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Exploiting Machine Learning For Improving In-memory Execution of Data-intensive Workflows on Parallel Machines

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Abstract: Workflows are largely used to orchestrate complex sets of operations required to handle and process huge amounts of data. Parallel processing is often vital to reduce execution time when complex data-intensive workflows must be run efficiently, and at the same time in-memory processing can bring important benefits to accelerate execution. However, optimization techniques are necessary to fully exploit in-memory processing avoiding performance drops due to memory saturation events. This paper proposes a novel solution, called Intelligent In-memory Workflow Manager (IIWM), for optimizing the in-memory execution of data-intensive workflows on parallel machines. IIWM is based on two complementary strategies: 1) a machine learning strategy for predicting memory occupancy and execution time of workflow tasks; 2) a scheduling strategy that allocates tasks to a computing node taking into account the (predicted) memory occupancy and execution time of each task, and the memory available on that node. The effectiveness of the machine learning-based predictor and the scheduling strategy are demonstrated experimentally using as a testbed Spark, a high-performance Big Data processing framework that exploits in-memory computing to speed up execution of large-scale applications. In particular, two synthetic workflows have been prepared for testing the robustness of IIWM in scenarios characterized by a high level of parallelism and a limited amount of memory reserved for execution. Furthermore, a real data analysis workflow has been used as a case study, for better assessing the benefits of the proposed approach. Thanks to high accuracy in predicting resources used at runtime, IIWM was able to avoid disk writes caused by memory saturation, outperforming a traditional strategy in which only dependencies among tasks are taken into account. Specifically, IIWM achieved up to 31\% and 40\% reduction of makespan and a performance improvement up to 1.45x and 1.66x on the synthetic workflows and the real case study respectively.

Keywords: Workflow, Data-intensive, In-memory, Machine Learning, Apache Spark, Scheduling.

1. Introduction

A data-intensive workflow is the description of a process that usually involves a set of computational steps implementing complex scientific functions, such as data acquisition, transformation, analysis, storage, and visualization [1]. Parallelism can be achieved by concurrently executing independent tasks by trying to make use of all computing nodes, even if, in many cases, it is necessary to execute multiple tasks on the same computing node [2]. For example, this occurs when the number of tasks is greater than the number of available nodes, or because multiple tasks use a dataset located on the same node. These scenarios are prone to memory saturation and moving data to disk may result in higher execution times, which leads to the need for a scheduling strategy able to cope with this issue [3,4].
In most cases, distributed processing systems use a-priori policies for handling task execution and data management. For example, in the MapReduce programming model used by Hadoop, mappers write intermediate results after each computation so performing disk-based processing with partial use of memory [5] through the exploitation of the Hadoop Distributed File System (HDFS). On the other hand, Apache Spark\(^1\), that is a state-of-the-art data analysis framework for large-scale data processing exploiting in-memory computing, relies on a Directed Acyclic Graph (DAG) paradigm and is based on: \(i\) an abstraction for data collections which enables parallel execution and fault-tolerance, named Resilient Distributed Datasets (RDDs) [6]; \(ii\) a DAG engine, that manages the execution of jobs, stages and tasks. Besides, it provides different storage levels for data caching and persistence, while performing in-memory computing with partial use of the disk. The Spark in-memory approach is generally more efficient, but a time overhead may be caused by spilling data from memory to disk when memory usage exceeds a given threshold [7]. This overhead can be significantly reduced if memory occupancy of a task is known in advance, to avoid running in parallel two or more tasks that cumulatively exceed the available memory, thus causing data spilling. For this reason, memory is considered a key factor for performance and stability of Spark jobs and Out-of-Memory (OOM) errors are often hard to fix. Recent efforts have been oriented towards developing prediction models for the performance estimation of Big Data applications, although most of the approaches rely on analytical models and only a few recent studies have investigated the use of supervised machine learning models [8–10].

In this work we propose a system, named Intelligent In-memory Workflow Manager (IIWM), specially designed for improving application performance through intelligent usage of memory resources. This is done by identifying clusters of tasks that can be executed in parallel on the same node, optimizing in-memory processing so avoiding the use of disk storage. Given a data-intensive workflow, IIWM exploits a regression model for estimating the amount of memory occupied by each workflow task and its execution time. This model is trained on a log of past executed workflows, represented in a transactional way through a set of relevant features that characterize the considered workflow, such as:

- Workflow structure, in terms of tasks and data dependencies.
- Input format, such as the number of rows, dimensionality, and all other features required to describe the complexity of input data.
- Type of the tasks, i.e., the computation performed by a given node of the workflow. For example, in the case of data analysis workflows, we can distinguish among supervised learning, unsupervised learning, and association rules discovery tasks, and also between learning and prediction tasks.

Predictions made for a given computing node are applicable to all computing nodes of the same type (i.e., having the same architecture, processor type, operating systems, memory resources), which makes the proposed approach effectively usable on large-scale homogeneous HPC systems composed of many identical servers. Given a data-intensive workflow, IIWM exploits the estimates coming from the machine learning model for producing a scheduling plan aimed at reducing (and, in most cases, avoiding) main memory saturation events, which may happen when multiple tasks are executed concurrently on the same computing node. This leads to the improvement of application performance, as swapping or spilling to disk caused by main memory saturation may result in significant time overhead, which can be particularly costly when running workflows involving very large datasets and/or complex tasks.

IIWM has been experimentally evaluated using as a testbed Spark, which is expected to become the most adopted Big Data engine in the next few years [11]. In particular, we assessed the benefits coming from the use of IIWM by executing two synthetic workflows specially generated for investigating

\(^1\) https://spark.apache.org/
specific scenarios related to the presence of a high level of parallelism and a limited amount of memory reserved for execution. The effectiveness of the proposed approach has been further confirmed through the execution of a real data mining workflow as a case study. We carried out an in-depth comparison between IIWM and a traditional blind scheduling strategy, which only considers workflow dependencies for the parallel execution of tasks. The proposed approach showed to be the most suitable solution in all evaluated scenarios outperforming the blind strategy thanks to high accuracy in predicting resources used at runtime, which leads to the minimization of disk writes caused by memory saturation.

The rest of the paper is organized as follows. Section 2 discusses related work. Section 3 describes the proposed system. Section 4 presents and discusses the experimental results. Section 5 concludes the paper.

1.1. Problem statement

The problem addressed in this study consists in the optimization of the in-memory execution of data-intensive workflows, evaluated in terms of makespan (i.e., the total time required to process all given tasks), and application performance. The main reason behind the drop in performance in such workflows is related to the necessity of swapping/spilling data to disk when memory saturation events occur. To cope with this issue, we propose an effective way of scheduling a workflow that minimizes the probability of memory saturation, while maximizes in-memory computing and thus performance.

A workflow $W$ can be represented using a DAG, described by a set of tasks $T = \{t_1, t_2, \ldots, t_n\}$ (i.e., vertices) and dependencies among them $A \subseteq (T \times T) = \{a_1, \ldots, a_m\}$: $a_i = (t_j, t_l), t_j, t_l \in T$ (i.e., directed edges). Specifically, data dependencies (i.e., all input data of a task have already been made available) have to be considered rather than control dependencies (i.e., all predecessors of a task must be terminated before it can be executed), as we refer to data-intensive workflows [12].

Formally, given a set of $q$ computing resources $R = \{r_1, \ldots, r_q\}$, workflow scheduling can be defined as the mapping $T \rightarrow R$ from each task $t \in T$ to a resource $r \in R$, so as to meet a set of specified constraints which influence the choice of an appropriate scheduling strategy [13]. Workflow scheduling techniques are often aimed at optimizing several factors, including makespan and overall cost that in turn depend on data transfer and compute cost [14]. In this study, a multi-objective optimization has been applied, jointly minimizing execution time and memory saturation. This is achieved by using a scheduling strategy that exploits a regression model aimed at predicting the behavior of a given workflow, in terms of resource demand and execution time (see Section 3). For the Reader’s convenience, Table 1 shows the meaning of the main symbols used in the paper.

| Symbol | Meaning |
|--------|---------|
| $T = \{t_1, t_2, \ldots, t_n\}$ | Set of tasks. |
| $A \subseteq (T \times T) = \{a_1, \ldots, a_m\}$ | Dependencies. $a_i = (t_j, t_l), t_j, t_l \in T$. |
| $d_t$ | Description of the dataset processed by task $t$. |
| $W = (T, A)$ | Workflow. |
| $N^{in}(t) = \{t' \in T \mid (t', t) \in A\}$ | In-neighbourhood of task $t$. |
| $N^{out}(t) = \{t' \in T \mid (t, t') \in A\}$ | Out-neighbourhood of task $t$. |
| $M$ | Regression prediction model. |
| $S = (s_1, \ldots, s_k)$ | List of stages. $s_j \subseteq T \mid \{t_k \parallel t_j\}\forall t_k, t_j \in s_j$. |
| $C$ | Maximum amount of memory available for a computing node. |
| $C_s = C - \sum_{t \in s} M.predict.mem(t, d_t)$ | Residual capacity of a stage $s$. |

Table 1. Meaning of the main symbols.
2. Related work

Recent studies have shown the effectiveness of machine learning-based prediction modelling in supporting code optimization, parallelism mapping, task scheduling, and processor resource allocation [10]. Moreover, predicting running times and memory footprint is important for estimating the cost of execution and better managing resources at runtime [11]. For instance, in-memory data processing frameworks like Spark can benefit from informed co-location of tasks [10]. In fact, if too many applications or tasks are assigned to a computing node, such that the memory used on the host exceeds the available one, memory paging to disk (i.e., swapping), data spilling to disk in Spark, or OOM errors can occur with consequential drops of performance.

Our work focuses on improving the performance of a Spark application using machine learning-based techniques. The challenge is to effectively schedule tasks in a data-intensive workflow for improving resource usage and application performance, by inferring the resource demand of each task, in terms of memory occupancy and time.

State-of-the-art techniques aimed at improving the performance of data-intensive applications can be divided into two main categories: analytical-based and machine learning-based. For each category, the main proposed solutions and their differences with respect to our technique are discussed.

2.1. Analytical-based

Techniques in this category use information collected at runtime and statistics in order to tune a Spark application, improving its performance as follows:

- Choosing the serialization strategy for caching RDDs in RAM, based on previous statistics collected on different working sets, such as memory footprint, CPU usage, RDDs size, serialization costs, etc. [15,16].
- Dynamically adapting resources to data storage, using a feedback-based mechanism with real-time monitoring of memory usage of the application [17].
- Scheduling jobs by dynamically adjusting concurrency through a feedback-based strategy. Taking into account memory usage via garbage collection, network I/O, and Spark RDDs lineage information, it is possible to choose the number of tasks to assign to an executor [18,19].

The aforementioned works use different strategies to improve in-memory computing of Spark that exploit static or dynamic techniques able to introduce some information in the choice of configuration parameters. However, no prediction models are employed and this may lead to unpredicted behaviors. IIWM, instead, uses a prediction regression model to estimate a set of information about a running Spark application, exploiting it to optimize in-memory execution. Moreover, unlike real time adapting strategies, which use a feedback-based mechanism by continuously monitoring the execution, the IIWM model is trained offline, achieving fast and accurate predictions while used for inferring the resource demand of each task in a given workflow.

2.2. Machine learning-based

These techniques are based on the development of learning models for predicting performance (mainly memory occupancy and execution time) of a large set of different applications in several scenarios, on the basis of prior knowledge. This enables the adoption of a performance-centric approach [8], based on an informed performance improvement, which can be beneficial for the execution of data-intensive applications, especially in the context of HPC systems.

Several techniques use collaborative filtering to identify how well an application will run on a computing node. For instance, Quasar [8] uses classification techniques based on collaborative filtering to determine the characteristics of the running application in allocating resources and assigning tasks. When submitted, a new application is shortly profiled and the collected information is combined with the classification engine, based on previous workloads, to support a greedy scheduling policy that improves throughput. Application is monitored throughout the execution to adjust resource allocation
and assignment if required, using a single model for the estimation. Adapting this technique to Spark can help to assign tasks to computing nodes within the memory constraints and avoid exceeding the capacity, thus causing spilling of data to disk. Another approach based on collaborative filtering has been proposed by Llull et al. [9]. In this case, the task co-location problem is modelled as a cooperative game and a game-theoretic framework, namely Cooper, is proposed for improving resource usage. The algorithm builds pairwise coalitions as stable marriages to assign an additional task to a host based on its available memory, and the Spark default scheduler is adopted to assign tasks. In particular, a predictor receives performance information collected offline and estimates which co-runner is better, in order to find stable co-locations.

Moving away from collaborative filtering, Marco et al. [10] present a mixture-of-experts approach to model the memory behavior of Spark applications. It is based on a set of memory models (i.e., linear regression, exponential regression, Napierian logarithmic regression) trained on a wide variety of applications. At runtime, an expert selector based on k-nearest neighbour (kNN) is used to choose the model that best describes memory behavior, in order to determine which tasks can be assigned to the same host for improving throughput. The memory models and expert selector are trained offline on different working sets, recording the memory used by a Spark executor through the Linux command "/proc". Finally, the scheduler uses the selected model to determine how much memory is required for an incoming application, for improving server usage and system throughput.

Similarly to machine learning-based techniques, IIWM exploits a prediction model trained on execution logs of previous workflows, however it differs in two main novel aspects: i) IIWM only uses high-level workflow features, without requiring any runtime information as done in [8] and [10], in order to avoid the overhead that could be not negligible for complex applications; ii) it provides an algorithm for effectively scheduling a workflow in scenarios with limited computing resources.

As far as we know, no similar approaches in literature can be directly compared to IIWM in terms of goals and requirements. In fact, differently from IIWM, Quasar [8] and Cooper [9] can be seen as resource-efficient cluster management systems, aimed at optimizing QoS constraints and resource usage. With respect to the most related work, presented in [10], IIWM presents the following differences.

- It focuses on data-intensive workflows while in reference [10] general workloads are addressed.
- It uses high-level information for describing an application (e.g. task and dataset features), while in reference [10] low-level system features are exploited, such as cache miss rate and number of blocks sent, collected by running the application on a small portion (100 MB) of the input data.
- It proposes a more general approach, since the approach proposed in [10] is only appliable to applications whose memory usage is a function of the input size.

3. Materials and Methods

The Intelligent In-memory Workflow Manager (IIWM) is based on three main steps:

1. Execution monitoring and dataset creation: starting from a given set of workflows, a transactional dataset is generated by monitoring the memory usage and execution time of each task, specifying how it is designed and giving concise information about the input.
2. Prediction model training: from the transactional dataset of executions, a regression model is trained in order to fit the distribution of memory occupancy and execution time, according to the features that represent the different tasks of a workflow.
3. Workflow scheduling: taking into account the predicted memory occupancy and execution time of each task, provided by the trained model, and the available memory of the computing node, tasks are scheduled using an informed strategy. In this way, a controlled degree of parallelism can be ensured, while minimizing the risk of memory saturation.

In the following sections, a detailed description of each step is provided.
3.1. Execution monitoring and dataset creation

The first step in IIWM consists of monitoring the execution of different tasks on several input datasets with variable characteristics, in order to build a transactional dataset for training the regression model. The proposed solution was specifically designed for supporting the efficient execution of data analysis tasks, which are used in a wide range of data-intensive workflows. Specifically, it focuses on three classes of data mining tasks: classification tasks for supervised learning, clustering tasks for unsupervised learning and association rules discovery. Using Spark as a testbed, the following data mining algorithms from the MLlib\(^2\) library have been used: Decision Tree, Naive Bayes, and Support Vector Machines (SVM) for classification tasks; K-Means and Gaussian Mixture Models (GMM) for clustering tasks; FPGrowth for association rules tasks.

3.1.1. Execution monitoring within the Spark unified memory model

As far as execution monitoring is concerned, a brief overview of Spark unified memory model is required. In order to avoid OOM errors, Spark uses up to 90% of the heap memory, which is divided into three categories: reserved memory (300 MB), used to store Spark internal objects; user memory (40% of heap memory), used to store data structures and RDDs computed during transformations and actions; spark memory (60% of heap memory), divided in execution and storage. The former refers to that used for computation during shuffle, join, sort, and aggregation processes, while the latter is used for caching RDDs. It is worth noting that, when no execution memory is used, storage can acquire all the available memory and vice versa. However, storage may not evict execution due to complexities in implementation, while stored data blocks are evicted from main memory according to a Least Recently Used (LRU) strategy.

The occupancy of storage memory relies on the persistence operations performed natively by the algorithms. Table 2 reports some examples of data caching implemented in the aforementioned MLlib algorithms. In particular, the `cache()` call corresponds to `persist(StorageLevel.MEMORY_AND_DISK)`, where MEMORY_AND_DISK is the default storage level used for the recent API based on DataFrames.

| MLlib algorithm | Persist call |
|-----------------|--------------|
| K-Means         | //Compute squared norms and cache them norms.cache() |
| DecisionTree    | //Cache input RDD for speed-up during multiple passes BaggedPoint.convertToBaggedRDD(treeInput,...).cache() |
| GMM             | instances.cache() ... data.map(_asBreeze).cache() |
| FP-Growth       | items.cache() |
| SVM             | InstanceBlock.bloifyWithMaxMemUsage(...).cache() |

Table 2. Examples of persist calls in MLlib algorithms.

According to the Spark unified memory model, the execution monitoring was made via the Spark REST APIs, which expose executor-level performance metrics, collected in a JSON file, including peak occupancy for both execution and storage memory along with execution time.

3.1.2. Dataset creation

Using the aforementioned Spark APIs, we monitored the execution of several MLlib algorithms on different input datasets, covering the main data mining tasks, i.e. classification, clustering, and

\(^2\) https://spark.apache.org/mllib/
association rules. The goal of this process is the creation of a transactional dataset for the regression
model training, which contains the following information:

- The description of the task, such as its class (e.g., classification, clustering, etc.), type (fitting or predicting task), and algorithm (e.g., SVM, K-Means, etc.).
- The description of the input dataset in terms of the number of rows, columns, categorical columns and overall dataset size.
- Peak memory usage (both execution and storage) and execution time, which represent the three target variables to be predicted by the regressor. In order to obtain more significant data, the metrics were aggregated on median values by performing ten executions per task.

For the sake of clarity, table 3 shows a sample of the dataset described above.

| Task Name      | Task Type  | Task Class    | Dataset Rows | Dataset Columns | Categorical Columns | Dataset Size (MB) | Peak Storage Memory (MB) | Peak Execution Memory (MB) | Duration (ms) |
|----------------|------------|---------------|--------------|-----------------|--------------------|---------------------|--------------------------|---------------------------|---------------|
| GMM Estimator  | Clustering | 1474971       | 28           | 0               | 87.0045            | 433.37              | 1413.5                   | 108204                    |               |
| K-Means Estimator | Clustering | 5000000       | 104          | 0               | 1239.78            | 4624.52             | 4112                     | 56322.5                   |               |
| DecisionTree Estimator | Classification | 9606       | 1921         | 0               | 84.9105            | 730.09              | 297.95                   | 39292                     |               |
| NaiveBayes Estimator | Classification | 269924     | 4            | 0               | 13.4986            | 340.92              | 6982.82                  | 16531.5                   |               |
| SVM Estimator | Classification | 5000000       | 129          | 0               | 1542.38            | 6199.11             | 106.6                    | 238594.5                  |               |
| FPGrowth Estimator | AssociationRules | 825095     | 180         | 180             | 697                | 9495.85             | 1371.03                  | 96071.5                   |               |
| GMM Transformer | Clustering | 165474        | 14           | 1               | 636604             | 2.34                | 1e-06                    | 62.5                      |               |
| K-Means Transformer | Clustering | 489431        | 42           | 3               | 648.887            | 2.23                | 1e-06                    | 35                       |               |
| DecisionTree Transformer | Classification | 195972      | 42           | 4               | 257.686            | 3.68                | 1e-06                    | 65.5                      |               |
| NaiveBayes Transformer | Classification | 347899     | 4            | 0               | 17.9982            | 4.26                | 1e-06                    | 92.5                      |               |
| SVM Transformer | Classification | 5000000       | 129          | 0               | 1542.38            | 2.36                | 1e-06                    | 55.5                      |               |
| FPGrowth Transformer | AssociationRules | 136073     | 34           | 34              | 13.5483            | 1229.95             | 63.5                     | 52449                     |               |

Table 3. A sample of the training dataset.

Starting from 20 available datasets, we divided them into two partitions used for training and testing respectively. Afterwards, an oversampling procedure was performed, aimed at increasing the number of datasets contained in the partitions. Specifically, a naive random sampling approach can lead to unexpected behaviors regarding the convergence of algorithms, thus introducing noise into the transactional dataset used to build the regression model. To cope with this issue, we used the following feature selection strategy:

- For datasets used in classification or regression tasks we considered only the $k$ highest scoring features based on:
  - analysis of variance (F-value) for integer labels (classification problems);
  - correlation-based univariate linear regression test for real labels (regression problems).
- For clustering datasets we used a correlation-based test to maintain the $k$ features with the smallest probability to be correlated with the others.
- For association rules discovery datasets no features selection is required, as the number of columns refers to the average number of items in the different transactions.

The described procedure has been applied separately on the training and test partitions, so as to avoid the introduction of bias into the evaluation process. Specifically, the number of datasets in the training and test partitions has increased from 15 to 260 and from 5 to 86 respectively. Subsequently, we fed these datasets to the MLlib algorithms, obtaining two final transactional datasets of 1309 and 309 monitored executions, used for training and testing the regressor, respectively.

3.2. Prediction model training

Once the training and test datasets with memory and time information were built, a regression model can be trained with the goal of estimating peak memory occupancy and turnaround time of a task in a given workflow.

As a preliminary step, we analyzed the correlation between the features of the training data and each target variable, using the Spearman index. We obtained the following positive correlations:
value of 0.30 between storage memory and the input dataset size, 0.46 between execution memory and the task class and 0.21 between execution time and the number of columns. These results can be seen in detail in Figure 1.

![Figure 1](image_url)

**Figure 1.** Correlation of target variables with the other features.

Afterwards we moved to the training of the regression model. Due to its complexity, the regression problem cannot be faced with a simple linear regressor or its regularized variants (e.g. Ridge, Lasso or ElasticNet), but a more robust model is necessary. We experimentally evaluated this aspect by testing the forecasting abilities of these linear models achieving poor results. For this reason, an ensemble learning model has been used in order to fit the nonlinear distribution of features. Specifically, the *stacking* technique (meta learning) [20] has been used by developing a two-layer model in which a set of regressors are trained on the input dataset and a *Decision Tree* is fitted on their predictions. The first layer consists of three tree-based regressors, able to grasp different aspects of input data: a *Gradient Boosting*, an *AdaBoost* and an *Extra Trees* regressor. The second layer exploits a single *Decision Tree* regressor, which predicts the final value starting from the concatenation of the outputs from the first layer. The described ensemble model has been set with the hyper-parameters shown in Table 4.

| Hyper-parameter | Value         |
|-----------------|---------------|
| n_estimators    | 500           |
| learning_rate   | 0.01          |
| max_depth       | 7             |
| loss            | least squares |

**Table 4.** Hyper-parameters.

Among 20 trained models, initialized with different random states, we selected the best one by maximizing the following objective function:

\[ O = R^2 - MAE \]

whose goal is to choose the model that best explains the variance of data, while minimizing the forecasting error. This function jointly considers the adjusted determination coefficient (\( R^2 \)), which guarantees robustness with respect to the addition of useless variables to the model compared to the classical \( R^2 \) score, and the mean absolute error (\( MAE \)), normalized with respect to the maximum.

The described model has been developed in *Python3* using the *scikit-learn*³ library and evaluated against the test set of 309 unseen executions obtained as described in Section 3.1.2. Thanks to the

³ [https://scikit-learn.org/stable/](https://scikit-learn.org/stable/)
combination of different models, the ensemble technique showed to be very well suited for this task, leading to good robustness against outliers and a high forecasting accuracy, as shown in Figure 2.

![Figure 2](image-url)

**Figure 2.** Meta-learner regression estimates for the different target variables.

These results are detailed in Table 5, which shows the evaluation metrics for each target variable, including the $R^2$ score and the Pearson correlation coefficient. In particular, the Mean Absolute Error (MAE) and the Root Mean Square Error (RMSE) for the storage and execution memory represent average errors in megabytes, while for the duration they represent errors in milliseconds.

|                | RMSE   | MAE    | Adjusted $R^2$ | Pearson Correlation |
|----------------|--------|--------|----------------|---------------------|
| Storage Memory | 108.23 | 26.66  | 0.96           | 0.98                |
| Execution Memory | 312.60 | 26.30  | 0.91           | 0.95                |
| Duration       | 4443.17| 2003.70| 0.95           | 0.98                |

**Table 5.** Evaluation metrics on the test set.

3.3. Workflow scheduling

The prediction model described in Section 3.2 can be exploited to forecast the amount of memory that will be needed to execute a given task on a target computing node and its duration, based on the task features listed in Section 3.1. These predictions are then used within the scheduling strategy described in the following, whose goal is to avoid swapping to disk due to memory saturation in order to improve application performance and makespan through a better use of in-memory computing.

The results discussed below refer to a static scheduling problem, as the scheduling plan is generated before the execution. In typical static scheduling the workflow system has to predict the execution load of each task accurately, using heuristic-based methods [21]. Likewise, in the proposed method the execution load of each task of a given workflow is predicted by the model trained on past executions. Moreover, we investigated how workflow tasks can be scheduled and run on a single computing node, but this approach can be easily generalized to a multi-node scenario. For example, a data-intensive workflow can be decomposed into multiple sub-workflows to be run on different computing nodes according to their features and data locality. Each sub-workflow is scheduled locally to the assigned node using the proposed strategy.

In IIWM, we modelled the scheduling problem as an offline Bin Packing (BP). This is a well-known problem, widely used for resource and task management or scheduling, such as load balancing in mobile cloud computing architectures [22], energy-efficient execution of data-intensive applications in clouds [23], DAGs real-time scheduling in heterogeneous clusters [24] and task scheduling in multiprocessor environments [25]. Its classical formulation is as follows [26]. Let $n$ be the number of items, $w_j$ the weight of the $j$-th item and $c$ the capacity of each bin: the goal is to assign each item to a bin without exceeding the capacity $c$ and minimizing the number of used bins. The problem is $\mathcal{NP}$-complete and a lot of effort went into finding fast algorithms with near-optimal solutions. We adapted the classical problem to our purposes as follows:
• An item is a task to be executed.

• A bin identifies a stage, i.e. a set of tasks that can be run in parallel.

• The capacity of a bin is the maximum amount $C$ of available memory in a computing node.

   When assigning a task to a stage $s \in S$, its residual available memory will be indicated with $C_s$.

• The weight of an item is the memory occupancy estimated by the prediction model. In the case of Spark testbed, it will be the maximum of the execution and storage memory, in order to model a peak in the unified memory. For what concerns the estimated execution time, it is used for selecting the stage to be assigned when memory constraints hold for multiple stages.

With respect to the classical BP problem two changes were introduced:

• All workflow tasks have to be executed, so the capacity of a stage may still be exceeded if a task takes up more memory than the available one.

• The assignment of a task $t$ to a stage $s$ is subjected to dependency constraints. Hence, if a dependency exists between $t_i$ and $t_j$, then the stage of $t_i$ has to be executed before the one of $t_j$.

To solve the BP problem, modelled as described above, in order to produce the final scheduling plan, we used the First Fit Decreasing algorithm which assigns tasks sorted in non-increasing order of weight. However, the introduction of dependency constraints in the assignment process may cause the under-usage of certain stages. To cope with this issue, we introduced a further step of consolidation, aimed at reducing the number of stages by merging together stages without dependencies according to the available memory. The main execution flow of the IIWM scheduler is shown in Figure 3 and described by Algorithm 1. In particular, given a data-intensive workflow $W$ as a DAG by its tasks and dependencies, and the prediction model $M$ as input, a scheduling plan is generated in two steps: i) building of the stages and task assignment; ii) stage consolidation.

The algorithm is divided into two main parts: in the first part (lines 1-23), the stages are built by iteratively assigning each task according to the estimates of the prediction model; in the second part (lines 25-34), a consolidation process is performed, trying to minimize the number of stages.

The first part (lines 1-23) starts with the initialization of an empty list of stages $S$, which will be filled according to a dictionary $Q$ that stores the in-degree of each task in the DAG, which is used for identifying the free tasks which can be scheduled. The prediction model $M$ is exploited to estimate the memory occupancy and execution time of each task in $T$, according to their dataset description (lines 3-4). The dictionary $P_{mem}$, which collects the predicted memory occupancies, is then used to sort tasks according to the First Fit Decreasing strategy (line 5). At each iteration, tasks that can be scheduled (i.e., assigned to a stage) are collected in the $T_{free}$ set. In particular, they are identified by a zero in-degree, as their execution does not depend on others (line 7). By virtue of the acyclicity of the DAG-based workflow representation, there will always exist a task $t \in T$ with a zero in-degree not yet scheduled, unless set $T$ is empty. Afterwards, the task with the highest memory occupancy is
selected from $T_{\text{free}}$ in order to be scheduled (line 8). At this point, a list of candidate stages ($S_{\text{sel}}$) for the selected task is identified according to the peak memory occupancy forecasted by the prediction model $M$ (lines 9-10). In particular, a stage $s_i$ belongs to $S_{\text{sel}}$ if it satisfies the following conditions:

- The residual capacity $C_{s_i}$ of the selected stage $s_i$ is not exceeded by the addition of the task $t$.
- There not exists a dependency between $t$ and any task $t'$ belonging to $s_i$ and every subsequent stage $(s_{i+1} \cup \cdots \cup s_k)$, where a dependency $(t',t)^n$ is identified by a path of length $n > 0$.

If there exist one or more candidate stages $S_{\text{sel}}$ (line 11), the best one is chosen based on the minimum marginal increase. Specifically, for each of these stages, the expected increase of the execution time is estimated (lines 12-13), assigning the task $t$ to the stage $s$ with the lowest value (lines 14-16). Otherwise (line 17), a newly created stage is allocated for $t$ and added to the list $S$ (lines 18-21). Once the task $t$ is assigned to the stage $s$, the residual capacity $C_s$ is updated (lines 15, 20). Then, the residual in-degree for every task in the out-neighbourhood of $t$ (line 22) is decremented by updating the dictionary $Q$, so as to allow the assignment of these tasks in the next iterations. Finally, the assigned task $t$ is removed from the set of workflow nodes $T$ (line 23).

**ALGORITHM 1: IIWM Scheduler**

**Input:** Workflow $W = (T, A)$, Prediction model $M$

**Output:** A list of stages $S$

1. $S \leftarrow \emptyset$
2. $Q \leftarrow \{ t : |N^\text{in}(t)|, \forall t \in T \}$
3. $P_{\text{mem}} \leftarrow \{ t : M.\text{predict_mem}(t,d), \forall t \in T \}$ ➨ Memory prediction for each task in $T$
4. $P_{\text{time}} \leftarrow \{ t : M.\text{predict_time}(t,d), \forall t \in T \}$ ➨ Time prediction for each task in $T$
5. $T \leftarrow \text{sort}\_\text{decreasing}(T, P_{\text{mem}})$
6. while $T \neq \emptyset$
7.     $T_{\text{free}} \leftarrow \{ t \in T | Q[t] == 0 \}$
8.     $t \leftarrow \text{get}\_\text{first}(T_{\text{free}})$
9.     $\text{mem}_t \leftarrow P_{\text{mem}}[t]$
10. $S_{\text{sel}} \leftarrow \{ s_i \in S | \text{mem}_t \leq C_{s_i} \text{ and } C_s(n) \in A, n > 0, \forall t' \in s_i \cup s_{i+1} \cup \cdots \cup s_k \}$
11. if $S_{\text{sel}} \neq \emptyset$ then
12.     duration $\leftarrow \{ s : \max_{t \in S} P_{\text{time}}[t], \forall s \in S_{\text{sel}} \}$
13.     increase $\leftarrow \{ s : \max_{t \in S} P_{\text{time}}[t], \text{duration}[s], \forall s \in S_{\text{sel}} \}$
14.     $s \leftarrow \text{argmin}_{s \in S_{\text{sel}}} \text{increase}$
15.     $C_s \leftarrow C_s - \text{mem}_t$
16.     $s \leftarrow s \cup \{ t \}$
17. else
18.     $s \leftarrow \emptyset$
19.     $s \leftarrow s \cup \{ t \}$
20.     $C_s \leftarrow C_s - \text{mem}_t$
21.     $S \leftarrow S \cup \{ s \}$
22.     $Q[t'] = Q[t] - 1$, $\forall t' \in N^\text{out}(t)$
23.     $T \leftarrow T \setminus \{ t \}$
24. // Consolidation step
25. $S_{\text{mem}} \leftarrow \{ s \in S | |N^\text{out}(t)| == 0, \forall t \in s \}$
26. if $S_{\text{mem}} \neq \emptyset$ then
27.     for $s_i \in S_{\text{mem}}$ do
28.         for $s_j \in S | j > i$ do
29.             $\text{mem}_{s_i,s_j} \leftarrow \sum_{t \in s_i \cup s_j} P_{\text{mem}}[t]$
30.         if $\text{mem}_{s_i,s_j} \leq C$ then
31.             $s_j \leftarrow s_i \cup s_j$
32.         $S \leftarrow S \setminus s_j$
33. break
34. return $S$
The second part of the algorithm (lines 25-34) performs a consolidation step with the goal of reducing the number of allocated stages by merging some of them if possible, with a consequent improvement in the global throughput. The stages involved in the consolidation step, namely the movable stages ($S_{mov}$), are those containing tasks with a zero out-degree (line 25). This means that no task in such stages blocks the execution of another one, so they can be moved forward and merged with subsequent stages if the available capacity $C$ is not exceeded. For each movable stage $s_i$ (line 27), another stage $s_j$ from $S$ is searched among the subsequent ones, such that its residual capacity is enough to enable the merging with $s_i$ (lines 28-30). The merging between $s_i$ and $s_j$ is performed by assigning to $s_j$ each task of $s_i$ (line 31), finally removing $s_i$ from $S$ (line 32). In the end, the list of stages $S$ built by the scheduler is returned as output. Given this scheduling plan, the obtained stages will be executed in sequential order, while all the tasks in a stage will run concurrently.

Compared to a blind strategy where the maximum parallelism is achieved by running in parallel all the tasks not subjected to dependencies, which will be referred to as Full-Parallel in our experiments, IIWM can reduce both delays of parallelization ($\epsilon_p$), due to context switch and process synchronization, and swapping/spilling to disk ($\epsilon_s$), due to I/O operations. Delay $\epsilon_p$ is always present in all scheduling strategies when two or more tasks are run concurrently, while $\epsilon_s$ is present only when a memory saturation event occurs. Given $\epsilon = \epsilon_p + \epsilon_s$, IIWM mainly reduces $\epsilon_s$, which is the main factor behind the drop in performance in terms of execution time, due to the slowness in accessing secondary storage.

As far as the Spark framework is concerned, the proposed strategy is effective for making the most of the default storage level, i.e. MEMORY_AND_DISK: at each internal call of the cache() method, data is saved in-memory as long as this resource is available, using disk otherwise. In this respect, IIWM can reduce the actual persistence of data on disk by better exploiting in-memory computing.

4. Results and Discussion

This section presents an experimental evaluation of the proposed system, aimed at optimizing the in-memory execution of data-intensive workflows. We experimentally assessed the effectiveness of IIWM using Apache Spark 3.0.1 as a testbed. In particular, we generated two synthetic workflows for analyzing different scenarios, by assessing also the benefits coming from the use of IIWM using a real data mining workflow as a case study.

In order to provide significant results, each experiment was executed ten times and the average metrics with standard deviations are reported. In particular, for each experiment, we evaluated the accuracy of the regression model in predicting memory occupancy and execution time.

We evaluated the ability of IIWM to improve application performance taking into account two different aspects:

- **Execution time**: let $m_1$ and $m_2$ be the makespan for two different executions. If $m_2 < m_1$ we can compute the improvement on makespan ($m_{imp}$) and application performance ($p_{imp}$) as follows:

$$m_{imp} = \frac{m_1 - m_2}{m_1} \times 100\% \quad p_{imp} = \frac{m_1}{m_2}$$

- **Disk usage**: we used the on-disk usage metric, which measures the amount of disk usage, jointly considering the volume and the duration of disk writes. Formally, given a sequence of disk writes $w_1, ..., w_k$ let $\tau_i^\prime$, $\tau_i^\prime\prime \in T$ be the start and end time of the $w_i$ write respectively. Let also $W : T \to \mathbb{R}$ be a function representing the amount of megabytes written to disk over time $T$. We define on-disk usage as:

$$\text{on-disk usage} = \sum_{i=1}^{k} \frac{1}{\tau_i^\prime - \tau_i^\prime\prime} \int_{\tau_i^\prime}^{\tau_i^\prime\prime} W(\tau)d\tau$$

Specifically, for each workflow we reported: i) a comparison between Full-Parallel and IIWM in terms of disk usage over time; ii) a detailed description of the scheduling plan generated by both
strategies; iii) the average improvement on makespan and application performance with IIWM; iv) statistics about the use of disk, such as the time spent for I/O operations and the on-disk usage metric; v) the execution of the workflow by varying the amount of available memory, in order to show the benefits of the proposed scheduler in different limited memory scenarios.

4.1. Synthetic workflows

We firstly evaluated our approach against two complex synthetic data analysis workflows, where the Full-Parallel approach showed its limitations due to a high degree of parallelism. The dependencies in these workflows should be understood as execution constraints. For instance, clustering has to be performed before classification for adding labels to an unlabelled dataset, or a classification task is performed after the discovery of association rules for user classification purposes.

The first test has been carried out on a synthetic workflow with 42 nodes. Table 6 provides a detailed description of each task in the workflow, while their dependencies are shown in Figure 4.

| Node | Task Name | Task Type | Task Class | Rows | Columns | Categorical Columns | Dataset size (MB) |
|------|-----------|-----------|------------|------|---------|---------------------|------------------|
| 0    | FPGrowth | Estimator | AssociationRules | 494156 | 180 | 180 | 417.03807 |
| 1    | NaiveBayes| Estimator  | Classification | 2939059 | 18 | 4 | 198.93904 |
| 2    | SVM       | Estimator  | Classification | 19213  | 1442 | 0 | 123.28475 |
| 3    | NaiveBayes| Estimator  | Classification | 2939059 | 18 | 4 | 198.93904 |
| 4    | DecisionTree | Estimator | Classification | 14410 | 1921 | 0 | 127.3811 |
| 5    | NaiveBayes| Estimator  | Classification | 2000000 | 104 | 0 | 252.72347 |
| 6    | NaiveBayes| Estimator  | Classification | 2939059 | 18 | 4 | 198.93904 |
| 7    | K-Means   | Estimator  | Clustering    | 2939059 | 18 | 4 | 198.93904 |
| 8    | NaiveBayes| Estimator  | Classification | 2939059 | 18 | 4 | 198.93904 |
| 9    | K-Means   | Estimator  | Clustering    | 2939059 | 18 | 4 | 198.93904 |
| 10   | NaiveBayes| Estimator  | Classification | 1959372 | 18 | 4 | 132.62437 |
| 11   | K-Means   | Estimator  | Clustering    | 2939059 | 18 | 4 | 198.93904 |
| 12   | NaiveBayes| Estimator  | Classification | 1000000 | 53 | 0 | 126.36286 |
| 13   | K-Means   | Estimator  | Clustering    | 2939059 | 18 | 4 | 198.93904 |
| 14   | NaiveBayes| Estimator  | Classification | 1959372 | 18 | 4 | 132.62437 |
| 15   | SVM       | Estimator  | Classification | 19213  | 1442 | 0 | 123.28475 |
| 16   | NaiveBayes| Estimator  | Classification | 1000000 | 53 | 0 | 126.36286 |
| 17   | K-Means   | Estimator  | Clustering    | 1000000 | 129 | 0 | 154.5775 |
| 18   | NaiveBayes| Estimator  | Classification | 2000000 | 104 | 0 | 252.72347 |
| 19   | K-Means   | Estimator  | Clustering    | 2939059 | 18 | 4 | 198.93904 |
| 20   | NaiveBayes| Estimator  | Classification | 1959372 | 18 | 4 | 132.62437 |
| 21   | NaiveBayes| Estimator  | Classification | 1000000 | 53 | 0 | 126.36286 |
| 22   | K-Means   | Estimator  | Clustering    | 2939059 | 18 | 4 | 198.93904 |
| 23   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |
| 24   | NaiveBayes| Estimator  | Classification | 1000000 | 53 | 0 | 126.36286 |
| 25   | K-Means   | Estimator  | Clustering    | 14410 | 1921 | 0 | 127.3811 |
| 26   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |
| 27   | NaiveBayes| Estimator  | Classification | 1000000 | 53 | 0 | 126.36286 |
| 28   | K-Means   | Estimator  | Clustering    | 14410 | 1921 | 0 | 127.3811 |
| 29   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |
| 30   | NaiveBayes| Estimator  | Classification | 1000000 | 53 | 0 | 126.36286 |
| 31   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |
| 32   | NaiveBayes| Estimator  | Classification | 1000000 | 53 | 0 | 126.36286 |
| 33   | NaiveBayes| Estimator  | Classification | 1000000 | 53 | 0 | 126.36286 |
| 34   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |
| 35   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |
| 36   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |
| 37   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |
| 38   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |
| 39   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |
| 40   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |
| 41   | SVM       | Estimator  | Classification | 14410 | 1921 | 0 | 127.3811 |

Table 6. Task and dataset descriptions (workflow 1).

The first test has been carried out on a synthetic workflow with 42 nodes. Table 6 provides a detailed description of each task in the workflow, while their dependencies are shown in Figure 4.

4.2. Performance evaluation of the prediction model

We firstly considered a configuration characterized by 14 GB available for running the workflow, which will be used up to 60% due to the Spark unified memory model. Table 8 shows an execution.
example with IIWM, focusing on its main steps: i) the scheduling of tasks based on their decreasing memory weight; ii) the allocation of a new stage; iii) the exploitation of the estimated execution time while computing the marginal increase. This last aspect can be clearly observed in iteration 17, where task \( t_{17} \) is assigned to stage \( s_7 \), which presents a marginal increase equal to zero. This is the best choice compared to the other candidate stage (\( s_0 \)), whose execution time would be increased by 12496 milliseconds by the assignment of \( t_{17} \), with a degradation of the overall makespan.

### Table 8. Example of execution of algorithm 1 at iteration level.

| Iteration | State | Stages |
|-----------|-------|--------|
| lt. 0     | \( T_{\text{free}}^0 = \{ t_0 \} \) | \( s_0 = \{ t_0 \} \) |
|           | Create \( s_0 \) and assign \( t_0 \) Unlock \([t_1,t_2,t_3]\) | |
| lt. 1     | \( T_{\text{free}}^1 = \{ t_1, t_3, t_2 \} \) | \( s_0 = \{ t_0 \}, s_1 = \{ t_1 \} \) |
|           | Create \( s_1 \) and assign \( t_1 \) Unlock \([t_4]\) | |
| lt. 2     | \( T_{\text{free}}^2 = \{ t_3, t_4, t_2 \} \) | \( s_0 = \{ t_0 \}, s_1 = \{ t_1 \}, s_2 = \{ t_3 \} \) |
|           | Create \( s_2 \) and assign \( t_3 \) Unlock \([t_7,t_8,t_9,t_{10}]\) | |
| lt. 3     | \( T_{\text{free}}^3 = \{ t_7, t_4, t_{10}, t_2, t_8, t_9 \} \) | \( s_0 = \{ t_0 \}, s_1 = \{ t_1 \}, s_2 = \{ t_3 \}, s_3 = \{ t_7 \} \) |
|           | Create \( s_3 \) and assign \( t_7 \) | |
| lt. 4     | \( T_{\text{free}}^4 = \{ t_4, t_{10}, t_2, t_8, t_9 \} \) | \( s_0 = \{ t_0 \}, s_1 = \{ t_1 \}, s_2 = \{ t_3, t_4 \}, s_3 = \{ t_7 \} \) |
|           | \( S_{\text{sel}} = \{ s_2, s_3 \} \) |
|           | \( \text{increase} = \{ 0, 0 \} \) |
|           | Assign \( t_4 \) to \( s_2 \) | |
| ...       | ...   | ...
| ...       | ...   | ...
| lt. 17    | \( T_{\text{free}}^{17} = \{ t_{17}, t_{23}, t_8, t_9 \} \) | \( s_0 = \{ t_0 \}, s_1 = \{ t_1, t_2 \}, s_2 = \{ t_3, t_4, t_5 \}, s_3 = \{ t_7, t_{10}, t_6 \} \) |
|           | \( S_{\text{sel}} = \{ s_6, s_7 \} \) | \( s_4 = \{ t_{12}, t_{11} \}, s_5 = \{ t_{15}, t_{18} \}, s_6 = \{ t_{19}, t_{22} \}, s_7 = \{ t_{24}, t_{16}, t_{17} \} \) |
|           | \( \text{increase} = \{ 12496,363,0 \} \) |
|           | Assign \( t_{17} \) to \( s_7 \) Unlock \([t_{25}]\) | |

At the end of the process, a consolidation step is exploited for optimizing throughput and execution time, by merging two stages with zero out-degree with some tailing stages, so as to avoid the sequential execution of the two stages in favour of a parallel one.

Figure 5 shows disk occupancy throughout the execution. As a consequence of memory saturation, the execution of Full-Parallel resulted in a huge amount of disk writes, while IIWM achieved a null disk usage since no swapping occurred thanks to intelligent task scheduling. Thus, this translates into better use of in-memory computing.
These results can be clearly seen also in Table 9, which shows the scheduling plan produced by the IIWM scheduler, together with some statistics about execution times and the use of the disk. In particular, given the curves representing disk writes over time shown in Figure 5, on-disk usage graphically represents the sum, for each disk write, of the ratio between the area under the curve identified by a write and its duration. Compared to the Full-Parallel strategy, IIWM achieved better execution times and an improvement in application performance, with a boost of almost 1.45x ($p_{imp}$) and a 31.15% reduction in time ($m_{imp}$) on average.

| Strategy   | Task-scheduling plan                                                                 | Number of stages | Time (min.) | Peak disk usage (MB) | Writes duration (min.) | On-disk usage (MB) |
|------------|---------------------------------------------------------------------------------------|-------------------|-------------|----------------------|------------------------|--------------------|
| Full-Parallel | $\{(t_0), (t_1 \parallel t_2 \parallel t_3), (t_4 \parallel t_5 \parallel t_6 \parallel t_7 \parallel t_8 \parallel t_9 \parallel t_{10})\}$, $\{(t_{11} \parallel t_{12} \parallel t_{13} \parallel t_{14})\}$, $\{(t_{15} \parallel t_{16} \parallel t_{17} \parallel t_{18} \parallel t_{19} \parallel t_{20} \parallel t_{21})\}$, $\{(t_{22} \parallel t_{23} \parallel t_{24} \parallel t_{25} \parallel t_{26} \parallel t_{27} \parallel t_{28})\}$, $\{(t_{29} \parallel t_{30} \parallel t_{31} \parallel t_{32} \parallel t_{33} \parallel t_{34} \parallel t_{35})\}$, $\{(t_{36} \parallel t_{37} \parallel t_{38} \parallel t_{39} \parallel t_{40})\}$ | 10                | 31.52 ± 0.6  | 356106.601  | 11.56                  | 126867.065         |
| IIWM      | $\{(t_0), (t_1 \parallel t_2), (t_3 \parallel t_4), (t_7 \parallel t_8 \parallel t_9)\}$, $\{(t_{12} \parallel t_{13} \parallel t_{14}, (t_{15} \parallel t_{16} \parallel t_{17} \parallel t_{18} \parallel t_{19} \parallel t_{20} \parallel t_{21})\}$, $\{(t_{22} \parallel t_{23} \parallel t_{24} \parallel t_{25} \parallel t_{26} \parallel t_{27} \parallel t_{28})\}$, $\{(t_{29} \parallel t_{30} \parallel t_{31} \parallel t_{32} \parallel t_{33} \parallel t_{34} \parallel t_{35})\}$, $\{(t_{36} \parallel t_{37} \parallel t_{38} \parallel t_{39} \parallel t_{40})\}$ | 16                | 21.70 ± 0.63 | 0           | 0                     | 0                   |

Table 9. Scheduling plan and statistics about execution times and disk usage with 14 GB of RAM.

With different sizes of available memory, the Full-Parallel approach showed higher and higher execution times and disk writes as memory decreased, while IIWM was able to adapt the execution to available resources as shown in Figure 6, finding a good trade-off between the maximization of the parallelism and the minimization of the memory saturation probability. At the extremes, with unlimited available memory, or at least greater than that required to run the workflow, IIWM will perform as a full concurrent strategy, producing the same scheduling of Full-Parallel.

The second synthetic workflow consists of the 27 tasks described by Table 10 and their dependencies, shown in Figure 7. This scenario is characterized by highly heavy tasks and very low resources, where the execution of a single task can exceed the available memory. In particular, task $T_{18}$ has an estimated peak memory occupancy higher than Spark available unified memory of 5413.8 MB (i.e., corresponding to a heap size of 9.5 GB): this will bring the IIWM scheduling algorithm to allocate the task to a new stage, but memory will be saturated anyway.

In such a situation, data spilling to disk cannot be avoided, but IIWM tries to minimize the number of bytes written and the duration of I/O operations. Even in this scenario, the prediction model achieved very accurate results, shown in Table 11, confirming its forecasting abilities.
Table 10. Task and dataset descriptions (workflow 2).

| Node | Task Name  | Task Type     | Task Class | Rows | Columns | Categorical Columns | Dataset size (MB) |
|------|------------|---------------|------------|------|---------|---------------------|-------------------|
| t0  | K-Means Estimator | Clustering   | 3918745    | 34   | 4       | 446.54932           |
| t1  | DecisionTree Estimator | Classification | 4000000   | 27   | 0       | 257.4918           |
| t2  | GMM Estimator | Clustering   | 2458285    | 28   | 0       | 120.24             |
| t3  | DecisionTree Estimator | Classification | 3918745   | 129  | 0       | 265.2537           |
| t4  | DecisionTree Estimator | Classification | 4984531   | 42   | 3       | 688.97             |
| t5  | DecisionTree Estimator | Classification | 2996959   | 42   | 4       | 386.53033          |
| t6  | K-Means Estimator | Clustering   | 2458285    | 56   | 0       | 278.75266          |
| t7  | GMM Estimator | Clustering   | 3000000    | 53   | 0       | 157.8823           |
| t8  | SVM Estimator | Classification | 4000000   | 53   | 0       | 505.45             |
| t9  | K-Means Estimator | Clustering   | 2996959    | 42   | 4       | 265.2537           |
| t10 | SVM Estimator | Classification | 2000000   | 53   | 0       | 372.75266          |
| t11 | K-Means Estimator | Clustering   | 2996959    | 9    | 2       | 371.93442          |
| t12 | NaiveBayes Estimator | Classification | 265924    | 3    | 0       | 10.3273            |
| t13 | K-Means Estimator | Clustering   | 2000000    | 78   | 0       | 371.93442          |
| t14 | DecisionTree Estimator | Classification | 3918745   | 26   | 4       | 466.54932          |
| t15 | FPGrowth Estimator | AssociationRules | 823593  | 180  | 180     | 692.75266          |
| t16 | DecisionTree Estimator | Classification | 3918745   | 26   | 4       | 265.2537           |
| t17 | DecisionTree Estimator | Classification | 2996959   | 42   | 3       | 688.97             |
| t18 | FPGrowth Estimator | AssociationRules | 164719   | 180  | 180     | 139.75266          |
| t19 | GMM Estimator | Clustering   | 3000000    | 27   | 0       | 193.119            |
| t20 | DecisionTree Estimator | Classification | 2000000   | 104  | 0       | 348.4647          |
| t21 | K-Means Estimator | Clustering   | 2458285    | 69   | 2       | 473.94803          |
| t22 | FPGrowth Estimator | AssociationRules | 494156  | 180  | 180     | 417.01007          |

Table 11. Performance evaluation of the prediction model.

| RMSE      | MAE      | Adjusted $R^2$ | Pearson Correlation |
|-----------|----------|----------------|---------------------|
| Storage Memory | 213.81   | 78.92          | 0.98                | 0.99               |
| Execution Memory | 29.86   | 11.56          | 0.98                | 0.99               |
| Duration   | 20086.80 | 9925.13        | 0.82                | 0.94               |

Figure 8 shows disk occupancy during the execution. As we can see, even IIWM cannot avoid data spilling, even though its disk usage was much lower considering peak value and writes duration compared to Full-Parallel.

Finally, Table 12 shows the statistics about disk usage and execution times. Again, IIWM achieved better results with a boost in performance of almost $1.30x (p_{imp})$ with respect to a Full-Parallel strategy and a 23% reduction in time ($m_{imp}$) on average. An interesting aspect that emerges from the behavior of IIWM scheduler, in the task-scheduling plan, is the similarity with priority-based scheduling in assigning tasks based on decreasing weights. In fact, tasks characterized by low memory occupancy may be assigned to tailing stages even if they are close to the root (e.g., in Full-Parallel, $t_1$ is executed in the second stage, while in IIWM it is executed in the seventh one). Hence, in a dynamic scheduling scenario where tasks can be added at runtime, IIWM may suffer from the starvation problem, as such tasks may experiment an indefinite delay as far as new tasks with a higher memory weight
are provided to the scheduler. Nevertheless, in the proposed work we dealt with a static scheduling problem, where all tasks are known in advance and the task-set is not modifiable at runtime.

| Strategy | Task-scheduling plan | Number of stages | Time (min.) | Peak disk usage (MB) | Writes duration (min.) | On-disk usage (MB) |
|----------|----------------------|------------------|-------------|----------------------|-----------------------|---------------------|
| Full-Parallel | $\{(t_1, t_1 \parallel t_2 \parallel t_3 \parallel t_4),\,$ $\{t_5 \parallel t_6 \parallel t_8 \parallel t_9 \parallel t_{10} \parallel t_{11} \parallel t_{12}\},\,$ $\{t_{12} \parallel t_{14} \parallel t_{15} \parallel t_{16} \parallel t_{17} \parallel t_{18} \parallel t_{19} \parallel t_{20}\},\,$ $\{t_{21} \parallel t_{22} \parallel t_{24} \parallel t_{25}\},\,$ $\{t_{26}\}\}$ | 7 | 29.42 ± 1.88 | 27095.837 | 20.6 | 10593.79 |
| IIWM | $\{(t_0), (t_4 \parallel t_2), (t_{11} \parallel t_7), (t_8 \parallel t_5),\,$ $\{t_{15} \parallel t_{10} \parallel t_{9} \parallel t_{10}\}, (t_{18}),\,$ $\{t_{17} \parallel t_{12}, (t_6 \parallel t_5), (t_{14}),\,$ $\{t_{13} \parallel t_{20} \parallel t_{19}, t_{22}\},\,$ $\{t_{23} \parallel t_{24}, (t_{25}), (t_{26})\}$ | 15 | 22.68 ± 1.65 | 304.5 | 3.6 | 60.82 |

Table 12. Scheduling plan and statistics about execution times and disk usage with 9.5 GB of RAM.

4.2. Real case study

In order to assess the performance of the proposed approach against a real case study, we used a data mining workflow [27] that implements a model selection strategy for the classification of an unlabelled dataset. Figure 9 shows a representation of the workflow designed by the visual language VL4Cloud [28]. A training set is divided into $n$ partitions and $k$ classification algorithms are fitted on each partition for generating $k \times n$ classification models. The $k \times n$ fitted models are evaluated by a model selector on a test set to choose the best model. Afterwards, the $n$ predictors use the best model to generate $n$ classified datasets. The following $k$ classification algorithms provided by the MLlib library were used: Decision Tree with C4.5 algorithm, Support Vector Machines (SVM), and Naive Bayes. The training set, test set and unlabelled dataset provided as input for the workflow have been generated from the Physical Unclonable Functions (PUFs) [29] simulation through a $n$-fold-cross strategy. In this scenario, IIWM can be used to optimize the data processing phase regarding the execution of the $k \times n$ classification algorithms (estimators first, transformers then) concurrently. The other phases, such as data acquisition and partitioning, are out of our interest. The red box in Figure 9 shows the tasks of the workflow that will be analyzed.

Figure 9. Ensemble learning workflow.

Figure 10 shows disk occupancy over time with 14 GB of RAM. Also in this case, IIWM avoided disk writes, while Full-Parallel registered a high level of disk usage. In particular, during the training phase, the parallel execution of the $k \times n$ models (with $k = 3$ and $n = 5$) saturates memory with 15 concurrent tasks and generates disk writes up to 124 GB.
The results are detailed in Table 13, which shows a boost in execution time of almost $1.66 \times (p_{imp})$ and a 40% time reduction ($m_{imp}$) with respect to Full-Parallel.

| Strategy | Task-scheduling plan | Number of stages | Time (min.) | Peak disk usage (MB) | Writes duration (min.) | On-disk usage (MB) |
|----------|----------------------|------------------|-------------|----------------------|-----------------------|-------------------|
| Full-Parallel | $(t_0 || t_2 || t_4 || t_6 || t_8 || t_{10} || t_{12} || t_{14} || t_{16} || t_{18} || t_{20} || t_{22} || t_{24} || t_{26} || t_{28} || t_{30})$ | 2 | 11.42 ± 0.27 | 124730.874 | 9.6 | 54443.186 |
| IIWM | $(t_{10} || t_{12} || t_{14} || t_{16} || t_{20} || t_{22} || t_{24} || t_{26} || t_{28} || t_{30})$, $(t_{18} || t_{0} || t_{2} || t_{4} || t_{6} || t_{8} || t_{10} || t_{12} || t_{14} || t_{16} || t_{18} || t_{20} || t_{22} || t_{24} || t_{26} || t_{28} || t_{30} || t_{32} || t_{34} || t_{36} || t_{38})$ | 3 | 6.88 ± 0.1 | 0 | 0 | 0 |

Table 13. Scheduling plan and statistics about execution times and disk usage with 14 GB of RAM.

The general trends varying the amount of available resources are also confirmed with respect to the previous examples, as shown in Figure 11.

Figure 11. Average peak disk usage and execution time, varying the size of available memory.

5. Conclusions and Final Remarks

Nowadays, data-intensive workflows are widely used in several application domains, such as bioinformatics, astronomy, and engineering. This paper introduced and evaluated a system, named Intelligent In-memory Workflow Manager (IIWM), aimed at optimizing the in-memory execution of data-intensive workflows on high-performance computing systems. Experimental results suggest that by jointly using a machine learning model for performance estimation and a suitable scheduling strategy, the execution of data-intensive workflows can be significantly improved with respect to state-of-the-art blind strategies. In particular, the main benefits of IIWM resulted when has been applied to workflows having a high level of parallelism. In this case a significant reduction of memory saturation has been obtained. Therefore it can be used effectively when multiple tasks have to be executed on the same computing node, for example when they need to be run on multiple immovable datasets located on a single node or due to other hardware constraints. In these cases, an uninformed scheduling strategy will likely exceed the available memory, causing disk writes and therefore a
drop in performance. The proposed approach also showed to be a very suitable solution in scenarios characterized by a limited amount of memory reserved for execution, thus finding possible applications in data-intensive IoT workflows, where data processing is performed on constrained devices located at the network edge.

IIWM has been evaluated against different scenarios concerning both synthetic and real data mining workflows, using Apache Spark as a testbed. Specifically, by accurately predicting resources used at runtime, our approach achieved up to 31% and 40% reduction of makespan and a performance improvement up to 1.45x and 1.66x for the synthetic workflows and the real case study respectively.

In future work additional aspects of performance estimation will be investigated. For example, the IIWM prediction model can be extended also to consider other common stages in workflows besides data analysis, such as data acquisition, integration and reduction, and other information about tasks, input data, and hardware platform features can be exploited in the scheduling strategy.

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