

Annual variation in prey composition of domestic cats in rural and urban environment

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Abstract The threat that domestic cats pose to wildlife has gained increased recognition by researchers and conservationists, and in this study, we investigated the seasonal variability and the effects of environment type (rural vs. urban) on the prey composition of free-ranging house cats in Poland. We analysed the variability in 307 monthly prey samples of different prey items killed by cats and brought to their owners (i.e., prey brought home by cats living in one home in one month) between 2002 and 2007 at 26 rural and urban sites. The variability in prey composition over time was analysed using additive models and canonical correspondence analysis. In total, we recorded 1348 prey items. Rodents were the most common prey in both environments, but shrews and reptiles were killed by cats more often in the rural environment while birds (mainly sparrows and pigeons) were more common in the urban environment. Additionally, prey composition changed seasonally. The pooled number of vertebrates killed by cats was largest in September and lowest in January, and rodents were killed most often in September, shrews and birds in June, and reptiles in April. The seasonal variation in the prey composition of cats was relatively high in the rural environment and more stable in the urban environment. Prey composition seemed to follow temporal and spatial variations in

prey availability, thus confirming a facultative feeding strategy in free-ranging house cats.

Key words *Felis catus* · Central Poland · Prey brought home · Season · Vertebrates · Rural to urban

Introduction

As medium-sized carnivores, domestic cats, *Felis catus*, are usually strictly associated with human settlements and have been introduced by humans all over the world. They are effective predators that mainly hunt small or medium-sized vertebrates (Fitzgerald and Turner 2000), and while the density of feral cats is directly correlated with prey abundance (Genovesi et al. 1995; Edwards et al. 2001), the population density of free-ranging house cats is more reflective of human density (due to the provision of supplementary food) than that of their prey (Sims et al. 2008). Consequently, cat populations reach high densities near human settlements, so they impose relatively strong hunting pressure on prey populations (Baker et al. 2005; Sims et al. 2008; Thomas et al. 2012). In urban areas in particular, cats are one of the most common avian predators (Sims et al. 2008), but their total impact on wildlife is difficult to estimate and varies greatly between studies (Fitzgerald and Turner 2000) and with study methodology (Krauze-Gryz et al. 2012a). Recently, however, cats have been estimated to kill between 100 and 350 million birds per year in Canada (Blancher 2013) and 2.4 billion birds and 12.3 billion mammals annually in the United States (Loss et al. 2013). In terms of overall proportions, they have been estimated to kill 2–7% of all birds in southern Canada (Blancher 2013) and to be responsible for at least 30% of sparrow deaths in an English village (Churcher and Lawton 1987). Introduced cats are

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responsible for the decline of native mammals in Australia (Risbey et al. 2000; Wheeler and Priddel 2009; Moseby et al. 2009; Frank et al. 2014), and several island studies have documented strong negative effects by cats on local endemic fauna (e.g., Fitzgerald and Veitch 1985; Medina and Nogales 2009; Bonnaud et al. 2012). These examples show, without a doubt, that both feral and free-ranging house cats should be considered a serious threat to biodiversity.

The evidence of the strong impact of cats on wildlife has focused the attention of conservation biologists towards possible methods of reducing that pressure. Consequently, many conservation initiatives have been aimed at the eradication of feral cats on islands (Nogales et al. 2004), but the threat that free-ranging house cats pose to wildlife is also serious and has gained recognition by both researchers and conservationists (e.g., Kays and DeWan 2004; Baker et al. 2005; Sims et al. 2008; Morgan et al. 2009; Bonnington et al. 2013). While there is a set of mitigation measures that can be considered (reviewed in Calver et al. 2011), the application of any of them requires cooperation with cat owners, who often fail to understand the detrimental effect their pets can have on wildlife (McDonald et al. 2015). Thus, studies showing the diversity of animal species that are killed by cats as well the magnitude of their hunting pressure are important; they both inform public opinion about the problem and can help raise financial support for mitigation. Most of the studies focused on the diet of free-ranging house cats have been conducted in the USA (Kays and DeWan 2004; Lepczyk et al. 2004), Australia and Oceania (Barrat 1997; Gillies and Clout 2003; Morgan et al. 2009; van Heezik et al. 2010). In Europe, where there is a relationship between high-density human populations and a high abundance of domestic cats, previous investigations were mainly located in the United Kingdom (i.e., Churcher and Lawton 1987; Woods et al. 2003; Baker et al. 2005; Thomas et al. 2012) but also in Sweden (Liberg 1984), Switzerland (Weber and Daily 1998; Tschanz et al. 2010), Poland (Krauze-Gryz et al. 2012a) and Finland (Kauhala et al. 2015). However, this topic has never been investigated in many central and eastern European countries.

Previous studies have focused on the diets of free-ranging house cats to better understand their spatial and temporal variations (e.g., Barrat 1997; Baker et al. 2005; van Heezik et al. 2010; Kauhala et al. 2015), but several knowledge gaps still exist. The dietary composition of domestic house cats may differ from the empirical patterns observed in wild carnivores as most of house cats are supplementary fed by their owners, so they can continue to hunt a certain prey category even if the hunt is no longer profitable in terms of the costs-to-energy gain ratio. Furthermore, because of their reliance on artificial food sources, cats can substantially surpass the carrying capacity of the environment (reviewed in Kays and DeWan 2004). Consequently, even though food provisioning may reduce individual predation rates, the total impact of cat

predation can be severe (reviewed in Baker et al. 2005). Furthermore, free-ranging house cats hunt in both natural habitats as well as close to human settlements (Kays and DeWan 2004; Goszczyński et al. 2008; Wierzbowska et al. 2012), and the latter group are typically avoided by many wild carnivores because of the presence of humans (Krauze-Gryz et al. 2012b). Finally, cats exhibit atypical hunting behaviour; i.e., they often kill and leave prey uneaten (Fitzgerald and Turner 2000; Krauze-Gryz et al. 2012a; Loyd et al. 2013). Because of these factors, the general picture of the variation in the prey composition of free-ranging cats is incomplete, especially in central and eastern Europe, leaving a high degree of uncertainty in terms of the total pressure of cats on wildlife.

In this study, we investigated the seasonal variability and the effects of two environments on the prey composition of free-ranging house cats in Poland. In previous studies, the diet composition and predation pressure of cats in natural and rural habitats have been contrasted with records from more transformed and built-up areas (e.g., Gillies and Clout 2003; Lepczyk et al. 2004; Kauhala et al. 2015), but few, if any, studies have addressed the seasonal variation in prey composition in this context (but see Kauhala et al. 2015). We attempted to do so and analysed the year-round variability in the different prey items killed by domestic cats at 26 sites in rural and urban environments, which provided us with the opportunity to separate the effects of time and the environment and to test the following three predictions. First, given the well-described shift in the diets of other generalist predators towards avian prey in urbanized areas, e.g., the tawny owl *Strix aluco* (Goszczyński et al. 1993; Zalewski 1994); the red fox *Vulpes vulpes*; or the stone marten *Martes martes* (reviewed in Bateman and Fleming 2012), we predicted that cats living in urban environments hunt birds with a higher frequency than rural cats. Second, we hypothesized that the prey composition of cats, which are generalist predators (Fitzgerald and Turner 2000), reflects changes in prey availability throughout the year, despite the supplementary feeding that these cats receive. Finally, we predicted that the annual variability in prey composition may differ between rural and urban environments as prey populations in these areas may follow different temporal dynamics.

Methods

Study area

The study was conducted in central Poland, a region that is affected by the mild oceanic climate of Western Europe and the harsh and dry continental climate of Eastern Europe and Asia. The duration of the growing season is c. 210 days; the total precipitation is 600 mm per year; and the mean ambient temperature ranges from -4°C in January to $+18^{\circ}\text{C}$ in July.

We conducted the research in two environments: rural versus urban. The rural areas represented a typical field-forest mosaic with a prevalence of arable lands, in which woods of a few hundred hectares are surrounded by a fine mosaic of different crops, pastures, fallow land and stands of trees. Arable lands constitute 60–90% of the area, and the forest cover is much lower (6–30%). Villages (primarily consisting of a row of settlements along a road) and single farmsteads are evenly distributed at distances of no more than a few hundred metres from each other. Farms are typically small (i.e., most are 1–10 ha in size; Central Statistical Office-GUS data), and the population density is below 100 inhabitants/km². The urban environment was mainly represented by small towns in the vicinity and outskirts of Warsaw (i.e., its peripheral quarters), which is characterized by low buildings and houses with yards that are adjacent to green areas, such as parks or forests. However, the more central part of the city is densely built-up with tenements or blocks of flats with little green space that is mostly in the form of lawns located between buildings and sparse trees or shrubs. The population density is between 1000 and 4000 inhabitants/km², but in the central districts of the city, it can reach up to 8600 inhabitants/km². On the whole, the built-up areas cover approx. 50%, arable lands and fallows 30%, and forests 16% of the whole area of Warsaw.

Collection of cat preys

We collected the prey of cats belonging to 16 owners in rural and 10 owners in urban environments. Cat owners were recruited through door-to-door surveys in selected villages and through Internet forums, and they were asked to collect and record all of the prey that their cats brought home (Churcher and Lawton 1987; Gillies and Clout 2003; Baker et al. 2005). From October 2002 to December 2007, we collected data from the owners who cooperated with our study regularly for particular periods (between 1 and 43 months with an average of 12). In total, cats from 26 sites were monitored, and there were between 1 to 6 cats (1.8 on average) in a single site. All of the cats were active hunters and could leave their own settlements.

The cat owners were trained to recognize the most common prey species, and they used a data sheet to record all of the prey brought home and either froze the remains for storage or took photos for further assessment. The owners were also regularly contacted and assisted, and the data were collected at least once a month. Prey items were classified to the lowest possible taxonomic level (usually species or genus), but in some cases, i.e., if prey was not stored and we were not confident about the owner's ability to correctly identify the species, they were identified to order or class (e.g., 'rodentia unidentified', 'birds unidentified', etc.). In total, 17 detailed prey categories were distinguished (Table 1), but for the purpose of additive mixed modelling (see below), we grouped all

Table 1 Prey brought home by free-ranging domestic cats in rural and urban environments

Prey category	Rural	Urban
Muridae	363	115
Cricetidae	181	24
Rodentia unident.	134	19
Soricomorpha	111	5
Lagomorpha	2	0
Carnivora	3	0
Mammal unident.	22	1
Passeriformes	73	63
Piciformes	1	0
Columbiformes	4	25
Galliformes	4	0
Bird unident.	23	16
<i>Lacerta agilis</i>	16	1
<i>Lacerta vivipara</i>	24	0
Lacertilia unident.	81	9
Amphibia	10	2
Cypriniformes	11	5
Total	1063	285

of the prey items into five general categories: (1) rodents, (2) Soricomorpha (moles and shrews pooled), (3) birds, (4) reptiles and (5) all vertebrates pooled.

Statistical analyses

The variability in the five general prey categories collected by the cats was analysed using generalized additive mixed models (GAMM) with a Poisson error distribution and a log link. In the GAMMs, the number of prey items belonging to a given prey category in a given month, year and site was used as a response variable, and separate GAMMs were performed for each of these five categories (referred to as GAMMs 1 to 5). The models were based on a total of 307 monthly prey collections (i.e., prey brought by a cat or cats living in one site during one month) from 26 sites (i.e., cat owners) collected in 2002–2007. We used two explanatory variables in each GAMM: month (continuous variable, range: 1–12) and environment (categorical variable with two levels: rural vs. urban). The effect of month was fitted with a cyclic penalized cubic regression spline, i.e., splines whose ends match (Wood 2006), following the assumption that the response variable has similar values on the 31st of December and the 1st of January. We fitted the effect of month separately for the two environments (rural vs. urban) because the temporal dynamics of cat prey composition can differ between two environments. The number of knots for the spline fit was estimated to 5 to maintain a relatively simple fit. In each GAMM, site and year were used as random effects (the latter was fitted with a ridge penalty spline,

which acts as a random effect but improves the robustness of the model, Wood 2006). Moreover, the total number of all preys varied between the monthly collections from different sites and years, reflecting both the willingness of the cats to hunt and the owners to cooperate with us. Thus, we used the total number of preys in a given collection as an offset (in GAMMs 1 to 4) to keep the response variable (e.g., numbers of rodents, birds, etc.), independent of the variation in collection sizes. In case of GAMM 5 (for which the total number of preys was used as response variable), the number of hunting cats in a given site, month and year was instead used as an offset (i.e., it was assumed to be a proxy of sampling effort). We based the parameter estimation on the full models computed using the “mgcv” package (Wood 2006) in R (R Core Team 2015).

The seasonal variation in prey was described according to four seasons: winter (December – February), spring (March – May), summer (June – August), autumn (September – November).

We performed and displayed a canonical correspondence analysis (CCA) to identify the major relationships between the composition of cat prey and the two environments. In this analysis, we used 17 detailed prey categories as the response variables, whereas environment (rural vs. urban) was used as an explanatory variable. The CCA controlled for the hypothetical differences between years, months and sites by including these three variables as conditions in the analyses; their effects were partitioned out similarly to those of random factors in linear models (Oksanen et al. 2015). We tested the significance of the global model and the first two canonical axes with permutation tests with 5000 permutations. The analysis was performed using the ‘vegan’ package (Oksanen et al. 2015) in R.

Results

In total, we recorded 1348 prey items brought home by cats. Rodents were most common ($n = 836$ individuals) followed by birds (209), reptiles (131) and Soricomorpha (116). Mammals dominated cat prey in the rural environment (76.8%) with birds and reptiles accounting for 9.9 and 11.4%, respectively. In the urban environment, mammals accounted for approximately half of the prey killed, while birds were the second most common group (36.5%). In both environments, rodents were the most common mammalian prey (accounting for 83.1% of mammals in the rural and 96.3% in the urban environment) while soricomorphs composed 13.6 and 3.0% of mammalian prey, respectively. Murids were more frequently cat prey in urban than in rural environments (72.8% vs. 53.5%) (Table 1).

Soricomorphs and reptiles were killed significantly more often by cats living in rural than by those living in urban environments, while the reverse pattern was observed for birds (Table 2). There were no significant differences in

the number of rodents and the pooled number of all vertebrates brought home by cats living in the two environments (Table 2).

Season was a significant predictor of the composition of prey brought home by domestic cats. Rodents were most frequently caught by cats in autumn; shrews, moles and reptiles were more frequent in summer; and birds were most commonly brought home by cats in spring. When vertebrates were pooled, the highest number killed by cats was observed in autumn and the lowest in winter, and the seasonal variation in the number of prey was always higher in the rural environment (Fig. 1). This was most distinctly the case for soricomorphs and birds; the shares of these prey categories showed no temporal variation in the urban environment (the splines for the urban environment were not significant, Table 2) but significant nonlinear variation in the rural environment (Table 2, Fig. 1).

Canonical correspondence analysis showed that most of the mammals killed by cats (except mice) were associated with the rural environment, and lizards also followed this pattern. The birds in the prey composition, including sparrows (*Passer* spp.) and pigeons (Columbiformes), were largely associated with the urban environment (Fig. 2).

Discussion

In our study, we confirmed all three of the predictions posed in the introduction. First, cats in urban areas hunted more birds compared to cats living in the rural environment. Second, temporal variations in the prey brought home by cats seemed to follow changes in prey availability throughout the year; rodents were most often brought home in the autumn (with a peak around September and October, when the highest densities were reached) and birds in the spring (with a peak around June, when the young leave the nests), while reptiles were caught least often during late autumn and winter (from November to February, when they are inactive). Third, the temporal dynamics of the prey brought home by rural and urban cats were clearly different. Therefore, our data suggest a facultative feeding strategy in free-ranging house cats.

However, it should be kept in mind that the methodological approach used in this study may be somewhat biased. When free-ranging cats hunt, they may injure, capture or kill their prey and either leave it in the field, consume it in the field or bring it home (Fitzgerald and Turner 2000), so methods that are based on consumed prey (scat/stomach content analyse) give different results than those based on prey brought home (Krauze-Gryz et al. 2012a). Indeed, cats usually kill more prey than they actually bring home (Lloyd et al. 2013), so our study addressed only part of their hunting activity. Nevertheless, prey brought home may be assumed to be an index representing a minimum number of animals

Table 2 Summary of generalized additive mixed models (GAMMs 1–5) explaining the numbers of the five prey categories brought home by free-ranging domestic cats. Estimated degrees of freedom (Edf) are given for splines, and significant effects are in bold

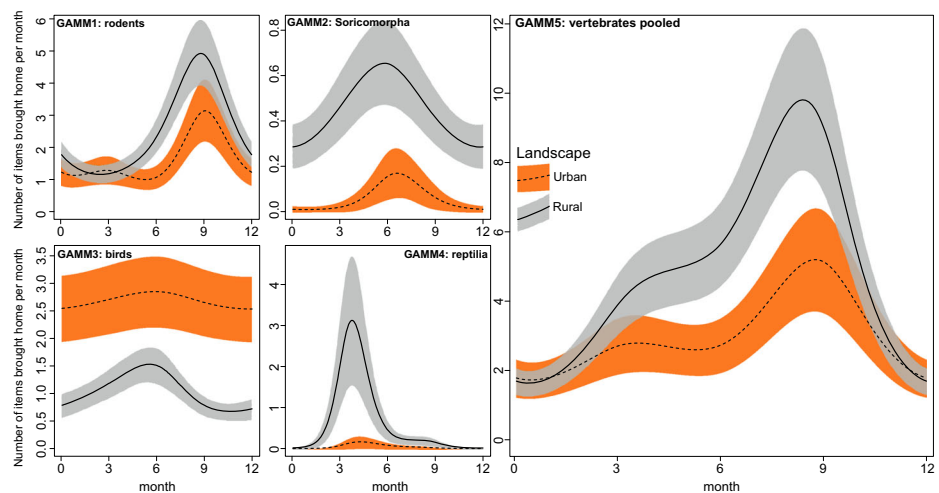
Model	Predictor	Estimate (SE)	Statistic	P-value
GAMM 1 Response: Rodents $R^2_{adj} = 0.34$	Intercept	-0.32 (0.28)	$t = 1.1$	0.2600
	Environment: rural	0.34 (0.35)	$t = 1.1$	0.2530
	Month _(rural)	Edf = 2.75	$F = 46.1$	0.0000
GAMM 2 Response: Soricomorpha $R^2_{adj} = 0.18$	Intercept	-3.93 (0.68)	$t = 5.8$	0.0000
	Environment: rural	2.37 (0.73)	$t = 3.3$	0.0013
	Month _(rural)	Edf = 1.85	$F = 3.0$	0.0052
GAMM 3 Response: Birds $R^2_{adj} = 0.14$	Intercept	-0.36 (0.10)	$t = 3.5$	0.0005
	Environment: rural	-0.97 (0.15)	$t = 6.5$	0.0000
	Month _(rural)	Edf = 1.78	$F = 2.4$	0.0151
GAMM 4 Response: Reptiles $R^2_{adj} = 0.12$	Intercept	-4.66 (0.98)	$t = 4.8$	0.0000
	Environment: rural	2.43 (1.11)	$t = 2.2$	0.0298
	Month _(rural)	Edf = 2.83	$F = 27.8$	0.0000
GAMM 5 Response: Vertebrates $R^2_{adj} = 0.16$	Intercept	0.58 (0.27)	$t = 2.2$	0.0313
	Environment: rural	0.40 (0.34)	$t = 1.2$	0.2372
	Month _(rural)	Edf = 2.95	$F = 82.8$	0.0000
	Month _(urban)	Edf = 2.75	$F = 11.5$	0.0000

killed and can show a general pattern in cat diets (Woods et al. 2003; Tschanz et al. 2010). The other limitation of this study is the division of the relatively small number of sites ($n = 26$) divided into the two environments (rural vs. urban). A larger number of sites distributed across different levels of urbanization would allow us to investigate variation over the full urbanization gradient and test for possible nonlinear relationships between local habitats and the hunting ecology of cats.

As in previous studies (Churcher and Lawton 1987; Barrat 1997; Fitzgerald and Turner 2000; Woods et al. 2003; Kays and DeWan 2004; Morgan et al. 2009; Tschanz et al. 2010; Krauze-Gryz et al. 2012a), mammals (mostly rodents) dominated the prey brought home by cats in both studied environments. However, their dominance was far lower in the urban

environment, where the share of birds in the prey clearly increased. Several former studies also indicate that pattern; in Finland, birds accounted for 24% of the prey of urban cats in contrast with 14% in the diet of rural cats (Kauhala et al. 2015). In the suburbs of Auckland (New Zealand), rodents dominated house cat kills in a habitat located on an urban/forest fringe, while birds were the most commonly caught urban vertebrate prey (Gillies and Clout 2003). The observed differences in the composition of prey brought home by cats living in urban and rural environments most likely reflected prey availability, which was driven by differences in land-use. Within the bird group, the share of pigeons and sparrows in the prey brought home by cats was higher in the urban environment. Sparrows were also the most frequent (Gillies and Clout 2003; Morgan et al. 2009) or among the most frequent

Fig. 1 The predicted number of the five prey categories brought home per month by free-ranging domestic cats living in rural (solid curve) and urban environments (dashed curve) throughout the year. Curves represent cyclic spline fits (accompanied by 95% confidence intervals indicated by transparent areas) as predicted by the GAMMs summarized in Table 2



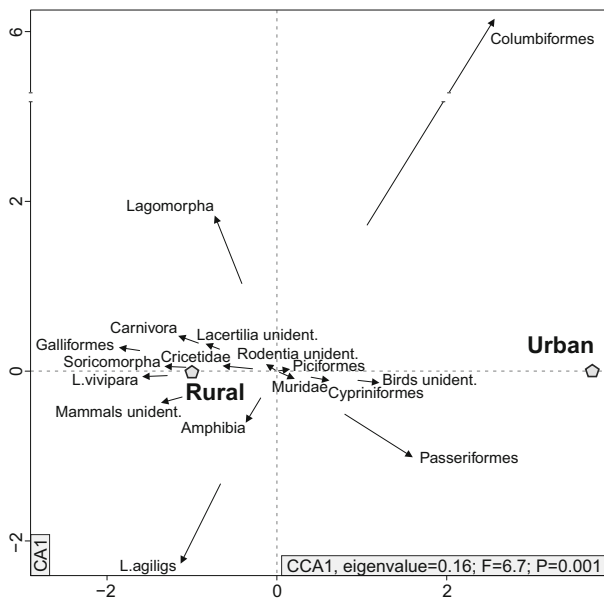


Fig. 2 Canonical correspondence analysis (CCA) showing the distribution of 17 prey categories brought home by free-ranging domestic cats in the two environments. The direction of the arrows indicates the correlation between prey category and environment type, and the length of the arrow indicates the strength of the correlation

(van Heezik et al. 2010) birds in urban and suburban habitats in New Zealand. Bird communities are usually less species rich in cities, but their abundance is relatively high compared to other environments (Rosin et al. 2016) with sparrows and pigeons being more common than in rural landscapes (Crooks et al. 2004). Thus, their share in the diet of opportunistic predators is usually higher in urban than rural environments, which was observed in studies of tawny owl (Goszczyński et al. 1993), martens and foxes (reviewed in Bateman and Fleming 2012).

Of the rodents killed by cats, the occurrence of mice was higher in the urban environment. Urbanization influences rodent species composition with the striped field mouse *Apodemus agrarius*, being a dominant species in considerably human-transformed habitats in Warsaw (Babińska-Werka et al. 1979; Gortat et al. 2014). Voles, on the other hand, are absent from the central quarters of Warsaw (Gryz et al. 2008), and their share in the rodent community decreases along an urbanization gradient (Andrzejewski et al. 1978; Gortat et al. 2014). Although populations of the striped field mouse reach very high densities in urban habitats, the species may be relatively difficult to catch due to dense vegetation (i.e., in parks, where rodents can hide in ivy cover), as observed for the tawny owl (Goszczyński et al. 1993). The two less common prey categories, soricomorphs and reptiles, also decreased in cat prey with increased urbanization. Our findings that the proportion of insectivores was higher in the rural than in the urban environment were similar to those of Kauhala et al. (2015).

Generally, shrews, being highly sensitive to fragmentation (Vergnes et al. 2013), are scarce in highly transformed, typically urban habitats (Andrzejewski et al. 1978; Gryz et al. 2008; Gortat et al. 2014).

We observed a clear temporal pattern in the composition of cat prey as different prey categories were caught in different proportions in different seasons. However, this seasonal variation was different in rural and urban environments, showing that urbanization affected the temporal patterns in the foraging ecology of cats. In our study, and similar to the results of other, rodents were most often caught in autumn and early winter (Barrat 1997; Weber and Daily 1998), and this variability is most likely driven by distinct changes in their abundance, which is the highest in autumn (Goszczyński 1977). Interestingly, this seasonal change in the number of rodents brought home by cats seemed to be smaller in the urban environment. According to Chernousova (2001), small mammal communities in cities are less dynamic, and rodent abundance remains relatively high compared to those in more natural areas. Another possible explanation for the autumnal increase in the numbers of rodents caught by cats in rural areas is their higher availability due to agricultural operations. After crops are harvested in the farmland (summer-autumn), rodents become much easier to hunt, and this is the time when cats more frequently penetrate fields (Goszczyński 1977; Krauze-Gryz et al. 2012b). The number of birds killed in rural areas fluctuated greatly, reaching a peak in spring (June) and a minimum in late autumn (November), while the number caught in the urban environment remained stable throughout the year. The peak in the number of birds killed by cats in rural areas, which primarily occurs in spring, probably reflects the killing of juveniles (e.g., Liberg 1984; Churcher and Lawton 1987; Barrat 1997; Baker et al. 2005; van Heezik et al. 2010), and several other generalist predators hunt birds most often during this season (e.g., Mirski et al. 2016). In winter, bird abundances are lower in rural areas due to seasonal migration, while in cities, most of the common species are sedentary (tree sparrow *Passer montanus*; house sparrow *Passer domesticus*; domestic pigeon *Columba livia domestica*), while migratory species are replaced by birds that winter in the city (e.g., Żmihorski et al. 2010); urbanization stabilizes winter bird communities (Suhonen et al. 2009). For example, birds can be hunted by cats next to bird feeders (Dunn and Tessaglia 1994). Woods et al. (2003) showed that cats living in households where birds were provided feed caught birds more often. The number of reptiles brought home by cats fluctuated greatly in rural areas. In the urban environment, where they were rather accidental prey, so the interseasonal change was rather small. Nevertheless, they appeared among the cat preys through spring and summer. Similarly, in Finland, reptiles were mainly brought home in the breeding season (Kauhala et al. 2015); in other seasons in the northern latitudes, when temperatures often drop

below zero Celsius degrees, reptiles remain inactive and thus unavailable to cats.

Our study suggests that the two critical impact periods of cats on native fauna is spring (for birds) and autumn (for rodents), while during winter, cats catch fewer prey (especially in the rural environment) and are less active (Goszczyński et al. 2008). Thus, their overall predation pressure is relatively low. Rural cats, which are mainly kept as ‘mousers’ (Krauze 2008), are perceived as effective rodent killers, but in addition to killing synanthropic rodents, such as house mice *Mus musculus*, and rats *Rattus* spp., they prey on numerous species of voles as well as legally protected or rare species (e.g., shrews; hares *Lepus europaeus*; red squirrels *Sciurus vulgaris*; least weasels *Mustela nivalis*) (Krauze-Gryz et al. 2012a; this study). The effect of predation by free-ranging domestic cats on prey populations can be severe as their numbers are kept artificially high by supplemental feeding (Sims et al. 2008). At the same time, they do not show normal numerical or functional responses to prey density (Coleman and Temple 1993) as they switch between household food and natural prey depending on accessibility (Liberg 1984; Weber and Daily 1998). As a result, cats can efficiently compete with wild predators (Krauze-Gryz et al. 2012b). Urban and suburban cats mainly focus on birds, which results in serious predation rates (Lepczyk et al. 2004) or increases in sub-lethal factors, such as a reduction in fecundity (Beckerman et al. 2007) or food delivery to chicks (Bonnington et al. 2013). Special attention should be paid to the influence of house cats on the fauna of nature reserves located in or adjacent to cities or suburbs as the presence of free-ranging domestic cats, which are likely the most abundant predator, can reduce the effectiveness of these protected areas as a tool for protecting nature (Wierzbowska et al. 2012).

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