Abstract

We propose networkmetrics, a new data-driven approach for monitoring, troubleshooting and understanding communication networks using multivariate analysis. Networkmetric models are powerful machine-learning tools to interpret and interact with data collected from a network. In this paper, we illustrate the application of Multivariate Big Data Analysis (MBDA), a recently proposed networkmetric method with application to Big Data sets. We use MBDA for the detection and troubleshooting of network problems in a campus-wide Wi-Fi network. Data includes a seven-year trace (from 2012 to 2018) of the network’s most recent activity, with approximately 3,000 distinct access points, 40,000 authenticated users, and 600,000 distinct Wi-Fi stations. This is the longest and largest Wi-Fi trace known to date. To analyze this data, we propose learning and visualization procedures that extend MBDA. These procedures result in a methodology that allows network analysts to identify problems and diagnose and troubleshoot them, optimizing the network performance. In the paper, we go through the entire workflow of the approach, illustrating its application in detail and discussing processing times for parallel hardware.

Keywords: Multivariate Big Data Analysis, Anomaly Detection, Big Data, Parallel Hardware, Dartmouth Campus Wi-Fi, Networkmetrics

1. Introduction

Multivariate exploratory analysis has been recognized as an outstanding approach in several domains, including industrial monitoring [1], network security [2], marketing [3], weather modeling [4], bioinformatics [5], food research [6], and so forth. In this methodology, visualization, interpretation and data interaction are the principal tools for an analyst to interact and understand data and the problem the data reflects. This is an alternative data-driven approach to the one currently dominating machine learning, e.g., deep-learning, in which the model is built as a black box to approximate an output of interest and little interpretation is left to the analyst.

There are advantages and disadvantages in the approach of multivariate exploratory analysis. A main shortcoming is that the analyst has to be proficient in the modeling approach used and knowledgeable about the type of data under analysis, just like a blacksmith needs to be proficient with the hammer and knowledgeable about the properties of metals. However, the benefit is worth the price. Multivariate exploratory analysis is useful to address specific questions, like “what is the best classification of an individual, given a set of classes”, or “what is the best prediction of a future outcome”, as in other machine-learning approaches. However, a major advantage of the former is that it also provides interpretable information about why a model provides a given answer. There are many situations in which an answer is not of practical use, without knowing the “why”. Network monitoring is an example: network analysts desire to detect unwanted events during network operation, but they also need to understand their root causes and troubleshoot them as soon as possible. An advantage of multivariate exploratory analysis is that, even if a model is created to respond to a specific question, the interaction of the analyst with the data through the model can bring much more information, like the derivation of new, unexpected findings. This is a useful property that black-box models do not normally provide.

In a recent paper, we introduced the notion of networkmetrics [7], the use of multivariate analysis to unravel network problems. The name is a derivation from other areas where multivariate analysis plays a central role in the scientific and professional communities, like in psychometrics, chemometrics or econometrics. In this paper we present an example of what these methods can bring to the network monitoring arena. Specifically, we present a case study of the Multivariate Big Data Analysis (MBDA) tool [8] as applied to monitoring and troubleshooting a campus-wide Wi-Fi network [9]. MBDA
is a complete multivariate anomaly detection and analysis approach, based on a workflow of five steps, that can handle large amounts of data from disparate sources. When an anomaly is identified, the output includes the log entries of raw information associated with it. These, in turn, can be presented to the analyst, so as to elucidate the root causes for the anomaly. In a context where the number of log entries is massive (i.e., Big Data), MBDA works as a magnifier glass, conveniently highlighting anomalous events.

Our contributions in this paper are as follows.

- We illustrate the application of MBDA to a real case study, showing what it can provide to network analysts and presenting the workflow in detail, including the parallelization of the code and processing time results.
- We enhance MBDA with several functionalities useful in real-life practice:
  - We propose an automatic feature-learning procedure, consistent with the MBDA methodology.
  - We incorporate state-of-the-art exploratory analysis visualizations within the central step of MBDA, to make it more data interactive.
  - We use gephi [10] network visualizations to improve the interpretation of diagnosis results, which are supposed to be condensed [8] – but can still be of large size in practical problems.
  - We discuss model updating after anomaly identification and diagnosis, to further refine the anomaly detection.

The rest of the paper is organized as follows. Section 2 introduces the data set under analysis. Section 3 presents the MBDA methodology in brief. Section 4 introduces the learning procedure contributed. Section 5 illustrates the five steps of MBDA in the Wi-Fi data. Section 6 presents the visualization on the anomalies and Section 7 discusses model update. Section 8 provides final conclusions.

2. The Dartmouth Wi-Fi network

Dartmouth College has a compact campus with over 200 buildings on 200 acres. The evolution of the network is documented in the series of papers [11, 12, 9]. The number of students, staff, and academic faculty reached near 6,500, 3,300 and 1,000, respectively, at the end of 2018, and the number of Access Points (APs) is above the 3,000. Researchers at Dartmouth have been capturing data about the usage of the network for many years, providing a perfect case study for tools like MBDA.

The paper analyses a data capture containing the connections of users to the network in a seven year time span: from 2012 to 2018. Data contains Simple Network Management Protocol (SNMP) traps [13] sent from wireless controllers to our collector. The capture reveals the statistics in Table 1. The data set contains a total of 5 Billion traps and 7 TB of data. A total of 38K authenticated users and an undetermined number of non-authenticated users have been connected to the network in the last seven years, using 600K devices. The network infrastructure supports several SSIDs, primarily Dartmouth Secure, the WPA2-Enterprise authenticated college network, Dartmouth Public, a public-access network, and eduroam, the world-wide roaming network for educational institutions [14]. Dartmouth Secure was entirely replaced by eduroam in the final months of the capture.

<table>
<thead>
<tr>
<th>Statistic</th>
<th>Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Capture period</td>
<td>Jan 1st 2012 - Dec 31st 2018 (2556 days)</td>
</tr>
<tr>
<td>log entries (SNMP traps)</td>
<td>5 Billion</td>
</tr>
<tr>
<td>Data Size (raw)</td>
<td>7 TB</td>
</tr>
<tr>
<td>Access points</td>
<td>3,330</td>
</tr>
<tr>
<td>Authenticated Users</td>
<td>38,096</td>
</tr>
<tr>
<td>Stations</td>
<td>624,903</td>
</tr>
<tr>
<td>SSIDs</td>
<td>20</td>
</tr>
</tbody>
</table>

To collect the trace, the Wi-Fi network controllers forwarded SNMP traps with a record of network activity to the Dartmouth team’s servers. Figure 1 shows an example of an SNMP trap as received. Each trap comprises a header, with timestamp and sender and collector information, followed by a variable number of triplets representing SNMP object identifiers (OIDs) with the format ‘<OID> = <type>: <value>’ and separated by hashes (#). OIDs are partly represented in ASN.1 notation, which can be translated into more meaningful OID names using the relevant Management Information Base (MIB). An important OID is the trap type (TT), in which the value is also an OID: ‘<TT> = OID: <OID>’. More details on the data capture can be found in [9].

3. Multivariate Big Data Analysis

We base our analysis on the Multivariate Big Data Analysis (MBDA) methodology [8], a recently proposed networkmetrics exploratory tool. MBDA makes use of two open software packages available on Github: the MEDA Toolbox [15, 16] and the FCPParser [17]. The FCPParser is a python tool for the parsing of both structured and unstructured logs. With the MEDA Toolbox, multivariate modeling and data visualization can be performed.

Intensive parsing requires a parallel processing hardware. We used the Anthill Compute Cluster hosted by the Computer Science Department at Dartmouth. It is a 100 node, 1200 core, 4,288GB distributed ram compute cluster, managed with the grid engine [18] as parallelization software. Matlab and Python scripts using the FCPParser and the MEDA Toolbox, respectively, run on top of the parallel hardware as grid jobs.

The MBDA approach consists of 5 steps:

1) Parsing: the raw data coming from structured and unstructured sources are transformed into quantitative features.

Table 1: Details of the SNMP trap capture at Dartmouth College.
2) Fusion: the features of the different sources of data are combined into a single data stream. In the example under analysis there is a single source of data: SNMP traps. Thus, fusion is not required.

3) Detection & Analysis: featured data is visualized and anomalies are identified in time using Principal Component Analysis (PCA) [4, 19] and Multivariate Statistical Network Monitoring (MSNM) [20, 21, 22, 23].

4) Pre-diagnosis: the features associated with an anomaly are found.

5) De-parsing: Using both detection and pre-diagnosis information, the original raw data records related to the anomalies are identified and presented to the analyst.

The first three previous steps are equivalent to what it is commonly done in other machine learning methodologies. However, steps 4 and 5, which perform the diagnosis of the anomalies, are a main advantage of the present proposal. These steps are possible thanks to the white-box, exploratory characteristics of PCA as the core of the MSNM approach in step 3). PCA is easy to interpret in terms of the connection between anomalies and features.

PCA transforms the original features into a lower number of uncorrelated features: the so-called principal components. The principal components are ordered by captured variance. PCA follows the expression:

$$ X = T_A \cdot P_A + E_A, $$

where $X$ represents the matrix of data, with $N$ rows and $M$ columns, $T_A$ is the $N \times A$ scores matrix containing the projection of the objects in the principal components sub-space, with $A$ the number of principal components, $P_A$ is the $M \times A$ loadings matrix containing the linear combination of the variables represented in each of the principal components, and $E_A$ is the $N \times M$ matrix of residuals.

Scores, loadings and residuals can be visualized using line and scatter plots to gain data understanding. Also, the data can be further compressed in a pair of statistics, the D-statistic and Q-statistic, where anomaly detection can be performed following the MSNM approach. The D-statistic and the Q-statistic for observation $n$ are computed with the following equations:

$$ D_n = t_n \cdot (\Sigma_T)^{-1} \cdot t_n^T $$

$$ Q_n = e_n \cdot e_n^T $$

where $t_n$ is a $1 \times A$ vector with the scores for observation $n$, $e_n$ is a $1 \times M$ vector with the residuals, and $\Sigma_T$ represents the covariance matrix of the scores.

As a summary, table 2 describes the general data pipeline. Steps 1) and 3) perform two steps of compression of the data, first from raw data to features and then from features to components/residuals using PCA and from these to statistics using MSNM. Once anomalies are found, steps 4) and 5) allow to identify the root cause in the raw information associated to them.

4. Learning Counts in Big Data

4.1. Feature-as-a-counter parsing

MBDA makes use of the feature-as-a-counter (FaaC) approach [2] in step 1). Each feature contains the number of times a given event takes place during a pre-defined time interval. Examples of suitable features are the counts of a given word in a log or the number of traffic flows with given destination port in a Netflow file. This general definition makes it possible to integrate, in a suitable way, most sources of information.

To implement the FaaC, the FCParse defines variables and features. Variables represent general entities in the data. In the previous two examples, the variables would be word and destination port. The Features are defined for a specific value of a variable. Examples of features would be word=’food’ and destination port=’80’). This representation in variables and features has the relevant advantage that allows for the definition of default features, e.g. word=<ANY >, useful to count the instances of a variable, regardless its value, in a data record. Variables and features are defined using regular expressions in configuration files. The FCParse applies this configuration to the data, in order to compute feature vectors for each interval of time. This is done using a multi-threading configuration to speed-up computation.
The default features in step 1) of MBDA play a similar role as residuals do in PCA and MSNM in step 3). Both transformations, steps 1) and 3), work as lossy compression steps, which are fundamental to visualize data when the data volume is massive. However, when doing anomaly detection, it is a good idea to include in the model a summary of what is left out in the compression. This allows the analyst to retain the ability to find uncommon patterns in the residual part, and it is the goal of the compression. This allows the analyst to retain the ability to find uncommon patterns in the residual part, and it is the goal of the residuals in eqs. (1) and (3).

A simplistic example of when default features can be useful follows. Imagine we capture traffic from a network where the main services are http (destination port 80) and smtp (destination port 53). To monitor the traffic, we define two features: destination port=’80’ and destination port=’53’, and a default feature destination port=< ANY >. During traffic monitoring, there is a sudden burst of Internet Relay Chat (IRC) traffic with destination port 6667. The two previous features cannot capture this burst, but the default one can. If something anomalous is detected in a default feature, the analyst can decide whether she needs to redefine features to account for the new situation, or simply look at the raw data for diagnosis.

### 4.2. Learning procedure

MBDA relies in the manual definition of the features in the configuration files of the FCParser. To write such configuration files, the analyst needs to get familiarized with the data. Unfortunately, in a practical Big Data problem like the one under analysis, the data capture is too massive for direct inspection. If we want to obtain a good description of the content, we need to apply an automatic feature derivation technique. This technique needs to be consistent with the posterior multivariate analysis, so that we minimize loss of information.

There are two basic properties we would like to meet in the learning procedure. First, main sources of variance need to be captured. Second, uncommon characteristics with low variance should also be modeled somehow, in a summary of residual information. The second feature is built-in in the FaaC methodology with the definition of default features, as already discussed.

We developed a learning algorithm to automatically identify a list of common FaaC features in a Big Data set, and included it in the FCParser repository at Github with the name `fclearning.py`. The learning algorithm extracts the features in the data ordered by their percentage of presence, measured as the portion of the log entries where a feature appears. It also defines default features and automatically writes the configuration files.

We used this algorithm in two steps to identify high variance OIDs in the Wi-Fi data. First, the algorithm was parallelized in 2556 processing jobs, one per different day in the capture. The resulting 2556 configuration files contain the FaaC features with a percentage of presence above the 5% during the day. Second, these configuration files were combined in a single configuration file, where we discarded all features with variance below a threshold\(^1\). This resulted in a total of 90 features, including default features. From their description we can conclude that we can use this data to identify who associated to the network, when the association took place, the APs involved in the connection, and thus the approximate location and movement of the user during it, and the device used.

The whole learning process using the parallel hardware and multi-threading (4 threads per processor) took 12 hours, during which a maximum of 150 jobs were processed in parallel. This means that the processing time could be reduced in 17-folds using a larger computer cluster. The combination of results in a regular server took another 30 minutes.

### 5. MBDA in action

In this section, we discuss the application of MBDA to the Wi-Fi data set using the learned features. Since there is one single source of data, the second step (fusion) is not necessary.

#### 5.1. Parsing

We use the FCParser to generate the feature vectors with the aforementioned configuration file learned. In agreement with the learning phase, we consider feature vectors for intervals of one day. These contain the number of traps, the total number of OIDs and the number of learned features. This makes a total of 92 features. Each day in the original SNMP capture is transformed into a feature vector, we call "observation", and each feature in the observation is the number of times the corresponding element is found in the traps of that day. This results in a compression of the data from 7TB to less than 1MB, yielding 2556 observations (days) of 92 features each in matrix X. The compression conveniently transforms a Big Data set into a handleable data set by any analysis package in a common computer.

The parsing was again parallelized in 2556 processing jobs, one per day, and the whole process using the Anthill Computer

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\(^1\)We used a threshold of 0.01%, taking the variance in the number of log entries per day as a reference
Cluster and multi-threading (again 4 threads per processor) took 15 hours. Again, a maximum of 150 jobs were processed in parallel, so the same reduction with a larger cluster discussed for the learning procedure applies here.

5.2. Detection & Analysis

Basically, PCA factorizes the data $X$ into a part for the observations (the scores) and a part for the features (the loadings), making the visualization simpler and interpretable. The remaining of this section discusses how to obtain and interpret this factorization. PCA has a number of interesting properties in terms of visualizing the data. First, the principal components are linear combinations of the original features and uncorrelated. In particular, distances between points in any scatter plot of the scores, commonly called a score plot, approximate the Mahalanobis distance in the original data. Second, since the principal components are ordered by variance; hence, we can be confident that the main patterns of high variance in the data will be observable in the first components. Therefore, by only inspecting a small set of principal components, we can gain an accurate insight into the data.

The number of principal components to use has to be determined. There are many methods to aid in that decision, and for a general description on which one is best, see [19, 24]. Fortunately, the data exploration is barely affected by selecting different numbers of components, specially because we also visualize a summary of the residuals. If in doubt, it is better to use larger numbers, since we may learn something else from additional principal components.

Still, it is useful to have an initial reference for a suitable number of principal components. The MEDA Toolbox, the software used in this third step of MBDA, provides of a plot with the residual variance per PC combined with the columnwise k-fold cross-validation (ckf) curve [25]. The corresponding plot for the Wi-Fi data is shown in Figure 2. To obtain this plot, we inputted the matrix $X$ of parsed data. The residual variance tells us the percentage of information not captured by the model, which should be low enough so that we do not miss much information in the summary of the residuals. The ckf gives information about the predictive ability of components, in terms of how well the structure among the original features is captured. A reasonable choice for the number of principal components should capture a large portion of variance and yield a reasonably good predictive model. In our case, 6 principal components seems like a reasonable choice, since models beyond that number do not improve in prediction ability and add very little variance.

Once we have selected the number of principal components, we can visualize the data using scatter plots of scores and loadings. A toy example of score plot is shown in Figure 3. We can see six points representing observations (rows in the data) scattered around the center of coordinates. We assume that data is mean centered, as data is commonly preprocessed before PCA, and note that interpretation changes for non-centered data. The interpretation is easier if we think of the observations not only as points, but also as vectors connecting the origin of coordinates to the points, as in the illustration. The relationship between observations is revealed by the distance between observations and the angle between vectors. Very close points reflect similarity of the observations in the principal components we are visualizing. However, we are mainly interested in those points far from the origin of coordinates, meaning that they present high scores. Observations with low scores are not well modeled by the inspected components. This is illustrated with observation 1, for which the plot does not provide any useful information other than it has a low score in the first two principal components. Vectors with the same direction indicates similarity, and the closer the points the more similar. This is the case for observations 2 and 6. Thanks to this property, score plots are useful to identify clusters of similar observations and other meaningful patterns, like trends and outliers. Vectors in the opposite direction show an inverse relationship, like observations 4 and 5. Vectors in different directions and distanced reflects they are somehow different (not directly or inversely related). Thus, observation 3, the group of 2 and 6 and the group of 4 and 5 are all different. This property is again useful because if the score plot highlights different groupings of observations, we can conclude that there is an underlaying different information content in the features of these groups. To obtain more information about those differences, we need to look at the co-

![Figure 2: Curves of residual variance and ckf [25] for the Wi-Fi data.](image)

![Figure 3: Illustration of interpretation in score and loading plots.](image)
responding loading plot.

Loading plots are similar to score plots, but the former represent the spacial distribution of features (columns in the data) instead of observations. Again, clusters, trends or outliers can be identified in the loading plot, and lead to interesting conclusions on the features distribution. We can also combine scores and loadings in a single plot, commonly called a biplot [26]. Well designed biplots allow similar comparisons in distance and angle between observations and features. Thus, if one observation is located close to a feature in the biplot, we expect this observation to have a high value of that feature. This is useful to draw connections between the patterns of observations and features: e.g. to identify what features make an outlier different from the rest of observations.

The plots corresponding to the first 6 principal components in the Wi-Fi data are depicted in Figure 4. Recall that matrix $X$ contains 2556 rows, representing days of the capture, and 92 features. We present score plots at the left of the figure and loading plots at the right. Visualizing score plots and loading plots together is useful to take a bird’s view of the data. In the score plots, points represent the 2556 days of the data capture and are colored according to the year. In the loading plots, points represent the 92 features and, in order to facilitate interpretation, we also plot a shadow of the scores, like in a biplot.

Figure 4(a) represents the score and loading plot for PC1 vs PC2. These two principal components represent 83% of the variance in the data, and therefore we can learn from them the main patterns of change in $X$. The score plot at the left shows that the dots with different colors are in different locations. This means there are relevant differences in the contents of the SNMP traps for different years. The loading plot shows that a large majority of the features are located far from the center of coordinates towards the right side. This can be interpreted in connection with the score plot: any day toward the right will have higher value in most of the features. Given that the features represent counts of events in the log entries, as we traverse from left to right in the score plot, the days will have more connection activity. Thus, busy periods are represented towards the far right of the plot, and vacations are clustered to the left, and we could say that the first PC (the horizontal direction in the score and loading plots) represents the general activity in the network. We annotated this in both plots using a horizontal arrow.

The loading plot in Figure 4(a) also shows that the variables are distributed from the bottom to top, and we see a similar distribution for the different years in the score plot: the first two years are in the bottom and the last two in the top, with middle years in between. We also see a separated cluster of days in 2018, highlighted with a circle. A closer look reveals that all the days in the cluster belong to the period from September to November, when eduroam replaced Dartmouth Secure. The vertical pattern in the loading and score plots shows that the distribution of traps has changed across the years: days towards the top have a higher content of traps represented by the features in the top and less of those in the bottom, and vice-versa. Again, we annotated this in the score and loading plots using a vertical arrow. Questioned about this difference, the network staff replied that there was an update in the controllers software, in which the types of traps that were collected changed. This variability in traps for different temporal periods makes the analysis of the data a real challenge.

Figure 4(b) represents PC3 vs PC4. Here we see again some differences among the years and the cluster of 2018. We also see a set of consecutive days from 2017 that depart from the rest of days, which is reflecting that something unusual took place during those days. This is sometimes referred to as an excursion of scores. Another 2-day excursion is shown in 2013. Both excursions are annotated with an arrow. The corresponding loading plot is difficult to interpret in connection with the observed deviations, but there are additional tools [27, 28, 29] that can be used to provide more information. This will be shown in the next step of the approach: the pre-diagnosis.

Figure 4(c), representing PC5 vs PC6, shows again the excursion of 2017 and another in 2012 and 2014, annotated with arrows. The loading plot allows to associate the excursions of 2012-2014 and 2017, also annotated with arrows, to specific features, also annotated with circles. The excursions of 2012-2014 are associated to a high number of failures in the RADIUS server, and the one in 2017 to a high number of restarts of APs. We will gain more detail afterwards.

No additional information was revealed from inspecting the following two principal components (not shown), so we decided to stop the initial analysis here.

Besides the inspection of score and loading plots, one can visualize a summary of the whole data distribution in one single plot using MSNM: a scatter plot of the observations in terms of the D-statistic and the Q-statistic. In this plot we can also show upper control limits (UCLs) to facilitate the detection of anomalies. UCLs leave below normal observations with a certain confidence level, e.g. 99%. They can be used as hypothesis tests, so that all observations above the limits reject the null hypothesis that they are normal. More information on how to compute these limits can be found in [22, 20].

The MSNM plot for the Wi-Fi data is shown in Figure 5. Anomalies are expected to surpass any of the two control limits. This plot is optimized for anomaly detection, and the excursions mentioned before are clearly observed. However, in the plot we miss other details, like the yearly and seasonal patterns, as well as the difference in traps contents. A main advantage of this plot is that it also includes residuals, which are not accounted for in the 6 principal components. The Q-statistic, which conforms a summary of the residuals, clearly identifies the excursion in 2013 and another anomaly in 2012.

The analysis is completely interactive in a regular computer, meaning that the processing time to obtain each of the plots shown in this section is in the order of seconds.

5.3. Pre-diagnosis

The plots discussed in the previous step provide a general view of the data and can assist in detecting patterns, like trends, outliers, excursions or clusters. However, if we want to have a closer detail on the interaction observations-features, multivariate diagnosis tools are more accurate and easier to interpret.
There is a large number of multivariate diagnosis tools, see [30, 31] for a comparison. The MEDA Toolbox includes the oMEDA plot for that purpose. It is a bar plot of the features, built to compare two groups of observations: e.g., observations in an excursion from normal observations. Each bar represents the contribution of the feature to the difference between both groups. A positive bar implies that the first group of observations presents a higher value in the corresponding feature than the second group. A negative bar reflects the opposite. A bar close to zero for a feature means that both groups of observations have a similar value in that feature.

To illustrate the use of oMEDA, we selected the excursions in 2013 and 2017, which were shown as the main outliers in the Q-statistic and in the D-statistic, respectively. The plots are
shown in Figure 6. Both of them reflect that each of the excursions are related to a higher value in a different set of features. The specific features are listed in Table 3. We determined that the first excursion is related to a large number of Authentication Fails, one order of magnitude higher than usual. The second excursion is related to an unprecedentedly high number of re-starts of APs, two orders of magnitude higher than usual.

As for 2018, the network staff did not have any registers for these old anomalies, but they suggested that the second one could be related to the installation of a security patch after the publication of a vulnerability. Effectively, October 16th of 2017, the famous KRACK attack against WPA2 [32] and the corresponding patch were released to the public. Even if a restart is necessary after a patch installation, the number and duration (15 days) of the event is remarkable, evidencing that a major management problem took place.

Again, the pre-diagnosis step is in the order of seconds and easily done in a regular computer.

5.4. Deparsing

While the visualizations already provided to the analyst are very informative, it is a good idea to identify the raw log entries related to the patterns found, in order to obtain more detail about them. Examples of findings that may be interesting for us are the specific APs and stations involved in the large number of authentication failures in 2013, or the APs involved in the long period of restarts in 2017. To obtain more detail, the FCParse is employed again in the deparsing step.

The deparsing algorithm [8] takes as inputs the timestamps of a given anomalous pattern, detected in step 3), and the features associated with it, identified in step 4). With this information, the FCParse matches the regular expressions of the features in the specific raw data files. The output is the set of raw log entries that matches at least one of the features, ordered by the number of features they match. With these log entries, the analyst can extract detailed information about the anomalies.

The deparsing algorithm was applied to the excursions in 2013 and 2017 in the Wi-Fi data set. We parallelized the processing using the Anthill Computer Cluster and multi-threading (4 threads per processor), an with as many parallel jobs as days in the excursions. The first excursion took 30 minutes to be processed, and the second one 135 minutes. Some statistics of the deparsing are provided in Table 4. We can see that the number of log entries and nodes involved in the excursion is very large. This means that it is hopeless to expect the analyst to manually inspect this data.

6. Visualization

Given the large number of log entries in the deparsing information, it is hard to extract useful information of the nodes (devices, APs) involved in the anomalies. To improve the detail of the analysis, we combined the output of the deparsing with state-of-the-art visualization techniques. Given that the data represents connections, Gephi [10] seems a suitable tool to visualize the deparsed data. Notice that even if Gephi can be applied to very large sets of connections and nodes, we can only visualize a little portion of the Wi-Fi Big Data set. Thus, the integration of MBDA and Gephi is necessary, and we make the most of the advantages of both approaches.
Table 3: Pre-diagnosis of the excursions of 2013 and 2017 with oMEDA.

<table>
<thead>
<tr>
<th>Timestamps</th>
<th>Features selected</th>
</tr>
</thead>
<tbody>
<tr>
<td>2013-12-14 – 2013-12-16</td>
<td>bsnDot11StationAuthenticateFail, bsnAuthenticationFailure, bsnDot11StationAssociateFail, bsnStationReasonCode, bsnAuthFailureUserType, bsnAuthFailureUserName</td>
</tr>
<tr>
<td>2017-10-16 – 2017-10-30</td>
<td>ciscoLwpAppIfUpNotify, ciscoLwpAppIfDownNotify</td>
</tr>
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</table>

Table 4: Deparsing of the excursions of 2013 and 2017 with oMEDA.

<table>
<thead>
<tr>
<th>Timestamps</th>
<th>log entries/tot</th>
<th>#APs</th>
<th>#Stations</th>
<th>#Users</th>
</tr>
</thead>
<tbody>
<tr>
<td>2013-12-14 – 2013-12-16</td>
<td>5.4M/8.4M (64%)</td>
<td>824</td>
<td>595</td>
<td>103</td>
</tr>
<tr>
<td>2017-10-16 – 2017-10-30</td>
<td>19.0M/64.1M (30%)</td>
<td>1,376</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

7. Model update

Once we have identified, diagnosed and troubleshooting a number of anomalies, we would like to correct the model to remove their effect. This will give us the opportunity to focus on additional, less distinct, anomalies, and also to use the model for the monitoring of future days (e.g. with MSNM).

We can think of two approaches to extract the already analyzed anomalies from the model. One choice is to discard the raw log entries de-parsed for the anomaly. This requires little processing time. We will call this approach log-wise extraction.

Another possibility, which requires even less processing, is discarding the complete feature vector corresponding to the anomalous observation. This has the implication that some non-anomalous raw log entries will also be discarded in the process.

We will call this approach observation-wise extraction.

In Fig. 9, the score plot for PC1 vs PC2 in the original model is compared to the models after log-wise and observation-wise extraction of the two anomalies analyzed in previous section. We can see that the log-wise method can have unexpected consequences. In our case, the two anomalies reflect a general network failure that prevented common connections from taking place. After extracting the anomalous log entries, the remaining activity is lower than in other regular days, and for this reason the observations deviate from the rest. If we extract the complete days, this problem is obviously avoided.

Fig. 10 compares the MSNM plots after both forms of extraction. The log-wise extraction leads to detect the same anomalies as before, which interfere with the performance of the method to detect new anomalies. Contrarily, the observation-wise extraction cleans the model and allows the identification of new anomalies, which can be subsequently diagnosed, deparsed, and visualized.

Clearly, this example points to the use of observation-wise extraction as the model update rule. However, it should be noted that if the anomaly has not such leverage on the log entries of a day, log-wise extraction may be suitable and even preferred.

Table 5 lists anomalies found in two iterations of MBDA. We can see that AP re-starts and RADIUS server problems are recurrent. Some of these may anomalies point to maintenance operations, in which case the analyst may directly discard the day in the monitoring model. More iterations lead to more anomalies.
8. Conclusion

In this paper, we show a case study of the application of Multivariate Big Data Analysis (MBDA) tool, a networkmetric tool optimized to analyze Big Data streams. The application is concerned with the detection, diagnosis and troubleshooting of communication failures in a Wi-Fi campus network. The results illustrate that multivariate methods can bring light into complex massive data sets.

MBDA can work on top of parallel hardware in order to speed up computation. We analyzed 7TB of data in a little more than a day, and this can be reduced to a couple of hours in a high-throughput cluster. While parallel computations are required to analyze Big Data, interaction with the data can be done in a regular computer, combining the advantages of interactive models with the power of parallel processing.

The core of MBDA in this paper is Principal Component Analysis. However, the MBDA framework can be easily extended to other exploratory models, more powerful to analyze specific data sources, like sparse methods, regression & classification methods, n-way models, constrained (e.g. non-negative) methods and many others.

Acknowledgement

This work was supported by Dartmouth College, and in particular by the many network and IT staff who assisted us in configuring the Wi-Fi network infrastructure to collect data, and who patiently answered our many questions about the network and its operation. We furthermore appreciate the support of research colleagues and staff who have contributed to our data-collection and data-analytics infrastructure over the years: most notably Wayne Cripps, Tristan Henderson, Patrick Procotor, Anna Shubina, and Jihwang Yeo. Some of the Dartmouth effort was funded through support from ACM SIGMOBILE and by an early grant from the US National Science Foundation un-
Figure 8: Gephi: AP re-starts in the period 2017-10-16 – 2017-10-30. For reference, the regular amount of re-starts is two orders of magnitude lower and would not be perceptible in this plot.

This work was also supported by the Ministerio de Educación, Cultura y Deporte under the Programa Estatal de Promoción de Talento y su Empleabilidad en I+D+i, Subprograma Estatal de Movilidad, del Plan Estatal de Investigación Científica y Técnica y de Innovación 2013-2016, grant number PRX17/00320 (Associated to a Fulbright Scholarship), and the Plan Propio de la Universidad de Granada, grant number PP2017.VS.02. Jose Manuel García-Giménez is acknowledged for his enthusiastic work on the FCParse.

References


Figure 9: PCA scores PC1 vs PC2: original model (a), after log-wise extraction of outliers (b) and after observation-wise extraction (c).

Figure 10: MSNM plot after extraction of outliers: log-wise extraction (a) and observation-wise extraction (b).


