The evolution of polymorphism in the warning coloration of the Amazonian poison frog *Adelphobates galactonotus*


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Abstract
While intraspecific variation in aposmatic signals can be selected for by different predatory responses, their evolution is also contingent on other processes shaping genetic variation. We evaluate the relative contributions of selection, geographic isolation, and random genetic drift to the evolution of aposmatic color polymorphism in the poison frog *Adelphobates galactonotus*, distributed throughout eastern Brazilian Amazonia. Dorsal coloration was measured for 111 individuals and genetic data were obtained from 220 individuals at two mitochondrial genes (mtDNA) and 7963 Single Nucleotide Polymorphisms (SNPs). Four color categories were described (brown, blue, yellow, orange) and our models of frog and bird visual systems indicated that each color was distinguishable for these taxa. Using outlier and correlative analyses we found no compelling genetic evidence for color being under divergent selection. A time-calibrated mtDNA tree suggests that the present distribution of dorsal coloration resulted from processes occurring during the Pleistocene. Separate phylogenies based on SNPs and mtDNA resolved the same well supported clades, each containing different colored populations. Ancestral character state analysis provided some evidence for evolutionary transitions in color type. Genetic structure was more strongly associated with geographic features, than color category, suggesting that the distribution of color is explained by localized processes. Evidence for geographic isolation together with estimates of low effective population size implicates drift as playing a key role in color diversification. Our results highlight the relevance of considering the neutral processes involved with the evolution of traits with important fitness consequences.

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Introduction

The presence of intraspecific variation in aposematic signals provides opportunities to examine the relative contributions of selection, drift, and geographic isolation to the evolution of this trait (Richards-Zawacki et al. 2013; Gehara et al. 2013; Kang et al. 2017; Lawrence et al. 2019). Because aposematic signals such as contrasting and bright colors advertise the presence of toxins or other defenses, the signal is under balancing selection and therefore is usually consistent in order to be effective as a deterrent (Endler and Greenwood 1988). Nevertheless, geographic variation in selective pressures, including differences in the local assemblage of predators, changes in predator behavior and Batesian and Mullarian mimicry complexes, can result in different colors being fixed in different populations (Endler and Mappes 2004; Darst and Cummings 2006; Amézquita et al. 2017). Adaptive potential is influenced by the distribution of genetic variation, and less effective aposematic coloration can be maintained if the level of gene flow is sufficiently low between populations with different colors (Lawrence et al. 2019). Furthermore, if different phenotypes experience the same level of selection, and effective population sizes are small enough, then diversification through random genetic drift is possible in accordance with Wright’s shifting balance theory (Wright 1932; Rudh et al. 2007; Chouteau and Angers 2012; Comeault and Noonan 2011).

Neotropical poison frogs (family Dendrobatidae) possess aposematic coloration in combination with skin toxins, which likely evolved in tandem (Summers and Clough 2001). Many species show variation in aposematic coloration and for this reason they are popular models to study the evolution of this trait (Chouteau and Angers 2012). Color differentiation in poison frogs has been a recent and recurrent evolutionary process for those species investigated thus far, often with intra-specific divergence in colors traced back to no earlier than the Pleistocene, 11,500–2.6 million years before present (Noonan and Gaucher 2006; Santos and Cannatella 2011; Gehara et al. 2013).

The Pleistocene is characterized by rapid environmental change and this seems likely to have altered both levels of connectivity and predation pressures. Disturbance during the Pleistocene has been characterized for some poison frogs as fine scaled habitat fragmentation, rather than large-scale retraction to refugia (Noonan and Gaucher 2006). Habitat disturbance during the Pleistocene resulted in fine-scaled divergence in the northern Amazonian dyeing poison frog *Dendrobates tinctorius*, which in concert with selection by predators, resulted in the establishment of populations that are monomorphic for different colors (Noonan and Gaucher 2006; Comeault and Noonan 2011). The process of isolation can also facilitate color diversification by hybridization following secondary contact. For example, hybridization has been implicated as the process generating color variation in the Andean harlequin poison frog *Oophaga histrionica* (Medina et al. 2013). Secondary contact and hybridization may be further facilitated by the likelihood of rapid habitat restructuring in the Amazon region (Sefc et al. 2017).

Isolation can also result from behavioral differences, potentially facilitating color differences at finer spatial scales than the effects of geographic isolation. For example, in parts of the strawberry poison frog (*Oophaga pumilio*) distribution, genetic distances between populations are more strongly associated with dorsal coloration than with landscape features, and this was attributed to selection against dissimilar immigrants (Wang and Summer 2010). Furthermore, in *O. pumilio*, sexual selection, coupled with drift in female preference is another of several mechanisms that have been proposed to explain color diversification (Mann and Cummings 2012; Gehara et al. 2013). Jointly analyzing whether genetic structure is associated with present day environmental features, or aposematic coloration, can provide clues about isolating processes and the scale at which these occur (Wang and Summer 2010). In addition, patterns of genetic structure associated with populations of different color types can be used to help infer whether signal variation is a consequence of dispersal or color becoming fixed within a population following de novo mutation.

Geographic isolation and drift has resulted in the diversification of color phenotypes for several poison frog species. Physical features that impede gene flow and influence the population sizes of poison frogs include rivers, topographic barriers, forest density and soil type because of their inherent variation in the water holding capacity (Kaefer et al. 2013; Simões et al. 2014; Ferreira et al. 2018). Where the habitat for *O. pumilio* is fragmented, low population sizes, increased drift, perhaps in combination with reduced predation pressure has allowed new phenotypes to increase in frequency and be fixed by selection (Rudh et al. 2007). The role of drift in generating color polymorphism has also been shown in the poison dart frog *Ranitomeya imitator*. Here it was shown that selection was relaxed at a point along an altitudinal gradient and this allowed new color phenotypes to be generated by drift (Chouteau and Angers 2012).

The Amazonian poison frog, *Adelphobates galactonotus*, varies in dorsal coloration throughout its distribution in eastern Amazonia, which comprises forest areas east of the Tapajós River to the Atlantic coast and south of the Amazon river to the southern boundary of the Amazon forest (Fig. 1; Hoogmoed and Avila-Pires 2012). Different dorsal colors (yellow, orange, red, blue, and brown) have not been found sympatrically and appear in a mosaic-like fashion across the species’ geographic range (Hoogmoed and Avila-Pires 2012).
The variation in the dorsal coloration of *A. galactonotus* is intriguing because it is non-mimetic and populations containing individuals with different color types are found in close proximity and within areas of similar environmental conditions.

Here we use field-collected dorsal color measurements, mtDNA sequence data, and a data set of several thousand single nucleotide polymorphisms (SNPs) from *A. galactonotus* to test the following hypotheses: (1) the time of divergence between populations containing different dorsal colors is consistent with the timing of color diversification in other Neotropical poison frogs, (2) the same dorsal coloration evolved independently in different locations, and (3) that dorsal coloration of *A. galactonotus* is under selection.

**Methods**

**Study area**

We sampled *Adelphobates galactonotus* across most of its known geographic distribution in eastern Amazonia, south of the Amazon River, in the Brazilian state of Pará. This region contains two large southern tributaries of the Amazon, the Xingu and the Tocantins rivers (Fig. 1). Our sampling effort was concentrated to the west of the Tocantins River. Because of the difficulty of sampling within large tracts of forest, individuals were sampled either in close proximity (<15 km) to rivers, or to the few roads that are in the region. Visual appraisal suggests that structural elements of the forest do not vary systematically with respect to the presence of rivers or roads.

**Tissue sampling**

A total of 220 individual samples were obtained from fieldwork carried out from 2012 to 2014 and from the tissue collection of Museu Paraense Emílio Goeldi (MPEG), Belém, Brazil (Table 1). Fieldwork was conducted in three regions from eastern Amazonia: Caxiuana–Anapu river basin, Tapajós–Jamanxim river basin, and Carajás (Fig. 1). Tissue samples consisted of toe clips, or muscle/river tissue when voucher specimens were collected. Voucher specimens were deposited at the Osvaldo Rodrigues da Cunha
Table 1 A summary of the data collected from localities in two regions (east and west of Xingu River)

<table>
<thead>
<tr>
<th>Region</th>
<th>Locality</th>
<th>Color</th>
<th>code</th>
<th>RS</th>
<th>COI</th>
<th>16S</th>
<th>SNPs</th>
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<tr>
<td>East</td>
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<td>Orange</td>
<td>OAR</td>
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<td>OAR</td>
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<tr>
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<td>BAR</td>
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<td>4</td>
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<td>3</td>
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<td></td>
<td>Cacoal, Atuá River right bank</td>
<td>Blue</td>
<td>BAR</td>
<td>6</td>
<td>2</td>
<td>4</td>
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<td></td>
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<td>BRAR</td>
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<td>Brown</td>
<td>BRAR</td>
<td>13</td>
<td>9</td>
<td>11</td>
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<td>BAL</td>
<td>13</td>
<td>10</td>
<td>10</td>
<td>16</td>
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<td>BC</td>
<td>2</td>
<td>2</td>
<td>6</td>
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<td>2</td>
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<td>YMV</td>
<td>5</td>
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<td>Porto de Moz</td>
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<td>OPM</td>
<td>2</td>
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<td>Caracol, UHE Belo Monte</td>
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<td>YBM</td>
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<td>OBA</td>
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<td>3</td>
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<td>Nazaré</td>
<td>Blue</td>
<td>BNA</td>
<td>2</td>
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<td></td>
<td>Serra das Andorinhas</td>
<td>Orange</td>
<td>OAN</td>
<td>2</td>
<td>2</td>
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<tr>
<td>West</td>
<td>Projeto ALPA</td>
<td>Red</td>
<td>RAL</td>
<td>2</td>
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<td></td>
<td>Tapajós MU</td>
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<td>OT</td>
<td>4</td>
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<td></td>
<td>Boca do Rato</td>
<td>Yellow</td>
<td>YT</td>
<td>7</td>
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<td>BT</td>
<td>6</td>
<td>22</td>
<td>17</td>
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<td>Mina do Palito</td>
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<td>BJ</td>
<td>7</td>
<td>6</td>
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<td>Jamanxim river left bank</td>
<td>Blue</td>
<td>BJ</td>
<td>6</td>
<td>5</td>
<td>3</td>
<td></td>
</tr>
</tbody>
</table>

Locality names and their code refer to the nearest geographic landmark (detailed location information is available on request from the corresponding author). Localities are grouped according to their position on each side (east/west) of the Xingu, a large southern tributary of the Amazon River. Color refers to the four dorsal coloration categories observed in this study. The four right-hand columns give the number of individuals used for color measurement (RS), Cytochrome Oxidase 1 sequencing (COI), 16S rDNA sequencing (16S), and genotyped at Single Nucleotide Polymorphisms (SNPs).

*The Betel sampling locality is divided by a river, the section furthest from Betel is designated M18 in Fig. 7d

Herpetological Collection of MPEG or at the Amphibian and Reptile Collection of Instituto Nacional de Pesquisas da Amazônia (INPA-H) in Manaus, Brazil. All tissue samples were stored in absolute ethanol prior to DNA extraction. We define a sampling locality as an area where all individuals were captured within 80 km of each other. These locations were consistently in continuous forest, without apparent biogeographic barriers (such as the presence of large rivers).

**Color data collection**

Adelphobates galactonotus has been collected at several of the sampling localities over a number of years and stored at MPEG (e.g., Caxiuana-Anapu locality has samples collected over ~20 years). These collections consistently found the same single dorsal color at each of the localities (Table 1; Hoogmoed and Avila-Pires 2012). The dorsum of *A. galactonotus*...
Fig. 2 Reflectance spectra for the dorsal coloration in four color categories: blue (a–e), brown (f), orange (g–i) and yellow (j). Images of live frogs representing these categories (correspondingly labeled a–j) are shown in Fig. 1. In each figure, the average reflectance spectrum value is shown with a solid black line while the shading shows the standard deviation. Figure titles provide information on the color category and sampling location where: East = east of the Xingu river; West = west of the Xingu river, and the localities are given using the codes provided in Table 1. At each locality, the number of individuals used for the measurement of reflectance spectra were as follows: OAL = 12; OAR = 20; BAR = 17; BRAR = 21; BAL = 13; BC = 2; OT = 4; YT = 7; BT = 6; BJ = 6

most often appears to have consistent coloration, though there are instances of color patterns in the form of spots or flecks (Table 1). To describe color variation, we measured the color of 111 live specimens at 20 localities distributed in the Caxiuanã-Anapu, Tapajós-Jamanxim and Carajás regions (1–13 individuals used per locality; Table 1) with an Ocean Optics USB200 spectrometer (Ocean Optics, Inc., Dunedin, FL, USA), an Ocean Optics DH-2000 BAL light source and Spectra Suite® software. The spectrometer was calibrated for both white and black prior to the individual measurements being taken. All of these measurements were obtained in the laboratory under equivalent lighting conditions. Individual reflectance spectra were obtained by averaging and smoothing two dorsal measurements, one at the dorsal surface of the head, the other on the lower dorsum (smoothing parameters: span = 0.2, negative values fixed to 0). The averages of these spectra datasets were used to represent the dorsal coloration reflectance for each locality (Fig. 2).

To categorize dorsal coloration and evaluate its variability, the mean reflectances were visualized and colorimetric variables (mean brightness, hue and chroma) from reflectance data were calculated (Endler 1990; Anderson and Prager 2006; Kemp et al. 2015) in the R package pavo (Maia et al. 2013). To test whether differences in dorsal coloration can be discriminated by conspecifics and potential predators (birds), we analyzed spectra with avian (tetrachromatic-V retina type) and frog (trichromatic) visual models using pavo (Vorobyev and Osorio 1998; Vorobyev et al. 2001). To model frog vision, we used the data from three cone classes of O. pumilio (Siddiqi et al. 2004). Specific predators of A. galactonotus are unknown, but birds are predators of many aposematic species including poison frogs (Lindström et al. 1999; Master 1999; Exnerová et al. 2008). Furthermore, in field experiments, birds were reported as the most frequent predators of artificial poison frog models (Noonan and Comeau 2009; Paluh et al. 2014; Rojas et al. 2015).

To test whether or not different color categories are perceived differently by conspecifics and birds, we estimated pairwise (localities) contrasts expressed in units of Just Noticeable Difference (JND). When JND values are above one it indicates that color distances would be discriminated by the visual system modeled (Siddiqi et al. 2004). All analyses were conducted in the R statistical computing environment (R Core Team 2014).

Mitochondrial sequence data

Total genomic DNA was extracted from preserved tissue samples using Promega® Wizard Extraction Kits, following
the manufacturer’s guidelines. We amplified a 559 bp fragment of the Cytochrome Oxidase I (COI) mitochondrial gene from 133 individuals using primers CHMf4 and CHMR4 (Che et al. 2012) and a 402 bp fragment of the 16S rDNA from 176 individuals using the primers 16Sar and 16Sbr (Palumbi 1996). PCR cycles were as follows: 30 s at 92 °C followed by 35 cycles of 92 °C for 10 s, 50 °C for 35 s, and 72 °C for 90 s, with a final extension of 10 min at 72 °C. Amplicons were purified for sequencing with ExoSAP-IT solution, then subjected to EtOH/EDTA precipitation and sequenced in the forward direction using an ABI sequencer 3031x (Applied Biosystems*).

**Development of SNPs**

Genomic DNA was extracted from 186 tissue samples (Table 1) using the GeneCatch™ Blood & Tissue Genomic Mini Prep Kit (Epoch Life Science, Inc.). Approximately 0.5 µg of DNA was sent to Diversity Arrays Technology Pty. Ltd. (Canberra, Australia—http://www.diversityarrays.com) where SNP discovery and genotyping was performed using the standard DartSeq™ protocol. DartSeq™ genotyping is a SNP-based genotyping-by-sequencing approach (Jaccoud et al. 2001; Kitanj et al. 2012; Lambert et al. 2013) where sequencing is carried out on an Illumina platform (Sansaloni et al. 2011) to genotype thousands of SNPs across the genome (Petroli et al. 2012).

**Library preparation**

Template DNA was incubated in a 1X solution of MultiCore™ restriction enzyme (RE) buffer (Promega) at 37 °C for 2 h to check genomic DNA quality. Approximately 100 ng per µL of each sample was digested with a combination of PstI and SphI restriction enzymes. PstI and SphI adapters and unique barcodes were ligated to each sample.

Each sample was amplified using PCR primers specific to barcode and adapter sequences. The PCR conditions were as follows: 1 min initial denaturation at 94 °C, followed by 30 cycles of 20 s denaturation (94 °C), 30 s annealing (58 °C) and 45 s extension (72 °C), and a final extension of 7 min at 72 °C. Using ~10 µL of each sample, all samples were pooled, diluted, and denatured using NaOH in preparation for hybridization to the flow cell. The library was sequenced on an Illumina HiSeq 2500 platform (single end read), resulting in 77 bp long fragments. A proportion of the samples (>40%) were processed again through the whole library preparation protocol and downstream analyses to create the set of technical replicates that were used to assess the reproducibility.

**Quality control and initial SNP calling**

Library sequences were converted to fastq format using the Illumina HiSeq®2500 software, and individuals were demultiplexed based on the ligated barcode using customized scripts developed by DARt. Each read was assessed using Phred quality scores (Q-scores; Ewing and Green 1998) and any reads containing Q-scores <25 were removed. All reads were checked against the DARt database and GenBank viral and bacterial sequences to assess for contamination. Following this primary workflow, SNPs were identified and called following the standard procedure in the DARt proprietary pipeline DArtSoft14™ (Diversity Arrays Technology). The workflow is technically similar to the STACKS pipeline (Catchen et al. 2013), but it differs by first calling sequence clusters from all pooled samples, prior to being called for each individual. All invariant sequence clusters were removed, and SNPs were called only if they were present in both homozygous and heterozygous forms across the whole data set. SNPs were retained in the data set if the average ratio of read depth between alleles ≤0.75, with reproducibility of 100% and a minimum read depth of five. We further filtered the dataset using the following criteria: We only included SNPs with a call rate of >98% (i.e., <2% missing data), where the average read depth for both alleles was >5 and the Minor Allele Frequency (MAF) >0.05. We retained only a single SNP per fragment to avoid creating a dataset containing closely linked loci.

**mtDNA analysis**

Sequences were aligned using the ClustalW algorithm (Thompson et al. 1994) implemented in BioEdit 7.2.5 (version updated in November 2013, Hall 1999). Substitutions were checked with the original chromatograms. In order to evaluate the genealogical relationships among haplotypes and overall haplotype distributions, a haplotype network for COI was built from the resulting alignment using statistical parsimony (Templeton et al. 1992) with the software TCS 1.21 (Clement et al. 2000). Analyses of genetic diversity at 16S rDNA and COI regions were carried out for each sampling location and collectively for individuals sampled on each side of the Xingu river using DnaSP v.5.10.01 (Librado and Rozas 2009).

**Estimating divergence times**

To estimate the divergence times for A. galactonotus we used BEAST 1.8.0 (Drummond et al. 2012) to analyze 59 unique mtDNA haplotypes recovered from a concatenated database of 16S rDNA and COI sequences (totaling 962 bp). To construct a gene tree, sequences for 16S rDNA and COI were obtained from GenBank for A. castaneoticus, A.
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...*quinevittatus* and *Phyllobates terribilis* to use as outgroups (accession numbers DQ502058, DQ502157, DQ502234, DQ502780, DQ502861, and DQ502906). Sequences were aligned with the MUSCLE algorithm (Edgar 2004) as implemented in MEGA 6.06 (Tamura et al. 2013). The most probable substitution model for each region was estimated independently in jModeltest 2.1.7 (Guindon and Gascuel 2003; Darnbta et al. 2012). We defined the approximate average rate of evolution for each region following Mueller (2006). As time calibration priors, we considered means and standard deviations previously estimated for the most recent common ancestor for *A. galactonotus* and *A. quinevittatus* (10.0 ± 2.5 million years ago—m.y.a.) and *A. galactonotus* and *A. castaneoticus* (8.0 ± 2.1 m.y.a.) (Santos et al. 2009). We followed the recommendations provided by the software developers for BEAST 1.8.0 to select the parameters and priors for this analysis (Drummond et al. 2012). We used the following priors: partitions with unlinked substitution and clock models, evolutionary rates (the number of substitutions per site every 100 million years for COI = 1.04 ± 0.27 and 16S = 0.70 ± 0.03 (Mueller 2006), a lognormal distribution and the tree prior was Yule. We applied a chain length of 80 million generations with samples taken every eight thousand generations. The first 10% of the trees were discarded as burn-in, resulting in 9000 sampled trees in the Markov Chain Monte Carlo (MCMC) run. The stationarity of the posterior distributions for all model parameters, including medians and 95% Highest Posterior Density intervals (HPD) of the nodes was verified on Tracer 1.6.0 (Rambaut et al. 2014). The final consensus tree was obtained from the MCMC output using Tree Annotator 1.8.0 (Drummond et al. 2012).

**Constructing a population tree with SNP data**

We generated a population tree with our SNP data using the Multispecies Coalescent (MSC) implemented by the SNAPP package (Bryant et al. 2012) and BEAST v. 2.4.7 (Drummond and Rambaut 2007; Bouckaert et al. 2014). The SNAPP analysis directly models incomplete sorting, which is expected in recently diverged species or populations, and can therefore be helpful in visualizing uncertainties around the majority-rule gene tree. For this analysis we randomly selected 2 individuals at each sampling location to reduce the computational time. Two independent runs were run for 10 million generations, sampling every 1000 generations. After stationarity was reached, we checked for convergence in Tracer and for consistency of the recovered topologies (Rambaut et al. 2014). The two runs were combined using LogCombiner, using a burnin of the first 3.5 million generations. The final tree was visualized in DensiTree v.2.2.6 where the individual lines represent trees sampled every 1000 generations (Bouckaert 2010).

**Ancestral state analysis**

To test whether evolutionary transitions in dorsal coloration occurred we conducted ancestral character reconstruction. Ancestral state reconstruction requires an accurate estimate of phylogenetic relationships, a model of trait evolution and accurate trait data (Duchêne and Lanfear 2015). Ancestral states were analyzed using the mtDNA consensus tree obtained using 16S rDNA and COI. To test whether this tree was influenced by incomplete lineage sorting, or undetected male gene flow we compared the topology with the population tree constructed using SNP data. Variation in dorsal coloration was collapsed into four character states according to the relative reflectance in short, medium, and long wave lengths, based on spectral data (in decreasing order: blue, yellow, and orange), and absolute light reflectance (brown); these color categories are distinguishable to a wide range of potential predators (Endler and Mappes 2004). A fifth state (patterned) was added for the outgroup species in the mitochondrial tree to indicate a general black dorsum crossed by white longitudinal stripes (*A. quinevittatus*) or blotches (*A. castaneoticus*). A black dorsum with yellow/whitish dorsolateral stripes represents the ancestral color pattern in the genus *Phyllobates*, although the species we used here (*P. terribilis*) loses this pattern as an adult and becomes uniformly yellow (see Fig. S8 for images of these three outgroup species). The use of parsimony for ancestral state reconstruction can be biased by rapid evolution, a problem that can be solved using a maximum likelihood or a Bayesian approach (Cunningham et al. 1998). We preferred a maximum likelihood approach rather than stochastic character mapping by Bayesian inference because of the inherent difficulties in visually representing the results (Revell 2013). The reconstruction of ancestral color states was implemented with a continuous-time Markov model (MK) in the R package ape (Paradis and Schliep 2018) and phytools (Revell 2012). The MK model assumes the random evolution of these color states and estimates the relative probability of each color category at each node. The analysis accounts for branch length, and transitions are independent of changes occurring elsewhere in the tree. Our model assumed that transitions occurred at equal rates for each color category. Equal transition rates were assumed because color type is highly variable in dendrobatid frogs and no directional changes were apparent.

**Genetic structure estimated with SNP data**

To describe patterns of genetic structure we performed a Principal Component Analysis (PCA) based on genotype
frequency data using the R package adegenet (Jombart and Ahmed 2011). Genetic structure was further investigated using the software fastSTRUCTURE (Raj et al. 2014), a fast algorithm for the approximate inference of the model implemented by STRUCTURE (Pritchard et al. 2000). We ran fastSTRUCTURE using K values ranging from 1 to 10, the logistic prior and five-fold cross-validation. Analyses on the entire dataset revealed a clear partitioning between individuals east and west of the Xingu river. We then repeated the analyses separately for individuals sampled east and west of the Xingu river. We considered smaller values of K with prediction errors within one standard error of the lowest estimate as useful models of the data (Raj et al. 2014). Results from fastSTRUCTURE were plotted in R using the package pophelper.

We then tested whether geographic distance and the presence of rivers explained the observed patterns of genetic differentiation. We constructed a matrix of river distances by summing river widths that were measured where intersected by a straight line between sampling locations. Genetic divergence was estimated with Weir and Cockerham’s FST calculated using GenAIEx v. 6.5 (Peakall and Smouse 2012). Spatial analyses were performed separately for sampling locations east and west of the Xingu river. Three different maximum-likelihood population-effects (MLPE) mixed models were tested, using the R packages corMLPE and nbme (Clarke et al. 2002). The first model included geographic distance alone, the second river distance alone, and the third used both distance matrices as predictors. The intercepts, slopes, and random effect terms were estimated using restricted maximum likelihood. Because marginal R2 values are biased towards more complex models, the best model was selected using Akaike information criteria (AIC) and Bayesian information criteria (BIC), following Row et al. (2017).

To assess whether loci putatively under the influence of selection were affecting patterns of genetic structure, we ran all of the analyses a second time with the data set excluding outlier loci identified via FST outlier testing in OutFLANK (see below; Whitlock and Lotterhos 2015). In all cases our analyses produced very similar results and we present genetic structure analyses using the SNP data excluding FST outliers.

**Effective population size**

For each sampling location we used NeEstimator v. 2.1 (Do et al. 2014) to calculate effective population size by linkage disequilibrium (LD). In a closed population with random mating and unlinked loci, LD results from genetic drift, occurring at a rate inversely proportional to the Ne. The random mating model under LD was chosen, and a critical value of 0.02 and 0.05 was selected to represent the minimum allele frequency cut off (O’Leary et al. 2013; Trask et al. 2017). Confidence intervals for Ne estimates were obtained by jackknifing (Do et al. 2014).

**Signatures of selection**

Divergent selection could result in significantly larger genetic distances than expected under neutrality for loci situated in parts of the genome responsible for the traits under selection. We evaluated this process by searching for FST outliers. For this approach we excluded localities with <5 individuals collected to ensure adequate representation of allele frequencies within each sampling locality. This resulted in data from 12 locations (with 5–26 individuals per location). Because of strong genetic divergence between populations separated by the Xingu River (see results), we conducted FST outlier analysis separately for populations east (K = 8 populations) and west (K = 4 populations) of the Xingu River. We used the software OutFLANK (Whitlock and Lotterhos 2015) and applied the default parameters for False Discovery Rate (FDR = 5%) and for the left and right trim factors of the FST distribution.

We also searched for signatures of selection on dorsal coloration using linkage disequilibrium (LD) network analyses developed by Kemppainen et al. (2015) and implemented in R package LDna. LD network clusters (single-outlier clusters, SOCs) represent networks of loci in high LD. Using loci from SOCs as an input for PCA, it is possible to discern whether these loci in LD represent population structure (if they group individuals by geography) or local adaptation (in which case they will group individuals by phenotype). This method has been successfully used to identify genomic regions under selection in distinct ecotypes of three-spined sticklebacks (Kemppainen et al. 2015) and parallel selection in ecotypes of Littorina saxatilis (Ravinet et al. 2016). LDna was first run using the data from all sampling localities and SNPs with frequencies >5%. This first analysis revealed that the strongest patterns of LD were associated with divergence between populations east and west of the Xingu river. Analyses were then run separately for populations on each side of the Xingu River. The two main parameters in LDna are E0 min (minimum number of edges in a cluster), and λ which represents the change in LD when two clusters merge. We retained only SOCs with a E0 min of ten and for which λ exceeded the median by a multiplier (φ) of five. These parameters were chosen interactively as outlined in the LDna tutorial and after consulting the developer of the software.

We looked for associations between color category and SNP genotypes using the latent factor mixed model approach implemented within the software LFMM (Frichot et al. 2013). The LFMM analysis considers genotypic matrix values as response variables in a linear regression model. The LFMM analysis accounts for the effects of neutral genetic structure by defining a number of latent
factors (LF), estimated beforehand by genetic clustering analysis (Fréchet et al. 2015). After accounting for genetic structure, significant associations between SNP allele frequencies and observed ecological (or morphological) variables may be interpreted as evidence for selection (Fréchet and François 2015). Compared to other outlier and environmental association analysis methods, LFMM reduces the type I error (i.e., false positives) in datasets with complex underlying genetic structure (Fréchet et al. 2013; de Villemereuil et al. 2014; Rollstup et al. 2015). We ran LFMM in R using the LEA package (Fréchet and François 2015). The number of LFs was based on the number of sampling locations, justified on the basis of the observed genetic and phenotypic structure (east: LF = 8; west: LF = 4). The program was run for 10,000 iterations with the first 5000 iterations discarded as burn-in. This was repeated 5 times and median z-scores were combined across runs, and p-values were adjusted for multiple tests using the Benjamini-Hochberg procedure (False Discovery Rate (FDR) = 5%) as outlined within the LEA package (Fréchet and François 2015). The median genomic inflation factor (GIF) was calculated across runs following Devlin and Roeder (1999) to further evaluate FDR. Successful control of FDR is indicated by a GIF close to, or slightly below 1.0 (Fréchet et al. 2015). A p-value threshold of $p < 10^{-6}$ was used to identify significant SNP-by-color associations.

DNA sequences containing SNPs identified as outliers or associating with color were ran through the BLASTN online database to search for matches with functionally relevant genes that may relate to color synthesis. We set the BLAST e-value acceptance threshold as $1 \times 10^{-6}$ with a sequence identity of more than 70% (Benestan et al. 2016).

Results

Analysis of dorsal coloration

The dorsal coloration of Adelphobates galactonotus could be placed into four categories, yellow, orange, blue, and brown with no individuals found to have intermediate coloration (Figs. 1, 2 and Table 1). Consistent with Hoogmoed and Avila-Pires (2012), we never observed more than a single color category at a locality, with individuals exhibiting only slight variation in color and pattern. Although we can statistically partition the color variation into four color categories, there appears to be considerable geographic variation within each of these categories (Figs. 1, 2 and S1). In particular, the blue color category showed a great deal of variation, not only with respect to reflectance spectra, but also the pattern of dorsal coloration (Fig. 1).

Models of bird and frog vision suggest that both of these taxa can discriminate between dorsal coloration in the blue, yellow, orange, and brown categories (JND > 1, Fig. S2) with the birds being capable of detecting subtler differences (JND values were higher for birds).

Timing the divergence into different color categories

We identified 37 variable sites and 35 haplotypes for 16S rDNA and 42 variable sites and 34 haplotypes for COI. Haplotype networks built using statistical parsimony were similar for both mtDNA regions, hence we show only the network using COI (Fig. 3; see Table S1 for additional data). No association was found between color categories and the genetic structure inferred from mtDNA markers (e.g., some individuals in different color categories have the same haplotypes). However, there were no haplotypes in common for A. galactonotus sampled east and west of the Xingu River, suggesting that the river has been a long-term barrier to female mediated gene flow.

The consensus tree based on mtDNA has well supported clades (bootstrap support > 0.70) that show instances of populations with different dorsal coloration sharing a common ancestor (Fig. 4). Samples collected from regions west and east of the Xingu River form two well-supported basal clades. The most recent common ancestor for these clades is estimated to be late Pliocene, most probably around 4.8 million years ago (Fig. 4). This is broadly consistent with the timing for range expansions reported for poison frogs in the Amazon basin (<10.0 m.y.a.) by Santos et al. (2009). The genetic divergence associated with the present day distribution of color categories occurred more recently, during the Pleistocene.

The population tree constructed by coalescence using SNP data with the software SNAPP showed a broadly similar topology as the consensus tree using mtDNA, there are several well supported clades, each with populations of different dorsal colorations (Fig. 5). The clade representing the basal split associated with the Xingu river, and two clades in the east, each have posterior probabilities of one and correspond to the three clades in the mtDNA consensus tree with bootstrap support > 90. In addition, in the east, the three nodes associated with populations BAR-YMV and the two nodes associated with populations OA-BC each had posterior probability support of 1 (Fig. 5). Similarly, these same nodes are well supported in the mtDNA consensus tree with bootstrap support > 0.88. This SNP based population tree demonstrates that incomplete lineage sorting, or male mediated gene flow, is not confounding our conclusion that populations with different colors share a common ancestor.

Ancestral state analysis

The reconstruction of ancestral coloration in A. galactonotus provides some support for transitions of color within the
two major clades, east and west of the Xingu river (Fig. 6). There are two instances of populations with frogs of yellow color having ancestors with high posterior probability for blue coloration, and one well supported instance of a population with yellow frogs having an orange colored ancestor in the western clade. The node in the mtDNA consensus tree representing the most recent common ancestor for the western clade, has high bootstrap support (0.99) and also a very high posterior probability of blue coloration. The tips of this clade represent populations with yellow, blue and orange coloration. In the eastern clade, there was strong support for a transition from blue to orange and orange to brown. For these two cases, there was also strong bootstrap support for the nodes associated with these transitions (blue to orange = 0.92 and orange to brown = 0.75) and the timing of these transitions was estimated to have occurred during the Pleistocene (Fig. 4). It can be seen that the ancestral colors could not be confidently assigned for several of the most recent common ancestors for population with different colors. The ancestral coloration for the eastern clade was also equivocal.

Neutral genetic structure based on SNPs

We identified a total of 7963 SNPs after filtering. Principal Components Analysis (PCA) resulted in two axes (PC1 and PC2) summarizing nearly 70% of the total genetic variation. Two genetic clusters corresponded to samples collected from the west and east of the Xingu River (Fig. S5b). The PCAs carried out separately for samples collected from each side of the Xingu River did not show patterns of divergence associated with color (Fig. 7a, b), but there were clear signals of geographic structure, with individuals clustering according to sampling locations. West of the Xingu River, the major differentiation is between blue populations (from Moraes de Almeida and the Tapajós River) and orange and yellow populations from the Tapajós River.

Analysis of genetic clustering using fastSTRUCTURE suggested values of $K$ that ranged from two to four (Fig. 7c–e) for both the east and west datasets. These genetic groupings contain populations of frogs with different categories of dorsal coloration, potentially reflecting the relatively recent diversification shown with mtDNA (Figs. 3 and 4). Nevertheless, limited gene flow among
Fig. 4 A time calibrated phylogenetic tree obtained using BEAST with sequences from 16S rDNA and COI. The two major clades recovered correspond to samples collected east (green shading) and west (yellow shading) of the Xingu River. Error bars on the nodes represent the 95% highest posterior density interval for node ages. The posterior probability of nodes lower than 0.70 are omitted. The timing of divergence (m.y.a) can be gauged using the scale given beneath tree. Tips are colored according to dorsal coloration in the four color categories; blue, orange, yellow, and brown. White represents samples obtained from specimens of unknown color. The black tips correspond to outgroup sequences (Pl = Phyllobates terribilis; Ag = Adelphobates quinquevittatus; Ac = Adelphobates castaneoticus). Codes indicate the sampling localities described in Table 1. The first letter in each code refers to color (B = blue, Y = yellow, B = brown, O = orange, U = unknown). Summary statistics for the mtDNA data at each locality is detailed in Table S1.

Sample locations is shown by significant pairwise $F_{ST}$ values between each of the sampling localities (east, $N=8$; west, $N=4$; Table S2). For localities with SNP data from a sufficient number of individuals, effective population sizes were low enough to suggest considerable levels of genetic drift ($N_e$ ranged from 45–213), though large or infinite confidence intervals bounded most of these estimates (Table S4).

There was no evidence of isolation by geographic or river distance in populations sampled west of the Xingu River, though this is likely to be because only four populations were sampled. East of the Xingu river both geographic ($p < 0.000001$) and river distance ($p < 0.005$) explained part of the observed genetic structure. The model including both river and geographic distance had the lowest AIC and BIC and explained the majority of the variance ($R^2 = 0.57$, Fig. 8, Table S5).

Detection of outlier and color-associated SNPs

The $F_{ST}$ outlier tests were carried out separately for samples collected east and west of the Xingu river. West of the river there were 13 SNPs (0.25 % of SNPs) identified as having
significantly high $F_{ST}$ values based on a model of neutral $F_{ST}$ distributions, east of the river no $F_{ST}$ outliers were detected. However, the model fit for the $F_{ST}$ distributions using OutFLANK was poor for both analyses (see $F_{ST}$ histograms: Figs. S3 and S4). This could be a result of strong genetic structure among many of the sampling localities (e.g., Fig. 6 and Table S2).

SNP-by-color association analyses using LFMM (Fröhlich et al. 2013) identified SNPs that were significantly associated with color, but the number of these SNPs differed east and west of the Xingu river (Table S3). The identity of SNPs that were associated with color also differed east and west of the Xingu River, with only 1.2% of these SNPs being in common. Furthermore, the color these SNPs were associated with differed when comparing associations found east and west of the Xingu. There were four SNPs associated with brown east of the Xingu that were associated with blue west of the Xingu, two SNPs associated with yellow in the east, were associated with orange in the west and two SNPs associated with yellow in the east, were associated with blue in the west. Of the 13 $F_{ST}$ outlier loci identified west of the Xingu river using OutFLANK, four of these SNPs were also identified as significantly associated with color using LFMM.

Linkage Disequilibrium (LD) network analysis identified several single outlier clusters (SOCs) of loci in high LD, most of which were consistent with the genetic structure associated with geographic distance and the presence of rivers, and none associated with dorsal coloration. When we analyzed the whole dataset, six SOCs were identified (Fig. S5a). Four of these SOCs grouped individuals east and west of the Xingu river (Fig. S5a, c–f) and the remaining two SOCs were consistent with genetic structure between two regions east of the Xingu River (Altamira and Serra Sul, respectively; Fig. S5g–h). A separate analysis of data collected east of the Xingu river with LDNA identified nine SOCs (Fig. S6a), eight of which were associated with geographic structure and none grouped individuals according to coloration (Fig. S6). The analyses of individuals west of the Xingu river identified only one SOC, which grouped individuals according to sampling location (Fig. S7).

We found no significant matches to functional genes with DNA sequences identified as outliers or associated with color using LFMM.
Fig. 7 Principal Component Analyses (PCA) based on SNP data from samples located east (a) and west (b) of the Xingu River. Data points are color-coded to reflect the color category. The figures show the prediction error from five-fold cross-validation with fastSTRUCTURE analysis carried out separately for samples collected east and west of the Xingu River. Histograms represent individual ancestry proportions that were calculated using fastSTRUCTURE. Results are given for individuals west (d) and east (e) of the Xingu River for values of K ranging from two to four.

Discussion

The dorsal coloration of the poison frog *Adelphobates galactomitus* can be categorized as yellow, orange, blue or brown, and individuals inhabiting the same region always belong to the same color category. Using both mitochondrial and SNP data, we show that populations with different colors are grouped in well supported clades and share a common ancestor. There is no obvious geographic pattern to color type, and our time calibrated phylogeny suggests that the present day distribution of coloration is a result of processes that most likely took place during the Pleistocene. We have some evidence from ancestral character state analysis that dorsal coloration has transitioned from one color category to another, but the ancestral color type could not be determined for many nodes. There are several possible explanations for the present day distribution of dorsal coloration. Broadly, these include founding events and hybridization,
or the effects of geographic isolation, where new mutations fixed by drift or selection might account for the color variation observed today.

Genetic divergence associated with the current distribution of colors occurred well within the Pleistocene, even allowing for the uncertainty around estimates of clade ages (Fig. 4). This was a period of high and repetitive climatic instability, especially in the eastern portion of the Amazon basin (Maslin and Burns 2000; Vonhof and Kaandorp 2011; Cheng et al. 2013). Such cyclic disturbances seem likely to have generated intermittent isolation and small population sizes through the contraction and expansion of adequate habitat, and the formation of new riverine barriers. Populations of A. galactonotus could be especially prone to isolation, partly because of their patchy distribution, a feature contributed to by their dependence on small water bodies for reproduction (Hoogmoed and Avila-Pires 2012).

Our estimates of low contemporary effective population sizes \( N_e \sim 50-250 \) (Table S4) and significant pairwise \( F_{ST} \) estimates (Table S2) are compatible with generally low levels of gene flow among populations.

The present day distribution of dorsal coloration could be a result of de novo mutations for color becoming fixed, from founding events distributing pre-existing color types, or hybridization and rapid selection on standing genetic variation (e.g., Barrett and Schluter 2008; Sefc et al. 2017). Our data suggests that localized color transitions are a more likely explanation for the current color variation than founding events distributing pre-existing color types. We have strong support from both SNP and mtDNA data that for several clades, populations with different dorsal coloration share a common ancestor (Figs. 4 and 5). We could also confidently infer transitions in dorsal coloration for at least a few lineages with ancestral character state analysis (Fig. 6). Alternately, if there was widespread movement of pre-existing color variation, we expected that genetic divergence would generally be greater between populations with different categories of dorsal coloration compared with those of the same color category. Contrary to this, we show with both mtDNA and SNP data that genetic structure is more strongly associated with geographic distance and the presence of rivers, rather than color type (e.g., Figs. 7 and 8). Furthermore, localized color transitions by de novo mutations might explain the geographic variation in dorsal coloration that we observed within each color category, especially among populations with the blue category (Figs. 2 and S1). Although strong genetic evidence for hybridization is lacking, we cannot discount it as a process generating color diversity, especially given that a brown color category appears intermediate to the blue and orange color categories on the right bank of the Anapu River (Fig. 7e). If the genetic distribution of dorsal coloration is at least partially a consequence of de novo mutations, or hybridization, the function of color as an aposematic signal suggests that selection has a role in driving these mutations to fixation.

In species with aposematic coloration, visually oriented predators are frequently thought to drive color diversification through natural selection (e.g., Jiggins et al. 2001). Our model of avian vision suggests that these predators can distinguish between the different color categories of A. galactonotus (Fig. S2). However, recent field experiments using A. galactonotus wax models failed to observe predatory bias toward particular colors (Rojas et al. 2015). It is possible that this result reflected the experimental design, because movement is relevant to prey detection by birds, which are considered predators of poison frogs (Pål et al. 2014). It is also possible that the color variation reflects selective pressures that are not current, or that the variation is a result of other forms of selection. These could include environmental differences, such as the lighting and background color, or differing predator assemblages (Endler 1992). It has also been demonstrated that sexual selection can effectively and rapidly drive a trait to fixation (Tazzaman and Iwasa 2010; Rudh et al. 2011) and thereby result in polyphyletic evolution. However, sexual dichromatism in A. galactonotus is not obvious to a human observer, so if sexual selection drives color divergence, it has not resulted in the magnitude of dichromatism reported in species where color plays a role in mate choice (Bell and Zamudio 2012).

Evidence for selection in our genetic data set was limited. We did identify SNPs potentially associated with directional selection, but believe these results should be cautiously interpreted. This is partially due to the following two reasons: first, the generally high \( F_{ST} \) between sampling localities, though accounted for to some degree in our analyses, may still compromise model outcomes (see Whitlock and Lotterhos 2015). Second, in cases where different localities had frogs assigned to the same color category, different
SNP identities were found to be associating with that color. Similarly, groups of loci in linkage disequilibrium were consistently associated with patterns of geographic structure rather than color. Future analyses could incorporate environmental variables and a more extensive sampling strategy to tease apart selection processes. In addition, a data set with higher density sequencing across the genome may be required, especially if a simple genetic mechanism underpins color in *A. galactonotus*.

Although the genetic mechanisms of color in anurans have received relatively little attention (Rudh et al. 2013), it is known that color is determined by three cell layers, the melanophores, the xanthophores and the iridophores, each containing specific pigments (Hofreiter and Schöneberg 2010). Conceivably, color variation could be due to one, or a few point mutations related to genes involved in color determination. These genes could be responsible for the synthesis or destruction of pigments like pteridines or carotenoids, which could be responsible for the yellow/orange/brown coloration. If the carotenoids and pteridines are removed, these colors disappear and the guanine crystals deeper in the epidermis are the main determinants of the iridescent, bluish color (Bagnara and Matsumoto 2007; Bagnara et al. 2007).

The maintenance of populations with different colors in close proximity suggests mechanisms that prevent recombination. Our model of frog vision suggests that *A. galactonotus* can discriminate between each of the categories of dorsal coloration (Fig. S2). Therefore, a plausible mechanism preventing recombination is that lower hybrid fitness selected for assortative mating with respect to color type. Narrow hybrid zones across unbroken rainforest landscapes have been characterized in other Amazonian frogs (e.g., Brown and Twomey 2009; Simões et al. 2012) suggesting that in these cases, hybridization lowers fitness. Further work could test for the presence of reproductive character displacement in *A. galactonotus* by taking color measurements at increasing distances from potential contact zones of populations with frogs of different dorsal coloration. Here, the prediction is that color contrast between the two populations will increase as the proximity between the two populations decreases (e.g., Lambert et al. 2013). Assortative mating that maintains reproductive isolation upon secondary contact is potentially a mechanism generating biodiversity both in the Amazon, and elsewhere, yet evidence for this process remains limited (Boul et al. 2007).

More generally, further knowledge of the processes resulting in spatial variation in trait variation is needed to test whether polymorphism is typically a precursor to speciation (McLean and Stuart-Fox 2014). The distribution of dorsal color variation and genetic variation that we describe here for *A. galactonotus* provides an ideal opportunity to further test these hypotheses.

**Data availability**

The mtDNA sequences can be accessed in GenBank (accession code KU597806 - KU597873). All mtDNA sequences, genotypes for SNP loci and summary statistics (call rate, heterozygosity, read depth, and reproducibility) are available from the Dryad Digital Repository (https://datadryad.org/stash/share/3fnquaEBQT_q4p7bH-A1WTH5YEVb4XZDn2XQLp1c). Trimmed sequence reads used to generate our SNP data set are held at Diversity Arrays Technology Pty. Ltd. (Canberra, Australia) and can be made available.

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**Author contributions** DR, AS, and APL designed the project. DR, APL, AS, and PIS collected samples. TCSA-P and MSH contributed useful discussion. TCSA-P, MSH and YOCB contributed samples and useful discussion. DR and AA measured, analyzed and interpreted the color data. AS, PIS, ILK, PM and RD analyzed and interpreted genetic data. DR and AS wrote the manuscript with contributions from all co-authors.

**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

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