01.2014 Context prediction (Algorithms, Applications)
- 29.01.2014 Networked Objects (Sensors and sensor networks, body area networks)
- 05.02.2014 Internet of Things (Sensors and Technology, vision and risks)

Selected Topics of Pervasive Computing

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Outline

Intro

- Recognition of patterns
- Bayesian decision theory
- Non-parametric techniques
- Linear discriminant functions
- Neural networks
- Sequential data
- Stochastic methods









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- Mapping of features onto classes by using prior knowledge
- What are characteristic features?
- Which approaches are suitable to obtain these features?

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Data sampling

- Record sufficient training data
 - Annotated! (Ground-truth)
 - Multiple subjects
 - Various environmental conditions (time of day, weather, ...)



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Data sampling

- Record <u>sufficient</u> training data
 - Annotated! (Ground-truth)
 - Multiple subjects
 - Various environmental conditions (time of day, weather, ...)

Example

- Electric supply data over 15 years covers 5000 days but only 15 christmas days
- Especially critical events like accidents (e.g. plane, car, earthquake) are scarce



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Feature subset-selection



- Pre-process data
 - Framing
 - Normalisation





Feature subset-selection

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Domain knowledge?
                 -> better set of
                     ad-hoc features
        Features commensurate?
                 -> normalise
    Pruning of input required?
                 -> if no, create disjunctive
                    features or weithted
                      sums of features
         Independent features?
                 -> construct conjunctive features
                     or products of features
             Is the data noisy?
                -> detect outlier examples
Do you know what to do first?
                -> If not, use a linear predictor
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- Pre-process data
 - Framing
 - Normalisation

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Feature extraction

- Identify meaningful features
 - remove irrelevant/redundant features



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Feature extraction

- Identify meaningful features
 - remove irrelevant/redundant features
- Features can be contradictory!



Feature subset-selection

Recognition

Simple ranking of features with correlation coefficients Example: Pearson Correlation Coefficient

$$\varrho(X,Y) = \frac{\operatorname{Cov}(X,Y)}{\sqrt{\operatorname{Var}(X)\operatorname{Var}(Y)}}$$
(1)

Identifies linear relation between input variables x_i and an output y

Feature subset-selection

How to do reasonable feature selection

- Utilise dedicated test- and training- data-sets
- Pay attention that a single raw-data sample could not impact features in both these sets
- Don't train the features. on the training- or testdata-set



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Training of the classifier

A decision tree classifier





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Training of the classifier

Evaluation of classification performance

- k-fold cross-validation
 - Standard: k=10



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Training of the classifier Evaluation of classification performance

Leave-one-out cross-validation

- n-fold cross validation where n is the number of instances in the data-set
- Each instance is left out once and the algorithm is trained on the remaining instances
- Performance of left-out instance (success/failure)



Training of the classifier

Evaluation of classification performance

0.632 Bootstrap

- Form training set by choosing n instances from the data-set with replacement
- All not picked instances are used for testing
- Probability to pick a specific instance:

$$1 - \left(1 - \frac{1}{n}\right)'' \approx 1 - e^{-1} \approx 0.632$$



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NN

Training of the classifier

Evaluation of classification performance

Classification accuracy

- Confusion matrices
- Precision
- Recall

	Classification							
	Aw	No	<u>1</u>	Sb	S	Sr	St	Σ
Aw	52		3	6	0	17	22	
No		436	25	7	6	17	9	
To	i.	40	59				1	
Sb	15	22		32	4	22	5	
SI	12	11	1	6	48	8	14	
Sr	4	15		6	1	67	7	
St	3	18	1	1	24	10	43	
\sum	92	551	86	65	94	129	83	

	Classification							
	AW	No	\mathbf{T}_{0}	æ	SI	Sr	St	recall
Aw	.58	.09		.13	.11	.05	.04	
No		.872	.05	.014	.012	.034	.018	
То		.4	.59				.01	
Sb	.15	.22		.32	.04	.22	.05	
SI	.12	.11	.01	.06	.48	.08	.14	
Sr	.04	.15		.06	.01	.67	.07	
St	.03	.18	.01	.01	.24	.1	.43	
prec	.630	.791	.686	.492	.511	.519	.518	

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Evaluation of classification performance

Information score

Let C be the correct class of an instance and $\mathcal{P}(C)$, $\mathcal{P}'(C)$ be the prior and posterior probability of a classifier We define.¹

$$I_{i} = \begin{cases} -\log(\mathcal{P}(C)) + \log(\mathcal{P}'(C)) & \text{if } \mathcal{P}'(C) \geq \mathcal{P}(C) \\ -\log(1 - \mathcal{P}(C)) + \log(1 - \mathcal{P}'(C)) & \text{else} \end{cases}$$
(2)

The information score is then

$$\mathsf{IS} = \frac{1}{n} \sum_{i=1}^{n} I_i \tag{3}$$

^{1.} Kononenko and I. Bratko: Information-Based Evaluation Criterion for Classifier's Performance, Machine 2000

Training of the classifier

Evaluation of classification performance

Brier score

The Brier score is defined as

Brier =
$$\sum_{i=1}^{n} (t(x_i) - p(x_i))^2$$
 (4)

where

$$t(x_i) = \begin{cases} 1 & \text{if } x_i \text{ is the correct class} \\ 0 & \text{else} \end{cases}$$
(5)

and $p(x_i)$ is the probability the classifier assigned to the class x_i .

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Training of the classifier

Evaluation of classification performance

Area under the receiver operated characteristic (ROC) curve (AUC)





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Pattern recognition and classification

Data mining frameworks

- Orange Data Mining (http://orange.biolab.si/)
- Weka Data Mining (http://www.cs.waikato.ac.nz/ml/weka/)





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Sequential

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Pattern recognition and classification

- From features to context
 - Measure available data on features
 - Context reasoning by appropriate method
 - Syntactical (rule based e.g. RuleML)
 - Bayesian classifier
 - Non-parametric
 - Linear discriminant
 - Neural networks
 - Sequential
 - Stochastic



Pattern recognition and classification

- Allocation of sensor value by defined function
 - Correlation of various data sources
 - Several methods possible simple approaches
 - Template matching
 - Minimum distance methods
 - 'Integrated' feature extraction
 - Nearest Neighbour
 - Neural Networks
- Problem
 - Measured raw data might not allow to derive all features required
 - Therefore often combination of sensors



Not enough features



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Pattern recognition and classification

- Methods Syntactical (Rule based)
 - Idea: Description of Situation by formal Symbols and Rules
 - Description of a (agreed on?) world view
 - Example: RuleML
- Comment
 - Pro:
 - Combination of rules and identification of loops and impossible conditions feasible

Contra:

- Very complex with more elaborate situations
- Extension or merge of rule sets typically not possible without contradictions

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Pattern recognition and classification

- Rule Markup Language: Language for publishing and sharing rules
- Hierarchy of rule-sub-languages (XML, RDF, XSLT, OWL)
- Example:
 - A meeting room was occupied by min 5 people for the last 10 minutes.

<atom></atom>	
<rel></rel>	occupied
<var></var>	meeting room
<ind></ind>	min 5 persons
<ind></ind>	last 10 minutes





Pattern recognition and classification

- Also conditions can be modelled
 - A Meeting is taking place in a meeting room when it was occupied by min 5 people for the last 10 minutes.



Pattern recognition and classification

- Logical combination of conditions
 - A Meeting is taking place in a meeting room when it was occupied by min 5 people for the last 10 minutes and the light is on.



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Recognition of patterns

Patterns can be described by a sufficient number of rules

Samples are inaccurate

Tremendous amount of rules to model all variations of one class

Therefore: Consider machine learning approaches



Recognition of patterns

Training set $x_1 \dots x_N$ of a large number of N samples is utilised

Classes $t_1 \dots t_N$ of all samples in this set known in advance

Machine learning algorithm computes a function y(x) and generates a new target t'

 $y(\underline{\sim}) \longrightarrow$

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Polynomial curve fitting

Example

A curve shall be approximated by a machine learning approach

Sample points are created for the function $\sin(2\pi x) + N$ where N is a random noise value



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Polynomial curve fitting

We will try to fit the data points into a polynomial function:

$$y(x, \overrightarrow{w}) = w_0 + w_1 x + w_2 x^2 + \dots + w_M x^M = \sum_{j=0}^M w_j x^j$$



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We will try to fit the data points into a polynomial function:

$$y(x, \overrightarrow{w}) = w_0 + w_1 x + w_2 x^2 + \dots + w_M x^M = \sum_{j=0}^M w_j x^j$$

This can be obtained by minimising an error function that measures the misfit between $y(x, \vec{w})$ and the training data set:

$$E(\overrightarrow{w}) = \frac{1}{2} \sum_{i=1}^{N} \left[y(x_i, \overrightarrow{w}) - t_i \right]^2$$

 $E(\vec{w})$ is non-negative and zero if and only if all points are covered by the function

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Bayesian

Recognition

One problem is the right choice of the dimension M

Non-parametric

When M is too small, the approximation accuracy might be bad

Linear discriminant

NN



Recognition

However, when ${\cal M}$ becomes too big, the resulting polynomial will cross all points exactly

When M reaches the count of samples in the training data set, it is always possible to create a polynomial of order M that contains all values in the data set exactly.



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Polynomial curve fitting This event is called overfitting

The polynomial now trained too well to the training data

It will therefore perform badly on another sample of test data for the same phenomenon

We visualise it by the Root of the Mean Square (RMS) of $E(\vec{w})$





With increasing number of data points, the problem of overfitting becomes less severe for a given value of M



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One solution to cope with overfitting is regularisation

A penalty term is added to the error function

This term discourages the coefficients of \overrightarrow{w} from reaching large values

$$\overline{E}(\overrightarrow{w}) = \frac{1}{2} \sum_{i=1}^{N} \left[y(x_i, \overrightarrow{w}) - t_i \right]^2 + \frac{\lambda}{2} ||\overrightarrow{w}||^2$$

with

$$||\overrightarrow{w}||^2 = \overrightarrow{w}^T \overrightarrow{w} = w_0^2 + w_1^2 + \dots + w_M^2$$

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Depending on the value of λ , overfitting is controlled



$$\overline{E}(\overrightarrow{w}) = rac{1}{2}\sum_{i=1}^{N}\left[y(x_i,\overrightarrow{w}) - t_i
ight]^2 + rac{\lambda}{2}||\overrightarrow{w}||^2$$

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Bayesian decision theory

With probability theory, the probability of events can be estimated by repeatedly generating events and counting their occurrences

When, however, an event only very seldom occurs or is hard to generate, other methods are required

Example:

Probability that the Arctic ice cap will have disappeared by the end of this century

In such cases, we would like to model uncertainty

In fact, it is possible to represent uncertainty by probability

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Linear discriminant

Conditional probability

Conditional probability

The conditional probability of two events χ_1 and χ_2 with $P(\chi_2) > 0$ is denoted by $P(\chi_1|\chi_2)$ and is calculated by

$$P(\chi_1|\chi_2) = \frac{P(\chi_1 \cap \chi_2)}{P(\chi_2)}$$

 $P(\chi_1|\chi_2)$ describes the probability that event χ_2 occurs in the presence of event χ_2 .

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Bayesian decision theory

With the notion of conditional probability we can express the effect of observed data $\overrightarrow{t} = t_1, \ldots, t_N$ on a probability distribution of \overrightarrow{w} : $P(\overrightarrow{w})$.

Thomas Bayes described a way to evaluate the uncertainty of \overrightarrow{w} after observing \overrightarrow{t}

$$P(\overrightarrow{w}|\overrightarrow{t}) = \frac{P(\overrightarrow{t}|\overrightarrow{w})P(\overrightarrow{w})}{P(\overrightarrow{t})}$$

 $P(\overrightarrow{t}|\overrightarrow{w})$ expresses how probable a value for \overrightarrow{t} is given a fixed choice of \overrightarrow{w}

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Bayesian decision theory

A principle difference between Bayesian viewpoint and frequentist viewpoint is that prior assumptions are provided

Example:

Consider a fair coin that scores heads in three consecutive tosses

Classical maximum likelihood estimate will predict head for future tosses with probability $\mathbf{1}$

Bayesian approach includes prior assumptions on the probability of events and would result in a less extreme conclusion



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Bayesian curve fitting

In the curve fitting problem, we are given \overrightarrow{x} and \overrightarrow{t} together with a new sample x_{M+1}

The task is to find a good estimation of the value t_{M+1}

This means that we want to evaluate the predictive distribution

$$p(t_{M+1}|x_{M+1}, \overrightarrow{x}, \overrightarrow{t})$$

To account for measurement inaccuracies, typically a probability distribution (e.g. Gauss) is underlying the sample vector \vec{x}

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Bayesian curve fitting

This means that we want to evaluate the predictive distribution

$$p(t_{M+1}|x_{M+1}, \overrightarrow{x}, \overrightarrow{t})$$

After consistent application of the sum and product rules of probability we can rewrite this as

$$p(t_{M+1}|x_{M+1},\overrightarrow{x},\overrightarrow{t})=\int p(t_{M+1}|x_{M+1},\overrightarrow{w})p(\overrightarrow{w}|\overrightarrow{x},\overrightarrow{t})d\overrightarrow{w}$$

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Bayesian curve fitting



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Example



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Histogram methods

Alternative approach to function estimation: histogram methods

- In general, the probability density of an event is estimated by dividing the range of N values into bins of size Δ_i
- Then, count the number of observations that fall inside bin Δ_i
- This is expressed as a normalised probability density

$$p_i = \frac{n_i}{N\Delta_i}$$



Histogram methods

Accuracy of the estimation is dependent on the width of the bins

Approach well suited for big data since the data items can be discarded once the histogram is created



Histogram methods

Issues:

Due to the edges of the bins, the modelled distribution is characterised by discontinuities not present in the underlying distribution observed

The method does not scale well with increasing dimension (Curse of dimensionality)



Assume an unknown probability density $p(\cdot)$

We want to estimate the probability density $p(\vec{x})$ of \vec{x} in a \mathcal{D} -dimensional Euclidean space

We consider a small region \mathcal{R} around \overrightarrow{x} :

$$P = \int_{\mathcal{R}} p(\overrightarrow{x}) d\overrightarrow{x}$$

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We utilise a data set of N observations

Each observation has a probability of P to fall inside \mathcal{R}

With the binomial distribution we can calculate the count K of points falling into \mathcal{R} :

$$\mathsf{Bin}(K|N,P) = \frac{N!}{K!(N-K)!} P^{K} (1-P)^{N-K}$$

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We utilise a data set of N observations

Each observation has a probability of P to fall inside $\mathcal R$

With the binomial distribution we can calculate the count K of points falling into \mathcal{R} :

$$\mathsf{Bin}(K|N,P) = \frac{N!}{K!(N-K)!} P^{K} (1-P)^{N-K}$$

For large N we can show

 $K \approx NP$

With sufficiently small $\mathcal R$ we can also show for the volume V of $\mathcal R$

 $P\approx p(\overrightarrow{x})V$

Therefore, we can estimate the density as

$$p(\overrightarrow{x}) = \frac{K}{NV}$$

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Non-parametric

Parzen estimator methods

We assume that ${\mathcal R}$ is a small hypercube

In order to count the number K of points that fall inside \mathcal{R} we define

$$k(\overrightarrow{u}) = \left\{egin{array}{cc} 1, & |u_i| \leq rac{1}{2}, & i=1,\ldots,D, \ 0, & ext{otherwise} \end{array}
ight.$$

This represents a unit cube centred around the origin

This function is an example of a kernel-function or Parzen window

$$k(\overrightarrow{u}) = \left\{ egin{array}{cc} 1, & |u_i| \leq rac{1}{2}, & i=1,\ldots,D, \\ 0, & ext{otherwise} \end{array}
ight.$$

When the measured data point $\overrightarrow{x_n}$ lies inside a cube of side *h* centred around \overrightarrow{x} , we have

$$k\left(\frac{\overrightarrow{x}-\overrightarrow{x_n}}{h}\right)=1$$

The total count K of points that fall inside this cube is

$$K = \sum_{n=1}^{N} k \left(\frac{\overrightarrow{\mathbf{x}} - \overrightarrow{\mathbf{x}_n}}{h} \right)$$

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The total count K of points that fall inside this cube is

$$K = \sum_{n=1}^{N} k\left(\frac{\overrightarrow{x} - \overrightarrow{x_n}}{h}\right)$$

When we substitute this in the density estimate derived above

$$p(\overrightarrow{x}) = \frac{K}{NV}$$

with volume $V = h^D$ we obtain the overall density estimate as

$$p(\overrightarrow{x}) = \frac{1}{N} \sum_{n=1}^{N} \frac{1}{h^{D}} \left(\frac{\overrightarrow{x} - \overrightarrow{x_{n}}}{h} \right)$$

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$$p(\overrightarrow{x}) = \frac{1}{N} \sum_{n=1}^{N} \frac{1}{h^{D}} \left(\frac{\overrightarrow{x} - \overrightarrow{x_{n}}}{h} \right)$$

Again, this density estimator suffers from artificial discontinuities (Due to the fixed boundaries of the cubes)

Problem can be overcome by choosing a smoother kernel function (A common choice is a Gaussian kernel with a standard deviation σ)

$$p(\overrightarrow{x}) = \frac{1}{N} \sum_{n=1}^{N} \frac{1}{(2\pi\sigma^2)^{\frac{D}{2}}} e^{-\frac{||\overrightarrow{x} - \overrightarrow{x_n}||^2}{2\sigma^2}}$$

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Density estimation for various values of $\boldsymbol{\sigma}$



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A problem with Parzen estimator methods is that the parameter governing the kernel width (*h* or σ) is fixed for all values \overrightarrow{x}

In regions with

...high density, a wide kernel might lead to over-smoothing ...low density, the same width may lead to noisy estimates



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NN-methods address this by adapting width to data density

Parzen estimator methods fix V and determine K from the data Nearest neighbour methods fix K and choose V accordingly

Again, we consider a point \overrightarrow{x} and estimate the density $p(\overrightarrow{x})$

The radius of the sphere is increased until K data points (the nearest neighbours) are covered

The value K then controls the amount of smoothing Again, an optimum value for K exists



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<u>Classification</u>: Apply KNN-density estimation for each class Assume data set of N points with N_k points in class C_k To classify sample \overrightarrow{x} , draw a sphere containing K points around \overrightarrow{x} Sphere can contain other points regardless of their class

Assume sphere has volume V and contains K_k points from C_k

<u>Assume:</u> Sphere of volume V contains K_k points from class C_k

We estimate the density of class C_k as

$$p(\overrightarrow{x}|C_k) = \frac{K_k}{N_k V}$$

The unconditional density is given as

$$p(\overrightarrow{x}) = \frac{K}{NV}$$

The probability to experience a class C_k is given as

$$p(C_k) = \frac{N_k}{N}$$

With Bayes theorem we can combine this to achieve

$$p(C_k | \vec{x}) = \frac{p(\vec{x} | C_k) p(C_k)}{p(\vec{x})} = \frac{K_k}{K}$$

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Nearest neighbour methods

$$p(C_k | \overrightarrow{x}) = \frac{p(\overrightarrow{x} | C_k) p(C_k)}{p(\overrightarrow{x})} = \frac{K_k}{K}$$

To minimise the probability of misclassification, assign \overrightarrow{x} to class with the largest probability

This corresponds to the largest value of

 $\frac{K_k}{K}$

To classify a point, we identify the K nearest points

And assign the point to the class having most representatives in this set

Choice K = 1 is called nearest neighbour rule

For this choice, the error rate is never more than twice the minimum achievable error rate of an optimum classifier $^2\,$

²T. Cover and P. Hart: Nearest neighbour pattern classification. IEEE Transactions on Information Theory, IT-11, 21-27, 1967 < □ > < ≧ > < ≧ > < ≧ > < ≧ > < ≧

Classification of points by the K-nearest neighbour classifier



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Nearest neighbour methods

Classification of points by the K-nearest neighbour classifier



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Sequential

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Nearest neighbour methods

The KNN-method and the Parzen-method are not well suited for large data sets since they require the entire data set to be stored

Stochastic

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In classification we assign \overrightarrow{x} to one of K discrete classes C_k The input is divided by decision boundaries

Here we assume that decision boundaries are linear functions of \overrightarrow{x} Data separable by linear decision surfaces are linear separable With high dimension, a set of two classes is always linear separable



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SVM pre-processes data to represent patterns in a high dimension Dimension often much higher than original feature space Then, insert hyperplane in order to separate the data



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Support vector machines (SVM) A pattern $\overrightarrow{x_k}$ is transformed to $\overrightarrow{y_k} = \varphi(\overrightarrow{x_k})$ Also, each $\overrightarrow{x_k}$ is associated with $z_k \in \{-1, 1\}$ A linear discriminant in an augmented \overrightarrow{y} space is $g(\overrightarrow{y}) = \overrightarrow{a} t \overrightarrow{y}$ A separating hyperplane ensures for $y_0 = 1, a_0 \ge 1$

 $z_k g(y_k) \geq 1$



The goal for support vector machines is to find a separating hyperplane with the largest margin b to the outer points in all sets

$$\frac{z_k g(y_k)}{||\overrightarrow{a}||} \ge b, \ k = 1, \dots, n$$

If no such hyperplane exists, map all points into a higher dimensional space until such a plane exists

Support vectors satisfy ' $\cdot = b$ '



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Intro Recognition Bayesian Non-parametric (Linear discriminant) NN Sequential Stoch

Support vector machines (SVM)

Simple application to several classes by iterative approach:

belongs to class 1 or not?

belongs to class 2 or not?

...
```

Search for optimum mapping between input space and feature space complicated (no optimum approach known)



Simple learning approach to find the correct hyperplane:

Starting from an initial separating hyperplane

Find worst classified pattern (on the wrong side of the hyperplane)

Design a new hyperplane with this pattern as one of the support vectors

Iterate until all patterns are correctly classified



Outline

Intro

- Recognition of patterns
- Bayesian decision theory
- Non-parametric techniques
- Linear discriminant functions
- Neural networks
- Sequential data
- Stochastic methods

(NN)

Sequential

Stochastic

Neural networks

Learn mapping from input to output vector

Representation by edge-weighted graph

Distinction between

- Input neurons
- Output neurons
- Hidden nodes



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Neural networks

Input neurons are only equipped with outgoing edges

Hidden nodes 'fire' (output value 1) when weighted inputs exceed threshold function Θ



$$y_i = \begin{cases} 1, & \text{if } \sum_{i=1}^n x_i w_i \ge \Theta \\ 0, & \text{else.} \end{cases}$$

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Learning with back-propagation (schematic):

- (Iterate until the error is sufficiently small)
 - Choose a training-pair and copy it to the input layer
 - Propagate it through the network
 - Salculate error between computed and expected output
 - Propagate the sum product of the weights back into the network in order to calculate the error in internal layers
 - Adapt weights to the error



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Neural networks

Single hidden layer sufficient to represent arbitrary multi-dimensional functions

- Well suited for noisy input data
- Implicit clustering of input data possible
- Complex to extend network (e.g. add new features)



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For the input layer, we construct M linear combinations of the input variables x_1, \ldots, x_D and weights w_1, \ldots, w_D

$$a_j = \sum_{i=1}^D w_{ji}^{(1)} x_i + w_{j0}^{(1)}$$

Each a_j is transformed using a differentiable, non-linear activation function

$$z_j = h(a_j)$$

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Input layer *M* linear combinations of x_1, \ldots, x_D and w_1, \ldots, w_D

$$a_j = \sum_{i=1}^{D} w_{ji}^{(1)} x_i + w_{j0}^{(1)}$$

Activation function: Differentiable, non-linear

 $z_j=h(a_j)$

 $h(\cdot)$ function is usually a sigmoidal function or tanh



Values z_i are again linearly combined in hidden layers:

$$a_k = \sum_{j=1}^M w_{kj}^{(2)} z_j + w_{k0}^{(2)}$$

with $k = 1, \ldots, K$ describing the total number of outputs

Again, these values are transformed using a sufficient transformation function σ to obtain the network outputs y_k

$$y_k = \sigma(a_k)$$

For multi-class problems, we use a function such as

$$\sigma(a) = \frac{1}{1 + e^{-a}}$$

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Combine these stages to achieve overall network function:

$$y_k(\overrightarrow{x},\overrightarrow{w}) = \sigma \left(\sum_{j=1}^M w_{kj}^{(2)} h\left(\sum_{i=1}^D w_{ji}^{(1)} x_i + w_{j0}^{(1)} \right) + w_{k0}^{(2)} \right)$$

(Multiple hidden layers are added analogously)

Neural networks

Activation functions of hidden units are linear \Rightarrow Always find equivalent network without hidden units

(Composition of successive linear transformations itself linear transformation)



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Number of hidden units < number of input or output units \Rightarrow not all linear functions possible

(Information lost in dimensionality reduction at hidden units)



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Neural networks are Universal approximators³ 4 5 6 7 8 9 10 \Rightarrow 2-layer linear NN can approximate any continuous function

 4 G. Cybenko: Approximation by superpositions of a sigmoidal function. Mathematics of control, signals and systems, 2, 304-314, 1989

 5 K. Hornik, M. Sinchcombe, H. White: Multilayer feed-forward networks are universal approximators. Neural Networks, 2(5), 359-366, 1989

 $^6{\rm N.E.}$ Cotter: The stone-Weierstrass theorem and its application to neural networks. IEEE Transactions on Neural Networks 1(4), 290-295, 1990

 7 Y. Ito: Representation of functions by superpositions of a step or sigmoid function and their applications to neural network theory. Neural Networks 4(3), 385-394, 1991

 $^8{\rm K}.$ Hornik: Approximation capabilities of multilayer feed forward networks: Neural Networks, 4(2), 251-257, 1991_

 $^9{\rm Y.V.}$ Kreinovich: Arbitrary non-linearity is sufficient to represent all functions by neural networks: a theorem. Neural Networks 4(3), 381-383, 1991

 $^{^3{\}rm K}.$ Funahashi: On the approximate realisation of continuous mappings by neural networks, Neural Networks, 2(3), 183-192, 1989

Remaining issue in neural networks

- Find suitable parameters given a set of training data
- Several learning approaches have been proposed



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Simple approach to determine network parameters: Minimise sum-of-squared error function

- Given a training set $\overrightarrow{x_n}$ with $n \in \{1, \dots, N\}$
- And corresponding target vectors $\overrightarrow{t_n}$
- Minimise the error function

$$E(\overrightarrow{w}) = \frac{1}{2} \sum_{n=1}^{N} (y(\overrightarrow{x_n}, \overrightarrow{w}) - \overrightarrow{t_n})^2$$

Neural networks - Classification

2 classes C_1 and C_2

• We consider a network with a single output

$$y = \sigma(a) \equiv rac{1}{1 + e^{-a}}$$

- Output interpreted as conditional probability $p(\mathcal{C}_1 | \overrightarrow{X})$
- Analogously, we have $p(\mathcal{C}_2|\overrightarrow{x}) = 1 p(\mathcal{C}_1|\overrightarrow{x})$

<u>*K*</u> classes C_1, \cdots, C_K

- Binary target variables $t_k \in \{0,1\}$
- Network outputs are interpreted as $y_k(\overrightarrow{x}, \overrightarrow{w}) = p(t_k = 1 | \overrightarrow{x})$

Introduction to self organising maps (SOM) Proposed by Teuvo Kohonen¹¹

As a model of the self-organisation of neural connections

Maps high dimensional input to low dimensional output

Based on neural network learning of the underlying mapping



Introduction to self organising maps

Present all points in a source space by points in a target space

Given a sequence of points in a sample space,

Create a mapping of these points into a target space that respects the neighbourhood relation in the sample space



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SOM is a topology preserving lattice of predefined number of nodes Represents topology of elements in input space.

Algorithm inherits self-organisation property

- Able to produce organisation starting from total disorder.
- Defines and preserves neighbourhood structure between nodes

Learning by two layer neural network

ant (NN)

Sequential

Stochastic

Introduction to self organising maps



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Introduction to self organising maps



When a pattern $\vec{\phi_i}$ is presented, each node (represented by outer neurons) in the target space computes its activation $\vec{\phi_i} \cdot \vec{w}$.

Most activated node y^* and weights to its neighbours are updated according to a learning rate $\rho(t)$

$$w_{ki}(t+1) = w_{ki}(t) + \rho(t)\Lambda(|y-y^*|)(\overrightarrow{\phi_i} - w_{ki}(t))$$

 $\Lambda(\cdot)$ defines a non-increasing neighbourhood function and $|y - y^*|$ describes the distance of nodes in the neighbourhood

SOM – Self organisation



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SOM – Definition

Condensed definition of SOM from Cottrell et al.¹²

Self organising maps

- Let $I = \{\overrightarrow{\eta_1}, \dots, \overrightarrow{\eta_{|S|}}\}$ be a set of *km*-dimensional vectors that are associated with nodes in a lattice.
- Neighbourhood structure provided by symmetrical neighbourhood function $d: I \times I \to \mathbb{R}$ which depends on the distance between two nodes $\overrightarrow{\eta_i}$ and $\overrightarrow{\eta_i} \in I$.
- State of the map at time t given by

$$\eta(t) = \left(\overrightarrow{\eta_1(t)}, \overrightarrow{\eta_2(t)}, \dots, \overrightarrow{\eta_{|S|}(t)}\right),$$

¹²M. Cottrell, J.C. Fort and G. Pages, *Theoretical aspects of the SOM algorithm*, Neurocomputing, pp. 119-138, vol 21, 1998.

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SOM – Definition

Self organising map algorithm The SOM algorithm is recursively defined by

$$\begin{split} i_{c}\left(\overrightarrow{v(t+1)},\overrightarrow{\eta(t)}\right) &= \arg\min\left\{\left\|\overrightarrow{v(t+1)}-\overrightarrow{\eta_{i}(t)}\right\|,\overrightarrow{\eta_{i}(t)}\in\eta(t)\right\},\\ \overrightarrow{\eta_{i}(t+1)} &= \overrightarrow{\eta_{i}(t)}-\varepsilon_{t}d\left[i_{c}\left(\overrightarrow{v(t+1)},\overrightarrow{\eta(t)}\right),\overrightarrow{\eta_{i}}\right]\\ &\cdot\left(\overrightarrow{\eta_{i}(t)}-\overrightarrow{v(t+1)}\right),\forall\overrightarrow{\eta_{i}}\in I. \end{split}$$

In this formula, $i_c\left(\overrightarrow{v(t+1)}, \overrightarrow{\eta(t)}\right)$ corresponds to the node in the network that is closest to the input vector.

Parameter ε_t controls the adaptability.

ant (NN)

Sequential

Stochastic

SOM – Operational principle

$\begin{array}{c} \text{Inputs} \\ (v_{s}(t)) & (v_{s}(t)) & (v_{s}(t)) \end{array}$

Input values $v_i(t)$ are to be mapped onto the target space

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SOM – Operational principle



Node with the lowest distance is associated with the input value:

$$i_{c}\left(\overrightarrow{v(t+1)},\overrightarrow{\eta(t)}
ight) = \operatorname{argmin}\left\{\left\|\overrightarrow{v(t+1)} - \overrightarrow{\eta_{i}(t)}\right\|,\overrightarrow{\eta_{i}(t)} \in \eta(t)
ight\}$$

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Linear discriminant

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SOM – Operational principle



Nodes in the neighbourhood of the associated node are moved closer to the input value

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Linear discriminant

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SOM – Operational principle



Nodes in the neighbourhood of the associated node are moved to the input value

$$\overrightarrow{\eta_i(t+1)} = \overrightarrow{\eta_i(t)} - \varepsilon_t d \left[i_c \left(\overrightarrow{v(t+1)}, \overrightarrow{\eta(t)} \right), \overrightarrow{\eta_i} \right] \\ \cdot \left(\overrightarrow{\eta_i(t)} - \overrightarrow{v(t+1)} \right), \forall \overrightarrow{\eta_i} \in I.$$

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Recognition

Sequential

SOM – Example application: TEA



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SOM – Example application: TEA



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SOM – Remarks

SOM algorithm always converges¹³

Normalisation of input vectors might improve numerical accuracy

Not guaranteed that self-optimisation will always occur (Dependent on choice of parameters)

Difficult to set parameters of the model since SOM is not optimising any well-defined function $^{14}\,$

If neighbourhood is chosen to be too small, the map will not be ordered globally

¹³Y. Cheng, Neural Computation, 9(8), 1997.

¹⁴E. Erwin, K. Obermayer, K. Schulten: Self-organising maps: Ordering, convergence properties and energy functions. Biological Cybernetics, 67, 47-55, 1992

Linear discriminant

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Sequential

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Problems of SOMs



Map created as target space might have several orientations

One part of the map might follow one orientation, while other parts are following other orientations

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Stochastic

Outline

Intro

Recognition

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