

A Distinct Visual Pathway Mediates High-Intensity Light Adaptation of the Circadian Clock in *Drosophila*

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To provide organisms with a fitness advantage, circadian clocks have to react appropriately to changes in their environment. High-intensity (HI) light plays an essential role in the adaptation to hot summer days, which especially endanger insects of desiccation or prey visibility. Here, we show that solely increasing light intensity leads to an increased midday siesta in *Drosophila* behavior. Interestingly, this change is independent of the fly's circadian photoreceptor cryptochrome and is solely caused by a small visual organ, the Hofbauer–Buchner eyelets. Using receptor knock-downs, immunostaining, and recently developed calcium tools, we show that the eyelets activate key core clock neurons, namely the *s-LN_vs*, at HI. This activation delays the decrease of PERIOD (PER) in the middle of the day and propagates to downstream target clock neurons that prolong the siesta. We show a new pathway for integrating light-intensity information into the clock network, suggesting new network properties and surprising parallels between *Drosophila* and the mammalian system.

Key words: circadian clock; *Drosophila*; photoreceptor

Significance Statement

The ability of animals to adapt to their ever-changing environment plays an important role in their fitness. A key player in this adaptation is the circadian clock. For animals to predict the changes of day and night, they must constantly monitor, detect and incorporate changes in the environment. The appropriate incorporation and reaction to high-intensity (HI) light is of special importance for insects because they might suffer from desiccation during hot summer days. We show here that different photoreceptors have specialized functions to integrate low-intensity, medium-intensity, or HI light into the circadian system in *Drosophila*. These results show surprising parallels to mammalian mechanisms, which also use different photoreceptor subtypes to respond to different light intensities.

Introduction

Circadian clocks have evolved to allow animals and plants to predict the changes of day and night and thereby increase their fitness (Daan and Tinbergen, 1980; DeCoursey et al., 2000). A crucial property of circadian clocks is the ability to adjust their physiology and behavior to light–dark cycles, in a process called entrainment (Golombek and Rosenstein, 2010).

In all clock cells, the transcriptional–translational feedback loop is believed to underlie these circadian rhythms of behavior. In short, Clock and Cycle activate *period* (*per*) and *timeless* (*tim*) transcription, respectively. PER and TIM proteins accumulate in the early night, reenter the nucleus toward the end of the night, and inhibit their own transcription (Hardin et al., 1990; Sehgal et al., 1994; Allada et al., 1998; Rutila et al., 1998).

In mammals, the circadian photoreceptor melanopsin in the retinal ganglion cells, as well as the rods and cones of the visual system, can transduce light information into the central brain clock, the suprachiasmatic nucleus (Ebihara and Tsuji, 1980; Benson et al., 2002; Hattar et al., 2002). Similarly, flies can use the circadian photoreceptor cryptochrome (CRY) within most of the clock neurons as well as their visual system to synchronize their clock (Stanewsky et al., 1998; Rieger et al., 2003; Benito et al., 2008; Yoshii et al., 2008). The latter consists of the two compound eyes, three ocelli, and two Hofbauer–Buchner (HB) eyelets (Hofbauer and Buchner, 1989).

The clock neurons determine the diurnal behavior pattern of the fly. In light–dark (LD) cycles, flies show a bimodal locomotor

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activity pattern with a morning (M) peak and an evening (E) peak, which are clearly separated by a siesta. Essential groups of lateral neurons include the s-LN_vs, which are important for morning anticipation (M activity) and are often referred to as M cells (Grima et al., 2004; Stoleru et al., 2004). The l-LN_vs are important arousal centers (Shang et al., 2008). Additional circadian lateral neurons include three CRY-positive LN_as, which dictate the E peak and are referred to as E cells.

In vitro experiments have investigated the connection of the visual system to the clock network. Whereas the compound eyes appear to be contacting the l-LN_vs via cholinergic interneurons, the HB eyelets directly innervate the accessory medulla and increase Ca²⁺ and cAMP in the s-LN_vs upon stimulation (Muraro and Ceriani, 2015; Schlichting et al., 2016). However, it is not understood how the visual system affects the rest of the central brain clock or how different light intensities affect these connections.

Studies in mammals and flies show that a complex visual system is required to adjust an animal clock to the light environment (Foster and Helfrich-Förster, 2001). At low intensity (LI), entrainment is predominantly performed by retinal rods, whereas melanopsin takes over at HI due to the saturation of the visual system by light (Lall et al., 2010; Lucas et al., 2012). In *Drosophila*, CRY is important for detecting LI due to its photon-integrating function (Vinayak et al., 2013) and four of the five rhodopsin-expressing photoreceptors of the compound eyes mediate entrainment at LI to moderate irradiances (Saint-Charles et al., 2016). Photoreception for HI is still unknown. However, flies lacking the whole visual system, as well as CRY, are unable to entrain to LD regimes of moderate light intensity (Helfrich-Förster et al., 2001).

In this study, we investigate how HI changes fly behavior and how it is incorporated into the clock network. HI significantly increases the fly siesta by delaying the E-activity onset. This phenotype is independent of CRY and the compound eyes, but does require the HB eyelets. We further show that the release of acetylcholine significantly increases *in vivo* Ca²⁺ and neuronal activity of the eyelet target neurons, the s-LN_vs. Our data further suggest that HI delays PER degradation in the s-LN_vs during the day. This change propagates throughout the network to change PER cycling of downstream target neurons, such as the DN₁s, which have been implicated in the control of the siesta (Guo et al., 2016).

Materials and Methods

Fly strains

The following fly strains have been described previously and were used in this study: WT_{CS}, WT_{ALA} (Sandrelli et al., 2007), WT_{LB} (Schlichting et al., 2014), *w¹¹¹⁸*, *cry⁰¹* (Dolezelova et al., 2007), *cli^{eya}* (Bonini et al., 1993), *rh6¹* (Cook et al., 2003), *rh5²* (Yamaguchi et al., 2008), *ninaE¹⁷* (Kumar and Ready, 1995), *sev^{LY3}*, *cli^{eya};rh6¹*, *norPA^{P24}*, *nocte* (Glaser and Stanewsky, 2005) *hdc^{JK910}* (BL 64203), *R6-GAL4* (Helfrich-Förster et al., 2007), *UAS-mAChRA-RNAi* (BL 44469, 27571), *UAS-nAChR-RNAi* (BL 31883, 28688), *pdf-GS-GAL4* (Depetris-Chauvin et al., 2011), *ActP-MKII::GAL4DBDo*, *ActP-p65AD::CaM* (UAS-Tric) (Gao et al., 2015), and *UAS-GFPS65T* (BL 1522). All experiments were performed in male flies.

Behavior recording

Regensburg system. Individual 2- to 5-d-old male flies were transferred into photometer half-cuvettes with water and sugar supply. The fly's activity was measured as IR beam crosses caused by the fly in 1 min intervals. Independent sets of flies were tested at LI (~10 lux, 1 week duration) or HI (~10,000 lux, 1 week duration). Independent sets of flies were used to exclude behavioral effects due to aging.

Würzburg system. Individual 2- to 5-d-old male flies were placed into glass tubes containing food (2% agarose, 4% sucrose). These glass tubes were placed in *Drosophila* activity monitors (DAMs) and the activity of the flies was measured in 1 min intervals. Flies were entrained for 1 week at LI (~10 lux) followed by 1 week of HI (~8000 lux). We were only able to reach 8000 lux due to a different position of the LEDs using the DAM system. Both light intensities were investigated within the same flies to follow changes in behavior within individual flies. All experiments were conducted within light boxes in climate-controlled chambers at a temperature of 20°C. To minimize fluctuations in temperature, each light box was equipped with a fan to constantly exchange the air within the light box.

Behavior analysis

All data were plotted as actograms using ActogramJ (Schmid et al., 2011). Subsequently, we calculated average activity and average sleep profiles using the last 4 d of each light condition as described previously (Schlichting and Helfrich-Förster, 2015). Sleep was measured as intervals of at least 5 min of inactivity. To determine the timing of M and E activity and onset/offset, we generated single-fly average days in 30 min bins. We then determined fly by fly the first time point when a fly's activity reached the siesta activity amount as the offset of M activity and the first time point of increased activity beyond siesta levels as E-peak onset. This analysis was done blindly by two independent investigators and we calculated the average value ± SEM. We further calculated a performance index (PI) to determine the flies' E-peak onset to overcome subjective determination of E-peak onset. The PI is calculated as follows: PI = sum of activity 3–6 h before E peak HI/sum of activity 3–6 h before E peak LI. A PI < 1 indicates a reduction of activity, which we interpret as a delay in E-peak onset. Data were compared using a Student's *t* test or a one-way *t* test comparing the PI with the value 1.

Immunostaining and confocal microscopy

Fifteen to 20 male flies were fixed in 4% PFA in PBST (0.5% Triton X-100) for 2 h and 45 min and subsequently rinsed with 0.5% PBST 4 × 10 min each. Brains were dissected in PBST and blocked in 5% NGS in PBST for 3 h. Primary antibodies were applied overnight at RT. The following antibodies were used: rabbit anti-PER (1:1000, provided by R. Stanewsky; Stanewsky et al., 1997), rat anti-TIM (1:1000, provided by J. Giebultowicz), mouse anti-pigment dispersing factor (anti-PDF, 1:2000, DSHB), and chicken anti-GFP (1:1500, Abcam). After rinsing 5 × 10 min each with PBST, secondary antibodies (Invitrogen) were applied for 3 h at room temperature. After washing the brains 5 × 10 in PBST, brains were mounted on glass slides using Vectashield mounting medium.

For comparing PER and TIM profiles at LI and HI, CS and *cli^{eya};rh6¹* flies were entrained to the respective light intensity for 5 d and collected around the clock in 2 h intervals. For imaging the dorsal arborizations, CS and *rh6¹* flies were entrained at the standard LD regime and collected at ZT6. To evaluate neuronal activity at LI and HI, *pdf-GS>Tric>GFP* flies were raised on standard cornmeal medium. Male flies between 2 and 5 d after eclosion were transferred on agar (2% agar, 4% sugar, and 200 μg/ml Ru486) and entrained for 3 d in LD 12:12 at LI and HI, respectively.

All brains were imaged using a Leica SPE or Leica SP5 confocal microscope in 2 μm sections. All settings were kept constant within the experiment. Staining intensity was determined by measuring the brightest 9 pixels of the cell of interest minus 3 different background intensities as described previously (Menegazzi et al., 2013).

Results

HI light delays the evening peak and lengthens the siesta

To investigate the effect of light intensity on fly behavior, we subjected three different WT strains to light dark cycles of LI and HI white light. All flies displayed the classic bimodal activity pattern with an M peak and an E peak of activity. These peaks were clearly separated by a siesta, during which flies tend to sleep (Fig. 1A). We did not observe any consistent changes in M- or E-peak amplitude or timing.

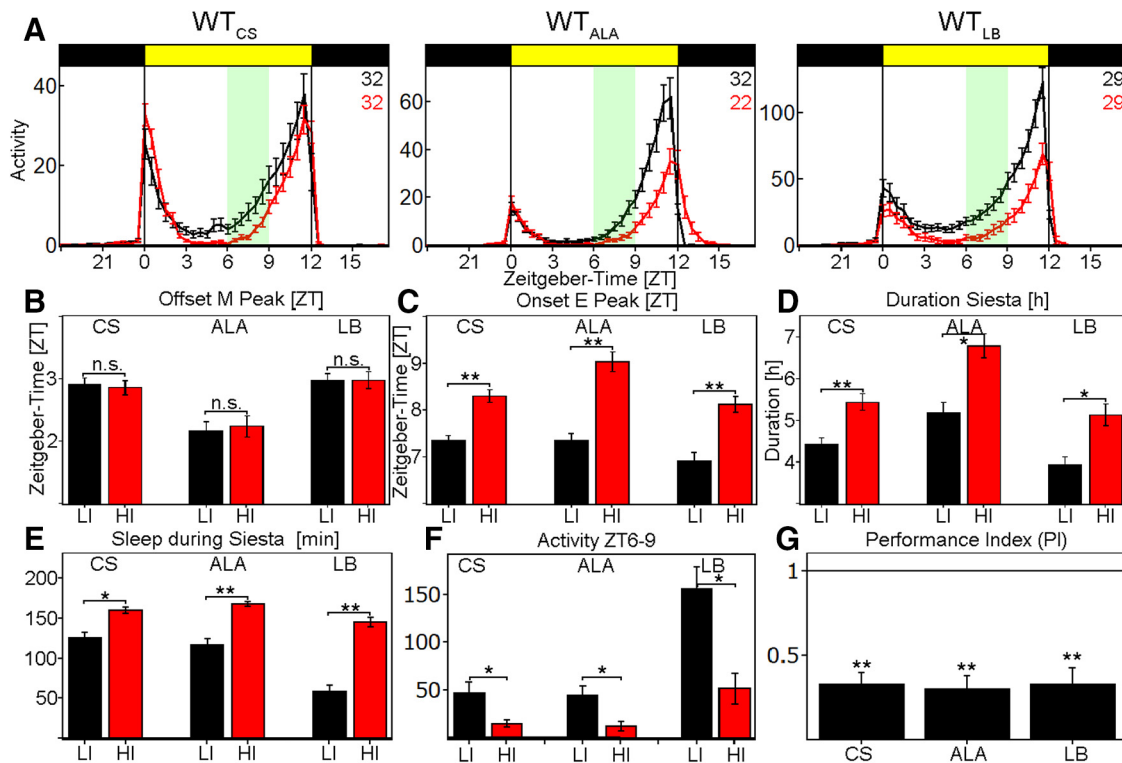


Figure 1. Behavior of WT-flies at LI (black line) and HI (red line). **A**, Average activity profile of WT_{CS} (left), WT_{ALA} (middle), and WT_{LB} (right) at LI (10 lux, black line) and HI (10,000 lux, red line) \pm SEM recorded in the Regensburg system. All flies show a bimodal activity pattern with reduced activity during the siesta. **B**, Quantification of M-peak offset at LI (black) and HI (red) in ZT \pm SEM. All strains show a M-peak offset \sim 2–3 h after lights on, but do not change the onset upon HI (WT_{CS}: $p = 0.9379$, WT_{ALA}: $p = 0.1859$, WT_{LB}: $p = 0.3862$). **C**, Quantification of E-peak onset at LI (black) and HI (red) in ZT \pm SEM. All flies significantly delay the E-peak onset by \sim 1 h upon HI simulation ($p < 0.0001$ for all genotypes). **D**, Duration of siesta in hours \pm SEM. All flies show a significantly longer siesta at HI compared with LI (WT_{CS}: $p = 0.0001$, WT_{ALA}: $p = 0.0015$, WT_{LB}: $p = 0.0012$). **E**, Total sleep amount at LI (black) and HI (red) between ZT6 and ZT9 in minutes \pm SEM. All flies significantly increase their sleep amounts at HI (WT_{CS}: $p = 0.0014$, WT_{ALA}: $p < 0.0001$, WT_{LB}: $p < 0.0001$). **F**, Sum of activity indicated in the green area in **A**. All flies significantly reduce their activity level at HI (WT_{CS}: $p = 0.0091$, WT_{ALA}: $p = 0.0114$, WT_{LB}: $p = 0.0006$). **G**, PI calculated by the sum of activity between ZT6 and ZT9 at HI divided by the sum of activity between ZT6 and ZT9 at LI. All genotypes show a PI significantly < 1 , which we interpret as a delayed E-peak onset ($p < 0.0001$ for all genotypes). * $p < 0.05$, ** $p < 0.001$.

We then considered the shape of the activity profiles. Upon careful inspection, all WT strains appeared to lengthen their siesta by delaying the E-peak onset by 1–2 h, whereas the M-peak offset remained unchanged (Fig. 1B–D). This change in behavior was accompanied by significantly reduced activity levels between ZT6 and ZT9 (green boxes in Fig. 1A, F). We calculated a PI from the activity levels during this period. A PI < 1 represents a reduction of activity at HI, whereas a $p > 1$ represents an increase. Because the PI is a more objective measurement of the E-peak onset, we focused on this value throughout the study. All WT strains showed a significant reduction of activity, which we interpret as a delayed E-activity onset (Fig. 1G). One explanation for this behavior is that HI induces a longer midday siesta. To address this possibility, we calculated sleep levels in all WT strains. Indeed, HI significantly increased the amount of sleep during the siesta by at least 30 min (Fig. 1E). HI therefore delayed the E activity onset by lengthening the siesta and increasing sleep amounts.

HB eyelets mediate light-intensity integration

To test which light input pathway mediates this siesta response, we analyzed the behavior of several photoreceptor mutants. *Cry*⁰¹ flies, like WT, showed a bimodal behavior with M and E peaks around lights on or lights off, respectively (Fig. 2A). The mutant strain also responded to HI similarly if not identically to the WT strain: it significantly delayed the onset of E activity ($p < 0.0001$; Fig. 2B) and showed a PI significantly < 1 ($p < 0.0001$;

Fig. 2C). These results suggest that HI responses are not mediated via CRY.

We also investigated mutants affecting the fly visual system. We first used the *cli*^{eya} mutant strain to investigate the contribution of the compound eyes for several reasons: (1) the mutant showed no residual eye structures; (2) we previously showed that ocelli and HB eyelets are still intact (Schlichting et al., 2014), and (3) several circadian studies have previously characterized this strain. As in earlier studies, *cli*^{eya} mutants showed a significantly reduced M-peak amplitude, as well as an advanced E peak in both LD conditions (Fig. 2A). A comparison between LI and HI in this strain indicated that HI still exhibits a delay of E activity onset ($p < 0.0001$; Fig. 2B) and a PI significantly < 1 in response to HI ($p < 0.0001$; Fig. 2C). This suggests that the siesta and E peak phenotypes are independent of the compound eyes.

We then investigated the effect of the HB eyelets. Because the lack of specific drivers precluded manipulating the eyelets without perturbing the compound eyes, we examined the effect of the *rh6*¹ mutant. The mutation eliminates the photopigment of the eyelets but also of 70% of R8 in the compound eyes (Sprecher and Desplan, 2008). The mutant exhibited a reduced M-peak amplitude and an advanced E peak under both LI and HI light (Fig. 2A). Importantly, the profiles did not differ between LI and HI light: they neither delayed the onset of E activity ($p = 0.2885$; Fig. 2B) nor significantly reduced the PI from 1 ($p = 0.5619$; Fig. 2C).

To verify that the effect is solely visible in *rh6*¹ mutant flies, we monitored the behavior of several other rhodopsin mu-

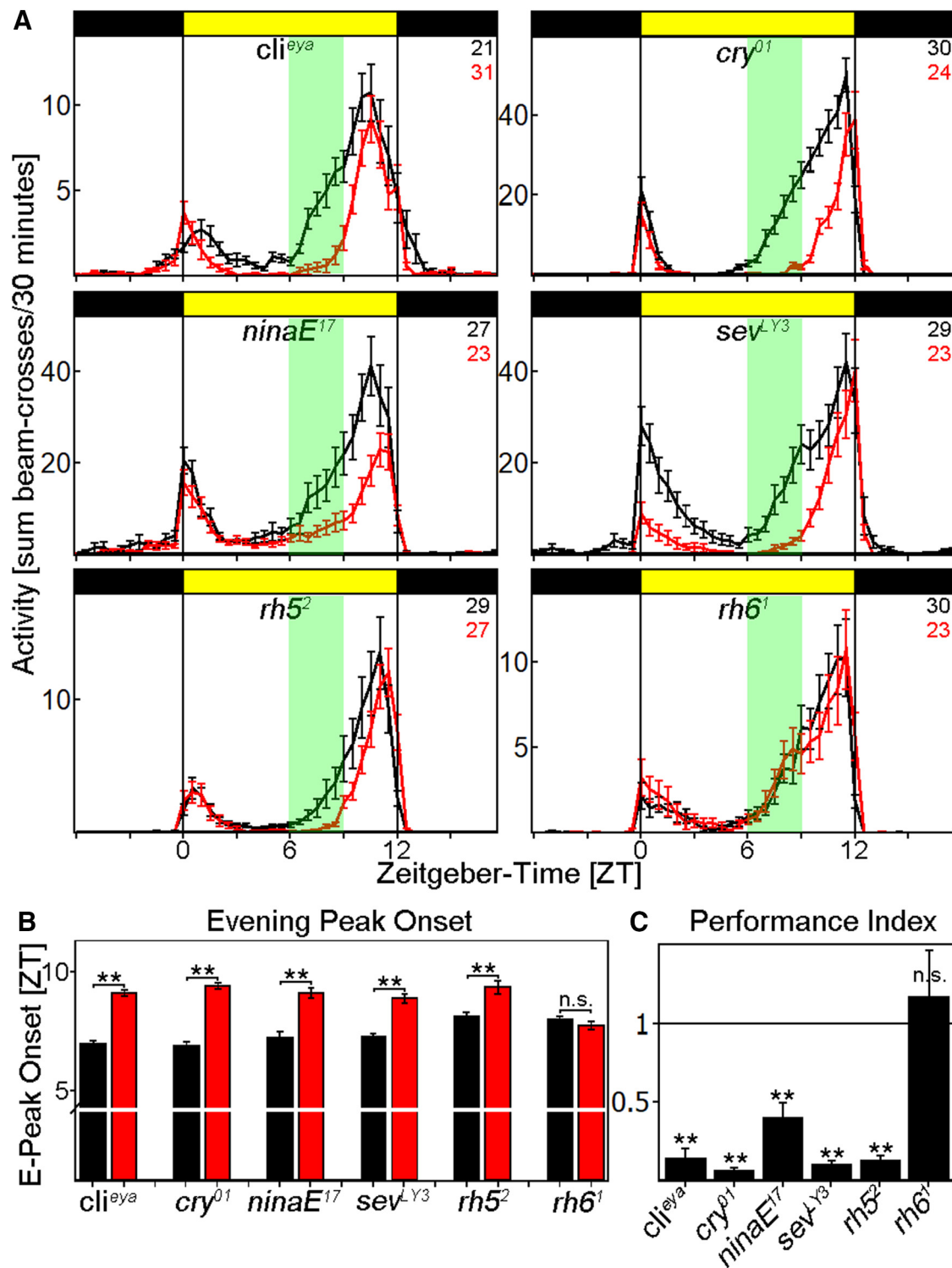


Figure 2. Behavior of photoreceptor mutants at LI and HI. **A**, Average activity profiles of flies exposed to LI (black line) and HI (red line) \pm SEM using the Regensburg system. Flies deficient of either compound eyes (*cli^{eya}*) or CRY (*cry⁰¹*) behave similar to WT_{CS}, whereas the loss of rhodopsin 6 renders the flies insensitive to light-intensity changes. Consistent with the *cli^{eya}* data, flies affecting the compound eyes (*ninaE¹⁷*, *sev^{LY3}*, and *rh5²*) behave like WT_{CS}. **B**, Determination of E-activity onset for all photoreceptor mutants. *cli^{eya}*, *cry⁰¹*, *ninaE¹⁷*, *sev^{LY3}*, and *rh5²* flies significantly delay the onset of the E activity at HI ($p < 0.0001$), whereas *rh6¹* mutants do not change the timing ($p = 0.2885$). **C**, *cli^{eya}*, *cry⁰¹*, *ninaE¹⁷*, *sev^{LY3}*, and *rh5²* flies show a PI significantly < 1 ($p < 0.0001$ for all), whereas the PI of *rh6¹* flies is indistinguishable from 1 ($p = 0.5619$). ** $p < 0.001$.

tants under LI and HI conditions: *ninaE¹⁷*, *sev^{LY3}*, and *rh5²*. They all significantly delayed the E-peak onset ($p < 0.0001$; Fig. 2*A, B*) and decreased their activity between ZT6 and ZT9 at HI as WT flies did, resulting in a PI significantly < 1 ($p < 0.0001$).

To exclude the possibility that heating of the flies at HI is the reason for the observed siesta phenotype, we tested *nocte* flies, which were previously shown to be deficient in temperature entrainment (Glaser and Stanewsky, 2005). As expected, these flies still showed a PI significantly < 1 (PI = 0.25 ± 0.06 , $p < 0.0001$),

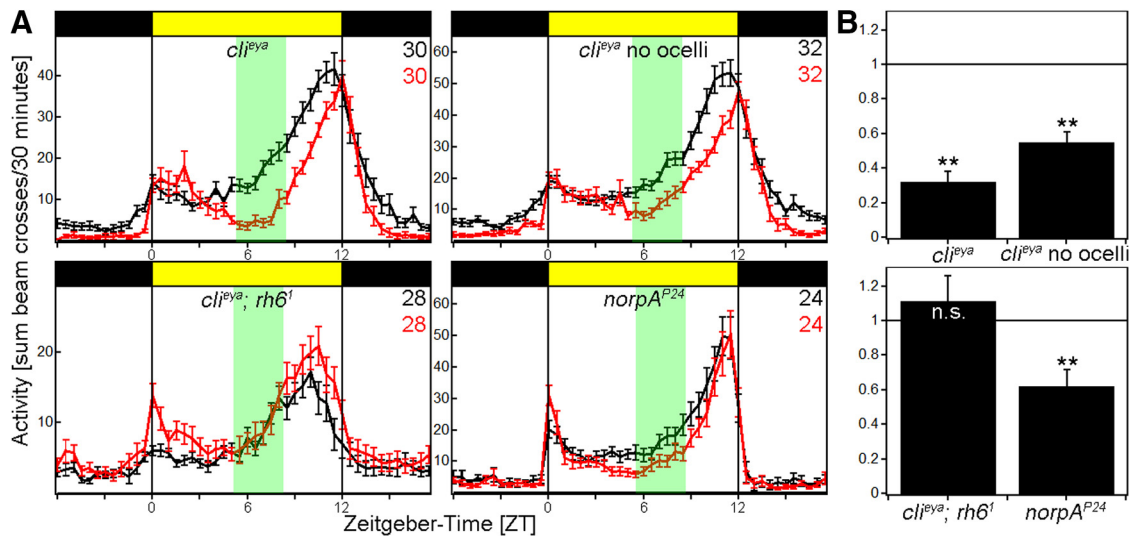


Figure 3. Behavior of photoreceptor mutants at LI and HI. **A**, Average activity profiles of flies exposed to LI (black line) and HI (red line) \pm SEM using the Würzburg system. Flies deficient of either compound eyes (*cli^{eya}*) or compound eyes and ocelli (*cli^{eya} no ocelli*) and flies lacking *norpA* (*norpA^{P24}*) still react to HI, whereas the loss of rhodopsin 6 in *cli^{eya}* background renders the flies insensitive to light-intensity changes. **B**, *cli^{eya}*, *cli^{eya} no ocelli* and *norpA^{P24}* flies show a PI significantly <1 (*cli^{eya}*, *cli^{eya} no ocelli*: $p < 0.0001$, *norpA^{P24}*: $p = 0.0014$), whereas the PI of *cli^{eya}; rh6¹* is indistinguishable from 1 ($p = 0.4613$). ** $p < 0.001$.

suggesting that light input is the source for the change in behavior (data not shown).

cli^{eya} mutant flies still have working ocelli and HB eyelets. To address the possibility of the ocelli contributing to HI entrainment, we painted the ocelli of *cli^{eya}* mutant flies to eliminate their function. *cli^{eya}* mutants with painted ocelli showed a significantly delayed E-peak onset and a PI significantly <1 at HI indistinguishably from *cli^{eya}* mutants with functional ocelli (Fig. 3A, B, top). This suggests that the ocelli do not or only marginally contribute to HI entrainment.

We further created *cli^{eya}; rh6¹* mutants to eliminate HB eyelet function in *cli^{eya}* background. These flies reproduced the *rh6¹* mutant phenotype: flies showed a reduced M-peak amplitude and an advanced E peak compared with WT flies in both light conditions; they also did not delay their E-peak onset at HI as evident by their PI, which was not different from 1 (Fig. 3A, B, bottom). These data further suggest that the HB eyelet is principally responsible for the adaptation to HI.

Rh6-positive photoreceptors were recently shown to use a *norpA*-independent phototransduction pathway (Ogueta et al., 2018). To address this issue, we monitored the behavior of *norpA^{P24}* flies at LI and HI. These mutant flies had a PI significantly <1 similar to WT flies (Fig. 3A, B, bottom), implicating the newly discovered *norpA*-independent pathway in the HB eyelets as being involved in mediating the adaptation to HI.

Acetylcholine mediates signal transduction to clock cells

The HB eyelets signal to the s-LN_vs via acetylcholine, whereas they appear to signal to the l-LN_s via histamine (Schlichting et al., 2016). We therefore investigated which of these two neurotransmitters transduces light-intensity information to the clock neuron network. As described previously, flies unable to synthesize histamine (*hdc^{JK910}*) showed a reduced M peak as well as an advanced E peak (Fig. 4A, top), but they also showed a PI significantly <1 ($p = 0.0012$; Fig. 4B, top); this suggests that histamine is not necessary for siesta adaptation to HI. Because histamine is the only neurotransmitter of the compound eyes (Pollack and Hofbauer, 1991), this agrees with the previous *cli^{eya}* data (Figs. 2A, 3A).

Acetylcholine plays a key role as an excitatory neurotransmitter in mammals and insects and the *Drosophila* genome encodes 10 nicotinic acetylcholine receptor subunits ($\alpha 1$ – $\alpha 7$ and $\beta 1$ – $\beta 3$); subpopulations of these subunits are implicated in *Drosophila* behavior. In addition, the genome encodes two muscarinic acetylcholine receptors (*mAChRA* and *mAChRB*). To investigate the contribution of these receptors to light-intensity integration, we expressed RNAi constructs against specific receptors or receptor subunits in the s-LN_vs with the R6-GAL4 driver.

We tested RNAi lines against two of the 10 existing *nAChR* subunits (Fig. 4A, B, bottom), and all flies showed a PI <1 ($p < 0.001$). In contrast, knocking down *mAChRA* completely reproduced the behavior of *rh6¹* mutants. Knock-down flies showed a significantly reduced M-peak amplitude and a significantly advanced E peak compared with both parental controls (Fig. 4A). Similar results were obtained in two independent RNAi lines (Fig. 4A, middle). These data suggest that mAChRA acts in the s-LN_vs to integrate cholinergic input. Interestingly, RNA-sequencing data suggest that the s-LN_vs are selective: whereas *mAChRA* is expressed in all circadian subpopulations investigated, only the PDF cells specifically exclude *mAChRB* expression, indicating an important role of *mAChRA* in these neurons (Fig. 4D; Abruzzi et al., 2017).

mAChRA is coupled to G_{q/11} and interacts with IP3/Ca²⁺ second messenger pathways (Ren et al., 2015). Because previous dissected brain imaging experiments showed increased Ca²⁺ in the s-LN_vs when the HB eyelets were artificially activated using the P2X2 system (Schlichting et al., 2016), we expressed TRIC-GFP in PDF neurons (Gao et al., 2015), a transcriptional reporter surrogate for Ca²⁺. To avoid expression during development and express Tric-GFP only in the adult, we used the gene-switch system (*pdf*-GS-GAL4) and measured GFP intensity in the s- and l-LN_vs separately after exposing the flies to LD 12:12 of LI or HI for 3 d. There was no significant difference between LI and HI in the l-LN_vs, but a 2-fold increase of GFP in the s-LN_vs with HI, suggesting that the eyelets activate the s-LN_vs at HI *in vivo* (Fig. 4C).

An increase in Ca²⁺ can reflect either an increase in intracellular signaling and/or an increase in neuronal activity. To address

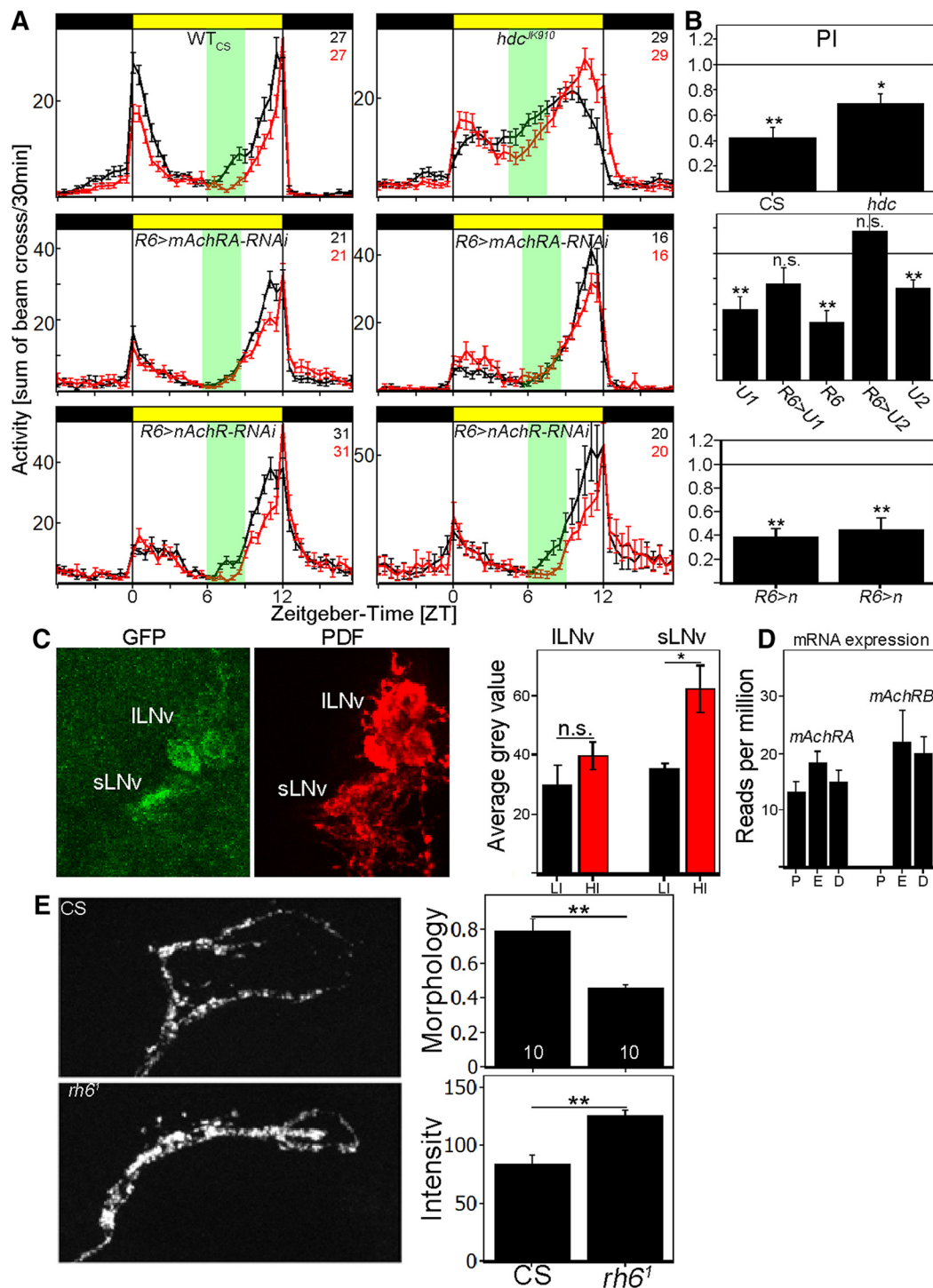


Figure 4. Cholinergic input to s-LN_s is responsible for HI adaptation. **A**, Average activity profiles at LI (black line) and HI (red line) ± SEM measured in the Würzburg system. Top, Average activity profile of WT_{CS} and the *hdc*^{JK910} mutant at LI (black line) and HI (red line) ± SEM. Flies deficient of histamine synthesis are still reacting to HI by a reduction of activity. Middle, Average activity profiles of *mAChRA* (BL: 27571; left) and *mAChRA* (BL: 44469) knock-down in sLN_s (right) at LI (black line) and HI (red line) ± SEM. Knock-down of *mAChRA* in the sLN_s reproduces *rh6*¹-mutant phenotype. The activity profiles do not differ between LI and HI. Bottom, Average activity profiles of 2 independent *nAChR* (left: BL 28688, right: BL 31883) knock-down in sLN_s at LI (black line) and HI (red line) ± SEM. Both genotypes show a delayed E-peak onset comparable to the GAL4 control. **B**, PIs calculated from **A**. WT_{CS} and *hdc*^{JK910} mutants significantly reduce their PI at HI (CS: $p < 0.0001$, *hdc*: $p = 0.0012$). *R6-GAL4* and both UAS control flies (U1: BL 27571; U2: BL 44469) show a PI significantly < 1 (*R6*: $p < 0.0001$, U1: $p = 0.0003$, U2: $p = 0.0002$), whereas the PI of *mAChRA* knock-down in sLN_s is indistinguishable from 1 (*R6>U1*: $p = 0.07134$, *R6>U2*: $p = 0.5529$). Knock-down of *nAChR* subunits does not affect HI adaptation and results in significantly reduced PI values ($p < 0.0001$). **C**, Brain images of *Pdf-GS>Tric-GFP* stained against GFP and PDF. GFP expression was measured in s-LN_s and l-LN_s as a correlate for Ca²⁺ levels in the individual neurons. s-LN_s significantly increase GFP levels at HI, whereas there was no change in l-LN_s. GFP levels, suggesting an increase of Ca²⁺ in s-LN_s at HI. **D**, Relative *AChR* mRNA expression in PDF (P) cells, evening (E) cells, or DN_s (D). Whereas *mAChRA* is expressed in all neuron clusters, the PDF cells do not show any *mAChRB* expression. **E**, Confocal images of s-LN_s dorsal projection in WT_{CS} and *rh6*¹ flies at ZT6 HI. WT_{CS} arborizations show a more open conformation compared with *rh6*¹ mutants ($p = 0.0003$), whereas *rh6*¹ mutants show a significantly higher staining intensity ($p < 0.0001$). * $p < 0.05$, ** $p < 0.001$.

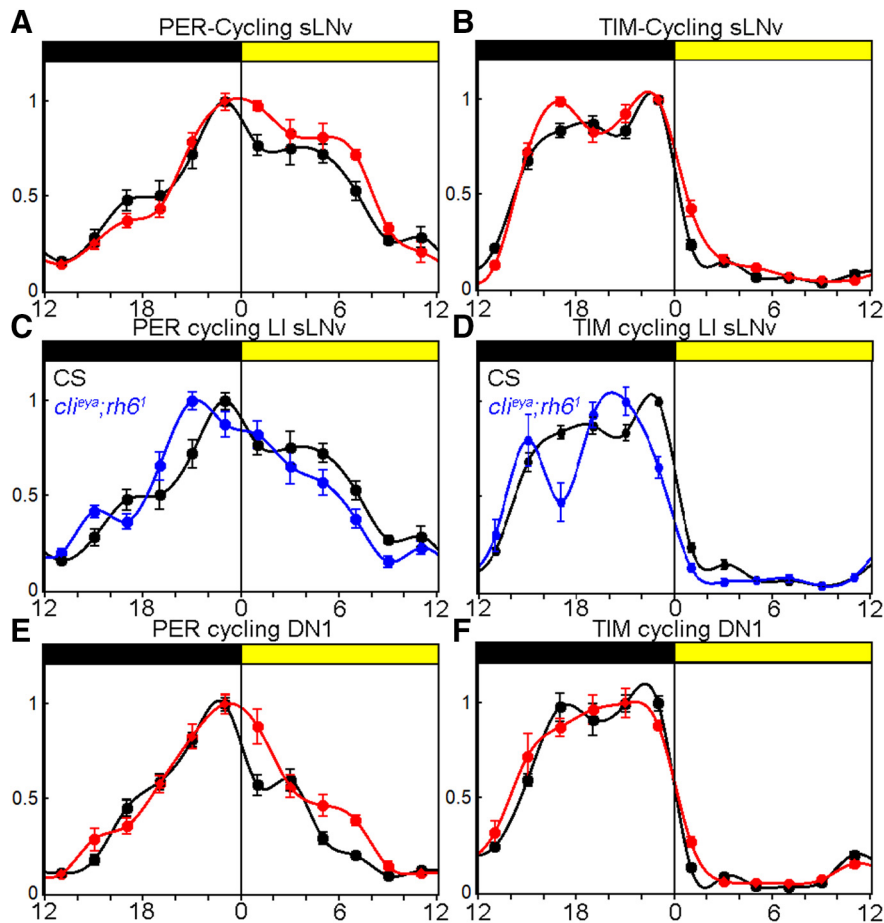


Figure 5. HI changes molecular clock synchronization. **A**, sLN_v PER cycling in WT_{CS} at LI (black line) and HI (red line) ± SEM. PER is stabilized during the daytime at HI. **B**, sLN_v TIM cycling in WT_{CS} at LI (black line) and HI (red line) ± SEM. TIM cycling is identical in both conditions. **C**, sLN_v PER cycling in WT_{CS} and *cli^{eya}; rh6¹* mutants at LI. PER decays faster during daytime and the PER staining intensity maximum is advanced in the mutant. **D**, sLN_v TIM cycling in WT_{CS} and *cli^{eya}; rh6¹* mutants at LI. TIM cycling staining intensity maximum is advanced in the mutant. **E**, DN₁ PER cycling in WT_{CS} at LI (black line) and HI (red line) ± SEM. PER is stabilized during the daytime at HI. **F**, TIM cycling in WT_{CS} at LI (black line) and HI (red line) ± SEM. TIM cycling is identical in both conditions.

neuronal activity, we assayed the s-LN_v dorsal projections, which undergo daily oscillations in fasciculation and volume that reflect changes in neuronal activity: they show an open conformation in the early daytime during times of relatively high activity, whereas the conformation is closed at nighttime during times of low activity (Fernández et al., 2008; Sivachenko et al., 2013). We compared the morphology and the staining intensity of these projections between WT_{CS} and *rh6¹* mutant strains in the middle of the day (ZT6), when they normally show an open conformation.

Indeed, the conformation appeared much more closed at ZT6 in *rh6¹* mutants (Fig. 4E). We quantified this phenotype with the complexity index, which represents the number of pixels stained by the PDF antibody divided by the circumference of the stained area; that is, a smaller index reflects a more closed conformation. We found a significantly lower complexity index in the *rh6¹* mutants compared with WT, suggesting that the s-LN_vs are less active in the mutant. Consistent with this interpretation, the mutants showed a significantly higher PDF staining intensity at ZT6 in the dorsal projections than the WT. Because neuronal firing is required for neuropeptide release, a higher neuropeptide level in the terminals at ZT6 also suggests less firing in the mutant.

Light input significantly alters PER, but not TIM cycling, in WT s-LN_vs

To investigate an effect of light intensity on the molecular circadian clock, we determined PER and TIM levels in clock neurons at 2 h intervals in WT flies under LI and HI conditions. As expected, TIM levels were extremely low during the day, started to accumulate at the beginning of the night, and reached their maximum around ZT20 (Fig. 5B). Also, as expected, PER reached its peak levels toward the end of the night and decayed more slowly than TIM during the day (Fig. 5A). There were no differences in TIM cycling between the two light conditions, most likely due to the extreme light sensitivity of CRY (Vinayak et al., 2013). However, there were significant differences in PER cycling in the s-LN_vs: whereas the increase of PER during the night was unaffected, PER disappeared more slowly at HI, suggesting relative PER stabilization during the daytime degradation phase. To address whether this PER stabilization is due to light input from the visual system, we compared WT PER cycling with *cli^{eya}; rh6¹* mutants at LI (Fig. 5C). As expected, PER was destabilized during the day compared with WT flies. Most interestingly, PER and TIM staining maxima appear to be advanced by ~2 h (Fig. 5C), which corresponds to the advanced E-peak timing in the mutant (Fig. 3C). These data suggest that light input from the visual system stabilizes sLN_v PER in an intensity-dependent manner.

Communication with dorsal neurons mediates the delay in E-peak onset

The s-LN_vs were previously shown to be essential for M activity and free run, but their role in governing the timing of the siesta via a delay in E-activity onset is quite unexpected. This suggests that communication of the s-LN_vs with downstream partners is necessary for proper siesta control and E-peak adjustment. A recent study showed that the DN₁s regulate the siesta via direct inhibition of the E cells (LN_{AS}; Guo et al., 2016). *In vitro* experiments showed that the DN₁s react to bath-applied PDF with an increase of neuronal firing and an increase of cAMP, similar to the positive effect of HB eyelet activation on the s-LN_vs (Seluzicki et al., 2014). In addition, the s-LN_vs send inhibitory PDF signals to the E cells (Liang et al., 2017). Given these strong interactions between the different clock neuron clusters, we expected the same change in PER oscillations in the DN₁s such as that shown above for the s-LN_vs. Indeed, there is more PER during the day the DN₁s at HI (Fig. 5E). Similar to the s-LN_vs, TIM staining remains largely unchanged (Fig. 5F). This indicates a HB eyelet to s-LN_v to DN₁/E cell connection that underlies the HI delay in the E-activity onset.

Discussion

Circadian clocks have evolved as an adaptation to the predictable changes of day and night. To optimally adjust physiology, the

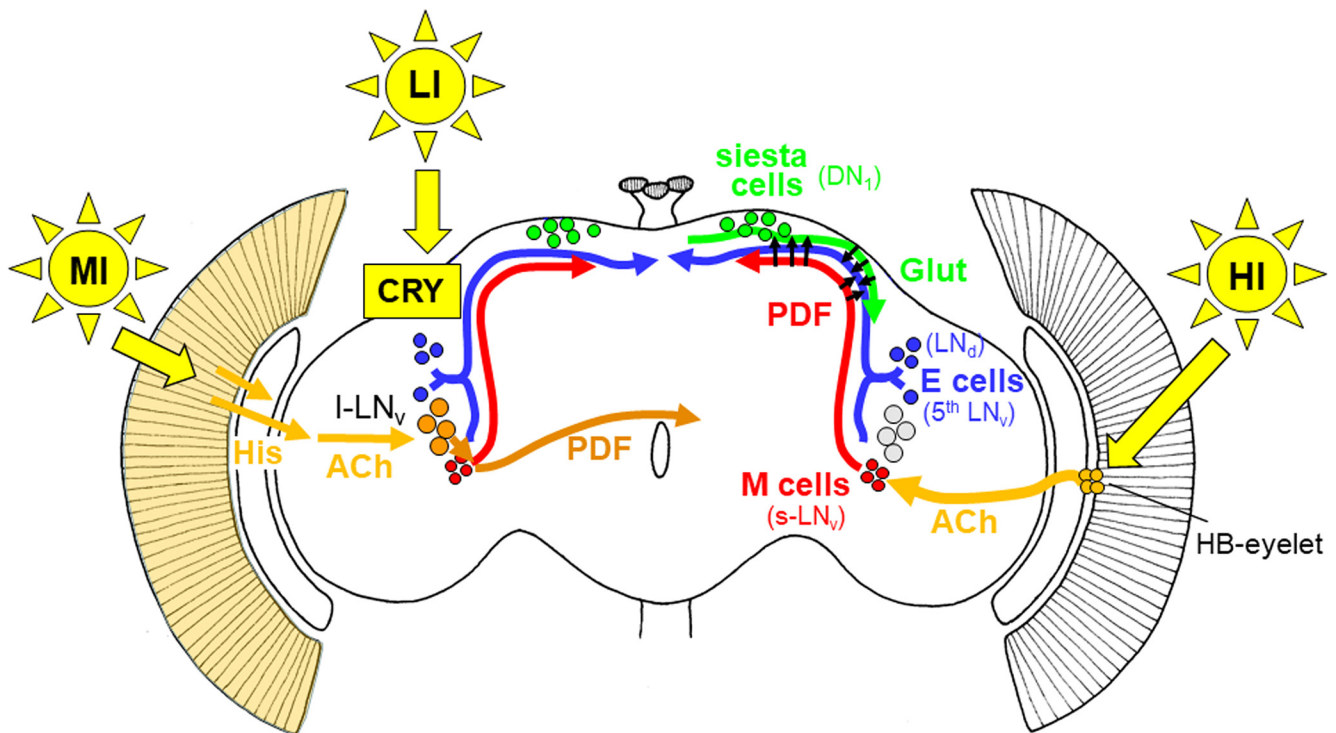


Figure 6. Model for light input to the circadian clock network of *Drosophila*. LI light is sensed by CRY in the clock neurons themselves (indicated in the left brain hemisphere) and can quickly reset their oscillations to light (Vinayak et al., 2013). MI light is perceived by the compound eyes, transferred via histamine (His) to the optic lobes and via acetylcholinergic (ACh) interneurons to the large ventrolateral neurons (I-LN_v; Muraro and Ceriani, 2015; indicated in the left brain hemisphere). The I-LN_v signal via the neuropeptide PDF to the morning (M) and evening (E) cells, respectively (short orange arrow), and to the contralateral hemisphere (long orange arrow). The signaling of HI is shown in the right hemisphere. HI light is perceived by the HB eyelets that signal via ACh to the M cells (s-LN_v; Schlichting et al., 2016). The M cells might signal via PDF directly to the E cells (shown by arrows) and inhibit E-cell activity or to the siesta cells (DN1) in the dorsal brain (Guo et al., 2016; Liang et al., 2017). The siesta cells then signal via glutamate (Glut) to the E cells (3 CRY-positive LN₄ and the fifth sLN₄) and delay the onset of E activity (Guo et al., 2016).

clock should be able to assess and respond to several external cues such as temperature, humidity and light. With respect to the latter, the clock can adapt to different levels of irradiance, at least in part to anticipate seasonal changes. Indeed, previous studies showed that the *Drosophila* clock is extremely light sensitive and can entrain to LD cycles of very low intensity of blue light (Vinayak et al., 2013); this is predominantly due to the photon-integrating function of CRY. In this study, we investigated the role of HI light by simulating LD cycles of 8000–10,000 lux during the day, which is comparable to the light intensity in the shade of a summer day. HI specifically alters the siesta of three WT strains; that is, the duration of the siesta increased by ~1 h and the flies increased their amount of sleep during the day. This behavioral change was accomplished by a delay in the timing of the E-activity onset, which was accompanied by a PI < 1, which represents a reduction of activity 3–6 h before the E peak at HI compared with LI, which we interpret as a delay in E-activity onset. Because all WTs show a reduction in E-peak amplitude, the PI likely reflects a combination of activity suppression and delayed E-peak onset at HI. However, the PI data correlate well with the manually determined timing of E-activity onset, making it an objective measure for fly behavior.

Communication from the s-LN_vs is necessary for this response to HI and the neuropeptide PDF is likely to play an essential role. As shown in Figure 6, it is released in the dorsal part of the brain, where PDF activates a subset of dorsal clock neurons (DNs; Seluzicki et al., 2014). In particular, a subset of the DN₁s release glutamate in response to stimulation, which directly inhibits the LN₄s, resulting in an enhanced siesta (Guo et al., 2016). In addition, PDF appears to inhibit the LN₄s directly (Liang et al.,

2017). In any case, the delayed onset of E activity is reproduced by adult-specific silencing of the E cells, suggesting that s-LN_v firing and PDF release ultimately affect LN₄ firing (Guo et al., 2017).

In contrast to the CRY-dependent extreme sensitivity of the fly clock to LI light, adaptation to HI light is strictly dependent on the visual system. This result parallels the role of the visual system in adapting fly behavior to natural conditions such as twilight or moonlight (Schlichting et al., 2014, 2015). The compound eyes are clearly involved in synchronizing the clock to a wide range of light intensity and to adapt daily behavior to medium-intensity (MI) light, but by investigating different photoreceptor mutants, we excluded the compound eyes as HI sensors; neither flies lacking the eyes (*cli^{eya}*) nor lacking the important visual system neurotransmitter histamine (*hdc^{JK910}*) fail to adapt to HI. In contrast, flies lacking Rh6 cannot respond to HI. Because this is the only rhodopsin expressed in the HB eyelets and given that the phenotype is reproduced in the *cli^{eya} rh6¹* background, the data indicate that it mediates HI light input to the clock (Sprecher and Desplan, 2008).

The HB eyelets are a major source of visual information in larvae and persist throughout the pupal stage into adulthood. They undergo changes in photopigment (Rh5 to Rh6) as well as in neurotransmitter (ACh to ACh and His) during metamorphosis and are conserved in other fly species such as the blowfly (Sprecher and Desplan, 2008). The HB eyelets directly innervate the adult accessory medulla, which is an important pacemaker center of other insects and contains dendrites of the LN_vs in *Drosophila* (Helfrich-Förster et al., 2002; Malpel et al., 2002).

Previous studies implicated the eyelets in adult entrainment. For example, flies lacking CRY and neurotransmitter output

from the entire visual system completely fail to re-entrain to a phase delay of 6 h, whereas functional eyelets alone allows 50% of the flies to re-entrain (Veleri et al., 2007). Similarly, our data here show that flies lacking HB eyelet function cannot delay the E-activity onset in response to HI; this delay may therefore resemble a more traditional circadian phase delay.

It is interesting to contrast these results with mammalian entrainment. Very low irradiance entrainment is predominantly performed by the rods due to their high light sensitivity. The circadian photoreceptor melanopsin functions predominantly at high intensities, which saturate rod and cone signaling (Lall et al., 2010). Here, we show that there are also different pathways for LI and HI light in *Drosophila*: CRY mediates entrainment to extremely low irradiances, whereas the HB eyelet mediates HI adaptation. This suggests some similarity if not an evolutionary relationship between the HB eyelet and the melanopsin-containing retinal ganglion cells of mammals.

Another goal of this study was to find the neuronal pathways downstream of the eyelets. In a previous study, we showed that eyelet activation with the P2X2 system leads to increases of Ca^{2+} and cAMP in the s-LN_vs (Schlichting et al., 2016). If this connection is relevant to HI adaptation, then we expected to see Ca^{2+} increases in the s-LN_vs in response to HI. To this end, we used the previously described Tric-GFP tool and showed that s-LN_v Ca^{2+} levels are indeed higher at HI than at LI.

We suggest that this Ca^{2+} increase reflects increased s-LN_v activation by the HB eyelet and HI illumination. This interpretation is buttressed by the fact that *rh6*¹ mutants retain a closed conformation of the s-LN_v dorsal projections, which is known to reflect relatively low s-LN_v activity (Fernández et al., 2008).

s-LN_v activation leads to release of neuropeptides/neurotransmitters at its axonal terminals. This also explains the behavioral phenotype of *rh6*¹ mutants: as described previously, they show a reduced M-peak amplitude and an advanced E-peak timing (Schlichting et al., 2014). The latter is present even more strongly in the complete loss-of-function *pdf*⁰¹ mutant strain (Renn et al., 1999). This suggests that *rh6*¹ mutant phenotype resembles that of a partial PDF loss-of-function; that is, the s-LN_vs do not release as much PDF in the dorsal brain as WT flies, as evident by still high PDF staining intensity in the dorsal brain of *rh6*¹ mutants in the middle of the day.

Circadian second messengers have previously been implicated in PER expression/stabilization, especially downstream of PDF signaling (Li et al., 2014). We therefore investigated changes of s-LN_v PER and TIM cycling in HI versus LI. Whereas TIM expression was not detectably different between the two conditions, PER staining was enhanced and shifted into the daytime by HI, consistent with PER stabilization. This delay of the PER staining profile nicely correlates with the ~1 h delayed onset of the E activity. The response to HI also fits with observations under long photoperiods: If one compares PER and TIM cycling in LD 12:12 and LD 18:6, only the maximum of PER staining intensity is significantly delayed under long day conditions. The E activity peak is similarly delayed (Menegazzi et al., 2013). Although this suggests that a similar mechanism regulates the integration of light intensity and exposure to long photoperiods, the molecular relationship between the PER cycle and behavior is unknown and awaits further study.

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