Radiation and hybridization of the Little Devil poison frog *(Oophaga sylvatica)* in Ecuador

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Abstract

Geographic variation of color pattern in the South American poison frogs (Dendrobatidae) is an intriguing evolutionary phenomenon. These chemically defended anurans use bright aposematic colors to warn potential predators of their inpalatability. However, aposematic signals are frequency-dependent and individuals deviating from a local model are at a higher risk of predation. The well-known examples of Batesian and Müllerian mimics, hymenopterans (wasps and bees) and *Heliconius* butterflies, both support the benefits of unique models with relatively high frequencies. However, extreme diversity in the aposematic signal has been documented in the poison frogs of the genus *Dendrobates*, especially in the *Oophaga* subgenus. Here we investigate the phylogenetic and genomic differentiations among populations of *Oophaga sylvatica*, which exhibit one of the highest phenotypic diversification among poison frogs. Using a combination of PCR amplicons (mitochondrial and nuclear markers) and genome wide markers from a double-digested RAD data set, we characterize 13 populations (12 monotypic and 1 polytypic) across the *O. sylvatica* distribution. These populations are mostly separated in two lineages distributed in the Northern and the Southern part of their range in Ecuador. We found relatively low genetic differentiation within each lineage, despite considerable phenotypic variation, and evidence suggesting ongoing gene flow and genetic admixture among some populations of the Northern lineage. Overall these data suggest that phenotypic diversification and novelty in aposematic coloration can arise in secondary contact zones even in systems where phenotypes are subject to strong stabilizing selection.

Keywords: Dendrobatidae, aposematism, population genomics, admixture, gene flow, phenotypic variation

Resumen

La variación geográfica de los patrones de color en las ranas venenosas de América del Sur (Dendrobatidae) es un fenómeno evolutivo intrigante. Estos anuros con defensas químicas utilizan colores brillantes aposemáticos para advertir a sus posibles depredadores de su inpalatabilidad. Sin embargo, las señales aposemáticas son dependientes de la frecuencia, y los individuos que se desvían de un modelo local están en mayor riesgo de depredación. Dos ejemplos bien conocidos de mimetismo Batesiano y Mülleriano, en himenópteros (avispas y abejas) y mariposas *Heliconius*, confirman los beneficios de que hayan modelos únicos con frecuencias relativamente altas. Sin embargo, diversidad extrema en la señal aposemática se ha documentado en las ranas venenosas del género *Dendrobates*, especialmente en el subgénero *Oophaga*. En este estudio investigamos la diferenciación filogenética y genómica entre poblaciones de *Oophaga sylvatica*, las cuales poseen una de las diversificaciones fenotípicas más grandes de entre las ranas venenosas. Usando una combinación de amplicones de PCR (marcadores mitocondriales y nucleares) y otros marcadores amplios del genoma de un set de datos de secuenciación RAD de digestión doble, caracterizamos 13 poblaciones (12 monótipicas y 1 polítipica) a lo largo de la distribución de *O. sylvatica*. Estas poblaciones están separadas principalmente en dos lineajes distribuidos en el norte y sur de su área de distribución en el Ecuador. Encontramos relativamente baja diferenciación genética dentro de cada linaje, a pesar de la variación fenotípica considerable, y evidencia de flujo de genes y mezcla genética entre algunas poblaciones del linaje del norte. En general estos datos sugieren que la diversificación fenotípica y novedades en la coloración aposemática pueden surgir en las zonas de contacto secundarias incluso en los sistemas en los que los fenotipos están sujetos a fuerte selección estabilizante.

Keywords: Dendrobatidae, aposematism, population genomics, admixture, gene flow, phenotypic variation
INTRODUCTION

Aposematism is an antipredator adaptation that has evolved in many animals as a defense strategy. This adaptation combines warning signals (e.g., vivid coloration) with diverse predator deterrents such as toxins, venoms and other noxious substances. Most groups of animals include at least one aposematic lineage such as insects (e.g., Heliconius (Jiggins & McMillan 1997) and Monarch (Reichstein et al. 1968) butterflies), marine gastropods (e.g., nudibranchs (Tullrot & Sundberg 1991; Haber et al. 2010), amphibians (Howard & Brodie 1973; Brodie 2008), reptiles (Brodie III 1993; Brodie III & Janzen 1995) and birds (Dumbacher et al. 1992, 2000). The aposematism strategy is dependent on the predictability of the warning signal for effective recognition as well as learning and avoidance by predators (Benson 1971; Mallet et al. 1989; Kapan 2001; Pinheiro 2003; Ruxton et al. 2004; Chouteau et al. 2016). However, some aposematic species present high levels of polymorphic coloration and patterning (Przeczek et al. 2008). The causes and consequences of intraspecific variation in warning signals remains a fundamental question in the evolution and ecology of aposematism.

Dendrobatid poison frogs are endemic to Central and South America and have evolved chemical defenses coupled with warning coloration 3-4 times within the clade (Santos et al. 2003, 2009, 2014). Interestingly, some poison frog genera/subgenera display a wide range of inter- and intra-specific color and pattern variability, including Dendrobates sensu lato (Noonan & Wray 2006; Noonan & Gaucher 2006; Comeault & Noonan 2011), Adelphobates (Hoogmoed & Avila-Pires 2012), Ranitomeya (Symula et al. 2001; Twomey et al. 2013, 2014, 2016) and Oophaga (Wollenberg et al. 2008; Wang & Summers 2010; Brusa et al. 2013; Medina et al. 2013). For example, some species like R. imitator have evolved variation in coloration and patterning in a unique vertebrate example of Müllerian mimicry (Symula et al. 2001). Another example is the Strawberry poison frog, Oophaga pumilio, which is extremely polymorphic within the geographically small archipelago of Bocas del Toro (Panamá), but has more continuous and less variable coloration in its mainland distribution from Nicaragua to western Panamá. Many factors might account for the origin and maintenance of O. pumilio color polymorphism including selective pressure from multiple predators, mate choice based on visual cues, and genetic drift among the islands of the archipelago (Gehara et al. 2013). However, it is unclear how the subgenus Oophaga as a whole has evolved such extreme diversity in coloration and patterning without evident geographic barriers.

Molecular phylogenies show that Oophaga as lineage includes many cryptic species that underlie its diversity. For example, O. pumilio populations include two distinctive mitochondrial lineages, each of which contains at least one closely related congeneric species O. speciosa, O. arborea or O. vicentei (Hagemann & Pröhl 2007; Wang & Schaeffer 2008; Hauswaldt et al. 2011). Mitochondrial haplotypes from both lineages are found co-occurring across a wide zone along the Panamá and Costa Rica border. These populations present variable levels of admixture that also might account for their phenotypic diversity. This complex phylogeographic pattern suggests a series of dispersals and isolations leading to allopatric divergence and then subsequent admixture and introgression among O. pumilio and other species of Oophaga (Hagemann & Pröhl 2007, Wang & Schaeffer 2008, Hauswaldt et al. 2011).

Species boundaries are sometimes difficult to delineate and extreme polymorphism can hide cryptic speciation events. Recent observations in the Oophaga subgenus (Posso-Terranova & Andrés 2016b) suggest a complex pattern of diversification correlated with a structured landscape and strong shifts in climatic niches. For instance, Harlequin poison frogs from Colombia were previously recognized as three nominal species, O. histrionica, O. occulter, and O. lehmanni (Myers & Daly 1976). Oophaga histrionica has long been suspected to be a species complex (Silverstone 1975; Lotters et al. 1999), and has recently been proposed to include three new species evolving along a diversifying landscape (Posso-Terranova & Andrés 2016a). However, these new species descriptions need further morphological, behavioral, toxicological, and genomic characterization to be clearly validated. The Oophaga subgenus is composed of nine species, which have extraordinary morphological and chemical diversity (Daly & Myers 1967; Daly et al. 1978; Daly 1995; Saporito et al. 2007a). Most research has been conducted in O. pumilio, including a number of studies in population genetics, phylogeography, behavior, diet specialization, and chemical defenses (Saporito et al. 2007b; a; Richards-Zawacki et al. 2012; Gehara et al. 2013; Stynoski et al. 2014; Dreher et al. 2015). Comparative work in other Oophaga species would lend insight into interesting speciation and diversification processes.

We explored population structure of the highly polymorphic Little Devil (or Diabirito) poison frog, O. sylvatica, which has an inland distribution in the Chocoan rainforest and ranges from southwestern Colombia to northwestern Ecuador (Funkhouser 1956). Two types of population structures exist within this taxon: one composed of many phenotypically distinctive populations with low within-population color and pattern variability, and another structure containing a unique population with extreme within-population variability. The color and pattern observed in this latter population may represent either an admixture or hybridization zone, with individuals
showing intermediate color patterns that combine traits from the surrounding monotypic populations. Previous research in other Oophaga (i.e., O. histrionica and O. lehmanni) suggests that hybridization may promote color polymorphism (Medina et al. 2013). However, whether such similar admixture or hybridization mechanisms also promote color polymorphisms within O. sylvatica is unknown, as genetic studies have never been conducted with this species.

To investigate the population structure of O. sylvatica, we used a set of PCR amplicons including mitochondrial and nuclear markers, and a collection of genome wide single nucleotide polymorphisms (double digested RAD sequencing) from 13 geographically distinct populations (12 low phenotypic diversity and 1 highly polymorphic). The aims of our research were: 1) explore the relationship between individuals and their geographic localization, 2) estimate population structure and differentiation and 3) evaluate if admixture or hybridization could account for color diversity within and among populations.

MATERIALS AND METHODS

Sample collection

We sampled 13 populations of Oophaga sylvatica (N = 200 individuals) in July 2013 and July 2014 throughout its Ecuadorian distribution (Figure 1). Frogs were collected during the day with the aid of a plastic cup and stored individually in plastic bags with air and leaf litter for 3–8 h. In the evening the same day of capture, individual frogs were photographed in a transparent plastic box over a white background. Frogs were anesthetized with a topical application of 20% benzocaine to the ventral belly and euthanized by cervical transection. Tissues were preserved in either RNAlater (Life Technologies, Carlsbad, CA, USA) or 100% ethanol. Muscle and skeletal tissue were deposited in the amphibian collection of Centro Jambatu de Investigación y Conservación de Anfibios in Quito, Ecuador (Suppl. Table 1). The Institutional Animal Care and Use Committee of Harvard University approved all procedures (Protocol 15–02–233). Collections and exportation of specimens were done under permits (001–13 IC-FAU-DNB/MA, CITES 32 or 17 V/S) issued by the Ministerio de Ambiente de Ecuador. Samples from O. histrionica (2 individuals from Colombia: Choco: Quibdo, La Troje; voucher numbers TNHCFS4985 [longitude: -76.591, latitude: 5.728] and TNHCFS4987 [longitude: -76.591, latitude: 5.728]) and O. pumilio (6 individuals from 3 different populations, El Dorado, Vulture Point and Almirante acquired from the USA pet trade) were also treated using the same protocol. In order to protect the vulnerable O. sylvatica populations that are highly targeted by illegal poaching, specific GPS coordinates of frog collection sites can be obtained from the corresponding author.

DNA extraction and amplification

Liver or skin tissue stored in RNAlater was homogenized in using Trizol (Life Technologies) in tubes with 1.5 mm TriplePure Zirconium Beads (Bioexpress, Kaysville, UT, USA). After tissue homogenization, DNA was purified by a standard phenol/chloroform procedure followed by ethanol precipitation according to the Trizol manufacturer’s instructions.

Three mitochondrial gene regions (cytochrome oxidase subunit 1 (CO1 or cox1), 16S ribosomal DNA (16S), and 12S ribosomal DNA (12S) flanked with tRNAVal) and three nuclear gene regions (recombination-activating gene 1 (RAG-1), tyrosinase (TYR) and sodium-calcium exchanger 1 (NCX)), were PCR-amplified using the following sets of primers for each gene: CO1 (CO1a-H: 5’-AGTATAAGCGGTCTGGGTAGTC-3’ and CO1f-L: 5’-CCTGCAGGAGGAGGAGAYCC-3’) (Palumbi et al. 1991), 16S (16sar-L: 5’-CGCCTGATTATCAAAAC and 16sbr-H: 5’-CCGGTCTGAACTCAGATCACGT) (Palumbi et al. 1991), 12S-tRNAVal (MVZ59-L: 5’-ATAGCACTGAAAYGCTDAGTG and tRNAval-H: 5’-GGTGTAAGCGARAGGCTTAKTGTAAG) (Santos & Cannatella 2011), RAG-1 (Rag1_OopF: 5’-CCATGAATCCAGCGAGCT and Rag1_OopR: 5’-CACGGTCAAATGCTCTGGGAC) (Hauswaldt et al. 2011), TYR (TYR_Oosyl_F: 5’-AACTCATCATTGGTCCAAAT and TYR_Oosyl_R: 5’-GAAGTTCTCATTCCCGTAAGC), and NCX (NCX_Oosyl_F: 5’-ACTATCAAGAAACCAATGTTGAAA and NCX_Oosyl_R: 5’-TGTGCTGTTAGGTGACCC). NCX and TYR primers were designed from publicly available O. sylvatica sequences (Genbank accession numbers HQ290747 and HQ290927) using the free online tool provided by Life Technologies.
DNA was amplified in a 30 µL PCR reaction containing 10 ng of genomic DNA, 200 nM of each primer and 1X Accustart II PCR Supermix (Quanta Biosciences, Gaithersburg, MD, USA). The thermocycling profiles comprised an initial denaturation (3 min at 95°C), followed by 40 cycles of denaturation (30 s at 95°C), annealing (30 s) with specific temperature for each primer set (see below), and elongation (72°C) with duration specific to each primer set (see below), and a final elongation step (5 min at 72°C). Specific parameters for annealing temperature (A) and elongation time (E) for each primer set is as following: 16S (A = 50°C, E = 45 s), CO1 (A = 54°C, E = 45 s), 12S (A = 46°C, E = 60 s), RAG1 (A = 62°C, E = 45 s), TYR (A = 55°C, E = 40 s), NCX (A = 55°C, E = 80 s). PCR products were analyzed on an agarose gel and purified using the E.Z.N.A Cycle-Pure Kit following the manufacturer protocol (Omega bio-tek, Norcross, GA, USA). Purified PCR products were Sanger sequenced by GENEWIZ company (South Plainfield, NJ, USA).

**Analysis of mitochondrial and nuclear markers**

Raw sequence chromatograms for each gene set was aligned, edited and trimmed using Geneious (version 8.0.5, BioMatters Ltd., Auckland, New Zealand). Alleles of nuclear genes containing heterozygous sites were inferred using a coalescent-based Bayesian method developed in PHASE 2.1 (Stephens et al. 2001; Stephens & Donnelly 2003) as implemented in DnaSP v5.10.01 (Librado & Rozas 2009). Three independent runs of 10,000 iterations and burn-in of 10,000 generations were conducted to check for consistency across runs. All DNA sequences have been deposited in GenBank (Accession numbers: 12S: KX553997 - KX554204, 16S: KX554205 - KX554413, CO1: KX574018 - KX574226, pending for NCX, RAG1 and TYR).

Mitochondrial genes (12S-IRNA^Val, 16S, CO1) were considered as a single unit and concatenated in a matrix of 2084 bp for each individual of the different populations of O. sylvatica (N=199, one sample was excluded as it failed to amplify for 12S-IRNA^Val), and for O. histrionica (N=2) and O. pumilio (N=6) individuals. Diversity indices were calculated using DnaSP v5.10.01 and Arlequin 3.5 (Excoffier & Lischer 2010). Estimation of genetic distances between mtDNA sequences was tested under different models for nucleotide substitution and the HKY+I model was selected according to the Bayesian information criterion implemented in jModeltest2 (Guindon & Gascuel 2003; Darriba et al. 2012). We retained the Tajima and Nei model (Tajima & Nei 1984) to infer the haplotype network in Arlequin 3.5, which was a representative consensus of the results obtained with the other models tested. Minimum spanning networks (MSN) were generated using Gephi (Bastian et al. 2009).
2009), under the ForceAtlas2 algorithm (Jacomy et al. 2014) in default settings.

We assessed population differentiation by calculating the conventional F-statistics from haplotype frequencies ($F_{ST}$) and the statistics from haplotype genetic distances based on pairwise difference ($\Phi_{ST}$) using Arlequin 3.5. P values for $F_{ST}$ and $\Phi_{ST}$ were estimated after 10,000 permutations and significance threshold level was fixed at $p = 0.05$.

We performed a Bayesian assignment test on the nuclear dataset to estimate the number of genetic clusters as implemented in the program STRUCTURE 2.3.4. The software assumes a model with K populations (where K is initially unknown), and individuals are then assigned probabilistically to one or more populations (Pritchard et al. 2000). We ran the admixture model with correlated allele frequencies (Falush et al. 2003) for 50,000 burn-in and 100,000 sampling generations for K ranging from one to the number of sampled population plus three ($K = 1-16$) with 20 iterations for each value of K. We determined the number of clusters (K) that best described the data using the delta K method (Evanno et al. 2005) as implemented in Structure Harvester (Earl & vonHoldt 2012), which was inferred at K=5. Then we ran the same model for 500,000 burn-in generations and 750,000 MCMC, from two to six clusters ($K = 2-6$) and 20 iterations and analyzed the results using CLUMPAK (Kopelman et al. 2015).

ddRADseq library generation and sequencing

We constructed double-digested restriction-site-associated DNA sequencing (ddRAD) libraries on a subset of 125 samples following the protocol in Peterson et al. (2012). Samples include three specimens randomly drawn within each sampling site from the monomorphic populations and all the specimens from the putative admixture zone (Otokiki locality, see Figure 1). DNA was extracted from skin tissues preserved at -20°C in RNA later using the Nucleospin DNA kit (Macherey-Nagel, Bethlehem, PA). Genomic DNA of each sample (1 μg) was digested using 1 μL of EcoRI-HF (20,000 U/mL) and 1 μL Sphi-HF (20,000 U/mL) (New England Biolabs, Ipswitch, MA) following the manufacturer’s protocol. Digested samples were then cleaned with Agencourt Ampure XP beads (Beckman Coulter, Danvers, MA). Purified digested DNA (100ng) was ligated to double stranded adapters (biotin-labeled on P2 adapter) with a unique inline barcode using T4 DNA ligase (New England Biolabs, Ipswitch, MA) and purified with Agencourt Ampure XP beads. Barcoded samples were pooled and size-selected between 250 and 350 bp (326-426 bp accounting for the 76 bp adapter) using a Pippin Prep (Sage Science, Beverly, MA) 2% agarose gel cassette. Sized-selected fragments were purified with Dynabeads MyOne Streptavidin C1 (Life Technologies, Carlsbad, CA). Samples were divided in three independent libraries and amplified using Phusion High-Fidelity DNA polymerase (New England Biolabs, Ipswitch, MA) for 12 cycles. Libraries were then pooled and cleaned with Agencourt Ampure XP beads. Paired-end sequencing (125bp) was conducted on an Illumina HiSeq 2500 at the FAS Bauer Core Facility at Harvard University. Raw reads are available on the Sequence Read Archive (SRA) database.

| Species/Population | N  | S  | H  | Hd | K  | π   |
|-------------------|----|----|----|----|----|-----|
| O. sylvatica      | 199| 83 | 61 | 0.94863 | 11.66327 | 0.0056 |
| Durango           | 14 | 8  | 6  | 0.85714 | 3.34066  | 0.00161 |
| Lita              | 6  | 8  | 4  | 0.86667 | 3.8       | 0.00183 |
| Alto Tambo        | 6  | 12 | 5  | 0.93333 | 6.06667  | 0.00292 |
| Otokiki           | 83 | 46 | 29 | 0.95386 | 8.36879  | 0.00402 |
| San Antonio       | 14 | 22 | 6  | 0.6044  | 5.82418  | 0.00281 |
| Felfa             | 9  | 16 | 4  | 0.69444 | 3.83333  | 0.00185 |
| Quingüe           | 8  | 1  | 2  | 0.42857 | 0.42857  | 0.00021 |
| Cube              | 7  | 2  | 3  | 0.66667 | 0.7619   | 0.00039 |
| Cristóbal Colón   | 10 | 6  | 4  | 0.77778 | 2.15556  | 0.00104 |
| Simón Bolivar     | 13 | 9  | 5  | 0.75641 | 3         | 0.00137 |
| Puerto Quito      | 8  | 4  | 3  | 0.67857 | 1.85714  | 0.0009  |
| Santo Domingo     | 8  | 2  | 3  | 0.60714 | 0.67857  | 0.00033 |
| La Maná           | 13 | 0  | 1  | 0       | 0         | 0      |
| O. histrionica    | 2  | 0  | 1  | 0       | 0         | 0      |
| O. pumilio        | 6  | 30 | 4  | 0.86667 | 15.8      | 0.00697 |
| Overall           | 207| 163| 66 | 0.95239 | 13.31143  | 0.00753 |

Table 1. Summary of species and within-population diversity for the concatenated mitochondrial genes (12S-rRNA™, 16S, CO1). N, number of individuals sequenced; S, number of segregating sites; H, number of haplotypes; Hd, haplotype diversity; K, sequence diversity; π, nucleotide diversity.
ddRAD sequence analysis

Raw fastq reads were demultiplexed, quality filtered to discard reads with Phred quality score < 20 within a sliding window of 15% of read length and trimmed to 120bp using the process_radtags.pl command from the Stacks v1.35 pipeline (Catchen et al. 2013). ddRAD loci were constructed de novo with the Stacks denovo_map pipeline (parameters: m = 5, M = 3, n = 3) and then corrected for misassembled loci from sequencing errors using the corrections module (rxstacks, cstacks and sstacks), which applies population-based corrections. In order to minimize the effect of allele dropout that generally leads to overestimation of genetic variation (Gautier et al. 2013), we selected loci present in at least 75% of the individuals of each of the 13 sampled populations and generated population statistics and output files using Stacks population pipeline (parameters: r = 0.75, p = 13, m = 5).

To test for admixture, we estimated the number of genetic clusters as implemented in the program STRUCTURE 2.3.4 (Pritchard et al. 2000). The data set was filtered using the Stacks population pipeline (parameters: r = 0.75, p = 13, m = 5) to contain only loci present in at least 75% of each of the 13 sampled localities. We retained only one SNP per RAD locus to minimize within-locus linkage. The data set contains 3,785 SNPs from concatenated pair-ended reads. We ran the admixture model with correlated allele frequencies (Falush et al. 2003) for 100,000 burn-in generations and 1,000,000 MCMC, for K = 2-16 with 10 iterations for each cluster. We determined the number of clusters K that best described the data (Evanno et al. 2005) as implemented in Structure Harvester (Earl & vonHoldt 2012) and analyzed the results using CLUMPAK (Kopelman et al. 2015).

We used a Discriminant Analysis of Principal Components (DAPC) to identify clusters of genetically related individuals, implemented in the R package adegenet v2.0.1 (Jombart et al. 2010; Jombart & Ahmed 2011). This multivariate method is suitable for analyzing large number of SNPs, providing assignment of individuals to groups and a visual assessment of between-population differentiation. It does not rely on any particular population genetics model and DAPC is free of assumption about Hardy-Weinberg equilibrium or linkage equilibrium (Jombart et al. 2010). The data set was built with the Stacks populations pipeline and is composed of 3,785 SNPs (5955 alleles) present in at least 75 % of the individuals from each of the 13 populations. To avoid over-fitting of the discriminant functions we performed stratified cross-validation of DAPC using the function xvalDapc from adegenet and retained 20 principal components, which gave the highest mean success and the lowest Root Mean Squared Error (MSE).

**Figure 2.** Haplotype network of 66 unique haplotypes of concatenated mitochondrial genes (12S-IRNA, 16S, CO1) of Oophaga sylvatica, O. histrionica and O. pumilio (2084 bp). Circles indicate haplotypes, with the area being proportional to the number of individuals sharing that haplotype. Colors refer to the geographic origin of the population and the pie charts represent the percentage of each population sharing the same haplotype. Line thickness between haplotypes is proportional to the inferred mutational steps (or inferred intermediate haplotypes). Inferred numbers of mutational steps are shown inside circles along the line when greater than four steps.
We also generated consensus sequences of the 125 individuals grouped by geographical sampling localities to assess their relationship at the population level. We used Bayesian inferences as implemented in Mr Bayes 3.2.6 (Ronquist et al. 2012) under Geneious to build a phylogenetic tree from a matrix of 41,779 SNPs (fixed within but variable among populations), with O. pumilio and O. histrionica as an outgroup. We used the default settings under the HKY85 + gamma model and ran a single MCMC with four chains (0.02 heated chain temp) for 1,100,000 generations, out of which the first 100,000 were discarded as burn-in.

RESULTS

Mitochondrial diversity and structure

After alignment of all 207 individuals, sequences show 163 segregating sites (S) that compose 66 haplotypes (H) (Table 1 and Figure 2). Among the 13 populations of O. sylvestica sampled, genetic diversity is relatively high, with 83 segregating sites, 61 haplotypes, and a mean haplotype diversity (Hd) of 0.948. Both O. histrionica and O. pumilio are well separated from O. sylvestica and link the network by two distinct branches (Figure 2). These branches constitute two remote clusters, one composed of haplotypes from Otokiki and Felfa populations that link to O. histrionica, and one composed of haplotypes from the San Antonio population that link to O. pumilio. Both branches link to the same node, a haplotype of one individual from Durango. Six short branches (2 to 4 inferred mutations) radiate from this central node: two are composed of private haplotypes from the Durango and San Antonio populations, and the other four have mixed origins belonging to the northern populations of Otokiki, Durango, Lita, Alto Tambo and San Antonio. These populations show the highest sequence (K) and nucleotide (π) diversity, as well as the highest haplotype diversity (except San Antonio population for Hd) (Table 1). One of the mixed clusters includes haplotypes from Durango, Lita, Alto Tambo and Otokiki populations; another includes haplotypes from Alto Tambo and Otokiki, and two different clusters are composed of haplotypes from the Otokiki and Lita populations. All of these clusters have inter-haplotype distances ranging from 1 to 4 inferred mutational steps. With 29 haplotypes, Otokiki is the most genetically diverse population (Table 1 and Figure 2), in addition to exhibiting the most variable color patterning (Figure 1B). Most haplotypes are unique to the Otokiki population, except for 6 that are shared with the populations of Alto Tambo, Lita and Durango. These are also the geographically the closest populations to Otokiki (Figure 1A). Genetically, Lita and Alto Tambo seem weakly separated from Otokiki, with the Lita population having no unique haplotype and only two out of five unique haplotypes in the Alto Tambo population. The southern and northern populations are separated by a unique haplotype from the Felfa population. Southern populations show less genetic diversity than northern populations, where 67 individuals collapsed in 15 haplotypes (Figure 2). One branch is composed of a small group of 3 haplotypes from Puerto Quito, Simón Bolívar and Cristóbal Colón. The second branch leads to one haplotype including 38 individuals from 6 populations: Quingüe, Cube, Simón Bolívar, Puerto Quito, Santo Domingo and La Maná. Radiating from this haplotype are 11 unique haplotypes (1 to 4 individuals each) with mostly 1 inferred mutational step. While closely related to the more southern populations, frogs from the Cristóbal Colón population do not share any haplotype with the rest of the southern populations.
### Table 2. Summary of species and within-population diversity for the concatenated nuclear genes (RAG-1, TYR and NCX).

| Species/Population | N  | S  | H  | Hd  | K  | π  |
|--------------------|----|----|----|-----|----|----|
| **O. sylvatica**   | 400| 10 | 16 | 0.584 | 1.51112 | 0.00067 |
| Durango            | 28 | 6  | 7  | 0.64815 | 2.32275 | 0.00103 |
| Lita               | 12 | 6  | 3  | 0.43939 | 1.68182 | 0.00075 |
| Alto Tambo         | 12 | 5  | 2  | 0.30303 | 1.51515 | 0.00067 |
| Otokiki            | 166| 7  | 9  | 0.53428 | 1.59927 | 0.00071 |
| San Antonio        | 28 | 5  | 3  | 0.57407 | 1.89153 | 0.00084 |
| Felfa              | 20 | 5  | 3  | 0.35789 | 1.53684 | 0.00068 |
| Quingüe            | 16 | 4  | 6  | 0.825  | 1.29167 | 0.00057 |
| Cube               | 14 | 2  | 3  | 0.67033 | 0.97802 | 0.00043 |
| Cristóbal Colón    | 20 | 1  | 2  | 0.1   | 0.1    | 0.00004 |
| Simón Bolívar      | 26 | 0  | 1  | 0     | 0      | 0      |
| Puerto Quito       | 16 | 2  | 3  | 0.34167 | 0.35833 | 0.00016 |
| Santo Domingo      | 16 | 4  | 6  | 0.8   | 1.23333 | 0.00055 |
| La Maná            | 26 | 3  | 4  | 0.76  | 1.14769 | 0.00051 |
| **O. histrionica** | 4  | 1  | 2  | 0.5   | 0.5    | 0.00022 |
| **O. pumilio**     | 12 | 7  | 6  | 0.84848 | 2.21212 | 0.00098 |
| **Overall**        | 416| 23 | 24 | 0.61527 | 2.06418 | 0.00092 |

**Population differentiation using mtDNA**

Levels of population differentiation are relatively high in *O. sylvatica*, with a mean $F_{ST}$ of 0.220 (ranging from -0.011 to 0.705) and a mean $\Phi_{ST}$ of 0.609 (ranging from 0.016 to 0.915), suggesting northern populations are well differentiated from southern populations. In the northern populations we observe very low values of both $F_{ST}$ and $\Phi_{ST}$ between the populations of Durango, Lita, Alto Tambo and Otokiki. These values are statistically non-significant ($p$-value > 0.05 after 10,000 permutations) for Otokiki versus Lita and Alto Tambo (both $F_{ST}$ and $\Phi_{ST}$), and for Durango versus Alto Tambo and Lita versus Alto Tambo ($F_{ST}$ only) (Suppl. Table 2). In the southern populations, frogs from Quingüe, Cube, Simón Bolívar, Puerto Quito, Santo Domingo and La Maná are genetically similar in haplotype clustering (Figure 3) and this observation is confirmed by low $F_{ST}$ and $\Phi_{ST}$ and non-significant $p$-values (Suppl. Table 2). Finally, based on both $F_{ST}$ and $\Phi_{ST}$ values, populations from San Antonio, Felfa and Cristóbal Colón appear to be different from every other northern and southern population.

**Nuclear DNA diversity and population structure**

Nuclear sequences (RAG-1, TYR, and NCX) were phased and analyzed separately. Across all three *Oophaga* species in this study, the NCX gene presents 10 segregating sites and 15 haplotypes, TYR has 9 segregating sites and 9 haplotypes, and RAG-1 has 4 segregating sites and 5 haplotypes (Suppl. Table 3). Among the nuclear genes, NCX is the most variable in *O. sylvatica*, with 7 segregating sites and 12 haplotypes, whereas RAG-1 has 2 segregating sites and 3 haplotypes, and TYR has only 1 segregating sites and 2 haplotypes. We then combined and phased the three sets of nuclear markers in a unique matrix for every individual (Table 2). This slightly increases the number of segregation sites ($S = 10$) and haplotypes ($H = 16$), as well as diversity indices for our *O. sylvatica* group. Similar to mitochondrial genes, sequence ($K$) and nucleotide ($\pi$) diversity are the highest for northern populations (Durango, San Antonio, Lita, Otokiki, Felfa, and Alto Tambo), but haplotype diversity ($Hd$) is higher in a subset of the southern group (Quingüe, Puerto Quito, and Santo Domingo). Within the minimum spanning network of haplotypes, inferred distances between nodes are very short, ranging from 1 to 5 mutation steps (Figure 4). At the species level, we can observe that *O. histrionica* connects *O. pumilio* and *O. sylvatica* nodes, which was not observed in the mitochondrial network. Interestingly, haplotype linkage seems to follow a geographic North to South gradient. Starting from *O. pumilio* through *O. histrionica*, *O. sylvatica* haplotypes are first only composed of northern populations and then, are gradually shared between northern and southern populations, and finally a unique haplotype includes every population of *O. sylvatica*.

**Population structure based on ddRAD markers**

A structure plot was drawn using Bayesian inferences with our ddRAD data (a subset of 125 individuals of *O. sylvatica*). The optimal number of clusters inferred by Evanno’s method was $K = 3$ (see Figure 5 for colors assigned to clusters). In *O. sylvatica* we can observe a clear genetic structure. One genetic cluster (blue) represents mostly populations from the northernmost part of the range (San Antonio, Lita, Alto Tambo, Durango and Otokiki). A second genetic cluster (orange) appears mostly in populations from the Southern part of the range (Felfa, Cristóbal Colón, Simón Bolívar, Quingüe, Cube,
with mixe
6
Durango
overlapping individuals from
discriminant analysis (DA)
each individual using the
population.
other and individual
In the south, populations are
Otokiki, Tambo and Durango overlap with individuals
north, individuals from the populations of Lita, Alto
follow the north/south split observed previously. In the

Information Criterion
number of cluster
(DAPC) on the ddRAD data
Discriminant Analysis of Principal Components

populations with intermediate range (Felfa and
proportions in every population. In addition, two
third
Puerto Quito, Santo Domingo and La Maná)
are indicated inside circles along the line when greater than four steps.
the inferred mutational steps (haplotypes). Inferred numbers of mutational steps
are indicated inside circles along the line when greater than four steps.

Puerto Quito, Santo Domingo and La Maná). Finally, a
third genetic cluster (purple) is present in small proportions in every population. In addition, two
populations with intermediate range (Felfa and Cristóbal Colón) have admixed proportion of both main
clusters (blue and orange).

Discriminant Analysis of Principal Components
(DAPC) on the ddRAD data
We used 3,785 SNPs from the ddRAD dataset
to conduct a DAPC analysis (Figure 6). The optimal
number of clusters inferred under the Bayesian
Information Criterion was K = 2. The two clusters
follow the north/south split observed previously. In the
north, individuals from the populations of Lita, Alto
Tambo and Durango overlap with individuals from
Otokiki, but San Antonio is well separated from them.
In the south, populations are separated from each other and individuals did not overlap with any other
population. By plotting the membership probability of
each individual using the compoplot function with two
discriminant analysis (DA), we can see that
overlapping individuals from Lita, Alto Tambo, Durango and Otokiki, suggestive of admixture (Figure
6). In addition, some individuals from the southern
populations of Cube and La Maná have been assigned
with mixed proportions to both populations.

Phylogenetic relationships derived from ddRAD data
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Little Devil poison frog population genetics
Using consensus sequences for
each geographical locality, we built a
population-based tree using Bayesian
inferences and setting O. pumilio as the
outgroup species (Figure 7). The San
Antonio population, which is located in
the northern-most part of the
Ecuadorian range, has diverged before
the rest of the O. sylvatica Ecuadorian
populations. Two major branches split
other populations with moderate support
(0.93 posterior probability), grouping
Alto Tambo, Lita, Durango and Otokiki
in one cluster, and Felfa, Cristóbal
Colón, Simón Bolívar, Puerto Quito,
Cube, Quingüe, Santo Domingo and La
Maná in another cluster.

DISCUSSION
In this study we investigated
genetic structure among populations of
Oophaga sylvatica in Ecuador. Our
results show that the genetic structure of
the populations follows their
geographical distribution and is split in
two main lineages, distributed in the
northern and the southern part of their
range. The existence of two distinct
lineages is supported by mitochondrial
haplotype network (Figure 2), Bayesian assignment of
individuals using Structure (Figure 5), DAPC analysis
(Figure 6) and a phylogeny (Figure 7) using ddRAD
data. Within each lineage, most populations share
mitochondrial haplotypes (with the exception of San
Antonio, Felfa and Cristóbal Colón) and have low
genetic differentiation (low or non significant F_{ST}
values). We hypothesize that gene flow from recent
range expansions might be driving the color diversity
in these frogs.

The northern lineage (San Antonio, Lita, Alto
Tambo, Durango and Otokiki) presents higher
mitochondrial diversity with a large amount of unique
haplotypes within populations. We observe that four of
these populations (San Antonio, Lita, Alto Tambo and
Durango) have unique haplotypes, and three
(Durango, Lita and Alto Tambo) share haplotypes with
the polymorphic population at Otokiki. These latter
populations are weakly differentiated based on genetic
data and are geographically close (distances from
Otokiki are approximately 4km, 5km and 20 km to Alto
Tambo, Lita and Durango, respectively). Otokiki
individuals present all color patterns found in the
surrounding populations in addition to intermediate
patterns (Figures 1A and 1B). The DAPC plot
highlights a genetic overlap between some individuals
of Otokiki and the three populations from Lita, Alto
Tambo and Durango; some individuals were assigned
with mixed proportions to these different groups.
probabilities for 125 individual frogs as inferred by STRUCTURE for K=3 clusters, each color

accentuated by decades of variation.

addition, this is the mitochondrial species and Dendrobatidae are not known to be migratory species, the low levels of differentiation observed at the mitochondrial markers is unlikely to occur through gene flow among most of these populations. In addition, this contrasting pattern of low genetic variation but high phenotypic diversity among geographically distant population suggests that the southern clade radiation has probably occurred relatively recently. This genetic pattern may be accentuated by decades of population bottlenecks and genetic drift, as southern populations have been subject to strong isolation due to human intervention and habitat destruction. Indeed, a similar pattern of recent expansion and isolation with reduced gene flow was proposed as a model to explain the phenotypic diversification within the southern lineage of O. pumilio, among the islands of Panamá (Gehara et al. 2013).

Repeated phylogeographical patterns of diversification among amphibians and reptiles across Northwestern Ecuador was recently described by Arteaga and colleagues (2016) and suggests that diversification follows thermal elevation gradients among the Chocoan region and the Andes. There are also several natural barriers to dispersion, including the Mira River in the northern most part of the O. sylvatica range. The Mira River geographically separates the San Antonio population from the rest of the northern lineage, which may explain its genetic differentiation. The Esmeraldas River also separates the distribution range of O. sylvatica in two main zones: the populations of San Antonio, Lita, Alto Tambo, Durango, Otokiki, Felfa and Cristóbal Colón in the north, and the populations of Simón Bolívar, Puerto Quito, Cube, Quingüé, Santo Domingo and La Maná in the south. Arteaga and colleagues (2016) have identified similar geographical patterns of differentiation for other frog species, such as Pristimantis nietoi and P. walker, and populations of the snake species Bothrops punctatus near the Esmeraldas and Guayllabamba Rivers (Arteaga et al. 2016). These large rivers may have acted as barriers precluding or reducing gene flow between the northern and southern lineages. Nevertheless, Felfa and Cristóbal Colón populations, distributed north of the Esmeraldas River, present conflicting genetic data suggesting that the barrier did not isolated these two lineages completely. Both populations have admixed proportions of the two lineages with high proportion of the southern lineage (Figure 5). Still, mitochondrial data support Felfa’s relationship to the northern lineage, but Cristóbal Colón likely belong to the southern lineage (Figure 2). Also, this genetic pattern observed for Cristóbal Colón could be supported by the presence of the Toisán Mountains extending in this area from west to east, which might have acted as a barrier, isolating the Cristóbal Colón population from the rest of the northern lineage.

Figure 5. Population structure inferred from ddRAD data. Bar plots showing Bayesian assignment probabilities for 125 individual frogs as inferred by STRUCTURE for K=3 clusters, each color depicting one of the putative clusters.

(Figure 6). Taken together, our results show a close genetic interaction between populations presenting low diversity and intermediate phenotypes among populations with overlapping distributions. There at least two scenarios that could explain these patterns. First, these features support the presence of an admixture zone occurring within Otokiki, which in turn promotes the dramatic color diversity of this population. Alternatively, Otokiki could be a large source population where surrounding populations represent recent expansions with less genetic and phenotypic diversity due to founder effects or other selection pressures. Although outside the scope of the current study, more analyses on population structure with more sampling in the northern populations is needed to disentangle these alternative hypotheses.

The southern lineage shows little mitochondrial diversity, in which a few variants radiate from a common haplotype shared by a majority of individuals across different populations (Figure 2). This lineage shows great color diversity among populations and little variation within, suggesting that variation in mitochondrial markers is ineffective to resolve the observed phenotypic diversity. Furthermore, these populations are geographically distant from each other (from 20km between Simón Bolívar and Puerto Quito to as much as 190km between Quingüé and La Maná), have low but significant FST values from ddRAD data (Suppl. Table 4) and are genetically well differentiated on the DAPC analysis. Since amphibians usually have low dispersal abilities (Vences & Wake 2007; Vences & Köhler 2008; Zeisset & Beebee 2008) and Dendrobatidae are not known to be migratory species, the low levels of differentiation observed at the mitochondrial markers is unlikely to occur through gene flow among most of these populations. In addition, this contrasting pattern of low genetic variation but high phenotypic diversity among geographically distant population suggests that the southern clade radiation has probably occurred relatively recently. This genetic pattern may be accentuated by decades of population bottlenecks and
A critical question is raised from this study: How is high phenotypic diversity in coloration and patterning maintained in the Otokiki population? A wide range of putative predators such as birds, reptiles or arthropods with distinct visual abilities and predatory strategies might act on color selection and diversity among O. sylvatica populations (Maan & Cummings 2012; Crothers & Cummings 2013; Dreher et al. 2015). In addition, sexual selection through non-random courtship within color morphs has been reported in O. pumilio (Reynolds & Fitzpatrick 2007; Maan & Cummings 2008) and Ranitomeya (Twomey et al. 2014), and can possibly promote diversification of color among populations, while decreasing the variation within. Although this phenomenon seems highly dependent on the environmental context and genetic background of the frogs (Richards-Zawacki et al. 2012; Meuche et al. 2013; Medina et al. 2013; Twomey et al. 2014). A recent study reported an example of hybridization promoting new coloration and patterning between two close species O. histrionica and O. lehmanni (Medina et al. 2013). This work also suggests complex roles played by sexual selection as hybrid females present non-random sexual preferences depending on the combination of available males. If similar processes occur in O. sylvatica at Otokiki, the extreme polymorphism within this admixture zone represents a unique opportunity to

Figure 6. DAPC scatterplot for ddRAD data. The scatterplot shows the first two principal components of the DAPC of data generated with 3,785 SNPs. Geographic samples are represented in different colors and inertia ellipses according to Figure 1, with individuals shown as dots. Lines between groups represent the minimum spanning tree based on the (squared) distances, and show the actual proximities between populations within the entire space. Right inset shows the inference of the number of clusters using the Bayesian information criterion (BIC). The chosen number of clusters corresponds to the inflexion point of the BIC curve (K=2). Left inset shows the number of PCA eigenvalues retained in black, and how much they accounted for variance. The bottom graph represents the membership probability of each individual to one or more populations. Geographic populations are represented with the same colors as in the DAPC plot.
further test the balance of selective pressure on the evolution of an aposematic trait by selection through predators or by conspecifics through mate choice.

**Summary**

By evaluating mitochondrial DNA variation and genome-wide SNPs, we have gained four important insights about *O. sylvatica*: (1) The Ecuadorian populations of *O. sylvatica* are composed of two clades that reflect their geographic distribution. Further behavioral, ecological and morphological information is required to determine whether the two geographical lineages observed in *Oophaga sylvatica* represent one polytypic or distinct species. A combination of climatic gradient and structured landscape generating geographic barriers to gene flow could explain the complex patterns of diversification observed in *Oophaga* and some other Dendrobatidae (Arteaga et al. 2016; Posso-Terranova & Andrés 2016ab). (2) The northern and southern populations show different amounts of structure, which may reflect more recent range expansions in the south. (3) Phenotypic variation in *O. sylvatica* has evolved faster than mitochondrial mutations can fix in the population. (4) A highly polymorphic population (Otoki) exists, which provides a unique opportunity for testing hypotheses about the selective pressures shaping aposematic traits. We hypothesize that this polymorphic population arose from either gene flow between phenotypically divergent populations at secondary contact zones or through a range expansion of the polymorphic Otoki population into surrounding regions. More data is needed to distinguish between these alternative scenarios.

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**Data Accessibility**

**DNA sequences:** Genbank accession numbers: 12S: KX553997 - KX554204, 16S: KX554205 - KX554413, CO1: KX574018 - KX574226, pending for NCX, RAG1 and TYR; **SRA for ddRADseq reads:** SRP078453

**Author Contributions:** ABR, JCS, LAC, and LAO designed the research; ABR, JCS, EET, LAC, and LAO collected samples in the field; ABR, BCC, and SNC performed the laboratory research; ABR analyzed data; ABR and LAO wrote the paper with contributions from all authors.
References

Arteaga A, Pyron RA, Peñaflie N et al. (2016) Comparative Phylogeography Reveals Cryptic Diversity and Repeated Patterns of Cladogenesis for Amphibians and Reptiles in Northwestern Ecuador (R Castiglia, Ed.). PLOS ONE, 11, e0151746.

Bastian M, Heymann S, Jacomy M (2009) Gephi: An Open Source Software for Exploring and Manipulating Networks. Third International AAAI Conference on Weblogs and Social Media, 361–362.

Benson WW (1971) Evidence for the Evolution of Unpalatability Through Kin Selection in the Heliconiinae (Lepidoptera). The American Naturalist, 105, 213.

Brodie ED (2008) An Experimental Study of Aposematic Coloration in the Salamander Plethodon jordani. Society, 1976, 59–65.

Brodie III ED (1993) Differential Avoidance of Coral Snake Banded Patterns by Free-Ranging Avian Predators in Costa Rica. Evolution, 47, 227.

Brodie III ED, Janzen F (1995) Experimental Studies of Coral Snake mimicry: Generalized Avoidance of Ringed Snake Patterns by Free-Ranging Avian Predators. Functional Ecology, 9, 186–190.

Brusa O, Bellati A, Meuche I, Mundy NI, Pröhl H (2013) Divergent evolution in the polymorphic granular poison-dart frog, Oophaga granulifera: genetics, coloration, advertisement calls and morphology (L Rocha, Ed.). Journal of Biogeography, 40, 394–408.

Catchen J, Hohenlohe PA, Bassham S, Amores A, Cresko WA (2013) Stacks: An analysis tool set for population genomics. Molecular Ecology, 22, 3124–3140.

Chouteau M, Arias M, Joron M (2016) Warning signals are under positive frequency-dependent selection in nature. Proceedings of the National Academy of Sciences of the United States of America, 113, 201519216.

Corneault AA, Noonan BP (2011) Spatial variation in the fitness of divergent aposematic phenotypes of the poison frog, Dendrobates tinctorius. Journal of Evolutionary Biology, 24, 1374–9.

Crothers LR, Cummings ME (2013) Warning Signal Brightness Variation: Sexual Selection May Work under the Radar of Natural Selection in Populations of a Polytypic Poison Frog. The American Naturalist, 181, E116–E124.

Crothers LR, Cummings ME (2015) A multifunctional warning signal behaves as an agonistic status signal in a poison frog. Behavioral Ecology, 00, 1–9.

Daly J (1995) The chemistry of poisons in amphibian skin. Proceedings of the National Academy of Sciences of the United States of America, 92, 9–13.

Daly JW, Brown GB, Mensah-Dwumah M, Myers CW (1978) Classification of skin alkaloids from neotropical poison-dart frogs (Dendrobatidae). Toxicon, 16.

Daly JW, Myers CW (1967) Toxicity of Panamanian poison frogs (Dendrobates): some biological and chemical aspects. Science, 156, 970–3.

Darriba D, Taboada GL, Doallo R, Posada D (2012) jModelTest 2: more models, new heuristics and parallel computing. Nature Methods, 9, 772–772.

Dreh CE, Cummings ME, Pröhl H (2015) An Analysis of Predator Selection to Affect Aposematic Coloration in a Poison Frog Species (D Osorio, Ed.). PLOS ONE, 10, e0130571.

Dumbacher JP, Beehler BM, Spande TF, Garraffo HM, Daly JW (1992) Homobatrachotoxin in the genus Phyllostomus: chemical defence in birds? Science, 258, 799–801.

Dumbacher JP, Spande TF, Daly JW (2000) Batrachotoxin alkaloids from passerine birds: a second toxic bird genus (Hirundo kowaldi) from New Guinea. Proceedings of the National Academy of Sciences of the United States of America, 97, 12970–5.

Earl DA, vonHoldt BM (2012) STRUCTURE HARVESTER: A website and program for visualizing STRUCTURE output and implementing the Evanno method. Conservation Genetics Resources, 4, 359–361.

Evanno G, Regnaut S, Goudet J (2005) Detecting the number of clusters of individuals using the software STRUCTURE: A simulation study. Molecular Ecology, 14, 2611–2620.

Excoffier L, Lischer HEL (2010) Arlequin suite ver 3.5: A new series of programs to perform population genetics analyses under Linux and Windows. Molecular Ecology Resources, 10, 564–567.

Falush D, Stephens M, Pritchard J (2003) Inference of population structure using multilocus genotype data: linked loci and correlated allele frequencies. Genetics, 164, 1567–1587.

Funk WC, Murphy MA (2010) Testing evolutionary hypotheses for phenotypic divergence using landscape genetics. Molecular ecology, 19, 427–30.

Gautier M, Gharbi K, Cezard T et al. (2013) The effect of RAD allele dropout on the estimation of genetic variation within and between populations. Molecular Ecology, 22, 3165–3178.

Gehara M, Summers K, Brown JL (2013) Population expansion, isolation and selection: novel insights on the evolution of color diversity in the strawberry poison frog. Evolutionary Ecology, 27, 797–824.

Guindon S, Gascuel O (2003) A Simple, Fast, and Accurate Algorithm to Estimate Large Phylogenies by Maximum Likelihood. Systematic Biology, 52, 696–704.

Haber M, Cerfeda S, Carbone M et al. (2010) Coloration and defense in the nudibranch gastropod Hypselodoris fontandraui. Biological Bulletin, 218, 181–188.

Hauswaldt JS, Ludewig A-K, Vences M, Pröhl H (2011) Widespread co-occurrence of divergent mitochondrial haplotype lineages in a Central American species of poison frog (Oophaga pumilio). Journal of Biogeography, 38, 711–726.

Hoogmoed M, Avila-Pires T (2012) Inventory of color polymorphism in populations of Dendrobates galactonotus (Anura: Dendrobatidae), a poison frog endemic to Brazil. Phylomedusa, 11, 95–115.

Howard RR, Brodie ED (1973) A Batesian Mimetic Complex in Salamanders: Responses of Avian Predators. Herpetologica, 29, 33–41.

Jacomy M, Venturini T, Heymann S, Bastian M (2014) ForceAtlas2, a Continuous Graph Layout Algorithm for Handy Network Visualization Designed for the Gephi Software (MR Muldoon, Ed.). PLoS ONE, 9, e89679.

Jiggins CD, McMillan WO (1997) The genetic basis of an adaptive radiation: warning colour in two Heliconius species. Proceedings of the Royal Society B: Biological Sciences, 264, 1167–1175.
Roland et al. final version for bioRxiv preprint

Little Devil poison frog population genetics

colour: Multivariate species delimitation and conservation of Harlequin poison frogs.

Posso-Terranova A, Andrés JA (2016b) Complex niche divergence underlies lineage diversification in Oophaga poison frogs. Journal of Biogeography.

Pritchard JK, Stephens M, Donnelly P (2000) Inference of population structure using multilocus genotype data. Genetics, 155, 945–99.

Przečzek K, Mueller C, Vamosi SM (2008) The evolution of aposematism is accompanied by increased diversification. Integrative zoology, 3, 149–156.

Reichstein T, von Euw J, Parsons J, Rothschild M (1968) Heart poisons in the Monarch butterfly. Science, 161, 861–866.

Reynolds RG, Fitzpatrick BM (2007) Assortative mating in poison-dart frogs based on an ecologically important trait. Evolution, 61, 2253–9.

Richards-Zawacki CL, Wang IJ, Summers K (2012) Mate choice and the genetic basis for colour variation in a polymorphic dart frog: inferences from a wild pedigree. Molecular ecology, 21, 3979–92.

Ronquist F, Teslenko M, Van Der Mark P et al. (2012) Mrbayes 3.2: Efficient bayesian phylogenetic inference and model choice across a large model space. Systematic Biology, 61, 539–542.

Ruxton GD, Sherratt TN, Speed MP (2004) Avoiding Attack: The Evolutionary Ecology of Crypsis, Warning Signals and Mimicry. Oxford biology, 17, 249.

Santos JC, Baquero M, Barrio Amorós CL et al. (2014) Aposematism increases acoustic diversification and speciation in poison frogs. Proceedings of the Royal Society B: Biological Sciences, 281, 20141761.

Santos JC, Cannatella DC (2011) Phenotypic integration emerges from aposematism and scale in poison frogs. Proceedings of the National Academy of Sciences of the United States of America, 108, 6175–6180.

Santos JC, Coloma LA, Cannatella DC (2003) Multiple, recurring origins of aposematism and diet specialization in poison frogs. Proceedings of the National Academy of Sciences of the United States of America, 100, 12792–7.

Santos JC, Coloma LA, Summers K et al. (2009) Amazonian Amphibian Diversity Is Primarily Derived from Late Miocene Andean Lineages (C Moritz, Ed.). PLoS Biology, 7, e1000056.

Saporito RA, Donnelly MA, Jain P et al. (2007a) Spatial and temporal patterns of alkaloid variation in the poison frog Oophaga pumilio in Costa Rica and Panama over 30 years. Toxicon, 50, 757–778.

Saporito RA, Zuercher R, Roberts M, Gerow KG, Donnelly MA (2007b) Experimental Evidence for Aposematism in the Dendrobatid Poison Frog Oophaga pumilio. Copeia, 2007, 1006–1011.

Silverstone PA (1975) A Revision of the Poison-Arrow Frogs of the Genus Dendrobates Wagler. Natural History Museum of Los Angeles County, Science Bulletin, 21, 1–55.

Stephens M, Donnelly P (2003) A Comparison of Bayesian Methods for Haplotype Reconstruction from Population Genotype Data. American journal of human genetics, 73, 1162–1169.

Stephens M, Smith NJ, Donnelly P (2001) A new statistical method for haplotype reconstruction from population data. American journal of human genetics, 68, 978–989.
Stynoski J (2012) Behavioral Ecology of Parental Care in a Dendrobatid Frog (Oophaga Pumilio).

Stynoski JL, Torres-Mendoza Y, Sasa-Marín M, Saporito RA (2014) Evidence of maternal provisioning of alkaloid-based chemical defenses in the strawberry poison frog Oophaga pumilio. *Ecology, 95*, 587–593.

Summers K, Bermingham E, Weigt L, McCafferty S, Dahlstrom L (1997) Phenotypic and genetic divergence in three species of dart-poison frogs with contrasting parental behavior. *The Journal of heredity, 88*, 8–13.

Summers K, Clough ME (2001) The evolution of coloration and toxicity in the poison frog family (Dendrobatidae). *Proceedings of the National Academy of Sciences of the United States of America, 98*, 6227–32.

Symula R, Schulte R, Summers K (2001) Molecular phylogenetic evidence for a mimetic radiation in Peruvian poison frogs supports a Mullerian mimicry hypothesis. *Proceedings of the Royal Society B: Biological Sciences, 268*, 2415–2421.

Tajima F, Nei M (1984) Estimation of evolutionary distance between nucleotide sequences. *Mol Biol Evol, 1*, 269–285.

Tullrot A, Sundberg P (1991) The conspicuous nudibranch Polycera quadrilineata: aposematic coloration and individual selection. *Animal Behaviour, 41*, 175–176.

Twomey E, Vestergaard JS, Summers K (2014) Reproductive isolation related to mimetic divergence in the poison frog *Ranitomeya imitator*. *Nature Communications, 5*, 4749.

Twomey E, Vestergaard JS, Venegas PJ, Summers K (2016) Mimetic Divergence and the Speciation Continuum in the Mimic Poison Frog Ranitomeya imitator. *The American Naturalist, 187*, 205–224.

Twomey E, Yeager J, Brown JL et al. (2013) Phenotypic and Genetic Divergence among Poison Frog Populations in a Mimetic Radiation (P Michalak, Ed.). *PLoS ONE, 8*, e55443.

Vences M, Köhler J (2008) Global diversity of amphibians (Amphibia) in freshwater. *Hydrobiologia, 595*, 569–580.

Vences M, Wake DB (2007) Speciation, species boundaries and phylogeography of amphibians. *Amphibian biology, 7*, 2613–2671.

Wang IJ, Summers K (2010) Genetic structure is correlated with phenotypic divergence rather than geographic isolation in the highly polymorphic strawberry poison-dart frog. *Molecular ecology, 19*, 447–58.

Wollenberg KC, Lötters S, Mora-Ferrer C, Veith M (2008) Disentangling composite colour patterns in a poison frog species. *Biological Journal of the Linnean Society, 93*, 433–444.

Zeisset I, Beebee TJC (2008) Amphibian phylogeography: a model for understanding historical aspects of species distributions. *Heredity, 101*, 109–119.