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► To cite this version:

Juan Antonio Lossio-Ventura, Clement Jonquet, Mathieu Roche, Maguelonne Teisseire. Biomedical term extraction: overview and a new methodology. Information Retrieval Journal, 2016, Medical Information Retrieval, 19 (1), pp.59-99. 10.1007/s10791-015-9262-2 . lirmm-01274539

HAL Id: lirmm-01274539

<https://hal-lirmm.ccsd.cnrs.fr/lirmm-01274539>

Submitted on 16 Feb 2016

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Biomedical Term Extraction: Overview and a New Methodology

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Received: date / Accepted: date

Abstract Terminology extraction is an essential task in domain knowledge acquisition, as well as for Information Retrieval (IR). It is also a mandatory first step aimed at building/enriching terminologies and ontologies. As often proposed in the literature, existing terminology extraction methods feature linguistic and statistical aspects and solve some problems related (but not completely) to term extraction, e.g. noise, silence, low frequency, large-corpora, complexity of the multi-word term extraction process. In contrast, we propose a cutting edge methodology to extract and to rank biomedical terms, covering the all mentioned problems. This methodology offers several measures based on linguistic, statistical, graphic and web aspects. These measures extract and rank candidate terms with excellent precision: we demonstrate that they outperform previously reported precision results for automatic term extraction, and work with different languages (English, French, and Spanish). We also demonstrate how the use of graphs and the web to assess the significance of a term candidate, enables us to outperform precision results. We evaluated our methodology on the biomedical GENIA and LabTestsOnline corpora and compared it with previously reported measures.

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Keywords Automatic Term Extraction · Biomedical Terminology Extraction · Natural Language Processing · BioNLP · Text Mining · Web Mining · Graphs

1 Introduction

The huge amount of biomedical data available today often consists of plain text fields, e.g. clinical trial descriptions, adverse event reports, electronic health records, emails or notes expressed by patients within forums [39]. These texts are often written using a specific language (expressions and terms) of the associated community. Therefore, there is a need for formalization and cataloging of these technical terms or concepts via the construction of terminologies and ontologies [52]. These technical terms are also important for Information Retrieval (IR), for instance when indexing documents or formulating queries. However, as the task of manually extracting terms of a domain is very long and cumbersome, researchers have strived to design automatic methods to assist knowledge experts in the process of cataloging the terms and concepts of a domain under the form of vocabularies, thesauri, terminologies or ontologies.

Automatic Term Extraction (ATE), or Automatic Term Recognition (ATR), is a domain which aims to automatically extract technical terminology from a given text corpus. We define technical terminology as the set of terms used in a domain. Term extraction is an essential task in domain knowledge acquisition because the technical terminology can be used for lexicon updating, domain ontology construction, summarization, named entity recognition or, as previously mentioned, IR.

In the biomedical domain, there is a substantial difference between existing resources (hereafter called *terminologies* or *ontologies*) in English, French, and Spanish. In English, there are about 9 919 000 terms associated with about 8 864 000 concepts such as those in UMLS¹ or BioPortal [44]. Whereas in French there are only about 330 000 terms associated with about 160 000 concepts [41], and in Spanish 1 172 000 terms associated with about 1 140 000 concepts. Note the strong difference in the number of ontologies and terminologies available in French or Spanish. This makes ATE even more important for these languages.

In biomedical ontologies, different terms may be linked to the same concept and are semantically similar with different writing, for instance “*neoplasm*” and “*cancer*” in MeSH or SNOMED-CT. Ontologies also contain terms with morphosyntactic variants, for instance plurals like “*external fistula*” and “*external fistulas*”, and this group of variants is linked to a preferred term. As one of our goals is to extract new terms to enrich ontologies, our approach does not normalize variant terms, mainly because normalization would lead to penalization in extracting new variant terms. Technical terms are useful to gain further insight into the conceptual structure of a domain. These may be: (i)

¹ http://www.nlm.nih.gov/research/umls/knowledge_sources/metathesaurus/release/statistics.html

single-word terms (simple), or (ii) multi-word terms (complex). The proposed study focuses on both cases.

Term extraction methods usually involve two main steps. The first step extracts candidate terms by unithood calculation to qualify a string as a valid term, while the second step verifies them through termhood measures to validate their domain specificity. Formally, unithood refers to the degree of strength or stability of syntagmatic combinations and collocations, and termhood is defined as the degree to which a linguistic unit is related to domain-specific concepts [26]. ATE has been applied to several domains, e.g. biomedical [34, 18, 63, 42], ecological [12], mathematical, [56], social networks [31], banking [17], natural sciences [17], information technology [42, 60], legal [60], as well as post-graduate school websites [48].

The main issues in ATE are: (i) extraction of non-valid terms (noise) or omission of terms with low frequency (silence), (ii) extraction of multi-word terms having various complex various structures, (iii) manual validation efforts of the candidate terms [12], and (iv) management of large-scale corpora. Inspired by our previously published results and in response to the above issues, we propose a cutting edge methodology to extract biomedical terms. We propose new measures and some modifications of existing baseline measures. Those measures are divided into: 1) ranking measures, and 2) re-ranking measures. Our ranking measures are statistical- and linguistic-based and address issues i), ii) and iv). Our two re-ranking measures the first one called *TeR-Graph* is a graph-based measure which deals with issues i), ii) and iii). The second one, called *WAHI*, is a web-based measure which also deals with issues i), ii) and iii). The novelty of the *WAHI* measure is that it is web-based which has, to the best of our knowledge, never been applied within ATE approaches.

The main contributions of our article are: (1) enhanced consideration of the term unithood, by computing a degree of quality for the term unithood, and, (2) consideration of the term dependence in the ATE process. The quality of the proposed methodology is highlighted by comparing the results obtained with the most commonly used baseline measures. Our evaluation experiments were conducted despite difficulties in comparing ATE measures, mainly because of the size of the corpora used and the lack of available libraries associated with previous studies. Our three measures improve the process of automatic extraction of domain-specific terms from text collections that do not offer reliable statistical evidence (i.e. low frequency).

The paper is organized as follows. We first discuss related work in Section 2. Then the methodology to extract biomedical terms is detailed in Section 3. The results are presented in Section 4, followed by discussions in Section 5. Finally, the conclusions in Section 6.

2 Related Work

Recent studies have focused on multi-word (n-grams) and single-word (uni-grams) term extraction. Term extraction techniques can be divided into four

broad categories: (i) *Linguistic*, (ii) *Statistical*, (iii) *Machine Learning*, and (iv) *Hybrid*. All of these techniques are encompassed in Text Mining approaches. Graph-based approaches have not yet been applied to ATE, although they have been successively adopted in other Information Retrieval fields and could be suitable for our purpose. Existing web techniques have not been applied to ATE but, as we will see, these techniques can be adapted for such purposes.

2.1 Text Mining approaches

2.1.1 Linguistic approaches

These techniques attempt to recover terms via linguistic pattern formation. This involves building rules to describe naming structures for different classes based on orthographic, lexical, or morphosyntactic characteristics, e.g. [20]. The main approach is to develop rules (typically manually) describing common naming structures for certain term classes using orthographic or lexical clues, or more complex morpho-syntactic features. Moreover, in many cases, dictionaries of typical term constituents (e.g. terminological heads, affixes, and specific acronyms) are used to facilitate term recognition [30]. A recent study on biomedical term extraction [21] is based on linguistic patterns plus additional context-based rules to extract candidate terms, which are not scored and the authors leave the term relevance decision to experts.

2.1.2 Statistical methods

Statistical techniques chiefly rely on external evidence presented through surrounding (contextual) information. Such approaches are mainly focused on the recognition of general terms [59]. The most basic measures are based on frequency. For instance, *term frequency* (tf) counts the frequency of a term in the corpus, *document frequency* (df) counts the number of documents where a term occurs, and *average term frequency* (atf), which is $\frac{tf}{df}$.

A similar research topic, called Automatic Keyword Extraction (AKE), proposes to extract the most relevant words or phrases in a document using automatic indexation. Keywords, which we define as a sequence of one or more words, provide a compact representation of a document's content. Such measures can be adapted to extract terms from a corpus as well as ATE measures. We take two popular AKE measures as baselines measures, i.e. *Term Frequency Inverse Document Frequency* ($TF-IDF$) [53], and *Okapi BM25* [49] (hereafter *Okapi*), these weight the word frequency according to their distribution along the corpus. *Residual inverse document frequency* ($RIDF$) compares the document frequency to another chance model where terms with a particular term frequency are distributed randomly throughout the collection, while *Chi-square* [37] assesses how selectively words and phrases co-occur within the same sentences as a particular subset of frequent terms in the document text. This is applied to determine the bias of word co-occurrences in the document

text, which is then used to rank words and phrases as keywords of the document; *RAKE* [50] hypothesises that keywords usually consist of multiple words and do not contain punctuation or stop words. It uses word co-occurrence information to determine the keywords.

2.1.3 Machine Learning

Machine Learning (ML) systems are often designed for specific entity classes and thus integrate term extraction and term classification. Machine Learning systems use training data to learn features useful for term extraction and classification. But the availability of reliable training resources is one of the main problems. Some proposed ATE approaches use machine learning [12, 62, 42]. However, ML may also generate noise and silence. The main challenge is how to select a set of discriminating features that can be used for accurate recognition (and classification) of term instances. Another challenge concerns the detection of term boundaries, which are the most difficult to learn.

2.1.4 Hybrid methods

Most approaches combine several methods (typically linguistic and statistically based) for the term extraction task. *GlossEx* [29] considers the probability of a word in the domain corpus divided by the probability of the appearance of the same word in a general corpus. Moreover, the importance of the word is increased according to its frequency in the domain corpus. *Weirdness* [1] considers that the distribution of words in a specific domain corpus differs from that in a general corpus. *C/NC-value* [18] combines statistical and linguistic information for the extraction of multi-word and nested terms. This is the most well-known measure in the literature. While most studies address specific types of entities, *C/NC-value* is a domain-independent method. It has also been used for recognizing terms in the biomedical literature [24, 22]. In [63], the authors showed that *C-value* obtains the best results compared to the other measures cited above. *C-value* has been also modified to extract single-word terms [40], and in this work the authors extract only terms composed of nouns. Moreover, *C-value* has also been applied to different languages other than English, e.g. Japanese, Serbian, Slovenian, Polish, Chinese [25], Spanish [4], Arabic, and French. We have thus chosen *C-value* as one of our baseline measure. Those baseline measures will be modified and evaluated with the new proposed measures.

Terminology Extraction from Parallel and Comparable Corpora

Another kind of approach suggests that terminology may be extracted from parallel and/or comparable corpora. Parallel corpora contain texts and their translation into one or more languages, but such corpora are scarce [9]. Thus parallel corpora are scarce for specialized domains. Comparable corpora are those which select similar texts in more than one language or variety [15].

Comparable corpora are built more easily than parallel corpora. They are often used for machine translation and their approaches are based on linguistics, statistics, machine learning, and hybrid methods. The main objective of these approaches is to extract translation pairs from parallel/comparable corpora. Different studies propose translation of biomedical terms for English-French by alignment techniques [16]. English-Greek and English-Romanian bilingual medical dictionaries are also constructed with a hybrid approach that combines semantic information and term alignments [28]. Other approaches are applied for single- and multi-word terms with English-French comparable corpora [14]. The authors use statistical methods to align elements by exploiting contextual information. Another study proposes to use graph-based label propagation [57]. This approach is based on a graph for each language (English and Japanese) and the application of a similarity calculus between two words in each graph. Moreover, some machine learning algorithms can be used, e.g. the logistic regression classifier [27]. There are also approaches that combine both corpora [38] (i.e. parallel and comparable) in an approach to reinforce extraction. Note that our corpora are not parallel and are far of being comparable because of the difference in their size. Therefore these approaches are not evaluated in our study.

2.1.5 Tools and applications for biomedical term extraction

There are several applications implementing some measures previously mentioned, especially *C-value* for biomedical term extraction. The study of related tools revealed that most existing systems that especially implement statistical methods are made to extract keywords and, to a lesser extent, to extract terminology from a text corpus. Indeed, most systems take a single text document as input, not a set of documents (as corpus), for which the *IDF* can be computed. Most systems are available only in English and the most relevant for the biomedical domain are:

- *TerMine*², developed by the authors of the *C-value* method, only for English term extraction;
- *Java Automatic Term Extraction*³ [63], a toolkit which implements several extraction methods including *C-value*, GlossEx, TermEx and offer other measures such as frequency, average term frequency, *IDF*, *TF-IDF*, *RIDF*;
- *FlexiTerm*⁴ [55], a tool explicitly evaluated on biomedical corpora and which offer more flexibility than *C-value* when comparing term candidates (treating them as bag of words and ignoring the word order);
- *BioYaTeA*⁵ [21], is a version of the YaTeA term extractor [2], both are available as a Perl module. It is a biomedical term extractor. The method used is based only on linguistic aspects.

² <http://www.nactem.ac.uk/software/termine/>

³ <https://code.google.com/p/jatetoolkit/>

⁴ <http://users.cs.cf.ac.uk/I.Spasic/flexiterm/>

⁵ <http://search.cpan.org/~bibliome/Lingua-BioYaTeA/>

- *BioTex*⁶ [32], only for biomedical terminology extraction. It is available for online testing and assessment but can also be used in any program as a Java library (POS tagger not included). In contrast to other existing systems, this system allows us to analyze French and Spanish corpora, manually validate extracted terms and export the list of extracted terms.

2.2 Graph-based approaches

Graph modeling is an alternative for representing information, which clearly highlights relationships of nodes among vertices. It also groups related information in a specific way, and a centrality algorithm can be applied to enhance their efficiency. Centrality in a graph is the identification of the most important vertices within a graph. A host of measures have been proposed to analyze complex networks, especially in the social network domain [7,8,3]. Freeman [19], formalized three different measures of node centrality: degree, closeness and betweenness. Degree is the number of neighbors that a node is connected to. Closeness is the inverse sum of shortest distances to all other neighbor nodes. Betweenness is the number of shortest paths from all vertices to all others that pass through that node. One study proposes to take the number of edges and their weights into account [45], since the three last measures do not do this. Another well known measure is PageRank [46], which ranks websites. Boldi [6], evaluated the behavior of ten measures, and associated the centrality to the node with largest degree. Our approach proposes the opposite, i.e. we focus on nodes with a lower degree. An increasingly popular recent application of graph approaches to IR concerns social or collaborative networks and recommender systems [43,3].

Graph representations of text and scoring function definition are two widely explored research topics, but few studies have focused on graph-based IR in terms of both document representation and weighting models [51]. First, text is modeled as a graph where nodes represent words and edges represent relations between words, defined on the basis of any meaningful statistical or linguistic relation [5]. In [5], the authors developed a graph-based word weighting model that represents each document as a graph. The importance of a word within a document is estimated by the number of related words and their importance, in the same way that PageRank [46] estimates the importance of a page via the pages that are linked to it. Another study introduces a different representation of document that captures relationships between words by using an unweighted directed graph of words with a novel scoring function [51].

In the above approaches, graphs are used to measure the influence of words in documents like automatic keyword extraction methods (AKE), while ranking documents against queries. These approaches differ from ours as they use graphs focused on the extraction of relevant words in a document and computing relations between words. In our proposal, a graph is built such that

⁶ <http://tubo.lirmm.fr/biotex/>

the vertices are multi-word terms and the edges are relations between multi-word terms. Moreover, we focus especially on a scoring function of relevant multi-word terms in a domain rather than in a document.

2.3 Web Mining approaches

Different web mining studies focus on semantic similarity, semantic relatedness. This means quantifying the degree to which some words are related, considering not only similarity but also any possible semantic relationship among them. The word association measures can be divided into three categories [10]: (i) *Co-occurrence measures* that rely on co-occurrence frequencies of both words in a corpus, (ii) *Distributional similarity-based measures* that characterize a word by the distribution of other words around it, and (iii) *Knowledge-based measures* that use knowledge-sources like thesauri, semantic networks, or taxonomies [23]. In this paper, we focus on co-occurrence measures because our goal is to extract multi-word terms and we suggest computing a degree of association between words composing a term. Word association measures are used in several domains like ecology, psychology, medicine, and language processing, and were recently studied in [47,61], such as *Dice*, *Jaccard*, *Overlap*, *Cosine*. Another measure to compute the association between words using web search engines results is the Normalized Google Distance [11], which relies on the number of times words co-occur in the document indexed by an information retrieval system. In this study, experimental results with our web-based measure will be compared with the basic measures (*Dice*, *Jaccard*, *Overlap*, *Cosine*).

3 Methodology

This section describes the baseline measures, their modifications as well as new measures that we propose for the biomedical term extraction task. The principle of our approach is to assign a weight to a term, which represents the appropriateness of being a relevant biomedical term. This allows to give as output a list ranked by their appropriateness. Our methodology for automatic term extraction has three main steps plus an additional step (a), described in Figure 1, and in the sections hereafter:

- (a) Pattern Construction,
- (1) Candidate Term Extraction,
- (2) Ranking of Candidate Terms,
- (3) Re-ranking.

Pattern Construction (step a)

As previously cited, we supposed that biomedical terms have a similar syntactic structure (linguistic aspect). Therefore, we built a list of the most com-

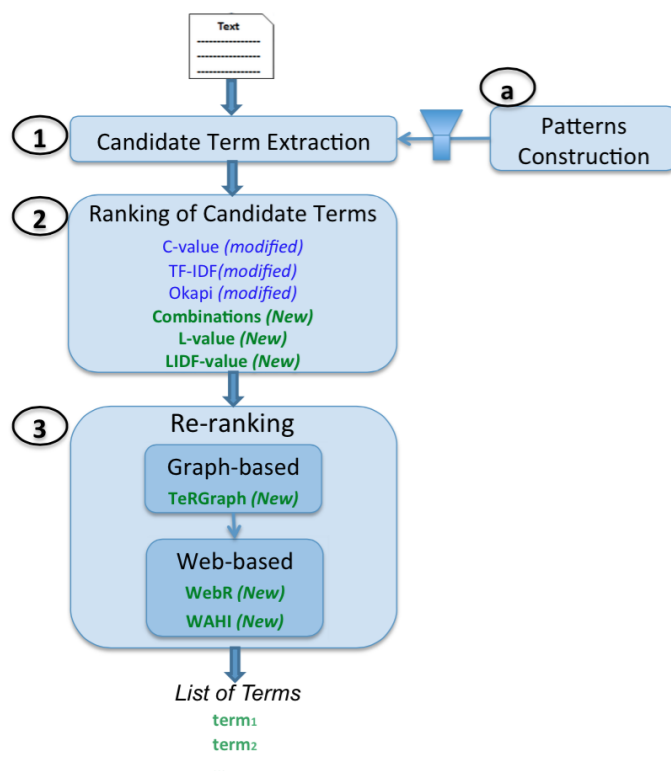


Fig. 1 Workflow Methodology for Biomedical Term Extraction.

mon linguistic patterns according to the syntactic structure of terms present in the UMLS⁷ (for English and Spanish), and the French version of MeSH⁸, SNOMED International and the rest of the French content in the UMLS.

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 in which it appears. This is highly time-consuming, so we conducted automatic part-of-speech tagging.

We evaluated three tools (TreeTagger⁹, Stanford Tagger¹⁰ and Brill's rules¹¹). This evaluation was carried out throughout the entire workflow with the three tools and we assessed the precision of the extracted terms. We noted that in general TreeTagger gave the best results for Spanish and French. Meanwhile, for English, the Stanford tagger and TreeTagger gave similar results. We finally chose TreeTagger, which gave better results and may be used for English, French and Spanish. Moreover, our choice was validated with regard to a recent

⁷ <http://www.nlm.nih.gov/research/umls>

⁸ <http://mesh.inserm.fr/mesh/>

⁹ <http://www.cis.uni-muenchen.de/~schmid/tools/TreeTagger/>

¹⁰ <http://nlp.stanford.edu/software/tagger.shtml>

¹¹ http://en.wikipedia.org/wiki/Brill_tagger

comparison study [58], wherein the authors showed that TreeTagger generally gives the best results, particularly for nouns and verbs.

Therefore, we carried out automatic part-of-speech tagging of the biomedical terms using TreeTagger, and then computed the frequency of the syntactic structures. Patterns among the 200 highest frequencies were selected to build the list of patterns for each language. From this list, we also computed the weight (probability) associated with each pattern, i.e. the frequency of the pattern over the sum of frequencies (see Algorithm 1), but this weight will only be used for one measure. The number of terms used to build these lists of patterns was 3 000 000 for English, 300 000 for French, and 500 000 for Spanish, taken from the previously mentioned terminologies. Table 1 illustrates the computation of the linguistic patterns and their weights for English.

Different terminology extraction studies are based on the use of regular expressions to extract candidate terms, for instance [18]. Generally these regular expressions are manually built for a specific language and/or domain [13]. In our setting, we prefer to (i) construct and (ii) apply patterns in order to extract terms in texts. These patterns have the advantage of being generic because they are based on defined PoS tags. Moreover, they are very specific because they are (automatically) built with specialized biomedicine resources. Concerning this last point, we can consider we are close to the use of regular expressions. There are two main reasons that we use specific linguistic patterns. First, we would like to restrict the patterns to the biomedical domain. For instance, biomedical terms often contain numbers in their syntactic structure, and this is very specific to the biomedical domain, e.g. “*epididymal protein 9*”, “*pargyline 10 mg*”. General patterns do not enable extraction of such terms. Our methodology is based on 200 significant patterns for English, French, or Spanish, yet different for each language. For instance, there are 55 patterns for English that contain numbers in the linguistic structure. Thus, this kind of pattern seems quite relevant for this domain. The second reason for using lexical patterns is that we assign a probability of occurrence to each pattern, which would not be possible with classical patterns and regular expressions.

3.1 Candidate Term Extraction (step 1)

The first main step is to extract the candidate terms. So we apply part-of-speech to the whole corpus using TreeTagger. Then we filter out the content of our input corpus using previously computed patterns. We select only terms whose syntactic structure is in the patterns list. The pattern filtering is specifically done on a per-language basis (i.e. when the text is in French, only the French list of patterns is used).

3.2 Ranking of Candidate Terms (step 2)

We need to select the most appropriate terms for the biomedical domain. Candidate term ranking is therefore essential. For this purpose, several measures

Algorithm 1: ComputePatterns (*Dictionary*, *np*)

Data: *Dictionary* = dictionary of a domain, *np* = number of patterns to use
Result: $HT_{patterns}(pattern, probability)$ = Hashtable of the first *np* linguistic patterns with its probability

```

begin
   $HT_{patterns} \leftarrow \emptyset$ ;
   $HT_{aux}(tag, freq) \leftarrow \emptyset$  // Hashtable of the tag of each term with its frequency ;
   $sizeHT \leftarrow$  number of terms in Dictionary;
   $freq_{total} \leftarrow 0$  ;
   $probability \leftarrow 0.0$  ;
  Tag of the Dictionary;
  for tag of each term  $\in$  Dictionary do
    if tag  $\in$   $HT_{aux}$  then
      | update  $HT_{aux}(tag, freq + 1)$ ;
    else
      | add  $HT_{aux}(tag, 1)$ ;
    end
  end
  Rank  $HT_{aux}(tag, freq)$  by the freq;
   $freq_{total} \leftarrow \sum_{i=1}^{np} freq(HT_{aux}(i))$ ;
  for  $i = 1; i \leq np; i++$  do
    |  $probability \leftarrow \frac{freq(HT_{aux}(i))}{freq_{total}}$  ;
    | add  $HT_{patterns}(tag(HT_{aux}(i)), probability)$ ;
  end
end

```

Pattern	Frequency	Probability
NN IN JJ NN IN JJ NN	3006	$3006/4113 = 0.73$
NN CD NN NN NN	1107	$1107/4113 = 0.27$
	4113	1.00

Table 1 Example of pattern construction (where *NN* is a noun, *IN* a preposition or subordinating conjunction, *JJ* an adjective, and *CD* a cardinal number)

are proposed and Figure 1(2) shows the set of available measures. We propose some modifications of the most known measures in the literature (i.e., *C-value*, *TF-IDF*, *Okapi*), and propose new ones (i.e., *F-TFIDF-C*, *F-OCapi*, *LIDF-value*, *L-value*). Those measures are linguistic- and statistic- based, they are also not very time-consuming. In this step, only one measure will be selected to perform the ranking. The measures of this section take a list of candidate terms previously filtered by linguistic patterns as input, which makes it possible to assess less invalid terms while dealing with the noise problem. In addition to the use of linguistic patterns to alleviate the problem of the extraction of multi-word terms having various complex structures. Moreover, the frequency decreases the number of invalid terms to evaluate (noise). The measures mentioned above are effective on large amounts of data [36,35,54], which overcomes the problem of large-scale corpora. Hereafter we describe all measures.

3.2.1 *C-value*

The *C-value* method combines linguistic and statistical information [18]. 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 the frequency of terms to compute the *termhood* (i.e. the association strength of a term to domain concepts). The *C-value* method aims to improve the extraction of long terms, and it was specially built for extracting multi-word terms.

$$C\text{-value}(A) = \begin{cases} w(A) \times f(A) & \text{if } A \notin \text{nested} \\ w(A) \times \left(f(A) - \frac{1}{|S_A|} \times \sum_{b \in S_A} f(b) \right) & \\ \text{otherwise} & \end{cases} \quad (1)$$

Where A is the candidate term, $w(A) = \log_2(|A|)$, $|A|$ the number of words in A , $f(A)$ the frequency of A in the unique document, S_A the set of terms that contain A and $|S_A|$ the number of terms in S_A . In a nutshell, *C-value* uses either the frequency of the term if the term is not included in other terms (first line), or decreases this frequency if the term appears in other terms, based on the frequency of those other terms (second line).

We modified the measure in order to extract all terms (single-word + multi-words terms), as also suggested in [4], but in a different manner.

The original *C-value* defines $w(A) = \log_2(|A|)$, and we modified $w(A) = \log_2(|A| + 1)$ in order to avoid null values for single-word terms, as illustrated in Table 2. Note that we do not use a stop word list or a frequency threshold as was originally proposed.

	Original <i>C-value</i>	Modified <i>C-value</i>
	$w(A) = \log_2(A)$	$w(A) = \log_2(A + 1)$
antiphospholipid antibodies	$\log_2(2) = 1$	$\log_2(2 + 1) = 1, 6$
white blood	$\log_2(2) = 1$	$\log_2(2 + 1) = 1, 6$
platelet	$\log_2(1) = 0$	$\log_2(1 + 1) = 1$

Table 2 Calculation of $w(A)$

3.2.2 *TF-IDF and Okapi*

These measures are used to associate a weight to each term in a document [53]. This weight represents the term relevance for the document. The output is a ranked list of terms for each document, which is often used in information retrieval so as to order documents by their importance for a given query [49].

Okapi can be seen as an improvement of the *TF-IDF* measure, while taking the document length into account.

$$TF-IDF(A, d, D) = tf(A, d) \times idf(A, d) \quad (2)$$

$$tf(A, d) = \frac{f(A, d)}{\max\{f(A, d) : w \in d\}}$$

$$idf(A, d) = \log \frac{|D|}{|\{d \in D : A \in d\}|}$$

$$Okapi(A, d, D) = tf_{BM25}(A, d) \times idf_{BM25}(A, d) \quad (3)$$

$$tf_{BM25}(A, d) = \frac{tf(A, d) \times (k_1 + 1)}{tf(A, d) + k_1 \times (1 - b + b \times \frac{dl(d)}{dl_{avg}})}$$

$$idf_{BM25}(A, d) = \log \frac{|D| - dc(A) + 0.5}{dc(A) + 0.5}$$

Where A is a term, considering d a document, D the collection of documents, $f(A, d)$ the frequency of A in d , $tf(A, d)$ the term frequency of A in d , $idf(A, D)$ the inverse document frequency of A in D , $dc(t)$ the number of documents containing term A , this means: $|\{d \in D : t \in d\}|$, $dl(d)$ the length of the document d in number of words, dl_{avg} the average document length of the collection.

As the output is a ranked list of terms per document, we could find the same term in different documents, with different weights in each document. So we need to merge the term into a single list. For this, we propose to merge them according to three functions, which respectively calculate the sum(S), max(M) and average(A) of the weights of a term. At the end of this task, we have three lists from *Okapi* and three lists from *TF-IDF*. The notation for these lists are $Okapi_X(A)$ and $TF-IDF_X(A)$, where A is the term, and 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. Figure 2 shows the merging process.

With aim of improving the term extraction precision, we designed two new combined measures, while taking the values obtained in the above steps into account. Both are based on harmonic means of two values.

3.2.3 Combinations: *F-OCapi* and *F-TFIDF-C*

Considered as the harmonic mean of the two used values, this method has the advantage of using all values of the distribution.

$$F-OCapi_X(A) = 2 \times \frac{Okapi_X(A) \times C-value(A)}{Okapi_X(A) + C-value(A)} \quad (4)$$

$$F-TFIDF-C_X(A) = 2 \times \frac{TFIDF_X(A) \times C-value(A)}{TFIDF_X(A) + C-value(A)} \quad (5)$$

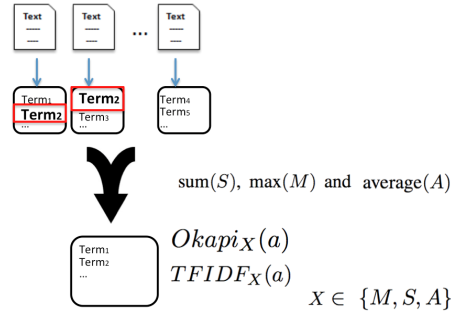


Fig. 2 Merging lists.

3.2.4 LIDF-value and L-value

In this section we present two new measures. The first one, called *LIDF-value* (**L**inguistic patterns, **IDF**, and **C-value** information). *LIDF-value* is partially presented in [34]. This is a new ranking measure based on linguistic and statistical information.

Our method *LIDF-value* is aimed at computing the termhood for each term, using the *linguistic* information calculated as described below, the *idf*, and the *C-value* of each term. The *linguistic* information gives greater importance to the term unithood in order to detect low frequency terms. So we associate the pattern weight (see Table 1) with the candidate term *probability*. That means the *probability* of a candidate term of being a relevant biomedical term. The *probability* is associated only if the syntactic structure of the term appears in the linguistic pattern list.

The inverse document frequency (*idf*) is a measure indicating the extent to which a term is common or rare across all documents. It is obtained by dividing the total number of documents by the number of documents containing the term, and then by taking the logarithm of that quotient. The *probability* and *idf* improve low frequency term extraction. The objective of these two components is to tackle the silence problem, allowing extraction of discriminant terms, for instance, in a biomedical corpus, “*virus production*” with low frequency being better ranked than “*human monocytic cell*”, which has a higher frequency. This means that for a low frequency candidate term, its score can be favored if its linguistic pattern is associated with a high probability and/or its *idf* value is also high. The *C-value* measure is based on the term frequency. The *C-value* (see formula 1) measure favors a candidate term that does not often appear in a longer term. For instance, in a specialized corpus (Ophthalmology), the authors of [18] found the irrelevant term “*soft contact*” while the frequent and longer term “*soft contact lens*” is relevant.

As an example, we implement the Algorithm 2, which describes the applied process. These different statistical information items (i.e. *probability* of linguistic patterns, *C-value*, *idf*) are combined to define the global ranking measure *LIDF-value* (see formula 6); where $P(A_{LP})$ is the probability of a

term A which has the same linguistic structure pattern LP , i.e. the weight of the linguistic pattern LP computed in Subsection *Pattern Construction*.

$$LIDF-value(A) = P(A_{LP}) \times idf(A) \times C-value(A) \quad (6)$$

Algorithm 2: ComputeLIDF-value ($Corpus$, $Patterns$, min_{freq} , num_{terms})

Data: $Corpus$ = set of documents of a specific-domain;
 $Patterns = HT_{patterns}(pattern, probability)$ //Hashtable of linguistic patterns with its probability;
 min_{freq} = frequency threshold for candidate terms;
 num_{terms} = number of terms to take as output
Result: L_{terms} = List of ranked terms
begin
 Tag the $Corpus$;
 Take the *lemma* of each tagged word;
 Extract candidate terms A by filtering with $Patterns$;
 Remove candidate terms A below min_{freq} ;
 for each candidate term $A \in Corpus$ **do**
 $LIDF-value(A) = P(A_{LP}) \times idf(A) \times C-value(A)$;
 add A to L_{terms} ;
 end
 Rank L_{terms} by the value obtained with $LIDF-value$;
 Select the first num_{terms} terms of L_{terms} ;
end

Note that $LIDF-value$ works only for a set of documents, mainly because the idf measure can only be computed on a set of documents (see formula 2). Therefore, for datasets composed of one document, we propose a new measure, $L-value$, as explained in the following paragraphs.

$L-value$ is a variant of $LIDF-value$, focused on one document with the goal of benefiting from the *probability* of linguistic patterns computed for $LIDF-value$. This measure does not contain the idf (see formula 7). $L-value$ is interesting to highlight the more representative terms of a single corpus without considering the discriminative aspects, e.g. idf . This measure gives another point of view and is complementary to those based on the idf weighting.

A single document can be considered as a free text without delimitation. For instance, a scientist article, a book, a document created with titles/abstracts from a library database. $L-value$ becomes interesting when it does not exist a considerable amount of data for a new subject, i.e. an emergent term in the community. For instance, the “Ataxia Neuropathy Spectrum” term appears only in 4 titles/abstracts of scientist articles from PubMed¹² between 2009 and 2015. PubMed is a free search engine accessing primarily the MEDLINE database of references and abstracts on life sciences and biomedical topics.

¹² <http://www.ncbi.nlm.nih.gov/pubmed>

$$L\text{-value}(A) = P(A_{LP}) \times C\text{-value}(A) \quad (7)$$

3.3 Re-ranking (step 3)

After the term extraction, we propose new measures to re-rank the candidate terms in order to increase the top k term precision. The re-ranking measures aim to improve the term extraction results of ranking measures. This involves positioning the most relevant biomedical terms at the top of the list. That provides more confidence that the terms appearing at the top of this list are true biomedical terms.

These re-ranking functions represent an extension of the measures presented in [33]. Therefore, as improvements, we propose to take graph-theoretic information into account to highlight relevant terms, as well as web information, as explained in the following subsections. These measures can be executed separately, but the graph construction is time consuming, and the number of search engine queries is limited. Therefore, we just apply these measures for a group of selected terms given by a ranking measure. Because the ranking measures have proved to be more efficient applied before than *TeRGraph* and *web*-based measures.

As these measures are applied to the list of terms obtained with a ranking measure, which tackles noise, silence and multi-word term extraction problems, so they also take into account those problems. As mentioned, the objective of re-ranking measures is to re-rank terms, so the manual validation efforts of the candidate terms decrease because the relevant biomedical term is allocated at the top of the list.

3.3.1 A new graph-based ranking measure: “*TeRGraph*” (*Terminology Ranking based on Graph information*)

This approach aims to improve the ranking (and therefore the precision results) of extracted terms. As mentioned above, in contrast to the above-cited study, the graph is built with a list of terms obtained according to a measure described in Section 3.2, where vertices denote terms linked by their co-occurrence in sentences in the corpus. Moreover, we make the hypothesis that the term representativeness in a graph, for a specific-domain, depends on its number of neighbors, and the number of neighbors of its neighbors. We assume that a term with more neighbors is less representative of the specific domain. This means that this term is used in the general domain. Figure 3 illustrates our hypothesis.

The graph-based approach is divided into two steps:

- (i) **Graph construction:** a graph (see Figure 5) is built where vertices denote terms, and edges denote co-occurrence relations between terms, co-occurrences between terms are measured as the weight of the relation in the initial corpus. This approach is statistical because it links all co-occurring

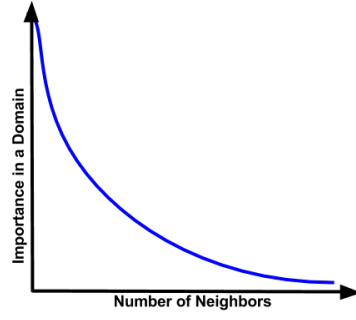


Fig. 3 Importance of a term in a domain

terms without considering their meaning or function in the text. This graph is undirected as the edges imply that terms simply co-occur, without any further distinction regarding their role. We take the *Dice coefficient*, a basic measure to compute the co-occurrence between two terms x and y , as defined by the following formula:

$$D(x, y) = \frac{2 \times P(x, y)}{P(x) + P(y)} \quad (8)$$

- (ii) **Representativeness computations on the term graph:** a principled graph-based measure to compute term weights (representativeness) is defined. The aim of this new graph-based ranking measure, *TeRGraph*, see Equation 9, is to derive these weights for each vertex, (i.e. multi-word term weight), in order to re-rank the list of extracted terms.

$$TeRGraph(A) = \log_2 \left(k + \frac{1}{1 + |N(A)| + \sum_{T_i \in N(A)} |N(T_i)|} \right) \quad (9)$$

Where A represents a vertex (term), $N(A)$ the neighborhood of A , $|N(A)|$ the number of neighbors of A , T_i the neighbor i of A , and k a constant. The intuition for Equation 9 is as follows: the more a term A has neighbors (directly with $N(A)$ or by transitivity with $N(T_i)$), the more the weight decreases. Indeed, a term A having a lot of neighbors is considered too general for the domain (i.e. this term is not salient), so it has to be penalized via the associated score.

The k constant affects the *TeRGraph* value, i.e. the set of values that *TeRGraph* takes when k changes. For instance, when $k = 0.5$, the set of values for *TeRGraph* is between -1 and 0 , (i.e., $TeRGraph \in [-1, 0]$), and when $k = 1$, $TeRGraph \in [0, 0.6]$. As the values taken by *TeRGraph* are different, then the slope of the curve is also different. Figure 4 shows the behavior of *TeRGraph* when k changes. According the experiments, we have chosen

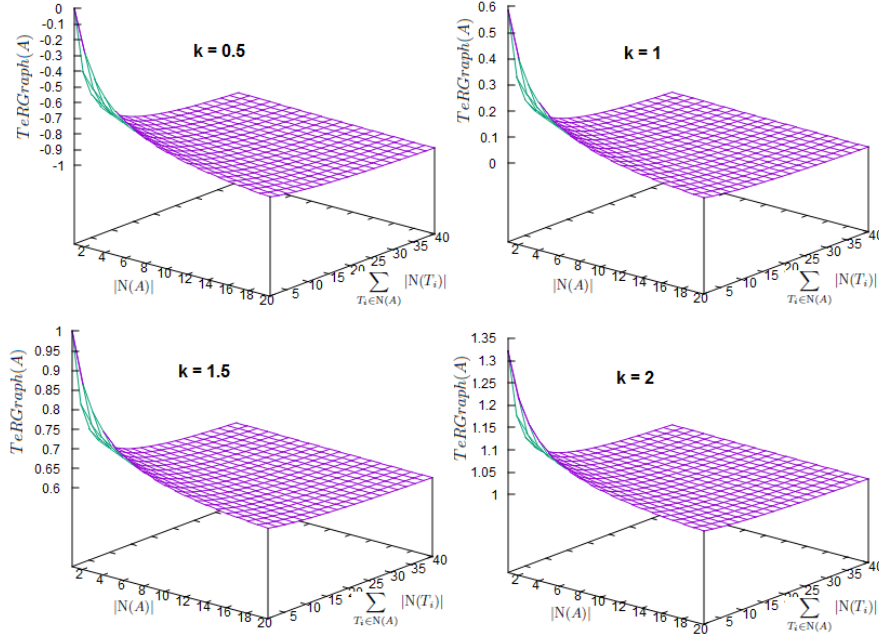


Fig. 4 TeRGraph's value for $k = \{0.5; 1; 1.5; 2\}$

$k = 1.5$. The main reason is that the slope of the curve is low, and the set of values for *TeRGraph* ranges from 0.6 to 1.

See Algorithm 3 for more details, it describes the entire process: (1) co-occurrence graph construction, (2) computation of the representativeness of each vertex.

Figure 5 shows an example to calculate the value of *TeRGraph* for a term in different graphs. These graphs are built with different co-occurrence thresholds (i.e. Dice's value between two terms). In this example, A_1 and A_2 represent the term *chloramphenicol acetyltransferase reporter* in Graphs 1 and 2, respectively.

3.3.2 WebR

The aim of our web-based measure, to predict with a better confidence if a candidate term is a valid biomedical term or not. It is appropriated for *multi-word* 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 to other web-based measures [11], *WebR* reduces the number of pages to consider by taking only web pages containing all words of the terms into account. In addition, our measure can be easily adopted for all types of multi-word terms.

Algorithm 3: ComputeTeRGraph (L_{terms} , num_{terms} , δ , k)

Data: L_{terms} = List of ranked terms;
 num_{terms} = number of terms to be evaluated;
 δ = threshold to create an edge between two terms;
 k = constant;
Result: RRL_{terms} = Re-Ranked List of terms
begin
 Select all possible pairs of terms of L_{terms} to compute $D(x, y)$ // in total
 $C_{num_{terms}}^2 = \frac{num_{terms}!}{2!(num_{terms}-2)!}$ possibilities ;
 Select pairs which $D(x, y) \geq \delta$ for creating an edge ;
 Select all terms of L_{terms} to compute $TeRGraph$;
 for each term $A \in L_{terms}$ **do**
 $N(A) \leftarrow$ neighborhood of A;
 $|N(A)| \leftarrow$ number of neighbors of A;

 $TeRGraph(A) = \log_2 \left(k + \frac{1}{1+|N(A)| + \sum_{T_i \in N(A)} |N(T_i)|} \right)$;

 add A to RRL_{terms} ;
 end
 Rank RRL_{terms} by the value obtained with $TeRGraph$;
end

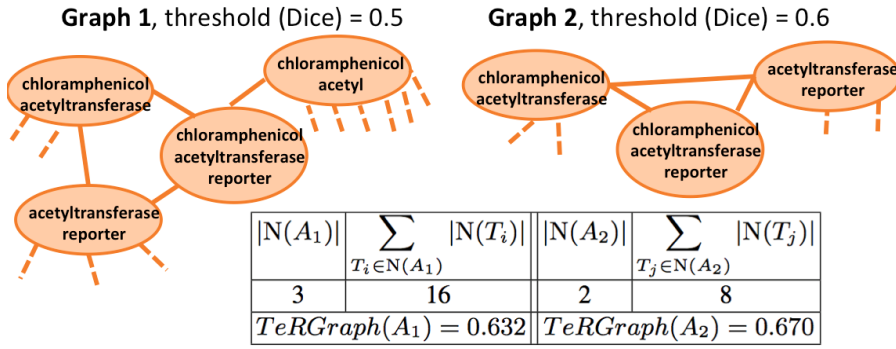


Fig. 5 $TeRGraph$'s value for *chloramphenicol acetyltransferase reporter*

$$WebR(A) = \frac{nb("A")}{nb(A)} \quad (10)$$

Where A = multi-word term, $a_i \in A$ and $a_i = \{noun, adjective, foreign word\}$.

Where A is the candidate term, $nb("A")$ the number of hits returned by a web search engine with exact match only with multi-word term A (query with quotation marks " A "), $nb(A)$ the number of documents returned by the search engine, including not exact matches (query A without quotation marks), i.e. whole documents containing words of the multi-word term A . For example, the multi-word term *treponema pallidum*, will generate two queries, the first

$nb(\text{"treponema pallidum"})$ which returns with Yahoo 1 100 000 documents, and the second query $nb(\text{treponema pallidum})$ which returns 1 300 000 documents, then $WebR(\text{treponema pallidum}) = \frac{1100000}{1300000} = 0.85$.

In our workflow, we tested Yahoo and Bing. *WebR* re-rank the list of candidate terms returned by the combined measures.

3.3.3 A new web ranking measure: WAHI (**W**eb **A**ssociation based on **H**its **I**nformation)

Previous studies of web mining approaches query the web via search engines to measure word associations. This enables measurement of the association of words composing a term (e.g. *soft*, *contact*, and *lens* that compose the relevant term *soft contact lens*). To measure this association, our web-mining approach takes the number of pages provided by search engines into account (i.e. number of hits).

Our web-based measure re-ranks the list obtained previously with *TeR-Graph*. We will show that this improves the precision of the k first terms extracted (see Section 4) and that it is specially appropriate for multi-word term extraction.

Formula 8 leads directly to formula 11¹³. The nb function used in formula 11 represents the number of pages returned by search engines (i.e. Yahoo and Bing). With this measure, we compute a *strict* dependence (i.e. neighboring words by using the operator ' ' ' of search engines). For instance, x might represent the word *soft* and y the word *contact* in order to calculate the association measure of the *soft contact* term.

$$Dice(x, y) = \frac{2 \times nb("x y")}{nb(x) + nb(y)} \quad (11)$$

Then we extend this formula to n elements as follows:

$$Dice(a_1, \dots, a_n) = \frac{n \times nb("a_1 \dots a_n")}{nb(a_1) + \dots + nb(a_n)} = \frac{n \times nb("A")}{\sum_{i=1}^n nb(a_i)} \quad (12)$$

This measure enables us to calculate a score for all multi-word terms, such as *soft contact lens*.

To obtain WAHI, we propose to associate Dice criteria with *WebR* (see formula 10). This only takes the number of web pages containing all the words of the terms into account by using operators " " and *AND*.

For example, *soft contact lens*, the numerator corresponds to the number of web pages with the query "*soft contact lens*", and for the denominator, we consider the query *soft AND contact AND lens*.

¹³ by writing $P(x) = \frac{nb(x)}{nb_{total}}$, $P(y) = \frac{nb(y)}{nb_{total}}$, $P(x, y) = \frac{nb(x, y)}{nb_{total}}$

Finally, the global ranking approach combining *Dice* and *WebR* is given by *WAHI* measure (**W**eb **A**ssociation based on **H**its **I**nformation):

$$WAHI(A) = \frac{n \times nb("A")}{\sum_{i=1}^n nb(a_i)} \times \frac{nb("A")}{nb(A)} \quad (13)$$

Algorithm 4 details the global web mining process to rank terms. We show in the next section that open-domain (general) resources, such as the web, can be tapped to support domain-specific term extraction. They can thus be used to compensate for the unavailability of domain-specific resources.

Algorithm 4: ComputeWAHI (L_{terms} , num_{terms} , LC)

Data: L_{terms} = List of ranked terms;
 num_{terms} = number of terms to be evaluated;
 $LC = \{noun, adjective, foreign\}$ // linguistic categories
Result: RRL_{terms} = Re-Ranked List of terms
begin
 Select the first num_{terms} terms of L_{terms} to compute *WAHI*;
 for each term $A \in L_{terms}$ **do**
 for all words a_i of $A \in LC$ **do**
 $n \leftarrow$ number of words in A ;
 $WAHI(A) \leftarrow n \times \frac{num-hits("A")}{\sum_{i=1}^n num-hits(a_i)} \times \frac{num-hits("A")}{num-hits(A)}$;
 end
 add A to RRL_{terms} ;
 end
 Rank RRL_{terms} by the value obtained with *WAHI*;
end

4 Experiments and results

4.1 Data, Protocol, and Validation

4.1.1 Data

We used two corpora for our experiments. The first one is a set of biological laboratory tests, extracted from LabTestsOnline¹⁴. This website provides information in several languages to patients or family caregivers about clinical lab tests. Each test that 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. The LabTestsOnline website was extracted totally

¹⁴ <http://labtestsonline.org/>

for English, French, and Spanish with a crawler created specifically for this purpose. These documents are available online¹⁵. Table 3 shows the details of LabTestsOnline corpus for different languages.

	Number of Clinical Tests	Number of Words
English	235	377 000 words
French	137	174 000 words
Spanish	238	396 000 words

Table 3 Details of LabTestsOnline corpus.

The second corpus is GENIA¹⁶, which is made up of 2 000 titles and abstracts of journal articles that were culled from the Medline database, with more than 400 000 words in English. The GENIA corpus contains linguistic expressions referring to entities of interest in molecular biology, such as proteins, genes and cells. GENIA is an annotated dataset, in which technical term annotation covers the identification of physical biological entities as well as other important terms. This is our *gold standard corpus*. Whereas the Medline indexes a broad range of academic articles covering the general or specific domains of life sciences, GENIA is intended to cover a smaller subject domain: biological reactions concerning transcription factors in human blood cells.

4.1.2 Protocol

As the measures described in step 2 of our workflow (i.e. *Ranking the Candidate Terms*) are not very time-consuming, and as they are easily applicable for large corpora, they were evaluated over the LabTestsOnline corpus for English, French, and Spanish, and over the gold standard corpus, GENIA. In contrast, as the measures described in step 3 (i.e. *Re-ranking*) are highly time-consuming, and they are used at the end of the process, to enhance the performance of the results, we evaluate them only over the GENIA corpus.

4.1.3 Validation

In order to automatically validate and cover medical terms, we use UMLS for English and Spanish, and the French version of MeSH, SNOMED International and the rest of the French content in the UMLS. For instance, if an extracted candidate term is found in the UMLS dictionary, this term will be automatically validated. The results are evaluated in terms of *precision* obtained over the top k extracted terms ($P@k$).

Biomedical terminologies or ontologies (e.g. UMLS, SNOMED, MeSH), contain terms composed of signs. Therefore, we cleaned these terminologies by eliminating all terms containing (; , ? ! : { } []), and we only took terms

¹⁵ www.lirmm.fr/~lossio/labtestsonline.zip

¹⁶ <http://www.nactem.ac.uk/genia/genia-corpus/term-corpus>

without signs. Table 4 shows the distribution in n -gram (i.e. n -gram is a term of n words, with $n \geq 1$) of biomedical resources for three languages, as well as the number of terms that we took after the cleaning task. For instance, the first cell means that 13.73% of terms are composed of one word (1-gram) in UMLS for English.

	1-gram	2-gram	3-gram	4+ gram	Number of Terms
English	13.73 %	27.65 %	14.44 %	44.18 %	3 006 946
French	13.17 %	25.82 %	17.08 %	43.93 %	304 644
Spanish	8.39 %	19.31 %	16.33 %	55.97 %	534 110

Table 4 Details of Available Resources for Validation.

4.2 Multilingual Comparison (LabTestsOnline)

In this section, we show results obtained only with all the ranking measures, i.e. step 2 (ranking) in Figure 1. In addition, we tested the measures for single- plus multi-word terms, or just for multi-word terms in English, French and Spanish. Table 5, 6, 7 show the results in English, French and Spanish, respectively. At the top of each table, the single-word + multi-word term extraction results are presented, while the multi-word term extraction results are presented at the bottom of the table.

These tables show that *LIDF-value* and *L-value* obtain the best results for both extraction cases and for the three languages. The combined measures based on the harmonic mean, and on the *SUM* and *MAX* (i.e. *F-TFIDF-C_M*, *F-TFIDF-C_S*), also give interesting results.

The single-word + multi-word term extraction results are better than just the multi-word term extraction results. The main reason for this is that the extraction of single-word terms is more efficient due to their syntactic structure (linguistic structure), i.e. usually a *noun*. In addition, this syntactic structure has fewer variations. The results are lower as compared to multi-word term extraction, which is more complicated and involves more variations.

We observe that *LIDF-value* and *L-value* obtain very close results. In most cases *LIDF-value* performs better than *L-value*. These two measures show that the probability associated with the linguistic patterns helps to improve the term extraction results. Note that the *idf* influences *LIDF-value*, for this reason *LIDF-value* has better results than *L-value*.

		Single- and Multi- Word Terms															
		P@100	P@200	P@300	P@400	P@500	P@600	P@700	P@800	P@900	P@1000	P@2000	P@5000	P@10000	P@20000		
<i>C-value</i>		0.930	0.935	0.927	0.943	0.938	0.930	0.920	0.916	0.904	0.892	0.802	0.629	0.480	0.318		
<i>TF-IDF_A</i>		0.7	0.715	0.697	0.663	0.636	0.637	0.616	0.603	0.6	0.588	0.515	0.421	0.350	0.322		
<i>TF-IDF_M</i>		0.910	0.920	0.917	0.898	0.868	0.843	0.824	0.811	0.794	0.781	0.688	0.542	0.448	0.358		
<i>TF-IDF_S</i>		0.970	0.955	0.960	0.960	0.960	0.950	0.943	0.936	0.917	0.906	0.822	0.659	0.511	0.370		
<i>Okapi_A</i>		0.570	0.390	0.4	0.378	0.366	0.347	0.341	0.329	0.336	0.335	0.295	0.314	0.310	0.326		
<i>Okapi_M</i>		0.910	0.915	0.917	0.878	0.824	0.793	0.711	0.685	0.677	0.692	0.613	0.513	0.438	0.355		
<i>Okapi_S</i>		0.940	0.945	0.927	0.930	0.926	0.920	0.909	0.9	0.882	0.873	0.810	0.656	0.513	0.363		
<i>F-OCapi_A</i>		0.690	0.645	0.603	0.570	0.546	0.545	0.527	0.519	0.522	0.527	0.509	0.462	0.414	0.333		
<i>F-OCapi_M</i>		0.970	0.955	0.920	0.895	0.9	0.888	0.874	0.859	0.849	0.842	0.757	0.615	0.465	0.340		
<i>F-OCapi_S</i>		0.940	0.945	0.930	0.935	0.922	0.923	0.917	0.896	0.887	0.879	0.807	0.653	0.515	0.345		
<i>F-TFIDF-CA</i>		0.710	0.715	0.7	0.670	0.642	0.638	0.626	0.613	0.596	0.596	0.532	0.421	0.346	0.323		
<i>F-TFIDF-C_M</i>		0.970	0.960	0.917	0.898	0.866	0.845	0.826	0.811	0.8	0.787	0.692	0.545	0.449	0.357		
<i>F-TFIDF-C_S</i>		0.970	0.960	0.960	0.960	0.960	0.948	0.943	0.931	0.914	0.906	0.823	0.659	0.510	0.369		
<i>L-value</i>		0.960	0.975	0.970	0.965	0.960	0.955	0.951	0.943	0.934	0.933	0.849	0.707	0.597	0.431		
<i>LIDF-value</i>		1.000	0.980	0.970	0.965	0.962	0.955	0.950	0.943	0.934	0.925	0.849	0.716	0.597	0.431		
		Multi-Word Terms															
		P@100	P@200	P@300	P@400	P@500	P@600	P@700	P@800	P@900	P@1000	P@2000	P@5000	P@10000	P@20000		
<i>C-value</i>		0.810	0.790	0.757	0.715	0.686	0.668	0.646	0.633	0.623	0.621	0.527	0.395	0.284	0.189		
<i>TF-IDF_A</i>		0.390	0.4	0.393	0.405	0.424	0.415	0.406	0.401	0.393	0.384	0.315	0.265	0.230	0.204		
<i>TF-IDF_M</i>		0.570	0.6	0.603	0.585	0.578	0.555	0.541	0.526	0.528	0.535	0.456	0.336	0.261	0.209		
<i>TF-IDF_S</i>		0.820	0.765	0.760	0.723	0.7	0.692	0.671	0.639	0.628	0.608	0.526	0.387	0.288	0.211		
<i>Okapi_A</i>		0.4	0.410	0.397	0.373	0.344	0.350	0.343	0.330	0.309	0.292	0.269	0.222	0.214	0.204		
<i>Okapi_M</i>		0.550	0.580	0.580	0.565	0.544	0.545	0.531	0.511	0.510	0.485	0.396	0.325	0.264	0.208		
<i>Okapi_S</i>		0.740	0.680	0.663	0.648	0.644	0.627	0.623	0.623	0.606	0.592	0.497	0.394	0.287	0.209		
<i>F-OCapi_A</i>		0.440	0.435	0.443	0.420	0.422	0.423	0.413	0.399	0.396	0.393	0.338	0.3	0.251	0.206		
<i>F-OCapi_M</i>		0.810	0.720	0.627	0.630	0.606	0.573	0.570	0.571	0.562	0.549	0.495	0.372	0.272	0.208		
<i>F-OCapi_S</i>		0.730	0.705	0.670	0.663	0.646	0.633	0.631	0.624	0.626	0.609	0.503	0.397	0.285	0.208		
<i>F-TFIDF-CA</i>		0.460	0.410	0.433	0.413	0.430	0.433	0.416	0.406	0.4	0.386	0.331	0.276	0.233	0.204		
<i>F-TFIDF-C_M</i>		0.820	0.735	0.630	0.608	0.590	0.565	0.561	0.539	0.529	0.535	0.462	0.349	0.262	0.209		
<i>F-TFIDF-C_S</i>		0.820	0.760	0.763	0.723	0.7	0.692	0.673	0.651	0.641	0.616	0.525	0.390	0.288	0.211		
<i>L-value</i>		0.860	0.760	0.760	0.725	0.704	0.692	0.687	0.651	0.654	0.625	0.536	0.428	0.326	0.235		
<i>LIDF-value</i>		0.860	0.795	0.757	0.743	0.718	0.682	0.667	0.653	0.637	0.626	0.537	0.428	0.327	0.235		

Table 5 Biomedical Term Extraction for English

Single- and Multi- Word Terms																
	P@100	P@200	P@300	P@400	P@500	P@600	P@700	P@800	P@900	P@1000	P@2000	P@5000	P@10000	P@20000		
<i>C-value</i>	0.560	0.610	0.607	0.605	0.594	0.595	0.589	0.584	0.567	0.565	0.469	0.302	0.198	0.121		
<i>TF-IDF_A</i>	0.630	0.575	0.550	0.525	0.486	0.430	0.413	0.395	0.394	0.388	0.291	0.199	0.163	0.145		
<i>TF-IDF_M</i>	0.8	0.745	0.703	0.648	0.626	0.603	0.583	0.566	0.540	0.507	0.419	0.260	0.195	0.156		
<i>TF-IDF_S</i>	0.810	0.780	0.723	0.698	0.662	0.650	0.637	0.625	0.613	0.606	0.510	0.334	0.226	0.161		
<i>Okapi_A</i>	0.580	0.415	0.383	0.315	0.270	0.242	0.229	0.205	0.207	0.220	0.190	0.145	0.146	0.150		
<i>Okapi_M</i>	0.8	0.740	0.683	0.645	0.542	0.532	0.527	0.479	0.432	0.399	0.344	0.256	0.198	0.156		
<i>Okapi_S</i>	0.530	0.455	0.523	0.530	0.558	0.547	0.564	0.574	0.564	0.566	0.5	0.338	0.230	0.159		
<i>F-OCapi_A</i>	0.6	0.525	0.457	0.418	0.386	0.345	0.324	0.308	0.296	0.272	0.214	0.158	0.155	0.149		
<i>F-OCapi_M</i>	0.880	0.735	0.703	0.668	0.654	0.618	0.593	0.574	0.559	0.532	0.408	0.274	0.193	0.153		
<i>F-OCapi_S</i>	0.520	0.470	0.527	0.550	0.550	0.553	0.573	0.568	0.578	0.566	0.5	0.342	0.217	0.153		
<i>F-TFIDF-CA</i>	0.640	0.575	0.557	0.528	0.486	0.453	0.417	0.404	0.396	0.388	0.298	0.199	0.160	0.144		
<i>F-TFIDF-C_M</i>	0.880	0.750	0.703	0.650	0.628	0.603	0.584	0.573	0.546	0.522	0.420	0.261	0.196	0.156		
<i>F-TFIDF-C_S</i>	0.820	0.775	0.720	0.693	0.666	0.647	0.639	0.619	0.613	0.603	0.510	0.334	0.224	0.160		
<i>L-value</i>	0.630	0.650	0.643	0.650	0.654	0.640	0.643	0.646	0.634	0.628	0.543	0.409	0.320	0.187		
<i>LIDF-value</i>	0.860	0.780	0.733	0.705	0.680	0.670	0.654	0.651	0.643	0.628	0.550	0.409	0.320	0.187		
Multi-Word Terms																
	P@100	P@200	P@300	P@400	P@500	P@600	P@700	P@800	P@900	P@1000	P@2000	P@5000	P@10000	P@20000		
<i>C-value</i>	0.450	0.470	0.460	0.425	0.398	0.377	0.359	0.353	0.338	0.315	0.233	0.168	0.091	0.062		
<i>TF-IDF_A</i>	0.330	0.280	0.240	0.245	0.250	0.235	0.209	0.205	0.2	0.195	0.133	0.103	0.087	0.074		
<i>TF-IDF_M</i>	0.460	0.430	0.4	0.340	0.328	0.333	0.314	0.310	0.282	0.258	0.184	0.120	0.098	0.076		
<i>TF-IDF_S</i>	0.610	0.495	0.480	0.438	0.410	0.397	0.386	0.358	0.342	0.333	0.240	0.150	0.106	0.076		
<i>Okapi_A</i>	0.320	0.270	0.230	0.203	0.196	0.173	0.174	0.165	0.152	0.145	0.120	0.095	0.082	0.074		
<i>Okapi_M</i>	0.430	0.420	0.383	0.330	0.328	0.312	0.304	0.275	0.248	0.242	0.168	0.120	0.095	0.075		
<i>Okapi_S</i>	0.420	0.435	0.437	0.405	0.392	0.370	0.350	0.346	0.334	0.326	0.248	0.150	0.103	0.075		
<i>F-OCapi_A</i>	0.330	0.290	0.250	0.258	0.252	0.250	0.229	0.213	0.201	0.196	0.137	0.101	0.085	0.074		
<i>F-OCapi_M</i>	0.560	0.410	0.403	0.395	0.356	0.337	0.326	0.309	0.294	0.281	0.2	0.127	0.096	0.074		
<i>F-OCapi_S</i>	0.420	0.445	0.440	0.415	0.390	0.380	0.364	0.344	0.342	0.334	0.251	0.149	0.103	0.074		
<i>F-TFIDF-CA</i>	0.330	0.295	0.257	0.255	0.248	0.250	0.233	0.221	0.216	0.209	0.136	0.102	0.087	0.074		
<i>F-TFIDF-C_M</i>	0.540	0.425	0.403	0.353	0.328	0.342	0.317	0.313	0.293	0.278	0.184	0.123	0.096	0.076		
<i>F-TFIDF-C_S</i>	0.610	0.475	0.483	0.445	0.422	0.393	0.387	0.368	0.350	0.330	0.242	0.151	0.106	0.076		
<i>L-value</i>	0.620	0.620	0.557	0.515	0.480	0.460	0.442	0.425	0.407	0.401	0.314	0.211	0.138	0.083		
<i>LIDF-value</i>	0.660	0.640	0.563	0.515	0.480	0.460	0.443	0.429	0.413	0.396	0.315	0.212	0.138	0.083		

Table 6 Biomedical Term Extraction for French

Single- and Multi- Word Terms																
	P@100	P@200	P@300	P@400	P@500	P@600	P@700	P@800	P@900	P@1000	P@2000	P@5000	P@10000	P@20000		
C-value	0.630	0.650	0.657	0.625	0.618	0.620	0.609	0.598	0.581	0.570	0.463	0.315	0.216	0.140		
TF-IDF _A	0.340	0.3	0.3	0.325	0.328	0.330	0.323	0.299	0.288	0.283	0.235	0.183	0.151	0.131		
TF-IDF _M	0.740	0.690	0.633	0.575	0.538	0.498	0.496	0.493	0.463	0.462	0.371	0.274	0.208	0.155		
TF-IDF _S	0.810	0.735	0.740	0.718	0.706	0.675	0.651	0.633	0.621	0.599	0.491	0.337	0.239	0.165		
Okapi _A	0.210	0.270	0.210	0.203	0.196	0.182	0.177	0.173	0.171	0.169	0.140	0.116	0.115	0.123		
Okapi _M	0.580	0.6	0.540	0.548	0.530	0.493	0.436	0.429	0.413	0.411	0.326	0.248	0.195	0.150		
Okapi _S	0.560	0.570	0.597	0.615	0.6	0.595	0.586	0.583	0.580	0.580	0.5	0.346	0.238	0.161		
F-OCapi _A	0.250	0.275	0.227	0.245	0.248	0.252	0.249	0.234	0.228	0.223	0.158	0.124	0.122	0.131		
F-OCapi _M	0.810	0.695	0.587	0.548	0.528	0.522	0.471	0.449	0.439	0.448	0.414	0.275	0.199	0.149		
F-OCapi _S	0.560	0.570	0.613	0.615	0.602	0.595	0.586	0.586	0.578	0.576	0.5	0.343	0.231	0.158		
F-TFIDF-CA	0.350	0.330	0.303	0.333	0.328	0.330	0.321	0.301	0.288	0.284	0.235	0.186	0.150	0.131		
F-TFIDF-C _M	0.820	0.705	0.640	0.583	0.552	0.497	0.497	0.494	0.473	0.467	0.375	0.274	0.210	0.155		
F-TFIDF-C _S	0.810	0.745	0.740	0.720	0.702	0.675	0.660	0.630	0.612	0.602	0.491	0.338	0.238	0.165		
L-value	0.660	0.660	0.623	0.630	0.610	0.608	0.597	0.590	0.573	0.557	0.467	0.339	0.250	0.176		
LIDF-value	0.810	0.755	0.730	0.710	0.696	0.682	0.677	0.663	0.653	0.645	0.512	0.436	0.324	0.248		
Multi-Word Terms																
	P@100	P@200	P@300	P@400	P@500	P@600	P@700	P@800	P@900	P@1000	P@2000	P@5000	P@10000	P@20000		
C-value	0.420	0.435	0.417	0.378	0.368	0.352	0.340	0.321	0.306	0.294	0.225	0.157	0.106	0.068		
TF-IDF _A	0.150	0.160	0.173	0.140	0.128	0.147	0.151	0.156	0.143	0.140	0.119	0.098	0.081	0.068		
TF-IDF _M	0.350	0.350	0.343	0.290	0.272	0.242	0.217	0.226	0.221	0.216	0.175	0.132	0.101	0.075		
TF-IDF _S	0.570	0.470	0.430	0.405	0.380	0.362	0.340	0.333	0.323	0.304	0.228	0.156	0.110	0.080		
Okapi _A	0.110	0.135	0.120	0.123	0.116	0.125	0.131	0.134	0.123	0.115	0.094	0.080	0.076	0.069		
Okapi _M	0.310	0.280	0.317	0.288	0.246	0.230	0.214	0.208	0.209	0.205	0.158	0.120	0.096	0.077		
Okapi _S	0.420	0.415	0.393	0.393	0.366	0.342	0.323	0.328	0.323	0.305	0.238	0.153	0.107	0.080		
F-OCapi _A	0.110	0.150	0.130	0.128	0.132	0.138	0.134	0.136	0.147	0.144	0.104	0.085	0.079	0.070		
F-OCapi _M	0.460	0.325	0.333	0.295	0.260	0.240	0.223	0.223	0.218	0.220	0.193	0.122	0.098	0.077		
F-OCapi _S	0.410	0.410	0.403	0.393	0.372	0.340	0.331	0.328	0.322	0.315	0.239	0.153	0.108	0.079		
F-TFIDF-CA	0.160	0.170	0.177	0.148	0.134	0.148	0.157	0.159	0.157	0.152	0.128	0.099	0.082	0.068		
F-TFIDF-C _M	0.480	0.375	0.347	0.298	0.280	0.245	0.221	0.228	0.223	0.223	0.188	0.133	0.1	0.075		
F-TFIDF-C _S	0.570	0.455	0.430	0.408	0.392	0.367	0.344	0.335	0.327	0.314	0.230	0.157	0.111	0.080		
L-value	0.470	0.490	0.460	0.438	0.410	0.387	0.370	0.359	0.349	0.337	0.266	0.189	0.144	0.090		
LIDF-value	0.530	0.510	0.460	0.438	0.418	0.392	0.370	0.368	0.349	0.337	0.274	0.189	0.144	0.090		

Table 7 Biomedical Term Extraction for Spanish

4.3 Evaluation of the global process (GENIA)

Since GENIA is the gold standard corpus, we conduct a detailed assessment of the experiments in this subsection. We evaluated the entire workflow of our methodology, i.e. steps 2 (ranking) and 3 (re-ranking) in Figure 1. As noted earlier, the multi-word term extraction results are influenced by the syntactic structure and their variations. So our experimentation in this subsection is focused only on multi-word term extraction.

In the following paragraphs, we also narrow down the presented results by keeping only the first 8 000 extracted terms for the graph-based measure and the first 1000 extracted terms for the web-based measure.

4.3.1 Ranking Results (step 2 in Figure 1)

Table 8 presents and compares the multi-word term extraction results with the best ranking measures, as shown earlier, i.e. *C-value*, *F-TFIDF-C_M*, and *LIDF-value*. The best results were obtained with *LIDF-value* with an 11% improvement in precision for the first hundred extracted multi-word terms. These precision results are also shown in Figure 6. The precision of *LIDF-value* will be further improved with *TeRGraph*.

	<i>C-value</i>	<i>F-TFIDF-C_M</i>	<i>LIDF-value</i>
P@100	0.690	0.715	0.820
P@200	0.690	0.715	0.770
P@300	0.697	0.710	0.750
P@400	0.665	0.690	0.738
P@500	0.642	0.678	0.718
P@600	0.638	0.668	0.723
P@700	0.627	0.669	0.717
P@800	0.611	0.650	0.710
P@900	0.612	0.629	0.714
P@1000	0.605	0.618	0.697
P@2000	0.570	0.557	0.662
P@5000	0.498	0.482	0.575
P@10000	0.428	0.412	0.526
P@20000	0.353	0.314	0.377

Table 8 Precision comparison of *LIDF-value* with baseline measures

Results of *n*-gram Terms

We also evaluated *C-value*, *F-TFIDF-C_M*, and *LIDF-value* in a sequence of *n*-gram terms (i.e. *n*-gram term is a multi-word term of *n* words), for this we require an index term to be a *n*-gram terms of length $n \geq 2$. We tested the performance of *LIDF-value* on the *n*-gram term extraction taking the first 1 000 *n*-gram terms ($n \geq 2$).

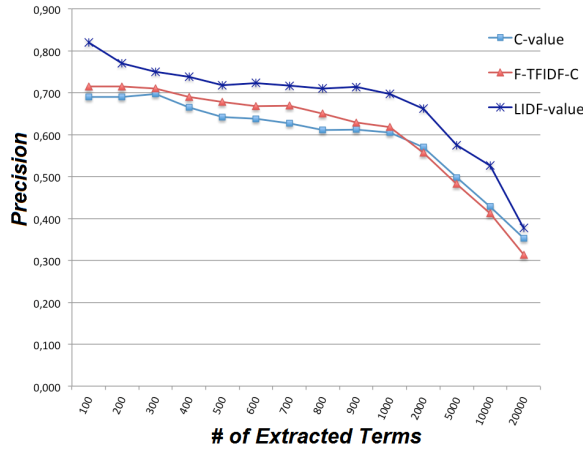


Fig. 6 Precision comparison with *LIDF-value* and baseline measures

Table 9 shows the precision comparison for the 2-gram, 3-gram and 4+ gram term extracted with *C-value*, *F-TFIDF-C_M*, and *LIDF-value*. We can see that *LIDF-value* obtains the best results for all intervals for any $n \geq 2$. These precision results are also shown in Figure 7 for the 2-gram terms, Figure 8 for the 3-gram terms, and finally Figure 9 for the 4+ gram terms.

Table 10 shows the top-20 ranked 2-gram terms extracted with the baseline measures and *LIDF-value*. *C-value* obtained 3 irrelevant terms, *F-TFIDF-C* obtained 5 irrelevant terms while *LIDF-value* obtained only 2 irrelevant terms for the top-20 ranked 2-gram terms.

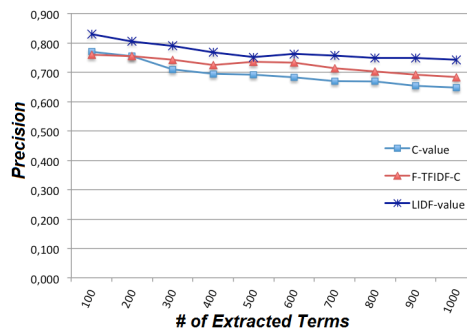
Similarly, Table 11 shows top-10 ranked 3-gram terms extracted with the baseline measures and *LIDF-value*. Finally, Table 12 shows the top-10 ranked 4+ gram terms extracted with the baseline measures and *LIDF-value*.

Note that in this context, “irrelevant” means that the terms are not in the above mentioned resources. These candidate terms might be interesting for ontology extension or population, however they must pass through polysemy detection in order to identify the possible meanings.

4.3.2 Re-ranking Results (step 3 in Figure 1)

Graph-based Results: our graph-based approach is applied to the first 8 000 terms extracted by the best ranking measure. The objective is to re-rank the 8 000 terms while trying to improve the precision by intervals. One parameter is involved in the computation of graph-based term weights, i.e. the *threshold* of Dice value which represents the relation when building the term graph. This involves linking terms whose *Dice value* of the relation is higher than *threshold*. We vary *threshold* (δ) within $\delta = [0.25, 0.35, 0.50, 0.60, 0.70]$ and report the precision performance for each of these values. Table 13 gives the precision performance obtained by *TeRGraph* and shows that it is well adapted for ATE.

<i>2-gram terms</i>			
	<i>C-value</i>	<i>F-TFIDF-C</i>	<i>LIDF-value</i>
P@100	0.770	0.760	0.830
P@200	0.755	0.755	0.805
P@300	0.710	0.743	0.790
P@400	0.695	0.725	0.768
P@500	0.692	0.736	0.752
P@600	0.683	0.733	0.763
P@700	0.670	0.714	0.757
P@800	0.669	0.703	0.749
P@900	0.654	0.692	0.749
P@1000	0.648	0.684	0.743
<i>3-gram terms</i>			
	<i>C-value</i>	<i>F-TFIDF-C</i>	<i>LIDF-value</i>
P@100	0.670	0.530	0.820
P@200	0.590	0.450	0.795
P@300	0.577	0.430	0.777
P@400	0.560	0.425	0.755
P@500	0.548	0.398	0.744
P@600	0.520	0.378	0.720
P@700	0.499	0.370	0.706
P@800	0.488	0.379	0.691
P@900	0.482	0.399	0.667
P@1000	0.475	0.401	0.660
<i>4+ gram terms</i>			
	<i>C-value</i>	<i>F-TFIDF-C</i>	<i>LIDF-value</i>
P@100	0.510	0.370	0.640
P@200	0.455	0.330	0.520
P@300	0.387	0.273	0.477
P@400	0.393	0.270	0.463
P@500	0.378	0.266	0.418
P@600	0.348	0.253	0.419
P@700	0.346	0.249	0.390
P@800	0.323	0.248	0.395
P@900	0.323	0.240	0.364
P@1000	0.312	0.232	0.354

Table 9 Precision comparison of 2-gram terms, 3-gram terms, and 4+ gram terms**Fig. 7** Precision comparison of 2-gram terms

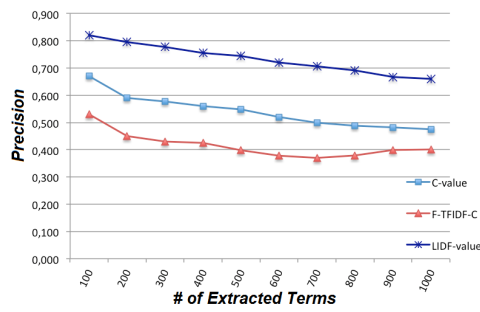


Fig. 8 Precision comparison of 3-gram terms

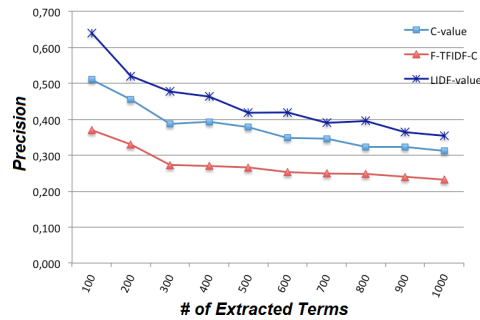


Fig. 9 Precision comparison of 4+ gram terms

	<i>C-value</i>	<i>F-TFIDF-C</i>	<i>LIDF-value</i>
1	t cell	t cell	t cell
2	nf-kappa b	nf-kappa b	transcription factor
3	transcription factor	kappa b	nf-kappa b
4	gene expression	b cell	cell line
5	kappa b	class ii	b cell
6	cell line	glucocorticoid receptor	gene expression
7	b cell	<i>b activation *</i>	kappa b
8	peripheral blood	<i>b alpha *</i>	t lymphocyte
9	t lymphocyte	reporter gene	dna binding
10	nuclear factor	endothelial cell	<i>i kappa *</i>
11	protein kinase	cell cycle	binding site
12	class ii	b lymphocyte	protein kinase
13	<i>b activation *</i>	<i>nf kappa *</i>	glucocorticoid receptor
14	human t	nf-kappab activation	tumor necrosis
15	tyrosine phosphorylation	u937 cell	binding activity
16	dna binding	<i>mhc class *</i>	tyrosine phosphorylation
17	<i>human immunodeficiency *</i>	<i>c ebp*</i>	<i>shift assay *</i>
18	binding site	il-2 promoter	immunodeficiency virus
19	<i>necrosis factor *</i>	monocytic cell	signal transduction
20	mobility shift	t-cell leukemia	mobility shift

Table 10 Comparison of top-20 ranked 2-gram terms (irrelevant terms are italicized and marked with *).

	<i>C-value</i>	<i>F-TFIDF-C</i>	<i>LIDF-value</i>
1	human immunodeficiency virus	<i>kappa b alpha *</i>	i kappa b
2	<i>kappa b alpha *</i>	nf kappa b	human immunodeficiency virus
3	tumor necrosis factor	jurkat t cell	electrophoretic mobility shift
4	electrophoretic mobility shift	human t cell	human t cell
5	nf-kappa b activation	mhc class ii	mobility shift assay
6	<i>virus type 1 *</i>	cd4+ t cell	<i>kappa b alpha *</i>
7	protein kinase c	<i>c-fos and c-jun *</i>	tumor necrosis factor
8	long terminal repeat	peripheral blood monocyte	nf-kappa b activation
9	nf kappa b	t cell proliferation	protein kinase c
10	jurkat t cell	<i>transcription factor nf-kappa *</i>	jurkat t cell

Table 11 Comparison of the top-10 ranked 3-gram terms (irrelevant terms are italicized and marked with *).

	<i>C-value</i>	<i>F-TFIDF-C</i>	<i>LIDF-value</i>
1	human immunodeficiency virus type 1	transcription factor nf-kappa b	i kappa b alpha
2	<i>human immunodeficiency virus type *</i>	<i>expression of nf-kappa b *</i>	electrophoretic mobility shift assay
3	<i>immunodeficiency virus type 1 *</i>	tumor necrosis factor alpha	<i>human immunodeficiency virus type *</i>
4	activation of nf-kappa b	normal human t cell	human t-cell leukemia virus
5	nuclear factor kappa b	primary human t cell	nuclear factor kappa b
6	tumor necrosis factor alpha	germline c epsilon transcription	tumor necrosis factor alpha
7	<i>human t-cell leukemia viru *</i>	gm-csf receptor alpha promoter	<i>t-cell leukemia virus type *</i>
8	<i>human t-cell leukemia virus type *</i>	il-2 receptor alpha chain	activation of nf-kappa b
9	<i>t-cell leukemia virus type *</i>	<i>transcription from the gm-csf *</i>	peripheral blood t cell
10	electrophoretic mobility shift assay	<i>translocation of nf-kappa b *</i>	major histocompatibility complex class

Table 12 Comparison of the top-10 ranked 4+ gram terms (irrelevant terms are italicized and marked with *).

	<i>TeRGraph</i>				
	$\delta \geq 0.25$	$\delta \geq 0.35$	$\delta \geq 0.50$	$\delta \geq 0.60$	$\delta \geq 0.70$
P@100	0.840	0.860	0.910	0.930	0.900
P@200	0.800	0.790	0.850	0.855	0.855
P@300	0.803	0.773	0.833	0.830	0.820
P@400	0.780	0.732	0.820	0.820	0.815
P@500	0.774	0.712	0.798	0.810	0.806
P@600	0.773	0.675	0.797	0.807	0.792
P@700	0.760	0.647	0.769	0.796	0.787
P@800	0.756	0.619	0.748	0.784	0.779
P@900	0.748	0.584	0.724	0.773	0.777
P@1000	0.751	0.578	0.720	0.766	0.769
P@2000	0.689	0.476	0.601	0.657	0.694
P@3000	0.642	0.522	0.535	0.605	0.644
P@4000	0.612	0.540	0.543	0.559	0.593
P@5000	0.574	0.546	0.544	0.554	0.562
P@6000	0.558	0.539	0.540	0.549	0.561
P@7000	0.556	0.540	0.540	0.545	0.552
P@8000	0.546	0.546	0.546	0.546	0.546

Table 13 Precision performance of *TeRGraph* when varying δ (*threshold* parameter for Dice)

Web-based Results: Our web-based approach is applied at the end of the process, with only the first 1 000 terms extracted during the previous linguistic, statistic and graph measures. For space reasons, we show only the results obtained with *WAHI*, which are higher than *WebR*.

We took the list obtained with *TeRGraph* and $\delta \geq 0.60$. The main reason for this limitation is the limited number of automatic queries possible in search engines. At this step, the aim is to re-rank the 1 000 terms to try to improve the precision by intervals. Each measure listed in Table 14 and Table 15 shows the precision obtained after re-ranking. We tested *WAHI* with *Yahoo* and *Bing* search engines.

Table 14 and Table 15 prove that *WAHI* (either using *Yahoo* or *Bing*) is well adapted for ATE and this measure obtains better precision results than the baselines measures for word association. So our measures obtain real terms of our dictionary with a better ranking.

	<i>WAHI</i>	<i>Dice</i>	<i>Jaccard</i>	<i>Cosine</i>	<i>Overlap</i>
P@100	0.960	0.720	0.720	0.760	0.730
P@200	0.950	0.785	0.770	0.740	0.765
P@300	0.900	0.783	0.780	0.767	0.753
P@400	0.900	0.770	0.765	0.770	0.740
P@500	0.920	0.764	0.754	0.762	0.738
P@600	0.850	0.748	0.740	0.765	0.748
P@700	0.817	0.747	0.744	0.747	0.757
P@800	0.875	0.752	0.746	0.740	0.760
P@900	0.870	0.749	0.747	0.749	0.747
P@1000	0.766	0.766	0.766	0.766	0.766

Table 14 Precision comparison of *WAHI* with *YAHOO* and word association measures

	<i>WAHI</i>	<i>Dice</i>	<i>Jaccard</i>	<i>Cosine</i>	<i>Overlap</i>
P@100	0.900	0.740	0.730	0.680	0.650
P@200	0.900	0.775	0.775	0.735	0.705
P@300	0.900	0.770	0.763	0.740	0.713
P@400	0.900	0.765	0.765	0.752	0.712
P@500	0.900	0.760	0.762	0.758	0.726
P@600	0.917	0.753	0.752	0.753	0.743
P@700	0.914	0.751	0.751	0.733	0.749
P@800	0.875	0.745	0.747	0.741	0.754
P@900	0.878	0.747	0.748	0.742	0.748
P@1000	0.766	0.766	0.766	0.766	0.766

Table 15 Precision comparison of *WAHI* with *BING* and word association measures

4.3.3 Summary

LIDF-value obtains the best precision results for multi-word term extraction, for each index term extraction (n -gram) and for intervals.

Table 16 presents a precision comparison of *LIDF-value* and *TeRGraph* measures. In terms of overall precision, our experiments produce consistent

results from the GENIA corpus. In most cases, *TeRGraph* obtains better precision with a δ of 0.60 and 0.70 (i.e. better precision in most $P@k$ intervals), which is very good because it helps alleviate the problem of manual validation of candidate terms. These precisions are also illustrated in Figure 10.

The performance of our graph-based measure somewhat depends on the value of the co-occurrence relation between terms. Specifically, the value of the co-occurrence relation affects how the graph is built (whose edges are taken), and hence it is critical for computation of the graph-based term weight. Another performance factor of our graph-based measure is the quality of the results obtained with *LIDF-value* due to the fact that the list of terms extracted with *LIDF-value* is required as input to re-rank *TeRGraph* in order to construct the graph, where nodes denote terms, and edges denote co-occurrence relations.

	<i>LIDF-value</i>	<i>TeRGraph</i> ($\delta \geq 0.60$)	<i>TeRGraph</i> ($\delta \geq 0.70$)
P@100	0.820	0.930	0.900
P@200	0.770	0.855	0.855
P@300	0.750	0.830	0.820
P@400	0.738	0.820	0.815
P@500	0.718	0.810	0.806
P@600	0.723	0.807	0.792
P@700	0.717	0.796	0.787
P@800	0.710	0.784	0.779
P@900	0.714	0.773	0.777
P@1000	0.697	0.766	0.769
P@2000	0.662	0.657	0.694
P@3000	0.627	0.605	0.644
P@4000	0.608	0.5585	0.593
P@5000	0.575	0.5538	0.562
P@6000	0.550	0.549	0.561
P@7000	0.547	0.545	0.552
P@8000	0.546	0.546	0.546

Table 16 Precision comparison of *LIDF-value* and *TeRGraph*

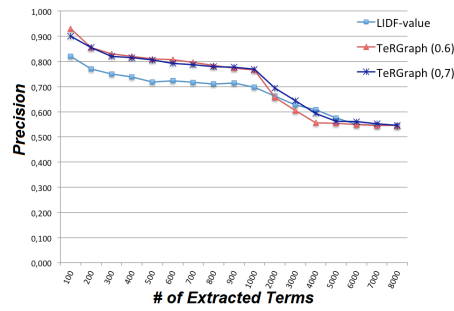


Fig. 10 Precision comparison of *LIDF-value* and *TeRGraph*

Table 17 presents the precision comparison of our three measures.

WAHI based on Yahoo obtains better precision for the first $P@100$ extracted terms with 96% precision whereas, in comparison, *WAHI* based on Bing obtains 90 precision. For the other interval, Table 17 shows that *WAHI* based on Bing generally gives the best results. This is very encouraging because it also helps alleviate the problem of manual validation of candidate terms.

The performance of *WAHI* depends on the search engine because algorithms designed for searching information on the web are different, so the number of hits returned will differ in all cases. Another performance factor is the quality of the re-ranked list obtained with *TeRGraph*, because this list is required as input.

Moreover, Table 17 highlights that re-ranking with *WAHI* enables us to increase the precision of *TeRGraph*. For all cases, our re-ranking methods improve the precision obtained with *LIDF-value*. The purpose for which this web-mining measure was designed has thus been fulfilled.

Note that these measures do not normalize the possible variants. This could be a limitation for researchers looking for a preferred term for a group of variants.

	<i>LIDF-value</i>	<i>TeRGraph</i> ($\delta \geq 0.60$)	<i>WAHI</i> (<i>Bing</i>)	<i>WAHI</i> (<i>Yahoo</i>)
P@100	0.820	0.930	0.900	0.960
P@200	0.770	0.855	0.900	0.950
P@300	0.750	0.830	0.900	0.900
P@400	0.738	0.820	0.900	0.900
P@500	0.718	0.810	0.900	0.920
P@600	0.723	0.807	0.917	0.850
P@700	0.717	0.796	0.914	0.817
P@800	0.710	0.784	0.875	0.875
P@900	0.714	0.773	0.878	0.870
P@1000	0.697	0.766	0.766	0.766

Table 17 Precision comparison *LIDF-value*, *TeRGraph*, and *WAHI*

5 Discussion

We discuss the effects of some parameters of our workflow. In the next sections, we explain the impacts of biomedical pattern lists, size of dictionaries, and the extraction errors.

5.1 Impact of Pattern List

In our methodology, we have shown that biomedical patterns directly affect the term extraction results. For instance, we can see that *L-value*, which is

a combination of *C-value* and the probability of pattern lists, gives better results than *C-value* for the three languages, and *LIDF-value* outperforms *L-value* in major cases. These pattern lists work specifically for the biomedical domain. If we use these biomedical patterns in another domain instead of using specific patterns of that domain, they will impact the term extraction results. To prove this, we have extracted terms from an agronomic corpus for English and French while taking biomedical patterns and agronomic patterns into account. We built the agronomic patterns using AGROVOC¹⁷, which is an agronomic dictionary. AGROVOC contains 39 542 and 37 382 English and French terms, respectively. Our corpus consists of titles plus abstracts extracted from the list of Cirad publications (French Agricultural Research Centre for International Development). Table 18 shows the details of the corpus formed for this comparison.

Table 19 presents a term extraction comparison while taking patterns built from two different domains into account. Again we note that *LIDF-value* obtains the best results. We also see that the results of terms extracted with agronomic patterns gives better results than when using biomedical patterns for English and French.

Note that even if the term extraction results obtained using agronomic patterns are higher than using biomedical patterns, these results are a bit close. The main reason is that the biomedical and agronomic terms overlap. It means that identical patterns exist in both domains. The results could be improved by using patterns of two completely different domains.

	Number of Titles/Abstracts	Number of Words
English	156	29 740 words
French	84	14 850 words

Table 18 Details of Cirad corpus.

5.2 Effect of Dictionary Size

Dictionaries play an important role in term extraction, specifically during the construction of pattern lists. Table 19 shows that a reduction in dictionary size degrades the performance of the precision results in comparison to Tables 5, 6, 8. For instance, for the agronomic and biomedical domain, Table 19 and Table 5 show the P@100 of 0.92 and 1.00 respectively, and this difference increases as the number of extracted terms increases (i.e. P@*k*).

¹⁷ <http://aims.fao.org/agrovoc>

	English (Single- and Multi- Word Terms)							
	With Agronomic Patterns				With Biomedical Patterns			
	P@100	P@200	P@1000	P@5000	P@100	P@200	P@1000	P@5000
<i>C-value</i>	0.910	0.825	0.631	0.255	0.870	0.790	0.527	0.223
<i>TF-IDF_S</i>	0.900	0.830	0.667	0.335	0.810	0.845	0.587	0.284
<i>Okapi_S</i>	0.910	0.865	0.680	0.331	0.870	0.845	0.625	0.281
<i>F-OCapi_M</i>	0.640	0.600	0.419	0.273	0.660	0.605	0.403	0.252
<i>F-OCapi_S</i>	0.900	0.845	0.672	0.304	0.870	0.840	0.612	0.260
<i>F-TFIDF-C_M</i>	0.740	0.610	0.412	0.261	0.760	0.610	0.402	0.270
<i>F-TFIDF-C_S</i>	0.900	0.835	0.664	0.323	0.810	0.845	0.600	0.272
<i>L-value</i>	0.700	0.660	0.542	0.338	0.840	0.795	0.688	0.320
<i>LIDF-value</i>	0.920	0.875	0.766	0.340	0.880	0.855	0.682	0.320

	French (Single- and Multi- Word Terms)							
	With Agronomic Patterns				With Biomedical Patterns			
	P@100	P@200	P@1000	P@5000	P@100	P@200	P@1000	P@5000
<i>C-value</i>	0.400	0.360	0.210	0.086	0.450	0.455	0.223	0.084
<i>TF-IDF_S</i>	0.430	0.380	0.248	0.114	0.500	0.450	0.293	0.119
<i>Okapi_S</i>	0.390	0.360	0.256	0.115	0.490	0.450	0.300	0.120
<i>F-OCapi_M</i>	0.310	0.225	0.154	0.100	0.340	0.245	0.167	0.115
<i>F-OCapi_S</i>	0.400	0.355	0.248	0.106	0.480	0.465	0.269	0.115
<i>F-TFIDF-C_M</i>	0.350	0.240	0.163	0.099	0.380	0.295	0.170	0.118
<i>F-TFIDF-C_S</i>	0.350	0.240	0.163	0.099	0.500	0.475	0.268	0.119
<i>L-value</i>	0.550	0.510	0.367	0.135	0.520	0.480	0.333	0.130
<i>LIDF-value</i>	0.560	0.535	0.367	0.135	0.510	0.510	0.336	0.130

Table 19 Precision comparison of Term Extraction with Agronomic and Biomedical Patterns

5.3 Term Extraction Errors

As explained in Section 3 (step a), the term extraction results are influenced by the Part-of-Speech (PoS) tagging tools, which have different results for different languages. Briefly, the tool “A” can perform very well for English, while for French the tool “B” gives the best results. For instance, the sentence “Red blood cells increase with ...” was tagged with the Stanford tool as “*adjective noun noun verb preposition ...*”, whereas the TreeTagger tool tagged it as “*adjective noun noun noun preposition ...*”. Therefore, in order to show the generality of our approach, we choose a uniform PoS tool, i.e. TreeTagger, as a trade-off for three languages (English, French, and Spanish), while understanding that it will penalize the results for the three languages.

6 Conclusions and Future Work

This paper defines and evaluates several measures for automatic multi-word term extraction. These measures are classified as *ranking measures*, and *re-ranking measures*. The measures are based on the linguistic, statistical, graphic and web information. We modified some baseline measures (i.e. *C-value*, *TF-IDF*, *Okapi*) and we proposed new measures.

All the ranking measures are linguistic- and statistic-based. The best ranking measure is *LIDF-value*, which overcomes the lack of frequency information with the *linguistic pattern probability* and *idf* values.

We experimentally showed that *LIDF-value* applied in the biomedical domain, over two corpora (i.e. LabTestsOnline, GENIA), outperformed a state-of-the-art baseline for extracting terms (i.e. *C-value*), while obtaining the best precision results in all intervals (i.e. $P@k$). And with three languages the *LIDF-value* trends were similar.

We have shown that multi-word term extraction is more complex than single-word term extraction. We detailed an evaluation over the GENIA corpus for multi-word term extraction. Moreover, in that case, *LIDF-value* improved the automatic term extraction precision in comparison to the most popular term extraction measure.

We also evaluated the re-ranking measures. The first re-ranking measure, *TeRGraph*, is a graph-based measure. It decreases the human effort required to validate candidate terms. The graph-based measure has never been applied for automatic term extraction. *TeRGraph* takes the neighborhood to compute the term representativeness in a specific domain into account.

The other re-ranking measures are web-based. The best one, called *WAHI*, takes the list of terms obtained with *TeRGraph* as input. *WAHI* enables us to further reduce the huge human effort required for validating candidate terms.

Our experimental evaluations revealed that *TeRGraph* had better precision than *LIDF-value* for all intervals. Moreover, our experimental assessments revealed that *WAHI* improved the results given with *TeRGraph* for all intervals.

As a future extension of this work, we intend to use the relation value within *TeRGraph*. We plan to include the use of other graph ranking computations, e.g. PageRank, adapted for automatic term extraction. Moreover, a future work consists of using the web to extract more terms than those extracted.

One prospect could be the creation of a regular expression for the biomedical domain from the linguistic pattern list. We plan to modify our measures in order to normalize the possible variants, looking towards for a preferred term for those variants.

Acknowledgements This work was supported in part by the French National Research Agency under the JCJC program, grant ANR-12-JS02-01001, as well as by the University of Montpellier, CNRS, IBC of Montpellier project and the FINCyT program, Peru. Finally, the author thanks DIST (Scientific and Technical Information Service) for the acquisition of Cirad corpus.

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