



ebony* Affects Pigmentation Divergence and Cuticular Hydrocarbons in *Drosophila americana* and *D. novamexicana

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Drosophila pigmentation has been a fruitful model system for understanding the genetic and developmental mechanisms underlying phenotypic evolution. For example, prior work has shown that divergence of the *tan* gene contributes to pigmentation differences between two members of the virilis group: *Drosophila novamexicana*, which has a light yellow body color, and *D. americana*, which has a dark brown body color. Quantitative trait locus (QTL) mapping and expression analysis has suggested that divergence of the *ebony* gene might also contribute to pigmentation differences between these two species. Here, we directly test this hypothesis by using CRISPR/Cas9 genome editing to generate *ebony* null mutants in *D. americana* and *D. novamexicana* and then using reciprocal hemizyosity testing to compare the effects of each species' *ebony* allele on pigmentation. We find that divergence of *ebony* does indeed contribute to the pigmentation divergence between species, with effects on both the overall body color as well as a difference in pigmentation along the dorsal abdominal midline. Motivated by recent work in *D. melanogaster*, we also used the *ebony* null mutants to test for effects of *ebony* on cuticular hydrocarbon (CHC) profiles. We found that *ebony* affects CHC abundance in both species, but does not contribute to qualitative differences in the CHC profiles between these two species. Additional transgenic resources for working with *D. americana* and *D. novamexicana*, such as *white* mutants of both species and *yellow* mutants in *D. novamexicana*, were generated in the course of this work and are also described. Taken together, this study advances our understanding of loci contributing to phenotypic divergence and illustrates how the latest genome editing tools can be used for functional testing in non-model species.

Keywords: CRISPR, Cas9, melanin, nanos, abdominal pigmentation, genome editing, virilis group

INTRODUCTION

Insect pigmentation is a well-studied trait that displays a variety of phenotypic differences within and between species (Wittkopp et al., 2003a; Kronforst et al., 2012). These differences have evolved over a wide range of divergence times and in a great diversity of ecological contexts. Differences in insect pigmentation often appear to be ecologically relevant, correlating with geographic and climatic factors and playing a role in phenomena such as mate recognition, camouflage, thermoregulation, and water balance (True, 2003; Wittkopp and Beldade, 2009). Studies of pigmentation differences within the genus *Drosophila* have emerged as a productive model for studying the evolution of development, exploiting the diversity of phenotypes as well as genetic tools available for working with *Drosophila* and a long history of research into the genetic and biochemical mechanisms controlling pigmentation development (Wittkopp et al., 2003a; Massey and Wittkopp, 2016; Rebeiz and Williams, 2017). Indeed, since the early 2000s, the genetic bases of dozens of pigmentation differences have been identified in varying levels of detail. Strikingly, in every case where a causal role has been directly attributed to a specific gene, the mechanism of change has been found to be a *cis*-regulatory change that affects gene expression rather than a change in the protein's function (Massey and Wittkopp, 2016). These case studies have also identified multiple independent instances of divergent expression for some pigmentation genes, suggesting that these genes are particularly tractable routes for the evolution of pigmentation in this genus (Massey and Wittkopp, 2016).

Changes in *cis*-regulatory sequences are thought to be a common mechanism of developmental evolution because they tend to be less pleiotropic than changes in protein function (Wray et al., 2003; Carroll, 2005). For example, a *cis*-regulatory change might alter a gene's expression in only a single tissue or a single point in development whereas changing its protein function is expected to impact the organism everywhere that protein is expressed. Genes controlling pigmentation development in *Drosophila* might be especially likely to evolve using this mechanism because the proteins encoded by these genes are also required for other biological functions. For example, genes required for pigment synthesis have also been shown to affect mating success, circadian rhythm, vision, and innate immunity (Nappi and Christensen, 2005; True et al., 2005; Suh and Jackson, 2007; Wittkopp and Beldade, 2009; Takahashi, 2013; Massey et al., 2019a). The pigmentation biosynthesis genes *ebony* and *tan* have also been found to affect the profiles of cuticular hydrocarbons on adult flies, which are hydrophobic lipids on the surface of insect cuticle that are involved in chemical communication, mate recognition, and water balance (Chung et al., 2014; Chung and Carroll, 2015; Massey et al., 2019b).

Here, we investigate genetic changes contributing to the evolution of novel body color in *D. novamexicana*. This species has evolved a much lighter and more yellow body color than its sister species *D. americana* during the approximately 400,000 years since these species diverged from their most recent common ancestor (Figure 1; Caletka and McAllister, 2004; Morales-Hojas et al., 2008). *D. novamexicana* and *D. americana*

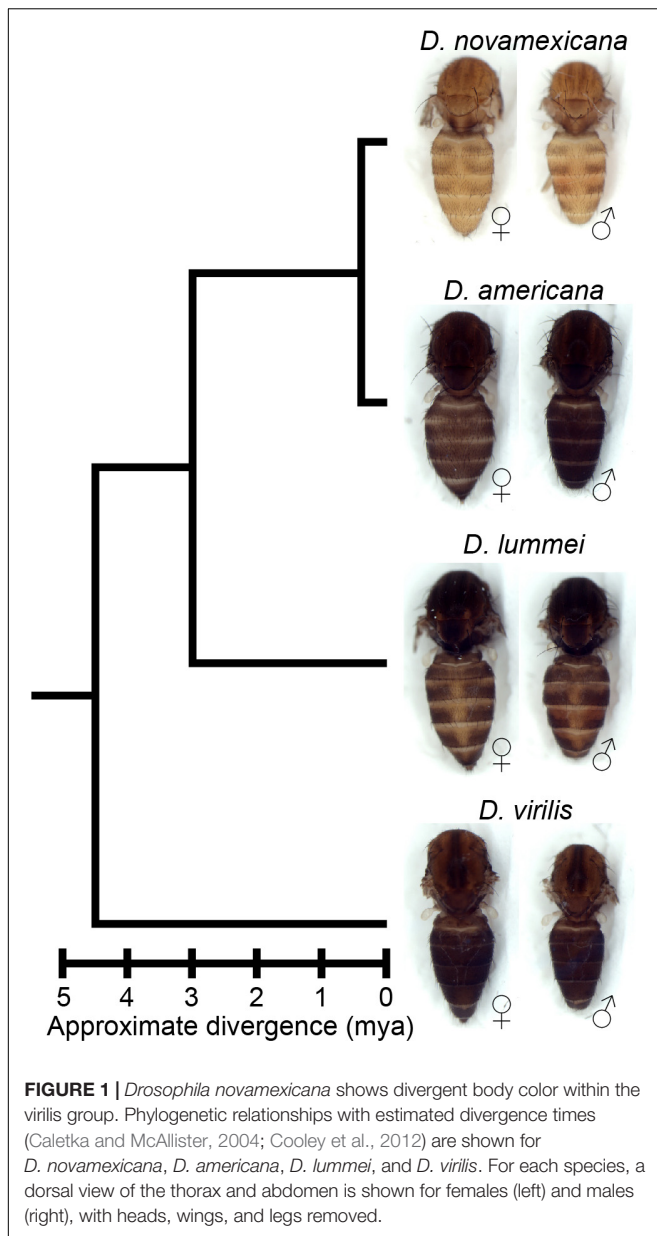
show signs of reproductive isolation (Patterson and Stone, 1949; Ahmed-Braimah and McAllister, 2012), but they are interfertile and can produce viable, fertile F₁ hybrids in the laboratory, allowing genetic analysis (Wittkopp et al., 2003b, 2009). Prior genetic mapping has identified two quantitative trait loci (QTL) that together account for ~87% of the pigmentation difference between *D. novamexicana* and *D. americana* (Wittkopp et al., 2009). Fine mapping and transgenic analysis revealed that the QTL of smaller effect was driven by divergence at *tan* (Wittkopp et al., 2009), a gene that encodes a hydrolase that catalyzes the conversion of N-B-alanyl dopamine (NBAD) to dopamine, a precursor for dark melanin pigment (True et al., 2005). The QTL of larger effect was linked to an inverted region containing the candidate gene *ebony*, but the presence of the inversion prevented fine mapping to separate the effects of *ebony* from linked loci (Wittkopp et al., 2009). *ebony* encodes a synthetase that catalyzes the conversion of dopamine into NBAD, a precursor for light yellow pigments (Koch et al., 2000), which is the opposite of the reaction catalyzed by *Tan*. *ebony* has also been shown to have expression differences between *D. novamexicana* and *D. americana* caused by *cis*-regulatory divergence (Cooley et al., 2012).

Despite these data suggesting that *ebony* contributes to pigmentation divergence between *D. novamexicana* and *D. americana*, the phenotypic effects of sequence divergence at *ebony* have not been demonstrated. Here, we show that divergence at *ebony* does indeed contribute to pigmentation divergence between these two species. We use CRISPR/Cas9 genome editing to mutate *ebony* in both species and use these mutant genotypes to directly test *ebony*'s contribution to pigmentation divergence through reciprocal hemizyosity testing (Stern, 2014). We find that the *D. novamexicana* *ebony* allele causes lighter pigmentation throughout the body than the *D. americana* *ebony* allele. We also find that allelic divergence at *ebony* is primarily responsible for a spatial difference in abdominal pigmentation between these species: the *D. novamexicana* *ebony* allele causes the absence of dark melanin along the dorsal midline of the abdomen seen in *D. novamexicana*. Finally, we show that *ebony* affects the cuticular hydrocarbon (CHC) profiles in *D. americana* and *D. novamexicana*, but does not contribute to the qualitative differences in CHC profiles seen between species. Taken together, our data show the power of using CRISPR/Cas9 genome editing to test functional hypotheses about evolutionary mechanisms. In addition, resources generated and lessons learned in the course of this work are expected to help other researchers perform CRISPR/Cas9 genome editing in *D. americana*, *D. novamexicana* and other *Drosophila* species.

MATERIALS AND METHODS

Fly Stocks and Husbandry

The following fly lines were used in this study: *D. americana* "A00" (National *Drosophila* Species Stock Center number 15010-0951.00), *D. novamexicana* "N14" (National *Drosophila* Species Stock Center number 15010-1031.14), *D. lummei*



(National Drosophila Species Stock Center number 15010-1011.08), *D. virilis* (National Drosophila Species Stock Center number 15010-1051.87), *D. melanogaster* $y^1 M\{w[+mC] = nos-Cas9.P\}ZH-2A w^*$ (Bloomington Drosophila Stock Center number 54591), and *D. melanogaster* Canton-S. All flies were reared on standard cornmeal medium at 23–25°C with a 12:12 hour light:dark cycle.

Transgenesis and CRISPR Mutant Generation in *D. americana* and *D. novamexicana*

To the best of our knowledge, prior to this work, the only transformation of *D. americana* or *D. novamexicana* resulted from the insertion of a piggyBac transgene (Wittkopp et al., 2009).

We therefore first used the CRISPR/Cas9 system to generate *white* mutants in both species to test the feasibility of CRISPR genome modification and to create lines that are easier to screen for common transformation markers that drive expression of fluorescent proteins or restore red pigmentation in the eyes by restoring *white* function. We successfully generated *white* mutant N14 and A00 lines used as transgenic hosts for future work, by injecting single guide RNAs (sgRNAs) targeting coding sequences in *white* conserved between *D. novamexicana* and *D. americana* in the second and third exons and screening for the loss of red eye pigment in male offspring of injected females (**Supplementary Figure 1**); *white* is on the X chromosome and thus only present in a single copy in males. These same guide RNAs were also used in *Drosophila virilis* to cut *white* and integrate an attP landing site potentially useful for site-directed transgene insertion (Lachowiec and Wittkopp, unpublished data), although the PhiC31 system does not seem to work well in *D. virilis* (Stern et al., 2017). For all CRISPR experiments, sgRNAs were *in vitro* transcribed from DNA templates using Invitrogen T7 MEGAscript Transcription Kit according to protocol described by Bassett et al. (2013). Oligonucleotides used to generate sgRNAs are listed in **Supplementary Table 1**. After transcription, sgRNAs were purified using RNA Clean and Concentrator 5 kit (Zymo Research), eluted with nuclease-free water, and quantified with Qubit RNA BR Assay Kit (Thermo Fisher Scientific). For CRISPR injections, sgRNAs were mixed with purified Cas9 protein (PNA Bio #CP01) with a final injected concentration of 0.05% phenol red to visualize the injection mix. CRISPR injections were performed in-house, using previously described methods (Miller et al., 2002).

To try to increase efficiency of CRISPR mutagenesis in these species, we next sought to generate transgenic lines expressing Cas9 in the germlines of *white* mutant *D. americana* (A00) and *D. novamexicana* (N14) flies using piggyBac transgenesis (Horn and Wimmer, 2000). Based on prior reports that the *nanos* (*nos*) promoter and 3'UTR drive expression in the germline of *Drosophila virilis* (Holtzman et al., 2010), a close relative of *D. americana* and *D. novamexicana*, we amplified the *nos-Cas9-nos* transgene from the pnos-Cas9-nos plasmid (Addgene #62208; Port et al., 2014) using Phusion High Fidelity Polymerase (NEB) with tailed primers and cloned the amplicon into pBac{3XP3-ECFPafm} (Horn and Wimmer, 2000) digested with AscI and Bsu36I restriction enzymes using Gibson Assembly Master Mix (NEB). Primers are included in **Supplementary Table 1**. We confirmed the insert was correctly incorporated and free of PCR-induced errors by Sanger sequencing. We sent the *white* mutant lines of *D. americana* A00 and *D. novamexicana* N14 that we generated to Rainbow transgenic services for piggyBac transgenesis¹ and screened offspring of injected adults for expression of the enhanced cyan fluorescent protein (ECFP) in the eye using a Leica MZ6 stereoscope equipped with a Kramer Scientific Quad Fluorescence Illuminator. Transformants were obtained from injections into *D. novamexicana* (N14) (PCR verified), but not from injections into *D. americana* (A00), despite multiple attempts.

¹www.rainbowgene.com

All subsequent CRISPR injections in *D. novamexicana* were performed using flies homozygous for the *nos-Cas9-nos* transgene, some with and some without the inclusion of commercially available Cas9 protein in the injection mix. CRISPR mutants were only obtained from injections containing the commercially available Cas9 protein, however, suggesting that the *nos-Cas9-nos* transgene might not drive expression of Cas9 in the germline of *D. novamexicana*. To test this hypothesis, we used western blotting to examine Cas9 protein expression in 3 transformed *D. novamexicana* N14 lines with independent insertions of the piggyBac transgene and in a *D. melanogaster* transgenic line carrying the original *pnos-Cas9-nos* transgene (Bloomington Drosophila Stock Center line 54591, transformed with Addgene plasmid #62208, Port et al., 2014). These experiments showed that the *nos-Cas9-nos* transgene in *D. novamexicana* N14 flies does not express Cas9 protein in the ovaries (**Supplementary Figure 2**). This conclusion was further supported when injection of sgRNAs targeting the *yellow* gene into the *D. novamexicana* line carrying the *nos-Cas9-nos* transgene also only produced *yellow* mutants when the Cas9 protein was co-injected with the sgRNAs (**Supplementary Figure 3**). Ability of the *nos* promoter to drive germline expression in the closely related species *D. virilis* has also been found to be variable among transgenic lines (Hannah McConnell, Aida de la Cruz, and Harmit Malik, personal communication), suggesting that other promoters should be used in the future to drive reliable germline expression in the virilis group.

To generate *ebony* mutant *D. americana* (A00) and *D. novamexicana* (N14), we synthesized five sgRNAs targeting conserved sites in the first coding exon of *ebony*. Because *ebony* is located on an autosome and *ebony* loss-of-function mutant alleles are generally considered recessive in *D. melanogaster* (Thurmond et al., 2019), we did not expect to be able to identify *ebony* mutants by simply screening progeny of injected flies for mutant phenotypes as we did for *white* and *yellow*. We therefore co-injected a donor plasmid containing the sequence of an eye-specific red fluorescent protein marker (3XP3-RFP) flanked by *ebony* sequences that could be inserted into *ebony* via homology-directed repair and used to screen for *ebony* mutants. Although we observed RFP expression in larvae injected with the homology-directed repair donor fragment, indicating that the reporter gene was functional in these species, injected individuals did not produce any offspring with red fluorescent eyes, suggesting that the donor plasmid was not integrated in the germline of injected individuals. Because non-homologous end joining occurs more frequently than homology directed repair following double-strand breaks (Liu et al., 2018), we also tried to identify flies that might be heterozygous for an *ebony* mutant allele by closely inspecting all offspring of injected (G_0) flies for any subtle changes in pigmentation. Specifically, we collected and mated (G_1) offspring of injected flies with any noticeably darker pigmentation, keeping them grouped by G_0 parent of origin. As further described in the results, we were ultimately able to identify homozygous *ebony* mutants among progeny from these $G_1 \times G_1$ crosses of relatively dark flies derived from two independent

D. novamexicana G_0 flies and one *D. americana* G_0 fly. Sanger sequencing these flies confirmed they were homozygous for *ebony* alleles containing deletions. We then crossed the mutated *ebony* alleles back into wild-type backgrounds of each parental species to generate homozygous *ebony* mutant lines with wild-type red eyes.

Western Blotting

For *ebony* western blotting, proteins were extracted from stage P14/15 pupae, identified by the following characteristics: black pigmentation present in wings and bristles, meconium visible in abdomen (Cooley et al., 2012). For each sample, five pupae were homogenized in 100 μ L of homogenization buffer (125 mM Tris pH 6.8, 6% SDS, 2.5X Roche cOmplete protease inhibitor cocktail, EDTA-free), then centrifuged for 15 min at 15000 rcf, and the supernatant transferred to a fresh tube with an equal volume of 2x Laemmli buffer (125 mM Tris pH 6.8, 6% SDS, 0.2% glycerol, 0.25% bromophenol blue, 5% Beta-mercaptoethanol).

For Cas9 western blotting, protein was extracted from ovaries dissected in ice cold PBS from the following lines: untransformed N14 *white* mutants (host line), three independently transformed lines of N14 *white* carrying the pBac{3XP3-ECFPafm-nosCas9nos} transgene, transgenic *D. melanogaster* carrying the *pnos-Cas9-nos* transgene, and wild-type (Canton-S) *D. melanogaster*. For *D. novamexicana* samples, we collected ovaries from 10 sexually mature flies, whereas for *D. melanogaster* samples, we collected ovaries from 18 sexually mature flies. Different numbers of flies were used for the two species because of differences in body size. In each case, ovaries were placed into microcentrifuge tubes on ice, spun down briefly in a tabletop centrifuge, and excess PBS was removed and replaced with 20 μ L of homogenization buffer. Samples were then treated as described for *ebony* western blots above. A positive control Cas9 sample was made by diluting purified Cas9 protein (PNA Bio CP01) in homogenization buffer, and mixing with 2X Laemmli buffer to a final concentration of 2.5 ng/ μ L.

Samples were heated at 95°C for 10 min before loading into 7.5% Mini-PROTEAN® TGX™ Precast Protein Gels (Bio-Rad) and running at 150V for approximately 90 min at 4°C in 1X tris-glycine running buffer. Separate gels were run for *ebony* and Cas9 blots. Samples were loaded in the following volumes: 35 μ L per pupa sample, 30 μ L per ovary sample, 10 μ L of Cas9 positive control (25 ng protein), 5 μ L PageRuler prestained protein ladder (Thermo Fisher Scientific). Gels were transferred onto PVDF membrane in tris-glycine transfer buffer, 10% MeOH, 0.01% SDS at 100 V for 1 h with stirring on ice at 4°C. Membranes were blocked in 3% nonfat dry milk in TBST for 30 min at RT with shaking, then divided in half using the prestained ladder as a guide just below the 100 kDa mark for the Cas9 membrane and just below the 70 kDa mark for the *ebony* membrane. The lower molecular weight halves of the membranes were placed in solutions containing primary antibodies to detect the protein used as a loading control (tubulin or lamin), whereas the halves of the membranes containing the higher molecular weight proteins were placed in solutions containing primary antibody solutions against the protein of

interest (Ebony or Cas9), each diluted in 3% nonfat dry milk in TBST. In all cases, membranes were incubated with the primary antibodies overnight at 4°C. Primary antibody solutions for *ebony* included rabbit anti-*ebony* 1:300 (Wittkopp et al., 2002) and rabbit anti-alpha tubulin 1:5000 (Abcam ab52866) as a loading control. Primary antibody solutions for Cas9 included mouse anti-Cas9 1:1000 (Novus NBP2-36440) and mouse anti-lamin 1:200 (DHSB adl67.10) as a loading control. Membranes were washed in TBST and transferred to secondary antibody solutions diluted in 3% nonfat milk in TBST for 2 h at RT. The following secondary antibodies were used: donkey anti-rabbit HRP 1:5000 (Amersham na934) or goat anti-mouse HRP 1:5000 (abcam ab97023). Membranes were finally washed in TBST and developed with SuperSignal West Pico Chemiluminescent Substrate (Thermo Fisher Scientific) and imaged using a Licor Odyssey FC imaging system.

Fly Crosses for Reciprocal Hemizygoty Testing and Cuticular Hydrocarbon Analysis

To generate F₁ hybrids carrying only one (*D. americana* or *D. novamexicana*) functional *ebony* allele, wild-type and *ebony* mutant flies from each species were collected as virgins and aged in vials for at least 12 days to reach sexual maturity and verify virgin female status by absence of larvae. Crosses were all set on the same batch of food on the same day and placed at 25°C. For most crosses, 4 virgin females and 4 males were used; however, 8 virgin females and 8 males were used in interspecific crosses with *D. novamexicana* females because of reduced mating success in these crosses. After 3 days, adult flies from these crosses that would be used for cuticular hydrocarbon (CHC) analysis were transferred to new vials with a fresh batch of food. Offspring from the first set of vials were used for imaging and pigmentation analysis, while offspring from the second set of vials were used for CHC analysis. Flies used for pigmentation phenotyping were aged 5–7 days after eclosion and preserved in 10% glycerol in ethanol before imaging (Wittkopp et al., 2011).

Imaging of Fly Phenotypes

Insect specimens were imaged using a Leica DC480 camera attached to a Leica MZ16F stereoscope equipped with a ring light attachment and Leica KL 1500 LCD lamp. Images were captured using Leica DC Twain software version 5.1.1 run through Adobe Photoshop CS6 version 13.0 X32. Prior to imaging, pupal cases and wings were mounted on slides in PVA mounting medium (BioQuip). Thorax, abdomen, and whole-body specimens were prepared from age-matched, preserved flies as described in the previous section. For imaging, thorax, abdomen, and whole-body specimens were submerged in 100% ethanol in custom wells composed of white oven-cured polymer clay (Sculpey).

Because the color of specimens spanned a wide range across genotypes, exposure was optimized for each sample type (e.g., whole body, thorax, abdomen, wing, pupal case) individually by placing specimens from the two phenotypic extremes in the same frame and adjusting exposure to avoid

over-exposing the lightest flies while capturing as much detail as possible from the darkest flies. Exposure time, lighting, white balance, background, and zoom were kept identical across all images of single tissue type. Minor color adjustments to improve visibility of phenotypes were performed simultaneously across all raw images of the same sample type in a single combined document using Photoshop CC 2019, ensuring that all images presented for direct comparisons were adjusted identically.

Cuticular Hydrocarbon Analyses

CHCs for each cross were extracted from 5-day-old females by soaking the flies for 10 min in 200 µl hexane containing hexacosane (C₂₆; 25 ng/ul) as an internal standard. Eight replicates were prepared for each cross. Extracts were directly analyzed by the GC/MS (7890A, Agilent Technologies Inc., Santa Clara, CA, United States) coupled with a DB-17ht column 30 m by 0.25 mm (i.d.) with a 0.15 µm film thickness (Agilent Technologies Inc., Santa Clara, CA, United States). Mass spectra were acquired in Electron Ionization (EI) mode (70 eV) with Total Ion Mode (TIM) using the GC/MS (5975C, Agilent Technologies Inc., Santa Clara, CA, United States). The peak areas were recorded by MassHunter software (Agilent Technologies Inc., Santa Clara, CA, United States). Helium was the carrier gas at 0.7 ml/min and the GC thermal program was set as follows: 100°C for 4 min, 3°C/min to 325°C. Straight-chain compounds were identified by comparing retention times and mass spectra with authentic standard mixture (C₆-C₄₀) (Supelco® 49452-U, Sigma-Aldrich, St. Louis, MO, United States). Methyl-branched alkanes, alkenes, dienes and trienes were then identified by a combination of their specific fragment ions on the side of functional groups (methyl branch or double bonds) and retention times relative to linear-chain hydrocarbon standards. Each individual CHC peak was quantified by normalizing its peak area to the peak area of the internal C₂₆ standard, converting each CHC peak area to ng/fly using the known internal standard concentration of 1000 ng/fly. Welch's *t*-tests with a Benjamini–Hochberg correction for multiple testing (Benjamini and Hochberg, 1995) were used to compare CHC amounts between pairs of genotypes. Because the effect of *ebony* on individual CHC abundance in *D. melanogaster* was recently shown to increase with CHC chain length (Massey et al., 2019b), we also compared the effects of *ebony* loss of function on different chain-lengths of CHCs. Eight biological replicates of homozygous *ebony* null measurements were divided by the mean measurement of the eight replicates of the matched *ebony* heterozygote for each individual CHC. The ratio of *ebony* null to heterozygote CHC abundance was plotted against CHC chain length. The relative effects of *D. americana* versus *D. novamexicana* *ebony* in a common F₁ hybrid background (described as F₁[*e*^A/*e*⁻] and F₁[*e*^N/*e*⁻], respectively), were also compared in this manner, with the replicates of the F₁[*e*^A/*e*⁻] divided by the mean F₁[*e*^N/*e*⁻] measurement for each CHC. We used Spearman's rank correlation (Spearman's rho) to test the relationship between CHC chain length and the effect of *ebony* on CHC abundance. The threshold for statistical significance was set at *alpha* = 0.05

for all tests. Datafile and R code used for this analysis are provided in **Supplementary Files 1, 2**, respectively.

RESULTS AND DISCUSSION

The reciprocal hemizyosity test is a powerful strategy for identifying genes with functional differences that contribute to phenotypic divergence [reviewed in Stern (2014)]. This test is performed by comparing the phenotypes of two hybrid genotypes that are genetically identical except for which allele of the candidate gene is mutated. Any phenotypic differences observed between these two genotypes are attributed to divergence of the candidate gene. Applying this test to identify functional differences between species requires loss-of-function (null) mutant alleles in both species and the ability for the species to cross and produce F₁ hybrids. Consequently, in order to use this strategy to test *ebony* for functional divergence between *D. novamexicana* and *D. americana*, we first needed to generate *ebony* null mutant alleles in both species.

Generating *ebony* Mutants in *D. americana* and *D. novamexicana* Using CRISPR/Cas9

We generated *ebony* null mutants in *D. novamexicana* and *D. americana* by using CRISPR/Cas9 to target double-strand breaks to five conserved sites within the first coding exon of *ebony*. As described more fully in the “Materials and Methods” section, we injected embryos of *white* mutants from both species with purified Cas9 protein and sgRNAs targeting all five sites simultaneously. BLAST searches showed that all of the sgRNAs targets were at least 5 bp different from all other sequences in genomes from two different strains of *D. americana*. Prior work in *D. melanogaster* has shown that heritable off-target mutations were never recovered in sequences with 3 or more mismatches to the sgRNA (Ren et al., 2014). To make it easier to identify *ebony* mutant alleles, we also injected a donor plasmid that would allow homology directed repair to integrate a transgene expressing red fluorescent protein in the fly’s eyes, but no progeny of injected flies were observed to express this transformation marker. However, we reasoned that although we were unable to insert a marker at *ebony*, the CRISPR machinery may still have induced double-strand breaks in the target sequence, and *ebony* mutants could have been generated by non-homologous end-joining resulting in deletions or insertions. Therefore, we also searched for *ebony* mutants by looking for changes in body pigmentation.

In *D. melanogaster*, *ebony* loss-of-function mutants have a much darker appearance than wild-type flies because they are unable to produce yellow sclerotin, causing an increase in production of black and brown melanins (Wittkopp et al., 2002). *D. melanogaster* *ebony* mutant alleles are commonly described as recessive to wild-type *ebony* alleles (Thurmond et al., 2019); however, in some genetic backgrounds, flies heterozygous for an *ebony* mutant allele are slightly darker than wild-type flies (Thurmond et al., 2019). Because *D. novamexicana* has such a light yellow body color (Figure 2A), we thought it possible

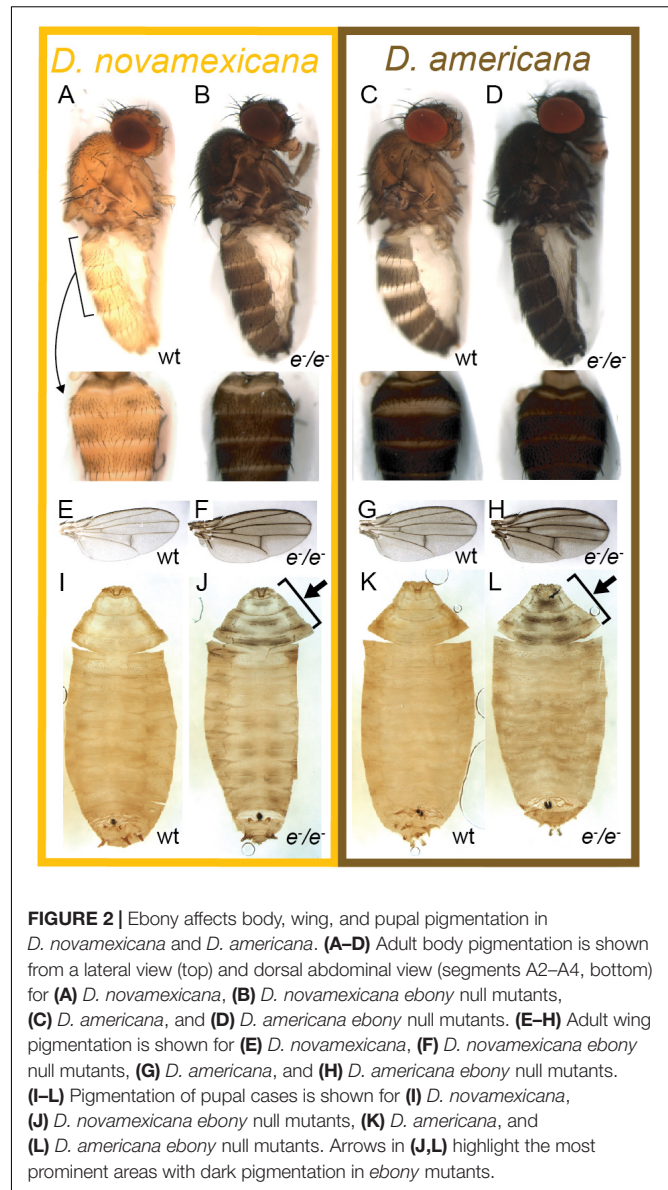


FIGURE 2 | Ebony affects body, wing, and pupal pigmentation in *D. novamexicana* and *D. americana*. (A–D) Adult body pigmentation is shown from a lateral view (top) and dorsal abdominal view (segments A2–A4, bottom) for (A) *D. novamexicana*, (B) *D. novamexicana* *ebony* null mutants, (C) *D. americana*, and (D) *D. americana* *ebony* null mutants. (E–H) Adult wing pigmentation is shown for (E) *D. novamexicana*, (F) *D. novamexicana* *ebony* null mutants, (G) *D. americana*, and (H) *D. americana* *ebony* null mutants. (I–L) Pigmentation of pupal cases is shown for (I) *D. novamexicana*, (J) *D. novamexicana* *ebony* null mutants, (K) *D. americana*, and (L) *D. americana* *ebony* null mutants. Arrows in (J,L) highlight the most prominent areas with dark pigmentation in *ebony* mutants.

that flies heterozygous for an *ebony* mutant allele might also show a detectable darkening of pigmentation; we were less optimistic about being able to detect heterozygous *ebony* mutants based on pigmentation in *D. americana* because its wild-type pigmentation is already very dark (Figure 2C). Nonetheless, we sorted through the progeny of injected *D. novamexicana* and *D. americana* flies, isolating any individuals that seemed to have darker pigmentation than their siblings and allowing these relatively dark flies to freely mate in vials segregated by injected parents, keeping individual “founder” mutations separate.

Two of the vials of darker pigmented *D. novamexicana* flies produced pupae with an unusual black pattern on the anterior end of the pupal case (Figure 2J). We moved these pupae to new vials and found that black-patterned pupae from both “founder” vials developed into adults with the much darker than wild-type body color expected for

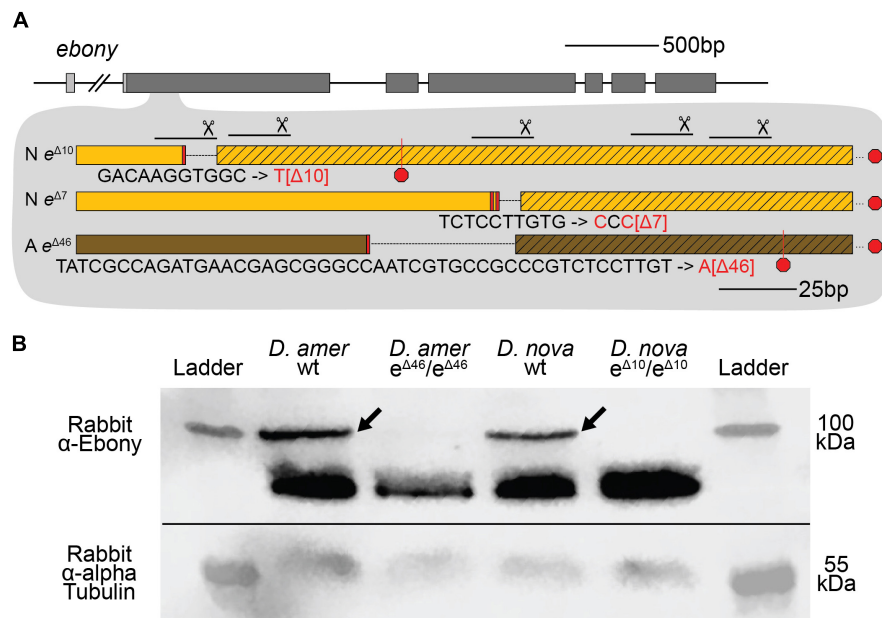


FIGURE 3 | CRISPR/Cas9-induced mutations created null alleles of the *D. novamexicana* and *D. americana* *ebony* genes. **(A)** A schematic of the *ebony* gene is shown with gray boxes indicating exons; coding sequence is indicated in the darker shade of gray. Locations of the five guide RNAs targeting the second exon of *ebony* are shown with solid lines below scissor symbols. Mutations observed in the two *ebony* mutants ($e^{\Delta 10}$ and $e^{\Delta 7}$) isolated in *D. novamexicana* ("N") and the one *ebony* mutant ($e^{\Delta 46}$) isolated in *D. americana* ("A") are shown. All three alleles included deletions that caused frameshifts. **(B)** Western blotting showed that the *D. americana* $e^{\Delta 46}$ and *D. novamexicana* $e^{\Delta 10}$ mutants (lanes 2 and 4, respectively), lacked a ~100 kDa protein (arrows) recognized by an antibody raised against *D. melanogaster* Ebony protein (Wittkopp et al., 2002) that is present in wild-type (wt) *D. americana* and *D. novamexicana* (lanes 1 and 3, respectively). Relative abundance of total protein loaded into each lane can be seen by the relative intensities of the shorter proteins also detected by the Ebony antibody (Wittkopp et al., 2002) as well as the relative intensities of ~55 kDa bands detected by an antibody recognizing alpha Tubulin (Abcam ab52866). The solid black line shows where the membrane was cut prior to incubation with primary antibodies during the western blotting procedure; the top half was incubated with anti-Ebony antibodies whereas the bottom half was incubated with anti-Tubulin antibodies. The two halves were realigned by hand for imaging, using the shape of the cut and the ladder staining as a guide. An un-annotated image of this blot is shown in **Supplementary Figure 4**.

homozygous *ebony* mutants in *D. novamexicana* (Figures 2A,B). Because pigmentation of the pupal case is very similar between *D. novamexicana* and *D. americana* (Figures 2I,K, Ahmed-Braimah and Sweigart, 2015), we also searched for pupae with similar pigmentation marks in the vials containing progeny of darker flies descended from injected *D. americana*. We found such pupae in only one of the *D. americana* vials (Figure 2L). Flies emerging from these pupal cases also showed darker pigmentation than wild-type *D. americana* (Figures 2C,D), as expected for homozygous *ebony* mutants, but this difference was much more subtle than in *D. novamexicana* (Figures 2A,B). Flies from both species emerging from pupal cases with abnormal pigmentation also showed increased levels of dark melanins in wings in a pattern similar to that seen in *D. melanogaster* *ebony* mutants (Figures 2E–H, Wittkopp et al., 2002), further suggesting that they were homozygous for *ebony* mutant alleles. Crossing putative homozygous *ebony* mutants from the same species to each other resulted in true-breeding lines of *D. americana* and *D. novamexicana* presumed to be homozygous for *ebony* mutant alleles.

To determine whether these true-breeding lines were indeed homozygous for *ebony* mutant alleles, we used Sanger sequencing to search for changes in the *ebony* sequence in the region

targeted for double strand breaks with CRISPR/Cas9. We found that the presumed *ebony* mutant lines of both species harbored deletions corresponding to the locations of sgRNA target sites in the first coding exon, with the two *D. novamexicana* mutant lines carrying deletions of 7 and 10 bases and the *D. americana* mutant line carrying a deletion of 46 bases (Figure 3A). Each of these mutations is expected to cause frameshifts, leading to multiple early stop codons. Further experiments described in this study using *D. novamexicana* *ebony* mutants were conducted with the 10 base deletion line, and any further description of *ebony* null *D. novamexicana* refers to this line.

To further assess whether these mutations caused null alleles, we used western blotting to examine the expression of the Ebony protein during late pupal stages when adult pigmentation is developing and the *ebony* gene is expressed in the developing abdomen (Wittkopp et al., 2002; Cooley et al., 2012). We performed western blots on protein extracts from P14/P15 stage pupae of both wild-type and homozygous *ebony* mutant flies of both *D. americana* and *D. novamexicana* using an antibody against *D. melanogaster* *ebony* (Wittkopp et al., 2002). This antibody recognizes a 94 kDa protein consistent with the predicted molecular weight of Ebony in pupal protein extracts from wild-type lines of both *Drosophila*

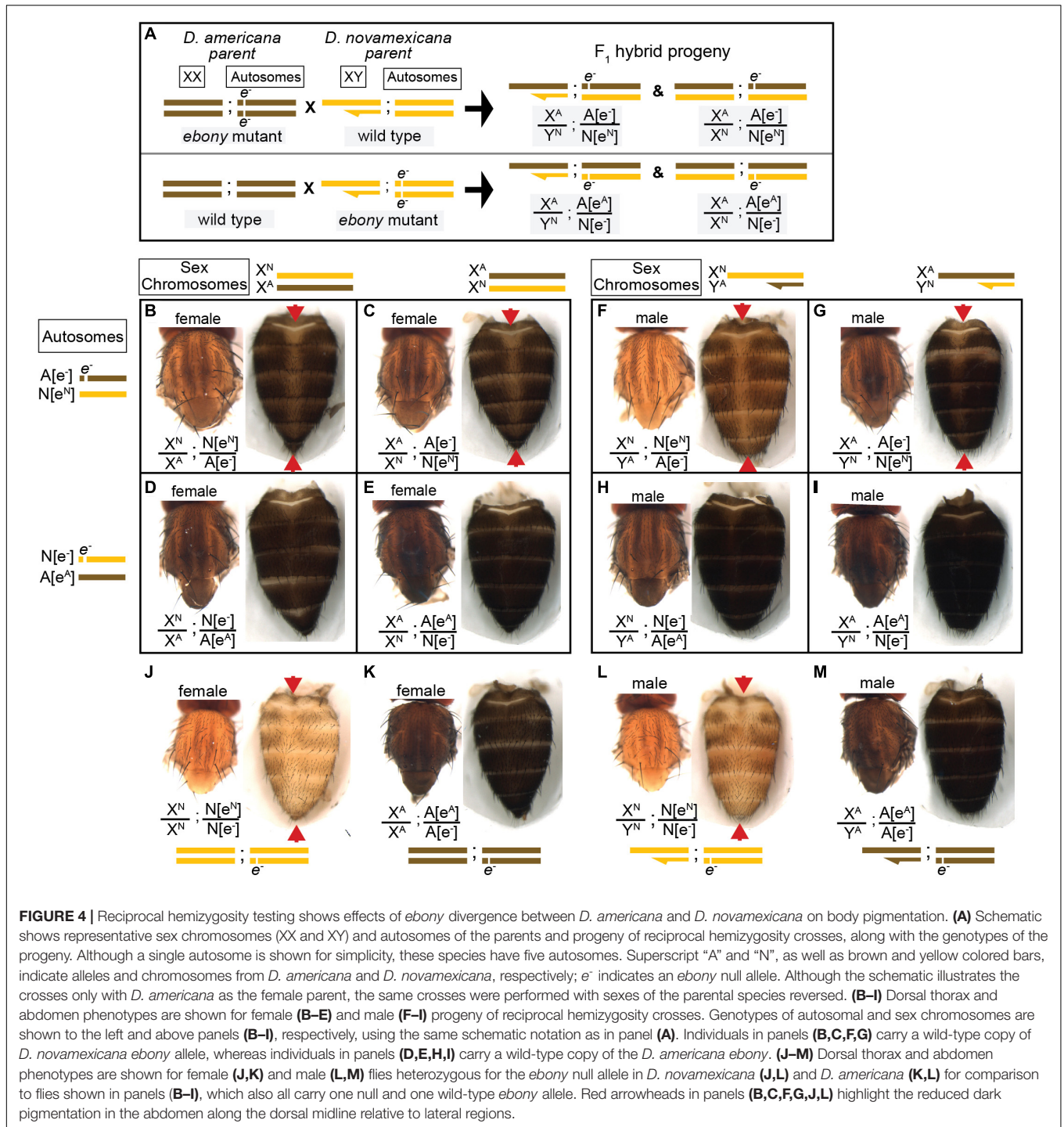


FIGURE 4 | Reciprocal hemizyosity testing shows effects of *ebony* divergence between *D. americana* and *D. novamexicana* on body pigmentation. **(A)** Schematic shows representative sex chromosomes (XX and XY) and autosomes of the parents and progeny of reciprocal hemizyosity crosses, along with the genotypes of the progeny. Although a single autosome is shown for simplicity, these species have five autosomes. Superscript "A" and "N", as well as brown and yellow colored bars, indicate alleles and chromosomes from *D. americana* and *D. novamexicana*, respectively; e⁻ indicates an *ebony* null allele. Although the schematic illustrates the crosses only with *D. americana* as the female parent, the same crosses were performed with sexes of the parental species reversed. **(B–I)** Dorsal thorax and abdomen phenotypes are shown for female **(B–E)** and male **(F–I)** progeny of reciprocal hemizyosity crosses. Genotypes of autosomal and sex chromosomes are shown to the left and above panels **(B–I)**, respectively, using the same schematic notation as in panel **(A)**. Individuals in panels **(B,C,F,G)** carry a wild-type copy of *D. novamexicana* *ebony* allele, whereas individuals in panels **(D,E,H,I)** carry a wild-type copy of the *D. americana* *ebony*. **(J–M)** Dorsal thorax and abdomen phenotypes are shown for female **(J,K)** and male **(L,M)** flies heterozygous for the *ebony* null allele in *D. novamexicana* **(J,L)** and *D. americana* **(K,L)** for comparison to flies shown in panels **(B–I)**, which also all carry one null and one wild-type *ebony* allele. Red arrowheads in panels **(B,C,F,G,J,L)** highlight the reduced dark pigmentation in the abdomen along the dorsal midline relative to lateral regions.

melanogaster and *Drosophila biarmipes*, but does not produce a 94 kDa band in pupal protein extracts of either *e1* or *In(3R)eAFA* *ebony* mutant lines of *D. melanogaster* (Wittkopp et al., 2002). Wild-type extracts of both *D. americana* and *D. novamexicana* produced presumptive Ebony bands while extracts from flies homozygous for *ebony* deletions did not produce a 94 kDa band for either species (**Figure 3B**). The nature of the frameshift deletions as well as the western

blot evidence together show that these *ebony* mutations cause null alleles.

ebony* Divergence Contributes to Body Color Differences Between *D. novamexicana* and *D. americana

We used the homozygous *ebony* mutant *D. novamexicana* and *D. americana* lines to perform a reciprocal hemizyosity test

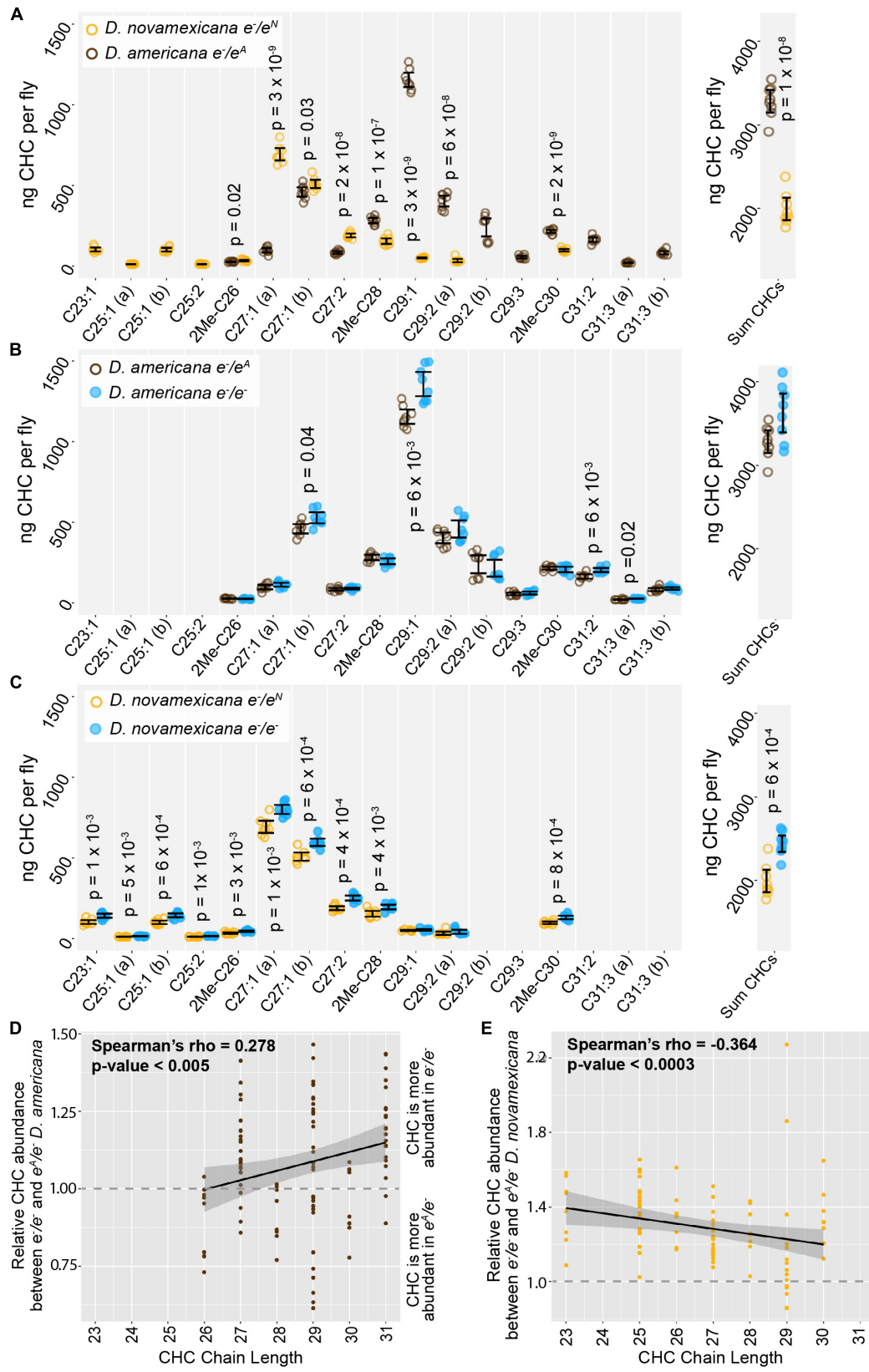


FIGURE 5 | Continued

FIGURE 5 | Cuticular hydrocarbons (CHCs) are affected by *ebony* and differ between *D. americana* and *D. novamexicana*. **(A–C)** Abundance of individual CHC compounds (ng/fly) and summed CHCs extracted from female flies are plotted for the following genotypes: **(A)** *D. americana* and *D. novamexicana*, each heterozygous for an *ebony* null (e^-) allele, **(B)** *D. americana* heterozygous and homozygous for an *ebony* null allele, **(C)** *D. novamexicana* heterozygous and homozygous for an *ebony* null allele. Eight biological replicates are shown for each genotype, with error bars representing 95% confidence intervals. For each comparison, the p -value from a Welch's t -test with a Benjamini–Hochberg multiple test correction ($\alpha = 0.05$) is shown when a significant difference in abundance was detected for a CHC present in both genotypes being compared. CHCs are shown from left to right with increasing chain length (represented by “C” followed by the chain length) with double-bond and methyl-branched structures indicated by notations after the colon or before the “C”, respectively. For example, C25:1 represents a 25-carbon monoene, C25:2 represents a 25-carbon diene, and 2Me-C28 represents a 28-carbon alkene with a methyl branch at the second carbon. **(D–E)** Abundance of each CHC in *ebony* null mutants relative to flies heterozygous for the *ebony* null allele is plotted by carbon chain length for **(D)** *D. americana* and **(E)** *D. novamexicana*. Black trendlines in panels **(D–E)** show linear regressions, with shaded areas representing the standard error and both Spearman's rho and p -values indicated on each plot.

by crossing *ebony* mutant *D. novamexicana* (e^-/e^-) to wild-type *D. americana* (e^A/e^A) and *ebony* mutant *D. americana* (e^-/e^-) to wild-type *D. novamexicana* (e^N/e^N) (**Figure 4A**). In order to observe the effects of the two species' *ebony* alleles in the presence of each species X chromosome, we conducted sets of reciprocal crosses (i.e., swapping the genotypes of the male and female parents). Female F_1 hybrids from reciprocal crosses are genetically identical except for the parent of origin of their one functional *ebony* allele (e^N or e^A). F_1 hybrid females carrying a functional *D. novamexicana* *ebony* allele ($F_1[e^N/e^-]$) developed a lighter body color than F_1 hybrid females carrying a functional *D. americana* *ebony* allele ($F_1[e^A/e^-]$) (**Figures 4B,C** vs. **Figures 4D,E**). These data demonstrate for the first time that functional divergence between the *D. novamexicana* and *D. americana* *ebony* alleles contributes to divergent body color between these two species.

To determine how *ebony* divergence interacts with divergent loci on the X-chromosome, we also compared the body color of male progeny from these reciprocal crosses. Like the F_1 hybrid females, these F_1 hybrid males differ for the parent of origin for their one functional *ebony* allele (e^A or e^N); however, they also differ for the parent of origin of all X-linked genes. Prior work has shown that divergence on the X-chromosome, particularly divergence in non-coding sequences of the *tan* gene, also contributes to differences in body color between *D. novamexicana* and *D. americana* (Wittkopp et al., 2003b, 2009). As expected, we found that body color differed between males carrying alternate species' X chromosomes (**Figure 4F** vs. **Figure 4G** and **Figure 4H** vs. **Figure 4I**) as well as between males carrying the same X chromosome but different species' functional *ebony* alleles (**Figure 4F** vs. **Figure 4H** and **Figure 4G** vs. **Figure 4I**). Consistent with prior findings demonstrating that divergence in the QTL containing *ebony* explained more of the difference in pigmentation than divergence at X-linked genes, we found that males with functional *D. americana* *ebony* alleles had the darkest phenotypes, regardless of their X-chromosome genotype (**Figures 4F–I**).

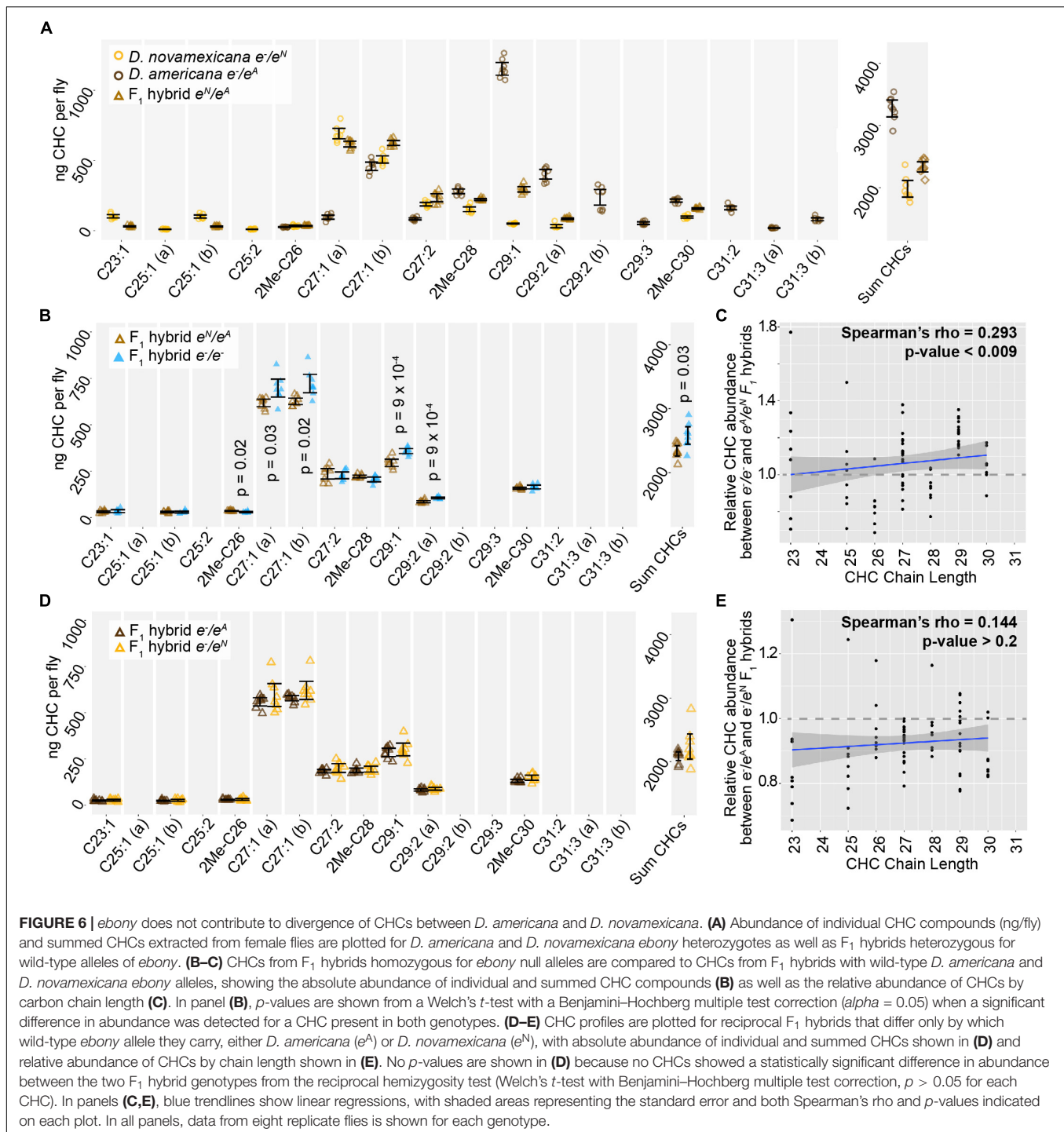
ebony* Divergence Also Contributes to a Difference in Abdominal Pigment Patterning Between *D. novamexicana* and *D. americana

Although the divergent overall body color is the most striking difference in pigmentation between *D. novamexicana* and

D. americana, there is also a difference in the distribution of pigments along the dorsal midline of the abdomen between these two species (**Figure 1**). This difference is also visible in individuals of both species heterozygous for an *ebony* null allele (**Figures 4J–M**). Prior work has shown that the absence of dark pigments seen in this region of *D. novamexicana* is dominant in F_1 hybrids to the presence of dark pigments seen in this region of *D. americana* (Wittkopp et al., 2003b). In addition, genetic mapping of this trait between *D. novamexicana* and *D. virilis* (which has a dark midline region similar to *D. americana*) has shown that the chromosome including *ebony* (chromosome 2) has a large effect on this trait (Spicer, 1991). We found that *D. novamexicana* *ebony* mutants showed even pigmentation across the width of each abdominal segment (**Figure 2B**), demonstrating that *ebony* is required for the development of lighter pigmentation along the dorsal midline in wild-type *D. novamexicana* (**Figure 2A**). In addition, comparing the pigmentation of this abdominal dorsal midline region between F_1 hybrid flies of both sexes from the reciprocal crosses described above (**Figure 4**) showed that divergence at *ebony* contributes to this trait difference between *D. novamexicana* and *D. americana*. Specifically, we observed less dark pigments in the dorsal midline region of the abdomen in F_1 hybrid individuals inheriting the wild-type *D. novamexicana* *ebony* allele ($F_1[e^N/e^-]$, **Figures 4B,C,F,G**) than the *D. americana* *ebony* allele ($F_1[e^A/e^-]$) (**Figures 4D,E,H,I**). Males carrying a functional *D. novamexicana* *ebony* allele ($F_1[e^N/e^-]$) showed reduced pigmentation in the dorsal midline relative to the lateral regions regardless of the origin of their X chromosome (**Figures 4F,G**), indicating that divergent loci on the X-chromosome (including *tan*) do not affect the presence of this phenotype.

Cuticular Hydrocarbon Profiles Differ Between *D. americana* and *D. novamexicana* and Are Affected by *ebony* Expression but Not Ebony Divergence

ebony expression was recently found to affect the relative abundance of cuticular hydrocarbons (CHCs) in *D. melanogaster* (Massey et al., 2019b). In addition, variation in *ebony* expression was also shown to correlate with variation in CHC profiles among natural isolates of *D. melanogaster* (Massey et al., 2019b). CHC profiles have been shown to vary among virilis group species as well, including between *D. novamexicana* and



the strain of *D. americana* used in this study (Bartelt et al., 1986). We therefore asked whether differences in *ebony* might contribute to differences in CHC profiles between these two species using a reciprocal hemizyosity test. Because this test compares phenotypes of reciprocal hemizygotes that each carry a single functional copy of *ebony*, we also examined CHC profiles in *D. americana* and *D. novamexicana* flies with a single functional copy of *ebony*. We found that *D. novamexicana*

flies hemizygous for *ebony* contained a distribution of CHCs biased toward shorter chain hydrocarbons relative to CHCs extracted from *D. americana* flies hemizygous for *ebony*. Prior work found similar profiles of CHCs for wild type *D. americana* and *D. novamexicana* (Bartelt et al., 1986): in both studies, CHCs with a chain length of 25 or fewer were only present in *D. novamexicana*, whereas multiple CHCs with a chain length greater than 30 were only present in *D. americana* (Figure 5A).

These data suggest that the wild-type *ebony* allele has a dominant effect on CHC profiles.

In order to test whether *ebony* affects CHCs in these species, we compared CHCs extracted from homozygous *ebony* mutants to those extracted from *ebony* heterozygotes. In both species, the loss of *ebony* function had no qualitative effect on which CHCs were produced by either species, but increased the abundance of some CHCs in both *D. americana* and *D. novamexicana* (Figures 5B,C). Because *ebony* loss-of-function mutants in *D. melanogaster* were recently shown to preferentially increase the abundance of long chain CHCs (Massey et al., 2019b), we compared relative abundance of individual CHCs between *ebony* null and heterozygous samples and plotted the results against CHC chain length (Figures 5D,E). We observed a similar pattern to *D. melanogaster* in *D. americana*, with *ebony* loss-of-function increasing the abundance of longer chain CHCs more strongly (Figure 5D). In *D. novamexicana*, we observed the opposite pattern, however: CHCs with shorter chain lengths showed greater increases in abundance in *ebony* null mutants (Figure 5E). The reason for this difference in how *ebony* affects CHCs in *D. americana* and *D. novamexicana* remains unclear, but might have to do with the different levels of *tan* expression in these two species (Cooley et al., 2012) given that *tan* was also shown to affect CHC profiles in *D. melanogaster* (Massey et al., 2019b). Specifically, *tan* expression is lower in *D. novamexicana* pupae relative to *D. americana* (Cooley et al., 2012), and *tan* loss-of-function has been shown to preferentially increase the abundance of shorter chain CHCs in *D. melanogaster* (Massey et al., 2019b). Further experiments exploring the mechanisms underlying CHC production in these species may provide more insight into the contrasting effects of *ebony* on CHCs in *D. americana* and *D. novamexicana*.

We also examined the CHC profiles of female F₁ hybrids produced by crossing *D. americana* females with *D. novamexicana* males. We found that these F₁ hybrid females showed a CHC profile that was distinct from both species, but more similar to *D. novamexicana* (Figure 6A): it contained some of the short chain CHCs unique to *D. novamexicana* and none of the long chain CHCs unique to *D. americana* (Figure 6A). As seen for both species, eliminating *ebony* function in F₁ hybrids by making them homozygous for *ebony* null alleles caused an increase in abundance of some CHCs but did not alter which CHCs were present (Figure 6B). Longer chain CHCs were more likely to show increased abundance than shorter chain CHCs (Figure 6C), but this relationship was not as strong as that seen for *D. americana* (Figure 5D). To determine whether divergence between the *D. americana* and *D. novamexicana* *ebony* alleles affected CHCs profiles, we compared CHCs extracted from females from the reciprocal hemizyosity test. These flies have only one functional *ebony* allele (*D. americana* or *D. novamexicana*) in the F₁ hybrid genetic background. The CHC profiles from these flies were not significantly different from each other (Figures 6D,E), indicating that allelic divergence at *ebony* does not have a detectable effect on CHCs in this species pair.

CONCLUSION

Identifying the genes responsible for phenotypic differences between species remains a significant challenge for evolutionary biology. This task is especially challenging when a gene contributing to phenotypic divergence is located in a region of the genome inverted between species, which precludes recombination-based mapping. Such is the case for the *ebony* gene in *D. americana* and *D. novamexicana*. Prior work suggested that *ebony* might contribute to differences in overall body color between these two species (Wittkopp et al., 2009; Cooley et al., 2012), but its location in an inversion made it difficult to directly test this hypothesis. In this study, we overcame this hurdle by using CRISPR/Cas9 genome editing to generate null mutants for *ebony* in *D. americana* and *D. novamexicana*, and then using these mutants to perform a reciprocal hemizyosity test (Stern, 2014), which directly compares the effects of the two species' alleles on pigmentation. We found that divergence at *ebony* does indeed contribute to differences in body color between *D. americana* and *D. novamexicana*.

Characterizing the phenotypes of *D. americana* and *D. novamexicana* *ebony* mutants, as well as flies from the reciprocal hemizyosity test, also identified effects of *ebony* on other phenotypes. For example, we found that differences in the activity of *ebony* alleles between *D. americana* and *D. novamexicana* are responsible for the absence of dark pigmentation seen along the dorsal abdominal midline of *D. novamexicana* but not *D. americana*. This trait has previously been described as derived in *D. novamexicana* (Spicer, 1991); however, we see a similar dorsal midline lightening in at least some lines of *D. lummei* (see Figure 1), another member of the virilis group, suggesting that the dorsal midline activity of *ebony* existed prior to the divergence of *D. americana* and *D. novamexicana*. An unexpected change in pupal pigment patterning was also seen in *D. americana* and *D. novamexicana* *ebony* null mutants. Although *ebony* is known to affect pupal case development in *D. melanogaster* (Sherald, 1980), its loss causes a pale white pupa color rather than the dark pigmentation we see in *D. americana* and *D. novamexicana* *ebony* null mutants. Because *ebony* is required for the production of yellow pigments, the dark markings seen in *ebony* mutant pupal cases likely result from expression of an enzyme required for synthesis of dark pigments, such as *tan*. Finally, we found that *ebony* null mutants showed significant changes in the abundance of some CHCs in each species, but divergence of *ebony* did not contribute to differences in the CHC profiles seen between species. These observations illustrate how *cis*-regulatory changes can cause divergence of some, but not all, traits affected by a pleiotropic gene.

Observations reported in this work were made possible by the ability to manipulate the *D. americana* and *D. novamexicana* genomes with CRISPR/Cas9 genome editing. While this technology has great potential for allowing functional hypothesis testing in species that have not historically been considered genetic model systems, this work was not always straightforward. We hope that the detailed descriptions of our genome editing efforts provided in the Materials and “Materials and Methods”

section of this paper will be helpful for other researchers striving to manipulate the genomes of non-model species.

DATA AVAILABILITY STATEMENT

All datasets generated for this study are included in the article/**Supplementary Material**.

AUTHOR CONTRIBUTIONS

AL and PW conceived of the experiments and wrote the manuscript. AL, PS, and ZW performed the experiments. HC and PJ provided funding, advice, and oversight. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fevo.2020.00184/full#supplementary-material>

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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