

Dynamics of rodent abundance and ground-nest predation risks in forest habitats of Central Europe: no evidence for the alternative prey hypothesis

Authors: Ježková, Martina, Svobodová, Jana, and Kreisinger, Jakub

Source: Folia Zoologica, 63(4) : 269-280

Published By: Institute of Vertebrate Biology, Czech Academy of Sciences

URL: https://doi.org/10.25225/fozo.v63.i4.a6.2014

BioOne Complete (complete.BioOne.org) is a full-text database of 200 subscribed and open-access titles in the biological, ecological, and environmental sciences published by nonprofit societies, associations, museums, institutions, and presses.

Your use of this PDF, the BioOne Complete website, and all posted and associated content indicates your acceptance of BioOne's Terms of Use, available at <u>www.bioone.org/terms-of-use</u>.

Usage of BioOne Complete content is strictly limited to personal, educational, and non - commercial use. Commercial inquiries or rights and permissions requests should be directed to the individual publisher as copyright holder.

BioOne sees sustainable scholarly publishing as an inherently collaborative enterprise connecting authors, nonprofit publishers, academic institutions, research libraries, and research funders in the common goal of maximizing access to critical research.

Dynamics of rodent abundance and ground-nest predation risks in forest habitats of Central Europe: no evidence for the alternative prey hypothesis

Martina JEŽKOVÁ¹, Jana SVOBODOVÁ^{1*} and Jakub KREISINGER²

¹ Department of Ecology, Faculty of Environmental Science, Czech University of Life Sciences Prague,

Kamýcká 1176, 165 21 Prague 6, Czech Republic; e-mail: svobodovajana@fzp.czu.cz

² Department of Zoology, Faculty of Sciences, Charles University in Prague, Viničná 7, 128 44 Prague 2, Czech Republic

Received 17 March 2014; Accepted 1 December 2014

Abstract. Birds' nesting success may vary significantly between years. Ample evidence exists that this variation is caused by temporal fluctuations in rodent populations, as rodents are important components in the diets of nest predators. The alternative prey hypothesis supposes that generalist predators switch to alternative prey (bird nests) when their main prey (rodents) is lacking, thus causing increased nest predation. According to the shared predation hypothesis, by contrast, predator density is enhanced at rodent population peaks and results in simultaneous increase in main and alternative prey predation. To evaluate these hypotheses, nest predation rate dynamics were examined using artificial nests (n = 560) and rodent abundance (2240 traps) during four breeding seasons in Central European (the Czech Republic) secondary forests. Although rodent abundance increased at the population peak by almost seven times compared to the baseline and nest predation rate also showed significant inter-year variation, the data support neither the alternative prey nor shared predation hypotheses. In rich ecosystems with complex trophic levels, predators can use many resources as alternative prey. Therefore, bird nest predation risk does not increase or decrease in periods of low rodent abundance.

Key words: edge effect, generalist predators, shared predation hypothesis, prey distribution

Introduction

Nest predation is one of the main factors limiting the reproductive success of birds (Ricklefs 1969, Wegge & Storaas 1990, Martin 1995, Kauhala & Helle 2002). This source of mortality may exhibit significant interyear variation (Summers 1986), although causes of these temporal fluctuations are poorly understood. It has been proposed that between-year variation in the nest predation rates is causally associated with temporal fluctuations in abundance of small mammals (i.e. voles, *Microtus* spp. and lemmings, *Lemmus* spp.; Bêty et al. 2001, Ackerman 2002), which constitute an important dietary component of many nest predator species. For instance, small mammals (including rodents) may comprise up to 70 % of food sources for some mammalian mesopredators, such as foxes, martens and weasels (Jedrzejewska & Jedrzejewski 1998, Martinoli et al. 2001, Panzacchi et al. 2008). Nevertheless, the effects on nest predation from the interplay of fluctuations in small rodents, foraging strategies and demographic responses of predators are

rather complex. One particular possible explanation, the so-called "shared predation hypothesis" (Norrdahl & Korpimäki 2000, Reif et al. 2004), assumes that predators increase in abundance when rodent density is peaking (Bêty et al. 2002) due to their increased survival and/or reproductive output (Tannerfeldt & Angerbjörn 1998) and because of migration to areas of high prey density (Korpimäki 1994). Consequently, the increase in predator abundance leads to a higher predation rate not only on the main prey (such as small rodents, i.e. those inducing the numerical response) but also on secondary prey (Norrdahl & Korpimäki 2000, Reif et al. 2004). Although this scenario may logically apply also for bird clutches, only a few studies report positive relationships between nest predation rates and rodent density (Table 1). Increases in predator abundance due to migration are expected to have immediate effect on predation risks for secondary prey. However, predation increases on secondary prey are likely to be delayed in comparison to the rodent peak if shared predation is induced via higher survival

* Corresponding Author

H). Studies were searched using the Web of Science, Scopus, ScienceDirect and Springer Link databases with various	edation", "rodent" or "lemming cycle", "bird-lemming hypothesis". A total 43 studies were found based on these parameters	irds and population fluctuations of small mammals (rodents, lagomorphs) were excluded. In addition, studies not providing	nce were not considered.
is (APH). Studies were searched using the Web of Science	nest predation", "rodent" or "lemming cycle", "bird-lemming hy	sting birds and population fluctuations of small mammals (roc	oundance were not considered.
Table 1. Studies directed to testing the alternative prey hypothesi	combinations of several keywords: "alternative prey hypothesis", "n	and evaluated. Studies not focused on nest success of ground-nes	data (or information as to the source of data) on small mammal ab

uata (ul IIIIulIIIatiuii as tu tite sut	lice ol uala) oli sil	וומוו ווומווווומו מטעו טמו וככ אכ	ale lior collsidered.				
Source	Location	Main prey	Secondary prey	Predator	Latitude	Duration	APH*
Ackerman 2002	California	Microtus californicus	Anas platyrhynchos	Mephitis mephitis	38°14′ N, 121°97′ W	3 years	supported
Angelstam et al. 1984	Sweden	Lepus timidus	Tetrao tetrix	Vulpes vulpes	59°40′ N, 15°25′ E	5 years	supported
Bêty et al. 2001	Canada	Lemmus sibiricus, Dicrostonyx groenlandicus	Anser caerulescens atlanticus	Alopex lagopus, Larus hyperboreus, Stercorarius parasiticus, Corvus corax	72° 53' N, 78°55' W	7 years	supported
Blomqvist et al. 2002	Sweden	Lemmus sibiricus, Dicrostonyx torquatus	Calidris canutus, Calidris ferruginea	Alopex lagopus	56°12′ N, 16°24′ E	50 years	supported
Brook et al. 2005	Canada	small mammals, <i>Lepus</i> americanus	Aythya affinis, Anas platyrhynchos	Vulpes vulpes, Mustela vison, Corvus corax, Accipiter gentilis, Circus cyaneus, Bubo virginianus	62°26′ N, 114°23′ W	10 years	supported
Haselmayer & Jamieson 2001	New Zealand	Oryctolagus cuniculus	Porphyrio porphyrio	Circus approximans, Mustela furo, Felis catus, Mustela erminea	45°52′ S, 170°30′ E	5 years	supported
Iles et al. 2013	Canada	small mammals	Somateria mollissima	Alopex lagopus, Larus smithsonianus	58°43' N, 93°24' W	41 years	unsupported
Kostin & Mooij 1995	Russia	Lemmus sibiricus, Dicrostonyx torquatus	Branta ruficollis	Alopex lagopus	71°23′ N, 90°14′ E	7 years	supported
Leconte et al. 2008	Canada	Lemmus sibiricus, Dicrostonyx groenlandicus	Anser caerulescens atlanticus and artificial nests	Vulpes lagopus, Stercorarius parasiticus, Larus hyperboreus, Corvus corax	72°53′ N, 79°54′ W	11 years	supported
Marcström et al. 1988	Sweden	Clethrionomys glareolus, Sorex araneus, Microtus agrestis, M. oeconomus	Tetrao urogallus, Tetrao tetrix, Bonasa bonasia, Lagopus lagopus	Vulpes vulpes, Martes martes, Corvus cornix, C. corax, Garrulus glandarius	66° N	9 years	supported

Kinnon et al. 2013	Canada	Dicrostonyx groenlandicus, Lemmus trimucronatus	<i>Chen caerulescens</i> <i>atlantica</i> and artificial nests	Alopex lagopus, Larus hyperboreus, Stercorarius parasiticus, Corvus corax	72°53' N, 78°55' W	3 years	supported
t al. 2013	Russia	Lemmus sibiricus, Dicrostonyx torquatus	Branta bernicla bernicla	<i>Alopex lagopus</i> , skuas and gulls	74°08′ N, 86°44′ E	48 years	supported
s et al. 2007	Canada	Lemmus spp., Dicrostonyx spp.	Arenaria interpres	Alopex lagopus, Stercorarius parasiticus	64°01' N, 81°47' W	4 years	supported
& Andersen 2011	Canada	Dicrostonyx richardsoni	Branta canadensis interior	Alopex lagopus	58°34' N, 93°11' W	12 years	unsupported
iers & Underhill 1987	Russia	Lemmus sibiricus, Dicrostonyx torquatus	Branta bernicla, Calidris alba, C. ferruginea, Arenaria interpres	Vulpes vulpes, Martes martes, Mustela erminea, Meles meles	72° N 96' E	18 years	supported
chkovskiy et al. 1991	Russia	Lemmini	Anser fabalis rossicus, Branta leucopsis, Anser albifrons, Cygnus columbianus bewickii	Alopex lagopus	70°15′ N, 58°47′ E	3 years	supported
et al. 2004	Czech Republic	Apodemus flavicollis, Clethrionomys glareolus, Microtus agrestis	artificial nests	Vulpes vulpes, Martes martes, corvids	50°40′ N, 13°36′ E	2 years	supported
hill et al. 1989	Russia	Lemmus sibiricus, Dicrostonyx torquatus	Calidris canutus	Alopex lagopus	74°00′ N, 98°00′ E	19 years	supported
e & Storaas 1990	Norway	Apodemus sylvaticus, Clethrionomys glareolus, Microtus agrestis, Myopus schistiocolor, Lepus timidus	Tetrao urogallus, Tetrao tetrix	Vulpes vulpes, Martes martes, Mustela erminea, Meles meles, Garrulus glandarius	60°10′ N, 12°30′ E	8 years	supported
n & Bromley 2001	Canada	Dicrostonyx groenlandicus, Lemmus trimucronatus	Anser albifrons frontalis, Branta canadensis hutchinsii	Alopex lagopus, Larus hyperboreus, Stercorarius parasiticus	68°21' N, 108°05' W	9 years	supported

APH* supported/unsupported, i.e. negative correlation/positive relationship between rodent abundance and nest predation.

rates and reproductive output in predators (Korpimäki et al. 1991).

The alternative, mutually exclusive scenario to shared predation is the "alternative prey hypothesis" (Angelstam et al. 1984). This indicates that increased main prey abundance results in decreased predations on such secondary prey as bird clutches due to the switching of predator foraging strategies (Wilson & Bromley 2001, Bêty et al. 2002, Korpimäki et al. 2005). Many mammalian mesopredators are primarily dependent on small mammals such as voles and lemmings (Ims & Fuglei 2005). If their main prey becomes scarce, they increase the breadth of their diet by using a limited amount of secondary prey (Bêty et al. 2001) and/or exploit habitats that would be suboptimal during rodent peaks (Norrdahl & Korpimäki 2005, Lecomte et al. 2008) in accordance with the predictions of the optimal foraging theory (MacArthur & Pianka 1966). The alternative prey hypothesis has mostly been tested in boreal and arctic ecosystems that are typified by low species diversity and low food web complexity consisting of 2-3 levels (Summers & Underhill 1987, Lecomte et al. 2008). However, the evidence for the alternative prey hypothesis seems rather weak at lower latitudes, which differ from boreal/arctic ecosystems in having greater prey diversity, more complex trophic cascades and, possibly, a less clear prey profitability hierarchy. The small number of pertinent studies as well as drawbacks associated with their experimental designs do not allow making any general assessment as to the validity of the alternative prey hypothesis at low latitudes. For example, only two studies have aimed at testing the alternative prey hypothesis in non-arctic/ non-boreal parts of Europe (Saniga 2002, Šálek et al. 2004). Unfortunately, Saniga (2002) did not sufficiently assess main prey density and Šálek et al. (2004) examined changes in predation rate and rodent density during only two consecutive nesting periods and did not directly test the association between rodent densities and predation risk.

The aim of this study was to test the alternative prey and shared predation hypotheses based upon longitudinal data on temporal fluctuations in artificial ground-nest predation and rodent abundance in secondary forest habitats (forest edge and forest interior) in Central Europe (the Czech Republic). Although fluctuations in rodent numbers do not commonly occur in regular cycles within Central Europe, rodent populations nevertheless exhibit irregular yet pronounced spatial and temporal variation in this region (Tkadlec & Stenseth 2001). At the same time, previous research has shown that both nest predation rates and mammalian predator activity fluctuate considerably between consecutive breeding seasons (Svobodová et al. 2011, Svobodová et al. 2012). The obtained data allowed for testing the main predictions of both the alternative prey hypothesis and the shared predation hypothesis. Whereas the alternative prey hypothesis predicts a negative relationship between nest predation rates and small rodent abundances, the shared predation hypothesis predicts a positive association between nest predation rates and rodent counts. Furthermore, since predators may increase their reproductive success in the year subsequent to the rodent peak, a positive relationship between predation rate and rodent abundance in the subsequent year would support the shared predation hypothesis with a oneyear delay in predator response. To achieve greater robustness in the conclusion, assessment of both temporal variations in nest predation risk and small rodent abundance was undertaken in two dominant habitat types: forest edges and forest interiors.

Material and Methods

The study area was located in South Bohemia (the Czech Republic) in the Písecké Mountains region (49°11'-49°18' 14°09'-14°22', 350 m a.s.l., 60.3 km²), which is formed by a complex of production forest stands (45 % of the study area) surrounded by farmland. Coniferous and deciduous growths are typically dominated by spruce (Picea abies), pine (Pinus sylvestris) and beech (Fagus sylvatica). Mixed growths bordering the forest complex are mostly composed of such broadleaved species as oaks (Quercus spp.) and limes (Tilia spp.) along with coniferous trees from production stands. Since the forest growths are intensively cultivated, they are frequently interrupted by clearcuts, plantations and road networks. The surrounding farmland (53 % of the study area) consists mainly of hay meadows, pastures and cultivated fields that are regularly partitioned by woodlots and a network of narrow (i.e. > 10 m wide) linear strips of uncultivated vegetation (2 % of the study area; see Šálek et al. 2009 for more details), such as growths along drainage channels, windbreaks and roads.

Habitat types where the monitoring of nest predation and rodent densities took place were classified into two categories: 1) forest edges, and 2) forest interiors (places where the distance from the nearest forest edge = 100 m). Forest edges were usually fringed with oaks, limes, such trees from coniferous stands as spruces and pines, poplars (*Populus* spp.) and birches (Betula pendula). The shrub layer was usually dense, consisting mainly of shrubs and saplings of canopy trees (hazels, Corylus avellana and blackthorns, Prunus spinosa). This habitat type was also characterized by dense and highly diversified herbaceous vegetation originating mainly from nearby meadows. The boundary between forest and adjacent habitats was sharp and clearly distinguishable. The tree canopies of forest interiors were typically dominated by conifers (spruces and pines) while broadleaved tree species such as oaks, birches and beeches formed occasional admixtures. Shrub and herbal undergrowth was very sparsely developed (Svobodová et al. 2012). Previous research had determined there to be a relatively broad community of potential nest predators inhabiting the study area (Svobodová et al. 2011, 2012). Carnivores include red fox (Vulpes vulpes), martens, Eurasian badger (Meles meles), and smaller carnivores from the family of mustelids (Mustelidae, Mustela nivalis and M. erminea). Avian nest predators present include carrion crow (Corvus corone), Eurasian magpie (Pica pica) and Eurasian jay (Garrulus glandarius). In addition, wild boar (Sus scrofa) has been shown to contribute markedly to predation on artificial ground clutches in the study area.

Artificial nests were constructed by digging small depressions in the ground and then lining these with small amounts of dry plant material. Each nest was baited with two brown chicken eggs (Yahner & Mahan 1996) to ensure against predation by small predators such as rodents (Picman 1988, Haskell 1995, DeGraaf & Maier 1996). One egg in the nest was filled with a mixture of beeswax and vegetable oil for the purpose of predator identification (Pasitschniak-Arts & Messier 1995). Each wax egg was anchored in the nest hole by a string and nail in order to prevent predators from carrying it away (Summers et al. 2004, Suvorov et al. 2012). In an earlier experiment conducted in exactly the same area, no differences had been found in predation between nests baited with quail versus chicken eggs (Svobodová et al. 2012), thus suggesting that artificial nest design has a low effect on experimental outcomes. Hence, we argue that the artificial nests used in this study may be representative for a heterogeneous group of ground nesting birds (e.g. Eurasian woodcock Scolopax rusticola, yellowhammer Emberiza citrinella, tree pipit Anthus triviallis) occurring in the Central European landscape.

Inter-year variation in the rate of nest predation was studied using artificial nests during breeding periods from 2006 to 2009. In 2006, 200 artificial nests were installed. In the subsequent years, 120 nests were placed. Further, to evaluate the edge effect hypothesis, half of the artificial nests were placed at the forest edge (i.e. 5 m from the forest border into the forest interior) and half within the forest interior (100 m distant from the forest edge). The distance between two adjacent nests was > 300 m in 2006. During subsequent years, however, nest pairs (i.e. one edge nest and one interior nest) were placed in closer proximity (hereafter together referred to as a "location"). The distance between nests within individual pairs was > 100 m and the distance between pairs, similarly as in 2006, was > 300 m, in order to minimize the effect of spatial pseudoreplication (e.g. Gehring & Swihart 2003). While that variation in study design resulted in lower nest density during 2006 compared with the other seasons, it nevertheless can be assumed that this fact had minimal effect on the results of the experiment. Nests were always placed in the same locations during late April of each year. They were checked after 14 days of exposure, which is equal to the main nesting and incubation period of most bird species in the study area (Hudec & Štastný 2005). A nest was considered depredated when at least one of the two installed eggs was damaged, removed from the nest bowl or missing. Nest predators were identified according to beak or tooth marks left on the wax eggs (Nour et al. 1993) and were assigned to four categories: carnivore (red foxes, martens, small mustelids), bird, wild boar and unidentified predator (see Svobodová et al. 2012 for further details).

Rodent capture took place during the same period as did the artificial nest experiment. Common snap traps baited with wicks dipped in grease and flour were used (Heroldová et al. 2008). The traps were laid in lines of 35 traps each with the traps 5 m apart. The total length of each line was thus about 175 m. As were the artificial nests, the lines of traps were placed in the two habitat types: 1) forest interior and 2) forest edge at the same eight locations every year (64 lines with 2240 traps in total). To eliminate the potential effect of spatial pseudoreplication, the spatial distribution of capture lines followed the same rules as in the case of the artificial nest experiment (see above). Since snap traps were used whose efficiency can be affected by capture from the previous day, the traps were exposed in each location only for 24 hours.

Nest fate (survived/depredated, i.e. 0/1) after 14 days of nest exposure was included as a binary response variable into a general linear mixed effect model (GLMM, binomial distribution, logit link function) to evaluate variation underlying nest predation probability. Habitat type (i.e. forest edge or forest interior) and year were included as categorical explanatory variables and location was included as a random effect.

The number of captured rodents *vs.* the number of empty traps in a given line (consisting of 35 traps) was included as a binomial response variable in the GLMM (binomial distribution of error, logit link function). As with the model above, habitat type, year and the interaction between these variables were fitted as fixed effects and the location was included as a random effect.

The best minimal adequate model (i.e. that model with the greatest parsimony and having all variables significant) was achieved by backward elimination of non-significant effects. The significance of a particular explanatory variable was calculated by the change in deviance (assuming a χ^2 distribution) between the model containing that variable and the reduced model (i.e. using likelihood-ratio tests; Crawley 2002). All analyses were performed in R 2.12.1 software (R Development Core team 2008). GLMMs were fitted using the lmer function implemented in the *lme4* R package (Bates et al. 2012). The significance of between-year differences in rodent abundance and nest predation risk was assessed by Tukey's HSD test using the multcomp package (Hothorn et al. 2008).

To test for the existence of association between rodent densities and predation risk, subsequent analysis was restricted to nests that surrounded the lines of traps (max. distance from lines = 900 m, total n = 247). A different maximum distance cut-off was tested as well (max. distance = 400 m, n = 127), but this analysis provided identical results. The proportion of predated nests in a given location and in a given year was considered to be a response variable and the numbers of captured rodents (log transformed) to be an explanatory variable in the GLMM (binomial distribution of error, logit link function). Sampling sites and years were considered random intercepts and the relationship between predation risk and numbers of captured rodents at individual sites and in individual years to be random slopes. The same approach was



Fig. 1. Mean probability (\pm 95 % confidence intervals) of a) rodent capture per trap, and b) nest predation risk in secondary forests of South Bohemia (2006-2009) based on the prediction of a general linear mixed effect model. Different letters above bars indicate significant differences in rodent capture probability and nest predation risk between corresponding years based on Tukey's HSD test (α = 0.05).

applied to test the hypothesis that rodent densities in a given location predict predation rates during the subsequent year (i.e. the explanatory variable was number of captured rodents in the previous year).

 Table 2. Occurrence of predation on experimental nests in South Bohemia (2006-2009). A total of 537 nests were randomly distributed within two forest habitat types. Predator categories were identified according to markings on wax-filled eggs.

	Forest interior				Forest edge				Total
	2006	2007	2008	2009	2006	2007	2008	2009	
Carnivores	8	6	3	4	22	8	4	2	57
Birds	2	1	0	0	4	0	1	2	10
Wild boar	3	0	2	4	1	1	4	0	15
Unidentified	9	15	25	13	5	18	25	20	130



Fig. 2. Relationship between proportion of nests predated in a given location (eight locations were sampled each year) during 14-day exposure and a) small rodent abundance (log transformed counts), and b) small rodent abundance during the previous year. Predictions are based on the general linear mixed effect model. Dashed lines correspond to 95 % confidence intervals. Individual observations are represented by either black dots or by converging segments in the case of more than one combination of predation rates and small rodent abundance.

Results

Nest predation on artificial nests significantly differed among years ($\chi^2 = 23.23$, $\Delta df = 3$, P < 0.001) in the study area. The lowest rate was recorded in 2006 (28.6 %, n = 189, Fig. 1a). Nest predation increased over the next two years (2007: 42.6 %, n = 115; 2008: 56.6 %, n = 113), while it decreased again during the final year of observation (37.5 %, n = 120). Tukey's HSD test showed significant differences in nest predation risk only between 2006 and 2008 and between 2008 and 2009 (P < 0.001 and P = 0.019, respectively, Fig. 1a). A marginally non-significant difference in predation rates was found between 2006 and 2007 (P = 0.060). No significant or temporally consistent difference was found in the probability of nest predation between the forest edge (mean predation rates across all years = 44 %) and the forest interior (mean predation = 38 %, χ^2 = 2.22, Δ df = 1, P = 0.14). The interaction between habitat type and year was also not significant ($\chi^2 = 0.45$, Δ df = 3, P = 0.91). The most common predators on artificial nests (by percentage of identified predators) were unidentified carnivores (37 %), wild boars (18



Fig. 3. Geographic distribution of studies on the alternative prey hypothesis (included in Table 1) A) visualised on a world map, and B) plotted as a histogram against the latitude (absolute values) of corresponding sampling locations.

%), small mustelids (17 %), red foxes (16 %) and birds (12 %). However, a large number of predators were not identified at all (Table 2).

In total, 117 individual rodents were trapped in 2240 traps. The most numerous species were the yellownecked mouse (Apodemus flavicollis) and wood mouse (Apodemus sylvaticus). As in the case of nest predation, rodent numbers differed significantly among years ($\chi^2 = 91.562$, $\Delta df = 3$, P < 0.001). A pronounced peak of rodent numbers was recorded in 2007 (65.0 %, n = 76, while in the remaining years rodent numbers were on average about one-sixth those in the peak year (P < 0.001 in all cases, Fig. 1b). Rodent abundance did not differ significantly between forest edge and forest interior ($\chi^2 = 2.798$, $\Delta df = 1$, P = 0.094), though rodents nevertheless tended to be more abundant in general at forest edges compared to forest interiors. The interaction between habitat type and year was also not significant ($\chi^2 = 0.757$, $\Delta df =$ 3, P = 0.860).

No association was found between the proportion of predated nests and rodent counts at the locations where rodent abundance was assessed (GLMM: slope \pm SE = 0.276 \pm 0.286, Δ df = 1, χ^2 = 0.814, P = 0.367, Fig. 2a). Similarly, rodent density in a given location did not predict nest predation in the following year (GLMM: slope \pm SE = 0.300 \pm 0.563, Δ df = 1, χ^2 = 0.098, P = 0.755, Fig. 2b).

Discussion

The data show that both rodent abundance and nest predation risk exhibit considerable temporal variation. The effect of habitat on predation risk and rodent abundance, however, is of relatively low importance. Many previous studies have implied that fluctuations in rodent abundance may cause marked effects on ecosystem functioning (e.g. Ims & Fuglei 2005), including nest predation risk (Ackerman 2002). In theory, both increases (Korpimäki & Norrdahl 1989) and decreases (Bêty et al. 2001) in nest predation rates might be expected when small rodent populations increase.

The alternative prey hypothesis, which predicts decreased nest predation risk during population peaks of small rodents, has received considerable support in many empirical studies. An extensive literature search (Table 1), turned up 20 studies which directly evaluated the alternative prey hypothesis and where small mammals (rodents, lagomorphs) and ground nests represented the main and the alternative prey, respectively. Most of these studies found support for the alternative prey hypothesis (n = 18). Nevertheless,

this nearly universal support for the alternative prey hypothesis might be artificially inflated by several factors. First, this set of studies exhibits a nonrandom geographical distribution (Fig. 3). While most data from testing the alternative prey hypothesis come from the Arctic (13 studies with latitude > 60°) and the north temperate zone (five studies with latitudes between 60° and 40°), only one study has been performed in the tropics or subtropics. In addition, only one study comes from the Southern Hemisphere. The mean latitude of the studies on the alternative prey hypothesis was 65°, which suggests a clear bias toward higher latitudes. The existing literature on the alternative prey hypothesis might be affected by a severe publication bias as well. In other words, mentioning the alternative prey hypothesis in the context of a certain type of result might be a more or less opportunistic ex post decision to make an article more appealing to potential readers, even when those results were not a priory collected for the purpose of testing the alternative prey hypothesis. Moreover, similar data that do not exhibit a pattern corresponding to the alternative prey hypothesis might possibly be less likely to be published. Nevertheless, a more extensive and rigorous review of the literature on temporal fluctuations in nest success would be necessary to account for these concerns.

The present study was situated at a latitude lower than those of 90 % of those studies which have tested the alternative prey hypothesis (Table 2, Fig. 3). Based on field data collected in the study area, it was possible to reject the alternative prey hypothesis. Contrary to the prediction of the alternative prey hypothesis, nest predation risk did not decrease during the rodent population peak. The lack of evidence for the alternative prey hypothesis and the fact that the results contradict those of most previous studies may be caused by several factors. The main predators of the artificial nests, such as red fox and martens, are habitat generalists (Svobodová et al. 2012) and so they may also respond to rodent abundance from adjacent farmland. We do not believe, however, that unobserved variance in rodent density on farmland would have a substantial effect on artificial nest predation in the study area. Data from the previous experiment indicate that rodent abundances in forest and farmland are strongly correlated and, in addition, rodent abundances are substantially lower in surrounding meadows than in forest habitats (Svobodová et al. 2011, Ježková M. unpublished data). In particular, the diversity of nest predator and rodent species is likely to be greater in the study area compared to higher latitudes, where

the evidence for the alternative prey hypothesis is rather strong. At the same time, lower latitudes may also offer a greater diversity of alternative prey that is of comparable or even higher profitability compared to bird nests. Many carnivores preying on artificial nests in the study area are obviously not dependent solely on bird nests as secondary prey, because during nesting periods a significant part of their diet is composed of birds, reptiles, amphibious insects (Jędrzejewska & Jędrzejewski 1998, Lanszki et al. 2007) and the young of larger mammals (e.g. the European hare, Lepus europaeus and roe deer, Capreolus capreolus; Lindström 1994, Kjellander & Nordström 2003, Panzacchi et al. 2008). In addition, a large number of the artificial nests were preyed upon by wild boar and corvids, whose diet is not dependent on rodents (Schley & Roper 2003, Hoyo et al. 2009). Consequently, for these predators the rodent population dynamic is unlikely to modulate profitability of bird clutches. At the same time, it is also possible that the results were affected by the fact that the population increase of small rodents in the area (ca 6.5 times the population baseline) was less pronounced compared to that of other studies focused on the alternative prey hypothesis. For example, main prey density was more than 30 times higher than the minimum during its population boom in some cases (Wegge & Storaas 1990, Ackermann 2002). Consequently, the change in main prey abundance observed in the study, although seemingly dramatic, might have had a negligible effect on predators' foraging tactics. Finally, irregular fluctuation of rodent density, which is typical for lower latitudes, might have contributed to the weak response by predators and consequently to the low effect on nest predation rates. In particular, it can be hypothesized that regular rodent population cycling at high latitudes may induce greater selective pressure on predators' foraging flexibility. On the other hand, the fact that the alternative prey hypothesis has been supported by several studies performed in areas where regular population cycling does not occur (Ackerman 2002, Šálek et al. 2004) suggests that the importance of this effect is rather limited.

Some previous studies used relatively short-term monitoring of nest predation and rodent density fluctuation to test the alternative prey hypothesis. For example, the contributions of Šálek et al. (2004) and Ackerman (2002) are based on data just from two and three years, respectively. The data from the present study demonstrate that relatively long-term data covering the periods both before and after the rodent peak are necessary for testing the alternative prey hypothesis. In particular, if the dataset had covered just the two breeding seasons in 2007 and 2008, it might have been erroneously found to support the alternative prey hypothesis, because, consistently with the predictions of this hypothesis, rodent abundance decreased and nest predation increased during these two years.

The opposite pattern of that predicted by the alternative prey hypothesis is expected under the "shared predation hypothesis". In particular, betweenyear differences in nest predation risk may correlate positively with small mammal density, due to an apparent competition-like effect (Holt 1977) induced by the numerical and dietary (functional) response of predators to the main prey (reviewed by Chalfoun et al. 2002). The shared predation hypothesis has been supported, however, mainly in the case of such predator specialists as weasels (Mustela nivalis) and stoats (Mustela erminea), for whose diets rodents comprise an important part (Jedrzejewski et al. 1995, Dupuy et al. 2009). It is nevertheless noteworthy that there also are studies providing support for the shared predation hypothesis in the case of generalist predators (Zalewski et al. 1995, O'Donoghue et al. 1997, Jedrzejewska & Jedrzejewski 1998). In any case, the present data do not support the "shared predator" scenario. We hypothesize that the rich Central European ecosystem probably hosts high densities of alternative prey which are of comparable or even higher profitability compared to bird nests. In addition, small rodents do not represent the main food sources in the area of this study for such important nest predators as wild boar and corvids (Schley & Roper 2003, Hoyo et al. 2009).

Many studies focused on nest predation have found nesting success to be decreased along the edges of various habitat types compared to within habitat interiors (e.g. Andrén 1992, Bayne & Hobson 1997, Malt & Lank 2007). Some predator species specifically exploit edge habitats because their main prey (rodents in the present study) is concentrated along these landscape structures (Ries & Sisk 2004, more in Koubová et al. 2011). The present study does not support the edge effect hypothesis, however, as the probability of nest predation and small rodent numbers did not differ substantially between forest interiors and forest edges. It is noteworthy in this regard that most studies supporting the edge effect on nesting success have been conducted in North America and Northwest Europe (reviewed in Batáry & Báldi 2004), which regions have different landscape compositions and predator communities compared to Central European conditions. It can therefore be hypothesized that, corresponding to the absence of the alternative prey hypothesis in the study area, the existence of the edge effect on nest predation is less obvious in a region with rich predator communities such as Central Europe's intensively managed landscape (Svobodová et al. 2012).

It may be argued that these results are ambiguous, as artificial nests were used for monitoring nest predation risk. Several concerns associated with the use of artificial nests have previously been raised (Storaas 1988, Willebrand & Marcström 1988), as they do not wholly mimic several features of real nests, such as parental behaviour (Weidinger 2002, Kreisinger & Albrecht 2008), scent (Whelan et al. 1994), microhabitat features (Yahner & Piergallini 1998), nest density (Major & Kendal 1996) and the construction of real nests (Báldi 2000). All of these characteristics are, in fact, crucial determinants of nesting success. Although these concerns have some merit, these key features of real nests may vary between breeding seasons. Using artificial nest experiments thus enables researchers to hold these invariant. The use of artificial nests, therefore, provides a considerable methodical advantage for longitudinal studies, as it allows for decreasing potential bias associated with this source of variation. In addition, it is worthy of note that artificial nests had been used in three previous studies on the alternative hypothesis and that all of them provided support for this hypothesis (Šálek et al. 2004, Lecomte et al. 2008, McKinnon et al. 2013). Therefore, despite wellknown methodological drawbacks, artificial nests are still, to a certain extent, useful tools for analysing temporal variation of complexities in trophic cascades, including predator *vs.* main *vs.* alternative prey relationships.

In conclusion, high inter-year variation was found in nest predation and rodent density, but the data suggest no straightforward link between these findings. The data allow for rejecting both the alternative prey hypothesis and the shared predation hypothesis. It can be supposed that in rich ecosystems having complex trophic levels, predators can use many sources of alternative prey, and therefore predation risk for bird nests is not substantially related to small rodent abundance. Hence, the possible importance of rodent abundances in relation to nest predation risk must be considered cautiously and with a view to specific local conditions.

Acknowledgements

We thank David Hořák, Ondřej Sedláček, Miroslav Šálek and two anonymous reviewers for valuable comments on earlier drafts of the manuscript. We are grateful to J. Ceplová, O. Častoral, E. Dýnková, P. Kukač, L. Mrštný and L. Soustružník for their help with fieldwork. This study was supported by the internal grant agencies of the Czech University of Life Sciences Prague (CIGA 20094205 and IGA 20134222).

Literature

Ackerman J.T. 2002: Of mice and mallards: positive indirect effects of coexisting prey on waterfowl nest access. Oikos 99: 469-480.

- Andrén H. 1992: Corvid density and nest predation in relation to forest fragmentation: a landscape perspective. *Ecology 73:* 794–804.
- Angelstam P., Lindström E. & Widén P. 1984: Role of predation in short-term population fluctuations of some birds and mammals in Fennoscandia. Oecologia 62: 199–208.
- Báldi A. 2000: Different depredation rates between daylight and twilight placed artificial nests. Folia Zool. 49: 157-159.

Batáry P. & Báldi A. 2004: Evidence of an edge effect on avian nest success. Conserv. Biol. 18: 389-400.

- Bates D., Maechler M. & Bolker B. 2012: lme4: linear mixed-effects models using S4 classes. R package version 0.999999-0. http:// CRAN.R-project.org/package=lme4
- Bayne E.M. & Hobson K.A. 1997: Comparing the effects of landscape fragmentation by forestry and agriculture on predation of artificial nests. *Conserv. Biol.* 11: 1418–1429.
- Bêty J., Gauthier G., Giroux J.F. & Korpimäki E. 2001: Are goose nesting success and lemming cycles linked? Interplay between nest density and predators. *Oikos 93: 388–400*.
- Bêty J., Gauthier G., Korpimäki E. & Giroux J.F. 2002: Shared predators and indirect trophic interactions: lemming cycles and arctic nesting geese. J. Anim. Ecol. 71: 88–89.
- Blomqvist S., Holmgren N., Åkesson S., Hedenström A. & Pettersson J. 2002: Indirect effects of lemming cycles on sandpiper dynamics: 50 years of counts from southern Sweden. *Oecologia 133: 146–158*.
- Brook R.W., Duncan D.C., Hines J.E., Carrière S. &. Clark R.G. 2005: Effects of small mammal cycles on productivity of boreal ducks. *Wildlife Biol.* 11: 3–11.
- Chalfoun A.D., Thompson F.R. & Ratnaswamy M.J. 2002: Nest predators and fragmentation: a review and meta-analysis. *Conserv. Biol.* 16: 306–318.
- Crawley M.J. 2002: Statistical computing. Wiley, Chichester.

DeGraaf R.M. & Maier T.J. 1996: Effect of egg size on predation by white-footed mice. Wilson Bull. 108: 535-539.

Dupuy G., Giraudoux P. & Delattre P. 2009: Numerical and dietary responses of a predator community in a temperate zone of Europe. *Ecography 32: 277–290.*

Gehring T.M. & Swihart R.K. 2003: Body size, niche breadth, and ecologically scaled responses to habitat fragmentation: mammalian predators in an agricultural landscape. *Biol. Conserv. 109: 283–295.*

Haselmayer J. & Jamieson I.G. 2001: Increased predation on pukeko eggs after the application of rabbit control measures. *New Zeal. J. Ecol.* 25: 89–93.

Haskell D. 1995: A revelation of the effects of forest fragmentation on rates of bird-nest predation. Conserv. Biol. 9: 1316–1318.

Heroldová M., Tkadlec E., Bryja J. & Zejda J. 2008: Wheat or barley? Feeding preferences affect distribution of three rodent species in agricultural landscape. *Appl. Anim. Behav. Sci. 110: 354–362.*

Holt R.D. 1977: Predation, apparent competition, and the structure of prey communities. Theor. Popul. Biol. 12: 197-229.

Hothorn T., Bretz F. & Westfall P. 2008: Simultaneous inference in general parametric models. Biometrical J. 50 (3): 346-363.

Hoyo J., Elliot A. & Christie D.A. 2009: Handbook of birds of the world, Vol. 14. Bush-shrikes to old world sparrows. *Lynx Edicions, Barcelona, Spain.*

Hudec K. & Šťastný K. (eds.) 2005: Fauna of the Czech Republic. Birds 2/2. Academia, Praha. (in Czech)

Iles D.T., Rockwell R.F., Matulonis P., Robertson G.J., Abraham K.F., Davies Ch. & Koons N. 2013: Predators, alternative prey, and climate influence annual breeding success of a long-lived sea duck. J. Anim. Ecol. 82: 683–693.

Ims R.A. & Fuglei E. 2005: Trophic interaction cycles in tundra ecosystems and the impact of climate change. BioScience 55: 311-322.

- Jędrzejewska B. & Jędrzejewski W. 1998: Predation in vertebrate communities. The Białowieża Primeval Forest as a case study. Springer Verlag, Berlin.
- Jędrzejewski W., Jędrzejewska B. & Szymura L. 1995: Weasel population response, home range, and predation on rodents in a deciduous forests in Poland. *Ecology 76: 179–195*.

Kauhala K. & Helle P. 2002: The impact of predator abundance on grouse populations in Finland – study based on wildlife monitoring counts. *Ornis Fennica* 79: 14–25.

- Kjellander P. & Nordström J. 2003: Cyclic voles, prey switching in red fox, and roe deer dynamics a test of the alternative prey hypothesis. *Oikos 101: 338–344.*
- Korpimäki E. 1994: Rapid or delayed tracking of multi-annual vole cycles by avian predators? J. Anim. Ecol. 63: 619-628.
- Korpimäki E. & Norrdahl K. 1989: Predation of Tengmalm's owls: numerical responses, functional response and dampening impact on population fluctuations of voles. *Oikos 54: 154–164.*
- Korpimäki E., Norrdahl K. & Rinta-Jaskari T. 1991: Responses of stoats and least weasels to fluctuating food abundances: is the low phase of the vole cycle due to mustelid predation? *Oecologia 88: 552–561*.
- Korpimäki E., Norrdahl K., Huitu O. & Klemola T. 2005: Predator-induced synchrony in population oscillations of coexisting small mammal species. *Proc. R. Soc. Lond. B* 272: 193–202.
- Kostin I. & Mooij J.H. 1995: Influence of weather conditions and other factors on the reproductive cycle of red-breasted geese *Branta ruficollis* on the Taimyr Peninsula. *Wildfowl 46: 45–54.*
- Koubová M., Suvorov P., Svobodová J., Albrecht T. & Kreisinger J. 2011: Fragmentation and loss of natural habitats. Chapter 7. In: Daniels J.A. (ed.), Advances in environmental research 14. *Nova Science Publishers Inc.: 183–207.*
- Kreisinger J. & Albrecht T. 2008: Nest protection in mallards *Anas platyrhynchos*: untangling the role of crypsis and parental behaviour. *Funct. Ecol. 22: 872–879.*
- Lanszki J., Zalewski A. & Horváth G. 2007: Comparison of red fox *Vulpes vulpes* and pine marten *Martes martes* food habits in a deciduous forest in Hungary. *Wildlife Biol.* 13: 258–271.
- Lecomte N., Careau V., Gauthier G. & Giroux J.F. 2008: Predator behaviour and predation risk in the heterogeneous Arctic environment. J. Anim. Ecol. 77: 439–447.

Lindström E.R. 1994: Large prey for small cubs - on crucial resources of a boreal red fox population. Ecography 17: 17-22.

MacArthur R.H. & Pianka E.R. 1966: On the optimal use of a patchy environment. Am. Nat. 100: 603–609.

Major R.E. & Kendal C.E. 1996: The contribution of artificial nests experiments to understanding avian reproductive success: a review of methods and conclusions. *Ibis 138: 298–307.*

Malt J. & Lank D. 2007: Temporal dynamics of edge effects on nest predation risk for the marbled murrelet. *Biol. Conserv. 140: 160–173.*

- Marcström V., Kenward R.E. & Engren E. 1988: The impact of predation on boreal tetraonids during vole cycles: an experimental study. *J. Anim. Ecol.* 57: 859–872.
- Martin T.E. 1995: Avian life-history evolution in relation to nest sites, nest predation, and food. Ecol. Monogr. 65: 101–127.

Martinoli A., Preatoni D.G., Chiarenzi B., Wauters L.A. & Tosi G. 2001: Diet of stoats (*Mustela erminea*) in an Alpine habitat: the importance of fruit consumption in summer. *Acta Oecol.* 22: 45–53.

- McKinnon L., Berteaux D., Gauthier G. & Bêty J. 2013: Predator-mediated interactions between preferred, alternative and incidental prey in the arctic tundra. *Oikos 122: 1042–1048*.
- Nolet B.A., Bauer S., Feige N., Kokorev Y.I., Popov I.Y. & Ebbinge B.S. 2013: Faltering lemming cycles reduce productivity and population size of a migratory Arctic goose species. J. Anim. Ecol. 82: 804–813.
- Norrdahl K. & Korpimäki E. 2000: Do predators limit the abundance of alternative prey? Experiments with vole-eating avian and mammalian predators. *Oikos 91: 528–540*.

Norrdahl K. & Korpimäki E. 2005: Survival through bottlenecks of vole cycles: refuge or chance events? Evol. Ecol. 19: 339-361.

Nour N., Matthysen E. & Dhondt A.A. 1993: Artificial nest predation and habitat fragmentation: different trends in birds and mammal predators. *Ecography 16: 111–116*.

O'Donoghue M., Boutin S., Krebs C.J. & Hofer E.J. 1997: Numerical responses of coyotes and lynx to the snowshoe hare cycle. *Oikos* 80: 150–162.

- Panzacchi M., Linnell J.D.C., Serrao G., Eie S., Odden M., Odden J. & Andersen R. 2008: Evaluation of the importance of roe deer fawns in the spring-summer diet of red foxes in south-eastern Norway. *Ecol. Res.* 23: 889–896.
- Pasitschniak-Arts M. & Messier F. 1995: Risk of predation on waterfowl nests in the Canadian prairies: effects of habitat edges and agricultural practices. *Oikos 73: 347–355*.
- Perkins D.E., Sith P.A. & Gilchrist H.G. 2007: The breeding ecology of ruddy turnstones (*Arenaria interpres*) in the eastern Canadian Arctic. *Polar Rec.* 43: 135–142.
- Picman J. 1988: Experimental study of predation on eggs of groundnesting birds: effects of habitat and nest distribution. Condor 90: 124–131.
- R Development Core Team 2008: R: a language and environment for statistical computing. Vienna, Austria. Available at http://www. Rproject.org
- Reif V., Jungell S., Korpimäki E., Tornberg R. & Mykrä S. 2004: Numerical response of common buzzards and predation rate of main and alternative prey under fluctuating food conditions. *Ann. Zool. Fenn.* 41: 599–607.
- Reiter M.E. & Andersen D.E. 2011: Arctic foxes, lemmings, and Canada goose nest survival at cape Churchill, Manitoba. *Wilson J. Ornithol.* 123: 266–276.
- Ricklefs R.E. 1969: An analysis of nesting mortality in birds. Smithson. Contrib. Zool. 9: 1-48.
- Ries L. & Sisk T.D. 2004: A predictive model of edge effects. Ecology 85: 2917-2926.
- Saniga M. 2002: Nest loss and chick mortality in capercaillie (*Tetrao urogallus*) and hazel grouse (*Bonasa bonasia*) in West Carpathians. *Folia Zool.* 51: 205–214.
- Schley L. & Roper T.J. 2003: Diet of wild boar *Sus scrofa* in Western Europe, with particular reference to consumption of agricultural crops. *Mammal Rev.* 33: 43–56.
- Storaas T. 1988: A comparison of losses in artificial and naturally occurring capercaillie nests. J. Wildlife Manage. 52: 123-126.
- Summers R.W. 1986: Breeding production of dark-bellied brent geese *Branta bernicla bernicla* in relation to lemming cycles. *Bird Study 33: 105–108.*
- Summers R.W. & Underhill L.G. 1987: Factors related to breeding production of brent geese *Branta b. bernicla* and waders (*Charadrii*) on the Taimyr Peninsula. *Bird Study 34: 161–171*.
- Summers R.W., Green R.E., Proctor R., Dugan D., Lambie D. & Moncrieff R. 2004: An experimental study of the effects of predation on the breeding productivity of capercaillie and black grouse. J. Appl. Ecol. 41: 513–525.
- Suvorov P., Svobodová J., Koubová M. & Dohnalová L. 2012: Effect of quail and chicken eggs on predation by European black-billed magpie *Pica pica. Acta Ornithol.* 47: 55–61.
- Svobodová J., Koubová M., Mrštný L., Albrecht T. & Kreisinger J. 2012: Temporal variation in nest predation risk along habitat edges between grassland and secondary forest in Central Europe. *Eur. J. Wildlife Res.* 58: 315–323.
- Svobodová J., Kreisinger J., Šálek M., Koubová M. & Albrecht T. 2011: Testing mechanistic explanations for mammalian predator responses to habitat edges. *Eur. J. Wildlife Res.* 57: 467–474.
- Syroechkovskiy Y.V., Litvin K.Y. & Ebbinge B.S. 1991: Breeding success of geese and swans on Vaygach Island (USSR) during 1986-1988; interplay of weather and Arctic fox predation. *Ardea 79: 373–382*.
- Šálek M., Kreisinger J., Sedláček F. & Albrecht T. 2009: Corridor vs. hayfield matrix use by mammalian predators in an agricultural landscape. *Agric. Ecosyst. Environ.* 134: 8–13.
- Šálek M., Svobodová J., Bejček V. & Albrecht T. 2004: Predation on artificial nests in relation to the numbers of small mammals in the Krušné hory Mts, the Czech Republic. *Folia Zool. 53: 312–318*.
- Tannerfeldt M. & Angerbjörn A. 1998: Fluctuating resources and the evolution of litter size in the arctic fox. Oikos 83: 545-559.

Tkadlec E. & Stenseth N.C. 2001: A new geographical gradient in vole population dynamics. Proc. R. Soc. Lond. B 268: 1547-1552.

- Underhill L.G., Waltner M. & Summers R.W. 1989: Three-year cycles in breeding productivity of knots *Calidris canutus* wintering in southern Africa suggest Taimyr Peninsula provenance. *Bird Study 36: 83–87*.
- Weidinger K. 2002: Interactive effects of concealment, parental behaviour and predators on the survival of open passerine nests. J. Anim. Ecol. 71: 424–437.
- Wegge P. & Storaas T. 1990: Nest loss in capercaillie and black grouse in relation to the small rodent cycle in southeast Norway. *Oecologia 82: 527–530.*
- Whelan C.J., Dilger M.L., Robson D., Hallyn N. & Dilger S. 1994: Effects of olfactory cues on artificial-nest experiments. Auk 11: 945–952.
- Willebrand T. & Marcström V. 1988: On the danger of using dummy nests to study predation. Auk 105: 378-379.
- Wilson D.J. & Bromley R.G. 2001: Functional and numerical responses of predators to cyclic lemming abundance: effects on loss of goose nests. Can. J. Zool. 79: 525–532.
- Yahner R.H. & Mahan C.G. 1996: Effects of egg type on depredation of artificial ground nests. Wilson Bull. 108: 129-136.
- Yahner R.H. & Piergallini N.H. 1998: Effects of microsite selection on predation of artificial ground nests. Wilson Bull. 110: 439-442.
- Zalewski A., Jędrzejewski W. & Jędrzejewska B. 1995: Pine marten home ranges, numbers and predation on vertebrates in a deciduous forest (Białowieża National Park, Poland). Ann. Zool. Fenn. 32: 131–144.