INTRODUCTION

A variety of bio-materials and proteins of different sizes and shapes have been studied using transmission electron microscopy (TEM). In this technique, high voltage electrons are used as an illumination source to produce a projected image containing details of the internal and external features of biological structures. The electron beam interacts with the sample under a vacuum as it passes through the entire thickness of the specimen, and contrast is generated by the scattering of electrons (Nagayama & Danev, 2004; Massover, 2008; Murata et al., 2010; De Carlo & Harris, 2011). Biological structures are easily distorted by dehydration caused by the vacuum of the microscope and radiation damage caused by the electron beam; hence, negative staining, a special specimen preparation technique, is commonly used to obtain sufficient contrast and preserve the native structure as much as possible (Brenner & Horne, 1959; Monroe & Brandt, 1970).

Using negative staining, the background is stained rather than the sample leaving it unstained; therefore, the image is formed from the difference between the non-stained area and the surrounding stained area. Good contrast is achieved when the stain surrounds the sample completely. Scattering of the electron beam by an object creates contrast; however, protein structures comprised of light atoms scatter the beam weakly and give poor contrast. Negative staining acts as a support to stabilize the biological structures, and enhances contrast to produce a well-defined electron density of the sample. Consequently, the electrons are scattered strongly by the stain and the TEM image is generated as a result of the contrast between differential scattering by the stain and the sample during the period of illumination.
The advantages of negative staining over high resolution electron microscopy (EM) approaches, such as cryo-EM, are its convenience, simplicity, rapid applicability to biological objects ranging from individual proteins to large macromolecular assemblies, and ability to detect structural homologies (Anderson et al., 1989; Boekema et al., 2009; Zhang et al., 2013); hence, negative staining is used widely to detect the structural features of bio-materials and small proteins. To obtain the best contrast between stain and protein, the electron beam exposure time, which is related to the thickness of the support film, is an important parameter to consider.

Here, we investigated the effects of short and long electron beam exposure times on the TEM outcomes of negatively stained samples. We also describe technical procedures to identify the optimal exposure time that produces higher electron densities during TEM imaging. Single particle image averaging (Umeki et al., 2009; Jaitly et al., 2010) was used to process and compare the structural details obtained from a set of TEM images taken at three different times. Overall, the results suggest that longer exposures (2–4 seconds) provide higher electron densities.

MATERIALS AND METHODS

Negative Staining Electron Microscopy

Negative staining was performed using purified thyroglobulin prepared from a protein stock (T9145; Sigma-Aldrich, USA). The stock protein was dissolved in phosphate-buffered saline (10 mM phosphate buffer, 2.7 mM KCl, and 137 mM NaCl; pH 7.4) and diluted with 20 mM Tris-HCl buffer (pH 7.5) to a final concentration of 100 nM. A 5 μL aliquot of the diluted protein sample was applied to a carbon-coated grid that had been glow-discharged (Harrick Plasma, USA) for 3 minutes in air. The grid was immediately negatively stained using 1% uranyl acetate (Jung et al., 2008; Jung et al., 2011; Elliott et al., 2012; Lee et al., 2015), and then examined using a Tecnai G2 Spirit Twin TEM system (FEI, USA) that was fitted with an anti-contaminator and operated at 120 kV. The analysis was performed using instrumentation at the Korea Basic Science Institute. Images were recorded using a 4×4 K Ultrascan 895 CCD (Gatan, USA) at a magnification of 30,000 (0.36 nm/pixel). Micrograph images were collected from the same sample area, but different exposure times were used to produce three micrograph datasets. The total dose of the sample exposed to the electron beam for 1 second was approximately 10^5 electrons/nm^2 at a nominal underfocus of 2.5 μm. Micrographs were also taken after longer exposure times of 2 and 4 seconds.

Comparative Analysis of Single Particle Image Processing

Images of individual particles on the micrograph were selected interactively, windowed out, and imported into the SPIDER program suite (Health Research Inc., USA). To compare the electron densities produced after different exposure times, the same coordination profile obtained from the selection of particles in the 1 second exposure time micrograph was used to window out particles from the 2 and 4 seconds exposure time micrographs. A total of 200 well-separated particles were used in the alignment and classification, which were performed using the reference-free method described previously (Burgess et al., 2004). All subsequent single particle image processing was carried out using windowed datasets.

![Fig. 1. Transmission electron microscopy images of the same sample site after three different electron beam exposure times. Negatively stained fields of protein samples exposed to the electron beam for 1 second (A), 2 seconds (B), 4 seconds (C). The arrowheads indicate representative molecules. Isolated thyroglobulin was used for the specimen preparation and 200 particles in each field were windowed for further analysis. Scale bar=100 nm.](image-url)
with the same alignment and classification parameters. A subtraction procedure was used to compare the structural differences between two 100×100 pixel averaged images processed from the same particles but obtained from electron densities resulting from different exposure times. The shorter exposure time image was subtracted from the longer exposure time image; the resulting subtracted image showed lighter shading in regions where the shorter exposure time image contained more density than the second image.

RESULTS AND DISCUSSION

TEM Imaging of Negatively Stained Bio-Materials Exposed to an Electron Beam for Different Lengths of Time

TEM images of negatively stained thyroglobulin (approximately 660 kDa) were collected after the same sample site was exposed to the after electron beam for 1, 2, or 4 seconds (Fig. 1). In all three images, the molecules existed as appeared egg-shaped forms with lengths of 24~26 nm and widths of 11~12 nm (You et al., 2013). The appearances of the individual molecules after the different exposure times were indistinguishable (indicated by arrowheads in Fig. 1).

Topographical Comparisons of the Averaging of Molecules

Single particle image processing, digitized at a pixel size corresponding to 0.36 nm, was used to compare the electron densities of the TEM images obtained using the three different exposure times. Individual molecules in the TEM images were selected interactively and windowed out into 100×100 pixels for processing, including alignment and classification. The

![Fig. 2. Iterative alignments of transmission electron microscopy images after three different electron beam exposure times. Nine iterative alignments (left to right) of datasets after exposure for 1 second (A), 2 seconds (B), or 4 seconds (C). The alignments were performed with 200 particles in each dataset. Scale bar=40 nm.](image)

![Fig. 3. (A-C) Windowed particles used in single particle analysis and their averaged images. Selected individual particles windowed from negatively stained fields (upper panels) and averaged images (bottom panels) from the three different exposure time datasets. The asterisks indicate the averaged image showing detailed densities that were selected for further comparative analyses. Each class average contained 15~25 individual particles. Scale bar=40 nm.](image)
datasets were independently aligned with iterative alignments using the same parameters. The aligned images appeared very similar, but subtle differences between them were identified (Fig. 2). Fig. 3 shows windowed individual particles in each dataset and class averages processed from their alignments. A comparison of the raw images of the individual particles with the averaged images revealed a vast improvement in the level of detail after averaging. The averaged images from the 1 second exposure time dataset did not correspond well to those from the 2 and 4 seconds datasets (Fig. 3), indicating that the aligned particles provided different electron densities resulting in different class averaging details.

Detection of Additional Electron Densities Induced by Longer Exposure Times
To clarify and detect recognizable differences between the electron densities of the averaged images taken from the 1, 2, and 4 seconds datasets, the constituent images of selected averages showing structural details with similar oriented views (asterisk-marked averages in Fig. 3) were segregated from their original datasets. The same particle numbers and the same molecules were reassembled into the averages (Fig. 4), with the aim of avoiding subjectivity in determining the differences between the electron densities. Although the same molecules were used from each dataset, structural details between the averaged images were identified.

To reveal additional electron densities, averaged images of longer exposure time samples were subtracted from those of the shorter exposure time samples (Fig. 5). The resulting subtracted images showed lighter shading distributed within the averages (indicated by arrowheads in Fig. 5B), reflecting more electron densities in the images from the 4 seconds exposure time samples. This finding indicates that a longer exposure time (4 seconds) resulted in additional electron densities in the averaged images, possibly resulting in better alignment and class averaging.

Single bio-materials, such as protein molecules, have low molecular weights; hence, the information obtained from TEM imaging of each particle is limited. Previous studies (Henderson, 1995) using theoretical considerations demonstrated that unstained molecules larger than approximately 100 kDa can be reconstructed to provide structural details. This molecular weight limit does not apply to negative staining TEM of proteins as well as cryo-EM (Kim et al., 2015); therefore, the use of correct instrumentation and optimized analytical parameters can maximize the quality of
the structural densities obtained, potentially aiding structural
determination of bio-materials at the molecular level.

CONCLUSIONS

The results presented here suggest that longer electron beam
exposure times provide more electron densities of bio-
materials analyzed by TEM imaging, ultimately resulting in
optimal visualization of their detailed structural features.
In general, generating raw images containing sufficient
contrast and electron density is a fundamental requirement
of morphological and structural analyses; thus, the evaluation
procedures described here are a useful tool for maximizing
the electron densities of TEM images for structural determi-
nations of bio-materials at the molecular level.

CONFLICT OF INTEREST

No potential conflict of interest relevant to this article was
reported.

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