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Array Optimizations for High Productivity Programming Languages

by

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ABSTRACT

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While the HPCS languages (Chapel, Fortress and X10) have introduced improvements in programmer productivity, several challenges still remain in delivering high performance. In the absence of optimization, the high-level language constructs that improve productivity can result in order-of-magnitude runtime performance degradations.

This dissertation addresses the problem of efficient code generation for high-level array accesses in the X10 language. The X10 language supports rank-independent specification of loop and array computations using regions and points. Three aspects of high-level array accesses in X10 are important for productivity but also pose significant performance challenges: high-level accesses are performed through Point objects rather than integer indices, variables containing references to arrays are rank-independent, and array subscripts are verified as legal array indices during runtime program execution.

Our solution to the first challenge is to introduce new analyses and transformations that enable automatic inlining and scalar replacement of Point objects. Our solution to the second challenge is a hybrid approach. We use an interprocedural rank analysis algorithm to automatically infer ranks of arrays in X10. We use rank analysis information to enable storage transformations on arrays. If rank-independent
array references still remain after compiler analysis, the programmer can use X10’s dependent type system to safely annotate array variable declarations with additional information for the rank and region of the variable, and to enable the compiler to generate efficient code in cases where the dependent type information is available. Our solution to the third challenge is to use a new interprocedural array bounds analysis approach using regions to automatically determine when runtime bounds checks are not needed.

Our performance results show that our optimizations deliver performance that rivals the performance of hand-tuned code with explicit rank-specific loops and lower-level array accesses, and is up to two orders of magnitude faster than unoptimized, high-level X10 programs. These optimizations also result in scalability improvements of X10 programs as we increase the number of CPUs. While we perform the optimizations primarily in X10, these techniques are applicable to other high-productivity languages such as Chapel and Fortress.
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Chapter 1

Introduction

Chapel, Fortress and X10, the three languages initially developed within DARPA's High-Productivity Computing System (HPCS) program, are all parallel high-level object-oriented languages designed to deliver both high-productivity and high performance. These languages offer abstractions that enable programmers to develop efficient scientific applications for parallel environments without having to explicitly manage many of the details encountered in low level parallel programming. These languages' abstractions provide the mechanisms necessary to improve productivity in high-performance scientific computing. Unfortunately, runtime performance usually suffers when programmers use early implementations of these languages. Compiler optimizations are crucial to reducing performance penalties resulting from their abstractions. By reducing, or in some cases eliminating the performance penalties, these compiler optimizations should facilitate future adoption of high-productivity languages for high-performance computing by the broad scientific community.

This dissertation focuses on developing productive and efficient implementations of high-level array operations and loop iteration constructs for high-productivity parallel languages. Arrays are important because they are a principal data structure used in scientific applications. Loops are important because they are the primary control structure in scientific applications and they tend to dominate execution time. Our compiler enhancements are designed for high-productivity languages in general and are applicable to all three HPCS languages. This dissertation reports our work on optimizing array accesses in X10 (Chapters 4 and 5) and implementing loop iterators in Chapel (Chapter 6). We take advantage of language features to develop the object
inlining work in part in Fortress (Chapter 4), enabling us to make contributions to all three high-productivity languages. While we detail each contribution in the context of a specific HPCS language as a proof of concept, we want to emphasize that each contribution will be applicable to other high-productivity languages as well.

This work addresses several productivity and performance issues related to high-level loop iteration and array operations. We begin by discussing high-level array accesses. We develop a variant of the object inlining compiler transformation to produce efficient representations for points. A point object identifies an element in a region, and can be used as a multi-dimensional loop index as well as an index into a multi-dimensional array. Object inlining is a transformation designed to replace an object by its inlined fields, resulting in direct field accesses and elimination of the object's allocation and memory costs. We employ a variant of type analysis to discover the dimensionality of points before applying this transformation. We extend object inlining to handle all final object types in scientific computing. We also detail a transformation to generate an efficient array implementation for high-level arrays in high-productivity languages. We evaluate the array transformation that converts high-level multidimensional X10 arrays into lower-level multidimensional Java arrays, when legal to do so. In addition, this thesis makes important advancements to the array bounds analysis problem. We highlight the importance of high-level language abstractions which help our compilation techniques discover and report superfluous bounds checks to the Java Virtual Machine.

We then turn our attention to iterator implementation. An iterator is a control construct that encapsulates the manner in which a data structure is traversed. We illustrate why iterators are important for productive application development and we demonstrate how to efficiently implement iterators in a high-productivity language. While most of our contributions target single-thread performance, we demonstrate the impact that our optimizations have on parallel performance. In particular, this thesis illustrates the effect these transformations have on scalability as we increase
the number of CPUs.

This research highlights the advantages of providing abstractions for iterating over data structures in a productive manner. It addresses both implementation details and optimization opportunities to leverage support for these abstractions in a scientific computing environment. The dissertation presents experimental results that demonstrate the importance of this work. Subsequent discussion summarizes our results and provides insight into possible future extensions of this research.

**Thesis Statement:** Although runtime performance has suffered in the past when scientists used high productivity languages with high-level array accesses, our thesis is that these overheads can be mitigated by compiler optimizations, thereby enabling scientists to develop code with both high productivity and high performance. The optimizations introduced in this dissertation for high-level array accesses in X10 result in performance that rivals the performance of hand-tuned code with explicit rank-specific loops and lower-level array accesses, and is up to two orders of magnitude faster than unoptimized, high-level X10 programs.
Chapter 2

Background

Since the transition from assembly language to Fortran [7, 74] and subsequent higher level languages for scientific computing, programmers have always been concerned about the tradeoff between programmability and performance. As architectures become increasingly complex, high-level languages and their programming models will need to provide abstractions that deliver a sufficient fraction of available performance on the underlying architectures without exposing too many low-level details. If they do, these languages should be attractive to the broader scientific computing community. The following sections introduce a non-exhaustive list of some of the languages and libraries that play a role in solving this challenging problem.

2.1 Message Passing Interface

The dominant parallel programming paradigm in high-performance scientific computing currently is some combination of Fortran/C/C++ with the message passing interface (MPI) [97]. MPI is a well-defined library that has long been the de facto standard in parallel computing for processor communication. Because MPI is a library, programmers don’t have to learn a whole new language to write parallel programs. However, using the single program multiple data (SPMD) execution model with two-sided communication inhibits the programmer’s productivity potential by enforcing per-processor application development.

This model places the burden of designing data distribution, computation partitioning, processor communication, and synchronization on the programmer. As a result, programmers must manage many of the low-level details of parallel program-
ming, thereby reverting back to assembly-like programming. Even expert programmers with this low-level responsibility are prone to introducing subtle parallel bugs in the code [24]. The early communication binding to traditional MPI limits the opportunities to take advantage of architectures which support one-sided communication. While MPI-2 [45] supports one-sided communication, it is currently unsuitable for parallel languages [102]. Another limitation of MPI is that it inherits the weaknesses of the programmer's language of choice for application development. For example, when programming in C/C++, compiler optimizations may be limited due to the complications arising from pointer analysis.

2.2 Data-Parallel Languages

Data-parallel languages enable programmers to develop sequential applications with annotations to specify data distributions. Next we introduce some of the data-parallel languages used in high-performance computing.

2.2.1 High Performance Fortran

HPF [33, 61, 63] is a global-view, data-parallel language that essentially extends Fortran 90 by adding array distributions. The compiler and runtime handle the mapping of the distributed arrays to the hardware. Clearly, programming in a data-parallel language improves productivity by shifting the burden of processor communication and synchronization to the compiler and runtime. A limitation of utilizing data-parallel languages like HPF tends to be the lack of language support for more general distributions to suitably implement computations with sparse data structures [61]. Another limitation of pure data-parallel languages is the lack of support for nested parallelism. As a result, parallel solutions to problems like divide and conquer can be challenging. These limitations combined with HPF portability and performance tuning issues [61] factored in the decline in popularity for data-parallel languages such as HPF for high-performance scientific computing.
2.2.2 ZPL

ZPL [90] is a global-view, data-parallel language. Similar to HPF, ZPL does not expose to the programmer the details of processor communication and synchronization. However, ZPL does support language abstractions which make processor communication visible to the programmer [24]. ZPL's sparse array and region structural language abstractions improve programmability by separating algorithmic specification from implementation. As a result, the programmer can focus on implementing the computation. Factoring out the specification also enables programmers to easily interchange specifications without impacting the algorithm's implementation.

Chamberlain et al. [24] show they can develop parallel applications in ZPL and still achieve results comparable to Fortran + MPI. They provide results for the NAS Parallel CG and FT benchmarks [9]. These results show that its possible to program in high-level languages without incurring severe performance penalties. Limiting the generality of ZPL is the lack of support for user-defined distributions to facilitate natural implementations of irregular computations.

2.2.3 CM Fortran

CM Fortran [95] is an extension of Fortran 77 with array processing features for the Connection Machine. In CM Fortran, operations on array elements can be performed simultaneously on a distributed memory system by associating one processor with each data element. One drawback of many CM Fortran codes is that, because they were tied to the CM-2 and CM-5 machines, when the Thinking Machine Corporation stopped supporting the hardware, those codes had to be ported to other languages such as HPF [88].
2.3 Task-Parallel Languages

Task-parallel languages support spawning of tasks to work on asynchronous program blocks. The next sections introduce some of the task-parallel languages in scientific computing.

2.3.1 OpenMP

OpenMP [35, 81] is a task-parallel language that focuses on the parallelization of loops. Programmers develop sequential applications and the compiler provides support to parallelize the loops. However, because OpenMP provides no support for data distributions, it does not scale well for distributed memory or non-uniform memory access (NUMA) architectures.

2.3.2 Java

Over the past decade, Java [46] has become one of the most popular programming languages. Java, primarily developed for the internet, is not well-suited to support high-performance computing for a variety of reasons. Java does not support multi-dimensional arrays. As a result, a programmer has to create arrays of arrays to simulate a multi-dimensional array. Because this extra level of indirection carries a performance penalty, programmers often provide confusing hand-coded one-dimensional representations using complex array indexing schemes to eliminate it. Consequently, while performance improves, productivity and readability suffer.

An additional concern for Java is the lack of support for sparse arrays. Programmers often use multiple one-dimensional Java arrays to model array sparsity. Another issue facing Java is the concurrency model. The principal way programmers develop applications with concurrency in Java is through threads, though the usage of the Java Concurrency Utilities is now on the rise [83]. While threads allow Java to address task-parallelism, they ignore locality opportunities due to its flat memory model. One final issue worth mentioning is the Java Virtual Machine (JVM). Because
the JVM interprets or dynamically compiles Java byte codes, Java applications often run slower than those that are statically compiled to native code. While having a portable JVM is an attractive feature for internet computing, it is undesirable for high-performance scientific computing if it leads to degradations in performance.

2.4 Partitioned Global Address Space Languages

As the popularity of data-parallel languages in high-performance scientific computing declined, new parallel partitioned global address space (PGAS) languages emerged. Titanium, UPC, and Co-Array Fortran are all PGAS languages with a single program multiple data (SPMD) memory model. These languages make developing parallel applications easier than developing with MPI because of the global address space.

2.4.1 Titanium

Titanium [103], a dialect of Java, leverages many of Java’s productivity features such as strong typing and object-orientation. As a result, a broad base of programmers will already be familiar with a core subset of Titanium’s features targeting productivity. Compared to Java, Titanium has more language features to support scientific computing. For example, Titanium provides support for multi-dimensional arrays. Titanium’s multi-dimensional arrays enhance programmability and eliminate the need for complex array indexing schemes common to Java due to Java’s lack of support for multi-dimensional arrays. Titanium also supports data distributions for arrays, trees, and graphs where the data-parallel languages described earlier provided distribution support for arrays only. However, to naturally express irregular computations such as adaptive mesh refinement (AMR), Titanium’s distributed data structures require an additional array of pointers [102]. Each element of the array points to a local section of distributed data resulting in a sacrifice of programmability to express more general computations. In addition, due to its approach of static compilation to C code, many dynamic features of Java are not supported in Titanium.
2.4.2 Unified Parallel C

UPC [23, 96], a parallel extension to C, is designed to give the programmer efficient access to the hardware. UPC sacrifices programmability (due to the use of C as its foundation and a user-controlled memory consistency model) for programmer control over performance. UPC views the machine model as a group of threads working cooperatively in a shared global address space. Similar to Titanium, programmers may specify data as local or global. However, by default, UPC assumes data is local whereas, in Titanium, data is global by default. This model gives programmers explicit control over data locality. In addition, programmers may specify whether a sequence of statements has a strict or relaxed memory model. The former ensures sequential consistency [64] with respect to all threads while the latter ensures sequential consistency with respect to the issuing thread [23]. Compiler analysis and optimization of UPC applications can be challenging due to the use of pointers.

2.4.3 Co-Array Fortran

Co-Array Fortran [78], an extension to Fortran 95, is designed to provide a minimal set of additional language abstractions to Fortran 95 to develop parallel applications. Each replication of a Co-array Fortran program is called an image. Co-array Fortran introduces the co-array, a language abstraction enabling programmers to distribute data on different images. One benefit of co-arrays is that they give programmers an explicit control and view over how data is distributed across images. The co-array's co-shape determines the image communication topology. Co-spaces are limited to expressing only Cartesian topologies. However, there are applications for which a Cartesian topology is not ideal. Programmers circumvent this problem by using neighbor arrays. Dotsenko [38] discusses the limitations of using neighbor arrays to express arbitrary communication topologies.
2.5 High Productivity Computing Languages

Chapel, Fortress, and X10 are all parallel object-oriented global address space languages emerging from the Defense Advanced Research Projects Agency (DARPA) challenge to the scientific community to increase productivity by a factor of 10 by the year 2010. These languages aim to increase productivity in high-performance scientific computing without sacrificing performance.

2.5.1 X10

X10 [25] is an object-oriented global address space language designed for scalable, high-performance computing. As with Titanium, Java developers will already be familiar with a core subset of X10 features, facilitating a natural transition to parallel program development. In fact, programmers can compile sequential Java 1.4 programs in X10, an attractive feature when attempting to migrate developers from preexisting languages with a broad user base. X10 provides language abstractions to improve programmability in high-performance computing. The point, range and dist abstractions provide programmers the opportunity to express distributed array computations in a productive manner. Programmers may omit rank (dimensionality) information when declaring X10 arrays, encouraging the development of rank-independent computations. X10 introduces the place abstraction to enable developers to exploit data locality by co-locating data with a place. In addition, X10 gives developers control over task-parallelism with the async and future constructs. Programmers can utilize these constructs to explicitly spawn asynchronous activities (light-weight threads) at a given place. Another productivity feature of X10 programs is that they are deadlock-free, if restricted to a (large) subset of X10 constructs.

The X10 team has shown the productivity benefits of X10 by parallelizing serial code [26, 40] and the performance benefits of programming in X10 on an SMP architecture [10] for the Java Grande benchmark suite [20]. While these results compared Java versus X10, in the future, X10 is expected to show results comparable to C/
Fortran with MPI, the dominant parallel programming paradigm currently utilized in high-performance scientific computing.

2.5.2 Chapel

Chapel [22, 34] is an object-oriented parallel language promoting high-productivity in high-performance computing. Chapel introduces the domain, a language abstraction influenced by ZPL regions that, when combined with a distribution, supports dynamic data structures useful for irregular computations such as adaptive mesh refinement. Similar to X10, Chapel allows programmers to exploit data locality with the locale abstraction while the cobegin statement allows programmers to express task-parallelism.

The design of Chapel is guided by four key areas of programming language technology: multithreading, locality-awareness, object-orientation, and generic programming. The object-oriented programming area, which includes Chapel’s iterators, helps in managing complexity by separating common function from specific implementation to facilitate reuse. Traditionally, when programmers want to change an array’s iteration pattern to tune performance (i.e. such as converting from column major order to row major order when migrating code from Fortran to C), the algorithm involving the arrays would be affected; even though the intended purpose is to change the specification, not the algorithm itself. Chapel iterators achieve the desired effect by factoring out the specification from implementation. Chapel supports two types of simultaneous iteration by adding additional iterator invocations in the loop header. Developers can express cross-product iteration in Chapel by using the following notation:

\[
\text{for (i,j) in [iter1(),iter2()]} \text{ do } ...
\]

which is equivalent to the nested for loop:

\[
\text{for i in iter1()} \text{ do }
\]
Zipper-product iteration is the second type of simultaneous iteration supported by Chapel, and is specified using the following notation:

\[
\text{for (i,j) in (iter1(), iter2()) do ...}
\]

which, assuming that both iterators yield \( k \) values, is equivalent to the following pseudocode:

\[
\text{for p in 1..k {
    var i = iter1().getNextValue();
    var j = iter2().getNextValue();
    ...
}}
\]

In this case, the body of the loop will execute each time both iterators yield a value. Similar to X10, the Chapel programming language is expected to show performance results comparable to C/Fortran with MPI to persuade scientists that Chapel is a suitable alternative to commonly utilized languages in high-performance computing.

### 2.5.3 Fortress

Fortress [3] is an object-oriented component-based parallel language that, along with X10 and Chapel, seeks to improve productivity in high-performance computing without sacrificing performance. Fortress introduces a parallel programming paradigm distinct from the other high-productivity computing languages. In Fortress, the \textit{for} loop is parallel by default, forcing the programmer to be aware of parallelism inside loops from the beginning of the development cycle. Fortress introduces the \textit{trait}, an abstraction specifying a collection of methods [3] which an object may extend. Traits simplify the traditional object-oriented inheritance of Java. In Fortress, objects cannot extend other objects or have abstract methods. One programmability advantage
that Fortress promotes is the natural expression of mathematical notation. As a result, Fortress eliminates learning programming language syntax as a prerequisite to expressing mathematical formulas. Because Fortress is built on libraries, these libraries must be efficient with respect to parallel performance for the adoption of the language by the scientific community.
Chapter 3

Related Work

We highlight in this section the related work in the areas of high-level iteration, object inlining, optimization of array accesses, and type inference.

3.1 High-Level Iteration

Iteration over data structures has long been a programmability concern. General iterator abstractions are essential to increasing productivity in high-performance computing. Iterators can facilitate a natural separation of concerns between data structures and algorithms. They separate the data structure iteration pattern from the actual computation, two problems that are orthogonal to each other. In addition, providing implicit language support for parallel iteration is important for parallel environments. The following sections discuss several language iterator implementations. We later compare these language iteration approaches to our work on Chapel iterators.

3.1.1 CLU

CLU [68, 69] iterators are semantically similar to those in Chapel. Unlike Chapel, the CLU language only permits invocation of CLU iterators inside the loop header. Each time the iterator yields a value, the body of the loop is executed. Both Chapel and CLU support nested iteration. Nested iteration occurs when, for each value that iterator $i$ yields, iterator $j$ yields all of its values. In CLU, only one iterator can be called in the loop header. As a result, CLU does not provide support for zippered iteration [59]; a process of traversing through multiple iterators simultaneously where each iterator must yield a value once before execution of the loop body can begin.
3.1.2 Sather

In contrast to CLU iterators, Sather iterators [76] can be invoked from anywhere inside the loop body. As a result, Sather iterators can support zippered iteration by invoking multiple iterator calls inside the loop body. Since Sather iterators may appear inside the loop body, iterator arguments may be reevaluated for each loop iteration. The semantics of Sather iterators are similar to both Chapel and CLU iterators. Sather iterators support zippered and nested iteration. However, Chapel's focus on high-level iteration in a parallel environment by factoring iteration implementation details out from the loop body separates itself from Sather.

3.1.3 Coroutine

A coroutine [48] is a routine that yields or produces values for another routine to consume. Unlike functions in most modern languages, coroutines have multiple points of entry. When encountering the \texttt{yield} in a coroutine, execution of the routine is suspended. The routine saves the return value, program counter, and local static variables in some place other than a stack. When the routine invocation occurs again, the execution resumes after the yield. We use zippered iteration in Chapel to provide this producer-consumer relationship in a modern language.

3.1.4 C++, Java, and Python

C++ [91] STL, Java [46], and Python [86] provide high level iterators that are not tightly coupled to loops like Chapel, CLU, and Sather iterators. These iterators are normally associated with a container class. These languages support simultaneous iteration on containers. Within these languages, Java and Python provide built-in support to perform iteration using special \texttt{for} loops that implicitly grab each element in the container, thereby separating the specification of the algorithm from its implementation. However, Java and Python's special \texttt{for} loops do not support zippered iteration since they may call only one iterator in the loop header.
3.1.5 Sisal and Titanium

Sisal [43] and Titanium [103] also provide some support for iterators using loops. Titanium has a \textit{foreach} loop that performs iteration over arrays when given their domains. Sisal supports 3 basic types of iterators using a \textit{for} loop. The first type iterates over a range specified by an lower and upper bound. The second type of iterator returns the elements of an array or stream (a stream is a data structure that is similar to an array). The third type of iterator returns tuples of a dot- or cross-product constructed from two range iterators. Sisal and Titanium iterators are limited when compared to Chapel iterators. They don't support zipperred iteration or general iteration such as column-major order or tiled iteration.

3.2 Optimized Compilation of Object-Oriented Languages

Broad adoption of high-level languages by the scientific community is unlikely without compiler optimizations to mitigate the performance penalties these languages' abstractions impose. The following sections detail optimizations employed in object-oriented languages to improve performance.

3.2.1 Object Inlining

\textit{Object inlining} [16, 19, 36, 37] is a compiler optimization for object-oriented languages that transforms objects into simple data, and conversely the rest of the program code that operates on objects into code that operates on inlined data. It is closely related to "unboxing" [65] for functional languages. Budimlić [16] and Dolby [36] introduced object inlining as an optimization for object-oriented languages, particularly for Java and C++. General form of object inlining requires complex escape analysis [29, 32] and concrete type inference [1], and the transformation is irreversible (once unboxed, objects cannot be boxed again in general). In past work, we extended the analysis to allow more objects and arrays of objects to be inlined in scientific, high performance
Java programs [18, 57]. The object inlining approach for X10 \textit{points} presented in this dissertation is more broadly applicable to point and value objects (\textit{all} point objects can be boxed and unboxed freely) than traditional inlining of mutable objects.

Zhao and Kennedy [104] provide an array scalarization algorithm for Fortran 90 which reduces the generation or size of temporary arrays, improving memory performance and reducing execution time. We also improve memory performance and reduce execution time by generating more efficient array computations. In our case, we generate efficient representations of high level array operations by inlining temporary \textit{point} objects and performing an array transformation on general X10 arrays.

3.2.2 Semantic Inlining

Wu et al. [99] presented \textit{Semantic Inlining} for Complex numbers in Java, an optimization closely related to object inlining. Their optimization incorporates the knowledge about the semantics of a standard library (Complex numbers) into the compiler, and converts all Complex numbers into data structures containing the real and imaginary parts. Although this optimization achieves the same effect as object inlining for Complex numbers, it is less general since it requires compiler modifications for any and all types of objects for which one desires to apply this optimization.

3.2.3 Point Inlining in Titanium

The point-wise \textit{for} loop language abstraction is not unique to the X10 language. Titanium [103], a Java dialect, also has \textit{for} loops which iterate over points in a given domain. There are two important advantages to using point-wise iteration for arrays. First, it prevents programmer errors induced by complicated iteration patterns. Second, the compiler can recognize that iterating over domain \( d \) eliminates the need for array bounds checking when the programmer accesses an array with domain \( d \).

The Titanium compiler also performs an optimization to remove points appearing
inside for loops. However, there are several differences between our approach and the one applied in Titanium. First, our work on object inlining is directly applicable to all value objects, not just points, and thus is a more general optimization. Transformation of points in Titanium, as far as we are aware, is designed specifically to convert loops involving points into loops with iterator variables and does not apply to other point objects. Second, because in X10 the rank specification of both points and arrays is not required at the declaration site, we also need to employ a type analysis algorithm to determine the rank for all X10 arrays. Omitting the rank when declaring an array allows the programmers to perform rank independent array calculations and increases their productivity.

3.2.4 Optimizing Array Accesses

In this section we discuss past work in array access optimization. We begin with optimizations aiming to reduce array bounds checking costs. Bodík et al. [13] reduce array bounds checks in the context of dynamic compilation. They focus their optimization on program hot spots to maximize benefits and to amortize the cost of performing the analysis on a lightweight inequality graph. Results from a prototype implementation in Jikes RVM [5] show that their analysis on average eliminates 45% of dynamic array bounds checks. Rather than modifying the JVM, we follow an alternate strategy in which the X10 compiler communicates with the JVM when it determines that array bounds checking is unnecessary. As a result, the X10 runtime and JVM don't have to perform array bounds analysis. Performing compile-time array bounds checking without adding runtime checks prevents the additional runtime costs resulting from array bounds analysis. Suzuki and Ishihata [94] provide an intraprocedural array bounds checking algorithm based on theorem proving which is costly. Most JIT compilers also perform array bounds analysis to eliminate bounds checks. However, the analysis is generally intraprocedural since the JIT is performing the analysis dynamically. We actually propagate interprocedural information which
enables us to potentially remove array bounds checks involving formal parameters, a case that JIT compilation would miss.

Aggarwal and Randall [2] use related field analysis to eliminate bounds checks. They observe that an array $a$ and an integer $b$ may have an invariant relationship where $0 < b < a.length$ for every instance of class $c$. Proving this invariant holds on every method entry and exit enables them to remove array bound checks in the program. To find related fields, they analyze every pair $[a,b]$ where $a$ is a field with type array(1-Dimensional) and $b$ is a field with type integer in class $c$. By contrast, we examine every array, region, point, and integer variable. As a result, we can catch useless bound checks for multi-dimensional arrays that Aggarwal and Randall would miss. We reduce the related variable analysis cost by limiting integer variable analysis to only those variables with a region relationship. Recall, an integral has a region relationship if the program assigns it a region bound or program execution assigns it a variable that represents a region or region bound. Consequently, Aggarwal and Randall may eliminate more 1-D array accesses since they analyze every $[a,b]$ pair.

Heffner et al [52] extend Aggarwal and Randall. by addressing the overhead required to prove program invariants for field relations at each point in the program. Thread execution in between two related fields during object construction can invalid invariants in multi-threaded code. Heffner observes that, in general, program execution accesses object fields in a structured way in concurrent environments. Proving that objects with related fields are modified atomically guarantees that the invariants hold in concurrent programs.

Gupta [49] uses a data-flow analysis technique to eliminate both identical and subsumed bounds checks. Ishizaki et al. [53] extends Gupta’s work by showing when bounds checks with constant index expressions can be eliminated. For example, when Ishizaki’s analysis encounters array accesses $a[i]$, $a[i+1]$, and $a[i+2]$, it will subsequently eliminate the array bounds checks for $a[0]$, $a[1]$, and $a[2]$. This algorithm relies on the assumption that all arrays have a lower bound of 0. This technique
would need to be extended for X10 arrays since the lower bound can be non-zero. The Array SSA form [62] work is related to our redundant array access analysis since it also demonstrates how to optimize code in the presence of multiple accesses to the same array element by providing a framework that enables parallel execution of code with output dependences.

3.2.5 Type Inference

FALCON [85], a compiler for translating MATLAB programs into Fortran 90, performs both static and dynamic inference of scalar (e.g. real, complex) or fixed array types. Statically, FALCON's compiler analysis attempts to derive precise intrinsic types when possible, resolving type ambiguity by choosing the more general type. Dynamically, when a variable's type is unknown, the compiler inserts a runtime check to determine if the type is real or complex, cloning the code for both possible cases. Since we do not define a partial order for ranks using a subtype relationship, when ambiguity cannot be resolved using specialization, we resolve the rank to $\perp$ (bottom). Because FALCON is a batch compiler, it doesn't have calling context information for the function it compiles. FALCON addresses this limitation by looking at its input files to get type information [4]. MAJIC [4], a MATLAB just-in-time compiler, compiles code ahead of time using speculation. MAJIC performs interprocedural analysis by using the source code to speculate about the runtime context. If speculation fails at runtime, the JIT recompiles the code using runtime type information. Both the FALCON and MAJIC type inference schemes are limited compared to our precise type inference with type jump functions since neither uses symbolic variables to resolve types.

The use of equivalence sets in our type analysis algorithm builds on past work on equivalence analysis [6, 66] and constant propagation [89, 98]. As in constant propagation, we have a lattice of bounded height $\leq 3$. By computing the meet-over-all-paths, our type inference may be more conservative than Sagiv's [87] type inference
dynamic programming algorithm for finding the meet-over-all-valid-paths solution. Other type inference algorithms such as Joisha’s [56] provide novel solutions to type problems with lattices of unbounded height (e.g., array shape).

The idea of creating specialized method variants based on the calling context is related to specialized library variant generation derived from type jump functions [27]. McCosh’s [73] type inference strategy generates pre-compiled specialized variants for MATLAB [72]. This strategy then replaces procedure calls with calls to the specialized variants based on the calling context. If a variable resolves to more than one type, McCosh generates a specialized variant for the general type. The context in which we apply our algorithm differs from McCosh since we perform type inference in an object-oriented environment on rank-independent type variables that we must map to rank-specific types. Our type inference algorithm requires that the formal parameters converge to a precise type (rank) since we translate X10 rank-independent code into Java rank-specific code. During rank analysis, we can use return jump functions to identify precise ranks for array computation involving formal parameters with multiple ranks. However, without function cloning [31] during rank analysis, formal parameters with multiple ranks resolve to ⊥. In practice, we have frequently generated the more efficient lower-level rank-specific version of the X10 arrays since programmer’s often do not take advantage of developing rank-independent code within the same program. However, to generate efficient code when applying rank-independent functions to arguments of different ranks within the same program, we could extend our approach using function cloning during rank analysis to obtain precise types. This extension would include a heuristic to ensure that we only clone paths that lead to better performance.

Plevyak and Chien [84] developed a type inference algorithm targeting object-oriented languages. The complexity of their algorithm depends of the imprecision of the type variables. Our algorithm is independent of the type imprecision in the program. Pechtchanski and Sarkar’s [82] type inference strategy combines the ad-
vantages of both static analysis and dynamic optimization. As a result, they can
use a more optimistic approach compared to whole-program static analysis. They
invalidate and reanalyze methods when their optimistic assumptions are false. Their
approach could be advantageous for programs that pass arrays of different ranks to
a method's formal parameter.

There are differences between this work and past work on APL analysis and opti-
mization [28, 42]. APL, a dynamic language, requires a runtime system with support
for untyped variables (and incurs the overhead of such a system). In contrast, X10
is statically typed, except that an array's rank/region is treated as part of its value
rather than its type. Further, a major thrust of past work on APL optimization has
been to identify scalar variables. The X10 type system differentiates between scalars
and arrays. Hence, performance improvements for X10 must come from sources other
than scalar identification.
Chapter 4

Efficient High-Level X10 Array Computations

The DARPA High Productivity Computing Systems (HPCS) program has challenged supercomputer vendors to increase development productivity in high-performance scientific computing by a factor of 10 by the year 2010 (the start of the HPCS program was in 2002). Participants in the HPCS program recognized that introducing new programming languages is important for meeting this productivity goal, and three languages have emerged as a result of this initiative: Chapel (Cray), X10 (IBM), and Fortress (Sun). These languages have significantly improved the programmability of high-performance scientific codes through the use of higher-level language constructs, object-oriented design, and higher-level abstractions for arrays, loops and distributions [41]. Chapter 6 demonstrates the programmability benefits of performing high-level loop iteration in Chapel. Unfortunately, high programmability often comes at a price of lower performance. The initial implementations of these higher-level abstractions in the HPCS languages can sometimes result in up to two orders of magnitude longer execution times when compared to current languages such as C, Fortran, and Java.

This chapter outlines the novel solutions necessary to generate efficient array computations for high productivity languages, particularly X10. Figure 4.1 shows the X10 compiler structure assumed in our research in the Habanero project [50]. The chapter focuses on compiler analyses and optimizations that improve the performance of high level array operations in high productivity languages — compilers for other high productivity languages have a similar structure to Figure 4.1.

In this chapter, we focus on the X10 language abstractions pertinent to high level
array accesses. There are two aspects of high level array accesses in X10 that are important for productivity but that also pose significant performance challenges. First, the high level accesses are performed through Point objects\(^1\) rather than integer indices. Points support an object-oriented approach to specifying sequential and parallel iterations over general array regions and distributions in X10. As a result, the Point object encourages programmers to implement reusable high-level iteration abstractions to productively develop array computations for scientific applications without having to manage many of the details typical for low-level scientific programming. However, the creation and use of new Point objects in each iteration of a loop can be a significant source of overhead. Second, variables containing references to Points and arrays are rank-independent \(i.e.,\) by default, the declaration of an array reference variable in X10 does not specify the rank (or dimension sizes) of its underlying array. This makes it possible to write rank-independent code in X10, but poses a challenge for the compiler to generate efficient rank-specific code.

Our solution to the first challenge is to extend the X10 compiler so as to perform automatic inlining and scalar replacement of Point objects. We have a hybrid solution to the second challenge that uses automatic compiler support to infer exact

\(^1\)Points are described later in Section 4.1
ranks of rank-free variables in many X10 programs and programmer support via X10’s *dependent type system* to enable the programmer to annotate selected array variable declarations with additional information for the rank and region of the variable. Subsequently, using dependent type information, the compiler automatically generates efficient code.

The novel contributions to generating efficient X10 array computations are the following:

- **Object Inlining for Points and Value Types.** We utilize the value type property of points to effectively perform object inlining on all rank-independent points anywhere in the code. Recall, the value type property prevents objects from being modified once initially defined. Past work [16, 19, 36, 37, 99, 103] was more conservative due to potential aliasing of mutable objects or more restrictive by only allowing inlining of a specific class or enabling object inlining in certain code regions.

- **Runtime Performance.** Our compiler optimizations improve X10 applications with general X10 arrays by 2 orders of magnitude, relative to the open source reference implementation of X10 [101]. In addition, we demonstrate that our compiler techniques enable better scalability when scaling from 1 CPU to 64 CPUs.

- **Safety Analysis and Array Transformation.** The X10 general array supports a rich set of operations that are not supported by Java arrays. As a result, before we can convert X10 arrays into Java arrays to boost runtime performance, we must perform safety analysis. Safety analysis ensures that, for each operation on an optimized X10 array, there is a semantically equivalent operation for the Java array.
4.1 X10 Language Overview

In this section, we summarize past work on X10 features related to arrays, points, regions and loops [25], and discuss how they contribute to improved productivity in high performance computing. Since the introduction of arrays in the FORTRAN language, the prevailing model for arrays in high performance computing has been as a contiguous sequence of elements that are addressable via a Cartesian index space. Further, the actual layout of the array elements in memory is typically dictated by the underlying language e.g., column major for FORTRAN and row major for C. Though this low-level array abstraction has served us well for several decades, it also limits productivity due to the following reasons:

1. *Iteration*. It is the programmer's responsibility to write loops that iterate over the correct index space for the array. Productivity losses can occur when the programmer inadvertently misses some array elements in the iteration or introduces accesses to non-existent array elements (when array indices are out of bounds).

2. *Sparse Array accesses*. Iteration is further complicated when the programmer is working with a logical model of sparse arrays, while the low level abstraction supported in the language is that of dense arrays. Productivity losses can occur when the programmer introduces errors in managing the mapping from sparse to dense indices.

3. *Language Portability*. The fact that the array storage layout depends on the underlying language (e.g., C vs. FORTRAN) introduces losses in productivity when translating algorithms and code from one language to another.

4. *Limitations on Compiler Optimizations*. Finally, while the low-level array abstraction can provide programmers with more control over performance using constructs like COMMON blocks and pointer aliasing, there is a productivity
loss incurred due to the interference between the low-level abstraction and the compiler's ability to perform data transformations for improved performance (such as array dimension padding and automatic selection of hierarchical storage layouts).

The X10 language addresses these productivity limitations by providing higher-level abstractions for arrays and loops that build on the concepts of *points* and *regions* (which were in turn inspired by similar constructs in languages such as ZPL [90]). A *point* is an element of an $n$-dimensional Cartesian space ($n \geq 1$) with integer-valued coordinates, where $n$ is the *rank* of the point. A *region* is a set of points, and can be used to specify an array allocation or an iteration construct such as the point-wise for loop. The benefits of using points inside of *for* loops include: potential reuse of common iteration patterns via storage inside of regions and simple point references replacing multiple loop index variables to access array elements. We use the term, *compact region*, to refer to a region for which the set of points can be specified in bounded space\(^2\), independent of the number of points in the region. Rectangular, triangular, and banded diagonal regions are all examples of compact regions. In contrast, sparse array formats such as compressed row/column storage are examples of non-compact regions.

Points and regions are first-class value types [8] — a programmer can declare variables and create expressions of these types using the operations listed in Figure 4.2 — in X10 [80, 100]. In addition, X10 supports a special syntax for point construction — the expression, \([a,b,c]\)”, is implicit syntax for a call to a three-dimensional point constructor, \(\text{point.factory}(a,b,c)\)” — and also for variable declarations. The declaration, \(\text{point } p[i,j]\)” is exploded syntax for declaring a two-dimensional point variable p along with integer variables i and j which correspond to the first and second elements of p. Further, by requiring that points and regions be value types,

\(^2\text{For this purpose, we assume that the rank of a region can be assumed to be bounded.}\)
Region operations:

R.rank ::= # dimensions in region;
R.size() ::= # points in region
R.contains(P) ::= predicate if region R contains point P
R.contains(S) ::= predicate if region R contains region S
R.equal(S) ::= true if region R and S contain same set of points
R.rank(i) ::= projection of region R on dimension i (a one-dimensional region)
R.rank(i).low() ::= lower bound of i-th dimension of region R
R.rank(i).high() ::= upper bound of i-th dimension of region R
R.ordinal(P) ::= ordinal value of point P in region R
R.coord(N) ::= point in region R with ordinal value = N
R1 && R2 ::= region intersection (will be rectangular if R1,R2 are rectangular)
R1 || R2 ::= union of regions R1 and R2 (may or may not be rectangular,compact)
R1 - R2 ::= region difference (may or may not be rectangular,compact)

Array operations:

A.rank ::= # dimensions in array
A.region ::= index region (domain) of array
A.distribution ::= distribution of array A
A[P] ::= element at point P, where P belongs to A.region
A | R ::= restriction of array onto region R (returns copy of subarray)
A.sum(), A.max() ::= sum/max of elements in array
A1 <op> A2 ::= returns result of applying a point-wise op on array elements,
when A1.region = A2. region
( <op> can include +, -, *, and / )
A1 || A2 ::= disjoint union of arrays A1 and A2
(A1.region and A2.region must be disjoint)
A1.overlay(A2) ::= array with region, A1.region || A2.region,
with element value A2[P] for all points P in A2.region

Figure 4.2 : Region operations in X10
the X10 language ensures that individual elements of a point or a region cannot be modified after construction.

A summary of array operations in X10 can be found in Figure 4.2. A new array can be created by restricting an existing array to a sub-distribution, by combining multiple arrays, and by performing point-wise operations on arrays with the same region. Note that the X10 array allocation expression, “new double[R]”, directly allocates a multi-dimensional array specified by region R. In its full generality, an array allocation expression in X10 takes a distribution instead of region. However, we will ignore distributions in this chapter and limit our attention to single-place executions although it is straightforward to extend the algorithm to handle distributed arrays.

As an example, consider the Java and code fragments shown in Figure 4.3 for the Java Grande Forum [54] SOR benchmark\(^3\). Note that the i and j loops in the Java version involve a lot of manipulation of explicit array indices and loops bounds that can be error prone. In contrast, the rank-specific X10 version uses a single for loop to iterate over all the points in the inner region (R.inner), and also uses point expressions of the form “t+[-1,0]” to access individual array elements. One drawback of the point-wise for loop in the X10 version is that (by default) it leads to an allocation of a new point object in every iteration for the index and for each subscript expression, thereby significantly degrading performance. Fortunately, the optimization techniques presented in this chapter enable the use of point-wise loops as in the bottom of Figure 4.3, while still delivering the same performance as manually indexed loops as in the top of Figure 4.3.

Figure 4.3 also contains a rank-independent X10 version. In this case, an additional loop is introduced to compute the weighted sum using all elements in the stencil. Note that the computation performed by the nested t and s for loops in this version can be reused for different values of R.inner and stencil with different ranks. Although

\(^3\)For convenience, we use the same name, G, for the allocated array as well as the array used inside the SOR computation, even though the actual benchmark uses distinct names for both.
Java version:

```java
double G[][] = new double[M][N];
int Mml = M-1; int Nml = N-1;
for (int p=0; p<num_iterations; p++) {
    for (int i=1; i<Mml; i++) {
        double[] Gi = G[i];
        double[] Giml = G[i-1];
        double[] Gipl = G[i+1];
        for (int j=1; j<Nml; j++)
                 + one_minus_omega * Gi[j];
    }
}
```

X10 version (rank-specific):

```java
region R = [0:M-l,0:N-1]; double[] G = new double[R];
region R_inner = [1:M-2,1:N-2]; // Subregion of R
for (int p=0; p<num_iterations; p++) {
    for (point t : R_inner) {
        G[t] = omega_o_four * (G[t+[-1,0]] + G[t+[1,0]]
                          + G[t+[0,-1]] + G[t+[0,1]])
             + one_minus_omega * G[t];
    }
}
```

X10 version (rank-independent):

```java
region R_inner = ...; // Inner region as before
region stencil = ...; // Set of points in stencil
double omega_factor = ...; // Weight used for stencil points
for (int p=0; p<num_iterations; p++) {
    for (point t : R_inner) {
        double sum = one_minus_omega * G[t];
        for (point s : stencil) sum += omega_factor * G[t+s];
        G[t] = sum;
    }
}
```

Figure 4.3: Java Grande SOR benchmark
stencil reuse improves productivity, it introduces performance overheads. However, compiler optimizations [55] can reduce this overhead.

### 4.2 Improving the Performance of X10 Language Abstractions

This section has two areas of focus. First, we discuss a compiler optimization we employ to reduce the overhead of using *points* in X10. Second, we describe our rank analysis which can be augmented with X10's *dependent type system* to further improve code generation. As an example, Figure 4.4 contains a simple code fragment illustrating how X10 arrays may be indexed with points in lieu of loop indices. Figure 4.5 shows the unoptimized Java output generated by the reference X10 compiler [101] from the input source code in Figure 4.4. The *get* and *set* operations inside the *for* loop are expensive, and this is further exacerbated by the fact that they occur within an innermost loop.

To address this issue, we have developed an optimization that is a form of *object inlining*, specifically tailored for value-type objects. Object inlining [16, 19, 36, 37] is a compiler optimization for object-oriented languages that transforms objects into primitive data, and the code that operates on objects into code that operates on inlined data. Budimlić [16] and Dolby [36] introduced object inlining as an optimization for Java and C++. General object inlining requires complex escape analysis and concrete object and rank-specific array type inference, and the transformation is irreversible (once unboxed, objects in general cannot be "reboxed").

However, because points in X10 are value types, we can safely optimize all array accesses utilizing point objects by replacing them with an object inlined point array access version. A value object has the property that once the program initializes the object, it cannot subsequently modify any of the object's fields. This prevents the possibility of the code modifying *point p* in Figure 4.4 in between the assignments – a situation that would prevent the inlining of the point. As a result, we can inline the
region arrayRegion1 = [0:datasizes_nz[size]-1];
...
//X10 for loop
for (point p : arrayRegion1) {
    row[p] = rowt[p];
    col[p] = colt[p];
    val[p] = valt[p];
}

Figure 4.4: X10 source code of loop example adapted from the Java Grande sparse-matmult benchmark

//X10 for loop body translated to Java
for ... {
    ... // Includes code to allocate a new point object for p
    (row).set(((rowt).get(p)),p);
    (col).set(((colt).get(p)),p);
    (val).set(((valt).get(p)),p);
}

Figure 4.5: Source code of loop following translation from unoptimized X10 to Java by X10 compiler

point object declared in the for loop header. In addition, we can also perform reboxing on an inlined point when a method invocation expects a point object. Figure 4.6 shows the results of applying this point optimization to the loop we introduce in Figure 4.4, and Figure 4.7 shows the resulting Java code.

4.3 Point Inlining Algorithm

We perform a specialized version of object inlining [16] to inline points. There are two main differences between points and the objects traditionally considered as candidates
for object inlining. First, a point variable can have an arbitrary number of fields because a programmer may use points to access arrays of different rank. Second, a point variable may appear in an X10 loop header. Consequently, the specialized object inlining algorithm must transform the X10 loop header by using the inlined point fields as loop index variables. As a result, this may lead to nested for loops if the point variable is a multi-dimensional point.

Figures 4.8 and 4.9 show the rank inference and point inlining algorithms. We first use the rank inference algorithm to discover the rank of as many X10 points in the program as possible. Recall, developers may omit rank information when declaring
Input: X10 program

Output: rank, a mapping of each X10 array, region and point to its rank

begin
  // initialization
  worklist = ∅, def = ∅
  foreach n ∈ Region, Point, Array do
    rank(n) = T
    worklist = worklist + n
  foreach assign a do
    if a.rhs ∈ constant then
      rank(a.lhs) = a.rhs
      def(a.rhs) = def(a.rhs) ∪ a.lhs
  foreach call arg c → param f do
    if c ∈ constant then
      rank(f) = c
      def(c) = def(c) ∪ f
  // infer ranks
  while worklist ∉ ∅ do
    worklist = worklist − n
    foreach v ∈ def(n) do
      if rank(n) < rank(v) then
        rank(v) = rank(n)
        worklist = worklist + v
        foreach e in def(v) do worklist = worklist + e
      else if rank(n) ≠ rank(v) then
        rank(v) = ⊥
        worklist = worklist + v
        foreach e in def(v) do worklist = worklist + e
  end

Figure 4.8: Rank Analysis Algorithm
Input: X10 program
Output: inlined points

begin
  // flow-insensitive point inlining algorithm
  // inlined points
  foreach AST node n do if get.rank(n) == CONSTANT then
    switch typeof(n) do
      case pointdeclaration
        inline(n)
      case methodcallarg
        reconstruct_point(n)
      case pointreference
        inline(n)
      case X10loop
        convert_loop(n)
  end

Figure 4.9: Algorithm for X10 point inlining

X10 points. However, we need to infer rank information to inline the point. We obtain rank information for points from both point assignments, array accesses, and array domain information found in X10 loop headers. Then we use safety analysis to discover which points we can safely inline. Because points have the value type property, we inline/unbox every safe point with an inferred rank. When encountering method calls that need a point as an actual parameter, we reconstruct the inlined point by creating a new point instance, but ensure that this overhead is only incurred on paths leading to the method calls by allowing the code to work with both original and unboxed versions of the point. Finally, when possible, we convert a point-wise X10 loop into a set of nested for loops using the X10 loop's range information for each dimension in the region.
// X10 array declarations with dependent type information

// rank==1 ==> array is one-dimensional
// rect ==> array's region is dense (rectangular)
// zeroBased ==> lower bound of array's region is zero

double[:, rank==1 && rect && zeroBased] row = ... ;

region arrayRegion1 = [0:temp-1];

// X10 for loop
for (point p : arrayRegion1) {
    row[p] = rowt[p];
    col[p] = colt[p];
    val[p] = valt[p];
}

Figure 4.10: X10 for loop example from Figure 4.4, extended with dependent type declarations

4.4 Use of Dependent Type Information for Improved Code Generation

When examining the Java code generated for the optimization example discussed in the previous section (Figure 4.7) we see that even though the point object has been inlined, significant overheads still remain due to the calls to the get/set methods. These calls are present because by default, the declaration of an array reference variable in X10 does not specify the rank (or dimension sizes) of its underlying array. This makes it possible to write rank-independent code in X10, but poses a challenge for the compiler to generate efficient rank-specific code. In this example, all regions and array accesses are one-dimensional, so it should be possible for the compiler to generate code with direct array accesses instead of method calls. One solution is to use the dependent type system [51] of the X10 language specification [79] to enable the programmer to annotate selected array variable declarations with additional informa-
Figure 4.11: Source code for loop body translated from X10 to Java by X10 compiler.

```java
//X10 optimized for loop translated to Java
for (int i = 0; i <= temp-1; i +=1) {
    ((DoubleArray_c)row).arr_[i]=((DoubleArray_c)rowt).arr_[i];
    ((DoubleArray_c)col).arr_[i]=((DoubleArray_c)colt).arr_[i];
    ((DoubleArray_c)val).arr_[i]=((DoubleArray_c)valt).arr_[i];
}
```

To illustrate this approach, Figure 4.10 contains an extended version of the original X10 code fragment in Figure 4.4 with a dependent type declaration shown for array `row`. Similar declarations need to be provided for the other arrays as well. The X10 compiler ensures the soundness of this type declaration i.e., it does not permit the assignment of any array reference to `row` that is not guaranteed to satisfy the properties. We extended the code generation performed by the reference X10 compiler [101] to generate the optimized code shown in Figure 4.11 for array references with the appropriate dependent type declaration. One drawback to relying solely on the dependent type solution is the performance costs remaining due to our compiler introducing the casts and indirect field accesses to the backing array `arr_` as shown in Figure 4.11. Ideally, this dependent type solution should be used to augment a compiler that deduces rank information automatically (e.g., by propagating rank information from the array's allocation site to all its uses). We present our automatic compiler interprocedural rank inference technique in the next section.
4.5 X10 General Array Conversion

The algorithm for converting general X10 arrays into an efficient lower-level implementation consists of three phases. The first phase, Rank Analysis, infers the concrete ranks of all the X10 arrays in the program and is described in Section 4.6. The second phase, Safety Analysis, determines which X10 arrays can be safely converted into Java arrays, using the rank information computed in Phase 1, and is described in Section 4.7. Extensions to safety analysis that were designed but not used in our experimental results are summarized in Section 4.8. The last phase of the algorithm is the actual conversion of the code manipulating X10 arrays into code operating directly on the underlying Java arrays (Section 4.9).

4.6 Rank Analysis

This section describes the type inference algorithm that we use to discover the ranks of X10 arrays. Recall, the generality of X10 arrays enables programmers to develop rank independent code by omitting array dimensionality at the declaration site. Our whole-program analysis first uses intraprocedural analysis to capture local rank information from array assignments. We then perform interprocedural analysis to obtain rank information arising from both X10 array method arguments and methods returning X10 arrays. Figure 4.8 shows our rank inference algorithm.

The rank information flows from right to left in the rank inference algorithm. That is to say, in an assignment, the inferred rank of the left hand side is the lower (in the type lattice sense) of the rank of the right hand side and the previous rank of the left hand side. Similarly for a method call (in which the parameter passing can be conceptually thought of as assignments of actual parameters to formal parameters), the rank information flows from actual to formal parameters.

The rank inference algorithm can be implemented to run in $O(|V| + |E|)$ time, where $V$ is the set of array, point, and region variables in the whole program and $E$ is the set of edges between them. An edge exists between two variables if one defines the
other. Theorem 4.6.1 and its proof demonstrate that this algorithm has complexity $O(|V| + |E|)$ and preserves program correctness:

**Definition** A graph is a pair $G=(V, E)$ where:

1. $V$ is a finite set of nodes.
2. $E$ are edges and are a subset of $V \times V$.

**Definition** A lattice is a set $L$ with binary meet operator $\wedge$ such that for all $i, j, k \in L$:

1. $i \wedge i = i$ (idempotent)
2. $j \wedge i = i \wedge j$ (commutative)
3. $i \wedge (j \wedge k) = (i \wedge j) \wedge k$ (associative)

**Definition** Given a lattice $L$ and $i, j \in L$, $i < j$ iff $i \wedge j = i$ and $i \neq j$

**Definition** Given a program $P$, let $T$ be the set containing point, region and array types in $P$ and $N$ be the set of variables in $P$ with type $t \in T$ such that for all $m \in N$:

1. $\text{DEF}(m)$ is the set of variables in $P$ defined by $m$.
2. $\text{RANK}(m)$ is the dimensionality associated with $m$. Each $\text{RANK}(m)$ has a lattice value. There exists a precise rank for $m$ iff $\text{RANK}(m) \neq \top$ or $\bot$.

**Theorem 4.6.1.** Given a directed graph $G$ where $V$ is the set of program variables of type array, region, or point, there exists an edge $(i, j)$ between variables $i, j \in V$ iff $j \in \text{DEF}(i)$. The rank analysis algorithm runs in time $O(V+|E|)$ and preserves program correctness.

**Proof.** Initially each node $n \in V$ is placed on the worklist with lattice value $\top$. Once node $n$ is taken off the worklist, $n$ can only be put back on the list iff $n \in \text{DEF}(m)$ and $m < n$ or there $\exists$ a precise rank for both $n$ and $m$ and $\text{RANK}(n) \neq \text{RANK}(m)$. In the latter case $\text{RANK}(n) \leftarrow \bot$ before we place $n$ back on the worklist. Figure 4.12 shows the rank lattice. Since the lattice is bounded and a node $n$ can only have its
lattice value lowered, each node can only be placed on the worklist a maximum of 3 times. Because we traverse source node edges when lattice value changes, each edge will be traversed a maximum of 2 times. Therefore, because \( V \) is a finite set of nodes, the algorithm must eventually halt. Since each node \( n \) is placed on the worklist a maximum of 3 times and its edges are traversed a maximum of 2 times, the complexity is \( O(V+E) \). Assuming the whole program is available to the rank analysis algorithm, the algorithm preserves program correctness. The rank algorithm will produce an incorrect program iff the algorithm assigns an incorrect precise rank to a program variable with type array, region, or point. This would only occur when the variable can have multiple ranks. However, when a variable has multiple ranks, the rank analysis algorithm assigns the variable \( \perp \). Therefore, the rank analysis algorithm produces a correct program.

While rank information, in general, flows from right to left in our rank algorithm,
Figure 4.13: X10 code fragment adapted from JavaGrande X10 montecarlo benchmarks showing when our rank inference algorithm needs to propagate rank information left to right.

we provide an extension allowing rank information to flow from left to right. This bi-directional propagation is useful in this context because the extended X10 compiler performs code generation by translating X10 code into Java. As a result, our X10 array rank analysis extension propagates left hand side assignment rank information to the right since we are performing code generation by translating rank-independent code to rank-specific code; thereby requiring the rank on left and right side of the assignment to be equal. Assuming a compiler performs dead code elimination, our extended analysis will not discover the precise ranks for more arrays than right to left rank propagation. Figure 4.13 shows an example of rank information flowing from left to right.
4.7 Safety Analysis

In addition to gathering precise rank information, our type inference algorithm also employs a safety analysis algorithm to ensure that it is safe to transform an X10 general array into a more efficient representation. The alternate representation used in this dissertation is the Java array. An X10 array is marked as unsafe if an operation is performed on it that cannot be supported by Java array operations.

Figure 4.15 shows the high-level description of the safety analysis algorithm we perform before transforming X10 arrays to Java arrays. The safety analysis algorithm can be implemented to run in $O(|V| + |E|)$ time, where $V$ is the set of array, point, and region variables in the whole program and $E$ is the set of edges between them. An edge exists between two variables if one defines the other. Theorem 4.7.1 and its proof illustrate that this algorithm has complexity $O(|V| + |E|)$ and preserves program correctness:

Definition $SAFE(i)$ is $\top$ iff $i \in V$ and we can either:

1. convert $i$ into a Java array if $i$ is an X10 array
2. convert $i$ into a set of size variables to potentially initialize a Java array if $i$ is a region
   otherwise, it is $\bot$.

Theorem 4.7.1. Given a directed graph $G$ where $V$ is the set of program variables of type array or region, there exists an edge between $i, j \in V$ where $i$ is the source and $j$ is the sink iff $j \in DEF(i)$. The safety analysis algorithm runs in time $O(V+E)$ and preserves program correctness.

Proof. Initially each node $n \in V$ is placed on the worklist with $SAFE(n) = \top$. Once node $n$ is taken off the worklist, $n$ can only be put back on the list iff $n \in DEF(m)$ and $SAFE(m) < SAFE(n)$. Since the lattice is bounded (i.e. can only be $\top$ or $\bot$) and a node $n \in V$ can only have its lattice value lowered, each node can only placed on the worklist a maximum of 2 times. Therefore, because $V$ is a finite set
of nodes, the algorithm must eventually halt. The complexity is $O(V+E)$ since we place only the sink nodes of a source node whose lattice value is lowered on the worklist. Assuming the whole program is available to the safety analysis algorithm, the algorithm preserves program correctness. The safety algorithm will produce an incorrect program iff the algorithm assigns a final lattice value of $(\text{safe})T$ to an unsafe program variable. This would only occur when the lattice value of a variable was not updated. However, since all edges are updated when the lattice value changes, all variables will have the correct lattice value. Therefore, the safety analysis algorithm produces a correct program.

One detail worth mentioning is that our algorithm performs a bi-directional safety inference. We utilize safety information on the left hand side of an assignment to infer safety information for the right hand side and vice versa, thereby reducing safety analysis to an equivalence partitioning problem. Figure 4.14 highlights the importance of the bi-directional safety inference. Our algorithm incorporates this bi-directional strategy for method arguments and formal parameters as well.

4.8 Extensions for Increased Precision

The Rank and Safety analysis algorithms as presented in this section are fairly easy to understand and implement as linear-time flow-insensitive and context-insensitive algorithms. We have also designed more complex flow-sensitive and context-sensitive versions of these algorithms summarized in this section that can potentially compute more precise rank and safety information, leading to better optimization.

For the set of applications we used as benchmarks in this paper these extensions do not produce more precise results, thus we chose to omit a more detailed discussion of these extensions and only include a brief summary here.

Use of SSA Form: The Rank Analysis and Safety Analysis algorithms described on Figures 4.8 and 4.15 are flow insensitive. Thus, if an array variable $a$ is reassigned an array of a different rank than before, it will get $\bot$ as its rank, which can further
// code snippet adapted from JavaGrande X10 Montecarlo
// benchmark to show benefit of bi-directional safety inference
int this.nTimeSteps = 1000;
region r1 = [1:nTimeSteps-1]; // non-zero based
region r2 = [0:nTimeSteps-2];
double[.] pathVal1;
double[.] pathVal2;
...
this.pathVal1 = new double[r1]; // r1 is not safe
this.pathVal2 = new double[r2]; // r2 is safe
...
// propagate safety info left to right in
// set_pathValue to ensure pathVal2 is marked unsafe
set_pathValue(pathVal2);
...
public void set_pathValue(double[.] pv3) {
    this.pathVal1 = pv3; // pv3 not safe
}

Figure 4.14: X10 code fragment adapted from JavaGrande X10 montecarlo benchmarks showing when our safety inference algorithm needs to propagate rank information left to right.
Input: X10 program

Output: safe, maps each X10 array, region and point to safe to transform lattice value

begin

// initialization
worklist = ∅, def = ∅

foreach n ∈ Region, Point, Array do
-safe(n) = T
-worklist = worklist + n

foreach a ∈ Region, Array do
-if a ∉ rect ∧ zero then
- safe(a) = ⊥

foreach array access with array p ∈ Point do
-if index i ∉ constant then
-safe(p) = ⊥

foreach assign a do def(a.rhs) = def(a.rhs) ∪ a.lhs

foreach call arg c → param f do def(c) = def(c) ∪ f

// infer X10 safety transform value

while worklist ≠ ∅ do
-worklist = worklist − n

foreach v ∈ def(n) do
-if safe(n) < safe(v) then
-safe(v) = safe(n)
-worklist = worklist + v

foreach e in def(v) do worklist = worklist + e

end

Figure 4.15 : Safety Analysis Algorithm
get propagated to other variables involved in computation with \( a \). Similarly, if a variable is marked unsafe for conversion into a Java array, it will prevent conversion of all occurrences of that variable into a Java array, even if they could potentially be safely converted in different regions of the code. This source of imprecision can be eliminated by converting the code into SSA form [14]. The \( \phi \) nodes in the SSA form are treated similarly to an assignment: the rank of the variable on the left hand side gets assigned a \( \text{merge}() \) of the ranks of all the argument variables to the \( \phi \) function. Since rank analysis does not involve any code reorganization, conversion from the SSA form back into the original form is simple and doesn’t involve any copy coalescing [17].

**Type Jump Functions:** The two algorithms, as described here, can propagate rank and safety information through infeasible paths in the call graph. If a method is called at one site with an argument of rank 2, and at another site with an argument of rank 1, the formal array parameter will receive \( \bot \) as its rank, and it may then propagate this lower type through the return variable back into the caller code.

This imprecision can be avoided by using *type jump functions* [27] for method calls. The idea behind type jump functions is to encapsulate the relation between the types of actual arguments to a method and the type of the return argument. Since rank and safety information are essentially types, this method generalization can be used to increase the precision of the rank and safety analysis algorithms. If a type jump function describes a method \( m \) as accepting the argument of rank \( R \) and returning a value of rank \( R - 1 \), then this method can be analyzed independently at different call sites and will propagate the correct values for the rank, even if the ranks of the arguments at different call sites are different.

During the conversion of X10 arrays into Java arrays, a method with polymorphic rank arguments has to be cloned to variants with the specific ranks that are determined by the call site. The most aggressive approach is to convert as many X10 arrays as possible by generating as many variants of the method as there are call sites with
different sets of ranks for actual arguments. Alternatively, to avoid code explosion, the compiler can generate a limited set of variants for the most profitable call paths, and leave the default variant that uses unconverted X10 arrays for the general case.

Type jump functions for the safety analysis, while similar to those for rank analysis, are simpler since the only two “types” a variable can have are safe and unsafe.

4.9 Array Transformation

Once we have completed the array rank and safety analysis, we begin the transformation from X10 arrays to the more efficient representation (Java array). There are two main steps in this process. First, we convert each declared X10 array to our analyzed precise type. Second, we must convert each such X10ArrayAccess AST node into a Java ArrayAccess AST node. The X10 compiler makes the distinction between these two types of nodes so that only the X10ArrayAccess can accept a point expression as an argument. As a result, during the conversion process, we must also convert any point valued subscript expression into equivalent integer-valued expressions since we cannot perform a Java array access with a point object. We use a variant of the Object Inlining optimization (Section 4.3) to convert the X10 points into integer values [58].

4.10 Object Inlining in Fortress

In Fortress [3], we extend our X10 point inlining algorithm to inline objects of any type. We aggressively inline all variables whose declared object type has not been omitted by the programmer since all object types in Fortress represent leaves in the Fortress type hierarchy (i.e. an object cannot be extended). There are a couple of differences worth highlighting between the point inlining algorithm and the extended Fortress object inlining algorithm. In X10, our point inlining algorithm

\[\text{AST node refers to the Polyglot Abstract Syntax Tree used in the X10 compiler}\]
performs object reconstruction of all inline point method arguments. This solution is effective since points have the value type property (i.e. once defined, points cannot subsequently be modified). In Fortress, instead of reconstructing inlined method arguments, our algorithm synthesizes new methods with inlined formal parameters. In addition, we extend the X10 point inlining algorithm in Fortress to inline arrays of objects by replacing the object array with a set of inlined arrays. Future object inlining work in Fortress includes adding type analysis to identify types for variables with omitted object types to enable optimizations such as object inlining to be more effective.
Chapter 5

Eliminating Array Bounds Checks with X10 Regions

Many high-level languages perform automatic array bounds checking to improve both safety and correctness of the code, by eliminating the possibility of an incorrect (or malicious) code randomly "poking" into memory through an out of bounds array access or buffer overflow. While these checks are beneficial for safety and correctness, performing them at run time can significantly degrade performance especially in array-intensive codes. Two main ways that bounds checks can affect performance are:

1. The Cost of Checks. The runtime may need to check the array bounds when program execution encounters an array access.

2. Constraining Optimizations. The compiler may be forced to constrain or disable code optimizations in code region containing checks, in the presence of precise exception semantics.

Significant effort has been made by the compiler research community to statically eliminate array bounds checks in higher-level languages when the compiler can prove that these checks are unnecessary [13, 77, 94]. In this thesis, we take advantage of the region language construct in X10 to help determine statically when array bounds checks are not needed in accesses to high-level arrays. In such cases, we annotate the array access with a no Bounds Check annotation to signal to a modified version of the IBM J9 Java Virtual Machine [67] ¹ that it can skip the array bounds check for those

¹Any JVM can be extended to recognize the no Bounds Check annotation, but in this thesis our experiences are reported for a version of the IBM J9 JVM that was modified with this capability.
particular array accesses.

X10 regions are particularly suitable for static analysis since they have the value type property (once defined, they cannot subsequently be modified). This simplifies the compiler task since the region remains unchanged over (say) an entire loop iteration space, even if the loop contains unanalyzed procedure calls. For example, consider the following two loops:

```java
double[] a = new double[[b.low,b.high]];  
loop1: for (n=b.low, n <= b.high, n++) {
    foo(b);
    a[n] = ...
}

region r = [b.low : b.high];  
loop2: for (point p : r) {
    foo(r);
    a[p] = ...
}
```

In loop1, in addition to proving that there are no modifications to \( n \) inside of the loop other than those imposed by loop iteration itself, one must also prove that neither \( low \) or \( high \) are changed inside the loop body (e.g., as a result of a call to \( foo() \)) in a manner that might introduce an illegal array access. However, in loop2, this additional analysis is unnecessary since the region bounds are immutable. Figure 5.3 illustrates how X10 regions help array bounds analysis with a code fragment taken from the Java Grande Forum \texttt{sparsematmult} benchmark [54]. In the \texttt{sparsematmult} example, the \texttt{kernel} method performs sparse matrix multiplication. Because our analysis discovers that \texttt{row} and \texttt{col} have the same region, the compiler can apply a transformation that adds an annotation around \texttt{col}'s subscript to signal to the Virtual Machine to skip the bounds check. Inserting this annotation is possible due to the immutability of X10 regions. A standard Java compiler cannot perform this optimization because it
depends on the knowledge that regions are immutable. In addition, determining that col and row share the same region would require interprocedural analysis, which may be challenging for a JIT compiler to perform.

In our approach, we insert the noBoundsCheck annotation around an array subscript appearing inside the loop if the compiler can establish one of the following properties:

1. **Array Subscript within Region Bound.** If the array subscript is a point that the programmer is using to iterate through region \( r_1 \) and \( r_1 \) is a subset of the array's region, then the bounds check is unnecessary.

2. **Subscript Equivalence.** Given two array accesses, one with array \( a_1 \) and subscript \( s_1 \) and the second with array \( a_2 \) and subscript \( s_2 \): if subscript \( s_1 \) has the same value number as \( s_2 \), \( s_1 \) executes before subscript \( s_2 \) and array \( a_1 \)'s region is a subset of \( a_2 \)'s region, then the bounds check for \( a_2[s_2] \) is unnecessary.

In Section 7, we show the effects of applying this transformation to a set of benchmarks. The novel contributions to eliminating bounds checks are the following:

- **Building Array Subset Region Relationships.** Programmers often define arrays in scientific codes over either the same domain or a domain subset. Our array region analysis enables us to discover when the domain of one array is a subset of the other. This information is useful in eliminating bounds checks when indexing two arrays with the same index value. Even if we cannot prove that the first check is superfluous, we can establish redundancy for the second one.

- **Region Algebra.** The idea of introducing region algebra is to expose computations involving variables with inherent region associations; thereby proving that the result of the computation may also become a region association. A defined variable reference has a region association if the variable satisfies one of the following properties:
1. The variable has type \textit{region} or is an X10 general array.

2. The variable has type \textit{point} and appears in the X10 loop header.

3. Program execution assigns the variable a region bound or an offset of the region bound (e.g. \texttt{int i = rl.rank(0).high()+1}, where \texttt{rl} is a region, 0 indicates that the bound will be taken from the first dimension, and the offset is 1, \texttt{i} in this example has a region association).

Only variables of type point, region, X10 array, and integer can have a region association. We use interprocedural analysis to propagate these region associations, allowing us to catch opportunities to eliminate bounds checks that the JIT compiler would miss due to both a lack of knowledge about region immutability semantics and the absence of interprocedural bounds analysis in today’s JIT compiler technology. We principally use the region inequalities \( n-k \geq l \) and \( n+k \leq h \) where \( n, k \) are variables with region associations, \( k \geq 0 \), \( l \) and \( h \) are respectively the low and high bounds of a region. We apply these inequalities to cases when \( k \) resolves to either a constant or a region where the rank is 1. In practice, we often discover cases that enable us to further simplify the inequality expressions such as when \( n \) is a region high bound in the first inequality and the region lower bound is 0. In this case, all we must prove is that \( k \) represents a sub region of \( n \)'s region to establish a resulting region association.

- \textit{Array Element Value Ranges}. Discovering an array’s range of values can expose code optimization opportunities. Barik and Sarkar’s [12] enhanced bit-aware register allocation strategy uses array value ranges to precisely determine how many bits a scalar variable requires when it is assigned the value of an array element. In the absence of sparse data structures [71], sparse matrices in languages like Fortran, C, and Java are often represented by a set of 1-D arrays that identify the indices of non-zero values in the matrix. This representation usu-
ally inhibits standard array bounds elimination analysis because array accesses often appear in the code with subscripts that are themselves array accesses. We employ value range analysis to infer value ranges for arrays. Specifically, our array value range analysis tracks all assignments to array elements. We ascertain that when program execution assigns an array's element a value using the \textit{mod} function, a loop induction variable, a constant, or array element value, we can analyze the assignment and establish the bounds for the array's element value range. \footnote{Note: when array \textit{a1} is an alias of array \textit{a2} (e.g. via an array assignment), we assign both \textit{a1} and \textit{a2} a value range of \( 1 \), even if \textit{a1} and \textit{a2} share the same value range, in order to eliminate the need for interprocedural alias analysis. In the future, value range alias analysis can be added to handle this case.}

- \textit{Value Numbering to Discover Redundant Array Accesses.} We use a dominator-based value numbering technique \cite{15} to find redundant array accesses. We annotate each array access in the source code with two value numbers. The first value number represents a value number for the array access. We derive a value number for the array access by combining the value numbers of the array reference and the subscript. The second value number represents the array's element value range. By maintaining a history of these array access value numbers we can discover redundant array accesses.

- \textit{Using Multiple Code Views to Enhance Bounds Elimination Analysis.} We maintain two code views. The first is the source code view which the compiler uses to perform code generation. The second is the analysis code view which we employ as an abstraction to derive array access bounds checking information. The second view is helpful to both prune useless source code information and ease the burden of assigning region value numbers to variables during the analysis phase. For example, X10 loops are transformed into straight line code blocks.
We generate a loop header point assignment to the loop header region and place it as the first statement in the code block. When a programmer assigns an array element a value using the $mod$ function, we transform the analysis code view by replacing the original assignment with an assignment to a region constructor where the low bound is 0 and the high bound is the expression on the right hand side of the $mod$ function - 1. Figure 5.1 provides an example displaying both the source view and analysis view of the code. Altering the analysis code view conveniently enhances our elimination bounds analysis without modifying the source code and affecting code generation.

- **Interprocedural Region Analysis with Return Jump Functions.** Using the idea of return type jump functions taken from McCosh [27], we can uncover cases when a method returns a region that the program passes as a method argument. We transform the analysis code view by replacing the method call with the region argument. As a result, even though the method call's formal parameter region can be $\bot$, the variable on the left hand side of the assignment may resolve to a more precise region.

- **Demonstrating Array View Productivity Benefits.** We illustrate the development productivity benefit of using array views with a *hexahedral cell* code example [47, 60]. Array views give the programmer the opportunity to work with multiple views of an array. In practice, programmer's commonly utilize multiple array views when they want to manipulate a single array row. We show the productivity benefit of using array views to switch between a multi-dimensional and linearized view of the same array.

- **Interprocedural Linearized Array Subscript Bounds Analysis.** In general, programmers create linearized arrays to avoid the performance costs incurred when

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3As discussed later, array views are different from source and analysis code views.
using a multi-dimensional array representation. However, the linearized subscripts can be an impediment to bounds check elimination. To mitigate this issue, we have the compiler automatically reconstruct multi-dimensional arrays from the linearized array versions. This "delinearization" transformation can enable array bounds analysis using the multi-dimensional array regions. Delinearization has also been proposed in past work on dependence analysis [70]. However, due to the difficulty in automatically converting some linearized arrays to their multi-dimensional representation, we must perform array bounds analysis on linearized array subscripts to glean domain iteration information which we subsequently employ to reduce the number of bounds checks. We extend this analysis interprocedurally by summarizing local bounds analysis information for each array.

- **Improving Runtime Performance.** Using our static array bounds analysis and automatic compiler annotation insertion to signal the VM when to eliminate a bounds check, we have improved sequential runtime performance by up to 22.3% over JIT compilation.

### 5.1 Intraprocedural Region Analysis

Our static bounds analysis first runs a local pass over each method after we translate the code into a static single assignment form (SSA) [14]. Using a dominator based value numbering technique [15], we assign value numbers to each point, region, array, and array access inside the method body. These value numbers represent region association. Upon completion of local bounds analysis, we map region value numbers back to the source using the source code position as the unique id. Figure 5.6 shows the algorithm for the intraprocedural region analysis.

To perform the analysis and transformation techniques described above, we use the Matlab D framework developed at Rice University [27, 44]. We generate an XML file
Source view:

```java
1 region r = [0:99];
2 double[] a = new double[r];
3 double[] b = bar(r);
4 for (point p1 : r)
5   b[p1] = foo(new Random()) % 100;
6 for (point p2 : r)
7   a[p2] = b[p2];
8 //generates random number
9 int foo(Random rand) {
10   return rand.nextInt();
11 }
12 double[] bar(region r) {
13   return new double[r];
14 }
```

Analysis view:

```java
1 r = [0:99];
2 a = r;
3 b = r; //replaced call with returned region argument r
4 p1= r; //next array access: subscript, array share value number
5 b[p1] = [0:99]; //determines value range for b
6 p2 = r; //next array access: subscript, array share value number
7 a[p2] = b[p2]; //determines value range for a
8 //generates random number
9 int foo(Random rand) {
10   return rand.nextInt();
11 }
12 region bar(region r) {
13   return r; //returns formal parameter
14 }
```

Figure 5.1: Example displaying both the code source view and analysis view. We designed the analysis view to aid region analysis in discovering array region and value range relationships by simplifying the source view.
Figure 5.2: X10 region analysis compiler framework
from the AST of the X10 program, then read this AST within the Matlab D compiler, convert it into SSA, perform the value numbering based algorithms presented in this chapter to infer the regions associated with arrays, points and regions in the program, then use the unique source code position to map the analysis information back into the X10 compiler. Figure 5.2 summarizes the compiler framework we use for region analysis.

We build both array region and value region relationships during the local analysis pass. An array will have a value region if and only if we can statically prove that every value in the array lies within the bounds of a region. For example, in Figure 5.3, assuming that the assignment of array values for row is the only row update, analysis will conclude that row's value region is reg1. Our static bounds analysis establishes this value region relationship because the mod function inherently builds the region \([0:\text{reg1.high}]\). Figure 5.4 shows this analysis code view update for array element assignments to row and col.

We use an implicit, infinitely wide type lattice to propagate the values of the regions through the program. The lattice is shown on Figure 5.5. In the Matlab D compiler [44], a \(\phi\) function performs a meet operation (\(\land\)) of all its arguments and assigns the result to the target of the assignment.

### 5.2 Interprocedural Region Analysis

If a method returns an expression with a value number that is the same as a formal parameter value number, analysis will give the array assigned the result of the method call the value number of the corresponding actual argument at the call site.

Static interprocedural analysis commences once intraprocedural analysis completes. During program analysis, we work over two different views of the code. The first is the standard source view which affects code generation. The second is the analysis view. Changes to the analysis view of the code do not impact code generation. In Figure 5.3, program execution assigns array \(x\) the result of invoking method
//code fragment is used to highlight
//interprocedural array element value
//range analysis

... 

region reg1 = [0:dm[size]-1];
region reg2 = [0:dn[size]-1];
region reg3 = [0:dp[size]-1];
double[.] x = randVec(reg2);
double[.] y = new double[reg1];
int[.] val = new double[reg3]
int[.] col = new double[reg3];
int[.] row = new double[reg3];
Random R;

for (point p1 : reg3) {
    //array row has index set in reg3 and value range in reg1
    row[p1] = Math.abs(R.Int()) % (reg1.high()+1);
    col[p1] = Math.abs(R.Int()) % (reg2.high()+1);...
}

kernel(x,y,val,col,row,..);

double[.] randVec(region r1){
    double[.] a = new double[r1];
    for (point p2: r1)
        a[p2] = R.double();
    return a;
}

kernel(double[.]x,double[.]y,int[.]val,int[.]col,int[.]row,..){
    for (point p3 : col)
        y[row[p3]]+= x[col[p3]]*val[p3];
}

Figure 5.3: Java Grande Sparse Matrix Multiplication kernel (source view).
//code fragment is used to highlight
//interprocedural array element value
//range analysis

reg1 = [0:dm[size]-1];
reg2 = [0:dn[size]-1];
reg3 = [0:dp[size]-1];
x = reg2; //replaced call with region argument reg2
y = reg1;
col = reg3;
row = reg3;

Random R;...
p1 = reg3; //replaced X10 loop with assignment to p1
row[p1] = [0:reg1.high()]; //followed by loop body
col[p1] = [0:reg2.high()]; //created value range from mod
kernel(x,y,col,row,..);

region randVec(region r1){
a = r1;
p2 = r1; //replaced X10 loop with assignment to p2
a[p2] = R.double(); //followed by loop body
return r1; //returns formal parameter
}

kernel(double[.]x,double[.]y,int[.]col,int[.] row,..){
p3 = col; //replaced X10 loop with assignment to p3
y[row[p3]]+= x[col[p3]]*val[p3]; //followed by loop body
}

Figure 5.4 : Java Grande Sparse Matrix Multiplication kernel (analysis view).
Random Vector. Because our analysis determines that the method will return the region the program passes as an argument (assuming region has a lower bound of 0), we will modify the analysis view by replacing the method call with an assignment to the argument (reg2 in our example). Figure 5.4 shows this update. When encountering method calls which our interprocedural regions analysis is not currently analyzing, we assign each formal argument to the actual argument if and only if the actual argument has a region association. Each actual argument can have one of the following three region states:

- If the method argument is a X10 array, region, or point, then the argument will be in the full region state.

- The method argument has a partial region state when it represents the high or low bound of a linear region.
Input: X10 program

Output: region, a local mapping of each X10 array, region and point to its region value number

begin

// initialization
foreach CFG node c do
    foreach n ∈ Region, Point, Array do
        region(n) = T

// infer X10 region mapping
foreach a ∈ assign do
    if a ∈ φ function then
        region(a.def) ← \land_{i=0}^{a.numargs} region(a.arg(i))
    else if a.rhs ∈ constant then
        region(a.lhs) = a.rhs
    else
        region(a.rhs) = region(a.rhs)

foreach a ∉ assign do
    region(a) = a;

end

Figure 5.6: Intraprocedural region analysis algorithm builds local region relationships.
• If the method argument does not fall within the first two cases, then we assign \( \bot \) to the argument (no region association). This distinction minimizes the number of variables that we need to track during region analysis.

In addition to analyzing the code to detect region equivalence, we augment the analysis with extensions to support sub-region relationships. Inferring sub-region relationships between arrays, regions and points is similar in structure to region equivalence inference analysis, but is different enough to warrant a separate discussion. As with the interprocedural region equivalence analysis, there is an implicit type lattice, but this time the lattice is unbounded in height as well as in width. The lattice is shown on Figure 5.7.

The lattice is defined as follows: there is an edge between regions \( A \) and \( B \) in the lattice if and only if the two regions are of the same dimensionality, and region \( A \) is completely contained within region \( B \). During our analysis, we compute on demand an approximation of the relation on Figure 5.7; if we cannot prove that a region \( A \) is a sub-region of region \( B \), then \( A \wedge B = \bot \).

In addition, our analysis is flow-insensitive for global variables. When static analysis determines that a global variable might be involved in multiple region assignments involving different regions, the region for the variable becomes \( \bot \). In the future, we can extend the algorithm to assign the variable the region intersection instead of \( \bot \). Figure 5.8 presents pseudo code for the static interprocedural region analysis algorithm. The interprocedural region analysis algorithm can be implemented to run in \( O(|V| + |E|) \) time, where \( V \) is the number of array, point, and region variables in the whole program and \( E \) is the number of edges between them. An edge exists between two variables if one defines the other. Theorem 5.2.1 and its proof shows that this algorithm has complexity \( O(|V| + |E|) \) and preserves program correctness:

**Definition** A graph is a pair \( G=(V, E) \) where:

1. \( V \) is a finite set of nodes.
2. \( E \) are edges and are a subset of \( V \times V \).
Figure 5.7: Type lattice for sub-region relation
**Definition** A lattice is a set $L$ with binary meet operator $\land$ such that for all $i, j, k \in L$:

1. $i \land i = i$ (idempotent)
2. $j \land i = i \land j$ (commutative)
3. $i \land (j \land k) = (i \land j) \land k$ (associative)

**Definition** Given a lattice $L$ and $i, j \in L$, $i < j$ iff $i \land j = i$ and $i \neq j$.

**Definition** Given a program $P$, let $T$ be the set containing point, region and array types in $P$ and $N$ be the set of variables in $P$ with type $t \in T$ such that for all $m \in N$:

1. $\text{DEF}(m)$ is the set of variables in $P$ defined by $m$.
2. $\text{REG}(i)$ is the region associated with $i$. There $\exists$ precise region for $i$ iff $i \in V$ and $\text{REG}(i) \neq \top$ or $\bot$.

**Theorem 5.2.1.** Given a directed graph $G$ where $V$ is the set of program variables of type array or region, there exists an edge $E$ between $i, j \in V$ where $i$ is the source and $j$ is the sink iff $j \in \text{DEF}(i)$. The region analysis algorithm runs in time $O(V+E)$ and preserves program correctness.

**Proof.** Initially each node $n \in V$ is placed on the worklist with lattice value $\top$. Once node $n$ is taken off the worklist, $n$ can only be put back on the list iff $n \in \text{DEF}(m)$ and $m < n$ or there $\exists$ precise regions for both $n$ and $m$ and $\text{REG}(n) \neq \text{REG}(m)$. In the latter case $n \leftarrow \bot$ before we place $n$ back on the worklist. Since the lattice is bounded and a node $n$ can only have its lattice value lowered, each node can only be placed on the worklist a maximum of 3 times. Because we traverse source node edges when lattice value changes, each edge will be traversed a maximum of 2 times. Therefore, because $V$ is a finite set of nodes, the algorithm must eventually halt. Since each node $n$ is placed on the worklist a maximum of 3 times and its edges are traversed a maximum of 2 times, the complexity is $O(V+E)$. Note that $i \land j = \bot$ even when $i \subset j$. Assuming the whole program is available to the region analysis algorithm,
the algorithm preserves program correctness. The region algorithm will produce an incorrect program iff the algorithm assigns an incorrect precise region to a program variable with type array or region. This would only occur when the variable can have multiple regions. However, when a variable has multiple regions, the region analysis algorithm assigns the variable \( \bot \). Therefore, the region analysis algorithm produces a correct program.

5.3 Region Algebra

Often in scientific codes, loops iterate over the interior points of an array. If through static analysis we can prove that loops are iterating over sub-regions of an array, we can identify the bounds checks for those array references as superfluous. We use the example on Figure 5.9 to highlight the benefits of employing region algebra to build variable region relationships. Figure 5.10 shows the algorithm for region algebra analysis.

When our static region analysis encounters the `dgefa` method call with a region high bound argument in Figure 5.9, analysis will assign `dgefa`'s formal parameter \( n \) the high bound of `region1`'s second dimension and \( a \) the region `region1`. We shall henceforth refer to the region representing `region1`'s second dimension as `region1.2dim`. Inside `dgefa`'s method body, analysis will categorize `nm1` as a region bound and `region3` as a sub-region of `region1.2dim` when inserting it in the region tree.

Next, we assign array `colk` the region `region1.2dim` and categorize `kp1` as a sub-region of `region1.2dim`. When static region analysis examines the binary expression \( n-kp1 \) on the right hand side of the assignment to `var1`, it discovers that the \( n \) is `region1.2dim.hbound()` and `kp1` is a sub region of `region1.2dim`. As a result, we can use region algebra to prove that this region operation will return a region \( r \) where: \( r.lbound() \geq region1.2dim.lbound() \) and \( r.bound() \leq region1.2dim.hbound() \). Consequently, `var1` will be assigned `region1.2dim`. 
Input: X10 program

Output: region, a mapping of each X10 array, region and point to its region

begin

// initialization
worklist = ∅, def = ∅

foreach n ∈ Region, Point, Array do
    region(n) = T
    worklist = worklist + n

foreach assign a do
    if a.rhs ∈ constant then
        region(a.lhs) = a.rhs
        def(a.rhs) = def(a.rhs) U a.lhs

foreach call arg c → param f do
    if c ∈ constant then
        region(f) = c
        def(c) = def(c) U f

// infer X10 region mapping
while worklist ≠ ∅ do
    worklist = worklist − n
    foreach v ∈ def(n) do
        if region(n) < region(v) then
            region(v) = region(n)
            worklist = worklist + v
            foreach e in def(v) do worklist = worklist + e
        else if region(n) ≠ region(v) then
            region(v) = ⊥
            worklist = worklist + v
            foreach e in def(v) do worklist = worklist + e

end

Figure 5.8: Interprocedural region analysis algorithm maps variables of type X10 array, point, and region to a concrete region.
Finally, analysis determines that var2's region is a sub-region of region1_2dim. As a result, when analysis encounters the daxpy call it will assign daxpy formal parameter dx the region region1_2dim and formal parameter dax_reg the same region as var2 enabling us to prove and signal to the VM that the bounds check for the array access dx[p2] in daxpy's method body is unnecessary.

5.4 Improving Productivity with Array Views

In the Habanero project [50], we have proposed an extention to X10 arrays called array views. A programmer can exploit the array's view to traverse an alternative representation of the array. Prevalent in scientific codes is the expression of the form $a = b[i]$ which often assigns the variable $a$ row $i$ of array $b$ when $b$ is a two-dimensional array. Array views can extend this idea by providing an alternate view for the entire array. The following code snippet shows an array view example:

```java
double[.] ia = new double[[1:10,1:10]]; double[.] v = ia.view([10,10],[1:1]); v[1] = 42; print(ia[10,10]);
```

The programmer declares array $ia$ to be a 2-dimensional array. Next, the programmer creates the array view $v$ to represent a view of 1 element in the array $ia$. This essentially introduces a pointer to element $ia[10,10]$. Subsequently, when the programmer modifies the array $v$, array $ia$ is also modified resulting in the print statement yielding the value 42. We will use a hexahedral cells code [47] as a running example to illustrate the productivity benefits of using array views in practice. Figure 5.11 shows the initialization of multi-dimensional arrays $x$, $y$, and $z$. Note: Only 1 for loop header would be needed (for point $p : reg_{mesh}_{3D}$) if statements appear in only the innermost loop.
// code fragment is used to highlight
// interprocedural region analysis using region algebra
int n = dsizes[size];
int ldaa = n;
int lda = ldaa + 1;
...
region region1 = [0:ldaa-1,0:lda-1];...
double[] a = new double[region1]...
info = dgefa(a, region1.rank(l).high(), ipvt);
// dgefa method, lufact kernel
int dgefa(double[] a, int n, int[] ipvt){...
  nml = n - 1;...
  region region3 = [0:nml-1];...
  for (point p1[k] : region3) {
    col_k = RowView(a,k);...
    kp1 = k + 1...
    int var1 = n-kp1;
    region var2 = [kp1:n];...
    daxpy(var1,col_k,kp1,var2,...);...
  }
}
...
// daxpy method
void daxpy(int n,double[] dx,int dx_off,region dax_reg,...){...
  for (point p2 : dax_reg)
    dy[p2] += da*dx[p2];...
}

Figure 5.9 : Java Grande LU factorization kernel.
Input: X10 program

Output: region, a mapping of each X10 array, region, point and int to its region association

begin

// initialization
worklist = ∅, def = ∅

foreach n ∈ Region, Point, Array, int do
    regAssoc(n) = ⊤
    worklist = worklist + n

foreach assign a do
    if a.rhs ∈ constant ∨ bound then
        regAssoc(a.lhs) = a.rhs
        def(a.rhs) = def(a.rhs) U a.lhs

foreach call arg c → param f do
    if c ∈ constant ∨ bound then
        regAssoc(f) = a.rhs
        def(c) = def(c) U f

// infer X10 region mapping

while worklist ≠ ∅ do
    worklist = worklist - n

foreach v ∈ def(n) do
    if regAssoc(n) < regAssoc(v) then
        regAssoc(v) = regAssoc(n)
        worklist = worklist + v
        foreach e in def(v) do worklist = worklist + e
    else if regAssoc(n) ≠ regAssoc(v) then
        regAssoc(v) = ⊥
        worklist = worklist + v
        foreach e in def(v) do worklist = worklist + e

end

Figure 5.10: Region algebra algorithm discovers integers and points that have a region association.
Figure 5.12 illustrates one problem that arises when programmers utilize an array access as a multi-dimensional array subscript. Since the subscript returns an integer, the developer cannot use the subscript for multi-dimensional arrays. As a result, the programmer must rewrite this code fragment by first replacing the 3-dimensional arrays $x$, $y$ and $z$ with linearized array representations. Subsequently, the developer needs to modify the array subscripts inside the innermost loop of Figure 5.11 with the more complex subscript expression for the linearized arrays. While this solution is correct, we can implement a more productive solution using X10 array views as shown in Figure 5.13. This solution enables programmers to develop scientific applications with multi-dimensional array computations in the presence of subscript expressions returning non-tuple values.

Figure 5.14 shows the result of applying our array transformation described in Section 4.9 to the hexahedral cells code example. The process converts the 3-dimensional X10 arrays into 3-dimensional Java arrays when analysis determines it is safe to do so. This compilation pass does not transform the X10 arrays $x$, $y$, $z$, $xv$, $yv$, and $zv$ because of their involvement in the X10 $array\_view()$ method call. There is not a semantically-equivalent Java method counterpart for the X10 $array\_view()$ method. One drawback of array views as presented is that safety analysis marks the view's target array as unsafe to transform. The array transformation pass does convert the X10 general arrays $p1$ and $p2$ in Figure 5.14 into 3-dimensional Java array representations. Although 3-dimensional array accesses in Java are inefficient, this transformation still delivers more than a factor of 3 speedup over the code version with only X10 general arrays.

We can achieve even better performance by linearizing the 3-dimensional Java arrays. Figure 5.15 displays the code after Java array linearization. The $LinearViewAuto$ call indicates where the compiler has automatically linearized a multi-dimensional X10 array whereas the $LinearViewHand$ method invocation indicates where the programmer has requested a linear view of a multi-dimensional region.
// code fragment is used to highlight
// array view productivity benefit
// create uniform cube of points
region reg_mex = [0: MESH_EXT-1];
region reg_mex_3D = [reg_mex, reg_mex, reg_mex];
double[.] x = new double[reg_mex_3D];
double[.] y = new double[reg_mex_3D];
double[.] z = new double[reg_mex_3D];...
for (point pt3[pz] : reg_mex) {...
  for (point pt2[py] : reg_mex) {...
    for (point pt1[px] : reg_mex) {
      x[pz, py, px] = tx;
      y[pz, py, px] = ty;
      z[pz, py, px] = tz;
      tx += ds;
    }
    ty += ds;
  }
  tz += ds;
}...

Figure 5.11: Hexahedral cells code showing the initialization of multi-dimensional arrays x, y, and z.
// code fragment highlights array view productivity benefit
region reg_mex = [0:MESH_EXT-1];
region reg_mex_linear=[0:MESH_EXT*MESH_EXT*MESH_EXT-1];
double[] x = new double[reg_mex_linear-];
double[] y = new double[reg_mex_linear];
double[] z = new double[reg_mex_linear];...
for (point pt3[pz] : reg_mex) {
    for (point pt2[py] : reg_mex) {
        for (point pt1[px] : reg_mex) {
            // using less productive linearized array access
            x[px+MESH_EXT*(py + MESH_EXT*pz)] = tx;
            y[px+MESH_EXT*(py + MESH_EXT*pz)] = ty;
            z[px+MESH_EXT*(py + MESH_EXT*pz)] = tz;
            tx += ds;
        }
        ty += ds;
    }
    tz += ds;
}
region reg_br = [0:MESH_EXT-2];
region reg_br_3D = [reg_br, reg_br, reg_br];
int[] pl,p2 = new int[reg_br_3D];...
// would be invalid if x, y, and z were 3-D arrays
for (point pt7 : reg_br) {
    ux = x[p2[pt7]] - x[p1[pt7]];
    uy = y[p2[pt7]] - y[p1[pt7]];
    uz = z[p2[pt7]] - z[p1[pt7]];
}

Figure 5.12: Hexahedral cells code showing that problems arise when representing arrays x, y, and z as 3-dimensional arrays due to programmers indexing into these arrays using an array access returning integer value instead of a triplet.
//code fragment highlights array view productivity benefit

region reg_mex = [0:MESH_EXT-1];
region reg_mex_3D = [reg_mex, reg_mex, reg_mex];
double[,] x,y,z = new double[reg_mex_3D];...
for (point pt3[pz] : reg_mex) {...
  for (point pt2[py] : reg_mex) {...
    for (point pt1[px] : reg_mex) {
      x[pz,py,px] = tx; // use productive multi-D
      y[pz,py,px] = ty; // access with array views
      z[pz,py,px] = tz;
      tx += ds;
    }
    ty += ds;
  }
  tz += ds;
}...
region reg_br = [0:MESH_EXT-2];
region reg_br_3D = [reg_br, reg_br, reg_br];
int[,] p1,p2 = new int[reg_br_3D];...
region reg_linear=[0:MESH_EXT*MESH_EXT*MESH_EXT-1];
double[,] xv = x.view([0,0],[0:reg_linear]);
double[,] yv = y.view([0,0],[0:reg_linear]);
double[,] zv = z.view([0,0],[0:reg_linear]);
for (point pt7: reg_br) {
  ux = xv[p2[pt7]] - xv[p1[pt7]];
  uy = yv[p2[pt7]] - yv[p1[pt7]];
  uz = zv[p2[pt7]] - zv[p1[pt7]];
}

Figure 5.13: Array views xv, yv, and zv enable the programmer to productivity implement 3-dimensional array computations inside the innermost loop.
//code fragment highlights X10 to Java array translation

region reg_mex = [0:MESH_EXT-1];
region reg_mex_3D = [reg_mex,reg_mex,reg_mex];
double[] x,y,z = new double[reg_mex_3D];...
for (point pt3[pz] : reg_mex) {...
  for (point pt2[py] : reg_mex) {...
    for (point pt1[px] : reg_mex) {
      x[pz,py,px] = tx; //use productive multi-D
      y[pz,py,px] = ty; //access with array views
      z[pz,py,px] = tz;
      tx += ds;
    }
    ty += ds;
  }
  tz += ds;
}...
region reg_br = [0:MESH_EXT-2];
region reg_br_3D = [reg_br, reg_br, reg_br];
to int[][][] p1,p2 = new int[reg_br_3D];...
region reg_linear=[0:MESH_EXT*MESH_EXT*MESH_EXT-1];
double[] xv = x.view([0,0],[0:reg_linear]);
double[] yv = y.view([0,0],[0:reg_linear]);
double[] zv = z.view([0,0],[0:reg_linear]);
for (point pt7[i,j,k]: reg_br) {
  ux = xv[p2[i][j][k]] - xv[p1[i][j][k]] ;
  uy = yv[p2[i][j][k]] - yv[p1[i][j][k]] ;
  uz = zv[p2[i][j][k]] - zv[p1[i][j][k]] ;...
}

Figure 5.14 : We highlight the array transformation of X10 arrays into Java arrays to boost runtime performance. In this hexahedral cells volume calculation code fragment, our compiler could not transform X10 arrays x, y, z, xv, yv, zv into Java arrays because the Java language doesn't have an equivalent array view operation.
//code fragment shows array linearization ...
region reg_mex = [0:MESH_EXT-1];
region reg_mex_3D = [reg_mex,reg_mex,reg_mex];
double[.] x,y,z = new double[reg_mex_3D];
for (point pt3[pz] : reg_mex) {
    for (point pt2[py] : reg_mex) {
        for (point pt1[px] : reg_mex) {
            x[pz,py,px] = tx; //use productive multi-D
            y[pz,py,px] = ty; //access with array views
            z[pz,py,px] = tz;
            tx += ds;
        }
        ty += ds;
    }
    tz += ds;
}
region reg_br = [0:MESH_EXT-2];
region reg_br_3D = [reg_br, reg_br, reg_br];
int[] p1,p2 = new int[LinearViewAuto(reg_br_3D)];
region reg_linear = [0:MESH_EXT*MESH_EXT*MESH_EXT-1];
double[.] xv = x.view([0,0], [LinearViewHand(reg_mex_3D)];
double[.] yv = y.view([0,0], [LinearViewHand(reg_mex_3D)];
double[.] zv = z.view([0,0], [LinearViewHand(reg_mex_3D)];
for (point pt7[i,j,k] : reg_br) { //sub M for MESH_EXT
    ux=xv[p2[k+(M-1)(j+(M-1)*i)]-xv[p1[k+(M-1)(j+(M-1)*i)];
    uy=yv[p2[k+(M-1)(j+(M-1)*i)]-yv[p1[k+(M-1)(j+(M-1)*i)];
    uz=zv[p2[k+(M-1)(j+(M-1)*i)]-zv[p1[k+(M-1)(j+(M-1)*i)];
}

Figure 5.15: We illustrate the array transformation of X10 arrays into Java arrays and subsequent Java array linearization. Note that LinearViewAuto is a method our compiler automatically inserts to linearize a multi-dimensional X10 array and LinearViewHand is a method the programmer inserts to linearize an X10 region.
Automatic linearization on the hexahedral code fragment further decreased the execution time by 12%. However, there are opportunities to realize faster execution times by optimizing away the X10 `array.view()` methods, enabling the array transformation strategy to convert and linearize the remaining X10 general arrays. We observe that the array views in this code fragment are themselves X10 linearized representations of the view target X10 array. If there are no other conditions preventing our compiler from performing array conversion and linearization on these X10 general arrays, linearizing these X10 general array at the declaration site introduces redundant linearization operations, namely these X10 `array.view()` in Figure 5.15. As a result, we can optimize away the array views by replacing them with an assignment to the whole array. Figure 5.16 provides the final source output for the hexahedral cells code fragment. Performing this optimization enables us to achieve an additional factor of 7 speedup relative to the previous best execution time and an order of magnitude improvement over the code version with only X10 general arrays.

5.5 Interprocedural Linearized Array Bounds Analysis

Our array bounds analysis algorithm as described in Section 5.1 and Section 5.2 makes heavy use of X10 points and regions to discover when bounds checks are superfluous. In general, the programmer iterates through the elements in an array by implementing an X10 for loop whose header contains both a point declaration \( p1 \) and the region \( r1 \) containing the set of points defining \( p1 \). As a result, when encountering an array access with subscript \( p1 \), if our array bounds analysis can establish a subset relationship between the array’s region and region \( r1 \), our analysis can signal the VM that a bounds check for this array access is superfluous.

In scientific codes, application developers typically linearize multi-dimensional array representations to deliver improved runtime efficiency. One drawback to this scheme is that to support this approach, programmers must often introduce complex subscript expressions when accessing elements in the linearized array to ensure
//code fragment shows opt with array linearization

region reg_mex = [0:MESH_EXT-1];
region reg_mex_3D = [reg_mex, reg_mex, reg_mex];
double[.] x, y, z = new double[LinearViewAuto(reg_mex_3D)];...
for (point pt3[pz] : reg_mex) {...
  for (point pt2[py] : reg_mex) {...
    for (point pt1[px] : reg_mex) {
      x[pz,py,px] = tx; //use productive multi-D
      y[pz,py,px] = ty; //access with array views
      z[pz,py,px] = tz;
      tx += 'ds;
    }
    ty += 'ds;
  }
  tz += 'ds;
}

//use productive multi-D
//access with array views

region reg_br = [0:MESH_EXT-2];
region reg_br_3D = [reg_br, reg_br, reg_br];
int[] p1,p2 = new int[LinearViewAuto(reg_br_3D)];...
region reg_linear=[0:MESH_EXT*MESH_EXT*MESH_EXT-1];
double[] xv = x;
double[] yv = y;
double[] zv = z;
for (point pt7[i,j,k]: reg_br) { //sub M for MESH_EXT
  ux=xv[p2[k+(M-1)(j+(M-1)*i)]-xv[p1[k+(M-1)(j+(M-1)*i)];
  uy=yv[p2[k+(M-1)(j+(M-1)*i)]-yv[p1[k+(M-1)(j+(M-1)*i)];
  uz=zv[p2[k+(M-1)(j+(M-1)*i)]-zv[p1[k+(M-1)(j+(M-1)*i)];
}

Figure 5.16 : We show the final version for the Hexahedral cells code which demonstrates the compiler's ability to translate X10 arrays into Java arrays in the presence of array views.
correctness. As a result, our array bounds analysis loses the ability to make the straightforward comparison between an array's region and its point subscript comprising the loop iteration space to discover unnecessary bounds checks for linearized arrays. Ideally, from our compiler's perspective, we should convert the linearized arrays back into the multi-dimensional representation, enabling the bounds analysis to treat linearized and multi-dimensional array accesses in the same way. However, automatically converting linearized arrays to a multi-dimensional representation is certainly not trivial and in some cases may not be possible.

Figure 5.17 illustrates an MG code fragment where the application developer linearizes a 3-dimensional array to boost runtime performance. This example shows why our current array bounds analysis cannot rely on the compiler automatically converting linearized arrays to X10 multi-dimensional arrays because the range for each dimension in this case cannot be established. As a result, our bounds analysis must be extended if we want to analyze linearized array accesses to discover useless bound checks. Figure 5.18 highlights another extension to the array bounds analysis we previously described in Section 5.1 and Section 5.2. Studying the MG code fragment reveals that all the accesses to array \( r \) inside method \( psinv \) are redundant. Our array bounds analysis adds the following requirements to prove that \( r \)'s bounds checks are redundant:

- The array region summary for \( psinv \)'s formal parameter \( r \) is a subset of the region summary for \( zero3 \)'s formal parameter \( z \). The region summary for a given array and procedure defines the valid array index space inside the procedure for which a bounds check is useless. The region summary contains only an index set that must execute when the programmer invokes this method. We do not include array accesses occurring inside conditional statements in the region summary.

- The region representing the actual argument of \( psinv \)'s formal parameter \( r \) is a subset of the region representing the actual argument for \( zero3 \)'s formal
// MG code fragment is used to highlight
// challenge of converting linearized
// arrays to a multi-dimensional representation
...
// create linearized array
int nm = 2+(1<<1m);
int nv = (2+(1<<ndim1))*(2+(1<<ndim2))*(2+(1<<ndim3));
int nr = (8*(nv+nm*nm+5*nm+7*lm))/7;
region reg_nr = [0:nr-1]; // non-trivially 3-D reconstruction
double[.] u = new double[reg_nr];...
zero3(u, 0, n1, n2, n3);
...
void zero3(double[.] z, int off, int n1, int n2, int n3) {
for (point p1[i3,i2,i1]: [0:n3-1,0:n2-1,0:n1-1])
z[off+i1+n1*(i2+n2*i3)] = 0.0;
}

Figure 5.17: Array u is a 3-dimensional array that the programmer has linearized to improve runtime performance. Converting the linearized array into an X10 3-dimensional array would remove the complex array subscript expression inside the loop in zero3's method body and enable bounds analysis to attempt to discover a superfluous bounds check. However, this example shows it may not be possible to always perform the conversion.

- The program must call zero3 before calling psinv.
- Since our analysis modifies psinv's actual method body, the previous requirements must hold on all calls to psinv.

These requirements enable our interprocedural region analysis to delinearize array accesses into region summaries and to propagate the region summary information to discover redundant bounds checks.
//MG code fragment highlights opportunity to eliminate
//bound checks with procedure array bound summaries
int nm = 2+(1<<1m);
int nv = (2+(1<<ndim1))*(2+(1<<ndim2))*(2+(1<<ndim3));
int nr = (8*(nv+nm*nm+5*nm+7*lm))/7;
region reg_nr = [0:nr-1];
double[] u = new double[reg_nr]; //create linearized array
zero3(u, 0, n1, n2, n3);
psinv(u, 0, n1, n2, n3);

... void zero3(double[] z, int off, int n1, int n2, int n3) {
  for (point p1[i3, i2, i1]: [0:n3-1, 0:n2-1, 0:n1-1])
    z[off+i1+n1*(i2+n2*i3)] = 0.0;
}

void psinv(double[] r, int off, int n1, int n2, n3) {
  for (point p40[i3, i2]: [1:n3-2, 1:n2-2]) {
    for (point p41[i1]: [0:n1-1]) {
      r1[p41] = r[roff+i1+n1*(i2-1+n2*i3)]
        + r[roff+i1+n1*(i2+1+n2*i3)]
        + r[roff+i1+n1*(i2+n2*(i3-1))]
        + r[roff+i1+n1*(i2+n2*(i3+1))];
      r2[p41] = r[roff+i1+n1*(i2-1+n2*(i3-1))]
        + r[roff+i1+n1*(i2+1+n2*(i3-1))]
        + r[roff+i1+n1*(i2-1+n2*(i3+1))]
        + r[roff+i1+n1*(i2+1+n2*(i3+1))];
    }
  }
}

Figure 5.18: This MG code fragment shows an opportunity to remove all array r
bounds checks inside the psinv method because those checks are all redundant since
the programmer must invoke method zero3 prior to method psinv.
Chapter 6

High Productivity Language Iteration

Novice programmers are taught that they should separate the specification of their algorithms from the data structures used to implement them, in order to create code that is more robust in the face of changes to either. Unfortunately, scientific computing has a history of mixing the specification of algorithms with their implementations, due in part to the need for performance and in part to the languages that are traditionally used for such applications.

Scientific programmers targeting uni-processors take great care to iterate over their data structures in a manner that will maximize performance by generating loops that will walk through memory in a beneficial order, take advantage of the cache, enable vectorization, and so forth. Since C and Fortran are the most prevalent languages used in this domain, iterations are typically expressed using carefully-architected scalar loop nests. As an example, programmers who wish to iterate over their array elements in a tiled manner will typically need to intersperse all the details associated with tiling (extra loops, bounds calculations, etc.) in with their computation, even though the algorithm probably does not care about these implementation details.

As a scientific code evolves or is ported to new machines, each of these loop nests may need to be rewritten to match the new parameters. One typical scenario involves porting a multidimensional array code from C to Fortran and changing all of its loops to deal with the conversion between arrays allocated in row-major and column-major order. Other porting efforts may require the loops to change due to new cache parameters or vectorization opportunities. In the worst case, every loop nest that contributes to the code's performance may need to be considered and rewritten.
during this porting process.

When coding for a parallel environment, the problem tends to be even more difficult due to the fact that data structures are potentially distributed among multiple processors. As a result, loops tend to be cluttered with additional details, such as the specification of each processor's local bounds, in addition to the traditional uniprocessor concerns described above. By embedding such details within every loop that accesses a distributed data structure, a huge effort is typically required to change the distribution or implementation of the data structure, resulting in code that is brittle and difficult to experiment with. In short, our community has failed to separate algorithms from data structures in high performance computing as intended.

This chapter describes our attempts to address this fragility within scientific codes by introducing an iterator abstraction, developed by the Chapel team, within the Chapel parallel programming language [22]. An iterator is a software unit that encapsulates general computation, defining the traversal of a possibly multidimensional iteration space. Iterators are used to control loops simply by invoking them within the loop header. Moreover, multiple iterators may be invoked within a single loop using either cross-product or zipped semantics [34, 59]. Just as functions allow repeated subcomputations to be factored out of a program and replaced with function calls, iterators support a similar ability to factor common looping structures away from the computations contained within the bodies of those loops. Changes to an iterator's definition will be reflected in all uses of the iterator, and loops can alter their iteration method either by modifying the arguments passed to the iterator or by invoking a different iterator. The result is that users (and in some cases the compiler) can switch between different iteration methods without cluttering the expression of the algorithm or requiring changes to every loop nest.

The novel contributions are as follows:

- We provide in Section 6.2 examples of using iterators that suggest their productivity benefits within larger scientific codes.
• We describe in Section 6.4.2 a nested function-based Chapel iterator implemen-
tation, which extends the capability of the sequence-based approach and
addresses its limitations.

• We describe in Section 6.5 different implementation strategies for zippered iter-
ation to support producer-consumer iteration patterns not commonly supported
in most modern languages.

6.1 Overview of Chapel

Chapel is an object-oriented language that, along with Fortress [3] and X10 [39], is
being developed as part of DARPA's High-Productivity Computing Systems (HPCS)
program, challenging supercomputer vendors to increase productivity in high perfor-
mance computing. The design of Chapel is guided by four key areas of program-
ming language technology: multithreading, locality-awareness, object-orientation,
and generic programming. The object-oriented programming area, which includes
Chapel's iterators, helps in managing complexity by separating common function
from specific implementation to facilitate reuse. The common function or specifica-
tion in scientific loops is how to specify the traversal a multi-dimensional the iteration
space for the data structures referenced inside loops in a way that maximizes reuse
and minimizes clutter within the algorithm. This specification can be reused if it is
factored away from the implementation of the algorithm. The benefit comes from
saving programmers from having to rewrite the specification alongside their com-
putations each time the code traverses those data structures. The separation also
allows the programmer to focus on the iteration and computation separately. Chapel
iterators provide a framework to achieve this goal effectively.
6.2 Chapel Iterators

Chapel iterators are semantically similar to iterators in CLU [68]. Chapel implements iterators using a function-like syntax, although the semantic behavior of an iterator differs from that of a function in some important ways. Unlike functions, instead of returning a value, Chapel iterators typically return a sequence of values. The yield statement, legal only within iterator bodies, returns a value and temporarily suspends the execution of the code within the iterator. As an example, the following Chapel code defines a trivial iterator that yields the first \( n \) values from the Fibonacci sequence:

```chapel
iterator fibonacci(n):integer {
    var i1 = 0, i2 = 1;
    var i = 0;
    while i <= n {
        yield i1;
        var i3 = i1 + i2;
        i1 = i2;
        i2 = i3;
        i += 1;
    }
    return;
}
```

Chapel invokes iterators using a syntax similar to function calls. Chapel iterator calls commonly appear in loop headers to model the idea of executing the loop body's computation once for each element in a data structure's iteration space. In Chapel, the ordering of a loop's iterations is specified by the iterator call located in the loop header. As a result, all the developer has to do to change the iteration space ordering is to modify the iterator invocation. As an example, the following loop invokes our Fibonacci iterator to generate 10 values, printing them out as they are yielded:

```chapel
for val in fibonacci(10) do
```
write(val);

Conceptually, control of execution switches between the iterator and the loop body. The actual Chapel iterator implementation, as we discuss in Section 6.4.1, may store all the yielded values in a list-like structure and subsequently execute the loop body once for each element in the list. Semantically, the loop body executes each time a `yield` statement inside the iterator executes. Upon completion, the loop body transfers control back to the statement following the `yield`. However, control of execution does not switch to the loop body when a `return` statement inside the iterator executes. Figure 6.1 provides a more detailed view of how iterators in Chapel may be utilized, using an example based on the NAS parallel benchmark FT [9], where we use the simplicity of iterators to experiment with tiling. This example shows three iterators that might be used to traverse a 2D index space, and shows that the `evolve` client code can switch between them simply by invoking a different iterator.

Chapel's iterators may be invoked using either sequential `for` loops, as shown above, or parallel `forall` loops. The iterator's body may also be written to utilize parallelism, potentially yielding values using multiple threads of execution. In such cases, the `ordered` keyword may be used when invoking the iterator in order to respect any sequential constraints within the iterator's body. Figure 6.2 illustrates this utilizing two Chapel iterators for the Smith-Waterman algorithm, a well-known dynamic programming algorithm in scientific computing that performs DNA sequence comparisons. Figure 6.3 shows, using an example similar to one found in the Chapel language specification [34], an parallel iterator traversing through an abstract syntax tree (AST) until it reaches all the leaf nodes. For more details, the reader is referred to the Chapel language specification [34]. This chapter focuses primarily on the implementation of sequential iterators, which represent a crucial building block for efficiently supporting parallel iterators and iteration.
iterator rmo(d1,d2): 2*integer do //row major order
  for i in 1..d1 do
    for j in 1..d2 do
      yield (i,j);

iterator cmo(d1,d2): 2*integer do //column major order
  for j in 1..d2 do
    for i in 1..d1 do
      yield (i,j);

iterator tiledcmo(d1,d2): 2*integer { //tiled col major order
  var (b1,b2) = computeTileSizes();
  for j in 1..d2 by b2 do
    for i in 1..d1 by b1 do
      for jj in j..min(d2,j+(b2-1)) do
        for ii in i..min(d1,i+(b1-1)) do
          yield (ii,jj);
}

function evolve(d1,d2) do
  for (i,j) in {rmo|cmo|tiledcmo}(d1,d2) { u0(i,j) = u0(i,j)*twiddle(i,j); u1(i,j) = u0(i,j); }

Figure 6.1: A basic iterator example showing how Chapel iterators separate the specification of an iteration from the actual computation.
Figure 6.2: A parallel excerpt from the Smith-Waterman algorithm written in Chapel using iterators. The `ordered` keyword is used to respect the sequential constraints within the loop body.
class Tree {
  var isLeaf : boolean;
  var left : Tree;
  var right : Tree;
}

class Leaf implements Tree {
  var value : integer;
}

iterator Tree.walk(): {
  if(isLeaf)
    yield(this);
  else
    cobegin {
      left.walk();
      right.walk();
    }
}

Tree t;
...

//print value of all leaves in tree
for leaf in t.walk()
  print leaf.value;

Figure 6.3: An iterator example showing how to use Chapel iterators to traverse an abstract syntax tree (AST).
6.3 Invoking Multiple Iterators

Chapel supports two types of simultaneous iteration by adding additional iterator invocations in the loop header. Developers can express cross-product iteration in Chapel by using the following notation:

```chapel
for (i,j) in [iter1(),iter2()] do ...
```

which is equivalent to the nested `for` loop:

```chapel
for i in iter1() do
  for j in iter2() do
    ...
```

Zipper-product iteration is the second type of simultaneous iteration supported by Chapel, and is specified using the following notation:

```chapel
for (i,j) in (iter1(),iter2()) do ...
```

which, assuming that both iterators yield \( k \) values, is equivalent to the following pseudocode:

```chapel
for p in 1..k {
  var i = iter1().getNextValue();
  var j = iter2().getNextValue();
  ...
}
```

In this case, the body of the loop will execute each time both iterators yield a value. However, recall that the semantics of the Chapel iterators, differing from normal functions, require that once program execution reaches the last statement in the loop body, control resumes inside the iterator body on the statement immediately following the `yield` statement for each iterator. Zippered iteration would be implemented
naturally using coroutines [48], which allow for execution to begin anywhere inside of a function, unlike functions in most current languages. However, without coroutines, zipper-product iteration may still be implemented using techniques we describe in Section 6.5.

6.4 Implementation Techniques

Chapel has two iterator implementation techniques, an iterator approach using sequences and an alternate approach using nested functions. The original approach was the sequence based implementation. Our contribution to Chapel iterators is the nested function based implementation. The motivation for our nested function based approach was to overcome the limitations of the sequence based approach. In the next sections we first describe the original Chapel iterator implementation and subsequently introduce our nested-function based solution to address the limitations of the original technique.

6.4.1 Sequence Implementation

The Chapel compiler’s original implementation approach for iterators uses sequences to store the iteration space of the data structures traversed by the loop. Subsequently, the loop body is executed once for each element in the sequence.

Sequences in Chapel are homogeneous lists which support iteration via a built-in iterator. Chapel supports declarations of sequence variables and iterations over them using the following syntax:

```chapel
var aseq: seq(integer) = (/ 1, 2, 4 /);
for myInt in aseq do ... 
```

where `integer` in this example can be replaced by any type.

In the sequence-based implementation, Chapel first evaluates the iterator call and builds up the sequence of yielded values before executing the loop body. Each time
// Illustration of compiler transformation

function tiledcmo(d1,d2): seq(2*integer) {
    var resultSeq: seq(2*integer);
    var (b1,b2) = computeTileSizes();
    for j in 1..d2 by b2 do
        for i in 1..d1 by b1 do
            for jj in j..min(d2,j+(b2-1)) do
                for ii in i..min(d1,i+(b1-1)) do
                    resultSeq.append(ii,jj);
    return resultSeq;
}

function evolve(d1,d2) {
    var resultSeq = tiledcmo(d1,d2);
    for (i,j) in resultSeq {
        u0(i,j) = u0(i,j)*twiddle(i,j);
        u1(i,j) = u0(i,j);
    }
}

Figure 6.4: An implementation of tiled iteration using the sequence-based approach.

the iterator yields a value, instead of executing the loop body, Chapel appends the value to a sequence. When execution reaches either the end of the iterator or a return statement, the iterator returns the constructed sequence of yielded values. Once the iterator returns its sequence of values, Chapel begins executing the loop body once for each element in the sequence returned from the iterator. Figure 6.4 illustrates the compiler rewrite that would take place using the sequence-based iteration approach for the tiled iterator of Figure 6.1.

The advantage to using the original approach is its simplicity. The Chapel compiler can use the language's built-in support for sequences to capture the iteration
space and to control how many times the loop body executes. Another advantage is that the iterator function only needs to be called once. As a result, this approach saves the cost of transferring control back and forth between the iterator and the loop body.

The chief disadvantage to this approach is that it is not general. It can only be applied when the compiler can ensure that no side effects exist between the iterator and loop body. Chapel must impose the side effect restriction because the sequence gathers the iteration space before loop body execution begins. If there was a side effect inside the loop body, such as changing the bounds of the iteration space, incorrect code could be produced. A second disadvantage to this approach is the space overhead required to store the sequence. The next section details our nested function-based Chapel iterator implementation approach, which addresses these limitations.

6.4.2 Nested Function Implementation

Our novel contribution to the Chapel compiler is the alternative iterator implementation strategy using nested functions [59]. Currently, this approach works well on a \textit{for} loop containing one iterator call in its loop header. We provide insight on extending this approach to handle zipper-product iteration in \textit{Section 6.5}. Implementing zipper-product iteration is a subject for future work.

There are two steps to implementing Chapel iterators with nested functions. The first step involves creating a nested function within the iterator’s scope that implements the \textit{for} loop’s body and takes the loop indices as its arguments. The second step creates a copy of the iterator, converting it to a function and replacing each \textit{yield} statement in the body with a call to the nested function created during the first step. The transformation passes the value of each yield statement as arguments to the nested function. Once the transformation completes this process, it replaces the original \textit{for} loop with the cloned iterator call, previously located in its loop header. Figure 6.5 demonstrates how the Chapel compiler implements iterators using nested
// Illustration of compiler transform

function evolve(d1, d2) {
    function tiledcmo(d1, d2) {
        function loopbody(i, j) {
            u0(i, j) = u0(i, j) * twiddle(i, j);
            u1(i, j) = u0(i, j);
        }
        var (b1, b2) = computeTileSizes();
        for j in 1..d2 by b2 do
            for i in 1..d1 by b1 do
                for jj in j..min(d2, j+(b2-1)) do
                    for ii in i..min(d1, i+(b1-1)) do
                        loopbody(ii, jj);
        }
    }
    tiledcmo(d1, d2);
}

Figure 6.5: An implementation of tiled iteration using the nested function-based approach.
functions for the tiling example.

Since the body of the nested function inside the iterator is small, it is often beneficial to inline it. Chapel inlines the nested function calls appearing inside the iterator to eliminate the costs of invoking the nested function every time the iterator yields a value. This optimization is not possible with the sequence-based approach since the iterator must yield all its values before preceding to execute the loop body.

Another advantage of using the nested function approach for iterators is generality: side effects between the iterator and the for loop’s body do not have to be identified in fear of producing incorrect code. As a result, this approach is more broadly applicable than using the sequence-based approach. The execution behavior of this approach is closer to that of CLU [69] and Sather [76] iterators. In addition, an advantage over the sequence-based approach is Chapel does not need to use storage for the iteration space. Consequently, the nested-function implementation is more practical in environments where large iteration spaces may be in danger of overflowing memory.

6.5 Zippered Iteration

Zipper-product iteration is the process of traversing through multiple iterators simultaneously where each iterator must yield a value once before execution of the loop body can begin. Figure 6.6 shows an example of zippered iteration in Chapel. This section describes possible zipper-product implementation approaches that we are exploring as we go forward. Chapel’s semantics define that zippered iteration is performed by requiring the iterators involved in the loop to each yield values before the loop body is executed. Recall that semantically, when an iterator yields a value, execution suspends from inside the iterator until the loop body has completed once. When execution resumes inside the iterator, Chapel will execute the statement immediately following the yield statement.

In modern languages, the only point of entry for functions is at the top. Coroutines are functions that can have multiple entry points and properly simulate the producer/-
iterator fibonacci(n):integer {
  var i1 = 0, i2 = 1;
  var i = 0;
  while i <= n {
    yield i1;
    var i3 = i1 + i2;
    i1 = i2;
    i2 = i3;
    i += 1;
  }
}

iterator squares(n):integer {
  var i = 0;
  while i <= n {
    yield i * i;
    i += 1;
  }
}

for i, j in fibonacci(12), squares(12) do
  writeln(i, " , ", j);

Figure 6.6: An example of zippered iteration in Chapel.
consumer relationship that simultaneous iteration between two iterators introduces. However, because most modern languages do not support coroutines, programmers must utilize other methods to properly simulate the producer/consumer relationship. Here we consider two techniques, one that uses state variables and one that uses multiple threads via synchronization variables.

Figure 6.7 shows one technique for implementing zipper-product iteration. The example implements the zippered iteration using state variables. Both iterators use Chapel's `select` statement with `goto` statements to enable simulation of a coroutine, similar to checkpointing in the porch compiler [92]. The state is preserved via the class that is passed into the function. The semantic execution behavior of the iterators is preserved by ensuring that the statement immediately following the `yield` is executed when the iterators are invoked on subsequent calls. Once the last yield is executed, the iterator will not be called again. The advantage of using this approach is that it eliminates the synchronization costs that are associated with our second approach. Also, by having the compiler simulate the coroutine, dead variables do not need to have their state saved. For example, an optimization could be performed to eliminate \( i3 \) from the state class for the Fibonacci iterator. The disadvantage of this approach is the overhead associated with entering and exiting the routine. This could be especially significant in recursive iterators where the stack would result in a large saved state class.

Our second implementation approach for zippered iteration uses multiple threads and synchronization (sync) variables. A `sync variable`[34] transitions to an undefined state when read. When a sync variable is undefined and a computation tries to read from it, the computation will stall until the sync variable is defined. As a result, sync variables allow us to model the producer/consumer relationship of coroutines that is needed to support zippered iteration. Note that the multi-threaded solution requires analysis which determines whether the iterators are parallel-safe or semantics which imply that iterators in a zippered context are executed in parallel.
In Figure 6.8, the sync variables are initially undefined. Each sync variable can transition to the defined state inside an iterator. Chapel utilizes the \textit{cobegin} statement to indicate that both iterators should be executed in parallel. The \textit{while} loop inside the \textit{cobegin} statement will stall until each iterator defines its sync variables. A sync variable is created for each iterator and a sync variable assignment replaces each \textit{yield} statement inside the iterator. The chief disadvantage to using this approach lies in the synchronization costs associated with the sync variables. Both approaches enable the support of zippered iteration in Chapel.
// Illustration of compiler transform

class ss_fib_state {var i1, i2, i3, i: integer; var jump = 1;}

function ss_fibonacci(n, ss): integer {
    select ss.jump {when 1 do goto lab1; when 2 do goto lab2;}

    label lab1 ss.i1 = 0;
    ss.i2 = 1; ss.i = 0;
    while ss.i <= n {
        ss.jump = 2;
        return ss.i1;

        label lab2 ss.i3 = ss.i1 + ss.i2;
        ss.i1 = ss.i2; ss.i2 = ss.i3;
        ss.i += 1; }
    ss.jump = 0;
    return 0;
}

class ss_sq_state { var i: integer; var jump = 1; }

function ss_squares(n, ss): integer {
    select ss.jump {when 1 do goto lab1; when 2 do goto lab2;}

    label lab1 ss.i = 0;
    while ss.i <= n {
        ss.jump = 2;
        return ss.i * ss.i;

        label lab2 ss.i += 1; }
    ss.jump = 0;
    return 0;
}

var ss1 = ss_fib_state(); var ss2 = ss_sq_state();
while ss1.jump and ss2.jump do {
    var i = ss_fibonacci(12, ss1); var j = ss_squares(12, ss2);
    writeln(i, ",", j);
}

Figure 6.7: An implementation of zippered iteration using state variables.
// Illustration of compiler transform

class mt_fib_state{sync flag:bool; sync result:integer;}
function mt_fibonacci(n, mt) {
  var i1 = 0, i2 = 1, i = 0;
  while i <= n {
    mt.flag = false;
    mt.result = i1;
    var i3 = i1 + i2;
    i1 = i2;
    i2 = i3;
    i += 1; }
  mt.flag = true;
}

class mt_sq_state{sync flag:bool; sync result:integer;}
function mt_squares(n, mt) {
  var i = 0;
  while i <= n {
    mt.flag = false;
    mt.result = i * i;
    i += 1; }
  mt.flag = true;
}

var mt1 = mt_fib_state(); var mt2 = mt_sq_state();
cobegin {
  mt_fibonacci(12);
  mt_squares(12);
  while not mt1.flag and not mt2.flag do
    writeln(mt1.result, ", " , mt2.result);
  }

Figure 6.8: A multi-threaded implementation of zippered iteration using sync variables.
Chapter 7

Performance Results

We ran the first set of experiments on a 1.25 GHz PowerPC G4 with 1.5 GB of memory using the Sun Java Hotspot VM (build 1.5.0_07-87) for Java 5. We measured performance results on the Java Grande benchmarks written in X10. These results are obtained using the class A versions of the benchmark. We report results for 3 different versions of the benchmark suite. Version 1 is an unoptimized direct translation of the original Java version obtained from the Java Grande Forum web site [54] renamed with the .xl0 extension, with all Java arrays converted into X10 arrays and integer subscripts replaced by points. Version 2 uses the same input X10 program as in Version 1 but turns on point inlining and uses programmer inserted dependent types to improve performance. Version 3, containing only Java arrays, can be considered as the baseline. We refer to Version 3 as X10 Light. These results include runtime array bounds checks, null pointer checks and other checks associated with a Java runtime environment.

Table 7.1 shows the impact unoptimized high-level X10 array computation has on performance by comparing Versions 1 and 3. The unoptimized X10 version runs up to almost 84 times slower. Table 7.2 shows the impact of the inlining points and generating efficient array accesses with dependent types by comparing the performance of Versions 1 and 2. While performance improvements in the range of 1.6× to 5.4× were observed for 7 of 8 benchmarks in Table 7.2, there still remain opportunities to employ automatic compiler interprocedural rank inference to replace high-level X10 arrays with more efficient representations; thereby leading to even better performance. Note: we observed no improvement in the series benchmark because its
performance is dominated by scalar (rather than array) operations.

<table>
<thead>
<tr>
<th>Benchmarks</th>
<th>Sequential Runtime Performance in seconds</th>
<th>Performance Slowdown</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Unopt. X10 (Version 1)</td>
<td>X10 Light (Version 3)</td>
</tr>
<tr>
<td>sparsematmult</td>
<td>57.97</td>
<td>9.75</td>
</tr>
<tr>
<td>crypt</td>
<td>8.14</td>
<td>4.60</td>
</tr>
<tr>
<td>lufact</td>
<td>52.87</td>
<td>1.38</td>
</tr>
<tr>
<td>sor</td>
<td>508.49</td>
<td>6.06</td>
</tr>
<tr>
<td>series</td>
<td>19.01</td>
<td>19.01</td>
</tr>
<tr>
<td>moldyn</td>
<td>2.39</td>
<td>0.57</td>
</tr>
<tr>
<td>montecarlo</td>
<td>7.59</td>
<td>3.00</td>
</tr>
<tr>
<td>raytracer</td>
<td>2.27</td>
<td>1.28</td>
</tr>
</tbody>
</table>

Table 7.1: Raw runtime performance showing slowdown that results from not optimizing points and high-level arrays in sequential X10 version of Java Grande benchmarks.

The second set of performance results reported in this section were obtained using the following system settings:

- The target system is either an IBM 64-way 2.3 GHz Power5+ SMP with 512 GB main memory or an IBM 16-way 4.7 GHz Power6 SMP with 186 GB main memory. Assume the former unless otherwise specified. The 16-way machine was used for the bounds check elimination results.

- The Java runtime environment used is the IBM J9 virtual machine (build 2.4, J2RE 1.6.0) which includes the IBM TestaRossa (TR) Just-in-Time (JIT) compiler [93]. The following internal TR JIT options were used for all X10 runs:
  - Options to enable classes to be preloaded, and each method to be JIT-compiled at a high ("very hot") optimization level on its first execution.
  - An option to ignore strict conformance with IEEE floating point.
<table>
<thead>
<tr>
<th>Benchmarks</th>
<th>Sequential Runtime Performance in seconds</th>
<th>Speedup Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>spsematmult</td>
<td>57.97</td>
<td>13.83</td>
</tr>
<tr>
<td>crypt</td>
<td>8.14</td>
<td>4.79</td>
</tr>
<tr>
<td>lufact</td>
<td>52.87</td>
<td>18.86</td>
</tr>
<tr>
<td>sor</td>
<td>508.49</td>
<td>93.41</td>
</tr>
<tr>
<td>series</td>
<td>19.01</td>
<td>18.95</td>
</tr>
<tr>
<td>moldyn</td>
<td>2.39</td>
<td>1.19</td>
</tr>
<tr>
<td>montecarlo</td>
<td>7.59</td>
<td>3.49</td>
</tr>
<tr>
<td>raytracer</td>
<td>2.27</td>
<td>1.43</td>
</tr>
</tbody>
</table>

Table 7.2: Raw runtime performance from optimizing points and using dependent types to optimize high-level arrays in sequential X10 version of Java Grande benchmarks.

- A special skip checks option was used for some of the results to measure the opportunities for optimization. This option directs the JIT compiler to disable all runtime checks (array bounds, null pointer, divide by zero).

- Version 1.5 of the X10 compiler and runtime [101] were used for all executions. This version supports implicit syntax [100] for place-remote accesses. In addition, all runs were performed with the number of places set to 1, so all runtime “bad place” checks [25] were disabled.

- The default heap size was 2GB.

- For all runs, the main program was extended with a three-iteration loop within the same Java process, and the best of the three times was reported in each case. This configuration was deliberately chosen to reduce the impact of JIT compilation time, garbage collection and other sources of perturbation in the performance comparisons.

The benchmarks studied in the second set of experiments are X10 ports of bench-
Figure 7.1: Comparison of the optimized sequential X10 benchmarks relative to the X10 light baseline

marks from the Java Grande [54] suite. We compare three versions of each benchmark:

1. The *light* versions use X10 concurrency constructs such as `async` and `finish`, while directly using low-level Java arrays as in [11]. While this version does not support the productivity benefits of higher-level X10 arrays, it serves as a performance target for the optimizations presented in this thesis.

2. The *unoptimized* versions use unoptimized X10 programs with high-level array constructs, obtained using the X10 reference implementation on SourceForge [101].

3. The *optimized* versions use the same input program as the *unoptimized* cases, with the optimizations introduced in this thesis applied.

Figure 7.1 shows the performance gap between “X10 Optimized” and “X10 Light”
(Version 1). The gap is at most 16% for MonteCarlo), but is under 1% in most other cases. In Figure 7.1, the reason why “X10 Optimized” delivers better performance than “X10 Light” for LUFact is because the address arithmetic present in the “X10 Light” version was naturally factored out in the “X10 Optimized” version due to the use of region iterators and points. We could modify the “X10 Light” version in this case to match the code that would be generated by the “X10 Optimized” version, but we instead chose to be faithful to the original Java Grande Forum benchmark structure when creating the “X10 Light” versions.

Next, we discuss the impact of our transformations on parallel performance. Table 7.3 shows the relative scalability of the Optimized and Unoptimized X10 versions. Since the biggest difference was observed for the \textit{sparsematmult} benchmark, we use Figure 7.2 and 7.3 to further study this behavior for \textit{sparsematmult}. Figure 7.2 illustrates that the optimized \textit{sparsematmult} benchmark scales better than the un-optimized version with an initial minimum heap size of 2 GB. Figure 7.3 shows that decreasing the initial minimum heap size to the default (4MB) value further increases the gap in scalability, suggesting that garbage collection is a major scalability factor in the Unoptimized case. These results are not surprising since the Unoptimized version allocates a large number of \textit{point} objects with short life times. Figures 7.4, 7.5, 7.6, and 7.7 show, using \textit{lufact} and \textit{sor}, that this behavior is not limited to \textit{sparsematmult}. The Optimized version mitigates this problem by inlining \textit{point} objects. We demonstrate with Figures 7.8 and 7.9 that Unoptimized X10 can scale as well as Optimized X10 in the absence of \textit{point-intensive} loops. Figures 7.10, 7.11, 7.12, 7.13 provide additional examples to highlight our optimization’s scalability impact using a minimum 2 GB heap size. Note that in all these results, the Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.

Table 7.4 shows the raw execution times for the unoptimized and optimized versions, and Figure 7.14 shows the relative speedup obtained due to optimization as we
Figure 7.2: Relative Scalability of Optimized and Unoptimized X10 versions of the sparsematmult benchmark with initial minimum heap size of 2 GB (and maximum heap size of 2GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.

Figure 7.3: Scalability of Optimized and Unoptimized X10 versions of the sparsematmult benchmark with initial minimum heap size of 4 MB (and maximum heap size of 2GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.
Figure 7.4: Relative Scalability of Optimized and Unoptimized X10 versions of the luFact benchmark with initial minimum heap size of 2 GB (and maximum heap size of 2GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.

Figure 7.5: Scalability of Optimized and Unoptimized X10 versions of the luFact benchmark with initial minimum heap size of 4 MB (and maximum heap size of 2GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.
Figure 7.6: Relative Scalability of Optimized and Unoptimized X10 versions of the sor benchmark with initial minimum heap size of 2 GB (and maximum heap size of 2GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.

Figure 7.7: Scalability of Optimized and Unoptimized X10 versions of the sor benchmark with initial minimum heap size of 4 MB (and maximum heap size of 2GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.
Figure 7.8: Relative Scalability of Optimized and Unoptimized X10 versions of the series benchmark with initial minimum heap size of 2 GB (and maximum heap size of 2GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.

Figure 7.9: Scalability of Optimized and Unoptimized X10 versions of the series benchmark with initial minimum heap size of 4 MB (and maximum heap size of 2GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.
Figure 7.10: Relative Scalability of Optimized and Unoptimized X10 versions of the crypt benchmark with initial minimum heap size of 2 GB (and maximum heap size of 2 GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.

Figure 7.11: Relative Scalability of Optimized and Unoptimized X10 versions of the montecarlo benchmark with initial minimum heap size of 2 GB (and maximum heap size of 2 GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.
Figure 7.12: Relative Scalability of Optimized and Unoptimized X10 versions of the moldyn benchmark with initial minimum heap size of 2 GB (and maximum heap size of 2GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.

Figure 7.13: Relative Scalability of Optimized and Unoptimized X10 versions of the raytracer benchmark with initial minimum heap size of 2 GB (and maximum heap size of 2GB). The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance.
Figure 7.14: Speedup of Optimized X10 version relative to Unoptimized X10 version.

scale from 1 to 64 CPUs. Figure 7.15 zooms in on Figure 7.14. As can be seen in Table 7.4 and Figure 7.14, the performance improvements due to optimization can be very significant, reaching as a high as a factor of 266×. The reason “series” behaves differently in Table 7.4 is due to the fact that its frequently-executed code regions are dominated by scalar computations. As a result, our array optimizations have very limited opportunities to impact performance for this benchmark. However, as shown in Table 7.4, the optimization opportunities are much larger for other scientific applications that are array intensive (as is the norm).

The benchmarks studied in the next set of experiments are X10 sequential ports of benchmarks from the Java Grande [54] suite. We compare two versions of each benchmark. The first version is the ported code. The second version, through static analysis, inserts noBoundsCheck calls around an array index when the bounds check is unnecessary.
Figure 7.15: Speedup of Optimized X10 version relative to Unoptimized X10 version (zoom in of Figure 7.14).

<table>
<thead>
<tr>
<th>Benchmarks (unopt/opt)</th>
<th>Runtime Performance speedup (relative to 1 CPUs)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
</tr>
<tr>
<td>sparsemm</td>
<td>1.0/1.0</td>
</tr>
<tr>
<td>crypt</td>
<td>1.0/1.0</td>
</tr>
<tr>
<td>lufact</td>
<td>1.0/1.0</td>
</tr>
<tr>
<td>sor</td>
<td>1.0/1.0</td>
</tr>
<tr>
<td>series</td>
<td>1.0/1.0</td>
</tr>
<tr>
<td>moldyn</td>
<td>1.0/1.0</td>
</tr>
<tr>
<td>montecarlo</td>
<td>1.0/1.0</td>
</tr>
<tr>
<td>raytracer</td>
<td>1.0/1.0</td>
</tr>
</tbody>
</table>

Table 7.3: Relative Scalability of Optimized and Unoptimized X10 versions with heap size of 2 GB. The Optimized speedup is relative to the 1-CPU optimized performance, and the Unoptimized speedup is relative to the 1-CPU unoptimized performance. The Optimized X10 version does not include the bounds check optimization.
Table 7.4: Raw runtime performance of Unoptimized and Optimized X10 versions as we scale from 1 to 64 CPUs. The Optimized X10 version does not include the bounds check optimization.

In Table 7.5, we report the dynamic counts for the Java Grande, hexahedral, and 2 NAS parallel (cg, mg) X10 benchmarks. We compare dynamic counts for potential general X10 array bounds checks against omitted general X10 array bounds checks using our static analysis techniques. We use the term "general X10 array" to refer to arrays the programmer declares using X10 regions. In several cases our static bounds analysis removes over 99% of potential bound checks.

In Figure 7.16, we report the execution times for the Java Grande, hexahedral, and 2 NAS parallel (cg, mg) X10 benchmarks both with and without the automatically generated noBoundsCheck annotations with runtime checks enabled. These annotations alert the IBM J9 VM when array bounds checking for an array access is unnecessary. Performing static array bounds analysis and subsequent automatic program transformation, we increase runtime performance by up to 22.3%. These results demonstrate that our static no bounds check analysis helps reduce the performance impact of programmers developing applications in type-safe languages. Table 7.6 shows that we may further improve runtime performance in some cases by eliminating other types of runtime checks such as null checks and cast checks.

Finally, in Table 7.7, we compare Fortran, Unoptimized X10, Optimized X10, and
Java execution times for the 2 NAS parallel (cg, mg) benchmarks. The Optimized X10 significantly reduces the slowdown factor that results from comparing Unoptimized X10 with Fortran. These results were obtained on the IBM 16-way SMP. Note: the 3.0 NAS Java mg version was run on a 2.16 GHz Intel Core 2 Duo with 2GB of memory due to a J9 JIT compilation problem with this code version. In the future, we will continue to extend our optimizations to further reduce the overhead of using high-level X10 array computations.

<table>
<thead>
<tr>
<th>Benchmarks</th>
<th>Dynamic Counts for X10 Array Bounds Checks (ABCs)</th>
<th>total X10 ABCs</th>
<th>total X10 ABCs eliminated</th>
<th>percent eliminated</th>
</tr>
</thead>
<tbody>
<tr>
<td>sparsemm</td>
<td></td>
<td>2,513,000,000</td>
<td>2,513,000,000</td>
<td>100.0%</td>
</tr>
<tr>
<td>crypt</td>
<td></td>
<td>1,000,000,312</td>
<td>100,000,130</td>
<td>10.0%</td>
</tr>
<tr>
<td>lufact</td>
<td></td>
<td>5,425,377,953</td>
<td>5,375,370,956</td>
<td>99.1%</td>
</tr>
<tr>
<td>sor</td>
<td></td>
<td>4,806,388,808</td>
<td>4,798,388,808</td>
<td>99.8%</td>
</tr>
<tr>
<td>series</td>
<td></td>
<td>4,000,020</td>
<td>4,000,002</td>
<td>99.9%</td>
</tr>
<tr>
<td>moldyn</td>
<td></td>
<td>5,955,209,518</td>
<td>4,023,782,878</td>
<td>67.6%</td>
</tr>
<tr>
<td>montecarlo</td>
<td></td>
<td>779,845,962</td>
<td>419,887,788</td>
<td>53.8%</td>
</tr>
<tr>
<td>raytracer</td>
<td></td>
<td>1,185,054,651</td>
<td>1,185,054,651</td>
<td>100.0%</td>
</tr>
<tr>
<td>hexahedral</td>
<td></td>
<td>35,864,066,928</td>
<td>32,066,331,077</td>
<td>89.4%</td>
</tr>
<tr>
<td>cg</td>
<td></td>
<td>3,044,164,220</td>
<td>1,532,754,859</td>
<td>50.4%</td>
</tr>
<tr>
<td>mg</td>
<td></td>
<td>6,614,097,502</td>
<td>383,155,390</td>
<td>5.8%</td>
</tr>
</tbody>
</table>

Table 7.5: Dynamic counts for the total number of X10 array bounds checks (ABC) in sequential JavaGrande, hexahedral benchmark and 2 NAS Parallel X10 benchmarks compared with the total number of eliminated checks we introduce using static compiler analysis and compiler annotations which signal the JVM to remove unnecessary bounds checks.
Figure 7.16: Comparison of the X10 light baseline to the optimized sequential X10 benchmarks with compiler inserted annotations used to signal the VM when to eliminate bounds checks.
Table 7.6: Raw sequential runtime performance of JavaGrande and 2 NAS Parallel X10 benchmarks with static compiler analysis to signal the JVM to eliminate unnecessary array bounds checks. These results were obtained on the IBM 16-way SMP because the J9 VM has the special option to eliminate individual bounds checks when directed by the compiler.

Table 7.7: Fortran, Unoptimized X10, Optimized X10, and Java raw sequential runtime performance comparison (in seconds) for 2 NAS Parallel benchmarks (cg, mg). These results were obtained on the IBM 16-way SMP machine.
Chapter 8

Conclusions and Future Work

Although runtime performance has suffered in the past when scientists used high productivity languages with high-level array accesses, our thesis is that these overheads can be mitigated by compiler optimizations, thereby enabling scientists to develop code with both high productivity and high performance. The optimizations introduced in this dissertation for high-level array accesses in X10 result in performance that rivals the performance of hand-tuned code with explicit rank-specific loops and lower-level array accesses, and is up to two orders of magnitude faster than unoptimized, high-level X10 programs.

In this thesis, we discussed the Point abstraction in high-productivity languages, and described compiler optimizations that reduce their performance overhead. We conducted experiments that validate the effectiveness of our Point inlining optimization combined with programmer specified dependent types and demonstrate that these optimizations can enable high-level X10 array accesses written with implicit ranks and Points to achieve performance comparable to that of low-level programs written with explicit ranks and integer indices. The experimental results showed performance improvements in the range of 1.6× to 5.4× for 7 of 8 Java Grande benchmark programs written in X10, as a result of these optimizations.

This thesis provides an algorithm to generate rank-specific efficient array computations from applications that use productive rank-independent general X10 arrays. The algorithm propagates X10 array rank information to generate the more efficient Java arrays with precise ranks. Our results demonstrate that we can generate efficient array representations and come within 84% of the baseline for each benchmark
and within 99% in most cases. This thesis introduces novel contributions to array bounds analysis by taking advantage of the X10 region language abstraction along with tracking array value ranges. We introduce an interprocedural static elimination bounds analysis algorithm with algebraic region inequality equations for points and integrals to establish region and sub region relationships; thereby aiding in the discovery of superfluous bounds checks when programmers utilized these variables in an array subscript or during region construction. We illustrate the benefits of array value ranges which are particularly useful in applications with sparse matrix computations. Experimental results show we experience runtime execution improvements of up to 22.3%. Our dynamic counts illustrate the elimination of over 99% of bounds checks in several cases with this optimization. In addition, we provide an optimization that provides a way to generate efficient array computations in the presence of array views resulting in a factor of 7 speedup. Another contribution is the analysis of how our optimizations impact scalability. The optimized version of the benchmarks scales much better than the unoptimized general X10 array version.

We also calibrated the performance of our optimizations for the two benchmarks for which equivalent Fortran programs were available, CG and MG. For CG, we improved the performance of the Unoptimized X10 by 3.15 × but there still remains a performance gap of 3.3× relative to the Fortran version. For MG, we improved the performance of the Unoptimized X10 by 3.42× but there still remains a performance gap of 13.66× relative to the Fortran version. In both cases, a large part of the performance gap can be attributed to the differences between the Java version and the Fortran version that have been studied in past work [75]. The remainder of the performance gap can be attributed to the cases where our optimization was not able to convert X10 arrays to Java arrays. These cases are challenging because the Java version includes hand-coded redundancy elimination that will require advanced compiler techniques such as redundancy elimination [21, 30] of array accesses through loop carried dependences to enable them to be performed automatically.
This thesis shows that Chapel iterators can effectively separate the specification of an algorithm from its implementation, thereby enabling programmers to easily switch between different specifications while also allowing them to focus on the algorithm's implementation. Using iterators to handle specifications such as iteration space ordering allows programmers to reuse specifications instead of having to write a specification for an algorithm each time the programmer implements an algorithm. We describe a novel strategy we have implemented in the Chapel compiler to support Chapel iterators which addresses the limitations of the original strategy. The first approach was sequence-based Chapel iterators and the second approach was to implement Chapel iterators with nested functions. The second strategy eliminates some of the imposed restrictions and spatial overhead of the first strategy.

In the future, we plan to extend our bounds analysis to discover sub region relationships between array views and to identify when regions share equivalent dimensions to reduce the cost for a multi-dimensional array access. We plan to extend array value range analysis with alias analysis to update array alias value ranges. We also plan to analyze and potentially eliminate other types of runtime checks such as cast checks and null checks. In addition, as part of the Habanero multicore software research project at Rice University [50], we plan to demonstrate on a wide range of applications that the techniques presented in this thesis can enable programmers to develop high productivity array computations without incurring the additional runtime costs that is usually associated with utilizing higher level language abstractions. Overall, these results emphasize the importance of the optimizations we have presented in this thesis as a step towards achieving high performance for high productivity languages.
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