1	Title: A framework for the identification of hotspots of climate
2	change risk for mammals
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20 Abstract

21 As rates of global warming increase rapidly, identifying species at risk of decline due to 22 climate impacts and the factors affecting this risk have become key challenges in ecology and 23 conservation biology. Here we present a framework for assessing three components of 24 climate-related risk for species: vulnerability, exposure and hazard. We used the relationship between the observed response of species to climate change and a set of intrinsic traits (e.g., 25 weaning age) and extrinsic factors (e.g., precipitation seasonality within a species geographic 26 27 range) to predict, respectively, the vulnerability and exposure of all data-sufficient terrestrial 28 non-volant mammals (3953 species). Combining this information with hazard (the magnitude of projected climate change within a species geographic range) we identified global hotspots 29 30 of species at risk from climate change that includes the western Amazon basin, south-western 31 Kenya, north-eastern Tanzania, north-eastern South Africa, Yunnan province in China, and 32 mountain chains in Papua-New Guinea. Our framework identifies priority areas for 33 monitoring climate change effects on species and directing climate mitigation actions for biodiversity. 34

37 Global mean temperature is projected to rise in excess of 2°C by 2050 relative to 38 preindustrial levels, which is expected to severely impact ecosystems, and cause an extensive loss of ecosystem functions and services (IPCC, 2014). Climate change is expected to impact 39 40 severely on animal populations and to drive massive extinction of species in the future (Isaac, 41 2009), but its effects are also already documented for several taxa. For example, marine 42 species have shifted the poleward edge of their range by 72 km per decade and terrestrial species by 6 km per decade (Scheffers et al., 2016), while both mammal and bird species have 43 44 been negatively affected in terms of reductions in population and range size, survival and reproductive rates (Pacifici et al., 2017). 45

Not all species are expected to respond to climate change alike. Life-history traits, and 46 to a lesser extent taxonomy and geographic distribution, mediate species responses to climate 47 change (Pacifici et al., 2017). Different mammal species, in particular, have shown variable 48 responses to recent climate change according to their life-history traits and the breadth of 49 50 environmental and climatic conditions they experience (Boutin & Lane, 2014; Santini et al. 2016; Pacifici et al., 2017). The above evidence is the basis for the identification of species at 51 higher risk of decline due to climate impacts, and of the factors affecting this risk. This is 52 53 done through a family of methods collectively named climate change vulnerability 54 assessments (CCVA; Foden & Young, 2016). CCVA methods include correlative approaches 55 (which project species niches into the future based on forecasts of climate change), 56 mechanistic approaches (that model individual behaviour in response to climate change), and 57 trait-based analyses. Trait-based CCVAs use species-specific intrinsic life-history traits (e.g., body mass, habitat and diet specialization, reproductive rates) to estimate species 58 59 vulnerability to climate change (Pacifici et al., 2015a).

60	Existing CCVA techniques have drawbacks (Pacifici et al., 2015a). The vast majority
61	of correlative CCVAs (e.g., Hughes et al., 2012; Songer et al., 2012; Baker et al., 2015) use
62	the relationship between the observed geographic distribution of a species and the current
63	climate to infer potential climatically suitable areas for a given species in the future, without
64	considering the intrinsic characteristics of a species that make it more or less predisposed to
65	be impacted by changing climatic conditions (but see Pearson et al., 2014; Visconti et al.,
66	2015). Trait-based CCVAs use a deductive (expert-based) approach rather than an inductive
67	(evidence-based) approach to infer which traits are more important in determining species
68	vulnerability. This introduces an unquantifiable error in the process (Pacifici et al., 2015a).
69	Most trait-based CCVAs also lack a spatial component to identify the areas in which the
70	climatic hazard will be more severe (Laidre et al., 2008; Moyle et al., 2013). Consensus is
71	growing over the benefits of combining different CCVA approaches to overcome some of
72	these limitations outlined above (Pacifici et al., 2015a; Willis et al., 2015; Foden & Young,
73	2016).

74 Here we propose a framework that combines elements of several CCVA approaches 75 under one single statistical model, to assess the overall likelihood of adverse effects of climate 76 change on species based on observed impacts on animal populations. Our framework 77 represents a biodiversity-relevant interpretation of the IPCC general framework for climate 78 impact assessment (IPCC, 2014). The IPCC framework uses the interaction of climate-related hazards, vulnerability (the predisposition to be adversely affected) and exposure (the 79 80 probability to experience the hazard) to derive an overall risk of climate-related impacts to 81 human and natural systems. Here we operationalize the IPCC framework for species as follows: i) vulnerability is the intrinsic predisposition of a species to be adversely affected by 82 83 climate change (given its life-history traits); ii) exposure is the likelihood that climatic refugia 84 exist for the species (related to the span of geographic and climatic variables currently

85	experienced by the species within its natural geographic range); and iii) hazard is the
86	magnitude of projected climate change within the species geographic range.

We applied our framework to 3953 species of data-sufficient terrestrial non-volant 87 mammals, using the multinomial logistic model recently developed by Pacifici et al. (2017). 88 89 Mammals are an ideal taxonomic group for our analysis since they include relatively few species and abundant information is available on their life-history traits (e.g., Jones *et al.*, 90 91 2009; Tacutu et al., 2013). While Pacifici et al. (2017) found species that could have been 92 already negatively impacted by climate change, here for the first time we identified areas with 93 the highest numbers of mammals likely to be at risk in the near future (hotspots of climate 94 risk). Our predictions for the future are based on observed impacts of climate change on 95 mammals, which has never been done before at a global scale. In addition, our framework 96 quantifies the relationship between life-history traits and selected spatial variables and the 97 response of species to climate change, in order to identify hotspots of species likely to be at 98 risk, thus overcoming some of the important limitations associated with trait-based and 99 correlative approaches outlined above. Finally, we propose a set of actions that can be applied 100 according to the risk element/s analysed. This work can be the basis for planning ad hoc 101 conservation actions for those species and areas likely to be more impacted by future climatic 102 changes.

103

104 Materials and methods

105 Hazard

Hazard is usually calculated using climatic scenarios. We used climate predictions
from 11 GCMs (global circulation models) and 3 representative concentration pathway (RCP)
scenarios (RCP2.6, RCP6.0 and RCP8.5) from the WorldClim database at 10 km resolution

(http://www.worldclim.org), to compute the difference between the current and the future
(2050) mean annual temperature in each grid cell. These greenhouse gas concentration
scenarios display different trends in radiative forcing during the 21st century and high human
population growth energy consumption, from the most optimistic scenario (RCP2.6) to the
most severe one (RCP8.5). We then averaged these values across the entire species range
provided by the IUCN, in order to obtain a mean value of temperature change in the areas
where the species is currently distributed.

116

117 *Hotspots of climate risk*

118 For our analysis, we adopted the database and multinomial logistic model used by 119 Pacifici *et al.* (2017), in which species response to recent climate change (negative, positive, 120 unchanged or mixed) was used as dependent categorical variable. The negative response was 121 assigned to a species if >50% of its populations were reported to have undergone declines in 122 the extent of the geographic range, population size, survival or reproductive rates, and body 123 mass. On the contrary, the positive response was assigned if the majority of the populations of 124 a species experienced expansions in the above mentioned metrics. The unchanged response 125 was attributed if no response was observed despite the recorded change in climate. Finally, the mixed response was attributed to species that exhibited a combination of positive and 126 127 negative impacts (see Pacifici et al. 2017 for further details). We used as independent variables those identified as most important for determining an overall climate risk for the 128 129 species in Pacifici et al. 2017, i.e. non-fossorial behaviour, restricted dietary breadth, reduced 130 precipitation seasonality and high minimum altitude, the latter two computed within the current species range (Table 1). In Pacifici et al. (2017), large differences in temperature 131 132 between the present and the past were also important determinant of negative responses. Since

the expected increase in temperatures is likely to be 3-4 times higher than that of the past 100
years, we included the difference in mean annual temperature between the present and the
future as predictor representing the hazard component. In addition, we considered taxonomy

as a fixed effect, in order to control for the non-independence of observed responses across

137 species (Table 1).

138

Variable	Description	Main data sources	Climate risk component
Dietary breadth	Number of categories of food items eaten by a species. In order to identify the species with the highest dietary specialization, the categories of food items were defined as fruit, flowers/nectar/pollen, wood/bark/stems, leaves, seeds/grains/nuts, roots/tubers, sap or other plant fluids, bryophytes, fungi, mammals, birds, reptiles, amphibians, fishes, molluscs, aquatic crustaceans, insects, terrestrial non-insect arthropods, terrestrial worms	Nowak (1999); Wilson & Reeder (2005); IUCN (2016)	Vulnerability
Fossoriality	YES or NO, referred to mammal species adapted/non-adapted to digging and life underground	Wilson & Reeder (2005)	Vulnerability
Order	Taxonomic order to which the species belongs	IUCN (2016)	Vulnerability
Precipitation seasonality	(mm), represents the difference in mean precipitation between the wettest and the driest quarters within a species current distributional range	http://www.worldclim.org/	Exposure
Minimum altitude	(meters), is the minimum altitude at which a species currently lives	Jarvis et al., (2008)	Exposure
Δ temperature	Mean difference between the current (mean 1980-2009) and the future (2050) temperature within a species range	http://www.cru.uea.ac.uk/data	Hazard

Table 1 Description of the independent variables used in the model.

140

141	By using the function <i>predict</i> in R, we predicted the likely response of all species
142	(3953) to projected scenarios of climate change (climate risk). In order to account for the
143	uncertainty in our models, we first predicted the probabilities associated with each of the four
144	options of the response variable, and then we sampled the response category assigned to each
145	species from the multinomial distribution 100 times to derive the mean and standard deviation
146	of the richness of species with a negative response. We then produced richness maps for each
147	climatic scenario to find hotspots of species that are likely to be at greatest risk from climate
148	change in 2050, defined as those species having a higher probability of showing a negative
149	response in the future, according to model predictions. In addition, to determine which areas
150	will be more severely affected in terms of potential loss of the overall local mammalian
151	biodiversity, we divided the number of species at risk in each grid cell by the total number of
152	species in the cell, to obtain a map of the areas hosting the highest proportion of species at
153	greatest risk from climate change.

154

155 Vulnerability and exposure

In order to consider the three components of climate risk both together and 156 157 independently, we also identified the areas with the highest concentration of vulnerable and exposed species. To do that, we ran two different models by using the predictors of the 158 159 multinomial logistic model for climate risk, but held the variables related to vulnerability (i.e., 160 dietary breadth and fossoriality) and exposure (i.e., precipitation seasonality and minimum altitude recorded within the current geographic range), respectively, constant at their mean 161 (numeric variables) or mode (nominal variables), calculated among all 3953 species 162 considered. In both the vulnerability and exposure models, in order to control for the latent 163

164 variables that may affect the responses to climate change that are phylogenetically conserved, 165 we did not change the values for taxonomic order. For example, in order to look at the effect of intrinsic life-history traits, we held the mean difference in temperature, minimum altitude 166 167 within a species range, and precipitation seasonality constant, while we used species-specific 168 values for dietary breadth, fossoriality and taxonomic order. We then stacked the distribution 169 ranges of these species obtained from the IUCN database (www.iucn.org) to derive a richness 170 map of global vulnerability. We applied the same procedure to identify the areas hosting the 171 highest numbers of species likely to be exposed to climate change, but holding constant only 172 the predictors not related to exposure.

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174 Results
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175 *Areas experiencing the greatest changes in temperature*

176 In all the three RCP scenarios considered, the minimum change in mean temperature 177 between the present and the future in a grid cell is $\geq 0.4^{\circ}$ C (0.4°C in RCP2.6, 0.6°C in RCP6.0 178 and 0.8°C in RCP8.5; Fig. 1). As expected, the most severe changes in climate is projected 179 under the RCP8.5 scenario (mean global temperature increase of 2.97°C), followed by the 180 RCP6.0 (2.16°C) and RCP2.6 (1.89°C). Despite the great differences in the magnitude of 181 climate change between the three scenarios, all the three converge in identifying areas that 182 would experience the highest changes in temperatures. These correspond to the northern polar 183 region (Fig. 1), where the predicted increase in temperature is always >1.7°C (RCP2.6: min 184 +1.7°C, mean +3.06°C, max +4.9°C; RCP6.0: min +1.8°C, mean +3.3°C, max +5.2°C; 185 RCP8.5: min $\pm 2.8^{\circ}$ C, mean $\pm 4.7^{\circ}$ C, max $\pm 7.5^{\circ}$ C). In the RCP8.5 scenario, 87.5% of the world will experience an increase in mean annual temperature $>2^{\circ}C$ by 2050 (while in 186 187 RCP6.0 and RCP2.6 these percentages stand at 50% and 37.5%, respectively), and additional

- areas that are likely to be considerably affected by these changes are the Kamchatka Peninsula
- in Russia, the Himalayas, West Russia, the territories on the border between Russia and
- 190 Kazakhstan, and the mountain chains in North America. From Fig. 1 it is evident that, in
- 191 general, the Northern Hemisphere is likely to be more subject to global climate change than
- the Southern Hemisphere, in particular in the RCP2.6 and RCP6.0 scenarios.

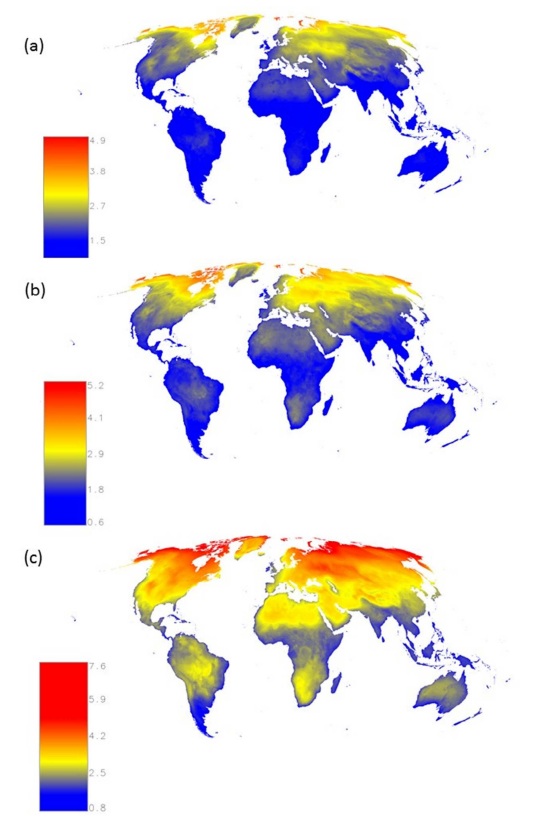


Figure 1 Mean annual temperature difference between the present and the future in 2050 in the a)
RCP2.6, b) RCP6.0 and c) RCP8.5 scenarios.

196 *Species most at risk under projected climate change*

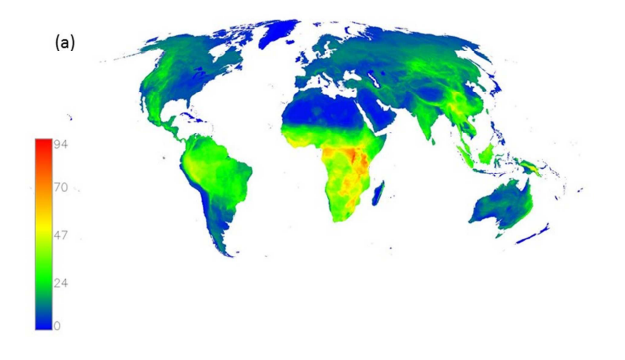
197	Our predictions of species responses to projected climate change were very similar
198	among the three different scenarios, with only 195 species out of 3953 (4.9%) showing
199	differing results. As expected, the number of species showing no response to climate change
200	decreased as the changes in mean annual temperatures became more intense, with 1.34% of
201	the species classified as 'unchanged' in the RCP2.6 scenario, 0.46% in the RCP6.0 and 0
202	species in the RCP8.5. The taxonomic orders having >85% of the species with a likely future
203	negative response in all scenarios were Proboscidea, Dasyuromorphia, Primates,
204	Diprotodontia, Cetartiodactyla, Perissodactyla and Lagomorpha (Tables S1, S2, S3).
205	Surprisingly, considering only the set of species having different responses in the
205 206	Surprisingly, considering only the set of species having different responses in the different scenarios, the majority of them were likely to benefit from increases in mean annual
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206 207 208	different scenarios, the majority of them were likely to benefit from increases in mean annual temperatures (133 species out of 195 had a negative or unchanged response in the RCP2.6 scenario and a positive one in the RCP8.5 scenario, Tables S1, S3). These species mostly
206 207 208 209	different scenarios, the majority of them were likely to benefit from increases in mean annual temperatures (133 species out of 195 had a negative or unchanged response in the RCP2.6 scenario and a positive one in the RCP8.5 scenario, Tables S1, S3). These species mostly included small mammals (belonging to the Rodentia and Eulipotyphla taxonomic orders). On

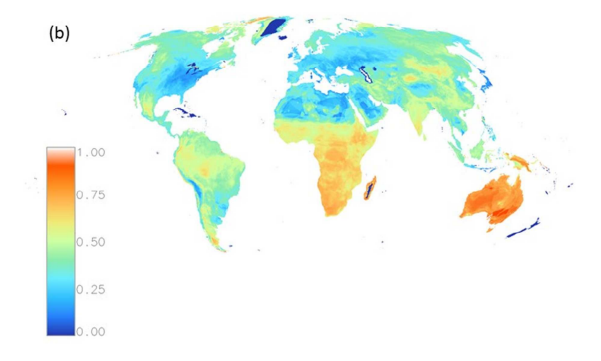
212

213 *Identifying hotspots of species at climate risk*

In general, we found the highest concentration of mammals most likely to be at risk from climate change in mountainous areas. In particular, hotspots of species richness are concentrated in the western Amazon River basin, south-western Kenya, north-eastern Tanzania, north-eastern South Africa, Yunnan province in China, and mountain chains in Papua-New Guinea (Figs. 2a & S1). It is interesting to note that they rarely overlap with areas identified as having the highest probability of experiencing significant changes in mean

220	annual temperatures. In fact, while most of the hotspots are found in the Southern
221	Hemisphere, the differences between mean annual temperature in the RCP scenarios between
222	2050 and the present show that the Northern Hemisphere will experience the greatest changes.
223	When we consider the number of species likely to be negatively impacted by climate
224	change in the future, with respect to the total number of species present in a grid cell, the
225	areas with the highest proportions of species at risk are found in northern Greenland, Tierra
226	del Fuego, lower altitudes of Madagascar, north-west Russia, Mongolia, north-western China,
227	Tibetan Plateau, Papua-New Guinea, Western Australia and New South Wales (Figs. 2b &
228	S2). These areas only partly overlap with the hotspots of species richness in central Africa.



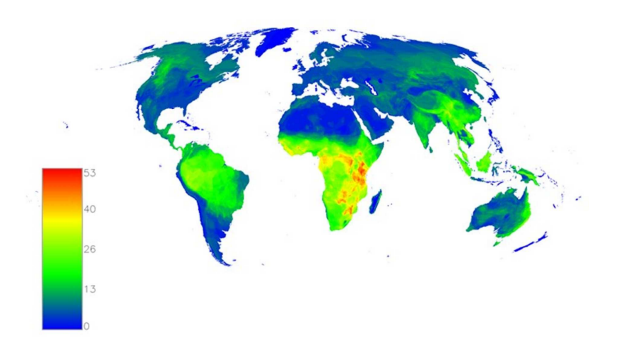




- **Figure 2** Maps of projected negatively impacted species by grid cell in the RCP8.5 scenario. a)
- 231 Richness and b) proportion of species with a negative predicted response.

232	Hotspots of vulnerable-only species
233	According to our models, the areas hosting large numbers of vulnerable species are
234	mostly concentrated in eastern Sub-Saharan Africa (Fig. 3a), specifically in the eastern side of
235	the Democratic Republic of Congo, southern South Sudan and Ethiopia, Kenia, Tanzania,
236	Zambia and north-eastern regions in South Africa.
237	
238	Hotspots of exposed-only species
239	The areas hosting high numbers of exposed species include almost all the hotspots of
240	vulnerable-only species, as well as the Rocky mountains side of Colorado and New Mexico in
241	the United States, central Namibia, the Cape province in South Africa, and the Altai
242	Mountains, a mountain system in central Asia extending through China, Russia, Mongolia
243	and Kazakhstan (Fig. 3b).





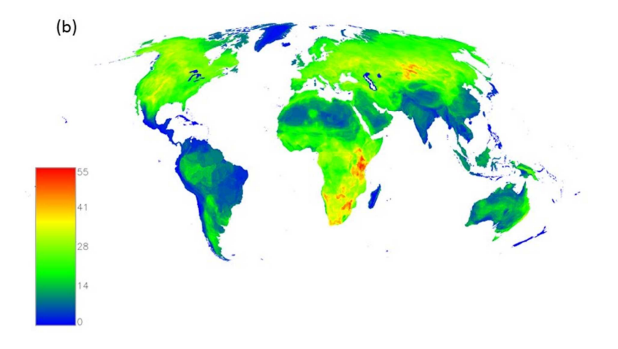


Figure 3 Relative richness of a) vulnerable and b) exposed species according to multinomial logistic

246 models.

247 Discussion

248 The Northern Hemisphere has outpaced the Southern Hemisphere in terms of 249 temperature increase since about 1980. This is mostly due to the fact that the Northern 250 Hemisphere has more land and less ocean than the Southern Hemisphere, and the general rate 251 of warming of the oceans is relatively slow (Friedman et al., 2013). The strong loss of Arctic sea ice and global ocean currents are another possible factor contributing to the Northern 252 253 Hemisphere greater warming. These currents transport heat away from southern oceans and 254 into the northern waters, helping to further warm nearby land areas in the north (Feulner et 255 al., 2013).

256 Physiographic factors such as slope, elevation and topographic convergence influence meteorological elements including precipitation, air temperature, wind, solar insolation, snow 257 accumulation and melt (Dobrowski, 2011). Microtopography can cause large temperature 258 259 differences within a short distance, conditions that would otherwise be observed only over 260 large altitudinal or latitudinal gradients. This suggests that for species living at high altitudes, which are often limited by dispersal, it is not always necessary to migrate several hundred 261 262 meters upslope to avoid warmer conditions (Patsiou et al., 2014). Microrefugia can be found in sites that are consistently decoupled from regional patterns because they are more likely to 263 264 support relict climates. In order to favour species persistence in areas that are likely to experience the greatest changes in climate, identifying potential microclimate refugia at local 265 266 scale would be fundamental for species that have limited ability to move elsewhere.

Despite the fact that the northern territories are warming faster than the southern ones, and that this trend is likely to exacerbate in the coming decades, we found that most species threatened by climate change mainly occur south of the Equator. Oceania is home to the vast majority of marsupials at risk from climate change. Specifically, the tropical montane forests

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271	have undergone major declines due to increasing land conversion in the last decades, and
272	global warming has already allowed changes in agricultural practices, such as a continuing
273	use of the same land with the same crop, with a reduced shortfall period (Brown, 2013). More
274	than 2/3 of the above mentioned species are currently listed in a threat category on the IUCN
275	Red List (IUCN, 2016). An important hotspot of species at risk is Papua-New Guinea.
276	Marsupials are very diversified in this island and are among the species with the larger body
277	mass. As demonstrated by McNab (1994), resource limitation on continental shelf islands
278	impacts larger-bodied mammals, often resulting in extinction. In the face of increasing global
279	warming, island species are unlikely to be able to track the change in climate and move into
280	new suitable climate space. Therefore, in addition to unrestricted logging and hunting (the
281	current most serious threats to marsupials in these areas; Grzimek et al., 2004), the very high
282	rates of temperature change that these mammals are likely to experience in the next decades
283	will probably have deleterious consequences for their survival, possibly leading to the
284	disruption and disaggregation of extant communities (Williams et al., 2007).
285	The second and largest hotspot of species at climate risk includes most of the central
286	and eastern Sub-Saharan Africa. Savannas and forests are home to many of Africa's most
287	prized species belonging to different taxonomic groups, including cheetahs, elephants, lions,
288	black and white rhinos, leopards, eastern gorillas and chimpanzees. Almost all these species
289	are currently threatened with extinction, predominantly due to hunting, habitat loss and
290	overexploitation (Schipper et al., 2008). Vegetation dynamic models predict that large

291 portions (>45%) of today's savannas will be replaced by deciduous woodlands under elevated

temperature and CO_2 concentrations (Scheiter & Higgins, 2009), with a consequent further

reduction of the species suitable habitat and geographic ranges, and increases in their risk of

extinction. In southern Africa, the east-west aridity gradient will probably induce to a

westward shift of the geographic ranges of species towards the moister and cooler areas at

higher elevations (Erasmus *et al.*, 2002; Thuiller *et al.*, 2006). Studies have shown that some
species of ungulates (e.g., roan antelope (*Hippotragus equinus*), tsessebe (*Damaliscus lunatus*), and kudu (*Tragelaphus strepsiceros*) among the antelopes) have suffered reductions
in abundance with decreases in mean annual precipitation in South African national parks
(Ogutu & Owen-Smith, 2003; Ogutu & Owen-smith, 2005), and this trend is likely to become
more severe when projecting increases in aridity conditions (Pacifici *et al.*, 2015b).

302 The third hotspot, which is also a crucial site for primates, is the western portion of the Amazon river, on the border of the Andes between Peru and Ecuador. This area is considered 303 304 one of the most biodiverse of the planet, and hosts more than 20 species of primates (Jenkins et al., 2013; Gouveia et al., 2014) in large portions of intact tropical moist forest (Killeen et 305 306 al., 2007). Despite assumptions on future climatic stability (Finer et al., 2008), we found that 307 relatively small changes in temperatures could have major impacts on the species living in the 308 western Amazon. This is probably due to the fact that these areas have been climatically 309 stable over time (Killeen et al., 2007), therefore it is unlikely that the species inhabiting this region will be able to adapt soon to these changes. Additional threats to these species are 310 311 linked to oil and gas extraction, which had a boom in the 1970s (Finer et al., 2008). The 312 intensification of these activities has already led to both direct and indirect impacts on species including deforestation for access roads, increased logging and hunting, and contamination 313 314 from oil spills and wastewater discharges (Finer et al., 2008), and it is likely to exacerbate 315 when acting in combination with climate change.

We found that the majority of species benefitting from an increase in severity of climate change (i.e., different responses in the different scenarios, changing from negative/unchanged in RCP2.6 to positive in RCP8.5) belong to the orders Rodentia and Eulipotyphla. All these species with a negative response in the RCP2.6 scenario are not adapted to living underground and exploit a relatively restricted number of dietary items.

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321 However, small mammals, in particular rodents, are typically associated with reduced 322 vulnerability and high capacity of adjusting to environmental changes due to the fact that they usually produce high numbers of offspring that need limited parental care, and that they are 323 324 often habitat generalists (Capizzi et al., 2014). Most of the small mammals with different 325 responses in our sample live in tropical areas (Figure S3) and are already adapted to living at 326 high temperatures. In this case, factors other than the life-history traits considered in our 327 analysis are probably more important in determining the response of species to climate 328 change. In addition, despite the general perception that climate change mitigation improves 329 water condition, it has been shown that increased demand for irrigation water for bioenergy 330 crops in mitigation scenarios might result in increased water stress (Hejazi *et al.*, 2015). We 331 can therefore hypothesize that mitigation strategies, as those planned in the RCP2.6 scenario, 332 might be worse than climate change if not designed with careful attention to water resources. 333 This will be particularly important for some of the species of carnivores listed in Table S4, 334 because water is a key element that has both direct (drinking water) and indirect (prey often 335 concentrate around water points) effects on their persistence.

It is interesting to note that high richness of impacted species and the highest 336 337 proportion of species at climate risk overlap only in part of central Africa and Papua-New 338 Guinea (Fig. 2). In contrast to what we found for the hotspots, these areas are generally found 339 in less inhabited regions (Doxsey-Whitfield et al., 2015), usually characterized by extreme 340 environments and climate, e.g., Greenland, deserts of the southern Hemisphere and Tierra del Fuego. Species living in these zones have probably been less subject to human threats like 341 342 hunting and land-use changes because in such extreme places human occupation may have 343 been intermittent (Johnson, 2002), but in the face of continuing climate change, the high 344 degrees of specialization that these species exhibit is likely to place them at great risk of 345 extinction. More importantly, in areas hosting high proportions of species at risk but low

absolute numbers of taxa, if all species decline or go extinct there will be no ecological
replacement, and there is a high probability of losing the ecosystem services and functions
these species provide.

One of the areas with a high proportion of species at climate risk is the Tibetan 349 350 Plateau. Since 1978 the Tibetan Plateau has warmed >0.04 °C/year, the greatest value for the whole China and East Asia. These changes in climatic conditions have been attributed to a 351 352 positive feedback between the increase in grazing pressure, with consequent degradation of pastures, and that of potential evapotranspiration (Du et al., 2004). Analyses of 353 354 meteorological stations, combined with projections of climate models, reveal that the prominent warming and the thawing of permafrost at higher elevations are likely to continue 355 356 in the next decades (Liu *et al.*, 2009), thus leading to a possible reduction of several biomes, 357 such as the temperate desert, alpine steppe, and ice/polar desert (Ni, 2000). These areas are 358 currently inhabited by several species of lagomorphs (pikas and hares), which are restricted to 359 mountainous habitats and cold climates, and by large threatened mammals. Some of these large threatened species are currently recovering thanks to habitat protection, like the Tibetan 360 361 antelope (Pantholops hodgsonii; IUCN SSC Antelope Specialist Group 2016), while many 362 others are already declining due to habitat destruction and overexploitation, for example the leopard (Panthera pardus; Stein et al. 2016) and the white-lipped deer (Cervus albirostris; 363 364 Harris, 2015). The Tibetan Plateau also hosts relatively large numbers of primates belonging 365 to the genus *Macaca*; here climate change is likely to further worsen the conservation status of these species acting in concomitance with hunting, habitat loss and human disturbance, 366 367 which are currently considered the major threats to the species (Boonratana et al., 2008).

When considering only the intrinsic component of climate risk, we found that areas hosting large numbers of vulnerable species overlap well with the sites of high mammalian diversity and endemism, in particular of Cetartiodactyla and Carnivores (Pimm *et al.*, 2014),

371 in central Africa. Here, human population size is expected to increase steeply by 2050, from a 372 minimum of 16% in South Africa, up to 117% in the Democratic Republic of Congo and 127% in Zambia (United Nations, 2015). This projected increase in human population, 373 374 associated with expected increases per-capita consumption, are likely to cause an acceleration 375 in the conversion of remaining forest habitats into human-dominated settlements and 376 agricultural areas (Barrie et al., 2007). This, in turn, is expected to endanger several mammal 377 species in Sub-Saharan Africa, with similar trends in insular Southeast Asia and the Amazon 378 rainforest (Visconti et al., 2015). In the areas inhabited by large numbers of vulnerable 379 species, priority actions should be aimed at mitigating the effects of other threats that may act concurrently with climate change. Temperature has clear effects on the behaviour of most 380 381 animal species, and this might translate in changes in the interactions between predators and 382 their preys (Creel et al., 2016), thus indirectly leading to possible increases in conflicts 383 between humans and wildlife.

384 As already observed in previous studies on other taxa (e.g. birds; Goodenough & Hart, 2013), hotspots of species under high climate threat exposure include areas with the highest 385 386 concentration of species with limited or no availability of climatic refugia. These obviously 387 include mountainous areas, where species are likely to be unable to move upslope to cope 388 with climate change. For example, in our study we found that the Rocky Mountains of 389 Colorado and New Mexico in the United States and the Altai Mountains in central Asia are 390 important centres for exposed species. The latter are home to several species of threatened 391 large mammals, like the snow leopard (Panthera uncia), the Asiatic Wild Dog (Cuon alpinus) 392 and the Siberian Musk Deer (Moschus moschiferus). For these species, translocations or 393 assisted colonization actions could be necessary in the coming decades if changes in climatic conditions exacerbate. The Siberian Musk Deer is already successfully bred in captivity at 394

musk deer farms in Russia and China (Nyambayar *et al.*, 2015), and these captive herds might
be used for future movements to wild populations.

397 Importance of combining intrinsic and extrinsic factors to identify species at high risk from 398 climate change

399 This work brings together information from life-history traits, distribution patterns and 400 predicted future levels of climate change to create a unique assessment of climate risk. Our 401 framework enables the first predictive assessment of the mammal species most at risk from 402 climate change, on the basis of empiric data and considering all the three components of risk. 403 With this work we have demonstrated that considering only the intrinsic characteristics that 404 make a species more vulnerable to climate change is not enough to predict its real climate 405 induced risk. Species displaying similar life-history traits may be differently affected by 406 global warming simply because they live in areas experiencing different rates of climate change. In our analyses, we found that >55% of the species of terrestrial non-volant mammals 407 408 may potentially be negatively impacted by climate change, depending on the climatic 409 scenario. In the Fourth IPCC Report a projected increase of 2-3°C was expected to lead to 410 high risk of extinction approximately 20 to 30% of the species assessed so far (IPCC, 2007). However, in the Fifth and most recent Report (IPCC, 2014), it has been estimated that the 411 412 expected increase in temperatures is likely to exceed the threshold of 4°C by 2100. This 413 translates into increased threats for species. Accounting for both the life-history traits that 414 make a species more vulnerable and the environmental conditions of its geographic range, 415 could in many cases help prioritization actions and make the difference between species 416 extinction or survival. However, due to the paucity of life-history trait data for numerous species, it is often difficult to provide a rapid assessment of their vulnerability to climate 417 418 change. Although mammals are among the best-known taxa, the number of new recognised 419 species is still increasing. For instance, newly described mammals have been found mostly in

420 Madagascar and the Amazon, while the Congo basin still remains scarcely explored (Schipper 421 *et al.*, 2008). These new species are usually poorly known and severely threatened, in part due 422 to their restricted geographic range, and may therefore become rapidly extinct. These new 423 data-deficient species are generally found in tropical forests (Schipper *et al.*, 2008), that well 424 overlap with the hotspots of species at risk from climate change.

425 Changes in the distribution or phenology of species are often detected via long-term surveys. Establishing or expanding systematic monitoring of abundance and distribution of 426 427 species is a very high priority in order to validate species assessments and monitor 428 effectiveness of adaptation. For species identified as most vulnerable, exposed, and that live in places where impacts are likely to occur sooner, intensive monitoring is essential to 429 430 increase our understanding of the ongoing process of climate change, and possibly respond to 431 it. This is not only true for species living in the hotspots, but also for those occupying areas 432 we identified as having the highest rates of mammals at risk. Although these areas often host 433 a reduced number of species, it is essential to monitor their status in order to avoid the loss of entire mammalian communities characterized by peculiar specializations. 434

Life-history traits have been often used in conservation biology to identify the most vulnerable species to environmental changes. The identification of traits specifically related to climatic risk will further strengthen species risk assessments, thus helping to plan appropriate conservation actions in the differently exposed areas and prioritize intervention targets. This is particularly important for many species of mammals living in climatic hotspots, which are almost unknown, to draw attention to them and begin fill some knowledge gaps.

441

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444	Supporting Information captions
445	Table S2 Percentage of predicted species responses by taxonomic group for the RCP2.6
446	scenario
447	Table S2 Percentage of predicted species responses by taxonomic group for the RCP6.0
448	scenario
449	Table S3 Percentage of predicted species responses by taxonomic group for the RCP8.5
450	scenario
451	Table S4 Species showing a change from an "Unchanged"/ "Negative" response in the
452	scenario RCP2.6 to a "Negative" response in the scenario RCP8.5
453	Figure S1 Richness of species with a negative predicted response by grid cell in the a)
454	RCP2.6 and b) RCP6.0 scenarios
455	Figure S2 Proportion of species with a negative predicted response by grid cell in the a)
456	RCP2.6 and b) RCP6.0 scenarios.
457	Figure S3 Richness of species showing a change from an "Unchanged"/ "Negative" response
458	in the scenario RCP2.6 to a "Negative" response in the scenario RCP8.5.
459	
460	

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