Genomic divergence across ecological gradients in the Central African rainforest songbird (Andropadus virens)

Ying Zhen1,2 | Ryan J. Harrigan2 | Kristen C. Ruegg2,3 | Eric C. Anderson4 | Thomas C. Ng5 | Sirena Lao2 | Kirk E. Lohmueller1,6 | Thomas B. Smith1,2

1Department of Ecology and Evolutionary Biology, University of California, Los Angeles, CA, USA
2Center for Tropical Research, Institute of Environment and Sustainability, University of California, Los Angeles, CA, USA
3Department of Ecology and Evolutionary Biology, University of California, Santa Cruz, CA, USA
4Fisheries Ecology Division, Southwest Fisheries Science Center, National Marine Fisheries Service, Santa Cruz, CA, USA
5Department of Biomolecular Engineering, University of California, Santa Cruz, CA, USA
6Department of Human Genetics, David Geffen School of Medicine, University of California, Los Angeles, CA, USA

Correspondence
Ying Zhen, Center for Tropical Research, Institute of Environment and Sustainability, University of California, Los Angeles, CA, USA.
Email: zhen@ucla.edu

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Abstract
The little greenbul, a common rainforest passerine from sub-Saharan Africa, has been the subject of long-term evolutionary studies to understand the mechanisms leading to rainforest speciation. Previous research found morphological and behavioural divergence across rainforest-savannah transition zones (ecotones), and a pattern of divergence with gene flow suggesting divergent natural selection has contributed to adaptive divergence and ecotones could be important areas for rainforests speciation. Recent advances in genomics and environmental modelling make it possible to examine patterns of genetic divergence in a more comprehensive fashion. To assess the extent to which natural selection may drive patterns of differentiation, here we investigate patterns of genomic differentiation among populations across environmental gradients and regions. We find compelling evidence that individuals form discrete genetic clusters corresponding to distinctive environmental characteristics and habitat types. Pairwise $F_{ST}$ between populations in different habitats is significantly higher than within habitats, and this differentiation is greater than what is expected from geographic distance alone. Moreover, we identified 140 SNPs that showed extreme differentiation among populations through a genome-wide selection scan. These outliers were significantly enriched in exonic and coding regions, suggesting their functional importance. Environmental association analysis of SNP variation indicates that several environmental variables, including temperature and elevation, play important roles in driving the pattern of genomic diversification. Results lend important new genomic evidence for environmental gradients being important in population differentiation.

KEYWORDS
adaptation, ecotone, environmental gradient, genomic divergence, RADseq, rainforest

1 INTRODUCTION

Rainforests are heralded for their exceptionally high biological diversity, yet the evolutionary mechanisms for the generation and maintenance of this diversity have been debated for decades (Beheregaray, Cooke, Chao, & Landguth, 2015; Haffer, 1969; Hoorn et al., 2010; Martin, 1991; Mayr & O’Hara, 1986; Moritz, Patton, Schneider, & Smith, 2000; Ogden & Thorpe, 2002; Price, 2008; Ribas, Aleixo, Nogueira, Miyaki, & Cracraft, 2011; Schluter, 2009; Schneider, Smith, Larison, & Moritz, 1999; Smith, Wayne, Girman, & Bruford, 1997; Smith et al., 2014). Models of rainforest speciation abound. Some emphasize the importance of neutral processes, for example genetic drift in allopatric populations isolated by historical refugia (Haffer, 1969), and some favour processes such as landscape change (Hoorn et al., 2010; Ribas et al., 2011) or dispersal (Smith et al., 2014), while others point towards a dominant role of divergent natural selection...
across ecological gradients and ecotones (Beheragary et al., 2015; Ogden & Thorpe, 2002; Schluter, 2009; Schneider et al., 1999; Smith et al., 1997, 2005, 2011). Each process is expected to shape the genomes of natural populations in different ways, leaving a signal that provides insights into the evolutionary mechanisms that may have led to divergence. Such information is of importance not only to evolutionary geneticists interested in understanding the processes involved in speciation, but also to conservation decision-makers, who are interested in preserving biodiversity and prioritizing new regions for protection in the face of rapid anthropogenic and climate change.

In this study, we explore the roles that population-level processes play in shaping biodiversity in Central Africa by examining the genomic diversity in a common songbird, the little greenbul (Andropadus virens). The little greenbul provides a particularly useful taxon for this enquiry because it has a broad geographic distribution across sub-Saharan Africa where it occurs in ecologically diverse habitats and has been the subject of long-term studies of intraspecific diversity and speciation. In the case of A. virens, as well as some other rainforest taxa, the rainforest–savannah transition zones (ecotones) have been shown to drive phenotypic divergence and likely speciation (Freedman, Thomassen, Buermann, & Smith, 2010; Kirschel, Blumstein, & Smith, 2009; Mitchell, Locatelli, Sesink Clee, Thomassen, & Gonder, 2015; Nadis, 2016; Smith et al., 1997, 2005). Compared to the central rainforest, ecotone habitats differ dramatically in numerous ways. For example, ecotones have less tree cover, lower levels of precipitation and greater intra-annual variation in environmental variables. These ecological differences may lead to distinctive food resources, pathogens, acoustic environments and predation levels (SlabbeKoorn & Smith, 2002; Smith et al., 2005, 2013). Consequently, these differences in both abiotic and biotic environments are hypothesized to result in divergent selection in ecotone and rainforest populations, leading to locally adapted populations (Freedman et al., 2010; Kirschel et al., 2009, 2011; Sehgal et al., 2011; Smith et al., 1997, 2005). This hypothesis is supported by the fact that parapatric A. virens populations across rainforest–ecotone gradients have undergone significant divergence in morphological (i.e., body mass, wing, tail, tarsus and beak length) and vocal characteristics despite significant levels of gene flow (Kirschel et al., 2011; SlabbeKoorn & Smith, 2002; Smith et al., 1997, 2005, 2013). This pattern of divergence with gene flow and the role of ecotones in driving adaptive divergence is further supported by the fact that allopatric rainforest populations of A. virens that were geographically isolated in West and Central Africa for two million years had much lower levels of phenotypic divergence in these traits compared to the level of divergence observed across a narrow (often 100 km) rainforest–ecotone gradient (Smith et al., 2005). Together, results for A. virens and those from other species suggest that strong divergent natural selection across the rainforest–savannah ecotone transition contributes to adaptive phenotypic divergence despite high levels of ongoing gene flow (Smith, Schneider, & Holder, 2001; Smith et al., 1997, 2005). Evidence for divergence with gene flow in A. virens is also consistent with models of ecological speciation where natural selection caused by shifts in ecology or invasions of new habitats can result in divergence in fitness-related traits and might play a prominent role in speciation (Beheragary et al., 2015; Ogden & Thorpe, 2002; Rundle & Nosil, 2005; Schluter, 2009; Schneider et al., 1999). Opportunities for this kind of divergence are possible across the little greenbul range, as they occur across a wide diversity of habitats, including mountains and islands, which are also known as hot spots of diversification and speciation (Darwin, 1859; Myers, Mittermeier, Mittermeier, da Fonseca, & Kent, 2000; Orme et al., 2005). Previous research has found that, compared to A. virens populations in mainland rainforests, mountain and island populations also show significant divergence in morphological traits typically related to fitness in birds, including body mass, wing length, tail length, tarsus length and bill size (Smith et al., 2005). Moreover, both habitats have considerable gene flow with mainland rainforest populations in Lower Guinea (Smith et al., 2005), suggesting natural selection may play an important role in divergence of mountain and island populations in A. virens.

To date, the paucity of high-resolution genomic data for rainforest species such as A. virens hinders a full exploration of the evolutionary mechanisms that may be important for diversification. Previous genetic studies on A. virens population structure utilized a handful of mtDNA markers (Smith et al., 2001) and microsatellite loci (Smith et al., 2005). These limited resources were unable to differentiate ecotone and forest populations at the genetic level; therefore, debates still remain whether the observed phenotypic divergence might be the result of plasticity in traits in response to varying environmental conditions, or strictly genomic divergence between populations in ecotone and rainforest. Recent development of next-generation sequencing techniques (NGS), especially restriction site-associated DNA (RAD) sequencing, enables one to de novo assemble hundreds of thousands of RAD loci across the genome in hundreds of samples without a reference genome. This cost-effective method to produce genomewide population data provides unprecedented opportunities to assess the patterns of diversity with much greater resolution, to find potential population structure and to identify candidate loci under local selection in nonmodel species such as A. virens.

Here, we take a population genomic approach leveraging single nucleotide polymorphism (SNP) data generated from RAD sequencing to survey the genomewide diversity of A. virens across multiple ecological habitats in Cameroon and Equatorial Guinea, including rainforests, ecotones, mountains, as well as an island. Our specific objectives for this approach were to (i) estimate overall levels of genetic diversity in A. virens; (ii) determine population structure and differentiation across habitats; (iii) identify candidate loci that are potential targets of selection; (iv) understand the biological functions of these candidate loci using transcriptome data; and (v) characterize genetic turnover across environmental gradients.

## MATERIALS AND METHODS

### 2.1 Sampling, DNA extraction and RADseq library construction

For RAD sequencing, blood samples from adult Andropadus virens were collected in Central Africa and stored in Queens lysis buffer.
Overall, 217 samples were collected from 15 geographically distant sampling sites (Figure 1a), representing 15 populations. Sampling sites were classified into one of four habitat types by researchers in the field and had previously been confirmed using remote sensing data (Slabbeekoon & Smith, 2002; Smith et al., 2005, 2013). Low-quality samples were removed by filtering, resulting in a total of 182 samples included in the final analysis (see RADseq data bioinformatics processing below). This included seven rainforest populations (83 samples), five ecotone populations (59 samples), two mountain populations (18 samples) and one population from the island of Bioko (12 samples). Each population was represented by 5–22 samples, with a mean representation of 12 samples (Table S1) (Nazareno, Bemmels, Dick, & Lohmann, 2017; Willing, Dreyer, & van Oosterhout, 2012).

Restriction site-associated DNA library preparation followed the methods for traditional RAD as described in Ali et al. (2016) that were slightly modified from the original RAD protocol as described in Baird et al. (2008). In short, genomic DNA (50 ng) for each sample was digested with 2.4 units of SbfI-HF (New England Biolabs, NEB, R3642L) at 37°C for 1 h in a 12 μl reaction volume buffered with 1× NEBuffer 4 (NEB, B7004S). Samples were heated to 65°C for 20 min, and then, 2 μl indexed SbfI P1 RAD adapter (10 nM) was added to each sample. Ligation of inline barcoded P1 adaptors was performed by combining 2 μl of the ligation mix (1.28 μl water, 0.4 μl NEBuffer 4, 0.16 μl rATP [100 mM, Fermentas R0441] with 0.16 μl T4 DNA Ligase [NEB, M0202M]) and heating at 20°C for 1 h followed by incubation at 65°C for 20 min. Following the ligation, half the per sample volume or 5 μl of each of the 96 samples was pooled into a single tube and cleaned using 1× Agencourt AMPure XP beads (A63881; Beckman Coulter); the remainder of the sample was stored for use in an additional library preparation if needed. The pooled DNA was then resuspended in 100 μl low TE and sheared to an average fragment size of 500 base pairs using a Bioruptor NGS sonicator (Diagenode). Sheared DNA was then concentrated to 55.5 μl using Ampure XP beads and used as the template in the NEBNext Ultra DNA Library Prep Kit for Illumina (NEB E7370L; version 1.2). The NEBNext protocol for library prep was followed apart from the fact that we used custom P2 adaptors which were created by annealing an NEBNext Multiplex Oligo for Illumina (NEB, E7335L) to the oligo GATCGGAAGACCGACGTCTGAACTCC AGTCACIIIATCAGAAAGA (the * represents a phosphorothioate DNA base). In addition, instead of the USER* enzyme step, we used a universal P1 RAD primer (AATGATACGGCACACCGAGATCTAC ACCTTTTTCCCTACAGACG*G) and a universal P2 RAD primer (CAAGCAGAAGACCGACGTCACTG A) during final amplification. The final RAD library was cleaned using AMPure XP beads and sequenced at the UC Berkeley QB3 Vincent J Coates Genome Sequencing Laboratory (GSL) on an Illumina HiSeq2000 (Illumina, San Diego, CA, USA) using single-end 100-bp sequence reads.

### 2.2 RADseq data bioinformatics processing

We used FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) to assess overall data quality of each RADseq sequencing run. To remove the lowest quality bases, we trimmed all raw sequencing reads (100 bp) by 14 bp at the 3′ end using seqtk (https://github.com/lh3/seqtk). We processed RADseq reads using the Stacks pipeline version 1.32 (Catchen, Amores, Hohenlohe, Cresko, & Postlethwait, 2011; Catchen, Hohenlohe, Basham, Amores, & Cresko, 2013) in the following manner. First, we demultiplexed the trimmed data by P1 barcodes and removed low-quality reads and those containing adapter sequences using process_radtags. After demultiplexing, reads were 80 bp in length (without barcodes) and data from different runs were combined together. These reads were used to de novo assemble RAD loci using denovo_map.pl (parameter settings: \( m = 3 M = 4 n = 4 \)). The parameters for de novo assembly were determined empirically to limit the impact of oversplitting of loci following methods described in Ilut et al. (2014) and Harvey et al. (2015). Specifically, we chose one sample that had a depth of coverage close to the median depth coverage of all samples and ran the de novo assembly over a wide range of values of \( M \) (1–8; \( n = M \) ) using ustacks. The percentage of homozygous and heterozygous loci

![FIGURE 1](image-url)  
Sampling and population structure. (a), Sampling locations. Each point is a sampling location, and habitat types are indicated by the same colour indicated in (b), (b–c), PCA using SNPs that have a minor allele frequency higher than 2%. Each point presents a sample, and samples are coloured by habitat type (b) or by population (c).
plateaued at $M = 4$, suggesting this value appropriately minimized oversplitting of alleles (Fig. S1). Thus, we used $M = 4$ for the final run on all samples. Stacks implements a multinomial-based likelihood model for SNP calling, by estimating the likelihood of two most frequently observed genotypes at each site and performing a standard likelihood ratio test using a chi-square distribution (Catchen et al., 2011; Hohenlohe et al., 2010). We used the default alpha (chi-square significance level) of 0.05. Paralogous loci that were stacked together were identified and removed by later quality control steps built into Stacks (e.g., max number of stacks per loci = 3; Ilut et al., 2014; Harvey et al., 2015). After the first round of assembly using denovo_map.pl, we ran stacks’ correction mode (rxstacks-cstacks-stacks) using the bounded SNP model with a 0.05 upper bound for the error rate (bound high = 0.05). The rxstacks program made corrections to genotype and haplotype calls based on population information, rebuilt the catalog loci and filtered out loci with average log likelihood $< -8.0$ (http://catchenlab.life.illinois.edu/stacks/).

We then identified a set of high-quality RAD loci for downstream population genetic analysis using the following steps. (i) We only kept RAD loci if they were present in at least 80% of all samples, because loci that only assembled in small subset of samples had limited utility in downstream analyses as well as possibly higher error rates. (ii) We filtered out RAD loci that had more than 40 SNPs along the 80-bp RAD loci sequence, as these likely represented sequencing errors or overclustering of paralogous loci. In the final data set, a maximum of 25 SNPs were recovered from a single RAD locus. Because the alignments look reasonable for the RAD loci that have higher number of SNPs, we did not apply any additional filters to avoid introducing additional biases. (iii) We mapped the RAD loci sequences from *A. virens* to the closest reference genome available, the zebra finch genome (version 3.24), using BLAT, and removed RAD loci that mapped to multiple positions in the zebra finch genome. (iv) We used BLAT to compare RAD loci sequences against each other and removed ones that had a match. This step further removes oversplitting RAD loci.

Following these filters, we obtained our final consensus set of RAD loci (Table S2). Samples that were missing more than 20% of the final consensus RAD loci were identified in a preliminary run and were removed from final analysis because they likely had low-quality DNA, low-quality libraries or low sequencing coverage. A total of 182 samples were included in the final data set (see above). Genotypes were called and filtered using methods implemented in the Stacks pipeline (Hohenlohe et al., 2010). We exported genotypes for the final consensus RAD loci in VCF format using Stacks populations program (only biallelic SNPs). Additional filters based on genotype calls were performed in vcftools (https://vcftools.github.io/index.html) or using custom scripts, which includes removing SNPs from the last seven bp of the RAD loci as this part of the locus was enriched for erroneous SNPs due to the lower sequencing quality at the 3’ end of reads, and filtering sites that have genotyping rate <80% of all samples.

We used the resulting full SNP data set with SNPs from all frequencies to estimate genetic diversity statistics such as number of segregating sites ($S$), average pairwise differences ($\theta$) and Waterson’s $\theta$ ($\theta_w$) in each population (Table S1). Rare SNPs that had a minor allele frequency (MAF) < 2% in the whole sample set were subsequently removed using vcftools, and the remaining SNPs were used for downstream analyses such as PCA, pairwise $F_{ST}$ calculations, BAYESCAN outlier analysis, and gradientForest analysis.

### 2.3 | RNA extraction, RNAseq library preparation and transcriptome de novo assembly

*Andropodus virens* lacks a reference genome. To help determine which of the RAD loci are transcribed, we collected RNAseq data and made a de novo assembly of the *A. virens* transcriptome. Fresh tissue samples were collected from 10 live individuals in the field (five tissue types: blood, brain, breast tissue, heart and liver). Tissue samples were stored in either PAXgene (Blood RNA Tubes; PreAnalytix/Qiagen, Switzerland) or Allprotect (Tissue Reagent, Qiagen, Germany) buffer and shipped to laboratory facilities at UCLA. RNA was extracted from each sample and tissue type separately using an RNeasy kit (Qiagen, Germany), and based on quality of extractions (both overall concentration and 260/280 ratio), three RNA samples from three tissue types (brain, heart and liver) were chosen to perform library preparations. RNAseq libraries were prepared using Illumina TruSeq RNA Library Prep Kit v.2 (Illumina, San Diego, California) following the manufacturer’s protocol, and libraries were indexed, normalized, pooled and sequenced on a single lane on Illumina HiSeq 2500 (paired-end 100-bp reads, Rapid Run mode) at GSL.

We obtained one lane of paired-end RNAseq data pooled from three tissue types. We first removed bases with quality scores lower than 20 and minimum sequence length of 30 bp using trim_galore (http://www.bioinformatics.babraham.ac.uk/projects/trim_galore/). We then pooled the remaining paired-end reads from different tissues together for a de novo assembly of the transcriptome using the TRINITY pipeline (Grabherr et al., 2011). We assessed the quality of the assembly using scripts provided in the TRINITY package and predicted the coding regions in the assembled transcriptome using TransDecoder in TRINITY.

### 2.4 | Detecting population structure using genomic data

To detect underlying population structure among samples, we performed a principal component analysis (PCA) using the bioconductor package SNPRELATE (Zheng et al., 2012). A total of 47,482 SNPs with MAF $\geq 2\%$ were used in PCA. The first six principal components were visually examined to identify clustering patterns of samples and to determine whether these genetic clusters tend to segregate with ecological factors or geography. We also used the program ADMIXTURE (Alexander, Novembre, & Lange, 2009) to estimate the ancestry of individual genotypes, using only the first SNP of each RAD loci to limit the impact of linkage disequilibrium. The analysis was run for $K = 1$–15.
To quantify pairwise population differentiation, we calculated pairwise $F_{ST}$ between populations using SNPPreloite. The correlation of population genetic differentiation (pairwise $F_{ST}$) and geographic distance, in other words, the presence of isolation by distance (IBD), was estimated by a simple Mantel test with 999,999 permutations using VEGAN 2.2-1 in R (Mantel, 1967; Oksanen et al., 2017). Mantel tests are reported using both raw $F_{ST}$ and $F_{ST}/(1–F_{ST})$, as well as both raw Euclidian geographic distance and log-transformed distances (Rousset, 1997; Slatkin, 1995).

Moreover, we found that pairwise $F_{ST}$ between populations from different habitats were higher than pairwise $F_{ST}$ computed between populations from the same habitat (see Results). In principle, this pattern could solely be driven by isolation by distance as populations from the same habitat tend to be located closer geographically to each other compared to population from different habitats. To determine whether the elevated $F_{ST}$ between populations from different habitats (compared to $F_{ST}$ between populations from the same habitat) could be explained simply by the differences in geographic distance, we performed permutation tests that accounted for the fact that populations from different habitats tend to be further apart. Specifically, we divided the population pairs into five bins based upon their geographic distances from each other (i.e., <200 km, 200–400 km, 400–600 km, 600–800 km, >800 km). Then, within each bin, we permuted whether the pairwise $F_{ST}$ values are from a within-habitat comparison or a between-habitat comparison. We generated 10,000 such permutations and, for each permutation, performed a $t$ test on whether the $F_{ST}$ values for between-habitat comparisons were higher than those for within-habitat comparisons. From the permuted data, we built a null distribution of $t$-statistics, which accounted for the effect of geography. Our final empirical $p$-value for the observed data was calculated as the percentage of permuted data sets that had a $t$-statistic as large or larger than the one seen in the original data. Similar permutation analyses were applied to the data set including all habitats as well as to a data set that only considered rainforest and ecotone populations. In the null distribution of $t$-statistics for the test of whether $F_{ST}$ is higher between as compared to within habitats, we found that none of the 10,000 permuted data sets had a $t$-statistic of $F_{ST}$ as large or larger than the one seen in the original data, suggesting a $p$-value < 1e-04. However, for the null distribution of $t$-statistics for the test of whether distance is higher between or within habitats, 1,581 of the 10,000 permuted data sets had a $t$-statistic as large or larger than the one seen in the original data, suggesting a $p$-value = .158. This suggests that our null distribution of $t$-statistics accounts for the fact that populations from similar habitats tend to be closer together geographically.

As an alternative method to test whether habitat contributed to the observed pattern of population differentiation above and beyond geographic distance, we created a binary matrix that indicated whether a pair of populations was from the same habitat or not. We tested the correlation of genetic distance matrix and this matrix while controlling for geographic distance using a partial Mantel test using VEGAN 2.2-1 in R (Mantel, 1967; Oksanen et al., 2017). Partial Mantel tests were performed using both raw $F_{ST}$ and $F_{ST}/(1–F_{ST})$, as well as both raw Euclidian geographic distance and log-transformed distances (Rousset, 1997; Slatkin, 1995).

### 2.5 Identifying outlier SNPs under selection

We used BAYESCAN 2.1 (Foll & Gaggiotti, 2008) to identify highly differentiated SNPs that are candidates to be under natural selection. This program takes a Bayesian approach to search for SNPs exhibiting extreme $F_{ST}$ values that could be due to local adaptation. Outlier SNPs were identified using SNPs with MAF ≥ 2% across all samples, specifying all 15 populations or four habitats (see Data S1). We ran 5,000 iterations using prior odds of 10 and assessed the statistical significance of a locus being an outlier using a false discovery rate (FDR) of 5%.

To explore the spatial patterns of population differentiation across chromosomes, we mapped the consensus RAD loci to the zebra finch genome using BLAT with default parameters. For the uniquely mapped RAD loci, we plotted the $F_{ST}$ of each SNP by genome coordinates to examine spatial patterns of outlier SNPs. To interpret the potential biological function of the outlier SNPs identified by Bayescan analysis, we used a zebra finch genome annotation v3.2.4.78 to identify outlier SNPs mapped to annotated genomic regions.

We further examined whether candidate loci under selection were enriched in exonic (transcribed) or coding regions. To do this, we mapped RAD loci to the de novo assembled A. virens transcriptome using BLAT with default parameters. Any RAD locus that mapped to the transcriptome was considered to be in exonic regions of the genome, and the remaining RAD loci were labelled “nontranscribed” regions of the genome. Similarly, we mapped RAD loci to predicted coding sequences and categorized them into coding and noncoding RAD loci. We then used a one-sided Fisher’s exact test to examine whether there was significant enrichment of outlier loci in exon or coding regions of the genome. Finally, we cross-checked these outliers to see whether there were any significant associations with environment using latent factor mixed models (Frichot, Schoville, Bouchard, & François, 2013) (see Data S1 for more details).

### 2.6 Detecting environmental drivers of genomic variation

In addition to population structure, we also tested whether allele frequencies in different populations were associated with environmental variables across the range of A. virens using the package GRADIENTFOREST (Ellis, Smith, & Pitcher, 2012) in the R statistical framework (R Core Team, 2014). Gradient forests are an extension of random forests (Breiman, 2001) that treat response variables (in this case, individual SNP minor allele frequencies within each population) as members of a larger community (the total genome) and provide summary statistics based on ensembled forest runs to indicate an overall association of changes in allele frequency to particular
environmental variables (Ellis et al., 2012; Fitzpatrick & Keller, 2015). Gradient forests were run using the following changes to the default settings: number of trees run for each environmental variable = 500, number of SNPs included in each bin = 1,000. Allelic frequencies across the genome were predicted for unsampled geographic locations by generating a random set of 100,000 points across the range of A. virens. Then, we used our final gradient forest model to predict allele frequencies at each of those points, given their environmental characteristics. Ordinary Kriging (Oliver & Webster, 1990) was then used within ArcMap (ESRI, Redlands, CA) to generate a continuous surface across the known range of A. virens in Cameroon.

We used a suite of 17 environment variables (Table S6), including bioclim measures of temperature and precipitation (n = 9; any variables showing a Pearson’s correlation coefficient >0.7 were removed) downloaded from the Worldclim database (www.worldclim.org), measures of vegetation and tree cover captured using the NASA MODerate-resolution Imaging Spectroradiometer (MODIS, n = 4), elevation and slope captured via the Shuttle Radar Topography Mission (n = 2), and surface moisture estimates measures using the Quick Scatterometer (QuikSCAT, n = 2). In addition to these variables, and to account for purely geographic associations, we also included Euclidean distances (measured as latitude and longitude) as predictor variables in all models.

3 | RESULTS

3.1 | SNP discovery and overall genetic diversity

We used RAD sequencing to survey the genomewide diversity of Andropadus virens. The final sample set included 15 populations from four different habitats, including rainforests, ecotones, mountains and an island (Figure 1a; Table S1). After removing low-quality reads and samples, we obtained a total of 916 million reads for 182 A. virens samples (PRJNA390972). The number of raw sequence reads per sample ranged from 1.60 to 20.73 million. On average, 99.2% of these reads were utilized in the de novo assembly of the RAD loci. The mean coverage depth ranged from 16× to 136× per sample (mean = 38×, median = 32×, Fig. S2). Using this data set, we assembled and identified 34,657 high-quality RAD loci that passed our quality filters and were genotyped in more than 80% of all final samples. On these 34,657 consensus RAD loci, there were a total of 255,290 SNPs. The median number of SNPs per RAD locus is seven. With a minimum minor allele frequency filter of 2%, we retained 47,482 SNPs that were present on 23,882 RAD tags (Table S2; Data S1).

The number of segregating sites ranges from 25,936 to 70,598 per population. Waterson’s θ (θw) was estimated to be 0.0049–0.0076/bp (mean = 0.0064/bp) and π ranges from 0.0034 to 0.0037/bp (mean = 0.0036/bp) (Table S1), which is comparable to π estimated from other bird species (Nadachowska-Brzyska et al., 2013; Romiguier et al., 2014). Overall levels of genetic diversity are comparable in each habitat, including the island population (Table S1). The finding that θw is larger than π indicates an excess of low-frequency variants relative to the standard neutral model which could be driven by recent population expansions.

3.2 | Transcriptome assembly and annotation

Transcriptome assembly was performed using 169 million paired-end RNAseq data from three different tissue types (PRJNA390772). The assembled transcriptome had a GC content of 45%. The average contig length was 815 bp, and N50 was 1,619 bp. In total, trinity produced 237,226 genes and 286,494 transcripts and predicted 81,018 coding sequences from these transcripts (Table S5). Of the 34,657 RAD loci we genotyped, 8412 RAD loci (24.2%) were mapped to the de novo assembled A. virens transcriptome, and 3,618 RAD loci (10.4%) were mapped to the predicted coding sequences (Fig. S3). The RAD tags overlapping coding sequence tend to have fewer SNPs than those that do not overlap with coding sequences (Fig. S4), consistent with the fact that the coding regions are likely under stronger selective constraint.

3.3 | Population structure

We used PCA to identify population structure in little greenbuls. The first two PCs revealed a clear clustering pattern of individuals from the same habitats (Figure 1b). Populations from the island, mountains, rainforests and ecotones formed four discrete clusters, suggesting genomic divergence across ecological gradients and habitats. Island and mountain populations were most distinct (Figure 1b); however, samples from all four habitats separated on PC1, including those from ecotone and rainforest habitats. PC2 further separated island and mountain samples from rainforest and ecotone samples. Remarkably, results suggest that, within rainforest and ecotone habitats, individual populations could be distinguished solely on the basis of genomic markers, mostly by PC1, with individuals from the same sampling sites clustering together (Figure 1c). The level of separation of ecotone populations from rainforest populations along PC1 approximately followed a latitudinal gradient, corresponding to environmental and rainfall gradients that distinguish ecotone in the north from rainforest in the south of Cameroon (see environmental analyses below). Specifically, samples collected from sites Wakwa and Ngaoundaba Ranch, towards the more extreme edge of the ecotone habitat and having the most extreme ecotone environmental conditions, formed clusters that were most distant from the rainforest samples, while samples collected from Betare Oya, at lower ecotone that was closest to the central rainforests, formed a cluster closest to the rainforest (Figure 1c). The pattern of genomic differentiation across habitats was confirmed using the program ADMIXTURE (Fig. S5).

Pairwise $F_{ST}$ between populations ranged from 0.017 to 0.078 (mean = 0.038; Figure 2a–b; Table S3), indicating low overall levels of genomic differentiation across populations. There was significant correlation between pairwise $F_{ST}$ and geographic distance between the populations (Mantel $r = .34$; mantel simulated p-value = .003), suggesting isolation by distance contributes to population differentiation. However, pairwise $F_{ST}$ between populations from different...
habitats was significantly higher than pairwise $F_{ST}$ between populations within the same habitat (one tailed $t$ test, $p$-value = $1.36e-10$; Figure 2a). Pairwise geographic distances between populations from different habitats were also significantly higher than pairwise distances between populations within the same habitat (one tailed $t$ test, $p$-value = $1.015e-06$). To account for the fact that populations from different habitats were also geographically further apart, we performed a permutation test, where we randomized whether a population pair was from the same or different habitats in different bins stratified by their geographic distance. Using permuted data sets, we built a null distribution of these t-statistics (that already includes the effect of geographic distance), which we used to evaluate the significance of our observed value. The higher $F_{ST}$ value for between-habitat comparison was highly significant when compared to this improved null distribution ($p$-value < $1e-04$, Figure 2c and Fig. S6), suggesting that isolation by distance alone cannot explain the higher $F_{ST}$ between habitats than within habitats. Similarly, only considering rainforest and ecotone populations, pairwise $F_{ST}$ was significantly higher between habitats as compared to within habitats (one tailed $t$ test, $p$-value = $9.793e-05$). Application of the same permutation test shows the higher $F_{ST}$ between ecotone and rainforest populations.
mountain populations were exceptionally high (from one another and from lowland populations. Divergence was geographically very close to each other. The FST between mountain populations and lowland forest/ectone populations was larger than pairwise FST values between mountain populations and other populations (Figure 2b). Interestingly, FST between two mountain populations were exceptionally high (FST = 0.060) compared to other within-habitat pairwise FST (ranging from 0.017 to 0.040, Figure 2a), despite the fact that the two mountain populations were geographically very close to each other. The FST values between mountain populations and lowland forest/ectone populations were larger than pairwise FST values between lowland populations, suggesting mountain populations are highly differentiated both from one another and from lowland populations.

3.4 Candidate loci under selection

To further explore potential candidate loci under selection, we identified SNPs with extreme allele frequency differences across populations, which should be enriched by targets of local adaptation. We identified 140 outlier SNPs across all populations with a false discovery rate of 5% using BAYESCAN. These candidate SNPs are potential targets of divergent selection across different sampling sites (Fig. S7). The 140 outlier SNPs reside in 119 loci, and 40 of these loci mapped to the zebra finch genome (Fig. S9). Of these, 36 mapped to main scaffolds of known chromosomes and four mapped to the Z chromosome. Only 13 of these outlier loci mapped to annotated genomic regions on the zebra finch genome and nine mapped to genes with functional annotations (Table S4).

To uncover the functional significance of outlier loci, we used the de novo assembly of greenbul transcriptome to partition the RAD loci and SNPs into different categories depending on whether they mapped to coding regions or transcribed (exonic) regions (Fig. S10). This enabled us to test for enrichment of outlier SNPs in putatively functional regions. Of the 47,482 SNPs, 9677 mapped to the transcriptome, and 42 were outliers based on a Bayescan analysis. Using a one-sided Fisher's exact test, we detected significant enrichment of outlier loci in exonic regions of the genome (p = 0.0044; Table S2; Fig. S10). Using the predicted coding sequence from the transcriptome, 3,602 SNPs mapped to the predicted coding sequences and 21 of these were outliers. We again detected a significant enrichment of outlier SNPs in protein-coding sequences (p = 0.002; Table S2; Fig. S10). Taken together, these enrichment results provide additional confidence that the outlier loci found using Bayescan captured functionally important, biologically relevant genetic variants, which were not merely loci that fell within the tail of a neutral distribution.

3.5 Genomic turnover across environments

Because some environmental adaptation may involve shifts in allele frequency at many loci across the genome (e.g., polygenic selection involving many genes of small effect), we used the gradientForest approach to look for correlations in allele frequencies associated with environmental variables. A total of 7238 SNPs, ~15% of all SNPs, had R² values above 0 (0.0073–0.83) when testing for a correlation between frequency and an environmental variable. Of the 19 environmental and geographic variables included in models (Table S6), variables capturing temperature variation (Min Temp: minimum temperature of the coldest month, Temp Range: mean diurnal temperature range and Mean temp: mean annual temperature) and elevation were most important in explaining environmentally associated variation in SNPs (Figure 3a). In some cases, measures of surface moisture or tree cover were also important, but axes for these variables largely overlapped with temperature or elevation measures along PC plot (likely the result of colinearity in environmental variables) (Figure 3b). Results from LFMM analyses indicated these same variables were also associated with differentiation observed at hundreds of individual loci, although exact functions of these regions remain unknown (see Data S1).

Geographic variables alone were not as important in explaining variation in allele frequency, again suggesting that geographic distance cannot fully account for all variation in SNP frequencies across the range of little greenbuls. Above and beyond neutral processes, selective pressures imposed by differences in these environments best explains the observed genomic patterns of variation. Predictions across Cameroon suggest strong genomic turnover (defined as coordinated shifts in allele frequencies across the genome) throughout the forest, savannah and ectone regions, with diagnostic genomic
variation occurring in each of these habitats (Figure 3). Distinct SNP frequencies at high elevations (Figure 3b–c) and the fact that elevation explains a large proportion of variation of allele frequencies in the greenbul genome (largely allied with PC1, Figure 3b) also suggest a unique genetic signature in populations at elevation.

4 | DISCUSSION

In this study, we used genomewide RADseq SNPs to characterize the overall level of genetic diversity in Andropadus virens populations across four different habitats. We found evidence of population structure of A. virens consistent with habitat type and previously observed phenotypic divergence. We demonstrated that population differentiation across habitats cannot be explained solely by isolation by distance, suggesting local adaptation further contributes to genomic divergence among habitats. We identified 140 outlier SNPs that are potential targets of selection and the fact that they are significantly enriched in exonic and coding regions suggests they are functionally important. Environmental association analysis further supports this conclusion and shows environmental variables, including temperature and elevation, are highly associated with patterns of genomic variation across the range of the little greenbul.

In addition to the differences between rainforest and ecotone populations, other habitats were found to harbour distinct patterns of genetic variation. The population from Bioko Island formed a distinctive genetic cluster based on PCA and also was identified as distinct in environmental association models (Figures 1b and 3), consistent with previous studies (Smith et al., 2005). Bioko Island is 32 km off the coast of Africa, separated from mainland ~10,000 years ago and has an area of 2,017 km². Island populations and species may have smaller effective population sizes than mainland populations or sister taxa (Robinson et al., 2016), due to possible population bottlenecks and considerably smaller ranges. As a result, island populations may have lower genetic diversity compared to their mainland counterparts (Frankham, 1997). However, in our study, the estimates of genetic variation using genomewide SNP markers in the greenbul population on Bioko Island are comparable to those from mainland populations (Table S1). This is consistent with the recent findings that island populations do not always have lower genetic diversity, particularly in birds (Francisco, Santiago, Mizusawa, Oldroyd, & Arias, 2016; James, Lanfear, & Eyre-Walker, 2016), and the fact that Bioko island is a large island that only recently separated from the mainland.

Tropical mountains are well known to support disproportionately high biodiversity and are thought to be hotspots for avian speciation (Drovetski et al., 2013; Fjeldså, Bowie, & Rahbek, 2012; Fjeldså, Johansson, Lokugalappatti, & Bowie, 2007; Myers et al., 2000; Orme et al., 2005; Roy, 1997; Smith et al., 2000). Little greenbuls are found at elevations up to 2,400 m, where environmental variables, particularly temperature and vegetation, change rapidly along altitudinal gradients. Our two mountain populations have high F_{ST} despite being geographically close and from same habitat type. Although the Euclidean distance between these two mountain populations is short, the environmental changes along altitudinal gradients are steep, causing isolation between populations from different mountains and forming "sky islands", between which the level of gene flow probably is much lower than among lowland populations. Moreover, we found that the two different mountain populations exhibited the lowest within-population genetic variation among all sampled populations (Table S1). While this difference was not statistically significant (likely due to small sample sizes), this decreased level of variation can inflate $F_{ST}$, the relative measurement of population differentiation. It also suggests that mountain populations may have overall smaller effective population sizes (consistent with presumably smaller suitable habitat size for mountain populations) and/
or have experienced serial bottleneck/founder effects as range expansions occurred. These processes can further contribute to divergence due to stronger genetic drift within each subpopulation leading to faster changes in allele frequencies. The idea that elevation can drive genomic changes is supported by previous estimates of morphological divergence (Smith et al., 2005) and emphasizes the importance of preserving elevational gradients in tropical ecosystems in general (Thomassen et al., 2011).

Most of the genes containing outlier SNPs only have annotations predicted from human homologs, except two that have annotations from bird species. Both of these two genes are of particular interest. One outlier locus mapped to the SUTR/coding junction of EDIL3, a calcium-binding protein that has been found to function in avian eggshell biomineralization (Marie et al., 2015). Avian eggshells protect the developing embryo and keep the egg free from pathogens. Environmental factors such as temperature, humidity and partial oxygen pressure have been reported to affect avian eggshell structure, and previous studies documented rapid evolution of eggshell structure in response to colonization of novel environments in the house finch (Stein & Badyaev, 2011). The second outlier locus mapped to MLXIPl (MLX interacting protein-like), which is a coactivator of the carbohydrate response element binding protein that has been correlated with fat deposition in caged chickens (Li et al., 2015; Proszekowiec-Weglarz, Humphrey, & Richards, 2008). Interestingly, seven more genes that contain outlier SNPs have annotations linked with metabolic traits or diseases in humans. For example, outlier SNPs were found in RGS6 (Sibbel et al., 2011 p. 6), CSAD (Comuzzie et al., 2012) and the UNC13B intron (Trégouet et al., 2008), which were associated with dietary fat intake, food preference, adiposity/obesity and diabetes in humans. Although metabolic traits were not measured, adult little greenbuls from the rainforest have significantly smaller body mass and body size compared to ecotone, mountain and island populations (Smith et al., 1997, 2005), which could be the result of divergent selection of these genes associated with metabolic traits. Several recent studies have discussed the limitations of identifying FST outlier as loci under divergent selection and suggest results should be interpreted carefully, because many other factors, including demographic history, recombination rate heterogeneity and background selection, may also create FST outliers (Cruickshank & Hahn, 2014; Lotterhos & Whitlock, 2014; Roesti, Salzburger, & Berner, 2012). Current work to model the demographic history of A. virens should help examine these various possibilities in greater detail.

Numerous hypotheses have been proposed for how biodiversity is generated in rainforests (Matur & O’Hara, 1986; Moritz et al., 2000). With the rapid advances in genomics and environmental modelling in the last decade, it is now possible to examine these mechanisms in greater depth. Using more powerful genomewide data, we have shown, for the first time, strong patterns of population structure and genomic differentiation between rainforest and ecotone habitats in A. virens. Previously, no genetic differentiation was found between morphologically divergent populations in rainforest and ecotone habitats, leaving open the possibility that the observed morphological difference could simply be the result of a homogenized meta-population that differentially responds to environmental gradients. Although identifying the underlying genetic basis of morphological traits that differ between rainforest and ecotone populations was beyond the scope of this study, our results complement previous work by demonstrating that populations along the rainforest–ecotone gradient are diverging at the genomic level, and raise the possibility that local adaptation could account for the patterns of morphological variation previously observed across ecotone–rainforest gradients. Results also complement past research on reproductive behaviour, which found differences in song characteristics along the forest–ecotone gradient, and showed experimentally that singing males respond more aggressively to male songs from their own habitat, suggesting incipient reproductive isolation driven by habitat (Kirschel et al., 2011; Slabkekoorn & Smith, 2002; Smith et al., 2013). These patterns of differentiation are consistent with models of ecological speciation, where natural selection caused by shifts in ecology can promote speciation (Beheregaray et al., 2015; Hanson, Moore, Taylor, Barrett, & Hendry, 2016; Ogden & Thorpe, 2002; Orr & Smith, 1998; Price, 2008; Räsänen & Hendry, 2008; Rundle & Nosil, 2005; Schluter, 2000, 2009; Schneider et al., 1999). However, further research is necessary to more fully understand the evolutionary significance of divergence across ecological gradients and ecotones. In particular, studies investigating the underlying genetic basis of phenotypic differentiation and mate choice experiments would provide additional insights into their importance in divergence and speciation.

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DATA ACCESSIBILITY

RADseq data: NCBI SRA database BioProject ID PRJNA390986. RNAseq data: NCBI SRA database BioProject ID PRJNA390772. Data files including RAD loci consensus sequences, VCF file and sample information available at Dryad https://doi.org/10.5061/dryad.8n8t0.

AUTHOR CONTRIBUTIONS

T.B.S. and K.E.L conceived and supervised the project. S.L. and R.J.H conducted the laboratory work. Sequence assemblies, population structure and outlier analysis were primarily carried out by Y.Z. with assistance from T.N., K.R., E.C.A. and K.E.L. Environmental association analysis was performed by R.J.H. The manuscript was written by Y.Z., R.J.H., K.R., K.E.L. and T.B.S., with input from all authors.
REFERENCES


Nadis, S. (2016). Life on the edge: Saving the world’s hotbeds of evolution/New Scientist.


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