Mixture models and density estimation with PYTHON

Objective:

To understand the Gaussian mixture model; how Expectation-Maximization is used to estimate the model parameters; and how information criteria and cross-validation is used to choose the number of clusters. Also, you should understand how frequent itemsets can be extracted by the Apriori algorithm and be able to calculate and interpret association rules in terms of support and confidence.

Material: Lecture notes "Introduction to Machine Learning and Data Mining" C17, C18 as well as the files in the exercise 11 folder available from Campusnet.

Preparation: Exercises 1-10

Part 1: Group discussion (max 15 min)

For the group discussion, each group should have selected a *discussion leader* at the previous exercise session. The purpose of the discussion leader is to ensure all team members understands the answers to the following two questions:

Multiple-Choice question: Solve and discuss problem 18.1 from chapter 18 of the lecture notes. Ensure all group members understand the reason why one of the options is true and why the other options can be ruled out.

Discussion question: Discuss the following question in the group

• Explain the intuition behind a.r.d.. Why not simply use the density? Sketch an example where the a.r.d. is more appropriate and explain why

Part 2: Programming exercises

Piazza discussion forum: You can get help by asking questions on Piazza: https://piazza.com/dtu.dk/fall2016/02450

Software installation: Extract the Python toolbox from Campusnet. Start Spyder and add the toolbox directory (<base-dir>/02450Toolbox_Python/Tools/) to PYTHONPATH (Tools/PYTHONPATH manager in Spyder). Remember the purpose of the exercises is not to re-write the code from scratch but to work with the scripts provided in the directory <base-dir>/02450Toolbox_Python/Scripts/

Representation of data in Python:

	Python var.	Type	Size	Description
	Х	numpy.matrix	$N \times M$	Data matrix: The rows correspond to N data objects, each of which contains M attributes.
	attributeNames	list	$M \times 1$	Attribute names: Name (string) for each of the M attributes.
	N	integer	Scalar	Number of data objects.
	M	integer	Scalar	Number of attributes.
Regression	у	numpy.matrix	$N \times 1$	Dependent variable (output): For each data object, y contains an output value that we wish to predict.
Classification	у	numpy.matrix	$N \times 1$	Class index: For each data object, y contains a class index, $y_n \in \{0, 1,, C-1\}$, where C is the total number of classes.
	classNames	list	$C \times 1$	Class names: Name (string) for each of the C classes.
	С	integer	Scalar	Number of classes.
Cross-validation				All variables mentioned above appended with _train or _test represent the corresponding variable for the training or test set.
	$\star_{ extsf{-}}train$	_	_	Training data.
	*_test			Test data.

11.1 The Gaussian Mixture Model and the EM algorithm

In the exercise last week we considered k-means clustering and hierarchical clustering. Today we will consider clustering based on the Gaussian mixture model (GMM). We recall that the multivariate Gaussian distribution is given by

$$\mathcal{N}(\boldsymbol{x}_i|\boldsymbol{\mu}_{(k)},\boldsymbol{\Sigma}_{(k)}) = \frac{1}{\sqrt{\det(2\pi\boldsymbol{\Sigma}_{(k)})}} \exp\{-\frac{1}{2}(\boldsymbol{x}_i - \boldsymbol{\mu}_{(k)})^{\top}\boldsymbol{\Sigma}_{(k)}^{-1}(\boldsymbol{x}_i - \boldsymbol{\mu}_{(k)})\}.$$

In the Gaussian mixture model we use a mixture of K multivariate Gaussians to model the data. We give each Gaussian a mixture coefficient w_k between zero and one such that all coefficients sum to one, $\sum_{k=1}^{K} w_k = 1$. The probability of a data vector \boldsymbol{x}_i is then modeled as a weighted sum of Gaussian distributions,

$$p(\boldsymbol{x}_i|\boldsymbol{w}, \{(\boldsymbol{\mu}_{(1)}, \boldsymbol{\Sigma}_{(1)}), \dots, (\boldsymbol{\mu}_{(K)}, \boldsymbol{\Sigma}_{(K)})\}) = \sum_{k=1}^K w_k \cdot \mathcal{N}(\boldsymbol{x}_i|\boldsymbol{\mu}_{(k)}, \boldsymbol{\Sigma}_{(k)})$$

The parameters of the model, which comprises the mixture coefficients and the K means and covariance matrices, are found by the Expectation-Maximization (EM) algorithm that progress in the following way:

- 1. Initialize the mixture coefficient \boldsymbol{w} , mean and covariance of each Gaussian $\boldsymbol{\mu}_{(k)}$ and $\boldsymbol{\Sigma}_{(k)}$ by random.
- 2. (E-step) Calculate the expectation $P(k|\boldsymbol{x}_i, \boldsymbol{w}, \{(\boldsymbol{\mu}_{(1)}, \boldsymbol{\Sigma}_{(1)}), \dots, (\boldsymbol{\mu}_{(K)}, \boldsymbol{\Sigma}_{(K)})\})$ for each data point.
- 3. (M-step) Optimize \boldsymbol{w} and $\{(\boldsymbol{\mu}_{(1)},\boldsymbol{\Sigma}_{(1)}),\ldots,(\boldsymbol{\mu}_{(K)},\boldsymbol{\Sigma}_{(K)})\}$ by maximizing the expected likelihood.
- 4. Keep doing 2 and 3 until the clusters do not change or a maximum number of iterations have progressed.

The EM-algorithm is closely related to the k-means algorithm but rather than operating with hard assignment of each data point, each data point is assigned a given probability of belonging to each cluster based on Bayes rule,

$$P(k|\mathbf{x}_i, \mathbf{w}, \{(\boldsymbol{\mu}_{(1)}, \boldsymbol{\Sigma}_{(1)}), \dots, (\boldsymbol{\mu}_{(K)}, \boldsymbol{\Sigma}_{(K)})\}) = \frac{w_k \cdot N(\mathbf{x}_i | \boldsymbol{\mu}_{(k)}, \boldsymbol{\Sigma}_{(k)})}{\sum_{k=1}^K w_k \cdot N(\mathbf{x}_i | \boldsymbol{\mu}_{(k)}, \boldsymbol{\Sigma}_{(k)})},$$

while each cluster is not only described by its center (mean) but also by its covariance. In order to get hard assignments of the data points from the results obtained by the above EM-algorithm we assign the observations to the clusters with highest probability.

An important property of the estimated Gaussian mixture model is that the GMM density integrates to one, i.e.

$$\int \sum_{k} w_k \cdot \mathcal{N}(\boldsymbol{x}|\boldsymbol{\mu}_{(k)}, \boldsymbol{\Sigma}_{(k)}) d\boldsymbol{x} = \sum_{k} w_k \int \mathcal{N}(\boldsymbol{x}|\boldsymbol{\mu}_{(k)}, \boldsymbol{\Sigma}_{(k)}) d\boldsymbol{x} = 1$$

This follows from the fact that each Gaussian integrates to one $\int \mathcal{N}(\boldsymbol{x}|\boldsymbol{\mu}_{(k)},\boldsymbol{\Sigma}_{(k)})d\boldsymbol{x} = 1$ and the constraint $\sum_{k=1}^K w_k = 1$. An important consequence of this is that when additional clusters are introduced they have to take density mass from existing clusters.

11.1.1 Inspect and run the script $ex11_1_1.py$. The script loads the synth1 data from the file pata/synth1 with the loadmat function and fits a K=4 component Gaussian mixture model to the data. Notice how the script makes a scatter plot of the data and the clustering using the clusterplot function in the toolbox.

Script details:

- To fit a Gaussian Mixture Model in Python you can use the class GMM from the package sklearn.mixture. You will need two methods: fit and predict. Type help(GMM) to learn how to use the class.
- · When creating GMM object you can specify number of clusters, covariance type (full/diagonal), number of repetitions with different initial seeds.
- You can extract the fitted cluster means (centroids) by calling the method means_ of GMM class object.
- · Type clusterplot(X,clusterid,centroids,cov_matrices) to plot the data and clustering.
- · Type help(clusterplot) to learn more about how to use the clustering plot tool in the toolbox. Note that you can specify covars parameter to plot ellipsoids of the gaussians corresponding to GMM.

Sometimes the estimated models for the same value of K are not necessarily the same due to different initial placement of the centroids. In such cases you might try setting the parameter n_init to 10 or more. What will this do, and why should it solve the problem?

Try to change the structure of the covariance matrix to be a diagonal matrix by setting the parameter covariance_type to 'diagonal'. How does this affect the generated clusters? What do you think might be benefits and drawbacks of restricting the covariance matrices?

11.1.2 **Discussion:** In k-means based on Euclidean distance observations are assigned the centroids they are the closest to. Is this also the case when clustering by the Gaussian mixture model or is it possible that points are assigned a cluster that is further away than other clusters in terms of Euclidean distance?

Can the scaling of the variables seriously affect the results we get when clustering by the GMM or is the model able to take the scaling of the data into account?

11.1.3 **Discussion:** For supervised learning we used cross-validation to evaluate performance and estimate the number of parameters in our models. In last weeks exercise, we found that this approach could not be used in the k-means algorithm. What happens if we validate the number of clusters for Gaussian mixture model based on the EM algorithm using cross-validation, i.e., split the data into training and test data, train the model on the training data and evaluate how likely the test data points

are based on the learned parameters, \boldsymbol{w} and $\{\boldsymbol{\mu}_{(1)}, \boldsymbol{\Sigma}_{(1)}, \dots, \boldsymbol{\mu}_{(K)}, \boldsymbol{\Sigma}_{(K)}\}$ using the logarithm of the likelihood of the test data,

$$\log L^{\text{test}} = \sum_{i=1}^{N_{\text{test}}} \log[p(\boldsymbol{x}_i^{(\text{test})} | \boldsymbol{w}, \{(\boldsymbol{\mu}_{(1)}, \boldsymbol{\Sigma}_{(1)}), \dots, (\boldsymbol{\mu}_{(K)}, \boldsymbol{\Sigma}_{(K)})\})]?$$

Script details:

- Remember, that in the GMM each cluster is represented by a mean vector and a covariance matrix. In one dimension, it is the well known bell-shaped probability distribution.
- · If we have 100 data points and use 1 cluster, what would be the optimal solution for the GMM using the EM-algorithm? How well would this solution generalize to test data?
- · If we have 100 data points and use 100 clusters, what would be the optimal solution? How well would this solution generalize?
- · What if we use some intermediate number of clusters?

Apart from cross-validation the optimal number of clusters are sometimes derived by penalizing model complexity based on the Bayesian Information Criteria (BIC) or Akaike's Information Criteria (AIC). The two information criteria are defined by

$$BIC = -2 \log L + p \log(N), \quad AIC = -2 \log L + 2p$$

where p is the number of free parameters in the model, i.e., the total number of estimated variables in \boldsymbol{w} , $\{\boldsymbol{\mu}_1, \boldsymbol{\Sigma}_1, \dots, \boldsymbol{\mu}_K, \boldsymbol{\Sigma}_K\}$ whereas N is the number of observations and $\log L$ is the log likelihood of observing the data, i.e.,

$$\log L = \sum_{i=1}^{N} \log[p(\boldsymbol{x}_i | \boldsymbol{w}, \{\boldsymbol{\mu}_{(1)}, \boldsymbol{\Sigma}_{(1)}, \dots, \boldsymbol{\mu}_{(K)}, \boldsymbol{\Sigma}_{(K)}\})].$$

The two information criteria define a trade-off between modeling the data well, i.e., minimize $-2 \log L$, and penalizing complexity of the model, i.e., $M \log(N)$ and 2M respectively, such that the model with lowest AIC and BIC value indicates the model with best trade-off. Note, that AIC and BIC do not require splitting the data in test and training sets, but are computed directly on the whole training data.

- 11.1.4 **Discussion:** Which of the two information criteria BIC and AIC will in general penalize model complexity the most?
- 11.1.5 Inspect and run the script $ex11_1_5.py$. We will use BIC, AIC, and 10-fold crossvalidation to assess the best number of clusters for the synth1 data set. Use the script to compute the three measures for $K=1,\ldots,10$, and use 10 replicates to avoid bad solutions due to poor initial conditions. Script details:

- As before, use sklearn.mixture.GMM class to fit the models.
- · As usual, use the module sklearn.cross_validation to set up the crossvalidation folds.
- · GMM class has bic and auc methods to compute BIC and AUC scores automatically. Type help(GMM) to read more.
- To evaluate the model fit on the test set in terms of negative log likelihood, you can use the method score (after fitting the model). For instance:
 NLOGL = -gmm.score(X_test).sum().

What are the benefits and drawbacks of AIC and BIC versus cross-validation?

- 11.1.6 (Optional): Use a Gaussian mixture model to cluster the Old Faithful data set, Data/faithful, using AIC, BIC, or crossvalidation to select the number of clusters.
- 11.1.7 (Optional): Use a Gaussian mixture model to cluster the Iris data set, Data/iris.xls, using AIC, BIC, or crossvalidation to select the number of clusters.

11.2 Density estimation

We will apart from the Gaussian mixture model (we just used) now consider the following two approaches to density estimation: kernel density estimation, and knearest neighbors density estimation.

Kernel density estimation is a non-parametric method of estimating the probability density function of a random variable. Inference about the population are made, based on a finite data sample. The estimated kernel density for a variable x is given by

$$f(x) = \frac{1}{N} \sum_{i=1}^{N} K(x - x_i)$$
 (1)

where K is the kernel function that must integrate to one, N is the sample size, and f(x) is the estimated density (see also figure 1.)

We will presently consider density estimation based on the kernel formed by the normal distribution ¹, i.e.,

$$K(x - x_i) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left(-\frac{(x - x_i)^2}{2\sigma^2}\right).$$

We will initially consider an artificially generated one-dimensional data set formed by a mixture of three Gaussians defined as follows

$$p(x) = \frac{1}{3}\mathcal{N}(x|1,1) + \frac{1}{3}\mathcal{N}(x|3,0.5) + \frac{1}{3}\mathcal{N}(x|6,2) . \tag{2}$$

¹Notice, that for this choice of kernel function the kernel density estimation can be considered a Gaussian mixture model where the number of clusters is the same as the number of observations, i.e., there is a Gaussian around each observation, while the mixing coefficient is fixed to be the same for all classes, $w_n = 1/N$.

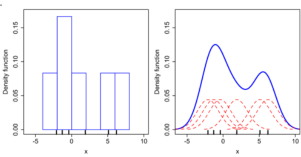


Figure 1: Estimation of the data density by histogram (left) and kernel density estimation (right). The Kernel specifies a shape around each data point (here a univariate normal distribution) and the density is estimated by summing over these Gaussians placed at each observation

11.2.1 Inspect and run the script $ex11_2_1.py$. Notice how the script generates 1000 data objects according to the model in equation (2), and plots a histogram of the data with 50 bins in the range -10 to 10.

Script details:

- · You can generate data from the Gaussian mixture model in the following way: First, you choose one of the components by random according to the mixture coefficients. Then, you generate the data according to the Gaussian distribution for the chosen component.
- Type help(np.random.multinomial) to learn how to generate a random variable from a discrete distribution.
- · You can use help(np.random.normal) function to draw samples from normal distribution (particular gaussian from the mixture).
- · The function numpy.hist can be used to plot a histogram. Use help(numpy.hist) to get help.
- To define the bins at which the histogram should be evaluated, you can define a vector of values and pass it to hist. To get 50 bins evenly spaced between -10 and 10, you can use the command x=np.linspace(-10,10,50).

Discussion:

- Can you identify each component of the mixture in the plot and its associated parameters (mean, variance, mixture coefficient?)
- ⋄ Try changing the parameters of the mixture to see the effect on the resulting density.
- ♦ Try varying the number of bins in the histogram.
- 11.2.2 Inspect and run the script $ex11_2_2.py$. Explain how the script estimates the density using a kernel density estimator with a Gaussian kernel and a kernel width of 1, and plot the density on the range -10 to 10.

Script details:

- Use gaussian_kde from the module scipy.stats.kde to fit a kernel density estimator.
- · To define at which x-values the KDE should be evaluated, you can define a vector of values and pass it to the evaluate method of the fitted KDE estimator. For example, to get 100 points evenly spaced between -10 and 10, you can use the command x=np.linspace(-10,10,100)

Discussion:

- ♦ Compare the result to the histogram.
- ♦ Try varying the kernel width in the KDE.
- ♦ How could you select an optimal kernel width?

Consider the following measure of density of the i^{th} observation x_i given based on its k-nearest neighbors (KNN)

density_{$$\mathbf{X}_{\setminus i}$$} $(\mathbf{x}_i, K) = \frac{1}{\frac{1}{K} \sum_{\mathbf{x}' \in N_{\mathbf{X}_{\setminus i}}(\mathbf{x}_i, K)} d(\mathbf{x}_i, \mathbf{x}')},$ (3)

where $N_{\boldsymbol{X}}(\boldsymbol{x},K)$ is the K observations in \boldsymbol{X} which are nearest to \boldsymbol{x} , and $\boldsymbol{X}_{\backslash i}$ is simply \boldsymbol{X} with observation i removed: $\boldsymbol{X}_{\backslash i}^T = \begin{bmatrix} \boldsymbol{x}_1 & \boldsymbol{x}_2 & \cdots & \boldsymbol{x}_{i-2} & \boldsymbol{x}_{i-1} & \boldsymbol{x}_{i+1} & \boldsymbol{x}_{i+2} & \cdots & \boldsymbol{x}_N \end{bmatrix}$. This estimates the density as the inverse of the average distance to the k nearest neighbors.

If the dataset contains regions of varying densities it can be useful to define a notion of density that is relative to the neighborhood of the object. The following average relative density is one such approach

$$\operatorname{ard}_{\boldsymbol{X}}(\boldsymbol{x}_{i}, K) = \frac{\operatorname{density}_{\boldsymbol{X}_{\setminus i}}(\boldsymbol{x}_{i}, K)}{\frac{1}{K} \sum_{\boldsymbol{x}_{j} \in N_{\boldsymbol{X}_{\setminus i}}(\boldsymbol{x}_{i}, K)} \operatorname{density}_{\boldsymbol{X}_{\setminus j}}(\boldsymbol{x}_{j}, K)}.$$
 (4)

11.2.3 Inspect and run $ex11_2_3.py$ and see how the script can be used to estimate the density using equation (3) as well as the average relative density using equation (4) with the 200-nearest neighbors based on the Euclidean distance measure, and plot the density on the range -10 to 10.

Script details:

- · To define at which x-values the nearest neighbor density estimator should be evaluated, you can define a vector of values, e.g., to get 100 points evenly spaced between -10 and 10, you can use the command x=np.linspace(-10,10,100).
- · To find the nearest neighbors and the distances needed to compute equation (3), you can fit the NearestNeighbors model from the module sklearn.neighbors and then use the method kneighbors() of to extract nearest neighbors distances.

Discussion:

♦ Try different values for the number of neighbors.

♦ Compare your results to the three previous exercises.

11.3 Outlier detection

11.3.1 Inspect and run the script ex11_3_1.py. Prior to running the script you need to execute the command X[-1,0]=-10 to add an outliner at -10 to the data. The script uses a kernel density estimator (as in exercise 11.2.2) with a Gaussian kernel to estimate the density at the x-values in the data set. Verify that the outlier you have introduced has the lowest density. Notice how the bar plot corresponds to the density of the 20 lowest-density points.

Script details:

- You originally had N = 1000 data objects. What is the index of the introduced outlier?
- · To get the KDE for each data object in the data set, you have to fit KDE model first (class gaussian_kde() from the module scipy.stats.kde), and then use evaluate() method.
- To find the indices and values of the 20 lowest-density points, you can use the methods argsort() and sort() respectively, on the output from evaluate() method.
- · To make a bar plot, use the function bar.

Discussion:

- ♦ Can the outlier be detected from this plot?
- ♦ Try different values of the kernel width.
- ♦ What happens when the kernel width is too large / too small?
- 11.3.2 The function <code>gausKernelDensity()</code> in the toolbox implements density estimation by the gaussian kernel density estimator using a very efficient implementation of leave-one-out cross-validation. I.e. the density of each observation is estimated from all other observations not including the observation itself in the estimate. Inspect and run the script <code>ex11_3_2.py</code> where it is used. Use the script to estimate the optimal kernel width by evaluating the estimated densities for a range of different kernel widths. Plot the leave-one-out density of the 20 lowest-density points in a bar plot. Script details:
 - · To get the leave-one-out KDE for each data object in the data set for a kernel width of 5, you can use the command f,log_f=gausKernelDensity(X,5).
 - · The log-density for all observations is given by logP=log_f.sum(). The optimal kernel width is the width with highest logP.
 - · To find the 20 lowest-density points, you can again use the function sort to sort the output from ksdensity.
 - · To make a bar plot, use the function bar.

11.3.3 (Optional): Repeat exercise 11.3.1 using the KNN-density estimator as well as the KNN-average relative density.

11.4 Hand written digits

We will in this part of the exercise investigate if some of the hand written digits of each class can be considered outliers. To do this, inspect and run the script ex11_4_1.py.

- 11.4.1 Load the hand written digits data set from the file Data/digits.mat with loadmat() function. Restrict your analysis to images of the digit "2" by the command X=X[y.A.ravel()==2,:].
 - 1) Use the function gausKernelDensity() from toolbox that implements density estimation by the gaussian kernel density estimator using the very efficient implementation of leave-one-out density estimation (as in exercise 11.3.2). Estimate the optimal kernel width evaluating a range of different widths. Make a bar plot of the leave-one-out densities of the 20 lowest-density data objects in the data set based on the optimal kernel width.
 - 2) Use the KNN density estimation method (as in exercise 11.2.4) based on the Euclidean distance measure and using K = 5 neighbors. Make a bar plot of the densities of the 20 lowest-density data objects in the data set.
 - 3) Use the KNN average relative density estimation method (as in exercise 11.2.4) also based on the Euclidean distance measure and using K=5 neighbors. Make a bar plot of the densities of the 20 lowest-density data objects in the data set.

Plot the images of the 20 data objects with the lowest density / highest outlier score.

Script details:

The images are 16 by 16 pixels stored as 256 dimensional vectors. To plot an image, you must reshape the vector to a 16 by 16 matrix and plot it using the imshow() function. For example, to plot the i'th image in X, use the command: imshow(np.reshape(X[i-1,:], (16,16)).T, cmap=cm.binary).

Discussion:

- ♦ Does the three bar plots reveal any possible outliers in the data set?
- ♦ Can you identify any data objects that really are outliers?
- ♦ Do the three methods agree?
- ⋄ Try plotting 20 random images of hand written 2's (for example the first 20 in the data set) for comparison. In what sense are the low-density 2's different from the randomly chosen ones?

- ♦ Try other digits than "2". Does the method in general identify digits that are outliers?
- ♦ Try the method on the whole data set with all digits (warning: this might take a long time to compute.)

11.5 Tasks for the report

Analyze your data by the GMM and use cross-validation to estimate the number of clusters. (If you encounter the problem "Ill-conditioned covariance" set the regularization parameter min_covar of gmm to a small constant, say 10^{-6}). Evaluate how well the clusters of the GMM model correspond to class labels using the cluster validity measures from last week by assigning observations to the cluster having highest probability. Apply the outlier scoring methods from last exercise in order to rank all the observations in terms of the Gaussian Kernel density (using the efficient leave-one-out density estimation approach), KNN density, and KNN average relative density for some suitable K. Discuss whether it seems there may be outliers in your data according to the three scoring methods.

References