Semantic Similarity in Medical Domain

Exposé of Master Thesis Bonn University & Fraunhofer IAIS

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The goal of this thesis is to use a neural network model to map patient records to semantically similar and potentially helpful research work. For training the model large volume of raw texts from PubMed [2] will be used.

The traditional approach for finding relevant documents for a given query is to count repetitions of query terms in the documents. Different weight schemes for these counts lead to a variety of TF-IDF ranking features. However, the basic forms of such a ranking system only consider query terms, under the assumption that non-query terms are less useful for document ranking. This makes such ranking systems incapable of capturing the deep semantic meaning of the text all by itself.

Neural network models have recently shown impressive results in capturing the underlying semantics of the documents[1]. The main stumbling block in creating such a semantic model is the lack of training labeled data. Another challenge is lack of freely available data, PubMed mostly provides abstracts of research articles for free.

The main idea of this thesis is to explore ways in which deep-learning can be used for solving the problem of finding semantically similar/helpful research articles given a patient record. The chosen direction of the research is to apply Doc2Vec model proposed in [8] and explore different ways in which the Doc2Vec model can be extended/improved.

Background & Literature Review

Embedding Models

Usage of deep learning in the context of NLP tasks requires representing the text as the input for neural networks. As of late, the most used and powerful representations are one of the following embeddings: Word2Vec [9] and GloVe [12], which are word level embeddings. An extension to Word2Vec known as Doc2Vec was proposed in [8].

We look closer at the Word2Vec and Doc2Vec models, as they are the most relevant to this work. Word2Vec model uses distributed vector representation of words, a well-known framework for learning word vectors as shown in the (Figure 1). The task is to learn to predict a word given other words in the context. More formally, given a sequence of training words $w_1, w_2, w_3, ..., w_T$, the objective of the word vector model is to maximize the average log probability

$$\frac{1}{T} \sum_{t=K}^{T-K} \log p(w_t \mid w_{t-1},, w_{t+1})$$
 (1)

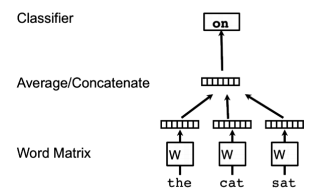


Figure 1: A framework for learning word vectors. Context of three words (the, cat, and sat) is used to predict the fourth word (on). The input words are mapped to columns of the matrix W to predict the output word.

Some commendable efforts have been made to go beyond word level representations [11] [17] [16] [5] [10]. A simple approach is to use the weighted average of all the words in the document. But weighted averaging of word vectors loses the word order. A more sophisticated approach is combining the word vectors in an order given by a parse tree of a sentence, using matrix-vector operations [15]. A drawback of such an approach is that it only works

on sentences as it relies on parse trees.

Doc2Vec is capable of constructing representations of input sequences of variable length. Unlike some of the previous approaches, it is general and applicable to texts of any length: sentences, paragraphs, and documents. In Doc2Vec framework (see Figure 2), every document is mapped to a unique vector, and every word is also mapped to a unique vector. The document vector and word vectors are averaged or concatenated to predict the next word in a context. The only difference to a Word2Vec model is the additional document token. It acts as a memory that remembers what is missing from the current context or the topic of the document. The document vectors and word vectors are trained using stochastic gradient descent and the gradient is obtained via back-propagation.

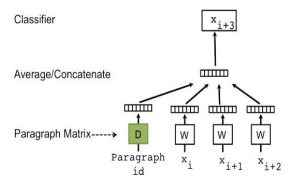


Figure 2: A framework for learning paragraph vector. This framework is similar to the framework presented in Figure 1; the only change is the additional paragraph token that is mapped to a vector via matrix D. In this model, the concatenation or average of this vector with a context of three words is used to predict the fourth word. The paragraph vector represents the missing information from the current context and can act as a memory of the topic of the paragraph.

Embedding Model Extension

Classical emitting methods such as Doc2Vec and Word2Vec are generally unsupervised requiring no domain information and as such has broad applicability. However, for highly specified domains with a moderately sized corpus, classical methods fail to find meaningful semantic relationships. [3] introduce a vocabulary driven Word2Vec method known as Dis2Vec which is used to generate disease-specific word embeddings from unstructured health-related news corpus. The input corpus D consists of a collection of word

context pairs. Based on the vocabulary V, we can categorize the word context pairs into three types as shown below:

- $D(d) = (w, c) : w \in Vc \in V$, i.e. both the word w and the context c are in V
- $D(\neg d) = (w, c) : w \notin Vc \notin V$, i.e. neither the word w nor the context c are in V
- $D(\neg d) = (w, c) : w \in V \oplus c \in V$, i.e. either the word w is in V or the context c is in V but both cannot be in V

Possible Directions

Some interesting directions which might improve the performance for the given task are the following

- Extended the query document by adding all the synonym and rephrases of the keywords.
- Focusing only on the features which are more descriptive than other.
- Learning out of vocabulary synonyms for key-words by using medical dictionaries.
- Learning word morphologies as suggested in [6]
- Using different distance metrics such as [14] [7] instead of [4] used by gensim [13].

Planned work

The work is planned to consist of following stages:

- Use the gensim [13] implementation of the Doc2Vec model as described in ([8]), and train it on the following
 - Full Wikipedia articles
 - Wikipedia info-boxes
 - A smaller subset of PubMed only containing only colorectal cancer abstracts

- Full PubMed abstracts
- PMC another Subset of PubMed which contains full articles
- PubMed uses special language/notation to describe different age groups. i.e elderly, middle aged, >60 years old and 50-60 years old. Similar language/notation is used while generating patient descriptions from a given database.
- Use different variants of Doc2Vec like DBOW, DM and DMC to see which performs the best.
- Extending the query document by adding rephrases of the key parts. For example, The query document "75 years old patient has rectal cancer" is extended to "Elderly patient, >70 years old, The patient has rectal cancer, cancer in rectum". The query document is generated in a structured way to allow easy extension. The basic structure of the query document is ."Age, Gender, Type of cancer, Type of metastasis".
- Extend the Dis2Vec [3] model to work on document level.
- The final results will compared against vanilla Doc2Vec model.

Technical Description

Implementation

Following tools are to be used in the project

- Gensim library [13] implementation will be used for the vanilla Doc2vec model and its variants.
- Pythons nltk library will be used for tokenization and stop-word removal
- Python library regex will be used for parsing the full PubMed abstracts
- To reduce the training time some elements will be implemented in Cython. Cython file will than be used to generate a .c file for faster computation
- Gcc 4.9 compiler will be used to compile the .c file generated by Cython

Datasets

For the basic research on the quality of the network, the following datasets can be used:

- Wikipedia articles corpus
- Wikipedia info-box corpus

For medical domain, the following can be used:

- Pubmed subset corpus on colorectal cancer
- Full Pubmed abstracts
- PMC another subset of PubMed which contains full articles

Validation

Wikipedia and Info-box corpus

Following steps describe the technique to be used as a proof of method on Full Wikipedia corpus and Wikipedia info-box corpus.

- Take a random target article and use the part of its information to infer a vector. Infer vector is a method to infer a vector for the out-of-sample documents. Inference starts with a low-magnitude random vector, that is then incrementally adjusted to be more predictive.
- Use this inferred vector to find 10 similar documents.
- The prediction task is to predict the article the given information belongs to.

This validation technique revealed an interesting correlation between the size of the document and the number of iterations (a.k.a steps) used to infer a vector.

PubMed corpus

For medical domain, we first infer a vector for a patient description and then find PubMed abstracts which are semantically closer to the given patient record. For result evaluation, we have 400 labeled abstracts. For each of these 400 abstracts, we have 10 other abstracts which have been labeled to be semantically close.

For a given patient record the task is to predict semantically similar PubMed abstracts which can be helpful for the doctors.

Evaluation Metric

Precision and recall are single-value metrics based on the whole list of documents returned by the system. For systems that return a ranked sequence of documents, it is desirable to also consider the order in which the returned documents are presented. Average Precision(AP) is thus define as:

$$AP = \frac{\sum_{k=1}^{n} (P(k) \times rel(k))}{\text{number of relevant documents}}$$
 (2)

where k is the rank in the sequence of retrieved documents, n is the number of retrieved documents, P(k) is the precision at cut-off k in the list and rel(k) is an indicator function equaling 1 if the item at rank k is a relevant document, zero otherwise.

For multiple queries, an extension of Average precision known as Mean average precision is used. Mean average precision for a set of queries is the mean of the average precision scores for each query. In this thesis Mean average precision(MAP) will be used as an evaluation metric.

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