Scalable Distributed Change Detection from Astronomy Data Streams using Local, Asynchronous Eigen Monitoring Algorithms

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Abstract
This paper considers the problem of change detection using local distributed eigen monitoring algorithms for next generation of astronomy petascale data pipelines such as the Large Synoptic Survey Telescopes (LSST). This telescope will take repeat images of the night sky every 20 seconds, thereby generating 30 terabytes of calibrated imagery every night that will need to be co-analyzed with other astronomical data stored at different locations around the world. Change point detection and event classification in such data sets may provide useful insights to unique astronomical phenomenon displaying astrophysically significant variations: quasars, supernovae, variable stars, and potentially hazardous asteroids. However, performing such data mining tasks is a challenging problem for such high-throughput distributed data streams. In this paper we propose a highly scalable and distributed asynchronous algorithm for monitoring the principal components (PC) of such dynamic data streams. We demonstrate the algorithm on a large set of distributed astronomical data to accomplish well-known astronomy tasks such as measuring variations in the fundamental plane of galaxy parameters. The proposed algorithm is provably correct (i.e. converges to the correct PCs without centralizing any data) and can seamlessly handle changes to the data or the network. Real experiments performed on Sloan Digital Sky Survey (SDSS) catalogue data show the effectiveness of the algorithm.

1 Introduction
Data mining is playing an increasingly important role in astronomy research [18] involving very large sky surveys such as Sloan Digital Sky Survey SDSS and the 2-Micron All-Sky Survey 2MASS. These sky-surveys are offering a new way to study and analyze the behavior of the astronomical objects. The next generation of sky-surveys are poised to take a step further by incorporating sensors that will stream in large volume of data at a high rate. For example, the Large Synoptic Survey Telescopes (LSST) will take repeat images of the night sky every 20 seconds. This will generate 30 terabytes of calibrated imagery every night that will need to be co-analyzed with other astronomical data stored at different locations around the world. Change point detection and event classification in such data sets may provide useful insights to unique astronomical phenomenon displaying astrophysically significant variations: quasars, supernovae, variable stars, and potentially hazardous asteroids. Analyzing such high-throughput data streams would require large distributed computing environments for offering scalable performance. The knowledge discovery potential of these future massive data streams will not be achieved unless novel data mining and change detection algorithms are developed to handle decentralized petascale data flows, often from multiple distributed sensors (data producers) and archives (data providers). Several distributed computing frameworks are being developed [12], [16], [17], [13] for such applications. We need distributed data mining algorithms that can operate on such distributed computing environments. These algorithms should be highly scalable, be able to provide good accuracy and should have a low communication overhead.

This paper considers the problem of change detection in the spectral properties of data streams in a distributed environment. It offers an asynchronous, communication-efficient distributed eigen monitoring (DDM) algorithm for monitoring the principal components (PCs) of dynamic astronomical data streams. It particularly considers an important problem in astronomy regarding the variation of fundamental plane structure of galaxies with respect to spatial galactic density and demonstrates the power of DDM algorithms using this example application. This paper presents the algorithm, analytical findings, and results from experiments. Experiments are performed using currently available astronomy data sets from virtual observatories. Our distributed algorithm is a first step in analyzing the astronomy data arriving from such high throughput data streams of the future. The specific contributions of this paper can be summarized as follows:

• To the best of the authors knowledge this is one of the
first attempts on developing a completely asynchronous and local algorithm for doing eigen analysis in distributed data streams.

- Based on data sets downloaded from astronomy catalogues such as SDSS and 2MASS, we demonstrate how the galactic fundamental plane structure varies with difference in galactic density.

Section 2 describes the astronomy problem. Section 3 presents the related work. Section 4 offers the background material and formulates the data mining problem. Section 5 describes the centralized version of the problem while Section 6 models the distributed version and explains the eigenstate monitoring algorithm. Section 7 presents the experimental results. Finally, Section 8 concludes this paper.

2 The Astronomy Problem

When the LSST astronomy project becomes operational within the next decade, it will pose enormous petascale data challenges. This telescope will take repeat images of the night sky every 20 seconds, throughout every night, for 10 years. Each image will consist of 3 gigapixels, yielding 6 gigabytes of raw imagery every 20 seconds and nearly 30 terabytes of calibrated imagery every night. From this “cosmic cinematography”, a new vision of the night sky will emerge – a vision of the temporal domain – a ten-year time series (movie) of the Universe. Astronomers will monitor these repeat images night after night, for 10 years, for everything that has changed – changes in position and intensity (flux) will be monitored, detected, measured, and reported. For those temporal variations that are novel, unexpected, previously unknown, or outside the bounds of our existing classification schemes, astronomers will want to know (usually within 60 seconds of the image exposure) that such an event (a change in the night sky) has occurred. This event alert notification must necessarily include as much information as possible to help the astronomers around the world to prioritize their response to each time-critical event. That information packet will include a probabilistic classification of the event, with some measure of the confidence of the classification. What makes the LSST so incredibly beyond current projects is that most time-domain sky surveys today detect 5-10 events per week; LSST will detect 10 to 100 thousand events per night! Without good classification information in those alert packets, and hence without some means with which to prioritize the huge number of events, the astronomy community would consequently be buried in the data deluge and will miss some of the greatest astronomical discoveries of the next 20 years (perhaps even the next “killer asteroid” heading for Earth – this time, it won’t be the dinosaurs that will go extinct!).

To solve the astronomers’ massive event classification problem, a collection of high-throughput change detection algorithms will be needed. These algorithms will need to access distributed astronomical databases worldwide to correlate with each of those 100,000 nightly events, in order to model, classify, and prioritize correctly each event rapidly. One known category of temporally varying astronomical object is a variable star. There are dozens of different well known classes of variable stars, and there are hundreds (even thousands) of known examples of these classes. These stars are not “interesting” in the sense that they should not produce alerts (change detections), even though they are changing in brightness from hour to hour, night to night, week to week – their variability is known, well studied, and well characterized already. However, if one of these stars’ class of variability were to change, that would be extremely interesting and be a signal that some very exotic astrophysical processes are involved. Astronomers will definitely want to be notified promptly (with an alert) of these types of variations. Just what is this variation? It is essentially a change in the Fourier components (eigenvectors) of the temporal flux curve (which astronomers call “the light curve”). This problem has several interesting data challenge characteristics: (1) the data streaming rate is enormous (6 gigabytes every 20 seconds); (2) there are roughly 100 million astronomical objects in each of these images, all of which need to monitored for change (i.e., a new variable object, or a known variable object with a new class of variability); (3) 10 to 100 thousand “new” events will be detected each and every night for 10 years; and (4) distributed data collections (accessed through the Virtual Astronomy Observatory’s worldwide distribution of databases and data repositories) will need to correlated and mined in conjunction with each new variable object’s data from LSST, in order to provide the best classification models and probabilities, and thus to generate the most informed alert notification messages.

Astronomers cannot wait until the year 2016 (when LSST begins its sky survey operations) for new algorithms to begin being researched. Those algorithms (for distributed mining, change detection, and eigenvector monitoring) will need to be robust, scalable, and validated already at that time. So, it is imperative to begin now to research, test, and validate such data mining paradigms through experiments that replicate the expected conditions of the LSST sky survey. Consequently, we have chosen an astronomical research problem that is both scientifically valid (i.e., a real astronomy research problem today) and that parallels the eigenvector monitoring problem that we have described above. We have chosen to study the principal components of galaxy parameters as a function of an independent variable, similar to the temporal dynamic stream mining described above. In our current experiments, the independent variable is not the time dimension, but local galaxy density.

The class of elliptical galaxies has been known for 20 years to show dimension reduction among a subset of physical attributes, such that the 3-dimensional distribution
of three of those astrophysical parameters reduce to a 2-dimensional plane. The normal to that plane represents the principal eigenvector of the distribution, and it is found that the first two principal components capture significantly more than 90% of the variance among those 3 parameters.

By analyzing existing large astronomy databases (such as the Sloan Digital Sky Survey SDSS and the 2-Micron All-Sky Survey 2MASS), we have generated a very large data set of galaxies. Each galaxy in this large data set was then assigned (labeled with) a new "local galaxy density" attribute, calculated through a volumetric Voronoi tessellation of the total galaxy distribution in space. Then the entire galaxy data set was horizontally partitioned across several dozen partitions as a function of our independent variable: the local galaxy density.

As a result, we have been able to study eigenvector changes of the fundamental plane of elliptical galaxies as a function of density. Computing these eigenvectors for a very large number of galaxies, one density bin at a time, in a distributed environment, thus mimics the future LSST dynamic data stream mining change detection (eigenvector change) challenge problem described earlier. In addition, this galaxy problem actually has uncovered some new astrophysical results: we find that the variance captured in the first 2 principal components increases systematically from low-density regions to high-density regions of space, and we find that the direction of the principal eigenvector also drifts systematically in the 3-dimensional parameter space from low-density regions to the highest-density regions.

3 Related Work

The work related to this area of research can broadly be subdivided into data analysis for large scientific data repository and distributed data mining in a dynamic networks of nodes. We discuss each of them in the following two sections.

3.1 Analysis of Large Scientific Data Collections The U.S. National Virtual Observatory (NVO)\(^1\) is a large scale effort to develop an information technology infrastructure enabling easy and robust access to distributed astronomical archives. It will provide services for users to search and gather data across multiple archives and will provide some basic statistical analysis and visualization functions. The International Virtual Observatory Alliance (IVOA)\(^2\) is the international steering body that federates the work of about two dozen national VOs across the world (including the NVO in the US). The IVOA oversees this large-scale effort to develop an IT infrastructure enabling easy and robust access to distributed astronomical archives worldwide.

There are several instances in the astronomy and space sciences research communities where data mining is being applied to large data collections [12][10][2]. Another recent area of research is distributed data mining [20][7] which deals with the problem of data analysis in environments with distributed data, computing nodes, and users. Distributed eigen-analysis and outlier detection algorithms have been developed for analyzing astronomy data stored at different locations by Dutta et al.[14]. Kargupta et al. [7] have developed a technique for performing distributed principal component analysis by first projecting the local data along its principal components and then centralizing the projected data. In both these cases, the data is distributed vertically (different full attribute columns reside at different sites), while in this paper, the data is distributed horizontally (different data tuple sets reside at different sites). Moreover, none of the above efforts address the problem of analyzing rapidly changing astronomy data streams.

3.2 Data Analysis in Large Dynamic Networks There is a significant amount of recent research considering data analysis in large-scale dynamic networks. Since efficient data analysis algorithms can often be developed based on efficient primitives, approaches have been developed for computing basic operations (e.g. average, sum, max, random sampling) on large-scale, dynamic networks. Kempe et al. [21] and Boyd et al. [8] developed gossip based randomized algorithms. These approaches used an epidemic model of computation. Bawa et al. [4] developed an approach based on probabilistic counting. In addition, techniques have been developed for addressing more complex data mining/data problems over large-scale dynamic networks: association rule mining [26], facility location [22], outlier detection [9], decision tree induction [7], ensemble classification [23], support vector machine-based classification [1], K-means clustering [11], top-K query processing [3].

A related line of research concerns the monitoring of various kinds of data models over large numbers of data streams. Sharfman et al. [24] develop an algorithm for monitoring arbitrary threshold functions over distributed data streams. And, most relevant to this paper, Wolff et al. [25] developed an algorithm for monitoring the L2 norm. We use this technique to monitor eigen-states of the fundamental plane concerning elliptical galaxies.

Huang et al. [19] consider the problem of detecting network-wide volume anomalies via thresholding the length of a data vector (representing current network volume) projected onto a subspace closely related to the dominant principal component subspace of past network volume data vectors. Unlike us, these authors consider the analysis of a vertically distributed data set. Each network node holds a sliding window stream of numbers (representing volume through time) and the network-wide volume is represented as a matrix with each column a node stream. Because of the

\(^1\)http://www.us-vo.org/
\(^2\)http://www.ivoa.net
difference in data distribution (vertical vs. horizontal), their
approach is not applicable to our problem. We assume that
each node is receiving a stream of tuples and the network-
wide dataset is matrix formed by the union of all nodes’ cur-
cently held tuples (each node holds a collection of rows of the
matrix rather than a single column as considered by Huang).

In the next section we present the notations and problem
definition that will be used throughout the rest of the paper.

4 Background

In order to analyze the data streams from the next genera-
tion of large scale astronomy systems such as the ones con-
structed by the LSST project, we need scalable infrastructure
for computing. It is generally agreed among the astronomy
community that the computing infrastructure will be a grid-
like environment comprised of a collection of desktop com-
pute nodes, high performance machines, and data sources
among others. We need data analysis algorithms that will
be able to work in this distributed heterogeneous computing
environment. This paper offers distributed eigen-analysis al-
gorithms that can handle data from distributed nodes (either
inherently distributed data or artificially distributed in order
to scale up the performance).

In the remainder of this section we first define the nota-
tions that will be used to discuss our distributed algorithms
and then formally state the problem definition.

4.1 Notations Let \( V = \{P_1, \ldots, P_n\} \) be a set of nodes
connected to one another via an underlying communication
infrastructure such that the set of \( P_i \)’s neighbors, \( \Gamma_i \), is
known to \( P_i \). Additionally, at any time, \( P_i \) is given a
time-varying data matrix \( M_i \) where the rows correspond
to the instances and the columns correspond to attributes
or features. Mathematically, \( M_i = [x_{i,1}, x_{i,2}, \ldots]^{T} \), where
each \( x_{i,t} = [x_{i,1}, x_{i,2}, \ldots, x_{i,d}] \in \mathbb{R}^d \) is a row vector. The
covariance matrix of the data at node \( P_i \), denoted by \( C_i \), is the
matrix whose \( (i, j) \)-th entry corresponds to the covariance
between the \( i \)-th and \( j \)-th feature (column) of \( M_i \). The
global data set of all the nodes’ data is \( G = \bigcup_{i=1}^{n} M_i \).

It can be shown that if the attributes of \( G \) are mean
shifted, \( i.e. \) the mean of each attribute is subtracted from
each value of that attribute, the covariance matrix can be
written as \( C = G^{T} G \) (we have ignored the scaling by the
number of points in \( G \)). Also under such conditions, it can
be shown that \( C = \sum_{i=1}^{n} C_i \).

4.2 Problem Formulation The identification of certain
correlations among parameters has lead to important discov-
eries in astronomy. For example, the class of elliptical and
spiral galaxies (including dwarfs) have been found to occupy
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5 Centralized Principal Components Analysis for the
Fundamental Plane Computation

For identifying the variability of fundamental plane on the
basis of galactic densities, we have used the SDSS and
2MASS data sets available individually through the NVO.
Since galactic density is not observed by the NVOs, we have
cross-matched the two data sets and computed the densities
based on other property values. In this section we describe
the data gathering procedure for this approach followed by the
PCA computation.

5.1 Data Preparation We create a large, aggregate
data set by downloading the 2MASS XSC extended
source catalog (http://irsa.ipac.caltech.edu/
aplications/Gator/) for the entire sky and cross-
match it against the SDSS catalog using the SDSS
Crossid tool (http://cas.sdss.org/astro/en/
tools/crossid/upload.asp) such that we select all
unique attributes from the PhotoObjAll and SpecObjAll ta-
tables as well as the photozd1 attribute from the Photoz2 table
which is an estimated redshift value. We filter the data based

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unique attributes from the PhotoObjAll and SpecObjAll ta-
tables as well as the photozd1 attribute from the Photoz2 table
which is an estimated redshift value. We filter the data based
on the SDSS identified type to remove all non-galaxy tuples. We then filter the data again based on reasonable redshift (actual or estimated) values (0.003 \leq z \leq 0.300).

For creating the new attribute, namely, galactic density, we transform the attributes \( cx, cy, cz \) (unit vectors), \( z \), and \( \text{photozd1} \) to 3D Euclidean coordinates using the transformation

\[
(X, Y, Z) = (Distance \times cx, Distance \times cy, Distance \times cz)
\]

where \( Distance = 2 \times \left[ 1 - \frac{1}{\sqrt{1+\text{redshift}}} \right] \). We finally use these Cartesian coordinates to compute the Delaunay Triangulation\(^3\) of each point (galaxy) in 3D space. To remove bias in the density calculation of the Delaunay cells, we identify all boundary points and remove them from the computation. This tessellation procedure is a data transformation step, which converts the spatial location of a galaxy (within the 3-D distribution of galaxies) into a single numeric attribute (local galaxy density). This parameter has astrophysical significance, even more than the actual spatial location information (i.e., galaxy properties are often modified and governed by the proximity of nearby galaxies, such as in high-density environments), and so we chose to use this new attribute — local galaxy density, as estimated through the triangulation step — because it has strong astrophysical relevance and scientific significance. This is a robust estimator and is as scientifically meaningful as any other attribute in the science database used in these experiments. Now using the output of the Delaunay triangulation the volumes of the Delaunay cells are computed using the expression

\[
\text{vol}(i) = \frac{1}{6} \cdot \det(\vec{a}_i - \vec{b}_i, \vec{b}_i - \vec{c}_i, \vec{c}_i - \vec{h}_i),
\]

where \( \vec{a}_i, \vec{b}_i, \vec{c}_i \) and \( \vec{h}_i \) are the vertices of the tetrahedron corresponding to the \( i \)-th point in 3D Euclidean space. The volume corresponding to the \( i \)-th point is the sum of the volumes of all tetrahedrons that contain the particular point. Using the DTFE formulation\(^4\), the density of the \( i \)-th cell is then computed as follows:

\[
\text{den}(i) = (D + 1) \times \frac{m_i}{\text{vol}(i)}
\]

where \( m_i = 1 \), since we have one object (galaxy) per cell and \( D = 3 \) for triangulation in 3D-space.

5.2 Binning and PCA The astronomy question that we want to address here is whether the fundamental plane structure of galaxies in low density regions differ from that of galaxies in high density regions. For this we take the above data set containing 155650 tuples and associate with each tuple, a measure of its local galactic density. Our final aggregated data set has the following attributes from SDSS: Petrosonian I band angular effective radius (\( Iaer \)), redshift (\( rs \)), and velocity dispersion (\( vd \)); and has the following attribute from 2MASS: K band mean surface brightness (\( Knsb \)). We produce a new attribute, logarithm Petrosonian I band effective radius (\( \log(Ier) \)), as \( \log(Iaer*rs) \) and a new attribute, logarithm velocity dispersion (\( \log(vd) \)), by applying the logarithm to \( vd \). We finally append the galactic density (\( \text{cellDensity} \)) associated with each of the tuples as the last attribute of our aggregated data set. We divide the tuples into 30 bins based on increasing cell density, such that there are equal number of tuples in each bin. For each bin we carry out the fundamental plane calculation or principal component analysis and observe that the percent of variance captured by the first two PCs is very high. This implies that the galaxies can be represented by the plane defined by the first two eigen vectors. It is also observed that this percentage increases for bins with higher mean galactic density. We report these results in Section 7.

As discussed earlier, analysis of very large astronomy catalogs can pose serious scalability issues, especially when considering streaming data from multiple sources. In the next section we describe a distributed architecture for addressing these issues and then show how the centralized eigen analysis of the covariance matrix can be formulated as a distributed computation and how it can be solved in a communication efficient manner.

6 Distributed Principal Component Analysis

When resources become a constraint for doing data mining on massive data sets such as astronomical catalogs, distributed data mining provides a communication efficient solution. For the problem discussed in the last section, we can formulate a distributed architecture where after cross matching the data using a centralized cross matching tool, we can store the meta data information in a central location. Such a service-oriented architecture would facilitate astronomers to query multiple databases and do data mining on large data sets without downloading the data to their local computing resources. The data set is downloaded in parts at a number of computing nodes (that are either dedicated computers connected through communication channels or part of a large grid) based on the meta data information maintained at the central server site. In such a computational environment, distributed data mining algorithms can run in the background seamlessly for providing fast and efficient solutions to the astronomers by distributing the task among a number of nodes. Figure 1 represents one such architecture.

Another distributed data mining scenario for large scale astronomy databases is the one described in Section 2 for the LSST project where high throughput data streams need to be
modeled and monitored for changes in an efficient manner. In the next few sections we describe a distributed formulation of our centralized eigen analysis and present a eigenstate monitoring algorithm for this purpose.

6.1 Problem formulation: Distributed Covariance Computation For the distributed setup, the entire data is not located at a central location. The data set of node \( P_i \) is \( \mathcal{M}_i \). Note that, \( \mathcal{G} = \bigcup_{i=1}^{n} \mathcal{M}_i \). It is true that, if the mean of each column of \( \mathcal{G} \) is subtracted from each value of \( \mathcal{G} \), i.e. \( \mathcal{G}_1 \) is mean-reduced \( \mathcal{G} \), then the covariance matrix of \( \mathcal{G} \) i.e. \( \mathcal{C} \) can be written as \( \mathcal{C} = \frac{1}{\# \text{points in } \mathcal{G}} \mathcal{G}_1^T \mathcal{G}_1 \). Now, in the distributed setup it is true that:

\[
\mathcal{C} = \frac{1}{\# \text{points in } \mathcal{G}} \mathcal{G}_1^T \mathcal{G}_1 = \frac{1}{\# \text{points in } \mathcal{G}} \sum_{i=1}^{n} \mathcal{M}_{1i}^T \mathcal{M}_{1i}
\]

where \( \mathcal{M}_{1i} \) is mean reduced \( \mathcal{M}_i \). Thus it turns out that if data is horizontally partitioned among \( n \) nodes and each column of data is mean shifted using the global mean, the covariance matrix is completely decomposable. With this formulation, we now describe certain notations for discussing our distributed eigen monitoring algorithm.

6.2 Preliminaries The goal of the PC monitoring algorithm is to track changes to the eigenvectors of the global covariance data matrix \( \mathcal{C} \). The crux lies in each node maintaining a current set of eigenvectors which it believes to be globally correct. We call it the knowledge of a node. Also each node checks if it is in agreement with all of its immediate neighbors with respect to the knowledge. It can be shown that if this is true for all nodes in the network, then the local eigenvectors of each node is indeed the correct global solution. Note that from our earlier discussion, \( \mathcal{G}^T \mathcal{G} = \mathcal{C} \) when \( \mathcal{G} \) is mean shifted. In the distributed setup, the mean of the global data is not known to each node. Therefore we decompose the PC monitoring algorithm in to (1) mean monitoring which maintains the correct mean of \( \mathcal{G} \), and (2) eigenvector monitoring of \( \mathcal{G}^T \mathcal{G} \). Given an eigenvalue or a mean as a model, each algorithm monitors changes to the corresponding model with respect to the global data using only its knowledge and agreement. If the data changes such that the models no longer fit the data, the algorithms raise a flag at each node. At this point, a sample of the data is centralized, new models are built and then disseminated to the network. The monitoring algorithms are then restarted with the new models and the process continues. Below we formally define these quantities and describe the algorithms.

6.2.1 Notations and Assumptions In this section we present certain notations necessary for the algorithms.

In the algorithm, each node sends messages to its immediate neighbors to converge to a globally correct solution. As already discussed, there are three kinds of messages: (i) monitoring messages which are used by the algorithm to check if the model is up-to-date, (ii) data messages which are used to sample data for rebuilding a model, and (iii) model messages which are used to disseminate the newly built model in the entire network. In this section we will discuss messages of the first type only. The other two will be discussed in the later sections since they are algorithm specific.

Let the model supplied to each of the monitoring algorithms be denoted by \( L \). For the mean monitoring algorithm, the model is a mean vector \( \overrightarrow{\mu_i} \); for the eigenvector monitoring, the model is a set of eigenvectors \( (\frac{1}{V}) \) and eigenvalues \( (\mathcal{G}_i) \). Let \( \mathcal{E}_i(\mathcal{M}_i, L) \) be the error between the model \( L \) and the data of node \( P_i \). Explicit computation of \( \mathcal{E}_i \) is problem specific and hence described in respective algorithm descriptions. The nodes jointly track if \( \mathcal{E}^V = \bigcup_{i=1}^{n} \mathcal{E}_i \) is less than a user-defined threshold \( \epsilon \).

Any monitoring message sent by node \( P_i \) to \( P_j \) contains information that \( P_i \) has gathered about the network which \( P_j \) may not know. In our case, the message sent by \( P_i \) to \( P_j \) consists of a set of vectors or matrix\(^5\) \( S_{i,j} \) with each row corresponding to observations and each column corresponding to features. Note that if each node broadcasts \( S_{i,j} = \mathcal{M}_i \), then each node would obviously be able to compute the correct result. However this would be communication intensive. Our next few sets of vectors allow us to compute the correct result in a more communication efficient manner.

Knowledge This is all the information that \( P_i \) has about the error:

\[
\mathcal{K}_i = \mathcal{E}_i \cup \bigcup_{P_j \in \Gamma_i} \mathcal{S}_{j,i}
\]

Agreement This is what \( P_i \) and \( P_j \) have in common:

\[
\mathcal{A}_{i,j} = \mathcal{S}_{i,j} \cup \mathcal{S}_{j,i}
\]

\(^5\)we use them interchangeably here
Held  This is what $P_i$ has not yet communicated to $P_j$
\[ \mathcal{H}_{i,j} = \mathcal{K}_i \setminus \mathcal{A}_{i,j} \]

These sets of vectors can be arbitrarily large. It can be shown that if vectors sent by $P_i$ to $P_j$ are never sent back to $P_i$, we can do the same computations using only the average vector of these sets and the size of the set. One way of ensuring this is to assume that communication takes place over a communication tree— an assumption we make here (see [25] and [7] for a discussion of how this assumption can be accommodated or, if desired, removed).

The following are the notations used for the set statistics — (1) average: $\overline{\mathcal{K}}_i, \overline{\mathcal{A}}_{i,j}, \overline{\mathcal{H}}_{i,j}, \overline{\mathcal{S}}_{i,j}, \overline{\mathcal{S}}_{j,i}, \overline{\mathcal{E}}_i, \text{ and } \overline{\mathcal{E}}^\Theta$, and (2) sizes: $|\mathcal{S}_{i,j}|, |\mathcal{S}_{j,i}|, |\mathcal{K}_i|, |\mathcal{A}_{i,j}|, |\mathcal{H}_{i,j}|, |\mathcal{E}_i|$, and $|\mathcal{E}^\Theta|$. With these notations, we can now write
\[ |\mathcal{K}_i| = |\mathcal{E}_i| + \sum_{P_j \in \Gamma_i} |\mathcal{S}_{j,i}| \]
\[ |\mathcal{A}_{i,j}| = |\mathcal{S}_{i,j}| + |\mathcal{S}_{j,i}| \]
\[ |\mathcal{H}_{i,j}| = |\mathcal{K}_i| - |\mathcal{A}_{i,j}| \]

Similarly for the average of the sets we can write,
\[ \overline{\mathcal{K}}_i = \frac{1}{|\mathcal{K}_i|} \left( |\mathcal{E}_i| \overline{\mathcal{E}}^\Theta + \sum_{P_j \in \Gamma_i} |\mathcal{S}_{j,i}| \overline{\mathcal{S}}_{j,i} \right) \]
\[ \overline{\mathcal{A}}_{i,j} = \frac{1}{|\mathcal{A}_{i,j}|} \left( |\mathcal{S}_{i,j}| \overline{\mathcal{S}}_{i,j} + |\mathcal{S}_{j,i}| \overline{\mathcal{S}}_{j,i} \right) \]
\[ \overline{\mathcal{H}}_{i,j} = \frac{1}{|\mathcal{H}_{i,j}|} \left( |\mathcal{K}_i| \overline{\mathcal{K}}_i - |\mathcal{A}_{i,j}| \overline{\mathcal{A}}_{i,j} \right) \]

Note that, for any node, these computations are local. For all the monitoring algorithms we assume that message transmission is reliable and ordered.

Since $\overline{\mathcal{E}}^\Theta$ is a vector in $\mathbb{R}^d$, the goal reduces to checking if $|\overline{\mathcal{E}}^\Theta| < \epsilon$. However, the quantity $\overline{\mathcal{E}}^\Theta$ is not available at any node. In the next section we state a key result which allow us to perform the same computation using only local vectors of a node.

6.2.2 Stopping Rule  The main idea of the stopping rule is to describe a condition for each node $P_i$ based on $\overline{\mathcal{K}}_i, \overline{\mathcal{A}}_{i,j},$ and $\overline{\mathcal{H}}_{i,j}$, which guarantee that $\overline{\mathcal{E}}^\Theta$ is greater than or less than $\epsilon$. In order to apply this theorem, we need to split the entire domain into non-overlapping convex regions such that the quantity $|\overline{\mathcal{E}}^\Theta| < \epsilon$ has the same value inside each of these convex regions. We denote the set of all such convex regions by $C_\omega$. Geometrically, checking if the L2-norm of a vector is less than $\epsilon$ is equivalent to checking if $|\overline{\mathcal{E}}^\Theta|$ lies inside a circle of radius $\epsilon$. Note that, by construction, the region in which the output is $0$ i.e. inside the circle is a convex region.

Let us denote it by $R_c$. The outside of the circle can easily be divided into convex regions by drawing random tangent lines to form half-spaces denoted by $(R_{h1}, R_{h2}, \ldots)$. The areas uncovered by $C_\omega$ denote the tie regions.

As stated by the Theorem below, if the following condition holds, the node can stop sending messages and determine the correct output based solely on its local averages.

**THEOREM 6.1.** [25] Let $P_1, \ldots, P_n$ be a set of nodes connected to each other over a spanning tree $G(V, E)$. Let $\mathcal{E}^\Theta, \mathcal{K}_i, \mathcal{A}_{i,j},$ and $\mathcal{H}_{i,j}$ be as defined in the previous section. Let $R$ be any region in $C_\omega$. If at time $t$ no messages traverse the network, and for each $P_i$, $\overline{\mathcal{K}}_i \in R$ and for every $P_j \in \Gamma_i$, $\overline{\mathcal{A}}_{i,j} \in R$ and either $\overline{\mathcal{H}}_{i,j} \in R$ or $\mathcal{H}_{i,j} = \emptyset$, then $\overline{\mathcal{E}}^\Theta \in R$.

**Proof.** For proof the readers are referred to [25].

Using this theorem, each node can check if $|\overline{\mathcal{K}}_i| < \epsilon$. If the result holds for every node, then we are guaranteed to get the correct result. If there is any disagreement, it would be between any two neighbors. In that case, messages will be exchanged and they will converge to the same result. In either case, eventual global correctness is guaranteed.

6.3 Algorithm  Both the mean monitoring algorithm and the eigenvector monitoring rely on the results of Theorem 6.1 to output the correct result. For the eigenvector monitoring, the model supplied to each node are the eigenvector $\overline{\mathcal{V}}$ and eigenvalue $\Theta$. Assuming that the mean of the data is zero, the goal is to check if:
\[
\left\| \mathbf{c} \cdot \overline{\mathcal{V}} - \Theta \overline{\mathcal{V}} \right\| \leq \epsilon
\]
\[
\implies \left\| \frac{1}{|G|} \left[ G^T G \right] \cdot \overline{\mathcal{V}} - \Theta \overline{\mathcal{V}} \right\| \leq \epsilon
\]
\[
\implies \left\| \frac{1}{\sum_{i} |M_i|} \sum_{i} \left[ M_i^T M_i \right] \cdot \overline{\mathcal{V}} - \Theta \overline{\mathcal{V}} \right\| \leq \epsilon
\]

Thus given $\overline{\mathcal{V}}$ and $\Theta$, each node can locally compute the vector $\left( \left[ M_i^T M_i \right] \cdot \overline{\mathcal{V}} - \Theta \overline{\mathcal{V}} \right)$. Let this instance of problem be denoted by $I_1$. We can write:
\[ I_1 \overline{\mathcal{E}}_i = \left( \left[ M_i^T M_i \right] \cdot \overline{\mathcal{V}} - \Theta \overline{\mathcal{V}} \right) \]
\[ I_1 \cdot |\mathcal{E}_i| = |M_i| \]

Thus for this problem, each node computes the vector $I_1 \overline{\mathcal{E}}_i$ which is then used as input to the eigenvector monitoring algorithm.

Similarly for the mean monitoring algorithm, the model supplied to each node is the mean $\overline{\mu} \in \mathbb{R}^d$. In this case, each node subtracts the mean $\overline{\mu}$ from its local average input.
vector $\mathcal{M}_i$. The goal is to check if:

$$\left\| \frac{1}{\sum_i |\mathcal{M}_i|} \sum_i \mathcal{M}_i |\mathcal{M}_i| - \mu \right\| \leq \epsilon$$

Note that the quantity $|\mathcal{M}_i| (\mathcal{M}_i - \mu)$ can be locally computed by a node. For this problem instance denoted by $I_2$, the following are the inputs:

1. $I_2, \mathcal{E}_i = (\mathcal{M}_i - \mu)$
2. $I_2, |\mathcal{E}_i| = |\mathcal{M}_i|$

Algorithm 6.1 presents the pseudo-code of the monitoring algorithm while Alg. 6.2 presents the pseudo-code for the algorithm which builds the model. The inputs to the monitoring algorithm are $\mathcal{M}_i, \mathcal{E}_i$ (depending on how it is defined), $\Gamma_i, \epsilon$ and $C_\omega$ and $L$. For each problem instance $I_1$ and $I_2$, each node initializes its local vectors $\overline{K}_i, \overline{A}_{i,j}$ and $\overline{C}_{i,j}$. Below we describe the monitoring algorithm with respect to only one instance $I_1$ and (hence drop the instance index $I_1$).

The other case is identical. The algorithm is entirely event-driven. Events can be of the following: (i) change in local data $\mathcal{M}_i$, (ii) on receiving a message, and (iii) change in $\Gamma_i$. In any of these cases, the node checks if the condition of the theorem holds. Based on the value of its knowledge $\overline{K}_i$, the node selects the active region $R_\ell \in C_\omega$ such that $\overline{K}_i \in R_\ell$. If no such region exists, $R_\ell = \emptyset$. If $R = \emptyset$, then $\overline{K}_i$ lies in the tie region and hence $P_i$ has sent all its data. On the other hand, if $R_\ell \neq \emptyset$ the node can rely on the result of Theorem 6.1 to decide whether to send a message. If for all $P_j \in \Gamma_i$, both $\overline{A}_{i,j} \in R_\ell$ and $\overline{C}_{i,j} \in R_\ell$, $P_j$ does nothing; else it needs to set $\overline{S}_{i,j}$ and $|S|$. Based on the conditions of the Theorem, note that these are the only two cases when a node needs to send a message. Whenever it receives a message ($\overline{S}$ and $|S|$), it sets $\overline{S}_{j,i} \leftarrow \overline{S}$ and $|S_{j,i}| \leftarrow |S|$. This may trigger another round of communication since its $\overline{K}_i$ can now change.

To prevent message explosion, in our event-based system we employ a “leaky bucket” mechanism which ensures that no two messages are sent in a period shorter than a constant $L$. Note that this mechanism does not enforce synchronization or affect correctness; at most it might delay convergence. This technique has been used elsewhere also [25][6].

**Algorithm 6.1. Monitoring Models**

**Input:** $\epsilon, C_\omega, \mathcal{E}_i, \Gamma_i$ and $L$.

**Output:** 0 if $|\overline{K}_i| < \epsilon$, 1 otherwise

**Initialization:** Initialize vectors;

if MessageRecvdFrom $(P_j, \overline{S}, |S|)$ then

$\overline{S}_{j,i} \leftarrow \overline{S}$ and $|S_{j,i}| \leftarrow |S|$

Update vectors

end if

if $\mathcal{M}_i, \Gamma_i$ or $K_i$ changes then

for all Neighbors $P_j$ do

if LastMsgSent > $L$ time units ago then

if $R_\ell = \emptyset$ then

Set $\overline{S}_{i,j} \leftarrow |K_i|$, $|S_{i,j}|$, $|S|$, $|S_{j,i}|$ /\* Tie Region */

Set $|S_{i,j}| \leftarrow |\mathcal{K}_i| - |S_{i,j}|$

end if

if $\overline{A}_{i,j} \notin R_\ell$ or $\overline{C}_{i,j} \notin R_\ell$ then

Set $\overline{S}_{i,j}$ and $|S_{i,j}|$ such that $\overline{A}_{i,j}$ and $\overline{C}_{i,j} \in R_\ell$

/\* Theorem Condition */

end if

MessageSentTo $(P_j, \overline{S}_{i,j}, |S_{i,j}|)$

LastMsgSent $\leftarrow$ CurrentTime

Update all vectors

end if

end for

end if

The monitoring algorithm raises a flag whenever either

$\left\| I_1, \mathcal{E}_i \right\| > \epsilon$ or $\left\| I_2, \mathcal{E}_i \right\| > \epsilon$.

Once the flag is set to 1, the nodes engage in a convergecast-broadcast process to accumulate data up the root of the tree, recompute the model and disseminate it in the network.

For the mean monitoring algorithm in the convergecast phase, whenever a flag is raised, each leaf node in the tree forwards its local mean up the root of the tree. In this phase, each node maintains a user selected alert mitigation constant, $\tau$ which ensures that an alert is stable for a given period of time $\tau$ for it to send the data. Experimental results show that this is crucial in preventing a false alarm from progressing, thereby saving resources. In order to implement this, whenever the monitoring algorithm raises a flag, the node notes the time, and sets a timer to $\tau$ time units. Now, if the timer expires, or a data message is received from one of its neighbors, $P_i$ first checks if there is an existing alert. If it has been recorded $\tau$, or more time units ago, the node does one of the following. If it has received messages from all its neighbors, $P_i$ first checks if there is an existing alert. If it has been recorded $\tau$ or more time units ago, the node does one of the following. If it has received messages from all its neighbors, it recomputes the new mean, sends it to all its neighbors and restarts its monitoring algorithm with the new mean. On the other hand, if it has received the mean from all but one of the neighbors, it combines its data with all of its neighbors’ data and then sends it to the neighbor from which it has not received any data. Other than any of these cases, a node does nothing.

For the eigenvector monitoring, in place of sending a local mean vector, each node forwards the covariance matrix $\mathcal{C}_i$. Any intermediate node accumulates the covariance matrix of its children, adds it local matrix and sends it...
to its parent up the tree. The root computes the new eigenvectors and eigenvalues. The first eigenstate is passed to the monitoring algorithm.

**Algorithm 6.2. Building Models**

**Input:** $\epsilon, C, \mathcal{M}, \Gamma, L, \tau$

**Output:** (i) $\overrightarrow{V}, \Theta$ such that $\|C \cdot \overrightarrow{V} - \Theta \cdot \overrightarrow{V}\| < \epsilon$, (ii) $\overrightarrow{\mu}$

such that $\|\overrightarrow{\mu} - \overrightarrow{\bar{\mu}}\| < \epsilon$

**Initialization:** Initialize vectors;

$\text{MsgType} = \text{MessageRecvdFrom}(P)$

if $\text{MsgType} = \text{Monitoring}_\text{-}Msg$ then

Pass Message to appropriate Monitoring Algorithm

end if

if $\text{MsgType} = \text{New}_\text{-}Model_\text{-}Msg$ {/*Broadcast*/} then

Update $\overrightarrow{V}, \Theta, \overrightarrow{\mu}$

Forward new model to all neighbors

Dataset=false

Restart Monitoring Algorithm with new models

end if

if $\text{MsgType} = \text{Dataset}_\text{-}Msg$ {/*Convergecast*/} then

if Received from all but one neighbor then

flag=Output Monitoring Algorithm()

if Dataset = false and flag = 1 then

if DataAlert stable for $\tau$ time then

$D_1=C_1 + \text{Recvd}_{\text{-}Dataset}$

$D_2=M_1 + \text{Recvd}_{\text{-}mean}$

Dataset=true;

Send $D_1, D_2$ to remaining neighbor

else

DataAlert=CurrentTime

end if

end if

end if

if Received from all neighbors then

$D=C_1 + \text{Recvd}_{\text{-}Dataset}$

$D_2=M_1 + \text{Recvd}_{\text{-}mean}$

$(\overrightarrow{V}, \Theta)=\text{EigAnalysis}(D)$

$\overrightarrow{\mu} = \text{mean}(D_2)$

Forward new $\overrightarrow{V}, \Theta, \overrightarrow{\mu}$ to all neighbors

Dataset=false

Restart Monitoring Algorithm with new models;

end if

if $M_1, \Gamma_i$ or $K_i$ changes then

Run Monitoring Algorithm

flag=Output Monitoring Algorithm()

if flag=1 and $P_i$ is Leaf() then

Execute the same conditions as

$\text{MsgType} = \text{Dataset}_\text{-}Msg$

end if

end if

**6.4 Correctness and Complexity Analysis**

The eigen monitoring algorithm is eventually correct.

**Theorem 6.2.** The eigen monitoring algorithm is **eventually correct.**

**Proof.** For the eigen monitoring algorithm, the computation will continue for each node unless one of the following happens:

- for every node, $\overrightarrow{K_i} = \overrightarrow{E\bar{\sigma}}$

- for every $P_i$ and every neighbor $P_j$, $K_i, A_{i,j},$ and $H_{i,j}$ are in the same convex region $R_\ell \in C_\omega$.

In the former case, every node obviously computes the correct output since the knowledge of each node becomes equal to the global knowledge. In the latter case, Theorem 6.1 dictates that $E\bar{\sigma} \in R_\ell$. Note that by construction, the output of the monitoring function (in this case L2-norm) is invariant inside $R_\ell$. In other words, the binary function $\|E\bar{\sigma}\| < \epsilon$ and $\|\overrightarrow{K_i}\| < \epsilon$ will have the same output inside $R_\ell$. Therefore in either of the cases, the eigen monitoring algorithm is correct.

Determining the communication complexity of local algorithms in dynamic environments is still an open research issue. Researches have proposed definitions of locality [6][25]. Note that for an exact algorithm as the eigen monitoring algorithm, the worst case communication complexity is $O(\text{sizeof network})$. This can happen, for example, when the each node has a vector in a different convex region and the global average is in another different region. However, as shown in this paper and also by several authors [25][6] there are several problem instances for which the resource consumption becomes independent of the size of the network. Interested readers are referred to [5] for a detailed discussion on communication complexity and locality of such algorithms.

**7 Results**

In this section we demonstrate the experimental results of both the centralized fundamental plane analysis and the distributed eigen monitoring algorithm. The centralized experiments show how the fundamental plane changes with variations in galactic density, while the distributed experiments show the performance of the eigen monitoring algorithm for a streaming scenario of the same experiment. Our goal is to demonstrate that, using our distributed eigen monitoring algorithm to compute the principal components and monitor them in a streaming scenario, we can find very similar results as were obtained by applying a centralized PCA. Even though our goal was not to make a new discovery in astronomy, the results are astronomically noteworthy. We argue
that our distributed algorithm could have found very sim-
ilar results to the centralized approach at a fraction of the
communication cost. Also, we want to emphasize that this
distributed eigen monitoring algorithm can be applied to a
number of change-detection applications in high-throughput
streaming scenarios (such as the LSST) for important astro-
nomical discoveries of many types. The importance and nov-
elty of this algorithm compared to existing distributed PCA
algorithms is that, this is an exact algorithm that determinis-
tically converges to the correct result.

7.1 Fundamental Plane Results
As noted in Section 5.1, we divide the entire dataset into 30 bins. The bins are
arranged from low to high density. In this section we present
the results of our fundamental plane experiments for only the
elliptical galaxies for 30 bins.

Our first experiment (Figure 2) shows the variance cap-
tured by the first PC (PC1) as the density of the galaxies
increase. The x-axis shows the mean density of each bin in
log-scale. As seen in the figure, the variance captured by
PC1 increases monotonically with increase in mean galactic
density.

Figure 2: Variance captured by PC 1 w.r.t. log of mean
density of each bin. Bin 1 has the lowest mean density and
Bin 30 the highest. The variance captured by PC1 increases
monotonically with increasing bin density.

Figure 3: Variance captured by PCs 1 and 2 w.r.t. log of
mean density of each bin. Bin 1 has the lowest mean density and
Bin 30 the highest.

Figure 3 provides the most significant scientific result. It
demonstrates the dependence of the variance captured by the
first 2 PC’s with respect to log of bin density. As seen, the
variance increases monotonically from almost 95% to 98% with increase in galactic bin density. This clearly demonstrates a new astrophysical effect, beyond that traditionally reported in the astronomical literature. This results from the application of distributed data mining (DDM) on a signifi-
cantly (by 1000 times) larger set of data. More such remark-
able discoveries can be anticipated when DDM algorithms of
the type reported here are applied to massive scientific (and non-scientific) data streams of the future.

To analyze more deeply the nature of the variation of
the first two PCs with respect to increasing galactic density, we plot the direction of the normal to the plane defined
by the first 2 PCs i.e. pc1 and pc2. Since each of these
PC’s are vectors in 3-d, so is the normal to the plane. The
normal vector is represented by its two directional angles: the spherical polar angles \( \theta \) and \( \phi \). Figure 4 shows a plot of \( \theta \) and \( \phi \) for 30 bins. Figure 4(a) shows the variation of \( \theta \) and \( \phi \) independently with log of mean galactic density. Figure 4(b)
shows the variation of both with log of mean density. The
systematic trend in the change of direction of the normal
vector seen in Figure 4(b) is a new astronomy result. This
represents exactly the type of change detection from eigen
monitoring that will need to be applied to massive scientific
data streams, including large astronomy applications (LSST) and large-scale geo-distributed sensor networks, in order to facilitate knowledge discovery from these petascale data collections.

Figure 5: Quality vs. number of nodes. Quality remains the same thereby showing good accuracy.

7.2 Results of Distributed PCA Algorithm

The distributed PCA implementation makes use of the Distributed Data Mining Toolkit (DDMT)\(^6\)– a distributed data mining development environment from DIADIC research lab at UMBC. DDMT uses topological information which can be generate by BRITE\(^7\), a universal topology generator from Boston University. In our simulations we used topologies generated according to the Barabasi Albert (BA) model. On top of the network generated by BRITE, we overlayed a spanning tree. We have experimented with network size ranging from 50 to 1000 nodes.

We have divided the data of the centralized experiments into 5 bins (instead of 30) sorted by galactic density. Each bin represents the data distribution at a certain time in the streaming scenario and the distribution changes every 200,000 simulation ticks which we call an epoch. This implies that every 200,000 simulation ticks we supply the nodes with a new bin of data. We stream the data at a rate of 10% of the bin size for every 10,000 simulation ticks. The two quantities measured in our experiments are the quality of the result and the cost of the algorithm.

We have used the following default values for the algorithm: size of leaky bucket \(L = 500\), error threshold \(\epsilon = 1.0\), alert mitigation constant \(\tau = 500\). Due to shortage of space we do not present an exhaustive analysis of the effect of all these parameters. We plan to report these in an extended version.

For the eigen monitoring algorithm, quality is the average L2 norm distance between the principal eigen vector and the the computed eigen vector in the distributed scenario over all the bins. Since we compute the principal eigen vector for each bin separately, we plot the average L2 norm distance between the centralized and distributed eigen vectors for every experiment. The experiment was repeated for 10 independent trials. Figure 5 shows the scalability results for the accuracy achieved by our algorithm. As shown in the figure, the proposed eigen monitoring algorithm produces results which are quite close to their centralized counterpart. Moreover, we can also observe that the quality does not degrade with increasing network size. Because our algorithm is provably correct, the number of nodes has no influence on the quality of the result.

Figure 6: L2 messages vs. number of nodes. Number of messages remain constant showing excellent scalability.

Figure 7: Number of convergecast rounds per epoch vs. number of nodes. In most cases the convergecast round is less than 3 per epoch.

Figures 6 and 7 show the number of messages exchanged per node when the number of nodes is increased from 50 to 1000. As shown in Figure 6, the normalized L2 messages per node is approximately 0.3. Normalized message per node means the number of messages sent by a node per unit of leaky bucket. Note that for an algorithm which uses broadcast as the communication model, its normalized messages will be 2.0, assuming two neighbors per node on average. Thus the proposed algorithm is highly efficient with respect to communication. Also as shown, the L2 messages remain a constant even if the number of nodes is increased. This demonstrates the excellent scalability of the algorithm.

Finally, we also plot the number of times data is collected per epoch. In most cases, the number of such convergecast rounds is 3 per epoch. Note that this can be re-
duced further by using a larger alert mitigation constant $\tau$, larger error threshold $\epsilon$ or larger local data set size.

8 Conclusion
This paper presents a local and completely asynchronous algorithm for monitoring the eigenstates of distributed and streaming data. The algorithm is efficient and exact in the sense that once computation terminates, each node in the network computes the globally correct model. We have taken a relatively well understood problem in astronomy — that of galactic fundamental plane computation and shown how our distributed algorithm can be used to arrive at the same results without any data centralization. We argue that this might become extremely useful when petabyte scale data repositories such as the LSST project start to generate high throughput data streams which need to be co-analyzed with other data repositories located at diverse geographic location. For such large scale tasks, distributing the data and running the algorithm on a number of nodes might prove to be cost effective. Our algorithm is a first step to achieving this goal. Experiments on current SDSS and 2MASS dataset show that the proposed algorithm is efficient, accurate, and highly scalable.

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