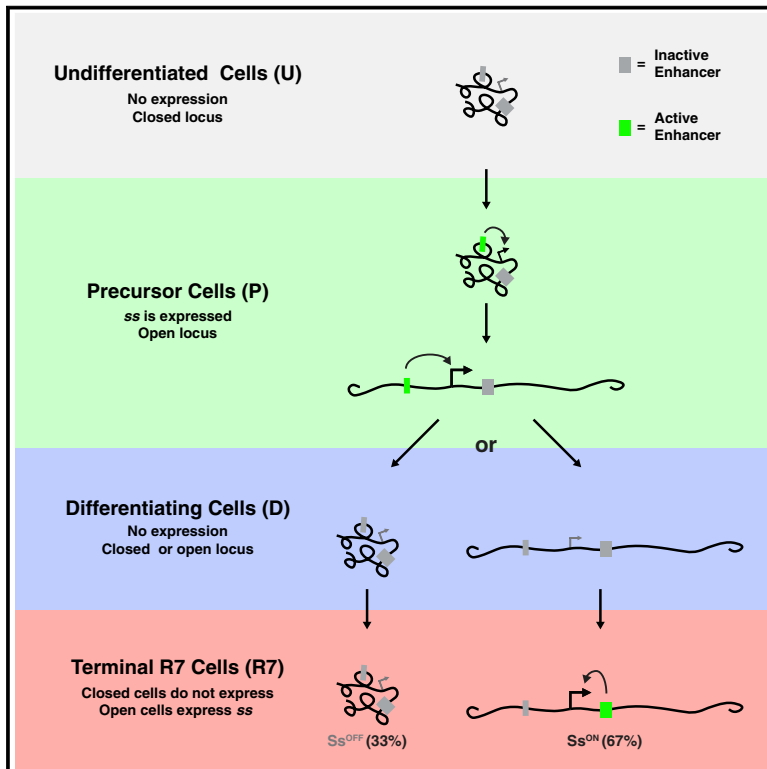


Developmental Cell

Temporally dynamic antagonism between transcription and chromatin compaction controls stochastic photoreceptor specification in flies

Graphical abstract



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In brief

Cells sometimes randomly choose between fates during development. In *Drosophila*, a random mosaic of photoreceptor subtypes is determined by the *spineless* gene. Voortman et al. find that *spineless* is regulated by a dynamic interplay between transcription and chromatin compaction during fly eye development.

Highlights

- *spineless* is off, and the locus is in a compact state in undifferentiated cells
- The *early enhancer* drives expression, and the locus opens
- Expression ceases, and the locus recompacts or remains open
- Repression in a subset of R7s limits expression driven by the *late enhancer*

Article

Temporally dynamic antagonism between transcription and chromatin compaction controls stochastic photoreceptor specification in flies

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SUMMARY

Stochastic mechanisms diversify cell fates during development. How cells randomly choose between two or more fates remains poorly understood. In the *Drosophila* eye, the random mosaic of two R7 photoreceptor subtypes is determined by expression of the transcription factor Spineless (Ss). We investigated how *cis*-regulatory elements and *trans* factors regulate nascent transcriptional activity and chromatin compaction at the *ss* gene locus during R7 development. The *ss* locus is in a compact state in undifferentiated cells. An early enhancer drives transcription in all R7 precursors, and the locus opens. In differentiating cells, transcription ceases and the *ss* locus stochastically remains open or compacts. In Ss^{ON} R7s, *ss* is open and competent for activation by a late enhancer, whereas in Ss^{OFF} R7s, *ss* is compact, and repression prevents expression. Our results suggest that a temporally dynamic antagonism, in which transcription drives large-scale decompaction and then compaction represses transcription, controls stochastic fate specification.

INTRODUCTION

Cell fate specification is controlled by lineage, signaling, and stochastic regulatory inputs, leading to highly precise developmental outcomes (Petkova et al., 2019). Stochastic mechanisms promote diversity in populations of photoreceptors (PRs), olfactory neurons, motor neurons, and immune cells (Alqadah et al., 2016; Bell et al., 2007; Dasen et al., 2003, 2005; Duffy et al., 2012; Johnston and Desplan, 2010; Miyamichi et al., 2005; Reissler et al., 1993; Vassar et al., 1993). Despite the importance of stochastic cell fate specification, how cells randomly choose between fates is poorly understood.

Stochastic cell fate specification is best understood in prokaryotes. One well-characterized example is the bet-hedging mechanism utilized by *Bacillus subtilis*. To minimize losses in a changing environment, populations of genetically identical bacteria maintain a subpopulation of cells that are competent for DNA uptake (Dubnau, 1999; Turgay et al., 1998; Nester and Stocker, 1963). The transient and random transition into the competent fate is controlled by expression of the transcriptional regulator ComK (Turgay et al., 1997, 1998). Though most cells maintain low expression of ComK, a subset will experience a pulse of ComK expression that exceeds a threshold and induces a transition to the competent fate (Maamar et al., 2007; Süel et al.,

2006). A similar mechanism occurs in the HIV life cycle, where transcription of the regulatory factor *trans*-activator of transcription (Tat) determines the switch from proviral latency to active replication (Hendy et al., 2017; Weinberger et al., 2008). Thus, stochastic cell fate specification often requires a pulse of expression of a critical regulator that determines a fate decision.

In addition to transcriptional dynamics, chromatin-mediated repression is a key mechanism mediating stochastic fate specification. In mice, each olfactory sensory neuron (OSN) expresses only one olfactory receptor (OR) gene from a battery of ~1,300 possibilities (Buck and Axel, 1991; Chess et al., 1994; Godfrey et al., 2004). Despite residing in numerous clusters across many chromosomes, all ~1,300 OR genes are repressed and coalesce into heterochromatic foci within the nucleus prior to OR selection (Clowney et al., 2012; Magklara et al., 2011; Sullivan et al., 1996; Zhang and Firestein, 2002). In mutants that impact chromatin modifications and nuclear organization, co-expression of multiple ORs is observed (Clowney et al., 2012). While the mechanism of selection remains elusive, a single OR allele escapes the repressive heterochromatic environment and is expressed in each OSN (Armelin-Correa et al., 2014; Lyons et al., 2013). Thus, chromatin-mediated silencing and selective de-silencing are paramount for the stochastic expression of a single OR gene.

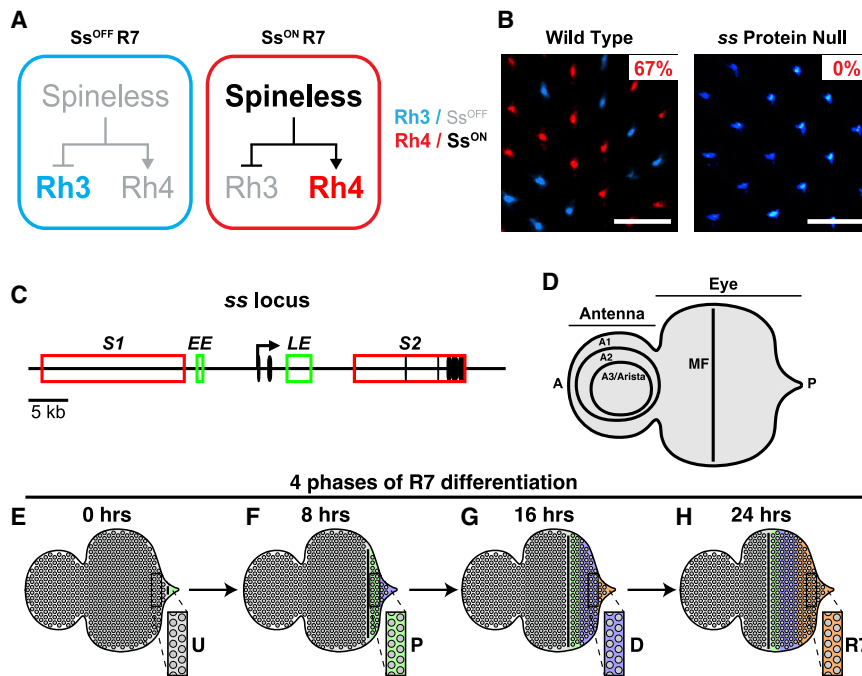


Figure 1. ss controls R7 subtype specification

(A) R7 subtype specification. Expression of Ss promotes the Rh4-expressing R7 fate. Absence of Ss yields the Rh3-expressing R7 fate.

(B) Wild-type retinas contain 33% Rh3/Ss^{OFF} R7s and 67% Rh4/Ss^{ON} R7s in a random pattern (left). ss protein null mutants contain only Rh3/Ss^{OFF} R7s and no Rh4/Ss^{ON} R7s (right). Scale bars, 20 μ m.

(C) ss gene locus. Black oval, exon; black arrow, promoter; red rectangle, silencer; green rectangle, enhancer; S1, silencer 1; S2, silencer 2; EE, early enhancer; LE, late enhancer.

(D) Schematized eye-antennal imaginal disc. Antenna is subdivided into the A1, A2, and A3/arista. A, anterior; P, posterior; MF, morphogenetic furrow.

(E–H) Schematized depiction of R7 maturation. Insets illustrate how cells proceed through development over time. Gray, undifferentiated cells/U; green, precursors/P; blue, differentiating cells/D; orange, R7.

The random mosaic of R7 PRs in the fly eye provides a paradigm to study the integration of transcription and chromatin-mediated repression in stochastic cell fate specification. In the fly eye, stochastic expression of the PAS-bHLH transcription factor Spineless (Ss) establishes the random pattern of two R7 subtypes across the retina. Ss^{ON} R7s express Rhodopsin 4 (Rh4), while Ss^{OFF} R7s express Rhodopsin 3 (Rh3) (Figures 1A and 1B; Bell et al., 2007; Duncan et al., 1998; Johnston and Desplan, 2014; Montell et al., 1987; Wernert et al., 2006). In wild-type flies, each R7 has a 67% chance of adopting the Ss^{ON} R7 fate and a 33% chance of assuming the Ss^{OFF} R7 fate, yielding a consistent ratio yet unique, random pattern of R7 subtypes across eyes (Figure 1B). In ss protein null mutants, all R7s express Rh3 (Figure 1B). The stochastic ON/OFF ss expression is controlled by an enhancer (late enhancer, LE) that drives expression in all R7s and silencers that limit expression to a subset of R7s (Figure 1C).

Here, we describe a mechanism that controls stochastic R7 subtype specification. Initially, the ss locus is compact in all undifferentiated cells. An early enhancer (EE) drives ss expression and the ss locus opens in all R7 precursors during larval development. Expression ceases and the ss locus randomly compacts or remains open. In R7s in which ss remains open, the LE drives ss expression and Ss^{ON} R7 fate. In R7s with compact chromatin, repression prevents expression driven by the LE, yielding the Ss^{OFF} R7 fate. Our data suggest that stochastic fate specification is controlled by the dynamic, intertwined relationship of transcription and chromatin: transcription opens chromatin then chromatin compaction represses transcription. We find that transcription is a source of stochasticity as modulating early transcription in precursors alters the proportions of alternative R7 fates.

RESULTS

ss expression is dynamic in developing R7s

PR identity, including R7 subtype, is specified during larval development in the eye-antennal imaginal disc (Figure 1D). Retinal differentiation begins at the posterior end and progresses in a wave anteriorly. An indentation called the morphogenetic furrow (MF) appears at the posterior end (Figure S1A). The MF progresses in a developmental wave from posterior to anterior (Figure S1). Behind the MF, PRs differentiate in a stereotypical progression: R8, R2/R5, R3/R4, R1/R6, and finally R7 (Figures S1B–S1F). As the eye develops in this spatiotemporal manner, individual discs provide information on all stages of PR specification, with undifferentiated cells in the anterior and the most differentiated cells in the posterior (Figure S1F; Gallagher et al., 2022; Ready et al., 1976; Tomlinson and Ready, 1987a, 1987b; Treisman, 2013; Wolff and Ready, 1991).

We defined four phases that R7s proceed through during development, including undifferentiated (U), precursor (P), differentiating (D), and differentiated R7 (R7) (Figures 1E–1H). In individual discs, we visualized all four phases (Figure 1H). Undifferentiated cells were anterior to the MF (Figures 1E–1H). Posterior to the MF, precursors were located at 0–10 μ m, differentiating cells were located at 10–30 μ m, and R7s were located at >30 μ m (Figures 1F–1H). In Figures 1E–1H and subsequent figures, we diagram only undifferentiated, precursor, differentiating, and R7 cells in the eye and developing cells in the antenna for simplicity.

To characterize ss expression, we performed nascent RNA fluorescence *in situ* hybridization (RNA FISH). We generated oligoprobes covering the entire ss transcript, including introns and exons (Figure S2A), and performed single-molecule RNA FISH (smFISH) (Beliveau et al., 2012; Little et al., 2013). This

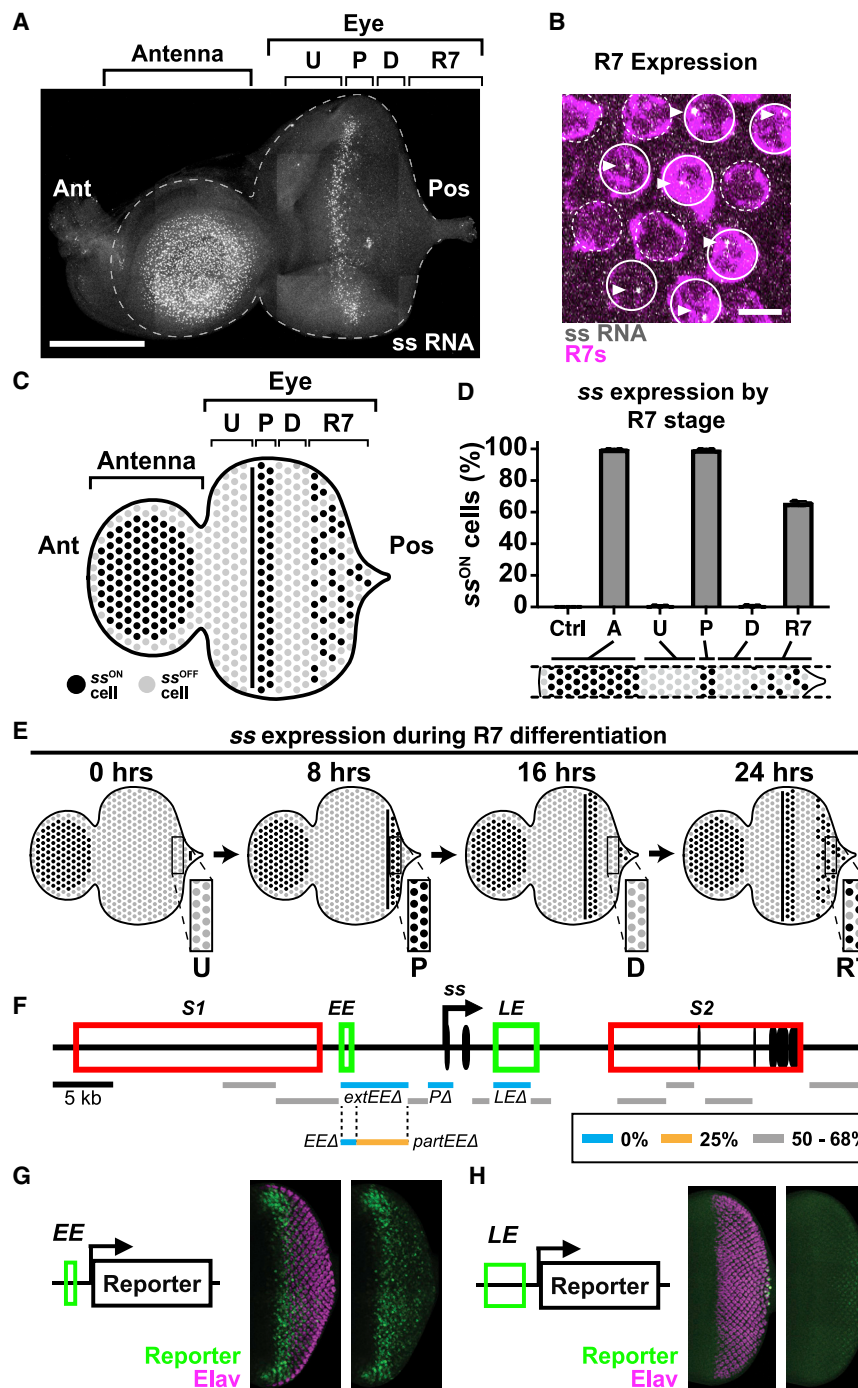


Figure 2. Two temporally distinct enhancers drive ss expression

(A–E) Ctrl, peripodial membrane; A, antennal cells; U, undifferentiated cells; P, precursors; D, differentiating cells; R7, R7s. Ant, anterior; Pos, posterior.

(A) ss RNA is expressed in antennal cells, precursors, and R7s. Gray, ss RNA. Scale bars, 100 μ m.

(B) Nascent ss RNA transcripts in a subset of R7s distinguished by *sev>Gal4, UAS>GFP*. Magenta, R7 reporter; gray, ss RNA; solid circles, Ss^{ON} R7s, dashed circles, Ss^{OFF} R7s. Scale bars, 5 μ m.

(C) Schematized eye-antennal imaginal disc.

(D) % cells expressing ss. Error bars denote standard deviation from the mean.

(E) Schematized ss expression across time. Insets illustrate ss expression dynamics.

(F) ss gene locus and CRISPR deletion screen. Black oval, exon; black arrow, promoter; red rectangle, silencer; green rectangle, enhancer; blue line, 0% Ss^{ON} R7s; orange line, 25% Ss^{ON} R7s; gray line, 50%–68% Ss^{ON} R7s; S1, silencer 1; S2, silencer 2; EE, early enhancer; LE, late enhancer.

(G and H) Green, reporter; magenta, Elav (neurons).

(G) The EE reporter is expressed in precursors.

(H) The LE reporter is expressed in R7s.

In the eye, ss is differentially expressed during R7 specification. ss is not expressed in undifferentiated cells, ss is strongly expressed in all precursors, ss is not expressed in differentiating cells, and ss is expressed in a subset of R7s (Figures 2A–2E and S2C). We distinguished R7s from other PRs using an R7 reporter line (*R7 > GFP*), allowing quantification of ss^{ON/OFF} R7s. The ss^{ON/OFF} ratio in larval R7s is similar to the ss^{ON/OFF} ratio in adult R7s (Figures 2D and S2C), consistent with this decision being made in larvae and maintained throughout the lifetime of the organism (Johnston and Desplan, 2014). As expression in an individual disc represents different temporal phases of R7 development, we conclude that ss expression is dynamic as R7s develop: off in the undifferentiated cell phase, on in the precursor phase, off in the differentiating cell phase, and

finally, on in a subset of R7s in the differentiated R7 phase (Figures 2A–2E).

strategy yielded single bright fluorescent punctae in ss-expressing nuclei, indicating sites of nascent transcription (Figures 2A and 2B). Our observation of one puncta per nucleus is consistent with chromosome pairing in close proximity in somatic cells of *Drosophila* (Stevens, 1908). This approach enabled quantification of ss transcription in each developmental context. ss is strongly expressed in all cells of the central antennal region, serving as a positive control (Figures 2A, 2C, and 2D, A). ss is not expressed in the peripodial membrane that overlies the eye-antennal disc, acting as a negative control (Figures 2D and S2B, Ctrl).

finally, on in a subset of R7s in the differentiated R7 phase (Figures 2A–2E).

Two temporally distinct enhancers drive ss expression in the developing eye

To identify *cis*-regulatory elements that regulate stochastic ON/OFF expression of ss in R7s, we used CRISPR to make a series of 1–5-kb deletions in the ss locus (Figure 2F; Table S1). As Rhodopsin expression faithfully reports Ss expression state in adults (Rh4 = Ss^{ON}, Rh3 = Ss^{OFF}), we examined Rh3 and Rh4

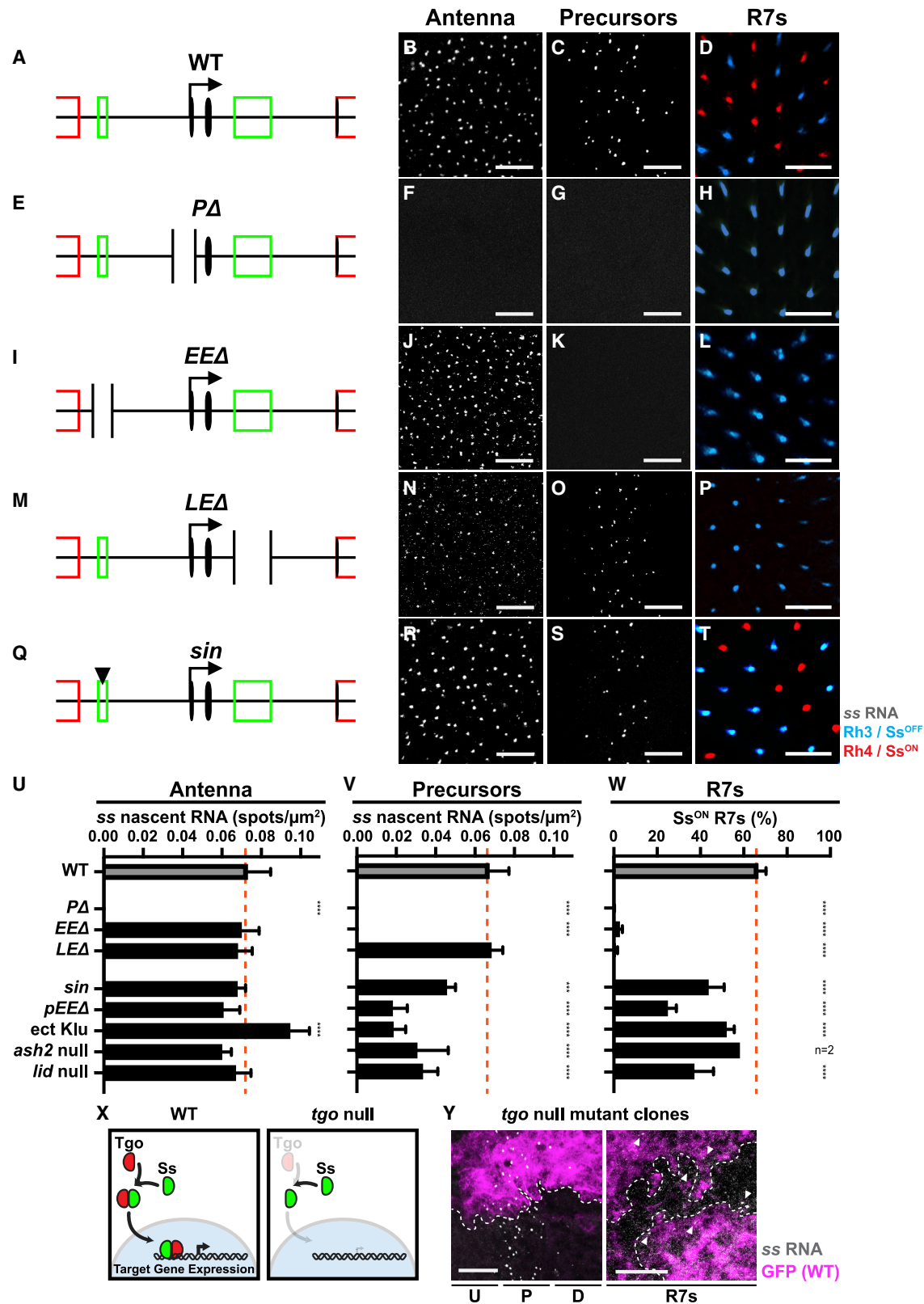


Figure 3. Decreasing early *ss* expression decreases % *Ss*^{ON} R7s

(A, E, I, M, and Q) Truncated schematized *ss* locus.

(B, F, J, N, and R) *ss* RNA in the antenna. Gray, *ss* RNA. Scale bars, 10 μ m.

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expression and determined the proportions of R7 subtypes. We identified three elements that are required for ss expression in R7s, including the promoter (*P*), a 5.4 kb upstream element (*extEE*), and the previously identified *LE* (Figure 2F; Table S1; Johnston and Desplan, 2014). Deletion of these regions reduced the proportion of Ss^{ON} R7s to 0% (Figure 2F; Table S1). Thus, these *cis*-regulatory regions are required for normal ss expression.

We conducted additional partial deletions of the *extEE* region to determine a minimal *cis*-regulatory region required for ss expression. Deletion of the 1.3-kb *EE* region caused a dramatic decrease of Ss^{ON} R7s to 0% (Figure 2F; Table S1), while deletion of the neighboring 4.1 kb *partEE* caused a partial reduction of Ss^{ON} R7s to 25% (Figure 2F; Table S1). As *EE* was strictly required for ss expression, we interrogated this region further.

To assess the spatiotemporal activity of *EE* and *LE* activities, we generated reporter constructs and examined expression in larval eye-antennal discs. The *EE* drove expression in precursors similar to ss RNA expression (Figure 2G). In contrast, the *LE* drove expression in all R7s (Figure 2H). We did not observe expression in the antenna for either construct, suggesting that *EE* and *LE* are eye-specific enhancers for ss. Thus, *EE* and *LE* are sufficient to drive expression in precursors and R7s respectively.

As chromatin accessibility is associated with enhancer activity, ATAC-seq can predict candidate enhancers (Buenrostro et al., 2015). We analyzed published scATAC-seq datasets (Bravo González-Blas et al., 2020). For antennal cells that express ss, accessibility peaks were observed at the promoter, but not at *EE* or *LE* (Figure S2D). For precursors that express ss, peaks occurred at the *EE* and *promoter*, but not *LE* (Figure S2D). For all PRs, of which only a subset of R7s express ss, peaks were observed at the *LE* and *promoter* and were significantly reduced for the *EE* (Figure S2D). As a small peak remains at the *EE* in R7s, some residual chromatin accessibility may remain at this later time point. Alternatively, some cells may have been incorrectly clustered into this cell type. These observations support roles for the *EE* and *LE* as enhancers that drive expression during distinct temporal phases of R7 development: *EE* drives early expression in precursors and *LE* drives late expression in R7s.

Early ss expression in precursors is required for ss expression in R7s

To test how the *EE* and *LE* regulate ss expression during PR development, we observed ss expression in mutant conditions. In the fly eye, automated identification and assignment of

nascent spots to individual cells is challenging in 3D and not necessary to describe the changes in expression observed here. Therefore, to quantify ss-expressing cells in the eye-antennal disc, we measured the density of nascent RNA spots per unit area (μm^2) (Figures S3A–S3C). To control for changes in tissue morphology, we measured the density of cells in the antenna and precursors (Figures S3D and S3E). We assessed Rh4/Ss^{ON} and Rh3/Ss^{OFF} in R7s in adult retinas (Figures 3D and 3W). ss expression in antennal cells in eye-antennal discs served as a positive internal control (Figures 3B and 3U). Promoter deletion (*PΔ*) mutants acted as a negative control, exhibiting a complete loss of ss expression (Figures 3E–3H and 3U–3W).

EEΔ mutants lost ss expression in precursors and Rh4/Ss^{ON} in R7s, while ss expression was maintained in antennal cells (Figures 3I–3L and 3U–3W). *LEΔ* mutants displayed a complete loss of Rh4/Ss^{ON} expression in R7s but showed normal ss expression in precursors and antennal cells (Figures 3M–3P and 3U–3W). *PΔ*, *EEΔ*, and *LEΔ* mutants displayed no differences in antennal or precursor cell densities (Figures S3D and S3E). Together, the *EE* is required for ss expression in precursors and R7s, whereas the *LE* is required for ss expression in R7s.

ss expression does not require Ss protein feedback in precursors or R7s

Early expression often affects later expression from the same gene locus through protein feedback (Maamar et al., 2007; Süel et al., 2006). In the fly eye, no detectable Ss protein is observed in precursors (Johnston and Desplan, 2014). Nevertheless, extremely low levels of Ss protein could trigger regulatory feedback. Ss protein requires heterodimerization with another PAS-bHLH transcription factor, Tango (Tgo) to enter the nucleus and regulate gene expression (Emmons et al., 1999; Thanawala et al., 2013; Ward et al., 1998). To test whether Ss/Tgo feedback activity affects early ss expression, we generated *tgo* null mutant clones and observed no effect on ss transcription in precursors or R7s (Figures 3X, 3Y, and S3I). This result suggests that (1) ss regulation in the eye does not require Ss protein feedback, consistent with our previous findings (Johnston and Desplan, 2014), and (2) the early transcription of ss activates late expression by a Ss/Tgo-protein independent mechanism.

Decreasing early ss expression decreases the proportion of Ss^{ON} R7s

The *EE* is required for specification of Ss^{ON} R7s, as knocking out the *EE* caused a complete loss of ss expression in R7s

(C, G, K, O, and S) ss RNA in precursors. Gray, ss RNA. Scale bars, 10 μm .

(D, H, L, P, and T) Adult Rh3/Ss^{OFF} and Rh4/Ss^{ON} expression in R7s. Scale bars, 20 μm .

(A–D) WT.

(E–H) *PΔ*.

(I–L) *EEΔ*.

(M–P) *LEΔ*.

(Q–T) Animals with *sin* variant.

(U–W) Orange line, mean WT expression. Error bars denote standard deviation from the mean. n.s. denotes $p > 0.05$; *** $p < 0.0005$; **** $p < 0.0001$.

(U) Quantification of ss in antennal cells.

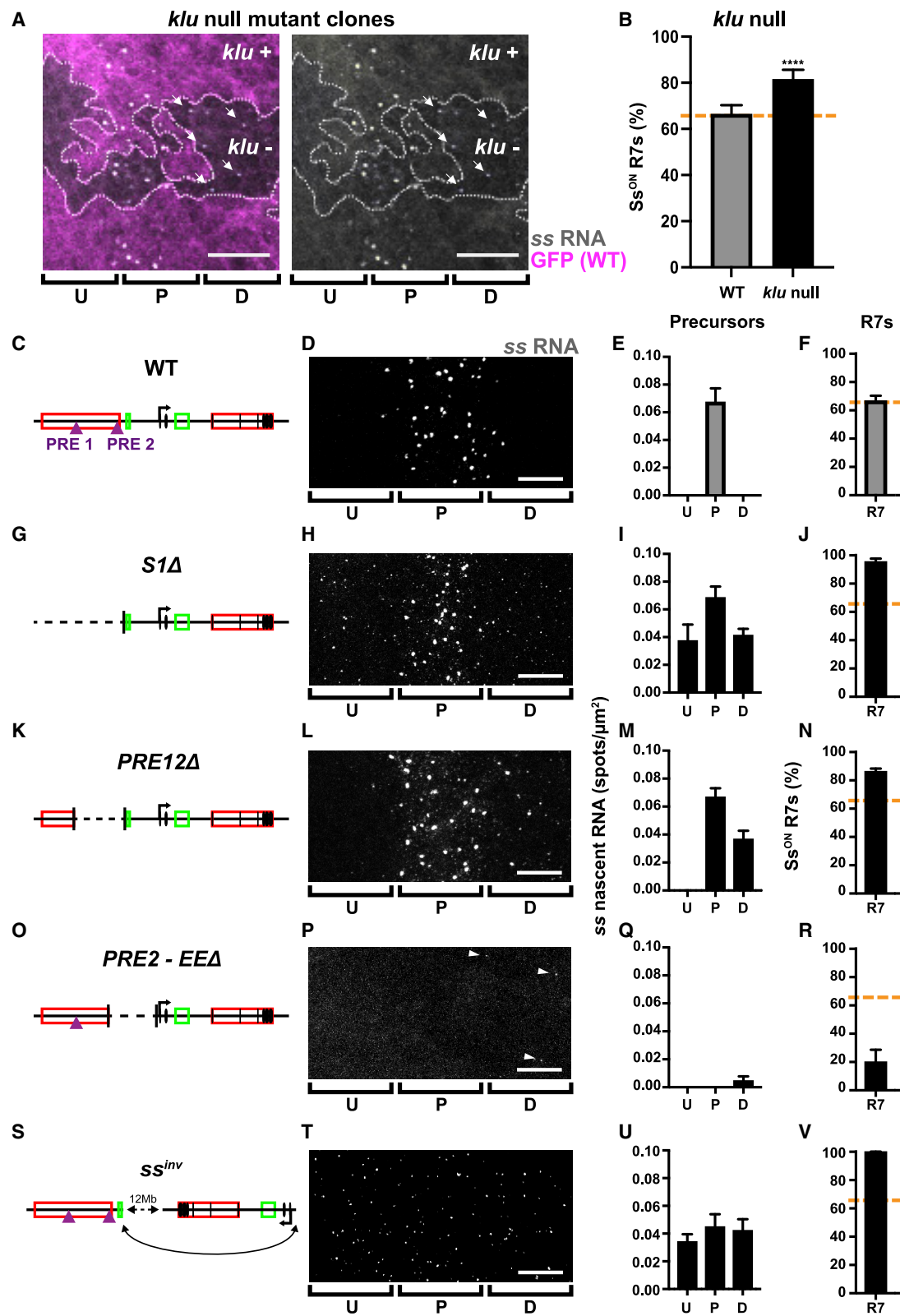
(V) Quantification of ss in precursors.

(W) Quantification of % Ss^{ON} R7s.

(X) Ss/Tgo mechanism in WT and breakdown in *tgo* mutants.

(Y) ss RNA in precursors and a subset of R7s in *tgo* null mutant clones. Dashed line, clone boundary.

GFP– = *tgo* null mutant; GFP+ = wild type. Gray, ss RNA; magenta, GFP; dashed line, clone boundary. Scale bars, 10 μm .



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(Figures 3I–3L and 3U–3W). We hypothesized that reducing activation by the *EE* would decrease the number of ss-expressing precursors and the proportion of Ss^{ON} R7s. Because 100% of precursors express ss in wild-type flies, we could identify changes in early and late expression as changes in the density of expression in precursors and the ratio of Ss^{ON/OFF} R7s in adults (Figures 2D and S2C).

The *EE* contains a binding site for the transcriptional repressor, Klumpfuss (Klu), which is expressed during all stages of R7 subtype specification (Figure S3F; Anderson et al., 2017). A single base insertion (“*sin*”) within the *EE* increases the binding affinity of Klu (Anderson et al., 2017). Flies with *sin* displayed a reduction in the number of ss-expressing precursors and a decrease in the proportion of Ss^{ON} R7s (45% Ss^{ON}) (Figures 3Q, 3S, 3T, 3V, and 3W). Flies with *sin* had no change in ss expression in the antenna (Figures 3R and 3U). Flies with *sin* displayed no differences in antennal or precursor cell densities (Figures S3D and S3E). Ectopic expression of Klu reduced ss expression in precursors and the ratio of Ss^{ON} R7s (51.8% Ss^{ON}) (Figures 3V, 3W, S3Q, and S3R; Anderson et al., 2017) with no effect on precursor cell density (Figure S3E). Ectopic expression of Klu in precursors caused an increase of ss expression in the antenna (Figures 3U and S3P) and an increase in antennal cell density (Figure S3D), consistent with differential regulation of ss by Klu across tissues through different enhancers (Klein and Campos-Ortega, 1997; Yang et al., 1997). Additionally, a partial deletion of the *EE* (*pEEΔ*) resulted in a reduction in expression in precursors and a reduction of Ss^{ON} R7s to 25% (Figures 3U–3W and S3M–S3O). *pEEΔ* mutants displayed no differences in antennal or precursor cell densities (Figures S3D and S3E). This deletion removes the sequence abutting the *EE* and may disrupt the binding of other *trans*-acting factors to the *EE*. These data suggest that decreasing *EE* activity by genetically altering *cis* or *trans* inputs reduces ss expression in precursors and leads to a reduction in Ss^{ON} R7s.

To identify regulators of R7 subtype specification, we screened flies with mutations or RNAi knockdowns in genes encoding chromatin modifiers for changes in the ratio of Ss^{ON} and Ss^{OFF} R7s (Table S2). Reducing activity of two genes encoding chromatin modifiers, *ash2* and *lid*, caused significant loss of Ss^{ON} R7s. Knockdown of the trithorax group gene *ash2* (Adamson and Shearn, 1996; Papoulas et al., 1998) caused a decrease in Ss^{ON} R7s in two independent RNAi lines (Table S2). *ash2*¹ null mutants displayed a reduction in Ss^{ON} R7s (38.4% Ss^{ON})

(Figures 3W and S3U). *ash2*¹ null mutants displayed a cell-autonomous decrease in ss expression in precursors (Figures 3V, S3H, and S3T), and no change in ss expression in antennal cells (Figures 3U and S3S). Similarly, a null mutation in the histone demethylase gene *lid* (Eissenberg et al., 2007; Secombe et al., 2007) caused a reduction in ss expression in precursors and the proportion of Ss^{ON} R7s (36.8% Ss^{ON}) but had no effect on ss expression in antennal cells (Figures 3U–3W and S3V–S3X). Though *ash2*¹ and *lid* mutants displayed a decrease in precursor cell densities (Figure S3E), the proportion of cells expressing ss was reduced after normalization to cell density (Figure S3Y). These data implicate a role for chromatin modifiers in ss regulation and suggest that decreasing ss expression in precursors decreases the proportion of Ss^{ON} R7s.

To evaluate the relationship of ss expression in precursors to the ratio of Ss^{ON} R7s, we normalized the densities of ss-expressing precursors and the ratios of Ss^{ON} R7s for each genotype to wild type. For expression in precursors and R7s respectively, *sin* mutants had 67.7% and 65.4%, *pEEΔ* mutants had 26.7% and 37.1%, *ash2* null mutants had 83.2% and 87.0%, and *lid* null mutants had 58.3% and 55.4% (Figure S3Y). These mutants had highly similar proportional changes between precursors and R7s (Figure S3Y). In contrast, flies with ectopic expression of Klu had 27.3% normalized expression in precursors and 77.9% Ss^{ON} R7s (Figure S3Y), suggesting multiple roles for Klu in this process or differences in levels and/or timing caused by the transgenic overexpression. Together, these data suggest that expression early in precursors is required for expression late in R7s.

Derepression of early ss expression increases the proportion of Ss^{ON} R7s

We next investigated how derepression of ss affected R7 subtype specification. We hypothesized that mutant genotypes with an increase in Ss^{ON} R7s will have an altered expression pattern earlier in development. For these experiments, we examined ss expression in undifferentiated cells (ss^{OFF}), precursors (ss^{ON}), and differentiating cells (ss^{OFF}) in larval eye-antennal discs as well as R7s (mix of ss^{ON} and ss^{OFF}) in adult retinas (Figures 4C–4F). As all precursors express ss, we did not observe an increase in the density of expression in these cells in these mutant conditions. Rather, we observed aberrant expression earlier in undifferentiated cells and/or later in differentiating cells during R7 specification.

Figure 4. Derepressing early ss expression increases % Ss^{ON} R7s

(A–V) U, undifferentiated cells; P, precursors; D, differentiating cells; R7, R7s.

(A) ss expression in precursors is extended into differentiating cells in *klu* null mutant clones. GFP[−] = *klu* null mutant; GFP⁺ = wild type. Gray, ss RNA; magenta, GFP; dashed line, clone boundary; arrows, ss RNA in differentiating cells. Scale bars, 10 μm.

(B, F, J, N, R, and V) Orange line, mean WT ss expression. Error bars denote standard deviation from the mean.

(B) % Ss^{ON} R7s increases in *klu* null mutants, similar to previous studies (Anderson et al., 2017). ****p < 0.0001. N = 3.

(C, G, K, O, and S) Schematized ss locus.

(D, H, L, P, and T) ss RNA in undifferentiated cells, precursors, and differentiating cells. Scale bars, 10 μm.

(E, I, M, Q, and U) Quantification of expression for (D), (H), (L), (P), and (T). Error bars denote standard deviation from the mean.

(F, J, N, R, and V) %Ss^{ON} R7s.

(C–F) WT.

(G–J) S1Δ.

(K–N) PRE12Δ.

(O–R) PRE2-EEΔ.

(S–V) ss^{inv}.

Increasing the binding affinity of a Klu site or increasing Klu levels reduced early ss expression and the proportion of Ss^{ON} R7s. In contrast, *klu* null mutant clones displayed a temporal extension of ss expression beyond the precursor state into the differentiating state, when ss is not normally expressed (Figure 4A). *klu* null mutants also exhibited an increase in the proportion of Ss^{ON} R7s (82% Ss^{ON}) without changing eye morphology (Figure 4B; Anderson et al., 2017). These data suggest that Klu is a cell-autonomous off switch for ss expression and that extended expression of ss leads to an increase in the probability of Ss^{ON} R7 fate.

Repressive *silencer* elements restrict expression of ss to a subset of R7s (Johnston and Desplan, 2014). We focused on the effects of a 36.4-kb deletion of *silencer1* (*S1Δ*) (Figure 4G; Thanawala et al., 2013). Heterozygous *S1Δ/+* mutants displayed ss expression in undifferentiated cells prior to the precursor stage and in differentiating cells after the precursor stage (Figures 4H, 4I, and S4B–S4D) and an increase in the proportion of Ss^{ON} R7s (95%) (Figures 4J and S4E). *S1Δ/+* mutants showed low level ss expression in most cells of the eye-antennal disc, including the peripodial membrane, which is normally ss^{OFF} (Figure S4H). *S1Δ/+* mutants did not display changes in cell density (Figures S4F and S4G). Together, these data indicate that *silencer1* is generally required for repression of ss.

As ss expression was diminished in *ash2* and *lid* mutants, we hypothesized that chromatin is playing a role in ss repression. We examined the region deleted in *S1Δ* mutants for Polycomb response elements (PREs), DNA elements bound by Polycomb group (PcG) proteins that nucleate repressive heterochromatin (Chan et al., 1994; Paro and Hogness, 1991; Simon et al., 1993; Strutt et al., 1997). ChIP-seq showed distinct peaks for PcG proteins, suggesting that two putative PREs (PRE1 and PRE2) fall within the region deleted in the *S1Δ* mutants (Figure S4J; Celniker et al., 2009; Schwartz et al., 2006). These putative PREs correspond to peaks in scATAC-seq datasets (Figure S2D; Bravo González-Blas et al., 2020), suggesting that these are binding sites for PcG proteins. We validated the activity of PRE1 and PRE2 using pairing sensitive silencing assays (Figures S4K–S4O; Kassiss, 1994; Kassiss et al., 1991). Together, these data suggest that PRE1 and PRE2 are functional PREs.

To test the roles for PRE1 and PRE2, we generated a deletion that removed 13 kb containing both PREs (*PRE12Δ*) (Figure 4K). Hemizygous *PRE12Δ* mutants displayed a temporal extension of ss expression into differentiating cells and an increase in the ratio of Ss^{ON} R7s (86% Ss^{ON}) but did not exhibit ectopic expression in undifferentiated cells (Figures 4L–4N). These data suggest that the two PREs repress ss expression. Together, chromatin regulation at the ss locus is critical for R7 subtype specification and extending early expression increases the proportion of Ss^{ON} R7s.

Derepression restores Ss^{ON} R7 fate in *EE* mutants

Our data suggest that activation in precursors is necessary and precedes repression in differentiating cells during R7 subtype specification. To test the temporality of these steps, we examined mutants that impaired activation in precursors and repression in differentiating cells. We predicted that derepression in differentiating cells would offset loss of activation in precursors to restore Ss^{ON} R7 fate. We used imprecise P-element excision to generate an 11.8-kb mutant that deleted one PRE within

silencer1 and the *EE* (*PRE2-EEΔ*) (Figure 4O). In *PRE2-EEΔ* mutants, ss expression in precursors was completely lost (Figures 4P and 4Q), consistent with the loss of activation by the *EE*. *PRE2-EEΔ* mutants displayed low levels of ss expression in differentiating cells (Figures 4P and 4Q), consistent with a loss of repression. *PRE2-EEΔ* mutants contained 20% Ss^{ON} R7s (Figure 4R), suggesting that derepression in differentiating cells restores Ss^{ON} R7 fate in the absence of activation by the *EE*.

To further test these interactions, we examined flies with an inversion (*ss^{inv}*) that moves the ss promoter and *LE* ~12 Mb away from upstream regions, preventing regulation by the *EE* and *silencer1* (Figures 4S–4V and S4F–S4G; Thanawala et al., 2013). *ss^{inv}* mutants displayed weak ss expression in undifferentiated cells, precursors, and differentiating cells (Figures 4T and 4U), consistent with the loss of repression by *silencer1*. Strong ss expression in precursors was decreased (Figures 4T and 4U), consistent with the loss of activation by the *EE*. *ss^{inv}* mutants contained 100% Ss^{ON} R7s (Figure 4V), suggesting that derepression enables Ss^{ON} R7 fate when activation by the *EE* is lost. These data are consistent with activation driven by the *EE* in precursors before repression by *silencer* elements in differentiating cells during R7 subtype specification in wild-type conditions.

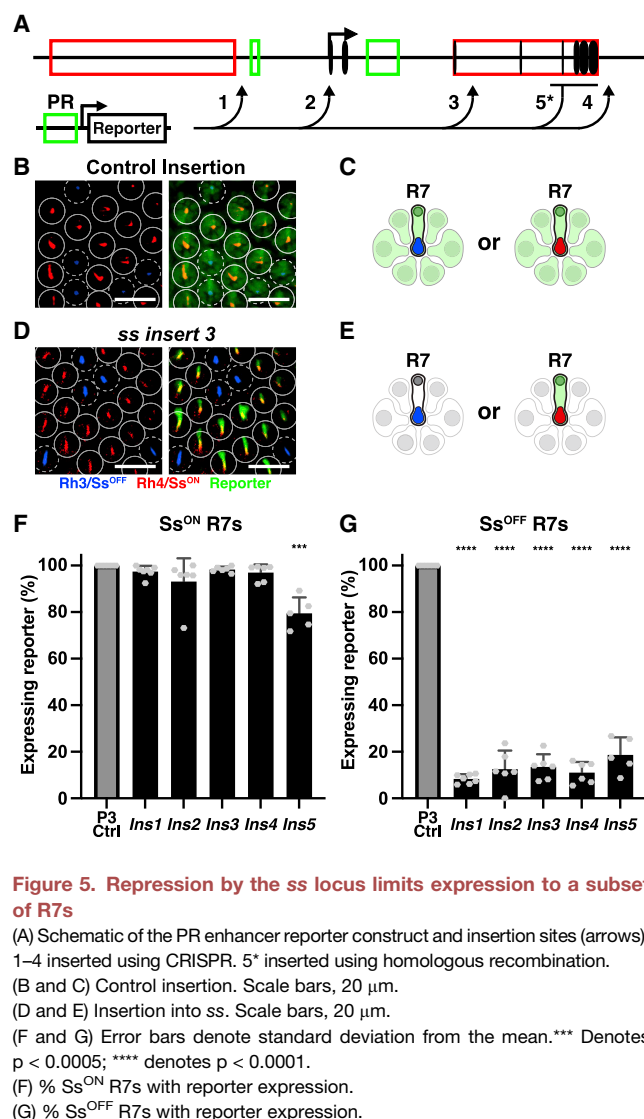
Repression limits ss expression to a subset of R7s

To test the hypothesis that repression limits ss expression to Ss^{ON} R7s, we developed a “repression reporter” strategy. A broad PR enhancer reporter (*[3xP3] > RFP*) drives expression in all PRs of the adult retina when inserted into a control locus on the X chromosome (Figures 5B, 5C, 5F, 5G, and S5A, P3 Ctrl) (Bischof et al., 2007). We used CRISPR to insert this reporter into different locations in the endogenous ss locus (Figure 5A, inserts 1–4). We hypothesized that, if the local chromatin environment at the ss locus was sufficient to repress expression, expression of the reporter would be limited to Ss^{ON} R7s. If the ss locus was not sufficient to repress expression, the reporter would be expressed in all PRs, including all R7s.

When inserted into four different locations in the ss locus, the reporter transgene was nearly perfectly repressed in Ss^{OFF} R7s and expressed in Ss^{ON} R7s in adult retinas (Figures 5D and 5E). The reporter was expressed in >93% of Ss^{ON} R7s and in <14% of Ss^{OFF} R7s (Figures 5F, 5G, and S5A). With the exception of *ins2*, inserted into the 5' UTR of ss, the reporter lines did not significantly affect the ratio of Ss^{ON/OFF} R7s (Figure S5B). These data suggest that repression at the ss locus limits expression to Ss^{ON} R7s.

When inserted into control loci, the reporter drives expression in the motion-detecting outer PRs (R1–6) (Figures 5B and 5C). When inserted into the ss locus, the reporter was not expressed in outer PRs (Figures 5D and 5E), suggesting that the ss locus represses expression in these cells.

In control insertion lines, the reporter drives expression in color-detecting R8 PRs. When inserted into the ss locus, the reporter remained expressed in a subset of R8s (Figure S5C). In these lines, Ss was expressed in the same subset of R8s (Figure S5C). These observations suggest that the broad PR enhancer reporter, inserted into the ss locus, was sufficient to overcome repression and ectopically drive ss and reporter expression in R8s.



Regulation of the “repression reporter” may be specific to this enhancer reporter inserted by CRISPR. We previously used homologous recombination to replace the last four exons of ss with a different broad PR enhancer reporter (*GMR > GFP*, Figure 5A, insert 5) (Thanawala et al., 2013). For this reporter, we examined $ss^{Ins5/+}$ heterozygous flies. Expression of this reporter was generally expressed in Ss^{ON} R7s (79.5% co-expressing), repressed in Ss^{OFF} R7s (18.7% expressing), repressed in outer PRs, and expressed in R8s (Figures 5F, 5G, and S5A).

Insertion of two different types of broad PR enhancer reporters by two different methods across five locations in the ss gene locus resulted in repression of the reporter in Ss^{OFF} R7s and expression in Ss^{ON} R7s. These data indicate that the ss gene locus represses expression and suggests a role for the local chromatin environment in repression.

Visualizing chromatin compaction at the ss locus

The repression reporter strategy showed that the ss locus restricts expression to a subset of R7s, likely through chromatin

remodeling of the ss locus. Additionally, two silencer elements are required for proper ss expression, consistent with a role for long range repressive interactions possibly through chromatin compaction.

We sought to characterize the compaction state of the ss gene locus during R7 subtype specification. The heterogeneity of cell types in the larval fly eye and limiting quantities of cells impede cell-type-specific analyses through ChIP-seq and ATAC-seq approaches. To examine chromatin compaction with single-cell resolution in intact tissue, we developed a 3-color DNA FISH strategy. We labeled a 50-kb upstream region, a 65-kb region encompassing ss, and a 50-kb downstream region with different fluorescently labeled probes (Figure 6A). We identified the center of the spheroid for each region and measured the 3D distance from the upstream region to the ss region (d_1) and from the ss region to the downstream region (d_2) in individual nuclei. We summed d_1 and d_2 to generate the total 3D distance (d_t) (Figures 6B, 6C, and S6A; Joyce et al., 2012; Rosin et al., 2018; Viets et al., 2019). Larger distances reflect a more open state, while smaller distances indicate a more compact state (Figures 6B and 6C). We hypothesized that transcribed ss loci would be more open compared with inactive ss loci, which would be more compact.

To test our method, we examined compaction in peripodial membrane cells and antennal cells. The ss locus was more compact in peripodial membrane cells (Ctrl) where ss is repressed, with a median compaction of 588 nm (lower quartile = 456 nm; upper quartile = 720 nm). The ss locus was significantly more open in antennal cells (A) where ss is expressed, with a median compaction of 809 nm (lower quartile = 643 nm; upper quartile = 985 nm) (Figure 6D). Thus, the DNA FISH method discerned differences in DNA compaction between cells with active or repressed ss.

Chromatin compaction is dynamic during R7 differentiation

We examined compaction at the ss locus in the developing eye. In single eye-antennal discs, we imaged all stages of R7 differentiation. As in previous experiments, we determined the differentiation state of cells based on their positions relative to the MF. Undifferentiated cells were anterior to the MF. Posterior to the MF, precursors were located at 0–10 μ m, differentiating cells were located at 10–30 μ m, and R7s were located at >30 μ m. R7s were also labeled by immunohistochemistry of a GFP reporter expressed in all R7s. The reporter did not alter ss expression in R7s or compaction dynamics (Figures S6M–S6O).

The ss locus was more compact in undifferentiated cells (ss^{OFF}), similar to peripodial membrane cells (ss^{OFF}) (Figure 6D). The ss locus was more open in precursors (ss^{ON}), similar to antennal cells (ss^{ON}) (Figure 6D). We predicted that differentiating cells (ss^{OFF}) would be compact but were surprised to observe intermediate compaction (Figure 6D). This intermediate compaction was also observed in R7s (a mix of ss^{ON} and ss^{OFF}) (Figure 6D). The intermediate compaction measurements suggested two main possibilities: (1) there are two distinct populations of cells, with compact or open chromatin at the ss locus, or (2) the ss locus is at an intermediate compaction state across all cells.

To discern between these hypotheses, we identified ss^{ON} and ss^{OFF} R7s using the reporter *Ins5*. ss^{ON} R7s were identified based on *Ins5* expression. ss^{OFF} R7s were identified based on

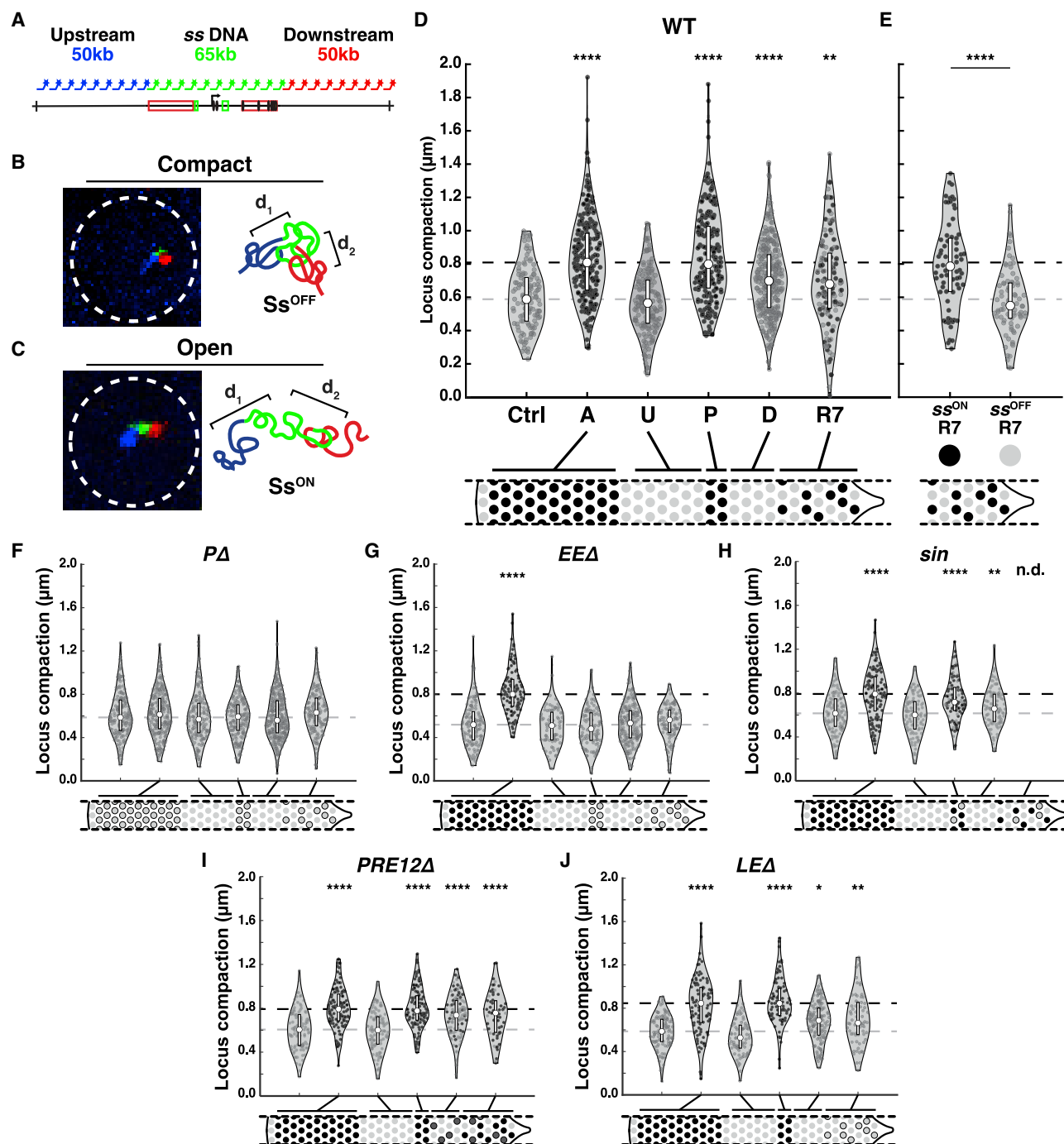


Figure 6. Dynamic chromatin compaction of the ss locus

(A) Schematic of DNA FISH probes used to label the upstream (blue), ss locus (green), and downstream (red) regions.

(B and C) Left, image; right, schematized model.

(B) Ss^{OFF} cell with compact chromatin.

(C) Ss^{ON} cell with open chromatin.

(D–J) Quantification. Ctrl, peripodial membrane cells; A, antennal cells; U, undifferentiated cells; P, precursors; D, differentiating cells; R7, R7s. Black circle, Ss^{ON} cell; gray circle, Ss^{OFF} cell; white rectangle, quartile; white circle, median; gray dashed line, Ss^{OFF} control median; black dashed line, Ss^{ON} control median. * $p < 0.05$; ** $p < 0.005$; **** $p < 0.0001$.

(D and F–J) Ctrl cells were compared with A, U, P, D, or R7 cells. For (E), Ss^{ON} R7s were compared with Ss^{OFF} R7s.

(D–J) $n > 70$ cells for each region.

(D) WT.

(E) Compaction in Ss^{ON} R7s and Ss^{OFF} R7s.

(legend continued on next page)

the absence of *Ins5* expression and their positions. The ss locus was more open in ss^{ON} R7s (median = 786 nm), similar to other ss^{ON} cells, whereas the ss locus was more compact in ss^{OFF} R7s (550 nm), similar to other ss^{OFF} cells (Figures 6D and 6E). These data suggested that the intermediate average compaction measurements observed for all R7s represented two distinct populations: ss^{ON} R7s with a more open ss locus and ss^{OFF} R7s with a more compacted ss locus.

Differentiating cells do not express ss and the variability in chromatin compaction prevented identification of two distinct cell populations. To characterize changes in compaction over time in differentiating cells, we examined “early” (at 10–20 μ m) and “late” (at 20–30 μ m) differentiating cells and observed no significant differences in compaction (Figures S6N and S6O). Considering (1) the temporal progression of development from differentiating cells to R7s and (2) the similarity of intermediate compaction between ss^{OFF} differentiating cells and the total population of R7s (including ss^{ON} and ss^{OFF} R7s), we surmise that differentiating cells also represented two populations: cells with a more open ss locus and cells with a more compacted ss locus. We cannot rule out that differentiating cells are comprised of cells with intermediate compaction states or a mix of cells with open, compact, and intermediate compaction states.

To determine differential compaction between regions of the locus, we evaluated d_1 and d_2 distances individually. We did not detect differences in compaction when evaluating d_1 or d_2 compared with d_t (i.e., $d_t = d_1 + d_2$) (Figures S6A and S6C–S6F), suggesting no asymmetries in local compaction. We also calculated the angles between d_1 and d_2 and observed similar distributions across the four phases of R7 subtype specification and controls (Figures S6B and S6G–S6L), suggesting no detectable changes in DNA looping at the locus across development.

Comparing the expression and compaction states for developing R7s over time, we find that the ss locus is (1) inactive and compact in undifferentiated cells, (2) active and open in precursors, (3) inactive and likely a mix of open and compact in differentiating cells, and (4) active and open, or inactive and compact in R7s. We next examined the relationship between transcription and chromatin compaction during R7 subtype specification.

Transcription in precursors is required for large-scale decompaction of the ss locus

The early expression of ss in precursors driven by the *EE* is required for expression later in R7s driven by the *LE*. As no discernible Ss protein is generated during the early expression in precursors, and Ss does not feedback to regulate its expression, we hypothesized that early expression in precursors was required to open the ss locus. To test this hypothesis, we examined promoter mutants ($P\Delta$) that do not express ss. In $P\Delta$ mutants, the ss locus was compact in undifferentiated cells and remained compact in precursors, differentiating cells, and

R7s (Figure 6F), suggesting that transcription plays a role in opening the ss locus in precursors. In $P\Delta$ mutants, the ss locus was similarly compact in antennal cells, which normally express ss and have open chromatin (Figure 6F). These data suggest that ss transcription is required for large-scale decompaction of the ss locus.

To test whether the *EE* is required to open the ss locus, we examined chromatin compaction in *EE* Δ mutants. In *EE* Δ mutants, ss was not expressed in precursors and R7s but remained active in the antenna. In *EE* Δ mutants, the ss locus was compact in undifferentiated cells, precursors, differentiating cells, and R7s but was open in antennal cells (Figure 6G). Thus, the *EE* is required for decompaction of the ss locus in precursors, differentiating cells, and R7s.

Decreasing *EE* activity reduced the proportion of precursors that expressed ss and the ratio of Ss^{ON} R7s. Since *EE* activity is required for chromatin decompaction in precursors, we hypothesized that decreasing ss expression in precursors would decrease the number of open cells in precursors and differentiating cells, resulting in a more compact state.

To test this idea, we assessed ss locus compaction for genetic conditions that reduced the number of ss-expressing precursors and the ratio of Ss^{ON} R7s. In flies with *sin*, the ss locus was compact in the control peripodial membrane cells (ss^{OFF}) and undifferentiated cells (ss^{OFF}) and open in antennal cells (ss^{ON}), similar to wild type (Figure 6H). In *sin* precursors (mix of ss^{ON} and ss^{OFF}), the ss locus displayed intermediate compaction and did not open to the same degree as the antenna (Figure 6H). The ss locus displayed intermediate compaction in differentiating cells (ss^{OFF}), but the ss locus was more compact than in precursors, consistent with fewer cells opening in precursors. Flies with overexpression of Klu displayed similar effects on ss locus compaction dynamics (Figure S6P). Though *ash2* and *lid* null mutant flies displayed decreases in cell densities (Figures S3D and S3E), limiting our analyses, ss locus compaction trended toward similar effects in these mutants (Figures S6Q and S6R). These data suggest that decreasing the proportion of precursors that expressed ss led to more compact chromatin in precursors. The further decrease in compaction in differentiating cells is consistent with the ss locus remaining open in some cells and closing in others.

Repression is required for ss locus compaction in differentiating cells

We next investigated how loss of repression affects ss locus compaction in mutants that delete PRE1 and PRE2 (*PRE12* Δ). In *PRE12* Δ mutants, ss was repressed in undifferentiated cells and expressed in precursors. We observed ectopic expression in differentiating cells and the proportion of Ss^{ON} R7s increased. ss was expressed in antennal cells and repressed in peripodial membrane cells. In homozygous *PRE12* Δ mutants, the ss locus was compact in undifferentiated cells and peripodial cells (ss^{OFF}) and open in precursors and antennal cells (ss^{ON}) (Figure 6I). The

(F) $P\Delta$.

(G) *EE* Δ .

(H) *sin*.

(I) *PRE12* Δ .

(J) *LE* Δ .

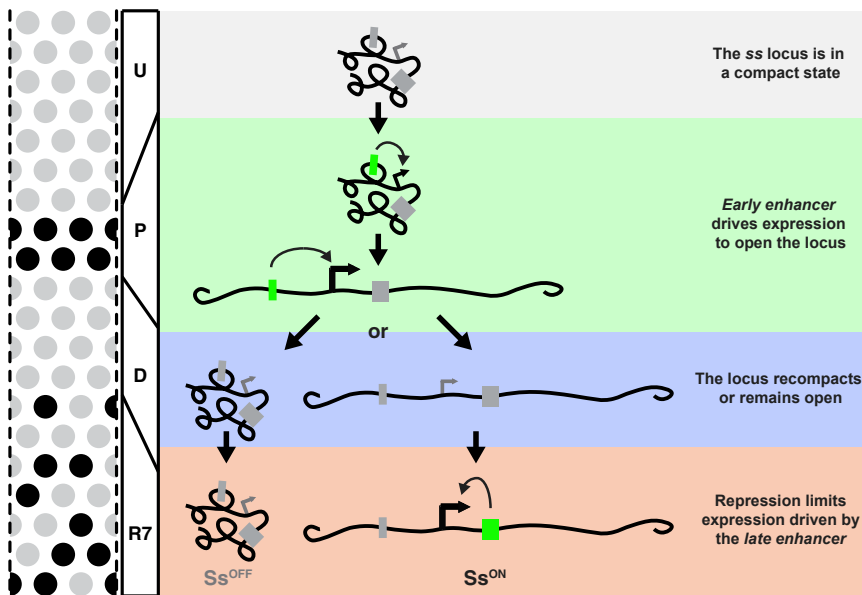


Figure 7. Proposed mechanism for stochastic R7 subtype specification

Gray box, inactive enhancer; green box, active enhancer.

As the cells mature, *ss* expression ceases, and the *ss* locus compacts during the transition from the precursor to the differentiating cell phase. Our observations that (1) the *ss* locus is open in *Ss*^{ON} R7s and compact in *Ss*^{OFF} R7s and (2) the similarity of median compaction in differentiating cells and all R7s (including *Ss*^{ON} R7s and *Ss*^{OFF} R7s) suggest that the *ss* locus assumes either an open or compact state in differentiating cells that is maintained until terminal R7 subtype specification. Our data are consistent with stable compaction states in differentiating cells, but

ss locus was more open in differentiating cells and R7s in *PRE12Δ* mutants compared with wild-type controls (Figure 6I), suggesting that repression by PREs is required to properly compact the *ss* locus and repress expression in differentiating cells and R7s.

Compaction is independent of LE activity

Finally, we tested the interaction of chromatin compaction and expression driven by the *LE*. In *LEΔ* mutants, *ss* was expressed in precursors, turned off in differentiating cells, and remained off in all R7s. We hypothesized that either (1) the *ss* locus would display intermediate compaction in R7s, similar to differentiating cells, or (2) the *ss* locus would be more compacted similar to other *ss*^{OFF} cells. In *LEΔ* mutants, the *ss* locus was compact in undifferentiated cells, open in precursors, and intermediate in differentiating cells and R7s (Figure 6J), like in wild-type flies (Figure 6D), suggesting that chromatin state in maturing R7s is not dependent on transcription driven by the *LE*.

Together, these data suggest that expression in precursors driven by the *EE* is required to open the *ss* locus, repression mediated by PREs is required to compact the *ss* locus in a subset of differentiating cells, and compaction state is independent of expression driven by the *LE*.

DISCUSSION

Temporally dynamic antagonism stochastically specifies R7 subtypes

We investigated how regulation of transcription and chromatin compaction at the *ss* locus controls stochastic R7 patterning in the fly eye. *ss* is initially in a compact, repressed state in undifferentiated cells. This compacted state is similar in other *Ss*^{OFF} cell types including peripodial cells and *Ss*^{OFF} R7s. As the eye develops, *ss* is transcribed in precursors and chromatin is opened. *ss* transcription and large-scale decompaction are lost in mutants deleting the *EE* or the *promoter*, suggesting that *ss* transcription drives the opening of the *ss* locus in precursors early.

they cannot rule out changes during this phase of R7 development.

ss expression and compaction during the transition from precursor to differentiating cell phases are critical processes that determine the stochastic R7 fate choice. Decreasing *EE* activity reduced *ss* expression in precursors and the proportion of *Ss*^{ON} R7s. Extending *ss* transcription into the differentiating cell phase increased the proportion of *Ss*^{ON} R7s. We propose that variable activation and duration of transcription in each precursor determines the probability of recompaction, which ultimately dictates the *Ss*^{ON} or *Ss*^{OFF} expression state in R7s.

In the last stage of R7 subtype specification, *ss* expression driven by the *LE* is repressed in a subset of R7s. The repression reporter strategy showed that repression at the *ss* locus limits expression to a subset of R7s. The chromatin compaction assays showed that the *ss* locus is open in *Ss*^{ON} R7s and compact in *Ss*^{OFF} R7s. Deletion of the *LE* ablated expression but did not alter compaction in R7s. Thus, the chromatin state is set and maintained independent of expression at this stage of R7 maturation. Further, open chromatin is not sufficient to activate *ss* expression. Together, our data suggest that open chromatin allows activation by the *LE* whereas compact chromatin represses *ss* expression.

Based on these findings, we propose a mechanism that controls stochastic R7 subtype specification (Figures 7 and S7). The *ss* locus is in a compact state in undifferentiated cells (Figure 7, U). The *EE* drives transcription and opens the *ss* locus in precursors (Figure 7, P). Early expression ceases and the *ss* locus randomly assumes an open or compact state in differentiating cells (Figure 7, D). R7s with open chromatin at the *ss* locus reactivate *ss* and take on the *Ss*^{ON} R7 fate, whereas R7s with compact chromatin at the *ss* locus repress *ss* and take on the *Ss*^{OFF} R7 fate (Figure 7, R7).

Prime and boost mechanisms controlling cell fate specification

A key aspect of this mechanism is the initial “priming” or opening of the *ss* locus during the early expression in precursors.

Transcription-based priming plays important roles in several stereotyped developmental programs (Anderson et al., 2016; Cochella and Hobert, 2012; Greenberg et al., 2017; Kaikkonen et al., 2013; Schmitt et al., 2005). A well-understood example has been described in *C. elegans*, where the bilateral pair of ASE gustatory neurons display asymmetric gene expression and function (Ortiz et al., 2006, 2009). Stereotyped specification of the left neuron ASEL is dependent upon the asymmetric expression of the microRNA *Isy-6* (Johnston and Hobert, 2003), achieved by a “prime and boost” mechanism. Several cell divisions prior to the birth of the terminal ASEL neuron, a pulse of *Isy-6* expression in the precursor cell promotes decompaction of the *Isy-6* locus. This decompacted state is maintained in the ASEL lineage throughout development, allowing for reactivation of *Isy-6* in the terminal ASEL neuron. In the ASER lineage that never experiences the early pulse of *Isy-6* expression, the locus remains in a repressed, compacted state, preventing later activation by transcription factors that are expressed in both ASE neurons (Charest et al., 2020; Cochella and Hobert, 2012). Thus, early transcription of a key regulator (*Isy-6*) promotes one cell fate (ASEL) by antagonizing chromatin-mediated repression important for the specification of the alternative fate (ASER).

The transcription-based prime and boost mechanism controlling ASEL/R sensory neuron specification in *C. elegans* has many similarities to the mechanism that we have identified for R7 subtype specification. In both systems, early expression of a key regulator in precursor cells opens a locus (prime) so that it can be reactivated later upon terminal specification (boost). A major difference is that the ASEL/R decision requires priming in only the ASEL lineage to reproducibly generate the ASEL fate, whereas the R7 subtype decision utilizes priming in all precursors, which opens the chromatin followed by variable chromatin compaction and repression that ultimately determines the Ss^{ON} or Ss^{OFF} R7 fate.

Both the ASEL/R and R7 subtype decisions also exhibit a window of inactivity between the early and late expression phases. However, this window appears to play two very different roles. In the ASEL/R decision in worms, the early priming of the *Isy-6* locus occurs several cell divisions prior to terminal differentiation. The time between the prime and boost is an obstacle that must be overcome to remember the early developmental event. In contrast, the window between the early and late stages of *ss* expression appears to enable chromatin compaction and repression that determine the Ss^{ON} or Ss^{OFF} expression states in R7s.

Shared features of stochastic fate specification

Though stochastic fate specification is an important feature of many cell fate programs, general features of these mechanisms have not been identified. In the bacterium *Bacillus subtilis*, transcriptional regulation is critical, as ComK transcription drives a stochastic cell fate switch to the “competent” fate. In both the competence decision in bacteria and R7 subtype specification in flies, all “precursor” cells express the key regulator, yet only a subset undergo the cell fate switch (Maamar et al., 2007; Mugler et al., 2016; Süel et al., 2006).

Stochastic R7 subtype specification in flies also shares mechanistic features with OR selection in mice, particularly in the

repression of alternative fates. In the olfactory system, OR genes are found in a compact heterochromatic region in the nucleus, with one gene that escapes repression and activates (Clowney et al., 2011; Magklara et al., 2011). Similarly, chromatin compaction and repression play key roles in determining ss^{ON} and ss^{OFF} R7 fates. Our studies in flies bridge the roles of transcription in bacteria and chromatin in mice for stochastic cell fate specification.

Stochasticity and the antagonism between transcription and chromatin

Our understanding of the relationship between transcription and chromatin is often a chicken and egg problem: it is unclear whether transcription state dictates large-scale chromatin state or vice versa. Here, we provide evidence that clearly identifies these cause-effect relationships and show how they change during development. The *EE* drives transcription to open chromatin in precursors. In differentiating cells, the *EE* ceases to function and transcription stops. Chromatin remains open or closes, marking the stochastic step. Finally, the *LE* turns on in mature R7s. In cells where the locus is open, transcription reinitiates, while in cells where the locus is closed, transcription is repressed. Thus, initially, transcription state regulates chromatin state and later, chromatin state controls transcription state.

Our studies not only outline this simple mechanism, but also identify how the stochastic step is regulated. The stochastic step occurs as cells cease *ss* transcription in the precursor phase and assume the open or compact chromatin state in differentiating cells. Decreasing or extending early transcription alters the probability of chromatin closing and ultimately, the proportion of R7 subtypes. Thus, variability in the duration of early transcription is likely a key input that determines the stochastic decision. Our findings provide an important step in understanding how transcription and large-scale chromatin states regulate one another to control how cells randomly assume fates.

Limitations of the study

Our data suggest that transcription drives large-scale chromatin decompaction and then compaction represses transcription, which controls stochastic cell fate specification. Our study focused on large-scale chromatin remodeling. We hypothesize that local changes of histone modifications at the enhancer and promoter likely precede transcription in precursors. The heterogeneity of the tissue limited testing this hypothesis in a cell-type-specific manner in intact tissue. Moreover, we concluded that chromatin compaction represses *ss* expression in a subset of R7s. The repression reporter experiments showed that the chromatin context at the *ss* locus is sufficient to repress expression in a subset of R7s. The DNA FISH experiments showed that the *ss* locus is open in ss^{ON} R7s and compact in ss^{OFF} R7s. Our experiments were limited as we could not identify conditions to artificially induce compaction and show sufficiency of compaction to repress *ss*. Additionally, our studies were conducted in fixed tissue, limiting the observation of the rapid temporal interplay between transcription and chromatin. Future studies could address this challenge with live imaging, enabling assessment of the transcriptional and chromatin dynamics of the

ss gene throughout the maturation of individual R7s during subtype specification.

STAR★METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.devcel.2022.06.016>.

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AUTHOR CONTRIBUTIONS

Conceptualization, L.V., C.A., and R.J.J.; methodology, L.V. and C.A.; software, L.V. and T.G.; validation, L.V. and C.A.; formal analysis, L.V.; investigation, L.V., C.A., E.U., R.Y., S.T., A.N.-F., and J.D.; writing—original draft preparation, L.V., C.A., and R.J.J.; writing—review and editing, L.V. and R.J.J.; visualization, L.V., C.A., and R.J.J.; supervision, R.J.J.; project administration, R.J.J.; funding acquisition, R.J.J.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Mouse anti-Lamin B antibody	DSHB	ADL67.10; RRID: AB_528336
Mouse anti-Lamin B antibody	DSHB	ADL84.12; RRID: AB_528338
Rabbit anti-GFP antibody	Invitrogen	A21311; RRID: AB_221477
Rabbit anti-RFP antibody	MBL International	PM005, RRID: AB_591279
Rabbit anti-Rh4 antibody	gift from C. Zuker, Columbia University	N/A
Mouse anti-Rh3 antibody	gift from S. Britt, University of Texas at Austin	N/A
Rat anti-Elav antibody	DSHB	7E8A10; RRID: AB_528218
Guinea pig anti-Ss antibody	gift from Y.N. Jan, University of California, San Francisco	N/A
Deposited data		
Predicted PRE data	modENCODE	FlyBase ID FBic0000414
scATAC-seq data	UCSC Genome Browser Custom Track	Available at: http://genome.ucsc.edu/s/cbravo/Bravo_et_al_EyeAntennalDisc
Experimental models: Organisms/strains		
<i>yw</i> ; +; <i>Df(3R)Exel6269</i>	Bloomington Stock Center	7736; RRID: BDSC_7736
<i>yw</i> ; <i>ey>Gal4</i> , <i>UAS>flp/+</i> ; <i>FRT82B ash2¹</i> , <i>e</i> , <i>red/GMR>hid</i>	Bloomington Stock Center	5253; RRID: BDSC_5253
<i>yw</i> ; <i>FRT40A lid¹⁴⁰</i> , <i>FRT40A GMR>hid</i> ; <i>ey>Gal4</i> , <i>UAS>flp/+</i>	Bloomington Stock Center	76954; RRID: BDSC_76954
<i>Ato(384)>Gal4/yw</i> ; <i>UAS>klu/+</i> ; <i>P[yf+t7.7] w[+mC]=20XUAS-6XGFP}attP2/+</i>	Bloomington Stock Center	52262; RRID: BDSC_52262
<i>yw</i> , <i>M{3xP3-RFP.attP}ZH-2A</i> ; +; +; <i>M{RFP [3xP3.PB] GFP[E.3xP3]=vas-int.Dm}ZH-102D</i>	Bloomington Stock Center	24480; RRID: BDSC_24480
<i>elav>Gal4/w</i> ; <i>UAS>Dcr2/+</i> ; <i>UAS>GFP RNAi/+</i>	Bloomington Stock Center	35786; RRID: BDSC_35786
<i>elav>Gal4/w</i> ; <i>UAS>Dcr2/+</i> ; <i>UAS>Cp190 RNAi/+</i>	Bloomington Stock Center	35078; RRID: BDSC_35078
<i>elav>Gal4/w</i> ; <i>UAS>Dcr2/+</i> ; <i>UAS>su(Hw) RNAi/+</i>	Bloomington Stock Center	33906; RRID: BDSC_33906
<i>elav>Gal4/w</i> ; <i>UAS>Dcr2/+</i> ; <i>UAS>mdg4 RNAi/+</i>	Bloomington Stock Center	33907; RRID: BDSC_33907
<i>elav>Gal4/w</i> ; <i>UAS>Dcr2/+</i> ; <i>UAS>trx RNAi/+</i>	Bloomington Stock Center	33703; RRID: BDSC_33703
<i>elav>Gal4/w</i> ; <i>UAS>Dcr2/+</i> ; <i>UAS>ash2 RNAi v20/+</i>	Bloomington Stock Center	64942; RRID: BDSC_64942
<i>elav>Gal4/w</i> ; <i>UAS>Dcr2/+</i> ; <i>UAS>ash2 RNAi v22/+</i>	Bloomington Stock Center	35388; RRID: BDSC_35388
<i>elav>Gal4/w</i> ; <i>UAS>Dcr2/+</i> ; <i>UAS>Mnn1 RNAi/+</i>	Bloomington Stock Center	35150; RRID: BDSC_35150
<i>elav>Gal4/w</i> ; <i>UAS>Dcr2/+</i> ; <i>UAS>Trl RNAi/+</i>	Bloomington Stock Center	67265; RRID: BDSC_67265
<i>elav>Gal4/w</i> ; <i>UAS>Dcr2/+</i> ; <i>UAS>Pc RNAi/+</i>	Bloomington Stock Center	36070; RRID: BDSC_36070
<i>elav>Gal4/w</i> ; <i>UAS>Dcr2/+</i> ; <i>UAS>lid RNAi/+</i>	Bloomington Stock Center	28944; RRID: BDSC_28944
Oligonucleotides		
Oligos for the ss locus deletion CRISPR	See Table S4 for homologous bridge sequence, mutation size, and phenotypic effect	N/A
Oligos for the repression reporter insert CRISPR	See Table S5 for homologous bridge sequence, mutation size, and phenotypic effect	N/A
Software and algorithms		
MATLAB_R2019b	MathWorks	https://www.mathworks.com
ImageJ	(Schindelin et al., 2012)	https://imagej.nih.gov/ij/

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Continued

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Script used to generate 19-bp barcoding primers for Oligopaints probe Design	(Viets et al., 2019)	Available at: https://github.com/kviets0913/Oligopaints-Primers-Custom-Script
Custom script used to analyze images and quantify the density of nascent RNA spots	Generated for this study	Available at: https://github.com/lvoortman/Automated_Image_Analysis

RESOURCE AVAILABILITY

Lead contact

All information queries or requests for resources can be directed to and will be fulfilled by the lead contact, Robert Johnston Jr. (robertjohnston@jhu.edu).

Materials availability

All reagents and fly lines are available upon request.

Data and code availability

The script used to generate 19-bp bar-coding primers for Oligopaints probe design is available at <https://github.com/kviets0913/Oligopaints-Primers-Custom-Script>. The custom script used to analyze confocal images and determine the density of nascent RNA expression is available at <https://zenodo.org/badge/latestdoi/504600646>.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Drosophila lines

Flies were raised on standard cornmeal-molasses-agar medium and grown at 25° C. All experiments in this study included both male and female flies. See [Table S3](#) for a full list of fly genotypes used.

METHOD DETAILS

Confocal image acquisition

All images were acquired using a Zeiss LSM 700 or LSM 980 confocal microscope. Adult retina images were acquired at a single Z plane at 20x magnification. Immunohistochemistry images at the pupal and larval stages were taken at 40x magnification as minimal Z stacks with a slice thickness of 500 nm. Image acquisition for larval DNA and RNA FISH experiments were taken at 63x magnification as large Z stacks encompassing the tissue with a slice thickness of 300nm.

CRISPR-mediated deletions

Deletions to the endogenous *ss* locus were generated using CRISPR (Gratz et al., 2014; Port et al., 2014). Sense and antisense DNA forward and reverse strands of the gRNA were designed to generate BbsI restriction site overhangs. The oligos were annealed and cloned into the pCFD3 cloning vector (Addgene, Cambridge, MA). A single stranded DNA homology bridge was generated with 60 bp homologous regions flanking each side of the predicted cleavage site. The gRNA construct (500 ng/μl) and homology bridge oligo (100 ng/μl) were injected into *Drosophila* embryos (BestGene, Inc.). Single males were crossed with a balancer stock (yw; +; TM2/TM6B), and F1 female progeny were screened for the deletion via PCR and sequencing. Single F1 males whose siblings were deletion-positive were crossed to the balancer stock (yw; +; TM2/TM6B) and the F2 progeny were screened for the deletion via PCR and sequencing. Information on all CRISPR oligonucleotides used for this study can be found in [Table S4](#).

CRISPR-mediated insertions

Insertion of the *[3XP3]>RFP* enhancer reporter construct into the endogenous *ss* locus was achieved using CRISPR. P3 is a Ey binding site from the *rh1* promoter. sgRNAs were designed using Chopchop and cctop (Labuhn et al., 2018; Labun et al., 2016), isolated from injection stocks, and amplified using *in vitro* transcription. A single stranded DNA homology bridge was generated with homologous regions flanking each side of the predicted cleavage site in the *[3XP3]>RFP* enhancer reporter construct using Gibson Assembly. The homology bridge was co-injected with Cas9 RNA (2 μg/μl) and sgRNA (1 μg/μl) into 300 *Drosophila* embryos (Qidong Fungene Biotechnology Co., Ltd.). Single males were crossed with a balancer stock (yw; +; TM2/TM6B), and F1 female progeny were screened for the deletion via PCR and sequencing. Single F1 males whose siblings were deletion-positive were crossed to the

balancer stock (yw; +; TM2/TM6B) and the F2 progeny were screened for the deletion via PCR and sequencing. Information on all CRISPR oligonucleotides used for this study can be found in [Table S5](#).

Antibodies

Antibodies and dilutions were as follows: mouse anti-lamin B (DSHB ADL67.10 and ADL84.12), 1:100; rabbit anti-GFP (Invitrogen), 1:500; rabbit anti-RFP (MBL), 1:400; rat anti-Elav (DSHB, 7E8A10), 1:50; rabbit anti-Rh4 (gift from C. Zuker, Columbia University), 1:50; mouse anti-Rh3 (gift from S. Britt, University of Texas at Austin), 1:50; guinea pig anti-Ss (gift from Y.N. Jan, University of California, San Francisco), 1:500; all secondary antibodies (Molecular Probes) were Alexa Fluor-conjugated and used at 1:400.

Immunohistochemistry

Adult, mid-pupal, and larval retinas were dissected as described ([Hsiao et al., 2012](#)) and fixed for 15 min with 4% formaldehyde at room temperature. Retinas were rinsed three times in PBS plus 0.3% Triton X-100 (PBX) followed by three 15 min washes in PBX. Retinas were incubated with primary antibodies diluted in PBX >2hrs at room temperature and then rinsed three times in PBX followed by three 15 min washes in PBX. Retinas were incubated with secondary antibodies diluted in PBX >2hrs at room temperature and then rinsed three times in PBX followed by three 15 min washes in PBX. Retinas were mounted in SlowFade Gold Antifade Reagent (Invitrogen). Images were acquired using a Zeiss LSM 700 or LSM 980 confocal microscope at 20x or 40x magnification.

Enhancer reporter

The *early enhancer* and *late enhancer* cassettes were amplified and inserted into the pJR20 plasmid. The *early enhancer* was amplified from chr3R:16,410,464 - 16,411,045. The *late enhancer* was amplified from chr3R:16,399,856 -16,396,676. These sequences were inserted upstream of a heat shock gene minimal promoter and the GFP gene coding sequence. All plasmids used were made through standard cloning procedures. Plasmids, plasmid maps, and cloning details are available on request. All constructs were sent to BestGene (Chino Hills, CA) for injection. Constructs were inserted via PhiC31 integration at the attP40 landing site.

Oligopaints probe design for RNA and DNA FISH

Probes for RNA and DNA FISH were designed using the Oligopaints technique ([Beliveau et al., 2012, 2013, 2015](#)). Target sequences were run through the bioinformatics pipeline available at <http://genetics.med.harvard.edu/oligopaints/> to identify sets of 50-bp optimized probe sequences (i.e. “libraries”) tiled across the DNA sequence of interest. Five 19-bp bar-coding primers, gene F and R; universal (univ) F and R, and either sublibrary (sub) F or random (rando) R, were appended to the 5' and 3' ends of each probe sequence. To ensure that all probes were the same length, an additional 8-bp random sequence was added to the 3' end of the probes. The gene F and R primers allowed PCR amplification of a probe library of interest out of the total oligo pool, and the univ F and R primers allowed conjugation of fluorophores, generation of single-stranded DNA probes, and PCR addition of secondary sequences to amplify probe signal. The ss 50-kb left and right extension libraries had a sub F primer between the gene and universal forward primers to allow PCR amplification of probes targeting a specific sub-region of the locus of interest. All other probe libraries had a rando R primer appended at the 3' end to maintain a constant sequence length between all probes. Bar-coding primer sequences were taken from a set of 240,000 randomly generated, orthogonal 25-bp sequences ([Xu et al., 2009](#)) and run through a custom script (available at <https://github.com/kviets0913/Oligopaints-Primers-Custom-Script>) to select 19-bp sequences with 15-bp homology to the Drosophila genome. Primers were appended to probe sequences using the orderFile.py script available at <http://genetics.med.harvard.edu/oligopaints/>. Completed probe libraries were synthesized as custom oligo pools by Custom Array (Bothell, WA), and fluorescent FISH probes were generated as described in references ([Beliveau et al., 2012, 2013, 2015](#); Viets et al., 2019).

RNA FISH

RNA FISH was performed using modified versions of the protocols described in references ([Beliveau et al., 2012, 2015](#)). 20-50 eye/-antennal discs attached to mouth hooks from third instar larvae were collected on ice and fixed in 129 μ L ultrapure water, 20 μ L 10X PBS, 1 μ L Tergitol NP-40, 600 μ L heptane, and 50 μ L fresh 16% formaldehyde. Tubes containing the fixative and eye discs were shaken vigorously by hand, then fixed for 10 min at room temperature with nutation. Eye discs were then given three quick washes in 1X PBX, followed by three 5-min washes in PBX with 0.5% (vol/vol) RNase inhibitor (Promega) at room temperature with nutation. Eye discs were then removed from the mouth hooks and blocked for 1 h in 1X PBX:Western Blocking Reagent (Roche) at room temperature with nutation. They were then incubated in primary antibody diluted in 1X PBX with 0.5 U/pL RNase inhibitor overnight at 4° C with nutation. Next, eye discs were washed three times in 1X PBX for 20 min and incubated in secondary antibody diluted in 1X PBX with 0.5 U/pL RNase inhibitor for 2 h at room temperature with nutation. Eye discs were then washed two times for 20 min in 1X PBX, followed by a 20-min wash in 1X PBS. Next, discs were given one 10-min wash in 20% formamide + 80% 2X SSCT (2X SSC+0.001% Tween-20), one 10-min wash in 40% formamide + 60% 2X SSCT, and two 10-min washes in 50% formamide + 50% 2X SSCT. Discs were then predenatured by incubating for 4 h at 37° C, 3 min at 92° C, and 20 min at 60° C. Primary probes were added in 36 μ L hybridization buffer consisting of 50% formamide + 50% 2X SSCT+2% dextran sulfate (w/v). All probes were added at a concentration of ≥ 5 pmol fluorophore/mL. 4 μ L of probe was added. After addition of probes, eye discs were incubated at 37° C for 16–20 h with shaking. Eye discs were then washed for 1 h at 37° C with shaking in 50% formamide + 50% 2X SSCT. 1 μ L of each

secondary probe was added at a concentration of 100 pmol/mL in 50 μ L of 50% formamide + 50% 2X SSCT. Secondary probes were hybridized for 1 h at 37° C with shaking. Eye discs were then washed twice for 30 min in 50% formamide + 50% 2X SSCT at 37° C with shaking, followed by three 10-min washes at room temperature in 20% formamide + 80% 2X SSCT and 2X SSCT with nutation. Discs were incubated in 2X SSCT with 300 μ M DAPI for 15 min at room temperature with nutation, followed by three 10-min washes at room temperature in 2X SSC with nutation. Discs were mounted in SlowFade Gold immediately after the final 2X SSC wash and imaged using a Zeiss LSM700 or Zeiss LSM980 confocal microscope at 63x magnification.

DNA FISH

DNA FISH was performed using modified versions of the protocols described in references (Beliveau et al., 2012, 2013, 2015; Viets et al., 2019). 20–50 eye/antennal discs attached to mouth hooks from third instar larvae were collected on ice and fixed in 129 μ L ultrapure water, 20 μ L 10X PBS, 1 μ L Tergitol NP-40, 600 μ L heptane, and 50 μ L fresh 16% formaldehyde. Tubes containing the fixative and eye discs were shaken vigorously by hand, then fixed for 10 min at room temperature with nutation. Eye discs were then given three quick washes in 1X PBX, followed by three 5-min washes in PBX at room temperature with nutation. Eye discs were then removed from the mouth hooks and blocked for 1 h in 1X PBX+1% BSA at room temperature with nutation. They were then incubated in primary antibody diluted in 1X PBX overnight at 4° C with nutation. Next, eye discs were washed three times in 1X PBX for 20 min and incubated in secondary antibody diluted in 1X PBX for 2 h at room temperature with nutation. Eye discs were then washed two times for 20 min in 1X PBX, followed by a 20-min wash in 1X PBS. Next, discs were given one 10-min wash in 20% formamide + 80% 2X SSCT (2X SSC+.001% Tween-20), one 10-min wash in 40% formamide + 60% 2X SSCT, and two 10-min washes in 50% formamide + 50% 2X SSCT. Discs were then predenatured by incubating for 4 h at 37° C, 3 min at 92° C, and 20 min at 60° C. Primary probes were added in 36 μ L hybridization buffer consisting of 50% formamide + 50% 2X SSCT+2% dextran sulfate (w/v), + 1 μ L RNase A. All probes were added at a concentration of \geq 5 pmol fluorophore/mL. For FISH experiments in which a single probe was used, 4 μ L of probe was added. For FISH experiments in which three probes were used, 1.3 μ L of each probe was added. After addition of probes, eye discs were incubated at 91° C for 3 min and at 37° C for 16–20 h with shaking. Eye discs were then washed for 1 h at 37° C with shaking in 50% formamide + 50% 2X SSCT. 1 μ L of each secondary probe was added at a concentration of 100 pmol/mL in 50 μ L of 50% formamide + 50% 2X SSCT. Secondary probes were hybridized for 1 h at 37° C with shaking. Eye discs were then washed twice for 30 min in 50% formamide + 50% 2X SSCT at 37° C with shaking, followed by three 10-min washes at room temperature in 20% formamide + 80% 2X SSCT and 2X SSCT with nutation. Discs were incubated in 2X SSCT with 300 μ M DAPI for 15 min at room temperature with nutation, followed by three 10-min washes at room temperature in 2X SSC with nutation. Discs were mounted in SlowFade Gold immediately after the final 2X SSC wash and imaged using a Zeiss LSM700 or Zeiss LSM980 confocal microscope at 63x magnification.

scATACseq

Regions of the eye/antennal disc containing open chromatin were obtained from publicly available data, http://genome.ucsc.edu/s/cbravo/Bravo_et_al_EyeAntennalDisc (Bravo González-Blas et al., 2020), and viewed on the UCSC Genome Browser (Kent et al., 2002). We analyzed published scATACseq datasets from developing fly eye-antennal discs that were clustered into cell types by integrating scATAC-seq and scRNA-seq data sets (Figure S2D). We mapped the cell type clusters expressing ss to the antennal cells, precursors, and mature photoreceptors of the eye/antennal imaginal disc and compared ATAC-seq profiles. We could not map clusters to the undifferentiated, differentiating, and peripodial membrane cells and did not evaluate chromatin accessibility for these cell types based on these datasets. All tracks were scaled to the same parameters for accurate comparisons.

QUANTIFICATION AND STATISTICAL ANALYSIS

Adult eye quantifications

The frequencies of Rh4- and Rh3-expressing R7s were scored manually for at least five eyes per genotype. R7s co-expressing Rh3 and Rh4 were scored as Rh4-positive (Mazzoni et al., 2008; Thanawala et al., 2013). 100 or more R7s were scored for each eye. RNAi lines were screened using the Gal4/UAS system, with the an elav driver (*elav>Gal4/w; UAS>Dcr2/+; +*). The co-expression of reporters and Rh4- or Rh3-expressing R7s were scored manually for at least five adult eyes per genotype. 100 or more R7s were scored for each eye. Ss^{ON}/Rh4 R7s and Ss^{OFF}/Rh3 R7s were scored independently. Due to the binary nature of the cell fate decision, this approach yielded two assessments of R7 subtype fate.

Density of expression quantification

The density of ss RNA punctae were calculated computationally using a custom written script in MATLAB. Due to homologous chromosome pairing between both copies of the endogenous chromosome, we observed a single dot for each chromosome. All images were acquired as 3D z-stacks with a slice thickness of 300 nm. The most highly expressed 25 slices within the antenna, and all slices containing punctae in R7 precursors were maximum intensity projected. Undifferentiated, precursor, and differentiating cells were demarcated in 10 μ m regions based on distance from the morphogenetic furrow. The punctae were then identified and the area was determined as a bounding box encompassing the identified spots. Nascent RNA spots were distinguished from mature transcripts using an intensity threshold, removing them from density calculations. The density was calculated as the number of punctae per unit area in μ m². To ensure high fidelity identification of spots, four images were quantified manually in parallel, and

the number of spots in each image were compared as the percentage of manual IDs. Automated identification had a mean %ID of 100.02% +/- 0.67% when compared to manual quantification, indicating high fidelity identification and density quantification.

Cell density quantification

All cell density quantifications were performed in 3D on z-stacks with a slice thickness of 300 nm. Quantifications were performed manually using Fiji (Joyce et al., 2012; Schindelin et al., 2012; Schneider et al., 2012). Three bounding boxes of variable area and a thickness of 33μm were drawn, and the number of cells was counted within each boundary. The density was calculated as the number of cells divided by the area demarked by the boundary.

Proportional expression changes quantification

To determine the proportional change from wild-type expression the density of expression in precursors and percentage of Ss^{ON}/Rh4 in R7s for each genotype was divided by the wild-type average of these values. Normalization first occurred for genotypes showing a significant change in cell density. A normalization factor was determined as the fractional change in cell density from wild-type. This value was multiplied to the raw density values for that genotype.

Compaction quantification

All compaction quantifications were performed in 3D on z-stacks with a slice thickness of 300 nm. Quantifications were performed manually using Fiji (Joyce et al., 2012; Schindelin et al., 2012; Schneider et al., 2012). Boundaries were drawn for each image to denote cell type. Undifferentiated, precursor, and differentiating cells were demarcated in 10 μm regions based on distance from the morphogenetic furrow. To determine the z position of each FISH dot, an encapsulating box was drawn around the dot and the Plot Profile tool was used to assess the stack in which the dot was brightest. Due to homologous chromosome pairing between both copies of the endogenous chromosome, we observed a single dot for each DNA region. The positional information was scored for several dots in a single channel before scoring dots in a different channel. This method ensured dot position information was gathered blindly relative to the position of other dots within individual cells. To determine the x-y-z distance between FISH dots, we used the multipoint tool to mark the center position for each spot within each nucleus. The distance between the FISH dots was then calculated in 3D as:

$$D_{total} = \sqrt{(X_1 - X_2)^2 + (Y_1 - Y_2)^2 + (Z_1 - Z_2)^2} + \sqrt{(X_2 - X_3)^2 + (Y_2 - Y_3)^2 + (Z_2 - Z_3)^2}$$

Compaction angle determination

Compaction angle was determined using the position and length information generated in the compaction quantification. The angle (γ) was calculated with the law of cosines as:

$$d_3^2 = d_1^2 + d_2^2 - 2 \cdot d_1 \cdot d_2 \cdot \cos(\gamma)$$

$$\gamma = \cos^{-1} \left(\frac{d_1^2 + d_2^2 - d_3^2}{2 \cdot d_1 \cdot d_2} \right)$$

Statistical analysis

All datasets were tested for a Gaussian distribution using a D'Agostino and Pearson omnibus normality test and a Shapiro-Wilk normality test. If either test indicated a non-Gaussian distribution for any of the datasets in an experiment, datasets were tested for statistical significance using a Wilcoxon rank-sum test (for single comparisons) or a one-way ANOVA on ranks with Dunn's multiple comparisons test (for multiple comparisons). If both the D'Agostino and Pearson and the Shapiro-Wilk tests indicated a Gaussian distribution for all datasets in an experiment, datasets were tested for statistical significance using an unpaired t-test with Welch's correction (for single comparisons) or an ordinary one-way ANOVA with Dunnett's multiple comparisons test (for multiple comparisons). These statistical tests were performed using GraphPad Prism. Statistical tests and p-values are described in the figure legends.