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The relative importance of clade age, ecology and life-history traits in mammalian species richness

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Abstract

Extant mammalian clades have very different degrees of species richness, as some groups contain a very large number of species while others are represented by only one. The causes for this asymmetry remain undetermined. We gathered data for 14 variables classified in three groups: biogeography, ecology, and life-history. We also included clade age as an additional variable, to test the hypothesis that older clades, having had more time to diversify, contain more species. By using several online databases, we were able to include 4514 mammal species belonging to 1096 genera, 127 families, and 27 orders, making this, to our knowledge, the most comprehensive study on the drivers of mammalian species richness. Our analysis, which consisted in a generalized linear model, was conducted at three taxonomic levels: genus, family, and order. This allowed us to verify if the effect of a given variable is consistent regardless of the taxonomic hierarchical level. Our results suggest that clade age is not related to species richness regardless of the taxonomic level of analysis. We found that biogeography, namely, geographic range size and latitude were the most important variables, but with different effects. We found geographic range size to have a negative relationship with mammalian species richness, while latitude has a positive relationship. The intragroup variability of adult body mass was also important in some cases, although this might be due to the correlation of body mass with other variables. Generation length appears more important than litter size, but its weak signal makes drawing conclusions difficult. Ecological variables were the least important ones, with the sole exception of trophic level diversity at the genus level. However, it is possible that the ecological variables used did not capture well the most important ecological factors influencing clade species richness. Overall, our study points the major role of biogeography at influencing diversification rate in mammals.

Keywords: species richness, clade age, life-history, ecology, biogeography

Resumo

Existe uma grande disparidade na riqueza específica entre os vários grupos animais. A maioria das espécies encontra-se distribuída por um número reduzido de ordens, famílias e géneros, tendo a maior parte dos grupos taxonómicos poucas espécies extantes. A idade dos clados (ou grupos monofiléticos) é uma explicação clássica deste fenómeno, através da hipótese de que clados mais antigos terão tido mais tempo para sofrer processos de especiação e, por conseguinte, que o número de espécies nesses clados seja maior. Contudo, esta hipótese ignora ou minimiza o papel da extinção, que reduz o número de espécies, e que terá tido mais oportunidades de ocorrência em clados mais antigos. O número de espécies extantes resulta, portanto do balanço entre especiação e extinção, normalmente designada por taxa de diversificação, que naturalmente pode não ser constante no tempo. Vários trabalhos têm procurado explicar a assimetria e heterogeneidade na riqueza específica entre diferentes clados recorrendo a diversas abordagens, mas frequentemente estudando a relação entre riqueza específica e um pequeno número de variáveis. Não existe ainda um consenso claro sobre as variáveis mais importantes na determinação da riqueza específica dos clados, pois são conhecidos exemplos contraditórios para várias hipóteses, sendo o mais provável que uma multiplicidade de factores esteja envolvida em simultâneo.

Neste estudo investigámos o efeito e a importância relativa de variáveis que podem afectar a taxa de diversificação e, consequentemente, o número de espécies de mamíferos em grupos taxonómicos supraespecíficos. A classe dos mamíferos é ideal para um estudo desta natureza porque a sua história evolutiva é bastante bem conhecida, existe bastante informação para espécies de mamíferos, e o número de espécies é elevado o suficiente para permitir inferir padrões globais, mas sem ser tão numeroso que torne excessivamente complexo e difícil o manuseamento dos dados.

Ao contrário da maioria da literatura sobre o assunto, procurámos realizar uma análise abrangente e assim incluímos três grupos de variáveis que podem influenciar a riqueza específica de mamíferos. Obtivemos os dados a partir de várias bases de dados disponíveis online, para um total final em análise de 4154 espécies de mamíferos (pertencentes a 1096 géneros, 127 famílias e 27 ordens) filtradas de um total de 5156 espécies existentes disponíveis na base de dados de referência, PanTHERIA. Para além da idade de cada clado, incluímos:

- Variáveis biogeográficas: tamanho da área de distribuição e a sua variação, latitude e longitude;
- Características biológicas e da história de vida: número de indivíduos por ninhada, tempo de geração, e massa corporal dos adultos, assim como o coeficiente de variação de cada um;
- Variáveis ecológicas: diversidade de níveis tróficos, diversidade de dieta, diversidade de actividade, e diversidade de estratégias de forrageamento, representadas pelo índice de Shannon de cada categoria em cada nível taxonómico.

Recorremos então a um modelo linear generalizado. Utilizando uma abordagem de inferência multimodelo obtivemos as variáveis mais importantes de cada grupo, a incluir num modelo final com variáveis de todos os grupos. Por fim, calculámos da mesma forma a importância relativa de cada um dos factores incluídos no modelo final.

Para verificar se o efeito de cada um destes grupos de variáveis é consistente independentemente do nível taxonómico a que é feita a análise, o processo foi realizado ao nível do género, família e ordem, utilizando sempre que necessário a média de uma dada variável por grupo taxonómico, ou o índice de diversidade de Shannon no caso das variáveis ecológicas.

As variáveis biogeográficas constituem o grupo de variáveis mais importante que identificámos a qualquer um dos níveis taxonómicos. A latitude, o tamanho da área de distribuição e o seu coeficiente de variação apresentam valores altos de importância. Ao nível da ordem, apesar de apenas a latitude surgir no topo da importância, o seu índice é mais baixo e nenhuma variável é identificada como estatisticamente significativa. Isto sugere que o seu efeito será real a qualquer um dos níveis, mas que o aumento da variância associado ao nível da ordem pode tornar o sinal mais difícil de obter. O tamanho da área de distribuição está negativamente relacionado com a diversidade, o que indica que terá um efeito negativo na taxa de especiação ou positivo na taxa de extinção, apesar desta segunda hipótese nos parecer menos provável. Por sua vez, verificámos que a variabilidade do tamanho da área de distribuição está de distribuição poderão permitir a persistência de um maior número de espécies numa dada área geográfica, e a uma elevada importância da especiação alopátrica. O facto de grandes áreas de distribuição poderem conferir maior resistência à extinção, pode não compensar o seu potencial impacto negativo na especiação.

As características da história de vida estudadas aparentam ter uma influência muito pequena na riqueza específica. A variação da massa corporal aparenta ter um efeito, especialmente aos níveis da família e ordem. No entanto, a massa corporal está correlacionada com várias outras características intrínsecas

das espécies que podem afectar a especiação e a extinção. Um resultado interessante, por ser contrário a resultados descritos em trabalhos anteriores em mamíferos, é o tempo de geração aparentemente contribuir mais para a diversidade de mamíferos do que o número de crias por ninhada. Estas características são altamente variáveis entre grupos taxonómicos e as diferentes importâncias relativas entre estudos podem ser devidas a diferenças nos *taxa* estudados. Será talvez devido a estas diferenças que obtivemos resultados diferentes relativamente a outros estudos. Contudo, tendo em conta a muito maior escala a que este trabalho foi efectuado, concluímos que, a nível global, o tempo de geração é a variável estudada da história de vida que mais impacto tem na diversidade de mamíferos.

Dos três grupos de variáveis, o das variáveis ecológicas foi o menos importante. A única variável ecológica a apresentar alguma importância foi a diversidade do nível trófico ao nível do género. Noutros trabalhos, foi encontrada uma relação entre o nível trófico e a especiação em mamíferos, tendo-se verificado que as espécies herbívoras têm em geral uma maior taxa de especiação, e as omnívoras uma menor taxa de especiação. Sugerimos que a importância da diversidade trófica que verificámos no estudo possa ser o resultado de duas causas. Por um lado, a diversidade de nível trófico contribui para a redução de extinções causadas por grandes alterações abióticas e bióticas que afectem a disponibilidade de recursos em diferentes níveis tróficos. Por outro lado, a diversidade de nível trófico tenderá a diminuir a competição entre espécies do mesmo género, o que por sua vez deverá ter um impacto negativo nas taxas de extinção e positivo nas taxas de especiação.

Os nossos resultados confirmam, tal como anteriormente concluído por vários autores, que a idade dos clados não é um factor explicativo da sua riqueza específica. A taxa de diversificação não será, portanto, constante no tempo para os diversos clados de mamíferos.

Neste estudo desenvolvemos uma abordagem o mais abrangente e exaustiva possível para tentar identificar variáveis importantes na determinação da riqueza especifica de *taxa* a diferentes níveis hierárquicos dentro de grandes grupos taxonómicos. Identificámos uma elevada importância de variáveis biogeográficas, que se sobrepõe a todas as outras variáveis em qualquer dos níveis taxonómicos que analisámos, na riqueza específica em mamíferos. O nosso trabalho sugere que, apesar da importância de certos factores variar entre clados, a importância da biogeografia é dominante globalmente.

Palavras-chave: riqueza específica, idade de clados, histórias de vida, ecologia, biogeografia

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1. Introduction

The causes of species richness in each clade of life have long been sought after by evolutionary biologists. The species richness of a clade results from the balance of the effects of speciation and extinction rates. The difference between speciation and extinction rates is usually called diversification rate (Stadler 2011). A constantly positive diversification rate results in a net increase in extant richness, as extinction does not offset speciation. On the other hand, a negative diversification rate implies that extinction prevails over speciation resulting in a net decrease in species richness.

The diversification rate is different across clades. In fact, one of the most prevalent and obvious patterns in the tree of life is the unevenness in the number of species among higher *taxa*. For example, in mammals, some families are comprised of several hundred species, an extreme example being the Muridae with 730 species, but most families are not particularly diverse, with many having a single species. This is the case of the Orycteropodidae whose only extant member is the aardvark (*Orycteropus afer*). Intuitively, species richness could positively correlate with clade age, as older clades have had more time to undergo speciation events and thus diversify (McPeek and Brown 2007). However, this relationship is far from linear, and several examples directly contradict it, with old clades having few species. This may be due to an inherent decline in speciation rates with time or an increase in extinction rates, which will ultimately result in the extinction of a clade. Conversely, in younger clades, speciation and extinction rates may reach equilibrium after an initial burst of speciation, which will transiently maintain the net number of species constant through time.

What, then, explains such diverse patterns of species richness? The debate on the exact dynamics of diversification rates is far from settled, with theoretical and empirical studies supporting both the importance of clade age (McPeek and Brown 2007; Bloom et al. 2014) or its decoupling from species richness and its causes (Ricklefs 2007; Rabosky 2009; Rabosky et al. 2012; Verde Arregoitia et al. 2013). The latter studies suggest that there may be several variables at play shaping species richness. Several variables have been proposed, most of which we can classify into three groups: **ecological**, biological **life-history traits**, and **biogeographical**.

Given the contrasting conclusions of the aforementioned studies, in this investigation we aimed to determine the relative importance of clade age and of some variables of those variable groups, as well as the relative importance of individual variables, in mammalian species richness.

Ecological traits are specific traits related to resource acquisition and coping with the environment. Resource acquisition is extremely important for the success of individuals and populations, and it can therefore play a part in a species' vulnerability to extinction and probability to persist and speciate. For example, species at higher trophic levels are generally considered more sensitive to environmental change, and hence more likely to become extinct (Purvis et al. 2000). On the other hand, trophic specialization is known to correlate with diversification rate, with higher specialization leading to greater diversification (Price et al. 2012). Additionally, analyses of diet diversity may provide finergrained insights into the importance of trophic ecology and its influence on speciation and extinction rates.

Biological and life-history traits are variables related to intrinsic characteristics of species. Many of such variables have a potential effect on the abundance, and consequently on the effective population size, of a clade, and thus on diversification rates. Indeed, researchers have found life-history traits to

have such an impact (Isaac et al. 2005). A short generation length and a high number of offspring per litter and smaller body size are correlated with abundance (Marzluff and Dial 1991; White et al. 2007). Abundance is the number of individuals in a population, or, in this case, in a species (Smith and Smith 2011). High abundance should provide more opportunities for speciation. From a genetic perspective, species with larger effective population sizes are expected to be more genetically diverse (Frankham 1996), and a greater amount of standing genetic variation can facilitate ecological speciation (Schluter and Conte 2009). Body size is negatively linked to abundance and is known to have a complex relationship with the prediction of extinction (Cardillo et al. 2005). In contrast, life-history traits, such as litter size and age of sexual maturity, have been found to be good predictors of extinction risk (González-Suárez and Revilla 2013).

Biogeographic characteristics of a species' distribution, such as range size, have long been identified as having an effect in species richness across the globe (Ricklefs et al. 2007; Rabosky 2009). For instance, species richness varies along a latitudinal gradient, with the tropics possessing the highest diversity (Brown and Lomolino 1998; Pyron and Wiens 2013). Moreover, species with a larger distribution are likely to occupy a greater number of different habitats. Such species may be more resistant to environmental change and as such, more likely to persist. However, if we consider allopatric speciation to be the main, or one of the most important, modes of speciation, it can be argued that broad and continuous distributions of ecologically flexible species lowers speciation rates and cladogenesis. Nevertheless, Cardillo and colleagues (2003) tested such hypotheses using Australian mammals and found a positive correlation between range size and diversification rates. Other studies identified small range sizes as being correlated with increased extinction risks (Purvis et al. 2000; Manne and Pimm 2001). Biogeographical variables are also linked with resource availability in a number of ways, an example being higher net primary productivity near the tropics (Gillman et al. 2015).

In this study, we investigated several variables belonging to the three types of variables described above, to identify the most relevant ones in shaping species richness across higher taxonomic levels in mammals. We aimed to determine if any group of factors is particularly crucial in explaining species richness patterns, and for that purpose we used both taxonomically informed and non-informed generalized linear models. We further assessed whether results were consistent across taxonomic ranks by repeating the analyses at the genus, family, and order levels.

We focused on the Mammalia class for three different reasons. First, its evolutionary history is generally well understood (Beck et al. 2006). Second, it contains a large enough number of species to allow reliable results without being unwieldy. Finally, restricting the analysis to a single high-taxonomic level clade makes variables more comparable between species. Our study included 4514 mammalian species and 14 variables from three groups of predictors. To our knowledge, this is the first research assessing simultaneously the relative importance of traits related to species' ecology, biology and life history, and biogeography, as well as clade age, in determining species richness in mammals.

2. Materials and methods

2.1 Data acquisition

We assembled a dataset of variables pertaining to the three major groups of potential predictors discussed above. Some of the variables result from calculations on the raw data. See Table 2.1.1 for a full list of variables, their short description, and data source databases.

We scaled and standardized all continuous variables and log transformed the response variable: number of species per taxonomic group (González-Suárez and Revilla 2013).

We retrieved species data on litter size, adult body mass in grams, geographic range size in km², maximum and minimum latitude and longitude in degrees from the PanTHERIA database (Jones et al. 2009). PanTHERIA is the most widely used database for mammals, comprising information for 5416 species, and it was the primary data source for this work. We also gathered activity, foraging strategy, trophic level, body mass, and diet information from the EltonTraits database (Wilman et al. 2014). EltonTraits has high data coverage and relatively little data extrapolation (81% of diet data, 89% of activity data, and 96% foraging information are considered certain). Whenever possible, we completed body mass data missing in PanTHERIA with information from EltonTraits. We further obtained data on generation length from the database Generation Length for Mammals (Pacifici et al. 2013), and on trophic level from the MammalDIET database (Kissling et al. 2014), with the latter also used to complete missing values in the PanTHERIA dataset. Some of these databases use imputation or extrapolation to complete missing values, and the percentage of such cases is given in Table 2.1.2. We computed missing data percentages for all species and removed from the study those with 20% or more missing values. Then, we checked if any family had a missing data percentage greater than 10%; whenever this was the case, the taxon was removed from the dataset. A list of removed taxa is given in Table S 2.-Table S 3. It would have been interesting to include other variables, such as age of sexual maturity, weaning age, abundance/density, and number of litters per year, but missing data for these variables was too high.

Clade ages were obtained from the TimeTree database (Hedges et al. 2015). TimeTree consists of a compilation of dating studies, with clade age estimates usually calculated by several methods and using different data (molecular and/or fossil), and the final value for each taxon corresponds to the mean of all studies. Occasionally, TimeTree estimates are the mean from a small number of studies with wildly different results. Given this, we took the following approach: if the difference between the mean and median of all studies for a given taxon was over 10% of the mean, we used the age estimate from either a single study or the mean of two studies, whenever two were available. Molecular studies on a single taxon were preferred over class wide studies and supertrees and over fossil dating (see Supporting Information Table S 1 for information on the literature used). We did not include clade age information in the analysis at the genus level due to the high number of genera (1230 in PanTHERIA) and the impossibility of obtaining quality data for all.

For taxonomy, we followed Wilson and Reeder's (2005) Mammal Species of the World and the IUCN Red List of Threatened Species (IUCN 2016). We compared and checked the taxonomy of the used databases and corrected inconsistencies.

We ended with three datasets for analysis arranged by taxonomic level, with species grouped by genus, family, and order, respectively. With the exception of latitude and longitude, we summarized continuous variables for each clade by calculating mean values across species. For each taxonomic level, we examined whether the mean was an appropriate descriptor of central tendency for all variables by

visually inspecting the cumulative mean (using randomly sorted values in three randomized plots per variable) to confirm that the values generally converged to a constant value. We then compared these results with those using the same procedure but with the median. Means were judged as a better representation of the data than medians, and therefore used in the analyses (see Supporting Information Figure S 1). For latitude and longitude, we calculated the difference between the maximum and minimum values within each clade. We hypothesized that the mode of the body mass could provide insight into the diversification potential of a clade since, for a similar geographic range, significantly distinct values of body mass may entail substantially different speciation and extinction rates. We divided the range of adult body mass of each clade into four modal classes of equal size, and the mean of the most frequent modal class was calculated and included in the analyses.

To obtain a single value per clade for activity, diet, foraging strategy, and trophic level, we used the Shannon-Wiener diversity index. This method has been applied to compare diets among species (Vezzosi et al. 2014). We also included in the analyses the coefficient of variation $\left(CV = \frac{\sigma}{\mu}\right)$ of each clade for adult body mass, generation length, litter size, and geographic range size, to test the importance of variability in these variables.

Table 2.1.1. Groups of predictors and variables. Data sources databases are coded as follows: P - PanTHERIA; M - MammalDIET, E - EltonTraits; G - Generation Length for Mammals

Variable group	Code	Variable	Calculated statistic	Sources
	Act	Activity diversity	Shannon diversity index	P, E
	Diet	Diet diversity	Shannon diversity index	E
Ecology	TrL	Trophic level diversity	Shannon diversity index	Р, М
	FrgS	Foraging strategy	Shannon diversity index	Ε
	BdM	Adult body mass	Average	P, E
	BdM_CV	Adult body mass CV	Coefficient of variation	P, E
Biology	BdM_M	Adult body mass mode	Average	P, E
and	GnL	Generation length	Average	G
life-history	GnL_CV	Generation length coefficient of variation	Coefficient of variation	G
	LS	Litter size	Average	Р
	LS_CV	Litter size CV	Coefficient of variation	Р
	GRS	Geographic range size	Average	Р
	GRS_CV	Geographic range size CV	Average	Р
Biogeography	Lat	Latitude	Difference between highest and lowest latitude values in degrees of a given taxonomic group	Р
	Lon	Longitude	Difference between highest and lowest longitude values in degrees of a given taxonomic group	Р

Table 2.1.2. Data source databases with number of species, and missing data and extrapolation percentages in the variables used.

Database	Number of species	Variable	Missing data percentage	Extrapolation percentage
		Litter size	53.82%	0%
		Geographic range size	13.81%	0%
		Latitude	13.81%	0%
	5416	Longitude	13.81%	0%
PanihekiA	5416	Trophic level diversity	60.08%	0%
		Adult body mass	34.60%	0%
		Activity diversity	69.37%	0%
		Foraging strategy	0%	3.81%
EltonTraits	5400	Adult body mass	0%	25.17%
		Diet diversity	0%	19.41%
		Activity diversity	0%	10.98%
MammalDIET	5365	Trophic level diversity	0%	62.10%
Generation Length for Mammals	5427	Generation length	0%	76.74%

2.2 Model fitting and selection

We modelled the number of species per clade as a function of the different predictors considered using negative binomial Generalized Mixed Effects Models (GLMMs). Taxa sharing an evolutionary history are not independent (Felsenstein 1985). For this reason, the model was informed taxonomically by including as a random effect the taxonomic level above that being used in the analysis. In the analysis at the taxonomic level of Order, the Superorder was used as a random effect. Afterwards, the resulting models were compared with those obtained using simpler negative binomial Generalized Linear Models (GLMs), not taking into account the relationships of taxa at each taxonomic level. As justified below, the GLM models were used in the final analysis and interpretation.

Multicollinearity among predictors was evaluated by calculating the variance inflation factors (VIF) separately for each group of variables using the function "vif" from the package "car" (Fox and Weisberg 2011). VIF is a measure of the impact of collinearity in the observed regression coefficients. Zuur et al. (2010) state that values as high as 10 have been used as a threshold for assessing multicollinearity, but recommended a more conservative value of three. We calculated pairwise Pearson correlations between selected predictors to check for the presence of highly correlated variables; whenever such cases were found, we removed one variable from each pair. Overdispersion was checked by calculating the ratio between the residual deviance and the residual degrees of freedom. A ratio < 2 was considered acceptable. Authors generally consider a dispersion parameter between 0,5 and 2 to be acceptable (Logan 2010).

After fitting all variable groups, we performed automated model selection followed by model averaging using all models. The relative importance of each variable within each variable group was calculated using the functions "dredge", "model.avg" and "importance" from the package MuMIn (Barton 2016). Any variable with an importance value greater than 0.60 within its respective group was included in the final global model. This approach enabled us to select only the most important variables for the final models in order to keep these as parsimonious as possible. The automatic model selection output was used to calculate averages of the estimates from all models with $\Delta AICc \leq 2$. The 95% confidence intervals of the estimates were computed by bootstrapping the data 1000 times using the function "confint". This whole procedure (Fig. 1) was carried out at each of the three taxonomic levels considered. All analyses were done in R (R Core Team (R Foundation for Statistical Computing) 2015) using the function "glmer.nb" from the package lme4 (Bates et al. 2015) and the function "glm.nb" from the package MASS (Venables and Ripley 2002) to fit the GLMM and GLM, respectively.



Figure 2.1. Flowchart of the GLM data analysis.

3. Results

After removing 902 species with high levels of missing data, the final dataset consisted of a total of 4514 species from 1096 genera, 127 families, and 27 orders. This corresponds to a coverage of 83% of all mammal species included in PanTHERIA. Most species that were removed were marine mammals, for which ecological data are largely unavailable (see Table S 2 and Table S 3 for a list of the excluded taxa). Fifteen (12%) of the families included had either no clade age estimate available in TimeTree or the difference between mean and median was over 10%. In these cases, ages were manually obtained from publications, so that in the end we did not have clade age information for only three of the 127 families (see Table S 1 for the list of families and respective clade age references).

Species were asymmetrically distributed among supraspecific taxa. Approximately 25% of the families had more than 30 species, but most families ($\approx 66\%$) had less than 20 species. A similar pattern was also observed at the order level (Figure 3.1 and Figure 3.2).

Latitude and longitude were highly correlated (r > 0.8) at all taxonomic levels considered, and therefore longitude was discarded from the final analysis to prevent collinearity issues. Other variables also showed correlation, namely body mass and generation length ($r \approx 0.7$). This may be partly due to the extrapolation used in the 'Generation Length for Mammals' database, the level of which is very high in several taxonomic groups. Nevertheless, because of the potential importance of both variables, we decided to include both in the analysis. Body mass mode was discarded as it was highly correlated with mean body mass ($r \approx 0.99$).

Some evidence of multicollinearity was found for the life-history variables at the Order level. Nonetheless, VIF values were always < 5 and Pearson's correlations were not extreme (e.g. body mass vs. generation length: r = 0.76), and so we chose to include both in the analyses. We found no overdispersion at any taxonomic level, as the ratio between residual deviance and degrees of freedom were low or close to one. Some models, however, showed a certain degree of underdispersion. We opted to include underdispersed models because the risk is to increase the type II error rate but not the type I error rate.



Distribution of species richness across families

Figure 3.1. Histogram of species richness (n = 4514) across 127 mammalian families.



Distribution of species richness across orders

Figure 3.2. Histogram of species richness (n = 4514) across 27 mammalian orders.

All GLMMs yielded zero or close to zero random effect variance. On the other hand, GLMs exhibited identical estimates and standard errors to those from GLMMs. This indicates that, for all taxonomic levels considered, informing the models with taxonomy does not result in an increase in explanatory power. Hence, we decided to conduct the analyses without mixed models.

We constructed models separately for each variable group to select the most important variables (importance > 0.60) in each group. The estimated importance values and selected variables (importance values in bold) are given in Table 2.2.1. We then used the most important variables selected in the group-level analyses to fit the final global model for each taxonomic level. The variables and estimated importance values from these analyses are shown in Table 2.2.2.

Among the biogeographical variables analysed, latitude and the coefficient of variation of geographic range were selected for the final models at all taxonomic levels. Geographic range was selected at the genus and family levels (importance ≈ 1 in both cases), but not at the order level, despite its reasonable correlation with latitude. Among the biological and life-history traits, the coefficient of variation of body mass was the only one selected at all taxonomic levels, while generation length and the coefficient of variation of litter size were only selected at the Genus and Family level, respectively. Trophic level diversity was the most selected ecological trait, and the only among these that was selected at all taxonomic levels. Foraging strategy was selected at the Genus and Family level, while diet and activity were only selected at the Order and Genus level, respectively. The importance value of clade age was always low, never exceeding 0.40, and therefore it was not included in any of the final models.

We found that the group of biogeographical variables generally included those with the highest relative importance (Table 2.2.2 and Figure 3.3). This was particularly evident in the analyses at the genus and family levels, where the importance values for biogeographical variables are almost always equal or very close to one. At the Order level, the importance values were low overall, with latitude having the highest value at 0.65, followed by litter size CV with 0.58 (Table 2.2.2 and Figure 3.3 C). Variables in the 'biological and life-history traits' group generally ranked between those of the 'biogeographical' and 'ecological' groups. Variables in the former group ranged from very low importance (0.31: litter size CV at the Genus level) to relatively important (0.58: body mass CV at the Family level). The ecological variables generally ranked lowest in the final models, with the exception of trophic level diversity, which, with an importance value of 0.62, ranked higher than any biological and life-history trait at the Genus level (Table 2.2.2 and Figure 3.3 A). This pattern was not observed at the family and order levels, where ecological variables always ranked lowest with values never exceeding 0.45.

Variable group	Variable	Genus	Family	Order
	Activity diversity	0.289	0.445	0.197
	Diet diversity	0.268	0.428	0.511
Ecology	Trophic level diversity	1.000	0.990	0.741
	Foraging strategy	0.972	0.890	0.246
	Clade age	_	0.271	0.189
	Adult body mass	0.630	0.358	0.278
	Adult body mass CV	0.771	0.909	0.861
	Generation length	0.280	0.713	0.220
Biology and life–history	Generation length CV	0.444	0.274	0.266
	Litter size	0.310	0.287	0.209
	Litter size CV	0.859	0.315	0.218
	Clade age	_	0.391	0.167
	Geographic range size	1.000	0.969	0.349
Biogeography	Geographic range size CV	1.000	0.671	0.648
	Latitude	1.000	1.000	0.991
	Clade age	_	0.385	0.182

 Table 2.2.1. Importance values obtained in the separate analyses for each variable group at the different taxonomic levels.

Variable group	Variable	Genus	Family	Order
Ecology	Trophic level diversity	0.618	0.302	0.366
	Foraging strategy	0.263	0.436	_
	Adult body mass	0.448	_	_
Biology and life–history	Adult body mass CV	0.369	0.582	0.582
	Generation length	_	0.460	_
	Litter size CV	0.313	_	_
	Geographic range size	0.999	0.958	_
Biogeography	Geographic range size CV	0.999	0.532	0.525
	Latitude	1.000	0.997	0.654

Table 2.2.2. Importance values obtained in the final global models for the different taxonomic levels.







Figure 3.3. Relative importance plots for the variables in the final global models. The vertical bars represent the relative importance of the variables in the analyses at the genus (A), family (B), and order (C) levels.

Finally, we calculated the averages and 95% confidence intervals of the best models obtained with the automatic model selection procedure. The model averaging plots show the degree and direction of the effect of the most important variables on species richness (Figure 3.4-Figure 3.6). Variables that were important at more than one taxonomic level had a consistent effect direction between levels.

Latitude was found to have a positive effect on species richness at the three taxonomic levels studied, while geographic range size was found to have a negative effect at both the genus and family levels. Moreover, the average estimate values suggest a positive effect of the CV of geographic range size at all taxonomic levels, but except for the genus level the lower boundaries of the confidence intervals overlapped zero. Similarly, the results indicated a positive effect of the CV of body mass at all taxonomic levels, but only for orders did the confidence intervals not overlap zero. The average estimates also pointed to a positive effect of trophic level diversity across taxonomic levels, but the lower boundaries of the confidence intervals always overlapped zero. All other variables were selected in only one taxonomic level (genus: body mass and CV of litter size; family: generation length and foraging strategy diversity) and in all cases the confidence intervals overlapped zero.

Model averaging estimates at the genus level



Figure 3.4. Estimates obtained by averaging the best models ($\Delta AICc \leq 2$) at the genus level. Horizontal bars represent 95% confidence intervals.

Model averaging estimates at the family level



Figure 3.5. Estimates obtained by averaging the best models ($\Delta AICc \leq 2$) at the family level. Horizontal bars represent 95% confidence intervals.



Figure 3.6. Estimates obtained by averaging the best models ($\Delta AICc \leq 2$) at the order level. Horizontal bars represent 95% confidence intervals.

1. Discussion

The aim of this study was to investigate possible causes for differences in species richness between mammalian groups, both in general and at different supraspecific taxonomic levels.

Clade age might seem one of the most intuitive factors shaping species richness. Older clades have had more time for speciation and this could lead to a net increase in the number of species over time. While this hypothesis has been supported in previous research (McPeek and Brown 2007), it has also been contradicted by some studies (Ricklefs 2007; Rabosky 2009; Rabosky et al. 2012; Verde Arregoitia et al. 2013). Our results agree with the latter reports in suggesting that clade age is decoupled from species richness. Although, unfortunately, we could not include clade age in the Genus-level analysis, due to the lack of reliable estimates for many genera, clade age was not important enough to be included in the final models at the Family and Order levels.

Several other variables had a much greater influence on species richness than clade age. Variables in the biogeographical group, namely latitude and geographic range and its CV, generally showed high relative importance values from Genus to Order. The consistent high ranking of biogeographical variables regardless of taxonomic level indicates a high influence of biogeography on species richness. The high significance of latitude is unsurprising, given the long know latitudinal gradient of species richness (Brown and Lomolino 1998; Pyron and Wiens 2013). We found a higher latitude range to correlate with species richness. A higher latitude range may result in a higher probability of occupying the more species rich tropical region. The negative effect of geographic range size at the Family and Genus levels suggests that range size, instead of contributing to a positive diversification rate, may slow speciation rates to an extent not compensated for by its potential protective effect against extinction (Jablonski and Roy 2003, but see Gaston 1998 for a discussion on the hypotheses regarding relationship between speciation, extinction, and range size). At the evolutionary scale, allopatric speciation may be more important for species richness than resilience to extinction provided by ecological generalism. Moreover, the positive effect of the CV of geographic range size, albeit the 95% confidence interval did not overlap with zero only at the Genus level, may indicate that range size diversity is also a driver of species richness. However, this may not necessarily be the case since the positive association between the CV of geographic range size and species richness could simply reflect the fact that taxa with more species are likely to have a greater diversity of species' geographic range sizes. In addition, it is possible that groups currently having high CV in geographic range size include species that were mostly widely distributed historically but which distributions have variably shrunk due to human impact (Boivin et al. 2016), thereby leading to an overestimation of the historical range size diversity in many groups. Nevertheless, the CV of geographic range size may have an effect. The Genus-level dataset has more data points, each derived from a smaller number of species and as such one would expect genera to have inherently lower CV values. However, we could detect an important and significant effect of the CV of geographic range size on the species richness of genera, whereas in the Order- and Family-level analyses the significance of the effect of the CV of geographic range size was unclear. This may suggest that the signal of the effect of the CV of geographic range size is not a statistical artefact resulting from the influence of taxa with high species richness. The observed positive association between the CV of geographic range size and species richness may indirectly reflect the effect of other factors, such as biological and ecological traits, but is not necessarily a by-product of high species richness. Geographic ranges appear to be highly heritable (Hunt et al. 2005; Waldron 2007), including in mammals (Jones et al. 2005). This implies that, typically, related species may tend to have similar ranges and hence variation in range size within taxa

should be due to other processes (for example, historical, ecological, etc.) and not simply a function of the species richness of taxa.

Due to the lack of data for many species, we unfortunately could not include dispersal ability as a variable in our study. High dispersal capacity may allow the exploration of a wide range of different habitats, and this could promote speciation, but low dispersal ability should lead to increased rates of allopatric speciation. The CV of geographic range size might be expected to be inversely correlated with dispersal capabilities. The negative effect of geographic range size together with the positive effect of the CV of geographic range size can be interpreted as indicating that susceptibility to vicariance, ecological or habitat specialism, and limited dispersal are the most influential factors increasing the net diversification rate (Bohonak 1999; Claramunt et al. 2012; Salisbury et al. 2012).

The biological and life history traits we studied appear to play a much smaller role than the biogeographical variables in shaping species richness. This is an interesting result since the previous most comprehensive analysis of the correlates of species richness in mammals (Isaac et al. 2005) mostly focused on biological and life history traits. That study, conducted at the Order level and examining four orders (Carnivora, Chiroptera, Marsupialia, and Primates), found species richness to be significantly correlated with litter size, gestation period and interbirth interval, while in our investigation neither litter size nor generation length were relatively important or statistically significant variables affecting species richness. Like ours, their study found no evidence that mammalian species richness is associated with adult body mass, but importantly we obtained some support for a positive effect of the CV of body mass, a variable they did not consider that is correlated with traits that may influence species richness (Cardillo et al. 2005, 2008). Conversely, Isaac et al. (2005) found some support for a correlation between species richness and abundance, a variable that we would have liked to have included in our analysis but for which data is lacking for many species. The different findings of the two studies are unsurprising given the differences in the number and nature of the variables considered, number and taxonomic level of the taxa analysed, sample sizes, and statistical approaches for data analysis (GLM vs. phylogenetically independent contrasts). For example, Isaac et al. (2005) did not use summary statistics of species data to characterize and represent supraspecific taxa as we did, but instead conducted their analyses at the species level. On the other hand, they analysed relatively narrow sets of taxa and variables and did not evaluate the importance and effects of variables at different taxonomic levels. Nevertheless, both studies support the idea that a host of factors influence species richness and that the major determinants may vary between taxonomic groups. For instance, Isaac et al. (2005) found that species richness is correlated with shorter gestation period in the carnivores and larger litter size in marsupials, while abundance correlates positively with species richness in primates and negatively in microchiropterans. Our Orderlevel analysis, where fewer variables were selected for the final model and their relative importance never exceeded 0.65, suggests that analyses at higher taxonomic levels may be confounded by high variability within taxa and may have difficulties in identifying important influencing factors.

The results for the ecological variables were remarkable in the sense that their importance was generally low. It is important to note, however, that a range of potentially important ecological variables could not be included due to a lack of data. Moreover, the ecological variables were coded in such a way that only the effect of the variable's variation, and not the presence of a particular variable's value, could be assessed. The only relatively important ecological variable was trophic level diversity, for which the final models indicated a positive effect on species richness at all taxonomic levels, albeit with weak statistical support. The importance of trophic level diversity may stem from two different causes. Firstly, trophic level diversity should help to reduce competition for resources between related species, especially those congeneric because, as mentioned above, they may tend to inhabit the same geographical regions (Hunt et al. 2005; Waldron 2007). Secondly, trophic level diversity may provide greater resilience to environmental and habitat changes, and thereby decrease extinction rates.

It should be noted that current geographic ranges can vary wildly from the ones found throughout most of a given species' evolutionary history, particularly given the impact of anthropogenic factors starting at the first human migrations up to the industrial revolution (Boivin et al. 2016). In fact, it has been found that accounting for the differences between historical and current ranges sizes yields different results when predicting extinction risk (Hanna and Cardillo 2013). This may be due to the different factors driving extinction currently, as opposed to the time before humans had a such a drastic impact on the environment, which could mean that current geographic ranges are a consequence and not a cause of extinction (Hanna and Cardillo 2013). Incorporating historical ranges will be required in future work. However, to dismiss that current biogeographical patterns are, at least in part, linked to a species' evolutionary history is impossible, as it must have been influenced by dispersal and by climatic events. The problem persists in how to acquire and incorporate historical biogeographical data. The uncertainty of such inferences can introduce unwanted variance which may end up hindering our understanding of speciation and extinction patterns. However, ignoring historical biogeography potentially misrepresents the effect of these variables on speciation and extinction rates.

The concept of species also raises important challenges on diversity and speciation/extinction studies that must be recognized. Most works on species richness use the number of species per taxonomic group, geographic location, or a similar grouping factor. However, the number of species can vary wildly according to which species concept is used (Hey 2001) and as more species are discovered and described. In 19 years (1982-2001), the number of described primate species grew by more than twofold (Isaac and Purvis 2004). To circumvent this issue we used standard databases which follow a popular reference taxonomy (Wilson and Reeder 2005). This provides a good degree of comparability with other publications on this topic. However, while comparisons between studies are possible, the true number of extant species is ever changing (as more species are discovered and taxonomy is refined), and depends on the particular species concept used.

On the same topic, count creep, that is, the revision of a previously described species into two or more new species, is another concern. Count creep does not occur randomly, being more frequent among emblematic species or the ones more easily available to systematists (Isaac and Purvis 2004), which can lead to a skewed perception of richness. Isaac and Purvis (2004) verified that there is no impact in regression slopes of several ecological and life history traits with primate taxonomy changes, but did find an effect on the significance values of their results. This means that the possibility of incorrectly interpreting the dependent variable is possible, especially by diluting the effect of factors which contribute only slightly to richness.

Finally, one should bear in mind that the analyses at the Order level are inherently more artificial. We expected a much weaker signal at this level, since each taxon is comprised of a higher number species, and possesses more within group diversity. However, a sufficiently strong effect of a particular variable may still be identified. The inclusion of data surmised at the Genus level attempts to further assess impact of intragroup variability, as well as confirm the general trends found in model fitting. We confirm that this method is applicable to identify variables with a very strong effect, as we encountered for biogeography.

The ambitious scope of the work, which included more than 80% of the species present in the PanTHERIA database and incorporated a wealth of different variables allows us confidently point to biogeography as the most important factors in predicting mammal species richness. Our approach

adequately identifies variables that are relevant across all levels of analysis. Similarly, it hints to other possibly more elusive variables, such as generation length. Further studies should focus on disentangling and measuring the effect of the weaker variables across mammal orders, to better quantify and understand why they are important only in some cases. Whenever possible, phylogenetic information should be included for dependence correction.

It must be noted that we used several databases to ensure that our data were as complete as possible, which raises some concern. On the one hand, a higher amount of data should lead to more accurate results, since we are able to sample from a much broader range of mammals when compared with previous studies on the subject. On the other hand, several of the databases we used to complement PanTHERIA use imputation, that is, several species' entries do not result from direct observation retrieved from the literature, but instead are extrapolations from related species. Since missing data in such databases does not occur at random, the validity of conclusions drawn from imputed values has been questioned (González-Suárez et al. 2012). Researchers have shown that predicting extinction risk from imputation heavy data leads to biased results. Due to time constraints, we assumed imputed data, resulting in a smaller, but more reliable dataset. In addition, the effect of missing data should be explored, to ensure that by removing imputed data we are not introducing bias but in a different way. We could asses this by randomly removing variables from the final dataset to gauge the effect of missing values.

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Supporting Information



Cumulative Means of Muridae

Figure S 1-12. Example of cumulative means and median plot of variables from the Muridae family.

Cumulative Medians of Muridae



Cumulative Means of Muridae



Cumulative Medians of Muridae



Cumulative Means of Muridae



Cumulative Medians of Muridae



Cumulative Means of Muridae



Cumulative Medians of Muridae



Cumulative Means of Muridae



Cumulative Medians of Muridae



Cumulative Means of Muridae



Cumulative Medians of Muridae



Table S 1. Data sources for Families with poor data on TimeTree.

Family	Reference 1	Reference 2
Rhinocerotidae	Steiner, C. C., & Ryder, O. a. (2011). Molecular phylogeny and evolution of the Perissodactyla. <i>Zoological Journal of the</i> <i>Linnean Society</i> , <i>163</i> (4), 1289–1303. http://doi.org/10.1111/j.1096- 3642.2011.00752.x	
Tapiridae	Steiner, C. C., & Ryder, O. a. (2011). Molecular phylogeny and evolution of the Perissodactyla. <i>Zoological Journal of the</i> <i>Linnean Society</i> , <i>163</i> (4), 1289–1303. http://doi.org/10.1111/j.1096- 3642.2011.00752.x	
Bradypodidae	Gibb, G. C., Condamine, F. L., Kuch, M., Enk, J., Moraes-, N., Superina, M., Delsuc, F. (2015). Shotgun mitogenomics provides a reference phylogenetic framework and timescale for living Xenarthrans. <i>Molecular</i> <i>Biology and Evolution</i> .	Delsuc, F., Vizcaíno, S. F., & Douzery, E. J. P. (2004). Influence of Tertiary paleoenvironmental changes on the diversification of South American mammals: a relaxed molecular clock study within xenarthrans. <i>BMC Evolutionary</i> <i>Biology</i> , <i>4</i> , 11. http://doi.org/10.1186/1471- 2148-4-11
Cyclopedidae	Gibb, G. C., Condamine, F. L., Kuch, M., Enk, J., Moraes-, N., Superina, M., Delsuc, F. (2015). Shotgun mitogenomics provides a reference phylogenetic framework and timescale for living Xenarthrans. <i>Molecular</i> <i>Biology and Evolution</i> .	Delsuc, F., Vizcaíno, S. F., & Douzery, E. J. P. (2004). Influence of Tertiary paleoenvironmental changes on the diversification of South American mammals: a relaxed molecular clock study within xenarthrans. <i>BMC Evolutionary</i> <i>Biology</i> , <i>4</i> , 11. http://doi.org/10.1186/1471- 2148-4-11

Abrocomidae	Opazo, J. C. (2005). A molecular timescale for caviomorph rodents (Mammalia, Hystricognathi). <i>Molecular Phylogenetics</i> <i>and Evolution</i> , <i>37</i> (3), 932–937. http://doi.org/10.1016/j.ympev.2005.05.002	Verzi, D. H., Olivares, A. I., Morgan, C. C., & Álvarez, A. (2015). Contrasting Phylogenetic and Diversity Patterns in Octodontoid Rodents and a New Definition of the Family Abrocomidae. <i>Journal of</i> <i>Mammalian Evolution</i> . http://doi.org/10.1007/s10914- 015-9301-1
Chinchillidae	Opazo, J. C. (2005). A molecular timescale for caviomorph rodents (Mammalia, Hystricognathi). <i>Molecular Phylogenetics</i> <i>and Evolution</i> , <i>37</i> (3), 932–937. http://doi.org/10.1016/j.ympev.2005.05.002	
Cricetidae	Steppan, S., Adkins, R., & Anderson, J. (2004). Phylogeny and Divergence-Date Estimates of Rapid Radiations in Muroid Rodents Based on Multiple Nuclear Genes. Systematic Biology, 53(4), 533–553. http://doi.org/10.1080/10635150490468701	
Cuniculidae	C. M. Voloch, J. F. Vilela, L. Loss-Oliveira, C. G. Schrago, Phylogeny and chronology of the major lineages of New World hystricognath rodents: insights on the biogeography of the Eocene/Oligocene arrival of mammals in South America. <i>BMC Res.</i> <i>Notes.</i> 6 , 160 (2013).	
Dinomyidae	Opazo, J. C. (2005). A molecular timescale for caviomorph rodents (Mammalia, Hystricognathi). <i>Molecular Phylogenetics</i> <i>and Evolution</i> , <i>37</i> (3), 932–937. http://doi.org/10.1016/j.ympev.2005.05.002	N. S. Upham, B. D. Patterson, Diversification and biogeography of the Neotropical caviomorph lineage Octodontoidea (Rodentia: Hystricognathi). <i>Mol.</i> <i>Phylogenet. Evol.</i> 63 , 417–429 (2012).
Dipodidae	Q. Zhang <i>et al.</i> , Tracing the Origin and Diversification of Dipodoidea (Order: Rodentia): Evidence from Fossil Record and Molecular Phylogeny. <i>Evol. Biol.</i> 40 , 32–44 (2013).	Q. Zhang <i>et al.</i> , Tracing the Origin and Diversification of Dipodoidea (Order: Rodentia): Evidence from Fossil Record and Molecular Phylogeny. <i>Evol.</i> <i>Biol.</i> 40 , 32–44 (2013).

Muridae	Steppan, S., Adkins, R., & Anderson, J. (2004). Phylogeny and Divergence-Date Estimates of Rapid Radiations in Muroid Rodents Based on Multiple Nuclear Genes. Systematic Biology, 53(4), 533–553. http://doi.org/10.1080/10635150490468701
Myocastoridae	J. C. Opazo, A molecular timescale for caviomorph rodents (Mammalia, Hystricognathi). <i>Mol. Phylogenet. Evol.</i> 37 , 932–937 (2005).
Petromuridae	 H. M. Sallam, E. R. Seiffert, M. E. Steiper, E. L. Simons, Fossil and molecular evidence constrain scenarios for the early evolutionary and biogeographic history of hystricognathous rodents. <i>Proc. Natl. Acad.</i> <i>Sci. U. S. A.</i> 106, 16722–7 (2009).
Thryonomyidae	 H. M. Sallam, E. R. Seiffert, M. E. Steiper, E. L. Simons, Fossil and molecular evidence constrain scenarios for the early evolutionary and biogeographic history of hystricognathous rodents. <i>Proc. Natl. Acad.</i> <i>Sci. U. S. A.</i> 106, 16722–7 (2009).

Table S 2. Species and corresponding families removed from the analysis due to missing data. Names in blue indicate the family corresponding to the species listed below.

Abrocomidae	Ctenomyidae	Muridae	Phocoenidae
Abrocoma budini	Ctenomys bergi	Acomys airensis	Neophocaena phocaenoides
Abrocoma famatina	Ctenomys brasiliensis	Acomys chudeaui	Phocoena dioptrica
Abrocoma shistacea	Ctenomys budini	Acomys cineraceus	Phocoena phocoena
Abrocoma uspallata	Ctenomys coludo	Acomys johannis	Phocoena sinus
Abrocoma vaccarum	Ctenomys coyhaiquensis	Acomys seurati	Phocoena spinipinnis
Cuscomys ashaninka	Ctenomys dorbignyi	Aethomys ineptus	Phocoenoides dalli
Cuscomys oblativa	Ctenomys famosus	Apomys gracilirostris	Phyllostomidae
Aotidae	Ctenomys flamarioni	Archboldomys musseri	Carollia colombiana
Aotus hershkovitzi	Ctenomys fochi	Arvicanthis ansorgei	Carollia sowelli
Atelidae	Ctenomys fodax	Arvicanthis rufinus	Micronycteris homezi
Alouatta coibensis	Ctenomys fulvus	Batomys russatus	Phyllonycteris major
Balaenidae	Ctenomys johannis	Bullimus gamay	Platyrrhinus recifinus
Balaena mysticetus	Ctenomys juris	Conilurus albipes	Vampyressa thyone
Eubalaena australis	Ctenomys lami	Coryphomys buehleri	Physeteridae
Eubalaena glacialis	Ctenomys mendocinus	Crateromys heaneyi	Kogia breviceps
Eubalaena japonica	Ctenomys osvaldoreigi	Crunomys suncoides	Kogia sima
Balaenopteridae	Ctenomys pilarensis	Dasymys alleni	Physeter catodon
Balaenoptera acutorostrata	Ctenomys pundti	Dasymys cabrali	Pitheciidae
Balaenoptera bonaerensis	Ctenomys roigi	Dasymys rwandae	Callicebus bernhardi
Balaenoptera borealis	Ctenomys scagliai	Dasymys sua	Callicebus discolor

Balaenoptera edeni	Ctenomys tulduco	Deomys ferrugineus	Callicebus lucifer
Balaenoptera musculus	Ctenomys viperinus	Dephomys eburneae	Callicebus lugens
Balaenoptera physalus	Ctenomys yolandae	Desmomys yaldeni	Callicebus purinus
Megaptera novaeangliae	Dasypodidae	Echiothrix centrosa	Callicebus regulus
Bathyergidae	Calyptophractus retusus	Echiothrix leucura	Callicebus stephennashi
Cryptomys amatus	Dasypus yepesi	Gerbillus bottai	Chiropotes chiropotes
Cryptomys anselli	Dasyuridae	Gerbillus dongolanus	Chiropotes israelita
Cryptomys kafuensis	Murexechinus melanurus	Gerbillus garamantis	Chiropotes utahickae
Bovidae	Paramurexia rothschildi	Gerbillus harwoodi	Platanistidae
Alcelaphus caama	Phascomurexia naso	Gerbillus jamesi	Platanista gangetica
Alcelaphus lichtensteinii	Sminthopsis boullangerensis	Gerbillus lowei	Platanista minor
Bos frontalis	Delphinidae	Gerbillus mackillingini	Potoroidae
Bos grunniens	Cephalorhynchus commersonii	Gerbillus maghrebi	Caloprymnus campestris
Bos taurus	Cephalorhynchus eutropia	Gerbillus mauritaniae	Potorous platyops
Bubalus bubalis	Cephalorhynchus heavisidii	Gerbillus mesopotamiae	Prolagidae
Capra hircus	Cephalorhynchus hectori	Gerbillus rupicola	Prolagus sardus
Cephalophus brookei	Delphinus capensis	Gerbillus somalicus	Pteropodidae
Damaliscus korrigum	Delphinus delphis	Gerbillus stigmonyx	Dobsonia chapmani
Damaliscus superstes	Feresa attenuata	Gerbillus syrticus	Dobsonia magna
Eudorcas rufina	Globicephala macrorhynchus	Gerbillus vivax	Epomophorus minor
Gazella erlangeri	Globicephala melas	Gerbillus zakariai	Paranyctimene tenax

Gazella saudiya	Grampus griseus	Grammomys kuru	Pteralopex acrodonta
Hemitragus hylocrius	Lagenodelphis hosei	Grammomys poensis	Pteropus brunneus
Hemitragus jayakari	Lagenorhynchus acutus	Hadromys humei	Pteropus loochoensis
Hippotragus leucophaeus	Lagenorhynchus albirostris	Hadromys yunnanensis	Pteropus pilosus
Ovis aries	Lagenorhynchus australis	Hydromys habbema	Pteropus subniger
Saiga borealis	Lagenorhynchus cruciger	Hydromys shawmayeri	Pteropus tokudae
Taurotragus derbianus	Lagenorhynchus obliquidens	Hylomyscus grandis	Rousettus linduensis
Taurotragus oryx	Lagenorhynchus obscurus	Leporillus apicalis	Rhinolophidae
Calomyscidae	Lissodelphis borealis	Limnomys bryophilus	Rhinolophus cornutus
Calomyscus grandis	Lissodelphis peronii	Lophuromys angolensis	Rhinolophus imaizumii
Camelidae	Orcaella brevirostris	Lophuromys ansorgei	Rhinolophus monoceros
Camelus bactrianus	Orcinus orca	Lophuromys aquilus	Rhinolophus siamensis
Camelus dromedarius	Peponocephala electra	Lophuromys brevicaudus	Rhinolophus ziama
Lama glama	Pseudorca crassidens	Lophuromys brunneus	Sciuridae
Canidae	Sotalia fluviatilis	Lophuromys dieterleni	Ammospermophilus insularis
Dusicyon australis	Sousa chinensis	Lophuromys dudui	Prosciurillus rosenbergii
Vulpes lagopus	Sousa teuszii	Lophuromys eisentrauti	Solenodontidae
Capromyidae	Stenella attenuata	Lophuromys huttereri	Solenodon arredondoi
Capromys gundlachianus	Stenella clymene	Lophuromys roseveari	Solenodon marcanoi

Geocapromys thoracatus	Stenella coeruleoalba	Lophuromys verhageni	Soricidae
Hexolobodon phenax	Stenella frontalis	Lophuromys zena	Anourosorex assamensis
Isolobodon montanus	Stenella longirostris	Malpaisomys insularis	Anourosorex schmidi
Isolobodon portoricensis	Steno bredanensis	Mastomys awashensis	Blarina peninsulae
Mesocapromys melanurus	Tursiops aduncus	Melomys bannisteri	Blarinella griselda
Mesocapromys sanfelipensis	Tursiops truncatus	Melomys caurinus	Blarinella quadraticauda
Plagiodontia araeum	Didelphidae	Melomys cooperae	Chodsigoa caovansunga
Plagiodontia ipnaeum	Caluromys lanatus	Melomys frigicola	Chodsigoa parva
Rhizoplagiodontia lemkei	Didelphis pernigra	Melomys fulgens	Chodsigoa sodalis
Caviidae	Gracilinanus agricolai	Melomys howi	Congosorex verheyeni
Cavia intermedia	Gracilinanus formosus	Melomys matambuai	Crocidura baluensis
Cavia porcellus	Gracilinanus ignitus	Melomys paveli	Crocidura brunnea
Kerodon acrobata	Hyladelphys kalinowskii	Melomys spechti	Crocidura caspica
Cebidae	Marmosa quichua	Melomys talaudium	Crocidura foetida
Callithrix aurita	Marmosops bishopi	Meriones grandis	Crocidura gmelini
Callithrix humilis	Marmosops dorothea	Micaelamys granti	Crocidura hilliana
Callithrix kuhlii	Marmosops incanus	Micaelamys namaquensis	Crocidura hutanis
Callithrix leucippe	Marmosops juninensis	Mus famulus	Crocidura ichnusae
Leontopithecus caissara	Marmosops neblina	Mus fragilicauda	Crocidura indochinensis

Leontopithecus	Marmosops paulensis	Mylomys rex	Crocidura
chrysopygus			jouvenetae
Saguinus graellsi	Marmosons ninheiroi	Myotomys sloggetti	Crocidura katinka
Sagamas graensi	Marinosops princitor	wyotomys stoggetti	Crocidara Katilika
Saguinus imperator	Micoureus phaeus	Nilopegamys plumbeus	Crocidura lepidura
Saguinus inustus	Monodelphis glirina	Notomys amplus	Crocidura malayana
Saguinus martinsi	Monodelphis palliolata	Notomys longicaudatus	Crocidura musseri
Saguinus melanoleucus	Monodelphis umbristriata	Notomys macrotis	Crocidura negligens
Saguinus niger	Philander andersoni	Notomys mordax	Crocidura orientalis
Saguinus pileatus	Philander frenatus	Otomys barbouri	Crocidura ramona
Saguinus tripartitus	Philander mcilhennyi	Otomys cuanzensis	Crocidura rapax
Cercopithecidae	Thylamys cinderella	Otomys dartmouthi	Crocidura
			shantungensis
Cercopithecus albogularis	Thylamys karimii	Otomys dollmani	Crocidura vosmaeri
Cercopithecus denti	Thylamys pusillus	Otomys jacksoni	Crocidura wuchihensis
Cercopithecus doggetti	Thylamys sponsorius	Otomys maximus	Cryptotis brachyonyx
Cercopithecus kandti	Thylamys tatei	Otomys orestes	Cryptotis colombiana
Cercopithecus lowei	Thylamys venustus	Otomys uzungwensis	Cryptotis equatoris
Cercopithecus roloway	Dipodidae	Papagomys theodorverhoeveni	Cryptotis goldmani
Cercopithecus wolfi	Allactaga williamsi	Paramelomys gressitti	Cryptotis griseoventris
Lophocebus opdenboschi	Salpingotus thomasi	Paramelomys steini	Cryptotis mayensis
Procolobus foai	Dugongidae	Pithecheirops otion	Cryptotis medellinia
Procolobus tephrosceles	Dugong dugon	Praomys degraaffi	Cryptotis mera

Procolobus tholloni	Hydrodamalis gigas	Praomys obscurus	Cryptotis merriami
Trachypithecus ebenus	Echimyidae	Praomys petteri	Cryptotis nelsoni
Cervidae	Boromys offella	Pseudomys calabyi	Cryptotis nigrescens
Rucervus schomburgki	Boromys torrei	Pseudomys glaucus	Cryptotis obscura
Chaeropodidae	Brotomys contractus	Pseudomys gouldii	Cryptotis orophila
Chaeropus ecaudatus	Brotomys voratus	Pseudomys laborifex	Cryptotis peregrina
Cheirogaleidae	Diplomys rufodorsalis	Rattus arfakienis	Cryptotis peruviensis
Microcebus griseorufus	Echimys semivillosus	Rattus arrogans	Cryptotis phillipsii
Chinchillidae	Heteropsomys antillensis	Rattus blangorum	Cryptotis tamensis
Lagostomus crassus	Heteropsomys insulans	Rattus macleari	Cryptotis tropicalis
Cricetidae	Isothrix negrensis	Rattus nativitatis	Myosorex kihaulei
Aepeomys reigi	Isothrix sinnamariensis	Rattus omichlodes	Myosorex zinki
Akodon aliquantulus	Makalata grandis	Rattus pococki	Nesiotites hidalgo
Akodon leucolimnaeus	Makalata obscura	Rattus salocco	Nesiotites similis
Akodon montensis	Makalata occasius	Rattus sanila	Notiosorex cockrumi
Akodon mystax	Mesomys leniceps	Rattus satarae	Notiosorex evotis
Akodon oenos	Mesomys occultus	Rattus tanezumi	Notiosorex villai
Akodon paranaensis	Phyllomys kerri	Rhabdomys dilectus	Sorex antinorii
Akodon pervalens	Phyllomys lundi	Rhagamys orthodon	Sorex arunchi
Akodon reigi	Phyllomys mantiqueirensis	Solomys spriggsarum	Sorex averini
Alticola olchonensis	Phyllomys medius	Sommeromys macrorhinos	Sorex gaspensis
Amphinectomys savamis	Phyllomys nigrispinus	Spelaeomys florensis	Sorex neomexicanus
Andalgalomys roigi	Phyllomys pattoni	Taeromys arcuatus	Sorex orizabae
Andinomys edax	Phyllomys unicolor	Taeromys celebensis	Sorex veraecrucis

Arvicola scherman	Proechimys chrysaeolus	Taeromys microbullatus	Sorex yukonicus
Brucepattersonius albinasus	Proechimys echinothrix	Taeromys punicans	Suncus aequatorius
Brucepattersonius griserufescens	Proechimys gardneri	Taterillus harringtoni	Sylvisorex camerunensis
Brucepattersonius guarani	Proechimys hoplomyoides	Taterillus tranieri	Sylvisorex konganensis
Brucepattersonius igniventris	Proechimys kulinae	Thamnomys major	Sylvisorex pluvialis
Brucepattersonius misionensis	Proechimys magdalenae	Uromys boeadii	Spalacidae
Brucepattersonius paradisus	Proechimys pattoni	Uromys emmae	Spalax carmeli
Brucepattersonius soricinus	Proechimys roberti	Uromys neobritannicus	Spalax galili
Calomys callidus	Proechimys urichi	Uromys porculus	Spalax giganteus
Calomys expulsus	Thrichomys inermis	Uromys siebersi	Spalax golani
Calomys tocantinsi	Thrichomys pachyurus	Vandeleuria nilagirica	Spalax judaei
Calomys venustus	Trinomys mirapitanga	Zyzomys palatalis	Tachyoryctes ankoliae
Chelemys delfini	Trinomys moojeni	Mustelidae	Tachyoryctes annectens
Chibchanomys orcesi	Trinomys myosuros	Hydrictis maculicollis	Tachyoryctes audax
Delomys collinus	Trinomys paratus	Lutra nippon	Tachyoryctes daemon
Eothenomys cachinus	Trinomys yonenagae	Meles anakuma	Tachyoryctes ibeanus
Eothenomys miletus	Elephantidae	Meles leucurus	Tachyoryctes naivashae
Eothenomys wardi	Loxodonta cyclotis	Neovison macrodon	Tachyoryctes rex
Euneomys fossor	Equidae	Mystacinidae	Tachyoryctes ruandae

Graomys centralis	Equus asinus	Mystacina robusta	Tachyoryctes ruddi
Habromys delicatulus	Equus burchellii	Neobalaenidae	Tachyoryctes spalacinus
Habromys ixtlani	Equus caballus	Caperea marginata	Tachyoryctes storeyi
Juliomys rimofrons	Equus quagga	Nesomyidae	Suidae
Juscelinomys candango	Erethizontidae	Beamys major	Babyrousa bolabatuensis
Juscelinomys guaporensis	Echinoprocta rufescens	Brachytarsomys villosa	Sus oliveri
Juscelinomys huanchacae	Sphiggurus ichillus	Cricetomys ansorgei	Sus salvanius
Loxodontomys pikumche	Sphiggurus roosmalenorum	Cricetomys kivuensis	Thylacinidae
Megalomys desmarestii	Erinaceidae	Dendromus leucostomus	Thylacinus cynocephalus
Megalomys luciae	Erinaceus roumanicus	Dendromus nyasae	Thylacomyidae
Megaoryzomys curioi	Hylomys megalotis	Eliurus antsingy	Macrotis leucura
Microakodontomys transitorius	Eschrichtiidae	Eliurus ellermani	Tragulidae
Microtus anatolicus	Eschrichtius robustus	Eliurus grandidieri	Tragulus versicolor
Microtus bavaricus	Felidae	Eliurus petteri	Tragulus williamsoni
Microtus brachycercus	Catopuma badia	Monticolomys koopmani	Trichechidae
Microtus dogramacii	Catopuma temminckii	Nesomys audeberti	Trichechus inunguis
Microtus irani	Felis bieti	Nesomys lambertoni	Trichechus manatus
Microtus liechtensteini	Felis catus	Steatomys bocagei	Trichechus senegalensis
Microtus paradoxus	Felis manul	Steatomys opimus	Vespertilionidae
Microtus qazvinensis	Leopardus braccatus	Voalavo gymnocaudus	Eptesicus kobayashii
Microtus savii	Leopardus pajeros	Nesophontidae	Histiotus humboldti
Microtus schidlovskii	Prionailurus iriomotensis	Nesophontes edithae	Kerivoula africana

Myodes imaizumii	Profelis aurata	Nesophontes	Kerivoula lenis
		hypomicrus	
Neacomys dubosti	Uncia uncia	Nesophontes longirostris	Lasiurus atratus
Neacomys minutus	Galagidae	Nesophontes major	Lasiurus salinae
Neacomys musseri	Galago alleni	Nesophontes micrus	Lasiurus varius
Neacomys paracou	Galago cameronensis	Nesophontes paramicrus	Lasiurus xanthinus
Necromys benefactus	Galago gabonensis	Nesophontes submicrus	Miniopterus africanus
Necromys lenguarum	Galago granti	Nesophontes superstes	Miniopterus macrocneme
Necromys punctulatus	Galago nyasae	Nesophontes zamicrus	Miniopterus paululus
Nectomys apicalis	Galago orinus	Ochotonidae	Miniopterus shortridgei
Nectomys magdalenae	Galago rondoensis	Ochotona turuchanensis	Murina grisea
Neodon forresti	Galago thomasi	Octodontidae	Murina silvatica
Neotoma anthonyi	Otolemur monteiri	Aconaemys porteri	Myotis abei
Neotoma bunkeri	Geomyidae	Octodon pacificus	Myotis alcathoe
Neotoma leucodon	Cratogeomys gymnurus	Pipanacoctomys aureus	Myotis australis
Neotoma macrotis	Cratogeomys neglectus	Salinoctomys loschalchalerosorum	Myotis bucharensis
Neotoma martinensis	Cratogeomys tylorhinus	Odobenidae	Myotis ciliolabrum
Nesoryzomys darwini	Cratogeomys zinseri	Odobenus rosmarus	Myotis davidii
Nesoryzomys indefessus	Gliridae	Otariidae	Myotis hosonoi
Noronhomys vespuccii	Dryomys niethammeri	Arctocephalus australis	Myotis ikonnikovi
Oecomys auyantepui	Eliomys munbyanus	Arctocephalus forsteri	Myotis mystacinus
Oecomys catherinae	Graphiurus angolensis	Arctocephalus galapagoensis	Myotis nipalensis
Oecomys paricola	Graphiurus johnstoni	Arctocephalus gazella	Myotis occultus

Oecomys trinitatis	Heptaxodontidae	Arctocephalus philippii	Myotis oxygnathus
Oligoryzomys brendae	Amblyrhiza inundata	Arctocephalus pusillus	Myotis ozensis
Oligoryzomys fornesi	Clidomys osborni	Arctocephalus townsendi	Myotis ricketti
Oligoryzomys stramineus	Elasmodontomys obliquus	Arctocephalus tropicalis	Myotis yesoensis
Oligoryzomys victus	Quemisia gravis	Callorhinus ursinus	Nycticeius aenobarbus
Oryzomys albigularis	Herpestidae	Eumetopias jubatus	Nyctophilus howensis
Oryzomys alfaroi	Crossarchus platycephalus	Neophoca cinerea	Phoniscus aerosa
Oryzomys angouya	Heteromyidae	Otaria flavescens	Pipistrellus bodenheimeri
Oryzomys auriventer	Heteromys oasicus	Phocarctos hookeri	Pipistrellus sturdeei
Oryzomys balneator	Heteromys teleus	Zalophus californianus	Plecotus alpinus
Oryzomys bolivaris	Hipposideridae	Zalophus japonicus	Plecotus kolombatovici
Oryzomys caracolus	Hipposideros scutinares	Zalophus wollebaeki	Plecotus sardus
Oryzomys chapmani	Hominidae	Peramelidae	Scotoecus albigula
Oryzomys couesi	Homo sapiens	Echymipera davidi	Scotoecus hindei
Oryzomys curasoae	Iniidae	Echymipera echinista	Scotophilus collinus
Oryzomys devius	Inia geoffrensis	Perameles eremiana	Viverridae
Oryzomys emmonsae	Lipotes vexillifer	Petauridae	Genetta bourloni
Oryzomys galapagoensis	Pontoporia blainvillei	Dactylopsila megalura	Genetta poensis
Oryzomys hammondi	Lemuridae	Phalangeridae	Prionodon linsang
Oryzomys keaysi	Eulemur albocollaris	Ailurops melanotis	Prionodon pardicolor
Oryzomys lamia	Lorisidae	Trichosurus arnhemensis	Ziphiidae
Oryzomys laticeps	Pseudopotto martini	Trichosurus cunninghami	Berardius arnuxii

Oryzomys legatus	Macropodidae	Trichosurus johnstonii	Berardius bairdii
Oryzomys levipes	Lagorchestes asomatus	Phocidae	Hyperoodon ampullatus
Oryzomys macconnelli	Lagorchestes leporides	Cystophora cristata	Hyperoodon planifrons
Oryzomys maracajuensis	Macropus greyi	Erignathus barbatus	Indopacetus pacificus
Oryzomys marinhus	Onychogalea lunata	Halichoerus grypus	Mesoplodon bidens
Oryzomys megacephalus	Manidae	Histriophoca fasciata	Mesoplodon bowdoini
Oryzomys melanotis	Manis gigantea	Hydrurga leptonyx	Mesoplodon carlhubbsi
Oryzomys meridensis	Mephitidae	Leptonychotes weddellii	Mesoplodon densirostris
Oryzomys nelsoni	Spilogale angustifrons	Lobodon carcinophaga	Mesoplodon europaeus
Oryzomys nitidus	Molossidae	Mirounga angustirostris	Mesoplodon ginkgodens
Oryzomys perenensis	Chaerephon aloysiisabaudiae	Mirounga leonina	Mesoplodon grayi
Oryzomys polius	Chaerephon ansorgei	Monachus monachus	Mesoplodon hectori
Oryzomys rhabdops	Chaerephon bemmeleni	Monachus schauinslandi	Mesoplodon layardii
Oryzomys rostratus	Chaerephon bivittatus	Monachus tropicalis	Mesoplodon mirus
Oryzomys russatus	Chaerephon bregullae	Ommatophoca rossii	Mesoplodon perrini
Oryzomys saturatior	Chaerephon chapini	Pagophilus groenlandicus	Mesoplodon peruvianus
Oryzomys scotti	Chaerephon gallagheri	Phoca largha	Mesoplodon stejnegeri
Oryzomys seuanezi	Chaerephon jobensis	Phoca vitulina	Mesoplodon traversii
Oryzomys subflavus	Chaerephon johorensis	Pusa caspica	Tasmacetus shepherdi

Oryzomys talamancae	Chaerephon leucogaster	Pusa hispida	Ziphius cavirostris
Oryzomys tatei	Chaerephon major	Pusa sibirica	
Oryzomys xanthaeolus	Chaerephon nigeriae		
Oryzomys yunganus	Chaerephon plicatus		
Oxymycterus amazonicus	Chaerephon pumilus		
Oxymycterus caparoae	Chaerephon russatus		
Oxymycterus	Chaerephon		
dasytrichus	shortridgei		
Oxymycterus josei	Chaerephon		
	solomonis		
Oxymycterus quaestor	Chaerephon tomensis		
Oryzomys tatei	Chaerephon major		
Oryzomys xanthaeolus	Chaerephon nigeriae		
Oryzomys yunganus	Chaerephon plicatus		
Oxymycterus amazonicus	Chaerephon pumilus		
Oxymycterus caparoae	Chaerephon russatus		
Oxymycterus	Chaerephon		
dasytrichus	shortridgei		
Oxymycterus josei	Chaerephon solomonis		
Oxymycterus quaestor	Chaerephon tomensis		
Pearsonomys annectens	Monodontidae		
Peromyscus beatae	Delphinapterus leucas		
Peromyscus fraterculus	Monodon monoceros		
Peromyscus hylocetes	Mormoopidae		
Peromyscus levipes	Mormoops magna		
Peromyscus pembertoni	Pteronotus pristinus		

Peromyscus sagax	Moschidae	
Phyllotis limatus	Moschus anhuiensis	
Punomys kofordi	Moschus cupreus	
Reithrodon typicus	Moschus leucogaster	
Rhagomys longilingua		
Rhipidomys caucensis		
Rhipidomys couesi		
Rhipidomys emiliae		
Rhipidomys gardneri		
Rhipidomys macrurus		
Rhipidomys modicus		
Salinomys delicatus		
Sigmodon hirsutus		
Sigmodon planifrons		
Sigmodon toltecus		
Sigmodon zanjonensis		
Tapecomys primus		
Thomasomys apeco		
Thomasomys		
Thomasonys		
cinereiventer		
Thomasomys cinereus		
Thomasomys		
cinnameus		
Thomasomys eleusis		
Thomasomys erro		
Thomasomys gracilis		
Thomasomys hudsoni		

Thomasomys macrotis		
Thomasomys		
monochromos		
Thomasomys onkiro		
Thomasomys		
popayanus		
Thomasomys praetor		
Thomasomys ucucha		
Thomasomys vulcani		

Table S 3. Complete families removed from the analysis due to missing data.

Family
Camelidae
Myzopodidae
Platacanthomyidae
Ptilocercidae
Thylacomyidae

Table S 4-7. Estimates obtained by GLMM and GLM methods at the genus level.

	GLMM			GLM	
Variable	Estimate	Std Error	Var Fixed effect	Estimate	Std Error
Adult body mass	-0.110	0.077		-0.110	0.078
Adult body mass CV	0.083	0.037		0.083	0.038
Generation length	0.031	0.051	2.322e-11	0.031	0.051
Generation length CV	-0.056	0.042		-0.056	0.042
Litter size	0.028	0.046		0.028	0.046
Litter size CV	0.090	0.040		0.090	0.040

** • 11	GLMM			GLM		
Variable -	Estimate	Std Error	Var Fixed effect	Estimate	Std Error	
Geographic range size	-0.217	0.048		-0.217	0.048	
Geographic range size CV	0.159	0.034	1.305e-11	0.159	0.034	
Latitude	0.313	0.038		0.313	0.038	

	GLMM			GLM		
Variable -	Estimate	Std Error	Var Fixed effect	Estimate	Std Error	
Activity diversity	0.062	0.038		0.015	0.033	
Diet diversity	0.054	0.042	_	0.004	0.035	
Trophic level diversity	0.332	0.028	1.305e-11 —	0.324	0.026	
Foraging strategy	0.081	0.024		0.074	0.022	
		GLMM		GL	Μ	
Variable –	Estimate	Std Error	Var Fixed effect	Estimate	Std Error	
Trophic level diversity	0.064	0.039		0.064	0.039	
Foraging strategy	0.005	0.038	-	0.005	0.038	
Adult body mass	-0.072	0.073	-	-0.072	0.072	
Adult body mass CV	0.034	0.039	0	0.032	0.039	
Litter size CV	0.024	0.040	-	0.024	0.040	
Geographic range size	-0.218	0.056	-	-0.218	0.056	
Geographic range size CV	0.158	0.040	-	0.158	0.040	
Latitude	0.254	0.047	-	0.254	0.047	

		GLMI		GLM	
Variable –	Estimate	Std Error	Var Fixed effect	Estimate	Std Error
Adult body mass	-0.056	0.121		-0.056	0.122
Adult body mass CV	0.144	0.065	-	0.144	0.066
Generation length	-0.142	0.100	_	-0.142	0.101
Generation length CV	0.002	0.069	7.287e-12	0.002	0.070
Litter size	0.013	0.076	_	0.013	0.077
Litter size CV	0.042	0.072	_	0.042	0.073
Clade age	-0.077	0.065	-	-0.077	0.065

Table S 7-10. Estimates obtained by GLMM and GLM methods at the family level.

		GLMM	(GLM	
Variable ⁻	Estimate	Std Error	Std Error Var Fixed effect		Std Error
Geographic range size	-0.248	0.090		-0.248	0.091
Geographic range size CV	0.130	0.067	2.08e-10	0.130	0.067
Latitude	0.293	0.063	-	0.293	0.064
Clade Age	-0.066	0.058		-0.066	0.059
X7 · 11		GLMM			GLM
variable —	Estimate	Std Error	Var Fixed effect	Estimate	Std Error

Activity diversity	0.126	0.078		0.081	0.061
Diet diversity	0.105	0.079		0.078	0.072
Trophic level diversity	0.205	0.069	1.305e-11	0.208	0.068
Foraging strategy	0.124	0.056		0.128	0.055
Clade Age	-0.040	0.065		-0.025	0.059

	GLMM			GLM	
Variable —	Estimate	Std Error	Var Fixed effect	Estimate	Std Error
Trophic level diversity	0.032	0.064		0.032	0.064
Foraging strategy	0.067	0.060		0.067	0.061
Adult body mass CV	0.070	0.062		0.0698	0.063
Generation length	-0.082	0.072	2.234e-11	-0.082	0.072
Geographic range size	-0.223	0.090	_	-0.223	0.091
Geographic range size CV	0.095	0.069		0.095	0.069
Latitude	0.239	0.068		0.239	0.068

	GLMM			GLM	
Variable -	Estimate	Std Error	Var Fixed effect	Estimate	Std Error
Adult body mass	-0.253	0.302		-0.253	0.328
Adult body mass CV	0.399	0.194		0.399	0.215
Generation length	0.011	0.279		0.011	0.302
Generation length CV	-0.037	0.185	1.868e-10	-0.037	0.203
Litter size	-0.132	0.162		-0.132	0.176
Litter size CV	0.108	0.117		0.108	0.118
Clade age	-0.069	0.156		-0.069	0.160

Table S 10-13. Estimates obtained by GLMM and GLM methods at the order level.

GLMM

GLM

X 7					
v ariable	Estimate	Std Error	Var Fixed effect	Estimate	Std Error
Geographic range size	-0.135	0.159		-0.135	0.160
Geographic range size CV	0.217	0.125	5.563e-12	0.217	0.126
Latitude	0.407	0.116	_	0.407	0.117
Clade Age	-0.077	0.145	—	-0.077	0.145

		GLMM	GLM		
Variable –	Estimate	Std Error	Var Fixed effect	Estimate	Std Error
Activity diversity	0.050	0.130		0.050	0.131
Diet diversity	0.172	0.156	-	0.172	0.158
Trophic level diversity	0.276	0.161	2.861e-10	0.276	0.163
Foraging strategy	0.082	0.130	-	0.082	0.132
Clade Age	0.042	0.136		0.042	0.136
Variabla —		GLMM		GLM	1
Variable –	Estimate	GLMM Std Error	Var Fixed effect	GLM Estimate	A Std Error
Variable – Trophic level diversity	Estimate 0.174	GLMM Std Error 0.159	Var Fixed effect	GLM Estimate 0.174	A Std Error 0.163
Variable – Trophic level diversity Adult body mass CV	Estimate 0.174 0.098	GLMM Std Error 0.159 0.154	Var Fixed effect 2.234e-11	GLM Estimate 0.174 0.098	4 Std Error 0.163 0.157
Variable – Trophic level diversity Adult body mass CV Geographic range size CV	Estimate 0.174 0.098 0.224	GLMM Std Error 0.159 0.154 0.126	Var Fixed effect 2.234e-11	GLM Estimate 0.174 0.098 0.224	4 Std Error 0.163 0.157 0.128

Table S 14-17 Pearon's r calculated at the genus level

Variable	Adult body n	nass	Litter size		Generation length
Adult body mass	1		-0.096		0.436
Litter size	-0.096		1		-0.434
Generation length	0.436		-0.436		1
Variable	Geographic 1	range size	Latitude		Longitude
Geographic range size	e 1		0.57	5	0.431
Latitude	0.57	5	1		0.654
Longitude	0.43	1	0.654		1
-					
Variable	Diet diversity	Activity divers	For	raging strategy diversity	Trophic level diversity
Diet diversity	1	-0.129		0.014	0.200
Activity diversity	-0.129	1		0.059	0.045
Foraging strategy diversity	0.014	0.059		1	0.011
Trophic level diversity	0.200	0.045		0.011	1

Table S 18-20. Pearson's r calculated at the family level.

Variable	Adult body mass	Litter size	Generation length
Adult body mass	1	0.986	-0.124
Litter size	0.986	1	-0.120
Generation length	-0.124	-0.120	1

Variable	Geographic range size	Latitude	Longitude
Geographic range size	1	0.355	0.267
Latitude	0.355	1	0.795
Longitude	0.267	0.795	1

Variable	Diet diversity	Activity diversity	Foraging strategy	Trophic level
Diet diversity	1	0.040	0.315	0.496
Activity diversity	0.040	1	0.103	0.143
Foraging strategy	0.315	0.103	1	0.264
Trophic level	0.496	0.143	0.264	1

Table S 20-22. Pearson's r calculated at the family level.

Variable	Adult body mass		Litter size	Generation length
Adult body mass	1		-0.200	0.763
Litter size	-0.2	00	1	-0.531
Generation length	0.7	63	-0.531	1
Variable	Geographic	range size	Latitude	Longitude
	Geographie	Tunge Size	0.127	
Geographic range size	1		0.137	0.029
Latitude	0.137		1	0.887
Longitude	0.029		0.887	1
Variable	Diet	Activity	Foraging strategy	Trophic level
Diet diversity	1	0.176	0.426	0.741
Activity diversity	0.176	1	-0.095	0.116
Foraging strategy	0.426	-0.095	1	0.340
Trophic level	0.741	0.116	0.340	1

Table S 23-25. Variance Inflation Factors calculated for all models.

Variable	Genus	Family	Order
Body mass	1.119	1.370	2.117
Body mass CV	1.073	1.778	4.738
Generation length	1.612	2.024	3.659
Generation length CV	1.087	1.478	3.634
Litter size	1.472	1.961	2.441
Litter size CV	1.217	1.553	1.126
Clade age	-	1.156	1.342

Variable	Genus	Family	Order
Geographic range size	1.546	1.522	1.513
Geographic range size	1.195	1.448	1.35
Latitude	1.641	1.454	1.179
Clade age	-	1.005	1.026

Variable	Genus	Family	Order
Activity diversity	1.038	1.025	1.068
Diet diversity	1.099	1.547	2.040
Foraging strategy	1.021	1.17	1.345
Trophic level diversity	1.091	1.415	1.915
Clade age	-	1.035	1.023