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SCN VIP Neurons Are Essential for Normal Light-Mediated Resetting of the Circadian System

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The suprachiasmatic nucleus (SCN) synchronizes circadian rhythms in behavior and physiology to the external light cycle, but the mechanisms by which this occurs are unclear. As the neuropeptide vasoactive intestinal peptide (VIP) is important for circadian light responses, we tested the hypothesis that rhythmic VIP-producing SCN neurons mediate circadian light responses in male and female mice. Using in vivo fiber photometry over multiple days, we found daily rhythms in spontaneous calcium events of SCN VIP neurons that peaked during the subjective day and were disrupted by constant light. The light-evoked calcium responses peaked around subjective dusk and were greater during the subjective night. Using novel VIP sensor cells, we found that the activity patterns in SCN VIP neurons correlated tightly with spontaneous and NMDA-evoked VIP release. Finally, in vivo hyperpolarization of VIP neurons attenuated light-induced shifts of daily rhythms in locomotion. We conclude that SCN VIP neurons exhibit circadian rhythms in spontaneous and light-responsive activity and are essential for the normal resetting of daily rhythms by environmental light.

Key words: calcium; circadian; light; neuropeptide; suprachiasmatic; vasoactive intestinal peptide

Significance Statement

Daily rhythms in behavior and physiology, including sleep/wake and hormone release, are synchronized to local time by the master circadian pacemaker, the suprachiasmatic nucleus (SCN). The advent of artificial lighting and, consequently, light exposure at night, is associated with an increased risk of disease due to disrupted circadian rhythms. However, the mechanisms by which the SCN encodes normal and pathological light information are unclear. Here, we find that vasoactive intestinal peptide (VIP)-producing SCN neurons exhibit daily rhythms in neuronal activity and VIP release, and that blocking the activity of these neurons attenuates light-induced phase shifts. We conclude that rhythmic VIP neurons are an essential component of the circadian light transduction pathway.

Introduction

Organisms have evolved endogenous circadian (~24 h) rhythms in behavior and physiology, including sleep/wake, hormone release, and metabolism (Kalsbeek et al., 2006), to anticipate reliable daily events such as the light cycle. In mammals, light at night, but not during the day, shifts a master circadian pacemaker in the suprachiasmatic nucleus (SCN) to entrain daily rhythms to local time (Gillette, 1996; Sakamoto et al., 2013). Phase-shifting environmental light is transduced to the SCN through melanopsin-containing, intrinsically photosensitive retinal ganglion cells through the release of glutamate and, perhaps, the neuropeptide PACAP to increase the firing rate of SCN neurons (Itaka et al., 1989; Meijer et al., 1993; Michel et al., 2006; Morin and Allen, 2006; Fernandez et al., 2016). This increase in firing rate increases intracellular calcium (Irwin and Allen, 2007), which activates a signaling cascade that shifts the neuron’s molecular clock (Nitabach et al., 2002; Yamaguchi et al., 2003; Jones et al., 2015), ultimately shifting behavioral and physiological rhythms. Thus, the SCN is able to encode changes in environmental light that occur during, for instance, seasonal changes in the wild (Lucassen et al., 2012), but is limited in its ability to track larger shifts associated with jet lag or shift work schedules (Sack et al., 2007). Importantly, chronic nocturnal light exposure disrupts circadian rhythms in the SCN and behavior (Granados-Fuentes et al., 2004; Ohta et al., 2005) and is associated with an increased risk of disease (Fonken et al., 2010; LeGates et al., 2012; Chang et al., 2015; Lucassen et al., 2016). We do not know, however, which cell types mediate normal and pathological phototransduction in the SCN.
The neuropeptide vasoactive intestinal peptide (VIP), produced by retinoenocrine neurons comprising ~10% of the SCN network, is the predominant contributor to synchrony in the SCN (Abrahamson and Moore, 2001; Colwell et al., 2003; Aton et al., 2005; Maywood et al., 2011) and has been implicated in photoentrainment. For instance, exogenous VIP administration can cause circadian phase shifts both in vitro and in vivo (Piggins et al., 1995; Reed et al., 2001; Atkins et al., 2010; An et al., 2011), and VIP−/− mice start their daily activity ~8 h earlier than wild-type mice (Colwell et al., 2003; Voako et al., 2015). Because phase-shifting light pulses upregulate markers of neuronal activation in VIP neurons (Romijn et al., 1996; Kuhlman et al., 2003), we tested the hypothesis that VIP neurons mediate light responses in the SCN. Furthermore, we sought to clarify whether VIP neuron activity varies with time of day in vivo given the equivocal in vitro evidence (Fan et al., 2015; Hermanstyn et al., 2016; Enoki et al., 2017). Finally, we developed a novel measure of VIP release to address the questions of if and when SCN neurons release VIP spontaneously and in response to stimulation. We found that SCN VIP neurons are circadian and excited by light in vivo, release VIP depending on time of day and excitatory input, and are necessary for the normal transduction of light signals to the SCN at times when light shifts daily rhythms.

Materials and Methods

**Animals.** All experiments were performed using approximately equal numbers of male and female mice in accordance with the Washington University Institutional Animal Care and Use Committee guidelines. No differences in results were observed between sexes. For in vitro fiber photometry and chemogenetics experiments, we used heterozygous VIP-ires-Cre knock-in mice (VIP<sup>tm1criZ220</sup>/); catalog #010908. The Jackson Laboratory: RRID:IMS_JAX:010908; Taniguchi et al., 2011) on a C57BL/6J/N background that were at least 6 weeks old at the time of virus injection and ~2-5 months old during recordings. To help identify VIP cells in some experiments, mice were crossed to Ai9 or Ai32 (B6.Cg-Gt(ROSA)26Sortm14(Cdx2<sup>Cre</sup>)/Elsv; flox/flox; B6.Cg-Gt(ROSA)26Sortm14(Cdx2<sup>Cre</sup>)/Elsv; flox/flox; B6.Cg-Gt(ROSA)26Sortm14(Cdx2<sup>Cre</sup>)/Elsv; flox/flox; B6.Cg-Gt(ROSA)26Sortm14(Cdx2<sup>Cre</sup>)/Elsv; flox/flox; B6.Cg-Gt(ROSA)26Sortm14(Cdx2<sup>Cre</sup>)/Elsv; catalog #007990 or #012569, The Jackson Laboratory; RRID:IMS_JAX:007990 or IMSR_JAX:0012569) mice to yield heterozygous VIP-ires-Cre; Ai9 or VIP-ires-Cre; Ai32 mice that expressed a tdTomato or enhanced yellow fluorescent protein reporter in VIP neurons. For in vitro SCN recordings, we collected SCN slices from wild-type or VIP−/− (founders were provided by C. Colwell, University of California, Los Angeles, Los Angeles, CA; RRID:IMS_JAX:009640) mice (~7-10 d old) on a C57BL/6J/N background. After weighing and before recording, up to 5 mice of the same sex were housed per cage in a 12 h light/dark LD (where lights on is defined as zeitgeber time (ZT) 0; light intensity ~2 × 10<sup>14</sup> photons/cm<sup>2</sup>/s) cycle with food and water available ad libitum and constant temperature (~22°C) and humidity (~40%).

**Virus injection and stereotaxic surgery.** Cre-inducible adeno-associated virus (AAV) vectors encoding GCaMP6s (AAV9-CAG-Flex-GCaMP6s-WPRE-SV40) and EGFP (AAV9-CAG-Flex-E-GFP-WPRE-BGH) were obtained from the University of Pennsylvania Vector Core (AV-9-ALL854 and AV-9-PV2818) and used at a final concentration of ~1 × 10<sup>12</sup> genome copies (GC)/ml. Cre-inducible AAV vectors encoding hM4Di (AAV8-hShyn-D10-hM4Di(G4)-mCherry) were obtained from Addgene (catalog #44362) and used at a final concentration of ~2 × 10<sup>12</sup> GC/ml. At the start of surgery, mice received preoperative analgesia (carprofen, 5 mg/kg), were anesthetized with isoflurane (induction dose, ~3%; maintenance dose, ~1.5%), and placed in a stereotaxic frame (Kopf) on a heating pad to maintain body temperature throughout the procedure. We then infused virus to the SCN (~0.46 mm posterior, ±0.15 mm lateral, ~5.65 mm ventral from bregma) at a rate of 0.05 µl/min through a 30 gauge needle attached to a 2 µl syringe (Neurog, Hamilton). The total volume injected for fiber photometry experiments was 0.3 µl unilaterally (GCaMP6s and EGFP) and, for chemogenetics experiments, was 0.3 or 0.5 µl bilaterally (EGFP or hM4Di, respectively). After infusion, the needle was left in place for at least 8 min before slowly being withdrawn. For fiber photometry experiments, we subsequently implanted a fiber-optic cannula [6 mm in length; 400-µm-diameter core; 0.48 numerical aperture (NA); male ferrule; Doric Lenses] immediately dorsal to the virus injection site. Cannulas were secured with a layer of adhesive cement (C&B Metabond, Parkell) followed by a layer of opaque black dental acrylic (Ortho-Jet, Lang). After surgery, skin was closed with sutures or surgical glue (Vetbond, 3M) and mice were allowed to recover on a heating pad until ambulatory. Mice were then transferred to their home cages for 2-4 weeks to allow for virus expression.

In vivo fiber photometry recording. After recovery and virus expression, mice were transferred to individual, open-top cages equipped with running wheels and bedding maintained in a light-, temperature-, and humidity-controlled chamber with food and water available ad libitum. Lights were set to 12 h LD, constant darkness (DD), or constant light (LL; light intensity, ~2 × 10<sup>14</sup> photons/cm<sup>2</sup>/s). Freely moving mice were chronically tethered to a fiber-optic cable (400 µm core; 0.48 NA; 1.5 m in length; Doric Lenses) connected to the implanted optic fiber by a zirconia mating sleeve (Doric Lenses). Running wheel activity was recorded in 6 min bins using Clocklab software (Actimetrics; RRID:SCR_014309). After at least 3 d of acclimation to tethering, we began fiber photometry recordings using established methods (Eban-Rothsheld et al., 2016). To control for motion artifacts, we excited GCaMP6s with its calcium-dependent excitation wavelength of 470 nm and at its calcium-independent isosbestic excitation wavelength of 405 nm (Lerner et al., 2015). Blue 470 nm and violet 405 nm LED lights (M470F9 and M405F11, Thorlabs) were sinusoidally modulated at 211 and 531 Hz, respectively, using a custom Matlab program (MathWorks; RRID:SCR_001622) and a multifunction data acquisition device (NI PCI-6110, National Instruments). The excitation lights passed through a fluorescence cube (FM04, Doric Lenses) containing two separate excitation filters (450–490 and 405 nm), reflected off a dichroic mirror and coupled into the tethered fiber-optic cable. The light intensity for each wavelength was set to ~30 µW at the termination of the fiber-optic cable. Sinusoidally modulated GCaMP6s fluorescence was collected through the same fiber-optic cable, passed through an emission filter (500–550 nm), and focused through a patch cord onto a femtowatt photoreceiver (model 2151, Newport). The photoreceiver signal was then sent to two lock-in amplifiers (30 ms time constant; model SR830, Stanford Research Systems) that were synchronized to 211 or 531 Hz for demodulation. Voltage signals from the amplifiers were collected at 10 kHz using a custom Matlab program and a multifunction data acquisition device and saved to disk as binary files. For spontaneous activity experiments, hourly 10 min recordings were obtained automatically using a custom Matlab program. For light-evoked activity experiments, hourly recordings were collected for 15 min, divided by the daily mean light exposure, and collected for 15 min. Chamber lights were turned on at 2 min after the 10 min data acquisition period. Animals were excluded from analysis if both the calcium-dependent and calcium-independent signals simultaneously fluctuated by ~100 mV (an indicator that the fiber was positioned below the SCN), if they did not exhibit circadian locomotor activity or if the fiber or virus was off-target as determined by histology.

**Fiber photometry data analysis.** Data files were opened in Matlab and downsampled to 1 kHz for further analysis as in the study by Zaloucysky et al. (2016). To account for different photobleaching dynamics, extracted calcium-dependent (470 nm) and calcium-independent (405 nm) signals were individually fit with an exponential filter, and the fitted signals were subtracted from the raw signals. Signals were then bandpass filtered between 0.016 and 2 Hz (spontaneous recordings) or 0.003 and 2 Hz (evoked recordings) and fit to one another using a linear-least-squares fit. The change in fluorescence over baseline fluorescence (ΔF/F) was calculated as (488 nm signal-fitted 405 nm signal)/(fitted 405 nm signal). Final values were adjusted so that a ΔF/F of 0 corresponded to the second percentile of the signal. ΔF/F values were converted to events by counting instances >1 s when ΔF/F exceeded the median plus 2 SDs of the trace or ~1.5% ΔF/F, whichever was larger. We report the number of events per minute and also the integrated calcium values, as defined by integrating the ΔF/F values >0% for >10 min. For light-evoked recordings, re-
response amplitude was defined as the peak ΔF/F during 2 min following the start of the light pulse divided by the median ΔF/F during the 2 min preceding the light pulse, and response duration was defined as the time from the end of the light pulse until the signal returned to the median ΔF/F of the entire recording.

**VIP sensor cell production.** The cognate VIP receptor in the SCN, VPAC2R, is preferentially coupled to a Gs-/adenylate cyclase–cAMP pathway (Langer, 2012). To generate reporter cells (Müller et al. 2014; Giswole et al. 2016) that respond to VIP through a Gs-βphosphoprotein-C–calcium pathway (Offermanns and Simon, 1995; Zhu et al., 2008), HEK293 cells (American Type Culture Collection; RRID:CVCL_0045) were sequentially transfected with pcDNA3.1 (+)·hVIPr2 (neomycin/kanamycin selection sequence replaced with puromycin sequence) and then pcDNA3.1 (+)·hGsbetaS (cDNA Resource Center). Sixteen microtiter of PuGene Transfection Reagent (Promega) and 1 μg of plasmid DNA were diluted in Opti-MEM (Thermo Fisher Scientific) to a final volume of 250 μL. Transfection medium in 2 mL of DMEM with 5% fetal bovine serum (FBS; Thermo Fisher Scientific) was added to HEK293 cells at 60–80% confluency and removed after 15–20 h. Transfected cells were maintained at 37°C and 5% CO2 in an atmosphere of 10% FBS and 1% penicillin/streptomycin (Thermo Fisher Scientific) for 2 d. On day 3, the antibiotic was replaced with 0.4 μg/mL puromycin (Thermo Fisher Scientific). Cells were selected for 10 d. Single-transfected and selected cells were then transfected with the second plasmid, as described. Double-transfected cells were selected with 550 μg/mL G418 sulfate (Thermo Fisher Scientific) for 10 d. Doses of puromycin and G418 were determined through a HEK293 cell curve using ranges of 0.36–0.44 and 200–800 μg/mL, respectively. Double-transfected and selected cells (HEK293 VIPr2·GhGS+) were kept in media with puromycin and G418 antibiotics until ready for reporter transfection and recording. To verify successful double transfection, cells were immunostained with the primary antibodies rabbit anti-VPAC2 (1:500; catalog #ab28624, Abcam; RRID:AB_778898) and mouse anti-Gαs (1:50; catalog #sc-393878, Santa Cruz Biotechnology; RRID:AB_2734138) and the secondary antibodies Alexa Fluor 488 donkey anti-rabbit and Alexa Fluor 594 donkey anti-mouse (1:500; catalog #711-545-152 and #715-585-150, Jackson ImmunoResearch; RRID:AB_2313584 and AB_2340854, respectively). Twenty-four hours before recording, 5 × 10^5 cells were plated on glass-bottom dishes coated with a mixture of poly-D-lysine, laminin, and collagen (Sigma-Aldrich). Three to four hours after plating, we transfected cells with the fluorescent calcium reporter pGp-CMV-NE-StRgaMP1b (Addgene; Dana et al., 2016). Before recording, the media were changed to recording media containing DMEM supplemented with 100 μM penicillin, 100 μg/mL streptomycin, 10 mM Hepes, 0.35 g/L NaHCO3, 4.5 g/L d-glucose, 2% 2-glutamine, and 1% B27 cell culture supplement (Thermo Fisher Scientific). To determine the responsibility and specificity of the reporter cell line, 1.1 × 10^4 (10 μM) or 1.0 × 10^5 (100 nM) gastrin-releasing peptide (GRP; Sigma-Aldrich) diluted in recording media was pipetted directly above the plated cells and imaged.

**Organotypic SCN culture and in vitro VIP sensor recording.** SCN slices were obtained using methods modified from the study by Abe et al. (2002). Mice were anesthetized with CO2 and decapitated. Brains were quickly removed and placed in cold oxygenated HBSS. Coronal sections (250 μm) containing the SCN were obtained using a vibratome (catalog #OTIS-5000, Electron Microscopy Sciences). SCNs were dissected and cultured individually on cell culture membranes (catalog #PICM0RG50, Millipore), which were then placed into a 35 mm Petri dish (Falcon) with 1 mL of recording medium, sealed with vacuum grease, and incubated at 37°C overnight. The following day, culture membranes containing the SCN were transferred to dishes containing cultured VIP sensor cells. We extrapolated subjective time of day in vivo from the prior light cycle. For time-lapse recordings, dishes were placed on an inverted microscope (Eclipse TE2000-S, Nikon) within an incubation chamber (In Vivo Scientific) held at 25°C. VIP sensor fluorescence was imaged with a 10x objective focused on the reporter cells, a TRITC filter cube (Chroma), and a digital EMCCD camera (iXon DU-497, Andor Technology), recorded at a frame rate of 1 frame/s for 24 h using a custom μManager script (http://micro-manager.org; RRID:SCR_000415). Recordings were stopped after 24 h before reporter cells reached confluence. For acute recordings, VIP sensor cells were prepared as above and kept at 25°C until they were imaged with SCN slices taken at either 11:00 AM to 1:00 PM (circadian time (CT)) or 6:00 to 7:00 PM (CT 12) the previous day. We imaged VIP sensor fluorescence for 5–8 min using a TRITC filter cube and a digital CCD camera (QIClick, QImaging) at 1 frame/s with QCapture (QImaging) and GemiStudio software. We then pipetted 1 μL of 1 or 10 nM NMDA (Sigma-Aldrich) directly onto the SCN slice and recorded for an additional 8–10 min. In all experiments, dishes were excluded from analysis if the VIP sensors did not respond to cognate VIP application.

**VIP sensor analysis.** Frames were extracted from movie files using Mlab and opened as TIFF stacks in ImageJ (http://imagej.nih.gov; RRID:SCR_003070) for analysis. Recordings were bleach corrected using an exponential fit and background subtracted using a rolling ball algorithm to diminish SCN autofluorescence. Regions of interest (ROIs) were selected using an SD projection and a diameter threshold of 5–30 pixels. The mean pixel intensity values for each ROI calculated for each frame were then processed in Matlab by first dividing by the mean pixel value of the entire frame to generate a background-corrected pixel intensity value. The ΔF/F value for each ROI for a given frame n was then calculated as (corrected pixel intensity value for frame n – median pixel intensity values across all frames)/(median pixel intensity values across all frames). The resulting ΔF/F values were smoothed by a 5 s moving average filter. For recordings of spontaneous activity, as there was no prestimulus or poststimulus epoch, data were analyzed by integrating the single-cell ΔF/F over each hourly 10 min recording. This single-cell integrated fluorescence was summed and divided by the number of cells analyzed in that hour to account for reporter cell proliferation to produce an average integrated fluorescence per cell. For evoked responses, the maximum ΔF/F for each cell was calculated before and after NMDA stimulation.

In vivo VIP neuron inhibition. After recovery, mice were transferred to wheel-running cages and housed in DD in a light–dark, temperature- and humidity-controlled chamber with food and water provided ad libitum. After allowing the mice to run free for multiple days, we injected them with doxapine-N-oxide (CNO; 1 mg/kg i.p.; Hello Bio) 30 min before activity onset (Grippo et al., 2017). Mice were returned to their cages and allowed to run free for multiple days and then were injected again with CNO at CT 11.5, 30 min before a 15 min light pulse [bright (ranging from 5 × 10^16 to 2 × 10^18 photons/cm^2 throughout the cage) or dim (≈4 × 10^18 photons/cm^2) light pulse], returned to their cages and allowed to free run for multiple days. Phase shifts were determined by using activity onset analysis in Clocklab, using at least 4 d of data per condition (baseline, after CNO, and after CNO + light pulse). The day of drug administration was excluded from analysis. A regression line was fit through activity onsets and extrapolated before and after each condition; phase advancement was quantified by integrating positive slope lines extrapolated to the days of drug administration (i.e., the time difference between the predicted and actual phase onsets). To determine light-evoked activation of VIP neurons, mice were injected with CNO at CT 11.5, received a 15 min bright-light pulse at CT 12, returned to DD, and perfused 1 h after the start of the light pulse, as described below.

**Histology and image analysis.** Mice were deeply anesthetized with isoflurane (>5%) and transcardially perfused with PBS followed by 4% paraformaldehyde. Brains were extracted, postfixixed overnight in 4% paraformaldehyde, and cryoprotected in 30% sucrose. Coronal sections (40 μm) containing the SCN were collected using a cryostat (Leica) and stored at −4°C in PBS. Immunostaining was performed as previously described (Grippo et al., 2017; Tso et al., 2017). The primary antibodies used were rabbit anti-mCherry (1:1000; catalog #ab167433, Abcam; RRID:AB_2571870) and goat anti-c-Fos (1:1000; catalog #sc-52-G, Santa Cruz Biotechnology; RRID:AB_2629503). Secondary antibodies used were DyLight 594 donkey anti-rabbit (1:500; catalog #SA5–10040, Thermo Fisher Scientific; RRID:AB_2556666), and DyLight 647 donkey anti-goat (1:500; catalog #SA460175, Sigma-Aldrich; RRID:AB_2734139). The 2048 × 2048 pixel images of the SCN were acquired on a Nikon Ai Confocal Microscope as a single z-stack.
Figure 1. SCN VIP neurons exhibit daily rhythms in spontaneous calcium activity in vivo. 
a. Fiber photometry experimental timeline and schematic. 

b. Representative image of a fiber-optic cannula (white dashed lines) positioned above a GCaMP6s 

SCN. OC, optic chiasm; 3V, third ventricle. Scale bar, 200 μm. 
c. Schematic of the two-color excitation system used to measure 
calcium-evoked (470 nm) or calcium-independent (405 nm) fluorescence. 
d. Illustration of calcium event detection. Blue, Raw 470 nm signal; purple, raw 405 nm signal; yellow, detrended 405 nm 
signal; green, ΔF/F; dashed line, event threshold value; red circles, events. 
e. Hourly ΔF/F traces taken from VIP neurons in transgenic mice with GCaMP6s or without EGFP a fluorescent calcium 
reporter in a daily LD cycle (yellow = light phase) or DD (light gray = subjective day). 
g. Double-plotted histogram of normalized event frequency per hour (green) and normalized locomotor activity 
counts per hour (black) recorded from an individual GCaMP6s 
mouse housed in LD (gray and yellow shading) and DD (gray shading). A 10 min light pulse (yellow) (Figure legend continues.)
(optical section thickness, 2 μm) using identical capture settings for each image, which was to be quantitatively compared. For quantitative c-Fos analysis, sections were z projected by their average intensity using ImageJ software. Regions of interest were selected automatically in ImageJ using Shanbhag thresholding with identical settings for each section followed by particle analysis with size and circularity thresholds of ≥200 pixels^2 and ≥0.1, respectively. Brightness and contrast settings were adjusted identically for sections depicted in representative images.

**Statistics.** Sample sizes were chosen to be sufficient for statistical analysis based upon similar techniques used in previous publications (Ehrlich-Rothschild et al., 2016). Data analysis was performed blind to genotype or condition. There were no methods to randomize mice to experimental groups or to blind investigators to genotype or condition during the experiments. All statistical analyses including one-way ANOVA, two-way ANOVA, one-way repeated-measures ANOVA, two-way repeated-measures ANOVA, Student's t test, Tukey’s multiple-comparison test (MCT), and Sidak’s MCT were performed in Matlab R2016a or Prism 7 (GraphPad; RRID:SCR_002798) with α defined as 0.05. Circadian rhythmicity was determined using cosinor analysis (Refinetti et al., 2007) in Matlab and independently using the TSK Cycle (Hughes et al., 2010) in R (RRID:SCR_001905). Shapiro–Wilks and Brown–Forsythe tests were used to test for normality and equal variances. Data are presented as the mean ± SEM.

**Results.**

SCN VIP neurons exhibit daily rhythms in spontaneous calcium activity in vivo

To determine the circadian regulation of SCN VIP neuron activity over multiple days, we used long-term in vivo fiber photometry to record hourly spontaneous calcium activity from the SCNs of freely moving mice (Fig. 1a–d). We injected a Cre-dependent AAV encoding the fluorescent calcium sensor GCaMP6s (AAV9-CAG-Flex-GCaMP6s) or a control fluorophore (AAV9-CAG-Flex-EGFP) into the SCNs of VIP-ires-Cre knock-in mice. We found daily rhythms in intracellular calcium levels and event frequency that persisted for as many days as we recorded, peaking during the daily minimum of locomotor activity in an LD cycle or in DD (Fig. 1e–j). The amplitude of the rhythm (LD day, 1.31 ± 0.38 events/min; LD night, 0.09 ± 0.15 events/min; DD subje-
Figure 3. VIP sensor cells reliably detect VIP and reveal day-night differences in acute VIP release from the SCN. a, Experimental outline and VIP sensor schematic. b, jRCaMP1b calcium fluorescence in VIP reporter cells stably transfected with jRCaMP1 + VPAC2R increases in the presence of VIP, but not of GRP (n = 189, 147 cells). c, VIP sensors exhibit a dose-dependent response to VIP concentration but do not respond to GRP or in the absence of VIPAC2R (n = 750 cells, 4 wells/group; one-way ANOVA: F(2, 19) = 242.2, p < 0.0001; with post hoc Tukey’s HSD). Error bars depict the mean ± SEM. Letters indicate statistically different groups. d, Representative 1 min binned average raw pixel intensity and average ΔF/F images of VIP sensors cultured under a acute SCN slice (top) or in the same dish but away from the SCN (bottom). White dashed outline delineates the boundary of the SCN. 3V, Third ventricle; OC, optic chiasm. e, ΔF/F traces of the cells depicted in d. Colors represent individual reporter cells; triangles designate cells indicated by white arrows in d. f, Representative heatmaps of ΔF/F responses from VIP sensors cultured under acute SCN slices explanted during the subjective day or night.

SCN VIP neurons exhibit daily rhythms in light-evoked calcium activity in vivo

We next tested how SCN VIP neurons responded to light around subjective dusk when light is known to delay daily rhythms in behavior. We found that a 10 min light pulse reliably increased intracellular calcium (Fig. 2a). We therefore tested the effects of 15 a light pulses at different times of day and found that calcium responses exceeded baseline at CTs 12, 18, or 24, but not during the middle of the subjective day (CT 6; Fig. 2b,c). These responses lasted multiple tens of seconds after the cessation of the light pulse, with daily peaks in amplitude and duration occurring at approximately CT 12 (Fig. 2d,e) and increased in amplitude with increasing light intensity (Fig. 2f,g). These results suggest that SCN VIP neurons exhibit in vivo daily rhythms in light-evoked calcium activity and can encode light intensity in a manner consistent with light-induced circadian phase shifts (Nelson and Takahashi, 1991).

SCN slices exhibit daily rhythms in spontaneous and evoked VIP release

To determine how SCN VIP neuron calcium activity correlates with the release of VIP, we engineered HEK293 reporter cells to selectively fluoresce in response to VIP (Fig. 3a). These cells reported VIP concentration in a dose-dependent manner (Fig. 3b,c) and fluoresced when cultured under acute SCN slices, but not when in the same dish remote from the SCN (Fig. 3d,e). VIP sensor cells exhibited more spontaneous calcium activity when placed under acute wild-type SCN slices during the subjective day than during the subjective night (Fig. 3f). Spontaneous calcium activity was circadian when recorded continuously under cultured wild-type SCN slices, but was arrhythmic and low amplitude when recorded under VIP−/− slices (Fig. 4a-c). Adding 1 or 10 mM NMDA to acute wild-type SCN slices at CT 12 to mimic light input (Ding et al., 1994; Mintz et al., 1999) also increased calcium activity in VIP sensors (Fig. 4d,e). Consistent with the inability of daytime light to shift the SCN, adding 1 mM NMDA to the SCN at CT 6 did not change VIP sensor fluorescence. Importantly, adding NMDA to reporter cells alone or to a VIP−/− SCN slice at CT 12 did not change reporter cell fluorescence. Intriguingly, 10 mM NMDA during the subjective day reduced VIP sensor fluorescence (which could be restored by adding VIP). These results collectively suggest that there are daily rhythms in spontaneous and NMDA-evoked VIP release from the SCN, consistent with a role for VIP in mediating circadian-, light-, and NMDA-induced shifts of the SCN and behavior.

SCN VIP neuron activity is necessary for normal light-induced phase shifts

Finally, to determine the contribution of SCN VIP neurons to the encoding of light that shifts daily rhythms, we used chemogenet-
ics to acutely silence these neurons in freely behaving mice. We recorded circadian locomotor behavior from VIP-ires-Cre mice injected with a SCN-targeted, Cre-dependent AAV encoding an engineered inhibitory G<i><i></i></i>, coupled receptor (AAV8-hSyn-DIO-
hM4Di) or a control fluorophore (AAV9-CAG-Flex EGFP; Fig. 5a). We then silenced VIP neurons around the daily time of locomotor onset (CT 12), when in vivo VIP neuron calcium activity is decreasing and maximally augmentable by light exposure. We found that intraperitoneal injections of the hM4Di-specific ligand CNO at CT 11.5 attenuated, but did not fully block light induction of c-FOS, an indicator of neuronal activation, in the SCN (Fig. 5b,c). CNO alone did not shift daily rhythms in locomotor activity in experimental or control mice (Fig. 5d), indicating that VIP neuron activity is not required at CT 12 for circadian rhythms to progress. Strikingly similar to its effects on photic induction of c-FOS, injection of CNO 30 min before a 15 min light pulse at CT 12 greatly attenuated the resulting phase delay from 2.10 ± 0.21 to 0.88 ± 0.07 h compared with control animals (Fig. 5d,e). Finally, CNO also reduced phase delays of circadian behavior after light pulses that were 10× dimmer (0.39 ± 0.04 h; compared with shifts of CNO-treated control mice, 0.91 ± 0.04 h; Fig. 5f,g). These results suggest that the depolarization of SCN VIP neurons is necessary for normal light-induced phase shifts.

Discussion

Together, these experiments demonstrate that SCN VIP neurons exhibit circadian rhythms in both spontaneous and light-evoked neuronal activity in vivo and in circadian and NMDA-evoked VIP release in vitro. Previous studies have reported circadian rhythms in basal calcium levels of VIP neurons in the isolated SCN in vitro (Enoki et al., 2017) and, recently, in vivo (Mei et al., 2018), but have differed on whether their firing rate is circadian (Webb et al., 2009; Fan et al., 2015; Hermanstyne et al., 2016). Here we provide the first evidence of the circadian regulation of calcium spikes in SCN VIP neurons, which likely relate to neuronal action potentials (Chen et al., 2013; Gunaydin et al., 2014). The regulation of these fast calcium events may be separable from the regulation of basal calcium levels under certain conditions (Ikeda et al., 2003; Enoki et al., 2012). Our results are consistent with light- and circadian-evoked VIP release from the SCN (Shinohara et al., 1995, 1999) which has not been not been detectable in vivo (Frands et al., 2010). Similarly, our results suggest that physiological levels of NMDA induce VIP release by elevating neuronal activity at times of day when neuronal activity is low but not when neuronal activity is already high, which is consistent with a maximal effect of NMDA application on SCN phase shifts (Ding et al., 1994) and on SCN calcium transients (Colwell, 2001) occurring during the
subjective night. High concentrations of NMDA, however, likely reduce VIP release during the subjective day by reducing elevated VIP neuron firing through depolarization block (Meijer et al., 1993).

Intriguingly, we also find that the in vivo circadian activity of VIP neurons rapidly becomes high and arrhythmic after acute exposure to constant light. This is consistent with results showing that chronic exposure to constant light increases levels of VIP (An et al., 2012) and disrupts the day-night difference in basal calcium levels in an in vitro SCN preparation (Colwell, 2000), but at apparent odds with the sustained circadian locomotor activity for multiple days in LL. We therefore postulate that aberrant lighting can elevate activity in SCN VIP neurons (some of which project to extra-SCN brain regions; Abrahamson and Moore, 2001) without immediate effects on circadian behavior, but with potential effects on behaviors such as mood (J. Gates et al., 2012).

We found that silencing SCN VIP neurons attenuate, but do not block, light-induced shifts in circadian behavior. Although nocturnal light upregulates the markers of activation in VIP neurons (Romijn et al., 1996; Kuhlman et al., 2003), our results indicate that all photic signaling does not pass through VIP neurons to entrain rhythms in the SCN and behavior. This is surprising because shifts following optogenetic stimulation of most SCN neurons in vitro can be blocked with VIP receptor antagonists (Jones et al., 2015), and blocking the acute induction of the core clock gene Per1, which is preferentially induced in VIP neurons in response to a light pulse (Kuhlman et al., 2003), greatly attenuates light-induced behavioral phase shifts in vivo (Akiyama et al., 1999; Tischkau et al., 2003). Our results are, however, consistent with the observation that VIP-deficient mice can still synchronize to a light cycle (although with an 8 h advanced-phase angle of entrainment; Colwell et al., 2003), and that phase response curves to VIP and light differ in their shape (for review, see An et al., 2011). It is possible that genetic deletion of VIP early in development allows for compensation and that our chemogenetic manipulations did not fully silence all VIP neurons. However, because retinal ganglion cells make contacts on numerous cell types throughout the SCN (Fernandez et al., 2016) and light induces c-FOS broadly in the SCN (Castel et al., 1997), we posit that
VIP neurons are a major, but not a unique, modulator of light entrainment of the SCN and behavior.

Interestingly, we found higher and longer light-induced VIP neuron calcium responses at subjective dusk compared with spontaneous VIP neuron calcium levels at any time, suggesting that light during the subjective night causes an increase in VIP neuronal activity, and, consequently, VIP release, that is above the normal daytime spontaneous activity levels occurring in these neurons. Because our results indicate that light activation of retinal afferents produces about half of the normal phase shift to SCN cells when SCN VIP neurons are silenced, we propose that light at night activates VIP and non-VIP SCN neurons. We predict that VIP release by VIP neurons enhances the non-VIP neuron response to light consistent with the antagonism or deletion of VIP signaling attenuating light induction of Per1 and c-FOS in both the VIP-enriched ventral SCN and VIP-deficient dorsal SCN (Vosko et al., 2015). Based on the in vitro phase-response curve to VIP (An et al., 2011) and on our observations that these neurons maximally respond to light at dusk, this VIP neuron-mediated phase shifting likely predominantly involves phase delays. Thus, we conclude that SCN VIP neurons show daily rhythms in spontaneous neuronal activity with peak VIP release around mid-day, and that light at night augments their relatively low activity and VIP release to amplify shifts of the daily rhythm in the SCN and behavior.

References


