

Only the largest terrestrial Carnivores increase their dietary breadth with increasing prey richness

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SCHOLARONE[™] Manuscripts **Title:** Only the largest terrestrial Carnivores increase their dietary breadth with increasing prey richness

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1	Tit	le: Only the largest terrestrial Carnivores increase their dietary breadth with increasing prey
2	ricł	nness
3	Ar	ticle type: Review
4	Ab	stract (maximum 300 words)
5	1.	Animals should adapt their foraging habits, changing their diet breadth in response to variation
6		in richness and availability of food resources. Understanding how species modify their diet
7		breadth according to variation in resource richness would help to predict their responses to
8		environmental changes that alter prey communities.
9	2.	We evaluated relationships between diet breadth of large terrestrial carnivores and local
10		richness of large prey. We tested alternative predictions suggested by ecological and
11		evolutionary theories: with growing prey richness, species could (i) show a more diverse diet,
12		thus broadening their diet breadth, or (ii) narrow their diet breadth, indicating specialisation on
13		a smaller number of prey.
14	3.	We collated data from 505 studies of the diets of 12 species of large terrestrial mammalian
15		carnivores to model relationships between two indices of dietary breadth and local prey
16		richness.
17	4.	For the majority of species, we found no evidence for narrowing diet breadth (i.e., increased
18		specialisation) with growing prey richness. Although snow leopard and dhole appeared to use a
19		lower number of large prey species with growing prey richness, larger sample sizes are needed
20		to support this result.
21	5.	With growing prey richness, the 5 largest carnivores (tiger, lion, spotted hyaena, jaguar, puma),
22		plus grey wolf and Eurasian lynx (which are usually top predators in the areas from which data
23		were obtained) showed broader diet breadth and/or used a greater number of large prey species
24		(i.e., increased generalism).
25	6.	Dominant carnivores should encounter little competition in expanding their diet breadth with
26		growing prey richness; conversely, the dietary niche of subordinate large carnivores would be

- limited by competition with larger, dominant predators. Over evolutionary time, resource 27
- partitioning would be more important in shaping the dietary niche of smaller, inferior 28
- competitors than those of dominant ones. 29
- 30
- 31
- Key-words: Canidae; food habits; large carnivores; Felidae; interspecific competition; predator-32
- prey relationships. 33
- Running head: Diet breadth of large terrestrial carnivores 34
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36 Introduction

Variation in richness and availability of food resources is expected to affect foraging habits and diet 37 composition of species (e.g., Jarman 1974; Macdonald 1983; Hofmann 1989), in turn influencing 38 their diet breadth (e.g., Birkhofer & Wolters 2012; Forister et al. 2015; Creel et al. 2018). 39 Environmental changes and anthropogenic alterations of natural communities have profound 40 ecological consequences (e.g., Walther et al. 2002, 2009), often influencing the richness and 41 availability of resources available to foragers. The extent to which species modify their diet 42 according to variation in food availability is influenced by anatomical, physiological and 43 behavioural traits, as well as ecological determinants (e.g., Hofmann 1989; Forister et al. 2015; 44 45 Creel et al. 2018). Understanding whether and how species modify their foraging habits according to variation in food resources should help to predict their responses and ability to adapt to the 46 effects of environmental changes (Forister et al. 2015). 47

Large terrestrial carnivores play fundamental roles in ecosystems and are ecologically, 48 culturally and economically important worldwide (Ripple et al. 2014). Changes in the abundance of 49 large terrestrial carnivores can trigger cascading effects with impacts on lower trophic levels (e.g., 50 Hebblewhite et al. 2005; Estes et al. 2011; Ripple et al. 2014). Additionally, many are threatened 51 with extinction (Macdonald et al. 2010). Usually, they depend on medium-large terrestrial 52 53 vertebrates for prey (Carbone et al. 1999, 2007), and this dependence tends to increase with carnivore body size (Carbone et al. 2010). Their food habits – and, thus, their diet breadth – are 54 expected to be strongly influenced by variation in the richness of large prey (see also Estes et al. 55 2011; Hatton et al. 2015; Creel et al. 2018). 56

Ecological and evolutionary theories, as well as empirical studies, invite opposing expectations about the dietary response of foragers to changing richness of their food resources. When a greater diversity of resources is available, foragers may respond by expanding their dietary breadth (Birkhofer & Wolters 2012). However, we might expect to observe a more diverse community of herbivores, which are prey to carnivores, in nutrient-rich areas (e.g. Danell et al.

62 1996; Prins & Olff 2002). In richer environments, optimal foraging theory predicts that species 63 should become increasingly specialised (MacArthur & Pianka 1966; Charnov 1976), narrowing 64 their niches to concentrate on the most profitable resources. Thus, when the richness of food 65 resources is greater, we would expect a narrower diet breadth (MacArthur & Pianka 1966; Schoener 66 1971; Forister et al. 2015).

Variations in prev communities can be triggered by natural or anthropogenic drivers (e.g., 67 Sinclair et al. 2003; Wegge et al. 2009; Creel et al. 2018). Poaching, intensive hunting and/or 68 habitat manipulation can reduce prey abundance and diversity, posing significant threats to 69 carnivore populations, as well as affecting their food habits (e.g. Estes et al. 2011; Sandom et al. 70 71 2017; Creel et al. 2018). Additionally, environmental/climatic changes can influence species' distributions (e.g. Walther et al. 2002, 2009), which is expected to trigger a re-arrangement of 72 communities, in the medium-to-long term. Natural or human-driven variations in prey communities 73 74 influence food ecology of carnivores (e.g., Meriggi & Lovari 1996; Bagchi & Mishra 2006; Khorozyan et al. 2015; Creel et al. 2018; Khan et al. 2018). Thus, assessing whether and how the 75 diet breadth of large terrestrial carnivores varies with changing prey diversity has important 76 theoretical and practical implications. Despite this, a global evaluation of how the diet breadth of 77 78 large carnivores varies with prey richness is lacking.

79 We reviewed information on the food habits of large terrestrial carnivores to evaluate intraspecific variation in diet breadth in relation to prev richness. In particular, we assessed support for 80 the following alternative predictions: (i) with increasing prey richness, carnivores use a larger 81 82 spectrum of prey, thus showing a broader diet breadth; or (*ii*) with increasing prey richness, carnivores tend to specialise on the most profitable prey, thus showing a narrower diet breadth. 83 Body size of carnivores influences their dependence on large prey (Carbone et al. 2010). From that, 84 we would expect that (iii) the strength of the relationships between dietary breadth and richness of 85 large prey increases with carnivore body size. 86

88 Methods

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90 *Study species and definitions*

We focused on large terrestrial carnivores (Mammalia: Carnivora), with a diet mainly based on 91 large terrestrial vertebrate prey. In particular, we included carnivores heavier than 14.5 kg (mean 92 body mass), the body weight threshold suggested for carnivores to switch to an obligate carnivorous 93 94 diet based on large prey (Carbone et al. 2007), excluding Ursidae, which typically have diets dominated by non-vertebrate foods. Considering mean adult body masses taken from the PanTheria 95 database (Jones et al. 2009), 15 carnivore species were selected initially: tiger Panthera tigris (for 96 97 body mass, see Table 2), lion Panthera leo, jaguar Panthera onca, spotted hyaena Crocuta crocuta, puma Puma concolor, leopard Panthera pardus, cheetah Acinonyx jubatus, brown hyaena Hyaena 98 bruna, striped hyaena Hyaena hyaena, snow leopard Panthera uncia, grey wolf Canis lupus, 99 100 African wild dog Lycaon pictus, Eurasian lynx Lynx lynx, dhole Cuon alpinus and clouded leopard *Neofelis nebulosa*. Among them, the brown hyaena (43.0 kg) and the striped hyaena (35.1 kg) were 101 not considered because they are mainly scavengers (e.g., Kruuk 1976; Owens & Owens 1978; Stein 102 et al. 2013; Yarnell et al. 2013), with food habits dependent on carrion left by other carnivores. 103 Additionally, the clouded leopard (14.9 kg) was excluded because of the very limited number of 104 105 studies on its food habits (Rabinowitz et al. 1987; Grassman et al. 2005). Thus, we based our review on 12 species. 106

Large terrestrial carnivores rely mainly on large terrestrial prey (Carbone et al. 2007). So, we considered the richness of large terrestrial vertebrate prey (hereafter "large prey") that we defined as heavier than 10 kg, following Carbone et al. (2007). Livestock types are often inconsistently reported across studies, being frequently pooled in a single 'livestock' category. To ensure a consistent treatment across studies, all livestock types were pooled into a single category of large prey. In some cases, several species of prey were reported in the literature as a cumulative prey category (e.g. 'deer', 'peccaries', or 'duikers'): in these cases, we considered the cumulative prey

114 category as a single 'species'. Wild prey smaller than 10 kg were pooled together into a single115 'Other' category.

116

117 *Literature review*

We searched ISI Web of Science and GoogleScholar with combinations of keywords that included 118 the common or scientific name of the focal carnivore species, and either 'diet', 'food habit', 'prev 119 120 use', 'prey selection' or 'predation'. We considered only documents published in English, up to January 2019. In particular, we considered peer reviewed scientific papers, books, PhD/MSc theses 121 and conference proceedings. We also considered literature cited in papers found through this search, 122 123 to locate potential papers not detected in the first step. Studies were included only when both the carnivore and prey communities were described. For each study, we defined the spectrum of 124 available potential large prey by considering information reported in primary sources. In some cases 125 - especially in prey-rich areas, such as African national parks - the description of the prey 126 community could be incomplete in primary articles. Therefore, we evaluated other official sources 127 (e.g., other contemporary papers or official check-lists) to assess the number of potential prey 128 species. Very large/mega-herbivores may not be prey for some carnivores (e.g., elephants 129 Loxodonta africana, for African wild dogs). Thus, for each carnivore we only considered those 130 131 species that have been reported - or known from literature – to be preved on by that carnivore as potential large prey (see Appendix S1 for a list of excluded prey species for each carnivore). We 132 only considered studies based on analyses of scat contents or kills. If a study reported both scats and 133 kills, we only used results obtained through the method based on the greatest sample size. 134

Several indicators can be used to quantify food habits of carnivores (e.g., absolute
occurrence, relative occurrence, volumetric estimates, biomass estimates; Kruuk & Parish 1981;
Ciucci et al. 1996; Chakrabarti et al. 2016). Among them, absolute/relative occurrences are the most
widely used methods and can be employed to evaluate dietary breadth (see below). Additionally,
besides uncertainties affecting the calculation of biomass consumed from scats (cf. Chakrabarti et

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al. 2016; Khorozyan et al. 2017; Lumetsberger et al. 2017), it is usually impossible to know (i) the 140 age class and sex that was preved upon; (ii) whether other carnivores also fed on the carcass; or (iii) 141 whether an individual fed alone or with conspecifics (e.g., a pair, a pack or a female with pups/cubs; 142 Bocci et al. 2017; Khan et al. 2018). Thus, to ensure consistent treatment across studies and species, 143 we considered studies: (i) which included absolute occurrences of given prey species in the diet, 144 i.e., the ratio of the total number of occurrences of that prey over the total number of scats/kills 145 analysed; (ii) for which absolute occurrence could be derived from relative occurrences (the ratio of 146 the total number of occurrences of that prey over the total number of prey items) reported in the 147 study; or (iii) reporting relative occurrences. 148

We defined as "study" an account of food habits of a carnivore species in a given study area. We selected an initial total of 505 studies (out of 351 sources) and compiled a dataset including the following information: paper title, year of publication, author(s), journal (if peer-reviewed), study area, study period (years/months), carnivore focal species, sample type (scats or kills), sample size (i.e., number of analysed scats or kills), absolute occurrence (or relative, if the absolute figure was not provided) of each prey reported in the primary source.

155

156 *Data analyses*

157 We (i) pooled together all livestock types and summed up their occurrences; and (ii) pooled together all smaller prey into the 'Other' category, and summed up their occurrences. For each 158 predator, multiple studies conducted in the same study area were pooled (i.e., absolute occurrences 159 of each prey, as well as sample sizes, were summed up across studies), to avoid pseudoreplication 160 of data obtained in the same study areas. Although changes in prey communities may occur 161 between different periods, the number of available prey species – which is the predictor used in the 162 analyses – is most unlikely to vary across several studies conducted in the same study area; 163 consequently, pooling should not affect our conclusions. Only studies conducted with the same 164 165 method (scats or kills) were pooled: in cases where the two methods had been used at different

times in the same area, only results obtained through the method based on the higher number of observations were considered. When necessary, we added additional prey species available in the study area, but not reported in the paper(s), and assigned them a '0' occurrence (n = 81 studies, after pooling). The list of studies selected for the analyses (N = 415) is reported in Appendix S2. A summary of the number of studies considered per species and sample type, before/after study filtering/pooling, is shown in Table 1.

We calculated relative occurrences (i.e., the relative use) of each prey by dividing their 172 absolute occurrence by the total number of prey items, which is necessary to calculate diet breadth 173 indices (see below). For each carnivore species, we considered two indices of diet breadth. First, we 174 175 considered the total diet breadth, inclusive of all prey. To estimate total diet breadth, we considered the Levins index (Levins 1968) as: $B = 1/\sum p_i^2$, where p_i is the proportion of items in the diet that 176 belong to the food category *j*. For this index, the 'Other' category was included in the calculation of 177 the overall dietary breadth. Large carnivores mainly depend on large prey for feeding and survival 178 (Carbone et al. 1999, 2007; see Introduction). Thus, we evaluated whether variation in the richness 179 of the guild of large prey would influence the use of large prey, prompting consideration of a 180 second, additional index focused on large prey only. Specifically, we determined the number of 181 frequently used resources (Krebs 1999) restricted to large prey showing a relative occurrence in 182 183 scats of at least 5% (this second index is the "Large Prey Index") (Krebs 1999). The Levins index and the Large Prey index can vary from 0 to the maximum number of available prey (with a 184 maximum of 20 for Large Prey index, which is theoretically possible if 20 prey are used, each with 185 a frequency of 5%). For both indices, increasing values indicate increasing dietary breadth. The 186 Standardised Levins index is frequently used to quantify diet breadth (Colwell & Futuyma 1971; 187 Hurlbert 1978; Krebs 1999). The calculation of this index includes both the proportional use of 188 resources and the number of available potential resources (Colwell & Futuyma 1971; Hurlbert 189 1978; Krebs 1999). For this index, values are expected to be influenced by the number of potential 190 191 resources available (i.e., resource states) and, thus, it is unsuitable for comparisons between

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192	populations using different resource matrices (Colwell & Futuyma 1971). Since our major focus
193	was the assessment of variations of dietary niche in relation to variations in prey richness, we did
194	not use the Standardised Levins index.
195	We used linear models to evaluate relationships between dietary breadth and richness of
196	large prey. For both indices, the following model was used for each species:
197	
198	log(Diet breadth index) ~ log(N available large prey) + Sample type
199	
200	To account for the greater reliability of studies based on larger samples, we weighted by sample
201	size [using "weights = sqrt(sample size)" in the model declaration]. To account for studies where no
202	large prey showed a frequency of use greater than 5%, we used a $log(x+1)$ transformation for the
203	Large Prey Index (where x represents this index for any given species in a given study).
204	All models were weighted by square root of sample size (i.e., N scats or kills analysed in each
205	study) because variance typically declines linearly with that measure.
206	To evaluate whether the relationship between diet breadth and prey richness was affected by
207	carnivore body mass, we considered the model coefficients for the relationships between prey
208	richness and each dietary breadth index. Specifically, for both indices, we evaluated the following
209	model:
210	
211	Model coefficient ~ log(body mass)
212	
213	Relationships were considered as statistically supported if the 0.95 confidence intervals of
214	coefficients did not include '0'. Analyses were conducted through the RStudio version 1.1.447
215	software (R Core Team 2018).
216	

217 Results

The Levins index (i.e., the total diet breadth) increased with growing prev richness for 6 carnivores 218 (lion, jaguar, spotted hyaena, puma, wolf and Eurasian lynx). Although the relationship between 219 this index and prey richness was negative for only two species, no association was supported for the 220 other 6 carnivores (Table 2; Fig. 1). 221 The Large Prev index (i.e., the diet breadth limited to the use of large prev) increased with prev 222 richness for 7 carnivores, including the five largest species considered (tiger, lion, jaguar, spotted 223 hyaena and puma), as well as the wolf and Eurasian lynx (Table 2; Fig. 2). This index showed a 224 decrease with growing prey richness for the dhole and the snow leopard, although 0.95 confidence 225 226 intervals were close to zero (Table 1).

227 We observed that the prey richness-dietary breadth relationship tended to increase with body mass,

but 0.95 confidence intervals always included zero (Table 3; Fig. 3). The fits, model coefficients

and 0.95 confidence intervals for linear models of both indices and predictors are shown in

230 Appendix S3.

231

232 Discussion

We assembled a large data set to relate the diet breadth of all large terrestrial mammalian carnivores 233 234 to the richness of their prey communities. In spite of the heterogeneity of sources, ecosystems and researchers, our results support the existence of relationships between body size, dominance in 235 guilds and niche-partitioning. In particular, our results strongly suggest that the five largest 236 237 carnivores in the world, as well as the wolf and the Eurasian lynx, use a greater number of large prey species with growing prey richness (our prediction *i*). Furthermore, our findings identify 238 directions for future research into the potential for large carnivores to adapt their foraging behaviour 239 in the face of changing prey communities. We discuss our findings in relation to those two broad 240 issues. 241

243 Diet breadth, niche partitioning and body mass

Optimal foraging theory suggests that, with growing environmental productivity, species will 244 specialise on the most profitable food resources, narrowing their diet breadth (MacArthur & Pianka 245 1966; Charnov 1976). As opposed to totally indiscriminate predation, prev selection has been 246 reported for large carnivores and it can be influenced by several factors, such as prev abundance, 247 accessibility and/or vulnerability, as well as body size and anti-predator behaviour (e.g., Karanth & 248 249 Sunquist 1995; Hayward & Kerley 2005; Jędrzejewski et al. 2012; Lovari et al. 2015). We found that most large carnivores do not reduce niche breadth when the richness of large prey increases. 250 The Levins index and the Large Prey index provide different information. The Levins index refers 251 252 to the whole dietary breadth, i.e., including all potential prey, whereas our estimate of Large Prey index is specifically relevant to prey heavier than 10 kg, i.e., the most substantial prey available to 253 large carnivores (Carbone et al. 2007, 2010). In addition, the Levins index accounts for evenness of 254 255 prey consumption. None of our focal carnivore species showed a decrease of diet breadth, as indexed by the Levins index, with the increase of large prey richness. We suggest that our results do 256 not provide support to our prediction *ii* (i.e., a narrowing of diet breadth with increasing prey 257 richness), which would have been expected under an 'optimal foraging' scenario (MacArthur & 258 Pianka 1966; Charnov 1976). 259

A major caveat to our interpretation is that richness may represent an array of suitable or 260 unsuitable prey with varying levels of population abundance across studies. Prey numerical or 261 biomass densities affect food habits of carnivores (e.g., Karanth & Sunquist 1995; Hayward & 262 263 Kerley 2005; Jedrzejewski et al. 2012; Lovari et al. 2015). Ecological factors such as predation, interspecific competition, climate or human-driven factors such as poaching, over-hunting and/or 264 habitat manipulation may influence prev populations differently across studies (e.g., Karanth & 265 Sunquist 1995; Arsenault & Owen-Smith 2002; Sinclair et al. 2003; Owen-Smith 2008; Ferretti et 266 al. 2015; Sandom et al. 2017), which may lead to non-linear relationships between prey richness 267 and density. Environmental productivity, which may influence herbivore diversity (Danell et al. 268

1996; Prins and Olff 2002), is not always associated with prey density or biomass (e.g., if one or 269 270 more prev are rare). Productive environments may have more finely-partitioned niches, i.e., so that 271 individuals in any given niche are no more abundant than they would be in environments of lower productivity. Moreover, productive environments could allow the persistence of species that are 272 more easily preved on (by allowing them to have higher reproductive rates that offset the mortality 273 caused by predation). Additionally, within the same prev species, individuals belonging to different 274 age/weight classes can also show different vulnerability to predation (e.g., Kruuk and Turner 1967; 275 Fuller 1989; Foster et al. 2010), implying that they may be considered as different 'resource states' 276 (Krebs 1999). However, data on use and availability of different age/weight classes is seldom 277 278 available, and the use by predators of individuals belonging to different age classes can be particularly difficult to assess in the field, especially for studies based on scat analyses. 279 Studies relating diet breadth and prey density across areas with different prey richness are 280 281 desirable. Densities of all potential large prey species should be needed to evaluate the relationships between diet breadth, prey density and prey richness. Unfortunately, estimates of the densities for 282 all potential prey species - especially for rare ones - are sparse and heterogeneous across studies. It 283 is seldom possible to describe the whole prey spectrum and define actual prey availability, 284 preventing us from accounting for that in our analyses. Nevertheless, a positive relationship 285 286 between productivity and herbivore abundance/biomass has been reported at local (Coe et al. 1976;

Danell et al. 1996) and continental scales (Pettorelli et al. 2009, but see Santini et al. 2018), which would support our conclusions. Additionally, prey diversity has been shown to be a fundamental determinant of food habits, potentially more than prey density (Service et al. 2019). Our review represents a first test of the relationships between diet breadth and prey richness for an important group of large mammals. Further work is necessary to explore relationships between environmental productivity, prey density, prey diversity and carnivore food habits and diet breadth, especially in the light of globally increasing anthropogenic pressure and its potential effects on prey communities

294 (Sandom et al. 2017).

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Inferences from analyses of Levins index were largely confirmed by restricting the analysis 295 296 to the use of large prey. Potential exceptions were the snow leopard and dhole, for which the number of Large Prey index seemed to decrease with increasing richness of large prey. This finding 297 may indicate that these carnivores increase specialisation with growing prey richness. However, 298 caution is required in the interpretation of these results, because of the relatively small sample size 299 and because of the width of confidence intervals of estimated coefficients, which were very close to 300 301 including values of zero. The snow leopard is a solitary carnivore, adapted to hardly accessible habitats with low productivity and steep, rocky terrain where prey density is typically low 302 (McCarthy et al. 2017). Over evolutionary time, the presence of potential, superior competitors such 303 304 as the wolf (Bocci et al. 2017), leopard (Lovari et al. 2013a) and tiger (Wang & Macdonald 2009), may have forced this felid to "marginal" areas hardly accessible to competitors, and to specialise on 305 prey living in those areas (Lovari et al. 2013b; Lovari & Mishra 2016). This specialisation may be 306 307 associated to higher selectivity in prey-rich areas where, presumably, competitors are present; in turn, that would favour coexistence with other large predators. However, not surprisingly especially 308 earlier studies on this large cat were based on a relatively small number of samples (10 out of 22 309 studies included less than 50 scats: Schaller 1977; Schaller et al. 1988a-b; Bagchi and Mishra 2006; 310 Wegge et al. 2012), suggesting that more data are needed to support our results. The dhole usually 311 312 coexists with larger predators such as the leopard and the tiger (e.g., Karanth & Sunquist 1995; Andheria et al. 2006; Wang & Macdonald 2009) but has extensive overall in dietary preferences 313 with these two species (Hayward et al. 2014). Our analyses show that the tiger increases its use of 314 315 large ungulates in prey-rich communities. These large cats are dominant over dhole and can kill them (Schaller 1967; Johnsingh 1992). Thus, dhole may be expected to limit the risk of encounters 316 with dominant competitors by selecting different prey, thus increasing their specialisation, which 317 would ultimately favour interspecific coexistence (Karanth & Sunquist 1995). If so, the reduction in 318 the number of large prey used with growing prey richness would be a result of a greater 319 specialisation triggered by prey partitioning with larger, superior competitors. 320

No linear association was supported between carnivore mean body mass and the diet 321 322 breadth-prev richness relationship, thus not supporting our prediction (*iii*). This result suggests that, among large carnivores, body size per se was not the main determinant of the increase of diet 323 breadth with growing prey richness. Thus, other factors could be influential (e.g., carnivore 324 gregariousness, carnivore/prey habitat use, and prey abundance). Nevertheless, four out of the five 325 largest carnivore species showed an increase of dietary breadth (as estimated by the Levins index) 326 and all five showed an increase of the Large Prey index with growing prey richness. For the tiger, 327 results were inconsistent between indices but the sign of both estimated relationship was positive. 328 Apart from these five species, the wolf showed an increase of diet breadth with growing prey 329 330 richness: this result could be easily explained considering the behaviour and ecology of this predator, which is gregarious, very adaptable, and often the top-predator in the areas where it lives 331 (Mech 1970; Mech & Boitani 2004). The Eurasian lynx also showed an increase of diet breadth and 332 the Large Prey index with growing richness of large prey. The Eurasian lynx was often the top 333 predator in the study areas included in this review (65% of 17 studies). However, we recommend 334 caution in interpreting the results for this species, because (i) they are based on a sample of 17 335 studies, (ii) the range of available large prey was only one-to-five, and (iii) for values of one, two 336 and three large prey available, only one, three and one study were found, respectively. 337

338

339 Evolutionary implications and future research directions

Interactions between carnivores are largely shaped by behavioural interference, escalating up to the
killing of smaller species by larger competitors, and affecting behaviour, distribution and/or
numbers of inferior competitors (Palomares & Caro 1999; Linnell & Strand 2000; Donadio &
Buskirk 2006). Interspecific dietary overlap is expected to increase the probability of interspecific
encounters, thus promoting the occurrence of interspecific killing (Palomares & Caro 1999;
Donadio & Buskirk 2006). Our results strongly suggest that top predators use a greater number of
large prey with growing prey richness. These carnivores are expected to encounter little competition

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in selecting areas with high prey density, whereas movements, activity, space or prey of subdominant carnivores are expected to be influenced by those of larger, dominant ones (e.g., Schaller
1967, 1972; Durant 1998; Vanak et al. 2013). Apex predators seem to take advantage of growing
prey richness by increasing their realised diet breadth by taking both larger and smaller prey
(Gittleman 1985; Radloff and do Toit 2004), behaving as generalists, with different degrees of
intensity.

The other carnivores we studied showed relatively stable diet breadths with increasing prey 353 richness, with diet breadth possibly even decreasing for snow leopards and dhole. Interspecific 354 competition may strongly affect carnivore guilds (Palomares & Caro 1999, Linnell & Strand 2000). 355 356 Over evolutionary time, inferior competitors are expected to have developed behavioural tactics to limit encounters with superior ones, involving food habits, space/habitat use and/or temporal 357 activity patterns (e.g., Mills & Biggs 1993; Durant 1998; Karanth & Sunquist 2000; Odden et al. 358 359 2010; Vanak et al. 2013). For example, subordinate carnivores may be forced to select marginal areas with relatively low densities of top-predators, in spite of the local low density of prey (e.g., 360 African wild dog: Mills & Gorman 1997; cheetah: Durant 1998; leopard: Odden et al. 2010), or 361 show morphological/behavioural adaptations to capture rapidly consumable prey, maximising the 362 amount of meat that they can consume before kleptoparasites arrive (cheetah: Hayward et al. 2006). 363 364 Accordingly, our results show that subordinate large carnivores did not increase their diet breadth with growing prey richness, suggesting a role of larger competitors in limiting that increase. Loss of 365 large prey, e.g. through poaching, over-hunting or habitat manipulation, is a major threat to the 366 367 conservation of large carnivores (e.g., Carbone et al. 2010; Estes et al. 2011; Ripple et al. 2014; Sandom et al. 2017). Prey depletion may lead top predators to increase the use of secondary prey, 368 which would further increase "niche suppression" of subordinate carnivores (Creel et al. 2018). If 369 so, subordinate carnivores may be affected not only by the loss of their main prey, but also by that 370 of prey preferred by dominant, apex predators. Further work is required to integrate information on 371 prey diversity, prey abundance (where available), presence/numbers of competitors, spatially 372

explicit proxies for environmental productivity (Pettorelli et al. 2009) and carnivore diet breadth. 373 374 Moreover, our results suggest that different carnivores tend to be found in areas showing different ranges of prey richness. Future research should evaluate whether results observed at the intra-375 specific level are confirmed at the inter-specific one, i.e. whether carnivores living in prey-rich 376 areas tend to take advantage of this richness through a broader diet breadth or rather being more 377 selective, thus showing a narrower diet breadth. 378

We conclude that (i) there is no evidence that large carnivores increase their selectivity with 379 growing large prey richness except, possibly, the snow leopard and the dhole; (ii) the largest and 380 most dominant carnivores increase their dietary breadth with growing prey richness; and (*iii*) the 381 382 dietary breadth of subordinate large carnivores appears to be limited by interspecific competition with dominant predators. Consequently, the realised dietary niche of sub-dominant carnivores may 383 tend to show considerable "conservatism" (Wiens and Graham 2005). If so, we suggest that 384 resource partitioning would be more important in shaping the dietary niches of smaller, inferior 385 competitors than those of dominant, superior ones. 386

References 388

387	
388	References
389	Andheria AP, Karanth KU & Kumar NS (2007) Diet and prey profiles of three sympatric large
390	carnivores in Bandipur Tiger Reserve, India. Journal of Zoology 273: 169–175.

Angerbjörn A, Eide NE, Dalén L, Elmhagen B, Hellström P, Ims RA, Killengreen S, Landa A, 391

Meijer T, Mela M, Niemimaa J, Norén K, Tannerfeldt M, Yoccoz NG, Henttonen H (2013) 392

393 Carnivore conservation in practice: replicated management actions on a large spatial scale.

Journal of Applied Ecology 50: 59-67. 394

- Arsenault R, Owen-Smith N (2002) Facilitation versus competition in grazing herbivore 395 assemblages. Oikos 97:313–318. 396
- Bagchi S, Mishra C (2006) Living with large carnivores: predation on livestock by the snow leopard 397 (Uncia uncia). Journal of Zoology 268:217–224 398

399	Birkhofer K, Wolters V (2012) The global relationship between climate, net primary production and
400	the diet of spiders. Global Ecology and Biogeography 21: 100-108.

- 401 Bocci A, Lovari S, Zafar Khan M, Mori E (2018) Sympatric snow leopards and Tibetan wolves:
- 402 coexistence of large carnivores with human-driven potential competition. *European Journal*403 *of Wildlife Research* 63:92.
- Buisson L, Blanc L, Grenouillet G (2008) Modelling stream fish species distribution in a river
 network: the relative effects of temperature versus physical factors. *Ecology of Freshwater Fish* 17: 244–257.
- Carbone C, Mace GM, Roberts SC, Macdonald DW (1999) Energetic constraints on the diet of
 terrestrial carnivores. Nature 402:442.
- Carbone C, Teacher A, Rowcliffe JM (2007) The costs of carnivory. *PLoS Biology* 5:e22. https
 ://doi. org/10.1371/journ al.pbio.00500 22
- 411 Carbone C, Pettorelli N, Stephens PA (2010) The bigger they come, the harder they fall: body size
 412 and prey abundance influence predator-prey ratios. *Biology Letters* 7:312-315.
- Chakrabarti S, Jhala YV, Dutta S, Qureshi Q, Kadivar RF, Rana VJ (2016) Adding constraints to
 predation through allometric relation of scats to consumption. *Journal of Animal Ecology*85:660–670.
- Charnov EL (1976) Optimal foraging, the Marginal Value Theorem. *Theoretical Population Biology* 9: 129-136.
- Ciucci P, Boitani L, Pelliccioni ER, Rocco M, Guy I (1996) A comparison of scat-analysis methods
 to assess the diet of the wolf *Canis lupus*. *Wildlife Biology* 2: 37-48.
- Coe MJ, Cumming DH, Phillipson J (1976) Biomass and production of large African herbivores in
 relation to rainfall and primary production. *Oecologia* 22:341–354.
- 422 Colwell RK, Futuyma DJ (1971) On the measurement of niche breadth and overlap. *Ecology* 52:
- 423 567–576.

- 424 Creel S, Matandiko W, Schuette P, Rosenblatt E, Sanguinetti C, Banda K, Vinks M, Becker M
- 425 (2018) Changes in African large carnivore diets over the past half-century reveal the loss of

large prey. *Journal of Applied Ecology* DOI: 10.1111/1365-2664.13227

- Danell K, Lundberg P, Niemela P (1996) Species richness in mammalian herbivores patterns in the
 boreal zone. *Ecography* 19:404-409.
- 429 Donadio E, Buskirk SW (2006) Diet, morphology and interspecific killing in Carnivora. *American* 430 *Naturalist* 167:524–536.
- 431 Durant SM (1998) Competition refuges and coexistence: an example from Serengeti carnivores.
 432 *Journal of Animal Ecology* 67: 370–386.
- 433 Estes J, Terborgh J, Brashares JS, Power ME, Berger J, Bond W, Carpenter SR, Essington TE, Holt
- 434 RD, Jackson JB, Marquis RJ, Oksanen T, Paine RT, Pikitch EK, Ripple WJ, Sandin SA,
- Scheffer M, Schoener W, Shurin JB, Sinclair ARE, Soulé ME, Virtanen R, Wardle DA (2011)
 Trophic downgrading of Planet Earth. *Science* 333: 301-306.
- 437 Ferretti F, Corazza M, Campana I, Pietrocini V, Brunetti C, Scornavacca D, Lovari S (2015)
- 438 Competition between wild herbivores: reintroduced red deer and Apennine chamois.

439 *Behavioral Ecology* 26: 550-559.

- 440 Forister M L, Novotny V, Panorska AK, Baje L, Basset Y, Butterill PT, Cizek L, Coley PD, Dem F,
- 441 Diniz IR, Drozd P, Fox M, Glassmire A, Hazen R, Hrcek J, Jahner JP, Kama O, Kozubowski
- 442 TJ, Kursar TA, Lewis OT, Lill J, Marquis RJ, Miller SE, Morais HC, Murakami M, Nickel H,
- 443 Pardikes N, Ricklefs RE, Singer MS, Smilanich AM, Stireman JO, Villamarín-Cortez S,
- 444 Vodka S, Volf M, Wagner DL, Walla T, Weiblen GD, Dyer LA (2015) The global
- distribution of diet breadth in insect herbivores. *Proceedings of the National Academy of*
- 446 *Sciences* doi: 10.1073/pnas.142304211
- 447 Foster RG, Harmsen BJ, Valdes B, Pomilla C, Doncaster CP (2010) Food habits of sympatric
- jaguars and pumas across a gradient of human disturbance. *Journal of Zoology* 280:309-318.

449	Fuller TK (1989) Population Dynamics of Wolves in North-Central Minnesota. Wil	llife	
450	Monographs 105:3-41.		

- 451 Gittleman JL (1985) Carnivore body size: ecological and taxonomic correlates. *Oecologia* 67:540452 554.
- Grassman LI, Tewes ME, Silvy NJ, Kreetiyutanot K (2005) Ecology of three sympatric felids in a
 mixed evergreen forest in North-Central Thailand. *Journal of Mammalogy* 86:29-38.
- Hatton IA, McCann KS, Fryxell JM, Davies TJ, Smerlak M, Sinclair ARE, Loreau M (2015) The
 predator-prey power law: biomass scaling across terrestrial and aquatic biomes. *Science* 349
- 457 (6252), aac6284 DOI: 10.1126/science.aac6284
- 458 Hayward MW, Hofmeyr M, O'Brien J, Kerley GIH (2006). Prey preferences of the cheetah
- *Acinonyx jubatus*: morphological limitations or the need to capture rapidly consumable prey
 before kleptoparasites arrive? *Journal of Zoology 270*: 615-627. doi: DOI: 10.1111/j.
- Hayward MW, Kerley GIH (2005) Prey preferences of the lion (*Panthera leo*). Journal of Zoology
 267:309-322.
- Hayward MW, Lyngdoh S, Habib B (2014). Diet and prey preferences of dholes (*Cuon alpinus*):
- dietary competition within Asia's apex predator guild. *Journal of Zoology* 294:255-266.

465 Hebblewhite M, White CA, Nietvelt CG, McKenzie JA, Hurd TE, Fryxell JM, Bayley SE, Paquet P

466 (2005) Human activity mediates a trophic cascade caused by wolves. *Ecology* 86:2135–2144

- 467 Hofmann RR (1989) Evolutionary steps of ecophysiological adaptation and diversification of
 468 ruminants: a comparative view of their digestive system. *Oecologia* 78:443–457.
- Hurlbert SH (1978) The measurement of niche overlap and some relatives. *Ecology* 59:67-77.
- 470 Jarman PJ (1974) The social organisation of antelope in relation to their ecology. *Behaviour*
- 471 48:215-267.
- 472 Jędrzejewski W, Jędrzejewska B, Okarma H, Schmidt K, Zub K, Musiani M (2000) Prey selection
- 473 and predation by wolves in Bialowieza Primeval Forest, Poland. *Journal of Mammalogy*
- 474 81:197–212

- Johnsingh AJT (1992) Prey selection in three large sympatric carnivores in Bandipur. *Mammalia*56:517-526.
- 477 Jones KE, Bielby J, Cardillo M, Fritz SA, O'Dell J, Orme CDL, Safi K, Sechrest W, Boakes EH,
- 478 Carbone C, Connolly C, Cutts MJ, Foster JK, Grenyer R, Habib M, Plaster CA, Price SA,
- 479 Rigby EA, Rist J, Teacher A, Bininda-Emonds ORP, Gittleman JL, Mace GM, Purvis A
- 480 (2009) PanTHERIA: a species-level database of life history, ecology, and geography of extant
- and recently extinct mammals. *Ecology* 90:2648.
- 482 Karanth KU, Sunquist ME (1995) Prey selection by tiger, leopard and dhole in tropical forests.
- 483 *Journal of Animal Ecology* 64: 439–450.
- 484 Karanth KU, Sunquist ME (2000) Behavioural correlates of predation by tiger (*Panthera tigris*),
- leopard (*Panthera pardus*) and dhole (*Cuon alpinus*) in Nagarahole, India. *Journal of Zoology*250: 255–265.
- 487 Khan U, Lovari S, Ali Shah S, Ferretti F (2018) Predator, prey and humans in a mountainous area:
- loss of biological diversity leads to trouble. *Biodiversity and Conservation* 27: 2795-2813.
- Khorozyan I, Ghoddousi A, Soofi M, Waltert M (2015) Big cats kill more livestock when wild prey
 reaches a minimum threshold. *Biological Conservation* 192:268–275,
- Khorozyan I, Lumetsberger T, Ghoddousi A, Soofi M, Waltert M (2017) Global patterns in biomass
 models describing prey consumption by big cats. *Mammal Review* 47:124-132.
- 493 Krebs CJ (1999) Ecological methodology. Addison Wesley Longman, Boston
- Kruuk H (1976) Feeding and social behaviour of the striped hyaena (*Hyaena vulgaris* Desmarest).
 East African Wildlife Journal 14:91–111.
- 496 Kruuk H, Parish R (1981) Feeding specialization of the European badger *Meles meles* in Scotland.
- 497 *Journal of Animal Ecology* 50:773–788
- Kruuk H, Turner M (1967) Comparative notes on predation by lion, leopard, cheetah and wild dog
 in the Serengeti area East Africa. *Mammalia* 3:1-27.
- Levins R (1968) Evolution in changing environments. Princeton: Princeton University Press.

501	Linnell JDC, Strand O (2000) Interference interactions, co-existence and conservation of
502	mammalian carnivores. <i>Diversity and Distributions</i> 6: 169–176.

- Lovari S, Mishra C (2016) Living on the edge: depletion of wild prey and survival of the snow
 leopard. In: McCarthy T, Mallon D (eds) Snow leopards. Elsevier, pp 69-76.
- Lovari S, Minder I, Ferretti F, Mucci N, Randi E, Pellizzi B (2013a) Common and snow leopards
 share prey, but not habitats: competition avoidance by large predators? *Journal of Zoology*291: 127–135.
- 508 Lovari S, Ventimiglia M, Minder I (2013b) Food habits of two leopard species, competition,
- climate change and upper treeline: a way to the decrease of an endangered species? *Ethology Ecology and Evolution* 25: 305–318.
- Lovari S, Pokheral CP, Jnawali SR, Fusani L, Ferretti F (2015) Coexistence of the tiger and the
 common leopard in a prey-rich area: the role of prey partitioning. *Journal of Zoology* 295:
 122-131.
- Lumetsberger T, Ghoddousi A, Appel A, Khorozyan I, Walter M, Kiffner C (2017) Re-evaluating
 models for estimating prey consumption by leopards. *Journal of Zoology* 203:201–210
- 516 MacArthur RH, Pianka ER (1966) On optimal use of a patchy environment. *The American*
- 517 *Naturalist* 100: 603-699.
- 518 Macdonald D (1983) The ecology of carnivore social behaviour. *Nature* 301:379-384.
- 519 Macdonald DW, Loveridge AJ, Nowell K (2010) *Dramatis personae*: an introduction to wild felids.
- 520 In Biology and conservation of wild felids. (eds. D.W. Macdonald & A. J. Loveridge), pp.
- 521 563–584. Oxford University Press, Oxford, UK.
- McCarthy T, Mallon D, Jackson R, Zahler P, McCarthy K (2017) *Panthera uncia*. The IUCN Red
 List of Threatened Species 2017: e.T22732A50664030.
- 524 http://dx.doi.org/10.2305/IUCN.UK.2017-2.RLTS.T22732A50664030.en. Downloaded on 16
- 525 February 2019.

Mech LD, Boitani L (2003) Wolves: behaviour, ecology and conservation. The University of
Chicago Press, Chicago

- 528 Mech LD, Peterson RO (2003) Wolf-prey relations. In: Mech LD, Boitani L (eds) Wolves:
- behaviour, ecology and conservation. The University of Chicago Press, Chicago, pp 131–157
- 530 Meriggi A, Lovari S (1996) A review of wolf predation in southern Europe: does the wolf prefer
- wild prey to livestock? *Journal of Applied Ecology* 33:1561–1571
- Mills MGL, Biggs HC (1993) Prey apportionment and related ecological relationships between
 large carnivores in Kruger National Park. *Symposium of the Zoological Society of London*
- **534 65:253-268**.
- Mills MGL, Gorman ML (1997) Factors affecting the density and distribution of wild dogs in the
 Kruger National Park. *Conservation Biology* 11:1397-1406.
- 537 Odden M, Wegge P, Fredriksen T (2010) Do tigers displace leopards? If so, why? *Ecological*538 *Research* 25: 875–881.
- Olff H, Ritchie ME, Prins HHT (2002) Global environmental controls of diversity in large
 herbivores. *Nature* 415: 901-904.
- 541 Owens MJ, Owens DD (1978) Feeding ecology and its influence on social organization in brown
- 542 hyenas (*Hyaena brunnea*, Thunberg) of the Central Kalahari Desert. *East African Wildlife*543 *Journal* 16:113-135.
- Owen-Smith N (2008) Changing vulnerability to predation related to season and sex in an African
 ungulate assemblage. *Oikos* 117:602-610.
- Palomares F, Caro TM (1999) Interspecific killing among mammalian carnivores. *American Naturalist* 153: 492–508.
- Pettorelli N, Bro-Jørgensen J, Durant SM, Blackburn T, Carbone C (2009) Energy availability and
 density estimates in African ungulates. *American Naturalist* 173: 698-704.
- 550 R Core Team (2018). R: A language and environment for statistical computing. R Foundation for
- 551 Statistical Computing, Vienna, Austria. URL <u>https://www.R-project.org/</u>.

552	Rabinowitz A, Andau P, Chai PPK (1987) The clouded leopard in Malaysian Borneo. Oryx 21:
553	107-111
554	Radloff FGT, du Toit JT (2004) Large predators and their prey in a southern African savanna: a
555	predator's size determines its prey size range. Journal of Animal Ecology 73:410-423.
556	Ripple WJ, Estes JA, Beschta RL, Wilmers CC, Ritchie EG, Hebblewhite M, Berger J, Elmhagen
557	B, Letnic M, Nelson MP, Schmitz OJ, Smith DW, Wallach AD, Wirsing AJ (2014) Status and
558	ecological effects of the World's largest carnivores. Science https://doi.org/10.1126/scien
559	ce.12414 84
560	Sandom CJ, Williams J, Burnham D, Dickman AJ, Hinks AE, Macdonald EA, Macdonald DW
561	(2017) Deconstructed cat communities: quantifying the threat to felids from prey defaunation.
562	Diversity and Distributions 2017:1-13 DOI: 10.1111/ddi.12558
563	Santini L, Isaac NJB, Maiorano L, Ficetola GF, Huijbregts MAJ, Carbone C, Thuiller W (2018)
564	Global drivers of population density in terrestrial vertebrates. Global Ecology and
565	Biogeography 27:968-979.
566	Schaller GB (1967) The deer and the tiger. Chicago: University of Chicago Press.
567	Schaller GB (1972) The Serengeti lion: a study of predator/prey relations. Chicago: University of
568	Chicago Press.
569	Schaller GB (1977) Mountain monarchs: wild sheep and goats of the Himalaya. University of
570	Chicago Press, Chicago, USA.
571	Schaller GB, Hong L, Talipu, Junrang R, Mingjiang Q (1988a) The snow leopard in Xinjiang,
572	China. <i>Oryx</i> 22:197-204.
573	Schaller GB, Junrang R, Mingjiang Q (1988b) Status of the snow leopard Panthera uncia in
574	Quinghai and Gansu Provinces, China. Biological Conservation 45:179-194.
575	Schoener TW (1971) Theory of feeding strategies. Annual Reviews of Ecology and Systematics
576	2:369-404.

- 577 Service CN, Bateman AW, Adams MS, Artelle KA, Reimchen TE, Paquet PC, Darimont CT (2019)
- Salmonid species diversity predicts salmon consumption by terrestrial wildlife. *Journal of Animal Ecology* 88:392-404.
- Sinclair ARE, Mduma S, Brashares JS (2003) Patterns of predation in diverse predator-prey system.
 Nature 425:288–290
- Stein AB, Fuller TK, Marker LL (2013) Brown hyaena feeding ecology on Namibian farmlands. *South African Journal of Wildlife Research* 43:27-32.
- Vanak AT, Fortin D, Thaker M, Ogden M, Owen C, Greatwood S, Slotow R (2013) Moving to stay
- in place: behavioral mechanisms for coexistence of African large carnivores. *Ecology* 94:

586 2619–2631.

- Walther GR, Post E, Convy P *et al.* (2002) Ecological responses to recent climate change. *Nature*416:389-395.
- Walther GR, Roques A, et al (2009) Alien species in a warmer world: risks and opportunities.
 Trends in Ecology and Evolution 24:686-693.
- 591 Wang SW, Macdonald DW (2009) Feeding habits and niche partitioning in a predator guild
- composed of tigers, leopards and dholes in a temperate ecosystem in central Bhutan. *Journal of Zoology* 277:275-283.
- Wegge P, Odden M, Pokharel CP, Storaas T (2009) Predator-prey relationships and responses of
 ungulates and their predators to the establishment of protected areas: a case study of tigers,
 leopards and their prey in Bardia National Park, Nepal. *Biological Conservation* 142:189-202.
- 597 Wegge P, Shrestha R, Flagstad Ø (2012) Snow leopard *Panthera uncia* predation on livestock and
- wild prey in a mountain valley in northern Nepal: implications for conservation management. *Wildlife Biology* 18:131-141.
- Wiens JJ, Graham C (2005) Niche conservatism: integrating evolution, ecology, and conservation
 biology. *Annual Review of Ecology, Evolution and Systematics* 36:519-539.

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602	Y arnell RW, Phipps WL, Burgess LP, Ellis JA, Harrison SWR, Dell S, MacTavish D, MacTavish
603	LM, Scott DM (2013) The influence of large predators on the feeding ecology of two African
604	mesocarnivores: the black-backed jackal and the brown hyaena. South African Journal of
605	Wildlife Research 43:155-166.

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Table 1. Number of studies per species of large terrestrial carnivores (ordered by increasing body
 weight) and sample type.

			Before	After filtering/pooling			
Species	Body mass (kg)	filte	ring/pool				
		Kills	Scats	Total	Kills	Scats	Total
Dhole	15.8	1	19	20	0	14	14
Eurasian lynx	19.3	13	7	20	11	6	17
African wild dog	22.0	21	1	22	16	0	16
Wolf	31.8	10	86	96	8	81	89
Snow leopard	32.5	0	24	24	0	22	22
Cheetah	50.6	21	3	24	14	3	17
Leopard	52.4	20	71	91	13	62	75
Puma	54.0	20	47	67	19	40	59
Spotted hyaena	63.3	10	17	27	7	16	23
Jaguar	83.9	2	25	27	1	22	23
Lion	158.6	39	9	48	29	6	35
Tiger	161.9	2	37	39	0	25	25
TOTAL		159	346	505	118	297	415

- **Table 2.** Model coefficients (and 0.95 confidence intervals, CIs) for relationships between indices
- of diet breadth and number of available large prey for large terrestrial carnivores. In bold, supported
- relationships (with confidence intervals not including zero).
- 614

	ichness rela	ationships			
Species	Levins		Large Prey index		
	В	B CIs		CIs	
Dhole	-0.214	±0.468	-0.413	±0.396	
Eurasian lynx	0.465	±0.358	0.465	±0.407	
African wild dog	-0.223	±0.614	0.180	±0.339	
Wolf	0.367	±0.125	0.447	±0.098	
Snow leopard	0.007	±0.222	-0.194	±0.180	
Cheetah	0.443	±0.688	0.370	±0.486	
Leopard	0.149	±0.162	0.117	±0.146	
Puma	0.279	±0.213	0.319	±0.204	
Spotted hyaena	0.505	±0.221	0.381	±0.122	
Jaguar	0.520	±0.312	0.386	±0.263	
Lion	0.467	±0.195	0.240	±0.158	
Tiger	0.150	±0.310	0.276	±0.187	

- **Table 3.** Effects of carnivore body mass on the relationships between diet breadth of large
- 618 terrestrial carnivores and prey richness. Model coefficients (*B*), their standard errors (SE) and 0.95
- 619 confidence intervals (CIs), as well as R^2 value of models, are shown.
- 620

Index	Predictor	Predictor <i>B</i> S.E. 0.95 CIs		CIs	<i>R</i> ²	
 Levins	Intercept	-1.790	0.975	-2.963	1.383	0 108
	Log(body mass)	0.228	0.208	-0.235	0.692	0.108
 Large Prey index	Intercept	-1.181	1.119	-3.674	1.312	0.105
	Log(body mass)	0.298	0.239	-0.234	0.830	0.135

623 Figure legends

- **Fig. 1** Relationship between the total diet breadth of large carnivores (Levins index) and the Number of
- 625 species of potential large vertebrate prey available (Prey richness). Fitted relationships and relevant 0.95
- 626 confidence intervals are shown for supported relationships, i.e. those with confidence intervals not including
- 627 '0' values (see Methods for explanations). Red: studies based on kills; black: studies based on scats.
- **Fig. 2** Fitted relationship between the dietary breadth of large carnivores (Large Prey index, i.e. the number
- of frequently used large vertebrate prey) and the Number of species of potential large vertebrate prey
- available. Fitted relationships and relevant 0.95 confidence intervals are shown for supported relationships,
- 631 i.e. those with confidence intervals not including '0' values (see Methods for explanations). Red: studies
- based on kills; black: studies based on scats.
- **Fig. 3** Estimated coefficients (\pm 0.95 confidence intervals) concerning the relationships between dietary
- breadth (a: Levins index; b: Large Prey index) and the number of species of potential large vertebrate prey
- available, for large carnivores, as a function of logged body mass.

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Figure 2





LOG BODY MASS