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PREDICTIVE REVIEW

# Testing for changes in rate of evolution and position of the climatic niche of clades

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#### ABSTRACT

- 1. There is solid recognition that phylogenetic effects must be acknowledged to appreciate climatic niche variability among species clades properly. Yet, most currently available methods either work at the intra-specific level (hence they ignore phylogeny) or rely on the Brownian motion model of evolution to estimate phylogenetic effects on climatic niche variation. The Brownian motion model may be inappropriate to describe niche evolution in several cases, and even a significant phylogenetic signal in climatic variables does not indicate that the effect of shared ancestry was relevant to niche evolution.
- 2. We introduce a new phylogenetic comparative method which describes significant changes in the width and position of the climatic niche at the interspecific (clade) level, while making no *a priori* assumption about how niche evolution took place.
- 3. We devised the R function phylo.niche.shift to estimate whether the climatic niches of individual clades in the tree are either wider or narrower than expected, and whether the niche occupies unexpected climates. We tested phylo.niche.shift on realistic virtual species' distribution patterns applied to a phylogeny of 365 extant primate species.
- 4. We demonstrate via simulations that the new method is fast and accurate under widely different climatic niche evolution scenarios.phylo.niche.shift showed that the capuchin monkeys and langurs occupy much wider, and prosimian much narrower, climatic niche space than expected by their phylogenetic positions.
- 5. phylo.niche.shift may help to improve research on niche evolution by allowing researchers to test specific hypotheses on the factors affecting clades' realised niche width and position, and the potential effects of climate change on species' distributions.

#### INTRODUCTION

The climatic niche of a species is defined by the set of temperature and precipitation regimes it experiences (Holt & Gaines 1992). Although the climatic niche is not a phenotypic trait per se, climatic preferences depend directly on the species' physiology and lifestyle (e.g. thermal tolerance limits, dietary preferences, body size), so that the evolution of climatic niches can be studied as if they were phenotypes (Holt & Gaines 1992, Rolland et al. 2018). Starting from this broad assumption, several investigations aimed to evaluate the importance of phylogenetic effects vs. contingency (e.g. competition, local adaptation, human disturbance) at determining the distribution of species (Freckleton & Jetz 2009, Corro et al. 2021), or whether climatic niche variability correlates with species diversity (Kozak & Wiens 2010, Rolland & Salamin 2016). Many studies apply the calculation of the phylogenetic signal (Blomberg et al. 2003) in niche variability as a measure of the degree of phenotypic resemblance between species. In the context of climatic niche evolution, high phylogenetic signal would indicate that phylogenetically close species live under more similar climates than distant species, provided the Brownian motion model (BM) is an appropriate descriptor of climatic niche evolution (Freckleton 2009, Kamilar & Cooper 2013, Boucher et al. 2014). Studies consistently indicate a generally weak yet irregular phylogenetic signal in climatic variables (Song et al. 2016, Perez & Feeley 2021); that the upper (warm) thermal tolerance limit is more phylogenetically structured than lower (cold) limit (Diamond & Chick 2018, Lancaster & Humphreys 2020, Perez & Feeley 2021); and that niche breadth correlates with both geographic range size and diversity (Kozak & Wiens 2010, Rolland & Salamin 2016).

Although it is widely used to infer the imprint of shared ancestry on climatic niche evolution, the phylogenetic signal has several limitations. It ignores potential differences between clades in the rate of climatic niche evolution, and describes a pattern rather than a process, meaning that high phylogenetic signal may derive from spatial proximity alone, with no effect of shared ancestry (Freckleton & Jetz 2009). Still, geologically rapid changes in the position of the climatic niche occupied by a clade may result from the emergence of key innovations, dispersals, or changes in the competition regimes that allow for the successful colonisation of previously unexplored habitats (Lancaster & Humphreys 2020). For instance, species belonging to clades distributed at low latitudes are often reported to exploit narrower climatic niches than temperate species clades, suggesting a slower rate of climatic niche evolution in the Tropics (Kozak & Wiens 2010, Rolland et al. 2018). Middle Pleistocene human species were shown to deviate significantly from their ancestors' habitual climatic niche

thanks to cultural innovation (Mondanaro et al. 2020). Panicoideae grasses (e.g. maize, switchgrass) expanded their niches to colonise regions with higher growing season temperatures by exploiting the C4 photosynthetic pathway (Aagesen et al. 2016). Simulation experiments show that latitudinal gradients in niche breadth may occur even without any change in the fundamental species' niche (Saupe et al. 2019), which suggests that the rate of climatic niche evolution and its position should be investigated together.

We devised a method, embodied in the R function phylo.niche.shift, which locates major changes in the rate of evolution and in the position of the climatic niche of clades contained within a phylogenetic tree. The method builds on phylogenetic ridge regression as implemented in the RRphylo R function (Castiglione et al. 2018). Contrary to BM, RRphylo does not assume a single rate of evolution operating through the phylogeny, or that trait (here bioclimatic variables) means should not change over evolutionary time. By using realistic species' distribution patterns, we demonstrate via simulations that phylo.niche.shift is especially powerful at pursuing both goals under widely different models of climatic niche evolution. We further applied the method to a phylogeny of 365 extant primate species.

#### **METHODS**

#### Simulating virtual species' distributions

Our aim was to test phylo.niche.shift on a realistic representation of climatic niche evolution on phylogenetic trees, covering different situations, from negligible to important climatic niche similarity within the clades, while reproducing genuine macroecological patterns of species' distribution on Earth.

We started by locating, on continental Eurasia (the sampling area ranges from -20° to 180° longitude, and from 0° to 90° latitude), a number of virtual species, randomly chosen between 100 and 200. The species are given a certain number of geographic occurrences each, drawn from a negative exponential distribution with rate parameter  $\lambda = 0.1$ , and bound between 100 and 5 occurrences. This way, a few species will be abundant whereas the majority will be rare, in keeping with the well-known shape of the species' abundance-frequency distribution (Gaston & Blackburn 2007). For each species, its first occurrence is randomly placed in Eurasia. A square mask ranging from  $10^6$  to  $1.1 \times 10^7$  square metres is centred around this first occurrence, setting the size of the mask to be proportional to the total number of occurrences (imposing a correlation coefficient between the number of occurrences and the size of the mask r = 0.8). All other occurrences are then randomly placed within the mask.

After all occurrences for all species are located, they are 'compressed' towards the equator by applying an exponential transformation with  $\lambda = 0.8$  to the occurrences' latitude, so that the whole set of species mimics the latitudinal diversity gradient (Gaston & Blackburn 2007). Finally, the occurrences falling outside the landmass of Eurasia are discarded. For each species, we calculated the area of the minimum convex polygon enclosing its occurrences to represent the species' geographic range size.

At the geographic location of each individual occurrence, we extracted 19 bioclimatic variables values from the Worldclim database (WorldClim version 2.1; Fick & Hijmans 2017) with the native 5-min spatial resolution. To avoid potential problems of multicollinearity, the full set of 19 bioclimatic variables was reduced considering a variance inflation factor  $\leq 5$  (Zuur et al. 2010; see Table 1 for the variables that were selected).

# Simulating different scenarios of climatic evolution on the tree

For each species, we calculated the mean of each variable over all its occurrences to create a bioclimatic vector representing the 'average' realised climatic conditions experienced by the species. Then, we constructed three different hypothetical phylogenetic trees by means of hierarchical clustering, using the angle  $\phi$  between bioclimatic vectors (which is equivalent to Pearson's correlation coefficient) as the distance metric to guide the clustering algorithm. The trees represent three abstract, widely different scenarios of climatic niche evolution:

1. Anticlimatic Evolution (AE). Under AE, phylogenetic proximity is determined by the difference  $90-\phi$ . Since only few bioclimatic variables (e.g. BIO6, minimum temperature of coldest month) can attain negative values, angles >90° (which would represent negative correlation

coefficients) are inappropriate, and were in fact never produced in the simulations. By clustering species via their climatic dissimilarity, under AE, each clade in the tree will have roughly equal chance to include climatically distant species (Fig. 1a). Therefore, the clades in the tree will be formed by species with widely differing climatic preferences and evolve at similar rates of climatic niche change. The phylogenetic signal in climatic variables is expected to be weak.

- 2. Random Climatic Evolution (RCE). Under RCE, species' phylogenetic proximity is disconnected from climatic proximity (Boucher et al. 2014). To produce RCE, we first computed the angle  $\phi$  between any pair of species' bioclimatic vectors. Then, we arbitrarily substituted  $\phi$  with a new value  $\phi_1$  sampled between  $\phi$  and 90° to feed the agglomeration algorithm in hierarchical clustering by using the 'unweighted pair group method with arithmetic mean'. This way, any signal of climatic resemblance between species is weakened randomly during the tree construction process (Fig. 1b), resulting in the lowest phylogenetic signal of any scenario.
- 3. Climate-driven evolution (CDE). Under CDE, climatic proximity (that is  $\phi$ ) is used to guide the agglomeration algorithm. Therefore, most clades in the tree will be bound to include species with strongly similar climatic preferences, so that climatic variability represents a strong determinant of species' geographic distribution (Fig. 1c). The phylogenetic signal is expected to be large, and climatic preferences will be apportioned between, rather than within, clades, generating presumably large evolutionary rate differences between them.

We ran each scenario 1000 times. At each iteration, we first tested the existence of the latitudinal diversity gradient. To this aim, we partitioned all species' occurrences over consecutive, 5° wide latitudinal bands poleward, and counted the number of species present with at least one occurrence in each band. Then, we regressed species richness per band against the mean latitude of

**Table 1.** A real-case representation of the transformations applied to simulate climatic niche evolution rate shifts to specific clades in the tree. Variable values represent the range of values sampled by the species belonging to the selected clade, pooled together. The bioclimatic variables are from the WorldClim database (WorldClim version 2.1; Fick & Hijmans 2017)

	Temperature			Precipitation				
	Mean diurnal range	Isothermality	Wettest quarter	Driest quarter	Driest month	Seasonality	Warmest quarter	Coldest quarter
	BIO2	BIO3	BIO8	BIO9	BIO14	BIO15	BIO18	BIO19
<u>у</u> 0	9.025	28.852	11.874	-4.518	0	50.92	4.328	9.488
- 0	15.717	61.136	29.078	26.885	18.575	128.305	587.316	127.374
<i>Y</i> <sub>plus</sub>	6.89	20.105	6.545	-16.09	-2.146	28.658	-70.026	-11.691
• plus	16.929	68.53	32.352	31.014	25.717	144.736	804.455	165.138
<i>Y</i> <sub>minus</sub>	10.434	34.626	15.39	3.119	1.416	65.612	53.402	23.465
- minus	14.918	56.255	26.917	24.159	13.861	117.46	444.004	102.449

575



**Fig. 1.** The three different scenarios of species' distributions within clades, representing evolution proceeding according to anticlimactic (a); random (b); or climate-driven (c) preferences in climatic conditions across species.

all the occurrences within each band. We further tested for the existence of the desired positive relationship between geographical range size and latitude (Gutiérrez-Pesquera et al. 2016). We regressed species' range sizes against the mean latitude of the species' occurrences. Both negative and significant diversity-latitude and positive and significant range size–latitude relationships are expected to occur in the data to prove that the simulations were based on realistic species' distribution patterns.

# The phylo.niche.shift algorithm for rate shifts

phylo.niche.shift calculates the evolutionary rate for each branch in the phylogeny, according to phylogenetic ridge regression as implemented in the RRphylo function, embedded in the namesake R package (Castiglione et al. 2018). A function linked to RRphylo, search.shift, still part of the RRphylo toolkit, is called from within phylo.niche.shift to locate on the tree shifts in the absolute evolutionary rates by using a randomisation procedure (Castiglione et al. 2018). A significant shift as attached to a particular clade in the phylogeny indicates that clade has more or less climatic variability than expected from its phylogenetic position. search.shift allows researchers to test for the existence of the rate shifts in climatic niche evolution either automatically, or by indicating specific clades to be tested.

# Simulating rate shifts as applied to a specific clade

To simulate a rate shift in climatic niche evolution as applied to a particular clade in the tree, we started identifying all the subtrees in the phylogeny  $CC_{can}$  including no less than one-tenth and no more than one half of the tree tips. A clade  $CC_{sel}$  was randomly selected among  $CC_{can}$ . The trace of the evolutionary rate matrix (whose diagonal elements indicate the rate of evolution for individual climatic variables; Revell & Harmon 2008) of  $CC_{sel}$  was computed to represent a measure of its

'multivariate' rate of climatic niche evolution. An xs scalar was used to transform each bioclimatic variable for each species belonging to  $CC_{sel}$ , according to the equation:

$$xs * y - (xs - 1) * \overline{y}$$

where *y* is the bioclimatic variable, and  $\overline{y}$  is its mean computed over the CC<sub>sel</sub> species. For *xs*>1 the range of the bioclimatic variable values realised by CC<sub>sel</sub> species will increase, simulating a faster rate of climatic niche evolution, and the other way around at *xs*<1, whereas the variables mean will not change. Table 1 depicts the actual effect of applying a relatively large *xs* (*xs* is bound in between -2 and 2 in the simulations) on the original range of  $y_0$  values for each bioclimatic variable of the species belonging to CC<sub>sel</sub>, either simulating an increase in the rate (with *xs* = 1.5,  $y_{\text{plus}}$  in Table 1) or its inverse (*xs* = 0.67,  $y_{\text{minus}}$  in Table 1).

Although some variables occasionally take unrealistic values (e.g. negative precipitation values in Table 1), our goal was to simulate feasible and proportionally small contractions (or increases) in climatic variability within the selected clade, which is well represented by the potential range of *xs* values we designated. To account for rate heterogeneity and skewed distribution of rates among  $CC_{can}$ , we transformed *xs* to a metric *effect* so that at *effect* = 1,  $CC_{sel}$  evolves at the same rate as the rest of the tree, and with *effect*  $\neq$  1, the impact of rate transformation is symmetric for rate decreases and increases (see Appendix S1). We further calculated Type I error at *effect* = 1.

# The phylo.niche.shift algorithm for niche drifts

To test for potential drift (a change in the position of the climatic niche of the clade) for each  $CC_{can}$ , phylo. niche.shift computes the Mahalanobis distance between the bioclimatic vectors of the species belonging to the focal clade and the mean bioclimatic values of all the other species in the tree as well as their covariance. The Mahalanobis distance is a scale-invariant

generalisation of the Euclidean distance for correlated variables. Therefore, it is best suited to measure the 'distance' between each  $CC_{can}$  and the rest of the tree. In phylo.niche.shift, a family of 100 random Mahalanobis distances is created by extracting from the tree bioclimatic vectors a random subset of *n* species for any clade of size *n* within  $CC_{can}$ , and computing the Mahalanobis distance between this set of *n* species and the rest of the tree. Significance is assessed by comparing the real Mahalanobis distance to the random distances and accepting as significant a real distance outside the range of random distances.

For any clade found to drift  $\mathrm{CC}_{\mathrm{drift}}$  (or for a clade specified by the user), phylo.niche.shift assesses whether individual bioclimatic variables drift from their expected position. The basic idea is to simulate the bioclimatic variables at  $CC_{drift}$  under BM (which produces no drift) and then compare the simulated values to the real values. The algorithm works as follows. First, given the most recent common ancestor  $(CC_{mrca})$  of the species belonging to CC<sub>driff</sub>, the function calculates the bioclimatic vector of ancestral state estimates at  $\mathrm{CC}_{\mathrm{mrca}}$  using both the RRphylo function (Castiglione et al. 2020) and the phyEstimate function in the R package picante (Kembel et al. 2010). With phyEstimate, we calculated the bioclimatic vector at CC<sub>mrca</sub> according to BM, obtained via phylogenetic imputation (Garland & Ives 2000), assuming CC<sub>drift</sub> is represented by a single species with unknown bioclimatic values. To be conservative, for each bioclimatic variable, the ancestral state estimate is chosen between RRphylo and phyEstimate valuations as the one closest to the mean value of the tips descending from CC<sub>mrca</sub>. The resulting bioclimatic vector of estimates at  $\mathrm{CC}_{\mathrm{mrca}}$  is retained and set as the vector of phylogenetic means to simulate 100 new BM distributions of the variables on CC<sub>drift</sub> proceeding with the Brownian rate  $\sigma^2$  calculated for the rest of the tree. A bioclimatic variable is deemed to drift if the mean of the real values falls outside 95% of the range of means generated by the 100 BM distributions.

# Simulating drifts in the position of the climatic niche as applied to a specific clade

To simulate a displacement in the average climatic niche position of a clade, we started sampling a *d* value within a family of 100 values ranging from 0.3 to 3. The scalar *d* was multiplied by the mean bioclimatic vector of the species belonging to the selected clade  $(y_0)$  so as to produce a set of drifted bioclimatic variables. A threefold change in  $y_0$  would, for instance, effectively transform a clade of warm-loving, Tropical species into high-latitude species experiencing cold temperatures, and *vice versa*  (e.g. a change from 5 to 15 °C in mean annual temperature), covering most of the known climatic variability in Eurasia.

Overall, for each tree and each scenario, we produced a starting matrix of bioclimatic vectors, y (each row representing a species), a matrix where a shift in the rate is applied to a selected clade  $CC_{sel}$ , yS, and a matrix where a drift is applied to  $CC_{sel}$ , named yD. We performed six different phylo.niche.shift runs, on y either nonindicating (automatic mode, y-auto), or indicating the  $CC_{sel}$  (node mode, y-node), the same on yS (yS-auto and yS-node) and on yD (yD-auto and yD-node).

To assess the power and sensitivity of phylo. niche.shift in finding rate shifts, we regressed the P-value produced by the function (which is two-tailed so that significance is accepted for either P < 0.025 or P > 0.975; at  $\alpha = 0.05$ ) on *yS-node* against the logarithm of the effect value, using logistic growth function in R. This approach correctly allows the gradual interpretation of P-values (Muff et al. 2021). The 'goodness-of-fit' of the logistic growth is approximated by calculating pthe model's Efron's pseudo-R<sup>2</sup> (Efron 1978), that is the squared correlation between the predicted and actual values. The P-values of effect predicted by the logistic model at P = 0.025 and P = 0.975, represent the level of *effect* necessary to produce significant rate shifts as assessed by phylo. niche.shift. We further counted the percentage of shifts across the simulations on y-node and the Type I error on *y*-node at *effect* = 1 (see Appendix S1). The phylo.niche.shift function repeats, upon indication by the user, the rate shift test assuming the BM model of evolution. To this end, we used function brownie.lite in phytools (Revell 2012) which applies the noncensored approach described by O'Meara et al. (2006) to test for the existence of multiple rates on the tree. We replicated the assessment of falsepositive incidence rate under BM to y-node.

To assess the power and sensitivity of phylo.niche. shift in finding climatic niche drifts, we counted the number of times the bioclimatic vector of  $CC_{sel}$  was found to drift in *yD-node* simulations and calculated the minimum *d* value necessary to provide significance at  $\alpha = 0.05$  for the simulations with both d>1 and d<1. The number of times the bioclimatic vector of  $CC_{sel}$  was found to drift by using *y-node* provides the incidence of Type I errors. We further counted the average number of bioclimatic variables included in each simulation and those found to drift under both *yS-node* and *y-node*.

We estimated power and Type I error in finding shifts by using *yD-node* and in finding drifts by using *yS-node*, to assess how often applied rate shifts provide evidence of drifts and vice versa, since the two are expected to influence each other (Saupe et al. 2019).

#### Applying phylo.niche.shift to climatic niche evolution in primates

We collected 122950 occurrences belonging to 453 records of extant primate species. The occurrences were downloaded from the Global Biodiversity Information Facility online database (https://www.gbif.org). We excluded individual occurrences reported as 'preserved specimens' in the database, and corrected taxonomic and geographic errors, taking the International Union for Conservation of Nature's Red List and range maps as a reference. After this manipulation, the number of accepted occurrences reduced to 89269, and the number of valid species to 398. *Homo sapiens* was excluded from the tree because our climatic niche depends on technology, rather than physiology.

The phylogenetic tree is an informal supertree derived by integrating phylogenetic information from different sources (Appendix S1). We started from the phylogeny available in Melchionna et al. (2020) and updated it with missing species by means of the *tree.merger* function (Castiglione et al. 2022), embedded in RRphylo. Node ages for calibration were derived from TimeTree (Kumar et al. 2017), and adjusted according to known fossil information. The final tree and data included 365 extant species. We applied phylo.niche.shift to the primates' tree and data under the automatic mode, by setting the minimum size of the clades to be scanned for rate shift (*f*) at 15 species.

#### RESULTS

#### Virtual species' distributions: power and accuracy of phylo.niche.shift in detecting rate shifts in climatic niche evolution

The average tree size is 107.8 species. Both the latitudinal diversity and range-size gradients are reproduced in the data, indicating that virtual species' distributions conform to well-known macroecological patterns (Table 2). Under AE, the phylogenetic signal K in the bioclimatic variables distributions falls in the 0.41–0.71 range. Under RCE and CDE, the corresponding figures are and 0.10–0.21 and 0.91–1.20, respectively, confirming our expectation that the signal grows from RCE towards CDE.

The logistic growth model provides very high Efron's  $R^2$  for AE (0.982, Table 3; Fig. 2) and still large  $R^2$  for both RCE and CDE (0.897 and 0.887, respectively, Table 3; Appendix S1 Figs. S1.2, S1.5). Despite the profound differences between the three scenarios, the significance levels

**Table 2.** The proportion of iterations where a significant negative latitudinal diversity gradient and significant positive latitudinal gradient in range size apply under each climatic evolution scenario (anticlimactic evolution – AE, random climate evolution – RCE, and climate-driven evolution – CDE). The average phylogenetic signal is calculated across all the selected bioclimatic variables when no rate shift is applied (K y), when the shift is applied to a specific clade (K yS), and when a drift in climatic niche position is applied to a specific clade (K yD)

	AE	RCE	CDE
Latitudinal diversity gradient	0.992	0.992	0.993
Latitudinal range size gradient	0.797	0.815	0.805
Кy	0.41 (0.37–0.66)	0.098 (0.09–0.12)	0.91 (0.79–1.5)
K <i>y</i> S	0.42 (0.37–0.67)	0.098 (0.089–0.13)	0.87 (0.69–1.6)
K yD	0.71 (0.49–1.2)	0.21 (0.12–0.46)	1.20 (0.89–2.3)

of *effect* are remarkably constant (Table 3), providing evidence that phylo.niche.shift is not very sensitive to the underlying mode of evolution of the climatic niche. The percentage of iterations where no shift is applied and yet a shift is returned is low under AE, reaches some 30% under RC0E and raches almost 100% under CDE. This properly reflects the finding that rate variation among  $CC_{can}$  is much larger than would be expected under a single-rate (BM) model (Appendix S1). The incidence of Type I errors on *y*-node provided by phylo.niche. shift is low under all models (Table 3). By using *y*-auto under AE, 33% of the simulations showed at least one significant rate shift. However, large this figure could appear, it must be considered that the entire set of  $CC_{can}$  is tested specifying automatic node. Since each tree contains

**Table 3.** The Efron pseudo-R2 of the logistic regression between the simulated rate shift (*effect*) and the *P*-value returned by phylo.niche.shift. '*effect* down' and '*effect* up' indicate the *effect* value at which 95% of the phylo.niche.shift runs correctly identify the rate shift with a simulated decrease or increase, respectively. No shift is expected to occur at *effect* = 1. The last three rows report: (a). the proportions of rate shifts found across all simulation when no shift is simulated; (b) the same figure calculated assuming the Brownian motion model of climatic niche evolution; and (c) the figure when a drift, rather than a shift, is simulated. Anticlimactic evolution – AE, random climatic evolution – RCE, climate-driven evolution – CDE

	AE	RCE	CDE
Efron pseudo-R2	0.982	0.897	0.887
effect down	2.22	2.4	2.09
effect up	1.43	1.51	1.53
Type I error	0.003	0.071	0.082
a. Incidence of shifts found, y-node	0.044	0.299	0.976
b. Incidence of shifts found, BM, y-node	0.012	0.159	0.471
c. Incidence of shifts found, yD-node	0.66	0.797	0.805



**Fig. 2.** Sensitivity of phylo.niche.shift to the simulated shifts in the rate of climatic niche evolution applied (*y*-axis) under the anticlimatic evolution scenario. Filled dots represent instances of significant results, for either instances of simulated rate decrease (*P*-value < 0.025) or increase (*P*-value > 0.975). The vertical dashed lines represent the values of the logarithm of the simulated effect where the fitted logistic curve (S-curve) reaches predicted *P*-values of 0.025 and 0.975.

12.2  $CC_{can}$  on average, the number of expected simulations finding at least one shift at  $\alpha = 0.05$  is 34% according to the binomial distribution, which confirms that the Type I error rate is below the  $\alpha$  nominal level. Under RCE, most of the iterations (92%) found at least one shifting clade by using *y-auto*.

With an incidence of rate shifts found at 0.159 (Table 3) and 12.2 clade in  $CC_{can}$  on average, the expected figure is 0.875. Finally, under CDE, the figure is higher still (96%) and this is to be explained by the very wide rate variation among  $CC_{can}$  clades under this scenario, that is two orders of magnitude larger than under BM (Appendix S1).

#### Virtual species' distributions: power and accuracy of phylo.niche.shift in detecting drifts in climatic niche evolution

The application of a drift was successfully recognised by phylo.niche.shift in most cases. A mere change in the mean value of the selected variables by some 15% (Table 4; Appendix S1, Figs. S1.1, S1.3, S1.6), under all scenarios and whether a reduction or an increase in variables values is imposed by applying d, is sufficient to achieve significance. Type I error is extremely low under both AE and RCE (Table 4). In contrast, under RCE, the drift is common even without imposing a drift (Table 4).

The precision of phylo.niche.shift obtained using automatic mode is high. The clades set to drift corresponds to 95% of the clades found to drift by using *yD-auto* under AE, 95% of the clades are found to drift in RCE. Under CDE, the figure is much lower (57%), which must be explained by the wide rate variation intrinsic to  $CC_{can}$  with this scenario.

For the simulations where d was selected to be either above 'd up' or below 'd down' (Table 4), the number of bioclimatic variables found to drift under AE by using yD-auto is 3.8 times larger than with y-node (Table 4; Fig. 3). The same figure under RCE is 2.7 (Appendix S1, Fig. S1.4).

Under the last CDE scenario, the number of bioclimatic variables found to drift was as low as 50% of the total number of variables (Appendix S1, Fig. S1.7), a figure consistently lower than with AE (80%) and RCE (87%, Table 4). When using non-drifted bioclimatic vectors (i.e. with *y*-node), the number of bioclimatic variables found to drift is in the 21% to 32% range (Table 4).

#### Climatic niche evolution in primates

By scanning the primate tree in search of climatic niche rate shifts, we found a negative shift pertaining to Strepsirrhini (rate difference = -37.76, *P*-value <<0.001).

**Table 4.** The percentage of drifts found as applied to the selected clade per different scenario when a drift is not applied (*y*-*nod*e), when it is applied (*y*-*nod*e) and when a rate shift is applied (*y*-*nod*e). 'd down' and 'd up' indicate the d value at which 95% of the phylo.niche.shift applications correctly identify the drift when a decrease or increase, respectively, in d is simulated. No drift is expected to occur at d = 1. Anticlimactic evolution – AE, random climatic evolution – RCE, climate-driven evolution – CDE

	AE	RCE	CDE
Incidence of drifts found, <i>y-node</i>	0.003	0.008	0.413
Incidence of drifts found, <i>yS-node</i>	0.366	0.406	0.55
Incidence of drifts found, <i>yD-node</i>	0.85	0.843	0.897
<i>d</i> down	0.84	0.84	0.87
d up	1.17	1.18	1.08
Average length of bioclimatic vector	8.66	8.69	8.73
Average number of drifting bioclimatic variables, <i>y-node</i>	1.841 (21.3%)	2.771 (31.8%)	2.194 (25.2%)
Average number of drifting bioclimatic variables, <i>yS-node</i>	1.820 (21.0%)	2.812 (32.4%)	2.322 (26.7%)
Average number of drifting bioclimatic variables, <i>yD-node</i>	6.942 (80.2%)	7.537 (86.7%)	4.456 (51.0%)



**Fig. 3.** The sensitivity of phylo.niche.shift to the intensity of the drift applied (*x*-axis), under the anticlimatic evolution scenario. On the *y*-axis, we report the logarithm of the Euclidean distance from the bioclimatic vector of means calculated over the entire tree, and the vector of means of the selected clade before (dots around the bottom dashed line) and after (dots forming the V-shaped distribution) multiplying the clade bioclimatic variables for a scalar *d*. Still on the *y*-axis, we also report the mean number of individual bioclimatic variables found to drift after (bottom dashed horizontal line) and before applying the *d* transform (top dashed horizontal line) vector of bioclimatic means. The broken solid lines connect the mean numbers of drifting bioclimatic variables calculated over 50 consecutive intervals in *d* after (top) and before (bottom) applying the *d* transform.

The clades including grey langurs and lutung (*Semnopithecus* and *Trachypithecus*) and capuchin monkeys (both robust and gracile, *Cebus* and *Sapajus*) both show significant and positive rate shifts (langurs+lutung: rate difference = 33.94, *P*-value = 0.998; capuchins: rate difference = 78.14, *P*-value = 1).

We found a significant niche drift in Catarrhini. Old World monkeys occupy more seasonal climates than other major primate clades, as testified by significantly higher than expected precipitation in the wettest month (BIO13), significantly lower than expected precipitation of the driest month (BIO14) and coldest quarter (BIO19), and low isothermality (BIO3). On the 365 species primate tree, phylo.niche.shift required 3.5s to run.

#### DISCUSSION

Phylogenetic groups of species differ widely in their realised climatic niche. These differences may result from selective changes promoting niche widening or its converse (Aagesen et al. 2016, Velasco et al. 2018, Mondanaro et al. 2020, Frost et al. 2022), or be essentially neutral with regards to the fundamental niche (Bonetti & Wiens 2014, Boucher et al. 2014, Saupe et al. 2019). Whether or not the fundamental niche changes, identifying such differences has a huge impact on our understanding of species diversity patterns (Rolland & Salamin 2016, Olalla-Tárraga et al. 2017, Rolland et al. 2018) of the link existing between climatic niche breadth and extinction risk (Wiens et al. 2019, Taheri et al. 2021) and of how climate change is impacting extinction and will continue to do so (Hanson et al. 2020, Trisos et al. 2020). Key to appreciating how species face environmental variation is understanding whether their climatic niche shifts (i.e. expands or retracts), drifts (i.e. moves along the niche dimension), or both. Although there is solid recognition that phylogenetic effects must be considered to understand these changes under a proper comparative context (Kellermann et al. 2012, Gutiérrez-Pesquera et al. 2016), most researchers just estimate the phylogenetic signal in bioclimatic variables, or rely on standard models of evolution such as BM (Münkemüller et al. 2015, Corro et al. 2021) or the Ornstein–Uhlenbeck process (Velasco et al. 2018, Frost et al. 2022). Yet, BM resides on assumptions that can be unrealistic for describing climatic niche evolution (Diniz-Filho et al. 2012, Kamilar & Cooper 2013), and even a significant signal does not indicate that the effect of shared ancestry on species' climatic niche is relevant (Freckleton & Jetz 2009, Cooper et al. 2011, Münkemüller et al. 2015). The Ornstein–Uhlenbeck process does not allow evolutionary rates to change across the tree, reducing to specified sets of purported evolutionary regimes.

We propose a new method to test for both niche shift and drift under a phylogenetically explicit scenario making no *a priori* assumptions about the rate and direction of niche evolution. The method, running under the R algorithm phylo.niche.shift, proved to be precise and very accurate, returning low Type I error rate and pervasively finding instances of rate shift (hence changes in the niche width) with 1.5-fold rate change (2.2-fold for rate decreases) and for drifts as low as ca. 15% displacement from the original niche position, regardless of the simulated scenario of niche evolution. phylo.niche.shift effectively finds 'global' drifts (i.e. those pertaining to the bioclimatic vectors as a whole) accurately and with very low error, although the incidence of false-positives among individual variables is large, suggesting caution against emphasising drifts into individual variables within the niche. We found remarkably high incidence of drifts under simulated shifts, and vice versa. This was expected, because a positive shift stretches the sampled bioclimatic niche space of clades to otherwise unexplored values, and suggests that instances of drifts and shifts should be investigated at once.

A number of methods developed to test niche shifts use the entire climatic variation experienced by the species within its range to represent its niche, and contrast the sampled climate against the background climate. The consideration of background climatic variability provides better delineation of the species' preferred conditions than presence-only data (Hengl et al. 2009, Broennimann et al. 2012). We relied on presence-only data, further reducing the sampled climates to bioclimatic vectors of average values. However, there is no formal impairment in deriving the bioclimatic vectors otherwise (i.e. by considering absence/pseudo-absence data). Furthermore, whereas the focus of such methods is studying niche shifts at the species level, phylo.niche.shift compares clades rather than species and accounts for phylogenetic effects. We deem the approach presented here to be best suited for largescale investigations testing macroevolutionary hypotheses about climatic niche evolution, such as whether a particular trait conferred greater environmental tolerance to a clade (e.g. C4 photosynthesis in grasses, Aagesen et al. 2016; or brown fat in mammals, Oelkrug et al. 2013); or whether niche width correlates with rates of taxonomic diversification (Rolland & Salamin 2016). Similarly, testing for climatic niche drifts with phylo.niche.shift means addressing whether trait evolution allows a clade to reside into a previously unexplored niche space (e.g. antifreeze proteins and life in Polar water fishes, Chen et al. 1997).

We addressed primate climatic niche evolution. We found that the clade including lutungs and the grey langur, and especially, the clade including capuchin monkeys proved to exploit unexpectedly large climatic niches. Gracile capuchins Cebus spp. are typical inhabitants of the wet and equable climates in the Amazon rainforest and forest savanna mosaic to the north (in the Roraïma area). In contrast, robust capuchins Sapajus spp. occur in cooler, drier, and more seasonal climates such as the Cerrado (a tropical savanna) and Caatinga (a xeric shrubland). Capuchins split into a robust and a gracile clade at the end of the Miocene, and Sapajus later greatly differentiated during the Pleistocene cooling (Lynch Alfaro et al. 2012), originating the great environmental variation (Appendix S1, Fig. S1.9) we found (Cáceres et al. 2014). The dichotomy between lutungs and grey langurs is very similar to the relationship between Cebus and Sapajus. Langurs Semnopithecus spp. in particular, inhabit exceptionally diverse habitats (from deserts to the Himalayas to dense forest habitats). Strong effect of habitat filtering (i.e. shared climatic preferences among species) is a common feature of strepsirrhine communities (Kamilar et al. 2014), testifying how narrow the climatic niche of strepsirrhine clades is compared with that of haplorrhine clades. Finally, the drift of Catarrhini as a whole is probably to be explained by their presence in highly seasonal climates in Eurasia, which is unparalleled by the other major primate clades. All these results indicate that phylo.niche.shift effectively found known, somewhat expected patterns of climatic niche change within primates.

In this study, we present a new R tool, phylo.niche. shift, meant to find instances of climatic niche shifts and drifts on phylogenetic trees. The tool proved exceptionally fast, accurate and sensible, under widely different scenarios of climatic niche evolution. phylo.niche. shift may help to improve research on niche evolution, as it allows users to find deviations from the expected niche among clades in the tree.

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#### DATA AVAILABILITY STATEMENT

Data and R codes are available at 10.5281/zenodo.6143630

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#### SUPPORTING INFORMATION

Additional supporting information may be found in the online version of this article at the publisher's website.

Appendix S1. Supplementary methods.