Accelerating Irregular Computations with Hardware Transactional Memory and Active Messages

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ABSTRACT

We propose Atomic Active Messages (AAM), a mechanism that accelerates irregular graph computations on both sharedand distributed-memory machines. The key idea behind AAM is that hardware transactional memory (HTM) can be used for simple and efficient processing of irregular structures in highly parallel environments. We illustrate techniques such as coarsening and coalescing that enable hardware transactions to considerably accelerate graph processing. We conduct a detailed performance analysis of AAM on Intel Haswell and IBM Blue Gene/Q and we illustrate various performance tradeoffs between different HTM parameters that impact the efficiency of graph processing. AAM can be used to implement abstractions offered by existing programming models and to improve the performance of irregular graph processing codes such as Graph500 or Galois.

Categories and Subject Descriptors

D.1.3 [Concurrent Programming]: Parallel Programming

General Terms

Performance, Design

1. INTRODUCTION

Big graphs stand behind many computational problems in social network analysis, machine-learning, computational science, and others [26]. Yet, designing efficient parallel graph algorithms is challenging due to intricate properties of graph computations. First, they are often *data-driven* and *unstructured*, making parallelism based on partitioning of data difficult to express. Second, they are usually *fine-grained* and have *poor locality*. Finally, implementing synchronization based on locks or atomics is tedious, error prone, and typically requires concurrency specialists [26].

Recent implementations of hardware transactional memory (HTM) [14] promise a faster and simpler programming for parallel algorithms. The key functionality is that complex instructions or instruction sequences execute in *isola*-

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tion and become visible to other threads atomically. Available HTM implementations show promising performance in scientific codes and industrial benchmarks [40, 36]. In this work, we show that the ease of programming and performance benefits are even more promising for fine-grained, irregular, and data-driven graph computations.

Another challenge of graph analytics is the size of the input that often requires distributed memory machines [27]. Such machines generally contain manycore compute nodes that may support HTM (cf. IBM Blue Gene/Q [36]). Still, it is unclear how to handle transactions accessing vertices on both local and remote nodes.

In this paper we propose a mechanism called Atomic Active Messages (AAM) that accelerates graph analytics by combining the active messaging (AM) model [35] with HTM. In AAM, fine units of graph computation (e.g., marking a vertex in BFS) are *coarsened* and executed as hardware transactions. While software-based coarsening was proposed in the past [18], in this paper we focus on developing high performance hardware-supported techniques to implement this mechanism on both shared- and distributed-memory machines, on establishing principles and practice of the use of HTM for the processing of graphs, and on illustrating various performance tradeoffs between different HTM parameters in the context of graph analytics. Figure 1 motivates AAM by showing the time to perform each phase in a synchronized BFS traversal using traditional fine-grained atomics and AAM based on coarser hardware transactions.



Figure 1: Comparison of the duration of an intra-node BFS traversal implemented with Blue Gene/Q fine-grained atomics and coarse hardware transactions (AAM-HTM). One transaction modifies 2^7 vertices. We use 64 threads and a Kronecker graph [23] with a power-law vertex degree distribution.

Another key insight of our work is that AAM constitutes a hierarchy of atomic actives messages that can be used to accelerate graph computations on both shared- and distributed-memory machines. We analyze this hierarchy in detail and conclude that AAM can be used to improve the

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performance of generic graph analytics tools such as Galois or Graph500. The key contributions of our work are:

- We design the generic AAM mechanism that uses state-ofthe-art HTM implementations to accelerate both sharedand distributed-memory graph computations.
- We establish the principles and practice of the use of HTM for graph computations. Specifically, we develop protocols for spawning remote/distributed hardware transactions.
- We introduce a performance model and we conduct a detailed performance analysis of AAM based on Intel Haswell HTM [40] and IBM Blue Gene/Q HTM [36] to illustrate various performance tradeoffs between different HTM parameters in the context of graph analytics. Specifically, we find optimum transaction sizes for x86 and PowerPC machines that accelerate Graph500 [30] BFS code by >100%.
- We show that AAM accelerates the processing of various synthetic and real-world graphs.

2. BACKGROUND

We now describe active messages, atomics, transactional memory, and how they are used in graph computations.

2.1 Active Messages

In the active messaging (AM) model [35] processes exchange messages that carry: the address of a user-level handler function, handler parameters, and optional payload. When a message arrives, the parameters and the payload are extracted from the network and the related handler runs at the receiver [38]. Thus, AMs are conceptually similar to lightweight Remote Procedure Calls (RPCs).

Active messages are often used to implement low-level performance-centric libraries that serve as a basis for developing higher-level libraries and runtime systems. Example libraries are Myrinet Express (MX), IBM's Deep Computing Messaging Framework (DCMF) for BlueGene/P, IBM's Parallel Active Message Interface (PAMI) for BlueGene/Q, GASNet [2], and AM++ [38].

2.2 Active Messages in Graph Computations

A challenging part of designing a distributed graph algorithm is managing its data flow. One way is to use a distributed data structure (e.g., a distributed queue) that spans all of its intra-node instances. Such structures are often hard to construct and debug [8]. A BFS algorithm that uses a distributed queue is presented in Listing 1.

```
if (source is local) Q.push(source);
while (!Q.empty()) {
   for (Vertex v : Q)
        if (v.visited == false) {
            v.visited = true;
            for (Vertex w : v.neighbors()) {Q.add(w); } } }
```



Another approach uses active messages to express the data flow of the program dynamically. When a process schedules computation for a vertex, it first checks whether it is the *owner* of this vertex. If yes, it performs the computation. Otherwise, the computation is sent in an active message to a different node for processing in a remote handler [39]. Thus, no distributed data structures have to be used. We illustrate BFS using this approach in Listing 2. struct bfs_AM_handler {
 bool operator()(const pair<Vertex, int>& x) {
 if (x.second < x.first.distance) {
 x.first.distance = x.second;
 send_active_message(x.first, x.second + 1); } };</pre>

Listing 2: Distributed BFS using active messages [39] (§ 2.2)

2.3 Atomic Operations

Atomic operations appear to the rest of the system as if they occur instantaneously. Atomics are used in lock-free graph computations to perform fine-grained updates [11, 30]. Yet, they are limited to a single word and thus require complex protocols for protecting operations involving multiple words. We now present relevant atomics:

Accumulate(*target, arg, op) (ACC): it applies an operation op (e.g., sum) to *target using an argument arg.

Fetch-and-Op(*target, arg, op) (FAO): similar to Accumulate
 but it also returns the previous value of *target.

Compare-and-Swap(*target, compare, value, *result) (CAS): if

*target == compare then value is written into *target and the function sets *result to true, otherwise it does not change *target and sets *result to false.

2.4 Transactional Memory

Transactional Memory (TM) [14] is a technique in which portions of code (*transactions*) are executed in isolation and their memory effects become visible atomically. Thus, such code portions are linearizable and easy to reason about. The underlying TM mechanism records all modifications to specific memory locations and commits them atomically. It also detects dependencies between transactions accessing the same memory locations and solves potential *conflicts* between such accesses by rolling back any changes to the data. TM can be based on software emulation [34] (software TM; STM) or native hardware support [14] (HTM).

Several vendors introduced HTM implementations: IBM, Sun, and Azul added HTM to Blue Gene/Q (BG/Q) machines [36], the Rock processor [6], and the Vega CPUs [5], respectively. Intel implemented two HTM instruction sets in the Haswell processor: Hardware Lock Elision (HLE) and Restricted Transactional Memory (RTM) that together constitute Transactional Synchronization Extensions (TSX) [40]. HLE allows for fast and simple porting of legacy lock-based code into code that uses TM. RTM enables programmers to define transactional regions in a more flexible manner than that possible with HLE [40].

There are few existing studies on STM in graph computations [16]. Using HTM in graph processing has been largely unaddressed and only a few initial works exist [7, 37].

3. ATOMIC ACTIVE MESSAGES

Atomic Active Messages (AAM) is a mechanism motivated by recent advances in deploying transactional memory in hardware. An atomic active message is a message that, upon its arrival, executes a user-specified handler called an *operator*. A *spawner* is a process (or a thread within this process, depending on the context) that issues atomic active messages. An *activity* is the computation that takes place as a result of executing an operator. Activities run speculatively, isolated from one another, and they either *commit* atomically or do not commit at all. We distinguish between operators and the activities to keep our discussion generic. To use AAM, the developer specifies the operator code that modifies elements (vertices or edges) of the graph. We use single-element operators for easy and intuitive programming of graph algorithms. Still, multiple-element *coarse* operators can be specified by experienced users. The developer also determines the structure of a vertex or an edge and defines the *failure handler*, an additional piece of code executed in certain types of algorithms (explained in § 3.2).

Our runtime system executes algorithms by exchanging messages, spawning activities to run the operator code, running failure handlers, and optimizing the execution. An activity can be *coarse*: it may execute *several* operators atomically. Note that operators are (optionally) coarsened by the developer while activities are coarsened by the runtime.

The implementation determines how activities are isolated from one another. An activity can execute as a critical section guarded by locks, or (if it modifies one element) as an atomic operation (e.g., CAS in BFS). However, we argue that in many cases running activities as hardware transactions provides the highest speedup; we support this claim with a detailed performance study in Sections 5 and 6.

3.1 Definitions and Notation

Assume there are N processes $p_1, ..., p_N$ in the system. A process p_i runs on a compute node $n_i, 1 \ge i \ge N$ and it may contain up to T threads. Then, we model the analyzed graph G as a tuple (V, E); V is a set of vertices and $E \subseteq V \times V$ is a set of edges between vertices. Without loss of generality we assume that G is partitioned and distributed using a one-dimensional scheme [4]: V is divided into N subsets V_i and every $V_i \subseteq V$ is stored on node n_i . We call process p_i the owner of every vertex $v \in V_i$ and every edge (v, w) such that $v \in V_i, w \in V$. We denote the average degree in G as \overline{d} .

3.2 Types of Atomic Active Messages

AAM accelerates graph computations that run on a single (N = 1) or multiple (N > 1) nodes. If N = 1 then messages only spawn intra-node activities. If N > 1 then a message may also be sent over the network to execute a remote activity. Now, we identify two further key criteria of categorizing messages: *direction of data flow* and *activity commits*. They enable four types of messages; each type improves the performance of different graph algorithms.

3.2.1 Direction of Data Flow

This criteria determines if an activity has to communicate some data back to its spawner. In some graph algorithms the data flow is *unidirectional* and messages are *Fire-and-Forget* (FF): they start activities that do not return any data. Other algorithms require the activity to return some data to the spawner to run a failure handler. We name a message that executes such an activity a *Fire-and-Return* (FR) message (the flow of data is *bidirectional*).

3.2.2 Activity Commits

In some graph algorithms messages belong to the type Always-Succeed (AS): they spawn activities that have to successfully commit, even if it requires multiple rollbacks or serialized execution. An example such algorithm is PageR-ank [3] where each vertex v has a parameter rank that is augmented with the normalized ranks of v's neighbors. Now, if we implement activities with transactions, then such transactions may conflict while concurrently updating the rank of the same vertex v, but finally each of them has to succeed

to add its normalized rank. The other type are *May-Fail* (MF) messages that spawn activities that may also fail ultimately and not re-execute after a rollback. An example is BFS in which two activities, which concurrently change the distance of the same vertex, conflict and only one of them succeeds. Note that we distinguish between rollbacks of activities at the algorithm level, and aborts of transactions due to cache eviction, context switches, and other reasons caused by hardware/OS. In the latter case the transaction is reexecuted by the runtime to ensure correctness.

Our criteria entail four message types: FF&AS, FF&MF, FR&AS, FR&MF. We now show examples on how each of these types can be used to program graph algorithms.

3.3 Example Case Studies

In AAM, a single graph algorithm uses only one type of atomic active messages. This type determines the form of the related operator and the existence of the failure handler. Here, we focus on the operator as the most complex part of graph algorithms. We show C-like code to implement the operator in isolation. Our implementation utilizes system annotations to mark atomic regions in C. We present the code of four single-element operators (one per message type); more examples can be found in the technical report¹. When necessary, we discuss the failure handlers. We describe multiple-element operators at the end of this section.

3.3.1 PageRank (FF & AS)

PageRank (PR) [3] is an iterative algorithm that calculates the rank of each vertex $v \in V$: $rank(v) = \frac{1-d}{|V|} + \sum_{w \in n(v)} (d \cdot \frac{rank(w)}{out.deg(w)})$. n(v) is the set of v's neighbors, d is the dump factor [3] and $out_deg(w)$ is the number of links leaving w. Depending on the operator design, PR may be either vertex-centric and edge-centric.

The pseudocode of the vertex-centric variant is presented in Listing 3. The operator increases the ranks of v's neighbors with a factor $d \cdot \frac{rank(v)}{out_{.}deg(v)}$. It also adds $\frac{1-d}{|V|}$ to rank(v). The copies of stale ranks from a previous iteration are kept and used for calculating new ranks. Assuming that each vertex v is processed by one activity, this PR variant uses AS messages: each activity has to successfully add the factors to the ranks of respective vertices (which may require serialization). Data flow is unidirectional (messages are FF) because activities do not have to communicate any results back to their spawners. Thus, the operator returns void.

```
void Operator(Vertex v) {
    v.rank += (1 - d) / vertices_nr;
    for(int i = 0; i < v.neighbors.length; i++) {
        v.neighbors[i].rank += d * v.old_rank/v.out_deg; } }</pre>
```

Listing 3: The operator in the vertex-centric PageRank variant (§ 3.3.1)

There exist other PR variants. Specifically, one can analyze incoming edges to dispose of conflicts. We will later (Section 6) show that a careful AAM design outperforms such approaches used in various codes such as PBGL.

3.3.2 Breadth First Search (FF & MF)

Breadth First Search (BFS) uses FF & MF messages. Spawners do not have to wait for any results, but some activities may fail when concurrently updating vertices using different distance values. Such a conflict is solved at the

 $^{^{1} {\}tt http://spcl.inf.ethz.ch/Research/Parallel_Programming/AAM/}$

node owning the vertex and no information has to be sent back to any of the spawners, thus the operator returns void. We present the operator pseudocode in Listing 4.

<pre>void Operator(Vertex v, int if(v.distance > new_dist)</pre>	<pre>new_distance) { {v.distance = new_dist;} }</pre>							
Listing 4: BFS operator (§ 3.3.2)								

3.3.3 ST Connectivity (FR & AS)

ST connectivity [31] determines if two given vertices (s and t) are connected. First, the algorithm marks each vertex as "white". Then, it starts two concurrent BFS traversals from s and t. Both traversals use different colors ("grey" and "green") to mark vertices as visited. Each activity returns the information on the colors of visited vertices. In case of "white" no action is taken and the operator returns false. If the found color is used by the other BFS, then s and t are connected, the operator returns true, and the runtime executes a failure handler at the spawner that terminates the algorithm. The operator is presented in Listing 5.

<pre>bool Operator(Vertex v, Color new_col) {</pre>		
<pre>if(v.color != WHITE && v.color != new_col)</pre>	return	true;
v.color = new_col; return false ; }		

Listing 5: ST Connectivity operator (§ 3.3.3)

3.3.4 Boman Graph Coloring (FR & MF)

Graph coloring proposed by Boman et al. [1] is a heuristic algorithm that minimizes the number of colors assigned to graph vertices. In this algorithm as expressed using AAM (see Listing 6), an activity changes the color or vertex v to X. Then, if any of v's neighbors has color X, either v or the neighbor has to change its color; the choice is random. Activities are spawned by MF & FR messages because multiple processes trying to update v's color may conflict and the spawners have to be notified if they need to assign new colors to v's neighbors in failure handlers.

```
int Operator(Vertex v, Color X) {
  v.Color = X;
  if(v.hasNeighborWithColor(X)) {
    //return the ID of a vertex to be recolored
    if(rand([0;1]) < 0.5) return v.neighborWithCol(X).ID;
    else return v.ID;
  } else { //NO_VERTEX_ID means no vertex is recolored
    return NO_VERTEX_ID; }</pre>
```

Listing 6: Boman graph coloring operator (§ 3.3.4)

3.4 Discussion

The introduced AAM operators modify single vertices. Thus, they enable intuitive developing and reasoning about graph computations that are also fine-grained by nature. Still, some users may want to specify coarser operators to use additional knowledge that they have about the graph structure for higher performance. Here, the user determines the number of elements to be modified in the operator and the policy of their selection (e.g., the operator may choose each vertex randomly, or try to modify elements stored in a contiguous block of memory to avoid HTM aborts).

Manual coarsening of operators may be challenging. Our runtime system automatically coarsens activities for easier AAM programming. We now discuss the implementation details of coarsening and other optimizations. While singleelement operators can be implemented with atomics or finegrained locks, we argue that a more performant approach is based on coarse transactions.

4. IMPLEMENTING ACTIVITIES

We now discuss the details of implementing activities; we skip most of the issues related to the runtime as they were properly addressed in other studies [38, 8, 39].

4.1 Implementing Activities with HTM

In this paper we advocate for using HTM to implement activities. However, locks and atomics would also match the activity semantics (atomics can implement fine activities that modify single words). We thus compared the performance of all the three mechanisms to illustrate HTM's advantages. Locks consistently entailed generally lower performance and we thus skip them due to space constraints; a brief discussion can be found in the technical report.

Transactions can implement an activity of any size. We use Intel Haswell HLE and RTM $ISAs^2$ and IBM BG/Q HTM. RTM provides two key functions: XBEGIN that starts a transaction and XEND that performs a commit. Yet, it does not guarantee progress. Thus, we repeat aborted transactions and we use exponential backoff to avoid livelock. The HTM in BG/Q automatically retries aborted transactions and it serializes the execution when the number of retries is equal to a certain value; we use the default value (10). HLE performs serialization after the first abort.

4.2 Optimizing the Execution of Activities

Two most significant optimizations applied by the runtime are *coarsening* and *coalescing* of activities. First, in the intra-node computations, the runtime coarsens activities by atomically executing more than one operator; an example is presented in Listing 7. We denote activities that are not coarse as *fine*. Coarsening amortizes the overhead of starting and committing an activity; it also reduces the amount of fine-grained synchronization. Second, activities targeted at the same remote node are sent in a single message, i.e., coalesced. This reduces the overhead of sending and receiving an atomic active message and saves bandwidth. Finally, we also use various optimizations that attempt to reduce the amount of synchronization even further. For example, the runtime avoids executing the BFS operator for each vertex by verifying if the vertex has already been visited.

```
void Activity(Vertex vertices[], int new_distance) {
  forall(Vertex v: vertices) {
    //call the BFS operator from Listing 5
    Operator(v, new_distance); } }
```

Listing 7: A BFS coarse activity (§ 4.2)

4.3 A Protocol for Distributed Activities

The ownership protocol enables activities implemented as hardware transactions that access or modify data from remote nodes. The basic idea behind the protocol is that a handler running such an activity has to first physically relocate all required vertices/edges to the memory of the node where the activity executes. This approach is dictated by the fact that a hardware transaction cannot simply send a message because it would not be able to rollback remote changes that this message caused. In addition, most HTM

 $^{^2}$ We verify the correctness of all the results to ensure that the limitations of TSX [15] do not affect our evaluation and the conclusions drawn.

implementations prevent many types of operations (e.g., system calls) from being executed inside a transaction [36].

Our protocol assumes that each graph element has an ownership marker that can be modified atomically by any process. Each marker is initially set to a value \perp different from any process id. When a transaction from a node n_i accesses a remote graph element, it aborts and the runtime uses CAS or a different mechanism (e.g., an active message) to set the marker of this element to the id of process p_i . If the CAS succeeds, the marked element is transferred to node n_i and the transaction restarts. If the CAS fails, the handler sets all previously marked elements to \perp and backs off for a random amount of time. If a local transaction attempts to access a marked element, it aborts. This mechanism is repeated until all remote elements are cached locally. Finally, after the transaction succeeds, the elements are sent back to their original nodes and their markers are set to \perp .

5. PERFORMANCE MODEL & ANALYSIS

We now introduce a simple performance model that shows the tradeoffs between atomics and HTM. Then, we analyze the performance of AAM and answer the following research questions: (1) what are HTM's advantages over atomics for implementing AAM activities, (2) what are performance tradeoffs related to various HTM parameters, and (3) what are the optimum transaction sizes for analyzed architectures that enable highest speedups in selected graph algorithms.

5.1 Experimental Setup

We compile the code with gcc-4.8 (on Haswell) and with IBM XLC v12.1 (on BG/Q). We use the following machines:

ALCF BG/Q Vesta (BGQ) is a supercomputing machine where each compute node contains 16 1.6 GHz PowerPC A2 4-way multi-threaded cores, giving the total of 64 hardware threads per node. Each core has 16 kB of L1 cache. Every node has 32 MB of shared L2 cache and 16 GB of RAM. Nodes are connected with a 5D proprietary torus network. This machine represents massively parallel supercomputers with HTM implemented in the shared last-level cache.

Trivium V70.05 (Has-C) is a commodity off-the-shelf server where the processor (Intel Core i7-4770) contains 4 3.4 GHz Haswell 2-way multi-threaded cores, giving the total of 8 hardware threads. Each core has 32 KB of L1 and 256 KB of L2 cache. The CPU has 8 MB of shared L3 cache and 8 GB of RAM. This option speaks for commodity computers with HTM operating in private caches.

Greina (Has-P) is a high-performance cluster that contains two nodes connected with InfiniBand FDR fabric. Each node hosts an Intel Xeon CPU E5-2680 CPU with 12 2way 2.50GHz multi-threaded cores; the total of 24 hardware threads. Each core contains 64 KB of L1 and 256 KB of L2 cache. The CPU has 30 MB of shared L3 cache and 66 GB of RAM. This machine represents high-performance clusters deploying HTM in private caches.

5.2 Considered Hardware Mechanisms

For Haswell we compare the following mechanisms: RTM (Has-RTM), HLE (Has-HLE), GCC (Has-CAS), __sync_bool_compare_and_swap GCC and __sync_add_and_fetch (Has-ACC). We select CAS and ACC because they can be used in miscellaneous graph codes such as BFS (a FF&MF algorithm), PR (a FF&AS algorithm), and ST Connectivity (a FR&AS algorithm) [30]. For BG/Q we analyze: IBM XLC __compare_and_swap (BGQ-CAS) and GCC __sync_add_and_fetch (BGQ-ACC). We compare two modes of HTM in BG/Q: the *short running mode* [36] (BGQ-HTM-S) that bypasses L1 cache and performs better for shorter transactions, and the *long running mode* [36] (BGQ-HTM-L) that keeps speculative states in L1 and is better suited for longer transactions [36].

5.3 Performance Model

Our performance model targets graph processing and we argue in terms of activities and accessed vertices. We predict that an activity implemented as a transaction that modifies one vertex is more computationally expensive than an equivalent single atomic. Yet, the transactional overheads (starting and committing) may be amortized with coarser transactions and respective activities would outperform a series of atomics for a certain number of accessed vertices.

We now model the performance to determine the existence of crossing points; out model includes both the execution of the operations and fetching the operands from the memory. The total time to execute an activity that modifies N vertices (using either atomics or HTM) can be modeled with a simple linear function with N as the argument. We denote the slope and the intercept parameters of a function that targets atomics as \mathcal{A}_{AT} and \mathcal{B}_{AT} ; the respective parameters for HTM are \mathcal{A}_{HTM} and \mathcal{B}_{HTM} . We predict that $\mathcal{B}_{HTM} > \mathcal{B}_{AT}$ due to high transactional overheads. On the contrary, we conjecture that $\mathcal{A}_{HTM} < \mathcal{A}_{AT}$ because HTM overheads will grow at a significantly lower rate (determined by accesses to the memory subsystem) than that of atomics.

We illustrate the model validation for CAS in Figure 2; we plot only the results for RTM on Has-C and the long mode HTM on BGQ because all the other results differ marginally and follow similar performance patterns. We use linear regression to calculate \mathcal{A}_{AT} , \mathcal{B}_{AT} , \mathcal{A}_{HTM} , and \mathcal{B}_{HTM} . The analysis indicates that the model matches the data. While a more extended model is beyond the scope of this paper, our analysis illustrates that it is possible to amortize the transactional overhead with coarser activities. We now proceed to a performance analysis that illustrates various tradeoffs between respective HTM parameters.



5.4 Single-vertex Activities

First, we analyze the performance of single-vertex activities. The results are illustrated in Figure 3. Has-C and Has-P follow similar performance trends and we show only the former (denoted as Has); we thus illustrate the results for both a multicore off-the-shelf system and a manycore high performance machine (BGQ).



624 (BGO)

		Aborts due to:						
		Memory conflicts	Buffer overflows	Other reasons				
10 ops	Has-RTM BGQ-HTM-L BGQ-HTM-S	2 802 1,118	2 3 46	0 1 180				
100 ops	Has-RTM BGQ-HTM-L BGQ-HTM-S	$2 \\ 1,539 \\ 2,242$	2 5 13	0 1 2				

(c) Marking a vertex as visited: details of the aborts in the HTM implementations (\S 5.4.1).

	Aborts due to:						
	Memory conflicts	Buffer overflows	Other reasons				
^{SC} Has-RTM O BGQ-HTM-L BGQ-HTM-S	$ \begin{array}{r} 1,520 \\ 624 \\ 623 \end{array} $	1 62 62	$\begin{array}{c} 0 \\ 614 \\ 613 \end{array}$				
^{SCI} Has-RTM BGQ-HTM-L BGQ-HTM-S	$\begin{array}{c} 18,952 \\ 6,374 \\ 6,392 \end{array}$	33 637 639	$\begin{array}{c} 0 \\ 6,360 \\ 6,380 \end{array}$				

(f) Incrementing a vertex' rank: details of the aborts in

(a) Marking a vertex as visited 10 times (§ 5.4.1). (b) Marking a vertex as visited 100 times (§ 5.4.1). HTM implementations (§ 5.4.1).

2.1

6 3k

Numbers are total aborts

433

5k

568

18

ar data noint

(d) Incrementing a vertex' rank 10 times (§ 5.4.2). (e) Incrementing a vertex' rank 100 times (§ 5.4.2).

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Dtal 0.010

0.00

time [

Figure 3: The analysis of the performance of intra-node activities implemented with atomics and HTMs (\S 5.4). Figures 3a and 3b illustrate the time it takes to mark a vertex as visited. Numbers in figures are sums of HTM aborts for a given datapoint (we report values for T = 4 for Has/BGQ; we also plot numbers for T = 8 (Has) and T = 64 (BGQ) to illustrate the numbers of aborts generated by all the supported hardware threads). Indicates the point where the latency of Haswell atomics stops to grow. Table 3c shows the distribution of the reasons of aborts for T = 64 (BGQ) and T = 8 (Haswell). We skip Has-HLE as it does not provide functions to gather such statistics. A similar performance analysis for incrementing the rank of a vertex is presented in Figures 3d-3e and Table 3f.

Threads per node (T)

5.4.1 Activity 1: Marking a Vertex as Visited

1e+0

ဖ်ာ^{1e_0}

1e-02

1e-03

1e-04

10 (HLE

43 (RTM)

4 8 16 Threads per node (T)

Numh

Total

Here, each thread uses a CAS or an equivalent HTM code to atomically mark a single vertex; see Fig. 3a-3b, Table 3c. This activity may be used in BFS or any other related algorithm such as Single Source Shortest Path (SSSP). We analyze a negligibly contended scenario that addresses sparse graphs (Fig. 3a; a vertex is marked 10 times to simulate low contention) and a more contended case for dense graphs with high \bar{d} (Fig. 3b; a vertex is marked 100 times). We repeat the benchmark 1000 times and derive the average total time to finish the operations.

total aborts per data poir

30

Figure 3a shows that Has-CAS finishes fastest and is slightly impacted by the increasing $T \ (\approx 50\%$ of difference between the results for T = 4 and T = 8). This is because Has-CAS locks the respective cache line, causing contention in the memory system. Both Has-RTM and Has-HLE have 1.5-3x higher latency than Has-CAS, with Has-RTM being 5-15% faster than Has-HLE. Their performance is not influenced by the increasing T as they rarely abort. Then, BGQ-HTM-S and BGQ-HTM-L are more sensitive to the growing T and their performance drops 11x when switching from T = 1 to T = 64due to expensive aborts. As expected, BGQ-HTM-S is faster than BGQ-HTM-L, but as T increases it also aborts more frequently, and becomes $\approx 2x$ less efficient (T = 64) with 37.5% more aborts. BGQ-CAS is least affected by the increasing T.

Figure 3b shows that Has-RTM, BGQ-CAS, BGQ-HTM-S, and BGQ-HTM-L follow similar performance patterns when threads access the vertex 100 times. The performance of Has-HLE drops rapidly as it always performs the costly serialization after the first abort and thus forces all other transactions to abort. The latency of Has-CAS grows proportionally to the contention in the memory system. It stabilizes at T = 8 as for T > 8 no more operations can be issued in parallel.

5.4.2 Activity 2: Incrementing Vertex Rank

This activity can be used to implement PR. Here, each thread increments the rank of a single vertex 10 times (Figure 3d) and 100 times (Figure 3e) with an ACC or an equivalent HTM code; see Table 3c for details. The most significant difference between the previous and the current benchmark is that the total time and the number of aborts of Has-RTM and Has-HLE grow very rapidly in both scenarios as T scales. This is because in the HTM implementation of ACC, the rank of the vertex is modified by each transaction, generating a considerable number of conflicts and thus aborts. On the contrary, the HTM implementation of CAS generates few memory conflicts: once the vertex id is swapped, other threads only read it and do not modify it. BGQ-HTM-S and BGQ-HTM-L follow a similar trend, with $\approx 3x$ more aborts than in the previous CAS benchmark.

Discussion We present the details of the above analysis in Tables 3c and 3f. We show that the considered singlevertex activities are in most cases best implemented with atomics. HTM is faster only in processing dense graphs with algorithms that use CAS (e.g., BFS) on Haswell. We also conclude that while atomic CAS is more expensive than ACC, HTM implementation of single ACC is slower ($\approx 100x$ for RTM and $\approx 10x$ for BG/Q HTM) than that of CAS as it generates more memory conflicts and thus costly aborts.

5.5 Multi-vertex Activities

The performance analysis of single-vertex intra-node activities illustrates that in most cases a transaction modifying a single vertex is slower than an atomic operation. We now analyze if it is possible to amortize the cost of starting and aborting transactions by enlarging their size, i.e., *coarsening.* This section extends the model analysis (§ 5.3) by



Figure 4: (§ 5.5) The analysis of the performance of Graph500 OpenMP BFS implemented with hardware transactions on BGQ (Figures 4a-4d), Has-C (Figures 4e-4h), and Has-P (Figures 4i-4l). In each figure we vary the size of the transactions M (i.e., the number of vertices visited). We also present the results for BFS implemented with atomics (horizontal lines). For BGQ, the percentages indicate the ratios of the numbers of serializations caused by reaching the maximum possible number of rollbacks to the numbers of all the aborts. For Haswell, the percentages are the ratios of the aborts due to HTM buffer overflows to all the aborts. Bolded numbers indicate the points with the minimum runtime per figure. We do not include the numbers for Haswell HLE because it does not enable gathering more detailed statistics [40]. Figures 4d, 4h, and 4l present the total number of HTM events (transactions, aborts, buffer overflows) for every analyzed M.

introducing effects such as memory conflicts or HTM buffer overflows. We perform the analysis for the highly-optimized OpenMP BFS Graph500 code [30]. We modify the code so that a single transaction atomically visits M vertices and we evaluate the modified code for M between 1 and 320 with the interval of 16. We present the results for three scenarios: a single-threaded execution (T = 1 for BGQ, Has-C, and Has-P), a single thread per core (T = 16 for BG/Q, T = 4 for Has-C, and T = 12 for Has-P), and a single thread per SMT hardware resource (T = 64 for BG/Q, T = 8 for Has-C, and T = 24 for Has-P). Other sets of parameters are illustrated in the technical report. We use Kronecker graphs [23] with the power-law vertex degree distribution and $|V| = 2^{20}$, $|E| = 2^{24}$. The results are shown in Figure 4.

5.5.1 BG/Q (Supercomputer)

Figures 4a-4d present the analysis for BG/Q. For T = 1 the runtime of both HTM-Long-Mode and HTM-Short-Mode is always higher than that of Atomic-CAS and it decreases ini-

tially with the increasing M because higher M reduces the number of transactions required to process the whole graph and thus amortizes the overhead of starting and committing transactions. The runtime of HTM-Short-Mode becomes higher with the increasing M > 32 because this mode is better suited for short transactions. The runtime of HTM-Long-Mode decreases as expected and it stabilizes at $M \approx 240$. For T = 16, initially the runtime drops rapidly for both HTM modes to reach the minimum (obtained for $M_{min} = 80$ in HTM-Short-Mode). Again, this effect is caused by amortizing the overheads of commits/aborts with coarser transactions. Beyond M_{min} the runtime slowly increases with M due to more frequent serializations caused by reaching the maximum number of allowed rollbacks (BGQ does not enable gathering more detailed statistics but we predict that these serializations are due to the higher number of HTM buffer overflows and memory conflicts). HTM-Long-Mode is never more efficient than Atomic-CAS. HTM-Short-Mode becomes more effi-



Figure 5: (§ 5.5 & § 5.6) The comparison of the percentage of the reasons of aborts on Has-C and Has-P (Figures 5a and 5b) and the analysis of the performance of inter-node activities on BG/Q and Has-P. The results for BG/Q are following: marking a remote vertex as visited (Figures 5c and 5d), incrementing a vertex' rank (Figures 5e and 5f), and executing distributed transactions (Figure 5i). The results for Haswell are following: marking a remote vertex as visited (Figure 5g) and incrementing a vertex' rank (Figure 5h).

cient than Atomic-CAS for M = 32 and achieves the speedup of 1.11 at $M_{min} = 80$. A similar performance pattern can be observed for T = 64; this time $M_{min} = 144$ in HTM-Short-Mode with the speedup of 1.49 over Atomic-CAS. The runtime becomes dominated by aborts for M > 144; cf. Figure 4d with more detailed numbers of aborts.

5.5.2 Has-C (Commodity Machine)

The results of the analysis for Has-C are presented in Figures 4e-4h. In each scenario (T = 1, 4, 8) the performance of both HTM-RTM and HTM-HLE decreases with increasing M. Several outliers are caused by disadvantageous graph data layouts that entail more aborts due to the limited associativity of L1 cache (8-way associative cache) that stores speculative states in Haswell [40]. We perform a more detailed analysis for $M \in \{1, ..., 16\}$ to find out that $M_{min} = 2$. HTM-RTM becomes less efficient than HTM-HLE at M≈200 because the cost of serializations due to the HTM buffer overflows dominates the runtime of HTM-RTM beyond this point (serializations in HTM-HLE are implemented in hardware [40], while in HTM-RTM they have to be implemented in software).

5.5.3 Has-P (High-Performance Server)

The analysis of Has-P is presented in Figures 4i-4l. The performance trends are partially similar to the observations for Has-C; especially for lower thread counts ($T \leq 4$). A distinctive feature is a significantly lower number of HTM buffer overflows than in Has-C. To gain more insight we performed an additional analysis to compare the number of memory conflicts and HTM buffer overflows with varying T for fixed M = 2. We present the results in Figures 5a-5b. Surprisingly, we observe Has-C has significantly more buffer overflows than memory conflicts for the increasing T; a reverse trend is observed on Has-P. This interesting insight may help improve the design of future HTM architectures. Discussion Our analysis shows that RTM is more vulnerable to aborts than BG/Q HTM. The difference between the number of transactions and aborts never drops below

25% for HTM in BG/Q for any analyzed M (cf. Figure 4d), while for RTM this threshold is achieved for M = 144 (Has-C). Another discovery is that Has-P is only marginally impacted by buffer overflows (<1% of all the aborts for T = 24and M = 320). On the contrary, aborts in Has-C are dominated by HTM buffer overflows that constitute more than 90% of all the aborts for M > 64. The only exception are the data points where the number of overflows drops rapidly as aborts become dominated by the limited L1 cache associativity (a similar effect is visible for Has-P). This effect is not visible in BG/Q because it stores its speculative states in its L2 16-way associative cache [36], while both Has-P and Has-C have 8-way associative L1s.

We conclude that the coarsening of transactions provides significant speedups (up to 1.51) over the Atomic-CAS baseline on BGQ and Has-C; Has-P does not offer any speedups due to the overheads generated by memory conflicts. We find the following optimum transaction sizes for PowerPC in BG/Q: $M_{min} = 80$ (T = 16), $M_{min} = 144$ (T = 64). For x86 (Has-C) $M_{min} = 2$ for $T \in \{4,8\}$. We present M_{min} for the remaining values of T in the technical report. We will use these values in Section 6 to accelerate Graph500 [30] for different types of graphs.

5.6 Activities Spawned on a Remote Node

We now analyze the performance of activities spawned on a remote node. We implement such activities as hardware transactions triggered upon receiving an atomic active message. We again test both the long and the short running mode (on BG/Q) and RTM/HLE (on Haswell). To reduce the overhead of sending and receiving an atomic active message and save bandwidth, we use *activity coalescing*: activities flowing to the same target are sent in a single message.

We run the benchmarks on BG/Q and Greina (Has-P); we skip Has-C because the Trivium server is not a distributed memory machine. On BG/Q, we compare inter-node activities to optimized remote one-sided CAS and ACC atomics provided by the generic function PAMI_Rmw in the IBM PAMI



(a) The performance of BFS on BG/Q for Kronecker graphs. Figure 6: (§ 6.1) The overview of the performance of intra-node Graph500 BFS implemented with atomics (Graph500-BGQ, Graph500-Haswell), AAM RTM (AAM-Haswell), and the short mode BG/Q HTM (AAM-BGQ). We vary |V| and \bar{d} ; T = 64 (on BG/Q) and T = 8 (on Has-C).

communication library [19]. On Has-P we compare activities to remote atomic operations provided by MPI-3 RMA [29] implemented over the InfiniBand fabric. We evaluate the performance of marking a remote vertex as visited (addressing distributed BFS computations) and incrementing the rank of a remote vertex (addressing distributed PageRank).

5.6.1 BG/Q (Supercomputer)

We first measure the time it takes a process p_i to mark 2^{13} vertices stored on a node n_j as visited (targeting distributed BFS). The results are presented in Figure 5c. Without coalescing, HTM activities (Inter-node-HTM-L for the long and Inter-node-HTM-S for the short mode) are \approx 5x slower than PAMI atomics (Inter-node-CAS). Still, for $C_{cross} = 16$ Inter-node-HTM-S becomes more efficient. Second, we scale the number of nodes N. Figure 5d shows the time to mark a vertex stored in process p_N 's memory by N-1 other processes. We use the short HTM mode. Coalesced AAMs (Inter-node-HTM-C) outperform Inter-node-CAS \approx 5-7 times.

We also evaluate an inter-node activity that increments the rank of a vertex (targeting distributed PR). We perform analogous benchmarks as for the remote CAS; we present the results in Figures 5e-5f. Implementing ACC using HTM again generates costly aborts that dominate the runtime; however coalescing enables a speedup of $\approx 20\%$ (for the short HTM running mode) over highly optimized PAMI atomics.

5.6.2 Has-P (High-Performance Server)

Here, we test the performance of inter-node activities implemented on Has-P. Our testbed has two nodes, thus we only vary C. We present the results in Figures 5g (CAS) and 5h (ACC). Setting C = 2 enables AAM to outperform remote atomics provided by MPI-3 RMA.

5.7 Distributed Activities

Finally, we test the ownership protocol for executing activities that span multiple nodes (see Figure 5i for BGQ results). Each process issues x transactions; each transaction marks a local and b remote randomly selected vertices. We compare four scenarios: 0-1 ($x = 10^3, a = 5, b = 1$), 0-2 ($x = 10^4, a = 5, b = 1$), 0-3 ($x = 10^3, a = 7, b = 3$), and 0-4 ($x = 10^4, a = 7, b = 3$). We measure the total time to execute transactions. 0-1 finishes fastest, 0-3 is slower as more remote vertices have to be acquired. 0-2 and 0-4 follow the same performance patterns; additional overheads are due to the backoff scheme. If no time is spent on backoff, then the protocol may livelock and may make no progress. We conclude that AAM can be used in various environments (e.g., IBM networks or InfiniBand) to enable remote transactions and to accelerate distributed processing.

6. EVALUATION

We now use AAM to accelerate the processing of large Kronecker [23] and Erdős-Renyi [10] (ER) graphs with different vertex distributions (power-law, binomial, Poisson). We also evaluate real-world SNAP graphs³. We evaluate BFS and PR because they are the basis of various data analytics benchmarks such as Graph500 and because they are proxies of many algorithms such as Ford-Fulkerson.

6.1 BFS: Massively-Parallel Manycores

We first evaluate the speedup that AAM delivers in highly-parallel multi- and manycore environments.

Comparison Baseline: Here, we use the OpenMP Graph500 highly optimized reference code [30] (Graph500-BGQ, Graph500-Haswell) based on atomics as the comparison baseline. The baseline applies several optimizations; among others it reduces the amount of fine-grained synchronization by checking if the vertex was visited before executing an atomic.

We compare the Graph500 baseline with the coarsened variants that use the short mode HTM in BG/Q (AAM-BGQ) and RTM in Haswell (AAM-Haswell). We only use Has-C (denoted as Haswell) because it provides higher speedups over atomics than Has-P as we show in Figure 4. The long mode and HLE are omitted as they follow similar performance patterns and vary by up to 10%. We set T = 64 (for BG/Q) and T = 8 (for Haswell) for full parallelism.

6.1.1 Processing Kronecker Power-Law Graphs

Here, we use the results of the analysis in Section 5 and set $M_{min} \in \{2, 80, 144\}$ for the most advantageous size of transactions on BG/Q and Haswell. We present the results in Figure 6. We scale |V| from 2^{20} to 2^{28} , and we use $\bar{d} \in \{1, 2, ..., 256\}$; highest values generate graphs that fill the whole available memory. For BG/Q, AAM-BGQ outperforms Graph500-BGQ by up to 102% for a graph with ≈ 2 millions vertices and $\bar{d} = 4$. For higher \bar{d} AAM-BGQ becomes comparable to Graph500-BGQ. This is because adding more edges for fixed |V|generates more transactions that conflict and abort more often. For Haswell, AAM consistently outperforms Graph500 by up to 27%. The speedup does not change significantly when increasing \bar{d} . This is because we use smaller transac-

 $^{^3\}mathsf{Available}$ at https://snap.stanford.edu/data/index.html.

Input graph properties					BG/Q analysis		Haswell analysis						
Type	ID	Name	V	E	$ \begin{array}{c} S \text{ over } \text{g500} \\ (M = 24) \end{array} $	M	$\begin{array}{c} S \ { m over} \\ { m g500} \end{array}$	$ \begin{vmatrix} S & \text{over g500} \\ (M = 2) \end{vmatrix} $	$ S \text{ over Galois} \\ (M = 2) $	M	$\begin{array}{c} S \ { m over} \\ { m g500} \end{array}$	S over Galois	S over HAMA
Comm. networks (CNs)	$\begin{vmatrix} cWT \\ cEU \end{vmatrix}$	wiki-Talk email-EuAll	$_{ m 2.4M}^{ m 2.4M}$	5M 420k	$2.82 \\ 3.67$	$\begin{vmatrix} 48 \\ 32 \end{vmatrix}$	$\begin{array}{c} 3.35\\ 4.36\end{array}$	0.91 0.76	$\begin{array}{c} 1.22 \\ 0.88 \end{array}$	$\begin{vmatrix} 6\\ 4 \end{vmatrix}$	$0.96 \\ 0.97$	$\begin{array}{c} 1.28 \\ 1.12 \end{array}$	$\begin{array}{c} 344 \\ 1448 \end{array}$
Social networks (SNs)	sLV sOR sLJ sYT sDB sAM	soc-LiveJ. com-orkut com-lj com-youtube com-dblp com-amazon	4.8M 3M 4M 1.1M 317k 334k	69M 117M 34M 2.9M 1M 925k	$1.44 \\ 1.22 \\ 1.44 \\ 1.67 \\ 1.33 \\ 1.14$	$\begin{vmatrix} 12 \\ 20 \\ 12 \\ 8 \\ 8 \\ 8 \\ 8 \end{vmatrix}$	$1.56 \\ 1.27 \\ 1.54 \\ 1.84 \\ 1.80 \\ 1.62$	$1.05 \\ 1.06 \\ 1.03 \\ 0.96 \\ \approx 1 \\ 1.04$	$1.1 \\ 0.69 \\ 1.03 \\ 1.1 \\ 2.5 \\ 1.64$	$\begin{vmatrix} 3 \\ 4 \\ 4 \\ 5 \\ 2 \\ 2 \end{vmatrix}$	1.07 1.13 1.04 0.98 ≈ 1 1.04	$1.12 \\ 0.74 \\ 1.04 \\ 1.11 \\ 2.53 \\ 1.64$	$> 10^4$ $> 10^4$ 603 670 2160 1426
Purchase network (PNs)	pAM	amazon0601	403k	3.3M	1.45	8	1.91	≈1	1.25	3	1.03	1.30	618
Road networks (RNs)	rCA rTX rPA	roadNet-CA roadNet-TX roadNet-PA	1.9M 1.3M 1M	5.5M 3.8M 3M	$pprox 1 \ pprox 1 \ pprox 1 \ pprox 1 \ pprox 1$	$\begin{vmatrix} 2\\ 2\\ 2 \end{vmatrix}$	$1.59 \\ 1.53 \\ 1.52$	$ \begin{array}{c c} 1.33 \\ 1.29 \\ \approx 1 \end{array} $	1.74 1.89 2.00	8 6 9	$1.38 \\ 1.42 \\ 1.07$	$1.80 \\ 2.08 \\ 2.16$	$> 10^4$ $> 10^4$ $> 10^4$
Citation graphs (CGs)	ciP	cit-Patents	$3.7\mathrm{M}$	$16.5 \mathrm{M}$	1.16	8	1.57	1.01	1.26	2	1.01	1.26	1875
Web graphs (WGs)	wGL wBS wSF	web-Google web-BerkStan web-Stanford	875k 685k 281k	5.1M 7.6M 2.3M	$1.78 \\ 1.91 \\ 1.89$	$\begin{vmatrix} 12\\24\\24 \end{vmatrix}$	$2.08 \\ 1.91 \\ 1.89$	0.98 0.93 0.98	$1.26 \\ 1.31 \\ 1.54$	6 5 5	$1.06 \\ 1.07 \\ 1.07$	$1.35 \\ 1.40 \\ 1.58$	$365 \\ 755 \\ 1077$

Table 1: (§ 6.1.2) The performance of AAM for real-world graphs. S and g500 denote speedup and Graph500. \approx 1 indicates that the given $S \in (0.99; 1.01)$.

tions in AAM-Haswell (M=2) than in AAM-BGQ (M=144) and thus they do not incur considerably more memory conflicts when \bar{d} is increased.

6.1.2 Processing Real-World Graphs

Next, we evaluate AAM for real-world graphs (see Table 1). For this, we extend Graph500 so that it can read graphs from a file. We selected directed/undirected graphs with |V| > 250k that could fit in memory and we excluded graphs that could not easily be loaded into Graph500 framework (e.g., amazon0505).

BlueGene/Q: The tested graphs are generally sparser than the analyzed Kronecker graphs. We discovered that the optimum M is smaller than 144 (we set it to 24). This is because in dense graphs more data is contiguous in memory and thus can be processed more efficiently by larger transactions. The results show that graphs with similar structure entail similar performance gains. The highest S (speedup) is achieved for CNs (up to 3.67) and WGs (up to 1.91). SNs. PNs. and CGs offer moderate S (1.14-1.67). RNs entail no significant change in performance. We also searched for optimum values of M for specific graphs; this improves S across all the groups. The results indicate that respective groups have similar optimum values of M. The differences are due to the structures of the graphs that may either facilitate coarsening and reduce the number of costly aborts (CNs and WGs) or entail more significant overheads (RNs).

Haswell: We compare AAM to several state-of-theart graph processing engines: Galois [18] (represents runtime systems that exploit amorphous data-parallelism), SNAP [22] (represents network analysis and data mining libraries), and HAMA [33]⁴ (an engine similar to Pregel [27] that represents Hadoop-based BSP processing engines). We do not evaluate these engines on BG/Q due to various compatibility problems (e.g., BG/Q does not support Java required by HAMA). BFS in Galois only returns the diameter. We modified it (with fine locks) so that it constructs a full BFS tree, analogously to AAM and Graph500.

First, we set M = 2. While AAM is in general faster than Graph500 (up to 33% for rCA), several inputs entail longer AAM BFS traversals. AAM is up to a factor of two

faster than Galois but is slower for two inputs (cEU and sOR). There is some diversity in the results because AAM on Haswell is significantly more sensitive to small changes of M than on BG/Q. Thus, we again searched for the optimum M for each input separately which resulted in higher AAM's speedups. The performance of HAMA and SNAP is generally much lower than AAM (results for SNAP are consistently worse than for HAMA and we exclude them due to space constraints). HAMA suffers from overheads caused by the underlying MapReduce architecture and expensive synchronization. The analyzed real-world graphs have usually high diameters (e.g., 33 for sAM) and thus require many BSP steps that are expensive in HAMA. This is especially visible for RNs that have particularly big diameters (554 for rCA) and accordingly long runtimes. As we will show in the next section, processing Kronecker graphs with lower diameters reduces these overheads. We also investigated SNAP and we found out that it is particularly inefficient for undirected graphs and it does not efficiently use threading. Our final discovery is that, similarly to BG/Q, respective groups of graphs have similar optimum values of M.

6.1.3 Evaluating the Scalability of AAM

Finally, we evaluate the scalability of AAM by varying T. The results are presented in Figure 7a (BG/Q) and 7b (Haswell). We use a Kronecker graph with 2^{21} vertices and 2^{24} edges. We vary T between 1 and the number of available hardware threads. The BG/Q results indicate that AAM utilizes onnode parallelism more efficiently than Graph500. For Haswell, the performance patterns for AAM and Graph500 are similar; both frameworks deliver positive speedups for any T and outperform other schemes by ≈ 20 -50% (Galois) and ≈ 2 orders of magnitude (HAMA). We skip SNAP for clarity; it is consistently 2-3x slower than HAMA.

6.2 PR: Distributed Memory Machines

As the last step, we provide an initial large-scale evaluation of AAM in a distributed environment. We select PR to illustrate that expensive and numerous aborts generated by the HTM implementation of ACC (cf. \S 5.4.2) can be amortized with the coalescing of activities. We compare AAM to a version of Parallel Boost Graph Library (PBGL) [11] based on active messages. The utilized variant of PBGL

 $^{^4\}mathrm{Update}$ (31.07.15): we use HAMA 0.6.4



(a) Scalability of AAM; BG/Q. (b) Scalability of AAM; Haswell. (c) PageRank, ER = 0.0005. (d) PageRank, ER = 0.0005. (e) PageRank, ER = 0.005. Figure 7: The analysis of the performance of: BFS (when varying T; § 6.1.3 and Figures 7a-7b) and distributed PR (§ 6.2, Figures 7c-7e).

applies various optimizations; for example it processes incoming edges to reduce the amount of synchronization and to limit the performance overheads caused by atomics. We run the benchmarks on BG/Q to enable large-scale evaluation. We use ER graphs with the probability parameter $ER \in \{0.005, 0.0005\}$ and the number of vertices up to 2^{23} . PBGL does not support threading, we thus spawn multiple processes per node and an equal number of threads in AAM; we scale T until PBGL fills in the whole memory.

The results of the analysis are presented in Figure 7. We scale N (Figure 7c), T (Figure 7d), and $|V_i|$ (Figure 7e). In each scenario AAM outperforms PBGL \approx 3-10 times thanks to the coalescing of activities and more efficient utilization of intra-node parallelism.

7. RELATED WORK AND DISCUSSION

The challenges connected with the processing of graphs are presented by Lumsdaine et al. [26]. Example frameworks for parallel graph computations are Pregel [27], PBGL [11], HAMA [33], GraphLab [24], and Spark [41]. There exist several comparisons of various engines [32, 12, 25, 9]. AAM differs from these designs as it is a mechanism that can be used to implement abstractions and to accelerate processing engines. It uses HTM to reduce the amount of synchronization and thus to accelerate graph analytics.

GraphBLAS [28] is an emerging standard for expressing graph computations in terms of linear algebra operations. AAM can be used to implement the GraphBLAS abstraction and to accelerate the performance of graph analytics based on sparse linear algebra computations.

The Galois runtime [17] optimizes graph processing by coarsening fine graph updates. AAM can be integrated with Galois. In AAM, we focus on scalable techniques for implementing coarsening with HTM. First, we provide a detailed performance analysis of HTM for graph computations, a core paper contribution. Instead, Galois mostly addresses locking [18]. Second, contrary to Galois, AAM targets both shared- and distributed-memory systems. Third, our work performs a holistic extensive performance analysis of coarsening. Instead, coarsening in Galois is not evaluated on its own. We conclude that AAM's techniques and analysis can be used to accelerate the Galois runtime.

Active Messages (AM) were introduced by Eicken et al. [35]. Various AM implementations were proposed [8, 38, 39, 19, 2]. Our work enhances these designs by combining AM with HTM. We illustrate how to program AAM and we conduct an extensive analysis to show how to tune AAM's performance on state-of-the-art manycore architectures.

Transactional memory was introduced by Herlihy et al. [14]. Several implementations of HTM were introduced, but their performance was not extensively analyzed [40, 36, 7, 6]. Yoo et al. [40] present performance gains from using Haswell HTM in scientific workloads such as simulated annealing. Our analysis generalizes these findings, proposes a simple performance model, and provides a deep insight into the performance of both BG/Q and Haswell HTM for a broad range of transaction sizes and other parameters in the context of data analytics.

We envision that the potential of AAM could be further expanded by combining it with some ideas related to code analysis. For example, one could envision a simple compiler pass that pattern-matches each single-vertex transaction against the set of atomic operations to transform it if possible to accelerate graph processing. However, such an analysis is outside the scope of this paper.

Finally, AAM can be extended with algorithms for the online selection of M. Here, as our study exhaustively illustrates performance tradeoffs in the available HTM implementations, it would facilitate the runtime decisions on how to select M. For example, the runtime can prune the space of all the applicable values of HTM parameters depending on which HTM is utilized. In addition, our performance model can be further extended and combined with data mining techniques to enable effective online decisions based on graph sampling. We leave this study for future research.

8. CONCLUSION

Designing efficient algorithms for massively parallel and distributed graph computations is becoming one of the key challenges for the parallel programming community [18]. Graph processing is fine-grained by nature and its traditional implementations based on atomics or fine locks are error-prone and may entail significant overheads [18].

We propose Atomic Active Messages (AAM), a mechanism that reduces the amount of fine-grained synchronization in irregular graph computations. AAM is motivated by recent advances towards implementing transactional memory in hardware. AAM provides several high performance techniques for executing fine-grained graph modifications as coarse transactions, it facilitates the utilization of state-ofthe-art hardware mechanisms and resources, and it can be used to accelerate highly optimized codes such as Graph500 by more than 100%.

AAM targets highly-parallel multi- and manycore architectures and distributed-memory machines. It provides a novel classification of atomic active messages that can be used to design and program both shared- and distributedmemory graph computations. AAM enables different optimizations from both of these worlds such as coarsening intra-node transactions and coalescing inter-node activities. We illustrate how to implement AAM with HTM; however, other mechanisms such as distributed STM [21], flatcombining [13], or optimistic locking [20] could also be used. Finally, to the best of our knowledge, our work is the first detailed performance analysis of hardware transactional memory in the context of graph computations and the first to compare HTMs implemented in Intel Haswell and IBM Blue Gene/Q. Among others, we conjecture that implementing HTM in the bigger L2 cache (BG/Q) enables higher performance than in the smaller L1 cache (Haswell). We believe our analysis and data can be used by architects and engineers to develop a more efficient HTM that would offer even higher speedups for irregular data analytics.

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