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Galois: A System for Parallel Execution of Irregular Algorithms

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Galois: A System for Parallel Execution of Irregular Algorithms

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To my family

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Galois: A System for Parallel Execution of Irregular Algorithms

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A programming model which allows users to program with high productivity and which produces high performance executions has been a goal for decades. This dissertation makes progress towards this elusive goal by describing the design and implementation of the Galois system, a parallel programming model for shared-memory, multicore machines. Central to the design is the idea that scheduling of a program can be decoupled from the core computational operator and data structures. However, efficient programs often require application-specific scheduling to achieve best performance. To bridge this gap, an extensible and abstract scheduling policy language is proposed, which allows programmers to focus on selecting high-level scheduling policies while delegating the tedious task of implementing the policy to a scheduler synthesizer and runtime system. Implementations of deterministic and prioritized scheduling also are described.

An evaluation of a well-studied benchmark suite reveals that factoring programs into operators, schedulers and data structures can produce significant performance improvements over unfactored approaches. Comparison of the Galois system with existing programming models for graph analytics shows significant performance improvements, often orders of magnitude more, due to (1) better support for the restrictive programming models of existing systems and (2) better support for more sophisticated algorithms and scheduling, which cannot be expressed in other systems.

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Chapter 1

Introduction

Computing devices play a central role in society. With the rise of data centers, which accounted for 2% of domestic energy consumption in 2012 ([Natural Resources Defence Council, 2014](#)), and the ubiquity of mobile devices, there is increasing need to improve the efficiency of computation to reduce energy consumption. The best way to improve efficiency is to exploit parallelism and to minimize data movement.

This dissertation addresses performance through the lens of programming models; it investigates what software program abstractions lead to high performance programs. One traditional and successful model for programmer productivity is thinking of a program as an algorithm over data structures or, in the words of Niklaus Wirth, Program = Algorithm + Data Structure ([Wirth, 1978](#)). This model improves productivity because it divides writing a program into two parts: algorithms that are specific to the problem at hand, and data structures that are more general and can be reused among different programs.

This dissertation argues that parallel programs require a more refined model. The algorithm itself should be divided into parts, Algorithm = Operator + Schedule. The operator is the core computation that is specific to a problem, and the schedule is how the computation is mapped to particular hardware resources in time. In the same way data structures are reused among sequential programs, schedulers should be reused among parallel programs.

Given a decomposition of a program into *activities* or tasks, the *scheduling problem* is the assignment of these activities to processors, and the specification of an order in which each processor should execute the activities assigned to it. Scheduling is important for both sequential and parallel implementations of algorithms since it may affect locality and load balance; it may even effect the total amount of work performed by some programs, as will be shown shortly.

Scheduling can be done either statically by a compiler or dynamically by a runtime or operating system. Static scheduling can be used when dependences between activities are known statically and the execution time of each activity can be estimated accurately at compile-time. Stencil computations are the classic examples of algorithms amenable to static scheduling (for example see (Stock et al., 2014)). Dynamic scheduling is useful for problems in which (1) dependences between activities cannot be elucidated statically, or (2) new work is created dynamically so the number of activities is not known statically, or (3) accurate estimates of the time required to execute each activity are not available. The vast majority of algorithms, including almost all *irregular* algorithms (algorithms where the key data structures are sparse graphs) require dynamic scheduling.

For the most part, prior work on dynamic scheduling has focused on problems in which there are no dependences between activities, and new work is not created dynamically, so the only problem is that the time required to execute an activity cannot be determined accurately. Self-scheduling of DO-ALL loops in OpenMP is the classic example. Activities in this case correspond to iterations of the DO-ALL loop; the number of iterations is known before the loop is executed, and it is assumed that there are no dependences between iterations. However, different iterations may take different and unpredictable amounts of time to execute, so to ensure good load balance, OpenMP provides scheduling policies such as chunked dynamic self-scheduling, in which a processor gets a chunk of k iterations every time it needs work, and guided self-scheduling, in which the chunk size decreases steadily as the loop nears completion (Dagum and Menon, 1998). Chunking reduces the

overheads of scheduling and may improve locality.

More recently, attention has shifted to task-parallelism in which new activities are created dynamically, although it is still assumed that all activities are independent except for fork-join control dependences. In OpenMP 3.0, there is support for different dynamic scheduling policies such as breadth-first and work-first policies (Duran et al., 2008). Another popular technique is *work-stealing*. In work-stealing, each thread has a local deque that contains activities to execute. When a thread's local deque is empty, it selects the local deque of another thread, the victim, and tries to steal activities from it. Work-stealing is parameterized by the order maintained in the local deque (usually LIFO) and how a thread selects a victim (usually at random). Work-stealing was implemented in MultiLisp (Halstead, 1985) and was later popularized by the Cilk language (Blumofe et al., 1995), where it is used to implement fork-join parallelism. It is now available in many programming environments: Intel Threading Building Blocks (TBB) (Reinders, 2007) the Java library (Lea, 2000) and the .Net library (Leijen et al., 2009).

The key assumptions in such work on task-parallelism are that any dependences between activities are captured by fork-join control dependences and are known statically. While these assumptions are reasonable for regular (i.e., dense-array) algorithms and divide-and-conquer algorithms, they do not hold for most irregular graph algorithms because dependences in these algorithms are complex functions of runtime values (e.g., the shape of the graph and the values on nodes and edges), which may themselves change during execution.

An abstract description of parallelism in irregular algorithms is the following. At each step of the algorithm, there are certain *active nodes* in the graph where computation needs to be performed. Performing the computation at an active node may require reading or writing other graph nodes and edges, known collectively as the *neighborhood* of that activity. The neighborhood is usually distinct from the neighbors of the active node. In general, there are many active nodes in a graph, so a sequential implementation must pick

one of them and perform the appropriate computation.

In *unordered algorithms*, which are the focus of this dissertation, the implementation is allowed to pick *any* active node for execution. In contrast, *ordered algorithms* have a specific order in which active nodes must be executed. For unordered algorithms, the final output may be different for different orders of executing active nodes, but all such outputs are acceptable, a feature known as *don't-care non-determinism*. A parallel implementation of such an algorithm can process active nodes simultaneously, provided their neighborhoods do not overlap. This condition can be relaxed but is sufficient for correct execution. In general, the neighborhood of an activity is not known until the activity has finished execution.

The parallelism that results from processing activities in parallel subject to neighborhood and ordering constraints is *amorphous data-parallelism* (ADP) (Pingali et al., 2011). Efficient exploitation of amorphous data-parallelism requires far more sophisticated runtime support than fork-join parallelism or DO-ALL parallelism for the following reasons.

1. In most irregular algorithms, nodes become active dynamically, so the number of activities is not known statically.
2. In general, the neighborhood of an activity may be known only after the activity completes execution. Therefore, it may be necessary to use optimistic or speculative parallelization.
3. Most importantly, the number of activities that are executed by an algorithm may be different for different schedules. In some cases, the amount of work may differ by an asymptotic factor, as shown in the following chapter. If this is the case, it is critical to capture the scheduling of the more work-efficient scheduling.

For these reasons, even *sequential* implementations of irregular algorithms often use handcrafted, algorithm-specific scheduling policies; for example, some mesh refinement algorithms process triangles or tetrahedra in decreasing size order since this can reduce the

total amount of refinement work (Miller, 2004). Section 2.3 gives examples of the policies used in the literature. However, following these orders strictly can dramatically reduce parallelism, so parallel implementations of irregular algorithms often use more complex scheduling policies that trade off extra work for increased parallelism. These schedulers are themselves concurrent data structures and add to the complexity of parallel programming. In addition, they cannot easily be reused for other applications.

This dissertation introduces a flexible and efficient approach for specifying and synthesizing schedulers for sequential and parallel implementations of irregular algorithms. It distinguishes between *scheduling policies*, which are informal descriptions of the order in which activities should be processed (e.g., LIFO, FIFO, etc.), *scheduling specifications*, which are formal descriptions of scheduling policies, and *schedulers*, which are concrete implementations of scheduling specifications. A *schedule* is a specific mapping of activities to processors in time.

Of course, to address performance, efficient scheduling must be combined with scalable data structures. The main contribution of this dissertation is the design and implementation of a parallel programming model for unordered algorithms based on an extensible scheduling policy DSL and a library of parallel data structures. This system is called the Galois system.

Chapter 2 describes the core program abstraction in Galois, the operator formulation, and it also shows how the operator formulation is a natural abstraction for many algorithms. Chapter 3 summarizes existing programming models for parallelism. Chapter 4 introduces the design principles that guide the implementation of the Galois system. Chapter 5 and Chapter 6 describe the implementation concretely in terms of data structures and scheduling, respectively. To address one concern with unordered algorithms, their non-determinism, Chapter 7 presents a deterministic scheduling algorithm that permits unordered algorithms to be run non-deterministically or deterministically as desired.

To support the utility of the Galois system, Chapter 8 evaluates the Galois system

in two ways. First, it shows that the design principles introduced in [Chapter 4](#) are sufficient conditions for scalability by showing that their manual application can substantially improve the performance of the STAMP benchmark suite, a well-studied but until now poorly performing benchmark suite. Second, [Chapter 8](#) shows that the Galois system is a significant improvement over existing parallel programming models because (1) existing programming models can be reimplemented in Galois and obtain better performance than their original implementations and (2) the Galois system can express more sophisticated algorithms beyond the capabilities of previous systems, which result in orders of magnitude performance improvements for many graph analytics problems.

Chapter 2

A Data-Centric View of Parallelism and Locality

This chapter introduces the operator formulation of programs. [Section 2.1](#) describes a model problem that illustrates the key ideas, and [Section 2.2](#) generalizes the basic issues in the model problem to develop the operator formulation. [Section 2.3](#) shows how various algorithms can be expressed with this formulation.

2.1 Model Problem: SSSP

Given a weighted graph $G = (V, E, w)$, where V is the set of nodes, E is the set of edges, and w is a map from edges to edge weights, the single-source shortest-paths (SSSP) problem is to compute the distance of the shortest path from a given source node $s \in V$ to each node in the graph. Edge weights can be negative, but it is assumed that there are no negative weight cycles.

In most SSSP algorithms, each node is given a label that holds the distance of the shortest known path from the source to that node. This label $\text{dist}(v)$ is initialized to 0 for

Portions of this chapter have previously appeared in [\(Pingali et al., 2011\)](#), where the TAO classification was originally described.

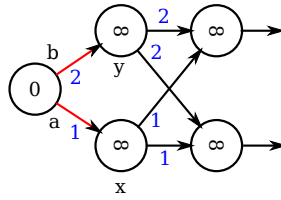


Figure 2.1: Example of SSSP (edge weights shown in blue)

s and ∞ for all other nodes. The basic SSSP operation is *edge relaxation* (Cormen et al., 2009): given an edge (u, v) such that $\mathbf{dist}(u) + w(u, v) < \mathbf{dist}(v)$, the value of $\mathbf{dist}(v)$ is updated to $\mathbf{dist}(u) + w(u, v)$. Each relaxation, therefore, lowers the \mathbf{dist} label of a node, and when no further relaxations can be performed, the resulting node labels are the shortest distances from the source to the nodes, regardless of the order in which the relaxations were performed. When relaxations are applied arbitrarily, this algorithm is called chaotic relaxation (Chazan and Miranker, 1969).

Nevertheless, some relaxation orders may converge faster and are therefore more work-efficient than others. For example, consider the graph in Figure 2.1. Edges in the graph where edge relaxation can be performed are shown in red. If edge b is relaxed, it will create new opportunities for edge relaxation at all the outgoing edges of node y . If those newly enabled edges are processed before edge a , those edges will be processed again once edge a is relaxed. However, if edge a is processed before edge b , processing edge b will not create any new opportunities for edge relaxation, and the total number of relaxation operations is reduced. Dijkstra's SSSP algorithm (Dijkstra, 1959) applies edge relaxation to all the outgoing edges of a node and relaxes each node just once by using the following strategy: from the set of nodes that have not yet been relaxed, pick one that has the minimal label.

However, Dijkstra's algorithm does not have much parallelism due to its reliance on a centralized priority queue, so some parallel implementations of SSSP use this rule only as a heuristic for priority scheduling: given a choice between two edges with different \mathbf{dist} labels on their sources, they pick the one with the smaller label, but they may also

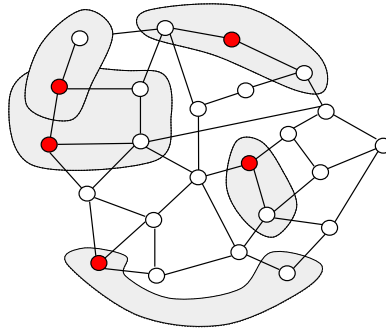


Figure 2.2: Illustration of the operator formulation

execute some edges out of priority order to exploit parallelism. One such algorithm is delta-stepping SSSP (Meyer and Sanders, 1998). The price of this additional parallelism is that some nodes may be relaxed repeatedly. A balance must be struck between controlling the amount of extra work and exposing parallelism.

2.2 Operator Formulation

To discuss common issues in parallel programs, it is convenient to use the terminology of the *operator formulation* (Pingali et al., 2011), a data-centric programming model for expressing parallelism in regular and irregular algorithms. The basic concepts of the operator formulation are illustrated in Figure 2.2.

- *Active nodes* are nodes in the graph where computation must be performed; they are shown as red dots in Figure 2.2.
- The computation at an active node is called an *activity*, and it results from the application of an *operator* to the active node. In some algorithms, it is more convenient to think in terms of active *edges* rather than active nodes. Without loss of generality, we will use the term active nodes. The operator is a composition of elementary graph operations with other arithmetic and logical operations.

Note that graphs themselves are general data structures; any other data structure can

be expressed as a graph. An array is a node with ordered neighbors. A pointer-based data structure is a graph of memory locations with edges corresponding to pointers to other memory locations.

- The set of graph elements read and written by an activity is its *neighborhood*. The neighborhood of the activity at each active node in [Figure 2.2](#) is shown as a “cloud” surrounding that node. If there are several data structures in an algorithm, neighborhoods may span multiple data structures. In general, neighborhoods are distinct from the set of immediate neighbors of the active node, and neighborhoods of different activities may overlap. In a parallel implementation, the semantics of reads and writes to such overlapping regions must be specified carefully. The general term for what happens when two activities cannot proceed in parallel due to their neighborhoods is a *conflict*.

The SSSP algorithms described in the previous section can be expressed in the operator formulation. In chaotic relaxation and Dijkstra’s algorithm, the operator is the edge relaxation operator. The active edges are edges where edge relaxation can be applied, and the neighborhood is the active edge and the corresponding endpoints of the edge.

In general, there may be multiple active nodes, so an algorithm must specify which order of executing active nodes is valid. There are two classes of ordering. In *unordered algorithms*, any order of processing active nodes is valid. The chaotic relaxation algorithm for SSSP is an example of an unordered algorithm. In *ordered algorithms*, the algorithm has a specific order in which active nodes are processed. Dijkstra’s algorithm is an example. In that algorithm, active edges must be processed in priority order.

The operator formulation leads to a natural definition of parallelism.

Definition 2.1. *Given a set of active nodes and an ordering on it, amorphous data-parallelism (ADP) is the parallelism that arises from simultaneously processing active nodes subject to neighborhood and ordering constraints.*

Amorphous data-parallelism generalizes many common notions of parallelism. ADP with no neighborhood or ordering constraints is data parallelism (Hillis and Steele, 1986). ADP with no neighborhood constraints but where activities are ordered according to fork-join dependencies is nested data-parallelism (Blelloch, 1992). One can go even further. Instruction-level parallelism (Hennessy and Patterson, 2003) can be seen as an instance of ADP where (1) activities are processor instructions, (2) activities are ordered according to program instruction order, and (3) conflicts only occur when neighborhoods overlap and at least one activity writes to the overlapping region.

ADP also captures many models of consistency. If (1) conflicts only occur when neighborhoods overlap and at least one activity writes to the overlapping region and (2) active nodes can be processed in any order, then ADP generates serializable executions (Papadimitriou, 1986). If conflicts only occur when two activities have overlapping neighborhoods and both activities write to the overlapping region, then executions satisfy snapshot isolation (Berenson et al., 1995).

The operator formulation and ADP permit an abstract description of algorithms that highlights similarities between algorithms and parallelization techniques across application domains. At first glance, the chaotic relaxation algorithm for SSSP and the relabel-to-front algorithm for maximum flow, described in Section 2.3, seem very different, but it turns out they share many of the same properties in the operator formulation, and optimizations like ordered execution with priorities (e.g., Dijkstra’s algorithm) that are used for SSSP also can be used with the maximum flow problem (e.g., HL ordering (Cherkassy and Goldberg, 1995)).

The next section gives a baseline execution model for programs in the operator formulation, and Section 2.2.2 and Section 2.2.3 describe two techniques to analyze and optimize programs based solely on their structure in the operator formulation.

2.2.1 Baseline Execution Model

The baseline execution model is speculative execution. Shared data structures like graphs are stored in shared-memory, and active nodes are processed by some number of threads. A thread picks an active node from a workset¹ and speculatively applies the operator to that node, making calls to a graph library to perform operations as needed. The neighborhood of an activity grows incrementally as graph methods touch areas of the graph. To detect conflicts, the graph maintains logical locks associated with each node or edge of the graph. These locks are acquired by a thread before it can access that element. Locks are held until the activity terminates. If a thread acquires a logical lock for writing that has been already acquired by another thread (for reading or writing), a conflict is reported to the runtime system, which rolls back one of the conflicting activities. Lock manipulation is performed entirely by the methods in the graph class. In addition, to support rollback, each graph method that modifies the graph makes a copy of the data before modification.

If active elements are unordered, the activity commits when the application of the operator is complete, and all acquired locks are then released. If active elements are ordered, active nodes can still be processed in any order, but they must appear to commit in serial order. This can be implemented using a data structure similar to a reorder buffer in out-of-order processors. In this case, activities that have been executed out-of-order keep their locks and are held in a reorder buffer until they reach the head of the buffer or are aborted. Alternatively, a dependence graph can be used to schedule ordered tasks; although, whether dependence graph scheduling is possible depends on what the order is and how tasks behave (Hassaan et al., 2015).

For unordered active elements, transactional memory (Harris and Fraser, 2003; Herlihy and Moss, 1993) can be used to accelerate conflict detection and rollback in hardware, see Section 8.1.

What constitutes a neighborhood conflict can be refined or coarsened. A more re-

¹Throughout this dissertation, the colloquial term workset is used, although more formally, these objects behave as bags or multisets because they may contain duplicate items.

```

1 Workset ws(G.nodes())
2 foreach Node p in ws:
3   // Phase 1: reading neighborhood
4   int s = 0
5   for Node n in G.neighbors(p):
6     s += G.getData(n)
7   // Failsafe point
8   // Phase 2: writing to neighborhood
9   // elements written to were read in Phase 1
10  for Node n in G.neighbors(p):
11    G.getData(n) += s

```

Figure 2.3: Example of a cautious operator

finer conflict detection scheme would allow activities to proceed in parallel as long as the corresponding method calls commute with respect to the logical operations they are implementing (Kulkarni et al., 2011). Accesses in disjoint areas of neighborhoods are presumed to commute. A more coarse conflict detection scheme would allow activities to proceed only if their neighborhoods are disjoint. Two activities reading or writing the same graph element would result in a conflict. This scheme can be implemented with exclusive logical locks that use a compare-and-set instruction to mark a graph element with the id of the activity that touches it (see Section 5.2).

This speculative executor is sufficient to execute any program in the operator formulation, but it may be inefficient in practice. For instance, the polyhedral model (Feautrier and Lengauer, 2011) is a methodology that can optimize and schedule array programs with affine subscripts at compile time without speculation. One way to address the performance concerns of the baseline execution model is to identify specific program properties that make programs amenable to certain analysis or execution strategies and use a specialized executor instead of the baseline executor for these cases.

As an example, sometimes tasks are *cautious* (Méndez-Lojo et al., 2010), which means they read their entire neighborhood before writing to any element of it (see Figure 2.3 for an example). For unordered cautious tasks, conflict detection and correction can be done using lightweight mechanisms because the synchronization problem reduces to the

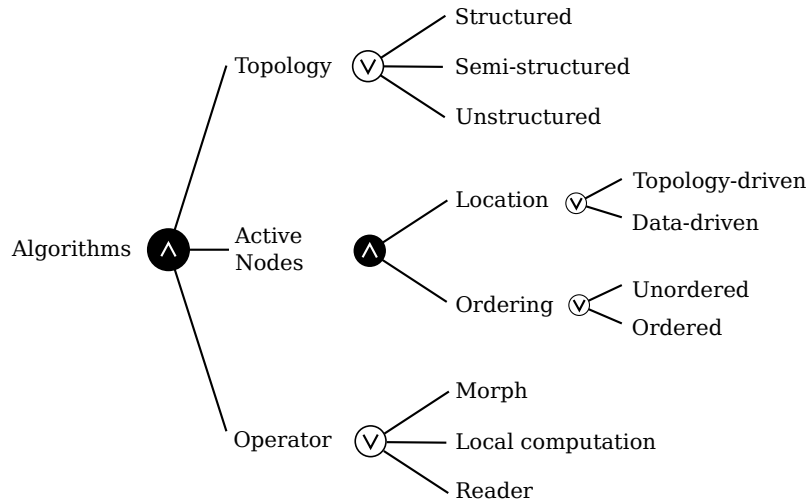


Figure 2.4: TAO analysis of algorithms

well-known dining philosopher’s problem (Chandy and Misra, 1984). Conceptually, each abstract location can be *acquired* by an owner. The execution of a task can be divided into two phases: in the first phase, a task reads locations but does not write to any of them, acquiring ownership of these locations, and in the second phase, the task writes to some locations, but it does not write to any location that it did not read in the first phase, because it is cautious. The point between the first and second phase is called the *failsafe point*. For cautious tasks, conflicts are detected in the first phase, and rollback is implemented simply by releasing ownership of all locations. Once the failsafe point has been crossed, global data structures can be updated in place without the need for backup copies of modified data.

In this spirit, the following two sections describe methods of classifying programs with an eye towards identifying properties that are useful for optimized execution. The first is TAO analysis (Pingali et al., 2011) (see Section 2.2.2), which classifies programs along three dimensions: topology, active nodes and ordering. The second method focuses on properties of iterative fixpoint algorithms (see Section 2.2.3).

2.2.2 TAO Analysis

TAO analysis (Pingali et al., 2011) is a method for structural analysis of algorithms with respect to their possible parallelizations. It is based on classifying algorithms and data structures along three dimensions.

1. *Topology*: graph topologies are classified according to the Kolmogorov complexity of their descriptions. Highly structured topologies can be described concisely with a small number of parameters, while unstructured topologies require verbose descriptions. The topology of a graph is an important indicator of the kinds of optimizations available to algorithm implementations; for example, algorithms in which graphs have highly structured topologies may be more amenable to static analysis and optimization.

- *Structured*: an example of a structured topology is a graph consisting of labeled nodes and *no* edges. This is isomorphic to a set or multiset; its topology can be described by a single number, the number of elements in the set or multiset. If the nodes are totally ordered, the graph is isomorphic to a sequence of stream. Cliques, i.e., graphs in which every pair of nodes is connected by a labeled edge, are isomorphic to square dense matrices with row/column numbers coming from the total ordering of the nodes. Their topology is completely specified by a single number, the number of nodes in the clique.
- *Semi-structured*: trees are classified as semi-structured topologies. Although trees have useful structural invariants, there are many trees with the same number of nodes and edges.
- *Unstructured*: general graphs fall in this category. Even among general graphs, some may be considered more structured than others. For instance, graphs whose nodes can be divided into partitions with a small edgecut versus graphs whose partitions have a large edgecut value.

2. *Active nodes*: This dimension describes how nodes become active and the order in which they must be processed.

- *Location*: nodes can become active in a *topology-driven* or *data-driven* manner. In topology-driven algorithms, the execution of the operator at some active node does not cause other nodes to become active. Common examples are algorithms that iterate over all the nodes or edges of a graph. In data-driven algorithms, an activity at one node may cause other nodes to become active, so nodes become active in a data-dependent and unpredictable manner. An example is the chaotic relaxation algorithm for SSSP.
- *Ordering*: As discussed in above, active nodes in some algorithms are ordered whereas in others they are unordered.

3. *Operator*: This final dimension describes how operators modify the graph.

- *Morph*: a morph operator may modify its neighborhood by adding or deleting nodes and edges, and it may also update values on nodes and edges. The Delaunay mesh refinement operator described in [Section 2.3](#) is an example.
- *Local computation*: a local computation operator may update values stored on nodes and edges in its neighborhood, but it does not change the graph connectivity. Finite-difference computations are a classic example. The chaotic relaxation algorithm is another.
- *Reader*: an operator is a reader for a data structure if it does not modify it in any way. For example, the ray tracing operator is a reader for the scene being rendered.

These definitions can be generalized in the obvious way for algorithms that deal with multiple data structures. In that case, neighborhoods span multiple data structures, and the classification of an operator is with respect to a particular data structure. For example, in matrix

multiplication, $C = AB$, the operator is a local computation for C and a reader for matrices A and B .

TAO analysis can be used to organize programs into classes that share the same parallelization concerns. For topology-driven active nodes, if the topology and operator are known at compile-time, which is the case for many dense matrix codes, parallelization can also occur at compile-time in principle. In practice, to adapt to the variance in the execution time of tasks, high performance parallelizations of dense matrix codes may also include a runtime component for load balancing, for instance see the DAGuE system (Bosilca et al., 2012). A similar trend occurs in data-parallel codes, which also can be parallelized at compile-time, and often use a work-stealing scheduler (Blumofe et al., 1995) to balance work among threads at runtime, for instance see (Baskaran et al., 2009).

For sparse matrix codes, the topology (i.e., the structure of the sparse matrix) is not known until the input is read by the program. Compiler-based parallelization cannot be used, and the earliest time that parallelization can be attempted is *just-in-time*, after the input is read but before the bulk of computation begins. This is called the inspector-executor (Das et al., 1995) approach. If the matrix is discovered to be relatively dense or if it has dense subregions (e.g., nearly block-diagonal), the executor in the inspector-executor approach can apply dense matrix subroutines for parts of the sparse matrix; however, in terms of the classification of techniques, the earliest point at which the decision to apply these dense matrix subroutines is when the input is read even though the dense subroutines themselves may be parallelized at compile-time.

At the most extreme, parallelization may be done at *runtime*, interleaved with the parallelized computation itself. This is the case with data-driven active nodes and with most morph computations. Data-driven active nodes require runtime parallelization because active nodes are not known without executing the activity.

A similar conclusion holds for morph operators, but one special case is when the modification performed by the morph can be efficiently simulated without running the op-

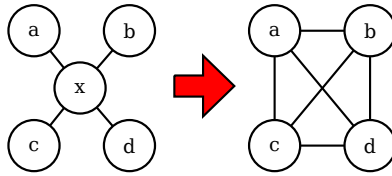


Figure 2.5: Example of node elimination of node x

erator itself. Examples are sparse matrix factorization codes like Cholesky and LU decomposition. In these codes, the operator performs a morph called *node elimination*, in which a node is removed from a graph and edges are inserted as needed between its erstwhile neighbors to make a clique (see [Figure 2.5](#) for an example). This operator can simulated by a just-in-time analysis with simple arithmetic operations although the operator itself requires floating-point operations to compute the factored matrix values. In this application area, the simulation of the factorization operator is called *symbolic factorization* and is an important step in high-performance, parallel implementations ([Gilbert and Schreiber, 1992](#)). Symbolic factorization builds a dependence graph called an elimination tree. The actual factorization, called *numerical factorization*, uses the elimination tree to schedule tasks. LU has an analogous operator, but *pivoting* is often done to improve numerical stability. In terms of TAO analysis, the active nodes are data-driven, which means runtime techniques must be used for parallelization. To facilitate just-in-time parallelization, parallel LU implementations use *partial pivoting* instead, in which the operator is coarsened to multiple nodes and pivoting only occurs within a coarsened operator.

TAO analysis is useful for understanding which parallelization strategies are feasible given a program. The following section describes another type of analysis that explores possible ways of implementing a particular class of programs, iterative fixpoint algorithms.

2.2.3 Analysis of Iterative Fixpoint Algorithms

Iterative fixpoint algorithms are programs that consist of operators that repeatedly read and write memory locations until some convergence property is met. A special case of iterative

fixpoint algorithms are asynchronous fixpoint algorithms (Bertsekas and Tsitsiklis, 1989). They are asynchronous because values read may not correspond to the value most recently written. Asynchronous algorithms² are amenable to parallel and distributed computation because they can tolerate long commutation delays. One example of an asynchronous fixpoint algorithm already introduced in this chapter is the chaotic relaxation algorithm for SSSP.

Iterative algorithms as a class tend to benefit from the same kinds of transformations. Since some of these transformation change the operator or the asymptotic behavior of the algorithm, they are not typical optimizations in the compiler community sense of the term, but nevertheless, they are techniques that application programmers use to improve the performance of programs.

The foremost transformation is tolerating asynchrony. Whether a program can tolerate stale updates is a deep algorithmic property, but once known, parallelizing systems can exploit this property to restructure communication to follow more efficient patterns at the machine level. For instance, as originally developed, using stochastic gradient descent (SGD) for solving linear support vector machines (SVMs) requires reading the most recent values for the weight vector, but the recently introduced Hogwild approach (Recht et al., 2011) eschews a serializable locking policy for racy reads and writes. This is possible because the algorithm tends to still converge even in the presence of noisy or stale data (see Section 5.4).

The remaining transformations can be summarized as follows: *what* does the operator do, *where* in the graph is it applied, and *when* is the corresponding activity executed?

What does the operator do? In general, the operator expresses some computation on the neighborhood elements. In some graph problems such as SSSP, operators can be implemented in two general ways called here *push* style or *pull* style. A push-style operator reads

²In contrast to asynchronous algorithms which will converge for any communication delay, partially asynchronous algorithms only converge when communication delays are bounded. For the purpose of the discussion in this section, asynchronous and partially asynchronous algorithms are treated interchangeably.

the label of the active node and writes to the labels of its neighbors; information flows from the active node to its neighbors. A push-style SSSP operator attempts to update the `dist` label of the immediate neighbors of the active node by performing relaxations with them. In contrast, a pull-style operator writes to the label of the active node and reads the labels of its neighbors; information flows to the active node from its neighbors. A pull-style SSSP operator attempts to update the `dist` label of the active node by performing relaxations with each neighbor of the active node. In a parallel implementation, pull-style operators require less synchronization since there is only one writer per active node.

Usually, the operator represents the smallest logical unit of parallel computation, but in some cases, the convergence of a fixpoint algorithm can be sped up by coarsening the graph to speed up the flow of information across the graph. In its simplest form, coarsening may simply be scheduling multiple activities as one unit or treating subgraphs as a single node or edge, but the transformations used in practice may also incorporate algorithmic performance improvements that change the behavior of the operator or the representation of subgraphs for the coarsened algorithm. The multigrid method for solving linear systems is a classic example as well as elimination-based dataflow analysis of programs (Allen and Cocke, 1976). The basic idea is to transform the graph into a smaller subproblem, solve the subproblem and interpolate the results back onto the original graph. The transformation, subproblem solving and interpolation steps are application-specific. A related idea is preconditioning, which is used in iterative linear solvers to preprocess an input to improve convergence or performance; in this case, the core operation remains the same whether or not preconditioning is used.

Where is the operator applied? As in TAO analysis, active nodes can be *topology-driven* or *data-driven*.

In a *topology-driven* computation, active nodes are defined structurally in the graph, and they are independent of the values on the nodes and edges of the graph. The Bellman-Ford SSSP algorithm is an example (Bellman, 1958; Ford and Fulkerson, 1962); this al-

gorithm performs $|V|$ supersteps, each of which applies a push-style or pull-style operator to all the edges. Practical implementations terminate the execution if a superstep does not change the label of any node. Topology-driven computations can be parallelized by partitioning the nodes of the graph between processing elements.

In a *data-driven* computation, nodes become active in an unpredictable, dynamic manner based on data values, so active nodes are maintained in a workset. In a data-driven SSSP program, only the source node is active initially. When the label of a node is updated, the node is added to the workset if the operator is push-style; for a pull-style operator, the neighbors of that node are added to the workset. Data-driven implementations can be more work-efficient than topology-driven ones since work is performed only where it is needed in the graph. However, load-balancing is more challenging, and careful attention must be paid to the design of the workset to ensure it does not become a bottleneck.

When is an activity executed? When there are more active nodes than threads, the implementation must decide which active nodes are prioritized for execution and when the side-effects of the resulting activities become visible to other activities. There are two popular models that are called here *autonomous scheduling* and *coordinated scheduling*.

In autonomous scheduling, activities are executed with transactional semantics, so their execution appears to be atomic and isolated. Parallel activities are serializable, so the output of the overall program is the same as some sequential interleaving of activities. Threads retrieve active nodes from the worklist and execute the corresponding activities, synchronizing with other threads only as needed to ensure transactional semantics. This fine-grain synchronization can be implemented using speculative execution with logical locks or lock-free operations on graph elements. The side-effects of an activity become visible externally when the activity commits.

Coordinated scheduling, on the other hand, restricts the scheduling of activities to rounds of execution, as in the Bulk-Synchronous Parallel (BSP) model (Valiant, 1990). The execution of the entire program is divided into a sequence of supersteps separated by barrier

synchronization. In each superstep, a subset of the active nodes is selected and executed. Writes to shared-memory, in shared-memory implementations, or messages, in distributed-memory implementations, are considered to be communication from one superstep to the following superstep. Therefore, each superstep consists of updating memory based on communication from the previous superstep, performing computations, and then issuing communication to the next superstep. Multiple updates to the same location are resolved in different ways as is done in the varieties of PRAM models, such as by using a reduction operation (JaJa, 1992).

Application-specific priorities Of the different algorithm classes discussed above, data-driven, autonomously scheduled algorithms are the most difficult to implement efficiently. However, they converge much faster than algorithms that use coordinated scheduling for some problems (Bertsekas and Tsitsiklis, 1989). Moreover, for high-diameter graphs like road networks, data-driven autonomously scheduled algorithms may be able to exploit more parallelism than algorithms in other classes; for example, in BFS, if the graph is long and skinny, the number of nodes at each level will be quite small, limiting parallelism if coordinated scheduling is used.

Autonomously scheduled, data-driven graph analytics algorithms benefit from application-specific priorities and priority scheduling to balance work-efficiency and parallelism. One example is delta-stepping SSSP (Meyer and Sanders, 1998), the most commonly used parallel implementation of SSSP. The workset of active nodes is implemented, conceptually, as a sequence of bags, and an active node with label d is mapped to the bag at position $\lfloor \frac{d}{\Delta} \rfloor$, where Δ is a user-defined parameter. Idle threads pick work from the lowest-numbered non-empty bag, but active nodes within the same bag may execute in any order relative to each other. The optimal value of Δ depends on the graph.

The general picture is the following. Each task t is associated with an integer *priority*(t), which is a heuristic measure of the importance of that task for early execution relative to other tasks. For delta-stepping SSSP, the priority of an SSSP relaxation task is the value

$\lfloor \frac{d}{\Delta} \rfloor$. A task t_1 has *earlier priority* than a task t_2 if $\mathbf{priority}(t_1) < \mathbf{priority}(t_2)$. It is permissible to execute tasks out of priority order, but this may possibly lower work efficiency.³ A good parallel runtime system must permit application programmers to specify such application and input-specific priorities for tasks, and the system must schedule these fine-grain tasks with minimal overhead and minimize priority inversions.

2.3 Parallel Algorithms

This section introduces several algorithms using the concepts of the operator formulation, TAO analysis and analysis of fixpoint algorithms. Pseudocode for these algorithms is written using the Galois programming model (see [Chapter 4](#)), which is a sequential, object-oriented programming model augmented with a *Galois unordered-set iterator*, which is similar to set iterators in C++ or Java but permits new items to be added to a set while it is being iterated over.

- **foreach** $T\ e\ \mathbf{in}\ S: B(e)$ — The loop body $B(e)$ is executed for each item e of type T in set S . The order in which iterations execute is indeterminate and can be chosen by the implementation. There may be dependences between the iterations. An iteration may add items to S during execution.

2.3.1 Delaunay Triangulation

Finding the Delaunay triangulation (DT) of a set of points is a classic computational geometry problem. There are many algorithms for finding the triangulation; this section describes the incremental algorithm of Bowyer and Watson ([Bowyer, 1981](#); [Watson, 1981](#)). Initially, there is one large triangle that covers all the points. Then, point p calculates the triangle that contains it and calculates all triangles whose circumcircles include p . This is the cavity

³If the priority order must be strictly followed, the algorithm is ordered and different schedulers must be applied ([Hassaan et al., 2015](#)). This dissertation focuses on implementing unordered algorithms.

```

1 Workset ws(points)
2 foreach Point p in ws:
3   Triangle t = findTriangleContaining(p)
4   Cavity c(t)
5   c.expand()
6   c.retriangulate()
7   G.update(c)

```

Figure 2.6: Pseudocode for Delaunay triangulation

of p . Finding the triangle that contains a point can be accomplished with a spatial acceleration structure like a kd-tree or oct-tree over triangles. The cavity is re-triangulated, p is removed, and the next point is processed. This process continues until there are no more points to process. [Figure 2.6](#) shows the pseudocode. G is the graph representing the triangulation. [Figure 2.7](#) shows an example of processing one point. To reduce the amount of time spent updating the acceleration structure, it can be built over a subset of triangles and a geometric search within the mesh can be used to find the enclosing triangle.

All orders of processing points lead to the same Delaunay triangulation. Clarkson and Shor have shown that selecting points at random is optimal ([Clarkson and Shor, 1989](#)). Amenta et al. present an algorithm called biased randomized insertion order (BRIO) that takes advantage of spatial locality while still maintaining the optimality of randomness ([Amenta et al., 2003](#)). Briefly, let n be the number of points to triangulate. Points are processed in $\log n$ rounds. The probability that a point is processed in the final round is $\frac{1}{2}$. For the remaining points, the probability that they will be processed in the next-to-last round is $\frac{1}{2}$, and so on until the first round. For the first round, all remaining points are processed with probability one. Within a round, points are processed according to the spatial divisions of an oct-tree.

2.3.2 Delaunay Mesh Refinement

Delaunay mesh refinement (DMR) ([Chew, 1993](#)) is an algorithm related to Delaunay triangulation. Given a Delaunay triangulation, triangles may have to satisfy additional quality

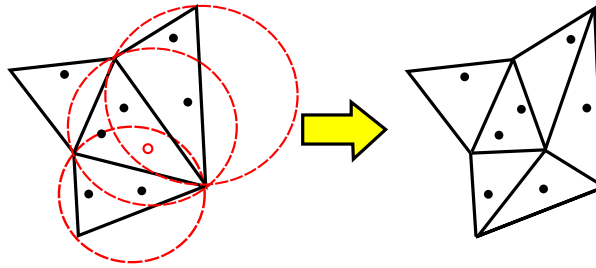


Figure 2.7: Example of processing an active point (hollow and red) for Delaunay triangulation. Circles are circumcircles of triangles containing the active point.

constraints beyond that guaranteed by triangulation. To improve the quality of a triangulation, Delaunay mesh refinement iteratively fixes “bad” triangles, which do not satisfy the quality constraints, by adding new points to the mesh and re-triangulating. Refining a bad triangle may itself introduce new bad triangles, but it can be shown that, at least in 2D, this iterative refinement process will terminate and produce a guaranteed-quality mesh. In 3D, naive refinement may terminate with tetrahedral elements with large aspect ratios (Li and Teng, 2001), which adversely affects the convergence and stability of numerical algorithms like the finite element method (Strang and Fix, 1973).

Figure 2.8 shows the pseudocode for this algorithm. It is similar to Delaunay triangulation, except that activities are centered on triangles rather than points. In both cases, a cavity is expanded and re-triangulated. However, in DMR, new bad triangles can be created that must be processed as well. They are tracked in a workset. Additionally, different orders of processing bad triangles lead to different meshes, but all such meshes satisfy the quality constraints and are acceptable outcomes of the refinement process (Chew, 1993). In contrast, for Delaunay triangulation, different orders still produce the same triangulation.

Naive implementations of DMR have quadratic worst-case running times (Ruppert, 1993) although they perform well in practice. Miller proved sub-quadratic worst-case time of a modification of DMR that processes triangles in decreasing circumcircle diameter together with other changes (Miller, 2004). In Shewchuk’s Triangle program, bad triangles are placed into buckets according to their minimum angle, each bucket stores triangles in


```

1 Workset ws(G.badTriangles())
2 foreach Triangle t in ws:
3     if !G.contains(t):
4         continue
5     Cavity c(t)
6     c.expand()
7     c.retriangulate()
8     G.update(c)
9     ws.addAll(c.badTriangles())

```

Figure 2.8: Pseudocode for Delaunay mesh refinement

FIFO order, and buckets are processed in increasing angle order (Shewchuk, 1996). Kulkarni et al. showed that a parallel implementation of DMR that distributes the initial bad triangles among threads and uses thread-local stacks for newly created bad triangles performs well in practice (Kulkarni et al., 2008).

2.3.3 Inclusion-Based Points-to Analysis

Inclusion-based points-to analysis (PTA), also known as Andersen’s algorithm (Andersen, 1994), is a flow and context-insensitive static analysis that determines the points-to relation for program variables. PTA is a fixpoint algorithm that computes the least solution to a system of set constraints. The basic algorithm maintains a workset of program variables whose points-to relations need to be computed. For each variable in the workset, the algorithm examines the system of constraints to see if the current variable satisfies the constraints. If so, the algorithm continues processing the remaining variables. If not, some set of program variables are modified to satisfy the constraints. These modified variables are then added to the workset, and the algorithm continues until the workset is empty. Hardekopf and Lin showed how the basic fixpoint algorithm augmented with sophisticated cycle detection can scale to large problem sizes (Hardekopf and Lin, 2007). From this algorithm, Méndez-Lojo et al. produced the first parallel implementation of this algorithm (Méndez-Lojo et al., 2010). The results in Section 6.2.5 are based on this implementation.

Since this is a fixpoint algorithm, all orders of processing variables will produce

the same solution. Many heuristics have been proposed for organizing the workset, such as processing variables in least recently fired (LRF) order (Pearce et al., 2003) or dividing the workset into current and next parts (Nielson et al., 1999). Variables are processed from the current part, but newly active variables are enqueued onto the next part. When the current part is empty, the roles of the current and next parts are swapped. Hardekopf and Lin report that the divided workset approach performs better in practice (Hardekopf and Lin, 2007).

2.3.4 Breadth-First Search

Given an unweighted graph $G = (V, E)$ and a starting node $s \in V$, breadth-first search (BFS) numbering is the problem of labeling each node with the length of the shortest path from s to that node. BFS is a special case of the SSSP problem, where all edge weights are one. Depending on the structure of the graph, there are two important optimizations. For low-diameter graphs, it is beneficial to switch between push and pull-based operators, which reduces the total number of memory accesses (Beamer et al., 2012). For high-diameter graphs, it is beneficial to use autonomous scheduling. Coordinated execution with high-diameter graphs produces many rounds with very few activities per round, while autonomous execution can exploit parallelism among rounds.

BFS algorithms apply the relaxation operator until convergence. For a push-based operator, the active node is a labeled node, and the operator assigns labels to unlabeled neighbors of the active node. For a pull-based operator, the active node is an unlabeled node, and the operator assigns it a label if it can find a labeled neighbor. At the beginning of the computation, it is more efficient to use a push-based operator since there are few labeled nodes and each edge relaxation propagates information through the graph; conversely, it is advantageous to switch to a pull-based implementation towards the end of the computation when most nodes are labeled, particularly for low-diameter graphs. It is possible to blend coordinated and autonomous scheduling as well to create a hybrid algorithm. Initially, the algorithm uses coordinated scheduling of the push and pull-based operators.

After a certain number of rounds of push-based traversals, the algorithm switches to prioritized autonomous scheduling with a priority function that favors executing nodes with smaller BFS numbers.

2.3.5 Approximate Diameter

The diameter of a graph is the maximum length of the shortest paths between all pairs of nodes. One exact algorithm is to compute all-pairs shortest-paths and return the maximum distance found. The cost of computing this exactly is prohibitive for any large graph, so many applications call for an approximation of the diameter (DIA) of a graph.

One algorithm is based on finding pseudo-peripheral nodes in the graph. The eccentricity $\text{ecc}(v)$ of a node v is the maximum shortest distance between v and any other node. A node is pseudo-peripheral if for every node u with distance $\text{ecc}(v)$ from v , $\text{ecc}(u) = \text{ecc}(v)$. The algorithm begins by computing a BFS from an arbitrary node. Then, it computes another BFS from the node with maximum distance, discovered by the first BFS. In the case of ties for maximum distance, the algorithm picks a node with the least degree. It continues this process until the maximum distance does not increase.

Another algorithm is to use the coordinated execution of BFS from k starting nodes at the same time. The k parameter is often picked such that the search data for a node fits in a single machine word so that it can be updated using machine atomic instructions. A bit-vector records whether the node has been visited by a BFS from starting node $i < k$. Edge relaxation performs logical-or on bit-vectors. The diameter is estimated by the maximum distance reached by the k breadth-first searches, which is a lower-bound on the diameter.

Another possibility is to use probabilistic counting (Flajolet and Martin, 1985), which estimates the number of unique vertex pairs with paths with a distance at most k . When the estimate converges, k is an estimation of the diameter of the graph.

2.3.6 Betweenness Centrality

Given a graph $G = (V, E)$ and a pair of nodes s, t , the betweenness score of a node v is the fraction of shortest paths between s and t that pass through v . The betweenness centrality (BC) of v is the sum of all its betweenness scores for all possible pairs s, t in G . A popular algorithm by Brandes (Brandes, 2001) computes the betweenness centrality of all nodes by using forward and backward breadth-first graph traversals. There are two major dimensions of parallelization. One dimension is to compute the scores for multiple source nodes at a time (outer loop parallelism). This is completely data-parallel. The other dimension is to parallelize the computation of the scores with respect to a single source node (inner loop parallelism). Inner loop parallelization can be accomplished by using the same techniques as breadth-first search (Proutzos and Pingali, 2013).

2.3.7 Connected Components

In an undirected graph, a connected component (CC) is a maximal set of nodes that are reachable from each other. One algorithm to compute the connected components of a graph is to iteratively apply BFS, choosing as a starting node any unvisited node in the graph until there are no more unvisited nodes. This algorithm is $O(|V| + |E|)$ but has a sequential dependency on the results of previous breadth-first searches. A more parallel algorithm is based on a concurrent union-find data structure. It is a topology-driven computation where each edge of the graph is visited once to add it to the union-find data structure. Another algorithm is based on iterative label propagation. Each node of the graph is initially given a unique id. Then, each node updates its label to be the minimum value id among itself and its neighbors. This process continues until no node updates its label, and it will converge slowly if the diameter of the graph is high. The complexity of this algorithm is $O(d(|V| + |E|))$ where d is the diameter of the graph.

2.3.8 Preflow-Push

Given a directed graph $G = (V, E)$, a capacity function $c : E \rightarrow \mathbb{R}^+$ mapping edges to non-negative values, and source and sink nodes $s, t \in V$, the preflow-push algorithm computes the maximal flow from source to sink. Unlike in maxflow algorithms based on augmenting paths, nodes in preflow-push can temporarily have more flow coming into them than going out. Each node n maintains its excess inflow $\text{excess}(n)$. Each node n also has a label called height, which is an estimate of the distance from n to t in the residual graph induced by unsaturated edges. Nodes with non-zero excess that are not the source nor sink are contained in a workset. These nodes are called active nodes (Goldberg and Tarjan, 1988). The preflow-push algorithm repeatedly selects a node from the workset. Each node tries to eliminate its excess by pushing flow to a neighbor (see Figure 2.9). Pushing flow may cause a neighbor to become active. A node can only push flow to a neighbor at a lower height. If a node is active but no neighbors are eligible to receive flow, the node relabels itself, increasing its height to one more than its lowest height neighbor.

Cherkassy and Goldberg show the importance of two heuristics named global relabeling and gap relabeling (Cherkassy and Goldberg, 1995). Global relabeling is a technique that periodically reassigns heights by performing a breadth-first traversal from the sink. The frequency of global relabeling is determined empirically. Gap relabeling is a technique that preemptively removes from the workset any nodes that cannot push flow to the sink. The key insight is that if no node has height h , all nodes with height greater than h cannot push flow to the sink. Cherkassy and Goldberg also consider two orders for processing active nodes: HL order, where nodes are processed in decreasing height order, and FIFO order.

```

1 Workset ws({s})
2 foreach Node u in ws:
3   L1:
4   while u.excess > 0:
5     for Node v : u.neighbors():
6       float cap = G.edgeData(u, v)
7       if cap > 0 && u.height == v.height + 1:
8         pushFlow(u, v, min(cap, u.excess))
9         if v != s && v != t:
10          ws.add(v)
11        if u.excess == 0:
12          break L1
13      relabel(u)
14  if *:
15    globalRelabel()

```

Figure 2.9: Pseudocode for preflow-push

2.3.9 PageRank

PageRank is an algorithm for computing the importance of nodes in an unweighted graph.

At its core is the following update rule

$$\mathbf{w}^{(i+1)}(v) = \alpha + (1 - \alpha) \sum_{u \in I(v)} \frac{\mathbf{w}^{(i)}(u)}{|N(u)|}$$

where $\mathbf{w}^{(i)}(v)$ is the current PageRank value for v at iteration i , $I(v)$ and $N(v)$ are the incoming and outgoing neighbors of v respectively, and $0 \leq \alpha < 1$ is some fixed damping parameter. The update rule is applied until the PageRank values converge.

Algorithms differ in how this update rule is scheduled. Topology-driven algorithms update all nodes. In the early implementations of the Google search engine, PageRank was computed using power iteration (Brin and Page, 1998), which is a topology-driven approach. Data-driven algorithms update only nodes whose neighbors' PageRank value has changed significantly. Another data-driven approach is to take samples from the graph (Leskovec and Faloutsos, 2006). Algorithms also vary in the consistency model used; in some cases, very weak consistency models have been used, like unsynchronized updates to shared

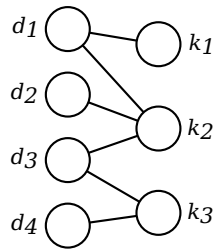


Figure 2.10: Bipartite graph of documents and features

PageRank values (Low et al., 2010). Convergence can vary significantly with scheduling and consistency model. A possible priority function is to prefer earlier execution of nodes with the greatest change in value.

The PageRank algorithm can be reduced to a sparse matrix-vector multiply. Instead of working on the input graph, the algorithm works on the transpose of the graph with edge weights added corresponding to the number of outgoing neighbors in the original input graph. That is, given an input graph G and its adjacency matrix representation A , the algorithm processes a matrix T , such that $T_{ji} = A_{ij} \cdot |N(i)|$. The topology-driven and coordinated sweep of the PageRank update above to all the nodes in the graph is equivalent to the following matrix product, $w^{(i+1)} = T \mathbf{w}^{(i)}$.

2.3.10 Support Vector Machines

Given a set of documents, a set of keywords (i.e., the *features*), and a partial function on documents indicating whether a document belongs to a class (i.e., binary classification), the problem is to learn a classifier for all documents. These inputs can be modeled as a bipartite graph with documents on one side and keywords on the other side; if a keyword k appears in a document d , there is an undirected edge (d, k) whose weight is the frequency of that keyword in that document, as shown in Figure 2.10 (weights are not shown in this figure).

One way to solve this problem is to train a support vector machine (SVM). Many algorithms for SVMs associate a value with each keyword node in the bipartite graph (these are called the *model parameters*), and iteratively update these parameters until some conver-

gence criterion is met. Stochastic gradient descent (SGD) is particular way of implementing this idea. An *update loop* iterates over each document, updating the model parameters at the immediate neighbors (keywords) of that document; for example, in [Figure 2.10](#), the values at k_1 and k_2 are updated when d_1 is processed (the value at d_1 is read-only). There is no particular order in which documents need to be visited in each iteration, so this is an example of an *unordered* algorithm. An outer loop is used to repeat the update loop until convergence.

One way to parallelize this algorithm is to recognize that if two documents do not have keywords in common (such as documents d_1 and d_4 in [Figure 2.10](#)), they can be processed in parallel. The inspector-executor approach ([Das et al., 1995](#)) can be used to implement this parallelization strategy; given the graph, the inspector finds a conflict-free schedule. Unfortunately, in realistic data sets, most keywords occur in most documents, so the bipartite graph is fairly dense and there are usually very few documents that can be processed in parallel with this parallelization strategy. Similarly, speculative parallelization will not find much parallelism either.

One way to cut this Gordian knot is to change the semantics of loads and stores to values at keywords to permit more parallelism in the update loop. The machine learning community has explored three variations of this theme.

- Local ([Agarwal et al., 2014](#)). Documents are partitioned between threads but each thread has a local copy of values associated with all keywords. Threads update keyword values independently. Periodically, keyword values from different threads are merged together using an application-specific merge function.
- Hogwild ([Recht et al., 2011](#)). Documents are partitioned between threads and there is only one copy of keyword values. Threads update keyword values without synchronization, and the cache-coherency protocol in hardware serializes writes to the same cache line in some order.
- Stale synchronous ([Ho et al., 2013](#)). This is a bulk-synchronous strategy. The program

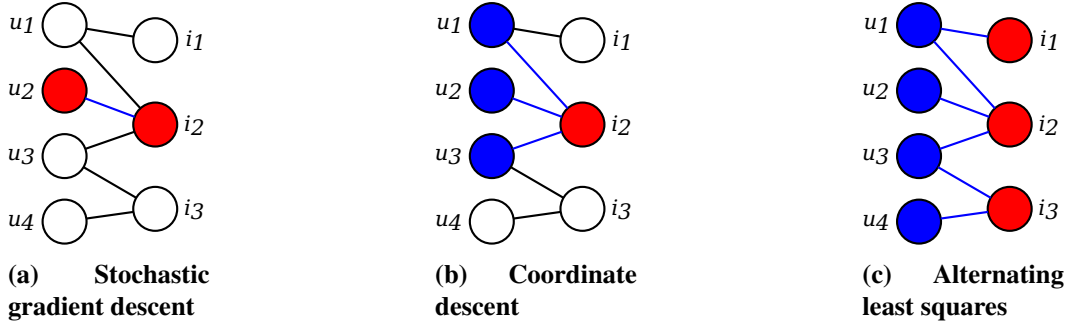


Figure 2.11: Data access patterns for the operator in different matrix completion algorithms. Red indicates values being updated. Blue indicates values being read.

is executed in rounds; in each round, threads read keyword values computed in the last round. Updates to keyword values are accumulated locally to each thread, and at the end of the round, all updates for a given keyword are merged for use in the next round.

2.3.11 Matrix Completion

The following matrix completion problem underlies many modern recommender systems. One is given a partially observed $m \times n$ ratings matrix $A \in \mathbb{R}^{m \times n}$, where m denotes the number of users and n the number of items. Let $\Omega \subseteq \{1 \dots m\} \times \{1, \dots, n\}$ denote the observed entries of A , i.e., $(i, j) \in \Omega$ indicates that user i gave item j a rating of A_{ij} . The goal is to predict accurately the unobserved ratings.

One popular “latent factor” model for matrix completion finds matrices $W \in \mathbb{R}^{m \times k}$ and $H \in \mathbb{R}^{n \times k}$ with $k \ll \min(m, n)$ such that $A \approx WH^\top$ by minimizing the following objective function

$$f(W, H) := \frac{1}{2} \sum_{(i,j) \in \Omega} \left\{ \left(A_{ij} - \mathbf{w}_i^\top \mathbf{h}_j \right)^2 + \lambda \left(\|\mathbf{w}_i\|^2 + \|\mathbf{h}_j\|^2 \right) \right\}, \quad (2.1)$$

where \mathbf{w}_i and \mathbf{h}_j denote the i -th and j -th row of W and H respectively, and λ is a scalar parameter.

In the operator formulation, the matrix A is considered to be a bipartite graph in which the nodes consist of either users or items, and an edge with weight A_{ij} indicates that the i -th user has given a rating of A_{ij} to the j -th item (see [Figure 2.11](#)). Three common ways of solving this matrix completion problem can be expressed with the operator formulation.

Stochastic Gradient Descent (SGD) SGD when applied to [Equation 2.1](#) performs the following update operations:

$$\begin{aligned}\mathbf{w}_i &\leftarrow \mathbf{w}_i - \eta \left((A_{ij} - \mathbf{w}_i^\top \mathbf{h}_j) \mathbf{h}_j + \frac{\lambda}{|\Omega_i|} \mathbf{w}_i \right) \\ \mathbf{h}_j &\leftarrow \mathbf{h}_j - \eta \left((A_{ij} - \mathbf{w}_i^\top \mathbf{h}_j) \mathbf{w}_i + \frac{\lambda}{|\bar{\Omega}_j|} \mathbf{h}_j \right)\end{aligned}$$

where η is a scalar step-size, and $|\Omega_i|$ (resp. $|\bar{\Omega}_j|$) denotes the number of observed entries in the i -th row (resp. j -th column) of A . All edges of the graph are the active, and they can be scheduled in different orders such as cyclic, randomized or prioritized order. The operator can be applied in parallel to any set of edges which do not share a vertex in common. For more rapid convergence, a variety of priority functions can be used. For example, users who have rated a lot of items can be processed before others.

Unlike SGD when applied to the SVM problem, which accesses a document and all its keywords, the SGD update for matrix completion only needs to access data associated with an edge and its endpoints. This is because the model parameters for the matrix completion problem are much sparser than for the SVM problem (i.e., \mathbf{w} versus $\mathbf{w}_i, \mathbf{h}_j$).

Coordinate Descent (CD) CD will apply the following update operator to every node w_{it} of \mathbf{w}_i and h_{jt} of \mathbf{h}_j respectively:

$$w_{it} \leftarrow \frac{\sum_{j \in \Omega_i} ((A_{ij} - \mathbf{w}_i^\top \mathbf{h}_j) + w_{it} h_{jt}) h_{jt}}{\lambda + \sum_{j \in \Omega_i} h_{jt}^2}$$

$$h_{jt} \leftarrow \frac{\sum_{i \in \bar{\Omega}_j} ((A_{ij} - \mathbf{w}_i^\top \mathbf{h}_j) + h_{jt} w_{it}) w_{it}}{\lambda + \sum_{i \in \bar{\Omega}_j} w_{it}^2}$$

In the most common variant, vertices are cycled through repeatedly. The operator can be applied in parallel to any set of active nodes and their neighbors as long as they don't share a vertex that has to be updated.

Alternating Least Squares (ALS) The ALS operator for \mathbf{w} is

$$\mathbf{w}_i \leftarrow \left(H_{\Omega_i}^\top H_{\Omega_i} + \lambda I \right)^{-1} H^\top \mathbf{a}_i$$

for $i = 1, 2, \dots, m$, where H_{Ω_i} denotes the sub-matrix of H formed by selecting the rows $j \in \Omega_i$, and I is the identity matrix, and \mathbf{a}_i is the i -th row of A . A symmetric update can be derived for \mathbf{h}_j . The algorithm repeatedly cycles through the vertices of the graph and applies the updates. The operator can be applied in parallel to all users but it requires barrier synchronization after cycling through the users before it can be applied to all the items.

Chapter 3

Parallel Programming Models

Abstractions for parallelism have a rich history. One of the oldest is the dependence graph. The idea of a dependence or task graph goes back to at least the 1950s when the US Navy used them for project management (Jarnagin, 1960). Central to a dependence graph is that dependencies are ordered or directed. Task a depends on task b . This directed representation appeared in software program abstractions as early as the control flow graph (CFG) (Allen, 1970) in the early 1970s. Around the same time, but independently, the database community developed systems based on serializability, which is an unordered abstraction for dependencies. Task a and task b should appear to execute in some order. These models of computation only began to intersect in the early 1990s with the proposal of transactional memory (TM) (Herlihy and Moss, 1993), a mechanism designed to provide the database notion of transactional execution to programs in general, but it would be at least a decade before practical TM systems became available.

The 1990s also saw sustained interest in auto-parallelization, which is the idea that a compiler would take a program with sequential semantics and find opportunities for parallelization automatically. Prior to this, parallelism was usually expressed explicitly either through direct use of low-level machine primitives (e.g., vector instructions) or through a program abstraction like data-parallelism (Hillis and Steele, 1986). Due to the difficulties

of producing good parallelizations automatically, the most visible auto-parallelizing system, HPF (Kennedy et al., 2007), never gained any traction. Performance of HPF programs depended on careful data placement. Users annotated data structures with placement policies, and a compiler generated communication schedules. However, the placement policies available were drawn from a small set and not extensible.

The mid-2000s saw a new growth (Charles et al., 2005; Allen et al., 2007; Chamberlain et al., 2007) of parallel programming models as the DARPA High Productivity Computing Systems project sought ways to achieve high performance with high programmer productivity. To avoid reliance on the compiler to find parallelism, these new languages required programmers to indicate parallelism explicitly. The underlying model of parallelism was generally a fusion of data-parallelism with task dependencies.

As no parallel programming model has become dominant, this expansion of parallel programming models continues to this day with proposals for general (Dean and Ghemawat, 2004; Bauer et al., 2012) and application-specific (Kale and Krishnan, 1993; Low et al., 2010; Kepner and Gilbert, 2011) models for parallelization. In most parallel programming models, the system takes sole responsibility for scheduling parallel tasks, and there is one fixed scheduling algorithm that it uses. For instance, Cilk (Blumofe et al., 1995) uses randomized workstealing with thread-local deques. In systems that do provide a number of schedulers like OpenMP (Dagum and Menon, 1998), the choices are from a small fixed menu (e.g., block, block-cyclic). From the discussion in Chapter 2, it is clear that more flexibility in scheduling is needed to cover the diversity of parallel algorithms. For instance, the different algorithms for SSSP can be seen as the same operator with different scheduling policies.

If we want a variety of scheduling policies, how should this be accomplished? Since schedulers are complex pieces of concurrent code, one cannot expect general-purpose programmers to implement them directly. There should be an intermediate point between implementing schedulers from scratch, which is error-prone, and selecting from a small set of

schedulers, which is insufficient for many algorithms.

Additionally, the implementation of the operator and its data structures can be influenced by which schedules are possible, or in other words, some data consistency models are more efficient to implement than others. E.g., it is much easier to implement a shared counter under bulk-synchronous consistency than to implement one under linearizability. How should these opportunities be exploited when schedules fall in these special cases?

The Galois system, which is described in the following chapters, is one answer to these questions. To address the need for a variety of schedulers, the Galois system provides an extensible library of scheduling policies and a scheduler synthesizer to produce a concurrent scheduler from a high-level specification. To address the implementation of the operator and its data structures, the Galois system uses exclusive locking for serializable schedules. For more relaxed consistency models, it uses a methodology called *diffracted state* to produce efficient data structure implementations even in the presence of frequent sharing.

Chapter 4

The Galois System

The Galois system is software library and runtime system for parallelizing programs in shared-memory based on the operator formulation ([Section 2.2](#)). Its main goal is the efficient parallelization of general unordered programs with high programmer productivity (i.e., a small amount of additional programmer effort). This chapter describes the principles that underlie the design of the Galois system, and the next two chapters describe how those principles are put into practice in the implementation of parallel data structures ([Chapter 5](#)) and scheduling ([Chapter 6](#)).

The Galois system is designed for shared-memory multicore systems. This means that there is a single address space of memory and there are multiple threads of execution. Threads are assigned to cores of a multicore processor, and there may be multiple processors per machine. Communication between processors may be slower than communication between cores.

4.1 Principles of High-Performance Parallelism

This section articulates two design principles that must be embodied in scalable parallel programs. These principles arise from the fact that the most common limiting factor on

scalability is data movement. Communication adds costs that increase as the number of threads or machines increases, so the best way to improve the scalability of a program is to (1) reduce communication and (2) to be tolerant of communication costs. Concretely, this means the following for parallel programming models.

Principle 4.1 (Disjoint accesses). *Tasks that are disjoint at the logical level should be disjoint at the physical level.*

This principle is a guide for reducing or eliminating conflicts between tasks that logically should be able to execute concurrently: it says that concurrent tasks should not conflict if they operate on disjoint data. For example, tasks that add and remove different items to a bag should not interfere because they operate on disjoint data, even though they operate on the same data structure. An implementation of a concurrent bag that uses a single lock, for example, violates this principle. Here and in the remainder of this section, the term conflict is used in a general sense. A conflict can mean introducing synchronization to preserve program semantics or it can mean writing to a shared cache line which causes an invalidation at the cache coherency protocol level.

Principle 4.2 (Virtualized tasks). *Tasks should be virtualized.*

Tasks are virtualized if their execution is not tied to a particular thread or schedule. This principle is a guide for ensuring that threads do useful work even in the presence of conflicts or communication delays. When this happens, implementations have flexibility in choosing when tasks should execute. For instance, a thread can set aside task and perform other work instead of waiting for a response for a remote memory request.

These principles may not be surprising, but parallel programs do not necessary satisfy either principle. Consider the STAMP benchmark suite (Cao Minh et al., 2008), a widely used suite of programs for evaluating parallel transactional memory (TM) programs, and to illustrate the key ideas, consider the yada benchmark in STAMP, which performs Delaunay mesh refinement of a triangular mesh. Figure 4.1 shows a simplified version. A


```

1  Workset wl
2
3  void func(int thread_id):
4      while true:
5          Task t = atomic { wl.pop() }
6          if !t:
7              break
8          Tasks newTs = atomic { work(t, thread_id) }
9          for Task nt in newTs:
10             atomic { wl.push(nt) }
11
12  thread_run(func, num_threads)

```

Figure 4.1: STAMP programming model

workset tracks the initial work and work generated during the progression of the algorithm. Conceptually, each task represents a bad triangle. A workset of bad triangles, implemented using a linked-list, is populated by walking over the initial mesh and testing each triangle for badness using simple geometric tests on its vertices.

To process and eliminate a bad triangle, a small neighborhood of triangles surrounding the bad triangle is identified (i.e., the cavity) and deleted from the mesh. The region previously occupied by these triangles is then re-triangulated, and the new triangles are added to the mesh. Some of these newly created triangles may be bad; if so, they are added to the workset. In the pseudocode of [Figure 4.1](#), this functionality is implemented by the function *work*.

Parallelism in this algorithm arises from the fact that each cavity is usually a small region of the overall mesh, so bad triangles whose cavities do not overlap can be processed in parallel. However, it is difficult to tell a priori whether or not the cavities of two bad triangles will overlap, so static parallelization does not work. Instead, we can consider the processing of each bad triangle to be a transaction and execute transactions speculatively, leaving it to the TM system to detect and recover from conflicts on the fly.

In yada, therefore, there are two concurrent data structures: the mesh and the workset of bad triangles. In the pseudocode shown in [Figure 4.1](#), each thread executes the func-

tion *func*. The body of this function is a while loop that terminates when the workset is empty. A thread pops a work-item from the workset using a transaction to synchronize with other threads that may be manipulating the workset at the same time. The function *work* is then called to perform the computation; in *yada*, this is the processing of the bad triangle, which must be performed transactionally since the mesh is being updated by multiple threads simultaneously. Finally, any newly created bad triangles are added to the workset using another transaction.

The programming model and data structures in the STAMP implementation of *yada* violate both principles for scalable parallel performance introduced earlier.

Workset push and pop operations from different threads conflict because a linked-list is used to represent the workset. At the logical level however, two threads that pop work items from the workset touch different data items, so they should not conflict according to the disjoint access principle. Similar considerations apply to the representation of the graph data structure: as long as cavities do not overlap, transactions should not conflict. However, the graph representation used in STAMP, which uses a linked-list to store nodes, introduces spurious conflicts, violating the principle of disjoint access.

In addition, the explicitly threaded programming model of [Figure 4.1](#) violates the principle of virtualized transactions. Once a transaction is attempted by a thread, there is no way for the thread to put aside that transaction if it aborts and do some other work; instead, the same thread must keep trying to finish that transaction until it commits. This limits scheduling freedom and reduces processor utilization.

While this discussion has focused on *yada*, similar issues arise in the other STAMP benchmarks. And while STAMP uses transactions to implement synchronization, a more traditional parallel program using locks and threads would follow a similar structure, and similar issues arise in parallel programs written by non-performance experts.

So, what can be learned from this example? First, most parallel programming models focus on scheduling parallelism, but addressing the performance problems in *yada* re-

quire addressing issues with scheduling *and* data structures. Second, addressing these issues is not the intended use of TM but solving them is important for performance. Once yada was modified to conform to the two performance principles, its execution time was reduced by a factor of 62 on a 32 thread execution (see [Section 8.1](#)). Following these principles is important for improved performance.

Now, rewriting programs by hand to conform the scalability principles introduced here may be feasible for a small number of performance critical and widely used applications, but how can these performance insights be applied to the large number of parallel programs being written today? The solution proposed by this dissertation is a programming model approach where judicious application of abstraction encourages programmers to write scalable programs.

4.2 Separation of Concerns

The goal of the Galois system is efficient and high-productivity parallelization. Productivity is achieved through abstraction. Certain information is hidden from the programmer and its implementation becomes the responsibility of the programming system, while other information is made explicit and must be provided by the programmer. When a system requires very little from the programmer, it promotes high productivity.

High productivity tends to be in tension with high performance. The less information given by the programmer, the more a system must infer and the more likely that the inferred information is suboptimal with respect to performance. Conversely, a dedicated programmer with a sufficient amount of time can produce a program from scratch that performs as well as or outperforms any one created by a programming system. The scientific and engineering question becomes choosing where to draw the line between programmer and system in such a way that balances productivity with performance.

One classic division of labor between programmer and system is the abstract data type (ADT) ([Liskov and Zilles, 1974](#)). Programs are written against interfaces (e.g., list, set,

map) that specify a set of methods and their behavior (e.g., list append, list remove), but the implementation details are hidden, and programmers can interchange different ADT implementations (i.e., data structures) with, at most, modest program changes. Niklaus Wirth’s aphorism “Algorithms + Data Structures = Programs” (Wirth, 1978) memorably alludes to the importance of ADTs to programming.

One reason for the success of ADTs is due to the fact that they factor “what” is done from “how” it is done. Programmers can focus on solving their problems without getting bogged down in the details. Another reason for the success of ADTs is due to the fact that they can grow organically. Language designers cannot select in advance all the components that programmers will need. ADTs provide a way for programmers to develop components themselves, as the need arises, without having to leave the programming system.

The common programming abstractions for parallelism—threads, data-parallel loops, task graph scheduling—do not satisfy these criteria. Threads closely match the underlying machine semantics, so users must deal with low-level concerns like memory models and data races. This contravenes the principle of abstraction. Not all parallel programs can be expressed solely as data-parallel loops or task graphs (Hassaan et al., 2015), so programmers must look outside these abstractions to parallelize certain codes. This contravenes the principle of growth.

The operator formulation offers one solution to this problem by abstracting a parallel task from its implementation and is complete with respect to the applications studied in this dissertation.

The Galois system is an implementation of the operator formulation for unordered algorithms within a sequential programming language. Application programmers write parallel programs without explicit parallel programming constructs like threads and locks. In the current system, the sequential language is C++. Key features of the system are the following.

- Application programmers specify parallelism *implicitly* by using an unordered-set

```

1 Galois::Graph::FirstGraph<Node,Edge> G;
2
3 struct Operator {
4     void operator()(Point p, Galois::UserContext<Point>& ws) {
5         Triangle t = findTriangleContaining(p);
6         Cavity c(t);
7         c.expand();
8         c.retriangulate();
9         G.update(c);
10    }
11 };
12
13 void main() {
14     std::vector<Point> points = readPointsFromFile();
15     G = constructInitialGraph();
16     Galois::for_each(points.begin(), points.end(), Operator());
17 }

```

Figure 4.2: Example Galois program in C++. This is the concrete implementation of pseudocode in [Figure 2.6](#).

iterator which iterates over a workset of active nodes. The workset is initialized with a set of active nodes before the iterator begins execution. The execution of a iteration can create new active nodes, and these are added to the workset when that iteration completes execution.

- The body of the iterator is the implementation of the operator, and it is an imperative action that reads and writes global data structures. Iterations are required to be *cautious*: an iteration must read *all* elements in its neighborhood before it writes to *any* of them. This is not a significant restriction since the natural way of writing applications tends to produce cautious iterations.
- The relative order in which iterations are executed is left unspecified in the application code; the only requirement is that the final result should be identical to that obtained by executing the iterations sequentially in some order. An optional application-specific priority order for iterations can be specified (see [Section 6.2](#)), and the implementation tries to respect this order when it schedules iterations.

- The system exploits parallelism by executing iterations in parallel. To ensure serializability of iterations, programmers must use a library of built-in concurrent data structures for graphs, worksets, etc. (see [Chapter 5](#)). These library routines expose a standard API to programmers, and they implement lightweight synchronization to ensure serializability of iterations (see [Section 5.2](#)).

[Figure 4.2](#) shows an example Galois program in C++ for the pseudocode given in [Figure 2.6](#). The program is implicitly parallel. Locks and synchronization are performed on the programmer's behalf by the data structure and scheduling library. Since data structures are abstract and synchronization and scheduling are implicit, the Galois system is free to use scalable implementations without changing the program; compare this model with the STAMP program described earlier ([Figure 4.1](#)). There, important decisions about scheduling and data structure implementations are fixed by the programmer and cannot be changed without changing the meaning of the program.

Chapter 5

Parallel Data Structures

The Galois system is based around a library of data structures and scheduling policies. This chapter describes the basic data structures used in the Galois system. The main challenge in implementing data structures is following the disjoint access principle ([Principle 4.1](#)). For example, a red-black tree may be a reasonable implementation of an associative map for sequential programs, but performing tree operations in parallel requires careful synchronization, and shared accesses on the root of the tree can be a performance bottleneck. A more scalable implementation is a hashtable, which is a more decentralized alternative, since logically disjoint operations can be implemented with operations on disjoint memory areas.

Of course, some data structure operations require communication to implement; e.g., finding the minimum value in a set in parallel requires some communication between threads. When communication must be done, Galois data structures try to accomplish it through the lowest cost communication pathways in the machine. In shared-memory multi-core machines, memory is organized hierarchically (see [Figure 5.1](#)), and the communication cost between two threads is a decreasing function of the height of their least common ancestor in the memory hierarchy, where the height of the root is zero. Communication between

Portions of this chapter have previously appeared in ([Nguyen et al., 2013](#)), where the Galois memory allocator was originally described.

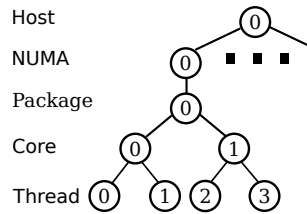


Figure 5.1: Example memory hierarchy for a multicore machine

threads that share the same core is faster than communication between threads that only share the same package, which in turn is faster than communication between threads that have nothing in common except that they are running on the same machine. Here, communication refers to reading and writing the same cache line. Large-scale non-uniform memory access (NUMA) machines may have more levels of hierarchy than shown in the figure, although the general cost pattern remains the same. Thus, when global communication must be done, the cheapest way to implement it is to use a communication tree that follows the memory hierarchy. This is similar to how distributed reduction algorithms are implemented.

All Galois data structures are built on top of a scalable memory allocator that is described in the next section. [Section 5.2](#) describes how Galois data structures implement transactional execution by using an exclusive locking discipline. [Section 5.3](#) introduces a methodology called *diffracted state* for reducing communication costs by following a machine’s communication hierarchy. Finally, [Section 5.4](#) and [Section 5.5](#) go in depth into the implementation of two specific data structures, an approximate value store and a sparse graph.

5.1 Memory Allocation

Galois data structures are based on a custom scalable memory allocator. While there has been considerable effort towards creating scalable memory allocators ([Berger et al., 2000](#); [Michael, 2004](#); [Schneider et al., 2006](#)), existing general solutions do not scale to large-scale multi-threaded workloads that are very allocation intensive nor do they directly address non-

uniform memory access (NUMA) concerns, which are important for even modestly sized multi-core architectures. Recently, there has been some effort in designing NUMA-aware data structures specifically for local computations over graphs (Zhang et al., 2015).

Providing a general, scalable memory allocator is a large undertaking, particularly because Galois supports morph applications that modify graphs by adding and removing nodes and edges. For most applications, memory allocation is generally restricted to two cases: allocations in the runtime (including library data structures) and allocations in an activity to track per-activity state.

For the first case, the Galois runtime system uses a slab allocator, which allocates memory from pools of fixed-size blocks. This allocator is scalable but cannot handle variable-sized blocks efficiently due to the overhead of managing fragmentation. The second case involves allocations from user code, which may require variable-sized allocation but also have a defined lifetime, i.e., the duration of an activity. For this case, the Galois system uses a bump-pointer region allocator.

The slab allocator has a separate allocator for each block size and a central page pool, which contains huge pages allocated from the operating system. Each thread maintains a free list of blocks. Blocks are allocated first from the free list. If the list is empty, the thread acquires a page from the page pool and uses bump-pointer allocation to divide the page into blocks.

The page pool is NUMA-aware; freed pages are returned to the region of the pool representing the memory node they were allocated from.

Allocating pages from the operating system can be a significant scalability bottleneck (Yoo et al., 2009; Clements et al., 2012), so to initialize the page pool, each application preallocates some number of pages prior to parallel execution; the exact amount varies by application.

The bump-pointer allocator manages allocations of per-activity data structures, which come from temporaries created by user code in the operator. This allocator supports stan-

dard C++ allocator semantics, making it usable with all standard containers. The allocator is backed by a page from the page pool. If the allocation size exceeds the page size (2 MB), the allocator falls back to `malloc`.

Each activity executes on a thread, which has its own instance of the bump-pointer allocator. The allocator is reused (after being reset) between iterations on a thread. Since the lifetimes of the objects allocated are bound to an activity, all memory can be reclaimed at once at the end of the activity.

5.2 Exclusive Locking

The Galois system uses exclusive locking to implement transactional execution. This contrasts with transactional memory (TM) systems, which ensure transactional execution by monitoring reads and writes to memory. One challenge tackled by the TM community is supporting a high amount of concurrency by sometimes allowing a read in one transaction to proceed in the presence of a concurrent read or write to the same address in another transaction.

In multicore architectures, writing to a shared cache line is a potential scalability bottleneck, and scalable programs are typically written (or rewritten) to not have such sharing (Clements et al., 2013), e.g., replacing red-black trees with hashtables. Having a conflict detection scheme that permits activities to write to a shared cache line enables a level of concurrency at the task level that is not scalable at the cache coherency level.

An alternative is to simply not permit activities to proceed if they access the same memory locations. One such scheme is *exclusive locking*. Whenever an activity reads or writes memory, it marks that location with its activity id. If another activity has already marked the location, the current activity *aborts*, rolling back its state changes and clearing its marks. When an activity finishes, it also clears its marks.

Figure 5.2 shows pseudocode for this mark functionality, which must be called before reading or writing a memory location. The value `-1` is used to indicate that the location

```

1  struct ExclusiveLocation :
2      T value
3      T oldValue
4      int owner
5      ExclusiveLocation* next
6
7  ThreadLocal<int> myId
8  ThreadLocal<ExclusiveLocation*> myLocs
9
10 void mark(ExclusiveLocation* l):
11     while true:
12         if l->owner == myId:
13             return
14         else if l->owner != -1:
15             rollback()
16             // returns to scheduler to reschedule
17             raiseAbort()
18         else if compareAndSwap(&l->owner, -1, myId):
19             // if l->owner was -1 and it is
20             // successfully updated to myId
21             l->oldValue = l->value
22             l->next = myLocs
23             myLocs = l
24     return

```

Figure 5.2: Marking a location with exclusive locking

has not been marked by any activity. Exclusive locations acquired by an activity are maintained in a linked-list. To clear its marks, an activity walks its list of marked locations, sets the owner fields back to the unacquired value, and resets the next fields to null. Rollback is similar to clearing marks except that the activity additionally restores the value to the previously stored old value. One optimization is to use transactional boosting (Herlihy and Koskinen, 2008) and use undo logs to rollback state. In the Galois system, an activity is assumed to be *cautious*. That is, it reads all its locations before modifying any of them. In this case, there is no state to rollback when a conflict occurs because no writes have happened yet.

Exclusive locking *fails fast* on activities that other conflict detection schemes would permit to continue. However, if a program follows the disjoint access and virtualization principles, the number of conflicts should be small because (1) activities access mostly disjoint

```

1  struct Graph :
2      // ...
3      NodeData& getData(GraphNode n, MethodFlag m = ALL):
4          // Return data associated with graph node n
5
6      Graph g
7      GraphNode n
8
9      g.getData(n)          // Get data with full transactional execution
10     g.getData(n, NONE) // Get data without transactional execution

```

Figure 5.3: Using method flags to indicate desired support for transactional execution

data and (2) aborted activities can be rescheduled to reduce conflicts. Exclusive locking also reflects the realities of modern hardware. Writes to shared cache lines are scalability bottlenecks, and exclusive locking reflects the performance model of the underlying hardware directly in the programming model. This makes potential problems more obvious, which encourages programmers to address them early.

A useful optimization is to bypass exclusive locking when an operator only performs a simple update to a machine word or when transactional execution is not needed at all. For these cases, all Galois data structure methods have an optional parameter that indicates whether an operation always or never acquires locks. Experienced users can disable locking and use machine atomic instructions if desired (see [Figure 5.3](#)).

An objection to exclusive locking is that it produces conflicts with read-only workloads. Locations that are always read-only can usually be determined statically (e.g., ([Lattner et al., 2007](#))). For situations where a location is sometimes read and sometimes written, TMs that allow concurrent reads typically use timestamp-based validation ([Dice et al., 2006](#); [Riegel et al., 2006](#)), which has its own scalability cost because a global counter is atomically updated by each thread on commit. The cost of synchronization can be eliminated by using a hardware clock ([Riegel et al., 2007](#)) or hardware performance counters ([Ruan et al., 2013](#)), but in these cases, transactions sometimes must wait to commit. One could instead turn to thread-local clocks ([Avni and Shavit, 2008](#)) but at the cost of false conflicts.

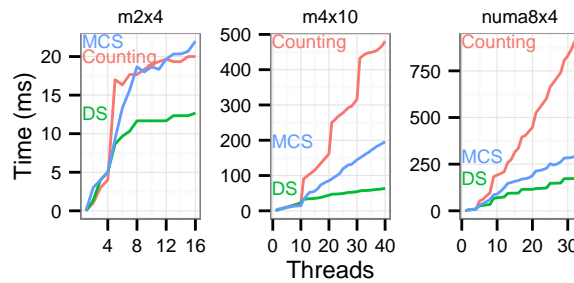


Figure 5.4: Execution time of different barrier implementations for executing $16 \cdot 1024$ invocations

5.3 Diffracted State

Diffracted state is a methodology used by Galois data structures to reduce communication costs for highly shared data structures like schedulers and approximate value stores (see [Section 5.4](#)). The basic idea is to associate state (replicas) at the various nodes of the memory hierarchy of a machine (see [Figure 5.1](#)). Data structure operations refer to replicas by node and strive to ensure that the most frequently executed operations refer to nodes closest to the executing thread. If an operation wants a more global view of state, it *merges* multiple replicas together with an operation-specific merge function.

As an example, consider the implementation of a barrier on a multicore machine. The standard implementation in the pthread library uses a mutex and a single count variable. A more efficient implementation would be to distribute the single count variable into a tree of variables with a variable for each thread in the machine. Each variable is a flag indicating whether the children of that node are waiting in the barrier. A thread only accesses flag variables of itself and its children in the tree. This is the approach taken by the classic MCS tree barrier ([Mellor-Crummey and Scott, 1991](#)). One implementation following diffracted state would be to create replicas for each thread and package of a machine. In contrast to the MCS tree barrier, state is explicitly mapped to the memory hierarchy.

[Figure 5.4](#) shows the execution time of the counting, MCS and diffracted state (DS) barrier. The machines used for the evaluation are: (1) m2x4, a two processor, four cores per

processor Intel system, (2) m4x10, a four processor, ten cores per processor Intel system, and (3) numa8x4, an eight processor, four cores per processor Intel system where every two processors are packaged into boards and boards are connected using a NUMA interconnect. The diffracted state implementation used by Galois is about twice as fast as the classic MCS tree at scale. This is due to the fact that the communication pattern more closely matches the actual machine hierarchy. Also, the MCS barrier uses a binary tree, while the typical number of threads per package is usually greater than two, so the overall tree height of the barrier has been reduced in the diffracted state implementation.

A barrier is a simple example of the principle of diffracted state. The next section introduces a more complex example: the implementation of approximate value stores.

5.4 Approximate Value Stores

A *value store* is a collection of locations that can be read and written. An *asynchronous update* is a procedure that reads a set of locations and writes to a set of locations. It is asynchronous because the value read at a location may not correspond to the most recent value written to that location; instead some previously written value may be returned. An *approximate value store* is a value store that may return values that have not been previously written. A value store implementation may provide progress guarantees that bound what values may be returned.

Many problems can be solved with the iterative application of asynchronous updates (Bertsekas and Tsitsiklis, 1989), and such asynchronous algorithms are attractive because they are robust to communication delays. For instance, a value store for distributed memory may buffer updates locally and only periodically exchange updated values with other distributed nodes. When there are multiple updated values for a location, those values are reduced to a single value based on an algorithm-specific reduction function.¹ This ap-

¹ Given a set S of multiple values, if the algorithm-specific reduction function returns an element of S (e.g., minimum value), the algorithm is simply asynchronous. If the function may return a value not in S (e.g., mean value), the algorithm is asynchronous and approximate.

proach underlines recent software frameworks for graph analytics (Malewicz et al., 2010; Gonzalez et al., 2012; Kyrola et al., 2012).

Although asynchronous algorithms are robust to communication delays, the overall speed of convergence depends on how quickly information propagates through the value store. Using value stores that reduce communication by propagating updates slowly cause asynchronous algorithms to converge slower than using value stores that propagate updates faster.

Finding the right amount of communication that balances cost with convergence is a challenge, but one simple heuristic is to allow communication when it is cheap relative to the average cost. With respect to the machine topology, this means allowing frequent communication within a package but restricting communication between packages. One possible implementation is to associate a value store with each package of the machine. Threads read and update their per-package value store, and periodically the per-package stores are merged with an algorithmic-specific reduction function. Since the updates are asynchronous, they do not need to be explicitly synchronized, and the underlying hardware coherence protocol can resolve concurrent writes to the same store.

Compared to strictly thread-local updates, this per-package value store allows updates between nearby threads to propagate quickly without waiting for the periodic reduction step. An alternate topology mapping would be to have a single value store. This eliminates the need for a periodic reduction step and, at first glance, improves the propagation of updates because updates between packages are sent without buffering. However, the overall communication cost increases because communication between packages is not controlled. Depending on the algorithm, the increased communication cost may outweigh any improvement to the convergence rate.

To evaluate the trade-off between communication and convergence, consider a popular asynchronous algorithm, stochastic gradient descent (SGD), which is general technique for solving an optimization problem by taking small steps along an approximate gradient.

SGD itself can be used to solve many different problems. For this case study, consider the problem of learning a linear support vector machine (SVM) model for use in binary classification (see [Section 2.3](#)). Briefly, the input to the algorithm is a set of documents $\mathbf{x}_i \in X$, which are vectors of length M , and a label $y_i \in \{-1, 1\}$ for each document indicating whether the document is a positive or negative example of classification being learned. The output is a vector w also of length M such that $\mathbf{w} \cdot \mathbf{x}_i \cdot y_i > 0$ for as many documents as possible. The vector \mathbf{w} can be used to classify new documents by computing the product $w \cdot \mathbf{x}_j$. If this value is greater than zero, the document is a positive example; otherwise, it is a negative example. SGD solves this problem by starting at an initial guess for $\hat{\mathbf{w}}$ and processing documents individually. Each document processed produces an update to $\hat{\mathbf{w}}$, and eventually $\hat{\mathbf{w}}$ converges to \mathbf{w} . Parallelism exists in this algorithm when documents are sparse, which means not all elements of $\hat{\mathbf{w}}$ need to be updated for each document.

Since SGD is an asynchronous algorithm, it can be implemented with many different value stores. Consider four possible implementations, three of which have been previously considered for parallelizing SGD. The first is thread-local value stores with periodic merging ([Agarwal et al., 2014](#)). The second is “hogwild” updates ([Recht et al., 2011](#)) where there is a single value store updated by all threads, and the hardware coherence sorts out concurrent updates. The third is stale synchronous ([Ho et al., 2013](#)) in which execution proceeds in rounds. Locations read in one round return values written in the previous round. Writes to locations are accumulated. When a round ends, the accumulated writes are merged to form the values to be read in the following round. The new implementation is a store using the diffracted state methodology introduced in the previous section. It uses per-package value store replicas with high value updates communicated between packages in a ring topology. In contrast to previous value store implementations, the diffracted state store explicitly maps data to the memory hierarchy.

[Figure 5.5](#) shows how accuracy (i.e., prediction quality) improves with time for a small input (news20 ([Keerthi and DeCoste, 2005](#)): 20 K documents, 1.4 M features) and a

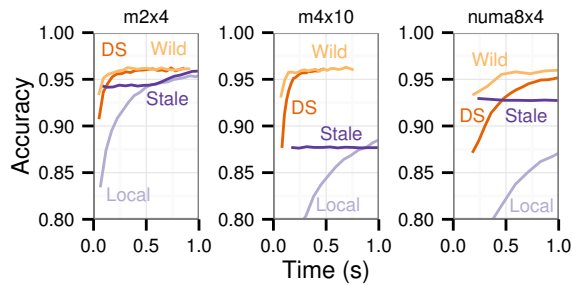


Figure 5.5: Convergence of SVM-SGD training on small input on three different machines with maximum number of threads on each machine

large data set (webspam (Webb et al., 2006): 350 K documents, 16.6 M features). The machines used for the evaluation are the same as in Section 5.3. Prediction accuracy is measured by training on four-fifths of the documents and testing on the last fifth. For reference, the widely used liblinear library (Fan et al., 2008) takes about 0.833 seconds to achieve an accuracy of 0.965 on the small input on machine m4x10 and 93.9 seconds to achieve an accuracy of 0.993 on the large input. However, it uses a different training algorithm, GLMNET, which will be discussed shortly.

Value store implementations that communicate frequently like the diffracted state (DS) and hogwild (Wild) approaches improve their accuracy more quickly than approaches that communicate less frequently like bounded staleness (Stale) and thread-local stores (Local). Convergence results for the large input are omitted because all of the implementations except for stale synchronous converged to the same accuracy (≈ 0.985) after one round of SGD updates. The stale synchronous approach failed to converge. Thus, for the large input, performance is determined by how fast each implementation can do one round of SGD updates.

Figure 5.6 shows speed-up for executing one round of SGD, where the baseline is the time of the best implementation for one thread. For reference, the time to execute one SGD round with the diffracted state value store on machine m4x10 with 40 threads is 0.048 seconds for the small input and 1.3 seconds for the large input. Comparing Figure 5.5 and

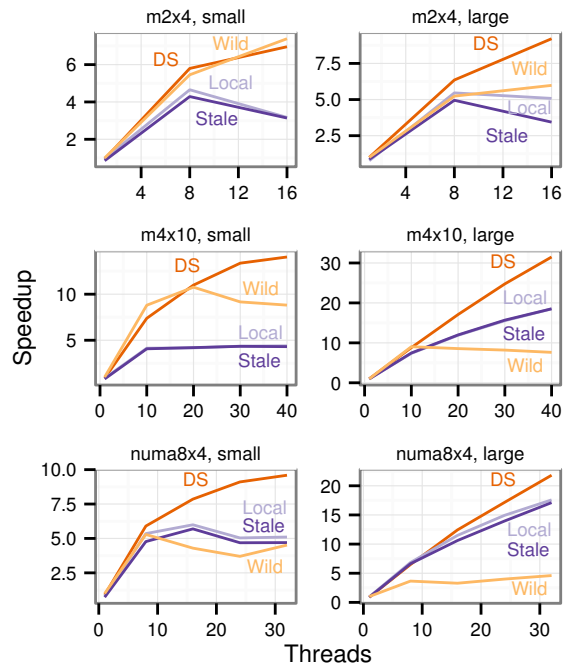


Figure 5.6: Speedup of SVM-SGD training for one SGD round

Figure 5.6 for the small input, one sees that although the hogwild approach produces higher quality results, it is doing so at increasing cost, as shown by the lower speedup numbers for an SGD round. The thread-local and bounded staleness stores also have low speedups because the overhead of merging values between rounds. On the large input, the increased communication of the hogwild approach is not worth the cost, and it is the diffracted state implementation that achieves the best performance.

To show the generality of the diffracted state store, consider logistic regression, an alternate machine learning method for classification. Although SGD can be used in this case as well, it requires frequent evaluation of transcendental functions, which can be avoided by using a different method called GLMNET (Friedman et al., 2010). The core of the method is computing an approximate Hessian, which has a similar data access pattern as SGD except that iterations are over keywords nodes and updates are to values on adjacent documents (in SVM-SGD iterations are over documents and updates are to values on adjacent keywords).

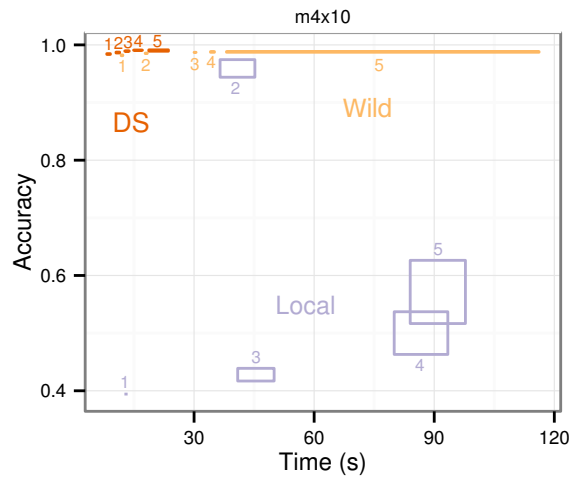


Figure 5.7: Convergence of GLMNET training on large input on machine m4x10 with 40 threads. Boxes indicate the min and max values observed over three runs. Boxes are labeled with the number of Newton iterations executed so far.

Another important difference is that in SGD-SVM, the value store maintains the model parameters themselves, while in GLMNET, the value store maintains elements of the Hessian, which will then be used by subsequent processing stages to update the model parameters. This added level of feedback has the potential to amplify the effect of small approximation errors.

Figure 5.7 shows the convergence of GLMNET with different approximate value stores. The bounded staleness store timed out and is not shown. Compared to SVM-SGD, there is more performance and quality variation as indicated by the width and height of the boxes. On this input and number of threads, the hogwild approach does worse than diffracted state due to the uncontrolled error introduced by racy reads and writes. This error causes subsequent phases of GLMNET to take longer to converge, which is why the fifth iteration with hogwild has such a wide variation in time. An unlucky interleaving of reads and writes can have a large performance impact on downstream phases.

These results show that having a good approximate value store implementation requires balancing communication costs based on the application and machine.

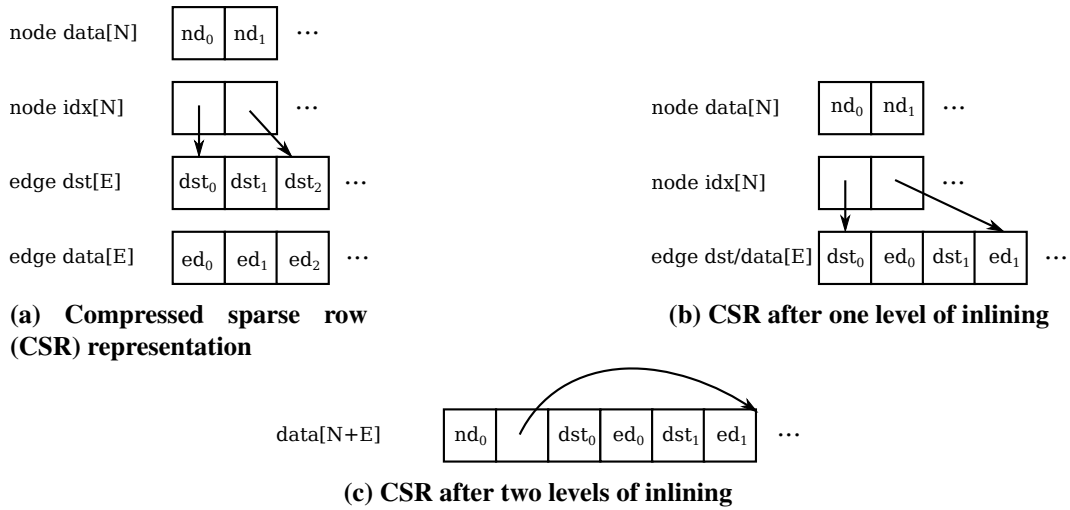


Figure 5.8: Illustration of inlining graph data

5.5 Sparse Graphs

A common data structure is a sparse graph. The Galois system provides a variety of sparse graph implementations based on whether the operator is a local computation or a morph. In this way, sparse graphs can be specialized according to their expected use.

For morph operators, Galois provides a general morph graph that supports concurrent node and edge addition and removal. The graph is implemented as a collection of node objects. Each node contains an array of edges that can grow dynamically. Each edge maintains the label data on the edge and a pointer to the neighboring node. The array of edges stores both edges from neighbors (in-edges) and edges to neighbors (out-edges). For undirected graphs, an in-edge stores a pointer to the corresponding out-edge so that updates to an edge label on one edge are reflected in the matching edge. The memory for this graph is allocated from the memory allocators described in [Section 5.1](#).

The exclusive lock locations described in [Section 5.2](#) are implemented as an additional field on each node of the graph.

When the operator is a local computation, the general morph graph implementation

wastes space on metadata like the resizable number of edges, which is not necessary for local computations. In this case, the sparse graph behaves like a sparse matrix and similar implementations can be used. [Figure 5.8](#) shows one popular sparse matrix representation, compressed sparse row (CSR). It uses four fixed-size arrays: one array for the node data (labels), one index array indicating where the edges for a node can be found, one array for the ids of the neighboring nodes, and one array for the edge data (labels). In local computations, the size and structure of the graph is known in advance, so the memory for these arrays can be allocated all at once from the operating system. One small optimization for graphs that have random access behavior is to interleave the allocation across NUMA nodes to improve memory bandwidth.

Another optimization is to inline the arrays to increase spatial locality. The common pattern of accessing a node and its neighbors in CSR representation requires accessing entries in four different arrays. Inlining as is shown in [Figure 5.8](#) can reduce the number of memory accesses by reducing the number of arrays accessed. Users of the Galois system can select different local computation graphs based on their desired level of inlining.

Chapter 6

Scheduling

Parallel programming models like Cilk ([Blumofe et al., 1995](#)) or OpenMP only provide a small number of scheduling policies, but [Section 2.3](#) illustrates the need for a variety of scheduling policies for algorithms. Efficient algorithms often require application-specific scheduling strategies. This chapter describes how the Galois system supports a variety of scheduling policies through compositional scheduler synthesis.

6.1 Scheduler Building Blocks

This section describes two example building blocks for a scheduler: (1) a topology-aware bag of tasks ([Section 6.1.1](#)), called distributed chunked LIFO, and (2) a topology-aware priority scheduler ([Section 6.1.2](#)), called obim. While these blocks can be used as task schedulers by themselves, the intention is to use them as well as other schedulers in the compositional scheduler synthesis described in [Section 6.2](#) to implement programmer-supplied scheduling policies.

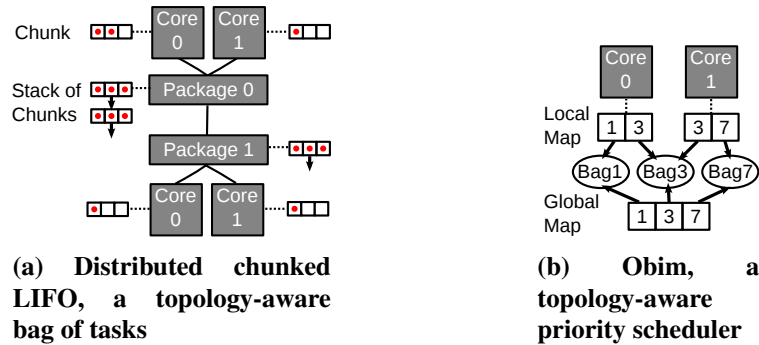


Figure 6.1: Organization of distributed chunked LIFO and obim schedulers

6.1.1 Topology-Aware Bag of Tasks

A common scheduling abstraction is a data structure that allows concurrent insertion and retrieval of unordered tasks. This data structure is called a bag, and it is a basic building block for dynamic scheduling of tasks. There are many possible implementations. One of the most well-known is per-thread dequeues with randomized workstealing, which was popularized by the Cilk system (Blumofe et al., 1995). This section describes another bag implementation, a distributed chunked LIFO, which unlike the original Cilk scheduler, attempts to optimize communication to follow the communication topology of a multicore processor. This is another application of the diffracted state methodology (see Section 5.3). It is possible to manipulate the bag in FIFO-style as well, but to keep this description simple, that possibility is omitted here.

Figure 6.1a outlines the structure of the distributed chunked LIFO and Figure 6.2 gives pseudocode to implement its two main operations: push, which adds a task to the bag, and pop, which retrieves a task from the bag if available.

This chapter assumes throughout that threads are bound uniquely to cores.

- Each thread has a data structure called a *chunk*, which is a ring-buffer that can contain 8–64 tasks (size chosen at compile time). The ring-buffer is manipulated as a stack

Portions of this chapter have previously appeared in (Nguyen and Pingali, 2011), where the Galois synthesis procedure was originally described.

```

1  class DistributedChunkedLIFO:
2      // One chunk per thread
3      PerThread<Chunk*> perThread
4
5      // Chunk embeds a next pointer for the linked list
6      PerPackage<ThreadSafeLinkedList<Chunk*>> perPackage
7
8      // Add a task
9      void push(Task t):
10         // Get thread-local chunk
11         Chunk*& cur = perThread.local()
12         // Push task to local chunk if exists
13         if cur && cur->push(t):
14             return
15         // Otherwise, chunk is full or does not exist
16         if cur:
17             perPackage.local().push(cur)
18         // Create a new chunk and use it
19         cur = new Chunk
20         cur->push(val)
21
22         // Get next task
23         Task pop():
24             Chunk*& cur = perThread.local();
25             // Try the local chunk first
26             if cur && !cur->empty():
27                 return cur->pop()
28             // Delete empty chunk
29             if cur:
30                 delete cur
31             // Next, try the per-package list
32             cur = perPackage.local().pop()
33             if cur:
34                 return cur->pop()
35             // If no task is found, probe other packages
36             for lst in perPackage:
37                 // Lock-free test for tasks
38                 if !lst.empty():
39                     cur = lst.pop() //synchronized pop
40                     if cur:
41                         return cur->pop()
42             return FAIL

```

Figure 6.2: Pseudocode for distributed chunked LIFO

(LIFO). New tasks are pushed onto the ring buffer, and tasks are popped from it when the thread needs work.

- Each package has a list of chunks. This list is manipulated in LIFO order.
- When the chunk associated with a thread becomes full, it is moved to the package-level list.
- When the chunk associated with a thread becomes empty, the thread probes its package-level list to obtain a chunk. If the package-level list is also empty, the thread probes the lists of other packages to find work. To reduce traffic on the inter-package connection network, only one hungry thread hunts for work in other packages on behalf of all hungry threads in a package.

This implementation is *topology-aware* because the most frequent operations use the least cost communication paths, while less frequent operations may use more costly communication. Threads can usually satisfy pushing or popping tasks by manipulating thread-private chunks. Only when a per-thread chunk becomes full or empty is the per-package linked-list manipulated. Communication between threads sharing the same package costs more than thread-private communication but costs less than communication between packages. Only in the rare case when (1) a thread is trying to pop a task, (2) its thread-private chunk is empty, and (3) its per-package linked-list is empty does a thread use expensive inter-package communication to find tasks in other packages.

Similar topology-aware optimizations have been proposed for per-thread dequeues (Guo et al., 2010; Wang et al., 2012b; Drebes et al., 2014), and the compositional scheduler synthesis described in Section 6.2 can use these schedulers, with slight modification (see Section 6.2.3), in addition to the topology-aware bag described here. However, most prior work in scheduling considers the problem of implementing a single, monolithic scheduler that is used for all parallel tasks and does not consider schedulers as blocks out of which more sophisticated schedulers are built. In the compositional scheme introduced later in this chap-

ter, the complete task scheduler may be composed of a basic scheduler like the distributed LIFO bag instantiated multiple times. In this context, it is important for the bags to support efficient queries for emptiness (i.e., the pop operation when returning failure should be fast) to allow the complete task scheduler to quickly dispatch operations among different bags. For p per-thread dequeues, checking for the absence of all tasks requires $O(p)$ work while the distributed chunked LIFO requires $O(\frac{p}{c})$ where c is the number of cores per package. When there is only one bag instance, this cost is negligible in practice, but when there are multiple bags and they are repeatedly queried, this cost can be significant.

6.1.2 Topology-Aware Priority Scheduler

The distributed chunked LIFO scheduler can be used as a component in a topology-aware priority scheduler.

Priority scheduling is used extensively in operating systems, but relatively simple implementations suffice in that context because tasks are relatively coarse-grained: operating system tasks may execute in tens or hundreds of milliseconds, whereas tasks in parallel programs may take only microseconds to execute. Therefore, the overheads of priority scheduling in the operating system context are masked by the execution time of tasks, which is not the case in many parallel programs, so solutions from the operating systems area cannot be used here. Another possibility is to use a concurrent priority queue like a lock-free skip-list (Shavit and Lotan, 2000). However, this has high overheads. These alternatives are described in more detail at the end of this chapter. This section describes a machine-topology-aware, physically distributed data structure called *obim* that exploits the fact that priorities are “soft,” so the scheduler is not required to follow them exactly.

Overview

Unlike the scheduler of Section 6.1.1 which implements an unordered collection of tasks, the *obim* scheduler uses a *sequence* of bags, where each bag is associated with one priority

level. To be concrete, this section assumes that each bag is implemented as the distributed chunked LIFO described above, but in practice, the scheduler synthesizer can use any bag implementation. Tasks in the same bag have identical priorities and can therefore be executed in any order; however, tasks in bags that are earlier in the sequence are scheduled preferentially over those in later bags. This is shown pictorially as the *Global Map* in [Figure 6.1b](#). This map is sparse since it contains bags only at entries 1, 3 and 7. Threads work on tasks in bag 1 first; only if a thread does not find a task in bag 1 does it look for work in the next bag (bag 3). If a thread creates a task with some priority and the corresponding bag is not there in the global map, the thread allocates a new bag, updates the global map, and inserts the task into that bag.

The global map is a central data structure that is read and written by all threads. To prevent it from becoming a bottleneck and to reduce coherence traffic, each thread maintains a software-controlled lazy cache of the global map, as shown in [Figure 6.1b](#). Each local map contains some portion of the global map that is known to that thread, but it is possible for a thread to update the global map without informing other threads.

The main challenge in the design of obim is getting threads to work on early priority work despite the distributed, lazy-cache design. This is accomplished as follows.

Implementation of global/local maps

The thread-local map is implemented by a sorted, dynamically resizable array of pairs. Looking up a priority in the thread-local map is done using a binary search. Threads also maintain a version number representing the last version of the global map they synchronized with. The global map is represented as a log-based structure which stores bag-priority pairs representing insert operations on the logical global map. Each logical insert operation updates the global version number.

Updating the global map When a thread cannot find a bag for a particular priority using only its local map, it must synchronize with the global map and possibly create a new

```

1  class Obim:
2      struct Data:                // Per-thread data
3          Bag* current            // Current bag
4          int curPriority         // Current priority
5          int lastVersion        // Last version synced
6          OrderedMap<int, Bag*> localMap // Copy of global state
7
8      PerThread<Data> perThread
9      ThreadSafeVector<Pair<int, Bag*>> masterLog
10     int masterVersion
11     Spinlock masterLock
12
13     // Add a task with a priority
14     void push(Task t, int priority):
15         Bag* b
16         Data& d = perThread.local()
17         // Check cache
18         if d.curPriority == priority && d.current:
19             b = tld.current
20         else:
21             b = updateLocalOrCreate(d, priority)
22         b->push(t)
23         // Update priority if necessary
24         if priority < tld.curPriority:
25             d.current = b
26             d.curPriority = priority
27
28     Task pop():
29         Data& d = perThread.local()
30         if d.current && (++d.popCount % scanPeriod) != 0:
31             Task t = tld.current->pop()
32             if t != FAIL:
33                 return t
34         replayLog(d) // Failed, update log
35         return scan(d)

```

Figure 6.3: Pseudocode for obim; auxiliary functions in [Figure 6.4](#)

```

1  class Obim:
2      // ...
3
4      // Search for earlier priority work
5      Task scan(Data& d):
6          int scanStart
7          if usingBackScanPrevention:
8              scanStart = min([x.curPriority for x in perThread.packageLocal()])
9          else:
10             scanStart = 0
11         for p in range(scanStart, d.localMap.maxKey + 1):
12             Bag* b = localMap.find(p)
13             if b:
14                 Task t = b->pop()
15                 if t != FAIL:
16                     d.curPriority = p
17                     d.current = b
18                     return t
19         return FAIL
20
21     // Find entry in local cache or create entry in global map
22     Bag* updateLocalOrCreate(Data& d, int priority):
23         while true:
24             replayLog(d)
25             if d.localMap.find(priority):
26                 return d.localMap.find(priority)
27             if masterLock.tryLock():
28                 break
29             // Serialization point
30             replayLog(d)
31             if d.localMap.find(priority):
32                 return d.localMap.find(priority)
33             // Update log
34             int v = masterVersion + 1
35             Bag* b = new Bag
36             masterLog[v] = Pair(priority, b)
37             memoryBarrier()
38             masterVersion = v
39             d.localMap.insert(priority, b)
40             masterLock.unlock()
41             return b
42
43     // Update local cache from log
44     void replayLog(Data& d):
45         int m = masterVersion
46         for i in range(d.lastVersion, m):
47             d.localMap.insert(masterLog[i])
48         d.lastVersion = m

```

Figure 6.4: Auxiliary functions for Figure 6.3

mapping there. This is accomplished in the *updateLocalOrCreate* method (see [Figure 6.4](#) line 22). A thread replays the global log from the point of the thread's last synchronized version to the end of the current global log. This inserts all newly created mappings into the thread's local map. If the right mapping is still not found, the thread will acquire a write lock, replay the log again, and append a new mapping to the global log and its local map. The write lock ensures that only one mapping exists for any priority value. Some care must be taken with the implementation of the global log to ensure that the log can be appended in the presence of concurrent readers without requiring locks.

Pushing a task A thread pushing a task uses its local map to find the correct bag into which to insert. Failing that, the thread updates its local map from the global map, as above, possibly creating a new mapping, and it uses the found or created bag for the push operation.

Retrieving a task To keep close to the ideal schedule, all threads must be working on important (earliest priority) work. When a task is executed, it may create one or more new tasks with earlier priority than itself because priorities are arbitrary application-specific functions. If so, the thread executes the task with the earliest priority and adds all the other tasks to the local map. Threads search for tasks with a different priority only when the bag in which they are working becomes empty or if a heuristically defined number of pops has occurred; the threads then scan the global map looking for important work. This procedure is called the *back scan*.

Because a scan over the entire global map can be expensive, especially if there are many bags, an approximate consensus heuristic, called *back scan prevention*, is used to locally estimate the earliest priority work available and to prevent redundant scans for earlier priority work. Each thread publishes the priority it is working at by writing it to shared memory. When a thread needs to scan for work, it looks at this value for all threads that share the same package and uses the earliest priority it finds to start the scan for work. To propagate information between packages, in addition to scanning all the threads in its

package, one leader thread per package will scan the other package leaders. This restriction allows most threads to incur only a small amount of local communication. Once a thread has a starting point for a scan, it simply tries to pop work from each bag from the scan point onwards.

Back scan prevention is effective because, in many uses of priority scheduling, tasks generate work at the same or later priority, and back scan prevention can limit the scan for earlier priority tasks to just a few bags.

This scheduler is topology-aware because push and pop operations are most likely satisfied by accessing thread-private chunks in a distributed chunked LIFO via operations on the bag at the current priority. The next most frequent operation is using the ordered map to find the bag into which to insert new work. The local map cache allows this operation to also be core-private with high probability. Finally, threads must from time to time find an earlier priority bag to retrieve work from. This requires some communication between threads to find the global minimum. Although this could be done on mostly read-only data, adding some communication in the form of a topology-aware, approximate, autonomous consensus algorithm greatly reduces the total time spent looking for the least bag in the system and reduces cases where a thread works on one priority even though earlier priority work has been enqueued (i.e., priority inversions).

Alternatively, one may use a more inductive argument to show that this scheduler is topology-aware. The push and pop operations mainly consist of finding an early priority bag and applying the appropriate operation to the bag. The bag is already known to be topology-aware, so one must only show that finding early priority bags is topology-aware.

Evaluation of Design Choices

This section evaluates the design choices of the obim scheduler by comparing the performance of several de-optimized variants. [Figure 6.5b](#) lists the variants, which focus on two main optimizations: (1) the use of per-package linked-list of chunks and (2) back scan

prevention. Recall that the distributed chunked LIFO uses per-package linked-lists to reduce inter-package communication. This optimization can be disabled by replacing these per-package linked-lists with a single lock-free linked-list shared by all threads. Back scan prevention can be disabled by always starting scans from the earliest priority.

The machines used for the evaluation are: (1) m2x4, a two processor, four cores per processor Intel system, (2) m4x10, a four processor, ten cores per processor Intel system, and (3) numa8x4, an eight processor, four cores per processor Intel system where every two processors are packaged into boards and boards are connected using a NUMA interconnect.

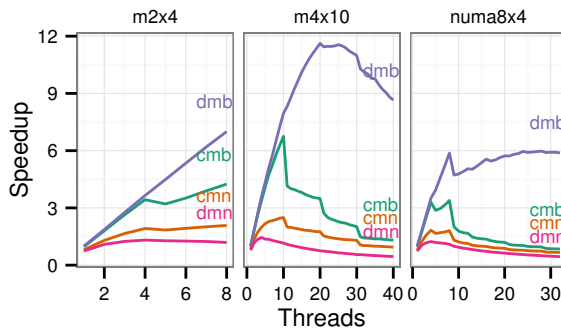
[Figure 6.5b](#) shows the speedup of SSSP relative to the best overall single-threaded execution time on a road graph of the United States (23.9 M nodes, 57.7 M edges). Back scan prevention is critical for performance; without this optimization (cmn and dmn), speedup is never more than 2.5 on any machine for any thread count, but with this optimization (cmb and dmb), speedup rises to about 12 on 20 threads on the m4x10 machine.

Using distributed bags is also important for performance: without this optimization, speedup is never more than 5 on any machine. It is interesting to note that without back scan prevention, a distributed bag is less efficient than a centralized one on this input. This is because it is more efficient to check that a (single) centralized bag is empty than it is to perform this check on a (per-package) distributed bag.

6.2 Compositional Scheduling Policies

[Section 2.3](#) suggests that many algorithms benefit from scheduling strategies that are more complex than simple strategies like LIFO and FIFO. This section shows how the informal policies described in [Section 2.3](#) can be encoded in a simple but flexible specification language.

Scheduling specifications in the Galois system are built from *ordering rules*, where an ordering rule specifies a total order on activities or items. Given two items a and b , an ordering rule R may specify that a should be processed before b (written as $a <_R b$) or



(a) Speedup of variants on road graph

	Back Scan Prevention	
	No	Yes
Centralized Bag	cmn	cmb
Distributed Bag	dmn	dmb (obim)

(b) Obim variants

Figure 6.5: Scaling of obim and its variants for SSSP application

vice versa ($b >_R a$), or it may leave the order unspecified ($a =_R b$). For example, if items have integer priorities, an ordering rule A_1 may order them in ascending priority order; the relative order of items with the same priority is left unspecified by A_1 . Ordering rules can be composed sequentially in a manner similar to lexicographic ordering: a sequence $D = R_1 R_2 R_3 \dots$ is itself an ordering rule that first orders items according to R_1 ; if two items are not strictly ordered by R_1 , they are ordered according to R_2 , etc. For example, if F_1 orders items in FIFO order, then $A_1 F_1$ denotes the order in which items are processed in increasing priority order and items with the same priority are processed in FIFO order. For the synthesis procedure described in Section 6.2.1, it is convenient to distinguish between *final rules*, which are rules that appear last in an ordering sequence, and *non-final rules*.

As discussed in Section 2.3, some implementations of irregular algorithms maintain separate global and thread-local worksets, so it is natural to use different ordering rules for them. A scheduling specification can have both a global ordering rule and a local ordering rule. The global rule is applied to the initial set of items, and the local rule is applied to each

thread-local workset, which holds items created dynamically by the corresponding thread. A thread accesses the global workset only when its local workset is empty. Continuing the previous example, if L_1 is last-in-first-out (LIFO) order, then global order A_1F_1 and local order L_1 specify that global items are processed as before, and local items are processed in LIFO order.

Figure 6.6 gives the syntax rules for scheduler specifications. \mathbf{T} is the type of items. Figure 6.7 gives the semantics. The meaning of an ordering rule R is given as a function over items a and b that is true iff $a <_R b$. The relation $<$ is the standard order on integers and reals.¹ For the FIFO and LIFO rules, we define an auxiliary function `time` that maps an item to an integer according to when the item is added to the scheduler. For the first item x_1 added to the scheduler, `time`(x_1) = 0; for the second item x_2 , `time`(x_2) = 1, and so on. f_U is an injective function mapping items to integers; this function essentially encodes a random permutation of items. The function f_D should be consistent with a total order.

Figure 6.8 shows the specifications of the scheduling policies discussed in Section 2.3. When a specification has an empty local ordering, the figure omits the global and local tags. The BRIO specification assumes that items have already been assigned rounds according to the random distribution described in Section 2.3. In the original delta-stepping algorithm (Meyer and Sanders, 1998), edges relaxations are divided into light (i.e., $< \Delta$) and heavy requests, and for a particular bucket, light requests are processed first, including any light requests enabled by processing a request, before moving on to the heavy requests. To process light requests before heavy ones, the delta-stepping specification splits each bucket into two: one part for the light requests and one part for the heavy requests.

¹ Instead of binary relation on pairs of items $\mathbf{T} \times \mathbf{T} \rightarrow \mathbf{bool}$, the semantics of a rule could be a function that maps a set of items to a sequence of sets of items $\mathbf{Set} \mathbf{T} \rightarrow \mathbf{Seq} \mathbf{Set} \mathbf{T}$. This allows a cleaner formal development (e.g., the non-syntactic a and b terms can be removed). We adopt the current approach because it seems more intuitive for programmers.

P	::= Global: D Local: D	<i>Specification</i>
D	::= R_{NF}^* $R_F^?$	<i>Ordering rule</i>
R_F	::= FIFO	<i>Final rule</i>
	LIFO	
	Random	
R_{NF}	::= ChunkedFIFO(k)	<i>Non-final rule</i>
	ChunkedLIFO(k)	
	Ordered(f_D)	
	OrderedByMetric(f_M)	
k		<i>Integer</i>
f_D		$\mathbf{T} \times \mathbf{T} \rightarrow \mathbf{bool}$
f_M		$\mathbf{T} \rightarrow \mathbb{R}$

Figure 6.6: Scheduling specification syntax

6.2.1 Synthesis

It is straightforward to implement sequential schedulers for policies specified in the language described in [Section 6.2](#). Each rule can be implemented by a *workset*, which is an object with the following methods:

- **void** push(Task t) — adds an item to the workset
- Task pop() — removes and returns the next item to execute; if there are no items left, returns a **FAIL** value distinct from all items added to the workset

Items are added to the workset by invoking the push method, which returns the value **void** when it completes. To get items from the workset, the pop method is invoked; if this method invocation does not find any items in the workset, it returns a unique **FAIL** value.

The goal of this section is to synthesize *concurrent* worksets that implement this functionality. One approach is to compose a set of library components, each of which is a workset by itself. There is a workset for each final rule; non-final rules are implemented by worksets parameterized by a function that constructs instances of the next workset in the ordering sequence, the *inner* workset.

$$\begin{aligned}
\llbracket \text{FIFO} \rrbracket(a, b) &= \mathbf{time}(a) < \mathbf{time}(b) \\
\llbracket \text{LIFO} \rrbracket(a, b) &= \mathbf{time}(a) > \mathbf{time}(b) \\
\llbracket \text{Random} \rrbracket(a, b) &= f_U(a) < f_U(b) \\
\llbracket \text{ChunkedFIFO}(k) \rrbracket(a, b) &= \lfloor \mathbf{time}(a)/k \rfloor < \lfloor \mathbf{time}(b)/k \rfloor \\
\llbracket \text{ChunkedLIFO}(k) \rrbracket(a, b) &= \lfloor \mathbf{time}(a)/k \rfloor > \lfloor \mathbf{time}(b)/k \rfloor \\
\llbracket \text{Ordered}(f_D) \rrbracket(a, b) &= f_D(a, b) \\
\llbracket \text{OrderedByMetric}(f_M) \rrbracket(a, b) &= f_M(a) < f_M(b) \\
\llbracket R_1 R_2 \dots \rrbracket(a, b) &= \begin{cases} \llbracket R_2 \dots \rrbracket(a, b) & \text{if } a =_{R_1} b \\ \llbracket R_1 \rrbracket(a, b) & \text{otherwise} \end{cases}
\end{aligned}$$

Figure 6.7: Scheduling rule semantics

However, a naive implementation along these lines can be incorrect in a concurrent setting, as will be seen in [Section 6.2.2](#), and the result may not satisfy any intuitive notion of correctness such as linearizability ([Herlihy and Wing, 1990](#)). To address this problem, [Section 6.2.3](#) proposes a relaxed correctness condition that requires modifications to the semantics of worksets and to how they are used by clients.

[Section 6.2.4](#) discusses two important consequences of this relaxed condition: (1) all final scheduling policy rules in [Section 6.2](#) have implementations that satisfy the relaxed condition, and (2) non-final worksets satisfy the relaxed condition assuming only that their inner worksets satisfy the relaxed condition. This permits compositional construction of worksets.

[Section 6.2.5](#) discusses a preliminary implementation of scheduling policies in Java in an early version of the Galois system and several optimizations to improve the performance of the synthesized worksets.

6.2.2 Problems with Naive Composition

To understand the issues that arise in composing worksets, consider the implementation of the bucket-based scheduler in lines 1–18 of [Figure 6.9](#). This is one possible implementation of the `OrderedByMetric` rule (the `obim` scheduler of [Section 6.1.2](#) is another). Lines 20–25

	Specification	Used by
DMR	OrderedByMetric($\lambda t. \text{minangle}(t)$) FIFO	Triangle angle (AS2) (Shewchuk, 1996)
	Global: ChunkedFIFO(k) Local: LIFO	Local stack (AS1) (Kulkarni et al., 2008)
DT	OrderedByMetric($\lambda p. \text{round}(p)$) ChunkedFIFO(k)	BRIO (AS1) (Amenta et al., 2003)
	Random	Random (Clarkson and Shor, 1989)
PFP	FIFO	FIFO (Goldberg and Tarjan, 1988)
	OrderedByMetric($\lambda n. - \text{height}(n)$) FIFO	HL order (AS1) (Cherkassy and Goldberg, 1995)
PTA	FIFO	LRF (Pearce et al., 2003)
	(empty)	Split worklists (BS-F) (Nielson et al., 1999; Hardekopf and Lin, 2007)
SSSP	FIFO	Bellman-Ford (Bellman, 1958; Ford and Fulkerson, 1962)
	OrderedByMetric($\lambda n. \lfloor 2 * w(n) / \Delta \rfloor + (\text{light}(n) ? 0 : 1)$) FIFO	Delta-stepping (AS1) (Meyer and Sanders, 1998)
	Ordered($\lambda a, b. w(a) \leq w(b)$)	Dijkstra (AS2) (Dijkstra, 1959)

Figure 6.8: Application-specific scheduling specifications

```

1  class BucketedScheduler:
2      Vector<Bag> buckets
3      int cursor
4
5      void push(Task t):
6          int index = floatToInt( $f_M(t)$ )
7          buckets[index].push(t)
8          if index < cursor: cursor = index
9
10     Task pop()
11         Task t = FAIL
12         while cursor < buckets.size():
13             t = buckets[cursor].pop()
14             if t == FAIL:
15                 cursor++
16             else:
17                 break
18         return t
19
20     BucketedScheduler scheduler
21     ThreadPool.fork(N) // spawn N threads
22     Task item
23     while (item = scheduler.pop()) != FAIL:
24         operator.call(item, scheduler)
25     ThreadPool.join(N)

```

Figure 6.9: Naive bucketed scheduler

show how this scheduler might be used in a parallel runtime system. The runtime system manages threads and assigns them work. The workset creates an array of inner worksets and processes each inner workset in ascending order. This workset is essentially an implementation of a priority queue in which the range of keys is known *a priori*; there is one inner workset (bucket) for each key value. Similar worksets have been used in a variety of sequential (Amenta et al., 2003; Cherkassy and Goldberg, 1995; Shewchuk, 1996) and parallel implementations (Meyer and Sanders, 1998) of irregular algorithms.

Unfortunately, the workset in Figure 6.9 can exhibit incorrect behavior because it is possible for items to be inserted into the workset but never retrieved. Consider two threads T_1 and T_2 , where T_1 is executing the pop method and T_2 is executing the push method. The following sequence of events may take place:

1. T_1 executes line 13. The cursor value is i , and the pop method on `buckets[i]` returns **FAIL**.
2. T_2 executes lines 6–8. The value of `index = i`, so an item is added to `buckets[i]`.
3. T_1 executes lines 14 and 15 of the pop method, incrementing cursor.

Clearly, the item added by T_2 is now lost. The race exists even if each line of the implementation is atomic, so that reading and updating the cursor on line 15 or incrementing its value on line 15 is performed atomically. To use this implementation correctly in a concurrent context, one must ensure that when poll moves to the next bucket, no thread is adding an element to an earlier priority bucket.

A second problem is that even if the methods of a workset are linearizable, the workset cannot be used to directly control the execution of threads in the client. In the client code of Figure 6.9, a thread stops processing items when the workset returns **FAIL**, and then waits for the rest of the threads to join it. However, one possible scenario is (1) all threads but one are waiting at the barrier, (2) the workset is empty, and (3) while processing the last item, the last thread adds several items to the workset. The waiting threads now need to wake up and re-enter the parallel loop, but this will not happen in the client code of Figure 6.9. Abstractly, the problem is one of termination detection: when should parallel execution stop? An eager termination detection algorithm risks having threads idle when there is work to be done. A lazy algorithm will wait needlessly when there is no work. A naive client of a concurrent workset will generally be too eager, so parallel runtime systems need to have separate mechanisms for termination detection.

These observations can be summarized as follows.

- In general, simple compositions of worksets do not produce correct concurrent worksets.
- Composition of concurrent worksets is problematic even in absence of having to maintain a particular order of items.

- Worksets are used by parallel runtime systems. In some cases, runtime systems themselves separately implement stronger properties like termination detection that overlap with some workset functionality.

6.2.3 Relaxed Concurrent Semantics

This section describes a solution to the problem of composing worksets that takes advantage of the fact that scheduling specifications for unordered algorithms are inherently “fuzzy” and are intended as suggestions to the runtime system rather than as commands that must be followed exactly. At a high level, the idea is the following.

- Relax the behavior of the pop method so that in a parallel setting, it may return a different item than the one it would have returned in a sequential setting. In addition, pop may return **FAIL** even when there are still items in the workset. These modifications permit us to implement pop with low overhead.
- To compensate for the relaxed behavior of pop, introduce a new method pop-s that is similar to pop but is never executed concurrently with other invocations. It returns **FAIL** only when the workset is truly empty.

Intuitively, if most items are retrieved from the workset using the pop method, and pop-s is used infrequently to determine if the workset is truly empty, the resulting scheduler is both correct and efficient. This section formalizes this behavior, and [Section 6.2.4](#) shows how this behavior is closed under composition.

A workset is modeled by its history H , which is a finite sequence of *events*, where an event is either (1) a method invocation, (2) a response to a method invocation, or (3) a special termination event, $\langle \text{term} \rangle$. The invocation on object o of method m with arguments a, b, \dots by thread T is written as $\langle o.m(a, b, \dots) T \rangle$; the response to method m with return value r is written as $\langle o.m(a, b, \dots)/r T \rangle$; The unit return value is written as **void**. An invocation $\langle o_1.m_1(a_1, b_1, \dots) T_1 \rangle$ *matches* a response $\langle o_2.m_2(a_2, b_2, \dots)/r T_2 \rangle$ if $o_1 =$

$o_2, m_1 = m_2, a_1 = a_2, b_1 = b_2$ and so on, and $T_1 = T_2$. Objects and/or threads are omitted if they are clear from context. Terms x_1, x_2, \dots are variables over arguments or return values.

An invocation is *pending* in H if it has no matching response. A history is *whole* if it has no pending invocations and it contains exactly one termination event and that is the last event in the history. A history *restricted to* object o or thread T is the subsequence with only events on o or by T respectively and possibly a termination event. Without loss of generality, items are assumed to be unique. The notation $a \rightarrow_H b$ denotes that event a precedes event b in history H ; when the history is clear from context, this is written $a \rightarrow b$.

Property 6.1 is a formal description of the behavior of pop and pop-s.

Property 6.1 (Weak Bag). *A history H models a weak bag if the following are true:*

- B1. There is an injective function M from non-FAIL response events $e_1 = \langle \text{pop}() / x_1 \ T_1 \rangle$ or $e_1 = \langle \text{pop-s}() / x_1 \ T_1 \rangle$ to invocation events $e_2 = \langle \text{push}(x_2) \ T_2 \rangle$ such that (1) $M(e_1) = e_2$, (2) $x_1 = x_2$, and (3) $e_2 \rightarrow e_1$.*
- B2. For each invocation event $e = \langle \text{pop-s}() \ T \rangle$ of pop-s in $H = H_1, e, H_2$, there are no pending invocations in H_1 .*
- B3. For each FAIL response event $e = \langle \text{pop-s}() / \text{null} \ T \rangle$ of pop-s in $H = H_1, e, H_2$, H_1 satisfies condition B1 and M is a bijective function.*

Condition B1 states that (1) items returned by pop and pop-s must have been added earlier by the push method and (2) a given item can only be returned once. Both requirements are captured by the injective function M . Condition B2 states that pop-s cannot be invoked when there are pending method invocations. Condition B3 states that if pop-s returns **FAIL**, all previously pushed items have been retrieved by pop or pop-s, and the workset is truly empty. This is captured by requiring M to be a bijective function.

A workset is *correct* if it only generates whole histories satisfying **Property 6.1**. It is the responsibility of the client to use the workset properly by never invoking pop-s

concurrently with other methods. This form of correctness may seem particularly weak since it does not refer to the sequential ordering semantics. However, we have found it useful because it includes many natural compositions of worksets as well as most hand-written schedulers. A linearizable bag is a correct workset if one considers pop-s the same as pop. Likewise, a bag with a single lock guarding all its methods is also a correct workset.²

One correct workset and its proper use by a runtime system is the following modification of Figure 6.9. Before line 25, the runtime system should call pop-s; if the returned value is a non-FAIL item, the system should process that item and go to line 23 to continue execution. The pop-s method should walk the bucket array calling poll-s on each inner workset and return the first non-FAIL item if it exists.

The implementations in the Galois system of the final rules in Figure 6.6 satisfy Property 6.1 since the LIFO and FIFO rules are implemented by a linearizable stack and queue respectively, and the Random rule is implemented with a resizable array and all method invocations are protected by a single lock.

6.2.4 Workset Composition

This section summarizes how the implementations of the non-final rules in Figure 6.6 satisfy Property 6.1, assuming they are parameterized by correct worksets, and gives a detailed description for one non-final rule, OrderedByMetric. The correctness of other worksets is briefly summarized.

Theorem 6.1 models the behavior of the OrderedByMetric workset by its history with respect to its inner worksets. Theorem 6.2 shows how objects that generate such histories are correct worksets.

Theorem 6.1 (OrderedByMetric). *Let o be an instance of the OrderedByMetric workset in Figure 6.9 modified so that each thread maintains a thread-local cursor variable and pop-s walks all the buckets calling pop-s on each inner workset. Let $W = \{w_1, w_2, \dots, w_n\}$ be*

²It is possible to introduce deadlock when arbitrarily composing worksets with locks. However, compositions based on Theorem 6.2 whose implementations are themselves wait-free do not introduce deadlocks.

the set of correct worksets contained in the buckets. The workset o only generates whole histories containing the following non-overlapping sequences when restricted to thread T for all threads:

D1. $\langle o.push(x) \rangle$,
 $\langle w.push(x) \rangle$, $\langle w.push(x)/\mathbf{void} \rangle$,
 $\langle o.push()/\mathbf{void} \rangle$.

D2. $\langle o.pop() \rangle$,
 $(\langle w*.pop() \rangle, \langle w*.pop()/\mathbf{FAIL} \rangle)^*$,
 $\langle w.pop() \rangle$, $\langle w.pop()/x \rangle$,
 $\langle o.pop()/x \rangle$ where $x \neq \mathbf{FAIL}$.

D3. The above with $pop()$ replaced with $pop-s()$.

D4. $\langle o.pop() \rangle$,
 $(\langle w*.pop() \rangle, \langle w*.pop()/\mathbf{FAIL} \rangle)^*$,
 $\langle o.pop()/\mathbf{FAIL} \rangle$.

D5. $\langle o.pop-s() \rangle$,
 $\langle w_1.pop-s() \rangle$, $\langle w_1.pop-s()/\mathbf{FAIL} \rangle$, \dots ,
 $\langle w_n.pop-s() \rangle$, $\langle w_n.pop-s()/\mathbf{FAIL} \rangle$,
 $\langle o.pop-s()/\mathbf{FAIL} \rangle$.

Proof. Note that the only objects shared between threads executing methods of o are the inner worksets in W , represented in [Figure 6.9](#) as the variable buckets. Thus, it is sufficient to only consider sequential executions of o .

The push method clearly satisfies clause D1.

The pop method may either return **FAIL** or a non-**FAIL** value. In the case of **FAIL**, each inner workset visited returns **FAIL**, and o satisfies clause D4. In the case of a non-**FAIL** value, there are some number of inner worksets that return **FAIL** and exactly one that returns a non-**FAIL** value. This satisfies clause D2.

The pop-s method is the same as the pop method except that it visits all the inner worksets in W . By reasoning similar to the pop method, the pop-s method must satisfy clauses D3 or D5. \square

Theorem 6.2 (OrderedByMetric is correct). *Whole histories H satisfying [Theorem 6.1](#) model a weak bag.*

Proof. Consider each condition of [Property 6.1](#) in turn.

First, H satisfies condition B1. Without loss of generality, consider events of pop; events of pop-s behave similarly. From clauses D2 and D4, the only time the workset o produces a non-FAIL response $e_1 = \langle o.\text{pop}() / x_1 \rangle$ is when one of its correct worksets $w \in W$ produces a response $f_1 = \langle w.\text{pop}() / x_1 \rangle$. There is exactly one event e_1 for each event f_1 and vice versa. Let M_1 be the bijective function from events of the form e_1 to events of the form f_1 . From clause D1, there is exactly one event $f_2 = \langle w.\text{push}(x_2) \rangle$ for each event $e_2 = \langle o.\text{push}(x_2) \rangle$ and vice versa. Let M_2 be the bijective function from events of the form f_2 to events of the form e_2 .

We now show there exists an injective function M_C such that $M_C(e_1) = e_2$, $x_1 = x_2$ and $e_1 \rightarrow e_2$. By definition, $M_1(e_1) = f_1 = \langle w.\text{pop}() / x_1 \rangle$. Since w is a correct workset, there exists an injective function M_w such that $M_w(f_1) = f_2 = \langle w.\text{push}(x_1) \rangle$ and $f_2 \rightarrow f_1$. Let M be the expansion of M_w over the range of all $w \in W$. The ranges of M_{w_1}, \dots, M_{w_n} are disjoint and each M_w is an injective function so M is also an injective function (see [Figure 6.10](#)). By definition, $M_2(f_2) = e_2 = \langle o.\text{push}(x_1) \rangle$. Let M_C be the composed function $M_2 \circ M \circ M_1$. M_C is injective because M is injective and M_1 and M_2 are bijective. $M_C(e_1) = e_2$ by function composition. From clauses D1 and D2, $e_2 \rightarrow f_2$ and $f_1 \rightarrow e_1$, and by the correctness of w , $f_2 \rightarrow f_1$; so, by transitivity, $e_2 \rightarrow e_1$.

Second, H satisfies condition B2. Clients of o do not invoke $o.\text{pop-s}()$ concurrently. From clauses D3 and D5, it is clear that if there are no pending invocations immediately before $o.\text{pop-s}()$ is invoked, then there will be no pending invocations immediately before $w.\text{pop-s}()$ for all $w \in W$.

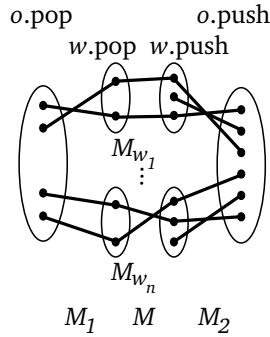


Figure 6.10: Relationship between M_1 , M and M_2 in the proof of [Theorem 6.2](#)

Finally, H satisfies condition B3. From above, M_C satisfies condition B1 and is an injective function from non-**FAIL** pop and pop-s responses to push invocations on o . We now show that when o produces the **FAIL** response $\langle o.pop-s()/\mathbf{FAIL} \rangle$, M_C is a bijection as well. From clause D5, if o produces a **FAIL** response, all $w \in W$ have produced a **FAIL** response as well. Thus, M is a bijection, and correspondingly, M_C is one as well. \square

The ChunkedFIFO rule is implemented with a single linearizable (global) queue whose elements (chunks) are instances of the inner workset. Each inner workset contains at most k items. Each thread maintains a thread-local chunk to pop from. When the chunk is empty, the thread pops from the global queue for the next chunk. Each thread also maintains a thread-local chunk to push to. When the chunk is full, the thread pushes it to the global queue and creates a new empty chunk to push to. The pop-s method walks each thread-local chunk and the global queue calling pop-s on each.

[Theorem 6.2](#) does not immediately apply because chunks are created and discarded dynamically. One modification would be to keep track of all the chunks ever created. However, one observation is that chunks are accessed by at most one thread at a time, and they are discarded when they are empty, which can be determined by invoking pop-s on the chunk. Empty chunks will never contain any more items. Thus, discarded chunks do not affect the eventual correctness of the workset. Only chunks that may contain items matter, which are precisely those traversed by pop-s.

The ChunkedLIFO rule is implemented similarly to ChunkedFIFO except with a linearizable stack.

The Ordered rule is implemented with a concurrent heap with additional locks to protect the inner worksets. Showing the correctness of its implementation is beyond the scope of this chapter because it uses commutativity conditions to reduce the granularity of the locking (Kulkarni et al., 2011).

Although not a rule *per se*, the composition of a global and local rule, used to implement global and local orders in specifications, is implemented with worksets as well. There is one workset that implements the global rule and thread-local worksets that implement the local rule. Initial work is pushed to the global workset, while newly created work is pushed to the thread-local workset. New work is retrieved from the thread-local workset, if possible, and from the global workset otherwise. For proving correctness, this implementation can be viewed as a refinement of a chunked workset where chunks are never discarded but are instead refilled from the global workset.

6.2.5 Preliminary Implementation and Evaluation

This section describes a preliminary version of scheduler synthesis built on top of an early version of the Galois system (version 2.0.1), which used the Java programming language. Since that version, the Galois system (version 2.1.0 and above) has been written in C++. The new versions use the same scheduling techniques introduced in this section, but the performance results are not directly comparable between the versions.

Implementation

The specification language is a library-based domain-specific language. Each rule is represented by a Java class that implements the corresponding workset. Figure 6.11 gives an example. Figure 6.12 gives the general form. In the newer versions of the Galois system, there are some syntactic changes to accommodate differences between the Java and C++

```

1 Lambda<T,Integer> indexer = new Lambda<T,Integer>() {
2     public Integer call(T item) {
3         return item.height;
4     }
5 }
6 Priority.first(OrderByMetric.class, indexer)
7     .then(FIFO.class)

```

Figure 6.11: Concrete syntax of HL order (AS1) scheduling policy for PFP application

```

1 Priority.first( $G_1$ .class, args).then(...)
2     .thenLocally( $L_1$ .class, args).then(...)

```

Figure 6.12: Concrete syntax of Global: $G_1 \dots$ Local: $L_1 \dots$

languages. In either case, the sequence of method calls produces an AST that is passed to the workset synthesizer.

Based on the semantics of scheduling rules, the synthesizer can choose the following optimized workset implementations.

- **Use Serial:** As mentioned in [Section 6.2.4](#), the inner worksets used by ChunkedFIFO and ChunkedLIFO are thread-local. The worksets generated from the local part of a specification are also thread-local. Thread-local worksets can be implemented with non-concurrent data structures that are typically more efficient than concurrent ones.
- **Ignore Size:** In certain cases, worksets require inner worksets to maintain an estimate of the number of items they contain. The chunked worksets use this to keep track of when a chunk is full. The Ordered workset uses these sizes to implement commutativity conditions. This overhead may be significant in concurrent worksets because keeping track of sizes may require atomic increments. When sizes are not needed, the size metadata and effort maintaining it may be removed.
- **Use Bounded:** When a chunked workset is used, each inner workset can be no larger than the chunk size. The inner worksets can be optimized for a bounded size rather than using dynamically sized data structures.

Ignore Size	Use Serial	Use Bounded	$t = 1$	$t = 8$
+	+	+	0.0	0.0
-	+	+	0.8	12.1
+	-	+	2.4	5.5
-	-	+	7.8	7.7
+	+	-	3.6	3.5
-	+	-	11.3	11.5
+	-	-	5.0	16.8
-	-	-	2.9	17.5

Figure 6.13: Relative difference in percent (%) of the runtime of PFP application with BASE scheduler on m4x4 machine varying synthesizer optimizations (+: on, -: off) relative to all optimizations on for one and eight threads

The synthesizer applies rewrite rules over the AST to detect the above cases and selects, if possible, implementations that are non-concurrent, do not keep track of their size or are bounded.

To implement these optimizations, the synthesizer introduces new rules (classes) to represent serial implementations of all the rules and bounded size implementations of the FIFO, LIFO and Random rules. It also adds an ignore size parameter to each rule, which determines if the implementation keeps track of its size. Then, the synthesizer traverses the AST (1) rewriting any rule after a chunked rule to use its serial implementation, (2) rewriting only rules after an Ordered rule to keep track of their sizes, and (3) rewriting rules after a chunked rule to use bounded implementations if possible, using the chunk size as the bound.

Figure 6.13 summarizes the impact of these optimizations for a preflow-push (PFP) application and a synthesized scheduler called BASE (described in more detail in the following section). Positive numbers indicate how much slower a combination is relative to all optimizations on. The PFP application was chosen because all the optimizations described above can be applied and the amount of work done per workset item is small, which increases the relative impact of an efficient workset implementation. From the figure, it is

DMR	Triangle mesh of 550,000 triangles of which 261,100 are initially bad
DT	1,733,360 points generated from edge detection of a photograph
PFP	A flow network of 526,904 vertices arranged in 14 consecutive 194x194 frames with uniformly random capacities
PTA	Analysis of the gimp program
SSSP	Road network of western USA, weights are distances between locations: 6,262,104 vertices, 15,119,284 edges

Figure 6.14: Datasets used in scheduler synthesis evaluation

clear that these optimizations on worksets have a significant and mostly beneficial impact on single-threaded and multi-threaded performance.

Evaluation

This section evaluates the scheduler synthesizer on a suite of applications described in [Section 2.3](#). [Figure 6.14](#) shows the datasets used for each application. Each application was run with the following set of schedulers.

- **BASE**: This is the default scheduler used by the Galois system. It is a synthesized ChunkedFIFO with a chunk-size of 32.³ Each chunk is a thread-local LIFO.
- **FIFO, LIFO, RAND**: These schedulers are synthesized from the final rules FIFO, LIFO and Random. Application-specific schedulers typically use one of these schedulers as their lowest-level (final) scheduler.
- **WS-L, WS-F**: These schedulers are work-stealing with local LIFOs and FIFOs respectively. These schedulers were ported directly from the Fork-Join implementation in JSR166 and appear in the Java JDK 7. WS-L is widely used in many parallel systems.

³In the more recent C++ version of Galois, the default scheduler is a *distributed* chunked FIFO (see [Section 6.1.1](#)).

- **BS-L, BS-F:** These schedulers use a bulk-synchronous strategy with global LIFOs and FIFOs respectively. A barrier is used to safely swap between queues concurrently.
- **AS1, AS2:** These schedulers are synthesized from the application-specific specifications in [Figure 6.8](#).

The evaluation uses three machines.

- **m2x4:** a Sun Fire X2270 machine running Ubuntu Linux 8.04.4 LTS 64-bit. It contains two 4-core 2.93 GHz Intel Xeon X5570 (Nehalem) processors. The two CPUs share 24 GB of main memory. Each core has a 32 KB L1 cache and a unified 256 KB L2 cache. Each processor has an 8 MB L3 cache that is shared among the cores.
- **m4x4:** a machine also running Ubuntu Linux 8.04.4 LTS 64-bit. It contains four 4-core 2.7 GHz AMD Opteron 8384 (Shanghai) processors. Each core has a 64 KB L1 cache and a 512 KB L2 cache. Each processor has a 6 MB L3 cache that is shared among the cores.
- **uma4x8:** a Sun T5440 machine running SunOS 5.10. It contains four 8-core 1.4 GHz Sun UltraSPARC T2 Plus (Niagara 2) processors. Each processor has a 4 MB L2 cache that is shared among the cores.

The Sun JDK v1.6.0_21 was used to compile and run the programs with a heap size of 20 GB. To control for JIT compilation, each application was run four times within the same JVM instance and only the last run is reported.

The Galois system uses speculative parallelization (see [Section 2.2.1](#)), which introduces overheads from (1) using concurrent implementations of data structures and schedulers rather than their sequential counterparts, (2) acquiring locks to guarantee disjointness of neighborhoods, and (3) recording undo actions to implement rollback. The *serial* version of an application uses sequential data structures only and does not acquire locks or perform undo actions. The difference in performance between the serial version and the parallel,

	BASE	RAND	LIFO	FIFO	WS-L	WS-F	BS-L	BS-F	AS2	AS1
m2x4										
DMR	12.88	14.80	11.45	13.09	11.51	13.27	12.76	13.17	15.56	11.62
DT	25.04					25.42				14.78
PFP	110.93	109.77	169.86	115.40	173.47	116.44	110.18	118.59		45.94
PTA	13.87	-	-	12.58	-	12.74	20.26	12.84		
SSSP	-	-	-	-	-	-	-	-	7.66	4.96
m4x4										
DMR	16.29	19.52	13.55	16.76	13.74	16.76	16.25	16.71	19.59	13.64
DT	43.40					43.55				27.86
PFP	237.04	210.57	320.24	237.17	314.53	234.13	216.50	217.67		74.26
PTA	19.99	-	-	18.80	-	18.79	26.44	18.82		
SSSP	-	-	-	-	-	-	-	-	11.08	9.53
uma4x8										
DMR	61.76	68.10	54.79	63.51	53.84	63.31	62.86	64.17	77.81	60.33
DT	178.21					179.00				149.42
PFP	787.05	734.27	1264.61	741.01	1297.71	775.04	720.20	827.07		342.41
PTA	59.17	-	-	57.73	-	57.30	76.16	56.99		
SSSP	-	-	-	-	-	-	-	-	33.84	23.35

Figure 6.15: Runtimes of serial versions in seconds. In bold are the best serial times, the basis for the speedup numbers in Figure 6.16. Entries with - timed out. Blank entries indicate invalid or redundant combinations.

one-threaded version is the overhead of enabling speculative execution but never using it. This overhead can be significant for applications with short activities like PFP and SSSP.

Figure 6.15 shows the runtimes for serial applications. Figure 6.16 shows the parallel speedup over the best performing serial scheduler (shown in bold in Figure 6.15). The runtimes for PTA exclude time to read input, perform offline-cycle detection and write results. This differs from the methodology of Méndez-Lojo et al., which includes the time to perform offline-cycle detection (Méndez-Lojo et al., 2010). For all other applications, runtimes exclude time to read input data or write results but may include sections of the application that are not parallelized. This portion of time is usually negligible, but for DT with the AS1 scheduler, this includes time to construct the oct-tree.

	BASE	RAND	LIFO	FIFO	WS-L	WS-F	BS-L	BS-F	AS2	AS1
m2x4 ($t \leq 8$)										
DMR	5.70	4.82	0.95	3.81	4.35	5.13	2.64	3.53	2.01	6.15
DT	2.21					2.09				2.35
PFP	1.30	0.71	0.20	1.15	0.72	2.30	0.37	0.89		3.35
PTA	2.83	-	-	3.53	-	2.05	2.37	3.77		
SSSP	-	-	-	-	-	-	-	-	0.61	3.16
m4x4 ($t \leq 16$)										
DMR	7.85	3.43	0.95	3.74	6.94	7.53	1.91	3.83	2.32	10.45
DT	2.64					2.65				2.53
PFP	1.28	0.62	0.20	1.00	0.65	2.19	0.37	0.74		2.56
PTA	3.69	-	-	3.63	-	3.08	3.25	5.03		
SSSP	-	-	-	-	-	-	-	-	0.80	3.04
uma4x8 ($t \leq 32$)										
DMR	18.77	5.95	0.89	6.81	11.47	18.53	3.60	5.89	3.59	21.53
DT	5.43					5.48				3.29
PFP	2.30	1.25	0.32	2.84	2.18	4.46	0.80	2.13		5.92
PTA	4.20	-	-	4.49	-	5.42	4.62	6.16		
SSSP	-	-	-	-	-	-	-	-	0.50	2.33

Figure 6.16: Speedup over serial versions. In bold are the best speedups for each (application, machine) pair. Entries with - timed out. Blank entries indicate invalid or redundant combinations.

Empty entries indicate combinations of schedulers and applications that would either be redundant or perform significantly worse (by an order of magnitude or more) than the best serial version. For DT, the performance without randomizing the initial points is much worse than with randomization. Since the application does not create any new tasks, the BASE and WS-L runs include an initial timed phase that randomizes the input points. The other non-random schedulers are omitted because they perform similarly after including this phase. For PTA, the RAND, LIFO and WS-L schedulers timed out. For SSSP, only the AS1 and AS2 schedulers are competitive.

An important result is that the best scheduler for serial execution tends to be the best for parallel execution as well. This supports the use case where users experiment with

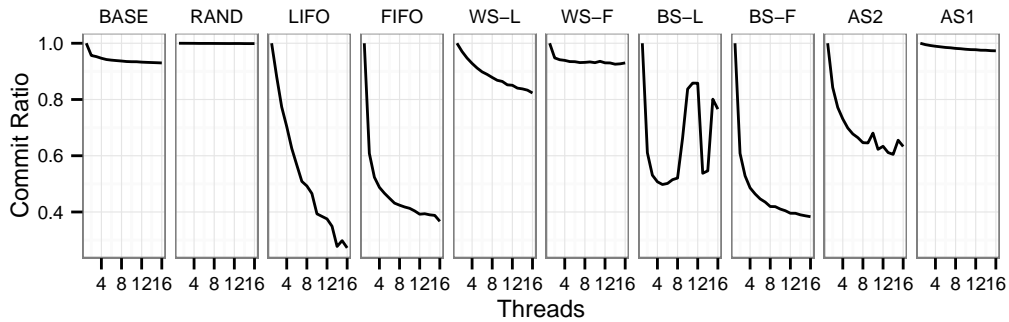


Figure 6.17: Relative number of committed to total iterations for DMR on m4x4

scheduling specifications within a sequential programming model and rely on a synthesis routine to generate efficient, concurrent schedulers that faithfully maintain the desired scheduling. Also, note the large swings between best and worst schedulers and recall that the missing entries for PFP, PTA and SSSP correspond to combinations that perform significantly worse than the recorded times. Choosing the wrong scheduler can have a drastic impact on performance.

Overall, the AS1 schedulers perform as well or better than any fixed-function scheduler for the same application. Importantly, the BASE scheduler, which is implemented by a global queue with fixed-size local stacks, performs relatively well across applications. The DMR application benefits from LIFO policies, while PFP, PTA and SSSP benefit from FIFO policies.

While direct comparison is not possible, these results are similar to previously reported results of application-specific schedulers on similar inputs for PFP (Bader and Sachdeva, 2005), PTA (Méndez-Lojo et al., 2010) and SSSP (Madduri et al., 2006).

Now, let us turn to each application in more detail.

Delaunay mesh refinement Figure 6.15 shows that for the serial implementation, the best performance is obtained by the LIFO scheduler. When a bad triangle is fixed, it may create a set of new bad triangles whose cavities overlap with the cavity of the original bad

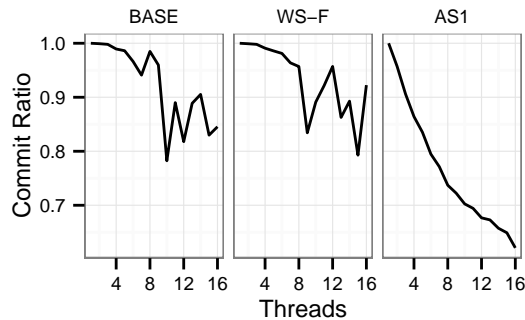


Figure 6.18: Relative number of committed to total iterations for DT on m4x4

triangle. The LIFO scheduler exploits this potential temporal and spatial locality. However, both of the global LIFO scheduling policies, LIFO and BS-L, perform poorly in a parallel setting because the probability of conflicts increases if triangles close to each other in the mesh are processed speculatively in parallel. Figure 6.17 shows the relative number of committed to total iterations on m4x4. The trends are similar for the other two machines. Not surprisingly, global FIFO schedulers behave the same way. The BASE scheduler, which uses a global FIFO, ameliorates this problem to some extent by distributing chunks of work to each thread.

The best performance is obtained with the AS1 scheduler, which is implemented by a global workset processed in chunked FIFO order and local worksets maintained in LIFO order. This enables exploitation of locality while controlling the commit ratio, as can be seen in Figure 6.17.

Delaunay triangulation This application does not create any new work. Its performance is governed by the initial work order, which should be randomized for best algorithmic performance. With randomization, the BASE and WS-L schedulers perform similarly. AS1 is significantly faster than the other two schedulers serially, but the performance difference dissipates as the number of threads increases. Recall that the AS1 scheduler is designed to increase spatial locality between activities. In speculative parallel execution, this scheduling

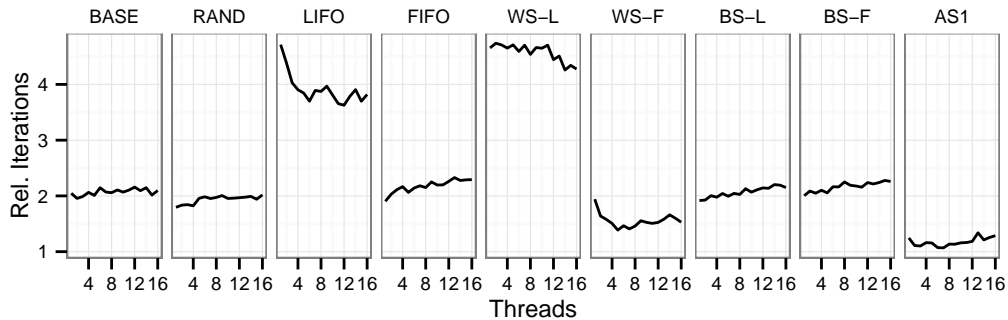


Figure 6.19: Relative number of committed iterations to the best performing serial version for PFP on m4x4

strategy causes the commit ratio to decrease, as can be seen in [Figure 6.18](#).

Preflow-push This application is an example of an algorithm whose performance is highly schedule-dependent because different schedulers result in dramatically different amounts of work. [Figure 6.19](#) shows the number of iterations committed relative to the best performing serial version on m4x4. Most schedulers result in twice as many iterations as the best serial version. LIFO and WS-L perform four times more iterations in some cases. [Figure 6.15](#) shows the impact of the varying work on the serial versions. The runtime of the fastest and slowest schedulers differ by a factor of more than three.

For a hand-parallelized implementation of PFP using the heuristics described in [Section 2.3](#), Bader and Sachdeva reported a maximum speedup of about 2 with eight processors on an UltraSPARC II architecture with an input similar to that used here, which is referred to as an RMF graph ([Bader and Sachdeva, 2005](#)).

Inclusion-based points-to analysis This application is another example of a highly schedule-dependent algorithm. The PTA application performs a fixpoint computation, and for these computations, FIFO policies usually perform well because a variable gets to accumulate several updates before its value is propagated down-stream. LIFO policies perform

poorly, and most versions time-out. BS-F, which alternates between two queues, does the best on this input. However, on other inputs, not shown here, BASE outperforms BS-F.

As noted earlier, these results are based on the implementation of Méndez-Lojo et al., the first parallel implementation of PTA (Méndez-Lojo et al., 2010).

Single-source shortest-Path This application is a case where the generally accepted best serial scheduler, which is AS2 and is based on Dijkstra’s algorithm, does not perform well in parallel. It has better theoretical algorithmic complexity, but the concurrent priority queue limits performance on most inputs. The difficulty in implementing such queues is one motivation for the delta-stepping order (AS1), which seeks to balance good order with efficient concurrent implementation. From experiments not shown here, it can be shown that the delta-stepping order performs only 1.2 times more work than Dijkstra’s algorithm for this input, an overhead that is modest enough to overcome through parallelism.

Madduri et al. produced an implementation of delta-stepping for the Cray MTA-2 architecture (Madduri et al., 2006). On the same input used here, a road network of the western USA, they reported a maximum speedup of about 2 on sixteen processors.

6.3 Exploiting Data Locality

Locality is exploited in the Galois system by scheduling activities according to the physical layout of data structures.

Although Galois data structures present a single logical view of data, they are often physically partitioned by thread or by package or by NUMA node. For instance, the morph graph described in Section 5.5 has nodes partitioned by thread. Users are not exposed to this partitioning directly, but through a scheduling policy, they can request that active nodes be initially assigned to the thread that owns the corresponding data. This increases the locality when the neighborhood of an operator is just the active node, which is common in data-parallel loops. Since partitions may be unbalanced, once a thread runs out of active

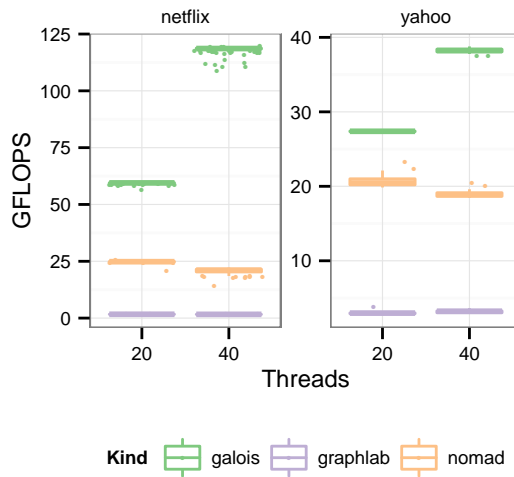


Figure 6.20: Throughput of matrix completion operators on machine m4x10 (theoretical peak throughput: 363.2 GFLOPS)

nodes that it owns the data for, it uses workstealing to find new work corresponding to other threads' partitions. In this way, locality can be exploited but full thread utilization is preferred over working on non-local data.

When neighborhoods consist of multiple nodes of a partitioned data structure, it is not obvious in general how to assign active nodes to threads. The owner-computes rule is one such strategy (Rogers and Pingali, 1989). For maximum effectiveness, the neighborhood of an operator should be contained to a single partition. Space-filling curves can be applied for geometric graphs, and graph partitioning algorithms (Karypis and Kumar, 1997) can be applied to arbitrary sparse graphs to increase the likelihood that a node's neighbors are assigned to the same partition, but general graph partitioning algorithms can be expensive relative to simple graph analytics algorithms. The well-known Metis graph partitioner can be expressed and parallelized in the Galois system (Sui et al., 2011), which raises the possibility of exploiting locality for more heavyweight algorithms. Recently more lightweight matrix reordering algorithms have been implemented in Galois (Karantasis

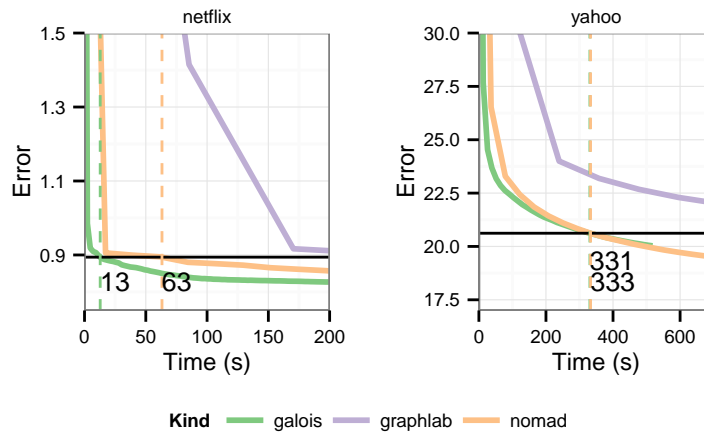


Figure 6.21: Convergence of matrix completion with 20 threads on machine m4x10. Horizontal line indicates error threshold of 90% of minimum observed error. Numbers indicate time to reach threshold.

et al., 2014), which also can be used to increase the likelihood that the neighborhood of an activity is localized to a small region of the physical data structure. Since the Galois system provides parallel data structures and scheduling, integrating these reordering and partitioning algorithms should not require significant changes in user programs, but a comprehensive solution has not yet been implemented.

Some locality optimizations can be framed as different scheduling policies. As an example, consider the matrix completion problem described in Section 2.3. When using SGD, the operator is over an edge and its two endpoints in a bipartite graph. In practice, the data associated with each endpoint is a reasonably sized vector ($k \approx 100$), which means that a high locality schedule is one that assigns edges to threads in such a way that the edges assigned to a thread share as many endpoints as possible. In this way, activities on edges reuse node data as much as possible. If the graph were complete, this schedule would resemble classical tiling for matrix-matrix multiply. However, in this case, the graph is sparse and the operator writes to its endpoints, which means that activities that share endpoints cannot be scheduled concurrently.

That being said, this sparse tiling is still a useful optimization. [Figure 6.20](#) shows the computational throughput of SGD for matrix completion on two inputs, netflix (17 K items, 480 K users, 99 M ratings) and yahoo (624 K items, 1 M users, 253 M ratings). The label Galois refers to an application in Galois using sparse tiling, which is simply a scheduling policy applied to the basic SGD algorithm; GraphLab refers to an application written in the GraphLab parallel programming model and does not do any locality optimizations; Nomad is a hand-written parallel code using MPI and pthreads ([Yun et al., 2014](#)). It tries to reduce communication (increase locality) by cyclically scheduling block diagonals of the matrix representation of the bipartite graph. The results show that even the simple block diagonal tiling done by Nomad is a significant improvement over no tiling at all and that the sparse tiling implemented in Galois can provide further performance improvements.

[Figure 6.21](#) shows the impact of the improved throughput on the end-to-end computation which runs until a certain error level is reached or for a certain amount of time. In this case, there are additional randomization steps taken by the Nomad implementation that reduce the initial throughput advantage of the Galois implementation for the yahoo input as time goes on, which is why 50% throughput improvement does not translate into a 50% reduction in overall time.

6.4 Coordinated Scheduling

The discussion so far has focused on autonomous scheduling (see [Section 2.2.3](#)), where activities are scheduled without global communication. Coordinated scheduling, where activities are executed in bulk-synchronous rounds, is also possible in the Galois system. An accumulating collection is a collection of elements that supports concurrent insertion of new elements but does not need to support concurrent reads of the collection. Coordinated scheduling policies can be built from multiple autonomously scheduled loops and an accumulating collection data structure, which is provided by the Galois library. For example, loop 1 executes, populating a collection with work that should be done by loop 2. Then,

loop 1 finishes, and loop 2 iterates over the collection generated by loop 1, and so on. Control logic can be placed between loops, allowing the expression of sophisticated coordinated strategies like the pull versus push operators required for breadth-first search (see [Section 2.3](#)).

6.5 Related Work

Although the Galois system synthesizes schedulers from specifications, this is not the standard concurrent program synthesis problem considered in the literature ([Vechev et al., 2007](#); [Solar-Lezama et al., 2008](#)). Scheduling specifications for unordered algorithms are inherently “fuzzy” (even FIFO scheduling of the workset can result in different executions depending on the speed of processors). Therefore, a scheduling specification is advice to the runtime system about how to bias scheduling decisions for efficiency, whereas in conventional program synthesis, implementations must satisfy specifications exactly.

Some work has shown that more efficient workset implementations may be possible if the application is written so that it can tolerate duplicate work items ([Michael et al., 2009](#); [Leijen et al., 2009](#)). Exploiting this idempotence property would require changes to the definitions in [Section 6.2](#) and [Section 6.2.1](#), but the implementation would be straightforward for final worksets. However, some non-final worksets, such as the chunked worksets, depend on the correspondence between adds and polls, which makes taking advantage of idempotence a challenge in these cases.

Previously, Kulkarni et al. ([Kulkarni et al., 2008](#)) explored different handwritten scheduling policies in the Galois system. However, the policies were a small number of fixed forms. In contrast, the schedulers described here are synthesized (not handwritten), they include policies studied in the literature such as delta-stepping, and they can be composed arbitrarily, which is important in practice.

There are several concurrent priority queues implementations ([Hunt et al., 1996](#); [Shavit and Lotan, 2000](#); [Sundell and Tsigas, 2005](#); [Bronson et al., 2010](#)). Preliminary

studies using concurrent skip-lists (Shavit and Lotan, 2000), a common implementation, revealed poor performance in the Galois system. Improved performance can be achieved by using bounded priorities (Shavit and Zemach, 1999), but the basic problem of lack of scalability remains. Chazelle investigated approximate priorities (Chazelle, 2000) but only considered sequential implementations.

Another possibility is to use a concurrent priority queue for each thread with work-stealing from other priority queues if the local priority queue becomes empty. Variations of this idea have been used previously in the literature (Papaefthymiou and Rodrigue, 1994; Bertsekas et al., 1996), and it is also used in GraphLab (Low et al., 2010). However, the work efficiency of the resulting implementation is often poor because early priority work generated by one thread does not diffuse quickly enough to other threads.

Yet another possibility is to use a concurrent priority queue for each thread, with logically partitioned graphs and the owner-computes rule (Rogers and Pingali, 1989) for task assignment. When task a creates a new task b , it looks at the partition assigned to task b to determine which priority queue to push task b on. This policy is used in GraphLab and other systems (Tang et al., 2008; Pearce et al., 2010). This policy is well-suited for distributed systems and has been used in distributed graph traversal algorithms (Pearce et al., 2010) but will perform poorly when work is localized to a subset of partitions.

Chapter 7

Deterministic Scheduling

In the optimistic scheduling described in [Chapter 6](#), when there is a conflict between two tasks¹, there is no particular order in which those tasks are executed, which can result in non-deterministic program execution. Non-deterministic scheduling is sufficient for many applications, but it is also possible to use a deterministic scheduler to guarantee that task conflicts are resolved in a reproducible way if desired.

The way non-determinism appears in the operator formulation is as follows:

- P is a pool of tasks that can be performed in *any* order, i.e., an unordered algorithm. The program terminates when all tasks have been executed.
- When task t is completed, it may create a set of new tasks $S(t)$, which are added to the task pool. Task t is the *parent* of the tasks it creates; the transitive closure of the parent relation is called the *ancestor* relation.
- Each task performs computation and reads and writes shared-memory locations. The set of locations read $R(t)$ and written $W(t)$ by a task t is said to constitute its *neighborhood*, which is denoted by $L(t) = R(t) \cup W(t)$. Tasks are required to be *cautious*:

This chapter draws from ([Nguyen et al., 2014](#)), where deterministic scheduling for the Galois system was originally described.

¹This chapter uses task interchangeably for activity to emphasize the applicability of these techniques for all programs including those not written in the operator formulation.

that is, a task must read all of the locations in its neighborhood before it can write to any of them.

- A *conflict* occurs between tasks t_1 and t_2 if (1) neither task is an ancestor of the other, and (2) one of them writes to the neighborhood of the other ($W(t_1) \cap L(t_2) \neq \emptyset$). If there can be no conflicts between tasks, parallel execution is straightforward since the program is a generalized data-parallel loop in which the iteration range can grow dynamically. In the presence of conflicts, a correct parallel schedule for the program should be serializable: it must appear as if all tasks were performed atomically in some order that respects the ancestor relation.

Non-determinism arises in this formulation because the serialization order between conflicting tasks is not defined. Schedulers that guarantee the same serialization order between conflicting tasks are *deterministic*. In practice, there may be other sources of non-determinism in the system, e.g., the tasks themselves use random variables or read non-deterministic external state; in this chapter, we will restrict ourselves to systems where the only source of non-determinism is the order in which conflicting tasks execute.

[Section 7.1](#) outlines a method of deterministically scheduling any unordered algorithm, but it requires runtime analysis to deal with potentially changing runtime dependencies. Under certain program restrictions, this deterministic runtime scheduling can be refined to use more efficient schedulers (see [Section 7.2](#)).

7.1 Interference Graph Scheduling

One general deterministic scheduling technique is based on successive construction and scheduling of *interference graphs*.

Definition 7.1. *Given a set of tasks P , an interference graph for P is an undirected graph $G_P = (V_P, E_P)$ in which there is a distinct node in V_P representing each task in P , and*

there is an undirected edge $(v_1, v_2) \in E_P$ if the tasks represented by v_1 and v_2 have a conflict.

The interference graph for a set of tasks can be built by executing each task up to its failsafe point (see [Section 2.2.1](#)) while tracking its neighborhood and putting a conflict edge between two tasks if their neighborhoods overlap. This is a conservative approach since it puts a conflict edge between two tasks even if they both read a location that neither of them writes to. A more precise technique which distinguishes reads from writes is described later in [Section 7.1.2](#).

Interference graphs can be used to schedule tasks as follows. The tasks in the task pool P are executed in *rounds*. In each round, the scheduler performs the following activities:

- *inspect*: builds an interference graph G_P for the tasks in P ,
- *select*: finds an *independent set* \mathcal{I} of nodes in G_P and removes the corresponding tasks from P , and
- *execute*: executes the tasks in \mathcal{I} in parallel, adding any newly created tasks to P .

Scheduling is completed when all tasks have been executed. During the *select* phase, it is desirable but not necessary to find a maximal independent set of nodes in the graph.

A subtle point is that the interference graph must in general be rebuilt from scratch each round since the neighborhood of a task is relative to the global state, which is modified by tasks in the *execute* phase. For instance, consider three tasks: task t_1 , which writes 1 to location l_1 ; task t_2 , which reads l_1 and if it is 1, writes to l_2 otherwise it does nothing; and task t_3 , which reads from l_2 . Let the initial value of l_1 be 0. The initial interference graph is $G_P^0 = (V_P^0 = \{v_{t_1}, v_{t_2}, v_{t_3}\}, E_P^0 = \{(v_{t_1}, v_{t_2})\})$. Assume that only task t_1 executes. In the following round, the interference graph is $G_P^1 = (V_P^1 = \{v_{t_2}, v_{t_3}\}, E_P^1 = \{(v_{t_2}, v_{t_3})\})$. Note that the edge between v_{t_2} and v_{t_3} only appears after executing task t_1 .


```

1  Tasks cur , next , todo
2  todo = P;
3  while ||todo|| > 0:
4    // order tasks in todo deterministically
5    next = sort(todo)
6    todo = {}
7    // execute sorted tasks
8    while ||next|| > 0: // for each round...
9      calculateWindow()
10     barrier.wait()
11     // get prefix of size window from next
12     cur , next = getWindowOfTasks(next)
13     // compute neighborhoods of tasks in cur
14     inspect(cur)
15     barrier.wait()
16     // execute successful tasks , move
17     // failed tasks to next , and add any
18     // newly created tasks to todo set
19     todo = todo  $\cup$  selectAndExec(cur , next)
20     barrier.wait()

```

Figure 7.1: Deterministic scheduler

There is an enhancement to this basic scheme that is useful for reducing the overhead of interference graph construction. Note that the scheduling strategy works correctly even if, in each round, the interference graph is constructed only for a subset of tasks in the pool; the remaining tasks are simply delayed to later rounds. This *windowing* scheme can reduce the overhead of interference graph construction when the number of tasks is much larger than the number of threads because the conflict rate monotonically increases with the number of tasks inspected a round. For the windowed scheduler to be deterministic, one must ensure the following in each round: (1) tasks for the current window are chosen deterministically from the task pool, and (2) during the select phase, the independent set of nodes is chosen deterministically. [Section 7.1.1](#) describes an implementation of deterministic interference graph (DIG) scheduling.

```

1  Tasks sort(Tasks todo):
2    // sort tasks in set todo
3
4  void inspect(Tasks cur):
5    doall t in cur:
6      writeMarksMax(id(t), L(t))
7
8  Tasks selectAndExec(Tasks cur, Tasks next):
9    Tasks newWork = {}
10   doall t in cur:
11     if readMarks(L(t)) == {id(t)}:
12       // all reads equal id, so execute task t
13       t()
14       // add new work if any to newWork
15       newWork = newWork  $\cup$  S(t)
16     else:
17       next = next  $\cup$  t
18       writeMarks(id(t), 0, L(t))
19   return newWork
20
21 bool writeMarks(Id expected, Id id, Set<Loc> locs):
22   for loc in locs:
23     atomic:
24       if Mark(loc) == expected:
25         Mark(loc) = id
26       else:
27         return false
28   return true
29
30 void writeMarksMax(Id id, Set<Loc> locs):
31   for loc in locs:
32     atomic:
33       if Mark(loc) < id:
34         Mark(loc) = id
35
36 Set<Id> readMarks(Set<Loc> locs):
37   // return set of ids in mark locations

```

Figure 7.2: Auxiliary functions for deterministic scheduler

7.1.1 Deterministic Interference Graph Scheduling

Figure 7.1 shows the pseudocode for the implementation of deterministic scheduling; auxiliary functions are shown in Figure 7.2. In the pseudocode, **doall** indicates a loop whose iterations are run in parallel. Instead of explicitly building an interference graph, this code directly finds an independent set of tasks by using marks on locations.

A summary of what the scheduler does is the following. The task set *todo* is initialized to the initial set of tasks *P*. These tasks are ordered deterministically to form a sequence *next*. This sequence of tasks is executed over several rounds; in each round, a prefix *cur* of tasks in *next* is tried for execution. Some of these will succeed and others may fail. Tasks created by successful tasks are added to *todo*; these are executed after *next* becomes empty. Failed tasks are added back to *next* and retried in later rounds. Execution terminates when *todo* and *next* become empty.

The *inspect* operation uses *writeMarksMax* to mark the neighborhood of a task, stealing ownership of neighborhood locations from tasks with lower ids. While the mark on a given location may be updated non-deterministically depending on how the tasks in *cur* are scheduled, the final mark values will be the same regardless of the order in which tasks were processed in the *inspect* phase. This is because the maximum (or minimum) element of a set with a total order is deterministic (the set in question is the set of ids of the tasks that read or wrote to a particular location in the current round).

The non-deterministic Galois execution uses exclusive locking (see Section 5.2) to identify conflicts. The behavior of exclusive locking is like the *writeMarks* function in Figure 7.2. One important difference between *writeMarks* and *writeMarksMax* is that *writeMarks* can fail early if it cannot update a mark location, but in order to be deterministic, *writeMarksMax* must attempt to update all mark locations even if it failed to update some of them. If a task skips some mark locations, it changes the set that *writeMarksMax* is computing the maximum of, and if the mark locations skipped depend on a scheduling choice, then the resulting maximums are non-deterministic.

In the second phase, the scheduler selects and executes an independent set of tasks (line 19 in Figure 7.1 and function `selectAndExec` in Figure 7.2). A task is selected if all of its neighborhood locations are still marked with its id at the end of the inspection phase. Tasks selected this way form an independent set in the interference graph. This set is unique because of the total order on ids. If any of the neighborhood locations of a task does not contain its id, the task is not part of the independent set, and it is placed in the *next* set to be executed in a future round. In either case, the marks written by a task are cleared in preparation for the next round.

Execution continues in rounds until there are no tasks left in *next*. If there are no tasks in the *todo* set, the scheduler terminates; otherwise these tasks are moved to *next*, and execution continues. Note that in each round, the task in *cur* with maximum id is guaranteed to execute, so each round executes at least one task.

Before enqueued tasks can be scheduled, they must be assigned a unique id. The assignment of ids must also be deterministic. Ids are assigned as follows. The initial tasks are given ids based on the iteration order of the C++ iterator that contains the tasks. When task t creates task u , the scheduler stores with task u the id of the task that created it $\text{id}(t)$ and a number k indicating whether it was the first, second, third, etc. task created by t . In the sort function, tasks are sorted lexicographically based on the pair $(\text{id}(t), k)$, and the scheduler uses the position in the total order defined by the sort as the id for the new tasks.

The performance of this scheduler depends critically on the window size, so the scheduler uses an adaptive algorithm that grows and shrinks the window size each round depending on the number of tasks that successfully committed in the previous round. The `getWindowOfTasks` and `calculateWindow` functions in Figure 7.1 implement this functionality. The `calculateWindow` function computes the window size for the current round based on the fraction of tasks that committed in the previous round. If the commit ratio is less than some target threshold (0.95 in these experiments), the next window size is scaled down proportionally. If the commit ratio is above the threshold, the window size is doubled. The

`getWindowOfTasks` function simply returns this prefix of tasks in *cur* and postpones the remainder to *next*. Since the number of tasks that commit in a round is independent of the number of executing threads, this heuristic is portable across machines.

To implement the deterministic marking scheme in the Galois system, there are two changes that need to be made to the non-deterministic scheduling system.

First, the default mark values in the Galois system are not ordered. The marking code needs to be modified to keep track of the id of a task and to use that value appropriately when writing mark values.

Second, neighborhoods are not explicitly maintained by the Galois system. Marks are acquired incrementally during execution via user code calls to a data structure library. The only way to get the neighborhood of a task is to execute the task and observe which marks are acquired. To implement the inspect phase, tasks are simply executed, which, by their normal execution, mark locations in their neighborhoods. When a task reaches its failsafe point (the first write to a global location), it immediately returns. To implement the `selectAndExec` phase, tasks are re-executed from the beginning, and instead of writing marks, they check whether the marks that would have been written match the values that have been written. This implements line 11 of [Figure 7.2](#). If a task reads a mark value that is not its id, the scheduler goes to line 17.

This baseline implementation is sufficient to deterministically schedule any un-ordered algorithm in the operator formulation.

7.1.2 DIG Optimizations

The baseline deterministic scheduler described in [Section 7.1.1](#) contains several inefficiencies, which are addressed in an optimized scheduler.

First, it redundantly executes the prefix of a task up to its failsafe point when a task is selected and executed. A more efficient method would be to suspend execution of a task at the failsafe point during the inspect phase and to resume execution in the commit phase.

On resumption, the task must check that all the mark values still match its id (Figure 7.2 line 11). The capability to pause and resume execution can be achieved generally using additional threads or creating continuations. The optimized scheduler uses a more ad-hoc approach, which simulates the effect of forming a continuation without implementing a full compiler transform. The scheduler provides a library function that allows users to allocate objects in the inspect phase which can be recalled during the commit phase. Programmers can use this functionality to manually achieve the same effect as task suspend and resume.

To make sure that resumed tasks are valid to commit, the optimized scheduler makes a small change to the protocol in the inspect phase. Instead of just writing the maximum mark value, a task t checks if the previous value of the mark location is not 0 and not $\text{id}(t)$; if so, by writing its mark, task t will prevent the task u that corresponds to the current mark value from committing. Normally, task u detects this case when the scheduler executes line 11, or in the case of the baseline scheduler, when task u is executed a second time. When using the continuation optimization, t is now responsible for preventing u from executing. It does this by writing to a flag variable that u checks before resuming execution.

Second, the performance of the scheduler is very sensitive to initial task order. Applications that exploit temporal locality execute tasks with overlapping neighborhoods close in time. This typically translates to those tasks being close together in iteration order, which, in the baseline scheduler implementation, means that they typically will be executed in the same round, where they will certainly conflict with each other. This leads to the perverse situation where the scheduler needs to reduce locality to improve performance. The optimized scheduler addresses this issue by assuming that tasks placed close together in iteration order have high locality and places those tasks in separate rounds if possible.

Third, the cost of sorting enqueued tasks can be large relative to the application time. There is a common special case where a task enqueues tasks, but those tasks are drawn from a fixed set of tasks. In this case, tasks can be assigned unique ids before parallel execution, and the programmer can pass these ids to the scheduler, which uses them directly

instead of generating new ids via the sort function.

Two possible optimizations not yet included in the optimized scheduler are the following.

First, instead of sorting tasks to generate ids for enqueued tasks, any deterministic hash function can be used instead. One way to generate unique ids with a deterministic hash function is to use a variation of linear probing with priority writes (Shun and Blelloch, 2014).

Second, the interference graph has an edge whenever two tasks have overlapping neighborhoods, but if both tasks only read locations the overlapping region, these tasks can be allowed to proceed in parallel. This situation does not frequently occur in the applications evaluated in Section 7.1.3, but allowing such tasks to proceed in parallel only requires a small modification to the deterministic interference graph scheduling algorithm. Each task t now has two ids, $\mathbf{wid}(t)$ which is the id it marks when it wants to write to a location, and $\mathbf{rid}(t)$ which is the id it marks when it wants to read a location. Marking a location to write is the same as before. When marking a location to read and the previous owner is a write id, the behavior is the same as a write. When marking a location to read and the previous owner u is a read id, the task t updates a disjoint-set data structure to record that t and u both have a read dependency on the same location. After the inspect phase, another phase is added to check if any location read was subsequently marked by a writer, and if so, the tasks that have a read dependency on that location are disabled for the current round. To ensure progress, the read and write id the greatest task (in a window) should be greater than the read and write ids of any other task.

Even with these implemented and proposed optimizations, there are some inherent inefficiencies that are introduced by any DIG scheduling implementation compared to non-deterministic scheduling.

1. The deterministic scheduler executes many more instructions than non-deterministic scheduling.

2. The deterministic scheduler introduces a concept of rounds that is not present in the original program. These rounds are implemented using global synchronization. Rounds extend the critical path length of a program because the scheduler cannot proceed to the next round until all of the tasks are processed for the current round.
3. The scheduler executes tasks according to a particular schedule, but that schedule may not be the best performing one among possible program schedules.
4. The execution of a task is broken into two parts, the inspect phase and the execution phase, separated by a barrier. The memory locations accessed during the inspect phase of a task are very likely to be accessed by the execution phase of the same task, but under DIG scheduling, these two phases are temporally separated by a factor that is a function of number of tasks attempted during a round, which is typically very large. Conversely, increasing locality by reducing the number tasks attempted in a round increases the number of rounds executed, which increases the critical path length of the program.

7.1.3 Evaluation

To evaluate DIG scheduling, applications were drawn from three different sources: the PARSEC (v2.1) benchmark suite (Bienia et al., 2008), the problem based benchmark suite (PBBS) (v0.1) (Blleloch et al., 2012), and the Lonestar (v2.1.5) benchmark suite (Kulkarni et al., 2009).

The PARSEC benchmark suite has been used in previous evaluations of deterministic scheduling (Bergan et al., 2010a; Devietti et al., 2011; Liu et al., 2011). It contains twelve applications or kernels. Most are parallelized using the pthread library. The evaluation uses the three benchmarks that have OpenMP implementations: **blackscholes**, **bodytrack** and **freqmine**. The blackscholes and freqmine results are for the simlarge input, while the body-track results are for the native input.

The PBBS programs (Blelloch et al., 2012) are organized by problem, and each problem has one or more solution programs, at least one of which is deterministic. There are a total of sixteen problems, but many of these programs are data-parallel or nested data-parallel, and their performance depends largely on factors like good load balancing and not on scheduling and therefore were excluded from this evaluation. The remaining deterministic programs solved the five remaining problems: breadth-first search (**BFS**), Delaunay triangulation (**DT**), Delaunay mesh refinement (**DMR**), maximal independent set (**MIS**), and maximal matching. Maximal matching was excluded from the evaluation due to its similarity to maximal independent set. In these codes, determinism is ensured by application-specific techniques customized to each application, and they typically involve bulk-synchronous execution in rounds. The PBBS maximal independent set program is data-parallel, but it is included in this evaluation for comparison with a non-deterministic maximal independent set program that exists in the Lonestar suite.

From the Lonestar benchmark suite, the evaluation contains four programs that solve the same problems as those included from PBBS, using the same algorithms, and an implementation of the preflow-push algorithm (**PPF**) that uses the global relabeling heuristic to improve convergence (see Section 2.3). The deterministic implementations of all Lonestar programs are automatically generated by applying the DIG scheduler (Section 7.1.1) and its optimizations (Section 7.1.2).

There is one small difference between the PBBS and Lonestar implementations of Delaunay triangulation. The algorithmic complexity of Delaunay triangulation depends on the order in which points are inserted, and random insertion order has been shown to be optimal (Clarkson and Shor, 1989). In the PBBS implementation, points are randomized offline. In the Lonestar implementation, points are reordered online using the biased randomized insertion order algorithm (Amenta et al., 2003). For comparison purposes, the evaluation does not include the reordering time in either implementation.

The performance of the PBBS and Lonestar benchmarks can vary significantly with

the type of input. From experience, the behavior across random inputs for an application is largely similar, so the evaluation uses a single representative input for each application. These inputs are largely drawn from the evaluation of Blelloch et al. (Blelloch et al., 2012). The BFS results use a random graph of 10 million nodes where each node is connected to five randomly selected nodes. The DMR results use a Delaunay triangulated mesh of 2.5 million randomly selected points from the unit square. The DT results use 10 million points randomly selected from a unit square. The MIS results use the same input as BFS. The PFP results use a random graph of 2^{23} nodes with each node connected to 4 random neighbors.

In the experimental results, the variant **g-n** denotes the original non-deterministic Lonestar application, and the deterministic variant generated from DIG scheduling is called **g-d**. The variant **PBBS** denotes the PBBS version of the application.

The evaluation uses three machines and runtimes are the average of at least three runs for each application/machine/thread-count combination. The three machines are

1. **m4x10**, a machine running Ubuntu Linux 10.04 LTS 64-bit (Linux 2.6.32) with four ten-core Intel Xeon E7-4860 (2.27 GHz) processors;
2. **m4x6**, a machine running Ubuntu Linux 10.04 LTS 64-bit (Linux 2.6.32) with four six-core Intel Xeon E7540 (2.0 GHz) processors; and
3. **numa8x4**, an SGI UV machine (ccNUMA) running SuSE Enterprise 11 SP1 64-bit (Linux 2.6.32.24) with eight four-core Intel E7520 (1.87 GHz) processors. The processors of numa8x4 are divided into blades of two processors each and enclosures of two blades each. Inter-blade communication uses SGI NUMALink 5.

Programs are compiled with icc version 12.1 with the -O3 optimization flag. For the PBBS programs, the Cilk runtime is used to manage and load balance threads. For the Lonestar programs, the Galois runtime system is used.

The evaluation is divided into four parts.

1. The first part (**Application Characteristics**) describes applications characteristics useful for understanding the performance results.
2. The second part (**Deterministic Thread Scheduling**) compares the performance of non-deterministic programs with and without CoreDet (**Bergan et al., 2010a**), a system that provides determinism by modifying how threads are scheduled. Because the C++ language primitives used in the Lonestar programs are not supported by CoreDet, the evaluation uses non-deterministic versions of the PBBS programs as representative of the non-deterministic benchmarks. The evaluation shows that with CoreDet, the non-deterministic PBBS programs do not perform well and have a median slowdown of 3.7X (min: 1.3X, max: 55X) compared to running without CoreDet. These experiments show that systems like CoreDet that provide determinism through deterministic thread scheduling are not suitable for irregular applications, which have relatively fine-grain tasks.
3. The third part (**DIG Scheduling**) compares the performance of non-deterministic Galois programs (g-n), generated deterministic implementations of these Galois programs (g-d), and handwritten deterministic PBBS programs for the same problems. Overall, the results show that at the maximum number of threads on each machine, (1) g-n variants achieve a median improvement of 4.2X compared to g-d, (2) g-n variants are 2.4X faster than the PBBS variants, and (3) g-d variants are only 0.62X slower than the PBBS variants.
4. These results show that the automatically generated deterministic Galois programs are comparable in performance to the handwritten PBBS programs and that there is a significant performance penalty for deterministic execution. Finally, a study with performance counters (**Determinism and Locality**) reveals that, for the most part, non-deterministic programs perform better than deterministic ones because they exploit more locality.

		$p = 1$			$p = 40$	
	Variant	Rounds	Abort Ratio	Tasks per μs	Abort Ratio	Tasks per μs
bfs	g-d	1700	0.08	0.45	0.08	9.76
bfs	g-n	0	0	1.32	0	39.92
bfs	pbbs	11	0	1.24	0	24.27
dmr	g-d	1287	0.11	0.13	0.11	2.53
dmr	g-n	0	0	0.26	< 0.01	8.98
dmr	pbbs	1165	0.03	0.18	0.03	2.90
dt	g-d	35213	0.27	0.12	0.27	1.78
dt	g-n	0	0	0.24	< 0.01	7.47
dt	pbbs	1330	0.10	0.11	0.10	2.48
mis	g-d	100	0.08	0.77	0.08	21.05
mis	g-n	0	0	3.98	< 0.01	79.69
mis	pbbs	29	0.05	14.59	0.05	143.12
pfp	g-d	21047	0.04	0.26	0.04	2.58
pfp	g-n	0	0	0.67	< 0.01	14.99

Figure 7.3: Abort ratio and task execution rates on machine m4x10

Application Characteristics

Previous evaluations of deterministic scheduling have focused mostly on applications with coarse-grain tasks that communicate relatively infrequently. Deterministic scheduling for these kinds of applications can be supported using relatively heavyweight mechanisms since the overhead of the system is a small fraction of the overall execution time. However, these mechanisms may not be useful for applications with very lightweight tasks that communicate frequently. On a shared-memory system, the concept of communication is less well-defined compared to a distributed system, but one approximation is the number of atomic updates an application performs. Figure 7.3 shows task execution rates and abort ratios on machine m4x10 with 1 and 40 threads. Figure 7.4 shows atomic update rates. For

		$p = 1$		$p = 40$	
	Variant	Count	Rate	Count	Rate
mis	pbbs	< 1	< 0.01	< 1	< 0.01
freqmine		< 1	< 0.01	< 1	< 0.01
bodytrack		< 1	< 0.01	15	0.07
dt	pbbs	1445	0.55	1522	0.66
blackscholes		< 1	< 0.01	48	0.77
bfs	pbbs	7191	1.24	7162	1.36
dmr	pbbs	2360	1.00	2634	1.37
pfp	g-n	4622	2.24	1519	27.57
dmr	g-n	707	1.59	583	36.21
dt	g-n	1376	3.10	920	79.94
mis	g-n	19292	10.27	4628	100.17

Figure 7.4: Atomic updates by application measured by binary instrumentation on machine m4x10. Variant has been g-d omitted. Count is atomic updates per million instructions executed. Rate is atomic updates per microsecond.

the deterministic variants, the number of rounds is also shown. For PBBS variants, this is the number of bulk-synchronous rounds of the handwritten deterministic scheduling.

Figure 7.3 shows that PBBS and Lonestar benchmarks have very fine-grain tasks. For example, the g-n variant of DMR, running on one thread, commits 0.26 tasks per microsecond, which translates to 3.8 microseconds per task (this is the parallel version of the code with synchronization, running on one thread), which is on the order of a thousand cycles. On 40 threads, this parallel program commits roughly 9 tasks per microsecond, which translates to a throughput of roughly 0.11 microseconds per task.

The figure also shows that the abort ratios of the g-n variants of all applications are essentially zero even at 40 threads. Conflicts between tasks in the non-deterministic variants are very rare: this is because there are a large number of tasks compared to the number of threads. The deterministic variants g-d and PBBS have larger abort ratios because in each round, the number of tasks whose neighborhoods are inspected is typically larger than the

number of threads. Conflicts can also happen with only one thread when two tasks with overlapping neighborhoods are inspected in the same round.

Figure 7.4 shows that the PARSEC benchmarks—blackscholes, bodytrack and freqmine, which are frequently used to evaluate deterministic schedulers—have orders of magnitude fewer atomic updates than the irregular algorithms of the PBBS and Lonestar suites. For example, blackscholes at 40 threads performs atomic updates at a rate of about 1 update per microsecond, while the MIS g-n variant performs atomic updates at the rate of 100 updates per microsecond.

These qualitative differences in application characteristics significantly impact the design of deterministic schedulers, which is quantified in the next section.

Deterministic Thread Scheduling

This section presents the performance results from using CoreDet, a deterministic thread scheduler, on the benchmark applications. Unlike DIG scheduling, CoreDet runs on unmodified pthread programs.

Ideally, CoreDet could be directly applied to run the PARSEC and g-n non-deterministic programs deterministically. Unfortunately, the CoreDet compiler is based on the older LLVM v2.6 compiler, and it is unable to compile any of the g-n programs. To get around this problem, the evaluation exploits the fact that BFS, DMR and DT in PBBS are deterministic implementations of non-deterministic algorithms whose program structure is similar to **Figure 7.1**. The programs are transformed by hand to be non-deterministic, and these programs are run with CoreDet. The MIS benchmark is left as a data-parallel program.

CoreDet supports many different conflict detection implementations. This evaluation uses CoreDet in its low-overhead, synchronization-only mode, which reduces the system to an implementation of the Kendo algorithm ([Olszewski et al., 2009](#)) and requires all synchronization between threads to use the pthread library. To fairly compare with the PARSEC and PBBS benchmarks, which use OpenMP or Cilk runtimes for parallelization,

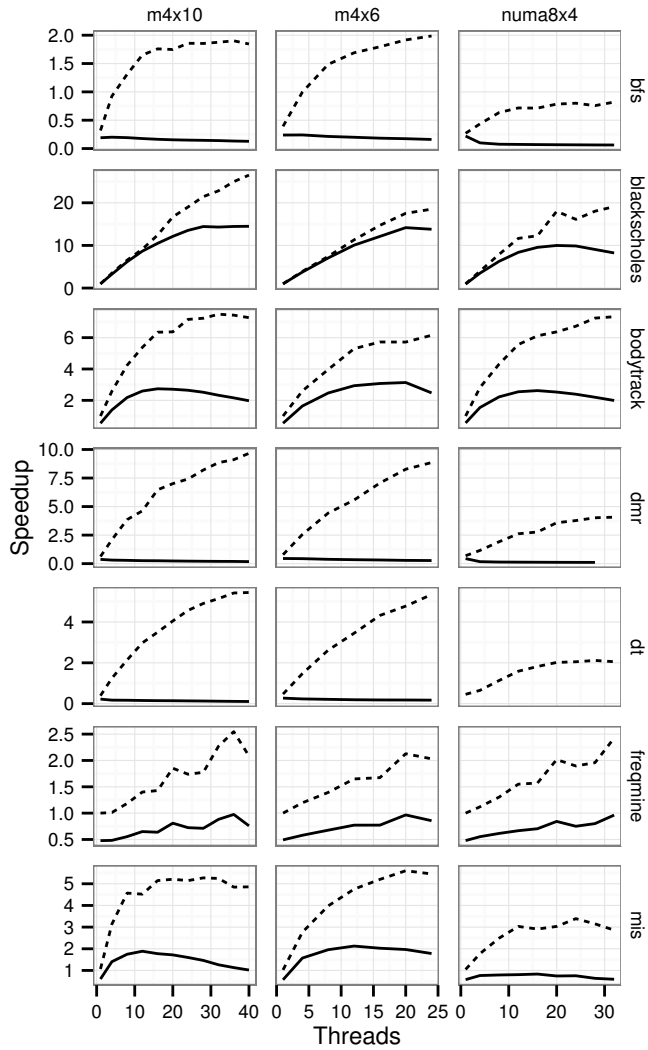


Figure 7.5: Speedup with (solid lines) and without (dotted lines) CoreDet system on non-deterministic programs. Speedup baselines are in [Figure 7.7](#). Some DMR and DT runs on numa8x4 timed out after 10 minutes.

all results comparing with CoreDet use programs where the calls to the OpenMP or Cilk runtimes are replaced with calls to a simple pthread-only runtime. The simple pthread-only runtime likely introduces minor inefficiencies compared to the more optimized Cilk and OpenMP runtimes, but they are dwarfed by the overheads of CoreDet. The results in this section use the LLVM v2.6 compiler with the -O3 optimization flag to compile programs.

Figure 7.5 summarizes the results using the CoreDet system to make PARSEC and modified PBBS programs deterministic. CoreDet works well for blackscholes: the performance with CoreDet is almost the same as without CoreDet for a small number of threads. As the number of threads increases, the gap between using CoreDet and not using CoreDet increases, hinting at a serialization bottleneck in the deterministic scheduler. The bodytrack and freqmine applications show more limited speedups. For the modified PBBS programs, the performance with CoreDet is poor except for MIS, the data-parallel code. The BFS, DMR and DT applications perform substantially more synchronization than the PARSEC applications and the MIS code (see **Section 7.1.3**). Overall, at the maximum number of threads on each machine, the benchmarks in this suite experience a median slowdown of 3.7X (min: 1.3X, max: 55X) compared to non-CoreDet runs.

Although this evaluation uses CoreDet, the other deterministic thread schedulers such as Kendo and DThreads have similar scheduling algorithms and differ mainly in how they deal with racy data accesses, which none of the modified PBBS programs have.

These results make the case that a different approach than deterministic thread scheduling is needed to handle applications that perform orders of magnitude more synchronization than more conventional programs like the PARSEC benchmarks.

DIG Scheduling

Figure 7.6 shows the speedups of g-n, g-d and PBBS relative to the best performing serial implementations shown in **Figure 7.7**. For BFS, the baseline is the code of Schardl and Leiserson. It uses data structures customized to the BFS problem (**Leiserson and Schardl, 2010**).

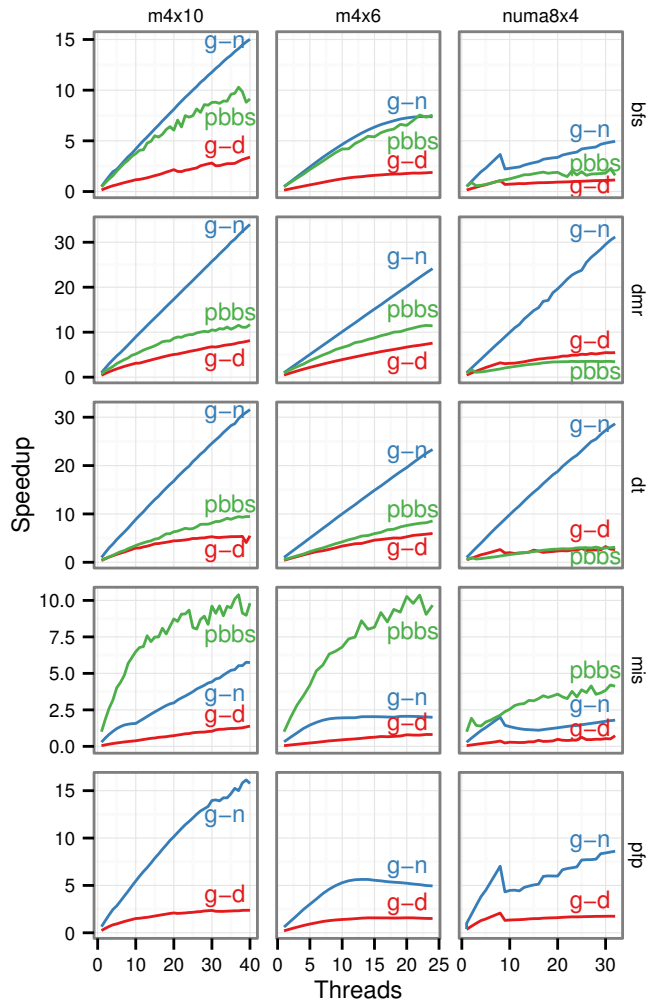


Figure 7.6: Speedup of selected deterministic and non-deterministic variants. Speedup baselines are in Figure 7.7.

	Machine	Var.	Time (s)		Machine	Var.	Time (s)
bfs	m4x10	cilk	3.76	dt	m4x10	g-nd	42.35
bfs	m4x6	cilk	4.36	dt	m4x6	g-nd	56.37
bfs	numa8x4	cilk	4.85	dt	numa8x4	g-nd	61.15
bs	m4x10		8.42	fm	m4x10		7.95
bs	m4x6		11.27	fm	m4x6		10.57
bs	numa8x4		12.01	fm	numa8x4		11.33
bt	m4x10		164.84	mis	m4x10	pbbs	0.72
bt	m4x6		216.31	mis	m4x6	pbbs	0.90
bt	numa8x4		249.07	mis	numa8x4	pbbs	0.91
dmr	m4x10	g-nd	44.48	pfp	m4x10	hi_pr	13.64
dmr	m4x6	g-nd	60.04	pfp	m4x6	hi_pr	14.64
dmr	numa8x4	g-nd	63.29	pfp	numa8x4	g-nd	26.17

Figure 7.7: Baseline times in seconds for speedup calculations (bs: blackscholes, bt: bodytrack, fm: freqmine). These are the best times for any variant with one thread. Cilk is a parallel BFS code (Leiserson and Schardl, 2010). hi_pr is a sequential implementation of PFP (Goldberg and Tarjan, 1988).

For preflow-push, the baseline is the highly optimized hi_pr implementation from Goldberg and Tarjan (Goldberg and Tarjan, 1988). For the other benchmarks, the best performing versions were from the benchmark suites considered in this evaluation.

The figure shows that the best performing variant overall is g-n, which has a median improvement of 2.4X over corresponding PBBS programs at the maximum number of threads on each machine (from Figure 7.8). The benefit is largest on the numa8x4 machine where the scalability of the PBBS variant is particularly poor, but there are positive benefits for almost all non-deterministic variants. Figure 7.6 also suggests that there are also significant scalability advantages to the non-deterministic variants compared to the deterministic ones. The g-n variants are able to achieve at least a 15X speedup on m4x10 for four of the five applications.

The main outlier in these results is the behavior of MIS. As mentioned above, the PBBS variant of MIS is a data-parallel program, and its execution characteristics are significantly different than the g-n or g-d variants. The main conclusion from this benchmark

is that if one has a deterministic algorithm for some problem, it may be better to use that algorithm rather than deterministically schedule a non-deterministic algorithm for the same problem. However, [Section 7.1.3](#) shows that the PBBS variant of MIS is more sensitive to input ordering than the g-n one.

The sharp drop in performance at eight threads in some of the numa8x4 runs is caused by inter-NUMA node communication. Runs of eight threads or less are scheduled to run on a single NUMA node. Runs with more than eight threads use more than one NUMA node, and remote memory accesses are significantly more expensive than local memory accesses. The g-n variants for DMR and DT are able to tolerate the transition to inter-node communication due to locality optimizations in the Galois runtime system. Fully exploiting locality may be difficult in deterministically scheduled programs due to the multiple parallel phases needed to execute a task. [Section 7.1.3](#) attempts to quantify the locality lost in these benchmarks.

These results suggest that for irregular applications, determinism comes at a significant price in performance, even if the determinism is obtained through hand-optimized, application-specific code.

A previous study by Blelloch et al. ([Blelloch et al., 2012](#)) reached the opposite conclusion; for example, they found that the deterministic PBBS version of DT was substantially faster than the non-deterministic DT program in the Lonestar suite. Unfortunately, their study did not ensure that the same algorithm was used for a given problem; in particular, the DT algorithm in the PBBS suite is different (and more efficient) than the one that was used in the Lonestar suite at the time their study was performed. For this evaluation, the DT algorithm in the Lonestar suite was reimplemented to match the algorithm used in PBBS, so the performance differences between non-deterministic and deterministic programs for a given problem are not entangled with algorithmic differences.

DIG scheduling vs. determinism by construction How good is the general-purpose deterministic scheduler described in [Figure 7.1](#) compared to the application-specific, hand-

		m4x10				m4x6				numa8x4			
Variant		Mean	Max	I ₁	I _{max}	Mean	Max	I ₁	I _{max}	Mean	Max	I ₁	I _{max}
bfs	g-n	1.28	1.68	1.07	1.64	1.10	1.20	1.00	0.98	2.23	3.48	1.00	3.09
bfs	g-d	0.31	0.37	0.33	0.37	0.29	0.32	0.28	0.25	0.61	1.03	0.30	0.71
bfs	pbbs	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00
dmr	g-n	2.12	2.99	1.39	2.90	1.63	2.11	1.11	2.11	5.45	9.12	1.18	9.12
dmr	g-d	0.62	0.71	0.57	0.70	0.60	0.66	0.52	0.66	1.38	1.75	0.55	1.59
dmr	pbbs	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00
dt	g-n	2.81	3.34	2.34	3.34	2.43	2.76	1.92	2.73	6.70	9.30	2.07	9.26
dt	g-d	0.72	0.95	0.87	0.58	0.77	0.89	0.79	0.70	1.15	2.08	1.11	0.86
dt	pbbs	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00
mis	g-n	0.40	0.64	0.29	0.59	0.27	0.35	0.31	0.21	0.48	0.94	0.28	0.44
mis	g-d	0.09	0.15	0.05	0.14	0.07	0.09	0.05	0.08	0.12	0.18	0.05	0.18
mis	pbbs	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00

Figure 7.8: Performance of variants relative to PBBS variant. Let $\text{time}_{\text{PBBS}}(p)$ and $\text{time}_{\text{var}}(p)$ be the times for variant PBBS and var respectively with p threads. The performance number shown is $\text{time}_{\text{PBBS}}(p)/\text{time}_{\text{var}}(p)$. I_1 and I_{max} show performance at 1 and the maximum number of threads respectively.

optimized deterministic code in the PBBS programs? Figure 7.8 shows the performance for different variants relative to the same baseline, the PBBS variant.

Across all machines and benchmarks and at the maximum number of threads (I_{max}), the median performance of the g-d variant relative to PBBS is 0.62X. If the benchmark mis is dropped, the median performance is 0.70X. These results show that the general-purpose deterministic scheduler described in Figure 7.1 provides reasonable performance compared to application-specific, hand-optimized determinism by construction code, although there is room for improvement.

For DMR and DT, the PBBS variants correspond to a handwritten version of DIG scheduling of the g-nd variants. The performance difference between the g-d and PBBS variants is largely due to the application-specific implementation of resuming tasks and the handtuned window selection policy used in PBBS.

		m4x10			
Variant		Mean	Max	I_1	I_{\max}
bfs	g-d	0.31	0.37	0.33	0.37
bfs	without	0.27	0.32	0.30	0.32
dmr	g-d	0.62	0.71	0.57	0.70
dmr	without	0.48	0.54	0.43	0.52
dt	g-d	0.72	0.95	0.87	0.58
dt	without	0.56	0.71	0.68	0.49
mis	g-d	0.09	0.15	0.05	0.14
mis	without	0.08	0.13	0.05	0.12

Figure 7.9: Performance without continuation optimization relative to PBBS variant on machine m4x10. I_1 and I_{\max} show performance at 1 and the maximum number of threads respectively.

Impact of continuation optimization The g-d variants use the continuation optimization described in [Section 7.1.2](#), which requires some user input to form proper continuations. This transformation could be done by a compiler, but the DIG scheduling system does not implement this yet. To weigh the effect of this optimization, [Figure 7.9](#) shows the performance of programs without this optimization. Overall, the continuation optimization provides a median improvement of 1.14X for the deterministic programs and provides a significant improvement only for the relatively more complicated DMR and DT programs.

Determinism and Locality

[Section 7.1.2](#) describes several inherent limitations of DIG scheduling. This section quantifies two of those costs. First, DIG scheduling can reduce existing locality, and second, DIG scheduling can make it more difficult to exploit locality.

Intra-task locality DIG scheduling decreases locality by splitting a task, which might have significant intra-task locality, into two phases well-separated in time. The impact of

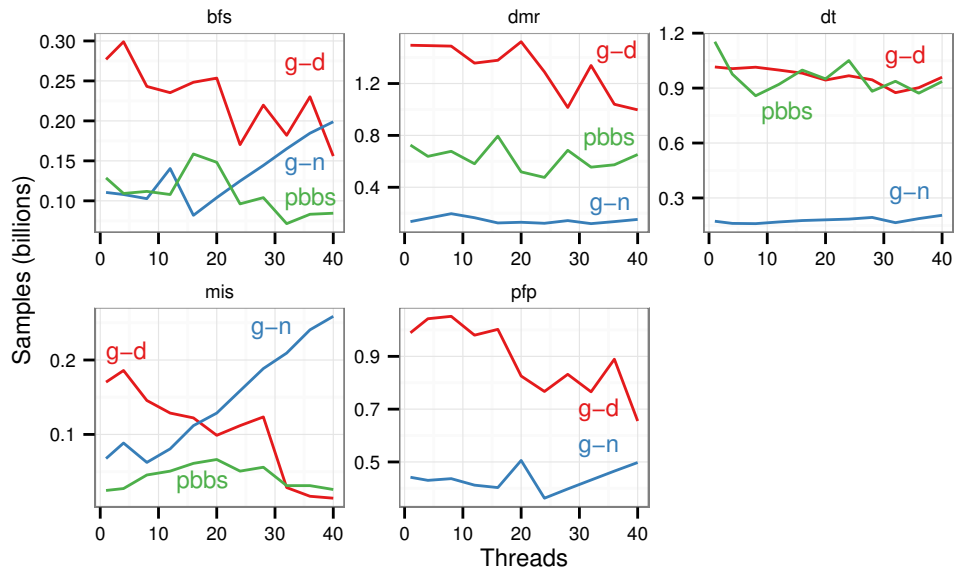


Figure 7.10: Samples of DRAM access performance counter on machine m4x10

this transformation can be quantified by measuring performance counter information about memory-level events.

Figure 7.10 gives the number of data requests satisfied from DRAM for the g-n, g-d, and PBBS application variants. One reason for the difference between g-d and PBBS for DMR is due to the increased number of memory accesses needed to sort new tasks. The DT application has many of the same trends as DMR, but since DT does not create any new tasks, the g-d variant has about the same number of DRAM accesses as PBBS. Overall, the non-deterministic variants typically have far fewer samples than the deterministic ones, but is the change in samples enough to explain the difference in performance?

One way to answer this question is to see how the observed data fits a simple model of performance. Let efficiency be speedup normalized by the number of threads. One simple model is that there is a linear relationship between the change in efficiency and the change of some performance counter. Symbolically, let eff_{var} and PC_{var} be the efficiency and performance counter value, respectively, of some application variant with some number of threads on a machine, and let eff_{ref} and PC_{ref} be the likewise for a particular reference

variant of the same application with the same number of threads. We would like to know how well the following linear model fits the observed data

$$\text{eff}_{\text{var}} = B_0 + B_1(\text{PC}_{\text{ref}}/\text{PC}_{\text{var}})\text{eff}_{\text{ref}}.$$

Fitting the above linear model to our observed data on machine m4x10 reveals that the change in DRAM accesses significantly predicts the change in performance, $\beta = 0.35$, $t(108) = 16.8$, $p < 0.001$. The change in this performance counter also explained a significant portion of the variance in change of performance, $R^2 = 0.72$, $F(1, 108) = 282$, $p < 0.001$. There are performance counters that are more highly correlated ($R^2 \geq 0.75$), clock cycles for instance, but that relationship is trivial.

DRAM accesses are not strongly correlated ($R^2 < 0.9$) to performance only for the MIS variants. In these cases, performance counters that measure operations closer to the processor such as instruction length decoder stalls and L1 cache misses are more closely correlated to performance. This suggests the behavior of MIS is qualitatively different than the other variants, which is not surprising given that the deterministic MIS and non-deterministic MIS variants are two different algorithms.

Figure 7.11 shows the fitted linear model along with observed and predicted efficiencies according to the model. The g-d variants across applications are well-predicted by the model. The main exception is MIS where there is not much variation in DRAM accesses or observed performance (i.e., MIS points are clustered along small vertical and horizontal bands). But among DIG scheduling implementations (i.e., the BFS, DMR, DT and PFP applications with the PBBS or g-d variants), there is a strong correlation between change in DRAM accesses and change in performance.

Among g-n variants, the outliers for the model are the DMR and DT applications. Here, the predicted performance is much higher than the observed performance, which means that the g-n variants have fewer DRAM accesses but that is not resulting in a corresponding increase in performance. One possible explanation is that factors that are not

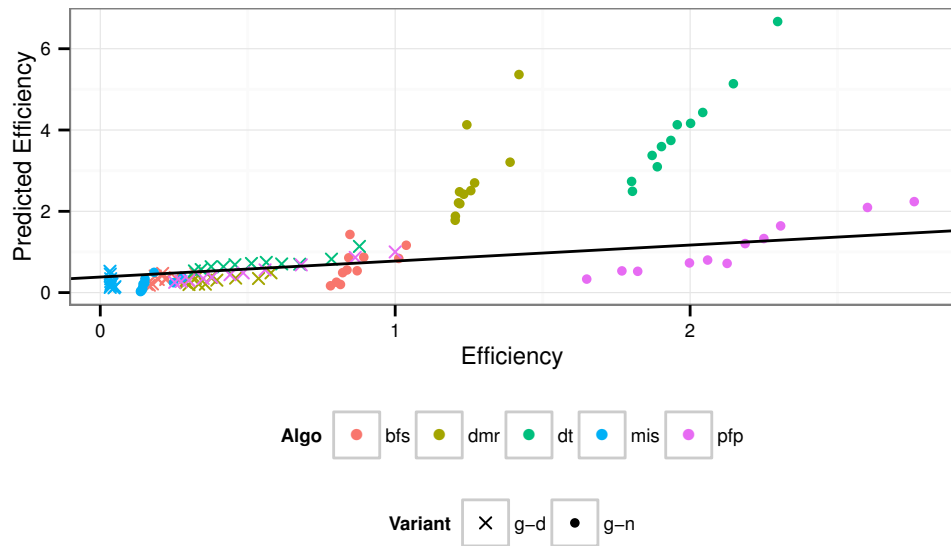


Figure 7.11: Correlation between predicted efficiency and measured efficiency on machine m4x10 across applications, variants and thread counts. Line indicates least-squares fit for linear model.

modeled such as memory bandwidth or memory allocation performance are now playing a stronger role.

Inter-task locality Figure 7.12 shows how the performance of MIS can vary on the same input, depending on whether the input is randomized or sorted. The input graph is a 2D mesh which is ordered by sorting nodes according to a space-filling curve. The left plot (ordered) shows that the g-n variant is able to effectively exploit the locality in the input data and obtains far better performance than the g-d or PBBS variants. When the input graph is randomized, there is no locality to be exploited, and the PBBS version performs slightly better than the g-n version.

In summary, non-deterministic programs can more readily exploit both intra-task and inter-task locality. The execution of a single task is not divided into phases separated in time, so they can exploit intra-task locality better. Furthermore, locality in the input data, which leads to inter-task locality, is easier to exploit.

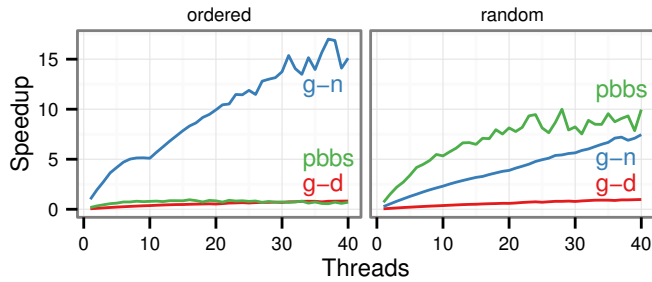


Figure 7.12: Effect of reordering input for MIS. Speedup is relative to the best variant with one thread among the two inputs (g-n, 4.3 seconds). The ordered input is a graph of a 2D mesh. The nodes are sorted according to a space-filling curve.

7.2 Refining Interference Graph Scheduling

Deterministic interference graph scheduling is general but heavyweight. One source of overhead is that the neighborhood of a task t must be computed and then recomputed if a task u with an overlapping neighborhood executes before t . In some cases, the neighborhood of a task does not change due to the execution of another task. An example is a topology-driven, local computation algorithm with an operator that just accesses the direct neighbors of a node in a graph. Since the algorithm is topology-driven, whether a node is active does not depend on the execution of another task, and since the algorithm is a local computation, the graph structure does not change, so the direct neighbors of a node also do not change. In this case, an interference graph of tasks can be built once and reused over multiple rounds of computation. Deterministic scheduling then looks like traditional task graph scheduling. Hassaan et al. present a number of sufficient conditions for using task graph scheduling for the operator formulation (Hassaan et al., 2015).

When unordered local computations operate on undirected graphs and their operators read the direct neighbors of an active node and only write to the active node, the interference graph of tasks is a subgraph of the application graph that is being manipulated by the algorithm. Instead of constructing an interference graph at runtime, scheduling can be done just-in-time by constructing independent sets or graph coloring on the application graph itself when it is read by the program (Kaler et al., 2014).

7.3 Related Work

The majority of deterministic parallel systems work by executing tasks in rounds and deterministically resolving conflicts when two tasks access the same resource in a round. A common way to resolve conflicts is to buffer updates privately and then deterministically merge updates to form the new state for the next round. Hardware systems like RCDC (Devietti et al., 2011) and Calvin (Hower et al., 2011) work this way, as well as runtime replacements like DThreads (Liu et al., 2011) and Kendo (Olszewski et al., 2009), compiler-based systems like CoreDet (Bergan et al., 2010a), OS systems like dOS (Bergan et al., 2010b) and Determinator (Aviram et al., 2010), and some parallel programming models like Grace (Berger et al., 2009).

Ideally, a deterministic parallel system would provide the following three features.

- **On-demand determinism.** It should be possible to turn deterministic execution on and off without much effort. Deterministic execution often imposes a substantial runtime overhead, particularly for parallel programs with fine-grain tasks. This overhead may be acceptable in some cases, but it should be possible to turn off determinism when desired.
- **Portability.** The output of a deterministic program should be the same regardless of the machine that it runs on. At the very least, this means that the output should not depend on the number of executing threads. Portability ensures that programs enjoy the benefits of determinism even when moving between machines.
- **Parameter-freedom.** If there are scheduling parameters that must be tuned to achieve good performance, they should not affect the output state. Since optimal values for such parameters vary by machine, such scheduling parameters hinder portability by providing an incentive for producing different results on different machines.

The deterministic interference graph scheduler described in Chapter 7 provides all these three features. Prior systems have not.

In systems that are deterministic by construction such as DPJ (Bocchino et al., 2009), nested data-parallel programs (Blumofe et al., 1995), stream programs (Thies et al., 2002) and commutativity-based techniques (Burckhardt et al., 2010; Blelloch et al., 2012), tasks have no conflicts. However, it is not possible to write non-deterministic programs in these approaches. Bocchino et al. have shown how to extend DPJ to compose safely with non-deterministic programs (Bocchino et al., 2011), but as yet, there is no system to make the resulting program deterministic on demand.

For round-based systems, an important characteristic of the system is how tasks are determined. The hardware systems and Kendo use the number of instructions executed (or a similar proxy) to divide sequences of instructions into tasks. Depending on how conflicts are detected, task boundaries may also have to be formed at every memory fence, synchronization instruction, store buffer completion, etc. RCDC and the bounded mode of Calvin form tasks based on when the store buffer is full, which is a micro-architectural event. This means that executions may not be reproducible across different processor implementations. CoreDet, Kendo and the unbounded mode of Calvin form tasks based on the number of executed instructions, so their results should be the same between processor implementations.

The determinism guarantee of these systems is still quite fragile, because the insertion of a single instruction will produce a program that generates different outputs. Also, performance is sensitive to the task length. Devietti et al. show that system overheads can vary between 160%–250% depending on the task size parameter (Devietti et al., 2011).

In contrast, systems like Grace and DThreads form their tasks based on synchronization instructions, which means that adding non-synchronization instructions will not change the decomposition of the program into tasks. However, this flexibility comes at a cost as tasks are now quite long, and load balancing becomes an issue. DThreads uses a sequential token passing algorithm to deterministically process synchronization events, so the entire sequence of instructions bounded by synchronization instructions is blocked waiting for the token. Kendo, which breaks tasks up into smaller pieces, can extract more paral-

lelism by executing a prefix of instructions before the synchronization instruction. More recently, Cui et al. have proposed that users add performance hints akin to thread barriers to improve the load balancing of a deterministic scheduler (Cui et al., 2013).

Kendo, CoreDet, Determinator and some PBBS programs (Blelloch et al., 2012) have a tunable parameter that controls the task or round size, but they have no method to adaptively set that parameter based on observed execution. dOS uses instruction-based task formation, but it uses an adaptive algorithm like the one described in Section 7.1.1 to deterministically adjust the task size based on observed parallelism. Calvin uses a standard hardware two-bit predictor to dynamically increase task size when there is no synchronization in a task (Hower et al., 2011).

Chapter 8

Comparison with Other Parallel Programming Models

This chapter tries to answer two questions: (1) how necessary is a new parallel programming model, and (2) if one is necessary, why should the Galois system be preferred over existing systems?

To address the first question, consider the current state of parallel programming for shared-memory machines. Parallel programs are written using a data-parallel programming model (e.g., MapReduce, OpenMP, Cilk), or they are written using the operating system thread primitives directly. The summary of algorithms in [Section 2.3](#) shows that many useful algorithms are not strictly data-parallel, so programmers must use threads, which are an error-prone and low-productivity programming abstraction.

The next natural point of comparison is transactional memory (TM), a hardware mechanism designed to improve the productivity of parallelizing arbitrary code. TM still requires spawning threads, but once programmers identify atomic regions of execution and delineate them for the TM system, TM implementations take care of the error-prone synchronization automatically. [Section 8.1](#) shows that the performance of TM, even with hardware implementations, on the well-studied STAMP benchmark suite is still poor.

To get a sense of how much room there is for performance improvements, [Section 8.1](#) shows how to rewrite the STAMP benchmarks to improve their performance. To ground the scope of the rewriting, the section focuses on transformations that enforce the disjoint accesses ([Principle 4.1](#)) and virtualization ([Principle 4.2](#)) principles introduced in [Section 4.1](#). By following these principles, the median *improvement over STAMP* with 32-thread runs of an Intel Westmere machine is 3.94X (min: 0.25X, max: 169X, geomean: 5.3X). These results support the claim that parallel codes will have to be rewritten in some form to achieve scaling on parallel hardware. Additionally, these results support the claim that these scalability principles are sufficient conditions on scalable implementations.

Now, the question becomes if programs should be rewritten for scalable parallelism, what is the appropriate programming model? [Section 8.2](#) addresses this question in two ways for the graph analytics application domain. First, it shows that existing graph analytics programming models are restricted versions of the operator formulation, and programs written in other graph analytics programming models can be run using Galois with the same and often better performance. Second, since the operator formulation is strictly more expressive than the graph analytics programming models, there are some programs that they cannot express efficiently. This expressibility gap has practical performance implications as [Section 8.2](#) shows that orders of magnitude performance improvements can be gained by implementing better algorithms, which are not possible in previous graph analytics programming models.

8.1 Rewriting Programs to Conform to Scalability Principles

One of the most important arguments for transactional memory (TM) is that it simplifies the writing of scalable parallel programs because it gives programmers the concurrency benefits of fine-grain locking without requiring them to write fine-grain locking code, which is usually difficult to debug, port and maintain. TM has been evaluated using mostly microbenchmarks, such as implementations of traditional data structures like stacks and trees,

and benchmark suites, such as the STAMP benchmarks (Cao Minh et al., 2008), that use these data structures. Evaluations using these benchmarks have focused mostly on small-scale¹ systems, but the speedups obtained for the STAMP benchmarks are very limited on all existing TM systems. For example, with 64 threads on the IBM Blue Gene/Q, the median speedup over sequential code is 4.1X on a state-of-the-art software transactional memory (STM), and the median speedup is 1.4X using the Blue Gene hardware transactional memory (HTM).

Are these scalability issues due to the benchmarks or existing TM implementations? This section shows that the scalability issues of STAMP can be addressed by using the right data structures and parallelization primitives. By systematically modifying STAMP to follow the disjoint access (Principle 4.1) and virtualization (Principle 4.2) principles and exploiting the now simpler access patterns, which permit a simpler conflict detection scheme, the median improvement over STAMP with 64-thread runs of a Blue Gene/Q machine is 4.4X (min: 0.19X, max: 37X, geomean: 3.9X). This new benchmark suite is called Stampede. These changes were not particularly difficult to implement; roughly 90% of the application code is unchanged, but they illustrate the kinds of changes that produce scalable programs. The following sections detail the changes made.

8.1.1 Applying the Disjoint Access Principle

The disjoint access principle requires that transactions accessing disjoint *logical* data structure elements should access disjoint *physical* memory locations. This is a general principle for scalability even for programs without transactions. For the STAMP benchmark suite, this is achieved by examining and changing the data structures used in each program.

Four general techniques were applied.

First, given an abstract data type (ADT), a scalable implementation was chosen. There are many different data structures that can satisfy an abstract data type, but some are

¹A few studies have investigated transactions for very specialized code patterns such as decoupled software pipelining (Kim et al., 2010) or microbenchmarks (Bocchino et al., 2008).

	Original	Scalable Alternative
bayes	linked-list	workstealing scheduler
genome	hashtable	reduction of hashtables
kmeans	shared-counter	workstealing scheduler
intruder	dynamic buffer	per-iteration buffer
labyrinth	growable array	workstealing scheduler
ssca2		
vacation	red-black tree	hashtable
yada	eager root update; linked-list	lazy update; workstealing scheduler

Figure 8.1: Original STAMP data structures and their scalable alternatives

more scalable than others. For instance, both a red-black tree and a hashtable can be used to implement a map, but a hashtable is more scalable because operations on different keys typically access different portions of the hashtable. Accesses to different keys in a red-black tree will have overlapping accesses on shared path from the root.

Second, for a given behavior, a scalable ADT was chosen. Supporting certain combinations of operations may be more scalable than others. For instance, a counter that supports concurrent modification and provides access to intermediate values requires more communication than a counter that does not. The latter can be implemented using a reduction tree while the former requires read-modify-write updates.

Third, scalable memory allocation patterns were chosen. Memory allocation is a common scalability bottleneck. Allocating new pages may be serialized in the operating system, and techniques for recycling and coalescing memory regions requires communication between threads. To avoid contention overheads, the Galois memory allocator was used ([Section 5.1](#)).

Fourth, transactions were assumed to have exclusive ownership of the data they access. Under the disjoint access principle, most transactions should access disjoint memory regions, so a strengthening of all accesses to be exclusive should not impact performance too much, especially when applying the scheduling optimizations available under the virtu-

alization principle (see [Section 8.1.2](#)). Tracking exclusive accesses enables a low-overhead, exclusive conflict detection policy (see [Section 5.2](#)).

Although the data structure analysis is manual, the code changes required were small because in many cases one ADT implementation was simply swapped for another. [Figure 8.1](#) shows the original STAMP data structures and their scalable alternatives. Some of the important changes are described below.

In four applications (bayes, kmeans, labyrinth, yada), a sequential data structure with transactions is used to implement a work-stealing scheduler, but it is more efficient to use a data structure designed from the ground up to be a scheduler than to use a shared counter or linked-list.

In genome, a central hashtable is used to find duplicate strings. Since the goal is to produce a hashtable without duplicates, a more scalable alternative is to create a hashtable for disjoint subsets of strings and then merge them in a reduction tree.

In intruder, dynamically allocated buffers are used to communicate values between processing pipeline stages. These allocations can be replaced with per-iteration allocation, which is highly scalable when downstream pipeline stages are nested in the current stage. The vacation application uses a red-black tree to maintain database relations, but a more scalable implementation of the same abstract data type is a hashtable.

Finally, the yada application maintains a pointer to a mesh element to verify connectivity of the final mesh. Each transaction checks if it is removing the element referenced by this pointer, and if so, it updates the pointer to another element. Since this pointer is used only during verification, it is possible to simply search for a valid element just before the verification step rather than eagerly updating the pointer during a transaction.

8.1.2 Applying the Virtualization Principle

The virtualized transaction principle requires that transactions be decoupled from threads and schedules. STAMP programs violate this principle in two ways. First, since STAMP

programs use threads directly, transactions are tied to the thread that issued them. Second, transactions issued by a thread are processed in-order, which unnecessarily restricts the space of possible schedules. A thread cannot execute its next transaction until its current transaction has committed.

Parallel programs can be written in a form that separates scheduling from the core computational operator like in the operator formulation. Given this form, the most natural granularity for a transaction is the complete execution of the operator or equivalently, an iteration of a parallel loop. These are called *loop-based transactions* in this chapter. Loop-based transactions are naturally virtualized. The mapping of transactions to threads is implicit, and the iteration space of the loop gives the entire set of transactions to execute.

In STAMP programs, transactions can appear anywhere, and in fact, one optimization is to reduce the granularity of transactions to be smaller than the computational operator. These are called *fine-grain transactions* in this chapter. Fine-grain transactions are akin to fine-grain locking. They can significantly improve performance by reducing transactional state, but they require sophisticated reasoning to ensure correct behavior. For instance, one must prove facts of the form: “atomic { A; B }” is equivalent to “atomic { A }; atomic { B }.” In contrast, loop-based transactions are easier to reason about. The behavior of a parallel loop with coarse-grain transactions is equivalent to executing iterations in some sequential order. They can also have performance benefits over fine-grain transactions. For instance, since transactions are the same as parallel tasks, many of the techniques used to schedule parallel tasks like workstealing can be directly applied to schedule transactions.

Fine-grain transactions can be made into loop-based ones by applying a continuation-passing transformation to divide loop iterations into units matching the fine-grain transaction boundaries (see [Figure 8.2](#)). These new coarse-grain transactions will have the same granularity as the original fine-grain ones, but unlike the originals, there is overhead from forming the continuation and a possible loss of data locality due the possible rescheduling of two previously sequentially composed transactions.

```

1 foreach Item i in I:
2   int v
3   atomic { b = fn0(i) }
4   atomic { fn1(i, v) }
5 }

```

(a) Original fine-grain transaction program

```

1 struct Task { Item i; int v; int s; }
2
3 foreach Task t in T:
4   atomic:
5     switch t.s:
6       case 0:
7         t.v = fn0(t.i); t.s = 1; T.push(t);
8         break;
9       case 1:
10        fn1(t.i, t.v);
11        break;

```

(b) New loop-based transaction program

Figure 8.2: Example of converting fine-grain transactions into loop-based ones

If expert parallel programmers still want to use fine-grain transactions, they can mimic most of their effect in loop-based code by annotating particular reads or writes in an iteration as protected by the transaction or not. In Stampede programs, this is accomplished by a special API call before shared data accesses.

Virtualization provides an opportunity for rescheduling work to improve performance. Figure 8.3 shows two possible schedulers for transactions. The first scheduler (Figure 8.3a) uses static work assignment and immediately retries aborted transactions. This is roughly the behavior of all the transactions in STAMP programs. The second scheduler (Figure 8.3b) uses workstealing to initially distribute transactions and uses a *serialization tree* to guarantee forward progress by gradually serializing aborted transactions on fewer and fewer threads. If thread n executes transaction t and it aborts, thread n tries the activity again (in case the conflict is transient), and if transaction t aborts again, thread n gives transaction t to thread $\lfloor n/2 \rfloor$ to execute, and so on.

The second scheduler is only possible when transactions are virtualized and is but

```

1 void execute():
2   for int i in range(begin, end):
3     while true:
4       task[i].execute()
5       if !task[i].aborted:
6         break

```

(a) Scheduling for unvirtualized transactions

```

1 ThreadLocal<int> myId
2 TerminationDetection term
3 WorkstealingQueue q
4 Queue aborted[numThreads]
5
6 void execute():
7   Task t
8   while !term.allDone():
9     if (t = q.pop())
10      || (t = aborted[myId].pop()):
11      term.notDone()
12      t.execute()
13      if t.aborted:
14        aborted[myId].push(t)
15      while (t = aborted[myId].pop()):
16        t.execute()
17        if t.aborted:
18          aborted[myId/2].push(t)
19    else:
20      term.done()

```

(b) Improved scheduling for virtualized transactions using workstealing and serialization tree

Figure 8.3: Scheduling virtualized transactions

one of many schedulers that could be used. CAR-STM (Dolev et al., 2008) previously investigated serializing aborted transactions in the context of STM, but it technically breaks the TM programming model by potentially scheduling a transaction on a thread that is different than the thread that reached the transactional code block. More importantly, this work did not investigate improving the program data structures; scalable programs result when both the virtualization and disjoint access principles are followed.

8.1.3 Evaluation

Starting with the STAMP benchmark suite (v0.9.10), each program was modified according to the disjoint access and virtualization principles to produce the **Stampede** benchmark suite. To give a sense of the change from STAMP to Stampede, STAMP has 41586 lines of code. Stampede has 43563 lines of code, of which 38677 lines (88%) are exactly the same as STAMP. The bulk of the changes are due to changing the way STAMP data structures iterate through elements to support the Stampede memory allocator and shifting code to conform to loop-based transactions. Of the changes, only 326 lines are due to new code; in this case, these are new data structures that do not have any analogue in the original STAMP code. This is a reasonable amount of effort as even applying TM to an existing threaded program using compiler support requires some level of human intervention and significant non-local changes (Ruan et al., 2014).

Each application was run with its largest input on two machines: an Intel Xeon E7-4860 (Westmere) machine, which has four 10-core processors², and a single Blue Gene/Q compute node. The Blue Gene/Q compute node has sixteen cores. Each core has four hardware execution contexts, which gives a total of 64 execution contexts per node. Westmere does not support hardware transactional memory, but Blue Gene/Q does. It is enabled by annotating programs with compiler pragmas that denote transaction boundaries. Transactional execution is achieved via the coordination of the compiler, kernel and TM runtime.

On Blue Gene/Q, programs are compiled with the IBM XLC compiler (v12.1) with the following compiler options: `-O -qsmpt=noopt3 -qalias=noansi`. On Westmere, programs are compiled with GCC (v4.8.1) with `-O3`.

Figure 8.4 compares the performance of STAMP programs using an STM system (**STAMP+STM**) with Stampede programs using several possible execution systems. There

²Although this machine has 40 cores, the experiments only run with threads in powers of two due to a restriction in the STAMP benchmark suite

³This option disables auto-parallelization in the XLC compiler; the other standard compiler optimizations are not affected.

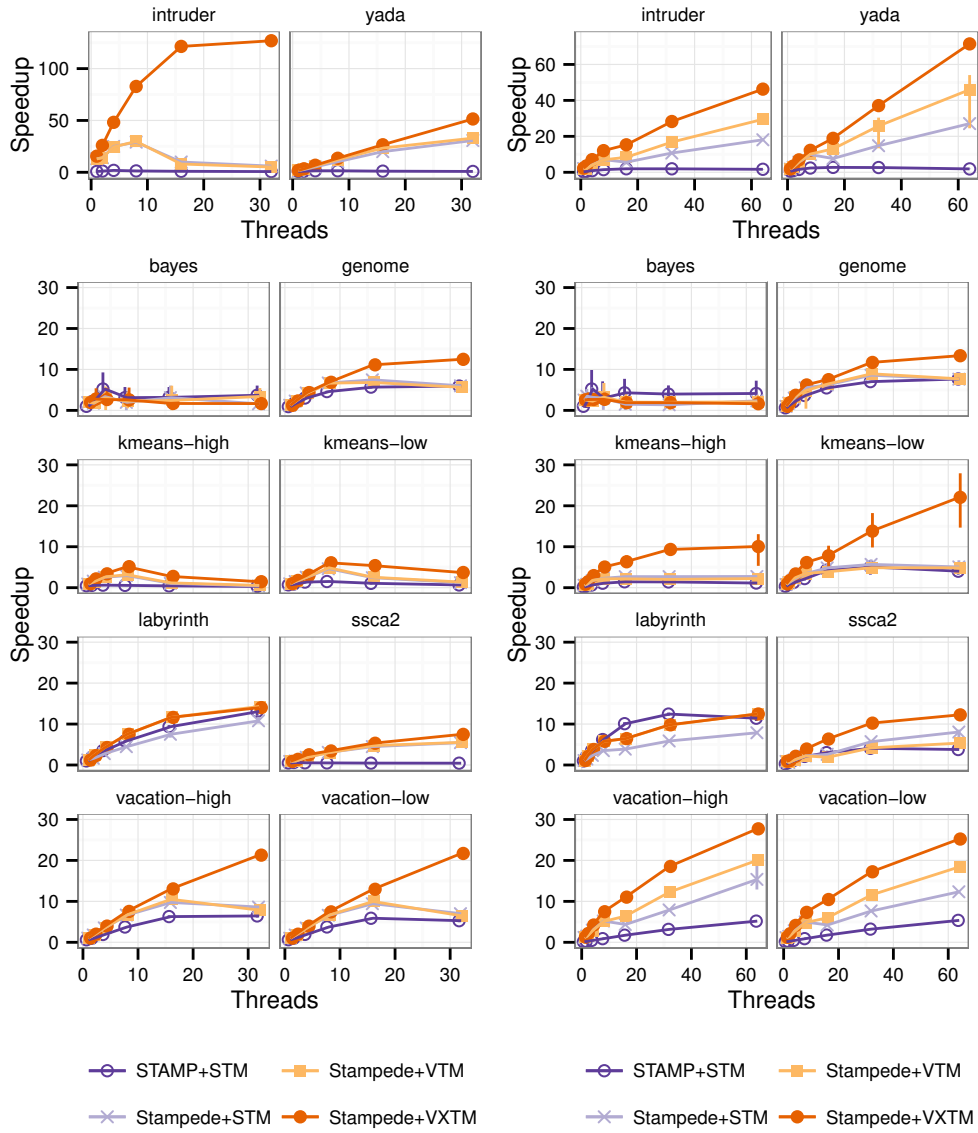


Figure 8.4: Comparison of programs on two architectures using improved data structures and scheduling (Stampede) and original STAMP programs. Speedup relative to STAMP sequential baseline (see Figure 8.5). Points at mean value of at least 5 runs. Vertical bars indicate min and max observed values.

Application	Xeon Westmere		Blue Gene/Q	
	Speedup	Time	Speedup	Time
bayes	1.94	7.07	2.58	67.79
genome	1.23	6.84	0.98	60.10
intruder	15.53	1.81	2.31	43.46
kmeans-high	0.96	2.78	0.84	14.78
kmeans-low	0.98	15.42	0.93	78.26
labyrinth	1.21	63.61	1.09	524.00
ssca2	0.89	7.82	0.91	29.99
vacation-high	1.03	25.19	1.26	70.67
vacation-low	1.02	19.39	1.23	51.77
yada	1.74	7.05	1.87	54.77

Figure 8.5: Speedup over STAMP sequential baseline and execution times in seconds for Stampede+VXTM programs with one thread

are many software transactional memory systems to choose from. This comparison uses TinySTM (Felber et al., 2008) (v1.0.4) as a representative system. It supports several different policies for conflict detection and resolution. It is used in its default configuration, which is encounter-time locking with write-back of transactional state on commit.

To run Stampede with HTM (**Stampede+HTM**), loop bodies are marked with compiler pragmas to indicate a transaction. The transaction only includes the loop body itself and does not include scheduling code. To run with STM (**Stampede+STM**), transactions boundaries are marked as in the HTM case, and shared values (i.e., exclusive locations) are marked as transactional variables. To access their values, the program writes the currently executing thread id to an owner field, relying on the underlying STM system to detect conflicts.

Although Stampede programs have virtualized transactions, Stampede with HTM or STM will behave as if transactions are not virtualized because the underlying TM is not aware of the virtualization and it will execute transactions like in Figure 8.3a. To measure the impact of virtualization, Stampede transactions are modified to abort if they are re-

executed by monitoring a non-transactional memory location. The aborted transactions are placed on a serialization tree and processed according [Figure 8.3b](#). This variant is called **Stampede+VTM**. A similar modification can not be applied to Stampede+HTM because the HTM provides strong atomicity and does not indicate if a transaction aborts.

Finally, **Stampede+VXTM** schedules virtualized transactions using workstealing and uses a serialization tree like Stampede+VTM, but instead of using an off-the-shelf STM, it uses the exclusive locking implementation described in [Section 5.2](#).

Overall, [Figure 8.4](#) shows that Stampede programs outperform the corresponding STAMP+STM programs with Stampede+VXTM being the fastest overall. On Westmere, the median speedup with 32 threads over sequential code is 2.25X for STAMP+STM, 5.98X for Stampede+STM and 13.24 for Stampede+VXTM. On Blue Gene/Q, the median speedup with 64 threads is 3.6X for STAMP+STM, 6.7X for Stampede+STM and 12.76X for Stampede+VXTM.

Two exceptions to this trend are bayes and labyrinth. The behavior of bayes is highly variable, but the mean STAMP+STM time is usually faster than the Stampede time across threads. The exclusive access policy taken by Stampede programs is a performance detriment on bayes. [Section 8.1.3](#) examines this effect further. For labyrinth on Westmere, STAMP+STM is faster than Stampede+STM, and on Blue Gene/Q, it is even faster than Stampede+VXTM for some number of threads. Here, the STAMP program implements a lazy transaction validation scheme that is not available in the Stampede program.

Another general trend is that Stampede+VTM is typically the same or faster than Stampede+STM. The difference between the two indicates the performance opportunity of virtualization. On Westmere, the difference is negligible for most programs, but on Blue Gene/Q, intruder, yada, vacation-high and vacation-low see significant performance gains with Stampede+VTM. Moreover, virtualization is an enabling transformation that facilitates optimizations like exclusive conflict detection. For example, [Section 8.1.3](#) shows how applying exclusive conflict detection to STAMP programs directly can produce dramatically

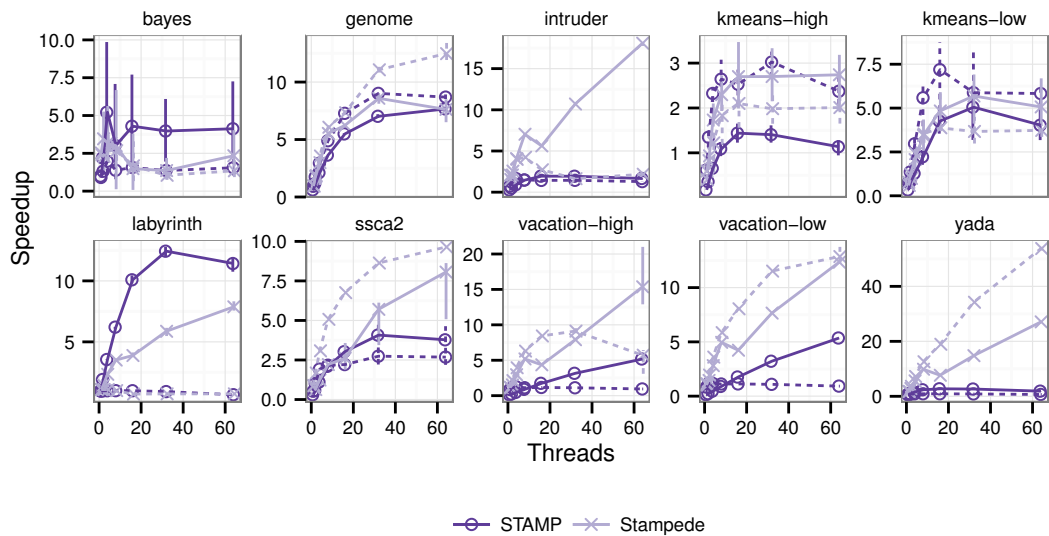


Figure 8.6: Speedup of STAMP and Stampede programs with STM (solid lines) and with HTM (dotted lines) on Blue Gene/Q

worse performance.

Figure 8.5 gives the absolute running times and the speedup of Stampede+VXTM programs with one thread over the STAMP sequential baseline, which is the application without any support for TM or parallelism. For most applications, the performance of Stampede+VXTM is close to the original sequential STAMP programs. The main exception is intruder. The loop-based Stampede version has significantly better locality than the pipeline-based STAMP version.

Hardware Transactional Memory

Figure 8.6 shows the speedup of STAMP and Stampede programs with an STM or HTM. A comparison of STAMP programs using STM and HTM on the Blue Gene/Q was previously done by Wang et al. (Wang et al., 2012a). The results here for the HTM and STM programs broadly match their reported results with one exception. Wang et al. report a maximum speedup of 12X for vacation-low and 16X for vacation-high.

The performance of HTM on vacation-high (and on vacation-low) strongly depends on the memory allocator used. The results in [Figure 8.6](#) use standard malloc while the results of Wang et al. use a different memory allocator that preallocates thread-local pools but does not track freed memory, causing a memory leak. The similarity of results between the two different evaluations excluding the vacation programs suggests that improving memory allocation by itself does not typically impact performance. [Section 8.1.3](#) shows results when switching the ad-hoc pooled allocator for a scalable malloc replacement. Only the yada benchmark was significantly improved (maximum speedup of 12X on Blue Gene/Q) compared to using standard malloc.

For STAMP programs, STAMP+HTM is faster than STAMP+STM for three out of ten benchmarks (genome, kmeans-high and kmeans-low), and for most STAMP benchmarks, the performance difference between STAMP+HTM and STAMP+STM is not more than a factor of two. The large exception is labyrinth where the large working set exhausts the transactional state of the hardware, which then serializes execution. For Stampede programs, Stampede+HTM is faster than Stampede+STM for four benchmarks (genome, ssa2, vacation-low and yada).

The existence of benchmarks with a large difference between STAMP+HTM and Stampede+HTM performance like ssa2, vacation-high, vacation-low and yada show that some of the poor performance with HTM can be alleviated by starting with scalable programs.

Commit Ratios

[Figure 8.7](#) shows the commit ratio of STAMP and Stampede programs. STAMP+STM has a higher commit ratio than Stampede+VXTM for only three benchmarks (bayes, vacation-high, vacation-low), which illustrates how the application of the disjoint access and virtualization principles can improve the effectiveness of transactional programs. Virtualization, that is moving from Stampede+STM to Stampede+VTM, usually improves the commit ra-

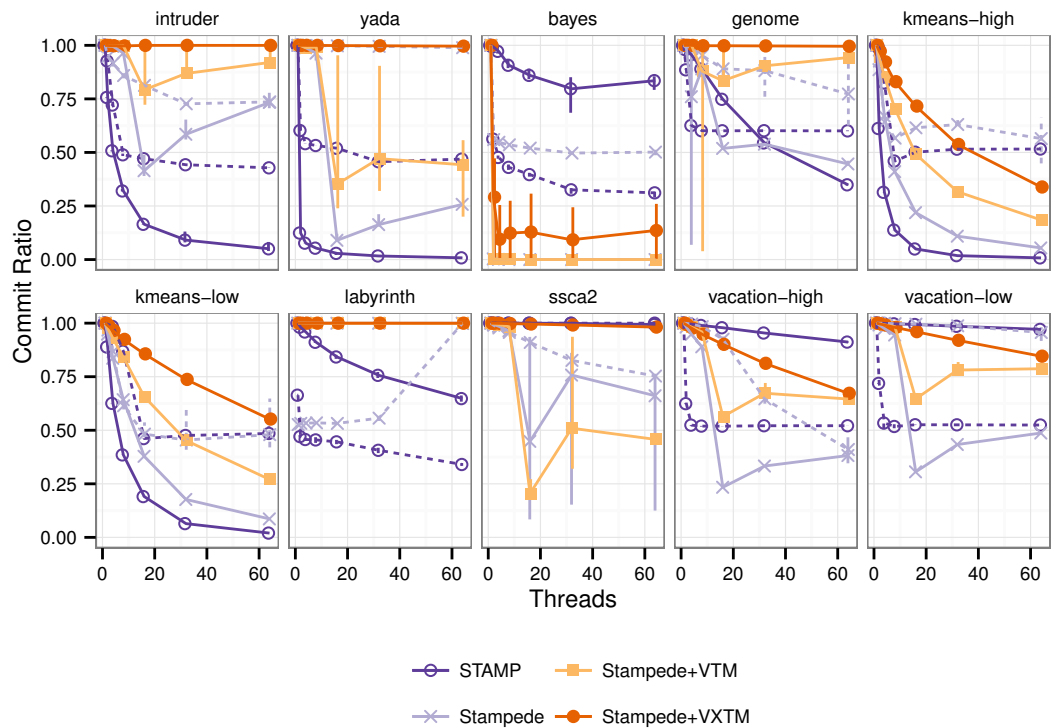


Figure 8.7: Commit ratios of STAMP+STM (solid line), STAMP+HTM (dotted line), Stampede+STM (solid line), Stampede+HTM (dotted line), Stampede+VTM and Stampede+VXTM programs on Blue Gene/Q

tion and is one reason that Stampede+VTM programs are faster than Stampede+STM ones.

Typically, moving from Stampede+VTM to the exclusive locking scheme of Stampede+VXTM also improves the commit ratio. This is because Stampede+VXTM detects conflicts at the memory word granularity and does not suffer from false sharing conflicts like many STMs do.

Even with the less precise exclusive conflict detection, the commit ratio of Stampede+VXTM programs exceeds that of Stampede+HTM for seven benchmarks (intruder, yada, genome, kmeans-low, labyrinth, ssc2, vacation-high). This suggests that some amount of capacity or false conflicts is inhibiting the performance of HTM programs. This is certainly true for labyrinth, which has a large working set that exhausts the capacity of the

HTM. Even when the commit ratio of Stampede+HTM is equal to or greater than Stampede+VXTM (bayes, kmeans-high and vacation-low), the performance is worse than Stampede+VXTM, suggesting that the cost of implementing more precise conflicts outweighs the benefit of having less conflicts.

The flattening of the commit ratio around 50% for Stampede+HTM is due to the fact that, during high conflict situations, the HTM runs a transaction once speculatively, and on abort, it is run in “irrevocable mode,” which gives an expected 50% commit ratio.

There are three general reasons why one conflict detection implementation performs differently than another on the same application.

1. The schedule produced by one implementation creates more total work than another implementation. This can happen in the bayes application because, depending on the scheduling, different operations will be attempted, which changes the total amount of work done.
2. Assuming that the total amount of work is the same between program executions, the number of conflicts can vary due to different definitions of a conflicts (e.g., word or cache line-based detection).
3. And finally, even if one implementation produces less conflicts than another, the cost of implementing more precise conflict detection may outweigh the benefit of the lower number of conflicts.

The first reason is strongly dependent on the application, but the latter two reasons are due to general properties of conflict detection systems.

Conflict detection in Stampede+VXTM only allows exclusive access to locations, so a read before a write will always be treated as a conflict for the writer. Both HTM and STM systems allow for more precise classification of reads and writes, but the granularity of detection varies. The HTM on the Blue Gene/Q detects conflicts at cache line granularity, which may result in false conflicts when state from two transactions shares the same cache

line. TinySTM is word-based, but since TinySTM is used only to track exclusive locations, which are cache-line aligned, the detection granularity of Stampede+STM is the same as Stampede+VXTM.

TM systems can suffer from *capacity conflicts* when there is no conflict but the system reports one due to limitations in the implementation. For instance, TM systems may use Bloom filters or address hashing to improve the performance of conflict detection but that produces some number of false positives. HTM systems have a finite amount of storage for transactional state, and exceeding that limit is treated as a conflict. The Blue Gene/Q HTM is further hampered by the fact that STMs allow control over which memory locations are transactional, but in the Blue Gene/Q HTM, all memory accesses within a transaction are considered part of the transactional state.

With respect to the cost of implementing conflict detection, conflict detection in Stampede+VXTM is very simple; it uses exclusive locking. General TM implementations are more complicated because they support state rollback and more precise read-write conflicts. For instance, to support state rollback, the default configuration of TinySTM uses a write-back cache, which means that accesses of modified state must go through a level of indirection. To support lightweight read transactions, TinySTM uses a time-based mechanism that requires shared access to a global clock. The Blue Gene/Q HTM implementation stores transaction metadata in the L2 cache, and transactions must either bypass or flush the L1 cache to ensure consistent updates of metadata (Wang et al., 2012a).

A question to ask is whether commit ratios are correlated with performance. Let efficiency be speedup normalized to the number of threads. There is a correlation between the change in commit ratio of Stampede+HTM and Stampede+VXTM programs and the change in efficiency of the two, $\beta = 0.63$, $t(58) = 4.58$, $p < 0.001$. The change in commit ratio also explained a significant portion of the variance in changes in efficiency, $R^2 = 0.25$, $F(1, 58) = 20.98$, $p < 0.001$. The same does not hold when comparing the change in commit ratio between Stampede+VTM and Stampede+VXTM programs, $t(58) = 0.630$,

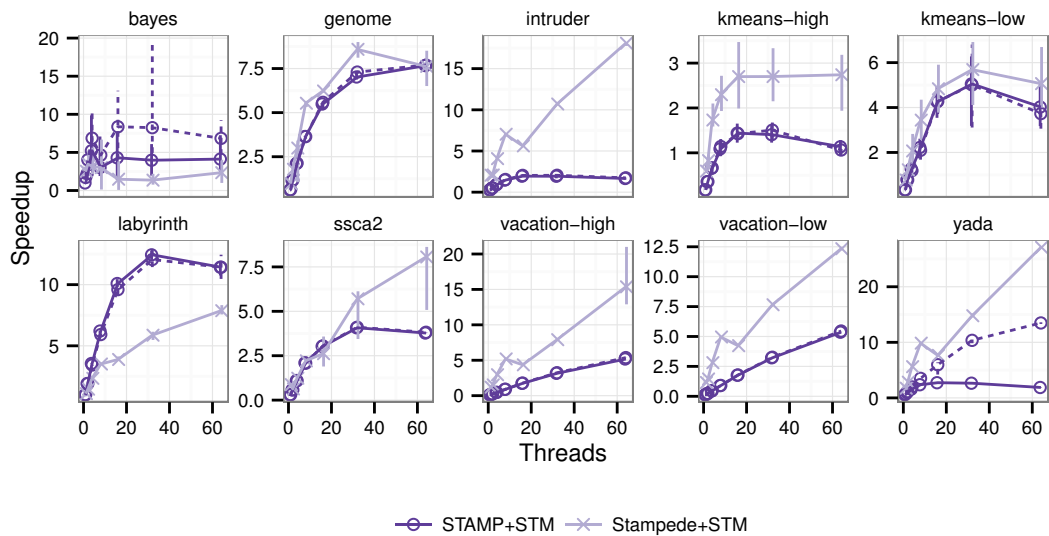


Figure 8.8: Performance of STAMP programs without (solid line) and with scalable malloc (dotted line) on Blue Gene/Q. Stampede+STM results have been included for reference.

$p > 0.1$ or between Stampede+STM and Stampede+VXTM programs, $t(58) = 1.923$, $p > 0.1$, suggesting that their performance is mostly related to the implementation of conflict detection rather than the number of conflicts detected.

Scalable Malloc

One of the transformations mentioned in Section 8.1.1 is to use a scalable memory allocator. Considering that applying the disjoint access principle might require refactoring an existing program, is it possible to achieve similar results by just linking in an off-the-shelf scalable memory allocator? Figure 8.8 shows the performance of STAMP benchmarks with and without TCMalloc (Ghemawat and Menage, 2014), a scalable malloc replacement. Only the yada benchmark is significantly improved by using the scalable malloc alternative, which suggests that the program refactoring done in Stampede may be necessary.

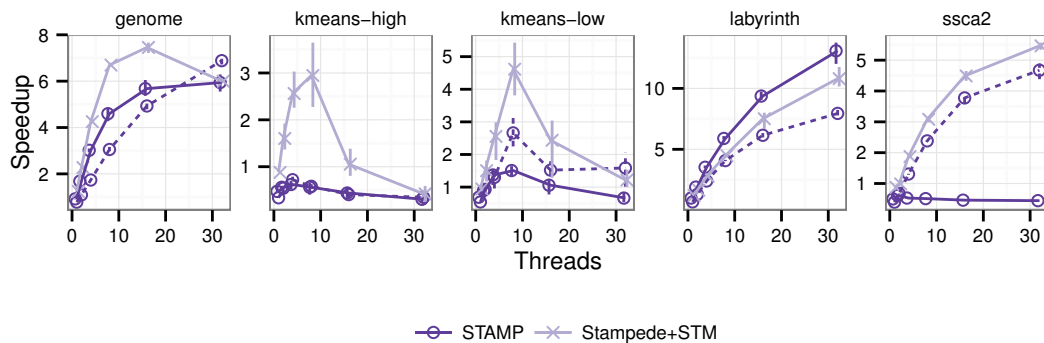


Figure 8.9: Performance of STAMP programs using STM (solid line) and using XTM (dotted line). Stampede+STM results have been included for reference.

XTM

Figure 8.9 gives the performance results of STM and using the exclusive locking (XTM) directly on STAMP programs. In these runs, when a transaction aborts, it is immediately retried by the currently executing thread.

On two of the ten benchmarks (kmeans-low and ssca2), STAMP+XTM performs as well as or better than STM. These are instances where the benefit of lower overhead conflict detection outweighs the need for higher concurrency. On three benchmarks (i.e., genome, kmeans-high, labyrinth), STAMP+XTM is worse than STAMP+STM.

On the other five benchmarks not shown in the figure (i.e., bayes, intruder, vacation-high, vacation-low, yada), the performance of STAMP+XTM is much worse. In these benchmarks, transactions tend to read and write the same data structure, and the number of aborted transactions grows very large, slowing the overall execution and potentially introducing livelock. Memory can also be exhausted if a transaction allocates memory and then aborts without freeing it.

In bayes, the central data structure is a graph and a workset used to schedule work. In vacation-high and vacation-low, the central data structure is a red-black tree used to store relational data entries. In yada, the central data structure is a root pointer, which points to

some element of a mesh, and is used to verify the correctness of the benchmark.

As noted above, these kinds of program behaviors are unlikely to scale irrespective of transactional memory. Highly concurrent transactional memory systems can ameliorate the scalability problem, but solving it requires changing the program itself, and [Section 8.1.1](#) discusses how these centralized access patterns can be addressed by modifying data structures.

Other Benchmarks

These scalability principles apply to other benchmarks as well. This section gives preliminary results on the PARSEC suite. The PARSEC suite ([Bienia et al., 2008](#)) (v2.1) is a collection of parallel programs that cover a number of parallelization techniques. However, none of the programs use TM, and as reported by others, most of the programs are data-parallel ([Best et al., 2011](#)). One that is not is canneal, which simulates cache-aware annealing to optimize routing costs.

Three variants of this program were evaluated. The first is the original PARSEC program that uses fine-grain locking, PARSEC+FGL. The second is a manual transformation to use TinySTM instead of locks, PARSEC+STM. The third is a new program that follows the disjoint access and virtualization principles and uses exclusive locking, NewPARSEC+VXTM. To satisfy disjoint access, the only change made was to allocate the main graph data structure in a NUMA-interleaved fashion to avoid congestion on physical memory controllers. Satisfying virtualization was simply a matter of converting explicit threading to parallel loops. On Westmere with 32 threads and using the largest input size, PARSEC+FGL is 22.3X faster than with 1 thread. On 32 threads, PARSEC+STM is 7% faster than PARSEC+FGL, and NewPARSEC+VXTM is 16% faster. These preliminary results support the applicability of our scalability principles beyond STAMP and programs initially written to use transactions.

8.2 Parallel Programming Models

The previous section described how programs can be rewritten to be more scalable by following the disjoint access and virtualization principles. If programs are rewritten, which parallel programming model should they use? This section evaluates the performance of several parallel programming models or domain-specific languages (DSLs) for graph analytics.

Graph analytics DSLs usually constrain programmers to use a subset of features described in [Section 2.2](#).

GraphLab ([Low et al., 2010](#)) is a shared-memory programming model for topology or data-driven computations with autonomous or coordinated scheduling, but it is restricted to *vertex programs*. A vertex program has a graph operator that can only read from and write to the immediate neighbors of the active node. There are several priority scheduling policies available, but the implementation of the priority scheduler is very different from the one used in the Galois system (see [Section 6.1.2](#)).

PowerGraph ([Gonzalez et al., 2012](#)) is a distributed-memory programming model for topology or data-driven computations with autonomous or coordinated scheduling, but it is restricted to *gather-apply-scatter* (GAS) programs, which are a subset of vertex programs. Graphs are partitioned by edge where the endpoints of edges may be shared by multiple machines. Values on shared nodes can be resolved with local update and distributed reduction. On scale-free graphs, which have many high-degree nodes, this is a useful optimization to improve load balancing. PowerGraph supports autonomous scheduling, but the scheduling policy is fixed by the system and users cannot choose among autonomous policies.

GraphChi ([Kyrola et al., 2012](#)) is a shared-memory programming model for vertex programs that supports out-of-core processing when the input graph is too large to fit in

[‡]This section draws from ([Nguyen et al., 2013](#)), where the evaluation of Galois versus graph analytics DSLs was originally reported.

memory. GraphChi relies on a particular sorting of graph edges in order to provide I/O-efficient access to both the in and out edges of a node. Since computation is driven by the loading and storing of graph files, GraphChi only provides coordinated scheduling.⁴

Ligra (Shun and Blelloch, 2013) is a shared-memory programming model for vertex programs with coordinated scheduling. A unique feature of Ligra is that it switches between push and pull-based operators automatically based on a user-provided threshold.

Linear algebra formulations (Kepner and Gilbert, 2011) are a methodology for exploiting the substantial expertise in parallelizing dense matrix kernels and sparse matrix-vector multiply operations to implement graph algorithms. The basic idea is to express graph operations as matrix operations over some algebraic semiring. For instance, in standard linear algebra, operations are over the field $(\mathbb{R}, +, \cdot)$. The shortest path can be found by iterative application of matrix-vector multiply over the semiring $(\mathbb{R} \cup \infty, \min, +)$.

Section 2.3 shows programs can use rich programming models. However, most existing graph DSLs have restricted themselves to supporting a simple programming model, and they do not support the more complex features such as autonomously scheduled, data-driven computation. Next, we discuss how these simpler models can be layered on top of the more expressive Galois system.

8.2.1 Other Domain Specific Languages in Galois

Graph analytics DSLs such GraphLab, GraphChi and Ligra can be simply layered on top of Galois. This section describes how to implement features of the GraphLab, GraphChi, and Ligra APIs on top of the Galois system. The Galois implementations of GraphLab and Ligra are called **GraphLab-g** and **Ligra-g** respectively. Also, to demonstrate the ease with which new DSLs can be implemented, a DSL called **LigraChi-g** is developed that combines features of the Ligra and GraphChi systems. Figure 8.10 gives the approximate lines of code required to implement these features on top of the Galois system.

⁴GraphChi takes a program that could be autonomously scheduled but imposes a coordinated schedule for execution.

Feature	LoC
Vertex Programs	0
Gather-Apply-Scatter (synchronous engine)	200
Gather-Apply-Scatter (asynchronous engine)	200
Out-of-core	400
Push-versus-pull	300
Out-of-core + Push-versus-pull (additional)	100

Figure 8.10: Approximate lines of code for each DSL feature

Vertex Programs These are directly supported by Galois. Granularity of serializability can be controlled through the use of Galois data structure parameters. For example, to achieve the GraphLab edge consistency model, a user can enable logical locks when accessing a vertex and its edges but not acquire logical locks when accessing a neighboring node.

Gather-apply-scatter The PowerGraph implementation of GAS programs has three different execution models: coordinated scheduling, autonomous scheduling without consistency guarantees, and autonomous scheduling with serializable activities. The Galois implementation of PowerGraph is called **PowerGraph-g**. The two autonomous scheduling models can be implemented in Galois by concatenating the gather, apply and scatter steps for a vertex into a single Galois operator and either always or never acquiring logical locks during the execution of the operator.

The coordinated scheduling model can be implemented by a sequence of loops, one for each phase of the GAS programming model. The main implementation question is how to implement the scatter phase, since it must handle the case when multiple nodes send messages to the same neighbor. The simplest implementation is to accumulate all messages for the same node in place, using a lock to protect concurrent updates. In practice, this does not scale well for many applications. Instead, one could gather all messages for a node in a list and have the receiving node reduce the list during the subsequent initialization phase,

but this requires significant memory allocation for applications that send many messages to the same node. The implementation used in PowerGraph-g is to have per-package message accumulation, protected by a spin-lock. The receiver accumulates the final message value by reading from the per-package locations.

PowerGraph supports dividing up the work of a single gather or scatter operation for a node among different processing units, but PowerGraph-g does not yet have this optimization.

Out-of-core The GraphChi implementation of out-of-core processing for vertex programs uses a carefully designed graph file format to support I/O-efficient access to both the incoming and outgoing edge values of a node. For the purpose of understanding how to provide out-of-core processing using general reusable components, this section focuses on supporting only a subset of GraphChi features.

The implementation of out-of-core processing is based on incremental loading of the compressed sparse row (CSR) format of a graph and the graph's transpose. The transpose graph represents the incoming edges of a graph and stores a copy of the edge values of the corresponding outgoing edges. Since these values are copies, it does not support updating edge values like GraphChi does; however, none of the applications described in this chapter require updating edge values. To reduce the waiting time on I/O operations, loading portions of the graph is double-buffered.

Push-versus-pull The push-versus-pull optimization in Ligra can be implemented as two vertex programs that take an edge update rule and perform either a push or pull-based traversal according to some threshold of active nodes to all the nodes in the graph. The Galois implementation of Ligra is called **Ligra-g**. It uses the same threshold heuristic as Ligra. In order to perform this optimization, the graph representation must store both incoming and outgoing edges.

	GraphLab	Ligra	PowerGraph	Galois
bfs	A; prioritized	C; push/pull	C	C/A; push/pull
cc	label propagation	label propagation	label propagation	union-find
dia	pseudo-peripheral	k -BFS	probabilistic counting	pseudo-peripheral
pr	push	push	push	pull
sssp	A; prioritized	C; no priority	C; no priority	A; prioritized
bc		C; push/pull		A; pull

Figure 8.11: Summary of application differences due to varying support for programming model features: coordinated (C), autonomous (A), coordinated and autonomous (C/A) scheduling.

Push-versus-pull and out-of-core The push-versus-pull optimization generates a vertex program, and it is possible to apply the out-of-core processing described above to the new program. The DSL that combines both these optimizations is called **LigraChi-g**. This highlights the utility of having a single framework for implementing DSLs.

8.2.2 Evaluation

This section compares the performance of applications in the Ligra, GraphLab (v1), PowerGraph (v2.1) and Galois (v2.2) systems. The main evaluation machine is a four processor Intel (E7-4860) machine with each processor having ten cores. The machine has 128 GB of RAM.

PowerGraph is a distributed-memory implementation, but it supports shared-memory parallelism within a single machine. Ligra, GraphLab and Galois are strictly shared-memory systems. Ligra requires the Cilk runtime, which is not yet available with the GCC compiler. Ligra applications are compiled with the Intel ICC 12.1 compiler. All other applications are compiled with GCC 4.7 with the -O3 optimization level.

All runtimes are an average of at least two runs. For out-of-core DSLs, the runtimes include the time to load data from local disk. For all other systems, this time is excluded.

Graph analytics algorithms can use rich programming models, but most existing

graph DSLs support only a simple set of features. In light of these restrictions, different applications are used to solve the same analytics problems. The following is a brief description of the application differences and is summarized in [Figure 8.11](#). Most of the implementations are provided by the DSL systems themselves, except for GraphLab, for which most implementations were developed from scratch.

- **Breadth-first search (BFS)**. The Galois BFS application blends coordinated and autonomous scheduling. Initially, the application uses coordinated scheduling of the push and pull-based operators. After a certain number of rounds of push-based traversals, the application switches to prioritized autonomous scheduling. The priority function favors executing nodes with smaller BFS numbers. The Ligra application uses coordinated scheduling and switches between push-based and pull-based operators automatically. Since PowerGraph does not provide a BFS application, one was created based on its SSSP application. GraphLab does not provide a BFS application, so one was created based on prioritized autonomous scheduling.
- **Connected components (CC)**. Galois provides a parallel connected components application based on concurrent union-find. PowerGraph, GraphChi and Ligra include applications based on iterative label propagation. This will converge slowly if the diameter of the graph is high. GraphLab does not provide an algorithm for finding connected components; one was implemented based on the label propagation algorithm.
- **Approximate diameter (DIA)**. The Galois application is based on finding pseudo-peripheral nodes in the graph. GraphLab does not provide an application for this problem; one was created based on the pseudo-peripheral algorithm. Ligra uses k -BFS. PowerGraph uses probabilistic counting.
- **PageRank (PR)**. GraphLab, GraphChi, PowerGraph and Ligra use topology-driven push-based operators. GraphChi has both a vertex program application as well as a

	$ V $	$ E $	MB	Weighted MB
rmat24	17	268	1207	2281
rmat27	134	2141	9637	18218
twitter40	42	1469	6207	12080
twitter50	51	1963	8262	16114
road	24	58	422	653

Figure 8.12: Input characteristics. Number of nodes and edges is in millions. MB is the size of CSR representation in megabytes.

gather-apply-scatter application. The latter is used because it is slightly faster. Galois provides a pull-based PageRank application that reduces the memory overhead and synchronization compared to push-based applications.

- **Single-source shortest-paths (SSSP).** The Galois SSSP application uses the data-driven, autonomously scheduled delta-stepping algorithm, using auto-tuning to find an optimal value of Δ for a given input. GraphLab does not provide an SSSP application, so one was created based on the Galois application, using the priority scheduling available in GraphLab. PowerGraph and Ligra use Bellman-Ford, which uses coordinated scheduling.
- **Betweenness centrality (BC).** The Galois application is based on a priority-scheduled, pull-based algorithm for computing betweenness centrality. The priority function is based on the BFS number of a node. The Ligra application switches between pull and push-based operators with coordinated execution, which can have significant overhead on large diameter graphs.

Figure 8.12 summarizes the graph inputs used. The rmat24 ($a = 0.5, b = c = 0.1, d = 0.3$) and rmat27 ($a = 0.57, b = c = 0.19, d = 0.05$) graphs are synthetic scale-free graphs. Following the evaluation done by Shun and Blleloch (Shun and Blleloch, 2013), the graphs are made symmetric. The **twitter40** (Kwak et al., 2010) and **twitter50** (Cha et al., 2010) graphs are real-world social network graphs. From twitter40 and twitter50, only the

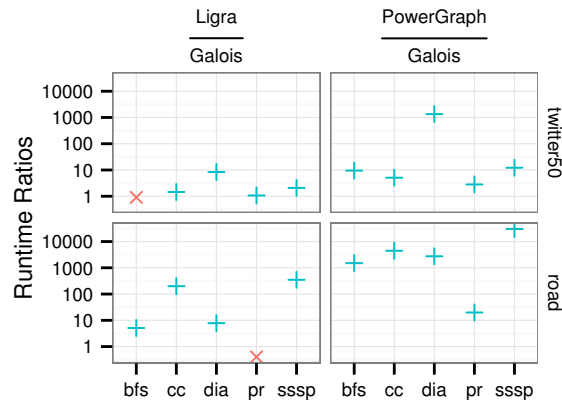


Figure 8.13: Ratio of Ligra and PowerGraph runtimes to Galois with 40 threads. Values greater than 1 (shown as blue crosses) indicate how many times faster the denominator system is than the numerator.

largest connected component is used. Finally, **road** is a road network of the United States obtained from the DIMACS shortest paths benchmark.

The road graph is naturally weighted; the weights are removed when an unweighted graph is needed. The other four graphs are unweighted. To provide a weighted input for the SSSP algorithms, unweighted graphs are given random edge weights in the range $(0, 100]$.

Figure 8.13 shows the runtime ratios of the Ligra and PowerGraph applications compared to the Galois versions on the twitter50 and road inputs for five applications. When the Galois version runs faster, the data point is shown as a cross; otherwise it is shown as an x (i.e., BFS on twitter50 and PR on road). The values range over several orders of magnitude. The largest improvements are on the road graph and with respect to PowerGraph.

Figure 8.14 shows the runtime ratios of the Ligra and PowerGraph applications compared to the Ligra-g and PowerGraph-g versions (that is, the implementations of those DSLs in Galois). The performance of Ligra-g is roughly comparable to Ligra. PowerGraph-g is mostly better than PowerGraph. This shows that much of the huge improvements in Figure 8.13 come not so much from the better implementations of the DSL in Galois *per se* but from the better programs that can be written when the programming model is rich

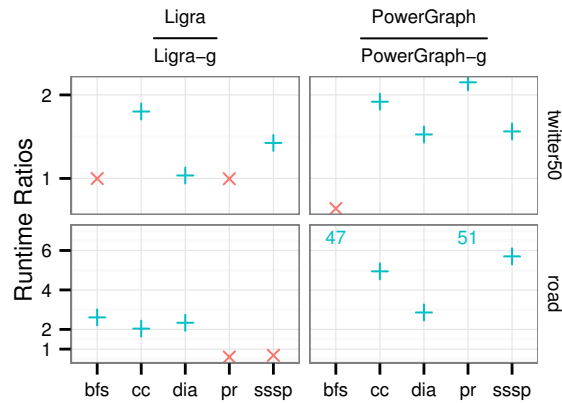


Figure 8.14: Ratio of Ligra and PowerGraph runtimes relative to Ligra-g and PowerGraph-g runtimes. Values greater than 1 (shown as blue crosses) indicate how many times faster the denominator system is than the numerator. Larger ratios shown as numbers rather than points.

enough.

Comparison between the two figures can also be illuminating. For example, most of the ratios in [Figure 8.13](#) are greater than those in [Figure 8.14](#), but one notable exception is the behavior of PowerGraph with PageRank on the road graph. The Galois improvement is about 10X while the PowerGraph-g improvement is about 50X. This suggests that the Galois application of PageRank, which is pull-based, is not as good as the push-based algorithm used by PowerGraph, on the road graph. Thus, Galois is faster than PowerGraph on PageRank because of a more efficient implementation of a worse algorithm.

The following sections dig deeper into the performance results.

Overall results

[Figure 8.15](#) gives the complete runtime results with 40 threads. The PageRank (PR) times are for one iteration of the topology-driven algorithm. The betweenness centrality (BC) times are for computing results with respect to one source node. Ligra-g and PowerGraph-g results will be discussed in the next section.

Overall, there is a wide variation in running times across different programming

		rmat24	rmat27	twitter40	twitter50	road
bfs	Galois	0.5	1.5	0.7	2.5	0.5
bfs	Ligra-g	0.3	1.3	0.8	2.3	1.1
bfs	PowerGraph-g	10.8	84.2	28.0	37.7	17.5
bfs	GraphLab	12.4	83.9	26.7	60.5	4092.7
bfs	Ligra	0.4	1.5	1.2	2.3	2.8
bfs	PowerGraph	7.0	30.8	16.9	24.1	821.6
cc	Galois	7.3	17.9	13.9 [†]	39.6 [†]	0.6
cc	Ligra-g	1.3	11.1	16.6	31.9	62.3
cc	PowerGraph-g	21.8	120.3	58.8	105.0	572.9
cc	GraphLab	14.1	89.6	36.0	64.5	1033.5
cc	Ligra	2.5	22.2	31.7	57.5	127.0
cc	PowerGraph	39.0	129.5	115.5	201.5	2831.5
dia	Galois	1.1	5.1	2.8	5.5	2.6
dia	Ligra-g	2.3	21.4	19.7	44.3	8.6
dia	PowerGraph-g	2029.7	oom	3816.1	4841.9	2466.6
dia	GraphLab	84.8	478.2	192.0	257.1	21363.3
dia	Ligra	1.7	11.8	19.3	45.8	20.1
dia	PowerGraph	1239.0	oom	5376.0	7390.5	7047.5
pr	Galois	1.3	10.3	6.5	10.7	0.5
pr	Ligra-g	1.1	15.6	4.6	11.5	0.4
pr	PowerGraph-g	2.1	21.0	11.7	14.0	0.2
pr	GraphLab	4.9	47.6	45.8	30.7	14.6
pr	Ligra	1.0	11.6	8.7	11.5	0.2
pr	PowerGraph	8.4	38.8	20.4	30.2	10.6
sssp	Galois	1.9	6.0	11.6	8.6	0.6
sssp	Ligra-g	2.8	9.1	10.0	12.5	320.7
sssp	PowerGraph-g	22.8	100.0	43.3	66.8	3317.3
sssp	GraphLab	28.8	153.9	60.9	87.6	28.6
sssp	Ligra	2.3	12.3	15.9	17.8	219.4
sssp	PowerGraph	34.4	78.8	52.9	104.4	18919.2
bc	Galois	1.3	13.7	13.0	12.0	1.3
bc	Ligra-g	1.4	7.6	5.3	12.9	5.1
bc	Ligra	1.2	5.5	6.8	13.9	6.6

Figure 8.15: Runtime in seconds of applications with 40 threads. The label oom indicates the application ran out of memory. In bold are the best times for each input and graph problem pair. (†) indicates that the best time on CC occurred with eight threads: twitter40 (13.8 s), twitter50 (13.6 s).

	rmat24	rmat27	twitter40	twitter50	road
Galois	17	10	14	14	8440
Ligra	10	6	15	15	6262
PowerGraph	9		7	8	>100

Figure 8.16: Approximate diameters computed

models solving the same problem. The variation is the least for PageRank, which is coordinated and topology-driven. For the other graph problems, the performance difference between programming models can be several orders of magnitude and is quite stark for the road input, whose large diameter heavily penalizes coordinated scheduling of data-driven algorithms.

The performance differences can be broadly attributed to three causes.

In some cases, a poor algorithm was selected even though a better algorithm exists and is expressible in the DSL. The PowerGraph diameter application is an example of this. The probabilistic counting algorithm just takes too long on these inputs and gives worse results than the Ligra algorithm, which also can be expressed as a gather-apply-scatter program. [Figure 8.16](#) gives the approximate diameters returned by each algorithm. The PowerGraph algorithm quits after trying diameters up to 100. Both Galois and Ligra algorithms give strict lower-bounds on the true diameter. The PowerGraph algorithm gives a probabilistic estimate.

In some other cases, the same algorithm is expressed in multiple DSLs, but one programming model just has a better system implementation. All the PageRank applications are largely implementations of the same algorithm. Differences between implementations are due to differences in the runtime systems for each programming model.

Finally, a DSL may be unable to capture an important algorithmic optimization—such as when an important optimization cannot be expressed in a DSL or when it can be expressed but the implementation of the DSL cannot adequately exploit it.

An example of not being able to express an optimization is the lack of priority

	rmat24		rmat27		twitter40		twitter50		road	
	8x16	64x16	8x16	64x16	8x16	64x16	8x16	64x16	8x16	64x16
bfs	29.2	21.8	73.0	28.6	73.2	38.5	81.5	50.8	161.5	821.6
cc	114.5	53.5	270.5	71.5	270.0	90.0	406.0	112.0		
pr	11.6	9.9	43.2	14.8	30.5	17.6	42.3	16.4	4.1	5.8
sssp	112.0	76.9	173.9	63.0	175.2	111.0	321.9	127.5		

Figure 8.17: Runtime in seconds of PowerGraph applications on a distributed system with eight or 64 machines

scheduling for the Ligra and PowerGraph applications for SSSP. GraphLab supports priority scheduling, so although the GraphLab SSSP application is worse on scale-free inputs, it performs much better than Ligra and PowerGraph on the road input due to its support for priority scheduling. Thus, in some cases, it is preferable to have inefficient support for priority scheduling than no support at all.

Another example is the push-versus-pull optimization implemented in Ligra. In principle, this optimization can be implemented in any DSL that supports coordinated scheduling of vertex programs, like GraphLab, but GraphLab does not provide any support for user-visible concurrent bag or worklist objects, so it is not possible to efficiently switch between push and pull traversals.

An example of the inability to exploit an optimization is the GraphLab diameter application. The faster pseudo-peripheral algorithm was implemented in GraphLab, but because of large overheads in starting and stopping parallel execution, which are required for the sequential composition of the parallel breadth-first searches, the overall application has very poor performance.

Figure 8.17 shows the performance of PowerGraph when run on a distributed system, the Stampede cluster at the Texas Advanced Computing Center (TACC). Each machine used is an instance of a two processor Intel (E5-2680) machine with each processor having eight cores. Each machine has 32 GB of RAM. Given the poor performance of the con-

		rmat24	rmat27	twitter40	twitter50	road
bfs	Galois	1.4	4.3	2.2	4.8	0.8
bfs	Ligra-g	1.0	4.5	2.8	5.4	1.0
bfs	PowerGraph-g	7.2	47.2	22.7	37.7	5.8
bfs	GraphLab	20.3	90.2	52.3	60.6	3177.2
bfs	Ligra	0.6	2.1	1.8	2.9	2.1
bfs	PowerGraph	11.9	58.7	29.4	42.8	452.8
cc	Galois	1.7	12.9	13.8 [†]	13.6 [†]	1.0
cc	Ligra-g	4.3	33.5	35.8	61.9	236.8
cc	PowerGraph-g	46.4	194.8	144.2	320.1	1336.3
cc	GraphLab	36.1	130.7	89.3	91.5	1512.6
cc	Ligra	2.7	24.9	27.1	49.4	141.0
cc	PowerGraph	70.0	280.5	226.5	384.0	2846.5
dia	Galois	3.5	13.5	7.4	9.8	3.7
dia	Ligra-g	8.8	55.8	54.1	89.4	21.7
dia	PowerGraph-g	2678.4	oom	3969.8	6094.5	3151.5
dia	GraphLab	177.5	810.0	259.6	230.8	20047.4
dia	Ligra	5.1	35.6	39.1	68.0	37.2
dia	PowerGraph	3528.0	oom	14877.0	20161.0	9617.5
pr	Galois	2.2	15.6	8.1	13.1	1.0
pr	Ligra-g	2.7	25.0	12.2	22.9	0.9
pr	PowerGraph-g	2.1	24.5	10.5	13.0	0.7
pr	GraphLab	13.3	92.5	30.8	30.2	28.2
pr	Ligra	2.9	34.7	20.6	34.0	0.4
pr	PowerGraph	12.8	67.2	37.0	51.0	16.6
sssp	Galois	3.4	15.4	12.2	14.8	1.1
sssp	Ligra-g	7.2	25.2	17.9	24.7	1154.2
sssp	PowerGraph-g	38.8	114.6	68.0	140.8	9071.7
sssp	GraphLab	84.0	548.1	199.9	276.2	60.1
sssp	Ligra	4.7	22.0	17.7	29.9	440.8
sssp	PowerGraph	54.2	141.4	90.7	189.1	23556.2
bc	Galois	4.3	27.0	21.2	18.9	1.9
bc	Ligra-g	4.0	20.6	17.7	30.7	3.6
bc	Ligra	2.2	11.3	11.2	17.8	5.4

Figure 8.18: Runtime in seconds of applications with 8 threads. The label oom indicates the application ran out of memory. In bold are the best times for each input and graph problem pair; in all but two cases, best times are with 40 threads (see Figure 8.15).

nected components and SSSP PowerGraph implementations on the road graph on shared-memory machines, they were not run on the distributed machine. Even with 64 machines ($64 \cdot 16 = 1024$ cores), the performance is worse than that of the best implementation on a single machine with 8 cores and 4 times the RAM for all but one application-input combination (see [Figure 8.17](#) and [Figure 8.18](#)). The one slower combination is PageRank on rmat27 where Galois takes 15.6 seconds and PowerGraph takes 14.8 seconds.

Comparison of Implementations

[Figure 8.15](#) also shows the performance results of Ligra and PowerGraph compared to the implementations of their programming models with the Galois system, Ligra-g and PowerGraph-g respectively.

Overall, the Galois implementations of graph DSLs do better than the original DSL implementations, although this varies from DSL to DSL. If pairs of Ligra and Ligra-g runtimes are considered for each graph problem, input and number of threads, in $18/30 \approx 60\%$ of the pairs, the Galois version is faster. However, due to compiler incompatibilities, a different compiler was used for each version of the application. Considering pairs of PowerGraph and PowerGraph-g, runtimes, $18/24 = 75\%$ of the pairs favor the Galois version. As noted earlier, PowerGraph supports distributed-memory execution as well, so some portion of the performance gap is due to the additional overhead of supporting distributed-memory execution and not using it. For GraphLab and Galois, all of the pairs favor Galois, although in this case, the Galois applications include optimizations that could not be implemented in GraphLab, like push-versus-pull.

Some improvements can be made to the Galois versions of these DSLs. For instance, the Ligra version of the diameter application tends to be faster than the Ligra-g version, and the PowerGraph version of BFS tends to scale better than the PowerGraph-g version, but overall, the results suggest that the Galois infrastructure is a reasonable substrate on which graph DSLs can be built.

		rmat24	rmat27	twitter40	twitter50	road
bfs	LigraChi-g	9	133	187	960	12
cc	LigraChi-g	17	205	175	310	169
cc	GraphChi	223	1164	870	1179	120
dia	LigraChi-g	21	192	265	697	29
pr	LigraChi-g	16	143	90	114	6
pr	GraphChi	38	308	154	220	13
sssp	LigraChi-g	36	1127	3227	4873	790
bc	LigraChi-g	18	237	251	1561	14

Figure 8.19: Runtime in seconds of applications on small memory machine with eight threads

Evaluation of Out-of-core DSLs

To evaluate the out-of-core DSLs, GraphChi and LigraChi-g (the combination of Ligra and GraphChi in Galois), a machine with less memory is used. Instead of the machine with 128 GB RAM used in the previous figures, this machine has 24 GB of RAM. It is a two processor Intel (X5570) machine. Each processor has four cores. To test the out-of-core capability, each DSL is given a memory budget of 2 GB of RAM to store graph data. This includes graph adjacency information and edge values, but it does not include user data allocated for a node nor any additional user or runtime-allocated structures. The entire road graph fits in this memory budget.

[Figure 8.19](#) gives the performance of the out-of-core DSLs. Inputs were stored on a 7200 RPM SATA drive. GraphChi allows separate configuration of load threads, which read the graph file, and execute threads, which run the vertex program. For these experiments, the number of threads refers to the number of execute threads. Two load threads are always used.

These out-of-core experiments highlight the impact of having enough memory for graph analytics applications. Ignoring differences in processors but keeping the number of threads the same, on the larger inputs, i.e., rmat27, twitter40 and twitter50, running in a memory-constrained environment with LigraChi-g (see [Figure 8.19](#)) is between 3.4X and

197X slower than performing the same algorithm with Ligra-g on machine m4x10 (compare with [Figure 8.18](#)), an unconstrained memory environment.

These results provide more context for the claim by Kyrola et al. that out-of-core execution of graph analytics only incurs a modest performance penalty ([Kyrola et al., 2012](#)).

The slowdown is between 3.4X and 7.8X for the connected components, approximate diameter and PageRank applications. Considering a 5X reduction in memory, these results suggest a reasonable trade-off between space and time for connected components and PageRank. For the approximate diameter application, there are additional gains from switching to the more expressive Galois programming model.

For the other applications, the slowdown ranges between 11.5X and 197X, not including the additional slowdown of Ligra or Ligra-g versus Galois. The out-of-core DSLs impose a particular scheduling of activities that optimizes I/O operations, but that order may not be efficient from the application standpoint. For more effective out-of-core implementations of these applications, more attention should be paid towards the joint optimization of application and I/O-level scheduling.

8.3 Related Work

There have been several performance evaluations of hardware transactional memory ([Dice et al., 2009](#); [Dalessandro et al., 2011](#); [Wang et al., 2012a](#); [Yoo et al., 2013](#); [Diegues et al., 2014](#)), and as noted in [Section 8.1.3](#), some part of the evaluation in this dissertation reproduces the results of Wang et al. Intel has also released HTM support in their fourth generation Intel Core i7 (Haswell) processors, but it has since been disabled due to reports of an implementation bug ([Intel, 2014](#)). Prior to its disabling, there were several evaluations of the Haswell HTM ([Yoo et al., 2013](#); [Diegues et al., 2014](#)). The hardware is targeted towards short-running, fine-grain transaction programs, which are not the kinds of transactions that arise in STAMP programs. The results of Yoo et al. on STAMP programs using four threads show abort rates significantly higher than using an STM, which suggests a large number of

capacity or false conflicts even at low levels of parallelism. The highest speedup over sequential reported by Diegues et al. for HTM is 3.5X on 8 threads for kmeans. In any event, the goal of this dissertation is not strictly to evaluate HTM implementations but to show how simple mechanisms are sufficient to efficiently execute programs with transactional semantics.

SwissTM (Dragojević et al., 2009) has been shown to perform slightly better than TinySTM, which was used as a representative STM in Section 8.1, but the STAMP benchmarks where SwissTM outperforms TinySTM the most (by 1.2X–1.5X)—intruder, kmeans-high, yada—also have poor scalability, so the actual performance difference is small.

There are several proposals for parallel programming models that support transactional behavior (Adl-Tabatabai et al., 2006; Carlstrom et al., 2006; Chamberlain et al., 2007; Ni et al., 2008), but in these cited works, transactions are fine-grain, and no separation is made between schedulers and user data structures, which is crucial to simplifying transaction implementation. Automatic Mutual Exclusion (AME) (Abadi et al., 2011) introduced a higher level abstraction for transactional execution where code blocks are transactional by default. This matches the spirit of loop-based transactions described here, but the work on AME is mainly concerned with semantics rather than performance.

Chapter 9

Conclusion

The growth of data center and mobile computing has renewed interest in computing efficiency and performance. Prior work on parallelism assumed that activities are independent or that dependences are subsumed by fork-join control dependences, so current general-purpose runtime schedulers embody only a few simple scheduling policies. In contrast, handwritten schedulers for parallel implementations of irregular algorithms often use carefully crafted policies that trade-off excess work for increased parallelism, but they increase the burden of parallel programming and may be difficult to reuse for other algorithms. Additionally, prior programming models often did not clearly distinguish the computational operator from its scheduling and from its data structures.

This dissertation described the design and implementation of a new parallel programming system, Galois, that surmounts these challenges and argues that a distinction between computational operators from scheduling and from data structures is necessary for high performance, high productivity programming models. To support this point, it showed that the performance of a well-studied benchmark suite can be significantly improved by factoring programs along these lines. To show that these benefits can be achieved without much programmer effort, the Galois system was evaluated against existing programming models for graph analytics, and the results showed orders of magnitude performance im-

provements. The benefit is mainly due to better support for more efficient algorithms, but if a simpler programming model is desired, it can be layered on top of the Galois system. The Galois system also facilitates efficient deterministic scheduling.

Overall, these results suggest a promising new direction for the design of programming models—one that sees a parallel program as operator + schedule + parallel data structure.

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