Neighborhood outlier detection

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ABSTRACT

KNN (k nearest neighbor) is widely discussed and applied in pattern recognition and data mining, however, as a similar outlier detection method using local information for mining a new outlier, neighborhood outlier detection, few literatures are reported on. In this paper, we introduce neighborhood model as a uniform framework to understand and implement neighborhood outlier detection. Furthermore, a neighborhood-based outlier detection algorithm is also given. This algorithm integrates rough set granular technique with outlier detecting. We propose a neighborhood-based metric on outlier detection, and compare neighborhood outlier detection with DIS, KNN and RNN. The experimental results show that neighborhood-based metric is able to measure the local information for outlier detection. The detected accuracies based on neighborhood outlier detection are superior to DIS, KNN for mixed dataset, and a litter better than RNN for discrete dataset.

1. Introduction

In contrast to traditional pattern recognition question that aims to construct a general pattern map to the majority of data, outlier detection targets to find the rare data whose behavior is very exceptional when compared with rest large amount of data. One of the most popular outlier detection techniques is distance-based outlier, introduced by Knorr and Ng (1998, 1999). A distance-based outlier in a dataset D is a data object with p% of the objects in D having a distance of more than dmin away from it. This notion generalizes many concepts from distribution-based approach and enjoys better detected accuracy. What is more, it is extended based on the distance of a point from its kth nearest neighbor, which is called KNN method (Ramaswamy, Rastogi, & Kyuseok, 2000). It ranks the top k points by the distance to its kth nearest neighbor as the outliers. Efficient algorithms for mining top-k outliers are also studied. Furthermore, in the algorithm proposed by Angiulli and Pizzuti (2002), the outlier factor of each datum point is computed as the sum of distances from its k nearest neighbors, which obtained better result comparing with traditional KNN. However, as KNN outlier detections computing all the dimensional distances of the points from one another, it is time-consuming if the available objects are of very great size. Besides, the direct application of KNN methods to high dimensional problems often results in unexpected performance and qualitative costs due to the curse of dimensionality.

With increasing awareness on outlier detection in literatures, more concrete meanings of outliers are defined for solving problems in specific domains (Breunig, Kriegel, Ng, & Sander, 2000; Jain, Murty, & Flynn, 1999; Jiang, Sui, & Cao, 2009; Johnson, Kwok, & Ng, 1998; Kovacs, Vass, & Vidacs, 2004; Rousseeuw & Leroy, 1987). In addition to distance-based outlier approach, the other approaches to outlier detection can be classified into five categories, which are distribution-based approach, depth-based approach, clustering approach, density-based approach and RST-based approach (Kovacs et al., 2004). Distribution-based approach is the classical method in statistics. It is based on some standard distribution model (Normal, Poisson, etc.) and those objects which deviate from the model are recognized as outliers (Rousseeuw & Leroy, 1987). Its greatest disadvantage is that the distribution of the measurement data is unknown in practice. Depth-based approach is based on computational geometry and compute different layers of k–d convex hulls and flags objects in the outer layer as outliers (Johnson et al., 1998). However, it is a well-known fact that the algorithms employed suffer from the dimensionality curse and cannot cope with large k. Clustering approach classifies the input data. It detects outliers as by-products (Jain et al., 1999). Since the main objective is clustering, it is not optimized for outlier detection. Density-based approach was originally proposed by Breunig et al. (2000). A local outlier factor (LOF) is assigned to each sample based on their local neighborhood density. Samples with high LOF value are identified as outliers. The disadvantage of this solution is that it is very sensitive to parameters defining the neighborhood. Rough set theory (RST) is proposed by Pawlak

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A distance metric is a distance function on a set of points, mapping pairs of points into the non-negative real numbers. In general, there are three metric functions that are widely used. Consider that \( x_1 \) and \( x_2 \) are two objects in \( n \)-dimensional space \( A = a_1, a_2, \ldots, a_n \), \( f(x,a_i) \) denotes the value of sample \( x \) in the \( i \)-th dimension \( a_i \), then a general metric, named Minkowsky distance, is defined as

\[
D_p(x,y) = \left( \sum_{i=1}^{n} |f(x,a_i) - f(y,a_i)|^p \right)^{1/p}
\]

where (1) it is called Manhattan distance \( D_1 \) if \( p = 1 \); (2) Euclidean distance \( D_2 \), if \( p = 2 \); (3) Chebychev distance \( D_{\infty} \), if \( p = \infty \).

**Example 1.** Given an information system \( IS = (U,A,V,f) \), where \( U = x_1, x_2, x_3, x_4, x_5, A = \{a\} \), as shown in Table 1.

Using the Manhattan distance, supposed \( q = 0.1 \), we can have the following neighborhoods for objects of \( U \):

\[
\begin{align*}
D_1(x_1, x_2) &= 0.2, \\
D_1(x_1, x_3) &= 0.1, \\
D_1(x_1, x_4) &= 0.3, \\
D_1(x_1, x_5) &= 0.7,
\end{align*}
\]

\[
\begin{align*}
D_2(x_1, x_2) &= 0.2, \\
D_2(x_1, x_3) &= 0.1, \\
D_2(x_1, x_4) &= 0.3, \\
D_2(x_1, x_5) &= 0.7,
\end{align*}
\]

\[
\begin{align*}
D_{\infty}(x_1, x_2) &= 0.2, \\
D_{\infty}(x_1, x_3) &= 0.1, \\
D_{\infty}(x_1, x_4) &= 0.3, \\
D_{\infty}(x_1, x_5) &= 0.7.
\end{align*}
\]

**Table 1.** An example of neighborhoods.

<table>
<thead>
<tr>
<th>( U )</th>
<th>( A )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( x_1 )</td>
<td>0.1</td>
</tr>
<tr>
<td>( x_2 )</td>
<td>0.2</td>
</tr>
<tr>
<td>( x_3 )</td>
<td>0.3</td>
</tr>
<tr>
<td>( x_4 )</td>
<td>0.7</td>
</tr>
<tr>
<td>( x_5 )</td>
<td>0.8</td>
</tr>
</tbody>
</table>

Obviously, neighborhood relations are one class of similarity relations, which satisfy reflexivity and symmetry. Specially, \( n(x) \) is an equivalent class and \( R \) is an equivalence relation if \( q = 0 \), this case is applicable to discrete data. Neighborhood relations draw the objects together for similarity or indistinguishability in terms of distances.

3. Neighborhood-based outlier detection

3.1. The value difference metric under the neighborhood relation

The value difference metric (VDM) was introduced by Stanfill and Waltz (1986) to provide an appropriate distance function for nominal attributes. A simplified version (without the weighting schemes) of the VDM is defined as:

\[
VDM(x,y) = \sum_{f \in F} d_f(x_f, y_f)
\]

where \( F \) is the set of all features in the problem domain, \( x \) and \( y \) are any two objects between which we shall calculate the distance and \( d_f(x_f, y_f) \) denotes the distance between two values \( x_f \) and \( y_f \) of feature \( f \), where \( x_f \) is the value of object \( x \) on feature \( f \) and \( y_f \) is the value of object \( y \) on feature \( f \).

For any feature \( f \in F \), \( d_f(x_f, y_f) \) is defined as follows:

\[
d_f(x_f, y_f) = (P(x_f) - P(y_f))^2
\]

where \( P(x_f) \) is the probability of object \( x \) on feature \( f \) and \( P(x_f) \) is the probability of object \( y \) on feature \( f \).
Since traditional rough set theory is suitable to discrete data, it deals with not only discrete data but also continuous data if we introduce neighborhood relation to rough set. Next we give the revised definition of VDM in rough set theory under the neighborhood relation.

**Definition 1.** Given an information system $IS = (U, A, V, f)$, where $U$ is a non-empty finite set of objects and $A$ is non-empty finite set of attributes. Let $x, y \in U$ be any two objects between which we shall calculate the distance. The value difference metric in rough set theory under the neighborhood relation $VDM_{B}: U \times U \rightarrow [0,\infty)$ is defined as $VDM_{B}(x, y) = \frac{|n_{B}^{1}(x) - n_{B}^{0}(y)|}{|U|}$ where $n_{B}(x, y)$ denotes the distance between two objects on attribute $a$, and $n_{B}$ is the value of object $x$ on attribute $a$. For any $a \in A$, let $n_{a}$ is a neighborhood parameter, define

$$d_{a}(x_{a}, y_{a}) = \left( \frac{|n_{B}^{1}(x) - n_{B}^{0}(y)|}{|U|} \right)^{2}$$

where $n_{B}^{1}(x)$ is a neighborhood of object $x$ on attribute $a$ and $n_{B}^{0}(y)$ is a neighborhood of object $y$ on attribute $a$.

If values on attribute $a$ are discrete, we set the neighborhood parameter $n_{a} = 0$, otherwise, set $q \in (0, \infty)$. Obviously, $d_{a}(x, y)$ is similar to $P(x|y)$ in the above definition.

**Example 2.** Given an information system $IS = (U, A, V, f)$, where $U = \{x_{1}, x_{2}, x_{3}, x_{4}, x_{5}\}$, $A = \{a, b, c\}$, as shown in Table 2.

The second column and the fourth column are continuous data. The third column is discrete data. Let $q_{a} = 0.1$, $q_{b} = 0$, $q_{c} = 0.1$. Using the distance metric defined by Definition 1, we can calculate the distance for every pair of objects in $U$. Because of the limitation of space, we just present the procedure for calculating the distance between $x_{1}$ and $y_{2}$.

**Initialization:**

$VDM_{B}(x_{1}, x_{2}) = d_{a}(0.1, 0.2) + d_{b}(1.0) + d_{c}(0.3, 0.4)$

**Step 1:** Calculate $d_{a}(0.1, 0.2)$:

$$n_{a}^{1}(x_{1}) = \{x_{1}, x_{2}\}$$

$$d_{a}(0.1, 0.2) = \left( \frac{|n_{a}^{1}(x_{1}) - n_{a}^{0}(x_{2})|}{|U|} \right)^{2}$$

$$= \left( \frac{|x_{1} - x_{2}|}{|U|} \right)^{2}$$

$$= \left( \frac{2}{5} \cdot \frac{3}{5} \right)^{2} = \frac{1}{25}$$

**Step 2:** Calculate $d_{b}(1.0)$:

$$n_{b}^{1}(x_{1}) = \{x_{1}\}$$

$$d_{b}(1.0) = \left( \frac{|n_{b}^{1}(x_{1}) - n_{b}^{0}(x_{2})|}{|U|} \right)^{2}$$

$$= \left( \frac{|x_{1} - x_{2}|}{|U|} \right)^{2}$$

$$= \left( \frac{1}{5} \cdot \frac{3}{5} \right)^{2} = \frac{4}{25}$$

**Step 3:** Calculate $d_{c}(0.3, 0.4)$:

$$n_{c}^{1}(x_{1}) = \{x_{1}, x_{2}\}$$

$$d_{c}(0.3, 0.4) = \left( \frac{|n_{c}^{1}(x_{1}) - n_{c}^{0}(x_{2})|}{|U|} \right)^{2}$$

$$= \left( \frac{|x_{1} - x_{2}|}{|U|} \right)^{2}$$

$$= \left( \frac{2}{5} \cdot \frac{3}{5} \right)^{2} = \frac{1}{25}$$

Repeating the above calculation, we can finally obtain distances for all the other pairs of objects in $U$. By then, we define a neighborhood-based object outlier factor (NOOF), which indicates the degree of outlier for every object in an information system.

**Definition 2 (Neighborhood-based Object Outlier Factor).** Let $IS = (U, A, V, f)$ be an information system, where $A = \{a_{1}, a_{2}, \ldots, a_{m}\}$ and $U = \{x_{1}, x_{2}, \ldots, x_{n}\}$. For any $x_{i} \in U$, let neighborhood parameter $q = \{q_{a_{1}}, q_{a_{2}}, \ldots, q_{a_{m}}\}$. The neighborhood-based object outlier factor of $x_{i}$ in $IS$ is defined as follows:

$$NOOF_{i}(x_{i}) = \sum_{j=1}^{n} VDM_{B}(x_{i}, x_{j})$$

**Step 4:**

$$VDM_{B}(x_{i}, x_{j}) = \frac{1}{25} \cdot \frac{4}{25} \cdot \frac{1}{25} = 0.24$$

**3.2. Algorithm**

**Neighborhood outlier detection (NED)**

**Input:** an information system $IS = (U, A, V, f)$, where $|U| = n$ and $|A| = m$; neighborhood parameter $q = q_{a_{1}}, q_{a_{2}}, \ldots, q_{a_{m}}$; threshold value $\mu$.

**Output:** a set $O$ of neighborhood-based outliers.

1. For every $a \in A$
2. {  
3. For every $x \in U$
4. {  
5. Calculate $n_{B}^{a}(x)$;  
6. }
7. }
8. For every $x \in U$
9. {  
10. For every $y \in U$
11. {  
12. For every $a \in A$
13. {  
14. Calculate $d_{a}(x_{a}, y_{a}) = \frac{|n_{B}^{a}(x_{a}) - n_{B}^{a}(y_{a})|}{|U|}$;  
15. }
16. Calculate $VDM_{B}(x, y)$  
17. }
18. Calculate $NOOF(x)$
19. If $NOOF(x) > \mu$, then $O = O \cup x$
20. }

<table>
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<th>a</th>
<th>b</th>
<th>c</th>
</tr>
</thead>
<tbody>
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<td>0.3</td>
</tr>
<tr>
<td>x_{2}</td>
<td>0.2</td>
<td>0</td>
<td>0.4</td>
</tr>
<tr>
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<td>0.6</td>
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<td>0</td>
<td>0.7</td>
</tr>
<tr>
<td>x_{5}</td>
<td>0.8</td>
<td>0</td>
<td>0.5</td>
</tr>
</tbody>
</table>
4. Experimental analysis

In this section, following the experimental setup in He, Deng, and Xu (2005), we shall use two real life data sets (Annealing and Cancer) to demonstrate the performance of neighborhood-based outlier detection algorithm (NED) against traditional distance-based method (Knorr & Ng, 1998, Knorr, Ng, & Tucakov, 2000) and KNN algorithm (Ramaswamy et al., 2000). Furthermore, in our experiment, for the KNN algorithm, the results were obtained by using the fourth nearest neighbor (Ramaswamy et al., 2000) and the overlap metric in rough set theory defined above.

4.1. Annealing data

The first is the Annealing data set, which can be found in the UCI machine learning repository (Bay, 1999). It contains 798 instances (or objects) with 38 attributes (including the class attribute). The 798 instances are partitioned into five classes. Class 3 has the largest number of instances. The remained classes are regarded as rare classes for they are small in size.

Aggarwal and Yu (2001) proposed a practicable way to test the effectiveness of an outlier detection method (Angiulli & Pizzuti, 2002; He et al., 2005). That is, we can run the outlier detection method on a given data set and test the percentage of points which belonged to one of the rare classes (Aggarwal considered those kinds of class labels which occurred in less than 5% of the data set as rare labels (Angiulli & Pizzuti, 2002)). Points belonged to the rare class are considered as outliers. If the method works well, we expect that such abnormal classes would be over-represented in the set of points found.

In our experiment, data in the Annealing data set is input into an information table \(SL = (U; A; V; f)\), where \(U\) contains all the 798 instances of Annealing data set and \(A\) contains 37 attributes of Annealing data set (not including the class attribute). Since the neighborhood parameters are needed by NED, the corresponding parameters are illustrated in Table 3. These parameters were determined based on a small number of preliminary runs. The experimental results are summarized in Table 4.

Table 4 shows the results produced by the NED algorithm against the KNN algorithm and DIS algorithm. Here, the top ratio is ratio of the number of objects specified as top-\(k\) outliers to that of the objects in the dataset. The coverage is ratio of the number of detected rare classes to that of the rare classes in the dataset. For example, we let NED algorithm find the top 80 outliers with the top ratio of 10%. By examining these 80 points, we found that 51 of them belonged to the rare classes. In contrast, when we ran the KNN algorithm on this dataset, we found that only 21 of 80 top outliers belonged to rare classes.

From Table 4, the performance of the NED algorithm outperformed that of the other two methods. This is especially the case when the top ratio is relative small, the NED algorithm worked much better. Anneal dataset has not only discrete data, but also continuous data. The experiment shows that the NED algorithm is suitable to mixed data.

Table 5 lists the results produced by the NED algorithm against the KNN algorithm and DIS algorithm on the Wisconsin breast cancer data set. The Wisconsin breast cancer data set consists of 699 instances (or objects) with 9 attributes (including the class attribute). The 699 instances are partitioned into 5 classes.

In the worst case, the time complexity of algorithm NED is \(O(m \times n^2)\), and its space complexity is \(O(m \times n)\), where \(m\) and \(n\) are the cardinalities of \(A\) and \(U\) respectively.
4.2. Wisconsin breast cancer data

The Wisconsin breast cancer dataset is found in the UCI machine learning repository (Bay, 1999). The data set contains 699 instances with 9 attributes. Here we follow the experimental technique of Harkins et al. by removing some of the malignant instances to form a very unbalanced distribution (Angiulli & Pizzuti, 2002). The resultant dataset had 39 (8%) malignant instances and 444 (92%) benign instances.

Data in the Wisconsin breast cancer data set is also input into an information table \( SW = (U; A; V; f) \), where \( U \) contains all the 483 instances of the data set and \( A \) contains nine attributes of the data set (not including the class attribute). We consider detecting outliers (malignant instances) in SW. Since the dataset is discrete, the parameters for NED are set to 0. The experimental results are summarized in Table 5.

Table 5 is similar to Table 4. From Table 5, we can see that for the Wisconsin breast cancer dataset, the NED performs better than RNN method, and a litter weaker than KNN and DIS. In fact, the performance of NED is more suitable to continue dataset than discrete dataset.

5. Conclusion and future work

Outlier detection is becoming critically important in many areas. In order to deal with not only discrete data but also continue data set, we proposed a new definition of the traditional distance metrics by considering neighborhood information. A measure for identifying the significance of an outlier is also presented. Furthermore, we give the neighborhood-based algorithm for discovering outliers. The experimental results show that our approach outperformed existing methods on identifying meaningful and interesting outliers for mixed dataset.

In the future work, for the neighborhood-based outlier detection algorithm, we shall consider using rough set feature select method to reduce the features while preserving the performance of it. For the performance of the computation of our method, we will sort all objects according to a given order on values of feature to improve the computational complexity.

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References


