ABSTRACT
How differential levels of gene expression are controlled in post-mitotic neurons is poorly understood. In the Drosophila retina, expression of the transcription factor Defective Proventriculus (Dve) at distinct cell type-specific levels is required for terminal differentiation of color- and motion-detecting photoreceptors. Here, we find that the activities of two cis-regulatory enhancers are coordinated to drive dve expression in the fly eye. Three transcription factors act on these enhancers to determine cell-type specificity. Negative autoregulation by Dve maintains expression from each enhancer at distinct homeostatic levels. One enhancer acts as an inducible backup (‘dark’ shadow enhancer) that is normally repressed but becomes active in the absence of the other enhancer. Thus, two enhancers integrate combinatorial transcription factor input, feedback and redundancy to generate cell type-specific levels of dve expression and stable photoreceptor fate. This regulatory logic may represent a general paradigm for how precise levels of gene expression are established and maintained in post-mitotic neurons.

INTRODUCTION
Genes are expressed at distinct cell type-specific levels at different times during development. Expression is often transient, arising for short periods of time to trigger downstream regulatory pathways. For example, expression driven by the eve stripe 2 enhancer, perhaps the best-understood regulatory DNA element, is very short-lived, persisting for only ∼15 min after the mature stripe is fully formed during embryonic development in flies (Bothma et al., 2014). By contrast, gene expression in post-mitotic neurons must be maintained on long timescales, often for the lifetime of the organism. Establishing and maintaining distinct levels of transcription factors is particularly important for neuronal fate and function across species. For example, in worms, low levels of the transcription factor MEC-3 specify the elaborate dendritic patterning of PVD pain-sensing neurons, whereas high MEC-3 determines the simple morphology of AVM and PVM touch neurons (Smith et al., 2013). Similarly, flies use differences in levels of the homeodomain transcription factor Cut to control dendritic branching complexity in sensory neuron subtypes (Grueber et al., 2003). In mice, the Hox accessory factor FoxP1 acts as a dose-dependent determinant of motor neuron subtype identity (Dasen et al., 2008). Beyond these cases, there are numerous examples of differential transcription factor expression in neuronal subtypes, such as the unique expression levels of Brn3b in ipRGC subtypes (Chen et al., 2011).

Establishing and maintaining distinct levels of gene expression for the lifetime of a neuron presents specific challenges. Regulatory mechanisms must ensure that expression levels remain within a narrow range for days and even years while providing robustness against acute perturbations caused by activity and environment. In some cases, the transcription factors that dictate cell type-specific expression levels have been identified (Corty et al., 2016), but how these regulatory inputs are interpreted by DNA elements has not been characterized. Furthermore, it is unclear how transcription factor feedback and cis-regulatory redundancy contribute to ensuring proper expression levels in neurons.

Expression of transcription factors at cell type-specific levels is required for the terminal specification of motion- and color-detecting photoreceptors in the Drosophila retina. The Drosophila compound eye consists of approximately 800 ommatidia, or unit eyes, each containing eight photoreceptors (PRs) (Wolff and Ready, 1993) (Fig. 1E). The outer PRs (R1-R6) express the broad spectrum-sensitive Rhodopsin 1 (Rh1) and detect motion (Hardie, 1985), whereas the inner PRs (R7 and R8) express color-sensitive Rhodopsin proteins (Rh3-Rh6) (Gao et al., 2008; Yamaguchi et al., 2010). Two ommatidial subtypes, pale (p) and yellow (y), are randomly distributed in the retina at a ratio of 35:65 (Bell et al., 2007; Franceschini et al., 1981) (Fig. 1A-D). The p subtype contains UV-sensitive Rh3 in pR7 and blue-sensitive Rh5 in pR8, whereas the y subtype contains UV-sensitive Rh4 in yR7 and green-sensitive Rh6 in yR8 (Fig. 1A-C) (Chou et al., 1996; Fortini and Rubin, 1990; Johnston and Desplan, 2010). The specification of these photoreceptor subtypes is controlled by a complex network of transcription factors and other regulators (Hsiao et al., 2013; Jukam and Desplan, 2011; Jukam et al., 2013, 2016; Mikeladze-Dvali et al., 2005; Viets et al., 2016; Wernet et al., 2006).

Differential expression of the K50 homeodomain transcription factor Defective proventriculus (Dve) is crucial for terminal specification of photoreceptors in the fly eye (Johnston et al., 2011; Thanawala et al., 2013). Dve is expressed in a unique pattern, with high levels in the outer PRs, low levels in yR7s and no expression in pR7s or R8s (Fig. 1K). High Dve in motion-detecting outer PRs represses expression of color-detecting Rh3, Rh5 and Rh6. Low levels of Dve in yR7s repress Rh3 to maintain exclusive expression of Rh4 in yR7 subtypes in the main ventral region of the retina (Fig. 1L,
Changes in levels of Dve expression have a dramatic impact on Rhodopsin expression and photoreceptor fate. In dve null mutants, Rh3 is derepressed in all R7s, and Rh3, Rh5 and Rh6 are variably expressed in outer PRs (Fig. 1O-Q) (Johnston et al., 2011; Sood et al., 2012). In dve hypomorphic mutants, where levels of Dve are lowered but not completely lost, Rh3 is still derepressed in all R7s, but only Rh6 is expressed in outer PRs (Johnston et al., 2011). When Dve levels are subtly lowered upon mutation of upstream regulators, the dorsal region of Rh3 and Rh4 co-expression is expanded from one-third of the retina to the entire dorsal half (Thanawala et al., 2013). The misexpression of Rh5s in dve mutants causes defects in low-intensity light discrimination (Johnston et al., 2011). Deleterious effects are also seen when Dve levels are increased: raising levels of Dve in yR7s causes loss of Rh3/Rh4 co-expression in the specialized dorsal third region (Mazzoni et al., 2008; Thanawala et al., 2013), whereas overexpression in R8s represses Rh5 and Rh6 completely (Johnston et al., 2011). Thus, the differential expression of dve in photoreceptors is important for proper Rh expression and visual function.

Cell type-specific levels of Dve are achieved through regulation by the K50 homeobox transcription factor Orthodenticle (Otd), the zinc-finger transcription factors Spalt major and Spalt related (referred to collectively as Spalt (Sal)), and the PAS-bHLH transcription factor Spineless (Ss). Otd activates Dve in all PRs (Fig. 1F), Sal represses Dve in the inner PRs (Fig. 1G-H) and Ss re-activates Dve in yR7s (Fig. 1I-J) (Johnston, 2013; Johnston et al., 2011).

To determine how these transcription factors dictate cell type-specific levels of Dve expression, we analyzed the cis-regulatory logic controlling dve and identified two enhancers, yR7 enh and outer enh, that together induced expression recapitulating endogenous Dve expression. yR7 enh is activated by Ss, Sal and Otd in yR7 cells, whereas outer enh is activated by Otd in all PRs and repressed by Sal in inner PRs. Negative feedback by Ove onto both enhancers maintains proper levels of Dve expression. This autoregulation is particularly important for yR7 enh, which is dramatically upregulated in yR7s when Ove feedback is ablated. Interestingly, we also observed derepression of yR7 enh in outer PRs in dve mutants, suggesting that yR7 enh serves as an inducible backup or ‘dark’ shadow enhancer in these cells. Shadow enhancers are DNA elements that drive redundant expression patterns and ensure robust gene expression in cases of genetic and environmental perturbation (Bothma et al., 2015; Frankel et al., 2010; Hong et al., 2008; Miller et al., 2014; Nolte et al., 2013; Perry et al., 2010; Wunderlich et al., 2015). yR7 enh represents an unusual ‘dark’ shadow enhancer as it is normally repressed and only becomes active when Dve driven by the primary outer enh is compromised. Together, the yR7 enh and outer enh integrate combinatorial transcription factor input, negative feedback and redundancy to ensure distinct cell type-specific levels of dve expression required for stable photoreceptor specification.

**RESULTS**

**Two enhancers determine yR7- and outer PR-specific expression of Dve**

The dve gene locus is ~65 kb with two alternative transcriptional starts driven by the dve-A promoter or dve-B promoter (Fig. 2A). Deletion of the dve-A promoter caused derepression of Rh3 in yR7s in the dorsal half of the retina (Fig. S1A,B), while Rh5 and Rh6 expression were unaffected (Fig. S1C). This incomplete dve
phenotype is consistent with a decrease in Dve levels in yR7s (Thanawala et al., 2013), suggesting that the dve-A promoter is required for normal Dve expression. To test the role of the dve-B promoter, we employed a CRISPR strategy to delete a ∼1.5 kb region encompassing the dve-B promoter and first exon. Deletion of the dve-B promoter did not alter Dve-regulated Rh expression (Fig. S1D-F), suggesting that the dve-B promoter is not required for Dve expression. As the dve-A promoter is required for normal Dve expression, we used this promoter as the minimal promoter in enhancer reporters.

To identify cis-regulatory elements controlling Dve expression, we generated transgenes containing 3-6 kb DNA fragments from the dve locus and the dve-A promoter driving nuclear GFP (Fig. 2A, dve enh>GFp). The dve-A promoter alone drove extremely weak GFP expression in pigment cells and R4 PRs, and therefore did not recapitulate normal Dve expression in all outer PRs and yR7s (Fig. S1H).

Two constructs drove GFP expression that together recapitulated endogenous Dve expression in midpupation [i.e. ∼48 h after puparium formation (APF)]. outer enh drove expression in outer PRs (Fig. 2A,E), and yR7 enh drove expression specifically in a subset of R7s (Fig. 2A,B). This subset corresponded to yR7 fate, as 68% of R7s had strong GFP expression and perfectly co-expressed Ss (i.e. yR7s), whereas 32% had weak or no GFP and lacked Ss (i.e. pR7s) (Fig. 2B-D).

Additionally, weak yR7 enh drove weak expression in yR7s (Fig. 2A, Fig. S1M-O, described further below), and dorsal R7 enh drove expression in dorsal posterior R7s (Fig. 2A, Fig. S1P-Q). Four enhancers drove weak expression in all PRs (all PRs enh 1-4) (Fig. 2A, Fig. S1I-L).

Janelia Research Campus and the Vienna Drosophila Resource Center (VDRC) both generated lines that express Gal4 driven by fragments of the dve locus (Fig. 2A). Expression driven by these fragments was consistent with results from our dve enh reporter constructs. GMR40E08, a ∼3 kb fragment that overlaps with outer enh, drove strong GFP expression in outer PRs, whereas other constructs that either did not overlap or only partially overlapped with outer enh or yR7 enh did not show significant expression (Fig. 2A).

As yR7 enh and outer enh recapitulated endogenous Dve expression, we further characterized the temporal dynamics of these two enhancers. At midpupation, Dve protein is expressed strongly in outer PRs and weakly in yR7s (Johnston et al., 2011), similar to GFP expression driven by yR7 enh and outer enh (Fig. S2B,F,J). In third instar larvae, analysis of Dve protein expression was obscured by non-specific antibody staining (Fig. S2A) (Johnston et al., 2011). Although outer enh was not expressed, yR7 enh was expressed in a subset of R7s (Fig. S2E,I), suggesting that Dve is expressed in larval yR7s. In adults, Dve protein is expressed in yR7s and outer PRs (Fig. S2C,D). Similarly, outer enh drove GFP expression in outer PRs in adults (Fig. S2K,L). yR7 enh drove expression in all R7s in adults (Fig. S2G,H), suggesting that additional activators present only in the adult stage induce yR7 enh expression in all R7s, and that this enhancer is missing DNA elements that prevent ectopic Dve expression in adults.

Together, the spatiotemporal dynamics of these enhancers are consistent with endogenous dve expression. Next, we tested how upstream transcription factors control expression of these two enhancers.

yR7 enh is activated by Ss, Sal and Otd

yR7 enh drives expression in yR7 cells (Fig. 3A). Dve is expressed at lower levels in yR7s in the dorsal third, allowing IroC-induced activation of Rh3 and co-expression of Rh3 and Rh4 (Johnston et al., 2011). In yR7 enh, yR7 enh is expressed at lower levels in dorsal third (DT) yR7s when compared with the rest of the retina (Fig. 3D).
Otd is required for Dve expression in yR7s (Johnston et al., 2011). yR7 enh failed to induce expression in yR7s in otd mutants, suggesting that Otd is required for activation of this enhancer (Fig. 3B).

Ss induces expression of Dve in yR7s (Johnston et al., 2011). Expression of yR7 enh was lost in ss mutants (Fig. 3C). Ectopic expression of Ss in all PRs induced strong yR7 enh expression in all R7s and weak expression in all other PRs (Fig. 3E), suggesting that another factor acts with Ss to activate strong yR7 enh expression.

As Sal is important for R7 fate (Mollereau et al., 2001), we posited that Sal may work with Ss to activate yR7 enh. Expression of yR7 enh was completely lost in sal mutants (Fig. 3F), whereas ectopic expression of Sal in all PRs induced yR7 enh expression in a random subset of R1 and R6 outer PRs (Fig. 3G). We showed previously that ectopic Sal induced Ss in a random subset of R1 and R6 outer PRs (Johnston and Desplan, 2014). These data suggest that Ss and Sal function together to activate expression of yR7 enh.

Supporting our hypothesis, ectopic expression of both Ss and Sal induced strong yR7 enh expression in all PRs (Fig. 3H), suggesting that Ss and Sal both activate expression of yR7 enh. As Sal induces expression of Ss, and Ss together with Sal induces yR7 enh, Ss, Sal and yR7 enh form a coherent feed-forward loop (Fig. 3I).

To further elucidate these combinatorial regulatory interactions, we truncated yR7 enh to a 0.8 kb fragment (yR7 enh*) that recapitulated yR7 enh expression driven by the entire yR7 enh fragment (Fig. 2A; Fig. 3J,L, Fig. S3A). Three other truncations that encompass the 0.8 kb region also recapitulated yR7 enh expression, whereas two truncations and four GAL4 lines generated by Janelia Research Campus and VDRC that excluded yR7 enh* failed to drive GFP expression, consistent with the role of yR7 enh* in driving yR7 enh expression (Fig. 3J). yR7 enh* contains three conserved Ss binding sites (called Xenobiotic Response Elements/XREs) (Fig. 3K), consistent with regulation by Ss.

weak yR7 enh drive weak GFP expression in yR7s, colocalizing with Ss expression (Fig. S1M-O). weak yR7 enh and yR7 enh* share a ~250 bp overlap that contains one of the three Ss XRE binding sites (Fig. 3K), suggesting that while the shared XRE site can drive GFP in yR7s, strong expression requires the presence of additional XRE sites. The Janelia enhancer GMR42E10 shares a ~75 bp overlap with yR7 enh but does not contain any Ss XRE binding sites (Fig. 3K). This construct failed to drive GFP expression, suggesting that at least one Ss XRE binding site is required for yR7-specific expression.

To further test the roles of Ss XRE binding sites, we generated a yR7 enh* construct that replaces all GCGTG Ss XRE binding sites with AAAA. This construct showed a near complete loss of yR7 GFP expression, indicating the importance of these sites for Ss activation (Fig. 3M). Very low-level expression of this reporter suggests the presence of additional cryptic Ss sites within yR7 enh* (Fig. 3M). Searching yR7 enh* for low-affinity Ss binding motifs (Zhu et al., 2011), we identified two putative sites (GTCTGA and GTGTGA), one of which is conserved (GTCTGA), suggesting that these cryptic/low-affinity sites may drive very low level expression in the absence of core conserved (GCGTG) sites. Together, these data suggest that Ss directly binds the XRE sites in yR7 enh* to regulate expression. However, we cannot rule out possible indirect mechanisms.

Although yR7 enh* has three Ss XRE sites, this enhancer contains no predicted Sal sites (Barrio et al., 1996; Sanchez et al., 2011), suggesting that Sal regulates yR7 enh* either directly via binding to cryptic sites or indirectly through regulation of other intermediary factors. The longer yR7 enh contains a Sal binding site, which may contribute to regulation. Genetic epistasis analysis supports an indirect mode of regulation by Sal (Fig. S4; see below).

yR7 enh* is required for expression of endogenous Dve in yR7s, as CRISPR-generated deletion of yR7 enh* caused a loss of Dve expression specifically in R7s (Figs 2A, 3N) and a corresponding upregulation of Rh3 in all PRs (Fig. 3O). Similarly, the larger dveA deleletion, covering yR7 enh and the dve-A promoter, also resulted in Rh3 upregulation in R7s (Fig. 2A, Fig. S1G). Together, these results suggest that yR7-specific expression of Dve requires yR7 enh, which is activated by Ss, Sal and Otd.

Negative feedback onto yR7 enh determines homeostatic levels

Expression levels of Dve are precisely controlled to determine region-specific activation or repression of Rh3 in yR7s (Thanawala et al., 2013) (Fig. 1D). Negative feedback is a mechanism that ensures precise, homeostatic levels of gene expression. As Dve is a transcriptional repressor, we hypothesized that Dve feeds back onto yR7 enh to control expression levels. To test Dve for negative regulation of yR7 enh, Dve was expressed in all PRs at high levels causing a complete loss of yR7 enh expression (Fig. 4A). yR7 enh was expressed at higher levels in yR7s in dve mutant clones compared with wild-type clones (Fig. 4B,C,F), suggesting that Dve driven by yR7 enh feeds back to control levels of expression in yR7s (Fig. 4H).

yR7 enh is a ‘dark’ shadow enhancer for outer PR expression

In addition to yR7s, expression of yR7 enh occurred in outer PRs in dve mutant clones (Fig. 4D,E,G), suggesting that outer enh induces Dve expression to completely repress yR7 enh in outer PRs in normal conditions (Fig. 4J). As yR7 enh was never expressed in pR7s or R8s in wild type or in dve mutants (Fig. 4B-E), yR7 enh is only competent to drive expression in yR7s and outer PRs, where Dve is normally expressed.

As outer enh drives expression in outer PRs in normal conditions and yR7 enh drives expression in outer PRs in dve mutants, we predicted that deleting outer enh would cause yR7 enh to drive expression of endogenous dve in outer PRs (Fig. 4J). Flies with a CRISPR-mediated deletion of outer enh displayed expression of Dve in outer PRs (Fig. 4K) and repression of Rh3, Rh5 and Rh6 (i.e. Dve target genes) in outer PRs in 1-week-old adults (Fig. 4L,N), suggesting that yR7 enh drives expression in the absence of functional outer enh. Although Rh3 expression remained unchanged (Fig. 4M), variable derepression of Rh5 and Rh6 occurred in 4-week-old adults (Fig. 4O), suggesting that expression driven by yR7 enh is not sufficient to completely rescue Dve expression due to differences in levels or timing.

As yR7 enh can drive expression in outer PRs, yR7 enh is a shadow enhancer (i.e. redundant regulatory DNA element) for outer enh, the primary enhancer for outer PR expression. Unlike typical shadow enhancers, the yR7 enh shadow enhancer is repressed (‘dark’) in outer PRs under normal conditions due to negative feedback from the primary enhancer (Fig. 4I). We therefore define yR7 enh as a ‘dark’ shadow enhancer, as its expression in outer PRs only occurs when outer enh function is lost (Fig. 4J).

Otd/Dve sites play context-dependent roles in yR7 enh

As Otd activates and Dve represses yR7 enh, we next tested the regulatory roles of canonical Otd/Dve binding sites (also called K50 sites; TAATCC). yR7 enh* contains two Otd/Dve sites, which are perfectly conserved across at least five out of six Drosophila species (Fig. 3K). Replacing these two sites with AAAA caused increased levels of GFP expression in yR7s (Fig. 4P), suggesting that these sites mediate repression by Dve but not activation by Otd in yR7s. As Otd is required for expression of yR7 enh, the expression of GFP in yR7s in the absence of optimal Otd binding
Fig. 3. *yR7 enh* is activated by Otd, Sal and Ss. (A-C,E-H,L-M) Yellow circles indicate yR7s; solid white circles indicate pR7s. Dashed white circles indicate outer PRs and R8s. Light green in ommatidium schematics indicates strong GFP expression; dark green indicates weak expression; crosshatch indicates variable expression; black indicates lack of expression. Images were acquired at mid-pupation. (A) *yR7 enh* is expressed in yR7s. (B) Expression of *yR7 enh* is lost in *otd* mutants. (C) Expression of *yR7 enh* is lost in *ss* mutants. (D) Quantification of GFP intensity in R7 cells shows three distinct intensity levels corresponding to pR7 [including pR7 and dorsal third (DT) pR7], yR7 and DT yR7 expression. Data are means±s.d., *n* = 22 for pR7s, 16 for DT pR7s, 31 for yR7s and 31 for DT yR7s. ****P < 0.0001, ns indicates P > 0.05 and not significant (unpaired *t*-test with Welch’s correction). All measurements were internally controlled within a single mid-pupal retina. (E) *yR7 enh* is strongly expressed in all R7s and weakly expressed in all PRs when Ss is ectopically expressed in all PRs. (F) Expression of *yR7 enh* is lost in *sal* mutants (white circle indicates presumptive R7). (G) *yR7 enh* is expressed in random R1s and R6s when Sal is ectopically expressed in all PRs. (H) *yR7 enh* is expressed in all PRs when Ss and Sal are ectopically expressed in all PRs (yellow circle indicates presumptive yR7). (I) The regulatory interactions governing *yR7 enh*. Otd, Ss and Sal activate *yR7 enh*, whereas Sal activates stochastic expression of Ss in yR7s (denoted by dashed arrow). (J) A truncated 0.8 kb fragment of *yR7 enh*, indicated by *yR7 enh**, was sufficient to recapitate GFP expression in yR7 cells. Larger truncations encompassing *yR7 enh* also expressed GFP in yR7 cells, while truncations excluding *yR7 enh* did not drive GFP expression. weak *yR7 enh* shares a ~250 bp overlap with *yR7 enh**, including one of the three Ss XRE binding sites (Fig. 3K). GMR42E10, a construct generated by Janelia that contains a fragment of dve driving Gal4, shares a ~75 bp overlap with *yR7 enh* that does not contain any Ss XRE binding sites (Fig. 3K). This construct failed to drive GFP expression in yR7 cells. Light-green fragments drive strong GFP expression; dark-green fragments drive weak GFP expression; gray fragments do not drive GFP expression. (K) *yR7 enh** contains three Ss binding sites and two Otd/Dve binding sites. Capitalized black text indicates perfect conservation across six Drosophila species. Capitalized gray indicates conservation across five out of the six species. Light-green fragments drive strong GFP expression; dark-green fragments drive weak GFP expression; gray fragments do not drive GFP expression. (L) *yR7 enh* is expressed in yR7s, similar to Dve and *yR7 enh*. (M) Knocking out Ss XRE binding sites in the *yR7 enh* construct resulted in a near complete loss of GFP expression. BS KO, binding site knockout. (N) CRISPR-mediated deletion of *yR7 enh* from the endogenous dve locus resulted in loss of Dve specifically in yR7s. Dashed white circles indicate outer PRs and R8s; solid white circles indicate R7s. Red in ommatidium schematic indicates Dve expression. (O) Loss of Dve in yR7s resulted in derepression of Rh3 in adults. Yellow circles indicate yR7s; white circles indicate pR7s; black circles indicate no expression.
sites suggests that Otd may act through additional Otd-specific cryptic sites or that activation is mediated by another activator downstream of Otd. Mutation of these sites did not cause derepression in outer PRs, suggesting that these sites mediate both repression by Dve and activation by Otd in outer PRs.

To test whether Dve directly binds the two Otd/Dve sites in \textit{yR7 enh*}, we conducted \textit{in vitro} electrophoretic mobility shift assays (EMSAs). Dve bound sequences containing the Otd/Dve sites, and mutation of these sites dramatically decreased binding (Fig. 4Q), suggesting that Dve directly binds the two Otd/Dve sites in \textit{yR7 enh*} to repress expression.

As regulation of \textit{yR7 enh*} is dependent on Otd/Dve sites, Otd likely directly binds these sites to regulate expression. However, we cannot rule out possible indirect mechanisms.

\textbf{outer enh is activated by Otd and repressed by Sal}

We next characterized \textit{outer enh}, the primary enhancer for Dve expression in outer PRs (Fig. 5A). The \textit{dve\textsuperscript{del}} deletion, which removes the first exon of \textit{dve}, the \textit{dve-A} promoter, and \textit{yR7 enh}, showed no derepression of Dve target genes (Rh3, Rh5 and Rh6) in outer PRs (Fig. 2A, Fig. S1G), suggesting that \textit{outer enh} is sufficient to drive Dve expression in outer PRs.

Otd activates Dve expression in all PRs, and Sal represses Dve expression in inner PRs (Johnston et al., 2011). \textit{outer enh} expression was completely lost in \textit{otd} mutants, consistent with a general requirement of Otd for \textit{dve} expression (Fig. 5B). In \textit{sal} mutants, \textit{outer enh} was derepressed in inner PRs (Fig. 5C), suggesting that Sal represses this element in inner PRs. Ectopic expression of Ss in all PRs did not affect \textit{outer enh} expression, consistent with

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Fig. 4. See next page for legend.
Fig. 4. Dve feeds back to control yR7 enh. (A-E) Expression analysis was conducted on mid-pupal retinas. (A) Expression of yR7 enh is lost when Dve is ectopically expressed in all PRs. Dashed white circles indicate outer PRs and R8s; solid white circle indicates R7. In schematic, black circles indicate no GFP expression. (B-E) Yellow circles indicate yR7 cells; white circles indicate pr7 cells. Dashed white circles are outer PRs. Solid gray lines represent the boundary between dve mutant clones (indicated by the absence of GFP) and wild-type clones (indicated by the presence of GFP). Green in ommatidia schematic indicates strong GFP expression; crosshatching indicates variable expression; black indicates lack of expression; red spot indicates GFP expression. (B,C) In yR7s, yR7 enh is upregulated in dve mutant clones compared with wild-type clones. (D,E) In outer PRs, yR7 enh is upregulated in dve mutant clones compared with wild-type clones. (F) Quantification of yR7 enh intensity of outer PRs in dve mutant and wild-type clones. yR7s in dve mutants show greater GFP intensity than in wild-type clones. R7 cells that are GFP positive indicate yR7s. n=37 for wild-type yR7s and n=37 for dve mutant yR7s. ****P<0.0001, unpaired t-test with Welch’s correction. All measurements were internally controlled within a single mid-pupal retina. (G) Quantification of GFP intensity of outer PRs in dve mutant and wild-type clones. In wild-type clones, outer PRs are GFP off, whereas dve mutant clones show a much greater distribution of GFP expression states. n=84 for wild-type outer PRs and n=54 for dve mutant outer PRs. ****P<0.0001, unpaired t-test with Welch’s correction. All measurements were internally controlled within a single mid-pupal retina. (H) yR7 enh induces Dve expression that negatively feeds back onto yR7 enh to maintain homeostatic Dve levels in yR7 cells. (i) outer enh induces Dve expression that negatively feeds back onto yR7 enh to completely repress yR7 expression in outer PRs. (J) When outer enh function is impaired, yR7 enh is derepressed in outer PRs. (K) Dve remains expressed in outer PRs upon deletion of outer enh. Dashed white circles indicate outer PRs. (L,N) Expression of downstream Dve targets (Rh3, Rh5 and Rh6) is unaffected in outer enh deletion mutants in 1-week-old adults. Dashed white circles indicate outer PRs. (M,O) Variable derepression of Rh5 and Rh6 in outer PRs is observed in outer enh deletion mutants in 4-week-old adults. Expression of Rh3 is unaffected in outer enh deletion mutants in 4-week-old adults. Dashed white circles indicate outer PRs. (P) Knocking out Otd/Dve K50 binding sites resulted in an increased level of GFP in yR7s, suggesting that these sites mediate repression by Dve but not activation by Otd in yR7s. Solid yellow circles indicate yR7s that express GFP; solid white circles indicate pr7s that do not express GFP; dashed white circles indicate outer PRs and R8s. BS KO, binding site knockout. (Q) EMSAs illustrating that the binding of Dve is dependent on K50 Otd/Dve sites in yR7 enh. WT, wild-type sequence; M, mutation of K50 Otd/Dve site. Arrows indicate the bands shifted upon Dve binding. Multiple bands are observed likely due to the presence of multiple functional DNA binding domains within Dve (Johnston et al., 2011), yielding higher-order DNA/protein structures.

regulation of this element independent of Ss (Fig. 5D). Thus, combinatorial regulation involving activation by Otd in all PRs and repression by Sal in inner PRs yields the outer PR-specific expression of outer enh (Fig. 5E).

We truncated outer enh to a 1.3 kb fragment (outer enh*) that recapitulated the expression of the entire outer enh fragment (Figs 2A, 5A,F,H, Fig. S3B). Two larger truncations and a Janelia Gal4 construct (GMR40E08) that encompass this 1.3 kb region also recapitulated expression, whereas fragments that exclude outer enh* failed to drive GFP, consistent with the role of outer enh* in driving outer PR-specific expression (Fig. 5F).

outer enh* has four K50 homeodomain consensus sites (TAATCC) for Otd and Dve (Fig. 5G) (Chaney et al., 2005). all PR enh 4 shares a 390 bp overlap with outer enh*, including one of the Otd/Dve binding sites, suggesting that its weak expression in all PRs may be due to the single Otd/Dve binding site functioning independently of the repressive Sal input that regulates the entire outer enh*.

We generated an outer enh* construct that removes all TAATCC Otd/Dve binding sites by replacing them with AAAAAA (Fig. 5I). This construct showed a near complete loss of GFP expression in outer PRs, consistent with our model that Otd is required for outer enh activation. As regulation of outer enh* is dependent on Otd/Dve sites, Otd likely directly binds these sites to regulate expression. However, we cannot rule out possible indirect mechanisms.

Although outer enh* has four Otd/Dve sites, this enhancer contains no predicted Sal sites (Barrio et al., 1996; Sanchez et al., 2011), suggesting that Sal regulates outer enh* either directly via binding to cryptic sites or indirectly through regulation of other intermediary factors. The longer outer enh contains a Sal binding site, which may contribute to regulation.

Feedback onto outer enh determines homeostatic levels

As yR7 enh is controlled by negative autoregulation, we next tested whether feedback also determines expression levels driven by outer enh. As outer enh (and Dve) are highly expressed in outer PRs, we expected that dve mutants may exhibit subtle increases in expression from outer enh. Indeed, in dve mutant clones, outer enh was expressed at higher levels in outer PRS compared with wild-type clones (Fig. 6B-D). To confirm negative feedback onto outer enh, Dve was ectopically expressed in all PRs at high levels (all PRs>dve), causing a complete loss of outer enh expression (Fig. 6A). Thus, Dve driven by outer enh feeds back onto this enhancer to autoregulate and ensure homeostatic levels of expression in outer PRs (Fig. 6E).

To test whether Dve directly binds the four Otd/Dve sites in outer enh*, we conducted EMSAs. Dve bound sequences containing the Otd/Dve sites, and mutation of these sites dramatically decreased binding (Fig. 6F), suggesting that Dve directly binds the four Otd/Dve sites in outer enh* to repress expression.

Sal represses outer enh to allow Ss-mediated activation of yR7 enh

yR7 enh is highly sensitive to levels of Dve feedback, particularly in outer PRs where Dve levels are high. Ss alone is sufficient to induce yR7 enh expression at high levels in all R7s but not outer PRs (Fig. 3E). Ss and Sal together are sufficient to induce yR7 enh at high levels in outer PRs (Fig. 3H). As Dve driven by outer enh feeds back to repress yR7 enh in outer PRs (Fig. 4D,E,I,L) and Sal represses Dve expression from outer enh (Fig. 5C), Sal may activate yR7 enh by repressing outer enh.

One prediction of this model is that ectopic Ss should be sufficient to activate yR7 enh at high levels in outer PRs in the absence of Dve. Indeed, when Ss is expressed at high levels in all PRs in otd mutants that lack Dve (Johnston et al., 2011), yR7 enh is activated in all PRs (Fig. S4A).

This result highlights two facets of yR7 enh regulation. First, Ss activates yR7 enh, whereas Sal represses outer enh to allow expression of yR7 enh, suggesting that Sal interacts indirectly with yR7 enh (Fig. S4B). Second, Ss requires Otd to activate yR7 enh in wild-type conditions (Fig. 3B) where Ss levels are low, whereas high levels of Ss are sufficient to override the requirement for Otd (Fig. S4A).

DISCUSSION

Dve is expressed in an intricate pattern with distinct levels in different photoreceptors. The regulation required to achieve this pattern is complex, involving two enhancers controlled by three main mechanisms: combinatorial transcription factor input, negative feedback and enhancer redundancy (Fig. 7). PR-specific Otd, inner PR-specific Sal and yR7-specific Ss work together to induce expression of yR7 enh in yR7s (Fig. 7A). By contrast, Otd activates outer enh whereas Sal represses this enhancer to yield Dve expression in outer photoreceptors (Fig. 7B).
Once these cell type-specific patterns are set, negative feedback by Dve maintains expression of the two enhancers at distinct levels important for regulation of downstream rhodopsin genes (Fig. 7C,D). This negative feedback appears especially crucial for the yR7 enh, the expression levels of which determine activation or repression of Rh3 in different regions of the retina. Gene regulatory network motifs involving negative feedback minimize variation in expression levels. With negative feedback, high concentrations of a regulator repress its expression, whereas low levels allow its activation. Negative feedback thus ensures homeostatic levels of expression (Alon, 2007; Becskei and Serrano, 2000; Irvine et al., 1993; Stewart et al., 2013).

As an additional layer of regulation, outer enh drives high levels of Dve that repress yR7 enh in outer PRs (Fig. 7E). When outer enh function is lost, yR7 enh becomes active in outer PRs, functioning as a shadow enhancer to provide redundancy and robustness to expression (Fig. 7F). Complex multi-enhancer systems enable genes to integrate multiple regulatory inputs, yielding intricate expression patterns. Although some enhancers account for distinct aspects of regulation, others drive overlapping patterns. Shadow enhancers can compensate for removal of a primary enhancer, resulting in mostly unaltered gene expression (Hong et al., 2008; Miller et al., 2014; Nolte et al., 2013; Perry et al., 2012). These shadow enhancers provide reliability and robustness in pattern formation, allowing crucial patterning genes to be buffered against environmental and genetic variation (Barolo, 2012; Bothma et al., 2015; Frankel et al., 2010; Perry et al., 2010).

We define yR7 enh as a dark shadow enhancer, as it is normally repressed in outer PRs but becomes active when the function of the primary enhancer is impaired. We were able to identify the yR7 enh dark shadow enhancer because we were characterizing how a complex pattern was controlled by combinatorial transcription factor input and feedback acting on two enhancers. Similar to the generality of shadow enhancers (Cannavo et al., 2016), dark shadow enhancers may be a common mechanism to ensure gene expression. However, they would be challenging to identify as they are active only upon genetic or possibly environmental perturbation.

Dve is a transcriptional repressor (Johnston et al., 2011) that acts directly on yR7 enh in outer PRs to repress expression (Fig. 4Q). Generally, transcriptional repressors would likely act directly on dark shadow enhancers to repress them, poised them as backup systems. For transcriptional activators, more complex indirect mechanisms would be required. For example, the primary enhancer could induce the activator to activate expression of a...
transcriptional repressor, which in turn could repress the dark shadow enhancer. As dark shadow enhancers require feedback, they would likely only be found in genes encoding regulatory factors.

A key aspect of regulation by primary enhancers and dark shadow enhancers is their differential responsiveness to repression. For outer enh, normal Dve levels induce a slight decrease in expression. However, for yR7 enh, these same levels completely turn off expression. Fig. 6. Dve feeds back to control outer enh. (A-C) Dashed white circles indicate outer PRs and R8s; solid white circles indicate R7s. Expression analysis was conducted on mid-pupal retinas. In schematics: green circles indicate GFP expression; black circles indicate no GFP expression; red spots indicate RFP expression. (B,C) Solid gray line represents boundary between dve mutant clones (indicated by absence of RFP) and wild-type clones (indicated by presence of RFP). (A) Expression of outer enh is lost when Dve is ectopically expressed in all PRs. (B,C) Autoregulatory feedback: in outer PRs, outer enh is upregulated in dve mutant clones compared with wild-type clones.

Fig. 7. Combinatorial transcription factor input, feedback and redundancy govern dve expression. (A) In yR7 cells, yR7 enh is activated by Otd and Ss, while outer enh is repressed by Sal. (B) In outer PRs, Otd activates outer enh. (C) In yR7 cells, yR7 enh induces Dve expression that negatively feeds back onto yR7 enh to maintain homeostatic levels. (D) In outer PRs, outer enh induces Dve expression that negatively feeds back onto outer enh to maintain homeostatic levels. (E) In wild-type outer PRs, outer enh induces Dve expression that negatively feeds back onto outer enh to completely repress expression. (F) Upon loss of outer enh function, yR7 enh is derepressed and drives expression in outer PRs.
expression in outer PRs. The difference may lie in activation by Otd: outer enh contains four Otd/Dve sites, whereas yR7 enh contains two (Figs 3K, 5G). As these sites mediate both activation by Otd and repression by Dve, cooperative action by the four sites in outer enh may drive stronger expression and prevent repression. Generally, the primary enhancer is expressed and must be significantly less susceptible to repression than the dark shadow enhancer, which is off.

Expression of Dve in outer PRs is seen in the mosquitoes Anopheles gambiae and Aedes aegypti (Johnston et al., 2011), suggesting a conserved role in Rh regulation that has been maintained over 250 million years of evolution. However, expression of Rhs in R7s of mosquito species is regionalized in contrast to the stochastic pattern in Drosophila (Hu and Castelli-Gair, 1999), suggesting that different mechanisms have arisen to regulate Dve and Rh expression in R7s. Dark shadow enhancers may be an ancestral mechanism to ensure gene expression despite evolutionary changes. Furthermore, they may allow the evolution of new functions such as the expression of yR7 enh in R7s.

Dark shadow enhancers appear to provide robustness to gene expression and may act as additional mechanisms of canalization (i.e. the ability for individuals in a population to produce similar phenotypes regardless of environmental or genetic perturbation) (Waddington, 1942). Buffering of gene expression occurs at the levels of cis-regulatory logic (Dunipace et al., 2013; Frankel et al., 2010; Hong et al., 2008; Staller et al., 2015; Wunderlich et al., 2015) and gene networks (Cassidy et al., 2013; Lott et al., 2007; Manu et al., 2009). Dark shadow enhancers are an interesting integration of these mechanisms, whereby a primary enhancer induces expression of a factor that feeds back to repress a dark shadow enhancer. When expression from the primary enhancer is perturbed, this feedback is broken and the dark shadow enhancer becomes active. Thus, dark shadow enhancers are poised as backup mechanisms for proper gene regulation. As our understanding of complex multi-enhancer systems increases, it will be interesting to see the generality of dark shadow enhancers.

In conclusion, our studies show how two enhancers integrate combinatorial transcription factor input, negative autoregulation and redundancy in cis-regulatory elements to determine robust levels of gene expression in photoreceptor neurons. These mechanisms likely play roles in the establishment and maintenance of gene expression levels in other neuronal subtypes.

MATERIALS AND METHODS

Generating dve enh-GFP constructs
Fragments (3-6kb; Fig. 2A) were cloned into GFP reporter constructs and injected into flies. Transgenic flies were isolated and stocks were generated (see supplementary Materials and Methods and Table S1).

Drosophila strains
Flies were raised on standard cornmeal medium and grown at room temperature (25°C) (see supplementary Materials and Methods and Tables S2-S4 for complete descriptions of Drosophila genotypes).

CRISPR-generated deletions
dve-B promoter, outer enh and yR7 enh deletions were generated using CRISPR (see supplementary Materials and Methods and Table S5 for further details).

Otd/Dve binding site knockout
Otd/Dve binding site knockouts for outer enh and yR7 enh were generated using site-directed mutagenesis (see supplementary Materials and Methods for further details).

Electrophoretic mobility shift assay
Binding assays were performed as described previously (Johnston et al., 2011; Li-Kroeger et al., 2008) (see supplementary Materials and Methods for further details).

Antibodies
Antibodies and dilutions used were as follows: mouse anti-prospero (1:10, DSDB), rat anti-Elav (1:50, DSDB), sheep anti-GFP (1:500, Bio-Rad, 4745-1051), mouse anti-Rh3 (1:100; a gift from S. Britt, University of Colorado, Boulder, CO, USA), rabbit anti-Rh4 (1:100; a gift from C. Zuker, Columbia University, New York, USA), mouse anti-Rh5 (1:200; Tahayato et al., 2003), rabbit anti-Rh6 (1:2000; Tahayato et al., 2003), guinea pig anti-Ss (1:200; a gift from Y. N. Jan, University of California, San Francisco, CA, USA) and rabbit anti-Dve (1:500; Nakagoshi et al., 1998). All secondary antibodies were Alexa-conjugated (1:400; Molecular Probes).

Retina dissection and immunohistochemistry
Retinas were dissected and stained as described previously (Hsiao et al., 2012) (see supplementary Materials and Methods for further details).

Quantification
Fluorescence intensity of nuclear GFP expression of single retinas was quantified using the ImageJ processing program. A small region in the center of each nucleus was selected for fluorescence intensity measurement. Images were taken under subsaturating conditions and comparisons of GFP intensity were drawn between cells of the same retina. Column scatterplots were generated using Graphpad Prism.

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Competing interests
The authors declare no competing or financial interests.

Author contributions

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References
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