Functional and Anatomical Variations in Retinorecipient Brain Areas in *Arvicanthis niloticus* and *Rattus norvegicus*: Implications for the Circadian and Masking Systems

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Functional and anatomical variations in retinorecipient brain areas in *Arvicanthis niloticus* and *Rattus norvegicus*: Implications for the circadian and masking systems

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Functional and anatomical variations in retinorecipient brain areas in *Arvicanthis niloticus* and *Rattus norvegicus*: Implications for the circadian and masking systems

Daily rhythms in light exposure influence the expression of behaviour by entraining circadian rhythms and through its acute effects on behaviour (i.e., masking). Importantly, these effects of light are dependent on the temporal niche of the organism; for diurnal organisms, light increases activity, whereas for nocturnal organisms, the opposite is true. Here we examined the functional and morphological differences between diurnal and nocturnal rodents in retinorecipient brain regions using Nile grass rats (*Arvicanthis niloticus*) and Sprague-Dawley (SD) rats (*Rattus norvegicus*), respectively. We established the presence of circadian rhythmicity in cFOS activation in retinorecipient brain regions in nocturnal and diurnal rodents housed in constant dark conditions to highlight different patterns between the temporal niches. We then assessed masking effects by comparing cFOS activation in constant darkness (DD) to that in a 12:12 light/dark (LD) cycle, confirming light responsiveness of these regions during times when masking occurs in nature. The intergeniculate leaflet (IGL) and olivary pretectal nucleus (OPN) exhibited significant variation among time points in DD of both species, but their expression profiles were not identical, as SD rats had very low expression levels for most timepoints. Light presentation in LD conditions induced clear rhythms in the IGL of SD rats but eliminated them in grass rats. Additionally, grass rats were the only species to demonstrate daily rhythms in LD for the habenula and showed a strong response to light in the superior colliculus. Structurally, we also analysed the volumes of the visual brain regions using anatomical MRI, and we observed a significant increase in the relative size of several visual regions within diurnal grass rats, including the lateral geniculate nucleus, superior colliculus, and optic tract. Altogether, our results suggest that diurnal grass rats devote greater proportions of brain volume to visual regions than nocturnal rodents, and cFOS activation in these brain regions is dependent on temporal niche and lighting conditions.

Keywords: diurnality, visual system, circadian, light exposure, magnetic resonance imaging
Introduction

Temporal niche refers to the time-of-day during the light-dark cycle in which an organism is active and is defined by the unique levels of ambient light and temperature of the environment present (Kronfeld-Schor & Dayan, 2008). These variations in ambient light and temperature can produce a range of activity patterns, with the two extremes restricting activity primarily to either the day (diurnal; Fogo et al., 2018) or night (nocturnal; Refinetti, 2006). Circadian rhythms and acute responses to light (i.e., masking) have uniquely adapted to regulate activity to these two temporal niches. Mechanisms underlying temporal niche preference are hypothesized to lie downstream of or independent from the suprachiasmatic nucleus (SCN, mammalian biological clock), as rhythms within the SCN are virtually identical in diurnal and nocturnal species (Smale, Nunez & Schwartz, 2008), even though behaviours occur at opposite times of day. With respect to masking patterns, light increases activity in diurnal species, whereas light suppresses activity in nocturnal species (Shuboni et al., 2012). Light is one of the most important and powerful entraining stimuli (i.e., Zeitgeber), and in mammals, light information is solely communicated to the brain via the visual system (Ibuka, Inouye, & Kawamura, 1977; Nelson & Zucker, 1981). Therefore, it is important that we understand how light affects the brain of both diurnal and nocturnal organisms.

Structurally, the visual systems of diurnal and nocturnal animals have evolved differently due to evolutionary pressure to accommodate the challenges of each temporal niche (Ankel-Simons & Rasmussen, 2008). As compared to diurnal animals, nocturnal animals have developed larger eyes (Garamszegi et al., 2001; Kirk, 2006), their retinas contain fewer cone receptors and more rod receptors (Ahnelt and Kolb, 2000; Peichl et al., 2000; Peichl, 2005; Solovei et al., 2009), and their optic nerves are significantly smaller (Stephan et al., 1984; Kirk & Kay, 2004). All of these characteristics allow nocturnal animals to adapt adequate vision for environments with
low light levels. Diurnal animals, on the other hand, have evolved vision for high light levels and have retinas rich in cone receptors (Gaillard et al., 2008; Peichl, 2005). In the Nile grass rat (*Arvicanthis niloticus*), a rodent model that is predominantly diurnal in the field and in the laboratory (Blanchong et al., 1999; Blanchong & Smale, 2000), the retina contains ten times the number of cones compared to nocturnal mice and rats (Gaillard et al., 2008) and electroretinograms show several visual acuity features that more closely resemble human retinal physiology (Gilmour et al., 2008). Grass rats also differentiate from their nocturnal counterparts in the projections of intrinsically photosensitive retinal ganglion cells (ipRGC), the photoreceptor responsible for circadian entrainment and masking. Specifically, as compared to nocturnal rodents, grass rats exhibit significantly less innervation from ipRGCs in the lateral geniculate nucleus and the olivary pretectal nucleus, two brain regions critical for vision (Langel et al., 2015). Grass rats, therefore, are an ideal model to study the relationship between the visual system and both circadian rhythms and masking.

Grass rats are a murid rodent indigenous to the grasslands of Kenya (Delany & Monro, 1986). In the laboratory, they are predominantly diurnal as measured by general activity patterns and body temperature (Katona & Smale, 1997; McElhinny, Smale, & Holekamp, 1997) and are considered a reliable diurnal rodent model (Refinetti, 2006) under these conditions; significant variations in chronotype only exist when grass rats are given a running wheel (Blanchong et al., 1999). Circadian and daily rhythms in cFOS activation within grass rat brain have been compared to nocturnal counterparts in many subregions, including within the circadian system (Nunez et al., 1999; Mahoney, Bult & Smale, 2001; Schwartz, Nunez & Smale, 2004), regions associated with sleep and wakefulness (Martínez, Smale & Nunez, 2002; Novak, Smale & Nunez, 2000; Nixon & Smale, 2004; Schwartz & Smale, 2005) and regions associated with reward
(Castillo-Ruiz et al., 2010). However, these studies have not examined all retinorecipient regions, and it has been recently shown that some of these brain regions exhibit distinct neuronal activation in response to light that are unique from nocturnal rodents (Gall et al., 2014; Langel et al., 2014; Shuboni et al., 2015). Previous work has brought attention to two structures, the geniculate complex and the olivary pretectal nucleus, as possible components of the masking neural mechanism. Lesions of both areas alter the behaviours of the grass rat, both in masking response to light and circadian/daily rhythms in activity (Gall et al., 2013, 2014, 2017). We do not, however, yet know the circadian patterns of neuronal activity within these regions nor how daily light exposure masks these rhythms under normal lighting conditions. Many of these retinorecipient brain regions have been shown to contribute to circadian rhythm regulation in nocturnal species (reviewed in Morin & Allen, 2006), but we do not yet know the pattern of expression in diurnal species, such as grass rats, or how FOS expression changes when animals are exposed to light during the day. Examining how light affects these brain regions differently in diurnal and nocturnal rodents under normal lighting conditions will allow us to better understand the different mechanisms by which these brain regions strengthen daily rhythms in behavior between temporal niches.

Here we examined the functional and structural differences of the visual system between the diurnal grass rat and the nocturnal Sprague Dawley rat. The expression of the immediate early protein, cFOS, was used to examine neuronal activation within visual regions of the brain. Rhythms in cFOS expression under constant darkness provided insight into circadian rhythmicity within the brain regions. We then compared these rhythms to 12:12 light/dark conditions to measure the impact of light on the rhythmic expression of neuronal activation across the day and directly at the two
timepoints where light is presented, which highlighted how nocturnal and diurnal rodents mask to the presentation of light under natural daily light exposure. Finally, to understand the structural differences of the visual system between temporal niches, we used high resolution magnetic resonance imaging (MRI) to measure the three-dimensional volumes of visual regions and compared the differences in relative size. Both the functional and anatomical studies demonstrated the differences between temporal niches.

**Methods**

**Animals**

All experiments were approved by the Institutional Animal Care and Use Committee (IACUC) at Michigan State University and are in accordance with the NIH Guide for the Care and Use of Laboratory Animals. A total of seventy-six adult male diurnal grass rats (breeding colony, Michigan State University) and sixty-three adult male nocturnal SD rats (Charles River Laboratories, Wilmington, MA) were used in the study. All animals were housed in standard 12:12 light/dark conditions prior to experimentation and were provided food and water ad libitum throughout the duration of the study. Two experiments were conducted to compare the visual systems of the diurnal grass rat and nocturnal SD rat. In Experiment 1, we examined the circadian rhythms of brain activation, using cFOS, within several retinorecipient brain regions from slides produced for a previous publication (Schwartz et al., 2004). Tissue was collected from both species at six timepoints (Zeitgeber time (ZT)1, 5, 13, 17, 20, and 23; grass rat: n=35; SD rat: n=29) in 12:12 light/dark (LD) conditions and at six timepoints in constant darkness (DD; Circadian time (CT)1, 5, 13, 17, 20, and 23; grass rat: n=35; SD rat: n=30). For the 65 animals sacrificed in DD, they were initially housed in 12:12 LD
using cage-top infrared motion detectors to detect general activity patterns for 1-2 weeks, and then they were placed in DD for 21-22 days for grass rats, or 16-17 days for SD rats. At the end of this time frame in DD, activity data were visualized using actograms, and onsets were eye-fitted independently by two investigators. Perfusion times were randomly assigned, with CT0 indicating activity onset for grass rats, and CT12 indicating activity onset for SD rats (see Schwartz et al., 2004). In Experiment 2, we examined the total volume of several retinorecipient brain areas using ex vivo high-resolution magnetic resonance imaging (MRI; grass rat: n=6; SD rat: n=4; see Ex vivo MRI Protocol below for details). All animals in Experiment 2 were males, singly-housed, and between 6 months and 1 year old when perfused. All perfusions for Experiment 2 occurred during the lights-on phase, between ZT4 and ZT8.

**Immunohistological Procedure**

Tissue was collected and stained as previously described in Schwartz et al. (2004). Briefly, animals were transcardially perfused using 0.01 M PBS followed by 4% paraformaldehyde. Brains were removed and post-fixed for 1-2 h and then transferred to a 20% sucrose solution for at least 48 h until sectioning. Coronal sections were cut on a freezing microtome at 30 µm and stored in cryoprotectant at -20 °C until further processing.

Free floating sections were rinsed three times in PBS for 10 min then blocked for 1 h in 5% normal goat serum (Vector Laboratories, Burlingame, CA). Tissue was incubated in the primary antibody, rabbit anti-cFOS (1:25,000; Santa Cruz Biochemistry, Santa Cruz, CA), for 48 h at 4 °C then moved into the secondary antibody, biotinylated goat anti-rabbit (1:200; Vector Laboratories), for 1 h at room temperature. An avidin-biotin peroxidase complex kit (ABC Vectastain Kit; Vector...
Laboratories) was used before visualizing the protein with 0.5 mg/ml diaminobenzidine (DAB; Sigma, St. Louis, MO). Sections were washed three times with PBS and then mounted on gelatinized slides. Slides were dehydrated with increasing concentrations of alcohol and xylenes, then coverslipped with Permount (Fisher Scientific, Hampton, NH).

**cFOS rhythms in the Visual System**

Images were acquired with a Zeiss light microscope (Zeiss, Gottingen, Germany) equipped with a digital camera (CX900, MBF Bioscience, Williston, VT). All images were combined into one composition file using Adobe Photoshop 7 (Adobe Systems, Mountain View, CA). Regions of interest were identified within the images, either by using counting boxes (superior colliculus (SC), 300 µm x 400 µm) or were outlined by a trained researcher blind to condition; olivary pretectal nucleus (OPN), lateral habenula (LHb), medial habenula (MHb), dorsal lateral geniculate nucleus (dLGN), intergeniculate nucleus (IGL), and ventral lateral geniculate nucleus (vLGN) (see Figure 1 for outlined brain areas). The number of cells positive for cFOS (cFOS+) was counted bilaterally for one section using the Particle Analysis tool and thresholding in the ImageJ Program (NIH, Bethesda, MD). Thresholding was performed by converting the photomicrograph into a monochrome image, and the threshold was set manually using the slider bars until the maximum number of cells containing cFOS were pixelated. Finally, the number of cFOS+ cells were counted automatically using ImageJ.

**Ex vivo MRI Protocol**

Perfused whole brains were transferred into a 15mL tube filled with Fombilin® Y (Sigma-Aldrich, St Louis, MO), a perfluorocarbon solution that produces no MRI signal and is the “gold standard” for *ex vivo* imaging biological samples. Tubes were secured
to a 2x2 surface receive array within a volume transmit coil. Images were acquired with a 7T Bruker Biospec 70/30 USR using a 2D T2-weighted TurboRARE sequence (TE:33 ms, TR:2654.1 ms, Rare Factor:8, 50 μm x 50 μm) for 1 h 1 min (30 repetitions) with 10 slices for grass rats and 12 slices (1 mm thickness) for SD rats.

One brain from each species was soaked for 48hr in a PBS solution doped with 0.1M gadopentetate dimeglumine (Gd-DTPA, Magnevist®). These brains were transferred, prior to scanning, to fresh PBS in a 15 mL tube which was cut to fit into a home-made 17 mm solenoid coil. Images were also acquired with a 7T Bruker Biospec 70/30 USR using 3D T1/T2-weighted Flash sequence (TE:10.44 ms, TR:31 ms, 100 μm x 100 μm x 100 μm) for 47 min (2 repetitions).

**Volume Analysis of Visual System**

Volume analysis for the high resolution T2-weighted images was conducted using the Measure tool in the ImageJ Program (NIH, Bethesda, MD). Three components of the visual system were clearly visible with MRI: SC, lateral geniculate nucleus (LGN), Habenula (EPI), and optic tract (opt). Two non-visual areas were also outlined as controls, the cortex (CTX) and hippocampus (HPC). The whole brain was then outlined to correct for total volume differences between the two species. A percentage was calculated by dividing the volume of each area by the whole brain volume and then multiplying by 100. Prior to statistical analysis, percentage data was arcsine transformed. The 3D MRI were used to create 3D volume rendering of the regions of interest using 3D Slicer 4.8 ([https://www.slicer.org/](https://www.slicer.org/); Fedorov et al., 2012).

**Statistical Analyses**

The histological data was compared separately for lighting condition (LD or DD) and for species (grass rats or SD rats) using one-way ANOVAs with time of day as the
independent variable and the number of cFOS+ cells as the dependent variable. For each region of interest, the number of cFOS+ cells were compared across time followed by post hoc analysis using t-tests (Tukey HSD). The presence of 24-h rhythms was detected using cosinor analysis, with data represented by the following function: $x_i = M + A \cos \left(\frac{2 \pi ZT}{24}\right)$, where $M$ denotes MESOR and $A$ denotes amplitude of the oscillation. Linear regression by method of least squares was used to test for rhythmicity, and the probability that $A$ is significantly different from zero was calculated using an F-test with 2 and $N$-3 degrees of freedom (Cornelissen, 2014; Nelson et al., 1979; Tong, 1976; Refinetti, Lissen, & Halberg, 2007). Additionally, within species CT and ZT were compared at two timepoints, 1 and 5, to examine the direct effects of light on cFOS using a two-way ANOVA. For the MRI volume data, independent samples t-tests were used to compare the size of each region between grass rats and SD rats. All analyses were performed with SPSS Statistic 23 software (IBM Corp., Armonk, NY) and significance for all tests was $p<0.05$.

**Results**

*Circadian Rhythms of cFOS in Constant Dark Conditions*

We examined significant variation among time points of neuronal activity within retinorecipient regions using cFOS activation across time in constant dark conditions, including the geniculate complex (e.g., dLGN, IGL, vLGN), OPN, habenula (e.g., MHb and LHb), and SC. Figure 1 presents photomicrographs of cFOS in the geniculate complex, OPN, and habenula (EPI); photomicrographs of cFOS in the SC can be found in Supplemental Figure 1. Grass rats exhibited significant variation among time points in cFOS protein expression within the OPN, $F(5,31)=3.840$, $p=0.011$, with post-hocs revealing higher levels of activation at CT5 and CT23 (Figure 2, left panel). When the
OPN was analyzed using cosinor analyses, however, a significant circadian rhythm was not detected (Supplemental Table 2). Within the geniculate complex, grass rats had two regions that trended toward significance from the ANOVA; the IGL, $F(5,31)=2.604, p=0.051$, and the vLGN, $F(5,31)=2.364, p=0.068$. The IGL cFOS expression was highest at CT17 and CT20 during the beginning of subjective night. Of these two brain regions, only the IGL exhibited a significant circadian rhythm following cosinor analyses (Supplemental Table 2). cFOS activation for the dLGN did not express a significant variation among time points from the ANOVA ($F(5,32)=1.209, p=0.421$) or a circadian rhythm from cosinor analyses. Neither the SC ($F(5,31)=0.604, p=0.697$) nor either portion of the grass rat habenula (LHb, $F(5,31)=0.789, p=0.566$; MHb, $F(5,31)=1.904, p=0.126$) exhibited a significant variation among time points in cFOS activation from the ANOVA. Cosinor analyses supported these findings, as a significant circadian rhythm was not detected in the SC or habenula of grass rats.

In the SD rats, there was a significant effect of time on cFOS activation in the IGL, $F(5,28)=3.023, p=0.031$, however the pattern of activation was low and exhibited the lowest cFOS expression at CT20 (Figure 2, right panel), and was not significant following cosinor analyses. The OPN of the SD rat also expressed a significant variation among time points in constant conditions ($F(5,27)=3.807, p=0.012$), but again the levels of cFOS were low with only CT17 having a heightened level of activation. Again, cosinor analyses did not reveal a significant circadian rhythm in the OPN (Supplemental Table 2). No rhythms were expressed within the dLGN ($F(5,28)=1.221, p=0.331$), vLGN ($F(5,28)=2.104, p=0.101$), SC ($F(5,28)=1.106, p=0.392$), LHb ($F(5,28)=0.991, p=0.445$), or MHb ($F(5,28)=0.855, p=0.526$), according to the ANOVA or cosinor analyses (Supplemental Table 2).
**Daily Rhythms of cFOS in 12:12 Light/Dark Conditions**

We examined the daily rhythms of neuronal activity within same retinorecipient regions using cFOS activation across time in a 12:12 light/dark cycle (Figure 3 for geniculate complex, OPN, & EPI; Supplemental Figure 1 for SC). Remarkably, in the grass rat, all regions that exhibited significant circadian rhythms in cFOS expression in constant darkness became arrhythmic in 12:12 LD conditions (IGL, $F(5,33)=0.711$, $p=0.621$; vLGN, $F(5,33)=0.563$, $p=0.727$; OPN, $F(5,34)=1.542$, $p=0.210$; Figure 4, left panel), because cFOS activation was heightened in the light (see Figure 5). However, both regions of the habenula expressed significant variation among time points in cFOS expression in LD conditions (LHb, $F(5,33)=4.633$, $p=0.003$; MHb, $F(5,34)=3.018$, $p=0.026$), and cosinor analyses revealed that these variations fit a sinusoidal wave. The SC ($F(5,33)=1.702$, $p=0.167$) remained arrhythmic as it was in DD.

In SD rats, two regions of the geniculate complex expressed significant variations among time points in cFOS activity in 12:12 LD conditions: the dLGN ($F(5,29)=4.409$, $p=0.005$) and the IGL ($F(5,29)=9.974$, $p<0.001$). However, cFOS expression patterns were different between these two regions. In the IGL, post-hocs revealed that levels were significantly higher during the active period and the beginning of rest period, while in the dLGN, two peaks occurred at ZT1 and ZT17. Cosinor analyses revealed significant rhythms only in the IGL (Supplemental Table 2). The pattern of expression within the vLGN, which was not significant $F(5,29)=1.212$, $p=0.334$, more closely resembled the dLGN. The OPN of the SD rat did not exhibit a significant daily rhythm in cFOS activation, $F(5,29)=1.008$, $p=0.436$ (Figure 4, right panel). No other regions expressed significant daily rhythms in cFOS activation (SC, $F(5,29)=1.860$, $p=0.143$; LHb, $F(5,29)=0.548$, $p=0.738$; MHb, $F(5,29)=0.254$, $p=0.934$).

We compared cFOS levels between ZT and CT groups at the two timepoints
during the day, ZT/CT1 & 5, to examine the direct effect of light on the cFOS expression (Figure 5 and Supplemental Table 1). In grass rats, light during the day in ZT conditions increased cFOS levels with main effects of lighting condition for the SC (F(1,20)=17.61, p<0.001), dLGN (F(1,20)=6.46, p=0.019) and a trend toward significance in the IGL (F(1,20)=4.22, p=0.053). The OPN had a significant interaction (F(1,20)=7.673, p=0.012), when the time points were compared between CT and ZT there was a 5.6x increase in expression one hour after lights on in ZT conditions when compared to CT, t(10)=2.621, p=0.026. SD rats had a similar directional pattern with dramatic increases in cFOS expression with main effects of light exposure in three regions: dLGN (F(1,16)=10.84, p=0.005), IGL (F(1,16)=54.57, p<0.001) and OPN (F(1,15)=17.23, p=0.001). The OPN had a 5.2X increase in expression at ZT1 and a 7.9X increase at ZT5. The SC for the SD rat did not have significant increase in cFOS expression as a ME of lighting F(1,15)=0.09, p=0.770 or an interaction (F(1,15)=2.27, p=0.153). Altogether, grass rats exhibited significant increases in light-induced FOS in the dLGN, IGL, OPN, and SC at ZT1, whereas SD rats exhibited significant increases at the same time point in only the dLGN, IGL, and OPN. We observed similar effects at ZT5, except for the OPN, which no longer exhibited a significant increase in light-induced FOS in grass rats. These results demonstrate that grass rats and SD rats express differential activation to the presentation of light within our nuclei of interest.

**MRI Volume Analysis**

We assessed the high resolution 2D images taken across the brains of grass rat and SD rat at 7 levels (Figure 6). The four components of the visual system (SC, LGN, opt, and EPI) are highlighted in representative sections (Figure 7). Clear size differences between the two species are visible in overall size and proportion of most of the visual
areas. Quantification of total brain volume of grass rats (526.2±6.7mm³) showed a significantly smaller brain than SD rats (1198.7±13.7mm³; t(8)=48.672, p<0.001), with a 56.1% decrease in total volume. Both control regions assessed did not differ in total percent volume between grass rats and SD rats, HPC (t(8)=0.162, p=0.875) and CTX (t(8)=1.087, p=0.309). There were significant differences in percent volume of 3 visual regions examined, LGN (t(8)=12.503, p<0.001), SC (t(8)=5.610, p<0.001) and opt (t(8)=7.411, p<0.001), but not in the EPI (t(8)=1.163, p=0.278; Figure 7). Within the LGN, SC, and opt grass rats had larger percent volume when compared to SD rats, with an increase of 47.0%, 47.4%, 48.8% respectively. When these regions were visualized in 3D MRI scans, the dramatic increase in size within the grass rats when compared with the SD rat can be clearly observed from coronal, sagittal and horizontal sections (Figure 8).

Discussion

Circadian rhythms and masking work synergistically in the presence of light (12:12 LD cycle) to define the daily expression of activity rhythms (Aschoff, 1999). There are clear differences within the neural mechanisms that drive these behaviours between species that occupy different temporal niches (Yan, Smale, & Nunez, 2018). The present experiments demonstrate the differences within the visual system between diurnal and nocturnal rodent species in both (1) circadian and masking responses in neuronal activation and (2) the morphology of the structural components.

The IGL has been shown to play a crucial role in defining masking behaviour between the temporal niches (Gall et al., 2013). Specifically, in grass rats, lesions of the IGL alter the direction of masking behaviour in response to the presentation of light from positive to negative, therefore causing the diurnal rodent to respond to light similar
to a nocturnal species by suppressing activity. Understanding the activation patterns based on time of day will provide insight into how the region works to promote diurnal behaviour through masking and circadian rhythms in natural 12:12 LD conditions by demonstrating the natural activity in DD and how light alters responses in LD. We found here that while grass rats showed a trend toward significant variation among time points of cFOS activation in the IGL under constant dark conditions and significant circadian rhythms following cosinor analyses, SD rats did not. For grass rats, higher activation was observed during the night (CT17 & CT20). The magnitude of cFOS activation was generally low in SD rats and did not have a clear rhythmic pattern. Although cFOS expression in the IGL was low overall, cosinor analyses did detect a rhythm in grass rats. In addition, these results are similar to the number of light induced FOS cells in the IGL of grass rats in previous reports (Shuboni et al., 2015). In 12:12 LD conditions, we found no daily rhythm in cFOS expression broadly within the IGL for grass rats, whereas SD rats did exhibit a significant rhythm in LD. Increased cFOS activation specifically during the light phase (Figure 5) masked the rhythm that is normally present in constant darkness. For grass rats, this is similar to previous findings that reported no significant effect of time when comparing the daily rhythms in the region at ZT4 and ZT16 (Smale et al., 2001). However, these authors found a difference in cFOS within NPY+ cells, which project to the SCN, between the day and night, suggesting a conduit for masking effects that merits further examination. In nocturnal rodents, similar patterns of IGL activation have been observed. Edelstein et al. (2000) showed that Wistar rats exhibited markedly low levels of cFOS expression in the IGL under the DD conditions but had clear and significant rhythms in 12:12 LD cycle. These rhythms were attributed to light activation of neurons within the IGL, with low activation in constant darkness and high activation during both LL and the light portion
of the 12:12 LD cycle. Light pulses at night have also been shown to induce increases of cFOS in the IGL in Fisher rats (Caldelas et al., 1998; Prichard et al., 2002) and blind mole-rats (Oelschlager et al., 2000). This increase in cFOS to a light pulse, however, was not observed in CD1 mice at ZT14 (Shuboni et al., 2015). Here we observed lower levels of cFOS during subjective day in constant dark conditions that were activated following the presentation of light in 12:12 LD cycles; this pattern of increased activation in both grass rats and SD rats during the light portion of the LD cycle was also observed in the OPN. Altogether, our results show that the IGL responds differently in 12:12 LD conditions in grass rats as compared to SD rats, suggesting that it plays an important role in temporal niche differences. Indeed, this is supported by lesion data of the IGL in diurnal and nocturnal species, which strongly suggests its involvement in temporal niche differences in masking behaviour (Gall et al., 2013; Redlin et al., 1999; Johnson et al., 1989; Pickard, 1989; Edelstein & Amir, 1999).

The OPN has also been shown to play a major role in the masking behaviour of diurnal grass rats, eliminating the positive masking response to light and causing a positive masking response to darkness post-lesion (Gall et al., 2017). We report here that grass rats and SD rats expressed significant variation among time points in cFOS activation in the OPN under constant dark conditions, with heightened activation in the subjective night; however, in a LD cycle, daily rhythms were eliminated in both species. In both species, this elimination of rhythms was caused by an increase in the level of activation observed during the light phase. We have previously demonstrated that the OPN is responsive to the presentation of light at night in the grass rat (Shuboni et al., 2015) and that when masking behaviour in grass rats was inverted to a nocturnal phenotype with IGL lesioned animals, the OPN’s responsiveness to light was no longer present (Gall et al., 2014). For the SD rat, the OPN had relatively low expression levels
across DD and were more dramatically increased in LD conditions. Other studies in nocturnal rodents, showed that light pulses induce an increase of cFOS in the OPN of albino Fisher F344 rats (Caldelas et al., 1998; Prichard et al., 2002) and blind mole-rats (Oelschlager et al., 2000). Altogether, these results suggest that light induces cFOS expression in both the IGL and OPN of grass rats and SD rats, but there are clear differences between the species.

To better examine the effects of light on cFOS in our data set we compared the two timepoints during light phase, ZT1 and ZT5, to the constant dark, CT1 and CT5. Grass rats have been shown to respond more dramatically to the presentation of light with increases in activity across the 24hr period, while mice only decrease activity at one time (CT14; Shuboni et al., 2012). Here we observed cFOS activation to light in all brain regions examined, except the habenula and vLGN. Interestingly, both components of the habenula in grass rats were the only regions examined to show a daily rhythm in cFOS expression in LD conditions. A study has recently shown a marked difference in the level of GABAergic cells in the LHb between mice and grass rats (Langel et al., 2018) suggesting a possible role for the nuclei in masking behaviour. Similar findings of light inducing the expression of cFOS were observed for SD rats in the entire geniculate complex (e.g., DLG, IGL, OPN), but not for the SC. Grass rats, in stark contrast, had a significant light-induced increase in cFOS activation in the SC. The SC has been shown to be responsive to brief flashes of light and to moving and stationary visual patterns (Craner et al., 1992; Montero and Jian, 1995), but it not yet known how the SC responds to sustained periods of light. Our data show that the SC is more light-responsive in grass rats than in SD rats following prolonged light presentation, suggesting a potential important species-difference in light functionality in this brain region. The clear functional difference found within these regions may have led to
structural adaptations based on temporal niche of the population.

It should be noted that one limitation of our study was only examining two time points during the day, and four time points at night. Because previous work found significant changes in the lower subparaventricular zone in grass rats as compared to SD rats (Nunez et al., 1999), we were more interested in the changes that occurred at night. A future study is needed to more thoroughly examine changes that happen throughout the day, particularly towards the end of the lights-on period.

Evolution between temporal niches has led to the adaptations of many components of the visual system (Ankel-Simons et al., 2008). Since grass rats are more functionally sensitive to the presentation of light, here we also examined the structural differences in the visual system between the niches using MRI. Within primates, there have been many studies comparing variation in the size of eye structures (Kirk, 2004; Kirk, 2006) and visual regions within the brain, particularly the cortex and geniculate (Heesy, Kamilar & Willms, 2011). Studies of adaptation within the rodent visual system between temporal niche switches, have demonstrated alterations in visual cortex (Heimel, Van Hooser, & Nelson, 2005; Campi & Krubitzer, 2010; Campi et al., 2011). We were particularly keen to examine differences between the niches in regions that were associated with masking behaviour. This study confirms that there is alteration in the relative size of optic tract, superior colliculus and geniculate complex between diurnal grass rats and nocturnal SD rats, with larger structures found in grass rats. These findings are in line with a previous histological study in grass rats which showed a larger SC, dLGN, and vLGN (Gaillard et al., 2013). Importantly, we further demonstrate that the input from the optic tract is also larger in grass rats, suggesting that for the diurnal grass rat these regions, (1) receive more visual information from the retina via the optic tract and (2) the devote a greater proportion of the brain to regions
that are critical for driving diurnal behaviour. These anatomical results support our
cFOS data and suggest that visual input into the non-image forming retinorecipient
brain regions in grass rats plays an important role in promoting diurnal behaviour. This
study is also the first to use high resolution MRI to compare the anatomical size of
visual system structures between species of different temporal niches. The technique is
a powerful tool for measuring in three-dimensions regions of interest within the brain
without sacrificing tissue for histology.

It is important to note that there are many ways to assess rhythmicity in cFOS
expression (Refinetti, Lissen, & Halberg, 2007). The most common way that has been
reported in multiple reports is using ANOVAs to detect significant variation among
time points (Caldelas et al., 1998; Prichard et al., 2002). We also analyzed our data
using cosinor analyses, which can be used to detect rhythms that fit a sinusoidal wave.
We included both types of analyses here, and we note some similarities and differences
between the ANOVAs and cosinor analyses. In constant conditions, the only rhythms
detected using cosinor analyses were within the IGL in grass rats; none were significant
in SD rats. In contrast, using ANOVAs, we detected significance in the IGL and OPN
of both species. With respect to LD conditions, rhythms detected using cosinor analyses
included only the IGL in SD rats and the habenula in grass rats; ANOVAs revealed
significance in the IGL and dLGN of SD rats, along with the LHb and MHb of grass
rats. Because rhythms do not always fit a sinusoidal wave, it is important to also include
results from the ANOVA.

**Conclusion**

Grass rats are an optimal model for studying the relationship between the visual
system and both circadian and masking mechanisms in a diurnal organism. Here we
have used this model to examine circadian rhythms in neuronal activation within brain regions that receive direct ipRGC projections and how activation in these regions is altered in the presence of a light/dark cycle. Circadian rhythms of cFOS activation were observed in several retinorecipient brain areas in grass rats and SD rats housed in constant darkness (e.g., geniculate complex, OPN). The expression of cFOS activation in the geniculate complex and OPN was significantly higher at ZT1 and ZT5 as compared to CT1 and CT5, indicating light-induced cFOS expression in these brain areas. The masking response to light, which was defined here as FOS expression induced by a LD cycle that is above and beyond FOS expression in DD, also has similar effects in many regions when comparing between species, including the IGL, dLGN, and OPN, confirming the responsiveness of these regions during times when masking occurs in nature. Interestingly, whereas light-induced cFOS activation in the SC was observed in grass rats, it was not observed in SD rats, suggesting a species-specific difference in light responsiveness in this brain region. Importantly, whereas light increases behaviour in diurnal species such as grass rats, light suppresses behaviour in nocturnal species such as SD rats. Since most retinorecipient brain regions respond similarly in both species, we hypothesize that these brain regions must affect downstream circuitry differently, or that the differences in masking behaviour arises from different visual pathways.

Additionally, we use high resolution MRI for the first time to compare visual system morphology between temporal niches. These findings further confirm the adaptation of structure in the visual system in a diurnal rodent and mirror changes observed between diurnal and nocturnal organisms in avian and primate species (Schmitz & Motani, 2010). We demonstrated that the optic nerve/tract, geniculate complex and the SC all are proportionally larger in grass rats which suggests a possible
driver for the temporal niche specific heightened sensitivity to light in our diurnal species. In summary, we have demonstrated significant functional and morphological differences within the visual system between diurnal and nocturnal rodents.
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Disclosure of Interest

The authors report no conflict of interest.

Data Availability Statement

The data that support the findings of this study are available from the corresponding author, [AJG], upon reasonable request.
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Figure 1. Photomicrographs of the cFOS expression at Circadian Time 5 and 17. In grass rats (A), photomicrographs of the lateral geniculate nucleus (LGN, top row), olivary pretectal nucleus (OPN, middle row), and habenula (EPI, lower row) show stained cFOS+ nuclei in constant dark conditions at Circadian Time (CT)5 and 17. In SD rats (B), photomicrographs of the LGN (top row), OPN (middle row) and habenula (lower row) also show cFOS+ nuclei in constant dark conditions at CT5 and 17.
Quantification of cFOS expression within retinorecipient regions across Circadian Time. The total number of cFOS+ cells was compared across constant darkness for grass rats (left column, black bars) and SD rats (right, grey bars) in several visual structures. In the grass rats, the OPN exhibited significant differences across the day, and the IGL and vLGN were trending toward significance. SD rats had significant circadian rhythms of cFOS expression within the IGL and OPN. Different letters indicate significance with p<0.05.
Figure 3. Photomicrographs of the cFOS expression at Zeitgeber Time 5 and 17. In grass rats (A), photomicrographs of the lateral geniculate nucleus (LGN, top row), olivary pretectal nucleus (OPN, middle row), and habenula (EPI, lower row) show stained cFOS+ cells in 12:12 LD conditions at Zeitgeber Time (ZT)5 and 17. In SD rats (B), photomicrographs of the LGN (top row), OPN (middle row) and habenula (lower row) also show cFOS+ nuclei in 12:12 LD conditions at ZT5 and 17.
Figure 4. Quantification of cFOS expression within retinorecipient regions across Zeitgeber Time. The total number of cFOS+ cells was compared across 12:12 LD conditions for grass rats (left column, black bars) and SD rats (right, grey bars) in several visual structures. In the grass rats, the OPN and IGL no longer express significant daily rhythms, however both the LHb and MHb have daily rhythms under LD conditions. In contrast, SD rats exhibited significant circadian rhythms of cFOS expression within the dLGN and IGL. Different letters indicate significance with p<0.05.
Figure 5. Quantification of masking effects of light between temporal niche. The total number of cFOS+ cells was compared for grass rats (left column) and SD rats (right column) in several visual structures at two timepoints in DD (black bars) and LD (white bars). Grass rats had a significant increase in cFOS+ cells during at least one time point for all regions depicted. SD rats had increases in expression during light for the dLGN, IGL, and OPN. * indicates a significant main effect of lighting with p<0.05, # indicates a main effect trend toward significance with p=0.053, † indicates a significant interaction with p<0.05.
Figure 6. High resolution T2-weighted MRIs across both the grass rat and SD rat brains. Slices include sections at the levels of the VLPO, SCN, habenula (EPI), LGN, OPN, and the SC. Only some structures are clearly discriminated in the MRI and include the EPI, geniculate complex, and SC. Additionally, control regions the hippocampus and cortex are easily identified in both species.
Figure 7. Volume analysis of visual system structures between species. The regions of interest within the visual system were delineated for both species (left columns). The grass rat had significantly larger relative SC, LGN and opt when compared to the SD rat (right column). However, the EPI between the two species were not different from one another. * indicates p<0.05.
Figure 8. Three-dimensional volume rendering of visual regions in grass rat and SD rat from 3D MRI. Volumes for the superior colliculus (green), geniculate complex (red) and the optic nerve/tract (blue) are shown in three orientations. The brains of the two species are represented in identical proportions to demonstrate the relative size differences between visual structures. Grass rats have significantly larger regions for the three visual regions illustrated.