Circadian-clock driven cone-like photoreceptor phagocytosis in the neural retina leucine zipper gene knockout mouse

Arthur Krigel, Marie-Paule Felder-Schmittbuhl, David Hicks

Département de Neurobiologie des Rythmes, CNRS UPR 3212, Institut des Neurosciences Cellulaires et Intégratives, Strasbourg, France

**Purpose:** Whereas much information is available on rod outer segment phagocytosis by the retinal pigmented epithelium (RPE), corresponding data for cones are quite limited, especially in laboratory models of normal rats and mice with very low cone numbers. To characterize the light and circadian control of cone photoreceptor phagocytosis in mice, we capitalized on the blue cone-like phenotype of neural retina leucine zipper gene (Nrl) null mice (Nrl−/−).

**Methods:** Nrl−/− mice were maintained under standard cyclic light (12h:12h light-dark [LD] cycle; light=300 lux) for one month, then divided into two groups: 1) continued maintenance in LD (36 mice); or 2) transferred to constant darkness (DD; 21 mice) for 36 h. Animals were sacrificed every 3 h over 24 h, and their eyes were rapidly enucleated and fixed. Cryosections were stained using specific cone short-wavelength opsin antibodies. Phagosome numbers in the RPE were quantified with a morphometric system. We monitored the expression of c-mer proto-oncogene tyrosine kinase (MerTK) in wild-type and knockout mice using a specific MerTK antibody.

**Results:** In LD, cone phagocytosis showed a statistically significant peak of activity 1 h after light onset, 2–3 fold higher than at other times. In constant darkness, the temporal phagocytic profile resembled that of LD (significant peak at 1 h of subjective day), but the number of phagosomes was decreased at all time points. Immunostaining of MerTK in wild-type and Nrl−/− mice showed expression at the apical surface of the RPE.

**Conclusions:** Cone-like outer segment phagocytosis in Nrl−/− mice shows a similar profile to that of rods in normal mice and other species. These data are the first to quantify blue cone-like photoreceptor phagocytosis under different lighting conditions in mice, and suggest this model may constitute a valuable system for investigating circadian regulation of cone function.

Retinal photoreceptors (RP) are composed of two different populations, rods, and cones. Rods are used for nocturnal vision because of their high light sensitivity [1], whereas cones require a relatively high light level to activate them, and are used for diurnal vision [2]. The latter are responsible for color discrimination and high-acuity vision, and are of utmost importance in human vision. In humans and Old World primates, there are three types of cones, containing blue, green, and red light-sensitive pigments (or short wavelength sensitive (S) cones [3], and middle/long wavelength sensitive cones, respectively). In the great majority of mammals, there are only two cone types: a short wavelength-sensitive and a mid/long wavelength-sensitive population [4].

Both rods and cones are known to undergo continuous cyclic turnover, involving the addition of new membranes at the base of the outer segment (OS) and the removal of aged membranes at the distal pole [5]. This removal is achieved by the opposing retinal pigmented epithelium (RPE), which phagocyte and digest the shed OS membranes [6]. In recent years, much progress has been made in identifying molecular components of the phagocytic pathway, including membrane-bound receptors such as MerTK and αVβ5 integrin, and ligands [7-9]. Mutations in the Mertk receptor lead to inherited retinal degeneration in animals [7,10] and humans [11]. The great majority of data have been obtained for rods, in part because conventional laboratory rodent (mouse and rat) retinas are composed of 97% rods and only ~3% cones [12, 13]. A genetically modified mouse line has become available for the study of rod differentiation, the neural retina leucine zipper gene (Nrl) knockout mouse (Nrl−/−) [14]. Deletion of the transcription factor Nrl results in the complete absence of rods, as revealed by histology, immunocytochemistry, electrophysiology, and gene expression analysis [14,15]. Morphological, molecular, and electrophysiological features of the Nrl−/− photoreceptors seem to be identical to blue or short wavelength light-sensitive S cones [14]. Hence, this retina provides a potential means for the investigation of blue cone function and cone-specific genes [16].

In the present study, we have used homozygous Nrl−/− mice to ask whether cone phagocytosis shows daily rhythms, if any such rhythms resemble or differ from those known for rods, and whether these rhythms are maintained under constant darkness. We show that rhythmic phagocytosis does
indeed occur, in both cyclic light and constant darkness, and that it resembles rod behavior.

METHODS
Animals and handling: This study was conducted with Nrl−/− mice, raised in our animal facilities. The strain was originally obtained from Dr. C. Grimm at the Laboratory of Retinal Cell Biology, University Hospital, Switzerland, with permission from Dr. A. Swaroop (NIH, Bethesda, MD). The colony was maintained on a 12 h:12 h light-dark (LD) cycle in an ambient temperature of 22 °C for four weeks. Young adult wild-type C57Bl/6 mice (n=3), the strain on which the knockout line was created, were used as a positive control for antibody binding and specificity. All experiments were conducted according to ethical guidelines in operation at the institute, and adhered to the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. Two experimental series were made (shown schematically in Figure 1): in the first group, a total of 36 knockout mice were used, composed of mostly young adults (2 months, n=30), with some individuals of 5 months (n=6). Older individuals were spread equally among sample points to eliminate any age-related bias. Animals were euthanized every 3 h during the LD cycle (n=4 for each time point). Lights were turned on at 7 AM, defined as zeitgeber time, hour zero (ZT0). Animals were killed at ZT1, 4, 7, 10, 13, 16, 19, and 22, and at an additional point in the following 24 h cycle (ZT24+1). The second group of animals (n=21) was raised in LD until they reached young adulthood (2 months). The animals were then placed in total darkness (constant darkness [DD]), for 36 h, and therefore experienced a complete cycle of subjective day and night before sample collection. Euthanasia commenced at circadian time (CT) 1 (i.e., 1 h after subjective light onset), and continued every 3 h at CT4, 7, 10, 13, and 16 (n=3 per time point).

In both groups during the night time (i.e., ZT12–24 for LD cycles and permanently for DD), darkness was total (no background dim red light), and euthanasia and enucleation were performed using night vision goggles. For every time point and in both groups, eyes were rapidly removed by dissecting the surrounding ocular muscle and optic nerve. A small hole was made with a 25-gauge hypodermic needle at the level of the ora serrata, and eyeballs were fixed overnight in 4% paraformaldehyde in buffered phosphate saline (0.1 M PBS; l. 91, NaCl 7.34 g/l, KCl 270 mg/l, Na$_2$HPO$_4$ 10 mg/l, NaH$_2$PO$_4$ 3 mg/l, pH 7.3), at 4 °C.

Immunohistochemistry and quantification: Eyes were rinsed in cold PBS for 1 h, then transferred to an ascending series of sucrose solutions (10%, 20%, and 30%, each for 2 h) and embedded in optimal cutting temperature resin (Tissue Tek; Sakura Fintek, Tokyo, Japan), and 10 μm thick cryostat sections were prepared and stored at −20 °C until ready for use for staining. Immunostaining and quantification of phagosomes was performed as published previously by us [17]. Briefly, sections were permeabilized and then saturated with PBS containing 3% BSA, 0.1% Tween-20, and 0.1% sodium azide (buffer A) for 30 min. Sections were incubated overnight at 4 °C with polyclonal anti-mouse S-cone opsin.

![Figure 1. Schematic diagram showing schedule of lighting conditions and sampling points. Arrow in series 2 indicates times at which animals were switched to constant darkness condition. Thin vertical bar joined by a horizontal bar indicates the beginning and end of the sampling period. Series 1 is the control condition, with animals maintained under light-dark conditions (alternating white bars [12 h light=300 lux] and black bars [12 h dark]) throughout the experiment. Series 2 represents total darkness (alternating black dots on a white background [subjective day] and black bars).](http://www.molvis.org/molvis/v16/a308)
antibody [18] (the generous gift of Dr. Cheryl Craft, Doheny Eye Institute, University of California, Los Angeles, CA) at a ~1 μg/ml final concentration in buffer A. Antibody binding was visualized using secondary fluorescent conjugated anti-rabbit IgG-Alexa 594 (Molecular Probes Ltd., Eugene, OR). Since the Nrl−/− retina exhibits partial degeneration with a scalloped appearance of the outer nuclear layer and the formation of rosettes [14], quantitative measurements were performed only in areas devoid of rosettes and degenerated cells. Any immunopositive structure with a diameter of ≥1 μm lying within the RPE sub-cellular space (visualized by faint background lighting to show pigmentation) was scored as a phagosome. Counting of phagosomes was performed by aligning a 150×150 μm grid placed within the eyepiece, parallel with the RPE layer [17]. The phagosome counts are expressed as the sum of all four sections per eye.

For comparative studies on phagocytic activity among different species, we also processed frozen sections of adult Wistar rats and adult C57Bl6 mice, from animals killed at ZT1. These sections were only stained with anti-rhodopsin (rhop4D2) antibody [19] as described above, and analyzed for rod phagosome numbers using the same morphometric grid procedure.

We also performed immunostaining for MerTK using a rabbit polyclonal MerTK antibody (the generous gift of Dr. G. Lemke, Scripps Institute, La Jolla, CA) [20] to verify the expression and distribution of MerTK in the RPE of Nrl−/− mice, compared to wild-type mice. Wild-type mouse retinal sections were double-immunolabeled with MerTK and monoclonal anti-rhodopsin antibody Rho-4D2 antibody [19] as described above, and analyzed for rod phagosome numbers using the same morphometric grid procedure.

RESULTS

Figure 2 shows a representative image of Nrl−/− mouse retinal sections stained with anti-S-cone antibody. Heavy staining of the cone OS (COS) region is seen, corresponding to concentration of S-cone pigment in this cellular compartment. In sections made from mice killed at ZT1 under LD conditions, there are scattered discrete fluorescent dots visible within the pigmented band of RPE (arrows, Figure 2A). In sections made at ZT19, COS staining is unchanged, but there are fewer fluorescent inclusions seen in the RPE (Figure 2B). Similarly, in sections made from mice maintained in DD and killed at CT1, there are also immunopositive spots in the RPE (arrows, Figure 2C), whereas retinal sections from mice killed at CT19 contain fewer such profiles within the RPE layer (Figure 2D). Immunostaining of wild-type mouse retinas with anti-S-cone opsin shows the occasional labeled COS (Figure 2E).

Quantification of fluorescent inclusions within the RPE layer showed marked fluctuations as a function of time of day. Under LD conditions, there was a distinct peak at ZT1, with phagosome numbers between 2 and fourfold higher than at all other time points (Figure 3A). This value was highly significantly different from all other time points in the day or night. Phagosome numbers fell quickly to baseline levels, to roughly 50% lower at ZT4 and 70% lower by ZT7. There was a tendency toward a smaller second peak at ZT13 (1 h after lights-off), but this did not attain statistical significance. Thereafter, phagosome numbers were minimal (30%–40% the values at ZT1), before rising steeply again following the night-day transition. A very similar temporal profile was seen when Nrl−/− mice were maintained under DD conditions for 36 h before sampling. Although the amplitude of phagocytosis was reduced by approximately 40%–50% at all time points, the shape matched closely that seen in LD, with a pronounced peak at CT1, some 2-4 fold higher than other points (Figure 3B).

To see whether Nrl−/− mice expressed the MerTK receptor, we also performed immunolabeling of wild-type and knockout mice retinal sections with a specific MerTK antibody. Wild-type mouse retinas underwent double immunolabeling against rhodopsin and MerTK. The rod OS were intensely immunolabeled against rhodopsin antibody (Figure 4A). The MerTK immunostaining pattern was similar, with staining of the sub-retinal space corresponding to the apical microvilli of the RPE (Figure 4B). Merged images showed that the MerTK distribution extended almost the length of the OS (Figure 4C). In Nrl−/− mice retinas, as before, S-cone opsin staining was prominent in the shortened cone OS (Figure 4D). MerTK was localized to the interface between the RPE monolayer and outer nuclear layer (ONL), the staining being limited to a narrow band and not extending into the sub-retinal space as in the wild-type retina (Figure 4E).
The general morphology of the posterior pole in Nrl−/− mice is shown in Figure 4F.

To see whether the overall phagocytic activity of blue cones in Nrl−/− mice resembled values for “normal” rods and cones from other rodent species and strains under similar conditions, we calculated the number of phagosomes (rods and cones) seen at peak amplitude (ZT1/CT1), and normalized this to the number of rod- or cone-cell bodies within the same region. To reduce variability from the use of different techniques, we analyzed samples from normal wild-type mice, laboratory Wistar rats, and the diurnal rodent Arvicanthis ansorgei, all assayed with our immunohistochemical procedure. We also included data from published reports on two other species in which cat and...
Figure 3. Cone shedding in Nrl−/− mice shows rhythmic activity in both cyclic light and constant darkness. Phagosome counts were made every 3 h throughout both the light and dark periods under light-dark (LD) conditions (A) and through subjective light and dark periods under constant darkness (DD) conditions (B). A burst in the number of cone phagosomes was seen at ZT1 (LD) and at circadian time (CT1; total darkness), significantly different from all other values (***p<0.001). Phagosome numbers were lowest in late daytime.
chicken cone phagocytosis has been examined. Table 1 expresses these values per 100 cell bodies. In four of the five species examined, peak cone activity was quite similar (8–12 phagosomes/100 cones); only in the cat were numbers considerably higher (20 phagosomes/100 cells). Rod phagosome frequencies in normal mice and rats were very close to the values for cone phagosomes in Nrl<sup>−/−</sup> mice, whereas rod activity was much higher in Arvicanthis (38 phagosomes/100 rods).

Table 1. Comparison of phagosome numbers between rods and cones and between different species.

| Species       | Peak cone value | Peak rod value | References |
|---------------|-----------------|----------------|------------|
| cat           | ~20 per 100 cones | -              | [25]       |
| chick         | ~12 per 100 cones | -              | [21]       |
| Arvicanthis   | ~9 per 100 cones  | ~38 per 100 rods | Present study |
| Wistar rat    | -                | ~7 par 100 rods  | Present study |
| C57Bl6        | -                | ~6 par 100 rods  | Present study |
| Nrl<sup>−/−</sup> | ~8 per 100 cones  | -              | Present study |

To compare numbers between different retinas, these were calculated as a fraction of 100 cell bodies, either from our own data (rats, mice, Arvicanthis) or from previously cited information (cat, chicken). The percentage of phagosomes is quite similar between most of the different examples, except for the elevated rod phagosome number in Arvicanthis.
DISCUSSION

We have used the all-cone Nrl\(^{-/-}\) mouse to document the temporal profile of cone phagocytosis under two lighting conditions—standard cyclic light and permanent darkness. The data show that mouse S cone-like photoreceptors, like rods, exhibit a daily rhythm in OS debris shedding/uptake by the RPE, and that this process shows a marked peak in activity shortly after light onset.

Cyclic light and/or circadian-driven rhythms in photoreceptor OS phagocytosis have been extensively explored in many species. Peak rod phagocytosis seems to be invariant, and occurs shortly after light onset in every species (nocturnal and diurnal) that has been studied, (e.g., chickens [21], mice [22], rats [23], ground squirrels [24], cats [25], and rhesus monkeys [26]). Data on cone phagocytosis are less abundant but show more irregularity. Most studies show maximal cone shedding in the night, as in diurnal chickens [21], fence lizards [27], tree squirrels [28], ground squirrels [24], and rhesus monkeys [26]. Some studies show the maximal cone peak at the beginning of the day, both in nocturnal species such as cats [25] and diurnal species such as Tupai [29] and Arvicanthis ansorgei [17,30]. Since mammals contain, in the great majority, mid/long wavelength cones [4,31], these studies have not provided information on the numerically minor S cones. Indeed, before the present investigation, it was not even known whether S cones undergo phagocytic turnover. The Nrl\(^{-/-}\) mouse thus represents a unique opportunity to analyze this population. It should be stated that cone identity in such mutant mice may not be strictly normal, and that the retina of this strain undergoes partial degeneration at an early age [14]. However, by a range of structural and functional criteria, they appear to be true cones [14,15].

Our results show that in Nrl\(^{-/-}\) mice, rhythmic phagocytosis occurs under normal cyclic conditions, where it may be regulated by light and/or an endogenous circadian clock. Under conditions of constant darkness, rod shedding continues with a profile similar to that of cyclic light [22,23,30,32]. Data for cone phagocytosis under constant darkness are scarce. In the cone-dominated retinas of fence lizards, rhythmic shedding continued when animals were placed in prolonged darkness, and the amplitude of shedding was greatly reduced [27], as seen in the present study. In our own studies on the diurnal rodent, Arvicanthis ansorgei, cone shedding also continued with the same profile in DD as in LD, but overall activity was actually enhanced by about 50% [30]. In this latter case, only mid/long wavelength cones were studied, and so the contrasting results may have reflected the altered responses of the two spectral types. But rod shedding also shows variable responses to prolonged darkness in terms of amplitude, with some studies showing increased activity [30,33] and others showing decreased activity [32]. Regardless, the continued cyclic shedding during constant darkness demonstrates the functioning of an endogenous circadian clock, capable of generating rhythmic behavior in the absence of light cues. This clock is able to synchronize peak shedding to the approximate beginning of day, at least after a single 24 h period of complete darkness.

Some observations on S-cone shedding as a function of lighting conditions deserve comment. According to our observations in wild-type mice, rats, and Arvicanthis ansorgei, and using previously published data from cats [25] and chickens [21], the relative maximal rate of phagocytosis is overall quite similar between rods and cones, and between species. These values are approximate, given that our data collection points may not have coincided precisely with actual peak activity, that we did not take into account differences in cell body or OS size, and that actual turnover rates for rods and cones are not known for all these species. But given the reduced cell density of ~80%, compared to the wild type [14], the length of OS [14] and RPE microvilli (~5–7 μm in wild-type: [20]; ~3 μm in Nrl\(^{-/-}\): [14]) in Nrl\(^{-/-}\) retinas, it is perhaps surprising that the phagocytic amplitude is so high (comparable to that of rods in normal mice). This is different from Arvicanthis, in which there is a clear difference in the turnover rates of rods and cones. Cone phagocytosis cannot be quantified in wild-type mouse retinas, because of the low cone numbers (3%–4%) [12] and diffuse distribution. Furthermore, rods are absent from the Nrl\(^{-/-}\) retina, so we are unable to compare data between the two strains. For the moment, we can only speculate that phagocytosis rates are perhaps related to relatively high turnover in S cones, or to a particular feature of the mutant retina (e.g., ongoing degeneration). Finally, the variability between species in terms of daily cone phagocytosis profiles, with in some cases an evening peak [21,24,26-28], and in others a morning peak overlapping with rods [25,29,30], makes it difficult to predict what the corresponding temporal cone profile would be in normal mice. The question is important, since two theoretical scenarios are possible: (1) normal mice have an evening cone peak (similar to most species studied), in which case the developmental switch from rods to cones in the Nrl\(^{-/-}\) mouse leaves the S cone-like cells with a partial functional rod phenotype; or (2) normal mice also have a morning cone peak, and the Nrl\(^{-/-}\) strain has a more fully cone-like functional phenotype.

Although the level of expression is reduced compared to wild-type mouse eyes, MerTK is clearly present at the apical RPE surface in Nrl\(^{-/-}\) mice, where it presumably participates in the binding and uptake of shed cone membranes, as in rats [7] and wild-type mice [34]. The distribution of the staining patterns for MerTK and S-cone opsin, showing shortened microvilli and OS respectively, suggests that RPE/OS interactions must be less intimate than in wild-type retinas.

In summary, the Nrl\(^{-/-}\) mouse presents a unique opportunity to examine cone shedding (moreover, blue cone
sheding) in the otherwise highly rod-dominant mouse retina, and represents a valuable model to investigate the molecular mechanisms involved in cone recycling.

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REFERENCES

1. Rodieck RW, Rushton WA. Cancellation of rod signals by cones, and cone signals by rods in the cat retina. J Physiol 1976; 254:77-85. [PMID: 1255506]

2. Hestrin S, Korenbrot JI. Activation kinetics of retinal cones and rods: response to intense flashes of light. J Neurosci 1990; 10:1967-73. [PMID: 2355261]

3. Szél A, Diamantstein T, Rohlich P. Identification of the blue-sensitive cones in the mammalian retina by anti-visual pigment antibody. J Comp Neurol 1988; 273:593-602. [PMID: 3209737]

4. Jacobs GH. Evolution of colour vision in mammals. Philos Trans R Soc Lond B Biol Sci 2009; 364:2957-67. [PMID: 19720656]

5. Steinberg RH, Fisher SK, Anderson DH. Disc morphogenesis in vertebrate photoreceptors. J Comp Neurol 1980; 190:501-8. [PMID: 6771304]

6. Young RW, Bok D. Participation of the retinal pigment epithelium in rod outer segment renewal process. J Cell Biol 1969; 42:392-403. [PMID: 5792328]

7. D'Cruz PM, Yasumura D, Weir J, Matthes MT, Abderrahim H, LaVail MM, Vollrath D. Mutation of the receptor tyrosine kinase gene Mertk in the retinal dystrophic RCS rat. Hum Mol Genet 2000; 9:645-51. [PMID: 10699188]

8. Finnemann SC, Bonilha VL, Marmorstein AD, Rodriguez-Boulan E. Phagocytosis of rod outer segments by retinal pigment epithelial cells requires alphavbeta5 integrin for binding but not for internalization. Proc Natl Acad Sci USA 1997; 94:12932-7. [PMID: 9371778]

9. Sparrow JR, Hicks D, Hamel CP. The retinal pigment epithelium in health and disease. Curr Mol Med. [PMID: 21091424] In press

10. Nandrot EF, Kim Y, Brodie SE, Huang X, Sheppard D, Finnemann SC. Loss of synchronized retinal phagocytosis and age-related blindness in mice lacking alphavbeta5 integrin. J Exp Med 2004; 200:1539-45. [PMID: 15596525]

11. Gal A, Li Y, Thompson DA, Weir J, Orth U, Jacobson SG, Apfelstedt-Sylla E, Vollrath D. Mutations in the receptor tyrosine kinase gene Mertk in the retinal dystrophic RCS rat. Hum Mol Genet 2000; 9:645-51. [PMID: 10699188]

12. Jeon CJ, Strettoi E, Masland RH. The major cell populations of the mouse retina. J Neurosci 1998; 18:8936-46. [PMID: 9786999]

13. Szél A, Röhlisch P. Two cone types of rat retina detected by anti-visual pigment antibodies. Exp Eye Res 1992; 55:47-52. [PMID: 1397129]

14. Mears AJ, Kondo M, Swain PK, Takada Y, Bush RA, Saunders TL, Sieving PA, Swaroop A. Nrl is required for rod photoreceptor development. Nat Genet 2001; 29:447-52. [PMID: 11694879]

15. Daniele LL, Lillo C, Lyubarsky AL, Nikonov SS, Philp N, Mears AJ, Swaroop A, Williams DS, Pugh EN Jr. Cone-like morphological, molecular, and electrophysiological features of the photoreceptors of the Nrl knockout mouse. Invest Ophthalmol Vis Sci 2005; 46:2156-67. [PMID: 15914637]

16. Yu J, He S, Friedman JS, Akimoto M, Ghosh D, Mears AJ, Hicks D, Swaroop A. Altered expression of genes of the Bmp/Smad and Wnt/calcium signaling pathways in the cone-only Nrl−/− mouse retina, revealed by gene profiling using custom cDNA microarrays. J Biol Chem 2004; 279:42211-20. [PMID: 15292180]

17. Bobu C, Craft CM, Masson-Petev M, Hicks D. Photoreceptor organization and rhythmical phagocytosis in the nile rat Arvicanthis ansorgei: a novel diurnal rodent model for the study of cone pathophysiology. Invest Ophthalmol Vis Sci 2006; 47:3109-18. [PMID: 16799057]

18. Zhu X, Brown B, Li A, Mears AJ, Swaroop A, Craft CM. GRK1-dependent phosphorylation of S and M opsins and their binding to cone arrestin during cone phototransduction in the mouse retina. J Neurosci 2003; 23:6152-60. [PMID: 12853434]

19. Hicks D, Molday RS. Differential immunogold-dextran labeling of bovine and frog rod and cone cells using monoclonal antibodies against bovine rhodopsin. Exp Eye Res 1986; 42:55-71. [PMID: 2420630]

20. Prasad D, Rothlin CV, Burrola P, Burstyn-Cohen T, Lu Q, Garcia de Frutos P, Lemke G. TAM receptor function in the retinal pigment epithelium. Mol Cell Neurosci 2006; 33:96-108. [PMID: 16901715]

21. Young RW. The daily rhythm of shedding and degradation of rod and cone outer segment membranes in the chick retina. Invest Ophthalmol Vis Sci 1978; 17:105-16. [PMID: 624604]

22. Besharse JC, Hollyfield JG. Turnover of mouse photoreceptor outer segments in constant light and darkness. Invest Ophthalmol Vis Sci 1979; 18:1019-24. [PMID: 478775]

23. LaVail MM. Rod outer segment disk shedding in rat retina: relationship to cyclic lighting. Science 1976; 194:1071-4. [PMID: 982063]

24. Long KO, Fisher SK, Fariss RN, Anderson DH. Disc shedding and autophagy in the cone-dominant ground squirrel retina. Exp Eye Res 1986; 43:193-205. [PMID: 3758219]

25. Fisher SK, Pfeffer BA, Anderson DH. Both rod and cone disc shedding are related to light onset in the cat. Invest Ophthalmol Vis Sci 1983; 24:844-56. [PMID: 6683265]

26. Anderson DH, Fisher SK, Erickson PA, Tabor GA. Rod and cone disc shedding in the rhesus monkey retina: a quantitative study. Exp Eye Res 1980; 30:559-74. [PMID: 7409012]

27. Bernstein SA, Breeding DJ, Fisher SK. The influence of light on disc shedding in the lizard, Sceloporus occidentalis. J Cell Biol 1984; 99:379-89. [PMID: 6746734]

28. Tabor GA, Fisher SK, Anderson DH. Rod and cone disc shedding in light-entrained tree squirrels. Exp Eye Res 1980; 30:545-57. [PMID: 7409011]

29. Immel JH, Fisher SK. Cone photoreceptor shedding in the tree shrew (Tupaia belangerii). Cell Tissue Res 1985; 239:667-75. [PMID: 3986885]

30. Bobu C, Hicks D. Regulation of retinal photoreceptor phagocytosis in a diurnal mammal by circadian clocks and ambient lighting. Invest Ophthalmol Vis Sci 2009; 50:3495-502. [PMID: 19234351]
31. Peichl L. Diversity of mammalian photoreceptor properties: adaptations to habitat and lifestyle? Anat Rec A Discov Mol Cell Evol Biol 2005; 287:1001-12. [PMID: 16200646]
32. Goldman AI, Teirstein PS, O'Brien PJ. The role of ambient lighting in circadian disc shedding in the rod outer segment of the rat retina. Invest Ophthalmol Vis Sci 1980; 19:1257-67. [PMID: 7429762]
33. Terman JS, Remé CE, Terman M. Rod outer segment disk shedding in rats with lesions of the suprachiasmatic nucleus. Brain Res 1993; 605:256-64. [PMID: 8481775]
34. Duncan JL, LaVail MM, Yasumura D, Matthes MT, Yang H, Trautmann N, Chappelow AV, Feng W, Earp HS, Matsushima GK, Vollrath D. An RCS-like retinal dystrophy phenotype in mer knockout mice. Invest Ophthalmol Vis Sci 2003; 44:826-38. [PMID: 12556419]