

Table 1. Sample Data Set By Protein Class

Cytosolic	Nucleic	Secreted	Mitochondrial
row4	row4	row4	row4

of these residues form a helix, although generally these consist of alternating positively charged and hydrophobic amino acids.

3. Into The Nucleus

Proteins that require importing into the nucleus often contain a *Nuclear Localization Sequence*, which may consist of a sequence amino acids that are positively charged (often arginine and lysine) exposed on the surface of the protein. As a result these proteins can be anywhere in the chain.

4. Out Of the Nucleus (Into The Cytoplasm)

Proteins that are being exported out of the nucleus and into the cytoplasm are often tagged with a *Nuclear Export Signal*. Often these peptide sequences consist of hydrophobic residues (such as leucine), interspersed with other amino acids.

2.2 Bagging & Boosting

This investigation compares two machine learning models to a benchmark random classifier, selecting the best. These two models are examples of common machine learning technique.

2.2.1 Random Forests

This is a sentence about random forests, how they work and such. They are quite clever and I might even have a maths equations about them and how clever they are.

2.2.2 AdaBoost

This is a sentence about boosting. I'm not fully clear on what that is but when I know I'll be sure to put it here - with an equation!

3 Method

3.1 Preprocessing

Data was obtained from four files containing protein sequences in Fasta format. Each file contained proteins according to its type of subcellular localization: cytosolic, secreted, nuclear and mitochondrial. The number of proteins in each file is presented in Table 1. These data files contained no homologues, and therefore could be assimilated, shuffled and split into a training and validation set randomly. This was done, and a training-validation split of 70%-30% was set.

The data was then fed through a pipeline to extract a number of features from the sequence. These features all corresponded to floating point numbers, and so were concatenated to form a 75-dimensional feature vector, that would be used to train the models. An analysis of the features themselves is present in section 4.

3.2 Features

The open source 'BioPython' *reference* module was used to extract features from the sequence data, allowing us to augment raw residue sequence data with important characteristics of the amino acids, such as isoelectronic point or molecular weight. In total 75 features were created, although many of these could be grouped into the same category. These features were:

- **Protein Length** - An integer representing the length of the protein sequence, in residues.
- **Amino Acid Composition** - A float created for each amino acid, corresponding to the percentage composition of that amino acid in the protein.
- **Aromaticity** - A corresponding to a measure of the aromaticity of the molecule according to Lobry et al/1994. This is measured by adding the frequencies of residues Phenylalanine, Tryptophan and Tyrosine, in the sequence of the protein.
- **Isoelectric Point** - A float representing the isoelectric point of the protein according to values taken by Bjellqvist et al ref.
- **Molecular Weight** - A float simply molecular weight of the protein.
- **Secondary Structure Features** - Three separate features were created indicating the fraction of amino acids which tend to be either α -helixes, β -sheets, or turns/loops. These amino acids were: α - Val, Ile, Tyr, Phe, Trp, Leu; β - Glu, Met, Ala, Leu; *loops* - Asn, Pro, Gly, Ser
- **Start/End Amino Acid Composition** - Two floats were created for each amino acid, according to the percentage composition of that amino acid in the first and last 20 residues of the protein.

These features were chosen as they had a high impact on the on the remaining the accuracy of the classifier. The importance of each of these features to various classifications is examined in section 4.

3.3 Classifier Training & Tuning

A number of classifiers were trained to investigate the best possible model in predicting the subcellular location. These models were:

- Logistic Regression
- Random Forests Classifier - MultiClass
- Random Forests Classifier - One-Vs-Rest (4 Models)
- Adaboost Classifier

All classifiers were implemented using the open source 'scikit-learn' package. In all cases, model parameters were fit with a 5-fold cross validation. Hyperparameters were tuned using a grid search **plot**, and choosing the parameters that had the best 5-fold cross validation score of accuracy.

In some cases, (Logistic Regression & Random Forests (One-Vs-Rest)), multiple models were trained, each solely to predict one class. In these cases, 'scikit-learn' provided the best means for ensembling these methods, using it's OneVsRestClassifier, to provide estimates of scores.

3.4 Benchmarks

A raw, bottom level benchmark for this investigation was a random classifier. This classifier was simply an implementation of a categorical distribution, with the probability vector learnt from the frequency of each category in the training dataset.

$$x \sim \text{Categorical}(\mathbf{p}) \text{ where } \mathbf{p} = [0.25, 0.25, 0.25, 0.25]$$

3.5 Ablation Study & Feature Evaluation

To investigate and evaluate the performance of certain features, an ablation study was performed on the each of the above categories. An ablation study was performed, by removing one feature at a time and testing the 5-fold cross validation score of the model in predicting these data. The results are presented *fig xyz*

4 Results

4.1 Final Hyperparameter

For each model, grid search found the following hyperparameters provided the best test cross validation score.

table hyper params

Table 2. Hyperparameters of Model

Model	Hyperparameter	Settings
Categorical (Benchmark)	N.A.	N.A.
Random Forests Classifier (MC)	estimators	450
Random Forests Classifier (OvM)	estimators	450
Adaboost Classifier	estimators	200
Adaboost Classifier	learning rate	0.25

4.2 Model Scores

All models were evaluated on final held-out test set. Calculations were made of the accuracy, the F1 score and Area Under Curve of the Returning Operator Characteristic (AUCROC)

Table 3. Hyperparameters of Models

Model	Cross Val Acc	Test Acc	F1 Score
Categorical (Benchmark)	x	Test Acc	row1
Random Forests Classifier (MC)	y	Test Acc	row2
Random Forests Classifier (OvM)	z	Test Acc	row3
Adaboost Classifier	w	row4	row1

4.3 Plots

Finally, to investigate the importance of various features, the ROC curve was plotted, for the best model, along with the confusion matrix, and the importance weights for the random forests classifier. These are presented *here*.

4.4 Final Predictions

Given the above evaluation of the models, a Random Forest Model was selected to give a predictions of **YYY** hitherto unclassified proteins. This prediction, along with per centage confidence scores, is given in table *blah*

Table 4. Predictions of Test Proteins

Protein ID	Subcellular Location	Probabilty
row1	row1	row1
row2	row2	row2
row3	row3	row3
row4	row4	row4

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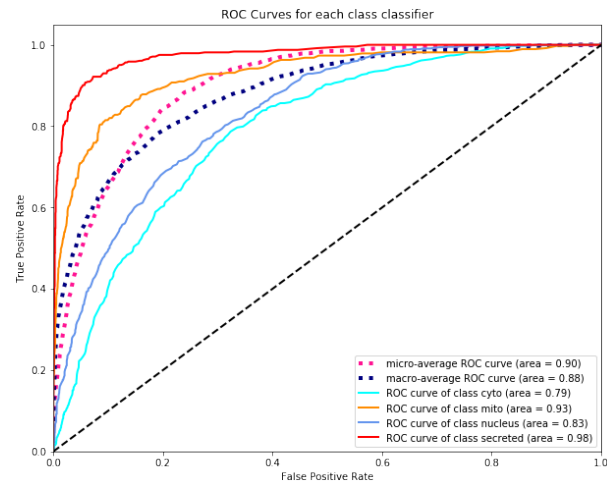
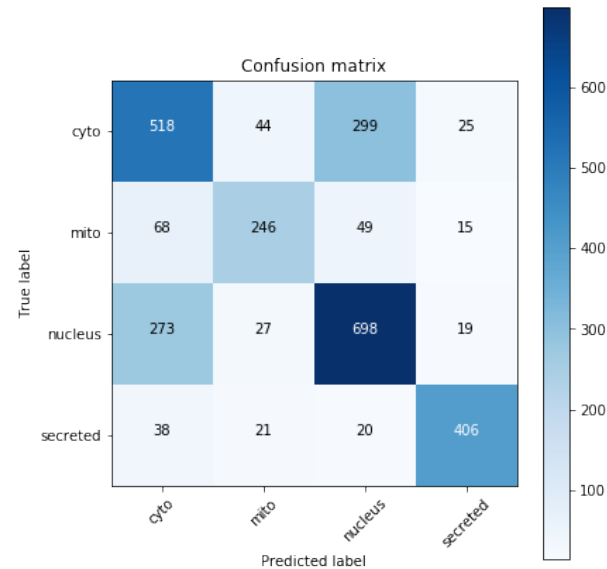
5 Discussion

5.1 Classifier Comparison

From the above results it is quite clear that both boosting and bagging methods far outperformed the benchmark method. The woeful performance of *X* accuracy is to be expected for a model that doesn't take into account any of the sequence data, or underlying features.

It is also clear that Random Forests outperforms AdaBoost, securing a decisive performance improvement of *X* compared to *Y*. In both cases this *who got more false positives, false negatives*.

This performance difference is likely down to *critique of bagging vs boosting*.



5.2 Error Analysis

In inspecting our best performing model, Random Forests, more closely, it is clear that the errors in the model were not distributed evenly across categories. Instead it appears that the greatest source error in fact come from misclassifying nuclear labels for cytosolic ones. This can be seen

clearly from the confusion matrix presented in *confusion matrix*. Almost $X\%$ of nuclear proteins are misclassified as cytosolic, and $Y\%$ as cytosolic as nuclear.

This misclassification is made all the more evident by looking at the Returning Order Characteristic (ROC) curves, where all the classifiers are plotted together. From these graphs, it is immediately clear that the best classifier is the secreted class, with the area under curve (AUC) of 0.98, followed by the mitochondrial class of 0.93. The worst performer is the cytosolic class (0.79) followed by the nuclear class (0.83).

There are a number of reasons why it is conceivable that these particular protein classifiers (cytosolic and nuclear) should perform worse than the other two categories (secreted and mitochondrial). For one thing, there are a number of proteins that frequently move between the nucleus and the cytoplasm, meaning that they may possess both *Nuclear Export Signals* and *Nuclear Localization Signals*. This mixup of very distinct features (in the form of target peptides) would likely lead the classifiers to struggle to distinguish the correct class in ambiguous cases.

Furthermore, it is likely that mitochondrial proteins and secreted proteins may have important properties that are unrelated to target peptides but still show up in our choice of features. For instance, since secreted proteins need to cross the cell membrane, it is possible features such as very large size or certain peptide make up, might make this action difficult. There may exist alternative restrictions on mitochondrial proteins, that need to be adapted to help in respiration, which might also restrict the protein's structure.

At any rate it is clear from the ROC curves in particular that secretion is particularly well detected, with *comment about false positives etc*, nuclear and cytoplasmic classes are easier to confuse. An analysis of the exact features that could be responsible for this are presented in the next section.

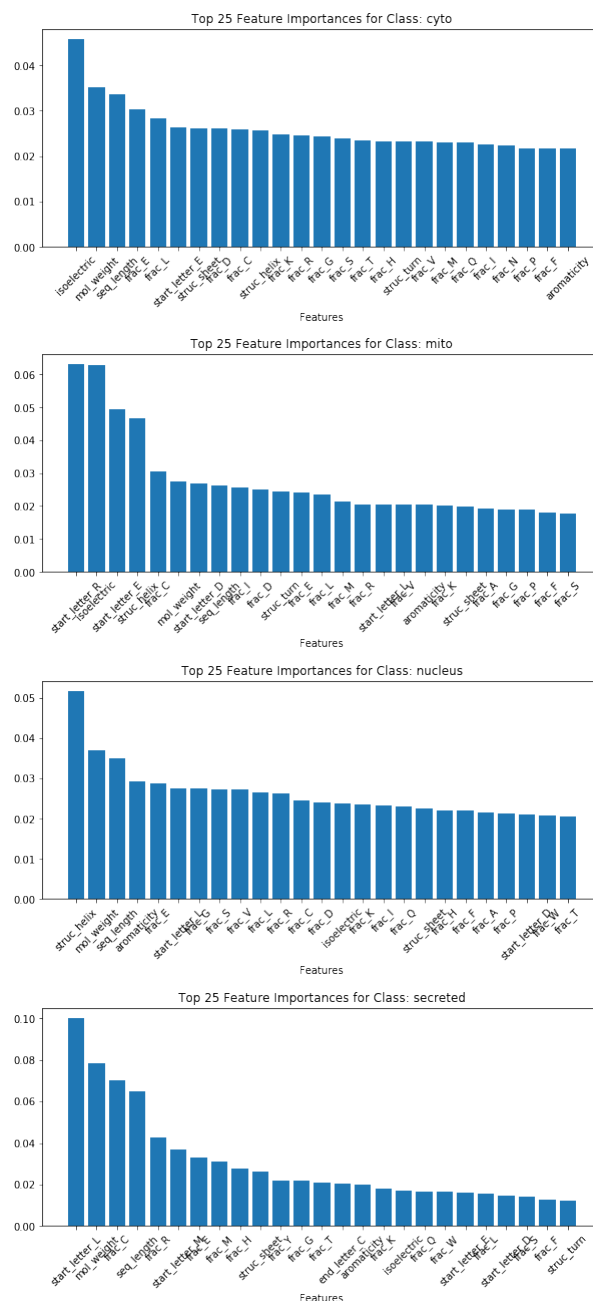
5.3 Features Analysis

Perhaps more interesting than the comparison of the two different models is the close comparison of the features that have had a significant impact on the model. Thankfully, our One-Vs-Rest Random Forest Model has an in built way of examining the features, via the feature importances. These are displayed in figure ref importances.

The first thing to notice is the relative scales of the four graphs, in comparison to one another. The secreted classifier has a much stronger set of "significant" features, than all the other classes. In the case of comparison to nucleus and cytoplasmic classes, secreted's top feature is over twice the importance of the latter classes highest feature. Furthermore its top 4 features features all have higher importances than the highest single feature for these classes. This indicates that there are distinct "killer features" for this class, and it appears they are highly tied to the fraction of Leucines and Cysteines at the start of the sequence, and the molecular weight and sequence length. **investigate data?**

Next to notice is that, as mentioned in the initial background, mitochondrial proteins also have importances supporting the target peptides alluded to. *Mitochondrial targeting signals* sometimes consist of alternating positively charged and hydrophobic amino acids forming a helix, so unsurprisingly our strongest features include features noting isoelectricity, structural helices, and the presence of positively charged Arginine at the start of the sequences.

When it comes to nuclear and cytosolic proteins, it's worth noting that the second and third best importance features are shared - indicating the difficulty these classes might have in distinguishing themselves. Furthermore these features - molecular weight and sequence length - are broad macroscopic features that do not really indicate a particular target sequence, and also show why it might be hard to classify these proteins. Even the "strongest feature" for these two classes is a macroscopic feature - isoelectric point, and percentage of "common helix" amino acids,



indicating our set of features haven't been able to pick up the nuclear target peptides.

Finally it is worth noting that none of the most important features correspond to peptides at the end of the sequence. This is fairly good evidence that something important happens at the N-terminus of a protein, and supports background theory that this is where target peptides are often found.

5.4 Further Improvements

Although this model performed well in distinguishing the secreted and mitochondrial proteins there are a number of additional features that may particularly improve future implementations of this model.

First, some measure of hydrophobicity would be a valuable additional feature. This wasn't added due to the lack of there being an available scale in the BioPython module, but it is likely this addition would help to pick

up the hydrophobicity of elements found in *either NLS or NES sections of the sequence*, helping to distinguish nuclear and cytoplasmic proteins.

yip

Furthermore it should be worth exploring and searching for specific target patterns within the model. I attempted to search for these by creating a feature for all possible "bigrams" and "trigrams" of residues, and looking at importances, but the high dimensionality made learning difficult. Instead, if one had specific examples of known target peptides this feature would likely greatly help classification, without adding too much to computational complexity.

Finally it may be worth trying gradient boosting algorithms or neural networks. Gradient boostin algorithms such as XGBoost *Explain why gradient boosting maybe better 2 lines*. Neural networks also have significant performance, but mainly on high data sets. Given the relatively small dataset (only a few *thousand points*), neural methods may not be suited to this particular small task.

6 Conclusions

Obtained data from X files, and mixed, shuffle and split into a train and validation split. Since the proteins are all non homologues, no need to arrange by family and could split anyway.

Then took these files and processed them into feature vectors, using the bipython module. These extracted, x features from the sequence.

Then trained a random forest classifier with X trees, and also tested a XGBboost classifier.

Cross validated grid search done to find best parameters for trees
Ablation study used to investigate salient features.

Trained a random forest with X trees.

What did you do? - extract features - train -test split - homologies - cross validate - compared with a model (gradient boosting?) - ablation study

- Results of accuracy - Confusion matrix - ablation study - results compared to xgboost

end of paper

7 Discussion

- Examine the features - Examine the ablation study - Examine comparison to XGBoost - Compare with existing literature and approaches

8 Further Work

- suggestions: more data? better features/

9 Conclusion

10 Theoretical Background

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