Attacked from two fronts: Interactive effects of anthropogenic and biotic disturbances generate complex movement patterns

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ABSTRACT
Anthropogenic and biotic disturbances have the potential to interact, generating cumulative impacts on animal movement or, alternatively, counterbalancing or masking each other. Despite their importance, those interactions have not been investigated thoroughly. Our study aimed to fill this knowledge gap by assessing the combined effects of a human activity—that is, military exercises—and a biotic disturbance—that is, insect harassment—on movement rates of free-ranging semidomesticated reindeer (Rangifer tarandus tarandus). From 2010 to 2012, we analyzed location data from fifty-one Global Positioning System (GPS)-collared female reindeer in the largest European military test range, situated in northern Sweden. In the presence of both military exercises and mosquito harassment, reindeer reacted by increasing their movement rates but not as much as when mosquito harassment occurred alone. Conversely, reindeer reduced their movement rates during military exercises performed with aircraft. Moreover, the effect of military exercises performed with vehicles was evident only when combined with mosquito harassment. These results stress the value of evaluating the effects of the interaction between biotic disturbances and human activities, especially in northern ecosystems, because of the predicted climate warming and the growing interest toward natural resource extraction and other forms of land use.

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Introduction

Today, most areas around the world are perturbed by human activities. For example, the contiguous United States are completely lacking areas free from anthropogenic noise (Barber et al. 2011). Similarly, the development and expansion of lighting technologies are making light pollution a spreading threat to natural systems around the world (Longcore and Rich 2004; Gaston et al. 2013). Moreover, the continuous expansion of the transport sector is generating levels of air and noise pollution that are turning into a global threat (Holden and Linnerud 2015; Lindgren and Wilewska-Bien 2016; Shannon et al. 2016). Human activities may displace and disturb animals by altering their habitat or affecting their behavior (Steidl and Powell 2006), with possible complex repercussions because animals can perceive human activities through diverse means, such as auditory, visual, and olfactory cues. The response is usually driven by a trade-off between the benefit of responding to the disturbance—that is, an increased chance of survival when the disturbance is perceived as a risk—and its cost; for example, displacement or reduced time available for parental care, mating, or feeding activities (Frid and Dill 2002). When the anthropogenic disturbance is strong and causes a reduction in the time spent feeding and resting, the cost of the response can be substantial and severely affect the animal’s survival and fitness (e.g., Luick et al. 1996). At the larger scale, human activities can end up affecting entire populations by displacing or confining them to low-quality habitats or habitats with higher predation risk (Frid and Dill 2002).

Anthropogenic disturbances can have long-lasting effects on arctic and subarctic ecosystems, due to the low biodiversity and harsh conditions of those systems (Forbes, Ebersole, and Strandberg 2001; Willard, Cooper, and Forbes 2007; Reynolds and Tenhunen...
areas used seasonally by humans, such as mountains, valleys, and plains. Traffic, helicopters and aircraft (Harrington et al. 1983, Legagneux et al. 2014), and other activities impose costs for the animals inhabiting them (Johnson et al. 2005). Moreover, climate change is manifested in the arctic and subarctic regions at faster rates than in any other area in the world (IPCC 2014). Arctic and subarctic ecosystems are shaped by their cold and harsh climate, and global warming is expected to have acute effects on the structure and functioning of their simple but sensitive food webs (Ims and Fuglei 2005; Stempniewicz, Blachowiak-Samołyk, and Węsławski 2007; Legagneux et al. 2014; Barthelemy et al. 2017). Even small disturbances directly affecting only one species may have widespread, indirect effects on a variety of organisms (Croll et al. 2005; Post et al. 2009). Direct anthropogenic disturbances and climate change have the potential to have interacting effects on species living in arctic and subarctic ecosystems, but that interaction has so far received little attention (but see Raynolds et al. 2014).

Reindeer (Rangifer tarandus, a species called caribou in North America) are the most abundant and important large herbivore in arctic and subarctic ecosystems. Reindeer regulate plant communities through grazing and are the main prey for several predator species (Pedersen et al. 1999; van der Wal 2006; Musiani et al. 2007; Andren et al. 2011; Hobbs et al. 2012; Bernes et al. 2015). Moreover, reindeer constitute the main source of subsistence for at least twenty indigenous peoples (Turi 2002; Forbes et al. 2006, 2009; Forbes 2013). Both wild and semidomesticated reindeer are integral parts of northern ecosystems (Uboni et al. 2016), because during most of the year semidomesticated reindeer range freely on natural pastures (Forbes and Kumpula 2009). Various human activities and infrastructures affect the behavior, movement, and habitat selection of wild and semidomesticated reindeer and caribou; for example, by increasing movement speed and distance traveled, by preventing animals from visiting high-quality pastures, or by increasing the time spent in vigilance behavior or traveling, which may decrease feeding time (Vistnes and Nellemann 2008; Skarin and Åhman 2014). Among those activities and infrastructures, we find tourism and hunting (e.g., Vistnes et al. 2008; Reimers et al. 2009; Skarin et al. 2010; Wilson and Wilmhurst 2019), vehicle traffic, helicopters and aircraft (Harrington 2003; Reimers and Colman 2006), and drill sites, mines, dams, and various kinds of power plants (e.g., Fancy 1983; Murphy and Curatolo 1987; Panzacchi et al. 2013; Skarin et al. 2015; Skarin, Sandström, and Alam 2018). Most studies so far have addressed only the short-term effects of those activities and infrastructures on reindeer behavior, but some have demonstrated that their impact goes beyond movement and habitat selection, affecting reproduction rates and calf survival (Luick et al. 1996; Vistnes and Nellemann 2008).

Harassment from insects is another well-known factor influencing reindeer behavior and fitness. Mosquitoes (especially Aedes spp., Culicidae), blackflies (Simulidae), horseflies (Tabanidae), warble flies (Hypoderma tarandi L., Oestridae), and nose bot flies (Cephenemyia trompe L., Oestridae) cause the strongest harassment, considerably affecting reindeer activity patterns (Hagemoen and Reimers 2002). Oestrid flies are indeed excellent and fast fliers and easily detect and follow reindeer (A. C. Nilssen and Anderson 1995), and reindeer often exhibit violent reactions to fly harassment, running for hours in search for relief (Hagemoen and Reimers 2002). Mosquitoes also have the ability to alter reindeer behavior (Fancy 1983; Downs, Theberge, and Smith 1986; Mörschel and Klein 1997; Noel et al. 1998; Raponi et al. 2018). In northern Scandinavia, oestrid activity occurs approximately between early July and early September (Anderson, Nilssen, and Folstad 1994; A. C. Nilssen 1997; Anderson et al. 2001), and mosquitoes are mainly active between June and August (Schäfer and Lundström 2001). During periods of intense insect harassment, reindeer search for relief in fresh and windy areas such as snow patches, hilltops, ridges, mires, open areas, roads, and stream estuaries (Fancy 1983; Helle and Aspi 1984; Downs, Theberge, and Smith 1986; Pollard et al. 1996; Noel et al. 1998; Skarin et al. 2004; Moen 2008; Vistnes et al. 2008). In those areas, reindeer are expected to move less, or slower, than in areas with high insect harassment. Overall, intense insect harassment may have dramatic consequences for reindeer health, body conditions, and fecundity (Hagemoen and Reimers 2002). In some cases, reindeer may even increase their tolerance to anthropogenic disturbances if insect harassment is severe. For example, reindeer are expected to avoid areas used seasonally by humans, such as mountain tourist huts and trails (Skarin 2007). However, in the Scandinavian Mountains, reindeer move to higher elevations in July and August to escape insect harassment even when those areas are heavily used by tourists (Skarin et al. 2004). Thus, the necessity to escape insects seems to override the need for human avoid- ance. Fancy (1983) determined that caribou movement
rates did not differ between areas characterized by high human disturbance (drill sites) and control areas under either low or high insect activity levels. Additionally, the effects of the disturbance generated by oilfields on caribou behavior are more pronounced in the absence of insects (Murphy and Curatolo 1987). Caribou may tolerate some level of human disturbance because drill sites are open and windy areas; that is, optimal insect relief areas (Fancy 1983; Pollard et al. 1996; Noel et al. 1998). Reindeer response to other types of human disturbance may also be overridden by insect harassment, and if the two disturbances are not analyzed simultaneously, their effects may be underestimated or misinterpreted.

Because of their essential role in arctic and subarctic ecosystems and their vulnerability to human encroachment in those ecosystems (Uboni et al. 2016), we chose free-ranging semidomesticated reindeer as a study species to investigate the potential combined effects of a human activity—that is, military exercises—and insect harassment on animal movements. Our study system was located in a military test range, which also overlapped the reindeer summer grazing grounds. Specifically, our aims were to assess whether (1) reindeer move away from their traditional summer range while military exercises occur and (2) such military exercises and insect harassment interact in affecting reindeer movement. We hypothesized that reindeer would react to human presence and loud noises due to military exercises by relocating from the test range. Moreover, we hypothesized that the effect of military exercises may be masked by insect harassment when the two disturbances occur simultaneously.

Methods

Study area

The study was conducted in the range of the Udtja reindeer herd, belonging to the Udtja reindeer herding community, Norrbotten County, northern Sweden (Figure 1). In Udtja, reindeer spend all year in the boreal forest. Nonetheless, they have distinct summer and winter ranges and range freely within the borders of the community, controlled by a fence toward the northwest and by active work of the herders in the southeast (Swedish Sami Parliament 2019; https://www.sametinget.se/8744). The summer range is dominated by old forest, constituted primarily by Scots pine (Pinus sylvestris) and Norway spruce (Picea abies), interspersed with mires, lakes, and, on mountain tops, downy birch (Betula pubescens) forest. Elevation ranges between 187 and 714 m.a.s.l., and roads are concentrated toward the southeast at rather low densities (gravel roads: 0.25 km/
km²; paved roads: 0.02 km/km²). Monthly temperatures in July–September averaged 10.13°C (±3.31) in the period 2006–2016, and the precipitation monthly sum for the same months averaged 69.78 mm (±37.74) in the period 2013–2016 (Swedish Hydrological and Meteorological Institute 2017; Tjåkaape weather station, http://opendata-downloadmetobs.smhi.se/explore/).

A military test range, the Vidsel Test Range, was established in 1959 within the Udtja spring and summer reindeer ranges. There, the main human activity is represented by military training exercises. With its 1,625 km² of ground space, this area is Europe’s largest military test range. Roads inside the test range are used only in conjunction with the military exercises. Outside the test range, main roads are used by regular traffic, and most secondary roads are predominantly used in conjunction with forestry activities. Inside the test range, 1,465 km² were designated in 1995 as a nature reserve where logging activities are not permitted (Figure 1). The main reindeer predator in the area is the brown bear (*Ursus arctos*), but European lynx (*Lynx lynx*), wolverine (*Gulo gulo*), and golden eagle (*Aquila chrysaetos*) also make up part of the predation (Sivertsen 2017).

**Data collection**

**Reindeer Global Positioning System data**

Between 2010 and 2012, we fitted seventy-one adult female reindeer (*n* = 19 in 2010, *n* = 27 in 2011, *n* = 25 in 2012) with Global Positioning System (GPS) collars (Followit AB, Tellus) in the Udtja reindeer herding community and downloaded the data manually in the field once a year during the winter gatherings of the animals. The collars were programmed to collect GPS locations at 2-hour intervals and the average GPS fix rate was 86.2 percent. Subsequently, we removed six individuals from the analysis because of collar failure (*n* = 2 in 2010, *n* = 3 in 2011, *n* = 1 in 2012) and fourteen individuals because their fix rate was <80 percent (*n* = 5 in 2010, *n* = 4 in 2011, *n* = 5 in 2012), which left fifty-one reindeer for analysis (*n* = 12 in 2010, *n* = 20 in 2011, *n* = 19 in 2012). Of those, forty-nine were fitted with collars for one year and two were followed for two years. We cleaned the GPS data set by removing obvious outliers (i.e., locations falling outside the study area, detected based on latitude and longitude), locations with a two-dimensional accuracy or a dilution of precision > 5, duplicates, and clear erroneous locations identified based on movement speed. Because herders gather their reindeer for calf marking prior to 12 July and insect harassment in northern Scandinavia occurs from early July up to early autumn (Anderson, Nilssen, and Folstad 1994; A. C. Nilssen 1997; Anderson et al. 2001; Skarin et al. 2010), we restricted the study period to 12 July to 10 September.

**Military exercises**

We obtained daily data on military exercises performed at the Vidsel Test Range in 2010–2012 by the Swedish Defense Materiel Administration in the form of occurrence of military exercises aiming at three locations (E,
Q, and T) within the test range (Figure 1), hereafter defined as activity points. Due to the classified nature of those data, the spatial extent of the military exercises was unknown. The exercises were performed with light (e.g., cars and trucks) and heavy terrestrial vehicles (e.g., tanks) or with aircraft. Because military exercises performed with light and heavy terrestrial vehicles always occurred simultaneously, we hereafter refer to those exercises as being performed with vehicles. Vehicles were usually driven toward one of the activity points from points C or D50 (Figure 1), whereas aircraft flew into the test range from a nearby military airport in the southeast. No mitigation measures were adopted in relation to disturbance to reindeer.

**Insect harassment indices**

Fly and mosquito harassment are positively correlated with air temperature and negatively correlated with wind speed (e.g., Hagemoen and Reimers 2002). Moreover, fly harassment is negatively correlated with cloud cover (Anderson, Nilssen, and Folstad 1994; Colman et al. 2003). The intensity of harassment by flies and mosquitoes on ungulates can be summarized by an insect harassment index (IHI; Weladji, Holand, and Almøy 2003) when the weather conditions of a certain area at a certain time and the threshold of insect activity (in terms of temperature, wind speed, and cloud cover) are known. For example, according to Weladji, Holand, and Almøy (2003), oestrid flies are active at air temperatures ≥13°C, wind speeds <6 m/s, and cloud cover <40 percent. Based on that information, Weladji, Holand, and Almøy (2003) developed an oestrid harassment index (OHI):

\[ OHI = PHD + S, \]

where PHD, the harassment day factor, indicates a day with midday temperature above the threshold of fly activity and wind speed below the threshold, and S, the severity factor, reflects cloud cover below the threshold.

Using hourly weather data provided by the Swedish Hydrological and Meteorological Institute (2017; Tjåkaape weather station, http://opendata-downloadmetobs.smhi.se/explore/), we calculated an OHI and mosquito harassment index (MHI) for our study period and area. However, in order to develop IHIs that are as precise as possible, we adapted Weladji, Holand, and Almøy’s (2003) index by averaging the wide range of values reported in the literature for temperature and wind speed thresholds of oestrid and mosquito activity (Table S1). We calculated the OHI (ranging from 0 to 1.5) based on PHD = 1 on a day with temperature >13°C and wind speed <9.6 m/s and S = 0.5 on days with cloud cover <40 percent. The MHI could only take a value of 1 when the temperature was between 7°C and 17°C and the wind speed was <6.12 m/s or a value of 0 otherwise, because there is no evidence of a relationship between cloud cover and mosquito activity. The IHIs were not validated in our study area. Because oestrid flies are obligate parasites of reindeer and reindeer are the most abundant large mammal that mosquitoes can feed upon in northern Scandinavia (except humans), when those insects are active, reindeer are likely one of their main targets. Moreover, our study covers the period of activity of adult oestrid flies and mosquitoes in northern Scandinavia (Anderson, Nilssen, and Folstad 1994; A. C. Nilssen 1997; Schäfer and Lundström 2001), and insect activity is highly dependent on meteorological conditions, which the indices take into account. Therefore, we assume that the IHIs are accurate proxies for insect harassment on reindeer.

**Environmental and anthropogenic variables**

Based on previous studies (Skarin et al. 2008, 2010), we collected six environmental and anthropogenic variables to assess drivers of reindeer movement (see models below): (1) vegetation type; (2) elevation; (3) slope; (4) ruggedness; (5) the Normalized Difference Vegetation Index (NDVI); and (6) the Euclidean distance between each reindeer location and the nearest road. We obtained maps of vegetation type, roads, and elevation (50-m-resolution digital elevation model) from Lantmäteriet (2018; www.lantmateriet.se). We derived slope and ruggedness from the digital elevation model using the slope and vector ruggedness measure (Sappington, Longshore, and Thompson 2007) in a geographic information systems (GIS) environment (QGIS 3.2.0; QGIS Development Team 2018). We obtained twelve sixteen-day composite NDVI maps (MODIS Terra Vegetation Index 250-m resolution) that covered the entire study period from NASA Earthdata Search website (2018; https://search.earthdata.nasa.gov/). Because the original NDVI maps had a 200-m resolution, we resampled them at 50-m resolution.

**Data analysis**

**Reindeer response to military exercises**

To assess whether reindeer responded to military exercises by moving away from the military activity points, we developed univariate regression models in which the daily average Euclidean distance (in meters) of each reindeer from the three activity points was the response variable (calculated in QGIS 3.2.0; QGIS Development Team 2018). The predictor categorical variable described the occurrence of military exercises
performed either with vehicles or with aircraft (Vehi_ALL and Air_ALL, respectively), each ranging from 0 (i.e., no military exercises performed in a given day) to 3 (i.e., military exercises performed at all three activity points). Due to collinearity between Vehi_ALL and Air_ALL (pairwise Pearson’s correlation coefficient: $r = 0.8$), we did not include both variables in the same models. For each type of military activity, we compared a set of six models: (1) a generalized least squares (GLS) model; (2) a generalized linear mixed effects model (GLMM) with reindeer ID as random intercept term; (3) a GLMM with year as a random term; (4) a GLMM with reindeer ID and year as nested random terms; (5) a GLS with an autoregressive correlation structure, AR1; and (6) a GLMM with reindeer ID as random intercept term and an autoregressive correlation structure, AR1. We evaluated model performance based on the Akaike information criterion (AIC; Burnham and Anderson 2002). Because no military exercises occurred in July, we limited this analysis to 1 August to 10 September. Furthermore, we only included those reindeer that spent at least part of the study period inside the test range ($n = 10$ in 2010, $n = 7$ in 2011, $n = 11$ in 2012).

Effects of insect harassment and military exercises on reindeer movement

To assess whether insect harassment and military exercises affected reindeer movement, we fitted GLS models and GLMMs using the Euclidean distance traveled by a reindeer between successive GPS locations (hereafter referred to as movement rate but also defined as step length) as the response variable. The predictor variables were Vehi_ALL, Air_ALL, OHI, MHI, temperature, elevation, slope, ruggedness, vegetation type, NDVI, distance from roads, and area (i.e., a binary variable indicating whether the reindeer was moving inside or outside the military test range). Thus, we did not measure insect harassment directly but inferred it from the IHIs. Movement rates were not estimated if one or more consecutive GPS locations were missing. Because reindeer movement rates were temporally autocorrelated and affected by circadian rhythm (Online Resource, Figures S1 and S2; but see also Erriksson, Källqvist, and Mossing 1981; van Oort et al. 2005), we followed Dray, Royer-Carenzi, and Calenge (2010) by using the nb2listw function in the R package “spdep” to define a spatial weighting object to represent the temporal structure of the data (Bivand, Hauke, and Kossowski 2013; Bivand and Piras 2015).

Subsequently, we detrended the data through the principal coordinate analysis of neighbor matrices (for details, see Borcard and Legendre 2002; Dray, Legendre, and Peres-Neto 2006) and used the detrended residuals as response variable in our models. We excluded from detrending twenty reindeer ($n = 7$ in 2010, $n = 9$ in 2011, $n = 4$ in 2012) because the procedure did not support the low temporal autocorrelation in their movement rates. Therefore, the final data set for this analysis included thirty-one female years (corresponding to twenty-nine female reindeer, because for two were followed two consecutive years).

We treated Vehi_ALL, Air_ALL, OHI, and MHI as continuous variables. We averaged OHI, MHI, and temperature on a 2-hour interval to match the GPS fix interval. We calculated elevation, slope, and ruggedness as the mean of the values along each step. We assigned temporally and/or spatially specific values of vegetation type, NDVI, and distance from roads to each step based on the first GPS location in the step. Lastly, we included interaction terms between the IHIs, and (1) the military exercises variables, (2) vegetation type, (3) elevation, and (4) distance from roads as well as area and (1) distance from roads and (2) the military exercises variables. This latter term aimed to assess whether reindeer reacted differently to military exercises whether the reindeer was close to or far from the areas where the exercises took place. The interactions between the IHIs, vegetation type, and elevation aimed to determine whether reindeer used insect relief areas in response to insect harassment. We screened all predictor variables for collinearity using a pairwise Pearson’s correlation coefficient (threshold for removal: $r \geq 0.8$), followed by the variance inflation factors (threshold for removal: variance inflation factor $\geq 3$; Zuur, Ieno, and Elphick 2010). We fitted four separate sets of models, one for each combination of IHI and type of military exercise, because of high correlation between OHI and MHI ($r = 0.81$; Online Resource, Table S2) and between the military exercises performed with vehicles and aircraft ($r = 0.83$). Each set of models included (1) a GLS model with no random term, fixed variance, or autocorrelation structure; (2) three GLMMs with either reindeer ID, year, or reindeer ID/year as random intercept terms; (3) two GLS models with either year or month as a fixed variance structure; and (4) a GLS model with year as a fixed variance structure and with an exponential spatial autocorrelation structure applied to the model residuals (range = 850, nugget = 0.4, sill = 1). We tested reindeer ID as a random intercept term to exclude the influence of differences in behavior and sample size among individual animals (Gillies et al. 2006). We added year and month as fixed variance structures to address heteroscedasticity and temporal variability (Gillies et al. 2006; Uboni, Smith et al. 2015; Uboni, Vucetich et al. 2015).
We tested the exponential spatial correlation structure to address spatial autocorrelation caused by landscape features such as topography and vegetation density, which could restrict or ease reindeer movement (Skarin et al. 2010; Panzacchi et al. 2016). We identified the best-fit model based on AIC and used restricted maximum likelihood estimation in all models for their AICs to be comparable (Zuur et al. 2009).  

Once the need for a random term, a variance structure, and a correlation structure were determined, we performed an automatic backward model selection procedure using the stepAIC function in the “MASS” package for R (Venables and Ripley 2002). In order to compare models with different fixed terms, we refitted the full model using maximum likelihood estimation. Then, once the most parsimonious fixed structure was found, we refitted the reduced model with restricted maximum likelihood (Zuur et al. 2009).

Results

Reindeer response to military exercises

According to the GLS with an autoregressive correlation structure (Online Resource, Table S3), the average distance between reindeer and activity points increased only when military exercises were performed with aircraft at one activity point ($\beta = 1.110.73$, SE = 257.26, $p = .0005$; all other $p s > .05$; Figure 2).

Effects of insect harassment and military exercises on reindeer movement

All best-fit models aimed to assess the effect of insect harassment and military exercises on reindeer movement rates included year as a variance structure and a spatial correlation structure but no random term (Online Resource, Table S4).

Reindeer movement rates increased in the predicted presence of mosquitoes, as they did, though to a lesser extent, when mosquito harassment and military exercises occurred simultaneously. On the contrary, reindeer movement rates decreased in response to military exercises performed with aircraft. Reindeer generally moved faster outside the test range and at higher elevations. Reindeer movement rates decreased with increasing distance to roads and steeper slopes, the former especially outside the test range (Table 1). Vegetation type, temperature, ruggedness, and NDVI did not seem to influence reindeer movement rates in the Udtja reindeer herding community.

Discussion

Movement rates of free-ranging, semidomesticated reindeer increased in the predicted presence of mosquito harassment, which was inferred from the MHI, whereas they decreased in response to the military exercises. When those two disturbances occurred simultaneously, reindeer response was intermediate. We hypothesize that in the presence of military exercises reindeer were not able to move as fast as they usually do when escaping harassing insects. At the same time, the effect of military exercises was masked when occurring simultaneously with insect harassment. These results stress the importance of assessing the impact of anthropogenic disturbances on animal movement behavior in relation to other biotic or abiotic stressors. Indeed, during our study, military exercises not only altered reindeer movement directly but also altered the reindeer response to insect harassment. A wide range of research has explored the ways in which human activities affect animal behavior (e.g., Longcore and Rich 2004; Steidl and Powell [2006] and references therein; Gaston et al. 2013; Shannon et al. 2016); similarly, many studies have assessed the effects of biotic and abiotic factors on animal movement (e.g., Avgar et al. 2013; Michel et al. 2013). However, few studies have investigated the co-occurrence of both sets of stressors on animal movement (Fancy 1983; Pollard et al. 1996; Noel et al. 1998), especially when their intensity varies over time. Our study reveals that the effects of those processes on animal movement can interact with each other and that analyzing them separately may lead to a misinterpretation of the complexity of their effects (Sih, Bell, and Kerby 2004). Moreover, the impact of human activities may be overlooked when those activities are performed at a time of high environmental stress for the animals. The biology of the animal should thus be considered in relation to the timing of an anthropogenic disturbance. In our study area, for example, the disturbance created by military exercises comes into play during a time when female reindeer are particularly susceptible to disturbance, because they are building up fat reserves for the winter and they are still lactating to feed their calves (Vistnes and Nellemann 2001; Harrington 2003; Skarin et al. 2008, 2015). Lactation requires a high amount of energy (White 1992); nonetheless, lactating female reindeer prefer an undisturbed environment at the expense of forage quality (Maier et al. 1998; Helle et al. 2012). If human disturbances occur at the time and in the areas of severe insect harassment, the consequences for female reindeer and their calves may be detrimental. Their increased movement rates translate into increased
Table 1. Effects of military exercises, insect harassment, and environmental and anthropogenic predictor variables on reindeer movement rates.

| Model          | Predictor                   | β    | SE   | p Value |
|----------------|----------------------------|------|------|---------|
| OHI–Vehi_ALL   | Intercept                  | 44.12| 63.57| .488    |
|                | OHI                        | −0.52| 24.31| .983    |
|                | Distance from roads        | −0.01| 0.00 | <.001***|
|                | Elevation                  | 0.15 | 0.11 | .173    |
|                | Slope                      | −11.46| 3.48 | <.001***|
|                | NDVI                       | −0.01| 0.00 | .087*   |
|                | Outside                    | 59.68| 29.00| .040**  |
|                | OHI: elevation             | 0.10 | 0.05 | .075*   |
|                | Distance from roads: Outside| −0.06| 0.01 | <.001***|
| OHI–Air_ALL    | Intercept                  | 37.76| 64.32| .557    |
|                | OHI                        | 2.79 | 24.50| .909    |
|                | Distance from roads        | −0.02| 0.00 | <.001***|
|                | Elevation                  | 0.18 | 0.11 | .119    |
|                | Slope                      | −11.46| 3.49 | <.001***|
|                | Air_ALL                    | −11.50| 9.59 | .230    |
|                | NDVI                       | −0.01| 0.00 | .086*   |
|                | Outside                    | 57.28| 29.26| .050**  |
|                | OHI: elevation             | 0.10 | 0.06 | .100    |
|                | Distance from roads: Outside| −0.06| 0.01 | <.001***|
| MHI–Vehi_ALL   | Intercept                  | −20.54| 59.32| .729    |
|                | MHI                        | 81.50| 11.13| <.001***|
|                | Distance from roads        | −0.02| 0.00 | <.001***|
|                | Elevation                  | 0.27 | 0.10 | .005**  |
|                | Vehi_ALL                   | −22.45| 12.69| .077*   |
|                | Slope                      | −11.25| 3.48 | <.001***|
|                | NDVI                       | −0.01| 0.00 | .113    |
|                | Outside                    | 54.08| 29.53| .067*   |
|                | Vehi_ALL: MHI              | 29.96| 14.72| .042**  |
|                | Distance from roads: Outside| −0.06| 0.01 | <.001***|
|                | Vehi_ALL: Outside          | 23.84| 14.83| .108    |
| MHI–Air_ALL    | Intercept                  | −19.51| 59.32| .742    |
|                | MHI                        | 79.71| 10.98| <.001***|
|                | Distance from roads        | −0.02| 0.00 | <.001***|
|                | Elevation                  | 0.27 | 0.10 | .004**  |
|                | Air_ALL                    | −29.61| 12.27| .016**  |
|                | Slope                      | −11.26| 3.48 | <.001***|
|                | NDVI                       | −0.01| 0.00 | .104    |
|                | Outside                    | 54.24| 29.49| .066*   |
|                | Air_ALL: MHI               | 39.24| 14.41| .007**  |
|                | Distance from roads: Outside| −0.06| 0.01 | <.001***|
|                | Air_ALL: Outside           | 31.24| 15.81| .048**  |

Note: Results refer to reduced regression models obtained with an automatic model selection procedure starting from the best-fit generalized least squares models listed in Online Resource, Table S4. All models included a variance structure fixed by year and an exponential spatial autocorrelation structure applied to the model residuals (range = 850, nugget = 0.4, sill = 1). Vehi_ALL and Air_ALL are continuous variables indicating the occurrence of military exercises performed at zero, one, two, or three military activity points, with either terrestrial vehicles or aircraft. Distance from roads = Euclidean distance in meters from the closest road. Outside = category of the dummy variable area, indicating that a reindeer was outside the military test range (reference category: inside). Statistically significant variables (p ≤ .05) are highlighted in bold.

***p ≤ .001. **< p ≤ .01. *p ≤ .10.

...as a consequence of the expansion of human activities and the increase in temperature predicted in northern ecosystems (Witter et al. 2012). Currently, insect harassment in arctic and subarctic regions is most severe in the central and hottest summer months, July and August. If the projected increase in temperatures occurs, the time of insect harassment may expand to late spring, when the calves are small and most vulnerable. Moreover, climate change is predicted to favor the expansion of new parasitic organisms into those regions (Mallory and Boyce 2018). Increased harassment may then start a negative feedback loop, because animals are even more sensitive to disturbances when they suffer nutritional stress. Coupled with increasing combined disturbances caused by various human...
activities (Vistnes and Nøllemann 2008; Skarin and Åhman 2014) and the negative effects of increased temperatures on reindeer physiology (K. J. Nilssen et al. 1984), increased harassment by insects and other parasites may have severe consequences for the species. Such disturbances may have complex repercussions on a wide range of species and environmental processes because of the key role that reindeer play in northern ecosystems (Pedersen et al. 1999; van der Wal 2006; Musiani et al. 2007; Andren et al. 2011; Hobbs et al. 2012; Bernes et al. 2015).

During the study period, female reindeer did not generally react to the military exercises by moving away from the areas where those activities were performed. On the one hand, reindeer may have traded off a peaceful environment for good quality pastures, especially because the disturbance created by military exercises was occasional and might not have been frequent enough to cause relocation of the reindeer (Gill, Norris, and Sutherland 2001). Woodland caribou (Harrington and Veitch 1991), peregrine and prairie falcons (Falco peregrinus and Falco mexicanus, respectively; Ellis, Ellis, and Mindell 1991), harlequin ducks (Histrionicus histrionicus; Goudie and Jones 2004), and red-cockaded woodpeckers (Picoides borealis; Delaney et al. 2011) tend to react similarly to military exercises. On the other hand, reindeer in the Utja reindeer herding community may have developed some level of habituation to the military exercises, especially those performed with vehicles. However, habituation should not be considered as a positive outcome, because animals that are habituated to vehicles are more likely to be victims of traffic accidents (Bejder et al. 2009). Additional studies are needed to determine whether reindeer have developed habituation to the military exercise but also to assess whether the effects of long and repeated military exercises on movement rates may translate into long-term consequences for the animal health, body conditions, and reproductive success, which could influence population growth and, in the case of reindeer husbandry, the local economy based on meat production.

Reindeer response to insect harassment (in terms of movement rates) did not differ depending on vegetation type, elevation, and distance from roads. Indeed, none of the interactions between the IHI and those variables were retained in the reduced regression models (Table 1). Reindeer often use open areas, higher elevations, and roads as insect relief areas (Helle and Aspi 1984; Downes, Theberge, and Smith 1986; Skarin et al. 2004; Moen 2008; Vistnes et al. 2008). In our study area, the maximum elevation is rather low (714 m.a.s.l.) and reindeer may not have access to many treeless hilltops. Indeed, in northern Scandinavia the treeline occurs at around 500–700 m. Moreover, roads have low density, which probably prevents reindeer from using those features to escape insect harassment. If roads had been used as insect relief areas, reindeer movement rates should have been lower in their proximity. On the contrary, reindeer movement rates increased with decreasing distance from roads (Table 1). We suggest that reindeer may use roads as movement corridors or they may perceive roads as a source of disturbance.

Female reindeer reacted to military exercises performed with aircraft by reducing their movement rates. This behavior may have originated as a response to a potential predator flying overhead. Golden eagles may indeed attack both adult females and calves (Nybakk, Kjelvik, and Kvam 1999; Nybakk et al. 2002; Gustine et al. 2006; Norberg et al. 2006; Nieminen, Norberg, and Majíjala 2011). Studies analyzing the antipredator response of reindeer to attacks by raptors are lacking in the literature, but attacks on other ungulates have been observed in some instances. Females of several ungulate species, such as Thomson’s gazelle (Gazella thomsonii), pronghorn (Antilocapra americana), bighorn sheep, mountain goat (Oreamnos americanus), chamois (Rupicapra rupicapra), and Dall’s sheep (Ovis dalli dalli), protect their young from raptor attacks by standing still (Nette, Burles, and Hoefs 1984; Locati 1990; Byers 1997; Bertolino 2003; Hamel and Côté 2009; Roberts 2014). Moreover, forest-dwelling reindeer are less gregarious than their conspecifics living in the tundra, which respond to predator attacks by grouping together (Baskin 1986; Bergerud 1988; Skarin and Åhman 2014). Dissimilarly, woodland and barren-ground caribou usually react to aircraft with a panic response, which results in overall increased activity levels and movement rates (Calef, DeBock, and Lortie 1976; Maier et al. 1998; Harrington 2003). Forest-dwelling reindeer might feel more protected from a predator flying above while hiding in the forest and move less to decrease the chances of being detected. Eagles usually attack and successfully kill ungulates in open areas (but see Kerley and Slaght 2013), and most predation events caused by eagles on reindeer occur in open areas (Nieminen, Norberg, and Majíjala 2011). Ultimately, the response of reindeer to aircraft observed in this study may also depend on the occasional herding of the reindeer with helicopter performed by the herders in the study area, which may have made the animals less prone to acute responses to flights, as well as on the domestication process (Reimers, Roed, and Colman 2012). Although the behavior observed in our study area does not necessarily imply increased energy expenditure due to increased movement, reindeer may
reduce feeding time by staying alert until the flight activity ends, which may negatively affect their body conditions (Åhman and White 2018).

Reindeer moved faster outside compared to inside the military test range. This result may be due to environmental differences between the two areas. Indeed, most of the military test range overlaps with a nature reserve where logging has been prohibited since 1995, making the forests denser and possibly animal movement slower. Additionally, the topography is more complex inside the test range compared to outside. Alternatively, the difference we detected in movement rates between inside and outside the test range may be due to the military exercises. Indeed, generally reindeer moved slower during military exercises performed with aircraft, except when outside the test range. In our study, the spatial extent of the military exercises was unknown. More precise data would allow for a better understanding of the effect of military exercises on reindeer movement, especially in the case of exercises performed with vehicles (where we did not detect any effect on reindeer movement), because they may be performed within a smaller area compared to aircraft exercises.

Based on the results of this study, we suggest that environmental impact assessments should always consider the combined and cumulative effects of planned and preexisting human activities and infrastructures, together with the conditions of the animals that might be affected, the location of insect relief areas, and the projected impacts of climate change. This is particularly important in arctic and subarctic ecosystems, because they are expected to experience some of the most pronounced changes in climatic conditions based on the current predictions (IPCC 2014), and the species inhabiting them are adapted to their harsh conditions, such as low nutrient availability and cold temperatures (Williams, Henry, and Sinclair 2015). Adding direct anthropogenic disturbance on top of the impacts of climate change may cause irreversible damage to northern species and ecosystems.

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