RESEARCH ARTICLE

Evaluating expert-based habitat suitability information of terrestrial mammals with GPS-tracking data

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Maarten J. E. Broekman¹ | Jelle P. Hilbers¹ | Mark A. J. Huijbregts¹ Thomas Mueller^{2,3} | Abdullahi H. Ali⁴ | Henrik Andrén⁵ | Jeanne Altmann⁶ | Malin Aronsson^{5,7} | Nina Attias^{8,9} | Hattie L. A. Bartlam-Brooks¹⁰ Floris M. van Beest¹¹ I Jerrold L. Belant¹² Dean E. Bever¹³ Laura Bidner¹⁴ Niels Blaum¹⁵ | Randall B. Boone¹⁶ | Mark S. Boyce¹⁷ | Michael B. Brown^{18,19} Francesca Cagnacci²⁰ | Rok Černe²¹ | Simon Chamaillé-Jammes²² Nandintsetseg Dejid² | Jasja Dekker²³ | Arnaud L. J. Desbiez^{9,24,25} Samuel L. Díaz-Muñoz²⁶ Julian Fennessy¹⁸ Claudia Fichtel²⁷ Christina Fischer²⁸ Jason T. Fisher²⁹ I Ilva Fischhoff³⁰ Adam T. Ford³¹ John M. Fryxell³² | Benedikt Gehr³³ | Jacob R. Goheen³⁴ | Morgan Hauptfleisch³⁵ | A. J. Mark Hewison^{36,37} A. J. Marco Heurich^{38,39,40} Lynne A. Isbell^{14,41} | René Janssen²³ | Florian Jeltsch¹⁵ | Petra Kaczensky^{40,42,43} Peter M. Kappeler²⁷ | Miha Krofel⁴⁴ | Scott LaPoint^{45,46} | A. David M. Latham^{17,47} | John D. C. Linnell^{40,42} | A. Catherine Markham⁴⁸ | Jenny Mattisson⁴² | Emilia Patricia Medici^{24,49} 💿 | Guilherme de Miranda Mourão⁵⁰ 💿 | Bram Van Moorter⁴² 💿 | Ronaldo G. Morato⁵¹ kilo | Nicolas Morellet^{36,37} kilo | Atle Mysterud⁵² kilo | Stephen Mwiu⁵³ | John Odden⁵⁴ | Kirk A. Olson⁵⁵ | Aivars Ornicāns⁵⁶ | Nives Pagon²¹ | Manuela Panzacchi⁴² | Jens Persson⁵ | Tyler Petroelje¹² | Christer Moe Rolandsen⁴² David Roshier⁵⁷ Daniel I. Rubenstein⁶ Sonia Saïd⁵⁸ Albert R. Salemgareyev⁵⁹ Hall Sawyer⁶⁰ | Niels Martin Schmidt^{11,61} | Nuria Selva⁶² | Agnieszka Sergiel⁶² Jared Stabach¹⁹ Jenna Stacy-Dawes⁶³ Frances E. C. Stewart^{29,64} Jonas Stiegler¹⁵ Jonas Viegler¹⁵ Jonas Viegler¹⁵ Jonas Stiegler¹⁵ Jonas Viegler¹⁵ Wiebke Ullmann¹⁵ | Ulrich Voigt⁶⁸ | Jake Wall⁶⁹ | Martin Wikelski^{70,71} | Christopher C. Wilmers⁷² | Filip Zięba⁷³ | Tomasz Zwijacz-Kozica⁷³ Aafke M. Schipper^{1,74} Marlee A. Tucker¹

¹Department of Environmental Science, Institute for Wetland and Water Research, Faculty of Science, Radboud University, Nijmegen, The Netherlands ²Senckenberg Biodiversity and Climate Research Centre, Senckenberg Gesellschaft für Naturforschung, Frankfurt (Main), Germany

³Department of Biological Sciences, Goethe University, Frankfurt (Main), Germany

⁴Hirola Conservation Programme, Garissa, Kenya

⁵Grimsö Wildlife Research Station, Department of Ecology, Swedish University of Agricultural Sciences, Riddarhyttan, Sweden

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⁶ Department of Ecology and Evolutionary Biology, Princeton University, Princeton, New	/ Jersey, USA		
⁷ Department of Zoology, Stockholm University, Stockholm, Sweden			
⁸ Ecology and Conservation Graduate Program, Federal University of Mato Grosso do Su	II, Campo Grande, Mato Grosso d	lo Sul, Brazil	
⁹ Instituto de Conservação de Animais Silvestres (ICAS), Campo Grande, Mato Grosso do	Sul, Brazil		
¹⁰ Structure and Motion Lab, Royal Veterinary College, London, UK			
¹¹ Department of Bioscience, Aarhus University, Roskilde, Denmark			
¹² Global Wildlife Conservation Center, State University of New York College of Environ	mental Science and Forestry, Syra	acuse, New York, USA	
¹³ Department of Fisheries and Wildlife, Michigan State University, East Lansing, Michiga	an, USA		
¹⁴ Department of Anthropology, University of California, Davis, California, USA			
¹⁵ Plant Ecology and Nature Conservation, University of Potsdam, Potsdam, Germany			
¹⁶ Department of Ecosystem Science and Sustainability, Colorado State University, Fort (Collins, Colorado, USA		
¹⁷ Department of Biological Sciences, University of Alberta, Edmonton, Alberta, Canada			
¹⁸ Giraffe Conservation Foundation, Eros, Namibia			
¹⁹ Conservation Ecology Center, Smithsonian National Zoo and Conservation Biology Ins	titute, Front Royal, Virginia, USA		
²⁰ Department of Biodiversity and Molecular Ecology, Research and Innovation Centre, F	Fondazione Edmund Mach, Trento	o, Italy	
²¹ Slovenia Forest Service, Ljubljana, Slovenia			
²² CEFE, Univ Montpellier, CNRS, EPHE, IRD, Univ Paul Valéry Montpellier 3, Montpellie	r, France		
²³ Bionet Natuuronderzoek, Stein, Netherlands			
²⁴ IPÊ (Instituto de Pesquisas Ecológicas; Institute for Ecological Research), São Paulo, Br	azil		
²⁵ Royal Zoological Society of Scotland (RZSS), Edinburgh, UK			
²⁶ Department of Microbiology and Molecular Genetics, University of California, Davis, C	California, USA		
²⁷ German Primate Center, Behavioral Ecology and Sociobiology Unit, Göttingen, German	ny		
²⁸ Faunistics and Wildlife Conservation, Department of Agriculture, Ecotrophology, and Bernburg, Germany	Landscape Development, Anhalt	University of Applied Sciences,	
²⁹ School of Environmental Studies, University of Victoria, Victoria, British Columbia, Ca	nada		
³⁰ Cary Institute of Ecosystem Studies, Millbrook, New York, USA			
³¹ Department of Biology, Faculty of Science, University of British Columbia, Kelowna, B	ersity of British Columbia, Kelowna, British Columbia, Canada		
³² Department of Integrative Biology, University of Guelph, Guelph, Ontario, Canada			
³³ Department of Evolutionary Biology and Environmental Studies, University of Zurich,	Zurich, Switzerland		
³⁴ Department of Zoology and Physiology, University of Wyoming, Laramie, Wyoming, U	ISA		
³⁵ Department of Agriculture And Natural Resources Sciences, Biodiversity Research Cent	re, Namibia University of Science	and Technology, Windhoek, Namibia	
³⁶ Université de Toulouse, INRAE, CEFS, Castanet-Tolosan, France			
³⁷ LTSER ZA Pyrénées Garonne. Auzeville-Tolosane. France			

LISER ZAT Viences Galonne, Auzevine Tolosane, Trance

³⁸Department of Conservation and Research, Bavarian Forest National Park, Grafenau, Germany

³⁹Chair of Wildlife Ecology and Management, Albert Ludwigs University of Freiburg, Freiburg, Germany

⁴⁰Department of Forestry and Wildlife Management, Inland Norway University of Applied Sciences, Koppang, Norway

⁴¹Animal Behavior Graduate Group, University of California, Davis, California, USA

⁴²Norwegian Institute for Nature Research, Trondheim, Norway

⁴³Research Institute of Wildlife Ecology, University of Veterinary Medicine Vienna, Vienna, Austria

⁴⁴Department of Forestry and Renewable Forest Resources, Biotechnical Faculty, University of Ljubljana, Ljubljana, Slovenia

⁴⁵Black Rock Forest, Cornwall, New York, USA

⁴⁶Lamont-Doherty Earth Observatory, Columbia University, Palisades, New York, USA

⁴⁷Wildlife Ecology and Management, Manaaki Whenua – Landcare Research, Lincoln, New Zealand

⁴⁸Department of Anthropology, Stony Brook University, Stony Brook, New York, USA

⁴⁹International Union for Conservation of Nature (IUCN) Species Survival Commission (SSC) Tapir Specialist Group (TSG), Campo Grande, Mato Grosso do Sul, Brazil

⁵⁰Embrapa Pantanal, Corumbá, Mato Grosso do Sul, Brazil

⁵¹National Research Center for Carnivores Conservation, Chico Mendes Institute for the Conservation of Biodiversity, Atibaia, Brazil

⁵²Centre for Ecological and Evolutionary Synthesis, Department of Biosciences, University of Oslo, Oslo, Norway

⁵³Wildlife Research and Training Institute, Naivasha, Kenya

⁵⁴Norwegian Institute for Nature Research, Oslo, Norway

⁵⁵Wildlife Conservation Society, Mongolia Program, Ulaanbaatar, Mongolia

⁵⁶Latvian State Forest Research Institute "Silava", Salaspils, Latvia

⁵⁷Australian Wildlife Conservancy, Subiaco, Western Australia, Australia

⁵⁸Direction de la Recherche et de l'Appui Scientifique, Office Français de la Biodiversité, Birieux, France

⁵⁹Association for the Conservation of Biodiversity of Kazakhstan (ACBK), Nur-Sultan, Kazakhstan

⁶⁰Western Ecosystems Technology Inc., Laramie, Wyoming, USA

⁶¹Arctic Research Centre, Aarhus University, Aarhus, Denmark

⁶²Institute of Nature Conservation Polish Academy of Sciences, Krakow, Poland

⁶³Conservation Science and Wildlife Health, San Diego Zoo Wildlife Alliance, Escondido, California, USA

⁶⁴Department of Biology, Wilfrid Laurier University, Waterloo, Ontario, Canada

⁶⁵Greater Yellowstone Coalition, Bozeman, Montana, USA

⁶⁶Carnivore Ecology Laboratory, Forest and Wildlife Research Center, Mississippi State University, Mississippi State, Mississippi, USA

⁶⁷Alaska Department of Fish and Game, Kodiak, Alaska, USA

⁶⁸Institute for Terrestrial and Aquatic Wildlife Research, University of Veterinary Medicine Hannover Foundation, Hannover, Germany

⁶⁹Mara Elephant Project, Nairobi, Kenya

⁷⁰Department of Migration, Max Planck Institute of Animal Behavior, Radolfzell, Germany

⁷¹Centre for the Advanced Study of Collective Behaviour, University of Konstanz, Constance, Germany

⁷²Center for Integrated Spatial Research, Environmental Studies Department, University of California, Santa Cruz, California, USA

⁷³Tatra National Park, Zakopane, Poland

⁷⁴PBL Netherlands Environmental Assessment Agency, The Hague, The Netherlands

Correspondence

Maarten J. E. Broekman, Department of Environmental Science, Institute for Wetland and Water Research, Faculty of Science, Radboud University, P.O. Box 9010, NL-6500 GL, Nijmegen, The Netherlands.

Email: m.broekman@science.ru.nl

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Abstract

Aim: Macroecological studies that require habitat suitability data for many species often derive this information from expert opinion. However, expert-based information is inherently subjective and thus prone to errors. The increasing availability of GPS tracking data offers opportunities to evaluate and supplement expert-based information with detailed empirical evidence. Here, we compared expert-based habitat suitability information from the International Union for Conservation of Nature (IUCN) with habitat suitability information derived from GPS-tracking data of 1,498 individuals from 49 mammal species.

Location: Worldwide.

Time period: 1998-2021.

Major taxa studied: Forty-nine terrestrial mammal species.

Methods: Using GPS data, we estimated two measures of habitat suitability for each individual animal: proportional habitat use (proportion of GPS locations within a habitat type), and selection ratio (habitat use relative to its availability). For each individual we then evaluated whether the GPS-based habitat suitability measures were in agreement with the IUCN data. To that end, we calculated the probability that the ranking of empirical habitat suitability measures was in agreement with IUCN's classification into suitable, marginal and unsuitable habitat types.

Results: IUCN habitat suitability data were in accordance with the GPS data (> 95% probability of agreement) for 33 out of 49 species based on proportional habitat use estimates and for 25 out of 49 species based on selection ratios. In addition, 37 and 34 species had a > 50% probability of agreement based on proportional habitat use and selection ratios, respectively.

Main conclusions: We show how GPS-tracking data can be used to evaluate IUCN habitat suitability data. Our findings indicate that for the majority of species included in this study, it is appropriate to use IUCN habitat suitability data in macroecological studies. Furthermore, we show that GPS-tracking data can be used to identify and prioritize species and habitat types for re-evaluation of IUCN habitat suitability data.

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1 | INTRODUCTION

Habitat is broadly defined as the entirety of environmental conditions that enable a species to survive and reproduce (Doligez et al., 2008; Stamps, 2008). Hence, information on species' habitats is crucial for evaluating the effects of environmental change on species and for designing and planning conservation (e.g., Brooks et al., 2019; Manly et al., 2002; Rondinini et al., 2011). Large-scale and multi-species assessments typically employ expert-based habitat suitability information, for example from the International Union for Conservation of Nature (IUCN; e.g., Crooks et al., 2017; Di Marco et al., 2017; Powers & Jetz, 2019; Santini et al., 2019). The IUCN created an internationally recognized habitat classification scheme that distinguishes 54 terrestrial habitat types (https://www.iucnredlist.org/resources/ habitat-classification-scheme; IUCN, 2020). Knowledge from more than 1,700 experts was used to identify suitable habitat types (i.e., habitat type in which a species occurs regularly or frequently) for as many species as possible (IUCN, 2020; Schipper et al., 2008). For some species marginal habitat types are also identified (i.e., habitat type in which a species only occurs irregularly or infrequently, or in which only a small proportion of individuals are found). All other habitat types are considered unsuitable and do not constitute habitat for a species. As the IUCN habitat suitability data are based solely on expert judgment, without direct empirical data input, the information is inherently subjective. This may bias the results of studies using these expert-based data and misinform conservation decisions (Campbell et al., 2020; Johnson & Gillingham, 2004).

An alternative to expert opinion is to generate information on habitat suitability directly from empirical data. For example, species distribution models or resource selection functions can be used to identify habitat for a species (e.g., Dellinger et al., 2020; Peterson et al., 2018). However, using expert-based data on habitat suitability

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is often the only option, particularly for large-scale analyses, as empirical data are typically scarce and spatially restricted for many species (Amano & Sutherland, 2013; Boakes et al., 2010). This points towards a clear need to evaluate the accuracy of expert-based habitat suitability information where possible, for example by comparing it with empirical data. A few studies have evaluated expert-based habitat suitability data by calculating the number of occurrences of species in suitable and unsuitable habitat types, using point locality data from, for example, the Global Biodiversity Information Facility (e.g., Ficetola et al., 2015; Jung et al., 2020; Rondinini et al., 2011). These studies found that 55-94% of the point locations occurred in suitable habitat types. However, the number of locations for each species in point locality datasets is often small and does not include information on the identity of the individual. It is therefore unknown whether observations in unsuitable habitat types come from individuals that consistently use unsuitable habitat types (i.e., IUCN habitat suitability is misclassified), or individuals that only traverse unsuitable habitat types transiently to reach suitable habitat types (Bever et al., 2010).

Tracking individuals fitted with GPS tags generates a large number of individual-specific po location data, which enables distinguishing between habitat types that are used consistently or only transiently. The recent increasing use of GPS-tracking data therefore offers opportunities to better evaluate expert-based habitat suitability data. To our knowledge, no study has yet systematically evaluated expert-based data on habitat suitability with GPS-tracking data for a large number of species.

In this study, we use GPS-tracking data from 1,498 individuals of 49 terrestrial mammal species to estimate two empirical measures of habitat suitability and compare these measures with expert-based habitat suitability data recorded by the IUCN. One measure is proportional habitat use, which is defined as the proportion of locations in each habitat type (Lele et al., 2013; Manly et al., 2002). The other measure is the selection ratio, which relates proportional habitat use to its availability (i.e., the proportion of a given habitat type that is accessible for the species; Johnson, 1980; Lele et al., 2013; Manly et al., 2002). As each measure has its strengths and limitations, they provide complementary information. For example, according to the IUCN definition of suitable habitat types, we would expect that suitable habitat types are characterized by a greater proportional use. However, a low availability of a habitat type might lead to a low proportional use, even when the habitat type is in fact suitable. The selection ratio accounts for habitat availability and thus does not have this problem. A disadvantage of studying the selection ratio is that there can be a functional response of the selection ratio to the availability of a habitat type, which in some cases could lead to a decrease in the selection ratio when the availability of this habitat type increases (Aarts et al., 2013; Holbrook et al., 2019; Mysterud & Ims, 1998; van Beest et al., 2016). There are several reasons that could explain the presence of functional responses. One of these reasons is that species may use a habitat type only up to a level sufficient to meet its requirements (Aarts et al., 2013; Johnson, 1980), and once this requirement is fulfilled, a higher availability of that

habitat type does not necessarily lead to higher proportional use. As a consequence, suitable habitat types with a high proportional use may have a low selection ratio (but see Van Moorter et al., 2013). Because the proportional habitat use and selection ratio provide complementary information, we studied both metrics in our study.

We also explored whether the probability of agreement between our empirical measures and the IUCN data was related to the species' body mass, habitat specialization, or IUCN Red List category. Because larger-bodied species are generally better studied than smaller species (dos Santos et al., 2020), they may have more accurate IUCN habitat suitability data, and hence we expect a greater probability of agreement between empirical measures and IUCN data on habitat suitability for larger-bodied species. We further expect that habitat specialists (i.e., species that only occur in a few habitat types) would have a greater probability of agreement than habitat generalists (i.e., species that occur in many different habitat types), because it is easier for experts to identify suitable habitat types when species only occur in a few habitat types. Finally, the IUCN Red List category may impact the probability of agreement between the IUCN data and empirical measures. First of all, threatened species may have a greater research prioritization (Rodrigues et al., 2006), which could lead to a greater probability of agreement. Secondly, a threatened species might have been forced to move to suboptimal habitat, due to high population densities of species (McLoughlin et al., 2010; van Beest et al., 2014) or high anthropogenic disturbances (Kerley et al., 2020), which makes it more difficult for experts to identify the suitable habitat types, potentially leading to a lower probability of agreement. This could be the case for nonthreatened species, as these species are generally more abundant and thus often have higher population densities. However, this could also be the case for threatened species, which have more restricted distributions due to anthropogenic disturbances and might thus only occur in suboptimal habitat. Identifying the role of species characteristics might help to identify species not included in our analysis for which expert-based suitability of some habitat types is misclassified. Research on habitat suitability could then be directed towards these groups of species.

2 | METHODS

2.1 | GPS-tracking data and habitat data

We compiled GPS-tracking data of 1,498 individuals from 49 species of terrestrial mammals tracked between 1998 and 2021 (see Supporting Information Appendices S1–S3). Captures and handling of all these individuals were approved by the appropriate national or regional authorities (Supporting Information Appendix S4). The individuals come from 114 populations, here defined as groups of individuals that occur in the same geographic area, and occurred on all continents (except Antarctica) and most terrestrial biomes (Figure 1). The species included belong to 19 families, have body masses ranging from 0.5 to 4,000kg, include species from





FIGURE 1 Distribution of locations of (a) all 114 studied populations, (b) populations in North America, (c) Europe, (d) southern Africa, and (e) eastern Africa. The projections of the maps are Mollweide equal-area projections

each IUCN Red List category (Least Concern, Near Threatened, Vulnerable, Endangered, Critically Endangered), and include both habitat specialists and generalists (Faurby et al., 2018, 2020; IUCN, 2021; Wilman et al., 2014).

We obtained habitat data from a global map of terrestrial IUCN habitat types in 2015 at a ~100-m resolution (Jung et al., 2020). Estimates of habitat suitability could be affected by a temporal mismatch between the habitat map based on data from 2015, and

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the individual GPS data collected between 1998–2021. Therefore, we only included individuals for which we estimated that less than 5% of the available habitat types had changed between the year in which an individual was tracked and the year 2015 (see Supporting Information Appendix S5).

For each individual, we removed locations from the first 7 days of tracking after capture (on average 3.9% of the data), to avoid possible effects of capture and handling on the individual's locations (Bergvall et al., 2021; Gese et al., 2019; Mayer et al., 2019; Morellet et al., 2009). Furthermore, we removed locations that could be considered outliers based on unrealistic distances or speed between successive locations (on average 0.5% of the data, see Supporting Information Appendix S5 for a detailed description of the data cleaning). All individuals had at least 100 locations (mean: 6,436 locations), were tracked for a duration of at least 30 days (mean: 405 days) and had a median number of 11 locations per day after data cleaning (Supporting Information Appendix S3).

2.2 | Proportional habitat use and selection ratios

We used the GPS-tracking data and the IUCN habitat map to estimate the proportional habitat use and selection ratio of each habitat type for each individual. We calculated the proportional habitat use as the proportion of GPS locations of an individual within each habitat type (Lele et al., 2013) and the selection ratio as proportional habitat use divided by the relative amount of a habitat type available to that individual (Manly et al., 2002). We calculated the relative amount of a habitat type available to each individual as the proportional area of habitat types within an individual's 100% Minimum Convex Polygon (MCP) (Mohr, 1947), a commonly used approach to measure habitat selection at the home range scale (i.e., third-order selection sensu Johnson, 1980). When there were distinct clusters of locations for an individual (e.g., winter and summer ranges), we fitted a separate MCP for each cluster of locations, with clusters delineated based on a species-specific threshold distance (see Supporting Information Appendix S5). We applied this multi-MCP approach when the locations of an individual occurred in multiple clusters, because a single MCP would lead to large areas being considered as available that do not contain any locations. A single MCP would thus not accurately define available habitat for that individual.

2.3 | Probability of agreement between IUCN and GPS-tracking data

For each species, we used the estimated proportional habitat use and selection ratios to calculate a probability of agreement between the IUCN habitat suitability data and each habitat suitability measure. To calculate the probability of agreement between the IUCN habitat suitability data and the proportional habitat use estimates, we compared the proportional use of each suitable habitat type with the proportional use of every unsuitable and marginal habitat type for each individual. Next, we calculated the log₁₀-transformed ratios of these proportional habitat use estimates to measure to what degree a suitable habitat type is used, compared to an unsuitable or marginal habitat type, using:

$$R_{use} = \log_{10} \left(\frac{p_{suitable}^u}{p_{unsuitable/marginal}^u} \right)$$
(1)

in which $p_{suitable}^{u}$ indicates the proportional use of a suitable habitat type, $p_{unsuitable/marginal}^{u}$ indicates the proportional use of a marginal or unsuitable habitat type, and R_{use} indicates the log-transformed ratio of these proportional use estimates. For each individual, the number of ratios depends on the number of suitable and unsuitable habitat types. For example, if an individual has two suitable habitat types and three unsuitable habitat types, six comparisons can be made between a suitable and an unsuitable habitat and six ratios are thus calculated.

We only included habitat types with a proportional use larger than zero. Individuals of some species only occurred in suitable habitat types or only in marginal and unsuitable habitat types. For these individuals no comparison could be made between suitable and unsuitable habitat types. To retain these individuals in the analyses, we compared their proportional habitat use estimates with the theoretical minimum proportional habitat use of the individual: the inverse of the individual's number of GPS locations. This proportion is equal to the proportional habitat use when there is only one location in a habitat type.

We combined the ratios of all individuals of the same species and fitted a linear mixed-effect model:

$$R_{use}^{\sim}$$
 suitability + (1 | population/individual) (2)

In this mo del, *suitability* indicates whether the ratio of a suitable and an unsuitable habitat type was calculated, or the ratio of a suitable and marginal habitat type. Nested random effects of population and individual were included to account for repeat sampling of the same population and individual. The intercept of this model indicates the species-specific log₁₀-transformed ratio of the proportional use of suitable compared to marginal habitat types, and the coefficient for *suitability* indicates the difference between this ratio and the species-specific log₁₀-transformed ratio of the proportional use of suitable compared to unsuitable habitat types.

We applied parametric bootstrapping using 1,000 simulations to obtain 1,000 estimates of the intercept and *suitability* coefficient of this model. These bootstrap results were used to calculate the probability of agreement between the IUCN habitat suitability data and the proportional habitat use estimates. This probability of agreement is the proportion of bootstrap simulations in which the model coefficients indicate $p_{suitable}^u > p_{marginal}^u > p_{unsuitable}^u$. The above condition is satisfied when the intercept estimate is larger than zero, indicating a higher proportional habitat use for suitable than for marginal habitat types. In addition, the *suitability* coefficient needs to be

larger than zero, indicating a larger difference in proportional habitat use between suitable and unsuitable habitat types than between suitable and marginal habitat types. For 17 species that did not occur in any marginal habitat type, we fitted an intercept-only model and calculated the probability of agreement as the proportion of bootstrap simulations in which the intercept is larger than zero, indicating $p_{uisuitable}^u > p_{uisuitable}^u$.

In an analogous way, we calculated the probability of agreement between the IUCN habitat suitability data and the selection ratio estimates by calculating the \log_{10} -transformed ratios (R_{SR}) of the selection ratios for a suitable habitat type ($SR_{suitable}$) and a marginal or unsuitable habitat type ($SR_{unsuitable/marginal}$) for each individual:

$$R_{SR} = \log_{10} \left(\frac{SR_{suitable}}{SR_{unsuitable/marginal}} \right)$$
(3)

To avoid selection ratios of zero, for which the log₁₀ is undefined, we substituted the zero selection ratios with the lowest possible selection ratios: the theoretical minimum proportional habitat use of the individual (the inverse of the individual's number of GPS locations) divided by the relative amount of available area of the habitat type. Because of this correction for zeros, we excluded habitat types for which the relative amount of available area was lower than the theoretical minimum proportional habitat use of the individual. If these habitat types were included, they would always have a selection ratio larger than one, even though they are not used. Furthermore, we also excluded these habitat types because selection ratios are unreliable for habitat types with a low availability. For habitat types with a low availability, the absence of any GPS location in this habitat type might not indicate habitat avoidance, but could just be the result of the scarcity of this habitat type. We note that some of the habitat types included in the selection ratio analyses have a proportional habitat use of zero; therefore, the number of included habitat types is larger than in the analyses of the proportional habitat use. For individuals where only suitable or only unsuitable or marginal habitat types were retained, we compared the selection ratios with the theoretical minimum selection ratio of these habitat types: the inverse of the individual's number of GPS locations, divided by the relative amount of available area of the habitat type [see Supporting Information Appendix S6 for a detailed calculation of the probability of agreement for the Impala (Aepyceros melampus)].

We note that a high probability of agreement (e.g., 100%) for a species does not necessarily imply that all suitable habitat types have a higher proportional habitat use or selection ratio than unsuitable habitat types. It is possible that the suitability of one habitat type is consistently misclassified, but because the selection ratio or proportional use of all the other habitat types is higher for suitable than unsuitable habitat types, the general probability of agreement for this species can still be high. To study whether the suitability of a habitat type might be misclassified, we also calculated a habitat type-specific probability of agreement for a few habitat types and species for which we suspect the suitability might be misclassified by the IUCN. These habitat types were found by looking for (a) suitable habitat types that were used less or that were selected for less than unsuitable habitat types in multiple individuals, or (b) unsuitable habitat types that were used or selected more than suitable habitat types. To calculate a habitat-type-specific probability of agreement, we performed the same analyses as described above, but only used the \log_{10} -transformed ratios that included the specific habitat type.

2.4 | Effects of species characteristics on probability of agreement

We evaluated whether the probability of agreement between our habitat suitability measures and the IUCN data were related to the species' body mass, habitat specialization and the IUCN Red List category. We derived body mass values from the literature or used the mean weight of the tracked individuals (Supporting Information Appendices S1 and S2). To retrieve information on habitat specialization, researchers who collected the data classified the species they tracked as either habitat specialists or generalists. With regard to the IUCN Red List category, we classified species as non-threatened (Least Concern, Near Threatened) or threatened (Vulnerable, Endangered, Critically Endangered) (IUCN, 2021). We then fitted beta regression models relating the probability of agreement based on proportional use or selection ratio to the three species characteristics. Because the response variable in beta regression models is bounded between zero and one, we transformed the probability of agreement with a Smithson-Verkuilen transformation to avoid zeros and ones (Smithson & Verkuilen, 2006):

$$y' = (y \cdot (n-1) + 0.5) / n \tag{4}$$

in which n indicates the sample size, that is, number of species.

To account for differences in the number of tracked individuals $(n_{individuals})$, number of populations $(n_{populations})$ and median tracking duration (t_{median}) between species, which could constrain the effects of species characteristics, we also included these factors as predictor variables in the models:

Probability of agreement%
$$\log_{10}(Body mass)$$

+ habitat specialization + IUCN category (5)
+ $n_{individuals} + n_{populations} + t_{median}$

We used a model selection approach based on the Akaike's information criterion corrected for small sample size (AIC_c) to identify which combination of the species' characteristics best explained the probability of agreement between our habitat suitability measures and the IUCN data (Burnham & Anderson, 2002). We included models containing all possible combinations of additive effects of the species' characteristics. The variance inflation factors of all predictor variables were lower than two, indicating no substantial collinearity problems (Zuur et al., 2010). Global Ecology

FIGURE 2 Probability of agreement between the International Union for Conservation of Nature (IUCN) habitat suitability data and the proportional habitat use estimates for each species. The dashed line indicates the average probability of agreement across all species. The numbers in parentheses indicate the number of populations and number of individuals for the species, respectively. * indicates species for which all individuals only occurred in unsuitable habitat types. ** indicates species for which all individuals only occurred in suitable habitat types

2.5 | Sensitivity analyses

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To assess whether our results were robust to methodological choices made in our analyses, we re-ran our analyses in three different ways. First, we repeated our analyses using longer minimum tracking durations (60–600 days) and greater minimum number of observations (100–20,000). Secondly, we also calculated the availability of each habitat type as the proportional area within an individual-specific buffer-distance around each GPS location, that is, calculating habitat availability using the buffer-approach (Montgomery et al., 2018). Finally, we repeated all analyses using GPS-tracking data subsampled to approximately constant time intervals for each individual. (See Supporting Information Appendix S5 for more information on these different methodological approaches.)

3 | RESULTS

The average probability of agreement across all species between habitat suitability data derived from the IUCN and empirical proportional use estimates was 76% (\pm 39% SD; Figure 2). Similarly, the average probability of agreement between IUCN habitat suitability data and selection ratio estimates across all species was 69% (±40% SD; Figure 3). Thirty-three (67%) and 25 (51%) species had a probability of agreement greater than 95%, based on the proportional habitat use and selection ratio, respectively. These numbers increased to 37 (76%) and 34 (69%) species when we used a probability of agreement greater than 50% as a threshold. There were also six species for the proportional habitat use and five species for the selection ratio with a very low probability of agreement (< 5%). In addition, for several species we found one or more individuals for which a suitable habitat type had a lower selection ratio or proportional use than an unsuitable or marginal habitat type (see Broekman et al., 2022). For example, for the gray wolf (Canis lupus) the suitable habitat type 'Pastureland' was often less used and less selected than unsuitable habitat types, such as arable land. The probability of agreement for this habitat type for C. lupus was therefore only 0.3% based on the proportional habitat use and 9.9% based on the selection ratio. Similarly, for several species we also found one or more individuals for which an unsuitable or marginal habitat was used and selected more than suitable habitat types. For example, for the jaguar (Panthera onca), the most used and selected habitat type in all four individuals was 'Forest Subtropical-tropical dry', which is classified as marginal by the IUCN. The probability of agreement for this habitat type for P. onca was zero based on both the proportional habitat use and selection ratio.

The most parsimonious model explaining the probability of agreement between the empirical habitat suitability measures and the IUCN habitat suitability data was an intercept-only model for both the proportional use and selection ratio (Supporting Information Appendices S7 and S8). Two models within 2 AIC_c units from the most parsimonious model for proportional habitat use contained the effects of the number of individuals and number of populations, respectively. For the selection ratio, three models within 2 AIC_c units from the most parsimonious model contained the effects of body mass, number of populations, and median tracking duration, respectively.

All the results were robust to methodological changes, including increments in minimum tracking duration and minimum number of observations per individual, data subsampling, and using the buffer-approach to estimate habitat availability (Supporting Information Appendix S9).

4 | DISCUSSION

4.1 | Interpretation

In this study, we showed how GPS-tracking data can be used to evaluate expert-based habitat suitability data. Our findings indicate that for the majority of species included in this study, it is appropriate to use IUCN habitat suitability data in macroecological studies. Nevertheless, caution should still be taken when using IUCN habitat suitability data, as for some species there might be a few misclassified habitat types. For example, for the vervet monkey (Chlorocebus pygerythrus) and the lowland tapir (Tapirus terrestris), the probability of agreement between IUCN habitat suitability data and our empirical habitat suitability measures was low, which may reflect misclassification of suitability for some habitat types. In addition, for some species, including several with a high probability of agreement, there were unsuitable or marginal habitat types with a greater proportional use or selection ratio than one or more suitable habitat types (see Broekman et al., 2022). Some other species had suitable habitat types with a lower proportional use or selection ratio than one or more unsuitable or marginal habitat types (see Broekman et al., 2022). These results also suggest that the suitability of these habitat types might be misclassified. We have not provided a list of habitat types that are potentially misclassified by the IUCN to avoid the use of arbitrary thresholds for identifying suitable and unsuitable habitat types. In fact, there could also be other reasons for a mismatch between the IUCN habitat suitability data and our habitat suitability measures (see below); our results might serve as a starting point to identify species and habitat types for which further research is necessary to evaluate suitability.



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FIGURE 3 Probability of agreement between the International Union for Conservation of Nature (IUCN) habitat suitability data and the selection ratio estimates for each species. The dashed line indicates the average probability of agreement across all species. The numbers in parentheses indicate the number of populations and number of individuals for the species, respectively. * indicates species for which all individuals only have unsuitable habitat types available. ** indicates species for which all individuals only have suitable habitat types available.

Our results also suggest that body mass, habitat specialization and Red List category are not informative characteristics for identifying species for which the IUCN habitat suitability data agree with inference from GPS-tracking data. These species characteristics were not included in the most parsimonious model predicting the probability of agreement between IUCN habitat suitability data and our empirical habitat suitability measures. Whether the IUCN classified habitat suitability for a species correctly most likely depends on various species-specific factors. For example, while larger-bodied species are, in general, better studied than smaller-bodied species, some large-bodied species may be highly elusive, and their habitat use may have been poorly known before the advent of GPS telemetry (e.g., jaguar, puma). Furthermore, smaller-bodied species often range over smaller areas, exposing them to fewer habitat types, given the spatial resolution of the IUCN habitat map (Jung et al., 2020), which may reduce the possibilities of misclassifying habitat suitability. Moreover, species living in large groups in open habitats (e.g., ungulates in grasslands) might have been studied more intensively than solitary, forest-dependent species. Because of the complexity of factors determining the accuracy of IUCN habitat suitability data, it is difficult for the moment to draw conclusions. In addition, inferences from the analyses are limited by our relatively small sample of species given the total number of terrestrial mammal species classified by the IUCN and several other limitations described in the next section. A larger number of species and more species-specific characteristics (e.g., home range size, diet, elusiveness of the species) should thus be investigated to identify characteristics that may explain for which species the IUCN habitat suitability data are likely to show a low probability of agreement with results from GPStracking data.

4.2 | Limitations

There are several limitations in the habitat maps, IUCN habitat suitability data, and GPS-tracking data that may explain why not all species have corresponding results on habitat suitability. First of all, the IUCN habitat types are sometimes difficult to match to vegetation or habitat types used in the field. This makes it difficult for experts to determine the suitability of a habitat type and might also have led to misassignments of habitat types to grid cells of the IUCN habitat map (Jung et al., 2020). For example, the IUCN defines 'plantations' as plantings of trees and shrubs (IUCN, 2020), which can be, for example, coffee plantations, but also mature forest plantations. For the Eurasian lynx (*Lynx lynx*) 'plantations' are classified as an unsuitable habitat type by the IUCN, which might be correct for coffee plantations. However, the lynx in our dataset are frequently located in 'plantations', which covers large parts of Europe (especially southern Sweden) and are more likely mature forest plantations, that is, commercially operated forests subject to cutting and replanting. From a lynx point of view, the suitability of mature forest plantations might be similar to, for example, temperate or boreal forest, which are suitable habitat types for the lynx, according to the IUCN. Most 'plantations' in Europe might therefore be better represented by one of the forest habitat types. Despite these possible misassignments, we still used the IUCN map as it currently contains the best available data for global analyses.

Second, individuals have intrinsic differences in habitat suitability (e.g., Leclerc et al., 2016; Lesmerises & St-Laurent, 2017; Montgomery et al., 2018), and the suitability of habitat types likely varies spatially and temporally for most species. Mismatches between the IUCN and GPS-tracking data might therefore arise, because for most species we only studied a few individuals that occurred on a small subset of the entire range or that were only tracked for a short time period. For example, the Eurasian lynx data in this study came from multiple studies across the European part of their distribution, but none from Asia. Estimated habitat suitability measures for these individuals might not be representative of the species as a whole. Regional differences in habitat suitability could arise due to other abiotic and biotic factors that we did not account for, such as weather, distance to water, human disturbances, or prevalence of prey/predator/competitor species (e.g., Attias et al., 2018; Dellinger et al., 2020; Jones et al., 2019; Mayer et al., 2019; Rivrud et al., 2019; Roever et al., 2012). For example, temperate grassland is listed as suitable habitat for the pronghorn (Antilocapra americana) by the IUCN, but this species tends to avoid roads (Jones et al., 2019). This habitat type might thus be unsuitable when it is located close to roads, but suitable otherwise. Temporal variability in habitat suitability could arise due to seasonal differences, such as between summer and winter (e.g., Rivrud et al., 2019) or between wet and dry seasons (e.g., Roever et al., 2012). Nevertheless, our results did not change when using longer minimum tracking durations as a threshold to select individuals. This indicates that our results were not influenced by individuals that were only tracked during a specific season.

Third, mismatches in habitat suitability might be attributed to taxonomic classifications that diverge from more recent phylogenetic assessments. For example, four species of giraffe have been classified based on phylogenetic analyses (Coimbra et al., 2021), whereas the IUCN only identifies a single species (*Giraffa camelopar-dalis*). We followed the IUCN taxonomy, as habitat suitability data are provided for this single species. However, there might be differences in habitat suitability among these four species. The taxonomic



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classifications of giraffe species are relatively new, and we are only now starting to understand the ecological implications of these new taxonomic classifications in terms of habitat use and conservation (O'Connor et al., 2019).

Fourth, mismatches between the IUCN habitat suitability data and our empirical measures of habitat suitability could occur because proportional habitat use and selection ratio do not always correctly indicate habitat suitability. For example, variability in habitat availability can lead to changes in selection ratios, that is, functional responses, even when the suitability of a habitat type does not change (Aarts et al., 2013; Holbrook et al., 2019; Mysterud & Ims, 1998; van Beest et al., 2016). Furthermore, the proportional use and selection ratio of a given habitat type depends on the other habitat types available (Johnson, 1980; Lele et al., 2013). More precisely, the selection ratio and proportional use of a habitat type will be greater when all other available habitat types are unsuitable, compared to when some of the other available habitat types are also suitable (see also Bever et al., 2010; Van Moorter et al., 2013). Moreover, a species might be forced to move to less suitable habitat types due to, for example, high population densities of species (McLoughlin et al., 2010; van Beest et al., 2014) or anthropogenic disturbances (Kerley et al., 2020), which might lead to an overestimation of the suitability for these habitat types. To limit the impact of individuals occurring only in suboptimal habitat on the calculations of the probability of agreement, we included individuals from as many different populations as possible, increasing the likelihood of including individuals that were not forced into suboptimal habitat. For example, for the giraffe, we included data from 21 different populations spread across its range. However, for some other species we had access to data from only one population. Nevertheless, by focussing on the overall rather than the species-specific results, we tried to limit the bias that might be generated by individuals that are forced to move to less suitable habitat types.

Finally, we note that our sample size of 49 species is relatively small compared to the total number of 5,480 mammal species for which the IUCN has habitat suitability data (IUCN, 2021). Furthermore, we only focused on mammal species, so our results are not representative of all species groups (e.g., birds, reptiles and amphibians). Our data also include some well-studied species, for which habitat suitability might be better known than for less-studied species, and it could thus be argued that our results represent a best case scenario. However, our data include also less-studied species (e.g., Beatragus hunteri, Priodontes maximus). In addition, for the red fox (Vulpes vulpes), a highly studied species, we found a probability of agreement of only 7% for the proportional habitat use and 6% for the selection ratio, which is well below the average. Thus, our results do not seem to be biased by unrepresentative sampling of mammal species. Given the difficulty of collecting GPS-tracking data, our dataset includes a very large number of individuals and species and is currently the best available. Nevertheless, the lack of data for the majority of mammal species highlights the need for more GPS-tracking studies, especially for species that are still poorly studied.

4.3 | Recommendations

Although our results show that for the majority of studied species it is appropriate to use IUCN habitat suitability data in macroecological studies, there might still be several habitat types for which suitability is misclassified by the IUCN. Reducing potential misclassifications of habitat suitability by the IUCN would benefit all studies that make use of these expert judgments. For example, when allocating new protected areas to prevent local extinction of species at risk, it is vital to have reliable information on species' habitat suitability, otherwise we risk making expensive but ineffective conservation decisions. Habitat suitability data can also be used to calculate the area of habitat for each species and subsequently assess its Red List status (e.g., Santini et al., 2019). More accurate habitat suitability information will therefore contribute to more accurate assessments of progress towards international targets related to nature conservation, including the Aichi targets and the targets within the post-2020 global biodiversity framework (SCBD, 2011, 2020).

We showed that GPS-tracking data can be used to identify species and habitat types for which re-evaluations of the suitability information might be necessary. In addition, GPS-tracking data can be used to account for several factors that constrain the habitat suitability assessment. For example, because the GPS data provide individual-specific information on habitat suitability, they can be used to derive separate habitat suitability data for different regions. Similarly, GPS-tracking data can be used to derive season-specific habitat suitability data. In this study, we did not calculate different habitat suitability metrics for, for example, different seasons, or regions, even though this might be appropriate for some species, to optimize comparability with the IUCN data. Nevertheless, future studies would benefit from using regional- and season-specific habitat suitability data, as this would lead to more accurate estimates of, for example, the area of habitat for a species within (potential) protected areas (e.g., Di Marco et al., 2017).

GPS-tracking data can also be used to distinguish more habitat suitability categories that may better reflect a species' habitat suitability. Currently, the IUCN only distinguishes suitable, marginal and unsuitable habitat types, and for many species marginal habitat types are not reported. However, using individual-specific GPStracking data we could rank habitat types for each individual from least to most used or selected. We could then classify habitat types as, for example, 'highly suitable' when they are always selected over all other habitat types, whereas we could classify habitat types as 'partly suitable' when they are only selected over unsuitable habitat types, but not other suitable habitat types. In this way, GPS-tracking information might also be used to accommodate inter-individual differences in habitat suitability, for example, 'highly suitable' habitat types are suitable for all individuals, and 'partly suitable' habitat types are only suitable for a subset of individuals, as a consequence of, for example, inherent individual differences or regional differences in habitat suitability. This will only be feasible when tracking data are available from a range of representative regions and seasons, as discussed in the previous paragraph. However, with the

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increasing availability of GPS-tracking data, these limitations may soon be overcome for several species. Because of these opportunities, we recommend using GPS-tracking data when updating IUCN habitat suitability data.

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CONFLICT OF INTEREST

The authors have no conflict of interest to declare.

DATA AVAILABILITY STATEMENT

Data is published on DANS (https://doi.org/10.17026/dans-xq3-6muu). These data contain the following information for each individual of each species: individual ID, population ID, species ID, proportional use of each habitat type, available proportions of each habitat type, selection ratios of each habitat type. Furthermore, the GPS-tracking data for some individuals are published on Movebank or are published in Eurommammals (euromammals.org) (see Appendix S4). In Supporting Information Appendix S4 we indicate the Movebank details for the individuals that have their GPS-tracking data published on Movebank. Data from Euromammals can be retrieved by logging into their website or via a contact form.

ETHICS APPROVAL

Captures and handling of all individuals were approved by the appropriate authorities (see Supporting Information Appendix S3).

ORCID

Maarten J. E. Broekman [®] https://orcid.org/0000-0002-0565-5220 Jelle P. Hilbers [®] https://orcid.org/0000-0002-9401-589X Mark A. J. Huijbregts [®] https://orcid.org/0000-0002-7037-680X Henrik Andrén [®] https://orcid.org/0000-0002-5616-2426 Nina Attias [®] https://orcid.org/0000-0002-9472-6763 Hattie L. A. Bartlam-Brooks [®] https://orcid.org/0000-0002-1507-1712 Floris M. van Beest [®] https://orcid.org/0000-0002-5701-4927 Laura Bidner [®] https://orcid.org/0000-0002-4198-6609 Niels Blaum [©] https://orcid.org/0000-0001-6807-5162 Randall B. Boone [®] https://orcid.org/0000-0003-3362-2976 Mark S. Boyce [®] https://orcid.org/0000-0001-5811-325X Francesca Cagnacci [®] https://orcid.org/0000-0002-4954-9980 Rok Černe [®] https://orcid.org/0000-0003-4111-4066 Simon Chamaillé-Jammes [®] https://orcid.

org/0000-0003-0505-6620

Nandintsetseg Dejid D https://orcid.org/0000-0002-5376-3808 Jasja Dekker () https://orcid.org/0000-0001-6453-4480 Arnaud L. J. Desbiez (b) https://orcid.org/0000-0001-5968-6025 Samuel L. Díaz-Muñoz D https://orcid.org/0000-0003-2967-9806 Claudia Fichtel D https://orcid.org/0000-0002-8346-2168 Christina Fischer b https://orcid.org/0000-0001-7784-1105 Jason T. Fisher () https://orcid.org/0000-0002-9020-6509 Benedikt Gehr b https://orcid.org/0000-0002-1044-9296 A. J. Mark Hewison ^(b) https://orcid.org/0000-0002-2276-4154 Robert Hering b https://orcid.org/0000-0001-6098-0387 Marco Heurich () https://orcid.org/0000-0003-0051-2930 Florian Jeltsch 🕩 https://orcid.org/0000-0002-4670-6469 Petra Kaczensky D https://orcid.org/0000-0001-5428-1176 Miha Krofel https://orcid.org/0000-0002-2010-5219 Scott LaPoint b https://orcid.org/0000-0002-5499-6777 John D. C. Linnell 🕩 https://orcid.org/0000-0002-8370-5633 A. Catherine Markham b https://orcid.org/0000-0001-9756-4870 Jenny Mattisson D https://orcid.org/0000-0001-6032-5307 Emilia Patricia Medici 🔟 https://orcid.org/0000-0003-1944-9249 Guilherme de Miranda Mourão 🕩 https://orcid.

org/0000-0002-8300-4191

Bram Van Moorter 💿 https://orcid.org/0000-0002-3196-1993 Ronaldo G. Morato D https://orcid.org/0000-0002-8304-9779 Nicolas Morellet b https://orcid.org/0000-0002-4274-7044 Atle Mysterud https://orcid.org/0000-0001-8993-7382 John Odden 💿 https://orcid.org/0000-0002-6275-8648 Kirk A. Olson (D) https://orcid.org/0000-0002-2543-3414 Aivars Ornicāns b https://orcid.org/0000-0002-1478-3080 Jens Persson () https://orcid.org/0000-0003-1405-7561 Tyler Petroelje D https://orcid.org/0000-0003-4957-6732 Christer Moe Rolandsen D https://orcid.org/0000-0002-5628-0385 David Roshier D https://orcid.org/0000-0002-8151-8447 Daniel I. Rubenstein Dhttps://orcid.org/0000-0001-9049-5219 Sonia Saïd https://orcid.org/0000-0003-1485-8297 Hall Sawyer 🕩 https://orcid.org/0000-0002-3789-7558 Niels Martin Schmidt 🕩 https://orcid.org/0000-0002-4166-6218 Nuria Selva D https://orcid.org/0000-0003-3389-201X Agnieszka Sergiel D https://orcid.org/0000-0002-3455-4218 Jared Stabach 💿 https://orcid.org/0000-0002-7896-1237 Jenna Stacy-Dawes b https://orcid.org/0000-0001-5828-6476 Jonas Stiegler D https://orcid.org/0000-0002-3860-8877 Nathan J. Svoboda 🗅 https://orcid.org/0000-0002-1939-0569 Ulrich Voigt https://orcid.org/0000-0002-9505-4612 Tomasz Zwijacz-Kozica 🕩 https://orcid.org/0000-0002-7488-975X Aafke M. Schipper b https://orcid.org/0000-0002-5667-0893 Marlee A. Tucker D https://orcid.org/0000-0001-7535-3431

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Maarten Broekman is a PhD candidate at the Radboud University in Nijmegen, the Netherlands. His research focuses on quantifying the effects of habitat conversion on biodiversity. Specifically, he studies terrestrial mammals at a global scale.

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