

University College London

Department of Genetics, Evolution and Environment

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

Adrienne Etard

Primary supervision: Dr. Tim Newbold

Secondary supervision: Dr. Alex Pigot

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Abstract

Key words: vertebrates; traits; missing values imputation; land-use change; functional diversity.

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List of abbreviations

BM	Body mass
BL	Body length
CR	Cropland
DA	Diel activity
DFR	Dendrogram-based functional richness
Di	Diet
DB	Diet breadth
FDis	Functional dispersion
FRic	Volume-based functional richness
GL	Generation length
HB	Habitat breadth
ISV	Intermediate secondary vegetation
L	Longevity
LCS	Litter/clutch size
TL	Trophic level
ITIS	Integrated Taxonomic Information System
LUCC	Land-use and climate change
MA	Maturity
MSE	Mean-squared error
MSV	Mature secondary vegetation
OOB	Out-of-bag
PA	Pasture
PD	Primary diet
PF	Plantation forest
PFC	Proportion of falsely classified
PREDICTS	Projecting Responses of Ecological Diversity In Changing Terrestrial Systems
PV	Primary vegetation
Q	Rao quadratic entropy
RS	Range size
SI	Supporting Information
UR	Urban
YSV	Young secondary vegetation

1 | Introduction

Anthropogenic activities are driving global biodiversity declines at unprecedented rates. Currently, habitat conversion and degradation – induced mainly by anthropogenic land-use change – are the primary causes of biodiversity loss (Newbold et al., 2015; Pereira et al., 2012). Climate change is projected to be one of the biggest driver of biodiversity loss by 2070, matching or exceeding the deleterious impacts of land-use change on ecological communities (Newbold, 2018). Understanding how land-use and climate change (LUCC) act on biodiversity, separately and in combination, is key to project the future responses of species, and to consequently put into place efficient policies for biodiversity conservation. Furthermore, biodiversity losses affect ecosystem properties, and can adversely impact the delivery of ecosystem services (Duraiappah et al., 2005; Hooper et al., 2012; Oliver et al., 2015). Assessing if and how biodiversity decreases link to the loss of ecosystem functions is a key research area (Lefcheck et al., 2015; Petchey and Gaston, 2006). Indeed, investigating how anthropogenic activities are likely to impact biodiversity, and how these impacts may relate to ecosystem processes, can help put into place mitigation measures aiming at protecting both biodiversity and ecosystem functioning.

Although there remains much uncertainty, it has now been established across diverse taxonomic groups that species traits mediate species responses to environmental changes, notably to LUCC (Angert et al., 2011; Estrada et al., 2018; Newbold et al., 2013; Pacifici et al., 2017; Pearson et al., 2014). McGill et al. (2006) defined traits as well-defined organismal characteristics, that can be measured at the individual level and that be used comparatively across species . Functional traits are those that particularly influence organismal fitness or performance (Violle et al., 2007). Functional traits relate to species’ abilities to exploit their biotic and abiotic environment. Functional traits can be divided into two groups. Functional traits that underpin species contributions to ecosystem processes have been termed ‘effect traits’ (Lavorel and Garnier, 2002; Wong et al., 2018). Effect traits determine how species use environmental resources and influence ecosystem processes. On the

other hand, ‘response traits’ shape how species respond to disturbances. Therefore, functional traits underpin both species’ aptitudes to cope with environmental changes and their role in ecosystem functioning. Certain functional traits can act as both effect and response traits. Conceptually, these are particularly interesting for investigating the impacts of environmental changes on ecosystem processes and services, as they provide a mechanistic understanding of how stressors affect both species’ responses and ecosystem processes (Hevia et al., 2017; Lavorel and Garnier, 2002; Luck et al., 2012).

Assessing the impacts of human activities on ecosystem functioning is increasingly important as pressures rise globally. Publications linking drivers of change and delivery of ecosystem services have increased exponentially since 2001 (Hevia et al., 2017); nevertheless, how species traits influence their responses to land-use and climate change, and how this relates to the loss of important ecosystem functions, remains to be largely explored. Notably, most studies investigating these questions have been conducted at local or regional scales on given taxonomic groups. Although response traits to LUCC have been identified in terrestrial vertebrates, there still remains uncertainty, for example as to which are most important to define species responses (Wheatley et al., 2017). In terrestrial vertebrates, no study has, to my knowledge, attempted to investigate how anthropogenic pressures are likely to disrupt ecosystem processes supported by terrestrial vertebrates at global scales.

The final aim of this PhD project is to investigate the effects of terrestrial vertebrate species’ traits on their responses to LUCC, at global scales. Specifically, my main goals are (1) to elucidate which traits are likely to put species at greater risk from land-use and climate change; (2) to investigate whether future biodiversity declines triggered by these anthropogenic threats are likely to disrupt important ecosystem functions. Unlike previous published studies, this work will investigate these questions at global scales, and simultaneously across the four terrestrial vertebrate classes – amphibians, birds, reptiles and mammals.

This report synthesizes the work I have achieved throughout the first year of my PhD. I start by reviewing the literature to present the questions I have addressed in the context of the past and current ecological research (Chapter 2). In Chapter 3 and 4, I expose the work achieved so far. Specifically, Chapter 3 focuses on the collation of extensive trait data across terrestrial vertebrates. Using these data, Chapter 4 presents a first analysis showing that land-use change promotes the functional homogenisation of local vertebrate communities. Finally, I present an outline of the questions that I plan to investigate in the upcoming years in Chapter 5.

2 | Literature review and hypotheses for the present work

2.1 Land-use and climate change, species traits and the functional composition of vertebrate communities

2.1.1 Land-use and climate change and species response traits

Currently, terrestrial land-use change is the most important driver of biodiversity declines (Chaudhary and Kastner, 2016; Maxwell et al., 2016; Newbold et al., 2015). With climate change projected to be catching up by 2070 (Newbold, 2018), it has become vital to understand how these threats will affect biodiversity, separately and in combination. By influencing species responses to environmental changes, response traits can provide a mechanistic understanding of how diverse threats shape ecological communities, an understanding particularly relevant for conservation policies.

There is now empirical evidence across taxonomic groups that species traits influence their responses to LUCC. For instance, response traits to LUCC have been identified in terrestrial plant (Díaz et al., 2001), fungal (Koide et al., 2014), invertebrate (Hall et al., 2019; Williams et al., 2010), and terrestrial vertebrate species (Table 2.1). It is important to point out that in some cases, contrasting results are found (for instance, in bees: larger body size having been found to influence species responses to land-use change both negatively (Larsen et al., 2005) and positively (De Palma et al., 2015), or having been found to have little effect (Forrest et al., 2015; Williams et al., 2010), see Bartomeus et al. (2018)). This highlights the fact that studies may be context-dependent, with contingent limitations. For vertebrates, most studies, conducted on different taxa and at different scales, tend to show that larger, longer-lived specialist species with a lower reproductive output are more likely to be impacted negatively by LUCC (Table 2.1). As such, a number of response traits to

land-use or climate change have been identified for vertebrate species belonging to diverse Classes (Table 2.1).

Despite this empirical evidence, there is still a need to refine our understanding of which traits significantly influence responses. As traits are commonly used to assess species vulnerability to threats or extinction risks (Böhm et al., 2016; Pacifici et al., 2015; Willis et al., 2015), it is particularly important to be confident about how they act on species responses. Trait-based approaches can be opposed to trend-based approaches (Pacifici et al., 2015), which rely on historic population trends (changes in abundance or shifts in distributions) to predict species vulnerability and extinction risks. Trend-based approaches require important field work effort to monitor species populations. Getting extensive information on all species population trends is virtually impossible. The appeal of trait-based approaches is that, by providing mechanistic insights, they diminish the amount of population information needed. If species' responses to a threat consistently relate to certain traits, it is possible to generalise patterns across species for which data is less available (Verberk et al., 2013). Nevertheless, for several reasons that I now expose, how species traits influence their responses to LUCC remains unclear.

First, there is a lack of comprehensive understanding about which traits are important in shaping species responses to climate change. Wheatley et al. (2017) compared different published climate change vulnerability assessment frameworks, some of which trait-based, some trend-based, and some incorporating elements of both (hybrid). They found that the different frameworks, applied to the same set of species, did not yield consensual outputs and classified species inconsistently into different risk categories. Their work underlines that currently, trend-based vulnerability assessments perform better at identifying species at risks from climate change than trait-based approaches. This study highlights the current lack of unanimous understanding as to which traits to consider, and how, in vulnerability assessments. More broadly, their study stresses the need to clarify our understanding of how response traits to climate change act across different taxa. The finding in Wheatley et al. (2017) that there is no consensus across assessment frameworks might be explained by the fact that frameworks were initially designed and tested for a particular taxon – generally at the Class level or lower ranks –, and do not hold when applied to other taxa. They nevertheless argue that frameworks should be universally applicable. Their findings put into question to our current ability to extrapolate the knowledge of response traits gathered for certain taxa to other taxonomic groups.

Table 2.1: Some references providing with empirical evidence for response traits to LUC in terrestrial vertebrates. Also see Hevia et al. (2017).

Pressure	Reference	Trait or property	Taxa	Study scale
Land-use change	Newbold et al. (2013)	Generation length	Birds	Pan-tropical forests
		Body mass		
		Migratory activity		
		Diet specialisation		
Urbanisation	La Sorte et al. (2018)	Habitat specialisation	Birds	Global (58 cities)
		Body mass		
		Range size		
		Diet specialisation		
Habitat modification	Nowakowski et al. (2017)	Foraging strata	Amphibians	Global
		Habitat specialisation		
		Range size		
		Habitat specialisation		
Land-use change	Flynn et al. (2009)	Litter size (Mammals)	Mammals Birds	America
		Diet (Mammals)		
		Body mass (Birds)		
		Diet (Birds)		
Human pressure history (human population density and land-conversion history)	Rapacciuolo et al. (2017)	Foraging habits (Birds)	Tetrapods	Western Hemisphere
		Body mass		
		Body mass		
		Body mass		
Land-use change	Tinoco et al. (2018)	Reproductive rate	Hummingbirds	Andes Mountains, southern Ecuador
		Habitat specialisation (Mammals)		
		Sexual maturity age (Mammals)		
		Habitat specialisation (Birds)		
Climate change (Range filling proxy)	Estrada et al. (2018)	Reproductive output (Birds)	Mammals Birds	Europe
		Habitat specialisation (Mammals)		
		Diet (Mammals)		
		Dispersal abilities (Birds)		
Climate change	Pacifci et al. (2017)	Generation length (Birds)	Mammals Birds	Global
		Altitudinal range (non-breeding) (Birds)		
		Occupied area		
		Generation length		
Climate change	Pearson et al. (2014)	Operational thermal range	Amphibians Reptiles	USA
		Body mass		
		Activity time		
		Dispersal ability		
Climate change	McCain and King (2014)	Diet breadth	Mammals	North America
		Migratory status		
		Reliance on open water		
		Reliance on open water		
Climate change	Schloss et al. (2012a)	Dispersal ability	Mammals	Western Hemisphere
		Diet breadth		
		Migratory status		
		Reliance on open water		
Climate change	Angert et al. (2011)	Dispersal ability	Birds (Passeriformes)	North America
		Diet breadth		
		Migratory status		
		Reliance on open water		

To my knowledge, comparative studies looking at whether response traits to LUCC differ across taxonomic groups (at ranks higher than Class), experiencing the same threat levels under similar conditions, are rare. The picture becomes even more complex when different studies find contradicting results within a taxon, such as was the case in some bee species (Bartomeus et al., 2018). Moreover, the importance of response traits may vary geographically. The work by Bartomeus et al. (2018) further emphasises the idea that unless similar response traits to a threat are identified consistently across different systems and taxa, our ability to use traits as predictors of vulnerability or extinction risk remains limited. For these reasons, it is necessary to conduct comparative analyses across taxa, to identify response traits, verify whether they are conserved across species and whether they have the same importance in shaping responses across taxonomic groups and geographical areas.

Second, another difficulty when identifying response traits is that different threats can be acting on the studied ecological community, so that observed modifications stem from the interactions of diverse response traits (Gonzalez-Suarez et al., 2013). Response traits must be identified for a single threat while controlling for others, before investigating potential interacting effects. Nevertheless, this is difficult to achieve when using global empirical data. For land-use change, this difficulty can be overcome by using data collected over sufficiently small-scale areas, over which other pressures can be assumed to be negligible.

To conclude, potential taxon-, threat- and geographical dependence of response traits to land-use or climate change makes it difficult to generalise patterns observed at local scales. This stresses the need to conduct global, cross-taxon studies to verify whether empirical evidence supports the generalisation of any response trait. Identifying response traits using global scale data, across the four terrestrial vertebrate classes, is one the goal of my PhD project.

2.1.2 Land-use and climate change, functional diversity and the disruption of ecosystem services

Response traits allow to understand and predict how environmental pressures are likely to modify ecological assemblages (changes in species richness and abundance). These alterations can lead to modifications in functional diversity (the diversity and variability of functional traits in a community). Several indices have been developed in the recent years to estimate diverse components of functional diversity (Laliberte and Legendre, 2010; Legras et al., 2018; Schleuter et al., 2010; Villéger et al., 2008). Functional diversity indices are interesting for at least two reasons. First, they can inform on how disturbances affect trait community composition. Second, as functional (effect)

traits relate to ecosystem functions, measures of functional diversity are often used as a proxy for ecosystem functioning. I will develop these two points in more detail further down, and I will mainly focus on land-use change as the disturbance of interest.

Impacts of land-use change on the functional diversity of vertebrate communities. Response traits determine whether a species is likely to be removed from a community due to the environmental filtering exerted by a pressure such as land-use change. Environment filtering is a major driver of community structure (but see Cadotte and Tucker (2017)); it refers to a process whereby environmental conditions select out species that cannot get established and cannot persist in a given area. As environmental filtering imposes barriers on establishment and survival, it is expected that species with similar traits, which render them able to persist in the altered conditions, are favoured (Cadotte and Tucker, 2017; Wong et al., 2018). Consequently, the trait composition of emergent communities is expected to be non-randomly impacted. The trait composition of a community can be assessed in different ways. When dealing with individual traits separately, community-weighted means are often employed. Trait distributions can also be compared across a land-use gradient. Finally, functional diversity indices allow to consider several traits simultaneously. As such, they provide with estimates summarising multivariate trait composition.

Various indices have been developed in the past years to estimate different facets of functional diversity. They have notably been reviewed in Schleuter et al. (2010) and in Legras et al. (2018). Some indices are, by construction, independent from species richness, while others are known to covary with species richness (Schleuter et al., 2010). Here, I will present two indices aiming at estimating the functional richness of ecological communities, as well as two indices quantifying multivariate trait dispersion.

Functional richness. Functional richness was initially assessed by grouping species together into functional groups: species sharing similar trait values were assumed to belong to the same functional group. Functional richness was then assessed as the total number of functional groups (Legras et al., 2018). As underlined by Legras et al. (2018), this approach is problematic for several reasons: the definition of functional groups depends on users' choices, notably to define trait boundaries between groups, which is particularly problematic with continuous traits. Consequently, other indices have been developed to estimate functional richness; in this work, I focus on two indices, which rely two different conceptual bases. These indices will be described in more details in Chapter 4 (see Figure 4.2 in Chapter 4).

FRic (Villéger et al., 2008). The FRic index, developed by Villéger et al. (2008), aims at estimating the amount of trait space that species occupy. This index relies on the projection of species in a multidimensional space, where each dimension is a trait (or a principal component, if the dimensionality of the trait dataset has been reduced). Species are placed in the multidimensional space according to their trait values. The functional richness is then estimated as the volume of the convex hull that encompasses all species of a given community (Villéger et al., 2008).

DFR (denominated ‘FD’ by Petchey and Gaston (2002)). The dendrogram-based functional richness index developed by Petchey and Gaston (2002) aims at estimating the total functional distance among species of a given community. Its calculation relies on the obtention of a functional dendrogram, from which functional richness is estimated as the sum of branch lengths for the species in a given community. The functional dendrogram is obtained by clustering a species×species distance matrix, derived from a species×trait dataset. In this work, dendrogram-based functional richness is referred to as DFR. DFR has notably been criticised for being sensitive to the choice of the clustering method (Legras et al., 2018).

Both FRic and DFR are conceptually, by construction, not independent from species richness. In experimental studies and natural communities, a positive correlation between these metrics is often found (Cadotte et al., 2011). For this reason, examining whether functional richness indices inform on community dynamics differently from species richness is an important question to elucidate. Indeed, if species richness is as informative as functional richness, the latter is not worth measuring: species richness is then a proxy for functional richness. This question was central to the study conducted by Cadotte et al. (2011). By reviewing the literature, they found that the relationship between functional richness and species richness is context-dependent, and that the shape of the relationship notably depends on the amount of functional redundancy in the community.

Functional redundancy aims at describing the degree to which species in a given assemblage share similar trait values, and, as such, sustain similar ecosystem processes (Mayfield et al., 2010; Ricotta et al., 2016; Rosenfeld, 2002). In communities with a high degree of functional redundancy, functions can be maintained despite species loss. On the other hand, the loss or gain of functionally diverse species can lead to marked variations in functional richness, despite small changes in species richness (Figure 2.1). Furthermore, Mayfield et al. (2010) showed that the relationship between species richness and functional richness could be affected in different ways by human land-uses. They proposed diverse mechanisms building upon community assembly processes to explain how

land-uses may influence species richness – functional richness trajectories.

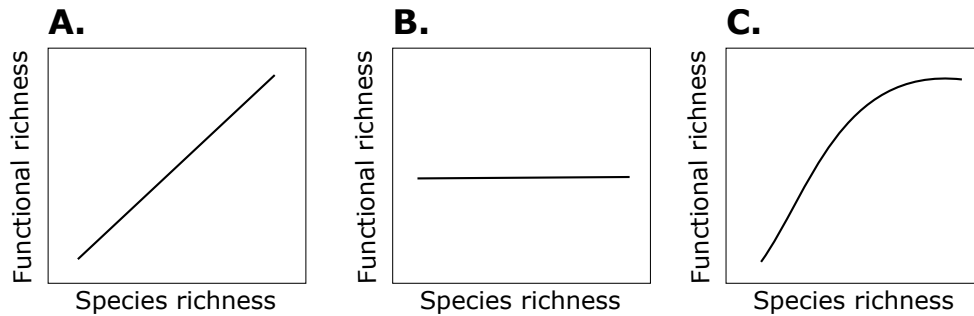


Figure 2.1: The species richness – functional richness relationship informs on functional redundancy. (A) High positive correlation between species richness and functional richness: functional richness increases at a constant rate with species richness. The gain or loss of species can lead to marked variations in functional richness. The rate of increase is higher than in (B), where functional richness remains constant despite variations in species richness. The functional redundancy of (B) is comparatively higher than that of (A). (C) The rate at which functional richness increases with increases in species richness is not constant. The functional redundancy increases with higher species richness.

The potential correlation of species richness with functional richness has two practical consequences.

First, as underlined above, the shape of the relationship can inform on the functional redundancy of a given community (Cadotte et al., 2011), and Figure 2.1. Thus, changes in the relationship between functional richness and species richness across a land-use gradient can provide insights into how land-use change impacts the functional redundancy of ecological communities. In assemblages with functionally redundant species, random species loss or addition are unlikely to lead to marked variation in functional richness.

Second, when species richness strongly correlates with functional richness (Figure 2.1 A), decreases in functional richness along a land-use gradient may be due to decreases in species richness. In other words, an observed change in functional richness may be driven by changes in species richness alone rather than by environmental filtering. As such, it becomes necessary to disentangle the effects of species richness from the effects of the environmental variable of interest. To that end, a common approach consists in generating the null expectation of functional richness given a species richness. Then, empirical values can be compared against null expectations. This can be achieved through simulations where the community composition is randomised, given a species richness, so as to generate a null distribution of functional richness (Flynn et al., 2009; Wong et al., 2018).

Functional dispersion. When a community is subject to environmental filtering, functional clustering is expected to be observed in the emergent community (Cadotte and Tucker, 2017; Wong et al., 2018). Functional clustering — also referred to as functional under-dispersion — qualifies

communities where species are more similar, in term of their traits, than expected by chance. Diverse indices have been developed to quantify the functional dispersion of ecological communities (for instance, Rao’s quadratic entropy, or functional dispersion FDis, developed by Laliberte and Legendre (2010)). Rao’s quadratic entropy and FDis aim at assessing the degree to which species resemble each other in terms of their traits. Chapter 4 provides more details on their definition and calculation (see Figure 4.3).

Rao’s quadratic entropy and functional dispersion are highly correlated. Both can be used to assess whether species in a given community are functionally clustered, as a consequence of environmental filtering. For instance, the observed dispersion in trait values can be compared to the null expectation of functional dispersion, obtained from a null model. As these indices are, by construction, independent from species richness, the effects of environmental filtering on functional dispersion can also be deduced directly from shifts in the values of the indices along an environmental gradient.

As anthropogenic land-uses globally negatively impact local species richness (Newbold et al., 2015), decreases in functional richness of local ecological communities are likely to take place, particularly in communities with low functional redundancy. Moreover, land-use change could also alter the functional richness of communities without altering local species richness. For instance, Flynn et al. (2009) showed that the functional richness (DFR) of avian, mammalian and plant communities located in the Western hemisphere decreased under agricultural intensification. Chapman et al. (2018) showed that the functional richness (DFR) of tropical bird communities declined along a land-use gradient of increasing human disturbance. Other studies have reported shifts in the distribution of trait values along land-use gradients (La Sorte et al., 2018; Rapacciuolo et al., 2017). Overall, all these studies show that land-use change alters the functional richness of local vertebrate assemblages. Nevertheless, most studies were conducted at local or regional scales.

Some studies have been conducted at global scales to understand patterns in the functional dispersion of avian assemblages (Cooke et al., 2019), or in the functional richness (DFR) of mammalian communities (Safi et al., 2011). Nevertheless such studies did not focus on land-use change or other anthropogenic pressures, but on latitudinal gradients of species diversity. As such, to my knowledge, no study has yet investigated how land-use change impacts the functional diversity of local vertebrate communities at global scales.

Studies looking at the effects of climate change on the functional diversity of terrestrial vertebrates are also rare. Barbet-Massin and Jetz (2015) investigated how future climate change was pro-

jected to impact the functional diversity of global avian assemblages through range shifts. Thuiller et al. (2014) investigated how both projected land-use and climate change affected the functional diversity of European avian assemblages. Both studies highlighted that future range shifts due to climate change led to uneven effects on functional diversity across space, with some areas projected to experience substantial loss of functional diversity. To my knowledge, no work has investigated how future climate change is likely to impact the functional diversity of other vertebrate taxa (herpetiles or mammals) at global scales.

To summarise, past empirical evidence has shown that (1) species sensitivity to LUCC depends on their traits; (2) LUCC is reshaping the functional composition of ecological assemblages, potentially disrupting important functions. Nevertheless, most studies have been conducted at local or regional scales, so that it is still unknown how global changes affect the functional diversity of local terrestrial vertebrate communities. Tackling this question, with land-use change at the disturbance of interest, is the aim of Chapter 4.

The recent development of many functional diversity indices, synthesising the diversity of functions in a community, reflects the importance of understanding how anthropogenic pressures are likely to modify ecosystem processes. In the field of biodiversity-ecosystem functioning relationships, it is now well established that higher species diversity is associated with higher ecosystem productivity and stability, better use of limiting resource, as well as better resistance to biological invasions (Tilman et al., 2014). I now explore the links between functional diversity indices and ecosystem functioning in more details.

Functional diversity and ecosystem functioning Early experiments investigating the relationships between functional composition and ecosystem functioning classified species in broad functional groups. Species belonging to similar groups were assumed to have similar effects on ecosystem processes (Legras et al., 2018). Ecosystem functioning was measured in various ways, depending on the studied system. For instance, plant biomass was used to measure primary productivity in Weisser et al. (2017). The field, overall, was dominated by experiments conducted on plant communities (Tilman et al., 2014), which are easy to manipulate. A higher number of functional groups was correlated to better ecosystem functioning and resilience (Hector et al., 1999; Tilman and Downing, 1994). Consequently, the consensus that emerged was that higher levels of diversity meant higher ecosystem stability and performance; this idea is now widely accepted (Hooper et al., 2005; Hooper et al., 2012; Oliver et al., 2015).

Conceptually, functional effect traits are the mechanistic links between biodiversity and ecosystem functioning (Lavorel and Garnier, 2002; Violle et al., 2007). A higher diversity of effect traits induces increases in ecosystem performance, through, for example, more interspecific complementarity in resource use, or greater use of limiting resources (Tilman et al., 2014). Empirically, functional diversity indices have been found to be better predictors of ecosystem functioning than species richness (Abonyi et al., 2018; Cadotte et al., 2011; Flynn et al., 2011). Most studies looking at patterns of functional diversity now invoke this argument to justify the use of functional diversity indices. Researchers often claim that decreases in functional diversity indices could reflect the imperilment of ecosystem processes. Although this holds true for plants, for which there is a wealth of empirical evidence, the picture is more complex for animal communities.

In animal communities, the use of functional diversity as a proxy for ecosystem functioning must be carefully justified. Indeed, there is to date little empirical evidence that functional diversity indices correlate with ecosystem processes supported by animal communities (Didham et al., 2016; Hatfield et al., 2018). For instance, in a meta-analysis of 24 studies looking at the effects of landscape change on functional diversity metrics, Hatfield et al. (2018) found that only five studies assessed whether functional diversity related to measures of ecosystem functioning. Moreover, these five studies overall found weak and contradictory associations between the functional metrics and the measures of ecosystem functioning. Hatfield et al. (2018) argued that, without empirical evidence of a strong link between functional diversity metrics and ecosystem processes, there is little incentive to continue to quantify the functional diversity of communities, in particular of animal assemblages. Cadotte et al. (2011) also emphasized the need to clearly justify the use of functional diversity indices by showing that there is a correlation between functional indices and ecosystem functioning. Indeed, they argued that functional indices are worth measuring only if they provide with novel insights, compared to species richness, and if they reflect ecosystem function. Here, I acknowledge the points Cadotte et al. (2011) and Hatfield et al. (2018). Nevertheless, as exposed in the previous section, I argue that functional indices are still useful to document how global changes are altering the composition of animal communities. However, understanding how functional diversity indices relate to ecosystem functioning in animal communities is vital, and remains to be largely explored.

In animal communities, the use of functional diversity indices as proxies for ecosystem functioning is complexified by several elements that are less problematic in plant communities. First, ecosystem processes supported by animals may involve more than one taxon: for instance, both birds and arthropods impact pest control through predatory activity. Ewers et al. (2015) showed

that, despite decreases in abundance in several invertebrate taxa between two land-uses, rates of decomposition, predation and seed consumption did not vary; vertebrate species compensated the loss of invertebrates by assuming similar functions at higher rates in altered land-uses. Therefore, the functional diversity of invertebrate species in this study could have been observed to decrease between the two land-uses, whereas no effect was observed on ecosystem functioning.

Second, a single ecosystem process could arise from a combination of different traits, some of which difficult to measure. Appropriate trait selection is vital to ensure that functional diversity indices reflect targeted ecosystem functions (Luck et al., 2012). As emphasised by Didham et al. (2016), the choice of functional traits must be mechanistically justified. However, the availability of trait data may be problematic, or traits could be difficult to measure. It may also be unclear which traits are more important in defining a given process. As such, the lack of correlation between functional diversity indices and ecosystem processes may arise from the difficulty to select and obtain appropriate traits in vertebrate communities.

Finally, experimental set-ups in controlled conditions are much more difficult to put into place for terrestrial communities. It is extremely difficult to manipulate the composition of vertebrate communities, as is done in plant communities. Data is therefore mostly obtained from field studies, which may have confounding factors, including diverse taxa whose functional roles are neglected (with possibly, compensatory effects as underlined above, shown in Ewers et al. (2015)).

Overall, it remains largely unclear how the diversity and variability of vertebrate functional traits relates to ecosystem processes, despite the ecological importance of terrestrial vertebrates. If carefully designed functional diversity indices consistently related to ecosystem functioning, they would be relevant measures for the conservation of ecosystem processes and species.

2.1.3 Vertebrate ecological roles

Vertebrate species play significant roles in ecosystem functioning, as they support a wide range of processes (Hocking et al., 2014; Sekercioglu, 2006; Severtsov, 2013). Vertebrates are also very important for human societies, both culturally and as sources of proteins (Albert et al., 2018; Alves et al., 2018; Hirons et al., 2016). Here, I briefly review the main ecological roles terrestrial vertebrate species participate in (mainly pollination, seed dispersal, predation, grazing and nutrient cycling).

First, vertebrate species participate in shaping global plant communities. The reproductive success of many plant species depends on vertebrates. Indeed, vertebrate species are significant pollinators (Ratto et al., 2018). Moreover, vertebrates are essential actors of seed dispersal. About

56% of angiosperm species rely on biotic seed dispersal, either obligatory (14%) or in complement to abiotic dispersal (42%), and 46% of gymnosperm species strictly rely on biotic seed dispersal (Tiffney, 2004). Vertebrates disperse seeds most frequently through endozoochory (ingestion of the disseminule or of part of the disseminule), and less frequently through exozoochory (where the disseminule gets attached to the surface of the disperser). Thus, through pollination and frugivory, vertebrates are important in maintaining gene flows among plant communities and impact the genetic diversity of plant assemblages (Calvino-Cancela et al., 2012).

By exerting top-down control, vertebrate grazers and herbivores regulate plant populations and influence global plant diversity patterns. Lin et al. (2018) and Zhang et al. (2018) both found global evidence that top-down interactions between vertebrates and plants shaped global plant communities. Mammalian seed predation contributes to the structure of tree communities (Paine et al., 2016). As ecosystem engineers (for example, through burrowing behaviours), vertebrates impact soil properties and influence the structure and the composition of plant assemblages (Sekericioglu, 2006; Severtsov, 2013).

Second, vertebrate species contribute to regulate animal populations through predatory activity (Barber et al., 2010; Letnic et al., 2012; Luck et al., 2012; Salo et al., 2010). Through both predatory and herbivory, vertebrates have a significant influence on the structure of food webs.

Finally, vertebrate species participate in energy transfers between the biota and the abiotic environment. For instance, they take part in nutrient cycling and matter decomposition through scavenging (Cunningham et al., 2018; Inger et al., 2016; Wilson and Wolkovich, 2011). Vertebrate excrements can modify nutrient availability in the soil, with cascading effects on the structure of plant communities (Severtsov, 2013).

To conclude, the range of ecosystem processes that vertebrate sustain is defined by their contribution to matter and energy flows at the ecosystem scale. Food webs are key to understand the transfer of matter and energy within the biota (interactions within and among trophic levels), from which multiple ecosystem properties emerge. Vertebrate species also contribute to mineralise organic matter, and as such participate in energy transfers between the biota and the abiotic environment.

2.1.4 Linking drivers of change and ecosystem functioning with the response-effect framework

Efforts to link drivers of change and ecosystem function responses have been disparate across taxonomic groups, with a major focus on plants and invertebrates in the past years. Indeed, Hevia

et al. (2017) showed in a metanalysis that most studies investigating how species traits mediate the impacts of stressors on ecosystem processes focused on plants and invertebrates, such that there is an existing taxonomic bias in this area. Vegetation and invertebrates both represented an approximate 40% of the sampled papers, whereas only 17% were dedicated to vertebrates. Their metanalysis also shed light on other biases, such as the spatial scale of the papers, with most sampled studies being conducted at local or national scales. Therefore, although terrestrial vertebrates have a major cultural, economic and functional importance and are over-represented in the overall biodiversity literature compared to other taxa (Titley et al., 2017), how disturbances affect the services they provide has not been extensively explored compared to other taxa. To understand how anthropogenic pressures may impact ecosystem processes sustained by vertebrate communities at global scales, there is a need to assess whether LUCC significantly affects the functional diversity of vertebrate communities, and, in particular, the effect trait composition; and to verify whether effect trait composition predicts ecosystem processes, as detailed in the previous section (or, alternatively, apply this idea the other way around).

The end-goal of the response-effect framework (Lavorel and Garnier, 2002; McGill et al., 2006; Naeem and Wright, 2003), initially developed for plants, is to understand how environmental changes alter ecosystem functioning using response and effect traits. Indeed, if effect traits inform on ecosystem processes, response traits mediate species responses to environmental change. As such, the link between environmental pressures and ecosystem functioning is conceptually realised with both response and effect traits, when they overlap. The response-effect framework relies on identified response and effect traits to provide a mechanistic understanding of how disturbances modify the trait composition of communities, and how these changes link to alterations in functioning, driven by changes in the effect trait composition.

The application of the response-effect framework to animal communities has been hindered by several issues (Bartomeus et al., 2018; Didham et al., 2016; Luck et al., 2012). For instance, there is a lack of empirical support for response and effect traits in animal communities; results may be contingent to a given taxon, hindering our ability to generalise predictions.

Luck et al. (2012) underlined the need to develop robust and broadly applicable methods for vertebrates. They readapted the response-effect framework, with the aim to make it applicable for vertebrate species and provide guidelines for its application. Currently, our knowledge of how anthropogenic changes will alter the global processes sustained by vertebrate species is extremely limited, due to the diverse reasons exposed previously. One of the end-goal of my PhD project is

to tackle this question at global scales and investigate whether the loss of certain trait combinations in response to LUCC may lead to the disruption of important ecosystem functions. As such, adaptations of the response-effect framework may be particularly relevant to this work.

To conclude, examining how vertebrate species traits influence their responses to LUCC is the first step to (1) elucidate which traits are likely to put species at greater risk, and find out whether it is possible to generalise patterns across vertebrate species (by working at global scales and comparatively across the four terrestrial vertebrate classes); (2) investigate whether future biodiversity declines triggered by these anthropogenic changes are likely to disrupt important ecosystem functions.

The work I have achieved so far focuses on land-use change at global scales and aims at investigating the questions detailed in the next section.

2.2 Questions and hypotheses investigated in the present report

Here, I briefly introduce the questions I tackled in this report. They will be developed in more detail in each corresponding Chapter.

2.2.1 Chapter 3: Collecting and imputing ecological traits across terrestrial vertebrates

As underlined in the introduction, functional traits can provide a mechanistic understanding of how environmental stressors affect both ecological assemblages and ecosystem processes. As such, they convey information most relevant to conservation policies. According to Hekkala and Roberge (2018), global assessments of how land-use change affects vertebrate functional diversity may have been limited so far by the amount of ecological information required to conduct such analyses, notably by the availability of species traits. There exist published databases of species traits, many of which quite recently published (see Table 3.1 in Chapter 3), but despite these collation efforts, some taxa are likely to remain under-sampled. For this project, I collate information on vertebrate traits prior to conducting any analysis (Chapter 3). I assess the gap in trait information across terrestrial vertebrates and investigate whether trait information present taxonomic, phylogenetic and spatial biases. Notably, I hypothesize that:

- Mammals and birds are, overall, better sampled than herptiles;
- Species with larger range sizes are more likely to have more complete trait information;

- Trait data is phylogenetically biased: closely related species are more likely to have a similar amount of available trait information than less related species.

After assessing the gaps in the availability of trait information, I impute missing trait values using random forests algorithms. All traits are used as predictors in the process. Phylogenetic information is incorporated as an extra predictor in the form of phylogenetic eigenvectors. I then evaluate imputation performance and congruence.

Next, the compiled trait dataset is used for further analyses (Chapter 4). In Chapter 4, I address the questions presented in the section below.

2.2.2 Chapter 4: Land-use change promotes the functional homogenisation of local vertebrate communities

The aim of this Chapter is to investigate how land-use change affects the functional diversity of vertebrate communities. Because obtaining longitudinal information on compositional changes can be difficult, the effects of land-use change throughout this project are studied using a ‘space-for-time’ substitution, whereby a spatial gradient is used as a proxy for temporal dynamics (De Palma et al., 2018). As such, the following analyses build upon the PREDICTS database, a large collated dataset of species occurrence and abundance around the world across different land-uses (Hudson et al., 2014, 2017). To date, this database constitutes the most comprehensive global collection of biodiversity samples across different land-uses. It comprises 666 studies, each of which recording the occurrence and/or abundance of species at different sites (abundance in most cases). Each site is classified into a land-use category; land-use categories encompass primary vegetation, secondary vegetation, plantation forest, cropland, pasture and urban. Primary vegetation refers to native vegetation undisturbed since its development under current climatic conditions. Where primary vegetation was destroyed (either by human actions or natural causes), recovering vegetation forms are referred to as secondary vegetation. Secondary vegetation is further divided into three categories: mature, intermediate and young, depending on the stage of recovery of the vegetation. Finally, plantation forest, cropland and pasture refer to agricultural areas (crop trees grown for human purposes, biofuels and herbaceous crops, and areas grazed by livestock).

Using this database, I aim to investigate how land-use change impacts the functional diversity of local vertebrate communities. I hypothesise that by reducing local habitat heterogeneity, human-dominated land-uses promote functional homogenisation and clustering, whereby the similarity in trait composition across assemblages increases through the loss of certain functions. This hypothesis

relies on the idea that strong environmental filtering will disproportionately remove certain functional types. To test this hypothesis, I use various indices of functional diversity. Specifically, I calculate two indices of functional richness (Legras et al., 2018): volume-based functional richness FRic (Villéger et al., 2008) and dendrogram-based functional richness DFR (Petchey and Gaston, 2002). Below, I state the hypotheses for these indices.

- Where functional richness indices are not correlated with species richness, I expect functional richness to decrease in more human-dominated land-uses, with habitat filtering reducing the amount of utilised trait space.
- When species richness is correlated with functional richness, I expect the slope of the species richness–functional richness relationship to be smaller in more disturbed land-uses (this hypothesis links to the hypothesis on functional redundancy presented further down; Figure 2.1).

I also calculate the functional dispersion (Laliberte and Legendre, 2010) of each local community. Finally, I use an index developed by Ricotta et al. (2016) to estimate functional redundancy. Specifically:

- I expect functional dispersion to decrease with increasing land-use disturbance, with species within human-dominated land-use communities presenting more similar trait values, due to functional clustering.
- As a consequence of the previous hypotheses, I expect more disturbed land-uses to have higher degrees of functional redundancy, as the similarity in functional composition increases across species.

These constituted the hypotheses for the work presented hereafter. In Chapter 5, I detail some questions that I aim to investigate in the future years of my PhD. All analyses and data collation were conducted using R (R Core Team, 2018).

3 | Collecting and imputing ecological trait data across terrestrial vertebrates

3.1 Introduction

Planetary anthropogenic threats are reshaping patterns of species diversity (Böhm et al., 2013; Schipper et al., 2008; Spooner et al., 2018; Stuart et al., 2004). Land-use change globally impacts local species richness (Newbold et al., 2015). In turn, species losses can negatively affect ecosystem functioning (Hooper et al., 2005, Hooper et al., 2012). Understanding how increasing pressures will affect ecosystem functioning and the services ecosystems provide is vital to put into place efficient mitigation measures.

The earliest experiments investigating the effect of species richness on the stability of ecosystem processes were conducted in the late twentieth century (Naeem et al., 1994; Tilman and Downing, 1994). Hundreds of grassplot experiments have since then confirmed that higher diversity promotes higher primary productivity and increased ecosystem stability (Balvanera et al., 2006; Tilman et al., 2014). Intuitive mechanistic explanations of this phenomenon include increased niche complementary and resource partitioning, favouring more efficient resource use. Specifically, species properties (or traits) shape how species interact with their biological and physical environment. As such, traits are central to elucidate how species diversity links to ecosystem functions.

Strictly, traits are defined as phenotypic characteristics measurable the level of an individual, with an effect on organismal fitness or performance (McGill et al., 2006; Violle et al., 2007). They can be physiological (e.g., metabolic rates), morphological (e.g., body mass), behavioural (e.g., activity time), phenological (e.g., anthesis); they can also relate to species life-history (e.g., longevity) or diet (e.g., trophic level). Traits shape species fundamental and realised niches; for instance, physiological traits influence species thermal tolerances, participating in defining their geographical distributions

(Calosi et al., 2010; Khaliq et al., 2017). Morphological attributes (e.g. body mass, eye position, etc.) participate in shaping the structure of food webs, and in determining the strength of inter- and intra-specific competition (Gravel et al., 2016; Laigle et al., 2018). As such, traits determine how species use and impact on their environment. Specifically, effect traits underpin species resource use and define organismal contributions to ecosystem functions. Response traits are those involved in determining species responses to environmental changes and can overlap with effect traits. This distinction between response and effect trait led to the development of a conceptual framework, the ‘response–effect’ paradigm (Lavorel and Garnier, 2002; Luck et al., 2012; see Chapter 1), which aims to understand how traits shape species responses to environmental changes, and how these changes in turn affect ecosystem functioning.

A looser, more flexible definition of trait is sometimes adopted. Some characteristics only measurable at the species level can be referred to as ‘ecological’ traits. Examples of ecological traits, only measurable in relation to species occurrence, include habitat preferences (breadth or thermal/moisture preferences). In this chapter, I use this more flexible definition of trait, and consider species ‘ecological’ traits.

Trait-based approaches are increasingly used to understand processes underpinning species co-existence and biodiversity–ecosystem functioning relationships. Notably, they are widely employed in the context of the response–effect paradigm, and publications in this field have increased exponentially since the 2000s (Hevia et al., 2017). Nevertheless, studies investigating how environmental changes are likely to affect species and ecosystem functions non-randomly with respect to species traits are both extremely taxonomically biased in favour of plants and invertebrates (more than 75% of analysed studies in the metanalysis conducted by Hevia et al. (2017)), and spatially biased towards local scales (about 60% of the studies in the metanalysis conducted by Hevia et al. (2017); more than 90% of the studies were conducted at local or national scales). Consequently, our understanding of biodiversity–ecosystem functioning relationships at various spatial scales need to be refined (Isbell et al., 2018; Thompson et al., 2018). Moreover, although terrestrial vertebrates have been extensively studied in the past (Titley et al., 2017), how environmental changes may affect their global contributions to ecosystem functions needs to be investigated further.

Indeed, vertebrates play diverse and important ecosystem roles. Through frugivory, they participate in seed dispersion (McConkey et al., 2012; Mokany et al., 2014; Wandrag et al., 2015). They are significant pollinators in numerous ecosystems (Ratto et al., 2018). Vertebrate herbivores impact global plant diversity patterns through top-down regulation (Lin et al., 2018; Zhang et al., 2018).

They also contribute in regulating animal populations through their predatory activity (Barber et al., 2010; Letnic et al., 2012; Luck et al., 2012; Paine et al., 2016; Salo et al., 2010). As scavengers, they participate in nutrient cycling and energy transfers (Cunningham et al., 2018; Inger et al., 2016; Wilson and Wolkovich, 2011). Moreover, they are culturally important (Albert et al., 2018; Hirons et al., 2016), and a source of protein for many people (Alves et al., 2018).

Understanding how environmental changes may affect their ecological roles is important to predict future ecosystem functioning, and to put into place appropriate mitigation measures. The end-goals of my PhD thesis include elucidating how species traits influence their responses to land-use and climate change at global scales, and how changes in community composition may affect ecosystem functions.

Addressing these questions requires to use extensive trait data. Despite vertebrates having been the focus of much research, and despite the growing interest for trait-based approaches, there exists no comprehensive database of vertebrate ecological traits encompassing all Classes. Consequently, collating trait data was a prerequisite for any further work. This constituted the aim of the current chapter: here, I collected and imputed trait data across the four terrestrial vertebrate Classes (mammals, birds, reptiles and amphibians).

In this chapter, I present the methods I used to collect and impute trait values across terrestrial vertebrates. Thanks to past and recent efforts to release data in the public domain, at least four ecological trait databases are now freely accessible (Table 3.1; Cooke et al. (2019)). Other trait datasets have been released on on-line platforms alongside published articles (e.g. Global Assessment of Reptile Distribution initiative, <http://www.gardinitiative.org/>), or can be downloaded from online databases (IUCN Red List (<https://www.iucnredlist.org/>), BirdLife data zone (<http://datazone.birdlife.org/home>)).

Data collection was constrained by the amount of information available in the literature. All primary sources offered a variety of traits, of which only a few were selected. Trait selection was motivated by two main reasons: (1) traits should be of ecological interest and be related to response or effect processes; (2) trait values should be available for many species, across the four terrestrial vertebrate Classes, allowing for cross-Classes comparative analyses. Targeted traits related to species life-history, morphology, behaviour and feeding habits (body mass; longevity; litter/clutch size; diel activity; trophic level; diet) and to their habitat preferences (habitat breadth and specialisation). Reptilian diet was not readily available in primary data sources, and one exception was made as I extracted diet data for the other Classes. Species mobility was hardly available across sources,

and no common variable could describe species mobility across Classes. Although species' abilities to move in their environment is key in understanding how species will respond to anthropogenic pressures (Barbet-Massin et al., 2012; Pearson, 2006; Schloss et al., 2012b), this trait was not considered for the above reasons. In this chapter, I detail the methodology I employed to collate targeted traits. I elaborate on some of the challenges met when compiling data across many species, such as inconsistency of taxonomy across sources, and problems posed by taxonomic inflation and synonymy.

Despite a wealth of information across primary sources, trait data was likely to be incomplete across terrestrial vertebrates. Many species were likely to present missing trait data for many traits; and taxonomic and geographical biases in the global trait knowledge were likely to exist (González-Suárez et al., 2012; Hortal et al., 2014). The gap in global trait knowledge was termed the 'Raunkiaer shortfall' by Hortal et al. (2014). Here, I assessed the Raunkiaer shortfall for terrestrial vertebrates. I investigated whether trait data presented taxonomic, phylogenetic and spatial biases.

After examining patterns in the gaps in trait data information, I imputed missing trait values. This chapter finally details imputation methodology and examines imputation performance and robustness.

3.2 Methods

3.2.1 Ecological trait data collection

Primary data sources.

I collated ecological trait data for terrestrial vertebrates from the sources figuring in Table 3.1. Information was compiled for the following target traits: body mass, longevity, litter or clutch size, trophic level, diel activity, diet, and habitat preferences. I also compiled traits that were potentially correlated to either body mass or longevity, to be used as potential predictors in imputations of missing trait values. As such, body length information was compiled when available, as well as generation length or age at sexual maturity. Most notably, longevity was chosen over generation length or age at sexual maturity as it was the only common currency across Classes reflecting generation turnover. In addition, species geographical range sizes were estimated from distribution data, extracted from the IUCN Red List of Threatened Species.

Table 3.1: Primary sources used for each compiled trait. Primary sources may contain more traits than shown here. **BM**: body mass; **BL**: body length; **L**: longevity or maximum longevity; **GL**: generation length; **LCS**: litter or clutch size; **TL**: trophic level; **Di**: diet; **DA**: diel activity; **RS**: range size; **H**: habitat data. Bolded abbreviations highlight target traits; other traits were added for potential correlations in further imputations.

Sources	Taxa	Traits									RS	H
		BM	BL	L	MA	GL	LCS	TL	Di	DA		
Oliveira et al., 2017	Amphibians	✓	✓	✓	✓		✓	✓	✓	✓		
Cooper			✓				✓				✓	
Senior			✓									
Bickford			✓								✓	
Wilman et al., 2014	Birds	✓							✓	✓		
Butchart		✓				✓						
Jones et al., 2009	Mammals	✓	✓	✓	✓		✓			✓		
Kissling et al., 2014								✓				
Gainsbury et al., 2018								✓				
Wilman et al., 2014		✓							✓	✓		
Pacifici et al., 2013		✓		✓	✓	✓						
Scharf et al., 2015	Reptiles	✓		✓	✓		✓	✓		✓		
Vidan et al., 2017										✓		
Stark et al., 2018		✓		✓			✓			✓		
Schwarz and Meiri, 2017							✓					
Novosolov et al., 2017		✓						✓			✓	
Novosolov et al., 2013							✓					
Slavenko et al., 2016		✓										
Myhrvold et al., 2015	Amniotes	✓	✓	✓	✓		✓					
IUCN Red List	Vertebrates										✓	✓

Compilation methods.

Continuous traits. All continuous traits were averaged within species when different sources provided estimates. Longevity and maximum longevity were assumed to provide the same information and were averaged within species. No measure of intra-specific variability was compiled and estimates were provided as a single measure for each species, despite intra-specific variation being increasingly recognised as .

Categorical traits.

Activity time. Species were described as being either nocturnal or non-nocturnal. Despite a higher resolution of activity time information in some of the primary sources (e.g. species being described as cathemereal, crepuscular or strictly diurnal), I adopted the classification of the primary source with the lowest resolution, in order to have consistent information across Classes.

Diet and diet breadth. For mammals and birds, diet information was compiled from the EltonTraits database (Wilman et al., 2014). Primary diet was available in the avian dataset and grouped into five categories: (1) plant or seed consumers; (2) fruit or nectar consumers; (3) vertebrate consumers, including fish and carrion; (4) invertebrate consumers; and (5) omnivores. Primary diet was not available for mammals. Instead, mammal diet was described as the percent use of different food items. I pooled these items together into the same five primary diet categories as for the avian dataset. Any food item for which percent use was equal to or above 50% was considered to be part of the primary diet. Species for which no food item had percent use above 50% were considered to be omnivores.

For amphibians, diet information was extracted from AmphiBIO (Oliveira et al., 2017). Diet information was available as binary variables for diverse food items (leaves, flowers, seeds, fruit, arthropods and vertebrates). Percent use was not recorded, so all items listed for a given species were considered to form species primary diet. I pooled amphibian diet into the five diet categories described above.

Trophic level. For amphibians and birds, trophic levels were partly inferred from the primary diet.

Habitat preferences. Species habitat preferences were compiled from habitat data files provided by the IUCN Red List. They were described as a binary variable recording whether a species was known to occur in a particular habitat. I calculated habitat breadth as the number of habitats a species was known to use. Weights were assigned to each habitat in this calculation depending on the recorded suitability and importance of the habitat; outcomes were not very sensitive to the presence of weights (compared to a non-weighted sum, see SI). Finally, a broad degree of habitat specialisation was produced. If any artificial habitat was recorded to be suitable, species were reported to be generalists; else, they were natural habitat specialists. More details on habitat preferences compilation are provided in the SI.

3.2.2 Phylogenetic information

I obtained phylogenetic trees for birds, amphibians, mammals and squamates from Hedges et al. (2015). Hedges et al. (2015) built a time-calibrated phylogenetic tree representing more than 50,000 species, using meta-analytic methods: results from 2274 phylogenetics and molecular evolution studies were assembled to build a ‘Super Time’ tree of life. Tree subsets for diverse clades are available at <http://www.biodiversitycenter.org/ttol> (downloaded 06/07/2018). The trees for vertebrate species were all ultrametric and fully resolved, except for the amphibian tree which presented polytomies. All trees contained a few branches of length 0 (193 branches for mammals, 136 for amphibians, 189 for birds and 284 for reptiles). Using trees that were built using molecular data, and not life-history traits, allows to avoid circularity in further imputations of missing trait values.

3.2.3 Tackling taxonomic synonymy

Across the different primary sources, similar species could appear under different binomial names. This was a problem when matching datasets by species. It was also problem when matching species to the PREDICTS database. Moreover, it is possible that within a primary source, a given species was appearing under two or more different names. As such, taxonomic synonymy created ‘pseudoreplicates’ of the same species, overall falsely increasing the total number of species and artificially inflating the amount of missing trait values. Taxonomic synonymy was hence a major issue; due to the large number of species across datasets, extensive manual checks could not be applied. The presence of typos in species names had the same effect as synonymy, erroneously duplicating species. I attempted to correct for taxonomy first by correcting for typos, and second by identifying species which were entered under a synonymic name and replacing these with the accepted name. To this end, I developed an automated procedure, complemented with a few manual entries. Obvious cases where vernacular names had been entered in the place of binomial names were also treated manually; that was the case for 44 PREDICTS species (when possible, I best assigned binomial names to species common names; unidentifiable species were left empty and assigned to a genus (5 species)).

Automated procedure and outputs.

Extracting names from the IUCN Red List and the Integrated Taxonomic Information System (ITIS). The automated procedure consisted of extracting species accepted and synonymic binomial names from the IUCN Red List or from the ITIS, using the `reddlist` and `taxize` R packages

(Chamberlain, 2018, Chamberlain and Szöcs, 2013). I started by generating a list of all names figuring across datasets (primary sources, phylogenies and PREDICTS). These ‘original’ names were corrected for typos (using `gnr_resolve` function, `taxize` package (Chamberlain and Szöcs, 2013)); then, the IUCN Red List was queried and synonyms and accepted names were stored when possible. When species were not found in the IUCN Red List, information was extracted from the ITIS. When species were not found in the ITIS either, corrected names (original names corrected for typos) were assumed to be accepted. Family and Order information was extracted using the same procedure and some entries were completed using the Global Biodiversity Information Facility taxonomic backbone (GBIF, <https://www.gbif.org/tools/species-lookup>).

NB: for species entered with the forms *Genus cf.*, *Genus aff.* or *Genus spp.*, the accepted binomial name was left empty.

Outputs. I generated a list of vertebrate species, recording whether species names were accepted or synonymic (for 14124, 8743, 6090, and 11183 names or identifiers found across datasets for birds, amphibians, mammals and reptiles respectively, including species names as they appeared in phylogenetic trees). For each name, the identified accepted name and the synonyms were stored when possible, as well as additional taxonomic information (Order, Family, Genus). When queries did not succeed, species accepted names were assumed to be the original names found in the datasets (corrected for typos).

Harmonising taxonomy in trait datasets. Taxonomy across datasets was finally homogenised by replacing recorded synonyms with their accepted scientific names. Overall, this procedure reduced the total number of species figuring in trait datasets (Figure 3.1). The species presenting the highest degree of ‘pseudoreplication’ was the East African mole rat (*Tachyoryctes splendens*), which was figuring under 12 names identified as being synonymic across primary sources (Figure 3.1B), highlighting the need for normalising taxonomy across sources.

Despite the automation efforts, taxonomic redundancy persisted to a degree in the trait datasets. Indeed, at this stage, not all species in PREDICTS matched a species in the trait datasets. Additional manual inputs were required to resolve taxonomic synonymy for these species. Verifying the presence of PREDICTS species in trait datasets was important for further analyses. Taxonomic synonymy was resolved manually for 91 PREDICTS species that did not match any species in the trait datasets; in that case, information was extracted from other diverse sources

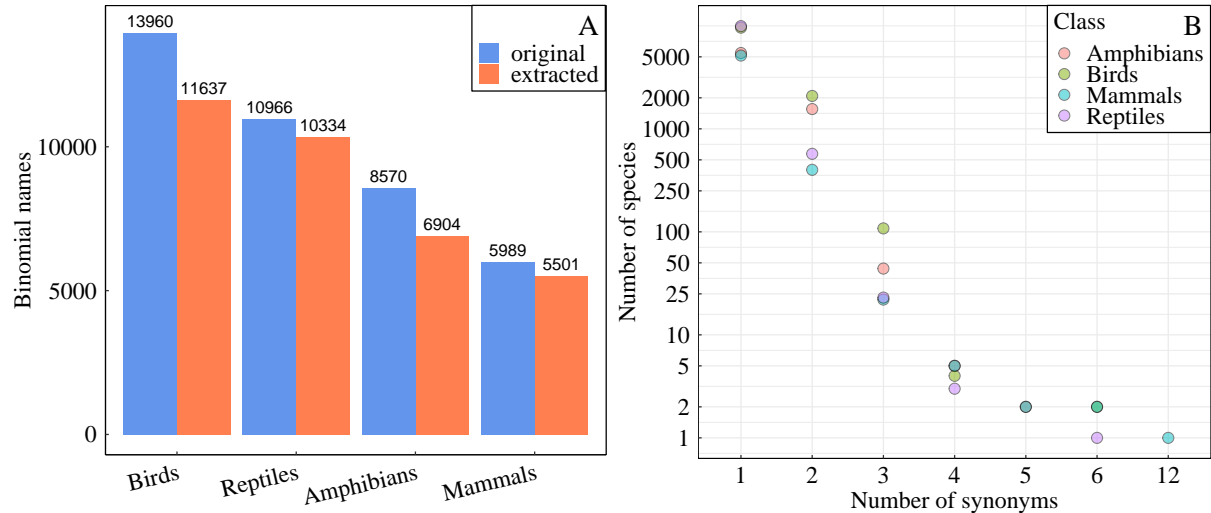


Figure 3.1: Difference in species number due to taxonomic correction (A) and distribution of number of synonyms across datasets (B). (A) shows the number of species across all primary sources (trait datasets and PREDICTS, excluding phylogenies), before and after correcting for taxonomy. Replacing identified synonyms by the extracted accepted name reduced the number of species in all Classes, with the most drastic reduction for birds (decrease by 2,323 unique binomial names). The diminution was of 632 unique identified species for reptiles, of 1,666 for amphibians and of 488 for mammals. (B) shows the distribution of the number of synonymic names. In all four Classes, more than 5,000 species (or binomial names) had no identified synonyms. Nevertheless, a large amount of species had two identified synonyms (range: 400 species for mammals - 2086 for birds). The most replicated species was the East African mole rat *Tachyoryctes splendens*, for which 11 synonyms were identified.

(such as the Reptile Database (<http://www.reptile-database.org/>); Avibase (<https://avibase.bsc-eoc.org/avibase.jsp?lang=EN&pg=home>); AmphibiaWeb (<https://amphibiaweb.org/>)). After adding manual inputs to the synonym datasets, all PREDICTS species were represented in trait datasets.

The need to apply additional manual inputs underlines the fact that the automated procedure was not optimal. The Red List and the ITIS were not comprehensive taxonomic sources, and for clades with high degrees of pseudoreplication in names, such as reptiles or amphibians, neither the Red List or the ITIS contained enough information. As I only applied manual checks for species in the PREDICTS database, ‘pseudoreplication’ and taxonomic errors are likely to have persisted to a degree. Moreover, certain species were entered using the format *Genus subspecies* rather than *Genus species*; for these, automated queries may have failed to identify the species.

Harmonising taxonomy in phylogenetic trees and increasing species phylogenetic representation.

Taxonomic correction across tip labels. Efforts to correct datasets for taxonomy created problems for a small proportion of species when dealing with phylogenies. The idea of the procedure described above was to replace two or more identified synonyms by a single accepted name, and then

collapsing dataset rows together by names. I applied the same method on phylogenies, replacing synonyms by their identified accepted names in trees' tip labels. Not unexpectedly, in some cases, the procedure ended up assigning the same accepted name to different phylogenetic tips. This was the case for 2.8% of mammalian, 1.7% of avian, 1.6% of amphibian and 1.7% of reptilian species, which then had multiple phylogenetic positions (most having two different positions, see SI). Because keeping several putative phylogenetic positions for a species was problematic in further analyses, I selected one tip to conserve and dropped other tips from the phylogenies (Figure 3.2). To briefly describe the procedure, 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. Further details on how replicated tips were dropped are available in the SI (with 3 examples for each case of Figure 3.2).

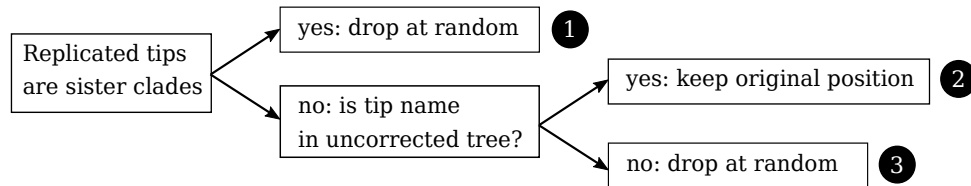


Figure 3.2: Procedure followed to drop replicated tips from phylogenies. Most of these were replicated twice. When replicated tips were sister clades, the tips to drop were chosen randomly, as it did not affect the ‘true’ phylogenetic position of the species (1). When replicated were not sister clades, I kept the tip whose position was closest to the position of the same tip in the uncorrected tree (2). 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 (3). Nevertheless, occurrences of that third case were rare (see SI).

Correcting for taxonomy in the phylogenies: conclusions. Figure 3.3 shows the phylogenetic representation of species figuring in the trait datasets. Overall, correcting for taxonomy improved species representation in the trees. For amphibian and reptilian species figuring in PREDICTS only, phylogenetic representation disproportionally increased (with a minimum representation of 76% for PREDICTS amphibians after correcting the trees for taxonomy, inset plot in Figure 3.3). Nevertheless, correcting phylogenetic tip labels generated replicates for a marginal number of tips, which then had to be dropped.

Species attachments to phylogenetic trees. Some species in the trait datasets were not represented in the phylogenies, even after taxonomic corrections (3.3). Maximising the number of species represented in the phylogenies was important for further trait imputations. Indeed, if traits were evolutionary conserved, species phylogenetic position could be an important predictor of trait values.

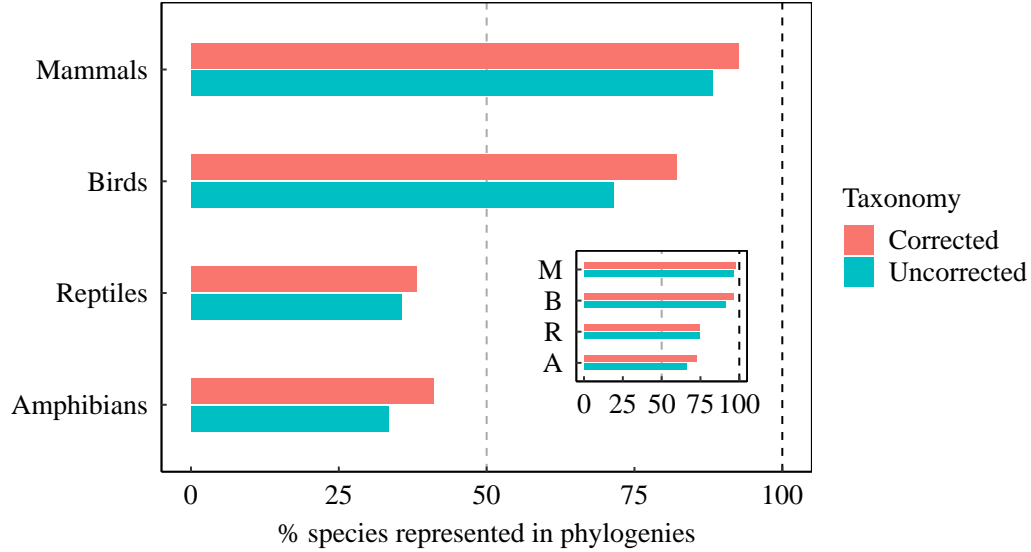


Figure 3.3: Percentage of species represented in the phylogenies, with and without taxonomic corrections. Overall, taxonomic correction increased species representation in phylogenetic trees. Representation for mammals and birds was high (after taxonomic correction: 82% of avian and 93% of mammalian species had a phylogenetic position). On the other hand, reptiles and amphibians were poorly represented (after taxonomic correction: only 38% of reptilian and 41% of amphibian species were placed in phylogenetic trees). The inset barplot shows representation for species figuring in PREDICTS. For these, species presence in phylogenetic trees after correction was high across all Classes, with a minimum representation of 76% for amphibians.

To maximise species representation, I attached non-represented species to the root of their genus, when possible (phytools package, Revell, 2016). Attaching species at the root of their genus created polytomies, which were resolved randomly (using multi2di in ape (Paradis and Schliep, 2018) and bifurcatr in PDcalc (Nipperess and Wilson, 2019)). Resulting trees contained additional branches of length zero. Such modifications of the phylogenetic trees could have altered the significance and the strength of trait phylogenetic signal. I further verified whether these alterations of the trees had impacted phylogenetic signal, by qualitatively comparing the strength and the significance of phylogenetic signal for each trait, estimated using both original trees and modified trees (see ‘Assessing phylogenetic signal in traits’).

Finally, a large number of species were attached to their genus in the trees (Table 3.2). For instance, only 38% of the species figuring in the reptilian trait dataset were initially found in the squamate phylogeny. After attaching non-represented species, 91% of the species were placed in the squamate phylogeny.

Table 3.2: Species representation in phylogenetic trees (datasets corrected for taxonomy). The number of species attached to the root of their genus ranged from 175 (mammals) to 5438 (reptiles). Finally, most species were represented in the phylogenies, whereas more than half reptilian and amphibian species initially had no known phylogenetic position.

Class	Initially not in tree	Of which randomly attached	No final representation in tree
Amphibians	59% (4040 of 6904)	96% (3883 of 4040)	2.3%
Birds	18% (2085 of 11637)	75% (1574 of 2085)	4.4%
Mammals	7.4% (407 of 5502)	43% (175 of 407)	4.2%
Reptiles	62% (6391 of 10334)	85% (5438 of 6391)	9.2%

3.2.4 Investigating biases in the coverage and completeness of trait information across Classes

Taxonomic biases.

Having normalised taxonomy and compiled trait data, I assessed trait coverage, defined as the percentage of species for which trait information was available for a given trait. I also estimated the amount of trait information available for a species by calculating trait completeness. For a species, trait completeness was defined as the proportion of traits for which information was available (number of non-missing trait values divided by total number of traits). In corrected datasets, species with 0% completeness in predictor traits were filtered out. I tested whether taxonomic Class impacted trait completeness using pairwise Kruskal-Wallis rank sum tests (the null hypothesis tested in each pair was that the distribution of completeness values were sampled from the same original distribution).

Phylogenetic biases.

Whether values are missing at random is likely to impact imputation errors, notably if some clades appear to be under-sampled. Further, I examined whether patterns in the distribution of missing values emerged within Classes, as particular clades or parts of the phylogenies could be under-sampled compared to other clades. To assess whether missing values presented patterns, I represented within-Family median completeness and within-Family median coverage values in each branch of phylogenetic trees built at the Family level. Tree branches were colour-coded to reflect median values in each Family (using contMap, phytools package, Revell, 2016). Specifically, within-Family trait completeness was calculated by aggregating species into their families and calculating the median trait completeness within each group.

Patterns of missing values in trait coverage were explored for each trait separately. Trait coverage was assessed within families as the number of species for which values were missing over the total number of species in each Family. As families represented by very few species might present higher percentages of missing values, reflecting Family size rather than randomness in sampling, I contrasted trait coverage plots against a plot showing how much each Family contributed to the total number of species (number of species in each Family over total number of species in the tree).

Spatial biases.

I finally investigated whether trait completeness was spatially biased. Specifically, I tested the hypothesis that bigger geographical range sizes were correlated with better trait completeness. To that end, I fitted a generalised linear model with a Poisson error distribution: trait completeness was treated as count data (number of sampled traits). Class was added as an explanatory factor, interacting with range size. The model was written as: $\text{Completeness} \sim \log(\text{RS}) + \text{Class} + \log(\text{RS}):\text{Class}$, and I specified a Poisson error distribution. I examined whether the fit of the model was good using a chi-squared test on the residual deviance.

NB. In all of the above, completeness was calculated over all predictor traits for each Class (target traits and supplementary traits). As such, the total number of traits sampled in each Class was not necessarily equal. The same analyses could be replicated on the set of target traits only. Analyses of spatial biases in trait sampling could also be developed in the future (see Chapter 5, Outline of future work).

Finally, I investigated whether correcting for taxonomy had an effect on trait completeness using Wilcoxon rank sum tests. I tested whether the median trait completeness was significantly higher for datasets corrected for taxonomy, than for uncorrected datasets, in each Class.

Conclusion: imputing missing values to increase coverage. Trait coverage was highly variable across Classes and traits (see Results). Trait coverage for species figuring in the PREDICTS database only overall improved compared to trait coverage for the whole set of species, particularly for reptiles and amphibians (see SI). Nevertheless, no trait reached 100% coverage in any Class. Obtaining trait estimates for all of PREDICTS species was important, as otherwise, each species for which trait values were missing would have to be dropped in further analyses. Moreover, within-

Class biases in availability of trait information appeared (see Results). Consequently, dropping missing-value species could skew trait distributions and generate biases in further analyses. As such, rather than dropping missing-value species, I aimed to fill coverage gaps by imputing missing trait values.

3.2.5 Imputing missing trait values

In order to achieve full coverage across Classes, I imputed missing trait values. Diverse imputation methods have been developed and used in published articles (Cooke et al., 2019, Molina-Venegas et al., 2018, Swenson, 2014). Penone et al., 2014 assessed the performance of four different imputation approaches (K-nearest neighbour (kNN, Troyanskaya et al., 2001), multivariate imputation by chained equations (mice, van Buuren and Groothuis-Oudshoorn, 2011), random forest algorithms as implemented in R by missForest (Stekhoven and Bühlmann, 2012, Stekhoven, 2016) and phylogenetic imputations implemented with PhyloPars (Bruggeman et al., 2009)). Their study showed that the kNN approach resulted in significantly higher imputation error rates than the three other approaches. Both missForest and phylopars were the best methods when phylogenetic information was included. Nevertheless, phylopars was much slower than missForest, and could only handle continuous traits. missForest was faster and could deal with mixed type data. Without phylogenetic information, mice was found to be the best method, with fast imputations of mixed-type data. Of all these methods, missForest was the only one that did not make assumptions about data distribution (being a non-parametric approach), or that did not require a prior knowledge of some tuning parameters. As such, missForest appeared to be a robust option for missing data imputation. To further assess whether to use random forests rather than multivariate chained equations, I estimated the amount of phylogenetic signal in traits. Strong phylogenetic signal in traits would indicate that missForest could perform better than mice.

Assessing phylogenetic signal in traits.

Measuring phylogenetic signal in continuous traits with Pagel's λ . Phylogenetic signal is a measure of the tendency of closely related species to resemble each other more than less related species. Diverse statistics have been developed to estimate phylogenetic signal, most of them applying to continuous traits (Münkemüller et al., 2012). Here, I used Pagel's λ (Pagel, 1999), estimated with the R function `phylosig` (phytools package, Revell, 2016), to assess the amount of phylogenetic signal in continuous traits. Pagel's λ is a scaling component that measures the transformation that

should be applied to the phylogenetic tree for a trait to have evolved under a pure Brownian motion model of evolution (Münkemüller et al., 2012). Under a Brownian motion model of evolution, changes in trait values happen at random along the branches and trait variance is proportional to evolutionary time. λ is then close to zero: the trait covariance matrix is scaled down and the tree loses its internal structure. When λ equals one, both the phylogeny and the trait covariance matrix remain unchanged and the structure of the tree explains trait evolution. As such, λ values close to one indicate that trait values are more similar in closer related species.

Using Pagel’s λ , I assessed the strength of the phylogenetic signal. The `phylosig` function (`phytools`) also allowed to test for signal significance (comparing the estimated λ to the null expectation of λ with a log-likelihood ratio test).

Measuring phylogenetic signal in categorical traits with δ (Borges et al., 2018). Very few methods have been developed to measure and test phylogenetic signal in categorical traits. Fritz et al., 2009 introduced the D -statistic; nevertheless, D is based on a discretisation of categorical traits, which reduces them to binary variables. Borges et al., 2018 introduced a new statistic, called δ , to measure phylogenetic signal in categorical traits of all types. Their approach uses Bayesian inferences to reconstruct trait evolution, that is, to infer trait values in ancestral nodes of the phylogeny. The underlying idea is that the better the phylogeny explains trait evolution, the lower the uncertainty in ancestral state inferences. As such, δ relies on the quantification of the uncertainty associated with the reconstruction of ancestral states. δ can take any positive number, with higher values indicating stronger signal. To test for the significance of the signal, the authors propose to compare the estimated value of δ with the null expectation of δ .

I estimated phylogenetic signal in categorical traits with the δ statistic; implementation used the R code provided by Borges et al., 2018 (https://github.com/mrborges23/delta_statistic). To test for the significance of the signal, I generated null distributions of δ for each trait by randomising trait vectors 50 times (simulating Brownian motion model of trait evolution), and calculating δ for each randomised vector. I then calculated the median of simulated δ values as well as 95% confidence intervals. I tested whether the null medians were significantly lower than the observed value of δ using one-sided Wilcoxon rank sum tests. I noted that the function developed by Borges et al., 2018 could not be implemented if phylogenetic trees contained branches of length 0. As both original and corrected phylogenies contained 0-length branches, I added a very small number to these (10^{-10}) to remedy to this issue and to test for phylogenetic signal.

Significant phylogenetic signal in all traits. All traits showed significant phylogenetic signal (Table 3.3 and SI for p-values of statistical tests), although the strength of the signal was variable across Classes and traits. Overall, modifying the original phylogenies by correcting for taxonomy and by attaching species to the root of their genus did not, qualitatively, have a strong impact on the signal (Figures 3.4 and 3.5), although differences were bigger in reptiles and amphibians, where more than 80% of missing species were added to phylogenetic trees. In mammals and birds, phylogenetic signals remained similar. On the other hand, the stronger effects were observed for reptilian body mass, where adding species to the tree lowered the strength of the signal, and for amphibian trophic level, where the opposite effect was observed.

Phylogenetic signals in categorical traits were all highly significant (Figure 3.5; p-values for Wilcoxon signed rank test: see SI). The strength of the signal differed across Classes and traits, with diel activity, trophic level and primary diet showing particularly strong signal in mammals and birds. Reptiles also showed strong signals for diel activity and trophic level. For amphibians, the results were more even across traits, and still highly significant. Overall, the signal for habitat specialisation was less strong.

Most mammalian continuous traits had very strong phylogenetic signal ($\lambda \geq 0.9$), except habitat breadth ($\lambda \approx 0.7$). In birds, both habitat and diet breadth showed weaker signal, but other continuous traits were highly conserved across closely related species. For amphibians and reptiles, signal strength was much more variable, which may be due to poorer initial trait coverage across phylogenetic tips (see Results). Nevertheless, body length showed high signal in both these Classes ($\lambda \geq 0.9$).

Table 3.3: Phylogenetic signal in continuous and categorical traits and in range size. 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 , after traits were log-10 transformed to improve normality. 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). '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, GL, SM, LCS, and BL being particularly strong (above 0.8 across the four classes). Here all calculations were conducted with the corrected phylogenies, after species were added at the root of their genus. See SI for phylogenetic signals computed with the original phylogenies.

Class	Continuous target traits, additional predictors and range size: λ									Categorical traits: δ			
	BM	L	LCS	HB	DB	GL	BL	SM	RS	TL	PD	DA	Sp
Mammals	0.99*	0.93*	0.97*	0.71*	0.99*	0.98*	0.99*	na	0.76*	17*	50*	19*	1.4*
Birds	0.98*	0.83*	0.93*	0.59*	0.49*	0.98*	na	na	0.70*	10*	18*	28·10 ³ *	1.6*
Reptiles	0.88*	0.93*	0.86*	0.45*	na	na	0.96*	0.93*	0.69*	4.3*	na	7.1*	1.5*
Amphibians	0.97*	0.83*	0.94*	0.81*	0.78*	na	0.95*	na	0.82*	18*	3.7*	2.9*	3.6*

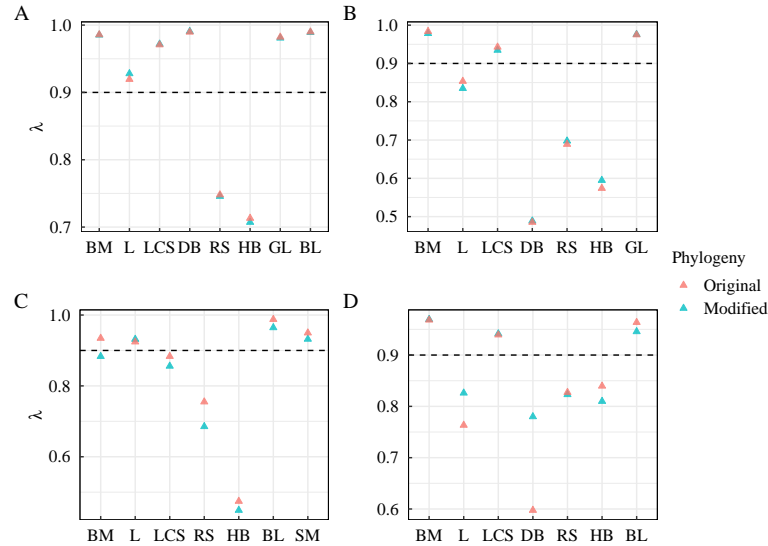


Figure 3.4: Phylogenetic signal in continuous traits (Pagel's λ) estimated with both original phylogenies and modified phylogenies. (A) Mammals; (B) birds; (C) reptiles and (D) amphibians. Overall, altering the phylogenies by correcting for taxonomy and by increasing species representation did not have an important effect on λ .

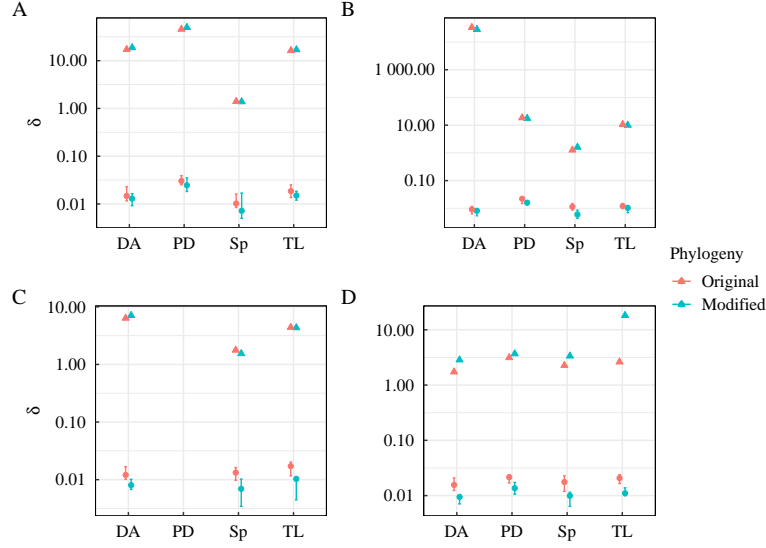


Figure 3.5: Phylogenetic signal in categorical traits (δ) estimated with both original phylogenies and modified phylogenies. (A) Mammals; (B) birds; (C) reptiles and (D) amphibians. Triangle-shaped points represent the estimated phylogenetic signal in each trait; round-shaped points represent the median null expectation of the phylogenetic signal ($\pm 95\%$ CI). Alterations of the phylogenies did not strongly impact δ .

Missing trait values: imputation implementation

Despite much variation in trait coverage across Classes (see Results), results indicated strong phylogenetic signal in many categorical and continuous traits (Table 3.3). I hence imputed missing trait values using random forest algorithms, implemented by missForest. As stated above, missForest was shown by Penone et al. (2014) to be the best method when including phylogenetic information for mixed-type variable imputations. Moreover, Penone et al. (2014) also showed that adding phylogenetic information did not, in any case, decrease the accuracy of imputations.

Phylogenetic relationships were included as additional predictors in the form of phylogenetic eigenvectors (Diniz-Filho et al., 2012), extracted from the phylogenies using the PVR package (Santos, 2018). In this package, phylogenetic eigenvectors were computed from a phylogenetic distance matrix, and calculated using principal coordinate analysis methods. Phylogenetic eigenvectors summarised the relationships among species. The first set of eigenvectors reflected larger distances among species, capturing divergences closer to the root (Diniz-Filho et al., 2012). Penone et al., 2014 showed that including the first 10 eigenvectors minimised the imputation error when imputing missing trait values with missForest. As such, I included the first 10 eigenvectors as additional predictors of missing trait values.

As not all species were represented in the phylogenies (Figure 3.3), I also added taxonomic Orders

as an extra predictor variable in the random forest algorithm. All traits in Table 3.1 were included in the imputations (except for primary diet and diet breadth in reptiles). Tuning parameters of `missForest` were set to 10 maximum iterations (if the stopping criterion was not met beforehand, see below) and to 100 trees grown in each forest. To further examine imputation robustness and consistency, I imputed eight datasets in parallel (eight imputed trait datasets for each Class: total of 32 imputed datasets).

Imputation error and robustness

Out-of-bag imputation error. To assess imputation accuracy, I used the ‘out-of-bag’ error (OOB error) returned by the `missForest` function. The `missForest` algorithm proceeds iteratively, training random forests on observed values first, then predicting missing values over several iterations. The stopping criterion is met when the difference in imputed values (Δ_{cont} and Δ_{cat} , see below) between the last imputed dataset and the previously imputed dataset increases for the first time. The penultimate imputed dataset is then returned. For continuous variables, the difference Δ_{cont} is defined as:

$$\Delta_{cont} = \frac{\sum_{j \in N} (X^{i,l} - X^{i,p})^2}{\sum_{j \in N} (X^{i,l})^2}, \quad (3.1)$$

where j is a continuous trait among N traits, $X^{i,l}$ is the last imputed dataset and $X^{i,p}$ is the penultimate imputed dataset. Δ_{cont} is a measure of the aggregated distance between two successive imputations across all continuous traits. For categorical variables, the difference Δ_{cat} is:

$$\Delta_{cat} = \frac{\sum_{k \in F} \sum_j J_{X^{i,l} \neq X^{i,p}}}{n(NA)}, \quad (3.2)$$

where k is a categorical trait among F categorical traits, $n(NA)$ is the number of missing values for k and J is the j^{th} imputed values for which the consecutive imputations predicted contradicting results. In other words, Δ_{cat} measures the proportion of values that were found to be different between two successive imputations (see Stekhoven and Bühlmann, 2012 for more details).

When the stopping criterion has been met, OOB imputation errors are estimated. OOB errors refer to errors estimated from sub-samples of the data (bootstrap datasets, on which models are trained). OOB errors are estimated from these bootstrap datasets and as such differ from ‘true’ imputation errors, which require previous knowledge of the full dataset. The true root-mean square

error (root-MSE) for continuous traits is defined as:

$$\sqrt{\frac{\text{mean}((X_t - X_i)^2)}{\text{var}(X_t)}}, \quad (3.3)$$

where X_t is a vector of the complete trait values and X_i a vector of the imputed trait values (Stekhoven 2011). With the OOB error, when the complete trait data is not provided, the MSE is calculated from the bootstrap datasets. For categorical traits, the OOB PFC is calculated as the PFC (Δ_{cat} , Equation 3.2), using the bootstrap sub-samples. Breiman, 2001 showed that OOB estimates provide accurate proxies of the true imputation error.

To assess imputation accuracy, I retrieved OOB imputation errors (OOB root-MSE and PFC) across the eight imputed trait datasets in each Class. I plotted the mean root-MSE and the mean PFC across the imputed datasets, as well as the range in errors (maximum error values and minimum errors values across all imputed datasets).

Imputation congruence. To further assess whether imputations were robust, I investigated whether similar values were imputed across the eight datasets in each Class, or in other words, whether results were congruent across the imputed datasets. My expectation was that, for a trait, values imputed independently in different rounds should be nearly identical if imputations were robust. As such, for a continuous trait, pairwise correlations coefficients should be high across the eight datasets (Pearson correlation coefficients for the same trait imputed in pairwise independent rounds, see Table 3.4). For categorical traits, the random forest should predict the same values across the eight datasets.

Table 3.4: Conceptual design for examining imputation congruence for continuous traits. For one trait, pairwise correlation coefficients across eight independent imputation rounds are expected to be high if imputation are robust. To assess imputation congruence across eight imputed datasets, pairwise correlation coefficients were averaged (and the spread assessed using the range).

	Imputed 1	Imputed 2	Imputed n
Imputed 1	1	-	-
Imputed 2	corr(1,2)	1	-
Imputed n	corr(n,1)	corr(n,2)	1

For continuous traits, I assessed imputation congruence across the eight imputed datasets by averaging pairwise Pearson correlation coefficients and plotting the mean (and range) for each trait. For categorical traits, I assessed congruence by assessing the percentage of species for which all eight

imputed values were identical.

3.3 Results

3.3.1 Outputs

I collected and imputed data for 10 traits across 11637 avian species, 5502 mammalian species, 10334 reptilian species and 6904 amphibian species. Datasets recording species accepted and synonymic binomial names are available alongside the trait data.

3.3.2 Biases in the availability of trait information: non randomness in coverage and completeness and patterns in missing trait values

Increases in coverage and completeness due to taxonomic corrections.

Figure 3.6 shows the trait coverage within each Class and for each trait, before and after correcting for taxonomy. Figure 3.7 shows the distribution of trait completeness before and after taxonomic corrections, as well as the median trait completeness for each Class. Across all Classes, correcting for taxonomy increased trait coverage (Figure 3.6). Nevertheless, the increase in coverage for reptiles was marginal, which may indicate that the procedure developed to extract and identify accepted names overall performed less well for reptilian species than for mammals, birds and amphibians. Similarly, correcting for taxonomy improved trait completeness in all Classes (Figure 3.7). Wilcoxon rank sum tests, testing the null hypothesis that uncorrected and corrected completeness distributions came from the same population, rejected this hypothesis across all Classes (alternative hypothesis: uncorrected medians were lower than corrected medians; mammals: $p\text{-value}=1.2\cdot 10^{-9}$; birds: $p\text{-value}<2.2\cdot 10^{-16}$; reptiles: $p\text{-value}=0.025$; amphibians: $p\text{-value}<2.2\cdot 10^{-16}$). To conclude, correcting for taxonomy had a significant impact on trait completeness, and increased coverage in most cases.

Taxonomic biases in the availability of trait information: exacerbated Raunkiær shortfall in reptiles and amphibians.

Trait coverage. Trait coverage was highly variable across Classes and traits. Trait coverage was initially good for most mammalian and avian traits, which had more than 50% coverage (Figure 3.6 A and B). Only longevity had a coverage lower than 50% for these Classes, although generation length was above 80% in both cases. Conversely, trait coverage was overall much poorer for reptiles

and amphibians (Figure 3.6 C and D). About two-thirds of amphibian and reptilian traits presented a coverage below 50%. Amphibians and reptiles appeared to be less sampled in all traits, except in body mass (reptiles) and in body length, range size and habitat variables (amphibians). As such, contrasting patterns of trait coverage appeared between, on the one hand, mammals and birds, and on the other hand, amphibians and reptiles. For species found in PREDICTS only, coverage increased disproportionally in reptiles and amphibians compared to the coverage for the full set of species (the figure for PREDICTS species only is available in the SI). To conclude, trait coverage revealed important taxonomic biases, with higher resolution of trait information across mammals and birds. A clear contrast in trait information appeared between mammals and birds versus herptiles, highlighting the existence of taxonomic biases in data collection.

Trait completeness. Trait completeness reflected similar biases as trait coverage (Figure 3.7). The median completeness with taxonomic correction was high for mammals and birds (92% and 82% respectively) but much lower for reptiles and amphibians (30% and 36% respectively). A pairwise Kruskal-Wallis rank sum test rejected the hypothesis that completeness distribution across Classes originated from the same distribution ($p\text{-values} < 2 \cdot 10^{-16}$ in all cases), showing that Class had a significant effect on the availability of trait information.

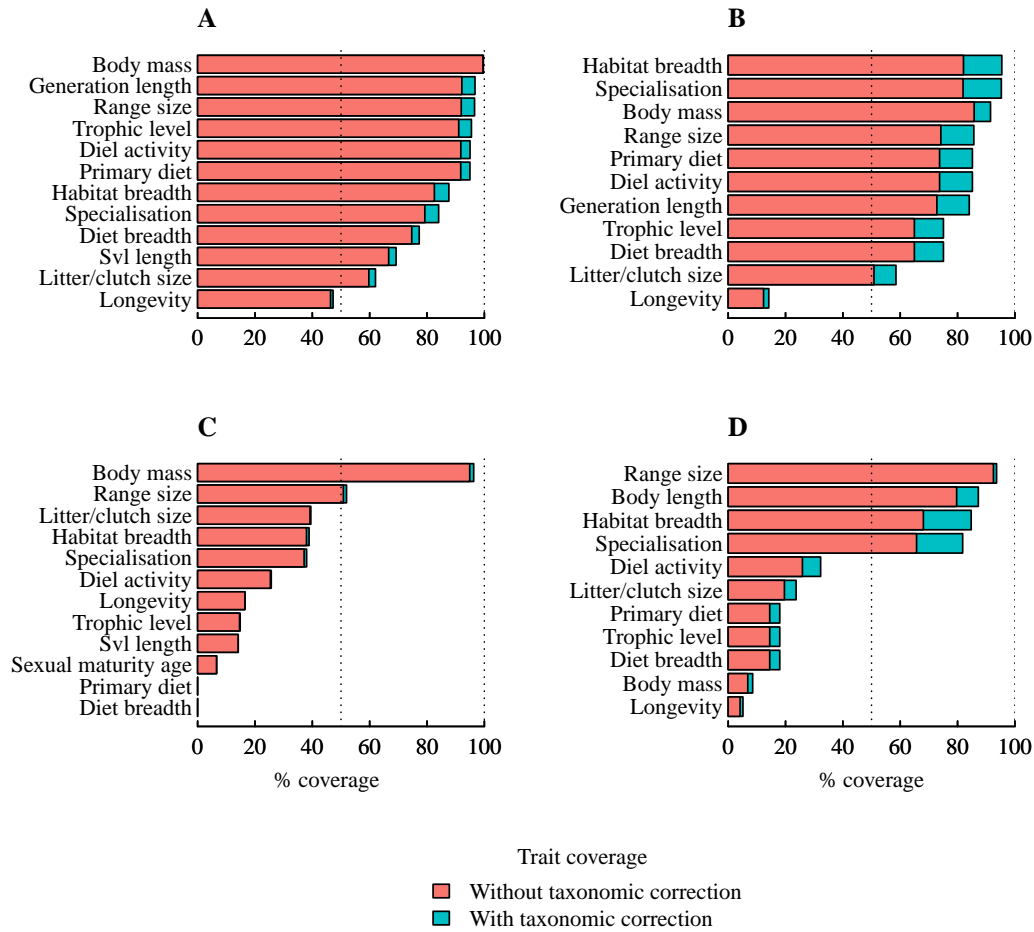


Figure 3.6: Trait coverage across all species before and after taxonomic correction. Here are shown target traits as well as a few other traits used in imputations as additional predictors (such as generation length for mammals and birds or body length for amphibians). **(A)** Mammals (5885 species before correction, 5502 after correction); **(B)** birds (13554 species before correction, 11637 after correction); **(C)** reptiles (10722 species before correction, 10334 after correction) and **(D)** amphibians (8643 species before correction, 6904 after correction). Trait coverage was calculated as the percentage of species for which trait information was available. Correcting for taxonomic synonymy improved coverage in most cases. For mammals and birds, all traits had an initial coverage of more than 50%, except longevity (but generation lengths were estimated for most species). On the other hand, trait coverage was poor (below 50%) for about two thirds of collected reptilian and amphibian traits.

Non-randomness in trait information: within-Class phylogenetic biases.

Within-Class patterns of trait completeness. Figure 3.8 shows within-Family trait completeness for each Class, colour-coded in the tree branches. For better visualisation, the trees are represented without tip labels. Figures providing tip labels are available in the SI (for each Class, tip label information includes taxonomic Order and Family). As expected from the distribution of completeness values for mammals and birds, within-Family completeness was high across most branches of the trees. In mammals, Chiropteras appeared to have lower median trait completeness than other

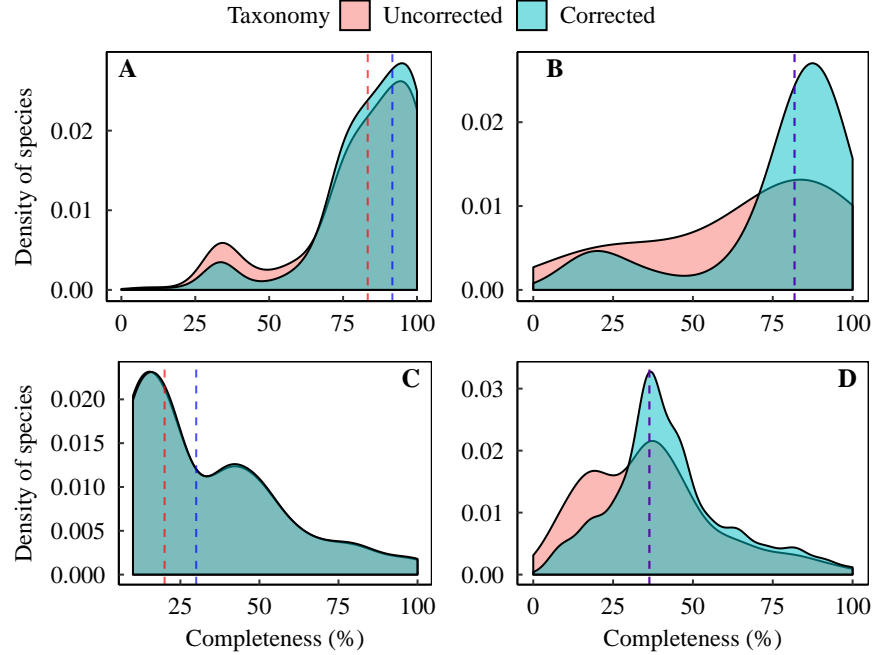


Figure 3.7: Distribution of completeness of trait information across species. (A) Mammals; (B) birds; (C) reptiles and (D) amphibians. Completeness was calculated here for the same set of traits shown in Figure 3.6 (all predictor traits). Correcting for taxonomy affected completeness, significantly shifting the distributions to the right (alternative hypothesis, Wilcoxon rank sum tests: uncorrected medians were lower than corrected medians; mammals: $p\text{-value}=1.2\cdot 10^{-9}$; birds: $p\text{-value}<2.2\cdot 10^{-16}$; reptiles: $p\text{-value}=0.025$; amphibians: $p\text{-value}<2.2\cdot 10^{-16}$). Class had a significant effect on median trait completeness (a pairwise Kruskal-Wallis rank sum test rejected the null hypothesis that completeness distributions across Classes originated from the same distribution ($p\text{-values}<2\cdot 10^{-16}$ in all cases)).

Orders (light blue cluster appearing in the middle of the tree, Figure 3.8 A). In birds, no particular structure seemed to emerge in within-Family median completeness (although the upper part of the phylogeny, corresponding to Procellariiformes, Charadriiformes, and Anseriformes appeared to be particularly well sampled, Figure 3.8 B). In herptiles, nevertheless, clusters of similar completeness appeared at Family levels. For reptiles, the lower part of the tree appeared to be particularly less well sampled than the above part of the tree (encompassing families such as Tropidophiidae, Lamprophiida or Typhlopidae: mostly, snakes; 3.8 C). In amphibians, groups of families in the Anura Order showed both the best and worst median completeness (Figure 3.8 D).

Overall, these results showed that trait completeness was not random with regard to the phylogenetic relatedness of families. Closely related families seemed to share more similar median trait completeness than less closely related families. As such, the availability of trait information for a species may be dependent on its phylogenetic history; many other factors may interplay with species evolutionary history to explain these patterns.

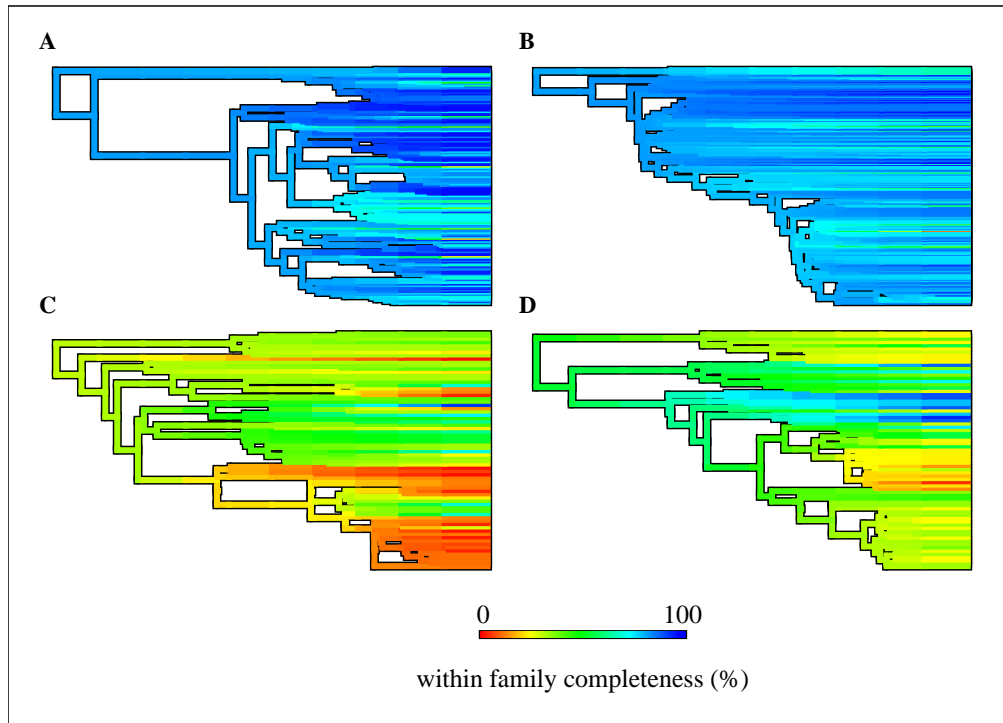


Figure 3.8: Median completeness across families. Tips labels are not shown here for better visualisation of the results; the same figures with tip labels are provided in the SI (zooming into the figure is necessary for mammals and birds); tip label information includes Order and Family. (A) Mammalian Family tree; (B) avian Family tree; (C) reptilian Family tree and (D) amphibian Family tree. Median trait completeness was calculated within families and colour-coded against tree branches. Family clusters of similar median trait completeness appear, particularly in reptiles and amphibians.

Within-Class patterns of trait coverage. Figures 3.9, 3.10, 3.11 and 3.12 show within-Family median trait coverage. In each figure, the subplots are ordered from the trait showing highest overall coverage to the trait showing lowest overall coverage (as in Figure 3.6). The last subplot represents the contribution of each Family to the total number of species in the phylogeny.

As trait coverage decreased, Family clusters of similar median trait coverage became more visible. In mammals, a cluster of families showed low median coverage for trophic level; most of these families were in the Cetartiodactyla Order, which contained marine and aquatic mammals. Families in the Chiroptera Order appeared to be less well sampled for three traits compared to other Orders (body length, litter size, longevity, subplots J to L, Figure 3.9). Families in the Primates Order also appeared to be less well sampled for certain traits (subplots E to I). Among the best sampled mammalian Orders were the Diprotodontia and the Carnivora. For birds, the patterns were less clear (Figure 3.10). Diet information was less resolved for Struthioniformes (subplots H and L: top of the tree); overall, no systematic bias emerged. In reptiles, the lower part of the phylogeny was systematically less well sampled, with a few exception for families such as Boidae and Pythonidae

(subplot E or F: green-blue areas within the red cluster). Overall, trait information for most snakes was systematically less well resolved than for other squamates. In amphibians (Figure 3.11), families in the Caudata Order were overall better sampled (second branch from the root) , as well as a group of families in the Anura Order (third branch from the root, above part). On the other hand, a large number of families in the Anura Order, and most families in the Gymnophonia Order (first branch from the root), were systematically less well sampled. These families nevertheless largely contributed to the total number of species.

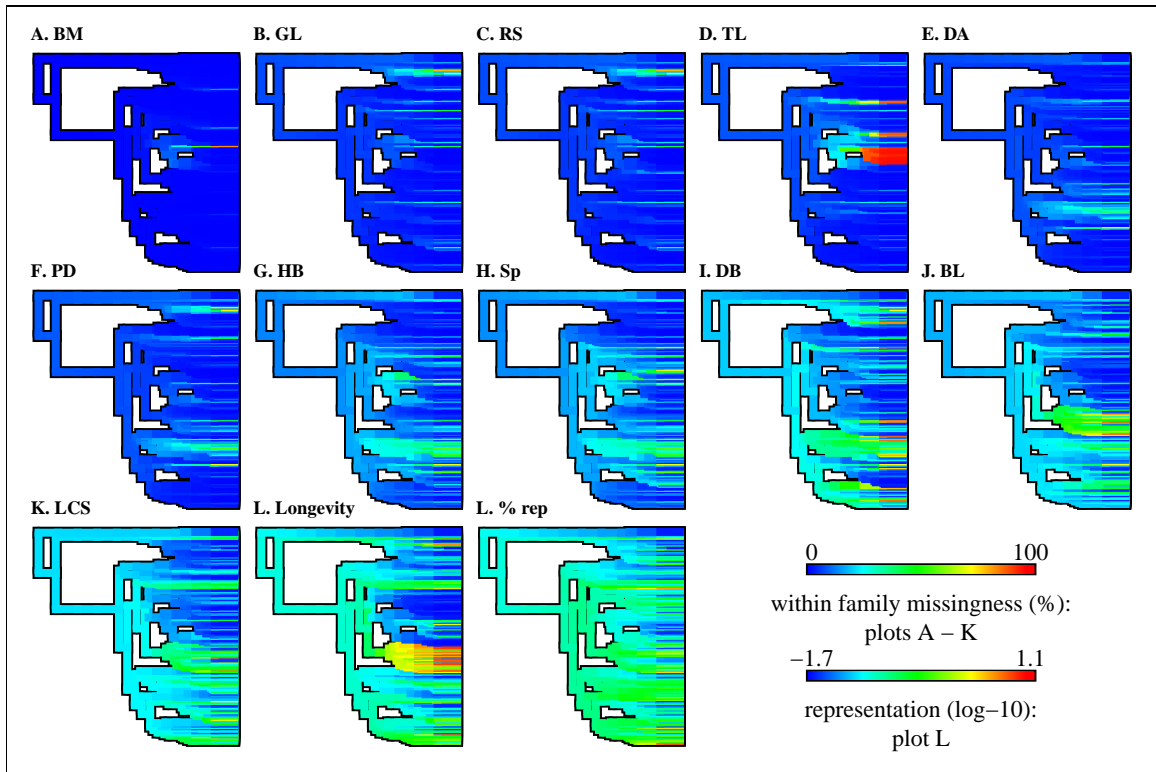


Figure 3.9: Within-Family median trait coverage in mammals.

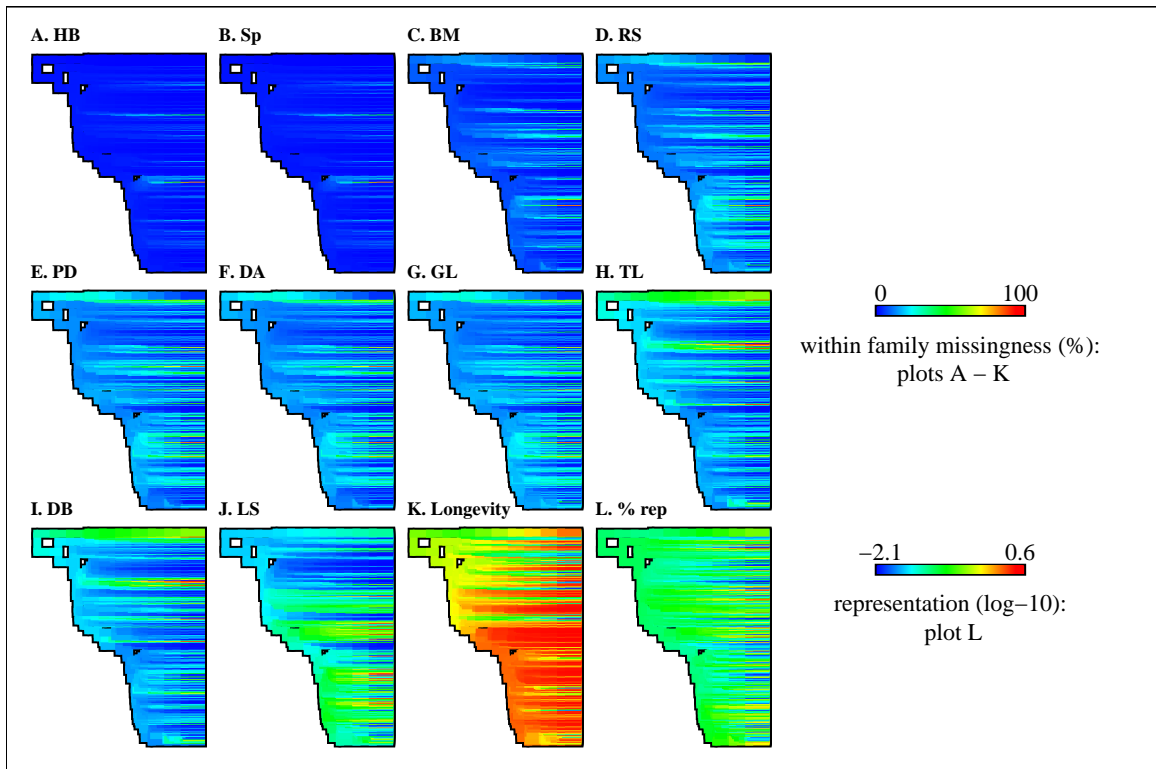


Figure 3.10: Within-Family median trait coverage in birds.

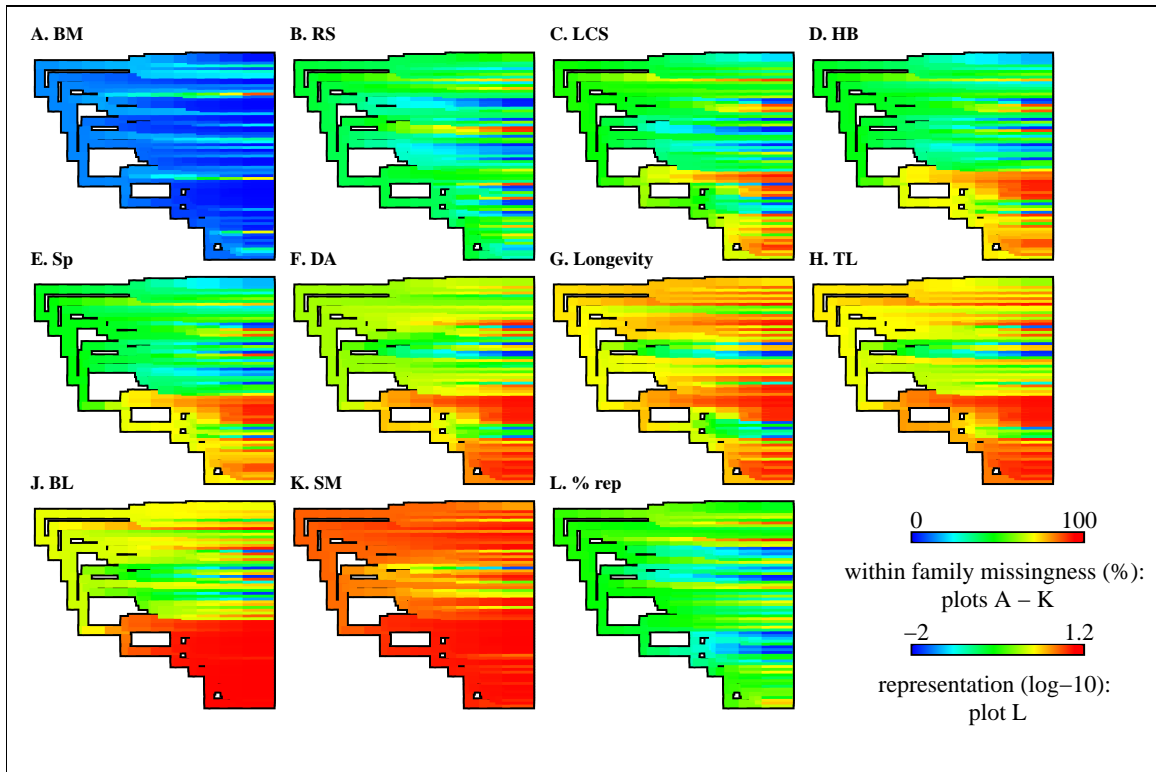


Figure 3.11: Within-Family median trait coverage in reptiles (squamates).

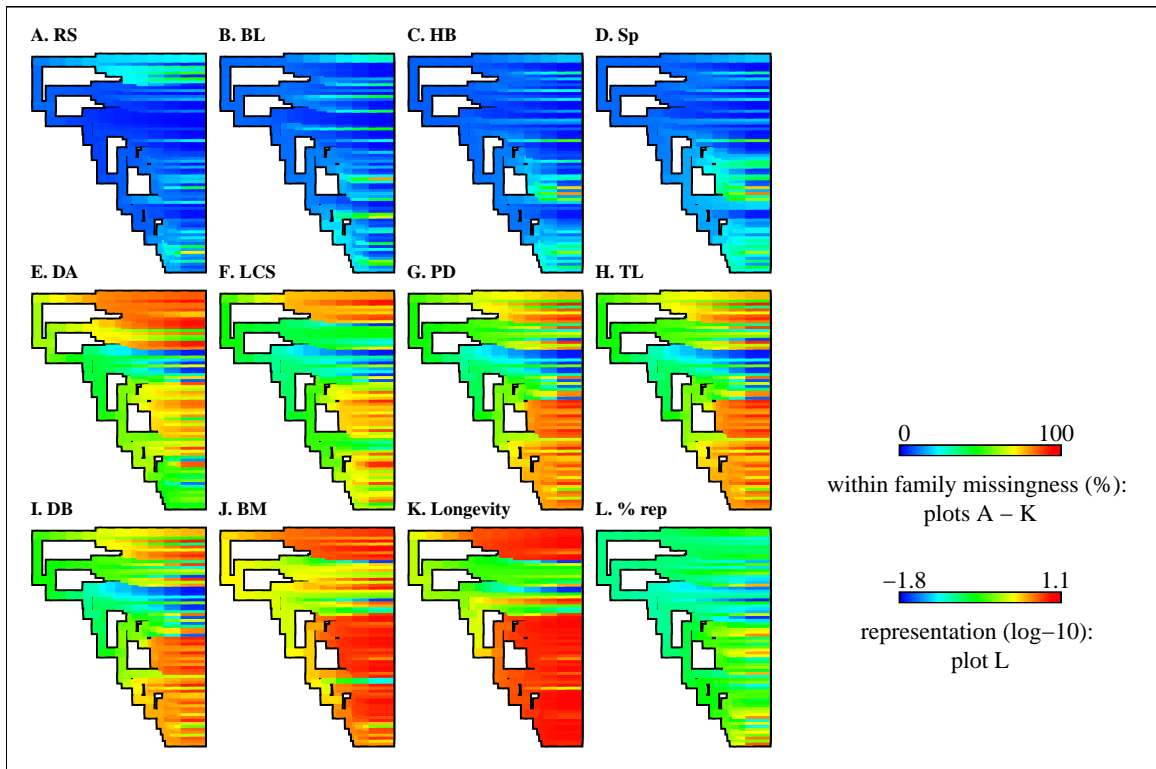


Figure 3.12: Within-Family median trait coverage in amphibians.

3.3.3 Spatial biases of trait completeness

Geographical range size had a significant effect on the number of sampled traits (Table ??, Figure 3.13). Class had a significant effect on the rate of increase in the number of sampled traits, except for reptiles (the rate of increase for reptiles was similar to the rate of increase for amphibians). Baseline rates were higher for mammals and birds than for herptiles, but rates of increase were higher for herptiles. A goodness-of-fit test on the residual deviance did not gather evidence that the model fitted badly ($p=1$, residual deviance: 14440 on 27105 degrees of freedom).

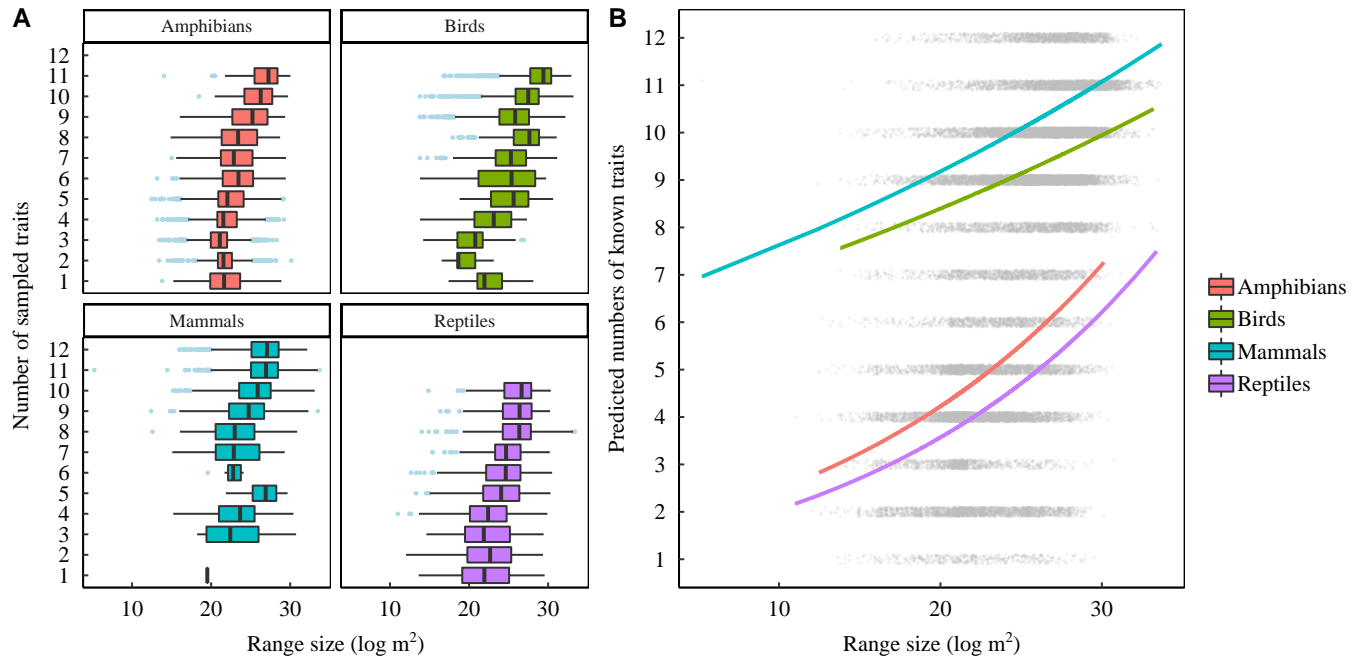


Figure 3.13: Relationship between trait completeness and species geographical range size. (A) Boxplots showing the number of traits sampled in each Class against species geographical range sizes (\log_{10}). **(B)** Regression lines for the fitted generalised linear model. Grey points represent empirical values (not colour coded for better visual clarity). The model was fitted using a Poisson error distribution. Class was added as an explanatory variable with interaction.

Conclusion. Trait data across terrestrial vertebrates is taxonomically biased, with better resolution of trait information in mammals and birds. Phylogenetic biases are strong in reptiles and amphibians, where entire clades appeared to be systematically less well sampled. Finally, across all Classes, species with bigger geographical range sizes are more likely to have more complete trait information. The effect of geographical range size is more pronounced for herptiles than for mammals and birds: the decrease in trait completeness corresponding to a decrease in range size is steeper for amphibians and reptiles.

Table 3.5: Model coefficients. A generalised linear model with a Poisson error distribution was fitted to the number of sampled traits, with range size and Class as interacting explanatory variables. All effects were significant, except for the interaction between reptiles and range size.

Independent variable	Estimate	Std. Error	z-value	Pr(> z)
Intercept (Amphibians)	0.37	0.046	8.05	8.07E-16
log(RS)	0.05	0.002	26.72	2.94E-157
Birds	1.42	0.055	25.66	3.13E-145
Mammals	1.47	0.058	25.16	1.04E-139
Reptiles	-0.21	0.066	-3.15	1.64E-03
log(RS):Birds	-0.04	0.002	-15.89	7.47E-57
log(RS):Mammals	-0.03	0.002	-14.27	3.14E-46
log(RS):Reptiles	0.00	0.003	0.69	4.92E-01

3.3.4 Imputation performance and robustness

Out-of-bag imputation errors.

Figure 3.14 A shows OOB root-mean-squared errors for each continuous traits (shown here for one randomly selected imputed dataset). Figure 3.14 B is the OOB proportion of falsely classified values for categorical traits (for the same imputed dataset). Estimated prediction errors for categorical traits were low to moderate (all below 20%). For continuous traits, estimated errors could be large (e.g., mammalian body mass, amphibian clutch size or range sizes). Nevertheless, such large errors were driven by high trait values in the dataset (see Figure 3.15, which shows the distribution of trait values after imputations; large prediction errors are estimated where traits can attain high values).

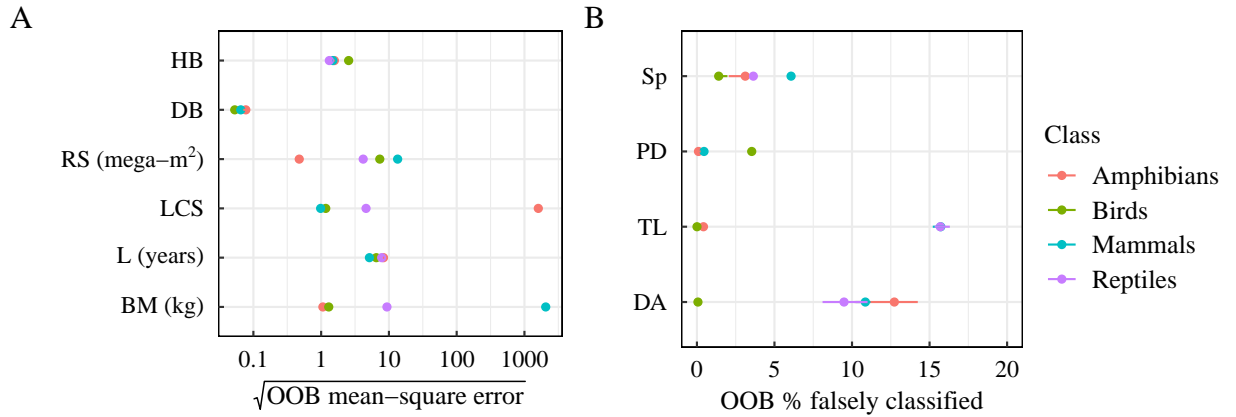


Figure 3.14: missForest out-of-bag root-mean-squared errors and proportion of falsely classified values. (A) Out-of-bag root-mean-square errors for continuous traits. (B) Out-of-bag proportion of falsely classified values.

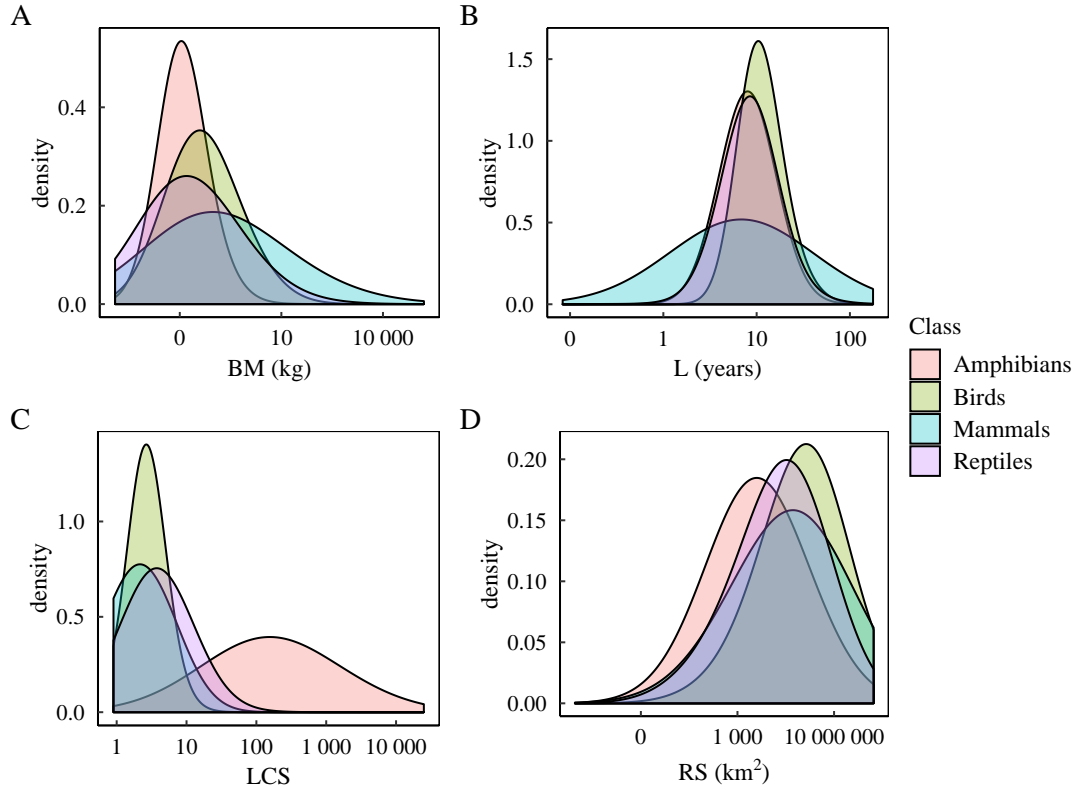


Figure 3.15: Distribution of trait values after imputation for body mass, longevity, litter/clutch size and distribution of range sizes.

Congruence of imputed values among eight imputed datasets.

Figure 3.16 A shows the range and mean of pairwise correlation coefficients obtained for each trait, across eight imputed datasets. Pairwise correlation coefficients were calculated for each trait, predicted in eight independent imputation rounds, so that high correlation values indicated more similar predictions for one trait across the eight datasets. Overall, imputation congruence was high for all continuous traits except habitat breadth. Imputation congruence was high across all Classes for longevity (minimum mean correlation coefficient of 0.87 for reptiles), but more variable in other traits depending on the Class. Figure 3.16 B shows the proportion of species for which imputed values were similar across the eight imputed datasets. At least 50% of all species had similar predicted values across all imputed traits. Imputation congruence was high for trophic level (above 86% in all Classes), and more variable in other traits depending on the Class.

Mammals had the best imputation congruence scores in both continuous and categorical traits (minimum mean correlation coefficient of 0.85 for continuous traits and minimum percentage of agreement of 85% for categorical traits). Imputation congruence for birds was also very good, though

scores were slightly lower for diet related variables (diet breadth and primary diet). For amphibians and reptiles, mean correlation coefficients were all above 0.60, except for habitat breadth. For amphibians in particular, imputation congruence on habitat breadth was poor. Overall, imputed results for amphibians were less congruent than for reptiles, birds and mammals.

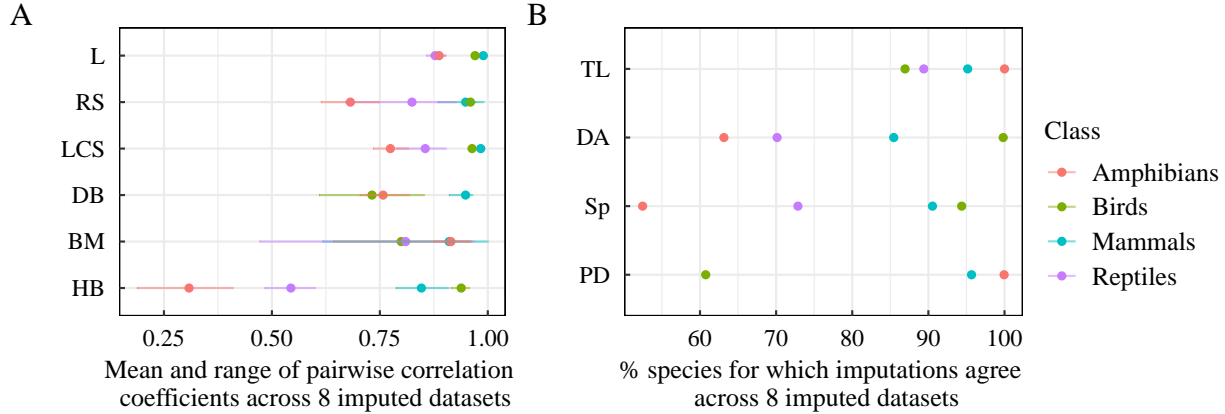


Figure 3.16: Imputation congruence across eight imputed datasets. (A) Mean of pairwise correlation coefficients for continuous traits; (B) proportion of species for which all imputed values were congruent. Imputation congruence was overall good for categorical traits (all above 50%). The lowest scores were obtained for primary diet in birds, as well as for diel activity and specialisation in both reptiles and amphibians (the four points below 85% in (B)). For continuous traits, the means of all pairwise correlation coefficients were above 50%, except for amphibian habitat breadth. The SI provide more detailed congruence results (plots for each Class and each trait).

Conclusion. OOB imputation errors and imputation congruence showed that predictions were overall robust. Habitat breadth was the only variable for which imputation congruence was highly variable across Classes, and below 50% for amphibians. Imputation accuracy may be impacted by the phylogenetic biases in trait completeness. Further work could investigate the impact of non-randomness in the sampling of trait values on imputation accuracy.

3.4 Discussion

In this work, I compiled and imputed data on 10 traits across 5502 mammalian, 10334 reptilian, 11637 avian and 6904 amphibian species. Traits related to species morphological characteristics (body mass), to their life-history (litter/clutch size, longevity, diel activity,), to their habitat preferences (habitat breadth, specialisation), and to their diet (trophic level; for mammals, birds and amphibians only, primary diet and diet breadth were also collated). To my knowledge, there is yet no published or freely available trait database encompassing all terrestrial vertebrates. As such, this work could constitute one of the first attempts to collate extensive trait information across all terres-

trial vertebrates, which was enabled by all past and recent efforts to release trait information in the public domain. Note that the current imputed dataset contains fossil species, as some of the primary sources provided estimates for these. Some marine and aquatic mammals are also represented. Both fossil and non-terrestrial species could be filtered out in the future.

Further developments could include enhancing the existing data to improve initial trait coverage. Alternatively, if novel primary sources were released, new variables could be added to the dataset. Even though the traits included in this work already encompass most of the ecological traits available in the literature across vertebrate Classes, one notable omission was species mobility. Species abilities to both move within their habitats (home range) and to disperse and colonise new areas has a major impact on their aptitudes to cope with anthropogenic changes and on ecosystem functions (Schloss et al., 2012b; Tucker et al., 2018). Nevertheless, traits relating to mobility in amphibians or reptiles were unavailable. The only readily available variable that could have been added was volancy. Other information that could further enhance the dataset include reptilian diet, foraging strata and terrestriality (species habitat preferences along a vertical gradient: e.g. above versus below ground preferences).

The data collection revealed important biases in the availability of trait information across terrestrial vertebrates. Mammals and birds were better sampled than reptiles and amphibians, even for species with similar range sizes. In herptiles, trait information was strongly phylogenetically biased. These results illustrate the biases in global biodiversity knowledge identified by Hortal et al. (2014). Identified gaps are consistent with biases found in González-Suárez et al. (2012) (study conducted on mammals only). Such biases have important consequences on macro-ecological studies. For instance, some analyses make inferences from certain taxa, for which data is available, to other missing-value taxa. Nevertheless, if the sample of studied species is not drawn at random, extrapolations may not be valid. As such, eliminating missing data can, not only, reduce sample sizes, but also bias estimates (Nakagawa and Freckleton, 2008). Imputing missing values therefore appeared to be an interesting option; nevertheless, non-randomness in missing values could bias imputation accuracy.

Penone et al. (2014) conducted a simulation study where missing values were introduced in a trait dataset (10 to 80% missing values). Values were removed in three different ways: completely at random; at random with respect to only one trait; and finally, at random with respect to phylogenies. Their results showed that differences in imputation error in these three cases were marginal and not significant. For some traits, there was a trend for bigger imputation error where missing data were clustered in closely related species. Nevertheless, Penone et al. (2014) showed that missForest

imputations were robust even when trait data was phylogenetically biased.

Using non parametric random forest algorithms to impute missing trait values, as implemented in R by the `missForest` function, presented several advantages over other imputation methods. First, random forests could deal with mixed type variables, and estimate OOB errors for each variable. Second, no underlying data distribution was assumed in the process. Third, `missForest` was computationally faster than other methods, which was an important criterion. Finally, `missForest` has been shown to outperform or perform as well as other approaches (Penone et al., 2014; Stekhoven and Bühlmann, 2012). Moreover, as stated above, Penone et al. (2014) showed that `missForest` imputations were robust even when missing values were not missing at random. Congruence results and imputation errors obtained here showed that `missForest` performed overall rather well. Habitat related variables showed less congruence and may as such be more difficult to impute.

This work highlighted several frequent issues met when working with a large number of species or when working with datasets from different origins. For categorical variables, the levels of the least resolved dataset had to be adopted across all Classes, even though more detailed information was available in another Class. Indeed, common denominators had to be found, at the expense of highly resolved data. One example was diel activity time, that I had to constrain to two categories (nocturnal or non-nocturnal).

I also did not compile any metric reflecting intra-specific variability in continuous traits. Intra-specific variability has been shown to have important effects on ecological systems (Bolnick et al., 2011; Des Roches et al., 2018; González-Suárez and Revilla, 2013). A growing body of literature encourages trait-based research to include intraspecific variability (Carmona et al., 2016; Violle et al., 2012). Here, metrics reflecting intraspecific variability were excluded due to both the scale of the data compilation and the lack of estimates across Classes.

One major issue in this work was the taxonomic ‘pseudoreplication’ of species due to the presence of similar species under diverse names, and other taxonomic errors. Here, taxonomic synonymy artificially increased the amount of missing trait values by creating pseudoreplicates of the same species, inflated the overall number of species and significantly lowered median trait completeness. Taxonomic uncertainty is a recurring problem in ecology and conservation (Isaac et al., 2004). For instance, Cardoso et al. (2017) showed that taxonomic inaccuracies and errors in species checklists lead to the overestimation of plant diversity in the Amazon. The lack of a universal, standardised database for species names complicates species identification. Nevertheless, the production of such a database is difficult to achieve, partly because different conceptual definitions of what a species is

can lead to different taxonomic systems (Isaac et al., 2004). Moreover, frequent updates would be necessary for the database to be in line with the most recent taxonomic revisions. Here, the procedure that I developed to tackle taxonomic redundancy built upon taxonomic information contained in the Red List and the ITIS. Overall, the procedure was not optimal, as these databases did not contain standardised taxonomic information. Nevertheless, it participated in reducing taxonomic mismatches and in increasing trait coverage. Initiatives such as the Taxonomic Name Resolution Service for plants (<http://tnrs.iplantcollaborative.org/>) or the Global Biodiversity Information Facility should encourage researchers to inspect taxonomic uncertainty when working with a large number of species.

Finally, Cooke et al. (2019) released a comprehensive dataset of six mammalian and avian traits. The methods they used to compile and impute trait data were very similar to the methods used in this work. The most notable divergences were the use of different imputation methods (multivariate chained equations) and the pre-selection of traits with more than 50% coverage in Cooke et al. (2019). Because very similar primary sources were used, I did not directly use their data in my work. Nevertheless, I compared the results of both data collection and imputation. The results figure in the SI for comparative purposes.

Conclusion. Future work will build upon the trait data compiled and imputed as presented in this chapter. Even though collation methods may be revisited in the future, the framework is likely to remain similar. More work could be dedicated to assess the impact of phylogenetic biases in trait completeness on imputation accuracy.

I illustrate the first use of this data in the next chapter, which investigates how land-use change impacts the functional diversity of vertebrate communities at global scales. In the last chapter, I detail some research questions that this data will allow to investigate in the future months on my PhD.

4 | Global land-use change promotes the functional homogenisation of local vertebrate communities

Abstract

Land-use change is a major driver of biodiversity loss. Land-use change globally impacts vertebrate ecological communities by reducing local species richness. Nevertheless, not all species respond similarly to land-use change. Species ability to cope with anthropogenic pressures is dependent on their functional traits. By preferentially removing certain functional types, land-use change is reshaping the functional composition of ecological communities. Nevertheless, there is no assessment to date of how global land-use change impacts the functional diversity of local vertebrate communities.

Here, I used a meta-analytic approach to investigate how land-use change impacts the functional diversity of terrestrial vertebrate communities at global scales. Specifically, I used seven vertebrate traits to estimate the functional richness, functional dispersion and functional redundancy of local vertebrate communities across eight land-uses. Using mixed-effect models, I estimated the effects of land-use on each functional diversity index.

Land-use change globally promotes the functional homogenisation of ecological assemblages by favouring species that are both less functionally dispersed and more functionally redundant. The breadth of functions decreases with increasing human land-use disturbance. As such, land-use change induces the functional clustering of local vertebrate assemblages, with potential impacts on local ecosystem processes remaining largely unknown.

4.1 Introduction

Global biodiversity is facing unprecedented challenges (Newbold et al., 2019; Pereira et al., 2012). In the past decades, vertebrate populations have declined by 60% (Grooten and Almond, 2018). Rates of ongoing extinctions have been estimated to be a thousand times higher than those inferred from fossil records (De Vos et al., 2015). This acceleration in biodiversity losses is driven by anthropogenic activities (Steffen et al., 2015). Currently, land-use change is the most important of the diverse anthropogenic pressures driving global biodiversity losses (Foley et al., 2005; Newbold et al., 2015). Future projections show that land-use change, alongside climate change, is likely to remain one of the major threats on biodiversity (Newbold, 2018). To enforce efficient conservation measures, it is vital to consolidate our understanding of how land-use change will impact the diversity of ecological communities. Moreover, it is now widely acknowledged that biodiversity correlates with ecosystem functioning and resilience (Hooper et al., 2012; Oliver et al., 2015; Tilman et al., 2014). As such, species losses are likely to provoke disturbances in ecosystem functioning, which could ultimately negatively impact the services ecosystems provide, many of which are fundamental for human well-being (Duraiappah et al., 2005).

In the recent years, diverse indices have been developed to quantify the functional diversity of ecological communities (Botta-Dukát, 2009; Laliberte and Legendre, 2010; Legras et al., 2018; Ricotta et al., 2016; Schleuter et al., 2010; Villéger et al., 2008). Functional diversity indices reflect the diversity and variability of species functional traits. Indeed, species can be viewed as a collection of traits that determine species fundamental and realised niches, and that underpin their ecological roles (Cadotte et al., 2011; Violle et al., 2007). The quantification of functional diversity can thus inform on processes beyond what other measures of biodiversity (such as species richness) can offer (Cadotte et al., 2011; Laureto et al., 2015). The diverse metrics that have been developed aim at capturing different facets of functional diversity. For instance, the functional richness index developed by Petchey and Gaston (2002) focuses on estimating the total functional distance among species in a community. The functional dispersion index proposed by Laliberte and Legendre (2010) quantifies the variability in the trait composition of a given community.

Traits are hence central to the calculation of functional diversity indices. In particular, traits that underpin ecosystem processes are termed ‘effect traits’ (Lamorel2002a). Modification in the effect trait composition of a community can lead to changes in ecosystem functioning (de Bello et al., 2010; Séguin et al., 2014). By measuring facets of biodiversity that can directly link to ecosystem

processes, functional indices are particularly relevant to understand how disturbances may impact ecosystem functioning. In that case, functional diversity is used as a proxy for ecosystem functioning. Studies investigating how changes in biodiversity relate to changes in ecosystem functioning, using species traits as a mechanistic link, have been mostly focused on plant and invertebrate communities (Hevia et al., 2017). As emphasized by Cadotte et al. (2011), for functional indices to be strictly informative of ecosystem functions, they should be based on traits that have explicit links with ecosystem functioning (or effect traits).

As opposed to effect traits, ‘response traits’ qualify traits that influence how species respond to disturbances (Lavorel 2002a). For instance, empirical evidence has shown that bigger tetrapods are more sensitive to land-use change pressures than smaller tetrapods (Rapacciuolo et al., 2017). As such, body mass has been identified as a response trait to land-use change in terrestrial vertebrates. Diverse studies have documented changes in the trait composition of vertebrate communities affected by land-use change by or habitat degradation (Colin et al., 2018; Flynn et al., 2009; La Sorte et al., 2018; Newbold et al., 2014; Tinoco et al., 2018). Such studies provide with empirical evidence that land-use change leads to non-random modifications of the functional composition of vertebrates. Functional diversity indices can therefore reflect changes in functional composition due to the uneven sensitivity of species, underpinned by their response traits.

Functional diversity indices are therefore useful to investigate how a disturbance affects the trait composition of ecological communities. Whether functional metrics explicitly link to ecosystem functions depends partly on the set of traits included in the calculation. When response and effect traits are coupled, changes in the functional composition of a community mirrors potential changes in ecosystem functioning. On the other hand, when effect and response traits are uncoupled, shifts in the functional composition of a community do not directly reflect changes in ecosystem processes (Luck et al., 2013). In terrestrial vertebrates, studies that investigated the impacts of land-use change on the functional diversity of ecological communities mostly used a set of traits that related to key ecosystem roles; traits notably related to species resource use and energy transfers. For instance, Flynn et al. (2009) showed that land-use intensification (from natural to agricultural) constricted the functional richness of over 25% mammalian and avian communities in the temperate Americas. Nevertheless, to my knowledge, no study has yet investigated how land-use change impacts the functional diversity of terrestrial vertebrates at global scales, across the four classes. Investigating this question is central to understand of global land-use change is reshaping the composition of ecological communities, and how these changes could relate to ecosystem functioning.

In this work, I aimed to assess how land-use change impacts the functional diversity of local terrestrial vertebrate communities at global scales. I used a meta-analytic approach, based upon the PREDICTS database (Hudson et al., 2014, 2017), to test the following hypotheses:

- **Land-use change leads to decreases in the functional richness of local vertebrate communities.** The range of ecological roles decreases in more disturbed land-uses. Land-use change homogenises the local environment and as such acts as an environmental filter. Species presenting certain traits are not able to persist in disturbed habitats and are selectively removed. Previous work has shown that larger, longer-lived species are more sensitive to land-use change (Newbold et al., 2013). Therefore, I hypothesise that the range of functions is likely to be constricted in disturbed habitats compared to natural habitats.
- **The functional dispersion (see Chapter 2) of vertebrate communities decreases with increasing land-use disturbance.** Species in disturbed habitats are more functionally similar than species in more pristine environments, as a consequence of hypothesis (1): species presenting more extreme trait values are being removed from disturbed land-uses, so that remaining species overall have more similar functional roles in disturbed habitats than in more pristine habitats.
- **The functional redundancy of vertebrate communities increases with increasing land-use disturbance.** As species in more disturbed habitats are more functionally similar than species in more pristine environments (hypothesis 2), more species are functionally redundant in disturbed habitats than in pristine environment. With an increase in species richness, new functional roles are introduced at higher rates in pristine environments than in disturbed habitats.

4.2 Methods

4.2.1 The PREDICTS database: using a ‘space-for-time’ substitution to assess the impacts of land-use change on local vertebrate communities

The impacts of land-use change on the functional diversity of vertebrate communities were assessed using a ‘space-for-time’ substitution (De Palma et al., 2018). A longitudinal approach would require data on the evolution of local community composition as the landscape changes from pristine to disturbed, with experimental designs such as ‘before-after-control-impact’ (De Palma et al., 2018).

Nevertheless, such data are difficult to obtain at large scales for diverse taxa and for multiple types of environmental change. With the space-for-time substitution, a spatial gradient is used as a proxy for temporal dynamics. This approach facilitates meta-analytic studies, as data on the community composition of local assemblages in various land-uses is both easier to collect and more abundant.

Such data were compiled in the PREDICTS database (Hudson et al., 2014, 2017). This database is the most comprehensive global collection of studies that sampled biodiversity across diverse land-uses. In all studies, species abundance (in most cases) or occurrence was recorded across a variety of sites, each characterised by a land-use (Table 4.1 and Table 4.3). Land-use was divided into six categories: primary vegetation, secondary vegetation, plantation forest, cropland, pasture and urban (Table 4.1). Secondary vegetation was further divided in three categories (mature, intermediate and young) according to the stage of recovery of the vegetation.

Table 4.1: Land-use categories in the PREDICTS database and their definition. See Hudson et al. (2014), Supplementary Information, for more details. I excluded sites in which land-uses were unknown, as well as secondary vegetation for which the successional stage could not be determined.

Land-use category	Definition
Primary vegetation	Native vegetation. Undisturbed since its development and stabilisation under current climatic conditions: no known alterations due to human activities or to extreme natural events.
Mature secondary vegetation	Recovering vegetation after complete destruction of primary vegetation (either human activities or natural events), where vegetation succession is complete. The architectural structure is close to that of primary vegetation.
Intermediate secondary vegetation	Recovering vegetation after complete destruction of primary vegetation (either human activities or natural events). The architecture is typical of a mid-successional state.
Young secondary vegetation	Recovering vegetation after complete destruction of primary vegetation (either human activities or natural events). The simple architecture is typical of an early successional state.
Plantation forest	Previously cleared areas planted with crop trees or shrubs grown and harvested for human consumption or for commercial purposes (includes wood, fruit, oil, biofuel, rubber,...).
Pasture	Areas grazed by livestock, permanently or regularly. Can be improved through cultivation techniques.
Cropland	Previously cleared areas planted with herbaceous crops and harvested for human or animal consumption (including animal feed and crops used in the food industry), or for commercial purposes (e.g., crops grown for the textile industry).
Urban	Previously cleared areas built by humans (housing, etc.). Vegetation is managed for civic or personal purposes.

I sub-setted the PREDICTS database to retrieve all the sites where vertebrate species were

sampled. Overall, 180 studies were selected (total of 6758 sites). Of these, abundance data were available for 132 studies, and occurrence data for the remaining 48 studies (Figure 4.1).

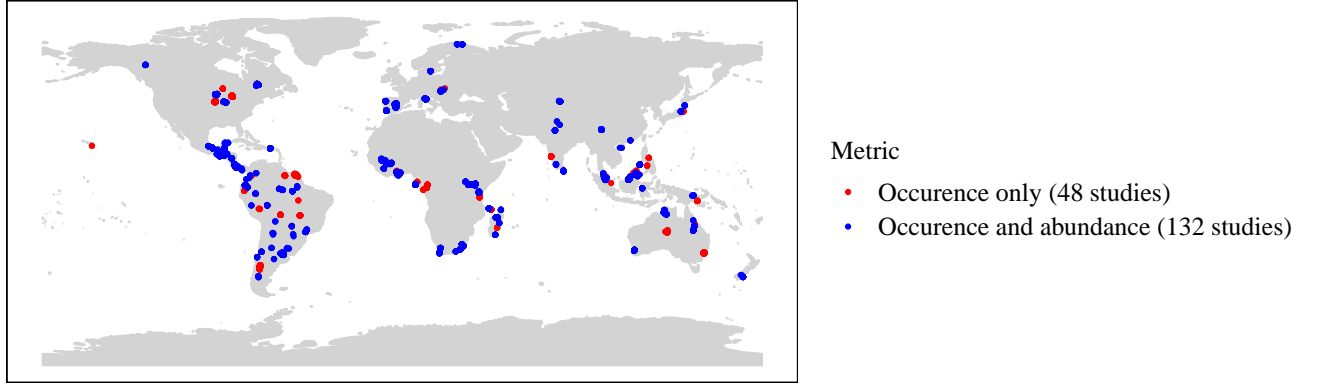


Figure 4.1: Location of selected PREDICTS sites. 180 studies containing occurrence or abundance data for terrestrial vertebrates were selected in the PREDICTS database. Of these, 132 studies provided species relative abundance (corresponding sites are shown in blue on the map). An additional 48 studies only recorded species presence-absence (corresponding sites shown in red).

As the PREDICTS database was built by collating data from independent studies, its design was imbalanced by construction. As such, nearly half of the selected studies only sampled birds, and nearly 80% of the studies considered one class only (Table 4.2). About 17% of the studies sampled species from different classes. Similarly, the number of sites sampled in each land-use category differed across studies (Table 4.3). As such, the PREDICTS database was imbalanced with respect to both taxonomy and land-use.

Table 4.2: Vertebrate classes considered in selected studies. As the PREDICTS database consists of a collation of independent studies, its design was imbalanced. Consequently, most studies focused on only one vertebrate class. Nearly half of the studies only sampled birds, whereas only 4% of the studies sampled only reptiles. About 17% of the studies sampled species from a variety of classes.

Class	Number of studies	% studies
Birds	87	48.3
Mammals	37	20.6
Amphibians	18	10
Amphibians, Reptiles	10	5.6
Birds, Mammals	9	5
Reptiles	8	4.4
Amphibians, Mammals, Reptiles	5	2.8
Amphibians, Birds, Mammals, Reptiles	2	1.1
Birds, Mammals, Reptiles	2	1.1
Mammals, Reptiles	2	1.1

Table 4.3: Sample size for each land-use category. The number of sites in each land-use is reported here across 180 selected studies. The sample size varied importantly with land-use. For instance, there were more than 10 times more sites sampled in primary vegetation than in mature secondary vegetation. Secondary vegetation and urban land-uses had the lowest number of sites sampled.

Land-use	Number of sites	% sites
Primary vegetation	2,569	38.0
Plantation forest	1,151	17.0
Cropland	888	13.1
Pasture	808	12.0
Young secondary vegetation	501	7.4
Intermediate secondary vegetation	350	5.2
Urban	292	4.3
Mature secondary vegetation	199	2.9

4.2.2 Definition of functional diversity indices

In this work, I focused on a few diversity indices aiming at (1) estimating the functional richness of a community; (2) assessing how functionally dissimilar the species in the community are; and (3) measuring the amount of functional redundancy within assemblages. Below, I provide details on the calculation of these indices.

Functional richness: volume-based index VS dendrogram-based index.

For functional richness only, at least four multivariate indices have been developed (Legras et al., 2018). These indices each estimate functional richness differently, and therefore explore a unique facet of the functional richness of a given community. Here, I used two indices, which build upon different conceptual approaches: a dendrogram-based functional richness index (DFR), developed by Petchey and Gaston (2002), and a volume-based richness index (FRic), proposed by Villéger et al. (2008). By construction, both DFR and FRic are likely to be correlated with species richness. Both measures are based on species presence-absence.

The estimation of dendrogram-based functional richness starts with the calculation of a species \times species distance matrix, given a species \times trait dataset (Petchey and Gaston, 2002). This distance matrix contains the pairwise dissimilarities among species, based on their traits. The distance matrix is then clustered to obtain a functional dendrogram, where each tip represents a species. The branch length of the functional dendrogram reflects the functional distance among species. For a given community, the functional richness is calculated as the sum of the branch lengths of the dendrogram

from root to tip, for the tips corresponding to the species in the community. DFR has been notably criticised for being sensitive to the choice of the clustering algorithm (Legras et al., 2018). Therefore, in addition to DFR, I used another index aiming at estimating the functional richness of ecological communities.

The FRic index is a multivariate analogue of the functional range (Legras et al., 2018; Schleuter et al., 2010; Villéger et al., 2008). It is based on the projection of species into a multidimensional space. This space can be a multidimensional trait space (each axis then corresponds to a trait, and trait values are species coordinates). It can also be an n -dimensional space where each axis corresponds to n principal components extracted after principal component analysis on the species \times species distance matrix, obtained from the *species**time**trait* dataset (approach used in this work). The relative position of the species in the space then constitutes the basis for the calculation of diverse indices, among which FRic. FRic is calculated as the volume of the minimum convex hull containing all the species in a given community. FRic hence measures the total volume in multivariate space occupied by the species of an assemblage. Therefore, FRic does not account for ‘holes’ in the convex hull, corresponding to areas within the hull that are not occupied by any species. Consequently, FRic is highly sensitive to the introduction or removal of outlier species, which will affect the definition of the minimum convex hull. FRic values are likely to capture disturbance effects on a given community only if species located at the extremity of the hull are affected.

Functional dispersion and Rao’s quadratic entropy.

Functional dispersion (Laliberte and Legendre, 2010) and Rao’s quadratic entropy (Botta-Dukát, 2009; Rao, 1982) are two highly correlated indices, that both aim to describe the spread of the species in the multidimensional trait space. Both indices are, by construction, independent from species richness (Schleuter et al., 2010). Both can take into account species relative abundance or presence-absence.

Functional dispersion is calculated as the mean distance, in the multidimensional trait space, of each species to the centroid of all species (Figure 4.3 and Equation 4.2). It is expressed as:

$$FDis = \frac{\sum_i p_i \cdot z_i}{\sum_i p_i}, \quad (4.1)$$

where p_i is the relative abundance of the i^{th} species and z_i the distance of the i^{th} species to the centroid of all species. See Laliberte and Legendre (2010) for more details.

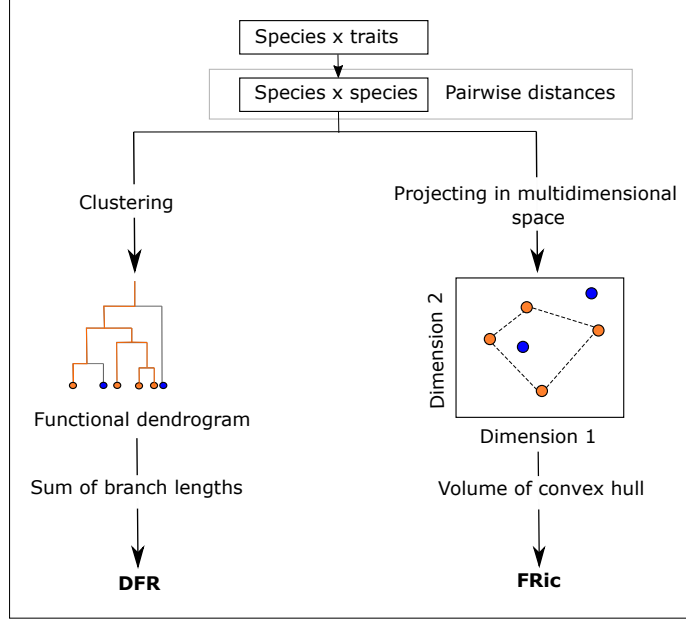


Figure 4.2: Conceptual frameworks for calculating two different functional richness indices: FRic (Villéger et al., 2008) and DFR (Petchey and Gaston, 2002). DFR (dendrogram-based functional richness) is based on the clustering of a pairwise distance matrix summarising functional dissimilarities across species. This results in the obtention of a functional dendrogram, where each tip is a species and branch lengths reflect the functional distance among all species. DFR is calculated as the sum of the branch lengths for species in a given community (for example, orange dots). FRic (volume-based functional richness) is based on the projection of species in a multidimensional space. Species relative position in the space reflects the functional distances. FRic is the volume of the convex hull that includes species in a given community.

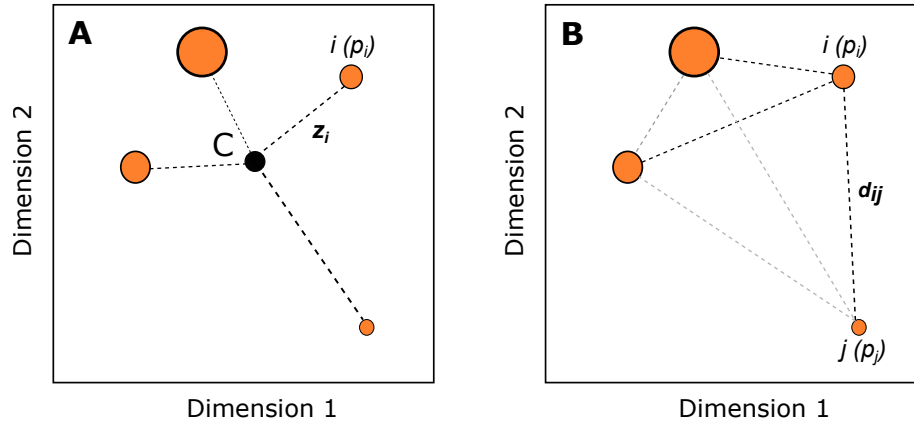


Figure 4.3: Conceptual framework for the calculation of functional dispersion and Rao's quadratic entropy. Each orange circle represents a species, positioned in a bi-dimensional trait space. **(A) Functional dispersion.** The black point C represents the centroid of all species (its position depends on whether the calculations are abundance-weighted or not; see for more details). For the i^{th} species, z_i is the distance to the centroid. The functional dispersion is the mean distance of all species to the centroid (and can be weighted by the relative abundance of each species, p_i). **(B) Rao's quadratic entropy.** $d_{i,j}$ is the pairwise distance between species i and j . Rao's quadratic entropy is the sum of pairwise distances weighted by the product of pairwise relative abundances.

Rao's quadratic entropy describes the average functional distance among species. It is conceptually similar to functional dispersion, but its calculation relies on the functional distance within each

species pair rather than on distances to the centroid. Rao's quadratic entropy is expressed as:

$$Q = \sum_{i,j} p_i \cdot p_j \cdot d_{i,j}, \quad (4.2)$$

where p_i is the relative abundance of the i^{th} species, p_j the relative abundance of the j^{th} species and $d_{i,j}$ is the functional distance between species i and j , obtained from the species×species distance matrix.

Functional redundancy.

Here, I used the framework developed by Ricotta et al. (2016) to calculate the functional redundancy of an ecological community. Ricotta et al. (2016) derived an index to estimate functional redundancy from Rao's quadratic entropy and from the Simpson's diversity index. Functional redundancy is expressed as:

$$R = 1 - \frac{Q}{D}, \quad (4.3)$$

where Q is Rao's quadratic entropy and D is Simpson's diversity index:

$$D = \sum_i p_i(1 - p_i), \quad (4.4)$$

where p_i is the relative abundance of the i^{th} species in the community.

4.2.3 Calculation of functional diversity indices

Trait selection.

In Chapter 3, I collected and imputed the values of nine traits across terrestrial vertebrates (as well as range size). I used these data here to calculate the functional diversity indices. A crucial step was to select the traits to include in the calculations. Indeed, functional diversity indices can be sensitive to the number of traits included (Cadotte et al., 2011; Mouillot et al., 2014). A larger number of traits allows to detect functional differences among species that could be missed with fewer traits (Petchey and Gaston, 2002). Not including enough traits may lead to missing important areas in the multidimensional trait space. On the other hand, the inclusion of correlated traits or neutral traits can inflate functional metrics or cause them to artificially converge with species diversity metrics (Cadotte et al., 2011; Naeem and Wright, 2003). Assessing the degree of multicollinearity among

traits is thus a necessary step before calculating functional diversity indices.

I randomly selected one imputed trait dataset among the eight imputed datasets (see Chapter 3). To improve normality, a log-10 transformation was applied to all continuous traits (except habitat breadth, which was square-rooted). Trait values were also centred and scaled to zero-mean and unit-variance across the four vertebrate classes. All traits were subsequently considered, except those relating to species diet (primary diet and diet breadth), as these were unavailable for reptiles. As such, the traits taken into consideration were: body mass; longevity; litter/clutch size; habitat breadth; habitat specialisation; diel activity; and trophic level.

Assessing the degree of multicollinearity across traits. To first assess whether multicollinearity could be a problem, I estimated Pearson’s pairwise correlation coefficients among continuous traits, as high correlation coefficients can be an indicator of collinearity. A threshold of 0.7 is usually used for detecting potential collinearity (Dormann et al., 2013). The determinant of the correlation matrix can also be assessed, with values close to 0 indicating high degrees of multicollinearity (Dormann et al., 2013).

Table 4.4 shows the pairwise correlation coefficients among continuous traits. Body mass and longevity were the two variables that had the highest correlation coefficient (0.51). The determinant of the correlation matrix was 0.67, thus indicating that the degree of multicollinearity was likely to be low among continuous traits.

Table 4.4: Pearson’s pairwise correlation coefficients among continuous traits. Overall, continuous traits were weakly or moderately correlated. Body mass and longevity had the highest correlation coefficient (0.51), which did not exceed the threshold value of 0.7 used across diverse field as an indicator of problematic collinearity. The determinant of the correlation matrix was 0.67, indicating that multicollinearity was likely not to be problematic among continuous traits.

	Body mass	Longevity	Litter/clutch size	Habitat breadth
Body mass	1			
Longevity	0.509	1		
Litter/clutch size	-0.146	-0.083	1	
Habitat breadth	0.167	0.134	0.194	1

Nevertheless, the previous diagnostics did not take into consideration categorical traits. Potential associations between categorical and continuous traits, or among categorical traits, also needed to be assessed. To that end, I used generalised variance inflation factors (GVIF) or variance inflation factors (VIF), as developed by (Fox and Monette, 1992), to detect multicollinearity across all traits.

Given a regression model, variance inflation factors quantify the overestimation in the variance of estimated regression coefficients due to multicollinearity among the predictors. A VIF or GVIF value of 5 or 10 is commonly used as a threshold to select out collinear predictors (Dormann et al., 2013). I used the function `stepwise.vif` of the R package `Rnalytica` (Tantithamthavorn and Jiarpakdee, 2019), in which a normally-distributed dummy variable was used as a dependent variable in a linear regression model where all traits were used as predictors. The VIF or GVIF of each predictor was then assessed. Multicollinearity across predictors was not detected to be problematically high, as all predictors had a VIF or GVIF value below 2 (Table 4.5). As such, all the traits figuring in Table 4.5 were included in the calculations of functional indices.

Table 4.5: Variance inflation factor of estimated regression coefficients for each trait treated as a predictor in a linear regression model. For categorical traits, the GVIF was calculated rather than the VIF. All traits had a VIF or GVIF below 2: multicollinearity was not problematic among all traits.

Predictor	VIF or GVIF
Diel activity	1.145
Litter/clutch size	1.267
Trophic level	1.288
Specialisation	1.391
Longevity	1.441
Habitat breadth	1.473
Body mass	1.584

Ecological relevance of the traits. See Chapter 1, 2 and Introduction to Chapter 3. All traits used here (4.5) were either response or effect traits, or both.

Calculation of functional diversity metrics across PREDICTS vertebrate communities.

Overview. Functional diversity indices were calculated for each local vertebrate community of the PREDICTS database (in other words, for each PREDICTS site, Figure 4.7A). Functional richness indices (DFR and FRic) were calculated across the 180 studies for which species occurrence was available. Functional dispersion, Rao’s quadratic entropy and functional redundancy were abundance-weighted, and calculated over the 132 studies which provided species relative abundance.

Implementation details. Previous to the calculations of the functional indices, a Gower distance matrix was computed from the species×trait dataset containing all terrestrial vertebrates,

using the `gowdis` function (FD package: Laliberte and Legendre (2010) and Laliberté et al. (2015)). Gower distances allowed to include mixed type variables in the computation. As such, the pairwise dissimilarities among 34,377 terrestrial vertebrates was calculated. This distance matrix was then sub-setted for the species figuring in the PREDICTS database (using the `dist_subset` function of the `usedist` package: Bittinger (2017)).

Dendrogram-based functional richness. For the calculation of DFR, the Gower distance matrix was clustered into a functional dendrogram, using the function `hclust` (base R). I selected the ‘UPGMA’ clustering algorithm (unweighted pair group method with arithmetic mean). This method provided the best correlation coefficient between cophenetic distances and original distances in the Gower distance matrix, of all the methods proposed in the `hclust` function. Then, for each PREDICTS community, the total branch length of the functional dendrogram corresponding to species in the assemblage was calculated using `treedive` (vegan package: Oksanen et al. (2019)).

Volume based functional richness, functional dispersion and Rao’s quadratic entropy. `FRic`, `FDis` and `Q` were calculated for each site using the FD package (Laliberté et al., 2015). The Gower distance matrix was directly passed as an argument in this function. In complement, I calculated the Simpson’s diversity index; I then combined Rao’s quadratic entropy and the Simpson’s index to estimate functional redundancy (Equation 4.3).

4.2.4 Assessing the impacts of land-use change on functional diversity indices

Functional diversity metrics, notably those aiming at estimating functional richness, can be correlated with species richness. For such indices, disentangling the effects of species richness from the effects of land-use is vital. Indeed, an observed decrease in the main effect of land-use on a functional richness index correlated with species richness may be driven by changes in species richness alone. For indices that do not correlate with species richness, the mean effects of land-use on the metrics can be directly estimated.

Functional indices independent from species richness.

As expected by construction, functional dispersion, Rao’s quadratic entropy and functional redundancy were not strongly correlated with species richness (Table 4.7). Surprisingly, volume-based functional richness was also not strongly correlated with species richness (Table 4.6, Figure 4.5A).

There was hence no need to disentangle the effects of land-use from the effects of species richness on these indices. The impact of land-use was assessed using mixed-effect models (lme4 package: Bates et al. (2015)), specified as follows: $\text{Metric} \sim \text{Land-use} + \text{RE}$, where **Land-use** was the recorded pre-dominant land-use in each site, and **RE** all random effects. To account for variation in experimental design across studies, the random effects included the identity of each study and block, as well as the Class of vertebrates considered in each study (Table 4.2).

Functional indices dependent on species richness.

Dendrogram-based functional richness was highly correlated with species richness (Table 4.6, Figure 4.5B). Consequently, it was not possible to directly assess the impacts of land-use change on DFR, as decreases in species richness along the land-use gradient could have confounding effects. To overcome this problem, different studies have investigated how the species richness–functional richness relationship is affected by a disturbance of interest. I used the same approach here.

Table 4.6: Pearson’s correlation coefficients between species richness and indices estimating functional richness. Volume-based functional richness (FRic) was not strongly correlated with species richness. On the other hand, the correlation between species richness and dendrogram-based functional richness (DFR) was higher. DFR and FRic were not strongly associated.

	DFR	FRic	log(SR)
DFR	1		
FRic	0.46	1	
log(SR)	0.71	0.26	1

Table 4.7: Pearson’s correlation coefficients between species richness and indices estimating multi-variate functional spread. As expected by construction, the correlation coefficients between functional dispersion, Rao’s quadratic entropy, functional redundancy and species richness were small. FDis and Q were expectedly strongly correlated. Here, all functional indices were abundance-weighted.

	FDis	Q	Redundancy	log(SR)
FDis	1			
Q	0.94	1		
Redundancy	-0.82	-0.91	1	
log(SR)	0.27	0.17	0.07	1

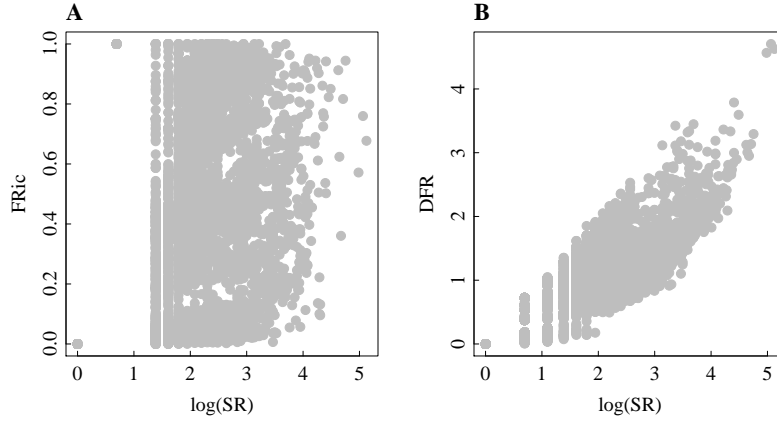


Figure 4.4: Functional richness against species richness. (A) FRic. FRic appeared to be independent from species richness. FRic was standardised so as to be constrained between 0 and 1. **(B) DFR.** DFR and species richness appeared to be positively associated. By definition, the functional richness for communities with only one species was 0.

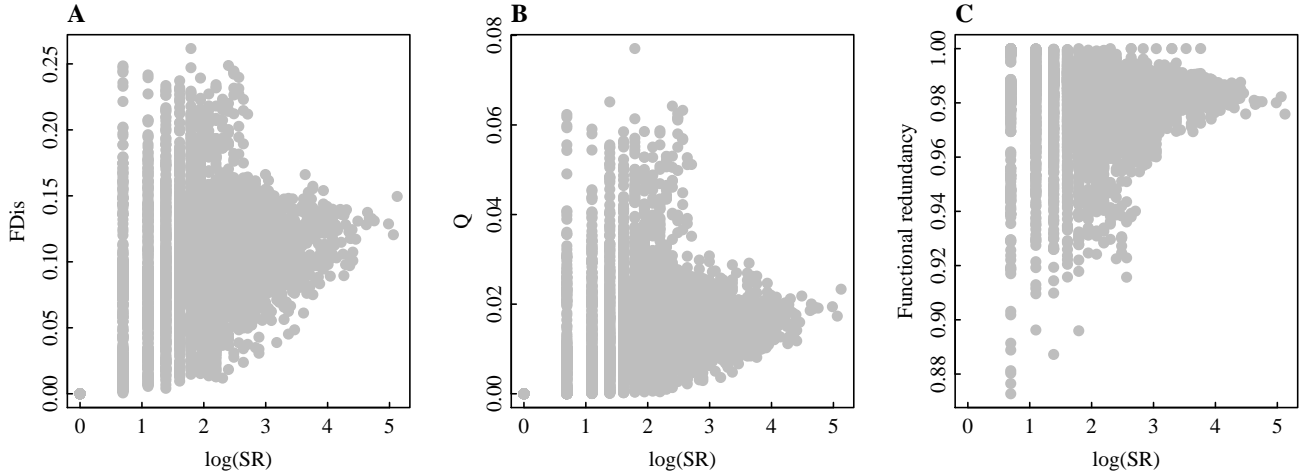


Figure 4.5: Functional dispersion, Rao's quadratic entropy and functional redundancy against species richness. (A) Functional dispersion. (B) Rao's quadratic entropy. (C) Functional redundancy. All three indices were not associated with species richness. Due to their conceptual similarity and high correlation, the behaviour of functional dispersion and Rao's quadratic entropy was similar against species richness. Functional redundancy was closely associated to both dispersion and Rao's quadratic entropy by construction.

How does land-use change impact the species richness–DFR relationship? Species richness was log-transformed to improve normality and linearity. I then investigated how land-use change affected the slope of the species richness–DFR relationship. To that end, I used a mixed-effect model written as follows: $DFR \sim \log(SR) + \text{Land-use} + \log(SR):\text{Land-use} + \text{RE}$. Land-use was added as a main effect with interaction with species richness: the model allowed land-use to affect both the estimated intercept and slope of the relationship between functional and species richness. By defini-

tion, the dendrogram-based functional richness of a community with a richness of one is 0. As such, I did not focus on interpreting the estimated intercepts, as they were not ecologically meaningful. The estimated slopes were the focus of this analysis. According to the hypotheses presented in the introduction, I expected to observe decreases in the slope of the relationship between species richness and DFR along the land-use gradient (Figure 4.6). Indeed, this would signify that with a similar increase in species richness, more disturbed environments gain new functions at lower rates than more pristine environments. In other words, this would mean that species are more functionally redundant in more disturbed land-uses. Moreover, for a given species richness, the theory would then predict that the functional richness in pristine habitats is higher than in perturbed habitats (given a similar intercept).

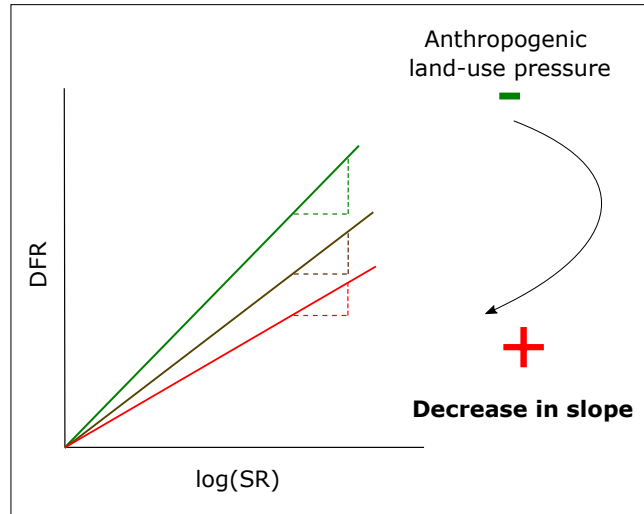


Figure 4.6: Expectations for the effect of land-use change on the DFR–species richness relationship. As by construction, the DFR of a community with only one species is 0, interpreting the effect of land-use on the estimated intercept of the DFR–species richness relationship was not relevant. On the other hand, by affecting the slope of the relationship, land-use change could impact on the functional redundancy of vertebrate communities. Indeed, higher slopes signify that new functions are gained or lost at higher rates with a variation in species richness. I hypothesized that land-use change leads to increases in the functional redundancy of vertebrate communities. As such, I expected decreases in the estimated slope of the DFR–species relationship along an increasingly disturbed land-use gradient.

Disentangling the effects of species richness from the effects of land-use on functional diversity indices through simulations: an adequate approach with PREDICTS?

Approach. Above, I detailed how investigating how land-use change affects the species richness – functional index relationship allows to overcome the ‘species richness problem’ stemming from a high correlation between species richness and the metric of interest. Here, I focus on an alternative

approach, classically used in diverse studies, to disentangle the effects of species richness from the effects of land-use change.

This approach is based on the randomisation of the species composition of local assemblages. It is based upon the formulation of null expectations. Given a species richness, randomising community composition m times allows to generate a distribution of null expectations for a given functional index. These null expectations can then be compared to the empirical (observed) values. Along a species richness gradient that relates to a land-use gradient, such an approach allows to disentangle the impacts of land-use independently from the impacts of species richness on the calculated metrics. Here, such an approach was not necessary *per se*, as most indices were independent from species richness by construction; and for DFR, I focused on the relationship between species richness and the index. Nevertheless, I implemented a simulation approach aiming at examining whether it would be a suitable method given the PREDICTS database.

Specifically, the community composition in each site was randomised by re-sampling species in the corresponding study’s species pool, maintaining the species richness of each site (Figure 4.7). Community composition was randomised 1000 times (for metrics calculated with dbFD) or 10,000 times (for DFR). For each randomised community in each site, the functional diversity indices were calculated. Null expectations of functional diversity indices were then generated for each site by taking the median value obtained across simulations.

Although only DFR was strongly correlated with species richness, I implemented this simulation approach for all functional indices that I considered. The effects of land-use were assessed using the mixed effect models detailed above. My expectations were that:

- For indices independent from species richness by construction, or uncorrelated with species richness, (FDis, Q and FRic), the mean values should be similar among land-uses. Specifically, the mean value expected for primary vegetation, the most pristine land-use type in the dataset, should not differ from the mean value expected in other land-uses (*expectation 1*).
- For the index that correlated with species richness (DFR), the slope of the metric–species richness relationship should be similar across land-uses (*expectation 2*).

NB: For the simulations, FDis and Q were not abundance-weighted, as the simulations were based upon the randomisation of species presence-absence.

Simulation results, and hypotheses as to why simulations may be inadequate. Simulation results showed that land-use was having an effect on null expectations, even for indices

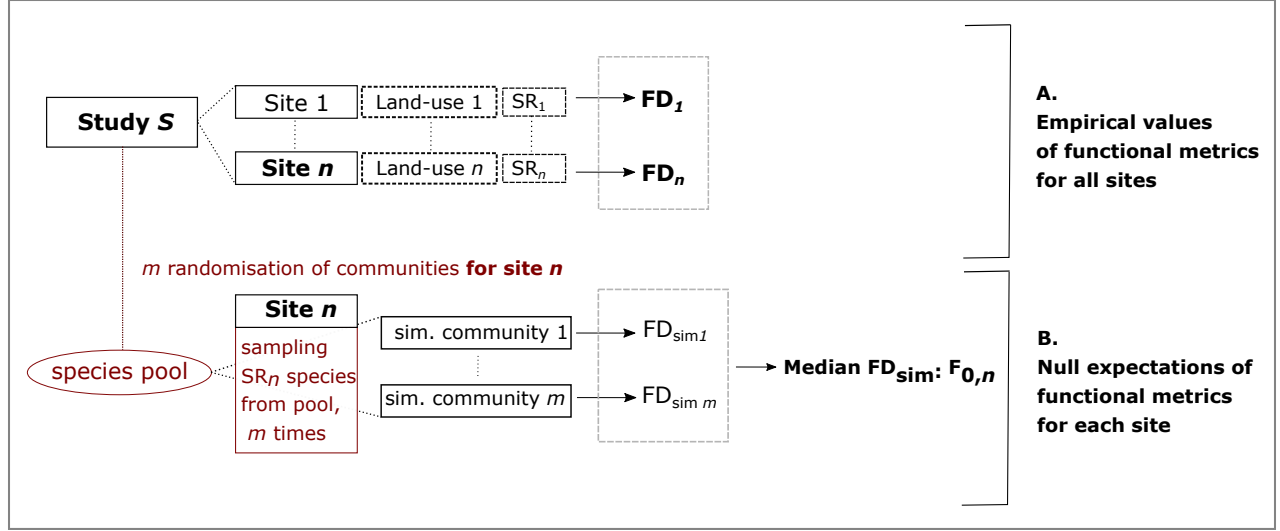


Figure 4.7: Design for calculating functional indices and null expectations for PREDICTS sites. (A) Empirical values for the functional diversity indices were obtained for each PREDICTS site, nested within studies. Each site was characterised by its species richness and its land-use. (B) For a given site n of richness SR_n , the community composition was randomised m times by drawing SR_n species from the study’s species pool. Null expectations for the site n were then obtained by taking the median value of the indices across simulations.

independent from species richness by construction (Figure 4.8). A significant decrease in the mean effect was observed for FRic, FDis and Q in a number of land-uses, contradicting *expectation 1*. Moreover, a significant decrease in the slope of the DFR–species richness relationship was observed for all land-uses except mature secondary vegetation, contradicting *expectation 2*.

Hereafter, I propose a mechanism that may explain why simulated results differed from *expectations 1* and *2*. Simulations were based on the randomisation of the species composition of each site. Species were drawn at random from the species pool, defined as the set of species in each study (equivalent to a ‘regional’ species pool). As such, simulations were sensitive to the composition of the species pool. Nevertheless, the PREDICTS database has an imbalanced design, such that each study do not have sites in all of the land-uses. This may be constricting species pools in some cases. For instance, for a site of land-use ‘Pasture’ belonging to a study where primary vegetation was also sampled, the species pool may be bigger than for a site of land-use ‘Pasture’ where only pasture and plantation forest were sampled. As such, biases in species pool may influence simulation results. Simulation results may capture trends reflecting differences in the size and composition of species pool (Figure 4.9), which may explain the patterns observed in Figure 4.8.

Simulation approach: conclusion. The imbalanced design of the PREDICTS database may be causing biases in species pool, which may render simulation approaches difficult to interpret. As

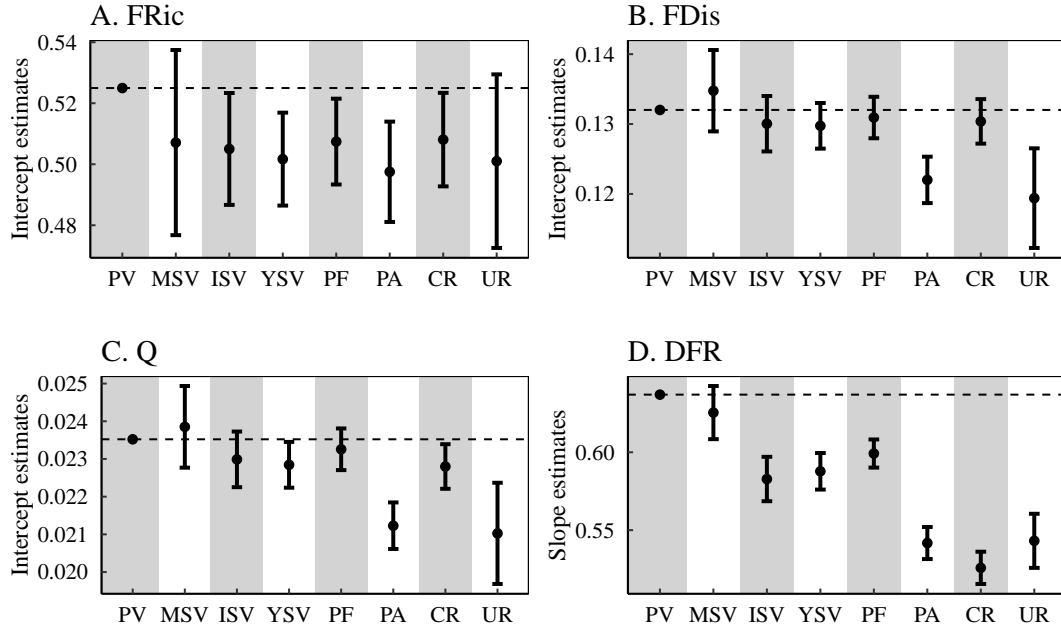


Figure 4.8: Simulation results. Here, null expectations for each index were generated by randomising species presence-absence in each site, maintaining the local species richness. Species were drawn at random from a ‘regional’ pool of species corresponding to all species sampled within studies. In theory, land-use should not have any effect on simulated values. Contrary to this expectation, land-use had a significant effect on the simulated values of all functional metrics for a number of land-uses. As such, I questioned the validity of the simulation approach employed here.

such, the simulation approach was not developed further here. For that reason, simulation results are not detailed in the Results section below.

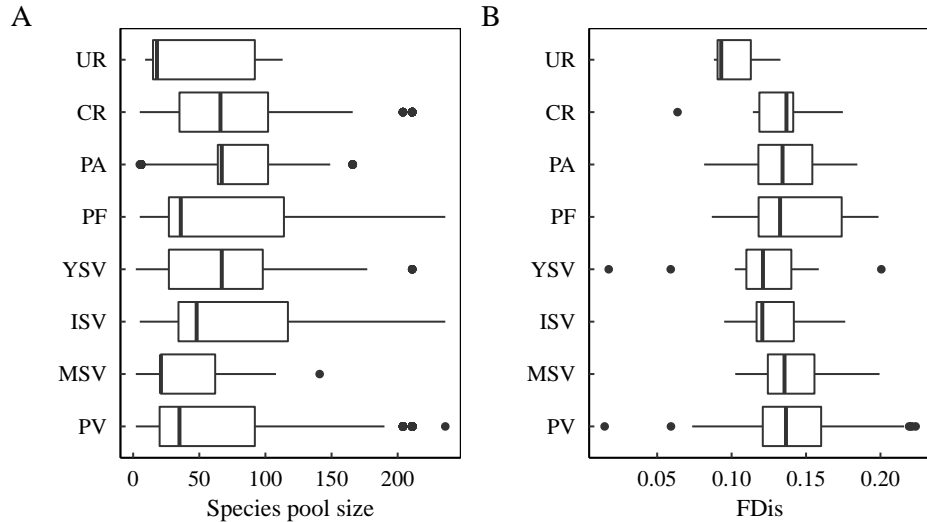


Figure 4.9: (A) Size distribution of species pools in each land-use and (B) distribution of the functional dispersion of the species pools. The trend observed in the simulation results may reflect differences in the composition of study’s species pools. For instance, the median functional dispersion of the species pools corresponding to urban land-uses was smaller than the median functional dispersion of the species pools in other land-uses.

4.3 Results

4.3.1 Land-use change constricted the functional richness of assemblages

Land-use had a significant effect on volume-based functional richness (Figure 4.10). For mature and intermediate secondary vegetation, the mean functional richness was similar to that of primary vegetation. For all other more disturbed land-uses, mean functional richness was significantly different from the mean functional richness of primary vegetation, and decreased with increasing land-use disturbance.

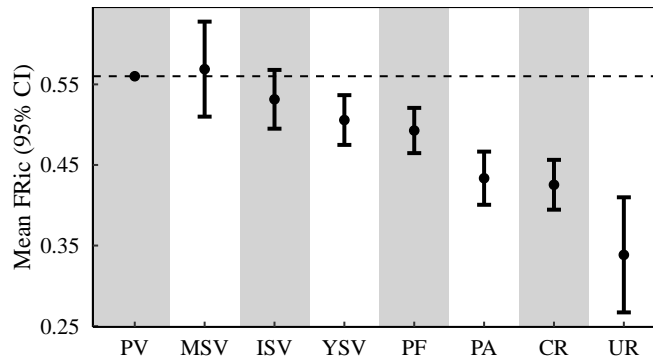


Figure 4.10: Mean effect of land-use on the volume-based functional richness of vertebrate communities (FRic). PV: primary vegetation; MSV: mature secondary vegetation; ISV: intermediate secondary vegetation; YSV: young secondary vegetation; PF: plantation forest; CR: cropland; UR: urban.

4.3.2 Land-use change promoted the functional homogenisation of local vertebrate communities

Land-use change reduced multivariate functional dispersion.

Land-use significantly impacted the functional dispersion of vertebrate communities: for all agricultural and urban land-uses, mean functional dispersion was significantly lower than for primary and secondary vegetation (Figure 4.11). For mature and intermediate secondary vegetation, mean functional dispersion, even though not significantly different, was higher than the mean functional dispersion for primary vegetation.

NB: The mean effect of land-use on Rao’s quadratic entropy was similar to the mean effect of land-use on functional dispersion. This was expected due to the high correlation between these two indices. The plot for Rao’s quadratic entropy figures in the SI.

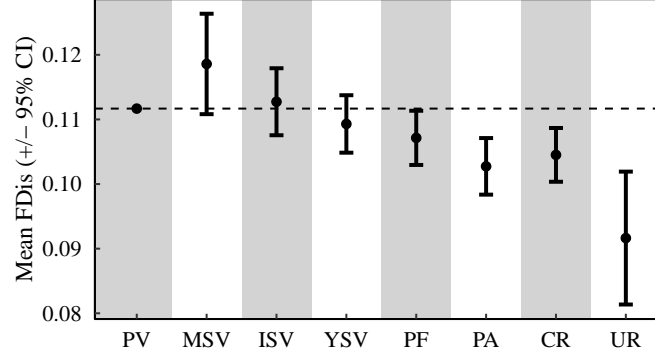


Figure 4.11: Mean effects of land-use on the functional dispersion of local vertebrate communities (FDis). PV: primary vegetation; MSV: mature secondary vegetation; ISV: intermediate secondary vegetation; YSV: young secondary vegetation; PF: plantation forest; CR: cropland; UR: urban.

Land-use change enhances the functional redundancy of local vertebrate communities.

Functional redundancy derived from Rao’s quadratic entropy and the Simpson’s diversity index. Here, functional redundancy was calculated from Rao’s quadratic entropy and the Simpson’s diversity index (Ricotta et al., 2016). The mean functional redundancy was only significantly higher than that of primary vegetation in urban land-uses ((Figure 4.12)). For mature and intermediate secondary vegetation, the mean functional redundancy was lower than the mean functional redundancy of primary vegetation, even though the difference was not significant.

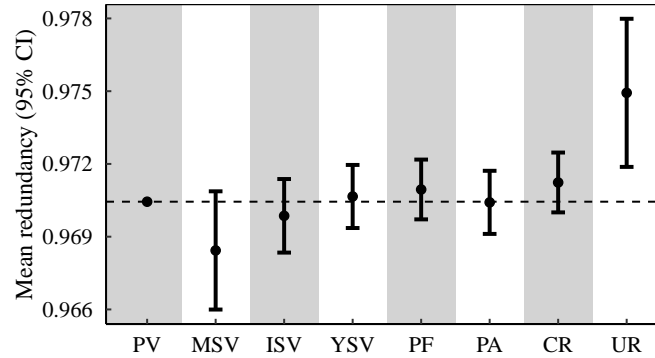


Figure 4.12: Mean effect of land-use on the functional redundancy of vertebrate communities. PV: primary vegetation; MSV: mature secondary vegetation; ISV: intermediate secondary vegetation; YSV: young secondary vegetation; PF: plantation forest; CR: cropland; UR: urban.

Functional redundancy inferred from the slope of the functional-richness – species richness relationship. Here, I investigated how land-use affected the slope of the DFR–species richness relationship. For all secondary vegetation, as well as for agricultural and urban land-uses,

the slope of the relationship was significantly lower than the slope estimated for primary vegetation (Figures 4.13 and 4.14). There was a trend for slopes to decline with increasing land-use disturbance (Figure 4.13).

Global land-use change hence impacted the rate at which the functional distance among species (measured from functional dendrograms) accumulated in local communities with increases in local species richness.

The two approaches used to estimate functional redundancy gave different insights. Notably, the functional redundancy estimated with the metrics developed by (Ricotta et al., 2016) was only detected to significantly increase in urban land-uses. On the other hand, functional redundancy as inferred from the species richness– was estimated to be significantly highest in pastures, using the second approach, and significantly different from that of primary vegetation in all land-uses except mature secondary vegetation.

Figure 4.14 also showed that, given a species richness, the functional richness of disturbed land-uses was predicted to be significantly lower than that of primary vegetation (given the differing slopes).

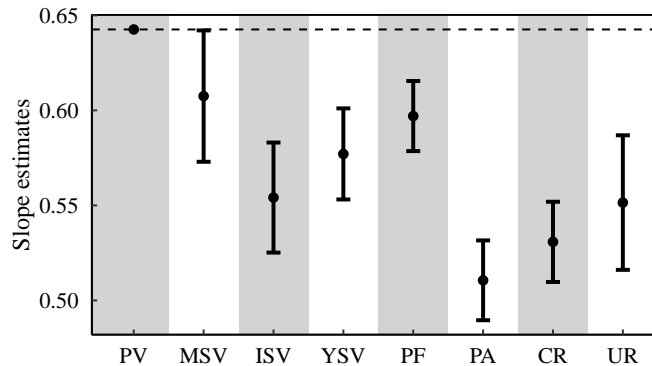


Figure 4.13: Effect of land-use on the slope of the relationship between species richness and dendrogram-based functional richness. **PV:** primary vegetation; **MSV:** mature secondary vegetation; **ISV:** intermediate secondary vegetation; **YSV:** young secondary vegetation; **PF:** plantation forest; **CR:** cropland; **UR:** urban. Land-use significantly affected the slope of the species richness–functional richness relationship.

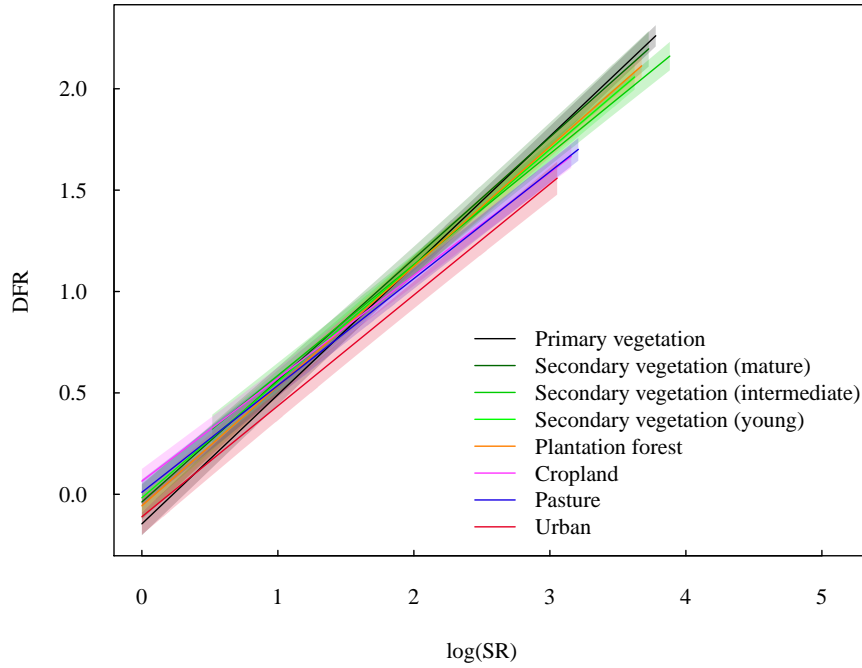


Figure 4.14: Regression lines for the estimated DFR–species richness relationships. **PV:** primary vegetation; **MSV:** mature secondary vegetation; **ISV:** intermediate secondary vegetation; **YSV:** young secondary vegetation; **PF:** plantation forest; **CR:** cropland; **UR:** urban. The regression line was significantly steeper for primary vegetation. Estimated intercepts were not meaningful, as by definition the DFR of communities with only one species is 0. Variations in the slope of the relationship indicated that the rate at which the functional distance among species increased with an increase in species richness was significantly affected by land-use.

4.4 Discussion

In this Chapter, I used a meta-analytic approach to assess how global land-use change impacted the functional diversity of local vertebrate communities. This work constituted, to my knowledge, the first attempt to tackle this question at global scales and across the four vertebrate classes simultaneously.

I showed that globally, land-use change had significant impacts on the functional diversity of vertebrate assemblages. Land-use change acted as an environmental filter that constricted the functional breadth of local assemblages and their mean functional dispersion.

Indeed, because volume-based functional richness is a multivariate analogue of the functional range, global land-use change significantly impacted the functional composition of local vertebrate communities by constricting the multivariate trait range (Figure 4.10). Species located at the periphery of the functional convex hull were more likely to be removed in disturbed land-uses. As such, land-use change significantly impacted the functional composition of local vertebrate communities

by reducing the breadth of functions. Land-use change acted as an environmental filter which excluded species having combination of trait values placing them at the extremity of the communities' convex hulls.

Global land-use change also impacted the trait composition of local vertebrate communities by reducing the mean functional distance of species to the functional centroid of all species (functional dispersion). Species were significantly closer to each other in the multidimensional trait space for agricultural and urban land-uses. These results were in agreement with results obtained for FRic: land-use change selectively removed species that were further away in the trait space, also more likely to be located at the periphery of the convex hulls. Global land-use change, by reducing multivariate functional dispersion, negatively impacted the breadth of functions in local vertebrate communities. These results show that global land-use change promoted the functional clustering of vertebrate assemblages.

Such results could be validated further with an appropriate simulation approach, allowing to generate null expectations of local functional diversity indices. Here, I showed that the simulation approach I attempted may have failed due to potential biases in regional species pool. This work could be developed further to try and design a pertinent and applicable simulation approach.

Overall, global land-use change promoted the functional homogenisation of local communities: species tended to be more functionally similar in perturbed land-uses, and the effect size tended to increase with increasing land-use disturbance. These results agree with studies conducted at local scales that showed, for example, that urbanisation impacted the range of trait values by cutting the tails of trait distributions (La Sorte et al., 2018). Nevertheless, these results contrast with one study that found no effect of urbanisation on the functional diversity of avian assemblages (Oliveira Hagen et al., 2017). There is, however, more and more empirical evidence showing that land-use change favours widespread, generalist species (Newbold et al., 2018), negatively impacts local functional diversity (Chapman et al., 2018; Flynn et al., 2009; Huijbers et al., 2015; Tinoco et al., 2018), and as such, enhances the biotic homogenisation of local communities (Newbold et al., 2019).

Here, I did not investigate the mechanisms behind changes in functional composition. Land-use change could filter out species with functional attributes rendering them unable to cope with the conditions of disturbed habitats; this would correspond to a mechanism known as functional nestedness (Baselga and Leprieur, 2015). On the other hand, novel species, not found in more pristine environments, could also settle in disturbed habitats: the establishment of novel species with functional attributes rendering them able to settle in more disturbed habitats is termed functional

turnover. Novel methods have been developed to assess functional nestedness and turnover (Baselga and Leprieur, 2015). They are notably based on the estimation of functional beta-diversity, which is then partitioned into a turnover and nestedness components. The present work could be developed in the future to investigate the mechanisms driving changes in the functional composition of local vertebrate communities.

By negatively impacting local functional diversity and promoting functional homogenisation, global land-use change could have deleterious effects on local ecosystem functioning (Olden et al., 2004). Functional redundancy is often presented as a safeguard for ecosystem processes. Cooke et al. (2019) found that, across mammals and birds, species-high regions had high functional redundancy and low functional dispersion. The results obtained in this work also underlined the existence of a trade-off between functional redundancy and functional dispersion. Undisturbed land-uses tended to have low redundancy coupled with high dispersion and high functional richness, whereas the opposite was observed for disturbed land-uses. As such, ecosystem processes driven by functionally vulnerable and rare species in pristine land-uses could be at risk if the landscape is modified for human purposes, given the overall lower functional redundancy of undisturbed land-uses.

Finally, in this Chapter, I did not explicitly link the functional diversity metrics to specific functional roles. Nevertheless, the traits that were included in the calculations of the indices were involved in the definition of certain ecosystem functions (for example, trophic level linked to food-web structure and energy transfers). Potential developments of this work could include building explicit links between functional diversity and specific ecosystem functions. The objective of the next Chapter is to develop future questions that I could investigate in the future years of my PhD.

5 | Future questions

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