CPSC 540 Assignment 3 (due February 27)

Density Estimation and Project Proposal

1 Discrete and Gaussian Variables

1.1 MLE for General Discrete Distribution

Consider a density estimation task, where we have two variables (d = 2) that can each take one of k discrete values. For example, we could have

$$X = \begin{bmatrix} 1 & 3 \\ 4 & 2 \\ k & 3 \\ 1 & k-1 \end{bmatrix}.$$

The likelihood for example x^i under a general discrete distribution would be

$$p(x_1^i, x_2^i | \Theta) = \theta_{x_1^i, x_2^i},$$

where θ_{c_1,c_2} gives the probability of x_1 being in state c_1 and x_2 being in state c_2 , for all the k^2 combinations of the two variables. In order for this to define a valid probability, we need all elements θ_{c_1,c_2} to be non-negative and they must sum to one, $\sum_{c_1=1}^k \sum_{c_2=1}^k \theta_{c_1,c_2} = 1$.

- 1. Given n training examples, derive the MLE for the k^2 elements of Θ .
- 2. Because of the sum-to-1 constraint, there are only (k^2-1) degrees of freedom in the discrete distribution, and not k^2 . Derive the MLE for this distribution assuming that

$$\theta_{k,k} = 1 - \sum_{c_1=1}^{k} \sum_{c_2=1}^{k} \mathcal{I}[c_1 \neq k, c_2 \neq k] \theta_{c_1, c_2},$$

so that the distribution only has $(k^2 - 1)$ parameters.

3. If we had separate parameter θ_{c_1} and θ_{c_2} for each variables, a reasonable choice of a prior would be a product of Dirichlet distributions,

$$p(\theta_{c_1}, \theta_{c_2}) \propto \theta_{c_1}^{\alpha_{c_1} - 1} \theta_{c_2}^{\alpha_{c_2} - 1}$$

For the general discrete distribution, a prior encoding the same assumptions would be

$$p(\theta_{c_1,c_2}) \propto \theta_{c_1,c_2}^{\alpha_{c_1}+\alpha_{c_2}-2}$$
.

Derive the MAP estimate under this prior.

Hint: it is convenient to write the likelihood for an example i in the form

$$p(x^i|\Theta) = \prod_{c \in [k]^2} \theta_c^{\mathcal{I}[x^i = c]},$$

where c is a vector containing (c_1, c_2) , $[x^i = c]$ evaluates to 1 if all elements are equal, and $[k]^2$ is all ordered pairs (c_1, c_2) . You can use the Lagrangian to enforce the sum-to-1 constraint on the log-likelihood, and you may find it convenient to define $N_c = \sum_{i=1}^n \mathcal{I}[x^i = c]$.

Student's Solution:

1. The MLE for a discrete distribution on k^2 elements $(j,\ell) \in [k] \times [k]$ is given by

$$\underset{0 \leq \Theta \leq 1}{\operatorname{arg\,max}} \, p(X \mid \Theta) = \underset{\Theta \in [0,1]^k}{\operatorname{arg\,max}} \, \prod_{i=1}^n p(x_1^i, x_2^i \mid \Theta) = \underset{\Theta \in [0,1]^k}{\operatorname{arg\,max}} \, \prod_{j,\ell \in [k]} \theta_{j,\ell}^{N_{j,\ell}}$$

where $N_{j,\ell} := \#\{i \in [n] : x_1^i = j \& x_2^i = \ell\}$ and $0 \le \Theta \le 1$ is multi-index notation for $\Theta \in [0,1]^k$.

2. From above we see that the assumption for this question yields, where $J:=\{(j,\ell)\in [k]^2: j,\ell\neq k\},$

$$\mathop{\arg\max}_{0\leq\Theta\leq1}p(X\mid\Theta) = \mathop{\arg\max}_{0\leq\Theta\leq1} \left(1 - \sum_{j,\ell\in J}\theta_{j,\ell}\right)^{N_{k,k}} \prod_{j,\ell\in J}\theta_{j,\ell}^{N_{j,\ell}}$$

Then the MLE can be determined by finding the critical point of the log-likelihood (since log is a concave function, hence does not change the maximizer). For $\lambda \in J$,

$$0 = \frac{\partial}{\partial \theta_{\lambda}} \log \mathcal{L}(\Theta \mid X) = N_{k,k} \frac{\partial}{\partial \theta_{\lambda}} \log \left(1 - \sum_{j,\ell \in J} \theta_{j,\ell} \right) + \frac{\partial}{\partial \theta_{\lambda}} \sum_{j,\ell \in J} N_{j,\ell} \log \theta_{j,\ell}$$
$$= \frac{-N_{k,k}}{1 - \sum_{j,\ell \in J} \theta_{j,\ell}} + \frac{N_{\lambda}}{\theta_{\lambda}}$$

Hence, letting $J(j,\ell) := \{(m,n) \in [k] \times [k] : (m,n) \neq (k,k), (m,n) \neq (j,\ell)\},\$

$$\frac{N_{\lambda}}{\theta_{\lambda}} = \frac{N_{k,k}}{1 - \sum_{j,\ell \in J} \theta_{j,\ell}} \quad \Longrightarrow \quad \frac{\theta_{\lambda}}{1 - \sum_{j,\ell \in J} \theta_{j,\ell}} = \frac{N_{\lambda}}{N_{k,k}}$$

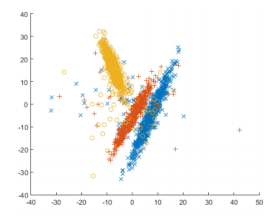
thereby implying, since selecting (k, k) above was arbitrary,

$$\theta_{\lambda} := \frac{N_{\lambda}}{\sum_{j,\ell \in [k]^2} N_{j,\ell}}$$

3.

1.2 Generative Classifiers with Gaussian Assumption

Consider the 3-class classification dataset in this image:



In this dataset, we have 2 features and each colour represents one of the classes. Note that the classes are highly-structured: the colours each roughly follow a Gausian distribution plus some noisy samples.

Since we have an idea of what the features look like for each class, we might consider classifying inputs x using a generative classifier. In particular, we are going to use Bayes rule to write

$$p(y = c|x, \Theta) = \frac{p(x|y = c, \Theta) \cdot p(y = c|\Theta)}{p(x|\Theta)},$$

where Θ represents the parameters of our model. To classify a new example \hat{x} , generative classifiers would use

$$\hat{y} = \underset{y \in \{1,2,\dots,k\}}{\arg\max} p(\hat{x}|y=c,\Theta) p(y=c|\Theta),$$

where in our case the total number of classes k is 3 (The denominator $p(\hat{x}|\Theta)$ is irrelevant to the classification since it is the same for all y.) Modeling $p(y = c|\Theta)$ is eays: we can just use a k-state categorical distribution,

$$p(y = c|\Theta) = \theta_c,$$

where θ_c is a single parameter for class c. The maximum likelihood estimate of θ_c is given by n_c/n , the number of times we have $y^i = c$ (which we've called n_c) divided by the total number of data points n.

Modeling $p(x|y=c,\Theta)$ is the hard part: we need to know the probability of seeing the feature vector x given that we are in class c. This corresponds to solving a density estimation problem for each of the k possible classes. To make the density estimation problem tractable, we'll assume that the distribution of x given that y=c is given by a $\mathcal{N}(\mu_c, \Sigma_c)$ Gaussian distribution for a class-specific μ_c and Σ_c ,

$$p(x|y=c,\Theta) = \frac{1}{(2\pi)^{\frac{d}{2}} |\Sigma_c|^{\frac{1}{2}}} \exp\left(-\frac{1}{2}(x-\mu_c)^T \Sigma_c^{-1} (x-\mu_c)\right).$$

Since we are distinguishing between the probability under k different Gaussians to make our classification, this is called Gaussian discriminant analysis (GDA). In the special case where we have a constant $\Sigma_c = \Sigma$ across all classes it is known as linear discriminant analysis (LDA) since it leads to a linear classifier between any two classes (while the region of space assigned to each class forms a convex polyhedron as in k-means clustering). Another common restriction on the Σ_c is that they are diagonal matrices, since this only requires O(d) parameters instead of $O(d^2)$ (corresponding to assuming that the features are independent univariate Gaussians given the class label). Given a dataset $\mathcal{D} = \{(x^i, y^i)\}_{i=1}^n$, where $x^i \in \mathbb{R}^d$ and $y^i \in \{1, \ldots, k\}$, the maximum likelihood estimate (MLE) for the μ_c and Σ_c in the GDA model is the solution to

$$\underset{\mu_1,\mu_2,\dots,\mu_k,\Sigma_1,\Sigma_2,\dots,\Sigma_k}{\arg\max} \prod_{i=1}^n p(x^i|y^i,\mu_{y^i},\Sigma_{y^i}).$$

This means that the negative log-likelihood will be equal to

$$-\log p(X|y,\Theta) = -\sum_{i=1}^{n} \log p(x^{i}|y^{i}|\mu_{y^{i}}, \Sigma_{y^{i}})$$
$$= \sum_{i=1}^{n} \frac{1}{2} (x^{i} - \mu_{y^{i}})^{T} \Sigma_{y^{i}}^{-1} (x^{i} - \mu_{y^{i}}) + \frac{1}{2} \sum_{i=1}^{n} \log |\Sigma_{y^{i}}| + \text{const.}$$

- 1. Derive the MLE for the GDA model under the assumption of common diagonal covariance matrices, $\Sigma_c = D$ (d parameters). (Each class will have its own mean μ_c .)
- 2. Derive the MLE for the GDA model under the assumption of individual scale-identity matrices, $\Sigma_c = \sigma_c^2 I$ (k parameters).

- 3. It's painful to derive these from scratch, but you should be able to see a pattern that would allow other common restrictions. Without deriving the result from scratch (hopefully), give the MLE for the case of individual full covariance matrices, Σ_c ($O(kd^2)$ parameters).
- 4. When you run example_generative it loads a variant of the dataset in the figure that has 12 features and 10 classes. This data has been split up into a training and test set, and the code fits a k-nearest neighbour classifier to the training set then reports the accuracy on the test data ($\sim 36\%$). The k-nearest neighbour model does poorly here since it doesn't take into account the Gaussian-like structure in feature space for each class label. Write a function generative Gaussian that fits a GDA model to this dataset (using individual full covariance matrices). Hand in the function and report the test set accuracy.
- 5. In this question we would like to replace the Gaussian distribution of the previous problem with the more robust multivariate-t distribution so that it isn't influenced as much by the noisy data. Unlike the previous case, we don't have a closed-form solution for the parameters. However, if you run example_tdist it generates random noisy data and fits a multivariate-t model (you will need to add the minFunc directory to the Matlab path for the demo to work). By using the multivariateT model, write a new function generativeStudent that implements a generative model that is based on the multivariate-t distribution instead of the Gaussian distribution. Report the test accuracy with this model.

Hints: you will be able to substantially simplify the notation in parts 1-3 if you use the notation $\sum_{i \in y_c}$ to mean the sum over all values i where $y^i = c$. Similarly, you can use n_c to denote the number of cases where $y_i = c$, so that we have $\sum_{i \in y_c} 1 = n_c$. Note that the determinant of a diagonal matrix is the product of the diagonal entries, and the inverse of a diagonal matrix is a diagonal matrix with the reciprocals of the original matrix along the diagonal. For part three you can use the result from class regarding the MLE of a general multivariate Gaussian. You may find it helpful to use the included logdet.m function to compute the log-determinant in more numerically-stable way.

Student's Solution:

1. Derive the MLE for the GDA model under the assumption of common diagonal covariance matrices $\Sigma_c = D$ (d parameters). (Each class will have its own mean μ_c .)

We first start to get the MLE in respect with μ_c . We take the gradient of the negative log-likelihood with respect to μ_c an set it to zero:

$$\begin{split} -\nabla_{\mu_c} \log p(X|y,\Theta) &= -\nabla_{\mu_c} \sum_{i=1}^n \log p(x^i|y^i|\mu_{y^i}, \Sigma_{y^i}) \\ &= \sum_{i=1}^n \mathcal{I}[y^i = c] \Sigma_{y^i}^{-1} (x^i - \mu_{y^i}) \\ -\nabla_{\mu_c} \log p(X|y,\Theta) &= 0 \implies \mu_c = \frac{1}{n_c} \sum_{i=1}^n \mathcal{I}[y^i = c] x^i \end{split}$$

We then can use this result for μ_c to get the MLE for the covariances matrices under the assumption of common diagonal covariance matrices. Using the derivation seen in class, we reparametrize the MLE in term of the precision matrix $\Phi = \Sigma^{-1}$ and the all samples covariance matrix $S = \frac{1}{n} \sum_{i=1}^{n} (x^i - \mu_{u^i})(x^i - \mu_{u^i})^T$:

 $-\log p(X|y,\Theta) = \frac{n}{2}Tr(S\Phi) - \frac{n}{2}log|\Phi|$

We now put in our assumption of common diagonal covariance matrices, we only to derive the diagonal

elements, the others are by assumption equal to 0, which implies:

$$\begin{split} -\nabla_{D^{-1} = \Phi} \log p(X|y,\Theta) &= \nabla_{D^{-1} = \Phi} \big(\frac{n}{2} Tr(S\Phi) - \frac{n}{2} log|\Phi|\big) \\ &= \frac{n}{2} diag(S) - \frac{n}{2} D \end{split}$$

Which gives us the following result:

$$\frac{n}{2}diag(S) - \frac{n}{2} = 0 \implies D = diag(S)$$

2. Derive the MLE for the GDA model under the assumption of individual scale-identity matrices, $\Sigma_c = \sigma_c^2 I$ (k parameters)

We will first notice that changing assumption on Σ_c does not change the MLE for μ_c . We then notice that here our assumption on the covariance matrix is strong, in the sense that we impose all the diagonal terms to be equal. We cannot use the reparametrization used earlier.

$$-\log p(X|y,\Theta) = \sum_{i=1}^{n} \frac{1}{2} \sigma_{y^{i}}^{-2} (x^{i} - \mu_{y^{i}})^{T} (x^{i} - \mu_{y^{i}}) + \frac{1}{2} \sum_{i=1}^{n} \log |\sigma_{y^{i}}^{2}| + \text{const.}$$
$$-\nabla_{\sigma_{c}^{2}} \log p(X|y,\Theta) = \frac{1}{2\sigma_{c}^{4}} \sum_{i=1}^{n} \mathcal{I}[y^{i} = c](x^{i} - \mu_{y^{i}})^{T} (x^{i} - \mu_{y^{i}}) + \frac{n_{c}}{2} \sigma_{c}^{-2}$$

which gives us the result:

$$-\nabla_{\sigma_c^2} \log p(X|y,\Theta) = 0 \implies \sigma_c^2 = \frac{1}{n_c} \sum_{i=1}^n \mathcal{I}[y^i = c](x^i - \mu_{y^i})^T (x^i - \mu_{y^i})$$

3. Without derivation, give the MLE for the case of *individual full* covariance matrices Your intuition in that case is:

$$\Sigma_c = \frac{1}{n_c} \sum_{i=1}^n \mathcal{I}[y^i = c](x^i - \mu_{y^i})(x^i - \mu_{y^i})^T$$

4. Write a function *generativeGaussian* that fits a GDA model to this dataset (using individual full covariance matrices). Hand in the function and report the test set accuracy.

```
function model = generativeGaussian(Xtrain,Ytrain,k)

[n,d] = size(Xtrain);

%training dataset
model.Xtrain = Xtrain;
model.Ytrain = Ytrain;
%number of class
model.K = k;

%Initialize cell array to store
%averages mu,
%probability of each class theta
%Covariances matrices SIGMA
```

```
mu = cell(k,1);
theta = zeros(k,1);
SIGMA = cell(k,1);
%For each class i
for i=1:k
    SIGMA{i} = zeros(d);
    ind = find(Ytrain == i);
    %Compute probability of the class theta_i
    theta(i) = sum(Ytrain == i)/n;
    %Compute average mu_i
    mu\{i\} = (1/(n*theta(i)))*sum(Xtrain(ind,:))';
    %Compute cavariance matrix SIGMA_i
    for j=1:size(ind,1)
        SIGMA\{i\} = SIGMA\{i\} + (1/(n*theta(i)))*((Xtrain(ind(j),:)'-\leftarrow
           mu{i})*(Xtrain(ind(j),:)'-mu{i})');
    end
end
model.predict = @(model, Xtest) predict(model, Xtest);
model.mu = mu;
model.theta = theta;
model.SIGMA = SIGMA;
end
%Predict function for test data
function yhat = predict(model, Xtest)
[nTest,d] = size(Xtest);
%Probability distribution
prob = @(x,mu,SIGMA) (1/(((2*pi)^(d/2))*sqrt(det(SIGMA))))*exp(-0.5* \leftarrow
   (x-mu)'*(SIGMA \setminus (x-mu));
ytemp = zeros(nTest, model.K);
%for each test data
for i=1:nTest
    %compute the probability to be in each class
    for j=1:model.K
        ytemp(i,j) = prob(Xtest(i,:)',model.mu{j},model.SIGMA{j});
    end
end
%Classify in the class with maximum probability
[",yhat] = max(ytemp,[],2);
end
```

Our test set accuracy is now 0.63.

5. Write a new function *generativeStudent* that implements a generative model that is based on the multivariate-t distribution instead of the Gaussian distribution. Report the test accuracy with this model.

```
function model = generativeStudent(Xtrain, Ytrain, k)
[n,d] = size(Xtrain);
%our model is a set of k multivariate T model
model.tdist = cell(k,1);
%For each class
for i=1:k
    %Fit a multivariate T model
    ind = find(Ytrain == i);
    model.tdist{i} = multivariateT(Xtrain(ind,:));
    model.tdist{i}.Xtrain = Xtrain(ind,:);
    model.tdist{i}.Ytrain = Ytrain(ind);
end
model.predict = @(model, Xtest) predict(model, Xtest);
model.K = k;
end
%Predict function for test data
function yhat = predict(model, Xtest)
[nTest,d] = size(Xtest);
ytemp = zeros(nTest, model.K);
%for each test data
for i=1:nTest
    %compute the probability to be in each class
    for j=1:model.K
        ytemp(i,j) = model.tdist{j}.pdf(model.tdist{j},Xtest(i,:));
    end
end
%Classify in the class with maximum probability
[",yhat] = max(ytemp,[],2);
end
```

Our test set accuracy is now 0.79.

Method	Accuracy
KNN	0.37
GDA	0.63
Student	0.79

1.3 Self-Conjugacy for the Mean Parameter

If x is distributed according to a Gaussian with mean μ ,

$$x \sim \mathcal{N}(\mu, \sigma^2),$$

and we assume that μ itself is distributed according to a Gaussian

$$\mu \sim \mathcal{N}(\alpha, \gamma^2),$$

then the posterior $\mu|x$ also follows a Gaussian distribution.¹ Derive the form of the (Gaussian) distribution for $p(\mu|x,\alpha,\sigma^2,\gamma^2)$.

Hints: Use Bayes rule and use the \propto sign to get rid of factors that don't depend on μ . You can "complete the square" to make the product look like a Gaussian distribution, e.g. when you have $\exp(ax^2 - bx + \text{const})$ you can factor out an a and add/subtract $(b/2a)^2$ to re-write it as

$$\exp\left(ax^2 - bx + const\right) \propto \exp\left(ax^2 - bx\right) = \exp\left(a(x^2 - (b/a)x)\right)$$
$$\propto \exp\left(a(x^2 - (b/a)x + (b/2a)^2)\right) = \exp\left(a(x - (b/2a))^2\right).$$

Note that multiplying by factors that do not depend on μ within the exponent does not change the distribution. In this question you will want to complete the square to get the distribution on μ , rather than x. You may find it easier to solve thie problem if you parameterize the Gaussians in terms of their 'precision' parameters (e.g., $\lambda = 1/\sigma^2$, $\lambda_0 = 1/\gamma^2$) rather than their variances σ^2 and γ^2 .

2 Mixture Models and Expectation Maximization

2.1 Semi-Supervised Gaussian Discriminant Analysis

Consider fitting a GDA model where some of the y^i values are missing at random. In particular, let's assume we have n labeled examples (x^i, y^i) and then another another t unlabeled examples (x^i) . This is a special case of semi-supervised learning, and fitting generative models with EM is one of the oldest semi-supervised learning techniques. When the classes exhibit clear structure in the feature space, it can be very effective even if the number of labeled examples is very small.

- 1. Derive the EM update for fitting the parameters of a GDA model (with individual full covariance matrices) in the semi-superivsed setting where we have n labeled examples and t unlabeled examples.
- 2. If you run the demo example_SSL, it will load a variant of the dataset from the previous question, but where the number of labeled examples is small and a large number of unlabeled examples are available. The demo first fits a KNN model and then a generative Gaussian model (once you are finished Question 1). Because the number of labeled examples it quite small, the performance is worse than in Question 2. Write a function generative Gaussian SSL that fits the generative Gaussian model of the previous question using EM to incorporate the unlabeled data. Hand in the function and report the test error when training on the full dataset.
- 3. Repeat the previous part, but using the "hard"-EM algorithm where we explicitly classify all the unlabeled examples. How does this change the performance and the number of iterations?

¹We say that the Gaussian distribution is the 'conjugate prior' for the Gaussian mean parameter (we'll formally discuss conjugate priors later in the course). Another reason the Gaussian distribution is important is that is the only (non-trivial) continuous distribution that has this 'self-conjugacy' property.

Hint: for the first question most of the work has been done for you in the EM notes on the course webpage. You can use the result (**) and the update of θ_c from those notes, but you will need to work out the update of the parameters of the Gaussian distribution $p(x^i|y^i,\Theta)$.

Hint: for the second question, although EM often leads to simple updates, implementing them correctly can often be a pain. One way to help debug your code is to compute the observed-data log-likelihood after every iteration. If this number goes down, then you know your implementation has a problem. You can also test your updates of sets of variables in this way too. For example, if you hold the μ_c and Σ_c fixed and only update the θ_c , then the log-liklihood should not go down. In this way, you can test each of combinationso of updates on their to make sure they are correct.

2.2 Mixture of Bernoullis

The function example_Bernoulli loads a binarized version of the MNIST dataset and fits a density model that uses an independent Bernoulli to model each feature. If the reports the average NLL on the test data and shows 4 samples generated from the model. Unfortunately, the test NLL is infinity and the samples look terrible.

- 1. To address the problem that the average NLL is infinity, modify the *densityBernoulli* function to implement Laplace smoothing based on an extra argument α . Hand in the code and report the average NLL with $\alpha = 1$.
- 2. Write a new function implementing the mixture of Bernoullis model with Laplace smoothing of the θ values (note that Laplace smoothing only change the M-step). Hand in the code and report the average NLL with $\alpha = 1$ and k = 10 for a particular run of the algorithm, as well as 4 samples from the model and 4 of the cluster images.