Abstract

The paper introduces a framework for representation and acquisition of knowledge emerging from large samples of textual data. We utilise a tensor-based, distributional representation of simple statements extracted from text, and show how one can use the representation to infer emergent knowledge patterns from the textual data in an unsupervised manner. Examples of the patterns we investigate in the paper are implicit term relationships or conjunctive IF-THEN rules. To evaluate the practical relevance of our approach, we apply it to annotation of life science articles with terms from MeSH (a controlled biomedical vocabulary and thesaurus).

1 Introduction

The ubiquity of methods for digital content publishing, processing and sharing has led to a lot of data being made globally available every day. Such an unprecedented worldwide availability of content is generally beneficial, yet it also poses big challenges. For instance, in as dynamic and voluminous domains as life sciences, it is virtually impossible for the users to utilise all the available relevant knowledge in a comprehensive and timely manner [17].

Mitigation of this problem (with a special focus on biomedical literature) has served as the main motivation for the research presented in this paper. As can be seen for instance in [9], a popular way of tackling the information overload in the context of biomedical literature is annotation of articles by terms from standardised biomedical vocabularies. Such annotations can in turn make the retrieval of relevant documents much more efficient. However, as providing the necessary annotations manually is very expensive, automated methods are desired [9], which is what we are going to address here.

The technical contribution of the presented work is two-fold. Firstly, we introduce a general framework for automated acquisition of knowledge from textual collections. The proposed framework builds on the principles of distributional [7] and emergent [5].
semantics, and allows for inference of complex knowledge patterns within simple co-
ocurrence statements extracted from articles. As a second contribution, we show how
the knowledge inferred from the text can be applied to unsupervised and parameter-free
annotation of biomedical articles.

The rest of the paper is organised as follows. Section 2 gives an overview of re-
lated work. The framework for emergent knowledge representation and acquisition is
described in Section 3. The application of the framework to document annotation is
detailed in Section 4 where we also discuss an experiment we performed to evaluate
our approach. Section 5 concludes the paper and outlines our future work.

2 Related Work

Our approach builds on and shares a lot of similarities with recent works in emer-
gent [5] and distributional [3] semantics. However, [5] is quite restrictive and applies
the notion of emergence [4] merely to complex patterns arising from simple inter-
actions of autonomous agents in distributed systems like P2P networks. We are more
general, focusing rather on inference and analysis of complex patterns emerging within
large amounts of simple statements being extracted directly from data. This is in accord-
dance with a recent approach to distributional semantics presented in [3]. We employ
similar tensor-based structures for representation of data and analysis of the knowledge
emerging from them. Yet we also augment the work [3] by an explicit representation
data provenance and, more importantly, by a method for mining rules out of the
distributional representations. The latter is related to the associative rule mining intro-
duced in [1], however, we generalise the state of the art method to make use of our
distributional (essentially vector-based) representation of the data.

Regarding the application of our framework to annotation of biomedical articles,
a body of more or less recent works like [10], [2], [13], [15] or [9] exists (the sec-
ond, third and fifth of the approaches are either used or considered for use as a support
service for the professional annotators of the articles on PubMed, a biomedical liter-
ature repository). The state of the art methods, however, often require at least an
indirect input from human users before they can produce annotations of new articles
automatically. For instance, [2] and [9] require a large corpus of previously annotated
articles for learning and ranking possible annotations of new resources. Other methods
like [13] require rather sophisticated tuning (e.g., experimenting with parameter set-
ings or with the processing pipeline composition) for optimum performance on new
data. This is not the case of our approach, as it can work in a purely unsupervised
manner off-the-shelf.

3 Distributional Framework for Emergent Knowledge
Acquisition

This section first describes how one can represent the knowledge emerging from textual
documents at various levels of complexity: (1) simple term co-occurrence statements
within the documents; (2) an integral view on the statements across the document cor-
pus; (3) different perspectives of the corpus-wide view for analysing various types of
emergent semantic phenomena. All levels of the representation are based on compact
tensor structures (tensor is a generalisation of the scalar, vector and matrix notions;
see http://en.wikipedia.org/wiki/Tensor for a more detailed overview).
The rest of the section deals with analysis of two particular types of emergent semantic phenomena that are relevant to document annotation, our motivating use case.

3.1 Source Representation

The first layer consists of a so called source representation $G$, which captures the co-occurrence of terms across a set of documents (a concrete way of extracting co-occurrence relationships is presented in Section 4). Let $A_l, A_r$ be sets representing left and right arguments of binary co-occurrence relationships (i.e., statements), and $L$ the types of the relationships. Furthermore, let $P$ be a set representing provenances of particular relationships (i.e., document identifiers). We define the source representation as a 4-ary labeled tensor $G \in \mathbb{R}^{|A_l| \times |L| \times |A_r| \times |P|}$. It is a four-dimensional array structure indexed by argument-link-argument-provenance tuples, with values reflecting the weight (e.g., frequency) of statements in the context of particular sources ($0$ if a statement does not occur in a source). For instance, if a statement (protein, different from, gene) occurs two times in a source $d_x$, then the element $g_{protein,different from,gene,d_x}$ of $G$ will be $2$. More details are given in Example 1.

Example 1 Let us consider documents $d_1, d_2, d_3, d_4$ and the following terms occurring in them: protein domain, protein, domain, gene, internal tandem duplications, mutations, juxtamembrane, extracellular domains (abbreviated as p.d., p., d., g., i.t.d., m., j., e.d., respectively, in the following). Let us further assume that the following statements were extracted from the documents:

- $d_1$: \{(p.d., D, p.), (p.d., T, d.)\}
- $d_2$: \{(g., D, p.)\}
- $d_3$: \{(i.t.d., T, m.), (i.t.d., I, j.), (i.t.d., I, e.d.)\}
- $d_4$: \{(p.d., D, p.)\}, where D, T, I are abbreviations for relation terms different from, type of, in. When omitting all zero values and representing a four-dimensional tensor as a two-dimensional table where the three first columns are for the tensor indices and the fourth one is for the corresponding tensor value, we can represent the source with the above statements as follows (using statement frequencies as values):

|   | 1 | 2 | 3 | 4 |
|---|---|---|---|---|
| d1 | p.d. | D | p. | 1 |
| d1 | p.d. | T | d. | 1 |
| d2 | g. | D | p. | 1 |
| d3 | i.t.d. | T | m. | 1 |
| d3 | i.t.d. | I | j. | 1 |
| d3 | i.t.d. | I | e.d. | 1 |
| d4 | p.d. | D | p. | 1 |

3.2 Corpus Representation

The source tensor is a low-level data representation merely preserving the association of statements with their provenance contexts. Before allowing for actual distributional analysis, the data have to be transformed into a more compact structure $C$ we call corpus representation. $C \in \mathbb{R}^{|A_l| \times |L| \times |A_r|}$ is a ternary (three-dimensional) labeled tensor providing for a universal and compact distributional representation of simple statements extracted from source documents. A corpus $C$ can be constructed from a source representation $G$ using functions $a : \mathbb{R} \times \mathbb{R} \rightarrow \mathbb{R}$, $w : P \rightarrow \mathbb{R}$, $f : A_l \times L \times A_r \rightarrow \mathbb{R}$. For each corpus element $c_{s,p,o}, c_{s,p,o} = a(\sum_{d \in P} w(d)g_{s,p,o,d,h(s,p,o)})$, where $g_{s,p,o,d}$ is an element of the source tensor $G$ and the $a, f, w$ functions act as follows: (1) $w$ assigns a relevance degree to each document $d \in P$; (2) $f$ reflects the relevance of the statement elements (e.g., mutual information score of the subject and object within the source); (3) $a$ aggregates the result of the $w, f$ functions’ application. This way of constructing the elements of the corpus tensor from the source representation aggregates the occurrences of statements within the input data, reflecting also two important things...
– the relevance of particular sources (via the \( w \) function), and the relevance of the statements themselves (via the \( f \) function). The specific implementation of the functions is left to applications – alternatives include (but are not limited to) ranking (both at the statement and document level) or statistical analysis of the statements within the input data.

**Example 2** A corpus corresponding to the source tensor from Example 1 can be represented as depicted below. The \( w \) values were 1 for all sources and \( a, f \) aggregated the source values using relative frequency (in a data set containing 7 statements in total).

\[
\begin{array}{ccccccc}
\text{s} & \langle p, o \rangle & \langle D, p. \rangle & \langle T, d. \rangle & \langle T, m. \rangle & \langle I, j. \rangle & \langle I, e.d. \rangle \\
\hline
p & 1/7 & 2/7 & 0 & 0 & 0 & 0 \\
p' & 1/7 & 2/7 & 0 & 0 & 0 & 0 \\
p'' & 1/7 & 2/7 & 0 & 0 & 0 & 0 \\
g & 1/7 & 0 & 0 & 0 & 0 & 0 \\
i.t.d. & 0 & 1/7 & 1/7 & 1/7 & 1/7 & 1/7 \\
\end{array}
\]

### 3.3 Corpus Perspectives

The elegance of the corpus representation lays in its compactness and universality that, however, yields for many diverse possibilities of the underlying data analysis. The analysis are enabled by the process of so called matricisation of the corpus tensor \( C \). Essentially, matricisation is a process of representing a higher-order tensor using a 2-dimensional matrix perspective. This is done by fixing one tensor index as one matrix dimension and generating all possible combinations of the other tensor indices within the remaining matrix dimension. In the following we illustrate the process on the corpus tensor from Example 2.

**Example 3** When fixing the subjects \( (A_i \text{ set members}) \) of the corpus tensor from Example 2 one will get the following matricised perspective (the rows and columns with zero values are omitted):

\[
\begin{array}{ccccccc}
\text{s} & \langle p, o \rangle & \langle D, p. \rangle & \langle T, d. \rangle & \langle T, m. \rangle & \langle I, j. \rangle & \langle I, e.d. \rangle \\
\hline
p & 1/7 & 2/7 & 0 & 0 & 0 & 0 \\
p' & 1/7 & 2/7 & 0 & 0 & 0 & 0 \\
p'' & 1/7 & 2/7 & 0 & 0 & 0 & 0 \\
g & 1/7 & 0 & 0 & 0 & 0 & 0 \\
i.t.d. & 0 & 1/7 & 1/7 & 1/7 & 1/7 & 1/7 \\
\end{array}
\]

The row and column index abbreviations correspond to Example 1. One can see that the transformation is lossless, as the original tensor can be easily reconstructed from the matrix by appropriate re-grouping of the indices.

The corpus tensor matricisations correspond to vector spaces consisting of elements defined by particular rows of the matrix perspectives. Each row vector has a name (the corresponding matrix row index) and a set of features (the matrix column indices). The features represent the distributional attributes of the entity associated with the vector’s name – the contexts aggregated across the whole corpus. This can be used for various types of analysis and for inference of more complex semantic features emerging within the simple statements extracted from the source data. In the following sections, we describe two particular types of analysis that are relevant to the motivating use case of this paper: (1) computation of related (semantically close) terms; (2) mining of conjunctive IF-THEN rules from the data.

### 3.4 Computing Related Terms

By comparing the row vectors in corpus tensor matricisations, one essentially compares the meaning of the corresponding label terms, as it is emerging from the underlying
data. For exploring the matricised perspectives, one can use linear algebra methods that have been proven to work by countless successful applications to vector space analysis in the last couple of decades\cite{16,6,12}. Large feature spaces can be reliably reduced to more manageable and less noisy number of dimensions by techniques like singular value decomposition or random indexing (see \url{http://en.wikipedia.org/wiki/Dimension_reduction}). After the (optional) dimensionality reduction, the perspective vectors can be compared in a well-founded manner by measures like cosine similarity (see \url{http://en.wikipedia.org/wiki/Cosine_similarity}) as illustrated in Example 4.

**Example 4** Let us add one more matrix perspective to the s/(p, o) one provided in Example 3. It represents the distributional features of right arguments (based on the contexts of relation terms and left arguments they tend to co-occur with in the corpus):

| x | s1 | s2 | s3 | s4 | s5 | o1 | o2 | o3 | o4 | o5 |
|---|---|---|---|---|---|---|---|---|---|---|
| y | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| y | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |

The vector spaces induced by the matrix perspectives s/(p, o) and o/(p, s) can be used for finding similar terms by comparing their corresponding vectors. Using the cosine vector similarity, one finds that \( \text{sim}_{v_1/(p, o)}(p.d., g.) = \frac{(1/7)(1/7)}{\sqrt{(1/7)^2+(2/7)^2}} = 0.2972 \) and \( \text{sim}_{o/(p, s)}(j., e.d.) = \frac{(1/7)(1/7)}{\sqrt{(1/7)^2+(1/7)^2}} = 1 \). These are the only non-zero similarities among the terms present in the corpus. This corresponds to the intuitive interpretation of the data represented by the initial statements from Example 1. Protein domains and genes seem to be different from proteins, yet protein domain is a type of domain and gene is not, therefore they share some similarities but are not completely equal according to the data. Juxtamembranes and extracellular domains are both places where internal tandem duplications can occur, and no other information is available, so they can be deemed equal (until more data comes).

It can be easily seen how the computation of related terms is relevant to the annotation use case that has motivated the paper. By computing MeSH terms related to the content of an article (i.e., terms that have been extracted from it), one can get annotations that are semantically related to the article even if they are not present in it and/or linked to it in any explicit way.

### 3.5 Rule Mining

Another type of emergent semantic pattern we can infer from the matricised corpus perspectives are IF-THEN rules. Rules are useful for our motivating use case due to their applicability to extension of the basic article annotations – once we know that an article has annotations that conform to a rule’s antecedent, we can also add annotations present in the rule consequent.

To simplify the presentation, let us consider conjunctive IF-THEN rules of type \((?x, l_1, r_1) \land (?x, l_2, r_2) \land \cdots \land (?x, l_k, r_k) \rightarrow (?x, l_{k+1}, r_{k+1}) \land (?x, l_{n}, r_{n})\) in the following, where ?x is a variable and \(l_i, r_i, i \in \{1, \ldots, n\}\) are concrete relation and (right) argument terms. An example of such rule is \((?x, \text{type of, domain}) \rightarrow (?x, \text{different from, protein}), \) which says that everything that is a type of domain is not a protein. The rule mining consists of two steps: (1) using the matrix perspective \(s/(p, o)\) for finding candidate sets of \((l_i, r_i)\) tuples that can form rules; (2) using the matrix perspective \(s/(p, o)\) for pruning the generated rules based on their confidence. Note that other types of single-variable conjunctive IF-THEN rules (i.e., the ones with variable
occurring in the second or third position of the rule statements) can be computed in the same way, only using different perspectives.

The first step corresponds to finding all frequent itemsets in a database as described in [1]. The row vectors of the \((p, o)/s\) matrix are essentially the ‘items’ – features of the rules, i.e., the concrete \((l_i, r_i), i \in \{1, \ldots, n\}\) tuples. By grouping close vectors, we can discover related features that may possibly form rules. Perhaps a simplest way of doing this is \(k\)-means clustering based on Euclidean distance [3] applied to the \((p, o)/s\) matrix. The \(k\) parameter is set so that the sizes of the generated clusters correspond to the desired maximum number of statements present in a rule. In practice, we recommend to apply dimensionality reduction to the columns of the matrix. This makes the clustering faster, while also leading to noise reduction and better representation of the features’ meaning in the sense of [6]. The described approach effectively replaces the process of finding frequent itemsets in [1]. Using our distributional representation, we find promising ‘itemsets’ not via support in discrete data transactions, but by exploiting their continuous latent semantics.

The second step involves pruning of the previously generated rules using measures of \(\text{support} (\text{supp})\) and \(\text{confidence} (\text{conf})\). Only rules with sufficiently high confidence are kept as a result of the mining process. The measures are computed on a matrix that is a transpose of the one used for generating the rules (\(sl/(p, o)\) in case of the discussed type of rules). We keep the original dimensions of the matrix this time, so that we can check for the confidence of the rules using the actual data without any transformations.

We base the rule pruning on the definitions of support and confidence provided in [1], however, we generalise the support so that we can fully exploit the power of our distributional representation. The classic definition of \(\text{supp}(X)\) for an itemset (set of features to form rule statements) is the relative frequency of rows in the data that contain the items in \(X\). This is due to the fact that the data representation in classical rule mining is crisp – the rows (transactions) contain only zeros and ones that indicate the lack and presence of an item in a transaction, respectively. Our data representation is more general – zeros in the matrix still mean lack of an item in the given row, however, the actual presence of items is represented in a more fluid way by real-valued weights. Therefore we define the generalised support as a function \(\text{supp} : 2^F \to \mathbb{R}\), where \(F\) is a set of rule features (i.e., the \((p, o)\) column labels of the corpus perspective matrix on which the rules are being tested – \(sl/(p, o)\) for the type of rules discussed above). The support of a feature set \(X\) on a perspective matrix \(M\) is computed as

\[
\text{supp}(X) = \frac{1}{||M||} \sum_{i \in I_X} \sqrt{\sum_{j \in X} m_{i,j}^2},
\]

where \(I_X\) is a set of all row indices of the matrix \(M\) where all the features from \(X\) are present (i.e., have a non-zero value), and \(m_{i,j}\) is an element of the matrix \(M\) with indices \(i, j\). \(||M||\) is a matrix norm (i.e., ‘size’) defined as \(||M|| = \sum_{i \in I} \sum_{j \in J} m_{i,j}\), where \(I, J\) are sets of row and column indices of \(M\), respectively. The confidence of a rule \(X \to Y\) is then computed as defined in [1], i.e., \(\text{conf}(X \to Y) = \frac{\text{supp}(X \cup Y)}{\text{supp}(X)}\), only using the generalised support. The process of rule mining is further illustrated in Example 5 at the end of this section.

The proposed definition of support essentially computes weighted relative frequency of the input feature set \(X\) in the matrix rows. Only rows that contain all features contribute to the absolute frequency count. The actual contribution is computed as a normalised Euclidean size of the row vector restricted only to the column indices from \(X\). The normalising factor is the size of the feature set (this to make the support value independent on the size of \(X\)). The absolute weighted frequency of the feature set is then divided by \(||M||\) to get the relative frequency (analogically to the classical def-
Articles were selected so that for each article present, the corpus also contained corresponding related articles as offered by the PubMed’s related articles service [11]. This fact was important for the evaluation later on. For the article annotation, we used the MeSH 2011 version (obtained at http://www.nlm.nih.gov/mesh/filelist.html).

4 Automated Document Annotation

This section illustrates the practical potential of the general framework introduced so far. First we describe its application to unsupervised annotation of biomedical articles with terms from the MeSH thesaurus. Then we present the evaluation of our approach and discuss the results obtained.

4.1 Data and Method

As a corpus of documents for annotation, we employed 2,003 articles from the PubMed repository (http://www.ncbi.nlm.nih.gov/pubmed/) that had their fulltexts available from PubMed Central (http://www.ncbi.nlm.nih.gov/pmc/). The articles were selected so that for each article present, the corpus also contained corresponding related articles as offered by the PubMed’s related articles service [11]. This fact was important for the evaluation later on. For the article annotation, we used the MeSH 2011 version (obtained at http://www.nlm.nih.gov/mesh/filelist.html).
We processed the data using the following high-level pipeline: (1) extraction of statements from the articles and from MeSH; (2) incorporation of the extracted statements into two separate knowledge bases for PubMed articles and for MeSH thesaurus; (3) construction of basic MeSH annotation sets for each article; (4) mining of rules from the MeSH knowledge base; (5) rule-based extension of the basic annotation sets; (6) evaluation of the initial and extended sets of annotations.

In the extraction step, we were focusing on simple binary co-occurrence statements. We tokenized the article text into sentences, then applied part of speech tagging and shallow parsing in order to determine noun phrases. Any two noun phrases $NP_1, NP_2$ occurring in the same sentence formed a statement $(NP_1, R, NP_2)$, where $R$ stands (here and in the following) for a related relationship expressing a general relatedness between the left and right arguments. Any synonyms of MeSH terms in the statements were converted to the corresponding preferred MeSH headings in order to lexically unify the data. 1,379,235 statements were generated from the 2,003 articles this way. From the MeSH data set, we generated statements from the articles and from MeSH; (2) incorporation of the extracted statements into two separate knowledge bases for PubMed articles and for MeSH thesaurus; (3) construction of basic MeSH annotation sets for each article; (4) mining of rules from the MeSH knowledge base; (5) rule-based extension of the basic annotation sets; (6) evaluation of the initial and extended sets of annotations.

The annotations for each article $d$ were computed using the article knowledge base as follows. First we constructed a set $TF = \{ (t, f_d(t)) | t \in d \}$, where $t$ are all terms extracted from $d$ and $f_d(t)$ is the absolute frequency of the term $t$ in $d$. For each $(t, f_d(t))$ tuple from $TF$, we computed another set $REL_1 = \{ (t', f_d(t) \cdot sim_{s/(p,o)}(t,t'))| sim_{s/(p,o)}(t,t') > 0 \}$. Rephrased in prose, the $REL_1$ set contained tuples of all terms similar to $t$ and the actual similarities multiplied by the $f_d(t)$ frequency (more frequent terms should generally produce terms with higher relatedness value). The $sim_{s/(p,o)}$ similarity function was defined as in Example 4. Eventually, we collated the particular term relatedness values across the whole document $d$ into an overall relatedness $rel(t') = \frac{1}{|W_t|} \sum w \in W_t w$, where $W_t' = \{ r | (t', r) \in \bigcup_{d \in d} REL_d \}$ and $W$ is a sum of all the relatedness values occurring in the $\bigcup_{d \in d} REL_d$ union. The final output of this step for each document $d$ was a set of all related terms $t'$ such that $t'$ is in MeSH. The $rel(t')$ values were used for ranking the set of MeSH annotations and taking only the top ones if necessary.

The rule mining part of the experimental pipeline was executed iteratively with different random initialisations of the clusters until no new rules were added in at least 10 most recent iterations. We obtained 33,384 rules with confidence at least 0.5 this
way. The rules were then used for extending the basic article annotation sets as follows. Let us assume an article $d$ has annotations $\{t_1, t_2, \ldots, t_n\}$. Then for any rule $(?x, R, e_1) \land (?x, R, e_2) \land \cdots \land (?x, R, e_k) \rightarrow (?x, R, e_{k+1}) \land (?x, R, e_{k+2}) \land \cdots \land (?x, R, e_m)$ such that $\{t_1, t_2, \ldots, t_n\} \subseteq \{e_1, e_2, \ldots, e_k\}$, we used the consequent set $\{e_{k+1}, e_{k+2}, \ldots, e_m\}$ as extended annotations for the article $d$. The relatedness measure of the extensions $e$ was computed as $\frac{1}{W} \sum_{w \in C_e} w$, where $C_e$ is a set of confidences of all rules that contributed with the extension $e$, and $W$ is a sum of all such confidences across all extensions computed. Similarly to the basic annotation sets, the relatedness of the extensions was used for their ranking and possible restriction to top-scoring ones.

Note that the data we have been working with, as well as the library and scripts we have implemented for the experiment, are available for reference at http://dl.dropbox.com/u/21379226/aaai

### 4.2 Evaluation and Discussion

To evaluate the annotation sets produced in the experimental pipeline, we used two methods. Firstly, we measured precision and recall of the basic and extended annotation sets based on their comparison with manually provided MeSH annotations of the corresponding articles (available through the PubMed’s Entrez API). For each article, we computed average precision, precision and recall [12] of all computed annotations and also of top $h$ ones, where $h$ is the number of human annotations for the given article.

The second evaluation method focused on the utility of the computed annotations, namely in the task of finding related articles. We used a standard vector space model [16] for determining the relatedness of documents, where features were formed by the sets of computed or manually assigned article annotations. For each document, we computed different sets of related documents (based on the human annotations and on the basic/extended ones generated by our framework). To determine their precision and recall, the computed sets were compared to corresponding sets of related articles provided by the dedicated PubMed service.

Similarly to the evaluation of annotations themselves, we measured average precision, precision and recall of all and of top $h$ related articles computed, where $h$ was the number of related articles in the gold standard.

The results of the evaluation are summarised in Tables 1 and 2. The mean average precision (MAP), precision and recall lines in the tables were computed as an arithmetic mean across the particular values for all 2,003 articles in the experimental corpus. The F-score ($F_1$ in particular) was computed from the mean precision/recall values. The columns in the tables correspond to the types of the article annotation sets described above. BASE, EXT. refer to the basic and extended annotations, while ALL, TOP refer to complete and top-$h$ only annotation sets. Note that we did not include the EXT./TOP annotations into the result summaries, since they were performing significantly worse than the other ones in most of the measured categories.

The comparison with the manually curated MeSH annotations in Table 1 does not look particularly impressive, with highest precision and recall values of 16.4% and 12.7%, respectively. On the other hand, the automatically computed annotations per...
formed much better than the ‘manual’ ones when using them as features for finding related articles. As can be seen in Table 2, there is a substantial improvement namely regarding precision and overall F-score. The only measure where the manually curated annotations perform slightly (ca. 1.1-times) better than the next-best automated method is recall. Especially notable is the difference in precision – the extended annotations achieve more than 91%, which is about two-times better than the human ones.

The results we obtain may have several interpretations. We believe that one of the more plausible ones is related to the nature of the manually provided MeSH annotations. As mentioned for instance in [13], the goal of PubMed annotators is to provide best MeSH ‘tags’ for the purpose of indexing in digital library collections. Thus they are motivated to select annotations that better discriminate papers from each other. This may, however, be rather detrimental when the task is to identify related papers using the annotations, as features used for identifying relatedness (i.e., similarity) are often dual to the features used for discrimination of entities [18]. This reasoning can in turn explain why our automatically computed article annotations, apparently very different from the manually curated ones, perform significantly better when used as features for finding related articles. The better performance (especially in case of the precision of extended annotations) may indicate that the automatically computed annotations are selected in a more fine-grained manner and from a more varied ‘vocabulary’ than the ones provided by human annotators, who can hardly grasp the scale of all the hypothetically available annotations (in addition to having different motivations as mentioned before). This is not to say that either kind of annotations is worse than the other, it much rather means that they simply serve slightly different purposes.

To conclude the discussion, we believe that despite of the low performance of our approach in terms of comparison with manually curated MeSH annotations, we can still offer potentially very beneficial results (especially in case of annotations augmented by emergent rules). This holds particularly for use cases where the annotations are supposed to be produced in a scalable and economical way in order to determine similarities between articles. Examples of such use cases include not only identification of related documents, but also question answering or automated linking of publications and supplementary data (e.g., biomedical data in the RDF format provided at http://linkedlifedata.com/sources) which we can easily incorporate as implied by [14].
5 Conclusions and Future Work

We presented an approach to acquisition of complex knowledge patterns emerging within simple statements extracted from textual data. The distinctive features of our approach are unification of the principles of emergent and distributional semantics, and a novel method for mining rules from the proposed distributional representation. To demonstrate the practical relevance of our work, we applied it to annotation of PubMed articles with terms from the MeSH thesaurus. After discussing our results, we identified areas where our approach can likely bring most benefits to users.

In future, we will explore more use cases and investigate other types of knowledge patterns (e.g., emergent formation of new candidate concepts and taxonomical relations to be recommended for inclusion into the MeSH thesaurus). Regarding the presented use case, we intend to look into possible combinations of our approach and relevant state of the art (namely the ranking-based methods like [9] or [15]). This is also related to deeper evaluation of our work that would utilise the state of the art approaches as a base-line (currently we were not able to do so comprehensively enough due to lack of publicly available and applicable implementations). Eventually, we want to perform a qualitative evaluation of the annotations produced by our system with an assistance of domain experts.

References

[1] Rakesh Agrawal, Tomasz Imieliński, and Arun Swami. Mining association rules between sets of items in large databases. *SIGMOD Rec.*, 22(2):207–216, 1993.

[2] A. R. Aronson, J. G. Mork, C. W. Gay, S. M. Humphrey, and W. J. Rogers. The NLM indexing initiatives medical text indexer. In *Studies in Health Technology and Informatics*, pages 268–272, 2004.

[3] M. Baroni and A. Lenci. Distributional memory: A general framework for corpus-based semantics. *Computational Linguistics*, 2010.

[4] David J. Chalmers. *Strong and Weak Emergence*. Oxford University Press, 2006.

[5] Philippe Cudré-Mauroux. Emergent semantics. In *Encyclopedia of Database Systems*. Springer, 2009.

[6] Scott Deerwester, Susan T. Dumais, George W. Furnas, Thomas K. Landauer, and Richard Harshman. Indexing by latent semantic analysis. *Journal of the American Society for Information Science*, 41(6):391–407, 1990.

[7] J.R. Firth. A synopsis of linguistic theory 1930-1955. *Studies in Ling. Anal.*, 1957.

[8] J. A. Hartigan and M. A. Wong. Algorithm AS 136: A k-means clustering algorithm. *Journal of the Royal Statistical Society. Series C (Applied Statistics)*, 28(1):100–108, 1979.

[9] M. Huang, A. Névéol, and Z. Lu. Recommending MeSH terms for annotating biomedical articles. *Journal of the American Medical Informatics Association*, 18(5):660–667, 2011.
[10] W. Kim, A. R. Aronson, and W. J. Wilbur. Automatic MeSH term assignment and quality assessment. In *Proceedings of AMIA Annual Symposium*, pages 319–323. AMIA, 2001.

[11] Jimmy Lin and W. John Wilbur. PubMed related articles: a probabilistic topic-based model for content similarity. *BMC Bioinformatics*, 8(1), 2007.

[12] Christopher D. Manning, Prabhakar Raghavan, and Hinrich Schuetze. *Introduction to Information Retrieval*. Cambridge University Press, 2008.

[13] A. Névéol, S. E. Shooshan, S. M. Humphrey, J. G. Mork, and A. R. Aronson. A recent advance in the automatic indexing of the biomedical literature. *Journal of Biomedical Informatics*, 42(5):814–823, 2009.

[14] Vít Nováček, Siegfried Handschuh, and Stefan Decker. Getting the meaning right: A complementary distributional layer for the web semantics. In *Proceedings of ISWC’11*. Springer, 2011.

[15] P. Ruch. Automatic assignment of biomedical categories: toward a generic approach. *Bioinformatics*, 22(6), 2006.

[16] G. Salton, A. Wong, and C. S. Yang. A vector space model for automatic indexing. *Commun. ACM*, 18(11):613–620, 1975.

[17] Junichi Tsujii. Refine and pathtext, which combines text mining with pathways. Keynote at Semantic Enrichment of the Scientific Literature 2009 (SESL 2009), March 2009.

[18] Amos Tversky. Features of similarity. *Psychological Review*, 84(2):327–352, 1977.