Three-dimensional reconstruction of the giant mimivirus particle with an X-ray free-electron laser


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We present a proof-of-concept three-dimensional reconstruction of the giant Mimivirus particle from experimentally measured diffraction patterns from an X-ray free-electron laser. Three-dimensional imaging requires the assembly of many two-dimensional patterns into an internally consistent Fourier volume. Since each particle is randomly oriented when exposed to the X-ray pulse, relative orientations have to be retrieved from the diffraction data alone. We achieve this with a modified version of the expand, maximize and compress (EMC) algorithm and validate our result using new methods.

INTRODUCTION

Free-electron lasers provide femtosecond X-ray pulses with a peak brilliance ten billion times higher than any previously available X-ray source. Such a large jump in one physical quantity is very rare, and can have far reaching implications for several areas of science. It has been suggested that such pulses could outrun key damage processes and allow structure determination without the need for crystallization[1]. In 2006 came the first verification of this “diffraction before destruction” method with the reconstruction of a silicon nitride nanostructure created with a focused ion beam (FIB) and exposed to the FLASH free-electron laser in Hamburg[2].

So far, imaging applications at FELs have mainly been limited to nanocrystallography and to two-dimensional projections of single particles while 3D reconstructions from single particles have remained elusive.

Nanocrystallography[1][3] is an extension to protein crystallography where the high intensity and short pulse duration of a FEL allow for the use of very small crystals. Some proteins only produce small crystals. However, the fundamental problem that some samples are hard or impossible to crystallize is still valid. For this reason, single-particle imaging was a key part of the scientific case for building X-ray free-electron lasers.

2D imaging with FELs such as the imaging of live cells[4][5], organelles[6] and viruses[7] is a promising method for imaging irreproducible samples. A resolution down to 21 nm have been achieved on Carboxysomes in a recent study[8]. There is also one application where a 2D image from a single-shot FEL experiment was compared to regular X-ray diffraction tomography performed at a synchrotron[8]. In a recent paper the structure of simple gold nanostructures were recovered in 3D from one single diffraction patterns[8]. This technique is however restricted to structurally simple and strongly scattering structures with a high degree of symmetry.

Several fundamental challenges exist for a general method of 3D single-particle imaging. First, 3D imaging requires the assembly of diffraction patterns from many identical copies of a reproducible object. Many of the applications of 2D imaging so far have been dealing with cells or other particles where each sample is structurally unique. Second, there is no way to directly measure the orientation of the sample when it was hit in the X-ray pulse. Instead the orientation of each sample particle has
to be recovered from the noisy signal of the diffraction patterns.

Solving these problems not only gives more information about the sample by presenting the structure in 3D, it is also a necessity for extending signal from weakly scattering samples such as proteins and small viruses. For these samples the scattering from a single particle may be too weak for reconstructing a 2D projection image and increasing the signal to noise ratio (SNR) by merging many patterns could allow for phasing even in this case.

A solution to the orientation problem was proposed by Duane Loh and Veit Elser in 2009 with the expand, maximize and compress (EMC) algorithm \[10\] which was verified for simulated diffraction patterns in the original publication. Later the algorithm was also tested for an artificial sample although at a resolution that was too low to allow for phase retrieval\[11\]. This paper presents the first application of the algorithm to a biological sample.

The sample used in this study was the Mimivirus (Acanthamoeba polyphaga mimivirus) particle\[12\]\[13\]\[14\]. Mimivirus is one of the largest known viruses. The viral capsid is about 450 nanometers in diameter and is covered by a layer of thin fibres. A 3D structure of the viral capsid exist\[14\] but the 3D structure of the inside is currently unknown.

**EXPERIMENTAL SETUP AND DATA PREPROCESSING**

Mimivirus particles were aerosolized and then focused to a narrow particle stream using an aerodynamic lens. The beam of particles was intersected with the pulse train of the Linac Coherent Light Source (LCLS). Diffracted light was collected on a detector placed 0.7 m downstream of the interaction region. At this distance the collected diffraction signal corresponds to the amplitude squared of a slice through the Fourier transform of the electron density of the sample. Mimivirus particles that were not hit by the FEL were shown to remain infectious after the injection process suggesting that they were not harmed by the injection process. A detailed description of the setup can be found in the Supplemental Material. A total of 198 diffraction patterns were selected and preprocessed and a subset of 25 of these are shown in FIG. 1. The selection preprocessing and selection is explained in supplemental material.

**ORIENTATION RECOVERY**

Three-dimensional structure determination requires the assembly of many 2D diffraction-patterns into an internally consistent 3D Fourier volume. A diffraction pattern represents an Ewald-sphere slice through the 3D Fourier-transform of the electron density. Since each particle is randomly orientated when exposed to the X-ray pulse, the relative orientations of the particles have to be retrieved from the diffraction data alone. This was done using a modified version of the EMC algorithm\[10\]. This algorithm has been verified for simulated data\[10\] and has been experimentally tested using artificial “nanorice” particles at a resolution too low to permit phase retrieval\[11\].

In the EMC algorithm a 3D diffraction space is iteratively updated to comply with the experimental data in the three steps expand, maximize and compress. In the expand step the current diffraction space is expanded into tomograms by taking slices through the diffraction space at a discrete sampling of all rotations. In the maximize step all tomograms are compared to all experimental diffraction patterns by calculating the probability of detecting the experimental pattern while treating the tomograms as expectation values. New tomograms are then created by summing together all diffraction patterns weighted by the respective calculated probability. In the compress step a new 3D diffraction space is assembled from the new tomograms.

For this study we introduce a new similarity function in the maximize step that is based on a Gaussian model:

\[
L(K, M) = \prod_i e^{-\frac{(M_i - K_i)^2}{2\sigma_i^2}}
\]

where \(K\) is the diffraction pattern, \(M\) is the slice through the 3D diffraction space, \(i\) is the pixel index and \(\sigma_i\) is the standard deviation of the Gaussian. We set \(\sigma_i = A\sqrt{M_i}\) where \(A\) is a constant. This similarity function balances well the contribution from the few but high-intensity central pixels and the numerous low-intensity outer pixels.
FIG. 2. The assembled three-dimensional diffraction space. (a): The first ten patterns are shown in their recovered best orientations. Each diffraction pattern represents a slice through the squared modulus of the 3D Fourier-transform of the electron density. (b): All 198 diffraction patterns plotted with a section cut out to show the central part of diffraction space. Diffraction symmetry and object symmetry can be directly recovered from the measured diffraction data in the EMC process.

The photon fluence at the particle is unknown in this type of experiment since neither the exact profile of the X-ray pulse nor the exact position of the particle in the beam is known. The fluence therefore needs to be recovered from the diffraction pattern in the EMC process just like the orientations of the particles.

We used a variation of the method described in with the two following key differences: (i) A new fluence is calculated for each comparison between a diffraction pattern and a slice through Fourier space instead of using one fluence per pattern. (ii) The calculation of the fluence maximizes the likelihood function under the new distance metric given in equation (1). The fluence is thus given as:

$$\phi(K,M) = \frac{\sum_i K_i^2 / M_i}{\sum_i K_i}$$

Regions that lacked data such as the beam stop area, had to be masked out for the analysis. We used a common mask for all diffraction patterns since the size and shape of the mask would otherwise bias the distance metric. The mask used was the union of the masks of the individual patterns.

FIG. 2 shows the three-dimensional assembly of the diffraction patterns in the orientations recovered from the data and intensities properly scaled by the recovered fluence. The probability of achieving a full coverage of fourier space from 198 diffraction patterns is calculated in Supplemental Material to be 99.999991%. To verify this full coverage all slices were also assembled giving each a thickness of one Shannon pixel. The assembled space contained no uncovered regions meaning that the number of diffraction patterns was enough.

**PHASE RETRIEVAL**

Non-crystalline objects produce oversampled diffraction patterns from which phases can be directly recovered in an interactive process where two constraints are sequentially enforced. The first constraint is that the Fourier amplitudes have to be consistent with the collected data. The second constraint is to enforce a known upper size limit of the sample.

We use an advanced version of the above algorithm called the Hybrid Input Output (HIO) algorithm implemented in the Hawk software package and enhanced by a positivity constraint. The support was handled by a Shrinkwrap algorithm with the constraint to have a specific area. The result was refined with 1000 iterations of the Error Reduction (ER) algorithm.

The average Fourier error was 0.019 and the average real-space error was 0.0048. The reconstruction did not suffer from weakly constrained modes meaning that the missing information in the center of the diffraction patterns could be completely recovered. This conclusion is based on an analysis method described in. The iterative phase retrieval was repeated 200 times with independent random starting phases. Real-space error, Fourier-space error and UPGMA clustering (unweighted pairgroup method with arithmetic mean) show only one outlier. The average of the 199 successful and similar 3D reconstructions is shown in FIG. 3. No symmetry was imposed during the assembly of the 3D data set. Object symmetry was instead recovered from the measured diffraction data in the EMC process. The map reveals an asymmetric internal structure with a shift of density to one side of the particle along a pseudo-five-fold axis.

The resolution is estimated from the phase-retrieval transfer-function (PRTF) which gives a full-period resolution of 125 nm (FIG. 4a). As expected, the PRTF drops where the signal is low. This behaviour also explains the oscillating nature of the PRTF that is common for nearly spherical objects.

**VALIDATION OF ORIENTATION RECOVERY**

Since this is the first 3D reconstruction from experimental data using the EMC algorithm, there were no validation methods available to assess the quality of our 3D orientation recovery. We have therefore developed two independent validation methods inspired by cryo-EM and X-ray crystallography.

A standard validation method in cryo-EM is to randomly split the data and analyse each set independently. A Fourier-shell correlation (FSC) is then calculated to quantify the differences between the two sets. We repeated the orientation and phase re-
FIG. 3. Reconstructed electron density. (a) The electron density of the mimivirus is recovered to a full-period resolution of 125 nm. The figure shows a series of iso-surfaces where blue represents denser regions and white represents lower density. The reconstruction shows a non-uniform internal structure and the line indicates the pseudo five-fold axis (b) A projection image of the recovered electron density. (c) A slice through the center of the recovered electron density.

retrieval independently on two disjoint sets of 99 randomly selected diffraction patterns. It can be shown that the 99 diffraction patterns cover reciprocal space with more than 99% probability. After assembly and phase retrieval, we compared the two electron density maps and plotted the FSC (FIG. 3b). Common thresholds for an acceptable FSC value range between 0.14 and 0.5 in cryo-EM literature. The FSC for our two reconstructions stays well above these values, even beyond the resolution according to the PRTF. This shows that the assembly of the 3D data was correctly performed and that the recovery of the phases was accurate. The results also indicate that the 198 mimivirus particles used in this experiment were identical to the resolution of the reconstruction.

In X-ray crystallography experiments, a subset of the recorded Bragg peaks are often excluded from the analysis and only used for validating the result. If the recovered structure matches the excluded data to a similar degree as to which it matches the included data, one can conclude that the structure is not overfitted to the data. Our second validation method is inspired by this analysis but it differs from it in two important ways: (i) entire diffraction patterns are excluded from the analysis instead of single Bragg peaks; (ii) the comparison is made between the omitted pattern and the recovered data.
intensity distribution, based on the similarity function in equation (1). We excluded 10% of the diffraction patterns from the analysis and calculated the similarity function given in equation (1) for both the data included in the assembly, $L$, and for the data excluded from the assembly, $L_{\text{free}}$:

\[ L = \frac{1}{N_{\text{inc}}} \sum_{\{K_i; \text{inc}\}} \max_{\{M_j\}} L(K_i, M_j) \quad (3) \]

\[ L_{\text{free}} = \frac{1}{N_{\text{exc}}} \sum_{\{K_i; \text{exc}\}} \max_{\{M_j\}} L(K_i, M_j) \quad (4) \]

Here, $\{K_i; \text{inc}\}$ and $\{K_i; \text{exc}\}$ are the sets of the included and excluded diffraction patterns respectively and $N_{\text{inc}}$ and $N_{\text{exc}}$ are the sizes of these sets. $\{M_j\}$ is the set of all model slices of the expanded model and the function $L$ is given in equation (1).

We calculate the average $L$ and $L_{\text{free}}$ values for 20 different random sets of excluded diffraction patterns and plot it in FIG. [4] as a function of iteration numbers in the EMC process. In contrast to the crystallographic measures $R_{\text{free}}$ and $R_{\text{cryst}}$, high values of $L$ and $L_{\text{free}}$ indicate a good fit. The fact that $L$ and $L_{\text{free}}$ closely follow each other in FIG. [4] indicates that the data are not overfitted.

CONCLUSION

In this article we have shown experimentally that 3D imaging of reproducible non-crystalline biological particles can indeed be performed at FELs. This was possible through an adapted version of the EMC algorithm. In this demonstration experiment we selected diffraction patterns with high signal. We developed two validation methods for use in this type of analysis. The validity of our reconstruction is supported by both new methods as well as by validation through the phase retrieval transfer function.

The main factor limiting the resolution is the small number of diffraction patterns that were available for this study. New experiments already provide much higher hit rates thanks to an improved sample injector.[6] This suggests that the resolution can be significantly improved in future applications. Also, further development of the EMC algorithm could allow for using individual masks for each diffraction pattern. This would make it possible to use patterns with large saturated regions that are currently thrown away, thus using more of the collected data.

There are many important reproducible biological objects with sizes of 30 - 300 nm. Three important pathogenic viruses, HIV, influenza and herpes are all in the 100 - 200 nm range. Furthermore, the EMC algorithm has been shown in simulation to be able to handle the much weaker signal strengths expected from single macromolecules or small viruses. It has been claimed that it is within the potential of free-electron lasers to image such objects at high resolution. This paper takes us one step closer to realizing this potential.

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[8] Marcus Gallagher-Jones, Yoshitaka Bessho, Sunam Kim,


