Coupling optogenetics and light-sheet microscopy, a method to study Wnt signaling during embryogenesis

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Optogenetics allows precise, fast and reversible intervention in biological processes. Light-sheet microscopy allows observation of the full course of Drosophila embryonic development from egg to larva. Bringing the two approaches together allows unparalleled precision into the temporal regulation of signaling pathways and cellular processes in vivo. To develop this method, we investigated the regulation of canonical Wnt signaling during anterior-posterior patterning of the Drosophila embryonic epidermis. Cryptochrome 2 (CRY2) from Arabidopsis Thaliana was fused to mCherry fluorescent protein and Drosophila β–catenin to form an easy to visualize optogenetic switch. Blue light illumination caused oligomerization of the fusion protein and inhibited downstream Wnt signaling in vitro and in vivo. Temporal inactivation of β–catenin confirmed that Wnt signaling is required not only for Drosophila pattern formation, but also for maintenance later in development. We anticipate that this method will be easily extendable to other developmental signaling pathways and many other experimental systems.

Dissections of signaling pathways and their downstream processes have centered on in vitro cell culture models and in vivo model organism studies using small molecule, knockdown and overexpression approaches. Although these approaches have been very useful, they do not lend themselves easily to the study of the spatial and temporal regulation of cell signaling pathways and cellular processes in the dynamic environment of a living organism. Normal genetic crosses using mutations and RNAi techniques are limited in their applications as they result in irreversible knockout or knockdown of a gene, which can be lethal or incomplete1. The recent discovery of optogenetic tools has provided a new and effective toolbox for spatial and temporal regulation of proteins in various in vitro and in vivo systems2–4. This is made possible by the rapid response of these proteins to light, such as the blue-light mediated activation of Arabidopsis thaliana cryptochrome 2 (CRY2)6. We chose to use the CRY2 system as it has proven effective in Drosophila embryos7.

The CRY2 system can be activated simply with 488 nm lasers, but this can result in high phototoxicity and bleaching which prevent collection of temporal developmental information. Hence, the application of optogenetic tools is only as good as the visualization approach that can be brought to bear. The latest advances in light-sheet microscopy facilitate high resolution optical sectioning and live imaging of embryos with low phototoxicity as compared to conventional laser confocal imaging8. Importantly, light-sheet microscopy enables selective activation of optogenetic insertions without significant out-of-plane effects. This contrasts with confocal activation, where optogenetic components above and below the focal plane (but inside the focal light cone) are activated as the system is typically highly light sensitive. Light-sheet illumination does result in some light scattering, but this effect is smaller than the light cone effect in confocal imaging. To reduce these effects, two-photon approaches have been developed9. In addition, new, stable fluorescent proteins allow imaging of various cellular processes at the same time10.

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Figure 1. Light-induced clustering of Arm-CRY2-mCh modulates the Wnt/β-catenin pathway in Drosophila. (A) Arm-CRY2-mCh puncta formation in S2R+ cells in response to 488 nm laser light. Scale bar represents 5 μm. (B) Quantification of puncta in Arm-CRY2-mCh expressing S2R+ cells after illumination with 488 nm laser light. One-way repeated measures ANOVA showed statistically significant difference in number of puncta between the different time points, ##p < 0.01. (C) S2R+ cells transfected with pAct-Arm-CRY2-mCh plasmid and the TOPFlash luciferase reporter showed reduced TOPFlash activity after 16hr exposure to light. Graph is representative of three independent experiments and the average of three replicates (mean ± SD). The variance was found to be equal as determined by the F-test (p > 0.05). Statistical significance relative to control samples was determined using the Student’s t-test. **p < 0.01 relative to control, ##p < 0.01. (D) Arm-CRY2-mCh puncta formation in Drosophila embryos overexpressing Arm-CRY2-mCh in a wild-type background upon exposure to 488 nm laser light at 2.5 min intervals. Puncta formation observed during illumination was reversible and oligomerization could be induced repeatedly. Scale bar represents 20 μm. (E) Quantification of mean intensity (used as an indicator of puncta formation) in Drosophila embryos overexpressing Arm-CRY2-mCh in a wild-
The Wnt pathway is a highly conserved signal transduction pathway functioning in development and disease in many organisms. In *Drosophila*, Wnt1 (Wingless or Wg) is a segment polarity gene that defines anterior-posterior patterning of the epidermis. Canonical Wnt signaling leads to inhibition of β-catenin (Armadillo or Arm in *Drosophila*) degradation, which subsequently activates downstream Wnt target genes in the nucleus. As β-catenin protein levels increase, it enters the nucleus where it regulates transcription of target genes by associating with the transcription factor TCF. This basic signaling mechanism is widely conserved, and critical in embryogenesis, but whether β-catenin signaling is needed in a constant widespread manner, versus in a temporally and/or spatially restricted manner was not known, because the tools needed to address this question were lacking. Here we describe our method for applying CRY2 protein fusions to investigate the spatial and temporal regulation of β-catenin protein in the developing embryo.

**Results**

**Light-induced oligomerisation of β-catenin-CRY2-mCh modulates Wnt signaling in vitro.** Blue light can induce oligomerization of *A. thaliana* CRY2 to form ‘photobodies’ in plant and mammalian cells. This leads to sequestration and hence functional inactivation of the protein. We tested this mechanism by fusing the photolyase homology region (PHR) of CRY2 and mCherry to the C terminus of *Drosophila* β-catenin or Arm protein (Arm-CRY2-mCh). To determine if the Arm-CRY2-mCh construct was responsive to light, we transfected it into S2R + Wnt responsive *Drosophila* cells. Distinct fluorescence puncta were observed with blue light illumination (Fig. 1A; full construct details in Supplementary Fig. 1). The puncta dispersed, but did not entirely disappear upon withdrawal of blue light (Fig. 1A,B).

We next investigated if inducible Arm-CRY2-mCh puncta formation functionally inactivated Arm activity. The effect of the Arm-CRY2-mCh construct on canonical Wnt signaling was determined by using a TOPFlash reporter assay, where the luciferase gene is attached downstream of multimerized TCF binding sites. Overexpression of Arm-CRY2-mCh in S2R + cells in the absence of light significantly increased TOPFlash activity relative to control transfected cultures (Fig. 1C). This activation was inhibited when transfected cells were illuminated with light. Hence, oligomerization of the Arm-CRY2-mCh construct upon illumination with light effectively blocked downstream Wnt signaling.

Having verified the responsiveness of the Arm-CRY2-mCh construct to blue light illumination and its ability to modulate Wnt signaling in vitro, we tested the oligomerization of this construct in *Drosophila* embryos. Expression in wild type embryos showed the cytoplasmic and diffuse localization of Arm as indicated by the mCh fluorescence (Fig. 1D, Supplementary video 1). Since CRY2 has a turn-off speed of 5 minutes, the embryo was illuminated with blue light at 2.5 minute intervals at the lowest effective laser power (6%) and exposure time (7.5ms), to ensure continuous clustering while at the same time preventing excessive heating of the embryo. Upon blue light illumination, distinct fluorescent puncta were observed (Fig. 1D, Supplementary video 1) which plateaued 15 minutes after exposure (Fig. 1E). These results show an in vivo, light-responsive aggregation of β-catenin during development of the *Drosophila* embryo.

**Arm-CRY2-mCh can rescue patterning in arm*XM19* mutant embryos.** In order to establish the ability of the Arm-CRY2-mCh construct to function in place of endogenous Arm, we expressed it in an arm*XM19* mutant background. We used arm*XM19* mutants, because these lack the C terminus required for transcriptional transactivation but do not lose adhesion (Fig. 2A). The basic test was to replace the mutated arm*XM19* with Arm-CRY2-mCh (arm*XM19*/FM6; armGal4 crossed to UAS-Arm-CRY2-mCh). As the arm gene is located on the X-chromosome, normal, non-balancer, males from this cross can only be the rescued progeny. We observed normal males eclosing (three separate experiments, 31 of 64 males) from vials kept entirely in the dark and no normal males appearing from vials left out in normal room light. This genetic rescue experiment showed that Arm-CRY2-mCh could rescue arm mutants to viability as long as the flies were not exposed to light.

When canonical Wnt signaling is turned off, all ventral epidermal cells produce denticles. The opposite is true when Wnt signaling is turned on ecptocally; most cells do not make denticles causing the naked phenotype (For a review of embryonic patterning). We made germline clones of *arm*1004 where the maternal and zygotic contribution of protein was removed, resulting in an embryonic phenotype identical to wg mutant embryos (Fig. 2B). In these embryos, we expressed Arm-CRY2-mCh and saw a rescue of the canonical signaling defect (Fig. 2B) where arm*XM19* mutants with Arm-CRY2-mCh showed a nearly wild type denticle pattern as long as the embryos were kept in the dark (Fig. 2B). When exposed to light, we observed no rescue of the canonical signaling defect as embryos showed the null signaling phenotype (Fig. 2B). Taken together, these results suggest that the Arm-CRY2-mCh construct can effectively rescue canonical Wnt signaling in *arm*1004 mutant embryos, and that this rescue can be blocked by exposure to light.

To confirm the Wnt dependence of the phenotype, we examined genes downstream of Wnt signaling using quantitative PCR. Expression of the embryonic Wnt target genes *en* and *wg* was significantly decreased in Arm-CRY2-mCh expressing arm*XM19* mutants exposed to light as contrasted with embryos kept in the dark both quantitatively (Fig. 2C), and qualitatively through immunostaining for En protein (Fig. 2D). In the absence of light, stripes of En protein were observed but these were disrupted with blue light illumination (Fig. 2D).

Next, we used light-sheet microscopy to live-image these embryos. arm*XM19* mutants expressing the Arm-CRY2-mCh construct (as observed by luminescence upon excitation with the 561 nm laser), showed
Figure 2. Arm-CRY2-mCh rescue of *arm<sup>xM19</sup>* mutants is reversible with blue light. (A) Structure of *arm<sup>xM19</sup>* truncated protein (adapted from<sup>49</sup>). The N-terminus of Arm is required for transactivation, phosphorylation-based and proteasome-mediated degradation, and α-catenin binding. The central repeats contain the binding sites for most of Arm's binding partners, such as APC, TCF, Cadherin, and Axin. The C-terminus is required for Cby and Teashirt binding and transactivation. *arm<sup>xM19</sup>* lacks the entire C-terminus. (B) Cuticle preparations of *arm<sup>xM19</sup>* mutant (germline clone, GLC) embryo with entire cuticle covered with denticles, *arm<sup>xM19</sup>* GLC expressing Arm-CRY2-mCh without light was rescued to a wild-type cuticle (male embryos were identified by the presence of the yellow mutation and sorted away from the female embryos which were y<sup>+</sup>). *arm<sup>xM19</sup>* GLC expressing Arm-CRY2-mCh with illumination showed defective Wg signaling (Dentine covered cuticle). Scale bar represents 50 μm. (C) Quantitative expression of genes downstream of Wg signaling in Arm-CRY2-mCh overexpressing *arm<sup>xM19</sup>* mutants. Pooled embryos exposed to light showed reduced *En* and *Wg* gene expression. Graph is representative of three independent experiments and the average of three replicates (mean ± SD). The variance was found to be equal as determined by the F-test (p > 0.05). Statistical significance relative to control samples was determined using the Student's t-test. *p < 0.05, **p < 0.01. (D) En antibody staining in Arm-CRY2-mCh overexpressing *arm<sup>xM19</sup>* mutant embryos illuminated or not illuminated with light. En stripes as in wild-type embryos were observed in embryos not illuminated with light whereas embryos illuminated with light showed only small patches of En expression. Scale bar represents 50 μm. (E) Stills from live imaging movies of Arm-CRY2-mCh overexpressing *arm<sup>xM19</sup>* mutants. Embryos not subjected to light showed diffused localization of Arm-CRY2-mCh (inset) as compared to embryos subjected to light (inset). Embryos not subjected to light showed normal development whereas embryos subjected to light showed various developmental defects. Scale bar represents 50 μm.
normal embryonic development (Fig. 2E, Supplementary video 2A) similar to embryos expressing Arm-GFP (Supplementary video 2C). To ensure that the embryos imaged were the correct mutant (armXM19 male embryos expressing the Arm-CRY2-mCh) we cuticle prepped the embryos post-imaging and scored for the yellow mutation. Only y- embryos expressing mCh are presented here. We next exposed these embryos to blue light and observed smaller embryos developing with a lawn of denticles (Fig. 2E, Supplementary video 2B), consistent with a loss of Wnt signaling. A troubling finding was that the puncta once formed did not disappear quickly raising the possibility that the effect was not reversible. To test this, we performed experiments where embryos were placed in the light sheet microscope and illuminated with blue light every 2.5, 5 or 10 minutes. armXM19 mutants expressing the Arm-CRY2-mCh construct illuminated every 2.5 or 5 minutes did not survive to the end of embryogenesis, but those illuminated only once every 10 minutes did not hatch. We propose that the persistence of puncta represents the cells’ inability to degrade aggregated Arm protein quickly, but this does not affect Wnt signaling as enough Arm protein is made during the dark phase to compensate for the aggregates as long as 10 minutes are allowed to elapse. The system, therefore, is reversible but may require new protein synthesis rather than dis-aggregation.

**Temporal regulation of Wnt signaling during development in armXM19 embryos.** Having established the efficacy of the Arm-CRY2-mCh transgene in embryos, we next performed a temporal experiment to answer the biological question of when Wnt signaling is required. Segment polarity genes are expressed downstream of pair rule genes early in development22. These genes turn on Wg expression24 through a regulatory loop with Engrailed25,26. Experiments with a temperature sensitive allele of wg established four phases of Wg signaling in establishment and maintenance of embryonic patterning27. These can be subdivided into early en activation and later shaven baby repression in combination with EGF signals25,26,29. We used eggs exposed to light at different developmental stages26, and observed the cuticle phenotypes. Staging was approximate as the experiment was carried out at 21°C. Exposure to light from stages 1 to 11 (From cleavage stages to Germ band elongation) showed a variable lawn of denticles similar to wg mutants (Fig. 3A). Strong Wnt pathway loss of function embryos show a lawn of denticles and are much smaller than wild-type. Restoring some Wnt pathway function rescues the small size phenotype12,31. The lawn of denticles began getting broader (size of the embryo increased) when light was applied at stage 12 (onset germ band retraction) with some distance developing between the denticle bands (Fig. 3A, stage 13 germ band retraction). This experiment showed that Arm-CRY2-mCh could be used to recapitulate previous findings with wg that showed effects into dorsal closure stages37.

**Spatial perturbation of Wnt signaling during development in armXM19 embryos.** We next wanted to test the spatial resolution of this approach using light sheet microscopy as a blue light source as well as a recorder for live imaging. Light sheet microscopy is not an ideal tool for this experiment as the light is not delivered as a point (For a better approach to spatial resolution see38). Half of an embryo overexpressing the Arm-CRY2-mCh transgene was illuminated with blue light and the whole embryo imaged in the red channel. The illuminated region showed the expected head involution and germ-band retraction defects with the unilluminated half remaining more wild-type (Fig. 3B, Supplementary Video 3A, compare to Supplementary Video 2A and 2B). Spatial manipulation of signaling was further tested by illuminating a portion of the embryo along the anterior-posterior axis. Illumination of the posterior region of the embryo showed distinct clustering and 2B). Spatial manipulation of signaling was further tested by illuminating a portion of the embryo along the anterior-posterior axis. Illumination of the posterior region of the embryo showed distinct clustering and

**Light-responsive properties of Arm-CRY2-mCh can modulate Wnt signaling in HEK293.** We have shown that Arm-CRY2-mCh functions in insect cells and in Drosophila embryos. We next tested if this construct could be applied to other experimental systems by looking at whether it could be used to modulate Wnt signaling in mammalian cells. We used HEK293 (Human embryonic kidney cells) stably transfected with the Wnt3a gene and TOPFlash vector (HEK293-Wnt3a-TOPFlash). Cells transfected with the construct showed fluorescent puncta formation upon blue light illumination (Fig. 4A). The puncta dispersed rapidly upon withdrawal of blue light (Fig. 4A).

The effect of inducible Arm-CRY2-mCh clustering on Wnt signaling in mammalian cells was quantified using a TOPFlash reporter assay. Overexpression of Arm-CRY2-mCh plasmid in HEK293-Wnt3a-TOPFlash cells in the absence of light significantly increased TOPFlash activity relative to control transfected cultures (Fig. 4B). This activation was inhibited when transfected cells were exposed to light. Our data suggest that oligomerization of the Arm-CRY2-mCh construct upon illumination with light could be applied to other experimental systems.

**Discussion**

Classical genetics tools allow tissue-specific manipulation of gene expression, but not precise temporal or spatial manipulation. Optogenetic tools can address this problem by allowing spatial and temporal regulation in a rapid, precise, and reversible manner23,34. Various optogenetic systems have been described to date such as the Cryptochrome 2 (CRY2), Phytochrome B (PHYB), LOV and Dronpa systems. However, only the CRY2 and LOV systems allow optogenetic control at a lower wavelength with live imaging at a higher wavelength. Both systems are activated within a few seconds, however, the CRY2 system has a slower turn off speed of approximately 5 minutes as compared to tens of seconds or minutes in the case of LOV17. This makes the CRY2 system ideal for live imaging with lower phototoxicity. Our study is the first to combine optogenetics with light-sheet microscopy for noninvasive, temporal and spatial control of cell signaling in vivo. Using these tools, we investigated the spatial and temporal regulation of β-catenin and Wnt signaling in vivo and in vitro proving the efficacy of this method. Wnt’s role in segment polarity is well known35,36, and we observed the same phenotypes by inactivating Wnt.
signaling during early stages of development (stage 1 to 11). Light-sheet microscopy is not designed for single cell conversion of optogenetic switches, but can be applied to sections of the embryo to study problems such as left-right asymmetry and anterior-posterior patterning by illuminating different regions of the embryo. Other microscopy approaches can be used to prevent light scattering, where the optical molecule could be activated in unintended parts of the embryo, but the advantage of light sheet microscopy is that it allows whole embryo imaging over the entire course of the 24 hour Drosophila embryogenesis.

Wnt signaling in segment polarity is well studied, but many additional functions have been observed. Recently, reported that intestinal formation during embryo development was disrupted in embryos treated with...
ionomycin, an inhibitor of Wnt/ß-catenin signaling. By coupling optogenetics with live imaging, we observed a similar defect in intestinal development in Arm-CRY2-mCh expressing armXM19 mutant embryos. Our experiment with spatial inactivation of Arm-CRY2-mCh in only a specified region of the embryo, also confirmed these findings. We see the application of optogenetics with light-sheet microscopy will allow the exploration of many more Wnt pathway effects. The main limitation appears to be the buildup of CRY2 puncta that do not disaggregate. In our experiments this leads to a buildup of mCh fluorescence in the later stages especially in hemocyte cells making imaging in the red channel difficult at late stages. We anticipate that this can be overcome by using separate fluorescent proteins for imaging and for optogenetics.

Our optogenetics approach is a non-invasive and versatile tool that allows for manipulation as well as live tracking of protein activity to interrogate complex phenotypes in vivo49. We foresee applying this tool to cell culture, organoid studies, vertebrate development in zebrafish, and many other systems40. It provides an alternative to small molecule studies which can only be performed on druggable targets. We have made a human ß-catenin-CRY2 version that functions similarly, and are applying the optogenetic methods to study adhesion by using embryos with a stronger loss of function armO43A01 allele that disrupts junctions but is rescued by Arm-CRY2-mCh41–43.

Figure 4. Light-induced clustering of Arm-CRY2-mCh modulates the Wnt/ß-catenin pathway in mammalian cells. (A) Arm-CRY2-mCh puncta formation in HEK293-Wnt3a-TOPFlash cells in response to 488 nm laser light, as well as dissociation after light withdrawal. Scale bar represents 10μm. (B) HEK 293-Wnt3a-TOPFlash cells transfected with pAct-Arm-CRY2-mCh plasmid and the TOPFlash luciferase reporter showed reduced TOPFlash activity after exposure to light. Graph is representative of three independent experiments and the average of three replicates (mean ± SD). The variance was found to be equal as determined by the F-test (p > 0.05). Statistical significance relative to control samples was determined using the Student’s t-test. **p < 0.01 relative to control, ***p < 0.01.
Materials and Methods

Transgenes and GAL4 driver lines. Two ubiquitous drivers were used for expression of transgenes: the weaker armadillo-GAL4 and the stronger daughterless-GAL4\(^{44}\). The Arm-CRY2-mCh constructs were made using the MultiSite Gateway\(^{3}\) Pro 2.0 cloning kit (Invitrogen). Full-length Armadillo was amplified from Drosophila embryo cDNA with attB1 and attB5r-flanked sites. This PCR product was recombined into pDONR P1-P5r vector. The CRY2-mCh was amplified from pCRY2PHR-mCherryN1 which was a gift from Chandra Tucker (Addgene plasmid # 26866)\(^{60}\). attB5 and attB2 sites were added to the CRY2-mCh PCR product and recombined into pDONR P5-P2 vector. Both pDONR vectors with Armadillo and CRY2-mCh insert were recombined by Gateway cloning (Invitrogen) into pUASg,attB with COOH-terminal 3XHA tag (A kind gift from J. Bischof and K. Basler, Zurich)\(^{45}\) to obtain the Arm-CRY2-mCh transgene for expression in Drosophila. Transgenes were injected into attP2 (Strain #8622) P\[CaryP\]attP2 68A4 by BestGene Inc. (California)\(^{46}\). For constitutive expression in S2R\(^+\) cells, the pDONR vectors were recombined into the Gateway destination vector, pAW (Drosophila Gateway\(^{39}\) Vector collection, Carnegie Institution). For constitutive expression in mammalian cell lines, the pDONR vectors were recombined into the mammalian Gateway destination vector, Gateway pCNA-pDEST40 (ThermoFisher Scientific, USA). EGFP-tagged protein lines were derived using the Minos mediated integration cassette (MiMIC) methodology described by Nagarkar-Jaiswal et al.\(^{47,48}\) and the Arm MiMIC line was obtained from the Bloomington Drosophila Stock Center (BDSC).

Crosses and expression of UAS constructs. The dominant female sterile technique was used to generate maternally mutant eggs\(^{22}\). Please see Flybase for details on mutants used (flybase.bio.indiana.edu). For mis-expression experiments, the ArmGAL4\(^{2}\) 2\(^{nd}\) chromosome was used. FRT 101 was used to generate the X chromosome mutant.

The following cross was conducted to generate germline clone embryos:

\[ y^{-}, \text{arm}^{X} \text{FRT101/FRT101} \text{; Arm-Gal4 x w\(^{-}\), y\(^{-}\)}; UAS-Arm-CRY2-mCh \]

Embryos were imaged in the light sheet microscope showing mCh expression, and were further cuticle prepped after imaging to confirm the genotype and phenotype as described for y\(^{-}\), arm\(^{X}\) mutation, an assay based on the pigmentation of the denticles\(^{41,42,49}\).

The following cross was conducted to generate zygotic mutant rescue flies:

\[ y^{+}, \text{arm}^{X} \text{FRT101/FM6; Arm-Gal4 x w\(^{-}\), y\(^{+}\)/Y-chromosome; UAS-Arm-CRY2-mCh} \]

In this cross, all males receive the X-chromosome from their mothers so only balance (FM6, Bar) males are expected unless the transgene can rescue the lethality of arm\(^{X}\). Identical crosses were set up with three kept entirely in the dark and three under normal laboratory light conditions for two weeks. The number and phenotype of males from all the crosses was scored.

Light-sheet microscopy. Embryos were dechorionated in bleach, washed in water and dried on a paper towel. They were then embedded into a capillary containing 1% agarose, low gelling temperature Type VII-A (Sigma) dissolved in water, using a fine wire such that the embryo was upright with the anterior-posterior axis aligned with the axis or perpendicular to the axis of the glass capillary. For imaging, the agarose was pushed out of the capillary and the sample was suspended freely in the sample chamber containing water as our lightsheet set-up utilized a water-immersion objective.

Live imaging of Drosophila embryos was carried out on the light-sheet Z.1 fluorescence microscope (Carl Zeiss, Germany) with the Lightsheet Z.1 10x/0.2 Illumination Optics and W Plan-Apochromat 20x/1.0 UV-VIS detection objective (Carl Zeiss, Germany). The lightsheet microscope was equipped with the Lightsheet Z.1 detection module “Standard”, 30 mW 488 nm and 20 mW 561 nm solid state lasers with BP505–545 and LP585 emission filters respectively. Whole embryos (unless otherwise stated) were imaged using dual-side illumination by a light-sheet modulated into a pivot scan mode. The 488 nm laser was used at 6% power with 7.5ms exposure time and the 561 nm laser was used at 13% laser power with 12.5ms exposure time. Since the oligomerization is reversible with removal of blue light, the whole embryo was excited with both laser lines every 2.5 min in Z-stack mode at 1 μm step size for 24 hrs. For control embryos not exposed to blue light, the same settings were used but with illumination using only the 561 nm laser line. All single plane illumination (SPIM) data was saved in the LSM format and processed using the ZEN 2014 SP1 software (Carl Zeiss, Germany). Background fluorescence was automatically calculated by the ZEN 2014 software using the background correction function (ZEN Software Guide, Carl Zeiss, Germany). Z-stacks that were deep in the embryo and out of focus were removed and maximum intensity projections of z-stacks in focus were generated for each embryo\(^{50}\). Images for figures were processed and assembled using Photoshop and Illustrator programs from Adobe.

Antibodies and Immunofluorescence. Embryos were dechorionated in bleach and fixed with heptane/4% formaldehyde in phosphate buffer (0.1 M NaPO4 pH 7.4) for 20 minutes\(^{41}\). The aqueous phase was removed and an equal amount of methanol was added to devitellinize the embryos. Antibody stainings were done in PBT (PBS, 0.1% Triton X-100, 1% bovine serum albumin, 0.1% Azide). The anti-Engrailed antibody mAbs (4D9) was obtained from the Developmental Studies Hybridoma Bank, created by the NICHD of the NIH and maintained at The University of Iowa, Department of Biology, Iowa City, IA 52242 and used at a dilution of 1:10. Alexa Fluor 488- anti-mouse secondary antibody was used (A-11001, ThermoFisher Scientific, USA) at a 1:1000 dilution. Embryos were mounted in Aquapalymount\(^{8}\) (Polysciences, Inc.). Images were acquired on the Zeiss AxioImager Z1 with Apotome (Carl Zeiss, Germany) and images were processed using the AxioVision Rel. 4.8 (Carl Zeiss, Germany).

Live-cell imaging. Time-lapse microscopy of activated CRY2 fusions in S2R\(^+\) and endogenously expressing WNT3A in a HEK 293 cell line that contains an integrated TOPFlash reporter (A kind gift from Dr Alan Prem
Kumar, Cancer Science Institute of Singapore) was performed using a Zeiss AxioImager Z1 microscope (Carl Zeiss, Germany). Clustering visualization was carried out at room temperature. Blue light exposure and mCherry imaging were performed simultaneously by imaging in both 488 nm and 561 nm laser channels. Images were acquired and processed using the AxioVision Rel. 4.8 (Carl Zeiss, Germany).

In order to quantify the number of CRY2 puncta, we used the Cell Counter plugin in ImageJ (NIH, https://imagej.nih.gov/ij/plugins/cell-counter.html). This tool allows manual and semiautomatic counting of user determined points within an image. The puncta numbers were determined for several images and graphed in Fig. 1.

**TOPflash assay.** TOPflash luciferase assays (TCF/LEF reporter assays) were performed to assess the effect of light on canonical Wnt-signaling in cells transfected with the Arm-CRY2-mCh plasmid. S2R+ cells were co-transfected with TOPflash32, Renilla luciferase-Pol III (Renilla luciferase-Pol III was a gift from Norbert Perrimon (Addgene plasmid # 37380))33 and the pAW-Arm-CRY2-mCh plasmid using lipofectamine 3000 relative quantification (2−ΔΔCT) normalized to the endogenous control, RPL32, followed by calibration to cultures not exposed to light using dissociation curve was generated to ensure specificity of amplification. For expression analysis, the mRNA data were mined points within an image. The puncta numbers were determined for several images and graphed in Fig. 1.

**RNA Extraction, cDNA Synthesis and qPCR.** Mixed stage and genotype embryos, 0 to 16 hours after deposition, were exposed to light or no light for 6 hours, dechorionated in bleach, washed in water and collected. Total RNA was extracted for each treatment using the RNeasy Mini Kit (Qiagen) as per the manufacturer's protocol. Total RNA concentration and purity was measured using the Cytation 3 Cell Imaging Multi-Mode Reader (BioTek). One μg of total RNA was reverse transcribed in a 20 μl reaction volume using the QuantiTect reverse transcription kit (Qiagen) according to the manufacturer's protocol. Gene specific primer sequences were obtained from Fly Primer Bank52. Quantification of mRNA was performed using SYBR® Green Assay (Thermo Fisher Scientific) on the PikoReal™ Real-Time PCR System (Thermo Fisher Scientific) and a PCR product dissociation curve was generated to ensure specificity of amplification. For expression analysis, the mRNA data were normalized to the endogenous control, RPL32, followed by calibration to cultures not exposed to light using relative quantification (2−ΔΔCT). Results were generated from 3 technical replicates for each mRNA. The average relative expression ± standard deviation (SD) was determined. Statistical analyses were performed using Student's t-test unless the data showed evidence of unequal standard deviation (F-test, p < 0.05). In these cases, the Mann–Whitney nonparametric test was used.

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Author Contributions
P.K. and N.S.T. designed and performed the experiments. N.S.T., P.K. and T.E.M. wrote the paper. All authors reviewed manuscript.

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