

Atomic structure of granulin determined from native nanocrystalline granulovirus using an X-ray free-electron laser

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To understand how molecules function in biological systems, new methods are required to obtain atomic resolution structures from biological material under physiological conditions. Intense femtosecond-duration pulses from X-ray free-electron lasers (XFELs) can outrun most damage processes, vastly increasing the tolerable dose before the specimen is destroyed. This in turn allows structure determination from crystals much smaller and more radiation sensitive than previously considered possible, allowing data collection from room temperature structures and avoiding structural changes due to cooling. Regardless, high-resolution structures obtained from XFEL data mostly use crystals far larger than $1 \mu\text{m}^3$ in volume, whereas the X-ray beam is often attenuated to protect the detector from damage caused by intense Bragg spots. Here, we describe the 2 Å resolution structure of native nanocrystalline granulovirus occlusion bodies (OBs) that are less than $0.016 \mu\text{m}^3$ in volume using the full power of the Linac Coherent Light Source (LCLS) and a dose up to 1.3 GGy per crystal. The crystalline shell of granulovirus OBs consists, on average, of about 9,000 unit cells, representing the smallest protein crystals to yield a high-resolution structure by X-ray crystallography to date. The XFEL structure shows little to no evidence of radiation damage and is more complete than a model determined using synchrotron data from recombinantly produced, much larger, cryocooled granulovirus granulin microcrystals. Our measurements suggest that it should be possible, under ideal experimental conditions, to obtain data from protein crystals with only 100 unit cells in volume using currently available XFELs and suggest that single-molecule imaging of individual biomolecules could almost be within reach.

XFEL | nanocrystals | structural biology | bioimaging | SFX

Imaging of biomolecules using radiation of short enough wavelength to resolve individual atoms is limited by radiation damage, which destroys the very structure being measured. Energy absorption is unavoidable because the ratio of elastic scattering to damaging photoabsorption is an inherent property of atoms. Absorbed dose is therefore proportional to the scattered intensity, and thus the measured signal, used to determine the structure. The maximum tolerable dose that the sample can withstand fundamentally limits atomic resolution structure determination (1). The tolerable dose, and therefore damage, is a function of spatial resolution, with fine detail being destroyed first (2, 3). Radiation damage is typically overcome by distributing the dose over many identical copies of the

same molecule, either using crystals containing many aligned copies of the same molecule, or by aligning images of individual molecules in the case of single particle electron microscopy. At room temperature, free radical production after a dose as small as 150 kGy (gray = $\text{J}\cdot\text{kg}^{-1}$ absorbed energy) causes rapid decay of biological samples (4). This decay can be reduced by cooling the sample to below ~ 120 K. Doses up to about 30 MGy at synchrotron beam lines are routinely used for atomic structure determination from cryocooled protein crystals larger than about $100 \mu\text{m}^3$ (3, 5, 6). Similar dose limits apply in single particle cryo-electron microscopy, where individual images must have sufficient contrast to allow alignment before averaging to improve signal-to-noise (7, 8).

Serial femtosecond crystallography (SFX) overcomes radiation damage by using ultrabright X-ray pulses that are shorter in

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Data deposition: The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.pdb.org [PDB ID code 5G0Z (SFX structure) and 5G3X (SYN structure)].

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Significance

The room temperature structure of natively formed protein nanocrystals consisting of 9,000 unit cells has been solved to 2 Å resolution using an unattenuated X-ray free-electron laser (XFEL) beam, representing, by far, the smallest protein crystals used for protein structure determination by X-ray crystallography to date. Radiation damage limits structure determination from protein crystals using synchrotron techniques, whereas femto-second X-ray pulses from free-electron lasers enable much higher tolerable doses, extracting more signal per molecule, allowing the study of submicrometer crystals. Radiation-sensitive features, such as disulfide bonds, are well resolved in the XFEL structure despite the extremely high dose (1.3 GGy) used. Analysis of signal levels obtained in this experiment indicates that structure determination from even smaller protein crystals could be possible.

duration than most damage processes (9–14), thereby vastly increasing the tolerable dose and thus the amount of information gathered before the specimen is destroyed. Commonly, crystals in a liquid stream are injected into the X-ray beam, and each crystal diffracts for only a few femtoseconds before being destroyed (12, 15). Each crystal is exposed to at most one X-ray pulse, and diffraction patterns from thousands of crystals in random orientations are combined to form a complete 3D dataset. After indexing and integration, hundreds or even thousands of observations of each individual reflection from different crystals contribute to the final dataset through an averaging and modeling process resulting in a single set of reflection intensities for crystallographic structure determination (16, 17).

Early SFX experiments performed at the Linac Coherent Light Source (LCLS) showed that useful structural information could be obtained from submicrometer-sized crystals (10) using long-wavelength X-ray pulses (2 keV photon energy) at a dose of 700 MGy and from micrometer-sized crystals at an energy and dose similar to cryogenic synchrotron data collection (11, 18, 19) (9.4 keV, 33 MGy). Simulations (20, 21) and experiments (12–14) suggest that it should be possible to outrun radiation damage with short enough pulses using much higher X-ray intensities. Calculations with pulse durations as short as 10 fs predict the possibility of high-resolution structure determination from

nanocrystals at doses greater than 10 GGy, and 1-TGy doses should be tolerable with subfemtosecond pulses (22). In this context, it has been proposed that the combination of intense X-ray laser pulses and the serial data collection method could be used to obtain macromolecular structures by single-molecule diffraction (9, 23). Despite being able to measure submicrometer-sized crystals or single molecules, most SFX experiments carried out to date, where structures have been solved to better than 3.5 Å resolution, have used protein crystals of about 1–10 μm in diameter and the full power of the XFEL source must be attenuated to avoid saturating or damaging the detector (11, 18, 19, 24, 25), at least when performing experiments in vacuum using the Cornell-SLAC Pixel Array Detector (CSPAD) detector.

Here, we use unattenuated XFEL pulses to study native occlusion bodies (OBs) of *Cydia pomonella* granulovirus (CpGV), which are spheroidal ~265 × 265 × 445 nm semicrystalline particles that each contain a well ordered ~0.01 μm² protein lattice encasing a virion. Granulovirus OBs are stable, do not aggregate, and are readily obtainable because the virus is commercially used in horticulture to control codling moth (*C. pomonella*) in orchards (26). Granuloviruses belong to the *Betabaculovirus* genus of the *Baculoviridae* family, a group of insect viruses characterized by viral OBs, infectious microcrystals that form within infected larvae and persist for long periods in the environment (27).

OBs contain membrane-bound virus particles embedded in a crystalline lattice of viral protein molecules (Fig. 1). The OB matrix protein of *Betabaculovirus* (granulin) is homologous to that of *Alphabaculovirus* (polyhedrin), sharing ~60% amino acid sequence identity. Much remains to be learned about OBs, such as how the crystals grow within cells, what controls their size and shape, how the polyhedrin/granulin lattice interacts with the embedded virus particles and the envelope layer that forms the outer surface, and how the OBs disassemble in the alkaline midgut of feeding larvae to release the infectious virus. Such studies would ideally be performed at physiological temperatures and conditions on native OBs rather than recombinantly expressed protein recrystallized into large crystals.

Atomic structures for Lepidopteran *Alphabaculovirus* polyhedrins have been determined using native polyhedral OBs purified from larvae and recombinant microcrystals prepared by expressing polyhedrin in insect cells (28, 29). These revealed a cubic body centered lattice with a 103-Å unit cell densely packed with polyhedrin molecules interconnected by extensive crystal contacts and intermolecular disulfide bonds. The solvent content

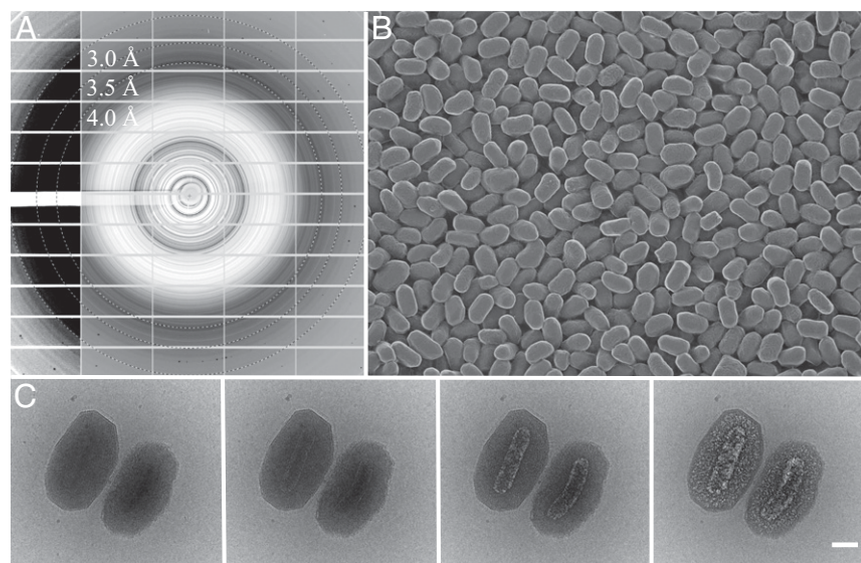


Fig. 1. Granulovirus OBs contain a single virion surrounded by a crystalline protein layer that diffracts to high resolution. (A) Powder X-ray diffraction from a pellet of granulovirus OBs at 100 K (*Materials and Methods*). Protein diffraction rings extend to a resolution between 3 and 3.5 Å. The detector panels on the left with enhanced contrast show evidence of diffraction at even higher resolution. Resolution rings are shown at 4, 3.5, and 3 Å. (B) Freeze etch electron micrograph showing the uniform size distribution of the particles (*Materials and Methods*). (C) Cryo-EM. The sequence of four 20 e/Å² exposures shows the effects of radiation damage on granulovirus OBs. The crystalline lattice is visible only in the first image and hydrogen gas bubbles produced by radiolysis eventually reveal the virion. (Scale bar, 100 nm.)

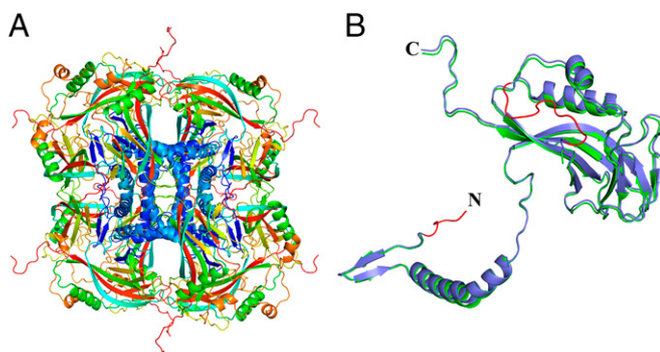


Fig. 3. (A) The structure of the biological unit of granulin building blocks forming the crystalline OB. (B) Granulin monomer. The SFX structure is displayed in blue and the SYN structure in green. Regions of granulin that were present in the SFX but not in the SYN structure are highlighted in red.

are well defined in the XFEL granulin structure (Figs. 3 and 4) but are completely absent in the WNPV polyhedrin structure.

Discussion

Using submicron-sized ($0.01 \mu\text{m}^3$) naturally occurring granulovirus OBs, where each particle contains only 9,000 unit cells, we have determined the room-temperature structure of granulin to 2-Å resolution using X-ray pulses that imparted a dose of up to 1.3 GGy per crystal. Despite the high dose applied and the small size of the crystals, we obtained a structure of CpGV granulin with good crystallographic statistics (Table 1), revealing features that are absent in the structure of recombinant granulin (Figs. 3 and 4). The final dataset contained 82,603 indexed diffraction patterns. Our analysis clearly shows that averaging over more diffraction patterns improves all figures of merit (Fig. S4), with an improvement of signal-to-noise ratio increasing with the square root of the number of patterns (Fig. 2C).

The 82,603 crystals contributing to the final dataset represent a maximum probed crystalline volume of less than $900 \mu\text{m}^3$, equivalent to a single cube-shaped crystal of $\sim 10 \mu\text{m}$ per side. Room-temperature measurement from a single large crystal of equivalent volume using conventional (rotation) data collection would be severely limited in resolution and completeness, because of radiation-induced damage accumulating over many exposures of a single crystal. Exposing many small crystals to femtosecond-duration X-ray pulses using serial crystallography is possible because of the tolerance of a much higher dose.

Given the high dose per crystal used in the SFX measurements, we looked for evidence of predicted local radiation damage (13, 14) in the vicinity of radiation-sensitive disulfide bonds. Fig. 4 shows that the disulfide bond between the two Cys135 molecules is largely intact in the SFX structure (occupancy 0.68), indicating that the structure is preserved even at sites of atoms that are strongly photo-absorbing, such as sulfur, despite the high dose used in the SFX experiment.

This 2-Å structure, obtained from $0.01 \mu\text{m}^3$ -size crystals, raises the following question: How small is the smallest crystal that could be studied at an XFEL facility? As the crystal size diminishes, so too does the intensity of the Bragg spots, and thus the size of the smallest crystal depends on the number of patterns that can be measured and indexed to build up a sufficiently accurate estimate of the integrated intensity. Assuming that indexing can be carried out, two other parameters that determine the ability to accurately estimate structure factors are the incident pulse flux density and the background level of the diffraction pattern (e.g., from scattering from the jet carrier liquid).

To further study this scaling, we simulated diffraction patterns from a perfect granulin crystal of 9,000 unit cells (*Materials and Methods*) and compared the average Bragg peak counts from individual diffraction patterns with our measured data. From this comparison we conclude that the incident flux density was of the order of 10^{12} photons per $1 \mu\text{m}^2$. A second simulation using the same beam parameters and zero background shows that a crystal of 123 unit cells would be sufficient to produce “indexable” diffraction patterns using the same dose of 1.3 GGy (Fig. S5) and that these data could be assembled into a complete high-resolution dataset given a sufficient number of measurements. Using the same beam parameters and focus size, it might therefore be possible to study crystals with 100 times smaller volume than those studied here, containing as few as 123 unit cells.

Going further, these rough estimates indicate that single-molecule imaging of a protein of molecular weight equal to one unit cell of granulin would require at least a 100–10,000-fold increase in X-ray peak flux density incident on a single unit cell to obtain continuous diffraction pattern with comparable numbers of scattered photons.

Additionally, single-molecule diffraction would require reducing the background scattering to much lower levels than achieved in serial crystallography measurements with a liquid microjet. Indeed, the goal is to achieve background-free injection, where photon-counting statistics are predicted to become the dominant source of error. Additionally, detector performance must be sufficiently well controlled to detect the presence of these particles and determine their orientation. A factor of 100 increase

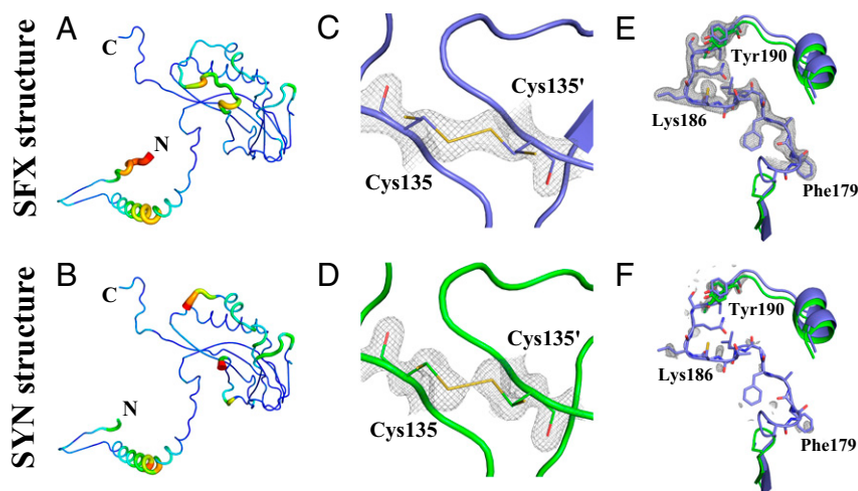


Fig. 4. Comparison of the final $2F_o - F_c$ electron density map and model for the SFX (blue) and SYN (green) structures of granulin. (A) SFX and (B) SYN model with color-map and thickness corresponding to local B-factor. (C and D) Close-up of Cys135 in (C) the SFX (blue) and (D) the SYN structure and (green) embedded in its electron (gray mesh, contoured 1.5σ). Cys135 adopts two distinct side chain conformations (shown as sticks): The major conformer forms an intermolecular disulfide bond with its counterpart in a neighboring molecule and has a refined occupancy of 0.68 and 0.87 for the SFX and SYN structure, respectively. (E) SFX and (F) SYN comparison of electron density for residues 168–201 (contoured at 1σ). Electron density defining residues 176–190 was only found for the SFX structure.

in X-ray intensity (which would impart a dose of 100 GGy) can in principle already be delivered in the 100-nm FWHM focus chamber of the CXI instrument under ideal conditions and perfect focus. Our simple scaling calculations indicate that at least a perfect focus, background-free measurements, and a noise-free detector capable of measuring single photons would be needed to perform single-molecule imaging of larger molecular weight macromolecules at LCLS, properties that are yet to be satisfied both individually and simultaneously.

Under the described limitations and assumptions, it should be possible to reach the border between crystallography and the field of single-particle X-ray imaging at ambient temperatures. Further increasing the flux density in parallel with improved signal-to-noise ratio of the data will open completely novel routes to structure determination of biological molecules at room temperature. The short pulses available from free-electron lasers allow structure determination from crystals much smaller or more radiation sensitive than previously considered possible for study by X-ray diffraction as well as time-resolved experiments on the ultrafast timescale.

Materials and Methods

A more detailed description of the sample preparation and diffraction experiments can be found in *SI Materials and Methods*.

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