SML: Exercise 2

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Summer Term 2020 Sheet 1

Task 1.1: Bayesian Decision Theory

In this exercise, we consider data generated by a mixture of two Gaussian distributions with parameters $\{\mu_1, \sigma_1\}$ and $\{\mu_2, \sigma_2\}$. Each Gaussian represents a class labeled C1 and C2, respectively.

1.1a)

The bayesian decision theory finds the a-posteriori probability (posterior) of a class C_k given a observation (feature) x by using the Bayes Theorem

$$p(C_k|x) = \frac{p(x|C_k)p(C_k)}{p(x)}.$$

The goal of bayesian decision theory is to minimize the misclassification rate. Let C_1, C_2 be two cases. The condition which holds at the optimal decision boundary is

$$p(C_1|x) = p(C_2|x).$$

We decide C_1 if

$$p(C_1|x) > p(C_2|x).$$

1.1b)

Let both classes C_1 and C_2 have equal probabilities $p(C_1) = p(C_2)$ and the same variance σ_1, σ_2 . In the following we will derive the decision boundary x^* analytically as a function of two means μ_1, μ_2 .

$$p(C_1|x) = p(C_2|x)$$

$$\frac{p(x|C_1)p(C_1)}{p(x)} = \frac{p(x|C_2)p(C_2)}{p(x)} \quad | \div p(C_1) \quad | \cdot p(x)$$

$$p(x|C_1) = p(x|C_2)$$

$$\frac{1}{\sqrt{2\pi}\sigma_1} \exp\left(-\frac{(x-\mu_1)^2}{2\sigma_1^2}\right) = \frac{1}{\sqrt{2\pi}\sigma_2} \exp\left(-\frac{(x-\mu_2)^2}{2\sigma_2^2}\right) \quad | \div \frac{1}{\sqrt{2\pi}\sigma_1} \quad | \ln()$$

$$-\frac{(x-\mu_1)^2}{2\sigma_1^2} = -\frac{(x-\mu_2)^2}{2\sigma_2^2} \quad | \cdot 2\sigma_1^2$$

$$(x-\mu_1)^2 = (x-\mu_2)^2$$

$$x^2 - 2x\mu_1 + \mu_1^2 = x^2 - 2x\mu_2 + \mu_2^2$$

$$\mu_1^2 - \mu_2^2 = (-2\mu_2 + 2\mu_1)x$$

$$x = \frac{\mu_1^2 - \mu_2^2}{-2\mu_2 + 2\mu_1}$$

$$x = \frac{\mu_1 + \mu_2}{2}$$

1.1c)

Let $\mu_1 < 0$, $\mu_1 = 2\mu_2$, $\sigma_1 = \sigma_2$ and $p(C_1) = p(C_2)$. The misclassification of a sample $x \in C_2$ as a class C_1 should be four times more expensive than the opposite. Clearly our boundary will change in such a way, that we classify x more often as in class C_2 then in class C_1 . In the following we will derive the boundary analytically.

$$R(\alpha_1|x) = 4p(C_2|x)$$
$$R(\alpha_2|x) = p(C_1|x)$$

We will get the decision boundary by looking at

$$R(\alpha_2|x) = R(\alpha_1|x)$$

$$p(C_1|x) = 4p(C_2|x) \text{ same computations as in 1.1b)}$$

$$p(x|C_1) = p(x|C_2)$$

$$\frac{1}{\sqrt{2\pi}\sigma_1} \exp\left(-\frac{(x-\mu_1)^2}{2\sigma_1^2}\right) = \frac{4}{\sqrt{2\pi}\sigma_2} \exp\left(-\frac{(x-\mu_2)^2}{2\sigma_2^2}\right) \quad | \div \frac{1}{\sqrt{2\pi}\sigma_1} \quad | \ln()$$

$$-\frac{(x-\mu_1)^2}{2\sigma_1^2} = \ln(4) - \frac{(x-\mu_2)^2}{2\sigma_2^2} \quad | \cdot 2\sigma_1^2$$

$$-(x-\mu_1)^2 = 2\sigma_1^2 \ln(4) - (x-\mu_2)^2$$

$$-x^2 + 2\mu_1 x - \mu_1^2 = 2\sigma_1^2 \ln(4) - x^2 + 2x\mu_2 - \mu_2^2$$

$$2\mu_1 x - 2\mu_2 x = 2\sigma_1^2 \ln(4) - \mu_2^2 - \mu_1^2$$

$$x = \frac{2\sigma_1^2 \ln(4) - \mu_2^2 + \mu_1^2}{\mu_1}$$

$$x = \frac{2\sigma_1^2 \ln(4) + 3\mu_2^2}{\mu_1}$$

Task 1.2: Density Estimation

We are given data C1 and C2, which we suppose to be generated by 2D-Gaussians with parameters μ_1, Σ_1 and μ_2, Σ_2 , respectively.

1.2a)

Assume we are given iid. datapoints x_i , i = 1, ..., n which are generated by a 2D-Gaussian. Following the max-likelihood principle, we maximize the log-likelihood function

$$l(\mu, \Sigma, x_1, ..., x_n) = \ln(\prod_{i=1}^n p(x_i | \mu, \Sigma)) = \sum_{i=1}^n \ln(p(x_i | \mu, \Sigma))$$

for the Gaussian probability density

$$p(x|\mu, \Sigma) = \frac{1}{\sqrt{(2\pi)^2 |\Sigma|}} \exp\left(-\frac{1}{2}(x-\mu)^T \Sigma^{-1} (x-\mu)\right) . \tag{1}$$

We receive

$$l(\mu, \Sigma) := l(\mu, \Sigma, x_1, ..., x_n) = \sum_{i=1}^{n} \left(-\ln(2\pi) - \frac{1}{2}ln(|\Sigma|) - \frac{1}{2}(x_i - \mu)^T \Sigma^{-1}(x_i - \mu) \right)$$
 (2)

$$= -n\ln(2\pi) - \frac{n}{2}ln(|\Sigma|) - \frac{1}{2}\sum_{i=1}^{n}(x_i - \mu)^T \Sigma^{-1}(x_i - \mu) .$$
 (3)

We compute the derivatives w.r.t. μ and Σ and set them equal to zero. This yields

$$\frac{d}{d\mu}l(\mu, \Sigma, x_1, ..., x_n) = \frac{d}{d\mu} - \frac{1}{2} \sum_{i=1}^n (x_i - \mu)^T \Sigma^{-1} (x_i - \mu)$$
$$= -\sum_{i=1}^n \frac{d}{d\mu} \frac{1}{2} (x_i - \mu)^T \Sigma^{-1} (x_i - \mu) .$$

Using the matrix identity $\frac{d}{dw} \frac{w^T A w}{dw} = 2Aw$ which holds if w does not depend on A and if A is symmetric, we get (with $w = (x - \mu), dw = -d\mu$)

$$0 \stackrel{!}{=} \frac{d}{d\mu} l(\mu, \Sigma, x_1, ..., x_n)$$
$$0 \stackrel{!}{=} -\sum_{i=1}^{n} \Sigma^{-1}(x_i - \mu) .$$

Finally, we use that Σ^{-1} is positive definite, so we can leave it out here and get

$$0 \stackrel{!}{=} n\mu - \sum_{i=1}^{n} x_i ,$$

which is solved for the MLE-estimate

$$\hat{\mu} = \frac{1}{n} \sum_{i=1}^{n} x_i \ . \tag{4}$$

Secondly, we need to compute the derivative w.r.t Σ . To do that, we will need some results from mathematical classes. The following is used without prove:

• Cyclic permutations of a matrix product do not change the trace of it:

$$tr[ABC] = tr[CAB]$$

• The trace of a scalar is the scalar itself. In particular: the result of a quadratic form $x^T A x$ is a scalar, such that:

$$x^T A x = tr \left[x^T A x \right] = tr \left[x^T x A \right]$$

- $\frac{d}{dA}tr[AB] = B^T$
- $\frac{d}{dA} \ln |A| = A^{-T}$

As a first result of these assumptions, we can show, that

$$\frac{d}{dA}x^TAx = \frac{d}{dA}tr\left[x^TxA\right] = \left[xx^T\right]^T = xx^T.$$

We now got the tools to re-write the log-likelihood function in (3) to

$$l(\mu, \Sigma) = -n \ln(2\pi) - \frac{n}{2} ln(|\Sigma|) - \frac{1}{2} \sum_{i=1}^{n} (x_i - \mu)^T \Sigma^{-1} (x_i - \mu)$$
$$= C + \frac{n}{2} ln(|\Sigma^{-1}|) - \frac{1}{2} \sum_{i=1}^{n} tr \left[(x_i - \mu)(x_i - \mu)^T \Sigma^{-1} \right]$$
$$= C + \frac{n}{2} ln(|\Sigma^{-1}|) - \frac{1}{2} \sum_{i=1}^{n} tr \left[\Sigma^{-1} (x_i - \mu)(x_i - \mu)^T \right]$$

for a constant C. Where in the last step we used that AB = BA for symmetric matrices A, B. Taking the derivative w.r.t Σ^{-1} yields

$$\frac{d}{d\Sigma^{-1}}l(\mu,\Sigma) = \frac{n}{2}\Sigma^{T} - \frac{1}{2}\sum_{i=1}^{n}(x_{i} - \mu)(x_{i} - \mu)^{T}$$

and plugging in $\hat{\mu}$ as an estimation of μ and setting equal to zero finally gives us

$$0 \stackrel{!}{=} \frac{d}{d\Sigma^{-1}} l(\hat{\mu}, \Sigma)$$
$$0 \stackrel{!}{=} \frac{n}{2} \Sigma^{T} - \frac{1}{2} \sum_{i=1}^{n} (x_i - \hat{\mu})(x_i - \hat{\mu})^{T}$$

which is solved for the (biased) MLE estimate

$$\tilde{\Sigma} = \frac{1}{n} \sum_{i=1}^{n} (x_i - \hat{\mu})(x_i - \hat{\mu})^T$$
(5)

1.2b)

We compute the prior probabilities of C1 and C2, using the following python code. We read the number of data points in each class and divide it by the sum of total data points in both classes.

import numpy as np

Calling

```
lengths = get_lengths()
print(get_priors(lengths[0],lengths[1]))
```

we get the following results for the prior probabilities: p(C1) = 0.239 and p(C2) = 0.761.

1.2c)

Having a data set X and an estimator $\hat{\theta}$ on the true parameter θ , we define the bias of an estimator as the expected deviation from the true parameter. We get the formula

$$bias(\hat{\theta}) = \mathbb{E}_X \left[\hat{\theta}(X) - \theta \right]$$

We call an estimator unbiased iff $bias(\hat{\theta}) = 0$ and biased otherwise.

From the lecture we know that the MLE of the mean of a Gaussian is unbiased, but the MLE of the variance of a Gaussian is biased. In fact, an unbiased estimator on the variance would be the sample covariance matrix

$$\hat{\Sigma} = \frac{1}{n-1} \sum_{i=1}^{n} (x_i - \hat{\mu})(x_i - \hat{\mu})^T .$$
(6)

To calculate the conditional distribution densities $p(x|C_i)$ we need to estimate the underlying parameters μ_i and Σ_i . For both classes we use the MLE-estimate of the mean, which is unbiased and calculated like in (4). We compute the biased MLE-estimate $\tilde{\Sigma}$ for the variance via (5) and the unbiased estimate $\hat{\Sigma}$ via (6). We wrote the following python code

```
def extract_data(t):
        a=np.empty((0,2),float)
        for line in t:
                v=np.array([[line.split()[0],line.split()[1]]],float)
                a=np.append(a, v, axis=0)
        return a
def get mean estimation(points):
        m=np.zeros(2)
        m[0]=sum(points[:,0])
        m[1] = sum(points[:,1])
        m=m/len (points)
        return m
def get_biased_var_estimation(mean_est, points):
        l=points-mean est
        s=np.zeros((2,2))
        for line in 1:
                 s=s+np.outer(line, line)
        s=s/len(1)
```

```
def get unbiased var estimation (mean est, points):
                                    l=points-mean est
                                    s=np.zeros((2,2))
                                   for line in 1:
                                               s=s+np.outer(line, line)
                                    s=s/(len(1)-1)
                        return s
                        def print results(link):
                                   a=extract data(open(link))
                                   mean=get_mean_estimation(a)
                                    print("mean=",mean)
                                   sigma=get_unbiased_var_estimation(mean,a)
                                    print("biased_Sigma=",get_biased_var_estimation(mean,a))
                                    print("sigma=",sigma)
which gives us the results for class C1
                        print results(link1)
                       mean= [-0.70681374 -0.81343083]
                        biased Sigma= [[9.01952586 2.67287085]
                                               [2.67287085 3.59633965]]
                        sigma= [[9.05742302 2.6841014 ]
                                   [2.6841014 3.61145033]]
and for class C2
                        print_results(link2)
                       mean= [3.98534252 3.98438364]
                        biased_Sigma= [[4.1753815  0.02758324]
                                    [0.02758324 2.75296323]]
                        sigma= [[4.18087542 0.02761954]
                                   [0.02761954 2.75658555]]
Transfered to the notation we introduced earlier, we get \hat{\mu_1} = \begin{pmatrix} -0.71 \\ -0.81 \end{pmatrix} with \tilde{\Sigma_1} = \begin{pmatrix} 9.02 & 2.67 \\ 2.67 & 3.6 \end{pmatrix} or with the unbiased
\hat{\Sigma_1} = \begin{pmatrix} 9.06 & 2.68 \\ 2.68 & 3.61 \end{pmatrix} for C1.
And \hat{\mu_2} = \begin{pmatrix} 3.99 \\ 3.98 \end{pmatrix} with \tilde{\Sigma_2} = \begin{pmatrix} 4.18 & 0.03 \\ 0.03 & 2.75 \end{pmatrix} or with the unbiased \hat{\Sigma_2} = \begin{pmatrix} 4.18 & 0.03 \\ 0.03 & 2.76 \end{pmatrix} for C2.
We can now just plug in into formula (1) to get
for the biased estimate p(x|C_i) = p(x|\hat{\mu_i}, \Sigma_i)
and for the unbiased p(x|C_i) = p(x|\hat{\mu}_i, \Sigma_i), where the right hand sides are given like in (1).
```

return s

1.2d)

Using the unbiased estimates $p(x|C_i) = p(x|\hat{\mu_i}, \hat{\Sigma_i})$ from last task, for each class we plot the Gaussian in a single graph with the data points: Therefore we added 2 new functions to our code:

```
def gaussian(x,mu, detsigma, sigmainv):
    z=np.array(x-mu)
    enum=np.exp(-0.5*np.dot(np.dot(z, sigmainv), np.transpose(z)))
```

```
denom=np.sqrt(np.power(2*np.pi,len(x))*detsigma)
        return enum/denom
def plot(link):
        a=extract_data(open(link))
        mean=get_mean_estimation(a)
        print("mean=",mean)
        sigma=get_unbiased_var_estimation(mean,a)
        print("biased_Sigma=",get_biased_var_estimation(mean,a))
        print("sigma=",sigma)
        detsigma=np.linalg.det(sigma)
        sigmainv=np.linalg.inv(sigma)
        x=np. linspace(min(a[:,0]), max(a[:,0]), 300)
        y=np.linspace(min(a[:,1]),max(a[:,1]),300)
        X, Y=np.meshgrid(x,y)
        Z=np.empty_like(X)
        i=0
        while i < len(X):
                j=0
                while j < len(Y):
                         xy=np.array([X[i,j],Y[i,j]])
                         ergb=gaussian(xy,mean,detsigma,sigmainv)
                         Z[i,j]=ergb
                         j=j+1
                i=i+1
        plt.contourf(X,Y,Z,25)
        plt.colorbar()
        plt.scatter(a[:,0],a[:,1],alpha=1,c="white",s=0.8)
        return
```

Calling "plot(link1)" gives us Figure 1 and "plot(link2)" gives Figure 2.

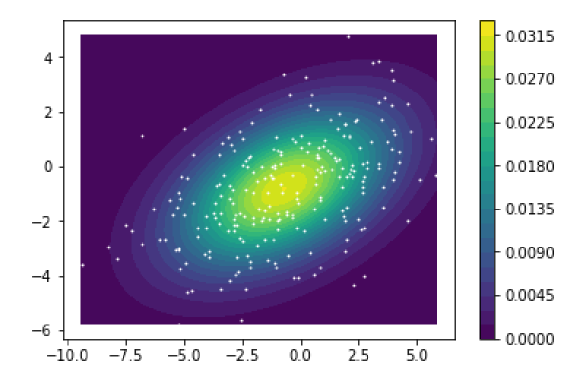


Figure 1: Gaussian $(\hat{\mu_1}, \hat{\Sigma_1})$ and data points C1 in white

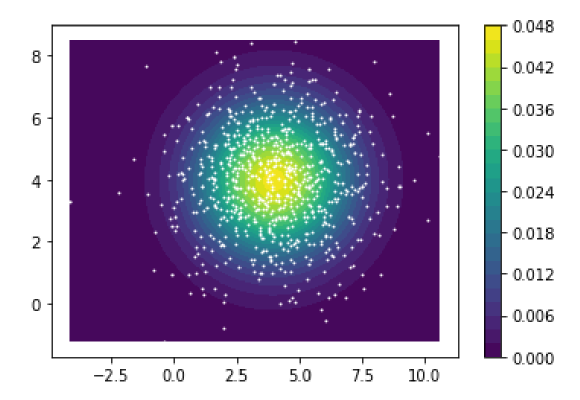


Figure 2: Gaussian $(\hat{\mu_2},\hat{\Sigma_2})$ and data points C2 in white

1.2e)

We are interested in $p(C_i|x)$. From Bayes' rule we know that this equals

$$p(C_i|x) = \frac{p(x|C_i)p(C_i)}{\sum_j p(x|C_j)p(C_j)}$$

which helps us a lot because we calculated all terms occuring in the right hand side. We add two more functions to our code:

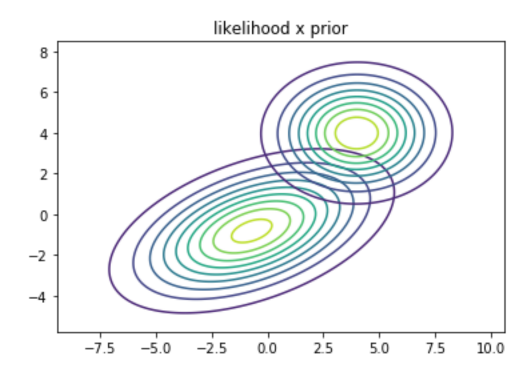
```
def postplot(link):
        g=extract_data(open(link3))
        a=extract_data(open(link))
        prior=len(a)/len(g)
        mean=get_mean_estimation(a)
        sigma=get_unbiased_var_estimation(mean,a)
        detsigma=np.linalg.det(sigma)
        sigmainv=np.linalg.inv(sigma)
        x=np. linspace(min(g[:,0]), max(g[:,0]), 300)
        y=np.linspace (\min(g[:,1]), \max(g[:,1]), 300)
        X, Y=np. meshgrid (x, y, sparse=False)
        Z=np.empty like(X)
        i = 0
        while i < len(X):
                 i=0
                 while j < len(Y):
                         xy=np.array([X[i,j],Y[i,j]])
                         ergb=prior * gaussian (xy, mean, detsigma, sigmainv)
                         Z[i,j]=ergb
                         j=j+1
                 i=i+1
        return [X,Y,Z,prior]
def actual_plotting():
        X,Y,Z1,p1=postplot(link1)
        Z2, p2=postplot(link2)[2:4]
        S=np.add(Z1,Z2)
        Z1neu=np.divide(Z1,S)
        Z2neu=np. divide (Z2,S)
        plt.contour(X,Y,Z1,10)
        plt.contour(X,Y,Z2,10)
        plt.title("likelihood_x_prior")
        plt.figure()
        plt.contour(X,Y,Z1neu,10)
        plt.colorbar()
        plt.title("p(C1|x)")
        plt.figure()
        plt.contour(X,Y,Z2neu,10)
        plt.colorbar()
        plt.title("p(C2|x)")
        plt.figure()
        dec=np.greater(Z1neu, Z2neu)
        plt.scatter(X,Y,dec)
        plt.title("blue=decideC1,_white=decideC2")
        mpoints=np.empty(shape=[0,2])
```

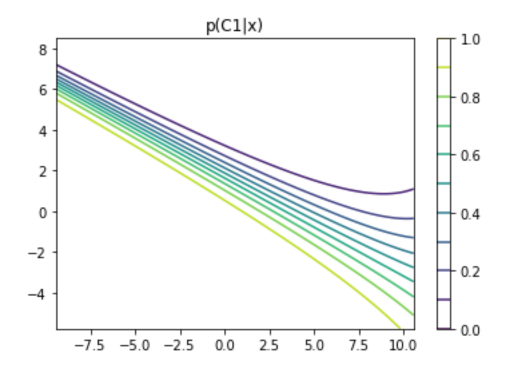
```
while j < len(X):
        i = 0
        while i < len(Y):
                 if not dec[i,j]:
                         xy=np.array([[X[i,j],Y[i,j]]])
                         mpoints=np.append(mpoints, xy, axis=0)
                         break
                 i=i+1
        j=j+1
plt.figure()
plt.scatter(mpoints[:,0], mpoints[:,1], s = 0.5)
plt.title("decision_boundary")
plt.figure()
plt.contour(X,Y,Z1,10)
plt.contour(X,Y,Z2,10)
plt.scatter(mpoints[:,0],mpoints[:,1],s=0.5)
plt.title("likelihood_x_prior_and_decbound")
```

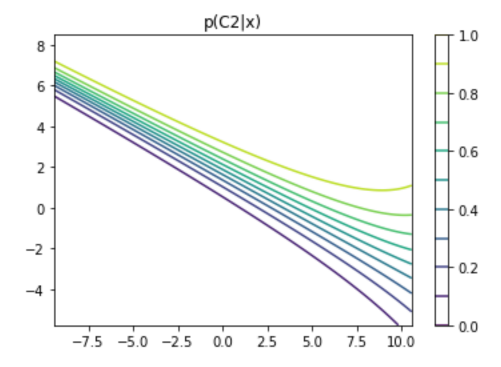
return

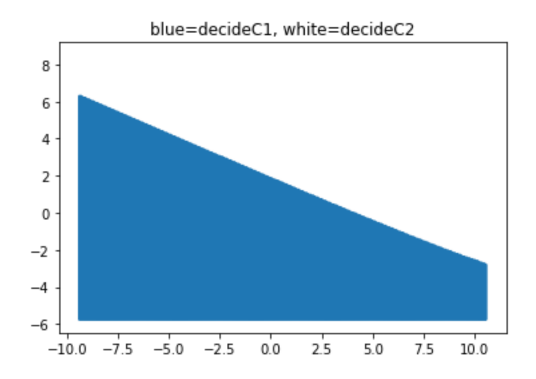
Where in link3 we stored a link to a .txt file containing all data from C1 and C2 combined. This can be generated by executing

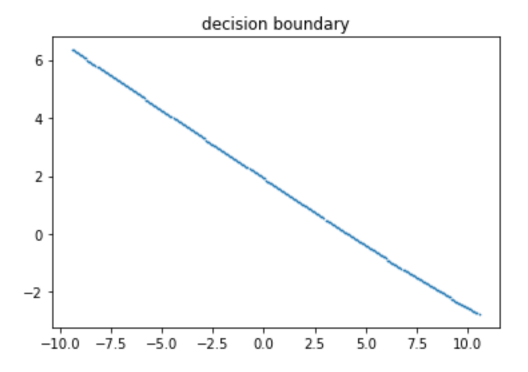
once, or just do it by hand. Calling actual_plotting() then gives us the following results:

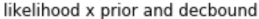


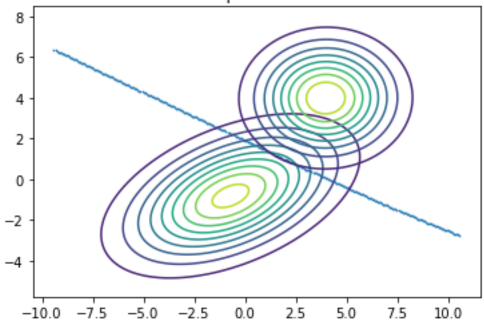










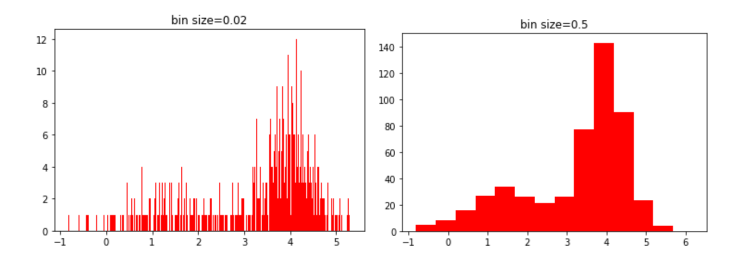


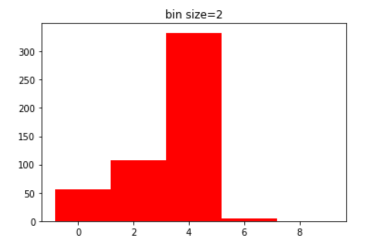
Task 1.3: Non-parametric Density Estimation

1.3a)

Histogram Given training data, we plot histograms with different binsizes. We therefore wrote a python function that plots a histogram given data and a binsize:

Calling the function for different binsizes with our training data gives us:





Intuitively, we suggest that the best bin-size from the three would be 0.5. Obviously, 0.02 is too small. Fixing a bin with that bin size, we got a very low probability of having even a single sample in the bin. We are very reliant on the randomly chosen data points, and therefore get a histogram where neighbored bins vary a lot. On the other hand, we see that a bin size of 2.0 gives us a result that is too smooth. We lose information of the shape of the underlying density by generalizing too much (we put too many, almost all, data points in one bowl, leaving no space for local changes in the density function). For bin size 0.5 we seem to be about right, because we see enough structure to guess the underlying distribution, and neighbored bins still seem related to another which emphasizes our idea of an underlying continuous density. This is referred to as the Bias-Variance Problem.

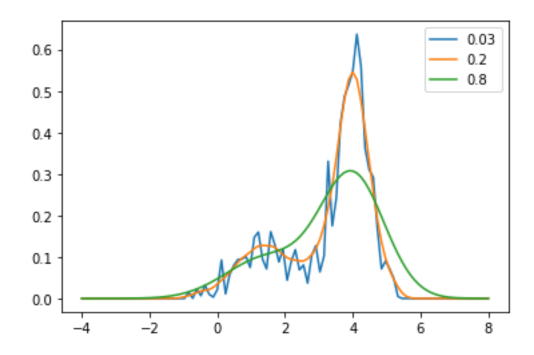
1.3b)

We compute the probability density estimates for Gaussian kernels with three different variances. Also we compute the log-likelihood of these estimates. Therefore we implement

```
def gaussian_kernel(data, x, sigma):
    result = 0
    for i in range(len(data)):
        result += m.exp(-(x-data[i])**2 / (2 * sigma**2))
    result = (result/(len(data) * m.sqrt(2 * m.pi * sigma**2)))
    return result
```

```
def log likehood gaussian(data, log like data, sigma):
         result = 0
         for i in range(len(log_like_data)):
                  result += np.log(gaussian kernel(data, log like data[i], sigma))
         return result
and calling
         # # settings
         data = nonParamTrain
         # plot densities
         x = np.linspace(-4,8,100)
         y = []
         for sigma in [0.03, 0.2, 0.8]:
                  for i in range(len(x)):
                           y.append(gaussian_kernel(data, x[i], sigma))
                  plt.plot(x,y,label=str(sigma))
                  plt.legend()
                  y = []
         plt.show()
         # log-likehood
         for sigma in [0.03, 0.2, 0.8]:
                  log_likehood_solution = log_likehood_gaussian(data, data, sigma)
print("sigma:", sigma, "log_likehood:", log_likehood_solution)
```

we get the following plot of the densities, labeled with the sigma value of the Gaussian kernel:



and the following results for the log likelihoods:

sigma: 0.03 log_likehood: -674.7279387090639 sigma: 0.2 log_likehood: -717.0216577444168 sigma: 0.8 log_likehood: -795.6632833459039

Interpretation: the bigger the log-likelihood number, the higher the probability that our data actually has the underlying density. Only taking this into account, we should decide on sigma=0.03. One could argue though, that the density for sigma=0.03 is not very smooth. So if we want to take this 'smoothness' into account without losing too much likelihood, we could select sigma=0.2 as a fair trade-off.

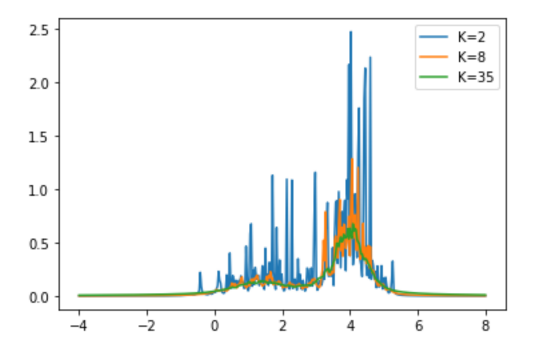
1.3c)

We estimate the probability density with the K-nearest neighbors method with K=2,8,35. In the plot we see that the bigger K gets, the smoother our estimate gets. This is, once again, because for big K we sum up a big amount of data points and therefore we 'lose' information on local peaks. We wrote the function

```
def k_nearest_neighbor(data,K,x):
    values = np.column_stack((data, data))
    values[:,0] = abs(values[:,0] - x)
    values = values[values[:,0].argsort()]
    result = values[K-1,0]
    v=2*result
    return (K / (v * len(data)))
```

and called with

we get the following graphs:



1.3d)

We compute the log-likelihood of testing data and our estimators (estimated with the training data).

method	log-likeli. on training	log-likeli. on testing
gaussian 0.03	-675	-inf
gaussian 0.2	-717	-2904
gaussian 0.8	-796	-3189
2-NN	-59	-2299
8-NN	-627	-2708
35-NN	-694	-2786

The result is pretty clear. kNN with K=2 gave the best results. In general, the kNN methods outperformed the Gaussians. Another mentionable result is the density recieved by 8-NN because it is way more smooth than the 2-NN but this is, at the cost of likelihood on both training and testing data.

This data was computed by calling

```
for sigma in [0.03, 0.2, 0.8]:
    log_likehood_solution = log_likehood_gaussian(data, nonParamTest , sigma)
    print("sigma:", sigma, "log_likehood_gaussian:", log_likehood_solution)

for K in [2, 8, 35]:
    log_likehood_solution = log_likehood_neighbor(data, nonParamTest, K)
    print("K:", K, "log_likehood_neighbor:", log_likehood_solution)

for K in [2, 8, 35]:
    log_likehood_solution = log_likehood_neighbor(data, data, K)
    print("K:", K, "log_likehood_neighbor training:", log_likehood_solution)
```

in our python file, where we only had to add one function to compute the log likelihood of a knn:

```
def log_likehood_neighbor(data, log_like_data, K):
    result = 0
    for i in range(len(log_like_data)):
        result += m.log(k_nearest_neighbor(data, K, log_like_data[i]))
    return result
```

Task 1.4: EM Algorithm

1.4a)

We assume a mixture of N=4 Gaussians. They all have to be modeled by their parameters $\{\mu_i, \Sigma_i\}$, i=1,..,4 and with an additional π_i for the estimated probability that Gaussian number i occurs. These parameters are updated in every iteration. To do this, we first do an 'E-step', where we calculate the posterior distribution for each Gaussian and for all data points. We store them in an array α with:

$$\alpha_{nj} = p(j|x_n) = \frac{\pi_j \mathcal{N}(x_n|\mu_j, \Sigma_j)}{\sum_{i=1}^{M} \pi_i \mathcal{N}(x_n|\mu_i, \Sigma_i)}$$

Then, in the next step, called 'M-step', we calculate:

$$N_{j} = \sum_{n=1}^{N} \alpha_{nj}$$

$$\mu_{j}^{new} = \frac{1}{N_{j}} \sum_{n=1}^{N} \alpha_{nj} x_{n}$$

$$\Sigma_{j}^{new} = \frac{1}{N_{j}} \sum_{n=1}^{N} \alpha_{nj} \left(x_{n} - \mu_{j}^{new} \right)^{2}$$

$$\pi_{j}^{new} = \frac{N_{j}}{N}$$

We implemented the following functions:

```
#imports
import numpy as np
import time
from matplotlib import pyplot as plt
#data
gmm data points = np.loadtxt("dataSets/gmm.txt")
N = len(gmm data points)
#initialization
gauss_1 = [np.array([0,0]), np.matrix([[1,0], [0,1]]), 0.25] # mu, sigma, pi
gauss 2 = [np.array([1,0]), np.matrix([[1,0], [0,1]]), 0.25]
gauss 3 = [np.array([2,0]), np.matrix([[1,0], [0,1]]), 0.25]
gauss_4 = [np.array([3,0]), np.matrix([[1,0], [0,1]]), 0.25]
gauss 1 = [gauss 1, gauss 2, gauss 3, gauss 4]
def log_likehood_gaussian(data, gauss):
        result = 0
        for i in range(len(data)):
                result += np.log(gauss[0][2]*gaussian_density_function(data[i], gauss
        return result
# evaluate gaussian function
def gaussian_density_function(x, mu, sigma):
        x = np.array(x)
        c1 = 1 / np. sqrt((2*np.pi)**2 * np. linalg. det(sigma))
        c2 = np.exp(-0.5 * np.matmul((x - mu), np.matmul(np.linalg.inv(sigma),(x - mu))
        y = c1 * c2
        return y.item()
#e step
def e_step(gauss):
        alpha = np.zeros((N, 4))
        for j in range(4):
                for i in range(N):
                         alpha[i,j] = gauss[j][2] * gaussian_density_function(gmm_da
        q=np.sum(alpha,axis=1)
        for i in range(len(q)):
```

alpha[i,:] = np. divide (alpha[i,:],q[i])

```
#m step
def m step(gauss, alpha):
        for j in range(4):
                # mu new
                # caclulate N_j
                N_j=np.sum(alpha[:,j])
                # calculate mu
                gauss[j][0] = np.array([0,0])
                for i in range(N):
                        gauss[j][0] = gauss[j][0] + alpha[i,j]*gmm_data_points[i]
                gauss[j][0] = gauss[j][0] / N_j
                # sigma new
                gauss[j][1] = np.zeros([2,2])
                for i in range(N):
                        gauss[j][1] = gauss[j][1] + alpha[i,j]*np.outer(gmm_data_poi
                gauss[j][1] = gauss[j][1] / N_j
                # pi new
                gauss[j][2] = N_j / N
        return gauss
def plot_all(gauss):
        delta = 0.25
        x = np.arange(-5.0, 5.0, delta)
        y = np.arange(-5.0, 5.0, delta)
        xx, yy = np.meshgrid(x, y)
        height = np.zeros((len(xx), len(xx)))
        height2 = np.zeros((len(xx), len(xx)))
        height3 = np.zeros((len(xx), len(xx)))
        height4 = np.zeros((len(xx), len(xx)))
        for i in range(len(xx)):
                for j in range(len(xx)):
                        height[i,j] = gaussian_density_function([xx[i,j], yy[i,j]],
                        height2[i,j] = gaussian_density_function([xx[i,j], yy[i,j]],
                        height3[i,j] = gaussian_density_function([xx[i,j], yy[i,j]],
                        height4[i,j] = gaussian_density_function([xx[i,j], yy[i,j]],
        plt.scatter(gmm_data_points[:,0], gmm_data_points[:,1], color="black")
        plt.contour(xx,yy,height, colors = 'red')
        plt.contour(xx,yy,height2, colors = 'blue')
        plt.contour(xx,yy,height3, colors = 'yellow')
        plt.contour(xx,yy,height4, colors = 'green')
        plt.show()
        return
result=np.zeros([31])
for i in range(31):
        a2=e step(gauss1)
        gauss1 = m_step(gauss1, a2)
        result[i] = log likehood gaussian(gmm data points, gauss1)
        if i in [1,3,5,10,30]:
```

return alpha

Calling

we get the following results:

