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# Emerging opportunities in structural biology with X-ray free-electron lasers

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X-ray free-electron lasers (X-FELs) produce X-ray pulses with extremely brilliant peak intensity and ultrashort pulse duration. It has been proposed that radiation damage can be 'outrun' by using an ultra intense and short X-FEL pulse that passes a biological sample before the onset of significant radiation damage. The concept of 'diffraction-before-destruction' has been demonstrated recently at the Linac Coherent Light Source, the first operational hard X-ray FEL, for protein nanocrystals and giant virus particles. The continuous diffraction patterns from single particles allow solving the classical 'phase problem' by the oversampling method with iterative algorithms. If enough data are collected from many identical copies of a (biological) particle, its three-dimensional structure can be reconstructed. We review the current status and future prospects of serial femtosecond crystallography (SFX) and single-particle coherent diffraction imaging (CDI) with X-FELs.

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## Introduction

The past decade has witnessed several important developments that potentially have significant impact on structural biology. First, the development of X-ray free-electron lasers (X-FELs) with extremely high peak intensity and ultrashort pulse duration allows data collection of biological samples with significantly reduced radiation damage. Radiation damage is presently a major factor limiting the attainable resolution in the imaging of biological material, in particular when using X-rays [1]. Initial radiation damage by X-rays in the energy range up to about 20 keV is dominated by the photoelectric effect. Upon absorption

of an X-ray photon, an electron is ejected, typically from an inner shell. The resulting core hole is filled by an outer shell electron. For biologically relevant elements such as C, N, O, S and P, the transition energy is transmitted primarily to another outer shell electron, which is then ejected as an Auger electron within a few to tens of femtoseconds (fs). Still on a femtosecond timescale, secondary electron ionization cascades ensue through electron-impact ionization. The removal of electrons and the formation of radicals lead to long timescale local electronic [2,3] and chemical changes [4,5] and will ultimately decrease the overall diffraction power of the sample. Cooling samples during data collection slows the long timescale damage since it reduces the mobility of the generated radicals. Protein crystals cooled to liquid nitrogen temperatures can tolerate a dose of ~30 MGy [6]. Solem predicted in 1986 that this dose can be significantly increased when using very intense X-ray pulses [7]. In this case, diffraction patterns with a diffraction-limited resolution can be captured before the obliteration of the sample by radiation damage [7]. With the emergence of X-FELs this concept was taken up by Hajdu and coworkers who calculated the pulse length dependence of the ionization induced atomic displacements and their effect on data quality using molecular dynamics simulations, suggesting that useable diffraction data can be obtained before explosion of the sample for very short FEL pulse lengths ('diffraction-before-destruction') [8\*\*].

Second, while X-ray crystallography is presently the primary methodology used for 3D structure determination of protein molecules at medium to atomic resolution, many biological systems are difficult to crystallize. In 1999, the methodology of X-ray crystallography was extended to allow structure determination of non-crystalline specimen and nanocrystals [9\*\*]. Compared to a regular crystal, the diffraction intensity from a non-crystalline specimen or nanocrystal is weak due to the loss or reduction of the amplification from the large number of unit cells in a sizable crystal. The diffraction pattern of a non-crystalline specimen or a nanocrystal can be sampled at an interval finer than the Bragg peak frequency, which in principle allows the direct phase determination by the oversampling method with iterative algorithms [10]. The combination of this methodology with X-FELs thus potentially opens a door to study biological systems without the requirement of sizable crystals.

## X-ray free electron lasers

X-ray free electron lasers are a new generation of X-ray sources that produce X-ray pulses with a peak brilliance

a billion times higher than synchrotron radiation and a pulse duration ranging from a few to hundreds of femtoseconds [11]. An X-FEL consists of a linear accelerator and a long undulator magnet. Intense electron bunches with low emittance are first accelerated to several to tens of gigaelectron volt (GeV) by the linear accelerator. The high energy electron bunches pass through the undulator with a sinusoidal magnetic field. When the electron bunches move along a sinusoidal, oscillating trajectory, they emit electromagnetic radiation. Initially because the size of each electron bunch is larger than the wavelength of the radiation, the radiation is incoherent and the intensity of the radiation is linearly proportional to the number of electrons in the electron bunch. If the undulator magnet is precisely tuned to match the phase and wavelength of the radiation, the radiation interacts with the electron bunch and the interaction accelerates part of the electrons and de-accelerates the others, such that the electrons form micro-bunches. The light emitted from these micro-bunches interferes constructively, resulting in an intensity proportional to  $\sim N^2$ , with  $N$  denoting the number of electrons in the micro-bunches. The increased intensity leads to an amplification of the micro-bunching process, which includes more and more electrons. This self-amplified process results in an exponentially growing intensity of the emitted light in the undulator until all electrons are part of a micro-bunch. At the end of the undulator, the electrons within micro-bunches produce X-FEL pulses with full transverse coherence, high peak intensity ( $\sim 10^{12}$  photons/pulse), and short pulse duration. The principle of this process is termed self-amplified stimulated emission (SASE) [12]. A more thorough review of the SASE X-FEL can be found elsewhere [11].

Presently, several soft and hard X-FELs are in operation worldwide, including the Free electron LASer in Hamburg (FLASH) [13], the Linac Coherent Light Source (LCLS) [14] at the SLAC National Accelerator Laboratory and the SPring-8 Angstrom Compact free electron Laser (SACLA) at the RIKEN Harima Institute in Japan [15]. In addition, the European X-FEL in Hamburg and the SwissFEL at the Paul Scherrer Institute in Switzerland are under construction and are expected to operate in 2016 and 2017, respectively. Furthermore, X-FEL facilities are under construction or planned in Italy, China and South Korea. Although SASE X-FELs are currently under rapid development worldwide, there are several issues associated with this kind of coherent X-ray sources. First, a SASE X-FEL is not a true laser due to the lack of temporal coherence [11]. Second, SASE starts with noise and is a stochastic process, which results in energy, intensity and position jitters of the X-ray pulses. As these issues will be resolved or alleviated by using different types of seeding technologies in the future [11], the present X-FELs with extremely intense and

ultrashort X-ray pulses already open exciting new opportunities in structural biology.

### Serial femtosecond crystallography

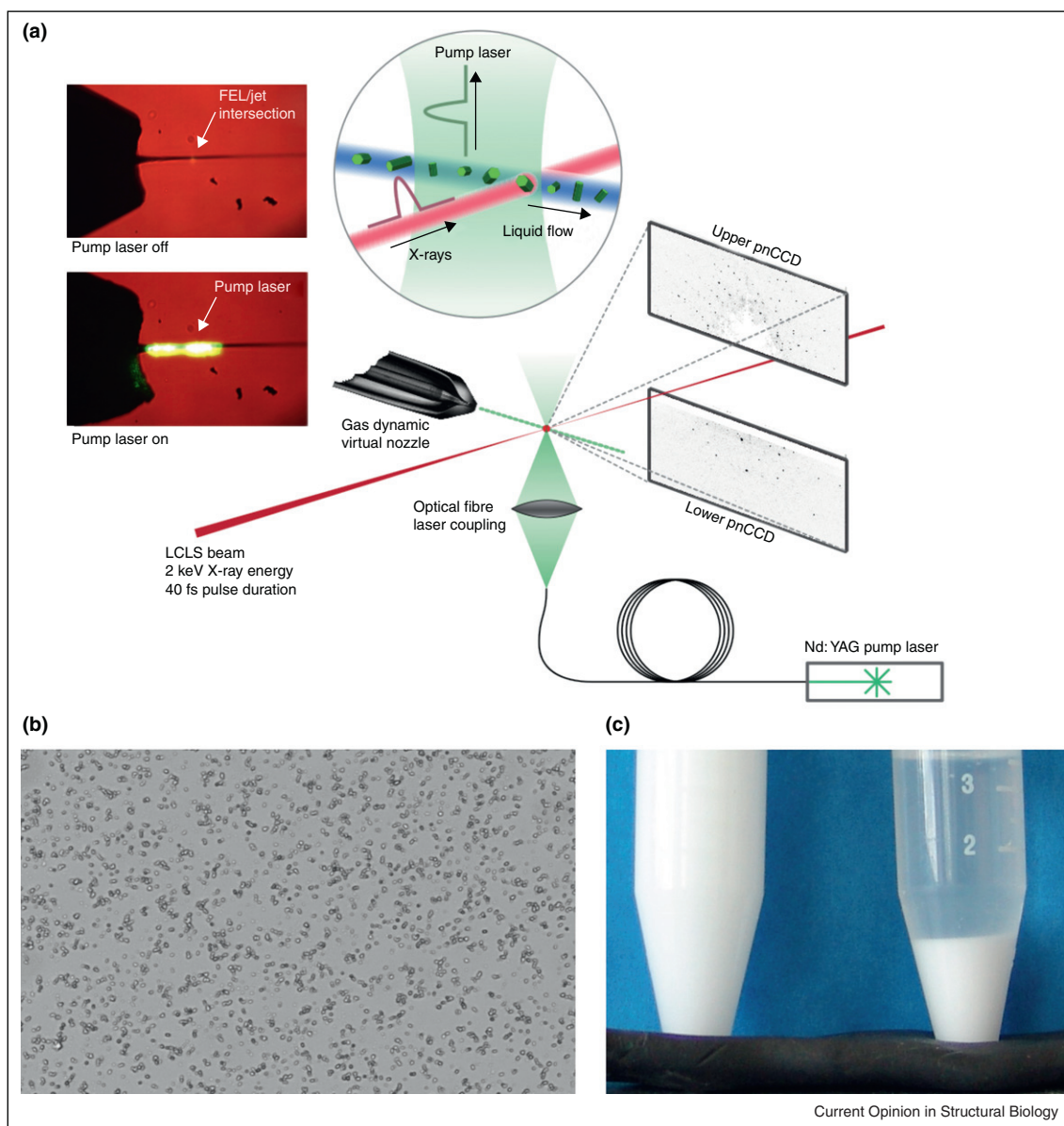
X-ray crystallography is the oldest and most successful method to determine high-resolution 3D structures of molecules of any size. Accordingly, the method's impact in biology and other areas of science is immense. A key feature of crystallography is the coherent addition of diffracted intensities of the molecules in the crystal into Bragg reflections with the absorbed dose being distributed among them. Thus, the resolution possible in a reconstructed image of the periodically averaged molecule in a crystal, for a given dose per molecule below the radiation damage threshold, depends on the number of molecules in the crystal [16]. This is the reason for the demand of large crystals for the collection of high-resolution data. For many systems, however, growing large, well ordered crystals is very difficult if not impossible. It is here that X-FELs are expected to make an impact in crystallography by allowing using tiny<sup>1</sup> crystals for data collection, beyond the conventional radiation damage limits [17\*\*].

Indeed, in a pioneering experiment, it has been demonstrated recently that tiny crystals of photosystem I, a megadalton membrane protein complex, yield diffraction data that are useful for structure analysis [17\*\*] (reviewed last year [18] in this journal). The experiment was performed at the LCLS at the AMO beamline [19] using the CAMP instrument [20\*] that houses two pairs of fast-readout large-area pnCCD detectors, positioned to capture both high-angle and low-angle diffraction. A gas-focused liquid microjet [21] was used to inject randomly oriented photosystem I microcrystals and nanocrystals into the X-FEL interaction region. There, they intersected with 300 fs long X-ray pulses with 1.8 keV photon energy. Since the crystals were destroyed upon exposure to a single femtosecond FEL pulse, the data were collected in a serial fashion, with the liquid jet providing a fast, gentle, and convenient means of replenishment as shown in Figure 1(a). Serial femtosecond crystallography (SFX) data were integrated using a Monte Carlo-like approach [22,23] which resulted in interpretable electron density of photosystem I, using the known structure for phasing. The resolution was limited to 8.5 Å by the wavelength of 6.9 Å and the geometry of the experimental setup [17\*\*].

High-resolution SFX data collection became feasible in February 2011 with the commissioning of the Coherent X-ray Imaging (CXI) instrument [24] at LCLS, providing access to 10-keV photons (1st harmonic). The prospects of serial femtosecond crystallography for high-resolution data collection and structure analysis were explored using

<sup>1</sup> 'Tiny' is used here as smaller than a few microns.

Figure 1



**(a)** Experimental setup for serial femtosecond crystallography. Tiny crystals are injected into an X-FEL beam using a liquid microjet. This setup is very convenient for pump-probe experiments using laser excitation. Shown is an experiment on the light-induced dissociation of photosystem I-ferredoxin cocrystals, taking place on a microsecond time-scale (from Ref. [43]). **(b)** Microcrystalline sample for serial femtosecond crystallography data collection. Lysozyme crystals ( $\leq 1 \mu\text{m} \times 1 \mu\text{m} \times 3 \mu\text{m}$ ) as seen through a conventional light microscope. **(c)** Microcrystalline lysozyme suspended in solution (left) and after settling (right) (from Ref. [42], reproduced with permission from the IUCr (<http://dx.doi.org/10.1107/S0021889812024557>)). It is important to note that the quantities depicted here are by no means excessive; several milliliters of very dense crystalline slurry are currently required for SFX experiments.

hen egg-white lysozyme as a well-characterized model system. SFX data were collected to 1.9 Å resolution of micrometer-sized lysozyme crystals (Figure 1(b)) using 40 fs and 5 fs X-ray pulses of 9.3-keV photons [25<sup>••</sup>]. Both SFX data sets scale reasonably with very low dose synchrotron data collected of crystals kept at room temperature. The SFX data yield clear molecular replacement solutions, the derived electron density maps show the

expected features, and the refinement statistics are similar to the synchrotron data. This establishes SFX as a viable and valuable complement to existing macromolecular crystallography techniques for the analysis of tiny crystals [25<sup>••</sup>].

Molecular replacement [26] has been used for phasing SFX data [25<sup>••</sup>,27<sup>\*</sup>]. So far, no *de-novo* phasing has been

demonstrated. Recently, the MAD method [28,29] has been theoretically generalized for phasing SFX data at high X-ray intensity by taking into account the detailed electronic damage dynamics of heavy atoms [30<sup>\*</sup>]. Although experimental results are needed to confirm the feasibility of this proposed generalization, in particular since the calculation only included the ionization of the heavy atoms, it may offer an *ab initio* structure determination method for SFX with X-FELs.

### Growing, detecting, and testing tiny crystals

Given the limitations imposed by radiation damage combined with signal-to-noise considerations, the emphasis has been on optimizing crystallization conditions that yield only very few large crystals instead of showers of small ones. Although finding initial crystallization conditions is largely driven by trial and error, that is, testing many different conditions, often using commercially available crystallization screens, once crystallization parameters have been identified, crystallization can be rationalized by a phase diagram that shows where liquid, crystalline or amorphous precipitate states are stable [31]. Crystallization involves two steps, nucleation and crystal growth. Nucleation takes place in a region of moderate supersaturation, while crystal growth occurs under conditions of lower supersaturation just below the nucleation zone where crystals are stable. Thus, for conventional large crystal growth, conditions are sought that lead the system into the so-called metastable zone where the concentration of the protein in the solution drops once nuclei have formed, resulting in crystal growth without the formation of further nuclei (see e.g. [www.emerald-biosystems.com/blog/post/protein-crystallization-phase-diagrams.aspx](http://www.emerald-biosystems.com/blog/post/protein-crystallization-phase-diagrams.aspx)). Therefore, to grow large amounts of tiny crystals, a higher degree of supersaturation has to be achieved, resulting in the formation of many nuclei in the 'labile zone' [32]. The most convenient way to achieve this is batch crystallization because of its simplicity, high reproducibility and scalability from a few microliters to hundreds of milliliters, allowing both screening and bulk preparation for the SFX experiments. Rapid and thorough mixing of precipitant and protein solutions can be essential for obtaining uniform crystal size distributions. Batch methods can also be used for lipidic sponge phase crystallization of membrane proteins for SFX as has been demonstrated recently for the photosynthetic reaction center of *Blastochloris viridis* [27<sup>\*</sup>].

Many macromolecules crystallize spontaneously if sufficiently concentrated. This can be achieved for example by centrifugation using centricons or by dialysis using appropriate molecular weight cut-offs. High local macromolecular concentrations can also occur in cells; indeed, microcrystals have been observed *in vivo*, providing eukaryotes and prokaryotes with a convenient and efficient way for storage or disposal [33]. Insect cells are interesting in this respect since many protein and virus crystals have been

observed *in vivo*. For example, polyhedrin forms a crystalline matrix coating the nuclear polyhedrosis baculovirus. Heterologous expression in the baculovirus Sf9 expression system under the polyhedrin promoter can be used for *in vivo* crystallization of proteins as has been shown for calcineurin [34] and recently for cathepsin from *Trypanosoma brucei* (TbCatB) and inosine monophosphate dehydrogenase from *T. brucei* [35<sup>\*</sup>]. In the case of TbCatB  $\sim 5 \times 10^5$  crystals were purified from  $10^6$  cells kept for 8 days in suspension culture. Since the crystals form inside the cell, they are limited in their size (10–15  $\mu\text{m}$  long and 0.5–1  $\mu\text{m}$  wide) and are very stable in regular buffers (pH > 4) and water. They were used for SFX experiments at the AMO beamline at the LCLS and diffracted to 8 Å resolution, limited by the geometry of the experimental setup and the available photon energy [35<sup>\*</sup>]. The advantages of *in vivo* crystallization are first, production of post-translationally modified proteins of interest, second, high stability and easy isolation by centrifuging the crystals after cell lysis, and third, a narrow crystal size distribution.

The identification of tiny crystals by light microscopy can be challenging, in particular when they occur as showers or in optically dense lipidic phases. *In situ* analysis of the crystallization setups exploiting UV fluorescence or birefringence may help in the identification of crystalline order as does the emerging application of Second-Order Non-linear optical Imaging of Chiral Crystals (SONICC) [36,37], which yields a strongly enhanced signal in the presence of chromophores. The latter two approaches, however, depend on the symmetry of the crystal lattice and the orientation of the crystals.

Electron microscopy is an excellent way to identify nanocrystals since it allows both imaging and diffraction studies, yielding information on both the size and crystallinity of the particles. However, sample transfer, in particular when using very small volumes, washing, blotting and, if desired, staining can be challenging especially when using crystallization solutions with high viscosity or high salt concentrations. Nevertheless, high throughput screening of 2D crystallization trials has been demonstrated recently [38] and there seems no reason that this cannot be transported to 3D crystals.

*In situ* (powder) diffraction is an option for screening tiny crystals [39], but the diffraction signal is likely not sufficient for nanoliter setups using standard beamlines. Nevertheless, powder diffraction does offer a convenient way of testing tiny crystals: crystalline slurries can be filled into quartz capillaries or mylar sleeves which are subsequently attached to conventional magnetic bases and stored in the associated plastic caps. Upon settling of the crystals, the pellet can be analyzed at room temperature by powder diffraction. Although the resulting diffraction pattern is unlikely to represent the diffraction

limit of a single crystal at an X-FEL, it demonstrates the crystallinity of the sample. Moreover, this approach can be used for optimization of crystallization conditions or of crystal storage solutions, similar to testing crystals at a home source before high-resolution data collection at a synchrotron. An important tool for improving the diffraction properties of large crystals is dehydration. At this point it is unclear whether similar approaches are applicable to tiny crystals.

### Practical aspects of serial femtosecond crystallography

Crystals of macromolecules require maintaining a certain hydration level during data collection. The liquid micro-jet [21] developed at Arizona State University allows keeping the sample in solution during injection and data collection which takes place in a vacuum. The crystalline liquid slurry is contained in a relatively large diameter inner capillary surrounded by a focusing sheath gas [40\*]. This design of a gas dynamic virtual nozzle (GDVN) provides both a means for reducing the probability of clogging of the capillaries while keeping the diameter of the liquid jet small (submicrometer to a few micrometer) and preventing of freezing of the jet [40\*].

The GDVN provides a stable liquid jet for a wide range of solution compositions before breaking up in droplets. Stable jets require typically sample flow rates of 5–25  $\mu\text{l}/\text{min}$  depending on the GDVN and solution composition, with the jet flowing at linear velocities of around 10 m/s. To achieve a high hit rate, defined as the fraction of detector frames collected containing a diffraction pattern of a crystal, while reducing the occurrence of double hits, the crystal concentration should be adjusted such that up to  $\sim 63\%$  of the X-ray interaction volume contains a crystal. With an FEL focus of  $1 \mu\text{m}^2$ , a diameter of the liquid jet of  $5 \mu\text{m}^2$  this corresponds to crystal densities of  $\sim 10^{11}/\text{ml}$ . For micrometer protein crystals with a density of  $1.35 \text{ g}/\text{cm}^3$  and 50% solvent content, this is equivalent to 130 mg/ml of protein. With a flow rate of 10  $\mu\text{l}/\text{min}$ , and an FEL repetition rate of 120 Hz (LCLS) or 10/30 Hz (SACLA), one out of 440,000 (LCLS) or 5.3 million/1.8 million (SACLA) crystals is hit by the FEL beam, emphasizing the need to develop methods that reduce sample consumption and high-lighting the importance of high repetition X-FELs, such as the European X-FEL. Possible solutions to decrease ‘unproductive’ sample consumption include approaches to slow the jet to an extent that the delivery rate of fresh crystals to the FEL interaction zone approaches the FEL repetition rate or to use fixed targets [41].

A high-performance liquid chromatography (HPLC) system can be used to supply the liquid pressure to deliver the sample into the nozzle. Such a setup also allows rapid switching between water and sample in case of clogging or problems with data collection, which can

save the sample. Recently, a propeller-mounted syringe pump was developed that minimizes sample settling (Figure 1(c)) during data collection by gently rotating a custom-made, temperature-controlled container storing the crystalline slurry [42]. The system is rated to 400 bar, allowing injection of very viscous samples, including various lipidic phases (e.g. [27\*]) used for membrane protein crystallization.

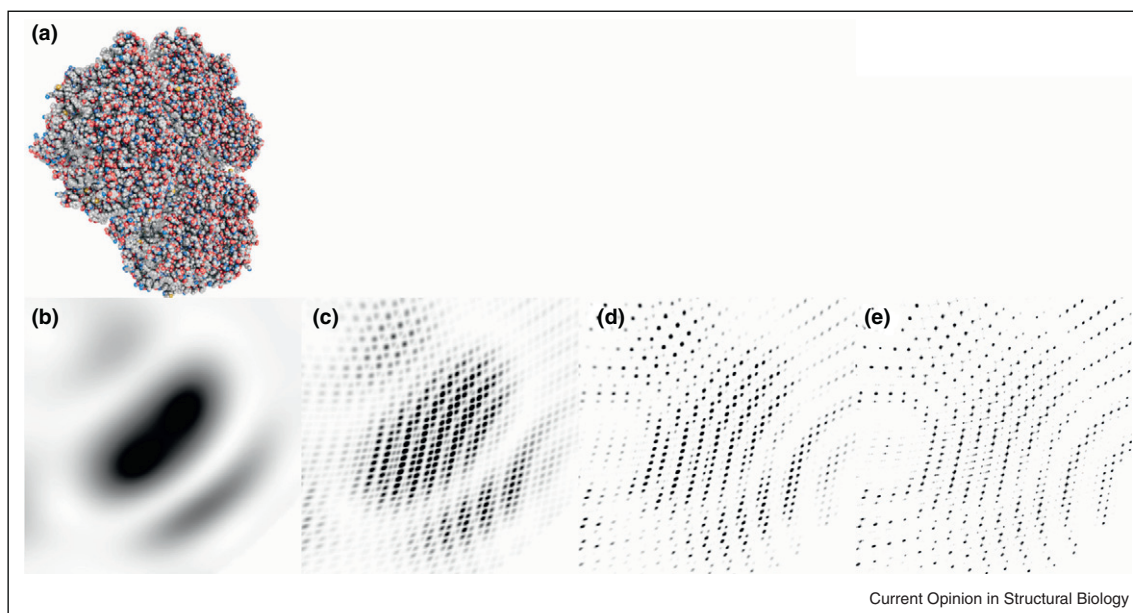
Crystals of macromolecules are characterized by a network of large solvent channels (solvent content typically 50%) allowing diffusion of small molecules to their binding sites in the crystallized macromolecule to induce chemical reactions or structural changes. Since diffusion times are significantly reduced for micron to submicron-sized crystals, they are very attractive for time-resolved mechanistic mixing experiments, using the liquid micro-jet. Because of their relatively low absorption, tiny crystals are also well suited for the study of light-induced reactions. A recent example is a pump probe experiment to study the light-induced dissociation of a ferredoxin–photosystem I complex on a microsecond timescale [43] (Figure 1a).

### Data evaluation in serial femtosecond crystallography

In SFX, the crystals are intersected by an X-FEL beam only fleetingly, and each exposure corresponds to a still image, representing a thin slice of the rocking curve of the Bragg reflections. The CrystFEL software suite [44] has been developed to analyze SFX data, using established procedures for indexing and a Monte Carlo like integration averaging out variations in crystal size and orientation and incident beam intensity when the sampling of the reflections is high enough [22\*]. Depending on the space group of the crystals, the independent indexing of the collected diffraction patterns may result in an equal probability of choosing one of multiple indexing possibilities, for example in the case of polar space groups (TA White *et al.*, Crystallographic data processing for free-electron laser sources, J Appl Crystallogr, 2012, Unpublished data). Therefore, the resulting dataset will appear to have higher symmetry than the actual crystal lattice and display perfect ‘twinning’. This feature complicates structure determination. The possible ambiguity in indexing is an open issue in data evaluation. The inclusion of profile fitting and post refinement is currently pursued by several groups. It will not only improve data quality but also likely reduce the number of diffraction patterns required for the convergence of the integrated reflection intensities, and thus the amount of sample required to collect high quality data sets.

For very small crystals, the assumption of an infinite lattice is no longer valid. This has direct consequences for the observed diffraction pattern which corresponds to the Fourier transform of the convolution of the electron

Figure 2



Simulated diffraction patterns of photosystem I (a). (b) The molecular transform of the molecule is continuous. The interference of several molecules arranged in a lattice of  $2 \times 2 \times 2$  (c),  $4 \times 4 \times 4$  (d) and  $8 \times 8 \times 8$  (e) results in the appearance of Bragg peaks and fringes between the Bragg peaks.

density distribution with the crystal lattice. In case of a finite lattice the intensities between Bragg peaks do not cancel out. Instead, each Bragg peak contains the Fourier transform of the crystal shape (shape transform) [17,45–47] and  $N-2$  interference fringes are present between adjacent Bragg peaks with  $N$  denoting the number of unit cells along the particular lattice direction (Figure 2). Currently, none of these finite-lattice features are taken into account in SFX data evaluation, but once they can be accurately measured, these features bear the potential for direct phase determination [45,47].

So far, 2D crystals have been within the realm of electron diffraction but this may change with the advent of X-FELs. Indeed, SFX diffraction data have been collected at the LCLS of beef catalase crystals which are typically used to calibrate transmission electron microscopes. The expected diffraction features were observed, with rod-like diffraction in one direction.

### Radiation damage analysis in serial femtosecond crystallography

The SFX experiment on photosystem I was performed using X-ray pulses of up to 300 fs duration. Strong Bragg peaks were observed at the edge of the detector, corresponding to 8 Å resolution [17\*\*]. Even when taken into account that the liquid jet surrounding the crystals may act as a tamper [48], this may seem at odds with predictions based on plasma models and hydrodynamics codes that predict 5 Å movements of the

generated ions in less than 100 fs. Thus, follow-up studies were performed on microcrystals of photosystem I [49\*] and lysozyme [50\*] to analyze the pulse-length dependence of radiation damage. For lysozyme, diffraction data were collected using nominal pulse durations of 70, 78, 85, 100, 150, 200, 250, 300 and 400 fs, with absorbed dose up to 3 GGy, two orders of magnitude higher than the conventional damage limit for protein crystals kept at cryogenic temperature during data collection [6]. To examine the quality of the X-FEL data, virtual powder patterns were computed by summing up the individual peaks and compared with conventional synchrotron powder patterns of identically prepared crystals. The correlation coefficients between intensities extracted by curve fitting from the X-FEL data and conventional powder data were >90% for all pulse lengths investigated, and all datasets can be used for structure determination by molecular replacement.

Radiation damage in SFX differs substantially from that observed in regular crystallography due to the difference in time-scales. The high irradiance of an FEL pulse results in very rapid formation of a hot electron gas by impact ionization, and impulse rearrangement of atoms and ions. The Coulomb repulsion of the ions and the rapid rise in electron temperature of the system causes displacement of both the atoms and ions during the pulse [51]. This heating leads to a high pressure that drives the explosion of the sample. The atomic ionization and rearrangement of atoms, ions and electrons manifests

itself by degradation of Bragg peak intensity and increase of diffuse scattering [52]. To analyze the effects of the X-ray pulse length on the data quality, all lysozyme SFX data sets were compared with the 70 fs one, which is assumed to be the least effected by radiation damage. A decrease of the intensities at high scattering angles was observed for pulse lengths longer than 100 fs. Given the stochastic nature of the experiment and the approximately Gaussian beam intensity profile which was significantly larger than the crystals, some crystals were in the center of the X-ray beam when exposed to pulses with high fluence, resulting in strong diffraction, whereas others were only grazed, or exposed to low fluence pulses, resulting in weaker diffraction. Since the lysozyme crystals had a uniform size distribution, diffraction intensity and dose can be correlated. By sorting the X-FEL data according to their relative intensity, it became apparent that the highest-intensity data show the strongest decrease of Bragg-intensity as a function of resolution with increasing pulse duration. Both dose and dose-rate dependent damage was observed even in the 70 fs data set.

When an X-FEL pulse interacts with a protein crystal, both time-dependent losses of electrons due to photo-ionization, Auger decay and impact ionization, and time-dependent uncorrelated disorder in the crystalline lattice take place. This has been modeled for events distributed evenly in the unit cell and a correction factor has been deduced [49<sup>o</sup>]. In order to understand whether non-uniform modifications of the electron density in the unit cell take place that cannot be corrected, further higher resolution studies are necessary [50<sup>o</sup>].

### Phase determination by the oversampling method

X-ray crystallography can nowadays be routinely used to determine the 3D structure of macromolecules, as long as sizable, well-ordered crystals are obtained. This remarkable achievement is partially attributed to the availability of powerful phase determination techniques such as direct methods [53], isomorphous replacement [54], multiple wavelength anomalous dispersion (MAD) [28,29] and molecular replacement [26]. However, when a crystal becomes very small or only has one unit cell (i.e. no crystallinity), a different class of phasing techniques exists, denoted the oversampling method. Partially based on the observation made by Boyes-Watson, Davidson and Perutz on haemoglobin [55], Sayre suggested in 1952 in his half-page paper that measuring the diffraction intensities at as well as in-between Bragg peaks may provide the phase information [56]. Following Sayre's idea, Bates proposed an explanation of the oversampling method in 1982 [57]. On the basis of the argument that the auto-correlation function of any function is exactly twice the size of the object itself, he concluded that the phase information can be recovered if and only if the diffraction

intensities are sampled at least twice as fine as the Bragg peak frequency.

In 1998, Miao *et al.* proposed a different explanation of the oversampling method and concluded that Bates' statement is inaccurate [58]. In their explanation, solving the phase problem is equivalent to solving the unknown electron density from a set of non-linear equations where each diffraction intensity point is considered as a non-linear equation. When the diffraction intensities are sampled at the Bragg peak frequency, the number of unknown variables is exactly twice the number of independent equations, where the independent equations are defined as those diffraction intensity points without any crystallographic or Friedel symmetry. When the diffraction intensities are sampled at an interval finer than the Bragg peak frequency, the number of independent equation is increased but the number of unknown variables remains the same. Equivalently, sampling the diffraction intensities finer than the Bragg peak frequency corresponds to surrounding the electron density with a 'no-density' region. The higher the sampling frequency, the larger the no-density region. To quantify the sampling degree, an oversampling ratio was introduced [58], defined as

$$\sigma = \frac{\text{volume of electron density region} + \text{volume of no-density region}}{\text{volume of electron density region}} \quad (1)$$

When  $\sigma \geq 2$ , the number of independent non-linear equations is larger than or equal to the number of unknown variables, and the phases can, in principle, be recovered from the diffraction intensities. However, it is a requirement but not a guarantee of a unique solution to have the number of independent equations larger than the number of unknown variables. By using the theory of polynomials, it can be shown that, while a finite number of multiple solutions may exist in one dimension, they are pathologically rare in two and three dimensions [59]. Physically, oversampling the diffraction intensities requires better spatial and temporal coherence of the incident beam than the Bragg peak sampling [60,61]. This is because the higher the oversampling frequency is, the finer the recording of the features in the diffraction pattern has to be. This explanation was confirmed by both numerical simulation and experiments [58,61].

Oversampling the diffraction intensities with  $\sigma \geq 2$ , in principle, makes the phases uniquely encoded in the intensities. However, it remains a challenging task to solve a large number of non-linear equations. A very effective way to find the solution (i.e. global minimum) is to use Fourier-based iterative algorithms. In 1972, Gerchberg and Saxton developed an iterative algorithm for retrieving the phases in electron microscopy [62]. The algorithm iterates back and forth between real and reciprocal space in which an electron micrograph in real space

and its diffraction pattern in reciprocal space are used as constraints. In 1978, Fienup further improved the iterative algorithm by using the finite size of an object (i.e. support) and positivity as constraints in real space [63]. Emphasis should be made that oversampling and support are two related, but different concepts. When the diffraction intensities are sampled at an interval slightly finer than the Bragg peak frequency, a support exists in real space, but the phases are not unique. The unique phases exist only when the sampling frequency is sufficiently fine with  $\sigma \geq 2$ . To date, several Fourier-based iterative algorithms have been developed, including error reduction [63], hybrid input–output [63], difference map [64], shrink-wrap [65], guided hybrid input–output [66], and relaxed averaged alternating reflections [67]. Each iteration in the Fourier-based iterative algorithms typically consists of the following five steps.

- (i) The measured Fourier magnitudes are combined with the phases obtained from previous iteration. In the first iteration, a random phase set is usually used.
- (ii) Applying the inverse fast Fourier transform (FFT), a new electron density map is calculated.
- (iii) A support is defined based on the oversampling ratio [58]. The electron density outside the support and the negative electron density are slowly pushed close to zero [63].
- (iv) Applying the FFT to the modified electron density map, a new Fourier transform is calculated and its phases are used for the next iteration.
- (v) An error metric is defined based on either the difference between the measured and calculated Fourier magnitudes or the electron density outside the support divided by that inside the support.

While the oversampling method with iterative algorithms works well for simulated data, the experimental realization of this method faced several obstacles: first, the high dynamic range of the diffraction intensities (typically spanning a few orders of magnitude), second, the missing data problem (i.e. usually the low frequency data cannot be measured due to the use of a beamstop), and third, the presence of different sources of noise in the diffraction pattern. It was not until 1999 that Miao and colleagues performed the first demonstration experiment and established the coherent diffraction imaging (CDI) method [9\*\*]. CDI has since been applied to a broad range of biological samples and inorganic materials [10,46,68,69]. The application of the oversampling method to phasing nanocrystals has also been pursued both theoretically [45,47] and experimentally [46,70]. By measuring intensities around the Bragg peaks, Robinson and colleagues were able to determine the 3D shape and the strain field of nanocrystals [70]. Although the oversampling method has been widely applied to determine the structure of non-crystalline specimens and the strain field of inorganic nanocrystals, it has not been

demonstrated experimentally for *ab initio* phasing of a 3D structure.

### Single-particle coherent diffraction imaging

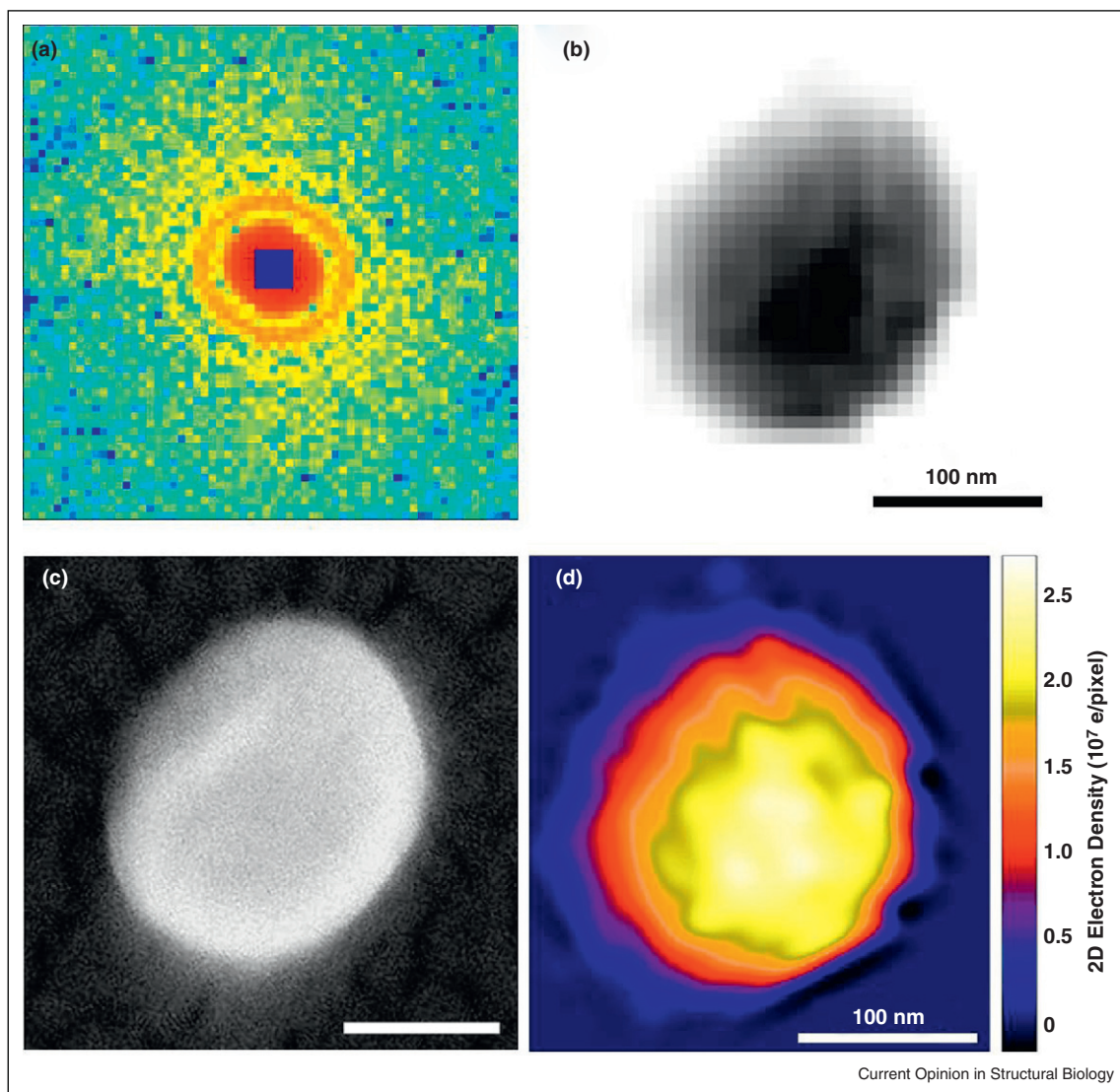
One of the most challenging applications of X-FELs is arguably the potential of imaging single large protein complexes in three dimensions. Unlike X-ray crystallography, single-particle CDI faces several obstacles. First, the X-ray diffraction signal from a single biomolecule is extremely weak which makes data acquisition a challenging task. Second, in order to enhance the signal-to-noise ratio and to obtain 3D structure information, the diffraction patterns of a large number of ‘identical’ particles with different orientations have to be measured. The heterogeneity of biomolecules poses a challenge. Third, compared to electrons, the interaction cross section of X-rays with atoms is about five to six orders of magnitude lower. This explains why cryo electron microscopy (cryo-EM) has been widely used to image large protein complexes and virus particles for 3D structure determination [71]. However, compared to electrons, X-rays have a much higher penetration depth and are not limited in their intensity by charging effects. Furthermore, X-FELs can provide intense X-ray pulses with  $\sim 10^{12}$  photons per pulse. When the X-ray pulses are shorter than tens of femtosecond, the diffraction pattern of a biomolecule can, in principle, be measured before the molecule is destroyed [8\*\*]. Follow-up studies suggest that in order to limit average atomic ionization in a biomolecule, X-FEL pulses may have to be shorter than 2–4 fs [72].

Over the past few years, a number of numerical simulations have been performed to explore the feasibility of single-particle CDI. It was first demonstrated through numerical simulations in 2001 that a molecular diffraction pattern at 2.5 Å resolution accumulated from  $3 \times 10^5$  copies of single rubisco biomolecules each generated by a femtosecond-level X-FEL pulse can in principle be successfully phased and transformed into an electron density map comparable to that obtained by more conventional methods [73\*]. In this study, however, the molecular orientation was assumed to be determined by other methods such as those developed in single-particle cryo-EM [71]. In subsequent years, several groups have developed more advanced methods to classify the weak diffraction patterns and then determine the molecular orientations based on the diffraction patterns [74–80]. This is an important step as the diffraction intensity of a single biomolecule from a single X-FEL pulse is very weak, and the signal to noise ratio of most detector pixels is likely much smaller than 1. After the classification and orientation determination, the 2D diffraction patterns can then be aligned, averaged and assembled to obtain a 3D diffraction volume, which can be directly phased to obtain the 3D electron density of the molecule using the oversampling method with iterative algorithms [10].

To perform a single-particle CDI experiment, two different schemes have been implemented: one with fixed targets and the other with particle injectors. The fixed target experiment was first conducted on single murine herpesvirus-68 virions at SPring-8 [81\*\*]. A herpesvirus virion has an asymmetric tegument and envelope outside of the icosahedrally symmetric capsid composed of defined numbers of subunits. The unstained herpesvirus virions were chemically fixed by 3% glutaraldehyde and supported on 30 nm thick silicon nitride membranes. Individual virions were illuminated by a coherent X-ray beam of 5 keV photon energy, and the diffraction patterns were captured by a CCD detector. Figure 3(a) shows a coherent diffraction pattern of a single, unstained

herpesvirus virion; no effects of radiation damage were observed at 22 nm resolution. The square at the center of the diffraction pattern represents the missing center problem, which is confined within the centro-speckle and allows for consistent phase retrieval [82]. By using the oversampling method with iterative algorithms, the diffraction pattern was directly phased to obtain an image with a resolution of  $\sim 22$  nm (Figure 3(b)) which is in good agreement with the scanning electron microscope image of the same virion (Figure 3(c)). By measuring the incident and diffracted X-ray intensities, the electron density of the reconstruction can be quantified. Figure 3(d) shows the quantitative electron density of the herpesvirus virion, where the yellow region represents the viral

**Figure 3**



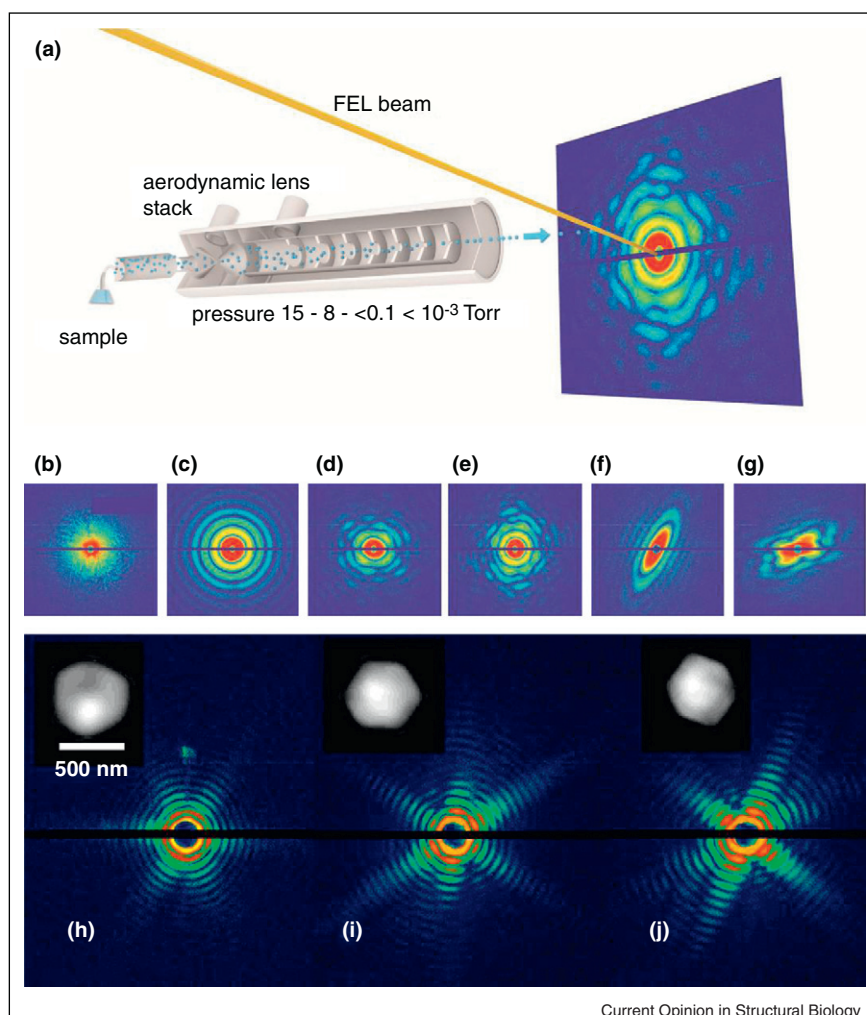
Single particle coherent diffraction imaging of herpesvirus virions. **(a)** Coherent X-ray diffraction pattern acquired from a single, unstained herpesvirus virion. **(b)** The diffraction pattern was directly phased to obtain an image with a resolution of  $\sim 22$  nm. **(c)** SEM image of the same virion. **(d)** Quantitative characterization of the reconstructed electron density map of the herpesvirus virion [from Ref. [81\*\*]].

capsid. Although this experiment was conducted at a synchrotron radiation source, the fixed target scheme of single particle CDI can be directly transferred to X-FELs.

The other scheme is to directly inject single particles into an X-FEL beam with an aerodynamic lens stack. Figure 4(a) shows a schematic layout of aerosolized particles injected into the X-FEL beam in random orientations [83]. The injected particles are intercepted by X-FEL pulses and the diffraction patterns are measured by a set of detectors. Figure 4(b–f) shows coherent X-ray diffraction patterns of a large aggregate, a water droplet, single T4 phage particles, a nanorice grain and two nanorice grains, measured with the LCLS pulses [83]. For virus particles smaller than 200–300 nm in diameter, it has been observed

that the size distribution of the particles estimated from the recorded diffraction patterns is larger than the actual particle size. This size increase is likely caused by aggregation of protein fragments, salts or a residual shell of ammonium acetate buffer around the virus particles. For larger virus particles such as mimiviruses, this effect has not been observed [84<sup>••</sup>], presumably because they have a similar size as the initial aerosol droplets. Figure 4(h–j) shows three representative coherent X-ray diffraction patterns measured from single mimivirus particles using LCLS pulses [85]. The LCLS pulses with photon energy of 1.2 keV and  $8 \times 10^{11}$  photons per pulse were focused to a spot of  $\sim 3 \mu\text{m}$  in diameter. The diffraction patterns exhibit symmetries reflecting the pseudo-icosahedral shape of the mimiviruses (Figure 4(h–j)). By using the

Figure 4



(a) Schematic layout of aerosolized particles injected into the X-FEL beam in random orientations. The injected particles are intercepted by the X-FEL pulses and the diffraction patterns are measured by a set of pnCCD detectors. Coherent X-ray diffraction patterns of a large aggregate (b) a water droplet (c), single T4 phage particles (d, e), a nanorice grain (f), and two nanorice grains (g), measured with the LCLS pulses. (h–j) Three representative coherent X-ray diffraction patterns measured from single mimivirus particles using LCLS pulses. The diffraction patterns were directly phased to obtain images with a resolution of  $\sim 32$  nm (insets) [from Refs. [83,85]].

oversampling method with iterative algorithms, the diffraction patterns were directly phased to obtain images with a resolution of  $\sim 32$  nm (insets in Figure 4(h–j)) [84<sup>\*\*</sup>,85]. In order to obtain the 3D structure, a series of diffraction patterns has to be measured from multiple identical copies of the particles. To date, a 3D reconstruction has been performed on nanorice particles in which 56 randomly oriented diffraction patterns were used [86,87].

Another potentially important direction is a newly developed CDI technique, denoted ankylography, which under certain circumstances can reconstruct the 3D structure of a small object (or a large object at low resolution) from its diffraction pattern sampled on the Ewald sphere [88<sup>\*</sup>,89,90]. Because of the size and resolution limitation, ankylography may be only applicable to certain classes of samples, but its advantage is the possibility of obtaining 3D structure without the requirement of averaging. The first ankylography paper has generated a huge debate in the scientific community. For those who are interested in ankylography, several ankylographic source codes have been posted on a public website and can be freely downloaded and tested (<http://www.physics.ucla.edu/research/imaging/Ankylography>). Finally, the Coherent X-ray Imaging Data Bank has recently been established ([www.CXIDB.org](http://www.CXIDB.org)) to make single-particle CDI data and reconstruction algorithms publicly available.

## Conclusions

X-FELs produce X-ray pulses with unprecedented peak flux and ultrashort pulse durations. The application of X-FEL radiation to structural biology has just begun, including the recent measurements of X-ray diffraction patterns from micron-sized protein crystals at near atomic resolution [25<sup>\*\*</sup>]. These experimental results suggest that the conventional radiation damage barrier in structure determination of biological systems can be significantly alleviated by using the ultra intense and short X-FEL pulses. Presently, however, several issues have to be resolved before serial femtosecond crystallography can be applied to routinely determine 3D structures using tiny crystals. First, sample consumption in serial femtosecond crystallography is currently a major bottleneck. It is thus an urgent need to develop sample delivery systems to reduce the sample amount required in the experiment. Second, *ab initio* phase determination methods need to be developed to phase the diffraction patterns in serial femtosecond crystallography. Third, indexing, post refinement and profile fitting have to be improved to reduce the number of hits required for structural determination. With the hard X-rays now available at LCLS and SACLA, great impact can be anticipated if increasingly smaller crystals can be used for structure determination, relaxing the requirements on crystal size. Smaller crystals will, however, demand for higher X-FEL flux to achieve sufficient signal levels and will be accompanied

by increased doses. This will bring the question whether radiation damage can be ‘outrun’ on the atomic level back into focus.

The other potentially important direction is the imaging of large protein complexes with X-FELs without the need of crystallization. The methodology of X-ray crystallography has been extended to allow imaging of non-crystalline specimens [9<sup>\*\*</sup>]. The continuous diffraction patterns from non-crystalline specimens allow the direct phase determination by the oversampling method with iterative algorithms [10]. With X-FELs diffraction patterns of single virus particles have been measured and then directly phased to obtain images [84<sup>\*\*</sup>,85]. In order to obtain high-resolution 3D structure information of single large protein complexes, there are several challenges that need to be overcome. First, the pulse fluence of X-FELs is not high enough to allow the measurement of high-resolution diffraction signals from single large protein complexes. This can, in principle, be overcome by improving the X-FEL peak intensity and using better focusing optics. Second, the dynamic range of the detectors presently used for single-particle imaging with X-FELs is  $\sim 10^3$ . This has to be increased by at least 1–2 orders of magnitude. Third, due to the weak signal from single particles, the background scattering has to be further reduced. Finally, sample preparation is critical for single particle imaging with X-FELs. It not only requires a very large quantity of identical particles, but also should enable to inject identical particles into an X-FEL beam without aggregation of protein fragments, salts or a residual shell of the buffer around the particles. As structural biology with X-FELs is a new, rapidly developing field, it is not surprising that many issues remain to be resolved and more developments need to be made. As more X-FELs worldwide start to operate within the next few years, it is safe to predict that the future of structural biology with X-FELs will be exciting, bright and also challenging.

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