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Impaired clock output by altered connectivity in the circadian network

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Substantial progress has been made in elucidating the molecular processes that impart a temporal control to physiology and behavior in most eukaryotes. In *Drosophila*, dorsal and ventral neuronal networks act in concert to convey rhythmicity. Recently, the hierarchical organization among the different circadian clusters has been addressed, but how molecular oscillations translate into rhythmic behavior remains unclear. The small ventral lateral neurons can synchronize certain dorsal oscillators likely through the release of pigment dispersing factor (PDF), a neuropeptide central to the control of rhythmic rest-activity cycles. In the present study, we have taken advantage of flies exhibiting a distinctive arrhythmic phenotype due to mutation of the potassium channel *slowpoke* (*slo*) to examine the relevance of specific neuronal populations involved in the circadian control of behavior. We show that altered neuronal function associated with the null mutation specifically impaired PDF accumulation in the dorsal protocerebrum and, in turn, desynchronized molecular oscillations in the dorsal clusters. However, molecular oscillations in the small ventral lateral neurons are properly running in the null mutant, indicating that *slo* is acting downstream of these core pacemaker cells, most likely in the output pathway. Surprisingly, disrupted PDF signaling by *slo* dysfunction directly affects the structure of the underlying circuit. Our observations demonstrate that subtle structural changes within the circadian network are responsible for behavioral arrhythmicity.

circadian circuitry | *Drosophila* | pigment dispersing factor | potassium channels | *slowpoke*

Rhythmicity in rest-activity cycles in *Drosophila* is under control of the circadian clock, which is based on self-sustaining, cell-autonomous transcriptional negative feedback loops. These feedback loops ultimately give rise to rhythms in the abundance, phosphorylation state, and nuclear localization of key intracellular proteins, such as period (PER) and timeless (TIM) (1). To date, several neuronal clusters have been shown to include a molecular oscillator. The one best understood encompasses the small ventral lateral neurons (LN_vs), comprised of five cells, of which four rhythmically release the neuropeptide pigment dispersing factor (PDF) at their dorsal terminals. Other oscillators within the fly brain include the dorsal lateral neurons (LN_ds) together with the dorsal neurons (DN1–3) (2). Ablation of all LN_vs by overexpression of proapoptotic genes, as well as null mutations on the *pdf* gene or its receptor, cause behavioral arrhythmicity a few days upon transfer to constant conditions (3–6) likely through the gradual loss of synchronization among the components of the small LN_v cluster (7).

The question of how the intracellular molecular oscillations taking place within specific neuronal clusters ultimately drive rhythmic locomotor activity has only recently been approached in *Drosophila* (7–11). Molecular oscillations must be somehow transduced into neuronal function to generate a rhythm in behavior and physiology. Increasing evidence places electrical activity as an essential element in the propagation of such

circadian oscillations. A possible mechanism to control membrane excitability is the circadian control of ion channel mRNA levels, which takes place in *Drosophila* (12, 13) and mammals (14). Free-running circadian rhythms in membrane conductance and K⁺ channel current have been observed in pacemaker neurons of the molluscan retina (15, 16) and in mammals (17, 18). Alternatively, activation of second messenger cascades (such as calcium-dependent signaling) could also serve this purpose. In the suprachiasmatic nuclei cytoplasmic calcium levels have been shown to oscillate in a circadian manner (19, 20), with high calcium levels during the day and low levels at night, again serving as a possible link to control active membrane properties.

Drosophila slowpoke (*slo*) is the first voltage-gated calcium-dependent potassium channel of the “Big K” family to be cloned and characterized (21). A null mutation in the *slo* locus strikingly alters the electrical properties of both neurons and muscles (22–24), affecting neurotransmitter release (25) and thereby causing a variety of behavioral defects among those in courtship behavior (26–28). We have previously demonstrated that a *slo* null mutant is behaviorally arrhythmic under free-running [constant darkness (DD)] conditions (13). In the present work, we show that the loss of *slo* function in neuronal tissues is responsible for the highly arrhythmic phenotype. Whole-mount adult brain immunohistochemistry indicates that molecular oscillations are still intact in the small LN_vs, whereas they are desynchronized in all dorsal clusters. Rhythmic PDF release is impaired in the null mutant in DD, thus explaining the lack of synchronicity between the ventral and dorsal oscillators. Remarkably, not only PDF levels, but also the proper arborization of the PDF terminals, are affected in the mutant. We propose that the electrical activity in the dorsal SLO⁺ clusters is impinging on the circadian control of the neuropeptide and the underlying PDF⁺ circuit, pointing to the SLO-expressing neurons as an intermediate relay circuit in the pathway relevant for rhythmic control of behavior.

Results

Lack of SLO Function in the CNS Accounts for Behavioral Arrhythmicity. *slo* is highly regulated at the transcriptional level, employing several alternative promoters and splice variants (29–31). To distinguish between a neuronal and/or a muscular defect associated with the strong impairment observed in *slo* mutants, we

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The authors declare no conflict of interest.

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Abbreviations: CT, circadian time; DD, constant darkness; DD3, 3 days after transfer to DD; DN_s, dorsal neurons; LD, light–dark; LN_ds, dorsal lateral neurons; LN_vs, ventral lateral neurons; PDF, pigment dispersing factor.

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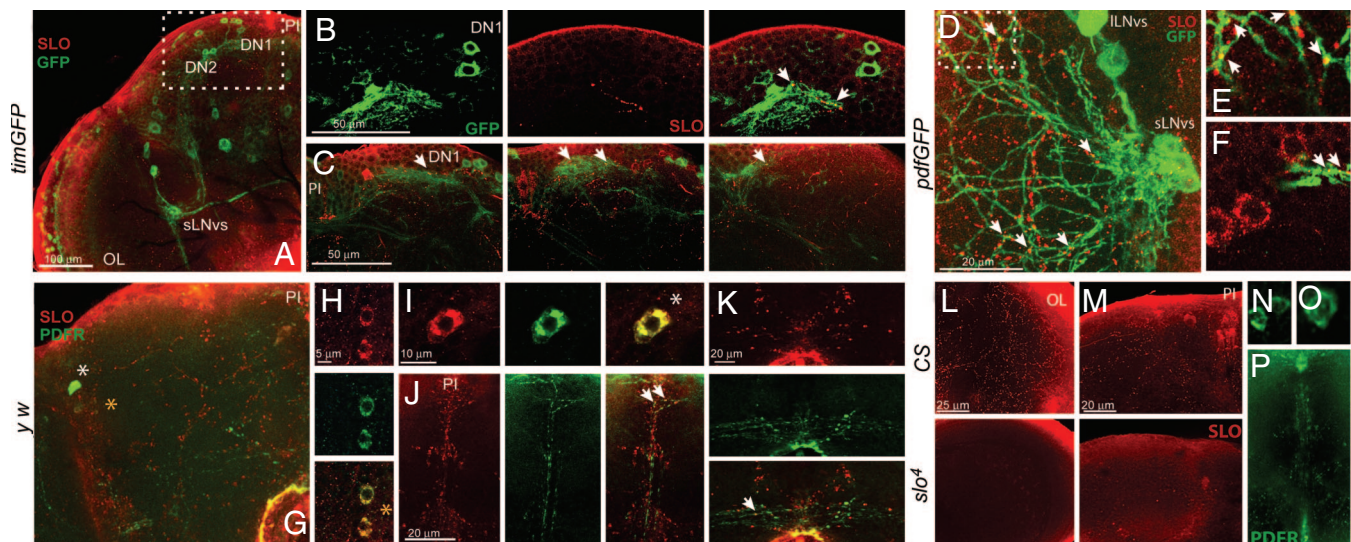


Fig. 2. SLO expression in the context of circadian structures. (A–F) *tim-Gal4/UAS-CD8GFP* (*timGFP*, A–C) and *pdf-Gal4/UAS-CD8GFP* (*pdfGFP*, D–F) brains were stained with anti GFP (green) and pSLO (red). Images shown in A, D, G, L, and M are projections, whereas the remaining ones are single optical scans. Arrows indicate colocalization. (A) SLO expression in a brain hemisphere displaying GFP+ *tim* neurons. (B) Inset from a dorsal region indicated in A, displaying axons from DN1 juxtaposed to SLO+ neurites. (C) Inset from the same region of a different brain displaying three consecutive scans to illustrate how GFP+ axons reach a region where a cluster of SLO+ somas is located. (D) Extensive ramification of PDF and SLO+ neurites in the optic lobe. (E) Inset from D showing colocalization between both signals. (F) A higher magnification of a dorsalmost segment of the small LNvs projections from another brain showing SLO+ somas in the region where these PDF+ projections reach their targets. (G–K) Wild-type brains stained for PDFR (green) and SLO (red), revealing a widely distributed punctate signal. Symbols depict the anatomical regions from which the single focal planes were taken. Each image corresponds to a different brain. (H and I) Magnified views of a region similar to that indicated in G, revealing colocalization of SLO and PDFR in some of the cell bodies in which the two are expressed. (J and K) Images display punctate-like signal along the midline and the main commissure, respectively. (L and M) The specificity of the pSLO antibody was assessed comparing the signal in the optic lobe (L) and the dorsal area (M) between CS and *slo*⁴. (N–P) PDFR signal in *slo*⁴ brains in the regions shown in H–J, respectively.

clearly indicating that this circadian-controlled behavior was unaffected in the mutant (SI Fig. 6B). However, only wild-type flies displayed a morning peak (SI Fig. 6A), suggesting that the molecular oscillators sustaining the morning and evening components of the rest-activity pattern under synchronized conditions were distinctly affected by the lack of *slo* function, which is also consistent with an impaired output from the central pacemaker cells (8, 9).

Molecular Oscillations Are Preserved in *slo*⁴ Pacemaker Cells. Abnormal excitability and firing pattern in the circuits underlying rhythmic behavior could alter the core molecular oscillations (37) and serve to explain the lack of rhythmicity observed in *slo*⁴.

To directly assess the pace of the intracellular molecular clock, whole-mount brain immunohistochemistry was carried out to monitor PER and TIM subcellular distribution in the small LNvs. Interestingly, *slo*⁴ samples were undistinguishable from Canton S (CS) throughout time courses performed in LD and 3 days after transfer to DD (DD3) (data not shown and Fig. 3, respectively). In the small LNvs of CS and *slo*⁴, PER was found in the nucleus at CT5 and CT11. By CT17, intense cytoplasmic PER was detected, although a mild nuclear signal could be seen. At CT23, PER protein was exclusively located within the nucleus. In contrast, TIM was undetectable until CT17, when it was mainly located in the cytoplasm, and it became nuclear by CT23. Remarkably, no differences were detected in the pace of these

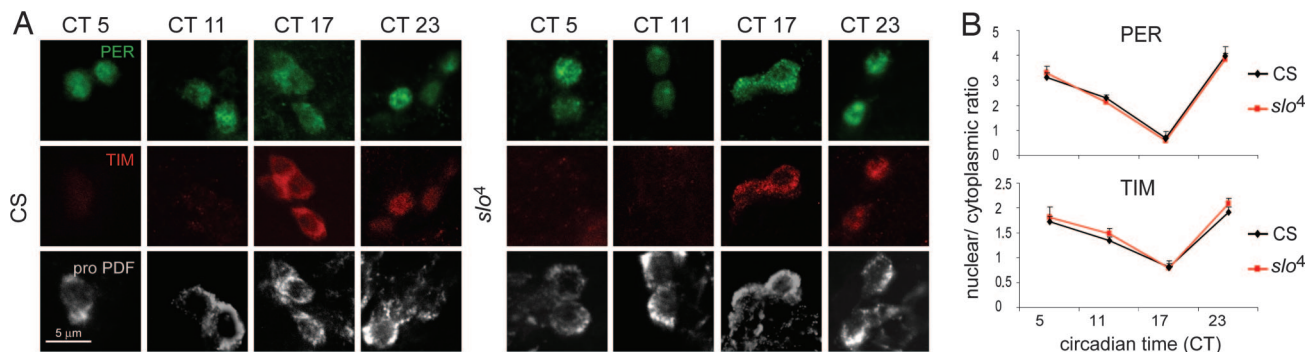


Fig. 3. Molecular oscillations in pacemaker neurons are not affected in *slo*⁴. Newly eclosed CS and *slo*⁴ adult flies were synchronized, and samples were taken in DD3. Whole-mount brain immunohistochemistry was performed to follow TIM (red) or PER (green) and proPDF (white) accumulation at CT5, CT11, CT17, and CT23. A minimum of 12 brains per genotype were analyzed at each time point. (A) Representative confocal stacks focused on the small LNvs in wild-type flies (Left) and *slo*⁴ (Right). Time courses were analyzed blind and repeated three to five times with identical results. (B) Ratio between nuclear and cytoplasmic PER/TIM signal. Error bars represent the SE of the mean. No statistical differences were found between both genotypes ($P > 0.05$). See Materials and Methods for details on quantitation methods and statistics.

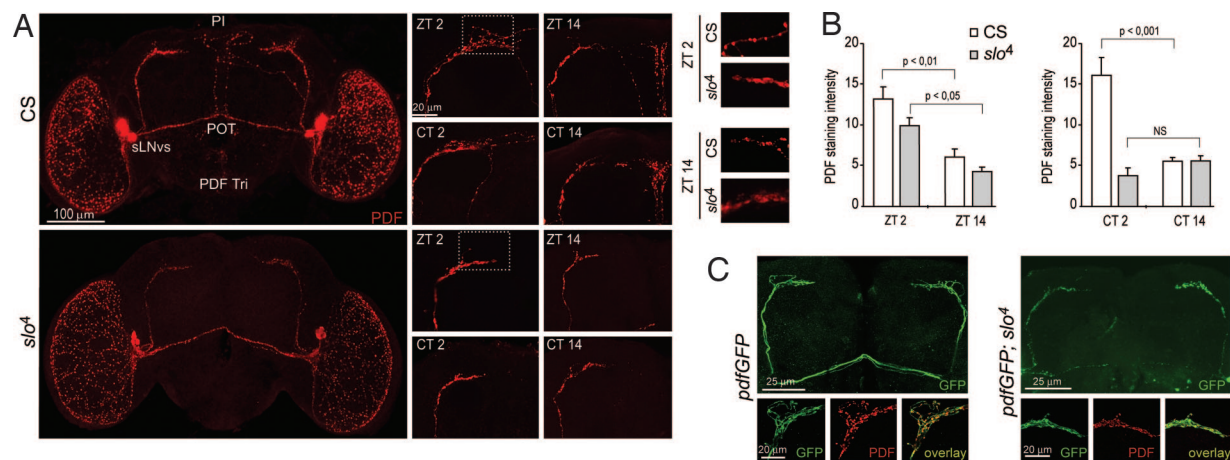


Fig. 5. Mutation in *slo* affects normal axonal arborization of pacemaker cells. (A) (Left) PDF secretion or accumulation is altered in *slo*⁴. Whole-mount brain immunohistochemistry of CS and *slo*⁴ flies stained for PDF. Brains <1 day old were fixed at the times indicated in the figure (CT samples were taken on DD2). Experiments were repeated at least three times, and a minimum of 50 brains were analyzed per time point. (Right) Magnified views of the PDF+ projections in the dorsal region. Box shows the area quantified in B. PI, *pars intercerebralis*; POT, posterior optic tract. (B) Quantitation of the average intensity of the LNvs dorsal projections. Error bars represent the SE of the mean. No circadian differences in PDF intensity in *slo*⁴ were found ($P > 0.05$). See *Materials and Methods* for details on quantitation methods and statistics. (C) (Upper) Evaluation of the structure of the PDF projections in *pdfGFP* and *pdfGFP;slo*⁴ flies stained for PDF (red) and GFP (green). (Lower) Higher magnification view of the dorsalmost segment.

PDF signals colocalized throughout the circuit and the PDF+ puncta were contained within GFP-labeled membranes of the dorsal termini. Strikingly, in *pdfGFP;slo*⁴ brains, both the PDF and GFP signals colocalized, indicating that loss of SLO function not only affected the accumulation of the neuropeptide, but also altered the axonal arborization of the PDF+ neurons (Fig. 5C).

Therefore, loss of SLO function specifically affected PDF axonal arborization, preventing a proper synchronization of the circadian network, and thus impacted on rhythmic behavior.

Discussion

Lack of Neuronal SLO Correlates with the Arrhythmic Behavior. We previously demonstrated that a null mutation in a clock-controlled gene that encodes the calcium-dependent voltage-gated potassium channel *slo* gives rise to a highly arrhythmic locomotor behavior phenotype (13). In the present work, we extended these original observations by using a transgenic line that expresses the most prevalent neuronal SLO version throughout the brain and has been shown to rescue other neuronal deficits of *slo*⁴ (32) to demonstrate that it is the lack of channel activity in CNS neurons that gives rise to the full phenotype (Fig. 1). Employing different entraining paradigms, we ruled out an inability to respond to light as the major component for the arrhythmicity (SI Fig. 6).

The Defect Lies Downstream of Pacemaker Function. The lack of rhythmicity characteristic of *slo*⁴ could derive from an abnormal core oscillator function, which would manifest at the level of the central pacemaker neurons and other peripheral tissues. We first investigated molecular oscillations in the small LNvs because their hierarchy over other brain oscillators in the rhythmic control of locomotor behavior is well documented (9, 10, 41, 42). We monitored PER and TIM oscillations on whole-mount brains and found neither changes in subcellular localization nor differences between the amplitude of the rhythmic oscillations in the *slo*⁴ background compared with wild-type controls (Fig. 3 and SI Fig. 7). These results clearly indicate that, regardless of the nature of the arrhythmicity, lack of SLO does not affect the underpinnings of the self-sustained molecular clock in every cell. Moreover, despite the potential defects in the excitability of the underlying circuits in *slo*⁴, no alterations or rundown of the molecular oscillations in pacemaker cells were evidenced (37, 43).

Lack of SLO Function Differentially Affects Dorsal Oscillators Under Free-Running Conditions. Not surprisingly, given that most circadian structures are light-responsive per se, no difference was observed between CS and *slo*⁴ when the pace of the oscillations in dorsal clusters was evaluated in LD, suggesting that they are not directly responsible for the weak rhythmicity of the rest-activity cycles in *slo*⁴. However, when molecular oscillations were evaluated under free-running conditions, a lack of synchronization was detected between the ventral and dorsal clusters (Fig. 4 and SI Fig. 8), in contrast to what has been reported for a null mutant in the channel narrow abdomen (44). Our data strongly suggest that dorsal oscillators are clearly taking part in the circuitry behind rhythmic control of rest-activity cycles under free-running conditions as previously suggested (38), and the hierarchy of the small LNvs is most likely exerted through the release of PDF at the dorsal terminals.

Abnormal PDF Projections Account for Lack of Rhythmicity Under Free-Running Conditions. *slo*⁴ mutants appeared to have intact PDF neurons. However, the dorsal projections toward the *pars intercerebralis* were clearly disrupted (Fig. 5A and B). We hypothesized that the pervasive arrhythmicity observed in this mutant was linked to the absence of proper PDF processes. We then evaluated its integrity in the neuronal rescue (*B52H;slo*⁴) and found that most brains showed strong PDF labeling (data not shown). Thus, distorted electrical activity in *slo*⁴ mutant flies is likely responsible for the abnormal PDF release and/or accumulation.

We also provide evidence that loss of SLO function altered the degree of axonal arborization within that region (Fig. 5A Inset and C). Labeling the PDF-expressing neurons by independent means (*pdfGFP;slo*⁴) showed no gross anatomical changes in *slo*⁴. That the lack of SLO function affects the connectivity of the PDF neurons suggests that its function is relevant in related circuits whose activity affects the maintenance or plasticity of PDF+ oscillators (or both). In any case, the observation that dorsal SLO neurons could potentially receive synaptic connections from projections of both ventral and dorsal oscillators (Fig. 2C and F) strongly suggests that the dorsal SLO-expressing neurons are indeed part of the network required for rhythmic behavior.

