Mining protein interactions affected by mutations using a NLP based machine learning approach

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Abstract—The knowledge on the protein/gene interactions that are affected by mutations helps understand phenotype-genotype associations and predict disease prognosis and responses to treatments. Such information is scattered around in scientific literature and its manual curation is very time and resource consuming. Although much research has been done in the past to extract protein-protein interaction (PPI) information automatically from literature, much less has been done on extracting PPIs affected by mutations. An important step towards extracting such information is automatically retrieving the relevant articles. In this study, we classify abstracts as relevant or irrelevant, where being relevant means the article contains information that at least one PPI is affected by a mutation. We started out with a bag-of-words and bigrams model, and then added some frequency statistics. Interactions are modeled by triplets, where a triplet is defined as two protein names and an interaction word in a sentence. The Stanford Dependency Parser, which helps find the dependency graph for each sentence, was used to further calculate the shortest path distances for each of the three pairs of words in a triplet. We also used a model to estimate the probabilities that a triplet forms a true interaction. The shortest paths between protein name and mutation word pairs in the sentences were computed as well. Features based on all these above were used in a Gradient Boosting Trees model. Our model achieved satisfactory performance in the training data of the Biocreative VI challenge.

Keywords—Protein-protein interactions; mutations; text mining; biomedical literature retrieval; protein interactions affected by mutations

I. INTRODUCTION

The publications in biomedical literature have been increasing at an accelerated speed. Reading all the literature in a particular domain of interest is less and less feasible in recent years. And the gap between the number of papers a researcher can read and the number of all the papers he/she needs to read will only become wider over time. Searching literature using keywords at literature databases such as PubMed has been a very popular approach for finding relevant scientific articles. However, such searches can often return hundreds or even thousands of papers, many of which are not highly relevant to the topics a user aims to find. Biomedical literature triage (or retrieval) has been a popular topic in recent years [1-12]. Most of the methods use machine learning methods to retrieve relevant articles related to a particular topic. Literature triage tasks vary widely depending on the subdomain of the literature and also on the particular information one aims to retrieve.

In this study, we address the problem raised in the Biocreative challenge VI track 4, mining protein interactions and mutations for precision medicine. This challenge consists of two subtasks: (1) Document triage: identify relevant PubMed citations describing genetic mutations affecting protein-protein interactions; and (2) Relation extraction: extract experimentally verified protein-protein interactions (PPI) affected by the presence of a genetic mutation. Much research has been done in the past to extract protein-protein interaction (PPI) information automatically from literature [13-24].Extracting PPIs affected by mutations is very challenging in that not only one needs to deal with information on PPIs, but also on whether they are affected by mutations. The information on whether a PPI is affected by mutations often occurs in sentences different from the one where PPIs are mentioned, which makes the tasks more challenging.

For document triage task, the training dataset consists of a set of ~4K PubMed articles. These articles are manually labelled as relevant/not relevant by BioGRID database [25] curators. The goal is to build automatic methods capable of receiving a list of PMIDs and return a relevance-ranked judgement of the test set for triage purposes.

For relation extraction task, a subset of the relevant articles in document triage task was manually annotated with relevant interacting protein pairs. Each PubMed article in this set has at least one interacting pair which is listed with the Gene Entrez ID of the two interactors. These protein-protein interactions have been experimentally verified and the analysis of natural occurring or synthetic mutations has identified protein residues crucial for the interaction. The goal is to build automated methods that are capable of receiving a set of PMID documents and return the set of interacting protein pairs (and their corresponding Gene Entrez IDs) mentioned in the text that are affected by a genetic mutation.

The validity of the text mining methods is evaluated using standard metrics such as average precision, f-measure, etc. In addition, we also used Area Under the Curve (AUC) to evaluate different models.

In the rest of the paper, we will first describe the method we developed for both tasks. We will then present our result on mainly the first task. We end the paper with conclusion and discussion.

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II. Methods

A. Data

The data used in this work are from PubMed articles. The dataset consists of titles and abstracts from 4082 papers, each manually labeled by BioGRID database curators as relevant or not relevant, for the training data, and 1500 papers for the test data.

B. Manual curation of mutation related words

Mutation related words were first manually curated from a list of words with a high TF-IDF between the true and false labels. When using cross validation to tune the model, mutation words not included in the training fold were excluded to avoid a potential information leak.

C. Feature extraction and model building

1) Triplet based modeling for protein-protein interactions

Protein-protein interactions (PPI) were modeled based on a triplet concept we developed in a previous study [26], where a triplet is defined as two protein names and an interaction word describing their interaction or relationship in the same sentence. The dictionary of interaction words were developed by a combination of manual and computational approach [26]. The method was later successfully used in knowledge discovery and other applications [27-30].

2) Bag-of-words and bigrams model

We started out with a Bag-of-words and bigrams model, and then added some frequency statistics from the title and body parts of the abstract, given that the title and the body have different information for triage. Treating the whole document as a bag, a word’s potential to become a feature for training classifiers will depend on its term frequency in each abstract compared with its occurrence through the whole document. This way the words that appear often in the whole bag will be weighted less when deciding features. Besides the features from Bag-of-words, some frequency statistics were proved to be contributive in our initial Support Vector Machine model. These include: the percentage counts of protein/gene names, mutation words and interaction words, the percentages of sentences in an abstract that have 1, 2 or 3 and above pairs of protein name and mutation word within the sentence, the percentage count of within-sentence protein name and mutation word within the sentence, the sentences in an abstract that have 1, 2 or 3 and above pairs of mutation words and interaction words, the percentages of these.

We computed the shortest paths in the dependency graphs between protein names and mutation words and added additional features based on these. For each protein word mutation word pair, we computed the shortest path between them. For each such pair, we computed the maximum probability given by our PPI extraction method over all triples containing the given protein. The probabilities were again put into bins [0 <= p < 1/3], [1/3 <= p < 2/3], [2/3 <= p <=1]. Shortest path lengths were again placed into the bins 1, 2, 3, 4-6, 7-11, 12 and above as before, then binned again according to

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<td>signaling</td>
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<td>show</td>
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<td>dependent</td>
<td>152</td>
<td>membrane</td>
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<td>dna</td>
<td>166</td>
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<td>129</td>
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<tr>
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<td>humans</td>
<td>145</td>
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<td>expression</td>
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<td>242</td>
<td>terminal</td>
<td>134</td>
<td>show that</td>
<td>119</td>
</tr>
</tbody>
</table>

3) Stanford Dependency Parser

The Stanford Dependency Parser was used to find the dependency graph for each sentence. For each triplet consisting of two protein names and an interaction word in a sentence, we computed the shortest path in the dependency graph between every pair of words in the triplet. Imagine each word is a node, each typed dependency is an edge that connects two nodes, then the parser can give the network graph of a sentence. Given two nodes, for example, Protein 1 and Protein 2, or Protein 1 and its interaction word with Protein 2, we need to find which path that connects these two nodes are the shortest. We can then obtained the Abstract-level features based on this information. For instance, the percentage counts of the triplets in an abstract that have the shortest path distances equaling 1, 2, 3, falling in the range 4 to 6, falling in the range 7 to 11, or equaling 12 and above, the percentage of sentences that have shortest path information in an abstract, etc.

We also used a model developed by another member of our group to extract true PPIs from the abstracts [26, 28, 31]. This model also uses information of the dependency graph generated by the Stanford Dependency Parser. Intuitively the shortest path features give us information to distinguish true PPIs among all the triplets. Triplets consisting of two proteins and an interaction word in the same sentence were then extracted from the abstracts and the model was used to estimate the probabilities that they form a true interaction. Therefore, by estimating the probabilities for triplets in an abstract to be true (PPIs), we added more features such as the percentage counts of the triplets with certain probability ranges. (We took the three equally sized bins [0, 1/3) (1/3, 2/3], (2/3,1].) We would then count the frequency of triplets with predicted probability falling into each bin.

The previous features give an idea of how many protein, interaction, and mutation words are contained in an abstract and an estimate for how many true PPIs are contained, but we have not yet done anything that connects the mutations to PPIs.

We computed the shortest paths in the dependency graphs between protein names and mutation words and added additional features based on these. For each protein word mutation word pair, we computed the shortest path between them. For each such pair, we computed the maximum probability given by our PPI extraction method over all triples containing the given protein. The probabilities were again put into bins [0 <= p < 1/3], [1/3 <= p < 2/3], [2/3 <= p <=1]. Shortest path lengths were again placed into the bins 1, 2, 3, 4-6, 7-11, 12 and above as before, then binned again according to
max PPI probability. For example, suppose there is a path connecting the protein tubulin with the mutation words SNP of length 8. This path will fit into the bin $[7 \leq sp \leq 11]$. If our PPI extraction model gives a maximum probability of 0.56 that a triplet containing tubulin is a true PPI then this path will go into a bin with $[7 \leq sp \leq 11]$ and $[1/3 < p < 2/3]$ since there are three probability bins and 6 shortest path length bins, there are a total of 18 bins a path length can go into. For each abstract, we take as a feature, the frequency of shortest paths lengths in each bin. For convenience we will call the probability bins: $[0 \leq p < 1/3]$ as bin 0, $[1/3 < p < 2/3]$ as bin 1, $[2/3 \leq p \leq 1]$ as bin 2.

We also employ a bag of words and bigrams approach to the words along each shortest path and to the dependencies along each path. If a path has a max PPI probability falling into bin 2, we append 2 to the end of each word in the path as follows: Tyrosines__2 recognized__2 trapped__2.

And similarly for dependencies. This shortest path had a max PPI probability falling into bin 1, so we have appended a 1 to each value. This is done so we can distinguish between paths connecting mutation words to proteins that have interactions from paths connecting to proteins that don’t have any listed interactions: Xcomp__1 dep__1 xcomp__1 acl:relcl__1 root__1 dep__1 advmod__1 acl__1.

Frequencies of protein words, interaction words, and mutation words were also computed along the shortest paths.

4) Gradient boosting trees model

We tested three classifier types, a Linear SVM, a Random Forest, and a Gradient Boosted Trees model. The first two implemented in Python’s scikit-learn. We used the popular XGBoost library[32] for gradient boosted trees. The number of words included in the bag of words and bigrams features was limited in the XGBoost model to avoid overfitting. The models were tuned using ROC-AUC scoring. The probability scores for the best model was then calibrated using Platt scaling[33]. Then cross validation was used to find a cutoff threshold to maximize the F1 score. A cutoff of $p = 0.35$ was chosen to predict abstracts as relevant. The model’s performance on the training data was reasonable, with a cross-validated ROC-AUC score of 0.793 (std 0.018) and F1 score of 0.709 (std 0.0090).

III. RESULTS

The following results were computed using ten-fold cross validation. Our baseline Linear SVM model using bag of words features and some count statistics received an ROC_AUC score of 0.7428. A random forest with 1000 estimators, min samples leaf set at 2 and a Gini splitting criterion (these parameter values were found through cross-validation) achieved an ROC-AUC score of 0.78825 with these features, giving significantly better results than the SVM. Applying a tuned XGBoost to this same set of features yielded an ROC_AUC score of 0.78686. With all features included, the best scoring model was a tuned XGBoost with an ROC_AUC score of 0.79304. In this case, the performance of the Random Forest barely improved with the addition of the new features.

When tested on test data set, our method gave an F-1 score of 0.6778, recall of 0.8877, a precision of 0.5482. We ranked the fourth among all the participating teams based on F-1 score.

IV. CONCLUSIONS AND DISCUSSIONS

In this study, we tackled the problem of retrieving abstracts which contains at least one protein-protein interactions (PPI) affected by a mutation. We developed natural language processing (NLP) based methods, extracted a set of diverse features, and experimented with several popular machine learning methods. The performance on the training data of Biocreative VI was quite satisfactory.

There is still much room for further improvement of our models, especially in terms of optimizing the set of features we will use for the machine learning models. These will be the subject of future studies.

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REFERENCES


