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Feature Transform

- Consider the following spaces:
  - $\mathbb{U}$ denotes the universe of data objects
  - $\mathbb{F} \subseteq \mathbb{R}^n$ denotes an $n$-dimensional feature space
- A feature transformation is a mapping $f : \mathbb{U} \rightarrow \mathbb{R}^n$ of objects from $\mathbb{U}$ to the feature space $\mathbb{F}$.

Similarity Model

- A similarity model $S : \mathbb{U} \times \mathbb{U} \rightarrow \mathbb{R}$ is defined for all objects $p, q \in \mathbb{U}$ as
  $$S(p, q) = \text{sim}(f(p), f(q))$$
  where $\text{sim} : \mathbb{R}^n \times \mathbb{R}^n \rightarrow \mathbb{R}$ is a similarity measure or a dissimilarity (distance) measure in $\mathbb{F}$. 
Comments:

- Often, dissimilarity (distance) is measured instead of similarity
- This is a small but important difference!
  - A similarity measure \((sim)\) assigns high values to similar objects
  - A dissimilarity measure \((dist)\) assigns low values to similar objects
- The design of \(f\) and the definition of \(sim/dist\) are important assumptions about the patterns we want to find later in the data
- As explained before, \(f\) and \(sim/dist\) can be derived manually (explicit transformation and coding versus implicit Kernels) or automatically (representation learning)
Dissimilarity

- Dissimilarity measures follow the idea of the geometric approach
  - objects are defined by their perceptual representations in a perceptual space
  - perceptual space = psychological space
  - geometric distance between the perceptual representations defines the (dis)similarity of objects

- Within the scope of Feature-based similarity
  - perceptual space = feature space $\mathbb{F}$ or feature representation space $\mathbb{R}^n$
  - geometric distance = distance function
Distance Functions

- The distance measure \( dist \) is a distance function if it is reflexive, non-negative, and symmetric.
- A distance function \( dist \) is a metric if it additionally satisfies the triangle inequality.
- Comments:
  - Sound mathematical interpretation
  - Allow domain experts to model their notion of dissimilarity
  - Metric distances allow to tune efficiency of data mining approaches
  - Long-lasting discussion of whether the distance properties and in particular the metric properties reflect the perceived dissimilarity correctly, see the following contradicting example:

![Images showing the example](Images/lecture_images)
Similarity versus Dissimilarity (again)

• Transformation
  • Let $\mathbb{F}$ be a feature space and $\text{dist} : \mathbb{F} \times \mathbb{F} \rightarrow \mathbb{R}$ be a distance function
  • Any monotonically decreasing function $f : \mathbb{R} \rightarrow \mathbb{R}$ defines a similarity function $s : \mathbb{F} \times \mathbb{F} \rightarrow \mathbb{R}$ as follows

\[
\forall x, y \in \mathbb{F} : s(x, y) = f(\text{dist}(x, y))
\]

• Some prominent similarity functions ($x, y \in \mathbb{F}$):
  • exponential:
    \[
s(x, y) = e^{-\text{dist}(x, y)}
    \]
  • logarithmic:
    \[
s(x, y) = 1 - \log(1 + \text{dist}(x, y))
    \]
  • linear: $s(x, y) = 1 - \text{dist}(x, y)$
Similarities: Examples (only very few)

- **Dot-Product** \((x, y \in \mathbb{F} \subseteq \mathbb{R}^d)\)
  \[
x \cdot y^T = \sum_{i=1}^{d} x_i \cdot y_i = \|x\| \cdot \|y\| \cdot \cos \angle(x, y)
  \]

- **Cosine** \((x, y \in \mathbb{F} \subseteq \mathbb{R}^d)\)
  \[
  \frac{x \cdot y^T}{\|x\| \cdot \|y\|}
  \]

- **Pearson Correlation** \((x, y \in \mathbb{F} \subseteq \mathbb{R}^d)\)
  \[
  \frac{\sum_{i=1}^{d} (x_i - \bar{x}_i)(y_i - \bar{y}_i)}{\sqrt{\sum_{i=1}^{d} (x_i - \bar{x}_i)^2} \cdot \sqrt{\sum_{i=1}^{d} (y_i - \bar{y}_i)^2}}
  \]
  where \(\bar{z}_i\) denotes the mean in attribute \(i\) over all data points

- **Random-Walk Kernel** (for graphs \(x, y\))
  - Count common (random) walks in \(x\) and \(y\)
  - Walks are sequences of nodes (connected by edges)
Distances: Examples (only very few)

- $L_p$-norm (aka Minkowski metric) ($x, y \in F \subseteq \mathbb{R}^d$)

\[ L_p(x, y) = \sqrt[p]{\sum_{i=1}^{d} |x_i - y_i|^p} \]

where
- $p < 1$: fractional Minkowski distance
- $p = 1$: Manhattan distance
- $p = 2$: Euclidean distance
- $p = \infty$: Chebyshev/Maximum distance

- Malahanobis distance

- Hamming distance $\text{HammingDist}(x, y) = \sum_{i=1}^{d} \begin{cases} 
1 & : x_i \neq y_i \\
0 & : \text{else}
\end{cases}$
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A Motivating Example

• Let’s play the baby shapes game (truly motivating for students ...): Group the items!!!

• What about grouping based on both shape and color?

• Lesson to learn: there may be different semantic concepts (and their corresponding patterns) hidden in the data (here: shape and color)
The good old days of data mining . . .

- Data generation and, to some extend, data storage was costly (hard to imagine but those were the days ...)
- Domain experts carefully considered which features/variables to measure before designing experiments/a feature transform/...
- Consequence: also data sets were well designed and potentially contained only a small number of relevant features
Nowadays, data science is also about integrating everything

- Generating and storing data is easy and cheap
- People tend to measure everything they can and even more (including even more complex feature transformations)
- The Data Science mantra is often interpreted as “we can analyze data from as many sources as (technically) possible, just record anything you can”
- Consequence: data sets are high-dimensional containing a large number of features but the relevancy of each feature for the analysis goal is not clear a priori
High-dimensional Data is NOT a Myth

• Example: Image data
  • Low-level image descriptors (color histograms, textures, shape information ...)
  • Regional descriptors: between 16 and 1,000 features
  • ...

• Example: Metabolome data
  • Feature = concentration of one metabolite (intermediates/results of metabolism)
  • Bavaria newborn screening (for each baby, the blood concentrations of 43 metabolites are measured in the first 48 hours after birth)
  • between 50 and 2,000 features
More High-dimensional Data

• Example: Microarray data (deprecated)
  • Features correspond to genes
  • Up to 20,000 features
  • Dimensionality is much higher than the sample size

• Example: Text data
  • Term frequency: features correspond to words/terms
  • Between 5,000 and 20,000 features (and even more)
  • Often, esp. in social media: abbreviations, colloquial language, special words

Excerpt from LMU website:
http://tinyurl.com/qhq6byz
Overview:

- Distances grow
- Contrast of distances diminish (concentration problem)
- Meaning of “neighborhood” concept
- Growing data space
- Growing hypothesis space
- Empty spaces and importance tails
- Different semantic layers
- ...

So let us have a closer look on these problems ...
The following example uses the Euclidean distance but holds for most distance measures:

- Consider 2D vectors \( a = (1, 2) \) and \( b = (4, 3) \)
- The Euclidean distance between \( a \) and \( b \) is

\[
L_2(a, b) = L_2((1, 2), (4, 4)) = \sqrt{(1 - 4)^2 + (2 - 3)^2} = \sqrt{10}
\]

which corresponds to the norm of the difference vector \( c = (3, 1) \):

\[
\|c\|_2 = \sqrt{3^2 + 1^2}
\]
With increasing dimensionality, distances grow, too:

- Example: $L_2((1,2),(4,3)) = \sqrt{10}$
- Now double the feature vector length (double the original features): $L_2((1,2,1,2),(4,3,4,3)) = \sqrt{3^2 + 1^2 + 3^2 + 1^2} = \sqrt{20}$
- Effect seems not so important, values might be only in a larger scale?
- NOPE:

  **Contrast of distances is lost in high dimensional data since distances grow more and more alike!**

This is know as the Concentration of Distances problem (see next)
Concentration of Distances

Concentration Phenomenon

• As dimensionality grows, distance values grow, too, such that the (numerical) contrast provided by usual measures decreases or even diminishes

• In other words, the distribution of norms in a given distribution of points tends to concentrate

• Example: Euclidean norm of vectors consisting of several variables that are (assumed to be) independent and identically distributed

\[ \| y \|_2 = \sqrt{y_1^2 + y_2^2 + \ldots + y_d^2} \]

• In high dimensional spaces this norm behaves unexpectedly . . .
Concentration of Distances

Theorem: Concentration of Distances

• Let $y$ be a $d$-dimensional vector $(y_1, \ldots, y_d)$ where all components $y_i (1 \leq i \leq d)$ are independent and identically distributed

• Then the mean and the variance of the Euclidean norm are:

$$
\mu_{\|y\|} = \sqrt{a \cdot d - b + O(d^{-1})} \quad \text{and} \quad \sigma_{\|y\|} = b + O(d^{-1/2})
$$

where $a$ and $b$ are parameters depending only on the central moments of order 1, 2, 3, 4.

Interpretation:

• The norm grows proportionally to $\sqrt{d}$, but the variance remains approx. constant for large $d$ (because $\lim_{d \to \infty} d^{-\mathrm{const}} = 0$)

• With growing dimensionality, the relative error made by taking $\mu_{\|y\|}$ instead of $\|y\|$ becomes negligible

Neighborhood Concept Become Meaningless

Implications from the concentration of distances:

- A lot of data mining methods use distances and neighborhoods to define patterns (e.g. kNN classifier, density-based clustering, distance-based outlier detection, ...)

- Using neighborhoods is based on a key assumption:
  - Objects that are similar to an object o are in its neighborhood
  - Object that are dissimilar to o are not in its neighborhood

- What if all objects are in the same neighborhood?
  - Consider the above effect on distances: kNN distances are almost equal to each other, i.e., the k nearest neighbors are random objects
Neighborhood Concept Become Meaningless

**Definition: Unstable Neighborhood**

- A NN-query is unstable for a given $\varepsilon$ if the distance from the query point to most data points is less than $(1+\varepsilon)$ times the distance from the query point to its nearest neighbor.

- It can be shown that with growing dimensionality, the probability that a query is unstable converges to 1.
Neighborhood Concept Become Meaningless

• Consider a $d$-dimensional query point $q$ and $n$ $d$-dimensional sample points $x_1, \ldots, x_n$ (independent and identically distributed)

• We define:
  
  \[ DMIN_d = \min \{ L_2(x_i, q) \mid 1 \leq i \leq n \} \] (dist to next neighbor)
  
  \[ DMAX_d = \max \{ L_2(x_i, q) \mid 1 \leq i \leq n \} \] (dist to farthest neighbor)

**Theorem**

• If \( \lim_{d \to \infty} \left( \frac{\text{VAR}_{L_2}(x_i, q)}{\mu_{L_2}^2(x_i, q)} \right) = 0 \)

• Then \( \forall \varepsilon > 0 : \lim_{d \to \infty} P(DMAX_d \leq (1 + \varepsilon) DMIN_d) = 1 \)

In other words: if the precondition holds, all points converge to the same distance from the query!

Neighborhood Concept Become Meaningless

Visually: Pairwise distances of a sample of 105 instances drawn from a uniform $[0, 1]$ distribution, normalized $(1/\sqrt{d})$. 
• Be clear about the precondition of the Theorem!!!
• Consider the feature space of $d$ relevant features for a given application (i.e., truly similar objects display small distances in most features)
• Now add $d \cdot c$ additional features being independent of the initial feature space
• With increasing $c$ the distance in the independent subspace will dominate the distance in the complete feature space
• So the question is:
  How many relevant features must be similar to indicate object similarity?
  (or: how many relevant features must be dissimilar to indicate dissimilarity?)
• With increasing dimensionality the likelihood that two objects are similar in every respect gets smaller.
Growing Data Space

- OK, the data space grows with increasing dimensionality
- But what are the problems?
- In low dimensional spaces we have some (intuitive) assumptions on the behavior of volumes (sphere, cube, etc.) and on the distribution of data objects
- However, basic assumptions do not hold in high dimensional spaces:
  - Spaces become sparse or even empty and the probability of one object inside a fixed range tends to become zero
  - Distribution of data has a strange behavior e.g. a normal distribution has only few objects in its center and the tails of distributions become more important

We will have a closer look on these issues ...
Growing Hypotheses Space

- The more features, the larger the hypothesis space
- The lower the hypothesis space is,
  - the easier it is to find the correct hypothesis
  - the less examples you need to properly test hypothesis
Growing Hypotheses Space

- Consider $f$ a unit multivariate normal distribution and normal kernel (KDE)
- The aim is to find an estimate $\hat{f}$ of $f$ at the point 0
- The relative mean square error should be fairly small, e.g.

$$\frac{\mu^2_{\hat{f}(0) - f(0)}}{f(0)^2} < 0.1$$

<table>
<thead>
<tr>
<th>Dim.</th>
<th>Req. sample size to achieve 0.1 error estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>2</td>
<td>19</td>
</tr>
<tr>
<td>5</td>
<td>768</td>
</tr>
<tr>
<td>8</td>
<td>43.700</td>
</tr>
<tr>
<td>10</td>
<td>842.000</td>
</tr>
</tbody>
</table>

Even with only 10 dimensions, we need nearly a million observations to estimate a distribution with an error less than 0.1!!!

Empty Spaces and Tails

• Consider a $d$-dimensional space with partitions of constant size $1/m$

• The number of cells $N$ increases exponentially in $d$: $N = m^d$

• Suppose $x$ points are randomly placed in this space

• In low-dimensional spaces there are few empty partitions and many points per partitions

• In high-dimensional spaces there are far more partitions than points there are many empty partitions

\[
\begin{align*}
&d = 1 \\
&N = 4 \\
&d = 2 \\
&N = 4^2 = 16 \\
&d = 3 \\
&N = 4^3 = 64
\end{align*}
\]
Empty Spaces and Tails

Analogously:

• Consider a simple partitioning scheme, which splits the data in each dimension in 2 halves
• For $d$ dimensions we obtain $2^d$ partitions
• Consider $n = 10^6$ samples in this space
• For $d \leq 10$ such a partition may make sense
• For $d = 100$ there are around $10^{30}$ partitions, so most partitions are empty (given the above $10^6$ points)
Empty Spaces and Tails

• Consider a hyper-cube range query with length $s$ in all dimensions, placed arbitrarily in the data space $[0, 1]^d$.

• $E$ is the event that an arbitrary point lies within the query cube.

• The probability for $E$ is $P(E) = s^d$.

⇒ with increasing dimensionality, even very large hyper-cube range queries are not likely to contain a point.
Empty Spaces and Tails

- The same holds of course for a spherical range query (instead of a cubical range query)
- Consequence: with increasing dimensionality the center of the hyper-cube (or more generally: of the data space) becomes less important and the volume of the data space concentrates in its corners (i.e. randomly distributed points tend to be on the border of the data space . . . )
- This seems to be a distortion of space compared to our 3D way of thinking — and that is actually what it is ...
And that also means, that the tails of a distribution become extremely important

- Consider standard density function \( f \)
- Consider \( \hat{f} \) with

\[
\hat{f}(x) = \begin{cases} 
0 & f(x) < 0.01 \\
 f(x) & \text{else}
\end{cases}
\]

- Rescaling \( \hat{f} \) to a density function will make very little difference in 1D, since very few data points occur in regions where \( f \) is very small
Empty Spaces and Tails

But for high dimensional data:

- More than half of the data has less then $1/100$ of the maximum density $f(0)$ (for $\mu = 0$)
- Example: 10-dimensional Gaussian distribution $X$:

$$
\frac{f(X)}{f(0)} = e^{-\frac{1}{2}X^T X} \approx e^{-\frac{1}{2} \chi^2_{10}}
$$

since the median of the $\chi^2_{10}$ distribution is 9.34, the median of $\frac{f(X)}{f(0)}$ is $e^{-\frac{9.34}{2}} = 0.0094$

- Thus, most objects occur at the tails of the distribution
- In other words, in contrast to the low dimensional case, regions of relatively very low density can be extremely important parts
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- Thus, most objects occur at the tails of the distribution
- In other words, in contrast to the low dimensional case, regions of relatively very low density can be extremely important parts
Empty Spaces and Tails

Example: $(\mu = 0, \sigma = 1)$

- 1D: 90% of the mass of the distribution lies between $-1.6$ and $1.6$
- 10D: 99% of the mass of the distribution is at points whose distance from the origin is greater than $1.6$
- Thus, it is difficult to estimate the density, except for enormous samples because in very high dimensions virtually the entire sample will be in the tails
Further Problems

• Patterns and models on high-dimensional data are often hard to interpret, e.g. long decision rules

• Efficiency in high-dimensional spaces is often limited because e.g. index structures degenerate and distance computations are much more expensive

• There may be different semantic layers so pattern might only be observable in subspaces or projected spaces (cf. the baby shape game)

• Cliques of correlated features dominate the object description
• Summarizing: the higher the dimensionality, the worse is the expected outcome of the mining algorithm (i.e., dimensionality is a curse, says Kröger)
• Well, not in general, the Kernel trick shows the opposite: through the extension of the data space with new attributes, the mining algorithm (e.g. a SVM classifier) gets more accurate (i.e., dimensionality is a blessing, says Tresp in his ML course)
• So: Who is right????????? – Both – What????
• Look at what we assumed for the curse: attributes are independent (and often even uniformly distributed)
• These attributes are likely to be irrelevant for the mining task
• And the blessing: a Kernel (if it works) adds relevant attributes (even more relevant than the original ones)
• Message: high-dimensional data is tricky and the curse can come by as several problems
  • Some are due to irrelevant attributes, so try to get rid of irrelevant attributes and keep the relevant ones
  • Some are instead of relevant attributes, so among the relevant attributes, try to get rid of redundant ones
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Feature Selection

• A task to remove irrelevant and/or redundant features
  • Irrelevant features:
    • Not useful for a given task
    • Probably decrease accuracy
  • Redundant features:
    • Strongly correlated with another relevant feature
    • Does not drop the accuracy, but may drop efficiency, explainability, etc.

• Deleting irrelevant and redundant features can improve the quality as well as the efficiency of the methods and the found patterns.

• New feature space: Delete all useless features from the original feature space.

Keep in mind...

Feature selection ≠ Dimensionality reduction
Feature selection ≠ Feature extraction
Irrelevant and Redundant Features (Unsupervised Case)

Irrelevance

Feature $y$ is irrelevant, because if we omit $x$, we have only one cluster, which is uninteresting.

Redundancy

Features $x$ and $y$ are redundant, because $x$ provides (appr.) the same information as feature $y$ with regard to discriminating the two clusters.

---

Irrelevant and Redundant Features (Supervised Case)

Irrelevance

Feature $y$ separates well the two classes. Feature $x$ is irrelevant. Its addition “destroys” the class separation.

Redundancy

Features $x_1$ and $x_2$ are redundant.

Individually irrelevant together relevant

Source: http://www.kdnuggets.com/2014/03/machine-learning-7-pictures.html
Problem Definition

• **Input:** Vector space \( F = d_1 \times \cdots \times d_n \), dimensions \( D = \{ d_1, \ldots, d_n \} \).
• **Output:** a minimal subspace \( M \) over dimensions \( D' \subseteq D \) which is optimal for a given data mining task.
  - Minimality increases the efficiency, reduces the effects of the curse of dimensionality and increases interpretability.

**Challenges:**

• Optimality depends on the given task.
• There are \( 2^d \) possible solution spaces (exponential complexity)
• This search space is similar to the frequent itemset mining problem, but:
  • There is often no monotonicity in the quality of subspace (which is important for efficient searching)
  • Features might only be useful in combination with other certain features.

\[ \Rightarrow \] For many popular criteria, feature selection is an exponential problem.
\[ \Rightarrow \] Most algorithms employ search heuristics.
Two Main Components (Steps)

1. Feature subset generation
   - Single dimensions
   - Combinations of dimensions (subspaces)

2. Feature subset evaluation
   - Importance scores like information gain, $\chi^2$
   - Performance of a learning algorithm

⇒ How to select/evaluate features? How to traverse the search space?
1. Filter methods
   – Explores the general characteristics of the data, independent of the learning algorithm.

2. Wrapper methods
   – The learning algorithm is used for the evaluation of the subspace.

3. Embedded methods
   – The feature selection is part of the learning algorithm.
Feature Selection/Evaluation Methods

• Filter methods
  – Basic idea: assign an “importance” score to each feature to filter out useless ones
  – Examples: information gain, $\chi^2$-statistic, TF-IDF for text...
  – Disconnected from the learning algorithm.
  – Pros:
    ○ Fast and generic
    ○ Simple to apply
  – Cons:
    ○ Doesn’t take into account interactions between features
    ○ Individually irrelevant features, might be relevant together
    ○ Too generic?
• Wrapper methods
  – A learning algorithm is employed and its performance is used to determine the quality of selected features.
  – Pros:
    ○ take feature dependencies into account
    ○ interaction between feature subset search and model selection
  – Cons:
    ○ higher risk of overfitting than filter techniques
    ○ very computationally intensive, especially if building the classifier has a high computational cost.
• Embedded methods
  – Such methods integrate the feature selection in model building
  – Example: decision tree induction algorithm: at each decision node, a feature has to be selected.
  – Pros:
    ◦ less computationally intensive than wrapper methods.
  – Cons:
    ◦ specific to a learning method
Search Strategies in the Feature Space

- **Forward selection**
  - Start with an empty feature space and add relevant features
- **Backward selection**
  - Start with all features and remove irrelevant features
- **Branch-and-bound**
  - Find the optimal subspace under the monotonicity assumption
- **Randomized**
  - Randomized search for a $k$ dimensional subspace
- ...
Kapitel 3: Feature Selection

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General Idea

**Input**

- Target dimensionality $k \leq d$
- Training set of $n$-dimensional feature vectors with features $d_1, d_2, \ldots, d_n$ and target variable $C$

**General Approach**

- Compute the quality $q(d_i, C)$ for each dimension $d_i \in \{d_1, \ldots, d_n\}$ to predict the correlation to $C$
- Sort the dimensions $d_1, \ldots, d_n$ w.r.t. $q(d_i, C)$
- Select the best $k$ dimensions

**Basic Assumption**

- Attribute independence (no correlations between features)
General Idea

Key Concept

- Quality of feature $d_i$: How suitable is the feature for predicting the value of class attribute $C$?
- Statistical measures
  - Rely on distributions over feature values and target values
  - How strong is the correlation between both value distributions?
  - How good does splitting the values in the feature space separate values in the target dimension?
Quality of Features

How to measure the distribution?

- For discrete values: determine probabilities for all value pairs.
- For real valued features:
  - Discretize the value space (reduction to the case above)
  - Use probability density functions (e.g. uniform, Gaussian,..)
- Example quality measures:
  - Information Gain
  - Chi-square $\chi^2$-statistics
  - Mutual Information
Quality of Features: Entropy

- Idea: Evaluate class discrimination in each dimension (Used in ID3 algorithm for decision trees)
- It uses entropy, a measure of pureness of the data set $S$ w.r.t. the class labels $c_i \in C$

\[
\text{Entropy}(S) = \sum_{c_i \in C} -p_{c_i} \cdot \log_2(p_{c_i})
\]

where $p_{c_i}$ is the relative frequency of class $c_i$ in $S$
Example

- Let $S$ be a collection of positive and negative examples for a binary classification problem, i.e., $C = \{+, -\}$
- Then $\text{Entropy}(S) = -p_+ \log_2(p_+) - p_- \log_2(p_-)$
  - $p_+$ is the percentage of positive examples in $S$
  - $p_-$ is the percentage of negative examples in $S$
- Example splits:
  - Let $S : [9+, 5-]$: $\text{Entropy}(S) = -\frac{9}{14} \log_2(\frac{9}{14}) - \frac{5}{14} \log(\frac{5}{14}) = 0.940$
  - Let $S : [7+, 7-]$: $\text{Entropy}(S) = -\frac{7}{14} \log_2(\frac{7}{14}) - \frac{7}{14} \log(\frac{7}{14}) = 1$
  - Let $S : [14+, 0-]$: $\text{Entropy}(S) = -\frac{14}{14} \log_2(\frac{14}{14}) - \frac{0}{14} \log(\frac{0}{14}) = 0$
- Obviously: Entropy is 0, when all samples belong to the same class while Entropy is 1, when there is an equal number of samples in all splits
Quality of Features: Information Gain

• The information gain $Gain(S, d_i)$ of a feature $d_i$ relative to a training set $S$ measures the gain reduction in $S$ due to splitting on $d_i$, i.e., the entropy of the data set $S$ before splitting minus the weighted sum of the entropies of all splits $S_j$ in a given feature $d_i$:

\[
Gain(S, d_i) = Entropy(S) - \sum_{S_j} \frac{|S_j|}{|S|} \cdot Entropy(S_j)
\]

• For nominal attributes: use attribute values for splitting, i.e. each possible value $v_j$ in $d_i$ defines one split and $S_j$ contains all objects having $v_j$ in $d_i$

• For real valued attributes: Determine a splitting position $\nu$ in the value set and split e.g. into $S_1$ containing all objects with values $\leq \nu$ and $S_2$ containing all objects with values $> \nu$ in $d_i$
Example

- Which dimension, “Humidity” or “Wind”, is better?

\[
\text{Gain (S, Humidity)} = 0.940 - 0.985 - 0.592 = 0.151
\]

- Larger values are better!
Quality of Features: Chi-square Statistics

- Idea: Measures the independence of a feature $d$ from the class variable $C$
- Contingency table: divide data based on a split value $s$ or based on discrete values
- Example: Does “liking science fiction movies” imply “playing chess”?

### Example Table

<table>
<thead>
<tr>
<th>Predictor attribute</th>
<th>Class attribute</th>
<th>Play chess</th>
<th>Not play chess</th>
<th>Sum (row)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Like science fiction</td>
<td>Play chess</td>
<td>250</td>
<td>200</td>
<td>450</td>
</tr>
<tr>
<td></td>
<td>Not play chess</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Not like science fiction</td>
<td>Play chess</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Not play chess</td>
<td>50</td>
<td>1000</td>
<td>1050</td>
</tr>
<tr>
<td>Sum(col.)</td>
<td></td>
<td>300</td>
<td>1200</td>
<td>1500</td>
</tr>
</tbody>
</table>

### Chi-square $\chi^2$ test

$$
\chi^2 = \sum_{i=1}^{C} \sum_{j=1}^{Values(d)} \frac{(o_{ij} - e_{ij})^2}{e_{ij}}
$$

- $o_{ij}$: observed freq. of value $j$ in class $i$
- $e_{ij}$: expected freq. of value $j$ in class $i$
Example

- Compute the $\chi^2$ values for the following table (numbers in parenthesis are expected counts calculated based on the data distribution in the two categories)

<table>
<thead>
<tr>
<th>Predictor attribute</th>
<th>Class attribute</th>
<th>Play chess</th>
<th>Not play chess</th>
<th>Sum (row)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Like science fiction</td>
<td></td>
<td>250 (90)</td>
<td>200 (360)</td>
<td>450</td>
</tr>
<tr>
<td>Not like science fiction</td>
<td></td>
<td>50 (210)</td>
<td>1000 (840)</td>
<td>1050</td>
</tr>
<tr>
<td>Sum(col.)</td>
<td></td>
<td>300</td>
<td>1200</td>
<td>1500</td>
</tr>
</tbody>
</table>

\[
\chi^2 = \frac{(250 - 90)^2}{90} + \frac{(50 - 210)^2}{210} + \frac{(200 - 360)^2}{360} + \frac{(1000 - 840)^2}{840} = 507.93
\]

- Smaller values are better!
• In general, the Mutual Information (MI) between two variables $x$ and $y$ measures how much knowing one of these variables reduces uncertainty about the other.

• In our case, it measures how much information a feature contributes to making the correct classification decision, i.e., $x$ is the dimension $d_i$ we want to evaluate and $y$ is the class variable $C$.

• MI is based on probability distributions:
  • $p(x)$ and $p(y)$ are the marginal probability distributions of $x$ and $y$, respectively
  • $p(x, y)$ is the joint probability distribution function
Quality of Features: Mutual Information

• Discrete case

\[ MI(x, y) = \sum_{x_i \in x} \sum_{y_i \in y} p(x_i, y_i) \cdot \log \frac{p(x_i, y_i)}{p(x_i)p(y_i)} \]

• Continuous case

\[ MI(x, y) = \int_x \int_y p(x, y) \cdot \log \frac{p(x, y)}{p(x)p(y)} \, dx \, dy \]

• Interpretation: if \( x \) and \( y \) are statistically independent, then
  • \( p(x, y) = p(x) \cdot p(y) \) and, thus, \( \log(1) = 0 \)
  • Or in other words: knowing \( x \) does not reveal anything about \( y \)
Advantages

- Efficiency: it compares each feature \(\{d_1, d_2, \ldots, d_n\}\) separately to the class attribute \(C\) (and takes the best \(k\)) instead of testing \(\binom{n}{k}\) subspaces
- Works already for rather small sample sizes

Limitations

- Independency assumption: Classes and features must display a direct correlation
- In case of correlated features: Always selects the features having the strongest direct correlation to the class variable, even if the features are strongly correlated with each other
1. Introduction to Feature Spaces

2. Challenges of High Dimensional Data

3. Supervised Feature Selection
   3.1 Forward Selection and Feature Ranking
   3.2 Backward Elimination and Random Subspace Selection
   3.3 Subspace Projections

4. Feature Reduction and Metric Learning

5. Clustering High Dimensional Data
**General Approach**

- Start with the complete feature space and delete redundant features
- Greedy Backward Elimination
  1. Generate the subspaces $R$ of the feature space $F$
  2. Evaluate subspaces $R$ with the quality measure $q(R)$
  3. Select the best subspace $R^*$ w.r.t. $q(R)$
  4. If $R^*$ has the target dimensionality, terminate else start backward elimination on $R^*$.

**Remarks**

- Useful in supervised and unsupervised setting (in the latter scenario, $q(R)$ measures structural characteristics)
- Greedy search if there is no monotonicity on $q(R)$; for monotonous measures, branch and bound can be employed
Supervised Quality Measure: Distance-based

- Idea: Subspace quality can be evaluated by the distance between the within-class nearest neighbor and the between-classes nearest neighbor.

- Quality criterion: For each object $o$ from the data set $S$, compute distance to the closest object having the same class $NN_{c_i=C(o)}^R(o)$ (within-class nearest neighbor distance) in subspace $R$, and to the closest object belonging to another class $NN_{c_j\neq C(o)}^R(o)$ (between-classes nearest neighbor distance), where $C(o)$ denotes the class label of object $o$ in subspace $R$:

\[
q(R) = \frac{1}{S} \cdot \sum_{o \in S} \frac{NN_{c_j\neq C(o)}^R(o)}{NN_{c_i=C(o)}^R(o)}
\]

- Remark: $q(R)$ is not monotonous: by deleting a dimension, the quality can increase or decrease.
Supervised Quality Measure: Model-based

• Idea: Directly employ the data mining algorithm to evaluate the subspace, e.g. by training a Naive Bayes classifier

• Practical aspects:
  • Success of the data mining algorithm must be measurable (e.g. class accuracy)
  • Runtime for training and applying the classifier should be low
  • The classifier parameterization should not be of great importance
  • Test set should have a moderate number of instances
Backward Elimination: Discussion

Advantages

- Considers complete subspaces (multiple dependencies are used)
- Can recognize and eliminate redundant features

Limitations

- Tests w.r.t. subspace quality usually requires much more effort
- All solutions employ heuristic greedy search which do not necessarily find the optimal feature space
Branch and Bound: General Idea

General Approach

• Given: A classification task over the feature space \( F \)
• Aim: Select the \( k \) best dimensions to learn the classifier
• Backward elimination approach “Branch and Bound” is guaranteed to find the optimal feature subset under the monotonicity assumption
• The monotonicity assumption states that for two feature subsets \( X, Y \in F \) and a feature selection criterion \( J \), if \( X \subset Y \) then
  • \( J(X) \leq J(Y) \) if \( J \) is maximized
  • \( J(X) \geq J(Y) \) if \( J \) is minimized
• Branch and Bound starts from the full set \( F \) and removes features using a depth-first strategy
• Nodes whose objective function are smaller (greater) than the current best are not explored since the monotonicity assumption ensures that their children will not contain a better solution
Example: Original dimensionality 4, <A,B,C,D>. Target dimensionality \( d = 1 \).

- selected feature
- removed feature

\[(\text{All}) = 1.0\]

- selected feature
- removed feature

A: $J(\text{A}) = 0.715$
B: $J(\text{B}) = 0.421$
C: $J(\text{C}) = 0.603$
D: $J(\text{D}) = 1.0$

(A) = 1.0

- Blue circle: selected feature
- Gray circle: removed feature

\[(\text{All})=1.0\]

- \(J(BCD)=1.0\)
- \(J(ACD)=0.715\)
- \(J(ABD)=0.421\)
- \(J(ABC)=0.603\)

- \(J(CD)=0.815\)
- \(J(BD)=0.5\)
- \(J(BC)=0.5\)
Example: Original dimensionality 4, <A,B,C,D>. Target dimensionality \( d = 1 \).

- **Selected feature**
- **Removed feature**

Diagram:

```
(All)=1.0

A: J(ACD)=0.715
B: J(ABD)=0.421
C: J(ABC)=0.603
D: J(BCD)=1.0

J(CD)=0.815
J(BD)=0.5
J(BC)=0.5

J(D)=0.62
J(C)=0.43
```

aktuBound = 0.62

Selected features: A, B, C, D

Removed features: None

Prune with $aktBound = 0.62$

- Blue: selected feature
- Grey: removed feature

\[ J(BCD) = 1.0 \]
\[ J(ACD) = 0.715 \]
\[ J(ABD) = 0.421 \]
\[ J(ABC) = 0.603 \]
\[ J(CD) = 0.815 \]
\[ J(BD) = 0.5 \]
\[ J(BC) = 0.5 \]
\[ J(AD) = 0.5 \]
\[ J(AC) = 0.32 \]
\[ J(D) = 0.62 \]
\[ J(C) = 0.43 \]

aktBound = 0.62
Subspace Inconsistency (IC)

- Given a data set $S$ (works best for categorical data)
- Idea: Having identical vectors $u, v$ ($u_i = v_i, 1 \leq i \leq d$) in subspace $R$ but the class labels are different ($C(u) \neq C(v)$), this subspace displays an inconsistent labeling
- Measuring the inconsistency of a subspace $R$
  - $X_R(u)$: Amount of all identical vectors $u$ in $R$
  - $X_R^c(u)$: Amount of all identical vectors $u$ in $R$ having class label $c \in C$
  - Inconsistency of $u$ in $R$: $IC_R(u) = X_R(u) - \max_{c \in C} X_R^c(u)$

Then, inconsistency of subspace $R$ is

$$IC(R) = \frac{\sum_{u \in S} IC_R(u)}{|S|}$$

- Monotonicity: $R_1 \subset R_2 \Rightarrow IC(R_1) \geq IC(R_2)$
Advantages

• Monotonicity allows efficient search for optimal solutions
• Well-suited for binary or discrete data (identical vectors are very likely with decreasing dimensionality)

Limitations

• Useless without groups of identical features (real-valued vectors)
• Worse-case runtime complexity remains exponential in the number of features $d$
Random Subspace Selection: General Idea

General Approach

- Idea: Select $n$ random subspaces having the target dimensionality $k$ out of the $\binom{d}{k}$ many possible subspaces and evaluate each of them.
- Needs quality measures for complete subspaces.
- Trade-off between quality and effort depends on $n$.
- Good alternative to forward selection if quality measure is not monotonic.

- Different randomization approaches exist (see next subsection):
  - Genetic algorithms
  - $k$-medoids feature clustering
  - ...

Prof. Dr. Peer Kröger: KDD2 (SoSe 2019) — Lecture 2 — High Dimensional Data — 3. Feature Selection
1. Introduction to Feature Spaces

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Genetic Algorithms: General Idea

General Approach

• Idea: Randomized search through genetic algorithms
• Genetic Algorithms encode individual states in the search space as bit-strings
• Population (of current solutions) is a subset of all possible $k$-dimensional subspaces
• Fitness function: quality measure for a subspace
• Algorithmic schema to find the best solution in the search space by mixing/changing the population in each iteration (stops e.g. if the best solution of the current population is less fit than the best solution in the previous population)
• Each iteration manages a specific population from which the next population is obtained
Genetic Algorithms: Population Generation

- Operators on the population ($k$-dim subspaces) to create candidates for the next population:
  - Mutation: dimension $d_i$ in subspace $R$ is replaced by dimension $d_j$ with a likelihood of $x\%$
  - Crossover: combine two subspaces $R_1$ and $R_2$, i.e., unite the features sets of $R_1$ and $R_2$ and delete random dimensions until dimensionality is $k$ again
- Selection for next population: All subspaces having at least a quality of $y\%$ of the best fitness in the current generation are copied to the next generation
- Free tickets: Additionally each subspace is copied into the next generation with a probability of $u\%$
- Remark: Many variants on the basic algorithmic schema, e.g. different operations, efficient convergence by “Simulated Annealing” (likelihood of free tickets decreases with the iterations), ...
Genetic Algorithms: Discussion

Advantages

• Can escape from local optima during the search
• Often good approximations of the optimal solutions

Limitations

• Runtime (is not bounded (in the original schema)
• Configuration depends on many parameters which have to be tuned to achieve good quality results in efficient time
Feature Clustering: General Idea

General Approach

- Given: A feature space $F$ and an unsupervised data mining task
- Target: Reduce $F$ to a subspace of $k$ (original) dimensions while reducing redundancy
- Idea: Cluster the features in the space of objects and select one representative feature for each of the clusters (this is equivalent to clustering in a transposed data matrix)
- Problem: often many more samples than features so transposed data matrix has many more features than samples
Feature Clustering: Example

- Typical example: item-based collaborative filtering
- E.g. features 3 and 4 are similar over all persons so they could be “merged” to one feature

<table>
<thead>
<tr>
<th></th>
<th>1 (Titanic)</th>
<th>2 (Braveheart)</th>
<th>3 (Matrix)</th>
<th>4 (Inception)</th>
<th>5 (Hobbit)</th>
<th>6 (300)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Susan</td>
<td>5</td>
<td>2</td>
<td>5</td>
<td>5</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>Bill</td>
<td>3</td>
<td>3</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Jenny</td>
<td>5</td>
<td>4</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>Tim</td>
<td>2</td>
<td>2</td>
<td>4</td>
<td>5</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Thomas</td>
<td>2</td>
<td>1</td>
<td>3</td>
<td>4</td>
<td>1</td>
<td>4</td>
</tr>
</tbody>
</table>
Feature Clustering: Example

• Work around for the “many features” problem: specialized feature similarity measures, e.g.
  • Cosine similarity
  • Pearson correlation

• Algorithmic schema
  • Cluster features with a $k$-medoid clustering method based on correlation
  • Select the medoids to span the target data space

• Remark
  • For group/cluster of dependent features there is one representative feature
  • Other clustering algorithms could be used as well, e.g. approximate clustering methods for performance reasons
# Feature Clustering: Discussion

## Advantages

- Depending on the clustering algorithm quite efficient
- Unsupervised method

## Limitations

- Results are usually not deterministic (partitioning clustering results depend on initialization)
- Representatives are usually unstable for different clustering methods and parameters
- Method captures pairwise correlations and dependencies among features but multiple dependencies are not considered
Summary: Feature Selection

- Forward-Selection examines each dimension separately and selects the $k$-best to span the target space
  - Greedy Selection based on Information Gain, $\chi^2$ statistics or Mutual Information
- Backward-Elimination start with the complete feature space and successively remove the worst dimensions
  - Greedy Elimination with model-based and nearest-neighbor based approaches
  - Branch and Bound Search (monotonicity required!) based on inconsistency
- $k$-dimensional Projections directly search in the set of $k$-dimensional subspaces for the best suited
  - Genetic algorithms (any quality measures possible, e.g. those from backward elimination)
  - Feature clustering based on correlation
Discussion: Feature Selection

- Many algorithms based on different heuristics
- There are two reasons to delete features:
  - Redundancy: Features can be expressed by other features
  - Missing correlation to the target variable
- Often even approximate results are capable of increasing efficiency and quality in a data mining task
- Caution: Selected features need not to have a causal connection to the target variable, but both might depend on the same mechanisms in the data space (hidden variables)
- Different indicators to consider in the comparison of before and after selection performance, e.g. model performance, time, dimensionality, ...
Feature Selection — Further Readings


• A.Blum and P. Langley: Selection of Relevant Features and Examples in Machine Learning, Artificial Intelligence (97), 1997.


1. Introduction to Feature Spaces

2. Challenges of High Dimensional Data

3. Supervised Feature Selection

4. Feature Reduction and Metric Learning
   4.1 Reference Point Embedding
   4.2 Principle Component Analysis (PCA)
   4.3 Singular Value Decomposition (SVD)
   4.4 Kernel PCA
   4.5 Further Measures
5. Clustering High Dimensional Data
Overview

• Idea: Instead of removing features, try to find a low dimensional feature space generating the original space as accurate as possible:
  • Redundant features are summarized
  • Irrelevant features are weighted by small values or are “erased” (in the best case of course, the new feature space should contain no irrelevant features anymore)

• Some sample methods (among lots of others):
  • Reference point embedding
  • Principal component analysis (PCA)
  • Singular value decomposition (SVD)
  • Fischer-Faces (FF) and Relevant Component Analysis (RCA)
  • Large Margin Nearest Neighbor (LMNN)
• **Goal**: Describe data with fewer features (reduce number of columns)
• Be clear: (like in feature selection) there will always be an information loss

• There are supervised and unsupervised methods
Kapitel 4: Feature Reduction and Metric Learning

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   4.5 Further Measures

5. Clustering High Dimensional Data
General Approach

• Idea: Describe the position of each object by their distances to a set of reference points

• Given: Vector space \( F = D_1 \times \ldots \times D_n \) where \( D = \{ D_1, \ldots, D_n \} \)

• Target: A \( k \)-dimensional space \( R \) which yields optimal solutions for a given data mining task

• Method: For each reference point \( R = \{ r_1, \ldots, r_k \} \) and a distance measure \( \text{dist} \), transform vector \( x \in F \) as follows:

\[
r_R(x) = \begin{pmatrix}
\text{dist}(r_1, x) \\
\vdots \\
\text{dist}(r_k, x)
\end{pmatrix}
\]
Diskussion

• Distance measure is usually determined by the application
• Selection of reference points can be important (use centroids of the classes or cluster-centroids, points on the margin of the data space, use random samples, ...)

**Advantages**

• Simple approach which is easy to implement
• The transformed vectors yields lower and upper bounds of the exact distances (What the hell is that good for???)

**Disadvantages**

• Even using $d$ reference points does not reproduce a $d$-dimensional feature space
• Selecting good reference points is important but very difficult
Kapitel 4: Feature Reduction and Metric Learning

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   4.4 Kernel PCA
   4.5 Further Measures

5. Clustering High Dimensional Data
Motivation

- Consider the grades of students in Physics and Statistics
- If we want to compare among the students, which grade should be more discriminative? Statistics or Physics?

Answer:
Physics because the variation along that axis is larger

Source: http://astrostatistics.psu.edu/su09/lecturenotes/pca.html
Motivation

• Suppose now the plot looks as below
• What is the best way to compare students now?

Answer:
We should take a linear combination of the two grades (that represents the direction of highest variance) to get the best results

Source: http://astrostatistics.psu.edu/su09/lecturenotes/pca.html
Introduction

Motivation

- PCA returns two principal components
- The first gives the direction of the maximum spread of the data.
- The second gives the direction of maximum spread perpendicular to the first

Source: http://astrostatistics.psu.edu/su09/lecturenotes/pca.html
A feature $X$ can be normalized by subtracting its values with the mean $\bar{X}$ and dividing by the standard deviation $s_X$, e.g. $\tilde{X} = \frac{X - \bar{X}}{s_X}$.

**Example:**

Consider the following body heights measured in different units:

<table>
<thead>
<tr>
<th></th>
<th>Person A</th>
<th>Person B</th>
<th>Person C</th>
<th>mean</th>
<th>sd</th>
</tr>
</thead>
<tbody>
<tr>
<td>body height (cm)</td>
<td>180.00</td>
<td>172.00</td>
<td>175.00</td>
<td>175.67</td>
<td>4.04</td>
</tr>
<tr>
<td>body height (m)</td>
<td>1.80</td>
<td>1.72</td>
<td>1.75</td>
<td>1.76</td>
<td>0.04</td>
</tr>
<tr>
<td>body height (feet)</td>
<td>5.91</td>
<td>5.64</td>
<td>5.74</td>
<td>5.76</td>
<td>0.13</td>
</tr>
</tbody>
</table>

After normalizing, we always obtain the normalized body height (no matter which unit we used):

<table>
<thead>
<tr>
<th></th>
<th>Person A</th>
<th>Person B</th>
<th>Person C</th>
<th>mean</th>
<th>sd</th>
</tr>
</thead>
<tbody>
<tr>
<td>normalized body height</td>
<td>1.07</td>
<td>-0.91</td>
<td>-0.16</td>
<td>0.00</td>
<td>1.00</td>
</tr>
</tbody>
</table>
Normalizing all features in a data set, can have several advantages:

- It puts all features into *comparable* units, i.e., we make sure that all normalized features have mean 0 and standard deviation of 1
- It can avoid numerical instabilities in several algorithms, e.g. if a feature has very low / high values
- It helps in computing meaningful *distances* between observations
- Finally, if we want to find directions of highest variances, it might be better to do this on normalized data
Normalizing: Covariance vs. Correlation

• Sure, there are many ways to do normalization
• here, we will use the notion common in Statistics, where the variance of a normalized feature is always 1, its mean is always 0
• The covariance of two normalized features \( \tilde{X} = \frac{X - \bar{X}}{s_X} \) and \( \tilde{Y} = \frac{Y - \bar{Y}}{s_Y} \) is the same as the correlation of the non-normalized features \( X \) and \( Y \).
• One can proof this with the help of

\[
s_{\tilde{X}\tilde{Y}} = \frac{1}{n-1} \sum_{i=1}^{n} (\tilde{x}_i - \bar{\tilde{x}})(\tilde{y}_i - \bar{\tilde{y}}) = \ldots = \frac{1}{n-1} \sum_{i=1}^{n} \frac{(x_i - \bar{x})}{s_X} \frac{(y_i - \bar{y})}{s_Y} = r_{XY}.
\]
Example I:

- Feature $x_1$ explains most of the variation
- Feature $x_2$ has a lower variance than $x_1$
- If we disregard $x_2$ and project the points into the 1-dimensional space of $x_1$, we do not lose much information w.r.t. variability
**Example II:**

- $x_1$ and $x_2$ are correlated and have similar variances.
- Find a new orthogonal axes (e.g. PC1 and PC2), where PC1 explains most of the variation.
- Rotate the points and consider PC1 and PC2 as new coordinate system (situation as in the previous example).
- We can now project points onto PC1 and disregard PC2 (hopefully without losing much information).
PCA Intuition

• PCA finds the optimal rotation such that the transformed data explains the variability of the data best
• The new axis are the principal components (also called “eigenvectors” because PCA is technically an Eigen-Decomposition); for a $d$-dimensional data set we always get $d$ principal components
• The variance along each eigenvector (called “eigenvalue”) is decreasing, i.e. the first eigenvector has the highest eigenvalue, while the $d$-th eigenvector has the smallest eigenvalue
• This can be used for dimensionality reduction: if we pick the $k$-th first eigenvectors as new axes and transform the $d$-dimensional data into the new $k$-dimensional space, this transformation is optimal w.r.t. loss of total variance
PCA Procedure

**General procedure**

1. Rotate the original \( p \)-dimensional coordinate system until the first PC that explains most of the variation is found.

2. Fix the first PC and proceed with rotating the remaining \( p - 1 \) coordinates until the second PC (which is orthogonal to the first PC) is found that explains most of the *remaining* variation, etc.

3. We can reduce the dimensions by projecting the points onto the first, say \( k < p \), PC.
PCA Intuition: Find first PC
Variance of projected points: 0.87

- rotated coordinate system
- original coordinate system
- projected points
PCA Intuition: Animation

Variance of projected points: 0.25

rotated coordinate system
original coordinate system
projected points
PCA Intuition: Animation

Variance of projected points: 0.08

rotated coordinate system
---
original coordinate system

projected points
PCA Intuition: Animation

Variance of projected points: 0.38

- rotated coordinate system
- original coordinate system
- projected points
PCA Intuition: Animation

Variance of projected points: 0.84

- rotated coordinate system
- original coordinate system
- projected points
PCA Intuition: Animation

Variance of projected points: 1.31

- rotated coordinate system
- original coordinate system
- projected points
Variance of projected points: 1.63

- rotated coordinate system
- original coordinate system
- projected points
PCA Intuition: Reduce dimensionality

Rotate the points and use PC1 and PC2 as new coordinate system.

Here, the PC1 axis explains most of the variance:
Dimensionality can be reduced by projecting the points onto the PC1 (and by disregarding PC2). The hope is that we won’t lose much information this way.
Idea: Transform an original set of correlated metric features to a new set of uncorrelated (orthogonal) metric features, called principal components (PC), that explain the variability in the data.

- The objective is to investigate if only a few PC account for most of the variability in the original data.
- If the objective is fulfilled, we can use fewer PCs to reduce the dimensionality.
- The PCs remove collinearity of the input variables as they are orthogonal to each other.
PCA Intuition: Final Remarks

- PCA is used for dimensionality reduction by disregarding dimensions with lower variability.
- There is always an information loss, especially for other criteria.
- **Attention**: dimensionality reduction can worsen the classification accuracy when the task is to classify two groups:
Aim: Find a new set of features (PC scores, eigenvectors) $\mathbf{pc}_1, \ldots, \mathbf{pc}_p$ based on the original data $\mathbf{X} = [\mathbf{x}_1, \ldots, \mathbf{x}_p]$ so that

- each PC score $\mathbf{pc}_1, \ldots, \mathbf{pc}_p$ is a linear combination of the original metric features with coefficient weights (so-called loading vectors) $\mathbf{a}_1, \ldots, \mathbf{a}_p$, i.e.

$$\mathbf{pc}_j = a_{j1}\mathbf{x}_1 + a_{j2}\mathbf{x}_2 + \ldots + a_{jp}\mathbf{x}_p = \mathbf{Xa}_j.$$  

- the set is mutually uncorrelated: $\text{Cov}(\mathbf{pc}_j, \mathbf{pc}_k) = 0$, $\forall j \neq k$.
- the variances (eigenvalues) of the PC scores decrease:

$$\lambda_1 > \lambda_2 > \ldots > \lambda_p,$$  

where $\lambda_k := \text{Var}(\mathbf{pc}_k)$. 

We look for the loading vector \( a_1 = (a_{11}, a_{21}, \ldots, a_{p1})^\top \) that maximizes the variance of \( pc_1 \):

\[
\max_{a_1} \ Var(pc_1) = Var(Xa_1) = a_1^\top \Sigma a_1
\]

subject to the normalization constraint \( a_1^\top a_1 = \sum_{k=1}^{p} a_{k1}^2 = 1 \).

The constraint is required for identifiability reasons, otherwise we could maximize the variance by just increasing the values in \( a_1 \).

Repeat this maximization step for the other PCs and additionally use the orthogonality constraint, i.e. for the second PC:

\[
a_2^\top a_1 = 0.
\]
Example: The Olympic Heptathlon Data

The heptathlon data set (e.g. available in the R package HSAUR3) contains the competition results of 25 athletes in 7 disciplines for the Olympics held in Seoul in 1988.

- **Aim**: Rank the athletes according to their overall performance in all 7 disciplines.
- **Idea**: Use PCA to reduce the dimensionality (i.e., reduce the results of the 7 disciplines to one dimension) and compare the scores of the first PC with the official scores.
Example: The Olympic Heptathlon Data

Features of the heptathlon data:

- **hurdles**: results 100m hurdles (in seconds).
- **highjump**: results high jump (in m).
- **shot**: results shot putt (in m).
- **run200m**: results 200m race (in seconds).
- **longjump**: results long jump (in m).
- **javelin**: results javelin (in m).
- **run800m**: results 800m race (in seconds).
- **score**: total score of the official scoring system.
Example: The Olympic Heptathlon Data

The features hurdles, run200m and run800m are time measurements, i.e. low values are better. For all other features high values are better.

Results of the best and worst participant:

<table>
<thead>
<tr>
<th></th>
<th>hurdles</th>
<th>highjump</th>
<th>shot</th>
<th>run200m</th>
<th>longjump</th>
<th>javelin</th>
<th>run800m</th>
<th>score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Joyner-Kersee (USA)</td>
<td>12.7</td>
<td>1.86</td>
<td>15.8</td>
<td>22.6</td>
<td>7.27</td>
<td>45.7</td>
<td>129</td>
<td>7291</td>
</tr>
<tr>
<td>Launa (PNG)</td>
<td>16.4</td>
<td>1.50</td>
<td>11.8</td>
<td>26.2</td>
<td>4.88</td>
<td>46.4</td>
<td>163</td>
<td>4566</td>
</tr>
</tbody>
</table>

We use negative time measurements so that higher values are better and therefore all features have the same direction:

<table>
<thead>
<tr>
<th></th>
<th>hurdles</th>
<th>highjump</th>
<th>shot</th>
<th>run200m</th>
<th>longjump</th>
<th>javelin</th>
<th>run800m</th>
<th>score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Joyner-Kersee (USA)</td>
<td>-12.7</td>
<td>1.86</td>
<td>15.8</td>
<td>-22.6</td>
<td>7.27</td>
<td>45.7</td>
<td>-129</td>
<td>7291</td>
</tr>
<tr>
<td>Launa (PNG)</td>
<td>-16.4</td>
<td>1.50</td>
<td>11.8</td>
<td>-26.2</td>
<td>4.88</td>
<td>46.4</td>
<td>-163</td>
<td>4566</td>
</tr>
</tbody>
</table>
Scatter Plot Matrix

<table>
<thead>
<tr>
<th></th>
<th>hurdles</th>
<th>highjump</th>
<th>shot</th>
<th>run200m</th>
<th>longjump</th>
<th>javelin</th>
<th>run800m</th>
</tr>
</thead>
<tbody>
<tr>
<td>hurdles</td>
<td>0.811</td>
<td>0.651</td>
<td>0.774</td>
<td>0.912</td>
<td>0.00776</td>
<td>0.779</td>
<td></td>
</tr>
<tr>
<td>highjump</td>
<td>0.441</td>
<td>0.488</td>
<td>0.782</td>
<td>0.00215</td>
<td>0.591</td>
<td></td>
<td></td>
</tr>
<tr>
<td>shot</td>
<td>0.683</td>
<td>0.743</td>
<td>0.269</td>
<td>0.033</td>
<td>0.42</td>
<td></td>
<td></td>
</tr>
<tr>
<td>run200m</td>
<td>0.817</td>
<td>0.333</td>
<td>0.617</td>
<td>0.7</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>longjump</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>−0.02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>javelin</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>run800m</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Perform PCA

- If features are on very different scales, PCA should be carried out on the correlation matrix (which is equivalent to the correlation matrix if normalized features are used).
- As the features of the heptathlon data are on different scales, we perform the PCA based on the correlation matrix.
- Alternatively, we could also perform the PCA based on the covariance matrix but on the normalized heptathlon data.
- The result contains:
  - The loadings $a_1, \ldots, a_p$,
  - The PC scores $pc_1, \ldots, pc_p$ and
  - The variance $\lambda_1, \ldots, \lambda_p$ (or standard deviation) of the PC scores.
The total variance of the $p$ PC scores is equal the total variance of the original features, i.e.,

$$\sum_{j=1}^{p} \lambda_j = s_1^2 + s_2^2 + \cdots + s_p^2,$$

where $\lambda_j$ is the variance of the $j$th PC and $s_j^2$ is the sample variance of variable $x_j$.

The proportion of explained variance of the $j$-th PC is

$$\frac{\lambda_j}{\sum_{j=1}^{p} \lambda_j}.$$

The first $k$ PCs account for a proportion

$$\frac{\sum_{j=1}^{k} \lambda_j}{\sum_{j=1}^{p} \lambda_j}.$$
Choosing the Number of PCs

Two simple rules of thumb for choosing the number of PCs:

1. Retain the first $k$ components, which explain a large proportion of the total variation, e.g., 80-90%.

2. Use a scree plot: Plot the component variances vs. the component number and look for an *elbow*. For components after the *elbow*, the variance decreases more slowly.

![Scree plot](image-url)
PC Scores vs. Official Scores

The first PC explains 63.72% of the variation of the heptathlon, the loadings of the first PC are:

<table>
<thead>
<tr>
<th>Feature</th>
<th>Loadings</th>
</tr>
</thead>
<tbody>
<tr>
<td>hurdles</td>
<td>0.4529</td>
</tr>
<tr>
<td>highjump</td>
<td>0.3772</td>
</tr>
<tr>
<td>shot</td>
<td>0.3631</td>
</tr>
<tr>
<td>run200m</td>
<td>0.4079</td>
</tr>
<tr>
<td>longjump</td>
<td>0.4562</td>
</tr>
<tr>
<td>javelin</td>
<td>0.0754</td>
</tr>
<tr>
<td>run800m</td>
<td>0.3750</td>
</tr>
</tbody>
</table>

Dimensionality reduction:

- Project all 8 features onto the first PC.
- Compare the scores of the first PC with the official scores used to rank the athletes.
The scores of the first PC $p_{c_1}$ have a similar ranking as the scores of the official scoring system:
Discussion

**Advantage**

- Considers arbitrary correlations between features
- Selected subspace is optimal w.r.t. loss of variance

**Disadvantage**

- Assumption: components with high variance are useful to discover the desired patterns
- Considers only linear correlations (work-around: Kernel-PCA, see later)
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5. Clustering High Dimensional Data
Idea

• PCA is an eigenvalue decomposition of the $d \times d$ covariance matrix $
abla = D^T D$ of the (normalized) data matrix $D$:

$$
\nabla = VEV^T
$$

such that

• $V = (pc_1, ..., pc_d)$, is a $d \times d$ matrix whose columns are the pairwise independent unit vectors, the eigenvectors

$$
\left( \begin{array}{ccc}
\lambda_1 & \cdots & 0 \\
\vdots & \ddots & \vdots \\
0 & \cdots & \lambda_d
\end{array} \right)
$$

• $E = \left( \begin{array}{ccc}
\lambda_1 & \cdots & 0 \\
\vdots & \ddots & \vdots \\
0 & \cdots & \lambda_d
\end{array} \right)$ is a $d \times d$ diagonal matrix, the diagonal elements are the eigenvalues of the corresponding eigenvectors

• The decomposition can be found e.g. based on numerical algorithms
Compute the SVD

- SVD is a generalization of the eigenvalue decomposition
- Let $D$ be the $n \times d$ data matrix ($n$ objects, $d$ dimensions) and let $k$ be its rank (max number of independent rows/columns)
- We can decompose $D$ into matrices $O, S, A$ with $D = OSA^T$ or

$$
\begin{pmatrix}
  x_{1,1} & \cdots & x_{1,d} \\
  \vdots & \ddots & \vdots \\
  x_{n,1} & \cdots & x_{n,d}
\end{pmatrix} =
\begin{pmatrix}
  o_{1,1} & \cdots & o_{1,k} \\
  \vdots & \ddots & \vdots \\
  o_{n,1} & \cdots & o_{n,k}
\end{pmatrix} \cdot
\begin{pmatrix}
  \lambda_1 & \cdots & 0 \\
  \vdots & \ddots & \vdots \\
  0 & \cdots & \lambda_k
\end{pmatrix} \cdot
\begin{pmatrix}
  a_{1,1} & \cdots & a_{1,d} \\
  \vdots & \ddots & \vdots \\
  a_{k,1} & \cdots & a_{k,d}
\end{pmatrix}
$$

such that
- $O$ is a $n \times k$ column-orthonormal matrix (each of its columns is a unit vector and the dot product of any two columns is 0)
- $S$ is a diagonal $k \times k$ matrix
- $A$ is a $k \times d$ column-orthonormal matrix. Note that we always use $A$ in its transposed form, so it is the rows of $A^T$ that are orthonormal
SVD: Example I

- \( D \) contains movie ratings by users
  - The corresponding SVD shows two concepts “science fiction” and “romance”
- \( S \) shows the strength of these concepts
- \( A \) relates movies to concepts

\[
D = \begin{pmatrix}
1 & 1 & 1 & 0 & 0 \\
3 & 3 & 3 & 0 & 0 \\
4 & 4 & 4 & 0 & 0 \\
5 & 5 & 5 & 0 & 0 \\
0 & 0 & 0 & 4 & 4 \\
0 & 0 & 0 & 5 & 5 \\
0 & 0 & 0 & 2 & 2 \\
\end{pmatrix} = \begin{pmatrix}
.14 & 0 \\
.42 & 0 \\
.56 & 0 \\
.70 & 0 \\
0 & .60 \\
0 & .75 \\
0 & .30 \\
\end{pmatrix} \cdot \begin{pmatrix}
12.4 & 0 \\
0 & 9.5 \\
\end{pmatrix} \cdot \begin{pmatrix}
.58 & .58 & .58 & 0 & 0 \\
0 & 0 & 0 & .71 & .71 \\
\end{pmatrix} \cdot \begin{pmatrix}
\end{pmatrix}
\]

SVD: Example II

- Now a slightly different $D$
  - The corresponding SVD shows three concepts “science fiction” and “romance” and ???

\[
\begin{pmatrix}
1 & 1 & 1 & 0 & 0 \\
3 & 3 & 3 & 0 & 0 \\
4 & 4 & 4 & 0 & 0 \\
5 & 5 & 5 & 0 & 0 \\
0 & 2 & 0 & 4 & 4 \\
0 & 0 & 0 & 5 & 5 \\
0 & 1 & 0 & 2 & 2 \\
\end{pmatrix}
\]

\[
\begin{pmatrix}
.13 & .02 & -.01 \\
.41 & .07 & -.03 \\
.55 & -.09 & -.04 \\
.68 & .11 & -.05 \\
.15 & -.59 & .65 \\
.07 & -.73 & -.67 \\
.07 & -.29 & .32 \\
\end{pmatrix}
\]

\[
\begin{pmatrix}
12.4 & 0 & 0 \\
0 & 9.5 & 0 \\
0 & 0 & 1.3 \\
\end{pmatrix}
\]

\[
\begin{pmatrix}
.56 & .59 & .56 & .09 & .09 \\
.12 & -.02 & .12 & -.69 & -.69 \\
.40 & -.80 & .40 & .09 & .09 \\
\end{pmatrix}
\]

(Source: http://infolab.stanford.edu/~ullman/mmds/ch11.pdf)
To reduce dimensionality, we can set the smallest singular values to 0 in $S$ and eliminate the corresponding columns in $O$ and rows in $A^T$ (check previous examples).

- How Many Singular Values Should We Retain?
  - Rule of thumb: retain enough singular values to make up 90% of the energy in $S$.
  - Energy is defined in terms of the singular values (matrix $S$).
  - In the previous example, the total energy is:
    \[(12.4)^2 + (9.5)^2 + (1.3)^2 = 245.70\]
  - The retained energy is: \[(12.4)^2 + (9.5)^2 = 244.01 > 99\%\]
Connection between SVD and PCA

- PCA is applying SVD on the covariance matrix $\Sigma = D^T D$
- SVD means: $D = OSA^T$
- Thus:
  \[
  \Sigma = D^T D = (OSA^T)^T OSA^T = AS^T (O^T O) SA^T
  \]
- Since $O$ is an orthonormal matrix, $O^T O$ is the identity:
  \[
  AS^T (O^T O) SA^T = A (S^T S) A^T
  \]
- $S$ is a diagonal matrix, so transposing has no effect:
  \[
  A (S^T S) A^T = AS^2 A^T = A \begin{pmatrix}
\lambda_1^2 & \ldots & 0 \\
\vdots & \ddots & \vdots \\
0 & \ldots & \lambda_k^2
\end{pmatrix} A^T
  \]
Connection between SVD and PCA

- Here: $A$ is a matrix of eigenvectors
- Eigenvalues of the covariance matrix = squared singular values of $D$
- Conclusion: Eigenvalues and eigenvectors of the covariance matrix $S$ can be determined by the SVD of the data matrix $D$ (or in other words: SVD is a method to perform PCA)
- SVD is sometimes a better way to perform PCA (Large dimensionalities e.g., text data)
- SVD can cope with dependent dimensions ($k < d$ is an ordinary case in SVD)
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5. Clustering High Dimensional Data
Consider the following scenarios:

- PCA will be effective since data is linearly correlated
- PCA may find the orange line as the first component
Basic Idea

Recall: the solution of linear classifiers (e.g. SVMs) for non-linear problems is “make them linear!” using a suitable feature mapping

- No linear separation of classes possible
- Mapping $\mathbb{R}^2 \rightarrow \mathbb{R}^3$ with $(x_1, x_2) \mapsto (x_1, x_2, x_1^2 + x_2^2)$
Kernel Trick

- Since a high-dimensional mapping can still have negative impact, the Kernel trick is used whenever possible (see KDD I lecture).
- Given the intended mapping $\Phi$, the Kernel is usually defined as $K(x, y) = \Phi(x)^T\Phi(y)$.
- Example: Degree-$d$ polynomials: $K(x, y) = (x^T y + c)^d$ with an arbitrary constant $c$, e.g. for $d = 2$:

\[
\Phi : R^2 \rightarrow R^3 \\
(x_1, x_2) \mapsto (z_1, z_2, z_3) := (x_1^2, \sqrt{2}x_1x_2, x_2^2)
\]

(Image source: http://i.stack.imgur.com/qZV3s.png)
Kernel PCA Using SVD

- Recall the SVD $D = OSA^T$
- $A$ is a $k$-dimensional basis of the eigenvectors of $DD^T$ (originally $d \times d$)
- Analogously, $O$ is a $k$-dimensional basis of eigenvectors of $DD^T$
- $DD^T$ is a Kernel matrix for the linear Kernel (i.e., no mapping made - cf. KDD I) or any other Kernel
- $A$ and $O$ are related as follows:

$$D = OSA^T \Rightarrow O^T D = O^T OSA^T = SA^T \Rightarrow S^{-1} O^T D = A^T$$

i.e. each $d$-dimensional eigenvector in $A$ is a linear combination of vectors in $D$ (original or mapped!) and the $n \ k$-dimensional eigenvectors in $O^T$ ($O$ is $n \times k$)
Kernel PCA Using SVD

- Let $K(x, y) = \Phi(x)^T \Phi(y)$ be a kernel for the non-linear transformation $\Phi$
- Assume: $K(x, y)$ is known, but $\Phi(x)$ is not explicitly given
- Let $K$ be the Kernel matrix of $D$ w.r.t. $K(x, y)$, i.e.

$$K = \begin{pmatrix}
K(x_1, x_1) & \ldots & K(x_1, x_n) \\
\vdots & \ddots & \vdots \\
K(x_n, x_1) & \ldots & K(x_n, x_n)
\end{pmatrix}$$

- The eigenvalue decomposition of $K$ is $K = VSV^T$ where $V$ is a $n$-dimensional basis from eigenvectors of $K$
- Dimensionality Reduction through mapping of $y \in D$ w.r.t $V$ to

$$\hat{y} = \begin{pmatrix}
\Phi(y)^T (\sum_{i=1}^{n} v_{i,1} \Phi(x_i)) \\
\vdots \\
\Phi(y)^T (\sum_{i=1}^{n} v_{i,k} \Phi(x_i))
\end{pmatrix} = \begin{pmatrix}
\sum_{i=1}^{n} v_{i,1} (\Phi(y)^T \Phi(x_i)) \\
\vdots \\
\sum_{i=1}^{n} v_{i,k} (\Phi(y)^T \Phi(x_i))
\end{pmatrix} = \begin{pmatrix}
\sum_{i=1}^{n} v_{i,1} K(y, x_i) \\
\vdots \\
\sum_{i=1}^{n} v_{i,k} K(y, x_i)
\end{pmatrix}$$
Matrix Factorization as Optimization Task

• BTW, SVD (and, thus PCA) is a matrix decomposition that can be formalized as optimization task

\[
D = OSA^T = \begin{pmatrix} O \left( \begin{array}{cc} \sqrt{\lambda_1} & \cdots & 0 \\ \vdots & \ddots & \vdots \\ 0 & \cdots & \sqrt{\lambda_k} \end{array} \right) \end{pmatrix} \begin{pmatrix} U \end{pmatrix} \begin{pmatrix} A^T \end{pmatrix} = UV^T
\]

• As an optimization problem: \( L(U, V) = \| D - UV^T \|_f^2 \)

subject to \( \forall i \neq j : \langle v_i, v_j \rangle = 0 \land \langle u_i, u_j \rangle \)

using the squared Frobenius Norm of an \( n \times m \) matrix \( M \):

\[ \| M \|_f^2 = \sum_{i=1}^{n} \sum_{j=1}^{m} |m_{i,j}|^2 \]
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5. Clustering High Dimensional Data
Fisher Faces

• Idea: Use examples from a training set (supervised!) to increase the discriminative power of the target space

• Minimize the similarity between objects from different classes (between class scatter matrix: \( \sigma_b \))
  Use covariance matrix of the class centroids for \( \Sigma_b \)

• Maximize similarity between objects belonging to the same class (within class scatter matrix \( \Sigma_w \))
  Use average covariance matrix of all classes for \( \Sigma_w \)

• Determine new basis vectors \( b_i \) by maximizing
  \[
  \frac{b_i^T \Sigma_b b_i}{b_i^T \Sigma_w b_i}
  \]

subject to \( \forall i \neq j : \langle b_i, b_j \rangle = 0 \)
Fisher Faces

Remarks on Fisher Faces

• The vector having the largest eigenvalue corresponds to the normal vector of the separating hyper plane in linear discriminant analysis or Fisher’s discriminant analysis. (cf. KDD I)
• Fischer Faces are limited due to the assumption of mono-modal classes: each class is assumed to follow one multivariate Gaussian
• Multi-modal or non-Gaussian distributions are not modeled well
• Many variants (e.g. Relevant Component Analysis (RCA), Large Margin Nearest Neighbor (LMNN))
Summary

- Linear basis transformation yield a rich framework to optimize feature spaces
- Unsupervised methods delete low variant dimensions (PCA und SVD)
- Kernel PCA allows to compute PCA in non-linear kernel spaces
- Basic assumption: direction of highest variance bear the most relevant information
- Supervised methods try to minimize the within class distances while maximizing between class distances (Fischer Faces and variants)
Further Readings


Kapitel 5: Clustering High-dim Data

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   5.2 Subspace Clustering
   5.3 Projected Clustering
   5.4 Pattern-based Clustering
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5. Clustering High Dimensional Data

5.1 Challenges

5.2 Subspace Clustering

5.3 Projected Clustering

5.4 Pattern-based Clustering

5.5 Correlation Clustering
Customer Recommendation / Target Marketing

- Data: customer ratings for given products
  - Rows (objects): customers (millions?)
  - Columns (features): products (hundreds to thousands)
  - Value $x_{ij}$ in the data matrix is the rating of product $i$ by user $j$

- Task: Cluster customers to find groups of persons that share similar preferences or disfavor (e.g. to do personalized target marketing)

- Challenge: customers may be grouped differently according to different preferences/disfavors, i.e. different subsets of products
  (See: baby shapes game)
Curse of Dimensionality (Revisited)

Relevant and irrelevant attributes

- Not all features, but a subset of the features may be relevant for clustering.
- Groups of similar (e.g., “dense”) points may be identified when considering only these features.
- Different subsets of attributes may be relevant for different clusters.
Effect on clustering

- Traditional distance functions give equal weight to all dimensions
- However, not all dimensions are of equal importance
- Adding irrelevant dimensions ruins any clustering based on a distance function that equally weights all dimensions
Curse of Dimensionality (Revisited)

- Example: different attributes are relevant for different clusters
It can even be a little bit more complex ...

- **Task:** Cluster test persons to find groups of individuals with similar correlation among the concentrations of metabolites indicating homogeneous metabolic behavior (e.g. disorder)

- **Challenge:** different metabolic disorders appear through different correlations of (subsets of) metabolites
Correlation among attributes

- A subset of features may be correlated.
- Groups of similar (e.g. “dense”) points may be identified when considering this correlation of features only.
- Different correlations of attributes may be relevant for different clusters.
Why not feature selection/reduction

• (Unsupervised) feature selection or feature reduction (e.g. PCA) is global, i.e., it transforms the original feature space into one new representation

• We face a local feature relevance/correlation: some features (or combinations of them) may be relevant for one cluster, but may be irrelevant for a second one, i.e., we need multiple representations
Example: use PCA (target dim = 1) before clustering
Example: cluster first, then find correlations (with PCA)
Problem Summary

• Feature relevance and correlation
  • Usually, no clusters in the full dimensional space
  • Often, clusters are hidden in subspaces of the data, i.e. only a subset of features is relevant for the clustering
  • E.g. a group of genes play a common role in a subset of experimental conditions but may behave completely different in other conditions

• Local feature relevance/correlation
  • For each cluster, a different subset of features or a different correlation of features may be relevant
  • E.g. different genes are responsible for different phenotypes other conditions

• Overlapping clusters (different semantic concepts)
  • Clusters may overlap, i.e. an object may be clustered differently in varying subspaces
  • E.g. a gene plays different functional roles depending on the environment other conditions
General Problem Setting

Search for clusters in (in general arbitrarily oriented) subspaces of the original feature space

Challenges

- Find the correct subspace of each cluster (Search space virtually infinite: all possible arbitrarily oriented subspaces of a feature space)
- Find the correct cluster in each relevant subspace (Search space depends on clustering algorithm)

Even worse: both challenges depend on each other:
- In order to determine the correct subspace of a cluster, we need to know (at least some) cluster members
- In order to determine the correct cluster memberships, we need to know the subspaces of all clusters
• Hmm, it is really not so easy, especially in practice ...

• Even if we found the relevant subspace, we still might have a hard time to find the correct cluster(s)

• Here, clusters (yellow, red, green, blue) and noise (gray) are not separable in the “correct” subspace . . .

• Rather, we would need a new cluster model (recall: cluster = group of similar objects)

• What is the concept of similarity that all members of a cluster share in this example (and does not include members from other clusters/noise)???
### A Taxonomy of Approaches

<table>
<thead>
<tr>
<th>Subspace Clustering (restricted to axis-parallel subspace)</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Find all clusters in all subspaces (allow overlaps)</td>
</tr>
<tr>
<td>• Usually bottom-up subspace search</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Projected Clustering (restricted to axis-parallel subspace)</th>
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</thead>
<tbody>
<tr>
<td>• Each point is assigned to one subspace cluster or noise</td>
</tr>
<tr>
<td>• Usually top-down subspace search</td>
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</tbody>
</table>

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<thead>
<tr>
<th>Correlation Clustering (explores arbitrarily oriented subspaces)</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Each point is assigned to one subspace cluster or noise</td>
</tr>
<tr>
<td>• Bottom-up and top-down subspace search approaches</td>
</tr>
</tbody>
</table>
1. Introduction to Feature Spaces

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Bottom-up Subspace Cluster Search

- Similar to Branch-and-Bound feature selection: Start with 1-D subspaces or subspace clusters and merge them to compute higher dimensional ones
- Most approaches transfer this problem into a frequent item set mining problem
- In this case, the cluster criterion must implement the downward closure (monotonicity) property:
  - If the criterion holds for a $k$-dimensional subspace $S$, then it also holds for any $(k-1)$-dimensional projection of $S$
  - Use the reverse implication for pruning: If the criterion does not hold for a $(k-1)$-dimensional projection of $S$, then the criterion also does not hold for $S$
- Some approaches use other search heuristics (especially if monotonicity does not hold) like best-first-search, greedy-search, ...
Downward-closure Property: Example

- Consider a simple cluster criterion (density of grid cells): If a cell $C$ of side length $s$ contains more than $m$ points, it represents a cluster.
- Monotonicity: if $C$ contains more than $m$ points in subspace $S$ then $C$ also contains more than $m$ points in any subspace $T \subseteq S$.

Cell $C$ contains more than $m=5$ points in subspace "$AB$"
$\Rightarrow$ Also in subspaces "$A \subseteq AB$" and "$B \subseteq AB$"

Cell $C$ contains less than $m=5$ points in subspace "$A$"
$\Rightarrow$ Also in subspace "$AB$"
CLIQUE: Idea

- Probably the first bottom-up algorithm
- It uses a density-grid-based cluster model (similar to previous slide)

- Clusters are “dense regions” in the feature space
- Partition the feature space into $\xi$ equal sized parts in each dimension
- A unit is the intersection of one interval from each dimension
- Unit $u$ is dense if it contains more than $\tau$ objects

- Clusters are maximal sets of connected dense units (e.g., $A \cup B$)
- Two-step approach (1. subspace search, 2. clustering)
CLIQUE: Subspace Search

Step 1: Find subspaces with dense units

- Explore Downward Closure property of dense cells (APRIORI-style search)

- Candidate generation
  - Based on $D_{k-1}$, the set of $(k-1)$-dimensional dense units, generate candidate set $C_k$ by self joining $D_{k-1}$
  - Join condition: units share first $k-2$ dimensions
  - Discard those candidates which have a $k-1$ projection not included in $D_{k-1}$
  - For the remaining candidates: check density
CLIQUE: Cluster Search

Step 2: Find clusters as maximal sets of connected dense units

- Given: a set of dense units $D$ in the same $k$-dimensional subspace $S$
- Output: A partition of $D$ into clusters $D_1, \ldots, D_k$ of connected dense units
- The problem is equivalent to finding connected components in a graph
  - Nodes: dense units
  - Edge between two nodes if the corresponding dense units have a common face (neighboring units)
  - Depth-first search algorithm: Start with a unit $u$ in $D$, assign it to a new cluster ID and find all the units it is connected to
  - Repeat if there are nodes not yet visited
CLIQUE: Discussion

- Input parameters: $\xi$ and $\tau$ specifying the density threshold
- Output: all clusters in all subspaces, clusters may overlap
- Simple but efficient cluster model
- Uses a fixed density threshold for all subspaces (in order to ensure the downward closure property)
  - To represent a cluster, a unit in 10D must contain as many points (or more) as in 2D . . .
  - For a cluster $C$ in subspace $S$, all clusters in all projections of $S$ are also reported as clusters (extremely high redundancy)
  - Which of the redundant information is more interesting? ($D$ in $S$ or $D'$ in $T \subset S$; $D'$ may contain more units/points)
- Worst case runtime?
CLIQUE: Variants

There are different variations of CLIQUE varying mainly in terms of ...

- ... different density definition, e.g. using Entropy for subspaces rather than simple counts of units
- ... different grid construction methods, e.g. adaptive intervals (but then, no downward closure property is given anymore)

Drawbacks of a grid-based clustering model:

- Positioning of the grid influences the clustering
- Selection of $\xi$ and $\tau$ is very sensitive
- Example: Either $C_2$ and $C_1$ are found as clusters or none of them
Use DBSCAN model (maximal density-connected sets): density is defined w.r.t. the location of points not w.r.t. the data space:

**Core Points**
- Points finding more than $MinPts$ other points in its $\varepsilon$-neighborhood

**Density connectivity**
- Core points may have core points in their $\varepsilon$-neighborhood
- Build transitive chains of such core points to find connected sets
\textbf{SUBCLU: Basics}

- Richer cluster model: detects clusters of arbitrary shapes and locations (in the corresponding subspaces)
- Naive approach: Apply DBSCAN in all possible subspaces (exponential runtime!)
- Idea: Exploit clustering information from previous step (subspaces)
- Density-connected clusters are not monotonic, but density connected sets are (see next slide)
  - If \( C \) is a density connected set in subspace \( S \) then \( C \) is a density connected set in any subspace \( T \subset S \)
  - But, if \( C \) is a cluster in \( S \), it need not to be a cluster in \( T \subset S \) because maximality might be violated, i.e., in \( T \) there may be additional points density connected to the points in \( C \)
  - A cluster in a higher-dimensional subspace \( S \) will be a subset of a cluster in the a projection \( T \)
- Thus, APRIORI-style subspace search is possible
Example (circles indicate $\varepsilon$-neighborhood):

$p$ and $q$ density connected in $\{A, B\}$, thus, they are also density connected in $\{A\}$ and $\{B\}$

$p$ and $q$ not density connected in $\{B\}$, thus, they are not density connected in $\{A, B\}$, although they are density connected in $\{A\}$
• Algorithm
  • All subspaces that contain any density-connected set are computed using the bottom-up approach (similar to CLIQUE/APRIORI)
  • Density-connected clusters are computed using a specialized DBSCAN run in the resulting subspace to generate the subspace clusters

• Discussion
  • Input: $\epsilon$ and $MinPts$ specifying the density threshold
  • Output: all clusters in all subspaces, clusters may overlap
  • Uses a fixed density threshold for all subspaces
  • Advanced but costly cluster model
Bottom-up Variants: FIRES

Uses a different bottom-up heuristic for subspace search:

- Starts with 1-dimensional clusters called base clusters (generated by applying any traditional clustering algorithm to each 1-dimensional subspace)
- Merges these clusters to generate subspace cluster approximations by applying a clustering of the base clusters using a variant of DBSCAN (similarity between two clusters $C_1$ and $C_2$ is defined by $|C_1 \cap C_2|$) in order to “jump” to maximal dimensional subspaces
- Refines the resulting cluster approximations using any traditional clustering algorithm on the points within the approximations
Similar idea to FIRES:

- Cluster cores are hyper-rectangular approximations of subspace clusters
- Subspace search is APRIORI-style: cluster cores are computed bottom-up from “significant” 1D intervals
- Significant 1D intervals are determined using a hypothesis test:
  - Hypothesis: no clusters, i.e. points are randomly distributed
  - If an interval contains significantly more points than expected, the hypothesis is rejected (significant 1D interval)
- Significant 1D intervals follow the downward closure property
- Postprocessing: Cluster cores initialize an EM fuzzy clustering of all data points
Find subspace cluster hierarchies (lower dimensional clusters embedded in higher dimensional ones):

- Integrate a proper distance function into hierarchical clustering
- Learns distance function instance-based bottom-up
Several different variants exist; most of them suffer from a key limitation: global density thresholds

- In order to ensure the downward closure property, the density threshold must be fixed globally
- Consequence: the points in an e.g. 20-dimensional subspace cluster must be as dense as in an e.g. 2-dimensional cluster
- This is a rather optimistic assumption since the data space grows exponentially with increasing dimensionality (see “curse” discussion)
- Consequences:
  - A strict threshold will most likely produce only lower dimensional clusters
  - A loose threshold will most likely produce higher dimensional clusters but also a huge amount of (potentially meaningless) low dimensional clusters
Kapitel 5: Clustering High-dim Data

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Top-down Subspace Search: Approaches

Several different variants for top-down subspace search:

• Cluster-based search
  • Learn the subspace of a cluster in the entire $d$-dimensional feature space
  • Start with full-dimensional clusters
  • Iteratively refine the cluster memberships of points and the subspaces of the cluster

• Instance-based search
  • Learn for each point its subspace preference in the entire $d$-dimensional feature space
  • The subspace preference specifies the subspace in which each point “clusters best”
  • Merge points having similar subspace preferences to generate the clusters

• Random search: learn the subspace preference of a cluster or a point from randomly sampled points
How should we learn the subspace preference of a cluster or a point?

- Most approaches rely on the so-called “locality assumption”: the subspace preference can be learned from the local neighborhood in the $d$-dimensional space.
- The subspace is usually learned from the local neighborhood of cluster representatives/cluster members in the entire feature space:
  - Cluster-based approach: the local neighborhood of each cluster representative is evaluated in the $d$-dimensional space to learn the “correct” subspace of the cluster.
  - Instance-based approach: the local neighborhood of each point is evaluated in the $d$-dimensional space to learn the “correct” subspace preference of this point (i.e. the subspace in which the cluster exists that accommodates this point).
- Random search approaches do not suffer from the locality assumption but usually need many sampling rounds.
• Cluster-based top-down approach: we learn the subspace for each cluster

• \(K\)-medoid cluster model
  • Cluster is represented by its medoid
  • To each cluster a subspace (of relevant attributes) is assigned
  • Each point is assigned to the nearest medoid (where the distance to each medoid is based on the corresponding subspace of the medoid)
  • Points that have a large distance to their nearest medoid(s) are classified as noise
PROCLUS

3-phase algorithm (input: number of clusters $k$, average dimensionality of subspaces $L$):

**Phase 1**: Initialization of cluster medoids

- Ideally we want a set of centroids, where each centroid comes from a different cluster
- We do not know which are these $k$ points though, so we choose a superset $M$ of $b \cdot k$ medoids such that they are well separated
  - Chose a random sample ($S$) of $a \cdot k$ data points
  - Out of $S$, select $b \cdot k$ points ($M$) by greedy selection: medoids are picked iteratively so that the current medoid is well separated from the medoids that have been chosen so far
- Additional input parameters $a$ and $b$ are introduced for performance reasons
- This procedure has nothing to do with subspace/projected clustering but could be applied for any $k$-partitioning algorithm
Phase 2: Iterative phase (works similar to any $k$-medoid clustering)

- $k$ randomly chosen medoids from $M$ (Phase 1) are the initial cluster medoids
- Replace the “bad” medoids with other points in $M$ if the quality of the clustering increases
- Procedure:
  - Find dimensions related to the medoids
  - Assign data points to the medoids
  - Evaluate the clusters formed
  - Find the bad medoids, and try to improve the result by replacing these bad medoids
PROCLUS: Find Cluster Dimensions

• For each medoid \( m_i \), let \( 2 \cdot \delta \) be the distance to its closest medoid

• All the data points within \( \delta \) will be surely assigned to the medoid \( m_i \) (this set \( L_i \) is the locality of \( m_i \))

• Intuition: to each medoid we want to associate those dimensions where the points are closed to the medoid in that dimension

• Compute the average distance \( X_{i,j} \) along each dimension \( j \) from the points in \( L_i \) to \( m_i \)

• Calculate for \( m_i \) the mean \( Y_{i,j} \) and standard deviation \( \sigma_{i,j} \) of \( X_{i,j} \)

• Calculate \( Z_{i,j} = (X_{i,j} - Y_{i,j}) / \sigma_{i,j} \)

• Choose \( k \cdot l \) smallest values \( Z_{i,j} \) with at least 2 chosen for each medoids (to ensure that cluster preferences are lower dimensional)

• Output: A set of \( k \) medoids and their associated dimensions
PROCLUS: Point Assignment and Cluster Evaluation

- Assign each data point to its closest medoid using Manhattan segmental distance (only relevant dimensions count)

Manhattan segmental distance: For any two points \(x_1, x_2\) and any set of dimensions \(D\):

\[
MSdist_D(x_1, x_2) = \frac{\sum_{i \in D} |x_{1i} - x_{2i}|}{|D|}
\]

- Evaluate a cluster \(C_i\) using average \(MSdist\) from the points in \(C_i\) to the centroid of \(C_i\) along dimension \(j\)

- Replace bad medoids with random points from \(M\)

- Terminate if the clustering quality does not increase after a given number of current medoids have been exchanged with medoids from \(M\) (it is not clear, if there is another hidden parameter in that criterion)
Phase 3: Refinement

• Reassign subspaces to medoids as above (but use only the points assigned to each cluster rather than the locality of each cluster, i.e., $C_i$ not $L_i$)

• Reassign points to medoids

• Points that are not in the locality of any medoid are classified as noise
• Use DBSCAN cluster model instead of \( k \)-medoid clustering

• Use DBSCAN with a distance that adopts to the subspace preferences of points

• Given the input parameters \( \delta \) and \( \kappa >> 1 \), for each point \( p \), the \( d \)-dimensional subspace preference vector \( w^p \) of \( p \) is defined using the following components:

\[
    w^p_i = \begin{cases} 
        1 & \text{if } \VAR_i > \delta \\
        \kappa & \text{if } \VAR_i \leq \delta 
    \end{cases}
\]

where

\[
    \VAR_i = \frac{\sum_{q \in N_\varepsilon(p)} \dist(p_i, q) \dist(p_i, q)}{|N_\varepsilon(p)|}
\]
• Preference weighted distance function

\[ dist_p(p, q) = \sqrt{\sum_{i=1}^{d} w_i^p \cdot (p_i - q_i)^2} \]

can be made symmetric:

\[ dist_{pref}(p, q) = \max\{dist_p(p, q), dist_q(q, p)\} \]

• Preference \( \varepsilon \)-neighborhood of a point uses \( dist_{pref} \)
• Preference weighted core points, direct density reachability, reachability and connectivity are defined based on preference neighborhood
• A subspace preference cluster is a maximal density connected set of points associated with a certain subspace preference vector
• Clusters may have arbitrary shape in the subspace projection
Variants

• **DOC**
  • Clusters are “dense” hyper-rectangles
  • Computes at most one cluster in each run
  • Random sampling for subspace learning

• **COSA**
  • Learns a distance matrix that can be used for clustering
  • Instance-based locality assumption
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• Challenges and Approaches, Basic Models
  • Constant Biclusters
  • Biclusters with Constant Values in Rows or Columns
  • Pattern-based Clustering: Biclusters with Coherent Values
  • Biclusters with Coherent Evolutions

• Algorithms for
  • Constant Biclusters Pattern-based Clustering: Biclusters with Coherent Values
General Idea

- Pattern-based clustering relies on patterns in the data matrix.
- Simultaneous clustering of rows and columns of the data matrix (hence “bi-clustering”).
- Data matrix $A = (X, Y)$ with set of rows (objects) $X$ and set of columns (features) $Y$.
- $a_{xy}$ is the element in row $x$ and column $y$.
- Submatrix $A_{IJ} = (I, J)$ with subset of rows $I \subseteq X$ and subset of columns $J \subseteq Y$ contains those elements $a_{ij}$ with $i \in I$ and $j \in J$. 
General Idea

General aim of biclustering approaches:

- Find a set of submatrices \( \{(I_1, J_1), (I_2, J_2), \ldots, (I_k, J_k)\} \) of the data matrix \( A = (X, Y) \) such that each submatrix (= bicluster) meets a given homogeneity criterion.

Some values often used by bicluster models:

- Mean of row \( i \): \( a_{i,J} = \frac{1}{|J|} \sum_{j \in J} a_{ij} \)
- Mean of column \( j \): \( a_{I,j} = \frac{1}{|I|} \sum_{i \in I} a_{ij} \)
- Mean of all elements:

\[
\begin{align*}
    a_{I,J} &= \frac{1}{|I||J|} \sum_{i \in I, j \in J} a_{ij} = \frac{1}{|J|} \sum_{j \in J} a_{I,j} = \frac{1}{|I|} \sum_{i \in I} a_{iJ}
\end{align*}
\]
General Idea

Different types (models) of biclusters

- constant biclusters
- biclusters with constant values on columns and/or constant values on rows
- biclusters with coherent values (aka. pattern-based clustering)
- biclusters with coherent evolutions
Cluster Model

- All points share identical value in selected attributes
- The constant value $\mu$ is a typical value for the cluster
- Thus, the cluster model is $a_{ij} = \mu$
- Obviously a special case of an axis-parallel subspace cluster
- Example:
Constant Biclusters

• Interpretation: points are located on the bisecting line of participating attributes

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<thead>
<tr>
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• Visualization using Parallel Coordinates: identical constant lines

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</table>
Cluster Model

- Cluster model for $A_{IJ} = (I, J)$ with $a_{ij} = \mu + c_j$
- $c_j$ is an adjustment value for column $j$
- Results in axis-parallel subspace clusters
- Example:

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Biclusters with Constant Column Values

• Interpretation: points are located on the same values (not necessary bisecting line)

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</table>
Biclusters with Constant Row Values

Cluster Model

- Cluster model for $A_{IJ} = (I, J)$ with $a_{ij} = \mu + r_i$
- $r_i$ is an adjustment value for column $i$
- Results in points building a sparse hyperplane parallel to irrelevant axes (axis-parallel subspace cluster)
- Example:

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<tr>
<td>P4</td>
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Biclusters with Constant Row Values

- Interpretation: points are accommodated on the bisecting line of participating attributes

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- Visualization using Parallel Coordinates: parallel constant lines

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</table>
Biclusters with Coherent Values

Cluster Model

- Based on a particular form of covariance between rows and columns: $a_{ij} = \mu + r_i + c_j$
- Special cases: constant rows ($c_j = 0$) and constant columns ($r_i = 0$)
- Results in points building a hyperplane parallel to axes of irrelevant attributes (axis-parallel subspace cluster)
- Example:

```
+-----+-----+-----+
|     | a1  | a2  | a3  |
|-----+-----+-----+-----|
| P1  | 1   | 2   | 3.5 |
| P2  | 2   | 3   | 2.3 |
| P3  | 4   | 5   | 0.2 |
| P4  | 5   | 6   | 0.7 |
+-----+-----+-----+-----+
```
Biclusters with Coherent Values

- Interpretation: increasing one-dimensional line in relevant subspace

- Visualization using Parallel Coordinates: parallel lines

\[\begin{array}{ccc}
\text{P1} & 1 & 2 \\
\text{P2} & 2 & 3 \\
\text{P3} & 4 & 5 \\
\text{P4} & 5 & 6 \\
\end{array}\]

\[\begin{array}{ccc}
\text{P1} & 1 & 2 & 3.5 \\
\text{P2} & 2 & 3 & 2.3 \\
\text{P3} & 4 & 5 & 0.2 \\
\text{P4} & 5 & 6 & 0.7 \\
\end{array}\]
Biclusters with Coherent Evolution

Cluster Model

- For all rows, all pairs of attributes change simultaneously, e.g.
  - Discretized attribute space: coherent state-transitions: similar to grid-based axis parallel approaches (see above)
  - Change in same direction irrespective of the quantity

- Example with coherent state-transitions:
Biclusters with Coherent Evolution

- Interpretation: points are in certain (grid-cell-like) half spaces
- Visualization using Parallel Coordinates: all lines cross border between states (in the same direction)
Biclusters with Coherent Evolution

• Change in same direction — general idea: find a subset of rows and columns, where a permutation of the set of columns exists such that the values in every row are increasing
• Clusters do not form a subspace but rather half-spaces
• Example with change in same direction:

<table>
<thead>
<tr>
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<th>a1</th>
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<td>P4</td>
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Biclusters with Coherent Evolution

• Interpretation: points are in certain (non axis-parallel) half spaces

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• Visualization using Parallel Coordinates: all lines increasing

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Bicluster Models: Big Picture

Matrix-Pattern
- no change of values
- change of values only on columns or only on rows
- change of values by same quantity (shifted pattern)

Bicluster Model
- Constant Bicluster
- Constant Columns
- Constant Rows
- Coherent Values
- Coherent Evolutions

Spatial Pattern
- axis-parallel, located on bisecting line
- axis-parallel
- axis-parallel sparse hyperplane – projected space: bisecting line
- axis-parallel sparse hyperplane – projected space: increasing line (positive correlation)

- state-transitions:
- grid-based axis-parallel
- change in same direction:
- half-spaces (no classical cluster-pattern)
Algorithms for Constant Biclusters

- Classical problem statement by Hartigan in 1972
- Quality measure for a bicluster: variance of the submatrix $A_{I J}$:
  \[
  \text{VAR}(A_{I J}) = \sum_{i \in I, j \in J} (a_{ij} - a_{I,J})^2
  \]
- Recursive split of data matrix into two partitions
- Each split chooses the maximal reduction in the overall sum of squares for all biclusters
- Avoids partitioning into $|X||Y|$ singularity-biclusters (optimizing the sum of squares) by comparing the reduction with the reduction expected by chance
Biclustering Algorithms

Algorithms for Biclusters with Constant Values in Rows or Columns

• Simple approach: normalization to transform the biclusters into constant biclusters and follow the first approach
• Some application-driven approaches with special assumptions in the bioinformatics community
• Constant values on columns: general axis-parallel subspace/projected clustering
• Constant values on rows: special case of general correlation clustering
• Both cases special case of approaches to biclusters with coherent values
Biclustering Algorithms

Pattern-based Clustering: Algorithms for Biclusters with Coherent Values

- Classical approach by Cheng and Church 2000
- Introduced the term biclustering to analysis of gene expression data
- Quality of a bicluster: mean squared residue value $H$ defined as follows
  \[
  H(I, J) = \frac{1}{|I||J|} \sum_{i \in I, j \in J} (a_{ij} - a_{i,J} - a_{I,j} + a_{I,J})^2
  \]
  - Submatrix $(I, J)$ is considered a bicluster, if $H(I, J) < \delta$
  - If $\delta = 0$, the biclusters are perfect
• The model for a perfect bicluster predicts value $a_{ij}$ by a row-constant, a column-constant, and an overall cluster-constant:

$$a_{ij} = a_{i,J} + a_{l,j} - a_{l,J} = \mu + r_i + c_j$$

with $\mu = a_{l,J}$, $r_i = a_{i,J} - a_{l,J}$, and $c_j = a_{l,j} - a_{l,J}$, i.e., each row and column exhibits absolutely consistent bias

• For a non-perfect bicluster, the prediction of the model deviates from the true value by a residue

$$a_{ij} = \text{res}(a_{ij}) + a_{i,J} + a_{l,j} - a_{l,J}$$

with $\text{res}(a_{ij}) = a_{ij} - a_{i,J} - a_{l,j} + a_{l,J}$ which is the optimization criterion $H(I, J)$ above
Related Algorithms

- $p$-cluster algorithm specializes the $\delta$-bicluster-property to a pairwise property of two objects in two attributes
- FLOC uses a randomized procedure
- MaPle introduces improved pruning
- CoClus follows a $k$-means-like approach
Biclustering: Summary

Related Algorithms

- Biclustering models do not fit exactly into the spatial intuition behind subspace, projected, or correlation clustering
- Models rather make sense in view of a data matrix
- Strong point: the models generally do not rely on the locality assumption
- Models differ substantially, so a careful selection is a non-trivial task
Kapitel 5: Clustering High-dim Data

1. Introduction to Feature Spaces

2. Challenges of High Dimensional Data

3. Supervised Feature Selection

4. Feature Reduction and Metric Learning

5. Clustering High Dimensional Data

5.1 Challenges

5.2 Subspace Clustering

5.3 Projected Clustering

5.4 Pattern-based Clustering

5.5 Correlation Clustering
• As we have seen before, there may be more than simple (independent) feature relevancy: feature correlation

• Example:
  • Cluster 3 exists in an axis-parallel subspace (so far so good)
  • However, clusters 1 and 2 exist in (different) arbitrarily oriented subspaces that represent different correlations among attributes

• Can the methods introduced so far capture those patterns?
Motivation

• Subspace/projected clustering methods
  • Obviously, these methods are restricted to axis-parallel clusters, i.e., assume feature independence

• Pattern-based methods
  • Can find simple pairwise positive correlations (coherent evolution)
  • However, negative correlations have no additive pattern:
Motivation

• Pattern-based methods (cont.)
  • Also, more complex relationships between several features are out of scope of pattern-based approaches
  • Example:

    Complex correlation among three attributes: \( a1 - 2 \cdot a2 + a3 = 0 \)

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</tr>
<tr>
<td>P5</td>
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</table>

Data matrix

Parallel coordinates

Data space
Correlation Clustering: General Idea

• To find clusters in subspaces of correlated attributes, a more general approach is necessary

• This more general approach is called oriented clustering aka. generalized subspace/projected clustering aka. correlation clustering in the literature \(^1\)

• Assumption: any cluster is located in an arbitrarily oriented affine subspace \(S + a\) of \(\mathbb{R}^d\)

---

Correlation Clustering: General Idea

- Affine subspace $S + a$ of $\mathbb{R}^d$ is interesting if a set of points clusters within this subspace.
- Points may exhibit high variance in perpendicular subspace $(\mathbb{R}^d S) + a$
  $\Rightarrow$ points form a hyperplane located in this subspace $(\mathbb{R}^d S) + a$.
Correlation Clustering: General Idea

• So in general: points on a hyperplane appear to follow linear dependencies among the attributes participating in the description of the hyperplane

• Another example with a 1D subspace / 2D hyperplane

• How can we capture that???
PCA Revisited

- Assume, we know the cluster members, how can we capture the relevant subspace?
- In order to find the directions of high/low variance of a set of points $D$, we could use PCA
- So we could compute the covariance matrix $\Sigma_D$ of $D$, i.e.,

$$\Sigma_D = \frac{1}{|D|} \sum_{x \in D} (x - \mu_D)(x - \mu_D)^T,$$

where $\mu_D$ is the mean (or centroid) of the points in $D$, and compute the eigen decomposition.
PCA Revisited

- The decomposition $\Sigma_D = V_D E_D V_D^T$ gives us
  - The $d$ eigen values in the diagonal matrix $E_D$ representing the variance along the direction of highest variance
  - The $d \times d$-dimensional eigenvectors in the matrix $V_D$ indicating the directions of highest variance sorted by decreasing eigen values

- $E_D$: $d \times d$ diagonal matrix with eigen values
- $V_D$: $d \times d$ orthonormal matrix with eigen vectors
PCA Computes the Correlation

• If points in $D$ forming $\lambda$-dimensional hyperplane, this hyperplane is spanned by the first $\lambda$ eigenvectors of $V_D$ (called “strong” eigenvectors), denoted by $\tilde{V}_D$

• The subspace where the points cluster is spanned by the remaining $d - \lambda$ eigenvectors (called “weak” eigenvectors), denoted by $\hat{V}_D$

• For the eigensystem, the sum of the smallest $d - \lambda$ eigenvalues is minimal under all possible transformations

• Thus, points in $D$ cluster optimally dense in this subspace

• Note again: the subspace where the points cluster is spanned by the weak eigenvectors $\hat{V}_D$
General Model for Correlation Clusters

• As a general model for correlation clusters, we consider a \(\lambda\)-dimensional hyperplane accommodating the points of a correlation cluster \(C\) defined by an equation system of \(d - \lambda\) equations for \(d\) variables and the affinity (e.g. the mean point \(\mu_C\) of all cluster members):

\[
\hat{V}_C^T x = \hat{V}_C^T \mu_C
\]

• The equation system approximately fulfilled for all points \(x \in C\)
• This is quantitative model for the cluster allowing for probabilistic prediction (classification)
• Note: correlations are observable, linear dependencies are merely an assumption to explain the observations — predictive model allows for evaluation of assumptions and experimental refinements

---

PCA-based Correlation Cluster Approaches

- Many algorithms include PCA for feature evaluation into the clustering process similar to (axis-parallel) variance analysis.
- But how do we know \( C \), the set of points we should apply PCA on in order to get the correct subspace?
- Similar to the axis-parallel versions, we need some assumptions (very often a locality assumption, see above).
- Examples:
  - ORCLUS\(^3\): integrates distance function into \( k \)-means, relies on a cluster-based locality assumption.
  - 4C\(^4\): integrates distance function into DBSCAN, relies on instance-based locality assumption.
  - Extensions to 4C: COPAC and ERiC.

---

\(^3\)Aggarwal, Yu: Finding generalized projected clusters in high dimensional space. In Proc. SIGMOD, Dallas, TX, 2000

ORCLUS: Basics

- A generalized projected cluster is a set of orthonormal vectors $E$ and a set of points $C$ such that the points in $C$ are closely clustered in the subspace defined by the vectors $E$ (where $|E| \leq d$)
- Input:
  - The number of clusters $k$
  - The average dimensionality of the subspace of the clusters, $l$
- Output:
  - A set of $k$ clusters and their associated subspaces of dimensionality $l$
- Main idea
  - To find the subspace of a cluster $C_i$, compute the covariance matrix $M_i$ for $C_i$ and determine the eigenvectors
  - Pick “approx.” $l$ eigenvectors with the smallest eigenvalues
ORCLUS: Overview of the Algorithm

- Very similar to PROCLUS but use PCA on locality of cluster representatives instead of variance analysis
- $k$-means like approach (uses centroids rather than medoids)
- Start with $k_c > k$ seeds
- Assign points to clusters according to distance function based on the eigensystem of the current cluster (starting with axes of data space, i.e. Euclidean distance)
- The eigensystem is iteratively adapted based on the updated cluster members (i.e., not only optimize the centroids but also the eigensystems of each cluster)
- Reduce the number of clusters $k_c$ in each iteration by merging best-fitting cluster pairs
ORCLUS: Merging Clusters

- Find best fitting pair of clusters: least average distance in the projected space spanned by weak eigenvectors of the merged clusters

- Two clusters \( C_i \) and \( C_j \) exist in possibly different subspaces

- Compute the subspace of their union \( C_i \cup C_j \) (take the smallest \( l \) eigenvalues)

- Check mean square error (MSE) of the points from the new centroid in this new subspace

- Assess average distance in all merged pairs of clusters and finally merge the best fitting pair (smallest MSE)
• Density-based cluster-paradigm, see PREDECON

• Extend a cluster from a seed as long as a density-criterion is fulfilled — otherwise pick another seed unless all data objects are assigned to a cluster or noise

• Density criterion: minimal required number of points in the neighborhood of a point

• Neighborhood: distance between two points ascertained based on the eigensystems of both compared points
4C: Distance Adaption

- Perform PCA on the local neighborhood $S$ of $p$ to find subspace correlations
- A parameter $\delta$ discerns large from small eigenvalues
- $CorDim(S) = \#\text{eigenvalues} > \delta$
- In the eigenvalue matrix $E_p$ of $p$, large eigenvalues are replaced by 1, small eigenvalues by a value $\kappa >> 1$
- The adopted eigenvalue matrix of $p$ is denoted by $E'_p$
4C: Distance Adaption

- Effect on distance measure:

  distance of \( p \) and \( q \) w.r.t. \( p \):
  \[
  \sqrt{(p - q) \cdot V_p \cdot E'_p \cdot V'_p \cdot (p - q)^T}
  \]
  relies on \( E'_p \) and \( V_p \)

  distance of \( p \) and \( q \) w.r.t. \( q \):
  \[
  \sqrt{(q - p) \cdot V_q \cdot E'_q \cdot V'_q \cdot (q - p)^T}
  \]
  relies on \( E'_q \) and \( V_q \)

- Distance is not symmetric:
4C: Distance Adaption

- Since DBSCAN needs a symmetric distance measure, use the maximum
- Then, \( p \) and \( q \) are “correlation-\( \varepsilon \)-neighbors” if

\[
\max \left\{ \sqrt{(p - q) \cdot V_p \cdot E'_p \cdot V_p^T \cdot (p - q)^T}, \sqrt{(q - p) \cdot V_q \cdot E'_q \cdot V_q^T \cdot (q - p)^T} \right\} \leq \varepsilon
\]

\( p \) and \( q \) are no \( \varepsilon \)-neighbors

\( p \) and \( q \) are \( \varepsilon \)-neighbors

(Circles indicate the \( \varepsilon \) threshold)
4C: Summary

- Finds arbitrary number of clusters
- Requires specification of density-thresholds
  - $\mu$ (minimum number of points): rather intuitive
  - $\varepsilon$ (radius of neighborhood): hard to guess
- Additional parameters:
  - Maximal dimensionality $\lambda$ of correlation clusters (required, otherwise, 4C can also find full-dimensional clusters like DBSCAN)
  - $\delta$ for distinction between strong and weak eigen vectors
- These two parameters are erased by the extensions such as COPAC: take the minimum number of strong eigen vectors that explain a given portion of the overall variance (see PCA)
Correlation Clustering with PCA

- PCA is a mature technique, allows construction of a broad range of similarity measures for local correlation of attributes.
- Can be replaced by similar techniques like relevant component analysis (RCA), independent component analysis (ICA), etc.
- Key limitation still: all approaches suffer from the locality assumption because PCA only works, if the set of points on which PCA is performed is a fairly good approximation of a subspace/correlation cluster.

We need a new principle to employ correlation clustering (coming up)
This new principle is the Hough transform (originally developed in image analysis)

- Transform each object into a so-called parameter space representing all possible subspaces accommodating this object (i.e. all hyperplanes through this object)
- This parameter space is a continuum of all these subspaces
- However, the subspaces are represented by a considerably small number of parameters
- This transform is a generalization of the Hough Transform (which is designed to detect linear structures in 2D images) for arbitrary dimensions
The Hough Transform in 2D

Check out the board ...
The General Hough Transform

- For each $d$-dimensional point $p$ there is an infinite number of
  $(d-1)$-dimensional hyper-planes through $p$
- Each of these hyper-planes $s$ is defined by $(p, \alpha_1, \ldots, \alpha_{d-1})$, where
  $\alpha_1, \ldots, \alpha_{d-1}$ is the normal vector $n_s$ of the hyper-plane $s$
- The function $f_p(\alpha_1, \ldots, \alpha_{d-1}) = \delta_s = \langle p, n_s \rangle$ maps $p$ and
  $\alpha_1, \ldots, \alpha_{d-1}$ onto the distance $\delta_s$ of the hyper-plane $s$ to the origin
- The parameter space plots the graph of this function
The General Hough Transform

Properties:

- Point in the data space = sinusoid curve in the parameter space
- Point in the parameter space = hyper-plane in the data space
- Points on a common hyper-plane in the data space (cluster) = sinusoid curves intersecting at one point in the parameter space
- Intersection of sinusoid curves in the parameter space = hyper-plane accommodating the corresponding points in data space
Using the Hough Transform to Detect Clusters

So we can use the parameter space to detect correlation clusters:

- Determine all intersection points of at least \( m \) curves in the parameter space (where \( m \) is a minimum number of points in a cluster threshold) to detect \((d-1)\)-dimensional clusters.
- Exact solution (check all pair-wise intersections) is usually too costly.
The CASH algorithm employs a grid-based bisecting search (input: minimum number of points $m$, and minimum number of splits $s$):

- Find dense regions in parameter space
- Define dense grid cells as cells still intersected by $m$ parametrization functions after $s$ splits
- Construct the candidate grid cells by recursively splitting the parameter space along one dimension (in fixed order)
- Always split the candidate cell with the highest number of intersecting functions next (best first)
- Prune candidate cells where less than $m$ functions intersect
- If a cell has been split $s$ times and still intersects $m$ functions, a cluster is found; delete corresponding functions (from other candidate cells) and continue splits until no candidate is remaining
CASH: Clustering in Arbitrary Subspaces Using the Hogh Transform

- Dense grid cell represents \((d-1)\)-dimensional linear structure
- Transform corresponding data objects in corresponding \((d-1)\)-dimensional space and repeat the search recursively to find lower dimensional clusters
- This is comparable to a bottom-up search — WHY?
An Empirical Comparison

(a) Data set DS2.

(b) CASH – Cluster 1 - 5.

(c) 4C – Cluster 1 - 8.

(d) ORCLUS – Cluster 1 - 5.
Summary

• Traditional clustering in high dimensional spaces is most likely meaningless with increasing dimensionality (curse of dimensionality)
• Clusters may be found in (generally arbitrarily oriented) subspaces of the data space
• The partitioning need not be unique (clusters may overlap)
• The subspaces may be axis-parallel or arbitrarily oriented
• Analysis of this general problem:
  • Sub-problem 1: search for clusters
  • Sub-problem 2: search for subspaces
Summary

- Analysis of the 2nd sub-problem (subspace search)
  - A naive solution would examine all possible subspaces to look for clusters
  - The search space of all possible arbitrarily oriented subspaces is infinite
  - We need assumptions and heuristics to develop a feasible solution
- What assumptions did we get to know here to solve the subspace search problem?
  - The search space is restricted to certain subspaces
  - A clustering criterion that implements the downward closure property enables efficient search heuristics
  - The locality assumption enables efficient search heuristics
  - Assuming simple additive models (“patterns”) enables efficient search heuristics
  - ...
• Remember: also for the clustering problem (1st sub-problem) we need assumptions and heuristics that have an impact on the algorithms’ properties
  • Cluster model (what patterns to look for
  • Model specific assumptions/parameters, e.g. number of clusters need to be specified, results are not deterministic e.g. due to randomized procedures, ...
Overview (classification according to sub-problem 2):

- **Matrix-Pattern**
  - Constant values in columns, change of values only on rows

- **Problem**
  - **Subspace / Projected Clustering**
  - **Pattern-based / Bi-Clustering**
  - **Correlation Clustering**

- **Spatial Pattern**
  - Axis-parallel hyperplanes
  - Special cases of axis-parallel to special cases of arbitrarily oriented hyperplanes
  - Arbitrarily oriented hyperplanes
Outlook

• Recall, (unsupervised) outlier detection is very similar (but often considered to be orthogonal) to clustering

• Outlier detection in high dimensional spaces is also often very hard and similar problems can occur, e.g. points are only outliers when considering specific subspaces

• There are several approaches to extend full-dimensional outlier detection methods analogously so that subspace outliers can be detected

• If you want, ask Prof. Kröger to sketch one ...