Finding patients using similarity measures in a rare diseases-oriented clinical data warehouse: Dr. Warehouse and the needle in the needle stack

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Objective: In the context of rare diseases, it may be helpful to detect patients with similar medical histories, diagnoses and outcomes from a large number of cases with automated methods. To reduce the time to find new cases, we developed a method to find similar patients given an index case leveraging data from the electronic health records.

Materials and methods: We used the clinical data warehouse of a children academic hospital in Paris, France (Necker-Enfants Malades), containing about 400,000 patients. Our model was based on a vector space model (VSM) to compute the similarity distance between an index patient and all the patients of the data warehouse. The dimensions of the VSM were built upon Unified Medical Language System concepts extracted from clinical narratives stored in the clinical data warehouse. The VSM was enhanced using three parameters: a pertinence score (TF-IDF of the concepts), the polarity of the concept (negated/not negated) and the minimum number of concepts in common. We evaluated this model by displaying the most similar patients for five different rare diseases: Lowe Syndrome (LOWE), Dystrophic Epidermolysis Bullosa (DEB), Activated PI3K delta Syndrome (APDS), Rett Syndrome (RETT) and Dowling Meara (EBS-DM), from the clinical data warehouse representing 18, 103, 21, 84 and 7 patients respectively.

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1. Introduction

There are approximately 7000 different types of rare diseases and disorders affecting a large population worldwide [1]. Collecting and analyzing patient data is critical aspect of rare and undiagnosed diseases programs that has the potential to increase the opportunity of discovering new knowledge. Because of their individual rarity, identifying patients that share similar phenotypes can be particularly challenging. To quote E. Ashley, the co-chair of the steering committee for the Undiagnosed Diseases Program, the challenge with rare disease data “is not so much finding the needle in the haystack as finding the right needle in a whole pile of needles” [2]. It may be helpful to detect patients with similar medical histories, diagnoses and outcomes from a large number of cases with automated methods. For example, when a new causal mutation for a disease is discovered, being able to find potential cases in a retrospective database by looking for patients similar to the 2 or 3 patients already diagnosed and genotyped could reduce their diagnosing journey and confirm the causal association.

The ability to retrospectively mine all patient records has proven beneficial [3]; and it has been made possible with the widespread adoption of clinical data warehouses such as i2b2 [4] and STRIDE [5] in hospitals. In most cases, the users query the data warehouse using a Boolean combination of criteria e.g., a list of signs and phenotypic traits. But in the case of rare complex disorders, it may be difficult to query the data warehouse in the quest for similar patients, with a formal and precise list of symptoms. The search for similar cases in the data warehouse based on similarity metrics would be more powerful and versatile. We have adapted the vector space approach to provide similarity metrics between rare disease patient medical records. In this paper, we present the method, its implementation and its evaluation on the Necker-Enfants Malades/Imagine data warehouse.

2. Background and significance

Both supervised and unsupervised machine learning methods [6] have been used to compute patient similarity, e.g., support vector machines. All these methods are based on learning models thus require training sets of a sufficient number of cases.

The Vector space model (VSM) was first proposed by Salton in 1968 [7]. Then it was implemented in SMART [8], an information retrieval system that computes similarity between documents represented as vectors of keywords. The matrix (documents by indexing terms) consists of binary values indicating the presence (1) or absence (0) of a term in a document. Salton demonstrated in 1968 the noticeable improvement in performance by using the term frequency weight [9] instead of binary values. Spärck Jones (1972) demonstrated the interest of the frequency of a term in a collection [10] and introduced the tf-idf (term frequency – inverse document frequency) weight.

Since SMART, the VSMs have been broadly used in information retrieval [11–17], in classification [18,19], and clustering [20]. The VSM has also been used in other applications. For example in social network analysis, Lee et al. used the VSM to find new network ties [21]. Santos et al. used Topic-based VSM approach to enhance a spam filter [22]. Castells et al. also used ontology in association with the VSM to improve the ranking of a search engine [23].

The VSM has also been used to compute similarity in the biomedical domain.

With the objective of identifying potentially related diseases based on genetic relationships, Sarkar et al. [24] proposed an adaptation of the VSM that bridges gene and disease knowledge inferred across three knowledge bases: Online Mendelian Inheritance in Man, GenBank, and Medline. In this study, the relatedness between diseases, via this network of gene-based relationships, was determined using a cosine similarity metric. The authors concluded that VSM was a potentially powerful method for exploring the complex landscape of polygenic diseases. Lee et al. [25] used a VSM and applied a cosine-similarity-based patient similarity metrics to an intensive care unit database to identify patients who are most similar to each index patient and predict their outcomes. They applied a VSM to MIMIC II structured data, including ICD-9 codes and quantitative data, and showed that their approach outperformed the standard severity scores usually used in the intensive care units.

In a study published in 2013, we applied a VSM approach to identify surgical site infections after neurosurgical procedures in full-text reports [26]. The method applied to patient narrative documents achieved a high recall score (92%) and a precision of 40%, much higher than the same approach based on ICD-10 codes (85% and precision 5%). These results are consistent with several studies that demonstrated that information extraction from unstructured clinical narratives is essential to most clinical applications [27]. For example, structured data alone is insufficient in resolving eligibility criteria for recruiting patients onto clinical studies [28].

All of these studies have suggested that the VSM approach can be effective at representing and computing similarity between patient reports. In the next section, we describe the VSM-based system that we have developed to search for similar patients attending the Imagine Institute, a research and healthcare institute focusing on genetic and rare diseases associated with the Necker-Enfants Malades Hospital in Paris.

3. Methods

The goal of this study was to explore the potential of using a VSM approach to identify potentially similar patients in a rare disease data warehouse.

3.1. Data warehouse

Necker Enfants Malades Hospital is an Assistance Publique-Hôpitaux de Paris (AP-HP) children’s hospital located in Paris, France. The hospital is specialized in rare diseases and is associated with a research institute, the Imagine Institute of Genetic Diseases. The hospital and the institute hold a joint clinical data warehouse,
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