

## CPSC 340 Assignment 2 (due 2021-05-24 at 9:25am)

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### Important: Submission Format

Please make sure to follow the submission instructions posted on Piazza. **We are entitled to deduct 50% of your marks if the submission format is incorrect.**

## 1 Training and Testing

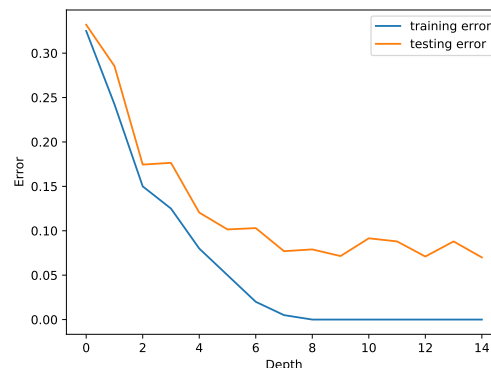
If you run `python main.py -q 1`, it will load the *citiesSmall.pkl* data set from Assignment 1. Note that this file contains not only training data, but also test data, `X_test` and `y_test`. After training a depth-2 decision tree with the information gain splitting rule, it will evaluate the performance of the classifier on the test data. With a depth-2 decision tree, the training and test error are fairly close, so the model hasn't overfit much.

### 1.1 Training and Testing Error Curves

Use decision tree classifier with information gain splitting rule to train and test your model. **Make a plot that contains the training error and testing error as you vary the depth from 1 through 15. How do each of these errors change with the decision tree depth?**

Note: it's OK to reuse code from Assignment 1.

Answer: The plot should look like this:



The training error goes down monotonically to 0 (quickly at first and then slowly). The test error also starts going down but after a depth of 9 it goes up and then stays flat.

## 1.2 Validation Set

Suppose that we didn't have an explicit test set available. In this case, we might instead use a *validation* set. Split the training set into two equal-sized parts: use the first  $n/2$  examples as a training set and the second  $n/2$  examples as a validation set (we're assuming that the examples are already in a random order). What depth of decision tree would we pick to minimize the validation set error? Does the answer change if you switch the training and validation set? How could we use more of our data to estimate the depth more reliably?

Answer: Using sklearn's implementation with `random_state=1` I got the best depth to be 8 using the first half for training, and 6 when using the second half for training. To make the estimate more reliable, you could use cross-validation.

## 1.3 Trustworthiness of Validation Error

Repeat the above experiment, now using **first fifteen (15) examples in the training data as a validation set** and the rest as a training set. Choose the best depth for the tree based on validation error. Report the best depth and the validation error. Do you trust this validation error? How does your answer change if you switch the training and validation sets? Explain.

Answer: With first 15 examples in validation set, I get depth=7 and validation error=0.00. The number of validation examples is very small, which makes the validation error not a reliable estimate of the test error. Hence we overfitted to the validation set. I get test error=0.085. With first 15 examples in training set, I get depth=3 and validation error=0.268. The number of training examples is very small, so even though I have a fairly large validation set, my model struggles to generalize to validation examples in a meaningful way. I shouldn't trust the validation error in both cases.

## 2 K-Nearest Neighbours

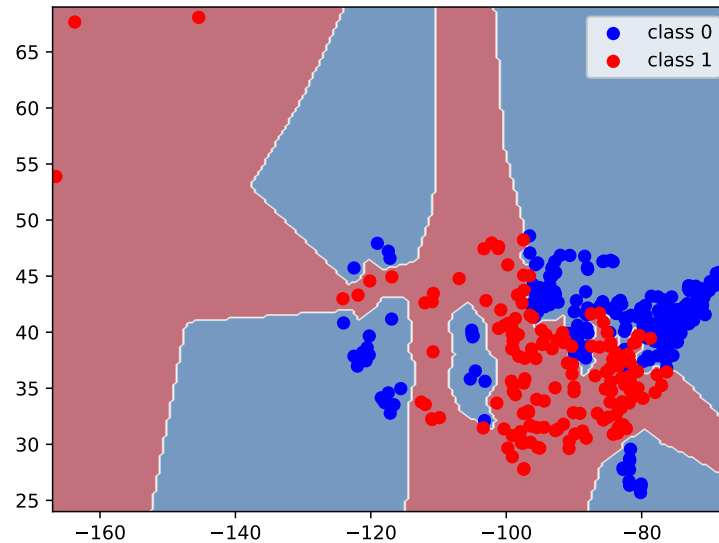
In the *citiesSmall* dataset, nearby points tend to receive the same class label because they are part of the same U.S. state. For this problem, perhaps a  $k$ -nearest neighbours classifier might be a better choice than a decision tree. The file *knn.py* has implemented the training function for a  $k$ -nearest neighbour classifier (which is to just memorize the data).

Fill in the `predict` function in *knn.py* so that the model file implements the  $k$ -nearest neighbour prediction rule. You should find Euclidean distance, and may numpy's `sort` and/or `argsort` functions useful. You can also use `utils.euclidean_dist_squared`, which computes the squared Euclidean distances between all pairs of points in two matrices.

1. Write the `predict` function.
2. Report the training and test error obtained on the *citiesSmall* dataset for  $k = 1$ ,  $k = 3$ , and  $k = 10$ . How do these numbers compare to what you got with the decision tree?
3. Hand in the plot generated by `utils.plot_classifier` on the *citiesSmall* dataset for  $k = 1$ , using both your implementation of KNN and the `KNeighborsClassifier` from scikit-learn.
4. Why is the training error 0 for  $k = 1$ ?
5. Recall that we want to choose hyper-parameters so that the test error is (hopefully) minimized. How would you choose  $k$ ?

Answer:

1. See knn.py.
2. The training/test errors are:
  - $k = 1$ : training is 0, test is 0.065.
  - $k = 3$ : training is 0.028, test is 0.066.
  - $k = 10$ : training is 0.072, test is 0.097.
3. The plot should look like this:



4. The training error is 0 for  $k = 1$  because every training example is 1-nearest neighbour of itself, so when you are predicting you just copy the labels from the training data. (Unless you have duplicate training examples, 1-nearest neighbour always obtains a training error of 0: this is why reporting low training errors is meaningless.)
5. The training error strictly goes down as  $k$  decreases, so you can't use the training error to choose  $k$ . Instead, you should split your data into a training and validation set, or use cross-validation.

### 3 Naive Bayes

In this section we'll implement naive Bayes, a very fast classification method that is often surprisingly accurate for text data with simple representations like bag of words.

## 3.1 Naive Bayes by Hand

Consider the dataset below, which has 10 training examples and 3 features:

$$X = \begin{bmatrix} 0 & 0 & 1 \\ 0 & 1 & 1 \\ 0 & 1 & 1 \\ 1 & 1 & 0 \\ 0 & 1 & 0 \\ 0 & 1 & 1 \\ 1 & 0 & 0 \\ 1 & 1 & 0 \\ 1 & 0 & 1 \\ 1 & 0 & 0 \end{bmatrix}, \quad y = \begin{bmatrix} \text{spam} \\ \text{spam} \\ \text{spam} \\ \text{spam} \\ \text{spam} \\ \text{spam} \\ \text{not spam} \\ \text{not spam} \\ \text{not spam} \\ \text{not spam} \end{bmatrix}.$$

The feature in the first column is <your name> (whether the e-mail contained your name), in the second column is “pharmaceutical” (whether the e-mail contained this word), and the third column is “PayPal” (whether the e-mail contained this word). Suppose you believe that a naive Bayes model would be appropriate for this dataset, and you want to classify the following test example:

$$\hat{x} = [1 \quad 1 \quad 0].$$

### 3.1.1 Prior probabilities

Compute the estimates of the class prior probabilities, which I also called the “baseline spam-ness” in class. (you don’t need to show any work):

- $p(\text{spam})$ .

Answer: 6/10.

- $p(\text{not spam})$ .

Answer: 4/10.

### 3.1.2 Conditional probabilities

Compute the estimates of the 6 conditional probabilities required by naive Bayes for this example (you don’t need to show any work):

- $p(<\text{your name}> = 1 \mid \text{spam})$ .

Answer: 1/6.

- $p(\text{pharmaceutical} = 1 \mid \text{spam})$ .

Answer: 5/6.

- $p(\text{PayPal} = 0 \mid \text{spam})$ .

Answer: 2/6.

- $p(<\text{your name}> = 1 \mid \text{not spam})$ .

Answer: 1.

- $p(\text{pharmaceutical} = 1 \mid \text{not spam})$ .

Answer: 1/4.

- $p(\text{PayPal} = 0 \mid \text{not spam})$ .

Answer:  $3/4$ .

### 3.1.3 Prediction

Under the naive Bayes model and your estimates of the above probabilities, what is the most likely label for the test example? (**Show your work.**)

Answer:

$$\begin{aligned} p(\text{spam} \mid x_1 = 1, x_2 = 1, x_3 = 0) &\propto p(x_1 = 1, x_2 = 1, x_3 = 0 \mid \text{spam})p(\text{spam}) \\ &= p(x_1 = 1 \mid \text{spam})p(x_2 = 1 \mid \text{spam})p(x_3 = 0 \mid \text{spam})p(\text{spam}) \\ &= (1/6)(5/6)(2/6)(6/10) \\ &\approx 0.028 \end{aligned}$$

$$\begin{aligned} p(\text{not spam} \mid x_1 = 1, x_2 = 1, x_3 = 0) &\propto p(x_1 = 1, x_2 = 1, x_3 = 0 \mid \text{not spam})p(\text{not spam}) \\ &= p(x_1 = 1 \mid \text{not spam})p(x_2 = 1 \mid \text{not spam})p(x_3 = 0 \mid \text{not spam})p(\text{not spam}) \\ &= (1)(1/4)(3/4)(4/10) \\ &\approx 0.075. \end{aligned}$$

Since  $p(\text{not spam} \mid x_1 = 1, x_2 = 1, x_3 = 0)$  is proportional to a bigger number, and the proportionality constants are the same ( $p(x_1 = 1, x_2 = 1, x_3 = 0)$ ), we would predict “not spam”.

### 3.1.4 Simulating Laplace Smoothing with Data

One way to think of Laplace smoothing is that you’re augmenting the training set with extra counts. Consider the estimates of the conditional probabilities in this dataset when we use Laplace smoothing (with  $\beta = 1$ ). Give a set of extra training examples that we could add to the original training set that would make the basic estimates give us the estimates with Laplace smoothing (in other words give a set of extra training examples that, if they were included in the training set and we didn’t use Laplace smoothing, would give the same estimates of the conditional probabilities as using the original dataset with Laplace smoothing). Present your answer in a reasonably easy-to-read format, for example the same format as the data set at the start of this question.

Answer: You could add the following examples:

$$X_\beta = \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 0 \\ 1 & 1 & 1 \\ 1 & 1 & 1 \end{bmatrix}, \quad y_\beta = \begin{bmatrix} 0 \\ 1 \\ 1 \\ 0 \end{bmatrix}.$$

## 3.2 Exploring Bag-of-Words

If you run `python main.py -q 3.2`, it will load the following dataset:

1. **X**: A binary matrix. Each row corresponds to a newsgroup post, and each column corresponds to whether a particular word was used in the post. A value of 1 means that the word occurred in the post.
2. **wordlist**: The set of words that correspond to each column.
3. **y**: A vector with values 0 through 3, with the value corresponding to the newsgroup that the post came from.

4. `groupnames`: The names of the four newsgroups.
5. `Xvalidate` and `yvalidate`: the word lists and newsgroup labels for additional newsgroup posts.

Answer the following:

1. Which word corresponds to column 41 of  $X$ ? (This is column 40 in Python.)

Answer: "honda"

2. Which words are present in training example 401?

Answer: "christian", "fact", "god", "studies", "university"

3. Which newsgroup name does training example 401 come from?

Answer: "talk.\*"

### 3.3 Naive Bayes Implementation

If you run `python main.py -q 2.3` it will load the newsgroups dataset, fit a basic naive Bayes model and report the validation error.

The `predict()` function of the naive Bayes classifier is already implemented. However, in `fit()` the calculation of the variable `p_xy` is incorrect (right now, it just sets all values to  $1/2$ ). [Modify this function so that `p\_xy` correctly computes the conditional probabilities of these values based on the frequencies in the data set. Submit your code. Report the training and validation errors that you obtain.](#)

Answer: The training error is approximately 0.20. The validation error is approximately 0.19.

### 3.4 Laplace Smoothing Implementation

Laplace smoothing is one way to prevent failure cases of Naive Bayes based on counting. Recall what you know from lecture to implement Laplace smoothing to your Naive Bayes model. [Do the following:](#)

- [Modify the `NaiveBayesLaplace` class provided in `naive\_bayes.py` and write its `fit\(\)` method to implement Laplace smoothing. Submit this code.](#)
- [Using the same data as the previous section, fit Naive Bayes models with \*\*and\*\* without Laplace smoothing to the training data. Use  \$\beta = 1\$  for Laplace smoothing. For each model, look at  \$p\(x\_{ij} = 1 \mid y\_i = 0\)\$  across all  \$j\$  values \(i.e. all features\) in both models. Do you notice any difference? Explain.](#)
- [One more time, fit a Naive Bayes model with Laplace smoothing using  \$\beta = 10000\$ . Look at  \$p\(x\_{ij} = 1 \mid y\_i = 0\)\$ . Do these numbers look like what you expect? Explain.](#)

Answer: See code. I have 12 zeros appear without Laplace smoothing, and there are no zeros with Laplace smoothing. When I set  $\beta = 10000$ , I see all of the numbers are converging to  $\frac{1}{4}$ , which is expected.

### 3.5 Runtime of Naive Bayes for Discrete Data

For a given training example  $i$ , the predict function in the provided code computes the quantity

$$p(y_i \mid x_i) \propto p(y_i) \prod_{j=1}^d p(x_{ij} \mid y_i),$$

for each class  $y_i$  (and where the proportionality constant is not relevant). For many problems, a lot of the  $p(x_{ij} \mid y_i)$  values may be very small. This can cause the above product to underflow. The standard fix for

this is to compute the logarithm of this quantity and use that  $\log(ab) = \log(a) + \log(b)$ ,

$$\log p(y_i | x_i) = \log p(y_i) + \sum_{j=1}^d \log p(x_{ij} | y_i) + (\text{irrelevant proportionality constant}).$$

This turns the multiplications into additions and thus typically would not underflow.

Assume you have the following setup:

- The training set has  $n$  objects each with  $d$  features.
- The test set has  $t$  objects with  $d$  features.
- Each feature can have up to  $c$  discrete values (you can assume  $c \leq n$ ).
- There are  $k$  class labels (you can assume  $k \leq n$ )

You can implement the training phase of a naive Bayes classifier in this setup in  $O(nd)$ , since you only need to do a constant amount of work for each  $X(i, j)$  value. (You do not have to actually implement it in this way for the previous question, but you should think about how this could be done.) [What is the cost of classifying  \$t\$  test examples with the model and this way of computing the predictions?](#)

Answer: For each of the  $t$  examples, the dominant cost is computing  $p(x_{ij}|y_i)$  for all  $d$  values of  $j$  and all  $k$  class labels. You can do this with three “for” loops (as in the naive Bayes `predict` function in the given code): one looping over the examples  $t$ , one looping over the features  $d$ , and one looping over the class labels  $k$ . Since each of the loops does a constant amount of work, the total time is  $O(tdk)$ . Note that this is much slower than using a depth  $m$  decision tree in the common case that  $m \ll dk$ . (However, the training and testing phases can be much faster if the examples are sparse, meaning that most values of  $x_{ij}$  are zero.)

## 4 Random Forests

### 4.1 Implementation

The file `vowels.pkl` contains a supervised learning dataset where we are trying to predict which of the 11 “steady-state” English vowels that a speaker is trying to pronounce.

You are provided with a `RandomStump` class that differs from `DecisionStumpInfoGain` in that it only considers  $\lfloor \sqrt{d} \rfloor$  randomly-chosen features.<sup>1</sup> You are also provided with a `RandomTree` class that is exactly the same as `DecisionTree` except that it uses `RandomStump` instead of `DecisionStump` and it takes a bootstrap sample of the data before fitting. In other words, `RandomTree` is the entity we discussed in class, which makes up a random forest.

If you run `python main.py -q 4` it will fit a deep `DecisionTree` using the information gain splitting criterion. You will notice that the model overfits badly.

1. Using the provided code, evaluate the `RandomTree` model of unlimited depth. Why doesn’t the random tree model have a training error of 0?

Answer: Even if you get 0 training error on the bootstrap sample that the random tree is being trained on, this likely does not correspond to 0 training error on the original training set.

2. For `RandomTree`, if you set the `max_depth` value to `np.inf`, why do the training functions terminate instead of making an infinite number of splitting rules?

Answer: At some point, we reach one of the termination criteria for every stump: (a) we have a single class left in the label vector or (b) we have only a single example left.

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<sup>1</sup>The notation  $\lfloor x \rfloor$  means the “floor” of  $x$ , or “ $x$  rounded down”. You can compute this with `np.floor(x)` or `math.floor(x)`.

3. Complete creating the `RandomForest` class in `random_tree.py`. This class takes in hyperparameters `num_trees` and `max_depth` and fits `num_trees` random trees each with maximum depth `max_depth`. For prediction, have all trees predict and then take the mode. Submit this code.
4. Using 50 trees, and a max depth of  $\infty$ , report the training and testing error. Compare this to what we got with a single `DecisionTree` and with a single `RandomTree`. Are the results what you expected? Discuss.

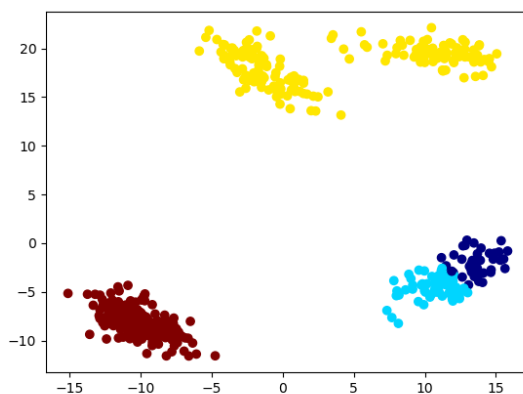
Answer: Train error is 0, test error is 0.163. This is better test error than a single tree, as expected.

5. Why does a random forest typically have a training error of 0, even though random trees typically have a training error greater than 0?

Answer: Each tree with infinite max depth will perfectly classify whatever data used for actual fitting, i.e. the tree-specific bootstrapped data. If each training example is in more than half the trees, there will be majority correct predictions across trees in the forest. Then the forest will predict the correct label for each training example.

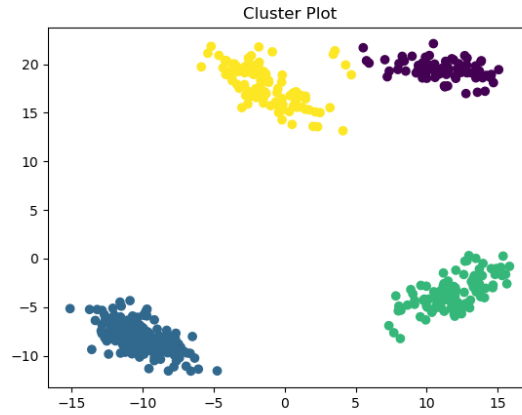
## 5 Clustering

If you run `python main.py -q 5`, it will load a dataset with two features and a very obvious clustering structure. It will then apply the  $k$ -means algorithm with a random initialization. The result of applying the algorithm will thus depend on the randomization, but a typical run might look like this:



(Note that the colours are arbitrary – this is the label switching issue.) But the ‘correct’ clustering (that was used to make the data) is this:





## 5.1 Selecting among $k$ -means Initializations

If you run the demo several times, it will find different clusterings. To select among clusterings for a *fixed* value of  $k$ , one strategy is to minimize the sum of squared distances between examples  $x_i$  and their means  $w_{y_i}$ ,

$$f(w_1, w_2, \dots, w_k, y_1, y_2, \dots, y_n) = \sum_{i=1}^n \|x_i - w_{y_i}\|_2^2 = \sum_{i=1}^n \sum_{j=1}^d (x_{ij} - w_{y_i j})^2.$$

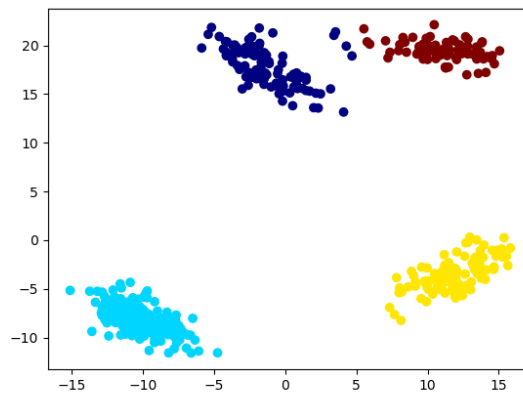
where  $y_i$  is the index of the closest mean to  $x_i$ . This is a natural criterion because the steps of  $k$ -means alternately optimize this objective function in terms of the  $w_c$  and the  $y_i$  values.

1. In the `kmeans.py` file, complete the `error()` method. `error()` takes as input the data used in fit (`X`), the indices of each examples' nearest mean (`y`), and the current value of means (`means`). It returns the value of this above objective function. Submit this code. What trend do you observe if you print the value of this error after each iteration of the  $k$ -means algorithm?

Answer: See code. It decreases monotonically.

2. Run  $k$ -means 50 times (with  $k = 4$ ) and take the one with the lowest error. Report the lowest error obtained. Visualize the clustering obtained by this model. Submit your plot.

Answer: I get 3071 as error. The clustering tends to be correct.



## 5.2 Selecting $k$ in $k$ -means

We now turn to the task of choosing the number of clusters  $k$ .

1. Explain why we should not choose  $k$  by taking the value that minimizes the **error** value.

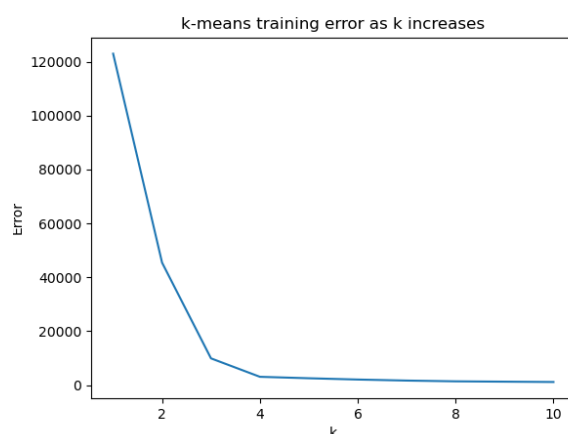
Answer: The error value always prefers the largest possible  $k$ , sort of like choosing the depth of a decision tree with training error.

2. Explain why even evaluating the **error** function on test data still wouldn't be a suitable approach to choosing  $k$ .

Answer: Since you have more clusters as  $k$  increases, the closest mean is still likely to be closer on new data with large values of  $k$ . So even on test data this objective function would likely prefer the largest value of  $k$ .

3. Hand in a plot of the minimum error found across 50 random initializations, as a function of  $k$ , taking  $k$  from 1 to 10.

Answer:



4. The *elbow method* for choosing  $k$  consists of looking at the above plot and visually trying to choose the  $k$  that makes the sharpest “elbow” (the biggest change in slope). What values of  $k$  might be reasonable according to this method? Note: there is not a single correct answer here; it is somewhat open to interpretation and there is a range of reasonable answers.

Answer: This will change based on a person's interpretation, but reasonable value might be 3 or 4.

## 6 Very-Short Answer Questions

Write a short one or two sentence answer to each of the questions below. Make sure your answer is clear and concise.

1. What is a reason that the the data may not be IID in the email spam filtering example from lecture?

Answer: Many answers are possible. For example, the spammers might modify their e-mail over time (in response to getting filtered out).

2. Why can't we (typically) use the training error to select a hyper-parameter?

Answer: Hyper-parameters (typically) control model complexity, and more complex models (typically) have lower training error (so this just leads to picking the most complex model we try rather than one

that is likely to have a small test error).

3. What is the effect of  $n$  on the optimization bias (assuming we use a parametric model).

Answer: As  $n$  increase the optimization bias decreases.

4. What is an advantage and a disadvantage of using a large  $k$  value in  $k$ -fold cross-validation?

Answer: Large  $k$  values let you use more data (to fit the model), but are more expensive (since you need to fit  $k$  models).

5. Recall that false positive in binary classification means  $\hat{y}_i = 1$  while  $\tilde{y}_i = 0$ . Give an example of when increasing false positives is an acceptable risk.

Answer: Many answers are possible. In general, when we really don't want false negative, e.g. COVID-19 test, detecting cancer, etc.

6. Why can we ignore  $p(x_i)$  when we use naive Bayes?

Answer: It's the same for all classes, so doesn't affect our decision.

7. For each of the three values below in a naive Bayes model, say whether it's a parameter or a hyper-parameter:

- (a) Our estimate of  $p(y_i)$  for some  $y_i$ .
- (b) Our estimate of  $p(x_{ij} | y_i)$  for some  $x_{ij}$  and  $y_i$ .
- (c) The value  $\beta$  in Laplace smoothing.

Answer: The probabilities are parameters and  $\beta$  is a hyper-parameter.

8. Suppose we want to classify whether segments of raw audio represent words or not. What is an easy way to make our classifier invariant to small translations of the raw audio?

Answer: Add versions of the training data with small translations.

9. Both supervised learning and clustering models take in an input  $x_i$  and produce a label  $y_i$ . What is the key difference?

Answer: In supervised learning we're given specific  $y_i$  values during training (so they have a meaning in the real world).

10. Suppose you chose  $k$  in  $k$ -means clustering (using the squared distances to examples) from a validation set instead of a training set. Would this work better than using the training set (which just chooses the largest value of  $k$ )?

Answer: Nope, still chooses the largest value (the distances get smaller as  $k$  increases).

11. In  $k$ -means clustering the clusters are guaranteed to be convex regions. Are the areas that are given the same label by KNN also convex?

Answer: No (you could have two points with the same label that have other points in between with different labels).