Photoreceptor Organization and Rhythmic Phagocytosis in the Nile Rat *Arvicanthis Ansorgei*: A Novel Diurnal Rodent Model for the Study of Cone Pathophysiology

Corina Bobu,1 Cheryl M. Craft,2,5 Mireille Masson-Pevet,1 and David Hicks1

PURPOSE. To characterize rod and cone distribution, organization, and phagocytosis in the diurnal mouse-like rodent *Arvicanthis ansorgei*.

METHODS. Retinas of adult *A. ansorgei* were processed for histology, electron microscopy and immunohistochemistry using rod- and mouse cone-specific antibodies. For phagocytosis studies, retinas were sampled every 3 hours under a 12-hour light–dark cycle and processed for double-label immunohistochemistry. The number of phagosomes in the retinal pigmented epithelium were quantified with a morphometric system.

RESULTS. *A. ansorgei* retinas were composed of 33% cones and 67% rods, approximately 10 times more cones than mice and rats. Cones were arranged in two cell layers at the sceral surface, distributed uniformly across the entire retina. Cone arrestin was distributed throughout the dark-adapted cones, from outer segments to synapses, whereas short- and mid-wavelength cone opsins were restricted to outer segments. Short-wavelength cone density was mapped in wholemounted retinas, in a significantly higher number in the central region. Rhodopsin immunopositive (rod) phagosomes showed a small peak late in the dark phase, then a large burst 1 to 2 hours after light onset, after decreasing to low baseline levels by 12 AM. Peak late in the dark phase, then a large burst 1 to 2 hours after light onset, after decreasing to low baseline levels by 12 AM. Mid-wavelength cone opsins immunopositive (cone) phagosomes were 10 times less numerous than rods, and demonstrated a broad peak 1 to 2 hours after light onset.

CONCLUSIONS. The diurnal rodent *A. ansorgei* possesses a large number of cones, organized in a strict anatomic array. Rod and cone outer segment phagocytosis and shedding can be monitored simultaneously and show similar profiles but different amplitudes. This species may constitute a valuable novel animal model for investigating cone pathophysiology. (Invest Ophthalmol Vis Sci. 2006;47:3109–3118) DOI:10.1167/iovs.05-1397

Phagocytosis of photoreceptor (PR) outer segment (OS) apical tips by the adjacent retinal pigmented epithelium (RPE) occurs throughout life and is an extremely elevated metabolic activity.1–2 In the case of rods, continual synthesis of OS proteins and their insertion into nascent membranes at the basal surface of OS leads to apical displacement of older disc membranes.4 To renew the phototransduction machinery while maintaining a constant OS length, apical packets of discs are shed and ingested by the RPE at a very high rate. It has been estimated that each RPE cell in a person 80 years of age has internalized some 200 million discs.5 In addition, phagocytic uptake of OS debris is not uniform, but exhibits a robust light-driven and/or circadian burst of activity near dawn.6–7 Furthermore, other key features of PR function, including visual pigment synthesis,6–7 OS turnover,9 and ion channel sensitivity10–11 exhibit strongly rhythmic activities. Several lines of experimental evidence, including temperature dependence, use of pharmacological inhibitors, and saturation curves of OS uptake,12–14 indicate that rod OS phagocytosis involves a ligand-receptor interaction, and several different candidate receptors have been identified on the RPE apical surface. Recent work has identified the receptor tyrosine kinase c-Mer (MerTK) as critically involved in rod OS internalization,15 and mutations in this protein are implicated in some forms of human retinal degeneration.16 In addition, αvβ3 integrin has been shown to be a component of the multimolecular complex controlling OS internalization.17–19 However, virtually all of this information has been generated for rod PR, and knowledge of cellular and molecular mechanisms involved in cone OS phagocytosis is very limited. Although it was initially proposed that cone OS did not undergo phagocytosis in the same way as rods,20 subsequent elegant morphologic studies have shown that these PRs did indeed shed apical packets of membrane that were ingested by the RPE.21–25 Cone PRs are responsible for color discrimination and visual acuity and are of utmost importance in human eyesight. The progressive loss of cone PRs represents the major source of visual handicap in a variety of retinal diseases, including retinitis pigmentosa and age-related macular degeneration (AMD).26–28 The factors influencing cone survival are unknown, although we recently identified a diffusible factor produced by rods that may be critical for cones.27 Perturbations in phagocytosis may well play a role in the development of AMD. It has been hypothesized that the progressive loss of vision associated with AMD could involve the impediment of phagocytosis through the accumulation of lipofuscin.28

One simple reason that has greatly hindered research on cone pathophysiology is the lack of a mammalian model permitting scrutiny under controlled experimental conditions. Conventional rodents (i.e., laboratory rats and mice) are nocturnal and have few cones (<3% of total PRs for mice29 and <1% total PRs for rats30), whereas animals possessing cone-rich retinas—ground squirrels (approximately 90% cones31), chickens (approximately 65% cones32), and pigs (15%–20% cones33)—exhibit various difficulties for experimentation, utilization of preexisting antibodies or databank sequences, or

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3109
routine maintenance. Cone-rich transgenic mouse models have recently become available, but the cones in such strains may not be fully normal (although recent data show that they display typical cone features by a wide range of structural and functional criteria). We thus sought a species that would facilitate experimental research into cone function and survival and would allow us to examine cone phagocytosis specifically, through exhibiting certain key features: (1) a large percentage of easily identifiable cones; (2) diurnal behavior patterns similar to the human’s; (3) genetic homology to mice for capitalizing on existing databases and antibody probes; (4) feasibility of monitoring behavior patterns and conducting in vivo experimentation; and (5) ease of maintenance and breeding in captivity. In this report, we present the initial characterization of such a species.

Materials and Methods

Animals

This study was conducted with Nile rats, Arvicanthis ansorgei, raised in our animal facilities from individuals captured in southern Mali in 1998. Since this time, the colony has been maintained on a 12-hour light–dark (LD) cycle in an ambient temperature of 22 ± 1°C. A total of 40 animals were used, all females, with most experiments involving young adults (5–7 months), with some individuals up to 26 months. The animals were supplied ad libitum with water and standard rat chow. Animal treatment and experimentation adhered to the guidelines in the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.

Histology and Retinal Ultrastructure

For examination of the A. ansorgei retina by light and electron microscopy for histologic analysis, eyes were removed from euthanatized animals; fixed in 4% paraformaldehyde, 0.1% glutaraldehyde, and 0.1 M phosphate-buffered saline (PBS; pH 7.4) for 3 hours; and washed in 0.1 M PBS overnight at 4°C. The tissue was postfixed for 1 hour at 4°C in 2% osmium tetroxide in rinse buffer. Dehydration, infiltration with Epon resin, and polymerization of samples was performed by conventional methods. Ultrathin sections (70–90 nm) were cut on a microtome (Ultrotome; Leica, Heidelberg, Germany), collected on grids, and counterstained successively in 1% aqueous uranyl acetate and 1% aqueous lead citrate for 2 minutes each. Sections were finally washed, rotary, cut on a microtome (Rotaput; Slidewax; Leica, Heidelberg, Germany), collected on grids, and stained with 1% toluidine blue in 1% aqueous sodium tetraborate for 1 minute.

Western Blot Analysis

To ensure correct cross-reactivity of the different antibodies, we ran Western blots for each on extracts prepared from A. ansorgei and rat (Rattus norvegicus) retinas (two retinas from each). Retinas were collected and homogenized in lysis buffer (20 mM Tris-Cl; pH 7.4, 150 mM NaCl, 1% Triton X-100, 0.1% SDS, 1 mM EDTA) containing a protease inhibitor cocktail (Roche Molecular Biochemicals, Meylan, France). Measurement of total protein concentrations in aliquots from each sample gave values of 18 mg/mL for the rat and 18.8 mg/mL for A. ansorgei. Proteins (5 μL/well) were separated by 10% SDS-polyacrylamide gel electrophoresis and transferred to nitrocellulose membranes with a semidy filter (Bio-Rad, Munich, Germany). The membranes were blocked with TBS containing 0.1% Tween 20 and 3% fat-free powdered milk (pH 7.3) for 1 hour at room temperature. Membranes were incubated with the following primary antibodies: monoclonal anti-rhodopsin antibody Rho-4D2, polyclonal anti-mouse mid-wavelength (MW)-cone and short-wavelength (SW)-cone opsins, and polyclonal anti-mouse cone arrestin. Each primary antibody was diluted to a final concentration of −0.1 μg/mL (1:10,000 dilutions of ascites fluid or antiserum, respectively) and incubated overnight at 4°C, the membranes were washed thoroughly and then incubated with goat anti-mouse or goat anti-rabbit IgG-horseradish peroxidase secondary antibodies (1: 20,000; Jackson ImmunoResearch Laboratories, West Grove, PA). Immunoreactive bands were visualized with chemiluminescence (Super Signal ECL kit; West Pico; Pierce, Rockford, IL), according to the manufacturer’s instructions. Apparent molecular masses were estimated by comparison to prestained molecular size markers (Invitrogen-Life Technologies, Gaithersburg, MD). Western blot analyses were performed for each antibody in two independent experiments, and representative images are shown.

Immunohistochemistry

Eyes were fixed overnight in 4% paraformaldehyde at 4°C, transferred to an ascending series of sucrose solutions (10%, 20%, and 30%, each for 2 hours) and embedded (Tissue-Tek; Sakura Finetek, Tokyo, Japan), and 18-μm-thick cryostat sections were prepared and stored at −20°C until ready for use. The sections were permeabilized with Triton X-100 (0.1% in PBS for 5 minutes) and then saturated with PBS containing 0.1% BSA, 0.1% Tween-20, and 0.1% sodium azide (buffer A) for 30 minutes. Sections were incubated overnight at 4°C with the following primary antibodies diluted in buffer A. Antibodies used were monoclonal anti-rhodopsin antibody Rho-4D2, polyclonal anti-mouse MW-cone and SW-cone opsins, polyclonal anti-mouse cone arrestin, and monoclonal anti-RPE65 (Chemicon International Ltd., Temecula, CA). Secondary antibody incubation was performed at room temperature for 2 hours with Alexa (488 or 594) goat anti-rabbit or anti-mouse IgG-conjugated antibodies (Invitrogen, Eugene, OR). Cell nuclei were stained with 4,6-diamino-phenylindolamine (DAPI; Invitrogen). Slides were washed thoroughly, mounted in PBS and glycerol (1:1), and observed with a confocal laser scanning microscope (LSM 510 ver. 2.5; Carl Zeiss Meditec, Jena, Germany), a scanning device mounted on an inverted microscope (Axiovert 100; Carl Zeiss Meditec), or a fluorescence microscope (Optiphot 2; Nikon, Melville, NY) equipped with differential interference contrast optics. Images were relayed by charge-coupled device (CCD) camera video capture to a dedicated computer containing image analysis software (Visiolab; Biocom, Lyon, France).

Retinal Wholemount Immunohistochemistry

Two eyes were used to prepare retinal wholemounts. Before excision, a small incision was made in the dorsal margin of the ora serrata for orientation. Retinas were carefully dissected to preserve orientation; fixed in 4% paraformaldehyde, PBS, and 5% sucrose for 4 hours at room temperature; washed with 0.1 M Tris-buffered saline (pH 7.4; TBS) and water (5 minutes each), and dip frozen in chilled acetone (−20°C for 5 minutes). After washes in water and TBS (5 minutes each), the retinas were incubated for 2 hours in 10% H2O2 in PBS with 1 drop of 30% ammonia to bleach pigment granules in the RPE and visualize the retina more clearly. The retinas were then washed with PBS and permeabilized in 0.3% Triton X-100 for 1 hour and saturated in blocking buffer overnight and incubated with either rho-4D2 or anti-SW-cone opsins antibodies for 1 day at ambient temperature. After extensive washing (8 hours with six changes of buffer), the retinas were incubated with the corresponding secondary antibody for 24 hours at 4°C, cut into four quadrants (dorsal, ventral, temporal, and nasal), using the initial incision as a reference, and flattened under coverslips in 1:1 PBS-glycerol. For quantification at each of the selected retinal locations, tissue was divided into near (within 1.5 mm of the optic nerve head), mid (1.5–3 mm from the optic nerve head), and far (>3 mm from the optic nerve head) regions for each of the four quadrants, and blue cone densities were estimated for each zone by counting the average number of SW cones in 20 randomly chosen 150 × 150-μm fields viewed with a ×40 objective lens. Because total cell-packing density did not differ greatly across the retina, these values are expressed as the mean number per field.
Quantification of Phagocytosis during the Light–Dark Cycle

Animals used to study temporal variations in phagocytosis were euthanatized every 3 hours during the 12-hour light–dark cycle (n = 4 for each time point). According to the 24-hour clock, lights were turned on at 0800 hours (8 AM) and off at 2000 hours (8 PM), and animals were killed at midnight (2400 hours) and 0300, 0600, 0900, 1200, 1500, 1800, and 2100 hours. Eyes were rapidly removed and fixed in 4% paraformaldehyde, as described earlier. For samples taken during the dark phase, enucleation was performed under dim red light illumination. Sections were obtained across the entire width of the retina, from the superior to inferior margin, passing within 1 mm of the optic nerve head. Four sections from one eye of each animal at each point were analyzed (i.e., a total of 16 sections per time point). They were immunostained as detailed herein, using either rho-4D2 (rods) or anti-MW-cone opsin (MW cones). OS staining of both PR populations was very regular, the leading edge forming a straight border. Taking this edge as a baseline, any immunopositive inclusion of ≥1 μm lying within the RPE subcellular space (visualized by faint background lighting) was scored as a phagosome. Counting of phagosomes was achieved by aligning a 150 × 150-μm grid placed within the eyepiece, parallel with the RPE layer. Taking the optic nerve head as reference, the grid was displaced dorsally and ventrally along the OS/RPE interface, from the posterior to the superior margin. In this way, the RPE was sampled across the entire width of each eye.

Statistics

For statistical analysis of SW-cone densities and phagosome uptake, we used a one-way analysis of variance (ANOVA) followed by an ad hoc Fisher least-significant difference test, with acceptance levels of P < 0.05.

RESULTS

Morphologic Characterization of the Retina of A. ansorgei

The retinal morphology of Arvicanthis represented in Figure 1 shows a typical tiered structure. The RPE is visible as a clear basal region containing nuclei and mitochondria, and a heavily pigmented apical area packed with melanin granules. Within the outer nuclear layer (ONL), nuclei of cone PRs can be distinguished from those of rods by the less-intense staining, corresponding to less-condensed heterochromatin.

At the ultrastructural level, the RPE–OS interface demonstrated an orderly arrangement of rod and cone OS apposed to the RPE apical surface (Fig. 2). Rod OS appeared as cylindrical structures, the stacked discs clearly visible. Cone OS were narrower and tapered, slightly more electron dense, with areas of clear cytoplasm and surrounding cone matrix sheath. RPE exhibited many melanin granules positioned within the apical region. Occasional phagosomes of rod or cone origin were equally visible (Fig. 2A, arrows). Individual RPE cells were in contact with both rod and cone OS.

Western Blot Analysis of PR Antibodies

We performed comparisons of the different PR antibodies between rat and Arvicanthis whole retinal extracts. As shown in Figure 3, each antibody gave similar results in terms of the apparent molecular mass species detected: MW-cone opsin immunoreactivity was observed at ~40 kDa, a faint band in the rat retina but a much more intense signal in the Arvicanthis retina (Fig. 3A; lanes 1, 2). SW-cone opsin immunoreactivity was seen at the same size range (~40 kDa) and was approximately of equal intensity in both rodents (Fig. 3A, lanes 3, 4); cone arrestin immunoreactivity was detected at ~48 kDa, being very faint in the rat and strong in the Arvicanthis (Fig. 3A, lanes 5, 6). Rho-4D2 anti-rod opsin immunoblots were performed in two different conditions, using heated and unheated samples. In both rodents, rod opsin immunostaining was generally comparable, although somewhat less intense in the Arvicanthis. Boiled samples contained immunoreactive bands at 54, 80, 115, and >140 kDa (Fig. 3B, lanes 1, 3), and unheated samples also showed strong staining of a band at ~34 kDa (Fig. 3B, lanes 2, 4). These immunoblot data provided evidence for correct cross-reactivity of the different antibodies between the rodent species.

Rod and Cone Immunohistochemistry

Immunolabeling of the A. ansorgei retina was performed with different rod and cone antibody markers. Anti-mouse MW-cone opsin stained abundant OS, forming a dense continuous band of label across the retina (Figs. 4A, 4B). There was no antibody binding detectable in other cellular locations. Anti-mouse SW-cone opsin labeled only a few small structures within the OS (Figs. 4C, 4D). By contrast, in sections obtained from dark-adapted animals, anti-mouse cone arrestin produced intense staining throughout the cone cells, filling the OS and cell bodies within the uppermost two layers of the ONL and extending down their axons to the synaptic pedicles (Figs. 4E, 4F). Rhodopsin immunostaining produced an intense signal in the OS, forming a somewhat broader band than that observed.
for MW-cone opsin, and in addition outlined rod cell bodies in the inner two-thirds of the ONL (Fig. 5). These data show that the ONL is separated into a distal region composed exclusively of cone cell bodies (two rows) and an inner region composed exclusively of rod cell bodies (four rows). This anatomy was apparent uniformly across the retina, from center to mid periphery, but toward the far periphery, the ONL thinned to four to five cell bodies, with only a single distal row of cones, and three to four rows of rods (not shown).

We also immunolabeled frozen sections of *A. ansorgei* eyes with anti-RPE65 monoclonal antibody. As seen in Figure 6, antibody staining was intense and specific within the RPE layer.

**Wholemount Immunolabeling of A. ansorgei Retina**

To obtain a more comprehensive distribution pattern of SW cones, we performed immunostaining of flatmounted retinas (Fig. 7A). Whereas rod opsin flatmount labeling revealed numerous immunopositive OS throughout the retinal fields (data not shown), SW cones were sparsely distributed across the retinal surface. The distribution pattern is shown in Figure 7B, mapped against a camera lucida depiction of the retina, and in histogram form for each of the 12 regions analyzed (Fig. 7C). There were no significant differences in the number of SW cones between left and right eyes (not shown) or between any of the four quadrants or the superior and inferior hemispheres, but cone density was significantly higher in the central annulus than in the far periphery (an overall threefold difference; *P* < 0.001). There were also significantly higher densities in the mid periphery than in the far periphery (*P* < 0.01).

**Daily Rhythms in Rod and Cone Phagocytosis**

Immunoreactive rod or cone debris were identifiable unambiguously within the RPE cytoplasm. An examination of sections obtained at different time points within the standard 12-hour light–dark cycle revealed the presence of inclusions labeled with rod opsin antibody at 0900 hours (1 hour after light onset), but not at 2100 (1 hour after light offset; Figs. 8A, 8B). Similar analyses using an MW-cone opsin antibody showed a similar profile: immunopositive inclusions were reproducibly seen at 0900 but not at 2100 hours (Figs. 8C–F). Simultaneous visualization of both rod and cone phagosomes at 0900 hours did not reveal any obvious association between the two populations, although they were clearly both present within individual RPE cells (Fig. 9). Given the low number of SW-cones we did not attempt to examine phagocytosis in this subpopulation.

**FIGURE 2.** Electron microscopic images of the PR–RPE interface in adult *A. ansorgei*. Shown are representative views of rod (ROS) and cone (COS) outer segments apposed to the RPE. COS were identified from their tapered shape and lower position, slightly greater electron density, and greater artifactual damage during tissue processing. In addition, traces of cone matrix sheath were visible surrounding them (open arrows). Several phagosomes (Ph) are visible within the RPE cytoplasm in both panels (tissue obtained at 0900 hours). Ch, choroid; M, melanin granules. Scale bar, 1 μm.

**FIGURE 3.** Western blot analysis of mouse rod- and cone-specific antibodies in retinal extracts from adult *A. ansorgei* and *R. norvegicus*. Total proteins isolated from retinas of either species were run in parallel and immunoblotted with cone (A) or rod (B) antibodies. (A) Lanes 1 and 2: MW-cone opsin; lanes 3 and 4: SW-cone opsin; lanes 5 and 6: cone arrestin. Lanes 1, 3, 5: rat retina; lanes 2, 4, 6: *A. ansorgei* retina. Whereas immunoreactive bands were of the same apparent mass in both species, MW-cone opsin and cone arrestin were much more highly expressed in *A. ansorgei* retina. SW-cone opsin intensity was equal in both species, MW-cone opsin and cone arrestin were much more highly expressed in *A. ansorgei* retina. SW-cone opsin intensity was equal in both species, MW-cone opsin and cone arrestin were much more highly expressed in *A. ansorgei* retina.

**FIGURE 4.** Western blot analysis of mouse rod- and cone-specific antibodies in retinal extracts from adult *A. ansorgei* and *R. norvegicus*. Total proteins isolated from retinas of either species were run in parallel and immunoblotted with cone (A) or rod (B) antibodies. (A) Lanes 1 and 2: MW-cone opsin; lanes 3 and 4: SW-cone opsin; lanes 5 and 6: cone arrestin. Lanes 1, 3, 5: rat retina; lanes 2, 4, 6: *A. ansorgei* retina. Whereas immunoreactive bands were of the same apparent mass in both species, MW-cone opsin and cone arrestin were much more highly expressed in *A. ansorgei* retina. SW-cone opsin intensity was equal in both species, MW-cone opsin and cone arrestin were much more highly expressed in *A. ansorgei* retina. SW-cone opsin intensity was equal in both species, MW-cone opsin and cone arrestin were much more highly expressed in *A. ansorgei* retina.
Quantification of rod and cone phagosomes was performed throughout the light–dark cycle. These analyses confirmed the rhythmic nature of phagocytic uptake of both populations and revealed that the two profiles were largely similar (Fig. 10). Both showed large peaks of phagosome numbers at around 0900 hours, significantly different from all other time points. A second smaller peak was also visible for rod phagosomes, centered on 0300 hours but detectable at both 2400 and 0600 compared with the minimum values observed at 1800 and 2100 hours (Fig. 10A). The number of cone phagosomes was always much lower than rods, by a factor of 10 (Fig. 10B). Phagosomal activity was, however, quite variable, since at any given time the number of phagosomes generally varied by a factor of 2 between individual animals and sometimes by as much as 10-fold. In addition, these differences were not consistent with respect to rods or cones (i.e., animals showing the lowest number of rod phagosomes did not necessarily also exhibit a low number of cone phagosomes; Table 1). There was some correspondence between a low number of phagosomes and age, since the two lowest values for both rod and cone phagosomes were observed in the two oldest individuals (Table 1).

**DISCUSSION**

We present the initial characterization of a diurnal murine rodent with respect to PR phenotype, distribution, and rhythmic phagocytosis. The data show that cones form a large percentage of total PRs, approximately 33%, and that they undergo cyclic phagocytic turnover in a general pattern similar to that of rods.

Rodent species typically used for experimental research, namely laboratory rats (*Rattus norvegicus*) and mice (*Mus*...
animals for studying cone PRs, such as chickens. These animals have aided in exploring PR differentiation, relationship between circadian clock gene expression, suprachiasmatic nucleus function, and diurnal behavior. Despite the implication of photic stimuli in regulation of activity, to the best of our knowledge, there are no published data referring to retinal neuroanatomy.

All tested antibodies, raised against either mouse or rat sequences of retinal proteins, cross-reacted fully with *Arvicanthis*. The anti-mouse cone arrestin antibody demonstrated a very similar subcellular distribution in dark-adapted *Arvicanthis* retinal sections (Fig. 4) compared with those obtained from dark-adapted mice. We conducted mapping studies of SW-cones across the entire retina, since it has been reported that the two cone spectral classes present in lower mammals are distributed in a heterogeneous pattern, with SW-cones predominantly located in the ventral field. Although we observed a significantly higher number of SW cones in the central field than in the far periphery, there was no detectable variation in their number as a function of retinal quadrant. Relatively homogenous distributions have also been seen in other murine genera. Although we did not perform similar fine mapping studies for MW cones, analysis of a large number of sections taken across all retinal areas did not reveal major differences (data not shown). Western blot comparisons of relative abundance of MW-cone opsin and cone arrestin between rats and *Arvicanthis* showed expected large excesses for the latter species; unexpectedly, the abundance of SW-cone opsin appeared virtually identical between the two. This similarity is presumably due to a low SW- to MW-cone ratio, a lower amount of SW-cone opsin per cell, or both. Electrophoretic separation of rhodopsin frequently produces multimeric aggregates, especially on heating of samples under non-reducing conditions which converts virtually all monomeric
forms into dimers or higher complexes. We used this anomalous electrophoretic migration behavior to underline that rhodopsin antibody specificity was retained for *Arvicola hisbis*. The intensity of bands was noticeably fainter in *Arvicola hisbis* preparations, consistent with the reduced rod population.

An important aspect that has been difficult to study in mammals concerns cone phagocytosis by the apposing RPE. Autoradiographic studies led to the conclusions that protein (mainly opsin) synthesis occurred in the inner segment and that newly synthesized membranes (i.e., discs) were displaced apically by continuous addition at the basal surface of the OS. Since this time, extensive data have been gathered concerning the chain of events linking OS shedding and their subsequent internalization by the RPE. Initially, OS bind to the RPE apical surface, triggering activation of an intracellular second-messenger cascade, which in turn activates the ingestion of bound OS. CD36, MerTK, and αVβ5-integrin receptors have all been implicated as playing crucial roles in phagocytosis. The macrophage scavenger receptor CD36 was found to regulate the rate of OS internalization. Binding of OS to the receptor tyrosine kinase c-Mer (MerTK) leads to phosphorylation and activation of the second-messenger cascade.

Binding of OS requires the activation of αVβ5-integrin receptors, which interact with MerTK via the focal adhesion kinase (FAK) signaling pathway. In brief, OS binding to αVβ5-integrin receptors activates FAK, which in turn phosphorylates MerTK, leading to activation of the second-messenger cascade. Finally, the specific ligand for MerTK is growth-arrest–specific protein 6, which is necessary for OS phagocytosis. Perturbation in the phagocytosis pathway can lead to retinal degeneration, as observed in the RCS rat and a subset of patients with retinitis pigmentosa.

These molecular details are only known for certain for rod uptake. Contrary to rods, cones appeared to be more static elements, since they did not exhibit a band of radioactivity—rather diffuse labeling of the OS. However, phagocytic uptake of shed cone membrane debris was described in the cone-dominated retinas of lizards and squirrels. This latter group also observed unequivocal cone OS uptake in cats, monkeys, and humans. An important goal of the present study was to quantify rod and cone phagocytosis as a function of daily time. Previous investigations have shown a strong rhythmic component of phagosome uptake, entrained by day/night alternation and/or circadian control. In all nocturnal and diurnal vertebrates examined, both cold- and warm-blooded, rod OS uptake occurs maximally at or shortly after light onset. In rats, this daily rhythm is established by 2 weeks of age and is irrespective of the lighting conditions during development. It persists for up to 14 days in constant darkness and is independent of the brain’s biological clock, since it is maintained after optic nerve section or destruction of the suprachiasmatic nucleus. Thus, an intrinsic oscillator exists within the eye, although feedback from the brain is needed to synchronize the circadian shedding rhythm with the day–night cycle. The timing of cone shedding is more variable, since in
most species (goldfish, lizards, chickens, and tree squirrels) it occurs during the night time, whereas in rhesus monkeys, tree shrews, and cats rod and cone OS shedding both occur maximally just after light onset. The data from *Arvicansita* resemble those obtained in cats in several respects: (1) Rod and cone shedding occur together shortly after light onset. Although there are large differences between the animals in lifestyles and retinal anatomy (there is no obvious area centralis in Nile rats), *Arvicansita* represents a further example of maximum cone shedding occurring at light onset. The four mammalian species in which such dynamics are observed are all active during the day. (2) There is a secondary, smaller increase in rod shedding late during the night period. This smaller peak occurred around the same time in both animals (~6 to 8 hours after light offset). (3) The maximum number of rod phagosomes greatly exceeds that of cones. In both species rod phagosomes outnumbered cone phagosomes by a factor of 10. This may reflect the different turnover rates of rods versus cones, because rod membranes are thought to be renewed more rapidly than cones. However, there is controversy over the issue, and estimates in the cat give similar values for both PR types. (4) Variability in the number of phagosomes between individuals was observed at any given time point. This variability was also seen in cats, and, as in this study, differences between individuals were not consistent (i.e., those showing a high number of rod phagosomes could exhibit a low number of cones, and vice versa). One source of variation could be age, since the eldest animals showed the lowest number of inclusions. We are currently performing similar analyses under constant dark conditions to see whether rod and/or cone phagocytosis are circadian or light driven.

The methodological approach used herein to quantify rod and cone phagosomes differs from that used in previous studies. Whereas purely morphologic criteria have been used by other workers, we exploited the availability of rod- and cone-specific immunologic probes to map simultaneously the appearance and distribution of both populations. In distinguishing rod from cone phagosomes, previous studies have relied on species exhibiting large cone OS, or regional specializations (macula or area centralis). Rodents possess small, narrow cone OS, and similar approaches would not be feasible. The use of

**Table 1.** Total Data from All Individuals Counted during the Experiments Shown in Figure 10

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<tr>
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<td>7</td>
<td>6</td>
<td>6</td>
<td>5</td>
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<tr>
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<td>445</td>
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<td>Age (mo)</td>
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<td>6</td>
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<td>11</td>
<td>107</td>
<td>20</td>
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<td>18</td>
<td>9</td>
<td>7</td>
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</tbody>
</table>

Data are replotted as time of day (0900–1800 represent daylight, D; 2100–0600 represent nighttime, N with average phagosome count from all four sections for a given individual. Also given are the ages of each animal. This representation illustrates the variation that was recorded between individuals for a given time point, and the inverse correlation between phagosome number and age.
antibodies directed against the integral membrane protein visual pigments has the advantage of allowing screening of a large number of sections and of performing ultrastructural immunohistochemistry. Rhodopsin antibodies have been used previously to examine rod-derived phagosomes in the RPE at the electron microscope level. An important caveat is that the epitopes recognized by each antibody will be destroyed during the enzymatic degradative process, and hence do not allow us to observe phagolysosome metabolism. We also noted that cone arrestin antibodies did not label phagosomes, presumably because the cytoplasmic antigen leaks rapidly from incipient phagosomes (data not shown). Finally, we made no distinction between ‘shedding’ and ‘phagosomes,’ since our counting procedure would have excluded incipient packets of disc present outside the RPE.

In conclusion, we present the initial characterization of a diurnal mouselike rodent with a large number of histologically and immunologically identifiable cones. We have mapped rod and cone distribution and composition and quantified rod and cone phagosome uptake as a function of daily time. Our future studies will focus on cone phagocytosis, of particular importance with respect to human macular function and health and possibly centrally involved in the progression of macular degeneration.

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References


