Communication Among Photoreceptors and the Central Clock Affects Sleep Profile

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Light is one of the most important factors regulating rhythmical behavior of Drosophila melanogaster. It is received by different photoreceptors and entrains the circadian clock, which controls sleep. The retina is known to be essential for light perception, as it is composed of specialized light-sensitive cells which transmit signal to deeper parts of the brain. In this study we examined the role of specific photoreceptor types and peripheral oscillators located in these cells in the regulation of sleep pattern. We showed that sleep is controlled by the visual system in a very complex way. Photoreceptors expressing Rh1, Rh3 are involved in night-time sleep regulation, while cells expressing Rh5 and Rh6 affect sleep both during the day and night. Moreover, Hofbauer-Buchner (HB) eyelets which can directly contact with s-LNvs and l-LNvs play a wake-promoting function during the day. In addition, we showed that L2 interneurons, which receive signal from R1-6, form direct synaptic contacts with l-LNvs, which provides new light input to the clock network.

Keywords: sleep, photoreceptors, Hofbauer-Buchner eyelets, Drosophila, peripheral clock

INTRODUCTION

Circadian rhythms in Drosophila are regulated by a system of oscillators, which includes the pacemaker located in the central brain and peripheral oscillators located in various cells, tissues, and organs. Peripheral oscillators such as the ones in glial cells, compound eyes, antennae, gustatory receptor neurons, or Malpighian tubules express clock genes and show circadian rhythms in their structure and physiological processes (Siwicki et al., 1988; Zerr et al., 1990; Hege et al., 1997; Krishnan et al., 1999; Chatterjee and Hardin, 2010; Chatterjee et al., 2010).

The compound eye consists of ommatidia and each of them contains eight photoreceptors. Six of them, R1–R6, are located in the distal retina and express rhodopsin 1 (Rh1), sensitive to a broad spectrum of light wavelengths (O’Toola et al., 1985; Hardie, 1987). R1–R6 terminate in the lamina, where they form tetrad synaptic contacts with L1, L2, L3, and amacrine cells (Meinertzhagen and O’Neil, 1991; Meinertzhagen et al., 1997). The R1–R6 photoreceptors are involved in motion detection and image formation (Rüster et al., 2007; Yamaguchi et al., 2008). Two other photoreceptors of each ommatidium, R7 and R8, are involved in color vision and detection of polarized light. They terminate in the second optic neuropil (medulla), where they contact transmedulla neurons (Tm5, Tm9, and Tm20; Gao et al., 2008; Melnattur et al., 2014). Tm5, Tm20 neurons receive also indirect input from R1–R6, through L3 (Gao et al., 2008; Melnattur et al., 2014). R7 forms additional synaptic contacts with amacrine cell Dm8 (Gao et al., 2008), and
moreover, R7 and R8 contact each other in the medulla through direct synaptic contacts (TakeMurra et al., 2015), most probably using histamine as a neurotransmitter (Hardie and Raghu, 2001; Pantazi et al., 2008; Schnaitmann et al., 2018). R7 photoreceptors express UV-sensitive rhodopsin 3 (Rh3) or blue light—absorbing Rh4, while R8 cells express Rh3, blue light-sensitive Rh5 or green light-absorbing Rh6 (Salcedo et al., 1999).

According to the rhodopsin type expressed in R7 and R8, three different subpopulations of photoreceptors have been described: “pale” ommatidia consist of Rh3-expressing R7 and Rh5-expressing R8, the “yellow” type is composed of Rh4 expressing R7 and Rh6-expressing R8, and “DRA” (dorsal rim area) expresses Rh3 in both R7 and R8 and is involved in the polarized light detection (Wernet et al., 2006). An additional photoreceptive structure in the visual system is the Hofbauer–Buchner (HB) eyelets (Hofbauer and Buchner, 1989; Yasuyama and Meinertzhagen, 1999) composed of 4 cells expressing Rh6, terminating in the accessory medulla (Helfrich-Förster et al., 2002).

Different photoreceptor types and photopigments seem to play different roles in the circadian rhythm and behavior regulation. R1–R6, expressing Rh1, play a role in dim light detection in motion vision and phototaxis; they are also important for nocturnal activity which increases in response to raising day-light activity (Schlichting et al., 2014). Rhodopsin 6, expressed in a population of R8 photoreceptors, plays a role in the integration of light signals received by the other photoreceptors (Saint-Charles et al., 2016), while Rh5-expressing cells seems to be involved in light entrainment, that is the clock ability to adapt to light cycles and phase shifts of the rhythm, which suggests that Rh5 may use NorpA-independent pathway (Saint-Charles et al., 2016). Finally, HB eyelets play a role in regulating evening onset under high intensity light conditions, as well as the length of the siesta (Schlichting et al., 2019).

The role of the visual system in circadian entrainment has already been studied (Nippe et al., 2017), however, the effect of different photoreceptors on sleep is still not fully recognized. In this study we examined the role of different photoreceptors and their post synaptic targets in the regulation of sleep and locomotor activity. We have shown that Rh1 and Rh3-expressing photoreceptors affect sleep during the night, while Rh5 and Rh6-expressing cells, both during the day and night. Moreover, we have presented that photoreceptors influence the pace of the molecular clock in pacemaker cells and that retinal oscillators are as much important for maintaining sleep as synaptic transmission from photoreceptors to target cells. Finally, we have described connections between the visual system and clock cells.

**MATERIALS AND METHODS**

**Fly Strains**
The following strains of *Drosophila melanogaster* were used in the present study: GMR-Gal4, Rh1-Gal4, Rh3-Gal4, sp;CyO; Rh5-Gal4, yw; Sp;CyO; Rh6-Gal4/TM6B, R8212AD;CyO; R75H08BD/TM6B (herein called L2-Gal4), yw; Pdf-Gal4, w; UAS-Acroc24, w; UAS-TeTxLC (herein called UAS-TeTx), w; Pdf-LexA,AS-GFP1;CyO; LexAop-GFP,11 (for GRASP experiment), yw,UAS-myrGFQUAS-HA::RFP; transTANGO (herein called transTANGO), w; UAS-CAmp5f, w; Pdf-LexA,LexAopGFP,11; UAS-Nrt::GFP,1;0 (for Nrx GRASP experiment).

L2-Gal4 strain was obtained from Janelia Research Campus, the others from Bloomington Drosophila Stock Center.

Flies were maintained under 12 h of light and 12 h of darkness (LD12:12) conditions and at a constant temperature of 25°C, unless the procedure required constant darkness (DD).

**Recording Locomotor Activity and Sleep**
The locomotor activity was recorded using a monitoring system (TriKinetics) composed of monitors equipped with infrared light-emitting diodes and detectors, connected to a computer. Each monitor houses 32 glass-tubes of a diameter just sufficient to maintain a single fly. Tubes are sealed at both ends: one by food and the other by a foam stopper. When the fly passes the emitter/detector pair, the infrared beam is interrupted resulting in a signal transmitted to the computer. To analyze circadian rhythms in locomotor activity, flies were maintained for 7 days under LD12:12 and next under constant darkness (DD) for next 7 days.

Activity was counted every 5 min (1 bin) and analyzed in Excel by using “Belly!” software (Department of Genetics, University of Leicester). Lomb–Scargle normalized periodogram was used to determine rhythmic flies; flies with period value lower than 10 (confidence level 0.05) were regarded as arrhythmic. Flies which did not survive until the end of experiments were removed from analyses. Every experiment was repeated three times, at least 60 flies in total were used.

The Anticipation Index (Morning and Evening) was calculated for individual flies under LD12:12, by determining the proportion of activity counts during the 3 h preceding the phase transition over the activity within 6 h preceding phase transition.

To study sleep pattern, activity of flies was analyzed in the second day in LD12:12. Sleep was measured as intervals of at least 5 min of inactivity.

Heterozygous parental strains were used as control. In case of gene silencing, progeny of driver line crossed with UAS-VALIUM10-GFP was used as additional control. This strain has expression of empty VALIUM10-GFP vector in targeted cells.

Statistical analysis was performed using ANOVA with a Tukey’s multiple comparison test for normally distributed data. To analyze rhythmicity of flies we used non-parametric Kruskal–Wallis test to compare percentage from three repetitions. GraphPad Software was used to performed statistical analysis.

**Immunohistochemistry**
Flies were decapitated and their heads were fixed in 4% paraformaldehyde in phosphate buffer saline (PBS; pH 7.4) for 4 h, then they were washed in PBS twice and cryoprotected by incubation in 12.5% sucrose for 10 min and in 25% sucrose at 4°C overnight. Material was embedded in Tissue Tek, frozen in liquid nitrogen, and cryostat 20 μm sections were cut. The sections were washed in PBS for 30 min and five times in phosphate buffer with an addition of 0.2% Triton X100 (PBT). After that,
sections were incubated in 5% normal goat serum (NGS) with an addition of 0.5% bovine serum albumin (BSA) for 30 min at room temperature. Next they were incubated with primary antibodies for 24 h. Afterwards, sections were washed six times in PBT/BSA, blocked in 5% NGS for 45 min and secondary antibodies were applied for overnight incubation at 4°C. Finally, sections were washed twice in BSA, six times in PBT, and twice in PBS. Then, cryosections were mounted in Vectashield medium (Vector) and examined with a Zeiss Meta 510 Laser Scanning Microscope.

In addition to sections of the brain, whole brains were also used for immunohistochemistry. They were isolated after 1 h of head fixation in 4% PFA, washed in PBS and fixed again for the next 45 min. The next steps of immunostaining were done according to the protocol described above for cryosections.

For immunohistochemistry the following antibodies were used: nc82 (against the presynaptic protein Bruchpilot) (1:20, Developmental Studies Hybridoma Bank), PDF C7 (against Pigment Dispersing Factor) (1:500, Developmental Studies Hybridoma Bank), anti-GFP (rabbit, 1:1000, Novus Biological), anti-GFP (mouse, 1:20, Sigma Aldrich), goat anti-mouse conjugated with Cy3 (1:500, Jackson ImmunoResearch Laboratories, Inc.), goat anti-rabbit conjugated with Alexa 488 (1:1000, Molecular Probes), goat anti-mouse conjugated with Cy2 (1:500, Abcam).

**Calcium Imaging**
L2→GCaMP6f flies were dissected on ice at specific time points, and brains were placed in PBS. Images of the brain were collected immediately with a confocal microscope. The fluorescence intensity was measured using ImageJ software. The ratio of fluorescence per area was calculated using ImageJ macro. Data obtained at different time points were compared.

**TransTANGO**
L2-Gal4 or GMR-Gal4 crossed with transTANGO flies were raised at 18°C, adult males were separated and aged for 15 days at 18°C. The ICCs procedure was the same as described above, except for the length of the incubation with primary antibody, which was extended to 5 days at 4°C. The following primary antibodies were used: rabbit anti-DsRed (1:250, Rockland), chicken anti-GFP (1:500, Aves Labs) and rat anti-PDF (1:250; Depetriz-Chauvin et al., 2011). The following secondary antibodies were used: Cy2-conjugated anti-chicken, Cy5-conjugated anti-rat, and Cy3-conjugated anti-rabbit (1:250, Jackson ImmunoResearch Laboratories, Inc.). Images were acquired with a ZEISS LSM 880 Confocal Laser Scanning Microscope.

**RESULTS**

**Retina Photoreceptors Are Involved in Sleep Regulation**
To analyze the contribution of specific photoreceptors to the regulation of rhythmic behavior we first examined the impact of blocking light input through the retina. In order to do so we took advantage of the GMR→TeTx strain, in which neurotransmission from cells expressing glass was totally blocked by expression of tetanus toxin light chain, which cleaves synaptobrevin protein and blocks neurotransmitter release. GMR→TeTx flies showed no defects in rhythmicity or periodicity, but the evening anticipation was increased (Table 1). Interestingly, these flies exhibited an altered sleep profile (Figure 1A) with increased sleep time during the day (Figure 1B), resulting in lower levels of total activity (Supplementary Figure S1A).

The retina photoreceptors are peripheral oscillators with rhythmic expression of clock genes. To check whether retinal clocks are involved in the network regulating sleep in flies, we used GMR→Acyc24 strain, in which expression of the dominant negative form of CYCLE causes disruption of the molecular clock in photoreceptors. We found that this manipulation triggered arrhythmicity in 55% of GMR→Acyc24 flies in constant darkness conditions (Table 1) and in rhythmic flies, the period was shorter (22.9 h) compared to the control (Table 1). Moreover these flies exhibited an increased morning anticipation index (Table 1), altered sleep pattern and length during the day and night (Figures 1C,D), and a decreased total activity (Supplementary Figure S1A). Knowing that cyc dominant negative overexpression in inn-Gald cells is lethal (Chow et al., 2016), we carefully examined the eye morphology and anti-BRP labeling in the brain to exclude a possibility that the observed results originate from some neuronal degeneration. However, we did not observe any changes in the examined individuals.

These two experiments showed that glass-expressing cells, which are mostly retinal photoreceptors, are involved in the regulation of sleep, while they are not necessary to maintain the locomotor activity rhythm, as it has already been shown (Grima et al., 2004).

**The Clock Located in the Retina Photoreceptors Regulates Their Own Circadian Output**
To investigate how oscillators located in the retina transmit rhythmic signals to the deep brain, we looked at the effect of peripheral clock disruption on the presynaptic protein Bruchpilot (BRP) cycling in the photoreceptor terminals in the lamina. Under LD12:12 BRP levels oscillate daily with two maxima observed at the beginning of the day (ZT1) and at the beginning of the night (ZT13) (Figure 2A; Górski-Andrzejak et al., 2013). Interestingly, in our study in GMR→Acyc24 flies the expression was changed, the BRP level at ZT1 was not significantly different than at ZT4 and ZT16, and only one peak at the beginning of the night (ZT13) was observed (Figures 2B,C). This result is in accordance to what was already described for DD conditions, reinforcing the notion that morning BRP peak is controlled by light (Górski-Andrzejak et al., 2013). However, the fluorescence intensity level at ZT1 in the experimental and control flies was similar, and enhanced at ZT4 and ZT16, which may suggest that BRP degradation rather than expression is affected.

We then explored the effect of peripheral clock located in the photoreceptors on the pace of the main oscillator, by blocking neurotransmission from the retina photoreceptors (GMR→TeTx) or disrupting the clock in these cells (GMR→Acyc24). Strikingly,
### TABLE 1 | Locomotor activity of flies with disrupted synaptic transmission (TeTx) or molecular clock (Δacyc24) in the specific type of cells in the visual system.

| Genotype          | Period [h] | % Rhythmic | MAI   | EAI   | Number of flies |
|-------------------|------------|------------|-------|-------|-----------------|
| UAS-cycA24/+      | 23.7       | 93         | 0.63  | 0.76  | 88              |
| UAS-TeTx/+        | 24.0       | 94         | 0.56  | 0.79  | 84              |
| UAS-ChATRNAi/+    | 23.3       | 100        | 0.69  | 0.65  | 58              |
| GMR-Gal4/+        | 23.9       | 99         | 0.54  | 0.74  | 92              |
| GMR>Δacyc24       | 22.9****   | 45*        | 0.64  | 0.91**** | 91             |
| GMR>TeTx          | 23.8       | 95         | 0.52  | 0.79  | 91              |
| GMR>Acyc24        | 23.7       | 77         | 0.7   | 0.79  | 92              |
| Rh1-Gal4/+        | 23.5       | 100        | 0.79**** | 0.8   | 91              |
| Rh1>Δacyc24       | 23.3       | 83         | 0.58  | 0.79  | 95              |
| Rh1>TeTx          | 24.2       | 75         | 0.73  | 0.81  | 95              |
| Rh1>Acyc24        | 23.9       | 45*        | 0.63  | 0.79  | 70              |
| Rh1>TeTx          | 24.3       | 89         | 0.57  | 0.71  | 70              |
| Rh1>Δacyc24       | 24.3       | 97         | 0.79  | 0.71  | 89              |
| Rh2-Gal4/+        | 23.7       | 78         | 0.63  | 0.7   | 78              |
| Rh2>Δacyc24       | 23.6       | 86         | 0.58  | 0.79  | 74              |
| Rh2>TeTx          | 23.6       | 66         | 0.6   | 0.72  | 74              |
| Rh2>Val10         | 23.4       | 96         | 0.58  | 0.78  | 55              |
| Rh6-ChatRNAi      | 23.7       | 86         | 0.52  | 0.88** | 73             |
| L2-Gal4/+         | 24.0       | 96         | 0.65  | 0.87**** | 91          |
| L2>Δacyc24        | 24.0       | 92         | 0.64  | 0.8   | 90              |

Columns present: genotype, period of locomotor activity, % of rhythmic flies, morning anticipation index (MAI), evening anticipation index (EAI), and total number of flies used for the experiments. Experimental flies which showed differences with both controls are highlighted with bold. Statistically significant differences marked with asterisks (*p ≤ 0.05; ****p ≤ 0.0001). Detailed statistics are presented in Supplementary Tables S1, S2.

### FIGURE 1 | Effects of glass-expressing cells on sleep.

(A) Sleep pattern of flies with blocked neurotransmission from photoreceptors (GMR>TeTx) measured as number of sleep bins per hour. (B) Total sleep time of GMR>TeTx flies measured as minutes per 12 h, separately for day and night-time. (C) Sleep pattern of flies with clock disruption in photoreceptors (GMR>Δacyc24). (D) Total sleep time of GMR>Δacyc24 flies. Heterozygous parental strains were used as control. Statistically significant differences marked with asterisk *p ≤ 0.05; **p ≤ 0.01; ****p ≤ 0.0001. Detailed statistics are presented in Supplementary Tables S3, S4.
we found that GMR>TeTx and GMR>Δyc24 flies display a clear dampening of PER oscillations in the small and large LNvs (Figures 3A–D, respectively). Thus, blocking neurotransmission from the retina photoreceptors and disruption of the clock in glass-expressing cells decreased the amplitude in PER cycling in essential pacemaker neurons (Figure 3).

**Specific Photoreceptor Types Regulate Sleep at Different Ways**

Because our experiments with GMR strain showed changes in sleep pattern and level, we focused on this behavior in the next experiments. GMR expression is not limited to the retina, however. In fact, GMR is expressed in some clock neurons (DN1p; Klarfeld et al., 2004), which are involved in the sleep regulation (Guo et al., 2018; Lamaze and Stanewsky, 2020). To investigate which cell types triggered the observed responses, we expressed tetanus toxin in different types of photoreceptors, using various rhodopsin drivers, which allowed us to exclude the effect of DN1p. Surprisingly, we obtained different effects depending on photoreceptor types. In general, blocking input from different photoreceptor types resulted in decreased total activity, but their effects on sleep and morning/evening peaks were different. No changes in the period of locomotor activity were observed (Table 1).

After blocking light transmission pathway from R1–R6 cells (Rh1-expressing; Figures 4A,B) we observed clear changes in the sleep pattern (Figure 4C) with significant longer sleep time during the night, no effect on day-time sleep (Figure 4D), and a subtly reduced evening peak (Table 2 and Supplementary Figure S2A). These effects were opposite to those observed in case of GMR; nevertheless, in this case a decrease of total activity was also observed (Supplementary Figure S1B).

Similar effects were obtained after disrupting the clock in the R1-6 photoreceptors (Figure 4E), despite the impact on night-time sleep was less pronounced (Figure 4F and Supplementary Figure S3A).

We next inquired the relevance of the R7-8 photoreceptors. The block of neurotransmission from the Rh3-expressing cells, reaching the medulla (Figures 5A,B), had strong effect on the activity patterns, with an increased morning peak and a decreased evening peak (Table 2 and Supplementary Figure S2B). The sleep pattern was also affected (Figure 5C), with increased night-time sleep (Figure 5D), and decreased total activity (Supplementary Figure S5C). Similarly, the clock disruption in these cells resulted in a decreased total activity, although the impact on night-time sleep was less pronounced (Figures 5E,F and Supplementary Figures S1C, S3B).

Next we examined the contribution of Rh5-expressing R8 “pale” ommatidia (Figures 6A,B). Rh5>TeTx flies exhibited reduced total activity (Supplementary Figure S1D) and both morning and evening activity peaks were decreased compared to controls (Table 2 and Supplementary Figure S2C). The sleep pattern changed (Figure 6C) with increased sleep time in both, day and night (Figure 6D). Overexpression of cycA24 in “pale” R8 cells recreated the effects linked to TeTx expression: decreased total activity (Supplementary Figure S1D), blunted morning and evening peaks (Table 2 and Supplementary Figure S3C) and changes in the sleep pattern (Figure 6E) and level during the day and night (Figure 6F).
Rh6 is expressed in the R8 “yellow” ommatidia as well as in the HB eyelets (Figures 7A,B). Blocking the synaptic transmission from these cells did not affect overall activity levels (Supplementary Figures S1E, S2D) and the sleep was slightly extended at the beginning of the day and during the night (Figures 7C,D). Clock disruption in this type of photoreceptors only mirrored the one triggered during the day (Figures 7E,F and Supplementary Figure S3D).

To distinguish between the contribution of R8 photoreceptors and the HB eyelets, both expressing Rh6, we knocked down the expression of choline acetyltransferase, necessary for the synthesis of acetylcholine, used as neurotransmitter by the HB eyelets. In Rh6>ChATRNAi flies changes in sleep pattern and increased sleep time became evident only during the day (Figures 7G,H).

Taking these data together, we can conclude that R1-6 cells, as well as Rh3 and Rh6-expressing retinal photoreceptors affect sleep during the night, while R8 expressing Rh5 is involved in both day- and night-time sleep regulation. Moreover, cholinergic HB eyelets play a wake-promoting role during the day.

**L2 Interneurons as an Additional Peripheral Clock Output**

Since light signals are transmitted to the pacemaker neurons from the retina photoreceptors via the lamina interneurons, we examined the behavior of flies with blocked neurotransmission in L2 interneurons, which receive light input from R1-R6 cells (Figure 8A). We focused on this cell type because it shows rhythmic changes in the size of dendritic trees, probably connected with daily changes in signal transmission. Moreover, L2 terminals are located in a close vicinity to s-LNv's (Figure 8B). These flies displayed behavioral changes reminiscent of GMR>TeTx flies, with a smaller morning peak of activity (Table 2). The sleep pattern of these flies was changed (Figure 8C) with increased sleep duration during the day and night (Figure 8D).

To correlate daily changes in the size of dendritic trees' activity with that of L2 cells, we carried out calcium imaging using L2>GCaMP6f transgenic strain. GCaMP6f is a calcium indicator, which allows to measure Ca^{2+} level correlated to the indicator fluorescence intensity. In our experiment we isolated brains at selected time points and measured the fluorescent signal immediately after dissection. The fluorescent intensity in the L2 terminals was measured in the medulla (Figure 8E) and calculated per terminal area, comparing Ca^{2+} level/area unit time points. In LD12:12, Ca^{2+} level was highest at night (ZT16, ZT20) and lowest in the middle of the day (ZT8; Figure 8F). In DD this pattern subtly changed, with the highest intensity signal at CT16 and CT20, but without the lowest level at ZT8 (Figure 8G).

The obtained results suggest that L2 interneurons play an important wake-promoting role. It is supported by the fact that calcium levels in the terminals are the lowest during siesta.

**Photic Inputs to the Pacemaker May Be Regulated by Synaptic Plasticity**

Some photoreceptor terminals are located next to the clock neurons in a region called the accessory medulla (Figures 9A,B). It was previously shown that the visual system can directly communicate with clock neurons and receive photic information through the HB eyelets (Muraro and Ceriani, 2015; Schlichting et al., 2016). Taking advantage of the GFP reconstitution across synaptic partners (GRASP) technique (Feinberg et al., 2008)
we found that the HB eyelets terminals differentially contact the LN\\textsubscript{vs} during the day, with stronger contacts during light phase (Figure 9C). To confirm that these results correlate with differential synaptic connectivity, we used Nrx GRASP to visualize only active synaptic contacts (Figures 9D,E). At ZT1 100% of brains showed reconstituted signal (n = 20), the
FIGURE 5 | Rh3-expressing cells affect night-time sleep. (A) Rh3 is expressed in R7 and R8 cells, which terminate in the medulla (cryosection of Rh3>GFP brain). (B) Graphical presentation of synaptic connections formed by Rh3-expressing cells which are blocked in Rh3>TeTx flies (Tm - transmedulla neurons). (C) Sleep pattern of Rh3>TeTx flies. (D) Sleep time during the day and night of Rh3>TeTx flies. (E) Sleep pattern of Rh3>hcyc24. (F) Amount of sleep during the day and night of Rh3>hcyc24. Heterozygous parental strains were used as control. Statistically significant differences marked with asterisks **p < 0.01; ***p < 0.001; ****p < 0.0001. Detailed statistics are presented in Supplementary Table S3.

The proportion of brains decreased at other time points, i.e., 43% at ZT4 (n = 21), 59% at ZT13 (n = 22), and 35% at ZT16 (n = 20), suggesting that the LNvs receive direct input from the HB eyelets in a plastic manner, preferentially during the early morning.

The HB eyelets terminals arborize in a close vicinity to the s-LNvs cell bodies. Taking into account the localization of the reconstituted signal, it is likely the HB eyelets communicate with the s-LNvs rather than the l-LNvs.

To gain insight into how the information from peripheral oscillators is transmitted to the main clock we took advantage of transTANGO (Talay et al., 2017) to uncover postsynaptic cells to photoreceptors (GMR-Gal4) and L2 interneurons (L2-Gal4; Figures 10, 11, respectively). This tool uses a modified signaling pathway to express RFP in the postsynaptic cells of a GAL4 of interest, while marking the cells recruited by that driver with GFP. In pursuit of the postsynaptic targets to GMR+ photoreceptors we observed red fluorescence in the lamina and medulla, probably coming from the lamina interneurons and amacrine cells. The signal was also detected in a few cell bodies in the accessory medulla, which project to the dorsal brain and to the contralateral side of the brain. Double staining with antibodies against PDF confirmed that a subset of RFP-expressing cells belong to both the s-LNvs and l-LNvs groups and to processes described as the posterior optic tract (POT), which connect the l-LNvs cluster located on both sides of the brain, and dorsal projections of the s-LNvs. Closer inspection of the GFP signal indicated that the cells contacting LNvs are those of the HB eyelets (Figure 10C), confirming what is depicted in the recently published connectome (Scheffer et al., 2020).

Despite no direct contact between L2 neurons and LNvs was detected through GRASP (Muraro and Ceriani, 2015), transTANGO mapping of L2 postsynaptic targets highlighted the l-LNvs, but not the s-LNvs, in the majority of brains (86% positive, n = 7) (Figure 11). This l-LNvs exclusive connection could help explain why disrupting the connectivity or the clock in the retinal peripheral oscillator has a stronger effect on the sleep pattern than in other circadian outputs examined.

DISCUSSION

The circadian clock relies on a self-sustained molecular mechanism, which is entrained to the daily changes of environmental conditions. The most powerful factor
synchronizing circadian clocks is light. The LNvs, essential circadian pacemakers, receive light information through the visual system and the deep brain photoreceptor CRY (Emery et al., 2000; Sheeba et al., 2008a).

The involvement of the visual system is still not completely understood. Pacemaker cells are responsible for the temporal organization of rhythmic behavior, such as locomotor activity and sleep but also contribute to other features such as sleep length during the day and night and total activity level (Parisky et al., 2008; Shang et al., 2009; Potdar and Sheeba, 2018). In this study we focused on the role of retinal and extra-retinal photoreceptors in the transduction of photic information to modulate sleep patterns. We uncovered that peripheral clocks in the eye contribute to maintaining sleep pattern with little effects on the period of locomotor activity rhythm, as expected (Grima et al., 2004).

Clock disruption in glass-expressing cells triggered arrhythmicity, and in the remaining rhythmic flies, a shorter period of the locomotor activity pattern. Surprisingly, blocking synaptic transmission from photoreceptors to postsynaptic cells did not affect rhythm or periodicity, however, observed changes in the sleep pattern and level were similar in both experiments. Subsets of clock neurons employ CRY for light entrainment, while the others use CUL-3 mediated mechanism for molecular light resetting (Ogutea et al., 2020). However, it seems that they need rhythmic inputs from the visual system, as arrhythmic signaling may affect the molecular mechanism of pacemaker neurons.

The visual system shows daily changes of sensitivity to light. In addition, the clock-controlled optomotor response in Drosophila is higher at night than during the day (Mazzotta et al., 2013; Damulewicz et al., 2017). In the first neuropil (lamina) of the optic lobe, circadian rhythms have been observed at the cellular (Weber et al., 2009) and molecular levels (Górski-Andrzejak et al., 2013) including the expression of several genes and proteins, for example the alpha subunit of the sodium/potassium pump in the lamina glia (Górski-Andrzejak et al., 2009; Damulewicz et al., 2013).

In the present study, we showed that not only sensitivity of photoreceptors, but also interneurons involved in light transmission changes during the day. L2 monopolar cells, one of the postsynaptic cells in tetrad synapses, which hyperpolarize in response to light, show daily rhythms in calcium levels. The highest Ca2+ concentration in the terminals was observed at night, similarly to the retina photoreceptors. It seems that the visual system is more sensitive to light during the dark phase to detect low intensity light, such as moonlight, and such sensitivity decreases at the end of the night to be prepared for high intensity...
of light in the morning (Shang et al., 2008; Sheeba et al., 2008b; Nippe et al., 2017).

We showed that rhythms observed in the lamina, like the presynaptic protein BRP expression, is controlled by both the pacemaker and the oscillators located in the retina. Moreover, the BRP expression pattern in flies in which the peripheral clock in the retina is disrupted, is unimodal, similar to that observed in constant darkness, when the morning peak of BRP is missing and only the evening peak is observed (Górski-Andrzejak et al., 2013). However, in our experimental model BRP level at ZT1 was similar to those observed in control, which suggests that clock disruption affects rather degradation of BRP than its expression.

Despite the strong effect on sleep of genetic manipulations in photoreceptors using the GMR-Gal4 driver, the glass gene is also expressed in some DN1s and in a group of cells located next to LNds (Yoshall and Young, 1995; Klarfeld et al., 2004), opening the possibility that the results obtained could not be exclusively eye-specific (Li et al., 2012; Ray and Lakhotia, 2015). To avoid
multi-cellular effects, we focused on data performed with drivers specific to different photoreceptor types, and we observed various effects on sleep, depending on the photoreceptor type.

The previous analysis employing the norpA mutant, which lacks phospholipase C in the canonical phototransduction pathway (Pearn et al., 1996), showed that light signals to the clock are transmitted not exclusively from the retina photoreceptors, but also by the circadian photoreceptor CRY and the HB eyelets (Emery et al., 2000; Helfrich-Förster et al., 2001). However, norpA mutants are not able to entrain to changes in light conditions, because of reduced circadian sensitivity to light (Emery et al., 2000). Moreover, in the double mutant norpA; cryB, PER cycling was maintained in s-LNv,s and DN1,s but abolished in l-LNv,s and LNds, suggesting that these clusters are photoentrained by an alternative pathway (Helfrich-Förster et al., 2001). This conclusion is further supported by the fact, that an alternative
A norpA-independent phototransduction pathway occurs in Rh1, Rh5, and Rh6-expressing photoreceptors (Shortridge et al., 1991; Szular et al., 2012; Ogueta et al., 2018). Flies with blocked synaptic transmission using GMR>TeTx resembled the norpA mutant. Since this genetic combination blocks synaptic transmission between photoreceptors, R1–R8, the HB eyelets, and their postsynaptic partners, the effect on sleep pattern was stronger compared to that in norpA mutants. More severe changes in PER expression were observed in l-LNv s than s-LNv s after impairing input from the photoreceptor cells, which is consistent with previous reports (Helfrich-Förster et al., 2001).

Lack of photic information from R1–R6 increased sleep duration at night. However, blocking synaptic transmission from L2 interneurons caused more complex effects on both day- and night-time sleep. L2 interneurons are important for front-to-back motion detection at intermediate pattern contrast (Rister et al., 2007). They are postsynaptic to R1–R6 photoreceptors and to L4 interneurons and presynaptic in feedback synapses, which are formed back to them (both to R1–6 and L4) and use acetylcholine and glutamate as neurotransmitters (Meinertzhagen and O’Neil, 1991; Kołodziejczyk et al., 2008; Raghu and Borst, 2011; Takemura et al., 2011, 2015; Hu et al., 2015). These feedback synapses play a role in preventing photoreceptors from saturation and improving signal quality (Zheng et al., 2006), however, they seem to be extremely important for the functioning of R1–R6 photoreceptors. R1–6 transmit signal to different types of cells, which belongs to pathways specialized to detect contrast increments (ON pathway, L1) or decrements (OFF pathway, L2, L3; Jessch et al., 2010). In effect, blocking R1–6 signaling affects both pathways, however, L2 cells can still receive signal from L4, and send information to deeper parts of the brain. On the other hand, lack of neurotransmission from L2 cells can disrupt not only downstream OFF pathway, but also proper R1–6 functioning, giving the effect on both, day- and night-time sleep. Moreover, according to our transTANGO data, L2 can also directly contact the l-LNv s. It is possible, however, that these synapses are formed only in a specific time of the day, or contacts are very weak and conventional GRASP is not strong enough to show positive results (Muraro and Ceriani,
FIGURE 10 | Glass-expressing cells contact both l-LN_v and s-LN_v. Immunostaining of GMR>transTANGO adult male brain. transTANGO marks presynaptic glass-expressing cells with GFP and the postsynaptic partners of those cells with RFP. Anti-PDF immunostaining confirms that both small and large LN_v are postsynaptic to GMR-Gal4 recruited cells. (A) Hemibrain. Asterisks were placed where PDF and RFP labeling co-localize: in posterior optic tract (POT) coming from l-LN_v, s-LN_v, and their terminals in the dorsal brain. (B) l-LN_v somata (labeled with PDF, blue) have postsynaptic mark (RFP, magenta). HB eyelets projections can be seen nearby (labeled with GFP, green). (C,D) s-LN_v somata and terminals have postsynaptic mark (RFP, magenta). Every brain analyzed (n = 9) showed similar staining. The images were acquired from different brains.

2015). This aspect needs to be addressed by more detailed experiments in the future.

Surprisingly, the lack of synaptic transmission from Rh3-expressing cells affected the morning peak of activity in a very specific way, opposite to that observed after blocking signaling from Rh5. Rhodopsin 3 is expressed in R7 in “pale” ommatidia and in both, R7 and R8, in the DRA type. Since the block of transmission from “pale” ommatidia, with R8 expressing Rh5, decreased the morning peak of activity, it is possible that Rh3, which absorbs UV, inhibits morning activity, while blue light, which is absorbed by Rh5, enhances the morning activity. This can also be an effect of the disruption in polarized light detection, which is received by DRA only.

Siesta or day-time sleep seems to be mostly regulated by R8 “pale” type of photoreceptors. The results obtained in this work may support the idea that blue light inhibits sleep during the day. The exposure to blue light seems to affect lifespan and induce neurodegeneration (Nash et al., 2019), and short sleep during the day allow avoiding UV and blue light during the day, thus protecting flies against harmful light exposure.

Rh6-expressing photoreceptors affect sleep in a specific way, since nap was longer after clock disruption but sleep during the night and day was increased after blocking transmission. Rh6, however, is expressed not only in the retina photoreceptors, but also in the HB eyelets. Our data suggest that R8 cells are involved in night-time sleep and HB eyelets in day-time sleep regulation. Acetylcholine expression silenced in Rh6-expressing cells increased sleep time only during the day, while tetanus toxin expression affected both day and night sleep time. As it is known that among Rh6-expressing cells only HB eyelets use...
FIGURE 11 | L2 interneurons form direct synaptic contacts with large but not small LNv. Immunostaining of L2-transTANGO adult male brain. (A) Hemibrain. Asterisks mark places where PDF and RFP labeling co-localize, which means l-LNv cell bodies and POT. (B) l-LNv somata have postsynaptic labeling, meaning that L2 interneurons form direct contacts with l-LNv. This was seen in 86% of brains (n = 7). A L2 projection can be seen in cyan, rounding a somata. (C,D) Neither somata nor projections of s-LNv have postsynaptic labeling. None of the brains analyzed showed RFP signal in these cells. The images were acquired from different brains.

acetylcholine as neurotransmitter we can conclude that observed effect is specific for these extracellular photoreceptors. It has been reported that the HB eyelets form direct synaptic contacts with pacemaker cells located in the aMe (Muraro and Ceriani, 2015; Li et al., 2018; Schlichting et al., 2019). Our GRASP results suggest that the s-LNv are main target cells, however, it has been shown that also L1-LNv, ITP-expressing LNv, DN1a and DN3a cells receive signals from the HB eyelets through terminals located in the aMe (Li et al., 2018), which was confirmed by transTANGO data. These contacts are the strongest at the beginning of the day, indicating that signaling between the HB eyelets and LNv takes place in the morning. Down-regulation of histamine receptor expression in the pacemaker PDF-immunoreactive cells (Pdf>HisCl) results in lack of the morning anticipation, which is typical of clock mutants. In turn disruption of histamine signaling from photoreceptors to pacemaker cells affects sleep by increasing its length during the day and night (Oh et al., 2013). This means that histaminergic neurotransmission from photoreceptors is involved in the regulation of LNv activity. It has been shown that the HB eyelets express histamine and acetylcholine (Pollack and Hofbauer, 1991; Yasuyama and Meinertzhagen, 1999; Yasuyama and Salvaterra, 1999) and LNv have receptors for both neurotransmitters (McCarthy et al., 2011; Lelito and Shafer, 2012). According to our NrG GRASP data the HB eyelets communicate with the s-LNv directly at the beginning of the day. Schlichting et al. (2016) suggest that HB eyelets may also contact the L1-LNv, as histamine bath had inhibitory effect on L1-LNv (Schlichting et al., 2016), nevertheless, there are also other histamine sources in the aMe (Hamasaka and Nassel, 2006). Light signal received by HB eyelets in the morning is transmitted via acetylcholine and excite s-LNv via nicotinic receptors (Wegener et al., 2004; McCarthy et al., 2011; Schlichting et al., 2016), causing increased cAMP level (Lelito and Shafer, 2012) and wake-promoting effect. This supports our results, which showed that down-regulation of acetylcholine expression in HB eyelets increases sleep time during the day.
Taking together, R1-6, R7 (Rh3-expressing photoreceptor), and R8 “yellow” (Rh6-expressing cells) seem to be important for night-time sleep regulation, and HB eyelets for day-time sleep, while the strongest effect, both during the day and night, has neurotransmission from R8 “pale” (Rh5-expressing cells). We observed a similar effect on night-time sleep when synaptic transmission was blocked from all types of photoreceptors, which means that during the night even weak light inputs received by retinal cells are important to keep flies awake. As a result of the decreased neurotransmission flies are less active and spend more time sleeping during the night. However, it was previously shown that the retinal photoreceptors play a role of peripheral oscillators and regulate daily changes in the visual system, we found that those oscillators also affect behavior. Disruption of the clock in a single type of photoreceptors decreased total activity, and in most cases, it affected sleep time and both morning and evening peaks of activity. These effects were similar to those observed after blocking transmission from specific photoreceptor types. This suggests that synaptic transmission is regulated by the photoreceptor clocks and light. In addition, we showed that HB eyelets, but also L2 interneurons can directly communicate with LNv’s cells, which provides new light pathway to the clock neurons.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

AUTHOR CONTRIBUTIONS

MD designed the study and wrote the manuscript with input from all co-authors. MD and JI performed the experiments and analyzed the data. All authors contributed to the article and approved the submitted version.

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REFERENCES

Chatterjee, A., and Hardin, P. E. (2010). Time to taste: circadian clock function in the Drosophila gustatory system. Fly 4, 283–287. doi: 10.4161/fly.4.4.13010
Chatterjee, A., Tanaeue, S., Heurt, J. H., and Hardin, P. E. (2010). Regulation of gustatory physiology and appetitive behavior by the Drosophila circadian clock. Curr. Biol. 20, 300–309. doi: 10.1016/j.cub.2009.12.055
Chow, E. S., Long, D. M., and Giebultowicz, J. M. (2018). Circadian rhythm in mRNA expression of the glutathione synthesis gene Gcl is controlled by peripheral glial clocks in Drosophila melanogaster. Physiol. Entomol. 41, 369–377. doi: 10.1111/phen.12164
Chung, R. Y., Kliman, V. L., Keith, J. R., Pitman, J. L., and Allada, R. (2009). The GABAA receptor RDL acts in peptidergic PDF neurons to promote sleep in Drosophila. Curr. Biol. 19, 386–390. doi: 10.1016/j.cub.2009.01.040
Damulewicz, M., Mazzotta, G. M., Sartori, E., Rosato, E., Costa, R., and Pyza, E. M. (2017). Cryptochrome is a regulator of synaptic plasticity in the visual system of Drosophila melanogaster. Front. Mol. Neurosci. 10:165. doi: 10.3389/fnmol.2017.00165

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fphys.2020.00993/full#supplementary-material

FIGURE S1 Effects of blocking neurotransmission on activity pattern, performed for (A) R1–R6 photoreceptors (Rh1), (B) Rh3-expressing cells, (C) Rh5-expressing cells, (D) Rh6-expressing cells, (E) L2 cells. Red arrows indicate statistically significant changes in morning or evening peak level. Detailed statistics is presented in Table 2.

FIGURE S2 Effects of clock disruption on activity pattern, performed for: (A) R1–R6 photoreceptors (Rh1), (B) Rh3-expressing cells, (C) Rh5-expressing cells, (D) Rh6-expressing cells. Red arrows indicate statistically significant changes in morning or evening peak level. Detailed statistics is presented in Table 2.

FIGURE S3 Effects of neurotransmission block (Tox) or disruption of the peripheral clock on total activity level, performed for (A) glass-expressing cells (GMR), (B) R1–R6 photoreceptors (Rh1), (C) Rh5-expressing cells, (D) Rh6-expressing cells, (E) Rh-expressing cells, (F) L2 interneurons. Heterozygous parental strains were used as control. Statistically significant differences marked with asterisks: *p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001. Detailed statistics is presented in Supplementary Table S4.

TABLE S1 Statistical analysis of the period of locomotor activity rhythm and percent of rhythmic flies. Each experimental strain was compared with control strains (Gal4 and UAS) using one-way ANOVA and Tukey’s test. Degrees of freedom [F(DFn, DFd)] are listed for every group.

TABLE S2 Statistical analysis of morning anticipation index (MAI) and evening anticipation index (EAI). Each experimental strain was compared with control strains (Gal4/+ and UAS/+) using one-way ANOVA and Tukey’s test (for period) or non-parametric Kruskal-Wallis (for % rhythmic). Genotypes with p < 0.05 with both controls are marked as statistically significant changes with bold. Degrees of freedom [F(DFn, DFd)] are listed for every group.

TABLE S3 Statistical analysis of day-time and night-time sleep. Each experimental strain was compared with control strains (Gal4/+ and UAS/+) using one-way ANOVA and Tukey’s test. Genotypes with p < 0.05 with both controls are marked as statistically significant changes with bold. Degrees of freedom [F(DFn, DFd)] are listed for every group.

TABLE S4 Statistical analysis of PER expression in the clock neurons s-LNv’s and l-LNv’s measured as fluorescence intensity at different time points.

TABLE S5 Statistical analysis of PER expression in the clock neurons s-LNv’s and l-LNv’s measured as fluorescence intensity at different time points.
Grima, B., Chelot, E., Xia, R., and Rouyer, F. (2004). Morning and evening peaks.

Górska-Andrzejak, J., Makuch, R., Stefan, J., Görlich, A., Semik, D., and Pyza, E. (2018). A circadian output

Hamasaka, Y., and Nassel, D. R. (2006). Mapping of serotonin, dopamine, and

Hofbauer, A., and Buchner, E. (1989). Does

Hardie, R. C. (1987). Is histamine a neurotransmitter in insect photoreceptors?

Helfrich-Förster, C., Winter, C., Hofbauer, A., Hall, J. C., and Stanewsky, R. (1997). Rhythmic

Hege, D. M., Stanewsky, R., Hall, J. C., and Giebultowicz, J. M. (1997). Rhythmic

Helfrich-Förster, C., and Winter, C. (2005). GLP-1 receptors mediate circadian modulation of the small ventrolateral pacemaker neurons of Drosophila clock mutant ClkJrk.

Joesch, M., Schnell, B., Raghu, S. V., Reiff, D. F., and Borst, A. (2010). ON and

Krishnan, B., Dryer, S. E., and Hardin, P. E. (1999). Circadian rhythms in olfactory

Lamaze, A., and Stanewsky, R. (2020). DNP1p or the "fluffy" cebers of clock

Leifsson, K. R., and Shafer, O. T. (2012). Reciprocal cholinergic and GABAergic modulation of the small ventrolateral pacemaker neurons of Drosophila's circadian clock neuron network: J. Neurophysiol. 107, 2096–2108. doi: 10.1152/jn.00914.2011

Li, M. T., Cao, L. H., Xiao, N., Tang, M., Deng, B., Yang, T., et al. (2016). Hub-organized parallel circuits of central circadian pacemaker neurons for circadian phototransduction in Drosophila. Nat. Commun. 7:4427. doi: 10.1038/ncomms5604-5

Liu, W. Z., Li, S. L., Zheng, H. Y., Zhang, S. P., and Xu, L. (2012). A broad expression profile of the GMR-GAL4 transgene in Drosophila, Genet. Mol. Res. 11, 997–2002. doi: 10.4238/2012.August.6.1

Mazzotta, G., Rossi, A., Leonardi, E., Mason, M., Bertocchi, C., Caccin, L., et al. (2013). Fly cryptochrome and the visual system. Proc. Natl. Acad. Sci. U.A. 110, 6163–6168. doi: 10.1073/pnas.1221721110

McCarthy, E. V., Yu, Y., deCarvalho, T., Brandt, C., Cao, G., and Nitschke, M. N. (2011). Synchronized bilateral synaptic inputs to Drosophila melanogaster neuropeptidergic rest/arousal neurons. J. Neurosci. 31, 8181–8193. doi: 10.1523/JNEUROSCI.17-10.2011

Meinertzhagen, I. A., and O’Neill, S. D. (1981). Synthetic activity of columnar elements in the lamina of the wild type in Drosophila melanogaster. J. Comp. Neurol. 205, 232–263. doi: 10.1002/cne.902050202

Meinertzhagen, J. A., and Sorra, K. E. (2004). Synthetic organization in the fly’s optic lamina: few cells, many synapses and divergent microcircuits. Proc. Brain Res. 151, 53–69. doi: 10.1016/S0166-2236(03)00107-5

Melnattur, K. V., Pursley, R., Lin, Y. T., Ying, C. Y., Smith, P. D., Pohida, T., et al. (2014). Multiple redundant modulla projection neurons mediate color vision in Drosophila. J. Neurogenet. 28, 374–388. doi: 10.1080/01677063.2014.981592

Munro, N., I., and Cerami, M. F. (2015). Acetylcholine from visual circuit modulates the activity of arousal neurons in Drosophila. J. Neurosci. 35, 16315–16327. doi: 10.1523/JNEUROSCI.0517-15.2015

Nesh, T. R., Chow, E. S., Law, A. D., Fu, S. D., Fuszara, E., Bilska, A., et al. (2019). Drosophila's circadian clock: development, elements in the lamina of the wild type in Drosophila melanogaster. J. Comp. Neurol. 205, 232–263. doi: 10.1002/cne.902050202

Nishio, M., Wada, A. R., Elliott, C. J. H., and Chand, S. (2017). Circadian rhythms in visual responsiveness in the behaviorally arrhythmic Drosophila clock mutant Clock. J. Biol. Rhyth. 32, 583–592. doi: 10.1177/0190821617735397

Ogut, M., Hardie, R. C., and Stanewsky, R. (2018). Non-canonical phototransduction mediates synchronization of the Drosophila melanogaster circadian clock and retinal light responses. Curr. Biol. 28, 1725–1735. doi: 10.1016/j.cub.2018.04.014

Ogut, M., Hardie, R. C., and Stanewsky, R. (2020). Light sampling via throttled visual phototransduction robustly synchronizes the Drosophila circadian clock. J. Biol. Rhyth. 30, 2551–2563. doi: 10.1086/699550-47588-6

O’Toole, J., Eisele, W., Martin, R. L., Hirth, J., Paik, W. L., and Applebury, M. L. (1965). The Drosophila ninae gene encodes an opsin. Cell 40, 839–850. doi: 10.1016/0006-8495(65)90003-X

Pantsak, A., Segaran, A., Liu, C. H., Nikolaeff, A., Rister, J., Thurn, A. S., et al. (2008). Distinct roles for two histamine receptors (hHa and hHb) in the Drosophila photoreceptor synapse. J. Neurosci. 28, 7250–7259. doi: 10.1523/JNEUROSCI.4070-08.2008

Parisky, R. M., Agasto, J., Pulver, S. R., Shang, Y., Kuklin, E., Hodge, J. J. L., et al. (2018). Drosophila’s circadian clock: development, elements in the lamina of the wild type in Drosophila melanogaster. J. Comp. Neurol. 205, 232–263. doi: 10.1002/cne.902050202

Perzynski, A. M., Randall, L. L., Shortridge, R. D., Burg, M. G., and Pak, W. L. (1996). Molecular, biochemical, and electrophysiological characterization of Drosophila ninae mutants. J. Biol. Chem. 271, 4937–4945. doi: 10.1074/jbc.271.9.4937-4945

Pollack, L., and Hofbauer, A. (1991). Histamine-like immunoactivity in the visual system and brain of Drosophila melanogaster. Cell Tissue Res. 266, 391–398. doi: 10.1007/BF00318495
Potdar, S., and Sheeba, V. (2018). Wakefulness is promoted during daytime by PDRF signaling to dopaminergic neurons in Drosophila melanogaster. *eLife* 5:e36964. doi: 10.7554/eLife.36964

Ray, M., and Lakhotia, S. C. (2015). The commonly used eye-specific sev-GAL4 and RAG8-GAL4 drivers in Drosophila melanogaster are expressed in tissues other than eyes as well. *J. Genet.* 94, 407–416. doi: 10.1016/j.jgenet.2014.05-0558

Rister, J., Pauls, D., Schnell, B., Ting, C. Y., Lee, C. H., Smuckler, E., et al. (2007). Dissection of the peripheral motion pathway in the visual system of Drosophila melanogaster. *Neuron* 55, 155–170. doi: 10.1016/j.neuron.2007.09.014

Saint-Charles, A., Michard-Vanhee, C., Alejevski, F., Chelot, E., Boivin, A., Szular, J., Sehadova, H., Gentile, C., Szabo, G., Chou, W.-H. H., Britt, S. G., et al. (2018). candidate glutamatergic neurons in the visual system of Drosophila melanogaster. *Neuron* 56, 155–170. doi: 10.1016/j.neuron.2017.10.033

Takamura, S. Y., Xu, C. S., Lu, Z., Rivlin, P. K., Parag, T., Olbris, D. J., et al. (2015). Synaptic circuits and their variations within different columns in the visual system of Drosophila. *Proc. Natl. Acad. Sci. U.S.A.* 112, 1371–1376. doi: 10.1073/pnas.1509821112

Talsky, M., Richman, E. B., Snell, N. J., Hartmann, G. G., Fisher, J. D., Sorkin, A., et al. (2017). Transsynaptic mapping of second-order taste neurons in flies by trans-tango. *Neuron* 96, 783–795. doi: 10.1016/j.neuron.2017.10.011

Vossius, L. B., and Young, M. W. (1995). Circadian rhythms in Drosophila can be driven by period expression in a restricted group of central brain cells. *Neuron* 15, 345–360. doi: 10.1016/0896-6273(95)90039-X

Weber, P., Kola-Iwrowe, E., and Pyza, E. (2009). Circadian control of dendrite morphology in the visual system of Drosophila melanogaster. *PLoS One* 4:e2420. doi: 10.1371/journal.pone.0004290

Wegener, C., Haranaska, Y., and Nüssel, D. R. (2004). Acetylcholine increases intracellular Ca2+ Via nicotinic receptors in cultured Drosophila clock neurons. *J. Neurophysiol.* 91, 912–923. doi: 10.1152/jn.00726.2003

Wernet, M. F., Muzzoni, E. O., Gallì, A., Duncan, D. M., Duncan, L. and Desplan, C. (2005). Stochastic spike emission creates the retinal mosaic for colour vision. *Nature* 430, 174–180. doi: 10.1038/nature02815

Yamaguchi, S., Wolf, R., Desplan, C., and Helsenberg, M. (2008). Motion vision in independent of color in Drosophila. *Proc. Natl. Acad. Sci. U.S.A.* 105, 4910–4915. doi: 10.1073/pnas.0711840105

Yasuyama, K., and Meinertzhagen, I. A. (1999). Extraretinal photoreceptors at the compound eye's posterior margin in Drosophila melanogaster. *J. Comp. Neurol.* 412, 193–202. doi: 10.1002/(sici)1096-9861(19990920)412:2<193::aid-jcn2>3.0.co;2-0

Yasuyama, K., and Salvaterra, P. M. (1999). Localization of choline acetyltransferase-expressing neurons in Drosophila nervous system. *Microsc. Res. Techniq.* 45, 65–79. doi: 10.1002/(sici)1097-0029(19990815)45:2<65::aid-jemt2>3.0.co;2-0

Zerr, D. M., Hall, J. C., Rosbash, M., and Siwicki, K. K. (1990). Circadian fluctuations of period protein immunoreactivity in the CNS and the visual system of Drosophila. *J. Neurosci.* 10, 2749–2762. doi: 10.1523/jneurosci.10-08-2749.1990

Zheng, L., de Polavieja, G. G., Wolfram, V., Asyali, M. H., Hardie, R. C., and Dammin, R. W. (2006). Feedback network controls photoreceptor output at the layer of first visual synapses in Drosophila. *J. Gen. Physiol.* 127, 495–510. doi: 10.1085/jgp.2005049470

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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