Electronic supplementary material for "Why is Amazonia a 'source' of biodiversity? Climate-mediated dispersal and synchronous speciation across the Andes in an avian group (Tityrinae)"

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Introduction:

Here we present our supplementary methods and results for genetic locus filtering, scoring of taxa for ancestral ecological analyses, bioinformatics, phylogenomic methods, macroevolutionary models, and additional discussion of biogeographic history and macroevolutionary patterns.

Supplementary Methods

Sequence capture of UCEs

We performed target-capture with the Tetrapods-UCE-2.5kv1 probe set (available at ultraconserved.org), which target 2386 UCEs. All individuals were previously extracted and many were previously sequenced for another study (Musher and Cracraft, 2018). For the remaining individuals, we sent extracts to Rapid Genomics (Gainesville, FL) for DNA shearing, library preparation, and UCE enrichment following the protocol detailed in Faircloth et al., (2012). Sequencing was performed on an Illumina HiSeq 2500.

Bioinformatic processing of UCEs

After sequencing, we cleaned data by removing low quality reads and trimming low quality base-pairs. We assembled reads using velvet (Zerbino and Birney, 2008) and matched these assembled contigs to the probes using prewritten python scripts from Phyluce (Faircloth et al., 2012). For one outgroup taxon, *Acanthisitta chloris*, we extracted UCE loci from assembled genomic read data available online (Jarvis et al., 2014). We matched the assembled contigs to UCE probes and incorporated the identified UCE markers into the sequence capture dataset explained above. Finally, we aligned orthologous loci for our final taxon set using MAFFT (Katoh and Standley, 2013). We chose to include all loci for which 95% of the individuals had sequence (henceforth, 95% complete dataset) for our molecular analyses.

Genetic locus filtering

We wrote custom scripts in R to quantify 1) the total number of parsimony informative sites, 2) the proportion of parsimony informative sites, and 3) two measures of clock-likeness for each marker. The latter were obtained by, first, using the chronopl rate-smoothing algorithm implemented in the R package "Ape" (Paradis et al., 2004) to estimate the optimal smoothing parameter (λ) using cross validation (compared multiples of 10 between 10^{-6} and 10^{2}), and then by performing likelihood ratio tests (clocklike to non-clocklike molecular models) at each locus

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(see below). λ is a measure of clock-likeness, with higher values at a locus representing gene trees that have relatively even terminal branch lengths (i.e., gene tree shape is close to ultrametric), and lower values representing gene trees with more uneven terminal branch lengths.

Following Doyle et al. (2015) our intention was not to accept or reject a molecular clock at each locus, but rather to take loci that most closely fit with a clocklike model of molecular evolution (i.e., lower likelihood ratio). We used RAxML (Silvestro and Michalak, 2012) to build maximum likelihood (ML) gene trees at each locus using an automated shell script. We then imported these trees into R and calculated the likelihood of the gene tree, which represents a non-clocklike model estimate where branch lengths are calculated independently. We then calculated the likelihood of the gene tree given a clocklike model (i.e., an ultrametric model). To do this we used the Likelihood ratio, *LR*, which is given by two times the difference between log-likelihood of the clocklike model and log-likelihood of the non-clocklike model:

$$LR = 2 \bullet (lnL_{clocklike} - lnL_{nonclocklike})$$

In order to identify the markers that were most informative for divergence dating, we built a multivariate model in R that modeled LR as a function of the total variable sites, proportion of variable sites, and $\log_{10}(\lambda)$. As we are interested in divergence dating, clocklike markers are desirable, and those that deviate from clocklike tendencies were identified as outliers using Cook's distances (D). D values that were greater than three times the mean D were considered outliers, and removed from downstream analyses. Stated alternatively, loci that have exceptionally high or low values of LR, proportion of parsimony informative sites, or λ overinfluence the model result in high values for D and are considered outliers. Because Bayesian estimation of divergence times can be intractable for large molecular datasets, after removing outliers, we chose the 15 most clocklike (lowest LR) loci to use in divergence dating analysis. Our filtering regime is similar to that of Smith et al., (2018) in that we sort our markers by clock-like tendency, however we did not filter based on gene tree topology, and instead identify outlier loci by measuring the Cook's distance deviation from a linear model.

Phylogenomic analyses

In order to obtain a tree topology for divergence dating (see below), we concatenated all loci for the 95% complete dataset, and estimated the topology using RAxML (Silvestro and Michalak, 2012). We performed 20 Maximum Likelihood (ML) searches, under a GTRCAT model of substitution. The GTRCAT model approximates a GTR + Gamma model of sequence evolution (Stamatakis, 2006). We then performed bootstrapping using the autoMRE option, which halts bootstrapping after replicates converge. In our previous study on *Pachyramphus*, concatenated and species tree methods gave nearly identical topologies but species tree approaches experienced difficulty in placing individuals with relatively poor-quality sequence (Musher and Cracraft, 2018; see also Hosner et al., 2015). Because we included several individuals sequenced from museum study skins, which are likely to have poorer quality sequence than fresh tissue and can bias species tree results (e.g., Hosner et al., 2015; Musher and Cracraft, 2018), we chose not to use species tree approaches on this dataset, as the relationships in this group seem robust to methodological choices.

Reanalysis of the Claramunt and Cracraft (2015) dataset

The dataset from Claramunt and Cracraft (2015) used slow evolving nuclear genes (RAG1 and RAG2) and a fossil-calibrated dataset for family representatives across all Neornithes. We sampled RAG1 and RAG2 sequences available on GenBank (Tello et al., 2009)

for four additional taxa not included in the original study: *Schiffornis turdina, X. albinucha, P. polychopterus, and P. aglaiae*. We then incorporated these sequences into the original RAG dataset, keeping all priors and parameters identical, including fossil calibrations, and used BEAST v2.4.3 (Bouchaert et al., 2014) to generate a time tree. We ran the Markov Chain Monte Carlo (MCMC) with thinning of 1,000 for 1.9 x 10⁸ generations. We assessed convergence using Tracer v1.6.0 (Rambaut et al., 2014), and then summarized the posterior distribution as a maximum clade credibility tree after a burnin of 25 percent.

MCMCTree Priors

We used the results from the Claramunt and Cracraft (2015) reanalysis to apply calibrations to our dataset of filtered UCE markers. We specifically obtained 95% credible intervals (CI) of node ages for crown Passeriformes, stem Tyranni, crown Tyrannides-Furnariides, crown Tyrannides, and crown Tityridae (taxonomic names *sensu* [35]). The locus substitution rate is defined by a gamma prior for mean substitution rates, $\Gamma(\alpha,\beta)$, where rate mean is equal to α/β and rate variance is equal to α/β^2 . We applied a prior distribution of $\Gamma(2,200000)$, which defines a mean rate of 10^{-5} substitutions MY⁻¹ assuming a one-year generation time. For the birth-death model, we applied uniform priors for birth and death rates of one, and sampling of 0.90. We then ran the MCMC for 1.5 x 10^6 generations following a burnin of 5 x 10^5 generations. We assessed convergence using Tracer v1.6.0 (Rambaut et al., 2014).

Ancestral ecological modeling and biogeographic inference

We quantified the amount of *in situ* speciation, lineage emigration, and lineage immigration for each biogeographic area in our Dispersal Extinction Cladogenesis (DEC) model - Amazonia, Atlantic Forest, Andes, arid diagonal, lowlands west of the Andes, and Central America. To do this, we defined total *in situ* speciation for each area as the total number of parent nodes whose maximum likelihood (ML) state contained at least one matching area in the maximum likelihood state of both of its daughter nodes. For example, if a node's maximum likelihood state was Amazonia, and both of its daughter nodes also contained Amazonia in their ML states, we considered this a single *in situ* speciation event for Amazonia. We defined total lineages emigrating from a given area as the total number of novel ML area states at daughter nodes stemming from all parent nodes whose ML state was the area of interest. For example, if Amazonia is the state of the parent node and one of its daughters contained an area other than Amazonia, we counted this as a single emigration event for Amazonia. And finally, we defined total immigration for each area as the number of daughter nodes whose parent node did not contain at least one of the same areas. For example, if a given daughter node contained Amazonia, but its parent node did not, we considered this a single immigration event for Amazonia.

For habitat affinity, we coded each species based on the its habitat codes found in Parker et al. (1996), and additional information where available (Fitzpatrick et al., 2004). Species found in F1 (Tropical lowland evergreen forest) or F4 (Montane evergreen forest) were coded as having an affinity for forest interior, while those found in other habitats, including F1E (edge) or F4E, were coded as having an affinity for semi-open habitat (this includes semi-open forests such as flooded, seasonally dry, white sands, and gallery). Taxa that use both types of habitat were coded as polymorphic. We constructed a model in RevBayes (Hohna et al., 2016) that allowed for equal probabilities of state addition and loss in habitat preference by employing a DEC model (Ree and Smith, 2008) that is implemented within the RevBayes language. We

chose a DEC model, designed for biogeographic analysis, rather than a discrete character evolution model because habitat is spatially distributed, and transitions occur in the same ways that dispersal into novel regions occurs – as a time-dependent process. Still, fitting a DEC biogeographic model, is akin to fitting a MK model of discrete character evolution. We put an exponential prior with mean of one state change per 20my per lineage. We ran the MCMC for 10,000 generations and summarized posterior probabilities of each ancestral state after a burnin of 25 percent. Scripts are available at https://github.com/lukemusher/Tityrinae biogeography.

Macroevolutionary rates

To estimate diversification rates through time and among lineages, and to identify potential speciation or extinction rate shifts, we used both Bayesian Analysis of Macroevolutionary Mixtures (BAMM; Rabosky et al., 2014) and TESS (Hohna et al., 2015). TESS uses a reversible-jump MCMC (rjMCMC) algorithm to estimate the timing of rate shifts in speciation and extinction under an episodic birth-death process. We assumed 85% complete taxon sampling for the subfamily, expected number of rate changes of three (boundaries between epochs since Miocene), and zero mass extinctions. We estimated empirical hyperpriors for speciation and extinction. We ran the TESS analysis for 10⁶ generations and assessed convergence in parameter estimates using Tracer v1.6.0 (Rambaut et al., 2014). BAMM also uses an rjMCMC to quantify diversification rate heterogeneity along a phylogeny, and we ran this analysis for 5 x 10⁷ generations with thinning of 10⁵ and a burn-in of ten percent. We optimized priors using the "setBAMMpriors" function implemented in the R package BAMMtools (Rabosky et al., 2014).

To test for diversity-dependent diversification, we also applied a model-fitting approach implemented in the R package, DDD (Etienne et al., 2012). We fit six models, four of which explicitly incorporate a parameter for carrying capacity (K): (1) birth-death constant rates without carrying capacity, (2) linear dependence of speciation rate on K without extinction, (3) linear dependence of speciation rate on K with extinction, (4) exponential dependence of speciation rate on K, and (6) exponential dependence of extinction rate on K.

We then used the results of the TESS analysis to quantify the effect of palaeotemperature on net-diversification rate (speciation rate – extinction rate). We first divided the posterior net-diversification rates from TESS into 50 evenly-spaced intervals between the basal split and tips, and quantified mean posterior net-diversification rate for each interval in TESS. Then, using palaeotemperature data available in RPANDA (Morlon et al., 2016; Condamine et al., 2013), we quantified the relationship between temperature and net-diversification. To obtain temperature values for each interval, we took the mean of all temperature measurements within each time interval. We then evaluated their correlation using linear regression and compared three generalized linear models where net-diversification is either dependent on time, temperature, or interactive effects of both time and temperature.

Finally, as a more sophisticated test of environmentally-driven diversification we applied one additional model selection implemented in RPANDA (Morlon et al., 2016; Condamine et al., 2013), which explicitly models speciation and extinction rates and optimizes parameters. We fit ten models to our divergence time results from MCMCTree. We specifically fit (1) pure birth constant rate, (2) birth-death constant rates, (3) speciation varies exponentially with time and no extinction, (4) speciation varies exponentially with time and constant extinction, (5) extinction varies exponentially with time and constant speciation, (6) speciation and extinction vary

exponentially with time, (7) speciation varies exponentially with temperature and no extinction, (8) speciation varies exponentially with temperature and constant extinction, (9) extinction varies exponentially with temperature and constant speciation, and (10) speciation and extinction vary exponentially with temperature. To evaluate the effect of using stem or crown ages, we also employed this model selection twice, once conditioning each model on stem ages and once on crown ages.

For all model selection sets we compared likelihoods of each model and chose the model with the lowest corrected Akaike Information Criterion (AICc). Because in some cases multiple models differ only slightly in AICc, the relative fit of each model was assessed using AICc weights (AIC ω), which allow a more intuitive interpretation of the fit of each model relative to other models. AIC ω represents the conditional probability for each model. All scripts are available at https://github.com/lukemusher/Tityrinae_biogeography.

Supplementary Results

Data filtering

Our 95% complete UCE dataset contained 1,160 loci with a total of 825,445 bps. Using this concatenated alignment, RAxML recovered a topology identical to our previous study (Musher and Cracraft, 2018), except for the position of *Pachyramphus minor*, which was recovered with weak support (ML bootstrap = 55). We constrained the position of *P. minor* for divergence dating based on the more robust relationship recovered in our previous study (Musher and Cracraft, 2018). Because UCE datasets may contain aberrant loci that can cause problems for downstream analyses, we quantified informativeness and clock-likeness of each locus via two measures, λ and LR, and explored their relationship in addition to their relationship with locus informativeness (total number and proportion of informative sites)(Figure S1).

The number of informative sites per locus ranged from 4 to 273 (mean \pm standard deviation = 99.22 \pm 46.45), the proportion of informative sites per locus ranged from 0.0053–0.39 (mean = 0.14 \pm 0.063), and the LR for each locus ranged from 2007.04–25800.57 (mean = 7360.096 \pm 3608.302). Locus informativeness was correlated with both clocklike measures, but informativeness predicted significantly more of the variance for LR than for λ (total informative sites with $\log_{10}\lambda$: Adj. R² = 0.05708, F= 71.16, p = 0.0; Proportion informative sites with $\log_{10}\lambda$: Adj. R² = 0.05731, F= 71.46, p = 0.0; Total informative sites with LR: Adj. R² = 0.1555, F= 214.4, p = 0.0; Proportion informative sites with LR: Adj. R² = 0.1812, F= 257.5, p = 0.0). LR and $\log_{10}\lambda$ were also weakly correlated (Adj. R² = 0.02058, F= 25.35, p = 0.0).

Our glm that modeled LR as a function of informativeness and λ did not find λ to be a significant predictor in the model (p=0.073) but removing lambda did not improve the model's fit to the data (with λ AIC = 22054, without λ AIC = 22055). Our filtering protocol identified 83 outlier loci, which were primarily either loci that were very un-clocklike (high LR) given their informativeness, or very uninformative given their low LR (relatively few informative sites). Removing these loci from the dataset improved the relationship between locus informativeness and clock-likeness (Total informative sites with LR: Adj. R² = 0.1791, F= 235.8, p = 0.0; Proportion informative sites with LR: Adj. R² = 0.2105, F= 287.8, p = 0.0; Figure S1), and significantly improved the quality of the model fit (AIC = 15,507). Figure S2 shows 25 randomly-chosen examples of gene trees of loci that were identified as outlier loci (Figure S2A) and the 15 markers used in our divergence dating analysis (Figure S2B). Loci identified as outliers tended to have high values for LR with relatively low informativeness and were aberrant

in shape (mix of relatively long and relatively short branch lengths across the tree). After removing outlier loci, our 15 lowest LR UCEs (Figure S2B) ranged in LR from 2475 to 2909.

Time calibrated phylogeny

We reconstructed a time-calibrated phylogeny using the Claramunt and Cracraft (2015) dataset and recovered a timeframe of diversification consistent with the previous study (Figure 1). The BEAST analysis converged after 1.9 x 10⁸ generations, except for a few parameters associated with the Birth-Death Model (see Claramunt and Cracraft, 2015), which had Effective Sample Size values slightly below 200. The analysis recovered a crown age of 53.65 (95% HPD = 49.18–58.02) mya for crown Passeriformes, 49.89 (95% HPD = 45.81–53.83) mya for stem Tyranni, 42.53 (95% HPD = 37.72–47.46) for crown Tyrannides-Furnariides, 38.75 (95% HPD = 27.44–39.5) mya for crown Tyrannides, and 26.14 (95% HPD = 19.75–32.66) mya for crown Tityridae, which were used as secondary node calibrations for the MCMCTree analysis. The BEAST phylogeny additionally recovered ages of 17.11 (95% HPD = 9.98–24.25) mya for the split between *Xenopsaris* and *Pachyramphus* (stem *Pachyramphus*), 14.76 (95% HPD = 7.68–22.1) mya for crown *Pachyramphus*, 19.97 (95% HPD = 12.81–26.91) for the split between *Tityra* and *Xenopsaris+Pachyramphus* (stem *Tityra*).

The results of our MCMCTree on filtered UCE loci converged, with all ages and parameter values that showed high ESS values above 500. We recovered an age of 22.9 (95% HPD = 18.80–26.27) mya for the crown split of Tityrinae, 8.53 (95% HPD = 5.33–12.33) mya for *Tityra*, 11.68 (95% CI = 8.84–14.88) mya for crown *Pachyramphus*, and 2.51 (95% HPD = 0.58–6.01) mya for crown *Iodopleura*. The split between *Xenopsaris* and *Pachyramphus* was recovered to be 13.44 (95% HPD = 10.29–16.89) mya (Figure 1). Higher-taxon ages from each tree and from an additional study (Prum et al., 2015) are summarized in Table S2.

Macroevolutionary rates

TESS analyses of episodic diversification found a trend of increasing speciation and decreasing extinction rates through time (Figure S3). In BAMM, the best shift configuration (maximum posterior probability of rates through time) found increasing speciation and net-diversification rates through time, and constant extinction rates through time (Figure S3). Our best model in DDD was a model without carrying capacity (AICc = 222.151, AIC ω = 0.354), but this model was only a marginal improvement over the five other models (AIC ω = 0.106–0.237). Four of these models, however, estimated values of K ranging from ~190,000 to infinity. The final model estimated K to be nearly 2000. Taken together these results suggest that there is no meaningful effect of K on net-diversification rates.

We additionally found a relationship between palaeotemperature and net-diversification rates (slope = -0.03279 ± 0.003222 , intercept = 0.2735 ± 0.02143 , p < 10^{-15} ; Adjusted R-squared = 0.6833, F = 103.6. df = 48, p < 10^{-12}). The best-fitting glm was one where net-diversification varies as a function of interactive effects of time and temperature (Table S4). Model selection in RPANDA, however, showed that a model where speciation rate varies exponentially with time (AICc = 231.9682, AIC ω = 0.2660) is marginally better-fitting than a model where speciation rate varies exponentially with temperature (AICc = 232.3937, AIC ω = 0.2151), though the difference in AIC ω was not significant (Table S5). Estimates of ω (exponential scaler for speciation rate) were negative for all RPANDA models indicating that speciation rate increases with decreasing temperature and with decreasing time before present.

Supplementary Discussion

A novel method for filtering of sequence capture data

We designed a novel model-based framework for identifying aberrant loci to remove from high-throughput sequencing datasets. Large molecular datasets, such as those resulting from sequence capture approaches, can be highly variable in locus informativeness and utility. While most modern techniques for estimating phylogeny and divergence times are robust to genetree discordances due to incomplete lineage sorting, deep coalescence, and (sometimes) hybridization, gene tree error can be a major source of noise and bias in molecular datasets (Richards et al., 2018). Similarly, high throughput datasets can be very difficult to analyze due to their size and analyzing these data is often intractable over reasonable time frames (Smith et al., 2018). Thus, filtering and subsampling data are often necessary steps for improving efficiency of analyses.

Our filtering protocol follows previous studies' attempts to quantify both informativeness and clock-likeness of molecular markers based on parsimony informative sites and likelihood ratios, respectively (Doyle et al., 2015; Manthey et al., 2016). We evaluated these metrics for our molecular dataset and found that informativeness and clock-likeness were correlated. We used this information to model LR as a function of informativeness and identified those loci which were most influential in the model, under the assumption that those loci could be a source of noise. Those loci contained primarily very un-clocklike genes, though a handful of genes were more clocklike, including two which were among the most clocklike markers in the dataset (Figure S1). Thus, identifying outliers was worthwhile, even though we only chose to use the 15 most clocklike markers in the filtered dataset. Had we not identified outliers first, some of the markers may have biased our results. More work is needed in order to evaluate the efficacy of this protocol in comparison to, for example, randomly sampling markers.

Macroevolutionary rates

Virtually all models estimated parameters that were consistent with positive net-diversification rates and thus increasing diversity through time (Figures 3, S3, S4, S5, and S6). Only TESS estimated negative net-diversification rates early in Tityrinae's history, but this result is contrary to the bulk of macroevolutionary modeling results. Although BAMM also showed an overall increase in rates of net-diversification over time, speciation rate was always higher than extinction rate. Similarly, all models in DDD estimated speciation rates to be greater than extinction rates, thereby supporting the observed pattern of increasing diversity through time, and negative values for α in all of the RPANDA models are also consistent with increasing net-diversification rates toward present (Table S5; Figure S6). Negative α in temperature-dependent models indicates that speciation rates increase with decreasing temperature. Because contemporary temperatures are lower close to the present, this result is consistent with increasing net-diversification at present. Similarly, negative estimates for α in time-dependent models show that speciation rate increases with decreasing time before the present. Therefore, speciation rates are highest at time = 0 (present) and lowest at time = \sim 23 (Tityrinae origin).

The best-fitting DDD model was one where K was not estimated, thereby rejecting models of diversity-dependent diversification. Interestingly, models that did incorporate K estimated K to be extremely high (Table S3). For example, all DDD models except one estimated K to be at least one order of magnitude higher than the total number of extant bird species (10,000–20,000 avian species in total; e.g., Barrowclough et al., 2016) and thus four orders of magnitude higher than all extant Tityrinae. Even the model with the lowest estimate of

K (the third-best model) estimates K to be nearly 2000 species. These abnormally high estimates indicate that K has no effect on net-diversification and that diversity-dependent mechanisms have had no influence on Tityrinae diversification.

We also statistically evaluated the effect of temperature on net-diversification. Although we find a strong negative relationship between palaeotemperature and net-diversification rates (Figures 3, S5, and S6), we lacked robust statistical support to show that temperature alone explains diversification rates better than time (Table S5; Figure S6). For example, although temperature explains nearly 68% of the variance in net-diversification rates through time estimated in TESS, time explains roughly 94%. Our best-fitting glm, on the other hand, was a model where net-diversification varies with interactions of time and temperature. That is, under the best glm, net-diversification varies with time, but the effect of time on net-diversification also varies with temperature. Therefore, based on the glms, time and temperature are both important parameters influencing diversification.

As an independent test of climate-mediated diversification, RPANDA resulted in statistical uncertainty. When conditioning on stem ages we showed that a model where speciation rate varied as a function of palaeotemperature was nearly as well-fitting as a model where rates only varied with time, and we find similar results when conditioning on the crown ages. Here, time may be the most important predictor of net-diversification rate, but this observation is still consistent with a hypothesis of barrier displacement if the rate of barrier formation and degradation is relatively constant through time (e.g., see Albert et al., 2017). While palaeotemperature has likely played some role in influencing barrier displacement, temperature is just one parameter out of a large number of variables that may influence barrier change. For example, additional unmeasured factors such as precipitation, erosion, and tectonics are also important parameters in a model of barrier displacement, but would be difficult (if not impossible) to measure and factor into our analyses.

We also suspect that imprecise estimates of divergence times in our dated phylogeny could bias the RPANDA results and limit the fit of the models to the data, and that our dataset lacks the power to distinguish among models due to relatively small sample size. For example, it would be very difficult to measure slight increases and decreases in speciation rate over relatively short periods on the order of 10⁴ years based on a tree with 50 terminal taxa with posterior densities of node ages in the order of 10⁶ years (Figure 1). However, that same time frame of 10⁴ years is the temporal scale in which we see many dramatic shifts in global temperature. Therefore, levels of temporal precision in the temperature dataset are far finer and more exact than levels of precision in our divergence times and subsequent net-diversification rate estimates. The problems of sample size and precision are also reflected in the wide posterior densities surrounding parameter estimates in TESS (Figures 3 and S3). Furthermore, on a spatial scale, temperature is likely imprecise because we use global measurements, which do not pick up local effects driven by, for example, Andean orogenic uplift.

Because of temporal imprecision, net-diversification rate estimates probably become somewhat smoothed through time. We attempted to circumvent the problem of disparate degrees of temporal precision by "smoothing" temperature measurements. To do this, we took mean temperatures during ~450,000-year intervals (Table S4). By averaging temperatures over longer periods of time, we reduce the temporal precision of the temperature dataset, thereby improving comparability of the two datasets (i.e., diversification rates and temperature). Remarkably, this resulted in a statistically significant result that showed that a model where net-diversification rate

varies in relation to interactions of time and temperature greatly outperforms the other models where net-diversification rate varies in relation to time or temperature alone (Table S4).

Nevertheless, more data are needed in order to more robustly determine the relationship between temperature and diversification. Even though we lack a robust causal relationship between temperature and net-diversification rate, we suggest that future studies should further investigate this relationship in detail. Specifically, studies that sample large clades (hundreds or thousands of species) at dense taxonomic scales (e.g., subspecies, phylogenetic species, or "coalescent" species) and use rigorous fossil-calibrated divergence dating based on genome-wide markers may provide increased precision of divergence times and rates of diversification through time. Additionally, modeling diversification as a function of climate at more local scales (i.e., South American instead of global) and including measures of precipitation, if possible, might assist in teasing apart the relative effects of time and palaeoclimate.

Diversification in Amazonia

Amazonia is known for its high species richness and debates over how this diversity was generated have been ongoing for decades (Haffer, 1969; Cracraft and Prum, 1988; Smith et al., 2014). All early divergences within Tityrinae appear to have occurred within proto-Amazonia but are not easily-explained by the presence of current barriers that separate extant clades. This is probably due the dynamic history of those barriers through landscape changes over the last ten million years. For example, the Aripuana River, which was likely an important historical barrier, has since become a less effective dispersal barrier (Fernandes et al., 2012, 2014; Ferreira et al., 2017). Within-Amazonian splits were clearly important for generating diversity in Tityrinae from the Miocene through the Pleistocene, as in situ speciation in Amazonia was exceptionally high (Figure 2). Although the timing of the establishment of the Amazon River drainage is still debated (Campbell et al., 2006; Hoorn et al., 2010; Nogueira et al., 2013), the existence of a massive mega-wetland system, called Pebas, in western Amazonia during the Miocene is wellknown based on fossil vertebrates, palynology, and sedimentology (Latrubesse, et al., 2010; Antoine et al., 2015; Salas-Gismondi et al., 2015; Tejada-Lara et al., 2015). Similarly, Miocene Amazonia saw a series of marine transgressions that may have separated populations of organisms in the Andean foothills, Guiana Shield, and Brazilian Shield (Aleixo and Rossetti, 2007). Transgressions are thought to have occurred at roughly 15-13 mya and possibly again at 10-5 mya (Hernandez et al., 2005; Aguilera et al., 2013; Jaramillo et al., 2017), which roughly corresponds to early splits in Tityrinae. In combination with a dynamic fluvial landscape that postdates the Miocene (Campbell et al., 2006; Latrubesse et al., 2010; Hoorn et al., 2010; Ribas et al., 2012), these marine transgressions may have contributed to high in situ speciation for Tityrinae in Amazonia (Figure 2).

Atlantic-Amazonian connections

Much variance exists in the timing of speciation between taxon-pairs between Atlantic and Amazonian forests (see main text). For example, splits between *P. m. marginatus* and *P. m. nanus* across this region are likely tied to relatively old Amazon-Atlantic forest connections that existed in the south, while younger splits within *P. polychopterus* in the south may be associated with more recent climatically-mediated gallery and dry forest connections and regressions between southern ecosystems (Arruda et al., 2018; Costa et al., 2018; Trujillo-Arias et al., 2018). Extensive wetlands connecting the western Amazonia and Paraná basins during the Miocene may have facilitated connectivity between river edge specialists and/or gallery forest birds until

the formation of the Fitzcarrald arch roughly 4 Ma (Espurt et al., 2010; Tagliacollo et al., 2015), which may have isolated populations of humid forest and edge taxa from east to west. However, previous work has demonstrated that speciation times across this region vary, rendering the effect of this arch on avian speciation uncertain (Claramunt, 2014). Even so, the relationship between Amazonian river edge specialists (e.g., *P. polychopterus nigriventris*) and southeastern South American taxa (e.g., *P. polychopterus spixii*) documented by a previous study is recapitulated by Tityrinae herein (Claramunt, 2014).

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Supplementary Tables:

Table S1: A list of all samples used in this study.

Tissue	Material	Institution	Tissue/Coll. #	Genus	Species	Subspecies	Coded as	Citation
Tissue	Tissue	Cantebury Museum		Acanthisitta	chloris		acachl	Jarvis et al., 2014
Tissue	Tissue	AMNH	DOT 12145	Elaenia	albiceps	chilensis	Elaenia albiceps chilensis DOT12145	This study
Tissue MPRG \$5056	Tissue	USNM	626253	Iodopleura	fusca		Iodop fusca S626253	NSF grant 1146248
Final	Tissue	LSUMNS	9597	Iodopleura	isabellae		Iodop isalae L9597	NSF grant 1146248
	Tissue	MPEG	55056	Lepidothrix	nattereri		Lep nat MPEG55056	This study
Figure CWBM 10442		FMNH	393859	Pachyramphus	aglaiae	albiventris	P a alb393859	Musher and Cracraft, 2018
Skin	FJ501700.1	AMNH	3688	Pachyramphus	aglaiae	latirostris	Pachyramphus agliael [RAG dataset]	
ISSUMNS	Tissue	UWBM	104042			sumichrasti		Musher and Cracraft, 2018
Tissue	Skin	AMNH	494147	Pachyramphus	albogriseus	albogriseus	P albogriseus 494147	This study
Tissue	Tissue	LSUMNS	33450	Pachyramphus		guayaquilensis	P a gua33450	Musher and Cracraft, 2018
Tissue	Tissue	LSUMNS	41620	Pachyramphus	albogriseus	ornatus	P a orn 41620	Musher and Cracraft, 2018
FINNH		LSUMNS	8114				Pac salv8114	
ISSUMNS	Tissue	FMNH	390007	Pachyramphus	castaneus	amazonus	P c ama390007	Musher and Cracraft, 2018
ISSUE ISLMNS	Tissue	FMNH	395729	Pachyramphus	castaneus	castaneus	P c cas395729	Musher and Cracraft, 2018
ISSUE ISUMNS 12174	Tissue	LSUMNS	46602	Pachyramphus	cinnamomeus	cinnamomeus	P c cin46602	Musher and Cracraft, 2018
ISSUE	Tissue	LSUMNS	8812	Pachyramphus	cinnamomeus	fulvidior	P c fulv8812	Musher and Cracraft, 2018
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Table S2: Age estimates for a set of clades based on our analyses in addition to a recently published phylogenomic study (Prum et al., 2015).

Dataset/citation	Crown Passeriformes (mya)	Stem Tyranni (mya)	Crown Tyrannides-Furnariides (mya)	Crown Tyrannides (mya)	Crown Tityridae (mya)
RAG (BEAST)	53.8 (49.65-57.82)	50.05 (46.47-53.60)	45.77 (37.88-47.24)	34.33 (28.86-40.59)	26.31 (19.67-33.64)
UCEs 95% (MCMCTree)	55.49 (50.74-58.10)	49.42 (46.38-53.33)	41.51 (37.97-46.16)	29.55 (27.97-32.34)	25.34 (21.35-27.82)
Prum et al., 2015	49 (41.5–55.5)	46.8 (38-53)	38.2 (27–46)	27 (17.5–36.5)	16 (5-25)

Musher et al. Supplementary Material for "Why is Amazonia a 'source' of biodiversity? Climate-mediated dispersal and synchronous speciation across the Andes in an avian group (Tityrinae)"

Table S3: Model fitting results for diversification models that explicitly incorporate carrying capacity (K) (Etienne et al., 2012). The best-fitting model is shown in bold.

Model	estimated parameters	logL	AICc	AICω	Speciation rate (λ)	Extinction rate (μ)	Carrying Capacity (K)
Birth-death constant rates	2	-108.942	222.151	0.339	0.340	0.119	NA
Linear dependence of speciation rate without extinction	2	-109.344	222.954	0.227	0.277	NA	1040588
Linear dependence of speciation rate with extinction	3	-108.942	224.430	0.108	0.340	0.118	189619
Exponential dependence of speciation rate with extinction	3	-109.013	224.572	0.101	0.390	0.149	Infinity
Linear dependence of extinction rate	3	-108.942	224.430	0.108	0.341	0.120	356419
Exponential dependence of extinction rate	3	-108.942	222.151	0.116	0.364	0.087	1742

Table S4: Results for generalized linear model test with averaged values for temperature and net-diversification across 50 intervals during the history of Tityrinae as estimated in TESS. The best model is M3, which models net-diversification rate as a linear function of interactive effects of time and temperature. Slopes are positive because we used values of negative time

Model	Adj. R- squared	slope	standard error	p-value (slope)	intercept	standard error	p-value (intercept)	AIC	ΑΙСω
M1: Time	0.943	0.013	0.000	0.000	0.210	0.005	0.000	-243.898	0.009
M2: Temp	0.677	-0.033	0.021	0.000	0.274	0.003	0.000	-157.074	0.000
		Time 0.009	0.002	0.000					
M3: Time and Temp	0.955	Temp 0.007	0.003	0.026	0.178	0.013	0.000	-253.386	0.991
		Time:temp 0.001	0.000	0.027					

Table S5: Model-fitting results for ten models: two constant net-diversification, four in which net-diversification varies with time, and four in which net-diversification varies with temperature. Models were conditioned on both stem and crown ages.

Model	Conditioning	Number of parameters	logL	AICc	ΑΙСω	speciation rate (λ)	exponential scaler for speciation rate (α)	extinction rate (μ)	exponential scaler for extinction rate (B)
Constant speciation	Stem	1	-116.41	234.90	0.06	0.2501			
Constant speciation and extinction	Stem	2	-114.49	233.24	0.14	0.3865		0.2423	
Speciation varies exponentially with time	Stem	2	-113.85	231.97	0.26	0.3432	-0.0706		
Speciation varies exponentially with time and constant extinction	Stem	3	-113.85	234.24	0.08	0.3432	-0.0706	0.0000	
Extinction varies exponentially with time and constant speciation	Stem	3	-114.21	234.97	0.06	0.3627		0.1633	0.0345
Speciation and extinction vary exponentially with time	Stem	4	-113.85	236.63	0.02	0.3432	-0.0706	0.0000	0.0208
Speciation varies exponentially with temperature	Stem	2	-113.96	232.18	0.23	0.3848	-0.0922		
Speciation varies exponentially with temperature and constant extinction	Stem	3	-113.96	234.46	0.07	0.3846	-0.0922	0.0000	
Extinction varies exponentially with temperature and constant speciation	Stem	3	-114.21	234.97	0.06	0.3627		0.1592	0.0306
Speciation and extinction vary exponentially with temperature	Stem	4	-113.96	236.85	0.02	0.3847	-0.0907	0.0000	0.0250
Constant speciation	crown	1	-115.01	232.11	0.01	0.2452			
Constant speciation and extinction	crown	2	-112.49	229.25	0.06	0.4030		0.2846	
Speciation varies exponentially with time	crown	2	-110.91	226.09	0.28	0.3683	-0.0962		
Speciation varies exponentially with time and constant extinction	crown	3	-110.91	228.37	0.09	0.3684	-0.0962	0.0000	
Extinction varies exponentially with time and constant speciation	crown	3	-110.60	227.74	0.12	0.3330		0.0679	0.1361
Speciation and extinction vary exponentially with time	crown	4	-109.54	229.58	0.05	0.3297	-0.0623	0.0003	0.4437
Speciation varies exponentially with temperature	crown	2	-111.43	227.12	0.17	0.4089	-0.1125		
Speciation varies exponentially with temperature and constant extinction	crown	3	-111.36	229.26	0.06	0.4108	-0.0809	0.0666	
Extinction varies exponentially with temperature and constant speciation	crown	3	-110.60	227.74	0.12	0.3330		0.0614	0.1206
Speciation and extinction vary exponentially with temperature	crown	4	110.54	230.01	0.04	0.3505	-0.0594	0.0006	0.3596

Supplementary Figures:

Figure S1: Regressions for clock-likeness (measured by likelihood ratio; LR) as a function of (A) the proportion of informative sites, (B) the number of informative sites, and (C) the $\log(\lambda)$ of per locus, where λ is the optimal smoothing parameter for the penalized likelihood analysis in ape (Paradis et al., 2004). Low values of λ are indicative of higher heterogeneity in branch rates, and thus lower clock-likeness. Gray points represent outlier loci. Gray lines represent each linear model before filtering outliers, and red lines represent the linear model after filtering outliers.

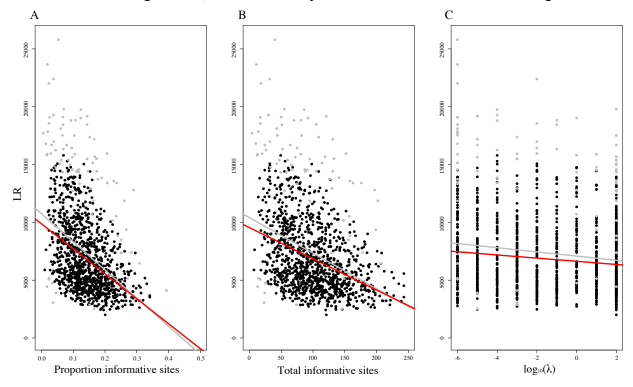


Figure S2: A random subsample of 25 gene trees for loci that were considered outliers and removed from downstream analysis (left), and the 15 most clocklike markers after filtering, which were used for molecular divergence dating analysis (right). The values for number of parsimony informative sites (IS) and the likelihood ratio (LR) are above each gene tree. Values in red represent low LR gene trees that were none-the-less identified as outliers.

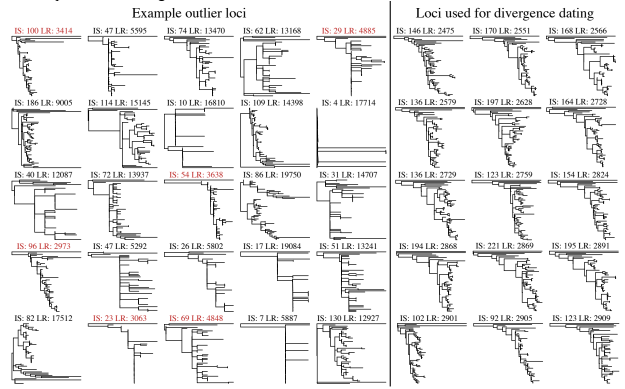


Figure S3: Left: speciation (purple) and extinction (red) rates through time from TESS. Right: The posterior probabilities of speciation (purple bars) and extinction (red bars) rate shifts through time.

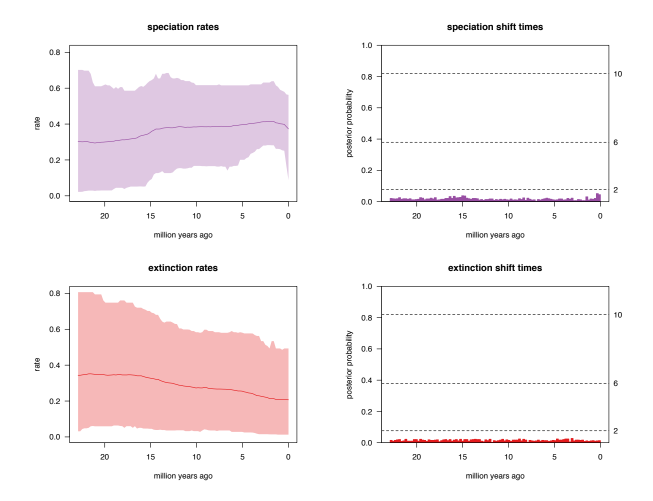


Figure S4: BAMM best shift configuration results for (left to right) speciation rates, extinction rates, and net-diversification rates.

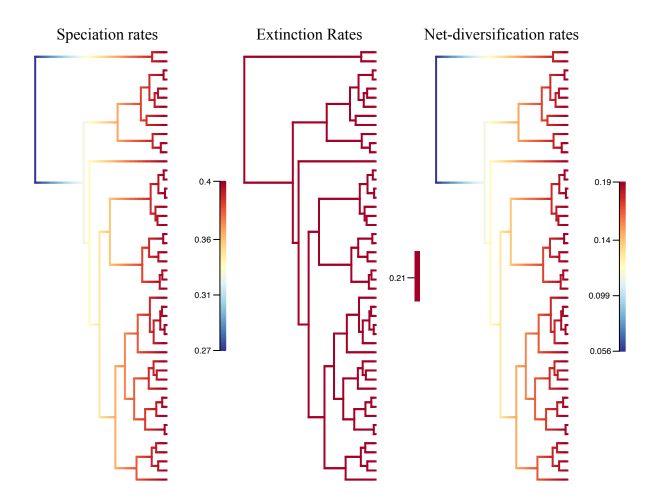


Figure S5: The relationship between net-diversification estimate in TESS and temperature (adj. R-sq. = 0.66813) for 50 intervals between the root and tips of the dated Tityrinae phylogeny.

Relationship between palaeotemperature and net-diversification rates through time

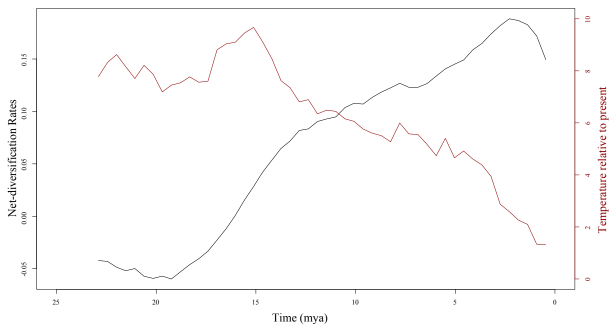


Figure S6: Results from best-fitting RPANDA models conditioned on the crown showing how speciation rate varies as a function of time and palaeoclimate (mean global temperature relative to present). Model-fitted net-diversification estimates for pure birth with exponential dependence on time (top row), and pure birth with exponential dependence on temperature (bottom row).

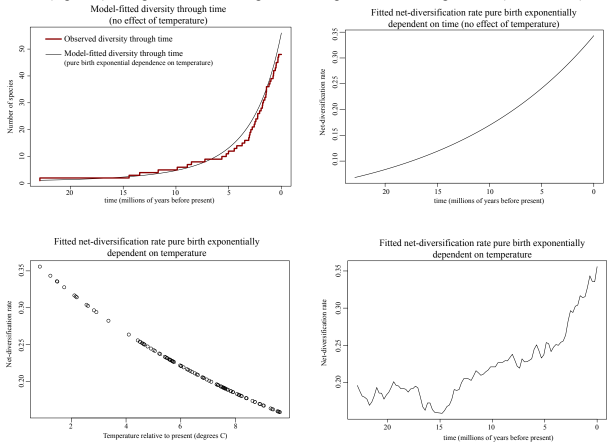


Figure S7: Results from RAxML on concatenated UCEs.

