Supporting information for

The influence of vertebrate species traits on their responses to land-use and climate change

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1 Trait data compilation

1.1 Habitat affinities and broad specialisation

Habitat preferences. IUCN habitat data records habitat types in which species occur. Habitats are classified into 96 categories, which I pooled into 13 broader habitat variables: Forest, Savanna, Grassland, Shrubland, Wetland, Rocky areas, Caves and subterranean, Desert, Marine, Marine intertidal or coastal/supratidal, Artificial, Introduced vegetation and Other/Unknown. Species habitat preferences were described using these variables as binary (taking 1 if a species was known to occur in the habitat and 0 otherwise).

Habitat breadth. Habitat breadth was calculated as the number of habitats recorded to be used by a species in the IUCN database. Given that information regarding habitat suitability and habitat importance was also available in the IUCN data files, I used a weighted sum to calculate habitat breadth. Suitability was declined in three categories in the IUCN files: 'suitable', 'marginal' or 'unknown'. Habitats were recorded to be either of major importance, not of major importance or of unknown importance. I used the weights provided in Table 1 to produce weighted sums of the number of habitats used by each species. A comparison of the distribution of habitat breadths calculated with and without weights shows that weighting did not have a strong impact on the results (Figure 1).

Table 1: Weights used in the calculation of habitat breadth. Habitat breadth was calculated as the weighted sum of the number of habitats used by a species. Weights were assigned to each habitat given its importance and its suitability.

Suitability	Major importance						
Surtability	Yes	No	Unknown				
Suitable	1	0.5	1				
Marginal	0.3	0.3	0.3				
Unknown	1	0.3	1				

Degree of specialisation. A broad classification was adopted for species degree of specialisation. Using IUCN habitat files, I determined whether species were strictly natural habitat specialists or generalists. Generalists were species for which any habitat, suitable or of unknown suitability, was recorded to be artificial. Else, species were considered to be natural habitat specialists. When a habitat of an unknown type was considered suitable or was of unknown suitability, missing data was introduced.

1.2 Tackling taxonomic synonymy

1.2.1 Dropping replicated phylogenetic tips

2.6% of mammalian, 1.5% of avian, 1% of amphibian and 1.5% of reptilian species had multiple multiple phylogenetic positions after taxonomic corrections. Most of these species had only two replicates (as most species with synonyms were found to have two identified synonym, Figure 2.1 in main text). Table 2 provides the number of replicates across replicated species. Replicated tips corresponding to species described in the format *Genus cf.*, *Genus sp.*, *Genus spp.* or *Genus aff.* were all conserved in the phylogenies.

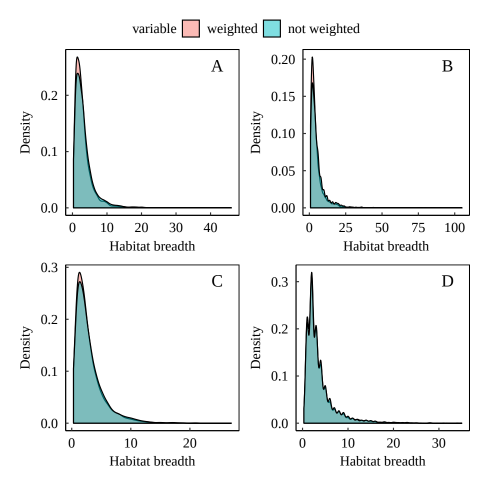


Figure 1: Distribution of habitat breadths across species for terrestrial vertebrates. The distribution is shown for calculations assigning weights to habitats according to their suitability and their importance (red surface area) or without weights (blue surface area). The distribution of habitat breadths slightly shifts to the left when a weighted sum is used (in **A**, **B** and **C**), as less importance is accorded to some habitats in the calculation (Table 1).

Figure 2 describes the procedure I used. If replicated tips were sister clades, the tip to conserve was chosen randomly among the replicates. Else, I chose to conserve the tree tip whose position was closest to the position of the same tip in the uncorrected tree, when present. In all other few cases, tips to drop were chosen randomly. See Figures 3, 4 and 5 for an example of each case.

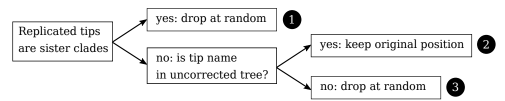


Figure 2: Procedure followed to drop replicated tips from phylogenies. Most redundant tip labels were replicated twice. (1) When replicated tips were sister clades, the tips to drop were chosen randomly, as it did not affect the 'true' phylogenetic position of the species; see case study 1 in Figure 3. (2) When replicated were not sister clades, I kept the tip whose position was closest to the position of the same tip in the uncorrected tree (Figure 4). (3) In a few cases, the corrected name did not appear in the original tree. Those were problematic cases, and the tips to drop were chosen randomly (Figure 5). Nevertheless, occurrences of that third case were rare.

Table 2: Number of replicated tip labels in phylogenies, number of which are sister clades and identified species for which dropping tips at random was problematic. Most redundant tips appeared twice. When replicated tips were sister clades, tips to drop were chosen randomly; this did not affect the species phylogenetic position. When replicated tips were not sister clades, I verified whether the corrected tip name appeared in the original, uncorrected tree. If so, I kept the tip in the corrected tree whose position was closest to the position of the tip in the original tree. Note that I used tip order in that case, which is sensitive to branch permutation. Finally, for replicated tip labels that were neither sister clades nor figuring in the original tree, tips to drop were chosen randomly. I identified these as problematic cases, although occurrences were rare. See Figure for a visual representation of each case.

Class	R	eplica	tes	Sister clades	In uncorrected tree	'Problematic'		
Class	2	3	>3	Sister clades	in uncorrected tree	Troblematic		
						Heterogeomys cherriei		
						Heterogeomys dariensis		
		8				$Hylopetes\ sagitta$		
Mammals	141		2	29	143	Marmosa paraguayana		
Wallimais	141			29	145	Neoromicia brunnea		
						Neoromicia malagasyensis		
						Plecturocebus discolor		
						Proechimys trinitatis		
						$Antrostomus\ arizonae$		
Birds	158	7	0	21	160	Calendulauda erythrochlamys		
Birds	156				100	Myiothlypis rivularis		
						Spermestes bicolor		
Reptiles	68	2	0	17	69	Salvator merianae		
Amphibians	41	4	2	8	44	$Lithobates\ berlandieri$		
Amphibians	41	4		0	14	Uperodon taprobanicus		

Case studies

Case study 1 (Figure 3): replicated tips are sister clades. In this example, Przewalski's toadhead agama (*Phrynocephalus przewalskii*) figures in the original, uncorrected tree under two names identified as being synonyms (*P. przewalskii* and *P. frontalis*). As such, two replicated tips with the same name appear in the tree after the taxonomic correction. As they are sister clades, one tip is randomly dropped in the final corrected tree. In that case, the 'true' phylogenetic position of the species is not affected.

Case study 2 (Figure 4): replicated tips are not sister clades, but the accepted name (tip label) figures in the original, uncorrected tree. In that example, the amphibian species Ambystoma californiense (California tiger salamander) was also found under its identified synonym A. tigrinum in the original tree. In the final tree, the tip closest to the position of the same tip in the uncorrected tree was kept.

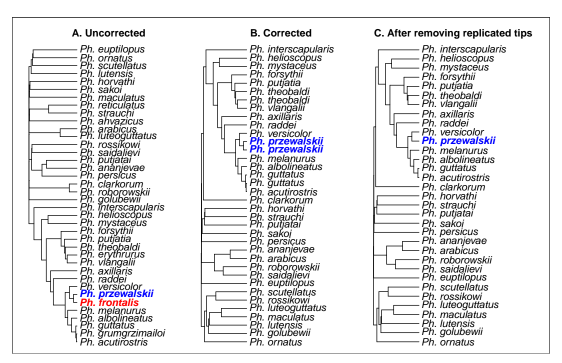


Figure 3: Phylogenetic trees before (A) and after (B) taxonomic correction, and after removing replicated tips (C) for species in the reptilian genus *Phrynocephalus*. The identified accepted name *P. przewalskii*, in blue (A), is identified as being a synonym for *P. frontalis*, which also figures in the uncorrected tree. As these clades are sister in the tree, one tip is dropped at random, and the final phylogenetic position of the species is not affected.

Case study 3 (Figure 5): replicated tips are not sister clades, and the accepted name (tip label) does not figure in the original, uncorrected tree. These were the problematic species listed in Table 2. In this example, the mammalian species *Petinomys sagitta* (arrow flying squirrel) and *Hylopetes lepidus* (gray-cheeked flying squirrel) have been identified to be synonymic, but the accepted name does not figure in the original tree. They are not sister clades; hence, one tip is randomly chosen to be dropped.

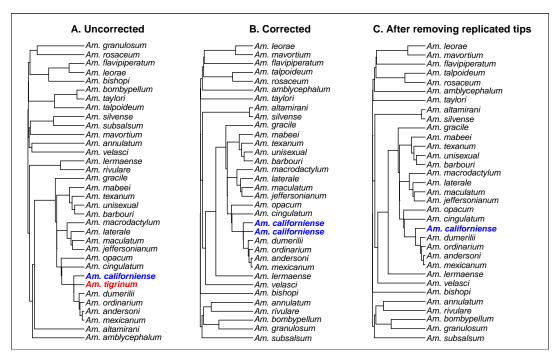


Figure 4: Phylogenetic trees before (A) and after (B) taxonomic correction, and after removing replicated tips (C) for species in the amphibian genus *Ambystoma*. The identified accepted name A. californiense, in blue (A), is identified as being a synonym for A. tigrinum, which also figures in the uncorrected tree. There are not sister clades; in the final tree, the tip closest to the position of the same tip in the uncorrected tree was kept.

- 2 Trait coverage and completeness for PREDICTS species
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- 3.1.4 Amphibians

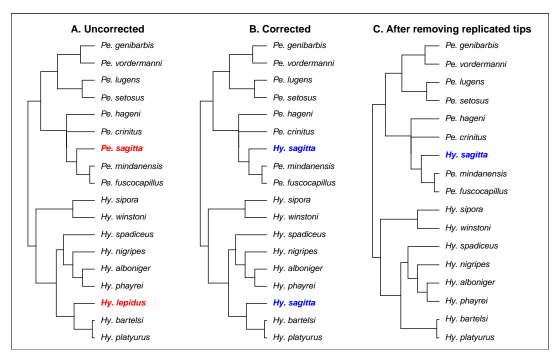


Figure 5: Phylogenetic trees before (A) and after (B) taxonomic correction, and after removing replicated tips (C) for species in the mammalian genera *Hylopetes* and *Petinomys*. The identified accepted name *H. sagitta*, in blue (B), does not figure in the uncorrected tree. It was identified to be the corresponding accepted name for *P. sagitta* and *H. lepidus*, which are not sister clades. One tip is randomly chosen to be dropped. Such cases were problematic but occurrences were rare (Table 2).

4 Missing values imputation

4.1 Phylogenetic signal

4.1.1 Significance

4.1.2 Phylogenetic signal computed with the original phylogenies

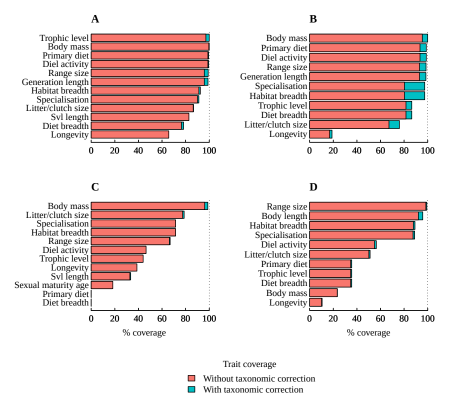


Figure 6: Trait coverage for species figuring in the PREDICTS database only.

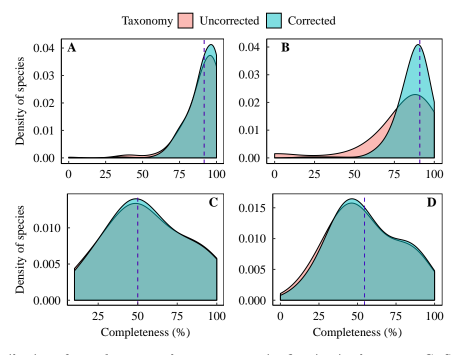


Figure 7: Distribution of completeness values across species figuring in the PREDICTS database only. Completeness for reptilian and amphibian species figuring in the trait dataset particularly improved compared to the completeness for the whole dataset.

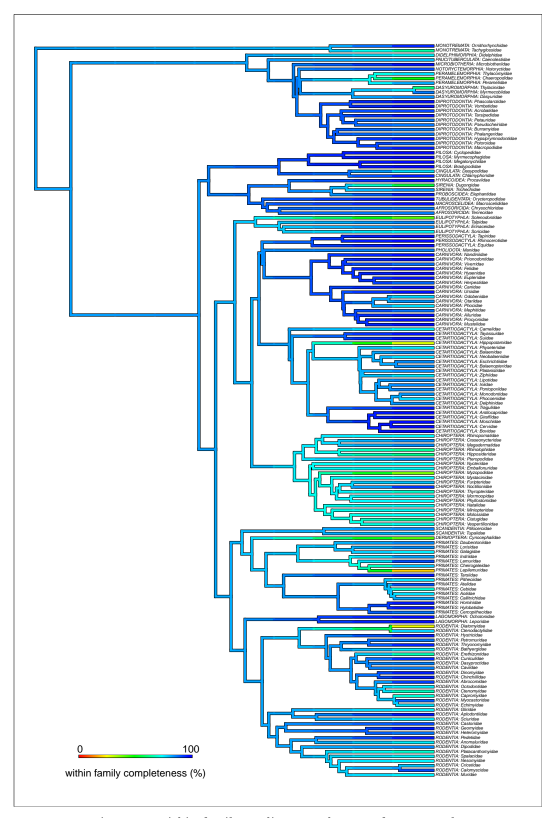


Figure 8: Within-family median completeness for mammals.

Table 3: Phylogenetic signal in continuous and categorical traits and in range size, calculated with original phylogenies. BM: body mass; L: longevity; LCS: litter/clutch size; HB: habitat breadth; DB: diet breadth; GL: generation length; BL: body length; SM: sexual maturity; RS: range size; TL: trophic level; PD: primary diet; DA: diel activity; Sp: specialisation. The phylogenetic signal in continuous traits was calculated with Pagel's λ . For categorical traits, the δ metric developed by Borges et al (2018) was used. A star indicates a significant signal (significant p-values scores for the log-likelihood ratio test in the case of λ ; and significant difference from the simulated null distribution of δ for categorical traits, see SI for test results). 'na' are introduced for traits that were not considered in a class but may have been used in another as a predictor in missing values imputations. All traits showed significant phylogenetic signal, with signals for BM, L, LCS, and GL being particularly strong in mammals and birds (above 0.9). Here all calculations were conducted with the original phylogenies.

	Continuous target traits,									Categorical traits:				
Class	additional predictors and range size: λ										δ			
	вм	L	LCS	нв	DB	\mathbf{GL}	BL	SM	RS	TL	PD	DA	Sp	
Mammals	1.0*	0.97*	0.96*	0.71*	0.99*	0.99*	1.0*	na	0.98*	16*	46*	17*	1.4*	
Birds	1.0*	0.89*	0.92*	0.48*	0.49*	0.99*	na	na	0.66*	10*	18*	33.103*	1.3*	
Reptiles	0.69*	0.79*	0.76*	0.40*	na	na	1.0*	0.86*	0.56*	4.4*	na	6.3*	1.8*	
Amphibians	0.72*	0.49*	0.58*	0.80*	0.52*	na	0.96*	na	0.62*	2.6*	3.1*	1.7*	2.3*	

4.2 Imputation robustness

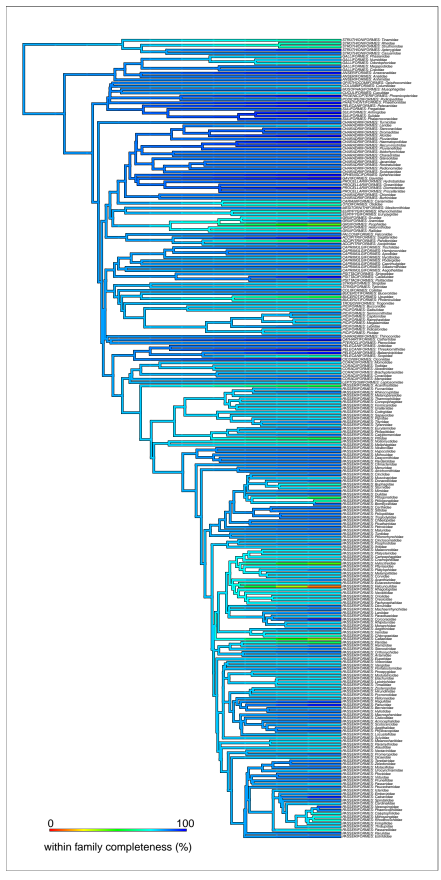


Figure 9: Within-family median completeness for birds.

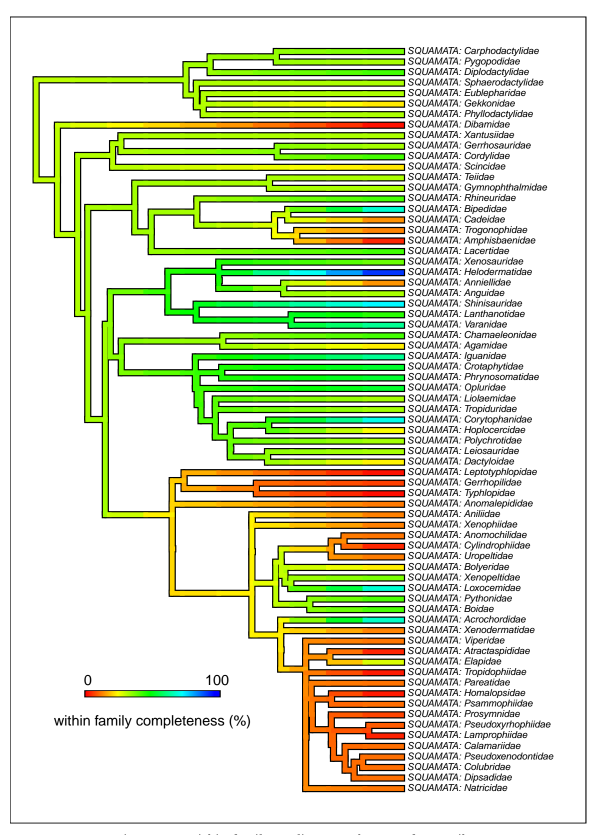


Figure 10: Within-family median completeness for reptiles.

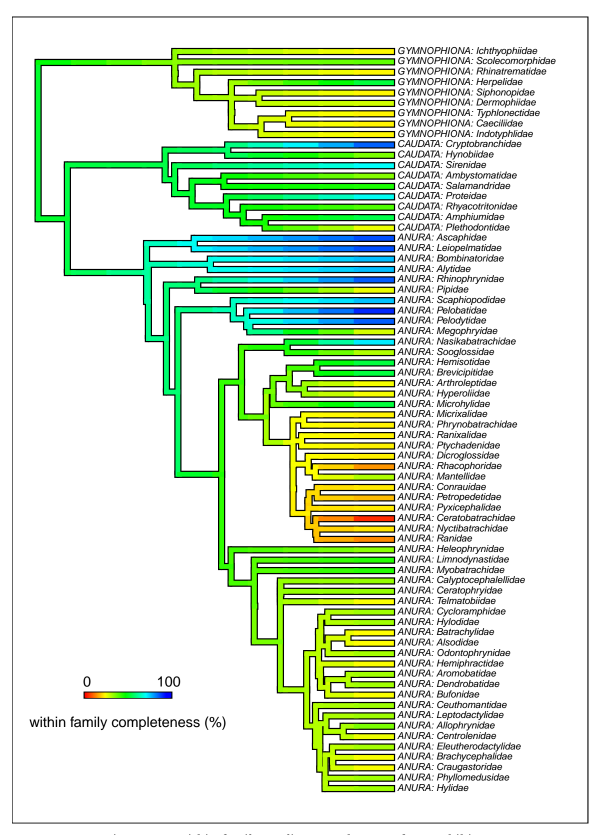


Figure 11: Within-family median completeness for amphibians.

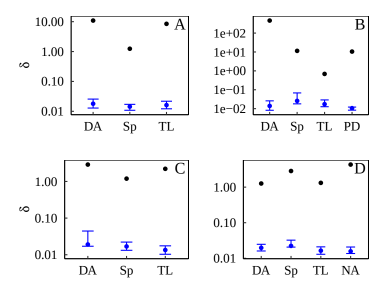


Figure 12: Observed and simulated phylogenetic signals in categorical traits. (A) Mammals; (B) birds; (C) reptiles; (D) amphibians. Black dots represent the observed phylogenetic signal in each categorical trait measured with δ , a metric developed by Borges et al (2018). The observed measured is compared to the null distribution of δ (phylogenetic signal calculated when trait values are randomised). Blue points represent the median value of null δ expectations, and the blue error bars are 95% confidence intervals. The null distribution of δ for each trait was generated by simulating 100 randomised trait vectors and calculating δ for each. As such, the median of null values represent the expected signal under a Brownian motion of trait evolution. Across all classes, all categorical traits presented significant phylogenetic signal. DA: diel activity; Sp: specialisation; TL: trophic level; PD: primary diet. Note that the phylogenetic signal of primary diet in mammals could not be calculated due to an implementation error.