ABSTRACT. When monitoring the breeding ecology of birds, the causes and times of nest failure can be difficult to determine. Cameras placed near nests allow for accurate monitoring of nest fate, but their presence may increase the risk of predation by attracting predators, leading to biased results. The relative influence of cameras on nest predation risk may also depend on habitat because predator numbers or behaviour can change in response to the availability or accessibility of nests. We evaluated the impact of camera presence on the predation rate of artificial nests placed within mesic tundra habitats used by Arctic-breeding shorebirds. We deployed 94 artificial nests, half with cameras and half without, during the shorebird-nesting season of 2015 in the East Bay Migratory Bird Sanctuary, Nunavut. Artificial nests were distributed evenly across sedge meadow and supratidal habitats typically used by nesting shorebirds. We used the Cox proportional hazards model to assess differential nest survival in relation to camera presence, habitat type, placement date, and all potential interactions. Artificial nests with cameras did not experience higher predation risk than those without cameras. Predation risk of artificial nests was related to an interaction between habitat type and placement date. Nests deployed in sedge meadows and in supratidal habitats later in the season were subject to a higher risk of predation than those deployed in supratidal habitats early in the season. These differences in predation risk are likely driven by the foraging behaviour of Arctic fox (Vulpes lagopus), a species that accounted for 81% of observed predation events in this study. Arctic fox prey primarily on Arvicoline prey and goose eggs at this site and take shorebird nests opportunistically, perhaps more often later in the season when their preferred prey becomes scarcer. This study demonstrates that, at this site, cameras used for nest monitoring do not influence predation risk. Evaluating the impact of cameras on predation risk is critical prior to their use, as individual study areas may differ in terms of predator species and behaviour.

Key words: nest survival; cameras; Arctic shorebirds; artificial nests; predation risk; Cox proportional hazard

RÉSUMÉ. Lors de la surveillance de l’écologie de reproduction des oiseaux, les causes et les périodes de défaillance des nids peuvent être difficiles à déterminer. Des caméras placées près des nids permettent une surveillance précise du sort des nids, mais leur présence peut augmenter le risque de prédation en attirant les prédateurs, ce qui entraîne des résultats biaisés. L’influence relative des caméras sur le risque de prédation des nids peut également dépendre de l’habitat, car le nombre ou le comportement des prédateurs peut changer en fonction de la disponibilité ou de l’accessibilité des nids. Nous avons évalué l’impact de la présence de caméras sur le taux de prédation des nids artificiels placés dans les habitats de la toundra mésique utilisés par les limicoles se reproduisant dans l’Arctique. Nous avons installé 94 nids artificiels, la moitié avec des caméras et l’autre moitié sans caméras, pendant la saison de nidification des limicoles de 2015 dans le Refuge d’oiseaux migrateurs de la baie Est, au Nunavut. Les nids artificiels étaient répartis uniformément dans les cariçaies et les habitats supratidaux généralement utilisés par les limicoles nicheurs. Nous avons utilisé le modèle à risques proportionnels de Cox pour évaluer la survie différentielle des nids en fonction de la présence de caméras, du type d’habitat, de la date d’installation des caméras et de toutes les interactions potentielles. Les nids artificiels équipés de caméras ne présentaient pas de risque de prédation plus élevé que ceux sans caméras. Le risque de prédation des nids artificiels était lié à une interaction entre le type d’habitat et la date d’installation des caméras. Plus tard dans la saison, les nids placés dans les cariçaies et dans les habitats supratidiaux ont fait l’objet d’un risque de prédation plus élevé que ceux des habitats supratidiaux en début de saison. Ces différences de risque de prédation sont probablement attribuables au comportement d’alimentation du renard arctique (Vulpes lagopus), une espèce qui représentait 81 % des événements de prédation observés dans cette étude. Le renard arctique se nourrit principalement d’arvicolinés et d’œufs d’oiseau sur ce site, et s’empare des nids de limicoles de manière opportuniste, peut-être plus souvent plus tard dans la saison, lorsque ses proies préférées se font plus rares. Cette étude démontre que, sur ce site, les caméras servant à la surveillance des nids n’influencent pas le risque de prédation. Il est essentiel d’évaluer l’impact des caméras sur le risque
INTRODUCTION
Cameras are a commonly used tool for monitoring the nests of breeding birds as they can provide information about nest attendance and provisioning behaviours, parental defense, nest fate, and the identity of nest predators and their foraging behaviours (Ball and Bayne, 2012; Ribic et al., 2012; Brautigam et al., 2016; Ellis et al., 2018). Cameras can be an efficient supplement or replacement for direct human observation (Ribic et al., 2012). However, when a novel object like a camera (or other conspicuous marker) is placed near a nest, it can influence the behaviour of predators by either attracting them to the nest location (Reynolds, 1985; Renfrew and Ribic, 2003) or dissuading them (Herranz et al., 2002; Richardson et al., 2009). Cameras can also potentially affect incubation behaviour of the monitored birds in similar ways, and birds have abandoned their nests following placement of a camera (Brown et al., 1998; Renfrew and Ribic, 2003; Liebezeit, 2013). The value of cameras as a research tool is in part dependent on whether they effect changes in predator behaviours, reproductive success, or parental behaviour.

Previous studies on the survival of artificial or real shorebird nests in association with camera presence have reported either no differences (Liebezeit and Zack, 2008; McKinnon and Béty, 2009), reduced (Liebezeit, 2013), and greater survival probability (Bentzen et al., 2017). Conflicting results among studies on the influence of cameras on shorebird nest survival may be caused by differences in methodology or predator-prey dynamics at local scales. Differences among studies in methodology and study sites justify the continued evaluation of cameras as an appropriate tool to monitor shorebird nests in diverse habitat conditions given the potential influence cameras may have on nest survival and subsequently the results of studies that use them (i.e., a potential observer effect) (McKinnon and Béty, 2009). Predation is the primary cause of nest failure for most birds (Ricklefs, 1969; Martin, 1993) and Arctic-breeding shorebird nests are especially vulnerable to predation from both Arctic fox (Vulpes lagopus) and Parasitic Jaegers (Stercorarius parasiticus). In particular, the Arctic fox is the most common predator of shorebird nests, especially during years of low lemming abundance (Liebezeit and Zack, 2008; McKinnon et al., 2013; Flemming et al., 2019a). The predation rate of shorebird nests with cameras may depend on the foraging efforts of Arctic fox within different habitats, in addition to the relative availability of primary prey items within different habitat types (i.e., lemmings: Dicrostonyx and Lemmus spp., or goose eggs: Anser, Chen, and Branta spp.).

Shorebird nest predation rates vary throughout the short breeding period of shorebirds (Smith and Wilson, 2010; Liebezeit et al., 2011) and across habitats (Smith et al., 2007). Both avian and mammalian predators must provide food for their developing offspring and hunt with increasing intensity as their offspring age, which may cause predation pressure on shorebird nests to increase as the breeding season progresses (Gilg et al., 2006; McKinnon et al., 2013). Arctic foxes are opportunistic specialist predators that prefer Arvicoline prey (Elmhagen et al., 2000). Foraging efforts of Arctic fox may be concentrated in areas where preferred prey such as collared lemmings (Dicrostonyx groenlandicus) and nesting waterfowl are found (Stickney, 1991; Samelius et al., 2007). Arvicoline rodents are not commonly found in areas near the coast with saline conditions (Banfield, 1974), and predator search efforts might be lower in these habitats. As a result of this foraging habitat preference, Arctic fox may be more likely to encounter and depredate shorebird nests within sedge meadow than in habitats that provide fewer prey items. If camera presence is used as a cue to find nests, cameras may have greater influence on nests in habitats that are not subject to intense predator foraging efforts. By contrast, nests in areas already subject to intense foraging efforts may not experience a similar increase in predation pressure. Additionally, using cameras as a cue to find shorebird nests may be a learned response by predators (Reynolds, 1985), and therefore the effect of cameras on nest survival may be greatest later in the breeding season or in years following the use of cameras at nests.

Predation rates on shorebird nests are influenced by both the habitat features of the nest itself (e.g., concealment) and the behaviour of the nesting bird(s), including both incubation and nest defense behaviours (Smith and Wilson, 2010; Smith et al., 2010, 2012). In the Arctic, where the diversity of nest predators is low, artificial nests offer a useful alternative to real nests for controlled experiments in measuring relative predation risk (McKinnon et al., 2010a; Bentzen et al., 2017). Artificial nests cannot accurately mimic the presence of adult shorebirds but may be effective proxies for assessing relative predation risk of real nests in designed experiments when the predator communities that prey upon each are similar (Pårt and Wretenberg, 2002; McKinnon et al., 2010b).

Here, we use artificial nests to evaluate whether camera presence influences the relative predation risk of shorebird nests at an Arctic breeding site. We predicted that camera presence would not influence artificial nest depredation.
rates in our study area because they have been successfully deployed without an enhancement of nest predation in another similar Arctic study site (McKinnon et al., 2010a). Because visual concealment could have an important influence on nest survival, we compared concealment measures of real and artificial nests to determine how closely our artificial nests mimicked the concealment of real shorebird nests. Finally, we tested for the influence of interactions between nest placement date, habitat type, and camera presence, to ensure that spatial and temporal variability in predation rates of artificial nests did not obscure the effects of cameras and to assess differences in predation risk across the study site and season.

METHODS

Study Site

Our study was conducted from 3–27 July 2015, at the East Bay Migratory Bird Sanctuary, Southampton Island, Nunavut (63°59’ N, 081°40’ W); a site previously described by Smith and Wilson (2010). The study site is composed of diverse tundra habitats: low-lying areas near the coast are characterized by sparse vegetation, rocky substrate, and many brackish ponds. Farther inland, habitat diversity increases with low-lying moss, sedge, and grass areas, drier habitats dominated by ericaceous shrubs and forbs, and gravel ridges with sparse vegetation (Smith et al., 2007; Smith and Wilson, 2010). Five shorebird species commonly nest at the study site: Red Phalarope (Phalaropus fulicarius), White-rumped Sandpiper (Calidris fuscicollis), Ruddy Turnstone (Arenaria interpres), Semipalmated Plover (Charadrius semipalmatus) and Black-bellied Plover (Pluvialis squatarola).

Artificial Nests

We placed a total of 94 artificial nests across the study area, with 48 nests placed in sedge meadow habitats and 46 in areas influenced by saline conditions near the coast (hereafter “supratidal”) (Fig. 1). While these areas have a saline influence from salt spray, they are all well above the high-tide mark. Sedge meadows are characterized by moss-covered peat, abundant graminoid vegetation, and small hummocks while supratidal habitat is characterized by sand and rock substrate, with sparse patches of heavily goose-grazed Puccinellia spp. (Smith et al., 2007). We placed cameras at half of the nests in each habitat. Artificial nests consisted of four Japanese quail (Coturnix japonica) eggs placed uncovered on the ground without added nest lining, in a divot resembling a shallow nest scrape just large enough to accommodate the eggs (< 10 cm diameter). A nail with a small piece (0.5 cm²) of brightly coloured flagging tape was driven into the substrate and hidden beneath the eggs to allow relocation of the nest following predation events. Artificial nests were placed a minimum of 70 m from all other (known) real or artificial nests initiated during the breeding season. When placing artificial nests, we selected locations that were similar to those selected by shorebirds at this site. Sedge meadow habitats in East Bay have short vegetative cover (~ 5 cm) and low hummock heights (~ 10 cm). Within sedge meadow habitats, site characteristics used by nesting shorebirds and chosen by us included relatively high vegetation and locations on or near hummocks. In supratidal habitat, both real and artificial nests were placed on dry, silty substrate between the rocks that cover much of the area.

Within each habitat type, Day 6 Outdoors Plotwatcher ProTM trail cameras were placed at half of the artificial nests. Cameras were placed approximately 10 m from the nests, at a random bearing from the nest, and set on camouflage-painted stands (Fig. 2) with a total height of 50 cm. Cameras were set to take continuous 24-hour time-lapse videos by capturing one photo every 5 sec. Following their placement, we checked artificial nests once every two days until failure or to 22 days (a typical incubation duration for shorebirds at this site), at which point the nest would be considered successful. We identified nest predators by watching videos and observing photos that contained predators with eggs in their mouths or with their head at the nest location on the nest failure date.

We placed nests in both sedge meadow and supratidal habitats throughout the period from 3–19 July 2015. As older nests were depredated and cameras became available, new nests were constructed and cameras were moved. When cameras were available, we deployed new nests evenly distributed between habitats and camera treatments on each date. Nests placed from 3–11 July were considered “early” and nests placed from 13–19 July were considered “late” nests. In sedge meadow habitat, we constructed 16
and 32 nests in the early and late periods, respectively. In supratidal habitat we constructed a total of 10 artificial early nests and 36 late nests.

Concealment

Microhabitat characteristics around shorebird-nesting sites can provide visual concealment from predators. White-rumped Sandpipers and Red Phalaropes construct nest scrapes with high concealment while the other species that nested within our study area rely on early detection of predators, with nests with little vegetative cover (Smith et al., 2007, Flemming et al., 2019b). We used three, 12 cm diameter disks, marked with a grid system and fastened together at right angles to create an identical silhouette in four directions and from above (Smith et al., 2007). Nest concealment was measured by placing the disk into a nest cup, observing the disks from a distance of 4 m and 40 cm above the ground (mimicking the observation height of the Arctic fox) from each cardinal direction and directly above the nest, and estimating the percentage of the disks’ grid system covered by the vegetation, rocks, or hummocks surrounding the nest cup.

Statistical Analysis

Cox proportional hazard regression was used to determine whether camera presence, habitat type, or date of nest deployment was related to the survival time for artificial nests (Cox, 1972; Fox, 2001). We selected Cox proportional hazard regression because it is a semi-parametric method for estimating hazard ratios without assuming a baseline hazard function. This method also allows for right censoring (i.e., unknown fate of individuals surviving beyond the end of the study). The Cox model is appropriate here because nests were placed across multiple days and some nests survived to the end of this study. The initial model contained camera presence, habitat type, and placement date with all possible interactions. Non-significant factors were removed until the final model was reached. The Cox model assumes that the hazard ratio is proportional to the survival function (i.e., differences in survival probability are proportional among groups across time). We assessed this assumption statistically and graphically using Schoenfeld residuals analysis (Fox, 2002). Survival estimates were generated from the fit Cox model following Laine and Reyes (2014). We compared concealment of artificial and real nests located within sedge meadow and supratidal habitats using an independent t-test. All statistical analyses were conducted in R version 3.2.2. The Cox proportional hazard model and figures were created using the “Survival” and “ggplot2” packages respectively (Therneau, 2015; Wickham, 2016; R Core Team, 2017).

RESULTS

Most of the artificial nests failed within two days (71.7%), and none (with or without cameras) survived beyond 14 days. Time-lapse footage from the 47 nests with cameras captured 43 predation events; four predation events were missed by the cameras. Arctic foxes were by far the most common predator of artificial nests (81%), with Herring Gulls (Larus argentatus) and Parasitic Jaegers accounting for 14% and 5% of recorded predation events, respectively (Table 1).

The proportional hazards assumption was met for all variables included in the model ($p > 0.05$), which indicates that survival probability was proportional among treatment groups (camera presence, placement date, and habitat). The model was also assessed for and satisfied the general linear model assumptions of influential values (i.e., individual points driving measured relationships) and non-linearity (i.e., residuals fit a normal distribution) graphically (Fox, 2002). Camera presence had no effect on the failure rate of artificial nests ($z = 1.17, p = 0.24, \beta = 0.80, CI = -0.53, 2.13$; Fig. 3). The interaction between habitat type and initiation

FIG. 2. A Day 6 Plotwatcher ProTM camera deployed on Southampton Island. Rocks are placed on the wooden stand to stabilize the camera in high winds. Cameras were placed 10 m from nests and the total height is approximately 50 cm.

FIG. 3. Survival estimates (± SD) of artificial nests placed with (solid line) and without (dotted line) cameras associated, as estimated by Cox proportional hazard regression. The probability of survival is estimated at 2-d intervals coinciding with nest checks.

FIG. 3. Survival estimates (± SD) of artificial nests placed with (solid line) and without (dotted line) cameras associated, as estimated by Cox proportional hazard regression. The probability of survival is estimated at 2-d intervals coinciding with nest checks.
TABLE 1. Number of artificial nests depredated by each of three predator species, within sedge meadow and supratidal habitats at East Bay, Southampton Island, Nunavut, in July 2015. Predators were identified from camera footage at nests.

| Predator           | Sedge meadow | Supratidal | Total |
|--------------------|--------------|------------|-------|
| Arctic fox         | 15           | 19         | 35 (81%) |
| Parasitic Jaeger   | 2            | 0          | 2 (5%)  |
| Herring Gull       | 5            | 1          | 6 (14%) |
| Total              | 22           | 20         | 43 (100%) |

date was a significant predictor of failure rate ($z = -3.30$, $p < 0.001$, $\beta = -0.15$, CI = $-0.24$, $-0.06$). Nests placed both early and late in July within sedge meadow habitat had similarly low survival estimates (Figs. 4, 1). Nests placed early in July within supratidal habitat had higher survival estimates while nests placed later within supratidal habitats had lower survival estimates, comparable to the survival estimates for sedge meadow nests (Fig. 4). Real shorebird nests were more concealed than artificial nests within supratidal ($t = 3.61, p < 0.001$; real (mean ± SD): 17 ± 3.4%, n = 11; artificial: 5 ± 2.5%, n = 46) and sedge meadow ($t = 4.46, p < 0.0001$; real (mean ± SD): 25 ± 4.2%, n = 25; artificial: 14 ± 2.1%, n = 48) habitats.

**DISCUSSION**

We did not detect an effect of camera presence on the survival of artificial nests. Despite the relatively conspicuous appearance of cameras placed on the tundra, their association with artificial nests did not affect the rate that nests were depredated. Thus, common shorebird nest predators such as Arctic fox apparently did not use cameras as cues to locate shorebird nests. Artificial nest survival was, however, dependent on placement date and the habitat in which nests were located.

Artificial nests placed in sedge meadow habitat experienced lower survival than those placed in supratidal habitats regardless of the date. Survival of artificial nests in supratidal habitat was dependent on the date they were set; nests placed early in July survived longer than those placed later. The differential survival of nests based on habitat and initiation date likely reflects the foraging efforts of Arctic fox at the study site, the most commonly identified predator of real and artificial shorebird nests at this site and others (Liebezeit and Zack, 2008; McKinnon and Béty, 2009). Lemmings are the primary prey item of Arctic fox, and foraging efforts of Arctic fox are more concentrated in sedge meadow where lemmings occur than in supratidal habitat where lemmings are generally absent (Lecomte et al., 2008). Because of this higher foraging effort, foxes probably encounter incidental prey such as artificial nests more frequently in sedge meadow habitat.

By contrast, foxes avoid habitats where numerous ponds make travel difficult for them (Lecomte et al., 2008). The supratidal habitats at East Bay are dominated by ponds. Despite their ability to swim, Arctic foxes avoid deepwater bodies while foraging (Strub, 1992; pers. observ.) and this standing water early in the season may inhibit or deter Arctic fox from extensive foraging in supratidal habitat. However, as the summer progresses, the ponds evaporate, increasing land connectivity. These drier conditions provide greater access to alternative prey for Arctic fox, including Brant (Branta bernicla) and Cackling Goose (Branta hutchinsii) nests (Peterson, 1990; Anthony et al., 1991). If foxes redirect foraging efforts to supratidal habitats later in the season, the probability that shorebird nests would be incidentally discovered would increase. Furthermore, progression of the breeding season is accompanied by increased energetic demands for all reproductively active adult predators as they must provide food for their offspring (McKinnon et al., 2013). Predator incursions into secondary foraging areas may also be driven by the increased energetic demands of parental care and the associated increase in foraging effort. Increased foraging efforts by Arctic fox in secondary habitats like supratidal areas may be especially important if feeding success on their primary prey (Arvicoline rodents) is low in the sedge meadow habitats.

This study necessarily assumes that artificial nests are a useful proxy for real nests when determining predation risk. Real shorebird nests are simple scrapes in the substrate and thus can be simulated more easily than passerine nests. The artificial nests used in this study were less concealed than real shorebird nests at the same site. Greater visual exposure of nests to predators may have resulted in elevated rates of predation in artificial nests. Predation rates of artificial nests are expected to be further elevated because artificial nests lack the benefit of parental defense behaviours (Nguyen et al., 2006). However, lack of parental activity at the artificial nests could provide fewer cues to nest predators at these nests than at real nests. Importantly, artificial nests cannot be used as proxies for measuring absolute real nest survival but they are effective for measuring relative predation risk, especially in habitats with relatively simple predator communities (Pärt and Wretenberg, 2002). Our results indicate that camera-monitored and control (no camera) artificial nests did not
differ in their survival; this pattern of no relative survival difference could apply to real nests. However, we cannot rule out that parental defense behaviours either alone or influenced directly by camera presence interact to influence predation of real nests.

Most artificial nests in the present study were depredated by Arctic fox, which is consistent with previously reported observations at this study site of predators on shorebird nests (Smith and Wilson, 2010). Both Herring Gulls and Parasitic Jaegers depredated nests in sedge meadow, but only one nest was taken by a Herring Gull in supratidal habitats (Table 1). Herring Gulls have not been documented as an important predator of real shorebird nests at this site (Smith et al., 2012) and the nest predation events recorded here may not be independent. Based on the camera evidence, five of the six predation events by Herring Gulls occurred within 3 d, and 0.9 km of each other; these predation events may be attributed to a single individual and thus increased the estimate of avian predation overall and in sedge meadow habitat (Table 1). Despite the possible inflation of Herring Gull predation events, the artificial nest predator guild closely corresponds to that of real nests at this study site (Smith and Wilson, 2010).

Our study was conducted during a year of low lemming abundance and low shorebird nest success (pers. observ.). Smith et al. (2007) found that shorebirds experienced increased nest mortality during a year of low lemming abundance, likely due to Arctic fox foraging for bird eggs when their primary prey was not available. Because of their increased reliance on alternative prey during this study, we expected that nest predators would be more sensitive to cues like cameras that may improve foraging efficiency during years of low primary prey abundance than during years when primary prey are abundant. Artificial nest survival was very low in our study and nearly three-quarter of nests we placed were depredated within two days. It is possible that the rapid predation of artificial nests could have limited our ability to detect an influence of camera presence on survival because of low exposure time. However, given that differences in survival between habitats and placement date were detected, it is unlikely that we failed to detect an influence of camera presence on artificial nest survival in our study.

The present study examined the effect of cameras on predation risk for artificial nests of Arctic-breeding shorebirds. We have shown that the risk of predation for artificial nests varied across two Arctic coastal habitats and across the breeding season in our study year with a higher risk of predation in sedge meadow throughout the breeding season. We cannot eliminate the possibility that this pattern could shift among years depending on the relative availability of prey and the predator responses to prey. While real nest survival was not assessed in this study, artificial nests can be effective proxies of real nests for measuring relative predation risk and we did not find a difference in risk associated with camera presence (McKinnon et al., 2010b). This study provides support for the continued use of cameras as a monitoring tool at this site. Our results also highlight that differences in study site and design may influence results (Bentzen et al., 2017). We recommend that future use of cameras at other study sites be preceded by pilot studies such as the present one to determine any influence of cameras on predator behaviour that could influence predation rate of real nests.

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