Three-Dimensional Analysis of Melanosomes Isolated from B16 Melanoma Cells by Using Ultra High Voltage Electron Microscopy

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ABSTRACT

Melanosomes, isolated by centrifugal separation from culture broth of B16 melanoma cells derived from mouse, were observed by scanning electron microscopy (SEM), and by transmission electron microscopy (TEM). Some interesting structural features were found inside and outside of the melanosomes. By SEM observation, the melanosomes were ellipsoid shape, their surface was not smooth and was covered with rough substructure, 10 to 20 nm particles. By TEM, uneven structure and micro particles were observed in the melanosomes. Furthermore, three-dimensional analysis was tried by using the ultra-high voltage electron microscopy (UHVEM). Micrographs of the melanosomes were taken at various tilted angles by UHVEM, after preparing 500 nm thickness specimens stained with lead citrate. From the micrographs collected, the three-dimensional structures were reconstructed by using i-mode software. Melanin stained by lead and non stained parts was clearly observed in the reconstructed structure. Non stained parts were round, regular size, and distributed widely in the melanosomes.

KEYWORDS

B16 Melanoma; Melanosome; Scanning Electron Microscopy; Transmission Electron Microscopy; Ultra High Voltage Electron Microscopy; Three-Dimensional Structure

1. Introduction

Melanosomes are specialized organelles in pigment cells called melanocyte controlling melanin biosynthesis. The matured melanin is transferred from a pigment cell to neighboring cells, such as keratinocytes. Melanocytes are neural crest-derived cells that exist in the substantia nigra of brain, uveal tract including iris of eye, cochlear duct of ear and basal layer/hair follicle of skin. Melanin contributes to the determination of hair and skin color, the biological defense to ultraviolet rays, etc. in humans.

As the surface of the melanosome is central to its functions [1,2], elucidation of the pigment composition of the surface is necessary. Some reports previously have described the three-dimensional ultrastructure of melanosomes [1,3-5]. It is thought that each melanosome consists of structural proteins such as melanin, enzymes related to melanization and carrier proteins of melanin [6]. Melanin can be classified into dark black eumelanin and light red pheomelanin. Melanosomes which synthesize predominantly eumelanin are called eumelanosomes, while those that synthesize predominantly pheomelanin are called pheomelanosome. Surface substructure, amino acid contents, metal composition and PTCA/PDCA com-
position in eumelanosomes/pheomelanosomes and enzymatic/acid/base were reported [4,5].

Since the molecular structure of melanin polymer was not clear, the detailed inside ultrastructure of melanosome was also still unidentified. In particular, it is unknown where biosynthesis of melanin is performed inside of melanosome. Controlling biosynthesis pathway of melanin is an important theme in the dermatological field and the beauty field. It is thought that the understanding of the inside ultrastructure may lead to the elucidation of the melanization mechanism.

We have examined the internal and external structure of melanosomes. For this purpose, we have used ultra-high voltage electron microscopy imaging (Osaka Univ.) besides normal transmission electron microscopy imaging. This huge electron microscope makes it possible to obtain a three-dimensional ultrastructure profile [7,8]. Reports about inside ultrastructure of melanosome using this three-dimensional electron microscopy have not been reported.

In our first attempt we focused on B16 melanoma cells (B16) derived from mouse.

As a part of investigating the characteristics of melanosomes themselves and their role in controlling the pigment system, we purified melanosomes and successively examined their ultrastructure. B16 cells were cultured, homogenized and melanosome containing fraction was isolated by density difference centrifugation method. We demonstrate characteristic features about the ultrastructural imaging of extracted and isolated melanosomes.

2. Materials and Methods

2.1. Regents

DMEM; Dulbecco’s modified Eagle’s medium (GIBCO).
FBS; Fetal bovine serum (Equitech-Bio).
EDTA; Ethylenediaminetetraacetic acid (Sigma).
Trypsin (Sigma)
Penicillin (GIBCO).
PBS; Ca-Mg-free phosphate buffer saline (Sigma).
HEPES; N-2-hydroxyethylpiperazine-N-2-ethane sulfonic acid (life technologies).

2.2. Strains and Culture Condition

The strain of B16 melanoma derived from mouse, kept as a stock strain at the department of tissue and organ development regeneration of Gifu University, Japan, was used in these experiments. The culture condition of B16 melanoma cells was referred to the Riken’s protocol (http://www2.brc.riken.jp/lab/cell/detail.cgi?cell_no=RCB0557&type=1). The culture of B16 melanoma cells was performed in a 10-cm dish containing DMEM supplemented with 10% FBS and penicillin by using CO₂ incubator (ASTEC, Japan) at 37°C. The cultured melanoma cells which formed a layer on the bottom of the dish could be recovered by addition of drops of 0.25% trypsin, 1 mM EDTA in 0.7× PBS. Same DMEM medium successively was added.

For the subculture, the cell suspension was diluted 3 to 6 times of the original concentration with the same DMEM medium. After several times of subcultures, melanoma cell layers on the bottoms are recovered as cell suspension by the same procedure as above. Medium was removed and collected by decantation and washed once by 1× PBS. Melanoma cells were treated with drops of a solution containing 0.25% trypsin, 1 mM EDTA in 0.7× PBS for removal of the cells from the dish.

The subcultured dishes finally increased from 30 to 50 dishes, before the following melanosome preparation step.

2.3. Preparation 1 of Melanosome

The melanoma cell suspension was centrifuged for 5 min. at 700× g, 4°C to remove supernatant. The melanoma cell pellet was washed with 0.25 M sucrose in 10 mM HEPES buffer, and was centrifuged for 10 min at 1000× g, 4°C.

Pellets were resuspended in HEPES buffer and homogenized with a glass/glass homogenizer (Dounce Tissue Grinder, WHETON, USA) in 0.25 M sucrose in 10 mM HEPES buffer, on ice using 25 strokes. Homogenates centrifuged 10 min at 1000× g, 4°C to remove pellets containing unbroken cells and nuclei. Supernatants were collected and ultra-centrifuged 45 min at 20,630× g, 4°C.

After centrifugation, pellets including melanosomes were precipitared [9,10]. Pellets were suspended twice by distilled water and preserved at -80°C, temporarily. Finally, these were freeze-dried and sealing in glass tubes following preservation long term.

2.4. Preparation 2 of Melanosome

Cultured medium was separately collected and centrifuged for 5 min at 10,000× g, 4°C to precipitate melanosomes released from B16 into its medium. These pellets were suspended twice by distilled water and preserved at -80°C. These also were freeze-dried and sealing in glass tubes following preservation.

2.5. Electron Microscopies

2.5.1. Scanning Electron Microscopy (SEM)

Field Emission-SEM JSM-6330F (JEOL, Japan) was
used to examine external structures in high vacuum mode. The freeze-drying samples were mounted on aluminum specimen mount using electrical conductive double-sided tape. The mounted freeze-drying samples were coated platinum sputtering using ESC-101 (ELIONIX, Japan) before observation.

2.5.2. Transmission Electron Microscopy (TEM)

TEM, JEM-1010 (JEOL, Japan) and H-7100 (HITACHI, Japan) were used to examine internal structures. The freeze-drying samples were embedded with Epoxy resin after fixation and dehydration. Usual thin-sections were stained with double-stain of Uranyl acetate dihydrate/Lead citrate or with Lead citrate only.

2.5.3. Ultra-High Voltage Electron Microscopy (UHVEM)

UHVEM was used to examine three-dimensional internal structures. UHVEM (H-3000, HITACHI, Japan) had 3 million volts for acceleration voltage and was located at Research center for Ultra-High Voltage Electron Microscopy, Osaka University. The freeze-drying samples were fixed with normal Epoxy resin. Thin-sections 1 μm thickness, and stained with Lead citrate in 10 min incubation at room temperature.

UHVEM acquired images while tilting thin-sections. Single tilt series were recorded over an angular range of 120 degrees (-60 to +60 degrees). And 60 TEM images sequentially acquired at 2 degrees intervals. Figure 1 gives the graphic representation of this principal for acquisition of three-dimensional Images [11]. TEM images were recorded using a 4096 × 4096 pixel slow scan CCD camera (TVIPS, Germany). These tilt series were aligned by the etomo in the IMOD software package (http://bio3d.colorado.edu) using the gold particles, and tomograms were generated. Three-dimensional projection images were generated by Image-J (http://rsbweb.nih.gov/ij/), image-analysis software. These softwares can make animation files (video clip, avi file) of tomograms and projection images.

The software (an auto contour and imodauto, which is a threshold-based algorithm) was generated the information of interface between one material and another material. This can visualize the difference of non-specific staining contrast in electron microscopy sample as internal structure and contents [8].

3. Results and Discussion

B16 melanoma cells were cultured and subcultured several times according to the Riken’s protocol. The cultured cells were harvested for the following preparation step.

The method for preparation of melanosomes from B16 was adapted from previously described method [9,10]. B16 cells were harvested with trypsin/EDTA and collected by centrifugation. After suspension in 0.25 M sucrose in 10 mM HEPES buffer, cells were homogenized with a Dounce glass/glass homogenizer. Unbroken cells and nuclei were removed by centrifugation. The supernatants were collected and ultra-centrifuged 45 min at 20,630 × g, 4°C.

Harvested black pellets were collected as a suspension of B16 melanosomes in water and freeze-dried (Figure 2(a)). Since the samples after freeze-drying are treated like powder, it is thought that samples looked gray or whitish by diffused reflection (Figure 2(b)). B16 secreted melanosomes fraction (Materials and methods 2.4.) are also observed that same color as isolated B16 melanosomes fraction (Figures 2(c), (d)).

Figure 3 shows SEM micrographs of melanosomes isolated from B16. Almost all particles (melanosomes) are ellipsoidal shape. Diameter of particles is long axis of 300 - 400 nanometers and short axis of 200 - 300 nanometers. Its surface is not smooth and uniform but uneven appearance. Diameter of each uneven substructure is 10 - 20 nanometers.

Figure 4 shows SEM micrographs of melanosomes released (secreted) naturally from B16 to cultured medium. The size and surface structure of released melanosomes are same range and same shape as isolated one with homogenized procedure. About its shape, there are many round shapes in released melanosomes compared with isolated melanosomes.

B16 melanoma cells naturally secrete mature melanosomes to outside of cells. The cultured melanoma cells release melanosome to own medium. The features of their shape and surface substructure were nearly same between isolated melanosomes and released melanosomes. Collection of medium was carried out at every subculture. Naturally released melanosomes are regarded as advanced stage IV. It is unknown whether same entire melanosomes release to keratinocyte in vivo. The yield of released melanosomes was very small, and fluctuated by every subculture. Application of stimulative factors (UV irradiation or chemicals) for stimulation of releasing melanosomes is required.

In this report, isolated melanosome containing fraction included amorphous materials around melanosomes, although these amorphous materials were able to be removed by ethanol treatment. These amorphous contaminations are considered to be residual substance consisting nucleus or proteins or unbroken parts of cell, etc.

Figure 5 shows TEM micrographs of melanosomes isolated from B16. Some unstained parts (small white
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Figure 1. A simplified schema showing the acquisition of sequential images to get three dimensional data by using Ultra-High Voltage Electron Microscopy (UHVEM) at Osaka Univ. Specimen: thin sections for UHVEM. Tilting angles: +60 to −60 degrees. Interval of acquisition: 2 degrees.

(a) Isolated B16 melanosomes pellets before freeze-drying. (b) Isolated B16 melanosomes samples after freeze-drying in the sealed glass tube. (c) Secreted B16 melanosomes pellets before freeze-drying. (d) Secreted B16 melanosomes samples after freeze-drying in the sealed glass tube.

(a) SEM micrographs of isolated B16 melanosomes fraction by the preparative method of 2.3. Preparation 1. Bar = 100 nm.
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Figure 4. (a), (b) SEM micrographs of B16 secreted melanosomes fraction by the preparative method of 2.4. Preparation 2. Bars: (a) = 100 nm, (b) = 1 μm.

Figure 5. (a), (b) TEM micrographs of isolated B16 melanosomes fraction by the preparative method of 2.3. Preparation 1. Thin sections staining: Lead citrate. Bars = 100 nm.

particles) are observed inside almost all melanosomes. These seem to exist uniformly from the central region to the peripheral region. Size of each unstained part is 10 - 20 nanometers.

SEM observation would be probing smaller ultrasubstructures along the surface of melanosomes. TEM observation would be probing unstained parts in the melanosomes. As both ultrasubstructure (size of 10 - 20 nanometers and uniform distributions) are comparatively similar, some kind of association may be suggested.

In the observations of TEM and UHVEM, the influence of amorphous contaminations (nucleus or proteins or unbroken parts of cell, etc.) which observed in SEM images was reduced by single stain of lead citrate. Lead citrate stains melanin only.

60 original photographs taken from UHVEM were processed with image-analysis software to generate three-dimensional images. These are outputted as projection animation files (video clip, avi file). Figure 6 shows projection images of melanosomes isolated from B16. These 6 still images ((a)-(f)) were picked out by regular intervals from sequential 60 original projection photographs. 10 to 20 nm size unstained parts exist widely throughout in melanosomes. Size of these is the same as that of normal TEM images.

Figure 7 shows tomogram (three-dimensional TEM images) of melanosomes isolated from B16. These 6 still images ((a)-(f)) were extracted at regular time intervals from whole tomogram animation (video clip, avi file). Small unstained parts exist every section of tomograms.
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Figure 6. (a)-(f) UHVEM, projection sequential micrographs of isolated B16 melanosomes fraction by the preparative method of 2.3. Preparation 1. Thin sections staining: Lead citrate. These 6 still images were picked out by regular intervals from sequential 60 original projection photographs. Bar = 100 nm.

Figure 7. (a)-(f) UHVEM, tomogram sequential images of isolated B16 melanosomes fraction by the preparative method of 2.3. Preparation 1. Thin sections staining: Lead citrate. These 6 still images were extracted by regular time intervals from whole tomogram animation. Bar = 100 nm.

Figure 8 shows three dimensional images that indicate the small unstained particles in one melanosome. These 3 still images ((a)-(c)) were extracted at random from whole contour animation. Red lines of coloring by auto contour software correspond to interface between melanosome and outside. Green lines of coloring by auto contour software correspond to interface between the unstained particles and another stained part. It is thought that two kinds of substances in melanosomes are different in their physical properties.

Equal and widespread distribution of unstained parts
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Figure 8. (a)-(c) UHVEM, three dimensional contour images of isolated one B16 melanosome generated using specific software. These 3 still images were extracted at random from whole contour animation.

inside the melanosomes was able to be confirmed by the animation (projection images and tomograms) of UHVEM. There was distributive variation in each melanosome. Some melanosomes had many intrinsic unstained parts. Such variation may be relating mature level of melanosome, difference of stage and/or eumelanin/pheomelanin ratio.

Since the details of the molecular structure of the melanin itself are unknown, little information about three-dimensional ultrastructure of melanosomes is available. Some reports previously have been described three-dimensional ultrastructure of melanosomes from iridal stroma, retinal pigment epithelium and human hair [1,3-5].

Free electron laser photoemission electron microscopy data is attributed to eumelanin on the surface of human iridal stroma melanosomes [1]. The surface of the iridal stroma melanosomes is comprised of smaller uneven ultrasubstructures of a few tens of nanometers. Another report shows that eumelanin premelanosomes comprise zigzagging longitudinal strands with cross-links that occur every 20 nm [12]. From this, they conclude that these surface ultrasubstructures of melanosomes have eumelanin coats. These substructures were found on the surface of the iridal stroma melanosomes all have eumelanin coats [1].

As for a hair, black and red human hairs were analyzed for their structural, chemical and physical properties [4,5]. The black-hair melanosomes are ellipsoidal and maintain structural integrity. The red-hair melanosomes are less regularly shaped and are generally smaller than black hair and tend to fall apart. A single black-hair melanosome have packed substructures forming a relatively smooth outer surface. Red-hair melanosomes exhibit a rougher surface, with larger substructures [4]. Eumelanosomes are rod-like and pheomelanosomes are round and more spherical [13]. New born melanosomes are short and round and grow to form rod shaped aging [14].

It is suggested that many B16 melanosomes are eumelanosome type since they are ellipsoidal shape and have small substructures forming uneven outer surface. But there are some more spherical shapes compared with human hair melanosomes which are sharp rod-like. It is necessary to examine ratio of existence of eumelanine and pheomelanin from now on. Recently, chemical analysis has been established [15].

For these examinations, comparative studies of other melanomas or normal melanocytes from various origins may be effective. Melanosomes are classified into four stages according to their maturity. It is said that internal structure of melanosome begins to be formed at stage I, and is completed at stage II. Tyrosinase activity appears and melanization begins at stage, and mature melanosome is completed at stage IV. In this report, it is thought that isolated melanosomes are mature melanosomes because inside melanin of isolated melanosomes was stained and outside structure was nearly the same as released melanosomes. Early stage (I, II) and mature stage (III, IV) of melanosomes could be collected separately using density-gradient centrifugation method [9]. We are considering examining the difference of stages, origins (ex. human races, aging change) about ultrastructure of melanosome.

The constituent of unstained parts has not been analyzed yet. Preliminary analysis by ICP-MS (PerkinElmer, ELAN DRC II) after acid digestion pretreatment showed the presence of Cu in melanosome containing fraction sample isolated from B16 (data not shown). Tyrosinase analogues are the rate limiting enzyme binding Cu for biosynthesis of Dopa. The minimum detectable mole fraction of Cu ion was detected in retinal pigment epithelium melanosomes of rats by energy dispersive X-ray microanalysis [16]. In human hair, abundant Cu ion was detected, and the content of black hair was two times higher than red hair [4].

Further analysis should be necessary for localization of
Culike specific contents in melanosome particle.

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