

Fear of the dark? Contrasting impacts of humans vs lynx on diel activity of roe deer across Europe

Nadège Bonnot, Ophélie Couriot, A. Berger, F. Cagnacci, S. Ciuti, J. de

Groeve, B. Gehr, M. Heurich, P. Kjellander, M. Kröschel, et al.

▶ To cite this version:

Nadège Bonnot, Ophélie Couriot, A. Berger, F. Cagnacci, S. Ciuti, et al.. Fear of the dark? Contrasting impacts of humans vs lynx on diel activity of roe deer across Europe. Journal of Animal Ecology, 2020, 89 (1), pp.132-145. 10.1111/1365-2656.13161 . hal-02609832

HAL Id: hal-02609832 https://hal.inrae.fr/hal-02609832

Submitted on 28 Jan 2022

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution - NonCommercial - NoDerivatives 4.0 International License

Author-produced version of the article published in Journal of animal ecology, Volume 89, n°1, 2020, Pages 132-145 The original publication is available at https://besjournals.onlinelibrary.wiley.com/doi/10.1111/1365-2656.13161, doi : 10.1111/1365-2656.13161 ©. This manuscript version is made available under the CC-BY-NC-ND 4.0 license http://creativecommons.org/licenses/by-nc-nd/4.0/

Fear of the dark? 1 Contrasting impacts of humans vs lynx on diel activity 2 of roe deer across Europe 3 4 Nadège C. BONNOT^{*a, b}, Ophélie COURIOT^{*c}, Anne BERGER^d, Francesca CAGNACCI^e, 5 Simone CIUTI ^f, Johannes DE GROEVE ^{e, g}, Benedikt GEHR ^h, Marco HEURICH ^{i, j}, 6 Petter KJELLANDER^a, Max KRÖSCHEL^{k,1}, Nicolas MORELLET^c, Leif SÖNNICHSEN^{m, d} 7 8 & A. J. Mark HEWISON^c 9 * the two first authors contributed equally to this work 10 11 12 Addresses: 13 ^a Grimsö Wildlife Research Station, Department of Ecology, Swedish University of Agricultural Sciences, 14 Riddarhyttan, Sweden 15 ^b UR EFNO, Irstea, Domaine des Barres, Nogent-sur-Vernisson, France 16 ^c CEFS, Université de Toulouse, INRA, Castanet-Tolosan, France 17 ^d Leibniz Institute for Zoo and Wildlife Research, Berlin, Germany 18 ^e Department of Biodiversity and Molecular Ecology, Research and Innovation Centre, Fondazione Edmund Mach, San Michele all'Adige, Italy 19 20 ^f Laboratory of Wildlife Ecology and Behaviour, School of Biology and Environmental Science, University College 21 Dublin, Ireland 22 ^g Department of Geography, Ghent University, Gent, Belgium 23 ^h Department of Evolutionary Biology and Environmental Studies, University of Zurich, Zurich, Switzerland

25	^j Bavarian Forest National Park, Department of Visitor Management and National Park Monitoring,
26	Germany
27	^k Division of Wildlife Ecology, Forest Research Institute of Baden-Württemberg, Freiburg, Germany
28	¹ Chair of Wildlife Ecology and Wildlife Management, University of Freiburg, Freiburg, Germany

ⁱ Faculty of Environment and Natural Resources, University of Freiburg, Freiburg, Germany

29 ^m Mammal Research Institute, Polish Academy of Sciences, Białowieża, Poland

30

24

31

32 **Contact information:**

- *Nadège Bonnot: bonnot.nadege@gmail.com (corresponding author)
- 34 *Ophélie Couriot: o.couriot@gmail.com
- 35 Anne Berger: berger@izw-berlin.de
- 36 Francesca Cagnacci: francesca.cagnacci@fmach.it
- 37 Simone Ciuti: simone.ciuti@ucd.ie
- 38 Johannes De Groeve: degroevejohannes@gmail.com
- 39 Benedikt Gehr: benedikt.gehr@ieu.uzh.ch
- 40 Marco Heurich: marco.heurich@npv-bw.bayern.de
- 41 Petter Kjellander: petter.kjellander@slu.se
- 42 Max Kröschel: max.kroeschel@wildlife.uni-freiburg.de
- 43 Nicolas Morellet: nicolas.morellet@inra.fr
- 44 Leif Sönnichsen: leif.soennichsen@gmx.de
- 45 A.J. Mark Hewison: mark.hewison@inra.fr

46

Grafenau,

47 Abstract

Humans, as super predators, can have strong effects on wildlife behaviour, including profound
 modifications of diel activity patterns. Subsequent to the return of large carnivores to human modified ecosystems, many prey species have adjusted their *spatial* behaviour to the
 contrasting landscapes of fear generated by both their natural predators and anthropogenic
 pressures. The effects of predation risk on *temporal* shifts in diel activity of prey, however,
 remain largely unexplored in human-dominated landscapes.

We investigated the influence of the density of lynx (*Lynx lynx*), a nocturnal predator, on the
diel activity patterns of their main prey, the roe deer (*Capreolus capreolus*), across a gradient
of human disturbance and hunting at the European scale.

3. Based on 11 million activity records from 431 individually GPS-monitored roe deer in 12
populations within the EURODEER network (http://eurodeer.org), we investigated how lynx
predation risk in combination with both lethal and non-lethal human activities affected deer
diurnality.

61 4. We demonstrated marked plasticity in roe deer diel activity patterns in response to spatiotemporal variations in risk, mostly due to human activities. In particular, roe deer decreased 62 their level of diurnality by a factor of 1.37 when the background level of general human 63 disturbance was high. Hunting exacerbated this effect, as during the hunting season deer 64 switched most of their activity to nighttime and, to a lesser extent, to dawn, although this 65 66 pattern varied noticeably in relation to lynx density. Indeed, in the presence of lynx, their main 67 natural predator, roe deer were relatively more diurnal. Overall, our results revealed a strong influence of human activities and the presence of lynx on diel shifts in roe deer activity. 68

69	5.	5. In the context of the recovery of large carnivores across Europe, we provide important insights				
70		about the effects of predators on the behavioural responses of their prey in human-dominated				
71		ecosystems. Modifications in the temporal partitioning of ungulate activity as a response to				
72		human activities may facilitate human-wildlife coexistence, but likely also have knock-on				
73		effects for predator-prey interactions, with cascading effects on ecosystem functioning.				
74						
75		Key-words: Accelerometers; Crepuscularity; Diurnality; Human footprint; Hunting;				
76		Landscape of fear; Nocturnality; Predator-prey interaction; Temporal partitioning;				
77		Urbanization				
78						
79						

80 Second abstract in native language (French): Résumé

1. Les humains, en tant que "super-prédateurs", peuvent avoir des effets importants sur le 81 comportement de la faune sauvage, y compris des modifications profondes de leurs rythmes 82 circadiens d'activité. A la suite du retour des grands carnivores dans les écosystèmes 83 anthropisés, de nombreuses espèces proies ont ajusté leur comportement spatial à ces paysages 84 de la peur contrastés, générés à la fois par les pressions liées aux risques anthropiques et à la 85 présence de leurs prédateurs naturels. Les effets du risque de prédation sur les modifications 86 temporelles des rythmes circadiens d'activité des proies restent cependant largement inconnus 87 dans les écosystèmes dominés par l'homme. 88

2. Ici, nous avons étudié l'influence de la densité de lynx (*Lynx lynx*), un prédateur nocturne, sur
les rythmes circadiens d'activité de leur proie principale, le chevreuil (*Capreolus capreolus*),
à travers un gradient de pressions anthropiques à l'échelle Européenne.

3. Sur la base de plus de 11 million de données d'activité issues de 431 suivis individuels de chevreuils équipés de colliers GPS provenant de 12 populations au sein du réseau
EURODEER (<u>http://eurodeer.org</u>), nous avons analysé comment le risque de prédation par le lynx, associé aux risques létaux et non-létaux des activités humaines, influence la diurnalité des chevreuils.

97 4. Nous avons démontré une forte plasticité des rythmes circadiens d'activité des chevreuils en
98 réponse aux variations spatio-temporelles du risque, et notamment face aux activités
99 humaines. Plus particulièrement, les chevreuils diminuent leur degré de diurnalité d'un facteur
100 de 1.37 lorsque le dérangement humain est important. La chasse accentue cet effet, puisque
101 durant la saison de chasse les chevreuils basculent la plupart de leur activité de nuit, et dans

une moindre mesure, durant l'aube également, bien que ce patron soit essentiellement variable
en fonction de la densité de lynx. En effet, en présence de lynx, leur principal prédateur, les
chevreuils sont relativement plus diurnes. Globalement, nos résultats révèlent une forte
influence des activités humaines et de la présence de lynx sur l'ajustement des rythmes
circadiens d'activité des chevreuils.

5. Dans le contexte du retour des grands carnivores en Europe, notre étude apporte de nouvelles
connaissances sur les effets des prédateurs sur la réponse comportementale de leur proie dans
des écosystèmes anthropisés. La modification de la répartition temporelle de l'activité des
ongulés en réponse aux activités humaines pourrait être un facteur facilitant la coexistence
homme-faune sauvage, avec toutefois des conséquences autres sur les interactions prédateursproies et leurs effets en cascade sur le fonctionnement des écosystèmes.

113 6.

Mots-clés : Accéléromètres; Crépuscularité; Diurnalité; Empreinte humaine; Chasse; Paysage
de la peur; Nocturnalité; Interaction prédateurs-proies; Répartition temporelle de l'activité;
Urbanisation

118 Introduction

Global changes linked to human activity are having increasingly marked impacts on many 119 wildlife populations, influencing their geographical range due to increasing urbanization and 120 landscape fragmentation (Dirzo et al. 2014) and constraining their behavioural repertoire (Sih, 121 122 Ferrari & Harris 2011). Recently, Tucker et al. (2018) reported a global decrease in the mobility of mammals living in human-disturbed environments. They suggested that animals living in built-123 up landscapes were confined to smaller ranges due to the prevalence of artificial barriers which 124 125 reduced the amplitude of their movements, although this may be accentuated by the availability of 126 supplementary food sources in anthropogenic environments. Gaynor et al. (2018) further showed that, irrespective of taxa, habitat or location, mammals were markedly more nocturnal in response 127 to human disturbance. Indeed, wildlife appears to associate anthropogenic activities with a 128 perceived risk of mortality (Frid & Dill 2002). Responses to human activities are particularly 129 common among hunted species (Stillfried et al. 2015), but have been documented even in the 130 absence of real risk (Creel & Christianson 2008; Clinchy et al. 2016). 131

As large carnivores are currently recolonising Europe (Chapron et al. 2014), many game 132 species are faced with the combined risks associated with human hunting and their natural 133 134 predators. Large carnivores may have significant impact on both the demography (Lehman et al. 2018) and behaviour (Lone et al. 2017) of prey populations in areas where they have become re-135 established. For instance, lynx (Lynx lynx) recolonisation lead to a marked fall in population 136 growth rate of roe deer (*Capreolus capreolus*) in Sweden (from λ =1.08 to 0.94; Andrén & Liberg 137 2015), whereas the presence of olfactory cues for lynx increased the levels of deer vigilance two-138 fold in Germany (Eccard, Meißner & Heurich 2017). Indeed, prey are expected to adopt 139 140 behavioural responses to reduce exposure to humans and predators which may be costly (Lima & 141 Dill 1990; Preisser, Bolnick & Benard 2005), generating a complex landscape of fear (Laundré,
142 Hernández & Altendorf 2001).

Large herbivores are both primary prey for large carnivores and are widely hunted across 143 Europe. Because prey commonly shift their activity patterns as a strategy to avoid predators 144 (Tambling et al. 2015), we might expect them to adjust their diel activity patterns in relation to 145 146 variation in these contrasting risk factors (Lone et al. 2017). Indeed, while human hunting and disturbance are concentrated into daylight hours, natural predators are mostly nocturnal or 147 crepuscular (Kusak, Skrbinšek & Huber 2005; Eriksen et al. 2011). The lynx, for instance, 148 primarily hunts during nighttime and twilight, notably during the first part of the night (Schmidt 149 1999; Heurich et al. 2014). Hence, while large herbivores frequently leave refuge habitat to feed 150 at night in human-dominated landscapes (e.g. Graham et al. 2009 on elephants Loxodonta 151 152 africana; Tolon et al. 2009 on wild boar Sus scrofa; Bonnot et al. 2013 on roe deer; Roberts, Cain III & Cox 2017 on elk *Cervus canadensis*), populations exposed to natural predators might be 153 expected to shift a substantial proportion of their activity to daytime. To minimize exposure to 154 both natural and human risks, we might therefore expect prey to be particularly crepuscular, 155 squeezing as much of their activity as possible into dawn and dusk. Such shifts in diel activity of 156 prey have the potential to mitigate human-wildlife conflicts by lowering the risk of collisions 157 158 between vehicles and wildlife (e.g. Murray & St Clair 2015), or by attenuating the negative impacts of climate change on water sensitive species (Levy et al. 2019). However, such behavioural 159 alterations may be energetically costly, substantially affecting predator-prey dynamics and, 160 ultimately, prey fitness (Creel & Christianson 2008; Kronfeld-Schor et al. 2017; Levy et al. 2019). 161 With the increasingly widespread cohabitation between large carnivores and human activities, it 162

therefore appears important to better understand how prey species respond behaviourally to thecontrasting mortality risks due to hunting and their natural predators.

In this study, using a unique data set generated from activity sensors deployed on 431 165 individual roe deer from 12 populations across Europe, we investigated variation in individual diel 166 activity patterns in relation to the landscapes of fear generated by a natural predator and human 167 168 activities (see Shamoon et al. 2018 for a comparable study at the population-level based on camera traps). We analysed data from populations distributed over a wide gradient of human disturbance, 169 with well-defined hunting seasons, and with contrasting density of lynx, a specialist predator of 170 roe deer (Andersen et al. 2007; Nilsen et al. 2009). We hypothesized that roe deer would adjust 171 their diel activity budgets in relation to variation in the level of human disturbance and lynx 172 predation risk. As human disturbance and hunting are least intense during nighttime, we predicted 173 174 that: 1/ roe deer would be more nocturnal in areas where human disturbance was higher, particularly during the hunting season. In contrast, we expected 2/ the degree of diurnality would 175 be higher in areas where lynx were present due to the higher risk of predation at night, particularly 176 outside of the hunting season. Finally, although large herbivores are routinely crepuscular, they 177 should partition their activity between dawn and dusk in relation to temporal variations in 178 predation risk. Therefore, we expected 3/ roe deer would be more pronouncedly crepuscular in 179 180 areas with high levels of both human disturbance (which is most intense during daytime) and lynx predation (which predominantly occurs at night). Notably, we expected roe deer to be particularly 181 crepuscular at dusk during the hunting season (because hunting occurs mainly during daytime), 182 but at dawn where lynx were present (as lynx hunt primarily during the first part of the night; 183 184 Heurich et al. 2014).

186 Materials and methods

187 Study areas and data collection

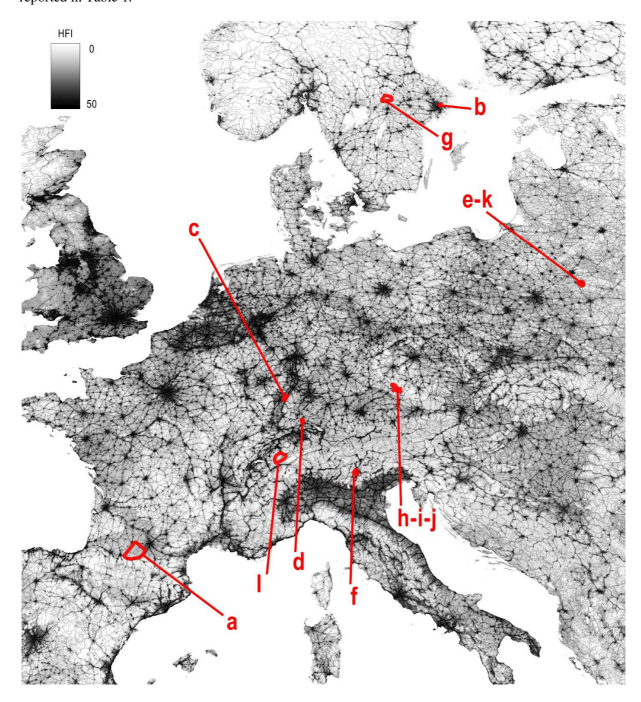
To investigate the influence of predation risk and human disturbance on roe deer diel activity 188 patterns, we analysed activity data obtained within the EURODEER project (http://eurodeer.org). 189 190 Data were collected for a period spanning from 2003 to 2015 and included 431 adult roe deer of more than one year old (254 females and 177 males) from 12 contrasted populations located in 9 191 geographical regions across Europe (Fig. 1). At each study site, roe deer were caught during 192 winter, sexed, aged and equipped with GPS collars (Lotek 3300, Lotek Small WildCell, Vectronic 193 194 GPS Plus or e-obs) carrying activity or acceleration sensors. Collars were programmed to record an activity measurement every 5 min and a GPS fix every 4 to 6 hours (depending on the study 195 areas and the year of monitoring). 196

The studied populations differed in terms of habitats, levels of human disturbance and 197 predation risk (Table 1). Lynx, the main natural predator of roe deer, was present on half of the 198 study sites at densities ranging from 1.0 to 2.5 lynx/100 km² (see Table 1). Wolf (Canis lupus) and 199 bear (Ursus arctos) are potential predators of roe deer, but were each present on very few study 200 sites (bear: Monte Bondone, Italy; wolf: Białowieża, Poland and Grimsö, Sweden). In contrast, 201 202 red fox (Vulpes vulpes) were widespread across the study sites, but exclusively attack neonates. 203 Therefore, we considered the presence of lynx as the predominant component of variation in predation risk. To account for the presence of lynx, we categorized each study site according to 204 relative lynx density: no lynx, low lynx density (sites with transient lynx and densities of 205 approximately 1 lynx/100 km²) and high lynx density (sites with densities of approximately 2 206 207 lynx/100 km²; see Table 1 for more details).

Roe deer were also hunted in all study areas, most commonly during daytime. Although the 208 start and end of the hunting seasons differed slightly among study sites, we could define a non-209 hunting season for both sexes ranging from 15th March to 30th April and a hunting season from 1st 210 211 October to 14th November which were common to all sites. Sit-and-wait hunting occurred on all sites. Drive hunting is also used for roe deer (mainly on three sites: Aurignac, Baden - Rhine 212 valley, Baden - Hegau, but also at Bernese), and for other species (such as wild boar, moose, red 213 214 deer) almost everywhere during the same period (i.e. Aurignac, Baden - Rhine valley, Baden -215 Hegau, Bogesund, Grimsö, Bavarian forest).

To quantify human disturbance, we used the human footprint index (HFI) which is a reliable 216 217 proxy of the overall level of human activities (Venter et al. 2016; Tucker et al. 2018). The HFI is 218 generated from nine global data layers related to the level of human pressure which describe spatial variation in population density, built-up areas, nighttime lights, land use/land cover ratio, 219 220 coastlines, roads, railroads and navigable rivers. At a global scale, values of HFI vary between 0 (the least disturbed areas) to 50 (the most disturbed areas). Using the human footprint map of 2009 221 (i.e. the most recent available, https://wcshumanfootprint.org/, Venter et al. 2016), and based on 222 all pixels (1 km²) within each individual's seasonal home-ranges, we calculated two values of 223 mean HFI for each roe deer, one for the hunting season and one for the non-hunting season. 224 225 Seasonal home ranges were calculated using the fixed kernel home range method at 95% with an 226 ad hoc factor. The mean HFI values varied substantially across populations (from 6.8 to 25.3). Within populations, the mean HFI values also varied substantially among individuals (see Table 227 1), but not between seasons (15.3 [6.9-24.8] during Spring vs 15.2 [6.2-26.2] during Autumn). 228

Fig. 1. Locations of the 12 roe deer populations plotted in red on the European map of the Human Footprint
Index (HFI), ranging from 0 (low HFI in white) to 50 (high HFI in black): a: Aurignac; b: Bogesund; c:
Baden Rhine Valley; d: Baden Hegau; e: Białowieża – open; f: Monte Bondone; g: Grimsö; h, i, j: Bavarian
forest (three populations); k: Białowieża – forest; l: Bernese. The main characteristics of the study sites are
reported in Table 1.



Study area id	Study area name	Sample size	Location (average coordinates)	Lynx presence (density in animals/100 km ²)	Mean individual HFI (range)	Hunting season (both sexes)	Habitat type	
a	Aurignac	209	France (43°29'20"N, 00°88'21"E)	-	11 (5–19)	Sep10 - Feb28	Hilly agricultural landscape with forest patches, meadows and croplands	
b	Bogesund	5	Sweden (59°39'73"N, 18°19'45"E)	-	18 (16–21)	Oct1 - Jan31	Mixed landscape with forest, bogs and croplands	
с	Baden - Rhine Valley	30	Germany (48°63'27"N, 07°97'74"E)	-	19 (13–27)	Sep1 - Jan31	Mixed agricultural landscape with forest patches, meadows and croplands	
d	Baden - Hegau	12	Germany (47°88'31"N, 08°72'93"E)	-	19 (16–23)	Sep1 - Jan31	Mixed agricultural landscape with forest patches, meadows and croplands	
e	Białowieża - open	4	Poland (52°44'49"N, 23°26'35"E)	-	21 (13–23)	Oct1 - Jan15	Agricultural landscape	
f	Monte Bondone	6	Italy (46°02'14"N, 11°01'14"E)	-	25 (13–34)	Sep1 - Oct30	Alpine mountain range	
g	Grimsö	9	Sweden (59°68'23"N, 15°40'17"E)	Lynx (1.0)	7 (5–9)	Oct1 - Jan31	Boreal forest	
h	Bavarian forest - FRG	22	Germany (49°03'56"N, 13°19'07"E)	Lynx (1.2)	9 (6–19)	Sep1 - Jan15	Mixed mountain forest	

Table 1: Characteristics of the study areas.

i	Bavarian forest - RLG	59	Germany (48°54'40"N, 13°28'09"E)	Lynx (1.2)	10 (6–20)	Sep1 - Jan15	Mixed mountain forest
j	Bavarian forest - PJR	14	Germany (48°54'05"N, 13°15'11"E)	Lynx (transient dispersers)	14 (12–15)	Sep1 - Jan15	Mixed landscape with mountain forests and croplands
k	Białowieża - forest	8	Poland (52°39'22''N, 23°29'12''E)	Lynx (2.5)	10 (6–17)	Oct1 - Jan15	Mixed landscape with forest and croplands
1	Bernese	53	Switzerland (46°33'36"N, 07°30'47"E)	Lynx (2.1)	20 (4–29)	Oct1 - Nov15	Mixed landscape with forest and meadows

237 Activity data

The activity sensors on the GPS collars measured the overall level of activity by recording 238 forward/backward and sideway motions (Vectronic, e-obs) or up/down and sideway motions 239 (Lotek) on two axes, X and Y. Because activity measured on the third Z-axis was only available 240 for 4 populations (28% of all individuals), we discarded data on this axis prior to analyses. Lotek 241 3300 collars measured activity as the count of contacts along the X- and Y-axes. For each 5-minute 242 243 interval, the sensors provided the mean value of all activity measurements, indexing the average 244 level of activity associated with the corresponding date and time interval, ranging from 0 to 255 for each axis. Lotek Small WildCell and Vectronic collars measured activity based on the true 245 246 acceleration in the X- and Y-axes by indexing the difference in acceleration between two consecutive measurements and averaging these values within 5-minute intervals ranging from 0 to 247 255 for each axis. E-obs collars sampled acceleration every minute in bursts of 9 seconds and 248 249 provided raw accelerometer readings for both axes. We calculated activity for e-obs collars as above, by averaging the difference in acceleration between two consecutive measurements within 250 5-minute intervals for each axis. Finally, for all collars, we used the sum of the values for the X-251 and Y- sensors as our measure of activity per 5-minute interval, with values ranging from 0 (no 252 activity) to 510 (high activity) (see Bonnot et al. 2016). Considering only the two 45-day seasons 253 254 analysed in this study, we obtained 10,866,096 activity records corresponding to an average of 76 255 ± 25 days of monitoring per individual.

256

257 **Diurnality index**

To evaluate our first hypotheses, that roe deer would be more nocturnal where human-related disturbance was high but more diurnal where lynx were present, we generated a diurnality index based on the relative level of activity during daylight compared to nighttime for each individual
on each given day (Hoogenboom et al. 1984). Because we wished to focus on the shift of activity
from daytime to nighttime in this analysis, we removed the twilight periods which we defined here
as the period of four hours centered on sunrise (dawn) and sunset (dusk). Date-specific times for
sunrise and sunset for each study site were obtained from the National Oceanic & Atmospheric
Administration (https://www.noaa.gov/). The diurnality index was calculated as follows (Eqn 1),

266
$$D_i = A_{DAYi} / (A_{DAYi} + A_{NIGHTi})$$
 (Eqn 1)

where A_{DAYi} is the mean activity value during daytime of day i and A_{NIGHTi} is the mean activity value during nighttime (from midnight to 2 hours before sunrise and from 2 hours after sunset to midnight) of day i for a given individual. D_i ranges between 0 (when a given deer was strictly nocturnal during day i) and 1 (when a given deer was strictly diurnal during day i). By using a diurnality index calculated as the ratio between daytime and nighttime activity levels per 24-h and per individual, we circumvented the need for standardizing the activity data.

273

274 Crepuscularity index

To evaluate our prediction that roe deer would be more pronouncedly crepuscular in areas with high levels of both human disturbance and predation risk, we calculated a crepuscularity index for both dawn and dusk. In order to restrict this index to the peak crepuscular period, we defined dawn and dusk as periods of two hours, comprising one hour each side of sunrise and sunset, respectively. Thus, the index of crepuscularity is a proxy of the relative level of activity during dawn (or dusk) compared to overall activity recorded during a given 24-hour cycle for each individual (Eqns 2 and 3).

283	$C_{DAWNi} = A_{DAWNi} / (A_{DAY'i} + A_{NIGHT'i} + A_{DAWNi} + A_{DUSKi})$	(Eqn 2)

 $284 \qquad C_{\text{DUSK}i} = A_{\text{DUSK}i} / (A_{\text{DAY}'i} + A_{\text{NIGHT}'i} + A_{\text{DAWN}i} + A_{\text{DUSK}i}) \tag{Eqn 3}$

where C_{DAWNi} and C_{DUSKi} are, respectively, the indices of crepuscularity during dawn and dusk and A_{DAWNi}, A_{DUSKi}, A_{DAY'i} and A_{NIGHT'i} are, respectively, the mean activity values during dawn, dusk, daytime and nighttime during day i, for a given individual. Crepuscularity indices may range between 0 (when a given deer was strictly inactive during dawn/dusk during day i) and 1 (when a given deer was strictly active during dawn/dusk during day i).

290

291 Statistical analyses

292 Overview

In a first step, we explored temporal variation in diel activity in relation to the risk of lynx predation and hunting using density functions following Ridout & Linkie (2009). We compared diel activity patterns by quantifying the degree of overlap between seasons (hunting vs. non-hunting) for each population. In a second step, we analysed variation in the indices of diurnality and crepuscularity using Generalized Linear Mixed Models (GLMMs). Because the indices ranged within the interval [0,1], we transformed both metrics using the equation proposed by Cribari-Neto & Zeileis (2010) (Eqn 4) so that they conformed to a beta distribution (i.e. comprised within the interval]0,1[).

$$300 \quad (Y_i (n-1) + 0.5) / n \tag{Eqn 4}$$

301 where Y_i represents the value of a given index (diurnality or crepuscularity) during the day i and 302 n is the sample size (i.e. the total number of observations for a given index). All analyses were performed in R version 3.3.3 (R Development Core Team 2017). Diel activity patterns and overlaps were estimated using the '*overlap*' package (Ridout & Linkie 2009) and GLMMs were fitted using the '*glmmTMB*' package (Brooks et al. 2017).

306

307 Diel activity patterns across seasons and lynx densities

We classified activity data into active *vs* inactive behaviour based on the frequency distribution of activity measurements (Gervasi, Brunberg, & Swenson 2006) for each collar type independently (for more details, see Appendix S1 in the Supporting Information file). This method uses a specific threshold to discriminate active and inactive behaviour which is, therefore, insensitive to variation in absolute values of activity among individuals.

313 Using the above mentioned approach, we described deer diel activity patterns for each study site and quantified the degree of overlap between the non-hunting and hunting seasons, based 314 315 on the observations where a given individual was active. To account for changes in the sun's position across seasons, instead of using clock time, we first scaled the time of day to sunrise and 316 sunset. We did so by respectively adjusting the time of each sunrise and sunset to $\pi/2$ and $3\pi/2$ 317 with the 'sunTime' function (see Nouvellet et al. 2012 and 'overlap' R-package). Once sunset and 318 sunrise times were synchronized across seasons and populations, we then assessed daily activity 319 patterns by fitting circular kernel density functions (Fig. 2 and Appendix S2) and calculated a 320 321 coefficient of overlap (i.e. the common area under the kernel density curves; Ridout & Linkie 2009). The coefficient of overlap ranges from 0, indicating total temporal independence between 322 323 activity patterns, to 1, indicating perfect synchronization. A high coefficient of overlap between the two seasons indicates that activity patterns are similar. 324

326 Variation in diurnality in relation to human disturbance, hunting and lynx

To evaluate our first hypotheses (H1 and H2) concerning the relative level of activity during 327 daytime compared to nighttime, we fitted GLMMs to analyse variation in the diurnality index in 328 relation to i/ the overall level of human disturbance, indexed by the HFI, ii/ the season, as a proxy 329 of hunting activity (non-hunting vs hunting), and iii/ the risk of predation indexed by lynx density 330 (three-modality variable: no-lynx, low lynx density, high lynx density). Because we expected the 331 332 impact of human disturbance and predation risk on diurnality to differ in relation to hunting 333 activity, the most complex model contained two two-way interactions between the HFI and the season, and between lynx density and the season. Sex was included in all models to control for 334 335 potential differences in the level of diurnality between males and females (Pagon et al. 2013). However, as we had no a priori reason to expect one sex to respond to risk more strongly than the 336 other in terms of temporal shifts in diel activity, we did not include any interactive effects of sex 337 338 with other terms in the models. We included individual identity as a random effect on the intercept in all models as we had repeated measures of the diurnality index (one measure per day) for each 339 340 individual. For model selection, we used Akaike's Information Criterion (AIC, Burnham & Anderson 2002), Akaike weights and the number of parameters to select the most parsimonious 341 model that best described the data. 342

343

344 Variation in crepuscularity in relation to human disturbance, hunting and lynx

To evaluate our H3 hypotheses concerning the relative level of activity during dawn and dusk compared to the rest of the day, we fitted GLMMs to analyse variation in the crepuscularity index in relation to the HFI, the season and lynx density, but including the crepuscular period (dawn *vs*. dusk) as a binary factor. Because we expected that the influence of both hunting and predation risk would differ between dawn and dusk, and that these sources of risk might be cumulative, the most complex model contained three two-way interactions between lynx density and the crepuscular period, between the season and the crepuscular period, and between lynx density and the season, with the HFI as an additional fixed effect. As before, we included sex as a fixed effect and individual identity as a random effect on the intercept in all models and used AIC criteria for model selection.

355

356 **Results**

357 Diel activity patterns across seasons and lynx densities

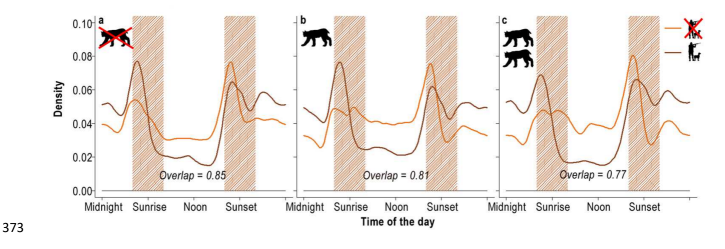
In all the 12 studied populations, we observed a clear bimodal diel activity pattern for roe deer 358 which was consistent across seasons (see Fig. 2 for an example of three populations with varying 359 lynx density; see Appendix S2 for the full representation of the 12 populations), as indicated by 360 the high coefficients of overlap between seasons (mean overlap of 0.81 [0.71-0.87]). That is, roe 361 deer expressed marked peaks of activity during the two crepuscular periods, with moderate levels 362 of activity during daytime and nighttime (Fig. 2). As predicted, deer were consistently less diurnal 363 during the hunting season compared to the non-hunting season in all populations (Fig. 2 and 364 365 Appendix S2). In contrast, during the non-hunting season and where lynx were present, roe deer were more active during daytime (Fig. 2b and c). Note, however, that there was considerable 366 variation among populations in diel activity (see Appendix S2 and the values of HFI on each plot). 367

369 Fig. 2. Graphical representation of diel activity patterns during the non-hunting (orange) and hunting

370 (brown) seasons for three roe deer populations with varying densities of lynx (a: Baden Rhine Valley, n=30,

371 no lynx; b: Bavarian RLG, n = 59, low lynx density, and c: Białowieża forest, n = 8, high lynx density).

372 The hatched shading represents the crepuscular periods as defined for the diurnality index.



374

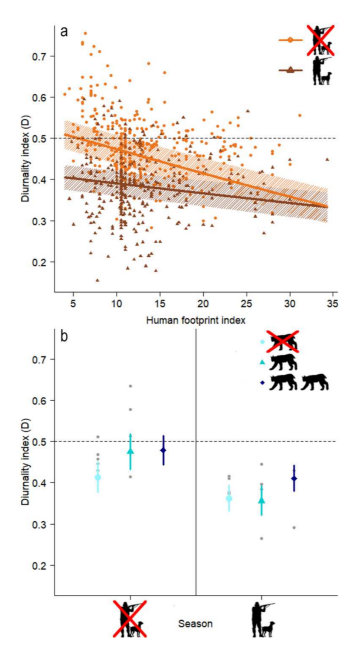
375 Variation in the level of diurnality

The diurnality index averaged per individual ranged from 0.28 to 0.75 during the non-hunting 376 377 season, and from 0.15 to 0.59 during the hunting season. Model selection revealed that the 378 diurnality index was best described by two two-way interactions between lynx density and season, and between HFI and season, plus the additive effect of sex (the AIC value was much lower than 379 that of any of the simpler models, i.e. $\Delta AIC \ge 240$, AIC weight = 1, see Appendix S3). In 380 accordance with our first hypothesis, the level of roe deer diurnality decreased, on average, by a 381 382 factor of 1.37 over the gradient of HFI (Fig. 3a). Indeed, roe deer were relatively less diurnal when 383 human disturbance was high, particularly during the non-hunting season (mean diurnality estimates \pm standard error: D = 0.51 \pm 0.01 in areas with low HFI vs D = 0.33 \pm 0.02 in areas with 384 high HFI). Roe deer were also consistently less diurnal during the hunting season compared to the 385

non-hunting season (Fig. 3), although this difference was not significant in areas with high human
disturbance (for HFI > 15).

In contrast, as predicted by our second hypothesis, roe deer were relatively more diurnal when lynx were present. On average, the level of diurnality was 1.2 higher where lynx were present at high density compared to areas without lynx ($D = 0.45 \pm 0.02$ and $D = 0.39 \pm 0.02$, respectively Fig. 3b). Interestingly, the effect of lynx density on the level of diurnality was attenuated during the hunting season, particularly where lynx were present at low density. Finally, males were only slightly less diurnal overall than females (see Appendix S3: Figure S3).

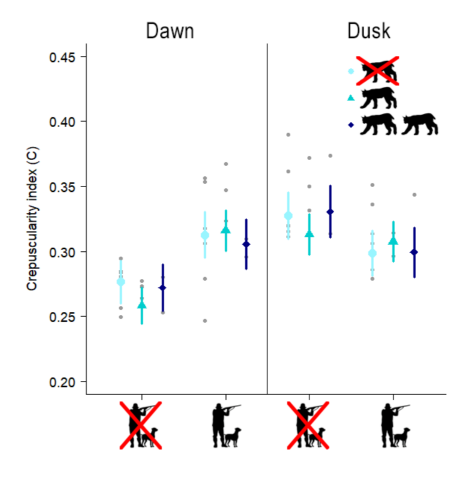
Fig. 3. Graphical representation of the best model describing variation in the level of roe deer diurnality (D) as a function of the two-way interactions between (a) the hunting season and the Human Footprint Index (HFI) and (b) the hunting season and lynx density. The dotted line represents an equivalent level of activity during daytime and nighttime (i.e. D = 0.5). 95% confidence intervals are represented (a) by the dashed areas and (b) by bars. The points correspond to the diurnality indices averaged (a) per season and per individual and (b) per season and per population.



402 Variation in the level of crepuscularity

On average, across all individuals, the crepuscularity index ranged from 0.29 [0.16-0.61] during 403 dawn to 0.32 [0.18-0.47] during dusk, indicating that roe deer expressed around 60% of their diel 404 405 activity during twilight periods (Fig. 4). The most parsimonious model that best described variation in the crepuscularity index included the three two-way interactions between lynx density and the 406 crepuscular period, between lynx density and the season, and between the season and the 407 408 crepuscular period, plus sex (AIC weight = 0.34, $\Delta AIC = 1.33$; see Appendix S4). Contrary to our 409 expectation, we found no marked difference in the global level of crepuscularity between the hunting and non-hunting seasons or among areas with contrasting lynx densities, irrespective of 410 411 potential differences between dawn and dusk (Fig. 4). However, we found an effect of hunting on 412 how deer partitioned their activity between dawn and dusk. During the hunting season, roe deer shifted their crepuscular activity to become relatively more active at dawn, but slightly less active 413 414 at dusk. More precisely, they increased their level of crepuscular activity at dawn by a factor of 1.19 compared to the non-hunting season (non-hunting: $C_{DAWN} = 0.26 \pm 0.02$; hunting: $C_{DAWN} =$ 415 416 0.31 ± 0.02), and decreased their level of crepuscular activity at dusk by a factor of 1.07 (hunting: $C_{DUSK} = 0.30 \pm 0.02$; non-hunting: $C_{DUSK} = 0.32 \pm 0.02$). Contrary to our expectations, we did not 417 find a marked effect of lynx density on the partitioning of activity between dawn and dusk (Fig. 418 419 4). Finally, although both sex and HFI featured in the two best models, there was no obvious relationship with the level of crepuscularity in either case (see Appendix S4: Figures S4). 420

Fig. 4. Graphical representation of the model describing variation in the level of roe deer crepuscularity (C) as a function of the three two-way interactions between the hunting season and lynx density, between the hunting season and the crepuscular period of the day (dawn *vs* dusk), and between lynx density and the crepuscular period of the day (dawn *vs* dusk). The predictions are plotted with their 95% confidence intervals. The grey points correspond to the crepuscularity indices averaged per season and per population.



430 **Discussion**

In the context of the increasingly widespread coexistence of large carnivores and humans in 431 Europe, our study provides important insights about the contrasting influence of predation risk and 432 anthropogenic activities on temporal partitioning of activity in their ungulate prey. By comparing 433 434 diel activity patterns among 12 populations of roe deer across Europe, our analyses revealed marked variation in diurnality in response to both lethal and non-lethal human activity and, to a 435 lesser degree, the risk of predation by lynx. These modifications in the temporal partitioning of 436 ungulate activity likely have knock-on effects on a variety of ecological processes. As a 437 438 perspective to our work, we discuss the implications of our results below, notably in terms of predator-prey interactions and human-wildlife coexistence. 439

440

441 Impact of lethal and non-lethal human activities on diurnality of roe deer

First, as expected (H1), the way in which roe deer partitioned their activity over the day was 442 strongly modified by the degree of anthropisation in the surrounding landscape. On average, deer 443 were globally 1.37 times less active during the day in areas with a high human footprint, and up to 444 1.52 times less active outside the hunting season (see Fig. 3a). Moreover, we found an additional 445 446 effect of hunting such that roe deer shifted their diel activity cycle by, on average, a factor of 1.20 to become predominantly nocturnal during the open season (Fig. 3b, Appendix S2). Overall, our 447 results are highly consistent with the recent meta-analysis of Gaynor et al. (2018) who showed that 448 wild mammals increased their degree of nocturnality by a factor of 1.36 in response to human 449 450 activity.

One important novelty of our study is that we were able to disentangle the effects of the 451 general background level of human disturbance on roe deer diurnality from lethal effects due to 452 hunting. In particular, we found that hunting had a greater impact on the level of diurnality for 453 animals living in relatively undisturbed areas (Fig. 3a). Because we evaluated the effect of hunting 454 at the seasonal scale, the observed response reflects behavioural plasticity (i.e. the array of 455 behavioural responses of an individual to variation in the environment, Komers 1997) of 456 457 individuals to a modification in their landscape of risk (Reebs 2002; Murray & StClair 2015). 458 Behavioural plasticity is likely one of the main keys behind the success of roe deer in humandominated environments (Andersen, Duncan & Linnell 1998). In these environments, non-lethal 459 460 human activities are often considered analogous to predation risk (Frid & Dill 2002) so that prey adopt comparable anti-predator responses to disturbance. Whereas plastic behavioural responses 461 are often considered adaptive, responses of prey to non-lethal stimuli could be maladaptive in 462 463 terms of the loss of time and energy that would otherwise be allocated to fitness-enhancing activities, generating an ecological trap. For instance, roe deer adjust their anti-predator behaviours 464 in relation to proximity to human settlements (e.g. their vigilance levels, see Benhaïem et al. 2008, 465 and flight distances, see Bonnot et al. 2017) which may potentially incur a fitness cost (Bonnot et 466 al. 2018). However, these effects also appear to be highly dependent on the availability of refuge 467 habitat and the period of the day (Benhaïem et al. 2008; Bonnot et al. 2013; Sönnichsen et al. 468 2013), indicating that animals may buffer human disturbance by adjusting both their space use and 469 temporal partitioning of activity. 470

471 However, the extent of plasticity is finite and our results further suggest that roe deer living in 472 the most human-disturbed areas had reached the upper limit of their potential plasticity with 473 respect to the degree of nocturnality. Indeed, these animals were more nocturnal year round in

comparison with roe deer living in relatively undisturbed areas and they did not modify their diel 474 475 activity patterns further during the hunting season (Fig. 3a). More specifically, at a threshold of approximately 15 for the human footprint index, roe deer diurnality no longer differed between the 476 hunting and non-hunting seasons. On the HFI scale from 0 for wild areas to 50 for very developed 477 areas, this value describes relatively undeveloped environments, with low levels of human 478 pressure. This suggests that, even in areas of relatively low human pressure, anthropogenic 479 480 activities may substantially modify the degree of nocturnality in prey. However, for the specific 481 case of our study species, we also know that roe deer are constrained to maintain a minimum level of activity to feed during daytime, even in the most human-disturbed landscapes. Indeed, all 482 483 ruminants must alternate feeding bouts with periods of rest and rumination (Hofmann 1989), but because the roe deer has a particularly small rumen, these cycles are relatively short (Duncan et 484 al. 1998). Roe deer also have highly flexible diets (Abbas et al. 2011), exploiting substantially 485 486 different foods in spring and autumn, which should affect their spatial behaviour (i.e. habitat selection, Godvik et al. 2009; Bonnot et al. 2018). Although we have no a priori reason why such 487 seasonal differences in the risk-resource trade-off should influence the partitioning of activity 488 between day and night, further work should investigate whether preferred habitat is also associated 489 with higher levels of risk. 490

491

492

Because large carnivores are predicted to influence the landscape of risk and the landscape of fear of their prey, prey should adjust their behavioural responses to spatial and temporal variation in the risk associated with their natural predators (Lima & Dill 1990; Manning, Gordon & Ripple

Impact of a natural predator on diurnality of roe deer

496 2009; Dröge et al. 2017). In support of this hypothesis, we found that roe deer shifted from a

predominantly nocturnal activity cycle to a more diurnal rhythm when lynx were present (Fig. 3).
In a similar manner, Tambling et al. (2015) showed that African ungulates were more likely to be
active during the day when cohabiting with lions (*Panthera leo*) and hyaenas (*Crocuta crocuta*),
thereby decreasing activity overlap with these nocturnal predators.

In our study, the behavioural modifications associated with the presence of lynx were mainly 501 confined to the spring, when no hunting occurred. During hunting, roe deer became predominantly 502 nocturnal, whether or not lynx were present (Fig. 3b). This result highlights the strong influence 503 of humans as a "super-predator" in shaping the behavioural responses of prey (Ciuti et al. 2012a, 504 505 b; Clinchy et al. 2016), with potential fitness consequences, notably in multi-predator landscapes 506 (e.g. Gehr et al. 2018). For example, humans kill mesocarnivores at more than four times the rate at which they are killed by non-human predators (Darimont et al. 2015). Moreover, human 507 508 activities create a well-defined landscape of risk, which is often highly predictable in time and 509 space, provoking direct and immediate behavioural responses of prey (Cromsigt et al. 2013). In contrast, the landscape of risk due to large carnivores may be more difficult for their prey to 510 predict. 511

Besides humans and lynx, wolves were also present in two of our study areas, which likely 512 created even more complex landscapes of risk for roe deer, although they are not their main prey 513 514 in these ecosystems (see Jedrzejewski et al. 2002; Sand et al. 2005). Wolf can also alter the spatial behaviour of their prey (e.g. Dellinger et al. 2019; Bongi et al. 2008), but there is less evidence 515 that they alter their diel activity patterns (Eriksen et al. 2011 on moose Alces alces, but see Kohl 516 517 et al. 2018 on elk *Cervus elaphus*). One explanation could be that the cues associated with risk of predation by ambush predators, like lynx, are generally more reliable than those for coursing 518 predators, like wolf (Preisser, Orrock & Schmitz 2007; Kohl et al. 2019). Further studies are 519

needed to understand the influence of predator hunting tactics on the activity of their prey, notablyin multi-predator environments (see also Kohl et al. 2019).

These landscapes of risk are also likely dynamic depending on the degree of interaction between humans and natural predators. For instance, large carnivores generally avoid humans (Oriol-Cotteril et al. 2015; Belotti et al. 2018), which may create a human-shield effect for their prey (Berger 2007), but also may result in higher kill rates (Smith, Wang & Wilmers 2015). Lynx, in particular, must tradeoff avoidance of human activities during daytime against prey availability (Basille et al. 2009; Gehr et al. 2017; Filla et al. 2017), which should accentuate the risk of predation for roe deer during the night.

529

530 Crepuscularity of roe deer in a multi-predator landscape

531 Large herbivores are frequently reported to be markedly crepuscular (e.g. Krop-Benesch et al. 2013), which has commonly been interpreted as an antipredator strategy (Kamler, Jedrzejewska & 532 Jędrzejewski 2007; Monterroso, Alves & Ferreras 2013; Swinnen, Hughes & Leirs 2015). This is 533 expected to be particularly the case in complex landscapes of risk composed of coexisting 534 predators with contrasting hunting methods and diel activity patterns (Gehr et al. 2018; Lone et al. 535 2014). However, we found little support for this hypothesis here, as the level of crepuscularity did 536 537 not appear to be linked to variations in either the human- or lynx-induced risk of predation. Overall, roe deer carried out around 60% of their diel activity during dawn and dusk, but, contrary to our 538 539 expectation, this proportion did not increase when they were exposed simultaneously to both the risk of predation from lynx and hunting (Fig. 4). In line with previous studies, our results rather 540 indicate a strong physiological and/or behavioural constraint promoting the maintenance of 541 crepuscular activity peaks in ungulates, irrespective of the risk context (Kronfeld-Schor et al. 2001; 542

Massé & Côté 2013; Bonnot et al. 2016; but see Loe et al. 2007 and Long et al. 2013). Another explanation could be that any further increase in crepuscular activity would not be an efficient anti-predator strategy (Kronfeld-Schor & Dayan 2003), as both humans and lynx can potentially hunt at twilight.

Finally, while markedly crepuscular, we still found that large herbivores may partition their 547 crepuscular activity differently between dusk and dawn, notably depending on hunting risk (Fig. 548 549 4). Indeed, our results suggest that the risk of predation by lynx did not markedly influence how 550 roe deer partitioned their activity between dawn and dusk, whereas deer became more active at dawn during the hunting season compared to spring. We suggest that, because roe deer were able 551 552 to feed more during the day when there is no risk of hunting, they were also less constrained to be 553 crepuscular. This could explain why roe deer were markedly less active at dawn during spring, when human daily activities may be prevalent in agro-systems, but maintained their activity peak 554 555 at dusk when human presence is generally lower (Fig. 4). Although the disturbing effects of hunting on prey behaviour are well-documented (Cromsigt et al. 2013; Gaynor et al. 2018), further 556 work will be required to quantify the variation in how large herbivores respond to varying hunting 557 methods and sources of disturbance, and the cascading effects of their resulting behaviours on 558 ecosystem functioning. 559

560

561 Conclusion and perspectives

562 Our study provides further evidence of the strong behavioural plasticity of large herbivores 563 which allows them to thrive in heavily anthropogenic landscapes. Behavioural plasticity plays a 564 key role in species adjustment to rapid environmental change due to human activities (Sih et al. 565 2011) and is also likely crucial in the context of the return of large carnivores. Prey may respond

in several ways to variations in the level of predation risk: moving to safer habitat during risky 566 times (Godvik et al. 2009), decreasing their movement rate (Picardi et al. 2019), adjusting their 567 levels of vigilance (Dröge et al. 2017) or escape decisions (Bonnot et al. 2015, 2017). With recent 568 advances in biologging, we will soon be able to investigate the fine-scale behavioural responses 569 of prey, as well as their ecological and energetic costs in a dynamic landscape of fear (Brown et 570 al. 2013; Kays et al. 2015; Williams et al. 2017; Kröschel et al. 2017). For example, the observed 571 572 shift of roe deer to nocturnal activity in response to hunting could result in a higher risk of lynx predation during the hunting season (Gehr et al. 2018), or in lower foraging efficiency due to an 573 increase in alternative anti-predator behaviours such as vigilance (Benhaïem et al. 2008), 574 575 potentially affecting predator-prey dynamics, ecological communities and ecosystem functioning 576 (Fortin et al. 2005). Indeed, as both prey and consumer of vegetation, large herbivores are key ecosystem engineers with marked impacts on a variety of ecological processes (Côté et al. 2004). 577 578 Therefore, shifts in the temporal partitioning of their activity in response to predation risk and/or human disturbance likely have knock-on effects on the frequency and spatial distribution of 579 important ecosystem services (e.g. seed and nutrient transfer, biodiversity) and disservices (e.g. 580 road traffic accidents, damage to saplings and crops, parasite abundance). For example, roe deer 581 is one of the main hosts for adult ticks (Ixodes ricinus) which are more active during the night 582 (Belozerov 1982; Mejlon 1997). Any shift to diurnal activity in deer populations exposed to 583 nocturnal predators could decrease their level of infestation and, hence, the dispersal of ticks and 584 tick-borne diseases over the landscape (Hofmeester et al. 2017). The modification of the activity 585 patterns of prey species to the contrasting pressures of human activities and large carnivores may 586 therefore result in behaviourally-mediated trophic cascades which urgently require further 587 investigation. 588

589 Authors' contributions

590 NCB, PK, MH, AJMH, FC conceived the ideas and designed the study. AJMH, NM, PK, MH, MK, AB, 591 FC, BG, LS provided the data with the critical help of JDG who managed the database. OC and NCB carried 592 out the statistical analyses with the help of AJMH and NM. NCB, OC and AJMH took the lead in writing 593 the manuscript. All authors contributed to the interpretation of the results, provided critical feedback on the 594 manuscript and gave final approval for publication.

595

596 Acknowledgements

This manuscript was conceived within the EURODEER collaborative project (paper #11 of the 597 EURODEER series; http://www.euromammals.org).The authors are grateful to all members for their 598 599 support for the initiative. The EURODEER spatial database is hosted by Fondazione Edmund Mach. We also thank three anonymous referees and the associate editor for their constructive comments which 600 contributed to markedly improve the manuscript. This project was supported by the Marie-Claire Cronstedts 601 foundation to NCB and PK, by the "Move-It" ANR grant ANR-16-CE02-0010-02 to AJMH and NM and 602 603 by the Institute for Zoo and Wildlife Research (IZW), the Mammal Research Institute - Polish Academy of Sciences, the Polish Ministry of Science and Higher Education (grant no N N304172536) to LS (data from 604 605 Białowieża, Poland). The authors declare that they have no conflict of interest.

606

607 Data accessibility

Data used in this study are available from the Dryad Digital Repository (Bonnot, Couriot et al. 2019)
at https://doi.org/10.5061/dryad.1zcrjdfnm. Raw data are also available through the EURODEER platform.
Anyone is welcome to join the EURODEER project and obtain an access to the database after contacting
the persons in charge (see <u>https://eurodeer.org/contacts/</u>).

613 **References**

- Abbas, F., Morellet, N., Hewison, A.J.M., Merlet, J., Cargnelutti, B., Lourtet, B., Angibault,
 J.-M., Daufresne, T., Aulagnier, S. & Verheyden, H. (2011) Landscape fragmentation generates
 spatial variation of diet composition and quality in a generalist herbivore. *Oecologia*, 167(2), 401411
- Andersen, R., Duncan, P., & Linnell, J.D.C. (Eds.). (1998). The European roe deer: thebiology of success.
- Andersen, R., Karlsen, J., Austmo, L.B., Odden, J., Linnell, J.D.C & Gaillard, J.-M. (2007)
- 621 Selectivity of Eurasian lynx *Lynx lynx* and recreational hunters for age, sex and body condition in
- 622 roe deer *Capreolus capreolus*. *Wildlife Biology*, 13(4), 467-474
- Andrén, H. & Liberg, O. (2015) Large impact of Eurasian lynx predation on roe deer
 population dynamics. *PloS One*, 10(3), e0120570
- Basille, M., Herfindal, I., Santin-Janin, H., Linnell, J.D.C, Odden, J., Andersen, R., Høgda,
- 626 K.A. & Gaillard, J.-M. (2009). What shapes Eurasian lynx distribution in human dominated
- 627 landscapes: selecting prey or avoiding people? *Ecography*, 32(4), 683-691
- 628 Belotti, E., Mayer, K., Kreisinger, J., Heurich, M., & Bufka, L. (2018). Recreational activities
- affect resting site selection and foraging time of Eurasian lynx (*Lynx lynx*). Hystrix, 29, 181–189
- Benhaiem, S., Delon, M., Lourtet, B., Cargnelutti, B., Aulagnier, S., Hewison, A.J.M.,
- 631 Morellet, N. & Verheyden, H. (2008) Hunting increases vigilance levels in roe deer and modifies
- 632 feeding site selection. *Animal Behaviour*, 76(3), 611-618
- Belozerov, VN. (1982). Diapause and biological rhythms in ticks. *Physiology of ticks* (eds
- 634 F.D. Obenchain & R. Galun), pp. 469-500 Oxford: Pergamon Press, Oxford

- Berger, J. (2007) Fear, human shields and the redistribution of prey and predators in protected
 areas. *Biology Letters*, 3(6), 620-623
- Bongi, P., Ciuti, S., Grignolio, S., Del Frate, M., Simi, S., Gandelli, D. & Apollonio, M. (2008)
 Anti-predator behaviour, space use and habitat selection in female roe deer during the fawning
- 639 season in a wolf area. *Journal of Zoology*, 276(3), 242-251
- Bonnot, N.C., Couriot, O., Berger, A., Cagnacci, F., Ciuti, S., De Groeve, J., Gehr, B.,
- Heurich, M., Kjellander, P., Kröschel, M., Morellet, N., Soennichsen, L. & Hewison, A.J.M.
- 642 (2019) Data from: Fear of the dark? Contrasting impacts of humans vs lynx on diel activity of roe
- 643 deer across Europe. *Dryad Digital Repository* https://doi.org/10.5061/dryad.1zcrjdfnm
- Bonnot, N.C., Goulard, M., Hewison, A.J.M., Cargnelutti, B., Lourtet, B., Chaval, Y. &
 Morellet, N. (2018) Boldness-mediated habitat use tactics and reproductive success in a wild large
- 646 herbivore. Animal Behaviour, 145, 107-115
- Bonnot, N.C., Hewison, A.J.M., Morellet, N., Gaillard, J.-M., Debeffe, L., Couriot, O.,
- 648 Cargnelutti, B., Chaval, Y., Lourtet, B., Kjellander, P. & Vanpé, C. (2017) Stick or twist: roe deer
- adjust their flight behaviour to the perceived trade-off between risk and reward. *Animal Behaviour*, 124, 35-46
- Bonnot, N.C., Morellet, N., Hewison, A.J.M., Martin, J.-L., Benhamou, S. & Chamaillé-
- Jammes, S. (2016) Sitka black-tailed deer (Odocoileus hemionus sitkensis) adjust habitat selection
- and activity rhythm to the absence of predators. *Canadian Journal of Zoology*, 94(6), 385-394
- Bonnot, N., Morellet, N., Verheyden, H., Cargnelutti, B., Lourtet, B., Klein, F. & Hewison,
- A.J.M. (2013) Habitat use under predation risk: hunting, roads and human dwellings influence the
- 656 spatial behaviour of roe deer. European Journal of Wildlife Research, 59(2), 185-193

657	Bonnot, N., Verheyden, H., Blanchard, P., Cote, J., Debeffe, L., Cargnelutti, B., Klein, F.,
658	Hewison, A.J.M. & Morellet, N. (2015) Interindividual variability in habitat use: evidence for a
659	risk management syndrome in roe deer? Behavioral Ecology, 26(1), 105-114
660	Brown, D.D., Kays, R., Wikelski, M., Wilson, R., & Klimley, A.P. (2013). Observing the
661	unwatchable through acceleration logging of animal behavior. Animal Biotelemetry, 1(1), 20
662	Brooks, M.E., Kristensen, K., van Benthem, K.J., Magnusson, A., Berg, C.W., Nielsen, A.,
663	Skaug, H.J., Maechler, M. & Bolker, B.M. (2017) glmmTMB balances speed and flexibility
664	among packages for zero-inflated generalized linear mixed modeling. The R Journal, 9(2), 378-
665	400
666	Burnham, K.P. & Anderson, D.R. (2002) Model selection and multimodel inference: a
667	practical information-theoretic approach. Second edition. Springer-Verlag, New York
668	Chapron, G., Kaczensky, P., Linnell, J. D., von Arx, M., Huber, D., Andrén, H., López-Bao,
669	J.V., Adamec, M., Álvares, F., Anders, O., Balčiauskas, L., Vaidas Balys, V., Bedő, P., Bego, F.,
670	Blanco, J.C., Breitenmoser, U., Brøseth, H., Bufka, L., Bunikyte, R., Ciucci, P., Dutsov, A.,
671	Engleder, T., Fuxjäger, C., Groff, C, Holmala, K., Hoxha, B., Iliopoulos, Y., Ionescu, O., Jeremić,
672	J., Jerina, K., Kluth, G., Knauer, F., Kojola, I, Kos, I., Krofel, M., Kubala, J., Kunovac, S., Kusak,
673	J., Kutal, M., Liberg, O., Majić, A., Männil, P., Manz, R., Marboutin, E., Marucco, F., Melovski,
674	D., Mersini, D., Mertzanis, Y., Mysłajek, R.W., Nowak, S., Odden, J., Ozolins, J., Palomero, G.,
675	Paunović, M., Persson, J., Potočnik, H., Quenette, PY., Rauer, G., Reinhardt, I., Rigg, R., Ryser,
676	A., Salvatori, V., Skrbinšek, T., Stojanov, A., Swenson, J.E., Szemethy, L., Trajçe, A., Tsingarska-
677	Sedefcheva, E., Váňa, M., Veeroja, R., Wabakken, P., Wölfl, M., Wölfl, S., Zimmermann, F.,
678	Zlatanova, D. & Boitani, L. (2014) Recovery of large carnivores in Europe's modern human-
679	dominated landscapes. Science, 346(6216), 1517-1519

- 680 Ciuti, S., Muhly, T.B., Paton, D.G., McDevitt, A.D., Musiani, M. & Boyce, M.S. (2012a)
- 681 Human selection of elk behavioural traits in a landscape of fear. *Proceedings of the Royal Society*
- 682 of London B: Biological Sciences, rspb20121483
- 683 Ciuti, S., Northrup, J.M., Muhly, T.B., Simi, S., Musiani, M., Pitt, J.A. & Boyce, M.S. (2012b)
- Effects of humans on behaviour of wildlife exceed those of natural predators in a landscape of fear. *PloS one*, 7(11), e50611
- Clinchy, M., Zanette, L.Y., Roberts, D., Suraci, J.P., Buesching, C.D., Newman, C. &
 Macdonald, D.W. (2016). Fear of the human "super predator" far exceeds the fear of large
 carnivores in a model mesocarnivore. *Behavioral Ecology*, 27(6), 1826-1832
- Côté, S.D., Rooney, T.P., Tremblay, J.-P., Dussault, C. & Waller, D.M. (2004) Ecological
 impacts of deer overabundance. *Annual Review of Ecology, Evolution, and Systematics*, 35, 113147
- 692 Creel, S. & Christianson, D. 2008. Relationships between direct predation and risk effects.
 693 *Trends in Ecology and Evolution*, 23, 194-201
- 694 Cribari-Neto, F. & Zeileis, A. (2010) Beta Regression in R. *Journal of Statistical Software*,
 695 34(2), 1–24
- 696 Cromsigt, J.P.G.M., Kuijper, D.P.J., Adam, M., Beschta, R.L., Churski, M., Eycott, A.,
- 697 Kerley, G.I.H., Mysterud, A., Schmidt, K. & West, K. (2013) Hunting for fear: innovating
- 698 management of human–wildlife conflicts. Journal of Applied Ecology, 50(3), 544-549
- Darimont, C.T., Fox, C.H., Bryan, H.M. & Reimchen, T.E. (2015) The unique ecology of
 human predators. *Science*, 349(6250), 858-860

- Dellinger, J.A., Shores, C.R., Craig, A., Heithaus, M.R., Ripple, W.J. & Wirsing, A.J. (2019)
 Habitat use of sympatric prey suggests divergent anti-predator responses to recolonizing gray
 wolves. *Oecologia*, 189(2), 487-500
- Dirzo, R., Young, H.S., Galetti, M., Ceballos, G., Isaac, N.J.B. & Collen, B. (2014)
 Defaunation in the Anthropocene. *Science*, 345(6195), 401-406
- Dröge, E., Creel, S., Becker, M.S. & M'soka, J. (2017) Risky times and risky places interact
 to affect prey behaviour. *Nature Ecology & Evolution*, 1(8), 1123
- 708 Duncan, P., Tixier, H., Hofmann, R.R. & Lechner-Doll, M. (1998) Feeding strategies and the
- physiology of digestion in roe deer. The European roe deer: the biology of success (eds R.
- 710 Andersen, P. Duncan & J.D.C. Linnell), pp. 91-116. Scandinavian University Press, Oslo
- Eccard, J.A., Meißner, J.K. & Heurich, M. (2017) European roe deer increase vigilance when
 faced with immediate predation risk by Eurasian Lynx. *Ethology*, 123(1), 30-40
- Eriksen, A., Wabakken, P., Zimmermann, B., Andreassen, H.P., Arnemo, J.M., Gundersen,
- H., Liberg, O., Linnell, J.D.C., Milner, J.M., Pedersen, H.C., Sand, H., Solberg, E.J. & Storaas, T.
- 715 (2011) Activity patterns of predator and prey: a simultaneous study of GPS-collared wolves and
- 716 moose. Animal Behaviour, 81(2), 423-431
- Filla, M., Premier, J., Magg, N., Dupke, C., Khorozyan, I., Waltert, M., Bufka, L. & Heurich,
 M. (2017) Habitat selection by Eurasian lynx (*Lynx lynx*) is primarily driven by avoidance of
 human activity during day and prey availability during night. *Ecology and Evolution*, 7(16), 63676381
- Fortin, D., Beyer, H.L., Boyce, M.S., Smith, D.W., Duchesne, T. & Mao, J.S. (2005) Wolves
 influence elk movements: behavior shapes a trophic cascade in Yellowstone National Park. *Ecology*, 86(5), 1320–1330

- Frid, A. & Dill, L. (2002) Human-caused disturbance stimuli as a form of predation risk. *Conservation Ecology*, 6(1), 11
- Gaynor, K.M., Hojnowski, C.E., Carter, N.H. & Brashares, J.S. (2018) The influence of
 human disturbance on wildlife nocturnality. *Science*, 360(6394), 1232-1235
- Gehr, B., Hofer, E.J., Muff, S., Ryser, A., Vimercati, E., Vogt, K. & Keller, L.F. (2017) A
 landscape of coexistence for a large predator in a human dominated landscape. *Oikos*, 126(10),
 1389-1399.
- Gehr, B., Hofer, E.J., Pewsner, M., Ryser, A., Vimercati, E., Vogt, K. & Keller, L.F. (2018)
 Hunting-mediated predator facilitation and superadditive mortality in a European
 ungulate. *Ecology and Evolution*, 8(1), 109-119
- Gervasi, V., Brunberg, S. & Swenson, J. (2006) An individual-based method to measure
 animal activity levels: a test on brown bears. *Wildlife Society Bulletin*, 34, 1314–1319
- Godvik, I.M.R., Loe, L.E., Vik, J.O., Veiberg, V., Langvatn, R. & Mysterud, A. (2009)
- 737 Temporal scales, trade-offs, and functional responses in red deer habitat selection. *Ecology*, 90(3),
 738 699-710
- Graham, M.D., Douglas-Hamilton, I., Adams, W.M. & Lee, P.C. (2009) The movement of
 African elephants in a human-dominated land-use mosaic. *Animal Conservation*, 12(5), 445-455
- Heurich, M., Hilger, A., Küchenhoff, H., Andrén, H., Bufka, L., Krofel, M., Mattisson, J.,
- 742 Odden, J., Persson, J., Rauset, G.R., Schmidt, K. & Linnell, J.D.C (2014) Activity patterns of
- Eurasian lynx are modulated by light regime and individual traits over a wide latitudinal range.
- 744 *PLoS One*, 9(12), e114143

745	Hofmeester, T.R., Jansen, P.A., Wijnen, H.J., Coipan, E.C., Fonville, M., Prins, H.H., Sprong,
746	H. & van Wieren, S.E. (2017) Cascading effects of predator activity on tick-borne disease
747	risk. Proceedings of the Royal Society B: Biological Sciences, 284(1859), 20170453
748	Hoogenboom, I., Daan, S., Dallinga, J.H., & Schoenmakers, M. (1984) Seasonal change in
749	the daily timing of behaviour of the common vole, Microtus arvalis. Oecologia, 61(1), 18-31
750	Hofmann, R.R. (1989) Evolutionary steps of ecophysiological adaptation and diversification
751	of ruminants: a comparative view of their digestive system. Oecologia, 78(4), 443-457
752	Jędrzejewski, W., Schmidt, K., Theuerkauf, J., Jędrzejewska, B., Selva, N., Zub, K. &
753	Szymura, L. (2002) Kill rates and predation by wolves on ungulate populations in Białowieża
754	Primeval Forest (Poland). Ecology, 83(5), 1341-1356
755	Kamler, J.F., Jędrzejewska, B. & Jędrzejewski, W. (2007) Activity patterns of red deer in
756	Białowieża National Park, Poland. Journal of Mammalogy, 88(2), 508-514
757	Kays, R., Crofoot, M.C., Jetz, W. & Wikelski, M. (2015) Terrestrial animal tracking as an eye
758	on life and planet. Science, 348(6240), aaa2478
759	Kohl, M.T., Stahler, D.R., Metz, M.C., Forester, J.D., Kauffman, M.J., Varley, N., White,
760	P.J., Smith, D.W. & MacNulty, D.R. (2018) Diel predator activity drives a dynamic landscape of
761	fear. Ecological Monographs, 88(4), 638-652
762	Kohl, M.T., Ruth, T.K., Metz, M.C., Stahler, D.R., Smith, D.W., White, P.J. & MacNulty,
763	D.R. (2019) Do prey select for vacant hunting domains to minimize a multi-predator
764	threat? <i>Ecology Letters</i> , 00, 1–10, https://doi.org/10.1111/ele.13319

Komers, P.E. (1997) Behavioural plasticity in variable environments. *Canadian Journal of Zoology*, 75, 161-169

- 767 Krop-Benesch, A., Berger, A., Hofer, H. & Heurich, M. (2013) Long-term measurement of
- roe deer (*Capreolus capreolus*) (Mammalia: Cervidae) activity using two-axis accelerometers in

769 GPS-collars. *Italian Journal of Zoology*, 80(1), 69-81

- 770 Kröschel, M., Reineking, B., Werwie, F., Wildi, F. & Storch, I. (2017) Remote monitoring of
- vigilance behavior in large herbivores using acceleration data. *Animal Biotelemetry*, 5(1), 10
- 772 Kronfeld-Schor, N. & Dayan, T. (2003) Partitioning of time as an ecological resource. Annual

773 *Review of Ecology, Evolution and Systematics*, 34(1), 153-181

Kronfeld-Schor, N., Dayan, T., Elvert, R., Haim, A., Zisapel, N, & Heldmaier, G. (2001) On

the use of the time axis for ecological separation: diel rhythms as an evolutionary constraint. *The*

- 776 *American Naturalist*, 158(4), 451-457
- 777 Kronfeld-Schor, N., Visser, M.E., Salis, L. & van Gils, J.A. (2017) Chronobiology of
- interspecific interactions in a changing world. *Philosophical Transactions of the Royal Society B:*
- 779 Biological Sciences, 372(1734), 20160248
- Kusak, J., Skrbinšek, A.M. & Huber, D. (2005) Home ranges, movements, and activity of
 wolves (*Canis lupus*) in the Dalmatian part of Dinarids, Croatia. *European Journal of Wildlife Research*, 51(4), 254-262
- Laundré, J.W., Hernández, L. & Altendorf, K.B. (2001). Wolves, elk, and bison:
 reestablishing the "landscape of fear" in Yellowstone National Park, USA. *Canadian Journal of Zoology*, 79(8), 1401-1409
- Lehman, C.P., Rota, C.T., Raithel, J.D. & Millspaugh, J.J. (2018) Pumas affect elk dynamics
- in absence of other large carnivores. *The Journal of Wildlife Management*, 82(2), 344-353

- Levy, O., Dayan, T., Porter, W.P. & Kronfeld-Schor, N. (2019) Time and ecological
 resilience: can diurnal animals compensate for climate change by shifting to nocturnal activity? *Ecological Monographs*, 89(1), e01334
- Lima, S.L. & Dill, L.M. (1990) Behavioral decisions made under the risk of predation: a
 review and prospectus. *Canadian Journal of Zoology*, 68(4), 619-640
- Loe, L.E., Bonenfant, C., Mysterud, A., Severinsen, T., Øritsland, N.A., Langvatn, R., Stien,

A., Irvine, R.J. & Stenseth, N.C. (2007) Activity pattern of arctic reindeer in a predator-free
environment: no need to keep a diel rhythm. *Oecologia*, 152, 617–624

- Lone, K., Loe, L.E., Gobakken, T., Linnell, J.D.C., Odden, J., Remmen, J. & Mysterud, A.
- 797 (2014) Living and dying in a multi-predator landscape of fear: roe deer are squeezed by contrasting

pattern of predation risk imposed by lynx and humans. Oikos, 123(6), 641-651

Lone, K., Mysterud, A., Gobakken, T., Odden, J., Linnell, J.D.C. & Loe, L.E. (2017)

Temporal variation in habitat selection breaks the catch-22 of spatially contrasting predation risk
from multiple predators. *Oikos*, 126(5), 624-632

- Long, E.S., Jacobsen, T.C., Nelson, B.J. & Steensma, K.M.M. (2013) Conditional diel and seasonal movement strategies of male Columbia black-tailed deer (*Odocoileus hemionus columbianus*). *Canadian Journal of Zoology*, 91(10), 679–688
- Massé, A. & Côté, S.D. (2013) Spatiotemporal variations in resources affect activity and
 movement patterns of white-tailed deer (*Odocoileus virginianus*) at high density. *Canadian Journal of Zoology*, 91(4), 252–263
- Manning, A.D., Gordon, I.J. & Ripple, W.J. (2009) Restoring landscapes of fear with wolves
 in the Scottish Highlands. *Biological Conservation*, 142(10), 2314-2321

- Mejlon, H.A. (1997). Diel activity of *Ixodes ricinus* Acari: Ixodidae at two locations near
 Stockholm, Sweden. *Experimental & Applied Acarology*, 21(4), 247-256
- Monterroso, P., Alves, P.C. & Ferreras, P. (2013) Catch me if you can: diel activity patterns
 of mammalian prey and predators. *Ethology*, 119, 1044-1056
- 814 Murray, M.H. & St. Clair, C.C. (2015) Individual flexibility in nocturnal activity reduces risk
- of road mortality for an urban carnivore. *Behavioral Ecology*, 26(6), 1520-1527

816 Nilsen, E.B., Gaillard, J.-M., Andersen, R., Odden, J., Delorme, D., Van Laere, G. & Linnell,

J.D.C. (2009) A slow life in hell or a fast life in heaven: demographic analyses of contrasting roe

- deer populations. *Journal of Animal Ecology*, 78(3), 585-594
- Nouvellet, P., Rasmussen, G.S.A., Macdonald, D.W. & Courchamp, F. (2012) Noisy clocks
 and silent sunrises: measurement methods of daily activity pattern. *Journal of Zoology*, 286(3),
 179-184
- Oriol-Cotterill, A., Valeix, M., Frank, L.G., Riginos, C., & Macdonald, D.W. (2015) Landscapes of coexistence for terrestrial carnivores: the ecological consequences of being downgraded from ultimate to penultimate predator by humans. *Oikos*, 124(10), 1263-1273
- Pagon, N., Grignolio, S., Pipia, A., Bongi, P., Bertolucci, C. & Apollonio, M. (2013) Seasonal
 variation of activity patterns in roe deer in a temperate forested area. *Chronobiology International*, 30(6), 772-785
- Picardi, S., Basille, M., Peters, W., Ponciano, J. M., Boitani, L. & Cagnacci, F. (2019)
 Movement responses of roe deer to hunting risk. *The Journal of Wildlife Management*, 83(1), 4351
- Preisser, E.L., Bolnick, D.I. & Benard, M.F. (2005) Scared to death? The effects of
 intimidation and consumption in predator–prey interactions. *Ecology*, 86(2), 501-509

833	Preisser, E.L., Orrock, J.L. & Schmitz, O.J. (2007) Predator hunting mode and habitat domain
834	alter nonconsumptive effects in predator-prey interactions. Ecology, 88(11), 2744-2751
835	R Core Team. (2017) R: a language and environment for statistical computing. R Foundation
836	for Statistical Computing, Vienna, Austria. Available from https://www.r-project.org/
837	Reebs, S.G. (2002). Plasticity of diel and circadian activity rhythms in fishes. Reviews in Fish
838	Biology and Fisheries, 12(4), 349-371
839	Roberts, C.P., Cain III, J.W. & Cox, R.D. (2017) Identifying ecologically relevant scales of
840	habitat selection: diel habitat selection in elk. Ecosphere, 8(11), e02013
841	Ridout, M. & Linkie M. (2009) Estimating overlap of daily activity patterns from camera trap
842	data. Journal of Agricultural, Biological and Environmental Statistics, 14(3), 322-337
843	Sand, H., Zimmermann, B., Wabakken, P., Andrèn, H. & Pedersen, H.C. (2005) Using GPS
844	technology and GIS cluster analyses to estimate kill rates in wolf-ungulate ecosystems. Wildlife
845	Society Bulletin, 33(3), 914-925
846	Schmidt, K. (1999). Variation in daily activity of the free-living Eurasian lynx (Lynx lynx) in
847	Białowieża Primeval Forest, Poland. Journal of Zoology, 249(4), 417-425
848	Shamoon, H., Maor, R., Saltz, D. & Dayan, T. (2018) Increased mammal nocturnality in
849	agricultural landscapes results in fragmentation due to cascading effects. Biological Conservation,
850	226, 32-41
851	Sih, A., Ferrari, M.C. & Harris, D.J. (2011) Evolution and behavioural responses to human-
852	induced rapid environmental change. Evolutionary Applications, 4(2), 367-387
853	Smith, J.A., Wang, Y. & Wilmers, C.C. (2015) Top carnivores increase their kill rates on prey
854	as a response to human-induced fear. Proceedings of the Royal Society B: Biological
855	Sciences, 282(1802), 20142711

- Sönnichsen, L., Bokje, M., Marchal, J., Hofer, H., Jędrzejewska, B., Kramer-Schadt, S. &
 Ortmann, S. (2013). Behavioural responses of European roe deer to temporal variation in predation
 risk. *Ethology*, 119(3), 233-243
- 859 Stillfried, M., Belant, J.L., Svoboda, N.J., Beyer, D.E. & Kramer-Schadt, S. (2015) When top
- 860 predators become prey: black bears alter movement behaviour in response to hunting pressure.
- 861 Behavioural Processes, 120, 30-39
- 862 Swinnen, K.R.R., Hughes, N.K. & Leirs, H. (2015) Beaver (*Castor fiber*) activity patterns in
- a predator-free landscape: what is keeping them in the dark? *Mammalian Biology* 80, 477-483
- Tambling, C.J., Minnie, L., Meyer, J., Freeman, E.W., Santymire, R.M., Adendorff, J. &
- Kerley, G.I. (2015) Temporal shifts in activity of prey following large predator reintroductions. *Behavioral Ecology and Sociobiology*, 69(7), 1153-1161
- 867 Tucker, M.A., Böhning-Gaese, K., Fagan, W.F., Fryxell, J.M., Van Moorter, B., Alberts, S.C.,
- Ali, A.H., Allen, A.M., Attias, N., Avgar, T., Bartlam-Brooks, H., Bayarbaatar, B., Belant, J.L.,
- Bertassoni, A., Beyer, D., Bidner, L., van Beest, F.M., Blake, S., Blaum, N., Bracis, C., Brown,
- 870 D., de Bruyn, P.J.N., Cagnacci, F., Calabrese, J.M., Camilo-Alves, C., Chamaillé-Jammes, S.,
- 871 Chiaradia, A., Davidson, S.C., Dennis, T., DeStefano, S., Diefenbach, D., Douglas-Hamilton, I.,
- 872 Fennessy, J., Fichtel, C., Fiedler, W., Fischer, C., Fischhoff, I., Fleming, C.H., Ford, A.T., Fritz,
- 873 S.A., Gehr, B., Goheen, J.R., Gurarie, E., Hebblewhite, M., Heurich, M., Hewison, A.J.M., Hof,
- 874 C., Hurme, E., Isbell, L.A., Janssen, R., Jeltsch, F., Kaczensky, P., Kane, A., Kappeler, P.M.,
- 875 Kauffman, M., Kays, R., Kimuyu, D., Koch, F., Kranstauber, B., LaPoint, S., Leimgruber, P.,
- 876 Linnell, J.D.C., López-López, P., Markham, A.C., Mattisson, J., Medici, E.P., Mellone, U.,
- 877 Merrill, E., de Miranda Mourão, G., Morato, R.G., Morellet, N, Morrison, T.A., Díaz-Muñoz, S.L.,
- 878 Mysterud, A., Nandintsetseg, D., Nathan, R., Niamir, A., Odden, J., O'Hara, R.B., Oliveira-Santos,

- 879 L.G.R., Olson, K.A., Patterson, B.D., de Paula, R.C., Pedrotti, L., Reineking, B., Rimmler, M.,
- 880 Rogers, T.L., Rolandsen, C.M., Rosenberry, C.S., Rubenstein, D.I., Safi, K., Saïd, S., Sapir, N.,
- 881 Sawyer, H., Schmidt, N.M., Selva, N., Sergiel, A., Shiilegdamba, E., Silva, J.P., Singh, N.,
- 882 Solberg, E.J., Spiegel, O., Strand, O., Sundaresan, S., Ullmann, W., Voigt, U., Wall, J., Wattles,
- B83 D., Wikelski, M., Wilmers, C.C., Wilson, J.W., Wittemyer, G., Zięba, F., Zwijacz-Kozica, T.,
- 884 Mueller, T. (2018) Moving in the Anthropocene: Global reductions in terrestrial mammalian
- 885 movements. *Science*, 359(6374), 466-469
- Tolon, V., Dray, S., Loison, A., Zeileis, A., Fischer, C. & Baubet, E. (2009) Responding to
- spatial and temporal variations in predation risk: space use of a game species in a changing
- landscape of fear. *Canadian Journal of Zoology*, 87(12), 1129-1137
- 889 Venter, O., Sanderson, E.W., Magrach, A., Allan, J.R., Beher, J., Jones, K.R., Possingham,
- 890 H.P., Laurance, W.F., Wood, P., Fekete, B.M., Levy, M.A. & Watson, J.E.M. (2016) Global
- terrestrial Human Footprint maps for 1993 and 2009. Scientific Data, 3, 160067
- Williams, H.J., Holton, M.D., Shepard, E.L., Largey, N., Norman, B., Ryan, P.G., Duriez, O.,
- 893 Scantlebury, M., Quintana, F., Magowan, E.A., Marks, N.J., Alagaili, A.N., Bennett, N.C. &
- Wilson, R.P. (2017) Identification of animal movement patterns using tri-axial magnetometry. *Movement ecology*, 5(1), 6
- 896

897 Supporting information provided:

Appendix S1. Classification of the activity data into active *vs* inactive behaviour for
describing diel activity patterns
Appendix S2. Graphical representation of the diel activity patterns of roe deer over a 24-h
cycle and overlap between the non-hunting and hunting seasons

- 902 Appendix S3. Summary and results of the top-ranked candidate models explaining variation
 903 in the level of diurnality (D)
- 904 Appendix S4. Summary and results of the top-ranked candidate models explaining variation
- 905 in the level of crepuscularity (C)

906