# An Evaluation of Data-Parallel Compiler Support for Line-Sweep Applications

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# ABSTRACT

Data parallel compilers have long aimed to equal the performance of carefully hand-optimized parallel codes. For tightly-coupled applications based on line sweeps, this goal has been particularly elusive. In the Rice dHPF compiler, we have developed a wide spectrum of optimizations that enable us to closely approach hand-coded performance for tightly-coupled line sweep applications including the NAS SP and BT benchmark codes. From lightly-modified copies of standard serial versions of these benchmarks, dHPF generates MPI-based parallel code that is within 4% of the performance of the hand-crafted MPI implementations of these codes for a  $102^3$  problem size (Class B) on 64 processors. We describe and quantitatively evaluate the impact of partitioning, communication and memory hierarchy optimizations implemented by dHPF that enable us to approach handcoded performance with compiler-generated parallel code.

### 1. INTRODUCTION

A significant obstacle to the acceptance of data-parallel languages such as High Performance Fortran [10] has been that compilers for such languages do not routinely generate code that delivers performance comparable to that of carefullytuned, hand-coded parallel implementations. This has been especially true for tightly-coupled applications, which require communication within computational loops over distributed data dimensions [8]. Loosely synchronous applications, which only require communication between loop nests, can achieve good performance without sophisticated optimization. Without precise programmer control over communication and computation or sophisticated compiler optimization, it is impossible for programmers using dataparallel languages to approach the performance of handcoded parallel implementations with any amount of sourcelevel tuning. Since high performance is the principal motivation for developers of parallel implementations, scientists have preferred to hand-craft parallel applications using lower level programming models such as MPI [18], which can be tuned to deliver the necessary level of performance, despite the higher programming effort required.

The Rice dHPF compiler project [1, 2, 13] has focused on developing data-parallel compilation technology to enable generation of code whose performance is competitive with hand-coded parallel programs written using lower-level programming models. The dHPF compiler supports compilation of applications written in a Fortran 77 style augmented with HPF data distribution directives. dHPF implements a broad spectrum of novel data-parallel optimization techniques that enable it to produce high-quality parallel code for complex applications written in a natural style.

This paper uses the NAS SP and BT computational fluid dynamics application benchmarks [4] as a basis for a detailed performance study of code generated by dHPF for tightly-coupled line sweep applications. First, we compare the performance of several versions of parallel code generated by dHPF to other compiler-based and hand-coded parallelizations. These comparisons show that dHPF generates parameterized parallel code with scalable performance that closely approaches that of the hand-coded parallelizations by NASA. The remainder of the paper focuses on a quantitative evaluation of the performance contribution of dHPF's key optimizations. We determine how each key optimization contributes to the performance of dHPF's generated code by comparing overall application performance with and without that optimization.

Section 2 provides some brief background about the dHPF compiler and the NAS SP and BT benchmarks. Section 3 provides an overall comparison of the performance of dHPF-generated code against hand-coded and other compiler-based parallelizations of SP and BT. Section 4 compares the performance impact of several different compiler-based partitioning strategies suitable for SP and BT. Section 5 describes and evaluates communication optimizations in dHPF. Section 6 describes and evaluates an optimization to improve memory hierarchy utilization for off-processor data. Section 7 presents our conclusions and outlines open issues.

<sup>\*</sup>This work has been supported by the Los Alamos National Laboratory Computer Science Institute (LACSI) through LANL contract number 03891-99-23, as part of the prime contract (W-7405-ENG-36) between the Department of Energy and the Regents of the University of California.

# 2. BACKGROUND

# 2.1 The dHPF Compiler

The Rice dHPF compiler has several unique features and capabilities that distinguish it from other HPF compilers. First, it uses an abstract equational framework that enables much of its program analysis, optimization and code generation to be expressed as operations on symbolic sets of integer tuples [2]. Because of the generality of this formulation, it has been possible to implement a comprehensive collection of advanced optimizations that are broadly applicable.

Second, dHPF supports a more general computation partitioning model than other HPF compilers. With few exceptions, HPF compilers use simple variants of the *ownercomputes* rule [16] to partition computation among processors: partitioning of the instances for each statement (or loop iteration [3]) is based on a single (generally affine) reference. Unlike other compilers, dHPF permits independent computation partitionings for each program statement, where the partitioning for a statement maps computation of its instances onto the processors that own data accessed by one (or more) of a *set* of references. This computation decomposition model enables dHPF to support sophisticated partially-replicated partitionings, which have proven crucial for achieving high performance with the NAS SP and BT benchmarks as we describe in later sections.

Finally, the dHPF compiler provides novel compiler support for multipartitioning [6, 7], a family of sophisticated skewed-cyclic block distributions that were originally developed for hand-coded parallelization of tightly-coupled multidimensional line-sweep computations [5, 11, 14]. Multipartitioning enables decomposition of arrays of d > 2 dimensions among a set of processors so that for a line sweep computation along any dimension of an array, all processors are active in each step of the computation, load-balance is nearly perfect, and only coarse-grain communication is needed. We have extended HPF's standard data distribution directives with a new keyword, MULTI, that specifies a multipartitioned distribution. The MULTI keyword must be used in at least two dimensions and our implementation currently only supports \* in other dimensions of a multipartitioned data distribution.

## 2.2 NAS SP and BT Benchmark Codes

The NAS SP and BT application benchmarks [4] are tightlycoupled computational fluid dynamics codes that use linesweep computations to perform Alternating Direction Implicit (ADI) integration to solve discretized versions of the Navier-Stokes equation in three dimensions. SP solves scalar penta-diagonal systems, while BT solves block-tridiagonal systems. The codes both perform an iterative computation. In each time step, the codes have a loosely synchronous phase followed by tightly-coupled bi-directional sweeps along each of the three spatial dimensions. These codes have been widely used to evaluate the performance of parallel systems. Sophisticated hand-coded parallelizations of these codes developed by NASA provide a yardstick for evaluating the quality of code produced by parallelizing compilers.

The NAS 2.3-serial versions of the SP and BT benchmarks each consists of more than 3500 lines of Fortran 77 sequential code (including comments). To these, we added HPF data layout directives which account for 2% of the line count. To prepare these codes for use with dHPF, we manually inlined several procedures as we describe below. For SP, we inlined procedures lhsx and ninvr into the source code for procedure x\_solve; lhsy and pinvr into y\_solve; as well as lhsz and tzetar into z\_solve. For BT, we inlined lhsx, lhsy, and lhsz into x\_solve, y\_solve, and z\_solve respectively. The purpose of this inlining in SP and BT was to enable the dHPF compiler to globally restructure the local computation of these inlined routines so it could be overlapped with line-sweep communication. The inlining was necessary to eliminate a structural difference between the hand-coded MPI and serial versions that would have required interprocedural loop fusion to achieve automatically. In BT, one additional small inlining step was needed to expose interprocedurally carried dependences in the sweep computation to avoid having to place communication within the called routine.

## 2.3 Experimental Framework

All experiments reported in this paper were performed on on a dedicated SGI Origin 2000 parallel computer with 128 R10000 250MHz processors. Each processor possesses 4MB of L2 cache, 32KB of L1 data cache and 32KB of L1 instruction cache. All Fortran code generated by source-to-source compilation with dHPF along with the reference hand-coded versions of the NAS SP and BT benchmarks (version 2.3) were compiled using SGI MIPSpro compilers and linked with the SGI MPI native library. Detailed metrics, such as cache misses and MPI operations were measured using hardware performance counters under control of SGI's ssrun utility. In our experiments, we measured SP and BT executions using both class A  $(64^3)$  and class B  $(102^3)$  problem sizes. To see the impact of individual optimizations on scalability, we measured appropriate metrics for 16 and 64 processors. Metrics were collected for executions with only 10% of the standard number of iterations to keep execution time and trace file size manageable. This approach helps show differences in optimization impact for small and large data and processor sizes.

# 3. PERFORMANCE COMPARISON

In this section we compare the resulting performance of *four* different versions of the NAS SP and BT benchmarks<sup>1</sup>:

- NAS SP & BT Fortran 77 MPI hand-coded version, implemented by the NASA Ames Research Laboratory
- NAS SP & BT (multipartitioned) compiled with dHPF from sequential sources
- NAS SP & BT (2D block) compiled with dHPF from sequential sources
- NAS SP & BT (1D block, transpose) compiled with the Portland Group's pgHPF from their HPF sources

The Portland Group's (PGI) versions of the NAS SP and BT benchmarks were obtained directly from the PGI WWW site [15].

<sup>&</sup>lt;sup>1</sup>The source and generated code used for our experiments is available at www.cs.rice.edu/~dsystem/dhpf/pact2002.



Figure 1: Parallel efficiency for NAS SP (class 'B').



Figure 2: Parallel efficiency for NAS SP (class 'A').

PGI's HPF versions of these codes use a 1D BLOCK data distribution and perform full tranposes of the principal arrays between the sweep phases of the computation. Since their compiler does not support array redistribution, their implementation uses *two* copies of two large 4D arrays, where the copies are related by a transpose. The PGI versions of the NAS SP and BT computations were written from scratch to avoid the limitations of the PGI compiler. A discussion of this topic can be found elsewhere [1].

We ran these code versions on 1–64 processors for the class 'A'  $(64^3)$  problem size and 1–81 processors for the larger class 'B'  $(102^3)$  problem size. This range of problem sizes and processor counts is intended to show the performance of different variants of the benchmarks for a range of perprocessor data sizes and communication-to-computation ratios.



Figure 3: Parallel efficiency for NAS BT (class 'B').



Figure 4: Parallel efficiency for NAS BT (class 'A').

Figures 1 and 2 compare the efficiency of four different parallelizations of the NAS SP benchmark for the class 'B' and 'A' problem sizes respectively. Figures 3 and 4 present the efficiency comparisons for the NAS BT benchmark's class 'B' and 'A' problem sizes.

For each parallelization  $\rho$ , the efficiency metric is computed as  $\frac{t_s}{P \times t_p(P,\rho)}$ . In this equation,  $t_s$  is the execution time of the original sequential version implemented by the NAS group at the NASA Ames Research Laboratory; P is the number of processors;  $t_p(P,\rho)$  is the time for the parallel execution on P processors using parallelization  $\rho$ . Using this metric, perfect speedup would appear as a horizontal line at efficiency 1.0. We use efficiency rather than speedup or execution time as our comparison metric because it enables us to accurately gauge the relative performance of multiple versions across the *entire* range of processor counts. The graphs show that the efficiency of the hand-coded MPIbased parallelizations based on a multipartitioned data distribution is excellent, yielding an average parallel efficiency of 1.20 for SP class 'A', 0.94 for SP class 'B', 0.97 for BT class 'A' and 1.02 for BT class 'B'. Thus, the hand-coded versions achieve nearly linear speedup.

The dHPF-generated multipartitioned code achieves similar parallel efficiency and near-linear speedup for most processor counts, demonstrating the effectiveness of our compilation and optimization technologies. Its average efficiency across the range of processors is 1.11 for SP class 'A', 0.99 for SP class 'B', 0.94 for BT class 'A' and 1.04 for BT class 'B'. The dHPF compiler achieves this level of performance for code generated for a symbolic number of processors, whereas the hand-coded MPI implementations have the number of processors compiled directly into them.

The remaining gaps between the performance of the dHPFgenerated code and that of the hand-coded MPI can be attributed to two factors. First, the dHPF-generated code has a higher secondary cache miss rate on large numbers of processors due to interference between code and data in the unified L2 cache of the Origin 2000. The memory footprint of dHPF's generated code is substantially larger than that of the hand-coded version due to its generality. Second, on 81 processors the class 'B' benchmarks suffer from load imbalance on tiles along some of the surfaces of multipartitioned arrays. dHPF uses a fixed block size (namely,  $\left\lceil \frac{s}{t} \right\rceil$ , where s is the extent of the dimension and t is the number of tiles in that dimension) for tiles along a dimension; using this scheme, tiles at the rightmost end may have fewer elements. For the class 'B' benchmarks on 81 processors, this leads to partially-imbalanced computation with 8 tiles of size 12 and a tile of size 6 at the right boundary. In contrast, the tiling used by the hand-coded MPI ensures each processor has either  $\lfloor \frac{s}{t} \rfloor$  and  $\lceil \frac{s}{t} \rceil$  elements, which leads to more even load balance.

The performance of the dHPF-generated 2D block partitioning using coarse-grain pipelining (CGP) [1] is respectable, with an average efficiency of 0.80 for SP class 'A' and 0.65 for class 'B'. Due to a compiler limitation, the CGP version of SP uses overlap regions for storing off-processor data; this places it at a disadvantage with respect to the multipartitioned version, which uses its data directly out of communication buffers. Even if this limitation were eliminated, it would not boost the performance to the level achieved by the multipartitioned codes; we discuss this issue in more detail in the next section. Due to a limitation in our dependence analysis, we were not able to obtain results for BT using this partitioning scheme.

We also compare the efficiency of the version of the NAS SP and BT benchmarks written by PGI which use 1D block distributions with transposes between sweep phases. These codes, compiled with version 2.4 of the *pghpf* compiler, achieve an average efficiency of 0.69 for SP class 'A', 0.53 for SP class 'B', 0.87 for BT class 'A' and 0.99 for BT class 'B'. Despite good average efficiency for BT, Figures 3 and 4 show that the PGI version has considerably lower scalability than the dHPF and MPI versions. The full array transposes cause communication volume proportional to the full 3D array *vol*-

*ume*, whereas communication volume for multipartitioning is proportional to the *surface area* of the partitioned tiles.

# 4. DATA PARTITIONING SUPPORT

The previous section showed considerable variations in performance between codes using different partitioning strategies. Here, we quantitatively evaluate the impact of the partitioning strategy on overall execution time and communication volume. The data partitioning strategy employed in a parallel code influences an application's communication pattern, communication schedule and communication volume, as well as its local node behavior. The data partitioning choice is a key determinant of an application's overall performance.

Here, we use the SP and BT benchmarks to compare the impact of three partitioning choices suitable for tightly-coupled line sweep applications. Tables 1 and 2 present ratios of execution time and communication volume that, respectively, compare the 2D-block partitioned versions of the codes and PGI's 1D-block partitioned versions of the codes with respect to the dHPF multipartitioned versions.

Benchmark	16 proc.	64 proc.
Time SP 'A' CGP Comm. Vol. SP 'A' CGP	$1.45 \\ 1.09$	$1.23 \\ 1.08$
Time SP 'B' CGP Comm. Vol. SP 'B' CGP	$1.76 \\ 1.09$	$1.58 \\ 1.12$

Table 1: 2D-block CGP vs. multipartitioning.

Benchmark	16 proc.	64 proc.
Time SP 'A' PGI	1.40	1.61
Comm. Vol. SP 'A' PGI	3.29	3.27
Time SP 'B' PGI	1.91	1.98
Comm. Vol. SP 'B' PGI	4.28	3.28
Time BT 'A' PGI	0.85	1.34
Comm. Vol. BT 'A' PGI	3.11	3.16
Time BT 'B' PGI	0.95	1.69
Comm. Vol. BT 'B' PGI	4.00	3.13

Table 2: 1D-block transpose (PGI) vs. multipartitioning.

From Table 1, we see that the relative increase in communication volume with respect to multipartitioning is modest for both problem sizes. The increase in execution time is not directly proportional to the increase in communication volume. Previous studies have shown that the principal factor driving the increase in execution time is that the coarsegrain pipelining version incurs more serialization [6].

From Table 2, it is clear that the communication volume of PGI's 1D-block transpose implementation is a factor of 3–4 larger than that of dHPF's multipartitioned version. For smaller numbers of processors, this strategy is reasonably competitive, but for larger numbers of processors, efficiency degrades. Without more detailed analysis of their implementation, the precise reasons for differences in scalability and performance in response to data and processor scaling are not clear.

# 5. COMMUNICATION OPTIMIZATIONS

On modern architectures, achieving high-performance with a parallel application is only possible if processors communicate infrequently. The high cost of synchronizing and exchanging data drives the need to reduce the number of communications. This is true on NUMA-style distributed shared memory machines, multicomputers and clusters.

We have designed and implemented a broad range of communication analysis and optimization techniques in the dHPF compiler. These techniques improve the precision of processor communication analysis, reduce the frequency of messages between processors and reduce the volume of data communicated. We also describe some language extensions that assist the compiler in generating high performance code.

## 5.1 HPF/JA-inspired Extensions

The Japanese Association for High Performance Fortran proposed several new HPF directives to enable users to fine-tune performance by precisely controlling communication placement and providing a limited means for partially replicating computation. These directives are known as the HPF/JA extensions [17]. dHPF implements variants of several of the HPF/JA extensions as we describe below.

To enable an HPF programmer to avoid unnecessary communication that can't be eliminated automatically by an HPF compiler without sophisticated analysis, dHPF supports the HPF/JA LOCAL directive. The LOCAL directive asserts to the compiler that communication for a set of distributed arrays (specified as parameters to LOCAL) is not needed in a particular scope.

Enabling an HPF programmer to partially replicate computation to reduce communication can be important for high performance. For this reason, we provide extended support in dHPF for the ON HOME directive to give users control over partially replicating computation in shadow regions. This support was inspired by the HPF/JA ON EXT\_HOME directive, but enables more precise replication control. The HPF/JA ON EXT\_HOME directive enables computation to be partiallyreplicated onto all elements in an array's shadow regions. A drawback of the ON EXT\_HOME is that it can cause computation for elements in an arrays shadow region that are not needed. To avoid this undesirable effect, we extended HPF's ON HOME directive to allow multiple ON HOME references to be specified; this provides more precise control over partial replication than ON EXT\_HOME and enables the manual specification of *partially replicated* computation partitionings similar to the ones generated semi-automatically by the dHPF compiler for localization [1]. Figure 5 shows how using our extended ON HOME directive enables precise control over partial replication of computation to selectively fill a portion of the shadow region (not all of the shadow region may be needed in a particular step of the computation). In the figure, each inner square represents an owned section for a processor; the enclosing squares represent the processor's shadow region. Our extended ON HOME directive enables us to include or exclude the "corners" that would be filled by the HPF/JA EXT\_HOME directive.

The HPF/JA  ${\tt REFLECT}$  directive was designed to support explicit data replication into shadow regions of neighbors.



Figure 5: Extended ON HOME vs. EXT\_HOME



Figure 6: Extended REFLECT directive

REFLECT was designed for use in conjunction with the LOCAL directive to avoid redundant communication of values. At the point a REFLECT directive occurs in the code, communication will be scheduled to fill each processor's shadow regions for a distributed array with the latest values from neighboring processors. In dHPF, we implemented support for an extension of the **REFLECT** directive that enables more precise control over the filling of the shadow regions. Our extension to REFLECT allows the programmer to specify the dimensions and the depth of an array's shadow regions to fill. By specifying a single dimension and width, we can selectively fill all or part of the shadow region along that dimension. If corner elements are needed, multiple dimensions can be specified at once in a single entry, which will include them. Figure 6 shows 2D array a's left and top shadow regions selectively filled in from the values owned by its left and top neighbors.

In the following sections, we describe how we use these directives to tune application performance.

# 5.2 Partially Replicated Computation

The dHPF compiler's ability to compute statements on multiple home locations enables it to partially replicate computation. This approach, used judiciously, can significantly reduce communication costs and even, in some cases, eliminate communication altogether for certain arrays.

With dHPF, computation can be partially replicated either automatically by the compiler [1], or manually using the extended ON HOME directive. Figure 7 illustrates the technique for a simple case: computation along the columns has been replicated on both processors (the owning processor and the non-owning one) by using an extended computation partitioning as described in section 5.1. The figure shows the locally-allocated sections of distributed array **a** for two different processors. The diagonally-striped regions represent the each processor's "owned" section; the squares enclosing



Figure 7: Partially Replicated Computation along a processor boundary

each owned section represent surrounding shadow regions. The shaded portion of processor p + 1's shadow region represents the replicated computation of one column owned by processor p, as shown by the shaded portion of processor p's owned region.

For SP and BT, partial computation replication enables us to completely eliminate boundary communication for several large distributed arrays, without introducing any additional communication. In the BT benchmark, partial computation replication eliminates communication for 5D arrays, fjac and njac in the lhsx, lhsy, and lhsz routines. In both the BT and SP benchmarks, partial computation replication completely eliminates communication for six 3D arrays in the compute\_rhs routine. The computation of these values can be replicated at boundaries using a 4D array that must be communicated anyway. The hand-coded parallelizations exploit this partial replication of computation too.

## 5.2.1 Performance Impact

Table 3 compares the relative performance of versions with and without partial replication of computation. We present the ratios of execution time and communication volume between the dHPF-generated multipartitioned versions of SP and BT without partial computation replication and those with partial replication. The partially replicated versions uses both compiler-derived and explicitly-specified replication using the extended ON HOME directive. The results in Table 3 show that without partial replication of computation, both execution time and communication volume increase significantly.

Benchmark	16 proc.	64 proc.
Time SP 'A'	1.18	1.36
Comm. Vol. SP 'A'	1.33	1.36
Time SP 'B'	1.12	1.21
Comm. Vol. SP 'B'	1.33	1.35
Time BT 'A'	1.35	1.58
Comm. Vol. BT 'A'	2.94	2.96
Time BT 'B'	1.13	1.62
Comm. Vol. BT 'B'	2.97	2.97

### Table 3: Impact of partial computation replication.

This optimization contributes significantly to scalability: its relative improvement in execution time is more significant when communication time is not dominated by computation; for this reason, we observe a higher impact for the smaller class 'A' problem size on a large number of processors. It is worth noting that our experiments were executed on a tightly-coupled parallel computer with a highbandwidth low-latency interconnect; this reduces the performance penalty associated with higher message volume.

### 5.2.2 Interprocedural Communication Elimination

dHPF's communication analysis and generation is procedurebased; dHPF does not currently support interprocedural communication analysis and placement. However, using our HPF/JA-inspired directive extensions enables us to eliminate communication of values that we know are available as a side effect of partially-replicated computation or communication elsewhere in the program.

We present results on the impact of using the HPF/JA extended directives to eliminate communication across procedures on the selected benchmarks. Table 4 presents ratios comparing the execution time and communication volume of dHPF-generated multipartitioned versions of SP and BT without using the HPF/JA directives to eliminate communication across procedures with respect to the versions that use them. Without the directives, communication volume and execution time increase 10–20%.

Benchmark	16 proc.	64 proc.
Time SP 'A'	1.05	1.15
Comm. Vol. SP 'A'	1.12	1.13
Time SP 'B'	1.05	1.08
Comm. Vol. SP 'B'	1.11	1.12
Time BT 'A'	1.12	1.09
Comm. Vol. BT 'A'	1.18	1.18
Time BT 'B'	1.06	1.20
Comm. Vol. BT 'B'	1.18	1.18

Table 4: Impact of using the HPF/JA extended directives to eliminate communication interprocedurally.

In SP, we use the LOCAL directive to eliminate communication for three 3D arrays in the lhsx, lhsy and lhsz routines for values that are available locally because their computation was partially replicated in the compute\_rhs routine using extended ON HOME.

In BT, we use the LOCAL directive to eliminate communication for a 3D array u in the lhsx, lhsy and lhsz routines. The required off-processor values for u had already been communicated into the shadow region in routine compute\_rhs by a REFLECT directive.

The combination of the HPF/JA LOCAL, extended ON HOME and REFLECT directives obtain all of the benefits of interprocedural communication analysis, without its complexity.

## 5.3 Coalescing

For best performance, an HPF compiler should minimize the number and volume of messages. Analyzing and generating messages for each non-local reference to a distributed array in isolation produces too many messages and the same values might be transferred multiple times.

```
CHPF$ distribute a(*, block), b(*, block) onto P
do j = 2, n
do i = 1, n
a(i, j) = b(i, j - 1) + c ! ON_HOME a(i, j)
a(i, j) = a(i, j) + d + b(i, j - 1) ! ON_HOME a(i, j)
end do
end do
```

Figure 8: Simple overlapping non-local data references.

```
CHPF$ distribute a(*, block), b(*, block) onto P
do j = 2, n
    do i = 1, n - 1, 2
        a(i, j) = b(i, j - 1) + c ! ON_HOME a(i, j)
        a(i + 1, j) = d + b(i, j) ! ON_HOME a(i + 1, j)
    end do
end do
```

# Figure 9: Complex overlapping non-local data references

In dHPF, a communication set is initially computed and placed separately for each individual non-local reference. A communication set is represented in terms of an ON HOME reference (corresponding the computation partitioning where the data is required) and a non-local reference. Both references are represented in terms of value numbers that appear in their subscripts (if any). Whenever possible, dHPF vectorizes communication and hoists it out of loops. When multiple communication events are scheduled at the same location in the code, dHPF tries to coalesce them to avoid communicating the same data multiple times.

Consider the code in Figure 8, an HPF compiler should generate a single message to communicate a single copy of the off-processor data required by both references to b(i, j - 1). However, detecting when sets of non-local data for multiple references overlap is not always so simple. References that are *not* syntactically equivalent may require identical or overlapping non-local data. In Figure 9, references b(i, j - 1) and b(i, j) require identical non-local values and can be satisfied by a single message. To avoid communicating duplicate values, overlap between sets of non-local data required by different loop nests should be considered as well, as shown in Figure 10.

### 5.3.1 Normalization

To avoid communicating duplicate values, dHPF uses a normalization scheme as a basis for determining when communication sets for different references overlap. To compensate for differences in computation partitionings selected for different statements, normalization rewrites the value numbers representing a communication set into a form relative to its ON\_HOME reference expressed in a *canonical* form. In our discussion of normalization, we refer to the ON\_HOME reference for a communication set as the *computation partitioning (CP) reference* and the non-local reference as the *data reference*.

dHPF's value-number based representation for communication sets has the disadvantage that non-local references that arise in different loops are incomparable because their sub-



Figure 10: Coalescing non-local data across loops.

scripts have different value number representations. To enable us to detect when such references may require overlapping sets of non-local data, we convert all data and CP references to use a canonical set of value numbers for the loop induction variables involved.

We apply our normalization algorithm to communication sets represented in terms of value numbers based on affine subscript expressions of the form ai + b, where *i* is an induction variable, *a* is a known integer constant and *b* is a (possibly symbolic) constant. This restriction comes from the need to compute a symbolic inverse function for such expressions<sup>2</sup>. If this restriction is not met, the communication set is left in its original form.

A communication set is in normal form if:

- The CP reference is of the form  $A(i_1, i_2, i_3, ..., i_n)$
- The data reference is of the form  $A(a_1i_1' + b_1, a_2i_2' + b_2, a_3i_3' + b_3, ..., a_ni_n' + b_n)$

where A is an n-dimensional array, each  $i_j$  is an induction variable or a constant, and each  $i'_j$  corresponds to a unique  $i_k$ . We say that a particular subscript expression in the CP reference is normalized if it is of the form  $i_j$ , where  $i_j$  is an induction variable or a constant.

If a communication set is not in normal form but meets our restriction of affine subscript expressions, we normalize it by computing symbolic inverse functions for each nonnormalized CP subscript position. We then apply this function to each subscript position in both the CP and data references. After this step, *only* the data reference has subscripts of the form  $a_j i_j + b_j$ . Each subscript position gets a canonical value number represented by an artificial induction variable that has the range and stride of the original  $a_j i_j + b_j$  subscript expression. We apply this normalization step to all communication events at a particular location in the code before attempting to coalesce them.

### 5.3.2 Coalescing Normalized Communication Sets

Following normalization, we coalesce communication events to eliminate redundant data transfers. There are two types

 $<sup>^2 \</sup>rm Normalization$  could be extended to support more complex affine expressions without too much difficulty.

of coalescing operations that can be applied to a pair of communication events:

- Subsumption: Identify and eliminate a communication event (nearly) completely covered by another.
- Union: Identify and fuse partially overlapping communication events, which do not cover one another.

Both operations eliminate communication redundancy, however these two cases are handled separately at compile time.

For both cases, the coalescing algorithm first tests to see if communication events are compatible. To be compatible, both communication events must correspond to the same distributed array, be reached by the same HPF alignment and distribution directives, have the same communication type (read or write) and be placed at the same location in the intermediate code.

### 5.3.2.1 Subsumption

The subsumption of one communication event by another requires that the non-local data of the subsuming event be a superset of that of the subsumed one. According to the model for normalized data and CP references, this implies that both messages represent data shifts of constant width in one or more dimensions. For example, a single-dimensional shift of a submatrix owned by a processor would correspond to sending a few rows or columns to a neighboring processor. A pair of shifts must be in the same direction and along the same dimensions for communication events to be compatible. For instance, trying to coalesce a shift that sends two rows and a shift that sends two columns of a submatrix is infeasible.

For two compatible shift communications, where the *shift* width of one is larger than that of the other, the volume of data transferred can be reduced by completely eliminating the smaller shift since its data values will be provided by the larger one. For dimensions not involved in a shift, the ranges of the data reference subscript value numbers in the subsuming communication event must be supersets of the corresponding ranges in the subsumed event. For data dimensions involved in a shift, subsumption does not require that their range be a strict superset of the corresponding ranges in the subsumed event. If the subsuming communication has a wider shift width, but doesn't have a large enough range for its loop induction variable, the coalescer synthesizes a new induction variable with extended range to cover the induction variable in the subsumed event as well. This range-extension technique is very effective for avoiding partially-redundant messages and generating simple communication code.

### 5.3.2.2 Union

Coalescing partially overlapping communication events requires less strict conditions than subsumption. Given normalized data and CP references, The coalescer will only try to union communication events that have a common shift dimensions and directions.

The dHPF compiler uses the Omega library [12] to implement operations on integer tuples for its communication analysis and code generation [2]. We apply integer set-based analysis to determine the profitability of unioning two communication events. We construct sets that represent the data accessed by the non-local references of each communication event. If these sets intersect, this implies that the communication sets for some pair of processors *may* intersect. In this case, the algorithm will coalesce the two communication events by unioning them. If the sets do not intersect, then the communication sets for any pair of processors also do not intersect, which implies there is no redundancy to eliminate, so coalescing is not performed.

As described in the previous section, generation of a coalesced communication set can require that new induction variables be synthesized with extended range. When unioning communication sets, this transformation is applied to an induction variable appearing in any dimension as necessary.

### 5.3.3 Performance Impact

Table 5 presents the relative performance of codes where coalescing was applied with and without normalization. (Coalescing without normalization may overlook opportunities for eliminating redundancy.) We present the ratios of execution time and communication volume between the dHPFgenerated multipartitioned versions of SP and BT with nonnormalized coalescing and those with normalized coalescing. The results show that without normalization, both execution time and MPI communication volume increase.

Benchmark	16 proc.	64 proc.
Time SP 'A'	1.41	2.10
Comm. Vol. SP 'A'	1.70	1.71
Time SP 'B'	1.28	1.65
Comm. Vol. SP 'B'	1.71	1.71
Time BT 'A'	1.02	1.24
Comm. Vol. BT 'A'	1.32	1.32
Time BT 'B'	1.04	1.08
Comm. Vol. BT 'B'	1.32	1.33

Table 5: Impact of normalized coalescing.

We found that the reduction in execution time due to normalized coalescing is mostly due to reduced communication volume in the critical tightly-coupled line sweep routines  $(x\_solve, y\_solve, z\_solve)$  of SP and BT. Loosely coupled communication is also reduced, but its impact is not as significant on machines with a high-bandwidth, low-latency interconnect such as the Origin 2000. The greater execution time of the non-normalized version for SP cannot be attributed completely to the extra communication volume, because non-normalized coalescing produces sets which are not rectangular sections; this leads to less efficient management of communicated data, as explained in more detail in Section 6.1.

After both partial replication of computation and normalized coalescing, we found that the communication volume for the dHPF-generated code for SP was within 1% of the volume for the hand-coded versions. For BT, we found that the volume in the dHPF-generated code was actually lower by 5% (class B) and 17% (class A) than that of the handcoded versions.

## 5.4 Multi-variable Aggregation

Often, applications access multiple arrays in a similar fashion. When off-processor data is needed, this can lead a pair of processors to communicate multiple variables at the same point in the program. Rather than sending each variable in a separate message, dHPF reduces communication overhead by shipping multiple arrays in a single message in such cases. dHPF's implementation strategy of aggregation is described elsewhere [6]. Here we focus on message aggregation's impact on performance. Table 6 presents ratios for execution time and communication frequency that compare the cost of of non-aggregated versions of the codes with respect to aggregated versions. We present the ratios of execution time and communication frequency between the dHPF-generated multipartitioned versions of SP and BT without aggregation and those with aggregation. Ratios larger than one indicate an increase in cost without aggregation. While aggregation reduces message frequency considerably for both SP and BT in all cases, the impact of aggregation on execution time for BT is small because of its high computation to communication ratio. However, the latency reduction aggregation yields eventually becomes important when scaling a computation to a large enough number of processors so that communication time becomes significant with respect to computation. SP has a higher communication/computation ratio and thus aggregation has a larger impact. For SP, as we drop back from the larger class 'B' to the smaller class 'A' problem size, and/or increase the number of processors, the execution time impact of aggregation increases. For the 64 processor execution of SP using the class 'A' problem size, aggregation cuts execution time by 10%.

Benchmark	16 proc.	64 proc.
Time SP 'A'	1.02	1.10
Comm. Freq. SP 'A'	2.12	2.31
Time SP 'B'	1.01	1.03
Comm. Freq. SP 'B'	2.12	2.31
Time BT 'A'	1.00	1.01
Comm. Freq. BT 'A'	1.37	1.43
Time BT 'B'	1.03	1.02
Comm. Freq. BT 'B'	1.37	1.43

Table 6: Impact of aggregation.

## 6. MEMORY HIERARCHY ISSUES

Today's computer systems rely on multi-level memory hierarchies to bridge the gap between the speed of processors and memory. To achieve good performance, applications must use the memory hierarchy effectively.

In the dHPF compiler, we have implemented many techniques to improve memory hierarchy utilization. These include padding of dynamically allocated arrays, inter-array tile grouping (which lays out corresponding tiles from different arrays consecutively in memory to reduce conflicts), and an arena-based communication buffer management scheme to reduce the cache footprint of communication buffers. Here we discuss a novel compiler-based technique for managing off-processor data that increases cache efficiency.

```
CHPF$ distribute a(*, block) onto P
CHPF$ shadow a(0, 1:0)
do j = 2, n
do i = 1, n
a(i, j) = a(i, j - 1) + c ! ON_HOME a(i, j)
end do
end do
```

Figure 11: An overlap region on an array in HPF.

```
do j = max(2, my_j_lo), min(my_j_lo + j_blocksize, n)
    do i = 1, n
        a(i, j) = a(i, j - 1) + c
        end do
end do
```

Figure 12: SPMD code using an overlap region.

## 6.1 Overlap Regions vs. Direct-Access Buffers

Overlap regions [9], in which a processor allocates additional storage around the boundary of data it owns to store neighboring off-processor data, are a commonly used technique by compilers and application developers. The dHPF compiler allows the use of overlap regions for distributed arrays. An HPF2 SHADOW directive specifies the extent of an overlap region for an array on a dimension-by-dimension basis. Figure 11 shows HPF source code with an overlap region specified for array a; the SHADOW directive specifies a lower overlap of width one in the second dimension.

Overlap regions are convenient because they enable uniform access to both local and off-processor data, which leads to simple code for partitioned loops. For example, in Figure 12 overlap regions make it possible to access a(i, j - 1) for j equal to the processor boundary  $(my_j_lo)$ .

While overlap regions lead to simple SPMD code, there are three ways that using them can degrade performance. First, any loop accessing an array that has overlap regions allocated, which does not use most of the overlap region in each of the dimensions for which it is provided, suffers from both reduced spatial reuse (values in the overlap region may be fetched into cache and not used) and less effective cache utilization (some cache sets may be underutilized because unaccessed overlap regions map to them). Second, if data is received into a message buffer and then copied into overlap regions, there will be two live copies of the data occupying space in the cache. Third, copying the data from a communication buffer into an overlap region can be costly, particularly if the data in the overlap region is non-contiguous.<sup>3</sup>

To avoid the cache inefficiency that comes with using multidimensional overlap regions for single-dimensional shift communication, the dHPF compiler supports accessing remote data directly out of communication buffers. This has the dual advantage of eliminating the unpacking phase for the receiving processor, as well as eliminating the need for overlap regions on the receiving processor's tile. Directly accessing the buffers, introduces two modes of access for array references: boundary remote data, accessed out of the buffer and interior data accessed out of the array.

 $<sup>^{3}</sup>Any$  data movement in modern machines is costly!

```
do j = max(2, my_j_lo), min(my_j_lo + j_blocksize, n)
    do i = 1, n
        if (j - 1 .lt. my_j_lo) then ! REMOTE data
        t1 = buffer_a(i, j - my_j_lo)
        else ! LOCAL data
        t1 = a(i, j - 1)
        end if
        a(i, j) = t1 + c
        end do
end do
```

Figure 13: SPMD code using a direct-access buffer.

```
do j = my_j_lo, my_j_lo + 1
    do i = 1, n
        a(i, j) = buffer_a(i, j - my_j_lo) + c
    end do
end do
do j = my_j_lo + 2, min(my_j_lo + j_blocksize, n)
    do i = 1, n
        a(i, j) = a(i - 1, j) + c
    end do
end do
```

#### Figure 14: Optimized use of a direct-access buffer.

If the compiler were to generate naive code for data that may reside either in a buffer or in a local array, the access would require a conditional test, which may be costly. Figure 13 shows naive code for this situation. To avoid the overhead of this approach, the dHPF compiler splits a loop nest that accesses non-local data out of buffers into a loop nest whose iterations *may* require remote data and those that *must* access data only from the local array.

Loop splitting in this manner eliminates conditionals from the interior section of the loop nest, but may not eliminate conditionals in the non-local section of the loop nest. Figure 14 shows a split loop with one iteration space accessing data for array **a** only out of the local array section, and the other iteration space accessing data out of a buffer. In this case, it was possible to eliminate all conditionals for the non-local reference, because the set of non-local iterations for the statement is a subset of the non-local iterations for the reference. In a more general case, the non-local iteration space may not access exclusively off-processor data; a conditional would be required in this case.

### 6.1.1 Aggregation and Direct Buffer Access

Direct buffer-access is very useful, especially in combination with communication aggregation. Incoming non-local data for different arrays or disjoint sections of the same array is laid out sequentially in the buffer. dHPF generates code for directly accessing such data by using pointers into each contiguous section in the buffer. With this scheme, the dHPF compiler supports direct access to buffers comprised of *unions of constantly-strided rectangular sections*.

### 6.1.2 Performance Impact

We compare the efficiency of using direct-access buffers and overlap regions for the selected benchmarks. Table 7 presents ratios that compare the execution time and L2 cache misses for versions using overlap regions with respect to versions using direct access buffers. We present the ratios of execution time and L2 data cache misses between the dHPF-generated multipartitioned versions of SP and BT without direct buffer access and those with it. Both the execution time and L2 cache misses increase without direct access buffers.

Benchmark	16 proc.	64 proc.
Time SP 'A'	1.13	1.40
L2 Misses SP 'A'	1.22	1.41
Time SP 'B'	1.19	1.10
L2 Misses SP 'B'	1.09	1.11
Time BT 'A'	1.02	1.03
L2 Misses BT 'A'	1.01	1.09
Time BT 'B'	1.04	1.04
L2 Misses BT 'B'	1.00	1.01

Table 7: Impact of direct-access buffers on SP and BT.

Direct-access buffers play a significant role in reducing L2 data cache misses by avoiding extra copies into shadow regions and reducing the memory footprint of large multidimensional arrays. In particular, they are very important for the tightly-coupled line sweep phases of the SP and BT benchmarks and the **lhs** and **rhs** arrays they use.

# 7. CONCLUSIONS

The results presented in this paper show that the dHPF compiler is able to virtually match the performance of sophisticated, carefully tuned, hand-coded parallelizations of the NAS SP and BT benchmarks. To our knowledge this is the first time that data-parallel compilers have been able to deliver performance at this level for such tightly-coupled line sweep applications. Achieving this level of performance was not a matter of just implementing a few "big-ticket" optimizations. Delivering hand-coded performance with a dataparallel compiler requires a surprisingly broad spectrum of analysis and code generation techniques. With the exception of support for the multipartitioning data distribution, the optimizations we implemented in dHPF have broad applicability. We believe that they will be useful for other parallel architectures, although their relative importance may differ depending upon the parameters of the architecture.

Our experience is that everything affects scalability. Excellent parallel performance requires not only a good parallel algorithm, but also excellent resource utilization on the target parallel machine. A fast, scalable program must make effective use of the processors, memory hierarchy and processor interconnect. For data parallel compilers, the implications are clear: discovering parallelism is only the beginning; exploiting it effectively is not necessarily as glamorous, but it is critically important. Optimizations must aim to effectively utilize all classes of resources in a parallel system. Only by targeting each potential source of inefficiency in compilergenerated parallel code can data-parallel compilers achieve the level of performance that will make them acceptable to application scientists.

Our ongoing work is focused on exploring data-parallel compiler issues for other types of numerical applications.

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