AUTOENCODED REDUCED CLUSTERS FOR ANOMALY DETECTION ENRICHMENT (ARCADE) IN HYPERSPECTRAL IMAGERY

THESIS

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DEPARTMENT OF THE AIR FORCE
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AIR FORCE INSTITUTE OF TECHNOLOGY

Wright-Patterson Air Force Base, Ohio

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AUTOENCODED REDUCED CLUSTERS FOR ANOMALY DETECTION ENRICHMENT (ARCADE) IN HYPERSONTIAL IMAGERY

THESIS

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Department of Operational Sciences
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Air Education and Training Command

In Partial Fulfillment of the Requirements for the
Degree of Master of Science in Operations Research

Brenden A. McLean, BS
Captain, USAF

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Abstract
Anomaly detection in hyper-spectral imagery is a relatively recent and important research area. The shear amount of data available in many hyper-spectral images makes the utilization of multivariate statistical methods and artificial neural networks ideal for this analysis. Using HYDICE sensor hyper-spectral images, we examine a variety of pre-processing techniques within a framework that allows for changing parameter settings and varying the methodological order of operations in order to enhance detection of anomalies within image data. By examining a variety of different options, we are able to gain significant insight into what makes anomaly detection viable for these images, as well as what impact parameter and methodology changes can have on the total classification effectiveness, false positive fraction and true positive fraction regarding classification.
I want to express my love and devotion to my wife and my three boys. Their support and encouragement in every area of life is beyond compare. I want to give full acknowledgement that everything good I have in life is from my Father in heaven through His son Jesus the Christ.
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Brenden A. McLean
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I. Introduction

General Issue

How does one determine when outliers or anomalies exist within data? Once outliers and anomalies are found, what confidence does one have that it actually represents an anomaly and what confidence does one have that it represents an anomaly of interest? In simple problems, we can often attain insight into whether a data point is anomalous by a plotted observation of the data or through relatively simple sets of calculations. However, when data reaches complexity beyond the capacity of the human mind, as is the case with hyperspectral imagery (HSI), this process becomes substantially more difficult.

For this particular problem set, we examine HSI data, which represent images taken utilizing a substantially broader region of the electromagnetic spectrum than normal photographs. Studies have examined these types of images since at least the 1970s, examining images to identify minerals [1]. The military began considering multispectral imagery in the 1980s for topography and terrain analysis [2]. This proceeded into the recognition that with high quality images the possibility arose for remote sensing and target acquisition [3], [4]. This eventually led to the applications of anomaly detection regarding the HSI images. Currently the field is pushing toward ever greater utility in finding these anomalies and the issue of finding the true, real world, anomalies within the HSI images remains.
**Problem Statement**

Specifically, our problem rests with whether military targets can be acquired utilizing HSI to determine threats within a particular area (the image). Upon determining the anomalies within a particular area, we desire to have a high degree of certainty that the anomaly detected is an anomaly warranting action. We wanted to develop a new algorithm that would decrease the false positive identification of an anomaly within an image.

**Research Objectives/Questions/Hypotheses**

While the specific problem of anomaly detection in HSI has been assessed previously [5]–[7], our objective is to design and test a new anomaly detection procedure that utilizes a combination of different algorithms and Artificial Neural Networks (ANNs) to improve upon the preprocessing of other anomaly detectors. Upon designing a functional algorithm, this would be the first time in this research stream that ANNs have been employed for anomaly detection.

Our research was specifically tested with HSI from the Hyperspectral Digital Imagery Collection Equipment (HYDICE) sensor [8]. We utilized blocked adaptive computationally efficient outlier nominators (BACON) [9], principal component analysis (PCA), k-means clustering, autoencoded ANNs, and reconstructive error calculations to zero in on the anomalies with the data, but in doing so, it became necessary to explore the appropriate parameters and settings of those algorithms. Our primary hypothesis that through a combination of a variety of dimensionality assessment and reduction techniques as well as a combination of data evaluation techniques, one can determine the
true anomalies in highly dimensional, enormous data arrays with a high degree of certainty, such that actionable determinations can be executed.

**Investigative Questions**

Our primary focus was to determine if this set of methods could be combined to more accurately detect and confirm the existence of critical anomalies within the aforementioned images. Specifically, we desired to confirm whether the algorithmic combination of these methods with defined parameters could be applied broadly to sets of images for anomaly detection with confidence. By examining a range of parameters, we hoped to get insight into the optimal settings of these parameters for the identification of anomalous data.

**Methodology**

By designing an experiment and combining a variety of both anomaly detection techniques and dimensionality reduction techniques, we hope to best determine outliers within HSI images that coincide with known anomalous objects within those images. In order to do this, we first reshape the image cube to fit on a single plane, and then we pre-process the image utilizing existing outlier detectors. This provides us with a set of pre-identified potential outliers based on specified criteria and parameters for the pre-processing algorithm. Essentially the goal of the initial procedure is to separate the specified images into two meta-classes, one which we know to contain background information and another which would primarily contain the potential anomalies. First, we complete PCA to re-dimensionalize, orthogonalize, and center the data using the meaningful scores as determined using Maximum Distance Secant Line (MDSL) [10].
This allows us to use a clustering algorithm to find like classes within the background class of image. Once the background class is clustered we autoencode each cluster employing ANNs. We are able to choose the best ANN by its performance as assessed in MATLAB. Thereby, we filter the anomalous class through each cluster’s optimal ANN, obtain reconstructive errors for each observation, and assess each point’s validity as an anomaly by assessing whether the point should belong to one of the clusters associated with the background class.

**Assumptions/Limitations/Implications**

Based on the literature, we assume the validity of each methodology for its specified purpose, when it is implemented in a vacuum free of other methods. We assume that the current truth data is accurate; therefore, a valid basis for comparison.

The primary limitation is computational time and efficiency. While we are testing ways in which to enhance this limitation, a hyperspectral image still represents a vast amount of data, and in order to process the entire methodology, it does represent a cumbersome process in terms of time.

Effectiveness of the method means an enhanced method for identifying potential outliers and assessing the validity of being concerned about those specified outliers. It further means that we can improve or validate pre-processing by other algorithms. Lastly, it indicates the viability of utilizing ANNs as an instrument for anomaly detection. This has the potential to greatly enhance anomaly detection as whole throughout a variety of fields and applications.
Overview

In Section II, we conduct a literature review of the various methodologies and techniques we employ, as well as other techniques associated with the field that have impacted anomaly detection, HSI, and multivariate statistical analysis. Section III provides a more in-depth view of our specific methodology, showing how the algorithms combine to form our solution space. We cover our analysis and results in Section IV describing the solution space and its implications. Finally, in Section V, we provide our conclusions based on the results of our research and analysis. In this final section, we also try to postulate on potential future research, such as the comparison of other additional pre-processes, improvements in computational efficiency, expansion to other images or data sets, and confidence enhancements.
II. Literature Review

Chapter Overview

Initially, as we began to examine our problem set our focus was on feature selection. Over time this shifted to hone in more specifically on anomaly detection, utilizing some of the ideas associated with feature selection. For the literature review, we examine anomaly detection, hyperspectral imagery (HIS), feature selection, multivariate techniques, principal component analysis (PCA), factor analysis, discriminant analysis, multiple outlier detection, clustering and artificial neural networks (ANNs).

Relevant Research

Over the years, it seems that just about every area where data exists has been explored. Bauer, Alsing, and Greene examined University of Wisconsin breast cancer Data, US Congressional voting records, and diabetes diagnosis [11], East, Bauer, and Lanning surveyed pilot mental workload [12], college admissions officers have sought “to determine which variables are most important in judging the potential success of student” [13], dietary intake as it relates to urine samples and brain function of rats [14], and many other areas have been explored.

Related to imagery specifically, there has also been an abundance of work. In 1973, texture was identified as “one of the important characteristics used in identifying objects or regions of interest in an image” [15]. Within the realm of the electromagnetic spectrum (Figure 1), HSI images provide a ripe opportunity for multivariate analysis, where there are spectral values (features) for each spatial location or pixel [16].
Given that magnitude of possible applications and some of the difficulties associated research has abounded. Smetek worked at “hyperspectral target detection by developing autonomous anomaly detection and signature matching methodologies that reduce false alarms relative to existing benchmark detectors” [18]. Essentially, he “adapts multivariate outlier detection algorithms for use with hyperspectral datasets containing tens of thousands of non-homogeneous, high-dimensional spectral signatures” [18]. Johnson “[employed] independent component analysis (ICA) to unmix HSI images. Via new techniques to fully automate feature extraction, feature selection, and target pixel identification” [19]. This was extended “for global anomaly detection on a variety of HSI, utilizing fusion of spatial and spectral information, factor analysis, clustering, and screening” [16].
Feature Selection

For decades, feature selection has been a critical component of synthesizing, interacting with, and analyzing vast amounts of data. Truly, “variable and feature selection have become the focus of much research in areas of application for which datasets with tens or hundreds of thousands of variables are available” [20]. In fact, “the remarkable development of computing power and other technology has allowed scientists to collect data of unprecedented size and complexity” [21]. Moreover, while it is certainly applied to datasets with tens or hundreds of thousands or even millions of variables, feature selection has been applied to datasets of much lower dimensionality as well [16]. “Feature selection may be employed to improve a classification model . . . by eliminating non-informative features . . . [and it] may also be used to gain further insight into the rationale underlying class divisions within a particular domain” [14]. Simultaneously, in feature selection the desire is to minimize the computational time to the greatest degree possible, recognizing that in some cases reduced time is the preferred objective as long as adequate accuracy is maintained [12]. Feature selection enhances the probability of ascertaining insight into varying situations, while minimizing the Type I and Type II errors associated with the predicted results of those situations. It is important to remember that as selecting the appropriate features will enhance the predictive nature of the models examined, but it is crucial to ensure that we are making selections based on the appropriate response. We have to determine the right question or questions; what is the crucial result or response and why. And we certainly do not want the “right answer for the wrong question” [22].
Anomaly Detection

This leads us to field of anomaly detection. “Anomaly detection refers to the problem of finding patterns in data that do not conform to expected behavior . . . often referred to as anomalies, outliers, discordant observations, exceptions, aberrations, surprises, peculiarities, or contaminants” [23]. The “goal is to discover the true outliers and avoid mistakenly marking normal points as abnormal. In other words, a good anomaly detector must have a high detection rate and low false alarm rate” [24]. And even a low false alarm rate does not necessarily provide all the information of importance regarding the anomaly detected, leading to anomaly classification. Which is “implemented in a three-stage process, first by anomaly detection to find potential targets, followed by target discrimination to cluster the detected anomalies into separate target classes, and concluded by a classifier to achieve target classification” [5].

Classifying anomalies holds importance, because it is not just important to find the anomalies, “it is often more important to make sure that those anomalies that are reported to the user are in fact interesting” [25]. Notably, “experiments show that anomaly classification performs very differently from anomaly detection” [5]. Consistent with [26], [27], Chandola et al. reminds us that “anomaly detection is related to, but distinct from noise removal and noise accommodation . . . [where] noise can be defined as a phenomenon in data that is not of interest to the analyst, but acts as a hinderance to data analyis” [23].

Dimensionality

Numerous methodologies have been applied to increase the quality of feature selection. Importantly, feature selection is not limited to one field, but has been a chief
concern in numerous industries, fields, and problem sets [11], [13], [16]. In applying feature selection methodologies, one recognizes that often times the use of all available data to establish the model can “[result] in poor classification accuracy due to the distracting effect of numerous redundant and/or unnecessary variables” [12]. Additionally, we know that, from using ANNs, “as the number of features grows, the number of training vectors required grows exponentially”, obviously having potentially deleterious effects on computational time [11]. Screening to reduce these redundancies adds robustness and prediction accuracy to our solution space. However, one of the great things about feature selection is that it helps provide a framework to determine which variables should stay in the model. There are even cases where “noise reduction and consequently better class separation may be obtained by adding variables that are presumably redundant”, and this seems to be especially true, when utilizing methods more commonly associated with ranking instead of prediction [20].

In the case of HSI, an understanding of dimensionality is truly critical because there are specific hyperspectral bands that tend to absorb the incident energy associated with natural materials [16]. By identifying these atmospheric absorption bands, one reduces the dimensionality of an image, by removing those bands in which the sensor detects random noise [18].

**Screening Methods and Saliency Measures**

Since a primary component to the validity of feature selection within a scenario is this ability to maintain the features which discriminate between classes, screening methodologies are a crucial component in the success of feature selection. Various methods have been proposed utilizing saliency measures. While “a number of feature
saliency measures can be used to evaluate and rank the relative importance of candidate features . . . a saliency measure alone does not indicate how many of the candidate features should be used,” once again returning to the importance of asking the right question [28]. The Belue-Bauer screening method utilizes “a confidence interval constructed around [the average saliency of injected noise allowing] for the identification of features that contribute little to classification” [29]. Demonstrably, the utilization of “an appropriate hypothesis test to account for naturally paired observations of feature saliency measures and the use of a Bonferroni-type test statistic to reflect a conservative degree of statistical confidence for joint hypothesis testing. . . indicates that the truly noisy features can be consistently identified” [28]. Still other methods “measure the relative size of the weight vector emanating from each feature” [30]. Additional evidence indicates that while various screening methods “typically require between 10 and 30 training runs” the use of signal-to-noise (SNR) saliency measures may only require one training run potentially resulting in a significant reduction of computational processing time [11]. Other saliency measures have “[evaluated] each feature with respect to relative changes in either the neural network’s output or the neural network’s probability of error” [30].

**Techniques**

ANNs have long been a technique tied to feature selection, recognizing, as previously indicated that, “the relevance of a given input feature in a neural network is important in classification and prediction problems. “ANNs are desirable because they provide a well-structured framework to discover non-linear relationships within data sets that are considered ‘noisy’ or complex” [31]. In fact quite often, one is faced with a large
number of candidate features which may or may not be useful for the problem at hand” [28]. Figure 2 shows the framework of the neural network.

Essentially, ANNs “are networks or systems formed out of many highly interconnected nonlinear memoryless computing elements or subsystems . . . [that] can be represented mathematically as a weighted, directed graph” [32]. “There is also an appealing quality to [their] ‘brain-like’ structure” [29]. In fact it has been recognized that:

Computational properties of use to biological organisms or to the construction of computers can emerge as collective properties of systems having a large number
of simple equivalent components (or neurons). . . . The collective properties of this model produce a content addressable memory which correctly yields an entire memory from any subpart of sufficient size [33].

“Intuitively an analyst would like to include only those features that make a significant contribution to the network”, and given the networks structure as features are extracted and noise is removed and reduced the addressable memory can actually yield greater predictability [29]. Three processes take place to maximize the predictability of the ANN, first the network is trained with some sample of data, then it is internally validated with another, typically smaller sample, before it is finally tested for classification accuracy on the remainder of the data [11]. With autoencoding in particular, the goal is for the ANN to learn “the underlying feature structure of the data” that makes the data distinctive, then that ANN can be used for determining the classification of other data [34]. Figure 3 shows how the input layer and output layer are the same size, with the goal of finding the weights between nodes that yield the minimum reconstructive error between the input and the output.

![Autoencoder Structure](image)

**Figure 3: Autoencoder Structure** [35]

Similar to ANNs, discriminant analysis trains on one set of data to be predictive for another sample of data. “Discriminant analysis is a statistical technique for
classifying individuals or objects into mutually exclusive and exhaustive groups on the basis of a set of independent variables” [13]. By having multiple methods available, it enhances the robustness of the analysis. Notably, classification accuracy is not the only concern, “we will also be particularly concerned with determining the dimensions on which the groups differ . . . [allowing for] both prediction and explanation” [13].

“Principal component analysis is a standard tool in modern data analysis . . . [where] it provides a roadmap for how to reduce a complex data set to a lower dimension to reveal the sometimes hidden, simplified structures that often underlie it” [36]. “PCA generates a set of orthogonal vectors, any subset of which can be used to project into a subspace and where each vector accounts for some portion of the variance found in the data” [16]. This allows for ranking of features or components within the data based on the eigenvalues associated with each portion of the variance, allowing the user “to reduce the dimensionality of the feature space . . . by selecting only the . . . principal components with the largest eigenvalues” [11].

Notably, different methodologies are not necessarily exclusive of one another. In fact, while many use factor analysis to reduce data dimensionality [13], “factor analysis [may also act] as a fusion device for [various] feature selection procedures” [12]. Crucially, the problem must be identified, when “we wish to extract features which represent the difference between one pattern class and another. These features do not necessarily coincide with the important features to represent the pattern classes” [37]. “Selecting subsets of features that are useful to build a good predictor . . . contrasts with the problem of finding or ranking all potentially relevant variables” [20]. By coupling factor analysis or principal component analysis (PCA) with feature selection techniques,
demonstrated that the possibility exists to “extract features that are important for classification while maintaining features with good discriminatory power”\cite{11}.

**Multiple Outlier Detection**

A key difficulty when examining large datasets remains identifying multiple outliers simultaneously or in larger batches. Especially important is “establishing a metric that is not itself contaminated by inhomogeneities by which to measure how extraordinary a data point is” \cite{9}. “Despite considerable research . . . algorithms implemented for the detection of outliers are sparse. Moreover, the few algorithms available are so time-consuming that using them may be discouraging” \cite{38}. Specifically, “all multiple outlier detection methods have suffered in the past from a computational cost that escalated rapidly with the sample size” \cite{9}. BACON, the blocked adaptive computationally efficient nominators algorithm, has demonstrated success in reducing the computational expense while achieving quality results \cite{9}, \cite{38}, \cite{39}.

Utilizing BACON for the multivariate data representing a hyperspectral image, an “initial subset [is] selected based on Mahalanobis distances”, computing Mahalanobis distance with

\[
d_i(\bar{x}, S) = \sqrt{(x_i - \bar{x})^T S^{-1} (x_i - \bar{x})}, \quad i = 1, \ldots, n, \tag{1}
\]

“where \(\bar{x}\) and \(S\) are the mean and covariance matrix of the \(n\) observations” (Figure 4) \cite{9}.

Figure 4 shows us that there are definitely observations with high Mahalanobis distance, when we sort those observations, we attain a clearer picture of that distribution.
Figure 4: Plot of the Mahalanobis Distances (ARES1D) (from BACON algorithm)

Therefore, “the initial basic subset is given by the \( m \) observations with the smallest Mahalanobis distances for the whole sample” [38]. “This start is not robust, but it is affine equivariant . . . [and] subsequent iterations tend to make up for the non-robustness of the start as long as the fraction of outliers is relatively small”; however, a subsequent option allows for the determination of the initial subset utilizing “distances from the medians,” which provides a robust start, but without affine equivariance [9].

Which methodology to use largely depends on the problem at hand, extending from [40], [41], Caulk et al. [42] notes that “certain outlier detection methods, such as [multivariate trimming] are known to be unreliable due to their use of the Mahalanobis distance in determining the initial mean vector and covariance matrix estimate”. To complete BACON, a new basic subset is determined iteratively using a chi square distribution, and
the algorithm continues “until the size of the basic subset no longer changes, [whereby we] nominate the observations excluded by the final basic subset as outliers” [9].

The Reed-Xiaoli (RX) algorithm is another algorithm that has the potential to partition data, it “was developed from the generalized likelihood ratio (GLR) test . . . based on the suggestion that most optical image clutter can be modeled as a Gaussian random process with possibly a rapidly fluctuating space-varying mean and a more slowly varying covariance” [43]. As shown by [44], [45], RX “is considered as a benchmark” detector for comparison due to its simplicity” [46].

**Clustering**

“Clustering is usually considered to be the problem of partitioning a single set of unlabeled points . . . [where] one of the most common iterative, [non-hierarchical] algorithms is the K-means algorithm, broadly used for its simplicity of implementation and convergence speed” [40]. “The k-means process was originally devised in an attempt to find a feasible method of computing such an optimal partition” [47], where it “produces relatively high quality clusters considering the low level of computation required” [40]. K-means

“requires that the number of clusters used to classify the dataset will be pre-determined. It is based on determining arbitrary centers for the desired clusters, associating the samples with the clusters by using a pre-determined distance measurement, iteratively changing the center of the clusters and then re-associating the samples” [48].

Figure 5 shows an example derived in MATLAB, where one sees the clustering of various points around the centroid of those points.
Summary

While the applications are essentially boundless, we have defined sets of tools and processes which can be implemented and refined to optimize the solution space or predictive nature of a models despite what to the human mind may seem like insurmountable amounts of data. As we define the problem we wish to examine, the potential exists to gain far more insight into the space associated with problem using a combination of feature selection, feature extraction, and ranking.
III. Methodology

Chapter Overview

The purpose of this chapter is to provide a description of the methodology (Figure 6) we utilized to develop our analysis and garner our results. As previously noted in Section I, we employed a variety of techniques to both appropriately reduce the dimensionality of the data and assess the data for application, decision-making, and executable action. While all of these techniques have been used previously, they have never been used with the specific coordination suggested here. Our goal was to develop an actionable way to assess the data representing HSI images with a heightened degree of accuracy in the determination of anomalous data.

Figure 6: Flow Chart of Methodology
Test Subjects

In order to test and develop our methodology, we utilized a set of test images maintained by the Air Force Institute of Technology’s (AFIT’s) Sensor Fusion Laboratory derived from the HYDICE sensor. This provided us the opportunity to test our methodology against truth data already available for the images tested. Our testing was initially conducted against the image ARES1F, prior to expanding to ARES1D and ARES2D.

Summary

As stated previously, we utilized various methods in combination and coordination to assess and analyze potential anomalies within HSI images. We will discuss the method in association with the numbered steps indicated in the upper left hand corners of the Flow Chart of Methodology (Figure 6).

Step 1 – Multivariate Data

We select the image for testing. In this case, HYDICE sensor images as discussed above beginning with image ARES1F prior to assessing both ARES1D and ARES2D.

Pre-Processing

Step 2 – Remove Absorption Bands

As discussed in Section II, we acknowledge that HSI images contain absorption bands that generate noise within the images; therefore, we reduced dimensionality using the pre-determined absorption bands. This reduces the dimensionality of the image from 210 dimensions (Hyperspectral Bands) to 145 [50].
**Step 3 – Reshape Data**

Then, we converted image cubes into a two-dimensional image (Figure 7), which yields a two dimensional matrix as exemplified in the left-most image of Figure 8 [46].

![Figure 7: Conversion of Image Cube to a 2-dimensional matrix](image)

**Step 4 – BACON algorithm**

Upon re-shaping, we ran the resulting matrix through the BACON algorithm to pre-identify potential outliers. Our initial parameter settings utilized an alpha value of 0.05 for the Chi-Squared test with 40 degrees of freedom. We will discuss additional testing parameters further in Section IV. This provided a subset of potential outliers by splitting the original data set into two classes as discussed in Section I (Figure 8). Additionally, it yielded a covariance matrix and mean vector for the data that was not pre-identified.
Step 5 – PCA

We then took the PCA of the covariance matrix for the data identified by the BACON algorithm as clean data. This allowed the original data, centered to the mean vector associated with BACON, to be projected by the PCA results to give scores for analysis.

Step 6 – Dimensionality Reduction

At this point, we implemented Maximum Distance Secant Line (MDSL) (Figure 9) to reduce dimensionality based on the eigenvalues resulting from the PCA [10]. Between BACON and the reduced dimensionality derived from PCA (Figure 8), we had a workable dataset with which to progress. This was influential because it focused purely on those hyperspectral bands, which with assessment show the greatest importance. And, this had substantial effects on computational efficiency going forward. Especially as we
began the process of autoencoding, computationally it was enormously critical to have the lowest dimensionality possible.

**Figure 9: MDSL for ARES1F, with BACON based on medians**

**Step 7 – Partition Data Classes**

At this point, we separated the data into the two classes identified by BACON, with the first class encompassing background data and the second class encompassing potentially anomalous data.

**Post-Processing**

**Step 8a (Cluster Non-Outliers) and 8b**

With the dimensionality reduced data, the background class went through a k-means clustering algorithm. In order to eventually analyze the establishment of an optimal cluster quantity, we processed the data for a variety of cluster numbers, 2 to 11. Desiring to ensure and observe separation with the clusters, we plotted different cluster
combinations by their principal components. Figure 10 shows the 2-dimension component comparisons for ARES1D, when it is broken into 3 clusters. As one observes, there is clear separation of clusters across various principal components. However, as shown in green, it is difficult in 2-dimensions to see the extent of the separation for the outlier class. Some separation obviously exists, but the 2-dimension graph does not provide a complete picture, as they consistently display overlap with the clusters.

![Figure 10: 2-Dimension Principal Component Comparisons by Cluster with Outliers](image)

Recognizing no significant separation from the clustered data when compared to the BACON-determined anomalous data class, we then plotted the principal components across three dimensions (Figure 11). When plotted in this fashion, there is noticeable between the various clusters of the background class, and further separation between those clusters and the anomalous class.
Step 9 – Random Sample Clusters

At this point, we utilized random number generation to select one-tenth of the data to represent a cluster. This ended up giving us multiple advantages going forward. Our initial reasoning was that it allowed us to reduce computational time and increase computational efficiency in training the ANNs. But simultaneously, it provided us with a training data set and validation data set for when the data was utilized by the ANNs. Additionally, it allowed for another comparison between the various clusters, without running the entire cluster through another cluster’s net.

Step 10 – Autoencode ANNs for Each Cluster

At this point, we were able to autoencode ANNs to determine the best net for mirroring a specific cluster. As previously hinted, we applied the ANN to a training set of the data using the randomly generated sample set, then we validated the ANN performance with another randomly generated validation set of equal size. For training the ANN, we used
10 hidden nodes and we trained the ANN as many as 20 times. Referencing the diagram from the flow chart (Figure 6), the autoencoder attempts to generate weights which will reconfigure the input layer, \( x_0 \ldots n \), to the output layer, \( z_0 \ldots n \), where \( n \) is the number of dimensions remaining after the dimensionality reduction, Step 6 (Figure 12).

**Figure 12: Autoencoder Structure (Reproduced from Figure 3) [35]**

**Step 11 – Determine the Best Net**

In this step, we determine the best network to use, by choosing the ANN which produces the lowest mean squared error value for the associated cluster.

**Step 12 – Run Potential Outliers through the Best Nets**

Knowing the best ANNs for each cluster, we ran the data that was class pre-identified by BACON as having potential outliers through each cluster’s optimal net.

**Performance Assessment**

**Step 13 – Determine Reconstructive Error for Cluster and Potential Outliers**

Next we calculated the reconstructive errors for each observation in regards to ANN performance within each cluster.

**Step 14 – Develop Receiver Operating Characteristic (ROC) Curves**

With the truth data and the maximum reconstructive errors for each cluster, we generated a ROC curve of the background data versus the anomalies. After completing
this for various images, we were able to determine a threshold which we were comfortable and classify the observations within the originally designated anomalous class as true anomalies or as background.
IV. Analysis and Results

Chapter Overview

We specifically examined three hyperspectral imagery (HSI) images in an effort to validate the algorithmic and methodological results as compared to the known anomalies within the specified images. Since the Autoencoded Reduced Clustering for Anomaly Detection Enrichment (ARCADE) parameters are modifiable, we could examine the each of the images using our base methodology independently. In conducting the analysis, we developed receiver operating characteristic (ROC) curves to measure our method’s anomaly prediction accuracy. Furthermore, we examine various parameter settings throughout the process to gain insight into how those parameters affect the overall methodology.

In addition to comparing the results between the images themselves, we also compared the results of the methodological variations for each image. This provided some insight into inelasticity of various parameters and the robustness of the ARCADE method. While our base methodology (Figure 6) was described in Section III, this section will examine excursions made from the basic process (Table 1).

Table 1: Image & Parameter Excursions (Base Methodology)

<table>
<thead>
<tr>
<th>Run</th>
<th>Image</th>
<th>Degrees of Freedom</th>
<th>ANNs #</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ARES1F</td>
<td>40</td>
<td>3</td>
</tr>
<tr>
<td>2</td>
<td>ARES1F</td>
<td>30</td>
<td>3</td>
</tr>
<tr>
<td>3</td>
<td>ARES1F</td>
<td>20</td>
<td>3</td>
</tr>
<tr>
<td>4</td>
<td>ARES1D</td>
<td>40</td>
<td>20</td>
</tr>
<tr>
<td>5</td>
<td>ARES2D</td>
<td>40</td>
<td>20</td>
</tr>
<tr>
<td>6</td>
<td>ARES1F</td>
<td>40</td>
<td>20</td>
</tr>
<tr>
<td>7</td>
<td>ARES1F</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>8</td>
<td>ARES1D</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>9</td>
<td>ARES2D</td>
<td>20</td>
<td>20</td>
</tr>
</tbody>
</table>
We initially examined the basic methodology (Figure 6), considering various parameter changes, while recognizing the possibility of using different pre-processing techniques. Table 1 shows the parameter changes that were explored in examining our basic methodology. Following a description of the results associated with the base methodology and the parameter excursions associated with the base methodology, we will describe five variations that were made regarding our pre-processing techniques. These variations and excursions provide substantial information to inform the quality of our results and conclusions. In our initial excursions from our base methodology, we examined the number of degrees of freedom associated with the BACON algorithm and we examined the number of ANNs. For degrees of freedom, we ranged from 20 to 40, and for ANNs, we ranged from 3 to 20 to determine if additional processing would impact the results.

Results of Base Methodology
Step 1 to 3 – Image Processing

After importing the images, we removed the absorption bands, and re-dimensioned the data into two dimensions as previously described.

Step 4 – BACON algorithm

As discussed in Sections II and III, BACON finds a pre-identified selection of potential outlier points, which we define as our anomalous class, while considering all other points as our background class. Figure 14 shows a visualization of the image on the left and a class separated visual of the image on the right.

![Figure 14: True Image versus BACON Anomalies (Run 1)](image)

By visually examining this pictorial comparison, it provided a level of confirmation that BACON was indeed picking up at least a majority of the truly anomalous data, promoting continuation of our method. Simultaneously, we were able to compare the difference with changes to the degrees of freedom, where we see that as degrees of freedom decreases the size of the anomalous class increases significantly (Figure 15).
Similarly, an equivalent representation of ARES1D (Figure 16) and ARES2D (Figure 17), yielded similarly positive confirmations. Furthermore, a direct comparison of the class sizes provided insight into the degree of separation between the classes (Table 2), similar to that of Figure 15. As one observes, increasing the degrees of freedom pulls substantially more pixels into the background class, but independent of degrees of freedom, pre-processing the data using BACON allows for quality separation of the data into the two meta-classes. The concern becomes whether the reduced degrees of freedom over-classifies the data into the anomalous class.

<table>
<thead>
<tr>
<th>Table 2: Class sizes varying BACON’s degrees of freedom</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>ARES1F</td>
</tr>
<tr>
<td>ARES1D</td>
</tr>
<tr>
<td>ARES2D</td>
</tr>
</tbody>
</table>
It is important to note that the lines through class separation images are most likely sensor artifacts, which if removed may enhance classification. This would be a great opportunity for future study. Either related to the detection of the sensor artifacts themselves or using some method to separate the anomalous class, similar to how we will utilize k-means clustering further into our methodology.

*Step 5 – PCA*

As shown in Figure 9, PCA reduces the dimensionality of ARES1F to 12 dimensions. Similarly, ARES1D’s (Run 4) dimensionality reduces to 10 dimensions and
ARES2D’s (Run 5) dimensionality reduces to nine dimensions (Figure 18). Thus, for these images, the worst case is a dimensionality reduction by a factor of 12.

![Figure 18: MDSL (Runs 4 & 5) [10]](image)

**Step 8a – Cluster Non-Outliers**

The data representing the background class was separated into clusters utilizing a k-means algorithm in MATLAB as referenced in Section II. We separated the data in anywhere from 2 to 11 clusters, allowing for optimality comparisons. Upon clustering the clean data (background class), one observes that in general the clusters tend to have relatively uniform size with a few assumptions (Table 3), which further supports our visual assessment of the components from Figure 10 and Figure 11. Additionally, we can observe differing levels of separation by implementing color-mapping (Figure 19). It is important to note that each color-map has utilized the dark blue to represent the anomalous class.
Step 10 and 11 – Autoencode and Select Best ANN

Given the various clusters, we autoencoded ANNs on the clusters using a 10 percent random sample of the data, then ran another random sample of equal size through the various ANNs, whereby we were able to determine the best ANN for each cluster by taking the ANN that produced the lowest mean squared error with the second sample (Table 4). As we can see, the best ANN does not necessarily occur within the first three (Run 1), but some reasonable performance occurred in the first three ANNs, so if computational efficiency was a major concern, it could be worthwhile for future research to attempt to optimize the number of ANNs for a full array of images.
Table 4: Reconstructive Error's for 2 Samples (Run 6)

<table>
<thead>
<tr>
<th></th>
<th>ANN 1</th>
<th>ANN 2</th>
<th>ANN 3</th>
<th>ANN 4</th>
<th>ANN 5</th>
<th>ANN 6</th>
<th>ANN 7</th>
<th>ANN 8</th>
<th>ANN 9</th>
<th>ANN 10</th>
<th>ANN 11</th>
<th>ANN 12</th>
<th>ANN 13</th>
<th>ANN 14</th>
<th>ANN 15</th>
<th>ANN 16</th>
<th>ANN 17</th>
<th>ANN 18</th>
<th>ANN 19</th>
<th>ANN 20</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1</td>
<td>753.2</td>
<td>703.8</td>
<td>734.7</td>
<td>1248.0</td>
<td>1298.5</td>
<td>720.6</td>
<td>718.5</td>
<td>715.4</td>
<td>788.4</td>
<td>730.5</td>
<td>1027.1</td>
<td>774.8</td>
<td>710.3</td>
<td>748.0</td>
<td>726.5</td>
<td>1308.2</td>
<td>726.3</td>
<td>744.8</td>
<td>724.1</td>
<td>725.9</td>
</tr>
<tr>
<td>Sample 2</td>
<td>941.0</td>
<td>946.8</td>
<td>874.3</td>
<td>1778.4</td>
<td>1586.4</td>
<td>883.1</td>
<td>802.5</td>
<td>716.5</td>
<td>879.7</td>
<td>913.2</td>
<td>2182[4]</td>
<td>4]</td>
<td>694.0</td>
<td>894.0</td>
<td>963.1</td>
<td>903.9</td>
<td>1522.1</td>
<td>952.0</td>
<td>990.2</td>
<td>895.3</td>
</tr>
</tbody>
</table>

*Step 12 and 13 – Assess outliers in light of reconstructive error*

After running the clusters through ANNs in order to autoencode each cluster, we were able to run the pre-identified background class through the ANN associated with each cluster; thereby yielding a reconstructive error for each BACON-identified outlier in regards to each cluster. As the histograms seem demonstrate and confirm (Figure 20), a reasonable proportion of the data within the anomalous class actually belongs to the background class, but it is difficult to distinguish the extent of the anomalous class that belongs to the background class. Additionally, it is difficult to visually distinguish the differences between the two histograms, so Figure 21 compares the classes more directly.

![Figure 20: Histograms of Reconstructive Error (Run 6)](image)

Based on this insight, we determined the maximum reconstructive error of the background class, then we projected all the observations in the anomalous class that had a greater reconstructive into another histogram (Figure 21) to attain greater insight into how the anomalous class was interacting with the background class in our post-processing procedures.
This separation was further represented when we placed all the observations on a scatter plot (Figure 22). We see that overall the background class fits well within the area encompassed by the anomalous class, further indicating that at some threshold we can predict or classify the anomalous class as true outliers.
Step 14 – ROC Curves

ROC curves of the respective reconstructive errors show an improved FPF statistics at a greater range of thresholds. Importantly for ARES1F at least running more ANNs 20 versus 3 for Runs 6 and 1 respectively (Figure 23), the method provides greater improvement across the thresholds with greater consistency.

![ROC Curves](image)

**Figure 23: ROC Curves (Runs 6 & 1)**

This provided the initial indication that ARCADE could have potential, in that while it does not seem like a major jump looking at the figure, a nearly three percent increase in predictive capacity is statistically significant.

Furthermore, we compared the results of the BACON algorithm with the various degrees of freedom. As we see with only three ANNs, the lower the predictive capacity of the pre-processor the more impact the post-processor has on improving prediction (Figure 24).
When comparing the results between images, we see that with 20 degrees of freedom too much data is included in the anomalous class for BACON to be effective independent of the image. This is further confirmed with 20 ANNs (Figure 25). In all of these cases, ARCADE shows the potential for improvement upon BACON’s pre-processing, but the final result is definitely affected by the initial class sizes. Figure 26 shows the results for ARES2D, where once again the degrees of freedom make as significant difference in the pre-processing quality.
Variation Summary

With the parameter excursions and results, it enabled the idea of varying the pre-processing techniques themselves to attain further information regarding the effectiveness of the post-processor. As we progressed through the variations, it enabled even more ideas regarding implementation of pre-processing, which yielded some very interesting results. While some parameter changes were made through the following variations, the primary goal was to compare different methodologies. Further research regarding parameters and even more pre-processing methods is definitely a possibility. The five variations we explored included variations of BACON and RX algorithms (Table 5).

Table 5: Variation from Base Methodology

<table>
<thead>
<tr>
<th>Variation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Base Methodology</td>
</tr>
<tr>
<td>1</td>
<td>Switched Steps 4 &amp; 5 to perform PCA (5) prior to BACON (4)</td>
</tr>
<tr>
<td>2</td>
<td>Switched Steps 4, 5 &amp; 6 to perform PCA (5), reduce dimensionality (6), then execute BACON (4)</td>
</tr>
<tr>
<td>3</td>
<td>RX replaced BACON</td>
</tr>
<tr>
<td>4</td>
<td>Threshold Variations on raw RX output</td>
</tr>
<tr>
<td>5</td>
<td>Threshold Variations on Mahalanobis Distance</td>
</tr>
</tbody>
</table>
Methodology Variation 1

Our first change in methodology was to implement the BACON classification after performing PCA. Essentially, the variation instituted the BACON algorithm against the scores rather than the raw data to determine classes. In this variation, the execution of the dimensionality reduction still occurred after BACON was implemented. This is impactful because it means that BACON is applied to the data centered and projected onto an orthogonal space. Table 6 shows the excursions made within this variation. This yielded a reduced dimensionalities of 15 dimensions (Figure 28), with MDSL [10]. Interestingly, this variation produced slightly higher dimensionality for ARES1F (Figure 28) which is likewise for ARES2D, but the dimensionality is slightly reduced for ARES1D (Figure 28).

![Diagram Showing Variation 1](Refer to Figure 6 or Figure 13)

Table 6: Image & Parameter Excursions (Using Scores Matrix)

<table>
<thead>
<tr>
<th>Run</th>
<th>Image</th>
<th>Degrees of Freedom</th>
<th>ANNs #</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ARES1F</td>
<td>40</td>
<td>3</td>
</tr>
<tr>
<td>2</td>
<td>ARES1D</td>
<td>40</td>
<td>3</td>
</tr>
<tr>
<td>3</td>
<td>ARES2D</td>
<td>40</td>
<td>3</td>
</tr>
<tr>
<td>4</td>
<td>ARES1F</td>
<td>40</td>
<td>20</td>
</tr>
<tr>
<td>5</td>
<td>ARES1D</td>
<td>40</td>
<td>20</td>
</tr>
<tr>
<td>6</td>
<td>ARES2D</td>
<td>40</td>
<td>20</td>
</tr>
</tbody>
</table>
Clustering was definitely affected by the variation, with cluster sizes shifting from one cluster to another, despite maintaining the same class sizes for ARES1F (Table 7). This truly demonstrates the impact of having centralized data and its effects on BACON.

Table 7: Cluster Sizes (Run 6)

<table>
<thead>
<tr>
<th>Cluster</th>
<th>2 Clusters</th>
<th>3 Clusters</th>
<th>4 Clusters</th>
<th>5 Clusters</th>
<th>6 Clusters</th>
<th>7 Clusters</th>
<th>8 Clusters</th>
<th>9 Clusters</th>
<th>10 Clusters</th>
<th>11 Clusters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cluster 1</td>
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<td>5062</td>
<td>6204</td>
<td>5624</td>
<td>4878</td>
<td>3578</td>
<td>2836</td>
<td>1661</td>
<td>1968</td>
<td>3058</td>
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<tr>
<td>Cluster 2</td>
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<td>9195</td>
<td>6527</td>
<td>5899</td>
<td>4057</td>
<td>3768</td>
<td>1530</td>
<td>2792</td>
<td>1802</td>
<td>3555</td>
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<tr>
<td>Cluster 3</td>
<td>10052</td>
<td>3018</td>
<td>4863</td>
<td>4571</td>
<td>1695</td>
<td>4032</td>
<td>2466</td>
<td>1081</td>
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<tr>
<td>Cluster 4</td>
<td>8600</td>
<td>5618</td>
<td>4591</td>
<td>2790</td>
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<td>2308</td>
<td>3107</td>
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<tr>
<td>Cluster 5</td>
<td>2845</td>
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<td>3686</td>
<td>3402</td>
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<tr>
<td>Cluster 6</td>
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<td>2870</td>
<td>3545</td>
<td>2824</td>
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<td></td>
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<tr>
<td>Cluster 7</td>
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<td>2830</td>
<td>3043</td>
<td>2677</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Cluster 8</td>
<td>3324</td>
<td>3217</td>
<td>3017</td>
<td>1972</td>
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<tr>
<td>Cluster 9</td>
<td>2251</td>
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<tr>
<td>Cluster 11</td>
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<td></td>
<td></td>
<td></td>
<td>1646</td>
</tr>
</tbody>
</table>
The ANN post-processing improved post-processing classification accuracy slightly with the use of 20 ANNs by 0.0024 percent despite the fact that no change occurred in the pre-processing classification for ARES1F (Figure 29), and this was further confirmed and enhanced with the use of only three ANNs at 0.0055 percent improvement.

**Methodology Variation 2**

In this variation, we once again assess the data utilized PCA followed by BACON, but in this particular variation, we performed our dimensionality reduction prior to the execution of the BACON algorithm. Therefore once again, BACON was run on the PCA scores, but in this case, it was intended to only run BACON against the dimensions identified through MDSL [10].
However, upon attempting this variation with the parameters associated with BACON for the previous runs, we were not able to execute the subsequent clustering because the class size of the anomalous class was too small or in some cases non-existent. Thus, we substantially adjusted three of the parameters. Notably, some adjustments led to all data observations being sent to the anomalous class, while leaving the background class empty. First, we made adjustments to the alpha values associated with the Chi-Squared test utilized in BACON. Previous runs had applied an alpha of 0.05, this was modified to span between 0.1 and 0.5. Second, we adjusted the degrees of freedom. Other experimentation had degrees of freedom ranging from 20 to 40; with this variation, we ranged degrees of freedom from four to ten. Third, the dimensionality given using MDSL did not provide enough dimensionality to successfully execute BACON. Initially, we contemplated applying either a Horn’s Curve or Kaiser’s Criterion using the correlation matrix of the data to determine dimensionality (Figure 31), but unfortunately both of these methods reduced the dimensionality of the data even more than MDSL; therefore, BACON was not left with enough data to clearly extricate one class from the other.

Figure 31: Horn's Curve (ARES1F)
Learning this, we experimented with an added dimensionality ranging from an additional 20 to an additional 50 dimensions. Again it is critical to note that in some cases, these parameter varieties did not include all of the known anomalies and in others they sent so much of the image data into the anomalous class that it did not have a positive effect. Based on these findings, we tried a variety of parameter combinations to see what the differences would be as the parameters shifted. Generally speaking, we saw some improved performance using our post-processing methodology, but we did not see across the board quality results with any specific set of parameter settings across the various images.

For ARES1F, we compared four different parameter settings (Table 8), but while our method showed demonstrable improvement over the BACON class identification accuracy, BACON’s predictive capacity for this image given the reduced dimensionality was extremely poor. Therefore even with our improved performance, the performance regarding this image was poor. One sees that BACON never achieves a classification rate much better than the flip of a coin (Figure 32). The best result of our method comes when the parameter settings are at their middle points, but the second best performance comes when the parameter settings were at their lowest points, leaving a lot to be discovered.
Table 8: Image & Parameter Excursions (BACON with Reduced Dimensionality)

<table>
<thead>
<tr>
<th>Run</th>
<th>Image</th>
<th>Alpha</th>
<th>Degrees of Freedom</th>
<th>Added Dimensionality</th>
<th>ANNs #</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ARES1F</td>
<td>0.1</td>
<td>4</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>2</td>
<td>ARES1F</td>
<td>0.1</td>
<td>4</td>
<td>50</td>
<td>20</td>
</tr>
<tr>
<td>3</td>
<td>ARES1F</td>
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<td>7</td>
<td>35</td>
<td>20</td>
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<td>4</td>
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<td>10</td>
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<td>20</td>
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<tr>
<td>5</td>
<td>ARES1D</td>
<td>0.1</td>
<td>4</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>6</td>
<td>ARES1D</td>
<td>0.1</td>
<td>4</td>
<td>50</td>
<td>20</td>
</tr>
<tr>
<td>7</td>
<td>ARES1D</td>
<td>0.3</td>
<td>7</td>
<td>35</td>
<td>20</td>
</tr>
<tr>
<td>8</td>
<td>ARES1D</td>
<td>0.5</td>
<td>10</td>
<td>50</td>
<td>20</td>
</tr>
</tbody>
</table>

Using ARES1D, we re-attempted the runs for each parameter setting to see if it yielded any important findings (Figure 33). Interestingly, while BACON performed better with the minimum settings, the higher parameters yielded a substantially better result utilizing ARCADE, so this may warrant more experimentation in the future.
Methodology Variation 3

Our third methodological variation was a significant shift in completely changing our pre-processor to utilize RX rather than BACON (Figure 34). This approach also maintained the same dimensionality adjustments as Variation 1 due to conducting PCA with a covariance matrix of the data, rather than some modified covariance matrix associated with the pre-processor of choice. This of course lends to the possible future study of extracting a covariance matrix from the background class identified by the RX algorithm and proceeding with PCA, dimensionality reduction, and post-processing from there. For RX, we ran RX then ran ARCADE through every image as shown in Table 9.
Fascinatingly, RX substantially shrinks the size of the anomalous class as compared to the BACON derived methods previously examined, leaving only 2,264 observations in the anomalous class for ARES1F, showing the impact of varying pre-processors. The truth mapping (Figure 35) regarding the RX identified anomalous class produced a far lower level of inclusivity for ARES1F background class as compared to the BACON methods (Figure 36), where not all of the known anomalous pixels were included by the anomalous class using the RX algorithm. Pre-processing utilizing RX incorporated 783 of the ARES1F images into the background class, which means that only 77.8 percent of the truly anomalous pixels were incorporated in the anomalous class for testing.
To further testing, we still desired to determine the effects of our post-processing despite the lack of all anomalous pixels being incorporated. Besides the corresponding change in cluster size (Table 10), there also exists a major change in the variance of the cluster sizes when compared to the BACON methods described earlier (Table 11).
We see that while there were slight changes in the variances and standard deviations between the Base Method and Variation 1, as much as a seven percent increase in standard deviation, RX promotes a significant increase in the variance and standard deviation independent of the number of clusters. It never falls below a 16 percent increase in standard deviation, while reaching as high as a 56 percent increase (Table 11).

### Table 10: Cluster Size (ARES1F)

<table>
<thead>
<tr>
<th>Clusters</th>
<th>2 Clusters</th>
<th>3 Clusters</th>
<th>4 Clusters</th>
<th>5 Clusters</th>
<th>6 Clusters</th>
<th>7 Clusters</th>
<th>8 Clusters</th>
<th>9 Clusters</th>
<th>10 Clusters</th>
<th>11 Clusters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cluster 1</td>
<td>11306</td>
<td>15447</td>
<td>2427</td>
<td>3407</td>
<td>4228</td>
<td>5285</td>
<td>6220</td>
<td>7138</td>
<td>8117</td>
<td>9177</td>
</tr>
<tr>
<td>Cluster 2</td>
<td>16990</td>
<td>2596</td>
<td>9096</td>
<td>5692</td>
<td>1523</td>
<td>222</td>
<td>4809</td>
<td>1155</td>
<td>4500</td>
<td>1167</td>
</tr>
<tr>
<td>Cluster 3</td>
<td>10259</td>
<td>11112</td>
<td>2373</td>
<td>1256</td>
<td>6799</td>
<td>2709</td>
<td>4089</td>
<td>4532</td>
<td>3525</td>
<td>3352</td>
</tr>
<tr>
<td>Cluster 4</td>
<td>5661</td>
<td>3588</td>
<td>6372</td>
<td>1075</td>
<td>1184</td>
<td>1076</td>
<td>1709</td>
<td>3026</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cluster 5</td>
<td>7236</td>
<td>8059</td>
<td>6397</td>
<td>4561</td>
<td>3927</td>
<td>2937</td>
<td>3503</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cluster 6</td>
<td></td>
<td>6918</td>
<td>5170</td>
<td>5752</td>
<td>8028</td>
<td>3349</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>Cluster 7</td>
<td></td>
<td>2201</td>
<td>4714</td>
<td>3817</td>
<td>4707</td>
<td>4516</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cluster 8</td>
<td></td>
<td></td>
<td>3422</td>
<td>4857</td>
<td>2201</td>
<td>3038</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cluster 9</td>
<td></td>
<td></td>
<td></td>
<td>4529</td>
<td>3110</td>
<td>1493</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Cluster 10</td>
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<td></td>
<td></td>
<td></td>
<td>1081</td>
<td>217</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cluster 11</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>4047</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

After running the ANNs on the background class clusters, we see minimal improvement in the FPF along the ROC curve when it comes to ARES1F, still showing a substantially lower prediction accuracy over the course of the curve than the BACON derived methods (Figure 37). Subsequently, RX showed improvement over BACON when it came to ARES1D and ARES2D, while our post-processing demonstrated a continued improvement in FPF for ARES1D, with no statistically significant improvement demonstrated for ARES2D (Figure 37). It is important to note that we did
not change any of the parameter settings built into the RX algorithm. But the use of RX led us to variation 4, which was to assess the peak point given the actual data from RX.

Methodology Variation 4

Deriving from Variation 3, we determined that it would be worth examining the actual values associated with the RX algorithm. In order to do this, we ran the RX algorithm as we did in Variation 3, but we set a variety of thresholds by which to test the images. First, we determined that 13.11 percent was a peak threshold for ARES2D. We utilized this image due to its high amount of noise. Then we ran the pre-processor for all the images, dividing the background class and the anomalous class using this threshold. So in effect, 86.89 percent of the data went into the background class, while 13.11

Figure 37: ROC Curves (Runs 1, 2 & 3)
percent was called anomalous. We compared this with thresholds of 0.311 and 0.05 (Table 12). Of note, all experimentation presented henceforth utilized 20 ANNs.

Table 12: RX Image & Threshold Excursions

<table>
<thead>
<tr>
<th>Run</th>
<th>Image</th>
<th>Threshold</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ARES1F</td>
<td>0.311</td>
</tr>
<tr>
<td>2</td>
<td>ARES1F</td>
<td>0.1011</td>
</tr>
<tr>
<td>3</td>
<td>ARES1F</td>
<td>0.05</td>
</tr>
<tr>
<td>4</td>
<td>ARES1D</td>
<td>0.311</td>
</tr>
<tr>
<td>5</td>
<td>ARES1D</td>
<td>0.1311</td>
</tr>
<tr>
<td>6</td>
<td>ARES1D</td>
<td>0.05</td>
</tr>
<tr>
<td>7</td>
<td>ARES2D</td>
<td>0.311</td>
</tr>
<tr>
<td>8</td>
<td>ARES2D</td>
<td>0.1311</td>
</tr>
<tr>
<td>9</td>
<td>ARES2D</td>
<td>0.05</td>
</tr>
</tbody>
</table>

This allowed us to pursue our post-processing as we had through the other variations. Assessing the HSI images using this methodology truly forces the user to reflect on and determine their goal for analysis. Similarly to its seeming lack of significant effect on the RX classification, ARCADE does not seem to show substantial improvement when considering various thresholds on the RX outputs. Importantly, lower thresholds demonstrated less improvement in post-processing. However, by examining various thresholds within RX, we see a definite shift between TPF and FPF for each of the images. RX-distance represents the class separation threshold when we use the LRX values themselves. RX-algorithm represents the classification based on RX’s nominal settings. For ARES1F, we can see how a higher threshold gives us a better TPF at its peak, but a trade-off definitely occurs (Figure 38). The RX classification generally does very well in limiting FPF for this image, but we see that depending on threshold, it is capable of producing a high TPF rate, which in some cases may be more beneficial.
Figure 38: ROC Curves (Runs 1, 2 & 3)

This holds true for ARES1D (Figure 39) and ARES2D (Figure 40), but ARES1D shows a pronounced positive return on investment (ROI) using the middle threshold, whereas with ARES1F we saw a more distinct trade-off. The trade-off accelerates at a threshold of 0.05, and we see a slightly reduced overall performance, but the FPF is reduced significantly.
Figure 39: ROC Curves (Runs 4, 5 & 6)

While ARES 2D does not yield a substantial positive ROI using a lower threshold, a high threshold conversely produces a substantial negative ROI (Figure 40). We see that at the 0.05 threshold a small tradeoff occurs, whereas at the 0.1311 threshold a tradeoff occurs in the opposite direction, but the 0.311 threshold significantly decreases classification performance.
Our fifth variation in methodology changes the pre-processing requirements of BACON to use a more optimal Mahalanobis distance value, similar to how Variation 4 was conducted, but utilizing BACON instead of RX (Figure 41).

Figure 40: ROC Curves (Runs 7, 8 & 9)

Methodology Variation 5

Figure 41: Diagram showing Variation 5 (Refer to Figure 6 or Figure 13)
The whole concept behind this was whether the parameters of the pre-processors actually provided a near-optimal solution space. It helped to approach the idea from a perspective of simplification. We determined the peak point by finding the Mahalanobis distance as calculated within the BACON algorithm, plotting a ROC curve and finding the point along the ROC curve nearest to perfection. This was initially conducted on ARES1D (Figure 42), which yielded a threshold of 0.1632 at an alpha of 0.05, which we then transitioned to 0.32 and finally to 0.05 for the threshold. Table 13 gives a comprehensive breakdown of the excursions taken.

Table 13: Image & Parameter Excursions (Mahalanobis Distance)

<table>
<thead>
<tr>
<th>Run</th>
<th>Image</th>
<th>Threshold</th>
<th>Alpha</th>
<th>Sample Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ARES1D</td>
<td>0.1632</td>
<td>0.05</td>
<td>10%</td>
</tr>
<tr>
<td>2</td>
<td>ARES1F</td>
<td>0.1632</td>
<td>0.05</td>
<td>10%</td>
</tr>
<tr>
<td>3</td>
<td>ARES2D</td>
<td>0.1632</td>
<td>0.05</td>
<td>10%</td>
</tr>
<tr>
<td>4</td>
<td>ARES2F</td>
<td>0.32</td>
<td>0.05</td>
<td>10%</td>
</tr>
<tr>
<td>5</td>
<td>ARES3D</td>
<td>0.32</td>
<td>0.05</td>
<td>10%</td>
</tr>
<tr>
<td>6</td>
<td>ARES3F</td>
<td>0.32</td>
<td>0.25</td>
<td>10%</td>
</tr>
<tr>
<td>7</td>
<td>ARES2D</td>
<td>0.32</td>
<td>0.25</td>
<td>10%</td>
</tr>
<tr>
<td>8</td>
<td>ARES3D</td>
<td>0.32</td>
<td>0.25</td>
<td>10%</td>
</tr>
<tr>
<td>9</td>
<td>ARES3F</td>
<td>0.32</td>
<td>0.25</td>
<td>10%</td>
</tr>
<tr>
<td>10</td>
<td>ARES2D</td>
<td>0.32</td>
<td>0.5</td>
<td>10%</td>
</tr>
<tr>
<td>11</td>
<td>ARES3D</td>
<td>0.32</td>
<td>0.5</td>
<td>10%</td>
</tr>
<tr>
<td>12</td>
<td>ARES2D</td>
<td>0.32</td>
<td>0.5</td>
<td>10%</td>
</tr>
<tr>
<td>13</td>
<td>ARES3F</td>
<td>0.05</td>
<td>0.25</td>
<td>10%</td>
</tr>
<tr>
<td>14</td>
<td>ARES2D</td>
<td>0.05</td>
<td>0.25</td>
<td>10%</td>
</tr>
<tr>
<td>15</td>
<td>ARES3D</td>
<td>0.05</td>
<td>0.25</td>
<td>10%</td>
</tr>
<tr>
<td>16</td>
<td>ARES3F</td>
<td>0.05</td>
<td>0.25</td>
<td>40%</td>
</tr>
<tr>
<td>17</td>
<td>ARES2D</td>
<td>0.05</td>
<td>0.25</td>
<td>40%</td>
</tr>
<tr>
<td>18</td>
<td>ARES3F</td>
<td>0.05</td>
<td>0.05</td>
<td>10%</td>
</tr>
<tr>
<td>19</td>
<td>ARES2D</td>
<td>0.05</td>
<td>0.05</td>
<td>10%</td>
</tr>
</tbody>
</table>

Figure 42: ROC Curve to Establish Initial Threshold (Run 1)
Applying this 0.1632 threshold to ARES1F demonstrated an significant improvement over the BACON’s classification utilizing the same parameters, displaying an increase of 3.6 percent in the AUC (Figure 43). Unfortunately, this was not replicated wholesale with ARES2D, as the FPF increases slightly at its peak following post-processing. However, the change is minimal (Figure 43).

Experimenting with a larger threshold indicated that alpha increased the post-processor showed improved classification, as compared to the standard BACON classification, but it further demonstrated the importance in correctly setting the threshold for the Mahalanobis distance. At a threshold of 0.32 and alpha value of 0.05, BACON far outperforms the threshold with post-processing for every image (Figure 44). As we increased the alpha value to 0.25 (Figure 45), and subsequently to 0.5 (Figure 46), the post-processor appropriately identified a higher proportion of the false positive to reduce the FPF, but this was due in part to the fact that BACON increased the size of the background class, generating more false positives. In conjunction with the previous research displayed, this provided some hope in the post-processing method regarding its ability to correctly identify false positives classified the pre-processor of preference.
Figure 44: ROC Curves (Runs 4, 5 & 6)
Figure 45: ROC Curves (Runs 7, 8 & 9)
Figure 46: ROC Curves (Runs 10, 11 & 12)
Conversely, reducing the threshold to 0.05 with an alpha value of 0.25 had the effect of substantially under-identifying the anomalous observations in two of the three images (Figure 47), indicating the need for further experimentation with between the thresholds of 0.1632 and 0.05 and amongst the various potential alpha values. This reduced threshold did show some promise for future research as it substantially reduced the FPF for ARES2D, while substantially increasing the AUC (Figure 47).

As a final trial, we desired to see if the sample size for autoencoding the clusters would have a significant effect on the post-processing. We re-ran the ROC curves with the 0.05 threshold on the Mahalanobis distance and the 0.25 alpha value for ARES1F and

![ROC Curve (Runs 13, 14 & 15)](image)

**Figure 47: ROC Curve (Runs 13, 14 & 15)**

As a final trial, we desired to see if the sample size for autoencoding the clusters would have a significant effect on the post-processing. We re-ran the ROC curves with the 0.05 threshold on the Mahalanobis distance and the 0.25 alpha value for ARES1F and
ARES1D utilizing a 40 percent sample from the clusters. While there was very minor improvement, neither example showed an improvement that would justify the substantial increase in computational time. However, this was at a threshold that had limited success as a whole, so it would be justifiable to conduct more testing to see if this change holds any promise. Finally, we maintained the 0.05 threshold, but changed the alpha value back to 0.05, showing that once again the lower alpha value enhances the performance quality across the board (Figure 48).

Figure 48: ROC Curves (Runs 16, 17, 18 & 19)
Investigative Questions

We examined six different variations of our method (Figure 13). In these variations, we compared two different pre-processors, two orders of processing and two outputs from each of the pre-processors. In addition to the comparisons between the pre-processors, we compared various parameter settings associated with those processors. Furthermore in post-processing, we compared a variety of thresholds by which to comparatively assess HYDICE images. This provided us with extensive scope into the overall methodology to optimally determine parameter settings and methodology by which to approach the analysis of HYDICE HSI images for classification and anomaly detection.

All of this experimentation allowed us to explore the effectiveness of pre-processing techniques, as well as the robustness of our post-processing method in enhancing data classification post pre-processing. It is important to note, as verified by experimentation, that our post-processing method really applies to reducing the FPF produced by a pre-processor. The ANNs as we have applied them are unable to take data that was pre-classified as background data and re-classify it as anomalous. This emphasizes the importance of ensuring a quality problem definition. If the most important thing is minimizing false positives, it may be reasonable for some of the true anomalies to be classified as background data, as long as it is not to the extent that it impinges upon the true classification of the background data. However, if the most important thing is to find the anomalies, the true positives, then it is critical that any pre-processing technique utilized pushes all of the true anomalies to the anomalous class,
while ensuring that the anomalous class is not so large as to make it nearly indistinguishable from the background class.

**Summary**

We tested a variety of pre-processing methods with our post-processing methodology, allowing for and making some parameter variations along the way. The different pre-processing methods changed how the post-processor interacted with the data sets, but this helped to gain insight into the overall validity of the post-processor as applied to data pre-processed using a variety of techniques. It implied future research and validity in applying this post-processing method to other pre-processors to further reduce Type I error associated with the classification of anomalies within data. Hopefully, in future research, we can enhance the number of true positives; thereby, reducing false negatives as well. But we seemingly have had the most success thus far in reducing the false positives associated with the Type I error. As we progress in this research, there is ample promise for the enhancement of this process, recognizing that in these images in particular, given the state of the world today, false positives are by far the most negative outcome. Therefore, by enhancing our thresholds, parameters and performance, we can amply improve performance in line with our desired outcome.
V. Conclusions and Recommendations

Chapter Overview

In assessing our base methodology, we observed some improved classification accuracy with ARCADE, but this discovery also led us to examine other pre-possessing possibilities. Originally, we executed BACON on the raw data. Our first and second excursions were to assess ARCADE’s variance executing BACON on the principal component scores, with full and reduced dimensionality. This was followed by the use of the RX algorithm and finally evaluations based on the Mahalanobis distances. This use of a variety of pre-processing techniques allowed us to experiment with ARCADE. Simultaneously, we experimented with parameter adjustments within the pre-processors themselves.

In this chapter, we first provide a more in-depth examination of how the BACON and RX algorithms interacted with the data, assessing their true ability in performing anomaly detection. We assessed the algorithmic classification, as well as the values which produced the partitioning of classes for the various algorithms. This allows us to further examine the variations associated with both algorithms and both algorithms outputs. Assessments across multiple pre-processing algorithms and parameters, allowed us to determine robustness of a variety of features within the pre-processors themselves and ARCADE.

By examining algorithmic components separately and together it provides great insights into the future potential and the options for future research. While we initially set out with a focus on post-processing through ARCADE, this research gave us great
insight into the use of the various pre-processors, impacts of those pre-processors, clustering and random number generation.

**Conclusions of the Research**

Interestingly, in using a variety of pre-processing techniques, we were able to compare the robustness of those various algorithms. Figure 49 shows the class separation using three primary pre-processing methodologies. While the threshold or parameters could be changed for each of them, we see that they all pick up a majority of the true anomalies, but in some, the amount of noise is certainly much higher. In fact RX did not pick up all of the truly anomalous data and it included substantial noise, whereas the Mahalanobis distance pre-processing seemed to primarily pick up that noise associated with sensor artifacts, which in truth is anomalous – just not the anomalous data important for action.

![Figure 49: Class Mapping Pre-Processing Comparison (ARES1F)](image)

As demonstrated in Section IV, we also see that parameter shifts have the potential to make fairly significant changes in the quality of classification (Figure 50). BACON, utilizing 40 degrees of freedom on the right, includes all of the true anomalies in the
anomalous class, but the noise is not overbearing, as the degrees of freedom are reduced
to 30 in the middle, and then with 20 degrees of freedom on the right, where it almost
consumes the entire image. When too much of the true background data is over-classified
into the anomalous class, it becomes nearly impossible for additional processing to
remove enough data from the anomalous class to produce a quality result.

![Figure 50: Class Mapping BACON Dimensionality (ARES1F)](image)

However, we must recall that, in the instance of ARCADE, post-processing is
only able to improve upon the false positive classification associated with the classes. It
does not impact those observations that were inadvertently consumed by the background
class. Therefore, if true anomalies are included in the background class it produces two
negatives. First, it reduces the ability of the algorithm to find all of the true anomalies.
Second, anomalous data included in the background class has the potential to alter the
clusters associated with k-means, and thereby it could alter the effectiveness of the ANNs
in encoding those clusters, which potentially could cause more of the true anomalies
associated with the anomalous class to move into the background class. This may be
intuitive, but it re-affirms the over-arching importance in not under-partitioning the anomalous class.

Simultaneously, the thresholding robustness made a significant difference as well. Figure 51 shows how as the threshold decreases, more and more of the data remains in the background class. As we saw in Section IV, a threshold of 0.32 allows for too much background within the anomalous class, while a threshold of 0.05 significantly over-limits the anomalous class (Figure 51). Like the shifting parameters in Figure 50 this can have negative effects on the overall classification capability associated with ARCADE.

![Class Mapping Comparison (ARES1D)](image)

**Figure 51: Class Mapping Comparison (ARES1D)**

How the clustering interacted with the data is definitely an issue that can and should be explored more with future research. While overall the number of clusters had relatively similar predictability, there were definitely instances where a specific number of clusters definitely under-performed or over-performed.

Overall the utilization of ANNs was both a novelty and a positive. The literature review revealed that this was a novel idea, and while dependent on the settings, it
definitely showed promise in reducing false positives. When examining other research, we note that other pre-processing algorithms have been shown to hold promise when compared to both RX and BACON, so additional study in the application of ARCADE in conjunction with these other algorithms would definitely be worthwhile [16].

**Significance of Research**

The major additive effects of this research include the noticeable classification capacity of Mahalanobis distance. With the appropriate thresholding, Mahalanobis distance is one of the easiest methods for classification, providing high computational efficiency, while providing a high level of predictability. Using Mahalanobis distance has been a component of numerous classification techniques [9], but our research indicates that, at least with the use of HSI images collected with the HYDICE sensor, it provides a high classification accuracy in and of itself.

Of additional importance, our research shows how the clusters achieve a high degree of effectiveness with only a relatively small proportion of the associated observations. When we compared the classification ability utilizing only a 10 percent sample of each cluster to that of a 40 percent sample, the overall algorithmic performance was only slightly affected, and not always with an improved result. While computational efficiency was not the overall goal of this research, this is notable in that the ARCADE ran fairly quickly despite having literally dozens of moving parts, which could in reality be scaled down, by slightly tightening the number of clusters or reducing the number of ANNs run upon each cluster. While current evidence suggests that three ANNs do not provide enough robustness in performance, 20 ANNs seems to be more than necessary.
Of similar importance to the computational efficiency, the 10 percent sample directly demonstrates the continuity of the clusters, confirming the general quality of k-means clustering.

Subsequently, this research provided a potentially viable methodology for post-processing using ARCADE. At minimum, it establishes a starting point into future research utilizing ANNs in post-processing classification. This combination of techniques had not been attempted before, but we observed a high level of fidelity when comparing the potential to other known methods. As ANNs expand in their use and application, the potential definitely exists to further enhance pattern classification and anomaly detection.

Lastly, this research firmly asserts that all of these methodologies have potential in decision-making based upon the goals associated with the analysis. While in a perfect world, one would classify data and immediately find all of the true positives without acquiring any false negative observations; this is not typically the case. However, if we could use these techniques in accordance with our goals, there are many ways which it could be immediately applicable. In some cases, a few false positives are not very problematic, but it can have dire consequences not to find all of the anomalies. In other cases, false positives are absolutely egregious, but if even half of the anomalies could be resolved, it would create a significant benefit.
Recommendations for Action

It would benefit numerous problem sets, especially within the realm of HSI, to utilize methodological capacities in conjunction with desired intent to either eliminate all false negatives or enhance the classification of true positives.

Recommendations for Future Research

We would recommend a number of future research options to include:

1) ANN Testing
   - Testing for the optimal number or range of middle nodes used by the ANNs
   - Testing for the optimal number or range of hidden layers in ANNs
2) Signature matching of the anomalous class following completion of ARCADE
3) Testing additional pre-processing methodologies
4) Testing across variations and excursions with Horn’s Curve
5) Further parameter testing for the pre-processing methodologies already tested
6) Optimize sample size
7) Optimize the number of clusters
8) Test various clustering techniques
9) Run replications

Summary

The development of ARCADE provided a first-in-its-class classification technique utilizing ANNs as a post-processor for data sets that have already been classified utilizing an already proven pre-processing technique. ARCADE strings together a variety of mathematical techniques to filter out false positives garnered during the original classification. While there is still testing to be done, ARCADE shows promise in this capacity. By developing and testing ARCADE, it provided additional insight into the robustness of the various classification methods used as a pre-processors, it gave tremendous insight into the validity and quality of Mahalanobis distance as classification tool, it demonstrated the robustness of k-means clustering and small sample
sizes based on the associated clusters, and it showed promise in the reduction of false positives. With certainty there is still work to be done, but the information attained will advance future research possibilities with the possibility of finding ever better anomaly detection techniques and procedures. Furthermore, this research affirms the necessity in setting quality goals for anomaly detection. If we can ascertain the intent of our anomaly detection, we correctly frame problems to have immediate, actionable insights into varying problem sets.
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Autoencoded Reduced Clusters for Anomaly Detection Enrichment (Arcade) In Hyperspectral Imagery

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14. ABSTRACT
Anomaly detection in hyper-spectral imagery is a relatively recent and important research area. The sheer amount of data available in many hyper-spectral images makes the utilization of multivariate statistical methods and artificial neural networks ideal for this analysis. Using HYDICE sensor hyperspectral images, we examine a variety of pre-processing techniques within a framework that allows for changing parameter settings and varying the methodological order of operations in order to enhance detection of anomalies within image data. By examining a variety of different options, we are able to gain significant insight into what makes anomaly detection viable for these images, as well as what impact parameter and methodology changes can have on the total classification effectiveness, false positive fraction and true positive fraction regarding classification.

15. SUBJECT TERMS
Anomaly Detection, Clustering, Reconstructive Error, Mahalanobis Distance

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