Example-based Outlier Detection with Relevance Feedback

Cui ZHU\(^\star\) Hiroyuki Kitagawa\(^\star\)
Spiros Papatdimitriou\(^\star\)
Christos Faloutsos\(^\star\)

Detecting outliers is an important, but tricky problem, since the preference of outlier-ness often depends on the user and/or the dataset. Existing techniques traditionally detect outliers based on some prescribed definitions of outlier. However, it is very difficult for a user to decide the definition of outliers in prior. We have developed a system of detecting outliers that match users' intentions implied by outlier examples in prior work. In this paper, we propose a new refined method of interactive outlier detection adaptive to users' intentions. Our experiments on both synthetic and real datasets show that the new interactive method surpass the former method in detecting outliers that match the users' intentions.

1. Introduction

In applications like fraud detection, medical analysis, etc, outlier detection is an important problem, since the rare events or exceptional cases are more interesting and useful than the common cases.

In the context of outlier detection, what makes the problem more difficult is that not everyone has the same idea of what constitutes an outlier. Intuitively, an object is an "outlier" or "abnormal" if it is in some way "significantly different" from its "neighbors". Different answers to what constitutes a "neighborhood", how to determine "difference" and whether it is "significant" would lead to various sets of objects defined as outliers.

In most circumstances, users are experts in their problem domain and not in outlier detection. Usually, they have a few outlier examples in hand, which may "describe" their intentions and want to find more objects that exhibit "outlier-ness" characteristics just like those examples.

We have presented in our previous work [13, 14] a novel method that detects outliers adaptive to users' intentions implied by the outlier examples. In the method, users are intended to provide some examples and also the fraction of outliers they want to be detected.

However, sometimes the fraction of outliers is hard to figure out in prior. To address the problem, we improve the system by interactive detection of outliers eliminating the need of the outlier fraction.

The remainder of the paper is organized as follows: In section 2, we discuss related work on outlier detection. In section 3, we introduce briefly the method of outlier detection based on examples. Section 4 presents the improved system. Section 5 reports the experimental evaluation on both synthetic and real dataset. Finally, Section 6 concludes the paper.

2. Related Work

In essence, outlier detection techniques traditionally employ unsupervised learning processes. The several existing approaches can be broadly classified into the following categories: (1) Distribution-based approach, (2) Depth-based approach, (3) Clustering approach, (4) Distance-based approach, (5) Density-based approach. All of the above approaches regard being an outlier as a binary property. They do not take into account both the degree of "outlier-ness" and where the "outlier-ness" is presented. (6) LOF, [9]. They introduced a local outlier factor (LOF) for each object, indicating its degree of "outlier-ness." We proposed the multi-granularity deviation factor (MDEF) in [10]. MDEF measures the "outlier-ness" of objects in neighborhoods of different scales. LOCI examines the MDEF values of objects in all ranges. Even though the definition of LOF and MDEF can capture "outlier-ness" in different scales, these difference of scales were not taken into account by the system.

In order to deal with the curse of high dimensionality, a quite different technique is proposed in [2], where outliers are found by studying the behavior of projections from the data set. The most sparse low-dimensional cubes in the data are found by GA algorithm, and all the objects in these cubes are reported as outliers.

Another outlier detection method was developed in [8], which focuses on the discovery of rules that characterize outliers, for the purposes of filtering new points later. This is a largely orthogonal problem. Outlier detection by SmartFilter are used to create labeled data, which are then used to find the outlier filtering rules.

In summary, none of the existing methods can directly incorporate user in the discovery process. We have proposed to detect outliers using user-provided examples [13, 14]. In this paper, we improve the method by interactive detection of outliers.

3. Outlier Detection by Examples

3.1 Measuring Outlier-ness

In order to understand the users' intentions and the "outlier-ness" they are interested in, a first, necessary step is measuring the "outlier-ness." We employ the multi-granularity deviation factor (MDEF) [10] for this purpose, which is capable of measuring "outlier-ness" of objects in the neighborhoods of different scales (i.e., radii).

Here we describe some basic terms and notation. Let the \( r \)-neighborhood of an object \( p \) be the set of objects within distance \( r \) of \( p \). Let \( n(p, r) \) and \( n(p, r) \) be the numbers of objects in the \( ar \)-neighborhood (counting neighborhood) and \( r \)-neighborhood (sampling neighborhood) of \( p \), respectively. Let \( \hat{n}(p, r) \) be the average of \( n(p, r) \), over all objects \( p \) in the \( r \)-neighborhood of \( p \).

\(^1\) In all experiments, \( \alpha = 0.5 \) as in [10].
Definition (MDEF). For any $p_i$, $r$ and $\alpha$, the multi-granularity deviation factor (MDEF) at radius (or scale) $r$ is defined as follows:

$$MDEF(p_i, r, \alpha) = \frac{n(p_i, r, \alpha) - m(p_i, r, \alpha)}{\hat{n}(p_i,\alpha, r)}$$ (1)

Intuitively, the MDEF at radius $r$ for a point $p_i$ is the relative deviation of its local neighborhood density from the average local neighborhood density in its $r$-neighborhood. Thus, an object whose neighborhood density matches the average local neighborhood density will have an MDEF of 0. In contrast, outliers will have MDEFs far from 0. And outliers with dissimilar density distribution will exhibit "outlier-ness" at different scales, or have different peaks of MDEF plots. In our paper, the MDEF values are examined (or, sampled) at a wide range of sampling radii $r, r_{\text{min}} \leq r \leq r_{\text{max}}$ [13, 14].

3.2 Detecting Outliers by Examples

Below we give a concise overview of the method according to [14]. The method detects outliers based on user-provided examples and a user-specified fraction of objects to be detected as outliers in the dataset. The method performs in two stages: feature extraction step and classification step.

Feature Extraction Step The purpose of this step is to map all objects into the MDEF-based feature space, where the MDEF plots of objects capturing the degree of "outlier-ness," as well as the scales at which the "outlier-ness" appears, are represented by vectors. Let $D$ be the set of objects in the feature space. In this space, each object is represented by a vector: $O_i = (m_{r_1}, m_{r_2}, \ldots, m_{r_m}), O_i \in D$, where $m_j = MDEF(p_i, r_j, \alpha)$, $0 \leq j \leq n$, $r_0 = r_{\text{min}}$, $r_n = r_{\text{max}}$, $r_j = \frac{r_{\text{max}} - r_{\text{min}}}{n}$.

Classification Step We use an SVM (Support Vector Machine) classifier to learn the "outlier-ness" of interest to the user and then detect outliers which match this. Concretely, the algorithm uses the marginal property of SVMs to learn from the examples and the unlabeled data (i.e., the rest of the objects in the dataset).

The classification step consists of the following five sub-steps. (1) Negative training data extraction sub-step. All objects are sorted in descending order of $\max_j(m_j)$. Then, from the objects at the bottom of the list, we select a number of (strong) negative training data equal to the number of examples. Let the set of strong negative training data be $\text{NEG}$. Also, let the set of examples be $\text{POS}$. (2) Training sub-step. Train a SVM classifier using $\text{POS}$ and $\text{NEG}$. (3) Testing sub-step. Use the SVM to divide the dataset into the positive set $\text{P}$ and negative set $\text{N}$. (4) Update sub-step. Replace $\text{NEG}$ with $\text{N}$, the negative data obtained in the testing sub-step. (5) Iteration sub-step. Iterate from the training sub-step to the updating sub-step until the ratio of the objects in $\text{P}$ converges to the fraction specified by the user. The objects in the final $\text{P}$ are reported to the user as detected outliers.

4. Proposed Method

The previously proposed method has the ability to detect outliers adaptive to users' intentions, given outlier examples. Aside from these examples, the algorithm uses as input the fraction of outliers to be determined. The fraction is used to determine the termination condition of iterations in the classification step.

In some cases, the fraction of outliers is hard to figure out in prior. Here, we further develop the method to eliminate the need to input outlier fraction.

4.1 Modification of the Former Method

Without the information of fraction of outliers, the former termination condition of iterations in classification step should be modified. Now we do iterations from the training sub-step to the update sub-step until the set of classified outliers $\text{P}$ is same as $\text{P'}$. $\text{P'}$ is the set of outliers in the previous iteration.

Accordingly, at the end of iteration, the hyperplane for separating outliers and normal objects is set as close as possible to the set of given examples. Therefore, the detected objects are those displaying stronger "outlier-ness" characteristics than the examples.

4.2 Proposed Method

In order to discover more outliers adaptive to users' intentions, we integrate the aforementioned modified detection method with a relevance feedback technique which enables the user to refine their preference by specifying relevant and non-relevant outliers (or normal objects). Based on users' feedback information, the system attempts to "guess" what are his intentions of outliers, and how strong the "outlier-ness" they want.

The framework of interactive detection is as follows:

Classification Step Use the abovementioned classification method to set the hyperplane for separating outliers and normal objects as close as possible to the set of given examples.

Feedback Step In order to provide valuable feedback, i.e., raise questions of promising user outliers, we select randomly objects within the negative margin of the hyperplane. Even though, these objects have been classified as "normal" by the SVM in the classification step, they still exhibit "outlier-ness" characteristics close to the examples. The MDEF plots of the feedback objects are also presented to help users to determine whether the objects are interesting outliers. The "yes" answers, or positive feedbacks, are new outlier examples, and are incorporated into the positive training data.

Convergence Step Iterate the classification and feedback steps until all the answers to feedbacks are "no". The hyperplane constructed at the end of the process will be close to the desired hyperplane which divides all the interesting outliers from normal objects. The objects separated by the last hyperplane are reported to the user as all detected outliers.

Figure 1 shows the flow of the proposed outlier detection method.

5. Experiments

In this section, we describe our experimental methodology and the results on both synthetic and real data.

5.1 Experimental procedure

Our experimental procedure is as follows:
To simulate interesting outliers, we start by selecting objects which represent “outlier-ness” at some scales under the conditions of the form \( \land_j(m_{i,j}, \max_{i,j}, \text{Cond}_{i,j}, K_j) \), where \( m_{i,j}, \max_{i,j}, \text{Cond}_{i,j}, K_j \) stands for the condition that \( m_{i,j} \leq j \leq \max_{i,j} \) and \( \text{Cond}_{i,j} \) is either “>” or “<”.

2. Then, we randomly sample 8% of the outliers to serve as examples that would be picked by a user, and “hide” the remainders.

3. Next, we detect outliers using the proposed method.

4. Finally, we compare the detected outliers with the (known) simulated set of interesting outliers. Precision (P), recall (R), and F1-measure (F1) defined below are adopted to measure the performance. F1-measure is a trade-off of precision and recall. A larger value of F1-measure indicates better performance.

\[
P = \frac{\# \text{ of correct positive predictions}}{\# \text{ of positive predictions}}
\]

\[
R = \frac{\# \text{ of correct positive predictions}}{\# \text{ of positive data}}
\]

\[
F1 = \frac{2 \times R \times P}{(R + P)}
\]

We use the LIBSVM [7] implementation for our SVM classifier. In all experiments, we use polynomial kernels and the same SVM parameters. Therefore, the whole processes can be done automatically.

4In all experiments, \( y = 10 \).

3For the parameter C (the penalty imposed on training data that fall on the wrong side of the decision boundary), we use 1000, i.e., a high penalty to mis-classification. For the polynomial kernel, we employ a kernel function of \((w' \cdot v + 1)^2\).

For feedback, each time we select randomly \( z \) objects among the negative margin as questions. 4 “Yes” answers are given to the objects which are among the simulated interesting outliers. “No” answers are given to those not in the set of interesting outliers. We terminate the convergence step when the number of “yes” answers is zero.

### 5.2 Datasets and Results

We do experiments on both synthetic and real dataset to evaluate the proposed method. Table 1 shows the descriptions of all datasets.

Table 2 shows all the sets of interesting outliers along with the corresponding discriminants used as the underlying outlier concept in our experiments. In the table, for instance, the discriminant (1, 40, >, 0.9) means that objects are selected as interesting outliers when their MDEF values are greater than 0.9 in the range of radii from 1 to 40. The number of interesting outliers is also shown in Table 2. We always randomly sample 10% (\( r = 10 \)) of the interesting outliers to serve as user-provided examples and “hide” the rest.

The results are also shown in Table 2. The former method is the method that detects outliers with inputs of fractions of outliers, and the improved method refers to interactive detection of outliers without fraction factors inputted. To compare performance of the two methods, we always use the same sets of user examples. And the results in Table 2 are average of five trials. In all cases, with a little sacrifice of recall measurement, the interactive method surpassed the former method in detecting interesting outliers so far as precision measurement is concerned. F1-measure metrics exceed those of the former method. Note also that the number of feedbacks is always small (less than 10).

### 6. Conclusion

The fact that the preference of outlier-ness often depends on the user and/or the dataset makes the problem of outlier detection difficult to resolve. We propose to solve this problem by allowing the user to give us some outlier examples and acting interactively with the system. Experiments on both real and synthetic data demonstrate that the interactive method can successfully incorporate these examples including positive examples in the feedback and detect outliers with “outlier-ness” characteristics like the given examples.

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Table 2: Interesting Outliers, Discriminants and the Performance of the Former and Improved Methods. Note that we always use the same set of user examples to compare the two methods. IO denotes interesting outliers. Precision (P), recall (R) and F1-measure (F1) are used to show the performance. The number of iterations (IT) for convergence in the improved method is also shown.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Cases</th>
<th>IO</th>
<th>Former Method</th>
<th>Improved Method</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>P</td>
<td>R</td>
</tr>
<tr>
<td>Ellipse Dataset</td>
<td>E-Fringe Fringe (5, 30, &gt;) 0.85)</td>
<td>214</td>
<td>84.02</td>
<td>89.77</td>
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<td></td>
<td>E-Long Long Ends (15, 25, &gt;) 0.8)</td>
<td>137</td>
<td>95.97</td>
<td>97.30</td>
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<tr>
<td></td>
<td>E-Short Short Ends (5, 15, &gt;) 0.8)</td>
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<td>83.26</td>
<td>89.94</td>
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<tr>
<td>Mixture Dataset</td>
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<td>86.51</td>
<td>93.09</td>
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<tr>
<td></td>
<td>M-Large Large Cluster (15, 40, &gt;) 0.9)</td>
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<td>89.13</td>
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<td></td>
<td>M-Small Small Cluster (1, 10, &gt;) 0.9)</td>
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<tr>
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<td>M-Origin Origin Part (0, 40, &gt;) 0.94)</td>
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<td>62.70</td>
<td>77.33</td>
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References: