Vapur: A Search Engine to Find Related Protein - Compound Pairs in COVID-19 Literature

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Abstract

Coronavirus Disease of 2019 (COVID-19) created dire consequences globally and triggered an enormous scientific effort from different domains. Resulting publications formed a gigantic domain-specific collection of text in which finding studies on a biomolecule of interest is quite challenging for general-purpose search engines due to terminology-rich characteristics of the publications. Here, we present Vapur, an online COVID-19 search engine specifically designed for finding related protein - chemical pairs. Vapur is empowered with a biochemically related entities-oriented inverted index in order to group studies relevant to a biomolecule with respect to its related entities. The inverted index of Vapur is automatically created with a BioNLP pipeline and integrated with an online user interface. The online interface is designed for the smooth traversal of the current literature and is publicly available at https://tabilab.cmpe.boun.edu.tr/vapur/.

1 Introduction

Coronavirus Disease of 2019 (COVID-19) outbreak had severe impacts on human health all around the world since December 2019, but also triggered an exceptional amount of scientific work. As of September 2020, PubMed recorded over 45K articles (Chen et al., 2020) related to COVID-19 since its inception, including works on diagnosis (Dorche et al., 2020), drug repurposing (Gao et al., 2020), and text-mining (Tarasova et al., 2020). As this tremendous literature keeps growing in the form of unstructured text, it also becomes more and more challenging for researchers to find the relevant information they need. Furthermore, the publications include named entities and chemical relations that challenge the general-purpose search engines. Therefore, it is of critical importance to build a search engine that can find relevant documents in this terminology-rich and domain-specific literature.

Biomedical named entity recognition (NER) and relation extraction can be utilized to semantically structure the publications around the biochemically related entities. When named entities and their relations are extracted, a document can be expressed as a set of triplets of the form (Entity1, Entity2, Relation). This formulation can be converted to a semantic index from the related entities to the publications in order to enable retrieving relevant documents to a query by entity and relation matching. If the same entities are referenced with different words (e.g. ACE2, Angiotensin-converting enzyme 2, Q9BYF1), we can use named entity normalization to identify different mentions of the same entity.

In this work, we present Vapur, an online search engine to find related protein - compound pairs in the COVID-19 anthology. Vapur is empowered with a biochemical relation-based inverted index that is created through named entity recognition and relation extraction on CORD-19 abstracts (Wang et al., 2020a). In order to obtain the index, we first identify and normalize the named entities in the documents using a pre-trained model, BERN (Kim et al., 2019). Afterward, we determine if the entity pairs in the same sentence are related to each other by training a binary relation extraction model on the ChemProt data set (Krallinger et al., 2017) and obtain a list of related entities for each abstract. We utilize these sets to construct an inverted index that maps related entities to the documents they are mentioned.
We illustrate our pipeline in Figure 1 and present Vapur at https://tabilab.cmpe.boun.edu.tr/vapur/. We publicly share the code and models at https://github.com/boun-tabi/vapur.

2 Related Work

Extracting information from the growing body of biomedical literature has become even more important with the explosion of scientific research. Efforts such as ChemProt (Krallinger et al., 2017) aim to promote research in this direction by presenting PubMed abstracts in which proteins, chemicals, and their relations are manually labeled. Liu et al., 2018 and Lim and Kang, 2018 developed sequential models for chemical and protein relation extraction on ChemProt, while Peng et al., 2018 used an ensemble of deep models with SVM. However, all these models depend on the named entities to be recognized beforehand.

Named entity recognition is a widely studied topic in biomedical text mining. Conditional Random Fields (CRF) was a popular approach in early biomedical NER studies (Leaman et al., 2015; Wei et al., 2015, 2018) and with the rise of deep learning, sequential models were also integrated into the models (Sachan et al., 2018). Recently, transformers-based NER models were popularized, including BERN (Kim et al., 2019) which is a state-of-the-art biomedical named entity recognition and normalization tool that builds on BioBERT (Lee et al., 2020b) to identify and normalize the entities in a sentence. BERN is adopted by recent general-domain biomedical text mining studies (Jang et al., 2019) as well as a model to answer COVID-19 related questions based on CORD-19 (Lee et al., 2020a).

CORD-19 (Wang et al., 2020a) is a data set comprising scientific publications related to COVID-19 and provided a valuable resource for text mining studies. Su et al., 2020 and Esteva et al., 2020 developed question answering models with integrated information retrieval and summarization modules, while Wang et al., 2020b focused on extracting the chemical entities in CORD-19 and presented an automatically annotated NER data set, CORD-NER. The entities in CORD-NER are annotated with multiple tools, but they are not normalized. Here, for the first time, we apply named entity normalization and relation extraction to CORD-19 in order to create a biochemical relation-aware search engine.

3 Methods

3.1 Data sets

CORD-19 We create an entity-oriented search engine for CORD-19 abstracts. CORD-19 is a regularly updated data set that contains studies related to COVID-19 and SARS-CoV-2. We use the CORD-19 snapshot of August 23, 2020, that contains ≈ 233K documents for which ≈ 143K unique abstracts are provided. Vapur indexes only these abstracts, but it is able to return the linked full-paper, as well.

ChemProt We trained the binary relation extraction model using the ChemProt data set (Krallinger et al., 2017), a curation of PubMed abstracts whose entities and relations are manually annotated. ChemProt consists of three file types: abstracts, entities, and relations. An abstract file contains a PubMed abstract ID and its text, whereas an entity file lists the chemicals and proteins in the linked abstract. Last, a relation file reports the relation type between entity pairs in the same sentence whose relation types can be inferred from the sentence.

The relations in ChemProt are categorized into 11, where the first 10 categories (CPR:0 to CPR:9)
<table>
<thead>
<tr>
<th>Relation ID</th>
<th>Description</th>
<th>Train Count</th>
<th>Dev Count</th>
<th>Test Count</th>
<th>Binary Label</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPR:0</td>
<td>No description provided by ChemProt</td>
<td>1</td>
<td>2</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>CPR:1</td>
<td>Part of</td>
<td>308</td>
<td>153</td>
<td>215</td>
<td>1</td>
</tr>
<tr>
<td>CPR:2</td>
<td>Regulator / Direct Regulator / Indirect regulator</td>
<td>1652</td>
<td>780</td>
<td>1743</td>
<td>1</td>
</tr>
<tr>
<td>CPR:3</td>
<td>Upregulator / Activator / Indirect upregulator</td>
<td>777</td>
<td>552</td>
<td>667</td>
<td>1</td>
</tr>
<tr>
<td>CPR:4</td>
<td>Downregulator / Inhibitor / Indirect Downregulator</td>
<td>2260</td>
<td>1103</td>
<td>1667</td>
<td>1</td>
</tr>
<tr>
<td>CPR:5</td>
<td>Agonist / Agonist activator / Agonist inhibitor</td>
<td>173</td>
<td>116</td>
<td>198</td>
<td>1</td>
</tr>
<tr>
<td>CPR:6</td>
<td>Antagonist</td>
<td>235</td>
<td>199</td>
<td>293</td>
<td>1</td>
</tr>
<tr>
<td>CPR:7</td>
<td>Modulator / Modulator Activator / Modulator inhibitor</td>
<td>29</td>
<td>19</td>
<td>25</td>
<td>1</td>
</tr>
<tr>
<td>CPR:8</td>
<td>Cofactor</td>
<td>34</td>
<td>2</td>
<td>25</td>
<td>1</td>
</tr>
<tr>
<td>CPR:9</td>
<td>Substrate / Product of / Substrate product of</td>
<td>727</td>
<td>457</td>
<td>644</td>
<td>1</td>
</tr>
<tr>
<td>CPR:10</td>
<td>Not relation</td>
<td>241</td>
<td>175</td>
<td>267</td>
<td>0</td>
</tr>
<tr>
<td>Other</td>
<td>No relation information between entities in this context</td>
<td>11664</td>
<td>7780</td>
<td>9987</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 1: Chemical - Protein Relations (CPR) in ChemProt. The entity pairs are annotated with 11 different relations in ChemProt and we refer to pairs whose relation information cannot be inferred from the context as Other. We report the instance count of each category and specify the label we assign to each category for binary relation extraction.

<table>
<thead>
<tr>
<th>Statistic</th>
<th>Train</th>
<th>Dev</th>
<th>Test</th>
</tr>
</thead>
<tbody>
<tr>
<td># documents</td>
<td>1020</td>
<td>612</td>
<td>800</td>
</tr>
<tr>
<td># sentences</td>
<td>18046</td>
<td>11294</td>
<td>15712</td>
</tr>
<tr>
<td># entities</td>
<td>25752</td>
<td>15567</td>
<td>20828</td>
</tr>
<tr>
<td># chemical mentions</td>
<td>13017</td>
<td>8004</td>
<td>10810</td>
</tr>
<tr>
<td># unique chemical entities</td>
<td>3710</td>
<td>2517</td>
<td>3442</td>
</tr>
<tr>
<td># protein mentions</td>
<td>12735</td>
<td>7563</td>
<td>10018</td>
</tr>
<tr>
<td># unique protein entities</td>
<td>4610</td>
<td>3018</td>
<td>3757</td>
</tr>
<tr>
<td># duplicate relations</td>
<td>98</td>
<td>86</td>
<td>158</td>
</tr>
<tr>
<td># positive labels</td>
<td>6143</td>
<td>3339</td>
<td>5459</td>
</tr>
<tr>
<td># negative labels</td>
<td>11903</td>
<td>7955</td>
<td>10253</td>
</tr>
</tbody>
</table>

Table 2: ChemProt summary statistics. We split abstracts to sentences via GENIA (Kim et al., 2003) and computed the statistics accordingly. We refer to relations in the same sentence with the same protein, chemical and relation type as duplicate relations and consider them only once during training. We observe that the # of mentions is considerably higher than the # of unique entities for both proteins and chemicals, emphasizing the need for normalization. In addition, negative entity pairs are significantly more frequent than the positive ones, but their distribution across the training, development, and test folds is similar.

denote various types of biochemical relations, while the last category (CPR:10) is reserved for “not relation” (i.e., the sentence explicitly states that there is no relation between the specified pair of entities). In this work, we focus on binary relation extraction, where we aim at determining whether a sentence states that there is a relation between the specified two entities or not. Therefore, we consider the entity pairs in the first 10 categories as positive samples. Likewise, since the sentences in the CPR:10 and the “Other” classes do not state that there is a relation between the corresponding pair of entities, we treat these two classes as the negative class. We describe each relation type and report their instance counts in the ChemProt data set in Table 1. We also provide summary statistics for ChemProt in Table 2.

3.2 Named Entity Recognition and Normalization

We used BERN (Kim et al., 2019), a neural NER architecture with an integrated normalizer that can perform in a multi-type setting, to identify and normalize the entities in CORD-19 abstracts. BERN is an ensemble of existing tools and models (Wei et al., 2015; Leaman et al., 2015; D’Souza and Ng, 2015; Wei et al., 2016, 2018; Lee et al., 2020b) and outputs a set of IDs for each recognized entity, given a sentence.

In this study, we first tokenized the abstracts with GENIA (Kim et al., 2003) and identified 1.23M
EGFR inhibitors currently under investigation include the small molecules gefitinib and erlotinib.

<table>
<thead>
<tr>
<th>Raw Sentence</th>
<th>Preprocessed Form I</th>
<th>Preprocessed Form II</th>
</tr>
</thead>
<tbody>
<tr>
<td>EGFR inhibitors currently under investigation include the small molecules gefitinib and erlotinib.</td>
<td>(&lt;e2&gt;\text{EGFR}&lt;/e2&gt;) inhibitors currently under investigation include the small molecules (&lt;e1&gt;\text{gefitinib}&lt;/e1&gt;) and erlotinib.</td>
<td>(&lt;e2&gt;\text{EGFR}&lt;/e2&gt;) inhibitors currently under investigation include the small molecules (&lt;e1&gt;\text{gefitinib}&lt;/e1&gt;) and (&lt;e1&gt;\text{erlotinib}&lt;/e1&gt;).</td>
</tr>
</tbody>
</table>

Table 3: An example of preprocessing. The raw sentence contains two chemicals (gefitinib, erlotinib) and one protein (EGFR), creating two protein - chemical pairs. Thus, we create two different forms of the sentence to encode each protein - chemical pair separately. We use \(<e1>\) and \(<e2>\) tags to enclose chemicals and proteins, respectively.

Figure 2: Relation Extraction Model. Our model is composed of a BioBERT and a binary classification layer. We tokenize the input sentences by BioBERT tokenizer and enclose the biochemical entities with special tags (\(<e1>\) and \(<e2>\)). BioBERT also adds \(\text{CLS}\) and \(\text{SEP}\) tags for fixed length-relation representation and sentence separation, respectively. We use BioBERT output of \(\text{CLS}\) token to train our binary classifier.

Sentences. We used BERN to extract named entities and their IDs in these sentences and recognized 1.58M entities in total, in which 171K are chemicals and 318K are proteins. With normalization, we computed the number of unique chemicals and proteins as 20K and 70K, respectively.

3.3 Relation Extraction

We propose a BioBERT-based (Lee et al., 2020b) model to identify related entities in CORD-19 abstracts. To this end, we first preprocessed the sentences to explicitly encode the entities in the input and then fine-tuned BioBERT with the preprocessed sentences in ChemProt. We used the binary labels in Section 3.1 as outputs.

Preprocessing: We surrounded the named entities in the sentences with opening and closing tags to mark their location. We tagged chemicals with \(<e1>\) and \(</e1>\) and proteins with \(<e2>\) and \(</e2>\) to encode entity type and location during training. When a sentence had multiple chemical - protein pairs, we considered each pair separately and created copies of the sentence with different tags. Table 3 provides an example of input and its two preprocessed forms.

Model: We used BioBERT to create a binary model that decides if two entities in the same sentence are related. We added a single-layer binary log-softmax classifier to BioBERT and trained the classifier with the inputs and outputs from the previous step. BioBERT creates a fixed-length relation representation for the starting tag \(\text{CLS}\) and we used this representation to vectorize the sentences. We trained the binary classifier with cross-entropy loss where we considered the pairs in CPR:1-9 as positive and pairs in CPR:10 or no reported relation (i.e., the “Other” Type in Table 1) as negative. Figure 2 illustrates our relation extraction model.

Experimental setup: We conducted a hyperparameter search with different optimizers, learning rates, and weight decays. We trained a model 10 times per parameter combination and selected the best one with respect to F-score on the development set. We found that the AdamW optimizer with a 0.00003 learning rate and 0.1 weight decay yields the best results. We selected the best setting for relation extraction and computed mean precision, recall, and F1-score as well as the standard deviation on the ChemProt development (dev) and test sets.

4 Vapur

Vapur is an online search engine with a focus on finding related proteins and chemicals from CORD-19, a compilation of COVID-19 literature (Wang et al., 2020a). Vapur uses an inverted index that is constructed through mining the relations and entities in CORD-19 and is able to retrieve relevant
Figure 3: A search scenario on Vapur. When Interleukin 1b is entered as the query, Vapur lists the biochemically related compounds and displays the publications that mention the relation. Unique IDs of each chemical entity are displayed to help the user for further research. Similar proteins to Interleukin 1b are also listed in order to let the user explore the index smoothly.

documents to the queried biomolecule as categorized by the relation extraction model. Vapur’s inverted index explicitly represents relations as related entity pairs and maps each relation to the documents in which the relation was mentioned.

We also interpret the relation-oriented index of Vapur as a graph to infer relations that cannot be inferred from the inverted index trivially. We construct a graph from the index such that the nodes represent biomolecules and the edges denote biochemical relations. The resulting graph satisfies the bipartiteness property, since we identify only the relations between proteins and chemicals during binary relation extraction. Table 4 demonstrates the summary statistics of this bipartite graph computed via networkx (Hagberg et al., 2008).

We leverage the bipartite graph structure to identify similar biomolecules of the same type. We use SimRank (Jeh and Widom, 2002) to compute pairwise node similarity and list the five most similar entities to the query. The aim is to encourage the user to explore the information in CORD-19 not only for exact matches to the query, but also similar entities.

The entities are represented as equivalence classes learned from named entity recognition and

<table>
<thead>
<tr>
<th>Graph Property</th>
<th>Statistic</th>
</tr>
</thead>
<tbody>
<tr>
<td># Nodes (Entities)</td>
<td>12384</td>
</tr>
<tr>
<td># Chemical Nodes</td>
<td>5018</td>
</tr>
<tr>
<td># Protein Nodes</td>
<td>7366</td>
</tr>
<tr>
<td># Edges (Relations)</td>
<td>17657</td>
</tr>
<tr>
<td># Connected Components</td>
<td>807</td>
</tr>
<tr>
<td># Nodes in the Largest Component</td>
<td>10194</td>
</tr>
<tr>
<td># Edges in the Largest Component</td>
<td>16226</td>
</tr>
<tr>
<td>Diameter of the Largest Component</td>
<td>18</td>
</tr>
</tbody>
</table>

Table 4: Summary statistics for the bipartite graph of the inverted index. We observe that number of chemicals and proteins are comparable to each other and the number of relations is close to the number of nodes, indicating the sparsity. In addition, the graph is formed of 807 connected components but the largest one is significantly larger than the others.
normalization steps. For instance, “Interleukin-1b” is represented as \{interleukin-1b, IL-1beta, IL1B, HGNC:5992, . . . , BERN:323737602\} in the index and Vapur retrieves related entities and relevant documents with a mention “Interleukin-1b” even if the query is “IL1B”. The equivalence classes cover a wide range of mention types from free-text to BERN IDs to present a flexible search experience in terms of search terms.

Vapur represents each mention type as a string and adopts a 3-gram based matching algorithm to search the queried entity in its index. Given a query, Vapur first creates a multi-set of all 3-grams of the query and computes the similarity of this set to all 3-gram multi-sets of the mentions in the index, which are pre-computed. Vapur measures the similarity of the query to a mention by generalized Jaccard similarity:

\[
J(Q, M) = \frac{\sum_i \min(Q_i, M_i)}{\sum_i \max(Q_i, M_i)}
\]

where \(Q\) and \(M\) are query and mention vectors that store the count of each 3-gram. Generalized Jaccard similarity makes Vapur more robust while retrieving results for a mistyped query.

When the related entities are found and their related abstracts are retrieved, Vapur ranks the related entities by the number of times they are co-mentioned with the queried entity. Each entity is displayed with its most frequent mention alongside the links to the papers. Figure 3 illustrates an example search scenario on Vapur, which is publicly available at https://tabilab.cmpe.boun.edu.tr/vapur/

5 Results and Discussion

In this work, we built Vapur, a CORD-19 search engine empowered by text-mining. We applied named entity recognition, named entity normalization, and relation extraction to CORD-19 abstracts and automatically created an inverted index from related entities to publications. Therefore, the performance of the named entity extraction and relation extraction modules is critical to identify relevant works. Since BERN was already evaluated for named entity recognition and normalization on the biomedical domain (Kim et al., 2019), here we focus on evaluating the relation extraction model.

We built the relation extraction model by fine-tuning a binary classifier on BioBERT and computed precision, recall, and F1-Score on the dev and test sets of ChemProt. We also fine-tuned the same classifier using English BERT based model and report the results in Table 5.

Table 5 demonstrates that BioBERT obtained higher scores in terms of all three metrics on both folds, indicating that BioBERT is superior to BERT as the pre-trained language model for relation extraction on ChemProt. We relate this with the fact that BioBERT is trained with a more domain-related text and BERT observed texts from a wider range.

We further investigated the performance of our relation extraction model by computing test accuracy for each chemical - protein relation (CPR) category and pairs with no assigned CPR label (i.e., Other). We observe that the relation extraction model achieved the highest accuracy in the Other category.
Table 7: Sample sentences with incorrectly labeled entities. The entity types of Alanine, carboxyl, and Ben are incorrectly predicted as chemical by BERN during the named entity recognition step. Consequently, the relation extraction model incorrectly identified a biochemical relation.

category, indicating that the model can successfully identify whether the context is sufficient to deduce a relation between the entities or not. We relate this with using BioBERT, which is a contextual model for relation representation. Context-awareness enables Vapur to eliminate the documents that mention the queried entity in irrelevant contexts and to retrieve a document only if it contains relation information for the query.

We analyzed 41 sample sentences that we predicted to be positive to discover the limitations of our methodology. Most of the incorrect relation labels are due to incorrect entity assignment by BERN. In some cases, parts of the protein sequence such as N-terminal, carboxyl terminal or residue names such as Asp238 are recognized as compound. Table 7 illustrates sample sentences with incorrectly labeled entities. More examples that are manually checked by a domain expert are presented on our GitHub page as an Appendix in the Readme file.

6 Conclusion

We presented Vapur, a search engine with an emphasis on identifying protein - chemical relations in the COVID-19 domain. Vapur is the first search engine we know of that uses relation extraction to construct an inverted index of related biochemical entities. Thanks to the relation extraction model, Vapur can categorize relevant documents to a query with respect to related chemical entities and present organized search results.

We evaluated the relation extraction model on ChemProt and observed that BioBERT outperforms English BERT on both development and test sets in terms of precision, recall, and F1-score. We further analyzed the model predictions and demonstrated the high specificity of the model at identifying contexts with biochemical relation information.

Although we focused on COVID-19 in this work, our approach can find application in any biomedical domain with a different choice of database to index. We believe that domain-specific scientific search engines will gain more interest in the future since COVID-19 is unlikely to be the last global health crisis for human-kind. In order too promote the development and use of search engines to help coping with such difficult times, we make the underlying pipeline and Vapur available at https://github.com/boun-tabl/vapur and https://tabilab.cmpe.boun.edu.tr/vapur/, respectively.

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