

# A Scalable Parallel Poisson Solver in Three Dimensions with Infinite-Domain Boundary Conditions

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## Abstract

*We present an elliptic free space solver that offers vastly improved performance over a previous variant of the algorithm. We currently scale up to 1024 processors of an IBM SP system, and we are planning to port the solver to Blue Gene/L. The solver employs a method of local corrections that avoids the need for costly communication, while retaining parallel scalability of the method. Communication costs are generally small: 25 percent of the total running time or less for runs on up to 512 processors and 37 percent of the total time on 1024 processors. The numerical overheads incurred are independent of the number of processors for a wide range of problem sizes. The solver currently handles infinite-domain (free space) boundary conditions, but may be reformulated to accommodate other kinds of boundary conditions as well.*

## 1. Introduction

Elliptic solvers for free space problems often scale poorly owing to the difficulty in treating the infinite-domain boundary conditions. While some of the underlying communication could be masked by overlapping it with useful computation [3, 4, 5, 20, 21], the approach is ultimately non-scalable, as the total cost of communication grows with the size of the problem. We present an alternative approach that algorithmically reformulates the solution, reducing the amount of data communicated at the expense of additional computation. This tradeoff is beneficial. The added computational overheads are purely local work, and the resultant algorithm employs a fixed number of communication and computation steps. We use a local corrections strategy that divides the solution into low and high resolution

components. The low resolution component carries far-field data, thereby reducing the amount of communication required. Our algorithm exploits elliptic regularity, an approach which is also employed by the fast multipole method [14], the method of local corrections for particle methods [2], the finite element method of Bank and Holst [9], and the two-dimensional method of local corrections for free space problems [6, 8]. Results presented by Holst [16] include a rigorous proof that these types of algorithms can produce accurate results with little communication.

We have previously reported early results with our solver, called Scallop. While that solver enabled us to avoid high communication overheads, computation of the infinite-domain boundary conditions became a bottleneck, and numerical overheads hampered scalability beyond 512 processors of an IBM SP system with POWER3 processors. We discuss algorithmic enhancements in our new version that greatly reduce the computational overheads and diminish the total running time required to reach a solution. Scallop solves elliptic partial differential equations with infinite-domain (free space) boundary conditions. Such boundary conditions are useful for particle in cell calculations [15] and for fast particle methods, particularly those which require the solution of the velocity field [1, 2]. While Scallop can be altered to handle other boundary conditions, we focus here on the infinite-domain case. For purposes of simplification, we restrict the discussion to the Poisson equation.

Unlike domain decomposition methods such as [19] which require multiple iterations between the local and non-local descriptions, Scallop does not perform repeated iterations between coarse and fine levels or several communication steps. Scallop reaches a solution to the Poisson equation in three steps and communicates data only twice. First, coarse grid data are communicated to generate a global coarse grid charge field. Second, coarse and fine boundary

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condition data are communicated once among neighboring regions. These communication costs are low in practice — at most 25 per cent on up to 512 processors — and come at the expense of computational (numerical) overhead. We show that the extra computation involved is reasonable over a wide range of problem sizes. As a result, we are currently able to demonstrate scalability on up to 1024 processors of an IBM SP system, and we plan future computations on thousands of processors.

Our contribution to our prior work with Scallop comes in two parts. First, we greatly improve the speed of our serial infinite-domain solution by using the fast multipole method to calculate the necessary boundary conditions. Second, we now calculate coarse grid values necessary for the method of local corrections simultaneously with the initial local solutions, also using the fast multipole method. By calculating coarse grid values in the correction radius in this way, we are able to scale up to a greater number of processors with much lower computational overhead.

Scallop is representative of a class of algorithms that employ sophisticated numerical techniques to reduce communication costs. The techniques in turn require an appropriate software infrastructure to manage the underlying details, in particular the bookkeeping. To this end, Scallop was prototyped [7] with KeLP [13], a rapid development C++ framework for implementing scientific applications on distributed memory parallel computers. KeLP provides geometric and communication abstractions that facilitated the development of Scallop without sacrificing performance. The Scallop prototype was subsequently ported to the Chombo Structured Adaptive Mesh Refinement Infrastructure [11], which provides many of the same geometric and communications abstractions as KeLP.

## 2. Preliminaries

The equation we solve is the Poisson equation in three dimensions with a charge distribution  $\rho$  with compact support, i.e. the charge is only nonzero in a finite region of space. Specifically, we seek the solution  $\phi$  to

$$\Delta\phi = \frac{\partial^2\phi}{\partial x^2} + \frac{\partial^2\phi}{\partial y^2} + \frac{\partial^2\phi}{\partial z^2} = \rho(x, y, z)$$

which has far-field behavior characterized by

$$\phi = -\frac{R}{4\pi|\vec{x}|} + o\left(\frac{1}{|\vec{x}|}\right), \quad |\vec{x}| \rightarrow \infty,$$

where  $R$  is the total charge:

$$R = \int_{\Omega} \rho(\vec{x}) d\vec{x},$$

and the region  $\Omega$  contains the support of the charge  $\rho$ .

Many applications for which our method would be useful, including particle in cell calculations [15] and low mach number incompressible flow simulations [10], are limited to  $O(h^2)$  accuracy. We likewise seek a solution which is accurate to  $O(h^2)$ .

To simplify our discussion, we will be solving problems on a unit cube, discretized into  $N$  cells ( $N + 1$  points) in each direction. We refer to this discretized computational domain as  $\Omega^h$ , and define  $h = 1/N$ . This computational domain corresponds to the index set of the discrete solution  $\phi^h$ , i.e. the indices of the underlying discrete mesh.

Since our goal is to solve the problem on parallel processors, we partition  $\Omega^h$  into a set of disjoint subdomains  $\Omega_k^h$ :

$$\Omega^h = \bigcup_k \Omega_k^h.$$

Our method entails solving local problems on each of the  $\Omega_k^h$  in parallel, as well as on a single coarsened global mesh  $\Omega^H$ . The spacing of this coarsened mesh is  $H = Ch$ , where  $C$  is a specified *coarsening factor*.

We choose the domain  $\Omega^h$  to be a rectangular region,  $\Omega^h = [\vec{l}, \vec{u}]$ , where  $\vec{l}$  and  $\vec{u}$  are the integer vectors corresponding to the lower and upper corners of the region. The coarsened domain is then defined as

$$\mathcal{C}(\Omega^h, C) = \Omega^H = [[\vec{l}/C], [\vec{u}/C]]$$

where the operators  $[\cdot]$  and  $[\cdot]$  represent the *floor* and *ceiling* operators, respectively.

Because our meshes are node-centered, the points of  $\Omega^H$  map directly onto corresponding points in  $\Omega^h$ , and no averaging is required to coarsen the mesh data. Thus, we can coarsen the mesh by sampling the mesh without having to interpolate. In particular, we coarsen a fine grid representation using the *sample* operator  $\mathcal{S}^H$ : for each point  $\vec{x}_C$ , we can find the coarse grid value  $\psi^H(\vec{x}_C)$  (where  $\psi^H$  has grid spacing  $H$ ) by finding the fine grid point  $\vec{x}$  at the corresponding position in  $\psi^h$  (with grid spacing  $h = H/C$ ):

$$\psi^H(\vec{x}_C) = (\mathcal{S}^H(\psi^h))(\vec{x}/C) = (\psi^h)(\vec{x})$$

For the discussion that follows, we also need one more bit of notation. The *grow* operation extends or shrinks an index domain by a uniform amount in each direction. If  $\Omega^h = [\vec{l}, \vec{u}]$  (where  $\vec{l} = (l_x, l_y, l_z)$  and  $\vec{u} = (u_x, u_y, u_z)$ ), we define *grow* as

$$\text{grow}(\Omega^h, g) = [\vec{l} - (g, g, g), \vec{u} + (g, g, g)].$$

When  $g < 0$ , *grow* returns a shrunken domain.

## 3. The Method

Our domain decomposition method is built upon a method for solving single-processor infinite-domain Pois-

son problems, as described previously in [7]. We will summarize the single-grid algorithm first, and then describe the domain decomposition algorithm.

### 3.1 A Serial Infinite-Domain Poisson Solver

Following the approach described in [17] and [18], we are able to calculate a solution to the Poisson equation with infinite-domain boundary conditions in four steps, using two solution grids. Given a charge on a grid  $\Omega^h$ , these are the *inner grid*  $\Omega^{h,g}$  and the *outer grid*  $\Omega^{h,G}$ , defined as

$$\begin{aligned}\Omega^{h,g} &= \text{grow}(\Omega^h, s_1); \\ \Omega^{h,G} &= \text{grow}(\Omega^{h,g}, s_2).\end{aligned}$$

The four steps required to calculate the solution are as follows.

1. Find the solution to the Poisson equation on the inner grid,  $\Omega^{h,g}$ , using Dirichlet boundary conditions.
2. Calculate a charge,  $q$ , along the inner grid boundary equal to the normal derivative of the solution from step 1 at the inner grid boundary.
3. Calculate boundary conditions at each point on the outer grid boundary by numerically integrating the effect of the charge at the inner grid boundary:

$$g(\vec{x}) = \int_{\partial\Omega^{h,g}} G(\vec{x} - \vec{y})q(\vec{y})dA_{\vec{y}},$$

where  $G$  is the Green's function.

4. Find the solution to the Poisson equation on the outer grid,  $\Omega^{h,G}$ , using the boundary conditions,  $g$ , just calculated.

Our approach here is identical to the approach described previously in [7] except in the way the integration is performed in step 3. Previously, we integrated the charge from the inner grid onto a coarsened version of the outer grid, with mesh spacing  $H = h/O(\sqrt{N})$ , and interpolated from the coarse grid to find necessary values on  $\partial\Omega^{h,G}$ . Our straightforward integration required  $O(N^3)$  work but significantly took more time to compute than the Dirichlet solutions on the inner and outer grids.

In our current implementation, we perform the integration required for the boundary calculation using the fast multipole method (FMM). Each face of  $\Omega^{h,g}$  is divided into patches of  $r \times r$  points. We then calculate the multipole moments of the charge up to order  $M$  on each patch. On each face of  $\Omega^{h,G}$ , for points on a mesh coarsened by  $r$  in each dimension and expanded by a coarse layer of points of width  $P$ , we add up the evaluations of multipole expansions

due to all the faces of  $\Omega^{h,g}$ . Finally, we interpolate polynomially, one dimension at a time from the coarse mesh values to the remaining fine mesh points on the face. (See Figure 1.)

Choosing  $r = \sqrt{N}$  provides sufficient accuracy for the solution and allows the integration step to be completed in  $O((M^2 + P)N^2)$  work. The values of  $M$  and  $P$  are chosen with regard to accuracy requirements and are independent from  $N$ , so for a given degree of accuracy, the integration step requires  $O(N^2)$  work.

We should also note the constraints required on  $s_1$  and  $s_2$ , the spacing between the grids  $\Omega^h$ ,  $\Omega^{h,g}$ , and  $\Omega^{h,G}$ . We have found that setting  $s_1 = 0$  has only small effects on the accuracy of our solutions and doing so allows us to minimize the size of the solution grids. Convergence requirements of the multipole method force us to choose  $s_2$  with more care, however. In order for the multipole expansions from a patch to converge, the distance from a patch center on  $\partial\Omega^{h,g}$  to the points on  $\partial\Omega^{h,G}$ , on which the expansion is evaluated, should be at least twice the radius of the patch. Here we define the radius of a patch as the maximum distance from the patch center to any point on the patch. Recall that we chose our patches to be  $r$  fine grid points square. Thus our patches have a radius of  $rh/\sqrt{2}$ , and the distance requirement becomes  $s_2h \geq 2(rh/\sqrt{2})$ . We also need the number of cells along the length of  $\Omega^{h,G}$  to be divisible by  $r$ . Combining these two requirements, we arrive at the following formula for  $s_2$ :

$$s_2 = \frac{r}{2} \lceil 2\sqrt{2} + \frac{N}{r} \rceil - \frac{N}{r}. \quad (1)$$

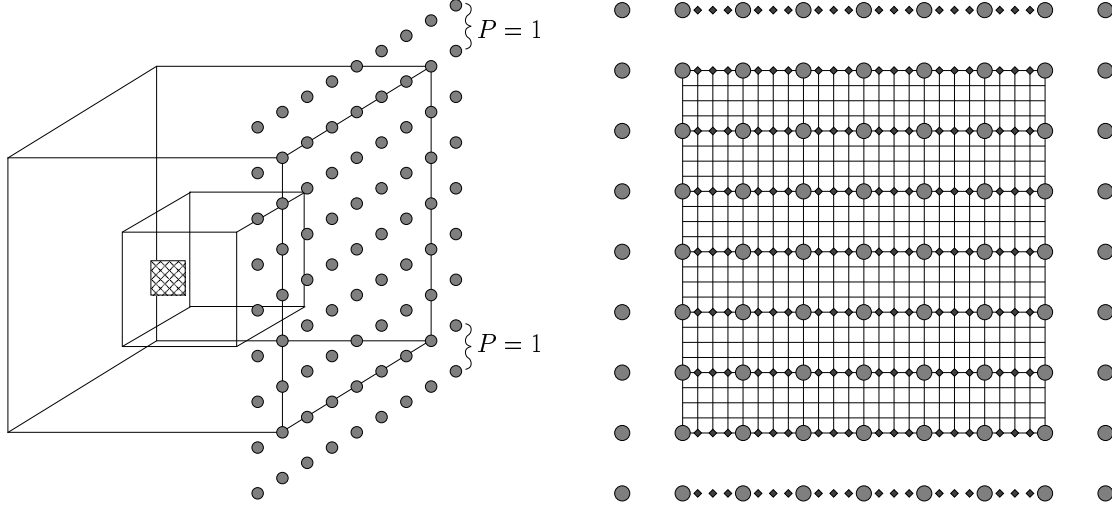
In order to demonstrate the effect of these requirements, we show in Table 1 the necessary values of  $s_2$  for grid sizes,  $N$ , ranging from 16 to 2048 by powers of 2. Values of  $r$  are chosen to be close to the square root of  $N$  but also multiple of four. Note that the ratio of  $N^G$  (the length of  $\Omega^{h,G}$ ) to  $N$  decreases as  $N$  increases. For serial solutions, this implies that overhead will be smaller for larger infinite-domain problems.

### 3.2 Domain Decomposition

The domain decomposition algorithm described here is a finite-difference analogue of Anderson's method of local corrections [2]. Our algorithm consists of three computational steps interspersed by two communication steps, as described previously in [7].

1. INITIAL LOCAL SOLUTION. We calculate a local infinite-domain solution on each local subdomain,  $k$ , augmented with an overlap region:

$$\Delta_{19}\phi_k^{h,initial} = \rho_k^h \text{ on } \text{grow}(\Omega_k^h, s + Cb).$$



**Figure 1.** In step 3, multipole moments are calculated for each patch on  $\partial\Omega^{h,g}$ , such as the one shown cross-hatched on the inner box. The multipole expansions are then evaluated at the coarse points of  $\partial\Omega^{h,G}$ , plus an additional layer of width  $P$ , indicated with gray circles for one face. These evaluations are interpolated to the fine points of  $\partial\Omega^{h,G}$ , located at intersections of the black lines, using two passes. The evaluation points of the first pass are shown as small gray diamonds.

$N$	$r$	$s_2$	$N^G$	$N^G/N$
32	8	12	56	1.75
64	8	12	88	1.38
128	12	20	168	1.31
256	16	24	304	1.19
512	24	44	600	1.17
1024	32	48	1120	1.09
2048	48	80	2208	1.08

**Table 1.** Values of the coarsening factor,  $r$ , annulus thickness,  $s_2$ , and resulting expanded grid size,  $N^G = N + 2s_2$ , for various input grid sizes  $N$ . The ratio of  $N^G/N$  decreases for increasing  $N$ .

and construct a coarsened version of the solution,  $\phi_k^{H,initial}$ , by sampling:

$$\phi_k^{H,initial} = \mathcal{S}^H(\phi_k^{h,initial}) \text{ on } grow(\Omega_k^H, s/C + b).$$

Here  $s$  is a correction radius,  $C$  is the coarsening factor, and  $b$  is the width of a layer for polynomial interpolation to be used in step 3. The  $\Delta_{19}$  operator represents the Laplacian calculated with a 19-point stencil of nearest neighbor points. The error characteristics of the 19-point stencil are essential for maintaining  $O(h^2)$  accuracy in the overall algorithm when combin-

ing the effects of coarse and fine grid data later on. (For further discussion of discrete Laplacian operators, see [12].)

2. GLOBAL COARSE SOLUTION. We couple the individual local solutions by solving another Poisson equation on a coarsened mesh covering the entire domain. We first construct coarsened local charge fields:

$$R_k^H = \begin{cases} \Delta_{19}\phi_k^{H,initial} & \text{on } grow(\Omega_k^H, s/C - 1), \\ 0 & \text{otherwise} \end{cases}$$

and then sum up these charge fields to form a global coarse representation of the charge:

$$R^H = \sum_k R_k^H.$$

Then we solve

$$\Delta_{19}\phi^H = R^H \text{ on } grow(\Omega^H, s/C + b)$$

with infinite-domain boundary conditions.

3. FINAL LOCAL SOLUTION. Using the discrete Laplacian  $\Delta_7$  calculated with a 7-point stencil at nearest neighbor points, we solve

$$\Delta_7\phi_k^h = \rho_k^h \text{ on } \Omega_k^h$$

with Dirichlet boundary conditions on  $\partial\Omega_k^h$ :

$$\phi_k^h(\vec{x}) = \sum_{k': \vec{x} \in grow(\Omega_{k'}^h, s)} \phi_{k'}^h(\vec{x}) + \mathcal{I}(\phi^{H,corr})$$

where  $\mathcal{I}$  is the same interpolation operator used in the serial infinite-domain Poisson solver and

$$\phi^{H,corr} = \phi^H(\vec{x}) - \sum_{k': \vec{x} \in grow(\Omega_{k'}^h, s)} \phi_{k'}^{H,initial}(\vec{x}).$$

Figure 2 depicts the regions from which data are taken to set boundary conditions on a face.

Note that the algorithm does not require fine grid data at all points in  $grow(\Omega_k^h, s + rb)$ . Specifically, the algorithm does not need fine grid data outside of  $grow(\Omega_k^h, s/r + b)$ . In order to complete steps 2 and 3, we only need the following data from step 1:

- the solution at coarse grid values,  $\phi_k^{H,initial}$ , on  $grow(\Omega_k^H, s/C + b)$ , necessary for step 2, and
- the solution at fine grid values,  $\phi_k^{h,initial}$ , on the faces of  $grow(face, s)$  for each face  $face$  of  $\Omega_k^h$  (here  $grow$  is a two-dimensional operator).

To ensure accuracy of the method, we need  $s = 2C$  and  $b = 1$ .

As before, communication is only required in two phases: first, in constructing the global coarse charge field, and second, to set the boundary conditions for the final local solution.

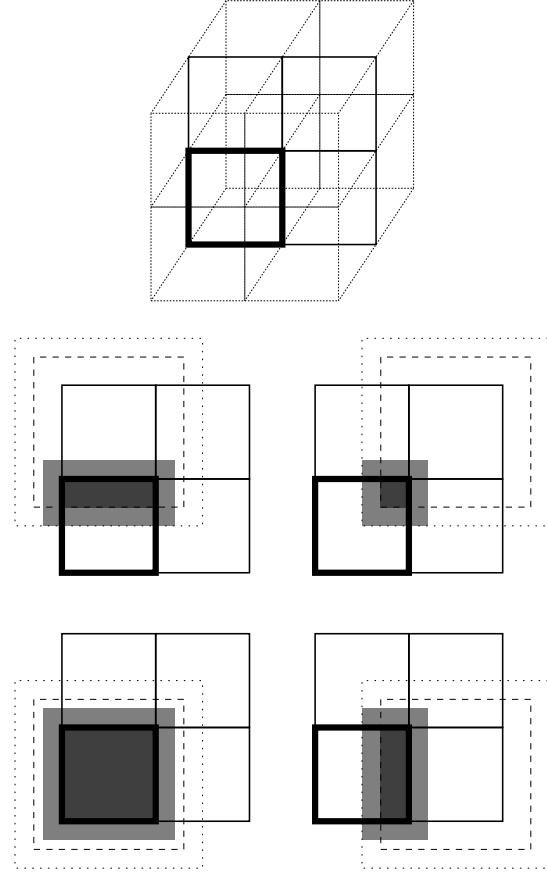
## 4. Performance Model

As mentioned previously, the principle behind Scallop is to trade off communication against computation. We next discuss these tradeoffs and show that they are reasonable. We describe a performance model, and use it to show that in theory the overheads are reasonable. In the following two sections we reconcile our predictions with practice.

In determining the computational overhead in Scallop, we will use the serial infinite-domain Poisson solver as a baseline. We will first show that the cost of our initial fine grid solutions is similar to the cost of a serial solution. The computational overhead in Scallop can be described as the sum of three costs: the extra computation required to calculate solutions on expanded local grids, the cost of the coarse grid solution, and the time required for the final local solutions on fine grid data. We will discuss each of these costs, and show that with the proper choice of a coarsening factor, Scallop should be able to scale to thousands of processors.

### 4.1 Serial Infinite-Domain Poisson Solver

Scallop reuses many of the same components as the serial infinite-domain Poisson solver. We will examine the computational costs involved in the serial solver first and



**Figure 2. Setting boundary values for a final local solution in step 3 of MLC. For the face outlined in bold in the top of the figure, in a layout of eight cubes, the lower diagrams depict the regions from which data are copied from faces of different neighboring boxes. Solid lines indicate the boundaries of the boxes  $\Omega_{k'}^h$ , dashed lines the boundaries of the boxes  $grow(\Omega_{k'}^h, s)$ , and dotted lines the boundaries of the boxes  $grow(\Omega_{k'}^h, s + Cb)$ . Fine-grid data are copied to the bold face from the nodes inside and on the edges of the regions shaded dark gray. Coarse grid data are copied from nodes inside and on the edges of the regions shaded both dark and light gray, and then interpolated to nodes on the bold face that are inside and on the edges of the regions shaded dark gray.**

compare this cost with the cost of the initial solutions in Scallop.

For simplicity, let us consider only cubical domains with edge length  $N$ . The operation counts for each step of the algorithm described in Section 3.1 are as follows.

1. Finding the solution to the Poisson equation on the inner grid using a fast (FFT) Poisson solver:  $O(N^3 \log N)$ .
2. Calculate a charge,  $q$ , along the inner grid boundary:  $O(N^2)$ .
3. Calculate boundary conditions at each point on the outer grid using FMM:  $O(N^2)$
4. Find the solution to the Poisson equation on the outer grid using a fast (FFT) Poisson solver:  $O(N^3 \log N)$ .

Thus the serial infinite-domain solver operation count is bounded by the Dirichlet Poisson solve, and the overall computational cost of an infinite-domain solution is  $O(N^3 \log N)$ .

## 4.2 Practical Work Estimates

Since the computation required by the Poisson solvers used in our algorithm is nearly proportional to the number of points for which a solution is being found (ignoring the weaker  $\log N$  term in the previous work estimates), we propose the following work estimates on the basis of these grid sizes. First, let us define  $W^{Dir}$  as an estimate of the work required for a Poisson solution with Dirichlet boundary conditions on a mesh  $\Omega^h$ :

$$W^{Dir} = \text{size}(\Omega^h),$$

where the *size* operator returns the total number of points in the mesh  $\Omega^h$ . Similarly, let us refer to  $W_k^{Dir}$  as the work required to compute a Dirichlet solution on a subdomain  $\Omega_k^h$ .

Recall that the infinite-domain boundary calculation requires the solution of a Dirichlet problem on an enlarged domain, as defined by equation (1) and shown by example in Table 1. After calculating the extents of  $\Omega^{h,g}$  and  $\Omega^{h,G}$  according to these requirements, we can define a work estimate on an infinite-domain solution as

$$W^{id} = \text{size}(\Omega^{h,g}) + \text{size}(\Omega^{h,G}),$$

and also  $W_k^{id}$  as the corresponding estimate for the work required to compute a local initial fine grid solution on  $\Omega_k^h$  within our MLC method.

Finally, as an estimate of the total work required for our MLC method for a particular processor  $P$ ,

$$W_P^{mlc} = W_{coarse}^{id} + \sum_{k \text{ assigned to } P} (W_k^{id} + W_k^{Dir})$$

where  $W_{coarse}^{id}$  is an estimate of the work required to calculate the infinite-domain solution on the global coarse mesh. Note that to allow for the possibility of overdecomposition, multiple subdomains  $k$  may be assigned to a single processor  $P$ .

## 4.3 Limiting the Cost of the Coarse Grid Solution

In comparing the computational cost of Scallop to a serial infinite-domain solver, the cost of computing the solution on the global coarse grid is overhead. We want to determine what range of problems can be solved with only small computational overhead due to the coarse grid calculation.

As before, let  $N$  be the length of a side, thus  $(N + 1)^3$  is the total number of points. Let  $q$  be the number of subdomains on a side. Then  $q^3$  is the total number of subdomains (the maximum number of processors) and  $N_f = N/q$  is the length of a local fine subdomain.

Let  $C$  be the coarsening factor, as defined previously, such that the size of coarse grid,  $N_c$ , is  $N/C$ . Since the coarse grid solution is not parallelized, we want  $N_c < N_f$  in order to reduce the overhead due to the coarse grid. Thus we have  $N/C < N/q$ , which reduces to  $q < C$ .

## 4.4 Limits of Parallelism for the Method

As in most numerical libraries, an important consideration is how to optimize parameter settings that affect performance. The performance of Scallop is most affected by the choice of two parameters:  $q$  and  $C$ . However, various factors constrain the choice of these parameters, as well as the intrinsic parallelism, and these constraints limit performance.

When choosing the coarsening factor,  $C$ , we may affect both the size of the coarse grid solution and the size of the initial local solutions. Recall that our MLC algorithm requires us to find the initial local solutions on grids expanded by  $2C$  in each direction. Thus increasing  $C$  may lead to extra work for the initial local solutions. As we have just seen in section 4.3, however,  $C$  needs to increase with  $q$  in order to keep the cost of the coarse solution in line with the local solutions.

Since the serial infinite-domain solver requires an annulus itself, there is a range of problems for which our algorithm is most suitable. We would like the coarsening factor for our MLC solver to be less than or equal to half the annulus size required by the infinite domain solver, i.e.  $C \leq s_2/2$ . The coarsening factor must also evenly divide the local grid size  $N_f$ . The maximum number of processors is then dependent on the choice of the ratio between  $q$  and  $C$ .

Table 2 shows the limits of parallelism in terms of the maximum number of processors,  $P$ , and maximum prob-

$q/C$	$N_f$	$s_2$	$q$	$P$	$N^3$
1/2	64	12	2	8	$128^3$
1/2	128	20	4	64	$512^3$
1/2	256	24	4	64	$1024^3$
1/2	512	44	8	512	$4096^3$
1	64	12	4	64	$256^3$
1	128	20	8	512	$1024^3$
1	256	24	8	512	$2048^3$
1	512	44	16	4096	$8192^3$
2	64	12	8	512	$512^3$
2	128	20	16	4096	$2048^3$
2	256	24	16	4096	$4096^3$
2	512	44	32	32768	$16384^3$

**Table 2. Limits of parallelism for our MLC method, showing maximum number of processors,  $P$ , and problem size,  $N^3$ , with fixed  $q/C$  and fixed local problem size  $N_f$ . The maximum value for  $P$  is  $q^3$ , the total number of subdomains.**

lem size  $N^3$  for various local problem sizes,  $N_f$ , and ratios of  $q$  and  $C$ . If we assume, for instance, that  $128^3$  problems will easily fit in local memory, users willing to expend twice the computational effort can reach problem sizes of  $1024^3$  using 512 processors. Users willing to expend eight times the computational effort could reach a problem size of  $2048^3$  on 4096 processors.

## 4.5 Future Improvements

The current implementation is limited by the relationship between the coarsening factor,  $C$ , and the number of subdomains per side,  $q$ . This restriction is due to computing the global coarse grid solution in serial. By parallelizing the global coarse solution, we can vary  $C$  and  $q$  independently and extract significantly more parallelism from our MLC method. Our current version of Scallop includes a parallel implementation of the multipole calculation on the coarse grid infinite-domain solution, and we are considering alternatives for efficiently parallelizing the Dirichlet solves on the coarse grid while keeping communication requirements low. If our efforts are successful, Scallop could efficiently use thousands of processors without incurring additional computational overhead on the local solutions.

In generating the data shown in the following section, we realized that the implementation of the Chombo library function called to handle nearest neighbor boundary communication performs far more communication than necessary. Rather than communicating two-dimensional planes of data, the function sends three-dimensional slabs of thick-

ness  $2C$  from processor to processor. Limiting the communication to only those planes required by the algorithm will reduce the overall amount of communication for the boundary condition phase of the method by a factor of between  $C$  and  $2C$ .

## 5. Results

In this section we present computational results which demonstrate the low communication overhead of Scallop on up to 1024 processors. We also compare our performance results with the estimates presented earlier in Section 4.

### 5.1 Hardware and System Environment

We ran on NERSC’s Seaborg IBM SP system, located at the National Energy Research Scientific Computing Center<sup>1</sup>. Seaborg contains POWER3 SMP High Nodes interconnected with a “Colony” switch. Each node is an 16-way Symmetric Multiprocessor (SMP) based on 375 MHz Power-3 processors<sup>2</sup>, sharing between 16 and 64 Gigabytes of memory, and running AIX version 5.1.

Scallop version 2 is written in a mixture of C++ and Fortran 77. We used the IBM C++ and Fortran 77 compilers, `mpCC` and `mpxlf`. C++ code was compiled with the IBM `mpCC` compiler, using options `-O2 -qarch=pwr3 -qtune=pwr3`. Fortran 77 was compiled with `mpxlf` with `-O2` optimization. We used the standard environment variable settings, and we collected timings in batch mode using `loadleveler`. The timings reported are based on wall-clock times, obtained with `MPI_Wtime()`. Each calculation was performed 3 times. The times reported are for the runs with the shortest total times. Timers were placed around large function calls rather than inner loops to reduce the effects of noise in the timing results. The bulk-synchronous nature of the algorithm allows us to fully separate computation times from communication times. Reported running times do not include a preprocessing phase for the serial Poisson solver that computes a matrix for obtaining outer-grid boundary conditions from multipole coefficients due to charges on the inner-grid boundary. This matrix depends only on the problem size and accuracy parameters, and its computation is considered a fixed overhead to be amortized over many calls to the solver.

### 5.2 Scalability

In order to measure performance, we scaled the work with the number of processors. In order to have grid sizes

<sup>1</sup><http://www.nersc.gov/nusers/resources/SP>

<sup>2</sup><http://www-1.ibm.com/servers/eserver/pseries/hardware/whitepapers/nighthawk.html>

Input Parameters				Times for Each Stage (seconds)						Total (sec)	Proc-time/pt. ( $\mu\text{sec}$ )	Percent Comm.
$P$	$q$	$C$	$N^3$	Local	Red.	Global	GC	Bnd.	Final			
128	8	6	$768^3$	36.92	6.39	15.73	2.55	9.61	4.95	73.83	20.78	25
256	8	8	$1024^3$	48.38	6.74	14.46	1.35	7.41	6.05	84.40	20.06	18
512	8	10	$1280^3$	49.23	4.90	16.35	3.31	7.66	7.63	86.16	20.99	18
1024	16	12	$1536^3$	49.07	6.44	15.99	2.88	30.72	4.96	107.33	30.27	37

**Table 3. Input parameters and timing breakdowns for runs. The Local and Global solutions require an infinite-domain solution, whereas the Final calculation solves a simpler Dirichlet problem. The time for Reduction (Red.) includes everything necessary to accumulate the coarsened local solutions into a single coarse grid for the Global solution. GC is the communication time within the Global solution. The time for Boundary (Bnd.) includes everything required to assemble correct boundary conditions for the calculation of the Final solution.  $P$  is the number of processors,  $q$  is the number of subdomains on a side, and  $C$  is the coarsening factor.  $N$  is the number of cells in one dimension ( $N+1$  the number of grid points in one dimension). Proc-time/pt. is the total time over all  $P$  processors divided by the number of solution points,  $(N+1)^3$ . Percent Comm. is the percent of time spent in the communication phases, Red., Bnd., and GC.**

with many small prime factors (important for the efficiency of FFTW), however, we were not able to keep the work per processor constant. The run parameters and timing results for the performance tests are shown in Table 3.

Ideally, the total processor-time taken per solution point would remain constant. In our results, the times per solution point are fairly stable, at around 20 to 21  $\mu\text{sec}$  for problems on up to 512 processors. On 1024 processors the boundary communication becomes a significant cost and the time per solution point is adversely affected as a result.

Communication overhead is relatively low in Scallop, no more than 25% on up to 512 processors. Nearest neighbor boundary condition communication, significant in all the runs, increases substantially on 1024 processors. After eliminating the extra communication performed in our current implementation, as discussed in Section 4.5, we expect the communication costs for the boundary conditions to be reduced by a factor of  $C$ , resulting in an overall communication cost in the range of 10–15%.

As can be seen in Table 3, time spent on the coarse grid solutions is approximately one third the time spent on initial fine grid solutions. Ideally, the time required for coarse grid solutions would be negligible, but these results match our expectations since we chose values of  $C$  closer to  $q$  than  $2q$ .

In comparing our timing results to our work estimates, we start from the simplest building block and work up. Let us define the *grind time* of a phase as the ratio of the total time spent over all processors divided by the work estimate for that phase from Section 4.2. For the final Dirichlet Poisson solve, the grind time ranges from 1.36 to 1.83  $\mu\text{sec}$ . We believe the variation in performance is largely due to inefficiencies of the FFTW solver on meshes sizes that are non-powers of 2.

Due to our choice of parameters, the global infinite domain solutions were performed on identical mesh sizes for all our problem sizes. Our work estimate for this stage of the computation,  $W_{coarse}^{id}$ , is  $7.18 \times 10^6$  mesh points. The grind times for this phase of the computation vary from 2.01 to 2.28  $\mu\text{sec}$ . The variation in these grind times may be due in part to the small amount of communication required within the global solution in our current implementation. Comparing these grind times with those for the Dirichlet solutions, we infer that the fast multipole method for the infinite-domain boundary calculation adds approximately 35% to the running time above what would be required for the two Dirichlet solutions. This is a significant improvement over our previous Scallop solver where the infinite-domain boundary calculation, performed by rather straightforward numerical integration, dominated the entire solution.

Grind times for the initial local solutions vary from 2.83 to 3.70  $\mu\text{sec}$ . We suspect that the unusual grid sizes required for these calculations and the resulting varying efficiency of the FFTW solver may be contributing to the variation we see here, but the connection is not as clear. The grind times for this phase of the calculation are larger than those for the global infinite domain calculation. In part, this may be due to the extra work required during the infinite-domain boundary calculation to find the extra coarse grid values required later for interpolation.

In summary, we are able to scale a problem up from 128 to 1024 processors with, at worst, a 51% increase in the time per solution point. Running times for each phase of our solver correlate fairly well with simple work estimates based on the number of points updated, with the global and local infinite-domain solutions taking slightly longer than



the final Dirichlet solutions due to the extra work required for boundary condition calculations. Communication times are reasonable for an elliptic partial differential equation solver, generally staying within 25% of the total running time but rising to 37% on 1024 processors. We expect the communication time to drop significantly when our implementation of the boundary condition communication is corrected.

### 5.3 Comparison to Other Methods

To our knowledge there are no other parallel finite-difference infinite-domain solvers with which we can compare our results. Recent related work on fast multipole methods has been for two-dimensional problems. Three-dimensional fast multipole methods, which would be reasonable comparisons for Scallop, are not widely used because the overhead (the constant in front of the  $O(N \log N)$  work estimates) is much larger in three dimensions than in two dimensions. Our approach in Scallop reduces the potential theory component of the problem to two dimensions, leading to an algorithm where the bulk of the time is spent in highly efficient serial fast Fourier transforms.

Our current results do show marked performance improvement over our previous Scallop prototype, however. The performance results for a 128-processor run are shown side by side in Table 4. The fast multipole method greatly reduces the cost of the infinite-domain boundary calculation in both the initial local solution and the global coarse grid solution. Communication times have increased somewhat, reaffirming that optimization is necessary within the Chombo communication routines. Overall, our current version of Scallop is more than three times as fast as the original Scallop prototype.

### 6. Conclusions and Future Work

We have presented a scalable 3D Poisson solver for free space problems that utilizes a method of local corrections that, in practice, reduces communication overheads. The method employs a philosophy for embracing technological change that substitutes relatively inexpensive computation for relatively expensive communication.

We described the design of the Scallop solver, which realizes our strategy. In practice, the performance of Scallop matches the expectations of our performance model quite well. Communication costs are generally less than one quarter of the total running time, and total computation time is dominated by the time required for the initial fine grid calculations. The benefit of little communication comes at the expense of added computation, but this overhead is reasonable, and it remains almost constant, independent of the number of processors. We expect that reimplementing the

boundary condition communication, which currently transfers roughly a factor of 10 more data than necessary, will reduce communication costs to 10–15% of the total running time.

We are currently investigating ways to parallelize the global infinite-domain solution at the center of Scallop algorithm. Even modest parallelism in this phase of the computation would enable significantly increased parallelism overall allowing us to scale performance to many thousands of processors. At the same time, parallelizing the global infinite-domain calculation would lift restrictions imposed on the initial local solutions, further reducing the computational overhead incurred by the method.

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Code Version	Input Parameters				Times for Each Stage (seconds)					Total (sec)	Proc-time/pt. ( $\mu$ sec)
	$P$	$q$	$C$	$N^3$	Local	Red.	Global	Bnd.	Final		
prototype	128	8	6	$768^3$	187.7	1.89	67.3	6.42	4.42	270.7	76.49
Scallop	128	8	6	$768^3$	36.92	6.39	15.73	9.61	4.95	73.83	20.78

**Table 4. Comparison of running times of prototype [7] and current version of Scallop.**

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