

mirror determines the far posterior domain in butterfly wings

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Reviewed Preprint

Published from the original preprint after peer review and assessment by eLife.

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Reviewed preprint version 1

May 8, 2024 (this version)

Posted to preprint server

February 20, 2024

Sent for peer review

February 15, 2024

 https://en.wikipedia.org/wiki/Open_access

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Abstract

Insect wings, a key innovation that contributed to the explosive diversification of insects, are recognized for their remarkable variation and many splendid adaptations. Classical morphological work subdivides insect wings into several distinct domains along the antero-posterior (AP) axis, each of which can evolve relatively independently. There has been little molecular evidence, however, for AP subdivision beyond a single compartment boundary described from *Drosophila melanogaster*. Here we show that the transcription factor *mirror* acts as a selector gene to differentiate a far posterior domain in the butterfly wing, classically defined as the vannus, and has wide-ranging effects on wing shape, scale morphology, and color pattern. Our results confirm that insect wings can have more than one posterior developmental domain, and support models of how selector genes may facilitate evolutionarily individuation of distinct AP domains in insect wings. Our results also suggest that the alula, a small *mirror*-dependent structure at the base of the *D. melanogaster* wing, may be an evolutionary derivative of the vannus, and therefore that the *D. melanogaster* wing blade is a solitary remigium that represents only a fraction of the archetypal insect wing.

eLife assessment

This article is a **valuable** addition to the growing literature on the developmental patterning of insect wings. Using CRISPR mutagenesis and localization of mRNA, the authors present **solid** evidence that the transcription factor Mirror is necessary for specifying the morphological identity of the most posterior regions of butterfly wings. The manuscript would benefit from more careful use of terminology and appropriate citation of related *Drosophila* literature, and there are also some concerns about whether the phenotype represents transformation or loss which might be clarified through a closer look at ultrastructure. With a clearer presentation of terminology, this paper would be of general interest to developmental and evolutionary biologists.

Results and discussion

Insect wings vary dramatically in their size and morphology. Butterflies and moths in particular have attracted attention for their rapidly evolving wings that show extreme variation in shape and color pattern. Interestingly, lepidopteran color pattern diversity appears to follow some simple rules where basic spot and stripe color pattern elements change size, color, and presence/absence relatively independently based on which domain along the antero-posterior (AP) axis they occur in.³ Analyses of wing pattern diversity across butterflies, considering both natural variation and genetic mutants, suggest that wings can be subdivided into at least five AP domains, bounded by the M1, M3, Cu2, and 2A veins respectively, within each of which there are strong correlations in color pattern variation and wing morphology (**Figure 1A**).⁴⁻⁶ The anterior-most of these domains, bordered by the M1 vein, appears to correspond to an AP compartment boundary originally described by cell lineage tracing in *Drosophila melanogaster*,⁷ and later supported in butterfly wings by expression of the Engrailed transcription factor.⁸ Interestingly, however, *D. melanogaster* work has yet to reveal clear evidence for additional AP domain boundaries in the wing. It has thus been a matter of significant interest and controversy whether additional AP wing domains might occur in other insects,⁹⁻¹¹ as their existence would provide an explanatory model for the modular diversification of insect wing morphology.

The goal of this study was to identify the molecular basis of AP domain specification in the butterfly wing, posterior of the Engrailed boundary. We initially identified the transcription factor *mirror* as a potential candidate gene based on previous mRNA-seq studies,¹² coupled with *D. melanogaster* work showing that *mirror* specifies the alula – a small membranous lobe at the posterior base of the wing in some dipterans (**Figure 1C,D**).^{2,13} We used *in situ* hybridization to visualize *mirror* mRNA localization in the last-instar imaginal discs of the common buckeye butterfly, *Junonia coenia*, and observed *mirror* expression throughout the wing domain posterior to the 2A vein boundary (**Figures 2A** and **S1**). This expression precisely marks the vannus, or anal region (**Figure 1E**), of the butterfly wing – a distinct posterior field of the wing blade populated by the anal (A) veins, and bordered anteriorly by the claval fold between the cubital (Cu) and A veins in the hindwing. This region was also previously proposed to represent the posterior-most color pattern domain in butterflies (**Figure 2B**).³

To functionally assess the role of *mirror* in wing development, we produced CRISPR/Cas9 deletion mosaic knockouts (mKOs) targeting the coding region of *J. coenia mirror*. The resulting mutants revealed vannus-specific phenotypes, consistent with the *mirror* expression patterns. The unique identity of the vannus was lost in mutants, so that the vannus resembled the remigium (**Figure 2C-E**; **Figure S2**). In brief, the vannus took on a morphology consistent with anterior wing domains, including loss or reduction of the vannal fold (along with an overall shape along the posterior wing margin), reduction of light silvery iridescent coloration, and continuation of normally truncated wing margin color patterns along the posterior distal margin of the wing (**Figure 2E**; **Figure S2**). Some mutants also displayed aberrations in vannal venation, including loss of the 3A vein, discontinuous and bifurcated 2A veins, and ectopic veins (**Figures 2C-E**; **Figures S2** and **S3**). Together, our expression and knockout data show that *mirror* acts as a selector gene necessary for defining the identity of the far posterior wing domain that corresponds to the vannus.

Our finding is of interest for several reasons. First, it provides a molecular explanation for how different AP domains of the butterfly wing may be individuated to independently evolve their own shape and color pattern variations. Previous authors have proposed the existence of such individuated domains, and speculated that they may be specified by selector genes.^{5,10} Our data provide experimental support for this model, and now motivate us to identify factors that specify other domain boundaries between the M1 and A2 veins. Second, the role of *mirror* in

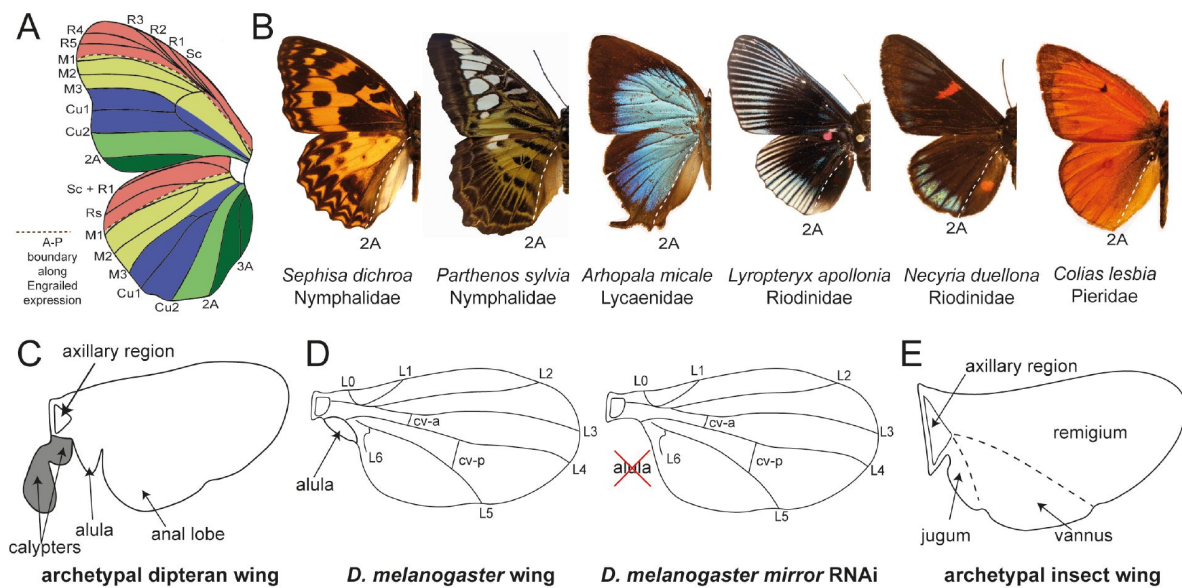


Figure 1.

Proposed AP domains of insect wings.

(A) Comparative morphology has identified boundaries associated with color pattern variation demarcated by the M1, M3, Cu2, and 2A veins, as summarized in McKenna et al.⁵ (B) Examples of the 2A vein marking a clear posterior color pattern boundary across multiple butterfly families. (C) Proximal posterior features of the archetypal dipteran wing include the calypters, which are only found in Calypterata clade, and the alula, a lobe at the base of the wing blade.¹ (D) *mirror* RNAi knockdowns result in highly specific loss of the alula in *D. melanogaster*.² (E) Snodgrass' model of the archetypal insect wing specifies three major domains along the anteroposterior axis: the remigium, the vannus, and the jugum. Homologies of these domains with the features of the dipteran wing (C), or butterfly wing pattern boundaries (A), has remained unclear.

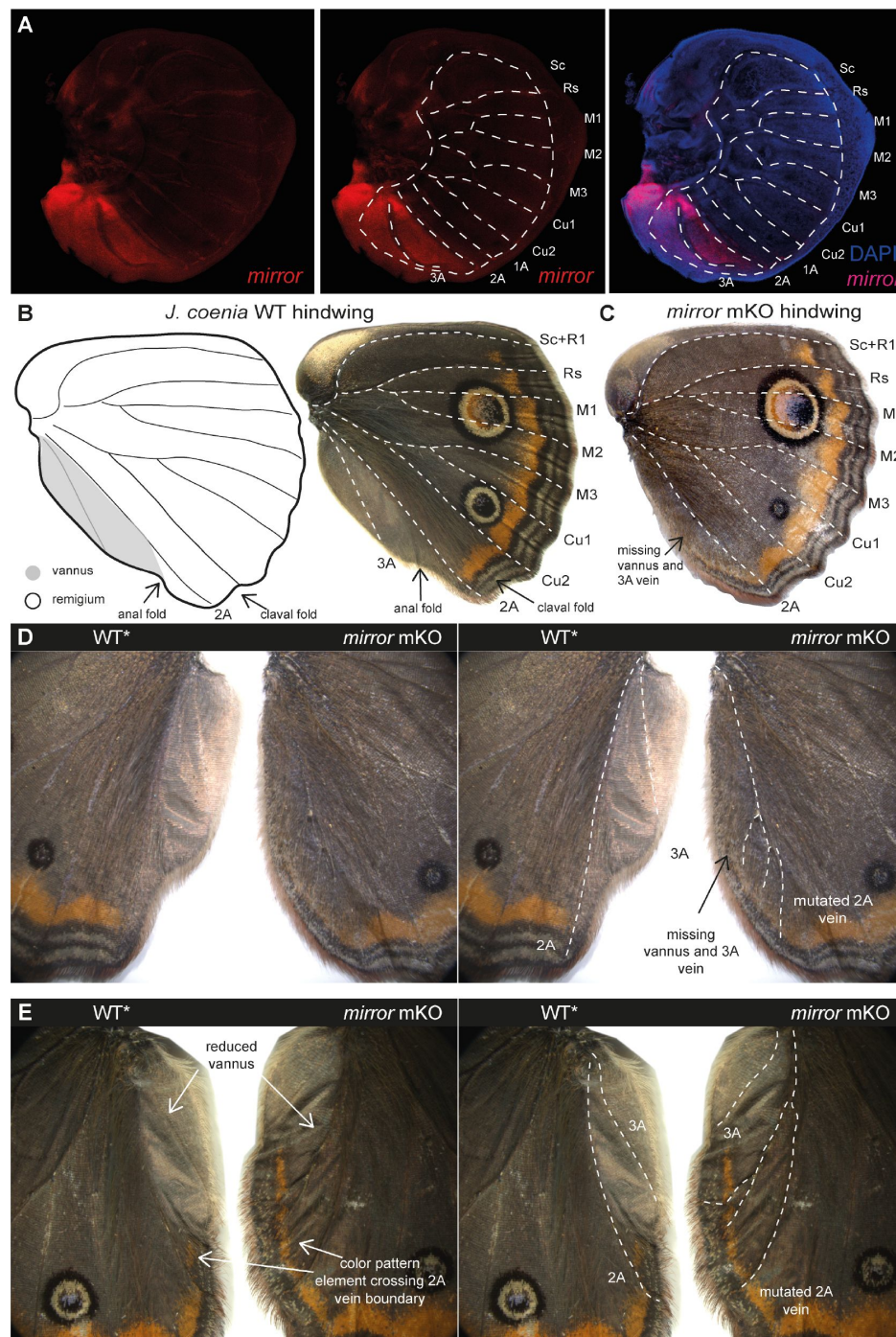


Figure 2.

***mirror* determines the identity of the far posterior vannus domain in *J. coenia*.**

(A) In *J. coenia* *mirror* (red) is expressed posterior to the 2A vein in the late last-instar hindwing imaginal discs, marking the region that will develop into the vannus. See Figure S1 for forewing expression and controls. (B) In adult butterfly wings the vannus is the wing field posterior to the 2A vein, as shown here in *J. coenia*. (C) Targeted mosaic knockouts (mKOs) of *mirror* results in loss of vannal identity in *J. coenia*, including (D) reduction of the anal fold, and (E) continuation of margin color patterns along the posterior wing margin. Some *mirror* mKOs (C-E) also show mutation in the 2A vein along with loss (D) or reduction (E) of the 3A vein, likely associated with mosaic clone boundaries. mKOs result in asymmetric effects observed in the wings where (D,E) one wing resembles a wild-type wing (WT*) more than the other wing from the respective individual. For more *mirror* mKOs see [Figures S2](#) and [S3](#).

specifying the vannus provides an important link between molecular developmental biology and Snodgrass' classical anatomical designations of the insect wing fields – i.e., the remigium, the vannus, and the jugum (**Figure 1E**).¹⁴ Importantly, *mirror* knockdown in the milkweed bug *Oncopeltus fasciatus* causes developmental defects in the claval and anal furrows in the posterior wing,¹⁵ which leads us to infer that *mirror*'s role in determining the vannus may be deeply conserved in insects.

Finally, the function of *mirror* in determining the alula in *D. melanogaster*² suggests that the alula may represent an evolutionarily reduced vannus. The ramifications of this are significant for reconstructing the history of the insect wing, as they suggest that the *D. melanogaster* wing blade is a lone remigium – only one-third of the archetypal insect wing (**Figure 1E**). The dipteran jugum appears to have been lost entirely, perhaps except as a calypter in the Calypterata lineage, as speculated by Snodgrass.¹⁴ Thus, while flies have been an important model for characterizing genes and processes that build insect wings, a more comprehensive understanding of the development and evolution of insect wings will require work in species that have wings more representative of the complete ancestral blueprint.

Acknowledgements

We thank Kate Siegel, Nigel Williams, and Rick Fandino for assistance with rearing butterflies; Johanna Dela Cruz for help with confocal microscopy at Biotechnology Resource Center (BRC) Imaging Facility (RRID:SCR_021741) at the Cornell Institute of Biotechnology. We also thank Dr. Eirene Markenscoff-Papadimitriou for use the Leica Stellaris 5 confocal microscope for imaging. This work was supported by the United States National Science Foundation grants NSF IOS-1753559 and IOS-2128164 awarded to R.D.R., and a Cornell Summer Experience Grant and an Office of Undergraduate Biology fellowship to X.Y.

Materials and methods

Identity of *Iroquois* family gene *mirror* ortholog in *J. coenia*

We performed a reciprocal BLAST of amino acid sequences of *D. melanogaster* *Iroquois* complex genes *araucan*, *caupolican*, and *mirror* to identify their orthologs in the latest *J. coenia* and *Heliconius erato lativitta* genome sequences on lepbases.org and *Tribolium castaneum* and *Apis mellifera* on NCBI. The amino acid sequences of the top hits – from *J. coenia* (JC_02265-RA, JC_02269-RA), *H. erato lativitta* (HEL_009655-RA, HEL_009656-RA), *A. mellifera* (LOC412840, LOC412839) and *T. castaneum* (LOC652944, LOC660345) were aligned to the *Iroquois* genes of *D. melanogaster* using MUSCLE.¹⁶ We used these alignments to build a maximum likelihood gene phylogeny on IQ-TREE 2¹⁷ using ModelFinder¹⁸ for estimating the most accurate substitution model for our amino acid sequences. We used FigTree to visualize the tree depicted in **Figure S4**.

HCR fluorescent *in situ* hybridization of *mirror* mRNA

This protocol was adapted and modified from Bruce et al. ([dx.doi.org/10.17504/protocols.io.bunznvf6](https://doi.org/10.17504/protocols.io.bunznvf6)). The coding sequence of JC_02269-RA was used by Molecular Instruments, Inc. (Los Angeles, CA, USA) to design a *mirror*-specific hybridization chain reaction (HCR) probe set (Lot #PRQ901). The probes are unique to the JC_02269-RA transcript as verified by BLAST search using the latest *J. coenia* genome assembly (v2) on lepbases.org, and do not share significant sequence similarity with any other transcripts, including the other *Iroquois* family member JC_02265-RA (see **Figure S4**).

Last (5th) instar larval wing imaginal discs were dissected in cold 1x PBS. The dissected discs collected from left and right sides of the larvae were randomly assigned to control or treatment batches. Both controls and treatment batches were fixed in 1x cold fix buffer (750μL PBS, 50mM EGTA, 250μL 37% formaldehyde) on ice. The discs were then washed three times in 1x PBS with 0.1% Tween 20 (PTw) on ice spending approximately 30sec – 2min for each wash. Tissues were then gradually dehydrated by washing in 33%, 66%, and 100% MeOH (in PTw) for 2-5min / wash on ice, and stored in 100% MeOH in -20°C for days until HCR protocol was started. On the day of the HCR protocol, the wing discs were progressively rehydrated in cold 75% MeOH, 50% MeOH, and 25% MeOH in PTw spending 2-5min per wash. After rehydrating, discs were washed once for 10min and twice for 5min with PTw. The discs were then permeabilized in 300-500μL of detergent solution (1.0% SDS, 0.5% Tween, 50mM Tris-HCL pH 7.5, 1mM EDTA pH 8.0, 150mM NaCl) for 30min at room temperature. While waiting, hybridization buffer (Molecular Instruments) was warmed to 37°C (200μL/tube). After permeabilizing with detergent solution, each tube of wing discs was incubated in 200μL of pre-warmed hybridization buffer for 30min at 37°C. Probe solution was prepared by adding 0.8pmol (0.8 μl of probe from 1uM stock solution) of each probe mixture to 200μL of probe hybridization buffer at 37°C. The pre-hybridization buffer was removed, and the probe solution was added. For negative control tissues, hybridization buffer without probes was added instead. All wing discs were incubated overnight (12-16hr) at 37°C. Before resuming the protocol, probe wash buffer (Molecular Instruments) was warmed to 37°C, amplification buffer (Molecular Instruments) was calibrated to room temperature, and a heat block was set to 95°C. The probe solution was removed and saved at -20°C to be reused. Wing discs were then washed four times in 1mL pre-warmed probe wash buffer at 37°C for 15min per wash. After the last wash step, discs were washed twice (5min/wash) with 1mL of 5x SSCT (5X sodium chloride sodium citrate, 0.1% Tween 20) at room temperature. The wing discs were pre-amplified with 1mL of equilibrated amplification buffer for 30min at room temperature. During this step, 2μL of each of hairpin h1 and 2μL of each hairpin h2 were mixed in 100μL of amplification buffer at 95°C for 90sec and cooled to room temperature in the dark for 30 min. For *mirror-only in situ* hybridizations, we used B1 hairpin amplifiers tagged with AlexaFluor 594 fluorophore (Molecular Instruments). For *mirror* and *wingless* double stains, we used *mirror* probes with B1 amplifier tagged with AlexaFluor 647 fluorophore and *wingless* probes (Lot #PRG129) with B3 hairpin amplifiers tagged with AlexaFluor 546 fluorophore. After 30min, the pre-amplification buffer was removed from the discs, hairpin solution added, and the tissues were then incubated in the dark at room temperature for 2-16hr. The hairpins were removed and saved at -20°C to be reused later. Excess hairpins were removed by washing 5 times (twice for 5min, twice for 30min and once for 5min) with 1mL of 5x SSCT at room temperature. The wing discs were then incubated in 50% glycerol solution (in 1X PBS) with DAPI (0.01μg/mL) overnight at 4°C. The wing discs were then mounted and visualized on a Zeiss 710 or Leica Stellaris 5 confocal microscope.

CRISPR-Cas9 mediated mutagenesis of *mirror* in *J. coenia*

Two single-guide RNAs (sgRNAs; sgRNA1-5'-GAATGGACTTGAACGGGCA; sgRNA2-5'-AGAAACAGGGTCGATGATGA) targeting the homeobox domain of *mirror* were mixed with 500ng/μl of Cas9 nuclease and injected to *J. coenia* eggs 0.5-4hr after oviposition (n = 1042 eggs) as previously described.¹⁹ Injected G0 individuals were reared on standard artificial diet, until they emerged and were immediately frozen in -20°C upon emergence. Wings of surviving G0 adults (n=99) were assayed for anomalies to detect mosaic knockout (mKO) phenotypes (n=29). Injection results and mutation phenotype frequencies are detailed in Table S1.

Validating *mirror* mutants by genotyping

DNA was extracted using E.Z.N.A. Tissue DNA Kit (Omega Bio-Tek) from the thorax of individuals that showed mutations in the wings. Extracted genomic DNA was amplified using a pair of primers flanking the sgRNA cut sites (Forward: 5'-CGCTGTGCCACCTTAAAC, Reverse: 5'-

GTATGGCTCGGGGGATTCTG). Amplified DNA was run on a 2% agarose gel and were excised and purified using the MicroElute Cycle-Pure Kit (Omega Bio-Tek). Purified DNA was Sanger sequenced by Cornell Institute of Biotechnology. Example mutant alleles of *mirror* are shown in **Figure S5** [↗](#).

Author contributions

Conceptualization, M.C and R.D.R; Methodology, M.C and R.D.R; Investigation, X.Y.Y, M.C, N.K.B and G.C.H; Writing-Original Draft, M.C, X.Y.Y and R.D.R; Writing-Review & Editing, M.C and R.D.R; Visualization, M.C and X.Y.Y; Funding Acquisition, R.D.R and X.Y.Y; Supervision, M.C and R.D.R

Supplemental information

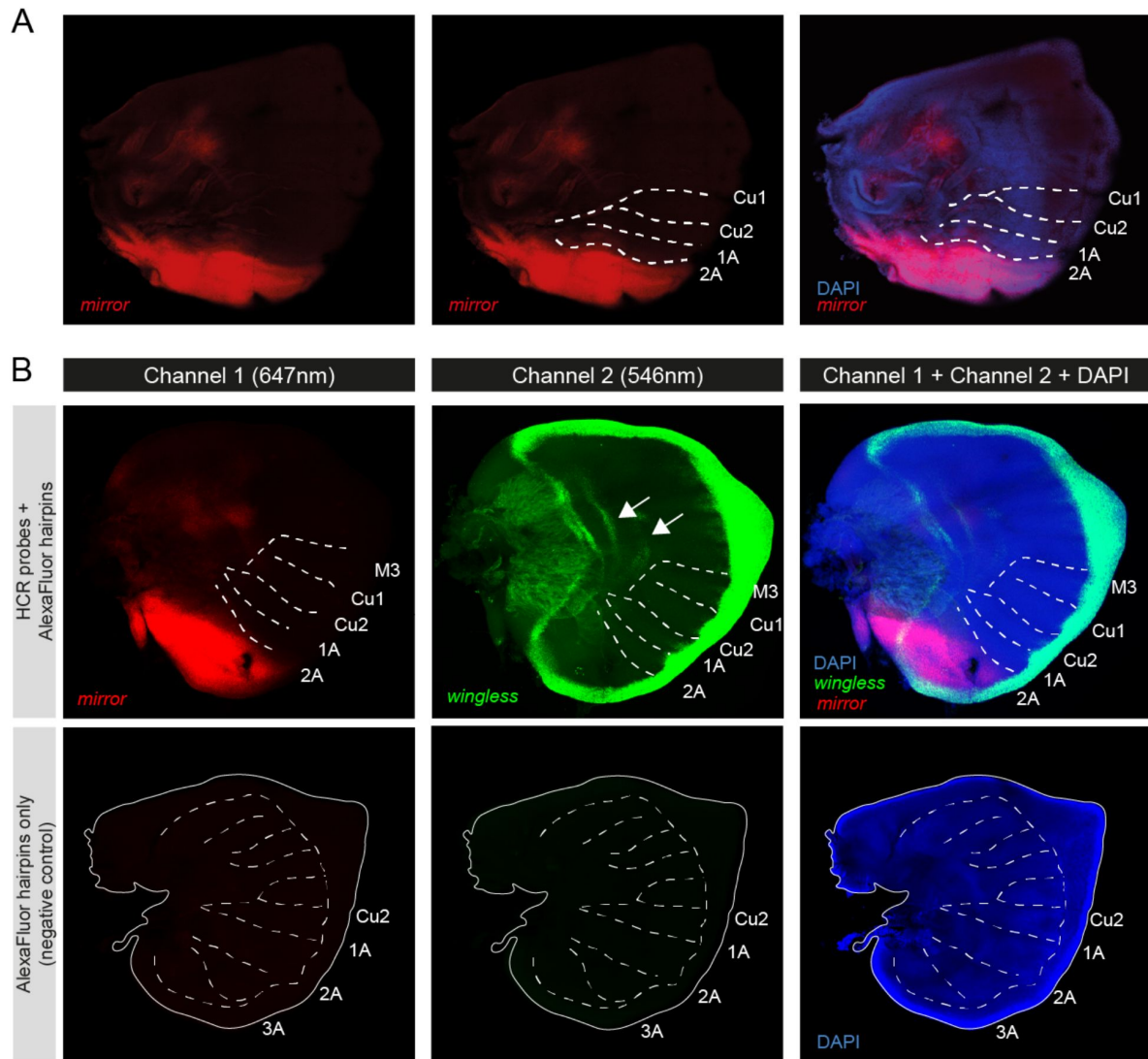


Figure S1.

HCR *in situ* hybridization of *mirror* highlight posterior domain expression in *J. coenia* wing imaginal discs. Related to Figure 2A.

(A) *mirror* is expressed posterior to the 2A vein in the last-instar forewing imaginal disc of *J. coenia*. (B) Double *in situ* of *mirror* (Channel 1: 647nm) and *wingless* (Channel 2: 546 nm) in last instar *J. coenia* hindwing imaginal discs. Expression of *wingless* observed by HCR precisely matches expression in wing margin peripheral tissue and discal bands (white arrow) as previously described from chromogenic *in situ*.⁵² Bottom panel: Negative controls, lacking HCR probes, showed no signal in the 647nm and 546nm channels. Dashed lines show position of posterior veins.

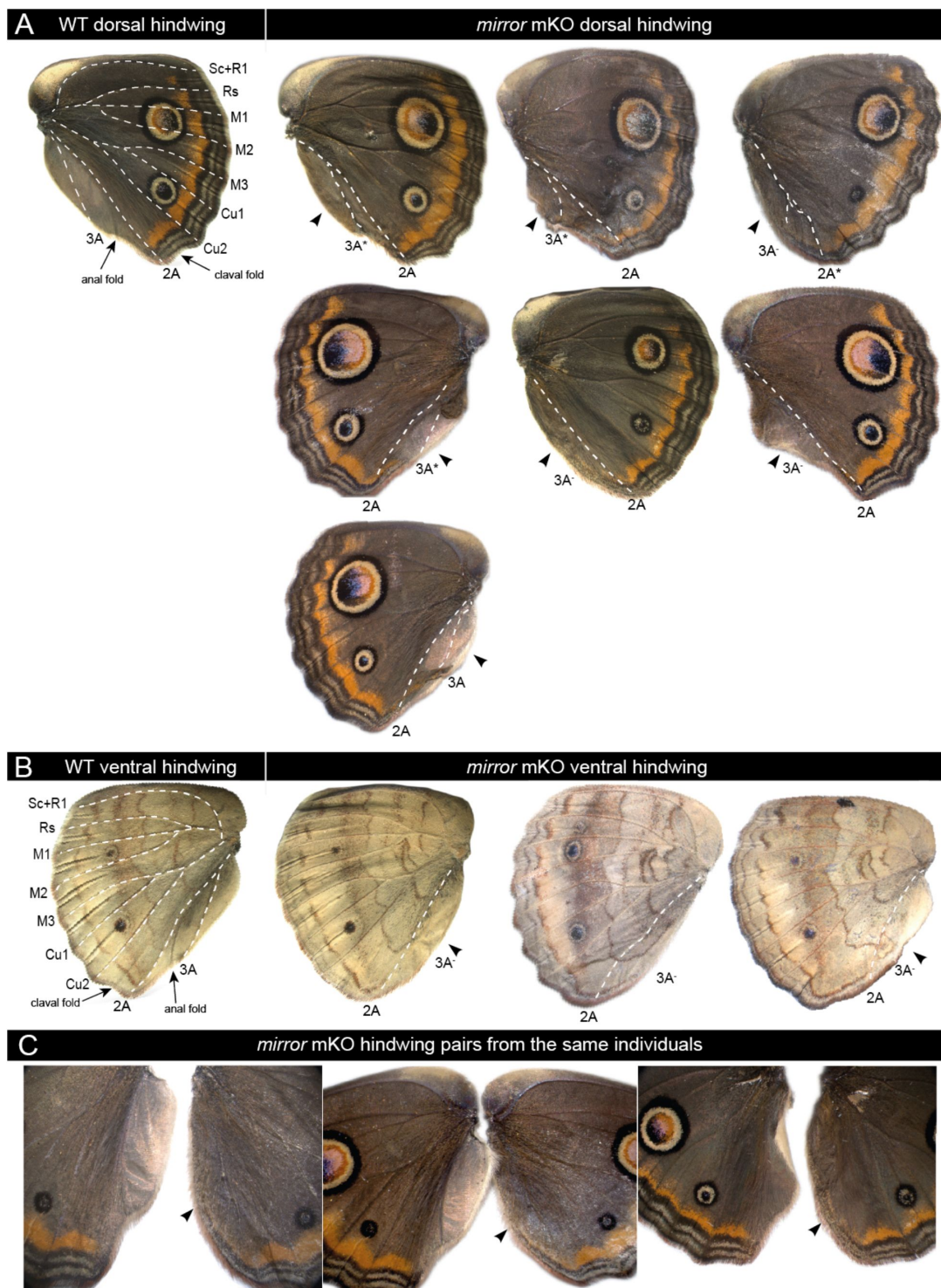


Figure S2.

mKOs of *mirror* result in partial or entire loss of vannus identity in *J. coenia* hindwings. Related to Figure 2B-E

Arrows highlight vernal reduction-related phenotypes in the (A) dorsal side and (B) ventral side of the hindwings. Dashed lines show positions of veins. Next to vein labels, asterisks denote vein anomalies, and "-" denote missing veins. (C) Comparison of left and right dorsal hindwings from the same individual, with vernal reduction phenotype highlighted with arrows.

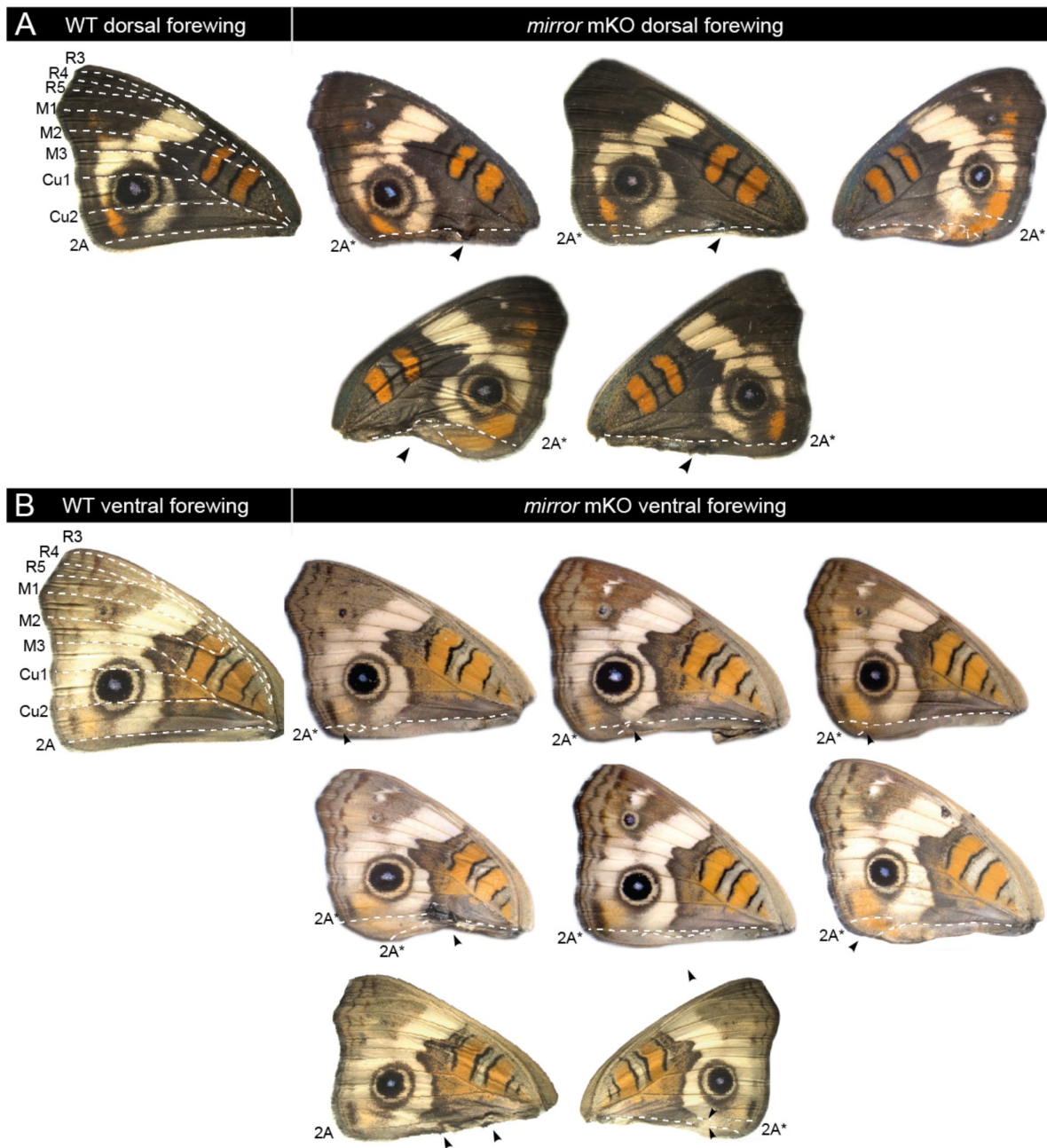


Figure S3.

***mirror* mKOs show post-2A morphological anomalies *J. coenia* forewings. Related to Figure 2B-E [↗](#).**

mKOs of *mirror* result in anomalies along the forewing 2A vein, including vein bifurcation and ectopic vein fragments, which are often associated with color pattern disruption. Dashed lines show the position of 2A vein; asterisks denote 2A vein anomalies.

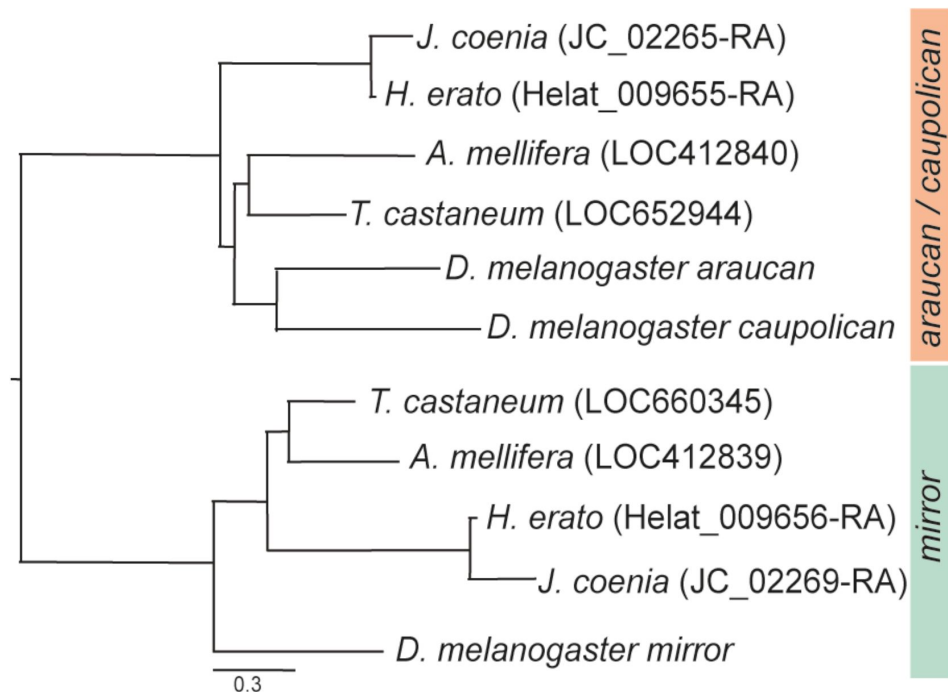


Figure S4.

Identification of *mirror* ortholog in *J. coenia*.

A maximum likelihood phylogeny of *J. coenia*, *H. erato lativitta*, *Tribolium castaneum*, *Apis mellifera*, and *D. melanogaster Iroquois* complex genes confirms that JC_02269-RA is the ortholog of *mirror*.

Individuals	Indels	sg RNA1	PAM
mKO 1	WT	GAATGGACTTGAACGGGGCAAGGAGAAAAAATGCAACCAGAGAGACCACA	
	-7	GAATGGACTTGAACGG-----AGAAAAAATGCAACCAGAGAGACCACA	
	-2	GAATGGACTTGAACGGG---AAGGAGAAAAAATGCAACCAGAGAGACCACA	
	+1	GAATGGACTTGAACGG--NGCAAGGAGAAAAAATGCAACCAGAGAGACCAC	
	-2	GAATGGACTTGAACGG---CAAGGAGAAAAAATGCAACCAGAGAGACCACA	
mKO 2	WT	GAATGGACTTGAACGGGGCAAGGAGAAAAAATGCAACCAGAGAGACCACA	
	-7	GAATGGACTTGAACGG-----AGAAAAAATGCAACCAGAGAGACCACA	
	-2	GAATGGACTTGAACGG---CAAGGAGAAAAAATGCAACCAGAGAGACCACA	
	-2	GAATGGACTTGAACGGG---AAGGAGAAAAAATGCAACCAGAGAGACCACA	
	+4	GAATGGACTTGAACGGGNNNNCAAGGAGAAAAAATGCAACCAGAGAGACC	
mKO 3	WT	GAATGGACTTGAACGGGGCAAGGAGAAAAAATGCAACCAGAGAGACCACA	
	-6	GAATGGACTTGAACGGG-----AGAAAAAATGCAACCAGAGAGACCACA	
	-7	GAATGGACTTGAACGG-----AGAAAAAATGCAACCAGAGAGACCACA	
	-3	GAATGGACTTGAACGG---AAGGAGAAAAAATGCAACCAGAGAGACCACA	
	-1	GAATGGACTTGAACGG--GCAAGGAGAAAAAATGCAACCAGAGAGACCACA	
	-6	GAATGGACTTGAACGG-----GAGAAAAAATGCAACCAGAGAGACCACA	
	-2	GAATGGACTTGAACGGG---AAGGAGAAAAAATGCAACCAGAGAGACCACA	

Figure S5.

Sanger sequencing confirms mutations at CRISPR sgRNA site in *mirror* mKOs.

Sanger sequencing results from three different *mirror* mutant individuals. Each line denotes a different wildtype (WT) or mutant allele with indels indicated in red, and the PAM site in green.

sgRNA concentration (ng/μL)	Eggs Injected	Hatched	Adults Emerged	Mutants
487 ng/μL	142	25	4	2
391 ng/μL	96	33	6	3
335 ng/μL	163	32	15	4
281 ng/μL	98	11	1	0
175 ng/μL	149	52	19	8
125 ng/μL	252	79	24	9
88.5 ng/μL	142	41	30	3
Total	1042	273	99	29

Table S1.

mirror CRISPR/Cas9 injection results.

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Reviewer #1 (Public Review):

Summary:

This short report shows that the transcription factor gene *mirror* is specifically expressed in the posterior region of the butterfly wing imaginal disk, and uses CRISPR mosaic knock-outs to show it is necessary to specify the morphological features (scales, veins, and surface) of this area.

Strengths:

The data and figures support the conclusions. The article is swiftly written and makes an interesting evolutionary comparison to the function of this gene in *Drosophila*. Based on the data presented, it can now be established that *mirror* likely has a similar selector function for posterior-wing identity in a plethora of insects.

Weaknesses:

This first version has minor terminological issues regarding the use of the terms "domains" and "compartment".

<https://doi.org/10.7554/eLife.96904.1.sa2>

Reviewer #2 (Public Review):

This is a short and unpretentious paper. It is an interesting area and therefore, although much of this area of research was pioneered in flies, extending basic findings to butterflies would be worthwhile. Indeed, there is an intriguing observation but it is technically flawed and these flaws are serious.

The authors show that *mirror* is expressed at the back of the wing in butterflies (as in flies). They present some evidence that is required for the proper development of the back of the wing in butterflies (a region dubbed the *vannus* by the ancient guru Snodgrass). But there are problems with that evidence. First, concerning the method, using CRISPR they treat embryos and the expectation is that the *mirror* gene will be damaged in groups of cell lineages, giving a mosaic animal in which some lines of cells are normal for *mirror* and others are not. We do not know where the clones or patches of cells that are defective for *mirror* are because they are not marked. Also, we do not know what part of the wing is wild type and what part is mutant for *mirror*. When the *mirror* mutant cells colonise the back of the wing and that butterfly survives (many butterflies fail to develop), the back of the wing is altered in some selected butterflies. This raises a second problem: we do not know whether the rear of the wing is missing or transformed. From the images, the appearance of the back of the wing is clearly different from the wild type, but is that due to transformation or not? And then I believe we need to know specifically what the difference is between the rear of the wing and the main part. What we see is a silvery look at the back that is not present in the main part, is it the structure of the scales? We are not told. There are other problems. *Mirror* is only part of a group of genes in flies and in flies both *iroquois* and *mirror* are needed to make the back of the wing, the *alula* (Kehl et al). What is known about *iro* expression in butterflies?

In flies, *mirror* regulates a late and local expression of *dpp* that seems to be responsible for making the *alula*. What happens in butterflies? Would a study of the expression of *Dpp* in wildtype and *mirror* compromised wings be useful?

Thus, I find the paper to be disappointing for a general journal as it does little more than claim what was discovered in *Drosophila* is at least partly true in butterflies. Also, it fails to explain what the authors mean by "wing domains" and "domain specification". They are not

alone, butterfly workers, in general, appear vague about these concepts, their vagueness allowing too much loose thinking.

Since these matters are at the heart of the purpose and meaning of the work reported here, we readers need a paper containing more critical thought and information. I would like to have a better and more logical introduction and discussion.

The authors do define what they mean by the vanner of the wing. In flies the definition of compartments is clear and abundantly demonstrated, with gene expression and requirement being limited precisely to sets of cells that display lineage boundaries. It is true that domains of gene expression in flies, for example of the *iroquois* complex, which includes *mirror*, can only be related to patterns with difficulty. Some recap of what is known plus the opinion of the authors on how they interpret papers on possible lineage domains in butterflies might also be useful as the reader, is no wiser about what the authors might mean at the end of it!

The references are sometimes inappropriate. The discovery of the AP compartments should not be referred to Guillen et al 1995, but to Morata and Lawrence 1975. Proofreading is required.

<https://doi.org/10.7554/eLife.96904.1.sa1>

Reviewer #3 (Public Review):

Summary:

The manuscript by Chatterjee et al. examines the role of the *mirror* locus in patterning butterfly wings. The authors examine the pattern of *mirror* expression in the common buckeye butterfly, *Junonia coenia*, and then employ CRISPR mutagenesis to generate mosaic butterflies carrying clones of *mirror* mutant cells. They find that *mirror* is expressed in a well-defined posterior sector of final-instar wing discs from both hindwings and forewings and that CRISPR-injected larvae display a loss of adult wing structures presumably derived from the *mirror* expressing region of hindwing primordium (the case for forewings is a bit less clear since the *mirror* domain is narrower than in the hindwing, but there also do seem to be some anomalies in posterior regions of forewings in adults derived from CRISPR injected larvae). The authors conclude that the wings of these butterflies have at least three different fundamental wing compartments, the *mirror* domain, a posterior domain defined by engrailed expression, and an anterior domain expressing neither *mirror* nor engrailed. They speculate that this most posterior compartment has been reduced to a rudiment in *Drosophila* and thus has not been adequately recognized as such a primary regional specialization.

Critique:

This is a very straightforward study and the experimental results presented support the key claims that *mirror* is expressed in a restricted posterior section of the wing primordium and that mosaic wings from CRISPR-injected larvae display loss of adult wing structures presumably derived from cells expressing *mirror* (or at least nearby). The major issue I have with this paper is the strong interpretation of these findings that lead the authors to conclude that *mirror* is acting as a high-level gene akin to engrailed in defining a separate extreme posterior wing compartment. To place this claim in context, it is important in my view to consider what is known about engrailed, for which there is ample evidence to support the claim that this gene does play a very ancestral and conserved function in defining posterior compartments of all body segments (including the wing) across arthropods.

(1) Engrailed is expressed in a broad posterior domain with a sharp anterior border in all segments of virtually all arthropods examined (broad use of a very good pan-species anti-En

antibody makes this case very strong).

(2) In *Drosophila*, marked clones of wing cells (generated during larval stages) strictly obey a straight anterior-posterior border indicating that cells in these two domains do not normally intermix, thus, supporting the claim that a clear A/P lineage compartment exists.

In my opinion, mirror does not seem to be in the same category of regulator as engrailed for the following reasons:

(1) There is no evidence that I am aware of, either from the current experiments, or others that the mirror expression domain corresponds to a clonal lineage compartment. It is also unclear from the data shown in this study whether engrailed is co-expressed with mirror in the posterior-most cells of *J. coenia* wing discs. If so, it does not seem justified to infer that mirror acts as an independent determinant of the region of the wing where it is expressed.

(2) Mirror is not only expressed in a posterior region of the wing in flies but also in the ventral region of the eye. In *Drosophila*, mirror mutants not only lack the alula (derived approximately from cells where mirror is expressed), but also lack tissue derived from the ventral region of the eye disc (although this ventral tissue loss phenotype may extend beyond the cells expressing mirror).

In summary, it seems most reasonable to me to think of mirror as a transcription factor that provides important development information for a diverse set of cells in which it can be expressed (posterior wing cells and ventral eye cells) but not that it acts as a high-level regulator as engrailed.

Recommendation:

While the data provided in this succinct study are solid and interesting, it is not clear to me that these findings support the major claim that mirror defines an extreme posterior compartment akin to that specified by engrailed. Minimally, the authors should address the points outlined above in their discussion section and greatly tone down their conclusion regarding mirror being a conserved selector-like gene dedicated to establishing posterior-most fates of the wing. They also should cite and discuss the original study in *Drosophila* describing the mirror expression pattern in the embryo and eye and the corresponding eye phenotype of mirror mutants: McNeill et al., *Genes & Dev.* 1997. 11: 1073-1082; doi:10.1101/gad.11.8.1073.

<https://doi.org/10.7554/eLife.96904.1.sa0>