Exploring Disease Association from the NHANES Data: Data Mining, Pattern Summarization, and Visual Analytics

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ABSTRACT

Finding associations among different diseases is an important task in medical data mining. The NHANES data is a valuable source in exploring disease associations. However, existing studies analyzing the NHANES data focus on using statistical techniques to test a small number of hypotheses. This NHANES data has not been systematically explored for mining disease association patterns. In this regard, this paper proposes a direct disease pattern mining method and an interactive disease pattern mining method to explore the NHANES data. The results on the latest NHANES data demonstrate that these methods can mine meaningful disease associations consistent with the existing knowledge and literatures. Furthermore, this study provides summarization of the data set via a disease influence graph and a disease hierarchical tree.

Keywords: Association Patterns, Clustering, Data Mining, Interactive Data Mining, Medical Informatics

INTRODUCTION

The National Health and Nutrition Examination Survey (NHANES) is a nationwide survey conducted by the National Center for Health Statistics and some other health agencies since 1971 (CDC, n.d.). It aims at providing nationally representative information on the health and nutritional status of the population and tracking changes over time.

NHANES data has been used to evaluate the prevalence and risk factors of diseases in the population and to provide health guidelines. The prevalence of a disease is the percentage of population having the disease. For example, in Beuther (2007) and Saydah et al. (2007), the NHANES data is used to study the prevalence of obesity and chronic kidney diseases over time and in different demographics groups (e.g., age, ethnicity and gender). A risk factor of a disease is a characteristic, condition or behavior that increases a person’s chance of developing the disease. The NHANES data has been used to verify the hypotheses of risk factors of chronic kidney (Saydah et al., 2007), obesity (Gangwisch et al., 2005), congestive heart failure (He et al., 2001) and some other diseases. The

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analysis results from the NHANES data have been used in the development of health related guidelines and public policies. For example, the early NHANES data revealed that the blood levels of lead among Americans were too high. The findings led to the federal regulations on reducing the amount of lead in gasoline, paint and soldered cans (Pirkle et al., 1998).

The NAHDES data contains a questionnaire component in which selected people are interviewed for their medical conditions and disease histories. It is a valuable data source for discovering disease associations among dozens of diseases. Disease associations can provide useful information in disease prevention, diagnosis and treatment.

There are some studies on evaluating correlated diseases by using statistical methods (He et al., 2001; Manjunath et al., 2003; Spence et al., 2003). The statistical methods focus on evaluating a number of pre-defined hypotheses of a set of risk factors or some associated diseases with respect to a particular disease. In contrast to the statistical methods, data mining methods aim at discovering the knowledge of associated diseases among a large number of diseases without any hypotheses. However, to the best of our knowledge, the NHANES data has not been systematically explored for mining associations among extensive diseases.

Is mining disease association patterns straightforward? One may think that association rule mining or association pattern mining (Agrawal et al., 2003) can provide an immediate solution. In an association rule about diseases \( A \Rightarrow B \), where \( A \) and \( B \) are two diseases, the probability that disease \( A \) appears in the population is called the support of the rule, and the probability that disease \( B \) appears in the condition of disease \( A \) appearing is called the confidence of the rule. Some other correlation measurements such as lift (Han et al., 2006), all-confidence (Omiecinski et al., 2003) and cosine (Han et al., 2006; Tan et al., 2002) are also proposed.

Since the number of people with diseases is usually much smaller than the number of healthy people, to mine association patterns of diseases, the support threshold often has to be set very low. Furthermore, diseases are very complex mechanisms. Different sub-types of a disease or people with different health conditions may have very different disease association patterns. Therefore, disease association patterns usually are not very strong. Consequently, a low confidence threshold has to be used in order to find many meaningful disease association patterns. Many other interestingness measures on association rules also meet some difficulties. For example, the lift for the patterns on high prevalence diseases is very different from the lift for patterns on low prevalence diseases.

Due to the complexity and diversity in disease association patterns, it is very difficult for a user to choose an appropriate threshold for a quality measure in the mining. If a user picks a low threshold in order to avoid missing some interesting patterns, the user may often be overwhelmed by a large number of rules and patterns which are hard to be analyzed and used.

To make disease association pattern mining practical and useful for health industry users, two problems need to be solved. First, to help users to understand the results, summarization of patterns and mining results should be provided. Second, a user should be able to interactively explore patterns interesting to the user.

In this paper, we tackle the problem of mining disease association patterns in the context of mining NHANES data. We develop a disease association pattern mining tool and present a detailed case study. The tool is publicly available at http://www.cs.sfu.ca/~zxing/personal/. We make the following contributions.

First, we give a direct pattern mining method to mine disease patterns and discuss the selection of measurements to rank the patterns. We propose a novel disease influence graph to summarize the top correlated patterns. The graph visualizes interactions among diseases and prevalence of diseases in the population.

Second, we develop an interactive disease pattern mining method by hierarchical clustering. It clusters the population based on disease
similarities and generates candidate disease association patterns. It also comes with a cluster visualization module. The hierarchical clustering structure groups related disease patterns together. The interactive disease pattern mining method enables users to browse the hierarchical clustering structure and interactively explore interesting sub-trees for the related disease association patterns. The hierarchical clustering structure also provides interesting insights on the relationships among diseases and disease sub-groups/sub-types.

Third, we present a detailed case study on the latest NHANES data, which demonstrates that our methods can mine meaningful disease association patterns consistent with existing knowledge and literatures.

The rest of the paper is organized as follows. In the second section of the direct pattern mining method is presented. In the third section, the interactive pattern mining method is described. An empirical study of the NHANES data is reported in the fourth section. The fifth section reviews the related work. The paper is concluded in the last section.

**DIRECT PATTERN MINING**

The NHANES data contains four components, namely demographic data, examination data, laboratory data and questionnaire data. The demographic component contains information such as age, gender and ethnic group. The disease status data is contained in the questionnaire component. To find disease association patterns, we use the demographic component and the questionnaire. The details of data preprocessing and feature selection will be described in the Section of Experiments.

After the preprocessing of the raw data set, the population in the survey is presented as a relational table $T$. Each tuple in the table represents a person. Each column represents a disease. Using diseases as features, a person $t$ is represented as a tuple $t = (t_1, t_2, ..., t_n)$, where $t_i = 1$ if $t$ had or has disease $d_i$, $t_i = 2$ if $t$ never has the disease, and $t_i = 0$ if whether the person $t$ has that disease is unknown.

To mine disease patterns, we do not consider patterns involving disease absence. A disease association pattern $D = <d_{i1}, d_{i2}, ..., d_{in}>$ represents that diseases $d_{i1}, d_{i2}, ..., d_{in}$ occur together.

To evaluate a disease association pattern, we consider two aspects. One is the prevalence of the pattern in the population, and one is the correlations among diseases in the pattern.

Given a population $T$ of size $N$, the prevalence of a pattern $D = <d_{i1}, d_{i2}, ..., d_{in}>$, denoted by $P(<d_{i1}, d_{i2}, ..., d_{in}>)$ is the percentage of people who have all the diseases in $D$. Formally, we have $P(<d_{i1}, d_{i2}, ..., d_{in}>) = |\{t \in T | t_{i1} = 1, t_{i2} = 1, ..., t_{in} = 1\}| / N$.

To evaluate the correlation among diseases in a disease association pattern, we need to choose a proper measurement. In Tan et al. (2002), they compared 21 interestingness measures for association patterns, including support, confidence, odds ratio, lift, cosine, mutual information and some others. The comparison shows that no single measurement is consistently better than the others in all application domains.

The number of people with diseases is much smaller than the number of healthy people. Moreover, we consider disease co-presence more important than co-absence. Therefore, the disease data is sparse. Han et al. (2006) and Tan et al. (2002) suggest that the correlation measurements with the null-invariance property are suitable for applications with sparse data. In our application setting, a correlation measure is null-invariant means for a disease pattern $D$, that the value of the correlation measure is not affected by the number of people who do not have diseases in pattern $D$.

For example, measure cosine is null-invariant and measurement lift is not. The measurement cosine and lift (Han et al., 2006; Tan et al., 2002) for a disease pattern of two diseases are defined as follow,
cosine \( <A, B> \) = \frac{P(<A, B>)}{\sqrt{P(<A>)P(<B>)}}

\[ \text{lift}(<A, B>) = \frac{P(<A, B>)}{P(<A>)P(<B>)} \]

Given a population, suppose disease \( A \) has prevalence 0.03, disease \( B \) has prevalence 0.02, and a pattern \( <A, B> \) has prevalence 0.01. We have

\[ \text{cosine}(<A, B>) = \frac{P(<A, B>)}{\sqrt{P(<A>)P(<B>)}} = 0.408 \]

and

\[ \text{lift}(<A, B>) = \frac{P(<A, B>)}{P(<A>)P(<B>)} = 16.67. \]

Suppose we remove 90\% of the population that do not have diseases \( A \) or \( B \) from consideration. Then, the prevalence of disease \( A \) increases to 0.3 and that of disease \( B \) becomes 0.2. The prevalence of pattern \( <A, B> \) is upgraded to 0.1. We have

\[ \text{cosine}(<A, B>) = \frac{(P(<A', B'>))}{\sqrt{P(<A'>)P(<B'>)}} = 0.408, \]

remained unchanged. This elaborates that cosine is null-invariant. On the other hand,

\[ \text{lift}(<A', B'>) = \frac{P(<A', B'>)}{P(<A'>)P(<B'>)} = 1.67 \]

Thus, lift is not null-invariant.

As shown in this example, a measure which is not null-invariant may give a higher score to patterns involving low prevalent diseases. If a user picks the top-\( k \) patterns, those patterns may mainly involve diseases with low prevalence.

Two null-invariant correlation measures, cosine and Jaccard distance, are discussed in (Tan et al., 2002). However, they are not effective in guiding disease association pattern mining since they are only applicable to patterns involving two diseases. Therefore, we want to find a null-invariant correlation measure that can handle multiple diseases in a pattern.

One possible measure in the literature is all-confidence (Omiecinski, 2003). A disease association pattern \( D \) can induce a set \( R(D) \) of \( x \) \(-2 \) association rules in the form of \( X \Rightarrow (D - X) \) such that \( X \subseteq D \). The all-confidence of \( D \) is the lowest confidence among all the rules \( r \in R(D) \).

Although all-confidence can measure the overall affiliation among all the diseases in a disease pattern, it captures the weakest correlation. For example, suppose we have a disease pattern \( <A, B> \) where \( A \) is rare with prevalence 0.03, \( B \) is common with prevalence 0.3. If all people having \( A \) also have \( B \), then \( P(<A, B>) = 0.03 \). Pattern \( <A, B> \) induces two rules, \( A \Rightarrow B \) with confidence 1 and \( B \Rightarrow A \) with confidence 0.1. The all-confidence of pattern \( <A, B> \) is 0.1. If a user sets the minimum all-confidence threshold to 0.4, the pattern, though interesting in disease association, is missed. In general, all-confidence may assign a low score to patterns involving the association between high prevalent diseases and low prevalent ones.

To serve our disease association pattern mining, we define a new null-invariant correlation measure. For a disease association pattern \( D \), the \( \text{any-cosine} \) is the highest cosine score among all the rules induced by \( D \). That is,

\[ \text{any-cosine}(D) = \max_{X \subseteq D} \left\{ \text{cosine}(X \Rightarrow (D - X)) \right\} \]

\[ \text{cosine}(X \Rightarrow (D - X)) = \frac{P(D)}{\sqrt{P(X)P(D - X)}}. \]

Easily, we can show that \( \text{Any-cosine is a null-invariant correlation measure}. \)

\( \text{Any-cosine} \) captures the most interesting rule induced from \( D \). We will compare the effectiveness of all-confidence and any-cosine in the experiments.

In our direct pattern mining, we use Apriori (Agrawal et al., 1993) to find all patterns passing a minimum prevalence threshold. Then, we rank those patterns by a correlation measure.
There may be many patterns passing the minimum support threshold. As discussed before, summarization of patterns is highly desirable. We summarize the top ranked patterns by a disease influence graph.

Figure 1 shows an example of the nodes in the graph being the diseases, and the size of each node is proportional to the prevalence of the corresponding disease. Two diseases are linked by an edge if they appear in a pattern. The thickness of the edge is proportional to the correlation score of the pattern. If two diseases appear in multiple top-ranked patterns, the thickness of the edge between them takes the highest correlation score in those patterns.

The visualization in a disease influence graph is intuitive. A disease influence graph provides an overview of the prevalence and the interactions among the diseases in a population. Moreover, in a disease influence graph, the hubs and the dense connected areas can be viewed as the influential diseases in association patterns. We will discuss more details in the Section of Experiments.

INTERACTIVE PATTERN MINING

More often than not, a user may only be interested in the disease patterns involving some certain diseases. For example, a clinical specialist may only be interested in the patterns involving diabetes. Can we provide an effective way to let the user interactively browse the disease patterns according to his/her own interests? In this section, we propose an interactive disease pattern mining method based on agglomerative hierarchical clustering (Murtagh, 1983).

Disease associations are sophisticated. People having different subtypes of a disease or people in different health conditions may have different disease co-occurrences. This motivates us to cluster people based on the similarity of their disease patterns. Intuitively, people in a cluster share some common diseases, such as a cluster of people with a kidney disease or a cluster of people with a heart disease.

There are many different clustering methods, such as partitioning based clustering and hierarchical clustering. In disease association pattern analysis, it is highly desirable to organize

Figure 1. The disease influence graph of people of age at least 20
disease associations in a hierarchical structure. For example, people having diabetes form a big cluster. Among those people, some may only have diabetes, and the others may have diabetes and some other complications. Therefore, we employ hierarchical clustering instead of partitioning based clustering.

Clustering is useful beyond pattern exploration. A cluster can be used to generate hypotheses of disease associations by summarizing its semantic meaning. For example, if we find a cluster of people with the characteristics of high blood pressure and high blood cholesterol level, a hypothesis of disease associations, \(< \text{High blood pressure, High blood cholesterol} >\) can be generated.

Figure 2 demonstrates our interactive disease pattern mining system. Starting from the whole hierarchical clustering structure of the population, a user can click on a sub-cluster. The detailed distribution of the diseases in the selected sub-cluster will be visualized. Moreover, using the dominating diseases in the sub-cluster, a hypothesis disease association pattern is generated. If the user is interested in such a pattern, he/she can further test the pattern against the whole population using various statistical measures. The analysis can be conducted recursively in the hierarchical structure.

In order to obtain interesting clustering structures, a proper similarity measure between people is important. Recall that we represent a person \(p\) as a disease vector \(\langle d_1, d_2, \ldots, d_n \rangle\), where \(d_i = 1\) if \(p\) had or has disease \(d_i\), \(d_i = 2\) if \(p\) never has the disease, and \(d_i = 0\) if whether the person has that disease is unknown. Given two persons

\[ p_x = \langle d_{x1}, d_{x2}, \ldots, d_{xn} \rangle \]

and

\[ p_y = \langle d_{y1}, d_{y2}, \ldots, d_{yn} \rangle \]

where \(d_{x1}, d_{x2}, \ldots, d_{xn}\) and \(d_{y1}, d_{y2}, \ldots, d_{yn}\) are the disease variable values of \(p_x\) and \(p_y\), respectively, we define the similarity between disease values as

\[
\text{Sim}(d_{xi}, d_{yi}) = \begin{cases} 
0 & \text{if } d_{xi} = d_{yi} \text{ and } d_{xi}, d_{yi} = 0 \\
1 & \text{if } d_{xi} = d_{yi} \text{ and } d_{xi}, d_{yi} = 0 \\
2 & \text{if } d_{xi} = d_{yi} 
\end{cases}
\]

Moreover, we define the similarity between \(p_x\) and \(p_y\) on the diseases known to them as

\[
\text{Sim1}(p_x, p_y) = \frac{\sum_{i \in \{d_{xi}, d_{yi} \neq 0\}} \text{Sim}(d_{xi}, d_{yi})}{|\{i \mid d_{xi} \neq 0 \land d_{yi} \neq 0\}|}
\]

\(\text{Sim1}(p_x, p_y)\) is the normalized sum of weighted similarities on all diseases whose values are not missing. We use the sum of weighted similarities because we consider two persons more similar if they share some common diseases rather than they both do not have some diseases.

The NHANES survey data contains a significant amount of missing data. The questions on diseases are presented to different age groups. Some diseases are specific for some age groups. For example, disease Glaucoma is only applicable to people over 40 years old. For people under 40 years, the data on this disease is missing. Furthermore, even if a person is interviewed for the status of a certain disease, the data may still be missing since the person may refuse or cannot answer the interview question. In order to handle missing data, in \(\text{Sim1}\), the sum of weighted similarities is normalized by the number of diseases without missing values.

The normalization may cause a bias of similarity on people with more missing data. In order to reduce the bias, we design \(\text{Sim2}(p_x, p_y)\), which is defined as the similarity between \(p_x\) and \(p_y\) on the diseases common to them that is,

\[
\text{Sim2}(p_x, p_y) = \frac{|\{i \mid d_{xi} = 1 \wedge d_{yi} = 1\}|}{n}
\]

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The distance between $p_x$ and $p_y$ is the combination of $Sim1()$ and $Sim2()$, which is defined as follows,

$$\text{Dist}(p_x, p_y) = (2 - Sim1(p_x, p_y)) + (1 - Sim2(p_x, p_y))$$

The range of $\text{Dist}(p_x, p_y)$ is between 0 to 3. The smaller the distance is, the more similar the two persons are.

Hierarchical clustering builds a hierarchical clustering tree of clusters. We use the average pair-wise distance of two clusters as the inter-cluster distance, since the average distance is more robust to noise data comparing to single link hierarchical clustering (Murtagh, 1983).

We visualize a cluster by a bar chart, which plots the prevalence of the diseases in the cluster. A cluster is summarized by its dominating disease pattern. A disease is a dominating disease of a cluster if the prevalence of the disease in the cluster passes a threshold. In this paper, we consider a disease in a cluster dominating if its prevalence in the cluster is higher than 75%. A dominating disease pattern is composed of all dominating diseases in the cluster. For example, in Figure 2, the dominating disease pattern of the picked cluster is <High blood pressure, High blood cholesterol level>). A cluster may be dominated by a single disease or multiple diseases. As a special case, if the prevalence of all diseases in a cluster is 0, the dominating disease pattern is <healthy people>. A cluster may not have a dominating disease pattern if it is too big, such as a cluster of the whole population.

At this point, the advantage of hierarchical clustering against partitioning based clustering in interactive mining of disease association patterns becomes clear. The partitioning-based clustering methods, such as k-mean, need the
number of clusters as a parameter. Setting the number of clusters improperly may produce a messy clustering result. Hierarchical clustering does not require a user to specify the number of clusters. By summarizing the clusters, hierarchical clustering automatically provides a natural partitioning of the population. When a user browses down the hierarchical clustering tree from the root, at the top levels, the clusters may not have dominating disease patterns. The descendant clusters closest to the root which have dominating disease patterns form a natural partitioning of the population. We call those clusters the natural clusters. Importantly, in a natural cluster, most of the dominating disease patterns contained in its sub-clusters are the extended patterns of the dominating disease pattern in the natural cluster. When browsing the hierarchical clustering tree by clicking the sub-trees, a user can view how the related dominating disease patterns change in a hierarchical structure.

A user may also be interested in extracting all dominating disease patterns in a hierarchical cluster tree. We can output all patterns by recursively traversing the hierarchical cluster tree from the root. We use two parameters, a size threshold and an entropy threshold of a cluster, as the stopping conditions for the recursive traversing to select only significant dominating disease patterns.

A very small cluster is statistically insignificant. The dominating disease patterns extracted from such small clusters are not interesting in statistical analysis. Therefore, we can use a size threshold to filter out small clusters.

Consider a cluster $C$ with $m$ diseases, denoted by $Fre(C) = \{f(d_1), f(d_2), ..., f(d_m)\}$. We define the entropy of a disease $d_i$ ($1 \leq i \leq m$) as

$$Entropy(d_i) = -f(d_i) \log(f(d_i)) - (1 - f(d_i)) \log(1 - f(d_i))$$

Moreover, the entropy of the cluster is

$$Entropy(C) = \sum_{i=1}^{m} \frac{Entropy(d_i)}{m}$$

The entropy of a cluster measures how coherent the cluster is in terms of the disease pattern. The smaller the entropy is, the more coherent the cluster is. For example, if in a cluster every disease has prevalence 0.5, the entropy is 1. The cluster does not have a dominating disease pattern and need to be further divided. In another cluster, if all the people do not have any disease, i.e., the cluster contains only healthy people, or all people share the exactly same disease(s), the entropy is 0. Using an entropy threshold, if a cluster is already very coherent, we do not further visit its sub-clusters.

Our interactive pattern mining tools is publicly available at http://cs.sfu.ca/~zxing/personal/. The tool is developed in C# using Microsoft Visual Studio 2005. As shown in Figure 1, the graphic user interface (GUI) contains the following functions.

1. Displaying the complete hierarchical clustering dendrogram;
2. Enabling clickable sub-clusters;
3. Visualizing the disease dominating pattern in a sub-cluster when the sub-cluster is clicked;
4. Computing the statistical measurements of a disease dominating pattern.

The graphic user interface is independent of particular hierarchical clustering methods. The GUI displays a dendrogram by taking an input file which represents the structure of a dendrogram. The GUI can easily be adopted for any applications which need the functions of dynamically exploring sub-clusters in a dendrogram and summarizing the dominating features in sub-clusters.

**EXPERIMENTS**

In this paper, we use the NHANES data of year 2005-2006. The data set contains in total...
10,348 people. In the questionnaire component, interviewees are asked about the histories on a collection of diseases, such as allergies, cardiovascular diseases, kidney diseases, and diabetes. The disease statuses are obtained from the self-reported answers in the following questions, “Has a doctor or other health professional ever told you/SP (for spouse) that you have/s/he/SP have...?”. We include the disease statuses obtained by the above questions. In total, the questionnaire covers 26 disease statuses/medical conditions. Some diseases such as heart disease are combined from several subtypes of appearing in the raw data such as congestive heart failure and coronary heart disease. We do not include any diseases about emotional health such as depression. The questions on diseases are presented to different age groups. For people of age 40 and older, all the 26 diseases are interviewed. To people of age between 20 and 40, 23 diseases are presented. To people younger than 20, only 9 diseases are presented.

In this report, we analyze the data on the people of age 20 and older. There are in total 4,979 people in this age group.

The Result of Direct Pattern Mining

The direct data mining method obtains 1,639 disease association patterns by setting the minimal support threshold to 0.5%

To compare the effectiveness of various correlation measures, Table 1 lists the top-10 patterns in three measures, all-confidence, any-cosine and any-lift. For a disease pattern, Any-lift is the highest lift among all the rules induced from the patterns using all the item. From Table 1, we can see that the top-10 patterns ranked by all-confidence are similar to the top-10 patterns ranked by any-cosine: 7 out of the 10 patterns are in common, as highlighted in the table. Any-lift only finds one pattern common with the top-10 patterns in all-confidence and any-cosine. Each of the top 10 patterns in any-lift involves at least one disease of low prevalence, such as chronic bronchitis and kidney diseases. As analyzed in the Section of Background, this is because lift is not null-invariant.

Among all the patterns in the top 10 lists in all-confidence and any-cosine, <Arthritis, High BP, High BC> is the only long pattern involving 3 diseases which is ranked high by any-cosine, but low by all-confidence. By examining the ranked lists in any-cosine and all-confidence, we find that the major difference between the two measures is that any-cosine ranks some long patterns high, while all-confidence always ranks short patterns high. For example, among the top 50 patterns ranked by any-confidence, there are only 7 long patterns and there are 12 long patterns in the top-50 list of any-confidence. This clearly shows that any-cosine can promote long patterns which induce interesting rules, while all-confidence cannot.

The top-ranked disease association patterns in Table 1 can be well explained by the existing medical knowledge. For example, pattern <High BP, High BC> is consistent with the well known fact that high blood pressure and high blood cholesterol level have a high chance to happen together. Pattern <Arthritis, High BP> is supported by (Biomedical.Org, n.d.; Forman et al., 2007), which showed that some pain reliever drugs used frequently by people of arthritis increase the risk of high blood pressure. Pattern <Arthritis, Overweight> is consistent with the finding in (Gill et al., 2005), which showed that overweight and obesity are strongly related to arthritis.

In addition to viewing the top patterns one by one, a user may also see an overall picture of the disease interactions on the population. In Figure 1 (the graphics is generated by the TouchGraph Navigator package http://www.touchgraph.com/navigator.html), the top 50 patterns ranked by any-cosine are visualized using a disease influence graph defined in Section 2. From the graph we can see that Hb (High blood pressure), Al (Allergies), Ov (Overweight), Hc (High blood cholesterol), Ar (Arthritis), Tr (Trouble seeing even with glasses) are the top six high prevalent diseases, and they influence each other intensively by forming a clique in
Table 1. Top-10 patterns by various measures

<table>
<thead>
<tr>
<th>All-confidence</th>
<th>&lt;High BP, High BC&gt;</th>
<th>&lt;Overweight, High BP&gt;</th>
<th>&lt;Arthritis, High BP&gt;</th>
<th>&lt;PSA, Prostate&gt;</th>
<th>&lt;Overweight, High BC&gt;</th>
<th>&lt;Arthritis, High BC&gt;</th>
<th>&lt;Overweight, Allergies&gt;</th>
<th>&lt;Arthritis, Allergies&gt;</th>
<th>&lt;Overweight, Arthritis&gt;</th>
<th>&lt;PSA, Arthritis&gt;</th>
<th>&lt;PSA, Asthma, High BP, High BC&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any-cosine</td>
<td>&lt;PSA, Prostate&gt;</td>
<td>&lt;High BP, High BC&gt;</td>
<td>&lt;Arthritis, High BP&gt;</td>
<td>&lt;Hay fever, Allergies&gt;</td>
<td>&lt;Overweight, High BC&gt;</td>
<td>&lt;Arthritis, High BC&gt;</td>
<td>&lt;Overweight, Allergies&gt;</td>
<td>&lt;Arthritis, Allergies&gt;</td>
<td>&lt;Overweight, Arthritis&gt;</td>
<td>&lt;PSA, Asthma, High BP, High BC&gt;</td>
<td>&lt;Hay fever, Allergies&gt;</td>
</tr>
<tr>
<td>Any-lift</td>
<td>&lt;PSA, Prostate&gt;</td>
<td>&lt;Emphysema, Chronic Bronchitis&gt;</td>
<td>&lt;Cancer, PSA, Prostate&gt;</td>
<td>&lt;Cancer, PSA&gt;</td>
<td>&lt;Heart Disease, Diabetes, Kidney&gt;</td>
<td>&lt;Heart Disease, Stroke&gt;</td>
<td>&lt;Heart Disease, Cardiovascular&gt;</td>
<td>&lt;Heart Disease, Stroke, Diabetess&gt;</td>
<td>&lt;Heart Disease, Emphysema&gt;</td>
<td>&lt;Heart Disease, Stroke, Cardiovascular&gt;</td>
<td>&lt;Cancer, PSA, Prostate&gt;</td>
</tr>
</tbody>
</table>

| Rank | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |

There are another 8 diseases which are directly connected with the clique through one or multiple edges. In other words, those diseases are 1 edge away from the clique. For example, Di (Diabetes) interacts with live diseases in the clique except for AI (Allergies). He (Heart disease) is linked with Di (Diabetes), Hb (High blood pressure), Ar (Arthritis), and Hc (High blood cholesterol) in the clique. There are several branches extending from the clique. For example, there is a branch of As (asthma) and Ch (Chronic bronchitis), which is a branch of allergy related diseases. There is also a branch of Cn (Cancer), Pr (Prostate diseases) and Ps (PSA test abnormal), which can be regarded as a cancer branch. Some diseases, such as Li (Liver condition) and Ki (Kidney diseases), are isolated in the graph. From this example, we can see that the disease influence graph provides an overview of the prevalence of diseases and their interactions measured by any-cosine.

The Result of Interactive Pattern Mining

Our interactive pattern mining method generates 23 natural clusters. In 22 of them, each is dominated by a single disease. The remaining natural cluster contains 905 healthy people who do not have histories on any diseases. There are in total 908 healthy people in the data set. 3 of them are merged into other clusters due to some missing data. This clearly shows that the hierarchical clustering method can naturally identify groups of people sharing similar disease patterns.

The largest natural cluster contains 1,392 people and is dominated by disease high blood pressure. The second largest cluster is the one of 905 healthy people.

Some diseases such as asthma do not dominate a natural cluster. The people with asthma are mainly within the cluster of high blood pressure and the cluster of allergies.

The sub-clusters as descendants of a natural cluster lead to disease dominating patterns closely related to the pattern of the natural cluster. For example, within the natural cluster of high blood pressure, the three biggest sub-clusters dominated by two diseases are the sub-cluster dominated by <Overweight, High BP>, the sub-cluster dominated by <High BP, High BC>, and the sub-cluster dominated by <Arthritis, High BP>. The hierarchical clustering tree groups related disease association...
patterns together, and helps a user to browse the tree according to her interests.

Figure 3 shows more examples of interesting sub-clusters. Among those examples, some disease association patterns in the clusters are consistent with the existing medical knowledge or literatures. For example, the cluster dominated by pattern <Arthritis, Allergies> is supported by (Panush et al., 1990) that some cases of arthritis are triggered by food allergies. Moreover, a cluster of people dominated by overweight and sleep disorder verifies the association between sleeping disorder and overweight reported in (Gangwisch et al., 2001).

Furthermore, some disease patterns captured by the clusters reveal new findings. In Figure 3, we list four clusters with dominating disease patterns of

<TroubleSeeing(MCQ140),
LiverCondition(MCQ160L),
Arthritis(MCQ160A),
HighBloodCholesterol(BPQ080)>,
<TroubleSeeing(MCQ140),
Arthritis(MCQ160A)>
and <Cancer(MCQ220),
HighBloodCholesterol(BPQ080)>

To the best of our awareness, those patterns are not well discussed by existing medical literatures. Interestingly, by searching those patterns on the internet, we found the associations of diseases in those patterns have already attracted attentions by people. For example, on an online discussion board (http://www.healthcentral.com/rheumatoid-arthritis/c/question/54342/29748), people asked “can RA(rheumatoid arthritis) cause high cholesterol? “ On a MSN online discussion board (http://health.msn.com/healthtopics/pain-management/arthritis/articlepage.aspx?cp-documentid=10020060), people asked “Are there vision disorders associated with rheumatoid arthritis?” Those new findings from NHANES data invite further investigation and verifications.

The hierarchical clustering tree conveys interesting information about disease subgroups and disease subtypes. For example, Figure 4 shows the sub-tree rooted at the natural cluster dominated by diabetes. The dominating diseases of some sub-clusters are tagged for easy understanding. The whole nature cluster, cluster 1, is composed of two sub-clusters, cluster 2 and cluster 3. Cluster 2 is dominated by heart disease and diabetes, while cluster 3 is dominated by diabetes.

Cluster 2 is further divided into clusters 4 and 5. Cluster 4 is dominated by heart disease, diabetes and stroke. This hierarchical cluster sub-tree suggests that many people with diabetes may also have heart disease. Among the people with diabetes and heart disease, stroke may appear. An article at the National Diabetes Information Clearinghouse website (NDIC, n.d.) suggests “people with diabetes are at least twice as likely as someone who does not have diabetes to have heart disease or a stroke”.

In cluster 3, a part of people form sub-cluster 7, which is dominated by diabetes, overweight and thyroid problem. The other people in cluster 3 form cluster 7, which is a diabetes cluster. Further dividing cluster 6, we can get cluster 8 and cluster 9. For cluster 9, the dominating diseases are diabetes and vision problem. It is confirmed by (Klein et al., 1995), “diabetes is the leading cause of new cases of blindness in people age 20-74 years in the United States”.

Diabetes and its treatments may cause many complications. The above hierarchical clustering sub-tree provides an insight on the sub-groups of patients with diabetes according to different situations of complications.

**Comparisons**

In addition to browsing the hierarchical clustering tree, our method can also output all disease association patterns by setting a minimum size threshold and a minimum entropy threshold. When setting the minimum size threshold to 5 (i.e., a cluster should contain at least 5 people)
Figure 3. Examples of sub-clusters

and the minimum entropy threshold to 0.01 (i.e., we do not visit sub-clusters of a cluster of entropy less than 0.01), our method outputs 432 disease dominating patterns.

Can those patterns obtained from the hierarchical clustering tree cover the strong correlation patterns found by the direct mining methods? Table 2 makes a comparison. In the table, the value 10 in the cell of the row named “Support” and the column named “1~10” indicates that among the top 10 patterns ranked by measure support in the direct mining method, all of them are contained in the set of patterns generated by the hierarchical clustering tree. The other cells in the table can be explained similarly.
The top 10 patterns ranked by support, any-cosine and all-confidence are all among the patterns generated by the hierarchical clustering tree. 68 patterns of the top 70 in support can be generated by the hierarchical clustering tree, which can be translated into a recall of 97%. 63 of the top 70 patterns in any-cosine can be generated by the hierarchical clustering tree, that is, a recall of 90%. Among the top 70 patterns in all-confidence, 59 can be generated by the hierarchical clustering tree. The recall is 84.3%.

The number of patterns generated by the hierarchical clustering tree is smaller than the number of patterns generated by the direct mining method, however, a large percentage of the top patterns ranked in support, any-cosine and all-confidence can be generated by the hierarchical clustering tree.

For some highly ranked patterns which are not among the patterns generated by the hierarchical clustering tree, the descendants of those missing patterns can usually be found in the tree. For example, pattern $D = \langle \text{Heart disease, Arthritis} \rangle$ is ranked the 17-th in any-cosine. No cluster can be found in the hierarchical clustering tree having $D$ as the dominating pattern. However, we find two clusters in the hierarchical clustering tree having dominating patterns $\langle \text{Heart disease, Arthritis, High BP.} \rangle$ and $\langle \text{Heart disease, Arthritis, Diabetes} \rangle$, respectively. Those descendant patterns can be used to generate hypotheses about the associations between heart disease and arthritis.

**RELATED WORK**

NHANES data has been used to find associated diseases using statistical methods. For example, by using statistical measuremen...
diabetes and overweighting are two independent risk factors for congestive heart. By using the Chi-square test, Manjunath et al. (2003) showed that the level of kidney function is associated with atherosclerotic cardiovascular disease. Spence et al. (2003) studied the association between arthritis and high blood pressure.

Comparing to statistical methods, our data mining methods have the following advantages.

1. The existing statistical methods on analyzing the NHANES data mainly focus on testing a small number of hypotheses of disease associations or risk factors of diseases. While data mining methods aims to mine many disease patterns among dozens of diseases.

2. As we discussed above, different statistical methods are used to evaluate different pre-defined disease patterns. There is a lack of comparisons by using one measurement to evaluate the disease patterns on a given population. Our data mining methods enable the users to compare many disease patterns based on one population by ranking the patterns using a particular correlation measurement.

3. Although statistical methods are good at evaluating a particular disease pattern, they often have difficulties to provide an overview of the disease interactions on a population. Our disease influence graph can provide a summarization and visualization of the overall structure of disease interaction on a population. If we have several different populations, such as people among different age ranges, we can easily compare the disease interactions structures in different population by using disease influence graph.

4. As shown before, our interactive disease pattern mining system can discover the hierarchical structure of diseases. The statistical methods cannot easily achieve that.

In summary, statistical methods focus on evaluating a particular disease pattern. Data mining methods are good at systematically generating many hypotheses of disease patterns, and providing different kinds of visualizations, comparisons and summarizations over the disease patterns. In our data mining system, we used several correlation measurements to evaluate disease patterns, such as all-confidence, all-cosine. Our system can also incorporate other statistical measurements to enhance our analysis.

Recently, some data mining methods have been applied to analyze the NHANES data. Walton et al. (2008) applied decision trees and linear regression to classify people into different categories of health conditions (e.g., excellent, fair or bad). Lee et al. (2008) used decision trees and association rules to find dependence among the laboratory and health condition variables in the latest NHANES data. Different from this study, the major findings in Lee et al. (2008) are on the dependence between the diet behaviors or medical conditions and disease absence. In this paper, we focus on the association among diseases.

Our direct pattern mining method uses null-invariant measures to rank disease patterns and summarize the top ranked patterns by a disease influence graph. The principles on choosing suitable measures to mine interesting patterns for different application domains are discussed in Tan et al. (2002). A method of mining risk patterns in medical data measured by relative risk is proposed in Li et al. (2005). Some methods of summarizing frequent patterns are proposed in Jin et al. (2008) and Yan et al. (2005).

In the interactively pattern mining method, hierarchical clustering is used. In bioinformatics, hierarchical clustering has been used in interactively exploring gene expression patterns, such as Saldaña et al. (2004) and Jiang et al. (2003).

**DISCUSSIONS**

In this paper, we tackled the problem of mining disease associations. We proposed the direct pattern mining method and the interactively
pattern mining method. We applied the proposed methods on NHANES data. Our findings are consistent with the existing medical knowledge and literatures. In addition, interesting summarizations through a hierarchical disease tree and a disease influence graph provide insights into the relationships among diseases. Carrying the success on analyzing the NHANES data, our methods can be applied to analyze other data about disease association as well. We already made our software available for public use. In the future, we plan to enhance our mining tool through more statistical analysis and better visualization.

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