The Role of Rab27a in the Regulation of Melanosome Distribution within Retinal Pigment Epithelial Cells

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Melanosomes within the retinal pigment epithelium (RPE) of mammals have long been thought to exhibit no movement in response to light, unlike fish and amphibian RPE. Here we show that the distribution of melanosomes within the mouse RPE undergoes modest but significant changes with the light cycle. Two hours after light onset, there is a threefold increase in the number of melanosomes in the apical processes that surround adjacent photoreceptors. In skin melanocytes, melanosomes are motile and evenly distributed throughout the cell periphery. This distribution is due to the interaction with the cortical actin cytoskeleton mediated by a tripartite complex of Rab27a, melanophilin, and myosin Va. In ashen (Rab27a null) mice RPE, melanosomes are unable to move beyond the adherens junction axis and do not enter apical processes, suggesting that Rab27a regulates melanosome distribution in the RPE. Unlike skin melanocytes, the effects of Rab27a are mediated through myosin VIIa in the RPE, as evidenced by the similar melanosome distribution phenotype observed in shaker-1 mice, defective in myosin VIIa. Rab27a and myosin VIIa are likely to be required for association with and movement through the apical actin cytoskeleton, which is a prerequisite for entry into the apical processes.

INTRODUCTION

The retinal pigment epithelium (RPE) is a monolayer that lies between the photoreceptors and the choroid. RPE cells control the neural retinal environment, phagocytose the tips of photoreceptor receptor outer segments, and are required for the regeneration of visual pigment and the maintenance of the blood-retinal barrier. As the name suggests, RPE cells exhibit melanin pigment contained within the cytoplasm in membrane-enveloped granules called melanosomes. The function(s) of pigment within the RPE are not entirely clear but are likely to include absorption of stray light, thus minimizing light scatter, and the absorption of free radicals and toxins. The basal surface of the RPE contacts a basement membrane called Bruch’s membrane and the choriocapillaris, a layer of fenestrated capillaries. The RPE apical surface forms numerous long processes, which reach up between the outer segments of the photoreceptors and partially envelop them. In fish and amphibians, the melanosomes of the RPE exhibit a dramatic redistribution from the cell body into the apical processes upon the onset of light, which is reversed in the dark. Melanosomes in the RPE of mammals are generally thought not to move with the light cycle (Arey, 1915; Burriside and Latties, 1979). Melanosomes in the RPE of mammals must, however, have the ability to move as some are found within the apical processes.

In melanocytes of the skin, melanosomes move bidirectionally along microtubules and are trapped in the cell periphery by interaction with the cortical actin cytoskeleton (Wu et al., 1998). This peripheral capture is essential for transfer of the granules to neighboring keratinocytes. Interaction of melanosomes with the actin cytoskeleton is mediated through a complex of Rab27a, melanophilin, and myosin Va, where Rab27a binds to the melanosome and myosin Va to actin and melanophilin is a linker between Rab27a and myosin Va (Hume et al., 2001, 2002; Wu et al., 2001, 2002; Fukuda et al., 2002; Provance et al., 2002; Strom et al., 2002). Studies of the regulation of melanosome movement in melanocytes have been greatly aided by studies of melanocytes of the ashen, leaden, and dilute mice which lack functional Rab27a (Wilson et al., 2000), melanophilin (Matesic et al., 2001), and myosin Va (Mercer et al., 1991), respectively, and are models for Griscelli syndrome in humans (Seabra et al., 2002).

The melanosome dynamics within the RPE is much less well characterized. A study of the RPE of ashen, leaden, and dilute mice has not been reported. The only known phenotype has been observed in the RPE of the shaker-1 mouse, which is defective in myosin VIIa (Gibson et al., 1995). In shaker-1 mice, the melanosomes are found exclusively in the cell body of the RPE and do not enter the apical processes (Liu et al., 1998). Shaker-1 is the mouse model for the human disease, Usher syndrome type 1B, whose patients suffer...
from progressive retinal degeneration and hearing defects (Weil et al., 1995; Pettit, 2001). The shaker-1 mice have hearing defects but do not suffer from retinal degeneration. Nevertheless, defects in the efficiency of degradation of phagocytosed rod outer segments (Gibbs et al., 2003) and defects in opsin transport from the inner to outer segment of photoreceptors (Liu et al., 1999) have been observed in shaker-1 retinas.

That Rab27a might have a function in the RPE was suggested by its identification as a possible trigger of retinal degeneration in X-linked choroideremia, a disease characterized by slow degeneration of RPE, choroid and photoreceptors resulting in blindness at middle age (Seabra et al., 1995; MacDonald et al., 1998). Patients suffering from this disease lack functional REP-1, one of two REP isoforms responsible for prenylation and activation of Rab GTPases (Seabra, 1996; van den Hurk et al., 1997). Although many Rab s are normally prenylated in this disease, Rab27a is not and is thus unable to associate with intracellular membranes and function properly (Seabra et al., 1995; Larijani et al., 2003). Rab27a appears to function through the recruitment of target proteins called effectors when it is activated upon GDP loading. Melanophilin is one such effector and forms part of a family of proteins that contain a Rab27a-binding domain (Kuroda et al., 2002; Strom et al., 2002). Mryptip is a member of the melanophilin gene family, which has been recently identified as a binding partner of both Rab27a and myosin VIIa (El-Amraoui et al., 2002). Myrtp is expressed in the RPE, suggesting that melanosome movement within the RPE is regulated by interaction with the actin cytoskeleton via Rab27a-Mryptip-myosin VIIa (El-Amraoui et al., 2002).

In this article, we tested the hypothesis that Rab27a and myosin VIIA regulate pigment granule movement in RPE by examining the pigment granule distribution within the RPE of ashen and shaker-1 mouse mutants. We have further investigated the motility of melanosomes within the RPE and investigated how Rab27a and myosin VIIA might regulate the distribution of the melanosomes within the RPE via interaction with the actin cytoskeleton.

MATERIALS AND METHODS

Materials

Biotinylated phallolidin was from Molecular Probes (Eugene, OR), rabbit antibiotin antibody from Toram Biologicals Ltd. (Peterborough, UK), protein A gold from University Medical Center (Utrecht) and rabbit anti-mouse intermediate antibody from Dako Ltd. (Ely, UK). Other primary antibodies were mouse anti-α tubulin (Kreis, 1987), anti-Rab27a mAb 4B12 and polyclonal antibody Q142 (Hume et al., 2002); antimyosin VIIA polyclonal antibody was a gift from Steve Brown (MRC Mammalian Genetics Unit, Harwell, UK; Todorcev et al., 2001).

Mice

All mice were bred and maintained on 12 h-light/12 h-dark cycle under UK project license PPL 70/5071 at the Central Biomedical Services of Imperial College London or at University Eye Hospital (Tubingen). Ashen mice (C57BL/6j; ash/ash) colony was produced in the Seabra laboratory as described previously (Ramalho et al., 2002). Shaker-1 mice on a modified Balb-c background containing white-bellied agouti (Aw) mice were obtained from MRC Mammalian Genetics Unit (Harwell, UK). To maintain a colony, heterozygous females (+/sh-1, C/c, A/a) were bred with homozygous males (sh-1/sh-1, c/c, A/a). Homozygous shaker-1 mice were albino given the linkage between both sh and c loci on chromosome 7. However, a spontaneous crossover event led to the generation of homozygous agouti shaker-1 mice that were used for this study. C57BL/6 wild-type mice were purchased from B&K Universal Limited (Hull, United Kingdom).

Tissue Histology

All the mice used for histology studies were perfused with phosphate-buffered saline (pH 7.4) before dissection. Sections were fixed in paraformaldehyde-glutaraldehyde fixative (4% paraformaldehyde, 5% glutaraldehyde, 0.1 M cacodylate buffer) overnight, except for the eyes, which were fixed only for 1 h at 4 °C. Eyes were cut in half and the anterior part was removed. Fixed organs were washed three times for 10 min at RT with phosphate-buffered saline. For histologic analysis, specimens were embedded in paraffin, sectioned to 3–5-μm thickness, and stained with hematoxylin and eosin.

Conventional Electron Microscopy

Mouse eyes were fixed in 1% paraformaldehyde, 3% glutaraldehyde in 0.07 M cacodylate buffer. The cornea was cut off and the lens was removed. The neural retina was then peeled from the eye cup leaving the RPE exposed at the apical surface and still attached to Bruch’s membrane on the choroid. The specimens were postfixed in 1% osmium 1.5% potassium ferricyanide in 0.1 M cacodylate and then dehydrated in alcohol. The specimens were then cut into pieces ~2-mm square and critical point dried, mounted on stubs with conductive silver paint, and sputter-coated with gold. Specimens were examined in a JEOL 1010 transmission electron microscope (Welwyn Garden City, United Kingdom).

Scanning Electron Microscopy

Mouse eyes were fixed in 4% paraformaldehyde in 0.1 M phosphate buffer. The cornea was cut off and the lens was removed. The eye cup was then cut into ~3-mm squares and embedded in 10% gelatin. After allowing the gelatin to solidify at 4 °C pieces of eye cup sandwiched between gelatin layers were excised and immersed for >2 h at 4 °C in 2.3 M sucrose. Eye cup sandwiches were then mounted on pins and frozen in liquid nitrogen. Sections (500 nm) were cut at −80°C using a Leica FCS cryo-ultramicrotome (Milton Keynes, United Kingdom). Sections were picked up in sucrose onto slides, quenched with 0.02 M glycine, blocked with 1% BSA, and then incubated with primary antibodies, followed by fluorescent secondary antibodies and/or fluorescent paint. Specimens were examined using a Bio-Rad Radiance 2000 confocal microscope (Richmond, CA).

Immunofluorescence of Tissue Sections

Mouse eyes were fixed in 4% paraformaldehyde in 0.1 M phosphate buffer. The cornea was cut off and the lens was removed. The eye cup was then cut into ~3-mm squares and embedded in 10% gelatin. After allowing the gelatin to solidify at 4 °C pieces of eye cup sandwiched between gelatin layers were excised and immersed for >2 h at 4 °C in 2.3 M sucrose. Eye cup sandwiches were then mounted on pins and frozen in liquid nitrogen. Sections (500 nm) were cut at −80°C using a Leica FCS cryo-ultramicrotome (Milton Keynes, United Kingdom). Sections were picked up in sucrose onto slides, quenched with 0.02 M glycine, blocked with 1% BSA, and then incubated with primary antibodies, followed by fluorescent secondary antibodies and/or fluorescent paint. Specimens were examined using a Bio-Rad Radiance 2000 confocal microscope (Richmond, CA).

Cryo-immunoelectron Microscopy

Mouse eyes were embedded as described for immunofluorescence above. Sections (100 nm) were cut at −120°C and picked up in 1:1 sucrose-methyl cellulose. Sections were then labeled using 10 nm protein A gold as described (Slot et al., 1991). For F-actin staining sections were labeled with biotinylated phallolidin, followed by rabbit antibiotin and protein A gold. For labeling with rabbit polyclonal antibodies primary antibody was followed by protein A gold or labeling with mouse monoclonal antibodies primary antibody was followed by rabbit anti-mouse intermediate antibody and then protein A gold.

Electroretinography

ERGs were obtained in 12-month-old mice according to previously reported procedures. Briefly, the mice were dark-adapted overnight and anesthetized with ketamine (66.7 mg/kg) and xylazine (11.7 mg/kg). The pupils were dilated with tropicamide eye drops (Mydriatikum Stulln, Pharma Stulln, Germany). The ERG setup featured a Ganzfeld bowl, a DC amplifier, and a PC-based control and recording unit (Multimicro Vision, Jager/Toennies, Hoechberg, Germany). Band-pass filter cut-off frequencies were 0.1 and 3000 Hz. Single flash recordings were obtained both under dark-adapted (scotopic) and light-adapted (photopic) conditions. Light adaptation before the photopic session was performed with a background illumination of 30 cd/m² for 10 min. Single flash stimuli were presented with increasing intensities, ranging from 10−4 candela second per m² (cds/m²) to 25 cds/m², divided into 10 steps of 0.5 and 1Log cds/m². Ten responses were averaged with an interstimulus interval (ISI) of either 5 or 17 s (for 1, 3, 10, 25 cds/m²).

RESULTS

The Role of Rab27a in the Regulation of the Distribution of Melanosomes within the RPE

To determine whether Rab27a regulates melanosome distribution in the retina, we examined the distribution of melanosomes within the RPE of homozygous ashen (Rab27a null) mice, heterozygous, and wild-type mice 2 h after light onset.
Histological sections observed under a light microscope revealed that the gross organization of the retina is normal. All retinal layers are present with normal thickness but the melanosome distribution within the ashen RPE is different from the wild type (Figure 1). Melanosomes in wild-type RPE are found in an area that extends clearly apical to the nucleus, whereas the distribution of melanosomes within ashen RPE is much more restricted in that

Daily Movement of Melanosomes within the RPE

Ever since the first microscopic studies of retinas over a century ago, it has been suggested that melanosome movement in mammals is minimal and very little is known about melanosome dynamics within mammalian RPE cells. The above results suggested that RPE melanosomes indeed move and that the movement may be regulated by at least two gene products, Rab27a and Myosin VIIa. To study RPE melanosome dynamics, we determined whether melanosomes of wild-type mice sacrificed at different times. Immediately before light onset, we observed only a small number of melanosomes in the apical processes and those are, in general, at or near the base below the outer segments (Figure 3A). Just >1 h after light onset, we observed a large number of phagosomes present in the apical cytoplasm. These phagosomes are predictably the product of light-induced phagocytosis of photoreceptor outer segments (Figure 3B).

At this time point, the number of melanosomes in the apical processes remains low but the number increases significantly 1 h later. Although many of these melanosomes are still in the base of the apical processes at the 2 h after light onset time point, a significant number of melanosomes extend into the interdigitating processes (Figure 3C). At later time points (3.5 and 5.5 h after light onset), there is a gradual reduction in the number of melanosomes in the apical processes (Figure 3, D and E). Quantitation of the number of melanosomes in the apical processes at different time points
indicates that there is a maximum of a threefold increase in
the number of melanosomes in the apical processes 2 h after
light onset, compared with the dark-adapted eye (Table 1).

**Ultrastructure of the Apical Processes of Mouse RPE**

Transmission EM of the wild-type mouse RPE suggested
there is very little space between adjacent outer segments
within which the melanosomes must move. This prompted
us to examine the melanosomes within the apical processes
at high magnification in order to further understand the
relationship between the melanosome, the cytoskeleton, and
the plasma membrane of the apical process. High magnifi-
cation shows multiple very thin apical processes extending
between the outer segments in the wild-type, *ashen*, and
*shaker-1* RPE (Figure 4, A–C). These images suggested that
the apical processes of the mutant *ashen* and *shaker-1* RPE
might be more disorganized than those of the wild-type
RPE, but this is difficult to resolve by thin-section EM, where
an entire apical process may not be within the section plane.
It is also not possible to assess the three-dimensional organ-
ization of the apical processes by thin-section EM. We
therefore performed scanning EM of the mouse RPE after
removal of the photoreceptor layer. The apical processes
of RPE cells are not finger-like projections but rather overlap-
ning leaf-like projections (Figure 4, E–G). Those projections
in the *ashen* and *shaker-1* RPE show no clear differences from
the wild type. Although the melanosomes that enter the
wild-type apical processes are elongated, their shortest di-
ameter is still greater than the width of the apical processes
when they do not contain melanosomes. When a melano-
some enters an apical process, the membrane of the apical
process appears to become distended around the granule

**Figure 2.** Electron microscopy shows that melanosomes do not move into the apical processes of RPE of *ashen* and *shaker-1* mouse. Transmission EM longitudinal sections of mouse retinas (A–C) show elongated melanosomes within the apical processes of wild-type RPE that extend between the rod outer segments (ROS), whereas in *ashen* and *shaker-1* mice melanosomes do not enter the apical processes. BM, Bruch’s membrane. Oblique sections (D and E) show that many melanosomes are above the level of the adherens junctions (arrows) and extend into the apical processes between the ROS in the heterozygous (+/−, phenotypically normal) mouse (D), whereas in the homozygous (−/−) *ashen* mouse (E) the melanosomes remain below the level of the junctions. Bar, 2 μm.
It is not possible even at the highest magnification to visualize cytoskeletal structures within the apical processes whether or not they contain melanosomes.

**Interaction of Melanosomes of RPE with Cytoskeleton**

Fluorescent phalloidin staining of wild-type RPE shows F-actin to be localized to the basal infoldings and to the apical processes (Figure 5, A and B). F-actin is also found in the circumferential actin ring in the apical region of the cell, which is associated with the adherens junctions. In these 0.5-μm sections that do not contain the entire cell, the circumferential actin ring appears as a short stretch of filament bundle only (Figure 5A). The apical processes are not rigid because after paraformaldehyde fixation the photoreceptors become partially or totally separated from the RPE. The RPE–photoreceptor interface is thus not as well preserved as in the conventional EM shown above, where glutaraldehyde fixation was used. Microtubules are confined to the cell body

(Figure 4G). It is not possible even at the highest magnification to visualize cytoskeletal structures within the apical processes whether or not they contain melanosomes.

**Table 1. Quantitation of the number of melanosomes within the apical processes at different times in the light cycle**

<table>
<thead>
<tr>
<th>Time after light onset (h)</th>
<th>% melanosomes in apical processes</th>
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<tr>
<td>0</td>
<td>5.0</td>
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<tr>
<td>1.25</td>
<td>5.7</td>
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<tr>
<td>2</td>
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<td>3.5</td>
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C57BL/6J mice were sacrificed at the indicated times after light onset and longitudinal sections of RPE examined by transmission EM. The point at which the apical process arises (which may be below the level of the outer segments) can be readily identified. Every melanosome within each section was scored for presence in the apical processes. More than 600 melanosomes were counted per time point.
and absent from the apical processes. Most melanosomes are found in the microtubule-rich cell body but some have penetrated the actin-rich cortex and into the actin-rich apical processes. Both melanosomes and actin are concentrated in the basal region of the apical processes and appear closely associated. In the ashen RPE, the organization of actin filaments and microtubules appears the same as the wild-type but the melanosomes are all within the microtubule-rich cell body and show no association with F-actin (Figure 5C). Rab27a is associated with the melanosomes within the wild-type RPE, particularly with the elongated melanosomes in the apical region of the cell but also with some spherical melanosomes in the cell body (Figure 5D). Myosin VIIa stains at least some of the melanosomes in the RPE by immunofluorescence and also shows some cytoplasmic staining (Figure 5E). Some bright staining is also observed above the cell body of the RPE, which may be on outer segments of photoreceptors that have remained associated with the RPE (Figure 5E). Myosin Va strongly stains structures that are clearly apical to the region containing the majority of the F-actin and melanosomes (Figure 5F). These structures are photoreceptor outer segments (see below), which sometimes remain associated with RPE after paraformaldehyde fixation.

Cryo-immuno-EM allowed the identification of the adherens junctions and revealed F-actin labeling in the infoldings of the basal membrane (unpublished observations) and in the circumferential actin ring, just beneath the apical plasma membrane, and within the apical processes (Figure 6). Melanosomes that were at the level of or above the adherens junctions had F-actin closely associated with them, whereas those below the adherens junctions were completely devoid of actin staining. Rab27a was found on all types of melanosomes within the RPE, whether present in the cell body or in the actin-rich apical regions and whether spherical or elliptical in shape (Figure 7, A and B). Although the majority of the Rab27a staining was around the perimeter membrane of the melanosome, some Rab27a staining was observed within the lumen of the granule. Myosin VIIa staining was found on some melanosomes within the RPE. The staining was observed on both spherical and elliptical melanosomes in the cell body, both above and below the adherens junctions and within the apical processes. Nearly all the staining was present around the perimeter membrane of the melanosome (Figure 7, C–E). Myosin Va localized to the plasma membrane of photoreceptor outer segments and both the apical processes, and melanosomes were largely devoid of myosin Va staining (Figure 7, F and G).

Electroretinography Analysis of Ashen Mice
The potential impact of the lack of Rab27a on retinal function was assessed with electroretinography (ERG). Under scotopic conditions with low stimulus intensities (up to \(10^{-2}\) cdes/m²), the rod system response can be determined, whereas responses of the cone system can be identified under photopic conditions. The scotopic ERG with stimulus intensities of \(10^{-1}\) cdes/m² and more represents a mixed response of rods and cones. Figure 8 summarizes the results obtained in ashen (Rab27a null) mice in comparison to wild-type controls.

A series of exemplary scotopic and photopic ERG traces obtained from an ashen and a wild-type control mouse is shown in Figure 8A. No significant differences in the shape of the waveforms or the size of the responses could be identified. A comparison between both groups regarding the photopic and scotopic b-wave amplitude is shown in Figure 8B. The upper and lower lines delimit the normal range.
and 5% quantiles of the control group), whereas the ashen data are shown as box plots (the gray boxes indicate 25 and 75% quantiles, the whiskers the 5 and 95% quantiles). The results demonstrate that the b-wave amplitude of the ashen mice was within normal limits for all stimulus intensities under both photopic and scotopic conditions. In summary, no functional differences between ashen and wild-type mice were found.

**DISCUSSION**

The results presented here clearly demonstrate a requirement for functional Rab27a in the light-dependent movement of melanosomes into the apical actin-rich cytoplasm and the apical processes within RPE cells. The phenotypic similarities observed in the RPE of mice lacking Rab27a or myosin VIIa suggest a model where Rab27a links melanosomes to the actin cytoskeleton via myosin VIIa to regulate melanosome distribution in the RPE.

**Melanosomes in the RPE Are Not Static but Move in and out of the Very Narrow Apical Processes**

Our data suggest that pigment granule distribution within the RPE is not random and that melanosomes do move in a manner which may be regulated by light. The early microscopic studies on mammalian retinas performed a century ago suggested little, if any, movement of melanosomes, in marked contrast with amphibians and fish (reviewed in Arey, 1915; Burnside and Laties, 1979). To our knowledge, this is the first detailed description of RPE melanosome movement in mammals. The present study suggests that melanosome movements are modest, both in terms of the number of melanosomes that move at any one time and in terms of the distance that they move. The correlation between melanosome movement and the light cycle is intriguing but it remains to be determined whether light onset is the primary trigger or secondary to other light-induced changes within the photoreceptor–RPE interface. It is possible, for instance, that the light-induced phagocytosis of the tips of the photoreceptor outer segments removes physical constraints, thus allowing melanosome movement up the apical processes. As the day progresses, the outer segments move back down to a position more closely juxtaposed to the RPE and may reimpose physical constraints resulting in the gradual movement of melanosomes down the apical processes. That physical constraints exist on melanosome movement within the apical processes is suggested by the very small diameter of the apical process compared with the smallest diameter of the elongated melanosome. In summary, our observations suggest that the speed and magnitude of melanosome movement in response to light-sensitive triggers (if they exist) is slowed by physical restraints.

**Figure 5.** Immunofluorescence localization of melanosomes, Rab27a, and cytoskeletal proteins. Wild-type (A, B, and D–F) and ashen (C) RPE and choroid were labeled with fluorescent phalloidin (red in A–C, E, and F), antitubulin (green in A–C, red in D), anti-Rab27a (green in D), antimyosin VIIa (green in E), and antimyosin Va (green in F). Melanosomes identified by transmitted light have been artificially colored blue in the merged images. Arrows indicate examples of coincidence between melanosomes and rab27a (D) and melanosomes and myosin VIIa (E). AP, apical processes; CB, cell body; BI, basal infoldings. Bar, 5 μm.
We have not been able to directly measure the kinetics of melanosome movement within mouse RPE, because pigmentation and the elaborate apical processes are not readily reproduced in cultured mammalian RPE cells.

We note that the extent and speed of melanosome movement within the RPE of some mammals may be more striking than in mice. The apical processes of the RPE of humans are more loosely packed between the photoreceptor outer segments, which may allow more movement of melanosomes within them (unpublished observations). We also show here that the apical processes of the mouse RPE are flattened leaflike, rather than fingerlike, projections, as has been reported in monkey RPE (Marmor et al., 1980). Movement of the melanosome within the narrow apical processes is likely to be facilitated by the fact that the apical processes are only narrow in two dimensions.

**How Does Rab27a Regulate Melanosome Movement within the RPE?**

In mammalian RPE cells, actin filaments are reported to be found in the infoldings of the basal plasma membrane, the lateral membrane cytoskeleton, the circumferential microfilament bundles which are associated with the adherens junctions and in the apical processes (Nguyen-Legros, 1978; Burnside and Bost-Usinger, 1998). In this study, we have...
found F-actin in all locations but the lateral membrane cytoskeleton. F-actin was clearly visible in the basal infoldings, the circumferential microfilament bundles, beneath the apical plasma membrane and within the apical processes of mouse RPE cells (Figures 5 and 6). Microtubules were completely absent from the actin-rich apical region, including the apical processes and were found exclusively in a meshwork in the cell body.

Melanosomes in melanocytes of the skin move bidirectionally along microtubules and are trapped by a Rab27a-dependent interaction with the cortical actin cytoskeleton (Hume et al., 2001; Wu et al., 2001). By analogy, melanosomes in the RPE may reach the actin-rich apical region by movement along the microtubules present in the cell body, although there is no obvious alignment of melanosomes with microtubules within the cell body of the RPE.

Our observations suggest that there are no melanosomes above the level of the adherens junctions, the circumferential actin ring and the actin-rich cortex in the two mutant RPE cells presented in this study. Similar results for shaker-1
Melanosomes in Retinal Pigment Epithelium

and stationary melanosomes that can be distinguished by the presence or absence of Rab27a/myosin VIIa, although we cannot rule out this possibility.

In the RPE of wild-type mice just before light onset, some of the melanosomes were found above the level of the adherens junctions in the actin-rich apical cytoplasm and a few had entered the base of the apical process. This suggests that the Rab27a-dependent association of melanosomes with apical F-actin may not be regulated by light, although subsequent movement of melanosomes up the apical processes may be light sensitive. In addition to the initial association of the melanosome with the apical actin cytoskeleton and/or movement through it, Rab27a/myosin VIIa may play a role in the subsequent movement within the apical processes. Melanosome movement along apical processes has been most extensively studied in teleost RPE, where both dispersion to the tips and aggregation within the cell body are dependent on actin filaments because they are disrupted upon cytochalasin treatment (King-Smith et al., 1997). Aggregation but not dispersion may also have a requirement for microtubules (Troutt and Burnside, 1989). Unlike the apical processes of teleost RPE, those of mammals have actin filaments but no microtubules (Burnside and Laties, 1979 and this study). In mammalian RPE the actin filaments are arranged somewhat differently from those of teleost RPE or apical microvilli of other epithelial cell types in that, rather than being arranged in a central bundle they are found in parallel arrays, which may provide greater mechanical support to the leaflike projections of mammalian apical processes (Burnside and Laties, 1976). The tight packing of the apical projections between the outer segments and the dispersion of the apical plasma membrane and the actin filaments around the melanosome suggest that force is required for movement of the melanosome up the apical process, and it is possible that Rab27a and myosin VIIa may participate in this movement along the actin filaments.

The interaction between Rab27a and myosin VIIa is probably not direct (unpublished observations; El-Amraoui et al., 2002). In skin melanocytes, Rab27a interacts indirectly with myosin Va through a bridging protein called melanophilin (Fukuda et al., 2002; Strom et al., 2002; Wu et al., 2002). Melanophilin contains an N-terminal domain that binds activated GTP-bound Rab27a and a medial domain that binds the melanocyte-specific splice isoform of myosin Va tail. Recently, a protein related to melanophilin called Myrip (for myosin and rab interacting protein) was discovered on the basis of binding to the tail of myosin VIIa (El-Amraoui et al., 2002). This protein also binds activated Rab27a and is expressed in the RPE, suggesting that it is the best candidate linker protein between Rab27a and myosin VIIa. It would be interesting to study Myrip knock-out mouse RPE, which is predicted to exhibit the same phenotype as ashen and shaker-1 mice.

Possible Additional Functions of Rab27a and Myosin VIIa in the RPE

We noted in this study that the distribution of melanosomes within the microtubule-rich cell body in ashen and shaker-1 mice was not substantially different from that in wild-type RPE. However, the melanosomes in the mutant cells appeared more randomly oriented both above and below the adherens junctions within the apical processes and in the cell body of the RPE. A possible mechanism whereby Rab27a would regulate granule orientation within the cell body below the adherens junctions of the RPE remains unclear because F-actin is largely absent from that area of the cell.
Myosin VIIa appears to be involved in adhesion between neighboring stereocilia of the inner ear and may play a similar role in the RPE (Self et al., 1998), although we did not observe significant staining of myosin VIIa on the apical processes. It is possible that melanosome movement plays a role in the stabilization of the apical processes by unknown mechanisms. Myosin VIIa has also been implicated in the transport of phagosomes within the RPE from the apical cytoplasm to the basal cytoplasm for fusion with lysosomes (Gibbs et al., 2003). We have not measured the effects of Rab27a deletion on this process, and this issue will be a focus for future studies. Furthermore, a role for melanosomes in the degradation of phagocytosed outer segments has been proposed (Thumann et al., 1999). Therefore, it is possible that Rab27a and myosin VIIa regulate the motility and fusion of melanosomes and phagosomes in the RPE.

We show here that myosin Va is not found on melanosomes within the RPE but is strongly expressed on the plasma membrane of the photoreceptor outer segments. In addition, myosin Va staining was previously reported to be enriched in inner segments and rod photoreceptor synapses, suggesting that myosin V may play important roles in photoreceptor physiology (Schlamp and Williams, 1996). We observed a normal gross retinal morphology and melanosome distribution in the RPE of dilute mice, defective in myosin Va (unpublished observations). The strain used, dilute-viral exhibits a coat-color defect resulting from melanosome transport defects in skin melanocytes similar to that observed in dilute-lethal mice. However, the retroviral insertion in the Myo5a gene affects primarily the melanocyte-specific, alternative splicing form and some myosin Va activity remains in other tissues. A more conclusive model to study would be the dilute-lethal strain, which is the null allele of mouse myosin Va, but unfortunately the early neonatal death of dilute-lethal mice hinders the examination of the RPE phenotype in these mice. As myosin Va has been clearly shown to be required for Rab27a-dependent movement of melanosomes in melanocytes of the skin, it is noteworthy that this myosin does not appear to be involved in pigment granule movement in the RPE.

Functional Consequences of Deficiency in Rab27a-dependent Movement of PGs within RPE Cells
In addition to the lack of detectable abnormalities in the ERGs of ashen mice, there is no evidence of retinal degeneration or visual defects (unpublished observations and Figure 8). Interestingly, the shaker-1 mutants also do not exhibit significant visual defects or retinal degeneration, whereas human patients suffering from Usher syndrome 1B who also lack functional myosin VIIa do show such changes and undergo retinal degeneration (Libby and Steel, 2001; Petit, 2001; our unpublished observations). However, the human eye disease features a slowly progressive, gradual retinal degeneration, and vision symptoms are usually not apparent before the second decade of life. The comparatively short lifespan of the mouse may prevent the onset of retinal degeneration in the shaker-1 model. Therefore, the failure of melanosomes to move into the apical cytoplasm/apical processes of RPE cells may contribute to gradual retinal degeneration in humans. A failure of movement of melanosomes could contribute to retinal degeneration because of reduced protection of the tips of outer segments from light damage. Alternatively, if Rab27a and myosin VIIa contribute to the phagosome maturation, this could lead to reduced photoreceptor degradation, which in turn would compromise photoreceptor function and lead to the gradual accumulation of undegraded material within the RPE, compromising its function.

This study highlights parallels and differences between melanosomes in RPE and skin melanocytes. Melanosomes of the RPE are longer lived, are frequently elliptical in shape, are not passed on to another cell type, and may be less motile, and the microtubular and actin cytoskeleton of the RPE has an organization very different from that of skin melanocytes. Future mechanistic studies on Rab27a-dependent movement of melanosomes in the different pigmented cell types should continue to contribute to a better understanding of organelle motility, intracellular form, and their pathological consequences in human disease.

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REFERENCES


Melanosomes in Retinal Pigment Epithelium


