Multithreaded and Spark parallelization of feature selection filters

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A B S T R A C T

Vast amounts of data are generated every day, constituting a volume that is challenging to analyze. Techniques such as feature selection are advisable when tackling large datasets. Among the tools that provide this functionality, Weka is one of the most popular ones, although the implementations it provides struggle when processing large datasets, requiring excessive times to be practical. Parallel processing can help alleviate this problem, effectively allowing users to work with Big Data. The computational power of multicore machines can be harnessed by using multithreading and distributed programming, effectively helping to tackle larger problems. Both these techniques can dramatically speed up the feature selection process allowing users to work with larger datasets. The reimplementation of four popular feature selection algorithms included in Weka is the focus of this work. Multithreaded implementations previously not included in Weka as well as parallel Spark implementations were developed for each algorithm. Experimental results obtained from tests on real-world datasets show that the new versions offer significant reductions in processing times.

1. Introduction

Massive amounts of data are generated on a daily basis nowadays. As an example, IBM estimated that in the time span between 2010 and 2012, 90% of the data worldwide were produced at a rate of 2.5 exabytes per day [1]. This data-generating trend has sparked interest in data analytics, which in turn has created the need for new tools, algorithms and methodologies that can cope efficiently with such massive amounts of data. Consequently, the dimensionality (samples × features) of datasets being analyzed in machine learning problems has been steadily growing in the last years. Taking the datasets posted in the popular libSVM Database [2] as a reference, their size has increased five hundredfold. This results in a challenge for traditional machine learning algorithms, as overfitting can negatively impact their performance, more complex models are harder to interpret and both speed and efficiency of these algorithms decrease as the dimensionality increases.

This situation has spawned a number of techniques designed to deal with big dimensionality datasets. This dimensionality can refer to samples, features or both. In the case in which we confront with datasets containing numerous features, feature selection techniques are mandatory. Feature selection consists in the process of determining the relevant features and trying to remove as much irrelevant and redundant information as possible, without leading to a degradation in the classification performance.

The Weka (Waikato Environment for Knowledge Analysis) suite [3] is a very popular machine learning platform that has been downloaded over six million times. It can be used as a stand-alone application or imported as a library from the user’s code. Feature selection is among its functionalities with several algorithms available to the user. This ample range of algorithms included in Weka makes its use widespread among data scientists for data analysis and for the development and testing of new algorithms. In addition, the fact that Weka runs on Java and is designed with single-machine setups in mind, makes it very suitable for the average user. Nevertheless, some of the implementations in Weka still struggle when processing large datasets, requiring very long execution times, effectively limiting the size of the datasets that can be analyzed with it. An improvement in the time efficiency of these algorithms will enable its many users to process large datasets that up to now were out of reach for these implementations.

On the other hand, and to specifically address the Big Data issue, new parallel programming solutions have been created in the last decade, such as MapReduce [4], that was implemented in the open-source solution Apache Hadoop [5] or, more recently, Apache Spark [6], which aimed at being a solution to Big Data analysis. The advent
of these new technologies led to the creation of parallel machine learning libraries: Mahout [7] that runs on top of Apache Hadoop, and MLLib [8] that uses Apache Spark, are some examples. Although these libraries contain a wide variety of machine learning algorithms, they do not provide many options when it comes to feature selection.

Spark is designed for distributed computing and can achieve great performance processing large amounts of data, but few implementations of feature selection algorithms are available. Moreover, to be able to use the Mahout or MLLib libraries, the user needs to have a Hadoop or Spark installation and, although they can run on single-machine environments, a cluster of computers would be needed to fully exploit these libraries, which is not always available for regular users.

The aim of this work is to obtain new implementations of these popular feature selection algorithms that are able to tackle sizable problems in different environments and find out the suitability of these implementations to different amounts of computing resources1. To this end, multithreaded implementations for Weka and distributed versions in Spark will be proposed. This will allow users to analyze larger datasets in shorter times and choose the most adequate implementation to the resources available to them.

This paper is organized as follows: Sections 2 and 3 are an overview of feature selection and parallelization approaches respectively. Section 4 describes the algorithms that are the object of this paper. The results of our tests are presented in Section 5 and in Section 6 we discuss our conclusions.

2. Feature selection

Feature selection is the name given to the process that detects relevant features and discards those that are redundant or irrelevant. The goal of this technique is to obtain a subset of features that has minimum degradation of performance when used by a classifier while describing the given problem properly. It simplifies the dataset both in size and in complexity of understanding [9], which leads to simpler and faster classification algorithms, better problem comprehension and reduced storage requirements.

2.1. Feature selection methods

Feature selection methods can be classified into two categories: individual evaluators or subset evaluators. Individual evaluators are also called rankers and they assign a weight to each attribute that represents its relevance. Subset evaluators, on the contrary, employ a search strategy to determine a candidate subset of features and have the advantage of removing redundant attributes at the cost of being more complex. According to the relationship with the learning method used, feature selection methods can also be divided as follows [10]:

- Filters are methods that are applied independently of the induction process. They are, in general, computationally inexpensive.
- Wrappers use the induction algorithm as a black box to evaluate the fitness of each candidate subset. This results in algorithms that are computationally demanding but more accurate.
- Embedded methods perform feature selection in the process of training and are typically specific to given learning algorithms.

In this paper, three of the most commonly used filter methods (InfoGain, RELIEF-F and CFS) and an embedded method (SVM-RFE) were selected for reimplementation using a parallel approach. The first two filters are rankers that return an order for the features to be discarded below a threshold of the user’s choice and they are included in the Weka suite:

- Information Gain (InfoGain) [11] is a filter that computes the mutual information of the different features with respect to the class and provides an ordered ranking of all the features according to this value.
- RELIEF-F [12] is a heuristic estimator built upon the RELIEF algorithm [13] that deals efficiently with noisy and incomplete datasets and with multiclass problems. It works by locating its nearest neighbors for each instance from the same and opposite class and updating the weights of each feature accordingly.

The remaining two algorithms are subset evaluators. To perform feature selection they search through the space of all possible attribute combinations for the set that offers a better score according to a heuristic method that depends on the algorithm.

- CFS [14] is a subset evaluator independent from the induction process that tries to identify correlations between attributes and the class.
- SVM-RFE [15], which stands for Support Vector Machine Recursive Feature Elimination is an embedded method that filters the attributes iteratively using a SVM at each stage to rank them.

The choice of these algorithms was made to obtain a set of tools that are well suited to a wide range of datasets. CFS and InfoGain perform well when the data has a large number of attributes when compared to the number of instances and are very fast, but they do not perform as well when there is noise in the inputs. RELIEF-F is very good at eliminating redundant and correlated features, even when there is noise in the inputs and attributes are non-linear, but it is much slower and does not perform well when few examples are available. Lastly, SVM-RFE detects correlation and redundancy even with few examples, but it performs poorly when there is noise in the inputs and is very time consuming [9].

Table 1 shows the theoretical complexity of the four methods described above.

3. Parallel approaches

The main purpose of this work is to parallelize the standard implementations of RELIEF-F, InfoGain, CFS and SVM-RFE. In order to empower Weka users, multithreaded implementations are proposed. Furthermore, to enable users that can access computational clusters, we developed and tested Spark versions of the algorithms.

3.1. Multithreaded processing

Multithreading allows users to take advantage of multicore systems without imposing the overhead of creating multiple processes and providing direct access to a common address space. However the creation and management of threads introduces a computational overhead that makes the use of threads suboptimal when the tasks parallelized have low complexity.

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Table 1: Theoretical complexity of the four feature selection methods focus of this work (where \( m \) is the number of examples and \( n \) is the number of attributes).

<table>
<thead>
<tr>
<th>Method</th>
<th>Complexity</th>
</tr>
</thead>
<tbody>
<tr>
<td>InfoGain</td>
<td>( nm )</td>
</tr>
<tr>
<td>RELIEF-F</td>
<td>( n^2m )</td>
</tr>
<tr>
<td>CFS</td>
<td>( n^2m )</td>
</tr>
<tr>
<td>SVM-RFE</td>
<td>( \max(n,m)nm^2 )</td>
</tr>
</tbody>
</table>
Java provides parallel programming support in the core of the language. This feature enables programmers to write code that exploits multithreading without the need to use any external library. Since Weka is written in Java, we use this support to implement our multithreaded parallel codes. We divide the feature selection algorithms in tasks that can be performed in parallel, which allows us to exploit the computational power of multicore machines.

### 3.2. Parallelization with Apache Spark

To alleviate the difficulties of developing distributed programs, a team of Google engineers developed the MapReduce framework [4] that handles the common aspects of distributed programs, providing the programmer with a tool to run parallel programs without having to worry about anything but the implementation of the algorithm.

The programming paradigm introduced by MapReduce requires the tasks to be divided into two separate steps: the Map phase, that applies a function given by the user to every element; and the Reduce phase, that combines the resulting values. Often times elements consist of key-value pairs and the Reduce phase merges results that have the same key, although this is not mandatory. The abstraction resulting of decomposing a job in simple Map and Reduce functions allows the framework to divide both data and code across the computing nodes, a task performed by a master node. Typically, the framework splits the data in as many chunks as nodes are available, distributes it among them so that each node can apply the Map function to the assigned elements. The results are then rearranged by the master node, using a key partitioning scheme, and distributed again back to the nodes so that they perform the Reduce phase.

MapReduce was implemented in the open-source framework Hadoop [5] and rapidly achieved great popularity for its reliability and scalability. Still, this direct implementation left room for an important improvement that was later implemented by the Spark [6] framework: the transition between the Map and Reduce phase requires data to be shuffled by the master node and redistributed to the nodes, in a time-consuming process that is unnecessary when several Map transformations need to be applied before the Reduce phase or in iterative algorithms. By avoiding unneeded data movement and introducing other optimizations Spark performs several times faster than Hadoop for certain applications [16].

Spark allows the programmer to manage work distribution by means of using Resilient Distributed Datasets (RDDs), an abstraction that represents a read-only set of objects that is distributed across multiple machines. RDDs can be transformed, performing an operation on each element, which can be done in parallel in each node, and they can be reduced, combining elements, which requires that the whole dataset is shuffled and redistributed to the nodes in a time-consuming process. Additionally, data can be sent to the nodes to work with by using broadcast variables, and the worker nodes can write increments to special variables named accumulators.

We decomposed the feature selection algorithms in independent tasks for a Spark implementation that will allow the user to take advantage of a computer cluster to process large datasets in reduced time.

### 4. Implemented algorithms

Four algorithms (listed in Table 2) were the object of this work. Of the 8 possible implementations (a Weka multithreaded and a Spark version for each algorithm), 2 were already available and 6 were developed as part of this work.

#### 4.1. RELIEF-F algorithm

The original RELIEF-F algorithm [12] loops through a set of instances R finding for each instance its k nearest neighbors from the same class, called nearest hits H, and the k nearest neighbors from each different class, which are denoted as nearest misses M(C). When all neighbors are found, the weight for each attribute W[A] is updated by subtracting the weighted average distance (computed with the diff function, that returns the Manhattan distance between two instances) of each hit H and adding, for each class C other than R's, the weighted average of the distance to each miss M(C). When computing averages, distances are weighted by the probability P of the class and divided by the total number of instances m.

Regarding the multithreaded implementation, the job was divided into as many tasks as threads we wanted to use, then a thread was created for each task. This approach avoided the need for a thread poole to manage the execution of threads. This process is detailed in Algorithm 1.

The process of finding the nearest neighbors for each instance (by means of the loop described between Lines 2 and 7 of Algorithm 1) is very time consuming since it requires comparing it with all other instances. This search can be executed independently for each instance and therefore it can be performed in parallel with no synchronization issues.

**Algorithm 1.** Pseudo-code for multithreaded RELIEF-F

| Input: R ← Set of instances having a set of attributes A classified in classes from a set C |
| Output: W ← vector storing the weight of each attribute |
| 1 set all weights W[A] ← 0.0 |
| 2 for i ← 1 to THREADS.AVAILABLE do in parallel |
| 3 Si ← disjoint subset of instances |
| 4 foreach I in Si do |
| 5 find k nearest hits H |
| 6 foreach class c ∈ C / c ≠ class(I) do |
| 7 from class c find k nearest misses M(c) |
| 8 endforeach |
| 9 foreach a in A do |
| 10 foreach R in R do |
| 11 W[a] ← W[a] − \( \sum_{j=1}^{k} \frac{diff(a, R_j, H_j) \cdot P(c)}{m+k} \) + \( \sum_{c \neq class(R_i)} \frac{P(c)}{1 - P(class(R_i))} \sum_{j=1}^{k} \frac{diff(a, R_j, M(c))}{m+k} \) |
| 12 end |
| 13 end |
| 14 end |

### Table 2

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>RELIEF-F</td>
<td>Time-efficient feature selection algorithm.</td>
</tr>
<tr>
<td>RELIEF+</td>
<td>An extension of RELIEF-F that considers the distances to the k nearest neighbors of each class.</td>
</tr>
<tr>
<td>CRELIEF</td>
<td>An extension of RELIEF-F that considers the distances to the k nearest neighbors of each class.</td>
</tr>
</tbody>
</table>

*Note: This table is a simplified representation and does not include all possible implementations.*
In our Spark implementation the work is split in the same way: each node computes the nearest neighbors to a subset of the examples. Every possible pairing of example indices is generated and stored in a Spark RDD, which is then distributed to the nodes. The whole dataset is sent to the nodes as a broadcast variable, so that they use it as a lookup table. This approach obtains a considerable speed gain, but effectively limits the size of the dataset to the maximum size a Spark broadcast variable can handle.

4.2. InfoGain algorithm

The InfoGain algorithm assigns the weight ($W$) of each attribute ($A$) by contrasting its information gain with respect to the class. To calculate this value, the entropy ($H$) of each class given the attribute in question is subtracted from the entropy of that class:

$$\text{InfoGain}(\text{Class}, \text{Attribute}) = H(\text{Class}) - H(\text{Class} | \text{Attribute})$$  \hspace{1cm} (1)

Entropy of a variable is defined as $−\sum p(i) \log(p(i))$, where $i$ loops through every possible value of the variable. The observed probability of a variable taking a value is represented by $p(i)$, and it is calculated as the ratio of cases where the variable takes that value divided by the total number of appearances of the given variable. If the variables are not discrete, the dataset needs to be preprocessed as described in further detail in Section 5.1.

Weka implements this calculation by looping through all the dataset counting the number of appearances of every possible value for each attribute, storing the counts in an array. Then this array is used to compute the information gain of each attribute. This process has linear complexity.

In our proposed multithreaded solution, detailed in Algorithm 2, the counting of every possible value is performed for a subset of the samples (Line 4). This requires an additional step, described in Line 7, that combines the counts of each thread into a global count. Since this division is performed on the number of instances, it will be more effective when the dataset has numerous instances. For small datasets, the additional accumulative step can take more time than the one that is gained from counting in parallel, but for large datasets the time required to add up the partial counts should be negligible when compared to the counting process. Lastly, the process of obtaining the information gain values from the counts can also be performed independently for each attribute, therefore it can be computed in parallel (Line 11). The functions $\text{Entropy}$ and $\text{ConditionalEntropy}$ shown in Line 14 represent the calculation of $H(\text{Class})$ and $H(\text{Class} | \text{Attribute})$ respectively.

```
Input: R ← Set of instances having a set of attributes A and classified in classes from set C
Output: W ← vector storing the weight of each attribute
1 set all counts ← 0.0
2 for t ← 1 to THREADS.AVAILABLE do in parallel
3     $S_t$ ← disjoint subset of instances
4     foreach $I$ in $S_t$ do
5         foreach $a$ in A do
6             $\text{count}_a, value_i(a), class_I$ ←
7                 $\text{partial}_\text{counts}_t, a, value_i(a), class_I$ +
8                 $\text{weight}(I)$
9         end
10     end
11 for t ← 1 to THREADS.AVAILABLE do in parallel
12     $A_t$ ← disjoint subset of attributes
13     foreach $a$ in $A_t$ do
14         $W[a] ← \text{Entropy}($\text{counts}_a$) −$
15         $\text{ConditionalEntropy}($\text{counts}_a$)
16     end
end
```

Again, the use of a thread pooler was avoided by creating as many tasks as threads are available.

The InfoGain algorithm is already included in the Spark Infotheoretic Feature Selection package [17] that implements several algorithms that share a common structure by the use of a framework [18]. This was the version tested in this paper.
### 4.3. CFS algorithm

CFS is a subset evaluator that uses the correlation between attributes to obtain a score for a group of attributes. The computational cost for this algorithm is greatly influenced by the need to obtain the matrix that contains the Pearson product–moment correlation coefficients between every possible pair of attributes. The time complexity of this process grows quadratically with the number of attributes and linearly with the amount of samples, which means that most of time of the CFS algorithm is spent in this process. Once the correlation matrix and the standard deviations of each attribute have been computed, CFS searches the space containing every possible attribute subset looking for one that obtains the highest score in its evaluation method.

The search algorithms used can vary in their complexity, from simple greedy algorithms as the one described in Algorithm 3 that simply adds to the set the best candidate at each step, to more complex backtracking ones like BestFirst, listed in Algorithm 4. This search method keeps a list with every candidate set that it encounters ordered by their score in the evaluating function. For each candidate, it explores every possible addition to the set, adding the resulting new set to the candidate list if its score is high enough. This process goes on until the examination of candidate sets renders no new candidates for a given number of iterations (named MAX_STALE in Line 6 of Algorithm 4).

#### Algorithm 3. Greedy stepwise search used in CFS

Input: A ← Set of all possible attributes  
Output: S ← Previously selected attributes  
Input: previous_merit ← Merit of S  
Output: S.out ← Selected attributes  

```plaintext
best_merit ← previous_merit  
best_set ← S  
for a ← A do in parallel 
  set_merit ← computeMerit((S, a))  
  if set_merit > best_merit then  
    best_merit ← set_merit  
    best_set ← (S, a)  
end  
if best_set != S then  
  return greedy_stepwise(A, best_set, best_merit)  
end  
else  
  return best_set  
end  

The evaluation function used by CFS is described in Algorithm 5. It increases when the attributes are highly correlated with the class and it decreases when any attribute is highly correlated with other attributes that are already in the set.

#### Algorithm 4. Best-first search used in CFS

Input: A ← Set of all possible attributes  
Output: S ← Selected attributes  

```plaintext
stale ← 0  
candidates ← OrderedList((empty, 0))  
merit_cache ← empty  
best_set ← empty  
best_merit ← 0  
while candidates.hasNextElements() and stale < MAX_STALE do  
  S, S_Merit ← candidates.popFirst()  
  added ← false  
  for a ← A do in parallel  
    if (S, a) in merit_cache then  
      set_merit ← merit_cache.getMerit((S, a))  
    else  
      set_merit ← computeMerit((S, a))  
      merit_cache.storeMerit((S, a), set_merit)  
    end  
    if set_merit > S_merit then  
      candidates.push((S, a), set_merit)  
      if set_merit > (S, a, set_merit)  
        added ← best_merit then  
          stale ← true  
          stale ← 0  
          best_merit ← set_merit  
          best_set ← (S, a)  
        end  
      end  
    end  
  end  
if not added then  
  stale ← stale + 1  
end  
```
Algorithm 5. Subset evaluation in CFS

Input: A ← Subset of attributes
Input: C ← Matrix containing the correlation between the ith and jth attributes in C[i][j]
Input: SDev ← Array containing the standard deviation for each attribute

Output: M ← Merit of subset

1 numerator ← 0
2 denominator ← 0
3 for a ← A do
4 numerator ← numerator + C[a][class] * SDev[a]
5 denominator ← denominator + SDev[a]^2
6 for b ← A where b < a do
7 denominator ← denominator + 2*SDev[a]*SDev[b]*C[a][b]
end
8 M ← \(\frac{numerator}{denominator}\)

In the existing Weka implementation, which is included by default in the Weka suite, the computation of the correlation matrix is performed in parallel by several threads, although this only occurs when the user chooses to precompute the correlation matrix. Otherwise the matrix is computed in an on-demand basis, which offers better performance.

Our proposed Spark implementation first performs the correlation matrix computation in parallel and then the search process (either BestFirst or GreedyStepwise) is performed, evaluating the different candidate subsets also in parallel.

4.4. SVM-RFE algorithm

To perform feature selection, the SVM Recursive Feature Elimination (SVM-RFE) algorithm makes use of support vector machine classifiers to assign a weight to each attribute. Starting with the whole set of attributes, an SVM is trained to classify binary datasets. The weights assigned to the features by the SVM are then examined and those with the lowest absolute value are removed from the set and added to the ranking in the lowest positions, as shown in Line 16 of Algorithm 6 (the number of elements added at each iteration can be configured with the STEP variable). Then the process is repeated for the remaining attributes until the ranking is complete (Line 12).

In order to work with multiclass datasets, a different ranking is obtained for each class (Line 3) using a one-vs-all approach, that is, assuming that those elements pertaining to a class other than the one being analyzed are negative examples. Then those rankings are combined by looping through them and adding to the final ranking the best of each list, then the second best and so on, in a loop described in Line 5. The process for obtaining the ranking for each class can be done in parallel, and this is the approach taken in our multithreaded Weka implementation. This allows the new version to take much less time when processing multiclass datasets, while not hindering the performance when used with binary datasets.

In the Spark implementation, by contrast, it is the process of training the SVMs that is done in parallel, allowing to save time both on multiclass and binary datasets. This can be done by using the existing SVM with stochastic gradient descent (SGD) implementation in Spark’s MLlib library. SGD is an incremental algorithm that is well suited for parallelization. Weka employs Sequential Minimal Optimization (SMO [19]), an analytical method that is generally faster, but much harder to parallelize. This change in the nature of the SVM training algorithm results in a selected set of features that can be different from that obtained with Weka.

Algorithm 6. Pseudo-code for multithr. SVM-RFE

Input: S ← Set of instances having a set of attributes A classified in classes from a set C

Output: ordered ← Ordered attributes

1 attributeScoresByClass ← empty
2 for c ← C do in parallel
3 attributeScoresByClass[c] ← RankBySVM(c, S)
end
4 ordered ← empty
5 for a ← A do
6 for c ← C do
7 if not ordered.contains(attributeScoresByClass[c][a])
8 ordered.add(attributeScoresByClass[c][a])
end
end
9 return ordered

Function RankBySVM(c, S)
10 numAttrs ← Number of attributes
11 ranking ← empty stack
12 while numAttrs > 0 do
13 weights ← new SVMClassifier(S,c).weights
14 for w in weights do
15 weights[w] = weights[w]^2
end
16 for i in 0 to STEP do
17 worstAttr ← findWorst(weights)
18 S.removeAttr(worstAttr)
19 ranking.add(worstAttr)
end
20 return ranking
5. Experimental results

The goal of this work is to take advantage of multithreaded and distributed processing to speed up feature selection. Hence, the features selected and the weights assigned by the new versions of the algorithms are the same as those obtained with the original versions, excluding any differences that may arise due to rounding or numeric processing (except in the case of SVM-RFE that obtains different results in Spark due to the change of the nature of the underlying SVM). Consequently, these new versions do not modify the classification accuracy, but aim at being able to perform feature selection in a reasonable, shorter time.

It is worth mentioning that the time complexity of the studied algorithms is very variable. Furthermore, their impact on the total time needed for the whole feature selection process can also be significantly different. Since one of the goals of this work is to provide a reference guide to help users select one implementation, we have decided to list the total execution time instead of just the time invested in the part of the algorithm that actually performs feature selection because this will give users a more accurate idea of what to expect from a certain implementation. There may be some use cases where the algorithm is used in a different context (for instance, loading a dataset once and then performing several iterations of a feature selection algorithm), that take more advantage from the gain associated with the parallel implementation. Nonetheless, the most common use case is performing feature selection on a dataset contained in a file.

In order to provide a variety of scenarios to test the proposed Weka and Spark implementations, seven high dimensional datasets were chosen (see their characteristics in Table 3). We used the Higgs dataset, which consists of 11,000,000 instances with 28 numerical attributes that represent kinematic properties of particles detected in an accelerator [20]. The second dataset used, from here on called Epsilon, was artificially created in 2008 for the Pascal Large Scale Learning Challenge [21]. A preprocessed version available on the LibSVM dataset repository [22] was used. This dataset consists of 500,000 instances that have 2000 numerical features each. Since both datasets mentioned above are binary datasets, one additional dataset with several classes was selected, KDD99 [23]. It contains close to 5 million samples of 41 connection parameters each that are categorized in 23 different classes. Also, SVMs require that datasets have numeric attributes only, so any non-numeric attribute needs to be transformed. Therefore, three multiclass datasets with numeric features were chosen: Isolet [24] consists of almost 8000 instances with 617 attributes each, divided in 27 classes. USPS [25] is a dataset containing over 7000 examples of elements with 256 attributes, representing handwritten characters, with 10 different labels. Lastly, the Poker dataset contains over a million elements with 10 features each, classified in 10 different classes, representing possible hands in the poker card game.

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Dataset description.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dataset</td>
<td>Features</td>
</tr>
<tr>
<td>Higgs</td>
<td>28</td>
</tr>
<tr>
<td>Epsilon</td>
<td>2000</td>
</tr>
<tr>
<td>KDD99</td>
<td>41</td>
</tr>
<tr>
<td>Isolet</td>
<td>617</td>
</tr>
<tr>
<td>USPS</td>
<td>256</td>
</tr>
<tr>
<td>Poker</td>
<td>10</td>
</tr>
<tr>
<td>KDDDB</td>
<td>29,890,095</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 4</th>
<th>Computer cluster description.</th>
</tr>
</thead>
<tbody>
<tr>
<td>16 nodes consisting of:</td>
<td></td>
</tr>
<tr>
<td>Processor:</td>
<td>2 × Intel Xeon E5-2660 Sandy Bridge-EP at 2.20Ghz</td>
</tr>
<tr>
<td>Cores:</td>
<td>8 per processor (16 per node)</td>
</tr>
<tr>
<td>Threads:</td>
<td>2 per core (total of 32 threads per node)</td>
</tr>
<tr>
<td>Hard drive:</td>
<td>1 × SSD 480GB SATA3</td>
</tr>
<tr>
<td>RAM:</td>
<td>64 GB DDR3 1600 Mhz</td>
</tr>
<tr>
<td>Network:</td>
<td>InfiniBand FDR &amp; Gigabit Ethernet</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 5</th>
<th>Execution times of the discretization algorithm implementations.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Runtime (s)</td>
<td>1 core</td>
</tr>
<tr>
<td>Higgs</td>
<td>1585</td>
</tr>
<tr>
<td>KDD99</td>
<td>316</td>
</tr>
<tr>
<td>Epsilon</td>
<td>1976</td>
</tr>
</tbody>
</table>

An additional larger dataset named KDDDB consisting of 19 million samples with 30 million attributes was included as an example of very high dimensionality [26].

The experiments were run on up to 8 nodes of a computer cluster. Each node has the specifications described in Table 4. The Weka version used was 3.7.12 running on OpenJDK 1.7.0_55. The OS installed in this machine was Rocks 6.1, based on CentOS 6.x. Spark applications were run using the MapReduce Executable (MREv) tool, that unifies the configuration of various distributed computing environments [27].

To measure the performance of the new versions of the algorithms comparatively to the original implementations we used the speed-up measure, defined as the ratio between the original sequential time and the parallel one.

5.1. On the preprocessing of the datasets: parallelization of a discretization algorithm

Some feature selection algorithms, such as InfoGain, require the attributes of the dataset to be discrete. This specification often forces the user to preprocess the dataset in order to obtain a modified version with discrete features. Weka provides an implementation of the Fayyad–Irani Minimum Descriptive Length (MDL) algorithm [28] that fulfills that purpose, although this process can be very time consuming. The goal of this algorithm is to transform real-valued attributes to discrete ones while maintaining as much information as possible. To achieve this, real values need to be assigned to different bins that cover the whole range of values of the attribute. The size, number, and distribution of the bins is decided by the algorithm in a long process that is performed independently for each attribute. This allows us to obtain better performance by using separate threads to compute different attributes, as described in Algorithm 7. A similar parallelization with Spark has not been addressed in this section as it was already available in Spark packages [29]. Table 5 shows the execution times for the sequential implementation compared to the multithreaded one when run on a 16 core machine using the three more general datasets (with and without numerical features, as explained at the beginning of this section).
**Algorithm 7.** Fayyad-Irani discretization

```
Input: A ← List of attributes
Input: S ← dataset
1 for a ∈ A do in parallel
2   orderedS ← S.orderBy(a)
3   bins[a] ← computeCutPoints(a)
   // computeCutPoints uses mutual
   information to obtain the bins in
   which to discretize the values for
   attribute a.
end
4 for i ← 1 to THREADS.AVAILABLE do in
   parallel
5   Si ← disjoint subset of instances
6   foreach I in Si do
7     for a ← A do
8       Si[a] ← bins[a].transform(Si[a])
end
end
```

Although the computing process is independent for each thread, a separate copy of the dataset needs to be allocated for each task, since its first step is to order it by the attribute being examined. The overhead created by copying the dataset can be quite large if the dataset is sizable, but in most cases it is not as large as the gain obtained by computing in parallel. In our experiments all datasets but one obtained a favorable speed-up, independently of their size. The new version performed worse than the sequential one for the Higgs dataset, due to its large size and few attributes, which amounts to costly copies of the dataset and less parallelism.

5.2. Analysis of the RELIEF-F implementations

The good adaptability of RELIEF-F to a parallel environment (which is often referred to as being “embarrassingly parallel”) translates into significant decreases in terms of execution time. Despite this improvement, RELIEF-F’s complexity grows quadratically with the number of samples and linearly with the number of features and this still makes it yield long times when the number of instances of the dataset is very high. However, our multithreaded implementation can take advantage of machines with a large number of cores, decreasing computational times.

In order to be able to make a comparison with the sequential version, we have used reduced versions of the largest general datasets (with numerical and non-numerical features) when analyzing the RELIEF-F implementation. For the Epsilon and the KDD99 datasets the top 10% of the instances were used, amounting to a total of 50,000 and almost 500,000 instances, respectively. The Higgs dataset had to be further trimmed, using the top 4%, consisting of 440,000 instances.

We performed tests with different number of threads processing the same datasets in order to illustrate the relation between the execution time and the number of threads employed. The results of these experiments are shown in Fig. 1. The node used to run the benchmarks offered 16 cores, each one capable of running two threads using HyperThreading. When 16 threads are used, they are mapped to different cores with exclusive use of resources, obtaining maximum performance. On the contrary, when we request the use of 32 threads, they are placed two on each core, competing for the core resources [30]. This results in a degradation of performance that, in our best case, barely improves on the use of 16 threads. Therefore, all subsequent experiments were made using just the 16 cores.

The left part of Table 6 lists the execution times of sequential and multithreaded Weka implementations. The multithreaded version was executed using the 16 cores available. A significant performance increase exists for all datasets. When the dataset being analyzed is large, the time taken to manage threads becomes irrelevant in comparison to the time gained by making computations in parallel. The multithreaded version of the algorithm was able to process the large datasets between 12.6 and 16.7 times faster than the sequential one. The good adaptability of this algorithm to a parallel paradigm reflects in the superlinearity of the speed-up obtained for the Higgs dataset.

For comparison purposes, Table 6 also shows the Spark execution times for different amount of cores. The Epsilon dataset was chosen for this comparison since its execution time was high on Weka and its size was suitable for the Spark implementation. As discussed in Section 4.1, the Spark implementation of RELIEF-F requires that the entire dataset is broadcast to all nodes. Good scalability is observed when more nodes are added and, even with one node (16 cores), the Spark implementation is more efficient than the Weka one. To assess the advantage of using Spark and a computer cluster vs Weka on a single machine, the speed-up shown as † is the best Spark result vs the multithreaded Weka result.

5.3. Analysis of the InfoGain implementations

The Weka implementation of the InfoGain feature selection algorithm requires the attributes to be discrete, so it performs a discretization process when needed before the feature selection is started. This discretization is independent from the InfoGain algorithm so, to eliminate its impact in the execution time and obtain a more accurate comparison of the two versions of the algorithm, all datasets used to test the InfoGain feature selector were discretized beforehand using the same algorithm employed by Weka [28]. This resulted in datasets that, in some cases, had several attributes with constant value. Additionally, to speed up this process for users, a multithreaded implementation of this algorithm is provided, as described in Section 5.1.
Table 6
Execution times of RELIEF-F implementations.a

<table>
<thead>
<tr>
<th># cores</th>
<th>Weka</th>
<th>Spark</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>16</td>
<td>32</td>
</tr>
<tr>
<td>Higgs</td>
<td>105,443</td>
<td>6328</td>
</tr>
<tr>
<td>KDD99 (10%)</td>
<td>154,305</td>
<td>10,517</td>
</tr>
<tr>
<td>Epsilon (10%)</td>
<td>84,149</td>
<td>6678</td>
</tr>
</tbody>
</table>

a Speed-ups listed are 16 cores vs 1 core for Weka and 128 cores vs 16 cores for Spark. † indicates the speed-up for 128 cores using Spark vs 16 cores using Weka, that is, the gain of the parallel approach.

Table 7
Execution times of InfoGain implementations.a

<table>
<thead>
<tr>
<th># cores</th>
<th>Weka</th>
<th>Spark</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>16</td>
<td>32</td>
</tr>
<tr>
<td>Higgs</td>
<td>204</td>
<td>192</td>
</tr>
<tr>
<td>Epsilon</td>
<td>458</td>
<td>424</td>
</tr>
<tr>
<td>KDD99</td>
<td>145</td>
<td>140</td>
</tr>
<tr>
<td>KDDB</td>
<td>200</td>
<td>192</td>
</tr>
</tbody>
</table>

a Speed-ups listed are 16 cores vs 1 core for Weka and 128 cores vs 16 cores for Spark. † indicates the speed-up for 128 cores using Spark vs 16 cores using Weka, that is, the gain of the parallel approach.

Table 8
Execution times of CFS implementations.a

<table>
<thead>
<tr>
<th># cores</th>
<th>Weka</th>
<th>Spark</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>16</td>
<td>32</td>
</tr>
<tr>
<td>Higgs</td>
<td>1350</td>
<td>1173</td>
</tr>
<tr>
<td>Epsilon</td>
<td>7183</td>
<td>8642</td>
</tr>
</tbody>
</table>

a Speed-ups listed are 16 cores vs 1 core for Weka and 128 cores vs 16 cores for Spark. † indicates the speed-up for 128 cores using Spark vs 16 cores using Weka, that is, the gain of the parallel approach.

The left part of Table 7 shows the comparison of the Weka execution times between both versions of the algorithm (sequential and multithreaded using 16 cores) when run on the different datasets that have been previously discretized.

When put in relation with the whole execution time, the speed improvement is negligible. Nevertheless, a deeper analysis of the implementation reveals that most of the time needed to perform InfoGain feature selection in Weka is spent getting the dataset ready, first reading it from disk and then checking that the attributes are fit for the algorithm. The feature selection process itself takes a short time when compared to the total execution time, so even a dramatic improvement in the time efficiency of the algorithm would lead to modest speed-ups for datasets that take a long time to process. Nevertheless, as discussed earlier, some use cases may take advantage of the speed-up obtained when just comparing the time devoted to the algorithm which, in the Weka implementation we are presenting, is close to the number of cores employed, around 16 in this case.

The Spark implementation tested was the one included in the InfoTheoretic Feature Selection Spark package [17]. Results can be seen in right part of Table 7. Instead of the KDD99 dataset, KDDB was used to illustrate how this method is capable of handling very high dimensional datasets.

Although performance increases when adding more cores, for the same number of cores the existing Spark implementation performs much worse than Weka. This results in the need of more nodes to achieve the same times than in Weka, being highly inefficient in terms of resources. For this particular algorithm and datasets it would be more advisable to use Weka on a single machine rather than the existing Spark implementation.

5.4. Analysis of the CFS implementations

The existing multithreaded implementation of the CFS algorithm included in Weka does not offer a significant improvement over the sequential one, being even slower in some cases. This is a result of the parallelization approach used, that requires that the entire correlation matrix is precomputed beforehand, in contrast with the sequential version, that only calculates each value when needed. Since the search method does not try every possible combination of attributes, oftentimes only a small fraction of the correlation matrix needs to be computed. Avoiding to compute these unnecessary values saves significant time that, in some cases, results in smaller computation times than the ones obtained by precomputing the entire correlation matrix with several cores. Our Spark implementation computes the entire correlation matrix every time, but it is still much more time-efficient than the Weka one, as shown in Table 8. The computation time decreases as more nodes are added which, when combined with the much better performance than the Weka algorithm obtained for the same number of cores, results in high speed-ups.

5.5. Analysis of the SVM-RFE implementations

Since the parallelization approach taken for the multithreaded Weka implementation divides the work among classes, multiclass datasets were needed for this experiment. Execution times are shown in Table 9 (please note that in this case times marked with – are executions that take more than three days). The different SVM training algorithm used in Weka and Spark makes a real difference regarding the kind of dataset that can be tackled with each
implementation. The Weka version (and thus our multithreaded version), which uses SMO (see Section 4.4), performs really well when there is a large number of attributes and, therefore, the SVM training process has to be repeated a large number of times. In this case the approach used by the Spark version takes much longer, as for every new training process the data needs to be shuffled. This, in some cases, makes its use unfeasible (for instance, for Isotet and USPS datasets). On the contrary, when datasets have fewer attributes (such as Poker), the SVM training process is repeated fewer times and SGD can be leveraged to train the model with a large number of examples in a much smaller time than SMO. This clearly differentiates both implementations in terms of the datasets that they handle efficiently. Table 9 shows how SGD is suitable for datasets with a large number of attributes and few instances (Isotet and USPS), whereas SMO performs better when there is a large number of instances and fewer attributes (such as Poker).

### 6. Conclusions

This work has explored new implementations of four popular feature selection algorithms. We have proposed new versions that take advantage of multithreaded processing to speed up the computation for their use in Weka and also distributed versions that use Apache Spark, enabling users to tackle bigger datasets in a reasonable time. For those implementations that already existed (see Table 2), tests were performed to assess their suitability for different kinds of datasets.

The experimental results obtained show a significant improvement in execution time for the RELIEF-F algorithm, achieving even superlinear speed-ups for large real-world datasets on a 16 core node, and scaling well in number of nodes for Spark. A considerable improvement was also obtained for a new distributed CFS implementation in Apache Spark that largely outperforms the existing multithreaded version included in Weka, and scales well when more cores are added. A new multithreaded InfoGain implementation was developed and compared to the existing Spark one, finding that its short execution times make the time gain obtained using a cluster less relevant, therefore advising the use of our proposed implementation on a single computer. Lastly, a new SVM-RFE multithreaded implementation enables users to process multiclass datasets up to four times faster than the sequential counterpart included in Weka, and a new Spark version allows the analysis of datasets that because of their dimensions could not be processed by Weka.

As future work, it would be interesting to explore different sampling techniques and their effects on the features selected for a variety of datasets, since this approach may offer a way to use algorithms that are computationally demanding on reduced versions of large datasets. Also pursuing a RELIEF-F implementation with Spark that could handle larger datasets than the ones at reach for the implementation presented in this paper would be advisable.

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### References


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