Target Capture and Massively Parallel Sequencing of Ultraconserved Elements for Comparative Studies at Shallow Evolutionary Time Scales

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Received 7 January 2013; reviews returned 1 April 2013; accepted 30 August 2013

Abstract—Comparative genetic studies of non-model organisms are transforming rapidly due to major advances in sequencing technology. A limiting factor in these studies has been the identification and screening of orthologous loci across an evolutionarily distant set of taxa. Here, we evaluate the efficacy of genomic markers targeting ultraconserved DNA elements (UCEs) for analyses at shallow evolutionary timescales. Using sequence capture and massively parallel sequencing to generate UCE data for five co-distributed Neotropical bird species, we recovered 776–1,516 UCE loci across the five species. Across species, 53–77% of the loci were polymorphic, containing between 2.0 and 3.2 variable sites per polymorphic locus, on average. We performed species tree construction, coalescent modeling, and species delimitation, and found that the five co-distributed species exhibited discordant phylogeographic histories. We also found that species trees and divergence times estimated from UCEs were similar to the parameters obtained from mtDNA. The species that inhabit the understory had older divergence times across barriers, contained a higher number of cryptic species, and exhibited larger effective population sizes relative to the species inhabiting the canopy. Because orthologous UCEs can be obtained from a wide array of taxa, are polymorphic at shallow evolutionary timescales, and can be generated rapidly at low cost, they are an effective genetic marker for studies investigating evolutionary patterns and processes at shallow timescales.

Highly multilocus data sets are becoming more widespread in phylogeography and population genetics. Population genetic theory and simulation studies suggest that increases in the number of unlinked loci included in analyses improve estimates of population genetic parameters (Kuhner et al. 1998; Beerli and Felsenstein 1999; Carling and Brumfield 2007). Under a Wright–Fisher model of evolution, variance in the topology and branch lengths among gene trees is large (Hudson 1990). Processes such as gene flow and natural selection add to this variance. To increase the precision of parameters estimated under the coalescent process, comparative genetic studies have transitioned from single-locus mitochondrial to multilocus data sets (Brumfield et al. 2003). Multilocus data permit the application of parameter-rich coalescent models that can simultaneously estimate multiple demographic parameters (Edwards and Beerli 2000), yet previous methods for collecting multilocus data from non-model taxa were limited.

The advent of massively parallel sequencing (MPS) and the development of generalized data collection and analytical methods allow biologists to scale data collection to thousands of unlinked loci, even without prior knowledge of the underlying genome sequence (McCormack et al. 2012b). Various approaches use reduced representation library preparation methods, such as ddRADs (Peterson et al. 2012), are relatively easy and cheap to perform, and analyses of SNP data sets derived from these methods are increasing (e.g., Emerson et al. 2010; Compert et al. 2010; McCormack et al. 2012c; Zellmer et al. 2012). However, as the genetic distance among study taxa increases, so does the challenge of collecting data from orthologous loci across species using PCR- or restriction digest-based approaches (Rubin et al. 2012).

The importance of using orthologous loci for comparative phylogeographic studies is unclear. Theory and simulations suggest that population genetic inferences from a non-intersecting set of loci will converge on the same answer, provided the data matrices comprise a sufficient number of independently segregating loci (Kuhner et al. 1998; Beerli and Felsenstein 1999; Carling and Brumfield 2007). It remains an open question, however, whether this is true in practice, especially because mutation rates are known to vary dramatically across the genome (Hodkinson and Eyre-Walker 2011). As a result, direct comparisons among divergent species are best accomplished by examining sets of orthologous loci.

A recently developed class of markers surrounding ultraconserved DNA elements (UCEs; Faircloth et al. 2012) can be used in conjunction with sequence

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capture and MPS to generate large amounts of orthologous sequence data among a taxonomically diverse set of species. UCEs are numerous in a diversity of metazoan taxa (Ryu et al. 2012), and over 5000 have been identified in amniotes (Stephen et al. 2008; Faircloth et al. 2012). UCEs have been used to resolve challenging phylogenetic relationships among basal avian (McCormack et al. 2013) and mammalian (McCormack et al. 2012a) groups, to determine the phylogenetic placement of turtles among reptiles (Crawford et al. 2012), and to understand the relationships among ray-finned fishes (Faircloth et al. 2013). Although UCEs are highly conserved across distantly related taxa, their flanking regions harbor variation that increases with distance from the conserved core (Faircloth et al. 2012). The conserved region allows easy alignment across widely divergent taxa, while variation in the flanks is useful for comparative analyses. Faircloth et al. (2012) suggested that because variation within human UCE flanking regions is abundant, UCEs would likely be useful for investigations at shallow evolutionary timescales. Here, we use empirical data from Neotropical birds to demonstrate that UCEs are useful genetic markers at shallow evolutionary timescales.

The lowland Neotropics harbor arguably the highest avian diversity on Earth and the origins of this diversity are contentious (Ehlo et al. 2010; Rull 2011; Brumfield 2012; Ribas et al. 2012; Smith et al. 2012a). The majority of phylogeographic studies on Neotropical birds have used mitochondrial DNA (mtDNA) to estimate divergence times and, in some cases, migration rates. Recent work suggests that ecology may play an important role in avian diversification in the lowland Neotropics (Burney and Brumfield 2009). Highly multilocus data sets in conjunction with highly parameterized models promise to provide further resolution to the processes underlying Neotropical diversity.

Here, we use UCE data to study genetic structure in five widespread Neotropical birds—Cymbilaimus lineatus, Xenops minutus, Schiffornis tundina, Querula purpurata, and Microcerulus marginatus. Cymbilaimus lineatus and Q. purpurata inhabit the canopy, whereas the other three species inhabit the under- and mid-story. Recent taxonomies treat the understory species as more polytypic than the canopy species. Dickinson (2003), for example, includes 3 subspecies within C. lineatus and treats Q. purpurata as monotypic, but lists 10, 5, and 6 subspecies for X. minutus, S. tundina, and M. marginatus, respectively. We use target enrichment followed by Illumina sequencing to collect population-level UCE data. In each species, we: (i) infer the species tree; (ii) use demographic models to estimate population genetic parameters; and (iii) identify cryptic species-level diversity within each of the five lineages. We compare and contrast these results with mtDNA (cytochrome b) data from the same populations. We examine whether canopy species and understory species display different divergence histories, effective population sizes, and migration rates.

## Methods

### Sequence Capture of UCEs from Genetic Samples

We sampled 40 individuals from five species of widespread Neotropical lowland forest passerines—Cymbilaimus lineatus, Xenops minutus, Schiffornis tundina, Querula purpurata, and Microcerulus marginatus (Supplementary Table S1; Supplementary Fig. S1). Understory birds have been shown to exhibit greater genetic differentiation across biogeographic barriers than more vagile canopy birds (Burney and Brumfield 2009). Our taxon sampling allowed us to address how dispersal affects genetic structure by including two canopy (C. lineatus and Q. purpurata) and three understory (X. minutus, S. tundina, and M. marginatus) species. Because each of the five species is distributed across three prominent biogeographic barriers—the Andes Mountains, the Isthmus of Panama, and the upper Amazon River—we were able to examine how these landscape features affected the geographic structuring of populations. For each species, we sampled two individuals from each of four areas of endemism that, collectively, span these barriers: Central America (CA), Chocó (CF), Napo (NA), and Inambari/Rondônia (SA) (Supplementary Table S1; Fig. 1).

We extracted total genomic DNA from vouched tissue samples using a DNeasy tissue kit (Qiagen). We fragmented genomic DNA using a BioRuptor NGS (Diagenode) and used Illumina libraries using KAPA library preparation kits (Kapa Biosystems) and custom sequence tags unique to each sample (Faircloth and Glenn 2012). To enrich targeted UCE loci, we followed an established workflow (Gnirke et al. 2009; Blumenstiel et al. 2010) incorporating several modifications to the protocol detailed in Faircloth et al. (2012). First, we prepared and sequence-tagged libraries using standard library preparation techniques. Second, we pooled eight samples within each species at equimolar ratios, prior to enrichment, and we blocked the Illumina TruSeq adapter sequence using custom blocking oligos (Supplementary Table S2). We enriched each pool using a set of 2560 custom-designed probes (MYcroarray, Inc.) targeting 2386 UCE loci (see Faircloth et al. 2012) and http://ultraconserved.org (last accessed September 24, 2013) for details on probe design). Prior to sequencing, we qPCR-quantified enriched pools, combined pools at equimolar ratios, and sequenced the combined libraries using a partial (90%) lane of a 100-bp paired-end Illumina HiSeq 2000 run (Cofactor Genomics).

last accessed September 24, 2013), and excluded reads containing ambiguous (N) bases. Following quality filtering, we generated consensus contigs, *de novo*, for all reads generated for each species using VelvetOptimiser (http://bioinformatics.net.au/software.velvetoptimiser.shtml last accessed September 24, 2013) and VELVET (Zerbino and Birney 2008). We aligned consensus contigs from VELVET to UCE probes using LASTZ (Harris 2007), and we removed any contigs that did not match probes or that matched multiple probes designed from different UCE loci. We used BWA (Li and Durbin 2009) to generate an index of consensus UCE contigs for each species and we mapped all reads to the index on an individual-by-individual basis. After mapping, we called SNPs and indels for each individual and exported individual-specific consensus sequences for each UCE locus, and we hard-masked low-quality bases (<Q20) within resulting consensus sequences using seqtk (https://github.com/lh3/seqtk last accessed September 24, 2013) and converted the masked FASTQ files to FASTA format.

We aligned hard-masked FASTA sequences within loci for each species using MAFFT (Katoh et al. 2005), and we used the alignments to generate additional input files for population genetic analyses in lociNGS (Hird 2012). We removed any alignments in a given species that did not contain data for at least one individual from each of the four areas of endemism. For each species, we generated two data matrices—one containing alignments of all loci recovered for the respective species (“single species full” data set) and another, smaller data set containing only the subset of loci that we also recovered in the other four species (“single species reduced” data set). We use these two data sets from each species in most subsequent analyses, we also generated interspecific alignments of individuals across all five species in MAFFT—one “all species full” data set containing all alignments, and one “all species reduced” data set containing only loci with data from at least one individual in each area of endemism in all five species. For clarity, we present details of these data sets in Table 1.

### Summary Statistics

We used alignments for each species to calculate a range of population genetic summary statistics (haplotype diversity, Watterson’s $\theta$, $\pi$, Tajima’s $D$) using the program COMPUTE (Thornton 2003). We calculated the number of variable positions, the distance of variable positions from the center of the UCE locus, and the frequency of variation by distance using the “get_smilogram_from_alignments.py” program within the PHYLUCE package (https://github.com/faircloth-lab/phyluce last accessed September 24, 2013). This program screens locus-specific UCE alignments for variability after excluding masked/ambiguous bases and unknown characters and stores locus- and position-specific variability data in a relational database (http://www.sqlite.org/ last accessed September 24, 2013). The program also tallies the count of variable and total positions by distance from the UCE center to compute a frequency of variable bases by distance and exports these summary data as a CSV file. We plotted these data in R using ggplot2 (Wikham 2009) after masking outlying positions with a frequency >0.04.
TABLE 1. UCE data sets used for computation of summary statistics and subsequent analyses

<table>
<thead>
<tr>
<th>Data set name</th>
<th>Number</th>
<th>Number of loci</th>
<th>Number of individuals in alignment</th>
<th>Description</th>
<th>Analyses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single species full</td>
<td>5</td>
<td>Variable</td>
<td>8 (4 for <em>M. marginatus</em>)</td>
<td>All loci within a given species that have data for at least one individual in each area of endemism</td>
<td>Summary statistics, G-PhoCS</td>
</tr>
<tr>
<td>Single species reduced</td>
<td>5</td>
<td>(one per species)</td>
<td>166 (4 for <em>M. marginatus</em>)</td>
<td>All loci within a given species that fulfill the criterion for the full data set for all five study species</td>
<td>Summary statistics, *BEAST, G-PhoCS, BPP</td>
</tr>
<tr>
<td>All species full</td>
<td>1</td>
<td>2219</td>
<td>36</td>
<td>All loci that are present in at least two individuals across species</td>
<td>Summary statistics</td>
</tr>
<tr>
<td>All species reduced</td>
<td>1</td>
<td>169</td>
<td>36</td>
<td>All loci that have data for at least one individual in each area of endemism for all species</td>
<td>Summary statistics</td>
</tr>
</tbody>
</table>

UCE Species Tree Construction

We used *BEAST (Heled and Drummond 2010)* in the BEAST package (Drummond et al. 2012) to generate species trees using the single species reduced data sets.

To assign samples to species/populations in *BEAST*, we used the endemic areas Central America (CA), Chocó (CH), Napo (NA), and Inambari/Rondônia (SA) as proxies for populations (Fig. 1). We split the two SA samples of *S. turdina* (Inambari = SA1 and Rondônia = SA2) for this and subsequent analyses because the SA2 sample was east of the Madeira River and in the Rondônia endemic area and showed high divergence from SA1 based on review of the alignments (Supplementary Fig. S1). We estimated the best-fit model of sequence evolution using CloudForest (https://github.com/ngcrawford/cloudforest last accessed September 24, 2013). To reduce the number of parameters in our species tree estimation, we used either the JC69 substitution model, by selecting the GTR model with all base frequencies equal and unselecting the ac, ag, at, cg, and gt operators, or the GTR model with all base frequencies equal and unselecting the ac, ag, at, cg, and gt operators, or the HKY model with base frequencies set to empirical. We used the following settings for all *BEAST* analyses: a strict molecular clock with a fixed rate of 1.0 for all loci, a Yule process on the species tree prior, and a lognormal distribution with ($\mu$ = 0.0012; SD = 2.0) for the species.popMean prior and an exponential distribution for the species.yule.birthRate prior ($\mu$ = 1000). We did not include monomorphic loci in the analyses. We conducted two runs of 2 billion generations for each species and sampled trees every 25,000 generations. We determined the number of burn-in replicates to discard and assessed MCMC convergence by examining ESS values and likelihood plots using Tracer v.1.5 (Rambaut and Drummond 2010).

Modeling Demographic History

We estimated $\tau$, $\theta$, and migration rates using the single species full data sets and the Bayesian coalescent program G-PhoCS (Gronau et al. 2011). G-PhoCS is a modified version of the program MCMCcoal (Yang 2002; Rannala and Yang 2003) that allows migration among unphased diploid genotypes. G-PhoCS requires a user-specified topology to estimate model parameters, thus we used the *BEAST* populations and species tree topology for each species. For each species, we ran four models examining the impact of migration and the inclusion of loci not present in all five species on parameter estimates. The models were: ($m_1$) single species reduced, no migration; ($m_2$) single species reduced, migration; ($m_3$) single species full, no migration; and ($m_4$) single species full, migration.

G-PhoCS uses a gamma ($\alpha, \beta$) distribution to specify the prior distribution for the population standardized mutation rate parameter ($\theta_0 = 4N\mu$; for a diploid locus, where $\mu$ is per nucleotide site per generation), the species divergence time parameter ($\tau = T\mu$; $T =$ species divergence time in millions of years), and the migration
rate per generation parameter \((m_{x} \times \theta_{s}/4 = M_{s})\), which is the proportion of individuals in population \(x\) that arrived by migration from population \(s\) per generation. We evaluated a series of prior distributions ranging from shallow to deep divergences, small to large theta values, and low to high migration rates by changing the shape parameter \((\alpha)\) and scale parameter \((\beta)\), which have a mean \(\alpha/\beta\) and variance \(\sigma^2 = \alpha/\beta^2\). For the final analyses, we used two different priors for \(\tau = 0\): (1, 30) and (1, 300). These priors represent wide ranges of divergence times and effective population sizes. The \(\tau = 0\) prior distributions had no apparent influence on the posterior estimates of \(\tau\) and extant \(\theta\), except in one case where there was uncertainty surrounding the ancestral theta of NA-SA in \(C.\ lineatus\). When we increased the mean of the migration rate prior distribution, posterior estimates of migration rate had larger uncertainty, poorer mixing, and in some cases runs failed. Despite these problems, the medians of the posterior estimates in the runs that finished were qualitatively similar across the different prior distributions we tested. Thus, to achieve adequate convergence we set the migration rate prior \(\alpha\) and \(\beta\) to 1.0, 10. We ran each analysis twice for 1,000,000 generations and collected samples from the posterior distribution every 500 generations. We assessed MCMC convergence and determined burn-in by examining ESS values and likelihood plots in the program Tracer v.1.5 (Rambaut and Drummond 2010).

In the absence of a fossil or geologic calibration, we converted raw parameter estimates using a relative scaling specification of the program. BPP generates a posterior distribution of speciation models containing differing numbers of species and estimates speciation probabilities from the sum of the probabilities of all models for speciation events at each node in the guide tree. To specify the guide tree for each species, we input a fixed species tree \((rjMCMC algorithm 0)\) using the topologies estimated by *BEAST, and we performed analyses using a range of priors that represented different population sizes and different ages for the root in the species tree (Leaché and Fujita 2010). BPP uses a gamma distribution where the shape parameter \((\alpha)\) and scale parameter \((\beta)\) are specified by the user. For the final analyses, we used the same prior distributions as our G-PhoCS and MCMCcoal analyses: (1,30) and (1,300) in BPP. We modified the G-PhoCS and MCMCcoal input files to fit the formatting specifications of the program. BPP generates a Bayesian modeling approach to generate speciation probabilities of closely related taxa from multilocus sequence data. The program takes into account gene tree uncertainty and lineage sorting and assumes no gene flow among species after divergence, no recombination among loci, and the JC69 finite-sites substitution model. Included in the speciation model are the divergence time parameter \(\tau\) and population standardized mutation rate parameter \((\theta = 4N_{e} \mu_{s}\) where \(N_{e}\) is the effective population size and \(\mu_{s}\) is the substitution rate per site per generation). To generate BPP input files, we modified “.ima” files for each species to fit the formatting specifications of the program. BPP generates a posterior distribution of speciation models containing differing numbers of species and estimates speciation probabilities from the sum of the probabilities of all models for speciation events at each node in the guide tree. To specify the guide tree for each species, we input a fixed species tree \((rjMCMC algorithm 0)\) using the topologies estimated by *BEAST, and we performed analyses using a range of priors that represented different population sizes and different ages for the root in the species tree (Leaché and Fujita 2010). BPP uses a gamma distribution where the shape parameter \((\alpha)\) and scale parameter \((\beta)\) are specified by the user. For the final analyses, we used the same prior distributions as our G-PhoCS and MCMCcoal analyses: (1,30) and (1,300) and assessed convergence using the same diagnostics as our previous analyses. We ran analyses with each prior twice for 1,000,000 generations, sampling every five, and specified a burn-in of the first 50,000 generations.

### Bayesian Species Delimitation

Current species limits in tropical birds often underrepresent actual species diversity because many species’ taxonomies have yet to be evaluated with genetic methods (Tobias et al. 2008; Smith et al. 2013). To delimit the number of species within each of the five taxa, we used the program BPP v.2.06 (Rannala and Yang 2003; Yang and Rannala 2010), which implements a Bayesian modeling approach to generate speciation probabilities of closely related taxa from multilocus sequence data. The program takes into account gene tree uncertainty and lineage sorting and assumes no gene flow among species after divergence, no recombination among loci, and the JC69 finite-sites substitution model. Included in the speciation model are the divergence time parameter \(\tau\) and population standardized mutation rate parameter \((\theta = 4N_{e} \mu_{s}\) where \(N_{e}\) is the effective population size and \(\mu_{s}\) is the substitution rate per site per generation). To generate BPP input files, we modified “.ima” files for each species to fit the formatting specifications of the program. BPP generates a posterior distribution of speciation models containing differing numbers of species and estimates speciation probabilities from the sum of the probabilities of all models for speciation events at each node in the guide tree. To specify the guide tree for each species, we input a fixed species tree \((rjMCMC algorithm 0)\) using the topologies estimated by *BEAST, and we performed analyses using a range of priors that represented different population sizes and different ages for the root in the species tree (Leaché and Fujita 2010). BPP uses a gamma distribution where the shape parameter \((\alpha)\) and scale parameter \((\beta)\) are specified by the user. For the final analyses, we used the same prior distributions as our G-PhoCS and MCMCcoal analyses: (1,30) and (1,300) and assessed convergence using the same diagnostics as our previous analyses. We ran analyses with each prior twice for 1,000,000 generations, sampling every five, and specified a burn-in of the first 50,000 generations.

### Results

The partial Illumina HiSeq lane produced 255 million reads from the 40 samples following initial
processing with Illumiprocessor. Four of the eight samples of *M. marginatus* (LSUMZ 2460, 9127, 11839, 106784) failed and we excluded these from additional analyses. From the remaining 36 samples we input 249 million trimmed reads into the assembly process (Supplementary Tables S1 and S3), and assembled a total of 29 million reads into contigs across the five study species (Supplementary Table S3). Across species, mean contig length varied from 258 bp to 371 bp, and mean read depth varied from 182× to 387× (Supplementary Table S3). Between 951 and 1796 contigs aligned to UCE loci in each species, and we retained 812–1580 of these in the MAFFT alignments (Supplementary Table S3). Many alignments did not contain all individuals—each individual was present in between 360 and 1499 alignments (Supplementary Table S1).

After filtering out poor alignments with limited overlapping sequence and alignments containing <1 individual in each population, 776–1516 loci remained in the single species full data sets for the five study species (Table 1; Supplementary Table S3). From these, we recovered 166 loci shared across all five species, and these comprised the single species reduced data sets. For the interspecific alignments, we recovered 2219 total UCE loci (all species full data set), of which 169 contained at least one individual in each area of endemism across all species (all species reduced data set; Table 1).

### UCE Summary Statistics

The percentage of polymorphic loci in each species ranged from 53% to 77%, and the average number of variable sites in each polymorphic locus ranged from 2.0 to 3.2 (Supplementary Table S4). Values of Watterson’s θ and π averaged across UCEs were lowest in *Q. purpurata* and *M. marginatus*, and highest in *S. turdina* (Table 2; Supplementary Fig. S2). Values of Tajima’s *D* within species ranged from −0.22±0.86 to −0.63±0.63, although there was considerable variation across loci (Table 2; Supplementary Fig. S2).

Mean values of Watterson’s θ, π, and Tajima’s *D* differed between the single species full and single species reduced data sets in only 3 of 15 independent comparisons (Mann–Whitney U-Test *P* < 0.05; Table 2; Supplementary Fig. S2). Summary statistics calculated from cytochrome *b* were outside the confidence limits of the equivalent statistics calculated from UCEs (single species full data sets) in 12 of 15 independent comparisons (Table 2). Variation in summary statistic estimates decreased as the number of UCE loci used increased (Supplementary Fig. S3). Plots of variation versus alignment position for each species indicated that the frequency of variant bases increased with distance from the center of UCEs (Fig. 2; Supplementary Fig. S4). Plots containing all taxa showed that the frequency of variant bases reached an asymptote around 300 bp from the UCE centers and subsequently decreased. This decrease may occur because fewer alignments extend past 300 bp from center and because those alignments having data in outer flanks contain fewer individuals (Supplementary Fig. S5). The decrease in variability in outer flanks may also result from reduced read depth across these outer positions.

### Species Trees

The best-fit substitution model for the majority of UCEs across species was JC69 (Supplementary Fig. S6). Species tree estimation using the single species reduced data sets required between 17 days and 27 days (Intel Core i7 3.4 GHz) before runs converged with ESS values >200. Species tree estimation recovered generally highly supported topologies (Fig. 3a; Supplementary Fig. S7), but discordance across individual gene tree topologies (Fig. 3b). The only parameters with ESS values >200 were four population size parameters in *C. lineatus*, which we attribute to an unresolved node in that tree. Species trees were highly congruent in topology with mtDNA gene trees from the same species (Supplementary Fig. S8). Species trees for *X. minutus*, *Q. purpurata*, and *M. marginatus* were fully resolved with high support values for all nodes (posterior probabilities, PP = 1.0). In these three species there was a basal divergence across the Andes separating a clade composed of Central America (CA) and Chocó (CH) populations west of the Andes from a clade comprising all other lineages found

<table>
<thead>
<tr>
<th>Scientific name</th>
<th>Full UCE</th>
<th>Reduced UCE</th>
<th>Cytochrome <em>b</em></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>θ</td>
<td>π</td>
<td><em>D</em></td>
</tr>
<tr>
<td><em>Cymbilaimus</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>lineatus</em></td>
<td>0.021 (0.0064)</td>
<td>0.0319 (0.0055)</td>
<td>-0.486 (0.7282)</td>
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<td>0.0211 (0.0045)</td>
<td>0.0019 (0.0040)</td>
<td>-0.221 (0.8660)</td>
</tr>
<tr>
<td><em>Sicyopus</em></td>
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<td>0.0225 (0.0018)</td>
<td>-0.6335 (0.6470)</td>
</tr>
<tr>
<td><em>Querandia</em></td>
<td>0.0014 (0.0045)</td>
<td>0.0013 (0.0048)</td>
<td>-0.5382 (0.7060)</td>
</tr>
<tr>
<td><em>Macrocranium</em></td>
<td>0.0015 (0.0021)</td>
<td>0.0015 (0.0021)</td>
<td>-0.3036 (0.6838)</td>
</tr>
</tbody>
</table>

*Significantly different from equivalent summary statistic calculated from non-orthologous UCEs at *P* < 0.05 level (Mann–Whitney U test).

*Value falls outside 95% confidence limits of equivalent summary statistic calculated from all UCEs.
FIGURE 2. Variability increases in flanking regions of ultraconserved regions within each of the five study species (single species full data sets) and in alignments of all individuals across all five study taxa (all species full data set). We removed data points with no variability and outliers for clarity of presentation. The apparent decrease in variability toward the edges in the interspecific alignments likely results from differences in alignment lengths and missing/reduced data near the ends of alignments, although reduced sequence coverage in these outlying areas may also reduce our ability to accurately call variants.
FIGURE 3. a) Maximum Clade Credibility species trees with posterior probability values for each node and b) cloudograms showing the Maximum Clade Credibility tree for each UCE from *BEAST analyses of the single species reduced data sets.

east of the Andes. In *X. minutus* and *Q. purpura*, the Napo (NA) and Inambari/Rondônia (SA) populations were sister. The topology for *C. lineatus* was similar, with a basal divergence that represented a cross-Andes break. Although the trans-Andean clade (CA+CH) clade (PP = 1.0) was well-supported in *C. lineatus*, the cis-Andean clade (SA+NA) was less well-supported (PP = 0.70) with some support for an alternative topology with NA sister to the trans-Andean clade (Supplementary Fig. S7). The species tree for *S. turdina* was fully resolved with high support values (PP > 0.99; Fig. 3), with the two SA populations (SA1 = Inambari and SA2 = Rondônia) sister (PP = 1.0) and NA sister to the (CA+CH) clade (PP = 1.0).

Comparative Divergence Estimates

G-PhoCS run lengths varied from 100 h to 150 h (Intel Core i7 3.4 GHz) and all runs converged, produced similar results using different priors, and ESS values for all parameters were >1000. The inclusion of migration and the analysis of full versus reduced UCE data sets across species had minimal effects on parameter estimates (Fig. 4; Supplementary Figs. S9 and S10; Supplementary Table S7). Parameter estimates were similar across all four models (m1–m4) examined in G-PhoCS. Tau (τ) estimates (from model m4) across the Andes suggest different divergences across the five species: *C. lineatus* with mean (95% HPD intervals) = $2.64 \times 10^{-4} (2.40 – 3.00 \times 10^{-4})$; *X. minutus* = $1.04 \times 10^{-3} (9.80 \times 10^{-4} – 1.08 \times 10^{-3}$); *S. turdina* = $5.63 \times 10^{-4} (5.00 – 6.10 \times 10^{-4})$; *Q. purpura* = $4.19 \times 10^{-4} (4.00 – 4.50 \times 10^{-4})$; and *M. marginatus* = $7.42 \times 10^{-4} (6.80 – 8.10 \times 10^{-4})$. After converting τ parameters to time in millions of years using relative substitution rates, we found that understory species had older divergences across the Andes than canopy species (Fig. 4a). Divergence times across the Isthmus of Panama and Amazon River were also older in understory species than canopy species (Fig. 4a). In comparison to the UCE estimates of divergence times, eight of the mtDNA divergence time estimates were older, three were younger, and four were approximately the same (Fig. 4a). For the majority of divergences, the mtDNA time estimates had larger credible intervals than the estimates from the UCE data sets (Fig. 4a).

Effective Population Sizes and Migration

Theta and effective population sizes estimated from UCEs provided insight into demographic patterns between understory and canopy birds (Fig. 4b; Supplementary Figs. S9 and S10; Supplementary Tables S6 and S8). We converted $\theta$ values ($\theta = 4N_e \mu$) using relative substitution rates (Supplementary Table S5) and a generation time of 1 year, and we found that understory species (*X. minutus*, *S. turdina*, and *M. marginatus*) had larger effective population sizes than canopy species.
FIGURE 4. Results from G-PhoCS demographic modeling including a) divergence times for each population divergence event in each study species based on output from G-PhoCS. Results of all four models are presented: (m3) single species full, no migration; (m4) single species full, migration; (m1) single species reduced, no migration; and (m2) single species reduced, migration. Divergence events are labeled by listing the populations included in both of the resulting clades. b) Effective population sizes (Ne) for all extant populations in each study species. Results of two models are presented: (m4) single species full, migration and (m2) single species reduced, migration. (C. lineatus and Q. purpurata) in all models (Fig. 4b; Supplementary Fig. 59). Ne estimates were similar with and without the inclusion of migration in the model (Supplementary Fig. 59), and Ne estimated from the single species reduced data sets produced values similar to those estimated from the single species full data sets (Fig. 4b).

Migration rates were non-zero in all analyses including migration, but migration rate estimates were sensitive to the prior distribution used. Understory species had higher mean values of migration parameters than canopy species, but credible intervals were large and included values near zero for all migration estimates (Supplementary Fig. S10). Inferred migration estimates were similar using both the single species reduced and single species full data sets (Supplementary Fig. S10).

Species Delimitation

BPP run lengths varied from 120 h to 200 h (Intel Core i7 3.4 GHz) with the independent runs producing similar results and all runs converging with high ESS values. The analyses were stable among independent runs and the different priors produced similar results for most speciation events. The number of inferred species within each species from the BPP analysis varied from two to five: C. lineatus = 2, X. minutus = 4, S. turdina = 5, Q. purpurata = 2, and M. marginatus = 2. Speciation probabilities were sensitive to prior distributions for some nodes in S. turdina and C. lineatus (Supplementary Fig. S11). Overall, speciation probabilities were higher in the understory species than in the canopy species (Supplementary Fig. S11).
phylogenetic markers (Crawford et al. 2012; Faircloth highlighted the utility of UCEs as a source of informative presented here complement previous studies that demographic histories for all species. The UCE results fewer species. We used these data sets to reconstruct the shared loci and loci that we recovered from four or fewer species. We used these data sets to reconstruct demographic histories for all species. The UCE results presented here complement previous studies that highlighted the utility of UCEs as a source of informative phylogenetic markers (Crawford et al. 2012; Faircloth et al. 2012, 2013; McCormack et al. 2012a, 2013), and extend the application of UCEs to studies of comparative phyleogeography at the species and population level. The success of UCEs in both population genetic inference and in resolving phylogenetic relationships at deep and shallow evolutionary time scales confirms their utility in both micro- and macro-evolutionary investigations.

Data Generation, UCE Variability, and Computation

We assembled 31.8% of the trimmed reads into contigs and only 27.4% of assembled contigs mapped to UCEs, suggesting that we recovered a high proportion of off-target reads. We were able to assemble nearly complete mitochondrial genomes for most samples using raw read data, suggesting that many of the off-target reads represent mtDNA “contamination”. This contamination may be due to the high copy number of mitochondria in the tissues we used. It is remarkable that despite this limitation, we were able to obtain data matrices of 166 shared loci from all five species. Looking forward, improved enrichment efficiency due to optimized lab protocols should lead to better capture of UCEs, and technological improvements in MPS may result in longer reads producing longer contigs.

Levels of variation in the flanking regions of UCEs are comparable to those found in sequences and introns recovered from Sanger sequencing of nuclear DNA within species. We found that 53–77% of UCEs in each species were polymorphic and UCEs contained 2.0–3.2 SNPs on average within species, or an average of one SNP every 222–550 bp. These SNP discovery rates are similar to intraspecific studies that found, for example, one SNP per 142 (Geraldes et al. 2011) or 518 (Bruneaux et al. 2013) bp, and one polymorphism every 16–157 bp in introns (Smith and Klicka 2013). Although 2.0–3.2 SNPs per locus may result in low gene tree resolution, methods that integrate low information content across many loci, along with longer contigs due to MPS improvements, may further improve the utility of UCEs for future studies.

To examine the effects of the number of loci on parameter estimation, we compared summary statistics estimated from subsamples of the UCE data sets containing different numbers of loci. As expected, confidence intervals generally decreased as the number of UCE loci used in the analysis increased (Supplementary Fig. S3). We also compared the genetic parameters estimated from the single species reduced data sets (166 loci) to those estimated from the single species full data sets (776–1516 loci) and mtDNA. Again, credible intervals generally decreased as the number of loci used to estimate parameters increased (Fig. 4a). The single species full and single species reduced UCE data sets produced similar summary statistics and mean parameter estimates, and the chronologic sequence of divergence events across the five species was consistent between data sets. It remains unclear how important it is that large UCE data matrices be composed exclusively of the same loci when drawing comparisons across species or data sets. Parameters estimated in G-PhoCS from the single species reduced and single species full data sets were similar. There were analytical limitations that did not allow us to compare parameters estimated from both data sets in *BEAST and BPP. For some analyses (e.g., G-PhoCS) we were able to efficiently and rapidly analyse the single species full data sets (~100 h on a 3.4 GHz Intel Core I7 processor). In contrast, preliminary analyses using the single species full data sets in *BEAST and BPP indicated that runs would take weeks to months to finish. Our results suggest that data sets composed of large numbers of non-intersecting loci may be sufficient for many comparative phyleogeographic applications, but additional data and faster computational methods are needed to explore this further.

With highly multilocus UCE data we were able to estimate θ with relatively high precision relative to mtDNA, but credible intervals on migration rates were large, highlighting the general difficulty of measuring dispersal parameters (Broquet and Petit 2009). Population divergence times estimated from mtDNA tended to be older than those estimated from UCEs (Fig. 4a), consistent with the idea that, because gene trees are embedded within species trees, analyses based on single-gene trees will overestimate divergence times (Edwards and Beerli 2000). Three divergence estimates are shallower in the mtDNA estimate than the multilocus UCE estimate (Fig. 4a), implicating either recent mitochondrial gene flow or the fact that, in rare cases, gene divergence is shallower than the confidence intervals associated with population divergence (Edwards and Beerli 2000). Information on theta and migration between populations in highly multilocus data sets, such as those generated using UCEs, provides another layer of information that complements what has been available historically using mtDNA.

Comparative Phylogeography in Neotropical Forest Birds

Our phylogeographic analyses based on UCEs indicated that co-distributed taxa exhibit discordant
evolutionary histories. Divergence time patterns inferred from UCEs are consistent with recent studies that show multiple speciation events across the Andes (Brumfield and Edwards 2007; Miller et al. 2008), Amazonian rivers (Naka et al. 2012), and the Isthmus of Panama (Smith et al. 2012b). Previous comparative phylogeographic studies on Neotropical birds have not assessed the influence of gene flow on inferred patterns. However, the inclusion of migration in our coalescent modeling had a negligible influence on divergence time estimates for these species. Since our migration rates were sensitive to the prior distribution and substitution rates for UCEs were low, it is unclear whether the consistency in divergence times across our models represents limited gene flow across barriers or insufficient information in the UCE data sets to accurately estimate gene flow. Despite uncertainty around the robustness of the migration rate estimates, the inclusion of migration in the coalescent modeling allowed us to relax the pure isolation assumption. Based on the BPP analyses, understory species typically had higher speciation probabilities or more inferred species. Some nodes had low (PP < 0.95) speciation probabilities, likely because of gene flow among endemic areas and/or shallow divergences among areas. Overall, divergence time and species delimitation estimates were largely concordant with the results of a previous mtDNA study that showed greater genetic differentiation in poorly dispersing understory species than in more vagile canopy species (Burney and Brumfield 2009).

These UCE data provide the first genomic insight into the effective population sizes of rainforest birds, a critical parameter for inferring how species have evolved in response to past events. We found that understory birds (X. nitidus, S. tundina, and M. marginatus) have larger effective population sizes than species that inhabit the canopy (C. lineatus and Q. purpurata). This finding, if corroborated with a larger sample of species, has important implications for understanding how species ecology influences genetic differentiation, turnover rates among ecological guilds, and the accumulation of diversity over time. Canopy species have larger territory sizes and occur at lower densities than understory species (Munn and Terborgh 1985; Terborgh et al. 1990). Canopy species may also have more dynamic or unstable distributions and population sizes (Greenberg 1981; Loiselle 1988), particularly Q. purpurata, which is a large frugivore and may track ephemeral sources of fruit across the landscape. The observation that C. lineatus and Q. purpurata have less divergent populations is consistent with the expectation that species with small or unstable population sizes will be less likely to accumulate genetic diversity over time (Leffler et al. 2012). Our results provide additional preliminary evidence that ecology is important in predicting the population histories of tropical birds. As precision in the estimation of population genetic parameters improves with multilocus data, so too will our ability to elucidate the linkage between ecological and evolutionary processes.

Supplementary Material

Data files and/or other supplementary information related to this paper have been deposited at Dryad (http://datadryad.org/) under doi:10.5061/dryad.qm4jI.

Funding

This research was supported by National Science Foundation (DEB-0841729 to R.T.B.).

Acknowledgments

We thank D. Dittmann (LSUMNS), D. Willard (FMNH), J. Klicka (UWB,M), M. Robbins (RU), and A. Aleixo (MEPG) for providing samples, S. Herke for assisting with laboratory work, L. Yan for assistance with de novo assembly on the LSU HPC Philip cluster, and I. Gronau for help running G-PhoCS. We thank S. Hird for writing custom scripts to process our data in lociNCS. We thank A. Cuervo and E. Rittmeyer for providing helpful assistance during data generation and analysis. F. Anderson, M. Charleston, and two anonymous reviewers provided comments that greatly improved this manuscript.

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We thank D. Dittmann (LSUMNS), D. Willard (FMNH), J. Klicka (UWBM), M. Robbins (RU), and A. Aleixo (MEPG) for providing samples, S. Herke for assisting with laboratory work, L. Yan for assistance with de novo assembly on the LSU HPC Philip cluster, and I. Gronau for help running G-PhoCS. We thank S. Hird for writing custom scripts to process our data in lociNCS. We thank A. Cuervo and E. Rittmeyer for providing helpful assistance during data generation and analysis. F. Anderson, M. Charleston, and two anonymous reviewers provided comments that greatly improved this manuscript.
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