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Biomedical Terminology Extraction: A new combination of Statistical and Web Mining Approaches

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Abstract

The objective of this work is to combine statistical and web mining methods for the automatic extraction, and ranking of biomedical terms from free text. We present new extraction methods that use linguistic patterns specialized for the biomedical field, and use term extraction measures, such as C-value, and keyword extraction measures, such as Okapi BM25, and TFIDF. We propose several combinations of these measures to improve the extraction and ranking process and we investigate which combinations are more relevant for different cases. Each measure gives us a ranked list of candidate terms that we finally re-rank with a new web-based measure. Our experiments show, first that an appropriate harmonic mean of C-value used with keyword extraction measures offers better precision results than used alone, either for the extraction of single-word and multi-words terms; second, that best precision results are often obtained when we re-rank using the web-based measure. We illustrate our results on the extraction of English and French biomedical terms from a corpus of laboratory tests available online in both languages. The results are validated by using UMLS (in English) and only MeSH (in French) as reference dictionary.

1. Introduction

The huge amount of data available online today is often composed of plain text field, for instances clinical trial description, adverse event report or electronic health records. These texts often contain the real language (expressions and terms) used by the community.
Although in the biomedical domain there exist hundred of terminologies and ontologies to describe such languages (Noy et al., 2009), those terminologies often miss concepts or possible alternative terms for those concepts. Our motivation is to improve the precision of automatic terms extraction process, the main reason for this, is that language evolves faster than our ability to formalize and catalog it. This is even more true for French in which the number of terms formalized in terminologies is significantly less important than in English.

NLP (natural language processing) tools and methods enable to enrich biomedical dictionaries from texts. Automatic Term Recognition (ATR) is a field in language technology that involves the extraction of technical terms from domain-specific language corpora (Zhang et al., 2008). Similarly, Automatic Keyword Extraction (AKE) is the process of extracting the most relevant words or phrases in a document with the propose of automatic indexing. Keywords, which we define as a sequence of one or more words, provide a compact representation of a document’s content; two popular AKE measures are Okapi BM25 (Robertson et al., 1999) and TFIDF (also called weighting measures). These two fields are summarized in Table 1.

<table>
<thead>
<tr>
<th></th>
<th>ATR</th>
<th>AKE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Automatic Term</strong></td>
<td><strong>Automatic Keyword</strong></td>
<td><strong>Recognition</strong></td>
</tr>
<tr>
<td><strong>Input</strong></td>
<td>one large corpus (i.e., not explicitly</td>
<td>single document within a dataset of documents</td>
</tr>
<tr>
<td></td>
<td>separated in documents)</td>
<td></td>
</tr>
<tr>
<td><strong>Output</strong></td>
<td>technical terms of a domain</td>
<td>keywords that describe the document</td>
</tr>
<tr>
<td><strong>Domain</strong></td>
<td>very specific</td>
<td>none</td>
</tr>
<tr>
<td><strong>Exemples</strong></td>
<td><em>C</em>-value</td>
<td><em>TFIDF</em>, <em>Okapi</em></td>
</tr>
</tbody>
</table>

*Table 1: Differences between ATR and AKE.*

In our work, we adopt as baselines an ATR method, *C*-value (Frantzi et al., 2000), and the best two AKE methods (Hussey et al., 2012), previously mentioned and considered state-of-the-art. Indeed, the *C*-value, compared to other ATR methods, often gets best precision results and specially in biomedical studies (Knoth et al., 2009), (Zhang et al., 2008), (Zhang et al., 2004). Moreover, *C*-value is defined for multi-word term extraction but can be easily adapted for single-word term and it has never been applied to French text, which is appealing in our case.

Our work follows two main steps: (a) we create new extraction methods by combining in different manners ATR and AKE measures, and we select the best list of ranked candidate terms, (b) we re-rank these extracted lists with a new web-based measure to obtain a new ranked list of candidate terms that maximize precision. Our results experiments present a great improvement of the precision with these new combined methods. We give priority to precision in order to focus on extraction of new valid terms (i.e., for a candidate term to be a valid biomedical term or not) rather than on missed terms (recall).
The rest of the paper is organized as follows: Section 2 describes the related work in the field of ATR, and specially the uses of the \textit{C-value}; Section 3 presents our combination of measures and the web-based measure for re-ranking candidate terms; Section 4 shows and discusses our experiment results; and Section 5 concludes the paper.

2. Related Work

ATR studies can be divided into four main categories: (i) rule-based approaches, (ii) dictionary-based approaches, (iii) statistical approaches, and (iv) hybrid approaches. Rule-based approaches for instance (Gaizauskas et al., 2000), attempt to recover terms thanks to the formation patterns, the main idea is to build rules in order to describe naming structures for different classes using orthographic, lexical, or morphosyntactic characteristics. Dictionary-based approaches use existing terminology resources in order to locate term occurrences in texts (Krauthammer et al., 2004). Statistical approaches are often built for extracting general terms (Eck et al., 2010); the most basic measure is frequency. XTRACT (Sma dja, 1993) is a statistical method, first it extracts the binary terms located in a window of ten words. The binary terms selected are those that exceed a statistically significantly frequency due to the chance. The next step is to extract the terms containing the binary terms found in the previous step. Another method is ACABIT (Daille et al., 1994) that performs a linguistic analysis to convert the nominal terms in binary terms. Then these terms are sorted according to statistical measures. \textit{C}/\textit{NC-value} (Frantzi et al., 2000), is another statistical method well known in the literature that combines statistical and linguistic information for the extraction of multi-word and nested terms. While most studies address specific types of entities, \textit{C}/\textit{NC-value} is a domain-independent method. It was also used for recognizing terms from biomedical literature (Hliaoutakis et al., 2009). The \textit{C}/\textit{NC-value} method was also applied to many different languages besides English (Frantzi et al., 2000) such as Japanese (Mima et al., 2001), Serbian, Slovenian, Polish, Chinese (Ji et al., 2007), Spanish (Barrón et al., 2009), and Arabic, however to the best of our knowledge not for French. An objective of this work is to combine this method with AKE methods and to apply them to English and French. We believe that the combination of biomedical term extraction and the extraction of keywords describing a document, could be beneficial since keywords techniques give greater importance to the actual terms of this domain. This combination has never been proposed and experimented in the literature.

3. Proposed Methodology for Automatic Biomedical Term Extraction

This section describes the baselines measures and their customizations as well as the new combinations of these measures and the new web-based measure that we propose for automatic biomedical terms extraction and ranking. Our method for automatic term extraction has five main steps; described in Figure 1:

1. Part-of-Speech tagging,
2. Candidate terms extraction,
3. Ranking of candidate terms,
4. Computing the new combined measures,
5. Re-ranking using web-based measure.

We execute those five steps taking either \textit{C-value} (right branch), and \textit{Okapi/TFIDF} (left branch) as the baseline method. Notice, that because \textit{C-value} is a method that deals with an unique corpus as input whereas weighting measure deal with several documents (cf. Table 1) then we need to do the union of documents of the corpus in the right branch case, in order to
consider the whole corpus as an unique document. A preliminary step not represented in Figure 1 is the creation of patterns for French and English, as described hereafter.

Figure 1: Workflow Methodology for Biomedical Term Extraction.
3.1. Part-of-Speech tagging

Part-of-speech (POS) tagging is the process of assigning each word in a text to its grammatical category (e.g., noun, adjective). This process is performed based on the definition of the word or on the context which it appears in.

We apply part-of-speech to the whole corpus. We evaluated three tools (TreeTagger, Stanford Tagger and Brill’s rules), and finally choose TreeTagger which gave the best results and is usable both for French and English.

3.2. Candidate Terms Extraction

3.2.1. Building biomedical patterns

As previously cited work, we supposed that biomedical terms have similar syntactic structure. Therefore, we build a list of the most common lexical patterns according the syntactic structure of biomedical terms present in the UMLS\(^1\) (for English) and the French version of MeSH\(^2\) (for French).

We also do a part-of-speech tagging of the biomedical terms using TreeTagger\(^3\), then compute the frequency of syntactic structures. We finally choose the 200 highest frequencies to build the list of patterns for each language. The number of terms used to build these lists of patterns was 2 300 000 for English and 65 000 for French. Examples of patterns are given in Table 2:

<table>
<thead>
<tr>
<th>English</th>
<th>French</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ProperNoun</td>
<td>Noun</td>
</tr>
<tr>
<td>2 Noun</td>
<td>Noun Adj</td>
</tr>
<tr>
<td>3 ProperNoun ProperNoun</td>
<td>Noun Prep Noun</td>
</tr>
<tr>
<td>4 Noun Noun</td>
<td>Noun Adj Adj</td>
</tr>
<tr>
<td>5 Adj Noun</td>
<td>Noun Prep:det Noun</td>
</tr>
<tr>
<td>6 Noun Noun ProperNoun</td>
<td>Noun Prep ProperNoun</td>
</tr>
<tr>
<td>7 Adj ProperNoun ProperNoun</td>
<td>Noun ProperNoun</td>
</tr>
<tr>
<td>8 Noun ProperNoun ProperNoun</td>
<td>Noun Noun</td>
</tr>
<tr>
<td>9 Noun Noun Prep Noun</td>
<td>Noun Prep Noun Adj</td>
</tr>
</tbody>
</table>

Table 2: 9 most frequent syntactic structures of biomedical terms.

3.2.2. Candidate terms extraction following patterns

Before applying any measures we filter out the content of our input corpus using patterns previously computed. We select only the terms which syntactic structure is in the patterns list.

3.3. Ranking of Candidate Terms

3.3.1. Using C-value

The C-value method combines linguistic and statistical information (Frantzi et al., 2000); the linguistic information is the use of a general regular expression as linguistic patterns, and the statistical information is the value assigned with the C-value measure based on frequency of terms to compute the termhood (i.e., the association strength of a term to domain concepts).

\[^1\] http://www.nlm.nih.gov/research/umls
\[^2\] http://mesh.inserm.fr/mesh/
\[^3\] http://www.cis.uni-muenchen.de/~schmid/tools/TreeTagger/
The aim of the C-value method is to improve the extraction of nested terms, it was specially built for extracting multi-word terms.

\[
C_{value}(a) = \begin{cases} 
    w(a) \times f(a), & \text{if } a \notin \text{nested} \\
    w(a) \times \left( f(a) - \frac{1}{|S_a|} \times \sum_{b \in S_a} f(b) \right), & \text{otherwise}
\end{cases}
\]

(1)

<table>
<thead>
<tr>
<th>Original C-value</th>
<th>Modified C-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(w(a) = \log_2(</td>
<td>a</td>
</tr>
<tr>
<td>antiphospholipid antibodies</td>
<td>(\log_2(2) = 1)</td>
</tr>
<tr>
<td>white blood</td>
<td>(\log_2(2) = 1)</td>
</tr>
<tr>
<td>platelet</td>
<td>(\log_2(1) = 0)</td>
</tr>
</tbody>
</table>

*Table 3: Calculation of \(w(a)\).*

3.3.2. Using Okapi - TFIDF

Those measures are used to associate each term of a document with a weight that represents its relevance to the meaning of the document it appears relatively to the corpus it is included in (and relatively to the size of the document int the case of Okapi). The output is a ranked list of terms for each document, which is often used in information retrieval, to order documents by their importance given a query (Robertson et al., 1999). Okapi can be seen as an improvement of TFIDF measure, taking into account the document length.

The outputs of Okapi and TFIDF are calculated with a variable number of data so their values are heterogeneous. In order to manipulate these lists, the weights obtained from each document must be normalized. Once values normalized we have to merge the terms into a single list unique for the whole corpus to compare the results. Clearly the precision will depend on the method used to perform such merging. We merged following three functions, which calculate respectively the sum(S), max(M) and average(A) of the measures values of the term in whole the corpus. At the end of this task we have three lists from Okapi and three lists from TFIDF. The notation for these lists are \(Okapi_X(a)\) and \(TFIDF_X(a)\), where \(a\) is the term, \(X\) the factor \(\in \{M, S, A\}\). For example, \(Okapi_M(a)\) is the value obtained by taking the maximum Okapi value for a term \(a\) in the whole corpus.

3.4. Computing the New Combined Measures

With aim of improving the precision of terms extraction we have conceived two new combined measures schemes, taking into account the values obtained in the above steps.

3.4.1. F-OCapi and F-TFIDF-C

Considered as the harmonic mean of the two used values, this method has the advantage of using all the values of the distribution.
We are extracting biomedical terminology.

\[ F - OCapi_x(a) = 2 \times \frac{Okapi_x(a) \times C - value(a)}{Okapi_x(a) + C - value(a)} \]  

(2)

\[ F - TFIDF - C_x(a) = 2 \times \frac{TFIDF_x(a) \times C - value(a)}{TFIDF_x(a) + C - value(a)} \]  

(3)

### 3.4.2. C-Okapi and C-TFIDF

For this measure, our assumption is that C-value can be more representative if the frequency, in the Equation (1), of the terms is replaced with a more significant value, in this case the Okapi’s or TFIDF’s values of the terms (over the whole corpus).

\[ C - m_x(a) = \begin{cases} 
  w(a) \times m_x(a), & \text{if } a \notin nested \\
  w(a) \times \left( m_x(a) - \frac{1}{|S_a|} \times \sum_{b \in S_a} m_x(b) \right), & \text{otherwise}
\end{cases} \]  

(4)

Where \( m_x(a) = \{ Okapi_x, TFIDF_x \} \), and \( X \in \{ M, S, A \} \).

Table 4 shows different ranking of terms with our system based on different measures. This example highlights specific and very relevant terms such as "antiphospholipid antibodies" and "platelet" which are explicitly better ranked with \( F-TFIDF-C_M \) (15,45); in comparison for "white blood" we get the opposite effect (796), because this candidate term is not a biomedical term by itself.

<table>
<thead>
<tr>
<th>Ranking of the terms</th>
<th>C-value</th>
<th>TFIDF_M</th>
<th>Okapi_M</th>
<th>F-TFIDF-C_M</th>
<th>F-OCapi_M</th>
<th>C-TFIDF_S</th>
<th>C-Okapi_S</th>
</tr>
</thead>
<tbody>
<tr>
<td>antiphospholipid antibodies</td>
<td>496</td>
<td>112</td>
<td>162</td>
<td>45</td>
<td>141</td>
<td>8</td>
<td>1770</td>
</tr>
<tr>
<td>white blood</td>
<td>129</td>
<td>745</td>
<td>387</td>
<td>796</td>
<td>356</td>
<td>679</td>
<td>754</td>
</tr>
<tr>
<td>platelet</td>
<td>159</td>
<td>112</td>
<td>112</td>
<td>15</td>
<td>59</td>
<td>219</td>
<td>800</td>
</tr>
</tbody>
</table>

Table 4: Ranking of terms based on different measures.

### 3.5. Re-ranking using Web-based Measure

After the extraction of terms we use a web-based measure to re-rank the candidate terms in order to augment the top \( k \) terms precision.

Different web mining studies focus on semantic similarity, semantic relatedness (Gracia et al., 2008). It means to quantify the degree in which some words or concepts are related, considering not only similarity but any possible semantic relationship among them. They are also used for multi-ontology disambiguation (Gracia et al., 2006). Web-based measures use web search engines to compute this similarity. One of the most well known web-based measure is the Normalized Google Distance (Cilibrasi et al., 2007).

Our web-based measure has for objective to tell us if a candidate term is a real biomedical term or not. It is specially appropriated for multi terms, as it computes the dependence
between the words of a term. In our case, we compute a “strict” dependence, which means the proximity of words of terms (i.e., neighboring words) is calculated with a strict restriction. In comparison with other web-based measures (Cilibrasi et al., 2007), WebR reduces the number of pages to consider by taking only into account web pages containing all the words of the terms. In addition, our measure can be easily adopted for all types of multi terms.

\[
\text{WebR}(a) = \frac{\text{num\_doc}("a")}{\text{num\_doc}(a)}
\]

(5)

Where \( a \) is the candidate term, \( \text{num\_doc}("a") \) the number of documents returned by the search engine with exact match only with multi term \( a \) (query with quotation marks “\( a \)”), \( \text{num\_doc}(a) \) the number of documents returned by the search engine including not exact match (query \( a \) without quotation marks), i.e., the whole documents containing the words of the multi term \( a \). For example, the multi term treponema pallidum, will generate 2 queries, the first \( \text{num\_doc}(\text{"treponema pallidum"}) \) which returns with Yahoo 1’100’000 documents, and the second query \( \text{num\_doc}(\text{treponema pallidum}) \) which returns 1’300’000 documents, then:

\[
\text{WebR}(\text{treponema pallidum}) = \frac{1\,100\,000}{1\,300\,000} = 0.85
\]

In our workflow we have tested Yahoo, Bing and Google, but WebR uses Yahoo because the results were the best. WebR re-rank the list of candidate terms returned by the combined measures. This is the final output of our workflow, on which we can evaluate precision taking the top \( k \) terms (P@k, in the following \( k = 60,300,900 \)), and compare them to results obtained either directly by baseline measures or by new combined measures without WebR.

In the following section, we evaluate a large list of extracted and ranked terms with our new measures and their different combinations.

4. Experiments and Results

4.1. Data and Experimental Protocol

We used biological laboratory tests, extracted from Lab Tests Online\(^4\) as corpus. This site provides information in several languages to patient or family caregiver on clinical lab tests. Each test which forms a document in our corpus, includes the formal lab test name, some synonyms and possible alternate names as well as a description of the test (at a glance, the test sample, the test and common questions), they are document that contain free text. Our extracted corpus contains 235 clinical tests (about 400 000 words) for English and 137 (about 210 000 words) for French.

In order to automatically validate our candidate terms we use as dictionaries UMLS for English and MeSH for French.

4.2. Experiments and Results

The evaluation was done automatically. Results are evaluated in terms of precision obtained over the top \( k \) terms (P@k, where \( k = 60,300,900 \)) at different steps of our workflow presented in previous section. Okapi and TFIDF provided three lists of ranked candidate terms (M, S, A). For each combined measure using Okapi or TFIDF, the experiments are

\(^4\)http://labtestsonline.org/
done with the three lists. Therefore, the number of ranked list to compare is \( C\)-value\( (1) + \) Okapi\( (3) + \) TFIDF\( (3) + \) F-OCapi\( (3) + \) F-TFIDF-C\( (3) + \) C-Okapi\( (3) + \) C-TFIDF\( (3) \) + WebR\( (1) = 20 \). In addition we experimented the workflow either for all (single and multi) or multi terms which finally gave 40 ranked lists.

The following paragraphs show part of the experiment results done for all (single- and multi-word terms) or multi terms. In the following we narrow down the presented results by keeping for the next workflow step only the best results.

### 4.2.1. Results obtained with baselines and new combined measures

<table>
<thead>
<tr>
<th></th>
<th>All Terms</th>
<th>Multi Terms</th>
<th>All Terms</th>
<th>Multi Terms</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>P@60</td>
<td>P@300</td>
<td>P@900</td>
<td>P@60</td>
</tr>
<tr>
<td>Okapi(_M)</td>
<td>0.96</td>
<td>0.95</td>
<td>0.82</td>
<td>0.68</td>
</tr>
<tr>
<td>Okapi(_S)</td>
<td>0.83</td>
<td>0.89</td>
<td>0.85</td>
<td>0.58</td>
</tr>
<tr>
<td>Okapi(_A)</td>
<td>0.72</td>
<td>0.31</td>
<td>0.27</td>
<td>0.48</td>
</tr>
<tr>
<td>TFIDF(_M)</td>
<td>0.97</td>
<td>0.96</td>
<td>0.84</td>
<td>0.71</td>
</tr>
<tr>
<td>TFIDF(_S)</td>
<td>0.96</td>
<td>0.95</td>
<td><strong>0.93</strong></td>
<td>0.82</td>
</tr>
<tr>
<td>TFIDF(_A)</td>
<td>0.78</td>
<td>0.74</td>
<td>0.63</td>
<td>0.50</td>
</tr>
<tr>
<td>C-value</td>
<td>0.88</td>
<td>0.92</td>
<td>0.89</td>
<td>0.72</td>
</tr>
<tr>
<td><strong>F − OCapi(_M)</strong></td>
<td>0.73</td>
<td>0.87</td>
<td>0.84</td>
<td>0.79</td>
</tr>
<tr>
<td><strong>F − TFIDF − C(_M)</strong></td>
<td><strong>0.98</strong></td>
<td><strong>0.97</strong></td>
<td><strong>0.86</strong></td>
<td><strong>0.98</strong></td>
</tr>
<tr>
<td>C − Okapi(_S)</td>
<td>0.88</td>
<td>0.86</td>
<td>0.80</td>
<td>0.61</td>
</tr>
<tr>
<td>C − TFIDF(_S)</td>
<td>0.96</td>
<td>0.95</td>
<td>0.86</td>
<td>0.85</td>
</tr>
</tbody>
</table>

Table 5: Extract of precision comparison for term extraction for English and French

Table 5 compares the precision between the best baselines measures and the best combined measures. Best results were obtained in general with \( F\)-TFIDF\(-C\(_M\)\) for English and \( F\)-OCapi\(_M\) for French. This table proves that the combined measures based on the harmonic mean are better than the baselines measures, and especially for multi word terms, for which the gain in precision reaches 16%. This result is particularly positive because in the biomedical domain it is often more interesting to extract multiword terms than single-word terms. However, one can notice that results obtained to extract all terms with C-Okapi\(_X\) and C-TFIDF\(_X\) are not better than Okapi\(_X\) or TFIDF\(_X\) use directly. The main reason for this is because the performance of those new combined measures are absorbed by the effect of extracting also single word terms. Definitively, all the new combined measures are really performing better for multi word terms.

Results of AKE methods for English show that TFIDF obtains better results than Okapi. The main reason for this, is because the size of the English corpus is larger than the French one, and Okapi is known to perform better when the corpus size is smaller (Lv et al., 2011).

In addition, Table 5 shows that C-value can be used to extract French biomedical terms with a better precision than what has been obtained in previous cited works with different languages. The precision of C-value for the previous work was between 26% and 31%.

### 4.2.2. WebR results

Our web mining approach is applied at the end of the process. With a number of terms small because the number of queries from an application to the search engines is limited, we took the lists with best results. The objective is re-rank the 300 terms of each list putting the “true”
terms at the top of the list, in this way the precision by intervals is improved. For this we had to choose the list, which got the best precision in the automatic validation.

Due to the restriction on the number of queries to search engines, it is more interesting for us first to evaluate the web measure with the French data. Table 15 shows the results between $F-OCapi_M$ and the WebR with automatic validation. We can see that WebR gets better results by intervals, this means true biomedical terms have a better ranking.

<table>
<thead>
<tr>
<th>Multi Terms with Automatic Validation</th>
<th>P@30</th>
<th>P@60</th>
<th>P@90</th>
<th>P@120</th>
<th>P@180</th>
<th>P@300</th>
</tr>
</thead>
<tbody>
<tr>
<td>$F - OCapi_M$</td>
<td>63.33%</td>
<td>65.00%</td>
<td>53.33%</td>
<td>49.17%</td>
<td>39.44%</td>
<td>34.67%</td>
</tr>
<tr>
<td>WebR (Yahoo)</td>
<td>80.00%</td>
<td>68.33%</td>
<td>61.11%</td>
<td>57.50%</td>
<td>47.22%</td>
<td>34.67%</td>
</tr>
</tbody>
</table>

Table 6: Precision comparison between $F-OCapi_M$ and WebR with automatic validation.

4.2.3. Discussion

Several terms proposed by our system are considered as irrelevant (i.e., false positive examples) with our automatic validation protocol because they are not present in known biomedical dictionaries, which does not mean that they are actually irrelevant. Indeed, elements that are not automatically validated can be considered relevant after the manual validation. For instance, they can represent new terms to add in biomedical ontologies or terminologies. Therefore, we proceed to a manual validation of the rest of the terms (i.e., the ones not found in the validation dictionary). For this, we gave a list of extracted terms to an user to validate manually. Table 7 shows the precision evaluated through human review for the best new combined measures for each language and for the web measure only for French. Note that the manual validation confirms that our ranking measure has a good behavior because the precision value is better for first terms. Table 7 also shows clearly that WebR gets better results by intervals than $F-Ocapi_M$.

<table>
<thead>
<tr>
<th>Multi Terms with Manual Validation</th>
<th>P@30</th>
<th>P@60</th>
<th>P@90</th>
<th>P@120</th>
<th>P@180</th>
<th>P@300</th>
</tr>
</thead>
<tbody>
<tr>
<td>English</td>
<td>100.00%</td>
<td>100.00%</td>
<td>99.17%</td>
<td>98.89%</td>
<td>96.67%</td>
<td>93.00%</td>
</tr>
<tr>
<td>$F - TFIDF - C_M$</td>
<td>100.00%</td>
<td>98.33%</td>
<td>95.56%</td>
<td>95.83%</td>
<td>95.00%</td>
<td>91.67%</td>
</tr>
<tr>
<td>French</td>
<td>100.00%</td>
<td>98.33%</td>
<td>97.78%</td>
<td>97.50%</td>
<td>95.56%</td>
<td>91.67%</td>
</tr>
<tr>
<td>WebR (Yahoo)</td>
<td>100.00%</td>
<td>98.33%</td>
<td>97.78%</td>
<td>97.50%</td>
<td>95.56%</td>
<td>91.67%</td>
</tr>
</tbody>
</table>

Table 7: Precision of $F-TFIDF-C_M$ for English and $F-OCapi_M$, WebR for French with manual validation.

We also have done experiments with two more corpus: (i) the Drugs data from MedlinePlus in English and, (ii) PubMed citations’ titles in English and French, we have verified that the new combined measures are performing better, particularly these based on the harmonic mean, $F-TFIDF-C_M$ and $F-OCapi_M$.

5. Conclusions and Perspectives

This work present a methodology for term extraction and ranking for two languages, French and English. We have adapted C-value to extract French biomedical terms, which was not proposed in the literature before.

We presented and evaluated two new measures obtained by combining three existing methods and another new web-based measure. The best results were obtained by combining C-value with the best results from AKE methods, i.e., $F-TFIDF-C_M$ and $F-OCapi_M$. 
Finally, *WebR* was applied to re-rank the best list of candidate terms to move “true” biomedical terms towards the top of list and thus to improve the P@k. The evaluation shows that *WebR* applied after *F-OCapiM* got the best precision for the extraction of French term.

The fact that we found false positive means that the term is not found in the validation data set. Then, we proposed a manual validation, for which the new precision results are very good and encouraging to use in terminology enrichment scenarios.

For our future evaluations, we will enrich our dictionaries with BioPortal’s terms for English and CISMeF’s terms for French. Our next task will be the extraction of relations between these new terms and already known terms, to help in ontology population. In addition, we are currently implementing a web application that implements these measures for the community.

Our work shows a comparison with some measures used in the literature, one of our objectives is to compare our work with a large number of measures on ATR for all domains and ATR applied to biomedicine in order to position our methodology regarding the others.

**Acknowledgments**

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