Dissecting and modeling photic and melanopsin effects to predict sleep disturbances induced by irregular light exposure in mice

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Abstract

Artificial lighting, day-length changes, shift work, and transmigrational travel all lead to sleep–wake disturbances. The rhythymethemmelanopsin sleep–wake cycle (SWc) is known to be controlled by output from the central circadian clock in the suprachiasmatic nuclei (SCN), which is entrained to the light–dark cycle. Additionally, intrinsically photosensitive retinal ganglion cells containing the photopigment melanopsin (Opn4), short-term light–dark alternations exert direct and acute influences on sleep and waking. However, the extent to which longer exposures typically experienced across the 24-h day exert such an effect has never been clarified or quantified, as disentangling sustained direct light effects (SDLE) from circadian effects is difficult. Recording sleep in mice lacking a circadian pacemaker, either through transgenesis (S5t1cre/m8a1fr) or SCN lesioning and/or melanopsin-based phototransduction (Opn4<sup>-/-</sup>), we uncovered, contrary to prevailing assumptions, that the contribution of SDLE is as important as circadian-driven input in determining SWc amplitude. Specifically, SDLE were primarily mediated (>80%) through melanopsin, of which half were then relayed through the SCN, revealing an ancillary purpose for this structure, independent of its clock function in organizing SWc. Based on these findings, we designed a model to estimate the effect of atypical light–dark cycles on SWc. This model predicted SWc amplitude in mice exposed to simulated transequatorial or transmeridian paradigms. Taken together, we demonstrate this SDLE is a crucial mechanism influencing behavior on par with the circadian system. In a broader context, these findings mandate considering SDLE, in addition to circadian drive, for coping with health consequences of atypical light exposure in our society.

Significance

Light pervasively affects physiology and behavior, especially the sleep–wake cycle (SWc), which is generated by clock signaling in the suprachiasmatic nuclei (SCN). Light synchronizes this clock and additionally exerts direct (circadian-independent) acute effects on sleep. However, it has never been clarified whether longer photic exposure, as experienced daily, might exert long-term sustained direct light effects (SDLE). Recording sleep in mice lacking a functional clock and/or the retinal photopigment melanopsin, we uncovered that circadian-driven input and SDLE equally contribute to SWc. SDLE were primarily mediated through melanopsin, with half relayed through SCN, redefining the latter’s role. Modeling allowed SWc prediction under simulated jet lags, further demonstrating SDLE as an important mechanism to consider when managing health consequences of light exposure.

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are thought to mediate light influence on physiology and behavior (13–15). Clock synchronization and circadian rhythm entrainment is mediated by projections to the SCN (16–18), whereas innervation of the sleep-promoting neurons of the ventrolateral preoptic area (VLPO) (19, 20) or the subparaventricular zone are thought to mediate light’s direct effects on sleep, yet it remains to be clarified whether the SCN might also participate in mediating those direct photic effects. Moreover, the loss of melanopsin does not abolish these responses, demonstrating that other photoreceptors involved in vision, the rods and cones, also play a role (12). Furthermore, the loss of ipRGCs results in dramatic loss of nonvisual effects, indicating that these cells receive inputs from rods and cones and are also the principal conduit for nonvisual light input from these photoreceptors (12, 18, 21–23). However, the degree of overlap between these two systems for regulating nonvisual responses is not yet certain.

In recent years, the development of transgenic mouse models targeting these phototransduction pathways has revealed the pronounced effects of light and dark pulses of short duration on sleep and waking (11, 12, 24). Moreover, we previously showed that this acute and direct photic influence can be observed at all times of day (12). These observations raise the question as to whether longer periods of light and dark exposure, such as those of the 24-h LD cycle, exert sustained direct noncircadian effects, that is, continuous acute effects that could be observed over longer periods of time. One could postulate that the complete absence of light (constant darkness [DD] experiments) would allow for the extrapolation of the influence of sustained direct photic effects; however, this would not consider the possibility that the absence of light would lead to circadian disruption. Thus, under standard LD cycles, the contribution of direct noncircadian photic regulation in shaping the sleep–wake cycle has to date never been properly quantified due to the difficulty in disentangling it from the circadian effects.

In the current study, we sought to address three related issues: 1) the quantification of the respective contribution of circadian effects (CE) and sustained direct light effects (SDL) in shaping the 24-h sleep–wake cycle; 2) the identification of the pathways underlying those effects (i.e., the respective role of the photoreceptive systems), melanopsin versus rods and cones, and the respective role of the SCN versus non-SCN brain relays as conduits for these direct photic effects; and 3) whether quantifying the respective contributions of CE and SDL allows for the prediction of the nychthemeral sleep–wake distribution under unnatural LD cycles, such as those experienced during jet lag. Recording sleep in mice lacking a functional central clock either through SCN lesioning or Bmal1 tissue-specific clock gene deletion and/or melanopsin-based phototransduction, we uncovered that CE and SDL contribute in equal proportion to shaping the nychthemeral cycle. Furthermore, analyzing sleep in melanopsin-deficient mice with or without disabled SCN revealed that SDL are primarily (>80%) mediated through melanopsin-based phototransduction, half of which passes via the SCN, implying a noncircadian function for the brain structure comprising the master circadian clock. To further validate our findings, these data were integrated into a model that accurately predicted 24-h sleep–wake distribution in mice exposed to a simulated “jet lag” or transequatorial travel paradigm.

**Results**

**Nychthemeral Sleep–Wake Cycle Amplitude Is Decreased by Half in the Absence of Light or when Light Is Equally Distributed Across 24 h.**

To estimate overall light influence on the 24-h sleep–wake cycle amplitude, we recorded sleep electrocorticograms (ECoG) in animals in the absence of light or when light is equally distributed across 24 h. Here, and below, we defined the nychthemeral sleep–wake cycle (SWC) amplitude as the difference in non-rapid eye movement sleep (NREMS) amounts between 12-h light (day) and 12-h dark (night) periods. Similar analyses were performed using waking or rapid eye movement sleep (REMS) for calculating SWC amplitude, leading to comparable observations. Of note, hourly NREMS amounts across the 24-h period follow a more square-like distribution, therefore taking the 12-h LD difference is appropriate as an amplitude estimate. Furthermore, under constant darkness, the term “subjective light” defines the ZT0–12 period corresponding to the 12-h light phase of the preceding days, and the term “subjective dark” corresponds to the ZT12–24 period, that is, the 12-h dark phase of the preceding days, given that a circadian phase shift would be negligible during a single day of DD. Under DD, we found that SWC amplitude was reduced by ~1 h (Fig. 1 B and C and SI Appendix, Fig. S1C), representing 45% when compared with LD12:12, indicating that the 55% remaining was generated by the circadian drive. Another way to estimate the overall effects of light is to record sleep and waking under ultradian light/dark cycles, in which light and dark pulses are evenly distributed throughout the nychthemeron with pulse-induced effects on sleep superimposing on the circadian modulation. Thus, mice were exposed to consecutive 1-h light and 1-h dark pulses (LD1:1), which we previously used to analyze the acute direct effects of light according to the time of day (12). Next, a group of animals was exposed to consecutive pulses consisting of 3.5 h light and 3.5 h dark (LD3.5:3.5), an alternative condition in which L/D pulses were not aligned to 24 h (24, 25). Under LD1:1, LD3.5:3.5, and DD, NREMS amounts were noticeably different (SI Appendix, Fig. S1 A–C). With these two ultradian LD cycles, we assumed that the direct effects of alternating light and dark pulses nullified one another over time and that the residual SWC amplitude represented the remaining circadian influence. Remarkably, the circadian contribution under these two ultradian LD regimes were found to be within the same range as was calculated previously under LD12:12 (LD1:1, 50 ± 11% and LD3.5:3.5, 52 ± 7% versus 55 ± 6% in DD; SI Appendix, Fig. S1D). Thus, this decrease by half of the SWC amplitude demonstrates a crucial role for the direct influence of light in driving the nychthemeral sleep–wake distribution and further suggests that it can be sustained for longer periods of time, with acute and sustained DLE having very similar effects. However, under those conditions, it cannot be ruled out that the circadian system is not altered in the absence of light or under alternating light signals. Therefore, this underlines that the reduction of SWC amplitude observed here may also result from an alteration of CE, a question that our step-by-step approach below addresses.

**Fig. 1.** Melanopsin-based SDL contributes to approximately one-third of the SWC amplitude. (A) Dynamics of the accumulated differences demonstrate that Opn4+/− mice lose 2 h of NREMS per day, with a maximum effect (1.5 h) during the light period (average of 2 consecutive days under LD12:12). (B) NREMS amount during light period under LD12:12 (Left) is decreased in Opn4+/− and similar to that during subjective light period under DD in both genotypes (Right). (C) SWC amplitude in Opn4+/− and Opn4−/− mice under DD did not differ. These observations demonstrate that sleep loss during the light period and subsequent SWC amplitude decreases in Opn4−/− mice result from a lack of Opn4-based SDL and not from a reduction in circadian signal. DD: sham Opn4+/− n = 5; sham Opn4−/− n = 7. Asterisks denote significant post hoc differences after ANOVA between LD conditions (blue) and the genotype (red).
Melanopsin has been shown to be the primary mediator of the acute direct influence of light on sleep and waking, with rods and cones playing only a minor role (for review, see ref. 7). To determine whether melanopsin-based direct photic influence would be sustained for longer periods, therefore contributing to the SWc, we assessed sleep in melanopsin-deficient mice (Opn4−/−). Under LD12:12, mice lacking melanopsin, in comparison with wild-type (WT) animals, displayed a 1.5-h reduction in NREMS, similar to what we previously observed in these mice (12). This NREMS decrease constantly accumulated across the entirety of the 12-h light phase (Fig. 1A). The reduction in NREMS also continued during the dark period but to a lesser extent (0.5 h), thereby reducing SWc amplitude and resulting in a 37% decrease compared with WT controls (Opn4+/+). In the context of SDLE, we interpreted this as a lack of melanopsin-driven input, assuming that CE were not impacted in these mice by a life-long loss of Opn4. We verified this latter point by confirming SWc amplitude under DD did not differ between mice either expressing melanopsin or not (Fig. 1B). Moreover, the accumulated NREMS loss during the 12-h light phase in Opn4−/− mice (Fig. 1A) was similar to the NREMS reduction observed during the subjective light period (CT0-12) under DD in both WT and Opn4+/− mice (Fig. 1C). Therefore, NREMS reduction in these animals resulted not from a circadian defect but a lack of direct photic influence, identifying that one-third (37%) of SWc amplitude was driven by Opn4-mediated SDLE.

One-Quarter of the Nychthemeral SWc Is Determined by SCN-Independent Sustained Direct Light Effects. The most straightforward method to eliminate the influence of CE on the sleep–wake distribution is to disable the central circadian pacemaker (SCN). To calculate the proportion of the SWc amplitude generated by SCN-signaling, we studied two groups of mice in which this structure had been either physically removed through thermolytic lesioning (SCNx) or rendered inert through the deletion of the clock gene Bmal1 in the mouse brain, as well as in the SCN (Syt10Cre/+). We verified this latter point by confirming SWc amplitude in animals lacking melanopsin (Opn4−/−) with or without a functional SCN (SCNx), respectively (Fig. 2D). Of note, about one-quarter of the SWc amplitude was conserved in these two models. This indicates that light exerts a sustained direct noncircadian effect, strong enough to uphold a 24-h organization of sleep and waking in the absence of SCN-generated circadian rhythms and that SCN-driven signaling is responsible for three-quarters of the SWc amplitude observed under LD12:12.

The SCN Acts Not Only as a Clock but Mediates Half of the Sustained Direct Alerting Effects. Thus far, we demonstrated that 77% of the SWc amplitude depends on SCN output, and we could therefore assume this to be the result of its inherent clock function. However, we could not rule out that part of this SCN-dependent SWc amplitude arose from direct effects of light mediated through the SCN independently of clock mechanisms. To separate melanopsin-based SDLE mediated through the SCN from pathways external to these nuclei, we examined SWc amplitude in animals lacking melanopsin photopigment (Opn4−/−) with or without a functional SCN (SCNs), which demonstrated that Opn4-based phototransduction accounted for 18% of the SWc amplitude through pathways outside of the SCN and 19% within (Fig. 3A and B). Thus, our observations confirmed that nearly half (49%) of SDLE were mediated through SCN pathways, identifying that this structure, beyond its clock function, significantly influenced vigilance states (Fig. 3C).

Interestingly, the flattened SWc amplitude observed in SCN-ablated mice resulted mainly from a decrease in waking during the dark phase. Indeed, in nocturnal animals such as mice, darkness is known to exert a wake-promoting effect, whereas in humans, light has been shown to acutely promote alertness, with the highest efficiency of blue light centered around 480 nm, within the spectral response peak of melanopsin (4, 30). Therefore, we calculated...
the power spectral density of the ECoG to quantify theta (7 to 10 Hz) and gamma (40 to 70 Hz) activities, known to be correlates of exploratory behavior and alertness (31–33). Remarkably, both were significantly decreased in each of the SCN-deficient models, specifically during the 12-h dark period of LD12:12 (SI Appendix, Fig. S3), indicating that the alerting effect of darkness in nocturnal animals (inverted in diurnal species) is primarily signaled though the SCN.

**SDLE and CE Equally Contribute to Shaping the 24h SWc.** Although we have shown that SDLE were primarily mediated by Opn4- dependent pathways, a possible contribution of rods- and cones- based photoreception could not be excluded. To address this, we calculated SWc amplitude in SCNx Opn4−/− mice, giving us an estimate of rod/cone-based SDLE mediated outside the SCN, which was found to be low (~5%; Fig. 3B). However, these values were not significantly different from zero (one-sample signed rank test, P = 0.22), signifying that their contribution is likely negligible. Summarizing these findings indicated that 42% of the SWc amplitude was determined by SDLE, that is, the SDLE contribution we previously calculated in the absence of light under DD in WT animals (45%), and we therefore attributed this 3% difference to a possible contribution of SCN-dependent rod/cone-based phototransduction (Fig. 3C). This confirms the results from our initial DD experiment on the role of light in determining the SWc and further demonstrates that optimal functioning of SDLE requires both melanopsin-based phototransduction and a functional SCN in addition to other brain relays such as the VLPO.

These results encompass different photic and circadian regulatory factors (CE versus SDLE, Opn4 versus rods/cones, and SCN versus non-SCN-mediated transmission), leading us to conclude that SDLE and CE are nearly equal in influencing 24-h sleep-wake distribution. This estimate was corroborated by analysis of WT mice under the previously mentioned ultradian cycles in which light and dark pulses were evenly distributed over the nychthemeral cycle, indicating approximate SWc amplitude values of 50 and 48% for LD1:1 and LD3:5:3.5, respectively (SI Appendix, Fig. S1). Finally, regarding other vigilance states, the contributions of CE and SDLE to REMS distribution were similar to those we calculated for NREMS (SI Appendix, Fig. S4). Overall, the remarkable coherence of our results across experimental conditions strengthens the robustness of our observations on an equal
direct contribution of CE and SDLE as well as on the respective role of the different underlying pathways.

We summarized our findings in a model presented in Fig. 3 C and D, integrating the respective contributions of the main phototransduction pathways regulating the natural SWc through use of SWc amplitude as a proxy. Briefly, the role of each component was calculated based upon intergroup differences of this measure as the degree of SWc amplitude reduction is dependent on the phototransduction element being removed in each. The contribution of the SCN (combined SDLE and CE) was estimated through the SWc amplitude difference between SCN-disabled mice and controls. In addition, the contribution of melanopsin-dependent SDLE was calculated based on the SWc amplitude reduction observed in melanopsin-deficient mice (given that CE were conserved in these mice). The percentage mediated through the SCN was estimated in SCNx animals with the difference between those either possessing or lacking melanopsin. These two latter calculations allowed us to derive the proportion of Opn4-mediated SDLE mediated through the SCN. The removal of both melanopsin and SCN (SCNx Opn4+/−) gave us the contribution of the rods/cones-dependent SDLE mediated through brain relays outside of the SCN (as it was the only pathway conserved), whereas the one mediated through the SCN was negligible. Finally, the calculations under DD, LD1:1, LD3.5:3.5, and from our intergroup comparisons under LD12:12 converged toward an estimation of the rods/cones-dependent SDLE mediated through brain relays outside of the SCN. The percentage mediated through the SCN was estimated in SCNx animals with the difference between those either possessing or lacking melanopsin.

Modeling Predicts 24-h Sleep–Wake Distribution under Simulated Jet Lag. We then integrated CE and SDLE contributions (55 and 45%, respectively) into a model aimed at estimating whether these respective effects could estimate overall sleep–wake distribution under other light/dark paradigms. As this was the first time SWc amplitude had been used to derive such a model, we chose two light/dark schedules simulating either transequatorial (i.e., rapid seasonal change) or transmeridian (intercontinental) travel, conditions commonly experienced in modern society. In WT mice exposed to a shortened photoperiod (transequatorial; LD8:16), we found the overall model estimation and shape of the nychthemeral SWc to be similar to values obtained from ECoG recordings (Fig. 4 A and B).

Next, a separate group of mice was subjected to a simulated westward transmeridian travel with an 8-h prolongation of the dark period followed by a shifted LD12:12 cycle (Fig. 4 C, Top). Whereas in the first experiment we assumed the SCN output not to be affected, this condition included an additional factor, as shifting the LD cycle induces a phase adjustment of the clock to
the new LD cycle (i.e., jet lag). Indeed, sleep recordings under this condition showed that resynchronization to the new LD12:12 cycle took ~96 h (Fig. 4C). This same time delay was observed when evaluating locomotor activity patterns, which fully resynchronized by day 5 following the jet lag (for example, see SI Appendix, Fig. S5). Therefore, we assumed stable entrainment to the new LD cycle across 4 d with a linear shift over this period of time of ~1.10/12 h (derived from a best fit of the observed shift based on NREMS data). To calculate the estimated NREMS time course, hourly values were multiplied by CE (55%) and SDLE (45%) contributions (as in Fig. 3D) and plotted in 3-h bins (Materials and Methods). Remarkably, when integrating this constant (phase shift of CE) into the model, the predicted 24-h NREMS distribution (Fig. 4C, purple line) closely followed the observed changes in NREMS across the entirety of the experiment (11 d; Fig. 4C, red line), although an overall increasing trend in NREMS over the days following resynchronization could not be fully captured by the model. Furthermore, the model predicted the SWc amplitude dynamics, substantiated by linear regression of predicted versus observed values of these changes across subsequent days (Fig. 4D). Indeed, it identified with greatest accuracy the SWc amplitude dynamics, which were initially attenuated as a result from CE and SDLE being out of phase and partially negating one another at certain times of the day before resynchronization (see blue and green lines in Fig. 4C, Bottom). Taken together, these results establish the robustness of this model, further validating our findings of an equal contribution of CE and SDLE in shaping nycthemeral sleep and wake distribution.

Discussion

In this study, we demonstrate, contrary to the prevailing assumption, that only half of the overall changes to SWc amplitude are generated by a circadian signal, with SDLE being an equally important mechanism. Indeed, these findings challenge our current understanding of how photic input influences the amplitude and pattern of the SWc. Furthermore, we show that SDLE is capable of compensating for the absence of clock-driven influence to help maintain sleep–wake distribution aligned with external LD cues.

This study was designed to investigate the sustained effects of light on sleep–wake distribution, but we acknowledge that other factors potentially influence the distribution of vigilance states. In the current study, we focus only on CE and SDLE, as the experiments were conducted under controlled laboratory conditions, which do not account for other environmental factors known to influence sleep and waking. Indeed, the homeostatic process, defined as an accumulation of sleep pressure with time spent awake and its release during sleep, is another key regulator (34). However, our results on SWc amplitude and the derived model were based on 12-h values of sleep/wake states, which likely contributes to its accuracy, with the kinetics of the homeostatic process being much faster than 12-h periods in mice (35), contrary to humans. Another limitation of this study is the lack of female animal inclusion. While it has been shown that young adult female mice display changes in overall sleep and waking behavior compared with males, circadian rhythms and 24-h sleep–wake distribution in animals of this age are unaffected (36). Furthermore, we acknowledge that these mice are recorded in socially isolated conditions, with identical luminance levels across the light period, and one should be aware that the relative contributions of CE and SDLE and resulting SWc amplitude could be influenced by behavior and social interaction such as mating, foraging, or other ecological conditions. Finally, the present quantification of these CE and SDLE contributions in shaping the SWc, and the resulting modeling, was designed based on controlled conditions to provide a unique framework intended to evolve with the integration of additional environmental factors (e.g., ambient temperature, noise, etc.), which may allow a more precise model to be designed and increase the prediction of sleep–wake distribution at smaller time scales, also facilitating its translation to humans and other species.

![Fig. 4. Prediction of nycthemeral NREMS distribution under simulated transequatorial and transmeridian jet lag. (A) Time course of NREMS in WT mice under 48 h of LD12:12 followed by 4 d under LD8:16, simulating a transequatorial travel, with observed values (blue; min/2 h) and predicted values (orange; min/2 h). (B) SWc amplitude from prediction (orange) and from recorded animals (blue) under baseline (first 2 d under LD12:12) followed by 4 d under LDB:16. (C) Time course of NREMS in WT mice under 48 h of LD12:12 followed by 7 d under a simulated 8-h westward “jet lag”. The predicted values of daily NREMS distribution (purple) based on the model integrating a shift of 1.1/12 h described in Fig. 3D, combining CE (blue) and SDLE (green), are shown. Daily NREMS distributions are obtained from sleep recordings (red) and are expressed as min/3 h. SH = shift (jet lag). (D) SWc amplitude under baseline (BL; LD12:12) and simulated 8-h westward and “jet lag” conditions across experimental (EXP) days. A linear regression of SWc amplitude is plotted to illustrate resynchronization to the new LD (R² = 0.99). The values in red obtained from recorded animals perfectly fit the predicted values in purple (one-way repeated measures ANOVA, Pgroup x time = 0.28). The dashed line represents the average baseline value at 100%. All values represent mean ± SEM, LDB:16: n = 7; jet lag: n = 9.](https://doi.org/10.1073/pnas.2017364118)
Our findings clearly redefine the role of the SCN, which should be seen not only as the orchestrator of circadian rhythms but a critical component of the direct photic regulation of alertness. Deciphering the molecular mechanisms behind circadian and direct photic processes within SCN cells will be crucial for our understanding of how environmental signals interplay with endogenous clock-driven mechanisms for optimal physiological and behavioral functioning. Furthermore, considering that numerous advancements in chronobiology have arisen from the interpretation that observed changes in animals with SCN lesions were solely attributed to the removal of the endogenous pacemaker, the present observations should encourage revisiting some of these conclusions (37). In addition, our findings emphasize the role of melanopin-based phototransduction, as compared with rods and cones, in mediating the direct effect of light on sleep–wake behavior. This corroborates current literature on the photic regulation of behavior and identifies melanopin phototransduction signaling as a potentially new pharmacological target for the manipulation of sleep and alertness (7).

This straightforward model emphasizes that optimal functioning requires CE and SDLE to be in phase with one another. Although these data and the proposed model are derived from experiments performed in nocturnal laboratory mice, the conservation of the circadian system and melanopin-based phototransduction across mice and humans allow us to provide a coherent framework for future studies in basic and clinical research. However, one should be aware that the interpretation of an equal contribution of CE and SDLE to shaping the 24-h behavior might differ in humans, as the weight of each of these two factors might be different in humans. Indeed, the present findings provide an explanation for the physiological and behavioral disturbances induced by the exposure to inappropriate lighting commonly observed in today's society, currently ascribed solely to a misalignment of the behavior relative to circadian time (1, 2). This could be used as a basis to optimize societal lighting and minimize light pollution as well as its application in certain medical conditions. Indeed, current therapeutics for circadian rhythm sleep disorders (CRSD) are based upon enhancing circadian re-entrainment of the clock (5, 38). Our present findings, however, suggest taking into account the direct noncircadian effects of light as a key factor of CRSD management, as the insufficient consideration for SDLE possibly participates in the high rate of unsuc-cessful treatment (38). Finally, these observations could be an explanation for the physiological and behavioral disturbances induced by the exposure to inappropriate lighting (1, 2). This corroborates current literature on the photic regulation of alertness (7).

### Materials and Methods

**Animals.** All experiments were performed on young adult male mice using 1) Opm4+/− and WT litterate controls (12) on a C57BL/6 × 129sv genetic background and 2) Syt10Cre/+Bmal1fl/− on a C57BL/6 background (26), animals rendered arrhythmic through a conditional deletion of the clock gene Bmal1 in the SCN using a Syt10 Cre driver (26) (from the Max Planck Institute for Biophysical Chemistry). Briefly, Syt10Cre/+ mice were crossed to a mouse line carrying a conditional allele of Bmal1, allowing Cre-mediated deletion of the exon encoding the BMAL1 basic helix–loop–helix domain (Bmal1fl/fl). To generate experimental animals (Syt10Cre/+Bmal1fl/−), Syt10Cre/+Bmal1fl/− females were crossed to Syt10Cre/+Bmal1fl/+ males. As controls, littermates of the genotypes Syt10Cre/+Bmal1fl/+ were used. To control for both the Cre driver and the heterozygosity of Bmal1, we verified that Syt10Cre/+Bmal1fl/+ and WT mice displayed similar locomotor activity, ECoG, and sleep–wake patterns. All procedures complied with the local and international rules for the Care and Use of Laboratory Animals and were approved by the ethical committees for animal research (protocol no. B6748225 registered at the French Research Ministry). The animals were maintained under environmentally stable conditions (LD12:12; 25 ± 0.5 °C, food and water ad libitum). The experimenters were blind to the lesion generator. SCNx and sham control mice were then implanted with ECoG and electromyography (EMG) electrodes (see below) for sleep recording. ECoG implantation. Before undergoing implantation, mice were anesthetized with an intraperitoneal injection of ketamine 80 mg/kg and xylazine 7 mg/kg. Male Opm4−/− (n = 16), Opm4+/− (n = 19), Syt10Cre/+Bmal1fl/− (n = 10), Syt10Cre/+Bmal1fl/+ (n = 7), and WT (n = 7) mice were implanted with a classical set of electrodes (two ECoG, one reference, and two EMG) in order to record vigilance states using previously published methods (12). Mice were given a minimum of 14 d to recover from surgery and habituate to the sleep recording. The animals were recorded simultaneously using commercially available hardware and software (Micromed France, SystemPLUS Evolution version 1092; Compumedics PSG, Compumedics).

**Experimental Design.** After surgery, different groups of mice were exposed to the following light/dark conditions: 1) LD12:12, 2) DD, 3) LD1:1, 4) LD3.5:3.5, 5) simulated transequatorial jet lag (LD18:6), and 6) simulated transmeridian jet lag (8-h prolongation of darkness followed by LD12:12). For anatomy experiments, mice were exposed to a either a 1-h light pulse from ZT 15 to 16 or darkness at the same time.

**Sleep and Wake Analysis.** Locomotor activity monitoring. General activity was monitored using a standard infrared motion detector, and data were analyzed using the ClockLab software package (Actimetrics). SCNx and Syt10Cre/+Bmal1fl/− mice were recorded for at least 10 d under LD12:12 followed by 2 wk under DD to measure their free running period to confirm the loss of circadian rhythm. Syt10Cre/+Bmal1ﬂ+/− mice were also recorded under the same conditions to confirm that they display a circadian pattern of locomotor activity similar to those of WT. The actimetry recordings were also performed (in parallel to ECoG recordings).

**Sleep Scoring and Spectral Analysis of the ECoG.** ECoG and EMG signals were acquired at 256 Hz and stored as continuous data streams. Vigilance states were scored into 4-s epochs using both ECoG and EMG signals based on standard criteria (32). The behavior in each of these 4-s epochs was classified as waking REMs or NREMS by visual inspection of the ECoG and EMG signals. The scoring was performed without information on the recording condition and genotype. The time spent in each of the vigilance states was averaged over 5 s and 1, 12, and 24 h in order to calculate intergroup differences. Spectral analysis was performed on raw signals every 4 s to which a Hanning window had been applied using a discrete Fourier transform. This produced power spectra between 0 and 128 Hz separated to 0.25 Hz bins. Dynamic changes in waking behavior were analyzed for theta (6 to 9 Hz) and gamma (40 to 70 Hz) frequencies and referenced to time periods where values were lowest (ZT 4 to 7).

**Model Parameters for Disturbed Light Condition Experiments.** The calculations for disturbed light conditions (4 to 7) were based on the percentages determined under LD12:12 (Figs. 2 and 3) for mice lacking either a functional SCN or melanopin. Briefly, under the light periods, the NREMS amounts are presumed to be due to photic influence of the circadian (55% CE) and noncircadian (45% SDLE) transduction systems. For baseline and experimental LD16:16 conditions, the following formula was used:

$$\text{NREMS}_{\text{baseline}} = \text{NREMS}_{\text{CE}} + \text{NREMS}_{\text{SDLE}}$$

Where

$$\text{NREMS}_{\text{CE}} = \frac{0.55 \times \text{NREMS}_{\text{baseline}}}{1 + 0.45}$$

$$\text{NREMS}_{\text{SDLE}} = \frac{0.45 \times \text{NREMS}_{\text{baseline}}}{1 + 0.55}$$
Light period = (NREMS_{total} × 0.55) + (NREMS_{min} × 0.45)

Dark period = (NREMS_{total} × 0.55).

For the jet lag experiment (Fig. 4 C and D), a CE shift of 1.1 h per 12 h was used. To achieve this, NREMS hourly values were split into 100 even bins (36 s). Subsequent days accounted for this change:

Light period = \sum (\text{NREMS}_{bin} × 0.55) + \sum (\text{NREMS}_{min} × 0.45)

Dark period = \sum (\text{NREMS}_{bin} × 0.55).

The values were then collapsed into hourly NREMS amounts. The model is denoted in purple in Fig. 4C and the CE and SDLE contributions in green and blue, respectively.

Anatomical Studies.

Immunostaining. After completion of the protocol, mice were deeply anesthetized with CO₂ and perfused with 0.1 M phosphate-buffered saline (PBS) pH 7.4 followed by transcardial fixation for 15 min with 4% paraformaldehyde in PBS, pH 7.4. In order to study c-Fos immunoreactivity in response to light in Syt10^{Cre/Cre}/Bmal1^{fl/+} (n = 3), Syt10^{Cre/Cre}/Bmal1^{fl/fl} (n = 4), and WT (n = 3) animals, the euthanasia occurred at the conclusion of a 1 h light pulse administered at ZT 15 and without a light pulse as a negative control. SCNx and sham animals were euthanized 48 h after intraocular injection of CtB (see Retinal Tract Tracing). To control for SCN lesion size and the conservation of retino-cerebral projections, immunohistochemistry was used in SCNx and sham-lesioned animals to stain AVP, DAPI, and CtB. To evaluate SCN cells reactivity to light in Syt10^{Cre/Cre}/Bmal1^{fl/fl} and their controls, c-Fos expression was quantified in the SCN with AVP staining. The procedure was carried out as described previously (12). After dehydration in 30% sucrose for 48 h, the brains were frozen and cut in a freezing microtome. Free-floating slices, brought to room temperature, were rinsed three times (5 min each) in 0.1 M PBS (pH 7.0), incubated in 5% normal donkey serum (Normal Donkey Serum, 017-000-121 Jackson Immunoresearch) and 0.25% Triton X-100 (TX) in PBS for 1 h, and then incubated overnight at 4 °C with primary antibodies (rabbit anti-c-Fos IgG, SC52 Santa Cruz Biotechnology, diluted 1:500, Sigma-Aldrich D9542). The omission of the primary antibody abolished all staining (control).

Photomicrographs and Quantification. Immunostainings were analyzed using a microscope equipped with appropriate filter settings for detecting Cy5, Alexa 555, and DAPI. Fluorescence images were obtained via a non-confocal microscope (DMRXA2, Leica Microsystems) equipped with Metamorph v 2.1.39 (Olympus). Light microscopy images were grabbed with a Leica DC200 camera using Leica DC200 software (Leica). The software program ImiEl was used to fuse the images, and the image editing software Microsoft publisher was used to combine the obtained images into panels. The reference of the various brain structures was made according to the Franklin and Paxinos Atlas “Mouse brain in stereotaxic coordinates” (39).

For the quantification of c-Fos immunoreactivity to light, the number of c-Fos-positive neurons in the SCN was counted by two independent experimenters blind to experimental conditions in an area of 300 × 500 μm at levels corresponding to bregma −0.30 to −0.70 mm (rostro caudal to the posterior of the SCN) (Fig. 2 A, viii). Approximately nine brain sections from each animal containing the SCN (with AVP containing) were analyzed, and c-Fos-positive cells were visually counted using a grid on each image. C-Fos immunoreactivity in response to light was analyzed using two-way ANOVA (light pulse and genotype conditions) followed by post hoc t tests.

Retinal Tract Tracing. After the completion of the recording protocol, SCNx and sham animals (sham Opn4<sup>−/−</sup> n = 9; sham Opn4<sup>−/−</sup> + n = 7; SCNx Opn4<sup>−/−</sup> n = 10; SCNx Opn4<sup>−/−</sup> + n = 8; WT n = 5; Syt10<sup>Cre/Cre</sup>/Bmal1<sup>fl/fl</sup> n = 6; and Syt10<sup>Cre/Cre</sup>/Bmal1<sup>fl/fl</sup> n = 8) were anesthetized (isoflurane 2.5%, applied during all procedure), and retinal transverse tracer GST (dissolved in 50 mM PBS, pH 7.4) was injected into the vitreal for over 2 min. After 48 h, the animals were perfused, and the histology procedure was processed as described below.

Statistical Methodology. All statistics were calculated using standard methods with Statistica (Statsoft v. 8), and graphics were generated either in SigmaPlot (Systat v. 11) or Microsoft Excel (v. 2013). The differences in n values between certain light/dark regimes were due to ECoG/EGM signal problems on the day of recording. For ECoG spectrum analysis, some animals were excluded due to the increased number of signal artifacts, which allowed for the quantification of sleep and wake distribution, but hampered Fourier signal transformation. The differences in sleep amounts and quantitative ECoG variables were determined by single- or multiple-way ANOVAs followed by post hoc t tests if 5% significance levels were reached. The differences in the number of c-Fos-positive neurons were assessed by two-way (light and genotype conditions) ANOVA followed by post hoc t tests.

Data Availability. Data used to generate the anatomy figures in this manuscript are accessible on a public FigShare repository (DOI: 10.6084/m9.figshare.14176520). All other study data are included in the article and/or SI Appendix.

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23. H. C. van Diepen, A. Ramkisoensing, S. N. Peirson, R. G. Foster, J. H. Meijer, Irradiance encoding in the suprachiasmatic nuclei by rod and cone photoreceptors. *FASEB J.* 27, 4204–4212 (2013).

24. C. M. Altimus *et al.*, Rods-cones and melanopsin detect light and dark to modulate sleep independent of image formation. *Proc. Natl. Acad. Sci. U.S.A.* 105, 19998–20003 (2008).

25. T. A. LeGates *et al.*, Aberrant light directly impairs mood and learning through melanopsin-expressing neurons. *Nature* 491, 594–598 (2012).

26. J. Husse, X. Zhou, A. Shostak, H. Oster, G. Eichele, Synaptotagmin10-Cre, a driver to disrupt clock genes in the SCN. *J. Biol. Rhythms* 26, 379–389 (2011).

27. J. A. Evans *et al.*, Shell neurons of the master circadian clock coordinate the phase of tissue clocks throughout the brain and body. *BMC Biol.* 13, 43 (2015).

28. X. Jin *et al.*, A molecular mechanism regulating rhythmic output from the suprachiasmatic circadian clock. *Cell* 96, 57–68 (1999).

29. M. Mieda *et al.*, Cellular clocks in AVP neurons of the SCN are critical for interneuronal coupling regulating circadian behavior rhythm. *Neuron* 85, 1103–1116 (2015).

30. S. A. Rahman *et al.*, Diurnal spectral sensitivity of the acute alerting effects of light. *Sleep (Basel)* 37, 271–281 (2014).

31. G. Buzsáki, E. I. Moser, Memory, navigation and theta rhythm in the hippocampal-entorhinal system. *Nat. Neurosci.* 16, 130–138 (2013).

32. C. S. Herrmann, M. H. Munk, A. K. Engel, Cognitive functions of gamma-band activity: Memory match and utilization. *Trends Cogn. Sci.* 8, 347–355 (2004).

33. X. J. Wang, Neurophysiological and computational principles of cortical rhythms in cognition. *Physiol. Rev.* 90, 1195–1268 (2010).

34. A. A. Borbély, A two process model of sleep regulation. *Hum. Neurobiol.* 1, 195–204 (1982).

35. P. Franken, D. Chollet, M. Tafti, The homeostatic regulation of sleep need is under genetic control. *J. Neurosci.* 21, 2610–2621 (2001).

36. P. Franken *et al.*, NPAS2 as a transcriptional regulator of non-rapid eye movement sleep: Genotype and sex interactions. *Proc. Natl. Acad. Sci. U.S.A.* 103, 7118–7123 (2006).

37. F. K. Stephan, I. Zucker, Circadian rhythms in drinking behavior and locomotor activity of rats are eliminated by hypothalamic lesions. *Proc. Natl. Acad. Sci. U.S.A.* 69, 1583–1586 (1972).

38. T. I. Morgenthaler *et al.; Standards of Practice Committee of the American Academy of Sleep Medicine, Practice parameters for the clinical evaluation and treatment of circadian rhythm sleep disorders. An American Academy of Sleep Medicine report. Sleep* 30, 1445–1459 (2007).

39. G. Paxinos, K. B. Franklin, *Paxinos and Franklin’s the Mouse Brain in Stereotaxic Coordinates* (Academic Press, 2019).

40. J. Hubbard *et al.*, Dataset for "Dissecting and modeling photic and melanopsin effects to predict sleep disturbances induced by irregular light exposure in mice." *FigShare.* http://dx.doi.org/10.6084/m9.figshare.14176520. Deposited 3 June 2021.