Tempo and mode of morphological evolution are decoupled from latitude in birds

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Abstract

The latitudinal diversity gradient is one of the most striking patterns in nature, yet its implications for morphological evolution are poorly understood. In particular, it has been proposed that an increased intensity of species interactions in tropical biota may either promote or constrain trait evolution, but which of these outcomes predominate remains uncertain. Here, we develop tools for fitting phylogenetic models of phenotypic evolution in which the impact of species interactions—namely, competition—can vary across lineages. Deploying these models on a global avian trait dataset to explore differences in trait divergence between tropical and temperate lineages, we find that the effect of latitude on the mode and tempo of morphological evolution is weak and clade- or trait dependent. Our results indicate that species interactions do not disproportionately impact morphological evolution in tropical bird families and question the validity of previously reported patterns of slower trait evolution in the tropics.

Introduction

In many groups of organisms, species richness increases toward lower latitudes—a pattern known as the latitudinal diversity gradient—inspiring generations of biologists to search for the causes and consequences of this gradient [1]. One hypothesis posits that species interactions are stronger in the tropics, and, therefore, play a more important role in many processes (e.g., diversification) in tropical lineages [2–6] (but see [7]). Previous tests of this “biotic interactions hypothesis” have generally focused on latitudinal gradients in the strength of ecological interactions between predator and prey, herbivore and plant, or pathogen and host [8–11]. Latitudinal gradients in the strength of competition between members of the same trophic level have been less explored, although they have been highlighted as one of the most important research directions for testing the biotic interaction hypothesis [5]. Competition among closely related species, such as those from the same taxonomic family, is often assumed to be
strong since their ecological and phenotypic similarity increases the likelihood of competition for access to resources or space \([12–16]\). Such interactions can influence selection on traits that mediate access to resources, influencing trait evolution either by promoting divergence between lineages via character displacement \([17,18]\) or, alternatively, by imposing constraints on geographical range overlap and ecological opportunity, reducing trait diversification as niches fill \([19–21]\).

Whether competition predominantly drives or constrains divergence, the impacts on trait evolution should leave a detectable phylogenetic signature \([22–25]\). In addition, this signature should be most prevalent in the tropics, where each lineage interacts with far larger numbers of potential competitors. As such, the biotic interactions hypothesis predicts differences between tropical and temperate taxa in the pace of evolution (the “tempo,” in the parlance of comparative studies) and/or the processes that drive trait diversification (the “mode”). In comparison with the wealth of studies that have investigated latitudinal gradients in rates of species diversification \([26–30]\), relatively few have tested for latitudinal gradients in the dynamics of phenotypic evolution and have mainly focused on tempo rather than mode. Their results so far suggest a potentially complex relationship between trait diversification and latitude. On the one hand, some studies have found greater divergence between sympatric sister taxa in body mass \([31]\) and in plumage coloration \([32]\) in the tropics, supporting the hypothesis that increased competition at lower latitudes drives character displacement \([5]\). On the other hand, some studies have found that species attain secondary sympatry after speciation more slowly in tropical regions \([33]\) or that evolutionary rates are lower in the tropics for climatic niches \([34]\), body size \([34,35]\), or social signaling traits \([34,36–39]\), implying that competition may limit ecological opportunity, and, therefore, constrain trait divergence in tropical regions.

Disentangling these opposing effects is challenging because previous macroecological studies have generally been restricted to either relatively few traits or limited samples of species. In addition, previous studies have been impeded by the lack of suitable methods for detecting the impact of species interactions on trait evolution \([40–42]\), although recent progress has been made in developing such methods for use in standard comparative analyses \([20,22,24,43]\). By incorporating species interactions directly into phylogenetic models of trait evolution, these developments overcome some of the issues faced by phylogenetic and trait approaches for studying community assembly that rely on overly simplistic comparisons to randomly assembled communities \([43–45]\). However, these developments have not yet been deployed in the context of latitudinal sampling, and, thus, the key prediction of a latitudinal gradient in trait diversification has yet to be tested.

Here, we begin by expanding existing phylogenetic models of phenotypic evolution, including models that incorporate competition between species—namely, diversity-dependent (DD) models \([19,20]\) and the matching competition (MC) model \([22,43]\)—such that the impact of interactions between co-occurring lineages on trait evolution can be estimated separately in lineages belonging to different, predefined competitive regimes (e.g., tropical and temperate). We note that we use “competition” to encompass all processes (both direct and indirect), whereby trait evolution is impacted by co-occurring lineages. The models we develop are designed to account for known intraspecific variability and unknown, nuisance measurement error, both of which can strongly bias model support and parameter estimates \([46]\). In particular, it has been suggested that intraspecific variability is lower in the tropics \([47]\), which could inflate estimates of evolutionary rates in the temperate biome. Next, we conduct a comprehensive test of the biotic interactions hypothesis using these new phylogenetic tools to model the effect of interspecific competition on the tempo and mode of morphological evolution based on 7 morphological characters describing variation in body size, bill size and shape, and locomotory strategies sampled from approximately 9,400 species representing more than 100 avian
families worldwide. These morphological characters have been demonstrated to predict diet and foraging behavior in birds [48], indicating that they are well suited as proxies for analyzing the dynamics of ecological divergence.

Results

Latitudinal variation in mode of phenotypic evolution

We tested whether modes of phenotypic evolution varied with latitude using 2 types of models. First, we tested whether support for various “single-regime” models that estimate a single set of parameters on the entire avian phylogeny [26] varied according to a clade-level index of tropicality. Second, we developed and used “2-regime” models with distinct sets of parameters for tropical and temperate species and tested whether these latitudinal models were better supported than single-regime models.

Across single-regime fits, we found no evidence for a latitudinal trend in the overall support for any model of phenotypic evolution (Fig 1A–1F and S4 Table), with one exception: There was an increase in model support for the MC model in tropical lineages for the locomotion phylogenetic principal component (pPC) 3 (Fig 1F and S4 Table). Similarly, there was no evidence that the overall support for models incorporating competition (i.e., MC or DD models) is higher in tropical clades (Fig 1G and S4 Table). Models with latitude (i.e., 2-regime models) were not consistently better supported than models without latitude, for any model or trait (S5 Table). Indeed, single-regime models were the best-fit models across 86% of individual clade-by-trait fits (S7 Fig).

Latitudinal variation in the effect of interactions on phenotypic evolution

We found no evidence for a latitudinal trend in the slope estimated from single-regime DD models (Fig 2C and 2D and S6 Table). However, the strength of repulsion estimated from single-regime MC models increased in more tropical families for locomotion pPC3 (Fig 2B and S6 Table). Parameter estimates from 2-regime models with competition (i.e., MC or DD models) do not support a stronger effect of biotic interactions on phenotypic evolution in the tropics (Fig 3B–3D)—in most traits, there is no consistent difference between estimates of the impact of competition on tropical and temperate lineages, and in one case (bill pPC2), there is evidence that competition impacts temperate lineages to a larger degree than tropical lineages (Fig 3B–3D and S7 Table). In all cases, there was substantial variation in the fits, and the overall magnitude of differences between tropical and temperate regions was rather small (Fig 3B–3D).

Impact of assuming continental scale sympatry

Phylogenetic models of competitively driven trait evolution rely on reconstructions of ancestral ranges to delimit the pool of potential species interaction at each point in the evolutionary history of a clade. Given the scale of our analyses and the computational limits of existing models of ancestral range estimation, we assumed that species occurring on the same continent were able to interact with one another. On average, species in our analyses are sympatric with 50% of clade members at the continental level, although there are differences across continents (mean range: 34% to 74%; S5 Fig and S9 and S10 Tables). Notably, we also found that temperate species are more likely to coexist in sympatry with family members than tropical species (S11 Table). To determine the impact of assuming continental scale sympatry, we investigated whether we would detect a latitudinal difference in the effect of competition on phenotypic evolution if it existed, even if competition occurs among only truly sympatric
species rather than among all species occurring on the same continent. Simulations examining the impact of the continental scale sympatry assumption on the statistical power of 2-regime
Fig 2. Parameter estimates from single-regime models reveal varying impacts of latitude. There is no impact of latitude on the effect of competition on trait evolution as measured by the slope of (a) DDexp models or (b) DDlin models. (c) The effect of competition on trait evolution as measured by the repulsion parameter ("S") from the MC models increases with the index of tropicality (the proportion of species in the clade with exclusively tropical breeding distributions) for locomotion pPC3 but not for other traits. (d) There is no relationship between the proportion of taxa in a clade that breed in the tropics and the estimated rate of trait evolution from BM models. Solid lines represent statistically significant relationships (S6 and S13 Tables). For (a–c), each point represents the mean across clade-by-trait fits to stochastic maps of biogeography (for all families with at least 50 species), and for (d), each point represents the MLE for each clade-by-trait fit (see S2 and S3 Datas). BM, Brownian motion; DD, diversity-dependent; DDexp, exponential diversity-dependent; DDlin, linear diversity-dependent; MC, matching competition; MLE, maximum likelihood estimate; pPC, phylogenetic principal component.

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MC models demonstrate that, even for relatively small clades, large but biologically plausible latitudinal differences in the effect of competition should be detectable, even when sympatry is overestimated (S8 Fig). Nevertheless, there is evidence that this assumption can impact the power to detect subtle differences between regions and for smaller trees, the estimated direction of the difference (S8 Fig). However, restricting our empirical analyses to large clades ($N \geq$...
100), we still find no support for a consistently stronger impact of competition on phenotypic evolution in tropical lineages (S8 Table).

**Latitudinal variation in tempo of phenotypic evolution**

Evolutionary rates estimated from single-rate models did not vary according to clade-level index of tropicality (Figs 2 and S9 and S13 Table). Similarly, estimates of rates from latitudinal models were neither consistently lower nor higher in tropical regions (Figs 3D and S10 Fig and S14 Table). We did find lower rates of locomotion pPC3 (Figs 3D and S10 and S14 Table) and bill pPC2 evolution in tropical lineages (S10 Fig and S14 Table), but the difference between regions was small, and the overall strength of this relationship was weak. Observational error contributed to these patterns: We found a significant negative correlation between observational error and the clade-level index of tropicality for body mass (S11 Fig and S15 Table), and we also found that there was a correlation between rates of body mass and locomotion pPC3 evolution in standard single-regime BM models excluding error (S12 Fig and S16 Table) and that the magnitude of the difference between tropical and temperate rates of trait evolution was higher in analyses of 2-regime fits excluding error (S12 Fig and S17 Table).

**Predictors of support for models with an effect of competition on phenotypic evolution**

We found no evidence that territoriality or diet specialization are useful predictors of support for models that incorporate the impact of co-occurring species on phenotypic evolution (S18 Table). We did, however, find that the maximum proportion of species co-occurring on a continent (i.e., the maximum number of extant lineages on a single continent divided by the total clade size) had a pronounced impact on model selection—clades with a high proportion of lineages occurring on the same continent were more likely to be best fit by the MC model, whereas clades with a low proportion of co-occurring lineages were more likely to be best fit by the exponential DD model (S13 and S14 Figs and S18 Table). In addition, we found that the MC model was less likely to be favored in clades with many members living in single-strata habitats (S18 Table).

**Discussion**

Contrary to what would be expected if the effect of competition on phenotypic evolution was stronger in the tropics, we did not find a consistent latitudinal gradient in the dynamics of phenotypic evolution across the entire avian radiation. Using novel methods for examining macroevolutionary signatures of the effect of competition on phenotypic evolution, we show that patterns of trait evolution across many clades are consistent with competition between clade members acting as an important driver of trait evolution. Nevertheless, we found no evidence that such competition has impacted the dynamics of trait diversification more in the tropics than in temperate regions. This lack of consistent latitudinal effect applied both to the support for specific models of phenotypic evolution and the parameters of these models. Our results contrast with several previous studies that have found a consistent signature of faster rates in the temperate biome [34,36–39,49].

The apparent absence of latitudinal patterns in support of phenotypic models with competition and estimates of competition strength did not arise from overall weak support for competition models, confirming previous findings that competition does leave a detectable signal in comparative, neontological datasets [22–25,50,51]. Indeed, models incorporating species interactions were the best-fit models in 25% of clade-by-trait combinations for single-regime fits. In sunbirds (Nectariniidae), for instance, the MC model was the best-fit model for body mass...
and 2 pPC axes describing variation in bill shape, suggesting that competition has driven trait divergence in this diverse clade. In owls (Strigidae), the exponential DD model was the best-fit model for body mass and several pPC axes describing bill shape and locomotory traits, suggesting that the rate of evolution in owls responds to changing ecological opportunity. The finding that interactions with co-occurring species commonly leave a signature on extant phenotypes in birds is echoed by a recent study showing that traits in a similar proportion of clades are best fit by competition models [50].

For both single-regime models and 2-regime models, we detected no systematic effect of latitude on the impact of competition on trait diversification. One possible explanation for this is that our approach was highly conservative since we assumed that species occurring on the same continent are likely to interact with one another, whereas they may be allopatric (with nonoverlapping geographical ranges) or exhibit low levels of syntopy within areas of sympatry [52]. However, previous work [23] and simulations exploring the impacts of assuming competition between potentially allopatric lineages suggest that the MC model is robust to some mis-specification of geographic overlap (e.g., allopatric species scored as sympatric). This robustness is likely explained by both the imprint of competition on ancestral, coexisting lineages and a formulation of competition where divergence occurs respective to mean phenotypic values across interacting species (the mean across all species within each continent may be a relatively good proxy for the mean across sympatric species). Nevertheless, the possibility remains that, if differences between regions in the impact of competition are sufficiently small, the 2-regime models may not have detected them. In aggregate, however, our results consistently point to a conspicuous absence of a latitudinal gradient in the effect of competition on trait diversification.

One plausible explanation for discrepancies between our results and other studies that examine gradients in the tempo of morphological trait evolution is that our study accounted for observational error. Indeed, we found that overall observational error for body mass increased with latitude, and, when we intentionally ignored observational error, Brownian motion (BM) models were more likely to pick up faster rates of trait evolution at high latitudes. This result makes sense in the light of previously reported higher trait variance for temperate taxa [47] and a positive correlation between such variance and rate estimates [53]. Our analyses demonstrate that accounting for observational error when testing for latitudinal trends in evolutionary rates is crucial and also suggest that previous analyses overlooking error may have detected spurious latitudinal gradients in trait evolution.

Another potential explanation for the discrepancy between this and previous studies is that many previous studies examined gradients in rapidly evolving plumage and song traits, which may vary latitudinally if sexual or social selection is more pronounced in temperate regions [54]. In contrast, divergence in ecological traits is likely more constrained, as they tend to evolve relatively slowly [55,56].

A third explanation for the discrepancy is that many previous studies used sister taxa approaches to estimate gradients in trait evolution [34,36,37,49]. Yet, avian sister taxa are younger in temperate regions [33,49], and how these age differences influence rate estimates if trait evolution has proceeded in a non-Brownian fashion is not clear. Analyses on sister taxa of different ages can recover different rates even though these rates are not representative of any process unique to any particular region. For example, given that rates of trait evolution have accelerated toward the present [57], we may expect sister taxa to recover a signature of faster rates in temperate regions (where sister taxa are younger), even if there are no clade-wide latitudinal differences in the overall tempo and mode of evolution.

Within the competition models, the MC model was more likely to be chosen as the best-fit model than DD models, which is consistent with the notion that competition promotes
divergence (e.g., via character displacement [17,18]) more often than it constrains divergence (e.g., via niche saturation [19]) at relatively shallow taxonomic scales [15,42,58]. Nevertheless, the possibility remains that other processes might generate patterns that are picked up by the MC and DD models. For instance, although the models we fit are designed to estimate the dynamics of trait evolution, competition can also generate patterns of divergence via its impacts on range dynamics (i.e., ecological sorting) when secondary sympatry is delayed by competitive interactions [21,59,60]. Therefore, although recent evidence suggests that the effects of competitive exclusion on community assembly is distinguishable from the action of character displacement in comparative datasets [25], the possibility remains that the MC model may detect a signal of ecological sorting of morphologically distinct lineages [21,61]—a process that is also fundamentally governed by competition—in addition to or instead of evolutionary divergence [25]. Further development of phylogenetic models that incorporate biotic interactions and simulation studies may help to clarify the processes that generate trait distributions which MC and DD models fit well.

In our analyses, we focused within clades, where we would expect competition to be strongest, owing to the phenotypic and ecological similarity of recently diverged taxa [16]. Nevertheless, in doing so, we excluded other competitors (e.g., nonfamily members with similar diets) that impose constraints on niche divergence. Such competitors have been shown to impact rates of trait evolution across clades of birds [53]. Future research could extend our approach by examining the impact of interactions between competitors from a wider diversity of clades.

We found evidence that support for the MC model was greater in clades with a higher proportion of lineages occurring on the same continent, suggesting that trait divergence may make coexistence possible [15,18]. The exponential DD model, on the other hand, was more likely to be the best-fit model in clades with relatively low levels of continental overlap, which may indicate that in these clades, niche saturation negatively impacts coexistence [62,63]. In addition, we found that model fits on clades with a high proportion of species living in single-strata habitats were less likely to favor the MC model, suggesting that opportunity for divergence may be limited in such habitats [64]. These relationships between ecological opportunity, trait evolution, and coexistence highlight the need for models that can jointly estimate the effects of diversification, range dynamics, and trait evolution [25,58]. Such models may identify an impact of competition on processes other than trait evolution, such as competitive exclusion, which may themselves vary latitudinally [21,33].

By including a suite of traits that capture functional variation in niches [48], we were able to identify patterns that would have been highly biased, or that we would have missed, by focusing on one specific trait, in particular body mass. Model support was distributed evenly across different traits, suggesting that the impact of competition varies both across clades and across different functionalities. For instance, while 31% (42/135) of clades exhibit some signature of competition acting on body size evolution in single-regime fits, 68% (92/135) of them exhibit some signature of competition acting on at least 1 of the 7 functional traits (body size, bill pPC axes, and locomotion pPC axes). These results further strengthen the notion that multiple trait axes are necessary to robustly test hypotheses about ecological variation [48,50,65].

We have extended various phylogenetic models of phenotypic evolution, including models with competition, to allow model parameters to vary across lineages (see also [51]) and to account for biogeography and sources of observational error. We then applied them to the case of latitudinal gradients, but they could be used to study other types of geographic (e.g., elevation), ecological (e.g., habitat and diet), behavioral (e.g., migratory strategy), or morphological (e.g., body size) gradients. Studies of gradients in evolutionary rates are often performed using sister taxa analyses, assuming BM or OU processes [66]. These analyses are useful
because they enable quantitative estimates of the impact of continuous gradients on rate parameters. However, by limiting analyses to sister taxa datasets (and therefore ignoring interactions with other coexisting lineages), they are unable to reliably detect signatures of species interactions [67] and so cannot be used to study competition. In addition, these approaches are not well suited to differentiating between different evolutionary modes. Applying process-based models of phenotypic evolution that incorporate interspecific competition and biogeography allow for such tests of evolutionary hypotheses about the mode of trait evolution across entire clades.

Focusing on the effect of competition between closely related species on phenotypic evolution, we did not find support for the biotic interactions hypothesis. Biotic interactions are multifarious; individuals face selective pressures arising from competition, but also from other types of interactions such as predator–prey and host–parasite interactions. Perhaps as a result of this complexity, pinning down clear empirical relationships between latitude and biotic interactions has yielded a complex and often inconsistent set of results [7], with empirical evidence ranging from stronger interactions in the tropics [8,10] to stronger interactions in temperate regions [9]. A challenge for future research on the biotic interactions hypothesis is, therefore, to more precisely identify the mechanisms that lead to latitudinal gradients in interactions, and, consequently, better predict the kinds of interactions that may shape latitudinal gradients in diversification.

**Materials and methods**

**Two-regime phylogenetic models of phenotypic evolution**

One approach to analyze gradients in phenotypic evolution is to fit phylogenetic models of phenotypic evolution that allow model parameters (e.g., evolutionary rates) to vary across the phylogeny; such models are already available for the simplest models of trait evolution such as BM and Ornstein–Uhlenbeck (OU) models [68,69]. To explore the effects of species interactions, we developed further extensions to early burst (EB), DD, and MC models, allowing parameters to be estimated separately in 2 mutually exclusive groups of lineages in a clade. Generalizing these new models to estimate parameters on more than 2 groups, or on non-mutually exclusive groups, is straightforward.

We began by developing a 2-regime version of the EB model in which rates of trait evolution decline according to an exponential function of time passed since the root of the tree [70]. We used this model here to ensure that the DD models, which incorporate changes in the number of reconstructed lineages through time, are not erroneously favored because they accommodate an overall pattern of declining rates through time. To estimate rates of decline separately for mutually exclusive groups, we formulated a 2-regime EB model with 4 parameters (Table 1): $z_0$ (the state at the root), $\sigma_0^2$ (the evolutionary rate parameter at the root of the tree), $r_A$ (controlling the time dependence on the rate of trait evolution in regime "A"), and $r_B$ (time dependence in regime "B"). This model can be written as follows:

$$dX_{ij}^{(0)}(t) = \begin{cases} \frac{1}{\sigma_0^2} \sigma_0^2 e^{r_A t} dW_i, & \text{if } j \text{ is in } A \text{ at time } t \\ \frac{1}{\sigma_0^2} \sigma_0^2 e^{r_B t} dW_i, & \text{if } j \text{ is in } B \text{ at time } t \end{cases}$$

(Eq 1)

where $X_{ij}^{(0)}$ is the trait value of lineage $j$ at time $t$, and $dW_i$ represents the BM process (S1 Fig). This model is the 2-regime equivalent of the EB model, where $\sigma^2(t) = \sigma_0^2 e^{rt}$; the (1/2) factor in Eq 1 comes from taking the square root of the rate.
DD models represent a process where rates of trait evolution respond to changes in ecological opportunity that result from the emergence of related lineages [19,20]. When the slope of these models is negative, this is interpreted as a niche filling process where rates of trait evolution slow down with the accumulation of lineages. We considered 2 versions of DD models, with either exponential (DD_{exp}) or linear (DD_{lin}) dependencies of rates to the number of extant lineages. The 2-regime model has 4 free parameters (Table 1): $z_0$ (the state at the root), $\sigma^2$ (the evolutionary rate parameter), $r_A$ (the dependence of the rate of trait evolution on lineage diversity in regime “A”), and $r_B$ (diversity dependence in regime “B”). For the exponential case, this model can be written as follows:

$$dX^{(i)}_t = \begin{cases} \frac{1}{\sigma_i} \exp r_A n^{(A)}_t dW_t, & \text{if } j \text{ is in } A \text{ at time } t \\ \frac{1}{\sigma_i} \exp r_B n^{(B)}_t dW_t, & \text{if } j \text{ is in } B \text{ at time } t \end{cases},$$

(Eq 2)

for the exponential case, where $n^{(A)}_t$ and $n^{(B)}_t$ are the number of lineages in regime $A$ and $B$ at time $t$. This model is the 2-regime equivalent of the DD_{exp} model, where $\sigma^2(t) = \sigma_0^2 e^{n^{(t)}_j}$; the (1/2) factor in Eq 2 comes from taking the square root of the rate. For the linear case, this can be written as follows:

$$dX^{(i)}_t = \begin{cases} \sqrt{\sigma_0^2 + b_A n^{(A)}_t} dW_t, & \text{if } j \text{ is in } A \text{ at time } t \\ \sqrt{\sigma_0^2 + b_B n^{(B)}_t} dW_t, & \text{if } j \text{ is in } B \text{ at time } t \end{cases},$$

(Eq 3)

This model is the 2-regime equivalent of the DD_{lin} model, where $\sigma^2(t) = \sigma_0^2 + b n_t$ and $b$ denotes the slope in the linear model. Standard DD models ignore whether lineages coexist.

Table 1. Parameters of models used in analyses.

| Model       | $k$ | $\sigma^2$ | $z_0$   | Other               |
|-------------|-----|------------|---------|---------------------|
| BM_single   | 2   | $\sigma^2$ | $z_0$   | —                   |
| BM_two      | 3   | $\sigma^2$ |         | $r\text{ trop}$     |
| OU_single   | 3   | $\sigma^2$ | $z_0$   | $r\text{ trop}$     |
| OU_two      | 4   | $\sigma^2$ | $z_0$   | $r\text{ trop}$ $r\text{ temp}$ |
| EB_single   | 3   | $\sigma^2$ | $z_0$   | $r$ (slope)         |
| EB_two      | 4   | $\sigma^2$ | $z_0$   | $r\text{ trop}$     |
| DD_{exp}     | 3 | $\sigma^2$ | $z_0$   | $r$ (slope)         |
| DD_{lin}    | 4   | $\sigma^2$ | $z_0$   | $b$ (slope)         |
| MC_single   | 3   | $\sigma^2$ | $z_0$   | $S$                 |
| MC_two      | 4   | $\sigma^2$ | $z_0$   | $S\text{ trop}$ $S\text{ temp}$ |

The subscripts “trop” and “temp” in the 2-regime versions of each model refer to parameters estimated separately for lineages with exclusively tropical breeding ranges and lineages with breeding ranges that include the temperate region. $k$ indicates the number of free parameters estimated in each model, $\sigma^2$ indicates the rate parameter describing the tempo of trait evolution, $z_0$ indicates the trait value at the root of the clade, and $\alpha$ describes the strength of the pull toward a stable optimum in the OU model. For descriptions of other parameters, see the main text. BM, Brownian motion; DD_{exp}, exponential diversity-dependent; DD_{lin}, linear diversity-dependent; EB, early burst; MC, matching competition; OU, Ornstein–Uhlenbeck.

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yet only those lineages likely to encounter one another in sympatry are able to compete with one another. Thus, we extended our model to incorporate ancestral biogeographic reconstructions to identify which species interactions are possible at any given point in time (i.e., which species co-occur [23]). With biogeography, these become

\[
d X_{i}^{(j)} = \begin{cases} 
\frac{1}{\sigma \sqrt{2} \tau A} \sum_{i=1}^{n^{(A)}} A_{i,j}^{(A)} dW_{i} & \text{if } j \text{ is in } A \text{ at time } t \\
\frac{1}{\tau B} \sum_{i=1}^{n^{(B)}} A_{i,j}^{(B)} dW_{i} & \text{if } j \text{ is in } B \text{ at time } t 
\end{cases}
\]  

(Eq 4)

for the exponential case and

\[
d X_{i}^{(j)} = \begin{cases} 
\sqrt{\sigma_{0}^{2} + b_{A} \sum_{i=1}^{n^{(A)}} A_{i,j}^{(A)} dW_{i}} & \text{if } j \text{ is in } A \text{ at time } t \\
\sqrt{\sigma_{0}^{2} + b_{B} \sum_{i=1}^{n^{(B)}} A_{i,j}^{(B)} dW_{i}} & \text{if } j \text{ is in } B \text{ at time } t 
\end{cases}
\]  

(Eq 5)

for the linear case, where \( A \) is a matrix denoting biogeographical overlap, such that \( A_{j,l} = 1 \) if lineages \( j \) and \( l \) coexist in sympatry at time \( t \), and 0 otherwise (S1 Fig).

The MC model is a model of competitive divergence [22,43], wherein sympatric lineages are repelled away from one another in trait space. We formulated the 2-regime MC model, which has 4 parameters (Table 1): \( z_{0} \) (the state at the root), \( \sigma^{2} \) (the evolutionary rate parameter), \( S_{A} \) (the strength of repulsion in regime “A”), and \( S_{B} \) (the strength of repulsion in regime “B”). This model can be written as follows:

\[
d X_{i}^{(j)} = \begin{cases} 
S_{A} \left( \frac{\sum_{i=1}^{n^{(A)}} X_{i}^{(j)}}{n^{(A)}} - X_{i}^{(j)} \right) + \sigma dW_{i} & \text{if } j \text{ is in } A \text{ at time } t \\
S_{B} \left( \frac{\sum_{i=1}^{n^{(B)}} X_{i}^{(j)}}{n^{(B)}} - X_{i}^{(j)} \right) + \sigma dW_{i} & \text{if } j \text{ is in } B \text{ at time } t 
\end{cases}
\]  

(Eq 6)

Incorporating biogeography, this becomes

\[
d X_{i}^{(j)} = \begin{cases} 
S_{A} \left( \frac{\sum_{i=1}^{n^{(A)}} A_{i,j} X_{i}^{(j)}}{\sum_{i=1}^{n^{(A)}} A_{i,j}} - X_{i}^{(j)} \right) + \sigma dW_{i} & \text{if } j \text{ is in } A \text{ at time } t \\
S_{B} \left( \frac{\sum_{i=1}^{n^{(B)}} A_{i,j} X_{i}^{(j)}}{\sum_{i=1}^{n^{(B)}} A_{i,j}} - X_{i}^{(j)} \right) + \sigma dW_{i} & \text{if } j \text{ is in } B \text{ at time } t 
\end{cases}
\]  

(Eq 7)

We developed inference tools for fitting the 2-regime MC and DD models to comparative trait data, following the numerical integration approach used previously [43,56]. For the EB model, we developed a branch transformation approach similar to the one used in mvMORPH [71]. In all model fits, we incorporated the possibility to account for deviations between measured and modeled mean trait values for each species [72–74] (see S1 Text for details). These deviations are of 2 types: the “known” deviation associated with estimating species means from a finite sample and the “unknown” deviation linked to intraspecific variability unrelated to the trait model (e.g., instrument errors and phenotypic plasticity). We follow the common practice of lumping these 2 sources of deviations (often called “measurement error”) and referring to
them as “observational error.” A simulation study demonstrated the reliability of estimates using these tools (S1 Text and S7 Data). Functions to simulate and fit these phenotypic models are available in the R package RPANDA [86].

**Phylogeny and trait data**

We obtained phylogenies of all available species from birdtree.org [26] and created a maximum clade credibility tree in TreeAnnotator [75] based on 1,000 samples from the posterior distribution (S13 and S14 Datas). Since the MC and DD models require highly sampled clades [43], we used the complete phylogeny including species placed based on taxonomic data [26] and the backbone provided by Hackett and colleagues [76]. We then extracted trees for each terrestrial (i.e., non-pelagic) family with at least 10 members (n = 108). As island species are generally not sympatric with many other members of their families (median latitudinal range of insular taxa = 1.28˚ and non-insular taxa = 15.27˚), we further restricted our analyses to continental taxa, excluding island endemics and species with ranges that are remote from continental land masses. We gathered data on the contemporary ranges of each species from shapefiles [77].

Mass data were compiled from EltonTraits [78] (n = 9,442). In addition, we used a global dataset based on measurements of live birds and museum specimens [48] to compile 6 linear morphological measurements: bill length (culmen length), width, and depth (n = 9,388, mean = 4.5 individuals per species), as well as wing, tarsus, and tail length (n = 9,393, mean = 5.0 individuals per species). These linear measurements were transformed into pPC axes describing functionally relevant variation in bill shape and locomotory strategies (S1 Text and S2 and S3 Tables and S1 Data).

**Biogeographic data and reconstruction**

Phylogenetic models that account for species interactions require identifying lineages that are likely to encounter one another [43]. To discretize the contemporary ranges of each species, we classified them as being present or absent in 11 different global regions [79]: Western Palaearctic, Eastern Palaearctic, Western Nearctic, Eastern Nearctic, Africa, Madagascar, South America, Central America, India, Southeast Asia, and Papua New Guinea/Australia/New Zealand. To assign each species to the global region(s) they occupied, we used several approaches. As a first pass, we used the maximum and minimum longitude and latitude for species’ (non-breeding) ranges. When the rectangle formed by these values fell entirely within the limits of a given global region, we assigned that region as the range for the focal species. Next, for species that did not fall entirely into one region, we compiled observation data from eBird.org [80] to identify all of the regions that a species occupies using country-level observations. Finally, for species whose ranges could not be resolved automatically using these techniques, we manually inspected the ranges.

We incorporated estimates of the presence/absence of each lineage in each range through time using ancestral range estimation under the dispersal–extinction–cladogenesis (DEC) model of range evolution [81]. We fit DEC models to range data and phylogenies for each family with the R package BioGeoBEARS [81,82]. Since the continents have changed position over the course of the time period of family appearance (clade age range = 12.84 to 71.88 Mya), we ran a stratified analysis with adjacency and dispersal matrices defined for every 10 my time slice [79]. Using the maximum likelihood (ML) parameter estimates for the DEC model, we then created stochastic maps for each family in BioGeoBEARS, each representing a single hypothesis for which ranges each lineage occupied from the root to the tip of the tree.
Tropical and temperate breeding habitats and reconstruction

To investigate the impact of latitude on trait evolution in 2-regime models, we assigned each species to either the “tropical” or “temperate” regime, based on its breeding range (i.e., a species that breeds exclusively in the temperate zones but migrates to the tropics when not breeding is assigned to the temperate zone). We focused on the breeding ranges of all species as they are likely to be the arena of strongest competition over territorial space and food. To do this, we first assigned each species to either “tropical,” “temperate,” or “both” based on breeding range limits extracted from range data in shapefiles and defining the tropics as the region between $-23.437^\circ$ and $23.437^\circ$ latitude. We then fit a continuous-time reversible Markov model where transitions between all categories were allowed to occur at different rates, using make.simmap in phytools [83] on the MCC tree. We then used the ML transition matrix to create a bank of stochastic maps under this model, each indicating a possible historical reconstruction of tropical versus temperate habitats through time from the root to tips (S1 Fig). In each stochastic map, we collapsed the “both” category and the “temperate” category to compare lineages with exclusively tropical ranges to lineages with breeding ranges that include temperate regions. Therefore, our “tropical” category indicates that a species breeds exclusively in the tropics, and our “temperate” category contains all species with breeding ranges that include the temperate zone (S4 Fig).

We note that this is a relatively simplistic way of categorizing tropical and temperate membership, and we hope that future methods will enable more sophisticated inferences of historical biogeography alongside paleolatitude and/or paleoclimate. However, given the scope of our analyses, and the emerging evidence that many tropical species ranges have shifted over the timescale of this study [84,85], we opted to keep the results of the historical biogeographical inference and the latitudinal regime reconstruction independent. Future extensions may accommodate the development of more sophisticated paleolatitude models, as well as interactions between various abiotic (e.g., global climate fluctuation [57]) and biotic factors.

Accounting for uncertainty in historical biogeography and latitude

We accounted for uncertainty in ancestral reconstructions by fitting phenotypic models on at least 20 stochastic maps of ancestral tropical/temperate range membership (for all 2-regime models) and/or biogeography (for all models incorporating competition, in both single- and 2-regime versions). For the single-regime model fits that included competition (i.e., DD and MC models), we computed model support and parameter estimates as means across fits conducted on stochastic maps of ancestral biogeography. For the 2-regime model fits, we computed model support and parameter estimates as means across fits conducted on stochastic maps of ancestral tropical/temperate range membership. For the 2-regime model fits with competition, these means also account for variation in estimates of ancestral biogeography (S1 Fig).

Given the scope of these analyses, we chose to account for uncertainty in the biogeographic reconstructions and in the ancestral reconstruction of tropical/temperate living while keeping the topology fixed under the MCC tree. A previous study with a similar model fitting approach found that results on MCC trees were highly concordant with results fit to trees sampled from the posterior distribution [56]. Moreover, there is no reason, to our knowledge, why basing inferences on the MCC tree would bias conclusions about latitude in any systematic way.

Latitudinal variation in mode of phenotypic evolution

We tested whether modes of phenotypic evolution varied with latitude in several ways. First, we used “single-regime” models (Table 1), i.e., models that estimate a single set of parameters
on the entire phylogeny regardless of whether lineages are tropical or temperate. We tested whether support for each of these single-regime models varied according to a clade-level index of tropicality (i.e., the proportion of species in each clade with exclusively tropical breeding ranges). Second, we used our newly developed “2-regime” models (Table 1) with distinct sets of parameters for tropical and temperate species and tested whether these latitudinal models were better supported than models without latitude.

We used ML optimization to fit several “single-regime” models of trait evolution to the 7 morphological trait values described above. For all families, we fitted a set of 6 previously described models [43] that include 3 models (BM, OU, and EB) of independent evolution across lineages, implemented in the R-package mvMORPH [71], and 3 further models (DD<sub>exp</sub>, DD<sub>lin</sub>, and MC) that incorporate competition and biogeography, implemented in the R-package RPANDA [86]. For details of reconstruction of ancestral biogeography, see S1 Text. In the DD models, the slope parameters can be either positive or negative, meaning that species diversity could itself accelerate trait evolution (positive diversity dependence), with increasing species richness driving an ever-changing adaptive landscape [4,67], or, alternatively, increasing species diversity could drive a concomitant decrease in evolutionary rates (negative diversity dependence), as might be expected if increases in species richness correspond to a decrease in ecological opportunity [87].

In cases where families were too large to fit because of computational limits for the MC model (>200 spp., n = 13), we identified subclades to which we could fit the full set of models using a slicing algorithm to isolate smaller subtrees within large families. To generate subtrees, we slid from the root of the tree toward the tips, cutting at each small interval (0.1 Myr) until all resulting clades had fewer than 200 tips. We then collected all resulting subclades and fitted the models separately for each subclade with 10 or more species separately, resulting in an additional 28 clades (n = 136 total).

In addition to this set of models, we fitted a second version of each of these models where the parameters were estimated separately for lineages with exclusively tropical distributions and lineages with ranges that include the temperate region (i.e., “2-regime” models; S1 Text and S2 Fig), limiting our analyses to clades with trait data for more than 10 lineages in each of temperate and tropical regions (S1 Fig; for details of ancestral reconstruction of tropical and temperate habitats, see S1 Text and S4 Fig). The BM and OU versions of these latitudinal models were fit using the functions mvBM and mvOU in the R package mvMORPH [71], and the latitudinal EB, MC, and DD models were fitted with the newly developed functions available in RPANDA [86].

We examined model support in 2 ways. First, we calculated the Akaike weights of individual models [88], as well as the overall support for any model incorporating species interactions and overall support for any 2-regime model. Second, we identified the best-fit model as the model with the lowest small-sample corrected Akaike information criterion (AICc) value, unless a model with fewer parameters had a ΔAICc value <2 [88], in which case we considered the simpler model with the next lowest AICc value to be the best-fitting model.

**Latitudinal variation in strength of interactions and tempo of phenotypic evolution**

We tested for latitudinal variation in the effect of species interactions on trait evolution using both our single- and 2-regime model fits. With the first class of model, we tested whether parameters that estimate the impact of competition on trait evolution (i.e., the slope parameters of the DD models and the S parameter from the MC model) estimated from our single-regime models varied according to the proportion of lineages in each clade that breed...
exclusively in the tropics. With the second class of models, we tested whether 2-regime models estimated a larger impact of competition on trait evolution in tropical than in temperate lineages.

Similarly, we tested whether lineages breeding at low latitudes experience lower or higher rates of morphological evolution compared to temperate lineages using our 2 types of models. First, we tested whether rates of morphological evolution varied according to the proportion of lineages in each clade that breed exclusively in the tropics. We estimated this rate directly as the $\sigma^2$ parameter from the single-regime BM model. For the single-regime EB and DD models, we calculated estimates of evolutionary rates at the present from estimates of the rate at the root and the slope parameters. Second, we compared rates estimated separately for tropical and temperate lineages from the 2-regime implementations of the BM, EB, and DD models. We also examined the impact of observational error on rate estimates by fitting single-regime and 2-regime BM models without accounting for observational error.

**Examining the potential impact of assuming continental scale sympathy**

Our biogeographical reconstructions add important realism into models of species interactions. Nevertheless, species that occur on the same continent do not necessarily interact with one another. We conducted a simulation analysis to determine how our ability to detect the impact of competition on trait evolution may be impacted by the fact that only a subset of the species occurring in a given continent are actually sympatric.

First, we determined the proportion of species that are sympatric within each continent. We calculated range-wide overlap for all family members that ever coexist on the same continent from BirdLife range maps [77] (S6 Data). We defined sympathy as 20% range overlap according to the Szymkiewicz–Simpson coefficient (i.e., overlap area/min(sp1 area, sp2 area)). We also determined if overall levels of sympathy vary latitudinally; to do so, we subset pairs of taxa whose latitudinal means are separated by less than 25˚ latitude [36] and calculated the midpoint latitude for each pair.

Next, we conducted a simulation study to determine whether competition unfolding between ‘truly’ sympatric species only (i.e., at a level finer than the course continental scale we employed) would systematically impact the fit (i.e., model selection) or performance (i.e., parameter estimation) of the 2-regime competition (MC) models for which we used continental-level sympathy (as in the empirical analyses). To do this, we selected 3 clades spanning the range of tree sizes, each with some traits best fit by single-regime MC model, but none best fit by 2-regime MC model (Cracidae.0 [N = 50, N$_tropical$ = 38, N$_temperate$ = 12], Nectariniidae.0 [N = 122, N$_tropical$ = 89, N$_temperate$ = 33], Picidae.1 [N = 190, N$_tropical$ = 86, N$_temperate$ = 104]). For each of these clades, we simulated 2 biogeographic scenarios reflecting empirical levels of sympathy (see above). In the first, we downsampled the continental biogeography such that 50% of tropical and 50% of temperate taxa that were estimated to occur in the same continent were sympatric (see S1 Text for more details). In the second scenario, to reflect the observed latitudinal variation in sympathy, we downsampled the continental biogeography such that 33% of tropical and 50% of temperate taxa that were estimated to occur in the same continent were sympatric (see S1 Text for more details).

With these downsampled biogeographic histories, representing hypothetical range overlap that is more realistic than our continental-level assumption of sympathy, we simulated trait evolution under the 2-regime MC model. For each clade, we used the mean $\sigma^2$ value estimated under the single-regime MC model in empirical fits of a trait that was best fit by the single-regime MC model. We then varied the ratio of the $S_{tropical}/S_{temperate}$ within the range of values in other trait-by-clade combinations where the 2-regime MC model was the best-fit model
(S12 Table). For each clade, parameter combination, and downsampled biogeographic scenario, we simulated 100 datasets, for a total of 3,000 simulated datasets. Finally, we fit the same 12 models that were used in empirical analyses. We conducted model selection to identify the best-fit model for each simulated dataset and assessed whether the estimated $\ln(|S_{\text{tropical}}|/|S_{\text{temperate}}|)$ had the sign expected given the simulated ratio of $S_{\text{tropical}}/S_{\text{temperate}}$ (S9 Data).

**Predictors of support for models with competition**

To identify factors other than latitude which influence whether models with competition were favored by model selection, we examined the impact of habitat (the proportion of species in single-strata habitats), territoriality (the proportion of species with strong territoriality), diet specialization (calculated as the Shannon diversity of diets among species in a clade), clade age, clade richness, and the maximum proportion of species co-occurring on a continent.

**Statistical approach**

We tested for an impact of the proportion of species in a clade that breed exclusively in the tropics on model support and parameter estimates in single-regime models by conducting phylogenetic generalized least squares (PGLS) using the pgls function in the R package caper [89], estimating phylogenetic signal ($\lambda$) using ML optimization, constraining values to $0 \leq \lambda \leq 1$. We tested support for the 2-regime versions of each model type (BM, OU, EB, DD, and MC) across families for a given trait by fitting intercept-only PGLS models with support for latitudinal models as the response variable. We conducted similar analyses to test overall support for latitudinal models across families for each trait and for differences in parameter estimates for tropical and temperate taxa. We found that statistical support for models incorporating competition was relatively rare in small clades (S6 Fig). As this pattern could be related to lower statistical power in smaller datasets [43], we focused all analyses of evolutionary mode (i.e., model support and parameter estimates from models incorporating competition) on clades with at least 50 species ($n = 66$ for single-regime fits and $n = 59$ for 2-regime fits).

For analyses of predictors of support for models with competition, we used the R package MCMCglmm [90] to fit phylogenetic generalized linear mixed models with categorical response variables indicating whether MC or $\text{DD}_\text{exp}$ models were chosen as the best-fit model (S12 Data).

**Supporting information**

S1 Text. Supporting information methods. (DOCX)

S1 Table. Parameters used for simulations generating datasets used to test 2-regime models. (DOCX)

S2 Table. Description of morphological variables. (DOCX)

S3 Table. Loadings for pPC axes of bill and locomotion measurements. pPC, phylogenetic principal component. (DOCX)

S4 Table. PGLS models of statistical support as a function of the latitudinal distribution (measured as the proportion of lineages with individuals that breed in tropical regions).
Statistical support was measured as the mean Akaike weights of single-regime models (i.e., calculated from pool of single-regime models only), and relative support for a model with competition, defined as the maximum Akaike weight for a model with competition divided by the sum of this value and the maximum Akaike weight for a model without competition \([\max(MC_{wi}, DD_{lin, wi}, DD_{exp, wi})/((\max(BM_{wi}, OU_{wi}, EB_{wi})+\max(MC_{wi}, DD_{lin, wi}, DD_{exp, wi}))])\), limiting analyses to clades with \(\geq 50\) tips \((n = 66)\). Values indicated in bold are those that are significant after controlling for multiple testing \((\alpha = 0.05/7)\). \(\lambda\) indicates the MLE of the phylogenetic signal. BM, Brownian motion; DD, diversity-dependent; DD_{exp}, exponential diversity-dependent; DD_{lin}, linear diversity-dependent; EB, early burst; MC, matching competition; MLE, maximum likelihood estimate; OU, Ornstein–Uhlenbeck; PGLS, phylogenetic generalized least squares.

**S5 Table.** Intercept-only PGLS models fit to indices of support for 2-regime models for each trait (for cases where \(N \geq 50\); \(n = 59\)). The index of relative support for any 2-regime model was calculated using \(\max(2\text{-regime Akaike weight})/(\max(2\text{-regime Akaike weight}) + \max(\text{single-regime Akaike weight}))\); other model specific indices were calculated using \(\max(2\text{-regime Akaike weight for specified model})/(\max(2\text{-regime Akaike weight for specified model}) + \max(\text{single-regime Akaike weight for specified model}))\). For each model, this index was transformed by subtracting 0.5 such that negative estimates indicate support for a single-regime model and positive values equal support for a 2-regime model. Values indicated in bold are those that are significant after controlling for multiple testing \((\alpha = 0.05/7)\). For all significant cases, the single-regime version of the model was supported over the 2-regime version. \(\lambda\) indicates the MLE of the phylogenetic signal. MLE, maximum likelihood estimate; PGLS, phylogenetic generalized least squares.

**S6 Table.** PGLS models comparing the observed latitudinal distribution (measured as the proportion of lineages with individuals that breed in tropical regions) of clade-by-trait level fits with the mean MLEs (across fits conducted on a bank of stochastic maps of ancestral biogeography) of the strength of species interactions in single-regime models incorporating competition. All comparisons were conducted on clades with \(\geq 50\) species \((n = 66)\). Note: One outlier was removed from the exponential diversity dependence analysis of bill pPC2. Values indicated in bold are those that are significant after controlling for multiple testing \((\alpha = 0.05/7)\). \(\lambda\) indicates the MLE of the phylogenetic signal. MLE, maximum likelihood estimate; PGLS, phylogenetic generalized least squares; pPC, phylogenetic principal component.

**S7 Table.** Intercept-only PGLS models linear regressions fit to tropical/temperate comparisons of ML parameter estimates of the strength of species interactions in 2-regime models (for cases where \(N \geq 50\) \((n = 59)\) for each trait. For each evolutionary model \((a: MC, b: DD_{exp}, \text{and } c: DD_{lin})\), the mean (across fits conducted on a bank of stochastic maps of ancestral biogeography and stochastic maps of breeding range) of the log-transformed ratio of the absolute value of parameter estimates for tropical taxa to that of temperate taxa \((\ln(|\text{par}_{\text{tropical}}|/|\text{par}_{\text{temperate}}|))\) was the response variable in the intercept-only PGLS model. Negative estimates, therefore, indicate that the impact of competition is estimated to be higher in temperate regions, whereas positive estimates indicate that competition is higher in the tropics. Values indicated in bold are those that are significant after controlling for multiple testing \((\alpha = 0.05/7)\). \(\lambda\) indicates the MLE of the phylogenetic signal. DD, diversity-dependent; DD_{exp}. 

No effect of latitude on avian morphological evolution
exponential diversity-dependent; $DD_{\text{lin}}$, linear diversity-dependent; MLE, maximum likelihood estimate; PGLS, phylogenetic generalized least squares.

**S8 Table.** Intercept-only PGLS models linear regressions fit to tropical/temperate comparisons of ML parameter estimates of the strength of species interactions in 2-regime models (for cases where $N \geq 100$) ($n = 34$) for each trait. For each evolutionary model ($a$: MC, $b$: $DD_{\text{exp}}$, $c$: $DD_{\text{lin}}$), the mean (across fits conducted on a bank of stochastic maps of ancestral biogeography and stochastic maps of breeding range) of the log-transformed ratio of the absolute value of parameter estimates for tropical taxa to that of temperate taxa ($\ln(|\text{par}_{\text{tropical}}|/|\text{par}_{\text{temperate}}|)$) was the response variable in the intercept-only PGLS model. Negative estimates, therefore, indicate that the impact of competition is estimated to be higher in temperate regions, whereas positive estimates indicate that competition is higher in the tropics. Values indicated in bold are those that are significant after controlling for multiple testing ($\alpha = 0.05/7$). $\lambda$ indicates the MLE of the phylogenetic signal. $DD_{\text{exp}}$, exponential diversity-dependent; $DD_{\text{lin}}$, linear diversity-dependent; ML, maximum likelihood; MLE, maximum likelihood estimate; PGLS, phylogenetic generalized least squares.

**S9 Table.** Zero-intercept mixed-effect linear model with a random effect for clade identity fit to the proportion of lineages pairs in each clade that are sympatric in each continent.

**S10 Table.** Intercept-only mixed-effect linear model with a random effect for clade identity fit to the proportion of lineages pairs that are sympatric in each clade.

**S11 Table.** Linear model fit to the proportion of lineages pairs that are sympatric as a function of the absolute value of midpoint latitude for species pairs.

**S12 Table.** Simulation parameters used in simulation study to explore the statistical power of 2-regime MC models under realistic levels of sympathy. Values were chosen based on MLEs from single-regime MC models. MC, matching competition; MLE, maximum likelihood estimate.

**S13 Table.** PGLS analyses of MLEs of evolutionary rates in single-regime model fits ($n = 135$) as a function of the latitudinal distribution (measured as the proportion of lineages with individuals that breed in tropical regions). For DD models, parameter estimates are the mean estimates across fits conducted on a bank of stochastic maps of ancestral biogeography. $\lambda$ indicates the MLE of the phylogenetic signal. DD, diversity-dependent; MLE, maximum likelihood estimate; PGLS, phylogenetic generalized least squares.

**S14 Table.** Intercept-only PGLS models fit to the difference between tropical and temperate ML parameter estimates of evolutionary rates in 2-regime models, fit separately for each trait ($n = 71$ for ln.mass and $n = 70$ for other traits). For DD models, the rate parameter was calculated as the mean comparisons between parameter estimates across fits conducted on a bank of stochastic maps of ancestral biogeography and stochastic maps of breeding range. Note: One outlier was removed from the linear diversity dependence analysis of locomotion pPC2 as it was $>2$ orders of magnitude larger than the next largest value. Values indicated in...
bold are those that are significant after controlling for multiple testing ($\alpha = 0.05/7$). $\lambda$ indicates the MLE of the phylogenetic signal. ML, maximum likelihood; MLE, maximum likelihood estimate; PGLS, phylogenetic generalized least squares; pPC, phylogenetic principal component.

(S15 Table) PGLS models comparing the observed latitudinal distribution (measured as the proportion of lineages with individuals that breed in tropical regions) of clade-by-trait level fits ($n = 135$) with the log-transformed error (calculated as the sum of the MLE error parameter and the clade-level mean squared standard error) in single-regime BM models. Values indicated in bold are those that are significant after controlling for multiple testing ($\alpha = 0.05/7$). $\lambda$ indicates the MLE of the phylogenetic signal. BM, Brownian motion; MLE, maximum likelihood estimate; PGLS, phylogenetic generalized least squares.

(S16 Table) PGLS models comparing the observed latitudinal distribution (measured as the proportion of lineages with individuals that breed in tropical regions) of clade-by-trait level fits ($n = 135$) with the ML parameter estimates of evolutionary rates in single-regime BM models that do not account for observational error. Values indicated in bold are those that are significant after controlling for multiple testing ($\alpha = 0.05/7$). $\lambda$ indicates the MLE of the phylogenetic signal. BM, Brownian motion; MLE, maximum likelihood estimate; PGLS, phylogenetic generalized least squares.

(S17 Table) Intercept only PGLS models fit to the mean difference (across stochastic maps of tropical and temperate living) in MLE estimates of tropical and temperate rates (from 2-rate BM models that do not account for observational error) ($n = 71$ for log-transformed body mass, $70$ for other traits). Values indicated in bold are those that are significant after controlling for multiple testing ($\alpha = 0.05/7$). $\lambda$ indicates the MLE of the phylogenetic signal. BM, Brownian motion; MLE, maximum likelihood estimate; PGLS, phylogenetic generalized least squares.

(S18 Table) The factors predicting which clades support models with competition, as revealed by PGLMMs fit to single-regime clade-by-trait fits ($n = 924$) with a categorical variable indicating (a) that the MC was the modal best-fit model (i.e., the most common best-fit model across fits conducted on a bank of stochastic maps of ancestral biogeography) ($n = 166$) or (b) that the DD$^{exp}$ model was the model best-fit model ($n = 66$) (S12 Data). The influence of the phylogeny was estimated from the random effect component of the PGLMM—the phylogenetic intraclass correlation coefficient is analogous to the $\lambda$ parameter (often referred to as “phylogenetic signal”) estimated from PGLS models [91]. To facilitate parameter exploration, we rescaled all predictor variables using $z$-transformations. We used an uninformative, inverse Wishart distribution as a prior for the random effects, a flat prior for the fixed effects, and fixed the residual variance at 1 [92]. To fit the models, we ran an MCMC chain for at least $5 \times 10^5$ generations, recording model results every 100 generations and ignoring the first $5 \times 10^3$ generations as burn-in. We fit each model 4 times and merged the 4 chains after verifying convergence both visually and using Gelman–Rubin diagnostics in the R-package coda [93,94]. Estimates and credibility intervals are therefore calculated from the pooled posterior distributions. The pMCMC (an MCMC derived $p$-value calculated as 2 times the proportion of estimates in either the positive or negative portion (whichever is smaller) of the posterior distribution) is presented from one chain. DD, diversity-dependent; DD$^{exp}$, exponential.
diversity-dependent; MC, matching competition; MCMC, Markov chain Monte Carlo; PGLMM, phylogenetic generalized linear mixed model; PGLS, phylogenetic generalized least squares; pMCMC, Markov chain Monte Carlo.

S1 Fig. Illustration of our model-fitting approach for clade-level model fits with different strengths of competition in tropical and temperate regions. We combine a matrix of the presence or absence of each lineage in tropical/temperate regions (“regime matrix”) with a matrix of biogeography (denoted “A”) to identify the competitive regime of each lineage and the identity of other lineages with which the focal lineage is able to interact with. Blue and red colors in the lower panel denote correspondence between the formula and the biogeography matrix (A) and the regime matrix, respectively.

S2 Fig. Results of the simulation study demonstrate the ML optimization returns reliable parameter estimates in 2-regime models. (a–d) Exponential time-dependent model. (e–h) DD_{exp} model. (i–l) DD_{lin} model. (m–p) MC model. In all plots, the red lines denote the parameters used to generate the simulated data (S7 Data). DD, diversity-dependent; DD_{exp}, exponential diversity-dependent; DD_{lin}, linear diversity-dependent; MC, matching competition; ML, maximum likelihood.

S3 Fig. Results of model selection depicting best-fitting models for data simulated under (a) 2-regime BM, (b) 2-regime OU, and (c) 2-regime EB models across a range of parameter values (S8 Data). BM, Brownian motion; EB, early burst; OU, Ornstein–Uhlenbeck.

S4 Fig. Clade-level distributions of tropical, temperate, and widespread breeding (a) sorted by clade name, (b) sorted by proportion of exclusively tropical breeding species, and (c and d) presented as separate histograms. The number following the family name indicates the subclade within that family (see Methods and S4 and S5 Data).

S5 Fig. Continental variation in the proportion of species that co-occur in sympatry (defined as 20% range overlap) (S6 Data).

S6 Fig. Clade size impacts the probability that a model incorporating competition is the modal best-fit single-regime model (i.e., the most common best-fit model across fits conducted on a bank of stochastic maps of ancestral biogeography and stochastic maps of breeding range) (S2 and S3 Datas).

S7 Fig. Best-fit models for each clade-by-trait combination shows that single-regime models generally outperform 2-regime models, although some clades (e.g., Meliphagidae and Phasianidae) do tend to support models with latitude across several traits. Shown is the modal best-fit model (i.e., the most common best-fit model across fits conducted on a bank of stochastic maps of ancestral biogeography across fits conducted on a bank of stochastic maps of ancestral biogeography and stochastic maps of breeding range). The number following the family name indicates the subclade within that family (see Methods and S4 and S5 Datas).
S8 Fig. Results from simulation analyses exploring the impact of assuming continental level sympatry for 3 clades. (a–c) Best-fit models for data generated under downsampled biogeographic scenario #1 (i.e., 50% of both tropical and temperate lineages set to allopatric at a continental scale). (d–f) Best-fit models for data generated under downsampled biogeographic scenario #2 (i.e., 50% of temperate lineages and 66.6% of tropical lineages set to allopatric at a continental scale). (g–i) The proportion of simulations for which MLEs of the ratio of competition from the 2-regime MC model (i.e., ln(|S_{tropical}|/|S_{temperate}|)) correctly identify the direction of the difference in the strength of competition (S9 Data). MLE, maximum likelihood estimate.

(TIFF)

S9 Fig. Evolutionary rates in other single-regime models (a: EB, b: DD_{exp}, and c: DD_{lin}) do not vary as a function of the proportion of lineages that breed in the tropics. For DD models, parameter estimates are the mean estimates across fits conducted on a bank of stochastic maps of ancestral biogeography (S2 and S3 Data). DD, diversity-dependent; DD_{exp}, exponential diversity-dependent; DD_{lin}, linear diversity-dependent; EB, early burst.

(TIFF)

S10 Fig. Differences between rates estimated separately on tropical and temperate taxa in 2-regime models (a: EB, b: DD_{exp}, and c: DD_{lin}). Shown are the mean comparisons between parameter estimates across fits conducted on a bank of stochastic maps of ancestral biogeography and stochastic maps of breeding range (i.e., tropical or temperate). Asterisks indicate statistical significance (S4 and S5 Data). DD, diversity-dependent; DD_{exp}, exponential diversity-dependent; DD_{lin}, linear diversity-dependent; EB, early burst.

(TIFF)

S11 Fig. The relationship between the total error (calculated as the log-transformed sum of the MLE nuisance error parameter from single-regime BM models and the clade-level mean squared standard error) and the proportion of tropical breeding lineages in a clade is negative for body mass, but not for other traits. Solid lines represent statistically significant relationships (S15 Table and S10 Data). BM, Brownian motion; MLE, maximum likelihood estimate.

(TIFF)

S12 Fig. BM models of trait evolution fit at a clade level when not accounting for observational error reveal a more pronounced relationship between rate and latitude for several traits. (a) There is a negative relationship between the proportion of taxa in a clade that breed in the tropics and the estimated rate of trait evolution from single-rate BM models for body mass and locomotion pPC3, but not other traits. Color of points indicate trait (as in panel b). (b) Differences between rates estimated separately on tropical and temperate taxa in 2-rate BM models are biased toward faster rates in temperate regions for body mass and locomotion pPC3, but not other traits. Shown are the mean comparisons between parameter estimates across fits conducted on a bank of stochastic maps of ancestral biogeography and stochastic maps of breeding range (i.e., tropical or temperate) (S11 Data). BM, Brownian motion; pPC, phylogenetic principal component.

(TIFF)

S13 Fig. Best-fit “single-regime” models for each clade-by-trait combination show that, while BM is most often the best model, several clades show evidence of MC (e.g., Cotingidae, Formicariidae, Malaconotidae, and Paridae) or diversity dependence (e.g., Strigidae, Fringillidae, and Columbidae subclade 2) acting on several traits. Shown is the modal best-
fit model across fits conducted on a bank of stochastic maps of ancestral biogeography. The number following the family name indicates the subclade within that family (see Methods and S2 and S3 Datas). BM, Brownian motion; MC, matching competition.

S14 Fig. Best-fit single-regime models (modal best fit across fits conducted on a bank of stochastic maps of ancestral biogeography), plotted as a function of total clade size and the number of species in each clade that occur on the same continent. (A) All models. (B) MC and DD$_{exp}$ models. Each point represents a clade-by-trait combination (i.e., each clade contributes a point for each of 7 traits). In both panels, points are jittered slightly to aid visualization (S2 and S3 Datas). DD$_{exp}$, exponential diversity-dependent; MC, matching competition.

S1 Data. Species-level trait data used in analyses.

S2 Data. Results of all individual single-regime fits.

S3 Data. Results of individual single-regime fits, summarized for each clade-by-trait combination.

S4 Data. Results of all individual 2-regime fits.

S5 Data. Results of individual 2-regime fits, summarized for each clade-by-trait combination.

S6 Data. Species range-wide overlap data calculated from BirdLife shapefiles.

S7 Data. Results from simulation exercise exploring the parameter estimability in newly developed 2-regime models.

S8 Data. Results from simulation exercise exploring model selection performance of 2-regime BM, OU, and EB models. BM, Brownian motion; EB, early burst; OU, Ornstein–Uhlenbeck.

S9 Data. Results from simulation exercise exploring the impact of assuming continent-scale sympatry on the performance of 2-regime MC models. MC, matching competition.

S10 Data. Total error (sum of the MLE nuisance error parameter from single-regime BM models and the clade-level mean squared standard error) for each clade-by-trait combination. BM, Brownian motion; MLE, maximum likelihood estimate.

S11 Data. Results of single-regime and 2-regime fits of BM models excluding observational error. BM, Brownian motion.
S12 Data. Data used for PLMM analyses of predictors for support for either MC or DD_{exp} models in single-regime fits. DD_{exp}, exponential diversity-dependent; MC, matching competition; PLMM, phylogenetic linear mixed model.

S13 Data. Species-level maximum clade credibility tree used during model fitting.

S14 Data. Clade-level maximum clade credibility tree used for PGLS and PLMM analyses. PGLS, phylogenetic generalized least squares; PLMM, phylogenetic linear mixed model.

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