Ultrastructural studies on actin-like filament in mung bean mitochondria and its potential functional significance

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(Received July 9, 2010; Accepted August 3, 2010)

ABSTRACT. This document reports that actin-like filaments, about 5 nm in width, were found in mung bean mitochondria after negative staining of mitoplast. The 5 nm filaments, composed of globular proteins, were visualized beneath the intramembranous particles by freeze-fracture electron microscope. Freeze-fracture immuno-labeling against anti-actin antibody suggested that actin-like filaments attached to the protoplasmic surface of mitochondrial inner membrane. These filaments were also found in the matrix and underneath of outer membrane. The presence of actin-like protein in mitochondria was also substantiated by immunolocalization in situ. Interestingly, some 5 nm filamentous bundles were often found between mitochondrion and its “bud”. It is likely that these images represent an undiscovered novel plant mitochondrial budding process which may involve some 5 nm filamentous structure in separating nascent mitochondrion to its mother mitochondrion. Treatment of cultured tobacco cells with F-actin depolymerization reagent, Latrunculin B (Lat B) might affect the shapes of mitochondria. Taken together, this study suggests that some 5 nm filaments, probably F-actin, may be localized in mitochondria and play a role in mitochondrial propagation and mitochondrial shaping.

Keywords: Actin/actin-like filament; Actin dynamics and mitochondrial morphology; Mitochondrial propagation; Vigna radiata.

INTRODUCTION

It has been well-investigated recently that actin-like filament in bacteria is responsible for bacterial shape, cell division and DNA segregation (Jones et al., 2001; ven den Ent et al., 2001; Carballido-Lopez, 2006; Graumann, 2007; Pogliano, 2008). In vitro biochemical study indicates that MreB can self-assemble into actin-like filaments. The progenitor of mitochondria was an α-proteobacterium and it is rational to assume that mitochondria might carry a MreB/actin-like protein involved in mitochondrial propagation and highly dynamic shape keeping. The presence of actin and tubulin in mitochondria and the association of actin with mitochondrial DNA has been reported but has not been further verified before 2011 (Etoh et al., 1990; Carre et al., 2002; Lo et al., 2002; Dai et al., 2005; Wang and Bogenhagen, 2006). The existence of actin inside mitochondria was verified by Reyes et al. (2011) and Lo et al. (2011) in human and mung bean mitochondria, respectively. After the issue of actin and actin filament’s presence in mitochondria was evidenced, a more focused effort in searching for its functional significance will contribute greatly to the largely unknown area on plant mitochondrion in terms of its DNA segregation, propagation, morphological alteration and macromolecular transportation inside mitochondrion.

FtsZ, a progenitor to tubulin, forms the cytoskeletal framework of cytokinetic ring in bacteria and plays a major role in constriction of the furrow at septation of prokaryotes. Arabidopsis chloroplasts were shown to import FtsZ in vitro (Osteryoung and Vierling, 1995), which could mediate their division. It appears that most mitochondria studied thus far have replaced FtsZ with dynamin-related proteins for fission (Smirnova et al., 1998; Bleazard et al., 1999, Sesaki and Jensen, 1999; Erickson, 2000; Arimura and Tsutsumi, 2002; Logan, 2006), though mitochondria of a few alga still use FtsZ for division (Beech et al., 2000). Unlike chloroplasts, mitochondria are dynamic organelles that often form a complex membrane system consisting of interconnected tubular structures, which undergo frequent fusion and fission.

We document in this report that a novel filamentous structure resembling actin filaments was found in mung bean seedling mitochondria. The 5 nm-wide filaments were visualized beneath the intramembranous particles (IMPs) by freeze-fracture electron microscope, thus placing the filaments on the protoplasmic surface of the mitochondrial membrane. This evidence for the presence of actin-like protein in isolated mung bean mitochondria was further
Mitochondria and mitoplasts preparation

Mung bean mitochondria were prepared from 3-day-old etiolated mung bean (*Vigna radiata*, TN-5) seedlings and purified by a sucrose gradient, as described by Dai et al. (2005). Mitoplast preparation for electron microscopic examination followed the methods of Kozlowski and Zagorski (1988).

**Fine structure and in situ immunolocalization analysis**

The procedure for immunolocalization in tissue sections was performed as described previously (Dai et al., 1998). Tissue sections (100-120 nm) were incubated with anti-actin (C4 clone, Chemicon) antibody followed by goat-anti-mouse antibody conjugated with 12 nm gold particles. The monoclonal antibody PM028 (GTMA) against a yet-unidentified maize mitochondrial protein, with no cross-reactivity to mung bean mitochondria, was used as a control in the immuno-localization study.

EM analysis on whole-mount mitoplasts was carried out by fixing the sample with 1% paraformaldehyde followed by 1% uranyl acetate (pH 4.5) staining for 1-2 min. For all fine structure analysis, the experiments were repeated 3-6 times and similar results were consistently obtained.

**Freeze Fracture and immunogold labeling study**

Purified mitochondria were placed on the specimen carrier and quick-frozen by nitrogen slush. The freeze fracturing was carried out in a BAF 400D freeze etch unit (Balzers Union, Liechtenstein) at -105°C. Replicas were made by evaporation of platinum-carbon from an electron-beam gun positioned at a 45° angle, followed by carbon coating from a 90° angle. The replicas were released from the specimen carrier by immersing in 0.1% paraformaldehyde followed by 0.05% SDS for 10 minutes at room temperature. Replicas were then washed four times with PBS. The immunogold labeling of anti-actin antibody was then carried out as described by Fujimoto et al. (1996) with minor modification. Anti-actin (C4 clone, Chemicon) antibody, followed by goat-anti-mouse antibody conjugated with 20 nm gold particles labelling, was carried out. Phalloidine (20 µM) was used to stabilize actin filaments before SDS treatment. 1 µM phalloidine was present in all washing solutions along labeling procedures. Fine structure examination was done under a Philips CM 100 electron microscope. This study was repeated more than 5 times and the same images were consistently observed.

**Fluorescent analysis on mitochondrial morphology in tobacco culture cell**

Wild-type control tobacco cells were stained with 250 nM MitoTracker Red CM-H2XRos (Molecular Probes) for 30 minutes at room temperature followed by 2X washing. Images were collected by a Zeiss LSM510 meta confocal microscope (Carl Zeiss MicroImaging, Inc.) with a C-Aprochromat 63x/1.2 W objective (Carl Zeiss MicroImaging, Inc.). The fluorescence of MitoTracker was excited using a 543 nm line of He-Ne laser, and fluorescence was collected by a 565-615 nm band pass filter.

**RESULTS**

Freeze-fracture electron microscopic analysis permits a direct visualization of mitochondrial structures on freshly-exposed surfaces beneath the outer membrane of randomly-fractured (preferentially cleaved along the hydrophobic inner plane of membranous structure). The arrow in Figure 1A/B and 1C points to an ellipsoidal area in which the upper face of the outer membrane appears to have been dislodged, exposing a discrete parallel-aligned filamentous structure and intramembranous particles (IMPs, marker of mitochondrial inner membrane, see Figure 1B). These particles are proteins/lipoproteins embedded in the intra-lipid bilayer of mitochondria (Lang, 1987). The fact that IMPs cast shadows on the surface of the 5 nm filament structure (see Figure 1B) suggests that this filament structure lies beneath the IMPs and presumably attaches to the protoplasmic surface or possibly to the external surface of the inner membrane. In the magnified image shown in Figure 1B, the width of the filaments measured at stretches with the least metal casting material is approximately 5 nm. This particular filament structure exhibits a distinct organization with an apparent periodicity, uniformly-sized constituting units and a possible inter-filament cross linking pattern (Figure 1B). These structural features bear a striking resemblance to an actin filament bundle. The 20 nm immunogold grains seen in Figures 1A and 1B originated from a low intensity background labeling of the COXIII protein as an internal control for the freeze-fracture procedure. A similar filamentous structure is shown in Figure 1C, but with the 5 nm filaments arranged into a meshwork. The same meshwork pattern was also present in the mitochondrial matrix (Figure 1D, arrow) but with a looser organization than in the membrane shown in Figure 1C.
Immunogold labeling of anti-actin antibody in freeze-fractured mitochondria (Figure 2) indicates that a high density of gold particles accumulate as a patch and arranging in lines on protoplasmic surfaces of the mitochondrial membrane (Figure 2A). Since IMPs heavily cover this dislodged and exposed area, actin (or actin-like) filaments are likely attached to the mitochondrial inner membrane (Figure 2A). Figures 2B and 2C show that gold-labeled antibody reacted with actin or actin-like proteins located under the mitochondrial outer membrane and in the matrix, respectively. In both cases, the linear arrangement of gold grains indicates the presence of actin or actin-like filaments with mitochondria. Figure 2D is the control of Figure 2A-C. Almost no cross reaction between the control antibody (PM 028) and the freeze-fractured mitochondria could be detected (Figure 2D and unpublished results).

In situ immuno-gold labelling on thin sections of mung bean mitochondria revealed the localization of actin in the

![Image](https://example.com/image1.jpg)

**Figure 1.** A 5 nm filamentous structure is visualized underneath the surface of mitochondria. Purified mitochondria were subjected to freeze-fracturing, and then immunogold labeling using anti-COX III antibody (for A and B) after laying the metal replica (see Materials and Methods). The arrow points to a fractured area in which the mitochondrial outer membrane appears dislodged, exposing the intramembranous particles (IMPs). The 5 nm-wide filaments lie adjacent to and just below these IMPs. Note that the width of the filaments and their constituting subunits display a good degree of uniformity and periodicity. The area indicated by the arrow in Panel A is amplified in Panel B. Panels C and D show a 5 nm filamentous structure with cross meshwork arrangements. Panel C shows filaments under the surface of mitochondrion as in Panel A. Panel D indicates cross meshwork and filament structure in mitochondrial matrix. All filaments are composed of globular protein. Bar equals 200 nm.
mitochondrial matrix (Figure 3, Panels A and B). Heavy gold particles representing actin frequently co-localized with the denser area of the mitochondrial matrix under EM examination. Most interesting finding was that nearly all the anti-actin immunogold grains observed inside the mitochondria clustered together and co-localized, rather than distributing evenly, in the denser area in the mitochondrial matrix. Some clusters localized in the center of the mitochondrial domain (Figure 3A) and some localized near the border (Figure 3B). In the same section, only a few immunogold grains scattered in the cytoplasm and virtually none appeared in the control sections (without the anti-actin antibody) (Figure 3C, see Materials and Methods).

After staining with 1% uranyl acetate, whole-mount electron microscopic analysis of mitoplasts (with the outer membrane removed by digtonin) revealed a filamentous bundle. Each filament measured about 5 nm wide and was composed of globular proteins, which are associated with mitoplasts (Figure 4, Panel A, arrow; Panel B is the enlargement of the area in Panel A indicated by an arrow). This filamentous bundle was obviously associated with the mitochondrial inner membrane since the mitoplast was repurified after its outer membrane was removed by the appropriate digitonin treatment.

**Figure 2.** Actin-like protein filaments detected in various locations of mitochondria by freeze-fracture immunogold labeling. A high density of gold grains (linearly-arranged) were accumulated as a patch underneath the protoplasmic surface of mitochondrial inner membrane (Panel A). The external face of mitochondrial outer membrane and matrix also show existing of actin-like filaments and hence the gold grains arrangement is mostly linear (Panels B and C, respectively). No gold particles were detected in the control shown in Panel D. Bar equals 200 nm.
During our ultra-structural study of freeze-fractured mitochondria, we often found that some filamentous bundles formed a belt around the neck between a “mother” mitochondrion and its “bud” (Figure 5, Panels A and B; B is the enlargement of arrow pointed area in A). A filamentous bundle composed of several 5 nm filaments circles around between mitochondrion and its “bud”. Each filament was about 5 nm in width (see arrow pointed enlarged area in Panel B). That the “mother” mitochondrion was heavily covered with IMPs indicates a typical inner membrane composition on the very active mitochondrion. No differentiated structure is shown on the smooth lipid membrane of the “bud”. Panel C presents a negative control of Panel A, the image showing a cleaved surface on the mitochondrial matrix that exhibits a “pinching out” of the inner membrane (small arrowhead) from an outer membrane (large arrowhead showing surrounding double membrane) and forming a bud-like structure. There is no 5 nm protein structure located at the border region between the “pinching out” inner membrane and the double membrane. This clearly shows that the bud in Panel A does not arise from the “pinched out” inner membrane but arises from its primordial mitochondrion. Panel D represents a possible “nascent” mitochondrion (small arrowhead) budding out from mother mitochondrion (large arrowhead) and still connected to its mother mitochondrion via a tubular structure (long arrow). It shows clearly that the “mother mitochondrion” encapsulated with a double membrane and the “nascent mitochondrion” shows few IMPs on its membrane.

Treatment of cultured tobacco cells with the F-actin depolymerization reagent, LatB, enhanced the formation of spherical type of mitochondria from 30% up to 48%, and reduced the population of tubular-shaped mitochondria from 70% down to 51% of the mitochondria investigated (Figure 6).

DISCUSSION

The dependency of mitochondria on actin for their intracellular movement and localization is well understood. In Saccharomyces cerevisiae, actin also appears to mediate mitochondrial inheritance (Yaffe, 1999). Though not widely known, actin (or actin-like) protein has been reported inside rat liver mitochondria (Etoh et al., 1990), pea chloroplasts (McCurdy and Williamson 1987) and mung bean mitochondria (Lo et al., 2002), respectively. Wang and Bogenhagen (2006) also reported that actin was found as a mitochondrial nucleoproteins as evidenced by proteomics analysis. In some previous studies, proteins attached to the surface of the organelles were not eliminated systematically prior to analysis. Close association of actin with the mitochondrial surface is to be expected. In order to demonstrate unambiguously that actin (or actin-like) protein exists inside mitochondria, a proper pretreatment of the sample seems necessary. Our study has shown that mung bean mitochondrial actin is protected by the outer mitochondrial membrane as well as by the inner membrane and is thus resistant to protease, high salt or a combination of both treatments. We also demonstrated that fluorescent actin without a mitochondrial presequence may import into plant mitochondria (Lo et al., 2011). This study aims to further understand the localization, organization and possible function of actin or actin-like filaments in higher plant mitochondria via a cytological approach.

Direct visualization of the 5 nm-wide actin-like filamentous bundles or meshwork beneath the outer mitochondrial membrane via freeze fracture microscopy and
whole mount negative staining was shown in Figures 1 and 4, respectively. Freeze-fracture analysis finds that the actin-like filament structure lies beneath the IMPs and presumably attaches to the inner membrane’s protoplasmic surface or possibly to its external surface (Figures 1B and 1C) and that it may also exist in the matrix (Figure 1D). The same meshwork pattern was shown in the mitochondrial matrix with a much looser organization than in the membrane of Figure 1C. We suspect that the matrix’s more loosely-oriented structure may be caused by the hydrophilic environment in the mitochondrial matrix. Based on this hypothesis, we believe that these 5 nm filaments are stable, well organized macromolecules which may move and assemble differently under different environmental conditions. This in vitro evidence is further substantiated by the immunolocalization of actin in the mitochondria (Figure 2A-C). In addition, the filamentous bundles revealed by whole-mount negative staining in the mitoplast (Figure 4) further supports the suggestion that the 5 nm filaments stabilize under hydrophilic conditions during mitoplast purification. Since the same meshwork was frequently found on inner mitochondrial membranes (Figures 1 and 4) during our freeze-fracture and whole-mount negative staining EM analysis, we suggest that these 5 nm filaments, composed of globular proteins, may co-localize with the mitochondrial membrane and play some role in its shape.

As a control for the in vitro experiments described above, we used intact tissue for the immunolocalization experiment in order to ascertain whether actin (or actin-like) protein could be be detected in mitochondria in situ. Using the same anti-actin antibody as for immunolocalization (shown in Figure 2), immunogold grains were localized inside mitochondria in thin tissue sections (Figure 3). Because of the random sectioning angle with respect to the alignment of actin filament bundles, only certain sections were expected to contain significant concentrations of gold grains. On those few sections containing a significant number of grains, they should have been clustered together reflecting the in situ geometry of the filament bundle relative to the section angle. This prediction was borne out in the actual result. Most or nearly all anti-actin immunogold grains observed inside mitochondria did not distribute evenly but were clustered together and co-localized with the denser area in the mitochondrial matrix (Figures 3 A and B). Only a few immunogold grains were scattered in the cytoplasm in the same section and virtually none in control sections, done in the absence of anti-actin antibody (Figure 3C, see Materials and Methods). This result reveals the presence of actin (or actin-like) protein inside mitochondria in situ in a non-random manner suggestive of a confined distribution of mitochondrial actin or actin-like filament bundles in the organelle.

The findings shown in Figure 5 indicates a possibility that an unknown budding process of mitochondrial propagation in mung bean seedlings. The presence of 5 nm filaments cable circle around the border area between “bud” and “mother” mitochondria (Figure 5A) may indicate that the “actin” cable is forming a mitochondrion-dividing ring and plays a role similar to that of FtsZ division ring (Beech et al., 2000; Momoyama et al., 2003; Osteryoung and Vierling, 1995; Miyagishima et al., 2001) or bacterial division ring (Lutkenhaus and Addinall, 2003). The “bud” gradually pinches out and forms a nascent mitochondrion.
Figure 5. A filamentous bundle composed of several 5 nm filaments circling around and between mitochondrion and its “bud” was found in our freeze-fracture analysis under EM (Panels A and B). Each filament is about 5 nm in width (see arrow pointing to enlarged area in Panel B). “Mother” mitochondrion heavily covered with IMPs indicates a typical mitochondrial inner membrane composition. No differentiated structure is shown on the “bud’s” smooth lipid membrane. Panel C presents a negative control image of Panel A. Mitochondrion with a different cleaved surface on mitochondrial matrix. Mother mitochondrion surrounded by a double membrane (large arrowhead) and a connected “pinching out” inner membrane entity (small arrowhead) surrounded by a single membrane extending from the inner membrane of mother mitochondrion. No 5 nm protein structure can be detected at the “pinching out” region. Panel D represents a possible “nascent” mitochondrion (small arrowhead) budding out from its mother mitochondrion (large arrowhead) and still connected to each other via a tubular structure (long arrow). It clearly shows that “mother mitochondrion’s” matrix is fractured and surrounded by a double membrane, the nascent mitochondrion shows few IMPs on its membrane. Bars equal 100 nm (Panels A and B), 200 nm (Panels C and D).
from the coding region of these cDNAs is 96.8% identical to those of ACT11 and ACT12 of Arabidopsis (Meagher et al., 1999) and contains no mitochondrial targeting sequence (unpublished data). This could mean that either the cDNA encoding an unique actin with a targeting sequence has not yet been identified or, that an actin (or actin-like) protein without a targeting sequence could be imported into mitochondria by an unknown mechanism. Nevertheless, our unpublished results demonstrated that a transgenic fluorescent actin without a targeting sequence can be imported into plant mitochondria (Lo et al., 2011).

The discovery of actin filaments in mitochondria raises additional questions: How are the actin molecules polymerized in the mitochondria? What is the role of polymerized actins or actin filaments inside the mitochondria? Based on the observation of the appearance of actin or actin-like filaments on mitochondrial membrane (Figures 1-5) and changes in mitochondrial morphology with the addition of an actin depolymerization reagent (Figure 6), we speculate that actin filaments play a role in maintaining mitochondrial shape or in its propagation. Considering the data shown above, we wonder whether our observa-

![Figure 6](image_url)
tion is in any way related to the mitochondria propagation model we proposed earlier, that is based on a “nascent mitochondria” population we found in mung bean seedlings (Dai et al., 1998). Taken together, we suggest that some 5 nm filaments surround the mitochondrion and its bud may play a role in mitochondrial shaping and propagation.

**Acknowledgments.** This research was supported by research grants from the National Science Council and from Academia Sinica, ROC.

**LITERATURE CITED**


以微細結構分析來研究似肌動蛋白細絲和綠豆粒線體的相關性
以及它潛在的功能性

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我們利用對 mitoplast 的 negative stainning 方法，在綠豆線粒體中發現 5 毫微米似肌動蛋白細絲的存在。由冷凍裂解電子顯微鏡觀察，球狀蛋白質組成的 5 毫微米似肌動蛋白細絲和粒線體的 intramembranous particles 共存。冷凍裂解的免疫標記法結果，建議似肌動蛋白細絲附在粒線體內膜 protoplasmic 內外表面。他們也可能位於外膜下層和基質中。由原位 immunolocalization 方法，也證實肌動蛋白或似肌動蛋白出現在線粒體基質。有趣的是，一些 5 毫微米似肌動蛋白細絲經常圍繞在線粒體和它的「芽體」之間。這也許代表一種新發現的由 5 毫微米似肌動蛋白細絲介入的一種新穎的植物粒線體發芽繁殖的過程。線狀肌動蛋白解聚作用試劑，Latrunculin B 處理煙草細胞，可影響粒線體形狀。綜合而論，這項研究建議一種 5 毫微米似肌動蛋白細絲，在高等植物粒線體中扮演著和粒線體繁殖和粒線體形態相關的角色。

關鍵詞：肌動蛋白細絲；肌動蛋白動力學和線粒體形態；線粒體繁殖；Vigna radiata。