Investigating eukaryotic cells with cryo-ET

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ABSTRACT The interior of eukaryotic cells is mysterious. How do the large communities of macromolecular machines interact with each other? How do the structures and positions of these nanoscopic entities respond to new stimuli? Questions like these can now be answered with the help of a method called electron cryotomography (cryo-ET). Cryo-ET will ultimately reveal the inner workings of a cell at the protein, secondary structure, and perhaps even side-chain levels. Combined with genetic or pharmacological perturbation, cryo-ET will allow us to answer previously unimaginable questions, such as how structure, biochemistry, and forces are related in situ. Because it bridges structural biology and cell biology, cryo-ET is indispensable for structural cell biology—the study of the 3-D macromolecular structure of cells. Here we discuss some of the key ideas, strategies, auxiliary techniques, and innovations that an aspiring structural cell biologist will consider when planning to ask bold questions.

INTRODUCTION
Electron cryomicroscopy (cryo-EM) is a family of methods that is used to study biological structure, from atoms to cells. To preserve a biological sample’s molecular details, the cryo-EM sample is kept free of the chemical fixation, dehydration, and heavy-metal staining that are commonly used in traditional EM. Furthermore, the sample is frozen so quickly that water molecules are immobilized in an amorphous state, which keeps the biological material “hydrated.” Once frozen, the sample must be kept colder than −135°C before and during image acquisition in a transmission electron microscope. Because biological matter is damaged rapidly by the microscope’s electron beam, cryo-EM imaging is done using a very limited electron dose. The resulting low-dose images are noisy and therefore require careful processing and interpretation. These stringent requirements ensure that the cryo-EM data represent biological structures in a minimally perturbed, lifelike state.

Two popular forms of cryo-EM are “single-particle” analysis (SPA) and electron cryotomography (cryo-ET). Structural biologists have used both these approaches to study macromolecular complexes, herein called complexes for brevity. SPA has revolutionized the structure determination of purified complexes. Because this method can routinely produce 3-D density maps in which protein side chains are resolved, SPA is a popular replacement for X-ray crystallography studies of complexes larger than ~100 kDa. In contrast, cryo-ET is a method that generates modest ~40 Å resolution 3-D density maps of complexes, organelles, and even cells. If multiple copies of a complex are identified, they can be analyzed by subtomogram averaging (STA). STA involves aligning and averaging multiple copies of the same complex, thereby increasing the signal-to-noise ratio and suppressing the missing-wedge artifacts (detailed later), resulting in higher-resolution maps (Asano et al., 2016; Oikonomou and Jensen, 2017; Hutchings and Zanetti, 2018; Schur, 2019).

This Perspective is written for cell biologists who are interested in learning and applying cryo-ET to their favorite eukaryotic systems. To cover a broad range of ideas, we have skipped some technical details, which are found in other literature and online training resources (Henderson, 1995; Koster et al., 1997; Frank, 2006a,b; Dubochet, 2007; Penczek, 2010; Shen and Iwasa, 2018; Vos and Jensen, 2018; Rodenburg, 2019). We discuss some nonintuitive principles of cryo-ET and our expectations from current and future technologies. We also review some studies that use artificially thin samples like cryosections and cryolamellae because such samples vastly expand the range of questions that can be asked. Finally, we explore how cryo-ET can be used to study large cellular machines that defy conventional approaches.
TOMOGRAPHY AND TOMOGRAMS

Because heavy-metal stains are not used, cryo-EM image features come from the atoms within proteins, nucleic acids, and other biological molecules. Cryo-EM images are approximations of 2-D “projections” of 3-D objects, meaning that the pixel values are proportional to the mass summed along parallel linear paths through the sample (Figure 1A). Real-world projections deviate from ideal projections because they are influenced by the transfer functions of the microscope and the camera system. Projection images encode information about the interior of the object, making them fundamentally different from shadows (Figure 1B). The pixel values, called densities are usually represented in grayscale. Note that the term “densities” is contextual and can also mean 3-D pixel (voxel) intensity values or groups of pixels (or voxels) that belong to the same complex.

A cryo-ET raw data set is a series of images, typically recorded over a tilt range from $-60^\circ$ to $+60^\circ$ ($\pm 60^\circ$), with tilt increments ranging from 1° to 4°. This data set has discrete angular sampling and is called a tilt series (Figure 1, C and D, and Supplemental Movie S1). Tilt-series images are mutually aligned to a common 3-D reference model. This alignment is done using image features from the cell itself or with the aid of fiducial markers, such as gold colloids that were added prior to sample freezing. The aligned images are then typically combined by an algorithm called “back projection” to generate a 3-D density map called a cryotomogram. The limited tilt range of ($\pm 60^\circ$) results in missing-wedge artifacts, which manifest as image distortions. These distortions include the triangular features at 6 and 12 o’clock, the spokelike feature protruding from the bottom of the U, and the poorly defined lower portion of the letter S.

FIGURE 1: The principles of cryo-EM and cryo-ET. (A) The example object is “NUS” in block letters, centered within a thin spherical shell. The object’s projection image (bounded by a trapezoid) is shown below the Cartesian coordinates symbol. Electrons (e −, white arrow) travel along the z-axis. To show the object’s orientation, the front of the shell has been removed. (B) Cryo-EM images are projections, not shadows. Density profiles (ρ) along the dashed lines (x-axis) are shown in the plots below. In a cryo-ET experiment, the sample is rotated around the tilt axis (y-axis in this example), typically over a tilt range from $-60^\circ$ to $+60^\circ$. Five possible tilted orientations are shown. (D) A tilt series consists of the set of cryo-EM images recorded over this tilt range. Each image corresponds to the object, as oriented directly above in C. The image gray levels are proportional to the samples’ projected mass along the z-axis. Together, the images encode the information needed for a 3-D reconstruction. Supplemental Movie S1 shows the full tilt series. (E) After alignment, the images (bounded by trapezoids) are oriented according to their corresponding tilt angles and then “back projected” to generate a 3-D density map called a cryotomogram. The limited tilt range of ($\pm 60^\circ$) results in missing-wedge artifacts, which manifest as image distortions. These distortions include the triangular features at 6 and 12 o’clock, the spokelike feature protruding from the bottom of the U, and the poorly defined lower portion of the letter S.
cryotomogram to appear distorted relative to the original structure (Figure 1, A and E).

When visualized in Fourier space, the missing cryo-ET data appear like a pair of empty wedges that intersect along a line that runs parallel to the tilt axis (Figure 2A). This missing wedge limits the resolution along the electron-optical “Z” axis and results in elongated and smeared density-map features. We illustrate these distortions using the Dam1C/DASH outer-kinetochore complex and the nucleosome, which we have simulated with 60° of data missing (Figure 2, B and C). Supplemental Movies S2 and S3 illustrate the bizarre appearance of cryotomograms when these complexes are rotated relative to a stationary missing wedge. Fortunately, many subtomogram-analysis programs have routines that account for the missing wedge during the alignment process. Within the cell, most known complexes are randomly oriented. When multiple copies of the complex are aligned, their respective missing wedges will be randomly oriented. The Fourier transform of the resulting average has the missing data filled in, meaning that the reconstruction no longer has missing-wedge artifacts.

How to learn cryo-ET
There are three major skill sets to cryo-ET. The first skill set is sample preparation, meaning the operation of rapid-freezing machines. Training in the two principal techniques of rapid freezing—plunge freezing and high-pressure freezing (Dahl and Staehelin, 1989; Dobro et al., 2010)—is typically offered by cryo-EM facility staff. The second skill set is the operation of cryo-EM hardware and software. Some cryo-EM facilities also offer hands-on training in cryomicroscope operation. The third skill set is image processing, which unlike the other two, can be self-taught with the assistance of online tutorials and “walk-throughs.” Recently, some academic organizations have started offering on-site courses in one or more of these critical skill sets. Training workshops have also been offered by cryo-EM centers such as Diamond eBIC (Diamond, 2019), NeCEN (Netherlands Centre for Electron Nanoscopy, www.necn.nl, 2019), and NYSBC (New York Structural Biology Center, https://nysbc.org, 2019). These intensive hands-on courses are usually announced on the cryo-EM mailing lists CCP-EM (CCP-EM, 2019) and 3DEM (Perkins, 2019) and the IMOD list (Mastronarde, 2019) and also on social media such as the Twitter platform.

New professionally produced web resources such as em-learning.com and cryoem101.org are making theoretical training more accessible (Shen and Iwasa, 2018; Vos and Jensen, 2018). In addition, the University of Colorado provides a step-by-step tomogram-reconstruction tutorial using their popular IMOD package (Mastronarde, 1997; O’Toole, 2018). This tutorial includes a tilt series of a negatively stained plastic section of a Chlamydomonas cell. The IMOD tutorial can be done in 1 day and is a gateway to the analysis of cryo-ET data, which are more challenging. Cellular cryo-ET data are accessible from the EMPIAR database (Jüldin et al., 2016) and from the Jensen lab’s ETDB-Caltech database (Ortega et al., 2019). Furthermore, we have deposited our surplus data in EMPIAR-10227 (Gan et al., 2019). Surplus data are tomograms or positions within tomograms (and their corresponding tilt series) that contain cellular structures not presented or analyzed for a research paper. Resources like these enable new practitioners to gain proficiency in the essential steps of cryo-ET image processing from the comfort of their office.

How to decide on the suitability and feasibility of cellular cryo-ET
The three most popular techniques of cellular imaging are light microscopy, traditional EM, and cryo-ET. Each of these approaches has

FIGURE 2: The missing wedge influences cryotomograms in unintuitive ways. The left subpanels correspond to a “complete” data set (180° tilt range) while the right subpanels correspond to an incomplete data set, with a 60° wedge of data missing. The tilt axis is parallel to the y-axis, and the electron beam is parallel to the z-axis.
(A) Volumetric representations of cryo-ET data in Fourier space; k_x, k_y, and k_z denote the Fourier-space coordinate system, which is aligned to the real-space coordinate system shown in the other panels. The right panel shows the two missing wedges, bounded by dashed lines. (B) Outer-kinetochore Dam1C/DASH ring densities simulated at 12 Å resolution. The rings in the bottom panel are rotated 90° around the x-axis, relative to an unchanged missing wedge. From PDB 6CFZ (Jenni and Harrison, 2018). (C) Nucleosome densities, also simulated at 12 Å resolution and shown in two orientations like in B. From PDB 1KXS (Davey et al., 2002). Supplemental Movies S2 and S3 show how the Dam1C/DASH ring and the nucleosome are distorted in other orientations relative to the missing wedge. Note that real cryotomograms are affected by additional factors not modeled here. These factors include defocus, radiation damage, and discrete angular sampling.
advantages (Table 1). The biggest advantage of cryo-ET is that it can provide structural information, which we define as the 3-D arrangement of image densities that correspond to the proteins, nucleic acids, and other biological material within the cell. Furthermore, cryo-ET is essential if the study requires 3-D localization at the nanometer level or if the complex being studied is either invisible or distorted in traditional and immuno-EM data. Superresolution light microscopy can in principle achieve 3-D localization at the nanometer level, but this positional information only applies to the fluorophore, which in the case of fluorescent proteins is displaced from the complex by a few nanometers. Note that many structural cell biology projects require two or more forms of imaging.

The next questions concern feasibility. First, can the sample be cooled to a frozen-hydrated state by plunge freezing? Samples that are smaller than approximately 10 µm can be frozen without the formation of crystalline ice when plunged into a cryogen like liquid ethane or liquid propane-ethane mixture (Tivol et al., 2008). To increase the chances of complete vitrification (the absence of crystalline ice), the plunge-frozen sample should be as thin as possible. Small cells like yeast are amenable to plunge freezing. Larger plunge-frozen cells have vitreous ice in their thinner extremities and crystalline ice in their thicker parts. Samples up to nearly 1 mm thick are high-pressure-frozen in the presence of 2000 bars of pressure to ensure the ice is vitreous throughout their entire volume. The upper limit of such “thick” samples include cell pastes, tissues, and small animals, which are sometimes high-pressure-frozen in the presence of a cryoprotectant (Harapin et al., 2015; Schaffer et al., 2019).

Next, one needs to ask, what complex needs to be observed? Can these complexes be seen in a cellular cryotomogram? How abundant must these complexes be to answer the project’s main question? Are the complexes large and monolithic enough to be unambiguously identified? These questions do not have simple answers, so we will make some calculations using a few well-studied complexes. As a cellular sample, we will consider a 100-nm-thick cryosection cut through the center of an early-interphase yeast nucleus, which has a ∼2 µm diameter. In this example, the total nuclear volume is ∼4.2 µm³ and the cryotomogram contains ∼0.3 µm³, or approximately 7% of the nucleus.

Our first example is the nucleosome, a 200-kDa complex shaped like a cylinder, 10 nm wide and 6 nm thick. The nucleosome is currently one the smallest complexes that can be detected in a cellular cryotomogram. Yeast cells have ∼12 megabases of genomic DNA and a ∼160-base-pair-long nucleosome repeat length, meaning that there are ∼75,000 nucleosomes per nucleus. We therefore expect a yeast nucleus cryosection to contain ∼5000 nucleosomes per cryotomogram, making nucleosomes amenable to identification by a purification in silico approach. Our second example is the 66-MDa nuclear pore complex (Rout and Blobel, 1993), which is more than 96 nm wide and 35 nm thick (Yang et al., 1998). There are fewer than 100 nuclear pore complexes per yeast cell, so it is not surprising that cryotomograms (in our experience) rarely contain more than one complete complex. Nuclear pore complexes can nevertheless be easily located and then identified because they are conspicuously positioned at sites of inner- and outer-nuclear membrane fusion. Both nucleosomes and nuclear pore complexes have a largely random nuclear distribution, meaning that cryotomograms that contain a piece of the nucleus usually contain these two complexes.

Rare complexes whose structures are poorly characterized in situ are more challenging to study. One such example is the replication origin, which hosts numerous interesting macromolecular complexes. Each yeast cell contains ∼300 replication origins (Nieduszynski et al., 2007). We estimate that a yeast cryosection cryotomogram should contain ∼20 origins. Let’s assume that these origins can be identified (a different challenge discussed later) and that the replication subassembly of interest is conformationally and constitutionally homogeneous. Then 50 cryotomograms will provide ∼1000 origin-containing subtomograms, which is enough to generate an average at better than 30 Å resolution. Extremely rare complexes (averaging fewer than 1 per cryotomogram) that have nonrandom locations, such as kinetochores and microtubule-organizing centers, require alternative strategies that are discussed in later sections.

**TABLE 1. Comparison of popular cell biology microscopy methods**

| Information               | Example                                                                 | Method          |
|---------------------------|-------------------------------------------------------------------------|----------------|
| Cytology                  | Architecture of mitochondrial network                                    | +              |
| Location                  | Ribosome assembly factor in nucleoli                                     | +              |
| Distance                  | Diameter of a membrane tubule                                           | +              |
| Volume                    | Enlargement of lipid bodies                                             | +              |
| Counts                    | Nucleosomes per unit volume                                             | +              |
| Positions                 | Ribosome redistribution under stress                                     | +              |
| Structure                 | 3-D structure of kinetochore subunits                                    | +              |
| Heterogeneity             | Oligomerization states of proteasomes                                    | +              |
| Higher-order structure    | Path and orientation of sequential nucleosomes and ribosomes;           | +              |
|                           | mammalian kinetochores                                                 |                |

LM, conventional and superresolution light microscopy; EM, traditional, immuno, and serial EM. Note that there are exceptions to these broad guidelines and that superresolution light microscopy may eventually be able to answer some of the questions that are now exclusive to cryo-ET.

**TABLE 1. Comparison of popular cell biology microscopy methods**

**WORKFLOWS AND STRATEGIES**

**Preparation of thin frozen-hydrated cell samples**

Cryo-ET samples typically are thinner than ∼500 nm, but may be thicker or much thinner depending on the resolution requirements and the microscope hardware (Grimm et al., 1998). Cryotomograms of ∼500-nm-thick samples have enough resolution to reveal the positions of membranes and large complexes such as ribosomes. STA generally requires samples thinner than 200 nm. Thin
cellular samples can be prepared by cryomicrotomy or cryo-focused-ion-beam (cryo-FIB) milling, which produce cryosections and cryolamellae, respectively. Cryosections are typically ~50–200 nm thick and are cut from a block of frozen-hydrated cells (or tissue) by a diamond knife cooled to ≤−140°C. Cryomicrotomy was once considered feasible only for those with extremely steady hands and years of training. Advanced micromanipulation techniques have now made cryomicrotomy easier and more reproducible (Ladinsky et al., 2006; Studer et al., 2014). FIB milling is routinely used in materials science and has been painstakingly adapted for frozen-hydrated life science sample preparation (Marko et al., 2006; Hayles et al., 2007; Rigort et al., 2010; Villa et al., 2013; Mahamid et al., 2015; Medeiros et al., 2018). In a cryo-FIB milling experiment, the operator uses a beam of gallium ions to remove a frozen-hydrated cell mass to produce a thin plank-like sample suspended between two unmounted positions.

Cryomicrotomy has a few advantages over cryo-FIB milling. First, the instrument and the consumables are an order of magnitude cheaper. Second, cryomicrotomy can deal with tissues and small animals. Third, cryomicrotomy can generate cryo-ET samples faster than cryo-FIB milling. While cryomicrotomy is a more established technique, cryo-FIB milling has several advantages that are making it the more popular choice of cellular cryo-ET students. First, cryo-FIB milling does not cause compression and cutting artifacts like in cryomicrotomy (Dubochet et al., 2007). Cryo-FIB milling is therefore less likely to damage biological structures. Second, cryo-FIB milling is done within the vacuum of a computer-controlled scanning electron microscope, meaning that it suffers less contamination from room humidity, is less dependent on dexterity, and has better potential for automation. Cryomicrotomy and cryo-FIB milling can be combined to generate compression-free cryolamellae from macroporous samples (Rigort et al., 2010; Zhang et al., 2019). Further advances in sample freezing (Ravelli et al., 2010; Zhang et al., 2019) will greatly increase the diversity of milled cellular samples that can be studied by cryo-ET.

Searching for targets of interest
Cryo-ET imaging experiments begin with a search for grid positions that contain the cell(s) or complex(es) of interest. This search is done at low magnification to maximize the field of view and at ultra-low dose to minimize radiation damage prior to tilt-series collection. If the complex of interest is abundant, the microscopist can image cells at random. Rarer complexes can first be localized by fluorescence light cryomicroscopy. These positions are then imaged by cryo-EM (correlative light and electron cryomicroscopy, cryo-CLEM). In a cryo-CLEM experiment, the complex is tagged with a protein or molecule that is fluorescent at temperatures colder than ~135°C (Hampton et al., 2017). Some fluorescent proteins are photoactivatable at these cryogenic temperatures, meaning that superresolution light cryomicroscopy is possible (Chang et al., 2014; Dahlberg et al., 2018; Tuietl et al., 2019). Cryo-CLEM can facilitate the identification of complexes for which no SPA or crystal structure exists, as shown in a study of a bacterial compartmentalization complex (Schlimpert et al., 2012). Recently, cryo-CLEM of a membrane-associated GFP-fusion protein enabled the classification of yeast plasma-membrane domains called eisosomes (Bharat et al., 2018). Cryo-CLEM was also used to sample different positions associated with the actin-actin phenomenon in Dictyostelium cells (Jasnin et al., 2019). Cryo-CLEM will facilitate even more localizations (and possibly identification of the complex; see below) once superresolution light cryomicroscopy extends to 3-D and the workflow becomes routine.

Data collection
Modern electron microscopes are highly reliable and automated, driven by state-of-the-art software (Mastronarde, 2005; Suloway et al., 2009; Zheng et al., 2009). Hardware advances have also kept pace, enabling more structural details to be extracted from cellular samples. One major advance is the complementary metal–oxide semiconductor–based direct detector, which operates in “movie mode” and thereby enables partial correction of beam-induced motion blur (Brilot et al., 2012). The direct detectors with the fastest readouts make it possible to record data in which each imaging electron is registered. Such electron-counted data have the highest signal-to-noise ratios, which are evident in comparisons of the cameras’ detective quantum efficiencies (Ruskin et al., 2013). A second major advance is the phase plate, a device that increases the phase difference between the scattered and the unscattered electrons (Daney et al., 2014). The phase plate increases the low-resolution contrast, which makes complexes easier to detect. New developments in laser phase plates could make this technology more user-friendly and higher throughput than possible with the current carbon film-based phase plates (Schwartz et al., 2019). Cryosections and cryolamellae are thick by cryo-EM standards, meaning that they scatter more electrons inelastically. Commercially available cryomicroscopes cannot focus these lower-energy electrons at the image plane, leading to worse chromatic aberration. To deal with this problem, cryo-ET imaging should be done with an energy filter, which can remove the inelastically scattered electrons.

Data processing
Tomogram reconstruction is a multistep process, including motion correction, contrast transfer-function correction, image alignment, low-pass filtering, 3-D reconstruction, and denoising. These steps are all automated to some degree in the popular packages IMOD (Mastronarde, 1997) and EMAN2 (Chen et al., 2019). Automated tilt-series alignment without fiducials is also being addressed by software such as ProtoMo (Noble and Stagg, 2015). Following cryotomogram reconstruction, subcellular image features can be annotated either manually or semiautomatically using new machine-learning approaches (Chen et al., 2017; Zeng et al., 2018). Known complexes can be annotated semiautomatically by template matching (Heumann, 2016; Castaño-Diez, 2017). These annotations (also called segmentations) are the starting point for quantitative analysis. Note that cryo-ET data are commonly stored as MRC-formatted files (Cheng et al., 2015). Cryo-ET practitioners should write format-interconversion scripts if they want to adapt analysis programs from other microscopy fields.

Identification of complexes
Cryotomograms are noisy, making the identification of complexes (or their subunits) a major challenge in cryo-ET of cells. Large, abundant complexes can be identified by classification and STA, which produce density maps that resemble known SPA or crystal structures. The best-studied examples are ribosomes and proteasomes (Mahamid et al., 2016; Albert et al., 2017). In contrast, the identification of rare complexes is facilitated with the knowledge of their structure and intracellular location. For example, we identified Dam1C/DASH outer-kinetochore complexes based on their localization to the plus ends of kinetochore microtubules, their absence from cytoplasmic microtubules, and their unmistakable structural similarity to recombinant complexes (Ng et al., 2019). To identify subunits within a complex, each subunit can be either deleted or fused with a protein tag and then analyzed by STA (Oda and Kikkawa, 2013; Chang et al., 2016). Density-map subtraction of the
mutant from wild-type densities will then reveal either a “positive” or a “negative” density feature that corresponds to the position of the deleted or tagged subunit, respectively.

Ideally, all copies of a complex should be identifiable without the need of STA or CLEM. This not-yet-invented technology is known colloquially as the “GFP of cryo-EM” and has several requirements. The tag must be a metal nanoparticle, which will be visible as a dense punctum. This tag must also be synthesized in situ without cell permeabilization. Finally, the GFP of cryo-EM must be small and inert enough that it does not compromise the function of the protein of interest. An example of such tags was recently demonstrated in fixed, plastic-embedded cells, which are compatible with in situ gold nanoparticle synthesis (Jiang et al., 2019). Some microorganisms can naturally synthesize metal nanoparticles (Komelli et al., 2006; Scheffel et al., 2006), meaning that the required chemistry is theoretically possible inside living cells. Reproducible, inducible nanoparticle synthesis in living cells will require ideas from inorganic chemistry and materials science. In short, there are exciting opportunities for breakthroughs in cryo-EM in situ labeling.

**STA, classification, and analysis**

Owing to radiation damage, the signal-to-noise ratio of cryotomograms beyond ∼40 Å resolution is very low. If multiple conformationally identical copies of a complex exist, they can be averaged together by STA, yielding a new map that has a higher signal-to-noise ratio. If the complex is conformationally heterogeneous, then some of the conformational states can be “purified in silico” by 2-D and 3-D classification. The combination of STA and classification bridges cell biology and structural biology and is being actively developed (Nicastro et al., 2006; Heymann et al., 2008; Bharat et al., 2015; Galaz-Montoya et al., 2015; Castaño-Diez, 2017; Himes and Zhang, 2018). Work done on HIV capsid subunits showed that subtomogram averages can exceed 4 Å resolution (Schur et al., 2016; Himes and Zhang, 2018). At these resolutions, amino acids with bulky side chains are recognizable and can thereafter be used as landmarks for building and/or refining an atomic model. Lower-resolution subtomogram averages are also valuable because they can be docked with atomic models from crystallography, NMR, or SPA.

Once a SPA or STA experiment produces an averaged structure, the relative translations and rotations between all contributing particles within an image become known. In SPA studies, these alignment data are rarely used for biological interpretations because in most ideal SPA samples, the complexes have random positions and orientations. In cell cryotomograms, however, the 3-D relationships between multiple complexes can encode biological information. To recover this information, the subtomogram average is copied and then remapped into each individual complex’s position and orientation within the cell (Forster et al., 2010). This method has revealed the interactions and distributions of complexes such as ribosomes (Mahamid et al., 2016) and proteasomes (Albert et al., 2017) and higher-order structures of polymers such as nucleosome chains (Cai et al., 2018a).

**Expecting the unexpected**

Cellular cryo-ET data are a rich source of biological surprises. In the recent study of fission yeast cytokinesis, cryo-CLEM clearly detected myosin at the division septum (Swulius et al., 2018). However, the cryotomograms of the myosin-positive positions did not reveal any structures with the size and morphology expected of myofilaments. This finding raises questions about how the actomyosin machinery pinches the two daughters apart. Our study of the Dam1C/DASH complex also revealed unexpected phenotypes, such as the absence of evidence of a globular inner-kinetochore complex coaxial with the kinetochore microtubule and the absence of contacts between Dam1C/DASH and the curved microtubule tips (Ng et al., 2019). Our data also showed that most Dam1C/DASH rings are incomplete, consistent with an earlier calibrated-fluorescence microscopy study (Dhatchinamoorthy et al., 2017). These findings challenged existing models of how the outer kinetochore anchors chromosomes to the plus ends of depolymerizing microtubules for chromosome segregation. As cryo-ET image quality improves, even more unexpected discoveries will be made in situ.

**Studying function in situ**

Because we know so little about the nanometer-level organization of eukaryotic cells, cryo-ET studies are often labeled as “descriptive”—they reveal new phenotypes and they inspire new hypotheses. To use cryo-ET as a more hypothesis-driven tool, one needs to perturb a cellular process and then characterize the phenotypic changes. Genetic perturbations in the form of deletions and temperature-sensitive mutants are available in model organisms. Such mutants may have unexpected phenotypes arising from adaptation to these mutations. To mitigate these potential artifacts, future studies will need to use conditional rapid mislocalization or knockdown technologies, such as anchor-away and the auxin-inducible degron (Haruki et al., 2008; Nishimura et al., 2009).

**STRUCTURAL INSIGHTS FROM THE DEPTHS**

Cryo-EM has come a long way since its birth four decades ago (Frank, 1975; Taylor and Gläscher, 1976; Dubochet and McDowell, 1981; Adrian et al., 1984; Henderson et al., 1986). Cryo-EM studies of eukaryotic cells were historically challenging because cells first had to be cryosectioned. Furthermore, the hardware and software needed for automation were still nascent at the turn of the century (Koster et al., 1997). Cryo-ET of eukaryotic cells only became feasible in 2002 (Hsieh et al., 2002; Medalia et al., 2002). Owing to the challenges of cryomicroscopy, early cryo-ET studies of eukaryotes were done primarily with plunge-frozen samples. These samples include picoplankton, which are the size of bacteria (Henderson et al., 2007), and cell parts thinner than 1 µm (Nicastro et al., 2005; Koning et al., 2008).

The eukaryotic chromatin field has been a major beneficiary of cryo-EM. Eukaryotic genes are packaged into chains of nucleosomes, each consisting of eight histone proteins that are wrapped by ∼147 base pairs of DNA (Kornberg, 1974; Luger et al., 1997). Sequential nucleosomes are thought to fold into higher-order structures that can either occlude or expose DNA sequences to the nucleoplasm. DNA accessibility controls important cellular functions such as transcription and replication, so the central question in the field is: what is the higher-order structure of chromatin (Maeshima et al., 2019)? The dominant model of higher-order structure was the 30-nm fiber, which portrayed nucleosome chains compacted into long helical fibers that have crystalline order. This model was conceived based on electron microscopy images of negatively stained chromatin fragments and X-ray scattering data of oriented chromatin fragments (Finch and Klug, 1976; Widom and Klug, 1985). The 30-nm fiber was not seriously challenged until cryo-EM of cells made it possible to visualize chromatin in situ, without traditional-EM-associated artifacts.

The first major challenge to the 30-nm-fiber model came from cryo-EM studies of sectioned CHO cells (Figure 3A). These pioneering analyses did not reveal any evidence that supported the 30-nm-fiber model in situ (McDowell, 1984; McDowell et al., 1986). Nevertheless, this model persisted, in part because X-ray scattering of
FIGURE 3: Structural cell biology of chromatin. (A) An early example projection image of a cryosectioned CHO cell (McDowall, 1984). Key cytological features include the nucleus (N), chromatin (Ch), nucleolus (Nu), and a nuclear pore complex (P). Cryomicrotomy artifacts such as knife marks (KM) and chatter (W) are also indicated. Image courtesy of A. McDowall. (B) Cryotomographic slice of a *Saccharomyces cerevisiae* nucleus, imaged with “defocus” phase contrast. The boxed region is enlarged twofold and shown in the top-right panel. This position is also rendered as isosurfaces in the bottom-right panel. Adapted from Chen et al. (2016). (C) Cryotomographic slice of a *Schizosaccharomyces pombe* nucleus imaged with “Volta” phase contrast. The bottom-left subpanel shows a threefold enlarged view of a representative nucleosome (white arrow) and a larger multi-MDa complex (megacomplex) in the top-left corner. Both complexes are from the position boxed in the main panel. The bottom-right subpanels are cartoons of the organization of nucleosomes and megacomplexes in G2 and prometaphase (PM) nuclei. Adapted from Cai et al. (2018b). (D) Cryotomographic slice of an embryonic *Drosophila* neuronal nucleus. Heterochromatin is marked out with white dashed lines and the nuclear pore complex (NPC) is labeled. Adapted from Eltsov et al. (2018). (E) Nucleosome remapping in a HeLa cell nucleus. Left: cryotomographic slice of a HeLa cell nucleus. The regions with higher local nucleosome concentrations are marked out with purple dashed lines. The white arrow indicates a nuclear pore complex. Right: 3-D annotation of the same cryotomogram after nucleosome remapping. (F) Arrangement and organization of di- and trinucleosomes in the HeLa cell nucleus. The two nucleosome conformational classes that were “purified” in silico are shaded magenta and blue. E and F were adapted from Cai et al. (2018a).
purified chromosomes showed diffraction features corresponding to regular 30-nm structure; these diffraction features have since been shown to be artifacts from ribosomes that copurified with and coated the chromosomes (Nishino et al., 2012). Furthermore, the data from early cryo-EM studies of cellular cryosections were limited by “1980s state-of-the-art technology,” including the microscope (poor beam stability) and image-recording media (film) (Chang et al., 1983; McDowall et al., 1983). The absence of evidence for abundant ordered helical mammalian chromatin was reproduced in HeLa cells using newer microscope hardware and newer image-processing techniques (Eltsov et al., 2008).

Using cryo-ET, we have shown that chromatin does not pack into any periodic arrays or monolithic masses inside picoplankton (Gan et al., 2013), budding yeast (Chen et al., 2016) (Figure 3B), and fission yeast cells (Cai et al., 2018b) (Figure 3C). Strangely, chromatin is more compact when released intact from cells, without any divalent-cation chelators added, than when confined within nuclei (Cai et al., 2018c). Our studies, considered together with a recent cryo-ET study of cryosectioned insect cells (Eltsov et al., 2018) (Figure 3D), firmly support the notion that chromatin’s irregular structure in situ is conserved. How, then, do nucleosomes chains fold up in situ? In the interphase HeLa cell nuclear periphery, we found that asymmetric nucleosomes follow paths consistent with irregular zigzags (Cai et al., 2018a) (Figure 3, E and F). Much more work is needed to explore the extent of chromatin folding in both transcriptionally active and silenced chromatin.

Cryo-ET studies of transient or rare cellular complexes are more feasible if they can be enriched by either pharmacological or genetic perturbation. For example, an arrest-and-release strategy was used to enrich for fission-yeast cells that have actively constricting actomyosin rings (Swulius et al., 2018). The division septa were then localized by cryo-CLEM (Figure 4A). In these dividing cells, dense bundles of actin filaments were located peripheral to—but not in contact with—the division septum (Figure 4, B and C). We have used both drugs and cell-cycle mutants to arrest picoplankton and budding yeast, respectively, in metaphase. In picoplankton, we found that the unusual spindle has fewer microtubules than chromatomes (Gan et al., 2011) while in budding yeast, we found that the outer-kinetochore Dam1C/DASH complex rarely (if ever) contacts the curved tips of the kinetochore microtubules (Figure 4, D–F) (Ng et al., 2019). The vast repertoire of well-characterized yeast mutants needed to explore the extent of chromatin folding in both transcriptionally active and silenced chromatin.

Many interesting complexes are extremely challenging to study in situ. Such complexes are too rare, or too transient, or have both conformational and constitutional heterogeneity, or are not monolithic, or a combination of the above. Nevertheless, these “biochemist’s nightmare” properties often underlie their function and mechanisms. Examples of such challenging complexes—if they can be called complexes at all—include signalosomes, focal adhesions, neuron synapses, membrane-membrane contact sites, chromatin, and kinetochores. To make such complexes easier to study, cryo-ET needs to be much faster. New tomography hardware and imaging workflows will make tilt-series collection ten times faster (Chrefi et al., 2019; Eisenstein et al., 2019), potentially without sacrificing resolution. Other steps in the workflow will need to speed up: cryo-amellae preparation, cryo-CLEM localization, and reconstruction. Recently, researchers have indeed reported advanced automation in cryo-FIB (Buckley et al., 2019; Zachs et al., 2019) and the integration of cryo-FIB milling and light cryomicroscopy in a single instrument (Gorelick et al., 2019). Image processing and analysis will also become faster as researchers exploit the massive parallelism of consumer graphics cards and multicore processors (Kimanius et al., 2016; Castaño-Díez, 2017; Himes and Zhang, 2018; Zivanov et al., 2018).

How would one use high-throughput cryo-ET to answer the most challenging questions? Consider the kinetochore—a large multi-functional complex central to chromosome segregation. Kinetochore complexes connect chromosomes to the spindle apparatus, detect and correct erroneous attachments, delay mitosis until all chromosomes are correctly attached to the spindle apparatus, and couple spindle-microtubule depolymerization to poleward-directed chromosome motion. Kinetochores have dimensions in the tens to hundreds of
nanometers and contain multiple elongated subassemblies (Jenni et al., 2017). Recently, many high-resolution cryo-EM structures of reconstituted kinetochore subcomplexes were published (Penta-kota et al., 2017; Chittori et al., 2018; Jenni and Harrison, 2018; Leber et al., 2018; Yan et al., 2018; Zhang et al., 2019; Hinshaw et al., 2019; Hinshaw and Harrison, 2019; Yan et al., 2019). Yet, very little progress on the kinetochore’s in situ structure(s) has been made. Using a combination of serial cryo-ET of cryosections, with kinetochore microtubules as macromolecular signposts, we recently characterized the Dam1C/DASH complex—just one piece of the yeast outer kinetochore (Ng et al., 2019). The in situ structure of the rest of the kinetochore remains unknown.

We now speculate on how high-throughput cryo-ET, combined with a “divide, conquer, and unite” subtomogram-analysis strategy, will reveal kinetochore structures in situ. In this exercise, we wishfully assume that the just-discussed microscopy bottlenecks will be solved by automation and improved workflows. Our starting point is a collection of thousands of cryotomograms of budding yeast cryolamellae, each containing a few kinetochores that were automatically localized by cryo-CLEM. The first step is to extract subtomograms that overlap with the light cryomicroscopy signals (Figure 6A). Identification of the kinetochore subassemblies will be guided by our knowledge of their relative positions (Joglekar et al., 2009; Aravamudhan et al., 2014). Small subtomograms that contain these
subassemblies will next be classified into their various conformational states (Figure 6B). We imagine that some kinetochore subassemblies have only a few translational and rotational degrees of freedom relative to kinetochore's center of mass, analogous to side-chain rotamers in protein structures. In contrast, there are also some flexible, elongated subassemblies such as the coiled–coil-rich Ndc80 complex, which gives the kinetochore its gossamer nature. Flexible subassemblies like Ndc80 are expected to have many conformations and would therefore produce the lowest-resolution models. Next, each kinetochore will be reassembled in silico by remapping the subtomogram averages of each subcomponent (Figure 6C). Kinetochores pieced together this way will have many components remapped with subnanometer accuracy. Remapped kinetochores from spindles with and without tension will reveal a wealth of information. We will learn which complexes undergo the largest conformational changes, what conformational changes are sensitive to spindle pulling forces, how checkpoint complexes dock in mitotically arrested cells, if detached kinetochores have the same conformation as tensionless kinetochores, and if anaphase kinetochores attach to the spindle the same way as metaphase ones. These in situ kinetochore structures will stimulate even more (presently unimaginable) questions.
FIGURE 6: Structural cell biology of floppy complexes. This schematic shows how a divide, conquer, and unite strategy can reveal kinetochore structures in situ. (A) Cartoons of cryotomograms (thin gray slabs; tomo1, tomo2, etc.) of thinned yeast (rounded gray bodies). In this example, the kinetochores are first localized by fluorescence cryomicroscopy (green signal). Cryo-CLEM greatly facilitates the identification of subtomograms that contain kinetochores. (B) Alignment and classification of smaller subtomograms that contain kinetochore subassemblies. Owing to the flexibility and conformational heterogeneity of the kinetochores, each subassembly must be windowed and then tracked throughout the alignment and classification process. Subassemblies that have similar conformations are aligned and averaged, producing a higher signal-to-noise ratio “class average.” In this schematic, the outer kinetochore, inner kinetochore, and centromere-associated complexes are colored green, violet, and blue, respectively. (C) These class averages can then be rotated and translated to the orientations and coordinates of each copy in their in situ context at the tips of kinetochore microtubules (gray). This remapping approach can deal with “floppy” complexes as long as some of the subassemblies are monolithic.

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