A supervised learning approach to biological question answering

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Abstract. Biologists rely on keyword-based search engines to retrieve superficially relevant papers, from which they must filter out the irrelevant information manually. Question answering (QA) systems can offer more efficient and user-friendly ways of retrieving such information. Two contributions are provided in this paper. First, a factoid QA system is developed to employ a named entity recognition module to extract answer candidates and a linear model to rank them. The linear model uses various semantic features, such as named entity types and semantic roles. To tune the weights of features used by the model, a novel supervised learning algorithm, which only needs small amounts of training data, is provided. Second, a QA system may assign several answers with the same score, making evaluation unfair. To solve this problem, an efficient formula for a mean average reciprocal rank (MARR) is proposed to reduce the complexity of its computation. After employing all effective semantic features, our system achieves a top-1 MARR of 74.11\% and top-5 MARR of 76.68\%. In comparison of the baseline system, the top-1 and top-5 MARR increase by 9.5\% and 7.1\%. In addition, the experiment result on test set shows our ranking method, which achieves 55.58\% top-1 MARR and 66.99\% top-5 MARR, significantly surpasses traditional BM25 and simple voting in performance by averagely 35.23\% and 36.64\%, respectively.

1. Introduction

When planning a research project, biological researchers are predominantly interested in relevant molecular pathways and underlying mechanisms [5]. Since molecular biology is a rapidly developing and evolving field, it is essential for researchers to be able to effectively search recently published papers. Currently, most of them use keyword-based search engines such as PubMed and Google [31]. However, with the tremendous amount of new biomedical literature being published and the increasing complexity of molecular pathway descriptions, it is becoming much harder to find specific and relevant information about molecular interactions using these tools. Keyword-based information retrieval (IR) is designed to find broadly related passages, not specific answers. When biologists are interested in exactly which protein is involved in a pathway, lots of manual effort is still required to locate the desired terms from search results. Question answering (QA) systems can offer more efficient and user-friendly ways of retrieving such information.

In this paper, a QA system is built, and aimed to answer questions about biomolecular events, such as gene and protein interactions. The answers to such questions mainly consist of short pieces of information such as protein, DNA, RNA, or cell names as well as times and locations. In addition to syntactic features, two effective semantic features, named entity types and semantic roles, are incorporated to help match the question with its corresponding answer phrases contained in retrieved documents. Accordingly, our system is focused on factoid questions, to which the answers are named entities.

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Ranking potential candidate entities is another important issue. Hu et al. [14] employed a linear model that combines several semantic features to score each candidate in their entity search system. They also proposed a supervised learning approach of estimating the weights associates with these features. Their experiment results showed that the supervised learning approach is much more effective in ranking candidates when the ranking influenced by these semantic features. Therefore, their method is enhanced to apply in the QA task.

In addition, devising an accurate performance measurement for QA is also a problem, especially when two answer candidates are scored equally. Traditional performance metric, such as the top-5 mean reciprocal rank (MRR), may be inaccurate in such cases. An efficient formula for an improved MRR measurement is proposed to reduce the computational complexity in this study. This measurement can also be applied to evaluate QA systems in other domains.

The remainder of this paper is organized as follows: Section 2 contains a review of related works. In Section 3, the system workflow of our QA system, a proposed ranking method, and an improved evaluation measurement are described. Section 4 depicts our datasets, the experiment design, and the experiment results. In Section 5, the experiment results are discussed and the proposed measurement with MRR is compared. Finally, Section 6 summarizes our conclusions.

2. Related works

2.1. Information retrieval systems

2.1.1. Traditional search systems

The objective of traditional IR systems is to identify documents or passages that are relevant to a query. In general, the criterion used to judge relevance is the existence of query terms in the documents or passages. The terms are usually weighted according to their occurrences in the documents or passages by using weighting models, such as TF-IDF [23], BM25 [22], and the Language Model [20], which are unsupervised and do not require labeled data for training.

In contrast, a relatively new trend in IR is to employ supervised learning methods to train ranking functions. Herbrich et al. [12] formulated the IR problem as an ordinal regression model, and proposed a method for training the model on the basis of SVM. Gao et al. [14] conducted discriminate training on a linear IR model and observed a significant improvement in the accuracy of document retrieval as a result.

Aside from the supervised approach introduced in the last paragraph, rule-based approaches are also widely used in many domains. Besnard et al. [1] use an ontology in the form of an IS-A hierarchy to capture explanations based on causal statements. Chen et al. [4] employed a rule-based dependency models for security protocol analysis. Tao et al. [25] proposed a novel computational model for solving retrieval problems by constructing and mining a personalized ontology based on world knowledge and a user’s Local Instance Repository. According to these recent achievements in using rule-based approach, this is another possible way to handle IR problems. There are also some works inspire us during our research. Gao et al. [10] describes a new approach to the use of clustering for automatic data detection in semi-structured web pages. Pera et al. [19] provide a correlation-based phrase matching (CPM) model and a fuzzy compatibility clustering (FCC) model. CPM can detect RSS news articles containing phrases that are the same as well as semantically alike, and dictate the degrees of similarity of any two articles. FCC identifies and clusters non-redundant, closely related RSS news articles based on their degrees of similarity and a fuzzy compatibility relation. Yusof et al. [32] proposed a model which combines the functional and structural information to facilitate software component search and retrieval.

2.1.2. Entity search systems

Before discussing factoid question answering, a similar task is described—entity search. Entity search (also known as expert search) systems try to identify entities that are strongly associated with query terms. The most studied type of entity is people, which has been addressed by [3,6]. However, existing entity search methods only exploit simple features and traditional ranking methods. Hu et al. [14] employed a supervised learning technique to train an entity search model. The results of experiments on several data sets indicate that the method significantly outperforms methods based solely on the co-occurrence of terms.

2.1.3. Question answering systems

The first large-scale evaluation of QA systems was hosted by the Text Retrieval Conference (TREC) in 1999 [30], where the task focused on responding to open domain questions with short passages of 50 to 250 words. In 2003, the evaluation task became more chal-
lenging because it required systems to provide exact answers without redundant information [30].

Given a collection of documents, a QA system should be able to retrieve answers for questions posed in natural language. QA systems are categorized according to the questions they can deal with. One type of question is the factoid question, for which the answer consists of a short factual tidbit of information, such as a date, a location, or a person/organization name.

Usually, a factoid QA system transforms a natural language question into keywords, which are sent to an IR engine to retrieve search results. It then extracts possible answers from the returned documents, and rank them using natural language processing (NLP) features like shallow/full parse, tokenization, and part-of-speech (POS) tagging.

The following are two examples of systems that incorporate the above techniques and steps, as well as several others. These systems, constructed by the Language Computer Corporation (LCC) and the National University of Singapore (NUS), were the most successful participants in recent TREC QA tracks. The LCC system [11] uses the COGEX Logic Prover to verify and extract any lexical relationships between a question and its candidate answers. It achieved the best top-1 MARR (71.3%) at TREC-14. The NUS system, developed by Sun et al. [7] utilizes syntactic information and semantic information generated by the MiniPar dependency parser [16] and the ASSERT semantic role labeler [21], respectively. Since the MiniPar parser does not work well with web documents, the system uses semantic role information to extract reliable answer candidates. Finally, it employs dependency-relation-based answer ranking to verify if the web answer is correct for the context. The NUS system achieved second place (66.6%) in top-1 MRR in TREC-14.

2.2. Traditional performance measurement

There are two commonly used measurements for QA system performance measurement: the top-1 accuracy and the top-5 mean reciprocal rank (MRR). For a question set Q, the, top-1 accuracy reports the average accuracy of the top-1 answers for all the questions. It is defined as follows:

\[
\text{top-1 accuracy} = \frac{\# \text{ of correct top-1 answers}}{|Q|}
\]

The top-5 mean reciprocal rank (MRR) [30] is calculated as follows:

\[
MRR(Q) = \frac{\sum_{q \in Q} RR(q)}{|Q|},
\]

where \(q_i\) is the \(i\)th question; and \(rank(q_i)\) is the rank of the first correct answer on the list of answer candidates for \(q_i\). In the first formula above, top-1 MRR uses 1 in place of 5.

However, there may be many answer candidates with the same score and all in the leading five places. This results in multiple answer candidate sequences with different RR scores for a question, but all based on the same score. Although selecting a candidate sequence at random and calculating its RR to represent all sequences may solve this problem, the true system performance may be overestimated or underestimated. To address this problem, a new measurement called the average reciprocal rank (ARR) is proposed and discussed in Section 3.3.

3. Biomedical question answering system

3.1. System workflow

The proposed QA system, BeQA, is comprised of four components, namely Question Processing, Passage Retrieval, Candidate Extraction and Feature Generation, and Answer Ranking, which are described in detail in the following sub-sections.

3.1.1. Question processing

Question processing transforms natural language questions into search keywords and extracts features for answer ranking. In our work, question processing involves five steps: named entity (NE) recognition (NER) [9,24,28,29], semantic role labeling (SRL) [15, 26], question classification, and query modification.

The NER step extracts named entities (NEs), such as protein and gene names, from the original question. Then, the SRL step extracts predicates (e.g., the main verb) and corresponding arguments (e.g., noun phrases) from the question. Both the NEs and the SRL information will be transformed into features and used by the answer ranking module, which is described in the Method section.

In the question classification step, hand-crafted patterns are used to identify the target NE type, such as protein, cell, DNA, and RNA, requested by the question (i.e., the answer’s NE type). The classified NE type is then sent to the ranking module, which filters out unmatched answer candidates. In the phrase chunking
step, questions are parsed by the GENIA Tagger [3]. Each word in the remaining phrases is then examined and eliminated if it appears on the stop word list. The remaining phrase segments are sent to the passage retrieval module as keywords for a Google search.

Query modification is used to improve recall for queries where Google returns pages. First, using WordNet [18] and Longman’s dictionary [2], queries are expanded to generate a list of synonyms and other tenses for the main verb of the question. Then, the web search with the expanded query terms is repeated. If there are still no returned pages, keywords begin to be removed to improve recall.

3.1.2. Passage retrieval
The passage retrieval module is a Google-interfacing program that sends queries to Google and retrieves a collection of web pages, which are sent to the answer extraction module. In the passage retrieval stage, to avoid unnecessary noise, only pages from Google’s index of the PubMed database on the NCBI website are retrieved.

3.1.3. Candidate extraction and feature generation
In this stage, two extraction technologies, NER and SRL, are utilized to extract candidate NEs and their corresponding features. NER extracts named entities for answer candidates, and generates features to help match a query with passages containing relevant NEs. The NERBio [28] is employed to identify four types of NE: protein, DNA, RNA, and cell. Biomolecular events in nominal form (e.g., protein expressions), in which the relevant NEs are involved, are also extracted. In our system, each candidate is output with the sentence that contains it, and the sentence is treated as its supporting evidence.

An SRL system – BIOSMILE [8,26] is developed to generate semantic features for answer ranking. SRL can recognize the predicate of a sentence and its corresponding argument phrases, such as the agent, recipient, and location. The argument types recognized by our SRL component and their descriptions are listed in Table 1.

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arg0</td>
<td>Agent</td>
</tr>
<tr>
<td>Arg1</td>
<td>Direct object / Theme / Patient</td>
</tr>
<tr>
<td>Arg2-5</td>
<td>Not fixed</td>
</tr>
<tr>
<td>ArgM-NEG</td>
<td>Negation marker</td>
</tr>
<tr>
<td>ArgM-LOC</td>
<td>Location</td>
</tr>
<tr>
<td>ArgM-TMP</td>
<td>Time</td>
</tr>
<tr>
<td>ArgM-MNR</td>
<td>Manner</td>
</tr>
<tr>
<td>ArgM-EXT</td>
<td>Extent</td>
</tr>
<tr>
<td>ArgM-ADV</td>
<td>General-purpose</td>
</tr>
<tr>
<td>ArgM-PNC</td>
<td>Purpose</td>
</tr>
<tr>
<td>ArgM-CAU</td>
<td>Cause</td>
</tr>
<tr>
<td>ArgM-DIR</td>
<td>Direction</td>
</tr>
<tr>
<td>ArgM-DIS</td>
<td>Discourse connectives</td>
</tr>
<tr>
<td>ArgM-MOD</td>
<td>Extent</td>
</tr>
<tr>
<td>ArgM-REC</td>
<td>Reflexives and Reciprocals</td>
</tr>
<tr>
<td>ArgM-PRD</td>
<td>Marks of secondary predication</td>
</tr>
</tbody>
</table>

3.1.4. Answer ranking
Each NE extracted in the previous step is treated as an answer candidate. The answer ranking module is responsible for calculating each candidate’s score. A linear model is employed to calculate a candidate’s score based on its features. To estimate feature weights more precisely, a supervised weight tuning procedure, which is described in the next section, is proposed.

3.2. Method
3.2.1. Linear answer ranking method
To calculate an answer candidate’s score, a new ranking method called linear answer ranking is proposed. It uses a linear function (combination of features) to calculate the weighted sum of the candidate’s features. Each candidate $c$ identified in the candidate extraction step is represented as a binary feature vector $f_c$. The $i$th dimension of $f_c$, $(f_{ci})$, indicates whether or not $c$ meets the criterion of the binary feature function $f_i$, which has a corresponding weight $w_i$. Therefore, the score of a candidate $c$ is calculated as follows:

$$\text{score}(c) = f_c \cdot w = \sum_i f_{ci} w_i,$$

where $w$ is the weight vector that corresponds to $f_c$.

3.2.2. Tuning feature weights
To improve the ranking results, a weight tuning procedure is applied. First, the procedure generates all possible weight combinations of the eight features, whose weights have integer values between 1 and 10. This yields $10^8$ different combinations. To avoid generating too many weight vectors with the same score, top-
5 MRR, rather than the top-1 accuracy, is utilized to measure the weight of each vector.

Next, for each of the top 20 weight vectors, new vectors are created by adjusting the weights upward or downward by 0.5, or by leaving the weight of a given vector unchanged. This produces $3^n - 1$ new vectors, ($n$ denotes the number of dimensions). The process is then repeated with an upward or downward adjustment of 0.25, and the algorithm iterates repeatedly until the weight decrement reaches 0.125. The combination of eight weights with the highest top-5 MRR scores is taken as the final feature weight set.

3.2.3. Features

The proposed BeQA system employs eight features: Verb_Match ($f_{VM}$), Argument_Match ($f_{ARGM}$), NE_Match ($f_{NEM}$), NE_Similarity ($f_{NES}$), KeyWord_Similarity ($f_{KWS}$), Consecutive_Word_Match ($f_{CWM}$), and Google_Reciprocal_Rank ($f_{GRR}$). Answer candidate is denoted as $c$, the query as $q$, the sentence containing $c$ as $s$, and the page containing $s$ as $p$. The eight feature types are defined as follows:

$$f_{VM}(c) = \begin{cases} 1 & \text{if } c\text{'s verb matches } q\text{'s main verb} \\ 0 & \text{otherwise} \end{cases}$$

$$f_{ARGM}(c) = \begin{cases} 1 & \text{if } c\text{'s semantic role matches} \\ \text{the target role} \\ 0 & \text{otherwise} \end{cases}$$

$$f_{NEM}(c) = \begin{cases} 1 & \text{if } c\text{'s NE type matches the} \\ \text{target type} \\ 0 & \text{otherwise} \end{cases}$$

$$f_{NES}(c, q, s) = \frac{\# \text{ of NEs in } s \text{ that match NEs in } q}{\# \text{ of NEs in } q}$$

$$f_{KWS}(c, q, s) = \frac{\# \text{ of keywords in } s \text{ that match}}{\# \text{ of keywords in } q}$$

$$f_{ARGS}(c, q, s) = \frac{\# \text{ args in } s \text{ that match arguments}}{\# \text{ args in } q \text{ excluding the target argument}}$$

$$f_{CWM}(c, q, s) = \frac{\# \text{ of consecutive words in } s \text{ that}}{\# \text{ of consecutive words in } q}$$

$$f_{GRR}(p) = \text{the Google reciprocal rank of } p$$

The first three features listed above are binary features. The next three represent the similarity between $q$ and $s$. The seventh feature denotes keywords adjacent to $c$ match that of $q$; and the last feature is $p$’s Google reciprocal rank. The values of the last five features range between 0 and 1.

3.3. An improved evaluation measurement

As mentioned in Section 2.2, the traditional QA measurement approach may overestimate or underestimate a QA system’s performance. In this section, a new measurement, top-$k$ ARR, which avoids the problem, is described.

The top-$k$ ARR score is the average of all possible ranked lists’ top-$k$ MRR scores. It is defined as follows:

$$r_i(s) = \begin{cases} \text{rank}(q_i), & \text{rank}(q_i) \leq k \\ \infty, & \text{rank}(q_i) > k \end{cases}$$

$$\text{top - } k \ \text{ARR}(q) = \frac{\sum_{q \in Q} r_i(s)}{|Q|}$$

where $S$ is the set of all possible ranked lists, including all the answer candidates for $q$; and $s$ is one list in $S$.

To further explain the differences between top-$k$ MRR and MARR, the following example is used to compare the results of the RR and ARR methods. Suppose there are three answer candidates A, B, and C, all of which have the highest score. However, only A is the correct answer candidate. Using the RR measurement, different ranked list containing A, B, and C are produced. There are $3! = 6$ lists, as shown in Table 2. The RR score for each list is shown in the last column, and the way to obtain it is introduced in Section 2.2. In this case, the QA system has a one-sixth probability of getting one of these sequences. Consequently, each run of multiple experiments may produce different evaluation results.

However, by using top-$k$ ARR, all the top-$k$ RR scores can be summed, and then divide by $|S|$. Therefore, the result is

$$\text{top-5 ARR}(q) = (1+1+1/2+1/3+1/2+1/3)/6 = 11/18,$$

which is a fixed value. In contrast to the RR method, ARR can evaluate the QA systems’ performance precisely.
However, the above ARR method has two limitations. (1) If the value of \(|S|\) is very large, calculating the ARR score directly is inefficient; for example, if 170 answer candidates have the highest score, 170! permutations need to be expanded totally. (2) Technologically, there are no numerical data types that can fit such a large value.

To solve the above problems in calculating the ARR score, the following efficient formula is proposed in the [17]:

\[
ARR(q_t) = \frac{\min(r+m-1,5)\ n(m-r)!(m-n)!}{r(m-n-r+1)!m!}
\]

where \(m\) represents the number of answer candidates with the same score; \(n\) denotes the number of correct answer candidates with the same score; and \(r\) indicates the highest rank over all correct answer candidates in all possible ranked list.

Although the above formula provides an efficient way to calculate the ARR score, it still has a problem. Consider the situation of \(r\) larger than 1 and \(m\) is equal to \(n\) (This is possible since maybe all of the answer candidates with same score are correct), then the item \(m - n - r + 1\) at the denominator will be less than zero, making its factorial undefined. To fix this problem, this formula is revised as follows:

\[
top - k\ ARR(q_t) = \frac{\min(r+m-n,k)\ n(m-n)!(m-(t-r)-1)!}{t(m-n-(t-r))!m!}
\]

Using the above formula, the top-5 ARR score is calculated as follows (\(r = 1\), \(m = 3\), \(n = 1\)):

\[
\sum_{t=1}^{3} \frac{1(3-1)!(3 - (t-1) - 1)!}{t(3 - 1 - (t-1))!} = 11/18.
\]

Here, the score is equal to that of previous ARR method. It provides a convenient way of calculating a large number of answer candidates with the same score.

### Table 2

<table>
<thead>
<tr>
<th>Sequence</th>
<th>Top 1</th>
<th>Top 2</th>
<th>Top 3</th>
<th>RR</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Δ</td>
<td>B</td>
<td>C</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>Δ</td>
<td>C</td>
<td>B</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>B</td>
<td>Δ</td>
<td>C</td>
<td>1/2</td>
</tr>
<tr>
<td>4</td>
<td>B</td>
<td>Δ</td>
<td>A</td>
<td>1/3</td>
</tr>
<tr>
<td>5</td>
<td>C</td>
<td>Δ</td>
<td>B</td>
<td>1/2</td>
</tr>
<tr>
<td>6</td>
<td>C</td>
<td>B</td>
<td>Δ</td>
<td>1/3</td>
</tr>
</tbody>
</table>

**4. Results**

### 4.1. Dataset

To the best of our knowledge, there are no well-established online factoid QA systems dedicated to the biomolecular domain. Hence, it is difficult to obtain a representative set of user queries for use as a benchmark. To create a question set, biologists in our laboratory referred to the TREC Genomics Track [13] to choose appropriate abstracts and generate candidate questions.

An independent committee composed of several other biologists then selected 400 questions from among the generated candidates. Next, the questions are divided into two sets, a development set and a test set, each containing 200 questions. The answer types of the 400 biomolecular event questions cover four NE classes, namely protein, DNA, RNA, and cell (including cell line and cell type). Furthermore, each question is based on one of 30 common biomolecular verbs selected by Chou et al. [27], shown in Table 3.

Six example selected questions with SRL annotations are shown as follows:

1. \([R−Arg0]Which protein\) \([predicate{increases}]\) \([Arg1]levels of active nuclear NF-kappa B complex\)?
2. \([R−AM−LOC]In which type of cell\) does \([Arg0]human immunodeficiency virus type 1 Nef protein\) \([predicate{inhibit}]\) \([Arg1]NF-kappa B induction\)?
3. \([R−Arg1]The transcription of which gene\) is \([predicate{enhanced}]\) \([Arg0]by recombinant OTF-2 protein\)?
4. \([AM−LOC]In human T lymphocytes,\) \([R−Arg1]Which protein\) is \([predicate{induced}]\) \([Arg0]by ALD\)?
5. \([R−Arg0]Which protein\) \([predicate{regulates}]\) \([Arg1]monocyte migration and activation\)
6. \([R−Arg1]Which mRNA\) is \([predicate{increased}]\) \([Arg0]by EBNA-2 expression in Daudi cells\)?
4.2. Experiment design and results

The following three experiments are designed to evaluate the performance of BeQA. In each experiment, the query results of all configurations are cached to eliminate the influence of updates on Google’s index.

The first experiment uses the development set to tune the best weight combination for the proposed features, and then applies the combination to the test set. The second experiment compares the proposed ranking method with two popular ranking methods. Finally, an experiment is conducted to assess the impact of the number of pages returned by a Google search.

4.2.1. Experiment 1

Verb Match ($f_{VM}$), NE Similarity ($f_{NES}$), NE Match ($f_{NEM}$), and Key-Word Similarity ($f_{KWS}$) are taken as the features of the baseline system. To assess the contribution of each feature, its related features are compared by adding Argument Match ($f_{ARGM}$), Argument Similarity ($f_{ARGS}$), Consecutive Word Match ($f_{CWM}$) and Google Reciprocal Rank ($f_{GRR}$) to the baseline configuration individually. The five configurations are Baseline, ARG, ARG, CWM, and GRR. Furthermore, all the features are incorporated into the sixth configuration, ALL. Table 4 shows the performance comparison of $f_{ARGM}$, $f_{ARGS}$, $f_{CWM}$, $f_{GRR}$ and ALL. When applied individually, $f_{CWM}$ and $f_{ARGM}$ are the most effective features; $f_{ARGS}$ can also improve the performance when used alone or with other features. By incorporating all the features, the top-1 MARR and top-5 MARR results are 74.11% and 76.68%, respectively.

For the development set, the actual tuned weights determined by our tuning procedure for the ALL configuration are shown in Table 5.

In Table 6, the actual tuned weights are applied to the test set, and the performance of the best configurations in the development and test sets are compared.

4.2.2. Experiment 2

For the comparison experiment, two traditional ranking functions are used: BM25 [22] and simple-voting [14]. BM25 is commonly used as an information retrieval function to rank passages. In this experiment, the function is used along with heuristics to rank NEs. The BM25 function is utilized to score the passages in the beginning. Then, the highest ranked passage containing at least one NE of the target type is examined, and the NE closest to the predicate is considered the top-1 answer. The simple-voting method ranks entities based on the number of supported passages. As shown in Table 7, our QA system significantly outperforms the compared methods.

4.2.3. Experiment 3

To explore the impact of maximum returned page (MRP), the performance of all configurations in MRP is examined, ranging from 2 to 14. As shown in Fig. 1, most target entities relevant to the query can be found in the first 13 pages. The top-1 MARR and top-5 MARR values increase slowly after the maximum number of returned pages reaches 13.
5. Discussion

Compared to BM25, which depends solely on co-occurrence statistics, our method ranks candidates more intuitively and naturally with semantic information. In this section, some examples are provided to demonstrate the effectiveness of each proposed syntactic or semantic feature, and an additional experiment that compares MRR with MARR is described.

5.1. The effects of using $f_{CWM}$

**Question:** Which protein inhibits the synthesis of Ig mRNA?

**Answer:**

<table>
<thead>
<tr>
<th>Baseline</th>
<th>CWM</th>
<th>Passage</th>
</tr>
</thead>
<tbody>
<tr>
<td>16 20.17</td>
<td></td>
<td>These findings demonstrate that [TGF-beta] decreases B lymphocyte Ig secretion by inhibiting the synthesis of Ig mRNA and inhibiting the switch from the membrane form to the secreted forms of mu and gamma mRNA.</td>
</tr>
<tr>
<td>16 18.50</td>
<td></td>
<td>Transforming growth factor-beta suppresses [human B lymphocyte Ig] production by inhibiting synthesis and the switch from the membrane form to the secreted form of Ig mRNA.</td>
</tr>
</tbody>
</table>

The $f_{CWM}$ feature reinforces the keyword match feature by considering consecutive matches between questions and passages. The experiment result shows that $f_{CWM}$ increases accuracy by approximately 7%. The above example is to demonstrate $f_{CWM}$’s effectiveness. The first and second columns show the scores of the Baseline and CWM configurations, respectively. The third column shows answer candidates (the phrases in brackets) and their passages. In the baseline configuration, both candidates achieve a score of 16. However, after adding $f_{CWM}$, the first candidate achieves a better score than the second. This is because the first passage’s CWM value is higher than that of the second. In the first passage, there is a consecutive five-word match, “the synthesis of Ig mRNA”, which is underlined. In the second passage, the length of the consecutive match “Ig mRNA” is only three words. This example demonstrates that $f_{CWM}$ is very useful for disambiguating candidates with similar contexts.

5.2. The effects of using $f_{ARGM}$

Here an example is given to illustrate how $f_{ARGM}$ significantly enhances the performance of the top-1 and top-5 MARR. In the question, “Which protein interacts with the alpha subunit of TFIIA?”, the semantic roles are tagged in addition to named entity tagging, so the question becomes:

[R→Arg0Which protein] [predicateinteracts] [Arg1 with the alpha subunit of TFIIA]?

In this question, the target role is Arg0 because “which” locates at R-Arg0. Therefore, our QA system searches for an Arg0 protein. The following is the answer sentence corresponding to the above question:

First, [Arg0Tax] was found to [predicateinteract] [Arg1 with the 35-kDa (alpha) subunit of TFIIA] [ArgMLOC in the yeast two-hybrid interaction system].

Although the requested answer is a protein, our QA system successfully identifies the “Tax” protein rather than “TFIIA” since “Tax” locates at an Arg0 argument, which matches the target role (Arg0).
5.3. The effects of using \( f_{ARGS} \)

The experiment results show that using \( f_{ARGS} \) alone can improve the performance by 4.52% in (top-1 MARR) and 6.09% in top-5 MARR). Using the feature cooperatively with other features (mainly \( f_{ARGM} \)) yields similar improvements. Compared to ARGM, ARGS tends to find answer sentences that have predicate-argument structures similar to the question. This finding suggests that these two features work independently. The following is a question and its corresponding answer that exclusively retrieved by employing \( f_{ARGS} \). The question and answer have three common arguments (Arg0, Arg1, and ArgM-LOC).

**Question:**
\[ R - Arg1 \] The expression of which protein is \[ predicate \] inhibited by \[ Arg0 \] IL-10 [ArgM-LOC in activated human monocytes]?

**Answer text:**
\[ Arg0 \] Interleukin-10 (IL-10), like IL-4, is known to \[ predicate \] inhibit \[ Arg1 \] cytokine expression [ArgM-LOC in activated human monocytes].

5.4. Comparing MRR with MARR

To compare MARR with MRR, using the same dataset, 30 additional experiments are conducted on the ALL configuration described in Section 4.2.1. From the results shown in Fig. 2, MARR yields a stable evaluation result, while MRR arbitrarily changes in response to each experiment. The results demonstrate that the proposed method can evaluate any QA system precisely and avoid the same score problem that makes MRR inaccurate.

6. Conclusion

Our proposed QA system provides biologists with another way to obtain the information they need. Compared with general IR systems, which retrieve all possible documents rather than an exact answer, a QA system retrieves a specific answer from a limited number of pages. In this paper, a more reliable method is designed to select the suitable answer from candidates, and incorporates syntactic and semantic features, including NE matching (NEM), verb matching (VM), SRL argument matching (ARGM), NE similarity (NES), keyword similarity (KWS), SRL argument similarity (ARGS), consecutive word matching (CWM), and Google reciprocal rank (GRR). To tune the optimal weight of each feature, a supervised learning algorithm is applied to adjust the feature weights reliably. Our experiments show that with the syntactic feature, CWM, the top-1 and top-5 MARR can be improved by 6.53% (from 57.94% to 64.47%) and 8.01% (from 58.07% to 66.08%,) respectively. Because the SRL system is used to label the semantic structures of input queries and candidate passages, our QA system can benefit from semantic information. For example, with the ARG feature, the top-1 MARR improves by 5.86% and the top-5 MARR by 7.3%. After combining all the syntactic and semantic features, the proposed BeQA system outperforms traditional ranking functions, such as BM25 and simple voting, by about 25.58% top-1 MARR and 26.99% top5-MARR on the test set. After combining all the syntactic and semantic features, the performance of the BeQA system achieves 74.11% top-1 MARR and 76.68% top-5 MARR.
Notwithstanding our system considers eight features with different weights, it still suffers from the same score problem that affects widely used measurement methods. To resolve the problem, a new measurement called the average reciprocal rank (ARR), which is the average of all possible RR score sequences, is proposed. However, expanding all permutations to calculate the ARR is inefficient, so an efficient formula is further proposed and the equality of the results is demonstrated.

In future work our plans are to (1) increase the variety of answer types by including more NE classes, such as diseases and viruses; (2) expand our corpus sources from short abstracts to full papers or other authoritative biomedical digital libraries; and (3) link the extracted answers directly to other databases or resources to provide biologists with related information in a fast and efficient manner.

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