Connecting the Navigational Clock to Sun Compass Input in Monarch Butterfly Brain

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Summary

Migratory monarch butterflies (Danaus plexippus) use a time-compensated sun compass to navigate to their overwintering grounds in Mexico. Although polarized light is one of the celestial cues used for orientation, the spectral content (color) of that light has not been fully explored. We cloned the cDNAs of three visual pigment-encoding opsins (ultraviolet [UV], blue, and long wavelength) and found that all three are expressed uniformly in main retina. The photoreceptors of the polarization-specialized dorsal rim area, on the other hand, are monochromatic for the UV opsin. Behavioral studies support the importance of polarized UV light for flight orientation. Next, we used clock protein expression patterns to identify the location of a circadian clock in the dorsolateral protocerebrum of butterfly brain. To provide a link between the clock and the sun compass, we identified a CRYPTOCHROME-staining neural pathway that likely connects the circadian clock to polarized light input entering brain.

Introduction

Remarkable progress has been made in understanding the molecular and cellular basis of circadian clocks in animals over the past decade (Stanewsky, 2002; Reppert and Weaver, 2002). One of the most intriguing functions of circadian clocks—their use in time-compensated sun compass navigation (von Frisch, 1967; Kramer, 1957)—has received only scant attention, however. From a functional vantage point, it is clear that time compensation is an absolute requirement for successful migration and, ultimately, survival, if celestial cues are used for long-distance navigation (Froy et al., 2003).

During their spectacular fall migration, North American monarch butterflies (Danaus plexippus) use a time-compensated sun compass to help them navigate to their overwintering sites in central Mexico (Perez et al., 1997; Mouritsen and Frost, 2002; Froy et al., 2003). Because this navigational capability is largely genetically determined (Brower, 1996), we are developing the monarch butterfly as a model to study the molecular and cellular basis of time-compensated sun compass navigation. Our ultimate goal is to understand the molecular and anatomical mechanisms for circadian clock interactions with the sun compass that enable migrants to maintain a fixed south/southwesterly flight bearing as the sun moves across the sky each day.

Previous studies suggest that monarch butterflies can use the angle of plane polarized light (e-vector) as an orientation (sun compass) cue (Hyatt, 1993; Reppert et al., 2004), and based on studies in hymenopterans (Brines and Gould, 1979; Edrich et al., 1979; Wehner, 1997), monarchs might also use the sun itself and/or spectral-intensity gradients. Because ultraviolet (UV) light is important for the initiation of oriented flight in monarch butterflies (Froy et al., 2003), it has been difficult to determine solely by behavioral assays whether polarized light in the UV range is important for orientation during sustained flight.

Like many insects (Lahbhart and Meyer, 1999), monarchs possess photoreceptors in the dorsal rim area (DRA) of the compound eye that are anatomically specialized for polarized-light detection (Reppert et al., 2004). Because the DRA of Drosophila melanogaster has a unique photoreceptor-specific pattern of opsin expression relative to the main retina (Wernet et al., 2003), we wondered whether a similar pattern of opsin expression occurs for the DRA of monarch butterflies.

Our previous work with monarch butterflies has also shown that disruption of the molecular circadian clockwork by constant light disrupts the time-compensated component of time-compensated sun compass orientation behavior (Froy et al., 2003). The molecular clock was monitored in that study by assessing the rhythmic expression of monarch period (per) mRNA levels in whole heads, because, as discussed below, per is an essential component of the circadian clock of Drosophila. It has not been known, however, where in monarch brain a circadian clock actually resides.

In Drosophila, the intracellular clock mechanism involves transcriptional feedback loops that drive persistent rhythms in mRNA and protein levels of key clock components (Stanewsky, 2002). The negative feedback
loop involves the dynamic regulation of the per and timeless (tim) genes. The resultant PER and TIM proteins heterodimerize and translocate back into the nucleus where PER inhibits transcription. TIM appears to regulate PER protein stability and nuclear transport and is also necessary for photic responses that reset (entrain) the circadian clock (see Williams and Sehgal, 2001). Cryptochrome (CRY) is colocalized with PER and TIM and is a blue-light photoreceptor involved in photic entrainment (Emery et al., 1998; Stanewsky et al., 1998; Emery et al., 2000).

On the basis of the Drosophila clockwork, we reasoned that the evaluation of clock protein expression in monarch butterfly brain (e.g., the expression patterns of PER, TIM, and CRY) could reveal the location of a circadian clock involved in navigational activities. Indeed, in another lepidopteran, the Chinese oak silkmoth Antheraea pernyi, PER and TIM are rhythmically expressed in brain in four large Ia neurons in each dorsolateral hemisphere that have been proposed to function as circadian clock cells (Sauman and Reppert, 1996). Interestingly, PER and TIM are primarily cytoplasmic throughout the 24 hr day in the silkmoth neurons, whereas in Drosophila pacemaker neurons these proteins undergo temporally regulated nuclear transport that is critical for clock function (Stanewsky, 2002).

In the current report, we set out to provide a firm link between the circadian clock and the sun compass. First, we determined the pattern of opsin expression in the DRA photoreceptors of the monarch to identify the spectral content of polarized light that might be involved in flight orientation. We cloned three opsin-encoding cDNAs and ascertained that all three are expressed uniformly in the main retina. The DRA, on the other hand, expresses only the UV opsin. We complemented these anatomical findings with behavioral studies and found that polarized light in the UV range is important for flight orientation.

Next, we examined the expression profiles of PER, TIM and CRY to identify the location of a circadian clock in monarch butterfly brain. We found strong evidence for a brain clock in the dorsolateral protocerebrum. We found that a cryptochrome-containing neural pathway may connect the circadian clock to polarized light input entering the brain from the DRA of compound eye. Taken together, the results provide insights into a “navigational clock” and its connection to a sun compass in monarch butterfly brain.

Results and Discussion

Opsin Sequences and Phylogeny

Based on the wavelengths for peak sensitivity, rhodopsins expressed in the photoreceptors of the insect compound eye are classified as short wavelength (indicating a peak sensitivity of 300–400 nm), middle wavelength (400–500 nm) and long wavelength (LW, 500–600 nm). Molecularly, these physiological classifications correspond to three major clades of rhodopsin apoproteins (opsins): UV, blue, and LW. Our use of the term ‘opsin’ throughout this paper refers to opsins that are homologous to those found in the photoreceptors of the insect compound eye.

Using monarch head cDNA, we cloned the full-length cDNAs for three opsin genes, which, based on phylogenetic analyses, represent a UV opsin (DpUVRh, GenBank AY605546), a blue opsin (DpBlueRhs, AY605544), and a long wavelength opsin (DpLWRh, AY605545) (see Figure S1 in the Supplemental Data available with this article online). The LW opsin likely has peak spectral sensitivity in the green (ca. 550 nm), based on study of the visual pigment absorption spectrum maxima of the closely related Queen butterfly, Danaus gilippus (Briscoe and Chittka, 2001). The monarch opsin cDNA sequences are highly homologous to the overlapping portions of the UV, blue, and LW (green) opsin mRNA transcripts that have been localized to the photoreceptor cells of the retina of Vanessa cardui, the only other butterfly of the family Nymphalidae so far examined (Briscoe et al., 2003) (data not shown).

The existence of only one LW opsin in these two nymphaid butterflies differs from the situation in butterflies of the family Papilionidae, which generally have several LW opsins expressed in retina (Briscoe, 2000) (Figure S1). In fact, phylogenetic analysis of the opsin gene family indicates that two duplications of an ancestral LW opsin gene occurred along the papilionid lineage (Briscoe, 2001). Importantly, the heterogeneous coexpression of filtering pigments with the LW opsin in monarch retina expands effectively the number of LW-sensitive receptors available for color vision (unpublished data). It is therefore likely that monarchs have tetrachromatic vision based primarily on three opsins and a lateral filtering pigment (also see below).

Uniform Pattern of Opsin Expression in the Main Retina

The basic unit of the butterfly compound eye, the ommatidium, contains nine photoreceptor cells (R1–R9) that express the light-sensitive visual (opsin) pigments embedded in microvillous membranes of a fused rhabdom (Figure 1). The monarch photoreceptor cells are arranged in a semi-tiered fashion (Figure 1B), in which the cell bodies of the R1 and R2 cells are widest near the crystalline cone and then taper considerably as they approach the basement membrane. The tiny R9 cell sits just above the basement membrane. To study opsin expression in R1–R9 throughout the monarch eye, we made digoxigenin-labeled antisense riboprobes to the UTRs of the opsin cDNAs (to avoid cross-hybridization) and examined their pattern of hybridization on cryostat-sectioned tissue.

In situ hybridizations on semitangential sections and longitudinal sections through the whole retina revealed a uniform dorsal-to ventral distribution of DpUVRh and DpBlueRh mRNA expression (Figure S2). Examination of monarch retina cross-sections, longitudinal sections, and dissected ommatidia showed that DpUVRh mRNA expression in the main retina was restricted to the R1 and R2 photoreceptor cells (Figure 2A). A survey of 102 ommatidia of the lateral retina revealed that 17% showed DpUVRh mRNA expression in both R1 and R2, 54% showed expression in either R1 or R2, and 29% did not express DpUVRh mRNA. DpBlueRh expression was also restricted to R1 and R2, which did not overlap with DpUVRh expression (Figure 2, compare [A] and
Figure 1. Anatomical Overview of the Compound Eye of the Monarch Butterfly
(A) Diagram of a longitudinal section through the compound eye. Black dots indicate the location of photoreceptor nuclei. R, retina; L, lamina; M, medulla.
(B) Schematic of a monarch ommatidium showing the R1–R9 photoreceptor cells in the main retina, as well as the presence of dark orange pupillary pigments (DO, black dots) that facilitate adaptation of the eye under different illuminant conditions. Heterogeneously expressed orange filtering pigments (O, gray line and circles) produce more than one spectral type of photoreceptor for long-wavelength light (unpublished data). Cross-hatched areas indicate the fused microvillous membranes of the rhabdomeres that contain the visual pigment protein. C, cornea; CC, crystalline cone; n, nuclei.
(C) DIC image of a tangential section of the retina showing the R1–R8 photoreceptor cells arranged in a radial fashion around the rhabdom.

Figure 2. Opsin Expression in the Main Retina
(A) Tangential section of ommatidia (14 μm) showing DpUVRh mRNA expression in R1 and R2 photoreceptor cells.
(B) Tangential section of ommatidia showing DpBlueRh mRNA expression in R1 and R2 that was nonoverlapping with DpUVRh expression. Dotted circles indicate identical ommatidia in adjacent sections.
(C) Tangential section showing DpLWRh mRNA expressed in the R3–R8 photoreceptor cells in all ommatidia in the main retina.

A survey of 102 ommatidia of the lateral retina revealed that 27% showed DpBlueRh mRNA expression in both R1 and R2, 52% showed expression in either R1 or R2, and 21% did not express DpBlueRh mRNA (Figure 2B). These percentages suggest that the majority, if not all of the R1 and R2 cells in the main retina, express either DpUVRh or DpBlueRh. DpLWRh mRNA was expressed in photoreceptor cells R3 through R8 in all ommatidia of the main monarch retina, which was apparent in cross-sections, longitudinal sections, and dissociated ommatidia (Figure 2C, data not shown). The DpLWRh mRNA transcript was also detected in the apical R9 photoreceptor in dissociated ommatidia (data not shown).

Collectively, the expression patterns of the three opsin mRNAs in monarch main retina indicate nonoverlapping opsin expression in each photoreceptor cell. Furthermore, the mutually exclusive expression of the UV and blue opsin mRNAs in R1 and R2 photoreceptors gives rise to three types of ommatidia in the main retina: those expressing the UV and LW opsins, those expressing the blue and LW opsins, and those expressing all three. The opsin expression patterns in monarch retina are similar to those in Vanessa, the only difference being the existence of a dorsoventral gradient in Vanessa; that is, the dorsal retina mainly consists of UV and LW opsin-expressing ommatidia (Briscoe et al., 2003).

The uniform dorsoventral distribution of UV and blue opsin expression in the main retina of monarch butterflies suggests that if skylight spectral-intensity gradients are used in sun compass orientation, the gradients would be sensed throughout the retina and would not be confined to a particular region.

The Dorsal Rim Area Expresses Only the UV Opsin
We next focused our studies of opsin expression on the DRA, which has ommatidia with rhabdomeres that are...
Figure 3. Opsin Expression in the Dorsal Rim Area
(A) Two to three rows of the ommatidia in the dorsal rim area (DRA) express DpUVRh mRNA in R1–R8 photoreceptor cells, in contrast to adjacent ommatidia, which express the DpUVRh mRNA in only the R1 and/or R2.
(B) Section adjacent to (A), showing the complete absence of DpBlueRh mRNA expression in the DRA ommatidia and expression of the same transcript in R1 and R2 in adjacent retina.
(C) Section adjacent to (B), showing the complete absence of DpLWRh mRNA expression in both the DRA and ommatidia in adjacent retina.
(D) Localization of the UV opsin protein in DRA by immunocytochemistry.

specialized for polarized-light detection (Reppert et al., 2004), to determine what spectral type of photoreceptor might be most important for detecting polarized skylight.

Strikingly, an area at the dorsalmost margin of the monarch eye stands out from the rest of the retina: cross-sections of this area showed that the R1 through R8 photoreceptor cells express only DpUVRh mRNA (Figure 3A). As observed in dissociated ommatidia, this strong expression is present throughout the entire length of the photoreceptor cells (not shown), although we were unable to determine conclusively that DpUVRh mRNA is expressed in R9 cells in the DRA. No DpBlueRh or DpLWRh mRNA was detected in the DRA ommatidia, though it is clearly present in the adjacent ommatidia of the main retina (Figures 3B and 3C). The DRA, as defined by these ommatidia expressing only the UV opsin, occurred within a distinct area 40 to 50 ommatidia in length and situated at the dorsal margin of the eye close to the antenna base. It is one to three ommatidia in depth from the eye margin (Figure 3A) and includes a total of approximately 100 ommatidia in each eye (n = 3).

The DRA ommatidia also showed strong UV opsin immunoreactivity with prominent staining of the rhabdoms (Figure 3D). We noticed that the rhabdoms of these ommatidia are thicker in diameter and more intensely immunostained than the rhabdoms of ommatidia in the main retina adjacent to the DRA (data not shown). This reflects UV opsin staining throughout the entire rhabdom because of UV opsin expression from all R1–R8 photoreceptor cells, compared to ommatidia in the rest of the retina in which the UV opsin is either not expressed (no immunofluorescence) or is expressed in only R1 and/or R2 photoreceptor cells (limited rhabdom staining). These variations in UV opsin staining in rhabdoms in main retina mirror our finding of UV opsin mRNA expression in ommatidia in main retina (Figure 2A). Because the rhabdoms of butterfly eyes are fused, it is difficult to distinguish by visual inspection of the pattern of rhabdom immunostaining those ommatidia with one versus two UV opsin-expressing cells.

Our assessment of opsin expression in the anatomically specialized DRA of monarch butterflies, which has previously been shown to contain orthogonally oriented microvilli (Reppert et al., 2004), indicates that all eight photoreceptor cells per ommatidium express only the UV opsin. Monochromatic receptors are required for efficient perception of polarized light in order to avoid interference with color information (Labhart and Meyer, 2002). DRA opsin expression has not been reported in any other butterfly.

Polarized UV Light and Monarch Butterfly Orientation
We then examined the importance of polarized light in the UV range for monarch butterfly orientation by placing butterflies in a flight simulator outdoors and using a broad-band linear polarizing filter (HNP'B, transmission 275–750 nm), as previously described (Reppert et al., 2004). The butterflies were positioned below the polarizing filter so that the angle of skylight vision through the filter was restricted to 80° and the sun was not directly visible. We then used a transparent filter that blocks >95% of light transmission below 397 nm (UV-interference filter) to cover the top of the simulator barrel. As our previous studies have shown that rotation of the polarizing filter by 90° causes a corresponding change in flight direction (Reppert et al., 2004), butterflies were challenged with a 90° rotation of the polarizing filter, either with or without the UV-interference filter in place. If polarized UV light is important for monarch butterfly orientation, then UV interference should dampen or substantially alter the orientation response to the 90° rotation in the polarizing filter.

Four butterflies were examined at consecutive 5 min intervals of flight in the simulator. Each 5 min orientation pattern was plotted as a circular histogram, and virtual flight paths were constructed. Four sequential flight records were examined (see Figures 4A–4D): The first record recorded flight orientation before rotation of the polarizing filter, while the second recorded flight after a 90° rotation of the polarizing filter to obtain a “baseline” response to a 90° change in the polarization axis. Between the second and third records, the UV-interference filter was positioned above the butterflies prior to a second 90° rotation in the polarizing filter, and flight orientation was recorded for the third record (colored red). The fourth record recorded flight direction after removal of the UV-interference filter, but with no further rotation of the polarizing filter.
flight orientation without the UV-interference filter (virtual flight path lengths in the records above and below the red records). Video monitoring showed that this decrease was due to near-continuous circling behavior by the butterflies that was a mixture of flight and struggling without flight (Movie S1). This behavior produced the increase in nondirectional baseline activity seen in the circular histograms (Figures 4A–4C, red records) and contrasts with the continuous flight and intermittent circling observed when the UV-interference filter was not in place. It is noteworthy that the butterfly in Figure 4C manifested clear, within-record bimodal flight patterns, but with a dominant unimodal direction, without UV interference (Figure 4C, top, second, and fourth records), which is a behavioral response to a linear polarization axis (von Frisch, 1967; Waterman, 1981; Wehner, 2001). The fourth butterfly also manifested an altered orientation pattern in the face of UV interference (see red flight path of Figure 4D). This individual initially maintained the same flight direction as that before UV interference, but by the end of the UV-interference flight period, the butterfly had altered its orientation substantially. For all four butterflies, removal of the UV-interference filter, with no further change in the plane of the polarizing filter, resulted in sustained directional flight that shifted in orientation relative to the direction before UV interference (compare change in flight paths between the second and the lower records of Figures 4A–4D), consistent with the orientation position of the polarizing filter (mean response of 109° ± 7.4°, n = 4). Examination of the orientation positions for three of these butterflies when exposed to the UV-interference filter without altering the position of the polarizing filter indicated that exposure to the UV-interference filter itself does not cause large changes in orientation position (Figure S3).

Our behavioral data show that UV interference severely blunts or alters orientation responses to a change in the plane of skylight polarization. Taken together with the anatomical data, we propose that polarized light in the UV range, sensed through specialized DRA photoreceptors, is important for flight orientation in monarch butterflies. UV polarized light has been shown to be the best celestial cue for diurnal animals under partly cloudy conditions (while blue receptors are best at dawn and dusk) (Barta and Horvath 2004), which is consistent with monarchs’ completely diurnal flight behavior and their tendency to cluster at dusk (Brower, 1996).

PER Staining Reveals Location of a Circadian Clock in Monarch Brain
How is information from DRA-derived polarized light input influenced by a circadian clock to produce time-compensated sun compass orientation? To begin to address this question, we sought to identify the location of a circadian clock in monarch brain that might be involved in navigational activities. This was approached by using PER immunocytochemistry to identify putative circadian clock cells. Two anti-PER antibodies (57/10w and 58/10w; Sauman and Reppert, 1996) were examined that gave identical staining patterns in monarch butterfly brain. The 58/10w antibody was used because of decreased background staining. There was consis-

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Figure 4. The Importance of Polarized UV Light for Flight Orientation
(A–D) Sequential records of flight orientation are shown as circular histograms in a column for each butterfly. All the butterflies started flight in the simulator with the polarizing filter plane placed parallel to the dominant skylight polarization axis viewed at the zenith. The polarization filter was rotated clockwise by 90° between each record of the top three records (curved arrows). Flight records in the presence of a UV-interference filter are colored red. The UV-interference filter was removed between the third and fourth record of each butterfly. Each circular histogram is a flight record of 5–7 min with sampling at 200 ms intervals. For each circular histogram, a calculated virtual flight path is depicted in the square to the right. We constructed virtual flight paths by starting in the center of the square and plotting each direction interval consecutively as one unit length (Mouritsen and Frost, 2002). 0° is North.

The baseline change in orientation induced by a 90° rotation of the polarizing filter was 77.3° ± 3.8° (mean ± SEM, n = 4) (Figures 4A–4D, two upper records). When the polarizing filter was rotated 90° again while UV-interference filter was applied, the orientation response was substantially altered (Figures 4A–4D, red highlighted records). Three of the butterflies exposed to the UV-interference filter showed at least a 70% decrease in directional flight (decreased length of the virtual flight paths in Figures 4A–4C, red flight paths) compared to
tent PER staining in a group of 8 to 12 large neurosecretory cells in the pars intercerebralis (PI) that was reduced during the day (compare Figures 5B and 5C) and in constant light (compare Figures 5H and 5I). These cells had stained axonal projections that projected laterally and ventrally (Figures 5A and 5H). There was also PER staining in two cells in the dorsolateral protocerebrum (pars lateralis, PL) of each hemisphere whose appearance was dependent on the phase of the 12 hr light:12 hr dark (LD) cycle examined; the stained cells were apparent at night (e.g., zeitgeber time [ZT] 18; Figure 5E), but were not detectable during the day (e.g., ZT 6; Figure 5D). Double-labeling experiments with both the anti-PER antibody and a monarch per cRNA probe demonstrated the colocalization of per transcript and protein in both the PI and PL (data not shown). There was also PER staining in a group of small cells at the base of the optic lobe (Figure 5F) and in cells in the corpora cardiaca (Figure 5G); in the optic lobe the staining was variable, independent of the lighting conditions. In all cases, cellular staining was cytoplasmic. There was no detectable PER staining in the compound eye, in contrast to the situation in both Drosophila and A. peryyi (Stanewsky, 2002; Sauman and Reppert, 1996). The conspicuous lack of detectable retinal PER staining argues against the existence of circadian clocks in monarch photoreceptor cells. The same PER staining pattern in brain was seen in both summer and migratory butterflies.

Because of the variation in PER staining in the PI and PL over the LD cycle, we quantified PER staining in those regions at 3 hr intervals over a 24 hr period in LD, constant darkness (DD), and constant light (LL). The daily oscillation in PER staining in the PI was broad in LD and did not persist in DD. In LL, there was an overall decrease in PER staining and no clear daily oscillation. In the PL, on the other hand, PER staining exhibited a robust 24 hr rhythm in LD, with peak staining at ZT 15. The rhythm in PER staining persisted in DD with a phase similar to that in LD, but the staining was eliminated at most times in LL.

The robust diurnal and circadian oscillation of PER staining in PL strongly suggests that a major circadian clock resides in these cells. We thus examined the expression of TIM staining in monarch brain by immunocytochemistry, as TIM should be coexpressed with PER in clock cells, on the basis of studies of Drosophila and A. peryyi. TIM-like immunoreactivity was detected in the cytoplasm of cells in the PI and in two cells in the PL of monarch brain, similar to the location of PER-positive cells (Figure S4). The modest intensity of TIM-like immunoreactivity in monarch brain together with high background levels, however, precluded definitive colocalization studies of PER/TIM staining and any assessment of oscillation in TIM staining in monarch brain.

CRY Staining in Monarch Brain
In Drosophila brain, all CRY-expressing cells also contain PER (Klarsfeld et al., 2004). In addition, CRY is a critical circadian photoreceptor in the fly (Emery et al., 2000). We therefore reasoned that PER and CRY should be colocalized in putative clock cells in monarch brain. We cloned the cDNA encoding monarch CRY (GenBank
There was also CRY staining in two cells in the PL of the posterior dorsal region of the medulla (section for both PER and TIM immunoreactivity in PI. Staining in PI appeared to be in the same location as cells in LD, DD, and LL; data not shown). The CRY staining did not vary with the lighting conditions (examinations for both PER-positive and TIM-immunoreactive cells in PI and the protocerebral bridge laterally toward the PI and the protocerebral bridge (Figure 5A). CRY immunoreactivity in two cells in the dorsolateral protocerebrum (PL). Double-labeling immunofluorescence of CRY (D) and PER (E) in the same cell (arrow) in PL. Brain section in (H) stained for the presence of cry mRNA demonstrates the colocalization of cry transcript and protein in the same cells. Note the absence of cry mRNA signal in the axonal projections forming the glomerular-like structure (arrow).

Figure 6. CRY Immunoreactive Cells and their Axonal Projections in Monarch Brain
(A) Schematic representation of CRY-positive neurons and their axons. (B) CRY staining in a restricted group of neurosecretory cells in pars intercerebralis (PI). (C) CRY immunoreactivity in two cells in dorsolateral protocerebrum (PL). (D and E) Double-labeling immunofluorescence of CRY (D) and PER (E) in the same cell (arrow) in PL. (F and G) Double-labeling immunofluorescence of CRY (F) and COR (G) in the two cells in PL. (H) A prominent group of small CRY-positive neurons in the dorsal region of the optic lobe (OL), forming a glomerular-like body of axonal projections (arrow). (I) Brain section in (H) stained for the presence of cry mRNA demonstrates the colocalization of cry transcript and protein in the same cells. Note the absence of cry mRNA signal in the axonal projections forming the glomerular-like structure (arrow).

AAC83828: Figure S5) so that we could generate monarch-specific anti-CRY antibodies (Figure S6). Then, using immunocytochemistry we examined the anatomical distribution of monarch CRY expression and its relationship to PER-positive cells.

There was consistent CRY staining in a group of large neurosecretory cells in PI (Figure 6B); the intensity of staining did not vary with the lighting conditions (examined in LD, DD, and LL; data not shown). The CRY staining in PI appeared to be in the same location as cells staining for both PER and TIM immunoreactivity in PI. There was also CRY staining in two cells in the PL of each hemisphere, the appearance of which was also independent of the lighting conditions (Figure 6C; data not shown). Moreover, double-labeling experiments with both the anti-CRY antibody and a monarch cry cRNA probe demonstrated the colocalization of the cry transcript and protein in the same cells in PL (data not shown). Importantly, double-labeling immunofluorescence revealed the colocalization of CRY and PER staining in the two cells in each PL (Figures 6D and 6E), as would be expected if these cells are indeed circadian clock cells in monarch butterfly brain.

The neuropeptide corazonin (COR), which appears to be involved in the initiation of ecdysis behavior (Kim et al., 2004), is co-expressed in the eight PER-positive cells in silkmoth brain previously proposed to function as circadian clock cells (Sauman and Reppert, 1996; Sauman and Hashimi, 1999; Qi-Miao et al., 2003; I. Sauman, unpublished data). PER-positive cells in the PL of the hawkmoth Manduca sexta that are homologous to the PER-positive cells in the silkmoth PL have been mapped as COR-expressing type-Ia, neurosecretory cells (Wise et al., 2002). Thus, COR is a marker of lepidopteran la, neurosecretory cells that should label homologous neurons in monarch brain. Indeed, COR-immunoreactivity was expressed in four cells in each PL of butterfly brain (data not shown) and was colocalized in the two CRY/PER-positive cells in each PL (Figures 6F and 6G). This finding provides further evidence that these CRY/PER-positive cells in monarch brain are bona fide circadian clock neurons.

In addition to those in the PI and PL, there was a prominent group of small CRY-positive cells in the dorsal region of the optic lobe that formed a glomerular-like body with axonal projections (Figures 6H and 6I). No PER staining was found in this region (see Figure 5). Double-labeling experiments with both the anti-CRY antibody and the monarch cry cRNA probe demonstrated the colocalization of cry transcript and protein in the same cells in dorsal optic lobe (Figures 6H and 6I). As for PER, there was no detectable CRY staining in the monarch eye. Similar CRY staining patterns were seen in both summer and migratory butterflies.

CRY Staining Reveals a Fiber Pathway from the Circadian Clock to the Optic Lobe
There were substantial CRY-immunoreactive fibers in the monarch brain (Figure 7) and, as discussed below, a CRY-containing pathway was delineated that could provide a vital link between the circadian clock in PL and axons originating from polarization-sensitive photoreceptors in the DRA. We identified CRY-positive fibers in the dorsolateral protocerebrum that were in direct contact with the two CRY-positive cells in PL (Figures 7A and 7B). This fiber pathway projected medially toward the PI and the protocerebral bridge and laterally toward the CRY-positive group of cells in the optic lobe. Fine projections from this optic lobe group of CRY-positive cells formed a glomerular-like arborization laterally from the cell bodies (Figures 6H and 7C). Additional CRY-positive fibers emanated from this glomerular structure and projected dorsolaterally along the surface of the medulla (Figure 7C). Finally, the fibers entered a posterior dorsal region of the medulla (sec-
Figure 7. CRY Immunoreactive Fibers in Monarch Brain
(A) Schematic diagram depicting the topography of CRY immunoreactivity in the monarch brain.
(B) Two CRY immunofluorescent cells and CRY-positive fibers in dorsolateral protocerebrum (PL).
(C) A glomerular-like arborization of CRY-positive cells in the optic lobe (OL) and axons projecting toward the dorsolateral medulla.
(D) A fine arborization of CRY immunofluorescent fibers in the dorsolateral medulla (ME).

Because the CRY-immunofluorescent fibers in the dorsolateral medulla are in the same location where axons from dorsal rim UV-expressing photoreceptors would be expected to project, on the basis of studies in flies (Hardie, 1984; Fortini and Rubin, 1991), we examined the relationship between the CRY-positive fibers and photoreceptor axons. This was evaluated by micro-injecting monarch DRA photoreceptor cells with fluorescent neuronal tracing dye.

We were able to show that the axons of these dorsal rim photoreceptors terminated in the posterior dorsolateral medulla of both summer and migratory butterflies. Although this was a consistent finding (in >40 injected animals), it was not seen following similar injection in the photoreceptor cells of more ventral or lateral ommatidia, which mostly terminated in the lamina (first-order neuropil). The injected dorsal rim photoreceptors (Figures 8A and 8B) projected proximally through the dorsal region of the lamina. The dye-labeled projections then passed through a cell body layer (rind) between the first- and second-order neuropils relatively close to the dorsal surface of the optic lobe. They entered the medulla via its distal surface and finally terminated in a single stratum of the posterior dorsolateral medulla (Figures 8C and 8D). This represents the identical region of medulla in which the CRY-immunofluorescent fibers from the optic lobe CRY-positive cells terminated (Figures 7C and 8E). Due to an incompatibility of tissue processing for the two methods (dye injections and CRY immunofluorescence), we were not able to directly assess the likely colocalization of the CRY and photoreceptor cell terminals on the same brain section.

The anatomical data provide compelling evidence of a CRY-containing pathway that could communicate circadian information from the circadian clock neurons in PL to axons arising from polarization photoreceptors in the dorsal rim region of monarch eye. This pathway has not been described in any other insect, and it may be a hallmark feature of butterflies that use a time-compensated sun compass.

Conclusions
The collective results provide strong evidence that a circadian clock important for monarch butterfly naviga-
tion resides in four cells in the dorsolateral region (PL) of monarch brain. The PER-positive cells in PI also appear to express TIM and CRY. It is thus possible that these cells contribute to the overall clock-regulated functions in monarch butterfly brain. In fact, studies in Drosophila have recently shown that the circadian network that controls locomotor behavior involves the activities of at least two distinct groups of circadian neurons (Stoleru et al., 2004; Grima et al., 2004). The lack of nuclear staining for PER and TIM seen in monarch brain has been found in most of the nondipteran insects that have been examined (Sauman and Reppert, 1996; Zavod ska et al., 2003), emphasizing the need for more extensive analysis (Rosato and Kyriacou, 2001).

Our most provocative finding is the existence of a CRY-positive fiber pathway from the clock neurons in PL to the optic medulla in the same location where the axons from dorsal rim photoreceptors terminate. This pathway may provide a critical link between the circadian clock and sun compass input into brain. Electrophysiological studies of polarization-sensitive retinulae in dorsal rim ommatidia in eye and of polarization-sensitive neurons in the optic lobe in brain should clarify whether the CRY-containing pathway provides the relevant time-compensated information (Labhart and Meyer, 2002). It is also possible that there are multiple pathways that connect the circadian clock to different aspects of the sun compass system, including the sun compass itself (which may reside in the central complex; see Labhart and Meyer, 2002) and/or at the interface between the sun compass and the motor system.

The CRY-positive fiber pathway between PI and PL may also have an important function. The decreasing length of the day in the fall appears to be an important determinant for stimulating migratory behavior in butterflies (Goehring and Oberhauser, 2002). A defining characteristic of the migratory state is adult reproductive diapause (Brower, 1996), which contributes to an increased longevity necessary for survival at the overwintering grounds. Adult diapause is secondary to a deficiency in juvenile hormone (see Herman and Tatar, 2001), the synthesis of which is probably controlled by insulin-like peptides originating from neurosecretory cells in the PI. Therefore, a CRY-containing fiber pathway from the clock cells in the PL to the PI may play a vital role in providing photoperiodic information to the neuroendocrine system and, as mentioned above, involve secondary clocks in the PI itself.

Experimental Procedures

Animals

“Summer” butterflies were reproductively active and were purchased from commercial sources throughout the year. To confirm the clock protein staining patterns in brain in migratory butterflies, animals were captured near Eagle Pass, Texas, from October 10–12, 2004 and shipped to Worcester. The migratory females were dissected under 10× magnification to determine reproductive status, and all were found to be in reproductive diapause (indicated by the absence of mature oocytes). All butterflies were housed in the laboratory with controlled temperature (21°C for summer butterflies and 18°C for migrants), humidity (70%), and lighting. The butterflies were fed 10%–15% sucrose.

Cloning and Sequence Analysis

cDNA fragments were cloned by degenerate polymerase chain reaction (PCR). cDNA templates for PCR were prepared from RNA purified from monarch butterfly brains or whole heads. The ends of the coding regions were obtained by rapid amplification of cDNA ends (RACE; Clontech). Complete open reading frames were obtained by Phusion (Stratagene) PCR from cDNA. Clones were sequenced at core facilities at the University of Massachusetts Medical School. Sequence analysis was facilitated by software from the Genetics Computing Group (GCG) and the National Center for Biotechnology Information website (http://www.ncbi.nlm.nih.gov/BLAST/).

Phylogenetic Analysis

To explore the possibility that we had recovered members of all paralogous opsin genes from Danaus plexippus, we searched for the only currently available lepidopteran genome resource, the Bombyx mori 3X (Mita et al., 2004) and 6X (Xia et al., 2004) assemblies in GenBank using the BLASTx search tool and whole genome sequence (wgs) setting. Full-length opsin cDNAs were downloaded, and the translated amino acids were aligned, using the ClustalW algorithm (Figure S1). Only first and second nucleotide positions were used in the tree reconstruction, as a test for composition homogeneity amongst lineages (Kumar and Gadagkar, 2001) was rejected for third positions (data not shown). The neighbor-joining algorithm with Tamura-Nei distance estimates with complete deletion of gaps was implemented in MEGA 3.0 (Kumar et al., 2004). Bootstrap analysis was performed using 500 replications.

Antibody Production

For generating monarch CRY antibodies, purified protein containing the C-terminal 113 amino acids of monarch CRY was used as an immunogen in rats and guinea pigs, as previously described (Lee et al., 2001). Antisera were affinity purified and designated DpCRY1-R and DpCRY1-GP (“R”, raised in rats; “GP”, raised in guinea pigs). Both anti-CRY antibodies gave identical results when used for either immunocytochemistry or for probing Western blots (Figure S6). Western blotting was performed as described previously (Lee et al., 2001). The V-5 antibody used for Western blots was a monoclonal mouse anti-V5 IgG purchased from Invitrogen.

Immunocytochemistry

Eye tissue was fixed as described in Briscoe et al. (2003), and cryostat-sectioned (14–16 μm slices). Prior to antibody incubation, the sections were washed in 100% cold acetone for 5 min, then washed for 30 min in 1x PBS, and finally washed for 5 min in 1x PBS/0.5% SDS. Dissected brain-suboesophageal ganglion complexes of adult monarchs were immediately processed for immunocytochemistry as described elsewhere (Sauman and Reppert, 1996). Stained sections were examined with the aid of the Zeiss Axioplan 2 or an Axioskop 2 plus microscope equipped with Nomarski (DIC) optics, epifluorescence, and a CCD camera.

The antibodies used for immunocytochemistry included: rabbit anti-PrPRH5 (UV), (1:200; Briscoe and Nagy, 1999; Lampel et al., 2005); rabbit anti-A. peryn PER (57/10w at 1:500 and 58/10w at 1:500; Sauman and Reppert, 1996); rabbit anti-CORAZONIN (from Makio Takeda; dilution, 1:1000), rat-TIM (307; from Michael W. Young; dilution, 1:1000) and anti-DpCRY1-R and anti-DpCRY1-GP. A Cy3-conjugated goat anti-rabbit secondary antibody (1:1000; Jackson Immuno Research Laboratories, Inc., West Corone, PA) incubated for 1–2 hr at room temperature was used to detect the anti-UV primary antibody.

To verify the specificity of immunological reactions, primary antibodies were replaced with normal goat serum. In additional controls of binding specificity, the anti-PER antibody and the anti-CORAZONIN antibody were preincubated with 100 molar excess of the respective original antigen prior to immunocytochemical staining. In all cases, no significant staining above background was observed.

In Situ Hybridization

Fragments of monarch DpUVRh, DpBlueRh, DpLWRh, and per and cry cDNAs were subcloned into pCR II vector (Invitrogen) and used as templates for antisense and sense (control) digoxigenin (DIG)-labeled riboprobe synthesis. The riboprobe yield was assessed by dot blots.

For the opsin in situ hybridizations, the fixation and detection
methods are as described in Briscoe et al. (2003). For per and cry, the methods were similar except that after fixation in paraformaldehyde, the tissue was embedded in paraplast and sectioned (10 μm). In situ hybridization was carried out using the mRNA locator kit (Ambion). The riboprobes were localized by incubation with an alkaline phosphatase-conjugated anti-digoxigenin antibody (Boehringer and Mannheim; 1:500 dilution overnight at 4°C) and visualized with BCIP and NBT (Perkin Elmer). DIG-labeled sense RNA probes were used in control experiments. In all cases, no signal was detected above background.

For double-labeling experiments, hybridized sections were incubated simultaneously with anti-DIG antibody and anti-PER or anti-CRY overnight at 4°C and detected with a fluorophore-conjugated secondary antibody (1 hr at RT). The AP activity was detected as described above.

For scoring of immunoreactive intensities, stained sections were coded and viewed under a microscope. Levels of staining were subjectively scored with an intensity scale ranging from 0 to 5 per cell. The time of collection was decoded after scoring.

**Microinjections**
Alexa Fluor-conjugated dextran neuronal tracer (10,000 MW, 10% in sterile water; Molecular Probes) was injected into photoreceptor and fluorescent traces. Injected animals were kept in constant darkness for 4–8 h before being killed.

**Behavioral Experiments**
Monarch butterflies migrating through Massachusetts were captured in the wild in the fall of 2003 and 2004. The migrants were housed outside in plastic mesh cages with free access to 10% sucrose in water. Butterflies were tethered as previously described (Mouritsen and Frost, 2002), and flight behavior was monitored using a modified Mouritsen and Frost flight simulator as described (Reppert et al., 2004). The four individuals studied were part of a group of 13 butterflies that initially flew under the parallel-placed polarizer in the southwesterly (migratory) direction, when corrected for bimodal orientation (α = 19°/199°; n = 13, r = 0.49, p < 0.05).

The linear polarizing filter was rotated by hand. The UV-interference filter was a long-wavelength pass filter with a photopic luminous transmission of 84% (E420 from Gentex, Carbondale, PA). Transmission values were as follows: 5% ≥ 397 nm; 50% ≥ 420 nm; 80% ≤ 436 nm. Data in each orientation histogram were analyzed to determine the significance of orientation and the mean direction using circular statistics (Batschelet, 1981).

**Supplemental Data**
Supplemental data include six figures and a movie and can be found with this article online at [http://www.neuron.org/cgi/content/full/46/3/457/DC1/](http://www.neuron.org/cgi/content/full/46/3/457/DC1/).

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