Deep Convergence, Shared Ancestry, and Evolutionary Novelty in the Genetic Architecture of Heliconius Mimicry

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ABSTRACT Convergent evolution can occur through different genetic mechanisms in different species. It is now clear that convergence at the genetic level is also widespread, and can be caused by either (i) parallel genetic evolution, where independently evolved convergent mutations arise in different populations or species, or (ii) collateral evolution in which shared ancestry results from either ancestral polymorphism or introgression among taxa. The adaptive radiation of Heliconius butterflies shows color pattern variation within species, as well as mimetic convergence between species. Using comparisons from across multiple hybrid zones, we use signals of shared ancestry to identify and refine multiple putative regulatory elements in Heliconius melpomene and its comimics, Heliconius elevatus and Heliconius besckei, around three known major color patterning genes: optix, WntA, and cortex. While we find that convergence between H. melpomene and H. elevatus is caused by a complex history of collateral evolution via introgression in the Amazon, convergence between these species in the Guianas appears to have evolved independently. Thus, we find adaptive convergent genetic evolution to be a key driver of regulatory changes that lead to rapid phenotypic changes. Furthermore, we uncover evidence of parallel genetic evolution at some loci around optix and WntA in H. melpomene and its distant comimic Heliconius erato. Ultimately, we show that all three of convergence, conservation, and novelty underlie the modular architecture of Heliconius color pattern mimicry.

KEYWORDS adaptation; cis-regulation; collateral evolution; genetic architecture

CONVERGENT evolution is a natural experiment in repeated evolution of similar traits, and offers unique insights into the evolutionary process (Blount et al. 2018). It is widespread across the tree of life, critical to the composition of ecosystems (Sage et al. 2012) (e.g., the repeated colonization of land/water/air by different taxonomic groups), and underpins the ability of organisms to exploit novel environments (e.g., the repeated evolution of drug/insecticide/drought resistance (Farhat et al. 2013). The genetic changes causing convergence can be categorized as (i) divergent genetic mechanisms, (ii) parallel genetic evolution, or (iii) collateral evolution (Stern 2013). With divergent genetic mechanisms, different loci cause the same phenotype in different lineages. In parallel genetic evolution, different alleles at the same locus cause trait convergence (this includes cases where the same mutation has arisen multiple times) (Tishkoff et al. 2007), whereas in collateral evolution, convergence results from the sharing of alleles that are identical by descent, either because the alleles were present in an ancestral population (Jones et al. 2012), or from the introgression of alleles from one species/taxon to another (Huerta-Sánchez et al. 2014).
Cis-regulatory evolution has been implicated in several examples of convergent evolution in vertebrates (Booker et al. 2016; Partha et al. 2017; Tollis et al. 2018; Feigin et al. 2019), suggesting that trait evolution proceeding via cis-regulatory changes to conserved regulatory pathways may be recurrent and predictable. Cis-regulatory evolution is a powerful mechanism that can result in rapid developmental and physiological changes (Wittkopp and Kalay 2012). This is because multiple enhancers at the same gene can each control the gene’s expression in different cell types or developmental times. In these cases, the modular architecture can isolate the effects of a mutation to a single trait (Feigin et al. 2019), circumventing the pleiotropic effects that might constrain adaptive evolution in the protein-coding sequence. Notable examples in which modular enhancers drive convergent evolution include the gain of melanin wing spots in Drosophila elegans and Drosophila tristis through enhancers of the gene yellow (Prud’homme et al. 2006), coat coloration phenotypes via regulation of the gene Agouti in Peromyscus mice (Steiner et al. 2007; Linnen et al. 2009, 2013), loss of Drosophila larval trichomes through mutations to regulatory regions of ovo/svb (Frankel et al. 2012), and pelvic reduction in sticklebacks due to enhancer deletions at the gene pitx1 (Chan et al. 2010). A modular cis-regulatory architecture has also been proposed as a flexible toolkit controlling wing color patterning in Heliconius butterflies (Wallbank et al. 2016; Van Belleghem et al. 2017).

Müllerian mimicry is ubiquitous among neotropical Heliconius butterflies, with multiple species evolving convergent, bright, aposematic wing color patterns. At the same time, in several species such as Heliconius erato and Heliconius melpomene, phenotypic divergence within species is also present in the form of geographic color pattern races (Figure 1), with color pattern loci easily identifiable in population genomic studies across hybrid zones as clear islands of divergence in the genome (Baxter et al. 2010; Counterman et al. 2010; Nadeau et al. 2013, 2014). Owing to the presence of repeated color pattern phenotypes, Heliconius butterflies are an excellent system for studying the genetic basis of convergent evolution (Baxter et al. 2008; Merrill et al. 2015). The small number of mimicry genes controlling the majority of color pattern elements, have been identified across Heliconius species using a combination of QTL mapping, genome-wide association studies across color pattern hybrid zones, and gene expression studies. Across multiple species, parallel genetic evolution at the genes optix, cortex, and WntA is known to control red-orange pattern elements (Baxter et al. 2008; Reed et al. 2011; Martin et al. 2014; Huber et al. 2015; Lewis et al. 2019), white and yellow pattern elements (Nadeau et al. 2016), and melanic patterning (Martin et al. 2012; Gallant et al. 2014; Mazo-Vargas et al. 2017; Moest et al. 2019; Morris et al. 2019; Van Belleghem et al. 2020), respectively.

Multiple putative regulatory elements have now been identified in H. erato around all three major wing patterning genes (optix, cortex, and WntA) using comparisons between phenotypically distinct races across multiple hybrid zones (Van Belleghem et al. 2017). In the H. melpomene clade the picture is less complete. ABBA-BABA comparisons and changes in phylogenetic topologies have shown that mimetic resemblance between some races of several H. melpomene-silvaniform clade species (H. melpomene, H. elevatus, H. timareta, and H. beskei; Figure 1B) are the result of collateral evolution, via the introgression of color pattern alleles at optix and cortex among the species (Dasmahapatra et al. 2012; Pardo-Díaz et al. 2012; Zhang et al. 2016). Association mapping across a number of H. melpomene and H. timareta taxa in conjunction with recombination breakpoint analysis (which included H. elevatus), was used to define both a 25 kb and an 11 kb regulatory element at optix associated with the presence and absence of the red hindwing rays and the forewing dennis phenotypes, respectively (Wallbank et al. 2016).

However, these genomic regions are still relatively large, and no regulatory element for red band has yet been found. Similarly, around cortex it has also been shown that introgression between races of H. melpomene and H. cydno has likely allowed these species to share variation in the hindwing yellow bar phenotype (Enciso-Romero et al. 2017). However, no yellow band regulatory element has been identified at cortex. Furthermore, patterns of introgression and any regulatory elements around WntA are thus far unknown in the H. melpomene-silvaniform clade.

In this study, we investigate the contributions of three genetic modes of evolution, divergent genetic mechanisms, parallel genetic evolution, and collateral evolution to explain the remarkable convergent wing color pattern phenotypes found in Heliconius butterflies. In particular, we use phylogenetic analysis across multiple hybrid zone comparisons to delimit narrow regions of the genome associated with color pattern elements due to shared ancestry (either from introgression or ancestral polymorphism). We do this by identifying genomic regions that show genotype-by-phenotype associations and particular phylogenetic histories consistent with controlling specific wing color pattern phenotypes. This allows us for the first time to look at the mechanism of convergence between Guianese H. melpomene and H. elevatus. We propose that these narrow regions are putative modular regulatory elements, with each controlling a specific wing pattern phenotype. We identify these around all three of the major wing patterning genes; optix, cortex, and WntA, in H. melpomene and its silvaniform mimics and determine the ancestral origins of each element. Finally, we investigate the homology and conservation of these regulatory elements between H. melpomene and H. erato, as well as across other Lepidoptera species.

**Materials and Methods**

**Sample collection and sequencing**

We used whole genome sequences of 53 individuals representing six species (H. melpomene, H. elevatus, H. beskei, H. pardalinus, H. ethilla, and H. hecale) and 15 races (Supplemental Materials, Table S1) from two hybrid zones in Peru.
and the Guianas, as well as two taxa from the Eastern Amazon in Brazil (H. m. nanna and H. besckei) (Figure 1). This includes data from newly sequenced samples of two H. elevatus tumatumari, two H. pardalinus butleri, and one H. elevatus butleri. For these new samples, RNA-free genomic DNA was extracted from thoracic tissue using a Qiagen DNeasy Blood and Tissue Kit. Libraries were prepared using Illumina TruSeq DNA PCR-Free Library Preparation Kits with an insert size of ~350 bp. Libraries were 100 or 125 bp pair-end sequenced to 30–40× coverage on an Illumina HiSeq 2500 instrument at the FAS Center for Systems Biology, Harvard (ENA accession number PRJEB37067).

**Variant calling**

We aligned sequences to the H. melpomene reference genome v2 (Davey et al. 2016; LepBase http://ensembl.lepbase.org) using BWA MEM (Li and Durbin 2009). We then sorted BAM files using Samtools (Li et al. 2009) and marked duplicate reads using PicardTools (http://broadinstitute.github.io/picard/). We then called genotypes in gVCF format with GATK's HaplotypeCaller with the parameters -baq CALCULATE_AS_NECESSARY, -hets 0.02 and–emitRefConfidence GVCF, -gt_mode DISCOVERY and–dontUseSoftClippedBases. Subsequently we combined GVCFS before genotyping them, using CombineGVCFs and GenotypeGVCFs respectively (Van der Auwera et al. 2013). Genotypes were then marked as missing (N) if minimum read depth was <5 or Q<20, while sites with a minor allele frequency lower than 2/53 across all samples were removed. Python scripts (available at https://github.com/simonhmartin) were used to parse variant call formats (VCFs) to prepare files for use in phylogenetic weighting analyses. SNPs with >10% missing data across taxa were removed.

**Identification of candidate regulatory modules based on shared ancestry**

Three major loci (cortex, optix, and WntA) control the main color pattern differences between the postman and dennis-rayed races of H. melpomene we examine here (Figure 1; Baxter et al. 2010). Previous studies have shown that mimicry...
between H. melpomene races and closely related silvaniform species such as H. elevatus and H. besckei is a consequence of shared alleles (Dasmahapatra et al. 2012; Zhang et al. 2016). Therefore, the signal of shared ancestry between comimics can be used to identify narrow genomic intervals that may control color pattern elements (Wallbank et al. 2016; Van Belleghem et al. 2017). We look for these patterns of shared ancestry around cortex, optix, and WntA. Where previous studies have delimited intervals (Wallbank et al. 2016), we constrain our search to these regions.

In order to identify regions that show shared ancestry among different comimetic species at these loci, we employed a descriptive phylogenetic weighting method called Topology Weighting by Iterative Sampling of Subtrees (Twisst; Martin and Van Belleghem 2017). This method provides a quantitative summary of a tree by weighting different subtree topologies according to their occurrence within the tree. Each of our Twisst comparisons used six taxa. We first used RAxML v8.2.4 (Stamatakis 2014) with model -GTRCAT to build maximum-likelihood trees for 100 SNP sliding windows (slide every 25 SNPs) across the entirety of chromosomes 10, 15, and 18 (which contain the major color pattern loci WntA, cortex, and optix, respectively). Trees were built only for windows where all samples had >30 SNPs. We used a dynamic threshold as implemented by Twisst to estimate weightings in all analyses, such that trees were sampled until the 95% binomial confidence interval around each weighting was <5%.

For each six taxon Twisst comparison there are 105 potential tree topologies, with just five topologies for each phenotype clearly indicative of shared ancestry related to phenotype (see Figures S2–S6). These trees show monophyly of the nonsister taxa with convergent pattern elements, while the other four taxa group as expected based on the species tree. A topology weighting of 0 indicates that none of the trees for that genomic window were of these five topologies, while a topology weighting of +1 indicates that all of the trees were among these five topologies. We used these six taxon Twisst comparisons to identify putative regions controlling wing pattern elements across three geographic contexts: Peruvian hybrid zone, Guianese hybrid zone, and H. besckei-H. melpomene taxa. These geographically distant comparisons allow semi-independent brid zone, and geographic contexts: Peruvian hybrid zone, Guianese hybrid zone, and H. besckei-H. melpomene taxa. The other two silvaniform species included H. ethilla, H. hecale, or H. pardalinus.

For comparisons with Guianese taxa, H. ethilla and H. hecale were used alongside Guianese H. elevatus taxa. However, for the Peruvian hybrid zone comparisons, H. ethilla and H. pardalinus butleri were used alongside H. e. pseudocupidineus. H. pardalinus butleri and H. e. pseudocupidineus are very closely related to each other across most of the genome (FST ~ 0 across 95% of the genome) (Kryvokhyzha 2014), and so including these taxa together with races of H. melpomene provides the sensitivity to differentiate shared variation that is associated with the phenotype (introgression or ancestral polymorphism) from shared variation that is unrelated to the phenotype (ancestry). If the mimetic silvaniform and H. melpomene taxa cluster together without the nonmimetic H. melpomene and silvaniform taxa, this is suggestive of shared variation associated with phenotype. Such a pattern should only occur at narrow color pattern regions due to our careful choice of taxa.

For comparisons with H. besckei, there is no nonmimetic H. melpomene race forming a hybrid zone with H. m. nanna available. Using H. m. nanna in Twisst analyses with other H. melpomene taxa (H. m. thlrixoeipia and H. m. aglaope) resulted in a noisy analysis due to divergence between the H. melpomene taxa. Therefore, we instead compared H. besckei to H. m. melpomene from the Guianese hybrid zone (alongside H. m. thlrixoeipia and H. m. aglaope).

**Identifying SNPs associated with phenotypes**

We also looked for SNPs on our three focal chromosomes that were “diagnostically fixed” between taxa in different phenotype groups (Table S8). We excluded H. e. tumatamari from the no-ray taxa grouping as it appears to have independently evolved its no-ray phenotype. We used the same SNPs as in the Twisst analysis, but with up to 20% of genotype calls across taxa allowed to be missing for each SNP. For optix phenotypes, genotype calls were required to be present in all H. melpomene meriana and H. besckei samples, as these taxa are essential in differentiating diagnostically fixed SNPs for rays—no-rays from those for dennis/no-band—no-dennis/band. At WntA we looked for SNPs “diagnostically fixed” between H. pardalinus in one group (with the silvaniform WntA phenotype) and all H. melpomene, H. elevatus, and H. besckei in the other group (with a nonsilvaniform WntA
phenotype). In this WntA analysis, genotypes had to be present in all *H. butleri* and *H. bescket* samples. As one of these groups is only made up of a single taxon (*H. pardalinus*), the results of the WntA analysis were noisier.

### Phylogenetic reconstruction at putative functional elements

At each of the putative functional elements we identified, we constructed unrooted trees using all 53 samples to determine the ancestral origin of each element. We included both variable and invariant sites within the boundaries of each element as delineated in our previous analyses using Twist and fixed differences. Bcftools was used to remove poor quality genotype calls, and mark genotypes as missing if minimum read depth was \(<5\) or \(GQ < 20\). In addition, we also constructed a phylogenetic tree to determine the overall species relationships using all sites from across chromosomes 1 and 2, which we expect to show the species phylogeny as they do not contain any of the main color loci. We used RAxML to build all trees, with the GTRCAT model and 100 maximum likelihood trees to determine all trees, with the GTRCAT model and 100 maximum likelihood trees to determine the best tree, followed by 1000 bootstrap pseudoreplicates.

### Conservation and homology

Functional elements controlling gene expression may be conserved across taxa. In order to identify whether there was sequence conservation across Lepidoptera and in particular with *H. erato* (a distant relative and mimic of *H. melpomene*) at the putative functional elements we have identified, we retrieved the genome sequences of nine additional species from Lepbase (http://ensemb.lepbase.org); *H. erato demophon*, *Junonia coenia* v1.0, *Bicyclus anynana* v1.2, *Danaus plexippus* v3, *Papilio machaon* v1.0, *Papilio polytes* v1.0, *Pieris napi* v1.1, *Amyelois transitella* v1, and *Bombix mori* GCA000151625.1. We then identified scaffolds corresponding to the WntA, *optic*, and *cortex* loci with BLAST (Altschul et al. 1990). Fine-scale sequence conservation between *H. melpomene* and *H. erato* was visualized using the Artemis Comparison Tool (Carver et al. 2005). We then calculated pairwise conservation between *H. melpomene* and each of the other species using mVISTA (Frazer et al. 2004), an mLAGAN (Brudno et al. 2003) alignment and a conservation cutoff of 70% sequence identity.

### Data availability

The authors state that all data necessary for confirming the conclusions presented in the article are represented fully within the article. All original raw sequence data files are available via the ENA (accession number PRJEB37067). Supplementary tables and figures referred to in the text are available via figshare: https://doi.org/10.25386/genetics.12911180.

### Results

To identify putative regulatory elements in the *H. melpomene*-silvaniform clade, we use Twist analyses across multiple hybrid zones to find signals of shared ancestry relating to wing pattern phenotypes and looked for diagnostically fixed SNPs between taxon groupings with these different phenotypes. We look for these patterns around loci known to control particular patterns, and where existing data have previously delimited an interval, we looked within that region. We first refine the rays and dennis loci, finding two putative loci for each of these phenotypes. Furthermore, we identify a 1.5 kb locus that we putatively delimit as the *optic* band region, as well as two loci at WntA that are associated with the full black discal forewing phenotype that *H. elevatus* shares with these *H. melpomene* races rather than the broken black of its sister species *H. pardalinus*. In contrast, our comparison using taxa from the Guianese hybrid zone shows no regions of shared derived ancestry between the broken band and no-rays forms of *H. elevatus* and *H. melpomene*, suggesting both these phenotypes evolved independently in mimetic *H. melpomene* and *H. elevatus* in the Guianas. We also identify a putative locus near the *cortex* gene associated with the yellow band phenotype. Finally, we use phylogenetic analysis to infer the ancestral origins of these putative regulatory elements across the *H. melpomene*-silvaniform clade, uncovering a complex history of introgression with different evolutionary origins for the various regulatory elements.

### Broader patterns of shared ancestry

On chromosome 18, our analyses identified only one additional \(\sim 4\) kb peak of shared ancestry (weighting of +1) between comimics that was consistent across hybrid zones and that was not located in the vicinity of *optic*. This was positioned \(\sim 30\) Mb from *optic*; at \(\sim 3,014,000\) bp on chromosome 18; and \(\sim 227,000\) bp on Hmel218005 (Figures S13 and S11) within the gene HML034250g1. As a tblastx search of this gene in the *Drosophila melanogaster* genome does not return any good hits we are unable to speculate about its function. No peaks of shared ancestry with a weighting of +1 were found outside of the vicinity of WntA on chromosome 10 (Figure S12), and no large peaks showing consistent shared ancestry between comimics across comparisons (Figures S13–S15) were found on chromosome 15 (which contains the *cortex* gene). Thus, we demonstrate the near absence of regions of consistent shared ancestry between comimics outside the proximity of WntA, *optic*, and *cortex* on chromosomes 10, 15, and 18, respectively.

### The ray loci

Inferred recombination breakpoints around shared haplotypes have previously been used to delimit a 37 kb ray region on chromosome 18, further narrowed to 25 kb using SNPs perfectly associated with the hindwing rays (Wallbank et al. 2016). Our Twist comparisons allow us to define two separable loci within this region with different evolutionary histories, a \(\sim 6\) kb region (Hmel218003 773301–779400) we call ray1 and another \(\sim 22\) kb region (Hmel218003 781427–803436) we call ray2 (Figure 2). Across both regions, we see a pattern of shared ancestry between rayed *H. elevatus* and
**H. melpomene** mimics from the Peruvian hybrid zone (Figure 2B). However, in our **H. besckei** comparison, we see shared ancestry only across ray2 (Figure 2C). Our phylogeny at ray2 (Figure 2F) also shows that both no-ray and rayed Guianese and Peruvian **H. elevatus** show shared ancestry with rayed **H. melpomene** taxa.

Our Twisst comparison using the Guianese taxa without rays does not show shared ancestry between no-ray **H. elevatus** and **H. melpomene** from the Guianas at either ray1 or ray2 (Figure 2D). This suggests that the Guianese no-ray phenotypes in **H. elevatus** and in **H. melpomene** have evolved independently and are not the result of shared ancestry, with the no-ray phenotype in **H. e. tumatumari** evolving after introgression of ray alleles. Other peaks of shared ancestry are also found in this comparison, some of these relate to dennis (Figure 3, B and C; peaks at 813370–814253 bp; 816686–819634 bp) and band (Figure 4, B and C; peak at 728308–729971) phenotypes found in both **H. m. meriana** and **H. e. tumatumari**. It is unclear what might cause the additional peaks; however, as these are not consistent across phenotypes/comparisons they do not contribute to the locations of putative functional elements.

Due to independent evolution of the no-ray phenotype, **H. e. tumatumari** was excluded from the “diagnostically fixed” SNP analysis as its inclusion in the no-ray group would wrongly remove all “diagnostically fixed” SNPs (Table S8). We find that ray1 contains 5 “diagnostically fixed” SNPs while ray2 contains 67. Our phylogenies inferred using all taxa (Figure 2, E and F) show that all rayed and no-ray

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**Figure 2** Twisst comparisons across the optix region of scaffold Hmel218003. (A) Location of optix and ray elements. Dark blue (ray1) and light blue (ray2) shading shows putative functional elements. Gray boxes show ray and dennis regulatory elements respectively as previously delimited in Wallbank et al. (2016). Diagnostic fixed SNPs between phenotypes shown with blue dots in (A). Twisst comparison (100 SNP windows sliding by 25 SNPs) using (B) Peruvian hybrid zone taxa, (C) **H. besckei** and Guianese hybrid zone **H. melpomene**, and (D) Guianese hybrid zone taxa. Black trees to left show species topology, while red trees shows groupings (taxa shaded in gray) that indicate shared ancestry between the heterospecific mimetic taxa. Weighting (black line) is the mean from all four overlapping windows for that region, light gray bars show weighting for each 100 SNP window. A weighting of +1 means 100% of trees at that genomic interval show shared ancestry between the heterospecific mimetic taxa. Mimetic phenotypes for taxa are shown by circles; red circles for **H. melpomene** clade and orange circles for silvaniform taxa. (E and F) Maximum likelihood phylogenies of the ray elements with red branches joining **H. melpomene** taxa and orange branches joining silvaniform taxa. Node bootstrap support; pink diamonds ≥ 95%, green diamonds 75–94%. Black branch (illustrative only) separates the silvaniform and **H. melpomene** clades (excluding those taxa where introgression appears to have occurred).
H. elevatus and all rayed H. melpomene group together at both loci, but that the no-ray H. besckei groups with no-ray H. melpomene only at ray2. These phylogenies also suggest the alleles in rayed taxa at these loci derive from different clades. The ray1 rayed allele appears to originate from the H. mel-
pomene clade (Figure 2E) as rayed individuals at ray1 are clustered within the melpomene samples. In contrast, the ray2 rayed allele appears to originate from the silvaniform clade, as rayed individuals at ray2 are clustered within the silvaniform samples (Figure 2F).

The dennis loci

Previously an ~11 kb dennis region had been defined (Wallbank et al. 2016). We narrow this region by including H. e. tumatumari and H. besckei in separate Twisst comparisons and comparing these to Guianese H. melpomene taxa. As H. e. tumatumari and H. besckei are separate species and allopatric to each other, we used separate comparisons so as not to introduce noise. We define our loci using “diagnostically fixed” SNPs between our two phenotype groups, and signals of shared ancestry that are both consistent across Twisst comparisons and within the previously delimited ~11 kb dennis region from Wallbank et al. (2016). On this basis, we delimit ~1 kb (Hmel218003 813370–814253) and ~3 kb (Hmel218003 816686–819634) regions that we term dennis1 and dennis2, respectively (Figure 3). We split these as a consequence of a dip in shared ancestry consistent across all our comparisons. Both regions show shared ancestry between H. elevatus and H. melpomene mimics with the dennis phenotype across both the Peruvian (H. e. pseudocupidineus and H. m. aglaope) and Guianese hybrid zones (H. e. tumatumari/bari and H. m. meriana/thelxiope) (Figure 3, B and C). Our diagnostic fixed SNP analysis found that the dennis1 region contained 20 “diagnostically fixed” SNPs with the correct pattern for dennis (or optix band as all red band taxa lack the dennis phenotype and so cannot be separated), while dennis2 contains 18 such SNPs. There were only 40 other “diagnostically fixed SNPs” across the whole of Chromosome 18, with nine of these coming from our putative optix band region as expected. However, some of these other SNPs are also clustered, 15 SNPs clustered just upstream (Hmel218003 822665–825712) but outside the dennis region defined by Wallbank et al. (2016) and therefore outside of our defined dennis region (Figure 3D), and eight SNPs were clustered at the peak at ~227,000 bp on Hmel218005. Phylogenies allowed us to define two separate phenotypic
groups: one containing all no-dennis *H. melpomene* and *H. besckei* and the other containing mimetic dennis *H. melpomene* and *H. elevatus*. This is consistent with Wallbank et al. (2016). As the dennis *H. melpomene* races are nested within the silvaniform clade, these phylogenies suggest that the dennis phenotype originated in the silvaniform clade (Figure 3, D and E) and introgressed into *H. melpomene*, while the no-dennis alleles appear to have introgressed from *H. melpomene* into *H. besckei*.

**The optix band locus**

Although our taxa do not allow us to tell shared ancestry due to band phenotypes from shared ancestry due to dennis phenotypes, we postulate that an optix band element would most likely be outside of the dennis (or ray) regions (Wallbank et al. 2016). This is based on (i) the existence of *H. timareta* races that lack red bands and dennis phenotypes (while other taxa have dennis phenotypes with rays but lack red bands) (Giraldo et al. 2008), and (ii) specimens of *H. melpomene* from the Guianese hybrid zone that have both dennis and red band phenotypes together (rather than having only one of these as seen in the main Guianese *H. melpomene* taxa) (J.M. personal observation). We hypothesize that any putative optix band element should also show shared ancestry across both Twisst comparisons. This is because the red band silvaniform *H. besckei* should show shared ancestry with red band races of *H. melpomene*, while *H. elevatus* races that have a yellow band should show shared ancestry with *H. melpomene* races that also have this phenotype. Using shared ancestry and diagnostically fixed SNPs we identify a single 1.5-kb region (*Hmel218003 728308–729971*), close to the 5' end of optix that we putatively delimit as the previously unidentified optix band region (Figure 4, A and B). This region spans the only consistent peak of shared ancestry (outside of the dennis and ray regions) across all hybrid zones and includes nine “diagnostically fixed” SNPs between mimetic red and yellow band taxa. Furthermore, the wider region around this 1.5 kb also contains windows that inconsistently show a signal of shared ancestry between mimetic red band taxa and mimetic yellow band taxa (shown as gray bars with weighting of +1 in Figure 4, B and C). However, because the red band is always found in the absence of dennis in our taxa (and vice-versa), it should be noted that we cannot rule out that this region is in fact involved in the control of the dennis phenotype, or that the dennis regions are not involved in the control of the red band phenotype. The phylogeny at this locus shows that *H. besckei* is nested within the clade containing the red band *H. melpomene* races. While *H. m. aglaope*, *H. m. meriana* and *H. m. thelxiopeia* which all lack the red fore-wing band group with the other silvaniform species (Figure 4D), suggesting an ancestral silvaniform clade origin for the allele causing loss of the red band.

**Two WntA loci**

Around WntA we identify two separate loci that we term WntA1 (*Hmel210004 1819577–1822844*) and WntA2 (*Hmel210004 1857021–1860913*), which show a signal of shared ancestry between *H. melpomene* aglaope/amaryllis...
and *H. elevatus pseudocupidinseus*, but not between *H. melpomene aglaope/amarilis* and *H. pardalinus butleri* (Figure 5B). This pattern is consistent with what we would expect in regions involved in controlling the full black discal forewing phenotype that *H. elevatus* shares with these *H. melpomene* races (and which replaces the broken black patterning found in *H. pardalinus* and other silvaniforms in this part of the wing). These two loci were the only peaks also supported by the results from our “diagnostically fixed” SNP analysis between *H. pardalinus butleri* and all taxa with a *H. melpomene* type phenotype (all *H. melpomene* and *H. elevatus* races and our *H. besckei*). In total *WntA1* and *WntA2* contained, respectively, 12 and 6 of the 251 “diagnostically fixed” SNPs on chromosome 10. Thus, *WntA1* and *WntA2* together contain 7% of these “diagnostically fixed” SNPs in just 0.04% of the length of the chromosome.

The phylogenies at *WntA1* and *WntA2* clearly show *H. elevatus* races grouping within (*WntA1*) or close to (*WntA2*) the *H. melpomene* clade. In both phylogenies, *H. besckei* was found outside the rest of the silvaniforms (excluding *H. elevatus*), rather than grouping with *H. ethilla* as it does in the species tree (Figure S16). This suggests that there may be some shared ancestry in these regions between *H. besckei* and the other *H. melpomene* races. Both phylogenies suggest that the full black discal forewing phenotype which replaces broken black in *H. e. pseudocupidinseus* originates from the *H. melpomene* clade. In contrast to our results with Peruvian taxa, our comparison using taxa from the Guianese hybrid zone (Figure S17) shows no regions of shared derived ancestry on chromosome 10 between the broken band forms of *H. elevatus* (*H. e. tumatumari* and *H. e. bari*) and *H. melpomene* in the Guianas (*H. m. meriana* and *H. m. thelxiopeia*), relative to *H. m. melpomene*, which lacks the broken band phenotype. Given previous research has shown that this phenotype is controlled by *WntA* in *H. melpomene* (Morris et al. 2019), our results here suggests that just like the no-rays phenotype, the broken band phenotype in the Guianas also evolved independently in mimetic *H. melpomene* and *H. elevatus*.

The cortex band locus

Only the yellow forewing band phenotype (and not the yellow hindwing bar) is found in both our Peruvian and Guianese hybrid zone taxa. We identified a ~3.5 kb region near cortex (*Hmel215006 1337470–1340886*) that shows shared derived ancestry between yellow band *H. e. pseudocupidinseus* and *H. m. aglaope* from the Peruvian hybrid zone (Figure 6B) and between red band *H. besckei* and *H. m. melpomene* (Figure 6C). This region also contained two of the three “diagnostically fixed” SNPs on chromosome 15, between red band *H. melpomene* and *H. besckei* and our yellow band *H. melpomene* and *H. elevatus*. These results suggest that this region may be involved in controlling the yellow band phenotype. We term this locus “cortex band.” Phylogenetic reconstruction at cortex band, shows that this region is also shared between all yellow band *H. elevatus* and *H. melpomene* taxa (Figure 6D), and that the putative yellow band allele appears to have originated in the silvaniform clade before then introgressing to...
into the *H. melpomene* taxa with yellow bands. However, *H. besckei* was not found on the branch with red band *H. melpomene*. Therefore, the signal of shared ancestry seen between *H. besckei* and red band *H. melpomene* in the Twisst comparison may not be due to shared ancestry between these two taxa in an allele that results in a lack of yellow band. Instead it may be an artifact of the shared ancestry between yellow band *H. melpomene* races and *H. elevatus*, which would make the *H. melpomene* haplotype in these races more closely related to *H. hecale* and *H. ethilla* haplotype than that in *H. besckei*. However, given that we only detect this phylogenetic signal at this region, and the location of our diagnostic fixed differences, this single peak is still our best candidate for a “cortex band” element, even if we cannot rule out other regions that we miss due to complex pleiotropic interactions that do not give such an expected phylogenetic signal of shared ancestry.

**Modular conservation across Heliconius and the Lepidoptera**

Fine-scale sequence conservation between *H. melpomene* and *H. erato* at and around *optix*, *WntA*, and cortex is shown in Figure 7. As expected, we find that these two genomes are largely colinear, with large sections of homology across the genomes. *Ray1* contains a substantial amount of conserved sequence between the *H. melpomene* and *H. erato*, as well as a 2000 bp region (Hmel18003:778000–780000; Figure S18) deeply conserved across the Lepidoptera, indicating the presence of a conserved functional element within *ray1* of both species (See Table S19 for scaffold locations in other Lepidoptera). Our analysis also finds narrow regions of homology (Table S20) between the *ray2* locus in *H. melpomene* and the *R* locus controlling the ray phenotype in *H. erato* (Van Belleghem et al. 2017). The *H. melpomene dennis1* element has very low sequence conservation with *H. erato* and no conserved sequence with other Lepidoptera. However, the *H. melpomene dennis2* element contains extensive sequence conservation with *H. erato* (as well as a short region of conservation with other nymphalids; Hmel18003: 819500–82000), and we also find a narrow region of homology between *dennis2* (Table S20) and the *D* locus controlling the dennis phenotype in *H. erato*. The *H. melpomene optix band* element identified here is located within 50 kb of the gene *optix*, but at a substantial distance from the *Y* element that controls the corresponding phenotype in the *H. erato* clade (Van Belleghem et al. 2017) in a region of low conservation with *H. erato*. The physical distance between the elements in the two species indicates that the evolution of the red band occurred by evolution of regulatory changes at unrelated loci. At the *WntA* locus, the *WntA1* and *WntA2* elements we identify contain homologous sequence with the *H. erato St* (a likely promoter of the *WntA* gene) and *Ly* elements respectively (Table S20). The *WntA1* element contains two peaks of conservation with multiple other lepidopteran species (Figure S18), corresponding to the two 3’ coding exons of the
WntA gene. At cortex, the H. melpomene cortex band element was located between the two large yellow bar-linked regions identified in H. erato (Van Belleghem et al. 2017). Overall we find that several of the putative regulatory elements we have identified as controlling patterning in H. melpomene and the silvaniform clade (ray2, dennis2, WntA1, and WntA2) appear to be at least partly homologous with regulatory elements proposed to control similar phenotypes in H. erato, a species from which they diverged ~12 MYA (Kozak et al. 2015).

Discussion

In this study, we have investigated the contributions of three genetic modes of evolution (divergent genetic mechanisms, parallel genetic evolution, and collateral evolution) to explain convergent wing color pattern phenotypes found in Heliconius butterflies (H. melpomene, H. elevatus, H. besckei, and H. erato). Using phylogenetic analyses, we have identified strong and narrow signals of shared ancestry related to wing pattern phenotypes in H. melpomene, H. elevatus, and H. besckei around all three main color pattern genes (optix, WntA, and cortex). This is indicative of collateral evolution of putative regulatory elements among these closely related species. In contrast, signals of consistent shared ancestry among these taxa were low outside of these regions. However, we also find that convergent phenotypes between H. melpomene and H. elevatus in the Guianas appear to have arisen independently and so are not a result of collateral evolution. We also show that four out of the seven putative regulatory elements around optix and WntA in H. melpomene also show some homology to regulatory elements controlling similar phenotypes in its distant comimic H. erato. Thus, convergent phenotypes between these more distantly related species appear to result from a combination of convergent parallel evolution and divergent genetic mechanisms. Overall, our results show that all three genetic modes of evolution underlie convergent phenotypes among mimetic Heliconius species, but that these likely operate at different evolutionary timescales.

Modularity of mimicry facilitates pattern switching

Recent studies in a variety of organisms have demonstrated the importance of combinatorial evolution, where ancient alleles are reused in novel combinations to generate new phenotypes and adaptive combinations (Marques et al. 2019). This can lead to adaptive convergent changes more rapidly than evolution via divergent genetic mechanisms or parallel genetic evolution. This can be seen in cichlids, where regulatory changes at the gene agouti-related peptide 2 are associated with the convergent stripe evolution across species (Kratochwil et al. 2018), and in sticklebacks, where recurrent deletions of the same pitx1 enhancer in different populations have led to reduced or lost pelvic structures (Chan et al. 2010). Another excellent example are the diverse wing patterns of Heliconius butterflies. These butterflies appear to have a flexible toolkit of cis-regulatory enhancers (Wallbank et al. 2016; Van Belleghem et al. 2017) through which gene
expression changes can rapidly alter phenotypes and drive adaptive evolution (Wray 2007), with a single mutation at an enhancer potentially enough to have major phenotypic effects (Chan et al. 2010; Frankel et al. 2012). Such genetic architecture combined with introgression can facilitate adaptive evolution through the swapping of these enhancers among lineages of Heliconius (Wallbank et al. 2016; Moest et al. 2019; Lewis and Van Belleghem 2020). For example, the evidence suggests that the ancestral sources of the ray and dennis elements were different, with the rays phenotype originating in the H. melpomene clade and the dennis phenotype originating in the silvaniform clade, before being brought together as the dennis-rayed phenotype in both H. melpomene and H. elevatus (Wallbank et al. 2016).

Our analysis, which narrows the dennis and ray elements and splits them into two loci each, is consistent with this finding of multiple origins, even finding separate origins for each ray locus. We also identify a putative optix band locus near optix that suggests the red band is ancestral to the H. melpomene clade, while its absence is ancestral to the silvaniform clade. The two putative loci near WntA that we propose control the full black melanic discal forewing phenotype of H. elevatus (which replaces the broken black pattern in other elevatus races) and the optix band, while the dennis phe-
notype originating in the silvaniform clade, before being brought together as the dennis-rayed phenotype in both H. melpomene and H. elevatus (Wallbank et al. 2016).

In contrast, the lack of rays in H. elevatus tumatumari appears to be due to having the no-ray allele introgressed from H. melpomene meriana being replaced by a no-ray allele through recombination. In contrast, the lack of rays in H. melpomene meriana appears to be due to having the no-ray allele introgressed from the rayed allele found in other rayed races of H. melpomene and H. elevatus, we hypothesize that H. elevatus tumatumari has secondarily and independently evolved a no-ray phenotype. This can occur via sequence duplications (Eichenlaub and Ettwiller 2011), transposable elements that lead to the trans-
location of regulatory elements from one gene to another (Daborn 2002; Domené et al. 2013), or the co-occurrence of regulatory sequences to derive novel expression patterns (Rebeiz et al. 2011). While our analyses cannot identify the elements that are not shared across taxa, we are able to show that the convergent no-ray phenotypes in mimetic H. mel-
pomene meriana and H. elevatus tumatumari appears to have resulted from independent evolution within each species because there is no signal of shared ancestry at the ray1 or ray2 regions between these taxa (Figure 2, E and F). Given our evidence that introgression of phenotypes between H. mel-
pomene and silvaniform taxa appears to have occurred in Peru, and that H. elevatus tumatumari appears to have the rayed allele found in other rayed races of H. melpomene and H. elevatus, we hypothesize that H. elevatus tumatumari has secondarily and independently evolved a no-ray phenotype. In contrast, the lack of rays in H. melpomene meriana appears to be due to having the no-ray allele introgressed from H. melpomene meriana then later gaining the dennis and yellow band phe-
notypes through introgression, or alternatively, a result of the ray allele of H. melpomene meriana being replaced by a no-ray allele through recombination with H. melpomene melpomene. In addition, our analyses also show that the forewing broken band phenotype found both in H. melpomene meriana/thel-xiopeia and H. elevatus tumatumari/bari in the Guianas must also have independent origins as we see no signal of shared ancestry between these taxa around WntA.

Our results indicate that mimicry via introgression between H. elevatus and H. melpomene has therefore not occurred consistently across their ranges. A possible scenario is that introgres-
sion first occurred in Peru allowing species to switch or perhaps create new mimicry rings, with these newly introgressed alleles and then simply persisting in the Guianas, where independent convergent evolution has refined local mimicry leading to H. elevatus losing the rays and melanic WntA phenotypes. This
means that both introgression and the independent convergent evolution of novel cis-regulatory elements, has been important in driving mimicry between these two species.

**Convergence and sequence conservation**

Our results further support the notion that cis-regulatory modularity is common across mimicry genes in *Heliconius*. Having refined and identified putative cis-regulatory elements, we investigated whether these intervals showed sequence conservation between *H. erato*, *H. melpomene* and other Lepidoptera, using sequence conservation as a proxy for cis-regulatory function. While the *band* element regulates the expression of *optix* in both *H. erato* and *H. melpomene*, this was achieved by divergent genetic mechanisms. On the other hand, we have found evidence of parallel evolution in the modification to the 5' noncoding region of *WntA* in both *H. melpomene* and *H. erato*; these modifications occurred in evidently homologous regulatory elements, despite their independent evolution.

The broad region around the *ray* and the *dennis* elements contains a high density of deeply conserved sequences, but appears to be a hotspot for the modification by selection.
Indeed, we observed this in the tations to occur close together in convergent species, and in the ancestor, then we would expect to (Blount et al. Tollis or not such a trend persists will be revealed. granularity, but, as more examples are characterized, whether been identi- examples of independent convergence in regulatory sequence have yet been identified. It is therefore yet to be seen whether there is a general trend of convergence across taxa at this level of granularity, but, as more examples are characterized, whether or not such a trend persists will be revealed.

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