Towards Predicting the Runtime of Iterative Analytics with PREDIcT

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ABSTRACT

Machine learning algorithms are widely used today for analytical tasks such as data cleaning, data categorization, or data filtering. At the same time, the rise of social media motivates recent uptake in large scale graph processing. Both categories of algorithms are dominated by iterative subtasks, i.e., processing steps which are executed repetitively until a convergence condition is met. Optimizing cluster resource allocations among multiple workloads of iterative algorithms motivates the need for estimating their runtime, which in turn requires: i) predicting the number of iterations, and ii) predicting the processing time of each iteration. As both parameters depend on the characteristics of the dataset and on the convergence function, estimating their values before execution is difficult.

This paper proposes PREDIcT, an experimental methodology for predicting the runtime of iterative algorithms. PREDIcT uses sample runs for capturing the algorithm’s convergence trend and per-iteration key input features that are well correlated with the actual processing requirements of the complete input dataset. Using this combination of characteristics we predict the runtime of iterative algorithms, including algorithms with very different runtime patterns among subsequent iterations. Our experimental evaluation of multiple algorithms on scale-free graphs shows a relative prediction error of 10%-30% for predicting runtime, including algorithms with up to 100x runtime variability among consecutive iterations.

1. INTRODUCTION

Today’s data management requirements are more complex than ever, going well beyond the traditional roll-up or drill-down operators proposed in OLAP systems [10]. Analytical tasks often include machine learning or graph mining algorithms [22, 28] executed on large input datasets. For instance, Facebook uses machine learning to order stories in the news feed (i.e., ranking), or to group users with similar interests together (i.e., clustering). Similarly, LinkedIn uses large scale graph processing to offer customized statistics to users (e.g., total number of professionals reachable within a few hops). These algorithms are often iterative: one or more processing steps are executed repetitively until a convergence condition is met [22].

Execution of iterative algorithms on large datasets motivates the need for predicting their resource requirements and runtime. Runtime estimates for such algorithms are a pre-requisite for optimizing cluster resource allocations in a similar manner as query cost estimates are a pre-requisite for DBMS optimizers. Operational costs associated to large cluster deployments are high, hence enterprises aim to maximize their utilization. In particular, schedulers and resource managers are used to optimize resource provisioning, reduce over-provisioning, while at the same time satisfying user contracted Service Level Agreements (SLAs) [40, 38].

Additionally, runtime prediction is a very useful mechanism for answering feasibility analysis questions: ‘Given a cluster deployment and a workload of iterative algorithms, is it feasible to execute the workload on an input dataset while guaranteeing user specified SLAs?’

Predicting the runtime of iterative algorithms poses two main challenges: i) predicting the number of iterations, and ii) predicting the runtime of each iteration. Additionally to the algorithm’s semantics, both iterations and per iteration runtime depend on the characteristics of the input dataset, and the intermediate results of all prior iterations. On one hand, the number of iterations depends on how fast the algorithm converges. Convergence is typically given by a distance metric that measures incremental updates between consecutive iterations. Unfortunately, an accurate closed-form formula cannot be built in advance, before materializing all intermediate results. On the other hand, the runtime of a given iteration may vary widely compared with the subsequent iterations according to the algorithm’s semantics and as a function of the iteration’s current working set [14]: Due to sparse computation, updating an element of the intermediate result may have an immediate impact only on a limited number of other elements (e.g., propagating the smallest vertex identifier in a graph structure using only point to point messages among neighboring elements). Hence, estimating the time requirements, or alternatively, the size of the working sets of each iteration before execution is difficult.

Existing Approaches: Prior work on estimating the runtime or the progress of analytical queries in DBMS (e.g., [8, 16, 2, 13, 27]) or more recent MapReduce systems (e.g., [21, 20, 31, 34]) do not address the problem of predicting the runtime of analytical workflows that include iterative algorithms. For certain algorithms theoretical bounds for the number of iterations were defined (e.g., [24, 22, 18, 4]). However, due to simplifying assumptions on the characteristics of the input dataset theoretical bounds are typically
sampling technique from one iteration to the next decreases below a user-defined threshold \( \tau \). We define \( \tau \) as the number of iterations when executing PageRank on sample graphs. The number of iterations is maintained as on the complete graph. During the sample run key input features are captured and used later as a basis for prediction. However, as some algorithm parameters are tuned to a certain dataset size, a sampling run cannot simply execute the same algorithm with the same parameters on a smaller dataset. We first have to identify the parameters that need to be scaled and then apply the transform function to obtain the suitable values for the sample dataset size. One such parameter is the convergence threshold used by PageRank [32] and other algorithms. We illustrate why we need to scale the threshold value with an example.

Example: PageRank is an iterative algorithm that computes the rank of all vertices of a directed graph by associating to each vertex a rank value that is proportional to the number of vertices, edges or in/out node degree ratios of samples S1-S3 as compared with G, the average PageRank change on the samples is not the same with the corresponding PageRank change on G. Computing the average PageRank change corresponding to the first iteration results in: \( \delta_{S1} = 3d/16 \), \( \delta_{S2} = d/8 \), \( \delta_{S3} = d/8 \), and \( \delta_G = d/16 \), where \( d=0.85 \) is the damping factor (for deriving these values please see Equation 1). For this example, for a convergence threshold \( \tau = d/16 \) the actual run converges after one iteration, whereas all sample runs continue execution. By applying the transformation \( T = (\tau_S = \tau_G \times 2) \) during the sample run on samples S2 or S3, the same number of iterations is maintained as on the complete graph. Hence, only by combining a transform function with a sampling technique (which maintains certain properties of the graph G: e.g., diameter), invariants can be preserved for the number of iterations.

PREDIcT proposes the methodology for providing transformation functions on a class of iterative algorithms that are operating on homogeneous graph structures, and have a global convergence condition: i.e., computing an aggregate at the graph level. Examples of such algorithms include: ranking (e.g., PageRank, top-K ranking), clustering on graphs (e.g., semi-clustering) or graph processing (e.g., neighborhood estimation). PREDIcT provides a set of default rules for choosing the transformations that work for a representative class of algorithms. At the same time, users can plug in their own set of transformations based on domain knowledge, if the semantics of the algorithm are not already captured by the default rules. Considering that a representative set of iterative, machine learning algorithms are typically executed repetitively on different input datasets [10, 22, 28], and that the space of possible algorithms is not prohibitive, deriving such a set of customized transformations is also practical and worthwhile.

As Figure 1 shows, after key input features (including iterations) are profiled during the sample run and extrapolated to the scale of the complete dataset, a cost model is required for translating key input features into runtime estimates. For this purpose, PREDIcT introduces a framework for building customizable cost models for network intensive iterative algorithms executing using the Bulk Synchronous Parallel (BSP) [36] execution model, in particular Apache Giraph implementation1. Our framework identifies a set of key input features that are effective for network intensive algorithms, it includes them into a pool of features, and then uses a feature selection mechanism for choosing the features that will be used in the cost model. The cost model is trained on the set of input features profiled during the sample run, and additionally, on the set of input features of prior actual runs of the algorithm on different input datasets (if such runs exist). Such historical runs are typically available for analytical applications that are executed repetitively over newly arriving data sets. Examples include: ranking, clustering, social media analytics.

1http://incubator.apache.org/giraph/
is the first paper that proposes the transform function for maintaining invariants among the sample run and the actual run in the context of iterative algorithms and demonstrates its practical applicability for prediction. We note that the methodology we propose for estimating key input features is conceptually not tied to Giraph, and hence, could be used as a reference for other execution models operating on graph structures such as GraphLab [28] or Grace [39]. To this end identifying the key input features that significantly affect the runtime performance of these engines is required. For some iterative algorithms (mappable to a dual problem operating on graphs) our approach for estimating iterations can be applied even to non-BSP frameworks like Spark [41] and Mahout [3].

This paper makes the following contributions:

- It proposes PREDIcT, an experimental methodology for predicting the runtime of network intensive iterative algorithms. PREDIcT was designed to predict not only the number of iterations, but also the key input features of each iteration, which makes it applicable for algorithms with very different runtime patterns among subsequent iterations.

- It proposes a framework for building customized cost models for iterative algorithms executing on top of Giraph. Although the complete set of key features and the cost model per se will vary from one BSP implementation to another (in a similar fashion as DBMS cost models vary from one DBMS vendor to another), our proposed methodology is generic. Hence, it can be used as a reference when building similar cost models on alternative BSP implementations.

- It evaluates PREDIcT on a representative set of algorithms using real datasets, showing PREDIcT’s practicality over analytical upper bounds. For a 10% sample, the relative errors for estimating key input features range in between 5%-20% , while the errors for estimating the runtime range in between 10%-30%, including algorithms with up to 100x runtime variability among consecutive iterations.

2. RELATED WORK AND BACKGROUND

In this section we present related work on runtime prediction techniques applied in the DBMS, and prior research on algorithmic approaches used for providing analytical upper bounds for the number of iterations an algorithm requires to converge. Then we introduce the key concepts of the BSP processing model [36] which is inherently iterative and emerges as the new paradigm for executing graph processing tasks at large-scale [30].

2.1 Prediction and Iterative Processing

Prior work on iterative algorithms mainly focuses on providing theoretical bounds for the number of iterations an algorithm requires to converge (e.g., [24, 22, 18]) or worst case time complexity (e.g., [4]). These parameters, however, are not sufficient for providing wall time estimates because of the following reasons: i) As simplifying assumptions about the characteristics of the input dataset are made, theoretical bounds on the number of iterations are typically loose [24, 4]. This problem is further exacerbated for a category of iterative algorithms executing sparse computation, where the processing requirements of any arbitrary iteration vary a lot as compared with subsequent/prior iterations [14, 28]. For such algorithms, per iteration worst case time complexities are typically impractical. ii) Per iteration processing wall times cannot be captured solely by a complexity formula. System level resource requirements (i.e., CPU, networking, I/O), critical path modeling and a cost model are additionally required for modeling runtime.

Estimating the runtime execution of analytical workloads was heavily studied in the DBMS context from multiple angles: initial runtime predictors [16, 2, 13, 34], progress estimators [8, 29, 31] or self-tuning systems [21, 20]. None of these approaches, however, is applicable for iterative pipelines, where particular operators (i.e., algorithms) are executed repetitively until a convergence condition is met. In contrast with algorithmic approaches, prediction approaches proposed in the DBMS context account for system level resource requirements, and use a cost model (either analytical, based on black box modeling or a hybrid) for translating them into actual runtime. For instance, [16] proposes a technique to predict query performance using a cost model based on machine learning that correlates queries with similar input features together. Other approaches [20] propose a hybrid of analytical models, simulation and controlled black box models for estimating the runtime of analytical queries that takes as input features: data characteristics, processing cost functions and system configuration parameters.

Although adaptive query processing techniques such as [8, 29, 31] can be used for updating the cost model at runtime (i.e., the processing cost factors), they cannot estimate or calibrate key input features in the context of iterative algorithms because: i) The processing requirements of consecutive iterations may vary a lot (e.g., connected components), and they are not necessarily monotonic as more iterations are executed. ii) Stopping condition cannot be evaluated before an iteration is completed. In DBMS terminology, iterative processing can be interpreted as a join aggregate query among a relation that does not change (i.e., graph structure) and a relation that gets updated in each iteration (i.e., the propagation function). Hence, building accurate statistics on the relation that is updated is not possible before execution. For the same reason above adaptive techniques for calibrating statistics at runtime [35, 12] are not applicable.

Iterative execution was also analyzed in the context of recursive query processing. In particular, multiple research efforts [5, 1, 6] discuss execution strategies (i.e., top-down versus bottom-up) with the goal of performance optimization. HaLoop [7] caches invariant input datasets among subsequent iterations when executing iterative algorithms on MapReduce execution model. Ewen et al. [14] optimize execution of incremental iterations that are characterized by few localized updates, in contrast with bulk iterations, that always update the complete dataset. Although highly related to our research, as performance optimization determines the runtime of the queries, all the aforementioned techniques are complementary to the goal of estimating the runtime of the algorithms. PREDIcT can be combined with previous work to perform cost-based optimizations when executing workflows of iterative algorithms.

In the context of resource allocation, both FLEX [40] and ARIA [38] require a prediction mechanism to find the optimal allocation of resources (i.e., number of map/reduce slots) and an ordering of jobs that satisfy user specified SLAs (e.g., deadlines). PREDIcT can be used in conjunction with such resource allocators to estimate the runtime of the algorithm for a given slot configuration.

2.2 The BSP Processing Model

Any algorithm executed on top of BSP is inherently iterative: It runs in a succession of supersteps (i.e., iterations) until a termination condition is satisfied. Each superstep
3.1 Modeling Assumptions

In our proposed prediction methodology we make the following assumptions:

- All the iterative algorithms we analyze in this paper are guaranteed to converge.
- Input datasets are graphs, and are amenable to sampling. The sample graph maintains its properties similar or proportional with those of the original graph.
- Both the sample-run and the actual-run use the same execution framework (i.e., Giraph) and system configuration parameters.
- All the worker nodes have uniform resource allocations, hence processing costs among different workers are similar.
- The dominating part of the runtime of the algorithms is networking: i.e., sending/receiving messages from other vertices.

Such assumptions hold for a class of algorithms implemented on top of BSP which are dominated by networking processing costs: Some of them have very short per vertex computation (e.g., PageRank), while some others have larger per vertex computation cost which is largely proportional with the size and the number of messages received (sent) from (to) the neighboring nodes (e.g., semi-clustering [30], top-k ranking [23]).

3.2 Sample Run

The sample run is the preliminary phase of the prediction approach that executes the algorithm on the sample dataset. As explained in section 1.1, two set of transformations characterize the execution of the algorithm during the sample run: the sampling technique adopted and the transform function. Once the set of transformations is determined, the algorithm is executed on the sample. During the sample run, per iteration key input features are profiled and used later in the prediction phase as a basis for estimating the corresponding features of the actual run.

3.2.1 Sampling technique

The sampling technique adopted has to maintain key properties of the sample graph similar or proportional with those of the original graph: Examples of such properties include in/out degree proportionality, effective diameter, clustering coefficient. Hence, we adopt similar sampling techniques with those proposed by Leskovec et al. [25], which show that such graph properties on the sample can be maintained similar to those on the complete graph.

Random Jump: We choose Random Jump (RJ) from the set of sampling methods proposed in [25], because it is the sampling method that has no risks of getting stuck into an isolated region of the graph, while maintaining comparable scores for all key properties of the graph with Random Walk and Forest Fire (as shown in Table 1 of [25]). RJ picks a starting seed vertex uniformly at random from all the input vertices. Then, at each sampling step an outgoing edge of the current vertex is picked uniformly at random and the current vertex is updated with the destination vertex of the picked edge. With a probability $p$ the current walk is ended and a new random walk is started from a new seed vertex chosen at random. The process continues until the number of vertices picked satisfies the sampling ratio. Such a sampling technique has the property of maintaining connectivity within a walk. Random jump achieves connectivity among multiple walks by returning to already visited vertices on
different edges. Returning to already visited nodes also improves the probability of preserving the in/out node degree proportionality.

**Biased Random Jump**: Based on the observation that convergence of multiple iterative algorithms we analyze is inherently dictated by high out-degree vertices (e.g., PageRank, top-k ranking, semi-clustering), we propose Biased Random Jump (BRJ), a variation of Random Jump. BRJ is biased towards high out degree vertices: Compared with RJ, BRJ picks $k$ seed vertices from the graph in decreasing order of their out-degree instead of using arbitrary seed vertices. Then, for each new random walk performed a starting vertex is picked uniformly at random from the set of seed vertices. The intuition of BRJ is to prioritize sampling towards the “core of the network”, that include vertices with high out degrees. Biased random jump trades-off sampling uniformity for improved connectivity: By starting random walks from highly connected nodes (i.e., hub nodes), BRJ has a higher probability of maintaining connectivity among sampled walks than RJ, where jumps to any arbitrary nodes are possible. We empirically find that BRJ has higher accuracy than RJ in maintaining key properties of the graph (such as connectivity), especially at small sampling ratios (the sampling ratio proposed for RJ in [25] is 25%). Hence, BRJ is used as our default sampling mechanism.

### 3.2.2 Transform function

The transform function $T$ is formally described by two pairs of adjustments: $T = (Conf_{fs} => Conf_{fc}, ConEv_{s} => ConEv_{c})$, where $Conf_{fs} => Conf_{fc}$ denotes configuration parameter mappings, while $ConEv_{s} => ConEv_{c}$ denotes convergence parameter mappings. For instance, the transformation $T = (dS = dC, rS = rC \times \frac{1}{2})$ for PageRank algorithm denotes: preserve the damping factor value on the sample run equal with the corresponding value of the actual run, and scale the convergence threshold.

While the transform function requires domain knowledge about the algorithm semantics, we provide a default rule which works for a set of representative algorithms operating on graphs and can be used as a reference when choosing alternative transformations: For the case that the convergence threshold is tuned to size of the input dataset (i.e., convergence is determined by an absolute aggregated value, as for PageRank): $T_{default} = (ID_{Conf}, rS = rG \times \frac{1}{2})$, while for the case that convergence threshold is not tuned to the size of the input dataset (i.e., convergence is determined by a relative aggregated value or a ratio that is maintained constant on a proportionally smaller dataset, as for top-k ranking): $T_{default} = (ID_{Conf}, rS = rG)$. Specifically, we maintain all the configuration parameters of the algorithm during the sample run (identity function over the configuration space) and we scale or maintain the convergence threshold for the sample run.

### 3.3 Key input features

We identify the key input features for the Giraph execution model based on a mix of domain knowledge and experimentation. Table 1 shows the set of key input features we identified for modeling the runtime of network intensive iterative algorithms. The number of iterations is not extrapolated, as the transform function targets to preserve the number of iterations during the sample run.

In order to understand the selection of key input features, consider Figure 3 that illustrates the execution phases of an arbitrary iteration of an iterative algorithm that uses BSP. Each worker executes three phases: compute, messaging, and synchronization, as explained in section 2.2.

**Figure 3**: BSP execution phases of an arbitrary iteration.

**Compute phase**: In this phase the user defined function that implements the semantics of the iterative algorithm is executed for every vertex of the input graph. For a large category of network intensive algorithms the cost of local, per vertex computation (executing the semantics of the algorithm) can be approximated by a constant cost factor, while the cost of initiating messages to neighboring nodes is proportional with the number of messages each vertex sends. Hence, the compute time of each worker (which has multiple vertices allocated to it) is proportional with the total number of active vertices (i.e., executing actual work), and the number of messages each worker sends.

**Messaging phase**: During this phase, messages are sent over the network and added into the memory of the destination nodes. Some BSP implementations can spill messages to disk. Hence, the runtime of this phase is proportional with the number of messages, their sizes, and the number and sizes of messages spilled to disk (if spilling occurs).

**Synchronization phase**: The synchronization time of a worker w.r.t. the worker on the critical path (the worker finishing its assignments the last) depends on the partitioning scheme adopted, which in turn may result in work allocation skew among workers. Instead of trying to model the synchronization time among workers explicitly, we model it implicitly by identifying the worker on the critical path, which has close to zero synchronization time.

![Table 1: Key Input Features](image)

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
<th>Extrapolation</th>
</tr>
</thead>
<tbody>
<tr>
<td>ActVert</td>
<td>Number of active vertices</td>
<td>yes</td>
</tr>
<tr>
<td>TotVert</td>
<td>Number of total vertices</td>
<td>yes</td>
</tr>
<tr>
<td>LocMsg</td>
<td>Number of local messages</td>
<td>yes</td>
</tr>
<tr>
<td>RemMsg</td>
<td>Number of remote messages</td>
<td>yes</td>
</tr>
<tr>
<td>LocMsgSize</td>
<td>Size of local messages</td>
<td>yes</td>
</tr>
<tr>
<td>RemMsgSize</td>
<td>Size of remote messages</td>
<td>yes</td>
</tr>
<tr>
<td>AvgMsgSize</td>
<td>Average message size</td>
<td>no</td>
</tr>
<tr>
<td>NumIter</td>
<td>Number of iterations</td>
<td>no</td>
</tr>
</tbody>
</table>

While the set of features illustrated in Table 1 is effective for network intensive algorithms, they should not be interpreted as complete. Given the generality of selecting input features into the cost model, our proposed methodology can be extended to include additional key input features in the pool of candidate input features. For instance, counters corresponding to spilling messages to disk during the messaging phase shall be also considered if spilling occurs. Giraph currently does not support spilling of messages to disk, hence such features were not required in our experiments.

### 3.4 Prediction

There are two phases in the prediction process: i) extrapolation of key input features profiled during the sample run; and ii) costing extrapolated features into runtime using a cost model.
Extrapolator: As shown in Figure 1, in the first prediction phase an extrapolator is used to scale-up input features profiled during the sample run. The input metrics that are used in the extrapolation phase are the number of edges and the number of vertices of the sample graph S, and the corresponding number of edges and vertices of the complete graph G. We use two extrapolation factors: i) for features that primarily depend on the number of vertices (e.g., ActVert), we extrapolate with a scaling factor on vertices: i.e., \( e_V = \frac{|V_G|}{|V_S|} \); ii) for features that depend both on the number of input nodes and edges (e.g., message counts depend on how many outbound edges a vertex has) we extrapolate with a scaling factor on edges: i.e., \( e_E = \frac{|E_G|}{|E_S|} \). Note that not all key input features require extrapolation: e.g., number of iterations is preserved during the sample run. Extrapolation of input features is done at the granularity of iterations: i.e., the input features of an arbitrary iteration of the sample run are extrapolated and then used to predict the runtime of the corresponding iteration of the actual run.

Customizable Cost Model: In the second phase a cost model is used to translate extrapolated input features into runtime. The cost model is invoked multiple times, on extrapolated input features corresponding to each iteration of the sample run. Hence, the number of iterations is used implicitly rather than explicitly in prediction.

Based on the processing model breakdown presented in sub-section 3.3, we propose a cost modeling technique for network intensive algorithms that uses multivariate linear regression to fit a set of key input features into per iteration runtime. Formally, given a set of input features \( X_1, \ldots, X_k \), and one output feature \( Y \) (i.e., per iteration runtime), the model has the functional form: \( f(X_1, \ldots, X_k) = c_1 X_1 + c_2 X_2 + \ldots + c_k X_k + r \) where \( c_i \) are the coefficients and \( r \) is the residual value. A modeling approach based on a fixed functional form was chosen for several reasons: i) For network intensive algorithms, each phase of the Giraph BSP execution model except the synchronization phase can be approximated by a fixed functional form (multivariate regression). The synchronization phase is modeled implicitly, as explained in section 3.3. ii) A fixed functional form can be used for prediction on input feature ranges that are outside of the training boundaries (e.g., train on sample run, test on actual run). In fact the coefficients of the multivariate linear model can be interpreted as the “cost values” corresponding to each input feature.

We use the set of features presented in Table 1 as candidates in the cost model. Customization of the cost model for a given iterative algorithm is done by selecting the actual input features that have a high impact on the response variable \( Y \), and yield a good fitting coefficient for the resulting model. In particular, selecting the actual key features from the above pool of features is based on an sequential forward selection mechanism [19] that selects the features that yield the best prediction accuracy on the training data.

Cost Model Extensions: For the cases where the compute phase is not linearly proportional with the number of active vertices, and the number and size of messages, our proposed cost model is extensible as follows: i) The compute phase and messaging phase are separately profiled; ii) A similar approach as above is used to model the messaging phase; iii) A non linear approach is used to model the compute function (e.g., decision trees). For this purpose, MART scale [27] can be used, as it was designed to be accurate even on key input features outside of the training boundaries. While such an extension is worthwhile, it is beyond the scope of this paper.

Modeling the Critical Path: In the BSP processing model the runtime of one iteration is given by the worker on the critical path (i.e., finishing its working assignments last). In a homogeneous environment where each worker has the same share of system resources, the worker on the critical path is the worker processing the largest part of the input graph. For a vertex centric partitioning scheme, non-uniform allocations may exist if some vertices are better connected than others, which in turn result into larger processing requirements. This observation holds for network intensive algorithms, where the number of outgoing edges determine the messaging requirements of the vertex, and in turn, the runtime. We adopt the following methodology for finding the worker on the critical path: For a given partitioning scheme of vertices to partitions, and a mapping of partitions to workers, the total number of outbound edges for each worker is computed. The worker with the largest number of outbound edges is considered to be on the critical path. Such a method for finding the latest worker can be piggybacked in the initialization phase of the algorithm, in the read phase, and can be exploited for prediction just before the algorithm starts its effective execution in the supervision phase.

Training Methodology: For training the cost model we use both sample runs, and additionally, historical actual runs of the algorithm on different input datasets (if such runs exist). Such a training scenario is applicable for the class of algorithms we target to address in the paper, as the underlying cost functions corresponding to each input feature: i.e., cost of sending/receiving messages, or the cost of executing the compute function, are similar when executing the same algorithm on different input datasets. Hence, once a cost model is built, it can be reused for predicting the runtime of the algorithm on different input datasets.

The cost model is trained at the granularity of iterations: Key input features are profiled and maintained on a per-worker basis for each iteration of the algorithm. Specifically, the code path of each BSP worker was instrumented with counters for all the input features potentially required in the cost model. Then, all counters are used to train the model.

3.5 Limitations

PREDICT was designed for a class of iterative algorithms that are operating on homogeneous graph structures and use a global convergence condition: i.e., computing an aggregate at the graph level (i.e., an average, a total, a ratio of updates). Algorithms for which convergence is highly influenced by the local state of any arbitrary vertex of the graph are not amenable to sampling, and hence, PREDICT methodology cannot be used for these cases. Similarly, PREDICT cannot be used on degenerated graph structures where maintaining key graph properties in a sample graph is not possible. In analogy with traditional DBMS, we cannot use a sample of a dataset to estimate outliers, but we can use it to produce average values. We note that the sampling requirements in our case are more relaxed, as we do not use sampling to approximate results. Instead, sampling is used as a mechanism to approximate the processing characteristics of the actual run. Examples of algorithms where our methodology is not applicable: collaborative filtering (heterogeneous graphs with two entity types: e.g., users and movies) or simulating advertisements in social networks [23] (the decision to further propagate an advertisement depends on the local interest of the node receiving the advertisement (i.e., his interest list)). Examples of datasets where our methodology is not applicable: e.g., degenerated, non uniform graph structures, e.g., lists.
4. END-TO-END USE CASES

In this section we show how to apply PREDiCT’s proposed methodology for predicting per iteration key input features for two categories of network intensive algorithms introduced in section 3: i.e., i) constant vs. ii) variable runtime among subsequent iterations. For the second category of algorithms, we consider two sub-cases: a) variable per iteration runtime caused by different message size requirements among iterations; b) variable per iteration runtime caused by a different number of messages sent among iterations. As end-to-end use cases, we choose PageRank as a representative algorithm for category i), semi-clustering is chosen for category ii) a), and top-k ranking is chosen for category ii) b).

4.1 PageRank

PageRank is an iterative algorithm proposed in the context of the Web graph, where vertices are web pages and edges are references from one page to the other. Conceptually, PageRank associates to each vertex a rank value proportional with the number of inbound links from the other vertices, and their corresponding PageRank values. In order to understand how is the rank transfer between vertices affecting the number of iterations, we introduce the formula used for computing PageRank [32]:

\[
PR(p_i)_{it} = \frac{1 - d}{N} + d \sum_{j \in M(p_i)} \frac{PR(p_i)_{it-1}}{L(p_j)}
\]

where \(PR(p_i)\) is the PageRank of the vertex \(p_i\), \(N\) is the total number of vertices, \(d\) is the damping factor (typically set to 0.85), \(M(p_i)\) are the vertices for which the rank is computed, \(L(p_j)\) is the set of vertices that link to \(p_j\), and \(PR(p_i)\) is the number of outbound edges of vertex \(p_i\). Before the iterative process starts, the initial rank value of each vertex is initialized to \(1/N\).

**Convergence**: PageRank algorithm converges when the average delta change of PageRank value per vertex is below a user defined threshold \(\tau\). Formally, the delta change of PageRank for an arbitrary vertex \(p_i\) is defined as: \(\delta_i = \frac{|PR(p_i)_{it} - PR(p_i)_{it-1}|}{PR(p_i)_{it-1}}\), and the average delta change of PageRank for any arbitrary vertex: \(\frac{1}{N} \sum \delta_i\). It can be shown that for a directed acyclic graph the maximum number of iterations required for PageRank to converge to a delta change of zero (i.e., \(\tau = 0\)) is the diameter of the graph \(D\). For real graphs, however, the DAG assumption does not hold as cycles between vertices are typical. Therefore, an additional number of iterations are required for the algorithm to converge to an average delta change threshold \(\tau > 0\).

**Sampling Requirements**: In order to take a representative sample that can maintain the number of iterations of the actual run similar with that of the sample run we make the following observations: i) Maintaining connectivity is crucial in propagating the PageRank transfer among graph vertices. Therefore, the sampling technique should maintain the connectivity among sampled vertices (i.e., the sample should not degenerate into multiple isolated sub-graphs). ii) The PageRank delta change per vertex depends on the number of incoming and outgoing edges. The sample should ideally maintain the in/out node degree ratio similar with the corresponding ratio on the original graph. iii) The diameter of the graph determines the number of iterations required to propagate the PageRank transfer among vertices located at the graph boundaries. Hence, ideally the diameter of the sample graph shall be similar with the diameter of the original graph. In practice, maintaining the effective diameter of the graph is more feasible, i.e., the shortest distance in which 90% of all connected pairs of nodes can reach each other.

**Transform function**: Consider the example introduced in Figure 2: It can be shown that for any arbitrary iteration, the average delta change of PageRank on graph S3 can be maintained in pair with the average delta change of PageRank on graph G by the following transform function: \(T = (1D_{Conf}, TS_{T})\), where \(Conf = (d, sr)\), and \(sr\) is the sampling ratio.

For a better understanding of transformation \(T\), we compute the PageRank of vertex 5 on graph G and then on graph S3 for the first iteration. On graph G, the PageRank of vertex 5 is given by: \((1-d)/(N/2)+d/(2*N)\). On graph S3: \((1-d)/(N/2)+d/(2* (N/2))\). We observe that the PageRank value of node 5 on the sample S3 is twice of the corresponding PageRank value on graph G (equal with the inverse of the sampling ratio), as the sample maintains the structure of the original graph (i.e., in/out node degree ratio and diameter). Similarly, it can be shown that the average delta PageRank change on the sample graph S3 is twice of \(G\). Hence, the transform function \(T\) maintains invariants for the number of iterations. In real graphs such symmetric structures cannot be assumed. Still, we can use such transformations as a basis for an heuristic approach that shows good results in practice.

4.2 Semi-Clustering

Semi-clustering is an iterative algorithm popular in social networks as it aims to find groups of people who interact frequently with each other and less frequently with others. A particularity of semi-clustering as compared with the other clustering algorithms is that a vertex can belong to more than one cluster. We adopt the parallel semi-clustering algorithm as described in [30]. The input is an undirected weighted graph while the output is an undirected graph where each vertex holds a maximum number of \(C_{\text{max}}\) semi-clusters it belongs to. Each semi-cluster has associated a score value:

\[
S_c = \frac{I_c - f_B * B_c}{V_c(V_c - 1)}/2
\]

, where \(I_c\) is the sum of the weights of all internal edges of the semi-cluster, \(B_c\) is the sum of the weights of all boundary edges, \(f_B\) is the boundary edge factor (i.e., \(0 < f_B < 1\), a user defined parameter) which penalizes the total score value, and \(V_c\) is the number of vertices in the semi-cluster. As it can be noticed, the score is normalized to the number of edges in a clique of size \(V_c\) such that large semi-clusters are not favored. The maximum number of vertices in a semi-cluster is bounded to a user settable parameter \(V_{\text{max}}\). After the set of best semi-clusters of each vertex are found, they are aggregated into a global list of best semi-clusters.

**Convergence**: The algorithm runs in iterations: In the first iteration, each vertex adds itself to a semi-cluster of size one which is then sent to all of its neighbors. In the following iterations: i) Each vertex \(V\) iterates over the semi-clusters sent to it in the previous iteration. If a semi-cluster \(sc\) does not contain vertex \(V\) and \(V_c < V_{\text{max}}\), then \(V\) is added to \(sc\) to form \(sc'\). ii) The semi-clusters \(sc\) that were sent to \(V\) in the previous iteration together with the newly formed semi-clusters \(sc'\) are sorted by score and the best \(S_{\text{max}}\) are sent out to \(V\)’s neighbors. iii) Vertex \(V\) updates its list of \(C_{\text{max}}\) best semi-clusters with the newly received / formed semi-clusters (i.e., the semi-clusters from the set: \(sc, sc'\)) that contain \(V\).

The algorithm converges when the list of all semi-clusters
that every vertex maintains stop changing. As such a stopping condition requires a large number of iterations an alternative stopping condition that considers the proportion of semi-cluster updates is more practical: More precisely: \( \frac{\text{updatedClusters}}{\text{totalClusters}} < \tau \), where \( \text{updatedClusters} \) represents the number of semi-clusters updated during the current iteration, while \( \text{totalClusters} \) represents the total number of semi-clusters in the graph.

**Sampling Requirements:** Semi-clustering has similar sampling requirements as PageRank: In particular, the sampling mechanism should maintain the connectivity among vertices (to avoid isolated sub-graphs) and the in/out node degrees proportionality, such that a proportionally smaller number of semi-clusters are sent along the edges of the sample graph in each iteration of the sample run.

**Transform function:** Semi-clustering’s convergence threshold is not tuned to the size of the dataset as a ratio of cluster updates decides convergence. Hence, we use the transform function: \( T = (1D_{\text{conf}}, T_S = \tau C) \), with \( D_{\text{conf}} = \{ f_{\text{ID}}, v_{\max}, C_{\max}, S_{\max} \} \), and \( sr \) is the sampling ratio. Intuitively, the total number of cluster updates on a sample that preserves the structure of the original graph is proportionally smaller than the total number of cluster updates on the complete graph. Similarly to the PageRank algorithm, such transformations assume perfect structural symmetry of the sample w.r.t. the original graph. Therefore, we adopt it as a heuristic, which shows good results in practice.

### 4.3 Top-K Ranking

Top-K ranking for PageRank [23] finds the top \( k \) highest ranks reachable to a vertex: Top-K ranking operates on output generated by PageRank and it proceeds for as follows: In the first iteration, each vertex sends its rank to the direct neighbors. In the following iterations, each vertex receives a list of ranks from all the neighboring nodes, it updates its local list of top-K ranks, and then it sends the updated list of ranks to the direct neighbors. A node that does not perform any update to its list of ranks in one iteration does not send any messages to the neighbors. As the number of messages and the message byte counts depend on the number of ranks stored per node, and whether the node has performed any updates to the list of top-k values, the runtime of consecutive iterations is not constant.

**Convergence:** Top-K ranking is executed iteratively until a fixed point is reached [23], or alternatively, until the total number of vertices executing updates goes below a user defined threshold: i.e., \( \frac{\text{activeVertices}}{\text{totalVertices}} < \tau \).

**Sampling Requirements:** There are two main requirements: i) maintaining connectivity, in/out node degrees and effective diameter among sampled vertices as for PageRank algorithm, and ii) maintaining the relative ordering of ranks for sampled vertices. Top-K ranking is executed on output generated by PageRank. Assuming an input sample that satisfies the sampling requirements of PageRank, the resulting output generated by PageRank preserves the connectivity and the relative order of rank values. Consider Figure 2, the rank of any node on \( S_h \) is twice the rank of the corresponding node on \( G \).

**Transform function:** We observe that the convergence condition is not tuned to the size of the input dataset as it uses a ratio of updates to decide convergence. For a sample that satisfies the sampling requirements, the ratio of rank updates on the sample is maintained in pair with the ratio of rank updates on the complete graph, hence, unlike PageRank algorithm, no scaling is required: \( T = (1D_{\text{conf}}, 1D_{\text{conv}}), \text{Conf} = \{ \text{topK} \}, \text{Conv} = \{ T_S \} \).

### 4.4 Labeling Connected Components

Labeling connected components is an algorithm that finds the number of connected components in a graph by mapping each vertex to a connected component identifier. The algorithm can be implemented in an iterative fashion as follows: Initially, the connected component value (i.e., CCV) of each vertex is initialized with the vertex identifier. In the first iteration, each vertex inspects the CCV of the neighboring vertices. If any of these values is smaller than the current CCV, the vertex changes its CCV with that one and broadcasts a message with the updated value to all of its neighboring vertices. In the following iterations, each vertex checks all the messages received from its neighbors. If any message includes a CCV smaller than the current identifier, the vertex changes its value and broadcasts a message with the updated CCV to all of its neighbors. The algorithm continues in a similar fashion until no new messages are being sent. A main characteristic difference between connected components and the previous algorithms is that the processing requirements of consecutive iterations may vary widely. Typically, a few long iterations are followed by multiple very short iterations.

**Convergence:** The total number of iterations required for running the connected components algorithm is bounded by the effective diameter [22].

**Sampling Requirements:** All of the three sampling requirements of the PageRank algorithm are equally important for connected components. We emphasize that vertices with a high-out degree (i.e., hub nodes) have a high impact on the convergence speed of the connected components algorithm. As such nodes are highly connected, their corresponding connected component identifier can be propagated towards other regions of the graph in a few steps. Hence, starting the sampling process from such nodes would be beneficial.

**Transform function:** Prior research showed that for uniform graphs, sampling mechanisms based on random walks typically maintain the diameter of the sample similar with the one of the complete graph [25]. For example, the sample graph \( G \) presented in Figure 2 has the same diameter with graph \( G \). Therefore, an explicit transform function is not required as for the other algorithms. In particular, \( T = (1D_{\text{conf}}, 1D_{\text{conv}}), \text{Conf} = \{ \}, \text{Conv} = \{ \} \).

### 4.5 Neighborhood estimation

Estimating the number of vertices reachable from a vertex \( \nu \) within \( h \) hops or shortly the neighborhood of \( \nu \) is used in social applications today. LinkedIn for instance provides information on the number of professionals reachable within \( h \) hops from any given user. We implement neighborhood estimation for all the vertices of an input graph using an iterative, probabilistic algorithm similar with estimating effective diameters and radii in large graphs [22]: Each vertex \( \nu \) of the graph stores the number of neighbors reachable from \( \nu \) in \( h \) hops as a set of \( k \) probabilistic Flajolet-Martin bitstrings \( b_k(h, \nu) \) [15]. In the first iteration, each vertex is initialized with a set of \( k \) random bitstrings. After initialization, each vertex sends its own bitstrings to the neighboring vertices. In the following iterations, each vertex updates its bitstrings using a bitwise OR operator among its bitstrings and the corresponding bitstrings received from the neighboring nodes. Only if the bitstrings are updated during the current iteration, the vertex sends again its updated bitstrings to the neighboring nodes. The algorithm has variable resource requirements per iteration as the number of messages sent, and the number of active vertices of each iteration depend on the actual number of vertices updating
their bitstrings.

Convergence: Unlike other algorithms, neighborhood estimation is executed until a fixed point is reached. The challenge stands in estimating per iteration key features such as active vertices and message byte counts as they vary from one iteration to the next. The neighborhood of a vertex \( v \) after \( h \) iterations is computed from the \( k \) Flajolet–Martin bitstrings by:

\[
N(h, v) = \frac{1}{\epsilon} \sum_{i=1}^{k} b_i(v),
\]

where \( b_i(v) \) is the position of leftmost 0 bit of the \( i \)-th bitstring of node \( v \), and \( k \) is the number of bitstrings stored at each node (a constant, typically 32 [22]).

Sampling Requirements: Maintaining connectivity among nodes, and effective diameter are primarily required. Preserving distances among sampled vertices contributes in propagating the bitstrings updates of the sample run at the same pace with those of the actual run.

Transform function: For a sample that satisfies the sampling requirements, the neighborhood function on the sample grows with the same rate as on the original graph: Consider vertex 1 in Figure 2: The number of vertices reachable within two hops on \( G \), is twice the number of vertices reachable within two hops on \( S_h \). Hence, the processing requirements during the sample run can be maintained proportionally with the processing requirements of the actual run using a sample that satisfies the sampling requirements. Considering that the neighborhood is estimated probabilistically starting from a set of \( k \) bitstrings that each vertex keeps updating, the only transformation required is to maintain the same set of initial bitstrings on the sampled nodes as on the complete graph. In particular, \( T = \{ID_{conf}, ID_{conv}\} \), where \( Conf = \{K, seed\} \), \( Conv = \{\}\). \( K \) is the number of bitstrings (e.g., 32), and \( seed \) is the seed used in generating the bitstrings of each node (each vertex sets its seed as the vertex id, such that the same initial bitstrings are generated for both the sample and the actual runs).

5. EXPERIMENTAL EVALUATION

Experimental Setup: Experiments are performed on a cluster of 10 nodes, each of the node having two six-core CPUs Intel X5660 @ 2.8GHz, 48 GB RAM and 1 Gbps network bandwidth. All experiments are run on top of Giraph 0.1.0, a library that implements the BSP model on top of Hadoop. We use Hadoop 1.0.3 as the underlying MapReduce framework. Unless specified otherwise each node is set with a maximum capacity of three mappers, each mapper having allocated 15GB of memory. Hence, our Giraph setup has a total of 30 tasks (i.e., 29 workers and one master).

Experimental Methodology: Four real datasets are used for evaluating PREDICT: Two of them are web graphs: Wikipedia, and UK 2002, and the remaining two are social graphs: LiveJournal and Twitter. The Wikipedia dataset is a subset of the online encyclopedia including the links among all English page articles. UK 2002 is the web graph of the .uk domain as crawled by UbiCrawler\(^2\) in 2002. LiveJournal graph models the friendship relationship among an online community of users\(^3\), while Twitter graph models the user base graph (i.e., the following relationships among users) as crawled in 2009. Table 2 is illustrating the characteristics of each dataset. A similar set of datasets were used in the context of optimizing iterative algorithms [14]. All datasets are directed graphs. For algorithms operating on undirected graphs we transform directed graphs into the corresponding undirected graphs. In Giraph, which inherently supports only directed graphs, to each edge its reverse edge is added.

Memory limits: The memory resources of our deployment are almost fully utilized when executing the algorithms on the largest datasets: i.e., Twitter and UK. In Giraph, per vertex state information and message buffers are allocated and stored into the memory of the cluster additionally to the input graph. Hence, the memory requirements to store the graph and its corresponding state are much larger than the size of the dataset itself. For instance, executing semi-clustering (which sends large number of large messages) on the UK dataset requires 90% of the full RAM capacity of our cluster, hence, the memory resources of our setup are almost fully utilized. As Giraph is currently lacking the capability of spilling messages to disk, we run out of memory when trying to run semi-clustering, top-k ranking, and neighborhood estimation on the Twitter dataset \(^5\).

<table>
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<tr>
<th>Name</th>
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<th># Nodes</th>
<th># Edges</th>
<th>Size [GB]</th>
</tr>
</thead>
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<td>68,993,777</td>
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<tr>
<td>Wikipedia</td>
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<td>97,652,232</td>
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<tr>
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<td>18,520,486</td>
<td>298,113,762</td>
<td>4.7</td>
</tr>
</tbody>
</table>

Table 2: Graph Datasets

Metrics of interest: For validating our methodology, we compute standard error metrics used in statistics that show the accuracy of the fitted model on the training data. In particular, we consider: the coefficient of determination (i.e., \( R^2 \)), and the signed relative error (i.e., negative errors correspond to under-predictions, while positive errors correspond to over-predictions).

Sources of error: There are two sources of error when providing end-to-end runtime estimates: i) misestimating key input features; ii) misestimating cost factors used in the cost model. Depending on the the error sign for key feature estimates and cost factors (over or under-prediction), the aggregated estimation errors can either accumulate or reduce the overall error. Hence, we first provide results on estimating key input features, then, we provide end to end runtime results.

Algorithms: We evaluate PREDICT on a set of representative algorithms: PageRank, semi-clustering, top-k ranking, connected components, and neighborhood estimation. Due to space constraints complete results for all algorithms are presented in the extended version of the paper [33].

5.1 Estimating Key Input Features

PageRank: This set of experiments shows the accuracy of predicting the number of iterations for PageRank algorithm as the size of the sampling ratio increases from 0.01 to 0.25. The convergence threshold value is set as \( \tau = 1/N \times \epsilon \), where \( N \) is the number of vertices in the graph, while \( \epsilon \) is the convergence tolerance level, a sensitivity parameter varied between 0.01 and 0.001. Figure 4 shows the results for all datasets when BRJ is adopted as the underlying sampling scheme. Sensitivity analysis w.r.t. the sampling method is deferred to section 5.3. For a sampling ratio of 0.1, and a tolerance level of \( \epsilon = 0.01 \) the maximum mis-prediction for the web graphs and Twitter datasets is less than 20%. LiveJournal has 40% relative error for the same sampling ratio. For this dataset, our results on multiple algorithms are consistently showing that the sampling method adopted \(^5\).
cannot capture a representative sample as for the other algorithms due to its underlying graph structure which is not scale-free. Lower errors correspond to a tolerance level of $\epsilon = 0.001$, when PageRank converges in a larger number of iterations. The relative errors for all datasets are maintained below 10% including LiveJournal. This is a desired outcome for a prediction mechanism, as accurate predictions are typically more useful for long running algorithms.

**Semi-clustering:** In this section we analyze the accuracy of predicting iterations for semi-clustering. The base settings we use in evaluation are: $C_{\max} = 1, S_{\max} = 1, V_{\max} = 10, f_R = 0.1, \tau = 0.001$. Figure 5 shows the accuracy results for all datasets but Twitter for two convergence ratios for $\tau = 0.01$, and $\tau = 0.001$. As explained in experimental methodology, as the memory footprint of semi-clustering algorithm on Twitter is much larger than the total memory capacity of our cluster we could not perform experiments on this dataset. For a sampling ratio of 0.1 the relative errors corresponding to all web graphs analyzed are below 20%.

Again, LiveJournal dataset shows higher variability in its error trend due to its underlying graph structure which is less amenable to sampling. We have performed sensitivity analysis w.r.t. $S_{\max}$ and $V_{\max}$ when running semi-clustering on LJ dataset, which has the highest relative error on the base settings. In particular, we analyzed two cases: i) increasing $S_{\max}$ from one to three, and ii) increasing $V_{\max}$ from ten to twenty. Compared with the base settings, for a sampling ratio of 0.1 (or larger) the relative errors were maintained in similar bounds for all sampling ratios.

**Top-K Ranking:** We analyze the accuracy of estimating iterations and the accuracy of estimating key input features (i.e., remote message bytes) in Figure 6. We execute sample runs on output generated by PageRank algorithm, and use a convergence threshold of $\tau = 0.001$. We observe that the relative errors for estimating iterations are below 35% for all scale free graphs analyzed, while the errors for estimating remote message bytes are below 10%. Similarly to our experiments on PageRank and semi-clustering, higher errors are observed for LiveJournal dataset: for a sampling ratio of 0.1, the number of iterations are over-estimated by a factor of 1.5, while the message byte counts by 40%. An interesting observation for top-k ranking is that the accuracy in estimating the message byte counts is more important than the accuracy of estimating the number of iterations per se. That is because the runtime of consecutive iterations varies and is proportional with the number of message byte counts and the number of active vertices of each iteration (results on estimating runtime are shown later, in Figure 9).

**Connected components:** For this experiment each cluster node was set with a maximum capacity of six mappers, each mapper having allocated 7GB of memory, accounting for a total of 60 tasks (i.e., 59 workers and one master). Due to the large processing variability among subsequent iterations, the number of iterations per se is not sufficient for predicting the runtime of connected components algorithm. Hence, we present the accuracy of estimating active vertices in addition to estimating iterations. Figure 7 a) shows the accuracy results when estimating iterations. For a sampling ratio of 0.1, the relative errors for all datasets but LJ are below 25%. Figure 7 b) shows the estimated total number of active vertices required for the execution of the algorithm (summed up for all iterations). For a sampling ratio of 0.1, the relative error for both web graphs is less than 10%. The reason that LJ highly over-estimates the total number of active vertices for a sampling ratio of 0.1 is that it is not scale-free, hence, the sample cannot capture the structure of the original graph. For Twitter, on the other hand, the sample of 0.1 is too small to capture key input features with a better accuracy than 81% due to the density of the graph (i.e., a very large number of incident edges per node): The sampling ratio of 0.1 vertices corresponds to a ratio of only 0.002 in terms of edges. Higher sampling ratios improve the accuracy results: For a sampling ratio of 0.25 the error
Figure 6: Top-k ranking key input features estimation: a) Estimating iterations (left), b) Estimating remote message bytes (right).

Figure 7: Predicting key input features for connected components: a) number of iterations (left), b) active vertices (middle), and c) active vertices for guided sampling (right).

decreases to 46%.

Sensitivity analysis w.r.t. sampling is showing that a smaller number of seed vertices used in BRJ sampling \((k = 100\) instead of \(k = 1\%\) of total vertices) improves the accuracy on Twitter dataset to 21\% relative error for a sampling ratio of 0.25 (Figure 7 c)). When using a smaller number of seed nodes in sampling, the average out degree of vertices increases (random walks return to already visited nodes more often, and more incident edges are picked), hence, the propagation of the connected components ids to all the vertices of the sample takes a fewer steps, as in the original graph. This result shows that additional information on the characteristics of the dataset and on the algorithm can guide the sampling process to achieve higher accuracy results. Similar trends are observed for predicting other input features such as message byte counters.

Neighborhood Estimation: We execute neighborhood estimation for a fixed number of iterations \(\text{numIter} = 10\): i.e., finding the number of vertices reachable within 10 hops. Figure 11 shows results for estimating remote message bytes. For a sampling ratio of 0.1 the relative errors for estimating remote message bytes are less than 19\% for all datasets analyzed. Compared with the other algorithms, we observe that the errors for LJ are much smaller for this case: As the number of iterations is fixed and the key input features variability among consecutive iterations is less pronounced than for algorithms like top-k ranking or connected components (a large number of vertices stay active and propagate messages to neighbors for the first 10 iterations), the overall estimations errors are reduced.

Upper Bound Estimates: In the following we analyze the accuracy of predicting iterations for PageRank when using analytical upper bound estimates. In particular, for PageRank iterations are approximated using the analytical upper bound as defined in the detailed survey of Langville et al. [24]: \(#\text{iterations} = \frac{\text{iter}_{\text{max}}}{\text{iter}_{\text{max}}^{\epsilon}}\), where \(\epsilon\) is the tolerance level as defined above, and \(d = 0.85\) is the dumping factor.

Note that the formula does not consider the characteristics of the input dataset, and as we show next, such bounds are loose: For instance, for a tolerance level of \(\epsilon = 0.001\) we obtain a number of 42 iterations using the above formula, whereas the actual number of iterations is less than 21 for all datasets (a factor of 2x misprediction). For a tolerance level of \(\epsilon = 0.1\) a misprediction of 3.5x is obtained for the Wikipedia dataset.

5.2 Estimating Runtime

In this section we show the accuracy of predicting the end-to-end runtime execution for semi-clustering, top-k ranking, connected components, and neighborhood estimation. As they show runtime variability among subsequent iterations, they are more challenging to predict than algorithms with constant per iteration runtime (i.e., PageRank). For training the cost model we show results for two cases: i) no prior executions of the algorithm exist (no history); ii) historical executions of the algorithm on different datasets exist. For the case that no history exists, sample-runs on samples of 0.05, 0.1, 0.15 and 0.2 are used for training. For the case that history exists, prior runs on all other datasets but the predicted one are additionally considered. We note that once a cost model is built it is used multiple times, for predicting the runtime of the same algorithm on different input datasets.

Semi-clustering: Figure 8 a) shows the accuracy of predicting runtime for the case that history does not exist. The coefficient of determination of the cost models corresponding to the three datasets on which predictions are made are as follows: \(R^2_{\text{LJ}} = 0.82\), \(R^2_{\text{WikiUK}} = 0.89\) and \(R^2_{\text{K}} = 0.84\), and are showing that each multi-variate regression model fits the training data (the closer the value to one, the better the model is). The key input features that achieve the highest correlation on the multi-variate model are the local and remote message byte counters. It can be observed that the error trend for each dataset is very similar with the cor-
Figure 8: Semi-clustering runtime prediction: a) Training with sample-runs (left), b) Training with sample- and actual-runs (right).

Figure 9: Top-k ranking runtime prediction: a) Training with sample-runs (left), b) Training with sample- and actual-runs (right).

responding error trend for predicting iterations (see Figure 5 for \( \tau = 0.001 \)). In contrast to predicting iterations, additional errors in estimating per-iteration input features (i.e., message byte counters) and cost model approximations are determining an error difference between the two graphs. For a sampling ratio of 0.1 the errors are less than 30\% for the scale free graphs and less than 50\% for LiveJournal.

Figure 8 b) shows similar results for the case that history exists. The corresponding coefficient of determination of the three models is improved: i.e., \( R^2_{LJ} = 0.95, R^2_{W_{iki}} = 0.95 \) and \( R^2_{UK} = 0.88 \). The error trends for Wikipedia and LiveJournal are similar as for the case that sample-runs are used for training. The cost factors for the UK dataset are improved and the errors are reduced to less than 10\% when using a sampling ratio of 0.1 or larger.

Top-K Ranking: We analyze the accuracy of estimating time in Figure 9. We observe that the error trends are less than 10\% for the scale free graphs analyzed. The key input features that achieve the highest correlation on the multivariate model are the local and remote message bytes and their corresponding message counts. For the case history is not used, the coefficient of determination of the models are as follows: \( R^2_{LJ} = 0.95, R^2_{W_{iki}} = 0.96 \) and \( R^2_{UK} = 0.99 \). Yet, the cost factors corresponding to the cost model for LJ dataset are over-predicted: That is due to the training phase which uses very short sample runs, especially for small datasets such as LJ. As the overhead of running very short iterations surpasses the actual processing cost associated to each key input feature, the coefficients of the cost model are over-estimated. Hence, the end to end relative errors are determined not only by over-predicting key input features, but also by over-predicting cost factors. In contrast to LJ, for larger datasets fairly accurate cost models can be built using sample-runs. For the case history is used, all the cost models are improved. The coefficient of determination of the models are: \( R^2_{LJ} = 0.99, R^2_{W_{iki}} = 0.99 \) and \( R^2_{UK} = 0.99 \). We observe that the error trends are in pair with the error trends for estimating message byte counts (Figure 6 b)).

Connected components: Figure 10 a) shows runtime results for the case that only sample-runs are used in training. Similar error trends as for the case of estimating active vertices are observed (see Figure 7 b)). We note that due to the variability among consecutive iterations, the number of active vertices and the message byte counts have a higher impact on runtime of connected components algorithm than the number of iterations per se.

The key input features that achieve the highest correlation on the multi-variate model are the number of active vertices, the local and the remote message byte counters. The coefficient of determination of the models corresponding to the four datasets on which predictions are made are as follows: \( R^2_{LJ} = 0.88, R^2_{W_{iki}} = 0.94, R^2_{UK} = 0.98, \) and \( R^2_{TW} = 0.99 \). For a sampling ratio of 0.1 the relative error for Wikipedia dataset is 28\% and for UK is -23\%. When historical runs are additionally used in training the corresponding errors decrease to 19\% and -8\% respectively. The high errors on LiveJournal datasets are determined in part by key input features over-predictions and in part by cost factors over-estimations (for a very similar reason as for top-k algorithms explained above). The causes of errors for Twitter are mainly coming from over-predicting key input features. Figure 10 c) is showing the corresponding results when the number of seed nodes used for BRJ sampling is set to 100 (guided sampling). While the web graphs are marginally affected by a smaller number of seed nodes, the accuracy on Twitter is improved by 30\% for a 0.25 sampling ratio.

Neighborhood Estimation: Figure 12 shows accuracy results for estimating runtime for neighborhood estimation. For a sampling ratio of 0.1 the relative errors for estimating runtime in the case that history is not used are less than 21\%, while for the case history is used, all errors are reduced to less than 10\% for the same sampling ratio. The key input features that achieve the highest correlation on the multi-variate model are the active vertices, the total vertices, and the local and remote message bytes, and the coefficient of determination of the models: \( R^2_{LJ} = 0.99, R^2_{W_{iki}} = 0.99, \) and \( R^2_{UK} = 0.98 \).
In this section we analyze the accuracy of predicting iterations when varying the underlying sampling technique. In order to analyze the impact of bias on maintaining key properties on the sample, we compare RJ with BRJ. Additionally, we select MHRW [17], another sampling technique based on random walks that in contrast with RJ, removes all the bias from the random walk, which is known to inherently have some bias towards high degree vertices. All sampling techniques use a probability \( p = 0.15 \) for restarting the walk, while the number of seed vertices for BRJ is \( k = 1\% \) of the total vertices of the graph.

Figure 13 shows sensitivity analysis for predicting iterations for PageRank, semi-clustering and top-k ranking on UK dataset. Figure 14 shows sensitivity analysis for predicting key input features for connected components (i.e., active vertices, iterations) and neighborhood estimation (i.e., remote message bytes). We observe that for a sampling ratio of 0.1, the relative error for BRJ sampling are generally better than for all the other sampling techniques. The result shows that the bias towards high out-degree vertices of BRJ contributes to a good accuracy in prediction for the algorithms we analyze in this paper. The reason is that convergence of these algorithms is inherently “dictated” by highly connected nodes: For instance, for PageRank such nodes contribute a large share to the average rank value, or for semi-clustering they contribute significantly to the ratio of semi-cluster updates.

While other iterative algorithms executing graph processing tasks such as: random walks with restart [22] (proximity estimation), or Markov clustering [37] are expected to benefit from similar sampling methods based on random walks, customized sampling methods may be required for other algorithms. In section 5.1 [33], we showed one example dataset for connected components algorithm, where guiding the sampling process can further improve the accuracy of results, given that more information about the input dataset is available.

Finally, in order to evaluate the consistency of the sampling method, we perform further sensitivity analysis: For each sampling ratio we take multiple samples (using different starting seeds for the random number generator), we run sample-runs on each of them and evaluate the standard deviation for estimating iterations. For a sampling ratio of 0.1, the largest deviations observed are as follows: For PageRank: 3% for scale free graphs and 0% for LiveJournal, for semi-clustering: 5% on scale free graphs and 14% on LiveJournal, and for connected components: 9% on scale free graphs and 10% on LiveJournal. While there is some inherent variability in the sampling process, the error trends are maintained similar among different sample instances.

5.4 Overhead Analysis

This section compares the runtime of the sample-run with that of the actual-run. Table 3 shows the runtime of all algorithms on multiple sampling ratios, on the largest graphs: Twitter and UK. For PageRank, the runtime of the sample-run on a 0.1 sample of the Twitter dataset accounts for 3.5% of the runtime of the actual-run. The reason is that the our sampling mechanism stops after a given ratio of vertices (not edges) is sampled. As Twitter graph is much denser than the others, the average number of incident edges per vertex is almost 9x smaller in the sample graph. For semi-clustering, the runtime of the sample-run on a 0.1 sample of the UK dataset accounts for 4.8% of the runtime of the actual-run for a similar reason as before. We note that the runtime
of the sample-run is much smaller than the runtime of the actual-run particularly for long running algorithms, where the runtime of the iterations dominate the runtime of the algorithm (i.e., the overhead of pre-processing the graph is relatively small). For algorithms where the overhead of pre-processing the graph dominates, the overhead of running sample-runs is higher. Connected components on Twitter is one such example: The actual time spent in running iterations is 19 seconds for the sample-run, which accounts for 4% of the time spent in running iterations for the actual-run (i.e., 465 sec). Yet, due to the overhead of reading, partitioning and outputting the result, that accounts for more than 80% of the sample-run time, the overall runtime of the sample-run relative to the runtime of the actual-run is higher, accounting for 12% of its time.

The data in Table 3 shows the runtime of sample-runs and actual-runs for PageRank (PR), semi-clustering (SC), connected components (CC), top-k ranking (TOP-K), and neighborhood estimation (NH), in seconds.

### Table 3: Runtime of sample-runs and actual-runs

<table>
<thead>
<tr>
<th>Sampling Ratio</th>
<th>PageRank (PR)</th>
<th>Semi-clustering (SC)</th>
<th>Connected Components (CC)</th>
<th>Top-k Ranking (TOP-K)</th>
<th>Neighborhood Estimation (NH)</th>
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</thead>
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<td>4192</td>
<td>861</td>
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</tr>
</tbody>
</table>

**5.5 Resource Allocation**

We present one experiment that demonstrates PREDiCT’s applicability for estimating runtime when a different resource allocation (i.e., number of slots) is used during the sample run: In particular, we use 15 workers for the sample-run, and 29 workers for the actual run. Figure 15 shows the results for estimating runtime for semi-clustering algorithm. In contrast with Figure 8 b) (where the same slot configuration was used for the sample and the actual runs) increased errors are observed (in particular for UK dataset) due to an additional level of critical path approximation: i.e., given a different number of slots to execute the algorithm, we use a uniform scaling factor to scale the two extrapolating factors (on edges and vertices). For higher accuracy results, our framework is extensible to support per worker extrapolating factors, according to the partition size of each worker (i.e., number of edges and vertices each worker is allocated with).

**6. CONCLUSIONS**

This paper presents PREDiCT, an experimental methodology for predicting the runtime of a class of iterative algorithms operating on graph structures. PREDiCT builds on the insight that the algorithm execution on a small sample can be transformed to capture the processing characteristics of the complete input dataset. Given an iterative algorithm, PREDiCT proposes a set of transformations: i.e., a sample technique and a transform function, that only in combination can maintain key input feature invariants among the sample run and the actual run.

Additionally, PREDiCT proposes an extensible framework for building customized cost models for iterative algorithms.
executing on top of Giraph, a BSP implementation. Our experimental analysis of a set of diverse algorithms: i.e., ranking, semi-clustering, and graph processing shows promising results both for estimating key input features and time estimates. For a sample ratio of 10%, the relative error for predicting key input features ranges in between 5%-35%, while the corresponding error for predicting runtime ranges in between 10%-30% for all scale-free graphs analyzed.

7. REFERENCES


[39] G. Wang, W. Xie, A. J. Demers, and J. Gehrke. Asynchronous Large-Scale Graph Processing Made
