Enron Submission Free-Response Questions

Question 1: Summarize for us the goal of this project and how machine learning is useful in trying to accomplish it. As part of your answer, give some background on the dataset and how it can be used to answer the project question. Were there any outliers in the data when you got it, and how did you handle those? [relevant rubric items: "data exploration", "outlier investigation"]

Answer: In this project, we aim to use machine learning to identify Enron employees who may have committed fraud based on the the Enron financial and email dataset. The dataset itself contains the details of **146** Enron employees, **18** of whom are persons of interest (POIs). Each employee is associated with **21** features, including salary, bonus, total stock value, and so on. From these features, we hope to be able to extract several useful ones, and use them to identify the POIs in the dataset.

The goal of this project is simple. We need to find the POIs out of many employees at Enron. We have wide range of variable of each employee and out of these we need to also find that which variables contributes most to the POI detection. Only a machine learning approach would be able to decide if the person is an POI or not because this is impossible for human eyes to find the pattern into the data. The dataset is very biased as it should be for all the financial frauds. Hence prediction for a machine learning algorithm is also not straight forward. We need a lot more feature engineering and data munging before we can pass it to the classifier. There is also a lot of hyper parameter tuning. While data was being explored, one outlier was found which is removed from Features base class. All the basic exploratory data analysis was given in eda.ipynb file.

The Enron dataset is not perfect. Firstly, many features, especially financial-related ones, have a lot of missing values. Removing observations with missing values is unwise because the full dataset itself is already small. Hence, what I did was to replace all **NaN** values with **zeros**. Secondly, there are two outliers in the dataset, namely "**Total**" and "**The Travel Agency in the Park**". Since both of them are not Enron employees, these two rows are irrelevant and were removed from the dataset. I recognised this while plotting the salary for each employee. We can see that one point is a lot more than others. With further investigation, I found that the name associated with this value is **TOTAL**. All other observations, which are Enron employees, are retained in the dataset.

Question 2: What features did you end up using in your POI identifier, and what selection process did you use to pick them? Did you have to do any scaling? Why or why not? As part of the assignment, you should attempt to engineer your own feature that does not come ready-made in the dataset -- explain what feature you tried to make, and the rationale behind it. (You do not necessarily have to use it in the final analysis, only engineer and test it.) In your feature selection step, if you used an algorithm like a decision tree, please also give the feature importance of the features that you use, and if you used an automated feature selection function like **SelectKBest**, please report the feature scores and reasons for your choice of parameter values. [relevant rubric items: "create new features", "intelligently select features", "properly scale features"]

Answer: There were two steps for feature selection. One is during classifier testing and the other is during fine tuning of the classifiers. During the classifier testing phase, 3 different types of feature selection is used. These feature selection is not done manually but rather using feature importance scores from classifiers like xgboost and random forest and feature selection methods like SelectKBest. XGBoost with cross validation was also used to using GridSearchCV with an evaluation metric for F1 score. These feature selection was pretty successful because even without any classifier tuning, a score of more than the prescribed score of 0.3 was achieved for almost every classifier for precision and recall. During fine tuning we have also used PCA and SelectKBest with cross validation with different types of scaling like MaxAbsScaler(), StandardScaler(), MinMaxScaler(). However it is worth nothing that most of the cases, MaxAbsScaler() performed best. 5 features of the following

The reasons for creating these features were mentioned in the actual code but to put in brief

poi_interaction - If a person receives and sends a lot of emails from and to the POIs, chances are that they are POI themselves income_ratio - Shows the ratio of an employee's total income standardized by the total payments. Lesser the value, more mismatch in his/her income and more chances of being POI

expenses_std, deferral_payments_std, other_std different payments standardized by the total payments.

Feature Rank	Features	Feature Score Random Forest
1	deferred_income	0.121053
2	poi_interaction	0.120613
3	restricted_stock	0.113415
4	income_ratio	0.111179
5	shared_receipt_with_poi	0.103416

Feature Rank	Features	Feature Score XGBoost without CV
1	other_std	579
2	restricted_stock	425
3	poi_interaction	394
4	shared_receipt_with_poi	346
5	expenses_std	335

Feature Rank	Features	Feature Score SelectKBest
1	poi_interaction	11.880088
2	total_stock_value	10.814634
3	shared_receipt_with_poi	10.669737
4	income_ratio	10.368613
5	exercised_stock_options	9.956167

Feature Rank	Features	Feature Score XGBoost with CV
1	poi_interaction	45
2	other_std	43
3	shared_receipt_with_poi	42
4	expenses_std	38
5	deferred_income	37

As we can see almost all the feature selection algorithms that I have used are giving at least of the features in top 5 list. To put that in more perspective, let's see how this feature creation and feature selection affects the precision and recall score.

For example,

Naïve Bayes without feature creation and without feature selection gives

Accuracy: 0.74700 Precision: 0.23548 Recall: 0.39950

Naïve Bayes with feature creation and with feature selection gives

Accuracy: 0.86740 Precision: 0.50479 Recall: 0.29000

As we can see that except the **recall** score all the other score improves with feature creation and feature selection.

One more example is with my best classifier NearestCentroid

NearestCentroid without feature creation and without feature selection gives

Accuracy: 0.83340 Precision: 0.35349 Recall: 0.30100 F1: 0.32514 F2: 0.31021

While **NearestCentroid** with feature creation and with feature selection gives

Accuracy: 0.77220 Precision: 0.31390 Recall: 0.59750 F1: 0.41157 F2: 0.50606

We can see that with compromising **Precision** a little bit, we gained a lot in other scores after feature creation and feature selection.

Further level of feature selection was done using cross validation with different types of scalars, feature selection and dimensionality reduction and with different parameter tuning for **NearestCentroid** and we received a score like below

Accuracy: 0.82367 Precision: 0.40778 Recall: 0.71300 F1: 0.51883 F2: 0.62016

We can see that we improved in every score after feature scaling, feature creation and feature selection.

Look at **eda.ipynb** for feature importance plots.

Question 3: What algorithm did you end up using? What other one(s) did you try? How did model performance differ between algorithms? [relevant rubric item: "pick an algorithm"]

Answer: I ended up using NearestCentroid(), a different type of KNN. This gave the best performance for both F1 and F2 score. I have also tried SVC with class penalty. This is because SVC assumes an unbiased class and this was not an unbiased class. Naïve Bayes was also used with PCA and cross validation. Because PCA would reduce the assumption that the features are related and keeps the Naïve Bayes as Independent as possible. The other algorithms that I tried is

- A. GradientBoostingClassifier
- B. AdaBoostClassifier

They perform poorly. For more info look at the **README.md** file.

Question 4: What does it mean to tune the parameters of an algorithm, and what can happen if you don't do this well? How did you tune the parameters of your particular algorithm? What parameters did you tune? (Some algorithms do not have parameters that you need to tune -- if this is the case for the one you picked, identify and briefly explain how you would have done it for the model that was not your final choice or a different model that does utilize parameter tuning, e.g. a decision tree classifier). [relevant rubric items: "discuss parameter tuning", "tune the algorithm"]

Answer: Tuning the parameters means adjusting the parameters of an algorithm so that it can handle a particular dataset better. Different parameter settings will result in different decision boundaries. If we don't tune the parameters well, the algorithm won't

decision boundaries. If we don't tune the parameters well, the algorithm won't be

able to generalize a dataset well and the final classification result might be less accurate.

While exploring for algorithms, I used **GridSearchCV** for parameter tuning. For

example, for **NearestCentroid**, I tried tuning two parameters: **shrink_threshold** [None, 0.1, 0.6, 0.7, 0.8, 0.9, 1, 2, 5, 10] **metric**': ["euclidean", "manhattan"]. With different number of features used with **PCA** and **KBest** and with different number of scalars and tuned the classifier for precision.

Question 5: What is validation, and what's a classic mistake you can make if you do it wrong? How did you validate your analysis? [relevant rubric items: "discuss validation", "validation strategy"]

Answer: Validation is a way to assess the performance of a machine learning algorithm using the given dataset. This is done by splitting a dataset into training and testing

datasets, and comparing our machine learning results with the labels in the testing

dataset. A classic mistake is to use all observations available in a dataset to train the

machine learning algorithm, and then to end up overfitting the given dataset. To assess the performance of my final algorithm, I used the Stratified Shuffle Split

cross-validation. This is because the Enron dataset is small and unbalanced [many

more non-POIs than POIs). Stratified shuffle split cross-validation could construct

new instances from the dataset to ensure better representation of POI class in both

training and testing datasets. Then, it does a randomized k-fold cross validation on

the dataset.

Question 6: Give at least 2 evaluation metrics and your average performance for each of them. Explain an interpretation of your metrics that says something human-understandable about your algorithm's performance. [relevant rubric item: "usage of evaluation metrics"]

Answer: For my final algorithm choice, **NearestCentroid**, the average performance is as follows:

Precision: 0.40778

This means that out of those identified as POIs by the model, only 40.8% are true POIs

• Recall: 0.71300

This means that the model is only able to point out 71.3% of POIs accurately.

• F1: 0.51883

This is a weighted average of true positives, false positives and false negatives scores. Higher F1 score means better prediction. My score was 51.8%

F2: 0.62016

This is a weighted average of precision and recall scores. Higher F1 score indicates higher precision and recall scores, and hence a better model. My score was 62%