Graph Partitioning for Distributed Shared Memory

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Abstract

The modern internet generates petabytes of data per day. Processing vast amounts of data is an increasingly common task both for scientists and modestly experienced programmers. Often this data is naturally represented as a graph, such as social media networks, webpage links or city networks, and requires clusters of machines to process. Concurrent trends in data centre architecture suggest that the rack is the new server, and shared memory is now a feasible interface between collocated rack servers. These trends made us wonder: How simple can fast graph processing be on a rack of servers?. We investigated the tradeoffs of the conventional Pregel "think like a vertex" programming model, and found its performance unacceptable. In contrast, we explored the merits of a "Think like a subgraph" model, which respects graph locality in common graphs, provides a more holistic programming interface, and runs faster!

1 Introduction

Big data processing is complicated. Scientists and experienced programmers alike struggle with managing and configuring clusters of machines to processes large amounts of data. Frameworks like Hadoop and Pregel have significantly eased the difficulty of big data processing, but they remain intimidating for the layman. We noticed a trend in scientists and researchers, finding that they wanted to "Just write python code that ran on a bunch of computers". For the benefit of science we investigated how to make this dream a reality.

Running all Python code on clusters of machines is impractical would lead to sluggish un-optimal code, due to immense network overhead. Instead, we concentrated our efforts on a common but difficult big data processing task: graph processing. Many frameworks exist for distributed graph processing [19, 6, 15, 18, 27, 9], and many general frameworks exist which are used for graph processing [26, 28, 13, 22]. These frameworks vary in their complexity, but none are "accessible" for programmers that

lack relevant experience.

By definition all programmers must be able to produce single threaded programs. Some researchers have argued and demonstrated that single threaded applications can outperform scalable distributed frameworks with little effort [20], and that such applications should be used as benchmarks for scalability. We evaluate these claims and propose a simple framework for multi-threading optimal single threaded applications.

With the exception of [15], the aforementioned systems suffer a common pitfall to accessibility; they expose the complexity of a distributed message passing system to the user. Extensive work has been done to hide this complexity in the abstraction of distributed shared memory (DSM) [14, 24, 21, 10, 12]. The benefits of DSM have been ignored in recent years due to its flaws, most notably fate sharing and sub optimal performance. Dismissing DSM may have been a shortsighted mistake. Ultra dense memory and the approach of terabit-level bandwidth within a rack give modern racks the appearance of a single machine, and has lead towards disaggregated architectures [1, 2, 3, 5, 11]. Such futuristic systems lend themselves naturally to DSM which motivates our proposal for a corresponding computation framework.

The largest disadvantage of DSM is performance. Programmers can write terribly performant programs by failing to reason about the location of memory, leading to memory thrashing. In computation where a high degree of consistency between shared resources is needed, DSM is the wrong tool for the job. In contrast, when large amounts of computation can be performed between memory synchronizations, DSM provides a simple and efficient programming model. Graph processing suffers from a lack of locality. In computations such as PageRank, a single iteration may perform edge updates which require the synchronization of all machines in a cluster. This problem can be largely avoided in practice by carefully pre-processing graphs into equally-sized partitions, where the minimum number of graph edges exist across machines. The cost of pre-processing a graph can be large; in some cases, the complexity of finding a good partition is greater than solving the initial problem! Here we demonstrate that the cost of graph partitioning is worth it for the benefits that DSM provides.

In this paper we attack the problem of developing a simple and efficient graph processing interface for DSM. Specifically we make the following contributions.

- An evaluation of single threaded applications as a baseline for scalability
- A simple framework for scaling single threaded applications
- A graph processing API for DSM
- A graph partitioning scheme optimal for DSM
- An evaluation of processing performance between partitioned DSM processing, and Pregel-style graph processing

The remainder of this paper is organized as follows. In Section 2 we discuss approaches and strategies for scalability. In Section 3 we methodology for automatically scaling up graph processing. Section 4 describes other systems similar to ours, Section 5 discusses future work, and Section 6 concludes the paper.

2 Scalability Measure

Before we could investigate advantages of any particular scalable framework we required a tool for measuring benefits gained from scalability. COST [20] proposed that the scalable benefits of any framework can be measured by comparing their performance to a single threaded program which perform an identical computational task. We began by attempting to duplicate COST's single threaded results to provide ourselves with a benchmark when scaling to multiple threads.

To obtain a baseline we ran COST's open source single threaded page rank on our own infrastructure. All of our experiments were run on a 60 core server composed of 4 Intel(R) Xeon(R) CPU E7-4870 v2 @ 2.30GHz NUMA nodes, and 128GB of DIMM DDR3 1600 MHz memory. Costs PageRank had an average runtime of 350s on our machine, most likely due to 2014 mac books having a higher averted clock speed of 3.5GHz.

Using simplicity as our guide we implemented single threaded PageRank [23] and processed the popular twitter_rv graph. We implemented PageRank in Go, and used it's built in map (key-value store) to store and iterate over edges in vertex order. We computed 20 iterations of PageRank on the 1.4 trillion edge graph in 374854s, approximately 4 days. These initial results were troubling as

COST boasted 300s processing using the same approach, a speedup of nearly 1071x.

To achieve higher performance in our single threaded application we carefully analyzed our data structures and data access path. Go's maps are unordered resulting in poor locality. To improve locality, we placed edges, and vertices in separate sequential arrays, and used a third array as a mapper between vertex id's and edge locations. Go's garbage collection, and runtime composed a large portion of our initial runtime. We minimized garbage collection by performing minimal memory allocations. These modification were non-trivial requiring a working knowledge of Go's runtime, and machine architecture. Post modification, our program executed in 1148.87s, 3.2x that of [20]. While we were unable to match the results of COST we consider our single threaded application a reasonable baseline for measuring scalability.

The page rank algorithm consists of two fundamental steps. First the rank of a vertex is divided onto it's outgoing edges, and propagated to other vertices, second vertices sum up propagated ranks on their incoming edges to calculate their rank for a given step. As values from previous iterations are used to generate the next, there is a dependency between iterations. Any asynchronous approach to page rank must prevent out of order data access or modification for correctness.

Arguably the simplest approach to synchronizing data access and update is to use a barrier. In which all threads complete edge updates before calculating their incoming rank. For the sake of simplicity will implemented such a barrier as a synchronization mechanism for multithreaded page rank. We copied the design of map reduce for managing our threads. First n worker threads are allocated. Edges are allocated to threads in continuous chunks of $num_e dges/n$. A master iteratively issues either update edges or calculate rank commands to threads. Upon receiving commands threads perform work on their range of edges, and finally signal back to the master. Multi threading our single threaded page rank application increased it's LOC from 20 to 48. Almost all of which was coordinator code.

Based on our own best effort single threaded application, our multi-threaded framework has a COST of 0. That is, we achieve faster than single threaded performance upon introducing a second thread. Figure ?? shows our PageRank runtime vs coarse used. While we attained a cost of 0 using our own single threaded application, we required 14 to match the the processing performance of COST. An interesting aspect of our runtime, is a steady decrease in runtime after each thread has been allotted a CPU. We speculate that this increase is due to some

threads blocking when updating edges, causing the OS scheduler to execute a collocated thread.

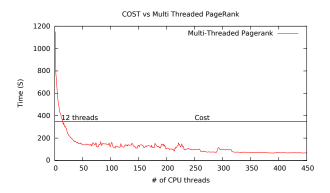


Figure 1: Page-Rank performance per core. COST benchmark reached at 14.

We measured the scalability of our multi-threaded page-rank by calculating the speedup/core against our single threaded implementation. Figure $\ref{eq:core}$ shows a near linear speedup to 60 cores.

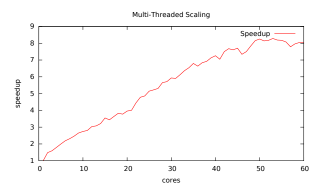


Figure 2: Scalability up to 60 cores

We compared our performance with frameworks analyzed by COST. Figure 3 plots the performance of each framework against our own for 20 iterations of PageRank. In all cases where our program was provisioned with the same number of cores as the framework (we were not able to run on 128 cores) our program outperformed the frameworks by nearly an order of magnitude.

It would be an extreme act of hubris to assume that our bench marking application endowed with a simple muti-thread manager are an order of magnitude better than existing graph frameworks. Instead we reflected on the differences between COST's single threaded application, our scalable application, and the graph processing frameworks. Both COST and our application are highly tuned

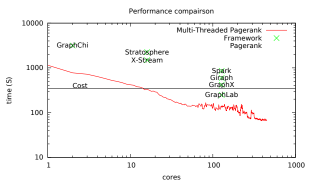


Figure 3: Page-Rank performance comparison against other frameworks. Closer to the red line is better. Our numbers past 60 core are an under approximation of runtime

for page-rank, and page-rank only. Specifically our highly tuned memory layout is designed for page rank memory access. This level of optimization is unavailable to frameworks which must have the expressiveness for the general space of graph computation. Based on this observation we anecdotal propose an additional requirement for the cost metric: Single threaded benchmarks and frameworks must implement an identical API. This addition would prevent any benefits gained from highly specific tuning, and only measure the overhead synchronization, and communication cost incurred by scalable frameworks.

3 Auto-Scaling

3.1 Why Distributed Shared Memory?

Given the results from the scalability analysis, we aim to build a system that transparently scales for practitioners that desire performance, yet want minimal changes to their single-threaded interface.

What options exist for this setting? Implementing a full MPI-style system would allow for the highest potential gains in performance, but is not generalizable and would alter the programmer's interface. One could also opt for a system such as Naiad [22] or Hadoop [26]. While simplier to reason about, this also requires significant setup and is still requires a large change in the programming interface.

Instead, we turn to the next-simplest abstraction: distributed shared memory (DSM). DSM comes with the advantage that its programming model is similar to the single machine through a key value interface, [24], and hides the complexities of the network from the developer. DSM

seems like an ideal solution for the problem presented, but has traditionally been shown to lack the required performance, and gives the developer little control over locality.

Thus, we propose BlueBridge, a locality aware DSM system. BlueBridge includes an API for access and placement of distributed shared memory, which aims to alleviate the concerns previously posed by DSM. With current trends in rack scale datacenters, disaggregated datacenters are evolving into dense memory systems with terabit-level internal bandwidth. These environments support the conceptual idea of DSM, in which a rack scale system poses as a single machine.

3.2 BlueBridge Design

BlueBridge is designed for a simple interface, that hides the complexity of shared memory from the developer. The intended users of BlueBridge are those who have a need to process large datasets, but do not want to reason about the complexities of distributed shared memory. We choose to provide an interface to BlueBridge through Python, using a shared dictionary interface. This API provides ability to perform put(key, value) and get(value) on a Python dictionary, while having all the required consistency and messaging capabilities hidden.

We simulate a DSM system by using Python 2.7, and its built-in *Manager* library. The *Manager* library allows for simple management of shared state between processes by providing interfaces to shared arrays, dictionaries and easy locking semantics.

Extending the theme from our COST investigation, we choose to target graph processing as an example use case for BlueBridge. We also use iGraph 0.6, which includes a flexible interface for graph processing in Python. iGraph performs its computation in C and is interfaced with Python, so it came to be a natural choice given the design constraints.

3.3 Experimental Setup

To evaluate the parallel graph processing of BlueBridge, we computed PageRank on three graphs, collected from the Stanford Large Network Dataset Collection. [17] These graphs will be referred to as:

- wikiVote: A Wikipedia voter network, with 7115 vertices and 103689 edges
- GrQc: A citation network of General Relativity/Quantum Cosmology papers, with 5242 vertices and 124496 edges

 condMat: A citation network of Condensed Matter papers, with 23133 nodes and 93497 edges

In all cases, we executed 20 iterations of PageRank on these three graphs, subject to a variety of programming models.

3.4 Partitioning

When processing graphs in distributed shared memory, portions of the graph are assigned to different machines, in order to balance workload and storage between machines. Pregel uses a hash of the vertex ID to assign vertices to machines, [19] which does not consider the graph structure. Given the performance and locality constraints concerning DSM, we believe that a locality-aware assignment of vertices to machines provides the opportunity for speedup, as it limits the number of network calls that must be performed during computation.

Graph partitioning occurs as a pre-processing step when performing graph processing on DSM in Blue-Brodge. In order to respect locality within DSM, our partitioning algorithm aims to provide an optimal environment for the parallel processing of graphs. We define an optimal partitioning scheme as one in which the number of vertices are roughly equal for each partition, while the number of edges that cross partitions is minimized. We leverage the METIS [16] library, an MPI implementation for parallel partitioning of large graphs, to determine an optimal partitioning for the system. METIS simply accepts an adjacency list, which can be built using iGraph library funcitons, and outputs the the graph partitioning scheme as a membership vector. METIS is built in and runs in C, and the corresponding Python binding, PyMetis [4] was used to determine the optimal graph partitioning once it is read into memory.

The three graphs explained previously all exhibit varying degrees of connectivity. GrQc can be split into 2,4 or 8 partitions with a relatively low degree of cutting, while wikiVote is much more difficult to cut. The degree of cutting for all three graphs can be observed in Table ??. The performance of the graph partitioning step was also measured and shown in, ?? and it was found that the time to partition the graph is several orders of magnitude less than the total PageRank processing time.

Applying techniques seen in [7] and [8], we use a "mirroring" technique for storing edges between partitions. For each edge that crosses partitions, only vertices on two separate partitions will be affected. The partition which contains the destination vertex will be considered the owner of the given edge. In the partition not containing the owner vertex, edges which connect to the owner

Graph	2 part.	4 part.	8 part.
GrQc	3.38%	4.93%	7.20%
condMat	6.56%	13.38%	18.69%
wikiVote	25.45%	50.05%	64.79%

Table 1: Percentage of edges cut when splitting into 2,4 or 8 partitions.

Graph	Time To Partition (avg.)
GrQc	0.027s
condMat	0.151s
wikiVote	0.050s

Table 2: Time taken to partition graph with PyMetis

vertex will instead connect to a proxy "mirror" vertex. The edges that connect to "mirrors" will be stored in shared memory, such that they can be written to and read from different machines. Owner vertices requires weights from all incoming edges and thus maintains references to the shared edges on which they depend. The partitioning procedure is shown in Figure 4.

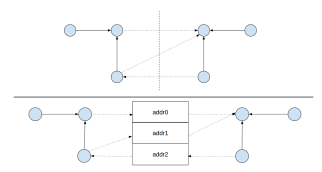


Figure 4: An example of the partitioning procedure. Shared edges are loaded into shared memory.

PageRank was evaluated on BlueBridge, both with the partitioning scheme described above, and a uniform random assignment of nodes, which models the partitioning scheme of Pregel. In a DSM environment, leveraging locality can lead to significant speedups. GrQc, which was the most easily partitioned graph, demonstrated a 10x speedup, while the wikiVote graph only showed speedups between 1-2x. These results are shown in Figure 5.

3.5 RDDs over Supersteps

Some graph processing algorithms rely on a "superstep" mechanic, where for each iteration, all vertices are synchronized together before proceeding to the next iteration.

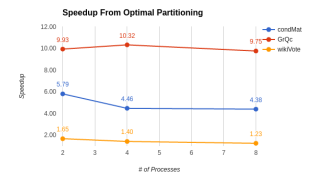


Figure 5: Speedups attained by using optimal partitioning. Speedups are relative to the performance of their corresponding random partitioning scheme.

When only a small subset of the edges in a partitioned graph are placed in distributed shared memory, creating a persistent record of edge weights on previous iterations becomes more feasible. This was explored by allocating additional addresses in the simulated shared memory for previous iterations, and storing them like resilient distributed datasets (RDDs). Access to this shared memory was performed in a publish and subscibe pattern of access. Thus, when performing a pagerank iteration, blocking only occurred on a read operation if the edge weight was not yet updated for the desired iteration.

We tackle the performance concerns of DSM by implementing this mechanism in BlueBridge. Speedups of approximately 1.5 - 2.1x wer observed, for both smart and random partitioning, varying degrees of graph partitioning, and different numbers of processes. The results of this experiment can be seen in Figure 6.

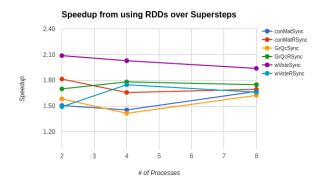


Figure 6: Speedups attained by using an RDD style of storage, instead of traditional supersteps.

Also not explored in this setting, the use of RDDs for

shared memory implies the potential for resilience in the face of machine failures. Failed instances of BlueBridge could potentially restart and pull its dependent values from the globally shared memory, preventing redundant computation.

4 Related work

The concept of using shared memory in parallel processing of graphs has been explored by Shun, who created Ligra: a graph processing framework for shared memory. [25] Ligra uses a vertex-subset level of consistency, and therefore fails to reason about the overall structure of the graph. One could use Shun's system in a fashion similar to BlueBridge by mapping the partitioning scheme onto the VertexMap design of Ligra. As Ligra processes vertices, it uses a frontier-based approach, performing vertex-centric computations in a breadth-first search matter. Ligra does no work in pre-partitioning the graph since it is not optimized for distributed shared memory and instead assumes that entire graphs can fit in a shared memory server. In settings that instead use distributed shared memory, applying Ligra's computation model would require constant swapping of memory pages, which would incur large amounts of network latency during computation.

GraphLab is another system for parallel processing of Graphs, that could also extend to a shared memory model. [18] GraphLab allows for a tunable consistency model (vertex-centric, edge-centric or full) while performing computations and does allow graph partitioning with ParMetis, the same library used by BlueBridge. However, GraphLab is a system with much more infrastructure, reading input files from a distributed storage system such as HDFS and compressing these partitions into binary files called atoms. GraphLab provides strong consistency guarantees and scales well but contains far too much overhead for the everyday scientist.

Piccolo is a distributed shared memory system that also performs pre-partitioning of data with respect to locality. Piccolo also uses a key-value table interface for defining access to shared memory. [24] Since Piccolo is also designed for DSM, locality is of high importance when accessing shared table entries, and Piccolo attempts to optimize the collocation of shared table entries. Unlike Blue-Bridge, Piccolo does not use a persistent RDD storage model, and instead relies on checkpointing of shared tables through a Chandy-Lamport distributed snapshot algorithm to provide fault tolerance.

5 Future Work

5.1 Implementing DSM

BlueBridge serves as a proof of concept for DSM, and has shown the potential merits of a DSM system for graph processing. Given the results shown in this analysis, BlueBridge should be implemented on an actual cluster, leveraging the potential of DSM. In order to do this, pages of memory in the DSM system would contain both local pages of memory, and shared global pages of memory. BlueBridge would load the external edges of the graph onto these shared global pages, and synchronization of these pages between iterations would be done by a shared state manager. One proposed implementation of this system is shown in Figure 7, in which outgoing external edge values are stored on shared pages and requested by other machines for their remote pages.

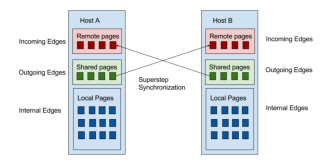


Figure 7: A proposed implementation of BlueBridge on distributed shared memory.

5.2 Offloading Work from Python

The present state of the BlueBridge prototype comes with many performance concerns. While Python provides a simple interface for the access of shared memory, use of Python's *Manager* library for access and control of shared state was unacceptable from a performance perspective. *cProfile*, the Python profiler, was used to investigate the performance of this library and it was found that up to 80-90% of the total runtime was spent in *Manager* library communication primitives, such as *send()* and *recv()*. We believe that writing a custom shared state manager to handle memory access in C would provide enormous performance benefits over the current BlueBridge prototype.

6 Conclusion

In this work we analyzed COST as a metric for analyzing distributed computation frameworks. We found COST inappropriate due to an en equality in expressiveness with these frameworks. We demonstrated the performance advantages of threading highly specific single threaded programs, and propose a framework for doing so. Further we propose a DSM framework for scaling out threaded graph computation to clusters of machines without changing a programs interface with the OS. In addition we measured the performance costs of graph processing on such a system, and evaluated the effect of graph partitioning, and RDD vs super step coordination on performance.

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