# Algorithmic Bombardment for the Iterative Solution of Linear Systems: A Poly-Iterative Approach* 

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#### Abstract

Many algorithms employing short recurrences have been developed for iteratively solving linear systems. Yet when the matrix is nonsymmetric or indefinite, or both, it is difficult to predict which method will perform best, or indeed, converge at all. Attempts have been made to classify the matrix properties for which a particular method will yield a satisfactory solution, but "luck" still plays large role. This report describes the implementation of a poly-iterative solver. Here we apply three algorithms simultaneously to the system, in the hope that at least one will converge to the solution. While this approach has merit in a sequential computing environment, it is even more valuable in a parallel environment. By combining global communications, the cost of three methods can be reduced to that of a single method.


## 1 Introduction

Many iterative methods for solving real nonsymmetric linear systems

$$
\begin{equation*}
A x=b \tag{1}
\end{equation*}
$$

have been proposed. A popular choice is GMRES [20], which in the absence of rounding error, guarantees convergence in $n$ steps for an order $n$ matrix, but its memory requirements often rule out solving large systems ${ }^{1}$. Variations for reducing the length of the recurrences have been proposed (using restarting or truncation), but this compromises the convergence theory. Also, in a distributed memory environment, the communication requirements of the increasing number of inner products that must be performed can severely slow the time to solution ${ }^{2}$.

Recent developments have made less memory and communication intensive algorithms more viable. Each offers theoretical justification for its convergence properties, and if the user has certain

[^0]information concerning the spectrum of the matrix, it is possible to select a method which should work well. However, when this is not the case, time (and expense) may be wasted when an algorithm terminates without convergence. For example, this is often true with nonlinear problems, where many linear systems are generated, and the properties of the matrix change. Also, an algorithm may stall, diverge, or break down even when the matrix properties indicate the method should converge.

Moreover, these properties may be so affected by rounding error that we lose the ability to predict their behavior. In fact, machines using different methods for performing the arithmetic may yield different results. For example, the IBM RS6000 performs the multiply and add before truncating, as opposed to say a Sun SPARCstation ipx which uses IEEE arithmetic, truncating after the multiply and again after the add. While the RS6000 is designed to be more accurate, it is still finite arithmetic, and with these iterative solvers, any inaccuracy may alter the behavior of the algorithm.

In response to this problem, we propose applying several iteration schemes, in parallel using the same data set, in the hope that at least one will yield an acceptable approximation to the true solution. Although the number of operations per iterative step equals the sum of the operations of the individual methods, we believe that when knowledge of matrix properties is lacking or incomplete, the extra floating point computation and memory requirements are outweighed by three factors:

1. An increased probability of finding the solution.
2. An efficient parallel implementation. By iterating in lock-step, i.e. the algorithms are always on the same iteration count, we gain time savings by combining overlapping communication (inner products, matrix-vector products, preconditioner solves).
3. Increased floating point performance. Depending upon the structure of the matrix, an efficient matrix-vector product may be constructed so as to make use of data locality. This may also be true for preconditioning.

The high cost of communication is the weak link in distributed memory architectures, so by combining the distribution of global data, we can improve upon the performance of the poly-iterative approach. For example, since each method is operating on the same matrix, and each matrix-vector multiplication requires communication to update the product vector, we may be able to minimize executation time by combining the $k$ product vectors, where $k$ is the number of methods whose matrix-vector product operands can be combined into one buffer for communication. Additionally, when the structure of a matrix allows it, we can multiply an $n$-dimensional matrix by a $n \times k$ matrix, rather than a single vector, thereby improving the floating point efficiency of the computation [6]. Moreover, we can combine other communication, such as that required for the computation and distribution of inner products and residual norms. This is discussed in more detail in section 3.3.

The other mathematical operations (vector updates, certain preconditioners, scalar operations, etc.) are computed in parallel, requiring no communication. The overall effect of the extra work is a function of the sparsity of the original coefficient matrix.

For our implementation, we have selected three iterative algorithms for the solution ${ }^{3}$ of (1):

- Quasi-Minimal Residual (QMR),
- Conjugate Gradient Squared (CGS), and
- Biconjugate Gradient Stabilized (BiCGSTAB).

[^1]
## 2 The Algorithms

In this section we give a brief description of the methods that make up our implementation of the algorithmic bombardment algorithm.

When the coefficient matrix of the linear system is a symmetric positive definite matrix, the traditional iterative algorithm of choice is the Conjugate Gradient (CG) method [18]. However, when the coefficient matrix is nonsymmetric, CG typically fails to find the solution. The Biconjugate Gradient method [12], [19], rather than relying on a single sequence of residuals (as does CG), creates another sequence $\{\tilde{r}\}_{j=0}^{n}$ using $A^{T}$, which is orthogonal to $\{r\}_{j=0}^{n}$, as follows:

$$
\tilde{r}_{j}=\tilde{r}_{j-1}-\alpha_{j} A^{T} \tilde{p}_{j}
$$

where

$$
\tilde{p}_{j}=\tilde{r}_{j-1}+\beta_{j-1} \tilde{p}_{j-1}
$$

The biorthogonality requirements between $r_{j}$ with $\tilde{r}_{j}$ and $p_{j}$ with $\tilde{p}_{j}$ (with respect to the $A$ inner product) are enforced by choosing

$$
\alpha_{j}=\frac{\tilde{r}_{j-1}^{T} r_{j-1}}{\tilde{p}_{j}^{T} A \tilde{p}_{j}}, \text { and } \beta_{j}=\frac{\tilde{r}_{j}^{T} r_{j}}{\tilde{r}_{j-1}^{T} r_{j-1}}
$$

### 2.1 Quasi-Minimal Residual (QMR)

BiCG can be erratic in practice, increasing in residual norm for several iterations. QMR is designed to smooth out this problem, and make progress or at worst stall when BiCG temporarily diverges. This algorithm was initially developed for complex symmetric linear systems [13], then later adapted to nonsymmetric systems [14].

Whereas GMRES constructs and solves an upper Hessenberg matrix consisting of an orthogonal Krylov subspace, the biorthogonality property of BiCG yields a tridiagonal matrix. Solving it in a least squares sense provides a quasi-minimization of the residual, which can overcome the instability that often occurs in BiCG , allowing for smoother convergence, while maintaining three term recurrences.

Further research into this algorithm has resulted in a number of improvements. A two term recurrence version has been developed [15]. Furthermore, van der Vorst has developed a relatively inexpensive recurrence relation for the computation of the residual vector, as well as a reduction in the number of preconditioning steps (from three to two) [2].

Note that as we have implemented it, QMR may break down ${ }^{4}$.

### 2.2 Conjugate Gradient Squared (CGS)

The goal of QMR is to further reduce the residual when the BiCG iteration stalls. In the case of convergence for BiCG , both $\left\|r_{j}\right\|$ and $\left\|\tilde{r}_{j}\right\|$ converge to zero, yet only the convergence of $r_{j}$ is exploited. Sonneveld showed [21] that by concentrating the effort on the $r_{j}$, the speed of BiCG convergence could be doubled.

If we write $r_{j}=P_{j}(A) r_{0}$ and $\tilde{r}_{j}=P_{j}\left(A^{T}\right) \tilde{r}_{0}$, we see that

$$
\left(r_{j}, \tilde{r}_{i}\right)=\left(P_{j}(A) r_{0}, P_{i}\left(A^{T}\right) \tilde{r}_{0}=\left(P_{i}(A) P_{j}(A) r_{0}, \tilde{r}_{0}\right)=0\right.
$$

[^2]for $i<j$. This implies that we could construct $\tilde{r}_{j}=P^{2}(A) r_{0}$. This is the basis for the Conjugate Gradient Squared Method (CGS). Note that the savings is not only that the $\tilde{r}$ 's are not formed, but we also do not require the transpose of matrix $A$. The result is that the Krylov subspace is built up twice as fast as BiCG, theoretically doubling the speed of convergence. Because of the "squaring" of the polynomial, when the BiCG iterate makes progress towards the solution, CGS doubles that progress. However, when the BiCG iterate turns away from the solution, that error is also doubled. This explains the erratic behavior of the residual norm.

### 2.3 Biconjugate Gradient Stabilized (BiCGSTAB)

Van der Vorst proposed [22] that instead of building the basis vectors for the $i-t h$ dimensional Krylov subspace $\mathcal{K}^{i}\left(\tilde{r}_{0}, A^{T}\right)$ using the same polynomial, i.e. $P_{i}(A)$, as does CGS, the residual could be smoothed using a different polynomial. He ruled out using Chebyshev polynomials since the optimal parameters were not easily obtainable. Instead, he selected a polynomial of the form $Q_{i}(A)=$ $\left(1-\omega_{1} A\right)\left(1-\omega_{2} A\right) \cdots\left(1-\omega_{i} A\right)$, which gives an easy recurrence relation for generating the $Q_{i}$ sequence by updates. The choice of $\omega$ would be such that $r_{n}=Q_{i}(A) P_{i}(A)$ is minimized. Experiments show that this often smoothes the peaks common to the residual norm in CGS, while maintaining the speed of convergence. Note that finite termination is maintained by the orthogonality property $\left(P_{j}(A) r_{0}, Q_{i}\left(A^{T}\right) \tilde{r}_{0}\right)=0$, for $i<j$.

## 3 The Algorithmic Bombardment Algorithm

As indicated earlier, none of the algorithms above are guaranteed to find the solution. They can diverge, stall out, or break down. Thus, we are led to the idea of using all algorithms simultaneously, on the same problem. As soon as one method has converged we stop the overall iteration; if a method breaks down we drop it from the iterative scheme. The resulting poly-iterative algorithm takes more time to converge than the best method, but it has an improved chance of finding the solution.

### 3.1 Parallel implementation

The poly-iteration requires the sum of the floating point operations of the included algorithms, but in a context of message-passing parallel computers, we can increase the efficiency of the approach.

The algorithms we consider are all based on some form of the conjugate gradient method, and thereby they have a very similar structure: they begin by computing an inner product, followed by vector updates, then a preconditioner solve, etc. The inner products, matrix-vector products, and preconditioner solves all require a communication stage. We make the poly-iterative method more efficient by aligning these methods at these operations and combining the communication stages.

Figure 1 illustrates the poly-iterative idea. The operations listed in the circles take advantage of combined communication. A listing of the other operations each algorithm performs in parallel (left to right: CGS, BiCGSTAB, QMR) is also provided.

### 3.2 Structure of the iteration

The global structure of an iteration of the poly-iterative method is as follows:

- In each parallel region, that is, a part of the algorithm where there is no communication, let each processor perform in sequence the operations of the individual methods on its part of the data.


Figure 1: Sequence of Operations
This figure illustrates the sequence of mathematical operations as performed by our implementation of algorithmic bombardment. The operations in the circles combine the communication required of all three methods into one message. The operations in the rectangles have all processors working in parallel (left to right: CGS, BiCGSTAB, and QMR).

- At the start of a communication stage, pack the data of all methods that is to be transmitted in one buffer ${ }^{5}$, then send this buffer in total.

Combining the communications amortizes the communication overhead over the methods. In the case of inner products where just a single floating point number per method is sent, this effectively divides the communication cost by the number of methods.

### 3.3 Cost Model

Obviously this approach requires the combined floating point operations and workspace of each method ${ }^{6}$. This limits the size of the linear system that may be solved, although the actual impact is a function of the sparsity of the matrix. Table 1 lists the the number of vector-sized operations and the amount of vector storage for each method, as well as for bombardment (excluding preconditioning). We ignore scalar storage and operations.

Table 1: Summary of operations and workspace requirements Summary of Operations for Iteration i. " $1 / 1$ " means iteration requires both a matrix times vector and matrix transpose times vector operation. The operations counts for algorithmic bombardment reflect the combining of communication. For example, when we are able to combine the computation of inner products, we count this as one inner product.

| Method | Amount of Work/Iteration |  |  |  | Storage |
| :--- | :---: | :---: | :---: | :---: | :---: |
|  | $\alpha \leftarrow x^{T} y$ | $y \leftarrow \alpha x+y$ | $y \leftarrow A x$ | $x \leftarrow M^{-1} y$ | Requirements |
| CGS | 3 | 6 | 2 | 2 | matrix $+8 n$ |
| Bi-CGSTAB | 6 | 6 | 2 | 2 | matrix $+8 n$ |
| QMR | 4 | 5 | $1 / 1$ | $1 / 1$ | $2 *$ matrix $+14 n$ |
| Bombardment | 3 | 17 | 2 | 2 | $2 *$ matrix $+28 n$ |

The scalar cost of an iteration of the poly-iterative method equals the sum of the costs of the individual methods. In the case where one method is more expensive than the others and this method is not the first to converge, we incur a relatively high cost. On the other hand, poly-iteration is more practical, and possibly much cheaper, than trying different methods in sequence, since this requires a decision when a method is to be abandoned as not likely to converge.

In addition to the storage of matrix $A$, bombardment requires 26 workspace vectors of length $n$. GMRES with restart parameter $m$ uses $(m+5) n=5 n+m n$, so the amount of workspace is equal when the restart parameter is 21 . The problem is that restarting voids the guaranteed convergence property of GMRES.

Another consideration is the amount of work per iteration. GMRES performs one matrix-vector product and one preconditioner solve per iteration compared to two each for each algorithm in bombardment. However, the number of inner products per iteration for GMRES grows linearly with the restart parameter, whereas bombardment requires three (in terms of communication). While it is true that it is possible to compute the GMRES inner products independently, this is known to cause a loss of stability [5].

Parallel architectures require global communication, and this remains the over-riding factor in the performance of these algorithms. For example, the computation of an inner product is an order $n$ operation, but each processor requires the global result. This requires the communication of a single scalar. Each processor computes and sends its local result to all other processors, and receives

[^3]the partial sums from all other processor. Our method computes three inner products locally, then packs them into one message for the same communication requirement as the single case. Our implementation performs three combined inner products per iteration (one is a Euclidean norm). Also, the two consecutive inner products in BiCGSTAB take advantage of combined communication, as they could be in an individual implementation.

To perform the matrix-vector product $A x$, we first collect the global multiplier vector $x$ on each processor ${ }^{7}$, then the resulting local product stays on that processor. This means that we combine the communication here by packing the three multiplier vectors $x_{1}, x_{2}$, and $x_{3}$ into one buffer $x$ which is broadcast to all participating processors. When the transpose of the matrix is explicitly stored, this is the same procedure for performing $A^{T} x$.

When the transpose is not stored, we can still combine communication as follows. First perform the local matrix-vector product $x^{T} A$. This results in a partial sum of the global product. Each processor needs the partial sums of the rows it is responsible for from each processor, so this is packed in the above buffer for collecting the global multiplier.

That is,

1. $x^{T} A$ is performed in parallel,
2. the buffer is packed and broadcast, then
3. the local matrix-vector products $A x$ are performed.

By combining these operations where possible, the bombardment scheme requires eight communications per iteration, compared with five for CGS, seven for BiCGSTAB, and six for QMR (ignoring preconditioning). The savings involved in the preconditioning step is a function of the structure of the preconditioner. For example, if we apply diagonal scaling or, more general, a block Jacobi method, no global communication is required, so there is no savings. However, if an incomplete factorization is used, a relative savings will occur, depending on the requirements of the solve.

## 4 Some Numerical Results

In this section we present some examples as justification for the bombardment approach. For comparison purposes, we define the best algorithm as the one that computes the solution in the least amount of elapsed time.

### 4.1 Implementation Details

- Software
- All codes were written in ANSI standard Fortran 77.
- The Distributed Iterative Linear System Solvers [10] research software was adapted to the bombardment algorithm.
- Also, we have adapted the PIM package [4]. In addition to writing the bombardment algorithm, we changed the communication interface to the BLACS [7]. This allows for portability of the code among the various platforms, while giving optimized communication patterns (especially useful for the global sums required by the inner products), at a negligible cost due to the added programming layer [8].
- Hardware

[^4]- Executed on an Intel iPSC860 Gamma Hypercube [9] at Oak Ridge National Laboratory (ORNL).
- Virtual parallel machines were formed using Sun SPARCstation IPX workstations using PVM [16] over ethernet.

For stopping criteria we use a tolerance TOL $<\left\|r_{k}\right\| /\|b\|$. Since we use the initial guess $x_{0}=0$, this is equivalent to TOL $<\left\|r_{k}\right\| /\left\|r_{0}\right\|$, i.e. we require that the initial residual is sufficiently reduced. We note that this is not necessarily the optimal stopping criteria since the actual accuracy of the reported solution is dependent upon the relationship between the norms of the matrix, the right-hand-side and the true solution. However, for the examples we offer here, this is a reasonable choice. For an overview of stopping criteria, see [2]. For the right hand side we use the unit vector $u=[1, \ldots, 1]^{T}$.

### 4.2 Distributed Memory Parallel Processing Experiments

In a distributed memory parallel processing environment, we can combine the communication of the three algorithms required for the matrix-vector products, preconditioner solvers, and inner products. The actual time savings depends on the structure of the matrix and preconditioner, and the resulting efficiency of the matrix-vector multiplier and preconditioner solver, as well as the latencies involved with message passing. The following experiments were run on the Intel iPSC860 multiprocessor machine at Oak Ridge National Laboratory [9] and clusters of workstations which communicate over ethernet using PVM. The overhead and latency of other machines, as well as floating point performance, will affect these results. Note that time, unless otherwise noted, refers to wall clock time.

## Example 1: Random sparse matrix

Our first example involves a matrix with random sparsity so that an efficient matrix-vector product cannot be designed. We have run the algorithm for a fixed number of iterations, during which there was no convergence of breakdown, and compared the times for each method individually and the time for the poly-iterative method. The respective times per iteration for CGS, BiCGSTAB, and QMR are $0.0274,0.0276$, and 0.0282 seconds. Bombardment took 0.0298 seconds per iteration, only $8.8 \%$ longer than CGS, $8.0 \%$ longer than BiCGSTAB, and $5.7 \%$ longer than QMR. These timings in some sense may be interpreted as the best case for bombardment since each processor must communicate with all the others, and the messages sent during the matrix-vector products are as long as they would ever be. Subsequent examples involve well-structured matrices so that the matrix-vector product can be optimized in order to minimize communication.

## Example 2: The Poisson Problem

Mathematicians have spent, and are spending, a great deal of time trying to identify the properties for which a particular method is optimal. For example

- CGS tends to quickly diverge when the initial guess is close to the exact solution. Therefore, this method should probably be avoided when solving time-dependent problems.
- BiCGSTAB tends to break down when the imaginary parts of the eigenvalues are large relative to the real parts.
- QMR is designed to avoid the breakdown situations that may arise with CGS and BiCGSTAB, but we have found that it is prone to stall.

Yet mysteries still remain, and careful analysis of the coefficient matrix may or may not provide clues as to which method to use. Additionally, even small perturbations may change these properties so that the method that worked well before no longer works at all. And even with this analysis, rounding errors may alter our prediction.

We illustrate this problem using the 2-D Poisson problem. In its basic form, the resulting symmetric positive definite matrix is easily solved by all three methods. Yet if we perturb the basic PDE, so that symmetry or definiteness is altered, a method that previously worked well may break down, stall, or diverge. Mathematical reasons could probably be found to explain this behavior, but when a user just wants the solution, the extra time and workspace needed by algorithmic bombardment may be justified. Below are some experiments run on distributed memory parallel machines as well as networks of work stations. They involve perturbations of the 2-D Poisson equation, solved using central differences on square grids. The goal of these experiments is to illustrate two things:

1. the difficulty in selecting the best algorithm, and
2. the use of the bombardment scheme is not much more expensive than using an individual routine.

We first consider the effects of the problem size on elapsed time. Suppose we wish to solve ${ }^{8}$

$$
\begin{equation*}
-\epsilon\left(\frac{\partial^{2} u}{\partial x^{2}}+\frac{\partial^{2} u}{\partial y^{2}}\right)+\cos (\alpha) \frac{\partial u}{\partial x}+\sin (\alpha) \frac{\partial u}{\partial y}=0 \tag{2}
\end{equation*}
$$

with $\epsilon=-1 / 10, \alpha=-\pi / 6$, on square grids ranging from dimension 100 (order 10,000 matrix) to 400 (order 160,000 matrix) on eight processors of the following parallel machines:

- The Intel iPSC860 (60 Mflop/s per node ${ }^{9}$ ) and
- SUN SPARCstation IPX workstations using PVM over ethernet.

As expected, as the size of the problem increases (and thus the number of floating point operations increases), the difference between executing the best algorithm (BiCGSTAB) and the bombardment algorithm increases (see Figures 2 and 3).

Because QMR takes many more iterations to converge than BiCGSTAB (see Table 2), the time to solution for QMR is greater than the time to find the solution using bombardment. This difference is of course more pronounced for the PVM implementation.

Again, the best algorithm is the one that gives us an accurate solution in the shortest amount of time, regardless of the number of iterations performed. This means the best algorithm could change based on the computing environment. For example, CGS takes more iterations to find the solution for these examples than does BiCGSTAB, and at first glance it appears that these two algorithms require about the same amount of work to perform an iteration. But BiCGSTAB requires an extra global communication step to accomplish the two extra inner products per iteration it must perform. In the Intel environment, where communication latencies are not high, BiCGSTAB is the fastest algorithm. However when the individual nodes are connected via ethernet, as is the case with the PVM experiments, the extra communication becomes significant. The gap closes, and in fact CGS converges faster for some matrix sizes. Although this result may be attributed to other network traffic, it is the nature of ethernet message passing. The time spent in communication provides the insight into why this is happening. Figures 4 and 5 show the proportion of the time to solution spent in message passing as opposed to floating point computation. As expected, the gap is a function of the interconnection network. As expected, the floating point operation requirements increase as the problem size increases, although the startup time to send a message remains constant.

[^5]

Figure 2: Times to solution on the Intel i860
Using 8 processors of the Intel iPSC860, with no preconditioning, we apply the individual algorithms and the bombardment algorithm to $-\frac{1}{10}\left(\frac{\partial^{2} u}{\partial x^{2}}+\frac{\partial^{2} u}{\partial y^{2}}\right)+\cos \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial x}+\sin \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial y}=0$, discretized on a square grid.


Figure 3: Times to solution on Sun SPARCstation IPX workstations
Using a parallel machine consisting of 8 SPARC IPX workstations connected with ethernet using PVM, with no preconditioning, we apply the individual algorithms and the bombardment algorithm to $-\frac{1}{10}\left(\frac{\partial^{2} u}{\partial x^{2}}+\frac{\partial^{2} u}{\partial y^{2}}\right)+\cos \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial x}+\sin \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial y}=0$, discretized on a square grid.


Figure 4: Percentage communication time on the Intel i860
Using 8 processors of the Intel iPSC860, with no preconditioning, we apply the individual algorithms and the bombardment algorithm to $-\frac{1}{10}\left(\frac{\partial^{2} u}{\partial x^{2}}+\frac{\partial^{2} u}{\partial y^{2}}\right)+\cos \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial x}+\sin \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial y}=0$, discretized on a square grid.


Figure 5: Percentage communication time on SPARC IPX workstations
Using a parallel machine consisting of 8 SPARC IPX workstations connected with ethernet using PVM, with no preconditioning we apply the individual algorithms and the bombardment algorithm to $-\frac{1}{10}\left(\frac{\partial^{2} u}{\partial x^{2}}+\frac{\partial^{2} u}{\partial y^{2}}\right)+\cos \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial x}+\sin \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial y}=0$, discretized on a square grid.

Table 2: Number of Iterations to Solution
This table lists the number of iterations required to find the solution to $-\frac{1}{10}\left(\frac{\partial^{2} u}{\partial x^{2}}+\frac{\partial^{2} u}{\partial y^{2}}\right)+$ $\cos \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial x}+\sin \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial y}=0$, discretized on a square grids (the order of the resulting matrix is the square of the grid size). The first line for a grid size is from 8 processors of the iPSC860 and the second line is from 8 Sparc IPX workstations connected with ethernet using PVM. "*" denotes a failure to converge. (However, for grid size 295, CGS converges in 970 iterations and for grid size 310, it converges after 1081 iterations.)

| Grid | CGS | BiCGSTAB | QMR |
| :---: | :---: | :---: | :---: |
| 100 | 181 | 145 | 296 |
|  | 181 | 156 | 296 |
| 150 | 356 | 205 | 459 |
|  | 271 | 220 | 479 |
| 200 | 427 | 289 | 581 |
|  | 480 | 269 | 587 |
| 250 | 1034 | 361 | 765 |
|  | 1157 | 377 | 765 |
| 300 | $*$ | 532 | 1164 |
|  | 658 | 423 | 911 |
| 350 | 1529 | 557 | 1142 |
|  | 1589 | 522 | 1287 |

We particularly note the difference in required iterations on the different machines (see Table 2). This is due to the way the arithmetic is performed by the floating point unit. The SPARCstation IPX uses IEEE arithmetic while the $i 860$ does not. The $i 860$ chip is designed to produce more accurate computations, but since these iterative solvers are not self-correcting, any inexact arithmetic alters convergence patterns, and more accurate does not necessarily correlate with fast convergence. In fact, experiments have shown that an algorithm may converge on one machine yet fails to converge on another [3]. This is illustrated here. For a grid size of 300, the IPX finds the solution, while the iPSC does not. (However, the iPSC does converge for grid sizes slightly smaller and slightly larger than 300.)

These experiments involved only 8 processors of the Intel machine so that results could be compared with a network of workstations. It is of interest, however, to see how our implementation performs on much larger problems, so we performed this experiment using 128 processors of the Intel iPSC860.

Again, we will apply bombardment to Equation (2) on square grids, ranging from dimension 400 (order 160,000 matrix) to 1500 (order $2,250,000$ matrix). The time to solution of the bombardment algorithm as well as the individual algorithms are shown in Figure 6.

We see that there are no surprises with respect to time to the solution for the winning algorithm (in this case BiCGSTAB) and bombardment. As the problem size increases, BiCGSTAB remains about twice as fast as bombardment. Again, the time spent in communication provides the insight into why this is happening. Figure 7 shows the proportion of the time to solution spent in message passing as opposed to floating point operations.

Notice the effects of the grid size upon convergence of CGS, which fails for these finer meshes. Looking back at the coarser meshes in the 8 processor experiments, this is not completely unexpected. For comparison purposes, we iterated for 2500,5000 , and 10000 iterations ${ }^{10}$ for grid sizes of $500 \times$ $500,1000 \times 1000$, and $1500 \times 1500$, respectively.

[^6]

Figure 6: Times to solution on the Intel 1860
Using 128 processors of the Intel iPSC860, with no preconditioning, we apply the individual algorithms and the bombardment algorithm to $-\frac{1}{10}\left(\frac{\partial^{2} u}{\partial x^{2}}+\frac{\partial^{2} u}{\partial y^{2}}\right)+\cos \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial x}+\sin \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial y}=0$, discretized on square grids ranging from dimension 400 to 1500 .


Figure 7: Percentage communication time on the Intel i860
Using 128 processors of the Intel iPSC860, with no preconditioning, we apply the individual algorithms and the bombardment algorithm to $-\frac{1}{10}\left(\frac{\partial^{2} u}{\partial x^{2}}+\frac{\partial^{2} u}{\partial y^{2}}\right)+\cos \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial x}+\sin \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial y}=0$, discretized on square grids ranging from dimension 400 to 1500 .

Next, we present some experiments in which bombardment finds the solution but one or more of the included algorithms fail.

## Example 3: BiCGSTAB is preferable.

Solving Equation (2) on a $200 \times 200$ grid ( 40,000 variables) and no preconditioning ${ }^{11}$, we set $\epsilon=1 / 100$ and $\alpha=-\pi / 6$. BiCGSTAB converges while neither CGS or QMR converge. (See Table 3 for timings, and residual norm histories in Figure 8.)

Table 3: Performance on Intel iPSC/860
Time (in seconds) to solution for solving perturbations of the Poisson equation. * means convergence not achieved.

| Example | CGS | BiCGSTAB | QMR | Bombardment |
| :---: | :---: | :---: | :---: | :---: |
| 1 | 1.37 e 2 | 1.38 e 2 | 1.41 e 2 | 1.49 e 2 |
| 3 | $*$ | 4.78 e 1 | $*$ | 1.26 e 2 |
| 4 | $*$ | $*$ | 7.98 e 0 | 1.32 e 1 |
| 5 | 4.01 e 2 | 5.02 e 2 | $*$ | 8.96 e 2 |



Figure 8: Parallel example: BiCGSTAB wins
Example 6: The residual norm history of each algorithm, using D-ILU preconditioning, applied to $-\frac{1}{100}\left(\frac{\partial^{2} u}{\partial x^{2}}+\frac{\partial^{2} u}{\partial y^{2}}\right)+\cos \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial x}+\sin \left(\frac{-\pi}{6}\right) \frac{\partial u}{\partial y}=0$, discretized on a 200 grid. ( 8 processors of Intel i860 gamma at ORNL.)

The next two examples perturb

[^7]\[

$$
\begin{equation*}
\frac{\partial^{2} u}{\partial x^{2}}+\frac{\partial^{2} u}{\partial y^{2}}+\alpha\left(\frac{\partial u}{\partial x}+\frac{\partial u}{\partial y}\right)+\beta u=1 \tag{3}
\end{equation*}
$$

\]

## Example 4: QMR is preferable.

On a $100 \times 100$ grid ( 10,000 variables) and using no preconditioning, we perturb $\beta(\alpha=0)$. QMR converges, yet CGS and BiCGSTAB fail to converge. (See Table 3 for timings.)

## Example 5: CGS is preferable.

Solving Equation (3) on a $400 \times 400$ grid (160,000 variables) and using block ILU preconditioning ${ }^{12}[1]$, we perturb $\beta(\alpha=0)$. BiCGSTAB converges, but CGS converges faster. QMR fails to determine the solution (even after 5000 iterations). (See Table 3 for timings.)

## 5 Sharing Information Between Algorithms

Since the individual algorithms are iterating in lock-step, it is tempting to share the information from the methods that appears to be working best with the others. That is, if for example CGS is closer to the solution that BiCGSTAB and QMR, why not restart them using the current iterate and residual from CGS? This would have the effect of projecting the iterate of one algorithm onto the Krylov subspace of another. Algorithmic Bombardment facilitaties the sharing of information between algorithms. Sample experiments are discussed below ${ }^{13}$.

## Example 6

A matrix provided by van der Vorst [23], which results from a 5 -point discretization, is easily solved by each method, fastest by BiCGSTAB (Figure 9). After 20 iterations, BiCGSTAB has the better solution approximation of the three, so we share its information with CGS and QMR. Also, every 20 iterations thereafter we share information from the algorithm with the best approximation. BiCGSTAB continues to have the best approximation, and therefore converges as before. However, CGS and QMR are irreparably damaged (figure 10). In fact, immediately after receiving the new information, CGS diverges quickly, then attempts to recover, but then is fed new information, interfering again. QMR merely stalls. Since CGS attempts to converge after temporarily diverging, we try giving it information only once (after 20 iterations). We see that CGS and QMR are still irreparably damaged (figure 11). In fact, this shows that by continuing to give CGS better information, we hid the fact that it's convergence pattern was destroyed. So information from BiCGSTAB is of no value to CGS and QMR. Next we try sharing information from QMR.

## Example 7

The nonsymmetric linear system comes from the discretization (on a $12 \times 12 \times 12$ grid) of the three-dimensional elliptic partial differential equation
$\left(e^{x y z} u_{x}\right)_{x}+\left(e^{-x y z} u_{y}\right)_{y}+\left(e^{-x y z} u_{z}\right)_{z}-250(x+y+z) u_{x}-250[(x+y+z) u]_{x}-(1+x+y+z)^{-1} u=F$
on the unit cube. (The exact solution is $u=e^{x y z} \sin (\pi x) \sin (\pi y) \sin (\pi z)$.) This problem has positive definite symmetric part $\left(A+A^{T}\right) / 2$ for any mesh size [11].

[^8]

Figure 9: All methods converge.
All methods easily find the solution even without preconditioning. This example will be used to show the effects of sharing information from algorithm with the current closest approximation.

Using block ILU, none of individual algorithms can find the solution. Yet with no preconditioning, QMR finds the solution in 382 iterations, while CGS and BiCGSTAB wander about, making no significant progress towards the solution. So every 20 iterations we decide to share information.

Again, the immediate effect is for the residuals of CGS and BiCGSTAB to diverge, then turn and try to converge. But since neither gets below QMR's residual, they are again restarted, and the pattern repeats. So we try restarting only once.

The effects are similar to the experiment where no information is exchanged, except that the CGS residual never dips below that of QMR as before. Hence information from QMR is of no value to CGS and BiCGSTAB. Next we try sharing information from CGS.

## Example 8

In the above example, the CGS residual makes a sharp dip below the QMR residual (around 88 iteration), temporarily giving it the best approximation. Here we give BiCGSTAB and QMR this information. BiCGSTAB gets more erratic, while QMR stalls. Then as before, the CGS residual jumps above the QMR residual, so that QMR has the best approximation, but as we saw above, this doesn't help CGS or BiCGSTAB. Since QMR stalls, this situation doesn't change.

There are two ways to understand these effects of sharing information. First, we could examine the effects on the various parameters of the algorithms. The problem is most easily seen in CGS with the computation of $\beta=\rho_{i-1} / \rho_{i-2}=\tilde{r}^{T} r_{i-1} / \tilde{r}^{T} r_{i-2}$. In the step immediately following the restart, $\beta$ will be smaller than it would have been without a restart. This causes a smaller than expected change in $p$ and $q$, carrying down to a smaller updating of the solution and residual. It is during the next step where the big problem occurs. Now the denominator in the computation of $\beta$ is smaller than expected, while the numerator is about the same size as it was during the previous step, causing $\beta$ to become too large. This cascades down to the approximation, where the updating


Figure 10: Sharing information from BiCGSTAB
Since BiCGSTAB is closest to the solution after 20 iterations, we give $x^{B i C G S T A B}$ and $r^{B i C G S T A B}$ to CGS and QMR. Then every 20 iterations we continuing sharing information.


Figure 11: Sharing information from BiCGSTAB (once)
Since BiCGSTAB is closest to the solution after 20 iterations, we give $x^{B i C G S T A B}$ and $r^{B i C G S T A B}$ to CGS and QMR. But we share information only once.
overshoots the solution. Since the choice of an initial guess doesn't matter, we might expect the algorithm to settle down and begin converging again. But these algorithms use information from all previous search directions, so the root cause of the problem is that we have interfered with that process, contaminating all previous work. Each algorithm builds up a different Krylov subspace in an attempt to find the solution, and while it is true that each algorithm operates on the same matrix, they do so in different ways.

This illustrates two important characteristics of these algorithms:

1. The initial guess doesn't matter (except with CGS, which is likely to diverge if $x_{0}$ is too close the true solution, and
2. each method must build up, and remain in, its own Krylov subspace, based upon the algorithm and the spectrum of the matrix.

A more thorough examination of appropriate sharing among the algorithms constitutes future research.

## 6 Conclusions

Many algorithms have been developed for solving large sparse nonsymmetric linear systems which use short recurrences, but their convergence is no longer guaranteed, nor predictable in practice. Therefore we have incorporated three of these algorithms into a poly-iterative scheme, so that we may apply them simultaneously to the same data set. We have shown through various experiments that this increases the chance of finding the solution, and in a parallel environment this does not increase the time to solution threefold. In fact, even when all three algorithms would have found the solution, bombardment may be faster than the slowest of the three.

The expected performance of a given application is dependent upon the combination of the structure of the matrix (sparsity, structure, etc.), the data structure used, the preconditioner, and these effects upon the performance of the matrix-vector product and preconditioner solver. For example, if the matrix is well-structured, a matrix-vector product can usually be implemented that requires a small amount of communication. Also, if the matrix has a large number of nonzeros, it may be possible to reduce the effects of the indirect addressing of the matrix-vector product.

Ultimately, the performance of the computing environment determines the performance. The new Cray T3D is expected to have much lower communication overhead and latency than the Intel iPSC/860. On the other hand, workstation clusters connected using PVM [16] exhibit high latency, and are dependent upon the traffic interconnection network, often the Internet. Regardless, the increased probability of convergence should justify using poly-iteration.

## 7 Future Work

The experiments presented above are frequently encountered in the scientific world, hence we believe the results justify our implementation of the poly-iterative idea, including the choice of algorithms as well as the scheme for performing the matrix-vector product and preconditioning. However, different approaches may be more appropriate depending upon the problem being solved. For example, some applications require solving many linear systems in a sequence of time steps. Since the matrix may not change significantly from one step to the next, it has been suggested that perhaps bombardment could be used during one such solve, then only the winning algorithm would be used for the next few solves, then back to bombardment, and so on.

Perhaps incorporating more GMRES concepts into the poly-iteration would be valuable in some cases. Since we originally ruled out using this valuable algorithm due to its linearly increasing
workspace requirements, we must find a way to overcome this limitation while still gaining performance. BiCGSTAB is actually the combination of BiCG and GMRES(1). Recent work [23] shows that increasing the effects of GMRES can be worthwhile, such as combining GMRES(2) or GMRES(4) with BiCG.

Additionally, different preconditioning schemes might prove valuable under some circumstances. Here we use the same preconditioner for each algorithm. Perhaps this is not optimal. For example, the performance of QMR was often degraded by ILU.

Further research into different matrix-vector product implementations may yield higher computational performance in some situations. For example, certain matrix structures may allow higher efficiency. One possibility would be to interleave the elements of the multiplier of the algorithms in order to force less indirect addressing, which slows the floating point performance. Dense matrix computations perform $O\left(n^{3}\right)$ operations on $O\left(n^{2}\right)$ data. But for sparse matrices, this is actually a vector-vector operation $(O(n)$ operations on $O(n)$ data), with the added degradation of indirect addressing. And since three such operations must be performed, the effect is magnified. This can be reduced in the bombardment scheme. Suppose the multipliers are $x^{C G S}, x^{B i C G S T A B}$, and $x^{Q M R}$. The obvious way to compute $A x^{C G S}, A x^{B i C G S T A B}$, and $A x^{Q M R}$ is to perform the operations sequentially. But the elements can be interleaved as

$$
x=\left[x_{1}^{C G S}, x_{1}^{B i C G S T A B}, x_{1}^{Q M R}, \ldots, x_{n}^{C G S}, x_{n}^{B i C G S T A B}, x_{n}^{Q M R}\right]^{T}
$$

reducing the effects of indirect addressing threefold. Note that this scheme will cause indirect addressing of some vector updates, so its overall effect is dependent upon the number of nonzeros in the matrix.

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    ${ }^{1}$ Indeed, there are examples in which the convergence of GMRES with any length recurrence less than $n$ will stagnate.
    ${ }^{2}$ It is possible to combine the communication of these inner products (using the unmodified Gram-Schmidt), but this is often unstable.

[^1]:    ${ }^{3}$ Although all are based on the idea of biconjugate gradients, and therefore theoretically converge or diverge together, experiments have shown that one may converge much faster than the others, may break down, or fail to converge.

[^2]:    ${ }^{4}$ A version of QMR that includes a "look-ahead" algorithm can avoid these problems, but for simplicity we do not use it.

[^3]:    ${ }^{5}$ In certain communication schemes such as PVM [16] this buffering is provided, in other schemes such as PICL [17] and the BLACS [7] it has to be implemented as part of the poly-iterative algorithm.
    ${ }^{6}$ less $2 n$ since the right-hand-side vector $b$ need only be stored once.

[^4]:    ${ }^{7}$ Actually, the structure of the matrix determines how much of the global multiplier vector is needed. For example, if the matrix is block tridiagonal, such as arises in five-point discretization methods, only nearest-neighborinformation may be needed.

[^5]:    ${ }^{8}$ This problem was used in Sonneveld's paper presenting CGS [21].
    ${ }^{9}$ Millions of floating point operations per second.

[^6]:    ${ }^{10}$ Actually, we iterated for far more iterations to convince ourselves that convergence would not be achieved.

[^7]:    ${ }^{11}$ The result is similar when D-ILU preconditioning is used.

[^8]:    ${ }^{12}$ Block ILU preconditioning requires no communication.
    ${ }^{13}$ Note that for experimental purposes, we continue until either all algorithms converge or some maximum number of iterations are performed. Of course, when an algorithm finds the solution to tolerance, we drop it entirely from the scheme.

