- 1 Title: The regulation of a pigmentation gene in the formation of complex color patterns in
- 2 Drosophila abdomens
- 3
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23 Abstract

Changes in *cis*-regulatory modules (CRMs) that control developmental gene expression 24 patterns have been implicated in the evolution of animal morphology¹⁻⁶. However, the 25 26 genetic mechanisms underlying complex morphological traits remain largely unknown. Here we investigated the molecular mechanisms that induce the pigmentation gene *vellow* 27 28 (y) in a complex spot and shade pattern on the abdomen of the quinaria group species Drosophila guttifera. We show that the y expression pattern is controlled by only one CRM, 29 which contains a stripe-inducing CRM at its core. We identified several developmental 30 31 genes that may collectively interact with the CRM to orchestrate the patterning in the pupal abdomen of *D. guttifera*. We further show that the core CRM is conserved among *D*. 32 guttifera and the closely related quinaria group species Drosophila deflecta, which displays 33 34 a similarly spotted abdominal pigment pattern. Our data suggest that besides direct activation of patterns in distinct spots, abdominal spot patterns in *Drosophila* species may 35 have evolved through partial repression of an ancestral stripe pattern, leaving isolated 36 spots behind. Abdominal pigment patterns of extant quinaria group species support the 37 partial repression hypothesis and further emphasize the modularity of the *D. guttifera* 38 39 pattern.

40

How complex morphological features develop and evolve is a question of foremost importance
in biology. To address this question, we identified genes underlying abdominal pigmentation
pattern development in *Drosophila guttifera* (*D. guttifera*). The abdomen is decorated with six
rows of black spots that run along the anterior-posterior axis, divided by a dark dorsal midline
shade. This color pattern shows four sub-patterns: a dorsal, median, and lateral pair of spot rows,

plus the dorsal midline shade (Fig. 1a, b). *D. guttifera* belongs to the quinaria species group, whose members display highly diverse abdominal pigmentation patterns^{7,8}. While *D. guttifera* shows the most complex pattern of this group, most other quinaria group species lack at least one of the four sub-patterns, illustrating the pattern modularity among species. Interestingly, the stripe patterns of certain species often separate into spots^{7,8}. In this study, we show that the abdominal pigment patterns of quinaria group members may be formed by a combination of localized spot induction and partial stripe repression.

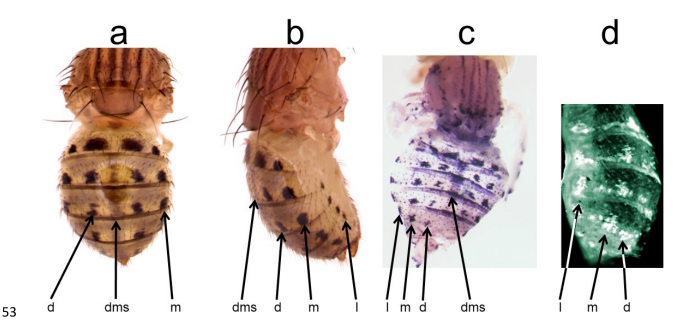
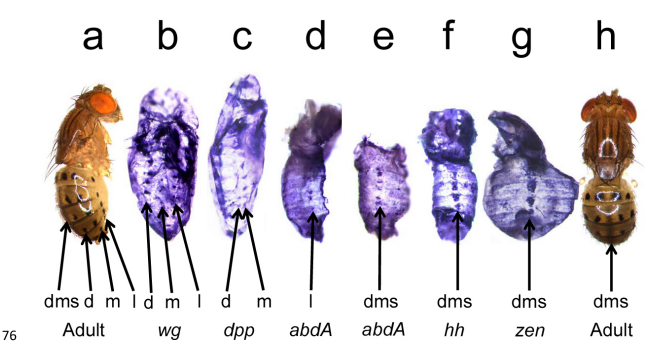


Figure 1: The *D. guttifera* abdominal color pattern is modular. a, Adult, dorsal view. b,
Adult, lateral view. c, *yellow* mRNA expression pattern in a pupal abdomen. d, Yellow protein
expression pattern in a pupal abdomen. dms = dorsal midline shade, d = dorsal, m = median, l =
lateral spot rows.

We focused on the regulation of the *yellow* (*y*) gene, which is required for the formation of black
melanin in insects⁸⁻¹⁴. Several *y* gene CRMs have been identified in various *Drosophila* species,

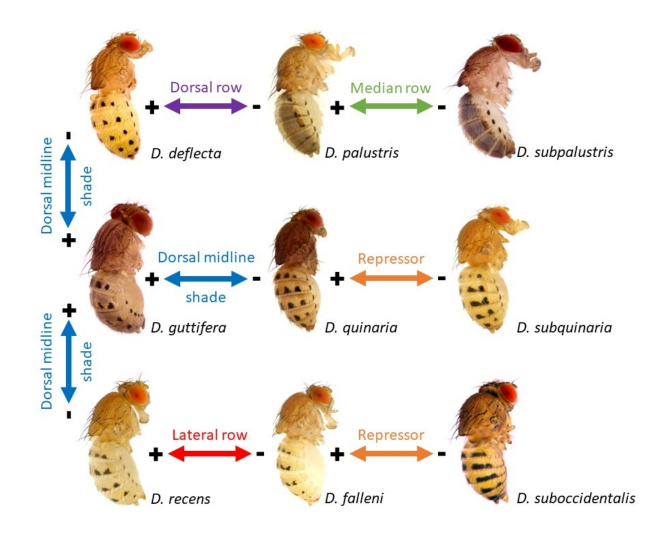
61	and changes in these CRMs and/or in the deployment of <i>trans</i> -factors that regulate y gene
62	expression have been implicated in the diversification of wing and body pigment patterns ^{12,15-19} .
63	In D. guttifera pupae, y gene expression and the location of the Y protein accurately prefigured
64	the complex adult abdominal pigment pattern (Fig. 1c, d). In order to identify putative upstream
65	activators of y , we performed an <i>in situ</i> hybridization screen for genes expressed in ways
66	prefiguring the y gene expression pattern. We found that wingless (wg) expression precisely
67	foreshadowed the six rows of black spots (Fig. 2b). Additionally, decapentaplegic (dpp)
68	expression foreshadowed the dorsal and median pairs of spot rows (Fig. 2c), while abdominal-A
69	(<i>abd-A</i>) expression correlated with the lateral pair of spot rows and the dorsal midline shade (Fig.
70	2d, e). <i>hedgehog (hh)</i> and <i>zerknullt (zen)</i> were additionally expressed along the dorsal midline of
71	the abdomen (Fig. 2f, g). Thus, the activation of the D. guttifera color pattern appears to be
72	induced in a modular fashion, which is in agreement with our observation that abdominal
73	pigmentation patterns within the quinaria group are variations of the D. guttifera pattern ground
74	plan (Fig. 3). This situation is reminiscent of the wing pattern ground plan in nymphalid
75	butterflies ^{20,21} .



77 Figure 2: The mRNA expression patterns of five developmental genes foreshadow the

78 *yellow* expression pattern. a, Adult, lateral view. b-g, *in situ* hybridizations in pupal abdomens.

h, Adult, dorsal view. dms = dorsal midline shade, d = dorsal, m = median, l = lateral spot rows.

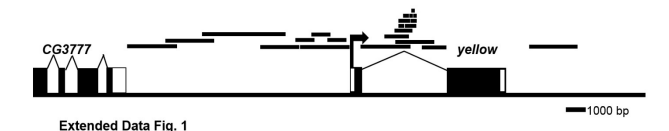


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Figure 3: Deviations from the *D. guttifera* ground plan create the diversity of quinaria
species' abdominal color patterns. + = gain, - = loss of a pattern element. "Repressor" suggests
stripes may be broken into spots by repressors of pigmentation and vice versa. The illustration does
not imply any evolutionary direction; it solely illustrates the modularity of these complex
patterns.

We hypothesized that the developmental candidate genes may activate the *y* gene through four
CRMs, each controlling one sub-pattern to assemble the complete melanin pattern. We searched
for these CRMs by transforming *D. guttifera* with *DsRed* reporter constructs containing non-

coding fragments of the 42 kb D. guttifera y gene locus¹² (Extended Data Fig. 1). Surprisingly, 91 only one 953 bp fragment from the y intron, the gut y spot CRM, drove expression closely 92 resembling all six spot rows on the developing abdomen (Fig. 4). To isolate possible sub-pattern-93 inducing CRMs, we subdivided the gut y spot CRM into 8 partially overlapping sub-fragments. 94 Unexpectedly, the 636 bp left sub-fragment displayed horizontal stripe expression along the 95 96 posterior edges of each abdominal segment, while the 570 bp right fragment was inactive (#1 & #2, Fig. 4.). Further dissection of this CRM revealed a 259 bp sub-fragment, which contained the 97 minimal gut y core stripe CRM with some additional dorsal midline shade activity (#7, Fig. 4). 98 99 These results suggest that the D. guttifera spots may have evolved from an ancestral stripe pattern that became partially repressed to isolate the spots. Currently, we cannot offer any direct 100 evidence for specific candidate repressor genes. Neither the *in situ* hybridization experiments nor 101 102 the bioinformatics analyses, using Jaspar, resulted in putative pigment stripe repressors. Although we identified 24 Engrailed (En)-binding sites and 19 Homothorax (Hth)-binding sites 103 in the gut y spot CRM (both are known repressors of pigmentation in *Drosophila*^{15,22}), these sites 104 were not enriched in the right half of the CRM, as we would have expected. However, our 105 transcription factor binding site analysis of the gut y spot CRM sequence revealed putative 106 107 transcription factor binding sites for most of the developmental genes that we identified as potential activators in our *in situ* hybridization screen, except for *hh*. This suggests that localized 108 109 spot activation by these developmental factors contributes to the formation of the pattern.



- 111 Extended Data Fig. 1: The y gene locus. The horizontal bars indicate the DNA fragments of the
- 112 *D. guttifera y* gene that were tested in transgenic *D. guttifera* for regulatory activity.

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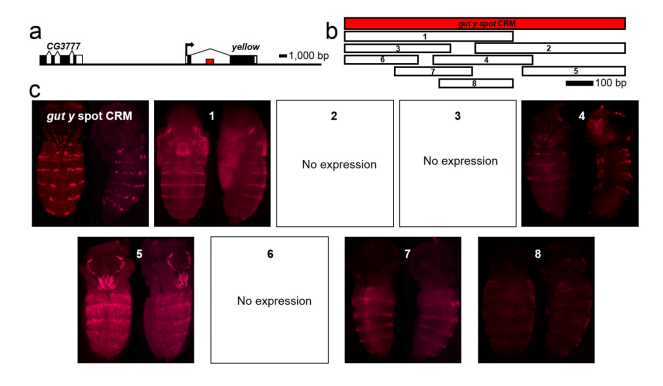


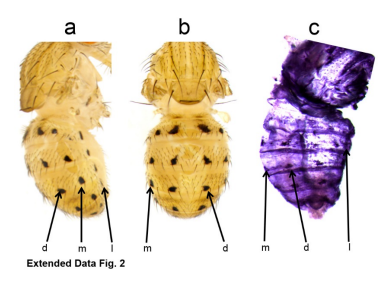
Fig. 4: The *gut y spot* CRM is harbored within the *y* intron. a, The *y* gene locus. The red bar indicates the relative position of the *gut y spot* CRM. b, Sub-dividing the *gut y spot* CRM revealed horizontal stripes on each abdominal segment. The white bars (1-8) represent subfragments of the *gut y spot* CRM, and the corresponding pupal *DsRed* expression patterns in transgenic *D. guttifera* are shown.

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121 Next, we asked whether the abdominal pigment spot pattern of a species closely related to *D*.

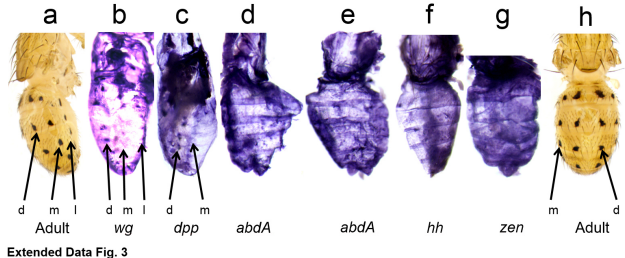
- 122 guttifera shares a similar developmental basis. We thus performed in situ hybridization
- 123 experiments in *Drosophila deflecta* (*D. deflecta*). This species displays six longitudinal spot rows

on its abdomen, but lacks the dorsal midline shade (Extended Data Fig. 2a, b). As in D. guttifera, 124 y mRNA in D. deflecta pupal abdomens was expressed in six rows of spots, except along the 125 dorsal midline (Extended Data Fig. 2c). Similarly, wg foreshadowed all six rows of spots, while 126 dpp expression matched all but the lateral spot rows (Extended Data Fig. 3b, c). In contrast to the 127 D. guttifera results, abd-A, hh, and zen were absent along the dorsal midline, which is in 128 129 agreement with the lack of pigment in *D. deflecta* adults (Extended Data Fig. 3d, e, f, g). However, abd-A expression was not detectable where the lateral spot rows will form (Extended 130 Data Fig. 3d), suggesting that these particular spots are controlled differently in D. deflecta. We 131 132 next cloned the 938 bp orthologous *def y spot* CRM and transformed it into *D. guttifera*, using the *DsRed* reporter assay. The *def y spot* CRM drove faint dorsal spot row and stripe expression, 133 especially along the dorsal spots (Extended Data Fig. 4). We further subdivided the def y spot 134 135 CRM into 8 sub-fragments and identified a minimal def y core stripe CRM (288 bp) (#7, Extended Data Fig. 4). This sub-fragment drove a striped pattern, but without the dorsal midline 136 shade activity seen in the D. guttifera minimal gut y core stripe CRM (#7, Fig. 4). We further 137 transformed the gut y spot and def y spot CRMs including all sub-fragments into D. 138 melanogaster to test if D. melanogaster trans-factors can bind to and activate these two quinaria 139 140 group species' spot CRMs. As a result, none of the reporter constructs showed any expression (data not shown). This suggests that the hypothetical ancestral stripe pattern of the quinaria 141 group and the pigment stripes found on the *D. melanogaster* abdominal tergites²³ have evolved 142 143 independently by changes in *trans*. As the spot CRMs from D. guttifera and D. deflecta are not orthologous to any sequences within the *D. melanogaster y* locus, changes in *cis* have also 144 contributed to the diversification of pigment patterns between D. melanogaster and the quinaria 145 146 species group.



Extended Data Fig. 2: The *y* gene expression pattern in *D. deflecta* foreshadows the black spot
pattern on the adult abdomen. a, Adult lateral view. b, Adult dorsal view. c, *y* mRNA in the pupal
epidermis. d = dorsal, m = median, l = lateral.

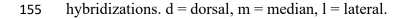


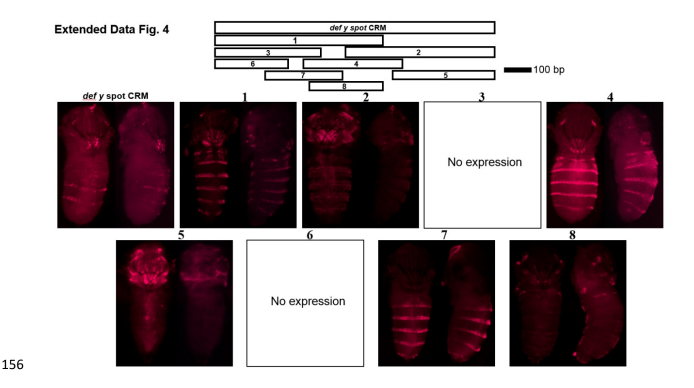


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153 Extended Data Fig. 3: Developmental gene expression patterns in *D. deflecta* foreshadow

distinct subsets of the adult abdominal color pattern. a, h, Adult, lateral view. b-g, Pupal *in situ*





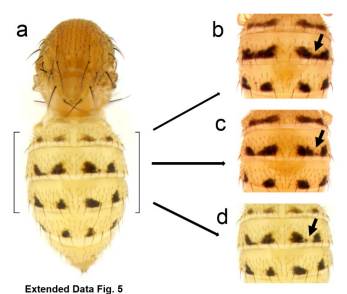
Extended Data Fig. 4: The orthologous *D. deflecta* region (*def y spot* CRM) analyzed in
transgenic *D. guttifera*. The white bars (1-8) indicate sub-fragments of the *def y spot* CRM that
were tested for reporter activity. The corresponding DsRed reporter expression patterns in
developing pupae are shown.

161

In contrast to the *D. guttifera* wing spot pattern¹², the abdominal pigment pattern develops in the 162 absence of visible physical landmarks. wg, dpp, and hh are homologous to known proto-163 oncogenes in humans²⁴, while *zen* and *abdA* are Hox genes. The abdominal color pattern of D. 164 165 guttifera appears to be regulated by multiple developmental pathways consisting of activators and repressors acting in parallel, possibly targeting pigmentation genes other than *y* as 166 well^{18,19,25,26}. Further evidence for the repression of stripes can be seen in *Drosophila falleni*'s 167 intraspecific pigment variation, another member of the quinaria species group (Extended Data 168 Fig. 5). Our multi-pathway model fits well with the observation that the abdominal pattern 169

- 170 variation presented by quinaria group members is largely due to modular derivations from the *D*.
- 171 *guttifera* ground plan (Fig. 3). This scenario is reminiscent of the modularity found in butterfly
- 172 wing patterns. Because insects use similar genes for color pattern development $^{21,27-30}$, the
- 173 quinaria group may serve as a valuable model to understand insect color pattern evolution.
- 174 Future work should aim to manipulate the genes involved in pigmentation to test if they interact
- according to the reaction-diffusion model, as predicted by Alan Turing 31 .





177

178 Extended Data Fig. 5: The abdominal pigment stripes of *D. falleni* break down into

179 pigment spots. Intraspecific variation, as illustrated here, is very common in *D. falleni*. a, Adult

abdomen. b-d, Adult abdominal pigment spots developing from stripe repression (arrows).

181

182

184 Materials and Methods

185 Molecular procedures

- 186 In situ hybridizations were carried out with species-specific RNA probes, as described in¹², but
- 187 with abdomens cut into halves and cleaned from the internal organs. At least three positive pupae
- 188 were observed for each result shown. Additional images for verification purposes are provided in
- 189 Extended Data Figs. 6-16. Immunohistochemistry for the Y protein in abdomens was performed
- according to¹⁵, with abdomens cut in half and cleaned with 1X PBS. *D. guttifera* CRMs were
- identified and tested in *D. guttifera* according to¹² and in *D. melanogaster* as described in²³.
- 192 Transgenic experiments were performed as outlined in 32 . Pupal stages were identified according
- 193 to^{33} .

194 Drosophila stocks

- 195 All fly stocks were a kind gift from the Sean B. Carroll Laboratory (University of Wisconsin -
- 196 Madison) and were cultured at room temperature. We used the *D. melanogaster* fly strain
- 197 VK00006 (cytogenic location 19E7), the *D. guttifera* stock no.15130–1971.10, and the *D*.
- 198 *deflecta* stock no. 15130-2018.00 for gene expression and transgenic analyses.

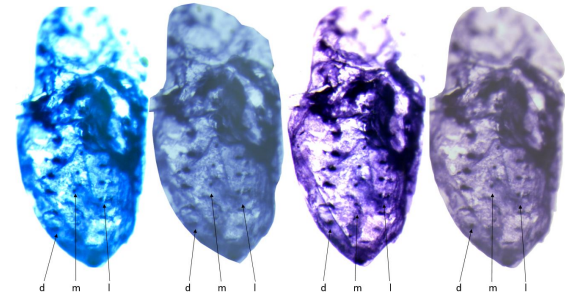
199 PCR primer sequences

- 200 We used the following primers to amplify the CRM sequences:
- 201 (iii) gut y spot CRM: Fwd: 5'-CAGCTGCGGTTGAGTACGAC-3' and Rvs: 5'-
- 202 GCCAACTCGACGGGAATTC-3'. Restriction sites: KpnI and SacII.
- 203 (iv) def y spot CRM: Fwd: 5'-CAGCTGCTGCGGTTCAGTAG-3' and Rvs: 5'-
- 204 GCTAGACACGTTGGTTTGCT-3'. Restriction sites: KpnI and SacII.

- 205 (v) gut y spot CRM sub-fragment #1: Fwd: 5'-CAGCTGCGGTTGAGTACGAC-3' and Rvs: 5'-
- 206 ACTGAATCTGATTTCGGCTCG-3'. Restriction sites: KpnI and SacII.
- 207 (vi) gut y spot CRM sub-fragment #2: Fwd: 5'-AGTTAATCGCCAGTCAATAATGGC-3' and
- 208 Rvs: 5'- GAATTCCCGTCGAGTTGGC-3'. Restriction sites: KpnI and SacII.
- 209 (vii) gut y spot CRM sub-fragment #3: Fwd: 5'-CAGCTGCGGTTGAGTACGAC-3' and Rvs:
- 210 5'-GCCATTATTGACTGGCGATTAAC-3'. Restriction sites: KpnI and SacII.
- 211 (viii) gut y spot CRM sub-fragment #4: Fwd: 5'-AAATGAAGCTCAGTGAGCCGC-3' and Rvs:
- 212 5'-ACTGAATCTGATTTCGGCTCG-3'. Restriction sites: KpnI and SacII.
- (ix) *gut y spot* CRM sub-fragment #5: Fwd: 5'-AGCATCTGAAACTTAAACGCCG-3' and Rvs:
- 214 5'-GAATTCCCGTCGAGTTGGC-3'. Restriction sites: KpnI and SacII.
- 215 (x) gut y spot CRM sub-fragment #6: Fwd: 5'-CAGCTGCGGTTGAGTACGAC-3' and Rvs: 5'-
- 216 CAGCGATATTAATTTTTTTTTTTTTTCAATGG-3'. Restriction sites: KpnI and SacII.
- 217 (xi) gut y spot CRM sub-fragment #7(gut y core stripe CRM): Fwd: 5'-
- 218 AAATGAAGCTCAGTGAGCCGC-3' and Rvs: 5'-GCGATTTGTTGTCAAGTCAAC-3'.
- 219 Restriction sites: KpnI and SacII.
- 220 (xii) gut y spot CRM sub-fragment #8: Fwd: 5'-AAATGAAGCTCAGTGAGCCGC-3' and Rvs:
- 221 5'-GTTGACTTGACAAACAAATCGC-3'. Restriction sites: KpnI and SacII.
- 222 (xiii) *def y spot* CRM sub-fragment #1: Fwd: 5'-CAGCTGCTGCGGTTCAGTAG-3' and Rvs:
- 223 5'-ATTGTCGCAGCTGCCTAACG-3'. Restriction sites: KpnI and SacII.
- 224 (xiv) *def y spot* CRM sub-fragment #2: Fwd: 5'-AACGAAGCTCACTGAGCTGC-3' and Rvs:
- 225 5'-AGCAAACCAACGTGTGTCTAGC-3'. Restriction sites: KpnI and SacII.

- 226 (xv) *def y spot* CRM sub-fragment #3: Fwd: 5'-CAGCTGCTGCGGTTCAGTAG-3' and Rvs:
- 227 5'-GTTAAAAGCAGCCAGTTGGCC-3'. Restriction sites: KpnI and SacII.
- 228 (xvi) def y spot CRM sub-fragment #4: Fwd: 5'-CAAAGAATCGAATTCGGAGACAG-3' and
- 229 Rvs: 5'-ATTGTCGCAGCTGCCTAACG-3'. Restriction sites: KpnI and SacII. (Clone name: *def*
- 230 *y* 1.1C2)
- 231 (xvii) *def y spot* CRM sub-fragment #5: Fwd: 5'-GAATGAGATTCGTTAGGCAGC-3' and Rvs:
- 232 5'-AGCAAACCAACGTGTGTCTAGC-3'. Restriction sites: KpnI and SacII.
- 233 (xviii) *def y spot* CRM sub-fragment #6: Fwd: 5'-CAGCTGCTGCGGTTCAGTAG-3' and Rvs:
- 234 5'-TTCAACGGATATTCGTTCAATTTC-3'. Restriction sites: KpnI and SacII.
- 235 (xix) def y spot CRM sub-fragment #7 (def y core stripe CRM): Fwd: 5'-
- 236 CAAAGAATCGAATTCGGAGACAG-3' and Rvs: 5'-GTCAGGCAATGTAAATGTTGTCG-
- 237 3'. Restriction sites: KpnI and SacII.
- 238 (xx) *def y spot* CRM sub-fragment #8: Fwd: 5'-AACGAAGCTCACTGAGCTGC-3' and Rvs:
- 239 5'-ATTGTCGCAGCTGCCTAACG-3'. Restriction sites: KpnI and SacII.
- 240 These forward and reverse primer sequences do not include restriction sites.

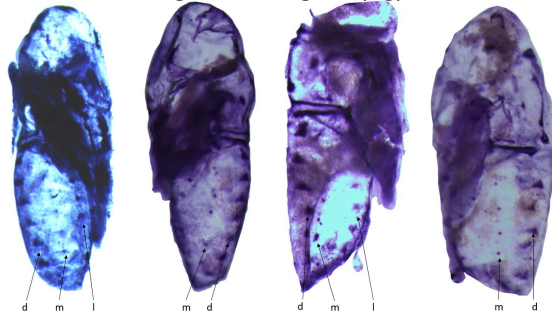
D. guttifera wingless (wg)



242 Extended Data Fig. 6: One *D. guttifera* pupa stained with a *wg* probe. d = dorsal,

243 m = median, l = lateral row of spots. Different image manipulations shown.

D. guttifera wingless (wg)

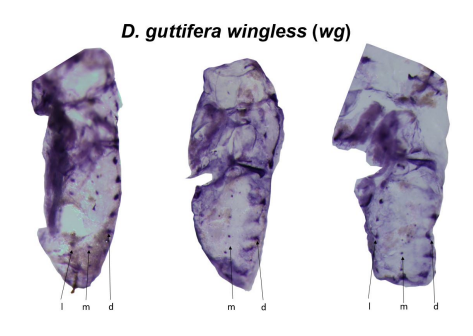


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Extended Data Fig. 7: *D. guttifera* **pupae stained with a** *wg* **probe.** d = dorsal, m = median,

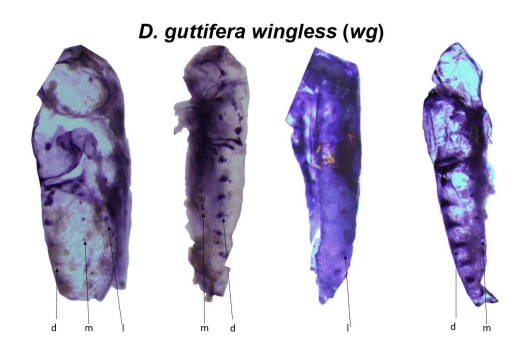
l = lateral row of spots.



- 248 Extended Data Fig. 8: D. guttifera pupae stained with a wg probe. d = dorsal, m = median,
- l = lateral row of spots.

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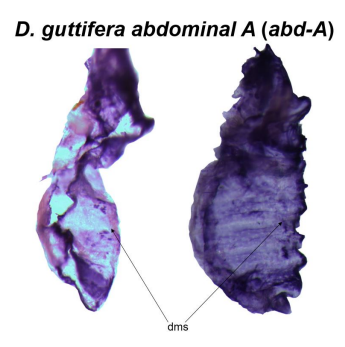
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252 Extended Data Fig. 9: *D. guttifera* pupae stained with a *wg* probe. d = dorsal, m = median,

l = lateral row of spots.



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255 Extended Data Fig. 10: *D. guttifera* pupae stained with an *abd-A* probe. dms = dorsal midline

shade.

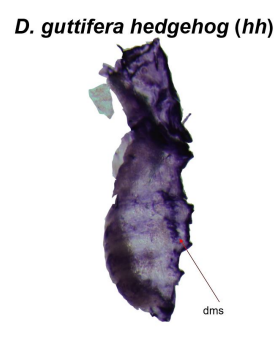
D. guttifera decapentaplegic (dpp)



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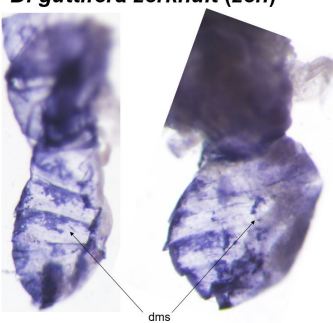
258 Extended Data Fig. 11: D. guttifera pupa stained with a dpp probe. d = dorsal, m = median

row of spots.



260

- 261 Extended Data Fig. 12: *D. guttifera* pupa stained with a *hh* probe. dms = dorsal midline
- shade.



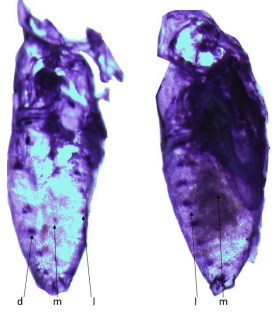
D. guttifera zerknult (zen)

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264 Extended Data Fig. 13: *D. guttifera* pupae stained with a *zen* probe. dms = dorsal midline

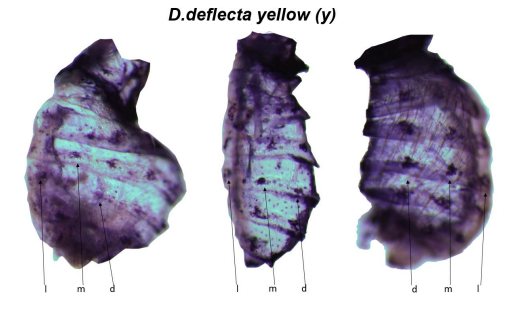
shade.



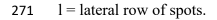


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- 267 Extended Data Fig. 14: *D. deflecta* pupae stained with a *wg* probe. d = dorsal, m = median,
- l = lateral row of spots.



270 Extended Data Fig. 15: *D. deflecta* pupae stained with a *y* probe. d = dorsal, m = median,



D.melanogaster yellow (y)





272

273 Extended Data Fig. 16: *D. melanogaster* pupae stained with a *y* probe.

274

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278

279 Author Contributions

- 280 K.K.B.R., M.S. and T.W. conceived and designed the experiments; K.K.B.R, P.M.E.N., T.E.S,
- E.B., P.P.K, A.McQ., E.M., A.A., A.N. and T.W. performed the experiments; K.K.B.R., S.M.
- and T.W. analyzed the data; T.W. obtained funding; K.K.B.R and T.W. wrote the paper;
- K.K.B.R, S.M., P.M.E.N., T.E.S., E.B., P.P.K., A.McQ., E.M., A.A., A.N. and T.W. edited the
- 284 paper.

285 Competing Financial Interests

286 The authors declare no competing financial interests.

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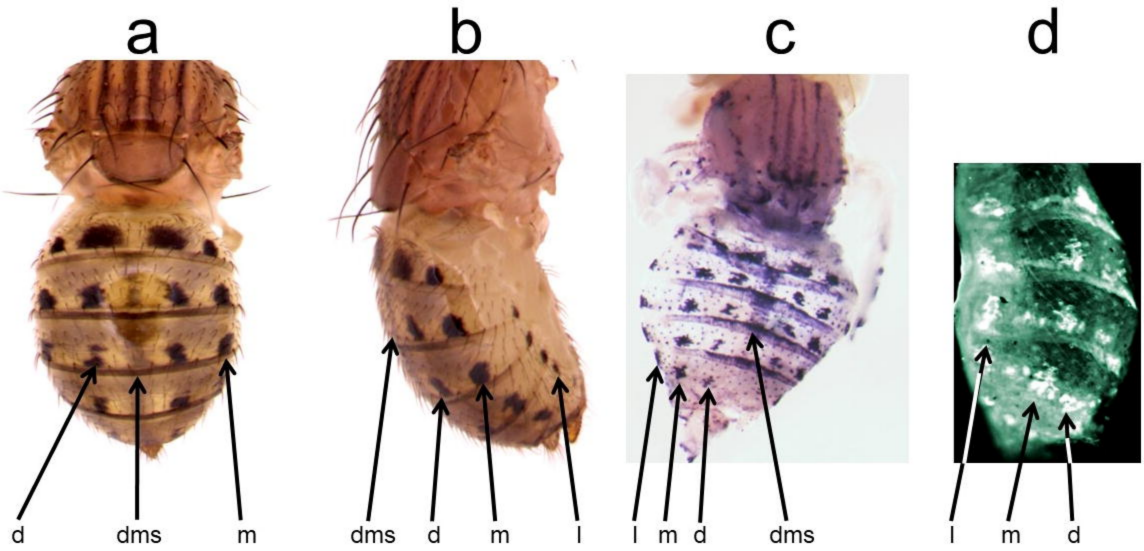
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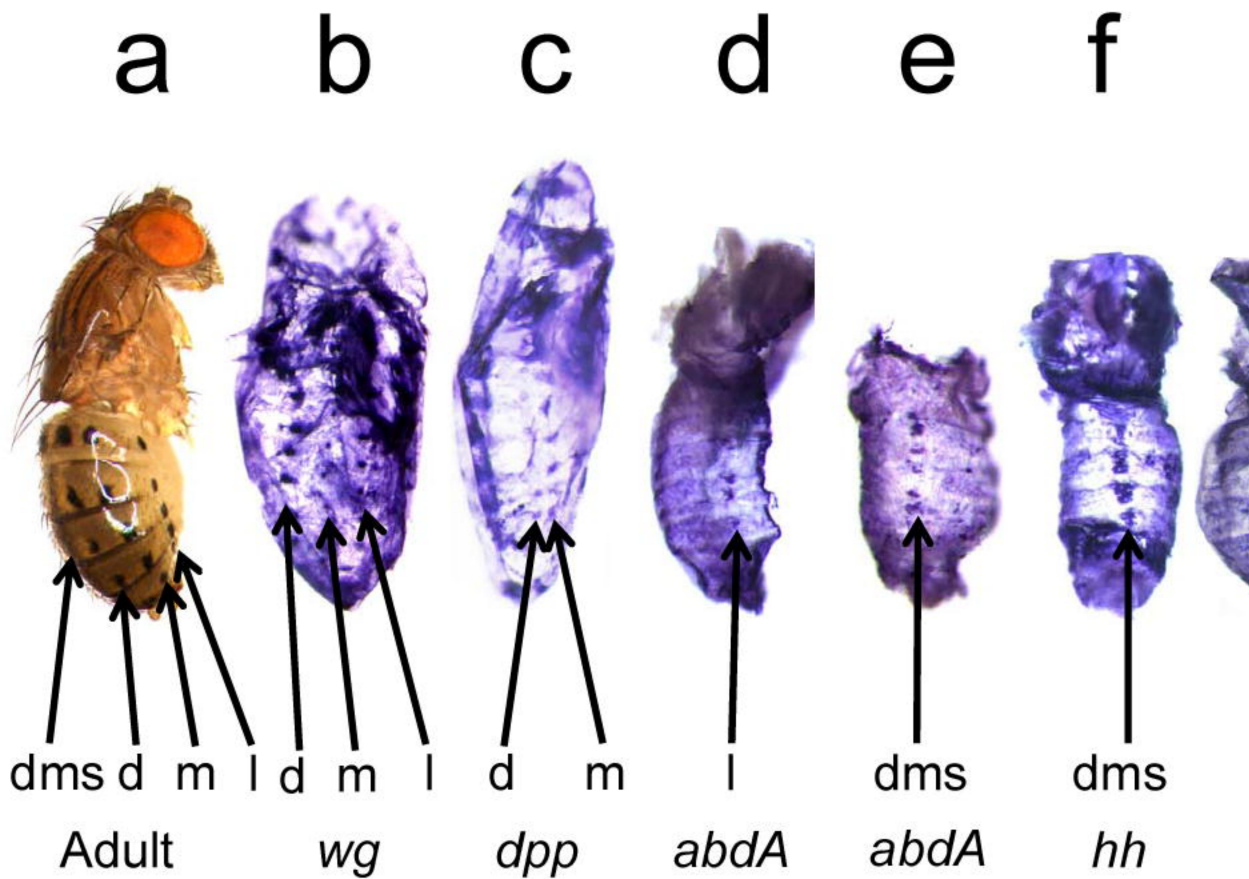
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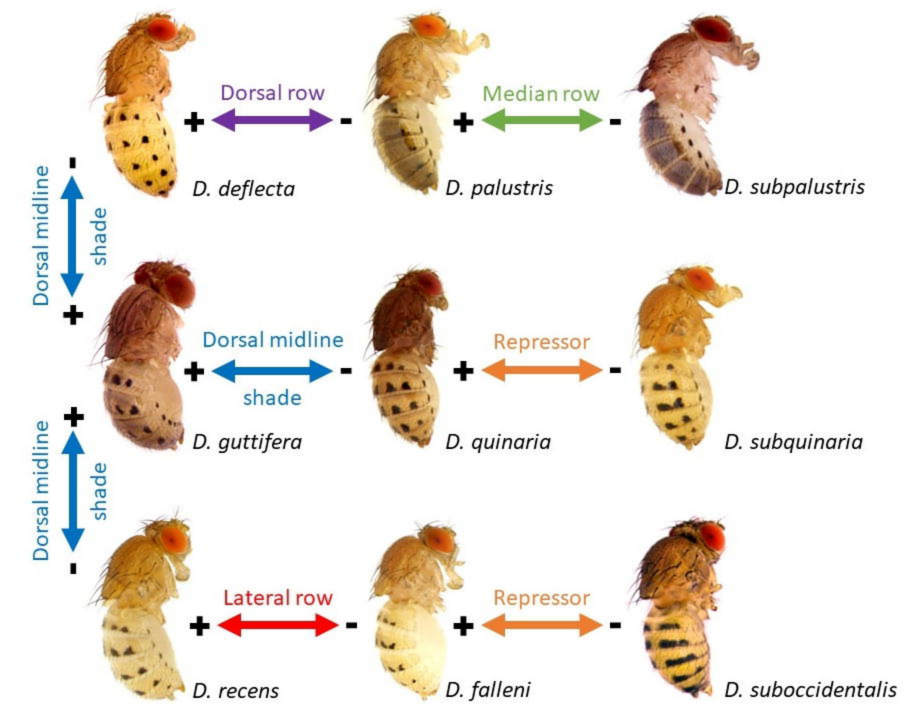


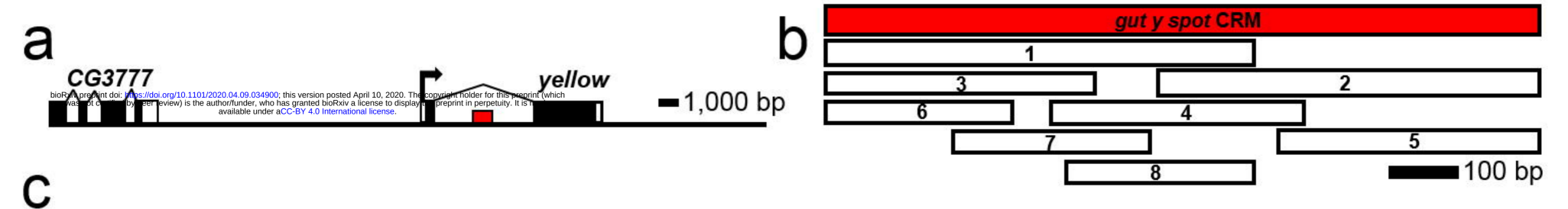
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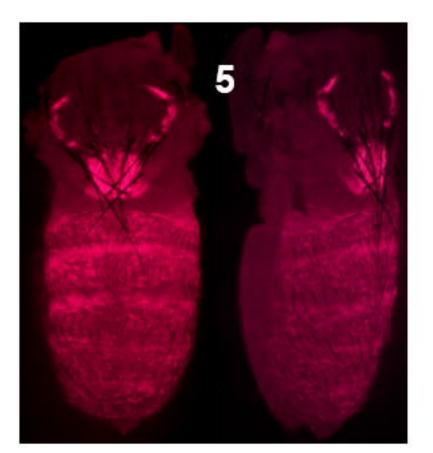
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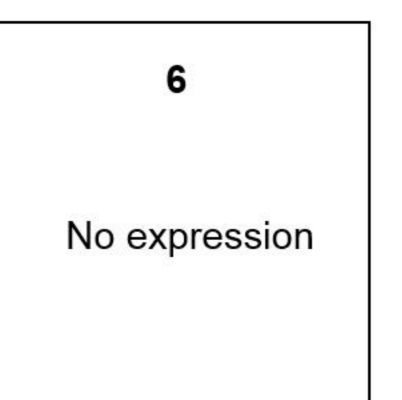


2

No expression



gut y spot CRM



3

No expression

