Human recreation affects spatio-temporal habitat use patterns in red deer (Cervus elaphus)

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Abstract

The rapid spread and diversification of outdoor recreation can impact on wildlife in various ways, often leading to the avoidance of disturbed habitats. To mitigate human-wildlife conflicts, spatial zonation schemes can be implemented to separate human activities from key wildlife habitats, e.g., by designating undisturbed wildlife refuges or areas with some level of restriction to human recreation and land use. However, mitigation practice rarely considers temporal differences in human-wildlife interactions. We used GPS telemetry data from 15 red deer to study the seasonal (winter vs. summer) and diurnal (day vs. night) variation in recreation effects on habitat use in a study region in south-western Germany where a spatial zonation scheme has been established. Our study aimed to determine if recreation infrastructure and spatial zonation affected red deer habitat use and whether these effects varied daily or seasonally. Recreation infrastructure did not affect home range selection in the study area, but strongly determined habitat use within the home range. The spatial zonation scheme was reflected in both of these two levels of habitat selection, with refuges and core areas being more frequently used than the border zones. Habitat use differed significantly between day and night in both seasons. Both summer and winter recreation trails, and nearby foraging habitats, were avoided during day, whereas a positive association was found during night. We conclude that human recreation has an effect on red deer habitat use, and when designing mitigation measures daily and seasonal variation in human-wildlife interactions should be taken into account. We advocate using spatial zonation in conjunction with temporal restrictions (i.e., banning nocturnal recreation activities) and the creation of suitable foraging habitats away from recreation trails.

Introduction

An increasing number of people are practicing nature-based tourism, with outdoor recreation activities generating pressure on the ecosystems in which they take place [1, 2]. An important
factor of how outdoor recreation affects ecosystems is the disturbance of wildlife by human recreation [3, 4], defined here as any effect on wildlife which is incurred by the presence of recreationists or infrastructure related to recreational activities, irrespective of possible—but mostly unknown—fitness consequences [5, 6]. Free-living animals often react to human presence in a similar way than to the presence of natural predators [7, 8]. This reaction can have a variety of facets [9, 10], ranging from physiological stress responses [11–14] to behavioural changes [8, 15] or a reduction in reproductive success [8]. Human disturbance might trigger short-term behavioural reactions (i.e. flushing or fleeing) [1, 16, 17] as well as long-term responses such as avoiding frequently disturbed areas [18, 19], e.g. recreational infrastructures such as hiking or skiing trails that are regularly used by humans [20]. Both types of reaction can involve direct energetic costs for the animal (e.g. due to fleeing or reduced food intake) which can affect fitness [14, 21, 22], and may even outweigh the effects of habitat conditions and natural predators [23].

In addition, the reactions of wildlife triggered by human recreational activities can cause conflicts with other forms of human land use, such as transportation, agriculture or forestry. For example, fleeing animals can trigger vehicle collisions [24], and foraging animals relocating to less disturbed areas might cause damage to crop or tree regeneration [25, 26].

To mitigate both the negative effects of human recreation on wildlife and the resulting conflicts with land use, spatial zonation schemes have become an important tool in wildlife management [27]. These schemes separate human activities from key wildlife habitats by designating undisturbed wildlife refuges and areas with different levels of restriction to human recreation, sometimes combined with habitat management or hunting regulations. The design of zonation schemes often takes spatial patterns of human-wildlife interactions into account but rarely considers temporal interactions, e.g. variation in diurnal and seasonal overlaps between habitat requirements and recreation activities. Using the red deer (Cervus elaphus) as an example organism, we investigated the temporal variation of its habitat use in relation to human recreation infrastructure and zones with different intensities of human disturbance, as established by a zonation scheme.

As one of the largest free ranging herbivores, and widely distributed across the globe [28, 29], the red deer is one of the focal species of wildlife management in Central Europe [30, 31]. Red deer are attractive to observe and are therefore highly valued for nature-based tourism [30] and also as a game species [28, 29]. They are considered an important vector species for seeds [32, 33] and invertebrates [34] and an essential prey for carnivores (e.g. wolf) [35]. With its browsing behaviour it can affect the vegetation structure [36–38] and there is some evidence for impacts on plant species richness [39]. At the same time however, deer browsing and bark stripping causes conflicts with forestry management [31, 40, 41]. In addition, thousands of individuals are injured or die in vehicle collisions every year, causing considerable property damage and fatal human injuries [42, 43]. The major objective of the red deer management in Europe is therefore to minimize the economic damage related to forestry and animal vehicle collisions while maximizing the economic benefits related to ecosystem services and hunting [44–46].

Outdoor recreation has been widely neglected within the management of free ranging ungulates [1], although red deer have been shown to be influenced by human recreationists [20, 26, 47, 48]. Direct reactions to disturbance include instant flight, relocation to areas with dense vegetation cover [26, 49] as well as a temporal abandonment of the disturbed area (i.e. for several hours or days) [26]. Sibbald et al. [20] found red deer avoided hiking trails, with larger distance to the trail kept during times of high human use compared to times of little use. In areas with high recreation pressure, red deer have been shown to increase their vigilance behaviour which might lead to a decrease in food uptake [47]. Animals can also adjust their
habitat use between hunting season and non-hunting season [50], which indicates their behav-
ioral plasticity. However, even though there are several examples where north American elk
(*Cervus elaphus canadensis*) have become habituated to human presence [51] and even use set-
tlements as habitat [52], this phenomenon is not known from free-ranging European red deer
[53, 54]. Human disturbance may therefore cause red deer to temporally or permanently aban-
don optimal habitat and forage in sub-optimal habitats [48]. Increased energy requirements
caused by fleeing, in conjunction with seeking cover [26] could result in damage to forestry
e.g. through bark-stripping in young, dense stands offering visual protection.

The most widely applied method of red deer management involves hunting, to regulate the
population and to gain trophies (antlers) and meat [28, 29]. Furthermore in many areas red
deer are provided with supplementary food during winter to reduce bark-stripping or because
of animal welfare reasons [55]. In the last decades, wildlife refuges have increasingly been des-
ignated, with the primary aim to reduce disturbance of deer by recreationists, land use man-
agement and hunters [25, 56, 57]. However, it has also been suggested that well-placed refuges
may help reduce human-wildlife conflicts [58] and contribute to decreasing damage to forestry
by reducing the browsing pressure on the surrounding forest stands [25]. To serve this pur-
pose, refuge systems have been extended to spatial zonation schemes that regulate recreational
activities but also hunting and forest management [27]. However, management schemes aim-
ing at furthering the coexistence of humans and wildlife must also consider temporal dimen-
sions of human-wildlife interactions [18, 59]. Given the seasonal and diurnal differences in
recreation activities and the behavioural plasticity of red deer, we expect that the spatial pattern
of habitat use in relation to human recreation infrastructure varies considerably between sea-
sons and between day and nighttime, which might also modify the relative importance (i.e.
intensity of use) of the zones of a static spatial zonation. To test this, we studied the habitat use
of free roaming red deer comparing daytime and nighttime activity in two different seasons
using GPS-telemetry. The study was conducted in a red deer management area in south-west-
ern Germany in which a spatial zonation scheme had been established, defining red deer ref-
uges (without human recreation), a core zone with limited recreational use and a border zone
with unrestricted recreation. The goals of our study were to determine if linear recreation
infrastructure (i.e. hiking, biking and skiing trails) and the zonation scheme affected red deer
habitat use and whether these effects varied daily or seasonally. From the results we derive rec-
ommendations for mitigating impacts of human recreation on ungulates in human-dominated
landscapes.

**Methods**

**Ethics statement**

Red deer capturing and tagging was carried out under the permit (No. 787.524) issued by the
ethical committee of the Regional Council of Freiburg, Baden-Württemberg (Regierungspräsi-
dium Freiburg, Baden-Württemberg). The ethical committee specifically approved this study.
GPS collars were attached under anesthesia (125 mg Xylazine + 100 mg Ketamine /ml).

**Study area**

The study was conducted in the Southern Black Forest, Baden-Württemberg, south-western
Germany (Fig 1). In the state of Baden-Württemberg it is official policy to try to keep red deer
in five specially designated areas, which are mainly state owned, to avoid conflicts with private
forest owners and farmers. Red deer leaving the management areas are shot at sight. The
Southern Black Forest red deer management area has a total surface of 17500 ha; our study
was performed in the central part of 5984 ha, located at elevations between 800 and 1300m
above sea level (a.s.l.). Most of the study area (77%) consists of intensively managed forest (for timber production) dominated by Norway spruce (*Picea abies*), European silver fir (*Abies alba*) and common beech (*Fagus sylvatica*) [60]. Extensively managed meadows prevail in the non-forested areas.

The study area is located between two major tourist attractions Lake Schluchsee and the Feldberg Mountain and is intensively and increasingly used for recreation all year round. This is reflected in a 24% increase of tourist visits to the region between 2004 and 2014 [61]. In the study area, a dense network of recreation trails has been established: in summer, a total of 162 km (2.71 km/km²) of paths are accessible, mainly for hiking and biking. During winter, trails for hiking (48 km; 0.8km/km²) and cross-country skiing (39 km; 0.65km/km²) are prepared, but there are also off-trail recreational activities (i.e. snowshoeing and back-country skiing) [62]. To assess the temporal patterns of human recreationists in the areas, automatic visitor counts were performed on hiking and skiing trails one year after the data collection for the telemetry study. Infrared trail counters (TRAFx), were placed along three designated hiking trails and three cross country skiing trails within the study area from 17.2.2010 to 14.4.2010,
showing a peak of recreation activities at noon, with an average number of six persons per hour recorded on hiking trails and two per hour on skiing trails (S2 Fig).

During the duration of our study (2007–2009), 600–700 free-roaming red deer were estimated to be present in winter within the total red deer management area (Forest Research Institute of Baden-Württemberg FVA, unpublished), which corresponded to a density of 3.43–4.00 individuals per km$^2$. Other ungulates present are roe deer (*Capreolus capreolus*), wild boar (*Sus scrofa*) and sika deer (*Cervus nippon*) (FVA, unpublished). Predators include red fox (*Vulpes vulpes*) and pine marten (*Martes martes*), but with lynx (*Lynx lynx*) and grey wolf (*Canis lupus*) absent, adult red deer have no natural predators in the area (FVA, unpublished).

### Zoning scheme

Beginning in 2003, a spatial zonation scheme was developed and implemented using a joint participative process, which included wildlife biologists, foresters, hunters and landowners. It was officially approved in 2008 by the local communities. The scheme includes different zones with regulations concerning not only recreational use, but also hunting, forestry and red deer habitat management (Fig 1, Table 1) [56]: A border zone—where no restrictions for recreation apply—surrounds a core zone where recreation is restricted to designated trails. Embedded within the core zone, refuge areas for red deer have been designated, where recreational use is totally banned. During winter the deer are fed at four feeding stations to minimize seasonal migration and thus reduce deer-vehicle collisions, but the animals are not fenced during any time of the year [56].

In the study area the hunting times are more restricted compared to the official state hunting regulations. In the border zone, hunting is only allowed from the 1st of August until 31st of December (i.e. banned in the summer hunting season between May and July). In the core zone hunting activities are additionally banned in December and restricted to interval hunting (i.e. short hunting intervals followed by several days without hunting with the goal to reduce disturbance). In the refuge areas hunting is restricted to driven hunts in three consecutive weeks in October. The aim of the hunting regime in the area is to limit the red deer population size (winter) to an overall number of 400 individuals (2.29 individuals per km$^2$).

In all zones, forestry is directed towards creating small openings during timber harvesting, to increase natural food resources for the deer. In the border zone, moderate protection measures, such as small scale fencing, can be implemented to avoid damage to forestry caused by deer. In the core zone, damage caused by deer to forestry is accepted. In the refuges and feeding stations no forestry measures (i.e. timber harvesting) are performed during the fawning season (i.e. May and June) and feeding times (i.e. snow conditions). The main goals of this zonation scheme are decreasing damage to forestry across the whole area by allowing the deer to retreat to undisturbed areas with sufficient food in summer and additional feeding in winter, while at the same time creating possibilities for human recreation which includes the possibility to observe and experience red deer [56].

### Red deer data

Our analysis was based on telemetry locations of 15 red deer (5 males and 10 females, all age classes, S1 Table) captured and surveyed between 2007 and 2009. Individuals were equipped with a GPS-collar (Vectronic Aerospace, Berlin, Germany; serial number 2000er, 3000er and 6000er) and located every 2 hours. The tracking period of individual animals ranged between 5 and 34 months (S1 Table), depending on the functional duration of the GPS collars and due to individual fatality events. We retained only locations if a minimum of 4 satellites were
available and the DOP (dilution of precision) value was smaller than 10 (corresponding to an estimated maximum location error of about 40 m [63]), resulting in 80% of the locations for further analysis. To model temporal differences in habitat use, each sample was allocated to a season (summer, winter) and a time of day (day, night). Since seasonal differences in habitat use patterns were assumed to be related to prevailing weather conditions rather than being determined by a predefined time period, seasons were defined using standard indicators of weather conditions: The “summer” season started with the flowering of dandelion (Taraxacum officinale) (18, 22 and 30 April in 2007, 2008 and 2009, respectively), as measured at the phenology reference station Bernau, 920 m.a.s.l., and ended with the start of the rutting season (15 September, all years). The “winter”-sample contained all locations taken between first of November and the beginning of the summer season in the following year, including only days where a continuous snow-layer was recorded at the nearby weather station (St. Blasien-Menzenschwand, 885 m.a.s.l.). Locations taken outside the defined seasons were discarded.

Among the retained locations, we distinguished between day and night. Day was defined as the time between sunrise and sunset, and night covered the time between the end of the nautical twilight. Due to failing fixes or fixes with too high DOP (i.e. low precision), the number of locations per day and time period varied greatly within and between individuals. To avoid an unbalanced sample, (i.e. some time periods being overrepresented by data showing high spatial and temporal autocorrelation) we adopted a conservative approach, randomly selecting only one location per time period and day for every individual.

Of the resulting 24259 locations which were retained for further analysis (S1 Table), 7384 locations pertained to summer, and 16875 to the winter season. The number of locations per individual varied between 244 and 3136 (S1 Table).

### Environmental variables

We distinguished three groups of environmental predictors, pertaining to land cover and topography, vegetation structure and human presence (Table 2). Topographic variables (altitude, slope and exposition) were calculated from the digital elevation model (DEM). Land cover characteristics (waterbodies, meadows, forest) were adopted from the Official Topographic and Cartographic Information System of Germany (ATKIS, www.atkis.de).

Vegetation was mapped in the field: forest stand type, canopy cover, tree-species mixture, successional stage, understory composition, cover of herbs and grass as well as bilberry cover (Vaccinium myrtillus) was recorded for forest stand units, which represent homogeneously structured patches with a mean size of 3.40 ha (min: 0.20, max: 48.20). The variable “visual protection” in summer and winter was recorded in a location where understory conditions were considered representative for the respective forest stand. Using a “chessboard”
(100x100cm) with a black and white grid (i.e. 100 10x10cm squares), placed upright at a distance of 30 m in all four cardinal directions from the observer, the amount of visual protection was then derived from the number of squares that were hidden by the vegetation. Covering an area of 10% of the average stand size, this measurement provides a rough estimation of the

Table 2. Predictor variables included in the models.

| Predictor type          | Variable Name | Description (unit)                                      | Min-Max | Type       |
|------------------------|---------------|--------------------------------------------------------|---------|------------|
| Landscape and          | D2H           | Altitude (m a.s.l.)                                    | 762–1314| continuous |
| topography             | SLOPE         | Slope (degree)                                         | 0–44    | continuous |
|                        | NORTHING      | Northness (cosine aspect)                              | -1–1    | continuous |
|                        | EASTING       | Eastness (sine aspect)                                 | -1–1    | continuous |
|                        | WATER         | Proximity to lakes, rivers and creeks (km)             | 0.005–0.704| continuous |
|                        | GREENL        | Proximity to greenland (i.e. meadows/ grassland) (km)  | 0–1.343 | continuous |
|                        | FOREST_250    | Forest cover within a 250m radius (%)                  | 0–100   | continuous |
| Vegetation             | CANOPY_TYPE   | Type of canopy trees                                   |         | categorical|
|                        | CAN_NO        | No forest (reference category)                         |         |            |
|                        | CAN_CON       | Coniferous >95%                                        |         |            |
|                        | CAN_CONMIX    | Conifer dominated mixed (conifers >50%)                |         |            |
|                        | CAN_DEC_MIX   | Deciduous dominated mixed (deciduous >50%)             |         |            |
|                        | CAN_DEC       | Deciduous >95%                                         |         |            |
|                        | CANOPY_COV    | Canopy cover (%)                                       | 0–100   | continuous |
|                        | SUC_OPEN      | Open (reference category)                              |         |            |
|                        | SUC_REGTHICK  | Regeneration & Thicket                                 |         |            |
|                        | SUC_POLE      | Pole stage                                             |         |            |
|                        | SUC_TREE      | Tree stage                                             |         |            |
|                        | SUC_OLD       | Old forest                                             |         |            |
|                        | UNDER_TYPE    | Type of understory trees                               |         | categorical|
|                        | UNDER_NON     | No understory (reference category)                     |         |            |
|                        | UNDER_CON     | Coniferous >95%                                        |         |            |
|                        | UNDER_DEC     | Deciduous >95%                                         |         |            |
|                        | UNDER_DEC MIX | Deciduous dominated mixed (deciduous >50%)             |         |            |
|                        | UNDER_CONMIX  | Conifer dominated mixed (conifers >50%)                |         |            |
|                        | UNDER_COV     | Cover of understory (%)                                | 0–90    | continuous |
|                        | BILBERRY      | Bilberry cover (%)                                     | 0–90    | continuous |
|                        | HERB_GRAS     | Cover of herbs and grass (%)                           | 0–100   | continuous |
|                        | PROTECTION_S/W| Protection from visibility in summer/winter (%)        | 0–75    | continuous |
| Human presence         | TOURI_S/W     | Proximity to summer tourism infrastructure in summer/winter (km) | S: 0–0.752| continuous |
|                        |               |                                                       | W: 0–1.824|            |
|                        | TOURI_DENS_S/W| Density of summer/winter tourism infrastructure within 250m (m/ha) | S: 0–129| continuous |
|                        |               |                                                       | W: 0–108|            |
|                        | ROAD          | Proximity to roads (km)                                | 0.006–2.321| continuous |
|                        | SETTLE        | Proximity to settlements (km)                          | 0–3.105 | continuous |
|                        | FEED          | Proximity to feeding stations (km)                     | 0–6.300 | continuous |
|                        | HUNT          | Proximity to hunter hides (km)                         | 0–3.918 | continuous |
|                        | MGT           | Different area-types of the red-deer management scheme (Table 1) |         | categorical|
|                        |               | MGT_BORDER = Border zone (reference category)          |         |            |
|                        |               | MGT_CORE = Core area                                   |         |            |
|                        |               | MGT_REFUGE = Refuge area                                |         |            |

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possibility for red deer to hide. As vegetation mapping was done in summer, protection in winter was estimated based on the understory type and density, i.e. subtracting the cover provided by broadleaved trees and bushes.

Human infrastructure (roads, settlements) was accessed from the Official Topographic and Cartographic Information System of Germany (ATKIS, www.atkis.de). In addition, we mapped tourism infrastructure in summer (hiking trails, mountain bike routes) and winter (cross country skiing and snowshoe trails, winter hiking paths), the location of the red-deer feeding stations in winter and the different zones of the zoning scheme. For all predictors we prepared raster maps with a 10 x 10 m resolution. To account for potential radio tracking errors, we performed a circular moving window analysis with a radius of 40m (corresponding to the maximum location error), assigning to the focal cell the mean value or, in case of categorical variables, the category that was most frequently present within the window. Variable maps were processed in ArcGIS 9.3 (ESRI 2009).

**Statistical approach**

To analyze habitat use we adopted a ‘used versus available’ design at two spatial habitat scales, comparing the presence data with two sets of random locations: First, to determine the factors influencing home range selection within the study area (second order habitat selection [64]), the presence locations of each individual were contrasted against the same number of random locations generated throughout the study area. Second, to analyze habitat selection within the home range (third order habitat selection [64]), we generated a second random sample selected from the individuals’ seasonal home ranges. Home ranges were calculated for each year and season separately, using the full data set (i.e. all available locations of the individual for the season) and the 100% minimum convex polygon (MCP) method. Habitat use was analyzed using Generalized Linear Mixed Effects Models (GLMM, R-package: lme4 [65]) with a logit link and binomial error structure, including the individual as a random factor. First, starting with the initial set of variables (Table 2), we identified pairs of strongly correlated variables (Spearman’s R > |0.5|), discarding the variable that explained less within a univariate model. Multivariate models with all possible combinations of the remaining variables were then fitted using the dredge function (R-package MuMin, [66]) in order to find the most parsimonious model according to Akaikes Information Criterion (AIC) [67]. Model averaging was applied if several “best models” did not differ significantly (ΔAIC < 2). For each season we fitted three models describing (1) home range selection within the study area (day and night pooled), as well as habitat use within the home range during (2) day and (3) night. In addition, we tested for differences in habitat use between day and night, identifying the environmental predictors that significantly discriminated the individual’s locations taken at the two different time periods. To assess multicollinearity in the final models (i.e. whether linear combinations of the fixed effects were correlated), we calculated the variance inflation factor (VIF) for all models, using the corvif function in the R-package AED [68]. For continuous variables we accepted an VIF of less than 10 [69], for factor variables the VIF was corrected for the number of degrees of freedom (VIF^2/(1/df)) [68]. The importance of individual variables was evaluated by fitting the final models while leaving out the respective variable. The change in AIC (ΔAIC) compared to the final model was then used as an indicator of the variable’s relative contribution to the final model. In addition, for every independent variable we calculated the odds-ratio and its 95% confidence interval using the Wald chi-square test [70] to approximate its effect on the dependent variable [70]. Model performance was evaluated using the area under the receiver operating characteristic (ROC) curve (AUC, R-package: AUCmodavg [71]). All statistical analyses were performed using the software R (R Version 2.15.1, www.rproject.org).
Results

Home range location

According to the classification of Hosmer and Lemeshow [72] our models performed well in explaining home range selection within the study area during both summer (AUC: 0.766 ± 0.003) and winter (0.919 ± 0.002). Home range selection was explained by variables describing landscape, vegetation structure and human presence (Table 3). According to the ΔAIC (Table 3), the zonation scheme was the strongest predictor for home range selection in both seasons: in summer the refuge areas were selected over the core area and the border zone, which served as a reference category (Table 3A), whereas winter home range selection was mainly located close to the feeding stations. The effect of human infrastructure differed between seasons: whereas in summer human settlements were avoided, winter home ranges were selected in closer vicinity to settlements and roads than expected from a random selection. The proximity and density of recreation infrastructure had no significant effect on home

Table 3. Selection of home range in study area.

| Type          | Variable     | Estimate | SE  | Sign | ΔAIC  | Estimate | SE  | Sign | ΔAIC |
|---------------|--------------|----------|-----|------|-------|----------|-----|------|------|
| Vegetation    | SUC_REGTHICK | 0.449    | 0.277 | 71   |       | 2.209    | 0.245 | ***  | 227  |
|               | SUC_POLE     | -0.083   | 0.270 |      |       | 1.026    | 0.210 | ***  |       |
|               | SUC_TREE     | 0.031    | 0.268 |      |       | 1.771    | 0.190 | ***  |       |
|               | SUC_OLD      | -0.200   | 0.272 |      |       | 1.898    | 0.198 | ***  |       |
|               | PROTECT_S/W  | <0.001   | 0.001 | 2    |       | -0.010   | 0.002 | ***  | 31   |
|               | BILBERRY     | -0.008   | 0.002 | ***  | 23    |          |      |      |      |
|               | CAN_CON      | -0.981   | 0.268 | ***  | 45    |          |      |      |      |
|               | CAN_DEC      | 0.404    | 0.413 |      |       |          |      |      |      |
|               | CAN_CONMIX   | -0.874   | 0.269 | **   |       |          |      |      |      |
|               | CAN_DEC MIX  | -0.761   | 0.276 | *    |       |          |      |      |      |
|               | HERB_GRAS    | 0.018    | 0.001 | ***  | 498   |          |      |      |      |
|               | CANOPY_COVER | -0.011   | 0.003 | ***  | 16    |          |      |      |      |
| Landscape     | WATER        | 1.146    | 0.171 | ***  | 48    | -0.856   | 0.286 | ***  | 9    |
|               | FOREST250    | 0.536    | 0.106 | ***  | 32    |          |      |      |      |
|               | SLOPE        | -0.004   | 0.003 | 4    |       |          |      |      |      |
| Human         | NORTHING     | -0.270   | 0.046 | ***  | 36    |          |      |      |      |
|               | EASTING      | -1.054   | 0.054 | ***  | 376   |          |      |      |      |
|               | MGT_CORE     | 1.930    | 0.664 | ***  | 1538  | -0.498   | 0.090 | ***  | 133  |
|               | MGT_REFUGE   | 2.723    | 0.077 | ***  | 529   | 0.125    | 0.67  | 67   |
|               | FEED         |          |      |      |       | 2.055    | 0.057 | ***  | 2098 |
|               | HUNT         | 1.762    | 0.051 | ***  | 1527  | 1.896    | 0.092 | ***  | 544  |
|               | SETTLE       | -0.347   | 0.028 | ***  | 137   | 0.368    | 0.068 | ***  | 67   |
|               | TOURI_S/W    | 0.147    | 0.132 | 2    |       |          |      |      |      |
|               | ROAD         | 0.684    | 0.102 | ***  | 31    |          |      |      |      |

Variables determining the home range selection of red deer within the study area in (a) summer and (b) winter. For all variables positive estimates indicate preference, negative estimates indicate avoidance. For predictor names see Table 2. Significance levels are indicated with: * p ≤ 0.05, ** p ≤ 0.01, and *** p ≤ 0.001. Relative variable importance is indicated by ΔAIC, which is the difference in AIC of a model discarding the respective variable compared to the full model.

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range selection. During the summer months deer home ranges were located in forest areas with a high proportion of openings and thickets rich in herbs and grasses whereas in winter older stands (pole and tree stage, and old forest) and south-eastern facing slopes were selected (cf. Table 3 for home range selection in the study area and S3–S7 Tables for further information regarding the model selection, VIF and odds ratios).

Habitat use within the home range

Habitat use within the home range during summer and winter was explained by vegetation, land use and human presence (Table 4). The zoning scheme also ranked among the most important predictors, with refuges and core areas being selected over the border zones in summer (Table 4, S2 Table, available online in Supporting Information). In winter red deer aggregated at the feeding sites during the day, whereas the refuges were predominantly selected during the night (Table 4). However, habitat use differed significantly between the day and nighttime, particularly with regard to the variables related to human presence (Fig 2, Tables 4 and 5). Both in summer and winter recreation trails were avoided during day, whereas a positive association could be found during night (Table 4). In addition, red deer selected areas with shallower slopes and in greater vicinity to water during night. In the summer season, red deer visited bilberry patches during night that were avoided during daytime, whereas in the winter season, they stayed more frequently in the vicinity of roads during nights compared to daytime (Table 5).

Discussion

Effects of human presence and outdoor recreation

Our study shows how adjustments of behaviour can result in oppositional patterns of wildlife habitat use at day and nighttime, when areas frequented by recreationists are avoided during the day and preferred during the night. Although it is suggested that animals become habituated to human presence [73] and might reduce flight-distances in areas with frequent human-wildlife contact [74], the deer avoided the areas close to the trails during daytime. Whereas in North America it is a widely known phenomenon that deer habituate to humans and even occur in settlements where they are not hunted [51], the deer in our study seem to actively avoid human recreationists. This might indicate that red deer are unable to distinguish recreational users and hunters, and therefore temporally avoid areas with high human use. The diurnal pattern was blurred when pooling day and night locations (S2 Table), which highlights the importance of accounting for temporal differences when analyzing human-wildlife interactions.

Linking spatiotemporal patterns of wildlife habitat use to human presence is an important prerequisite for designing efficient wildlife management concepts, even if reducing disturbance is not the primary management goal as it might be the case in hunted species like the red deer. Previous studies showed that red deer respond to the presence of recreationists by fleeing [43], moving to denser vegetation areas [43], increasing vigilance [40] and adjusting their foraging behaviour [42]. Sibbald et al. [41] found an avoidance of hiking trails by red deer, which was stronger during the day with higher visitor numbers.

Since the infra-red counter data (S2 Fig), collected shortly after our study on red deer show a strong diurnal variation of use (i.e. many visitors during day, little or none during night) and no other factor in the area which is spatially linked to recreational trails shows a diurnal pattern, we assume that the avoidance of recreational trails by day is caused by a the presence of recreationists. Sibbald et al. [20], also showed that red deer flexibly adjust their habitat use to the diurnal variation in human presence. Even though we could not directly link deer
### Table 4. Habitat use within the home range in summer and winter, day and night.

| Type       | Variable        | Summer Day (AUC: 0.684 ± 0.005) | Sign. | ΔAIC | Summer Night (AUC: 0.810 ± 0.005) | Sign. | ΔAIC |
|------------|-----------------|---------------------------------|-------|------|----------------------------------|-------|------|
|            |                 | SD (Individual): 0.329 AIC: 12040 |       |      | SD (Individual): 0.450 AIC: 7317 |       |      |
| **INTERCEPT** | -4.675 | 0.226 | *** | 37 | 2.190 | 0.188 | *** |
| **Vegetation** | CANOPY_COVER | -0.013 | 0.002 | *** | -0.036 | 0.002 | *** |
|             | SUC_REGTHICK | 2.412 | 0.224 | *** | 0.498 | 0.466 | 19 |
|             | SUC_POLe | 1.351 | 0.204 | *** | 0.743 | 0.437 | * |
|             | SUC_TREE | 0.853 | 0.199 | ** | 0.732 | 0.433 | . |
|             | SUC_OLD | 0.982 | 0.202 | *** | 0.203 | 0.436 | 19 |
|             | BILBERRY | -0.025 | 0.003 | *** | 59 | 0.022 | 0.003 | *** |
|             | PROTECT_S | -0.004 | 0.001 | ** | 6 | 6 |
|             | UNDER_CON | 0.731 | 0.100 | *** | 143 | 143 |
|             | UNDER_DEC | 1.850 | 0.245 | *** | 0.498 | 0.466 | 19 |
|             | UNDER_CONMIX | 0.674 | 0.079 | *** | 0.203 | 0.436 | 19 |
|             | UNDER_DECm | 0.148 | 0.088 | . | 0.203 | 0.436 | 19 |
|             | CAN_CON | -0.760 | 0.433 | . | 26 | 26 |
|             | CAN_DEC | 0.405 | 0.617 | . | 26 | 26 |
|             | CAN_CONMIX | -0.678 | 0.434 | . | 26 | 26 |
|             | CAN_DECm | -1.330 | 0.468 | ** | 26 | 26 |
| **Landscape** | WATER | -2.487 | 0.257 | *** | 160 | 1.175 | 0.361 | *** |
|             | EASTING | 0.083 | 0.042 | * | 2 | -0.409 | 0.055 | *** |
|             | SLOPE | 0.064 | 0.004 | *** | 213 | -0.042 | 0.006 | *** |
|             | NORTHING | -0.279 | 0.035 | *** | 60 | 60 |
|             | NORTING | 1.583 | 0.224 | *** | 51 | 51 |
|             | NORTHING | 1.159 | 0.118 | *** | 172 | 172 |
|             | MGT_CORE | 1.282 | 0.158 | *** | 73 | 1.371 | 0.136 | *** |
|             | MGT_REFUGE | 1.249 | 0.165 | *** | 73 | 1.371 | 0.136 | *** |
|             | TOURI_S | -1.616 | 0.189 | *** | 157 | 0.865 | 0.242 | * |
|             | HUNT | 1.159 | 0.118 | *** | 172 | 172 |
|             | SETTLE | 0.254 | 0.064 | *** | 33 | 33 |
|             | ROAD | 0.503 | 0.079 | *** | 27 | 27 |
| **Human** | MGT_CORE | 1.282 | 0.158 | *** | 73 | 1.371 | 0.136 | *** |
|             | MGT_REFUGE | 1.249 | 0.165 | *** | 73 | 1.371 | 0.136 | *** |
|             | TOURI_S | -1.616 | 0.189 | *** | 157 | 0.865 | 0.242 | * |
|             | HUNT | 1.159 | 0.118 | *** | 172 | 172 |
|             | SETTLE | 0.254 | 0.064 | *** | 33 | 33 |
|             | ROAD | 0.503 | 0.079 | *** | 27 | 27 |

### (Continued)
behaviour to the intensity of recreation activities on the trails, as visitor counts obtained with photo sensors (S2 Fig) were collected one year after the telemetry data, we assume that the diurnal pattern of recreation activities was similar during the time of our study. However, detailed information on the number of visitors per specific trail and time of the day would be favorable for quantifying the number of visitors that triggers an avoidance reaction in red deer.

Apart from human recreational infrastructure, habitat selection was based on a variety of factors related to forage quality (e.g. bilberry, deciduous trees) and essential resources (e.g. water). The predominant use of these habitat features during night but not during the day indicates that resources attractive to the deer are temporally not accessible due to human disturbance. In our study area this applied to the area along the lake, which is highly frequented by recreationists, as well as to clearings with abundant ground vegetation and bilberry patches.
which are mainly located in open forest with high visibility. During the day, particularly in summer, the deer was more frequently found in dense forest stands providing cover, i.e. thickets and pole stands (Tables 4 and 5). This finding supports the suggestions by previous studies \[44,47\] that human disturbance may contribute to reinforce possible conflicts with forestry: if clearings and open forest near trails are not usable for foraging during the day due to disturbance, the deer may be forced to relocate to dense forest stands with cover where they may cause forest damage by tree browsing and bark stripping as no alternative food is available in these stands.

### Zoning scheme

To reduce human-wildlife conflicts, zoning schemes might play an important role for wildlife managers confronted with combining varying interests in human dominated landscapes within central Europe. In our study, both seasonal home range selection within the study area, as well as habitat use within the home range, were closely linked to the zoning scheme: red deer selected the refuge areas over the core and border zone of the management scheme and—as expected—stayed close to the feeding stations in winter. We cannot prove a causal effect of zoning on red deer habitat selection though, as no systematically collected data before the establishment of management zones were available and we cannot exclude that the delineation of zones might have been influenced by pre-existing expert knowledge. It is therefore possible

**Table 5. Differences between diurnal and nocturnal habitat selection within the home range with regard to the relevant environmental predictors selected in the final models (Table 4).**

| Type       | Variable         | Estimate | SE   | Sign. | Estimate | SE   | Sign. |
|------------|------------------|----------|------|-------|----------|------|-------|
| STD (Individual): 0.967 | | | | | | | |
| Vegetation | INTERCEPT        | 6.661    | 0.322| ***   | 6.431    | 0.448| ***   |
| CAN_CON    | -1.194           | 0.448    |     |       | -1.373   | 0.310| ***   |
| CAN_DEC    | -0.557           | 0.739    |     |       | -4.467   | 2.293| *     |
| CAN_CONMIX | -1.074           | 0.447    |     |       | -1.312   | 0.322| ***   |
| CAN_DECMIX | -1.826           | 0.459    |     | *     | -3.483   | 0.615| ***   |
| SUC_REGTHICK| -2.172           | 0.461    |     | ***   |         |      |       |
| SUC_POLE   | -1.234           | 0.454    |     | ***   |         |      |       |
| SUC_TREE   | -0.323           | 0.453    |     | *     |         |      |       |
| SUC_OLD    | -0.341           | 0.464    |     | *     |         |      |       |
| BILBERRY   | 0.037            | 0.003    |      | ***   |         |      |       |
| FOREST250  | -2.207           | 0.250    |     | ***   |         |      |       |
| UNDERCOV   | -0.011           | 0.002    |     | ***   |         |      |       |
| Landscape  | SLOPE            | -0.065   | 0.005| ***   | -0.112   | 0.011| ***   |
| WATER      | 4.110            | 0.314    |     | ***   | 10.742   | 0.651| ***   |
| NORTING    | 0.258            | 0.044    |     | ***   |         |      |       |
| EASTING    | -0.203           | 0.047    |     | ***   |         |      |       |
| Human      | ROAD             | 0.740    | 0.068| ***   | 2.015    | 0.230| ***   |
|            | TOURI_S/W        | 1.733    | 0.222| ***   | 8.290    | 0.478| ***   |
|            | HUNT             | -1.087   | 0.139| ***   | -2.689   | 0.310| ***   |
|            | MGT_CORE         | -0.269   | 0.208|       |         |      |       |
|            | MGT_REFUGE       | 0.187    | 0.220|       |         |      |       |

Positive estimates indicate a relatively more frequent use of this variable in the night, while negative estimates indicate a relative more frequent use during the day. Significance levels are indicated with: * $p \leq 0.05$ and *** $p \leq 0.001$.

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that the deer had already preferred these areas prior to the establishment of the zonation scheme, due to other factors such as traditions or the distribution of forage.

Management implications

The diurnal avoidance of human recreation infrastructure by red deer, both in summer and winter, associated with an increased nocturnal use of temporarily inaccessible resources has several implications for the management of natural areas. As the avoidance of trails during the daytime renders some areas and resources inaccessible to the deer, it is important that the animals are not additionally disturbed during the night. Nocturnal sport events (i.e. torch-lit walks, nocturnal orienteering) should thus be strictly regulated in areas with disturbance-sensitive wildlife. In addition, patches of open forest, clearings and meadows, providing alternative food sources should be created within sufficient distance and with visual protection from hiking trails. Wildlife refuges, from which recreation is banned provide undisturbed areas during both, day and night, and are likely to benefit also other disturbance-sensitive wildlife.

Supporting information

S1 Data. all relevant data for this study is included in the data file.

S1 Fig. Relative probability of red deer presence in summer (upper panel) and winter (lower panel), during day (left) and night (right). Black and white represent high and low probability of presence respectively. Dashed lines indicate the presence of summer and winter recreation trails, respectively.

S2 Fig. Mean number of visitors per hour present on selected summer (black) and winter (grey) trails, for the different times of the day. The shaded area shows the time between sunrise and sunset for the studied time period. For three months (17.2.2010 14.4.2010) TRAFx Infrared trail counters were placed along three designated hiking trails and three cross country skiing trails within the study area. These count the number of times an individual passed the light sensor. Although we cannot exclude that red deer crossed the sensors, particularly during nighttime, we assume most of these crossings are humans since these were placed along designated recreation infrastructure. Most of the activity on the trails is during the time between sunrise and sunset.

S1 Table. Number of locations per individual used for analysis. Number of locations per individual per season and time of the day used for the analysis, and the period of tracking. The age was roughly estimated in three classes at the time of tagging (1 = 1–3 years, 2 = 3–5 years, 3 = >5 years old).

S2 Table. Models explaining habitat selection within the home range, when not discriminating between different times of the day. Left panel: summer, right panel: winter. Significance levels are indicated with: * p≤ 0.05, ** p≤ 0.01, and *** p≤ 0.001.

S3 Table. Results of the variable selection process to reach the final models presented in Tables 3 and 4. The first column shows all variables which were included in the model selection process. Each of the other columns represents one of the six final models: “HRinSA” = home range selection within study area, “inHR” = habitat selection within home range.
Variables denominated with a “+” are included in the final model, otherwise the reason for exclusion is indicated: A variable name indicates exclusion due to pairwise correlation (Spearman’s R > 0.5) with this variable, “-” indicates exclusion during the model selection process based on AIC as described in the methods part, “VIF” indicates this variable is excluded from the model due to a too high variance inflation factor value. Proximity of feeding stations (FEED) was not included in the summer models as no feeding was performed in summer.

**S4 Table.** Final models (provided in Tables 3 and 4) in comparison to the next-best candidate models as obtained during the model selection process. All candidate models with a ΔAIC < 2 to the final model (in bold) as well as the first model with ΔAIC > 2 are shown. Only for the model describing home range selection in the study area during summer (Table 3A) the next four candidate models were not significantly different to the final model (i.e. ΔAIC < 2), so model averaging was applied (S5 Table).

**S5 Table.** Model averaging of the model describing home range selection in study area during summer (Table 3A). The table shows the variables included in the five component models (with variable codes described below), as well as the relative importance of the variables.

**S6 Table.** Variation inflation factors (VIF) for the variables included in the models presented in Tables 3 and 4. For categorical variables the corrected VIF values (VIF^(1/2Df)) are provided.

**S7 Table.** Odd’s ratios with 95% confidence interval for the coefficients of the models provided in Tables 3 and 4. Odd’s ratios were obtained using the Wald chi-square method, the lower (2.5%) and upper (97.5%) boundary of the confidence interval are provided.

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