A Comparison of Graph Processing Systems

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Abstract—TODO

Index Terms—graphs, distributed computing, Galois, Ligra, Polymer, Giraph, Gluon, Gemini

I. Introduction

In recent years graph sizes have increased significantly [?] . Thus performance and memory efficiency of the graph analysis applications is now more important than ever.

Many applications like

Wachsende graphen, benötigt für folgende Applikationen, verwenden graph algorithmen, diese müssen schnell und verteilt laufen dafür wurden Frameworks entwickelt...

und das noch schön formulieren

We compare several non-uniform memory access (NUMA) aware systems in terms of their performance on three graph algorithms (PageRank, SSSP, BFS). The comparison is performed on both real world data sets and synthetic graphs.

To provide a comparison to a non-NUMA aware system, Giraph[7] is included in the testing. Giraph itself has often been compared to other state of the art systems like Pregel or GraphX.

To this end several non-uniform memory access (NUMA) aware systems were proposed like Polymer [3], Galois [1] or Ligra [5].

This paper makes the following contributions:

- Comparison of several state-of-the-art graph processing frameworks
- •
- •

II. PRELIMINARIES

a) Graphs and Paths: An unweighted graph is the pair G=(V,E) where the vertex set is $V\subseteq \mathbb{N}$ and the E is the edge set. The edge set describes a number of connections or relations between two vertices. Depending on these relations, a graph can be directed or undirected. For a directed graph the edge set becomes

$$E \subseteq \{(x,y) \,|\, x,y \in V, x \neq y\}$$

and in the undirected case E is a set of two-sets

$$E \subseteq \{\{x,y\} \mid x,y \in V, x \neq y\}.$$

The main difference is thus, that in the case of a directed graph a connection between s and t is not the same as a connection between t and s – the direction matters.

Independently of the graph being directed or not, a graph can be *weighted*. In this case the edge set is expanded by a numerical value $E_{\text{weighted}} = E_{\text{unweighted}} \times \mathbb{R}$ further describing the relation.

The size of a graph is often described in the number of edges |E| because this number is typically much larger than the number of vertices |V|.

A Path from start s to target t is a set of edges $P \subseteq E^n$ with

$$P = ((x_0, x_1), (x_1, x_2), \dots, (x_{n-1}, x_n))$$

where all $(x_i, x_{i+1}) \in E$ and $x_0 = s, x_n = t$.

Thus we call a target t accessible from s if a Path from s to t exists.

b) Single-Source Shortest-Paths: Single-Source Shortest-Paths (SSSP) describes the problem of finding a path along some edges of the input graph from a given start node to a target.

Input to the problem is a weighted graph G=(V,E) and a start node $s\in V$. Output is the shortest possible distance from s to each node in V. The distance is defined as the sum of edge weights w_i on a path from s to the target. In the case of a unweighted graph, the distance is often described in hops, i.e. the number of edges on a path.

The most common implementations are Dijkstra's algorithm or BellmanFord.

c) Breadth-First Search: Breadth-first search (BFS) is a search problem on a graph. It usually requires an unweighted graph and a start vertex as input. In some cases a target vertex is also given.

The output is usually a set of vertices that are accessible from the start vertex. In the case of a target being given, the output is true if a path from start to target exists or false otherwise.

- d) PageRank:
- e) Pregel Model:
- f) Congest Model:

III. OVERVIEW OF THE FRAMWORKS

A. Galois and Gluon

Galois[1] is a general purpose library designed for parallel programming. The Galois system supports fine grain tasks, allows for autonomous, speculative execution of these tasks and grants control over the task scheduling policies to the application. It also simplifies the implementation of parallel

applications by providing an implicitly parallel unordered-set iterators.

For graph analytics purposes a topology aware work stealing scheduler, a priority scheduler and a library of scalable data structures have been implemented. Galois includes applications for many graph analytics problems, among these are single-source shorthest-paths (sssp), breath-first-search (bfs) and pagerank. For most of these applications Galois offers several different algorithms to perform these analytics problems and many setting options like the amount of threads used or policies for splitting the graph. All of these applications can be executed in shared memory systems and, due to the Gluon integration, with a few modifications in a distributed environment.

Gluon[2] is a framework written by the Galois team as a middleware to write graph analysis applications for distributed systems. It reduces the communication overhead needed in distributed environments by exploiting structural and temporal invariants.

The code of Gluon is embedded in Galois. It is possible to integrate Gluon in other frameworks too, which the Galois team showed in their paper[2].

B. Gemini

Gemini is a framework for parallel graph processing[6]. In comparison to the other NUMA-aware frameworks we discuss here, Gemini does not support shared memory parallel processing. Computation is entirely message based using MPI. Gemini is lightweight and only provides basic functionality. A basic API hides data and computation distribution details from the user writing applications. There are five algorithms (which ones?) already implemented.

C. Giraph

Apache Giraph is an iterative graph processing framework, built on top of Apache Hadoop[7]. Hadoop as a large MapReduce infrastructure providing a reliable (fault tolerant) basis for large scale graph processing. Because of the underlying Hadoop MapReduce infrastructure, expanding single-node processing to a multi-node cluster is almost seamless.

Giraph is based on computation units that communicate using messages and are synchonized with barriers.

The input to a Giraph computation is always a directed graph. Not only the edges but also the vertices have a value attached to them. The graph topology is thus not only defined by the vertices and edges but also their initial values. Furthermore, one can mutate the graph by adding or removing vertices and edges during computation.

Computation is vertex oriented and iterative. For each iteration step called superstep, the Compute method implementing the algorithm is invoked on each active vertex, with every vertex being active in the beginning. This method receives messages sent in the previous superstep as well as its vertex value and the values of outgoing edges. With this data the values are modified and messages to other vertices are sent.

Communication between vertices is only performed via messages, so a vertex has no access to values of other vertices or edges other than its own outgoing ones.

Supersteps are synchronized with barriers, meaning that all messages only get delivered in the following superstep and computation for the next superstep can only begin after every vertex has finished computing the current superstep. Edge and vertex values are retained across supersteps.

Any vertex can stop computing (i.e. setting its state to inactive) at any time but incoming messages will reactivate the vertex again. A vote-to-halt method is applied, i.e. if all vertices are inactive or if a user defined superstep is reached the computation ends.

Each vertex outputs some local information (e.g. the final vertex value) as result.

Giraph being an Apache project makes it the most actively maintained and tested project in our comparison. Over the course of this project, several new updates were pushed to Giraph's source repository¹.

TODO: Facebook?

D. Ligra

Ligra[5] is a lightweight parallel graph processing framework for shared memory machines. It offers a programming inteface that allows expressing graph traversal algorithms in a simple way.

Algorithms can use the EdgeMap to make computations based on edges or the VertexMap to make computations based on vertices. Those mappings can be applied only to a subset of vertices. Based on the size of the vertex subset the framework automatically switches between a sparse and a dense representation to optimize speed and memory.

E. Polymer

Polymer is very similar to Ligra, in fact Polymer inherits the programming interfaces EdgeMap and VertexMap from Ligra as its main interface.[3]

Polymer is a vertex-centric framework, that tries to circumvent some of the random memory access drawbacks of such a design. It treats a NUMA machine as a distributed cluster and splits work and graph data accordingly between the nodes. Application-defined data is not distributed. Other runtime state data is allocated in a distributed way but only accessed through a global lookup table.

IV. AND

A. An Overview of Important Graph Formats

Since every frameworks uses different graph input formats, we supply a conversion tool capable of translating from EdgeList to the required formats. Data Sets retreived from KONECT can be directly read and translated.

The following sections explain the output formats of our conversion tool.

¹https://gitbox.apache.org/repos/asf?p=giraph.git

1) AdjacencyList: The AdjacencyList and WeightedAdjacencyList formats are used by Ligra and Polymer. The format was initially specified for the Problem Based Benchmark Suite, an open source repository to compare different parallel programming methodologies in terms of performance and code quality [?].

The file looks as follows

$$n, m, o_1, \ldots, o_n, t_1, \ldots, t_m$$

where commas are \n . First, n is the number of vertices and m the number of edges in the graph.

The o_k are the so-called offsets. Each vertex k has an offset o_k , that describes an index in the following list of the t_i . The t_i are vertex IDs describing target nodes of a directed edge. The index o_k in the list of target nodes is the point where edges outgoing from vertex k begin to be declared. So vertex k has the outgoing edges

$$(k, t_{o_k}), (k, t_{o_k+1}), \dots, (k, t_{o_{k+1}-1}).$$

For the WeightedAdjacencyList format, the weights are appended to the end of the file in an order corresponding to the target nodes.

2) EdgeList: The EdgeList format is the most intuitive and one of the most commonly used in online data set repositories. The KONECT database uses this format and thus it is the input format for our conversion tool.

An edge list is a set of directed eges $(s_1,t_1),(s_2,t_2),\ldots$ where s_i is a vertex ID representing the start vertex and t_i is a vertex ID representing the target vertex. In the format, there is one edges per line and the vertex IDs s_i,t_i are separated with any whitespace character.

For a WeightedEdgeList, the edge weights are appended to each line, again separated by a whitespace character.

3) Binary EdgeList: The binary EdgeList format is used by Gemini.

For s_i, t_i some vertex IDs and w_i the weight of a directed edge (s_i, t_i, w_i) , Gemini requires the following input format

$$s_1t_1w_1s_2t_2w_2\dots$$

where s_i, t_i have uint32 data type and the optional weights are float32. Gemini will derive the number of edges from the file size, so there is no file header or anything similar allowed.

4) Giraph's I/O formats: Giraph is capable of parsing many different input and output formats. All of those are explained in Giraph's JavaDoc². Both edge- and vertex-centric input formats are possible.

One can even define their own input graph representation or output format. For the purposes of this paper, we used an existing format similar to AdjacencyList but represented in a ISON-like manner.

In this format, the vertex IDs are specified as long with double vertex values, float out-edge weights. Each line in the graph file looks as follows

$$[s, v_s, [[t_1, w_{t_1}], [t_2, w_{t_2}]...]]$$

TABLE I: Size Comparison of the Used Graphs

Graph	# Vertices (M)	# Edges (M)
flickr	0.1	2
orkut	3	117
wikipedia	12	378
twitter	52	1963
rMat27	63	2147
friendster	68	2586
rMat28	121	4294

with s being a vertex ID, v_s the vertex value of vertex s. The values t_i are vertices for which an edge from s to t_i exists. The directed edge (s,t_i) has weight w_{t_i} .

There is no surrounding pair of brackets and no commas separating the lines as it would be expected in a JSON format.

V. TESTING METHODS

A. Hardware

For testing the graph processing systems, we used 5 machines with two AMD EPYC 7401 (24-Cores) and 256 GB of RAM each. One of those machines was only used as part of the distributed cluster, since it only has 128 GB of RAM. All five machines were running Ubuntu 18.04.2 LTS. Setup of each framework was performed according to our provided installation guides³.

B. Benchmark Software

All benchmarks were initiated by our benchmark script, that is available in our repository. Galois, Gemini and Giraph were benchmarked on both the 5-node cluster and a singe machine. Since Galois supports this parameter, we ran multiple tests comparing Galois' performance with different thread counts on a single machine.

The complete benchmark log files and extracted raw results are available in our repository.

C. Data Sets

The graphs used in our testing can be seen in detail in Table I. We included a variaty of different graph sizes, from relatively small graphs like the flickr graph with 100 thousand edges up to an rMat28 with 4.2 billion edges. All graphs except the rMat27 and rMat28 are exemplary real-world graphs and were retrieved from the graph database⁴ associated with the Koblenz Network Collection (KONECT)[8]. Both the rMat27 and rMat28 were created with a modified version (we changed the output format to EdgeList) of a graph generator provided by Ligra.

D. Measurements

For every framework, we measured the *execution time* as the time from start to finish of the console command.

For the *calculation time*, we tried to extract only the time the framework actually executed the algorithm. We came up with the following:

²http://giraph.apache.org/apidocs/index.html

 $^{^3{\}rm The}$ setup guides are available at https://github.com/SerenGTI/Forschungsprojekt/tree/master/documentation

⁴http://konect.uni-koblenz.de/

TABLE II: Tested Algorithms of Each Framework

Algorithm	Galois	Gemini	Giraph	Ligra	Polymer
PageRank	$\Delta { m Push}, \ \Delta { m Pull}$				Regular, Δ
SSSP	?, Push, Pull				
BFS	Yes		Yes*		

^{*} Algorithm not natively supported † Modified algorithm

^d Single node and distributed

 For Galois, we resulted to extracting console log time stamps. Galois outputs Reading graph complete.. Calculation time is the time from this output to the end of execution.

We know that this is not the most realiable way for measuring the calculation times. Not only due to unavoidable buffering in the console output we expect the measured time to be larger than the actual. First, it is not clear that all initialization is in fact complete after reading the graph. Second, we include time that is used for cleanup after calculation in the measurement.

- Polymer outputs the name of the algorithm followed by an internally measured time.
- Gemini outputs a line exec_time=x, which was used to measure the calculation time.
- Ligra outputs its time measurement with Running time : x.
- Giraph has built in timers for the iterations (supersteps) which we summed up to extract the computation time.

Furthermore, the *overhead* is the time difference between execution time and calculation time.

Each test case consisting of graph, framework and algorithm was run 10 times, allowing us to smooth slight variations in the measured times. Later on, we provide the mean values of the individual times as well as the standard deviation where meaningful.

E. Algorithms

The three problems Breadth-first search (BFS), PageRank (PR) and Single-source shortest-path (SSSP) were used to benchmark framework with every graph. For frameworks that support multiple implementations (e.g. PageRank in push and pull modes), we included the alternatives in our testing. Table II shows the tested algorithms in detail.

F. Comparison of setup

VI. RESULTS

A. The frameworks during setup and benchmark

We would like to raise some issues we encountered first while installing and configuring and second while running the different frameworks.

1) During setup and benchmark of Gemini, we encountered several bugs in the cloned repository. These include

non zero-terminated strings or even missing return statements.

The errors rendered the code as-is unable to perform calculations, forcing us to fork the repository and modify the source code. A repository with our changes can be found here⁵.

- 2) Furthermore, we would like to address the setup of Hadoop for Giraph. It requires multiple edits in xml files that aren't easily automized. This makes the setup rather time consuming, especially if reconfiguration is needed later on.
- 3) In order for Giraph to run, several Java tasks (the Hadoop infrastructure) have to be constantly running in the background. While we don't expect this to have a significant performance impact on other tasks, it is still suboptimal.

4)

On a plus side, setup of frameworks like Polymer or Ligra was straight forward and did not require any special treatment.

⁵https://github.com/jasc7636/GeminiGraph

B. Single-source Shortest-paths

- 1) Single-node: Figure 1 shows the average calculation times (time without initialization overhead), execution time and the normalized overhead for SSSP on the different frameworks. In these figures, Galois with 96 threads is shown. We show the impact of Galois' thread count in subsection VI-E.
- 2) Distributed: For the distributed scenario, Figure 2 shows the benchmark results as calculation and execution times.

Results are especially interesting for Giraph since it seems to not cope well with synthetic graphs. Analyzing the computation times in Figure 2a, we see that it is the fastest framework on our real-world graphs. And that with a considerable margin of other frameworks always taking at least 50% longer (Gemini on flickr) up to Galois Pull needing $18 \times$ longer on wikipedia. On both synthetic graphs however, Giraph is the slowest to compute. Giraph requires $12 \times$ or even $15 \times$ the computation time of Gemini on rMat27 or rMat28 respectively.

While Giraph's computation times are very competitive, when comparing the execution times in Figure 2b we see that Giraph is actually the slowest framework on 5 out of 7 graphs. For the other two, namely twitter and friendster, Giraph is second slowest with only Gemini taking longer to complete.

Giraph and Gemini's very long execution times are only due to their overhead being many orders of magnitude larger than Galois overhead (Figure 3). Overhead for Gemini is greater than that of Galois on every graph. From just a 20% increase on flickr up to friendster, where the overhead is $90\times$ that of Galois Push. For Giraph the overhead times are not as extreme but still generally worse. Even on flickr, Giraph's overhead time is already $23\times$ that of Galois. On friendster, where Gemini was worst, Giraph *only* requires $73\times$ the overhead time of Galois.

C. Breadth-first search

In these figures, Galois with 96 threads is shown. Again, we show the impact of Galois' thread count in subsection VI-E.

D. PageRank

E. Galois speedup

Analyzing the calculation time speedups for Galois, we can compare how or if the different algorithms benefit from increasing thread numbers.

1) Single-source Shortest-path: Starting with SSSP which is the algorithm that really is at an advantage when using many threads in Figure 8.

For all larger graphs, speedup is in most cases very close to optimal up to about 8 threads. Twitter has the best speedup, requiring only 38% the calculation time with 2 threads compared to one, 25% using 4 and 12.9% using 8 threads. Behaviour on friendster is similar with 52% at 2 threads, 29% at 4 and 16% at 8 threads compared to one.

Anything above 32 threads however no longer helps decrease the computation time, in some cases even the opposite e.g. calculation on rMat28 is actually slower with 48 (13% slower) or 96 threads (28% slower) compared to 40 threads.

Small graphs like flickr or orkut, neither benefit from more threads nor is the performance held up by synchronization overhead.

2) Breadth-first search: Flickr is sped up by about 5% with 4 threads, any more than that will actually decrease performance, making computation time up to 15% (96 threads) longer.

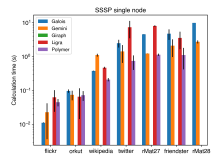
On all graphs, the speedup never exceeds $6\times$ even when using 96 thread. The initial speedup when switching from one to two threads is actually smaller than one, thus decreasing performance, on 4 of 7 graphs. Only flickr, twitter and rMat28 can benefit slightly by a speedup of 2%, 41% and 5% respectively.

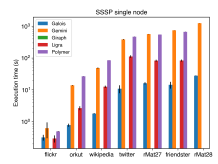
When comparing four threads to one, BFS on all but two graphs can be sped up by anywhere from 5% (flickr) to 68% (rMat28). The speedup is thus only possible to a very small degree. For the other two graphs, computation on friendster with 8 threads is just as fast as one thread and computation on rMat27 is actually 19% slower.

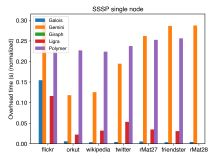
3) PageRank:

VII. DISCUSSION VIII. CONCLUSION

The conclusion goes here.

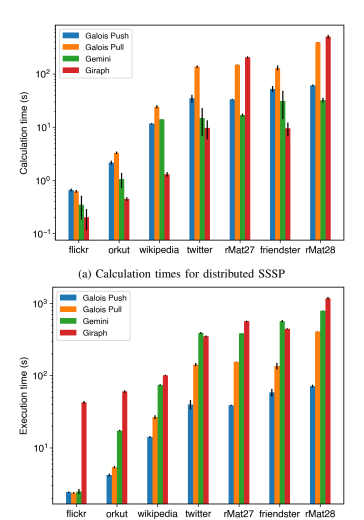






- (a) Calculation times for SSSP on a single node
- (b) Execution times for SSSP on a single node
- (c) Overhead time normalized by the graph size in million edges

Fig. 1: Average times on a single computation node, black bars represent one standard deviation in our testing. The runs on rMat28 for Ligra and Polymer failed and the frameworks were unable to complete the task.



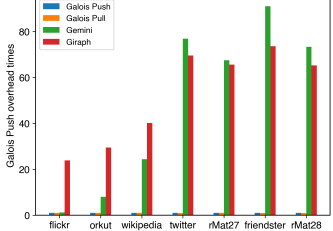
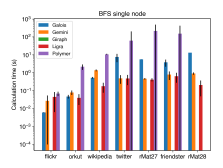
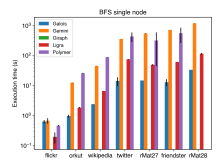


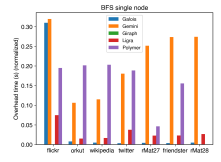
Fig. 3: Overhead times of each framework normalized by the overhead time of Galois Push

(b) Execution times for distributed SSSP

Fig. 2: Average times on the distributed cluster, black bars represent one standard deviation in our testing.





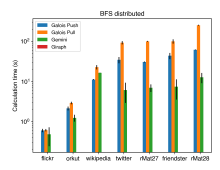


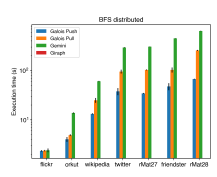
(a) Calculation times for BFS on a single node

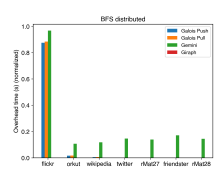
(b) Execution times for BFS on a single node

(c) Overhead time normalized by the graph size in million edges

Fig. 4: Average times on a single computation node, black bars represent one standard deviation in our testing





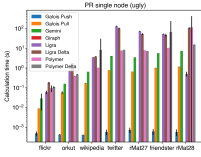


(a) Calculation times for distributed BFS

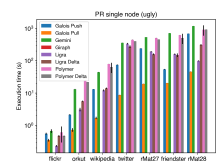
(b) Execution times for distributed BFS

(c) Overhead time normalized by the graph size in million edges

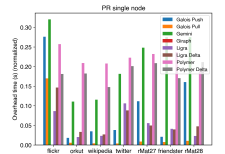
Fig. 5: Average times on the distributed cluster, black bars represent one standard deviation in our testing







(b) Execution times for PR on a single node



(c) Overhead time normalized by the graph size in million edges

Fig. 6: Average times on a single computation node, black bars represent one standard deviation in our testing

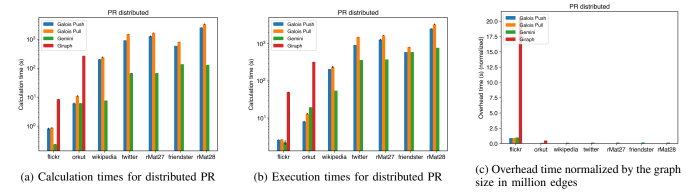


Fig. 7: Average times on the distributed cluster, black bars represent one standard deviation in our testing

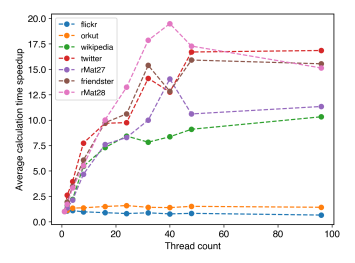


Fig. 8: Calculation time speedup with increasing thread count for Galois Single-source Shortest-paths

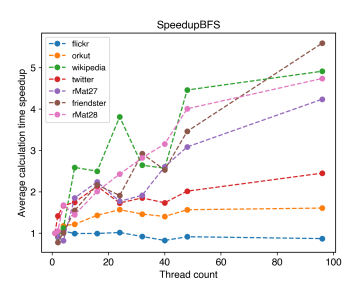
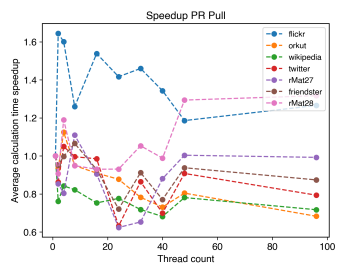


Fig. 9: Calculation time speedup with increasing thread count for Galois Breadth-first search



(a) PageRank Pull

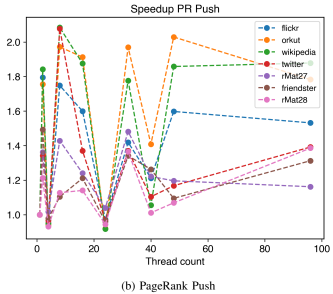


Fig. 10: Calculation time speedup with increasing thread count for Galois PageRank Push and Pull algorithms.

ACKNOWLEDGMENT

We are using the graph frameworks Galois [1], Ligra [5], Polymer [3], Gemini [6] as well as Apache Giraph [7].

Also we use Gluon [2] for the distributed Galois setups. Gemini [6]

SUPPLEMENTARY DATA

We have written a number of conversion tools and installation guides to help users or developers with the use of the tested frameworks.

Our GitHub repository: http://www.github.com/serengti/Forschungsprojekt.

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