

## École Polytechnique

BACHELOR THESIS IN COMPUTER SCIENCE

# Improving anomaly Explanation methods on time series data

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Academic year 2024/2025

#### **Abstract**

The increasing amount of data being created and processed in all sectors has led to the use of automatic methods for filtering and analysing this influx of information. These methods often rely on complex models whose methods are intractable and as such act as "Black Boxes". Such techniques are used for monitoring traffic on many different types of servers. Anomaly detection methods are important as they make it possible to know in real time when an issue has occured. However, in order to efficiently resolve any issue, as important as detecting an anomaly is understanding why the flagged interval is anomalous. This is where the field of Explainable AI comes into play. In this paper, we will improve Exstream, which is a method relying on the single feature entropy measure between normal points and anomalous points to determine features that contribute the most to the anomaly and the feature intervals where anomalies would happen. Our contribution is twofold, first we explored the possibility of sampling normal points using the latent space from the detection method. These points would be more representative of the current anomaly and would not be correlated with time. We found that the domain dependency of the data makes this technique unfeasible for our type of data. Secondly, we showed that the scoring for different features is not a submodular function due to correlated features but is simply increasing. We also generalized the entropy measure over multiple features which would enable the explanation of anomalies on more complex datasets where the interaction of two or more features is responsible for the anomaly.

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## 1 Introduction

An anomaly can be defined as "an observation which deviates so much from other observations as to arouse suspicions that it was generated by a different mechanism"[5]. As such, Anomaly detection is the task of identifying these observations. In recent years, anomaly detection has become a core part of the machine learning landscape. Its applications are numerous and can be found in many different sectors such as healthcare [3], finance [1] or security [14] [9]. These methods respond to the need to automate tasks that would be impossible to do manually due to the sheer amount of data on hand. In the medical domain, they can pick up on signs of a disease from medical imagery that would be impossible to detect for a human. In finance, fraudulent transactions can be pinpointed among the millions happening each day. In the domain of network and server security, they enable a real time monitoring of traffic.

However, in each domain detecting an anomaly is only the first step. This anomalous instance must then be acted upon. In most cases a human must intervene. It is therefore crucial to understand the reason why an anomaly was flagged. This also increases the confidence in the method and allows us to verify it is insight and provide solutions. Simple methods such as decision trees or single feature monitoring methods [6] can provide a list of features that contributed to the anomaly. However, with the advent of deep learning [16], anomaly detection methods have become more complicated and less interpretable. These methods are more accurate as they can capture more complex patterns but they are also more opaque. They rely on "Black box" models that were trained to detect anomalies but give no insight into the reason why an anomaly was detected. This is where the field of anomaly explanation comes into play. The objective of anomaly explanation is to provide a human readable and actionable explanation of the anomaly. In most cases, this will be a set features that contributed to the anomaly.

The objective is to provide "good" explanations. These must satisfy the following criteria:

- Conciseness: Explanations should be short in order to be easily understood and interpreted by a human.
- *Consistency:* The explanation must be consistent with the interpretation of it by a human. In practice, this means obtaining similar explanations for the same type of anomaly happening in the same context.
- *Prediction power:* Explanations should be able to predict future anomalies. This is especially important in the case of time series data.

## 2 Related works

In this section we will go over previous works in the field of anomaly explanation. We can separate Anomaly explanation methods into 2 categories: model explainers and data explainers [10] [17].

Model Explainers are methods that provide explanations for the output of a model. These methods use the model to determine the explanation. Such methods include SHAP [12], LIME [19] and Anchors [20]. LIME and Anchors perturb the data around the anomaly in order to determine the closest decision boundary and therefore the features that contributed most to the anomaly. LIME uses the perturb data and outputs from the model to train a simpler more interpretable model such as a decision tree or a linear regression. Anchors on the other hand uses the predictions on the generated data to create "if then" rules called Anchors that describe the conditions under which a model consistently makes the same prediction. The predictions of LIME depend heavily on the kernel width which defines the area of interest in the generation of synthetic data making the model unstable [23]. Anchors fixes this issue, however it becomes expensive on large datasets and is sensitive to feature selection. Overall, these two methods do not provide any theoretical guarantees as to the quality of the explanation.

SHAP on the other hand uses the SHAPLEY values concept from game theory to find a good set of features as explanation. This method treats the prediction task as a game, where each feature is a player that contributes to the final prediction. This method is computationally expensive and also assumes the independence of the

features which is not always the case.

**Data Explainers** on the other hand do not use the model in order to provide an explanation for anomalies. These methods takes anomalous points and normal points as input and from those provide an explanation. Therefore these methods are model agnostic. Such methods include Exstream, COIN, Macrobase and lookout. Lookout [4] is a method that attempts to find the pair of features where the anomaly is best explained in a visual manner. This is done by finding the two dimensional space where the anomalous points have the highest anomaly score as evaluated using an isolation forest. The features being returned would be the ones in which the anomaly stands out most. Macrobase [2] uses Relative Risk Ratio, a metric from epidemiology to find the features that are most responsible for the anomaly. However, this method is limited to categorical data in its explanations. It also does not take into account correlation in the features being explained. Finally, in order to have good explanations, relationships between entries in the data must be known. This is not suited for raw data. COIN [11] is a method that uses the nearest neighbors of an anomaly in order to find the best set of features to explain them. In order to do so, we find the closest cluster of normal points to our anomalous sample set. These will serve as our reference. We then fit a linear decision boundary between them. This line is constructed so that the number of non zero weights is small. These weights will indicate the features that most contribute to the anomaly. This method is reliant on a good selection of the nearest cluster which limits the local stability of the model. Moreover, the method uses a linear classifier, assuming the decision boundary is a line which is not always the case.

## 3 The Exathlon project

The Exathlon project was started in 2021 as an anomaly detection and explanation benchmark [8]. The github repository implements a pipeline that implements all needed steps from preprocessing of the data to training and evaluating methods for detecting and explaining anomalies. The full pipeline can be seen in Figure 14.

## 3.1 Dataset

The pipeline supports any dataset and custom preprocessing steps can be implemented. For the sake of this paper we will be using a datasets consisting of time series data in the form of traces taken from Apache Spark servers. These traces consist of real data collected from 93 repeated executions of 10 distributed streaming applications on a 4-node Spark cluster over a period of 2.5 months. Each of these executions includes 5 randomly selected applications running concurrently. In total, 2,283 metrics were monitored once per second creating a dataset totaling more than 24GB in size. For the sake of our experiments, the dataset went through some preprocessing, cutting it down to 237 features. It can be noted that this preprocessing step is done automatically in the pipeline. The dataset consists of 59 undisturbed traces and 34 disturbed traces constituting 97 anomaly intervals. There are 6 types of anomalies that were artificially produced. These are the following:

- T1: bursty input: temporarily increase the input rate by a given factor for a duration of 15-30 minutes.
- T2: bursty input until crash: Similar to T1 except the input rate is maintained at a high rate until experiencing a crash due to lack of memory.
- T3: stalled input: artificially set the input rates to 0 for about 15 minutes.
- T4: CPU contention : consume all CPU cores on a given Spark node.
- T5: driver failure: artificially failing the driver processes.
- T6: executor failure: artificially failing the executor processes.

• T7: a mix of unknown types of anomalies.

The data is processed and then turned into a *window dataset* which transforms the data into a sliding window format in order to detect the anomalies.

## 3.2 Anomaly detection

The anomaly detection consists of three steps. First training the *window model*. This model gives an anomaly score to a window of a certain size. Then following the "Unifying anomaly detection method" introduced by Vincent Jacob [7], we obtain an "Online" anomaly scoring function which given the scoring for windows, attributes a score for each point. This scoring function is used to determine a threshold above which points will be labelled as anomalies in the anomaly detection step.

Many different models belonging to various families of methods [22] are implemented and ready to be evaluated.

Such methods include among others: pca, xgboost, Autoencoder, Variational Autoencoder, LSTM, deep SVDD, deep SAD, iForest...

The most recent and best model on this dataset is Divad.

## **3.2.1** Divad

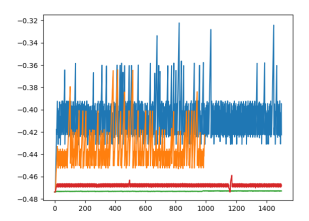


Figure 1: Plot of feature "driver BlockManager memory memUsed MB value" for 4 normal traces over time.

Divad is an anomaly detection method based around a VAE architecture [7]. This method was created to address the difference in the behaviour of traces based on the parameters and properties of the Spark application they were run on. We will call "Domain" the context in which the application is run. This context is characterised by the following elements: processing period, number of active executors, memory profile and input rate.

The different domains pose a real problem when trying to identify anomalies. A certain behavior can be considered anomalous in one domain but not in another. The limited amount of data also makes it impossible to train a model for each domain. Therefore we need a method that can detect anomalies and generalize to new domains. This is Divad.

The way it works is by assuming two independent priors that define the data z. The first prior is the class y, either anomaly or normal. The second prior is one that encodes the domain of the point Figure 2b. Therefore we are assuming that the points can be generated from their class and their domain and that these two priors are

## independent.

The objective is thus to approximate these prior spaces from the data. In order to do so, Divad employs a Variational Autoencoder structure with two independent encoders and one decoder. The first encoder is meant to capture the Class information and the second the Domain information Figure 2a Figure 15. This is done using well chosen error functions.

We therefore end up with two latent spaces:  $z_y$  and  $z_d$  that respectively contain information about the class and the domain of the data. We then use these two latent spaces to reconstruct the data.

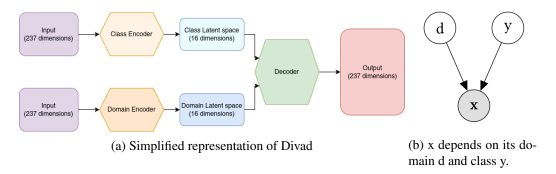


Figure 2: Divad architecture

We get an anomaly score by comparing the class latent space with what we would expect from the normal class. This is done by computing the KL divergence between the two distributions.

## 3.3 Anomaly Explanation

The anomaly explanation step will be the main focus of this paper. The objective of this step is to give a human readable and actionable explanation of the anomaly. This can be done in many ways such as lists of feature importance or plots as we have seen in the related works. In this paper, the focus will be on the Exstream method. For each anomaly, this method will output the features that cause the anomaly as well as the intervals where the anomaly is happening.

The explanation is given as a boolean expression in Conjunctive Normal Form. It consists of a conjunction of clauses, where each clause is a disjunction of predicates. Each predicate follows the form  $\{v \ o \ c\}$ , where v represents a feature, c is a constant, and o is one of the five operators:  $o \in \{>, <, >, <, =\}$ .

For example an explanation for an anomaly could be the following: ((feature 1 > 0.5)  $\lor$  (feature 1 < 0.8))  $\land$  (feature 2 = 0.7).

In practice, the explanation is given as a json file with one entry per anomalous interval in the following format:

```
end,
                       include start,
                       include end
                  ]
             ],
             "feature 2":[
                       start,
                       end,
                       include start,
                       include end
                  ],
                  Γ
                       start,
                       end,
                       include start,
                       include end
                  ]
    },
}
```

There previously existed two versions of Exstream. The original version was introduced by Haopeng Zhang, Yanlei Diao and Alexandra Meliou in 2017 [24]. It was then worked uppon by Mija Pilkaite during her Bachelor thesis [18] and finally refined and improved by Clement Martineau [13].

## 3.3.1 Original Exstream

Originally, Exstream was created as a rigorous and formal method for providing short and human readable explanations for anomalies. This method presents the challenge of finding the optimal explanation to an anomaly as a submodular optimization problem. The intuition behind this model is that adding features to our explanation has a diminishing return. This is because the more features we add, the less information new features will provide.

Submodular optimization being NP-hard, a heuristic was therefore introduced to solve the problem. This method relies on Entropy to compute single feature scores.

We start with two sets of points  $S_a$  and  $S_n$  containing respectively the anomalous and normal points. For each feature  $f_i$ , we will sort the points in  $S_a$  and  $S_n$  and merge them into one set. We will then define segments on this set as neighboring points of the same class.

We will compute a segmentation entropy defined as follows: If there are n segments, and  $p_i$  represents the ratio of data points included in the ith segment, the segmentation entropy is:

$$H_{segmentation} = \sum_{i=1}^{n} p_i \log(\frac{1}{p_i}) \tag{1}$$

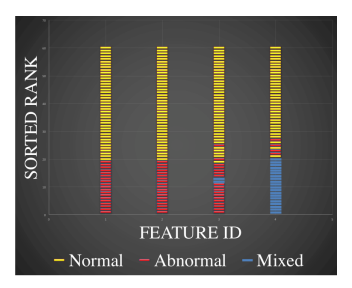


Figure 3: Visualization of the segments of 4 features. The red points are the anomalous points, the yellow points are the normal points and the blue are mixed. Segments are continuous points of the same colour.

Exstream was originally aimed at categorical data. Therefore there can be features with values that belong both to anomaly and to normal points. We need to penalize these features. This is done by giving the worst score to those points of mixed class. Let  $c_i$  be a mixed interval and  $c_i^*$  be the segment rearranged in its worst ordering.

Then the maximum entropy is defined as:

$$H_{max} = \sum_{i=1}^{n} H_{segmentation}(c_i^*)$$
 (2)

Finally, we normalize the score by the class entropy to get a value between 0 and 1. The class entropy is defined as follows: Let  $|S_a|$  and  $|S_n|$  be the number of points in  $S_a$  and  $S_n$  respectively.  $p_a = \frac{|S_a|}{|S_a| + |S_n|}$  and  $p_n = \frac{|S_n|}{|S_a| + |S_n|}$ . The class entropy is then:

$$H_{class} = p_a \log(\frac{1}{p_a}) + p_n \log(\frac{1}{p_n})$$
(3)

The final score for a feature is then:

$$score(f_i) = \frac{H_{segmentation} + H_{max}}{H_{class}} \tag{4}$$

This score reflects the segmentation of the feature with respect to anomaly and normal classes. The higher the score, the more separated normal and anomaly classes are. For example a feature with a score of 1 would have all the anomaly points in one segment and all the normal points in another. As we can see in Figure 3, the first two features have a score of 1. The third a score of 0.31 and the fourth 0.18.

Exstream selects the best features based on reward leap filtering. It also provides methods to filter out redundant features using correlation clusters. It also removes "false positive" features which are defined as those with excessively large standard deviation or that tend to only increase or decrease on both the normal and anomalous interval.

## 3.3.2 Exstream using Bins

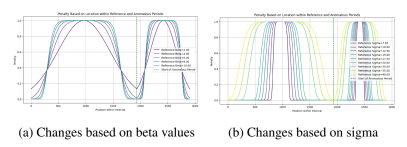


Figure 4: Normal distributions used to weigh points.

An attempt to improve Exstream was made by Mija Pilkaite during her Bachelor thesis [18]. The first idea was to use bins to discretize the data. This would make the method more suitable for continuous data. The values of each feature would be divided into b bins. Each bin would be marked as either Normal, Anomalous or Mixed. The entropy would then be computed on these bins instead of points. The second idea was to reduce uncertainty in the anomalous interval boundaries. Even when using ground truth segments with anomalous intervals marked by hand, there is always some uncertainty as to exact endpoints of the intervals.

In order to minimize this labeling uncertainty, the points would have different weights in the computation of the entropy depending on their position in the trace. This weighting would be done using a Gaussian distribution centered around the middle of the interval as such:

$$w_t = \begin{cases} \exp\left(-\left(\frac{|position - \frac{interval\_length}{2}|}{\sigma \times \frac{interval\_length}{100}}\right)^{\beta}\right), & \text{if } 0 \leq position \leq interval\_length \\ 0, & \text{otherwise} \end{cases}$$

This was further developed by Clement Martineau who corrected some issues and improved the method in higher dimensions.

### 3.3.3 Metrics

The Exathlon pipeline provides a set of metrics to evaluate the performance of the explanation models. These metrics are the following:

- 1. ED1: These metrics are based on local behavior of the anomaly explanation.
  - Time: The time taken to compute the explanation.
  - Size: The size of the explanation. This is the number of features contained in the explanation.
  - Perturbed Size: Size of the explanation after perturbation. A perturbation is defined as a random sampling of 80% of the data in the normal and anomalous intervals. The perturbed size is the number of features in the explanation of a trace having been perturbed.
  - Instability: A measure of the difference in the explanation before and after a perturbation. A high instability means that the explanation is not locally stable.
  - F1 Score, Precision, Recall: These metrics are computed by giving an explanation on a randomly sampled 80% of the normal and anomalous data and then testing the explanation on the remaining 20%. The testing is done on the intervals being returned as being anomalous. From them a Precision and Recall score are computed. The F1 Score is the harmonic mean of the two. Similarly as for the instability, the random sampling and evaluation are carried out 5 times and the average is taken.

- 2. ED2: These metrics are based on global behavior of the anomaly explanation.
  - Discordance: This is the degree of disagreement between the explanation for anomalies of the same type. This measure is computed for each type of anomaly as the entropy of the number of each feature found in all explanations of this type. The discordance is then normalized by the average number of features in an explanation.
  - F1 Score, Precision, Recall: these metrics are computed by getting an explanation on one sample and evaluating it on all the other test samples of the same trace. Similarly as for ED1, these scores are computed on the intervals being returned as being anomalous. The F1 Score is the average of the F1 Scores computed on each other test sample.

These metrics are all computed for each type of anomaly as well as for the whole dataset.

It is important to note the limitations of these metrics. For anomaly explanation contrary to anomaly detection, there is no ground truth. Therefore getting perfect Discordance and Stability does not imply that the explanation is correct. For example a method returning *feature1* as an explanation every time would get perfect scores for both metrics. This is especially important for explanation methods that do not provide intervals but only important features. This is why inspecting specific examples and determining if the explanation provided is relevant is important.

Overall, these metrics are very useful in comparing different methods. However, they should be taken with a grain of salt and the results should be interpreted with caution

### 3.3.4 Evaluating current methods

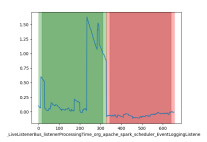
Methods	Time	Length	Instability	Discordance	ED1 F1 Score	ED2 F1 Score	
Exstream	1.5	1.6	2.26	7.71	0.88	0.87	
Exstream 95%	1.35	2.1	2.08	6.59	0.87	0.93	
Exstream 90%	1.2	2.4	1.88	6.04	0.88	0.92	
Exstream 80%	1.1	3.4	2.0	5.5	0.88	0.90	
Exstream Bins	140.7	4.3	1.5	6.16	0.79	0.72	

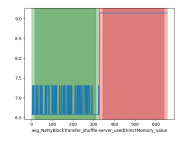
Table 1: Evaluation of Exstream methods

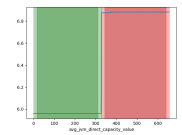
We start by evaluating the current existing Exstream methods and find little improvement in the Exstream Bins method over the orinal. The instability and discordance are reduced but so is the F1 Score in both ED1 and ED2 settings. The length of the explanation is also increased. The true bottle neck however is the runtime which is multiplied by a factor of 100. This limits its use in real time monitoring. The delay is due in part to poorly optimized code but also to the apparent complexity of the algorithm in processing the bins. The algorithm was also never evaluated on the entirety of the Exathlon data and only on a subset with shorted traces which limited the impact on performance.

In order to compare, we also evaluate the original Exstream while removing the first and last10%, 5% and 2.5% of normal and anomalous points. This in practice removes the uncertainty linked to the labeling at the boundaries of the intervals while keeping the algorithm simple and preserving speed. This new method improves the instability and discordance over Exstream while keeping a high F1 Score. The length of the explanation increases. This can be explained by the fact that labeling noise initially created segmentation in well separated features. Removing the noise leads to more well separated features to explain the data. It can also be noticed by looking at the examples that the explanations are more confident and more relevant. There are many more features that are selected with a confidence score of 1.

For example, trace 5\_5\_1500000\_5\_3\_9\_91 anomaly 1\_T6 as seen below. Using all points, the responsible feature is "driver\_LiveListenerBus\_listenerProcessingTime\_org\_apache\_spark\_scheduler\_EventLoggingListener\_stddev" with a score of 0.054. Meanwhile using 95% of the data it is a slightly longer explanation with features : "avg\_NettyBlockTransfer\_shuffle-server\_usedDirectMemory\_value" and "avg\_jvm\_direct\_capacity\_value" with both a score of 1.







- (a) Exstream responsible feature
- (b) Exstream 95% responsible feature 1
- (c) Exstream 95% responsible feature 2

Figure 5: Responsible features for anomaly at Trace 5\_5\_1500000\_5\_3\_9\_91 anomaly 1\_T6. Green are normal points, red are anomalous points. Lighter parts are the 5% removed.

Another element that shows the improvement when discarding 5% of the data is the length of the explanation file. When evaluating the length of the explanation, the metric is the number of relevant features being selected. This discards completely the number of intervals for each feature. The explanation file contains both features and intervals as seen previously subsection 3.3. The length of the explanation file is therefore a useful while imprecise metric when evaluating the method on the same anomalies. In the case using all points, this file has 22,434 lines. When using 95% of the data it has 3,720. This reflects the poor separation observed in many traces which require many intervals to describe their behavior. Using bins, the explanation file has 14,380 lines.

## 4 Objectives

The focus of this paper is to improve Exstream. The version we intend to work on will be the original version. This choice is motivated by a few reasons.

First, looking at the metrics and inspecting a few examples shows that the version using bins does not offer any significant improvements. Secondly, the time to run is multiplied close to 100 times. This performance issue is partly caused by unoptimized code and partly by an overcomplicated algorithm. The algorithm was also never previously evaluated as part of the Exathlon pipeline and only on a subset of the available data. The complexity of the algorithm makes it very rigid and hard to modify in relevant ways while at the same time the runtime is too long to be used in a real time setting.

There are two fundamental ways in which we can improve Exstream. The first is to improve the sample set. The second is to improve the algorithm being used. In this paper we will address both of these points with varying degrees of success.

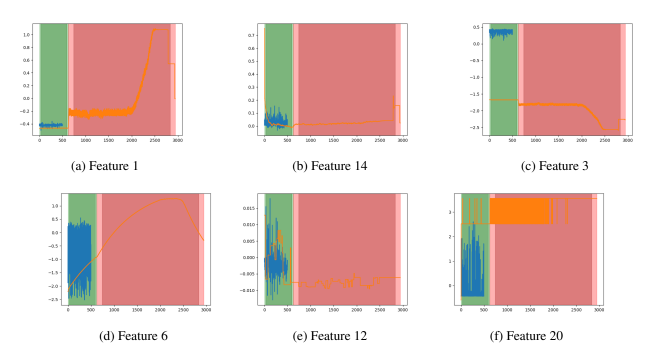


Figure 6: Plot of three features. Green is the normal interval and red the anomalous one. The blue points are reconstructed samples that are "close" to the anomaly, as detailed below.

## 5 Improving the sample set

A problem noticed when evaluating Exstream on the Exathlon data is that the normal points used are not necessarily the most representative of the current anomaly. This is due to the fact that the normal points selected are right before the anomaly. There is therefore a strong correlation between these points and time. The most obvious examples of this are features that are only increasing or decreasing over the normal and anomalous intervals. The segmentation of these features will be perfect and the score will be 1 while these features are not relevant explanations for the anomaly.

This specific case of time correlation is already addressed in the original Exstream paper using false positive filtering. However less obvious cases of correlation are not addressed and can lead to incorrect explanations. At the same time we will attempt to get sample of normal points "closer" to the anomaly. This sample should be more representative of the current anomaly and only the relevant features that induced the anomaly will significantly change. This should enable a better explanation.

## 5.1 Sampling using the latent space

The latent space of a VAE has a well defined structure. The normal points will be distributed following a normal distribution. The space is created such that relevant features to the detection of the anomaly are captures. Therefore the hope is that normal points close to anomalous points are similar to those. Moreover, the difference between those points will be marked more strongly by the features that caused the anomaly. Features unrelated to the anomaly will have similar values while those implicated will be different. This also enables a search on all points for a reference set and not only on preceding points.

The method is limited by the existence of different domains in the data. As stated above subsubsection 3.2.1, a datapoint might be normal in a specific domain while considered anomalous in another. Therefore the points closest to the anomaly in the latent space might belong to a different domain and therefore not be well suited to

interpret the anomaly. In order to address this issue, we will make use of the model to reconstruct the data in the correct domain.

As stated above, Divad uses two encoders, one to extract a latent space encoding the class and the other the domain. These two latent spaces are combined in the decoder to reconstruct the data. We can therefore extract the domain latent space of the anomaly and use it to reconstruct points in the right domain. These points will then be used as the normal points in the explanation.

In practice, some features are well reconstructed, while others not so well. This is due to the fact that the model is not perfect and that some features are more relevant to the anomaly than others. This can be seen in Figure 6. This is especially true in categorical data such as feature 20.

We evaluate the method on the Exathlon data similarly to the original Exstream by removing the first and last 0%, 2.5%, 5% and 10% of the data. The results are as follows:

The results of this method are as follows:

Methods	Time	Length	Instability	Discordance	ED1 F1 Score	ED2 F1 Score
Exstream Reconstruction 100%	5.2	1.6	2.77	8.56	0.79	0.81
Exstream Reconstruction 95%	4.3	1.8	3.17	7.08	0.78	0.87
Exstream Reconstruction 90%	4.3	2.2	2.87	5.59	0.79	0.86
Exstream Reconstruction 80%	4.2	2.6	2.04	5.65	0.79	0.84

Table 2: Evaluation of the Exstream method using reconstructed points

The results are disappointing. The instability and discordance are increased while the F1 Score is decreased. The length of the explanation is however similar. Using reconstructed points also raises another issue which is our confidence in the explanation. If the points being reconstructed are far from the truth, the method might miss some features that would otherwise be a relevant explanation. It is impossible to have a sense of the quality of the output of the algorithm as we are relying on the black box model that we aim to explain.

## **6** Generalizing Exstream to Higher dimensions

As seen earlier, exstream computes entropy on a single feature. This is a limitation as anomalies can be caused by the interaction of two or more features. This method was initially given as a way of finding an approximation to the problem of finding the best set of features to explain an anomaly. This problem was believed to be submodular.

In this section we will start by showing that it is not submodular. We will then present multiple ways the entropy measure can be generalized to multiple dimensions. We will then evaluate these methods and empirically show benefits that are derived from using them.

## 6.1 Submodular optimization

Let A be a set and  $\mathbb{A} = \mathcal{P}(A)$  be its powerset. The set function  $f : \mathbb{A} \to \mathbb{R}$  is submodular if  $\forall X, Y \subseteq \mathbb{A}$ ,

$$f(X) + f(Y) > f(X \cup Y) + f(X \cap Y)$$

It is also equivalent to saying  $\forall X \subseteq Y \subseteq \mathbb{A}, \forall x \in \mathbb{A} \setminus Y$ :

$$f(X \cup \{x\}) - f(X) \ge f(Y \cup \{x\}) - f(Y)$$

The submodularity of a function reflects the diminishing return of adding elements to a set. This property is very useful in optimization problems as it allows for the use of greedy algorithms to obtain an approximate of the solution [15].

This property has been shown to occur in feature selection problems but only in the case where features are independent [21]. This is not the case for our data.

Let us show visually that the submodularity property does not hold. We start by formally defining our setting: We denote the features by numbers from 1 to 237. Let D=[1,237] be the set of all features. Then  $\mathbb{D}=\mathcal{P}(D)$  be the powerset of D. We further annotate each interval of Anomalous and Normal points  $(N_i,A_i)$  by a number i. We then define the function  $f_i:\mathbb{D}\to[0,1]$  as a function computing the entropy of the segmentation of the features being passed for the pair  $(N_i,A_i)$ .

We need the function  $f_i$  to be submodular for each pair  $(N_i, A_i)$ . Therefore let us present an example for which this property does not hold, see Figure 7. Here in either dimension there will be very high segmentation. However, considering both features we see a clear separation between the anomalous and normal points. Using the tree method for multi feature scoring we respectively get 0.012, 0.012 and 0.55 considering  $[f_1]$ ,  $[f_2]$  and  $[f_1, f_2]$ .

Therefore the submodular property does not hold as  $f([f_1]) + f([f_2]) < f([f_1] \cup [f_2]) + f([f_1] \cap [f_2])$ .

This example was specifically chosen but we see that in practice the submodular property is also broken. This

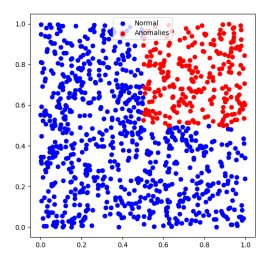


Figure 7: Plot of two features  $f_1$  and  $f_2$ . Blue points are normal  $N_i$  and red points are anomalies  $A_i$ .

property depends on the function being used to compute the entropy. Meaning that with a well chosen function it is possible that the submodular property holds. However, this function would lose a lot of information and would not choose the optimal selection of features.

In conclusion we have proven that the submodular property does not hold for our data. This implies that we do not have an efficient way of finding the best set of features even with a multi dimensional entropy measure.

## **6.2** Methods for generalizing Entropy

Generalizing the computation of entropy to higher dimensions is not trivial. The measure of entropy is computed after creating clusters. In one dimension this is simply done by taking segments containing the same type of points. The number of points in each segment is then used to compute the entropy.

In higher dimensions we wish to keep a similar method. In order to do so we will thus only need to determine

the optimal clustering method to be used.

There are certain properties that our clustering method must have in order to ensure it works properly.

- It must obtain a similar result to Exstream when applied to a single feature.
- It must classify discontinue points into different clusters. Meaning that if there exists an anomalous point in between two normal points, these two normal points should be in different clusters.
- The method must generalize to higher dimensions. The number of dimensions can be limited to  $\approx 10$  as, explained above any more would be too long and would not be useful.
- The method must fit any distribution of the data and work on few points.
- It must be possible to find intervals for the anomalies being clustered.
- The computational complexity of the method must not be "too" high. This is a vague requirement but it is important to keep in mind that the method will be used in a real time setting and will need to be evaluated on many sets of features. An exponential growth in complexity in the number of points would for example make the method unusable.

We will now present a few methods that could be used to cluster the data.

#### **6.2.1** Boxes

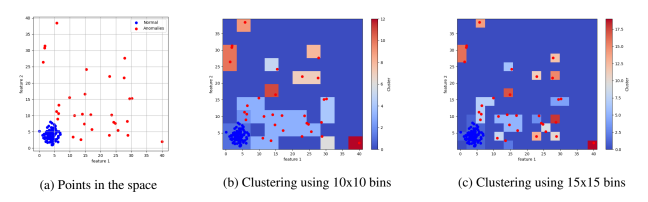


Figure 8: Plot of points being clustered by using the Box method. Same coloured squares are in the same cluster.

The first method is inspired by the bin method for Exstream [18] presented above. The idea of dividing the feature space into bins to discretize the data can be easily generalized to higher dimensions. We would use n dimensional cubes to divide the space. We would then check the points in each cube and notate them as either normal, anomalous or mixed. Having done that, we would then merge adjacent cubes of the same type unless they are mixed in which case it is not useful to merge them. This would give us the clusters we need to compute the entropy.

This method has the merit of being simple and computationally efficient running in O(n) time with n being the number of points considered. However, this method does not generalize well to higher dimensions and to different distributions of the data.

For example in the case with outlying points, as it is expected with anomalous data, the method would not be able to correctly merge the cubes. This would lead to a high number of clusters and a high entropy. The clusters are also highly dependent on the number of bins. The ideal number is impossible to know in advance and can

be different per dimension and per trace.

Overall this method can work on simple data but is not resilient enough for our use.

## 6.2.2 Nearest neighbours

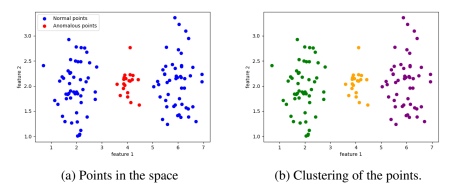


Figure 9: Plot of points being clustered by using the nearest neighbors method. Same coloured points are in the same cluster.

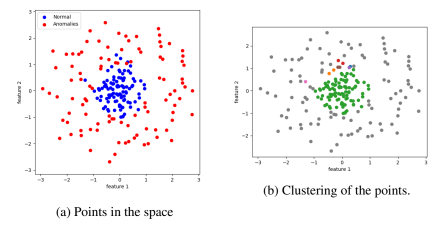


Figure 10: Plot of points being clustered by using the nearest neighbors method. Same coloured points are in the same cluster.

This method can be applied to single points or as an improvement to the post processing of the box method. It relies on the nearest neighbors of each points and is inspired directly by the property stating that we must "classify discontinue points into different clusters".

The method is as follows. For each point, we will iteratively get its next nearest neighbor until reaching a point of the other class. We will cluster these points together. On top of that we merge clusters if a point is already part of a cluster. This method makes it so if two clusters are discontinue, meaning a point of the other class is placed in between them, they will not be merged together. See Figure 9.

In practice we will use a KDtree for storing points and efficiently get the nearest neighbors. We will also restrict the maximum number of neighbors being considered in order to improve the efficiency.

Let n be the number of points and k a hyper parameter being the maximum number of neighbors. The KDtree

is created in  $O(n \log(n))$ , the nearest neighbors are then computed in  $O(kn \log(n))$  and the clustering is done in O(n). Overall the method runs in  $O(n \log(n))$  time. The method is therefore efficient and can be used in real time.

The method obtains the same results as Exstream on a single feature. It also generalizes well to higher dimensions and different distributions of the data as only the segmentation thereof is considered when clustering. The method is relatively stable with respect to the hyper parameter k, the only case where it would matter is for a group of isolated points of size greater than k. Those could be clustered together but not with the rest of the group.

The main concern with this method is the lack of intervals being determined. We only obtain points in clusters. These clusters are not based on intervals. For example Figure 10 shows a case where the gray class does not translate to a clear interval. This is an issue as intervals strengthen the explanation and give much more information about the anomaly.

#### **6.2.3 Decision Trees**

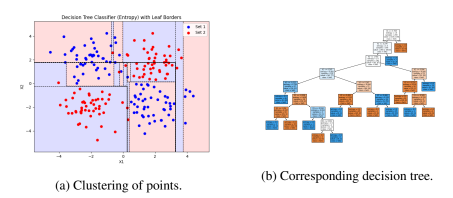


Figure 11: Clustering of points and corresponding decision tree. Red points are anomalous and blue are normal. The lines mark the separation between clusters.

The final method we will present is the use of decision trees to cluster points. The idea is to train a decision tree on the two sets of points we have. We then use the leaves of the tree as clusters. Intuitively, a decision tree splits the space into hypercubes. Each node splits the space in two along a certain feature. The leaves are a conjunction of these splits, in essence creating a hypercube subset of the space being considered.

In theory, the decision tree can split the space until each leaf is pure, i.e it contains only points of the same class. This would give us the optimal clustering. However, in practice we will allow the tree to split only while it has  $\geq k$  points in it's branch. Mixed clusters are then handled similarly as in Exstream and given the maximum segmentation score. This is done for efficiency purposes. It is also important to note that this limitation does not impact significantly the results. Reaching this limit would mean that the points are extremely segmented and that the entropy would be high in any case.

Now let us go over the properties of this method:

- The results are similar to those obtained using Exstream when run on a single feature. This is at least true for scores that are close to 1, with a limited number of splits in the branches.
- Points are classified using Hypercubes. Therefore two points being classified in the same cluster means there are no points of the other class in between them. At the same time, two normal clusters with anomalous points in between cannot be classified together as the hypercube containing them would be impure.

- As most models, decision trees suffer from the curse of dimensionality. However in our case the number
  of dimensions considered is low and the method would function correctly.
- A decision tree can fit any distribution. In our case we do not care about overfitting as we will not make predictions with it and only use it to cluster points.
- As stated previously, Decision trees create hypercubes. This in effect gives us intervals.
- We fit a decision tree on the data. This is done in  $O(mn \log(n))$  with m being the number of dimensions and n being the number of points. In practice, the algorithm runs slightly faster than the nearest neighbor one. Overall the time complexity is not too high.

This clustering method verifies all of the properties we needed making it a good candidate for our use. The construction of the decision tree case be done using Gini impurity or entropy. We will use entropy as this creates a link between the creation of the tree and the measure we are trying to compute. At each node of the tree, the best split is created by minimizing the entropy of the two leaves created. This suits us as we are trying to minimize the entropy of the clusters.

The use of hypercubes for clustering provides intervals for our explanation. However, this implies that some points that are adjacent will fall into multiple clusters. For example in Figure 7, the normal points will be split into two clusters. This can either be seen as a limitation or as a strength of this method. This penalizes the use of multiple dimensions when explaining the anomaly. Higher dimensions imply more cuts. This will prevent the incorporation of irrelevant features as these would either decrease the entropy or not change it. A byproduct of this is that getting a score of 1 for n features implies that there is a feature alone in that set that gets a score of 1. i.e let  $g_i$  be the entropy function on segment i.  $f_j$  is a feature for  $j \in [1, n]$ :

$$g_i([f_1, f_2, ..., f_n]) = 1 \Rightarrow \exists j \in [1, n], g_i([f_j]) = 1$$

This is due to the fact that getting an entropy score of 1 implies that the points are perfectly segmented, thus finding 2 clusters. Through the properties of decision trees, this implies a single cut along one dimension. Therefore the feature on which this cut was made is enough on its own to perfectly segment the points.

It is also possible to overcome this limitation by merging neighboring leaves into one cluster. The intervals would still be distinct but the clusters would be considered as one, lowering the overall entropy. This would be done by checking the neighboring leaves and merging them if they contain points of the same class. This would take place iteratively until no more merges are possible.

Decision trees also do not have any guarantees of finding the best clusters. The method is greedy and will only split the space along the best feature at each node. This can lead to suboptimal clustering as for example in Figure 11, the normal points on the right are split off early and are separated from the rest whereas they could have been clustered together. However, in practice the method works well and the results are satisfactory.

## **6.3** Incorporation into Exstream

We will now see how to incorporate any method mentioned above for computing entropy into Exstream. The intuitive way would be to find the set of features that obtains the best score through our method. However, as shown previously, finding such a set of features is not a submodular optimization problem. We therefore do not have an efficient way of finding the best set of features. There might be other ways of finding the best set of features using properties of the different functions. However, in this paper we will not explore these. And will limit ourselves to brute forcing the problem in two dimensions.

Another cause to motivate this decision of keeping the number of features considered to two is the complexity of our dataset and of the anomaly types. There are many traces where a single feature alone has perfect separation between the normal and anomalous points getting an entropy score of 1. Moreover, some traces contain multiple

such features that are not heavily correlated. In these cases it is useless to search for a combination of features as a single feature alone has a perfect score. It is also above the scope of this work to improve the selection of features to be returned in the above case. We will therefore rely on the same method as Exstream and when a feature gets a score above a certain threshold, we will not search in two dimensions and return all features with a score above that threshold subject that they are not correlated and that they are not false positives.

The returned format for the anomalous intervals will be the same as in the original exstream with  $I_i^i$  intervals:

$$((f_1 \in I_1^1 \lor f_1 \in I_2^1...) \land (f_2 \in I_1^2 \lor f_2 \in I_2^2...))...$$

The case where no single feature is enough to perfectly segment the normal and anomalous points, we will brute force the problem in two dimensions. After removing false positive features, we will compute the entropy for all pairs of features and return the pair that has the best score. Considering there are initially 237 features in the dataset, in the worst case (when there are not any false positive features), we will be computing the entropy for  $\binom{237}{2} = \frac{237 \times 236}{2} = 27966$  pairs of features. This is a substantial amount of computation especially considering the complexity for each pair is  $O(n \log(n))$ .

When using the decision tree method, we will obtain hypercubes as intervals. These hypercubes can be defined as the conjunction of intervals in each feature. Therefore the return format will be the following:

$$(f_1 \in I_1^1 \land f_2 \in I_1^2) \lor (f_1 \in I_2^1 \land f_2 \in I_2^2)...$$

This inversion between  $\land$  and  $\lor$  provides far more information about specific intervals where points are anomalous. This is due to the absence of negations in the Conjunctive normal form. It is also possible to return multiple pairs as explanation. These would be linked using  $\land$  clauses.

The new output format will be the following:

```
{
  "Trace":{
    "Anomaly number and type": {
      "feature_to_importance": {
          "feature_1, feature_2": score
      },
      "feature_to_intervals":{
           "feature_1, feature_2":[
               [
                   [
                        feature_1 start,
                        feature_2 start
                   ],
                   Γ
                        feature 1 end,
                        feature 2 end
                   ],
                   include starts,
                   include ends
               ],
          ]
      }
    },
    . . .
```

```
} ...
```

#### 6.4 Results

Methods	Time	Length	Instability	Discordance	ED1 F1 Score	ED2 F1 Score
Exstream Tree 80%	15.6	2.9	1.63	5.76	0.997	0.94
Exstream Tree 90%	29.3	1.9	1.91	6.91	0.997	0.96
Exstream Tree 95%	55.2	1.5	2.22	8.43	0.997	0.97

Table 3: Results of evaluating the new method

This new method is evaluated with the same metrics as the original Exstream and on the same traces. We also evaluate the method while removing the first and last 10%, 5% and 2.5% points of each normal and anomalous instance. The results are presented in Table 4.

An improvement in the length of the explanation along with a worsened discordance can be observed. This is due to the way the output is handled. A pair of features  $(f_1, f_2)$  is considered a single conjunction. Therefore, in disjunctive normal form it is counted as a single feature. Therefore whenever any single feature does not get a high enough score, the length of the explanation will be one. One the other hand, each pair is considered a unique new feature. This is why the discordance is higher.

The F1 score in ED1 and ED2 settings are both improved significantly using this new method. This is due to the fact that the method is able to capture the interaction between features and express intervals in a more precise way. This result is very satisfactory as it shows the benefit of using a multi dimensional entropy measure. This benefit can also be observed on traces that require multiple features in their explanation such as in Figure 12. Each feature individually has a low separation score and would not be selected as a satisfactory explanation. However both features together obtain a score of 0.75 and are chosen

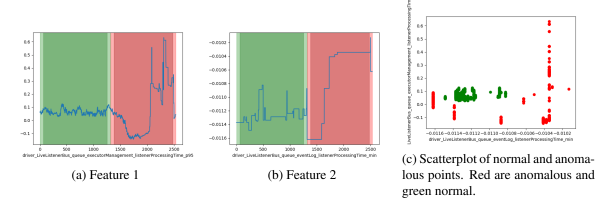


Figure 12: Example of Explanation using two features. Trace 8\_4\_1000000\_10\_2\_15\_77 3T4

## 7 Conclusion

We have first presented a method to improve the sampling of points. This method is restricted by the existence of domains in our dataset. We therefore used reconstructed points. This method does not provide the expected results. We then presented a method to generalize the entropy measure to higher dimensions. We showed that

Methods	Time	Length	Instability	Discordance	ED1 F1 Score	ED2 F1 Score
Exstream Bins	140.7	4.3	1.5	6.16	0.79	0.72
Exstream 100%	1.5	1.6	2.26	7.71	0.88	0.87
Exstream 95%	1.35	2.1	2.08	6.59	0.87	0.93
Exstream 90%	1.2	2.4	1.88	6.04	0.88	0.92
Exstream 80%	1.1	3.4	2.0	5.5	0.88	0.90
Exstream Tree 95%	55.2	1.5	2.22	8.43	0.997	0.97
Exstream Tree 90%	29.3	1.9	1.91	6.91	0.997	0.96
Exstream Tree 80%	15.6	2.9	1.63	5.76	0.997	0.94
Exstream Reconstruction 100%	5.2	1.6	2.77	8.56	0.79	0.81
Exstream Reconstruction 95%	4.3	1.8	3.17	7.08	0.78	0.87
Exstream Reconstruction 90%	4.3	2.2	2.87	5.59	0.79	0.86
Exstream Reconstruction 80%	4.2	2.6	2.04	5.65	0.79	0.84

Table 4: Compilation of all the results.

the submodular property does not hold for our data. Three methods to cluster points in higher dimensions were then introduced. The decision tree method was chosen as it fits all the requirements we had. We then showed how to incorporate this method into Exstream. The results show a significant improvement in the F1 score. The length of the explanation is also improved. The discordance is worsened but this is due to the way the output is handled. The methods introduced were evaluated while removing 0%, 2.5%, 5% and 10% of points on the extremities of the intervals. The results are consistent and show a significant improvement.

Overall, even though our first attempt in improving the sample of points was not successful, the second method of generalizing Exstream in higher dimensions was. The results are very satisfactory and show that the method is able to capture the interaction between features and provide a more precise explanation.

## 8 Further works

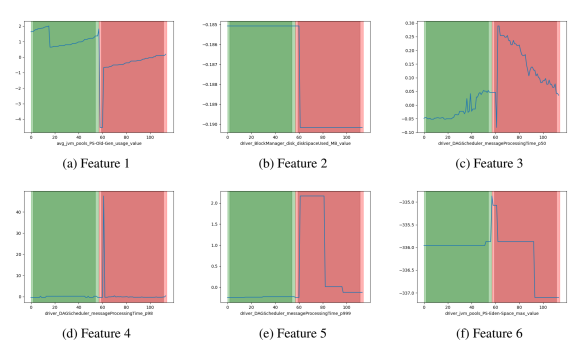


Figure 13: Example of Explanation using six features. Trace 5\_5\_1500000\_5\_3\_15\_92 1T5

The length of the explanations are greater than in the original Exstream. There are still explanations with 5 or 6 features. For example Figure 13, There are 6 features with each a score over 0.9. This number of features is too much in an explanation and some features might be correlated with each other while not being detected in the restricted sample which we have. Moreover, another limitation is that if features  $f_1$  and  $f_2$  are correlated, if  $f_1$  is returned as explanation in one anomaly and  $f_2$  in another, the discordance will increase. We could therefore find a way to always return the same features when they are correlated. This motivates the grouping of *globally* correlated features whith a set rule on which to return as explanation.

The method we introduced to sample using the latent space is not satisfactory. The features correlated with time are thus still not detected or removed. This is something to work on and improve in the future.

Our method using trees is currently limited to two dimensions. This is enough on our dataset that is simple. However, in a more complex dataset, the method would need to be generalized to more dimensions. The limitation is due to the way we search for the best set of features in a brute force manner. This is inefficient and limits the search in higher dimensions. Finding a faster way would enable the method to be used on more complex datasets.

A better handling of metrics for couples of features would also be needed. The discordance and length of explanation will need to be adapted to the new output format. Moreover, including a metric that includes the number of intervals being returned would be useful. This would allow for a better evaluation of the method.

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# A Appendix

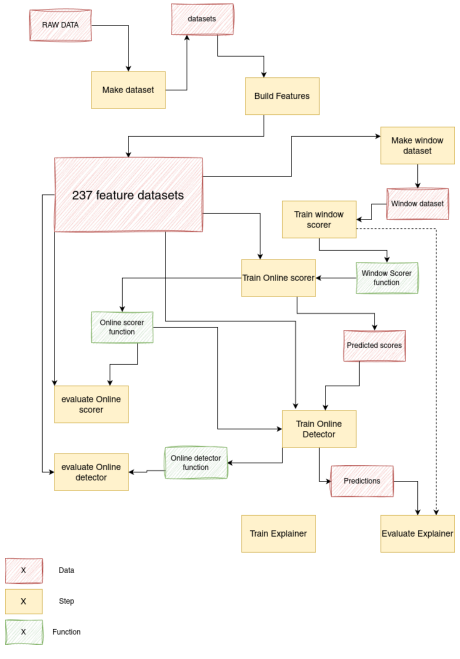


Figure 14: Pipeline of Exathlon

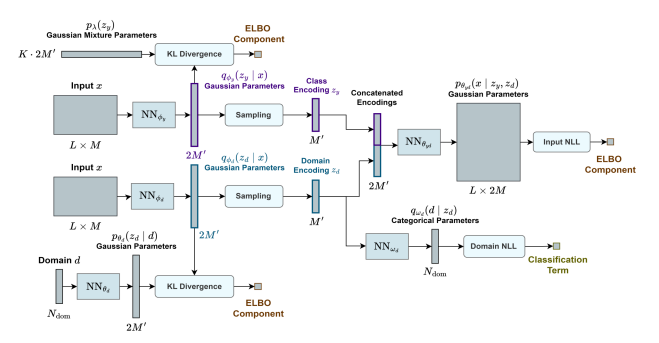


Figure 15: Divad model architecture