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STATISTICAL METHODS IN AI: RARE EVENT LEARNING USING ASSOCIATIVE RULES AND HIGHER-ORDER STATISTICS

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ABSTRACT:

Rare event learning has not been actively researched since lately due to the unavailability of algorithms which deal with big samples. The research addresses spatio-temporal streams from multi-resolution sensors to find actionable items from a perspective of real-time algorithms. This computing framework is independent of the number of input samples, application domain, labelled or label-less streams. A sampling overlap algorithm such as Brooks-Iyengar is used for dealing with noisy sensor streams. We extend the existing noise pre-processing algorithms using Data-Cleaning trees. Pre-processing using ensemble of trees using bagging and multi-target regression showed robustness to random noise and missing data. As spatio-temporal streams are highly statistically correlated, we prove that a temporal window based sampling from sensor data streams converges after n samples using Hoeffding bounds. Which can be used for fast prediction of new samples in real-time. The Data-cleaning tree model uses a nonparametric node splitting technique, which can be learned in an iterative way which scales linearly in memory consumption for any size input stream. The improved task based ensemble extraction is compared with non-linear computation models using various SVM kernels for speed and accuracy. We show using empirical datasets the explicit rule learning computation is linear in time and is only dependent on the number of leaves present in the tree ensemble. The use of unpruned trees (t) in our proposed ensemble always yields minimum number (m) of leafs keeping pre-processing computation to n x t log m compared to N^2 for Gram Matrix. We also show that the task based feature induction yields higher Quality of Data (QoD) in the feature space compared to kernel methods using Gram Matrix.

I. INTRODUCTION

We explore powerful label and label-less mechanisms for automatic rule extraction from streams which have rare event embedded in them using in-memory feature engineering. Data-intensive computing needs to process fast input streams with very less memory and without storing the samples. We compare two task induced feature induction models, explicit and non-linear kernels: first method uses task induced similarity metric to learn rules from the ensembles and, the second method learns the Gram matrix of dimensional R^d using different kernel window widths. The explicit representation uses Random Forest ensembles to generate the overlap proximity matrix. The similarity counts combines the k-similar trajectory paths ending up in the same terminal node. More trees are grown and added to the Random Forest till predictive accuracy is significantly higher. When learning the data driven Gram matrix the kernel computation model uses matrix factorization. At each iteration the best row of features are determined using the category labels to reduce the rank of the final kernel matrix. Finally a fast approximate in-memory data stream model using Hoeffding tree is induced to combine the computational and predictive performance of both the models.

Just as the web browser brought us click-stream data, GPS enabled mobile phone has created huge amounts of geo-tagged streams. Understanding of geo-intelligence and data mining actionable events from random data has not been studied until recently due to lack of scalable algorithms and computational models. The recent reference to "Data bonanza" (Krause, A., 2012c) in the knowledge discovery literature can be attributed to recent datasets and how it is impacting wireless data driven applications. New streams such as trajectories generated due to the proliferation of GPS enabled cabs (Zheng et al., 2009) in large metros and increase availability of Wi-Fi hotspots. The computation models needs to learn trajectories which can give insight to the behaviours of the users for designing future infrastructures. Just as designing a fast prefetch cache from web users’ click stream, the tree model needs to capture the branching of the paths by approximating from sparse segments of moving data in motion. When using a tree to represent the trajectories the memory savings could be enormous when compared to the size of the original trajectories due to potential overlaps. Electronically generated streams do not contain labeled data and thus not suitable for supervised algorithms. We propose a powerful overlap function (Prasad et al., 1991) metric to identify the actionable events. The stream pre-processing algorithm is resilient to noise and uses instance based learning and at the same time, works in unsupervised settings (it can have labels). The overlap function determines a proximity count induced by the tree ensembles as shown in Fig 1(a).

Non-linear learning methods are associated with Gram matrix and kernel methods. The kernel method uses a computational shortcut to compute its feature space. The resulting higher-order feature maps can be optimized for in-memory analytics and fast predictive models (Agrawal et al., 1993). We compare our explicit ensemble overlap feature induction framework with standard non-linear kernel function learning. Even though Gram matrix can be represent any input data and can learn the features by using higher-order polynomials we show the the number of support vectors (SV) do not scale well with large n. Computation of kernel methods are independent of the feature and hence does not have the “curse of dimensionality”, but still suffers from the “curse of support”.

This contribution has been peer-reviewed. The double-blind peer-review was conducted on the basis of the full paper.
We have successfully used Hoeffding bounds to guarantee that the baseline model is within a small probability $\delta$ from its true mean. We extend the Hoeffding principle to Data Streams (Bifet, 2009a), which have a property of infinite input size. For the in-memory model (Brooks and Iyengar, 1996) (Iyer et al. 2013) (Breiman, 2001) to fit the requirements to handle data in motion, the above rule extraction models needs to address the following resource constraints. To store sufficient statistics which are needed to model the stream that are to be held within node splits of the tree in-memory. Hoeffding bounds (Bifet and Kirkby, 2009) states that the estimated mean does not change very much after sufficient samples ($n$) are seen and this is independent of the distribution. To guarantee sufficient statistics for on-line classification and to guarantee its true mean we use Hoeffding tree. We also show that Hoeffsizing tree is able to compute sufficient statistics in memory and achieve a base line accuracy compared to the above two kernel methods tested with large data streams.

In the cases discussed so we assume the datasets have known events even though learning from them efficiently and its scaling have been satisfactorily addressed. Still there are rare events which are weekly labelled (due to unbalanced support) as they are buried inside noise and outlier present in large streams. A weekly labelled event (buried events) are illustrated in Fig. 1(b,c) which can occur due to big sample sizes. Data models for large data streams have successfully used to correct such outliers by using Bonferroni principle (Rajaraman and Ullman, 2011) to address false alarms. We further explore to mine actionable patterns of practical significance using power of SVM in the following ways: (i) One-class SVM for outliers and, (ii) soft-margin SVM for novel patterns.

The rest of the paper is organized into two major sections (i) low-level data-cleaning and (ii) task based feature extraction and statistical model building. Section II, discusses why rare event learning is an NP hard problem and seeks to use existing SVM to solve them. Section III, surveys related work in spatio-temporal domain to explore low-level data-cleaning reliability. Section IV, introduces how algorithms are applied to spatio-temporal data with a task based approach. As we will extend our low-level model with a computation module based on our original overlap function which is non-parametric. In Section V and VI we introduces kernel method computations and algorithms to build robust statistical models to deal with dynamic data. Section VII demonstrates the results for spatio-temporal datasets with in-memory models and compares preliminary streaming results with existing standard machine learning libraries. Section VII summaries and concludes the work.

II. RARE EVENT LEARNING

An example from weather dataset contains humidity, rain, temperature, and wind to predict the occurrence of fog, a rare event (Garimella and Iyer 2007), when rain attribute value happens to be false. We like to further understand rare events (Lin et al., 2013) and its prediction for label and label-less streams using similarity measures. We already know that non-parametric methods such as k-means clustering needs to know the number of clusters a priori. To overcome this limitation we use density based approaches in the feature space, which is explained in Section V. As an introductory example to illustrate this concept by using one-class SVM as illustrated in Fig. 1(b). We estimate the single class features and infer the properties of a rare event without assuming the number of clusters a priori. When using pattern recognition using Support Vector Machine (SVM) (Cortes and Vapnik, 1995) to classify rare events we use One-class and soft-margins SVMs as discussed in this section. The Bonferroni principle states that if the number of events in practice are know to occur can be calculated then the outliers can detected and avoided in random data sampling.

Looking at the same problem in terms of a probability density function (pdf), given a set of data embedded in a space, the problem of finding the smallest hypersphere containing a specified non-trivial fraction of the data is unfortunately NP-hard [kernel methods]. Hence, there are no known algorithms to solve this problem exactly. It can, however, be solved exactly for the case when the hypersphere is required to include all of the data. So we first consider the cases when all the data are available.

A. One-Class SVM

When estimating big samples sizes (Lin et al., 2013), one-class SVM can be used in a unsupervised learning setting. We generate two types of random data: one with ‘normal’ novel patterns with a fixed mean $\mu$ and, other with ‘abnormal’ novel patterns which are uniformly distributed. The effectiveness of the algorithms is illustrated in Fig. 1(c). The results shows that the SVM is unable to perfectly classify for the underlying unknown distribution as it has some outliers. Use of robust covariance estimator which assumes the data are Gaussian. The plot in Fig. 1(c) has no errors as it applies the Bonferroni principle to further filter the practical limits when applied to big samples.

B. Hinge-loss Cost Function with Soft Margins

When the data is completely random and cannot be completely separated using SVM hyperplane. The hidden events present which may be hard to classify needs and needs to be cross validated by splitting the training set into 80-20% splits and a soft-margin. The soft-margin is an automatic way to rejects a hard to classify sample by using cost function: the learning algorithm needs to balance the ratio of false positives occurring during testing (Lin et al., 2013) and help lower them. When a particular pattern has probability close to 0.5 in the case of a binary classification, then use of soft-margin SVM combined with a penalty function is more efficient to learn the hidden novel patterns we discussed earlier using the complete data. The SVM model can use the soft-margin tuning variables $C$ and $\gamma$ for their best fit (Hu and Hao, 2012; Iyer et al., 2012). The key is to tune the soft-margin hyperplane with the variable ($\xi$) for standard SVM hard hyperplane (Cortes and Vapnik, 1995). SVM equation as shown below:

$$y_i (x_i^T w + b) \geq 1 - \xi_i \quad \xi_i \geq 0$$

We define a convex cost-function such that

$$y_i (x_i^T w + b) \geq 1 - \xi_i \quad \gamma (\xi_i - 1)$$

The new estimate using reject option further helps to disambiguate hard to classify samples.

How to compute and estimate large sensor data streams to mine rare events:

- Feature Engineering - “Data driven learning”
  - Instance based label-less feature task-based induction
- Streams Models - “Data in motion”
  - Efficient in-memory sufficient statistics
- Rare Events - “Statistical effects of big sample sizes”
One-class novel pattern detector using Bonferroni correction
- Soft margin SVM using loss function to find hard to classify patterns

Current work focuses on feature extraction using explicit learning algorithms. Events can be learned using labels or sometimes hard to classify datasets. SVM based kernels can disambiguate hard to classify datasets using soft-margins. Most of the learning algorithms still have the training phase which make them unsuitable for real-time analytics. In the domain of sensor networks, distributed measurements have been successfully calibrated (Prasad et al., 1991) with fault tolerant (Prasad et al., 1993) algorithms to avoid false alarms. The techniques used in feature extraction and real-time fault tolerance does not address rare event learning directly. As rare events can be patterns, which may not necessarily be induced by basic feature occurring in the dataset. The ability to detect rare events is of at most importance, and needs the ability to predict such occurrence without training from past data. The proposed AI based rule learning technique allows to bridge the gap between feature extraction and real-time rare event mining.

III. RELATED WORK

It is a known fact that when dealing with big samples the p-value (Lin et al., 2013) linearly decreases causing false events detection even though they are not significant. In a similar way the learning algorithms have to deal with "curse of dimensionally" with respect to attributes and "curse of support" (Kar and Karnick., 2012) when determining support vectors using SVMS. There is a related problem of missing values in the dataset and how the in-memory model addresses it for a particular task. The work by Vens and Costa (Vens and Costa., 2011) on Random Forest a task based feature induction, deals with statistical and missing value sub-problems. A similar work by Bach and Jordan (Bach and Jordan., 2005) to compute a predictive low-rank decomposition for task based feature induction using kernel methods have addressed the same issue. In this work we use both methodologies to study the statistical properties and the quality of data related to the fitness needed in mobile data driven applications.

The parallel computation of large tree ensembles has been recently addressed by Breiman and Adele Cutler (Liaw and Wiener, 2002). Which combines individual tree representation (Chang et al., 2006) with a distributed feature table (big-memory scaling). New frameworks such as R Programming Studio, Mapreduce, and Spark MLlib, now support distributed processing just beyond < key, val > - pairs. The pre-processing language semantics support in the latest version of Spark further allows efficient feature extraction in real-time. Which enables existing batch algorithm to be ported to real-time sensor data stream environment effortlessly.

When using kernel matrix to approximate higher-dimensional feature space to lower-dimension space, further allows to address the huge number of SV’s. The optimization in (Kar and Karnick., 2012) helps to linearly scale (refer to detailed proof) with the size of input. The scalability of computation model is fast but it does not help the predictive performance when features are independent of their attributes. When modeling data of a user’s trajectory or other forms of synthetic data, the features cannot be generalized by using the basic patterns and may not form regular clusters. So one cannot use non-parametric clustering when the number of clusters are not known a priori. So we argue the use of explicit overlap function (Prasad et al., 1991) feature induction in these cases works well. We further extend sensor data streams for fast predictive overlap models which can handle real-time constraints.

On-line processing of noisy sensor measurements has been extended to distributed framework by Brooks-Iyengar (Brooks and Iyengar 1996). This seminal work balances precision and accuracy to estimate the non-faulty (Iyer et al., 2008) sensor’s confidence intervals. Real-time algorithm has been extended using Byzantine fault-tolerance principle to mitigate faulty sensors (Brooks and Iyengar., 1996), which are likely to trigger false alarms. The power-aware distributed computation is due to its simplicity and the protocol keeps communication overhead to a minimum using in-network processing and with minimum centralized coordination.

IV. AUTOMATIC RULE EXTRACTION

Computational data sciences has massive amounts of datasets generated from remote sensing and, periodic stream updates from several sites spread across many geographical regions. Much of the publicly available datasets have not been analyzed for the lack of computationally viable algorithms. Stream analytics cannot scale due to the following reasons: the known p-value statistical problem (Lin2013c) with large scale datasets; the statistical significance of a common feature increases linearly with number of samples.

A. Calculating Higher Order Side Information

In this paper we are not only interested in rare event prediction but also the temporal rules (slow moving in time) which attributes to it. Important patterns occurring many times in the data if there is enough support then it becomes a rule. Due to this large size datasets needs to estimate rare events in practice differently. One such estimation was suggested by the Bonferronis Principle. Which states that models ability to mine data for features that are not sufficiently rare in practice. Scientist in this domain have used raw sensor attributes which when combined with learned coefficients help to form linear ranking index functions such as Fire Weather Index (FWI) estimation. Using FWI ranking the streams from the weather model can track the time-series sensor stream and predict rare events before than can happen. Due to the number of datasets available for rare events being few for training, we propose an automatic methods to detect such events in a timely fashion. When mining massive datasets the rule extractor is a correlated side-information (Iyer et al., 2013b) present in the target concepts. Some of the feature extraction are discussed with sensor data (Table I) and ground truth examples (Zheng et al., 2009).

We generalize a predictive functions which has the following rules:

- Descriptive inputs att1, att2 → side-information

The predictive rule suggests that the inputs of the dataset are highly correlated to a few target attributes, which then uses as objective measures by tracking the side-information - especially in the case of unlabeled streams. As an example we use the Landsat dataset (Newman and Asuncion 2007) from University of Irvine, Machine learning repository. Landsat dataset attributes have the following attributes: channel-R; channel-G; Channel-invisible1, Channel-invisible1. For example, the extracted rule for our e.g. dataset is:

channel-R; channel-G → channel_invisible1, channel_invisible2

The input output mapping suggests that their exist a strong relation between visible channels (Statlog) (Newman and Asuncion,
2007) and the observed values of invisible channels, because the measured scenes are one and the same for the visible band and the satellites invisible band. There was no significant deviation in the normalized Root Mean Square Error (RMSE) values during training and testing, suggesting the use of the side-information are ideal in practice, when using regression trees.

The side-information concept can be extended (Iyer et al., 2013b) to rare events such as forest fire by computing the temporal FWI of that region. The above rule can then be written for FWI (Iyer and Iyengar et al., 2011) as:

\[
\text{temp}, \text{humidity}, \text{wind}, \text{precipitation}, \rightarrow \text{FWI scalar } (0-80)
\]

B. Performing Clustering Based On Data Cleaning Trees

The ensemble model which is illustrated in Fig. 1(a) uses Random Forest where the node splits are based on clustering target variables. The target variables help in finding unique clusters as discussed in this section. The prototype function (Vens and Costa 2011) used for clustering and node splitting helps to estimate the model density thresholds for the current window. The induction of such a tree with actionable events results in extracting their local temporal rules for further labeling and on-line event dash boarding.

V. TASK BASED DISTRIBUTED FEATURE ENGINEERING

Feature engineering as a domain, focuses on learning from data streams without a priori knowledge of the underlying process or the number of clusters. Source of the generating process are unknown and random and due to their high variance its validation can be only done solely by use of ground truth. Ground truth can be in the form of past events recorded by weather station logs and verifiable timestamps overlapping the same period as of the training dataset. Assuming that rare events are present in the data we focus on various task-based feature induction as described in Fig. 1(a). We discuss the advantages of task based learning compared to learning the kernel functions and how it handles missing values. The common metric used by the learning functions are non-parametric (distance and similarity) induced in the feature space, but we are interested in learning the fusion of spatio-temporal features. The task based explicit feature metric representation has other advantages. The computational advantage is that it does not need to learn the pair-wise distance and need not store quadruple amount of information such as the Gram Matrix. A tree based representation naturally allows to deal with missing values, compared to existing statistical methods.

We illustrate the stream model for both cases using static data-driven algorithms and mobile wireless application testbed as shown in Fig. 2(a). There are two forms of stream quality (Iyer et al., 2013a) non-parametric ranking for random streams which are shown in Fig. 2: kappa score for missing values and, finding actionable events of interest. Kappa score (Iyer 2013a) helps to compare how the learning algorithm perform using euclidean distance measure when dealing with spatial measurement. For temporal end-to-end trajectory streams, an explicit similarity measure is learned from Random Forest ensemble as shown in Fig 1(a). Which allows to learn similar trajectories during model training. Both these methods are cross-validated by the ground truth for selecting and refining the models for noisy, static and mobile streams. When growing the Random Forest ensembles the tree splitting at the root node is selected along one of these \(v\) attributes to minimize the number of misclassified training points if a majority vote is used in each node. The procedure is repeated until every node is pure and assumes the following assumptions such as \(n\) training samples and the samples are saved and can be read back many times. The current assumptions cannot be applied for real-time algorithms which cannot store stream samples and read the samples only once. Comparing the real-time Brooks-Iyengar’s algorithm the metric used for parameters’ confidence interval approximation are the weighted average of precision and its accuracy (Brooks and Iyengar 1996) over all the sensor measurements. As our task based distributed feature engineering needs to scale in a real-time window. The distributed framework need to extend the current feature splitting methodology to an iterative learning model, which can handle data in motion while computing in-memory.

Summary of the iterative stream learning model:

- Uses Hoeffding bounds for statistical sampling
- Uses task-induced similarity measure metric
- Handles missing value by randomization
- Distributed in-memory features representation using big-table
- Real-time computation guarantees to process fast streams

The main contribution of the paper is its evolution of static sensor network algorithms to address dynamic nature of spatio-temporal domain using Data Streams model. We divide the contribution into two parts the low-level overlap function and two the in-memory statistical computational model representation of actionable events. We show that our overlap algorithm work as a proof of concept using a simple one-dimensional case. Further experiments with
two-dimensional GPS datasets provided high accuracy in predicting actionable events with large datasets. Hoeffding tree and AI (Garimella and Iyer 2007) based ANOVA kernel computational model optimize resource requirements by one-tenth of other well proven models such as Naive Bayes and non-linear kernels.

VI. METHOD

Lower-level sensor measurements task are prone to errors due to noise and faulty (Iyer et al., 2008) sensors. Incorrect results can be obtained as the upper level Gram Matrix relies on tuning few parameters. Higher level data mining tasks should be descriptive that summarizes the underlying relationships in data. To design a data-driven application the low-level should be task driven that is independently supported by higher-level information. Our higher-order feature induction approach uses learning as follows: ensemble overlap similarity, Gram matrix using kernel methods, and moving window using an iterative Hoeffding tree model. Related work shows that kernel methods have higher predictive performance but does not allow to intuitively understand the rules extracted from them given the original input space. As we are analyzing data which has temporal attributes it is equally important to extract the thresholds which support them. Due to mobility based location context requirements, our emphasis is based on natural or explicit rule learning and does not rely on high-dimensional Gram matrix. Once the rule extraction process synchronizes with the low-level mining task, one can compare the performance over a moving window using on-line Hoeffding tree. The implementation methods discussed below addresses the following memory requirements for any data stream type. How learning natural rules from data can be learned in a task driven model.

A. Algorithm

Real-time algorithms needs to maintain sufficient statistics and the tree structure in memory. The incremental tree updates due to the changing statistics are performed by splitting using the best attribute. When referring to kernel methods an \( N \times N \) Gram matrix or an optimized lower rank is stored in memory.

1) Quality of Data and Big Matrix Representation

The use of tree structure allows efficient in-memory model and at the same time we like to extend the idea of instance based feature induction in our mobile applications. Random Forest ensemble builds many trees which prevents model over-fitting. The implementation of the distributed big-table (Chang et al., 2006) is by allowing the ensemble growth of the current working set to be more than the available fast memory. Internally the big table is implement as a shared distributed table across worker nodes. After the ensemble parameters are tuned and built with sufficient number of trees then the model is tested. We use the training set \((n)\) and the out-of-bag samples together, and traverse down the trees \((t)\) taking one input sample at a time. In the case of similar samples they will end up in the same terminal nodes \((m-leaves)\) using \(l(t \log n)\) steps. For those cases the model increases the proximity count by one. To get a number between \((0-1)\), the model divides the proximity counts for each case by the number of trees in the ensemble. The elements in the big-matrix will estimate the number of times instances \([m \text{ and } n]\) end up in the same node. This sparse measure is useful for upper-level trajectory learning as shown in Fig. 2(b) and other higher task based rule extractions from random and synthetic datasets. Computation overhead for the calculation of Gram matrix can have a worst case of \(N \times N\). The proximity computation of a Random Forest ensemble is always efficient. The framework can handle any number of attributes when performing proximity calculation in parallel with less than ten classes. We show the worst case \(N \times N\) complexity of calculating the primal Gram matrix can be efficiently computed using via its dual representation. The data-cleaning dual framework’s computation are only dependent on the number of spatio-temporal features and is given by \(O(N)\) (Shawe-Taylor and Cristianini, 2004) where \(f\) are linearly independent attributes found using kernel transforms.

2) Data in Motion

When estimating events from big sample data which can run into 10,000 or more observations. The p-value is inversely proportional to number of samples, the value goes quickly to zero as \(n\) increases. Relying on p-value estimation alone cannot find events of practical significance we use a confidence parameter \(\delta\) to determine the node splits. Due to model building with fixed size datasets is not possible with limited memory, we propose a real-time windowed data stream model using Hoeffding tree bounds (similar approach to Brook-Iyengar algorithm):

1. Data Streams cannot be stored and an emphasis of in-memory model is assumed to have low processing latency
2. Sufficient statistics are kept which are essentially a summary of the data and requires very small memory.
3. The distribution of the stream is expected to change over time which triggers recalculation of the statistical summaries.
The in-memory iterative model uses a window \( W \) which is partitioned into \( W_0 \cdot W_1 \) with lengths \( n_0, n_1 \), such that \( n = n_0 + n_1 \). To determine sufficient statistics for our iterative model we define \( \epsilon_{cut} \) which minimizes the error \( (1 - \delta) \). To further guarantee real-time performance we rely on harmonic mean defined by:

\[
m = \frac{1}{\frac{1}{n_0} + \frac{1}{n_1}}
\]

\[
\delta = \frac{1}{2m} \log \frac{4}{\delta}
\]

There are two cases which can be used for performance guarantees that estimate the change in distribution using a statistical test.

**Theorem 1.** The performance of our model depends upon on event rate processing. The data stream model performs in incremental time steps and we have:

1. **False positive rate bounds.** If the optimal window size is chosen for the Data Stream application then \( \mu \) remains constant within the windows \( W \). The probability that ADWIN will shrink the windows at this time step is at most \( \delta \).

2. **False negative rate bounds.** When the optimal window size \( W \) is not known a priori for the Data Stream applications some events may be ignored due to the overlapping settings \( \mu_0 - \mu_1 > 2\epsilon_{cut} \). Then with probability \( 1 - \delta \), \( W \) shrinks to \( W_1 \) in the next time step.

**Proof.** Now assume \( |\mu_0 - \mu_1| > 2\epsilon_{cut} \). We want to show that \( \Pr[|\mu_0 - \mu_1| > 2\epsilon_{cut}] \leq \delta \), which means that with probability at least \( 1 - \delta \) change is detected and the algorithm cuts \( W \) to \( W_1 \). AS before, for any \( k \in (0, 1) \), we can decompose \( |\mu_0 - \mu_1| \geq \epsilon_{cut} \) as

\[
\Pr[|\mu_0 - \mu_1| \geq \epsilon_{cut}] \leq \Pr[|\mu_0 - \mu_1| \geq |\mu_1 - \mu_0|] + \Pr[|\mu_1 - \mu_0| \geq |(1 - k)\epsilon_{cut}|]
\]

To see the first inequality, observe that if \( |\mu_0 - \mu_1| \geq \epsilon_{cut} \), and \( |\mu_0 - \mu_1| \leq (1 - k)\epsilon_{cut} \) hold, by the triangle inequality we have \( |\mu_0 - \mu_1| \leq |\mu_0 - \mu_1 - \mu_0 + \mu_1 - \mu_1 + (1 - k)\epsilon_{cut}| \leq |\mu_0 - \mu_1 - \mu_1| + |\mu_0 - \mu_1| + \epsilon_{cut} \leq 2\epsilon_{cut} \), contradicting the hypothesis. Using the Hoeffding bound, we have then

\[
\Pr[|\mu_0 - \mu_1| \geq \epsilon_{cut}] \leq 2 \exp(-2(1 - k)\epsilon_{cut}^2 n_1) + 2 \exp(-2((1 - k)\epsilon_{cut})^2 n_1)
\]

Now, choose \( k \) as before to make both terms equal. By the calculations in Part 1 we have

\[
\Pr[|\mu_0 - \mu_1| \geq \epsilon_{cut}] \leq 4 \exp(2\epsilon_{cut}^2 n_1) \leq \frac{\delta}{n} \leq \delta
\] as desired.

The windows management model assumes for a stable distribution, the error rate should decrease over time. We propose an algorithm ADWIN (Bület and Kirkbride, 2009), which uses a dynamic window to maintain in-memory statistics. The procedure is presented in Algorithm 1. At every step ADWIN outputs the value \( \mu_w \) which is an approximate of the value \( \mu_w \). Current models when mining static datasets the algorithm it able to revisit the input data many times. In the stream model this is not possible when dealing with fast stream processing, as not all the data needed can be held in memory. Moreover the distribution of the data source can change over time for the evolving streams. Due to this the existing mining techniques cannot be applied with the single pass approach. When the working set is a tree structure to represent data model, a single instance of a Very Fast Decision Tree (VFDT) is preferred. The memory overhead is overcome by using Hoeffding bounds. Hoeffding Tree which uses...
this technique for the representing the model does need a training phase; instead it learns incrementally and can be used to test new samples at any given time to keep up with high speed of stream processing. Hoeffding bound states that the true value of the mean $\mu$, of a random variable of range $R$, can be practically approximated for big samples after $n$ independent observations by more than (with a low probability of error):

$$
\epsilon = \sqrt{\frac{R^2 \log(1/\delta)}{2n}}
$$

The estimation using this bound is useful as it is independent of the distribution. Maintaining sufficient statistics are explained in (line 4-6) of Algorithm 1. Node splitting and how the tree grows are show in line (12-15) in Algorithm. The Hoeffding bounds allows to compute the true mean with in a reasonable margin of error. The growth of the Hoeffding tree can be show in Algorithm 1, is compatible with $p@0.01$ (Lin et al., 2013) which is desired for big sample data sizes.

Algorithm 4 Hoeffding Window Tree(Stream $\delta$)

1: procedure HWTreeGrow((x,y),HT, $\delta$)
2: \hspace{1em} Sort $(x,y)$ to leaf $l$ using HT
3: \hspace{1em} Update estimators $A_{ijk}$
4: \hspace{1em} at leaf $l$ and nodes traversed in the sort
5: \hspace{1em} if $l$ in $T_{alt}$ then
6: \hspace{1em} \hspace{1em} HWTreeGrow((x,y),T_{alt}, $\delta$)
7: \hspace{1em} end if
8: \hspace{1em} Compute G for each attribute
9: \hspace{1em} Evaluate condition for splitting leaf $l$
10: \hspace{1em} if $\text{Best Attr.}(\text{G[2nd best]} > \epsilon(\delta, ...)$ then
11: \hspace{1em} \hspace{1em} Split leaf on best attribute
12: \hspace{1em} end if
13: \hspace{1em} for each branch of the split do
14: \hspace{1em} \hspace{1em} Start new leaf
15: \hspace{1em} \hspace{1em} and initialize estimators
16: \hspace{1em} end for
17: \hspace{1em} if one change detector has detected change then
18: \hspace{1em} \hspace{1em} Create an alternate subtree $T_{alt}$ at leaf $l$ if there is none
19: \hspace{1em} end if
20: \hspace{1em} if existing alternate tree $T_{alt}$ is more accurate then
21: \hspace{1em} \hspace{1em} then replace current node $l$ with alternate tree $T_{alt}$
22: end if
23: end procedure

Algorithm 3 Hoeffding Window Tree(Stream $\delta$)

1: procedure Euclid($a, b$)
2: \hspace{1em} Let HT be a tree with a single leaf(root)
3: \hspace{1em} Init estimators $A_{ijk}$ at root
4: \hspace{1em} for example(x,y) in Stream do
5: \hspace{1em} \hspace{1em} HWTreeGrow((x,y),HT, $\delta$)
6: end for
7: end procedure

3) Cholesky Decomposition with Task based Side Information - CSI

Kernel based data driven learning has become popular as there are publicly available software libraries which support to solve complex matrix multiplication, pseudo inverse and convex optimizations needed in vector modeling (Kar and Karnick., 2012). One of the popular forms of factorization is Cholesky decomposition (Bach and Jordan., 2005). Since the Cholesky decomposition is unique, performing a Cholesky decomposition of the kernel matrix is equivalent to performing GramSchmidt orthonormalisation in the feature space, and hence we can view Cholesky decomposition as the dual implementation of the GramSchmidt orthonormalisation. The normal Gram matrix obtained from this is very large and in the order of $N^2$. The general optimization does not include side information which are in the form of class labels or response variables. Our previous discussion on using a sparse representation instead of full Gram matrix (Bach and Jordan., 2005) can be realized by this method. While the optimization steps pivots are generated during factorization an additional optimizations is used by CSI to include the remaining label information. Which allows to generate sparse and most of the time lower ranked matrices compared to the the general Cholesky methods. CSI optimization methods allows to reduce size of the final matrix allowing fast in-memory kernel models without loss of prediction accuracy. Similar to our previous approach of using task based Random Forest metric representation, we employ task based optimization to kernel matrix decomposition as described in CSI (Bach and Jordan., 2005) Algorithm 4. As previous discussed the kernel matrix are huge (Chang et al., 2006) and in the order of $N^2$ for real-time processing. To reduce the rank of the kernel matrix we use its labels (side-information) to mitigate pivots ($\gamma$) used during matrix factorization. The spatial feature and its residual characteristic of the significant Eigen values ($\lambda$) and its following Eigen vectors ($\nu$) are studied, the final rank obtained is always lower than the $N^2$ Gram matrix. Lines 5-9 in Algorithm 4 shows how the pivots are selected for a task based optimization. The above pre-processing step allows in-memory representation of large matrix.
VII. EXPERIMENTS ON UCI AND MSR DATASETS

The spatio-temporal datasets used in testing included forest cover (UCI), Statlog (UCI), and Geolife (Zheng et al., 2009) training and testing sets. We study the two task induced in-memory models: (i) QoD and explicit rule proximity matrix learning without kernel (bigrf-Table), and (ii) kernelized implementation in R based on labels (CSI-Kernel class). The performance of the predictive models are compared in three different memory scales (shown in x-axis of Fig 3): temporal windows in real-time (samples-per-sec), Original input space (Batch) ($R^p$), and $N \times N$ induced big-table feature space ($R^{p^2}$) (see Fig 5.)

A. Estimating Events Using Hoeffding Bounds for Big Sample - Real Variables

When estimating big samples the underlying source can be one of the following distributions:

- Normal
- Uniform

Algorithm 5 Cholesky decomposition with Side Information-CSI

1: procedure CSI($K$, $Y$)
2: Perform $\delta$ look-ahead steps of Cholesky and QR decomposition selecting pivots according to learning task
3: Initialization: $\eta = 2\epsilon$, $k = 1$
4: while $\min \eta > \epsilon$ and $k \leq m$
5: Compute estimated gains for the remaining pivots, and select best pivot
6: If new pivot not in the set of look-ahead pivots, perform a Cholesky and a QR step, otherwise perform the steps with a pivot optimization
7: Permute indices in the Cholesky and QR decomposition to put new pivot in position $k$
8: Compute exact gain $\eta$; let $k = k + 1$
9: end while
10: Output $G$ and its QR decomposition
11: end procedure

The accuracy of the ensemble model cannot be based on p-value alone as discussed in earlier sections. We introduce an reliability factor (Iyer2009b) which is used as a confidence measure to further distinguish the best feature to split on at the nodes. The sufficient statistics used by the in-memory model for prediction is based on Hoeffding bounds where $\delta \geq \epsilon$ (statistical significance).

$$\delta = \sqrt{\frac{1}{2m} \log \frac{4}{\delta}}$$

Where $: \delta \geq \epsilon$

1) Performance in Real-Time- Using Hoeffding bounds

To simulate a huge live streams we used the forest cover type dataset (Newman and Asuncion., 2007b), which has over 581,012 labeled samples. The predictive model builds a Hoeffding tree with 155 leaves nodes. Which took 12.69 seconds to build the tree in memory with a node splitting probability of $0.001$. The Hoeffding tree uses a very fast decision tree and closely approximates a full fledged decision tree. The estimated reduction of tree size from the original 53,904 leafs used by Random Forest classifier compared to 155 leaves used by Hoeffding tree. This fast decision tree implementation with incremental learning performs with a baseline accuracy of 79.5%.

<table>
<thead>
<tr>
<th>Sensors</th>
<th>$S_1$</th>
<th>$S_2$</th>
<th>$S_3$</th>
<th>$S_4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Observed</td>
<td>4.7 ± 2.0</td>
<td>1.6 ± 1.6</td>
<td>3.0 ± 1.5</td>
<td>1.8 ± 1.0</td>
</tr>
</tbody>
</table>

TABLE I A typical sensor dataset.

<table>
<thead>
<tr>
<th>$S_1$</th>
<th>$S_2$</th>
<th>$S_3$</th>
<th>$S_4$</th>
<th>Overlap</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>0.66</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>2</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
</tbody>
</table>

TABLE II Proximity counts after adding $S_4$ (good sensor) the ensemble overlap estimation now changes $S_1$ to tamely faulty.

B. Estimating Events Using Hoeffding Bounds Random Variables - Explicit Lower Dimension Representation

When mapping higher dimension space to an approximate linear lower dimension space, we use the following conversion:

- Euclidean distance
- Proximity distance

Optimization accuracy for linear case is:

$$\sup|Z(x) - Z(y) - K(x, y)| \leq \epsilon$$

1) Explicit Rule Extraction (Non-Gram matrix) in one dimension - $R$

A typical sensor measurement with an average and range that are provided as an base case example in Fig. 1. Using our proposed explicit feature metric which can be scaled between [0, 1]. This allows a uniform metric for all applications and to compare events of interest in any domain. In the example of four sensors we initially take three out of four to form an ensemble feature matrix. The matrix uses our proposed explicit feature representation as shown in this section. As shown in Fig. 3 (and tables) $S_1$ and $S_3$ fall into the same terminal node and their proximity count is incremented by one. Table I shows the overlapping rules on the right show the overlapped measure of reliability of $S_1$ - $S_3$ combined. If we use fuzzy interval definition of sensors to represent the varying levels of overlap then we represent its reliability by:

- Non faulty (= 1)
- Tamely faulty ($\geq 0.5$)
- Widely faulty ($\leq 0.5$)

Applying the rules to our dataset we find $S_2$ and $S_3$ are non faulty (as illustrated in table) as they have the maximum overlapping count. At the same time $S_1$ and $S_2$ is tamely faulty as $S_4$ is inclusive of $S_1$. The higher level application can now rely on measurements from $S_2$ and $S_3$ for its precision and range accuracy.

To further see the ensemble optimization we introduce $S_4$ and create the new feature matrix as formed in Table II. Reading from the data in the overlap column we see that $S_4$ has a maximum overlap of 1. The addition does not have any effect on $S_1$. The accuracy of the random subspace learned from the trees include $S_4$ as it has a narrow range compared to all others. The accuracy of the one dimensional case can then be measured as, 3 out of 4 sensors have a high degree of precision.
2) Explicit Rule Extraction (Non-Gram Matrix) in two dimension - $R^2$

Experiments are conducted Using the Geolife trajectory dataset similar to the one-dimensional case discussed as the base case example. The explicit learning algorithms use a tree structure to learn the user’s trajectories generated by mobility models of car, walking and riding bus. The matrix representation of the feature space shows that the mobile user’s walking class has the highest information gain as marked (green) in Table III. The predictive peak performance of bigrl-table is 96% for an ensemble size of 100 trees as shown in Fig. 5, when compared to the real-time streaming model. The added memory overhead of using a $N \times N$ in the two-dimensional case is solved by using distributed shared configuration across multiple nodes of a cluster.

### C. Estimating Events Using Hoeffding Bounds for Kernel Variables - Curse of Support

When dealing with support vector machine to transform non-linear relations to linear space we use the following kernels:

- ANOVA kernel $k(x; x') = \sum_{1 \leq i_1 < i_2 \leq N} \prod_{d=1}^{D} k(x_{i_d}; x'_{i_d})$
- Bassel Kernel $k(x; x') = (-Bessel_{p+1}^n \rho \|x - x'||)$
- Radial Basis Kernel $k(x; x') = exp(-\rho \|x - x'||^2)$
- Laplace Kernel $k(x; x') = exp(-\rho \|x - x'||)$

Optimization accuracy for non-linear case is:

$$sup |K(x, y)| \leq \epsilon$$

1) Data Representation Using Gram Matrix with Side Information Optimization - $FR^d$

For computational comparison, we perform feature space analysis using the Incomplete Cholesky (CSI) R package. The task based optimization uses pivoting to reduces the original [4434x4434] matrix. The final feature matrix rank is reduced from 4437 to 20 as show in Table IV. The lower rank outcome is based on the type of features induced using the kernel functions. ANOVA kernel uses the least number of samples and at the same time has superior prediction accuracy. The rest of the kernels RBF, Laplace and Bassel have comparable accuracies but the computation is $5 \times$ slower due to large matrix ranks. Further using visualization and projection as shown in Fig. 4. (Scholkopf et al., 1999) reveals that ANOVA kernel is able to induce higher-order features (up to 20) to separate all the six categories. This is due to ANOVA being able to learn statistical similarities from subspaces and represent them efficiently as factors. Earlier results verify from section IV on side-information using associates rules for the same dataset that the attributes are statistically correlated.

The same experiments are repeated without PCA like optimization and using the kernels directly with the input datasets. The following results for the non-optimized Gram matrix are shown in Table V. The overall reduction is accuracy is by 10%.

### 2) Data Representation Using $N \times N$ Gram Matrix - with No Memory Optimization

The standard kernels: ANOVA, Gaussian RBF and linear SVMs are used with full rank matrices (i.e. without memory optimization). The non-optimized model’s accuracies are moderate to base line variability, when compared to kernels with in-memory optimization. The comparative results are illustrated in the Fig. 4., ANOVA-kernel has a peak performance of 89.56% is significantly lower than the task based approach. The kernel KSVRM R package (Karatzoglou et al., 2004) allows to tune the SVM (Blanchard et al., 2004) soft margin parameters $C$ and $\gamma$ by use of cross-validation. The higher the $C$ the better would be the fit (see Section 2), but it may not generalize well. Using the cross-validation techniques discussed above, we arrive at $d=2, \sigma=0.015,C=50$, when using cross-validation (cross=4) for the spatio-temporal example UCI dataset (Newman and Asuncion., 2007).

We provide the optimization driven by real-time streaming requires with related cloud streaming analytics services. The details are in the plot as shown in Fig. 5. The on-line version of TREEFORESTCOVER supports multi-class and labeled samples in its in-memory implementation similar to the fault-tolerant algorithm of Brook-Iyengar. The approximate Hoeffding tree performs close to the best tree classifier with an accuracy of 79.5%. The results signifies the Hoeffding tree are among the best category of classifiers.

<table>
<thead>
<tr>
<th>Sensors</th>
<th>$S_1$</th>
<th>$S_2$</th>
<th>$S_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2.7</td>
<td>0</td>
<td>1.5</td>
</tr>
<tr>
<td>2</td>
<td>4.7</td>
<td>1.6</td>
<td>3.0</td>
</tr>
<tr>
<td>3</td>
<td>6.7</td>
<td>3.2</td>
<td>4.5</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sensors</th>
<th>$S_1$</th>
<th>$S_2$</th>
<th>$S_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>BUS</td>
<td>488</td>
<td>0</td>
<td>76</td>
</tr>
<tr>
<td>CAR</td>
<td>0</td>
<td>516</td>
<td>1</td>
</tr>
<tr>
<td>WALK</td>
<td>133</td>
<td>1</td>
<td>294</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Overlap</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.135</td>
</tr>
<tr>
<td>0.001</td>
</tr>
<tr>
<td>0.313</td>
</tr>
</tbody>
</table>

**Table III**: Feature induced matrix using two dimensional datasets.

**Fig. 3**: $S_1$ & $S_2$ end up in the same terminal node. The Big-Table proximity count is incremented by 1. (The color metric in the overlap columns specifies quality of data: Green=non faulty; Blue= tamely faulty; Red=widely faulty)
TABLE IV Reconstruction Error: Results for the UCI-Landsat dataset when pre-processing features with statistical properties of kernels (included Incomplete-Cholesky). Configuration: Intel i5-2435M Processor, 2.40GHz; 4GB DDR3. The timings included show feature extraction during pre-processing. Side-Info column indicates if it is supervised or unsupervised learning.

<table>
<thead>
<tr>
<th>Pivot selection</th>
<th>Training Accuracy</th>
<th>Rank</th>
<th>SV</th>
<th>Time (Batch in sec.)</th>
<th>Side-Info.</th>
</tr>
</thead>
<tbody>
<tr>
<td>ANOVA kernel</td>
<td>89.6%</td>
<td>20</td>
<td>20</td>
<td>40</td>
<td>Y</td>
</tr>
<tr>
<td>Laplace kernel</td>
<td>86.5%</td>
<td>100</td>
<td>1489</td>
<td>120</td>
<td>Y</td>
</tr>
<tr>
<td>RBF kernel</td>
<td>86.4%</td>
<td>100</td>
<td>1488</td>
<td>120</td>
<td>Y</td>
</tr>
<tr>
<td>RBF kernel</td>
<td>78.6%</td>
<td>100</td>
<td>1636</td>
<td>120</td>
<td>Y</td>
</tr>
<tr>
<td>Inclh RBF</td>
<td>-</td>
<td>3067</td>
<td></td>
<td></td>
<td>N</td>
</tr>
</tbody>
</table>

TABLE V Prediction Error: Results for the UCI-Landsat dataset when post-processing to predict or categorize the datasets using unknown test samples. The timings included show training time during learning the models.

<table>
<thead>
<tr>
<th>Models</th>
<th>Gram Matrix</th>
<th>Task Matrix</th>
</tr>
</thead>
<tbody>
<tr>
<td>Statistical Models</td>
<td>Train Test</td>
<td>t=25 t=50 t=75 t ≥ 100</td>
</tr>
<tr>
<td>Task Based Learning - Trajectory Dataset</td>
<td>91% 95% 96%</td>
<td>↑</td>
</tr>
<tr>
<td>Task Based Learning - Landsat Dataset</td>
<td>89% 91% 85%</td>
<td>↓</td>
</tr>
<tr>
<td>ANOVA-Kernel SVM</td>
<td>81.6% 82.87%</td>
<td></td>
</tr>
<tr>
<td>RBF-Kernel SVM</td>
<td>62.1% 52.2%</td>
<td></td>
</tr>
<tr>
<td>Linear SVM</td>
<td>69.1% 67.7%</td>
<td></td>
</tr>
<tr>
<td>Hoefdding Tree Learning - Forest Cover Type Dataset</td>
<td>100% (baseline)</td>
<td>8% (original RF)</td>
</tr>
</tbody>
</table>

The ensemble based rLBig explicit deep feature engineering framework performs significantly well for the synthetic datasets such as trajectories. The popularity of distance based kernel method shows how higher-order features helps to better predictive performance, without knowing a priori the underlying distribution, specially with few number of training data.

VIII. CONCLUSION

In this work we have extended an existing real-time sensor network algorithm using Hoeffding bounds to learn rare events of practical significance which can be applied to any stream distribution. The rare event feature learning which are computationally expensive have been adapted for resource constrained real-time streaming needs. The enhancements distinguishes the impact on model performance which sometimes can be biased due to the datasets. The spatio-temporal concepts learned shows that explicit rule learning has many advantages and can significantly increase performance when used with mobile trajectories. On the other hand when learning the kernel function and its Gram matrix, and using far more induced features helps the model to predict with higher accuracy. The use of spatio-temporal datasets brings out the art of pattern recognition and AI to life. The task based approach when applied to explicit and kernel based prediction has shown improved accuracy compared to earlier kernel versions. This AI framework’s enhancements will help develop geo-spatial applications which can reliably and efficiently find actionable events from satellite imagery. The learning of stream uses a non-parametric approach for outliers detection in time series streams and, a task based approach to improve QoD.

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Borchers R.R., Iyengar, S.S., 1996. Robust Distributed Computing and Sensing Algorithm. IEEE Computer 29, 6 (June 1996), 5360. DOI:http://dx.doi.org/10.1109/2.507632


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Fig. 4 Random project methods for kernel learning: PCA of stream data taken from four Statlog channels to detect six soil types. Soil types are separated using higher-order statistical properties in (a),(b) and (c). (a) and (b) are higher-order statistics while (c) are present in the original dataset. The hyperplane boundaries in (a) are able to separate all the six classes while in (b) a maximum of five are uniquely separable.


Random Forest Performance: (a) SATLOG (Spatial Landsat) (b) GEOLIFE Trajectory.

(c) Gram Matrix SATLOG (Spatial Landsat) (d) Hoeffding Tree.

Stream Learning combined performance plot.

Fig. 5 Average accuracy of randomForest, randomForest-Big, CSI-Kernel matrix and Hoeffding tree and over many datasets versus the maximum number of options per example. Accuracies were estimated to perform memory optimizations for the final model. The real-time streaming is shown in green which performs close to the best known batch tree model.