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LAB12 tutorial for Machine Learning Clustering with GMM

The document description are designed by Jla Yanhong in 2022. Nov. 21th

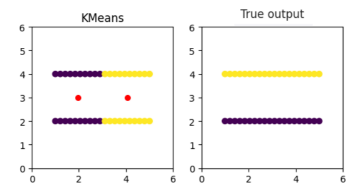
1 Objective

- · Understand GMM clustering algorithm theory
- Implement the GMM clustering algorithm from scratch in python
- · Complete the LAB assignment and submit it to BB or sakai.

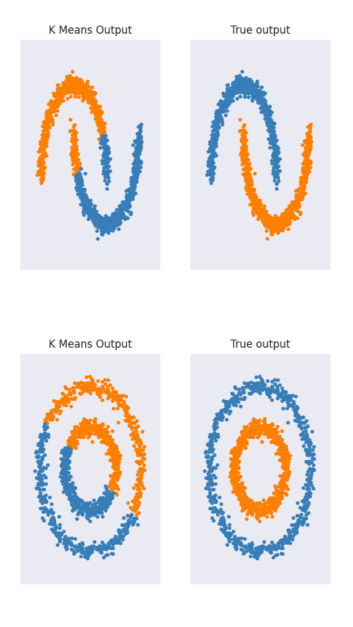
1.1 Drawbacks of k-means Clustering

The k-means clustering concept sounds pretty great, right? It's simple to understand, relatively easy to implement, and can be applied in quite a number of use cases. But there are certain drawbacks and limitations that we need to be aware of. K-means often doesn't work when clusters are not round shaped

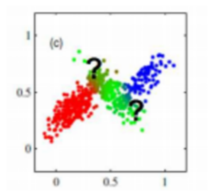
First, KMeans doesn't put data points that are far away from each other into the same cluster, even when they obviously should be because they underly some obvious structure like points on a line, for example.



Second, KMeans performs poorly for complicated geometric shapes such as the moons and circles shown below.



In addition,k-means doesn't work when clusters are may overlap.



Hence, we need a different way to assign clusters to the data points. So instead of using a distance-based model, we will now use a distribution-based model. And that is where <code>Gaussian Mixture Models</code> come into this lab!

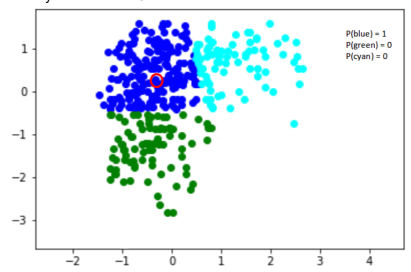
1.2 Gaussian mixture model (GMM)

Gaussian Mixture Models (GMMs) assume that there are a certain number of Gaussian distributions, and each of these distributions represent a cluster.

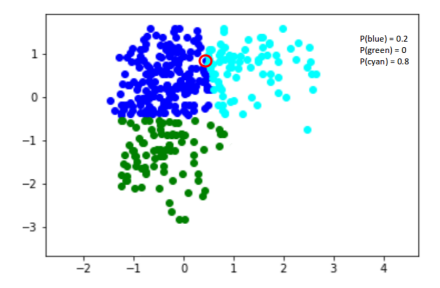
Gaussian Mixture Models are probabilistic models and use the soft clustering approach for distributing the points in different clusters.

Let us take an example that will make it easier to understand.

Here, we have three clusters that are denoted by three colors – Blue, Green, and Cyan. Let's take the data point highlighted in red. The probability of this point being a part of the blue cluster is 1, while the probability of it being a part of the green or cyan clusters is 0.



Now, consider another point – somewhere in between the blue and cyan (highlighted in the below figure). The probability that this point is a part of cluster green is 0, right? And the probability that this belongs to blue and cyan is 0.2 and 0.8 respectively.



Gaussian Mixture Models use the soft clustering technique for assigning data points to Gaussian distributions.

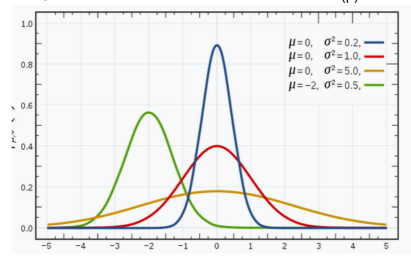
1.3 The Gaussian Distribution

In a one dimensional space, the probability density function of a Gaussian distribution is given by:

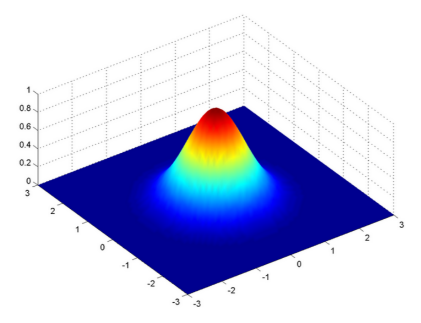
$$\mathcal{N}(X|\mu,\sigma) = \frac{1}{\sigma\sqrt{2\pi}}e^{-\frac{(x-\mu)^2}{2\sigma^2}}$$

where μ is the mean and σ^2 is the variance.

The below image has a few Gaussian distributions with a difference in mean (μ) and variance (σ 2).



But this would only be true for a single variable. In the case of two variables, instead of a 2D bell-shaped curve, we will have a 3D bell curve as shown below:



The probability density function would be given by:

$$\mathcal{N}(X|\mu, \Sigma) = \frac{1}{\sqrt{(2\pi)|\Sigma|}} \exp\left(-\frac{1}{2}(X-\mu)^T \Sigma^{-1}(X-\mu)\right)$$

where X is the input vector, μ is the 2D mean vector, and Σ is the 2×2 covariance matrix. The covariance would now define the shape of this curve. We can generalize the same for d-dimensions.

Thus, this multivariate Gaussian model would have X and μ as vectors of length d, and Σ would be a $d \times d$ covariance matrix.

1.4 Gaussian Mixture Models

Suppose there are K clusters (For the sake of simplicity here it is assumed that the number of clusters is known and it is K). So μ and Σ are also estimated for each k. Had it been only one distribution, they would have been estimated by the **maximum-likelihood method**. But since there are K such clusters and the probability density is defined as a linear function of densities of all these K distributions, i.e.

$$p(\mathbf{x}) = \sum_{k=1}^{K} \pi_k \mathcal{N}(\mathbf{x} | \mu_k, \Sigma_k) \Rightarrow p(x_1, x_2, \dots, x_N) = \sum_{k=1}^{K} \pi_k \mathcal{N}(x_1, x_2, \dots, x_N | \mu_k, \Sigma_k)$$

$$\begin{cases} \pi : \text{mixing coefficient} \\ \boldsymbol{\mu} : \text{means} \\ \boldsymbol{\Sigma} : \text{covariance matrix} \end{cases}$$

where π_k is the mixing coefficient for k-th distribution.

Assuming that data points are independent, for estimating the parameters by the maximum log-likelihood method, compute $p(\mathbf{x}|\mu, \Sigma, \pi)$.

$$ln \ p(\mathbf{x}|\mu, \Sigma, \pi) = \sum_{n=1}^{N} p(x_n) = \sum_{n=1}^{N} ln \sum_{k=1}^{K} \pi_k \mathcal{N}(x_n|\mu_k, \Sigma_k)$$

Now define a random variable $\gamma_k(x_n)$, such that $\gamma_k(x_n) = p(k|x_n)$. From Bayes' theorem,

$$\gamma_k(x_n) = \frac{p(x_n|k)p(k)}{\sum_{k=1}^{K} p(k)p(x_n|k)} = \frac{p(x_n|k)\pi_k}{\sum_{k=1}^{K} \pi_k p(x_n|k)} = \frac{\pi_k \mathcal{N}(x_n|\mu_k, \Sigma_k)}{\sum_{k=1}^{K} \pi_k \mathcal{N}(x_n|\mu_k, \Sigma_k)}$$

Now for the log-likelihood function to be maximum, its derivative of $p(x_n|\mu, \Sigma, \pi)$ with respect to μ , Σ and π should be zero. So equating the derivative of $p(x_n|\mu, \Sigma, \pi)$ to zero and rearranging the terms,

$$\mu_k = \frac{\sum_{n=1}^N \gamma_k(x_n) x_n}{\sum_{n=1}^N \gamma_k(x_n)}$$

Similarly taking derivative with respect to Σ and pi respectively, one can obtain the following expressions.

$$\Sigma_{k} = \frac{\sum_{n=1}^{N} \gamma_{k}(x_{n})(x_{n} - \mu_{k})(x_{n} - \mu_{k})^{T}}{\sum_{n=1}^{N} \gamma_{k}(x_{n})}$$

And

$$\pi_k = \frac{1}{N} \sum_{n=1}^N \gamma_k(x_n)$$

Note: $\sum_{n=1}^{N} \gamma_k(x_n)$ denotes the total number of sample points in the k-th cluster. Here it is assumed that there is a total N number of samples and each sample containing d features is denoted by x_i . So it can be clearly seen that the parameters cannot be estimated in closed form. This is where the **Expectation-Maximization algorithm** is beneficial.

1.5 Expectation-Maximization (EM) Algorithm

The Expectation-Maximization (EM) algorithm is an iterative way to find maximum-likelihood estimates for model parameters when the data is incomplete or has some missing data points or has some hidden variables. EM chooses some random values for the missing data points and estimates a new set of data. These new values are then recursively used to estimate a better first date, by filling up missing points, until the values get fixed. These are the two basic steps of the EM algorithm, namely **E Step or Expectation Step or Estimation Step** and **M Step or Maximization Step**.

- Estimation step (E step):
 - initialize μ_k , Σ_k and π_k by some random values, or by K means clustering results or by hierarchical clustering results.
 - Then for those given parameter values, estimate the value of the latent variables (i.e $\gamma_k(x_n)$)

$$\gamma_k(x_n) = \frac{\pi_k \mathcal{N}(x_n | \mu_k, \Sigma_k)}{\sum_{k=1}^K \pi_k \mathcal{N}(x_i | \mu_k, \Sigma_k)}$$

- Maximization Step(M step):
 - Update the value of the parameters(i.e. μ_k , Σ_k and π_k) calculated using ML method.

$$\mu_{k} = \frac{\sum_{n=1}^{N} \gamma_{k}(x_{n})x_{n}}{\sum_{n=1}^{N} \gamma_{k}(x_{n})}$$

$$\Sigma_{k} = \frac{\sum_{n=1}^{N} \gamma_{k}(x_{n})(x_{n} - \mu_{k})(x_{n} - \mu_{k})^{T}}{\sum_{n=1}^{N} \gamma_{k}(x_{n})}$$

$$\pi_{k} = \frac{1}{N} \sum_{n=1}^{N} \gamma_{k}(x_{n})$$

2 LAB Assignment

Please finish the **Exercise** and answer **Questions**.

2.1 Exercise (100 Points)

In this lab, our goal is to write a program to segment different objects using the **GMM and EM** algorithm. We also use <u>k-means clustering algorithm to initialize the parameters</u> of GMM. The following steps should be implemented to achieve such a goal:

- 1. Load image
- 2. Initialize parameters of GMM using K-means
- 3. Implement the EM algorithm for GMM
- 4. Display result

2.1.1 Import some libraries

```
In [1]:
```

```
1
   # Dependency
 2 import numpy as np
 3 from scipy.stats import multivariate_normal
   from sklearn.cluster import KMeans
   import tqdm
 5
 7
   from PIL import Image
 8
 9
   COLORS = [
10
        (255, 0, 0),
                       # red
11
        (0, 255, 0), # green
12
        (0, 0, 255),
                      # blue
13
        (255, 255, 0), # yellow
        (255, 0, 255), # magenta
14
    1
15
16
17
```

2.1.2 Load Image

What you should do is to implement Z-score normalization in <code>load()</code>:

```
In [ ]:
```

```
import cv2
    def load(image path):
 2
 3
        image = cv2.imread(image path)
 4
        h, w, c = image. shape
 5
 6
        # TODO: please normalize image pixl using Z-score
 7
        mean = None
        std = None
 8
 9
        image norm = None
10
11
12
        return h, w, c, image norm
```

k-means is used to initialize means, covariance matrices and mixing coefficients of GMM

In []:

```
def kmeans (n cluster, image pixl):
 1
 2
        kmeans = KMeans(n clusters=n cluster)# instantiate a K-means
 3
        labels = kmeans.fit_predict(image_pixl)# fit and get clustering result
 4
        initial_mus = kmeans.cluster_centers_# get centroids
 5
        initial_priors, initial_covs = [], []
        #Followings are for initialization:
 7
        for i in range (n cluster):
            datas = image_pix1[labels == i, ...].T
 8
 9
            initial covs. append (np. cov (datas))
            initial_priors.append(datas.shape[1] / len(labels))
10
11
        return initial_mus, initial_priors, initial_covs
12
```

2.1.4 Implement GMM algorithm

We use EM algorithm to refine GMM's parameters.

Although it may be not easy for some students to derive EM formula for GMM, GMM isn't very difficult to implement once you have the formula. Therefore, to help you understand GMM more, there are still some blanks for you to fill in.

$$E - step : \gamma(z_{nk}) = \frac{\pi_k \mathcal{N}(\mathbf{x}_n | \boldsymbol{\mu}_k, \boldsymbol{\Sigma}_k)}{\sum_{j=1}^K \pi_j \mathcal{N}(\mathbf{x}_n | \boldsymbol{\mu}_j, \boldsymbol{\Sigma}_j)}$$

$$M - step : \begin{cases} \boldsymbol{\mu}_k^{new} = \frac{1}{N_k} \sum_{n=1}^N \gamma(z_{nk}) \mathbf{x}_n \\ \boldsymbol{\Sigma}_k^{new} = \sum_{n=1}^N \gamma(z_{nk}) (\mathbf{x}_n - \boldsymbol{\mu}_k^{new}) (\mathbf{x}_n - \boldsymbol{\mu}_k^{new})^T \\ \pi_k^{new} = \frac{N_k}{N} \end{cases}$$

$$Loglikelihood: lnp(\mathbf{x}|\boldsymbol{\mu},\boldsymbol{\Sigma},\boldsymbol{\pi}) = \sum_{n=1}^{N} \ln\{\sum_{k=1}^{K} \pi_{k} \mathcal{N}(\mathbf{x}|\boldsymbol{\mu}_{k},\boldsymbol{\Sigma}_{k})\}$$

2.1.4.1 E-step

It is in inference().

In the following code, prob is $\pi_k \mathcal{N}(\mathbf{x}_n | \boldsymbol{\mu}_k, \boldsymbol{\Sigma}_k)$, gamma is γ . You need to implement log likelihood and γ .

```
def inference(self, datas):
    probs = []
    for i in range(self.ncomp):
        mu, cov, prior = self.mus[i, :], self.covs[i, :, :], self.priors[i]
        prob = prior * multivariate_normal.pdf(datas, mean=mu, cov=cov, allow_singular=Tr
ue)
    probs.append(np.expand_dims(prob, -1))
    preds = np.concatenate(probs, axis=1)

# TODO: calc log likelihood
    log_likelihood = None

# TODO: calc gamma
    gamma = None

return gamma, log_likelihood
```

2.1.4.2 M-step

It is in update()

You need to implement mean μ , covariance Σ and mixing coefficient π .

```
def update(self, datas, gamma):
    new_mus, new_covs, new_priors = [], [], []
    soft counts = np. sum(gamma, axis=0)
    for i in range (self. ncomp):
        # TODO: calc mu
        new mu = None
        new_mus. append (new_mu)
        # TODO: calc cov
        new cov = None
        new covs. append (new cov)
        # TODO: calc mixing coefficients
        new prior = None
        new_priors.append(new_prior)
    self.mus = np.asarray(new mus)
    self.covs = np.asarray(new_covs)
    self.priors = np. asarray(new priors)
```

2.1.5 Iteration

Iteration part is as you see in fit()

```
def fit(self, data, iteration):
    prev_log_liklihood = None

bar = tqdm.tqdm(total=iteration)
    for i in range(iteration):
        gamma, log_likelihood = self.inference(data)
        self.update(data, gamma)
        if prev_log_liklihood is not None and abs(log_likelihood - prev_log_liklihood) <
le-10:
        break
    prev_log_likelihood = log_likelihood

bar.update()
    bar.set_postfix({"log_likelihood": log_likelihood})</pre>
```

In []:

```
1
    class GMM:
 2
              <u>_init_</u>(self, ncomp, initial_mus, initial_covs, initial_priors):
        def
 3
 4
                                     the number of clusters
            :param ncomp:
 5
            :param initial mus:
                                     initial means
 6
            :param initial covs:
                                     initial covariance matrices
 7
            :param initial priors: initial mixing coefficients
 8
 9
            self.ncomp = ncomp
10
            self.mus = np.asarray(initial mus)
            self.covs = np.asarray(initial covs)
11
12
            self.priors = np. asarray(initial priors)
13
14
        def inference(self, datas):
15
16
            E-step
17
                             original data
            :param datas:
18
                             posterior probability (gamma) and log likelihood
            :return:
19
20
            probs = []
21
            for i in range(self.ncomp):
22
                mu, cov, prior = self.mus[i, :], self.covs[i, :, :], self.priors[i]
23
                prob = prior * multivariate normal.pdf(datas, mean=mu, cov=cov, allow singular=Tru
24
                probs. append (np. expand dims (prob, -1))
25
            preds = np. concatenate(probs, axis=1)
26
27
            # TODO: calc log likelihood
28
            log likelihood = None
29
30
            # TODO: calc gamma
31
            gamma = None
32
33
            return gamma, log_likelihood
34
35
        def update(self, datas, gamma):
36
37
            M-step
            :param datas:
38
                             original data
39
            :param gamma:
                              gamma
40
            :return:
41
            new mus, new covs, new priors = [], [], []
42
            soft counts = np. sum(gamma, axis=0)
43
44
            for i in range (self. ncomp):
                # TODO: calc mu
45
46
                new mu = None
                new mus. append (new mu)
47
48
                # TODO: calc cov
49
50
                new cov = None
51
                new covs. append (new cov)
52
53
                # TODO: calc mixing coefficients
54
                new prior = None
55
                new_priors.append(new_prior)
56
57
            self.mus = np.asarray(new_mus)
58
            self.covs = np. asarray(new covs)
            self.priors = np. asarray (new priors)
```

```
60
61
        def fit(self, data, iteration):
62
            prev log liklihood = None
63
64
            bar = tqdm. tqdm(total=iteration)
            for i in range (iteration):
65
                gamma, log_likelihood = self.inference(data)
66
67
                self.update(data, gamma)
                if prev log liklihood is not None and abs(log likelihood - prev log liklihood) <
68
69
70
                prev_log_likelihood = log_likelihood
71
72
                bar. update()
73
                bar.set_postfix({"log likelihood": log_likelihood})
```

2.1.6 Display

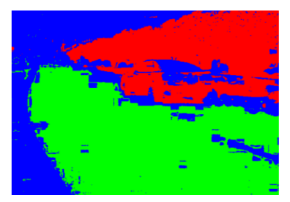
We use matplotlib to display what we segment, you can check the code in visualize()

In []:

```
from PIL import Image
 2
    import matplotlib.pyplot as plt
 3
 4
 5
   def visualize (gmm, image, ncomp, ih, iw):
 6
        beliefs, log_likelihood = gmm.inference(image)
 7
        map_beliefs = np. reshape(beliefs, (ih, iw, ncomp))
        segmented_map = np.zeros((ih, iw, 3))
 8
 9
        for i in range(ih):
10
            for j in range(iw):
                hard_belief = np.argmax(map_beliefs[i, j, :])
11
12
                segmented_map[i, j, :] = np.asarray(COLORS[hard_belief]) / 255.0
        plt.imshow(segmented_map)
13
        plt.show()
14
```

2.1.7 sample Result





2.2 Questions(3 points)

- 1. What are the strengths of GMM; when does it perform well?
- 2. What are the weaknesses of GMM; when does it perform poorly?
- 3. What makes GMM a good candidate for the clustering problem, if you have enough knowledge about the data?