USE OF DOMAIN KNOWLEDGE FOR DIMENSION REDUCTION
Application to Mining of Drug Side Effects

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Abstract: High dimensionality of datasets can impair the execution of most data mining programs and lead to the production of numerous and complex patterns, inappropriate for interpretation by the experts. Thus, dimension reduction of datasets constitutes an important research orientation in which the role of domain knowledge is essential. We present here a new approach for reducing dimensions in a dataset by exploiting semantic relationships between terms of an ontology structured as a rooted directed acyclic graph. Term clustering is performed thanks to the recently described IntelliGO similarity measure and the term clusters are then used as descriptors for data representation. The strategy reported here is applied to a set of drugs associated with their side effects collected from the SIDER database. Terms describing side effects belong to the MedDRA terminology. The hierarchical clustering of about 1,200 MedDRA terms into an optimal collection of 112 term clusters leads to a reduced data representation. Two data mining experiments are then conducted to illustrate the advantage of using this reduced representation.

1 INTRODUCTION AND MOTIVATION

In some domains such as biology, data complexity is ubiquitous and constitutes a major challenge for Knowledge Discovery from Databases (KDD) approaches. One can anticipate that the more complex the data, the more relevant and interesting the extracted knowledge is. However, human expertise is then heavily required to supervise the KDD process especially for the data preparation and the interpretation steps. Thus, an interesting research orientation consists in exploring how domain knowledge can be used to help the expert and contribute to the KDD process in an automated way.

Complex data are usually voluminous and high-dimensional. In a classical ObjectsXAttributes representation, the number of objects reflects the volume of data to handle and the number of attributes provides the number of dimensions in the dataset. Most data mining methods become rather inefficient when the dimensionality is too large. Moreover, the patterns extracted from the data may also reveal inappropriate for interpretation by the expert due to plethoric, redundant, or non informative attributes. Data reduction is therefore a crucial issue for data preparation in complex datasets. Several methods exist and are reviewed in section 2. An interesting situation is the case where attributes are terms of a controlled and structured vocabulary. In biology and biomedicine such vocabularies (e.g. the Gene Ontology) have been created to annotate biological objects in databases (genes, proteins, etc.) in order to facilitate the retrieval of objects sharing similar annotations. Actually, these vocabularies represent basic domain knowledge in the form of semantic relationships between terms which enhance subsequent data analyses such as classification or characterization.

The present case-study deals with drugs annotated with respect to their side effects. The objects are drugs retrieved from the Side Effects Repository (SIDER) database, which compiles all side effects described in the drug package inserts (Kuhn et al., 2010). The annotation terms used in SIDER pertain from the Medical Dictionary for Regulatory Activities (MedDRA) terminology. Not less than 1,200 MedDRA terms are used in SIDER to annotate the drugs. Dealing with annotation terms taken from a hierarchical vocabulary allows to use existing strategies for attribute reduction such as generalization (Han and
However, available domain vocabularies are often organized as a rooted Directed Acyclic Graph (DAG) rather than a tree structure in which generalization is intractable. In the present study, we propose alternatively to cluster annotation terms based on their similarity in the rooted DAG using the IntelliGO similarity measure that was initially defined on the Gene Ontology (Benabderrahmane et al., 2010).

The application of the IntelliGO measure to the MedDRA vocabulary resulted in the clustering of the 1,200 MedDRA terms into an optimal collection of 112 term clusters. These term clusters led to a reduced data representation in which drugs are annotated with term clusters instead of MedDRA terms (Section 3). Several data mining experiments are then conducted to show the advantage of using the reduced representation of the data (Section 4).

2 DATA REDUCTION FOR KDD: A BRIEF STATE OF THE ART

Methods for data reduction divide into two types: feature selection and dimension reduction. Feature selection (or attribute selection) is a means of data reduction without altering the original data representation (Guyon and Elisseeff, 2003). Feature selection methods fall into two categories: filters and wrappers. Filter methods perform feature selection as a pre-processing step based on a search in the feature space. Feature subset evaluation relies on measures that use the training data properties and is carried out independently of any classifier (John et al., 1994). A typical example is the correlation-based feature selection method which eliminates redundant and irrelevant attributes by selecting those that individually correlate with the class but have little inter-correlation (Witten et al., 2011). Concerning the wrapper methods, they evaluate candidate feature subsets on the basis of their predictive accuracy (classification performance) using a learning algorithm (Kohavi and John, 1997). Most feature selection methods work with supervised classification and use the class information of the training examples to select the relevant features. However, Wrapper methods were recently proposed for feature selection upstream unsupervised learning, namely clustering (Kim et al., 2000; Dy and Brodley, 2004). Such studies focus on how to evaluate the results of clustering for each candidate feature subset. Also, a recent study suggests a filter method independent of both the learning algorithm and any predefined classes by guiding the attribute selection with formalized domain knowledge (Coulet et al., 2008).

Alternative data reduction methods alter the data representation by encoding the data into a smaller representation space. Such dimension reduction is also called feature compression. The principal component analysis is a popular example of such methods which deals with numeric and continuous data. Clustering was proposed for grouping the attributes in order to improve the classification or the clustering of textual documents (Kyriakopoulou, 2008). Here the attributes are binary and correspond to words annotating textual documents. Word clustering then relies on comparing their joint distributions in the documents over the classes (Koller and Sahami, 1996; Slonim and Tishby, 2000). Thus, the similarity measures used for clustering the word attributes are corpus-based.

In this paper, we propose that when the attributes belong to a domain-specific structured vocabulary, a better clustering of these attributes could be achieved by using a suitable semantic similarity measure.

3 SEMANTIC CLUSTERING OF ATTRIBUTES

3.1 The MedDRA Terminology

The MedDRA medical terminology is used to classify adverse event information associated with the use of drugs and vaccines. MedDRA is a part of the Unified Medical Languages System (UMLS) and is often presented as a hierarchy consisting of five levels: System Organ Class, High Level Group Term, High Level Term, Preferred Term, Lowest Level Term (MedDRA, 2007). Lowest level terms correspond to different terms for the same preferred term. In the MedDRA terminology, each term has an identifier and all the paths to the root can be downloaded. For example, for the term C0000733, we have two paths: C1140263.C0017178.C0947761.C0947846 and C1140263.C0947733.C0021502.C0851837. A careful review of the parent-child relationships shows that the MedDRA is actually not a hierarchy: about 37% of the MedDRA terms have more than one direct parent. This together with the natural oriented of term-term relationship and the absence of cycle, confers to the MedDRA terminology the status of a rooted DAG.

3.2 Term-term Semantic Similarity Measure

Two approaches exist for term-term semantic similarity measures: structure-based approaches which
exploit the structure of the vocabulary (depth, path length) and corpus-based approaches which exploit the term distribution in a corpus (annotation frequency, information content). The IntelliGO measure is a structure-based approach in which the generalized cosine similarity measure proposed by Ganesan for hierarchical vocabularies has been adapted to rooted DAGs (Benabderrahmane et al., 2010). The IntelliGO measure was initially defined for quantifying similarity between Gene Ontology (GO) terms. For two terms \( t_i, t_j \), it takes into account the maximal depth of their common ancestors (CA) and the minimal path-length (SPL) between them:

\[
Sim_{IntelliGO}(t_i, t_j) = \frac{2 \cdot \text{MaxDepth}(\text{CA})}{\text{MinSPL}(t_i, t_j) + 2 \cdot \text{MaxDepth}(\text{CA})}
\]

To calculate the semantic similarity between two MedDRA terms, the algorithm starts by retrieving for each term all their paths to the root node. Then, the set of common ancestors is defined as the intersection between the two sets containing the ancestors of the two terms. In the next step, the algorithm identifies the common ancestors having the maximal depth from the root node (MaxDepth(CA)). Note that the Depth of a MedDRA term can be calculated as the maximal length of a path from this term to the root node. After that, the algorithm calculates the shortest path length (MinSPL) separating the two terms. Finally, the semantic similarity between the two terms is computed using the equation (1).

As the values of \( Sim_{IntelliGO} \) range from 0 to 1, we also define the distance \( Dist_{IntelliGO} \) as its complement to 1.

### 3.3 Clustering MedDRA Terms

A total of 1,288 terms from the 20,037 MedDRA terms are used in the SIDER database for annotating drug side effects. Pairwise distances were calculated for these 1,288 terms. We then used the Ward’s hierarchical agglomeration algorithm (Ward, 1963) with an optimization step necessary to select the best level where to cut the dendrogram in order to obtain a set of clusters (Kelley et al., 1996). This method defines a penalty value which is function of the cluster number and the intra-cluster distance. When this value is minimal, the resulting clusters are as highly populated as possible while simultaneously maintaining the smallest average intra-cluster distance. In our case the minimal penalty value was obtained with 112 clusters which were subsequently inspected and validated by two experts. In the rest of this paper, these clusters will be defined as the term clusters (TC) which will be used as attributes for data mining.

In order to label each TC with its most representative term, we introduce a function \( \text{AvgDist}_{TC} \) associating to each TC term its average distance to all the TC terms:

\[
\text{AvgDist}_{TC}(t_i) = \frac{1}{|TC|} \sum_{j=1}^{N_{TC}} \text{dist}(t_i, t_j)
\]

Then the representative element R of a TC is the term which minimizes \( \text{AvgDist}_{TC} \). For example in Table 1, Erythema is the representative element of the TC. The label of a given TC is the concatenation of the TC number and the representative term (e.g., 54_Erythema for the TC described in Table 1). Once TC are built, they can be used for dimension reduction.

### 3.3.1 Example of TC with the average distance function calculated for each term t

<table>
<thead>
<tr>
<th>Term Cluster Element t</th>
<th>AvgDist_{TC}(t)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Decubitus ulcer</td>
<td>0.35</td>
</tr>
<tr>
<td>Rash</td>
<td>0.35</td>
</tr>
<tr>
<td>Lichen planus</td>
<td>0.32</td>
</tr>
<tr>
<td>Parapsoriasis</td>
<td>0.32</td>
</tr>
<tr>
<td>Pruritus</td>
<td>0.35</td>
</tr>
<tr>
<td>Psoriasis</td>
<td>0.37</td>
</tr>
<tr>
<td>Sunburn</td>
<td>0.35</td>
</tr>
<tr>
<td><strong>Erythema</strong></td>
<td><strong>0.31</strong></td>
</tr>
<tr>
<td>Pityriasis alba</td>
<td>0.32</td>
</tr>
<tr>
<td>Photosensitivity reaction</td>
<td>0.37</td>
</tr>
<tr>
<td>Rash papular</td>
<td>0.32</td>
</tr>
<tr>
<td>Dandruff</td>
<td>0.37</td>
</tr>
<tr>
<td>Lupus miliaris disseminatus faciei</td>
<td>0.35</td>
</tr>
<tr>
<td>Vulvovaginal pruritus</td>
<td>0.35</td>
</tr>
</tbody>
</table>
In the first experiment we search for Frequent Closed Itemsets (FCIs) in order to compare the two data representations with respect to computation time, number and relevance of the extracted FCIs. In our context, a FCI of length $n$ and support $s$ corresponds to an association of $n$ terms, respectively term clusters, shared by the maximal group of drugs corresponding to a percentage value $s$ of the whole dataset. The Zart program was used for FCI extraction on the Coron platform (Szathmary et al., 2007). The experiment was ran on a 2.6GHz processor with 1GB memory.

As for the second experiment we use the CN2-SD subgroup discovery algorithm (Lavrac et al., 2004) with the two data representations in order to check the impact of term clustering on the computation time and the produced subgroups. Given a population of objects and a property of those objects that we are interested in, subgroup discovery allows to find subgroups of objects that are statistically most interesting, i.e., as large as possible and having the most unusual distributional characteristics with respect to the property of interest. In our case, two categories of drugs are investigated with this method for identifying subgroups of drugs sharing discriminative side-effects in one category versus the other. The CN2-SD algorithm implementation used is one of the Keel software (Alcala-Fdez et al., 2009) and was executed on a 8-core 1.86GHz processor with 8GB memory.

### 4.2 Datasets Description

The category of a drug refers to its therapeutic uses. The categories of the drugs present in the SIDER DB are available in the DrugBank DB (Knox et al., 2011). We chose to perform the data mining experiments on the drugs corresponding to the two largest categories: the Cardiovascular Agents (CA) and the Anti-Infective Agents (AIA) containing respectively 94 and 76 drugs.

For each category, we built two datasets: the All dataset has for attributes the 1,288 MedDRA terms annotating the drugs in SIDER and the TC dataset has for attributes the 112 TC (as described in section 3) note that a TC is assigned to a drug if at least one member of the TC is reported as annotating the drug in SIDER. This gives four datasets CAAll, AIAAll, CACTC, and AIA TC.

### 4.3 Frequent closed Itemset Extraction with and without Term Clustering

The Zart program was executed on each dataset with a minimal support varying from 50 to 100%. Table 2 summarizes the number of FCIs produced in each case.

The first observation is the increase in the number of FCIs for a given minimal support when term clusters are used. For the AIAAll and AIA TC datasets, this increase varies from more than 3-fold for minimal supports between 50 and 70% to about 1.5-fold for higher minimal supports. For CAAll and CACTC datasets this increase goes from about 14-fold at minimal support 50 and 60% to 6-fold for higher minimal supports. This increase clearly reflects the expected role of clustering in feature reduction, namely increasing the density of the binary (Object X Attributes) relation by aggregating object properties. Accordingly, the computation time of the program was two-fold longer with the TC than with the All representation.

Content analysis of the FCIs was done after ranking FCIs according to their support. The top five ranked FCIs (plus ex-aequo) are listed for each dataset in Figure 1. With the All representation, FCIs can be very redundant. For example in the AIAAll dataset (left panel), three from the five displayed FCIs contain very similar term: nausea, vomiting, nausea and vomiting symptoms. On the contrary with the TC representation (right panel), a unique FCI contains the attribute 64

#### Table 2: Number of FCIs for each dataset when varying the minimal support value.

<table>
<thead>
<tr>
<th>Minimal support</th>
<th>50%</th>
<th>60%</th>
<th>70%</th>
<th>80%</th>
<th>90%</th>
<th>100%</th>
</tr>
</thead>
<tbody>
<tr>
<td>AIAAll</td>
<td>178</td>
<td>41</td>
<td>9</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>AIA TC</td>
<td>654</td>
<td>154</td>
<td>30</td>
<td>3</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>CAAll</td>
<td>386</td>
<td>94</td>
<td>41</td>
<td>11</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>CACTC</td>
<td>5,564</td>
<td>1,379</td>
<td>256</td>
<td>62</td>
<td>6</td>
<td>0</td>
</tr>
</tbody>
</table>

Figure 1 also shows that the FCI supports are generally higher with the TC than with the All data representation. A correspondence can be established between individual terms in the All representation and the term clusters in the TC representation as illustrated in Figure 1 (remember that a term cluster is labeled with its number and its most representative term). For example, the {54_Erythema} FCI found in the AIA TC dataset with a support of 88% contains the term cluster labeled 54_Erythema that includes the Pruritus individual term and has therefore been matched with the {Pruritus} FCI found in the AIA All dataset with a support of 79%.
Figure 1: The five most frequent FCIs are extracted from the total list of FCIs obtained for each dataset. In case of ex-aequo, all itemsets with the same support are listed.

Furthermore, combined with the lack of redundancy, this increase of FCI support leads to the discovery in the \textit{TC} datasets of frequent term clusters for which all individual terms are less frequent and therefore not considered in the \textit{All} datasets.

4.4 Subgroup Discovery with and without Term Clustering

The second experiment aims at discriminating the drugs belonging to two categories in terms of presence or absence of side effects. Indeed, the absence of side effects may also be important for drug characterization. This is possible with the CN2-SD algorithm as the input data is in attribute-value format. We ran the CN2-SD program on the following unions of datasets in which an additional attribute was added for the category information: CA\textsubscript{TC} versus AIA\textsubscript{TC}, and CA\textsubscript{TC} versus AIA\textsubscript{All}.

The first observation concerns the computation time. When term clusters are used the execution time is less than ten minutes whereas it does not resume within six days when all side effects are used. Thus, data reduction is necessary for successful execution of the CN2-SD algorithm.

In a second stage, the rules extracted from CA\textsubscript{TC} versus AIA\textsubscript{TC} dataset were analyzed. The left part of a rule is verified for a number of drugs (support) among which a certain fraction (coverage) are of the category indicated in the right part of the rule. The resulting subgroup therefore identifies a subset of drugs from this category sharing a specific profile of side effects with regard to the other category. The best rules in terms of coverage are shown in Table 3.

Table 3: Best rules (with coverage / support) extracted by the CN2-SD program for CA\textsubscript{TC} versus AIA\textsubscript{TC}.

<table>
<thead>
<tr>
<th>Rule</th>
<th>Coverage</th>
<th>Support</th>
</tr>
</thead>
<tbody>
<tr>
<td>50_Angina_pectoris = T AND 93_Bacteraemia = F AND 52_Itchy = F AND 54_Erythema = T AND 49_Folate_deficiency = F =&gt; CA (0.96/56)</td>
<td>50</td>
<td>93</td>
</tr>
<tr>
<td>31_Splenic_infarction = T AND 41_Neutropenia = T AND 42_Fed_discharge = F AND 77_Facial_pain = T AND 79_Cachexia = T =&gt; AIA (0.85/26)</td>
<td>31</td>
<td>41</td>
</tr>
</tbody>
</table>

To sum up, these results show that the data reduction used allows subgroup discovery which was impossible with the extended data representation. Further investigation by domain experts is ongoing.

5 DISCUSSION AND PERSPECTIVES

In this paper we have reported a method for dimension reduction guided by domain knowledge. The method is based on attribute clustering using a semantic similarity measure. We took advantage of our recently defined \textit{IntelliGO} similarity measure which applies to the rooted DAG structure encountered in many vocabularies. We believe that our strategy can be applied in various other biomedical context (Leva et al., 2005; Pakhomov et al., 2007). We tested our method on a dataset of 170 drugs annotated with 1,288 terms taken from the MedDRA terminology and representing the drugs possible side effects. Using \textit{IntelliGO}-based term-term distances and hierarchical clustering, we reduced data representation from 1,288 individual terms down to 112 term clusters. In this work we adopted a binary representation for the reduced data representation, \textit{i.e.}, a TC is assigned to a drug if at least one of its elements has been associated with the drug in the SIDER database. This representation ignores the impact of multiple associations between a drug and TC elements. A many-valued relation could be produced to take into account such situations. Recently described extension of formal concept analysis may help us handling such data representation (Mes-sai et al., 2008; Kaytoue-Uberall et al., 2009).

The dimension reduction method we have developed was tested with two data mining algorithms: FCI extraction and subgroup discovery. The results show that FCIs extracted from the TC data representation are less redundant and display higher supports than from the \textit{All} representation. Another consequence of the data reduction is that the expert’s task is facilitated because more relevant and explicit side effects are found among FCIs displaying high support. As for subgroup discovery, dimension reduction revealed to play a crucial role. Indeed, the program was unable to resume with the \textit{All} data representation whereas it
provided, with the reduced TC representation, quite interesting rules characterizing subgroups of one drug category versus another one. Complementary experiments can now be carried out to identify rules specific of a given category versus all other categories.

REFERENCES


