Distinct Retinohypothalamic Innervation Patterns Predict the Developmental Emergence of Species-Typical Circadian Phase Preference in Nocturnal Norway Rats and Diurnal Nile Grass Rats

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ABSTRACT
How does the brain develop differently to support nocturnality in some mammals, but diurnality in others? To answer this question, one might look to the suprachiasmatic nucleus (SCN), which is entrained by light via the retinohypothalamic tract (RHT). However, because the SCN is more active during the day in all mammals studied thus far, it alone cannot determine circadian phase preference. In adult Norway rats (Rattus norvegicus), which are nocturnal, the RHT also projects to the ventral subparaventricular zone (vSPVZ), an adjacent region that expresses an in-phase pattern of SCN-vSPVZ neuronal activity. In contrast, in adult Nile grass rats (Arvicomys niloticus), which are diurnal, an anti-phase pattern of SCN-vSPVZ neuronal activity is expressed. We hypothesized that these species differences result in part from a weak or absent RHT-to-vSPVZ projection in grass rats. Here, using a developmental comparative approach, we assessed species differences in behavior, hypothalamic activity, and RHT anatomy. We report that a robust retina-to-vSPVZ projection develops in Norway rats around the end of the second postnatal week when nocturnal wakefulness and the in-phase pattern of neuronal activity emerge. In grass rats, however, such a projection does not develop and the emergence of the anti-phase pattern during the second postnatal week is accompanied by increased diurnal wakefulness. When considered within the context of previously published reports on RHT projections in a variety of species, the current findings suggest that how and when the retina connects to the hypothalamus differentially shapes brain and behavior to produce animals that occupy opposing temporal niches. J. Comp. Neurol. 000:000–000, 2012.

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INDEXING TERMS: suprachiasmatic nucleus; subparaventricular zone; wakefulness; presynaptic terminals; PACAP; VIP

Mammals exhibit sensory and thermoregulatory adaptations that are thought to have enhanced survival in a nocturnal niche (Crompton et al., 1978; Menaker et al., 1997). These adaptations, along with the fact that most mammalian species today are nocturnal, suggest that nocturnality is a primitive character of Mammalia (Park, 1940; Smale et al., 2003; Campi et al., 2011). Accordingly, diurnality is a derived character that has reemerged independently in some mammalian lineages (Smale et al., 2003; Campi et al., 2011). How the brain develops to support one or the other circadian phase preference is not known.

Circadian rhythms of sleep and wakefulness are regulated by a system that includes the retina, suprachiasmatic nucleus (SCN) of the hypothalamus, and their connection via the retinohypothalamic tract (RHT) (Hannibal, 2002a). All mammals studied thus far—whether diurnal or nocturnal—exhibit a rhythm of increased SCN activity during the day (Fuller et al., 2006; Challet, 2007; Houben et al., 2009). Consequently, interactions of the RHT and SCN with downstream structures are likely to account for species differences in circadian phase preference (Smale,

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et al., 2003). One candidate downstream structure, adjacent to the SCN and also important for the expression of circadian behavior (Lu et al., 2001; Schwartz et al., 2009), is the ventral subparaventricular zone (vSPVZ). In adult Norway rats (Rattus norvegicus), which are nocturnal, the SCN and vSPVZ exhibit in-phase activity such that the SCN and vSPVZ are both more active during the day than during the night (Nunez et al., 1999; Schwartz et al., 2004). In contrast, in adult Nile grass rats (Arvicanthis niloticus), a closely related Murid rodent (Campi et al., 2011) that exhibits behavioral and retinal hallmarks of diurnality (Blanchong et al., 1999; Gaillard et al., 2008), the SCN and vSPVZ exhibit anti-phase activity (Nunez et al., 1999; Schwartz et al., 2004). It is not known which features of the SCN or its downstream projections account for the different neural activity patterns in these two species (Schwartz et al., 2011).

Developmental investigations of sleep and wake behavior in altricial rodents, including Norway rats, are complicated by the fact that infants at the earliest ages, unlike adults, do not exhibit clear state-dependent electroencephalographic (EEG) activity (Gramsbergen, 1976; Seeke and Blumberg, 2008). However, sleep and wakefulness can be reliably studied in infant rats using electromyographic (EMG) and behavioral measures alone (Blumberg and Seeke, 2010). Using these methods, we recently examined the trajectory of day-night differences in sleep and wakefulness in developing Norway rats (Gall et al., 2008). Here, using a developmental comparative approach, we extend our analysis of circadian behavior to Nile grass rats. In infants of both species across the first two postnatal weeks, we identify day-night differences in behavior, Fos immunoreactivity (Fos-ir; a marker of neuronal activation) in the SCN and vSPVZ, RHT projections to the hypothalamus, and presynaptic terminals expressing pituitary adenylate cyclase-activating peptide (PACAP) or vasoactive intestinal peptide (VIP) in the vSPVZ. Our findings converge to support the hypothesis that developmental and species differences in RHT projections to the vSPVZ help to explain the emergence of circadian phase preference in these two species.

MATERIALS AND METHODS

All experiments were carried out in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals (NIH Publication No. 80-23) and were approved by the Institutional Animal Care and Use Committee of the University of Iowa.

Subjects

A total of 95 Nile grass rats from 53 litters and 84 Norway rats from 49 litters were used in this study. Males and females were distributed equally among all experimental groups. When littermates were used they were always assigned to different experimental groups. All pups were checked for clear milk bands before being used. Norway rats and grass rats were maintained in separate rooms in the animal colony at the University of Iowa. Litters of Norway rats were culled to eight pups within 3 days of birth (day of birth = Day 0). Because grass rat litters typically had fewer than eight pups, they were not culled. Grass rat litters averaged about six pups and no litters under four pups were used. All pups were housed with their mothers in standard laboratory cages and received food and water ad libitum. All rats were maintained on a 12-hour light-dark cycle, with lights on at 0700 for Norway rats and 0600 for grass rats.

Experiment 1: Development of day-night sleep and wake behavior in grass rats

Surgery

Twelve postnatal day (P)2 (body weights: 3.3–5.3 g), 12 P6–7 (hereafter P6; body weights: 6.4–10.7 g), and 12 P15 (body weights: 15.0–23.8 g) grass rats were used. Under isoflurane anesthesia, bipolar stainless steel hook electrodes (50 μm diameter; California Fine Wire, Grover Beach, CA) were implanted bilaterally in the nuchal muscle and secured with flexible collodion. A ground wire was looped through the back and secured with flexible collodion. Surgeries and anesthesia typically lasted 5–7 minutes. The pup was then transferred to the testing chamber and allowed at least 45 minutes to recover and acclimate. Surgeries performed at night were illuminated using a dim red light, with care being taken to ensure that pups were never exposed to white light.

Apparatus

The testing environment consisted of an electrically shielded, double-walled glass chamber (height = 17 cm, i.d. = 12.5 cm) with a Plexiglas lid. An access hole in the side of the chamber allowed for the passage of humidified air (flow rate: 300 ml/min) and EMG electrodes. Heated water circulated through the walls of the chamber to maintain air temperature at ≈35.5°C for P2 grass rats, 35°C for P6 grass rats, and 33.5°C for P15 grass rats. In preliminary studies, these temperatures were determined to be within the thermoneutral range for grass rats at these ages using similar methods for studying thermoneutrality in developing Norway rats (Spiers and Adair, 1986; Blumberg, 2001).

Recording procedure

During the day or night, freely moving pups were allowed to cycle undisturbed between sleep and...
were conducted at room temperature. Free-floating sections described previously (Todd et al., 2010). All reactions sliced in the coronal plane (40 μm) were incubated for 48 hours in 30% sucrose solution. After 5–10 days in 30% sucrose phosphate buffer (PB; pH 7.4), brains were removed and stored fused transcardially with phosphate-buffered saline (PBS; pH 7.4). Animals were killed with an overdose of Nembutal and perfused transcardially through the orbit. Approximately 10 ml of PBS was washed through the retina and vitreous cavity. At the point of the 6-hour test, tests during the day occurred from 0900 to 1500 and tests during the night occurred from 2100 to 0300. Daytime and nighttime tests were counterbalanced.

### Experiment 2: Development of day–night hypothalamic activity in Norway rats and grass rats

**Tissue preparation**

Twelve P8 (body weights: 17.7–25.8 g) and 12 P15 (body weights: 36.6–49.9 g) Norway rats and 12 P6 (body weights: 5.4–12.0 g) and 12 P15 (body weights: 19.3–30.5 g) grass rats were used. During the day or night, pups were removed from the litter and immediately sacrificed to ensure that these pups were never exposed to white light. Daytime and nighttime tests were counterbalanced.

### Experiment 3: Development of RHT projections in Norway rats and grass rats

**Eye injection**

Four P4 (body weights = 8.8–13.6 g), four P11 (body weights = 24.7–33.0 g), and four P17 (body weights = 49.0–54.2 g) Norway rats and four P2–3 (body weights = 4.7–8.3 g) and four P11 (body weights = 11.6–16.6 g) grass rats were used. Three additional adult grass rats (body weights = 90.0–135.0 g) were also used. Under isoflurane anesthesia, each rat received a unilateral intravitreal injection as follows: For P4 and P11 Norway rats and P2 grass rats, a 30G needle attached to a 10-μl Hamilton syringe (Reno, NV) was inserted through the skin between the sealed eyelids and into the orbit. For P17 Norway rats and P11 and adult grass rats, ages at which the eyelids have already opened, the procedure was the same except the needle was inserted directly into the orbit. Approximately 10 μl of Alexa Fluor 594 (Texas-Red)-conjugated CTB (1 mg/ml; Molecular Probes, Eugene OR) in PBS was slowly injected into the right or left eye.
left eye in a counterbalanced manner (Allen and Earnest, 2005). For adult grass rats, 10–15 μl of CTB solution was injected. Following the injection, the needle remained in the eye for 15 seconds before removal.

Tissue preparation and histology

Four days after the injection, P8, P15, and P21 Norway rats and P6 and P15 grass rats were overdosed and perfused as in Experiment 2. Adult grass rats were overdosed and perfused 4–6 days after the injection. Brains were postfixed, stored, and sliced as in Experiment 2. To preserve fluorescence, every other section was mounted and coverslipped using Vectastain Hard Set (Vector Laboratories). The remaining sections were mounted and Nissl-stained. They were then dehydrated with alcohol, cleared with citrus clearing solution, and coverslipped using Depex.

Experiment 4: Effect of light stimulation during the dark period on hypothalamic activity

Procedure

Twelve P8 (body weights = 14.3–21.8 g) and 12 P15 (body weights = 45.7–53.0 g) Norway rats were used. During the dark period, two littermates were removed and immediately placed—in separate rooms—into thermoneutral testing chambers identical to those described in Experiment 1. Chamber temperature was maintained at 35°C and 32°C for P8 and P15 rats, respectively. A dim red light was used for illumination in the dark. After 15 minutes of acclimation to the chamber, one littermate was maintained in the dark and the other was exposed to standard room lighting for 2 hours. The midpoint of the 2-hour test was timed to occur during the midpoint of the dark period, from 2400–0200.

Tissue preparation, immunohistochemistry, and histology

Immediately after the 2-hour exposure period, all subjects were overdosed and perfused as in Experiment 2. Brains were postfixed, stored, and sliced as in Experiment 2. Every other section throughout the hypothalamus was collected and Fos-immunohistochemistry was performed as previously described. The remaining sections were collected and Nissl-stained as in Experiment 3.

Experiment 5: Developmental expression of PACAP and VIP presynaptic terminals in the vSPVZ in Norway rats and grass rats

Tissue preparation

Six P8 (body weights = 19.4–23.8 g), six P11 (body weights = 22.8–30.8 g), six P15 (body weights = 36.0–48.1 g), and six P21 (body weights = 54.9–71.1 g) Norway rats and six P6 (body weights = 4.3–7.6 g), six P11 (body weights = 9.1–12.3 g), six P15 (body weights = 11.0–14.8 g), and six P21 (body weights = 22.8–25.4 g) grass rats were used. At the midpoint of their respective daytime photoperiods, pups were removed from the litter and immediately sacrificed. Accordingly, Norway rats were sacrificed at 1300 and grass rats were sacrificed at 1200. Animals were killed with an overdose of Nembutal and perfused transcardially with PBS followed by 2% PFA and 0.2% picric acid in PB. Brains were removed and stored in 30% sucrose solution for 5–10 days. Brains were then immersed in OCT compound (Sakura, Torrance, CA) and snap-frozen in isopentane cooled in a dry ice/ethanol bath. Using a cryostat, 18-μm sections were cut in the coronal plane and thaw-mounted onto gelatin-subbed slides.

PACAP and VIP immunohistochemistry

Sections were blocked in 2.5% bovine serum albumin and 0.1% Triton X-100 in PBS for 1 hour, followed by incubation with primary antibodies (Table 1) at 4°C overnight in the same solution. The monoclonal antibody against PACAP was collected from mice immunized with rat PACAP-38 conjugated to ovalbumin by glutaraldehyde. This antibody displays equal affinity for PACAP-38 and PACAP-27, recognizing an epitope between amino acids 6–10, but has no affinity with VIP (Hannibal et al., 1995). Preabsorption with PACAP-38 or PACAP-27 blocks staining (Hannibal et al., 1997). This antibody has been previously used to detect PACAP in the RHT and exhibits a characteristic pattern of staining in the SCN (Hannibal et al., 2000; Hannibal, 2002b).

The polyclonal antibody against VIP was collected from guinea pigs immunized with synthetic VIP (antigen sequence, H-His-Ser-Asp-Ala-Val-Phe-Thr-Asp-Asn-Tyr-Arg-Val-Leu-Arg-Lys-Asn-Val-Asp-Ser-Thr-Phe-Asp). Radiimmunoassay shows no crossreactivity with PHI-27, PHM-23, VIP (10-28), VIP (1-12), VIP guinea pig and chicken, PACAP-38, and substance P (manufacturer’s technical information). This antibody exhibits a characteristic pattern of staining in the SCN (Belenky et al., 2008; Vida et al., 2010).

Sections were washed in PBS and secondary antibodies (Alexa Fluor 488- or 568-conjugated; 1:200; Molecular Probes) were applied in PBS for 1 hour. Nuclei were counterstained with 4',6-diamidino-2-phenylindole (Sigma-Aldrich) added to the final wash. For sections serving as negative controls, the procedure was the same except primary antibodies were omitted. Slides were coverslipped with Fluorogel (Electron Microscopy Sciences, Hatfield, PA).
Confocal microscopy
Sections were imaged sequentially using a SP-5 Leica confocal laser scanning microscope (Leica Microsystems, Germany) with an argon 488 nm laser and a helium/neon 543 nm laser. Two sections containing the vSPVZ were selected from each animal at matching rostrocaudal levels across all groups. For each section, using an oil immersion 63× objective and a magnification factor of 4, a z-stack of optical images at 1-μm layer intervals was acquired and digitized into 512 × 512 pixel bitmaps. The area imaged corresponded to a 61.5-μm-wide square area within the vSPVZ. Laser settings were held constant across all subjects.

Data analysis
Sleep and wake behavior
EMG data from Experiment 1 were analyzed as described previously (Karlsson et al., 2005). Briefly, the records were dichotomized into bouts of sleep and wakefulness as follows: For each rat, the amplitudes of 5 1-second segments of both atonia and high-tone were averaged and the midpoint between the two was determined. Periods of at least 1 second in which muscle tone was below the midpoint were defined as atonia (indicative of sleep), whereas those above the midpoint were defined as high tone (indicative of wakefulness). An experienced individual blind to experimental condition scored all data.

Sleep and wake bout durations for all pups were imported into JMP 5.0 (SAS Institute, Cary, NC). Mean bout durations for sleep and wakefulness were determined for each pup and percentage of time awake was found by dividing the mean wake bout duration by the sum of the mean sleep and wake bout durations and multiplying by 100. A two-factor analysis of variance (ANOVA) was used to test for differences across age and time of day, and planned comparisons (unpaired t-tests) served to reveal day–night differences within groups. Additionally, data were pooled for each pup in order to produce survivor distributions of sleep and wakefulness, as described elsewhere (Blumberg et al., 2005). For all tests, alpha was set at 0.05.

Fos-immunohistochemistry
Images of 40-μm brain sections were captured using a Leica DM/LS microscope and imaging system with Firecam software (Leica). Images were imported into ImageJ (National Institutes of Health, http://imagej.nih.gov/ij/) and adjusted to binary values. Using methods similar to those described elsewhere (Todd et al., 2010), cells positive for Fos-ir were identified. Typically, cells were ∼10 μm in diameter. For each subject, 2–3 sections (in Experiment 2) or 3–4 sections (in Experiment 4) containing the SCN and vSPVZ were selected for analysis. Using a counting box of known dimensions, profiles of Fos-positive cells were counted unilaterally by an experienced individual blind to experimental group. An Abercrombie correction was performed on the counts for each area within each section (Abercrombie, 1946). The mean number of Fos-positive cells per section was then calculated for each pup.

The SCN and vSPVZ were identified using a stereotaxic atlas of the rat brain (Paxinos and Watson, 1998), as well as previous publications describing these structures in adult grass rats and Norway rats (Lu et al., 2001; Gooley et al., 2003; Schwartz et al., 2004, 2009). Although cell bodies were not counterstained in Experiment 2, brain areas were determined on the basis of adjacent anatomical structures (e.g., third ventricle, optic chiasm). When needed, archival Nissl-stained sections from separate animals at similar ages were used to help delineate boundaries between areas. Corresponding Nissl-stained sections from the same animals were used to determine nuclei in Experiment 4. Data were imported into JMP 5.0 and paired or unpaired t-tests were used to reveal group differences within each brain area. Alpha was set at 0.05 and Bonferroni corrections were performed to correct for multiple comparisons.

RHT projections
For Experiment 3, unstained sections were analyzed for the presence or absence of CTB labeling in the SCN and vSPVZ using a Leica DM/LS fluorescent microscope and imaging system with Firecam software. Light intensity and contrast levels were held constant for all sections across all subjects. Using a 20× objective, images of consecutive fluorescent and corresponding Nissl-stained sections were collected and then imported into Adobe Photoshop CS3 (Adobe Systems, San Jose, CA) to determine the locations of CTB labeling. To compare age differences in CTB labeling in the vSPVZ, two fluorescent sections were selected from each subject for further analysis. Images were scored by an individual blind to the age and species of each subject. Mean fluorescence intensity (MFI) in the red channel was quantified, contralateral to the injected eye, for each section using ImageJ software containing the RGB Measure plug-in (Bubier et al., 2007). MFI per section was then calculated for each subject. Data were imported into Statview 5.0 (SAS Institute). ANOVA was used to test for differences across ages within each species, and Fisher’s PLSD was used for post-hoc comparisons. For all tests, alpha was set at 0.05.

PACAP and VIP immunohistochemistry
To measure presynaptic terminals in the vSPVZ in Experiment 5, images of two confocal layers, evenly
spaced within each z-stack, were extracted using Leica LAS AF software and imported into ImageJ. Data were scored by an experimenter blind to the age and species of each subject. Red- and green-fluorescent presynaptic terminals were automatically counted using the ImageJ Puncta Analyzer plugin as described elsewhere (Ippolito and Eroglu, 2010). Mean values of PACAP and VIP terminals were calculated for each section and then means were calculated for each pup. Data were imported into Statview 5.0. ANOVA was used to test for differences across age within each species and planned comparisons (unpaired t-tests) were used to test developmental changes between P6/8 and P21. For all tests, alpha was set at 0.05.

**Photomicrographs**

Sizes of photomicrographs were adjusted and figure plates were prepared using Adobe Photoshop CS3. No other manipulations were performed on representative images.

**RESULTS**

**Development of circadian phase preference**

We first compared the developmental trajectory of day–night behavioral differences in Norway rats and grass F1 rats. Figure 1 presents log-survivor distributions for wake bouts and mean percentage of time awake (insets) in Norway rats and grass rats. The data for Norway rats have been published previously (Gall et al., 2008) and show that nocturnal wakefulness emerges between P8 and P15. Here, we recorded day–night differences in sleep and wakefulness in grass rats at P2, P6 (i.e., 1 day after eye opening), and P15. For mean percentage of time awake (insets in Fig. 1), ANOVA revealed significant main effects of age \( F(2,30) = 21.3, P < 0.0001 \) and time \( F(1,30) = 33.7, P < 0.0001 \), but not a significant age × time interaction \( F(2,30) = 0.1 \). Planned comparisons revealed that mean percentage of time awake was higher during the day at P15 \( (t(10) = 7.5, P < 0.0001) \). In grass rats at both P6 and P15 (Fig. 2A, top), the mean number of Fos-positive cells in the SCN was greater during the day than during the night \( (P8: t(10) = 7.5, P \leq 0.001; \text{P15: } t(10) = 5.8, P \leq 0.002) \). In the vSPVZ at P8, Fos values did not differ between day and night \( (t(10) = 1.0) \), whereas higher values were found during the day at P15 \( (t(10) = 6.3, P \leq 0.0001) \). In grass rats at both P6 and P15 (Fig. 2B, top), Fos values in the SCN were again greater during the day than during the night \( (P6: t(10) = 7.0, P \leq 0.0005; \text{P15: } t(10) = 13.8, P \leq 0.0001) \). In the vSPVZ at P6, Fos values did not differ between day and night \( (t(10) = 0.1) \), whereas higher values were found during the night at P15 \( (t(10) = 9.7, P \leq 0.0001) \). These patterns, at P15, of increased vSPVZ activity during periods of behavioral inactivity is typical of adults of these species (Schwartz et al., 2004).

**Norway rats, but not grass rats, develop a robust direct projection from RHT to vSPVZ**

In Norway rats, the RHT projects to the SCN as early as P1 (Hannibal and Fahrenkrug, 2004). Moreover, by adulthood the RHT of Norway rats also projects directly to the vSPVZ (Gooley et al., 2003). Based on the Fos-ir data described above, we hypothesized that the direct RHT connection to the vSPVZ would develop between P8 and P15. We tested this hypothesis using CTB tracing in P8, P15, and P21 Norway rats \( (n = 4 \text{ at each age}) \). At P8, as expected, CTB labeling was detected in the SCN but only sparse and inconsistent labeling was found in the vSPVZ (Fig. 3A–C). In contrast, at P15 strong labeling was found in both the SCN and the vSPVZ (Fig. 3D–F). Finally, 1 week later at P21, this vSPVZ labeling was not noticeably denser (Fig. 4A,B). As shown in Figure 5, MFI in the vSPVZ of Norway rats increased progressively between P8, P15, and P21. ANOVA revealed a significant main effect of age on MFI \( F(2,9) = 7.8, P < 0.05 \).

Based on the Fos-ir data described above, we hypothesized that the RHT of grass rats does not project directly
to the vSPVZ. We tested this hypothesis using CTB tracing in P6 and P15 subjects (n = 4 at each age) and in three adults. As predicted, we found intense CTB labeling of the SCN at P6 and P15 but little or no such labeling in the vSPVZ at those ages (Fig. 3G–I). We similarly found little or no labeling in the vSPVZ of the three adult grass rats.
As shown in Figure 5, MFI in the vSPVZ of grass rats was very low at P6 and did not increase significantly by P15 or adulthood [$F(2,8) = 3.1$]. Finally, to ensure that our tracing method was reliable in this species to structures beyond the SCN, we confirmed labeling in the supraoptic nucleus (SON) in all subjects.

**Development of PACAP and VIP presynaptic terminals in the vSPVZ**

We examined presynaptic terminal markers in the vSPVZ across development in Norway rats and grass rats. As shown in Figure 7, the mean number of PACAP-expressing terminals in the vSPVZ of Norway rats increased progressively from P8 to P21. ANOVA revealed a significant main effect of age [$F(3,20) = 6.7$, $P < 0.005$]. In grass rats, the mean number of PACAP-expressing terminals in the vSPVZ was very low at P6 but, in contrast to Norway rats, did not increase significantly with age [$F(3,20) = 1.4$]. Planned comparisons revealed that the number of PACAP-expressing terminals in the vSPVZ increased significantly from P6/8 to P21 in Norway rats ($t(10) = 4.3$, $P \leq 0.01$), but not grass rats ($t(10) = 0.3$). Representative images of PACAP-expressing terminals in a P21 Norway rat and grass rat are shown in Figure 8.

As also shown in Figure 7, the mean number of VIP-expressing terminals in the vSPVZ in Norway rats...
increased between P8 and P21. ANOVA revealed a significant main effect of age \(F(3,20) = 6.2, P < 0.005\). In grass rats, the mean number of VIP-expressing terminals in the vSPVZ also increased between P6 and P21; however, ANOVA did not reveal a significant main effect of age \(F(3,20) = 2.7, P = 0.07\). Planned comparisons revealed that, for both species, VIP-expressing terminals in the vSPVZ increased significantly from P6/8 to P21 \((t(10) > 3.1, P < 0.05)\).

**DISCUSSION**

We have shown here how developmental processes can differentially shape brain and behavior to alter circadian phase preference in Norway rats and Nile grass rats, two closely related Murid rodents (Campi et al., 2011). First, nocturnal Norway rats and diurnal grass rats undergo profound developmental changes in circadian wake behavior between the first and second postnatal weeks. Second, these behavioral changes are associated with the emergence of an in-phase pattern of SCN and vSPVZ activity in Norway rats and an anti-phase pattern in grass rats. Third, Norway rats develop an abundance of RHT projections to the vSPVZ between P8 and P15, whereas grass rats do not. Fourth, in support of our anatomical evidence, 2 hours of light stimulation during the dark period in Norway rats activates the vSPVZ at P15 but not P8. Finally, Norway rats, but not grass rats, exhibit significant increases in PACAP-expressing presynaptic terminals in the vSPVZ during the second postnatal week, thereby providing additional evidence of species differences in RHT connectivity. This converging evidence across...
five experiments suggests that the developmental pattern and timing of retinal projections to the hypothalamus contribute to the development of circadian phase preference in these two species.

Multiple lines of evidence support the vSPVZ as a fundamental component of the neural circuitry contributing to circadian phase preference. In adult Norway rats and grass rats, lesions of the vSPVZ selectively disrupt circadian rhythms of sleep and wake behavior (Lu et al., 2001; Schwartz et al., 2009). Also, the vSPVZ receives the majority of the SCN’s output and projects to the same downstream structures as the SCN, making it an ideal structure for mediating and modulating circadian information (Smale et al., 2003; Schwartz et al., 2011). Finally, as mentioned previously, adult Norway rats and grass rats also express in-phase and anti-phase profiles of SCN-vSPVZ activity, respectively (Nunez et al., 1999; Schwartz et al., 2004), which we show here are first expressed contemporaneously with developmental changes in circadian wakefulness.

Interestingly, it has been reported in adult mice that the SCN and subparaventricular zone (SPVZ) exhibit an anti-phase pattern of multiunit activity (Nakamura et al., 2008, 2011). Given that mice are nocturnal, this pattern of activity is opposite to what we would predict based on the present findings and those of others using Norway rats (Nunez et al., 1999; Schwartz et al., 2004). However, in the one Nakamura et al. (2008; supplemental data) article where recording sites were indicated, some sites appear to be dorsal and rostral to the vSPVZ and none was confirmed as being within areas targeted by the RHT. These issues may be important because the SPVZ is a heterogeneous structure with dorsal and ventral subregions that make separable contributions to locomotion, sleep, and thermoregulation (Lu et al., 2001).

Our understanding of the neurotransmitters involved in the RHT-SCN-vSPVZ circuit comes largely from work in adults. In Norway rats, the axon terminals of RHT projections co-store glutamate and PACAP, which have excitatory postsynaptic effects (Hannibal, 2002a). In contrast, the major projection from the SCN to the vSPVZ in Norway rats is composed of axon terminals that co-store γ-aminobutyric acid (GABA) and VIP and has been shown to have primarily inhibitory postsynaptic effects (Hermes et al., 2009). In adult grass rats, the vSPVZ also receives GABAergic and VIPergic projections from the SCN.
Mahoney et al., 2007); because there is only a sparse RHT projection to the vSPVZ in this species (Smale and Boverhof, 1999), we expected PACAP-expressing terminals in this region to be sparse as well, which is what we found (Fig. 8B). Figure 9 presents a model, based on our results, of the developmental and species differences in the neural circuitry underlying circadian phase preference in Norway rats and grass rats. According to this model, the vSPVZ of Norway rats does not display day–night differences in neural activity at P8, and receives little or no input from the retina or SCN at this age. By P15, however, the direct

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**Figure 5.** Developmental differences in mean fluorescence intensity (MFI) per section in the ventral subparaventricular zone (vSPVZ) following retinal tracing in P8, P15, and P21 Norway rats (left) and P6, P15, and adult Nile grass rats (right). n = 4 per group except for adult grass rats (n = 3). *Significant difference from two other groups, P < 0.05. Means + standard errors.

**Figure 6.** Effect of 2 hours of light exposure during the dark period on mean number of Fos-positive cells per section in the suprachiasmatic nucleus (SCN) and ventral subparaventricular zone (vSPVZ) of P8 and P15 Norway rats (n = 6 per group). Black bars represent control subjects maintained in the dark and hatched bars represent experimental subjects exposed to light. *Significant difference from control group, P < 0.005. Means + standard errors.

**Figure 7.** Developmental differences in the mean number of presynaptic terminals in the ventral subparaventricular zone (vSPVZ) in P8, P11, P15, and P21 Norway rats (left) and P6, P11, P15, and P21 Nile grass rats (right). n = 6 per group. Black bars represent terminals expressing pituitary adenylate cyclase-activating peptide (PACAP) and hatched bars represent terminals expressing vasoactive intestinal peptide (VIP). *Significant difference from P6/8, P < 0.05. Means + standard errors.
excitatory RHT projection to the vSPVZ has developed, as has the inhibitory projection arriving from the SCN. To produce the in-phase activity pattern in the vSPVZ, we surmise that the excitation provided by the direct RHT projection must exceed the inhibition provided by the SCN-vSPVZ connection. To better understand the relative contributions of the retinal and SCN connections to the vSPVZ, we would ideally want to selectively disable these connections to observe their effects on hypothalamic activity. However, such selective manipulations are not currently feasible.

As also depicted in Figure 9, grass rats possess an excitatory RHT projection to the SCN, but they do not develop the robust RHT projection to the vSPVZ exhibited by Norway rats. At P6, the vSPVZ does not display day-night differences in neural activity, presumably due to the relative lack of input from the SCN. By P15, however, the anti-phase pattern of SCN-vSPVZ activity develops and is associated with an increase in inhibitory projections to the vSPVZ.

For our functional test of RHT connectivity in Norway rats, we used Fos immunohistochemistry to measure...
neuronal activity changes in the SCN and vSPVZ in response to 2 hours of light stimulation during the dark period. As predicted, whereas P8 subjects exhibited increased activity only in the SCN, P15 subjects exhibited increased activity in both the SCN and vSPVZ. Although Fos immunohistochemistry is suited to detecting increases in neuronal activity, it is not suited to detecting short-term decreases because, once the Fos protein is produced, it takes several hours to degrade (Cirelli and Tononi, 2000). Nonetheless, for completeness and despite the likelihood of ambiguous results, we repeated the light-exposure test in a small group of grass rats at P15 (data not shown). As was found with Norway rats, 2 hours of light exposure during the dark period significantly increased Fos-positive cells in the SCN; however, in contrast to Norway rats, Fos activity in the vSPVZ of control subjects was already very high (due to the anti-phase SCN-vSPVZ relationship in this species) and light exposure neither increased nor decreased this activity. A failure to detect an increase in vSPVZ activity could have been due to a ceiling effect. But we believe it is more likely that the failure to detect a decrease in vSPVZ activity resulted from insensitivity of the c-fos method for detecting short-term functional inhibition. It should also be noted that the converse experiment—turning off the lights during the light period—would not resolve these interpretational difficulties. Direct measures of neurophysiological activity may be better able to detect the anti-phase relationship in grass rats.

The vSPVZ exhibited strikingly higher Fos-ir levels in grass rats at P6 than in Norway rats at P8 (Fig. 2). There are several possible causes of this species difference. One possibility is that vSPVZ neurons are more spontaneously active in grass rats than in Norway rats. Another possibility is that the vSPVZ in grass rats is activated by input from another neural structure. Whatever the cause, species differences in vSPVZ activity may provide a mechanism for effecting species differences in RHT connectivity. For example, the action of ephrins, which repel developing axons, is modulated by activity-dependent processes (Klein, 2009).

To our knowledge, it has not previously been hypothesized that RHT connectivity to the vSPVZ helps determine circadian phase preference. Keeping in mind the need for caution when comparing data from multiple laboratories using different techniques, published reports suggest that this hypothesis is supported by a variety of other species. As shown in Table 2, nocturnal species including hamsters (Speh and Moore, 1993; Morin et al., 2003), mink (Peytevin et al., 1997), mice (Cassone et al., 1988; Hattar et al., 2006), and opposums (Cassone et al., 1988) possess robust RHT projections extending beyond the SCN to the vSPVZ. In contrast to nocturnal species and in addition to grass rats, RHT afferents do not extend significantly beyond the boundary of the SCN in such diurnal species as humans (Dai et al., 1998), Cebus monkeys (Pinato et al., 2009), ground squirrels (Smale et al., 1991), and four-striped field mice (Schumann et al., 2006).
Table 2 also shows two notable exceptions to the above pattern. Specifically, both sheep (Torrealba et al., 1993) and degus (Goel et al., 1999) are diurnal but exhibit robust RHT projections extending dorsally and caudally outside the SCN. Interestingly, these two species are precocial in relation to all of the nocturnal species discussed above. Moreover, this direct projection develops postnatally in rats (Speh and Moore, 1993), hamsters (Speh and Moore, 1993), and mink (Peyetin et al., 1997). Thus, we hypothesize further that, for animals that develop the direct retina-to-vSPVZ connection, the precise timing of innervation may influence its subsequent effect on circadian phase preference. Although there are many behavioral and physiological factors that differ across the altricial–precocial spectrum that could potentially alter SCN function and connectivity, exposure to light may not be a critical factor since dark rearing of Norway rats has little or no effect on the development of RHT connectivity or circadian phase preference (Prichard et al., 2004, 2007).

Table 2 also indicates that RHT projections in guinea pigs and domestic cats are confined to the SCN (Cassone et al., 1988). However, substantial disagreement exists as to the circadian phase preference of these two species (Bowersox et al., 1984; Jilge, 1985; Kuwabara et al., 1988; Tobler and Scherschlicht, 1990; Tobler et al., 1993; Akita et al., 2001). Overall, our findings suggest that species-typical circadian phase preferences emerge through a developmental rewiring of the RHT to effect differential interactions among the retina, SCN, and vSPVZ. It bears noting that the mechanism identified here may be one of several; in other words, there may be multiple developmental paths to nocturnality and diurnality. Whatever the mechanism, once the system is established, most individuals will exhibit a species-typical bias toward one or the other (although it should be noted that this bias does not prevent individuals from shifting to the opposing temporal niche under certain ecological conditions; Mrosovsky and Hattar, 2005). Altogether, the present results indicate that a fuller understanding of the mechanisms underlying nocturnality and diurnality will require delineation, in a variety of species, of the developmental relations among RHT connectivity, hypothalamic activity, and behavior.

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