

A homeotic shift late in development drives mimetic color variation in a bumble bee

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Natural phenotypic radiations, with their high diversity and convergence, are well-suited for informing how genomic changes translate to natural phenotypic variation. New genomic tools enable discovery in such traditionally nonmodel systems. Here, we characterize the genomic basis of color pattern variation in bumble bees (*Hymenoptera*, *Apidae*, *Bombus*), a group that has undergone extensive convergence of setal color patterns as a result of Müllerian mimicry. In western North America, multiple species converge on local mimicry patterns through parallel shifts of midabdominal segments from red to black. Using genome-wide association, we establish that a *cis*-regulatory locus between the abdominal fate-determining *Hox* genes, *abd-A* and *Abd-B*, controls the red-black color switch in a western species, *Bombus melanopygus*. Gene expression analysis reveals distinct shifts in *Abd-B* aligned with the duration of setal pigmentation at the pupal-adult transition. This results in atypical anterior *Abd-B* expression, a late developmental homeotic shift. Changing expression of *Hox* genes can have widespread effects, given their important role across segmental phenotypes; however, the late timing reduces this pleiotropy, making *Hox* genes suitable targets. Analysis of this locus across mimics and relatives reveals that other species follow independent genetic routes to obtain the same phenotypes.

Bombus | Müllerian mimicry | evo-devo | *Abd-B* | *Hox*

Understanding how genomic change drives phenotypic evolution is a central goal of evolutionary biology. Such genotype-to-phenotype connections enable us to decipher the genetic code and together provide predictable trends, such as whether evolution tends to be mediated by few large-effect or many small-effect loci (1–3), targets the same genes and mutations for the same phenotypes (4–7), and modifies genes in *cis*- or *trans*- (8, 9), and whether certain types of genes are evolutionary hot spots (8, 10, 11). Adaptive traits that evolve independently in two or more lineages are especially suited for understanding these trends as they provide natural replicates (11).

Müllerian mimicry, in which two or more distasteful species share similar aposematic signals to avoid predation, presents exceptional examples of evolutionary convergence and thus potential for evolutionary genetics (12–15). Numerous insights have been gained, for example, in studying the genetic basis of wing pattern convergence in mimicry complexes of *Heliconius* butterflies. These studies have revealed that butterfly wing patterns are generated by *cis*-regulatory modifications to a few major developmental genes co-opted for novel function in the wing (16–20) and that diversity and convergence in phenotypes can involve different regulatory modifications to these same loci (21–23). Targeting of these genes has also revealed microevolutionary processes, showing that mimetic phenotypes can be differentially partitioned across populations and species through processes of ancestral allele sorting (24) and adaptive introgression (21, 25).

Bumble bees exhibit a similarly striking level of color pattern diversity partially as a result of Müllerian mimicry. The ~260 species in the genus (26) exhibit over 400 color patterns, making most bumble bee species polymorphic (27). Bumble bee color patterns are bright and contrasting, an aposematic signal of their toxic

sting (28, 29). Color is imparted by dense, setal pile coating the cuticle and is highly modular, tending to change color—switching between white, black, red-orange, brown, or yellow—by segmental sclerites of the head, thorax (mesosoma), and abdomen (metasoma) (27, 30). Many of the conceivable combinations of color across these segments have been achieved (refs. 30 and 31 and Fig. 14). Although multiple ecological factors may play a role in bumble bee color diversity, Müllerian mimicry is believed to drive much of their polymorphism (27). Color patterns have been documented to converge geographically onto over 24 global mimicry complexes, and species that span multiple mimicry complexes tend to converge onto each local pattern (27, 32, 33). This system, thus, has potential for revealing evolutionary developmental and genetic processes: it can inform processes driving sclerite identity and has abundant replicates across global mimicry complexes to assess whether similar mechanisms generate the same phenotypes and how these variants sort across populations and species.

Significance

Mimicry among bumble bees has driven them to diversify and converge in their color patterns, making them a replicate-rich system for connecting genes to traits. Here, we discover that mimetic color variation in a bumble bee is driven by changes in *Hox* gene expression. *Hox* genes are master regulators of numerous segment-specific morphologies and thus are some of the most conserved developmental genes across animals. In these bees, the posterior *Hox* gene *Abd-B* is upregulated in a more anterior location to impart phenotypic change. This homeotic shift happens late in development, when nonspecific effects are minimized, thus availing these genes for color pattern diversification. Similar mimetic color patterns were inferred to use different mutations, suggesting diverse routes to mimicry.

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Data deposition: The raw sequencing reads for genomic samples reported in this paper have been deposited in the National Center for Biotechnology Information BioProject database, <https://www.ncbi.nlm.nih.gov/bioproject> (accession no. PRJNA526235). The bioinformatic analysis protocols, codes, scripts, associated analysis files, and qPCR CT value data reported in this paper have been deposited in the Dryad Digital Repository: <https://doi.org/10.5061/dryad.3v6f25v>.

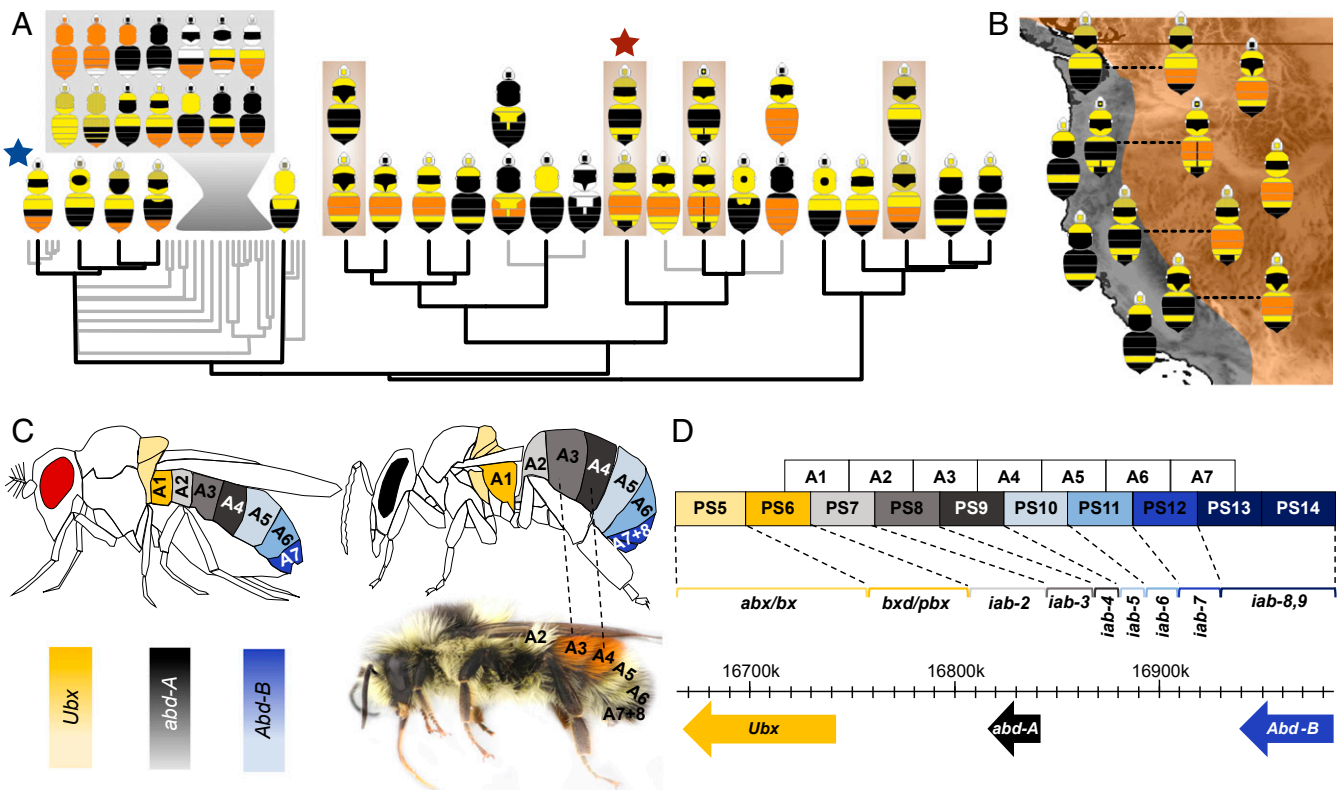
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Results and Discussion

Genetic Locus Driving Color Pattern Variation. Genome-wide association analysis between 21 red and black *B. melanopygus* males sampled from across the hybrid zone resulted in a single fixed ~50-kb peak of association in an intergenic domain between *Hox* genes *abd-A* and *Abd-B* of the bithorax complex (BX-C) (Fig. 2). In insects, the BX-C cluster involves three segmental *Hox* genes, *abd-A*, *Abd-B*, and *Ubx*, which specify posterior segmental phenotypes (Fig. 1C). Further fine-scale Sanger sequencing of 133 individuals across this block narrowed the associated region to a 4-kb, zero-recombinant interval closest to and ~50 kb downstream from the *Abd-B* transcript. In *Drosophila*, the intergenic region between *abd-A* and *Abd-B* (the *iab* region) is the primary regulatory region directing expression of both *abd-A* and *Abd-B*. The *iab* region is divided into several noncoding domains (*iab*-2 to -9) that direct the segmental expression of these genes in a collinear fashion, with the regions closest to *abd-A* directing the most anterior segmental expression of this gene (*iab*-2) and those closest to *Abd-B* directing the most posterior *Abd-B* expression (*iab*-9) (Fig. 1C and D) (41–43). Only a small proportion of the *B. melanopygus* *abd-A/Abd-B* region can be reliably aligned to *Drosophila*, including the *iab*-4/8 micro-RNA (44) and the protein-coding sequences of the two *Hox* genes (SI Appendix, Fig. S1). This is not unexpected, as the *iab* regions exhibit 12% sequence homology even across *Drosophila* species (45). The relative location of the *iab*-4 miRNA in this intergenic region, however, appears to be conserved across insect lineages, including *B. melanopygus* (46–48), suggesting some level of conservation in relative locations of regulatory domains. If in bumble bees *iab* regions were to function in the same collinear manner as in *Drosophila*, the relative location of the fixed color locus in *B. melanopygus* would most likely fall in *iab*-6/7, which in *Drosophila* would direct the expression of primarily *Abd-B* in the most posterior tergites (A6 and A7; bumble bee “tail”) (49) (Fig. 1C and D and SI Appendix, Fig. S1).

BX-C Gene Expression Across Bumble Bee Development. The location of the color locus between *abd-A/Abd-B* suggests that this phenotype is likely driven by shifts in expression of one of these *Hox* genes. Holometabolous insects undergo two major developmental transitions: (i) in the embryo to build larvae, and (ii) during late larval and pupal stages to build adult structures. The majority of developmental work focuses on how these *Hox* genes operate to establish segmental identity in embryos; however, *Hox* genes are likely to operate throughout development (50, 51). All three BX-C genes were shown to be expressed in the developing adult abdominal epidermis in *Drosophila* pupae in a spatial pattern similar to the embryo, although with more discrete domains (52). In bumble bees, setae develop in early white to pink-eyed pupal stages and are pigmented in the late pupal stages; thus, the relevant gene expression patterns for color should occur in pupae (53). To test how *Hox* genes fluctuate across metamorphosis in bumble bees and determine their potential to play a role in pupal stages, we performed qPCR in the model bumble bee *Bombus impatiens* at different developmental stages including whole embryos and larvae, and on metasomal terga of pupal and early adult stages (callow). mRNA of the three BX-C genes were detected across all developmental stages, even into adulthood (Fig. 3A). The level of mRNA expression, however, fluctuated, with peak levels of mRNA appearing at the major periods of metamorphosis, including in embryos and during the larval-pupal transition. This correlation with developmental change suggests that *Hox* genes continue to be players as fate-determining selector genes across development rather than relying on triggered effector genes downstream to regulate these phenotypes.

We further examined the location and timing of expression of each of the BX-C genes in *B. melanopygus* by metasomal segment. In pupal metasomal tergites of *B. melanopygus*, *Abd-B* is expressed primarily in metasomal segments T4–6 (Fig. 3B) [abdominal segments A5–7 (Fig. 1D)], *abd-A* in metasomal segments T1–6 (Fig. 3B) [abdominal segments A2–7 (Fig. 1D)], with highest levels in T3–4, and *Ubx* primarily in metasomal segment

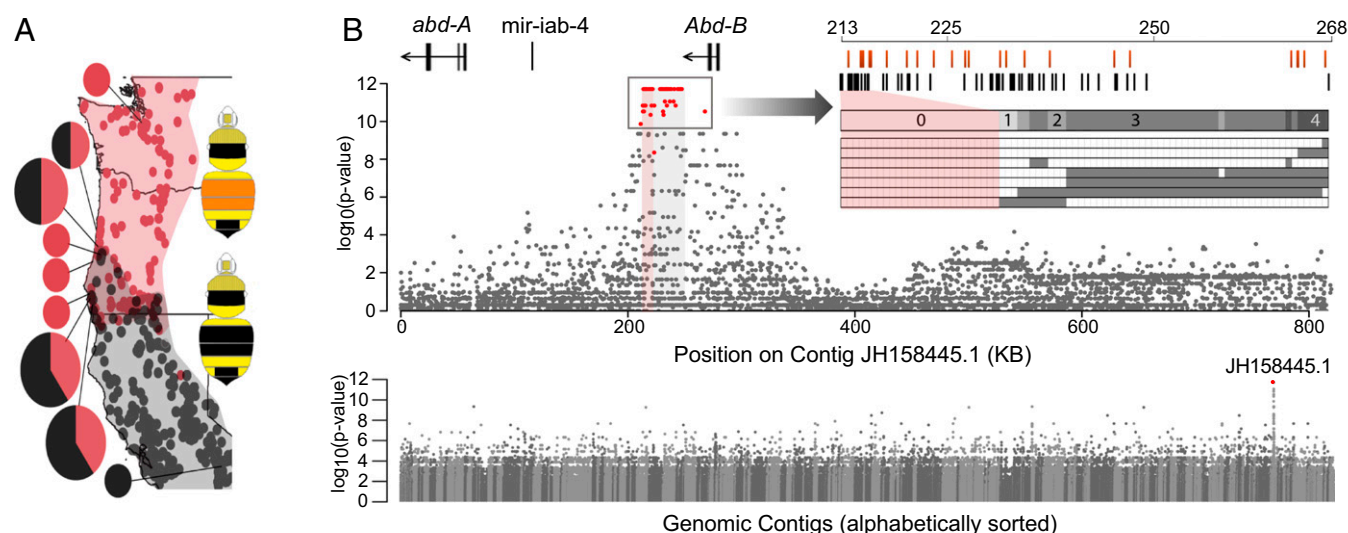


Fig. 2. The color locus is revealed using a genome-wide association study (GWAS). (A) Map of localities from which GWAS samples were collected. The dots on the map indicate the overall distribution of each color form from specimen data (34). The pie charts indicate sites where samples were collected for this analysis with the size and portion of these charts reflecting the number of samples per locality ($n = 1$ –5). (B) Whole-genome association (Bottom) showing that sites of fixed association (indicated in red) are confined to a single contig (contigs in alternating gray). The association within this contig (above) demonstrate that fixation (red dots) is confined to a ~50-kb block 3' of *Abd-B* (highlighted gray). Highlighted in red is the region that remains fixed with additional genotyping, further highlighted with the *Inset* SNP plot. The rows at the Bottom of the *Inset* plot represent ordered SNP/indel variants of recombinant individuals, with gray indicating SNP/indels belonging to the opposite phenotype in genomic samples and which thus break fixation. The number of individuals breaking fixation is indicated above these lines. At the Top, SNPs (black) and indels (red) and their position are indicated, and fixed sites are aligned with the respective sites in the plot below using highlighting.

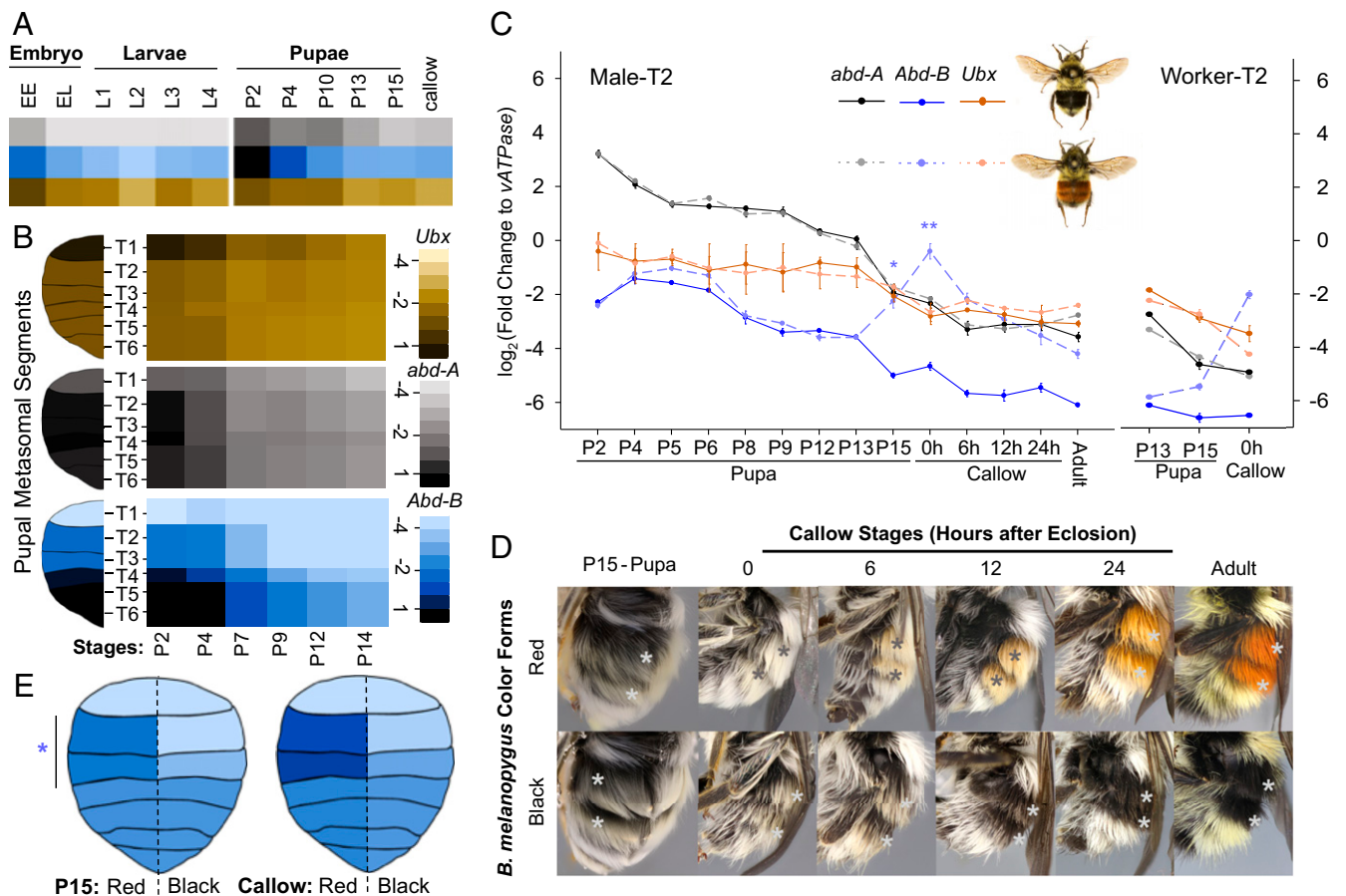


Fig. 3. Expression pattern of the bithorax complex (BX-C) homeotic genes during bumble bee development. (A) Relative expression of BX-C genes across *B. impatiens* developmental stages, including embryo, larvae (whole body), pupa, and early adult (metasomal tergites). (B) Spatial and temporal distribution of the BX-C genes in metasomal tergal segments of *B. melanopygus* pupae (worker, red form) from stage P2 to stage P14. Shades for both (A and B) represent log₂ relative fold change to vATPase. Right lane, scales for relative expression level. (C) Temporal expression pattern of the BX-C genes at the dimorphic metasomal tergal segment 2 (T2) of male *B. melanopygus*. Note that, in the callow adult stage, the relative abundance of *Abd-B* is higher than *abd-A* and *Ubx*. **P* < 0.01, ***P* < 0.001, Mann–Whitney *U* test. *Abd-B* expression was marginally significantly higher with <2× difference in the red form at P5E (*P* = 0.04) and P6 (*P* = 0.03); however, this difference is only seen in T2 but not T3 (SI Appendix, Fig. S2). A similar homeotic shift of *Abd-B* in the red form is also observed in polymorphic segments of workers (Right) (*P* = 0.057 at P15; no statistics applied at 0 h as only one sample is in black form). (D) Setal color development in late pupal and early adult stages of *B. melanopygus*. The stars indicate polymorphic segments. For P15 pupae, pupal cuticle was removed to reveal the developing adult setae underneath. The red and black setae are distinguishable in males starting from 0-h callow adult. Color continues to intensify in the first 24 h posteclosion. (E) *Abd-B* expression across abdominal tergites of *B. melanopygus* at stage P15 and in the newly eclosed callow adult showing up-regulation of *Abd-B* only in T2 and T3 for both of these stages, and only in the red form (segments on the Left side), not the other segments and the black form.

T1 (Fig. 3B) [abdominal segment A2 (Fig. 1D)]. The pupal expression pattern of the BX-C genes in bumble bees is thus consistent with their conserved spatial pattern in embryos across insects and other arthropods (i.e., *Ubx* anteriorly, *abd-A* medially, *Abd-B* posteriorly; Fig. 1 C and D) (40, 46, 51, 54). Consistently across segments, the expression of each BX-C gene was highest at the earliest stages of pupal development and declined thereafter (Fig. 3B).

BX-C Gene Expression Between Color Phenotypes. To determine whether setal color variation in *B. melanopygus* is associated with differential expression of BX-C genes during adult setal development, we performed qPCR on dissected pupal epidermal tissues of the dimorphic second and third metasomal terga (T2 and T3), run separately, on male pupae of the red and black phenotypes. As BX-C genes were determined to be expressed throughout early pupal to adult stages, we analyzed differential expression patterns across pupal–adult development to avoid missing any critical stages of gene regulation. Nine pupal and four callow adult stages, representing fine increments of pupal–adult development,

were sampled with staging based on morphological criteria (53). In both red and black bees, all three BX-C genes exhibit the highest levels of expression at the early pupal stage, with *abd-A* expression being higher than *Ubx* and *Abd-B* (Fig. 3C), as is expected given the typical dominance of *abd-A* in midabdominal segments. The expression of all three genes gradually decreases during pupal development, while their relative levels remain the same. Dramatic differences, however, were seen at the late pupal stages where a significant up-regulation of *Abd-B* was observed only in the red form in both dimorphic metasomal terga (T2 and T3), while *abd-A* and *Ubx* expression continued to decrease [T2 (Fig. 3C); T3 (SI Appendix, Fig. S2)]. In the red form, mRNA levels of *Abd-B* increase starting between stages P13 and P15 and peak with a 10–20× expression difference (*P* < 0.001) from the black form at the time of eclosion (0-h callow adult), returning to low background levels by 24 h posteclosion (Fig. 3C). Similar differentiation of *Abd-B* expression between the two color forms was also observed among the female worker caste (Fig. 3C). This *Abd-B* up-regulation was confined only to the dimorphic second and third metasomal segments, and not to the first segment or the more

posterior segments (Fig. 3E), supporting its role in generating the dimorphic differences observed in these segments. This is further supported by the timing of the differentiation, which is well aligned with the timing of differential setal pigmentation (Fig. 3D). Pigments for both red and black setae start to deposit as early as the late-P12/P13 pupal stage (~6 d after pupation in workers) as a pale greyish orange color; however, the differences in color between black and red setae are not macroscopically distinguishable until around stage P15 in queens and 0-h callow adults in male and workers (~8 d after pupation), when black setae exhibit a grayer tinge, while the red setae become more orange (Fig. 3D). Both red and black setae attain most of their color within the first 6 h posteclosion and complete pigmentation by 12–24 h (Fig. 3D). The expression of this gene concordant with the expression of the phenotype is in contrast to regulators of pigmentation in butterflies, which are turned on to premap the wing pattern early in development before wing pigmentation actually takes place (16, 55). The function of *Abd-B* is similar to its mode of action during embryonic development, where it controls onsite segment identification, and is typical for a selector gene (56, 57).

During this up-regulation, *Abd-B* mRNA has higher expression than that of the other BX-C genes, including *abd-A* (Fig. 3C). In *Drosophila*, *Abd-B* is considered to exhibit dominance over *abd-A*, leading to repression of *abd-A* translation and *Abd-B* segmental identity where it occurs (58). In other cases, combinatorial effects rather than complete dominance interactions have been observed (59, 60). The increase of *Abd-B* over *abd-A* expression suggests that regardless of the dominance mode, this uptick likely generated a homeotic shift late in development of the T2 and T3 segments from midabdominal to more posterior-abdominal identity.

These results suggest that shifting the expression of the major developmental gene *Abd-B* likely was the initial trigger driving a shift in melanin gene expression in *B. melanopygus*. Previous work examining genes driving natural and laboratory-generated differences in melanization in *Drosophila* support mutations occurring primarily in *cis*-regulatory regions of pigment enzymes (38). These studies have served as examples of how the genome tends to be targeted by changing downstream targets in *cis*-, where these mutations have reduced pleiotropic consequences (38, 61). Although they are not typically the evolutionary target, upstream players like *Hox* genes have been found to play a role in differential melanization in *Drosophila* (61). For example, sex-specific variation in the expression of developmental gene *bric-a-brac* impacts the ability of *Abd-B* to trigger tissue-specific differences in melanization by sex (39, 62). *Abd-B* is not known in this case to change its expression to mediate natural adaptive color pattern variation, but rather directs the posterior location of the dimorphism, acting as it typically does as a segment-specific selector. Shifts in the *Hox* genes themselves could be occurring in *Drosophila*, as minor effect mutations that could drive differences in *abd-A* and *Abd-B* expression have been linked to sex-specific pigment variation in some species (63).

Hox genes play diverse roles in driving the fate of all phenotypes in their respective segments; thus, changes in these genes can have highly pleiotropic consequences. Likely because of the deleterious effects of such mutations, the relative location of expression in these genes during embryogenesis is conserved across insects and other arthropods (51). In the case of bumble bees, the genetic changes in the *Abd-B cis*-regulatory region are not associated with any other apparent phenotype that would suggest segmental homeosis. This may be because the shift of *Abd-B* generated by these genetic changes occurs at the end of development, when the only marked changes in morphology involve melanization and sclerotization. Shifting *Abd-B* expression at these stages may thus switch setal pigmentation with minimal pleiotropic impacts on other tissues. These consequences can be even further reduced if up-regulation of *Abd-B* was cell-specific, as insect setae are single-celled structures (64).

If negative pleiotropic consequences can be avoided, upstream regulators are excellent targets for evolution and diversification, as these genes have already established numerous developmental genetic connections to impart change (8, 65).

Late developmental spatial shifts in expression of *Hox* and other upstream regulators can be an important mechanism for driving phenotypic change (57, 66). For example, in arthropods, localized shifts in *Hox* genes in later stages are responsible for variation in limb morphologies. Shifts in *Scr* expression influence male sex comb morphology in *Drosophila* forelimbs (67) and the formation of crustacean maxillipeds (68). Similarly, regional up-regulation of *Ubx* along insect hindlimbs drives growth and setal loss (69, 70), such as that observed in the expanded corbicular pollen basket in honey bees and bumble bees (71). Such variation during late development could be driven by *Hox* genes evolving to respond to downstream components of regulatory networks, further enabling developmental micromanagement (57, 67).

This leaves open the question as to how a homeotic shift generates an on-off switch for pigmentation in bumble bees. Many genes throughout the genome pick up binding sites for *Hox* genes to direct their segment-specific expression. At one point, *cis*-regulatory binding of *Abd-B* may have been picked up by a gene in the melanin pathway to turn on red pigmentation where it occurs in the posterior of the abdomen. The most common pattern across bumble bees is a yellow bee with a red-tipped abdomen (ref. 27 and Fig. 1A), an aposematic signal to frame its venomous sting. This is most likely the ancestral bumble bee pattern (e.g., it is common in the early diverging *Mendacibombus*) and is common throughout the Old World where bumble bees originated (31). The location of ferruginous stripes more anteriorly in bumble bees is a derived condition primarily found in the North American *Pyrobombus*, belonging to *B. melanopygus* and several of its close relatives (Fig. 1A). A more anterior expression of red color could have thus been achieved merely by relocating *Abd-B* expression from posterior to anterior segments, a mutation that, once generated, may have no other fitness consequences due to the loss of pleiotropy in these tissues late in development. Co-option of established gene networks in new locations to generate new phenotypes (heterotopy) is an emerging source of evolutionary novelties. For example, *Drosophila* wing spots were generated by the wingless gene being co-opted to turn on pigmentation in its original location of expression, followed by wingless shifting expression to multiple additional parts of the wing to generate spots (72). Similarly, the *optix* gene is involved in eye formation and pigmentation in insects and has been co-opted in butterflies to turn on the same pigments in the wing (19, 73). As *Hox* genes are major genes influencing the location of phenotypes, these genes could help promote rapid changes in color pattern.

Concerted Mutations Operate in the Regulatory Domain. Beyond the 4-kb locus, several recombinant red-form individuals were found bearing black haplotypes in the remaining block that is fixed in genomic samples (Fig. 2B and *SI Appendix*, Fig. S3). Although appearing primarily red by eye, detailed color analysis revealed that individuals with these recombinant haplotypes exhibit various levels of penetrance of the black phenotype, with more black setae mixed among the red. This suggests that, although the narrowed 4-kb fixed region contains sequence variants able to cause a nearly full color switch, there are variants within the interval outside this region that contribute to the phenotype. This explains why the fixed region occurs in a large block and lends support to adaptive phenotypes being driven by multiple linked SNPs that have evolved in concert as opposed to single mutations, as has been shown in regulatory regions of *Drosophila* (74–76). The red form was previously determined to have complete dominance, retaining red color in heterozygous individuals (35). With our sequencing of the color locus, we were able to identify and compare homozygous and heterozygous females,

and found that heterozygotes exhibited black hair penetrance similar to the recombinants (SI Appendix, Fig. S3), suggesting that gene expression is reduced in a similar manner for both. This indicates that dominance is partially incomplete.

The Color Locus and *Abd-B* Regulation. The function of the mutations in this color locus in driving *Abd-B* differential regulation remains unknown. The location of the color locus relative to known segmental regulatory domains in *Drosophila* suggests it may normally regulate more posterior segments than it does, in line with the idea that this region may have first evolved to regulate color in more posterior phenotypes and have been subsequently mutated to generate an anterior shift. Laboratory-generated *iab cis*-regulatory mutants in *Drosophila* can generate such an effect (77, 78), with homeotic shifts in segmental identity generated, for example, by altering insulators demarcating *iab* segmental regulatory regions (79, 80). Furthermore, in *Drosophila*, the entire region between *abd-A* and *Abd-B* is transcribed and spliced into a massive nc-RNA (*iab-8*-ncRNA) (41, 42, 49). This could be another source of *Abd-B* regulation (81). Further research on this regulatory locus would enable a more complete understanding of the regulatory mechanisms driving the *Hox* toolkit.

Implication of the Color Locus in Comimetic Species. To test whether the same genetic mechanism underlies parallel color variation displayed by comimetic species and decipher the ancestry of these phenotypes, we compared sequences of the fixed color locus from four dimorphic species displaying similar black and red color variation in metasomal segments T2 and T3 as well as in six monomorphic sister lineages bearing either red or black phenotypes in these segments. None of the fixed SNPs and indels

for *B. melanopygus* in an 18.5-kb genomic region including our fixed interval show the same association with color variation in other species, even those species most closely related to *B. melanopygus* (e.g., *B. sylvicola*) (Fig. 4). This supports the independent evolution of at least one of the two phenotypes from the ancestor in *B. melanopygus*. Mimetic color variation in these western North American mimicry complexes thus involves multiple, independent genetic mutations, rather than sorting of ancestral variation, even among close relatives with similar phenotypes. This lends support to the potential for mutation as opposed to alternative allele sorting in contributing to phenotypic diversity (82–84). This contrasts with *Heliconius* butterflies, where close relatives obtained their patterns from a common ancestral mutation, either through introgression (25) or ancestral polymorphisms (24). Although the same SNPs are not implicated, it is possible the same gene may be involved in these other species, but using different genetic targets. This was observed, for example, in *Heliconius* butterflies *H. erato* and *H. melpomene*, where comimicking forms were generated by the independent targeting of different parts of the *cis*-regulatory region of same gene, *optix* (23, 85, 86).

Most likely, the independently evolved phenotype is a black form from the red form. This is inferred because the red form tends to contain the same conserved ancestral sequences as the other taxa with most of the functional SNPs and indels derived in the black form (Fig. 4). The black form exhibits higher linkage disequilibrium (LD) and reduced nucleotide diversity across the color locus contig, suggesting that the region is under selection and more so in this color form (SI Appendix, Fig. S5). If black is derived, this seems counter to gene expression patterns, as red, not black, up-regulates *Abd-B*. Further gene expression analyses across bumble bee species are needed to examine when *Abd-B*

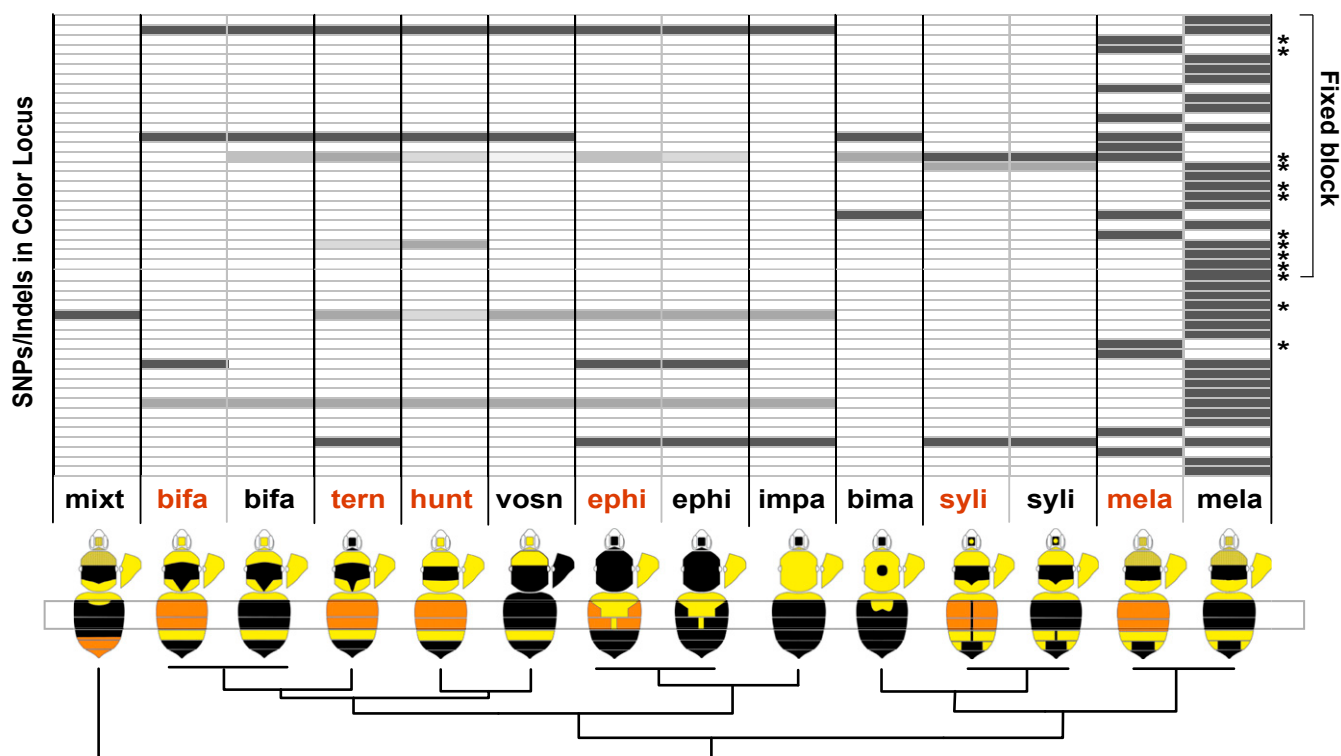


Fig. 4. Allelic variants in the color locus of *B. melanopygus* comimics and close relatives. No fixed SNP or indel are similarly differentiated by color in other species. At *Bottom* are species related to *B. melanopygus* (Right, "mela") that vary in the red/black color of metasomal segments T2 and T3 (segments indicated with a gray box) and their phylogenetic positions (Fig. 1). Species names and the respective color of the specimens are abbreviated above each picture. A map of the character state of each ordered SNP or indel (starred) across an 18.5-kb window that includes the narrowed fixed block (Fig. 2) is shown above. SNPs that are derived are shaded dark gray and ancestral alleles are white. Medium gray shading indicates alternative SNPs and indels to those in *B. melanopygus*.

up-regulation originated to determine whether this is a gain in the red form or a loss in the black form.

Conclusion and Future Directions. Our study demonstrates that evolution can target well-connected, highly conserved, pleiotropic developmental genes to generate phenotypic variation. Modifying regional selector genes, such as *Hox* genes, in *cis*-allows time-specific changes in expression, thus reducing pleiotropic consequences and making these genes suitable targets for driving region-specific variation in phenotypes. Targeting positional genes, such as *Hox* genes, may enable the diversification of segmental phenotypes seen in this lineage. Bumble bee coloration is a new system for evolutionary genetics. With this discovery, we can begin to explore how segmental variation is regulated and the multitude of ways evolution has achieved the same phenotypes. As a nonmodel group, functional genetics is more challenging, but the comparative potential provided by the extent of interspecific and intraspecific variation still enables strong candidates to be identified and can inform evolutionary processes. Studying abdominal *Hox* gene expression during pupal development across species will be important for understanding the evolutionary events leading to this phenotypic transition. Furthermore, few natural adaptive loci have been found in *cis*-regulatory regions of *Hox* genes. Deciphering how this regulatory region works in this system will help inform how *Hox* genes operate and evolve in insects beyond *Drosophila*.

Materials and Methods

Genomic Analysis. We sequenced genomes of red ($n = 11$) and black ($n = 10$) males of *B. melanopygus* from multiple localities throughout their morphological transition zone (Fig. 1 and *SI Appendix, Table S1*) including one individual per form outside this zone. Males are haploid, thus enabling improved alignment, assembly (87), and avoidance of dominance effects. DNA extraction was performed on thoracic muscle tissue using Qiagen DNeasy or EZNA (Omega Bio-tek) DNA extraction kits with RNaseA treatment. Samples were barcoded, and prepared for Illumina sequencing using Illumina TruSeq DNA Nano kits. Whole-genome sequencing was performed 5–10 samples/lane on an Illumina HiSeq 2500 sequencer in Rapid Run mode to generate 150-bp paired-end read libraries at the Penn State Huck Genomics Core Facility. Trimming was applied using Trimmomatic, version 0.38 (88), to remove low-quality bases, clip adaptors, and discard short-length sequences (<36 bp). Reads were then aligned to the published genome assembly (89) of closely related species *Bombus impatiens* (GCA_000188105.2, BIMP_2.0) [common ancestor with *B. melanopygus* ~ 10 mya (31)], using BWA-mem algorithm (90) of BWA, version 0.7.17 (91), aligner. Alignment postprocessing was conducted in SAMtools 1.8 (92) and Picard 1.119 (broadinstitute.github.io/picard/index.html). Multisample variant calling was performed using GATK Unified Genotyper (version 3.6-0) (93, 94) with parameters set for haploidy. After filtration of variable sites with substantial missing data ($>25\%$) with VCFtools 0.1.15 (95), a genotype–phenotype association analysis between case-control phenotypes (black and red) and the SNP dataset was performed in PLINK 1.9 (96) using Fisher's exact test.

To identify fixed insertions and deletions in the color locus, de novo genome assemblies of *B. melanopygus* including two for each color form were performed using MIRA assembler, version 4.9.6 (97). Indels were manually identified from assembled contigs and checked for fixation using IGV (98). Summary statistics for genomic samples are available in *SI Appendix, Table S1*. Genomic sequencing data are available under NCBI BioProject accession number PRJNA526235 (99). Additional data are available in the Dryad Digital Repository (100).

Genotyping. To narrow the locus regulating the phenotype, genotyping was performed on 80 red and 53 black *B. melanopygus*, including both males ($n = 108$) and females ($n = 10$ queens, 15 workers). These were sampled mostly from within the transition zone in Oregon and California, but also included areas where exclusively red or black morphotypes occurred in California, Oregon, Washington, and Utah (*SI Appendix, Table S2*). Individuals were subsampled for SNPs and indels across the fixed 50-kb interval to look for potential recombinants, which were then sequenced across all SNPs and indels to better define the window of fixation. DNA was extracted with Qiagen DNeasy or EZNA DNA kits, purified with ExoSAP-It (Thermo Scientific), and PCR amplified (HotStart Mastermix; NEB) with custom-designed

primers (*SI Appendix, Table S3*). Sanger sequencing was performed at the Penn State Huck Genomics Facility. SNPs were called from chromatograms with heterozygosity determined from double peaks. A subset of black males and females, red homozygous males and females, all recombinants, and red heterozygous females were phenotyped under the microscope to determine the extent of mixture of black hairs in red (*SI Appendix, Fig. S3*).

To examine whether the fixed SNPs in *B. melanopygus* are implicated in seemingly parallel color variation in closely related species, we sequenced all SNPs and indels in the narrowed fixed interval as well as an additional 4 kb flanking this from red and black T2/T3 polymorphic and monomorphic sister lineages, including nearly all of the closest relatives. SNPs were called as above and designated as ancestral or derived using parsimony principles. Data are available from the Dryad Digital Repository (100).

qPCR. Developmental stages of *B. impatiens* (Koppert Biological Systems) were used to assess general expression of *Ubx*, *abd-A*, and *Abd-B* across bumble bee development (experiment I; Fig. 3A). Sampling included early (no visible larval segments) and late (visible larval segments) embryos, four larval stages determined based on head capsule width (101), pupae from five stages (53), and a young adult callow, including a single specimen per stage. Embryos and larvae were sampled whole, while pupae and callow adults included dissected ($1\times$ PBS) abdominal tergite epidermis with fat body and muscle tissues removed.

We then compared *Ubx*, *abd-A*, and *Abd-B* gene expression by tergite throughout pupal development in *B. melanopygus* black color form workers (experiment II; Fig. 3B). Colonies of *B. melanopygus* were started from queens from California and Oregon. Abdominal tergites were dissected as above, from reared worker pupae of both red and black forms across six developmental stages (P2, P4, P7, P9, P12, and P14) (53) collecting four tergal regions: metasomal segment T1, T2+3, T4, and T5+6 with three to five replicates per condition.

To compare gene expression between red and black metasomal tergites T2 and T3 (experiment III; Fig. 3C), segments were dissected from male pupae and callow adults of both *B. melanopygus* color forms from nine pupal stages (*SI Appendix, Fig. S4*) covering time points fairly evenly across pupal development (~ 12 h/stage) and four callow adult stages (0, 6, 12, and 24 h posteclosion). Tergites of the polymorphic metasomal segment T2 and T3 were sampled separately. The same tissues were also collected from P13 and P15 pupae and 0-h callows of workers for comparison. For stage P15 male pupae and freshly eclosed male (0-h) callow adults, we also collected tergites from metasomal T1 and metasomal T4–7 (pooled) for comparison (Fig. 3E). Three to seven replicates/pupal stage were used. Pupae and callow samples were collected from five black queen and nine red queen colonies.

All bees were reared at 29°C and $60 \pm 5\%$ relative humidity in the dark. Samples were flash frozen and stored at -80°C . RNA was extracted using an RNAqueous total RNA kit (Invitrogen) for experiment I, RNAqueous Micro RNA isolation kits (Invitrogen) for experiment II, and the Direct-zol RNA MicroPrep (Zymo Research) kits for experiment III. RNA extractions, purification, and subsequent DNase treatment were performed following manufacturer recommendations, with purity and concentration of RNA confirmed using a Nanodrop (Thermo Scientific).

First-strand cDNA was synthesized from 500 ng of total RNA using the High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems). qPCRs were performed in triplicate using PowerSYBR Green PCR master mix (Applied Biosystems) on an ABI 7900 real-time PCR detection system. Primers and conditions for *abd-A*, *Abd-B*, and *Ubx* are provided in *SI Appendix, Table S4*. Stability analysis on five different housekeeping genes was performed in RefFinder (95), leading to the selection of vATPase. A standard curve ($5\times$ dilution series) per gene was used to quantify sample expression, followed by normalization with vATPase (target gene/vATPase) (96). Mann–Whitney U statistical tests were performed to test differences in expression between color forms by stage and gene. Data are available from the Dryad Digital Repository (100).

Population Genetics Analysis. Nucleotide diversity (π) and F_{ST} across the contig containing the color locus was assessed using custom Python scripts (https://github.com/simonhmartin/genomics_general). LD decay analysis was carried out using PopLDdecay (102).

Sequence Conservation Analysis. Comparison of sequence conservation was performed across the *abd-A/Abd-B* intergenic region between *B. impatiens* and *B. melanopygus* red and black forms, bumble bee *Bombus terrestris*, *Drosophila*, and corbiculate bees *Apis mellifera* and *Melipona quadrifasciata*, using mVISTA tools (103, 104) and LAGAN (105) sequence alignment.

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