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MicroED opens a new era for biological structure determination

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In 2013 we unveiled the cryo-electron microscopy (CryoEM) method of MicroED, or three-dimensional (3D) electron diffraction of microscopic crystals. Here tiny 3D crystals of biological material are used in an electron microscope for diffraction data collection under cryogenic conditions. The data is indexed, integrated, merged and scaled using standard X-ray crystallography software to determine structures at atomic resolution. In this review we provide an overview of the MicroED method and compare it with other CryoEM methods.

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Introduction

Whether being used to obtain images of individual biomolecules or using crystalline samples to collect high-resolution data through diffraction methods, cryoelectron microscopy, or CryoEM, is revolutionizing structural biology. CryoEM can be separated into four techniques for molecular structure determination (Table 1): cryo-tomography, single particle reconstruction, 2D electron crystallography, and Micro-electron diffraction, or MicroED. Cryo-tomography is generally used to study intact biomolecular complexes within the cellular environment. This provides mechanistic insights that are difficult to obtain by other methods. However, the structural data produced by cryo-tomography are relatively low resolution (1-5 nm resolution for thin samples). Single particle reconstructions are capable of producing biomolecular structures at near atomic resolution without the need of crystals as long as the samples are large (>200 kDa) homogenous and stable [1]. Both the 2D electron crystallography and MicroED techniques

use crystalline material; however due to the significant differences in samples (2D crystalline arrays vs. 3D crystals), differences in data collection (still diffraction vs. continuous rotation), and data processing (MRC suite of software vs. standard X-ray processing software), they should be considered distinct branches of CryoEM. In this review, we will focus on these two crystal-based CryoEM techniques, and describe recent progress in MicroED.

Electron diffraction in materials and in biology

Electron diffraction is a powerful tool for studying the atomic structures of materials and biological samples. Many methods have been developed over the years by a number of groups focusing on inorganic materials or small organic compounds that are relatively dose insensitive [2–6]. For biological samples that are dose sensitive and should be studied in cryogenic conditions with low dose methods, electron diffraction has traditionally been restricted to the use of two-dimensional (2D) crystals, which consist of a 1-3 layers of biomolecules ordered in a 2D crystalline lattice. These 2D crystals are capable of yielding atomic resolution structures of membrane proteins in the surrounding lipid environment [7,8]; however most structures solved by 2D crystallography are at modest resolutions (4–10 Å). It was not until 2013 that electron diffraction was used for the first time for determining the structure of a protein from 3D crystals by a newly established method called MicroED [9**]. This method has since been used to solve a number of macromolecular structures at cryogenic temperatures and with extremely low electron dose (Table 2).

Although many aspects of electron diffraction are analogous to X-ray crystallography, there are some important distinctions that should be kept in mind when comparing the two techniques. First, electrons interact much stronger with matter than X-rays and deposit less energy onto the sample [10]. This means that electron diffraction can extract meaningful high-resolution data from crystals that are orders of magnitude smaller in volume than what is needed from a conventional X-ray crystallography experiment. Second, the wavelength of electrons produced in an electron microscope is significantly smaller than the wavelengths used for X-ray crystallography. For example, the wavelength of the electrons from a microscope operated at 200 kV ($\lambda = 0.0251 \text{ Å}$) is over 60 times smaller than the X-rays from the commonly used Copper Kα source $(\lambda = 1.54 \text{ Å})$. This leads to an Ewald sphere that is much larger with electron diffraction, and the Ewald sphere is

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	Cryo-tomography	Single particle reconstructions	Electron crystallography	MicroED	
Samples typically used Strengths	Whole cells or organelles Directly observe molecular interactions	Purified biomolecules Material required is significantly less than in crystallography	2D crystals In the case of membrane proteins, the interactions with lipid bilayer can be seen	3D crystals Crystals used are several orders of magnitude smaller than those used in standard X-ray crystallography	
	Biomolecules are in the native cellular environment	Multiple conformations can be solved with a single sample	Protein dynamics probed	Data can be processed in standard X-ray crystallograph programs	
	No crystals necessary	No crystals necessary		Highest resolution cryo-EM technique	
Shortcomings	Relatively low resolution	Requires large proteins or protein complexes (size limitation)	Requires 2D crystals	Requires 3D crystals	
		Modest resolution	Incomplete data in the direction parallel to the electron beam		

essentially flat at the resolutions seen in macromolecular crystallography. Finally, because the scattered electrons can be focused by the microscope, images of the crystals can also be collected and accurate phase information can be recovered.

Biomolecular electron crystallography requires special set-up of the microscope because of the sensitivity of biological samples when compared with hard materials. Hydration is critical, and in order for the hydrated sample to withstand the high-vacuum within the TEM and still produce high-resolution data, samples are vitrified and loaded into a cryo-transmission electron microscope (cryo-TEM) [11]. Also, when compared to samples used in materials science, biological specimens are extremely radiation sensitive [12]. Therefore, very low doses are used when collecting MicroED data from biological material. We use approximately $0.01 \text{ e}^{-}/\text{Å}^{2}/\text{s}$ for biological samples, orders of magnitude smaller than what is used with more dose resistant samples that are typically exposed to approximately $1000 \text{ e}^{-}/\text{Å}^{2}/\text{s}$.

Electron crystallography of 2D protein arrays

The best-known examples of 2D electron crystallography involve work with membrane proteins that are crystallized within a lipid bilayer and produce planar 2D-crystals of protein and lipid. The pioneering work of Henderson and Unwin in 1975 produced 7 Å projection maps from naturally occurring 2D crystals of bacteriorhodopsin [13], which revealed how the α -helices are organized within a

Structures determined by MicroED								
Sample	Year	Data collection approach	Resolution	PDB	EMDB	SBGRIE		
Lysozyme	2013	Still diffraction	2.9 Å	3J4G [9**]	2945			
	2014	Continuous rotation	2.5 Å	3J6K [42**]	6342	185		
	2016	Continuous rotation	1.5 Å	5K7O	8217			
Catalase	2014	Continuous rotation	3.2 Å	3J7B [45]	6314	186		
	2015	Continuous rotation	3.2 Å	3J7U [53]				
Ca ²⁺ -ATPase	2015	Continuous rotation	3.4 Å	3J7T [53]				
α-Synuclein NACore	2015	Continuous rotation	1.4 Å	4RIL [46**]	3028	193		
α-Synuclein preNAC	2015	Continuous rotation	1.4 Å	4ZNN [46**]	3001			
Proteinase K	2016	Continuous rotation	1.75 Å	5I9S [54]	8077	262		
	2016	Continuous rotation	1.3 Å	5K7S	8221			
Prion Zn-NNQQNY	2016	Continuous rotation	1.0 Å	5K2E [55]	8196			
Prion Cd-NNQQNY	2016	Continuous rotation	1.0 Å	5K2F [55]	8197			
Prion GNNQQNY1	2016	Continuous rotation	1.1 Å	5K2G [55]	8198			
Prion GNNQQNY2	2016	Continuous rotation	1.05 Å	5K2H [55]	8199			
Tau peptide	2016	Continuous rotation	1.1 Å	5K7N	8216			
Xylanase	2016	Continuous rotation	1.9 Å	5K7P	8218			
Thaumatin	2016	Continuous rotation	2.11 Å	5K7Q	8219			
Trypsin	2016	Continuous rotation	1.5 Å	5K7R	8220			
Thermolysin	2016	Continuous rotation	1.6 Å	5K7T	8222			

transmembrane protein for the first time. Later work with bacteriorhodopsin 2D crystals yielded a high-resolution data where a complete atomic model of the protein could be built and refined [14]. Following the work on bacteriorhodopsin, several high-resolution structures of membrane proteins have been solved, including, but not limited to, the plant light-harvesting complex II [15] and several aquaporins [7,8,16,17]. The structure of taxol stabilized $\alpha\beta$ tubulin was also solved by this method although the protein is not membrane embedded [18]. In addition to producing high-resolution structures, 2D electron crystallography also provides quality structural information from relatively low-resolution structures, due to the high quality phase information obtained from imaging the crystals [19°,20–23].

2D crystals are typically grown from solutions when detergent solubilized lipids are mixed together with the detergent solubilized membrane protein of interest. Subsequent removal of the detergent by slow dialysis leads to protein reconstitution and crystallization within the newly forming lipid membranes [24]. Crystals are applied to a continuous-carbon coated EM grid, excess liquid is removed by blotting, and the preparation is then frozen in liquid ethane or nitrogen. After being loaded into the cryoEM, either images or diffraction data can be collected. To obtain a 3D dataset, diffraction patterns or images of the crystals must be collected at various tilt angles. Each crystal is imaged or diffracted at a known tilt angle. Radiation damage prohibits the collection of a full tilt series from a single 2D crystal. Therefore, full data sets are constructed by merging data from many crystals covering the tilt range. Because the 2D crystals all lie on the grid in a preferred orientation and the samples cannot be tilted beyond a certain angle, data collected from 2D crystals suffer from a systematic incompleteness often referred to as a 'missing cone' [25]. This leads to anisotropic data, with higher resolution data being obtained parallel to the plane of the membrane.

Collecting images of 2D crystals provides both amplitudes and phases, and are sufficient to determine a 3D density map of the protein. However, while the phases are extremely accurate, the amplitudes are affected by the contrast transfer function and are not as high-quality. Additionally, obtaining high resolution images can be difficult especially at high-tilt angles. Imaging difficulties arise in large part due to mechanical and thermal induced drifting of the sample and charging effects that reduce the obtainable resolution [26]. The collection of electron diffraction data is much less sensitive to these same sample drift problems, and can produce high-resolution and high-quality amplitude information. However, with diffraction all phase information is lost. To obtain phases, the high-resolution diffraction data can be combined with the high-quality phase information obtained from imaging. When the diffraction resolution is significantly higher

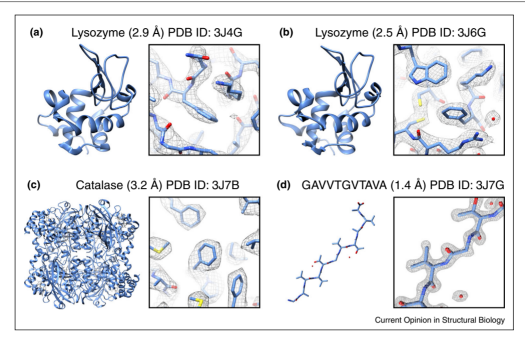
than that of the images, as is often the case, fragmentbased phase extension methods can be used [27]. Detailed procedures for performing 2D crystallographic experiments can be found in the many reviews written on the topic [24,28–30].

Electron crystallography of 2D arrays has been used effectively to study protein dynamics. This has been accomplished for a number of membrane proteins including: the light activated bacteriorhodopsin [31], the acetylcholine receptor embedded in tubular arrays [32], and more recently the voltage gated sodium channel [19°].

Electron crystallography goes 3D with MicroED

Although the initial work with 2D crystals was being performed in the 1970s, thin 3D microcrystals (a few hundred nanometers thick) were also being studied by EM, with the best example being bovine liver catalase [33–38]. In work by Dorset and Parsons, it was shown that hydrated catalase microcrystals could provide diffraction data beyond 3 A and that the diffraction was not affected by multiple scattering [35]. Not long after this work, Unwin and Henderson determined a 9 Å projection structure of catalase from 3D catalase crystals preserved with glucose [33]. More recently 3D crystals of lysozyme were imaged and the mosaic building blocks identified [39°]. Although these were important steps forward for electron microscopy, a full 3D structure was never determined because the existing methodology, specifically how collect and process 3D diffraction data, was not adequate for 3D crystals. Therefore, it was important that a new method for data collection had to be developed.

In 2013, a new technique called Microcrystal Electron Diffraction [9**,40], or MicroED, was developed based on the idea that if multiple diffraction patters were taken from a single crystal, a large enough region of reciprocal space could be sampled, allowing proper indexing and processing of the diffraction data set. The first iteration of MicroED used a series of still diffraction patterns from single crystals (approximately 2 million unit cells in size), with the crystal rotated in discreet angles between exposures. Each diffraction pattern was collected at liquid nitrogen temperature with an ultra-low dose rate of $\sim 0.01 \text{ e}^{-}/\text{Å}^{2}/\text{s}$. This extremely low dose rate still produced quality diffraction patterns to high resolution (up to 1.7 Å with lysozyme microcrystals initially) and allowed each crystal to be exposed up to 90 times [9**]. High-tilt cryoholders were used which allowed the samples to be tilted up to $\pm 70^{\circ}$ (140° total possible rotation range), and the crystals were tilted by 1° between exposures. This led to data sets which produced up to a 90° wedge of data from a single crystal. Because such a large amount of reciprocal space was sampled, the crystal orientation could be determined accurately and the diffraction patterns properly indexed. Each crystal data set was indexed



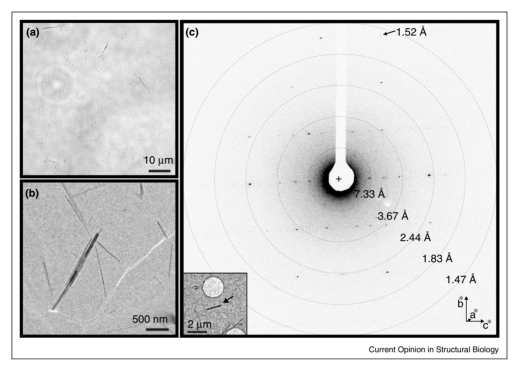
Example structures determined from MicroED. Structures are shown along with their PDB ID, resolution, full model (left) and representative region of the model and density map (right).

and integrated, followed by merging of all data into one final data set. The data from 3 crystals were integrated and merged using software developed specifically for still diffraction MicroED data [41]. This initial proof of concept work yielded a refined lysozyme structure to 2.9 Å (Figure 1a) with very good refinement statistics, and represented the first time electron diffraction had been used successfully to determine the structure of a protein from 3D microcrystals in cryogenic conditions.

Following this first proof of principle work, an improved method of MicroED data collection was developed [42**]. This new method, called 'continuous-rotation' MicroED data collection, uses the high frame rates achievable by a CMOS detector or a direct electron detector to collect diffraction data as the crystal is continuously rotated in the electron beam by the sample stage of the microscope. The methodology is similar to other techniques used for non-biological dose insensitive samples studied at ambient temperatures [2,4–6]. Continuous rotation MicroED led to several improvements in data quality and allowed for data processing using standard X-ray crystallographic software [43°]. The first benefit of continuous rotation data collection comes from a reduction in dynamic scattering, a process where the electron beam diffracts multiple times within the sample leading to a redistribution of reflection intensities. As was seen previously with precession electron diffraction, rotating the sample by continuous rotation (or the beam in the case of precession) is able to reduce dynamic scattering and intensity redistribution, which ultimately leads to more accurate intensity measurements [44]. The second advantage comes from the improved sampling of reciprocal space that occurs as the sample is rotated and data is collected. This allows the collection of full reflection intensities as opposed to the partial sampling obtained when the crystal remains still during data collection of individual frames. The third benefit of continuous rotation is that the movement of the crystal in the electron beam is similar to the crystal movements in X-ray crystallography. Because the experimental setups are so similar, MicroED data collected by continuous rotation can be processed by programs developed for X-ray diffraction data. Because of its many advantages over the collection of diffraction stills, 'continuous-rotation' data collection has become the standard method of MicroED data collection, and its first use resulted in a 2.5 Å structure of lysozyme using a single nanocrystal (Figure 1b). From this point forward, any mention of MicroED data collection in this review refers to continuous rotation data collection.

Following these studies the MicroED method was tested on a new sample, and for historical reasons mentioned above, we chose to study bovine liver catalase microcrystals [45]. Catalase (P2₁2₁2₁; a = 67.8 A, b = 172.1 A, $c = 182.1 \text{ Å}; \alpha = \beta = \gamma = 90^{\circ})$ presented a few unique challenges for MicroED, which included a larger unit cell and lower symmetry compared to lysozyme (P4₃2₁2; $a = b = 76.0 \text{ Å}, c = 37.2 \text{ Å}; \alpha = \beta = \gamma = 90^{\circ}), \text{ and crystals}$ which have a preferred orientation on the grid resulting

Figure 2



Crystals from the toxic core of α -synuclein. The crystals used to determine the α -synuclein structure were much too small to be seen by light microscopy (a). However, when visualized within the TEM many extremely small microcrystals could be seen (b), which diffracted to approximately 1.4 Å (c).

Source: Adapted from Ref. [46**].

in a missing cone of data in the direction of the c^* crystallographic axis. Data from a single nanocrystal (approximately $15 \times 7 \times 0.2 \,\mu\text{m}$) was collected while the stage rotated from approximately 53° to 0°. Subsequent data processing yielded a final scaled and merged diffraction data set from this single crystal that was 79.4% complete to 3.2 Å. With this amount of data, an acceptable molecular replacement solution could be found and the 3.2 Å resolution structure (Figure 1c) was refined with acceptable statistics ($R_{\text{work}}/R_{\text{free}} = 26.2\%/30.8\%$). Additional experiments with a molecular replacement search model that did not contain the bound NADP co-factor showed that the missing NADP could be found in the difference density maps. Furthermore, what was more exciting was that the subtle NADPH induced conformational change of phenylalanine-197 could be seen using the difference density maps, indicating the quality of MicroED data is in fact very high.

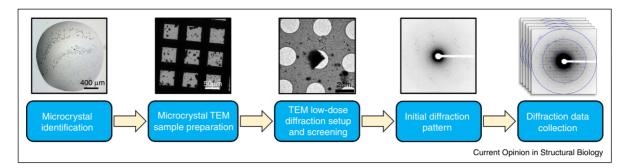
The first novel structures solved by MicroED were published in late 2015 [46°]. These structures were of peptide fragments that form the toxic core of α -synculein, the protein responsible for Parkinson's disease and lead to very important insights into how the protein forms toxic aggregates. The two α -synculein peptides formed very small and thin needle like nanocrystals (approximately

1–2 μm long, 20–50 nm wide and thick; Figure 2) and had resisted all other methods of structure determination, including those that utilize X-ray Free Electron Lasers (XFELs). The structures were solved by MicroED at 1.4 Å resolution and at the time represented the highest resolution structures of a biological molecules determined by any cryo-electron microscopy technique published to date (Figure 1d). More recent MicroED studies allowed direct phasing methods to be used for structure determination of 4 different samples at 1 Å resolution [55].

MicroED experimental setup and data collection

Samples prepared for MicroED are typically grown in standard hanging or sitting drop crystallization experiments as for X-ray crystallography. Microcrystals are transferred to a glow discharged grid by pipetting the drop directly onto the carbon surface. The excess sample is blotted and vitrified using automated vitrification devices in a procedure similar to what is used for single-particle or electron tomography cryo-EM work. Alternatively, the grids can be blotted by hand and frozen manually as for 2D crystals. Vitrified samples can be stored under liquid nitrogen until loading into a standard cryo-EM. Once a suitable crystal is found, the crystal is exposed with the beam in focus and diffraction data is

Figure 3



Flow diagram for MicroED data collection protocol. Source: Adapted from Ref. [47°].

continuously collected as the stage is rotated in the beam. See Figure 3 for a flow chart of the data collection process. Detailed protocols for sample preparation data collection have been recently published [47°].

MicroED data processing

Following data collection, the diffraction data set is indexed, integrated, merged and scaled using standard macromolecular crystallography programs originally developed for X-ray crystallography. Initially, data were processed in MOSFLM [48,49]; however, data has since also been processed in HKL2000 [50], DIALS [51], and XDS [52]. In depth descriptions and protocols on MicroED data processing have been previously published [43°], and we encourage readers to refer to this publication for detailed descriptions of the data processing procedures.

MicroED future directions

MicroED is still a relatively new method in CryoEM and there are many exciting areas of future research and improvements. One of the most interesting areas of MicroED research is the development of phasing methods. As was done with first structures solved by X-ray crystallography, the phase could be solved by the use of heavy atoms for isomorphous replacement could potentially be used to obtain phase information. A more unique interesting approach to obtain phases would be to incorporate imaging into MicroED. Images of 3D microcrystals have previously been collected and processed [39°], however integration into diffraction data has not been accomplished. The development of more accurate electron scattering factors promises to greatly improve refinement. There are also many other improvements to sample preparation, data collection, and data processing that will be developed when more samples are investigated with MicroED. It is important to note that many labs have been producing microcrystals for serial femtosecond crystallography (SFX), and many of these can be directly used for MicroED. Currently, the main advantages of MicroED over SFX are the availability of cryo-TEM

instrumentation for MicroED relative to the extremely limited access to XFELS for SFX, and the smaller amount of sample required for MicroED (1-10 crystals for MicroED vs. thousands to millions for SFX). As MicroED is further improved and refined, the technique promises to become a widespread tool for the determination of difficult biomolecular samples that form microcrystals. Together with other techniques, we expect that MicroED will play a major role in macromolecular structure determination in the coming decades.

Acknowledgements

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MicroED was used to determine the structure of the toxic core of α synuclein from peptide microcrystals that had resisted structure determination from all other methods. The final 1.4 Å structures represent the highest resolution by cryoEM and the first novel structures determined with MicroED.

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