

PDF Cells Are a GABA-Responsive Wake-Promoting Component of the *Drosophila* Sleep Circuit

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SUMMARY

Daily sleep cycles in humans are driven by a complex circuit within which GABAergic sleep-promoting neurons oppose arousal. *Drosophila* sleep has recently been shown to be controlled by GABA, which acts on unknown cells expressing the *Rdl* GABA_A receptor. We identify here the relevant *Rdl*-containing cells as PDF-expressing small and large ventral lateral neurons (LNvs) of the circadian clock. LNv activity regulates total sleep as well as the rate of sleep onset; both large and small LNvs are part of the sleep circuit. Flies mutant for *pdf* or its receptor are hypersomnolent, and PDF acts on the LNvs themselves to control sleep. These features of the *Drosophila* sleep circuit, GABAergic control of onset and maintenance as well as peptidergic control of arousal, support the idea that features of sleep-circuit architecture as well as the mechanisms governing the behavioral transitions between sleep and wake are conserved between mammals and insects.

INTRODUCTION

The regulation of sleep is of vast clinical importance. Insomnia and circadian disorders are costly in both economic and human health terms. Sleep is believed to be controlled by both circadian and homeostatic systems, which ensure that sleep needs are met. The heart of the mammalian sleep circuit is a switch consisting of reciprocally connected sleep and arousal centers (Fuller et al., 2006; Sakurai, 2007). The ventrolateral preoptic area of the hypothalamus contains inhibitory GABAergic sleep-promoting neurons, whereas arousal centers are more distributed and consist of both aminergic and cholinergic neurons; these cells additionally feed back to inhibit the hypothalamic sleep center. Hypothalamic neurons that release the peptide orexin/hypocretin also modulate the switch and stabilize the waking state. Loss of this peptide results in narcolepsy, an inability to maintain the

waking state. In summary, the organization of the human circuit is complex and not completely understood.

Drosophila has become a well-accepted behavioral model for sleep research, and we have recently shown that GABAergic transmission influences total sleep as well as sleep latency. In humans, both the initiation and maintenance of sleep are also controlled by GABAergic inputs to arousal centers, which explains why drugs that enhance transmission via GABA_A receptors are among the most widely used sleep-promoting agents (Roth, 2007). The conserved role of GABA even extends to this pharmacological level: the GABA_A receptor subunit encoded by the *Rdl* gene (defined by resistance to the insecticide and GABA_A antagonist dieldrin) has a role in the onset of fly sleep (Agosto et al., 2008). These findings indicate that part of the core circuitry controlling sleep in flies will consist of GABA_A-regulated, wake-promoting cells like in mammals.

In this study, we identify a population of circadian clock cells, the ventrolateral neurons or LNvs, that meet these criteria. Manipulation of *Rdl* levels within the LNvs indicates that they are a major target of sleep-promoting GABAergic neurons. Up- and downregulation of PDF-positive LNv activity demonstrates that they control both total sleep and sleep onset. Indeed, acute activation of the large LNvs alone is sufficient to block sleep in the early evening, indicating that this subset of the LNvs responds to arousal signals and is a target of homeostatic regulation. Moreover, we show that the peptide PDF and its receptor are required to mediate the wake-promoting effects of these cells and that PDF modulation of LNvs themselves (presumably small LNvs) can regulate sleep. These results indicate that in flies, as in mammals, the sleep circuit is intimately linked to the circadian clock and that the strategies used to govern sleep in the brain are evolutionarily ancient.

RESULTS

To identify the *Rdl*-expressing cells controlling sleep, anti-*Rdl* antisera were used to examine the pattern of *Rdl* protein expression in the adult brain. A strongly immunoreactive group of lateral brain neurons was identified by double staining as the PDF peptide-containing lateral clock neurons (LNvs). Figures 1A

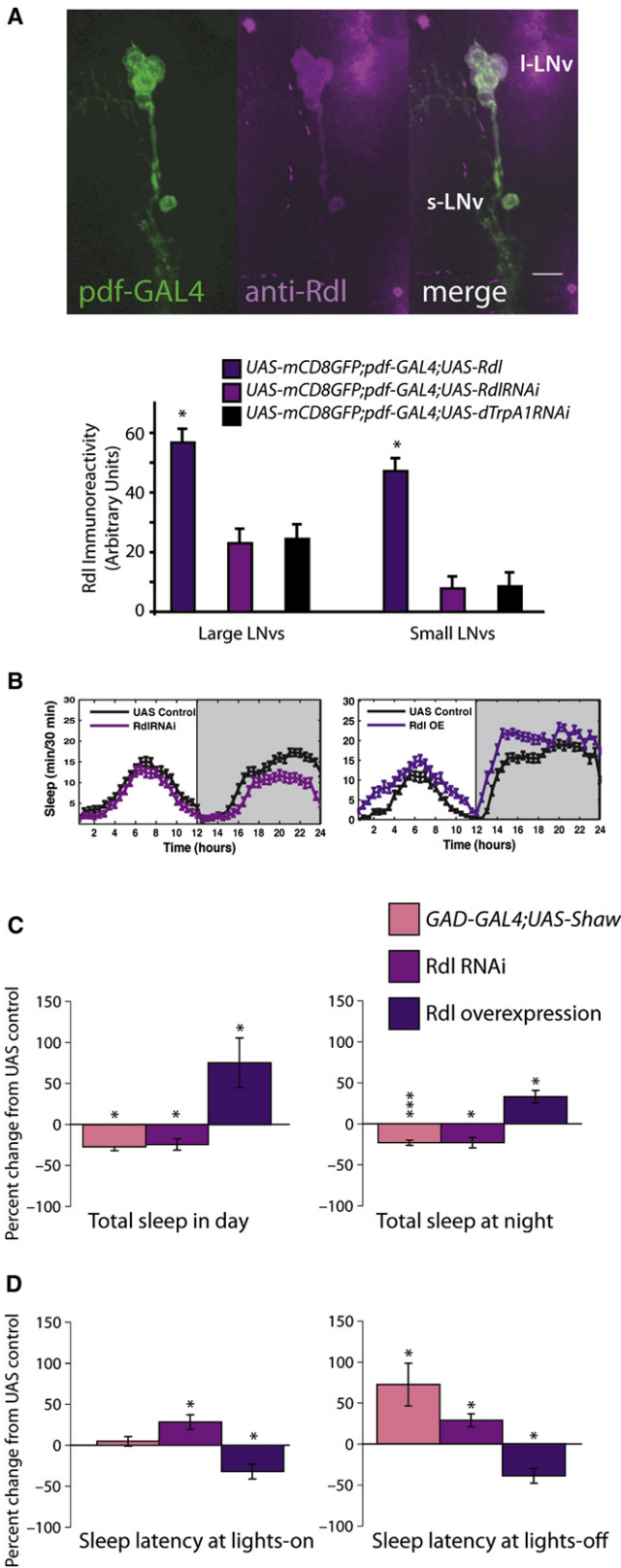


Figure 1. LNvs Express the *Rdl* GABA_A Receptor and Mediate GABAergic Effects on Sleep

(A) Top shows images of Rdl expression in wild-type LNvs. Adult brains from *pdf-GAL4;UAS-mCD8-GFP* animals were stained with anti-Rdl (1:100) and visualized with confocal microscopy. Rdl is shown in magenta, GFP in green, overlap in white. Scale bar, 10 μ m. Bottom shows quantification of somatic Rdl levels in LNvs expressing excess Rdl, RdlRNAi, or control dTrpA1RNAi.

(B) Standard sleep plots of control and experimental flies in 12 hr:12 hr light:dark (LD). Left panel shows the effects of reducing Rdl levels in LNvs: *pdf-GAL4;UAS-RdlRNAi*, right panel shows the effects of overexpressing Rdl in LNvs: *pdf-GAL4;UAS-Rdl*.

(C) GABA regulates total sleep. Twelve hour sleep from the light (left) or dark (right) period in LD was assessed for animals with decreased overall GABAergic transmission (*GAD-GAL4;UAS-Shaw*; n = 62), decreased LNV Rdl levels (*pdf-GAL4;UAS-RdlRNAi*; n = 93) or increased LNV Rdl levels (*pdf-GAL4;UAS-Rdl*; n = 21). Data are expressed as the percent change from the genetic control.

(D) GABA regulates sleep onset. The latency to first sleep bout during the light (left) or dark (right) period in LD was assessed for the same genotypes. Data are expressed as the percent change from the genetic control.

*p < 0.05, **p < 0.005, and ***p < 0.0005 for comparisons of experimental and control conditions using ANOVA with Tukey post hoc test. Data are presented as mean \pm SEM.

(top panel) and S1 (available online) show staining of whole adult brains from *pdf-GAL4;UAS-mCD8-GFP* animals, demonstrating that the *pdf-GAL4*-positive LNvs, i.e., both small (s-LNvs) and large LNvs (l-LNvs), express the *Rdl* GABA_A receptor. Strong Rdl expression was also detected in a nearby *pdf-GAL4*-negative cell, whose position and morphology suggest that it may represent the fifth small LNV (Figure S1). Preliminary experiments examining Rdl levels at ZT3 and ZT15 suggest that protein levels do not undergo circadian oscillations (data not shown).

As the circadian system and the suprachiasmatic nucleus (SCN) regulate mammalian sleep, the LNvs were attractive candidates for affecting *Drosophila* sleep. To determine whether *Rdl* activity within LNvs was relevant, we both overexpressed and downregulated the Rdl GABA_A receptor exclusively in LNvs (Figure 1A, bottom panel). Expression of RNAi for *Rdl* using *pdf-GAL4* slightly decreased somatic Rdl levels in both l- and s-LNvs, although the decrease was not statistically significant when compared to Rdl in LNvs expressing a control RNAi against dTrpA1, a channel protein expressed in a small number of non-clock adult neurons (Hamada et al., 2008). Expression of this control RNAi did not significantly affect sleep (Figure S2). Overexpression of Rdl cDNA significantly increased somatic Rdl immunoreactivity in both l- and s-LNvs. Rdl in LNV processes was also dramatically increased, going from undetectable in controls to bright in overexpressers (data not shown).

To assess the behavioral effects of altering Rdl levels in LNvs, we measured daytime and nighttime sleep in 12 hr light:12 hr dark cycles (LD). Remarkably, the nature and severity of the effects caused by LNV-specific knockdown of Rdl expression were virtually identical to the effects of reducing the excitability of GABA-producing neurons throughout the nervous system via overexpression of the hyperpolarizing Shaw potassium channel (Figures 1C and 1D; Agosto et al., 2008), a small but significant decrease in sleep. Only sleep latency after lights-off was more profoundly affected by the downregulation of GABAergic

tone with overexpression of *Shaw* (Figure 1D), perhaps reflecting the small change in Rdl levels or possibly a role for other GABA receptors; we note that cultured LNVs have been shown to have GABA_B receptors (Hamasaka et al., 2005). Not only was Rdl activity within the LNVs required for sleep, the level of Rdl activity appeared important in determining the quantity of sleep, as overexpression of Rdl within the LNVs caused flies to fall asleep faster and to remain asleep longer in both the day and at night (Figure 1B, right). These results indicate that the circadian LNVs are a major target for GABAergic control of sleep and suggest that the LNVs may act as wake-promoting cells.

To more directly test this hypothesis, we used transgenes that modulate the excitability of LNVs to increase or decrease neuronal activity. A similar approach has been previously used to demonstrate a role for LNV neuronal activity in coordinating the circadian clock circuit under constant (DD) conditions (Nitabach et al., 2002, 2006). Expression of the EKO potassium channel (White et al., 2001) is expected to hyperpolarize cells, reducing their ability to be stimulated by endogenous inputs. To chronically increase activity, we developed two new molecular tools. The response to inputs was increased by either expressing an RNAi construct (Figure 2A) against the ubiquitous leak channel *Shaw* (Hodge et al., 2005) or by expressing a dominant-negative Na⁺/K⁺-ATPase α subunit (Sun et al., 2001). Whole-cell current-clamp recordings from larval motor neurons expressing these transgenes indicate that they both increase resting membrane potential and the firing-rate response (Figures 2B and 2C). These manipulations amplify the effects of endogenous inputs and allowed us to interrogate the normal function of LNVs. This is unlike the widely used bacterial sodium channel NaChBac (Nitabach et al., 2006), which imposes a novel constitutive activity pattern on neurons (Sheeba et al., 2008b). The only other putatively activity-enhancing transgene, truncated Eag (Broughton et al., 2004), has not been characterized electrophysiologically.

To restrict the action of these activity modulators to LNV neurons, we expressed them under the control of *pdf-GAL4* and assayed sleep under standard LD conditions. Suppression of LNV activity increased both daytime and nighttime sleep (Figures 3A and 3B). In contrast, increasing LNV excitability using either the *Shaw* RNAi or the dominant-negative Na⁺/K⁺-ATPase transgene decreased daytime sleep significantly (Figures 3A and 3B). This enhancement of LNV activity also had a suppressing effect on total nighttime sleep, further supporting a role for LNVs in sleep regulation. Importantly, sleep latency was also bidirectionally modulated by alterations in LNV excitability: decreased LNV activity caused flies to fall asleep faster in both the day and night, whereas increasing excitability suppressed initiation of the first sleep bout (Figure 3C). We found no coherent effect of manipulation of LNV excitability on locomotor activity during active periods (Figure S3), demonstrating that the regulation of sleep is independent of basal locomotor activity, as has been previously demonstrated. We also found no effect of these manipulations on the circadian pattern of locomotor activity (Figure S4), indicating that their effects on sleep are not secondary to disruption of the clock.

Control flies of all of the genotypes shown have a long sleep latency during the daytime, maintaining wakefulness for about an hour after lights-on. This daytime sleep latency is strongly

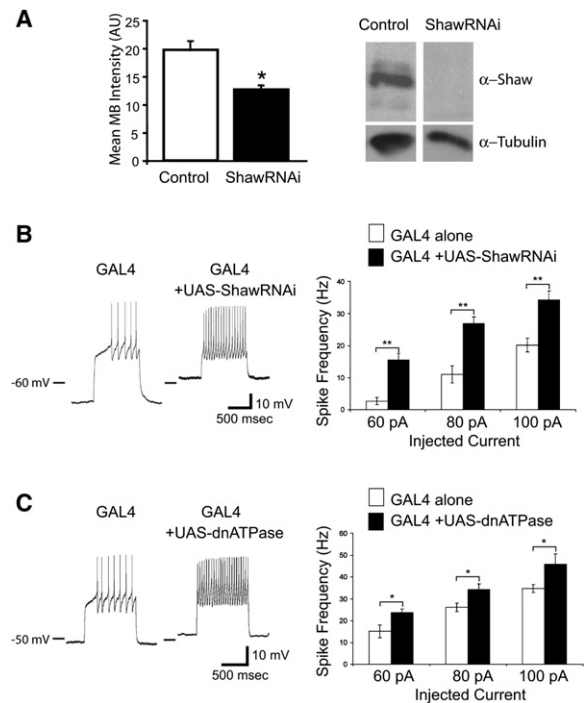


Figure 2. *Shaw* RNAi and Dominant-Negative Na⁺/K⁺-ATPase Increase Neuronal Excitability

(A) *Shaw* RNAi reduces endogenous *Shaw* expression. (Left) Expression of *Shaw* double-stranded RNA in adult central and motor neurons (*GAL4-C380/UAS-ShawRNAi*) causes reduction in *Shaw* levels. Endogenous *Shaw* levels were detected with the anti-C terminus *Shaw* antibody and quantified in the mushroom body calyx region with Leica confocal software. A significant reduction ($p < 0.05$) in intensity (arbitrary units) is seen in central neurons expressing *ShawRNAi*. (Right) Expression of interfering *Shaw* double-stranded RNA (*GAL4-24B/UAS-ShawRNAi*) decreases endogenous *Shaw* compared to controls (*+UAS-ShawRNAi*) when compared by immunoblot. Whole head lysates were separated by SDS-PAGE, transferred to nitrocellulose membranes, and detected with an antibody (1:1000) to the C terminus of *Shaw* that detects full-length *Shaw* protein (Hodge et al., 2005). Anti-tubulin (1:200,000) was used to assess protein loading.

(B) Expression of *Shaw* RNAi in larval motor neurons with *C380-GAL4* increases excitability. (Left) Traces of whole-cell current-clamp recording from MNISN-Is of control (*C380-GAL4* only) and experimental (*C380-GAL4;UAS-ShawRNAi*) animals injected with 60 pA current. (Right) Quantified data for the firing-rate response to various current injections. $n = 7$ for *C380-GAL4* alone, and $n = 10$ for *C380-GAL4;UAS-ShawRNAi*.

(C) Expression of dominant-negative Na⁺/K⁺-ATPase in larval motor neurons with *C380-GAL4* increases excitability. (Left) Traces of whole-cell current-clamp recording from MNISN-Is of control (*C380-GAL4* only) and experimental (*C380-GAL4;UAS-dnATPase*) animals injected with 60 pA current. (Right) Quantified data for the firing rate response to various current injections. $n = 8$ for *C380-GAL4* alone, and $n = 7$ for *C380-GAL4;UAS-dnATPase*. Data are presented as mean \pm SEM.

dependent on light, as flies fall asleep sooner after the start of the subjective day in DD ($p < 0.0001$ for a comparison of latency during the daytime in LD versus during subjective day in DD, Figures 3C and 4E). Suppressing LNV activity significantly blocks the wake-promoting effect of light (Figure 3C, left panel). Interestingly, enhancing LNV activity amplifies the effect of light on sleep latency (Figure 3C, right panel). The data are consistent

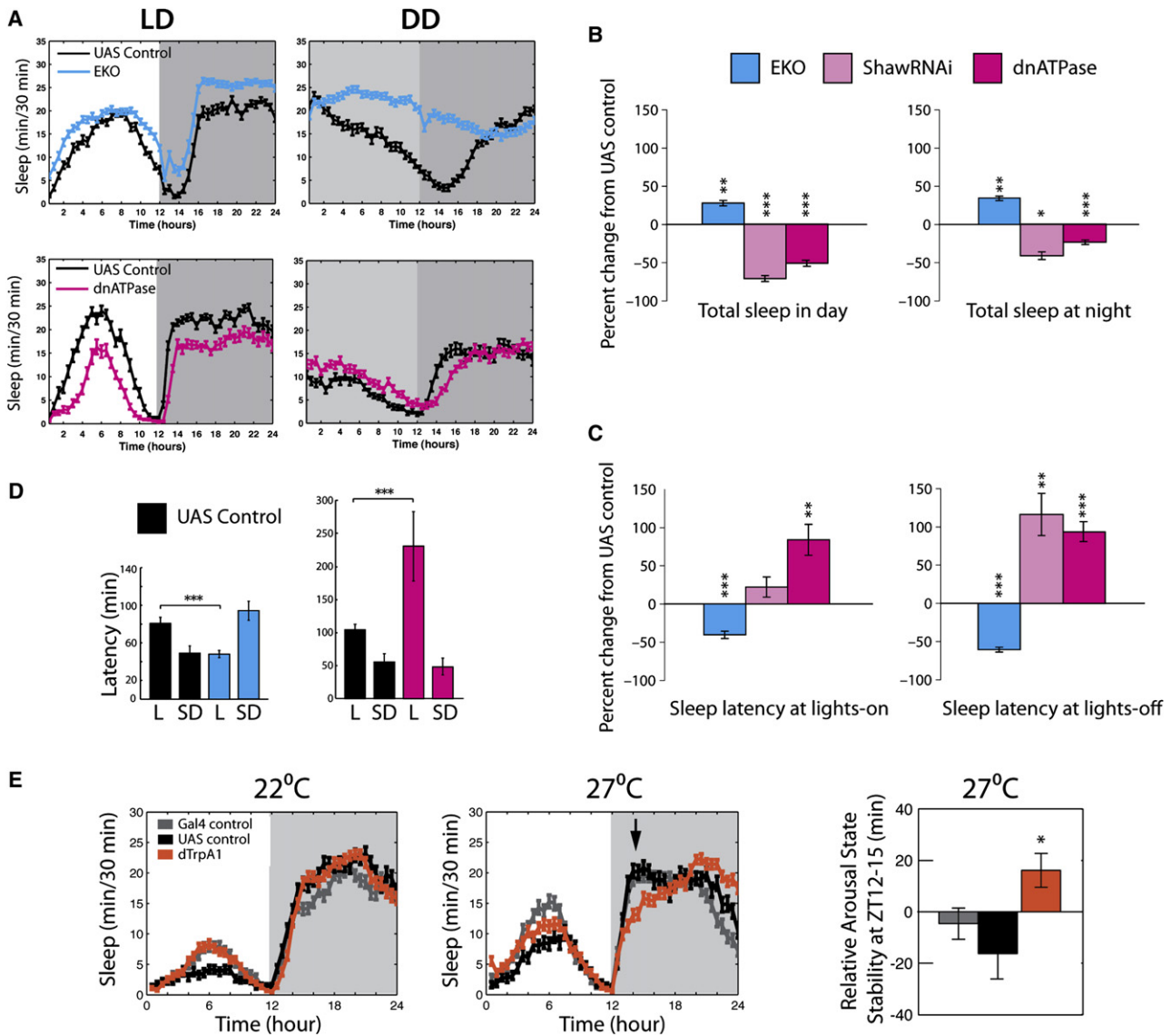


Figure 3. Excitability of LNvs Controls Sleep

(A) Standard sleep plots of control and experimental flies in 12 hr:12 hr light:dark (LD) or constant darkness (DD). Top panel show the effects of reducing neuronal activity levels in LNvs: *pdf-GAL4;UAS-EKO*. Bottom panels show the effects of enhancing normally patterned activity in LNvs: *pdf-GAL4;UAS-dnATPase*.

(B) LNv activity controls total sleep. Twelve hours sleep from the light (left) or dark (right) period in LD was assessed for animals with suppressed responsiveness to inputs (*pdf-GAL4;UAS-EKO*; $n = 55$) or increased responsiveness to inputs (*pdf-GAL4;UAS-ShawRNAi* and *pdf-GAL4;UAS-dnATPase*; $n = 32$ and 80). Data are expressed as the percent change from the genetic control.

(C) LNv activity controls sleep onset. The latency to first sleep bout during the light (left) or dark (right) period in LD was assessed for the same genotypes. Data are expressed as the percent change from the genetic control.

(D) LNvs mediate the wake-promoting effects of light. Latency to first sleep bout during the light period in LD (L) or subjective day in DD (SD) is shown for animals with reduced responsiveness to inputs (left, *pdf-GAL4;UAS-EKO*) or with increased responsiveness to inputs (right, *pdf-GAL4;UAS-dnATPase*).

(E) Acute activation of LNvs disrupts nighttime sleep. *pdf-GAL4;UAS-dTrpA1* and control animals ($n = 32$ for each genotype) were raised at the nonpermissive temperature of 22°C and entrained in LD at that temperature. Data were collected for 3 days and then temperature was increased to 27°C to activate dTrpA1. Left panel shows the 3 days immediately preceding the temperature increase. Middle panel shows 3 days after temperature increase. Left panel shows arousal state stability at 27°C for all genotypes in the early evening (ZT12–15; time marked by arrow in middle panel).

* $p < 0.05$, ** < 0.005 , and *** < 0.0005 for t test comparisons of experimental and control conditions in panels (B) and (C) and for Tukey post hoc test after ANOVA for panels (D) and (E). Data are presented as mean \pm SEM.

with light modulating LNv activity to control sleep onset and are in agreement with reports that LNvs are directly activated by light (Sheeba et al., 2007).

All of the manipulations documented above are chronic: the activity manipulation occurs throughout the lifetime of the cell. To rule out the possibility that developmental effects or circuit

rewiring were responsible for the sleep phenotypes we observed, we expressed the temperature-gated nonspecific cation channel dTrpA1 in LNvs. This channel is activated at temperatures above 25°C in *Drosophila* larval neurons (Hamada et al., 2008). In the adult brain, endogenous dTrpA1 is detectably expressed in only about a dozen cells. These cells are not known circadian cells, mushroom body cells, or other circuits believed to be involved in sleep. In contrast to the chronic manipulations used above that amplify or suppress the effects of native inputs to LNvs, activated dTrpA1 imposes a fast firing pattern on the cell. Animals raised at 22°C, a temperature at which the channel is not open, show normal sleep patterns in LD compared to both GAL4-alone and UAS-alone control animals (Figure 3E). Elevating the temperature to 27°C results in an immediate increase in overall sleep for all genotypes, especially in the day. Specific to flies with dTrpA1 expressed in LNvs is an increase in wakefulness in the early night. This is reflected in a significant increase in arousal state stability between ZT12 and 15, a measure of the relative length of wake and sleep bouts. Daytime sleep was largely unaffected by dTrpA1 expression in LNvs. However, the LNvs are normally activated by light (Sheeba et al., 2008b), potentially masking any additional effects of dTrpA1 on firing and wakefulness, particularly in females, which generally have lower levels of daytime sleep than males (Hendricks et al., 2003).

Previous studies of LNV function have uncovered roles for the LNV-specific circadian neuropeptide PDF and perhaps other transmitters released by LNvs in the regulation of locomotor behavior (Sheeba et al., 2008b). To determine whether PDF is involved in the LNV regulation of sleep, we examined the sleep behavior of *pdf⁰¹* mutant flies, which lack this neuropeptide transmitter. Compared to controls, mutant flies had significantly more daytime sleep (Figures 4A–4C) in LD and even under constant dark conditions (DD). Nighttime sleep was less affected, but this may be due to a ceiling effect. (The genetic controls for the *pdf⁰¹* mutant had a slightly higher basal level of nighttime sleep.) This increase in daytime sleep was due primarily to a decrease in wake duration/consolidation during the day, similar to mammalian narcolepsy. EKO flies also had a similarly decreased mean wake episode duration, whereas *Shaw* RNAi, *Rdl* RNAi, and dominant-negative Na⁺/K⁺-ATPase animals had the opposite effect: their mean wake episode duration increased (data not shown). Loss of PDF also had effects on sleep latency, i.e., how fast the fly fell asleep after a light transition (Figure 4E). The effects of *pdf⁰¹* on latency and on total sleep were similar in magnitude to the changes seen with expression of EKO or *Rdl* in LNvs (Figures 1 and 3).

As noted above, control flies have a longer sleep latency after lights-on in LD than in DD. This light-dependent latency is totally abolished in *pdf⁰¹* flies ($p > 0.05$ for comparison of latency during day versus subjective day; Figure 4E, right panel). This suggests that the comparable effect in animals with decreased LNV excitability (Figure 3D, left) is due to a decrease in PDF release, as opposed to some other LNV transmitter. There is also an increase in latency in subjective night compared with subjective day, which is also eliminated in the *pdf⁰¹* mutant (Figure 4E). The basis of this second alteration is unclear and could even be an indirect effect, i.e., it may mirror features of homeostatic sleep regulation and increased daytime sleep apparent in this mutant background.

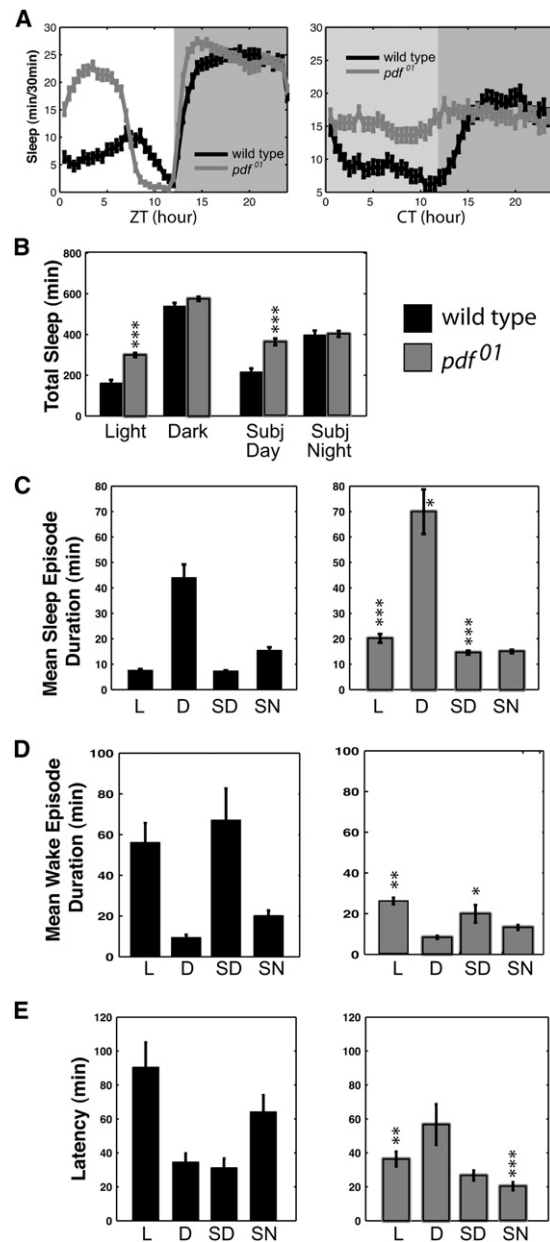


Figure 4. *pdf⁰¹* Mutants Have Increased Total Sleep and Decreased Sleep Latency

(A) Standard sleep plots of control and mutant flies in 12 hr:12 hr light:dark (LD, left) or in constant darkness (DD, right).

(B) Total sleep for controls (black bars) and *pdf⁰¹* mutants (gray bars) for the light period and dark period in LD and subjective day and subjective night in DD.

(C) Mean sleep episode duration, (D) mean wake episode duration, and (E) latency to first sleep bout, for control (left) and *pdf⁰¹* mutants (right). Data are shown for light (L) and dark (D) periods in LD and for subjective day (SD) and subjective night (SN) in DD.

Data are presented as means \pm SEM. * $p < 0.05$, ** $p < 0.005$, and *** $p < 0.0005$ for the comparison to control by ANOVA with Tukey post hoc test. $n = 106$.

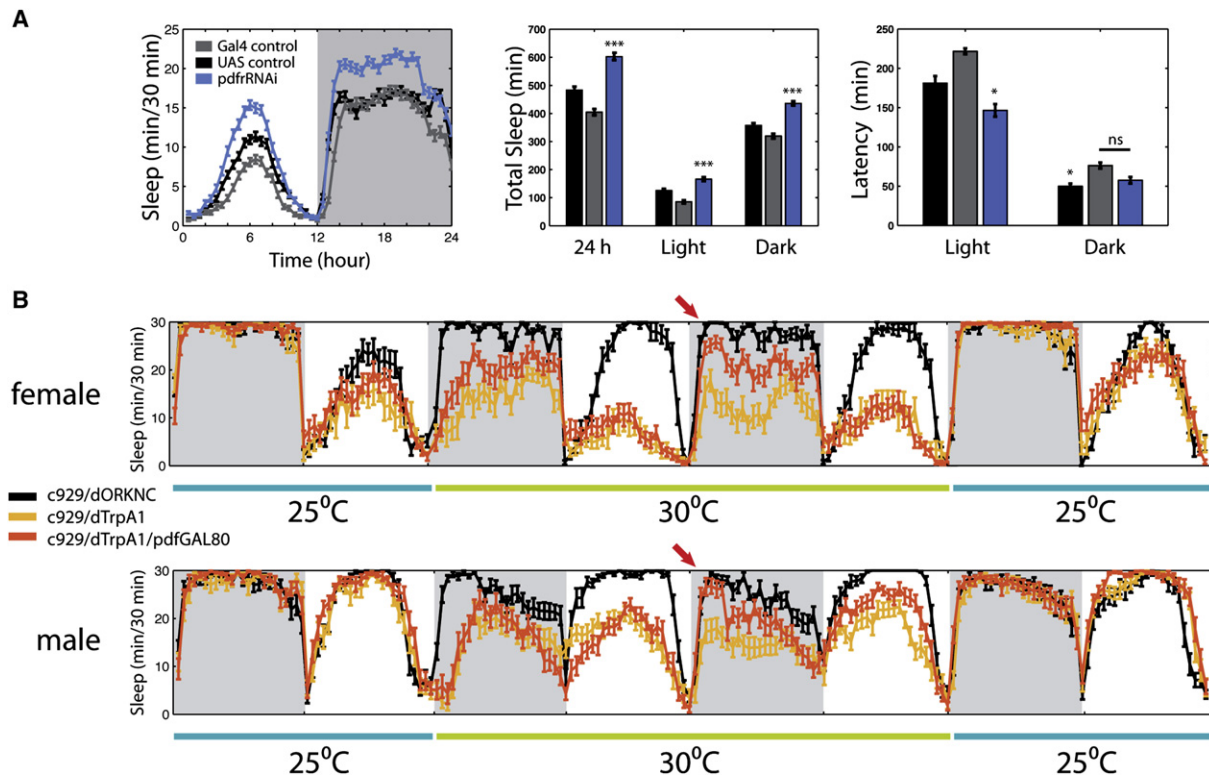


Figure 5. Both Large and Small LNvs Are Involved in Sleep Control

(A) Downregulation of the PDFR with *UAS-pdfRNAi* driven by *pdf-GAL4* in LNvs increases both daytime and nighttime sleep but only significantly affects daytime latency. Standard sleep plots of female flies in 12 hr:12 hr light:dark are shown. * $p < 0.05$, *** $p < 0.0005$, and ns indicates “not significant” for the comparison to other genotypes by ANOVA with Tukey post hoc test. $n = 70, 71$, and 75 for UAS alone (*UAS-pdfRNAi*), GAL4 alone (*pdfGAL4*), and experimental conditions (*pdf-GAL4;UAS-pdfRNAi*), respectively.

(B) Continuous sleep data from flies expressing the temperature-gated cation channel dTrpA1 in peptidergic neurons \pm I-LNvs. Flies were entrained in LD for 5 days at 25°C (last day is shown) and shifted to 30°C for 2 days, then back to 25°C. Females ($n = 16$ for control *c929-GAL4;UAS-dORKNC*), 14 for *c929-GAL4;UAS-dTrpA1*, and 21 for *c929-GAL4;pdf-GAL80;UAS-dTrpA1*) are shown at top, males ($n = 23$ for control, 20 for *c929*, and 19 for *c929+pdf-GAL80*) at bottom. Arrow indicates rescue of early evening sleep by suppression of dTrpA1 expression in I-LNvs by *pdf-GAL80* on day 2 of elevated temperature. Data are presented as mean \pm SEM.

These factors may also contribute to the increase in latency seen during subjective day in EKO flies (Figure 3D).

The *pdf-GAL4* driver expresses in both I-LNv and s-LNv cells. s-LNvs have been postulated to be “morning cells,” which time the onset of morning behavior (Grima et al., 2004), as well as the key pacemaker cells in constant darkness (Helfrich-Forster, 1998; Stoleru et al., 2005). The function of I-LNvs has been obscure, but it has recently been shown that they respond directly to light (Sheeba et al., 2008b) and promote activity during the day, i.e., they act as dawn photoreceptors for arousal (Shang et al., 2008; Sheeba et al., 2008a). To determine the relative roles of s- and I-LNvs in sleep, we asked whether PDF signaling between I- and s-LNvs was important for the wake-promoting role of these cells by downregulating PDFR. Since the majority of I-LNvs do not respond to PDF (Shafer et al., 2008), this manipulation should primarily test the function of s-LNvs. We find that loss of PDFR in these cells increases total sleep in both the daytime and the nighttime (Figure 5A). Interestingly, sleep latency is only decreased compared to both GAL4 and UAS controls during the light period, suggesting that there may be other targets of

PDF relevant to sleep. This is supported by the finding that a P element-generated partial deletion of the *pdfr* gene (Mertens et al., 2005) manifests a decrease in both daytime and nighttime sleep latency as well as increased total sleep (Figure S5).

To examine the specific role of I-LNvs, we first altered the temporal firing pattern of a broad set of peptidergic (PHM⁺) neurons by overexpressing dTrpA1 with the *c929-GAL4* driver (Park et al., 2008). The expression pattern of this driver includes I-LNvs but not s-LNvs. Flies were entrained for 5 days in LD at 25°C, switched to 30°C for 2 days and then back to 25°C to determine if effects were reversible. Figure 5B shows a continuous trace of sleep behavior starting during the last day of entrainment.

As in the *pdf-GAL4* experiment (Figure 3E), temperature elevation increased daytime sleep even for the control genotype. In contrast, however, activation of dTrpA1 in PHM⁺ neurons caused a dramatic decrease in both daytime and nighttime sleep ($p < 0.01$ compared to control for females, $p < 0.001$ compared to controls for males, ANOVA with post hoc t test). This indicates that some of these peptidergic neurons are part of the fly arousal system. Temperature elevation to 27°C also produced a decrease

in sleep in the *c929-GAL4;UAS-dTrpA1* animals (data not shown), but effects at 30°C were more robust.

Because PHM⁺ neurons include the I-LNvs and because *pdf-GAL80* completely suppresses GAL4 transcription activity in all LNvs (Stoleru et al., 2004), we also assayed the sleep phenotype of *c929-GAL4;UAS-dTrpA1;pdf-GAL80* flies (Figure 5A) to dissect out the specific role of I-LNvs. Immediately after temperature elevation, *c929-GAL4* female flies expressing *pdf-GAL80* slept significantly more than dTrpA1-expressing flies without GAL80 ($p < 0.0001$). On day 2 of elevated temperature, however, both male and female flies containing *pdf-GAL80* showed a markedly increased amount of sleep during the early night compared to *c929-GAL4;UAS-dTrpA1* flies (arrows on Figure 5A; $p < 0.0001$ for females, $p < 0.01$ for males). Late-night female sleep was also restored. This effect was harder to discern in males, perhaps because the M-peak of predawn activity causes control flies to sleep less in the late night. Because sleep homeostasis promotes rebound sleep after sleep deprivation, it seems likely that the enhanced sleep in the early night on day 2 reflects this process and that high levels of I-LNv rebound sleep were stimulated by the build up of sleep pressure on the second day. We were only able to visualize the effects of sleep deprivation with the dTrpA1 animals because only the chronic situation is visible with the dnATPase and ShawRNAi tools. In aggregate, the data suggest that persistent I-LNv firing keeps flies awake at night but that the effects are larger at the beginning of the night than at the end of the night. Therefore, activity and/or sleep circuits downstream of the I-LNvs and unknown wake-promoting non-clock peptidergic neurons may be gated differentially over the course of the night.

Although the effects on early evening sleep are almost identical, we see significant effects on daytime sleep in both *c929-GAL4;UAS-dTrpA1* and *c929-GAL4;UAS-dTrpA1;pdf-GAL80* flies that we did not see with *pdf-GAL4;UAS-TrpA1*. One possible explanation is that the innately higher levels of male and female daytime sleep in the *c929-GAL4* genetic background allow a bigger dynamic range for inhibition. A more interesting possibility is that this reflects different roles for I-LNvs versus s-LNvs in daytime sleep since the daytime loss is partially rescued on day 2 of 30°C in males ($p < 0.05$ for comparison of *c929-GAL4;UAS-dTrpA1* and *c929-GAL4;UAS-dTrpA1;pdf-GAL80* males).

Our data also imply that PDF receptors must be present on output cells downstream of the circadian/sleep circuit. The anatomical distribution of PDFR has been difficult to ascertain due to the lack of specific antibodies. Using a functional assay of cAMP accumulation, the only cells that have been definitively identified as PDF targets to date are other clock cells (Shafer et al., 2008). To determine if the PDFR might be expressed on cells more directly involved in control of motor activity, we fused 10 kilobases of genomic DNA upstream of the *pdf* gene to GAL4. Figure 6A (left) shows that this promoter region drives expression in a limited number of cells that parallel the distribution of PDF in the adult brain. GFP expression is seen in optic lobes and the dorsal and lateral brain. Lateral brain staining is partly due to expression of this GAL4 line in a subset of both I-LNvs and s-LNvs (example shown in Figure 6A, right). Thirteen brains were imaged and GFP was typically seen in at least one neuron per side in both the s-LNv and I-LNv groups. Dorsal brain staining is seen in the

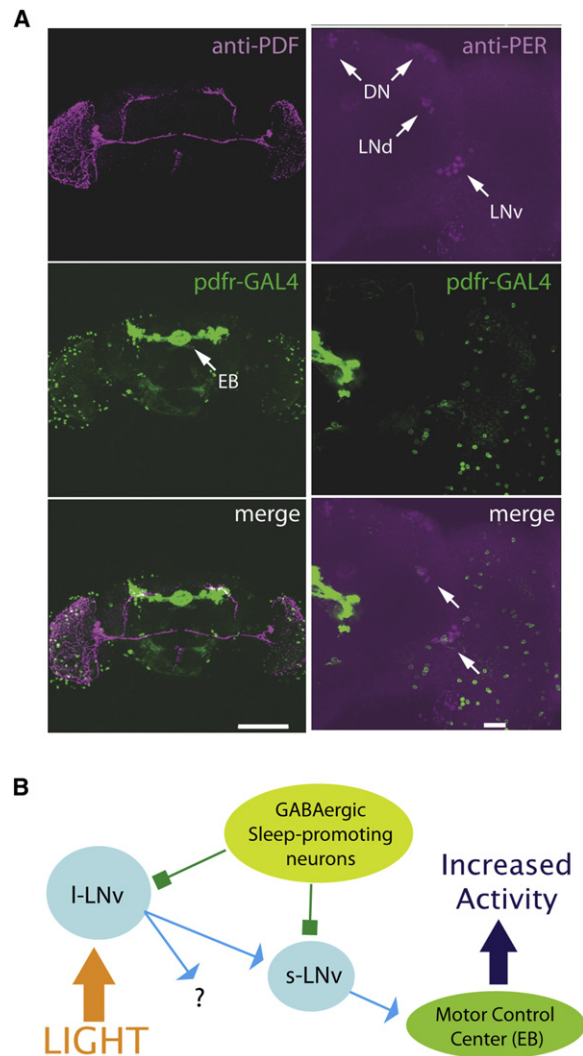


Figure 6. Output of the Sleep Circuit

(A) *pdf-GAL4* marks output cells of the LNv circuit. Left panels show reconstructed *pdf-GAL4;UAS-mCD8-GFP* adult brain stained with anti-PDF (1:1000) and visualized with confocal microscopy. PDF is shown in magenta, GFP in green, overlap in white. Scale bar, 150 μ m. Right panels show a 37 μ m section of a *pdf-GAL4;UAS-mCD8-GFP* adult brain stained with anti-Per (1:100) and visualized with confocal microscopy. Per is shown in magenta, GFP in green, overlap in white. Arrows indicate clock cells. Scale bar, 20 μ m. (B) Model of the *Drosophila* sleep circuit. Light, and perhaps other arousal cues, activates I-LNvs which release PDF onto s-LNvs that project to other clock cells and also send dorsal projections that pass by *pdf-GAL4*-positive cell groups, such as the ellipsoid bodies that are involved in control of activity. Both I- and s-LNvs express GABA_A receptors, allowing sleep-promoting GABAergic neurons to suppress wakefulness.

majority of LNds but not significantly in DNs, although GFP-positive processes from LNds project dorsally. This is consistent with our GAL4 line capturing a subset of the endogenous PDFR clock distribution that has been described. Interestingly, there is also very significant expression in neurons that innervate the ellipsoid bodies, a structure that is part of the central complex and has been implicated in motor control (Strauss, 2002). Ellipsoid

body expression was consistently strong in all three of the lines examined.

DISCUSSION

Using a variety of mutants and novel genetic strategies to manipulate chronic and acute circuit activity, we have shown that a small set of circadian clock cells in *Drosophila* has a critical role in the GABAergic initiation and maintenance of sleep. We have developed new genetic tools (dnATPase, ShawRNAi) that allow an increase in the chronic response of neurons to their endogenous inputs. This adds greatly to the arsenal of activity-manipulating tools, most of which suppress firing or neurotransmitter release. Bidirectional manipulation of activity provides much more information about circuit function and dynamics (c.f. Broughton et al., 2004). We have also shown the utility of a new tool for acute activity manipulation (dTrpA1), which can be used on small numbers of neurons deep within the fly brain. Our data suggest a model (Figure 6B) in which l-LNvs translate light inputs (and perhaps other arousal signals) into wakefulness. The release of PDF from these cells is required, and l-LNv PDF signals to s-LNvs. Our data demonstrating somnolence after downregulation of PDFR in LNvs indicate that s-LNvs participate in sleep control, although experiments in which they have been ablated suggest that they are not the only sleep-relevant l-LNv targets (Sheeba et al., 2008a). PDF signaling to PDFR-expressing neurons outside the clock that directly control activity is likely to be important (see below). GABA may modulate the ability of LNvs to suppress sleep by acting on either or both s- and l-LNvs.

In mammals, the role of the circadian clock in sleep is not completely understood. It is nonetheless clear that there are genetic (e.g., familial advance sleep phase syndrome) and environmental (e.g., jet-lag, shift work) conditions that disrupt sleep despite primarily affecting the circadian rhythms (Gottlieb et al., 2007). The clock has been shown to regulate both when an animal sleeps and how much sleep occurs. The current consensus view is that the mammalian clock is primarily wake-promoting (Edgar et al., 1993; Laposky et al., 2005; Naylor et al., 2000), acting along with the homeostatic sleep drive to shape sleep over the day and night (Dijk and Franken, 2005).

Our data indicate that in flies PDF and the circadian LNvs more generally regulate both the maintenance of sleep as well as the ability of flies to respond to the wake-promoting effects of light. Although these effects recall the role of the mammalian SCN in sleep regulation, there are few prior links between the *Drosophila* circadian clock and the regulation of fly sleep (Shaw et al., 2002). The almost complete elimination of the difference in total sleep between subjective day and subjective night in the *pdf⁰¹* background (Figure 4C) adds substantially to this connection, i.e., light regulation of sleep appears to be substantially circadian clock mediated. Therefore, the contribution of the circadian machinery and fly brain clock circuitry to the control of sleep will probably parallel the important role of the mammalian circadian clock and the SCN in sleep regulation (Borbely and Achermann, 1999; Edgar et al., 1993).

PDF neurons have been recently shown to be light responsive (Sheeba et al., 2007), like some neurons of the mammalian SCN

(Meijer et al., 1998). The l-LNvs also act as the dawn photoreceptor for the clock, sending a reset signal each morning to the rest of the clock (Shang et al., 2008; Sheeba et al., 2008a). There is also good evidence that fly cryptochrome responds directly to light in addition to influencing circadian timekeeping (e.g., Allada et al., 1998; Emery et al., 2000a, 2000b), and a *cry* mutant substantially decreases the PDF neuron acute light response (Sheeba et al., 2007). Therefore, some of the waking effects described here probably reflect a role of PDF cells on acute processes involving light stimulation. Indeed, the phenotypes of flies without PDF or with decreased LNv neuronal excitability resemble some of the acute effects of the loss of orexin/hypocretin in narcoleptic mice (Mochizuki et al., 2004). PDF neurons are also regulated by GABAergic inputs, analogous to those from the basal forebrain that regulate orexin/hypocretin neurons (Henny and Jones, 2006).

Despite these similarities, there are also important organizational differences between systems. Most notable is the wide distribution of sleep circuitry in mammals. There are for example many targets of sleep-promoting GABAergic neurons, and the role of the circadian clock may be largely modulatory (Mistlberger, 2005). The sleep circuitry of flies is almost certainly more circumscribed and simpler. Indeed, the surprisingly large effects of manipulating Rdl in the 16 LNvs argue that they are a principal target of sleep-promoting GABAergic neurons and constitute part of the “core” sleep circuitry. The fact that activation of a subset of these cells, the l-LNvs, has an effect on sleep homeostasis further suggests that these cells sit at the heart of the sleep circuit. The fly sleep circuitry may therefore have condensed mammalian stimulatory systems (e.g., histaminergic, cholinergic, and adrenergic, as well as orexin) into a simpler and more compact region, which may even largely coincide with the 16 PDF cells of the circadian circuit.

A limited number of other fly brain regions have been proposed to contribute to fly sleep. Our manipulations of PHM⁺ cells indicate that peptidergic neurons other than PDF neurons are wake promoting. An attractive hypothesis is that some of these other peptidergic cells reside in the pars intercerebralis, a group of neurohumoral cells shown to be an important sleep output center (Foltenyi et al., 2007). The targets of these cells may even overlap with the targets of LNvs, e.g., the ellipsoid bodies. The PDFR is a class II G protein-coupled receptor and is fairly promiscuous: PDF is the highest affinity ligand, but this receptor is also activated by DH₃₁ and PACAP-38 (Mertens et al., 2005). Since peptidergic modulation may occur by “volume” transmission instead of by direct synaptic contact (Zoli et al., 1999), both LNv peptides and peptides from the pars could together affect this motor center to regulate sleep and activity. The role of the pars may be to inform the sleep-generation machinery about nutritional and metabolic state, i.e., animals undergoing starvation exhibit hyperlocomotor activity that is believed to be evolutionarily useful as a method for finding food (Lee and Park, 2004), and alteration of this pars-generated locomotor program affects sleep (Mattaliano et al., 2007). The role of l-LNvs is clearly different from that of other PHM⁺ neurons, and their unique involvement in homeostatic sleep suggests they are central to sleep control.

The only other brain region that has been implicated in *Drosophila* sleep regulation is the mushroom bodies (Joiner

et al., 2006; Pitman et al., 2006). These studies showed that GAL4-driven manipulation of signaling or of neurotransmitter release in this neuropil had complex effects on sleep, not inconsistent with a modulatory role for this sensory integration center. The exact mechanism of these effects is not clear, however, especially since all of the mushroom body GAL4 lines we have examined also express in multiple subsets of clock cells (data not shown).

The small circuit we describe presents a tractable model system for understanding the circuit-level control of sleep, the relationship between homeostatic and circadian control as well as the dynamics of sleep-wake transitions; the latter are critical to an understanding of episodic and age-related insomnia.

EXPERIMENTAL PROCEDURES

Animals

Flies were raised under a 12 hr light:12 hr dark (LD) schedule at 24°C–25°C on cornmeal dextrose yeast food. Transgenic lines and mutants are as described: *pdf-GAL4* (Renn et al., 1999), *GAD-GAL4* (Mehren and Griffith, 2006), *C380-GAL4* (Packard et al., 2002), *c929-GAL4* (Park et al., 2008), *UAS-mCD8-GFP* (Lee and Luo, 1999), *UAS-dnATPase* (Sun et al., 2001), *UAS-EKO* (White et al., 2001), *UAS-ShawWT* (Hodge et al., 2005), *UAS-dTrpA1* (Hamada et al., 2008), *UAS-Rdl* and *UAS-RdlRNAi* (Liu et al., 2007), *UAS-NaChBacGFP* and *UAS-dORKNCGFP* (Nitabach et al., 2006), *pdf-GAL80* (Stoleru et al., 2004), *pdf⁰¹* (Renn et al., 1999), *UAS-pdfrRNAi* (Dietzl et al., 2007), *UAS-dTrpA1RNAi* (Hamada et al., 2008), and *pdfr^{P2-36}* and its revertant control (Mertens et al., 2005). The *UAS-ShawRNAi* transgene was generated by inserting a 720 bp fragment of the 3' end of the *Shaw* cDNA starting at nucleotide 881 in exon 8 through to the end of the coding region including approximately 110 bp of 3' untranslated sequence into Sym-pUAST-w (Giordano et al., 2002). A transgenic line containing inserts on both chromosomes 2 and 3 was generated by standard methods (Robertson et al., 1988). The *pdfr-GAL4* line was constructed by amplifying 10 kilobases upstream of the ATG of the *pdfr* gene (CG13758) by PCR, subcloning the fragment into the pPTGAL4 vector, and generating transgenic flies by standard methods. Expression patterns from three independent insertion lines were analyzed and found to be essentially identical. Primers used were *pdfr* forward, 5'CCGGCTTTTGTGTTTG TGTTTTG3', and *pdfr* rev, 5'GCCATCGACCGCATAGTAAATG3'. All other lines were obtained from Bloomington Stock Center.

For each LNV manipulation, experimental animals were compared to a control line that was generated by crossing the UAS line to *Df(1)w*, the background strain used to make transgenic lines. This is indicated in the figures as "UAS Control." We find that this type of control is very important to do since the genetic background of strains can have a big influence on basal sleep parameters (c.f. the controls in Figure 3A). Since *pdf-GAL4* is used as a common driver for all LNV manipulations, and therefore cannot contribute to differential phenotypes. Baseline data for this GAL4 line is shown in Figure 3E. *GAD-GAL4* has previously been shown to sleep normally (Agosto et al., 2008). For *pdf⁰¹*, the mutant was extensively outcrossed to *Canton S* wild-type and mutant and sibling control lines established using PCR genotyping. For *pdfr* the precise excision strain (Mertens et al., 2005) was used as a genetic background control. For *c929-GAL4* experiments, a control line expressing a dead channel protein (dORKNC) was used as a control line since the *c929-GAL4* genetic background has relatively high basal daytime sleep compared to other genotypes.

Sleep and Activity Assays

All behavior was done on female flies unless explicitly indicated. Five-day-old flies were placed in 65 mm × 5 mm glass tubes (Trikinetics, Waltham, MA) containing 5% agarose with 2% sucrose. Flies were acclimated in behavior tubes for at least 24 hr at 25°C (or 22°C where indicated) in 12 hr light/12 hr dark (LD) conditions before data collection. Flies were entrained at least 4 days in LD before switching to constant darkness (DD). Locomotor activity was collected

with DAM System monitors (Trikinetics) in 1 min bins as previously described (Agosto et al., 2008). Sleep was measured as bouts of uninterrupted 5 min of inactivity. Sleep parameters were analyzed using MATLAB software (Natick, MA) of averages over 4 days of LD. Total sleep duration, mean sleep and wake bout duration, and latency were analyzed for each 12 hr period of LD and DD and averaged over 3 days for each condition. Arousal state stability was calculated by subtracting the maximum sleep bout duration from the maximum wake bout duration. Values greater than one indicate a more wakeful state while values less than one are indicative of a stable sleep state.

Immunohistochemistry, Imaging, and Quantification

For determination of *Shaw* RNAi efficacy, adult brains from *C380-GAL4;mCD8-GFP* animals were processed and stained with anti-Shaw antibody (preabsorbed and used at 1:1000) and Cy5 secondary antibody (1:180; Jackson Labs) as described (Hodge et al., 2005). All preparations were processed in parallel and images acquired with identical settings using the 50× (zoomed 1–4×) objectives of a Leica TCS SP2 confocal microscope. Care was taken to keep all intensity readings within the linear range below saturation. Quantification was performed on 1 μm sections with pixel intensity readings taken in a given region of interest (in this case the mushroom bodies) for GFP and Cy5 using the Leica TCS SP2 quantification software. Quantification was performed blind to genotype. Statistical analysis was performed in Excel (Microsoft) and JMP (SAS). Significance levels in figures were determined by one-way ANOVA unless otherwise specified, and * indicates $p < 0.05$.

For Rdl localization, adult brains from *pdf-GAL4;mCD8-GFP* animals that had been entrained in a 12 hr L:12 hr D cycle for at least 3 days were dissected, fixed, and stained basically as described (Van Vector et al., 1991) with anti-Rdl (1:100) (Liu et al., 2007) and anti-GFP (1:200, Roche Applied Biosciences), and Alexa 635 and 488 secondary antibodies (1:200, Invitrogen). For costaining with *pdfr-GAL4*, *pdfr-GAL4;UAS-mCD8-GFP* animals were entrained in a 12 hr L:12 hr D cycle for at least 3 days and were dissected, fixed, and stained as described (Van Vector et al., 1991) with anti-GFP (1:300, Roche Applied Biosciences) + Alexa 635 secondary (1:200, Invitrogen) and either anti-PAP (PDF precursor) (1:1000) + Texas Red secondary (1:200, Jackson Labs) or anti-Per (1:1000) + Alexa 488 secondary antibody (1:200, Invitrogen). Images were acquired on a Leica TCS SP2 confocal system with Leica Confocal Software at 63×. Separate images were taken using 488 nm and 633 nm lasers and overlapped to avoid bleed through. Leica Confocal Software was used to quantify the images. Quantification was performed using the first scan taken at 633 nm excitation. After signal digitization RMS values of background were subtracted to get final values.

Immunoblotting

Age matched flies were frozen in liquid nitrogen and decapitated by vortexing. Extracts were prepared as described (Hodge et al., 2006), separated by SDS-PAGE, and analyzed by immunoblot using rabbit anti-C-terminal Shaw antibody (1:1000), monoclonal anti-tubulin (1:200,000; Sigma).

Electrophysiology

Whole-cell recordings were performed on third instar larvae using methods described previously (Choi et al., 2004). Larvae were cut open dorsally and pinned down onto a sylgard-lined dish in calcium-free solution consisting of (in mM) 128 NaCl, 2 NaOH, 2 KCl, 15 sucrose, 5 trehalose, 4 MgCl₂, and 5 HEPES, with pH 7.1–7.2. Sheath tissue surrounding dorsal motoneuron clusters was digested with 0.01% protease (type XIV, Sigma). Motor neuron MNISN-Is was targeted exclusively for all experiments. Pipette resistance was 5–10 MΩ, and solution contained (in mM) 130 potassium gluconate, 10 HEPES, 1 EGTA, 2 MgCl₂, 0.1 CaCl₂, 2 NaCl, 10 KOH, with pH adjusted to 7.2. An Axopatch 200B amplifier (Axon Instruments, Union City CA) was used to perform whole-cell recordings and acquisition and analysis performed with IgorPro (Wavemetrics, Oswego, OR). Two-tailed, unpaired t tests or ANOVA repeated-measures test was used to analyze significance of values when comparing two genotypes (Statview software package, Abacus Concepts, Cary NC). Multiple comparisons were done using ANOVA with the Tukey-Kramer post hoc test for pair-wise comparisons.

SUPPLEMENTAL DATA

The Supplemental Data include five figures and can be found with this article online at [http://www.neuron.org/supplemental/S0896-6273\(08\)00942-2](http://www.neuron.org/supplemental/S0896-6273(08)00942-2).

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