Are predation rates comparable between natural and artificial open-cup tree nests in boreal forest landscapes?

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Abstract

Nest predation studies often use artificial nests to secure sample sizes and nest distribution patterns that allow empirically testing differences in predation rates between ecological units of interest. These studies rely on the assumption that natural and artificial nests experience similar or consistent relative predation rates across ecological gradients. As this assumption may depend on several factors (for example differences in predator community, nest construction, parental care patterns), it is important to test whether artificial nests provide adequate and comparable estimates of predation rates to natural nests. In this study, we compare predation rates of above-ground natural open-cup nests, artificial nests and natural nests with artificial eggs along a forest gradient from edge to interior (interior, transition zone and edge) and within two nest visibility classes (visible and concealed). Our aim was to determine whether nest structure affects comparability between nest types along these ecological gradients in boreal forests. Our results indicated important contributions of nest type, nest visibility and location along the forest edge-interior gradient, but no variable had strong significant effects on predation rates, except exposure time that showed lower predation rates at longer exposure times. Predation rates in visible and concealed nests remained similar for all nest types, but not along the forest edge-interior gradient. Here, artificial nests showed much lower predation rates than natural nests, whereas natural nests with artificial eggs tended to have higher predation rates than natural nests. We conclude that artificial nests in boreal forests represent an adequate measure of relative nest predation risk in open-cup natural nests along some ecological gradients, but results on predation rates along forest edge-interior gradients obtained from artificial nests should be interpreted with care.

Introduction

Nest predation is the most important factor causing reproductive failure (e.g., [1,2]), and therefore has been studied extensively. Nest predation studies often rely on the use of artificial nests to secure sample sizes and nest distribution patterns that allow empirically testing differences in predation rates between habitats, predator removal treatments or other units of
interest [3–6]. Many of these studies rely on the assumption that natural and artificial nests experience similar predation rates, or that relative differences in predation rates between units are consistent between the two nest types [7]. However, the validity of this assumption has been questioned repeatedly (e.g., [8–15]). Although some studies comparing natural and artificial nests do suggest similarity in absolute predation rates [16–18], many studies show that artificial nests experience higher [4,15,19,20] or lower [9,10,12] predation rates than natural nests. These discrepancies in predation rates between natural and artificial nests have been attributed to several factors [3]. For example, different predator species may prey on artificial and natural nests [21], but also nest construction and visibility of artificial nests and characteristics of experimental eggs may increase or decrease the likelihood of predation [22,23]. Furthermore, absence of parental care in artificial nests can increase or decrease predator activity compared to natural nests [24], while increased nest density due to surplus of artificial nests may increase predation rates [25]. Finally, investigator activity may affect predation intensity of both artificial and natural nests [26,27]. Despite these biases, relative predation rates between natural and artificial nests can be comparable across ecological gradient, such as landscape fragmentation and forest patch size variation [4,28]. Regardless of the potential pitfalls of assuming artificial and natural nests provide similar information, artificial nests are still used in nest predation studies to ensure adequate sample sizes and nest distribution and to standardise conditions between nests. This highlights the need to identify situations where artificial nests provide adequate and comparable estimates of predation rates and where comparability between natural and artificial nests are misleading, as previously also suggested by Villard and Părt [29].

In many of the studies where predation rates are compared between artificial and natural nest, the focus seem to be on the inconsistencies in treatment or experimental effect on predation rates rather than identifying the source of the difference. The latter is important for determining in which contexts artificial nests are a useful tool for estimating predation impact on bird species. For example, nest predation may be comparable between artificial and natural nests during the incubation period [12], or predation rate responses to habitat or forest edge differ between artificial and natural nests (e.g., [30]). The studies that have identified why predation rates differ between artificial and natural nests often highlight that predator identity differs between nest types [11,21,28]. However, in areas where corvids are the main predators, very similar nest predation rate between natural and artificial nests have been reported [16].

Different methods are employed to make artificial nest more similar to natural, e.g., using smaller clay or plasticine eggs instead of quail eggs [3,22,31–34], using old natural nests [8,35,36], old nest sites [37], or nesting densities and locations based on natural nest patterns [4,35,37,38]. However, little emphasis has been put into determining whether simulating natural nests in different manners actually leads to more comparable predation rates compared to natural nests.

Here we compare predation rates after 10 and 25 days exposure between natural nests and two types of artificial nests to determine whether nest structure (natural nest versus wire basket dressed with dry grass) affects comparability between nest types in boreal forests, while controlling for exposure time. Artificial nest types include artificial nests placed systematically in a study site at the same time as natural nests and old natural nests from the corresponding breeding season equipped with artificial eggs. Avian predators mainly cause nest predation in the study region with corvids dominating the avian predator community [39]. If differences between nest types are smaller when corvids are the most common predator (see [16]), we predict similar nest predation rates between nest types in our study. The comparability of predation rates between nest types was tested along ecological gradients known to affect nest predation rates in natural nests, namely nest visibility, and location along a forest edge-interior.
gradient (edge, transition zone and interior). We do not a priory expect any differences in nest predation rates in response to the ecological factors between nest types.

Material and methods

Study area

All study areas are located in the western part of the mid-boreal forest zone in Norway. It is a mosaic landscape with forest, forest patches, farmland, lakes and ocean, mires and human habitations. Potential predators at the sites include different corvid species, mustelids, squirrels, foxes and badgers (see details in [39]). The study includes five sites located in forest patches, which were monitored between 2004 and 2006 (percentage forest cover and arable land within a 5.5 km² area at each site is presented in Table 1. Land cover percentages were obtained from OSM Landuse Landcover based on OpenStreetMap-data (osmlanduse.org [40])). Previous studies suggest that predation rates on both natural and artificial nests in this region are mostly similar between different study years [39], using data from multiple years should therefore not introduce too much bias due to between year variation in predation rates. Sites were distributed between 63.4 and 63.6°N, and 10.5 and 11.2°E within central Norway (Table 1, Fig 1).

The forests are dominated by Norwegian spruce *Picea abies* (59% of total forest in Trøndelag county between 2004 and 2008, data from Statistics Norway: www.ssb.no/en), with inclusion of Scots pine *Pinus silvestris* and deciduous tree species (22% pine and 19% deciduous trees between 2004 and 2008, data from Statistics Norway: www.ssb.no/en). We defined forest locations along a gradient from interior to edge: forest interior (> 50m from the forest edge), edge (within 2m from the forest edge) and transition zone between forest interior and edge (2 – 50m from edge inside the forest). The definition of forest interior at distance > 50 m was based on the meta-analysis results of Batáry and Báldi [30], which showed that edge effects were negligible beyond 50 m. The choice of including a transition zone was based on the same study, which showed that edge effects may be strong at least up to 25 m from the edge. Forest edges were mainly natural edges along agricultural land, with maximum 3 nests across sites located along clear-cuts.

Nest types include natural nests, systematically placed artificial nests (artificial), and natural nests with artificial eggs (nat w/artegg). Total nest numbers, position of sites on mainland or island and % cover of forest and arable land are also provided. Land use data are percentages of available land area, and excludes sea as water bodies, but includes lakes and rivers. Percentages are provided for a land area of ca 5.5 km² per site, which defines an area that provides representative information on surrounding land cover types while avoiding extensive overlap between sites. Percentage land cover was obtained from OSM Landuse Landcover based on OpenStreetMap-data (osmlanduse.org [40]) and excludes land areas that are not provided with landuse tags in OpenStreetMap. The reported values therefore represent approximate estimates and not absolute values.

Table 1. Distribution and number of nest types between sites.

| site no | % forest / arable | mainland/island | natural | artificial | nat w/artegg | total |
|---------|------------------|----------------|---------|------------|--------------|-------|
| 1       | 16.3 / 75.6      | mainland       | 1       | 30         | 27           | 58    |
| 2       | 72.7 / 1.6       | mainland       | 20      | 8          | 12           | 40    |
| 3       | 51.4 / 41.4      | island         | 12      | 14         | 11           | 37    |
| 4       | 44.3 / 49.7      | island         | 12      | 14         | 11           | 37    |
| 5       | 13.5 / 38.5      | mainland       | 12      | 14         | 11           | 37    |

https://doi.org/10.1371/journal.pone.0210151.t001
Experimental design

We monitored predation occurrence on natural and artificial eggs (see details below) after 10 and 25 days from three types of nests; natural nests, systematically placed artificial nests (hereafter artificial nests) and natural nests with artificial eggs. Exposure of 10 days represented the exposure time used in most artificial nest studies by the research team (see [39,41]), while exposure of 25 days represent the average incubation plus nestling period of natural nests. All nests used in this study were classified into three categories of visibility based on how visible nests were at different distances: 1) high cover, nests hardly visible at 0.5m distance, 2) medium cover, nests visible at 1m, and 3) low cover, nests visible at 1-5m. As there were very few natural nests (n = 5) and natural nests with artificial eggs (n = 4) that were classified as high cover, we combined the two classes “high” and “medium” cover to one class and operate now with concealed and visible nests. Only natural nests and natural nests with artificial eggs originating from birds of the families Turdidae and Fringillidae were used in this experiment, as they are the most common species in the habitat used for this study.

Fig 1. Map of study site locations relative to Northern Europe. Site numbers are provided in the map close up of the study area (see Table 1 for site details). For reference, the location of Trondheim city is represented by a star. Made with Natural Earth. Free vector and raster map data @ naturalearthdata.com.

https://doi.org/10.1371/journal.pone.0210151.g001
Natural nests were found before and during egg laying (nests active during May and June) and the fate of individual nests were monitored until fledging (see Table 2 for a complete species list). Natural nests located within fieldfare *Turdus pilaris* colonies were not included while fieldfares were breeding. This was done because i) their colonial breeding and intensive nest defence both against nest predators and human intruders can disturb experiments (e.g., [42,43]), ii) in most colonies we had data only on fieldfare nests, and iii) we did not have natural nests with artificial eggs within colonies or colonies within island sites. Most nests were located from 0.5 m to 6 m above ground (97%), with a few nests located below 0.5 meter or above 6 meter (3%).

Artificial nests were distributed in May, June or July with a minimum distance of 100 m between each nest. All nests were open-cup nests with 11 cm outer diameter placed on the in trees 1–1.5 m above the ground within all forest location categories, since all natural nests and natural nests with artificial eggs were located in trees. Nests were designed for a common bird species with similar sized nests as the artificial nests and do not model a specific bird species. This allows us to compare predation rates of most natural bird nests available at each site. Any potential effects of bird species on nest design are checked by adding natural nests with artificial eggs as a separate nest type (see description below). Nests were made of wire baskets lined with dry grass and/or moss, and were attached to tree branches with iron wire. The metal was visible from beneath in tree nests, but completely hidden by nesting material from above, and therefore the appearance was quite similar to a natural nest.

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**Natural nest with artificial eggs** were natural nests predated during or after egg laying or with successful fledging, in which we placed artificial eggs after the main breeding season (July, all natural nests used were active during May and June). Natural nests abandoned before egg-laying were not included since natural nests with artificial eggs might be abandoned for other reasons than nest predation.

In artificial nest and natural nests with artificial eggs, we put one fresh quail *Coturnix coturnix* egg and one plasticine egg. The plasticine eggs were a mixture of grey and green plasticine and had the similar size and shape as the quail eggs. We attached the plasticine egg with a metal wire with a barb inside the egg, and the metal wire was attached to a branch under the nest or in the surrounding vegetation. The plasticine and quail eggs were similar in size to many relatively common species in the areas surrounding the study sites, nesting in trees (e.g., thrushes and common wood pigeon *Columba palumbus*), and the most numerous nest predators in our areas will easily predate the actual egg sizes. A nest was considered predated if at least one of the eggs had been removed or damaged by a predator.
A few plasticine eggs had marks made by small mammals (different mouse or vole species), snails or tits, and a few quail eggs had small scratch marks without being destroyed. We did not consider these groups as predators because they cannot destroy quail eggs and are not potential predators for eggs at this size [22, 32, 33, 44]. Therefore, the few eggs with marks only from small mammals, snails or tits were recorded as not being predated. Thus, we also avoided the potential issue the great multi-annual variation in abundance of small mammals in boreal forests can have on predation rates (e.g., [45, 46]), especially in nests not guarded by parents. However, small mammals may predate small eggs from for example warblers [47], but the importance of small mammals as nest predators are probably overestimated in artificial nest experiments when plasticine is used [48]. Disregarding these nests as predated would potentially lead to smaller differences in predation rates between natural nests and artificial nests in our analyses, which makes our inferences conservative at worst. We have no grounds to believe that this would affect the response to ecological factors included in the analysis, and should therefore not affect the relative comparability between artificial and natural nests.

We did not use rubber boots or gloves to avoid leaving scents at the nest, since several studies show that there are small chances for predators to learn, by smell or visually, to associate human scent to artificial nests [44, 49]. This choice should not lead to large biases, since the same method was used in all sites and all years. Nest locations were not marked. No specific permissions were needed for the fieldwork since we only observe bird’s nests and use artificial nests to assess predation rates. The field work did not involve endangered or protected species.

**Data analyses**

Predation rates in natural and artificial nest types were compared using a logistic regression fitted as a generalised linear mixed model (GLMM) within a Bayesian framework (package **rstan**-**darm** [50]) in R statistical software (version 3.4.3, [51]). We use a logistic regression model for this analysis since the posterior resampled distribution of a binomial regression model with a complementary log-log link and exposure time as offset showed poor fit (see details on the model building procedure in S1 Appendix). Predation events (0 for survived and 1 for predated) recorded after 10 or 25 days exposure was entered as the dependent variable in the logistic regression. Some nests were checked after both 10 and 25 days exposure. Of these, nests predated before 10 days were defined to have 10 days exposure, whereas remaining nests were defined to have 25 days exposure. Nest type (natural, artificial and natural with artificial eggs), forest location (forest interior, edge and transition zone), visibility (extra cover, good cover and visible), and the interactions between nest type and the other main effects were added as predictor variables. To control for the potential confounding effects of site location on mainland or island, nest activity month and taxonomic family (thrushes and finches), island, month, family and the interaction between family and nest type were included as predictor variables. Finally, to control for differences in exposure time between nests, we added z-transformed exposure as a predictor variable. The variable exposure time was z-transformed to improve model convergence and obtain reliable parameter estimates (models with non-transformed exposure time provided predation estimates close to 1). Site no was used as a random effect in the model.

As weakly informative prior distributions for the intercept and the vector of regression coefficients we chose the default normal distribution with mean = 0 and standard deviation = 1. We also chose to use the default prior for the covariance structure, which is developed to be robust for common applied regression problems [52]. The covariance matrix is decomposed into a vector of variances and a correlation matrix, which summarises the covariation of slopes and intercepts in a standardised form. Default settings describe a joint uniform prior for the correlation matrix. The vector of variances is set equal to the product of a simplex vector
described by a symmetric Dirichlet prior. With the default settings of a concentration parameter at 1, this prior is jointly uniform over the space of simplex vectors.

Posterior distributions were obtained by fitting the model described above with function `stan_glmer` in package `rstanarm`, which uses a Hamiltonian Markov Chain Monte Carlo algorithm to obtain samples drawn from the posterior distribution. Default settings fits four Markov chains with 2000 iterations each (thinning = 1), half of which are discarded as a warm-up used to give the algorithm time to find the target posterior area. We assessed model fit by performing posterior predictive checks and sensitivity analysis following suggestions by Gelman et al. [53]. Model convergence, effective posterior sample size, Monte Carlo standard error and posterior predictive checks were performed using the `shinystan` R package [54], which provides a graphical user interface for exploring models fit using MCMC-techniques. All predictors had $R = 1$, Monte Carlo standard error $< 0.1$ and number of effective samples $> 1210$. In the sensitivity analysis, we tested the impact of choice for prior distributions on intercept and the vector of regression coefficients (see S1 Appendix for further details on the sensitivity analysis and model evaluation).

The means of simulated values from the joint posterior distribution of the model parameters were used as estimates, and the 2.5% and 97.5% quantiles as the lower and upper limits of the 95% credible intervals. Posterior probabilities of linear predictor effects being larger (or smaller) than zero were calculated by the proportion of simulated values from the posterior distribution being higher (or smaller) than zero. To simplify language in the results, we declare an effect to be significant if the posterior probability of being positive (or negative) is larger than 0.97, and we declare an effect to signify a trend if the posterior probability of being positive (or negative) is larger than 0.75. These values correspond to the limits of the 95% and 50% credible intervals, respectively, for the model not containing zero.

Results

Observed average predation rates were high (Table 3). The intercept value from the logistic regression model (Table 4) corresponded reasonably well with the observed predation rates with a back-transformed estimate of predation rate at 88.1%. Of the main predictor effects, only exposure time showed a significant negative effect with nests exposed for 25 days having lower predation rates than nests with a 10-day exposure time (Fig 2, Table 4). However, there was indications that nest predation rates depended on nest type, location along the forest gradient, nest visibility and observation month (Fig 2). Artificial nests had a probability of 0.77 of having lower nest predation rates than natural nests, and the probability of predation rates

| Nest type             | Forest gradient | Predation rates (mean ± se) in covered nests | Predation rates (mean ± se) in visible nests |
|-----------------------|-----------------|---------------------------------------------|---------------------------------------------|
| Natural               | Interior        | 0 ± 0                                        | 1 ± 0                                       |
|                       | Transition      | 0.833 ± 0.167                               | 0.667 ± 0.211                               |
|                       | Edge            | 0.864 ± 0.052                               | 0.857 ± 0.097                               |
| Natural w/art. egg    | Interior        | 0 ± 0                                        | 0.824 ± 0.095                               |
|                       | Transition      | 0.75 ± 0.25                                 | 0.844 ± 0.065                               |
|                       | Edge            | 0.857 ± 0.060                               | 0.571 ± 0.202                               |
| Artificial            | Interior        | 0.938 ± 0.063                               | 1 ± NA                                      |
|                       | Transition      | 0.563 ± 0.128                               | 0.733 ± 0.118                               |
|                       | Edge            | 0.813 ± 0.101                               | 1 ± 0                                       |

https://doi.org/10.1371/journal.pone.0210151.t003
being lower in the transition zone than in the forest interior was 0.83 (Table 4). Visible nests was 95% more likely to have higher predation rates than covered nests, and nest predation was 77% more likely to be higher in July compared with May (Table 4).

Relative differences in nest predation rates seemed to be similar between nest types between levels of nest visibility (Fig 2, Table 4). However, nest predation rates were not fully comparable within the forest gradient. Although nest predation rates seem to be similar between nest types when comparing predation rates in the forest interior and at the forest edge, both artificial nests and natural nests with artificial eggs showed different predation rates in the transition zone (Fig 2). Natural nests with artificial eggs tended to have higher predation rates than natural nests in the transition zone ($P(\beta > 0) = 0.94$), whereas predation rates in artificial nests were significantly lower compared with natural nests in the transition zone (Table 4). The only other interaction with evidence of significance was the interaction between nest type and taxonomic family. Natural nests with artificial eggs belonging to thrushes had a probability of 0.78 of being lower than natural nests (with artificial eggs) of finches (Table 4).

Table 4. Posterior effect estimates and credible intervals for fixed and random effects.

| Parameters | Mean | 2.5%   | 97.5%  | $P(\beta > 0)$ |
|------------|------|--------|--------|----------------|
| intercept (natural nest, forest interior, covered) | 1.974 | 0.208  | 3.671  | 0.984          |
| nesttype_nat.artegg | 0.127 | -1.206 | 1.466  | 0.579          |
| nesttype_artificial | -0.533 | -1.915 | 0.834  | 0.227          |
| forest.gradient_transition | -0.589 | -1.808 | 0.602  | 0.168          |
| forest.gradient_edge | 0.351 | -0.822 | 1.510  | 0.726          |
| visibility_visible | 0.822 | -0.157 | 1.805  | 0.947          |
| family_thrush | -0.130 | -1.159 | 0.962  | 0.400          |
| month_June | -0.060 | -1.090 | 0.962  | 0.461          |
| month_July | -0.469 | -1.701 | 0.742  | 0.226          |
| island_yes | 0.255 | -1.102 | 1.437  | 0.686          |
| z-transformed exposure | -1.463 | -1.971 | -0.995 | 0              |
| forest.gradient_transition:nesttype_nat.artegg | 1.065 | -0.283 | 2.391  | 0.940          |
| forest.gradient_transition:nesttype_artificial | -1.448 | -2.864 | -0.087 | 0.018          |
| forest.gradient_edge:nesttype_nat.artegg | -0.327 | -1.644 | 1.018  | 0.313          |
| forest.gradient_edge:nesttype_artificial | -0.001 | -1.359 | 1.435  | 0.501          |
| Visibility_visible:nesttype_nat.artegg | -0.004 | -1.259 | 1.318  | 0.492          |
| Visibility_visible:nesttype_artificial | 0.004 | -1.369 | 1.384  | 0.495          |
| Family_thrush:nesttype_nat.artegg | -0.489 | -1.679 | 0.679  | 0.219          |
| Random effect site 1 | 0.104 | -0.814 | 1.463  |                |
| Random effect site 2 | 0.423 | -0.369 | 1.893  |                |
| Random effect site 3 | -0.272 | -1.443 | 0.700  |                |
| Random effect site 4 | 0.404 | -0.479 | 2.083  |                |
| Random effect site 5 | 0.184 | -0.741 | 1.550  |                |
| Sigma | 0.603 | 0  | 3.388  |                |
| Mean_PPD | 0.806 | 0.743 | 0.862  |                |
| Log-posterior | -127.634 | -136.642 | -121.062 |                |

Mean effect estimates with 95% credibility interval of nest type, forest gradient location, nest visibility, taxonomic family, month, island and exposure time on nest survival rates. For fixed effects, we show the probability that effect slopes are above zero. Slopes of effect estimates with a probability between 2.5% and 97.5% of being above zero are highlighted with bold text; estimates with probability between 25% and 75% of being above zero are highlighted with bold italic text. The intercept represents baseline categories of predictor levels, which are listed in parentheses in order nest type, forest gradient location and visibility, for which other predictor levels (factorname_levelname) are compared with.

https://doi.org/10.1371/journal.pone.0210151.t004
Discussion

Predation is a major cause of nest failure in birds [1,2] and can have an enormous impact on life history evolution [55]. An interplay of several factors causes spatial and temporal variation in nest predation rates, involving characteristics of the birds themselves, the predators, and the surrounding habitat. Behaviour of prey bird species can affect predation rates through choice of nesting sites [37], nest density [49,56], nest concealment [57–59] and parental activity at the nest [60,61]. Predator characteristics important for nest predation risk include for example predator abundance and breeding phenology [49,62], community composition [48,63] and availability of alternative prey [64,65]. Yet again, factors related with landscape features, such as habitat differences, fragmentation or edge effects, can have strong impact on local predator abundances and predation pressure [30,66,67]. Often, patterns of variation in nest predation are evaluated based on studies using artificial nests instead of natural nests. In all the above-mentioned situations that determine nest predation rates, artificial nests may or may not experience the same patterns of predation as natural nests [12,15,17,18]. In this paper, we have evaluated how well artificial nests represent nest predation rates in natural nests according to nest visibility and location along a forest edge-interior gradient.

The findings of this study reveal that estimated nest predation rates were high, corresponding to high observed predation rates, but similar to predation rates observed for artificial tree nests in the study region in some years [39]. Nest predation rates tended to differ according to nest type (natural, natural with artificial eggs, and artificial), location along a forest gradient (interior, transition, edge), and nest visibility. Of the controlling factors, nest predation rates tended to differ according to month of observation (lower predation in July than May) and differed depending on exposure time (lower predation rates in nests exposed for 25 days than nests exposed for 10 days), whereas predation rates in natural nests with artificial eggs tended to be lower in nests of thrushes compared with nests of finches. When controlling for exposure
time, predation rates of artificial nests tended to be lower than predation rates in natural nests, which correspond to results from other studies [9,10,12]. Observed predation rates (exposure time not controlled for) were similar between natural and artificial nests, highlighting the importance of controlling for exposure time when assessing nest predation rates. Interestingly, natural nests with artificial eggs showed similar nest predation rates as natural nests, even though artificial eggs were added to natural nests in July, after the breeding season of the original residents. Relative timing of the treatment can affect the comparability of predation rates between natural and artificial nests due to seasonal variation in predation rates (e.g., [68,69]), and predation rates of artificial nests in our study system follow a seasonal pattern with peaks in June/July [41]. We could therefore expect lower predation rates in natural nests with artificial eggs compared with natural nests based on the timing of exposure. One possible explanation for our result is that some of the natural nests used in the experiment were predated before reuse, which could bias our results towards higher predation rates if predators are more likely to visit these nests later in the season. Although nest design can be an important feature determining predation rates (reviewed in [70]), the likelihood of predation in re-used nests that already experienced predation does not seem to have been investigated. Another option is that the expected pattern of lower predation rates due to season only apply to thrushes and not to finches, given the trend in the interaction effect between nest type and taxonomic family (technically between natural nests and natural nests with artificial eggs as artificial nests were not labelled according to taxonomic family). All nest types varied similarly with respect to nest visibility, but not according location along the forest edge-interior gradient.

We found predation rates to be significantly lower after 25 days exposure than after 10 days exposure, which seems counterintuitive. However, this result is partly caused by study design and data management choices related with categorisation of nests within 10- or 25-days exposure. Some nests were observed after both 10 and 25 days, of which nests predated before 10 days were categorised as being exposed for 10 days and the remaining nests were treated as being exposed for 25 days. This causes a skew in predation rate estimation towards being higher during the 10-day period as predated nests observed both exposure periods are censored out and not included in the estimations of predation rate after 25-days exposure. This is evident if we redefine all those nests as being exposed for 25 days, at which point the effect of exposure is largely reduced (S1 Fig). Therefore, our result may suggest that predation rates are higher during a time period representing the incubation period compared with predation rates during the nestling period, assuming if nests that survive during the first 10 days have lower risk of being predated later. However, most of the nests within the 10-day exposure category are artificial nests (or among nests predated before 10 days, S1 Table), whereas natural nests and natural nests with artificial eggs are more evenly distributed between the exposure time categories (S1 Table). Under this scenario, the results may indicate that artificial nests experience higher predation rates than natural nests when differences in exposure time are not controlled for. Note, however, that when redefining all double-observed nests as having exposure time of 25 days, predation rates of artificial nests do not differ significantly from rates in natural nests but the mean point estimate is positive rather than negative (S1 Fig). Alternatively, the remaining small effect of exposure time may relate with, for example, parental activity at natural nests that reduces the relative predation risk in these nests compared with artificial nests. Parental and nestling activity at nests can both increase and decrease predation rates in artificial nests compared with natural nests [24,71], and some bird species adjust their activity at the nest dependent on nearby predator activity [72].

Nest predation rates tended to be lower in visible nests compared with concealed nests. This is to be expected since nest visibility is an important determinant for predation risk in many systems [73–76], and is linked with higher predation rates of visually searching
predators [73], which are previously documented to dominate predation events in our study region [39,41]. This tendency for predation rates to depend on nest visibility probably reflect how easily predators can find the nests. However, the difference in predation rates between visibility classes remain similar between nest types, and the posterior estimated mean interaction effect was close to zero for both natural nests with artificial eggs and artificial nests. Also other studies have found relative nest predation rates between natural and artificial nests to respond similarly to nest concealment as we found in our study [20]. Relative predation rates between nest types thus seem to be comparable for different levels of this important ecological factor.

Along the forest edge-interior gradient, nest predation rates tended to be lower in the forest transition zone than the forest interior, but were similar between the forest interior and the edge. These results suggest weak or no evidence for edge effects in our study area when comparing the two extreme ends of the gradient. This result correspond with results from other studies in Europe [32,76,77], although some studies, in Europe and elsewhere, have found edge effects [66,73]. The discrepancies in results probably depend on regional or site-specific differences in main predator class and diversity [32,78,79], habitat type and type of edges investigated [30]. However, although predation rates for all nest types followed similar patterns between forest interior and forest edge, predation rates in artificial nests were markedly lower in the transition zone compared with natural nests and natural nests with artificial eggs tended to have higher predation rates than natural nests in the transition zone. Our transition zone (within 2–50 m distance from forest edge) correspond to an area where edge effects should be expected [30,73], suggesting that studies with artificial nests using distance within 50 meter from edge may show misleading patterns of predation rates compared with natural nests. Different conclusions regarding edge effects between studies could therefore, to some extent, be explained by the mixed use of natural and artificial nests in these studies, and different definitions of the edge zone.

**Conclusion**

This study indicates that both actual and relative predation rates were comparable along some of the ecological gradients explored, which correspond to the findings of other studies [4,16–18]. A clear exception to this concerned predation rates along the forest edge-interior gradient. Although natural nests with artificial eggs displayed similar absolute predation rates as natural nests, whereas artificial nests tended to have lower predation rates, both artificial nest types showed deviating predation rates from natural nests in the forest transition zone. We therefore suggest that studies using artificial nests to explore differences in predation rates along forest edge-interior gradients should be interpreted with care, especially at distances of ca 2–50 meter from edges. Comparability of predation rates between natural and artificial nests may still be valid when comparing the true forest edge (within 2 meters from the edge line) with the interior of a patch, although the comparability does not hold along the whole gradient for any artificial nest type tested here. Consistency and comparability of predation rates between natural and artificial open-cup nests often depend on an interaction between nest type and the ecological condition being tested [80], as we also show in this study. We therefore stress the importance of testing the comparability between artificial and natural nests when planning to use artificial nests to assess predation rates in new areas or in ecological contexts where this relationship has not been tested before.

**Supporting information**

**S1 Appendix. Model building and checking.** Model building procedure with model checking and evaluation from original binomial regression model with complementary log-log link and
exposure time as offset. Including r code used for analysis.

(SPDF)

**S1 File. Data file with all observations used for analysis.**

(CSV)

**S1 Fig. Dependency of exposure time effect on categorization of exposure time in nests observed at both 10 and 25 days.** Effect sizes of predictor variables explaining variation in nest predation rates based on dataset where all nests observed at both 10 and 25 days were categorized as having 25 days exposure time.

(ODCX)

**S1 Table. Distribution of nest types according to exposure time.** The table highlights how nest belonging to different nest type categories are distributed between 10 and 25 days exposure time in the dataset used for analysis in the paper, and for two alternative datasets where all nests observed after both 10 and 25 days are defined as having exposure time of either 10 days (Adjusted exposure time 10 days) or 25 days (Adjusted exposure time 25 days). See S1 Fig how the difference in categorization affects predictor estimates, and especially the effect of z-transformed exposure time.

(ODCX)

**Acknowledgments**

We thank Sylvia Aasen, Morten Einarsve, Espen Viken and Arve E. Hojem for their assistance in the field, and two anonymous referees for valuable comments.

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