Parallel 3D reconstruction of Assymmetric Virus Structure from Electron from Electron Micrographs: Algorithms and Experiments

Robert E. Lynch
Purdue University, rel@cs.purdue.edu

Dan C. Marinescu

Yongchang Ji

Hong Lin

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PARALLEL 3D RECONSTRUCTION OF ASYMMETRIC
VIRUS STRUCTURES FROM ELECTRON MICROGRAPHS:
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Robert E. Lynch
Dan C. Marinescu
Yongchang Ji
Hong Lin

Department of Computer Science
Purdue University
West Lafayette, IN 47907

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Robert E. Lynch, Dan C. Marinescu, Yongchang Ji, and Hong Lin
Department of Computer Sciences
Purdue University
West Lafayette, IN 47907
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Abstract

Recently we proposed a parallel 3D reconstruction algorithm based upon Fourier analysis using Cartesian coordinates. In this algorithm the computations to determine the values of the 3D Discrete Fourier Transforms of the density of an asymmetric object could be naturally distributed over the nodes of a parallel system. In this paper we discuss the accuracy of the algorithm and a method to improve the quality of the reconstruction based upon the CTF, the Contrast Transfer Function, of the electron microscope.

1. Introduction.

The 3D reconstruction of asymmetric objects from 2D projections is a data intensive problem. Indeed the reconstruction of an asymmetric virus structure, at high resolution, may require tens or even hundreds of thousand particle projections. Such projections are obtained experimentally by Cryo-Electron Microscopy.

There are several practical methods for reconstructing a 3D object from a set of its 2D projections. These include use of Fourier Transforms, 'back projection', and numerical inversion of the Radon Transform. See [Gor74] for a review of these and other methods. Also, for descriptions of (sequential) methods for 3D reconstruction and related tasks, see [Dea93], [Fra96], and [Gra96], three of several books containing clear explanations and many references.

In [Lyn99] an outline is given of our first parallel algorithm for 3D reconstruction which was based on algorithm developed by Crowther in early 70's. The sequential algorithm implemented by the 3D reconstruction programs used by the structural biology community uses Fourier Bessel transforms and can be used for reconstruction of symmetric particles [Cro70]. The parallel algorithm uses Cartesian coordinates and permits the reconstruction of objects that do not posses symmetries. The main idea of the algorithm is to decomposes a large linear system with $N^3$ unknowns into $N^2$ systems of linear equations each with $N$ unknowns, that can be solved independently on different processors of a parallel computer or on a cluster of workstations.

In this paper we discuss an improvement of the algorithm in [Lyn99] which, for reconstruction at points of an $N \times N \times N$ grid, uses $O(N^3)$ arithmetic operations instead of $O(N^5)$.

The results obtained with a parallel program based upon our 3D reconstruction algorithm are consistent with the ones produced by a sequential program used for many years for structural biology studies. We report the speedup and the load balancing results for processing cryo-EM data for several viruses. One iteration of the 3D reconstruction for the Bursalia Corella Virus that used to take about 4 hours using a sequential program was carried out in less than 3 minutes on 16 nodes, using the program based upon our algorithm. The algorithm is general and can be used for 3D reconstruction of asymmetric objects for applications other than structural biology.

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The paper is organized as follows. Section 2 provides an informal description of the 3D atomic structure determination and the 3D reconstruction algorithm. Theoretical basis for the algorithm for 3D reconstruction is given in Section 3. In Sections 4, 5, and 6 we present the method used by our algorithm to calculate the 3D Discrete Fourier Transform, DFT, of the electron density using the 2D DFTs of the particles projections. The effect of the zero-fill and the numerical errors in 3D reconstruction are analyzed in Sections 7 and 8. The performance of the parallel program implementing the algorithm is presented in Section 9. The symmetry and the motivation for a parallel algorithm for 3D reconstruction are discussed in Sections 14 and 11. In Section 12 we summarize preliminary performance data when running the parallel program on an IBM SP2, parallel system. In Section 13 we discuss the Contrast Transfer Function corrections. Finally, in Section 15 we present our conclusions.

2. 3D Atomic Virus Structure Determination and 3D Reconstruction.

X-ray crystallography, electron microscopy, EM, and Nuclear Magnetic Resonance, NMR, are techniques used for gathering experimental information about the 3D atomic structure of biological material. The atomic structure of viruses is of considerable interest for the design of anti viral drugs. Viruses are large macromolecules, small virus structures may have few thousands aminoacids, while larger ones have hundreds of thousands of aminoacids and millions of atoms.

X-ray diffraction and electron microscopy are the methods of choice for investigating virus structures. X-ray crystallography is used for high resolution structure determination, in the 2Å to 2.5Å range. Traditionally, electron microscopy produced low resolution maps to, say, 20-30Å; but more recently, researchers at Cambridge and NIH were able to produce maps to 7-7.5Å resolution [Bot97, Con97]. NMR methods can only be applied to smaller structure with few hundreds amino-acids.

The 3D atomic structure determination of macromolecules based upon electron microscopy is an important application of 3D reconstruction. The procedure for structure determination consists of the following steps:

Step a Extract individual particle projections from micrographs.

Step b Determine the center and the orientation of each projection.

Step c Carry out the 3D reconstruction of the biological macromolecule.

Step d Dock an atomic model into the 3D density map if the atomic structure is available.

Steps (b) and (c) are executed iteratively until the 3D electron density map cannot be further refined at a given resolution. The number of iterations for 3D reconstruction is in the 10-20 range and one step of 3D reconstruction for a medium size virus may take several hours on a sequential computer. It typically takes weeks or even months to obtain an electron density map using sequential programs for the orientation determination and for 3D reconstruction.

The maximum resolution for the 3D reconstruction process is limited by the size of a pixel in a particle projection and by the noise in the data. For example if the pixel size is 5.5Å then according to the Nyquist theorem the highest resolution we can possibly reach is 11Å. In practice, the noise in data limits the resolution to values considerably larger than the ones given by the Nyquist limit.

The development of parallel algorithms to carry out some of these computations is part of an ambitious effort to design an environment for 'real-time electron microscopy', where results can be obtained in hours or days rather than in weeks or months.

Algorithms for Step a, which include automatic identification of particle projections, the determination of the center and orientation of each virus particle projection are discussed elsewhere [Mar97], parallel algorithms to determine the orientation, Step b are presented in [Bak97]. We use the terms "projections" and "views" throughout this paper with essentially the same meaning; the first seems more suitable for the description of the algorithm, we associate a direction with one projection, the second one seems more meaningful in the context of the experiment, we have many views of a virus particle.
In this paper we are only concerned with Step c of the process outlined above, the 3D reconstruction of asymmetric objects. This process is carried out as follows:

**Step 1** Compute the 2D Discrete Fourier Transform of each projection.

**Step 2** Compute the 3D Discrete Fourier Transform of the electron density by interpolation knowing the set of 2D projections and their orientations.

**Step 3** Compute the inverse 3D transform to get the electron density.

The 2D Discrete Fourier Transform (DFT) of each projection is computed (Step 1). With the orientation information, the points of intersection of the plane of projection and 3D grid lines can be determined and then estimates of the 3D Fourier Coefficients of the density at non-integral grid points are determined. In the method outlined in [Lyn99], this step took $O(N^5)$ arithmetic operations; in this paper we show how this can be reduced to $O(N^3)$. An inverse 3D DFT is carried out to obtain estimates of the density at grid points (Step 3).

Some of these computations can be done independently from each other. For example in Step 1 each processor can be assigned a set of projections and carry out the 2D DFT concurrently. Data exchange among nodes is necessary to collect information for Step 2 and then each node calculates the Fourier Coefficients on its own set of 3D planes. Different portions of the 3D DFT of the electron density are stored on different nodes; where possible, we carry out 20 inverse transforms and then data are exchanged among nodes and so that the final set of 10 inverse transforms takes place to complete Step 3. Theoretical basis for this 3D reconstruction is given in Section 3.

To use efficiently a parallel computer or a cluster of workstations, we need parallel algorithms that partition the data and computations evenly among nodes to ensure load balance and, moreover, which minimize the communication among processors by maintaining a high level of locality of reference. Similar efforts have been reported in the past [Joh94], but the performance data available to us suggest that new algorithms have to be designed to reduce dramatically the computation time.

The algorithm for 3D reconstruction was designed to work well on workstation clusters. The program uses MPI (Message-Passing Interface) and consists of several phases including initialization, 2D Fourier analysis, estimating the 3D coefficients of the DFT, solving linear systems, and Fourier synthesis. In the first phase of the algorithm, pixel frames are distributed evenly among nodes and processed independently. The values of the 3D DFT are calculated by interpolation and collected for all three directions in a node. The systems are distributed among nodes so that a 20 Fourier synthesis for a plane can be done without an exchange. Then data must then be exchanged so that 1D syntheses can be done to obtain the electron density.

3. **3D Reconstruction by Fourier Transforms.** In this section we summarize the Fourier Transform method (see [Cro70]), the basis of our original algorithm.

The experimental information is gathered from digitized images of many identical macromolecules obtained with a cryo electron microscope. Individual particle projections are identified and the orientation of each 2D image, with respect to a standard XYZ Cartesian System, and the location on the image of the projection of the centroid of the molecule's electron density are determined. See [Lyn99] for an outline and see [Bak97] and [Mar97] for more details.

Each pixel frame is a square $M \times M$ array of pixel values $p$ which are taken as the values of the projection of the electron density, $\rho$, normal to the $(r, s)$-plane of the pixel frame:

$$p_t(r, s) = \int \rho_t(r, s, t) \, dt,$$

where the subscript $t$ indicates the unit vector normal to the $(r, s)$-plane of projection.

The electron density of the macromolecule is to be estimated at grid points of a 3D Cartesian coordinate system, we call the 'XYZ System'; the centroid of the density is at its origin. The Fourier Transform of the electron density is taken as

$$F(h) = \frac{1}{A^3} \int \rho(x) e^{-2\pi i h \cdot x/A} \, dx.$$
where \( T \) denotes transpose, \( x = (x, y, z)^T \), \( h = (h, k, \ell)^T \), and \( A \) denotes the edge-length of a cube containing the electron density. As is customary in crystallography and electron microscopy, we call \( x \) a point in real space and \( h \) a point in reciprocal space.

**Projection Theorem.** We sketch a proof of the projection theorem which states that the 3D Fourier Transform of \( p \) at points on a plane through the origin is equal to the 2D Fourier Transform of the projection of \( p \) onto that same plane; (also see [Cro70], [Den93], etc.).

A rotation about the origin of the XYZ System to an RST Cartesian System is accomplished by the transformation \( r = Ex \), where \( E \) is a 3 \( \times \) 3 orthogonal matrix; that is, \( E \) is real, invertible, and its inverse, \( E^{-1} \), is equal to its transpose, \( E^T \). It is easy to verify that lengths are invariant:

\[
\|x\| = \sqrt{x^T x} = \sqrt{r^T r}
\]

e.g., \( x^T x = x^T (E^T E) x = (Ex)^T (Ex) = r^T r \). Similarly, angles are invariant: if \( u = Eh \), then \( h^T x = u^T r = \|x\| \|h\| \cos \theta \). Thus, \( E \) defines a rotation of the XYZ System about its origin. Rotations of macromolecules are 'proper' so that the determinant of \( E \) is equal to \(+1\) and thus \( dx = dr \) ("improper" rotations have \( \det(E) = -1 \)).

Set \( p(x) = \hat{p}(r) = p(E^{-1} r) \), then, by direct substitution into (2),

\[
F(h) = \frac{1}{A^3} \int \hat{p}(r) e^{-2\pi i \frac{u^T r}{A}} dr = \tilde{F}(u)
\]

In particular, if \( u = (u, v, w)^T \) with \( w = 0 \) and \( (u, v, 0)^T = Eh' \), then

\[
F(h') = \frac{1}{A^2} \int \int \left\{ \frac{1}{A} \int \hat{p}(r, s, t) dt \right\} e^{-2\pi i \frac{(ur+vs)}{A}} dr ds = \tilde{F}(u, v, 0) = P(u, v)
\]

This shows that the 2D Fourier Transform, \( P \), of the 2D projection of the density onto the plane through the origin with normal parallel to the \( t \)-axis is equal to the 3D transform, \( F \), at points \( h' \) on the reciprocal space image of the plane of projection.

**Application of the Projection Theorem.** The macromolecule has finite volume, it is sufficient to use the limits \(-A/2 \leq u \leq A/2\) for each of the three integrations in (2). Then when \( h = (h, k, \ell) \) has integer components, \( F(h) \) is a Fourier coefficient of the periodic extension of \( p \), with period \( A \) in each coordinate direction. Thus its density can be represented as a 3D Fourier Series:

\[
\rho(x) = \sum_h F(h) e^{2\pi i h^T x/A}
\]

where \( h \) has integer components, \( h, k, \) and \( \ell \). Since \( ur + vs \) in (4) is equal to \( u^T r = (h')^T x \), use of the series representation of \( \rho \) yields

\[
F(h') = \frac{1}{A^3} \int \left\{ \sum_h F(h) e^{2\pi i h^T x/A} \right\} e^{-2\pi i h' x/A} dx = \sum_h F(h) \frac{1}{A^3} \int e^{2\pi i (h-h')^T x/A} dx
\]

where the components of \( h' \) are, typically, not integers. Since

\[
\frac{1}{A} \int_{-A/2}^{A/2} e^{2\pi i \ell t/A} dt = \text{sinc}(\ell) = \begin{cases} \sin(\pi \ell)/\pi \ell & \text{if } \ell \neq 0 \\ 1 & \text{if } \ell = 0 \end{cases}
\]

we have, as in [Cro70],

\[
F(h') = P(u, v) = \sum_{h, k, \ell} F(h, k, \ell) \text{sinc}(h - h') \text{sinc}(k - k') \text{sinc}(\ell - \ell').
\]

**Evaluation of \( F(h, k, \ell).** Similar to (5), the projection, \( p \), of the electron density onto the plane of the pixel frame can be represented as a 2D Fourier Series with period \( B \). The series and its coefficients, \( P \), are

\[
p(r, s) = \sum_{u, v = -\infty}^{\infty} P(u, v) e^{2\pi i (ur+vs)/B}, \quad P(u, v) = \frac{1}{B^2} \int_{-B/2}^{B/2} \int_{-B/2}^{B/2} p(r, s) e^{-2\pi i (ur+vs)/B} dr ds,
\]
Figure 1. One-dimensional interpolation on a line parallel to the axes of the 3D grid in the Fourier domain. The \( N \) values of the function at grid points (arrows pointing upwards) are unknown, the ones corresponding to arrows pointing downwards are the \( N \) known/observed values. We can determine by interpolation the values at grid points for all \( H \)-lines and construct the \( H \)-grid. Similarly we obtain the \( K \)-grid and the \( L \)-grid. Then we can average the three independent values of the function, for every grid point and obtain a final grid in the Fourier domain.

and \( P(u,v) \), with \( u, v \) integers, can be used in (8).

One limits each of the indices to a finite number, say \( N \); then there are \( N^3 \) unknown values \( F(h, k, \ell) \). There are \( M^2 \) known values of \( P \) for each projection and suppose there are \( \nu M \) projections, with \( \nu > (N/M)^3 \). For \( N = M \), [Ros98] gives a lower bound on \( \nu \) such that the number of experimental data is equal to the number of unknowns.

One can solve the resulting linear least squares problem using standard mathematical software, such as the LAPACK routine SGEQLS [And92], which uses the singular value decomposition. The order of magnitude of the number of arithmetic operations required to solve this system is the product of the number of equations and the square of the number of unknowns (see [GoI96], p. 263), so when \( M = N \), \( O(N^9) \) arithmetic operations are required.

**Reduction in Arithmetic.** In (8), \( u, v, h, k \), and \( \ell \), are integers and \( P \) is evaluated at grid points of the transformed pixel frame. [Cro70] points out that the computation can be significantly reduced if \( h', k' \) in (8) are the integers \( h, k \). But then the point at which \( P \) is evaluated is no longer a grid point. Instead, (8) reduces to

\[
P(u', v') = \sum_{\ell} F(h, k, \ell) \text{sinc}(\ell - \ell')
\]

where \( u' \) and \( v' \) are not integers. Because \( P \) is not evaluated at a grid point, some kind of interpolation must be used to estimate \( P(u', v') \) in terms of values of \( P(u, v) \) at grid points.

In (10), \( h \) and \( k \) are fixed and one has a single system of equations with \( N \) unknowns; for \( \nu N \) pixel frames, and the complexity of the algorithm to solve the system is \( O(N^3) \). There are \( N^2 \) such systems, one for each integer pair \((h, k)\). So there are \( O(N^3) \) operations required to estimate all the coefficients \( F(h, k, \ell) \), instead of the \( O(N^9) \) operations needed to solve (8).

The process can be repeated for each integer pair \((h, l)\) and \((k, l)\) as indicated in Figure 1. As a result we obtain three estimates for the value of each coefficient \( F(h, k, \ell) \) and by averaging the three estimates the accuracy of the algorithm can be improved. The three sets of values of the coefficients \( F(h, k, \ell) \) corresponding to the pairs \((h, k)\), \((h, l)\) and \((k, l)\) are called the \( L \), \( K \) and \( H \) grid respectively in Figure 1.

For each grid, the \( N^2 \) systems are independent from each other and they can be solved concurrently. This is the basis for the first method we described in [Lyn99].
4. **Projections onto Slabs.** We carried out numerical experiments with our program when (10) was reduced to a diagonal system. The quantity \( F(h, k, \ell) \text{sinc}(\ell - \ell') \) was included on the right side only if \(|\ell - \ell'| < 1/2\). For each \( \ell_j \) and the corresponding \( P(u'_j, v'_j) \), (10) reduces to

\[
P(u'_j, v'_j) = F(h, k, \ell) \text{sinc}(\ell - \ell'_j) + G(h, k, \ell'_j)
\]

such that all the other \( F \)'s are discarded and \( G(h, k, \ell'_j) \) is the error. The least square solution of the set of equations (11) chooses \( F(h, k, \ell) \) to minimize \( E(h, k, \ell) = \sum_j G(h, k, \ell'_j)^2 \) and since

\[
dE(h, k, \ell)/dF(h, k, \ell) = d \left[ \sum_j (P(u'_j, v'_j) - F(h, k, \ell) \text{sinc}(\ell - \ell'_j))^2 \right] /dF(h, k, \ell)
\]

the least square error is minimized

\[
F(h, k, \ell) = \left\{ \sum_j \text{sinc}(\ell - \ell'_j) P(u'_j, v'_j) \right\} / \sum_j \text{sinc}(\ell - \ell'_j)^2
\]

Solving (12) instead (10), reduces the amount of arithmetic by \( O(N^2) \).

In this method, values of \( P \) are given large weights when \( \ell' \) is very close to \( \ell \) because \( \text{sinc}(x) \) is an oscillating function decaying at the rate of \( 1/x \) and has its global maximum at \( x = 0 \).

The resulting electron density values at grid points obtained by a Fourier Synthesis were remarkably close to those obtained, with the same input, by the method described in [Bak88]; the program described there has been producing electron density values for several years which have been accepted as accurate. Based on these experimental results, we constructed a model of the approximations which leads to equations like (10), but having only one term on the right side.

**Piecewise Constant Model.** Variation of the value inside a pixel square cannot be measured, and thus we take \( P \) to be a piecewise constant function on the pixel frame:

\[
p_t(r, s) = p_t(i\Delta r, j\Delta s) \quad \text{for } |r - i\Delta r| < \Delta r/2, \quad |s - j\Delta s| < \Delta s/2, \quad \text{with } \Delta r = \Delta s,
\]

where \( i \) and \( j \) are integers, the subscript \( t \) denotes a unit vector normal to the plane of the pixel frame, and \((i\Delta r, j\Delta s)\) denotes grid points at the center of pixel squares.

If we regard this function to be defined on a plane, then we are lead to the system in (10) because, except at the origin, it is unlikely that any of the grid points of the XYZ system would be in the plane of the projection of a randomly oriented molecule and similarly for its transforms in reciprocal space.

We now regard the projection (1) as a function defined on a slab of thickness \( \Delta r \), rather than on a plane. This not only gives a formulation which is consistent in the three coordinate directions, but also leads to an algorithm using fewer arithmetic operations, namely, \( O(N^3) \) instead of \( O(N^4) \), and produces accurate results.

We take the slab to have thickness \( \Delta r \), use \( t \) to denote a point on the axis normal to the projection, and for each point \((r, s)\), we take the value of the projection to be independent of \( t \) so that the left side of (1) and (13) becomes

\[
p_t(r, s, t) = p_t(r, s, 0), \quad |t| < \Delta r/2.
\]

This function is piecewise constant (because of (13) and (14)) in the three coordinate directions.

5. **Grid Points in Slabs.** Recall that the HKL System is the reciprocal space of the XYZ System of the macromolecule. The UV System is the reciprocal space of the pixel frame and the \( W \)-axis is orthogonal to the UV-plane. The 3-by-3 matrix \( G = \sigma E \) is constructed to relate points in the UVW System and the HKL System:

\[
(u, v, w)^T = G(h, k, \ell)^T
\]
Figure 2. Slab projection simplified to the 2D case. The 1D 'pixel frame' shown on the left, is placed in its correct orientation in the 2D - Discrete Fourier Transform, DFT, \((h,k)\) domain on the right. The DFT at a grid point \((h_i,k_j)\) in the 2D domain is calculated by interpolating the value from the pixel frame, as indicated by the shaded square. In the original method presented in [Lyn99], the unknown DFT values along the \(h\)-line line formed a system of equations with \(N\) unknowns. The known values were the intersections of the \(h\)-line with all 1-D pixel frames.

We are only concerned with points \((u,v,w)^T\) in the reciprocal space of the slab and points \((h,k,\ell)^T\) in the reciprocal space of the molecule that satisfy

\[
|u|, |v| \leq M/2, \quad |w| \leq 1/2, \quad |h|, |k|, |\ell| \leq N/2.
\]

We call the sets of such points satisfying (15) and (16) the \(HKL\) and \(UVW\) systems, respectively. We want to find sets of integers, \(h,k,\ell\), in \(HKL\) and the corresponding \(u,v,w\) in \(UVW\).

Let \(G_{i,j}\) denote the entry that is in the \(i\)-th row and the \(j\)-th column of \(G\). Then the equation in (15) for \(w\) is

\[
w = G_{3,1}h + G_{3,2}k + G_{3,3}\ell.
\]

We choose a pair of integers, \(h,k\), which satisfy (16), and because of the restrictions on \(w\) in (16), we need \(\ell\) which satisfies

\[
-1/2 - G_{2,1}h - G_{2,2}k \leq G_{2,3}\ell \leq 1/2 - G_{2,1}h - G_{2,2}k.
\]

If this condition cannot be satisfied for any integer \(\ell\) with \(|\ell| \leq N/2\), then we reject the pair of integers \(h,k\) and choose another pair.

Suppose that the pair of integers \(h,k\) are such that (18) is satisfied for integers \(\ell_i\) with \(-N/2 \leq \ell_1 < \ldots < \ell_j \leq N/2\). Then for such an integer, \(\ell_i\), we have

\[
u_i = G_{1,1}h + G_{1,2}k + G_{1,3}\ell_i, \quad v_i = G_{2,1}h + G_{2,2}k + G_{2,3}\ell_i.
\]

If \(|u_i| > M/2\) or \(|v_i| > M/2\), then we reject the integers \(h, k, \) and \(\ell_i\) and try another value of \(\ell\).

When we find integers \(h,k,\ell_i\) and corresponding \(u_i, v_i\), such that (16) is satisfied, then we have found a grid point in the \(HKL\) System which corresponds to a point \((u_i,v_i,\ell_i)^T\) in the \(UVW\) System. In our model, which uses piecewise constant functions, we set the value of \(P(u,v,0)\) equal to \(p_t(u,v,t)\), where \(u\) and \(v\) are the integers nearest \(u_i\) and \(v_i\), respectively.

Because \(h,k,\ell_i\) are integers, the equation (10) with \(N\) terms on the right side reduces to

\[
F(h,k,\ell_i) = p_t(u,v,t),
\]

Here the right side is a known value of the transform of the pixel values.

As described above, one exams each of the \(N^2\) integer pairs \(h,k\); however this \(O(N^2)\) work can be reduced to \(O(N)\) because we know, by use of simple linear algebra, that acceptable pairs lie in a
specific slab of width $\Delta u$ containing the origin. Finding the set of acceptable integer pairs reduces to an integer programming problem but one for which we have very specific information about the location of its set of solutions.

Figure 2 illustrates the intuition behind the slab method simplified for the 2D case.

6. Estimate of the Electron Density. Because we are dealing with finite sets of points, Discrete Fourier Transforms (DFT) are used instead of Fourier Transforms.

Each pixel frame produces estimates of the $F(h,k,\ell)$ for a set of grid points, $(h,k,\ell)$, near a plane in the HKL System. These are obtained from (20) in terms of values of the DFT of the pixel frame. For a fixed grid point, $(h,k,\ell)$, suppose that there are estimates of $F(h,k,\ell)$ obtained from $S$ different pixel frames; denote these values by $F_s(h,k,\ell)$, $s = 1, \ldots, S$. The final estimate of $F(h,k,\ell)$ is obtained by averaging:

$$F(h,k,\ell) = \frac{1}{S} \sum_s F_s(h,k,\ell)$$

If there are no estimates of $F(h,k,\ell)$, then $F(h,k,\ell)$ is set equal to zero.

We accept these as the DFT of the the electron density and invert the transform to obtain the estimate of the electron density.

7. The Effect of Zero-Fill. We can put an $M_p \times M_p$ array of pixel values into a larger $M \times M$ array; the extra array entries are set equal to zero. We call this “zero-fill” and the amount of fill is called the “aspect ratio $k$”, defined by

$$\text{zero-fill aspect ratio } k = M/M_p$$

After the zero-fill, 2D DFT of the larger array is determined. Here we show the relationship between the grid spacings of the DFT of the pixel values and the spacing when the domain of the DFT is expanded to a larger domain with zero-fill (see [Bri95], p. 90 ff).

To be able to display informative figures, we simplify the discussion to the case of a 1D set of pixel values at $M_p$ grid points and a 2D density. As we now show, zero-fill results in interpolation (and scaling) of the Discrete Fourier Coefficients in reciprocal space; although the domain in real space increases by the zero-fill factor, the spacing in reciprocal space is reduced by this factor, so the length of the domain is not changed. An example with $k = 2$ is shown in Figure 3.

The spacing in real space is $\Delta A = A/M_p$. For even $M_p$, the DFT of $f$ can be written as

$$\hat{f}_u = \frac{1}{\sqrt{M_p}} \sum_{\mu = -M_p/2}^{M_p/2} f_p \exp(-2\pi i u \mu / M_p),$$

and its inverse as

$$f_p = \frac{1}{\sqrt{M_p}} \sum_{\mu = -M_p/2}^{M_p/2} \hat{f}_u \exp(2\pi i u \mu / M_p).$$

The transformed values are complex. Because $f$ is real valued, the complex valued transformed function, $\hat{f}$, is conjugate symmetric so that its real part is an even function and its imaginary part is an odd function. The real and the imaginary parts of the transform $\hat{f}$ are shown in Figure 3(1b) as piecewise linear interpolants to the values.

The arguments of the exponential can be written in terms of the spacings $\Delta r$ and $\Delta u$ of real space and reciprocal space, respectively, as

$$2\pi i u \mu / M_p = 2\pi i (u/A)(p A/M_p) = 2\pi i (u A)(p \Delta r)$$

so that the spacings in real and in reciprocal space are

$$\Delta r = A/M_p \quad \text{and} \quad \Delta u = 1/A.$$
respectively. The units of $\Delta r$ are Ångstroms and those of $\Delta u$ are reciprocal Ångstroms.

In both spaces, there are $M_p$ points in a fundamental period. The length of the fundamental domain of $f$ in real space is $M_p \Delta r = A$, and the length of the fundamental domain of $\hat{f}$ in reciprocal space is $M_p \Delta u = M_p / A = \Omega$.

Next consider extending the domain of $f$ from $M_p$ grid points to $2M_p$ points with zero-fill (see Figure 3 for an example with $M_p = 8$). The transform is given by

$$F_u = \frac{1}{\sqrt{2M_p}} \sum_{p=-M_p/2}^{M_p/2} f_p \exp(-2\pi i u p / 2M_p).$$

Because of the zero-fill, $f_p = 0$ for $p = \pm M_p/2, \pm (M_p/2 + 1), \ldots$, and we have

$$F_u = \frac{1}{\sqrt{2M_p}} \sum_{p=-M_p/2+1}^{M_p/2} f_p \exp(-2\pi i u p / 2M_p).$$

(24)

The argument of the exponential can be written in terms of $\Delta r$, $\Delta u$, and $\Delta U = \Delta u/2$ as

$$-2\pi i u p / 2M_p = -2\pi i (u/2A)(p\Delta r) = -2\pi i (u\Delta u/2)(p\Delta r) = -2\pi i (u\Delta U)(p\Delta r).$$

The grid spacing $\Delta r$ in real space does not change length when the length $A$ of the real domain is doubled to $2A$ and $f$ is set equal to zero in the extended domain. But, the grid spacing $\Delta U$ in reciprocal space, after the zero-fill, is half that of the spacing before the zero-fill. Consequently, the length of the fundamental domain in reciprocal space after the zero-fill is the same as without the zero-fill; specifically:

$$\Delta U = \Delta u/2, \quad 2M_p \Delta U = 2M_p \Delta u/2 = M_p \Delta u = \Omega,$$

where $\Omega$ is the length of the original reciprocal space fundamental domain as well as the fundamental domain after zero-fill.
The number of pixels on the edge of a \( P \times P \) pixel frame

\( D \)  The diameter of the sphere in pixel edges

\( V \)  The number of pixel frames or 'views'

\( k \)  Zero-fill aspect ratio; the pixel frame is put into a \( kP \times kP \) array

Table 1. The parameters for the test cases.

Comparing (22) and (24), one sees that \( \hat{f}_u = \sqrt{2} F_2 u, u = 0, \pm 1, \pm 2, \ldots \). Hence, \( \sqrt{2} F \) interpolates to \( \hat{f} \) at these points (see Figure 3).

In this example, we doubled the number of grid points and set \( f \) equal to zero at the points outside the fundamental period. It is easy to verify that if we had used \( M \) points, with \( M > M_p \), and set \( f \) equal to zero at the \( M - M_p \) additional points, and kept the spacing \( \Delta r \) in real space constant, then there is no change in the length of the domain in reciprocal space — it is \( \Omega \) with or without zero-fill. The new grid length, \( \Delta U \), in reciprocal space is a fraction of \( \Delta u \), the original reciprocal grid length:

\[ \Delta U = \frac{M_p}{M} \Delta u, \quad M \Delta U = M \frac{M_p}{M} \Delta u = M_p \Delta u = \Omega. \]

Similar to the example above, which considered \( M = 2M_p \), here a value \( F_u \) interpolates to \( \hat{f}_u \) at any point \( v \Delta U \) which is a grid point \( u \Delta u \) in the original domain.

Note that zero-fill does not extend the domain in reciprocal space — one does not obtain estimates of Fourier Transform values having longer wavelengths but, one does obtain estimates on a finer grid in reciprocal space.

These results extend to multi-dimensional transforms. Thus, when the \( M_p \times M_p \) domain in real space of the \( 20 \) pixel frame is enlarged to \( M \times M \) with zero-fill, then the grid spacing is reduced by the factor \( M_p/M \). But the length of the sides of the domain in reciprocal space is the same before as after the zero-fill.

Use of subsets of the computed \( F \)'s  The use of zero fill increases the number of approximate Fourier coefficients (see Figure 3). The pixel frames are \( P \times P \) and the arrays are \( kP \times kP \). In order to reconstruct a density with \( P \times P \times P \) (or fewer) density density values, one must select a portion of the \( kP \times kP \) values. Figure 4 shows the effect of using values only close to the origin — clearly such reconstruction gives inaccurate representation. To obtain a reason reconstruction, one must use value extended across the whole domain. For example in the case illustrated in Figure 4), one could use values at \( x = -2, 0, 2, \) and 4, but certainly not at \( x = -1, -1/2, 0, 1/2, \) and 1 (see Figure 3).

8. Numerical Errors in 3D Reconstruction. 3D reconstruction is subject to experimental as well as numerical errors. In this section we address only the problem of numerical errors. To determine the accuracy of the reconstruction method we take a known object and generate projections for randomly selected orientations.

Then we reconstruct the object from these projections and compare the original values with the computed ones. Our experiments are conducted using a uniform sphere with a constant density inside, equal to one, and zero outside, see Figure 5.

We used the piecewise constant model to compute estimates of the values of 3D DFT coefficients. The parameters for a test case are: the size of a pixel frame, the diameter of the sphere, the number of pixel frames, the aspect ratio for zero-fill, and the size of 3D DFT. The size of output grid is determined by the size of pixel frame, it is \( P \times P \times P \) for \( P \times P \) frames. The parameters for a test care are listed in Table 1.

We calculate the estimate of the maximum and minimum pointwise error and also the mean square error as a function of radius to determine the effect of the zero-fill and the number of views. For the mean square error, we average the square root of the sum of the squares of the errors in a set of annular regions inside the uniform sphere; we do not include grid points at the edge of the sphere where the jump discontinuity occurs.
Figure 4. Use of subsets of $F$. (a),(b),...,(f) show the 'reconstruction' obtained by synthesis of subsets of $F$. In each case we use a subset consisting of $F(0)$ and its nearest neighbors. This leads to inaccurate values of the density.

Figure 5. The original object: a uniform sphere with constant density inside and zero outside. The center of the sphere with diameter $aN$ with $a < 1$ coincides with the center of the cube with edge $N$. 
Figure 6. The transformation of a $P \times P$ pixel frame containing one projection consists of a translation of the origin and zero-fill. The frame is cut into 4 segments and rearranged in a larger frame of size of $kP \times kP$. $k$ is the aspect ratio of zero-fill ($k = 1.5$, in this example), and the centroid of the projection (point $O$) is moved to the upper left corner.

We also report some of these same values as obtained with a sequential program widely used by the structural biology community, EM3DR.

**The effect of zero-fill.**

Before 2D DFT is carried out on pixel frames, the pixel values in the frame are rearranged and put into a larger frame with the extra array entries set equal to zero. Figure 6 illustrates the process of zero-fill. Errors as a function of the zero-fill aspect ratio $k$ are listed in Tables 2 and 3. The mean square errors listed in Table 2 decrease from 20 - 25% for $k = 1$ to about 2% for $k = 4$. For a fixed $k$, the variation in the error is small as the size of the pixel frame changes with the ratio (diameter/frame edge) being kept nearly constant (about 0.79).

The values in Table 3 list the minimum and maximum of the density inside the sphere. The variation of the density becomes smaller when the aspect ratio increases.

**The number of views.** The effect of the number of views on the percent errors and on the minimum and maximum errors are summarized in Tables 4 and 5 for the aspect ratio $k = 4$. For fixed pixel size and diameter, the variation in the percent error varies very little as the number of views increases. Similarly, there is little change in the range of computed values inside the sphere.

Table 2. Mean square error (%) inside reconstructed uniform sphere. "FrameSize" denotes the number of pixels on each size of a pixel frame. "Diameter": denotes the diameter of the sphere in pixels. "NumberViews" denotes the number of projections used for reconstruction.

<table>
<thead>
<tr>
<th>Aspect ratio $k$</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FrameSize $41 \times 41$</td>
<td>FrameSize $61 \times 61$</td>
<td>FrameSize $81 \times 81$</td>
</tr>
<tr>
<td></td>
<td>Diameter 32</td>
<td>Diameter 48</td>
<td>Diameter 64</td>
</tr>
<tr>
<td></td>
<td>NumberViews 20100</td>
<td>NumberViews 45150</td>
<td>NumberViews 80802</td>
</tr>
<tr>
<td>1</td>
<td>20.16</td>
<td>23.38</td>
<td>24.41</td>
</tr>
<tr>
<td>2</td>
<td>6.92</td>
<td>7.97</td>
<td>7.77</td>
</tr>
<tr>
<td>4</td>
<td>1.63</td>
<td>2.11</td>
<td></td>
</tr>
</tbody>
</table>

Table 3. The effect of the zero-fill aspect ratio upon the minimum/maximum density values inside the sphere. "FrameSize" is size of the pixel frames in number of pixels and Diameter is the diameter of the sphere and em NumberViews is of projections used for reconstruction. The true values of the electron density are 1 inside the sphere and 0 outside.
Table 4. The effect of the number of projections upon the least square errors (%) in 3D reconstruction. FrameSize gives the dimensions of the pixel frames in number of pixels, Diameter is the diameter of the sphere in number of pixels, and NumberViews is the number of projections used for reconstruction.

<table>
<thead>
<tr>
<th>NumberViews</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FrameSize $21 \times 21$</td>
<td>FrameSize $41 \times 41$</td>
<td>FrameSize $61 \times 61$</td>
</tr>
<tr>
<td></td>
<td>Diameter 16</td>
<td>Aspect ratio 4</td>
<td>Diameter 32</td>
</tr>
<tr>
<td>300</td>
<td>1.68</td>
<td>2.01</td>
<td>2.21</td>
</tr>
<tr>
<td>1250</td>
<td>1.41</td>
<td>1.77</td>
<td>1.88</td>
</tr>
<tr>
<td>2775</td>
<td>1.39</td>
<td>1.78</td>
<td>1.90</td>
</tr>
<tr>
<td>5000</td>
<td>1.34</td>
<td>1.72</td>
<td>1.88</td>
</tr>
<tr>
<td>11250</td>
<td>1.67</td>
<td>1.89</td>
<td></td>
</tr>
<tr>
<td>20100</td>
<td>1.63</td>
<td>1.95</td>
<td></td>
</tr>
<tr>
<td>45150</td>
<td></td>
<td></td>
<td>2.11</td>
</tr>
</tbody>
</table>

Table 5. The effect of the number of projections upon the minimum/maximum density values inside the sphere. FrameSize gives the dimensions of the pixel frames in number of pixels, Diameter is the diameter of the sphere in number of pixels, and NumberViews represents the number of projections used for reconstruction.

<table>
<thead>
<tr>
<th>NumberViews</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FrameSize $21 \times 21$</td>
<td>FrameSize $41 \times 41$</td>
<td>FrameSize $61 \times 61$</td>
</tr>
<tr>
<td></td>
<td>Diameter 16</td>
<td>Aspect ratio 4</td>
<td>Diameter 32</td>
</tr>
<tr>
<td>300</td>
<td>0.97/1.00</td>
<td>0.96/1.00</td>
<td>0.96/1.00</td>
</tr>
<tr>
<td>1250</td>
<td>0.97/1.00</td>
<td>0.97/1.00</td>
<td>0.96/1.00</td>
</tr>
<tr>
<td>2775</td>
<td>0.97/1.00</td>
<td>0.97/1.00</td>
<td>0.96/1.00</td>
</tr>
<tr>
<td>5000</td>
<td>0.98/1.00</td>
<td>0.97/1.00</td>
<td>0.96/1.00</td>
</tr>
<tr>
<td>11250</td>
<td>0.97/1.00</td>
<td>0.96/1.00</td>
<td></td>
</tr>
<tr>
<td>20100</td>
<td>0.97/1.00</td>
<td>0.96/1.00</td>
<td></td>
</tr>
<tr>
<td>45150</td>
<td></td>
<td></td>
<td>0.96/1.01</td>
</tr>
</tbody>
</table>

We expect that increasing the aspect ratio of the zero-fill allows us to use fewer projections for the 3D reconstruction. Table 6 and Figure 7 show the results when the size of pixel frames is $41 \times 41$ and the number of pixel frames varies from 25 to 200 for zero-fill aspect ratio is 1 and 4. For both zero-fill aspect ratios, the results converge when the number of pixel frames goes beyond a certain value (approximately 75 for this case). However, when zero-fill aspect ratio is 4, with the same number of pixel frames, the results get close to the correct values.

Distribution of the errors as a function of radius. Figure 8 provides conclusive evidence that the magnitude of errors increases with the radius. This increase is quite obvious in Figures 8.1b, 8.2b, 8.3b where we display the density function of the radius for $k = 1$. The oscillation of the graph is quite obvious in this case. For $k = 4$ Figures 8 indicate again that the results are improved only slightly by increasing the number of views.

The effect of noise. Random numbers uniformly distributed between $-0.1$ and $+0.1$ were added to the constant value, $p = 1$, of the density inside the sphere to simulate the effect of the noise.

<table>
<thead>
<tr>
<th>$k$</th>
<th>$V$</th>
<th>25</th>
<th>50</th>
<th>75</th>
<th>100</th>
<th>150</th>
<th>200</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>8.84</td>
<td>4.47</td>
<td>2.08</td>
<td>1.68</td>
<td>1.85</td>
<td>1.97</td>
<td></td>
</tr>
</tbody>
</table>

Table 6. The effect of the zero-fill aspect ratio, $k$, and the number of projections, $V$, upon the errors (%) inside the sphere. The true values are 1 inside the sphere and 0 outside.
Figure 7. The effect of the number of views, $V$, upon the distribution of the density function of the radius of the sphere. For Figures (1a)-(1f), the aspect ratio of zero-fill $k = 1$; for Figures (2a)-(2f), $k = 4$. The size of pixel frames is $41 \times 41$, and the diameter of the sphere is 32.
Figure 8. The effect of the zero-fill upon the distribution of the density inside and outside the sphere function of the radius. The computed values for aspect ratio of zero-fill 1, 2 and 4. The size of pixel frames, the number of pixel frames, and diameter of the sphere differ: for (1a)-(1d) 41 x 41, 20100, and 32, for (2a)-(2d), 61 x 61, 45150, and 48, for (3a)-(3d), 81 x 81, 80902, and 64.
Figure 9. The effect of the number of views upon the distribution of density inside and outside the sphere, function at the radius. The computed values for different numbers of views when the aspect ratio of zero-fill is 4. The size of the pixel frames and the diameter of the sphere are: for (1a)-(1d) 21 x 21, and 16, for (2a)-(2d) 41 x 41, and 32, for (3a)-(3d), 51 x 61, and 48.
Figure 10. The effect of noise. Pixel values were computed for the 41 x 41 projections of the uniform sphere with diameter of 32. Then uniformly distributed random numbers equal to 5% of the maximum projected density were added to the pixel values.

Comparison of results of parallel 3D reconstruction program with the results produced by the sequential program EM3DR. Table 7 lists the mean difference (ε), the variance (σ) and coefficient of variation (δ) of the difference between the density computed by the parallel program based the algorithm described in this paper with the density computed by the sequential program, EM3DR.

Both versions of parallel 3D reconstruction program produced results close to those of EM3DR,
Table 8. The least square errors (%) for 3D reconstruction with a widely used sequential program that uses icosahedral symmetry. \textit{FrameSize} and \textit{NumberViews} are the size and number, respectively, of pixel frames. \textit{InsideError} is least square error inside the sphere.

\begin{table}[h]
\begin{tabular}{|c|c|c|}
\hline
\textit{FrameSize} & \textit{NumberViews} & \textit{InsideError} (%) \\
\hline
21 x 21 & 45 x 60 & 28.45 \\
41 x 41 & 198 x 60 & 51.29 \\
61 x 61 & 459 x 60 & 33.35 \\
81 x 81 & 831 x 60 & 32.46 \\
\hline
\end{tabular}
\end{table}

Table 9. The minimum and maximum density for 3D reconstruction with a widely used sequential program that uses icosahedral symmetry. \textit{FrameSize} and \textit{NumberViews} are the size and number, respectively, of pixel frames. \textit{InMin} and \textit{InMax} are the minimum and the maximum of the density value inside the sphere, \textit{OutMin} and \textit{OutMax} are the values outside the sphere, respectively. The true values are 1 inside and 0 outside.

\begin{table}[h]
\begin{tabular}{|c|c|c|c|c|c|}
\hline
\textit{FrameSize} & \textit{NumberViews} & \textit{InMin} & \textit{InMax} & \textit{OutMin} & \textit{OutMax} \\
\hline
21 x 21 & 45 x 60 & 0.4787 & 1.0000 & 0.0000 & 0.1615 \\
41 x 41 & 198 x 60 & 0.9225 & 1.8677 & -0.0099 & 0.4451 \\
61 x 61 & 459 x 60 & 0.3715 & 1.0000 & -0.0199 & 0.1870 \\
81 x 81 & 831 x 60 & 0.5659 & 1.5402 & -0.0859 & 0.3422 \\
\hline
\end{tabular}
\end{table}

a sequential program based upon the method described in [Cro70] and widely used for structural studies using cryo-EM data. The improved algorithm leads to slightly better agreement with the results produced by the sequential program.

Table 8 indicates that the least square errors are slightly larger than the ones reported above for zero-fill aspect ratio of 1. Table 9 indicates that the range of density values inside the sphere is slightly wider than the ones reported above even for \( k = 1 \). Figure 11 shows the distribution of the density for for several cases.

9. The performance of a parallel program implementing the 3D reconstruction algorithm. A program implementing the algorithm presented in this paper was written and tested using internal data as described in the previous section as well as several experimental data sets. The program is written in Fortran, uses the MPI library for communication, and was designed to run efficiently on a cluster of inexpensive PCs. We use a cluster of 16, 400 MHz Pentium II processors running SunOS5.6. Each processor has 256 MB of main memory and a 8 GB disk. The connectivity is provided by a 100 MBps Ethernet switch. The total cost of the system is about $40K. The actual performance of this system is comparable for this problem with the performance of a 16 processor SGI Origin 2000.

The program is based upon a data parallel execution model, all nodes perform essentially the same computation but on different data. The coordinator node reads the input files containing the set of projections and the orientation of each projection and then distributes the projections evenly among the set of available nodes. Then, each node processes the individual frames assigned to it; first it transforms each frame and expands it, if the zero-fill aspect-ratio \( k > 1 \) and then carries out a 2D DFT. A data exchange stage occurs at the end of the Fourier analysis phase, each node is assigned a set of linear equations. After solving the linear systems the nodes carry out a 2D DFT then a global exchange takes place and a 1D FFT completes the Fourier synthesis phase. Finally, the coordinator node gets individual sections of the 3D map from the other nodes and writes the electron density map out. We have opted for this solution because we do not have a parallel file system and several nodes reading the input data concurrently and then writing the output density maps concurrently would lead to an unacceptable performance degradation due to I/O contention.

We are primarily interested in the load balancing properties of the algorithm and in the speedup of the implementation. While the tests conducted with the internal data presented in Section 7 gave us enough confidence in the correctness of the algorithm and its implementation, we used actual data
collected in cryo-EM experiments as shown in Table 10, to further test the correctness of our program. We used only data for symmetric objects because our objective was to compare our results with the results produced by a sequential program widely used by the structural biology community. Figure 12 shows the picture of Paramedium Bursaria Chlorella Virus, type 1 (or PBCV₁), reconstructed with the parallel program using the algorithm presented in this paper, see Cases E and I in Table 10.

A first objective of our analysis is to profile the program and determine the time used for each execution phase. Table 11 shows that interpolation is the most intensive phase, followed by the 2D Fourier analysis, while solving the linear systems requires a relatively low amount of arithmetic operations. In [Lyn99] we reported that solving the linear systems was the most time-consuming phase of 3D reconstruction. The expected improvements of the algorithm described in this paper are confirmed by the measurements.

Our next objective is to study the load balancing properties of the algorithm and the speedup. Table 12 shows the time used in each node when the program solves one of the problems in multiple nodes. From the data in Table 12, we can see that the computation is evenly distributed among multiple nodes. We need to keep in mind that the coordinator is assigned extra duties in the initial and final phases of the algorithm.

Table 13 shows the speedups, for the nine problems presented above. The speedups are larger than 1.85 for two nodes, larger than 3.5 for four nodes, range from a low of 3.7 to a high of 6.9 in eight nodes and from 6.8 to 11.1 in 16 nodes. In case of problem H and I due to the problem size we were unable to run in one node and report only the speedups relative to the running time in two nodes.

14. Exploiting symmetry. The 3D reconstruction algorithm was specifically designed for asymmetric objects. However, when used for objects with known symmetry, such as dihedral, icosahedral, or other type of symmetry, the symmetry can be used to decrease the execution time and only a portion of the object has to be reconstructed.

Such a symmetric object is composed of a number $A$ asymmetric units, e.g., an object with
Table 10. Data for 9 (nine) problems used to test the parallel 3D reconstruction program. The virus family is indicated in parenthesis. The number of pixels, the number of views/projections and the types of symmetry are indicated.

<table>
<thead>
<tr>
<th>Problem</th>
<th>Virus</th>
<th>Pixels</th>
<th>Views</th>
<th>Symmetry</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Polyomavirus (Papovaviruses)</td>
<td>69 x 69</td>
<td>158 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>B</td>
<td>Papillomavirus (Papovaviruses)</td>
<td>99 x 99</td>
<td>60 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>C</td>
<td>Sindbis virus (Alphaviruses)</td>
<td>221 x 221</td>
<td>389 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>D</td>
<td>Sindbis virus (Alphaviruses)</td>
<td>221 x 221</td>
<td>643 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>E</td>
<td>Paramecium Bursaria Chlorella Virus, type 1</td>
<td>281 x 281</td>
<td>107 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>F</td>
<td>Ross River Virus (Alphaviruses)</td>
<td>131 x 131</td>
<td>1777 x 10</td>
<td>Dihedral</td>
</tr>
<tr>
<td>G</td>
<td>Bacteriophage Phi29</td>
<td>191 x 191</td>
<td>609 x 10</td>
<td>Dihedral</td>
</tr>
<tr>
<td>H</td>
<td>Aravivirus (Alphaviruses)</td>
<td>331 x 331</td>
<td>1940 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>I</td>
<td>Paramecium Bursaria Chlorella Virus, type 1</td>
<td>359 x 359</td>
<td>948 x 60</td>
<td>Icosahedral</td>
</tr>
</tbody>
</table>

Figure 12. The Paramecium Bursaria Chlorella Virus, type 1, reconstructed using the algorithm and the program discussed in this paper.

<table>
<thead>
<tr>
<th>Execution Phase</th>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initialization</td>
<td>0.26</td>
<td>0.13</td>
<td>0.13</td>
</tr>
<tr>
<td>2D Fourier analysis</td>
<td>1.75</td>
<td>1.2</td>
<td>75.9</td>
</tr>
<tr>
<td>Interpolation</td>
<td>11.49</td>
<td>8.34</td>
<td>270.2</td>
</tr>
<tr>
<td>Data Exchange for solvesys</td>
<td>0.0016</td>
<td>0.006</td>
<td>0.073</td>
</tr>
<tr>
<td>Solvesys and combine</td>
<td>0.019</td>
<td>0.053</td>
<td>0.64</td>
</tr>
<tr>
<td>2D Fourier synthesis</td>
<td>0.067</td>
<td>0.23</td>
<td>4.49</td>
</tr>
<tr>
<td>Data Exchange for 1D synthesis</td>
<td>0.010</td>
<td>0.027</td>
<td>0.33</td>
</tr>
<tr>
<td>1D Fourier synthesis</td>
<td>0.030</td>
<td>0.10</td>
<td>2.15</td>
</tr>
<tr>
<td>Gather data</td>
<td>0.0050</td>
<td>0.015</td>
<td>0.18</td>
</tr>
<tr>
<td>Write density map</td>
<td>0.2</td>
<td>0.45</td>
<td>3.11</td>
</tr>
</tbody>
</table>

Table 11. Time (in seconds) for each phase of the 3D reconstruction program for problems A, B and C running in one node.
Table 12. Time (in seconds) used by each node for problem D, Execution with 1, 2, 4, 8, and 16 nodes.

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>4</th>
<th>8</th>
<th>16</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td></td>
<td>1.82</td>
<td>2.79</td>
<td>1.16</td>
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<td>B</td>
<td></td>
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<td>2.86</td>
<td>1.54</td>
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<tr>
<td>C</td>
<td></td>
<td>1.82</td>
<td>3.57</td>
<td>6.18</td>
<td>8.39</td>
</tr>
<tr>
<td>D</td>
<td>1.76</td>
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<td>6.19</td>
<td>9.35</td>
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</tr>
<tr>
<td>E</td>
<td>1.73</td>
<td>3.25</td>
<td>4.31</td>
<td>5.98</td>
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<tr>
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<td>3.63</td>
<td>6.32</td>
<td>7.73</td>
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<tr>
<td>G</td>
<td>1.92</td>
<td>3.21</td>
<td>5.77</td>
<td>7.00</td>
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<tr>
<td>H</td>
<td></td>
<td>1.93</td>
<td>2</td>
<td>4.04</td>
<td>7.62</td>
</tr>
<tr>
<td>I</td>
<td>1.87</td>
<td>2.77</td>
<td>4.84</td>
<td>2</td>
<td></td>
</tr>
</tbody>
</table>

Table 13. The speedups attained by the 3D reconstruction program for the nine sample problems. The last two problems could only run in two or more nodes.

<table>
<thead>
<tr>
<th>Input \ Node Number</th>
<th>2</th>
<th>4</th>
<th>8</th>
<th>16</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>2.79</td>
<td>1.16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B</td>
<td>2.86</td>
<td>1.54</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>3.57</td>
<td>6.18</td>
<td>8.39</td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>3.50</td>
<td>6.19</td>
<td>9.35</td>
<td></td>
</tr>
<tr>
<td>E</td>
<td>3.25</td>
<td>4.31</td>
<td>5.98</td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>3.63</td>
<td>6.32</td>
<td>7.73</td>
<td></td>
</tr>
<tr>
<td>G</td>
<td>3.21</td>
<td>5.77</td>
<td>7.00</td>
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<tr>
<td>H</td>
<td>2</td>
<td>4.04</td>
<td>7.62</td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>2.77</td>
<td>4.84</td>
<td>2</td>
<td></td>
</tr>
</tbody>
</table>
Icosahedral symmetry consists of $A = 60$ asymmetric units, one with dihedral symmetry has $A = 10$. If a projection plane is normal to the orientation of one of the asymmetric units, then for each of the other asymmetric units there is an orientation normal to the same projection.

The symmetry of an object allows us to use fewer pixel frames for its 3D reconstruction. One particle projection can be used repeatedly. For example, in case of icosahedral symmetry, one projection has associated with it 60 orientations; in dihedral symmetry, one projection has 10 orientations. This allows us to carry out the Fourier Transform of a pixel frame only once and use the transform $A$ times in estimating the 3D DFT by interpolation. In case of a symmetric particle the number of arithmetic operations increases, because the number of projections is multiplied by $A$. At the same time we have a corresponding reduction in the amount of I/O operations and the input data size simply because we reduce the number of projections by $A$.

The symmetry in real space imposes symmetry in reciprocal space and this effect can be used to reduce the number of arithmetic operations required to compute the 3D DFT. For example to reconstruct an object with icosahedral symmetry, we only need to estimate the 3D DFT in one of the octants instead of four. This results in the reduction of the number of arithmetic operations for interpolation as well as in solving linear systems by a factor of four.

11. Motivations for using parallel algorithms for 3D reconstruction of asymmetric objects. The 3D reconstruction algorithm proposed by Crowther [Cro70] almost 30 years ago has been used extensively by the structural biology community ever since. The algorithm is based upon Fourier-Bessel Transforms and can be used for reconstruction of symmetric objects.

The protein shell of a spherical virus exhibits various degrees of symmetry, but the core of the virus consisting of genetic material does not. Structural studies of the virus core provide the motivation for 3D reconstruction algorithms of asymmetric objects.

The amount of experimental data for the reconstruction of an asymmetric object is considerably larger than the one for a symmetric one as discussed in Section 14. While a typical reconstruction of an icosahedral virus at say 20Å resolution may require a few hundreds projections, e.g. 300, the reconstruction of an asymmetric object with the same dimensions and at the same resolution would require 60 times more data, i.e. 18,000 projections.

X-ray crystallography is the only method to obtain high resolution (2-2.5Å) electron density maps for large macromolecules like viruses, while until recently electron microscopy was only able to provide low resolution (20Å) maps. Cryo-EM is appealing to structural biologists because crystallizing a virus is sometimes impossible and even when possible, it is technically more difficult than preparing samples for microscopy. Thus the desire to increase the resolution of cryo-EM methods to the 5Å range.

In the last years results in the 7-7.5Å range have been reported, [Bot97], [Con97]. But increasing the resolution of the 3D reconstruction process requires more experimental data. It is estimated that the number of views to obtain high resolution electron density maps from cryo-EM micrographs should increase by two order of magnitude from the current hundreds to tens of thousands.

The amount of experimental data may further increase because structural studies of even larger virus-antibody complexes may be necessary. Last but not least, using larger zero-fill aspect ratios to improve the accuracy of 3D reconstruction will increase the number of arithmetic operations and the amount of space needed for reconstruction. Thus it is not unrealistic to expect an increase in the volume of experimental data for high resolution asymmetric objects by three to four orders of magnitude in the next future.

Even though nowadays faster processors and larger amounts of primary and secondary storage are available at a relatively low cost, the 3D reconstruction of asymmetric objects at high resolution requires computing resources, CPU cycles, primary and secondary storage, well beyond those provided by a single system. Thus the need for parallel algorithms.

12. Preliminary results on a high performance system.

The parallel program implementing the algorithm presented in this paper can benefit from the ample computing resources of a parallel computing system. Faster processors, larger main memory and secondary storage space, higher communication bandwidth and reduced communication latency, lead to reduced execution time and better speedups.

We ported the parallel program to an IBM SP2 system with 64 nodes each with four processors.
Table 14. The 11 (eleven) problems used to test the parallel 3D reconstruction program on an IBM SP2. The virus type, the number of pixels, the number of views/projections and the types of symmetry are indicated.

<table>
<thead>
<tr>
<th>Problem</th>
<th>Virus Type</th>
<th>Pixels</th>
<th>Views</th>
<th>Symmetry</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Polyomavirus (Papovaviruses)</td>
<td>99 x 99</td>
<td>60 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>B</td>
<td>Paramecium Bursaria Chlorella Virus, type 1</td>
<td>281 x 281</td>
<td>107 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>C</td>
<td>Anivirus (Alphaviruses)</td>
<td>321 x 321</td>
<td>1940 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>D</td>
<td>CIV Chilo Iridescent Virus</td>
<td>342 x 342</td>
<td>697 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>E</td>
<td>Herpes Virus</td>
<td>511 x 511</td>
<td>173 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>F</td>
<td>Ross River Virus (Alphaviruses)</td>
<td>131 x 131</td>
<td>1777 x 10</td>
<td>Dihedral</td>
</tr>
<tr>
<td>G</td>
<td>Paramecium Bursaria Chlorella Virus, type 1</td>
<td>359 x 359</td>
<td>948 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>H</td>
<td>Bacteriophage Phi29</td>
<td>191 x 191</td>
<td>609 x 10</td>
<td>Dihedral</td>
</tr>
<tr>
<td>I</td>
<td>Sindbis virus (Alphaviruses)</td>
<td>221 x 221</td>
<td>389 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>J</td>
<td>Sindbis virus (Alphaviruses)</td>
<td>221 x 221</td>
<td>643 x 60</td>
<td>Icosahedral</td>
</tr>
<tr>
<td>K</td>
<td>Polyomavirus (Papovaviruses)</td>
<td>69 x 69</td>
<td>158 x 60</td>
<td>Icosahedral</td>
</tr>
</tbody>
</table>

Table 15. The execution time in seconds of the parallel 3D reconstruction program in 1, 2, 4, 8, 12, 16, 20, 24, 28, and 32 processors on the IBM SP2. The 4 processors in each node share the node's main memory and communicate using MPI with local processors and with processors in a different node. The execution time does not include the time to write out the 3D electron density map.

The processors are POWER II, running at 375 Mhz, each node has 4 GBytes of main storage and has a 36.4 GByte disk.

We ran the parallel program using 1 to 32 processors for the eleven problems in Table 14. The results are summarized in Table 15. The speedups are good for large problems, B-H. Memory limitations prevented us from running problems, C, D, E, and G in one processor.

To assess the quality of the results produced with the parallel program we compared several electron density maps produced by our program with the ones produced by the sequential program. In Figures 13 and 14 we show a central cross-section, section 171, of the electron density map obtained with the parallel and with the sequential program respectively. The result in Figure 14 is from reference [Yan00]. The virus is a lipid-containing, dsDNA icosahedral virus, the Chilo Iridescent Virus, CIV, with a diameter of 1850Å. The virus has a layered structure consisting of a dsDNA-protein core, surrounded by a lipid bylayer and icosahedral capsid shells consisting of thousands of subunits. The outer diameters of the virus capsid range from 1615Å along the two and three-fold axes to 1850Å along the five-fold axis [Yan00].

13. The effect of the CTF corrections. The relationship between the electron image of a specimen and the specimen itself is described by the Contrast Transfer Function, CTF, [Bak98]. The phase and the amplitude of the CTF are characteristic to a particular microscope, the specimen, and the conditions of imaging.

The CTF that enables the visualization of unstained specimens must be compensated in the
Figure 13. Crosssection 171 of the CIV, Chilo Iridiscent Virus at 27Å resolution. The reconstruction is done with the parallel program based upon the algorithm presented in this paper and running on an SP2 system. The time to produce the electron density map is 280 seconds using 32 processors. The size of a frame is 342 x 342 pixels. The zero fill aspect ratio is $k = 1.5$. The image for $k = 1.5$ is slightly better than that for $k = 1.0$.

Figure 14. Crosssection 171 of the CIV, Chilo Iridiscent Virus at 27Å resolution. The 3D reconstruction is done with the sequential program based upon an 3D algorithm due to Crowther. The size of a frame is 342 x 342 pixels and it is zero filled to 512 x 512. The image is from [Yan00]. The time to produce the electron density map is more than two hours using a 400 MHz Alpha processor with 500 MBytes of memory.
Figure 15. Crosssection 80-83 of phX174, a bacteriophage virus with icosahedral symmetry at 10 Å resolution without CTF correction.

Figure 16. Crosssection 80-83 of phX174, a bacteriophage virus with icosahedral symmetry at 10 Å resolution with CTF correction.
reconstruction on order to achieve a reliable representation of the structure. The transfer function reverses, removes, and attenuates data in the image. The effects of the transfer function are more important at higher resolution. One can correct the transfer function by Winner filtering.

Our algorithm allows an effective CTF correction. In the interpolation phase we weight experimental data points coming from different micrographs with the corresponding value of CTF.

Figures 15 and 16 show cross-section 80-83 of ph:X174, a bacteriophage virus with icosahedral symmetry at 10 Å resolution without and with CTF correction. The improvement in the quality of the reconstruction is visible.


In this paper we discuss an improvement of our 3D reconstruction algorithm for objects without symmetry based upon Fourier analysis using Cartesian coordinates [Lyn97], [Lyn99] which, for reconstruction at points of an \( N \times N \times N \) grid, uses \( O(N^3) \) arithmetic operations instead of \( O(N^5) \).

Though developed for structural biology studies the algorithm is general and can be used for any applications to reconstruct a 3D object from its 2D projections.

We report the results of an error analysis that shows that embedding the pixel frames into larger arrays, a technique we call "zero-fill", helps lower the numerical errors in the reconstruction process but increases the amount of space and the number of arithmetic operations. For example we can reduce the error inside a uniform sphere from about 25% for a zero-fill aspect ratio of \( k = 1 \) to less than 4.5% for \( k = 4 \). For the sake of completeness we report on errors both inside and outside a uniform sphere though for practical purposes only the errors inside matter.

The magnitude of the least square errors of a 3D reconstruction program based upon the algorithm presented in this paper is slightly lower than the ones for a sequential program based upon one of the algorithm described in [Cro70] when the zero-fill aspect ratio is one and significantly lower when we increase \( k \).

In practice, the data collected in cryo-electron microscopy is subject to experimental errors due to various factors e.g. variations of the intensity of the beam, the non-uniform layer of ice, and other sources of noise. Additional errors occur when extracting the individual projections from the micrographs, determining the center of each projection, etc. The traditional wisdom is that using a number of projections much larger than the minimum number required for reconstruction, see [Ros98], the effect of errors can be overcome. Indeed many structures have been solved along the years yet the reconstruction was carried out at relatively low resolution.

Our results confirm our intuition that errors have a non-uniform distribution, the further we are from the center of the sphere the larger are the errors. We studied also the effect of the number of projections upon the magnitude of errors. Since we are performing a Monte Carlo calculation, we expect that the least square error should decrease as \( 1/\sqrt{\text{number of views}} \). Interestingly enough, the least square errors seem to decrease even slower than the rate above, e.g. in one case, the error in the density inside a uniform sphere was only 4.59% for 1250, 4.51% for 5000, and 4.45% for 20100 projections.

We profiled the program and report results regarding the parallel aspects of our algorithm, namely load balance and speedup. As expected, the most time consuming phases of the program execution are: (a) the interpolation, (b) the 2D Fourier analysis, and (c) the initialization phase where input files containing the projections and the orientation of each projection are read in. Recall that in our previous experiments [Lyn99] we reported that the most time consuming phase of the program was solving linear systems.

The load balancing results are very good. In most cases the execution times of all but the coordinator node are within 1% of each other. The need to exchange data among nodes and to synchronize after each phase reduces somewhat the speedup on realistic problems. We report on the results of 3D reconstruction for 8 virus structures. The speedup in 4 nodes is about 3.5, ranges from a low of 3.7 to a high of 6.9 for 8 nodes and is in the range of 7 to 11 for 16 nodes. While the problem size may be too small in some of the cases we report on there is no doubt that further improvements in the implementation of the algorithm are needed.

The original experiments reported in this paper were conducted on a low-cost parallel system.
consisting of a cluster of 16 Pentium II processors running at 400 MHz, each with 256 MB of memory, interconnected by a 100 Mbps Ethernet.

Preliminary results on an IBM SP2 parallel machines are also reported. The speedup relative to a sequential program is impressive and the quality of the solution is comparable.

The results reported here are only for symmetric particles. No experimental data for asymmetric particles were available to us. The number of particle projections necessary for the reconstruction of an asymmetric particle is almost two orders of magnitude larger than for one with icosahedral symmetry. We attempted to reconstruct the CIV without imposing the known icosahedral symmetry, using the few hundred projections available to us but, as expected, the results were not very encouraging due to the insufficient amount of experimental information.

15. Acknowledgments. The authors are grateful for many insightful discussions with Timothy S. Baker and Michael G. Rossmann. Robert Aslunore from Baker's lab provided assistance with the sequential reconstruction program. We thank Wei Zhang and Xiaodong Yan, graduate students in the Biology Department at Purdue University, for very helpful information about the EM3DR program and for sharing their data with us.

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