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Abstract. Unsupervised anomaly detection is commonly performed using a distance or density based technique, such as K-Nearest neighbours, Local Outlier Factor or One-class Support Vector Machines. One-class Support Vector Machines reduce the computational cost of testing new data by providing sparse solutions. However, all these techniques have relatively high computational requirements for training. Moreover, identifying anomalies based solely on density or distance is not sufficient when both point (isolated) and cluster anomalies exist in an unlabelled training set. Finally, these unsupervised anomaly detection techniques are not readily adapted for active learning, where the training algorithm should identify examples for which labelling would make a significant impact on the accuracy of the learned model. In this paper, we propose a novel technique called Maximin-based Anomaly Detection that addresses these challenges by selecting a representative subset of data in combination with a kernel-based model construction. We show that the proposed technique (a) provides a statistically significant improvement in the accuracy as well as the computation time required for training and testing compared to several benchmark unsupervised anomaly detection techniques, and (b) effectively uses active learning with a limited budget.

Keywords: Anomaly detection \cdot Unsupervised learning \cdot Active learning.

1 Introduction

Anomaly detection is a key component of many monitoring applications, which aim to detect harmful rare events that can be subsequently controlled [8]. It has been used in a wide range of domains from cybersecurity [33, 7] to health and safety applications such as fall detection for elderly people [27, 35]. A key challenge for anomaly detection is the abundance of unlabelled data [23]. The high cost of labelling hinders the application of supervised anomaly detection techniques, which require labelled examples of anomalies and normal data [8]. Although one-class classification techniques mitigate this issue by building a normal profile given only normal data, they are not sufficiently robust to the

presence of unknown anomalies in the training set [32, 3, 16]. Even if the training set only comprises normal data but is noisy, one-class classification can deliver unsatisfactory results [16, 15]. Since one-class classification techniques such as One-Class Support Vector Machine (OCSVM) [30] and Support Vector Data Description (SVDD) [32] provide sparse solutions and are very fast during the testing phase, they have been enhanced to work in an unsupervised manner [3, 16, 15]. However, depending on the implementation and the characteristics of the dataset, training may require $O(n^2)$ to $O(n^3)$ operations, where n is the cardinality of the training data. Unsupervised anomaly detection techniques such as K-Nearest Neighbours (KNN) and Local Outlier Factor (LOF), have high computational requirements for processing new observations in a continuously monitored system. For scroring/labelling a new data point, anomaly scores of all or a subset of existing data points should be recomputed in a fairly large reference dataset. Therefore, these methods have limited scope for real-time anomaly detection. In other words, they do not learn an explicit model a priori, which can be later on used for timely evaluation of future observations [1]. iForest[25] is another unsupervised method that attempts to address these challenges by filtering anomalies through isolating trees that are trained on several subsets of the data. This way, iForest is not based on a density or distance measure and lowers the computational cost by sampling. However, the solution provided by iForest is not sparse like OCSVM and SVDD, and to score a test instance it must scan several trees.

In some applications, it might be possible to obtain expert feedback on whether an instance is normal or anomalous. Having that feedback on a small number of critical examples can make a substantial difference to the accuracy of the final model [23]. This process, known as Active Learning (AL), has been widely used in classification [34] and rare class discovery [20, 17] using supervised or semi-supervised learning. Using AL in unsupervised anomaly detection is an emerging trend [19, 1]. Sharma et al. [31] used active learning to train a two-class classifier for identifying operationally significant anomalies from insignificant ones in a flight trajectory dataset. They used the OCSVM algorithm first to identify top-ranked anomalies in an unsupervised manner. Given their scores, the top-ranked anomalies are then presented to an expert to generate a labelled set of operationally significant and insignificant anomalies. This labelled set is used to train a two-class SVM that distinguishes between interesting and unimportant anomalies. Pelleg and Moore [26] proposed a general framework for the same purpose that runs several loops for data modelling (using Gaussian mixtures) and labelling. The algorithm starts with an unlabelled dataset. After each modelling round, labels of 35 instances are asked from an expert. Three strategies are used to choose these instances: choosing instances with low likelihood under the current model, choosing instances that the model is not certain about them, and a combination of the two strategies. Our work is different as we aim to enhance the ability of the underlying unsupervised anomaly detection, which is used by these techniques to find interesting anomalies or discover rare classes. The ability of an anomaly detection technique to efficiently and effec-

tively use the AL budget is vital for adversarial tasks such as fraud detection, where anomalies can be similar to the normal data due to fraudsters mimicking normal behaviour [1].

In this paper, we present a novel approach called Maximin-based Anomaly Detection (MMAD). The contributions of this work are as follows. First, we use random sampling followed by the Maximin (MM) sampling technique [22] to select a representative subset of the input data, which achieves low constant computational cost for big datasets. Then, we use a cluster validity index (CVI) called the Silhouette index [29] on the representative samples that should contribute to defining a kernel-based model, which is subsequently used to score anomalies and normal data in nearly real-time. Second, we incorporate AL into MMAD. We show that with only a few labels, this enhancement of MMAD improves the accuracy of unsupervised anomaly detection. Our numerical experiments on benchmark anomaly detection datasets show that our proposed technique outperforms several state-of-the-art unsupervised anomaly detection techniques in terms of the time-complexity of training a model and testing the future data. Moreover, our technique provides statistically significant improvements in accuracy even when the AL budget is zero.

2 Definitions and Problem Specification

Let the normal data D^* be (an unknown) subset of a given unlabelled training set D, i.e., $D^* \subseteq \mathcal{D} = \{x_1, x_2, ..., x_n\} \subset \mathbb{R}^d$, drawn from an unknown probability distribution P on \mathbb{R}^d . The probability distribution P can be approximated by estimating the parameter values θ of P such that:

$$\theta = \underset{\theta \in \Theta}{\operatorname{arg\,min}} \sum_{x} [(1 - P(x;\theta)) \times I(x \in D^*) + P(x;\theta) \times I(x \notin D^*)], \quad (1)$$

where Θ represents the set of parameter values for the probability distribution, and I(.) is the indicator function. The cardinality and mean value of a set Dare respectively shown by |D| and \overline{D} , and for the training set |D| = n. In the unsupervised case, estimating θ is done without ground truth or a priori knowledge about the data. The estimated probability distribution P can be used to score data instances such that anomalies get low scores.

We assume that a limited budget B for AL is available, i.e., labels of B instances can be queried from an oracle. However, the training data might not be available after the query. Thus, the model should be updated after gaining a label from the oracle without having to build the model from scratch.

3 Methodology

An accurate description of a parametric probability distribution $P(x; \theta)$ for real datasets is usually unavailable. Therefore, instead of solving (1), MMAD esti-



Fig. 1. Demonstration of the three stages of the MMAD Algorithm on the toy example X_{toy} . A candidate set of representative samples CRS is chosen in Stage 1. By removing anomalies from CRS, the final set of representative samples RS is generated in Stage 2. A kernel-based model is learned in Stage 3.

mates a non-parametric probability density of the form:

$$P(x; \{w_{1..n}\}) = \sum_{i=1}^{n} w_i k(x, x_i), \text{ subject to } w_i \ge 0, \sum_{i=1}^{n} w_i = 1,$$
(2)

where k represents an arbitrary kernel function and w is the weight vector of the kernel. Our aim is to find a subset $\mathcal{X} \subseteq D$ of \mathbb{R}^d that represents the normal data D^* and divides \mathbb{R}^d into two subsets \mathcal{X} and $\mathbb{R}^d - \mathcal{X}$. Substituting θ with $\{w_{1..n}\}$ in (1), $P(x; \{w_{1..n}\})$ is viewed as a scoring function that assigns high scores to normal data and low scores to anomalies. The computational cost of testing an instance using $P(x; \{w_{1..n}\})$ scales with the number of non-zero w_i (i = 1..n), thus a sparse solution that minimises $\sum_{i=1}^n w_i$ is desirable.

To learn $P(x; \{w_{1..n}\})$, MMAD assigns the values of each w_i in three stages as shown in Figure 1 for a toy example X_{toy} , and explained in Algorithm 1. Figure 1(a) draws X_{toy} . The first stage includes the following steps. A subset of X_{toy} is selected at random as set S in Figure 1(b). MM sampling [22] is performed on S to generate a uniformly selected subsample from S shown in Figure 1(c). Choosing a large value for |S| and running this extra MM sampling step is necessary, because relying only on random sampling to secure a representative subset of the input data is not sufficient for two reasons. First, it does not guarantee the selection of representative samples from all clusters in the data, especially if there are very small clusters in the data. Second, random sampling does not provide a mechanism to eliminate anomalies from the model, especially when a high fraction of anomalies exists in the training data. After MM sampling, a set called *Candidate Representative Samples* (CRS) is built using representative MM samples. By evaluating each MM sample as a CRS object, in Figure 1(d) an optimal cardinality $|CRS|^*$ for this set is defined. In this step the first two MM samples are designated as cluster centers, and a crisp 2-partition U_2 of S,

Al	gorithm 1 MMAD	
	Input: training set $D = \{x_1, x_2,, x_n\}$, sample a	size $ S $
	budget B , metric δ	
	Output: model M: RSs , W , and γ	
	Stage $1.(a)$:	
1:	S = randSample(D, S)	\triangleright random sampling from D
	Stage 1.(b):	
2:	$[I_{CRS}, \Delta] = \text{Maximin}(S, [0.4 * S], \delta)$	\triangleright call Algorithm 2
3:	for $i = 2 CRS $ do	
4:	compute $Silh_i$ using (3) given $C = \{s_{I_{CPS}^{1i}}\}$	
5:	end for	
6:	$ CRS ^* = argmax \ Silh_i$	\triangleright maximising Silhouette index
-	r $r^{1 CBS *}$	
(:	$I_{CRS} = I_{CRS}^{-1}$	
8:	for $i = 1 CRS $ do	1
0	\triangleright Substitute CRSs with i	closet point to their cluster mean
9:	$c_i = \{s\}, \text{ where } I_{CRS}^\circ = \arg\min_i \delta(s, s_{I_{CRS}^j})$	
10:	$I^{i}_{CRS} = \arg\min_{i} \delta(\overline{c_{i}}, s_{j} \in c_{i})$	
11:	end for	
	Stage 2:	
12:	identify type of the dataset	
13:	if $type == WS$ then	
14:	$I_{RS} = \text{screen}(I_{CRS})$	▷ remove anomalous CRSs
15:	set γ using (4)	
16:	else	
17:	$I_{RS} = I_{CRS}$	
18:	set γ using (5)	
19:	end if	
20:	if $B > 0$ then	
21:	$I_{RS} = BS(I_{RS}, B, \Delta_{RS})$	\triangleright call Algorithm 3
22:	end if	
	Stage 3:	
23:	$cn_j = count(c_j), j = 1 RS $	\triangleright count each cluster's members
24:	$w_j = \frac{cn_j}{\sum cn_j}, j = 1 RS $	
25:	$\mathbf{M} = (s_{I_{RS}}, \gamma, W)$	

i.e., a partition into two mutually exclusive clusters, is built using the *nearest* neighbour prototype rule (NPR). Then, the value of a CVI called the Silhouette index [29] is computed on U_2 . After finding the third MM sample and designating it as a new cluster center, U_3 is built and the value of the Silhouette index is computed on it. This procedure stops when a predefined number of MM samples are assigned as cluster centers and used to generate NPR partitions of S. Then, $|CRS|^*$ is chosen as the number of MM samples that maximises the computed Silhouette index values. The first $|CRS|^*$ MM samples form the set CRS (Figure 1(e)). Removing anomalous instances from CRS is performed in Figure 1(f) and the new points are called *Representative Samples* (*RS*). Optionally, if an AL

Algorithm 2 Maximin

 $\begin{array}{l} \textbf{Input: set } S, \text{ number of objects } |CRS|, \text{ metric } \delta\\ \textbf{Output: Indexes } I_{CRS}, \text{ pairwise dissimilarities } \Delta\\ 1: \ \Delta = [\delta(s_i, s_j)] \text{ for all } (i, j) \in 1..|S| \qquad \qquad \triangleright \text{ Cost } O(|S|^2)\\ 2: \ I_{CRS}^0 = \arg\min_{1 \leq i \leq |S|} \delta(\overline{S}, s_i) \qquad \qquad \triangleright \text{ Closet point to the mean of } S\\ 3: \ \Gamma_0 = \Delta_{I_{CRS}^0}\\ 4: \ \textbf{for } p = 1..|CRS| \ \textbf{do}\\ 5: \quad I_{CRS}^p = \arg\max_{1 \leq p \leq |S|} \Gamma_{p-1}\\ 1 \leq p \leq |S|\\ 6: \quad \Gamma_p = [min(\Delta_{I_{CRS}^p}, \Gamma_{p-1})]\\ 7: \ \textbf{end for} \end{array}$

budget is available for securing labels from an oracle, it will be spent. In this figure we assumed that no AL budget is available. Finally in Figure 1(g), the model is built using RS data. The contour plot of the scores assigned to the data instances in this figure, shows that anomalies have scores close to zero. Next, we explain each of the steps in detail.

3.1 Stage 1: CRS Selection

To reduce the time-complexity for processing big data, MMAD first selects a subset S of the training set by shuffling it and randomly selecting |S| instances. The sample S is used as an input to the MM algorithm [22], which is shown in Algorithm 2. Our implementation of MM sampling starts by finding the index of the closest point to the mean of S (line 2 of Algorithm 2). At each iteration of its main loop (lines 4 - 7 of Algorithm 2), MM adds the index of the point that has maximum dissimilarity to the points already selected. We believe that this sampling technique guarantees that the indexes I_{CRS} of the representative object set CRS have at least one member from each of the unknown clusters in the data (see Proposition 1 in [18] for a theoretical guarantee in certain but not all - cases). Towards this end, we need to define the optimal cardinality $|CRS|^*$. One way to achieve this is to evaluate the appropriateness of the cluster centers $C = \{c_{1,|CRS|}\}$ that are defined on the CRS objects. CVIs can be used for this purpose [4]. However, the presence of anomalies in a dataset can mislead CVIs. We experimented with several popular indexes including Dunn [12], C-Index [21], Calinski-Harabasz [6], Davies-Bouldin [11], and Silhouette [29], and concluded that the Silhouette index provides the best estimate of $|CRS|^*$ such that most of the point anomalies are isolated as singleton cluster centers in S.

Given cluster centers $C = \{c_{1.|C|}\}$, each sample $s_i \in S$ is assigned to the closest cluster center c_p with the NPR, i.e., $||s_i - c_p|| < ||s_i - c_q|| \quad \forall q \in 1..|C|, q \neq p$. The set of points accumulated this way for each cluster center are the |C| crisp clusters in this partition of S. Let δ denote any metric on the input space. For each cluster center c_p , the Silhouette index uses the average within-cluster dissimilarity (cohesion) $In_i^{\delta} = \frac{1}{|c_p|} \sum_{s_i, s_j \in c_p} \delta(s_i, s_j)$ of s_i to the other members of c_p , which indicates how well s_i matches the other data in c_p . The smallest average

between-cluster dissimilarity (separation) $Out_i^{\delta} = min\{\frac{1}{|c_q|}\sum_{\substack{s_i \in c_p, \\ s_j \in c_q, q \neq p}} \delta(s_i, s_j)\}$

of s_i measures how suitable would be the assignment of s_i to its closest neighbouring cluster c_q . The Silhouette index combines the cohesion (In_i^{δ}) and separation (Out_i^{δ}) measures to derive a Silhouette value for each s_i as:

$$Silh_{s_i} = \frac{Out_i^{\delta} - In_i^{\delta}}{max(In_i^{\delta}, Out_i^{\delta})},\tag{3}$$

7

which is in the range [-1, 1], where a high value indicates that s_i fits well within its own cluster. When $Silh_C = \frac{1}{|S|} \sum_{i=1}^{|S|} Silh_{s_i}$ is close to 1, the corresponding partition is preferred to those with lesser values.

Given that $|CRS|^*$ is not known a priori, we need to examine the value of the Silhouette index by increasing the number of CRS objects selected by MM, and choose $|CRS|^*$ such that the Silhouette index is maximised. Therefore, MMAD first initialises |CRS| = [0.4 * |S|], and chooses |CRS| objects in S by MM sampling (Figure 1(c)). This value for |CRS| is chosen because it is assumed that majority of the data is normal. Therefore, in the worst case, at least 60%of data is normal. MMAD starts with $Silh_1 = 0$. Then, it picks the first two CRS objects and computes $Silh_2$ considering these objects as cluster centers. From there, MMAD computes $Silh_3$ after adding the third CRS as a new cluster center, and keeps repeating this until the last selected object by MM is added to the cluster centers. Let the Silhouette index be maximised at $Silh_m$. Then, $|CRS|^* = m$, and the size of CRS is reduced such that it comprises the m first CRS selected by MM sampling (Figure 1(d)). After this, a further adjustment is performed on CRS as follows (lines 8-11 of Algorithm 1). A crisp *m*-partition U_m is built by considering each of the *m* samples in *CRS* as a cluster center. Each of these m points is then replaced by the closest point to the cluster mean in its partition (Figure 1(e)). Since the representative samples are chosen using MM sampling, the cluster centers, i.e., CRS_s , are expected to be close to the border of their partitions. Therefore, this further adjustment is required to locate them close to the center of the partition.

We identified two types of datasets when using the Silhouette index to estimate $|CRS|^*$: **Not Well-Separated** (NWS) and **Well-Separated** (WS). If the value of the Silhouette index in $[Silh_{1..|CRS|}]$ has an increasing trend peaking at $Silh_{|CRS|}$, then anomalies and normal samples are not well-separated, otherwise anomalies and normal samples are well-separated. Examples of these types of datasets are shown later in Figure 3. This property is used in subsequent stages.

3.2 Stage 2: CRS Screening

This stage has two steps. First, an unsupervised heuristic approach is used to detect anomalies in the CRS set, which is generated in Stage 1. Removing potential anomalies from CRS results in creating the RS set, i.e., representative samples. Second, if a budget for AL is available, i.e., a small number of labels are allowed to be asked from an oracle, an optional active learning step is used to improve accuracy.

Algorithm 3 Budget Spending (BS)

Input: Representative Samples RS, budget B, dissimilarities Δ_{RS} Output: I_{RS} 1: $\{y_i^{RS} = 1\}, i = 1..|RS|$ 2: if $B \ge |RS|$ then 3: $y_i^{RS} = l_i, 1 \le i \le |RS|$ \triangleright get true label per RS_i 4: else $\Pi(I_{RS})$: same order as selected by MM sampling 5:6: relocate indexes of clusters with one member to end of $\Pi(I_{RS})$ 7: for i = 1..|B| do $\begin{array}{l} y_{\Pi(i)}^{RS} = l_{\Pi(i)}, 1 \leq i \leq |RS| \\ \text{if } y_{\Pi(i)}^{RS} = -1 \text{ then} \end{array}$ 8: \triangleright get true label of $RS_{\Pi(i)}$ \triangleright find indexes of other poor RSs 9: $NNs = find(e^{-\gamma \times \Delta(\Pi(i),j)} > 0.5)$ 10: $y_{NNs}^{RS} = -1$ 11: 12:end if end for 13:14: end if 15: $I_{RS} = \bigcup_i$ where $y_i^{RS} == 1$

Given that in unsupervised learning anomalies are a minority, it is expected that anomalous clusters in data have fewer members compared to normal clusters. Based on this intuition, if a dataset is classified as type WS (i.e., normal data and anomalies are well-separated according to the Silhouette index values), the set CRS is used to generate RS as follows. Samples in CRS are sorted in the reverse order of the number of their cluster members. Let π^{CRS} denote the permuted indexes for the ordered set and n_{π}^i denotes the corresponding number of members in the *i*th largest cluster indexed by π_i^{CRS} where i = 1...|CRS|. The RS set is initially empty. Then, samples from CRS in the order given by π^{CRS} are added to RS until n_{π}^i is less than a predefined threshold. In this paper, we evaluate values of n_{π}^i as follows: when $\frac{n_{\pi}^{i-1}}{n_{\pi}^i} \geq 2$, we stop adding the remaining samples from CRS to RS. In contrast, if the dataset is classified as type NWS (i.e., differentiating anomalous and normal objects is difficult because they are not well separated by their distance nor by density), we choose RS = CRS because there is insufficient information available to filter anomalies.

Active Learning Sub-stage In some application contexts, it may be possible to apply AL to assign a label for one or more selected points as either normal or anomalous by asking an oracle for the labels. Given that the budget for asking an oracle for labels is usually restricted to a small number of points, a key challenge is how to select the points that are likely to be most informative in terms of improving the accuracy if a label is given. If a budget B is available via AL, we can ask for a maximum of B labels from the oracle. If $B \ge |RS|$, all the RSs can be validated by the oracle to remove any anomalies that were left from the screening stage. Otherwise, labels are asked for clusters that have more than one member in the order that they were selected by MM sampling. Anomalous RSs are removed accordingly. This stage of removing anomalous RSs are shown in Algorithm 3 and can take place here or at any time after building the model.

3.3 Stage 3: Model Construction

For the choice of the kernel function, in this paper we use the *Radial Basis kernel* Function (RBF), $k(x, y) = e^{-\gamma ||x-y||^2}$ to localise the influence of *RS*s in the final model. The γ parameter is the kernel bandwidth and we set it based on the dataset type identified in Stage 1.

If the dataset is classified as type WS, anomalies and normal data are declared separable. The γ parameter in this case is estimated using the technique proposed by Evangelista et al. [13]. The reason is that this technique chooses a value for γ that maximises the dissimilarity of clusters in the data. We use a candidate set $\Gamma = \{2^{-6}, 2^{-5}, ..., 2^{6}\}$ for γ to extract the kernel matrix $K = [e^{-\gamma \times \Delta}], \ \Delta = [\delta(s_i, s_j)] \ \forall (i, j) \in 1... |S|$, and select a γ^* such that:

$$\gamma^* = \underset{\gamma}{\operatorname{argmax}} \frac{\sigma^2}{\overline{K}_{offDiag} + \varepsilon},\tag{4}$$

where σ^2 and $\overline{K}_{offDiag}$ are the variance and the mean of the off-diagonal kernel matrix entries, and ε is a small value to avoid division by zero.

If the dataset is classified as type NWS, potential anomalies are close to normal data. Therefore, γ^* is chosen so that the similarities in the feature space created by the RBF kernel are approximately the same as those in the input space:

$$\frac{e^{-\gamma \Delta_{max}^2}}{e^{-\gamma \Delta_{min}^2}} = \frac{\Delta_{min}}{\Delta_{max}} \Longrightarrow \gamma^* = \frac{-ln(\frac{\Delta_{min}}{\Delta_{max}})}{\Delta_{max}^2 - \Delta_{min}^2}.$$
(5)

where the values of Δ are an output of Algorithm 2. To build the final model, values of $\{w_{1..|RS|}\}$ are assigned for $RS = \{s_{I_{RS}^{1..|RS|}}\}$ and the rest of the data is deleted. For the *j*th *RS*, its weight is defined as $w_j = \frac{cn_j}{\sum_{t=1..|RS|} cn_t}$, where cn_j denotes the number of cluster members for the corresponding *RS*. The final model is $M = (RS, \{w_{1..|RS|}\}, \gamma)$ and a test point is evaluated using the following scoring function:

$$P(x; \{w_{1..|RS|}\}) = \sum_{i=1}^{|RS|} w_i \times k(x, RS_i).$$
(6)

If AL is used after constructing the model and deleting the training data, RSs labelled as anomalies by the oracle are deleted from the model, and the weight vector for the rest of the RSs is normalised to add up to one again.

4 Experimental Evaluation

We compare our MMAD method¹ to OCSVM [30] and its unsupervised extensions *Robust OCSVM* (ROCSVM) and η OCSVM [3], and also to iForest [25].

¹ Implementation of MMAD is available at https://github.com/zghafoori/MMAD.

Table 1. Description of the	datasets
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Dataset	DSA	HAR	MNIST	Cancer	Credit	Shuttle	NKH	Pen	Satimage2	Forest
#Instances	1,117,521	10,299	70,000	675	283,726	36,752	221,902	6,870	$5,\!801$	286,048
#Features	45	561	784	10	30	9	3	16	36	10
#Normal	60,000	1,777	$7,\!877$	439	283,253	34,108	221,830	6,714	5,732	$214,\!839$
#Anomaly	$1,\!057,\!521$	8,522	$62,\!123$	236	473	$2,\!644$	72	156	69	2,747

The default settings appearing in the referenced papers were used for each of these existing techniques. For iForest the default sample size is 256 and it trains 100 trees. For MMAD, the sample size |S| = min(200, [|D|/2]). We repeated our experiments by changing |S| in the range [100, 500], and observed that for |S| > 200, the result does not change significantly. Euclidean distance was used as the dissimilarity metric δ , but any other metric can be used when circumstances warrant a departure from this choice. To evaluate accuracy, we used the *Receiver Operating Characteristic* (ROC) curve and the corresponding *Area Under the Curve* (AUC). The reported AUC values were averaged over 100 runs. The experiments were conducted on a machine with an Intel Core i7CPU at 3.40 GHz and 16 GB RAM. The MATLAB LIBSVM toolbox [9] was used to implement OCSVM.

4.1 Datasets

We ran our experiments on four benchmark anomaly detection datasets from the *Outlier Detection DataSets* (ODDS) collection [28], namely *Forest Cover* (Forest), *Pendigits* (Pen), Satimage2, and Shuttle. From the *UCI Machine Learning Repository* [24], we selected the *Breast Cancer Wisconsin* (Cancer), MNIST, *Human Activity Recognition* (HAR), and *Daily and Sports Activities* (DSA) datasets. Table 1 shows that DSA, HAR and MNIST contain many more anomalies than normals. In the experiments we use a random subset of anomalies such that majority of data is normal.

For the Cancer dataset, the aim was to detect malignant breast cytology as anomalies. For the MNIST dataset, following Rayana [28], Bandaragoda et al. [5] and Amarbayasgalan et al. [2], digit zero was considered as the normal concept and instances of digit six were considered as anomalies. The HAR dataset included sensor signals of six different activities by a group of 30 volunteers within the age range [19, 48]. In this dataset, we used the sitting activity as the normal concept and walking in different ways, standing and laying as anomalies. The DSA dataset comprises sensor signals for 19 activities, each of which is performed by four females and four males within the age range [20, 30]. Again, the first activity (sitting) from all the 8 subjects in this dataset is considered as normal and the rest of activities from all subjects are considered as anomalies. This creates clusters of different shapes and cardinalities (Figure 2(a)) in order to evaluate the effectiveness and robustness of anomaly detection methods. We removed duplicates from normal and anomalous instances, which resulted in



Fig. 2. Visualisation of the datasets

generating a dataset with 60,000 normal and 1,057,521 anomalous instances for DSA. We also used the *Credit Card Fraud Detection* (Credit) [10] dataset that contains credit card transactions of European cardholders in September 2013. The goal was to detect fraudulent transactions as anomalies. Finally, from the NSL-KDD² dataset, we used HTTP data, which we refer to as *NSL-KDD*-*HTTP* (NKH). Attacks on HTTP services were regarded as anomalies in NKH.

The datasets' descriptions including the number of unique observations and features are summarised in Table 1. Down-sampling on anomalies and normal instances (if required) is used to generate training sets with anomaly fractions chosen from the set $\{0.01, 0.05, 0.1, 0.15, 0.2, 0.25, 0.3\}$. For each fraction of anomalies we randomly selected up to 10,000 instances from each dataset, and repeated the experiment 100 times. All datasets were re-scaled in the range [0, 1] based on maximum and minimum values observed in the training set. Test and training sets were randomly selected with a ratio of 1 to 4.

4.2 Results and Discussion

For each dataset, we took a subset of the data and used a variant of t-SNE called LN-SNE [14] to visualise it in Figure 2. The labels were used to draw the plots and were not provided to any of the anomaly detection techniques. This figure shows that there are different types of anomalies in the datasets. DSA and MNIST mainly have point anomalies, while shuttle and NKH have clusters of anomalies. Some of the anomalies in HAR and Credit occur inside the normal region. The density and distance similarities for anomalies is similar to the normal samples in Satimage2 and Forest. In the Pen dataset, anomalies have higher density compared to the normal samples and appear very close to them.

To see how the Silhouette index (3) changes by increasing the number of selected representative objects using MM sampling, in Figure 3, we plotted graphs of the Silhouette values for three datasets, namely DSA, HAR and Pen. The

² http://nsl.cs.unb.ca/NSL-KDD/



Fig. 3. Silhouette values obtained by increasing the number of objects selected by MM sampling for three representative datasets. For the well-separated datasets DSA and HAR, the the Silhouette value is maximised at an index less than 80, which indicates that separable clusters are identified in the data. However, for the Pen dataset, this index continues to increase to the maximum at 80, which means the data forms a single cluster.

other datasets exhibit a similar trend so we do not present their corresponding plots to save space. The samples size was |S| = 200. The fraction of anomalies was increased in the sub-figures from (a) to (g) to test the robustness of the proposed technique to the anomaly ratio in different types of datasets. The fractions of anomalies are (a) 0.01, (b) 0.05, (c) 0.1, (d) 0.15, (e) 0.2, (f) 0.25, (g) 0.3. The index drops for DSA and HAR but reaches its maximum at |CRS| = 80 for Pen. Therefore, the DSA and HAR datasets are categorised as type WS (i.e., wellseparated) based on the definitions in Section 3.1. The difference is that in DSA the normal data comprises several clusters and anomalies are point anomalies; each anomaly is treated as a separate cluster center to maximise the Silhouette index. However, in HAR there are two clusters in the data and one of them is anomalous. The Pen dataset is an example of a type NWS dataset (i.e., not wellseparated) because the value of the Silhouette vector is maximised at the end. It can be confirmed via the visualisation given in Figure 2 that anomalies and normal data in Pen are not well separated via distance or density dissimilarities when no ground truth is available.

Figure 4 depicts the average accuracy over 100 runs of each method for all the datasets for each anomaly fraction. This box-plot shows how the accuracy of the methods changes under different conditions of anomaly type and fraction. Each box-plot shows the variations of AUC for different fractions of anomalies including $\{0.01, 0.05, 0.1, 0.15, 0.2, 0.25, 0.3\}$. Increasing the fraction of anomalies can affect the accuracy of the corresponding technique in a nega-



Fig. 4. The average and standard deviation of AUC for MMAD and other techniques.

	Table 2	. Results	of Wilcoxon	test for	the accuracy	of MMAD-B0	vs other methods.
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MMAD*-B0 Vs	*-B1	*-B5	*-B10	*-B15	*-B20	OCSVM	ROCSVM	ηOCSVM	iForest
R^+	141	85	84	84	83	2296	2396	2391	1970
R^{-}	1344	1806	1807	1807	1808	189	89	94	515
<i>p</i> -value	2.23E-07	6.38E-10	6.09E-10	6.09E-10	5.82E-10	7.04E-10	1.47E-11	1.80E-11	2.07E-05

tive manner. For MMAD, the results are reported for different AL budget limits $B = \{0, 1, 5, 10, 15, 20\}$. MMAD-B0 means that no budget was available for AL, while MMAD-B* where $* \neq 0$ means that labels for * number of samples could be asked from the AL oracle. MMAD-B0 and iForest have better results than the different versions of OCSVMs. On average, MMAD-B0 works better than iForest considering all scenarios of the anomaly fraction over all the datasets, especially in HAR, Shuttle and Satimage2. Using an AL budget in the first 7 datasets provides limited advantage because MMAD-B0 effectively screens anomalies in the training phase for these datasets. However, for Pen and Forest, which had the most difficult type of anomalies, access to a limited AL budget B = 5 or even B = 1 in Forest, increases the accuracy to a great extent.

To assess the statistical significance of the differences in the accuracy shown in Figure 4 for each method, Table 2 lists the results of a Wilcoxon signed-rank test with a level of significance of $\alpha = 0.05$ on all the pairs of accuracy values per the anomaly fraction. In each comparison, the aim was to investigate to what extent the null hypothesis H_0 , which indicates that there is no difference between the first and second methods in terms of their accuracy, can be rejected. For each comparison, the test returns the sum of positive ranks (R^+) and negative ranks (R^-) of the first method, and the *p*-value. The *p*-value represents the lowest level of significance of a hypothesis that results in a rejection and if it is less than

Dataset		Train t	time (secon	ds) of			Test time	e (seconds j	per sample) of	
Databot	MMAD	OCSVM	ROCSVM	ηOCSVM	iForest	MMAD	OCSVM	ROCSVM	ETAOCSVM	iFOREST
DSA	0.093	26.450	29.323	32.439	0.19	3.15E-07	1.27E-04	1.32E-05	4.85E-06	1.38E-03
HAR	0.116	38.047	39.759	38.141	0.41	9.75E-06	$2.70\mathrm{E}\text{-}04$	1.27E-04	1.08E-04	1.37E-03
MNIST	0.127	269.453	281.008	289.487	0.67	3.43E-05	1.76E-03	4.76E-04	2.70E-04	1.47E-03
Cancer	0.060	0.054	0.060	0.114	0.10	1.79E-06	2.15E-05	1.70E-05	1.26E-05	1.37E-03
Credit	0.097	46.705	48.847	46.752	0.15	3.29E-07	9.26E-05	1.30E-05	8.61E-06	1.47E-03
Shuttle	0.090	18.605	19.171	34.718	0.16	7.88E-07	7.04E-05	1.71E-05	1.15E-05	1.36E-03
NKH	0.098	9.272	7.377	16.662	0.11	< 1E-12	7.44E-05	1.80E-05	9.42E-06	1.50E-03
Pendig	0.088	11.042	11.911	24.427	0.25	5.66E-06	6.40E-05	2.37E-05	1.58E-05	1.53E-03
Satimage2	0.091	9.588	10.379	10.867	0.20	1.21E-06	5.83E-05	1.74E-05	1.08E-05	1.52E-03
Forest	0.093	28.479	29.537	39.239	0.20	2.36E-06	5.31E-05	9.30E-06	7.48E-06	1.47E-03
Average	0.095	45.770	47.737	53.285	0.24	5.65E-06	2.59E-04	7.32E-05	4.59E-05	1.44E-03

Table 3. Training and testing CPU-times for MMAD compared to other methods.

 α , the null hypothesis H_0 can be rejected and the improvement is significant at the level α . Table 2 shows that spending even a limited budget B = 1 provides a statistically significant improvement and by increasing the budget, a better result can be achieved. This finding shows the importance of using a technique that can incorporate AL into its training. The small *p*-values for MMAD-B0 against OCSVM, ROCSVM, η OCSVM, and iForest, MMAD-B0 indicates that MMAD-B0 provides a statistically significant improvement compared to all of the comparison methods.

Table 3 reports the training and test CPU-times of the different techniques. MMAD and iForest have constant time given that they work on a fixed-size sample of the data. Given the size of the sample |S|, the training time-complexity for MMAD is $O(d \times |S|^2)$ and for iForest is $O(t \times |S| \times log|S|)$, where t is the number of trees. For testing, the time-complexity of MMAD is $O(d \times |RSs|)$ per instance, whereas it is $O(t \times log|S|)$ for iForest. The table shows that on average, the training time of MMAD is half of that for iForest, mainly because its default sample size is 200, whereas it is 256 for iForest. However, the testing time per sample for MMAD is more than 250 times less than iForest. For the HAR and Credit datasets, testing times per sample for MMAD are more than 4,000 times faster than iForest. Both MMAD and iForest are considerably faster than the OCSVM-based methods. The improvements by MMAD in accuracy and its capability of being used with AL, demonstrates that MMAD outperforms iForest on the examined datasets in this paper. To compare the scalability of MMAD and iForest to the different types of OCSVMs, we chose the DSA dataset with over a million instances, and ran all the methods several times by changing the size of the training set from 10,000 to 50,000 with a step size equal to 10,000. The results are shown in Figure 5, confirming that MMAD and iForest have constant time regardless of the size of the training set, while the training time



Fig. 5. Scalability of the five methods on DSA.

of the OCSVM variants depends on the size of the training set. This example shows the superiority of algorithms with constant time in processing big data.

5 Conclusion

We have proposed a constant time unsupervised anomaly detection technique called MMAD that can be used effectively with active learning to enhance the accuracy of unsupervised anomaly detection when anomalies mimic the behaviour of normal data. MMAD combines a representative subset selection with a cluster validity index and kernel-based model construction in a novel way that results in statistically significant improvement of the accuracy and training time for unsupervised anomaly detection on the examined datasets. In our future work, we will study the use of kernels other than RBF, distance measures other than Euclidean, and CVIs other than the Silhouette index. The information gained via active learning can be used in new ways to further identify the dataset characteristics and improve the accuracy. Finally, this technique can be extended to perform constant time clustering for big datasets that have point or cluster anomalies.

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- 16 Z. Ghafoori et al.
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