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Hadoop neural network for parallel and distributed feature selection

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Abstract

In this paper, we introduce a theoretical basis for a Hadoop-based neural network for parallel and distributed feature selection in Big Data sets. It is underpinned by an associative memory (binary) neural network which is highly amenable to parallel and distributed processing and fits with the Hadoop paradigm. There are many feature selectors described in the literature which all have various strengths and weaknesses. We present the implementation details of five feature selection algorithms constructed using our artificial neural network framework embedded in Hadoop YARN. Hadoop allows parallel and distributed processing. Each feature selector can be divided into subtasks and the subtasks can then be processed in parallel. Multiple feature selectors can also be processed simultaneously (in parallel) allowing multiple feature selectors to be compared. We identify commonalities among the five feature selectors. All can be processed in the framework using a single representation and the overall processing can also be greatly reduced by only processing the common aspects of the feature selectors once and propagating these aspects across all five feature selectors as necessary. This allows the best feature selector and the actual features to select to be identified for large and high dimensional data sets through exploiting the efficiency and flexibility of embedding the binary associative-memory neural network in Hadoop.

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1. Introduction

The meaning of “big” with respect to data is specific to each application domain and dependent on the computational resources available. Here we define “Big Data” as large, dynamic collections of data that cannot be processed using traditional techniques, a definition adapted from (Franks, 2012; Zikopoulos & Eaton, 2011). Today, data is generated continually by an increasing range of processes and in ever increasing quantities driven by Big Data mechanisms such as cloud computing and on-line services. Business and scientific data from many fields, such as finance, astronomy, bioinformatics and physics, are often measured in terabytes (10^{12} bytes). Big Data is characterised by its complexity, variety, speed of processing and volume (Laney, 2001). It is increasingly clear that exploiting the power of these data is essential for information mining. These data often contain too much noise (Liu, Motoda, Setiono, & Zhao, 2010) for accurate classification (Dash & Liu, 1997; Han & Kamber, 2006), prediction (Dash & Liu, 1997; Guyon & Elisseeff, 2003) or outlier detection (Hodge, 2011). Thus, only some of the features (dimensions) are related to the target concept (classification label or predicted value). Also, if there are too many data features then the data points become sparse. If data is too sparse then distance measures such as the popular Euclidean distance and the concept of nearest neighbours become less applicable (Ertöz, Steinbach, & Kumar, 2003). Many machine learning algorithms are adversely affected by this noise and these superfluous features in terms of both their accuracy and their ability to generalise. Consequently, the data must be pre-processed by the classification or prediction algorithm itself or by a separate feature selection algorithm to prune these superfluous features (Kohavi & John, 1997; Witten & Frank, 2000).

The benefits of feature selection include: reducing the data size when superfluous features are discarded, improving the classification/prediction accuracy of the underlying algorithm where the algorithm is adversely affected by noise, producing a more compact and easily understood data representation and reducing the execution time of the underlying algorithm due to the smaller data size. Reducing the execution time is extremely important for Big Data, which has a high computational resource demand on memory and CPU time.

In this paper, we focus on feature selection in vast data sets for parallel and distributed classification systems. We aim to remove...
noise and reduce redundancy to improve classification accuracy. There is a wide variety of techniques proposed in the machine learning literature for feature selection including Correlation-based Feature Selection (Hall, 1998), Principal Component Analysis (PCA) (Jolliffe, 2002), Information Gain (Quinlan, 1986), Gain Ratio (Quinlan, 1992), Mutual Information Selection (Wetscherek, 1994), Chi-square Selection (Liu & Setiono, 1995), Probabilistic Las Vegas Selection (Liu & Setiono, 1996) and Support Vector Machine Feature Elimination (Guyon, Weston, Barnhill, & Vapnik, 2002). Feature selectors produce feature scores. Some feature selectors also select the best set of features to use while others just rank the features with the scores. For these feature rankers, the best set of features must then be chosen by the user, for example, using greedy search (Witten & Frank, 2000).

It is often not clear to the user which feature selector to use for their data and application. In their analysis of feature selection, Guyon and Elisseeff (2003) recommend evaluating a variety of feature selectors before deciding the best for their problem. Therefore, we propose that users exploit our framework to run a variety of feature selectors in parallel and then evaluate the feature sets chosen by each selector using their own specific criteria. Having multiple feature selectors available also provides the opportunity for ensemble feature selection where the results from a range of feature selectors are merged to generate the best set of features to use. Feature selection is a combinatorial problem so needs to be implemented as efficiently as possible particularly on big data sets. We have previously developed a k-NN classification (Hodge & Austin, 2005; Weeks, Hodge, O’Keefe, Austin, & Lees, 2003) and prediction algorithm (Hodge, Krishnan, Austin, & Polak, 2011) using an associative memory (binary) neural network called the Advanced Uncertain Reasoning Architecture (AURA) (Austin, 1995). This multi-faceted k-NN motivated a unified feature selection framework exploiting the speed and storage efficiency of the associative memory neural network. The framework lends itself to parallel and distributed processing across multiple nodes allowing vast data sets to be processed. This could be done by processing the data at the same geographical location using a single machine with multiple processing cores (Weeks, Hodge, & Austin, 2002) or at the same geographical location using multiple compute nodes (Weeks et al., 2002) or even distributed processing of the data at multiple geographical locations.

Data mining tools such as Weka (Witten & Frank, 2000), Matlab, R and SPSS provide feature selection algorithms for data mining and analytics. However, these products are designed for small scale data analysis. Researchers have parallelised individual feature selection algorithms using MapReduce/Hadoop (Chu et al., 2007; Reggiani, 2013; Singh, Kubica, Larsen, & Sorokina, 2009; Sun, 2014). Data mining libraries such as Mahout (https://mahout.apache.org/) and MLlib (https://spark.apache.org/mllib/) and data mining frameworks such as Radoop (https://rapidminer.com/products/radoop/) include a large number of data mining algorithms including feature selectors. However, they do not explicitly tackle processing reuse with a view to multi-user and multi-task resource allocation. Zhang, Kumar, and Ré (2014) developed a database systems framework for optimised feature selection providing a range of algorithms. They observed that there are reuse opportunities that could yield orders of magnitude performance improvements on feature selection workloads as we will also demonstrate here using AURA in an Apache Hadoop (https://hadoop.apache.org/) framework.

The main contributions of this paper are:

- To extend the AURA framework to parallel and distributed processing of vast data sets in Apache Hadoop,
- To describe five feature selectors in terms of the AURA framework. Two of the feature selectors have been implemented in AURA but not using Hadoop (Hodge, Jackson, & Austin, 2012; Hodge, O’Keefe, & Austin, 2006) and the other three have not been implemented in AURA before,
- To theoretically analyse the resulting framework to show how the five feature selectors have common requirements to enable reuse.
- To theoretically analyse the resulting framework to show how we reduce the number of computations. The larger the data set then the more important this reduction becomes.
- To demonstrate parallel and distributed processing in the framework allowing Big Data to be analysed.

In our AURA framework, the feature selectors all use one common data representation. We only need to process any common elements once and can propagate the common elements to all feature selectors that require them. Thus, we can rapidly and efficiently determine the best feature selector and the best set of features to use for each data set under investigation. In Section 2, we discuss AURA and related neural networks and how to store and retrieve data from AURA, Section 3 demonstrates how to implement five feature selection algorithms in the AURA unified framework and Section 4 describes parallel and distributed feature selection using AURA. We then analyse the unified framework in Section 5 to identify common aspects of the five feature selectors and how they can be implemented in the unified framework in the most efficient way. Section 6 details the overall conclusions from our implementations and analyses.

2. Binary neural networks

AURA (Austin, 1995) is a hetero-associative memory neural network (Palin, 2013). An associative memory is addressable through its contents and a hetero-associative memory stores associations between input and output vectors where the vectors are different (Palin, 2013). AURA uses binary Correlation Matrix Memories (CMMs): binary hetero-associative matrices that store and retrieve patterns using matrix calculus. They are non-recursive and fully connected. Input vectors (stimuli) address the CMM rows and output vectors address the CMM columns. Binary neural networks have a number of advantages compared to standard neural networks including rapid one-pass training, high levels of data compression, computational simplicity, network transparency, a partial match capability and a scalable architecture that can be easily mapped onto high performance computing platforms including parallel and distributed platforms (Weeks et al., 2002). AURA is implemented as a C++ software library.

Previous parallel and distributed applications of AURA have included distributed text retrieval (Weeks et al., 2002), distributed time-series signal searching (Fletcher, Jackson, Jessop, Liang, & Austin, 2006) and condition monitoring (Austin, Brewer, Jackson, & Hodge, 2010). This new development will augment these existing techniques and is aimed at these same domains. It will couple feature selection, classification and prediction with the speed and storage efficiency of a binary neural network allowing parallel and distributed data mining. This makes AURA ideal to use as the basis of an efficient distributed machine learning framework. A more formal definition of AURA, its components and methods now follows.

2.1. AURA

The AURA methods use binary input I and output O vectors to efficiently store records in a CMM M as in Eq. (1) using the binary
Training (construction of a CMM) is a single epoch process with one training step for each input–output association (each $I_j O^T_j$ in Eq. (1)) which equates to one step for each record $j$ in the data set. Thus, the trained CMM $M$ represents $(I_1 \times O^T_1), (I_2 \times O^T_2), \ldots (I_n \times O^T_n)$ superimposed using bitwise or. $I_j O^T_j$ is an estimate of the weight matrix $W(j)$ of the synaptic connections of the neural network as a linear associator with binary weights. $W(j)$ forms a mapping representing the association described by the $j$th input/output pair of vectors. As a consequence of using unipolar elements (0,1) throughout, the value at each matrix component $w_{ij}$ means the existence of an association between elements $i$ and $j$. The trained CMM $M$ is then effectively an encoding (correlation) of the $N$ weight matrices $W$ for all $N$ records in the data set. Individual weights within the weight matrix update using a generalisation of Hebbian learning (Hebb, 1949) where the state for each synapse (matrix element) is binary valued. Every synapse can update its weight independently using a local learning rule (Palm, 2013). Local learning is biologically plausible and computationally simple allowing parallel and rapid execution. The learning process is illustrated in Fig. 1.

For feature selection, the data are stored in the CMM which forms an index of all features in all records. During training, the input vectors $I_j$ represent the feature and class values and are associated with a unique output vector $O_j$ representing a record. Fig. 1 shows a trained CMM. In this paper, we set only one bit in the vector $O_j$ indicating the location of the record in the data set, the first record has the first bit set, the second record has the second bit set etc. Using a single set bit makes the length of $O_j$ potentially large. However, exploiting a compact list representation (Hodge & Austin, 2001) (more detail is provided in Section 4.3.1) means we can compress the storage representation.

2.2. Data

The AURA feature selector, classifier and predictor framework can handle qualitative features (symbolic and discrete numeric) and quantitative features (continuous numeric).

The raw data sets need pre-processing to allow them to be used in the binary AURA framework. Qualitative features are enumerated and each separate token maps onto an integer ($Token \rightarrow Integer$) which identifies the bit to set within the vector. For example, a SEX_TYPE feature would map as ($F \rightarrow 0$) and ($M \rightarrow 1$).

Any quantitative features are quantised (mapped to discrete bins) (Hodge & Austin, 2012). Each individual bin maps onto an integer which identifies the bit to set in the input vector. Next, we describe the simple equi-width quantisation. We note that the Correlation-Based Feature Selector described in Section 3.2 uses a different quantisation technique to determine the bin boundaries. However, once the boundaries are determined, the mapping to CMM rows is the same as described here.

To quantise quantitative features, a range of input values for feature $F_j$ map onto each bin. Each bin maps to a unique integer as in Eq. (2) to index the correct location for the feature in $I_j$. In this paper, the range of feature values mapping to each bin is equal to subdivide the feature range into $b$ equi-width bins across each feature.

$$R_k \rightarrow bins_{f_k} \leftrightarrow Integer_{f_k} + offset(F_j)$$

where $F_j \in F, f_j$ is a value of $F_j$ and cardinality ($Integer_{f_k}$) (2)

$$\equiv cardinality(bins_{f_k}).$$

In Eq. (2), offset($F_j$) is a cumulative integer offset within the binary vector for each feature $F_j$, $\rightarrow$ is a many-to-one mapping and $\leftrightarrow$ is a one-to-one mapping. The offset for the next feature $F_{j+1}$ is given by offset($F_{j+1}$) = offset($F_j$) + nbins($F_j$) where nbins($F_j$) is the number of bins for feature $F_j$.

For each record in the data set.

For each feature.

Calculate bin for feature value.

Set bit in vector as in Eq. (2).

2.3. AURA recall

To recall the matches for a query (input) record, we firstly produce a recall input vector $R_k$ by quantising the target values for each feature to identify the bins (CMM rows) to activate as in Eq. (3). During recall, the presentation of recall input vector $R_k$ elicits the recall of output vector $O_k$, as vector $R_k$ contains all of the addressing information necessary to access and retrieve vector $O_k$. Recall is effectively the dot product of the recall input vector $R_k$ and CMM $M$, as in Eq. (3) and Fig. 2.

$$S^T = R_k^T \cdot M.$$
output vector multiplied by a weight based on the dot product of the input vector with itself. If the recall input \( R_k \) is not from the original training set, then the system will recall the output \( O_k \) associated with the closest stored input to \( R_k \), based on the dot product between the test and training inputs.

Matching is a combinatorial problem but can be achieved in a single pass in AURA. AURA can also exploit the advantages of sparse vectors (Palm, 2013) during recall by only activating regions of interest. If the input vector \( R_k \) has 1000 bits indexing 1000 CMM rows then only the rows addressed by a set bit in the input vector need be examined (as shown in Figs. 2 and 3). For a 10 bit set vector then only 10 of the 1000 rows are activated. The input pattern \( R_k \) would be said to have a saturation of \((10/1000 = 0.01)\). The total amount of data that needs to be examined is reduced by a factor that is dependent on this saturation providing that the data is spread reasonably evenly between the rows and the CMM is implemented effectively. Using smart encoding schemes can bring the performance improvement resulting from very low saturation input patterns to over 100-fold (Weeks et al., 2002).

The AURA technique thresholds the summed output \( S \) to produce a binary output vector \( T \) as given in Eq. (4).

\[
T_j = \begin{cases} 
1 & \text{if } S_j \geq \theta \\
0 & \text{otherwise.}
\end{cases}
\]

(4)

For exact match, we use the Willshaw threshold (Willshaw, Buneman, & Longuet-Higgins, 1969) to set \( \theta \). This sets a bit in the thresholded output vector for every location in the summed output vector that has a value higher than or equal to \( \theta \). The value of \( \theta \) varies according to the task. If there are ten features in the data and we want to find all stored records that match the ten feature values of the input vector then we set \( \theta \) to 10. Thus, for full match \( \theta = b^1 \), where \( b^1 \) is set to the number of set bits in the input vector. For partial matching, we use the L-Max threshold (Casasent & Telfer, 1992). L-Max thresholding essentially retrieves at least \( L \) top matches. Our AURA software library automatically sets \( \theta \) to the highest integer value that will retrieve at least \( L \) matches.

Feature selection described in Section 3 requires both exact matching using Willshaw thresholding and partial matching using L-Max thresholding.

3. Feature selection

There are two fundamental approaches to feature selection (Kohavi & John, 1997; Witten & Frank, 2000): (1) filters select the optimal set of features independently of the classifier/predictor algorithm while (2) wrappers select features which optimise classification/prediction using the algorithm. We examine the mapping of five filter approaches to the binary AURA architecture. Filter approaches are more flexible than wrapper approaches as they are not directly coupled to the algorithm and are thus applicable to a wide variety of classification and prediction algorithms. Our method exploits the high speed and efficiency of the AURA techniques as feature selection is a combinatorial problem.

We examine a mutual information approach Mutual Information Feature Subset Selection (MI) detailed in Section 3.1 that analyses features on an individual basis, a correlation-based multivariate filter approach Correlation-based Feature Subset Selection (CFS) detailed in Section 3.2 that examines greedily selected subsets of features, a revised Information Gain approach Gain Ratio (GR) detailed in Section 3.3, a feature dependence approach Chi-Square Feature Selection (CS) detailed in Section 3.4 which is univariate, and a univariate feature relevance approach Odds Ratio (OR) detailed in Section 3.5.

Univariate filter approaches such as MI, CS or OR are quicker than multivariate filters as they do not need to evaluate all combinations of subsets of features. The advantage of a multivariate filter compared to a univariate filter lies in the fact that a univariate approach does not account for interactions between features. Multivariate techniques evaluate the worth of feature subsets by considering both the individual predictive ability of each feature and the degree of redundancy between the features in the set.

All five feature selection algorithms have their relative strengths. We refer the reader to Forman (2003) and Varela, Martins, Aguiar, and Figueiredo (2013) for accuracy evaluations of these feature selectors. These papers show that the best feature selector varies with data and application. Using the CFS attribute selector, Hall and Smith (1998) found significant improvement in classification accuracy of k-NN on five of the 12 data sets they evaluated but a significant degradation in accuracy on two data sets. Hence, different feature selectors are required for different data sets and applications.

We note that the CFS as implemented by Hall (1998) uses an entropy-based quantisation whereas we have used equi-width quantisation for the other feature selectors (MI, GR, CS and OR). We plan to investigate unifying the quantisation as a next step. For the purpose of our analysis in Section 5, we assume that all feature selectors are using identical quantisation. We assume that all records are to be used during feature selection.

3.1. Mutual information feature selection

Wetttscherek (1994) described a mutual information feature selection algorithm. The mutual information between two features is “the reduction in uncertainty concerning the possible values of one feature that is obtained when the value of the other feature is determined” (Wetttscherek, 1994). MI is defined by Eq. (5):

\[
MI(F_j, C) = \sum_{i=1}^{b(f_j)} \sum_{c=1}^{n_{class}} p(C = c \land F_j = f_i) \cdot \log_2 \left( \frac{p(C = c \land F_j = f_i)}{p(C = c) \cdot p(F_j = f_i)} \right).
\]

(5)

To calculate \( p(C = c \land F_j = f_i) \), we use AURA to calculate \( \frac{n(BV_i \land BV_c)}{n} \).

AURA excites the row in the CMM corresponding to feature \( f_i \) of feature \( F_j \) and the row in the CMM corresponding to class value \( c \) as shown in Fig. 3. By thresholding the output vector \( S \) at Willshaw threshold value = 2, we obtain a thresholded output vector with a bit set for every co-occurrence. We can count these set bits to determine the co-occurrence count. Furthermore,
p (C = c) is the count of the number of set bits $n(BV_{C})$ in the binary vector (CMM row) for c and $p(f_i = f_i)$ is the count of the number of set bits $n(BV_{f_i})$ in the binary vector (CMM row) for $f_i$ as used by GR.

The MI calculated using AURA for qualitative features is given by Eq. (6) where $N$ is the number of records in the data set, $rows(F_j)$ is the number of CMM rows for feature $F_j$ and $nClass$ is the number of classes:

$$MI \left( F_j, C \right) = \sum_{c=1}^{nClass} \frac{n(BV_{f_i} \land BV_{C})}{N} \cdot \log_2 \left( \frac{n(BV_{f_i} \land BV_{C})}{nClass \cdot N} \right)$$

(6)

We can follow the same process for real/discrete ordered numeric features in AURA. In this case, the mutual information is given by Eq. (7):

$$MI \left( F_j, C \right) = \sum_{b=1}^{bins(F_j)} \sum_{f=1}^{nClass} \frac{n(BV_{f_i} \land BV_{C})}{N} \cdot \log_2 \left( \frac{n(BV_{f_i} \land BV_{C})}{nClass \cdot N} \right)$$

(7)

where $bins(F_j)$ is the number of bins (effectively the number of rows) in the CMM for feature $F_j$ and $BV_{f_i}$ is the CMM row for the bin mapped to by feature value $f_i$.

The MI feature selector assumes independence of features and scores each feature separately so it is the user’s prerogative to use this number of bits set in the vector (CMM row) for $f_i$ to determine the number of features to select. The major drawback of the MI feature selector along with similar information theoretic approaches, for example Information Gain, is that they are biased towards features with the largest number of distinct values as this splits the training records into nearly pure classes. Thus, a feature with a distinct value for each record has a maximal information score. The CFS and GR feature selectors make adaptations of information theoretic approaches to prevent this biasing.

3.2. Correlation-based feature subset selection

Hall (1998) proposed the Correlation-based Feature Subset Selection (CFS). It measures the strength of the correlation between pairs of features. CFS favours feature subsets that contain features that are highly correlated to the class but uncorrelated to each other to minimise feature redundancy. CFS is thus based on information theory measured using Information Gain. Hall and Smith (1997) used a modified Information Gain measure, Symmetrical Uncertainty, $SU$ given in Eq. (8) to prevent bias towards features with many distinct values (Section 3.1). $SU$ estimates the correlation between features by normalising the value in the range $[0, 1]$. Two features are completely independent if $SU = 0$ and completely dependent if $SU = 1$.

$$SU \left( F_j, G_i \right) = 2.0 \cdot \frac{\text{Ent} \left( F_j \right) - \text{Ent} \left( F_j \mid G_i \right)}{\text{Ent} \left( F_j \right) + \text{Ent} \left( G_i \right)}$$

(8)

where the entropy of a feature $F_j$ for all feature values $f_i$ is given as Eq. (9):

$$\text{Ent} \left( F_j \right) = -\sum_{f=1}^{n(f_j)} p(f_i) \log_2 (p(f_i))$$

(9)

and the entropy of feature $F_j$ after observing values of feature $G_i$ is given as Eq. (10):

$$\text{Ent} \left( F_j \mid G_i \right) = -\sum_{k=1}^{n(G_i)} \sum_{f=1}^{n(f_j)} p(g_k) p(f_i \mid g_k) \log_2 (p(f_i \mid g_k))$$

(10)

Any quantitative features are discretised using Fayyad and Irani’s entropy quantisation (Fayyad & Irani, 1993). The bin boundaries are determined using Information Gain and these quantisation bins map the data into the AURA CMM as previously.

CFS has many similarities to MI when calculating the values in Eqs. (8)–(10) and through using the same CMM (Fig. 3) as noted below.

In the AURA CFS, for each pair of features $(F_j, G_i)$ to be examined, the CMM is used to calculate $Ent (F_j)$, $Ent (G_i)$ and $Ent (F_j \mid G_i)$ from Eqs. (8)–(10). There are three parts to the calculation.

1. $Ent (F_j)$ requires the count of data records for the particular value $f_i$ of feature $F_j$ which is $n(BV_{f_i})$ in Eq. (6) for qualitative and class features and $n(BV_{f_i} \land BV_{G})$ in Eq. (7) for quantitative features. AURA excites the row in the CMM corresponding to feature value $f_i$ of feature $F_j$. This row is a binary vector (BV) and is represented by $BV_{f_i}$. A count of bits set on the row gives $n(BV_{f_i})$ from Eq. (6) and is achieved by thresholding the output vector $S_k$ from Eq. (4) at Willshaw value 1.

2. Similarly, $Ent (G_i)$ counts the number of records where feature $G_i$ has value $g_k$.

3. $Ent (F_j \mid G_i)$ requires the number of co-occurrences of a particular value $f_i$ of feature $F_j$ with a particular value $g_k$ of feature $G_i$ defined in Eq. (6) for qualitative features and $n(BV_{f_i} \land BV_{G})$ for quantitative features and between a feature and the class $n(BV_{f_i} \land BV_{C})$ and $n(BV_{f_i} \land BV_{C})$ for qualitative and quantitative features respectively. If both the feature value row and the class values row are excited then the summed output vector will have a two in the column of every record with a co-occurrence of $f_i$ with $g_k$ as shown in Fig. 3. By thresholding the summed output vector at a threshold of two, we can find all co-occurrences. We represent this number of bits set in the vector $n(BV_{f_i} \land BV_{G})$ which is a count of the set bits when $BV_{C}$ logically anded with $BV_{f_i}$.

CFS determines the feature subsets to evaluate using forward search. Forward search works by greedily adding features to a subset of selected features until some termination condition is met whereby adding new features to the subset does not increase the discriminatory power of the subset above a pre-specified threshold value. The major drawback of CFS is that it cannot handle strongly interacting features (Hall & Holmes, 2003).

3.3. Gain ratio feature selection

Gain Ratio (GR) (Quinlan, 1992) is a new feature selector for the AURA framework. GR is a modified Information Gain technique and is used in the popular machine learning decision tree classifier C4.5 (Quinlan, 1992). Information Gain is given in Eq. (11) for feature $F_j$ and the class $C$. CFS (Section 3.2) modifies Information Gain to prevent biasing towards features with the most values. GR is an alternative adaptation which considers the number of splits (number of values) of each feature when calculating the score for each feature using normalisation.

$$Gain \left( F_j, C \right) = Ent \left( F_j \right) - Ent \left( F_j \mid C \right)$$

(11)

where $Ent \left( F_j \right)$ is defined in Eq. (9) and $Ent \left( F_j \mid C \right)$ is defined by Eq. (10). Then Gain Ratio is defined as Eq. (12):

$$GainRatio \left( F_j, C \right) = \frac{Gain \left( F_j, C \right)}{IntrinsicValue \left( F_j \right)}$$

(12)

where $IntrinsicValue$ is given by Eq. (13):

$$IntrinsicValue \left( F_j \right) = \sum_{p=1}^{\psi} \frac{S_p \log_2 \left( \frac{S_p}{N} \right)}{N}$$

(13)
and \( V \) is the number of feature values \((n(f_i))\) for qualitative features and number of quantisation bins \( n(b_i) \) for quantitative features and \( S_p \) is a subset of the records that have \( f_i = f_j \) for qualitative features or map to the quantisation bin \( b_i(f_j) \) for quantitative features.

To implement GR using AURA, we train the CMM as described in Section 2.1. We can then calculate \( \text{Ent}(f_i) \) and \( \text{Ent}(f_j | c) \) as per the CFS feature selector described in Section 3.2 to allow us to calculate \( \text{Gain}(f_i, c) \). To calculate \( \text{IntrinsicValue}(f_i) \) we need to calculate the number of records that have particular feature values. This is achieved by counting the number of set bits \( n(BVf_i) \) in the binary vector (CMM row) for \( f_i \) for qualitative features or \( n(BVb_i) \) in the binary vector for the quantisation bin \( b_i \) for quantitative features. We can store counts for the various feature values and classes as we proceed so there is no need to calculate any count more than once.

The main disadvantage of GR is that it tends to favour features with low Intrinsic Value rather than high gain by overcompensating towards a feature just because its intrinsic information is very low.

3.4. Chi-square algorithm

We now demonstrate how to implement a second new feature selector in the AURA framework. The Chi-Square (CS) (Liu & Setiono, 1995) algorithm is a feature ranker like MI, OR and GR rather than a feature selector; it scores the features but it is the user’s prerogative to select which features to use. CS assesses the independence between a feature \((f_i)\) and a class \((c)\) and is sensitive to feature interactions with the class. Features are independent if CS is close to zero. Forman (2003) and Yang and Pedersen (1997) conducted evaluations of filter feature selectors and found that CS is among the most effective methods of feature selection for classification.

Chi-Square is defined as Eq. (14):

\[
\chi^2(f_i, c) = \sum_{i=1}^{n} \sum_{c=1}^{cClass} \frac{N ! (wz - yx)^2}{(w + y) (x + z) (w + x) (y + z)}
\]

where \( b(f_i) \) is the number of bins (CMM rows) representing feature \( f_i \), \( cClass \) is the number of classes, \( w \) is the number of times \( f_i \) and \( c \) co-occur, \( x \) is the number of times \( f_i \) occurs without \( c \), \( y \) is the number of times \( c \) occurs without \( f_i \), \( z \) is the number of times neither \( c \) nor \( f_i \) occur. Thus, CS is predicated on counting occurrences and co-occurrences and, hence, has many commonalities with MI, CFS and GR.

- Fig. 3 shows how to produce a binary output vector \((BVf_i \land BVc)\) for qualitative features or \((BVb_i \land BVc)\) for quantitative features listing the co-occurrences of a feature value and a class value. It is then simply a case of counting the number of set bits \((1s)\) in the thresholded binary vector \( T \) in Fig. 3 to count \( w \).

- To count \( x \) for qualitative features, we logically subtract \((BVf_j \land BVc)\) from the binary vector \((BVf_i)\) to produce a binary vector and count the set bits in the resulting vector. For quantitative features, we subtract \((BVb_j \land BVc)\) from \((BVb_i)\) and count the set bits in the resulting binary vector.

- To count \( y \) for qualitative features, we can logically subtract \((BVf_j \land BVc)\) from \((BVc)\) and count the set bits and likewise for quantitative features we can subtract \((BVb_j \land BVc)\) from \((BVc)\) and count the set bits.

- If we logically or \((BVf_i)\) with \((BVc)\), we get a binary vector representing \((f_i = f_j) \lor (c = c)\) for qualitative features. For quantitative features, we can logically or \((BVb_i)\) with \((BVc)\) to produce \((f_i = b_i(f_j)) \lor (c = c)\). If we then logically invert this new binary vector, we retrieve a binary vector representing \( z \) and it is simply a case of counting the set bits to get the count for \( z \).

As with MI and OR, CS is univariate and assesses features on an individual basis selecting the features with the highest scores, namely the features that interact most with the class.

3.5. Odds ratio

The third new feature selector is Odds Ratio (OR) (see Forman, 2003). OR is another feature ranker. Standard OR is a two-class feature ranker although it can be extended to multiple classes. It is often used in text classification tasks as these are often two-class problems. It performs well particularly when used with Naïve Bayes Classifiers. OR reflects relevance as the likelihood (odds) of a feature occurring in the positive class normalised by that of the negative class. OR has many commonalities with MI, CFS and GR but particularly with CS where it requires the same four calculations \( w, x, y, \text{ and } z \) (defined above in Section 3.4). Odds Ratio is defined by Eq. (15):

\[
\text{OR}(f_i, c) = \frac{b(f_i)}{w(x + z)}
\]

where \( b(f_i) \) is the number of bins (CMM rows) representing feature \( f_i \), \( w \) is the number of times \( f_i \) and \( c \) co-occur, \( x \) is the number of times \( f_i \) occurs without \( c \), \( y \) is the number of times \( c \) occurs without \( f_i \), \( z \) is the number of times neither \( c \) nor \( f_i \) occur. Thus, OR is predicated on counting occurrences and co-occurrences. To avoid division by zero the denominator is set to 1 if \( xy \) evaluates to 0.

4. Parallel and distributed AURA

Feature selection is a combinatorial problem so a fast, efficient and scalable platform will allow rapid analysis of large and high dimensional data sets. AURA has demonstrated superior training and recall speed compared to conventional indexing approaches (Hodge & Austin, 2001) such as hashing or inverted file lists which may be used for data indexing. AURA trains 20 times faster than an inverted file list and 16 times faster than a hashing algorithm. It is up to 24 times faster than the inverted file list for recall and up to 14 times faster than the hashing algorithm. AURA k-NN has demonstrated superior speed compared to conventional k-NN (Hodge & Austin, 2005) and does not suffer the limitations of other k-NN optimisations such as the KD-tree which only scales to low dimensionality data sets (McCallum, Nigam, & Ungar, 2000). We showed in Hodge et al. (2006) that using AURA speeds up the MI feature selector by over 100 times compared to a standard implementation of MI.

For very large data sets, the data may be processed in parallel on one compute node (such as a multi-core CPU) or across a number of distributed compute nodes. Each compute node in a distributed system can itself perform parallel processing.

4.1. Parallel AURA

In Weeks et al. (2002), we demonstrated a parallel search implementation of AURA. AURA can be subdivided across multiple processor cores within a single machine or spread across multiple connected compute nodes. This parallel processing entails “striping” the CMM across several parallel subsections. The CMM is effectively subdivided vertically across the output vector as shown in Fig. 4. In the data, the number of features \( m \) is usually much less than the number of records \( N, m \ll N \). Therefore, we subdivide the data along the number of records \( N \) (column stripes) as shown in the leftmost example in Fig. 4.

Splitting the data across multiple CMM stripes using columns means that the CMM can store data as separate rows within a single
If a CMM contains large data it can be subdivided (striped) across a number of CMM stripes. In the left hand figure, the CMM is striped vertically (by time) and in the right hand figure the CMM is striped horizontally (by feature subsets). On the left, each CMM stripe produces a thresholded output vector $T_n$ containing the top $k$ matches (and their respective scores) for that stripe. All $T_n$ are aggregated to form a single output vector $T$ which is thresholded to list the top matches overall. On the right, each stripe outputs a summed output vector $S_n$. All $S_n$ are summed to produce an overall summed output vector which is thresholded to list the top matches overall.

If the number of features is large then it is possible to subdivide the CMMs further. The CMM is divided vertically by the records (column stripes) as before and then the column stripes are subdivided by the input features (row stripes). Subdivision by input features (row stripes) is shown in the rightmost diagram in Fig. 4. Dividing the CMM using the features (row stripes) makes assimilating the results more complex than assimilating the results for column stripes. Each row stripe produces a summed output vector containing column subtotals for those features within the stripe. The column subtotals need to be assimilated from all row stripes that hold data for that column. Thus, we sum these column subtotals to produce a column stripe vector $C$ holding the overall sum for each column in that stripe. Row striping involves assimilating integer vectors of length $c$ where $c$ is the number of columns for the column subdivision (column stripe).

4.2. Distributed AURA

There are two central challenges for distributed feature selection: firstly, maintaining a distributed data archive so that data does not have to be moved to a central repository and secondly, orchestrating the search process across the distributed data. Different data and applications will have different criteria that they wish to optimise. These could be optimising communication overhead, processing speed, memory usage or combinations of these criteria. Hence, there is unlikely to be a single best technique for distribution.

To distribute AURA, we use the striping mechanisms detailed in the previous section. However, rather than spreading the stripes within the cores of a multicore processor, we distribute the stripes across computers within a distributed network. The stripes need to be distributed for maximum efficiency. This can be to maximise processing speed, to minimise memory usage, to minimise communication overhead or a combination of criteria. Distributing the stripes requires an efficient distribution mechanism to underpin the procedure.

Orchestrated search with minimal data movement is provided by the open source software project: Apache Hadoop (Shvachko, Hairong, Radia, & Chansler, 2010). Hadoop operates on the premise that “moving computation is cheaper than moving data” (Borthakur, 2008). Hadoop allows the distributed processing of large data sets across clusters of commodity servers. It provides load balancing, is highly scalable and has a very high degree of fault tolerance. It is able to run on commodity hardware due to its ability to detect and handle failures at the application layer. There are multiple copies of the stored data so, if one server or node is unavailable, its data can be automatically replicated from a known good copy. If a compute node fails then Hadoop automatically re-balances the work load on the remaining nodes. Hadoop has demonstrated high performance for a wide variety of tasks (Borthakur et al., 2011). It was initially aimed at batch processing tasks so is ideally suited to the task of feature selection where the feature selector is trained with the training data and feature selection is run once on a large batch of test data. Hadoop is currently developing real-time processing capabilities. In this paper, we focus on batch processing and the implementation details of the five feature selectors using AURA with Hadoop.

Hadoop is highly configurable and can be optimised to the user’s specific requirements, for example, optimising to minimise memory overhead, optimising for fastest processing or optimising to reduce communication overhead. Hence, we do not attempt to evaluate Hadoop here. Instead, we focus on describing how to map AURA CMMs to Hadoop to create a feature evaluation framework.

There are two parts of Hadoop that we consider here: YARN which assigns work to the nodes in a cluster and the Hadoop Distributed File System (HDFS) which is a distributed file system spanning all the nodes in the Hadoop cluster with a single namespace.

YARN (Kumar et al., 2013) supersedes MapReduce in Hadoop. YARN is able to run existing MapReduce applications. YARN decouples resource management and scheduling from the data processing. This means that data can continue to be streamed into the system simultaneously with MapReduce batch jobs. YARN has a central resource manager that reconciles Hadoop system resources according to constraints such as queue capacities or user-limits. Node manager agents monitor the processing operations of individual nodes in the cluster. The processing is controlled by an ApplicationMaster which negotiates resources from the central resource manager and works with the node manager agents to execute and monitor the tasks. The actual MapReduce procedure, divides (maps) the processing into separate chunks which are processed in parallel. The outputs of the processing tasks are combined (reduced) to generate a single result. The input and output data for MapReduce can be stored in HDFS on the same compute nodes used for processing the MapReduce jobs. This produces a very high aggregate bandwidth across the cluster. The user’s applications specify the input/output locations and supply map and reduce functions via implementations of appropriate interfaces and/or abstract-classes. The framework takes care of distributing the software/configuration, scheduling tasks, monitoring the tasks and re-executing any failed tasks.

HDFS links together the file systems on many local nodes to make them into one big file system. HDFS assumes nodes will fail, so it achieves reliability by replicating data across multiple nodes. Processing data in situ on local nodes is efficient compared to
moving the data over the network to a single processing node. This local processing architecture of Hadoop has resulted in very good performance (Rutman, 2011) on cheap computer clusters even with relatively slow network connections (such as 1 Gig Ethernet) (Rutman, 2011). Hence, Hadoop is ideal to underpin our distributed processing architecture.

4.3. Hadoop feature selection

Feature selection is a two part procedure. A training phase described in Section 2.1 trains the data into the CMMs. A test phase then applies test data to the trained CMMs and correlates the results to produce feature selections. Each compute node holds a CMM, CMM stripe or set of CMM stripes that stores all local data. During training, CMMs are not immutable as each association in Eq. (1) changes the underlying CMM so Hadoop MapReduce is not a suitable paradigm for CMM training. Hence, the CMMs are trained in a conventional fashion and uploaded to HDFS once trained. If the data stored in a node’s CMM exceed the memory capacity of that node then the CMM is subdivided into stripes as described in Section 4.1 and shown in Figs. 4 and 5. The set of all CMM stripes at a node stores all data for that node. Every CMM stripe across the distributed system has to be coordinated so that record identifiers (such as timestamps) are matched to allow the CMM sum and threshold. Sum and threshold is column-based and relies on columns representing the same datum. When the results from different CMMs are unified then the columns from the various CMMs need to be aligned. The system is very flexible; we only need to access relevant CMM stripes so we can access subsets of data. The approach is a combination of the stripping described above in Section 4.1 and the CMM distribution described in Section 4.2 with Hadoop orchestrating the search.

While the CMMs are being trained it is expedient to generate a MapReduce input file of input vectors to be used to produce the feature selections. These files will be split into batches by the MapReduce software and the results will be correlated to produce the feature selection scores. There is one input file per CMM stripe and the input vectors in each file represent the set of input vectors for recall to produce the feature selections.

Each CMM stripe that receives a search request, executes the recall process described in Section 2.3. The candidate matches are the set of stored patterns that are close to the query in the feature space. In Hadoop the processing is coordinated by MapReduce (Shvachko et al., 2010). Hadoop YARN schedules the MapReduce tasks independently of the problem being solved. There is one Map job for each input file. Therefore, we model feature selection as a series of MapReduce jobs with each job representing one CMM stripe and the tasks are batches of file iterations (batch processing subsets of records) from the test data. The tasks are processed in parallel on distributed nodes. Each CMM stripe is read into a job. The recall function for CMM stripes is written as a Map task. Each MapReduce job invokes multiple Map tasks, each task represents a batch of recalls for a subset of input records; the batches execute in parallel. The Hadoop Mapper keeps track of the output vector versus record ID pairs so we know which output vector is associated with which record. The Reduce tasks perform the integer output vector thresholding as described in Section 2.3 and write the data back into the file associated with the CMM stripe. Multiple feature selectors can be run in parallel, each executing as a series of MapReduce jobs. The CMMs for feature selection are immutable so subsequent iterations do not depend on the results (or changes) of the CMMs.

This whole MapReduce process has to be coordinated. If the MapReduce process is running at a single location then it can be coordinated as a Java class that initiates the individual jobs and then coordinates the results from all jobs to produce the feature selection scores. If the processing is geographically distributed then it needs a more complete coordinator. This can be achieved using for example the UNIX curl command and a monitor process that determines when curl has collected new data. Alternatively, it can be achieved using a distributed stream processor such as Apache Flume (https://flume.apache.org/) or Storm (https://storm.incubator.apache.org/). Essentially, whichever tool is used this is a three part process: initiate the feature selection process at each of the distributed nodes; retrieve the results data from the distributed nodes; and, monitor when the results have been returned from all nodes and combine them into a single unified result.
4.3.1. Stripe vectors

For Big data, the CMMs are too big to store in a one computer’s memory. Hence, they need to be striped across multiple computers as in Figs. 4 and 5. Each CMM stripe returns a vector representing the matching results for the input vector with respect to that CMM stripe. Palm (2013) has extensively analysed representations in associative memories and found that sparse representations are optimal because the number of matrix operations is proportional to the number of set bits in the vectors. A sparse pattern will have fewest set bits and require fewest operations. For our feature selector, each CMM stripe can return its results as

1. an integer vector \( S_k \) (un-thresholded),
2. a thresholded vector \( T_k \) or
3. a list of the set bits in the thresholded vector.

Option 1 is the least efficient as, potentially, every column could have an integer score so the vector would be an integer vector of length \( N \) where \( N \) is the number of data records stored. This integer vector can be thresholded for option 2 which produces a binary vector. A binary vector requires less storage capacity than an integer vector (1 bit per element for the binary vector compared to 16 or 32 bits per element for the integer vector). For option 3, we would return a list of the set bits. For this we can exploit a compact list representation for representing binary vectors (Hodge & Austin, 2001). This compact list representation is similar to the pointer representation used in associative memories (Benzt, Hagstroem, & Palm, 1997). It ensures that retrieval is proportional to the number of set bits in the thresholded output vector so is fast and scalable. The feature selection process produces a large set of output vectors from the CMM stripes; namely, all vectors necessary for all feature selectors. Option 3 allows AURA to be used for distributed processing with data sets of millions of records while using a relatively small amount of memory and with a massively reduced communication overhead. For example, if there were 10,000,000 records in the data set then a vector would have 10,000,000 elements. If only three records match (records; 8, 10 and 11) then processing (8, 10, 11) as indices requires much less time, memory and communication bandwidth compared to processing 10,000,000 binary digits. Hence, wherever possible we use option 3.

The results need to be amalgamated for each feature selector to produce the feature scores for that feature selector. The system maintains an index of what data are stored where and what each datum represents so the coordinating node can coordinate the matching, receive all matching data and determine the set of best matches across all searchable data. Each feature selector will have a separate amalgamate program running at the coordinating node. This program uses the required vectors and set bit counts returned from AURA to produce the feature score as described in Sections 3 and 5.

5. Analysis of AURA feature selection

We demonstrate theoretically using a worked example that our framework vastly reduces the number of required computations compared to processing the feature selectors separately. The worked example provides an easy and simple illustration of the method on a small data size. We envisage using the feature selector on Big Data sets where Big Data refers to data sets that require at a minimum multiple CPUs but more likely multiple compute nodes to process in tractable time for the application. The larger the data set and the more time critical the data processing then the more important our computation reduction will become. MI, CFS, CS, OR and GR can all use a single CMM representation for the data such as the CMM in Fig. 6. This overall CMM is amenable to striping across the processing nodes to allow Hadoop processing in a similar fashion to Figs. 4 and 5. The framework is underpinned by Hadoop which has been thoroughly evaluated in the literature (Kumar et al., 2013). Hadoop is highly configurable large data set framework that can be optimised to the user’s specific requirements, for example, optimising to minimise memory overhead, optimising for fastest processing or optimising to reduce communication overhead. Hence, we do not attempt to evaluate Hadoop itself here but just focus on how we minimise the number of feature selection computations to minimise processing. Users will use our framework to select the best feature selector for their data and application using their own specific criteria.

The feature selectors in Section 3 have many commonalities when implemented in the unified AURA framework. We can demonstrate the commonalities by analysing 12 records from the Iris data set (Fisher, 1936). The Iris data are illustrated in Fig. 6 (left) when trained into the CMM. The 12 records have been trained into a CMM using the four features and the class. Each feature is quantitative and has been subdivided into five quantisation bins of equal width. Fig. 6 (right) shows the same data divided into four CMM stripes (CMMStripe1, CMMStripe2, CMMStripe3 and CMMStripe4). The horizontal (row-based) striping means that the features “sepal len” and “sepal width” are in the top stripes and “petal len”, “petal width” and the class are in the bottom two stripes. The vertical (column-based) striping means that the first 6 data records are stored in the left two stripes and the other 6 records in the right two stripes. If the data were time-series or sequential, the column-based striping would form two time frames with the oldest data in the left two stripes and the newest data in the right two stripes. The input vectors are stored in a file for each CMM or CMM stripe. These files can then be batch processed in the Hadoop framework described. Within the evaluation, we consider how the data and CMMs would be accommodated in our Hadoop framework.

MI, CFS, CS, OR and GR all use \( BV_f \) (the binary vector where \( F_i = f_i \)), \( BV_b \) (the binary vector representing the quantisation bin \( bin(f_i) \)) and \( BV_c \) (the binary vector representing all records that have class label \( C \)). These only need to be extracted once and used in each feature selector as appropriate. For example in Fig. 6, if we want all records where \( 1.12 \leq petal width \leq 1.58 \) then we activate row 17 of the CMM. We can then Willshaw threshold the resultant integer output vector \( S \) (0000111110000) at level 1 and retrieve the binary thresholded vector \( T \) with a bit set for every matching record (bits 4, 5, 6, 7). For the Hadoop distributed version, only the relevant CMM stripes are queried in Fig. 6 (right). In this case, activating row 17 of CMMStripe3 and CMMStripe4 queries the relevant data. CMMStripe3 will output thresholded vector \( T_3 \) with bits 5 and 6 set and CMMStripe4 will output \( T_4 \) with bits 7 and 8 set. \( T_3 \) and \( T_4 \) can be concatenated to form a single vector thresholded vector \( T \) (as in Fig. 4) with bits 4, 5, 6 and 7 set. For the Hadoop distributed version, each CMM stripe CMMStripeX outputs a list of the indices of the set bits in \( T \) which are collected by the coordinator.

CFS, GR and MI all require \( nBV_f \) a count of the number of data records where a particular feature has a particular value \( f_i = f \) and \( BV_c \) a count of the number of records where the class has a particular label \( C = c \). To count the number of records where \( 1.12 \leq petal width \leq 1.58 \), we retrieve the binary thresholded vector as above and count the number of set bits (bits 4, 5, 6 and 7 are set giving 4 matching records). For the Hadoop approach, we coordinate the retrieval as above, concatenate the lists to produce a single overall list of set bits and count the list length. \( T_3 \) has bits 4 and 5 set and \( T_4 \) has bits 6 and 7 set giving 4 matching records in total.

CFS, CS, OR, GR and MI all use \( BV_f \wedge BV_b \) and \( BV_f \wedge BV_c \) for qualitative and quantitative features respectively. For example, we can find all records where \( 4.6 \leq sepal len \leq 5.1 \) and the
class is $A$ by activating rows 0 and 20 of the CMM, thresholding $S (1222000000)$ at Willshaw level 2 to give $T$ with three bits set: column 1, 2 and 3 in Fig. 6 (left). This takes more coordinating in the Hadoop framework as the data for the feature value may not be stored with the data for the class; they may be in different CMM stripes. In Fig. 6 (right), activate row 0 in $\text{CMMStripe1}$ and $\text{CMMStripe2}$ and then activate row 20 in $\text{CMMStripe3}$ and $\text{CMMStripe4}$. The coordinating program needs to correlate the sections of the vector for the feature value and correlate the sections of the vector for the class to form a single vector. $\text{CMMStripe1}$ needs to be added (summed) with the output integer vector of $\text{CMMStripe3}$ to give $S_1 + 3$ and $\text{CMMStripe2}$ needs to be added (summed) with the output integer vector of $\text{CMMStripe4}$ to give $S_2 + 4$. The summed vectors can then be thresholded at 2 to give $T_{1,3}$ with bits 1, 2 and 3 set (three matching records) and $T_{2,4}$ with no bits set (no matches). The two thresholded output vectors are concatenated to produce $T$ with bits 1, 2 and 3 set. This is simply a task of finding the common indices between the two vectors.

MI, CFS, CS, OR and GR all also need a count of the conjunction, that is $n(BVF_j \land BVc)$ and $n(BVb_i \land BVc)$ for qualitative and quantitative features respectively. Hence, we retrieve the binary thresholded vector $T$ as above and count the set bits. Rather than calculating these elements multiple times, we can take advantage of the commonalities by calculating each common value, binary vector or count only once and propagating the result to each feature selector that requires it. Following these common calculations, all necessary calculations will have been made for MI and GR. CFS just requires the pairwise feature versus feature analyses ($BVb_i \land BVb_k$). These are performed in the same way as the feature versus class analyses above. CS and OR require the manipulation of some of the binary vectors to produce the logical or vectors. This requires the coordination of the vectors. To find $(BVb_j) \lor (BVc)$, we combine the list of set bits for $(BVb_j)$ with the list of set bits for $(BVc)$ and count the resulting list length. By calculating the common elements first, the remainder of the calculations can be performed for each feature selector using either this CMM and processing the algorithms in series or by generating multiple copies of the CMM and processing them in parallel if sufficient processing capacity is available.

Once all of the binary vectors have been retrieved by the distributed Hadoop system, they need to be processed to calculate the feature scores as per Section 3 using the various feature selectors. A coordinator program organises this in parallel. There is one feature score calculation process per feature selector (currently five feature selectors are described here).

For the Iris data set, there are 20 feature row activations $20 \ast BVb_i$ and three class activations $3 \ast BVc$. To calculate $(BVb_i \land BVc)$ requires $20 \times 3 = 60$ calculations. Hence, there are 83 common calculations $(20 + 3 + 60)$ across all five feature selectors. CFS then needs to calculate $(BVb_i \land BVb_k)$ which would require 191 calculations if every feature value was compared to every other. However, CFS uses greedy forward search so that the number of comparisons is minimised (Hall, 1998) to a worst case of $(20^2 - 20)/2 = 190$. We have already extracted all $20 \ast BVb_i$ binary vectors so CFS needs 190 logical ands but no CMM accesses. We have saved a minimum of 20 CMM accesses for $BVb_i$ and a maximum of 190 CMM accesses for worst case forward search. Manipulating the binary vectors can be performed at the
coordinating node and in parallel as a Hadoop batch process. CS requires the logical or vectors $(BV_b \lor BV_c)$. Again, we already have all $20 \times BV_b$ binary vectors and all $3 \times BV_c$ binary vectors so there are $20 \times 3 = 60$ logical ors to perform. Thus, we have saved a minimum of $20 \times BV_b + 3 \times BV_c = 23$ CMM accesses and potentially $60$ CMM accesses if all 60 or operations were performed in the CMM. Thus MI requires 83 calculations, GR also requires 83, CFS requires 83 plus 190 and CS requires 83 plus 60. Without our reductions there would be $83 + 83 + 190 + 83 + 60$ calculations. We have reduced this to $83 + 190 + 60$. Additionally, $190 + 60$ of these can use vectors already extracted so there is no need to access the CMM. We have saved $3 \times 83 = 249$ recalls from the CMM by finding common aspects, have removed a minimum of $20 + 23$ further CMM recalls and have reduced the other calculations to logical operations on stored binary vectors. The minimum saving on CMM recalls is given by Eq. (16).

$$\text{Saving} = (3 \times (n \times BV_b) + n \times BV_c) + (n \times BV_b + n \times BV_c)) + ((2 \times n \times BV_b) + n \times BV_c). \tag{16}$$

6. Conclusion

Massive and complex data sources pose challenges for data mining but they also hold many opportunities. New information can be uncovered, vast timelines of data are available for analysis and the data models learned will be increasingly rich as the training data expands. How the data is represented needs to be carefully considered including careful preparation such as cleaning and selecting feature subsets. In this paper we have introduced a distributed processing framework for feature selection using the AURA neural network and Apache Hadoop. There are currently five feature selectors available which may be used independently or coupled with the AURA k-NN for classification or prediction.

All five feature selectors can use a single trained CMM. We have identified common aspects of the five feature selectors when they are implemented in the AURA framework and indicated how these common aspects may be processed as a common block. All remaining aspects of the feature selectors can then be implemented in parallel using duplicate copies of the trained CMM as compute resources allow. CMMs lend themselves to distributed processing as they can be striped (split) using both row-based and column-based striping. The CMM created for feature selection can be used directly for the AURA k-NN for classification or prediction and any unwanted features (those not selected by the feature selection) can simply be ignored (masked off). Alternatively, the CMM can be retrained with only the required data if processing speed and memory usage at recall time are the primary concern.

The AURA neural architecture has demonstrated superior training and recall speed compared to conventional indexing approaches such as hashing or inverted file lists (Hodge & Austin, 2001) and an AURA-based implementation of the MI feature selector was over 100 times faster than a standard implementation (Hodge et al., 2006). This is further augmented by using the scalability of Hadoop. This combined platform allows rapid processing of feature selectors on large and high dimensional data sets that cannot be processed on standard computers. We envisage using the method on data sets that require at a minimum multiple CPUs but more likely multiple compute nodes to process. The method is also best suited to data mining and analytics that processes a Big Data file in a longer term processing run such as overnight rather than on-line transaction processing which requires near real-time updating. The user can then evaluate the feature sets chosen by the feature selectors against their own data to determine the best feature selector and the best set of features. Additionally, each feature selector (MI, CFS, GR, CS and OR) generates scores for the features which can be used to weight the features during machine learning.

The technique is flexible and easily extended to other feature selection algorithms. By implementing a range of feature selectors in a single framework, we can also investigate ensemble feature selection where the results from a range of feature selectors are merged to generate a consensus overview of the best set of features to use.

We will investigate whether we can use Apache Spark, the in-memory data analytics and cluster computing framework (https://spark.apache.org/) to underpin the AURA feature selection framework. Apache Spark is closely coupled with Hadoop and allows YARN and MapReduce jobs to be run. Spark enables in-memory computing and is reputed to be up to 100 times faster than MapReduce (see https://spark.apache.org/). CMMs are optimised for in-memory processing so fit well with the Spark paradigm. A related development, Optimised RowColumnar (ORC) file format is currently being adopted by Spark. ORC is a file storage format that is tightly integrated with HDFS and provides optimisations for both read performance and data compression. An ORC file divides the data into groups of row data called stripes. This fits with the stripes used in AURA CMMs and would allow a direct mapping of ORC data file stripes to CMM stripes for optimised performance.

We plan to use the feature selection framework that we have developed in this paper in conjunction with the AURA k-NN for traffic analysis (Hodge et al., 2012; Hodge, Krishnan, Austin, & Polak, 2010; Hodge et al., 2011), condition monitoring (Austin et al., 2010) and railway infrastructure monitoring in the NEWTON Project (Hodge, O’Keefe, Weeks, & Moulds, 2015).

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