#### The gene *cortex* controls mimicry and crypsis in butterflies and moths

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30 The wing patterns of butterflies and moths (Lepidoptera) are diverse and striking examples of evolutionary diversification by natural selection<sup>1,2</sup>. Lepidopteran wing colour patterns are a 31 key innovation, consisting of arrays of coloured scales. We still lack a general understanding 32 33 of how these patterns are controlled and if there is any commonality across the 160,000 moth and 17,000 butterfly species. Here, we identify a gene, cortex, through fine-scale mapping 34 using population genomics and gene expression analyses, which regulates pattern switches in 35 multiple species across the mimetic radiation in *Heliconius* butterflies. *cortex* belongs to a 36 fast evolving subfamily of the otherwise highly conserved fizzy family of cell cycle 37 regulators<sup>3</sup>, suggesting that it most likely regulates pigmentation patterning through 38 regulation of scale cell development. In parallel with findings in the peppered moth (Biston 39 *betularia*)<sup>4</sup>, our results suggest that this mechanism is common within Lepidoptera and that 40 41 cortex has become a major target for natural selection acting on colour and pattern variation in this group of insects. 42

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44 In *Heliconius*, there is a major effect locus, Yb, that controls a diversity of colour pattern elements across the genus. It is the only locus in Heliconius that regulates all scale types and 45 colours, including the diversity of white and yellow pattern elements in the two co-mimics H. 46 melpomene (Hm) and H. erato (He), but also whole wing variation in black, yellow, white, 47 and orange/red elements in *H. numata*  $(Hn)^{5-7}$ . In addition, genetic variation underlying the 48 Bigeye wing pattern mutation in Bicyclus anynana, melanism in the peppered moth, Biston 49 betularia, and melanism and patterning differences in the silkmoth, Bombyx mori, have all 50 been localised to homologous genomic regions  $^{8-10}$  (Fig 1). Therefore, this genomic region 51 appears to contain one or more genes that act as major regulators of wing pigmentation and 52 patterning across the Lepidoptera. 53

Previous mapping of this locus in *He*, *Hm* and *Hn* identified a genomic interval of  $\sim 1 \text{Mb}^{11-13}$ 54 (Extended Data Table 1), which also overlaps with the 1.4Mb region containing the 55 carbonaria locus in *B. betularia*<sup>9</sup> and a 100bp non-coding region containing the *Ws* mutation 56 in *B. mori*<sup>10</sup> (Fig 1). We took a population genomics approach to identify single nucleotide 57 polymorphisms (SNPs) most strongly associated with phenotypic variation within the ~1Mb 58 Heliconius interval. The diversity of wing patterning in Heliconius arises from divergence at 59 wing pattern loci<sup>7</sup>, while convergent patterns generally involve the same loci and sometimes 60 even the same alleles<sup>14–16</sup>. We used this pattern of divergence and sharing to identify SNPs 61 62 associated with colour pattern elements across many individuals from a wide diversity of 63 colour pattern phenotypes (Fig 2).

In three separate *Heliconius* species, our analysis consistently implicated the gene *cortex* as 64 being involved in adaptive differences in wing colour pattern. In He the strongest associations 65 66 with the presence of a yellow hindwing bar were centred around the genomic region containing *cortex* (Fig 2A). We identified 108 SNPs that were fixed for one allele in He 67 68 favorinus, and fixed for the alternative allele in all individuals lacking the yellow bar, the majority of which were in introns of cortex (Extended Data Table 2). 15 SNPs showed a 69 similar fixed pattern for He demophoon, which also has a yellow bar. These were non-70 overlapping with those in *He favorinus*, consistent with the hypothesis that this phenotype 71 evolved independently in the two disjunct populations $^{17}$ . 72

Previous work has suggested that alleles at the *Yb* locus are shared between *Hm* and the
closely related species *H. timareta*, and also the more distantly related species *H. elevatus*,
resulting in mimicry between these species<sup>18</sup>. Across these species, the strongest associations
with the yellow hindwing bar phenotype were again found at *cortex* (Fig 2D, Extended Data
Fig 1A and Table 3). Similarly, the strongest associations with the yellow forewing band
were found around the 5' UTRs of *cortex* and gene *HM00036*, an orthologue of *D*.

79 melanogaster washout gene. A single SNP ~17kb upstream of cortex (the closest gene) was perfectly associated with the yellow forewing band across all Hm, H. timareta and H. 80 elevatus individuals (Extended Data Fig 1A, Fig 2 and Table 3). We found no fixed coding 81 82 sequence variants at *cortex* in a larger sample (43-61 individuals) of *Hm aglaope* and *Hm* amaryllis (Extended Data Figure 3, Supplementary Information), which differ in Yb 83 controlled phenotypes<sup>19</sup>, suggesting that functional variants are likely to be regulatory rather 84 than coding. We found extensive transposable element variation around *cortex* but it is 85 unclear if any of these associate with phenotype (Extended Data Figure 3 and Table 4; 86 87 Supplementary Information).

Finally, in *Hn* large inversions at the *P* supergene locus (Fig 1) are associated with different 88 morphs<sup>13</sup>. There is a steep increase in genotype-by-phenotype association at the breakpoint of 89 inversion 1, consistent with the role of these inversions in reducing recombination (Fig 2E). 90 91 However, the *bicoloratus* morph can recombine with all other morphs across one or the other inversion, permitting finer-scale association mapping of this region. As in *He* and *Hm*, this 92 93 analysis showed a narrow region of associated SNPs corresponding exactly to the *cortex* gene 94 (Fig 2E), again with the majority of SNPs in introns (Extended Data Table 2). This associated region does not correspond to any other known genomic feature, such as an inversion or 95 96 inversion breakpoint.

To determine whether sequence variants around *cortex* were regulating its expression we investigated gene expression across the *Yb* locus. We used a custom designed microarray including probes from all predicted genes in the *H. melpomene* genome<sup>18</sup>, as well as probes tiled across the central portion of the *Yb* locus, focussing on two naturally hybridising *Hm* races (*plesseni* and *malleti*) that differ in *Yb* controlled phenotypes<sup>7</sup>. *cortex* was the only gene across the entire interval to show significant expression differences both between races with different wing patterns and between wing sections with different pattern elements (Fig 3). 104 This finding was reinforced in the tiled probe set, where we observed strong differences in expression of *cortex* exons and introns but few differences outside this region (Extended Data 105 Table 2). cortex expression was higher in Hm malleti than Hm plesseni in all three wing 106 107 sections used (but not eyes) (Fig 3C; Extended Data Fig 4C). When different wing sections were compared within each race, *cortex* expression in *Hm malleti* was higher in the distal 108 section that contains the Yb controlled yellow forewing band, consistent with cortex 109 110 producing this band. In contrast, *Hm plesseni*, which lacks the yellow band, had higher cortex expression in the proximal forewing section (Fig 3F; Extended Data Fig 4J). Expression 111 112 differences were found only in day 1 and day 3 pupal wings rather than day 5 or day 7 (Extended Data Fig 4), similar to the pattern observed previously for the transcription factor 113 opti $x^{20}$ . 114

Differential expression was not confined to the exons of *cortex*; the majority of differentially 115 116 expressed probes in the tiling array corresponded to cortex introns (Fig 3). This does not appear to be due to transposable element variation (Extended Data Table 2), but may be due 117 118 to elevated background transcription and unidentified splice variants. RT-PCR revealed a diversity of splice variants (Extended Data Fig 5), and sequenced products revealed 8 non-119 120 constitutive exons and 6 variable donor/acceptor sites, but this was not exhaustive 121 (Supplementary Information). We cannot rule out the possibility that some of the differentially expressed intronic regions could be distinct non-coding RNAs. However, qRT-122 PCR in other hybridising races with divergent Yb alleles (aglaope/amaryllis and 123 rosina/melpomene) also identified expression differences at cortex and allele-specific splicing 124 differences between both pairs of races (Extended Data Figs 1 and 5, Supplementary 125 Information). 126

Finally, *in situ* hybridisation of *cortex* in final instar larval hindwing discs showed expressionin wing regions fated to become black in the adult wing, most strikingly in their

129 correspondence to the black patterns on adult *Hn* wings (Fig 4). In contrast, the array results
130 from pupal wings were suggestive of higher expression in non-melanic regions. This may
131 suggest that *cortex* is upregulated at different time-points in wing regions fated to become
132 different colours.

Overall, cortex shows significant differential expression and is the only gene in the candidate 133 region to be consistently differentially expressed in multiple race comparisons and between 134 differently patterned wing regions. Coupled with the strong genotype-by-phenotype 135 136 associations across multiple independent lineages (Extended Data Table 1), this strongly 137 implicates *cortex* as a major regulator of colour and pattern. However, we have not excluded the possibility that other genes in this region also influence pigmentation patterning. A 138 prominent role for *cortex* is also supported by studies in other taxa; our identification of 139 distant 5' untranslated exons of *cortex* (Supplementary Information) suggests that the 100bp 140 interval containing the Ws mutation in B. mori is likely to be within an intron of cortex and 141 not in intergenic space as previously thought<sup>10</sup>. In addition, fine-mapping and gene 142 expression also implicate *cortex* as controlling melanism in the peppered moth<sup>4</sup>. 143

It seems likely that *cortex* controls pigmentation patterning through control of scale cell 144 development. The *cortex* gene falls in an insect specific lineage within the fizzy/CDC20 145 146 family of cell cycle regulators (Extended Data Fig 6A). The phylogenetic tree of the gene family highlighted three major orthologous groups, two of which have highly conserved 147 functions in cell cycle regulation mediated through interaction with the anaphase promoting 148 complex/cyclosome  $(APC/C)^{3,21}$ . The third group, cortex, is evolving rapidly, with low amino 149 acid identity between D. melanogaster and Hm cortex (14.1%), contrasting with much higher 150 identities for orthologues between these species in the other two groups (fzy, 47.8% and 151 152 rap/fzr,47.2%, Extended Data Fig 6A). Drosophila melanogaster cortex acts through a

similar mechanism to fzy in order to control meiosis in the female germ line<sup>22-24</sup>. *Hm* cortex 153 also has some conservation of the fizzy family C-box and IR elements (Supplementary 154 Information) that mediate binding to the  $APC/C^{23}$ , suggesting that it may have retained a cell 155 cycle function, although we found that expressing *Hm cortex* in *D. melanogaster* wings 156 produced no detectable effect (Extended Data Fig 6, Supplementary Information). 157 Previously identified butterfly wing patterning genes have been transcription factors or 158 signalling molecules<sup>20,25</sup>. Developmental rate has long been thought to play a role in 159 lepidopteran patterning<sup>26,27</sup>, but *cortex* was not a likely *a priori* candidate, because its 160 *Drosophila* orthologue has a highly specific function in meiosis<sup>23</sup>. The recruitment of *cortex* 161 to wing patterning appears to have occurred before the major diversification of the 162 Lepidoptera and this gene has repeatedly been targeted by natural selection<sup>1,7,9,28</sup> to generate 163 both cryptic<sup>4</sup> and aposematic patterns. 164

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- 240 **Supplementary Information** is linked to the online version of the paper at
- 241 <u>www.nature.com/nature</u>.

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253	Author Contributions NJN performed the association analyses, 5' RACE, RT-PCR, qRT-
254	PCR and prepared the manuscript. NJN and CDJ co-ordinated the research. CP-D performed
255	and analysed the microarray and RNAseq experiments. AW performed the Hn association
256	analysis. MS assembled and annotated the HeCr BAC reference and the He alignments. SVS
257	performed in situ hybridizations. RWRW performed the transgenic experiments and analysis
258	of de novo assembled sequences and fosmids together with JJH. GW and LF initially
259	identified splicing variants of cortex. LM performed crosses between Hm races. HH screened
260	the HeCr BAC library. CS and RM provided samples. AD contributed to the Hm BAC
261	sequencing and annotation. R-fC, MJ, VL, WOM and CDJ are PIs who obtained funding and
262	led the project elements. All authors commented on the manuscript.

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Author Information Short read sequence data generated for this study are available from 264 265 ENA (http://www.ebi.ac.uk/ena) under study accession PRJEB8011 and PRJEB12740 (see Supplementary Table 1 for previously published data accessions). The updated Cr contig is 266 deposited in Genbank with accession KC469893. The assembled Hm fosmid sequences are 267 deposited in Genbank with accessions KU514430-KU514438. The microarray data are 268 deposited in GEO with accessions GSM1563402- GSM1563497. Reprints and permissions 269 270 information is available at www.nature.com/reprints. Correspondence and requests for materials should be addressed to n.nadeau@sheffield.ac.uk or c.jiggins@zoo.cam.ac.uk 271

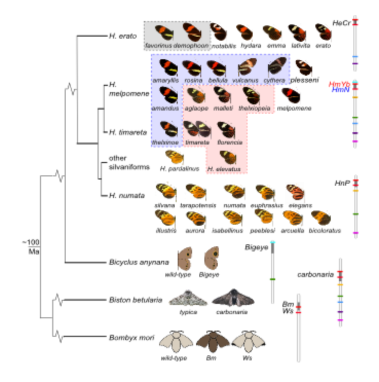




Figure 1. A homologous genomic region controls a diversity of phenotypes across the 275 Lepidoptera. Left: phylogenetic relationships<sup>29</sup>. Right: chromosome maps with colour pattern 276 intervals in grey, coloured bars represent markers used to assign homology<sup>5,8–10</sup>, the first and 277 last genes from Fig 2 shown in red. In He the HeCr locus controls the yellow hind-wing bar 278 phenotype (grey boxed races). In Hm it controls both the yellow hind-wing bar (HmYb, pink 279 box) and the yellow forewing band (HmN, blue box). In Hn it modulates black, yellow and 280 orange elements on both wings (*HnP*), producing phenotypes that mimic butterflies in the 281 genus Melinaea. Morphs/races of Heliconius species included in this study are shown with 282 283 names.

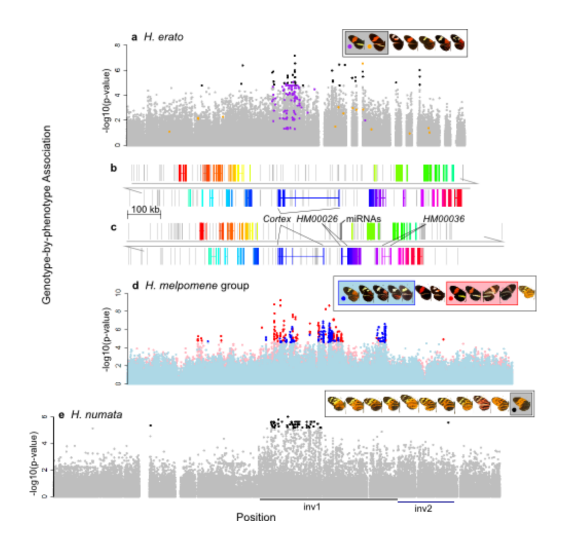


Figure 2. Association analyses across the genomic region known to contain major colour 285 286 pattern loci in *Heliconius*. A) Association in *He* with the yellow hind-wing bar (n=45). Coloured SNPs are fixed for a unique state in He demophoon (orange) or He favorinus 287 (purple). B) Genes in *He* with direct homologs in *Hm*. Genes are in different colours with 288 exons (coding and UTRs) connected by a line. Grey bars are transposable elements. C) Hm 289 genes and transposable elements: colours correspond to homologous He genes; MicroRNAs<sup>30</sup> 290 291 in black. D) Association in the Hm/timareta/silvaniform group with the yellow hind-wing bar (red) and yellow forewing band (blue) (n=49). E) Association in Hn with the bicoloratus 292 morph (n=26); inversion positions<sup>13</sup> shown below. In all cases black/dark coloured points are 293 above the strongest associations found outside the colour pattern scaffolds (*He* p=1.63e-05; 294 295 *Hm* p=2.03e-05 and p=2.58e-05; *Hn* p=6.81e-06).

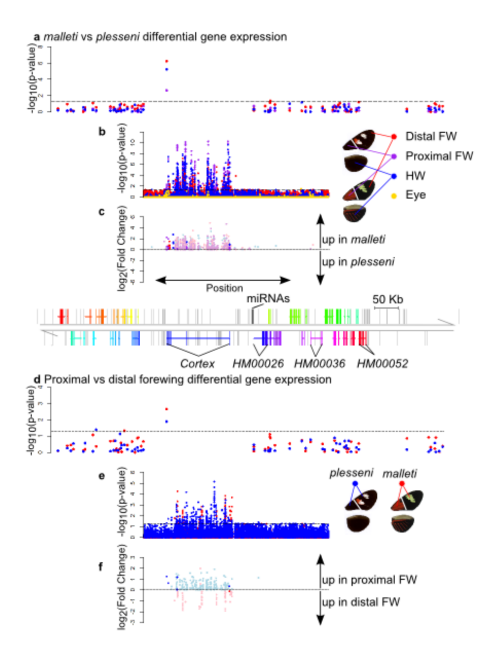
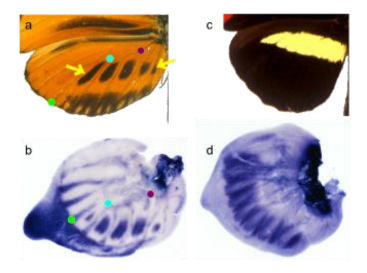


Figure 3. Differential gene expression across the genomic region known to contain major colour pattern loci in *Heliconius melpomene*. Expression differences in day 3 pupae, for all genes in the *Yb* interval (A,D) and tiling probes spanning the central portion of the interval (B,C,E,F). Expression is compared between races for each wing region (A,B,C) and between proximal and distal forewing sections for each race (D,E,F). C and F: magnitude and direction of expression difference (log<sub>2</sub> fold-change) for tiling probes showing significant differences ( $p \le 0.05$ ); probes in known *cortex* exons shown in dark colours. Gene *HM00052* 

- 304 was differentially expressed between other races in RNA sequence data (Supplementary
- 305 Information) but is not differentially expressed here.



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Figure 4. *In situ* hybridisations of *cortex* in hind-wings of final instar larvae. B) *Hn tarapotensis*; adult wing shown in A, coloured points indicate landmarks, yellow arrows highlight adult pattern elements corresponding to the *cortex* staining. D) *Hm rosina*; adult wing shown in C, staining patterns in other *Hm* races (*meriana* and *aglaope*) appeared similar. The probe used was complementary to the *cortex* isoform with the longest open reading frame (also the most common, Supplementary Information).

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### 315 Methods

316 He Cr reference

317 *Cr* is the homologue of *Yb* in *He* (Fig 1). An existing reference for this region was available

- in 3 pieces (467,734bp, 114,741bp and 161,149bp, GenBank: KC469893.1)<sup>31</sup>. We screened
- the same BAC library used previously<sup>11,31</sup> using described procedures<sup>11</sup> with probes designed

320 to the ends of the existing BAC sequences and the HmYb BAC reference sequence. Two BACs (04B01 and 10B14) were identified as spanning one of the gaps and sequenced using 321 Illumina 2x250 bp paired-end reads collected on the Illumina MiSeq. The raw reads were 322 323 screened to remove vector and E. coli bases. The first 50k read pairs were taken for each BAC and assembled individually with the Phrap<sup>32</sup> software and manually edited with 324 consed<sup>33</sup>. Contigs with discordant read pairs were manually broken and properly merged 325 using concordant read data. Gaps between contig ends were filled using an in-house 326 finishing technique where the terminal 200bp of the contig ends were extracted and queried 327 328 against the unused read data for spanning pairs, which were added using the addSolexaReads.perl script in the consed package. Finally, a single reference contig was 329 generated by identifying and merging overlapping regions of the two consensus BAC 330 331 sequences.

In order to fill the remaining gap (between positions 800,387 and 848,446) we used the overhanging ends to search the scaffolds from a preliminary *He* genome assembly of five Illumina paired end libraries with different insert sizes (250, 500, 800, 4300 and 6500bp) from two related *He demophoon* individuals. We identified two scaffolds (scf1869 and scf1510) that overlapped and spanned the gap (using 12,257bp of the first scaffold and 35,803bp of the second).

The final contig was 1,009,595bp in length of which 2,281bp were unknown (N's). The *HeCr* assembly was verified by aligning to the *HmYb* genome scaffold (HE667780) with mummer and blast. The *HeCr* contig was annotated as described previously<sup>32</sup>, with some minor modifications. Briefly this involved first generating a reference based transcriptome assembly with existing *H. erato* RNA-seq wing tissue (GenBank accession SRA060220). We used Trimmomatic<sup>34</sup> (v0.22), and FLASh<sup>35</sup> (v1.2.2) to prepare the raw sequencing reads, checking the quality with FastQC<sup>36</sup> (v0.10.0). We then used the Bowtie/TopHat/Cufflinks<sup>37–39</sup> pipeline 345 to generate transcripts for the unmasked reference sequence. We generated gene predictions with the MAKER pipeline<sup>40</sup> (v2.31). Homology and synteny in gene content with the Hm Yb346 reference were identified by aligning the Hm coding sequences to the He reference with 347 348 BLAST. Homologous genes were present in the same order and orientation in He and Hm (Fig 2B,C). Annotations were manually adjusted if genes had clearly been merged or split in 349 comparison to *H. melpomene* (which has been extensively manually curated<sup>12</sup>). In addition 350 *He cortex* was manually curated from the RNA-seq data and using *Exonerate*<sup>41</sup> alignments of 351 the *H. melpomene* protein and mRNA transcripts, including the 5' UTRs. 352

### 353 Genotype-by-phenotype association analyses

Information on the individuals used and ENA accessions for sequence data are given in 354 Supplementary Table 1. We used shotgun Illumina sequence reads from 45 He individuals 355 from 7 races that were generated as part of a previous study<sup>31</sup> (Supplementary Information). 356 Reads were aligned to an He reference containing the Cr contig and other sequenced He 357  $BACs^{11,31}$  with  $BWA^{42}$ , which has previously been found to work better than  $Stampy^{43}$ 358 (which was used for the alignments in the other species) with an incomplete reference 359 sequence<sup>31</sup>. The parameters used were as follows: Maximum edit distance (n), 8; maximum 360 361 number of gap opens (o), 2; maximum number of gap extensions (e), 3; seed (l), 35; maximum edit distance in seed (k), 2. We then used Picard tools to remove PCR and optical 362 duplicate sequence reads and GATK<sup>44</sup> to re-align indels and call SNPs using all individuals 363 as a single population. Expected heterozygosity was set to 0.2 in GATK. 132,397 SNPs were 364 present across Cr. A further 52,698 SNPs not linked to colour pattern loci were used to 365 366 establish background association levels.

For the Hm / Hn clade we used previously published sequence data from 19 individuals from enrichment sequencing targeting of the *Yb* region, the unlinked *HmB/D* region that controls the presence/absence of red colour pattern elements, and ~1.8Mb of non-colour pattern
genomic regions<sup>45</sup>, as well as 9 whole genome shotgun sequenced individuals<sup>18,46</sup>. We added
targeted sequencing and shotgun whole genome sequencing of an additional 47 individuals
(Supplementary Information). Alignments were performed using Stampy<sup>43</sup> with default
parameters except for substitution rate which was set to 0.01. We again removed duplicates
and used GATK to re-align indels and call SNPs with expected heterozygosity set to 0.1.

The analysis of the *Hm/timareta*/silvaniform included 49 individuals, which were aligned to v1.1 of the *Hm* reference genome with the scaffolds containing *Yb* and *HmB/D* swapped with reference BAC sequences<sup>18</sup>, which contained fewer gaps of unknown sequence than the genome scaffolds. 232,631 SNPs were present in the *Yb* region and a further 370,079 SNPs were used to establish background association levels.

The *Hn* analysis included 26 individuals aligned to unaltered v1.1 of the *Hm* reference genome, because the genome scaffold containing *Yb* is longer than the BAC reference making it easier to compare the inverted and non-inverted regions present in this species. We tested for associations at 262,137 SNPs on the *Yb* scaffold with the *Hn bicoloratus* morph, which had a sample of 5 individuals.

We measured associations between genotype and phenotype using a score test (qtscore) in the 385 GenABEL package in R<sup>47</sup>. This was corrected for background population structure using a 386 test specific inflation factor,  $\lambda$ , calculated from the SNPs unlinked to the major colour pattern 387 388 controlling loci (described above), as the colour pattern loci are known to have different population structure to the rest of the genome<sup>14,15,18</sup>. We used a custom perl script to convert 389 GATK vcf files to Illumina SNP format for input to genABEL<sup>47</sup>. genABEL does not accept 390 multiallelic sites, so the script also converted the genotype of any individuals for which a 391 third (or fourth) allele was present to a missing genotype (with these defined as the lowest 392

frequency alleles). Custom R scripts were used to identify sites showing perfect associations
with calls for >75% of individuals.

### 395 Microarray Gene Expression Analyses

We designed a Roche NimbleGen microarray (12x135K format) with probes for all annotated *Hm* genes<sup>18</sup> and tiling the central portion of the *Yb* BAC sequence contig that was previously identified as showing the strongest differentiation between *Hm* races<sup>45</sup>. In addition to the *HmYb* tilling array probes there were 6,560 probes tiling *HmAc* (a third unlinked colour pattern locus) and 10,716 probes tiling *HmB/D*, again distanced on average at 10bp intervals. The whole-genome gene expression array contained 107,898 probes in total.

This was interrogated with Cy3 labelled double stranded cDNA generated from total RNA 402 (with a SuperScript double-stranded cDNA synthesis kit, Invitrogen, and a one-colour DNA 403 404 labelling kit, Niblegen) from four pupal developmental stages of Hm plesseni and malleti. Pupae were from captive stocks maintained in insectary facilities in Gamboa, Panama. Tissue 405 was stored in RNA later at -80°C prior to RNA extraction. RNA was extracted using TRIzol 406 (Invitrogen) followed by purification with RNeasy (Qiagen) and DNase treated with DNA-407 free (Ambion). Quantification was performed using a Qubit 2.0 fluorometer (Invitrogen) and 408 purity and integrity assessed using a Bioanalyzer 2100 (Agilent). Samples were randomised 409 and each hybridised to a separate array. The *HmYb* probe array contained 9,979 probes 410 distanced on average at 10bp. The whole-genome expression array contained on average 9 411 probes per annotated gene in the genome  $(v1.1^{18})$  as well as any transcripts not annotated but 412 413 predicted from RNA-seq evidence.

Background corrected expression values for each probe were extracted using NimbleScan
software (version 2.3). Analyses were performed with the LIMMA package implemented in
R/Bioconductor<sup>48</sup>. The tiling array and whole-genome data sets were analysed separately.

417 Expression values were extracted and quantile-normalised, log<sub>2</sub>-transformed, quality

418 controlled and analysed for differences in expression between individuals and wing regions.

P-values were adjusted for multiple hypotheses testing using the False Discovery Rate (FDR)
 method <sup>49</sup>.

### 421 In situ hybridisations

Hn and Hm larvae were reared in a greenhouse at 25-30°C and sampled at the last instar. In 422 situ hybridizations were performed according to previously described methods<sup>25</sup> with a *cortex* 423 riboprobe synthesized from a 831-bp cDNA amplicon from *Hn*. Wing discs were incubated in 424 a standard hybridization buffer containing the probe for 20-24 h at 60°C. For secondary 425 detection of the probe, wing discs were incubated in a 1:3000 dilution of anti-digoxigenin 426 alkaline phosphatase Fab fragments and stained with BM Purple for 3-6 h at room 427 temperature. Stained wing discs were photographed with a Leica DFC420 digital camera 428 429 mounted on a Leica Z6 APO stereomicroscope.

# 430 De novo assembly of short read data in Hm and related taxa

In order to better characterise indel variation from the short-read sequence data used for the 431 genotype-by-phenotype association analysis, we performed *de novo* assemblies of a subset of 432 *Hm* individuals and related taxa with a diversity of phenotypes (Extended Data Figure 2). 433 Assemblies were performed using the *de novo* assembly function of CLCGenomics 434 Workbench v.6.0 under default parameters. The assembled contigs were then BLASTed 435 against the Yb region of the Hm melpomene genome<sup>18</sup>, using Geneious v.8.0. The contigs 436 identified by BLAST were then concatenated to generate an allele sequence for each 437 individual. Occasionally two unphased alleles were generated when two contigs were 438 matched to a given region. If more than two contigs of equal length matched then this was 439

440 considered an unresolvable repeat region and replaced with Ns. The assembled alleles were then aligned using the MAFFT alignment plugin in Geneious v.8.0. 441

#### Long-range PCR targeted sequencing of cortex in Hm aglaope and Hm amaryllis 442

We generated two long-range PCR products covering 88.8% of the 1,344bp coding region of 443 *cortex* (excluding 67bp at the 5' end and 83bp at the 3' end, further details in Supplementary 444 Information). A product spanning coding exons 5 to 9 (the final exon) was obtained from 29 445 *Hm amaryllis* individuals and 29 *Hm aglaope* individuals; a product spanning coding exons 2 446 to 5 was obtained from 32 Hm amaryllis individuals and 14 Hm aglaope. In addition, a 447 product spanning exons 4 to 6 was obtained from 6 Hm amaryllis and 5 Hm aglaope that 448 failed to amplify one or both of the larger products. Long-range PCR was performed using 449 450 Extensor long-range PCR mastermix (Thermo Scientific) following manufacturers guidelines with a 60°C annealing temperature in a 10-20µl volume. The product spanning coding exons 451 5 to 9 was obtained with primers HM25\_long\_F1 and HM25\_long\_R4 (see Supplementary 452 453 Table 2 for primer sequences); the product spanning coding exons 2 to 5 was obtained with primers HM25\_long\_F4 and HM25\_long\_R2; the product spanning exons 4 to 6 was 454 obtained with primers 25\_ex5-ex7\_r1 and 25\_ex5-ex7\_f1. Products were pooled for each 455 456 individual, including 5 additional products from the Yb locus and 7 products in the region of the *HmB/D* locus. They were then cleaned using QIAquick PCR purification kit (QIAgen) 457 before being quantified with a Qubit Fluorometer (Life Technologies) and pooled in 458 equimolar amounts for each individual, taking into account variation in the length and 459 number of PCR products included for each individual (because of some PCR failures, ie. 460 461 proportionally less DNA was included if some PCR products were absent for a given individual). 462

Products were pooled within individuals (including additional products for other genes not 463 analysed here) and then quantified and pooled in equimolar amounts for each individual 464 within each race. The pooled products for each race (Hm aglaope and amaryllis) were then 465 prepared as two separate libraries with molecular identifiers and sequenced on a single lane 466 of an Illumina GAIIx. Analysis was performed using Galaxy and the history is available at 467 https://usegalaxy.org/u/njnadeau/h/long-pcr-final. Reads were quality filtered with a 468 minimum quality of 20 required over 90% of the read, which resulted in 5% of reads being 469 discarded. Reads were then quality trimmed to remove bases with quality less than 20 from 470 471 the ends. They were then aligned to the target regions using the fosmid sequences from known races<sup>45</sup> with sequence from the Yb BAC walk<sup>12</sup> used to fill any gaps. Alignments were 472 performed with BWA v0.5.6<sup>42</sup> and converted to pileup format using Samtools v0.1.12 before 473 being filtered based on quality ( $\geq 20$ ) and coverage ( $\geq 10$ ). BWA alignment parameters were 474 475 as follows: fraction of missing alignments given 2% uniform base error rate (aln -n) 0.01; maximum number of gap opens (aln -o) 2; maximum number of gap extensions (aln -e) 12; 476 477 disallow long deletion within 12 bp towards the 3'-end (aln -d); number of first subsequences to take as seed (aln -l) 100. We then calculated coverage and minor allele frequencies for 478 each race and the difference between these using custom scripts in  $R^{50}$ . 479

### 480 Sequencing and analysis of Hm fosmid clones

481 Fosmid libraries had previously been made from single individuals of 3 *Hm* races (*rosina*,

482 *amaryllis* and *aglaope*) and several clones overlapping the Yb interval had been sequenced<sup>45</sup>.

483 We extended the sequencing of this region, particularly the region overlapping *cortex* by

- 484 sequencing an additional 4 clones from *Hm rosina* (1051\_83D21, accession KU514430;
- 485 1051\_97A3, accession KU514431; 1051\_65N6, accession KU514432; 1051\_93D23,
- accession KU514433) 2 clones from *Hm amaryllis* (1051\_13K4, accession KU514434;
- 487 1049\_8P23, accession KU514435) and 3 clones from *Hm aglaope* (1048\_80B22, accession

488 KU514437; 1049\_19P15, accession KU514436; 1048\_96A7, accession KU514438). These were sequenced on a MiSeq 2000, and assembled using the *de novo* assembly function of 489 CLCGenomcs Workbench v.6.0. The individual clones (including existing clones 1051-490 143B3, accession FP578990; 1049-27G11, accession FP700055; 1048-62H20, accession 491 FP565804) were then aligned to the BAC and genome scaffold<sup>18</sup> references using the 492 MAFFT alignment plugin of Geneious v.8.0. Regions of general sequence similarity were 493 identified and visualised using MAUVE<sup>51</sup>. We merged overlapping clones from the same 494 individual if they showed no sequence differences, indicating that they came from the same 495 496 allele. We identified transposable elements (TEs) using nBLAST with an insect TE list downloaded from Repbase Update<sup>52</sup> including known *Heliconius* specific TEs<sup>53</sup>. 497

### 498 5' RACE, RT-PCR and qRT-PCR

All tissues used for gene expression analyses were dissected from individuals from captive 499 stocks derived from wild caught individuals of various races of Hm (aglaope, amaryllis, 500 501 melpomene, rosina, plesseni, malleti) and F2 individuals from a Hm rosina (female) x Hm melpomene (male) cross. Experimental individuals were reared at 28°C-31°C. Developing 502 wings were dissected and stored in RNAlater (Ambion Life Technologies). RNA was 503 504 extracted using a QIAgen RNeasy Mini kit following the manufacturer's guidelines and treated with TURBO DNA-free DNase kit (Ambion Life Technologies) to remove remaining 505 genomic DNA. RNA quantification was performed with a Nanodrop spectrophotometer, and 506 the RNA integrity was assessed using the Bioanalyzer 2100 system (Agilent). 507

- Total RNA was thoroughly checked for DNA contamination by performing PCR for  $EF1\alpha$
- 509 (using primers ef1-a\_RT\_for and ef1-a\_RT\_rev, Table S2) with 0.5µl of RNA extract (50ng-
- 510  $1\mu g$  of RNA) in a 20 $\mu l$  reaction using a polymerase enzyme that is not functional with RNA

template (BioScript, Bioline Reagents Ltd.). If a product amplified within 45 cycles then theRNA sample was re-treated with DNase.

Single stranded cDNA was synthesised using BioScript MMLV Reverse Transcriptase
(Bioline Reagents Ltd.) with random hexamer (N6) primers and 1µg of template RNA from
each sample in a 20 µl reaction volume following the manufacturer's protocol. The resulting
cDNA samples were then diluted 1:1 with nuclease free water and stored at -80°C.

5' RACE was performed using RNA from hind-wing discs from one Hm aglaope and one 517 Hm amaryllis final instar larvae with a SMARTer RACE kit from Clonetech (California, 518 USA). The gene specific primer used for the first round of amplification was anchored in 519 exon 4 (fzl\_raceex5\_R1, Supplementary Table 2). Secondary PCR of these products was then 520 521 performed using a primer in exon 2 (HM25\_long\_F2, Supplementary Table 2) and the nested universal primer A. Other isoforms were detected by RT-PCR using primers within exons 2 522 and 9 (gene25\_for\_full1 and gene25\_rev\_ex3). We identified isoforms from 5' RACE and 523 524 RT-PCR products by cutting individual bands from agarose gels and if necessary by cloning products before Sanger sequencing. Cloning of products was performed using TOPO TA 525 (Invitrogen) or pGEM-T (Promega) cloning kits. Sanger sequencing was performed using 526 527 BigDye terminator v3.1 (Applied Biosystems) run on an ABI13730 capillary sequencer. Primers fzl ex1a F1 and fzl ex4 R1 were used to confirm expression of the furthest 5' 528 UTR. For isoforms that appeared to show some degree of race specificity we designed 529 isoform specific PCR primers spanning specific exon junctions (Extended Data Fig 2, 4, 530 Supplementary Table 2) and used these to either qualitatively (RT-PCR) or quantitatively 531 532 (qRT-PCR) assess differences in expression between races.

We performed qRT-PCR using SensiMix SYBR green (Bioline Reagents Ltd.) with 0.20.25µM of each primer and 1µl of the diluted product from the cDNA reactions. Reactions

535	were performed in an Opticon 2 DNA engine (MJ Research), with the following cycling
536	parameters: 95°C for 10min, 35-50 x: (95°C for 15sec, 55-60°C for 30sec, 72° for 30sec),
537	72°C for 5min. Melting curves were generated between 55°C and 90°C with readings taken
538	every 0.2°C for each of the products to check that a single product was generated. At least
539	one product from each set of primers was also run on a 1% agarose gel to check that a single
540	product of the expected size was produced and the identity of the product confirmed by direct
541	sequencing (See Supplementary Table 2 for details of primers for each gene). We used two
542	housekeeping genes (EF1 $\alpha$ and Ribosomal Protein S3A) for normalisation and all results
543	were taken as averages of triplicate PCR reactions for each sample.
544	$C_t$ values were defined as the point at which fluorescence crossed a threshold ( $R_{Ct}$ ) adjusted
545	manually to be the point at which fluorescence rose above the background level.
546	Amplification efficiencies (E) were calculated using a dilution series of clean PCR product.
547	Starting fluorescence, which is proportional to the starting template quantity, was calculated
548	as $R_0 = R_{Ct} (1+E)^{-Ct}$ . Normalized values were then obtained by dividing $R_0$ values for the
549	target loci by $R_0$ values for EF1 $\alpha$ and RPS3A. Results from both of these controls were
550	always very similar, therefore the results presented are normalized to the mean of $EF1\alpha$ and
551	RPS3A. All results were taken as averages of triplicate PCR reactions. If one of the triplicate
552	values was more than one cycle away from the mean then this replicate was excluded.
553	Similarly any individuals that were more than two standard deviations away from the mean of
554	all individuals for the target or normalization genes were excluded (these are not included in
555	the numbers of individuals reported). Statistical significance was assessed by Wilcoxon rank
556	sum tests performed in $R^{50}$ .

# 557 RNAseq analysis of Hm amaryllis/aglaope

RNA-seq data for hind-wings from three developmental stages had previously been obtained
for two individuals of each race at each stage (12 individuals in total) and used in the
annotation of the *Hm* genome<sup>18</sup> (deposited in ENA under study accessions ERP000993 and
PRJEB7951). Four samples were multiplexed on each sequencing lane with the fifth instar
larval and day 2 pupal samples sequenced on a GAIIx sequencer and the day 3 pupal wings
sequenced on a Hiseq 2000 sequencer.

Two methods were used for alignment of reads to the reference genome and inferring read 564 counts, Stampy<sup>43</sup> and RSEM (RNAseq by Expectation Maximisation)<sup>54</sup>. In addition we used 565 two different R/Bioconductor packages for estimation of differential gene expression, 566 DESeq<sup>55</sup> and BaySeq<sup>56</sup>. Read bases with quality scores < 20 were trimmed with FASTX-567 Toolkit (<u>http://hannonlab.cshl.edu/fastx\_toolkit/index.html</u>). Stampy was run with default 568 parameters except for mean insert size, which was set to 500, SD 100 and substitution rate, 569 which was set to 0.01. Alignments were filtered to exclude reads with mapping quality <30570 and sorted using Samtools<sup>57</sup>. We used the HT seq-count script in with HTseq<sup>58</sup> to infer counts 571 per gene from the BAM files. 572

RSEM<sup>54</sup> was run with default parameters to infer a transcriptome and then map RNAseq
reads against this using Bowtie<sup>37</sup> as an aligner. This was run with default parameters except
maximum number of mismatches, which was set to 3.

### 576 Annotation and alignment of fizzy family proteins

In the arthropod genomes, some fizzy family proteins were found to be poorly annotated based on alignments to other family members. In these cases annotations were improved using well annotated proteins from other species as references in the program Exonerate<sup>41</sup> and the outputs were manually curated. Specifically, the annotation of *B. mori fzr* was extended based on alignment of *D. plexippus fzr*; the annotation of *B. mori fzy* was altered 582 based on alignment of Drosophila melanogaster and D. plexippus fzy; H. melpomene fzy was identified as part of the annotated gene HMEL017486 on scaffold HE671623 (Hmel v1.1) 583 based on alignment of D. plexippus fzy; the Apis mellifera fzr annotation was altered based 584 585 on alignment of *D. melanogaster fzr*; the annotation of *Acyrthosiphon pisum fzr* was altered based on alignment of D. melanogaster fzr. No one-to-one orthologues of D. melanogaster 586 fzr2 were found in any of the other arthropod genera, suggesting that this gene is Drosophila 587 specific. Multiple sequence alignment of all the fizzy family proteins was then performed 588 using the Expresso server<sup>59</sup> within T-coffee<sup>60</sup>, and this alignment was used to generate a 589 neighbour joining tree in Geneious v8.1.7. 590

### 591 Expression of *H. melpomene cortex* in *D. melanogaster* wings

D. melanogaster Cortex is known to generate an irregular microchaete phenotype when 592 ectopically expressed in the posterior compartment of the adult fly wing<sup>24</sup>. We performed the 593 same assay using *H. melpomene cortex* in order to test if this functionality was conserved. 594 Following the methods of Swan and Schüpbach<sup>24</sup> a UAS-GAL4 construct was created using 595 the coding region for the long isoform of Hm cortex, plus a Drosophila cortex version to act 596 as positive control. The HA-tagged H. melpomene UAS-cortex expression construct was 597 generated using cDNA reverse transcribed (Revert-Aid, Thermo-Scientific) from RNA 598 extracted (Qiagen RNeasy) from pre-ommochrome pupal wing material. An HA-tagged 599 600 D.melanogaster UAS-cortex version was also constructed, following the methods of Swan and Schüpbach, (2007). Expression was driven by hsp70 promoter. Constructs were injected 601 into  $\phi$ C31-attP40 flies (#25709, Bloomington stock centre, Indiana; Cambridge University 602 603 Genetics Department, UK, fly injection service) by site directed insertion into CII via an attB site in the construct. Homozygous transgenic flies were crossed with w,y';en-GAL4;UAS-604 GFP (gift of M. Landgraf lab, Cambridge University Zoology Department) to drive 605

606	expression in the engrailed posterior domain of the wing, and adult offspring wings
607	photographed (Extended Data Fig 6B-D). Expression of the construct was confirmed by IHC
608	(standard Drosophila protocol) of final instar larval wing discs using mouse anti-HA and goat
609	anti-mouse alexa-fluor 568 secondary antibodies (Abcam), imaged by Leica SP5 confocal.
610	Successful expression of <i>Hm_Cortex</i> was confirmed by IHC against an HA tag inserted at the
611	N terminal of either protein (Extended Data Fig 6E).
612	
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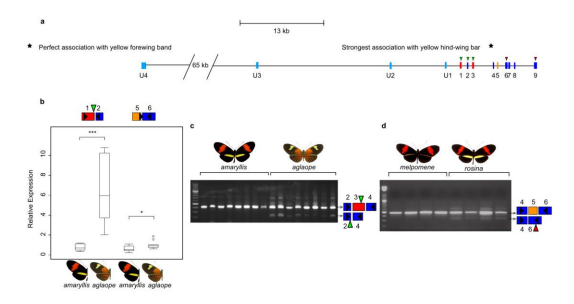
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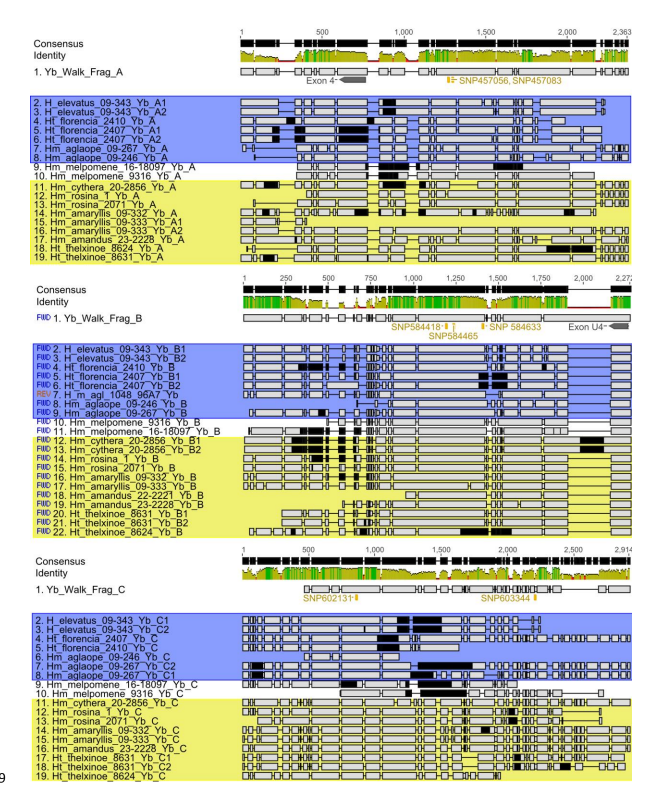
### 685 Extended Data



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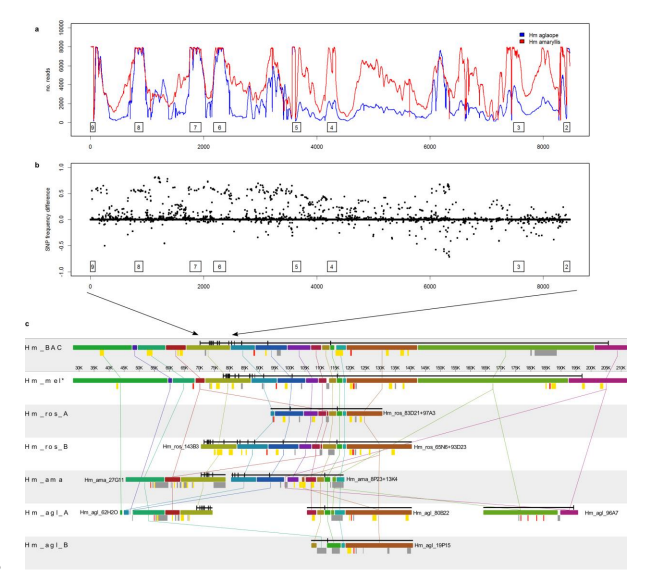
Extended Data Figure 1. A) Exons and splice variants of *cortex* in *Hm*. Orientation is
reversed with respect to figures 2 and 4, with transcription going from left to right. SNPs
showing the strongest associations with phenotype are shown with stars. B) Differential
expression of two regions of *cortex* between *Hm amaryllis* and *Hm aglaope* whole hindwings
(N=11 and N=10 respectively). Boxplots are standard (median; 75<sup>th</sup> and 25<sup>th</sup> percentiles;
maximum and minimum excluding outliers – shown as discrete points) C) Expression of a

693	<i>cortex</i> isoform lacking exon 3 is found in <i>Hm aglaope</i> but not <i>Hm amaryllis</i> hindwings. D)
694	Expression of an isoform lacking exon 5 is found in <i>Hm rosina</i> but not <i>Hm melpomene</i>
695	hindwings. Green triangles indicate predicted start codons and red triangles predicted stop
696	codons, with usage dependent on which exons are present in the isoform. Schematics of the
697	targeted exons are shown for each (q)RT-PCR product, black triangles indicate the position
698	of the primers used in the assay.





Extended Data Figure 2. Alignments of *de novo* assembled fragments containing the top
associated SNPs from *Hm* and related taxa short-read data. Identified indels do not show
stronger associations with phenotype that those seen at SNPs (as shown in Extended Data
Table 2), although some near-perfect associations are seen in fragment C. Black regions =

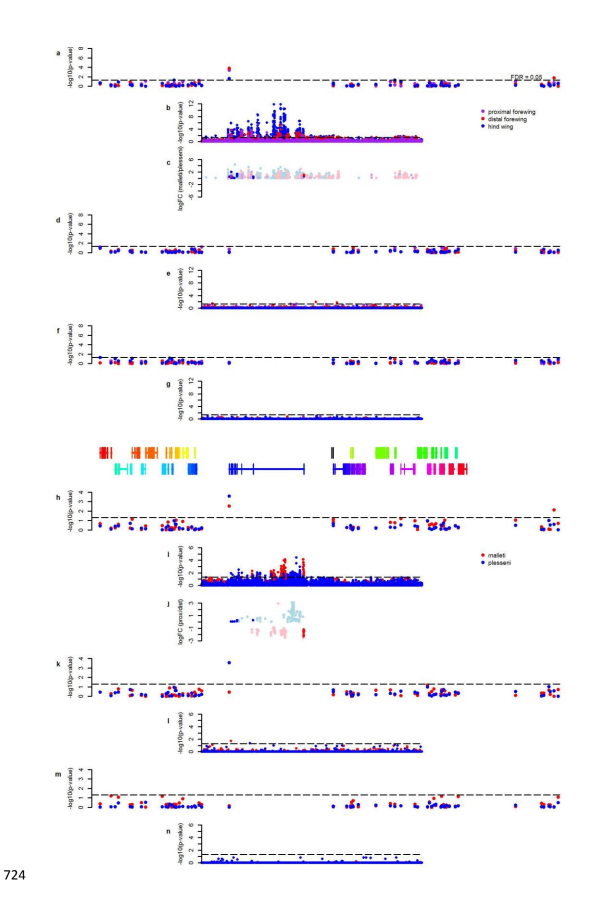


missing data; yellow box = individuals with a hindwing yellow bar; blue box = individuals
with a yellow forewing band.

Extended Data Figure 3. Sequencing of long-range PCR products and fosmids spanning *cortex*. A) Sequence read coverage from long-range PCR products across the *cortex* coding
region from 2 *Hm* races. B) Minor allele frequency difference from these reads between *Hm aglaope* and *Hm amaryllis*. Exons of *cortex* are indicated by boxes, numbered as in Extended
Data Figure 2. C) Alignments of sequenced fosmids overlapping *cortex* from 3 *Hm*individuals of difference races. No major rearrangements are observed, nor any major
differences in transposable element (TE) content between closely related races with different

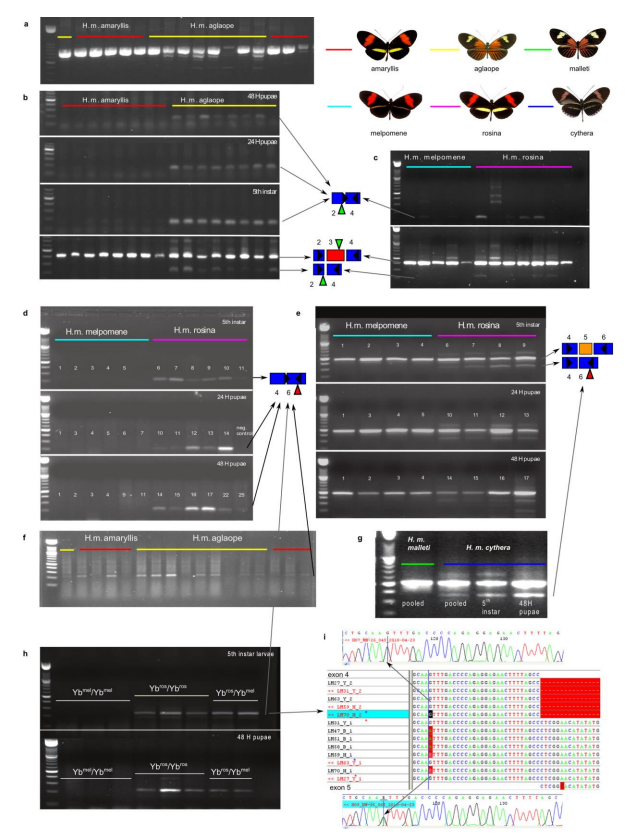
714 colour patterns (melpomene/rosina or amaryllis/aglaope). Hm amaryllis and rosina have the same phenotype, but do not share any TEs that are not present in the other races. Hm\_BAC = 715 BAC reference sequence, Hm\_mel = *melpomene* from new unpublished assembly of *Hm* 716 genome<sup>51</sup>, Hm ros = rosina (2 different alleles were sequenced from this individual), 717 Hm\_ama = *amaryllis* (2 non-overlapping clones were sequenced in this individual), Hm\_agla 718 = *aglaope* (4 clones were sequenced in this individual 2 of which represent alternative 719 alleles). Alignments were performed with Mauve: coloured bars represent homologous 720 genomic regions. *cortex* is annotated in black above each clone. Variable TEs are shown as 721 coloured bars below each clone: red = Metulj-like non-LTR, yellow = Helitron-like DNA, 722

723 grey = other.



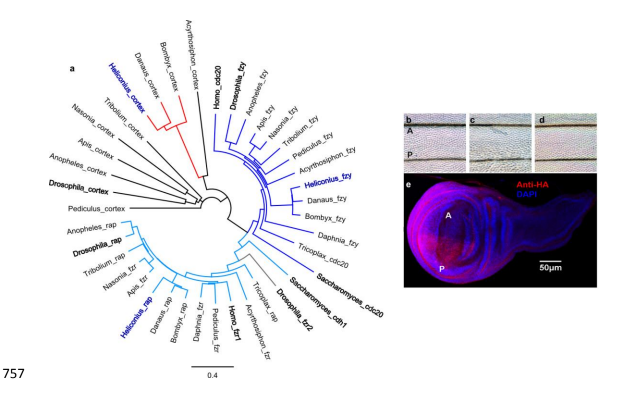
Extended Data Figure 4. Expression array results for additional stages, related to Figure 4. AG: comparisons between races (*H. m. plesseni* and *H. m. malleti*) for 3 wing regions. H-N:

- 727 comparisons between proximal and distal forewing regions for each race. Significance values
- 728  $(-\log 10(p-value))$  are shown separately for genes in the *HmYb* region from the gene array
- 729 (A,D,F,H,K,M) and for the *HmYb* tiling array (B,E,G,I,L,N) for day 1 (A,B,H,I), day 5
- 730 (D,E,K,L) and day 7 (F,G,M,N) after pupation. The level of expression difference (log fold
- change) for tiling probes showing significant differences ( $p \le 0.05$ ) is shown for day 1 (C and
- J) with probes in known *cortex* exons shown in dark colours and probes elsewhere shown as
- 733 pale colours.



Extended Data Figure 5. Alternative splicing of *cortex*. A) Amplification of the whole *cortex*coding region, showing the diversity of isoforms and variation between individuals. B)

737 Differences in splicing of exon 3 between H. m. aglaope and H. m. amaryllis. Products amplified with a primer spanning the exon 2/4 junction at 3 developmental stages. The lower 738 panel shows verification of this assay by amplification between exons 2 and 4 for the same 739 740 final instar larval samples (replicated in Extended Data Figure 2C) C) Lack of consistent differences between H. m. melpomene and H. m. rosina in splicing of exon 3. Top panel 741 shows products amplified with a primer spanning the exon 2/4 junction, lower panel is the 742 same samples amplified between exons 2 and 4. D) Differences in splicing of exon 5 between 743 H. m. melpomene and H. m. rosina. Products amplified with a primer spanning the exon 4/6 744 745 junction at 3 developmental stages. E) Subset of samples from D amplified with primers between exons 4 and 6 for verification (middle, 24hr pupae samples are replicated in 746 747 Extended Data Figure 2D). F) Lack of consistent differences between H. m. aglaope and H. 748 *m. amaryllis* in splicing of exon 5. Products amplified with a primer spanning the exon 4/6junction. G) H. m. cythera also expresses the isoform lacking exon 5, while a pool of 6 H. m. 749 *malleti* individuals do not. H) Expression of the isoform lacking exon 5 from an F2 H. m. 750 751 melpomene x H. m. rosina cross. Individuals homozygous or heterozygous for the H. m. rosina HmYb allele express the isoform while those homozygous for the H. m. melpomene 752 *HmYb* allele do not. I) Allele specific expression of isoforms with and without exon 5. 753 Heterozygous individuals (indicated with blue and red stars) express only the H. m. rosina 754 allele in the isoform lacking exon 5 (G at highlighted position), while they express both 755 756 alleles in the isoform containing exon 5 (G/A at this position).



758 Extended Data Figure 6. Phylogeny of fizzy family proteins and effects of expressing *cortex* 759 in the Drosophila wing. A) Neighbour joining phylogeny of Fizzy family proteins including functionally characterised proteins (in bold) from Saccharomyces cerevisiae, Homo sapiens 760 761 and Drosophila melanogaster as well as copies from the basal metazoan Trichoplax adhaerens and a range of annotated arthropod genomes (Daphnia pulex, Acyrthosiphon 762 pisum, Pediculus humanus, Apis mellifica, Nasonia vitripennis, Anopheles gambiae, 763 Tribolium castaneum) including the lepidoptera H. melpomene (in blue), Danaus plexippus 764 and Bombyx mori. Branch colours: dark blue, CDC20/fzy; light blue, CDH1/fzr/rap; red, 765 766 lepidoptran cortex. B-E) Ectopic expression of *cortex* in *Drosophila melanogaster*. Drosophila cortex produces an irregular microchaete phenotype when expressed in the 767 posterior compartment of the fly wing (C) whereas Heliconius cortex does not (D), when 768 769 compared to no expression (B). A, anterior; P, posterior. Successful Heliconius cortex expression was confirmed by anti-HA IHC in the last instar Drosophila larva wing imaginal 770 disc (D, red), with DAPI staining in blue. 771

# Extended Data Table 1. Genes in the *Yb* region and evidence for wing patterning control in

### 773 Heliconius

			Helio	conius I	melpor	nene						Н. е	rato		Hn	
Hm gene ID	He gene ID	Putative gene name	Yb <sup>l</sup>	Sb	A <sup>Yb</sup>	A <sup>N</sup>	E1	$E^{gw}$	E <sup>gr</sup>	Etw	E <sup>tr</sup>	Cr	Apet	Afav	P	Abic
HM00002	HERA000036	Acylpeptide hydrolase			2							x				
HM00003	HERA000037	HM00003										x				
HM00004	HERA000038	Trehalase-1B	x									х				
HM00006	HERA000038.1	Trehalase-1A	x									x				
HM00007	HERA000039	B9 protein	x									x				
HM00008	HERA000040	HM00008	x		2							x				
HM00010	HERA000041	WD40 repeat domain 85	x									x				
HM00012	HERA000042	CG2519	x					x				x				
HM00013	HERA000045	Unkempt	x									x				
HM00014	HERA000046	Histone H3	х									х				
HM00015	HERA000047	HM00015	x									x				
HM00016	HERA000048	HM00016	x									x				
HM00017	HERA000049	RecQ Helicase	x									x				
HM00018	HERA000051	HM00018	x									x				
HM00019	HERA000052	BmSuc2	x					x				x				
HM00020	HERA000053	CG5796	x									x				
HM00021	HERA000054	HM00021	x									х				
HM00022	HERA000055	Enoyl-CoA hydratase	x									x				
HM00023	HERA000056	ATP binding protein	x									x				
HM00024	HERA000057	HM00024	x									x				
HM00025	HERA000059	cortex	x	x	56	74	x	x	x	603	1796	x	2	99	x	51
HM00026	HERA000077	Poly(A)-specific ribonuclease (parn)		x	10					1	34	x			x	
HM00027	HERA000079	CG31320		x								x			x	
HM00028	HERA000080	ARP-like		x								x			x	
HM00029	HERA000081	CG4692		x								x			x	
HM00030	HERA000082	Proteasome 26S non ATPase subunit 4		x								x			x	
HM00031	HERA000083	HM00031		x					x			x			x	
HM00032	HERA000084	Zinc phosphodiesterase		x							1	x			x	
HM00033	HERA000085	Serine/threonine-protein kinase (LMTK1)		x							8	x			x	
HM00034	HERA000086	WD repeat domain 13 (Wdr13)			1	4					5	x			x	
HM00035	HERA000087	Domeless			1	2					0	x			x	
HM00036	HERA000061	WAS protein family homologue 1			5	36					37	x			x	
HM00038	HERA000062	Lethal (2) k05819 CG3054			U	00					0.	x	2		x	
HM00039	HERA000064	Mitogen-activated protein kinase (MAPKK)										x	-		x	
HM00040	HERA000064.1	DNA excision repair protein ERCC-6										x			х	
HM00040	HERA000065	Penguin										x			x	
HM00041	HERA000066	Thymidylate kinase										x			x	
HM00042	HERA000067	Caspase-activated DNase														
HM00043	HERA000068	Regulator of ribosome biosynthesis										x x			x x	
HM00045 HM00046	HERA000069 HERA000070	CG12659 CG33505										×			x	
HM00046 HM00047	HERA000070											x			x	
		Sr protein										x			x	
HM00048	HERA000073	HM00048										x			x	
HM00049	HERA000073.1	HM00049										x			x	
HM00050	HERA000074	Shuttle craft										x			x	
HM00051	HERA000075	HM00051										x			×	
HM00052	HERA000076	HM00052					×					x			х	

Yb<sup>I</sup>, within the previously mapped *Yb* interval<sup>12</sup>. Sb<sup>I</sup>, within the previously mapped *Sb*interval<sup>12</sup>. *Sb* controls a white/yellow hindwing margin and is not investigated in this study.
The *N* locus has not been fine-mapped previously. A<sup>Yb</sup>, number of above background SNPs

778	associated with the hindwing yellow bar in this study. A <sup>N</sup> , number of above background
779	SNPs associated with the forewing yellow band in this study. $E^1$ , detected as differentially
780	expressed between Hm aglaope and amaryllis from RNAseq data in this study
781	(Supplementary Information). E <sup>gw</sup> , detected as differentially expressed between forewing
782	regions in the gene array in this study. $E^{gr}$ , detected as differentially expressed between $Hm$
783	plesseni and malleti in in the gene array in this study. E <sup>tw</sup> , numbers of probes showing
784	differential expression between forewing regions in the tilling array in this study. E <sup>tr</sup> ,
785	numbers of probes showing differential expression between Hm plesseni and malleti in in the
786	tiling array in this study. $Cr^{I}$ , within the previously mapped <i>HeCr</i> interval <sup>11</sup> . A <sup><i>pet</i></sup> , number of
787	SNPs fixed for the alternative allele in <i>He demophoon</i> . A <sup>fav</sup> , number of SNPs fixed for the
788	alternative allele in <i>He favorinus</i> . P <sup>I</sup> , within the previously mapped P interval <sup>13</sup> . A <sup>bic</sup> , number
789	of above background SNPs associated with the <i>Hn bicoloratus</i> phenotype in this study.

# 791 Extended Data Table 2. Locations of fixed/above background SNPs and differentially

# 792 expressed (DE) tiling array probes

			<i>cortex</i> coding exons	<i>cortex</i> UTR exons	<i>cortex</i> introns (nonTE)	<i>cortex</i> flanking intergenic (nonTE)	TEs	Other genes (exons or introns)	Other intergenic	Total
	erato fav	vorinus fixed	2	0	96	8	2	0	0	108
	<i>erato de</i> fixed	emophoon	0	0	1	5	1	2	6	15
	above b	<i>bicoloratus</i> ackground	1	3	47	16	0	2	0	69
Posi prob		E tiling array	Known <i>cortex</i> coding exons	<i>cortex</i> UTR exons	<i>cortex</i> introns (nonTE)	miRNAs	TEs	Other gene exons	Other introns/ intergenic	Total
	esseni	Forewing proximal	8	7	323	0	13	1	7	359
ç	malleti vs plesseni	Forewing distal	12	2	327	0	8	0	8	357
Day3	E	Hindwing	5	14	378	0	9	1	6	413
	Proximal vs distal	malleti	0	1	68 222	0	0 10	0	12	81 242
		plesseni	2	4	222	0	10	0	4	242
	lesseni	Forewing proximal	1	0	22	0	3	0	7	33
Day1	malleti vs plesseni	Forewing distal	2	3	116	1	9	5	112	248
	E	Hindwing	9	10	500	1	20	2	80	622
	Proximal vs distal	malleti	0	12	95	0	1	0	0	108
	Ŀ,	plesseni	3	3	81	0	99	0	0	186

<sup>793</sup> 

Extended Data Table 3. SNPs showing the strongest phenotypic associations in the *H*.

796 *melpomene/timareta*/silvaniform comparison.

<sup>794</sup> 

Species	Race	Sample Code		SNP pos 457083† (p=6.07E- 10)	SNP pos 439063* (p=1.72E- 09)	SNP pos 602131‡ (p=2.42E- 09)	SNP pos 457056† (p=2.42E- 09)	FW band	SNP pos 584465§ (p=1.37E- 07)	SNP pos 584418§ (p=1.41E- 07)	SNP pos 584633§ (p=2.10E- 07)	SNP pos 603344‡ (p=2.19E 07)
H. melpomene	aglaope	09-246	0	A/A	A/G	A/A	C/C	1	T/T	A/A	NA	T/T
H. melpomene	aglaope	09-267	0	A/A	G/G	A/A	C/C	1	T/T	A/A	C/C	T/T
H. melpomene	aglaope	09-268	0	A/A	G/G	A/A	C/C	1	T/T	A/A	C/C	T/T
H. melpomene	aglaope	09-357	0	A/A	G/G	G/A	C/C	1	T/T	NA	C/C	T/T
H. melpomene	aglaope	aglaope.1	0	A/A	G/G	NA	C/C	1	C/T	T/A	T/C	T/T
H. melpomene	amandus	2221	1	A/A	NA	G/G	C/C	0	C/C	T/T	T/T	A/A
H. melpomene	amandus	2228	1	A/A	NA	G/G	C/C	0	C/T	T/A	T/C	A/A
H. melpomene	amaryllis	09-332	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. melpomene	amaryllis	09-333	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. melpomene	amaryllis	09-075	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. melpomene	amaryllis	09-079	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
2443 TP 52			1	T/T		G/G	T/T	0	C/C	T/T	T/T	A/A
H. melpomene	amaryllis	amaryllis.			A/A							
H. melpomene	bellula	228	1	T/T	NA	G/G	T/T	0	C/C	T/T	T/T	NA
H. melpomene	bellula	231	1	Т/Т	NA	G/A	Т/Т	0	C/T	T/A	T/C	NA
H. melpomene	cythera	2856	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. melpomene	cythera	2857	1	NA	NA	NA	NA	0	NA	NA	NA	NA
H. melpomene	malleti	17162	0	A/A	G/G	A/A	C/C	1	T/T	A/A	C/C	T/T
H. melpomene	melpomen	e18038	0	A/A	G/G	G/G	C/C	0	C/C	T/T	T/T	A/A
H. melpomene	melpomen	e18097	0	NA	G/G	NA	C/C	0	C/C	T/T	T/T	NA
H. melpomene	melpomen	em0.06	0	A/A	G/G	G/G	C/C	0	C/C	T/T	T/T	A/A
H. melpomene	melpomen	egen_ref	0	A/A	G/G	NA	C/C	0	C/C	T/T	T/T	A/A
H. melpomene	melpomen	e13435	0	A/A	G/G	A/A	C/C	0	C/C	T/T	T/T	A/A
H. melpomene	melpomen	e9315	0	A/A	G/G	A/A	C/C	0	C/C	T/T	T/T	A/A
H. melpomene	melpomen	e9316	0	A/A	G/G	A/A	C/C	0	C/C	T/T	T/T	A/A
H. melpomene	melpomen	e9317	0	A/A	G/G	A/A	C/C	0	C/C	T/T	T/T	A/A
H. melpomene	plesseni	9156	0	A/A	G/G	A/A	C/C	0	C/C	T/T	T/T	NA
H. melpomene	plesseni	16293	0	A/A	G/G	A/A	C/C	0	C/C	T/T	T/T	NA
H. melpomene	rosina	rosina.1	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. melpomene	rosina	2071	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. melpomene	rosina	531	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. melpomene	rosina	533	1	T/T	NA	G/G	T/T	0	C/C	T/T	T/T	NA
H. melpomene	rosina	546	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
(e.)	thelxiopeia		0	A/A	G/G	A/A	C/C	1	C/C	T/A	T/C	T/T
H. melpomene	vulcanus	14632	1	T/T	A/A	G/G	T/T	0	C/C	T/T	т/т	NA
H. melpomene			1									
H. melpomene	vulcanus	519	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. timareta	florencia	2403	0	A/A	G/G	A/A	C/C	1	T/T	A/A	C/C	T/T
H. timareta	florencia	2406	0	A/A	A/G	A/A	C/C	1	Т/Т	A/A	C/C	T/T
H. timareta	florencia	2407	0	A/A	A/G	A/A	C/C	1	T/T	A/A	C/C	T/T
H. timareta	florencia	2410	0	A/A	G/G	A/A	C/C	1	T/T	A/A	C/C	T/T
H. timareta	timareta	8533	0	A/A	G/G	A/A	C/C	1	C/T	T/A	T/C	T/T
H. timareta	timareta	9184	0	A/A	G/G	A/A	C/C	1	T/T	A/A	C/C	T/T
H. timareta	timareta	8520	0	A/A	G/G	A/A	C/C	1	T/T	A/A	C/C	T/T
H. timareta	timareta	8523	0	A/A	G/G	A/A	C/C	1	T/T	A/A	C/C	T/T
H. timareta	thelxinoe	09-312	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. timareta	thelxinoe	8624	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. timareta	thelxinoe	8628	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. timareta	thelxinoe	8631	1	T/T	A/A	G/G	T/T	0	C/C	T/T	T/T	A/A
H. elevatus		09-343	0	A/T	G/G	A/A	T/T	1	C/T	NA	C/C	T/T
H. pardalinus		09-326	0	A/A	A/A	A/A	NA	0	C/C	T/T	T/T	NA

\*downstream of *cortex*, †between exons 3 and 4 of *cortex*, ‡upstream of *cortex*, §between
exons U4 and U3 of *cortex*. None of these SNPs are within known TEs. Colours show
phenotypic associations: yellow = yellow hindwing bar; pink = no yellow hindwing bar;

- green = yellow forewing band; blue = no yellow forewing band; grey = allele does not match
- 802 expected pattern.

805

# 804 Extended Data Table 4. Transposable Elements (TEs) found within the *Yb* region.

BAC	Occurr			aal	_ Ne	TC name	Cupatomilu		Time
AC	mel	ros	ama	agl	No.	TE name	Superfamily		Туре
					1	BEL-1	BEL		LTR retrotransposon
					1	CR1-2	Jockey	LINE	Non-LTR retrotransposon
	1				1	Daphne-1	Jockey	LINE	Non-LTR retrotransposon
					1	Daphne-6	Jockey	LINE	Non-LTR retrotransposon
1					1	DNA-like-8			DNA transposon
					1	Helitron-like-14	Helitron_A		DNA transposon
	1	2			4	Helitron-like-12	Helitron_A		DNA transposon
	2				5	Helitron-like-12b	Helitron_A		DNA transposon
	1	1	1	1	7	Helitron-like-4a	Helitron_A		DNA transposon
						Helitron-like-4b	Helitron_A		DNA transposon
						Helitron-N2	Helitron_A		DNA transposon
					3	Helitron-like-7	Helitron_A		DNA transposon
	3	3	1	2	16	Helitron-like-6a	Helitron_B		DNA transposon
						Helitron-like-6b	Helitron_B		DNA transposon
						Helitron-like-11	Helitron_B		DNA transposon
2	2	1		1	11	Helitron-like-15	Helitron_B		DNA transposon
	5	3	1		18	Helitron-like-5	Helitron_B		DNA transposon
		1			2	Hmel Unknown 50			
	1		1		2	Hmel Unknown 174a/b			
	1				1	Hmel Unknown 187b			
	12		1	1	2	Hmel Unknown 230			
					1	Hmel Unknown 234a			
					1	Hmel Unknown 236a			
	1				1	Jockev-4	Jockey	LINE	Non-LTR retrotransposon
	1				1	LTR-3 gypsy	Gypsy		LTR retrotransposon
				1	1	Mariner-4	Mariner/Tc1		DNA transposon
R				3	29	Metuli-0	Metuli	SINE	Non-LTR retrotransposon
				5	20	Metulj-0	Metulj	SINE	Non-LTR retrotransposon
						Metulj-2	Metulj	SINE	Non-LTR retrotransposon
						Metuli-3	Metulj	SINE	Non-LTR retrotransposon
						Metulj-4		SINE	Non-LTR retrotransposon
						Metulj-4 Metulj-5	Metuli	SINE	Non-LTR retrotransposon
						Metulj-5 Metulj-6	Metulj Metulj	SINE	Non-LTR retrotransposon
						Metulj-6 Metulj-7	Metulj	SINE	Non-LTR retrotransposon
								SINE	
						nTc3-4	Mariner/Tc1	SINE	DNA transposon
	4				0	SINE-1	SINE Marinar/Tat	SINE	Non-LTR retrotransposon
	1				2	nMar-3	Mariner/Tc1		DNA transposon
			8		1	nMar-16	Mariner/Tc1		DNA transposon
			1		1	nMar-12/20	Mariner/Tc1		DNA transposon
				1	1	nPIF-3	PIF/Harbinger		DNA transposon
					1	nTc3-2	Mariner/Tc1		DNA transposon
					2	nTc3-3	Mariner/Tc1		DNA transposon
	1				2	R4-1	R2	LINE	Non-LTR retrotransposon
			1	1	6	Rep-1	REP	LINE	Non-LTR retrotransposon
		1		1	4	RTE-3	RTE	LINE	Non-LTR retrotransposon
				1	2	RTE-11	RTE	LINE	Non-LTR retrotransposon
	1				3	Zenon-1	Jockey	LINE	Non-LTR retrotransposon
			1		1	Zenon-3	Jockey	LINE	Non-LTR retrotransposon