

Altering the Temporal Regulation of One Transcription Factor Drives Evolutionary Trade-Offs between Head Sensory Organs

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SUMMARY

Size trade-offs of visual versus olfactory organs is a pervasive feature of animal evolution. This could result from genetic or functional constraints. We demonstrate that head sensory organ size trade-offs in *Drosophila* are genetically encoded and arise through differential subdivision of the head primordium into visual versus non-visual fields. We discover that changes in the temporal regulation of the highly conserved *eyeless/Pax6* gene expression during development is a conserved mechanism for sensory trade-offs within and between *Drosophila* species. We identify a natural single nucleotide polymorphism in the *cis*-regulatory region of *eyeless* in a binding site of its repressor Cut that is sufficient to alter its temporal regulation and eye size. Because *eyeless/Pax6* is a conserved regulator of head sensory placode subdivision, we propose that its temporal regulation is key to define the relative size of head sensory organs.

INTRODUCTION

The senses animals rely on have been shaped during evolution to better navigate and exploit the environment. As a result, even closely related species living in different ecological niches show variation in the sizes and shapes of their sensory structures. Adaptive variation in visual sensory organs is a fascinating case in point and ranges from almost complete loss of the eyes in darkness-adapted animals (Partha et al., 2017; Rétaux and

Casane, 2013) to the expansion of visual organs and processing areas in some other groups such as tree-dwelling mammals (Campi and Krubitzer, 2010; Campi et al., 2011) and predator insects (Elzinga, 2003). A striking, yet poorly understood feature of natural variation in eye size is that it often occurs as a trade-off between the visual organs and other head sensory structures such as olfactory organs. This was described in a large variety of animal groups including mammals (Nummela et al., 2013) and fishes (Rétaux and Casane, 2013). In arthropods as well, trade-offs between the size of the eyes and of the antennae, where most olfactory organs are located, are pervasive. Examples include beetle species with different life-styles (nocturnal versus diurnal; visual hunters or not (Bauer and Kredler, 1993); fireflies emitting or not emitting light signals (Stanger-Hall et al., 2018); surface and cave crustaceans (Protas and Jeffery, 2012); and millipedes (Liu et al., 2017). This is also the case between and within species of fruit flies, in which eye size often anti-correlates with the size of the face and/or of the antennae (Posnien et al., 2012; Arif et al., 2013; Norry and Gomez, 2017; Gaspar et al., 2019; Keesey et al., 2019). However, the developmental mechanisms that govern such trade-offs are essentially unknown.

A commonly observed property of sensory organ formation is the shared developmental origin of most head sensory structures—such as eyes and noses—that derive from the subdivision of a single multipotent primordium. In vertebrates, the olfactory and lens placodes derive from the subdivision of the anterior aspect of a multipotent preplacodal ectoderm (Grocott et al., 2012; Singh and Groves, 2016). Similarly, during *Drosophila* development, the ectodermal eye-antennal imaginal disk (EAD) gives rise to all external sensory, including the visual (compound eyes and ocelli) and olfactory (antennae and maxillary palps) sense organs, and non-sensory head cuticle. In vertebrates and in flies, antagonistic relationships between gene

regulatory networks (GRNs) and signaling pathways that promote different sensory identities regulate the subdivision of the multipotent primordium (Grocott et al., 2012; Singh and Groves, 2016; Wang and Sun, 2012; Weasner and Kumar, 2013). First active in the entire tissue, their expression segregates as the visual and non-visual territories become distinct (Bhattacharyya et al., 2004; Kenyon et al., 2003). In addition to promoting eye fate, the transcription factor (TF) Pax6 and its two *Drosophila* orthologues Eyeless (Ey) and Twin-of-Eyeless (Toy), play a key role in the growth and the subdivision of the multipotent primordium (Zhu et al., 2017). In *Drosophila*, at early developmental stages, Ey and Toy are co-expressed in the entire EAD with antennal TFs, such as Homothorax (Hth). The expression of these eye and antenna-promoting TFs progressively segregates along the EAD's anterior-posterior axis, delineating the posterior eye and anterior antennal compartments. Eye and antennal TFs mutually repress each other: the antennal TFs Hth and Cut (Ct) directly repress ey expression while Sine oculis, another eye promoting TF, represses ct (Anderson et al., 2012; Wang and Sun, 2012; Weasner and Kumar, 2013). Consequently, loss or gain of function of these selector TFs leads to the transformation of most of the head tissue into visual or olfactory organs at the expense of the other sensory structure (Halder et al., 1995; Czerny et al., 1999; Anderson et al., 2012).

Therefore, the subdivision of a single multipotent primordium into distinct territories through mutual repression by antagonistic TFs is a shared step of the development of head sensory organs across animals. It is thus tempting to speculate that evolutionary mechanisms have exploited this process leading to natural sensory size trade-offs between visual and olfactory organs. A hint in that direction comes from studies on *Astyanax* fishes, which live as cave or surface-dwelling morphs (Rétaux and Casane, 2013). Cave morphs have small lenses and large olfactory placodes, while surface-dwellers show the reciprocal ratio. Chemical manipulation of signaling pathways that regulate the subdivision of the lens versus olfactory territories mimics the differences observed between natural morphs (Hinaux et al., 2016). Whether this is a mechanism of natural variation in sensory trade-offs is unknown. Demonstrating a direct link requires the identification of naturally occurring causal genetic variants and the elucidation of their effect on the GRNs that regulate visual and olfactory sensory organ development. The paucity of model systems amenable to combining comparative, genetic, molecular, and developmental analyses has thus far hindered such an endeavor.

We reasoned that natural variation in eye size between and within *Drosophila* species may offer precisely such a model. We therefore used comparative analyses combined with developmental, molecular, and genome editing approaches to tackle this question. We show that differential subdivision of the EAD, resulting in different proportions of eye and antennal compartments, underlies eye size variation between and within *Drosophila* species. In both cases, this is associated with changes in the temporal regulation of the expression of ey during EAD subdivision. We also demonstrate that in *D. melanogaster* (*D. mel.*), this is caused by a non-coding single polymorphic nucleotide (SNP) present in most natural populations of *D. mel.* This SNP is located in a binding site for the antennal factor Ct

within the eye enhancer of ey. Using CRISPR/Cas 9 genome editing, we show that this SNP is causal to temporal changes in ey expression and to facet number variation. Thus, changes in the subdivision of a multipotent primordium, caused by subtle alterations of the mutual repression between distinct fates, underlies natural variation in sensory trade-offs.

RESULTS

Reciprocal Changes in the Sizes of Visual and Non-visual Head Structures

The insect compound eye is composed of a crystalline array of small units, named facets or ommatidia. In insects, and specifically in *Drosophilids*, eye size depends both on the number and diameter of the ommatidia and is often negatively correlated with face and/or antenna size (Posnien et al., 2012; Arif et al., 2013; Norry and Gomez, 2017; Gaspar et al., 2019; Keesey et al., 2019). In this study, we selected four *Drosophila* species, which presented a larger eye to face width ratio as compared to *D. mel.* (Figures 1A, 1A', and S1). All subsequent morphological measurements were performed on females raised in density-controlled conditions. We focused on *Drosophila pseudoobscura* (*D. pse.*), which had the largest difference in terms of ommatidia number, an increase of 35% as compared to *D. mel.* (Figures 1B and 1B') while sharing similar facet diameters (Figures 1C and 1C'). Interestingly, the third antennal segment, which hosts the olfactory sensillae, was thinner in *D. pse.* as compared to *D. mel.* (Figures 1D and 1E). As a control, we measured tibia length as a proxy to body size (Posnien et al., 2012; Arif et al., 2013) (Figure S2). In line with previous studies (e.g., Posnien et al., 2012; Arif et al., 2013; Gaspar et al., 2019; Keesey et al., 2019), our results suggest that variation in eye, face, or antennal size cannot be explained solely by variation in body size.

Increased facet number has been associated with higher visual acuity in predator flies (Elzinga, 2003; Gonzalez-Bellido et al., 2011). We thus tested whether a modest variation such as the one observed between *D. pse.* and *D. mel.* was potentially relevant to visual function. We measured the minimal angular distance between two successive vertical black stripes resolved by the flies as a read-out of their visual acuity (Figure 1F) (Götz, 1964; Buchner, 1976). *D. pse.* were able to distinguish between more closely juxtaposed stripes (minimal angle = 7.0°) as compared to *D. mel.* (minimal angle = 8.51°) (Figure 1F'). Thus, *D. pse.* have a better visual acuity correlating with an increased number of facets.

A Trade-Off between Eye and Non-eye Progenitor Fields

What is the developmental origin of facet number variation between *D. mel.* and *D. pse.* and why does it inversely correlate with the size of non-visual structures? All external structures of the head of the adult fly including the sensory organs develop from the EAD (Figure 2A). The eye field occupies most of the posterior EAD compartment and is marked by the expression of Eyes Absent (Eya; Figure 2B) (Roignant and Treisman, 2009). We measured the surface of the eye field in late, fully grown EADs (stage P0) using Eya and found that the eye field was 31% larger in *D. pse.* than in *D. mel.* (Figure 2B'). This is very close to the 35% difference in the number of adult eye facets.

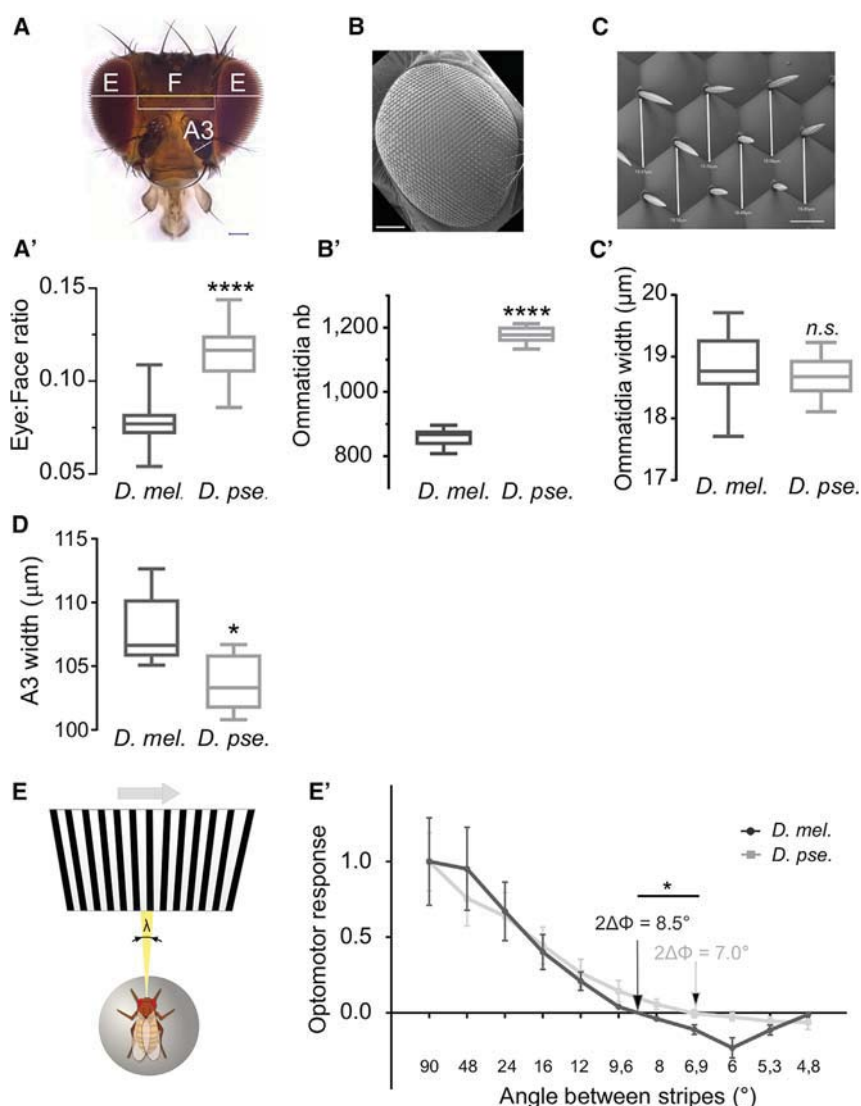


Figure 1. Natural Variation in *Drosophila* Facet Number

(A–C') Eye size comparisons between the *D. mel.* and *D. pse.* Boxes indicate interquartile ranges, lines medians, and whiskers data ranges. Scale bar for (A) and (B), 100 μ m; for (C), 10 μ m.

(A and A') Ratio between eye (E) and face (F) width. Sample sizes: *D. mel.* (n = 23), *D. pse.* (n = 26). Two-tailed unpaired t test: ****p < 0.0001.

(B and B') Ommatidia number. Sample sizes: *D. mel.* (n = 9), *D. pse.* (n = 8). Two-tailed unpaired t test: ****p < 0.0001.

(C and C') Ommatidia width. Sample size: n = 24. Two-tailed unpaired t test with Welch's correction: n.s. p = 0.1677.

(A and D) Third antennal segment (A3) width. Sample sizes: n = 5. Two-tailed unpaired t test: *p = 0.0409.

(E) Optomotor response set-up (see [STAR Methods](#)).

(E') Optomotor response (normalized to max) of *D. mel.* and *D. pse.* females in function of stripe width (spatial wavelength λ); mean \pm SEM. arrows: Spatial resolution (measured as the zero-crossing angle $2\Delta\Phi$). Sample size: n = 9. Two-tailed unpaired t test: *p = 0.014. See also [Figure S1](#).

disks, after the subdivision between the fields is completed ([Figure 2E](#)). The total EAD size was similar between the two species, confirming that it underwent similar growth during prior larval development ([Figure 2E'](#)). In contrast, already at this early stage, the eye field was proportionally larger in *D. pse.* than in *D. mel.* ([Figure 2E''](#)). Thus, the two species differ by the proportion of the multipotent EAD dedicated to the eye versus non-eye tissues, resulting in different proportions of the head structures in the adult. Therefore,

the species variation in eye size involves a developmental trade-off between eye and non-eye primordia.

Temporal Regulation of EAD Subdivision Governs the Trade-Off between Eye and Non-eye Fields

What are the regulatory mechanisms governing this developmental trade-off? EAD subdivision requires the temporally progressive restriction of selector TFs expression to the anterior "antennal" or posterior "eye" compartments, a process completed by mid to late second instar larval stage (L2) ([Kenyon et al., 2003](#)). At this developmental time point, the mutually exclusive expression domains of antenna and eye selectors define the relative sizes of the compartments. In *D. mel.*, a 3.2 kb *cis*-regulatory intron governs *ey* expression during eye development ([Figure 3A](#)). We cloned the orthologous intron from *D. pse.* based on the conservation of the flanking exons. The *D. pse.* intronic sequence is slightly shorter (3.0 kb) with 22% of the intron from *D. mel.* aligning to the corresponding sequences in *D. pse.* ([Figure 3B](#)). Nonetheless, when inserted at the same position in *D. mel.* genome, both *D. mel.* and *D. pse.* introns were

We therefore queried the developmental origin of the difference in eye field size between the two species and considered several possibilities. A first possibility is that the initial pool of embryonic cells forming the EAD differs between the two species. In the late embryo (stage 17), the EAD is composed of a few dozen closely juxtaposed cells located anterior to the brain. Using *Ey* as a marker, we quantified and compared the number of embryonic EAD cells between the two species ([Figures 2D and 2D'](#)) but found no significant difference in the number of EAD progenitors, ruling out this first possibility. Variation in eye field size could also originate from different rates of proliferation. However, the similar density of mitotic cells in the proliferating eye field in EADs of the two species did not support this hypothesis ([Figures 2F and 2F'](#)). In addition, the density of ommatidia progenitor cells in the eye field, characterized by the expression of the proneural factor *Atonal*, was similar between the two species ([Figures S3A–S3A''](#)). Finally, variation in eye field size could also derive from a change in the subdivision of the EAD between eye and non-eye fields. To test this possibility, we compared the proportion of the EAD occupied by the eye field in early L3 imaginal

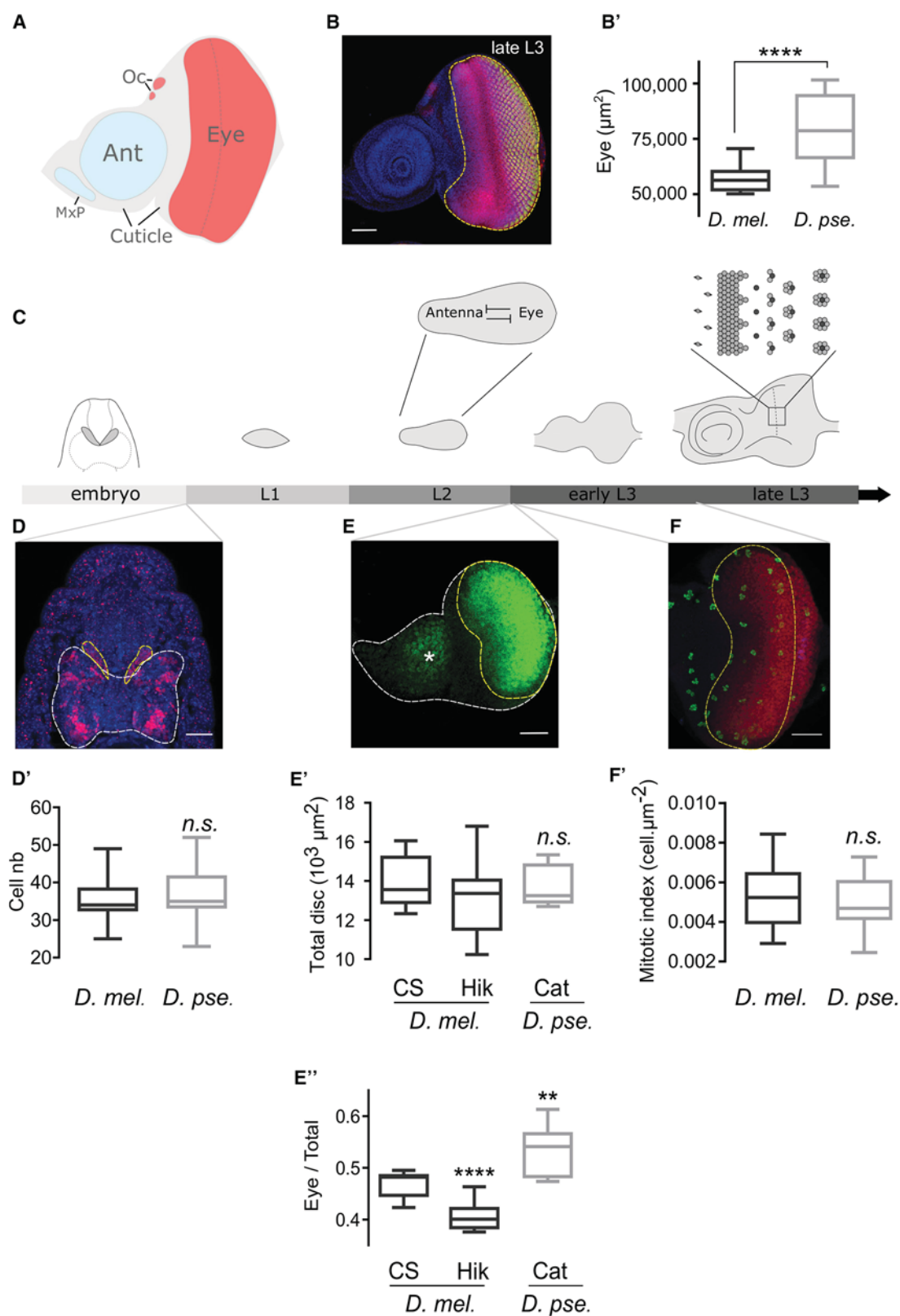


Figure 2. Developmental Origin of Eye Size Variation in *D. mel.* and *D. pse.*

(A) Schematics of the distinct fields of the EAD giving rise to the adult sensory and non-sensory head structures (Ant, antenna; MxP, maxillary palps; Oc, ocelli). Anterior is at the left.

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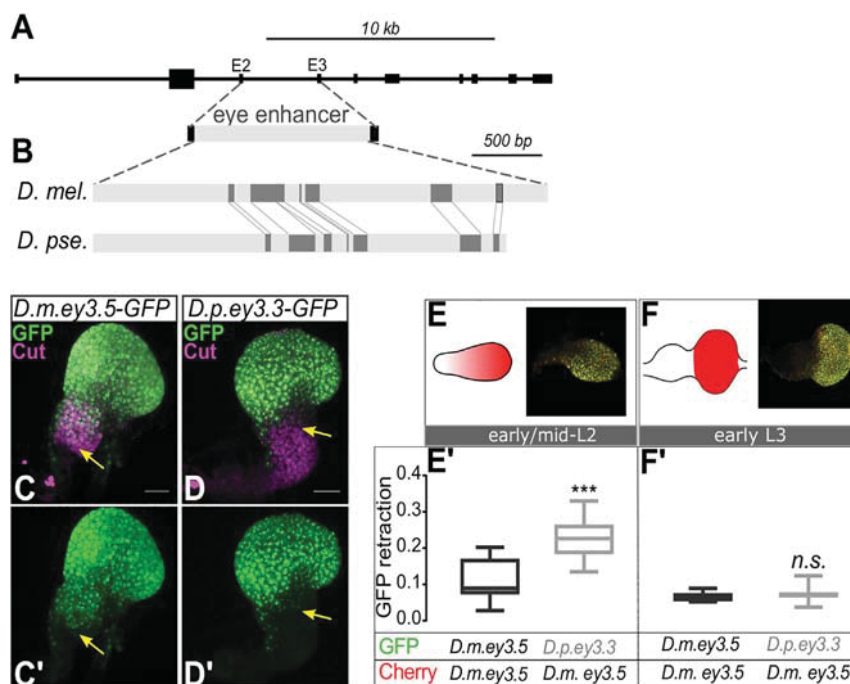


Figure 3. Different Temporal Regulation of EAD Subdivision Governs the Trade-Off between eye and Non-eye Fields

(A) Structure of the *ey* locus showing the location of the intronic *ey3.5* eye enhancer (Hauck et al., 1999). E2: exon 2; E3: exon 3.

(B) Alignment of the *ey* eye enhancer between *D. mel.* and *D. pse.* Dark gray boxes represent the fragments of the *D. mel.* intron sequence that aligned with the orthologous region of *D. pse.* Using BLAST. (C–D') GFP expression driven by *D. mel. ey3.5* and *D. pse. ey3.3* eye enhancers in mid-L2 EADs counterstained with anti-Cut. Green: GFP; magenta: anti-Cut. Yellow arrows indicate the anterior limit of GFP expression. Scale bars: 20 μ m. Anterior is at the bottom.

(E and F) Schematics and immunostainings showing ongoing (E, early/mid-L2 stage) and full (F, early L3 stage) posterior retraction of ey enhancer activity during EAD development.

(E' and F') Pairwise comparison of the difference in expression between GFP and mCherry, measured as the proportion of mCherry that was not colocalized with GFP, when driven by distinct combinations of ey enhancers during (E', early/mid-L2 stage) and after (F', early L3 stage) posterior retraction of ey enhancer activity. (*D.m.ey3.5*, *D. mel. ey* enhancer; *D.p.ey3.3*, *D. pse. Enhancer*).

(E') Early/mid-L2 stage. Sample sizes ($n = 8$, $n = 12$). Two-tailed unpaired t test *** $p = 0.0001$.

(D') Early-L3 stage. Sample sizes ($n = 10$, $n = 3$). Mann-Whitney test *n.s.* $p = 0.6643$.

able to drive GFP expression in the EAD throughout eye development, revealing global functional conservation (data not shown).

We tested whether, despite their overall functional conservation, subtle changes in ey regulation exist between *D. mel.* and *D. pse.* introns (Figures 3C–3D'). In early EADs (late embryos and in L1), both *D. mel.* and *D. pse.* enhancers drove GFP expression across the entire disk (Figure S4). At the L2 stage, we noted that GFP expression driven by the *melanogaster* enhancer (*D.m.ey3.5*) (Figures 3C and 3C') extended further anteriorly into the antennal compartment as compared to the *pseudoobscura* enhancer (*D.p. ey3.3*) (Figures 3D and 3D'). This means that the posterior retraction of expression driven by the two ey enhancers occurs at different velocities. To quantify this effect, we generated two lines of transgenic *D. mel.* flies.

The first line carries two transgenes driving the expression of the red fluorescent protein mCherry and the green fluorescent protein (GFP), respectively, both under the control of *D. mel. ey* enhancer. In this control line, any difference in the expression of mCherry and GFP driven by the same enhancer must only be caused by different dynamics of the two fluorescent proteins. In the second line, mCherry was driven by the *D. mel. ey* enhancer, while GFP was driven by *D. pse. ey* enhancer. In this case, the differences in expression between the GFP and mCherry is caused both by different dynamics of the fluorescent proteins as well as differences in their transcriptional regulation. Thus, to detect differences in the activity of *D. mel.* and *D. pse. ey* enhancers, we performed pairwise comparisons of the difference between GFP and mCherry expression in line 1 versus

(B) Late L3 EAD from *D. pse.* The eye field (yellow dashed line) is labeled with anti-eya (red) and committed photoreceptors are shown in yellow (anti-elav); blue: DAPI. Anterior is at the left. Scale bar: 50 μ m.

(B') Eye field surface in late (P0) eye-antennal disks. Sample sizes: *D. mel.* ($n=19$); *D. pse.* ($n=19$). Two-tailed unpaired t test with Welch's correction **** $p < 0.0001$.

(C) Schematics of eye-antennal disk development. Insets: (C') Segregation of "antennal" and "eye" compartments during L2; (C'') determination of ommatidia precursor cells during L3.

(D) Dorsal view of a late Canton-S embryo depicting the Ey-positive EADs (yellow dashed line) flanking the brain (white dashed line). Anterior is at the top. Scale bar: 25 μ m.

(D') Cell content of late embryonic EADs. Sample sizes: *D. mel.* ($n = 22$); *D. pse.* ($n = 33$). Two-tailed unpaired t test; N. S. $p = 0.0959$.

(E) Early L3 Canton-S EAD co-labeled by Eya (yellow dashed line) and Ct (asterisk). Anterior is at the left. Scale bar: 25 μ m.

(E') Total surface of the EADs (in μ m²). Ordinary one-way ANOVA ($p = 0.4520$) followed by Dunnett's multiple comparisons tests.

(E'') Ratio of eye field versus total EAD surface. Ordinary one-way ANOVA ($p < 0.0001$) followed by Dunnett's multiple comparisons tests. Sample sizes: CS ($n=11$); Hik ($n = 9$); and Cat ($n = 8$).

(F) Proliferative portion of the eye field (dashed yellow line). Red: eye field (anti-Eya); Green: mitotic cells (anti-phosphorylated histone 3). Anterior is at the left. Scale bar: 25 μ m.

(F') Mitotic index of the proliferative eye field in *D. mel.* and *D. pse.* Early L3 EADs. Sample sizes: *D. mel.* ($n = 20$); *D. pse.* ($n = 8$). Two-tailed unpaired t test *n.s.* $p = 0.6185$. Boxes indicate interquartile ranges, lines medians, and whiskers data ranges in all charts (B', D', E', and F').

See also Figure S3.

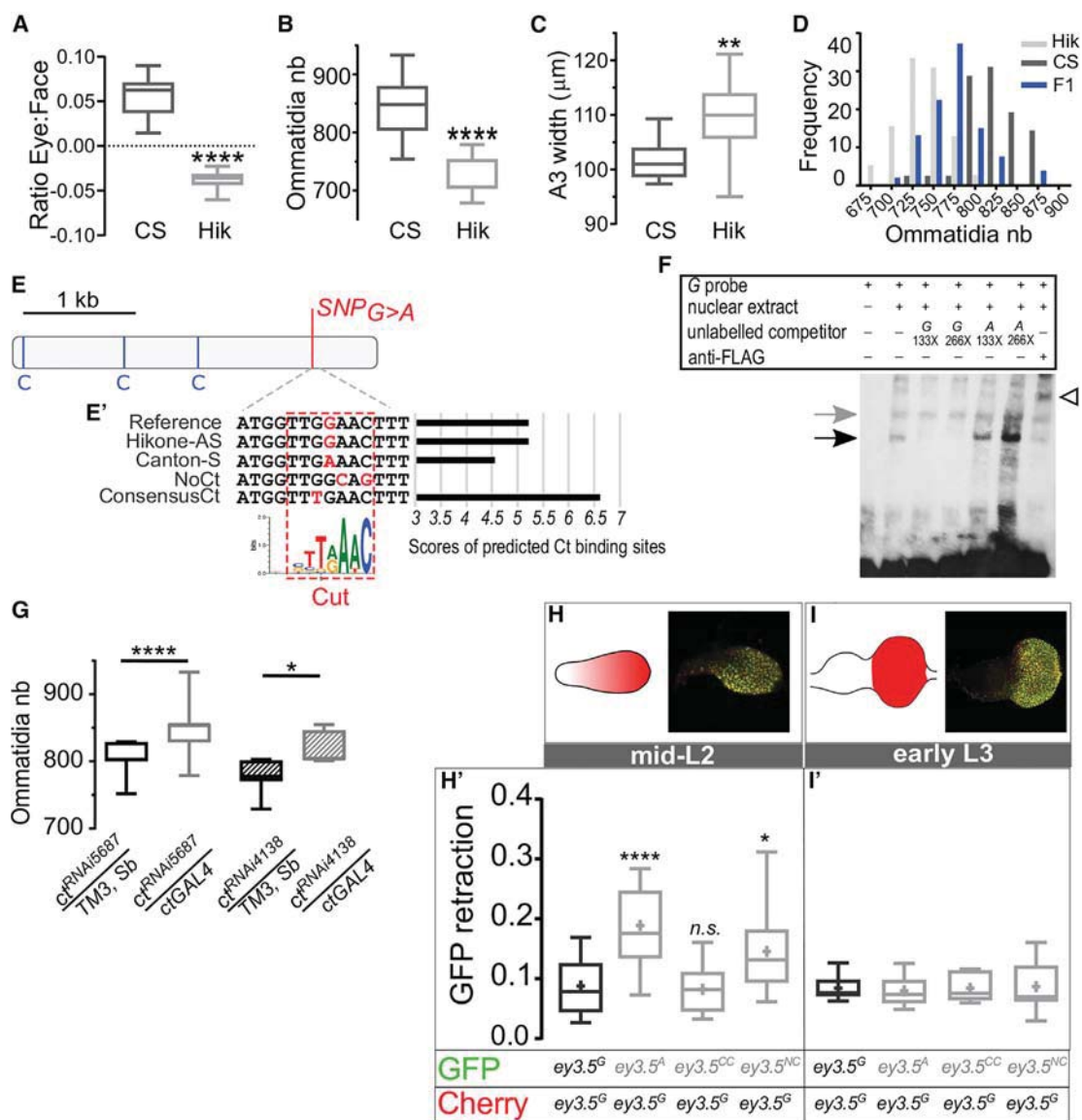


Figure 4. Developmental and Regulatory Origin of Intraspecific Eye Size Variation

(A) Eye versus face width ratio in Canton-S (CS) and Hikone-AS (Hik). Sample sizes: CS: n = 15, Hik: n = 16. Ordinary one-way ANOVA ****p < 0.0001 followed by Tukey's multiple comparisons: ****p < 0.0001. See full data set in supplement (Figure S6); see also Figure S1.

(B) Ommatidia number. Sample sizes: CS: n = 15, Hik: n = 16. Two-tailed unpaired t test: ****p < 0.0001.

(C) Third antennal segment (A3) width. Sample sizes: n = 11. Ordinary one-way ANOVA **p = 0.0035 followed by Tukey's multiple comparisons. **adjusted p = 0.0043. See full figure in supplement (Figure S6).

(D) Frequency distribution of ommatidia numbers in CS (n = 42), Hik (n = 39), and their F1 progeny (n = 54).

(E) Schematics of the *D. mel. ey* eye enhancer (*ey*^{3.5}) showing the localization of the G>A substitution (chr4: 710326) and of three published Ct binding-sites (C, blue lines) (Wang and Sun, 2012).

(E') The dashed red rectangle delineates the low-affinity putative Ct binding site overlapping the position of the G>A substitution (chr4: 710326) (Zhu et al., 2011). For each natural or synthetic enhancer allele, the best score of the Ct binding site as predicted by Cluster-Buster (Frith et al., 2003) is plotted. See also Figure S5; Table S2.

(F) Electrophoretic mobility shift assay. The Cut-FLAG expressing nuclear extract elicits a bandshift when incubated with the G-Probe (black arrow). Excess unlabeled G-probe, but not A-probe effectively competes with the binding, suggesting a higher affinity of Ct for the former. Incubation with an anti-FLAG induces a supershift (empty arrowhead). Gray arrow: non-specific binding.

(G) Ommatidia number variation following RNAi-mediated knockdown of *ct*. Sample sizes from left to right (n = 13; n = 23; n = 10; and n = 8). Ordinary one-way ANOVA ****p < 0.0001 followed by Sidak's multiple comparisons: ****p < 0.0001, *p = 0.0126. See also Figure S5.

(H-I') Pairwise comparison of the expression of the four *D. mel. ey* enhancer alleles: *ey*^{3.5^G} (Hikone-AS); *ey*^{3.5^A} (Canton-S); *ey*^{3.5^{CC}} (ConsensusCt); *ey*^{3.5^{NC}} (NoCt).

(G and H) Schematics and immunostainings of EADs with ongoing (F, early/mid-L2 stage) and full (G, early L3 stage) posterior retraction of *ey* enhancer activity.

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Table 1. Ommatidia Numbers

	Ommatidia Nb (Count)		Ommatidia Nb (Estimated)	
	Mean	St Dev	Mean	St Dev
Canton-S	869.8 ^a	17.96 ^a	828.0	34.00
Hikone-AS	742.4 ^a	20.17 ^a	736.0	27.09
<i>D. pse.</i>	1177 ^a	28.02 ^a	–	–
F1 CS/Hik	–	–	774.6	34.77
<i>ct</i> ⁵⁶⁸⁷ / <i>ct</i> -GAL4	–	–	835.7	25.88
<i>ct</i> ⁵⁶⁸⁷ /TM3,Sb	–	–	802.9	26.99
<i>ct</i> ⁴¹³⁸ / <i>ct</i> -GAL4	–	–	818.6	23.24
<i>ct</i> ⁵⁶⁸⁷ /TM3, Sb	–	–	777.6	22.99
Canton S BH	–	–	830.8	36.42
Canton S TP	–	–	775.4	21.69
Canton S RD	–	–	750.4	20.26
<i>ey3.5</i> ^G	713.8	29.77	745.1	43.76
<i>ey3.5</i> ^{G>A}	737.0	21.12	791	44.49
<i>ey3.5</i> ^G /Df(4)J2	–	–	822.9	34.44
<i>ey3.5</i> ^{G>A} /Df(4)J2	–	–	847.5	19.61

Ommatidia numbers are counted on SEM images (count) or estimated from light-microscopy images using an ellipse-based method (estimated, see STAR Methods and Figure S2).

^aFlies reared at 21°C.

line 2. In early L3 disks, when the antennal and eye compartments have already segregated, the co-expression of mCherry and GFP driven by either *D. mel.* or *D. pse.* regulatory sequences were indistinguishable (Figures 3F and 3F'). In contrast, at L2, during the process of *ey* retraction, the posterior retraction of the GFP was more posteriorly advanced when driven by the *D. pse.* than by the *D. mel.* enhancer (Figures 3E and 3E'). Therefore, the partitioning of the EAD into eye and non-eye fields occurs at an earlier time point in *D. pse.* compared to *D. mel.* Since *Ey* positive cells proliferate more than *Ey* negative cells (Zhu et al., 2017), earlier establishment of the two sensory fields would drive greater differential growth.

A Conserved Mechanism of Sensory Trade-Offs

To understand the genetic basis of sensory trade-off in *Drosophila*, we exploited the fact that such trade-offs have also been observed within single fly species (Cowley and Atchley, 1990; Posnien et al., 2012; Arif et al., 2013; Norry and Gomez, 2017; Gaspar et al., 2019). We found that two wild-type *D. mel.* laboratory strains, Canton-S and Hikone-AS, show different eye-to-face ratios. This is associated with changes in ommatidia number (12, 5% more facets in Canton-S) and diameter (Figures 4A, 4B, and S1A–S1D, Table 1) as well as variation of antennal width (Figure 4C). F1 progeny of Canton-S and Hikone-AS parents presented intermediate ommatidia numbers relative to their parents, demonstrating the heritable nature of

this trait (Figure 4D). We asked if facet number variation between Canton-S and Hikone-AS also originate from changes in the subdivision of the EAD into eye versus non-eye territories. We compared the subdivision of early L3 EADs between the two strains. While the size of the entire EAD was unchanged, the eye field was proportionally larger in Canton-S than in Hikone-AS (Figures 2E–2E'). These data suggest that despite 17–30 million years of separated evolution between the two species groups (Obbard et al., 2012), ommatidia number variation between *D. mel.* and *D. pse.* and between two *D. mel.* strains shares a common developmental logic.

A Single Nucleotide in a Ct Binding Site Distinguishes the Canton-S and Hikone-AS *ey* Regulatory Sequences

Does the difference in EAD subdivision between the Canton-S and Hikone-AS also result from a differential temporal regulation of *ey*? To answer this question, we cloned and aligned the *ey cis*-regulatory sequence from the Canton-S and Hikone-AS strains (Figure 4D). In contrast to the significant divergence observed between *D. mel.* and *D. pse.*, Hikone-AS and Canton-S intron sequences were nearly identical and differed only by a single nucleotide over the entire 3.2 kb intronic region, a G>A substitution at position chr4: 710326. *In silico* analyses revealed that this single nucleotide variant is located in a Ct binding site, distinct from the three sites previously described in the *ey cis*-regulatory sequence (Figure 4E) (Wang and Sun, 2012). Interestingly, the two variants are predicted to display different affinities for the repressor, in a manner that anti-correlates with the number of ommatidia: the A-allele presents a lower affinity score (4.56) and is associated with larger eyes (Canton-S) as compared to the G-allele (Hikone-AS; predicted affinity score 5.22) (Figures 4E' and S4). We thus performed electrophoretic mobility shift assays using a tagged recombinant Ct protein (Figure 4F), and found that Ct has the ability to bind to this sequence. Next, we tested the affinities of the two sequences by performing a competition assay in which unlabeled A-probes or unlabeled G-probes competed with the labeled G-probe. While unlabeled G-probes effectively suppressed the shift, the unlabeled A-probes did not, providing biochemical evidence that the two variants have different affinities for Ct. Put together, these results suggest that the strength of *ey* repression by Ct, a selector TF for antennal fate and a repressor of *ey* expression, influence eye size (Wang and Sun, 2012; Weasner and Kumar, 2013). Consistent with this, RNAi knockdown of *ct* expression during EAD development was sufficient to increase facet number in the adult eye (Figures 4G and S5). We note, however, that this did not consistently result in antagonistic trends in face and antennal width, possibly due to pleiotropic effects of *ct* loss of function on head development (Figure S5).

These findings raise two questions: first, is the G to A substitution in the *ey cis*-regulatory sequence sufficient to cause temporal changes in its activity; and second, if so, might such

(G' and H') Pairwise comparison of the difference in expression between GFP and mCherry, measured as the proportion of mCherry that was not colocalized with GFP, when driven by distinct combinations of *ey* eye enhancers during (F', early/mid-L2 stage) and after (G', early L3 stage) posterior retraction of *ey* enhancer activity. *ey* enhancer variants: *ey3.5*^G, G-variant (Hikone-AS); *ey3.5*^A, A-variant (Canton-S); *ey3.5*^{CC}, ConsensusCt-variant; *ey3.5*^{NC}, NoCt-variant.

(G') Mid-L2 stage. Sample sizes: (n = 16, n = 24, n = 20, and n = 12). Ordinary one-way ANOVA ****p < 0.0001 followed by Dunnett's multiple comparisons versus *ey3.5*^G: ****adjusted p = 0.0001, n.s. adjusted p = 0.9828; *adjusted p = 0.0167.

(H') Early-L3 stage. Sample sizes from left to right (n = 9; n = 10; n = 6; and n = 9). Ordinary one-way ANOVA n.s. p = 0.9652.

changes be caused by alterations in the regulation of the *ey* enhancer by Ct? To tackle these two questions, we used the same strategy described above for comparing the *D. pse.* and *D. mel.* enhancers using GFP and mCherry reporters. We first compared the activities of the Canton-S (A-allele) and the Hikone-AS (G-allele) of the *ey* 3.5 *cis*-regulatory sequences. At early/mid L2, during EAD subdivision, mCherry and GFP co-expression differed between the alleles such that the A-carrying variant (Canton-S; larger eyes) showed further posterior retraction of GFP expression than the G-carrying variant (Hikone-AS; smaller eyes) (Figures 4H and 4H'). In contrast, at early L3, after EAD subdivision is completed, the two alleles drove similar expression of GFP and mCherry (Figures 4I and 4I').

Could this differential temporal retraction of the *ey* enhancer be caused by changes in *ey* repression by Ct? To test this, we created two new synthetic *ey* enhancers, based on *in silico* predictions (Figure 4E'). The first, which we call the *NoCt* variant, is predicted to abolish Ct binding to the site harboring the G/A SNP (predicted affinity score <3). The second, which we call *ConsensusCt*, creates a Ct consensus-binding motif at that position (predicted affinity score 6.62). Remarkably, the *ConsensusCt* variant behaved similarly to the G-allele, while the *NoCt* variant mimicked the A-allele in that it caused faster posterior retraction of *ey* enhancer activity (Figures 4H–4I'). This further suggests that the Canton-S A-allele may constitute a lower affinity site for the Ct repressor as compared to the Hikone-AS G-allele.

Thus far, we showed that the changes in EAD subdivision between Hikone-AS and Canton-S and between *D. mel.* and *D. pse.* are both driven by differential temporal dynamics of the posterior retraction of *ey* expression. Between Hikone-AS and Canton-S, this is associated with a single nucleotide variant in *ey* eye enhancer, which introduces subtle changes in *ey* regulation, by affecting its repression by the antennal selector TF Ct.

A Common SNP in *D. mel.* Natural Populations Is Associated with Facet Number Variation

Because Hikone-AS and Canton-S flies have been in artificial lab culture conditions for decades, we asked if either of these two alleles is found in natural fruit fly populations. By investigating allele frequency patterns in whole-genome data of worldwide population samples, we found that most natural populations from Europe, North America, Asia, and Australia are polymorphic at this position. Thus, neither of the two alleles corresponds to a *de novo* mutation and variation at this position corresponds to a relatively frequent SNP (Figure 5A and 5A'; Table S2). Populations from sub-Saharan Africa are mostly fixed for the G-allele suggesting that the A-allele is a derived variant that appeared after *D. mel.* left Africa and colonized the rest of the world. In line with this hypothesis, we found statistical evidence (FET test, $p = 0.02$) that the few African populations carrying the A-variant are more likely to be admixed with the European genetic variation than the ones with the putatively ancestral G-allele (Table S3). Moreover, the frequency of the A-variant decreased from West to East in European populations (Figures 5A' and 5B). The slope of the longitudinal frequency cline of the *ey* SNP deviated significantly from that of 21,008 genome-wide SNPs in short introns that presumably evolved neutrally (Figure 5C), suggesting that the clinal pattern is not solely the result of neutral evolution or demography (see also Figure S7).

Causal Effect of the SNP on Eye Size

We further noted that natural populations from North-East America, where the Canton-S strain originated, are highly polymorphic for the *ey* SNP (Figure 5A; Table S2). By comparing Canton-S flies from three laboratories, we discovered that, while our Canton-S lab isolate (henceforth Canton-S^{BH}) carries the A-allele, two other strains from two different laboratories in Paris, France (T. Pr  at) and Florida, USA (R. Davis) were homozygous for the G-allele, similar to Hikone-AS. This strongly suggests that the original Canton-S population was polymorphic and that the two alleles were eventually segregated during the separate maintenance of different laboratory stocks (Colomb and Brembs, 2014). This provided a unique opportunity to quantify the contribution of the G/A SNP to eye size in a relatively homogeneous background. By comparing ommatidia numbers between the three stocks, we observed that Canton S^{BH} flies have significantly more facets than its two siblings (Figure 6A; Table 1), a difference that anti-correlates with their face and antennal width (Figure S6). These data suggest that the A-variant may be sufficient to drive larger facet numbers, possibly at the expenses of other head structures. To test this idea directly, we used CRISPR/Cas9 to introduce the A-allele in a G-homozygous stock. We recovered one transformant male carrying the A-allele and controlled that it bore no other mutation in the *ey* regulatory intron. Comparing ommatidia numbers between engineered G>A flies and the G-carrying control revealed an increase in eye size associated with the G>A substitution (Figure 6B; Table 1). It recapitulated up to 49% of the difference between Hikone-AS and Canton-S^{BH} and up to 86% of the variation observed between Canton-S^{BH} and Canton-SRD and Canton-S^{TP}, respectively. Three other A-homozygous lines deriving from the same transformant male were established to account for subtle differences in the genetic background of the flies. The three stocks followed the same trend, i.e., an increased facet number when compared to the G-homozygous controls, reaching statistical significance in two out of the three (Figure S6). This result suggests that the *ey cis*-regulatory SNP is causal to facet number variation. Next, we combined the two alleles with Df(4) J2, a large deficiency covering the entire *ey* locus. These flies recapitulated the phenotypes of the corresponding homozygous alleles, indicating that an undetected lesion outside of the *ey* locus did not cause the effect on eye size attributed to the A-allele (Figure 6C). In addition, in both cases, the antennae show a trend toward decreasing in size, but this trend did not reach statistical significance (Figure S6), likely because of the small size of the difference.

DISCUSSION

In 1987, Montgomery Slatkin proposed a mathematical model (Slatkin, 1987), which he referred to as “unrealistically simple,” predicting that mutations modifying the time at which “traits developing from the same tissue” begin to grow independently changes the relative size of the traits. Size trade-offs between head sensory organs represent precisely the types of traits referred to in Slatkin’s model. However, whether visual-olfactory sensory trade-offs follow a “Slatkin model” and if so, what the genetic basis of such a model are, remained unexplored.

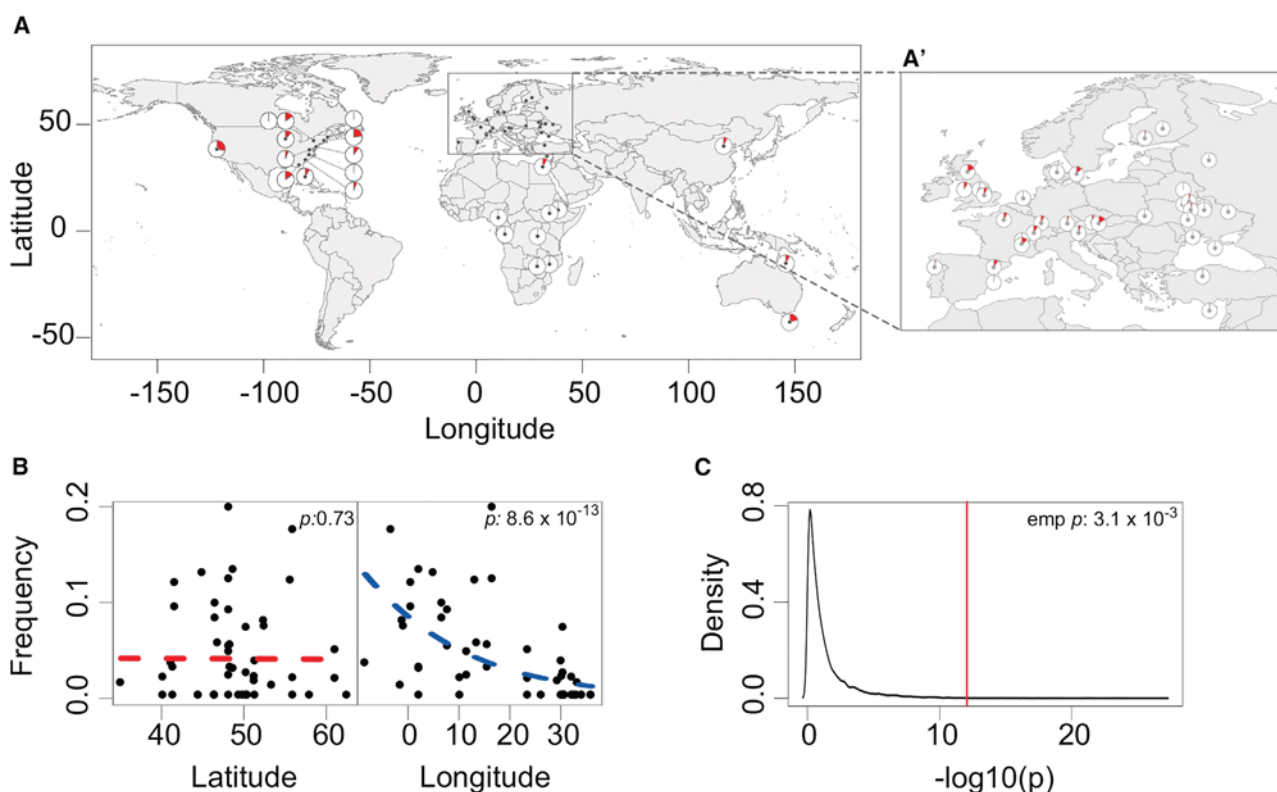


Figure 5. Worldwide Distribution of the G>A Substitution (chr4: 710326) in the Genome of *D. mel.* Natural Populations

(A) Worldwide allele frequency patterns. Frequencies of the A-variant are shown as red areas of the pie charts for worldwide populations with a sample-size ≥ 10 (see [Tables S2](#) and [S3](#) for details). The exact geographic location for each sample is indicated by a black dot.

(A') Frequency distribution of the A-variant in Europe revealing the longitudinal clinal distribution of the allele.

(B) Scatter plots showing allele frequencies of the A-variant along the latitudinal (left box; red regression curve) and the longitudinal axis (right box; blue regression curve) in Europe. The top-right p values show the significance from generalized linear models (GLM).

(C) The line plot shows the empirical cumulative density function (ECDF) from $-\log_{10}$ transformed p values of GLMs with longitude as the predictor variable for 21,008 neutrally evolving intronic SNPs (black curve). The vertical red line highlights the $-\log_{10} p$ value for the focal SNP ($p = 8.6 \times 10^{-13}$) which is inferior to the ones of 99.69% of the neutral SNPs, indicating its stronger correlation with longitude. The empirical p value ($p = 0.0031$) is calculated from the area confined by the p value of 4: 710326 and the tail of the ECDF.

See also [Figure S7](#).

In this study, we find that differential subdivision of the head primordium into eye and non-eye progenitor fields constitutes a developmental mechanism for creating different proportions of head structures in fruit flies, including trade-offs between the olfactory antennae and the eyes. We further demonstrate that this is associated with differential temporal regulation of the expression of the conserved eye selector transcription factor, *Ey/Pax6*. We propose a model ([Figures 6D–6F](#)) whereby early in development, the homogenous expression of *ey*, which promotes its proliferation ([Zhu et al., 2017](#)), causes homogenous growth throughout the entire EAD. Later, the progressive retraction of *ey* expression from the anterior antennal compartment creates an asymmetry in growth rate. Modulating the velocity of *ey* retraction through mutations affecting the bistable switch between GRNs governing antennal versus eye identity, changes the relative time during which the anterior and posterior compartments grow at different rates resulting in their different proportions. This provides direct biological evidence for mathematical models linking heterochrony in development to changes in adult traits ([Riska, 1986](#); [Slatkin, 1987](#); [Cowley and Atchley, 1990](#)). Our

observation of similar changes in *ey* temporal regulation between and within species further suggests that the temporal mechanism we identify may represent a “preferred route” to relative variation in sensory organ size because it results in no dramatic “pleiotropic” effects associated with changes to growth, patterning, or specification. We speculate that such variation of the temporal regulation of EAD partitioning could be caused by a variety of molecular changes acting either in *cis* or in *trans* on the GRNs governing eye and antennal compartment identities.

In this work, we uncovered an example of such a molecular change to the regulation of these GRNs. Specifically, within *D. mel.*, a naturally occurring SNP in the eye-enhancer of *ey* is sufficient to modulate the velocity of the posterior retraction of the enhancer activity and to vary facet numbers in the adult eye. The SNP, a G/A substitution, is located in a binding site for *ey* repressor Ct. *In-silico* predictions and EMSA experiments indicate that the G/A substitution modifies the affinity of Ct for the binding site. This suggests that different levels of *ey* repression by Ct are responsible for changing the dynamics of *ey* regulation

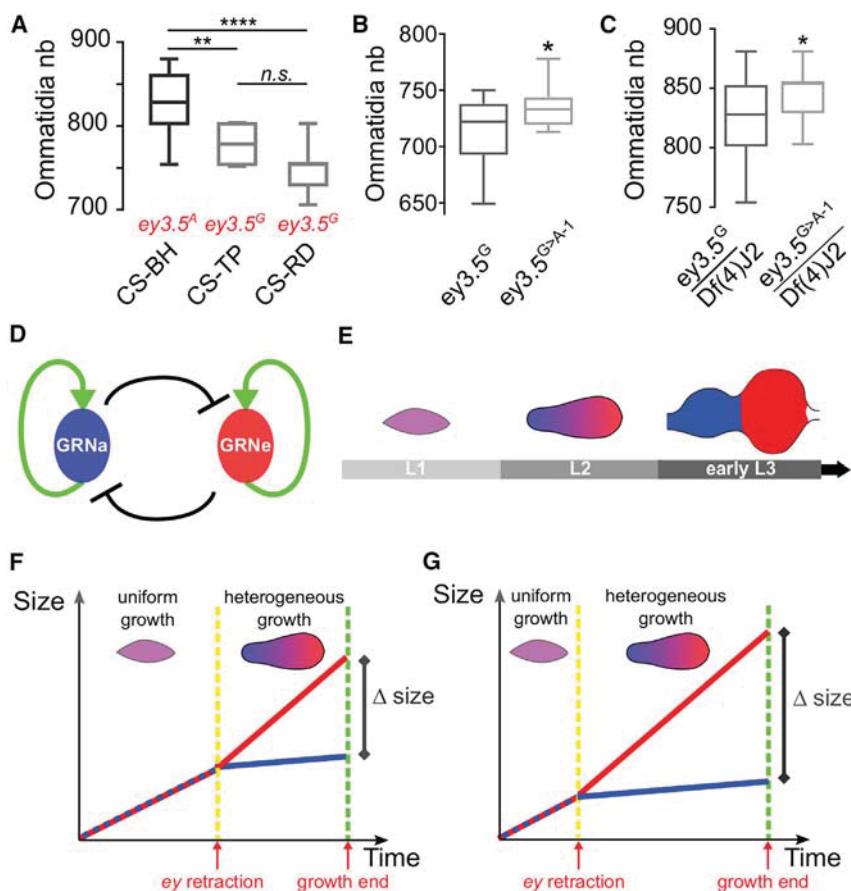


Figure 6. The Non-coding G>A SNP in *ey* Enhancer Causes Facet Number Variation

(A) Ommatidia numbers in three Canton-S strains with different *ey* SNP alleles (in red). Sample sizes: Canton-S^{BH} (CS-BH, n = 18), Canton-S^{TP} (CS-TP, n = 16), Canton-SRD (CS-RD, n = 19); Kruskal Wallis test ****p < 0.0001 followed by Dunn's multiple comparisons: **adjusted p = 0.0041; **** adjusted p < 0.0001; n.s. adjusted p = 0.1180. See also Figure S6.

(B) Ommatidia numbers in CRISPR A-variant and control G-variant homozygous fly eyes imaged by scanning electron microscopy. Sample sizes: *ey3.5^G*: n = 13; *ey3.5^{G>A-1}*: n = 12. Two-tailed unpaired t test: *p = 0.0356. See also Figure S6.

(C) Ommatidia numbers in CRISPR A-variant and control G-variant over a deficiency covering the entire *ey* locus. Sample sizes: n = 18. Two-tailed unpaired t test with Welch's correction: *p = 0.0138. See also Figure S6.

(A and B) Boxes indicate interquartile ranges, lines medians, and whiskers data ranges.

(C–F) Model of the developmental origin of the trade-off between EAD derived structures.

(C) The subdivision of the EAD into an anterior (“antennal”) and posterior (“eye”) compartments involves a bistable switch by which GRNs promoting eye (GRNe, in red) and antennal (GRNa, in blue) identity act antagonistically by activating their own and repressing the alternative GRN's activity.

(D) The bistable switch between the GRNa and GRNe results in the progressive segregation of the expression domains of TFs promoting eye (posterior, in red) versus antennal (anterior, in blue) fates during successive stages of EAD development. Anterior is on the left.

(E and F) Our model, inspired from Slatkin (Slatkin, 1987), proposes that different temporal dynamics of the posterior retraction of *ey*, a promotor of EAD proliferation, by changing the relative duration of uniform versus heterogeneous growth, modifies the proportion between the antennal and eye compartments. This could be caused by genetic changes affecting the dynamics of the bistable switch between GRNe and GRNa, like in the case of the *ey* G>A substitution.

and ultimately for causing morphological variation, a lower affinity of the binding site for Ct resulting into faster enhancer retraction and larger eyes and vice versa. This view is further supported by our findings that (1) synthetic mutations predicted to support or reduce Ct binding mimic the effect of the SNP on the velocity of the enhancer retraction; and that (2) knocking down Ct expression increases compound eye size. Interestingly, the highly divergent *ey* enhancers from *D. mel.* and *D. pse.* display similar differences in regulation, i.e., a faster retraction of the enhancer of the “larger eye” species. However, what feature of the two enhancer sequences causes their different temporal regulation is not known. A recent study identified a single nucleotide insertion that influences photoreceptor specification and ultimately color preference in *Drosophila* by modulating the affinity for a TF named Klumpfuss (Anderson et al., 2017). Together with our work, this suggests a general role for variation in suboptimal TF binding sites in sensory evolution (Crocker et al., 2016).

In vertebrates, antagonistic relationships between GRNs and signaling pathways that promote different sensory identities also regulate the subdivision of the multipotent sensory placode (Grocott et al., 2012; Singh and Groves, 2016). The anterior placodal region, which gives rise to the lens and olfactory placodes, expresses *pax6*. In the absence of *pax6*, both lens and olfactory placodes fail to thicken and to develop properly (Quinn et al.,

1996; Ashery-Padan et al., 2000; Collinson et al., 2000). Interestingly, the temporal regulation of the *pax6* placode enhancer is altered by manipulating suboptimal binding sites for one of its activators (Rowan et al., 2010). In addition, the *pax6* ectodermal enhancer shows evidence of accelerated evolution in subterranean mammalian species (Partha et al., 2017). This is consistent with our model whereby naturally occurring mutations that alter *pax6* regulation, either in *cis* or in *trans*, constitute a common genetic origin of the trade-offs between visual and olfactory organ size.

In *The Origin of Species* Charles Darwin referred to the evolution of the eye as a challenge to his theory (Darwin, 1872). He also discussed the importance of correlation between body parts concluding that it was “most imperfectly understood.” During the last decades, the common origin of animal eyes and their evolution over long evolutionary distances has been abundantly documented (Gehring, 2014). However, the developmental mechanisms by which small-scale variation in eye size or shape can take place without disrupting its organization and function remain largely elusive (Dyer et al., 2009). We have demonstrated that a single nucleotide change in a core regulator of eye development is sufficient to generate reciprocal sensory organ size variation, potentially providing a quick route to behavioral changes and perhaps adaptation. As predicted by Darwin,

adaptive variation in head derived structures, including the eye, can be produced by the accumulation of modest morphological changes, which our data suggest may be caused by a small number of genetic variants affecting the temporal regulation of core regulatory networks.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at <https://doi.org/10.1016/j.devcel.2019.07.027>.

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AUTHOR CONTRIBUTIONS

A.R. conceived the project. A.R. and B.A.H. designed the experiments. A.R. and J.Y. acquired SEM and light microscopy images. A.R., N.G., and S.W. performed and imaged immunostainings on imaginal disks, and A.R. and N.G. quantified and analyzed the imaging data. A.R. and A.C. generated the constructs and performed allele-specific PCR. J.Y. injected the CRISPR constructs. E.B. and R.W. performed and analyzed the optomotor tests. M.K. and T.F. performed population genetics analyses. D.P. performed Ct binding sites *in silico* predictions and synthetic enhancers design. D.C. and E.M.V. performed the EMSAs. A.R. and B.A.H. wrote the manuscript.

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-Drosophila Cut protein product Monoclonal Antibody, Unconjugated (1:10)	DHSB	DSHB Cat# 2b10; RRID:AB_528186
Mouse anti-Eyes Absent (Eya) protein antibody (1:75)	DHSB	DSHB Cat# eya10h6; RRID:AB_528232
Rat-Elav-7E8A10 anti-elav monoclonal antibody (1:100)	DSHB	Cat# 7E8A10; RRID: AB_528218
Rat anti-Eyeless (Ey) polyclonal antibody (1:300)	Laboratory of P. Callaerts	Halder et al. (1998)
Rabbit PhosphoDetect Anti-Histone H3 (pSer10) (7-20) Rabbit pAb antibody (1:1000)	Millipore	Cat# 382159-50UG; RRID:AB_565299
Sheep anti-Atonal (Ato) polyclonal antibody (1:250)	Laboratory of A. Jarman	Jarman AP; Development. 1995 Cat# ato; RRID:AB_2568143
Mouse Anti-Green Fluorescent Protein (GFP) Monoclonal Antibody, Unconjugated, Clone 3E6 (1:1000)	Molecular Probes	Molecular Probes Cat# A-11120; RRID:AB_221568
Rabbit Living Colors® DsRed Polyclonal Antibody (1:1000)	Takara Bio	Takara Bio Cat# 632496; RRID: AB_10013483
Donkey Anti-Rat IgG (H+L) Antibody, Alexa Fluor 488 Conjugated (1:200)	Thermo Fisher Scientific	Thermo Fisher Scientific Cat# A-21208; RRID:AB_141709
Goat Anti-Mouse IgG (H+L) Highly Cross-adsorbed Antibody, Alexa Fluor 488 Conjugated (1:200)	Thermo Fisher Scientific	Thermo Fisher Scientific Cat# A-11029; RRID:AB_138404
Donkey Anti-Mouse IgG (H+L) Antibody, Alexa Fluor 488 Conjugated (1:200)	Thermo Fisher Scientific	Thermo Fisher Scientific Cat# A-21202; RRID:AB_141607
Goat Anti-Rabbit IgG (H+L) Antibody, Alexa Fluor 488 Conjugated (1:200)	Thermo Fisher Scientific	Thermo Fisher Scientific Cat# A-11008; RRID:AB_143165
Donkey anti-Rabbit IgG (H+L) Highly Cross-Adsorbed Secondary Antibody, Alexa Fluor 555 (1:200)	Thermo Fisher Scientific	Thermo Fisher Scientific Cat# A-31572; RRID:AB_162543
Goat anti-Rat IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor 555 (1:200)	Thermo Fisher Scientific	Thermo Fisher Scientific Cat# A-21434; RRID:AB_2535855
F(ab)2-Goat anti-Mouse IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor 555 (1:200)	Thermo Fisher Scientific	Thermo Fisher Scientific Cat# A-21425; RRID:AB_2535846
Goat anti-Guinea Pig IgG (H+L) Highly Cross-Adsorbed Secondary Antibody, Alexa Fluor 555 (1:200)	Thermo Fisher Scientific	Thermo Fisher Scientific Cat# A-21435; RRID:AB_2535856
Monoclonal ANTI-FLAG® M2 antibody (1:200)	Sigma-Aldrich	Sigma-Aldrich Cat# F3165; RRID: AB_259529
Chemicals, Peptides, and Recombinant Proteins		
Qiagen Effecten Transfection reagent	Qiagen	Qiagen Cat# 301425
Vectashield	Vector Laboratories	Vector Laboratories Cat#H-1000
Critical Commercial Assays		
Qiagen DNeasy Blood and Tissue Kit	Qiagen	Qiagen Cat#69504
pENTR-D-TOPO kit	ThermoFischer Scientific	ThermoFischer Scientific Cat#K2400-20

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
DEST™-BLASTICIDIN SUPPORT KIT	ThermoFischer Scientific	ThermoFischer Scientific Cat#K515001
LightShift™ Chemiluminescent EMSA Kit	ThermoFischer Scientific	ThermoFischer Scientific Cat# 20148
Deposited Data		
<i>D. melanogaster</i> Reference Genome BDGP release 6	Berkeley Drosophila Genome Project	GCA_000001215.4
<i>D. pseudoobscura</i> release Dpse_3.0	Baylor College of Medicine	GCA_000001765.2
<i>Drosophila</i> Genome Nexus	http://www.johnpool.net/genomes.html	N/A
<i>D. melanogaster</i> European genomes	European Drosophila Population Genomics Consortium (DROSEU)	PRJNA388788
Experimental Models: Cell Lines		
<i>D. melanogaster</i> : Cell line S2: S2-DRSC	Laboratory of Norbert Perrimon	FlyBase: FBtc0000181
Experimental Models: Organisms/Strains		
<i>D. melanogaster</i> ; Canton-S ^{BH}	Laboratory of B. Hassan	N/A
<i>D. melanogaster</i> ; Canton-S ^{TP}	Laboratory of P. Callaerts	N/A
<i>D. melanogaster</i> ; Canton-S RD	Laboratory of P. Callaerts	N/A
<i>D. melanogaster</i> ; Hikone-AS	Kyoto Stock Center DGGR	RRID:DGGR_105668
<i>D. melanogaster</i> ; DGRP-208	Bloomington Drosophila Stock Center	RRID:BDSC_25174
<i>D. melanogaster</i> ; y ¹ M{w ^{+mC} =Act5C-Cas9.P}ZH-2A w*	Bloomington Drosophila Stock Center	RRID:BDSC_54590
<i>D. melanogaster</i> ; w ¹¹¹⁸ ; P{y ^{+t7.7} w ^{+mC} =GMR33F07-GAL4}attP2/TM3, Sb ¹	Bloomington Drosophila Stock Center	RRID:BDSC_45603
<i>D. melanogaster</i> ; w ¹¹¹⁸ ; P{w+mW.hs=GawB}hthNP5332 / TM6C, Sb1	Kyoto Stock Center DGGR	RRID:DGGR_104957
<i>D. melanogaster</i> ; UAS-RNAi ^{ct}	Vienna Drosophila Resource Center	VDRC Cat#v5687
<i>D. melanogaster</i> ; UAS-RNAi ^{ct}	Vienna Drosophila Resource Center	VDRC Cat#v5687
<i>D. melanogaster</i> ; y ¹ v ¹ ; P{y ^{+t7.7} v ^{+t1} .8=TRiP.JF01355}attP2	Bloomington Drosophila Stock Center	RRID:BDSC_31603
<i>D. melanogaster</i> ; <i>D.m.ey3.5^GGFP</i>	Laboratory of B. Hassan	this study
<i>D. melanogaster</i> ; <i>D.m.ey3.5^AGFP</i>	Laboratory of B. Hassan	this study
<i>D. melanogaster</i> ; <i>D.m.ey3.5^{NoCt}GFP</i>	Laboratory of B. Hassan	this study
<i>D. melanogaster</i> ; <i>D.m.ey3.5^{ConsensusCt}GFP</i>	Laboratory of B. Hassan	this study
<i>D. melanogaster</i> ; <i>D.m.ey3.5^Gmcherry</i>	Laboratory of B. Hassan	this study
<i>D. melanogaster</i> ; <i>D.m.ey3.5^Amcherry</i>	Laboratory of B. Hassan	this study
<i>D. melanogaster</i> ; <i>ey3.5^G</i>	Laboratory of B. Hassan	this study
<i>D. melanogaster</i> ; <i>ey3.5^{G>A-1}</i>	Laboratory of B. Hassan	this study
<i>D. melanogaster</i> ; <i>ey3.5^{G>A-2}</i>	Laboratory of B. Hassan	this study
<i>D. melanogaster</i> ; <i>ey3.5^{G>A-3}</i>	Laboratory of B. Hassan	this study
<i>D. melanogaster</i> ; <i>ey3.5^{G>A-4}</i>	Laboratory of B. Hassan	this study
<i>D. ananassae</i> ; isofemale WT	Drosophila Species Stock Center	DSSC Cat#14024-0371.30
<i>D. ananassae</i> ; isofemale WT	Drosophila Species Stock Center	DSSC Cat#14024-0371.31
<i>D. yakuba</i> ; WT	Drosophila Species Stock Center	DSSC Cat#14021-0261.00
<i>D. yakuba</i> ; WT	Drosophila Species Stock Center	DSSC Cat#14021-0261.01
<i>D. pseudoobscura</i> ; isofemale WT	Drosophila Species Stock Center	DSSC Cat#14011-0121.121
<i>D. pseudoobscura</i> ; isofemale WT	Drosophila Species Stock Center	DSSC Cat#14011-0121.118
<i>D. virilis</i> ; isofemale WT	Drosophila Species Stock Center	DSSC Cat#15010-1051.118
Oligonucleotides		
Primers, see Table S5	Laboratory of B. Hassan	this study

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Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Recombinant DNA		
Gateway™ pDONR™221 Vector	ThermoFischer Scientific	ThermoFischer Scientific Cat#12536017
pAWF vector	https://emb.carnegiescience.edu/Drosophila-gateway-vector-collection	DGRC Cat#1111
pStingerGFP vector	Aerts et al. 2010 ; Quan et al. 2016	N/A
pStingerMCherry vector	Aerts et al. 2010 ; Quan et al. 2016	N/A
pU6-BBS1-chiRNA vector	Gratz et al. 2013	Addgene #Cat45946
pU6gRNA ^{ey}	Laboratory of B. Hassan	this study
pAWF-Cut	Laboratory of B. Hassan	this study
pEntry-ey ^{3.3Pse}	Laboratory of B. Hassan	this study
pEntry-ey ^{3.5CSBH}	Laboratory of B. Hassan	this study
pEntry-ey ^{3.5Hik}	Laboratory of B. Hassan	this study
pEntry-ey ^{3.5NoCt}	Laboratory of B. Hassan	this study
pEntry-ey ^{3.5consensusCt}	Laboratory of B. Hassan	this study
pStingerGFP-ey ^{3.3Pse}	Laboratory of B. Hassan	this study
pStingerGFP- ^{3.5CSBH}	Laboratory of B. Hassan	this study
pStingerGFP- ^{3.5Hik}	Laboratory of B. Hassan	this study
pStingerGFP- ^{3.5NoCt}	Laboratory of B. Hassan	this study
pStingerGFP- ^{3.5consensusCt}	Laboratory of B. Hassan	this study
pStingerMCherry- ^{3.5CSBH}	Laboratory of B. Hassan	this study
pStingerMCherry- ^{3.5Hik}	Laboratory of B. Hassan	this study
Software and Algorithms		
Prism 8	Graphpad Software Inc.	https://www.graphpad.com/scientific-software/prism/
FIJI	Schindelin et al. 2012	http://fiji.sc/
ImageJ	Schneider et al. 2012	https://imagej.nih.gov/ij/
Other		
Code for mcherry/GFP colocalization macro	Oliva et al. 2016 .	https://github.com/rejsmont/FijiScripts/blob/master/mColoc3D.py

LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Bassem Hassan (bassem.hassan@icm-institute.org). Reagents generated during this study are available upon request from the authors.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Species

Drosophila Melanogaster

D. melanogaster stocks were cultured on standard cornmeal diet food at 25°C except mentioned otherwise in the corresponding figure legend. Developmental stages used in each experiment are indicated in the corresponding [Method Details](#) section. Morphological measures on adult flies were performed on females only. The fly strains used in this study were: Canton-S^{BH} (Hassan lab); Canton-S^{TP} (Pr  at lab, provided by P. Callaerts); Canton-SRD (Davis lab, provided by P. Callaerts); Hikone-AS (Kyoto DGRC 103421); DGRP-208 (*D.M.* 3, Bloomington #25174); Act5C-Cas9 ([Port et al., 2014](#)) (Bloomington # 54590); *ct-Gal4* ([Pfeiffer et al., 2008](#)) (Bloomington # 45603); *hth*^{NP5332}-Gal4 (DGRC Kyoto #104957); UAS-RNAi^{ct} (VDRC #v5687); UAS-RNAi^{ct} (VDRC #v4138); UAS-RNAi^{luc} (Bloomington #31603); *D.m.ey3.5^GGFP* (this study); *D.m.ey3.5^AGFP* (this study); *D.m.ey3.5^GmCherry* (this study); *D.m.ey3.5^AmCherry* (this study); *D.m.ey3.5^{NoCt}GFP* (this study); *D.m.ey3.5^{ConsensusCt}GFP* (this study); *ey3.5^G* (this study); *ey3.5^{G>A-1}* (this study); *ey3.5^{G>A-2}* (this study); *ey3.5^{G>A-3}* (this study); *ey3.5^{G>A-4}* (this study).

Drosophila Ananassae

Isofemale *D. ananassae* stocks were cultured on standard cornmeal diet food at 21°C. Morphological measures on adult flies were performed on females only. Stock origin: Kisangani, Congo (DSSc 14024–0371.30) and Mumbai, India (DSSc 14024–0371.31).

Drosophila Yakuba

D. yakuba stocks were cultured on standard cornmeal diet food at 21°C. Morphological measures on adult flies were performed on females only. Stock origin: Ivory Coast (DSSc 14021-8209.0261.00) and Liberia (Reference Genome strain DSSc 14021-0261.01).

Drosophila Pseudoobscura

Isofemale *D. pseudoobscura* stocks were cultured on standard cornmeal diet food at 25°C except when mentioned otherwise in the corresponding figure legend. Morphological measures on adult flies were performed on females only. Stock origin: Catalina Island, California, USA (Cat; DSSc 14011-0121.121) and Chiracahua Mountains, Arizona, USA (DSSc 14011-0121.118).

Drosophila Virilis

The isofemale *D. virilis* stock was cultured on standard cornmeal diet food at 21°C. Morphological measures on adult flies were performed on females only. Stock origin: Gikongoro, Rwanda (DSSc 15010-1051.118).

Developmental Stages

For selecting specific developmental stages, embryos were collected on grape fruit plates complemented with yeast paste changed every 2 hours. Freshly hatched L1 larvae were collected every two hours and transferred to corn meal food vials in a density-controlled fashion (20 larvae / vial). Staging was performed at 25°C. Correspondence of developmental stages between *D. mel.* and *D. pse.* was determined based on developmental transitions – larval hatching, L2 to L3 molt, pupa formation - and morphological features – embryo morphology, rows of differentiated photoreceptors in the EAD, size of the EAD.

Density-Controlled Culture Conditions

Morphological measurements were performed on flies raised in density-controlled conditions: batches of 20 young females and males (2-5 days old) were put together and cultured at 25°C. They were transferred in fresh vials every 24 hrs. For each vial, the individuals eclosing during the first two days of eclosion only were used for measurements.

METHOD DETAILS

Generation of Reagents

Constructs

Enhancer reporter constructs were generated using the Gateway Recombination Cloning Technology (ThermoFischer Scientific). *D. pse.* *ey3.3* and *D. mel.* *ey3.5* regulatory sequences were amplified respectively from *D. pse.* (from stock Cat; DSSc 14011-0121.121), Hikone-AS (for the G-variant) and Canton-S^{BH} (for the A-variant) genomic DNA (extracted using Qiagen DNeasy Blood and Tissue Kit #69504) and cloned into the Gateway pDONR221 entry vector (ThermoFischer Scientific #12536017) following the provider specifications. Primers for the enhancer amplifications are:

pEntry-*ey*^{3.3Pse}: *forward*:GGGGACAAGTTTGTACAAAAAAGCAGGCTAAGTGGTAGTGGACTAGG and *reverse*:GGGGACCACTTTGTACAAGAAAGCTGGGTCCTAGAAATTTTGCTAACGC;

pEntry-*ey*^{3.5CSBH} and pEntry-*ey*^{3.5Hik}:

forward:

GGGGACAAGTTTGTACAAAAAAGCAGGCTGGACTAGGCGGTATTGCT and *reverse*:GGGGACCACTTTGTACAAGAAAGCTGGGTTTGCTCACACATCCATTTG. The entry vectors with mutated forms of *ey3.5* enhancer, pEntry-*ey*^{3.5NoCt} and pEntry-*ey*^{3.5consensusCt} were generated by modifying the pEntry-*ey*^{3.5Hik} using primers carrying the corresponding mutations. These primers were (mutated nucleotides are in capital letters):

pEntry-*ey*^{3.5NoCt} :

forward: caataaaatggttg**Ca**Gtttttcgaactttcg

reverse: cgaagttcgaaaa**Ct**Gccaaccattttattg

pEntry-*ey*^{3.5consensusCt} :

forward: taaaatggt**T**gaacttttcgaactttcg

reverse: gaaaaagtt**A**aaccattttattg

Enhancer inserts were next transferred using Gateway recombination into mCherry- and GFP-expressing enhancer reporter vectors amenable to phiC31 integration –mediated transgenesis (Aerts et al., 2010; Quan et al., 2016).

pAWF-Cut: ct cDNA was kindly provided by I. Lohmann (U. Heidelberg). The full length cDNA was cloned, without its ATG, into a Gateway pEntry vector using the pEntr-D-TOPO kit (ThermoFischer Scientific K2400-20) following provider specifications. It was transferred using Gateway recombination into the pAFW vector (DGRC#1111), resulting in the addition of 3 x Flag tag coding sequence upstream ct cDNA.

pU6gRNA^{ey}: the following complementary phospho-oligomers were used to generate a double strand DNA sequence encoding the *ey* eye-enhancer guide RNA (gRNA): *forward*: phospho-CTTCGTCGAAAACAATAAAATGGT; *reverse*: phospho-AAACACCATTTTATTGTTTTCGAC. After hybridization, the resulting double-strand DNA was cloned into the pU6-BBS1-chiRNA plasmid (Addgene #45946)(Gratz et al., 2013).

Enhancer-Reporter Lines

Transformant flies carrying enhancer reporter constructs were generated by BestGeneInc. All constructs were integrated at the Attp2 landing site using phiC31 recombination.

CRISPR/Cas9 Engineering

For editing the *ey* eye enhancer, we injected SNP^G homozygous *D. mel.* Act5-Cas9 embryos (Port et al., 2014) with two constructs respectively encoding the guide RNA (pU6gRNA^{ey}) and the SNP^A-carrying *ey* eye enhancer sequence (pEntry-*ey*^{3.5CSBH}), each of them at a concentration of 500 ng/μl (Port et al., 2015). Candidates were screened using allele-specific PCR. We isolated one CRISPR modified male from which we established four CRISPR SNP^A lines. In parallel, a control line was established by mating non-injected Act-Cas9 flies following the same scheme their injected siblings. Sequencing the *ey* eye enhancer from the transformed SNP^A and of the non-injected SNP^G control stocks confirmed that they were differing only by this single nucleotide.

Allele-Specific PCR

SNP^A and SNP^G alleles were detected by allele-specific PCR using a common reverse primer (Ey-R3: AGAAATATCACATGGCCGAG) and one of two specific forward primers differing by the 3' most nucleotide (either A or G) and including a mismatch (underlined) to increase binding specificity (Ey-SNP^G-F: GGAATCGAAAACAATAAAATGGCTGG; Ey-SNP^A-F: GGAATCGAAAACAATAAAATG GCTGA).

Cut-FLAG Fusion Protein Expressing S2 Cells

S2 cells cultivated at 25°C in Schneider's medium supplemented with 10% fetal calf serum were co-transfected with 2μg of pAWF-Cut and 0,2 μg of pCoBlast vector (Thermo Fisher Scientific) using Effecten transfection reagent (Qiagen) according to the manufacturer's instructions (1/10 DNA-Effecten ratio). Blasticidin selection (10μg/μl) was applied three days after transfection. After one week of selection, cells were harvested, and expression of Cut-FLAG fusion protein was checked by western-blot using an anti-FLAG antibody (Sigma-Aldrich Cat# F3165). For long-term culture, cells were maintained in 2μg/μl blasticidin.

Imaging and Image Quantification

Image Processing

Except mentioned otherwise, all image processing was performed using ImageJ (versions 1.45 to 1.48)(Schneider et al., 2012).

Scanning Electron Microscopy

Whole flies were fixed overnight at 4°C in a 1:1 mix of 4% formaldehyde in phosphate buffer pH 7.2 and 100 % ethanol and dehydrated successively in graded ethanol series, hexamethyldisilazane (HMDS) and in a dessicator. Fly heads were mounted on specimen studs using silver paint in two distinct orientations: dorsal head up (for whole head imaging) and lateral (for ommatidia counts and measures). Samples were subsequently coated with platinum and images acquired in LEI mode with a JEOL JSM 7401F microscope at magnifications ranging from 120 (heads overviews) to 1900 times (ommatidia width) (Schneider et al., 2012).

Transmitted Light and Confocal Microscopy

Preparations of adult heads for the acquisition of light microscopy images were acquired from non-fixed, freshly cut adult heads glued laterally on glass slides. Images were acquired using a camera DFC295 (Leica) mounted on a DMRXA (Leica) microscope, operated via the open-source software Micro-Manager (Edelstein et al., 2014). Fluorescent preparations of embryos and imaginal discs were acquired using a Nikon A1R Eclipse Ti, a Leica TCS SP5 II or a Leica SP8 confocal microscope operated by the accompanying company software.

Measuring Adult Eye, Face and Antennae

All head and eye measurements were performed on female flies using ImageJ (Schneider et al., 2012). Adult Eye:Face ratio was expressed as E(Figure 1). Ommatidia width was measured on high magnification SEM images as the distance between one interommatidial bristle and the opposite angle of the facet. For each sample, measures of six adjacent ommatidia localized at the center of the eye were taken. To limit underestimation of the ommatidia width due to perspective projection distortion, samples were carefully oriented prior to image acquisition.

Ommatidia Numbers

Ommatidia numbers were manually counted on SEM images using the ImageJ plugin "Cell counter". We also developed an alternative method based on the approximation of the compound eye to an ellipse. With this method, the ommatidia number is calculated as the surface of an ellipse whose large and small axes correspond to the numbers of ommatidia along the compound eye anterior-posterior and dorso-ventral axes ($Area = \pi.a.b/4$ with a and b as the lengths of the large and small ellipse axes; Figure S4). This method accommodates lower resolution images and does not require the use of SEM. Bland-Altman method (Bland and Altman, 1986) was used to compare the outcome of the two methods applied on a common set of SEM images. The ellipse method results in an overestimation of approximately 20 ommatidia as compared to the manual counting (bias mean = 20. 29; SD = 11. 70). Importantly, this difference is independent of ommatidia number (Figure S2). Facet number estimation of Cut RNAi and CRISPR flies (Figures 4, 6, S5, and S6) were performed blind regarding to the genotype.

Measuring Embryonic and Larval EADs

Numbers of Ey-positive embryonic eye-antennal disc cells were counted manually. To measure the surface of the larval eye-antennal disc and eye progenitor field, regions of interest were selected manually using the ImageJ freehand selection tool. The number of mitotic pH3-positive cells was automatically counted using the Dead-Easy Mito-Glia ImageJ Plugin (Forero et al., 2010). The mitotic index was calculated as the number of mitotic cells per surface of the Eya-positive eye progenitor field.

MCherry and GFP Colocalization

Protocol for pixel-based quantifications of mCherry and GFP colocalization was adapted from (Oliva et al., 2016). We used Fiji/ImageJ2/ImgLib2 (Pietzsch et al., 2012; Rueden et al., 2017; Schindelin et al., 2012) macro implemented in Jython. Raw images were imported using BioFormats library (Linkert et al., 2010). EADs were manually segmented in each stack by the user. Stack

threshold levels for each channel were calculated using preselected auto-thresholding algorithms available in Fiji (Huang for both channels). Determined threshold levels were used to calculate Mander's overlap coefficient using Fiji implementation of the colocalization algorithm. Code for the macro is available on GitHub, <https://github.com/rejsmont/FijiScripts/blob/master/mColoc3D.py> (Oliva et al., 2016). The proportion of pixels expressing solely mCherry is used as a measure for GFP retraction.

Behavioral Measures of Visual Acuity. The experimental set-up is modified from (Buchner, 1976). It exploits the spontaneous tendency of fruit flies to adjust their trajectory to the surrounding landscape. Presented with rotating vertical black stripes, tethered flies spontaneously follow their movement. Narrowing the angular distance between the stripes beyond its resolving capacities makes the fly move in the opposite direction, due to an interference phenomenon similar to what we perceive when looking at the wheels of a starting train. It consists of two tracking balls and of two computer screens on which moving vertical black and white stripes are displayed in a window of 90° horizontal and 74° vertical extensions. The width of the stripes (spatial wavelength λ) as they move on the flat screen are adjusted such that they subtend a constant angle as seen from the fly 40 mm away of the screen. Pattern speed w is adapted to maintain the "contrast frequency" at 1 Hz. A positive optomotor response indicates the tendency of the flies to follow the direction of the movement of the stripes. Reduction of λ below the resolving power of the eye causes an inversion of the apparent direction of the movement of the stripes due to the geometrical interference between the fly's vertical columns of ommatidia and the vertical stripes and is accompanied by the inversion of the fly response towards negative values. The λ value at which the response of the fly is inverted (zero-crossing angle, $2\Delta\Phi$) provides a measure of spatial resolution (or visual acuity). Recordings of female *D. mel.* (Canton-S; n=9) and *D. pse.* (Cat; n=9) were performed simultaneously, with alternating assignments. We calculated the zero-crossing and its variance from the two average responses surrounding the zero-crossing (one positive, one negative) using linear interpolation and error propagation followed by t-test for differences between 2 means.

Immunostainings

Antibodies

We used the following primary antibodies: mouse anti-Ct (1:10, DHSB hybridoma supernatant 2B10, deposited by Rubin G. M.), mouse anti-Eya (1:75, DHSB hybridoma supernatant eya10H6, deposited by Benzer, S. and Bonini, N.M.), rat anti-elav (1:100, DHSB hybridoma supernatant Rat-Elav-7E8A10, deposited by Rubin G. M.), mouse anti-futsch (1:100; DHSB hybridoma supernatant 22C10 deposited by Benzer, S. and Colley, N.); rat anti-ey (1:300, received from P. Callaerts (Halder et al., 1998)); anti-phosphorylated histone 3 (1:1000, pSer10; Merck Millipore #382159); sheep anti-atonal (1:1000, received from A. Jarman and P. zur Lage (Jarman et al., 1995)); mouse anti-GFP 3E6 (1:1000, Invitrogen, #A11120); rabbit anti-DsRed (1:1000, Clontech, #632496). Secondary antibodies conjugated with Alexa 488, Alexa 555 and Alexa 647 were used at 1:200 (Molecular Probes). Nuclei were counterstained using Draq-5 (1:1000 in PBS, Abcam #ab108410). All samples were mounted in Vectashield (Vector laboratories #H-1000).

Procedure

Fixation and immunostainings were performed following standard procedures as described in (Patel, 1994) (embryos) and in (Blair, 2000) (EADs). Briefly, embryos were collected from grape juice agar plates and dechorionated with bleach (sodium hypochlorite 3%). They were fixed in a 1/1 mixture of n-heptane and 3.7% formaldehyde in PBS and devitellinized with methanol. Larval and pupal mouth complexes including the EADs, brain and pharynx were dissected in cold PBS and fixed in 3.7% formaldehyde in PBS for 20 minutes, then washed in PBT (embryos: PBS tween 0,1%; L1 and L2 EADs: PBS Triton-X 0,1%; L3 EADs: PBS Triton-X 0,3%) for 2 hours. Blocking was performed 1 hour at room temperature in blocking solution (5% normal goat serum in PBT) and incubation with primary antibodies diluted in blocking solution overnight at 4°C. After 2 hours of washes, tissues were incubated 2 hours at room temperature with the secondary antibodies diluted in blocking solution. After two hours of washes in PBT, tissues were rinsed in PBS, incubated with Draq5 (1/500 in PBS) 1 hour at room temperature or overnight at 4°C and mounted in Vectashield (Vector laboratories cat#H-1000). DAPI stainings were performed by adding DAPI in the mounting medium (final concentration 1,5 μ m).

In Silico Analysis

Genome Assemblies

All *D. mel.* Genome positions refer to BDGP release 6 assembly (GCA_000001215.4) and *D. pse.* Genome positions refer to genome release Dpse_3.0, Baylor College of Medicine (GCA_000001765.2).

Alignment of *D. mel.* *D. pse.* ey Enhancer

Pairwise alignment of *D. pse.* and *D. mel.* intronic ey eye enhancers was performed with BLASTn (McGinnis and Madden, 2004) (blast.ncbi.nlm.nih.gov) as described in (Swanson et al., 2011). Namely, the following parameters were selected: "somewhat similar sequences" and the lower complexity filter was off. The sequences used for the alignment were: for *D. pse.* chrU:2,848,595-2,851,564 and for *D. mel.* chr4:707,672-710,917. The 22 % of alignment corresponds to "query coverage" in BLASTn output and indicates the amount of *D. mel.* intron sequence included in blocks aligned to the *D. pse* ortholog by BLASTn.

Ct Binding Site Predictions

To predict possible Ct binding sites, the SNP surrounding region (500up and down) was scored with Cluster-Buster (Frith et al., 2003) (<https://github.com/weng-lab/cluster-buster> version, options -c0 -m3 -f5 -G0) using the 3 available *Drosophila* Ct PWMs (MA0128.1 from JASPAR (Mathelier et al., 2014), Ct_Cell_FBgn0004198 and Ct_SOLEXA_FBgn0004198 from FlyFactorSurvey (Zhu et al., 2011) for the 5 different alleles (Reference, Hikone-AS, Canton-S, NoCt, ConsensusCt). Predicted Binding sites overlapping our SNP are shown with either their corresponding score in Figure 4 for the best scoring PWM (Ct_SOLEXA_FBgn0004198), and in Figure S5 for all PWMs. Predictions for the entire region are shown in Table S2.

Population Genetics

We compiled whole genome sequencing data from multiple geographic samples collected in Africa, Europe, North America, Asia and Australia to investigate worldwide allele frequency patterns of the *eyeless* SNP at position Chr 4: 710326 in natural populations. This dataset consisted of single individual sequencing data (Campo et al., 2013; Grenier et al., 2015; Lack et al., 2015; Lack et al., 2016; Langley et al., 2012; Pool et al., 2012) and Pool-Seq data from various sources (Bastide et al., 2013; Bergland et al., 2014; Kapun et al., 2016; Orozco-terWengel et al., 2012; Reinhardt et al., 2014) (Table S2). For single individuals, we obtained genotypes of the focal SNP from the *Drosophila* Genome Nexus website (DGN; <http://www.johnpool.net/genomes.html>) and estimated allele frequencies based on the number of chromosomes carrying the A-variant for populations with at least ten sequenced individuals. For Pool-Seq datasets, we re-mapped quality-filtered raw data as described in (Kapun et al., 2016) and estimated allele frequencies based on read counts of the A-variant relative to the total coverage. To increase sequence coverage in Queensland and Tasmania, we merged libraries of multiple collections at the corresponding locations (Reinhardt et al., 2014). We further used a collection of Pool-Seq data from 48 population samples collected across Europe by the DrosEU consortium (Kapun et al., 2018) (accession number: PRJNA388788) for an in-depth analysis of spatial distribution of the A-variant. Specifically, we tested for clinal distribution along the latitudinal and longitudinal axes by means of generalized linear models (GLMs) in R based on allele counts to account for the biallelic nature of the focal SNP. We further contrasted the clinality of 4: 710,326 to 21,008 putatively neutral genome-wide SNPs located in short introns (<60bp) and in distance to chromosomal inversions (Clemente and Vogl, 2012; Parsch et al., 2010). To test if the observed p-value from a GLM at the focal SNP deviates from neutral expectation we empirically assessed significance. We therefore generated empirical cumulative density functions (ECDF) based on the $-\log_{10}$ transformed p-values of all neutral SNPs and calculated the area under the ECDF confined by the $-\log_{10}$ p-value of the focal SNP and the upper tail of the distribution by integration. This area corresponds to the percentile of neutral SNPs with p-values equal or smaller than the focal SNP and thus summarizes the significance of clinality for 4:710,326 relative to genome-wide neutral estimates. We further characterized chromosome-wide patterns of genetic variation by estimating the population genetics statistics π and Tajima's D for all 48 samples from the DrosEU dataset using Pool-Gen (Kapun et al., 2018) with implemented corrections for Pool-Seq data (Futschik and Schlötterer, 2010; Kofler et al., 2011). At last, we tested whether very rare occurrences of the A-variant in Sub-Saharan Africa may be due to admixture with non-African genetic variation. We therefore used admixture probability estimates from (Lack et al., 2015) (see Table S2) to classify African lines as admixed (>10% of the autosomes admixed) or non-admixed ($\leq 10\%$ of the autosomes admixed) and compared genotype counts for admixed and non-admixed lines by means of Fisher exact tests (FET) in R.

Electrophoretic Mobility Shift Assays

Nuclear extracts of stably transfected S2 cells expressing the Cut-FLAG fusion protein were prepared from cells pelleted and resuspended for 15 minutes at 4°C in the membrane lysis buffer (Tris-HCl pH7,4 20mM, NaCl 10mM, MgCl₂ 3mM, DTT 0,5mM, NP40 1%, protease inhibitors). Cell extracts were centrifuged 5 minutes at 13000 RPM. Pelleted nuclei were then rinsed twice with PBS 1X and lysed for 30 minutes at 4°C in the nuclear envelope lysis buffer (Tris-HCl pH7,4 20mM, NaCl 100mM, MgCl₂ 3mM, EDTA 1mM, glycerol 10%, DTT 0,5mM, SDS 0,1%, sodium deoxycholate 0,5%, Triton X-100 1%, protease inhibitors). After centrifugation for 5 minutes at 13000 RPM, the supernatant containing the nuclear extract was recovered and stored at -80°C.

Cold Electrophoretic Mobility Shift Assays (EMSA) were performed with 8 or 16µg of nuclear extract using the LightShift Chemiluminescent EMSA kit (ThermoFisher Scientific) according to the manufacturer's instructions, with a modified binding buffer (Tris-HCl pH7,5 10mM, KCl 50mM, DTT 1mM, MgCl₂ 5mM, EDTA 1mM, Ficoll 400 5%, BSA 0,1µg/µl, NP40 0,05%, poly dI-dC 50ng/µl or Tris-HCl pH7,5 10mM, KCl 100mM, DTT 1mM, MgCl₂ 5mM, EDTA 1mM, NP40 0,05%, poly dI-dC 50ng/µl)

5'-biotinylated and unlabeled probes corresponding to the G and A-alleles of the *ey* enhancer were generated. To this goal, labeled or non-labeled complementary oligonucleotides were synthesized and hybridized. Oligonucleotide sequences are:

G-allele: 5'-ACAATAAAATGGTTGGAACCTTTTCGAACCTTT-3'

A-allele: 5'-ACAATAAAATGGTTGAACTTTTCGAACCTTT-3'

The binding was performed for 20 minutes at room temperature, followed by electrophoretic migration onto native 5% polyacrylamide gel in TBE 0,5× buffer. The supershift experiment was performed adding 2µg of the M2 anti-FLAG antibody (Sigma-Aldrich Cat# F3165).

QUANTIFICATION AND STATISTICAL ANALYSIS

In all experiments, sample size was determined *a priori*. Data were excluded exclusively prior to quantification based on poor image quality or inadequate developmental stage, explaining differences in sample size between groups. Except where stated otherwise, all statistical tests and charts were performed using GraphPad Prism 7 (GraphPad Software Inc.). Normality of the data were systematically assessed and statistical tests selected accordingly. Details on statistical tests, sample sizes and p values are indicated in figure legends except where mentioned otherwise.

DATA AND CODE AVAILABILITY

The code for the macro used for quantification of GFP/RFP colocalization (Oliva et al., 2016) is available on GitHub at the following link: <https://github.com/rejsmont/FijiScripts/blob/master/mColoc3D.py>. Raw images are available upon request from the Lead Contact.