Supervised learning Unsupervised machine learning Statistical Natural Language Processing * The methods we studied so far are instances of supervised learning In supervised learning, we have a set of predictors x, and want to predict a response or outcome variable y Cağrı Cöltekin University of Tübingen Seminar für Sprachwissenschaft During training, we have both input and output variables
 Training consist of estimating parameters w of a model * During prediction, we are given x and make predictions based on model we ummer Semester 2021 Supervised learning: regression Supervised learning: classification 00 0 0 0 0 * The response (outcome) is a label. . The response (outcome) variable In the example: positive or negative (y) is a quantitative variable. . Given the features (x) we want to Given the features (x₁ and x₂), we want to predict the label of an unknown instance predict the value of u Unsupervised learning Supervised learning . In unsupervised learning, we do not have labels in our training data Our aim is to find useful patterns/structure in the data * The aim is to estimate a set of parameters w for exploratory study of the data
 for augmenting / complementing supervised methods . We define an objective function, and find the parameter values that minimize * Close relationships with 'data mining', 'data science / analytics', 'knowledge the objective The objective typically involves reducing the training error defined based on the true labels in the training data discovery' Most unsupervised methods can be cast as graphical models with hidder variables · Evaluation is difficult: we do not have 'true' labels/valu Today's lecture Clustering: why do we do it? * The aim is to find groups of instances/items that are similar to each of · Clustering: find related groups of instances Applications include Density estimation: find a probability distribution that explains the data - Clustering languag Dimensionality reduction: find an accuracy representation of the data rrate/useful lower dim Clustering (literary) texts, for e.g., authorship attribution
 Clustering woeds for e.g., better parsing
 Clustering documents, e.g., news into topics Unsupervised learning in ANNs (RBMs, autoencoders) Clustering in two dimensional space Similarity and distance

. Unlike classification, we do not have labels

. We want to find 'natural' groups in the data . Intuitively similar or closer data points are grouped toget

Distance measures in Euclidean space

· Euclidean distance

 Manhattan distance: $\|\alpha - b\|_1 = \sum_{i=1}^{k} |\alpha_i - b_j|$

 \bullet The notion of distance (similarity) is important in clustering. A distance measure D_{ν}

- is symmetric D(a, b) = D(b, a)non-negative: D(a, b) ≥ 0 for all a, b, and it D(a, b) = 0 iff a = b

obeys triangle inequality: $D(a,b) + D(b,c) \ge D(a,c)$

. The choice of distance is application specific

We will often face with defining distance measures between linguistic units (letters, words, sentences, documents, ...)

How to do clustering

Most clustering algorithms try to minimize the scatter within each cluster. Which is equivalent to maximizing the scatter between clusters.



 $\sum_{b=1}^{K} \sum_{a \in C} \sum_{b \in C} d(a, b)$

- K-means is a popular method for clustering. Randomly choose centroids, m₁,..., m_K, representing K clusters
- 2. Repeat until convergence Assign each data point to the cluster of the nearest centroid
 Re-calculate the centroid locations based on the assignments
- Effectively, we are finding a local minimum of the sum of squared Euclidean
- distance within each cluster

 $\frac{1}{2} \sum_{k=1}^{K} \sum_{\alpha \in \mathcal{C}_k} \sum_{b \in \mathcal{C}_k} \|\alpha - b\|^2$

K-means clustering: visualization

- · Assign data points to the closest

The data

Set cluster centroids randomly

- centroid Recalculate the centroids

K-means clustering: visualization



 The data · Set cluster centroids randomly

· Set cluster centroids randomly

centroid · Recalculate the centroids

Assign data points to the closest

 Assign data points to the closest . Recalculate the centroids

K-means clustering: visualization



- The data
- · Set cluster centroids randomly
- Assign data points to the closest centroid

K-means clustering: visualization



- * Set cluster centroids randomly Assign data points to the closest centroid

K-means clustering: visualization



- The data
- Set cluster centre
- Assign data points to the closest
- centroid

K-means clustering: visualization



- The data
- · Set cluster centroids rando Assign data points to the clos centroid
- · Recalculate the centroids

K-means clustering: visualization

K-means clustering: visualization



K-means: some issues

- The data
 - Set cluster cer · Assign data points to the clo
 - centroid

 - Recalculate the centroids



- Set cluster of
- Assign data points to the closes centroid
- Recalculate the centroids

How many clusters?

- . K-means requires the data to be in an Euclidean space K-means is sensitive to outliers
- The results are sensitive to initialization

 - There are some smarter ways to select initial points
 One can do multiple initializations, and pick the best
 (with lowest within-group squares)
- It works well with approximately equal-size round-shaped clusters
 We need to specify number of clusters in advance

ber of clusters is defined for some problems, e.g., classifying news

into a fixed set of topics/interests

- * For others, there is no clear way to select the best number of clusters . The error (within cluster scatter) decreases with increasing number of
- clusters, using a test set or cross validation is not useful either
- A common approach is clustering for multiple K values, and picking where there is an 'elbow' in the graph of the error function

How many clusters?

This plot is so times called a scrut plot

K-medoids

- · K-medoids algorithm is an alternation of K-means
 - Instead of calculating centroids, we try to find most typical data point (medoids) at each iteration
 - K-medoids can work with distances, does not need feature vectors to be in an Euclidean space
 It is less sensitive to outliers
 - . It is computationally more expensive than K-means

Hierarchical clustering

* Hierarchical clustering operates on distances (or similarities)

- The result is a binary tree called dendrogram
- · Dendrograms are easy to interpret (especially if data is hierarchical)
- The algorithm does not commit to the number of clusters K from the start, the dendrogram can be 'cut' at any height for determining the clusters

. There are two main 'modes of operation':

Bottom-up or age/foundative (utubering)

- starts with individual data points,

- starts with individual data points,

- merges the clasters until all data is in a single clus

Top-down or divisitve clustering

. Instead of a flat division to clusters as in K-means, hierarch builds a hierarchy based on similarity of the data points

r arcrisive cuistering

• starts with a single cluster,

• and splits until all leaves are single data points

Agglomerative clustering 1. Compute the similarity/distance matrix

Hierarchical clustering

- 2. Assign each data point to its own
- 3. Repeat until no clusters left to merge similar to each other

 - Merge them into a single cluster

Agglomerative clustering demonstration



How to calculate between cluster distances

Complete maximal inter-cluster distar Single minimal inter-cluster distance Average mean inter-cluster distance Centroid distance between the



How to calculate between cluster distances

Single minimal inter-cluster distance Average mean inter-cluster dist Centroid distance between the



How to calculate between cluster distan

Single minimal inter-cluster dista Average mean inter-cluster distance Centroid distance between the



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Single minimal inter-cluster dist Average mean inter-cluster dis roid distance between the



iote: we only need distances, (feat

Clustering evaluation

Evaluating clustering results is often non-trivial

- Internal evaluation is based a metric that aims to indicate 'good clustering': e.g., Dunn index, gap statistic, silhouette
- External metrics can be useful if we have labeled test data: e.g., V-measure, B³cd F-score
- The results can be tested on the target application: e.g., word-clusters
- evaluated based on their effect on parsing accuracy
- . Human Judgments, manual evaluation 'looks good to me'

objects in the same cluster b(i) average distance between object i and and objects in the closest cluster

Clustering evaluation mede vilhoust



· We want clusters that contain

Clustering evaluation

- members of a single gold-standard class (homogeniety)
- We want all members of a class to be in a single cluster (completeness)

Note the similarity with precision and recall

Density estimation

- K-means treats all data points in a cluster equally
 A 'soft' version of K-means is density estimation for Gaustine Company of the Co
- where
- We assume the data comes from a mixture of K Gaussian distributions
 We try to find the parameters of each distribution (instead of centroids) the maximizes the likelihood of the data
- · Unlike K-means, mixture of Gaussians assigns probabilities for each data
- point belonging to one of the clusters
- It is typically estimated using the expectation-maximization (EM) algorithm

- · Principal component analysis (PCA) is a method of dimensionality reduction
- · PCA maps the original data into a lower dimensional space by a linear
- transformation (rotation) · The transformed lower-dir
 - (=information) in the input

Principal component Analysis

- . PCA can be used for
 - visualization
 data compression



$$\begin{split} A = \begin{bmatrix} \cos\theta & -\sin\theta \\ \sin\theta & \cos\theta \end{bmatrix} = \begin{bmatrix} \frac{3}{2} & -\frac{4}{2} \\ \frac{3}{2} & \frac{5}{2} \end{bmatrix} \\ \text{p1} = A \times \begin{bmatrix} -5 \\ 0 \end{bmatrix} = \begin{bmatrix} -3 \\ -4 \end{bmatrix} \quad \text{p2} = A \times \begin{bmatrix} 0 \\ 0 \end{bmatrix} = \begin{bmatrix} 0 \\ 0 \end{bmatrix} \text{p3} = A \times \begin{bmatrix} 5 \\ 0 \end{bmatrix} = \begin{bmatrix} 3 \\ 4 \end{bmatrix} \end{split}$$

We can recover the original points p dimensionality of the data is only 1.

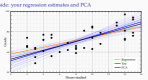
Why do we want to reduce the dimensionality

- - . If we use the data for other ML methods, we reduce the computation time
 we may avoid 'the curse of dimensionality

 - · Decorrelation is useful in some applications
 - We compress the data (in a lossy way)

 - . We eliminate noise (assuming a high signal to noise ratio)

Aside: your regression estimates and PCA



Clustering: some closing notes

- * We do not have proper evaluation procedures for clustering results (for unsupervised learning in general)
- Some clustering methods are unstable, slight changes in the data or parameter choices may change the results drastically Approaches against instability include some validation methods, or producing 'probabilistic' dendrograms by running clustering with different producing 'probabilistic' dendrograms by running clustering with different producing the prod

Density estimation using the EM algorithm

- . The EM algorithm (or its variations) is used in learning models with latent/hidden variables * It is closely related to the K-means algorithm
- 1. Initialize the parameters (e.g., randomly) of K multivariate normal distributions (μ, Σ)
- 2. Iterate until convergence:
- E-step Given the parameters, compute the membership 'weights', the probability of each data point belonging to each distribution M-step Re-scitmate the mixture density parameters using the calculated membershi
- weights in the E-step

PCA: a toy example



. How many dimensions do we have:

- How many dim
- · Short divergence: calculate the
 - $\Sigma = \begin{bmatrix} \frac{18}{3} & 8 \\ \frac{8}{8} & \frac{32}{3} \end{bmatrix}$
 - What is the correlation and x₂?



* What if the variables were not perfectly bu strongly correlated? We could still do a similar transformation

Discarding z₂ results in a small reconstruction error:

 $p1 - A \times \begin{bmatrix} -5 \\ 0 \end{bmatrix} - \begin{bmatrix} -3 \\ -4 \end{bmatrix}$ Note: z₁ (also z₂) is a linear combination of original variables

Different views on PCA



- Find the direction of the largest variance
- · Find the projection with the least
- · Find a lower dimensional latent
- Gaussian variable such that the
 - observed variable is a mapping of the latent variable to a higher dimensional space (with added

How to find PCs

- When viewed as maximizing variance or redu write the appropriate objective function and find the vectors that minimize it $\star\,$ In latent variable interpretation, we can use EM as in estimating mixtures of
- The principal components are the eigenvectors of the correlation matrix, where large eigenvalues correspond to components with large variation
- A numerically stable way to obtain principal components is doing singular value decomposition (SVD) on the input data

- PCA as matrix factorization (eigenvalue decomposition) • One can compute PCA by decomposing the covariance mat $\Sigma = X^T X$)
 - $\Sigma = U \Lambda U^T$
 - the columns of U are the principal components (eigenvectors) Λ is a diagonal matrix of eigenvalues
 - Another option is SVD, which factorizes the input vector $(k \text{ variables} \times n \text{ data points})$ as
 - X UDV
- $\begin{array}{ll} & \ U \ (k \times k) \ contains the eigenvectors \\ & \ D \ (k \times n) \ diagonal \ matrix \ D^2 = A \\ & \ V^* \ is \ a \ n \times n \ unitary \ matrix \end{array}$

Unsupervised learning in ANNs

- · Restricted Boltzmann machines (RBM) similar to the latent variable models (e.g., Gas
- Automodere train a constrained feed-forward network to predict its output
- representation learned by hidden layers as hidden variables (h), and learn p(x, h) that maximize the probability of the (unlabeled)data

The distribution defined by RBMs



$$p(h,x) = \frac{e^{h^TWx}}{Z}$$

This calculation is intractable (Z is difficult to calculate or calculate and conditional distributions are easy to calculate

$$p(h|x) = \prod_j p(h_j|x) = -\frac{1}{1+e^{\mathbf{W}_j x}}$$

$$p(x|h) = \prod_k p(x_k|h) = \frac{1}{1 + e^{\mathbf{W}_k^T h}}$$

Autoencoders

- · Autoencoders are standard feed-forward . The main difference is that they are
- trained to predict their input (they try to learn the identity function) . The aim is to learn useful
- representations of input at the hidden layer
- The weights are often shared/tied $(W^* W^T)$

Over-complete autoencoders



Unsupervised pre-training

- over-complete if there are more hidden units than inputs . The network can normally memorize the
- input perfectly
- This type of networks are useful if
- trained with a regularization term resulting in sparse hidden units (e.g., L1 regularization)

- A common use case for RBMs and autoencoders are as pre-tra for supervised networks Autoencoders or RBMs are trained using unlabeled data
- The weights learned during the unsupervised learning is used for initializing the weights of a supervised network
- This approach has been one of the reasons for success of deep networks

Some practical notes on PCA

- · Variables need to be centered * Scales of the variables matter, standardizing may be a good idea depending
 - on the units/scales of the individual variables * The sign/direction of the principal component (vector) is not important
 - If there are more variables than the data points, we can still calculate the principal components, but there will be at most n-1 PCs
 - . PCA will be successful if variables are correlated, there are extensions for
 - dealing with nonlinearities (e.g., kernel PCA, ICA, t-SNE)

Restricted Boltzmann machines (RBMs)



- * RBMs are unsupervised latent variable models, they learn only from unlabeled data
- * They are generative models of the joint probability $p(\mathbf{h}, \mathbf{x})$ · They correspond to undirected graphical
- models . No links within layers
- . The aim is to learn useful features (h)

Learning in RBMs

- ility the model assigns to the input, p(x), or equivalently minimize $-\log p(x)$ $\bullet\,$ In general, this is computationally ex
- * Contrastive divergence algorithm is a well known algorithm that efficiently finds an approximate solution

Under-complete autoencoders



- · An autoencoder is said to be under-complete if there are fewer hidden units than inputs
- The network is forced to learn a compa representation of the input (compress) An autoencoder with a single hidden
- layer approximates the PCA
- . We need multiple layers for learning non-linear features

Denoising autoencoders



- Instead of providing the exact input, we introduce noise by - randomly setting some inputs to 0 (dropout)
 - ng random (Gaussian) noisi
- Network is still expected to reconstruct the original input (without noise)

Summary

- In unsupervised learning, we do not have labels. Our aim is to find/exploit (latent) structure in the data . Unsupervised methods try to discover 'hidden' structure in the data
- Clustering finds groups in the data Density estimation estimates parameters of latent probability distributions
 Dimensionality reduction transforms the data in a low dimensional space
 while keeping most of the information in the original data
- More ML: sec
- Common CL tasks: tokenization, morphology, syntactic parsing, (lexical)
- . Some NLP applications: text classification, and maybe more

Derivation of PCA by maximizing the variance We focus on the first PC (z_1) , which maximizes the variance of the data onto not of the position of the direction, so we choose z_1 to be a unit vector $(z_1 -1)$. Remember that to project a vector onto another, we simply use dot product, So the projected data points are z_1 , for $1-1,\dots,N$. The variance of the projected data point (fine two wars to maximize) is, $\sigma_{x_1} = \frac{1}{N} \sum_{i=1}^N \sum_{i=2}^N z_i z_i z_i z_i^{-1} z_i^{-1} \sum_{i=1}^N z_i z_i z_i^{-1} z_i^{-1} z_i^{-1}$ where E_n is the covariance matrix of the unprojected data	Derivation of PCA by maximizing the variance (cont.) • The problem becomes maximize $z_1^2 z_2^2$ with the constraint $[z_1] - z_1^2 z_1 - z_1^2 z_1^2$ • Turning it this a succentrainted optimization problem with Lagrange multipliers, we minimize $z_1^2 z_1 + \lambda_1 (1 - z_1^2 z_1)$ • Taking the derivative and setting it to 0 gives us $\underline{z}_1 = \lambda_2 z_1 - \lambda_2 z_1$ • Taking the derivative and setting it to 0 gives us $\underline{z}_1 = \lambda_2 z_1 - \lambda_2 z_2$ Note: by definition, z_1 is an approximate $z_1 = z_1 - z_2 z_2$. Note by definition, z_1 is an approximate expensation of $z_1 = z_1 - z_2 z_2$. Since the consequence of $z_1 = z_2 z_2 z_2$ and $z_2 z_2 z_2 z_2 z_2$ and $z_1 = z_2 z_2 z_2 z_2 z_2$.
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