CS-345/M45 Lab Class 3 Total Marks: 5

This lab is about utilizing Linear Regression for continuous value prediction. It also looks at Principal Component Analysis and Linear Discriminant Analysis for feature dimension reduction. In this lab we also introduce the concept on training/testing/validation splits, and we will be training these algorithms on some example training data, before then carrying out the methods on a testing dataset. Packages used in this lab are numpy, matplotlib and scikit-learn.

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☐ Task 3.1 – Linear Regression with sklearn

This first task involves using Linear Regression to fit a predictive regression model to a collection of observed datapoints. To do this you will need to utilise LinearRegression class from within sklearn.linear_model. You are provided with the data files on Canvas.

 $x_values.npy$ corresponds to the regular variable, the magnitude along the x-axis. $y_values.npy$ corresponds to the target variable, i.e. the measurement we want to try and predict along the y-axis. In our Linear Regression model we will use $x_values.npy$ and $y_values.npy$ to train our model. Then, for a previously unseen value, we want to predict a corresponding y value.

We will first load in our data and visualise x values against y values. We then initialise a LinearRegression model and train it on the observed data. Linear Regression is a supervised approach, therefore you need to pass both the data (x) and the targets (y) when fitting the model. We will then predict an output \hat{y} value for an unseen input value.

- Load x_values.npy and y_values.npy, and visualise the data with a scatter plot.
- Create a LinearRegression object from sklearn and fit the model to the data.
- Load test_x_values.npy and predict the \hat{y} values for each of the x values it contains.
- Plot the training data x and y values using a scatter plot. On the same figure, plot the $test_x$ and predicted \hat{y} values using matplotlib's plot method (rather than scatter).
- Predict and print the \hat{y} value for x = 0.48. Note that the expected shape of input to the predict method is a matrix, not a scalar. Consider how to enter this value in the correct shape for the method. Note that reading the error message will help.

At this point in task 3.1, you should have something similar to Figure 1:

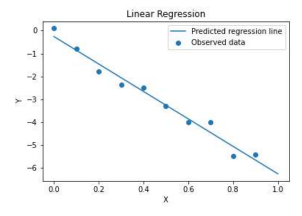


Figure 1: Visualisation of the linear regression model fit to the observed data.

□ Task 3.2 – Principle Component Analysis with sklearn

The second task is to use Principal Component Analysis to reduce the dimensionality of the Wine Dataset. Overall, we will: divide the dataset into a training and testing set, utilise a sklearn.decomposition.PCA object to reduce the dimensionality to two principle components, and then visualise the data in the new principle component space. You are provided with the data in the form of two numpy arrays, wineData.npy and wineLabels.npy, on Canvas. Download the files to your local directory.

- Load wineData.npy and wineLabels.npy, inspect the data, and visualise it. Use markdown to describe the data and it's attributes.
- Divide the data and labels into two sets: training and testing. Use slicing here to select some proportion of the data and assign them into two separate variables train_data and test_data. Use the same indices to slice your labels and create train_labels and test_labels, you need to make sure that the data and labels still match. A common split in literature is 80% training to 20% testing.
- Produce a scatter plot which shows your train:test split. For example, plot the training data with circle markers and the testing data with cross markers. Remember to colour them based on their respective labels. Plot feature 0 against feature 1.
- Initialise a PCA object, with an input argument to the constructor which will only keep the first 2 principle components. This will reduce dimensionality from 13 features, to 2.
- Fit the PCA model to the training data.
- Apply the dimensionality reduction transform to the training data. This takes in our 13-dimensional data and reduces it down to 2 dimensions. Hint: Check API for transform().
- Apply the dimensionality reduction transform to the testing data.
- Visualise the reduced-dimensionality training and testing data, using a scatter plot. Don't forget to use different markers for the two sets, and to colour the markers with the labels.

At this point in task 3.2, you should have something similar to Figure 2:

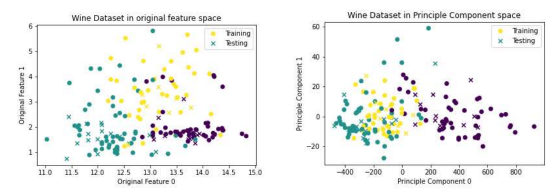


Figure 2: Left: Original feature space of the wine dataset, colored by class label. Right: first two principle components of the wine dataset.

☐ Task 3.3 – Linear Discriminant Analysis with sklearn

The third task is to use Linear Discriminant Analysis to reduce the dimensionality of the Wine Dataset. This time we will be using a supervised technique to reduce our dimensionality. In this task you will use the same train:test split you have identified in task 3.2, i.e. train_data, test_data, train_labels, and test_labels. LDA can also be used for class prediction, and has a predict method, but we will be using it for dimensionality reduction in this task by calling the transform method.

- Create a sklearn.discriminant_analysis.LinearDiscriminantAnalysis object. Again we want to provide an argument to the constructor which will only keep the first 2 components.
- Fit the model to the training data and training labels. Notice that unlike with PCA, we now need to provide the class labels for our training data (why?).
- Apply the dimensionality reduction transform to the training data.
- Apply the dimensionality reduction transform to the testing data.
- Visualise the reduced-dimensionality training and testing data, using a scatter plot. Don't forget to use different markers for the two sets, and to colour the markers based on the labels.

At this point in task 3.3, you should have something similar to Figure 3:

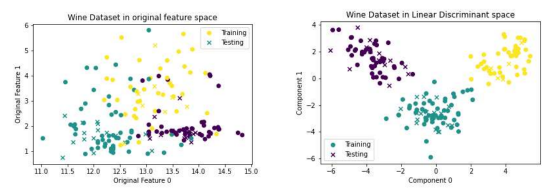


Figure 3: Left: Original feature space of the wine dataset. Right: First two linear discriminant analysis components of the wine dataset.

Use markdown cells to describe your understanding of the differences seen between PCA and LDA in these previous two tasks.

□ Task 3.4 – Principal Component Analysis by hand

The fourth task is to implement Principal Component Analysis by hand. We will perform the Singular Value Decomposition and project the data into the new principal component space, before visualising the data in the reduced dimensionality space. This task is more involved, and requires you to think about the equations presented and how the implementation of the method corresponds to the theory.

In this task you will use the same train:test split you have identified in task 3.2, i.e. train_data, test_data, train_labels, and test_labels

- Mean-centre the training data. To do this, you should identify the mean vector of the training data, and subtract that vector from the samples in the training data. You should save this mean vector, you will need it later for centring the test data.
- Calculate the Singular Value Decomposition of the mean-centred training data. You can do this with numpy.linalg.svd(), which returns the variables u, s and vh. The values returned by numpy's svd are analogous to U, S and V^T on slide 22 of the Dimensionality Reduction lecture. They form the linear system:

$$X = USV^T \tag{4}$$

where X is our mean-centred data, U is the left-singular vectors, S the singular values, and V the right-singular vectors.

• Project your data into a 2-dimensional Principle Component space. To do this you will need to select the first few significant eigenvectors of V^T (see slide 23 for reasoning) and project our training data through this matrix with a matrix multiplication. You should be doing something along the lines of:

```
projection_matrix = vh[slice indices]
projected_train_data = centred_train_data @ np.transpose(projection_matrix)
```

The slicing into vh selects our top significant eigenvectors, but why do we need the transpose? How can we tell that our dimensionality has been reduced?

- Now that you have projected your training data into the principle component space, we will project the testing data. To do this we need to mean-centre the test data using the mean vector from the training data (why?), and we need to use the same projection matrix from before (why?).
- Visualise your projected training and testing data with a scatter plot. It should look similar (but maybe slightly different) to the output from Task 3.2.

\square Task 3.5 – Apply your skill.

On Canvas there is the numpy datafile task3_5_data.npy. Download this file to your current working directory. Using the skills from the previous tasks, see if you can find the **minimum** number of principle components to keep whilst retaining at least 90% of the variance in the data. Hint: linear regression isn't needed here. We also provide task3_5_labels.npy in case you want to color the plots with the labels.

\Box Challenge Task 3.6

Some questions to consider:

- 1. Why do we not compute a projection matrix and mean value for the testing sets?
- 2. Why does LDA give us nice distinct clusters for our Wine Dataset when PCA does not?
- 3. What benefit does dimensionality reduction provide? What are the drawbacks?
- 4. How could you use LDA to predict class labels?