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Beyond protein structure determination with MicroED

Chi Nguyen¹ and Tamir Gonen^{1,2,3}

Microcrystal electron diffraction (MicroED) was first coined and developed in 2013 at the Janelia Research Campus as a new modality in electron cryomicroscopy (cryoEM). Since then, MicroED has not only made important contributions in pushing the resolution limits of cryoEM protein structure characterization but also of peptides, small-organic and inorganic molecules, and natural-products that have resisted structure determination by other methods. This review showcases important recent developments in MicroED, highlighting the importance of the technique in fields of studies beyond protein structure determination where MicroED is beginning to have paradigm shifting roles.

Addresses

¹ Department of Biological Chemistry, University of California Los Angeles, 615 Charles E Young Drive South, Los Angeles, CA90095, United States

² Department of Physiology, University of California Los Angeles, 615 Charles E Young Drive South, Los Angeles, CA90095, United States

³ Howard Hughes Medical Institute, University of California Los Angeles, Los Angeles, CA90095, United States

Corresponding author: Gonen, Tamir (tgonen@g.ucla.edu)

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Introduction

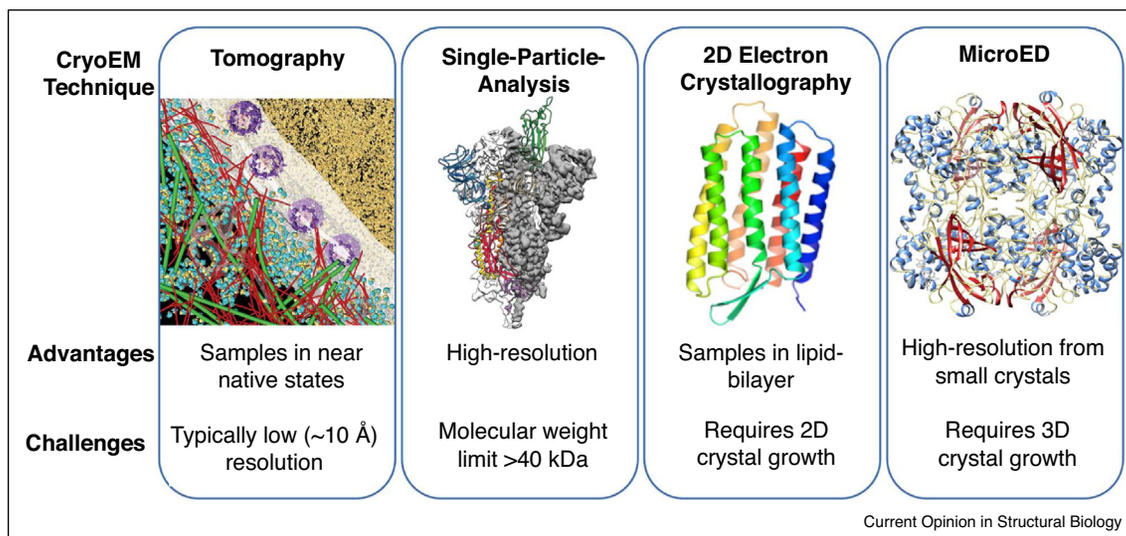
MicroED [1[•],2^{••}] has been pushing the limits of cryoEM in determining structures of macromolecular protein assemblies, peptides, and chemical compounds [1[•],3,4^{••},5[•]]. Prior the technological advancements in detectors [6–9] and software [10–13] that made the ‘cryoEM resolution revolution [14]’ possible, structure determination of biological assemblies, peptides, and chemical compounds has been dominated X-ray crystallography. To date, there are close to 150 000 depositions in the Protein Data Bank (PDB), comprised of roughly 90% X-ray, 8% NMR, and 2% EM structures. The first near atomic resolution structure by cryoEM reported in 2005 [15] set the stage for the highest growth in transmission electron microscopy (TEM) structure deposition that occurred in the last five years.

The field of cryoEM includes at least four major techniques: cryo-electron tomography (cryoET) [16,17], single-particle-analysis (SPA) [18–20], 2-dimensional (2D) electron crystallography [21,22], and MicroED [1[•],2^{••}] (Figure 1). All of these cryoEM techniques exploit the advantage that electrons interact orders of magnitude more strongly with materials than X-rays, allowing the application of samples that are not tractable by other methods [23]. While CryoET and SPA use imaging, the crystallographic cryoEM methods of 2D electron crystallography and MicroED also take advantage of electron diffraction. 2D-electron crystallography is typically used for structure determination of molecules in 2D arrays, which traditionally have been of membrane proteins that are crystallized within the native environment of the lipid bilayer [22]. In contrast, MicroED uses 3D crystals and data collection by continuous rotation to yield structures of a wide range of samples including soluble and membrane proteins, peptides, small organic and inorganic molecules, semi-conductors, and natural-products [5[•]].

During MicroED experiments, crystals are harvested and prepared in a number of ways for embedment on EM grids (discussed below) [1[•]] (Figure 2). Biological samples, which are more sensitive to radiation, are typically vitrified to protect from radiation damage and to withstand the high-vacuum within the electron microscope [1[•],27]. Once well-diffracting crystals are detected, MicroED datasets are collected by exposure of the sample to an electron beam in diffraction mode during continuous rotation of the stage [2^{••}] (Figure 2). MicroED data are then collected on a fast camera as a movie, where each frame contains a diffraction pattern that represents a wedge of the reciprocal space [27]. Because continuous rotation for MicroED is analogous to the rotation method in X-ray crystallography, the data collected can be directly processed by existing standard X-ray crystallography software such as Mosflm [28], XDS [29], DIALS [30], SHELX [31] and HKL2000 [32] (Figure 2).

After data-processing, the phases are determined and structures are built using the electron density maps [1[•]]. Given its wide-application and ability to extract structural information from nanocrystals, often a billionth the volume of those needed for X-ray crystallography, there are a growing number of structures determined by MicroED. Since its inception in 2013, there are close to 100 PDB entries produced by MicroED, with the highest growth in the just the past two years. To date, several laboratories have published MicroED studies and the number of practitioners are growing. Still, there are several challenges that lay ahead for MicroED including additional methodologies for sample preparation and technological developments for

Figure 1



The four major modalities of cryoEM.

From left to right. A model of the HeLa cell nuclear periphery by Cryo-ET (Reprint from Ref. [24] with permission from AAAS). Model of COVID-19 spike protein by SPA (Reprint from Ref. [25] with permission from AAAS). The structure of bacteriorhodopsin determined by electron crystallography (Reprint from Ref. [21]). The structure of catalase determined by MicroED (Reprint from Ref. [26]).

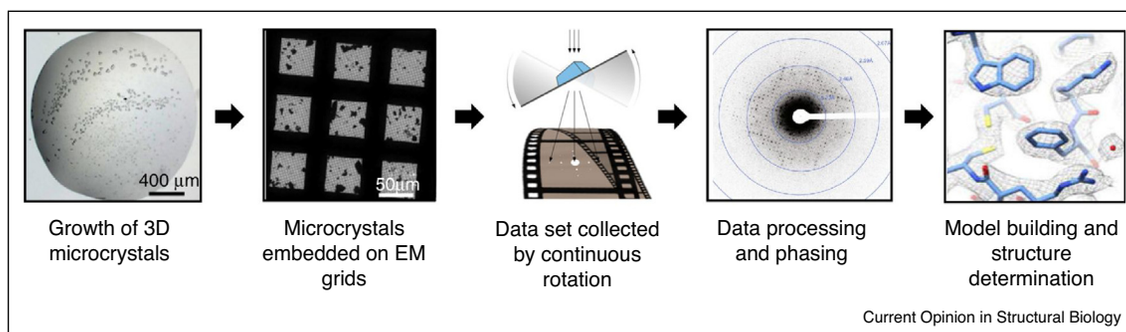
data collection that currently limit widespread usage of the technique. Below we discuss the most recent developments and strategies to expand the use of the MicroED.

Protein structure determination by MicroED

Formation of large crystals continue to be the most challenging and time-consuming step for X-ray crystallography, especially for membrane proteins and protein complexes [33]. The small crystals that are typically formed by membrane proteins and protein complexes can often diffract electrons using very low exposures to minimize radiation damage ($0.01 \text{ e}^-/\text{Å}^2$) [1]. One of the earliest membrane protein structures determined by

MicroED was the Ca^{2+} ATPase (PDB 3J7T/U) [34] (Figure 3a). The Ca^{2+} ATPase structure illustrated the utility of MicroED for generating Coulomb potential (charge density) maps to detail information about the charged-states of amino-acid sidechains, cofactors, metals, and ligands [34]. Since Ca^{2+} ATPase, there have been several important structures determined by MicroED including the non-selective sodium-potassium (NaK) channel (PDB 6CPV) [35] and the complex of the transforming growth factor beta paired type II (TGF- β m:T β RII) (PDB 5TY4) [36] (Figure 3a,b). The MicroED structure of NaK is similar to those previously determined X-ray crystallography [37]. However, like Ca^{2+}

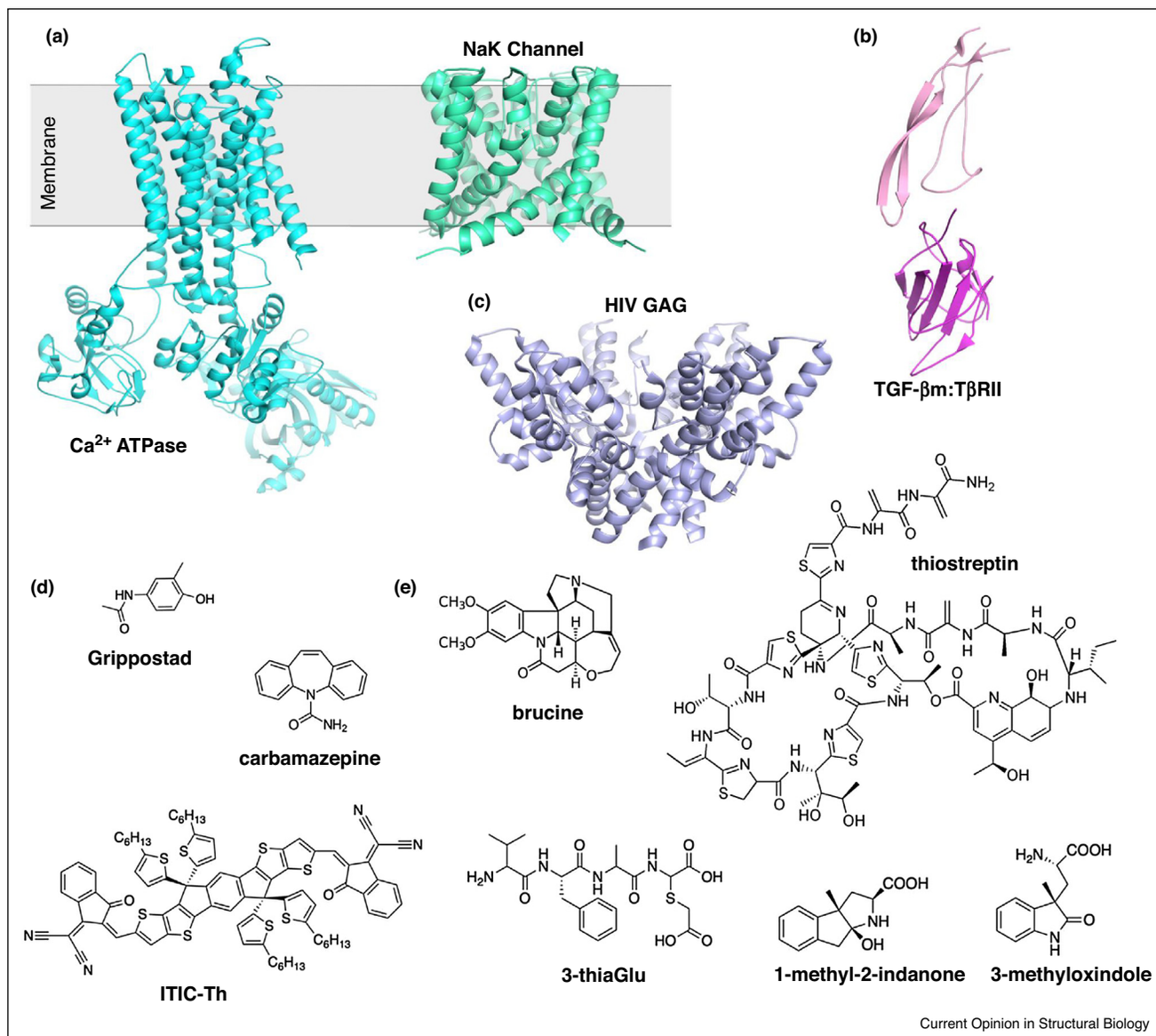
Figure 2



MicroED workflow.

From left to right. Crystals are grown, harvested, and placed directly on EM grids. Manual or automated screening to assess for electron diffraction of the crystals. When a well-diffracting crystal is identified, a dataset is collected by continuous rotation. The diffraction dataset is then processed using standard crystallography software to allow model building and structure refinement.

Figure 3



Representative MicroED structures of membrane proteins, protein complexes, small-molecules, and natural-products.

(a) MicroED structures of the membrane proteins rendered as cyan and green ribbon for Ca²⁺ ATPase (PDB 3J7T/U) [34**] and NaK (PDB 6CPV) [35**], respectively. (b) MicroED structure of the TGF-βm:TβRII protein–protein complex (PDB 5TY4) [36]. Protein rendered as pink and magenta ribbon for TGF-βm and TβRII, respectively. (c) MicroED structure of the HIV-GAG-bevirimat rendered as blue ribbon (PDB 6N3U) [39]. (d) A gallery of small-molecules with structures determined by MicroED. (e) Examples of natural-products with structures determined by MicroED.

+ ATPase, the structure of NaK by MicroED allowed generation of Coulomb potential maps to unambiguously place Na⁺ within the channel and to visualize a new transient state of the channel [35**]. The heterodimeric complex between TGF-βm and TβRII plays essential roles in the adaptive immune response and maintenance of the extracellular matrix [38]. Unlike Ca²⁺ ATPase and NaK, which formed nanocrystals, the structure of TGF-βm:TβRII was obtained from fragmentation of large, imperfect crystals (discussed below) [36]. Compared with

the imperfect, large parent crystals, this approach led to better MicroED data from well-ordered microcrystal fragments that ultimately yielded atomic-resolution structures [36]. This study expanded the application of MicroED to include a wider range of crystal sizes.

MicroED in drug discovery

MicroED has already made important contributions to drug discovery by determining structures of protein–drug complexes and supra-resolution of small-molecules and

natural-products, often directly from powders, bypassing crystallization experiments. The MicroED structure of HIV-GAG, which plays important roles in the life-cycle of HIV, was solved in complex with the antiviral drug, bevirimat (PDB 6N3U) [39^{*}] (Figure 3c). The HIV-GAG-bevirimat complex provided important information about the antiviral drug mechanism and was the first demonstration of drug discovery using MicroED.

MicroED was originally intended for studying protein assemblies [26], however, it was rapidly recognized that this technique is a powerful tool for the characterization of small-molecules and natural-products. In 2016, the structure of the sodium channel blocker carbamazepine was determined to ~ 1 Å resolution [40] (Figure 3d). In 2018, a method for small-molecule sample preparation using a "powder to structure" pipeline was described for carbamazepine [33] and later expanded to several small organic molecules [4^{**}]. The structure of MBBF4 [41^{*}], a methylene blue derivative with wide medical applications including its activity as a photo-activatable antimicrobial agent [42], was also solved. Since then, several structures of small-molecules have been reported by MicroED. These MicroED structures include Grippostad [41^{*}], an antiviral drug for the treatment of the common cold and the flu [43], and a recent example of the non-fullerene acceptor (NFA) semi-conductive material ITIC-Th (Figure 3d).

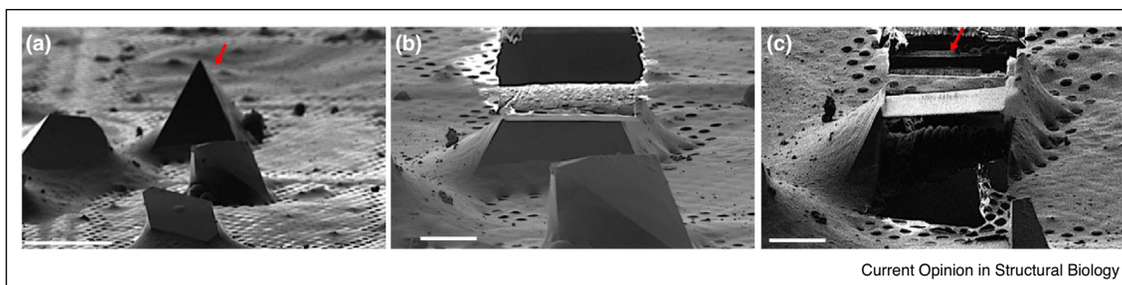
MicroED has also proven its usefulness for the structure characterization of several natural-products that have previously been challenging or, in some cases, impossible to determine by other techniques. Unlike their synthetic small-molecules counterparts, biosynthesized natural-products are typically larger, structurally dynamic, obtained in small amounts, and difficult to crystallize, posing considerable challenges for X-ray studies. Even when natural-products form lattices, these crystals are often too small and are not useful for X-ray diffraction [44^{**},45^{**}]. Brucine is an alkaloid toxin currently being

tested for its anticancer properties [46] (Figure 3e). The MicroED structure of brucine at 0.9 Å resolution allowed for definitive assignment of its two chiral centers, key for understanding its toxicity and anticancer properties [4^{**}] (Figure 3). Brucine, while large compared to small-molecules, is relatively small compared to amino-acid derived natural-products called ribosomally synthesized and post-translationally modified peptides (RiPPs), including 3-thiaGlu [44^{**}] and thioStreptin [47] (Figure 3e). Glutamylated thiols, similar to the peptide modification on 3-thiaGlu, have been shown to block jasmonate and ethylene signaling pathways [48]. ThioStreptin is an antibiotic currently used in veterinary medicine [47] (Figure 3e). When efforts failed by X-ray crystallography, MicroED readily provided a 0.9 Å resolution of the 3-thiaGlu peptide (PDB 6PO6) [44^{**}]. While thioStreptin has been studied by NMR [44^{**}] and X-ray crystallography [49] previously, the ease of its characterization speaks to the robustness of MicroED for structure determination of large, flexible natural-products (Figure 3e). Like the difficulties encountered for 3-thiaGlu, the structures of 3-substituted oxindole derivatives (Figure 3e), that contain a new stereocenter at the γ carbon installed by an enzyme through directed-evolution, was only solved with the application of MicroED [45^{**}]. These recent studies demonstrate that MicroED provides an additional strategy for more complete analysis of absolute configuration based on internal markers [44^{**},45^{**}].

Strategies for crystal preparation for MicroED

Electrons interact much more strongly with material than X-ray [23]. This phenomenon, however, results in high absorption and, thus, electrons can only penetrate very thin materials. Crystals that are greater than 500 nm in thickness must be thinned before MicroED data can be collected [36]. There are two strategies for trimming large crystals to thicknesses suitable for MicroED diffraction including mechanical fragmentation (typically by sonication, vigorous pipetting, or vortexing) [36] and milling with a focused ion beam (FIB) [50,51,52^{*}]. Mechanical

Figure 4



FIB milling of crystals for MicroED.

(a) Image of select proteinase K crystals at high magnification before milling. The arrow indicates the crystal that was milled. (b) FIB image after milling the top of the crystal. (c) FIB image after milling and cleaning both the top and bottom of the crystal leaving a lamella indicated by an arrow (Reprint from Ref. [52^{*}]).

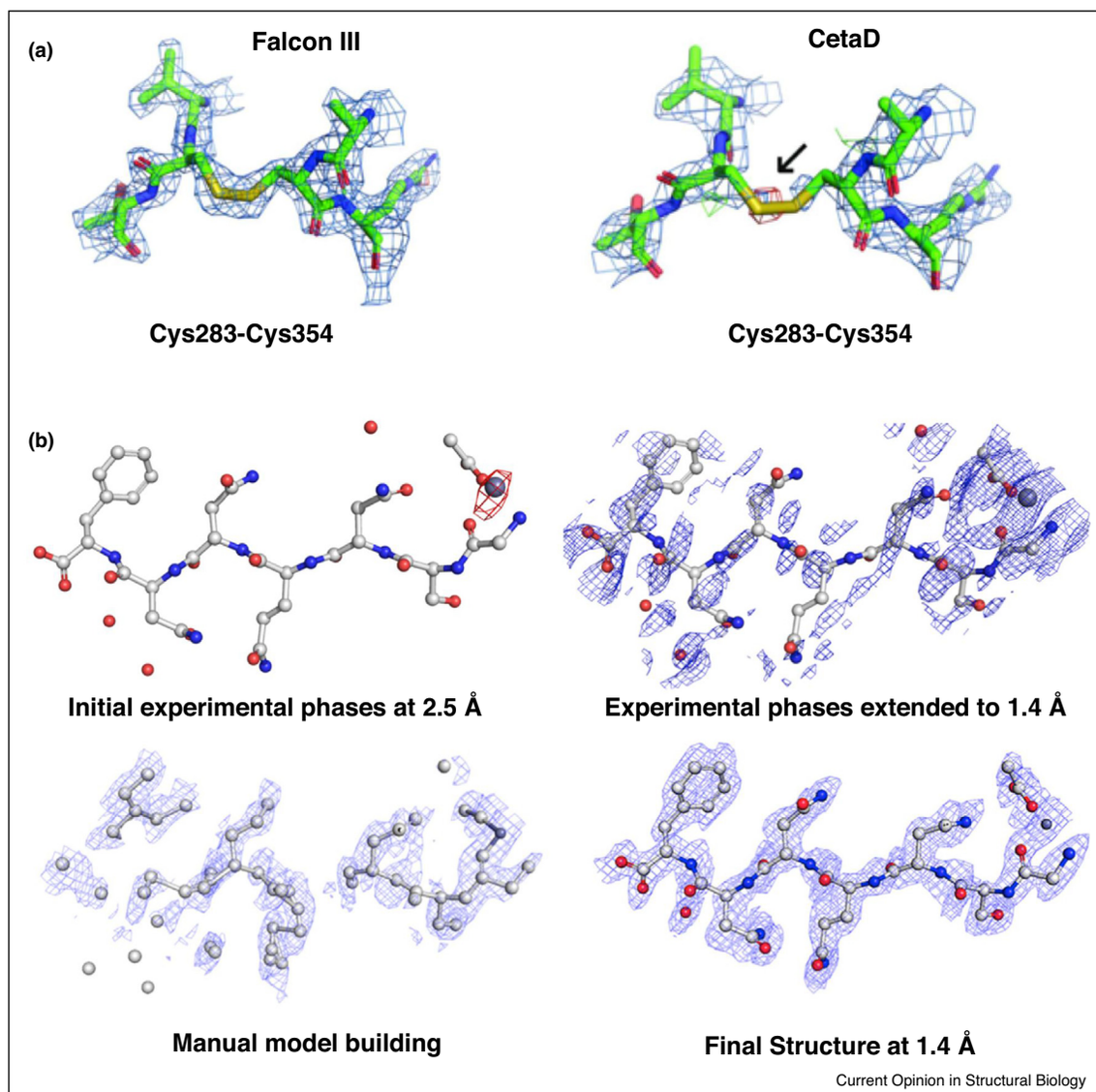
fragmentation has been successful for determining protein structures from large crystals of lysozyme, TGF- β m T β R11, xylanase, thaumatin, trypsin, proteinase K, thermolysin, and a segment of the protein tau [36]. Moving forward, the most current and promising technique for trimming large crystals for MicroED is FIB milling [50,51,52^{*}]. During FIB milling, a crystal is repeatedly exposed a gallium beam to trim away the surrounding materials and generate lamellas with controllable thicknesses. As proof of principle, the structures of several proteins, including lysosome and proteinase K, have been determined by FIB mill and MicroED [50,51,52^{*},53,54

(Figure 4). Currently, FIB milling crystals is a relatively slow process but, even then, about ten crystal lamellas can be prepared per day.

Outrunning radiation damage and phasing MicroED data

Radiation damage in structural studies continue to be a major challenge leading to poor processing statistics and map quality [55,56]. When electrons penetrate materials, they deposit energy that can deteriorate the samples, a process referred to as radiation damage. Radiation damage can be categorized into two forms: global and

Figure 5



MicroED radiation damage and experimental phasing.

(a) Disulfide bonds of the proteinase K structures determined from data collected on Falcon III and CetaD. The 2mFo-DFc densities (blue meshes) are contoured at 1.5 σ (Reprint of Ref. [63^{**}]). (b) Fourier difference maps between the damaged and undamaged structure of a peptide, contoured at 3 σ (Top left). Maps of the experimental phases of the peptide extended to 1.4 Å (Top right). Maps of an intermediate model-building step (Bottom left) to generate the final peptide structure at 1.4 Å (Bottom right) (Reprint from Ref. [64^{**}]).

site-specific. Global radiation damage typically results in the disruption of the crystal lattice which can be detected during data-processing when decreases in overall diffraction intensities and increases in B-factors are observed [57,58]. On the other hand, site-specific radiation damage is not uniform, is not typically detected during data-processing, and observable only during examination of the real-space map [59]. The degree of radiation damage depends, among other things, on the content of the sample, the surrounding solution, and is proportional to the amount of energy used during diffraction studies. For MicroED, site-specific radiation damage has been illustrated to occur on specific amino-acids including cysteines, glutamates, and aspartic acids [60]. To curb the effects of radiation damage, samples are often vitrified [61]. However, even the combination of vitrification and exposure to extremely low doses of electrons ($0.01 \text{ e}^-/\text{\AA}^2/\text{s}$) during MicroED experiments can still lead to detectable radiation damage [60].

TEMs for cryo-EM studies are typically equipped with highly-sensitive direct-electron detectors designed for imaging [9,14,62]. These highly sensitive cameras, however, have not been used extensively for MicroED because of concerns of damage to the sensors. As such, MicroED data are typically collected on indirect-electron detectors such as the complementary metal oxide semiconductor (CMOS)-based CetaD and TVIPS TemCam-F416 cameras. This strategy, however, limits the availability of MicroED because most facility TEMs are typically outfitted with top-of-the-line direct-electron detectors for imaging and not CMOS cameras. Recently, the Falcon III direct-electron camera was tested for MicroED data collection [63]. This study demonstrates that MicroED data collected at lower electron exposure, to avoid camera damage of Falcon III, lead to greater mean completeness relative to CMOS detectors and to higher quality maps. As proof of principle, examination of the maps of proteinase K from data collected on the Falcon III camera preserved the disulfide bonds, which are highly susceptible to radiation damage [63] (Figure 5a). A similar approach has been used recently with a Gatan K2 direct-electron detector in counting mode with an exposure 25 times less than for the Falcon III detector and a seemingly damage-free structure of Proteinase K has been determined.

Outrunning radiation damage using the direct electron detector Falcon III has been instrumental in determining the structures of samples that are highly susceptible to damage [44,45] and we believe that many more such examples would be forthcoming. Radiation damage was recently exploited to establish a pipeline for phasing MicroED data [64] (Figure 5b). A low-damage followed by a high-damage data sets were taken from the same crystal. A difference Patterson was calculated and allowed for generation of initial phases. Following cycles of

manual model building and refinement, the structure of a peptide was determined (Figure 5b). This study demonstrates the ability to extract meaningful phase information using radiation damage in MicroED.

Concluding remarks

MicroED is proving to be an important new tool in structural biology not only in determining structures of proteins but also of peptides, small organic and inorganic molecules, and natural-products. Continuous rotation MicroED is paradigm shifting because it has proven to be a robust, fast, and efficient method for structure determination of small molecules and natural products, even without crystallization and directly from mixtures [4]. To our knowledge, no other structural biology method is capable of determining atomic resolution structures directly from mixtures, making MicroED a useful and powerful tool for an array of problems that are yet to be explored. Moving forward, the application of FIB-milling and fast-cameras will certainly expand MicroED for structure determination of varying types of samples with a wide-range of crystal sizes and to facilitate time resolved studies. These advancements could ultimately be applied to establish automated pipelines that mirror those for X-ray crystallography to facilitate obtaining MicroED structures. Early examples of automation in MicroED has already reported in which several hundred data sets could be collected overnight to demonstrate similar throughput as synchrotrons [65].

Conflict of interest statement

Nothing declared.

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