ZK DrugResist
Automatic Extraction of Drug Resistance Mutations and Expression Level Changes from Medline Abstracts

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Abstract: Drugs are small molecules that generally work by binding to its target which is often a protein. This ligand molecule binding helps in the treatment of various diseases. Major obstacle to treat complex diseases is the phenomena underlying drug resistance mechanisms which are not fully understood so far. Previously reported literature has mentioned few of the motives behind this complex mechanism which dominantly include protein missense mutations and the changes in the expression levels of certain genes. A better understanding of these mechanisms is getting crucial for the researchers. Retrieving information on these processes can be challenging as scientific literature has huge pool of data and extracting the required information has always been a laborious task. We developed an online pipeline ZK DrugResist that automatically extracts PubMed abstracts of drug resistance paired with either mutation or expression for a given disease. Our classifier showed 97.7% accuracy with 93.5% recall and 96.5% F-measure. This system saves plenty of time in terms of data mining and also reduces efforts in retrieving information from online resources.

1 INTRODUCTION

The term drug also referred as dose or medication is used for treatment of various diseases. There are two ways to classify drugs, one named as the small molecule drugs which include proteins, biological medicinal product and vaccines which further used as therapeutic agents for the treatment of certain diseases. The second way of classification is based on how the drug is administered that is its specific mode of action following the therapeutic effects. The drug usually functions by binding to its target which is often a protein. Proteins are large biomolecules made up of amino acids. They are also visualized as large globular structures that have deep groves in it, which may have buried binding site that is good for druggability. The drug molecule will then fits in the binding site and the process is termed as ligand-molecule binding. In this way the drug performs its action and helps diagnosing and curing various diseases (Dean et al., 2005; Michael, 2002; Walsh, 2000).

Sometimes treatment phase has been passed through an obstacle “drug resistance” generally meaning the decrease in the efficacy of the drug in curing a disease. This is the major constraint to treat complex diseases. The underlying mechanisms are not very clear but still there are some notions about it. First theory states that drugs at their certain target sites are present in a decreased concentration caused by increased level of expression of drug molecules. Second involves the modification of drug targets which affects the protein-ligand binding complex (Remy et al., 2003). Drug resistance has a strong impact on disease treatment; it has been observed that in many of the cases this brings failure in treatment. This shows that rate of survival is proportional to how strongly the mechanism of drug resistance is being overpowered. The survival chances would increase if the drug resistance could be overcome (Longley and Johnston, 2005).

The current study focuses on evaluation of complex phenomena lying behind the drug resistance mechanism. From the literature it has been found that one of the major reasons behind this is the protein alteration which involves amino acid mutations at certain residue. These missense mutations affect the binding affinity of the protein with the ligand and hence results in making drug insensitive to the
treatment. For example as reported in previous studies that V299L, T315A, and F317I/L mutations are resistant against dasatinib while mutations like Y253F/H, E255K/V, and F359C/V are resistant for nilotinib, therefore making protein mutations as an important factor for drug resistance mechanisms (Chrisanthar et al., 2008; Hochhaus et al., 2011). Second important factor is the expression based drug resistance mechanism. The changes in expression level which either is the overexpression or down-regulation of certain genes induces enhanced resistance against various drugs. As one of the studies reported the overexpression of ANP32C creates enhanced resistance against FTY720 drug, hence makes it ineffective to treatment (Buddaseth et al., 2014).

In order to retrieve and comprehend drug resistance mechanisms, researchers either has to look for the online databases or read all freely available biomedical documents through online sources, which is of course a very time consuming task. Many computational biology/bioinformatics studies have focused in building automated pipelines to extract information from PubMed abstracts. There are some databases published in literature that stores different aspects of drug and gene relationship like BacMet which focuses on genetic alterations causing resistance against antibiotics (Pal and Larssoon, 2014). Moreover there is another tool named Biozyne P-gp Predictor which is based on SVM classifier that differentiates the substrates from efflux pumps (Levatic et al., 2013). Another similar reported database is CancerDR which focuses on the identification of the altered genes encoding drug targets (Kumar et al., 2013). Retrieving information from such kind of repositories is a laborious task. Making automated way of information retrieval is one solution to this. Previously published methods just focussed on general analytical tasks like mining genes and protein names or describing relationship of genes and drug. These methods don’t emphasize on combining all these information and placing them in one platform. Some of the tools on information mining are already been published, for instance Proux research group reported the syntactic parsing methodology for information extraction developed by (Proux et al., 1998). Similarly another method used statistical based information (Hishiki et al., 1998; Ohta et al., 1997). In the same way (Cutting and Kupiec, 1992; Aronson et al., 1994; Humphreys et al., 1998) also developed servers that used semantic analysis approach for information extraction. Thorough review of the literature revealed that there is another published tool EDGAR (Rindflesch et al., 2009) that overcomes the limitations of the previously existing information retrieval methods. This tool works in building relationship between genes and drugs relevant to cancer therapy. But unfortunately this tool is not available online yet and it is also not mentioned that how much accuracy authors have achieved in applying natural language processing on the abstracts. In another study reported by (Bui et al., 2010) the authors developed the method for combining drug and mutation level information for HIV. Again this method is only specific to HIV. Our proposed method has successfully benchmarked already existing methods. ZK DrugResist uses machine learning approach to retrieve drug resistance information. It provides one platform that gathers gene names, drug names, abstracts titles, link to the abstracts categorised by disease type. Our tool provides the most systematic way of information extraction for drug resistance abstracts available on PubMed. In this way it facilitates the researchers in mining desired information more robust and more accurate. The PubMed directory considered as a rich source of information as it has a huge collection of abstracts. Despite this fact, automated mining of worthy information remains a big challenge for researchers. Our study aims to develop an online tool to automatically extract all the abstracts from PubMed related to drug resistance. These abstracts and the related information are downloaded into a database. From this all the information about the mutation, gene and the expression status is processed and displayed on web. Furthermore the abstracts are also marked as cancer or other diseases based on the content provided in the abstract. We used MugeX and EnzyMiner approach developed by our computational biology group for implementing this classifier (Erdogmus and Sezerman, 2007; Yeniterzi and Sezerman, 2009).

2 METHODOLOGY

Abstracts available online queried by using search terms: “Drug resistance”, “amino acid mutation at drug resistance level”, “expression based drug resistance” and different combinations of these terms were downloaded from Medline which are many thousands in number. Out of them only those abstracts are filtered that has either the drug resistance and the protein mutations content present together or drug resistance and the expression level information present in a document. The downloaded abstracts were passed through variety of algorithms including
tokenization/sentence splitting followed by porter stemming. These algorithms are applied in order to break down the abstract into sentences and then into words making them easy to process.

2.1 Classification Modules

We applied two learning algorithms Naive Bayes and Rocchio algorithm both uses bag of words approach. After pre-processing the dataset, we applied our classifiers which for our case are based on four levels of classification. First stage is to separate the abstracts of drug resistance from the other ones. First the document is processed by tokenization and porter stemming algorithm. Further we applied TF-IDF weighting (term frequency inverse document frequency). It is the product of two statistics Term Frequency and Inverse Document Frequency, term frequency deals with the raw calculation of a term in a document while inverse document frequency deals with the significance of a word count. We observed how many times word “Drug Resistance” appears together in a document, if it is more common we labelled the document as drug resistance else it is labelled as others. Following this the next phase classification picks the drug resistance tagged abstracts to further classify them as either mutation or expression. The regular expressions are designed for this purpose. If regular expression matches any mutation related information in the document we marked it as mutation, on the other hand it is marked as expression based if the content displays the gene expression level changes for the drug resistance again TF-IDF is used for this purpose. Third step is to subcategorize the abstracts labelled as mutations. The mutation can be at amino acid level or at the nucleotide level our tool is interested only to pick the protein mutations. Those at DNA level are termed as ambiguous mutations, so this step targets to remove ambiguous mutations from the actual ones using the regular expressions defined earlier. The last module of our classifier is to divide the cancer related articles with the ones which are showing other diseases of metabolic, autoimmune and neurodegenerative. The documents cited by terms cancer, leukaemia and tumour belongs to cancer class while the rest are classified in others category. For term frequency we used TF-IDF as mentioned before. Figure 1 summarizes all the steps involved in ZK DrugResist.

For implementation Perl Regular Expressions were used, set of patterns were formed describing the protein mutations, for instance the mutation cited as L15V, Arg567Leu, Ala399->Asp and some are mentioned as full sentences substitution of Methionine with Valine at position 40. Following the mutations the gene names parallel to mutation stated in the abstracts were also downloaded and stored in the database.

2.1.1 Drug Resistance Vs. Others

As mentioned first stage of classification is to clearly mark the abstracts which are showing drug resistance mechanism from the others which are irrelevant to these. The total abstracts downloaded are 701 in number. For each of the downloaded abstract the feature vector is constructed. In order to distinguish them the frequency of each word is counted as a feature value. These words were then further processed using tokenization and porter stemming algorithms. After breaking the abstract into words and counting the frequency of term “Drug Resistance” it is marked as either drug or others.

2.1.2 Mutation Vs. Expressions

In the second category we picked these drug resistance documents and scanned them for the mutation level information. The documents cited using overexpression down regulation kind of terms are marked as expression abstracts while those which uses amino acid terminology are marked as mutation by our algorithm.

2.1.3 Protein Vs. DNA

The Perl regular expressions were applied to extract the amino acid level mutations from each document. The major hindrance is the mixing of some protein mutations with the DNA ones. For example the one letter code amino acid mutation like A456G can easily be misinterpreted with the nucleotide letters. We compiled regular expressions for this ambiguity to be solved. The documents are classified based on the content information.

2.1.4 Cancer Vs. Others

In the last module of classification those abstracts which are associated with cancer were separated with the abstracts which are related to other diseases which include neurodegenerative, autoimmune and metabolic disorders.

2.1.5 Gene/Inhibitors

The gene names following the protein mutation are also extracted from the abstracts. For this purpose the complete list of official gene names were being
downloaded from HUGO database http://www.genenames.org/. Any gene name mentioned in the abstract is programmed to match with the list of the genes stored and the results are displayed on web. We followed MugeX approach for this module.

2.2 Implementation

All the steps are implemented in The Perl Programming Language. Strawberry Perl version 5.20 was used. The regular expressions were compiled using PERL Regular Expression library. The necessary information from the articles including Title, Abstract and PubMed ID was downloaded and stored in XML format into MySQL database. After this the documents are processed first using tokenization using Perl module PPI::Tokenizer. Each sentence is broken into words and further porter stemming was applied to each document. This algorithm is used to remove the common words from English which are actually not contributing in classification for example “the”, “is”, “are” and similar words to that. After pre-processing all the documents, they are passed through Naïve Bayes and Rocchio classifiers in order to achieve four levels classification. The database is built on Xampp Server; database tables are stored in MySQL phpmyadmin of xampp. The web interface was designed in WordPress using html and Php. CGI, DBI and DBD modules of Perl were being used for retrieving the data from MySQL databases and displaying the output on the web page.

2.3 Testing

In order to test the classification results training, testing and k fold cross validation were employed. For all the four modules of classification 20% of the abstracts were being used as test set, while remaining abstracts were considered as training set. We performed 5 fold cross validation that means the whole data is being divided into 5 sets out of which 4 are used are training sets and the rest of one is as test set. These sets are being shuffled 50 times and average accuracy for both the training and test sets were being measured.

3 RESULTS

The classifier is tested with the entire pre-processing algorithms we implemented. Out of huge pool of data available online on drug resistance mechanisms we only filtered those which are showing either mutation level or expression level changes in causing drug resistance. This makes up to 701 documents in total. The first module of classification separates out 144 documents as drugs while the other 557 are the ones in which mutation is mentioned but not at drug level. These 144 documents are advanced to the second level of classification. This shows that out of 144 documents, 91 are those belonging to mutation category, 22 of them are the expression based resistance abstracts while rest of 31 did not show either the mutation or the expression based drug resistance. These 91 abstracts are then picked to distinguish the protein and DNA level mutations from each other. The results showed that 65 of them are the ones which are being labelled as amino acid mutations while 25 are the other ones. Last module of classification separates the cancer disease ones with the other diseases. The calculation shows that 53 are the drug resistance mutations at cancer level while 12 are the ones which are in other category. We compared the results of our classifiers and the results showed that Naïve Bayes classification outperformed Rocchio algorithm in precision, recall and accuracy as shown in Table1. The results of Naïve Bayes classifier are listed in Table 2, 3, 4 and 5 respectively. The graphical representation is illustrated in Figure 1.

| Table 1: Comparison of Naïve Bayes and Rocchio Algorithm. |
|-----------------|-----------------|-----------------|-----------------|
| Naive Bayes Classifier | Rocchio Algorithm |                  |
| Accuracy | Recall | Precision | Accuracy | Recall | Precision |
| 97%    | 96.5% | 95.9% | 90.5% | 83.0% | 89.4% |

| Table 2: Classification Results of Drug Resistance vs. Others in Training and Test Sets. |
|-----------------|-----------------|-----------------|-----------------|
| No of Abstracts | Accuracy Measure | Recall | F-measure |
| 660              | 96.4%           | 95.4% | 93.7%    |
| 140              | 96.7%           | 96.5% | 95.9%    |
### Table 3: Classification Results of Mutations vs. Expressions in Training and Test Sets.

<table>
<thead>
<tr>
<th>No of Abstracts</th>
<th>Accuracy Measure</th>
<th>Recall</th>
<th>Precision</th>
</tr>
</thead>
<tbody>
<tr>
<td>115</td>
<td>96%</td>
<td>95.4%</td>
<td>93%</td>
</tr>
<tr>
<td>28</td>
<td>97%</td>
<td>96.5%</td>
<td>95.9%</td>
</tr>
</tbody>
</table>

### Table 4: Classification Results of Protein Mutations vs. DNA Mutations in Training and Test Sets.

<table>
<thead>
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<th>Recall</th>
<th>Precision</th>
</tr>
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<td>94%</td>
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<td>18</td>
<td>96.7%</td>
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<td>95.9%</td>
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</table>

### Table 5: Classification Results of Cancer vs. Others in Training and Test Sets.

<table>
<thead>
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<th>No of Abstracts</th>
<th>Accuracy Measure</th>
<th>Recall</th>
<th>Precision</th>
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<tbody>
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<td>96%</td>
<td>94%</td>
<td>93%</td>
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<tr>
<td>13</td>
<td>96.7%</td>
<td>96.5%</td>
<td>95.9%</td>
</tr>
</tbody>
</table>

Figure 1: Flowchart of Methodology.

4 CONCLUSIONS

In this study we developed an online pipeline ZK DrugResist to find the PubMed abstracts of drug resistance combined with protein and expression level data. Our tool outperformed the already existing tools. ZK DrugResist is very proficient in mining drug resistance semantics in an automated way from literature without comprising the accuracy measure. It is freely available online and is a self-explanatory tool aimed to help researchers in finding desired information on one click. As for the future work we might extend this tool to also work on full articles rather just on the abstracts. As in some cases we found that the desired information is missing in the abstract but present in the remaining article. This will further increase our dataset size and may also improve the accuracy measure. Our tool is available at http://zkdrugresist.sabanciuniv.edu/.

REFERENCES


