Landscape structure, human disturbance and crop management affect foraging ground selection by migrating geese

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Abstract It is well known that agricultural intensification has caused severe population declines among bird species which use farmland for breeding and overwintering, while migrating bird species may benefit from intensive farming, but in turn damage crops. Knowledge of the habitat selection of migrating birds is important from both a conservation and agro-economic point of view. We investigated the habitat preferences of three common migrating goose species: White-fronted Goose *Anser albifrons*, Bean Goose *A. fabalis* and Greylag Goose *A. anser* during the autumn of 2009 in western Poland. A total of 24 flocks of these species were identified. Geese preferred large, elevated fields that were remote from forests and human settlements but in close proximity to a lake. Geese selected maize stubbles and avoided winter cereals. They selected sites in landscapes with a lower diversity of crops. Flock size was negatively correlated with the proportion of pastures in the landscape, but it increased with field size, distance to forest and distance to town. Our results are in contrast with the paradigm that less intensive farmland positively influences habitat use by birds during foraging. We advise the delayed ploughing of stubbles with the aim of creating appropriate foraging habitats for geese and minimizing damage to cereal crops.

Keywords Habitat preference · Intensive farming · Crop damage · Maize stubble · Human disturbance · *Anser* spp

Zusammenfassung

Landschaftsstruktur, menschliche Störung und Anbaumethoden beeinflussen die Nahrungsflächenwahl wandernder Gänse

Die Intensivierung der Agrarwirtschaft hat starke Populationsrückgänge sowohl brutender als auch überwinternder Vogelarten in landwirtschaftlich genutzten Flächen verursacht. Dennoch können ziehende Vogelarten von der intensiven Landwirtschaft profitieren, aber auch Kulturpflanzen schädigen. Sowohl aus Sicht des Naturschutzes, als auch aus agrarökonomischer Sicht ist es wichtig, Kenntnisse über die Habitatwahl der Gänse zu erlangen. Vor diesem Hintergrund untersuchten wir im Herbst 2009 in Westpolen die Habitatpräferenzen dreier häufiger und ziehender Gänsearten: Blässgans *Anser albifrons*, Saatgans *Anser fabalis* und Graugans *Anser anser*. Insgesamt fanden wir 24 Trupps dieser Arten. Die Gänse bevorzugten große, erhöhte Felder, die in der Umgebung eines Sees und abseits von Wäldern und menschlichen Siedlungen lagen. Die Gänse wählten Maisstoppeln und mieden Wintergetreide. Sie selektierten Gebiete in Landschaften mit einer geringen...
Diversität an Anbaupflanzen. Die Truppgröße war negativ korreliert mit dem Grünlandanteil der Landschaft. Truppgrößen erhöhten sich mit der Feldgröße und mit den Distanzen zu Wäldern und Siedlungen. Unsere Ergebnisse stehen im Gegensatz zum Paradigma, dass weniger intensive Landnutzung die Habitatnutzung der Vögel während der Nahrungssuche positiv beeinflusst. Wir empfehlen ein verspätetes Pflügen der Stoppeln, um geeignete Nahrungs- habitats für Gänse zu schaffen und damit die Schädigung von Getreidepflanzen zu minimieren.

Introduction

Many bird species which breed and overwinter in farmland habitats have suffered severe population declines in recent decades (Donald et al. 2006). It is generally acknowledged that agricultural intensification with associated breeding habitat loss and deterioration are the major causes of these declines in numbers (Nagy et al. 2009; Rodríguez-Teijeiro et al. 2009). However, much less attention has been paid to species that select various farmland habitats during migration (Lindström et al. 2010). Moreover, there are several bird species that might benefit from intensive agricultural landscapes during migration, such as the Golden Plover Pluvialis apricaria (Lindström et al. 2010), the Common Crane Grus grus (Leito et al. 2008) and geese (e.g., Fox et al. 2005; Van der Graaf et al. 2007; Jensen et al. 2008), and which in turn may have important impacts on ecosystem processes and the agro-economy. Consequently, understanding how such migratory birds select foraging sites during migration is desirable (Lindström et al. 2010; Sebastian-González et al. 2010; Batáry et al. 2011).

The selection of foraging habitat during migration is central to the life history of these birds and greatly influence their fitness (Cody 1985; Farmer and Wiens 1999; Drent et al. 2006; Duriez et al. 2009). Migrating birds depend mostly on the availability of resources at stopover sites that influence survival and recuperation (Bauer et al. 2008; Alerstam 2011). Although many studies have focused on the physiology of migration and strategies for energy refuelling (e.g., Tinkler et al. 2009; Guglielmo 2010), much less is known about site choice where birds forage (Newton 2006).

The selection of foraging sites by birds in agricultural landscapes is highly influenced by human-related factors (Olsson et al. 2002; Madsen and Boertmann 2008). Firstly, human disturbance may directly affect bird behaviour (Sirot 2010) and probably thereby their strategies of feeding site choice. Secondly, predation pressure may be mediated by human-related management of agricultural landscapes (e.g., Pita et al. 2009; Wang et al. 2011). Landscape composition also affects the foraging decisions of animals (Cresswell and Osborne 2004; Skórka et al. 2009). Migrating birds foraging in agricultural landscapes may seek foraging patches rich in resources, such as pastures and/or set-aside or unploughed fields (Olsson et al. 2002; Lindström et al. 2010). The identification of these sites and an understanding of which factors affect their selection by birds may be of key importance for land management, successful conservation and reduced crop damage (Jensen et al. 2008).

Most goose populations in the Western Palearctic have recovered substantially during the last three to five decades (Madsen et al. 1999; Andersson et al. 2001). This has led not only to the expansion of their breeding range, but also to an increase in the number and size of migrating and wintering goose flocks and, in turn, to damage to crops and other complaints of nuisance (e.g., Roomen and van Madsen 1992). However, current goose population trends vary in different areas (Lawicki et al. 2010). Additionally, large concentrations of goose flocks are recognized as one of the criteria to create Natura 2000 areas (Council Directive 2009). Among 174 Important Bird Areas of international importance in Poland, 21 were created especially to conserve migrating and wintering Bean Goose Anser fabalis and 13 for White-fronted Goose Anser albi-frons based on the criteria of BirdLife International (2004) and Wilk et al. (2010). However, in many European countries, the most numerous migrating goose species, namely, the White-fronted Goose, Bean Goose and Greylag Goose A. anser, are game species (Council Directive 2009). Thus, there are often conflicts between goose conservation and measures aimed at controlling their numbers in farmland (Jensen et al. 2008).

The aim of this paper was to further our understanding of foraging site selection by migrating geese in agricultural landscapes in western Poland. Using a general approach, we studied how agricultural methods, crop diversity at the landscape scale, human-related factors, forests and lakes, and site elevation affected the foraging habitat choice of three species of migratory geese.

Methods

Bird surveys

Eight large plots (100 km$^2$) were selected (through the generation of random geographical coordinates in QuantumGIS software) as study areas in agricultural landscapes in western Poland (Fig. 1). This is the most important agricultural region in Poland (Jankowiak et al. 2003; Tryjanowski et al. 2011) and is representative of the main
staging and wintering goose areas in Central Europe (Lawicki et al. 2010). The preferred type of agriculture activities in Wielkopolska is cereal crop farming and cattle and pig breeding, but areas of intensive cultivation have been expanding recently. On a national scale, the region dominates the production of sugar beet and other vegetable crops (Jankowiak et al. 2003). During the autumn of 2009 (October–November), the plots were surveyed from cars. Within each plot, observers stopped several times to search for geese using binoculars (10–12 × 50) and telescopes (30–60 ×). The observation points were arranged so that the entire plot area was covered by visual observation. One count, lasting approximately 5 h, was conducted in each plot. The surveys were conducted during favourable weather conditions (no rain or snowfall). When a goose flock was found, its species composition, the number and age of birds, as well as the geographical coordinates of the site were noted. We counted birds from a distance so as not to disturb their foraging and movements.

Habitat and landscape characteristics of the foraging sites

In the sites where geese were found we measured several variables that potentially could influence site-selection by the birds (Table 1).

In general, variables 1–9 (Table 1) describe foraging habitat (foraging patch size and availability of various food resources) since geese often forage in agricultural areas. Variables 10–13 were indicators of human disturbance/pressure. We distinguished between local roads and regional/national roads because the former are characterized by little traffic (usually much fewer than 100 vehicles per hour), whereas the latter represent roads with >500 vehicles per hour (authors’ unpublished observations). Variables 14–15 may be indicators of the site’s safety from predators, variables 16–17 describe the proximity of roosting areas or migration/dispersal corridors, as has been demonstrated for other waterbirds (Skórka et al. 2009; Maciusik et al. 2010). The lake size of 10 ha was chosen because it was the smallest lake where roosting geese were observed in the area (authors’ unpublished observations).

To check if the geese preferred any of these features we selected random sites equal in number to the number of geese flocks \(n = 24\). We selected random geographical coordinates within the plot boundaries. In a given plot, the number of random sites was equal to the number of geese flocks found there. At random points we measured the same variables as for geese flocks (Table 1).

Variables 1, 10–14 and 16–17 (Table 1) were measured from satellite images and calculated in ImageJ and QuantumGIS software. Variables 2–8 and 14 were derived by direct mapping in the field. We recorded crop type on the maps during field surveys and, later, scans of these maps were digitalized in QuantumGIS software. Variable 9 was calculated in the BioDiversity Pro Software (McAleece 1997).

Data processing and statistical analysis

To test whether geese had preferences for crop type during foraging, we used Fisher’s exact test and compared the number of observations of geese flocks in different crops with the equivalent from random points (expected numbers). If geese had no preferences for crop type, the number of flocks and number of random points in a given crop type should be similar. The same test was used to check if geese had preferences for an elevation category.

To compare whether sites where geese foraged differed in landscape composition from random sites, we used general linear mixed models (GLMM). We tested if the mean values of the environmental variables measured at sites selected by geese differed from mean values at random sites. The categorical independent variable was foraging or random site. Because observations were conducted in eight large plots, we assigned their identity as a random effect to account for spatial autocorrelation (Dormann et al. 2007).

Spatial autocorrelation was tested by Moran’s local indicator (Legendre and Legendre 1998). The spatial autocorrelation value at a given distance class indicates how predictable (positively or negatively) the measured variable (goose flock size in our study) was at a given point.
of the sampling framework. Moran’s index varies between -1 and 1, with non-significant values close to zero.

To test whether the number of geese was dependent on measured variables, we used modified Pearson correlation coefficients that take into account the pattern in spatial autocorrelation (Dutilleul 1993).

We did not build multivariate models because the number of geese flocks was low (in a statistical sense) compared to the number of variables that potentially affected site choice and flock size. Instead, we examined in detail Pearson spatial correlation coefficients between the continuous explanatory variables to identify possible confounding effects due to multicollinearity. This could lead to biased statistically significant results of GLMM resulting simply from strong correlations between environmental variables. When collinearity between two variables was found, we calculated a simple regression between them, and residuals (part of the variation of one variable not explained by the second) were included in the GLMM. The GLMMs were built in the same manner as described above.

The GLMMs and correlation analyses are sensitive to outliers; therefore, where necessary, data were \( \log_{10} \) or \( \log_{10}(x + 1) \) transformed to minimize effects of the outliers and to homogenize variances (Quinn and Keough 2002). The Grubbs tests (Grubbs 1969) indicated that the analyses based on raw and logarithmically transformed data (in the case of skewed distributions) were not affected by outliers (all \( P > 0.100 \)). Variables expressed as proportions were arcsin-transformed. Corrected means calculated on \( \log_{10} \)-transformed data and then back-transformed in the case of skewed distributions are presented throughout the paper (Quinn and Keough 2002). Means are given with their 95% confidence intervals (CIs) that are also calculated on \( \log_{10} \)-transformed data where necessary, and then back-transformed (and hence will be asymmetric).

All calculations were done in JMP 9 (2010) and SAM 4.0 (Rangel et al. 2010) software.

### Results

**Foraging flock size, species composition, and spatial autocorrelation**

We noted 24 goose flocks during the surveys. The mean number of flocks per plot was 3.0 (95% CI 1.4–4.4, median 2.5, range 1–6 flocks per plot). Three goose species were present in the flocks: White-fronted Goose, Bean Goose and Grey-lag Goose (Table 2). Overall, 40,926 birds were counted during surveys. The Bean Goose was the most

| Number | Code         | Characteristics                                                                 |
|--------|--------------|---------------------------------------------------------------------------------|
| 1      | FieldSize    | Field size (ha)                                                                  |
| 2      | PPloughed    | Percentage cover of ploughed land in a 500-m radius                              |
| 3      | PBeet        | Percentage cover of beet stubble in a 500-m radius                               |
| 4      | PRape        | Percentage cover of rape stubble in a 500-m radius                               |
| 5      | PMaize       | Percentage cover of maize stubble in a 500-m radius                              |
| 6      | PWinterCere  | Percentage cover of winter cereals in a 500-m radius                             |
| 7      | PPastures    | Percentage cover of pasture in a 500-m radius                                    |
| 8      | PSetAside    | Percentage cover of set-aside in a 500-m radius                                  |
| 9      | HabitatDiver | Simpson reciprocal diversity index (1/D) calculated from the percentage cover of |
|        |              | crops (Simpson 1949). The value of this index has 1 as the lowest possible figure. |
|        |              | This figure would represent a composition containing only one cover type. The    |
|        |              | higher the value, the greater the diversity of crop types.                       |
| 10     | DLocRoad     | Distance (m) to the nearest local road from the centre of the foraging flock     |
| 11     | DRegRoad     | Distance (m) to the nearest regional/national road                              |
| 12     | DHumanS      | Distance (m) to the nearest human settlement                                    |
| 13     | DTown        | Distance (km) to the nearest town with >5,000 citizens                           |
| 14     | DForest      | Distance (m) to the nearest forest                                               |
| 15     | Elevation    | Site elevation, coded as a simple categorical variable: -1 (a site located      |
|        |              | below the surrounding area), 0 (a site located at the same elevation as the      |
|        |              | surrounding landscape), 1 (a site at a higher elevation than the surrounding    |
|        |              | landscape). The classification was based on the prevailing elevation of          |
|        |              | landscape in a radius of 500 m from the centre of the foraging flock (e.g., if   |
|        |              | 60% of the landscape within 500 m was below elevation of the site selected by    |
|        |              | the flock it was classified as elevated and coded as (1)                         |
| 16     | DRiver       | Distance (m) to the nearest river longer than 50 km                              |
| 17     | DLake        | Distance (m) to the nearest lake >10 ha                                           |
numerous (80.8% of all individuals), with the White-
fronted Goose and Grey-lag Goose being much less
numerous (15.0 and 4.2% of all individuals, respectively; 
Table 2). Mean goose flock size was 1,166.4 birds (95% CI
787.0–1,728.6, median size 1,375, range 236–5,800). We
found a significant positive spatial autocorrelation in flock
size of geese at small distances (2 km) and a statistically
significant negative spatial autocorrelation at distances of
10–30 km (Fig. 2).

Habitat and landscape factors affecting foraging site
selection by geese

Sites selected by geese were more often located on maize
stubbles than expected by chance (from crop types found at 
random sites; Fisher exact test $P < 0.001$; Fig. 3). Sites
selected by foraging geese were more often located at
higher elevation in relation to the surrounding landscape
than expected by chance (Fisher exact test $P < 0.001$; 
Fig. 4), and fields selected by foraging geese were on
average larger than expected (Table 3). Sites selected by
geese were located in a landscape with a higher cover of
maize stubble and a lower cover of ploughed fields and
winter cereals (Table 3). When we took the cover of all
crop types and calculated their diversity within a land-
scape, we found that geese selected sites in landscapes with
a lower diversity of crops (Table 3). Sites selected by
foraging geese were on average more distant from human
settlements, local roads and forests than random sites
(Table 3). Furthermore, sites selected by foraging geese
were closer to lakes than randomly selected sites (Table 3).

Factors affecting flock size of foraging geese

The size of goose flocks was positively correlated with
field size, distance to nearest forest and distance to the
nearest town, but negatively correlated with proportion of
pasture in the landscape (Table 4; Fig. 5). When we

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**Table 2** Frequency of goose species in flocks and their abundance

| Goose species          | Frequency in flocks ($n = 24$) | Number of individuals per flock |
|------------------------|---------------------------------|--------------------------------|
|                        |                                 | Mean | Lower–upper 95% CI | Median | Range   |
| White-fronted goose   | 23                              | 175.9 | 114.0–271.6    | 200    | 30–1,200|
| Bean goose             | 22                              | 895.7 | 574.8–1,395.7  | 800    | 180–5,500|
| Grey-lag goose        | 7                               | 108.9 | 16.3–729.7     | 309    | 2–400   |

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*a* Both means and 95% confidence intervals (CIs) were calculated on log$_{10}$-transformed data and then back-transformed.
analysed numbers of particular species within foraging flocks, we found that the number of White-fronted Goose was positively correlated with distance to the nearest local road and distance to the nearest forest but negatively correlated with the proportion of pasture in the landscape (Table 4; Fig. 5). The number of Bean Goose was positively correlated with distance to the nearest regional road and distance to the nearest town (Table 4; Fig. 5), and the

Table 3 Habitat and landscape characteristics of the random points and of sites selected by geese for foraging during the autumn

| Variable                  | Random points | Foraging sites | GLMM F | df | P      |
|---------------------------|---------------|----------------|--------|----|--------|
|                           | Mean          | Lower–upper 95% CI | Mean   | Lower–upper 95% CI |        |
| FieldSize (ha)            | 7.2           | 4.6–11.1        | 21.5   | 14.0–32.8        | 16.177 |
| PPloughed (%)             | 9             | 7–12           | 3      | 1–5             | 13.351 |
| PBeet (%)                 | 1             | 0–2            | 3      | 0–7             | 3.012  |
| PRape (%)                 | 12            | 7–18           | 24     | 16–34           | 3.791  |
| PMaize (%)                | 5             | 1–9            | 16     | 10–21           | 4.126  |
| PWinterCere (%)           | 52            | 47–58          | 35     | 26–46           | 5.569  |
| PPastures (%)             | 5             | 2–8            | 4      | 1–8             | 0.786  |
| PSetAside (%)             | 3             | 1–6            | 1      | 0–2             | 4.244  |
| HabitatDiver (%)          | 2.57          | 2.33–2.82      | 2.11   | 1.94–2.32       | 5.485  |
| DLocRoad (km)             | 0.22          | 0.17–0.27      | 0.46   | 0.35–0.60       | 17.443 |
| DRegRoad (km)             | 1.75          | 1.16–2.496     | 2.11   | 1.60–2.72       | 1.656  |
| DHumanS (km)              | 0.33          | 0.25–0.42      | 0.51   | 0.42–0.62       | 4.046  |
| DTown (km)                | 6.87          | 5.25–8.49      | 7.95   | 6.21–9.69       | 2.802  |
| DForest (km)              | 1.02          | 0.67–1.36      | 2.31   | 1.89–2.73       | 22.568 |
| DRiver (km)               | 3.06          | 2.20–4.11      | 4.02   | 2.83–5.57       | 2.106  |
| DLake (km)                | 2.79          | 2.19–3.39      | 1.64   | 1.14–2.13       | 10.110 |

Data are the results of univariate generalized linear mixed models (GLMM)

For explanation of the variables: see Table 1

* Significant results at P < 0.05

a Both means and 95% confidence intervals (CIs) were calculated on log 10 –transformed data and then back-transformed

Table 4 Pearson correlation coefficients adjusted for spatial autocorrelation (P value in brackets) between number of individuals of all geese species, number of individuals of particular species and habitat and landscape features measured at sites selected by geese

| Effect                  | All species | White-fronted Goose (n = 23 flocks) | Bean goose (n = 22 flocks) | Grey-lag goose (N = 7 flocks) |
|-------------------------|-------------|-------------------------------------|-----------------------------|--------------------------------|
|                         | Effect      | Mean (95%)                          | Mean (95%)                  | Mean (95%)                     |
| FieldSize               | 0.460 (0.050)* | 0.283 (0.199)                       | 0.379 (0.094)              | 0.318 (0.178)                  |
| PPloughed               | -0.379 (0.105) | -0.081 (0.77)                        | -0.386 (0.138)             | 0.120 (0.687)                  |
| PBeet                   | -0.230 (0.275) | -0.269 (0.191)                       | -0.151 (0.449)             | -0.286 (0.203)                 |
| PRape                   | 0.060 (0.810)  | 0.050 (0.791)                        | 0.071 (0.768)              | 0.021 (0.920)                  |
| PMaize                  | 0.430 (0.167)  | 0.238 (0.225)                        | 0.399 (0.199)              | -0.065 (0.774)                 |
| PWinterCere             | -0.090 (0.591) | 0.092 (0.652)                        | -0.133 (0.353)             | 0.200 (0.326)                  |
| PPastures               | -0.467 (0.031)* | -0.381 (0.037)*                      | -0.387 (0.104)             | -0.135 (0.504)                 |
| PSetAside               | -0.149 (0.392) | -0.064 (0.785)                       | -0.125 (0.498)             | -0.153 (0.504)                 |
| HabitatDiver            | -0.391 (0.132) | -0.110 (0.630)                       | -0.381 (0.169)             | 0.010 (0.971)                  |
| DLocRoad                | 0.208 (0.243)  | 0.459 (0.035)*                       | 0.088 (0.643)              | 0.284 (0.212)                  |
| DRegRoad                | 0.366 (0.107)  | 0.300 (0.242)                        | 0.478 (0.044)*             | 0.360 (0.159)                  |
| DHumanS                 | 0.480 (0.107)  | 0.144 (0.439)                        | 0.477 (0.116)              | -0.130 (0.529)                 |
| DTown                   | 0.716 (0.015)* | 0.382 (0.195)                        | 0.650 (0.034)*             | 0.075 (0.797)                  |
| DRiver                  | 0.677 (0.002)* | 0.677 (0.003)*                       | -0.043 (0.858)             | 0.570 (0.045)*                 |
| DLake                   | 0.436 (0.171)  | -0.100 (0.699)                       | 0.496 (0.128)              | -0.245 (0.267)                 |

* Significant correlations at P < 0.05

For explanation of the variables: see Table 1
number of Grey-lag Goose was positively correlated with distance to the nearest forest (Table 4; Fig. 5). Multicollinearity between environmental variables and analysis of residuals

We found five statistically significant correlation coefficients between variables (Appendix). Cover of ploughed fields and cover of winter cereals were negatively correlated. The GLMM based on residuals not explained by the cover of winter cereals revealed that sites selected by geese had a lower cover of ploughed fields than random sites ($F_{1,39.8} = 16.55, P < 0.001$). We also found two statistically significant correlation coefficients between habitat diversity index and the cover of ploughed fields, and between diversity and the cover of winter cereals. However, since the

Fig. 5 Statistically significant correlations between goose flock size and field size (a), cover of pastures in a landscape (b), distance to nearest town (c) and distance to nearest forest (d). Analyses for individual species showed that flock size of White-fronted Goose Anser albifrons were significantly correlated with cover of pastures (e), distance to local road (f) and distance to forest (g). Flock size of the Bean Goose A. fabalis correlated with distance to local road (h) and distance to town (i). Flock size of Grey-lag Goose A. anser correlated with distance to forest (j).
index is composed from the other variables, this correlation was not unexpected. Therefore, it made little sense to calculate residuals between these variables and conduct a GLMM. We found two additional statistically significant correlation coefficients (Appendix); however these correlations did not lead to false significant results in our GLMMs because rape cover and distance to the river were similar in sites selected by geese and random sites (see Table 3; Appendix).

Discussion

Our results show that geese foraging in farmland during the autumn migration in western Poland select specific crops and sites located in areas guaranteeing low disturbance both by predators and humans. Contrary to most studies on declining farmland bird populations during the breeding season, geese selected those parts of farmland that were intensively managed, preferring larger field sizes and lower crop diversity. Flocks of foraging geese also had a predictable spatial structure, as indicated by spatial autocorrelations.

Crop preferences of geese

We found that geese avoided fields of winter cereals but were frequently found at stubbles of maize, beet, and rape (as revealed by Fisher exact tests). These preferred crop types were found much less often at randomly chosen sites. Moreover, a cover of maize stubbles in a landscape positively affected site choice by geese, whereas the opposite result was found for a cover of winter cereals and ploughed fields. Stubbles are favourite foraging sites even when their share in a landscape is low (Madsen 1985a; Jepsen 1991; Gill 1996). For example, geese have been found to prefer stubbles of beet (Gill 1996; Gill et al. 1997; Nilsson and Persson 2000), maize (Frederick and Klaas 1982) and rape (Nilsson and Persson 2000). The preference for these crop types may be easily explained by the high energy value of these food types, which are needed to build up the reserves during the autumn migration (Raveling 1979). During harvesting, many corn cobs drop to the ground, and some plant material is also left, providing a good source of energy for geese. Moreover, geese may forage on weeds growing among maize stems (Reed et al. 1977; Frederick and Klaas 1982). However, the preferences of geese for winter cereals differ between studies; for example, Nilsson and Persson (2000) considered this crop type to be an important food type for geese, whereas Jepsen (1991) found the opposite. These differences might be due to different crop availability and/or weather conditions since it has been proven that geese graze on winter cereals when stubbles are ploughed or the ground is frozen (Nilsson and Persson 1991). It has also been shown that the change from their natural food, *Scirpus* tubers, to agricultural areas was gradual and that geese first turned to crops similar to their natural food (pasture, winter cereals) but gradually learned to take advantage of various crop types rich in energy (Gill et al. 1997; Prop et al. 1998; Nilsson and Persson 2000).

Spatial pattern in flock size

We detected significant spatial autocorrelation in the size of flocks. Spatial autocorrelation may lead to increased type I error in analyses and the determination of too many significant effects (Dormann et al. 2007). However, we accounted for this problem in almost all our analyses. In a biological sense, spatial autocorrelation may lead to spatial synchrony that indicates coincident numbers of individuals (Liebhold et al. 2004). In our study, we found significant positive spatial autocorrelation for the flock size at distances up to 2 km. Spatial autocorrelation was also found in a study on Pink-footed Goose in Norway (Jensen et al. 2008). Taken together, these results indicate that there is some similarity in the size of flocks foraging close together in the landscape. This similarity may result from sites located close to each other having a more similar landscape structure than those further apart. If geese prefer specific landscape features (e.g. field size), their flocks may be spatially bound to these features which, in turn, limits flock sizes (Spilling et al. 1999), leading to the observed pattern of spatial autocorrelation. We believe that the odds of counting the same flock twice were minimal because we did not disturb foraging birds and one plot was visited only once within a restricted time framework. Therefore, the spatial autocorrelation found in our study likely did not result from the movements of individuals.

Predators and/or disturbance during foraging in farmland

In this study we documented that geese probably chose sites that were higher than the surrounding landscape. We believe that this result is linked with predation or disturbance avoidance. Foraging animals must cope with the potential risk of predation (Lima 1998; Jonker et al. 2010). Sites that allow the early detection of approaching predators should therefore be preferred (Lima 1987; McNamara and Houston 1992), and higher elevation inevitably enables geese to observe what is going on in their surroundings. The study of Madsen (1985b) also indicated that geese avoid sites with limited visibility. However, it has also

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been reported that some goose species, such as the Pink-footed Goose, avoid sites located at higher elevation in farmland (Jensen et al. 2008). Our landscape plots were located in a lowland region (50–152 m a.s.l) with little variation in elevation. It is possible that geese generally prefer regions located at low elevation, but within a given region, they choose sites slightly raised above the surrounding landscape. In our study, the scale at which relative elevation was measured was quite simple, but there is no other good method to measure relative elevation at the given sites. The effect of site elevation in relation to the surrounding landscape should be incorporated in a more sophisticated manner in future studies on the landscape ecology of geese.

Geese were found to avoid the proximity of forests. This result may also be explained by predator avoidance. In Poland, forest edges and small mid-field forests are a major habitat of the opportunistic Red Fox Vulpes vulpes (Tryjanski et al. 2002; Jankowiak et al. 2008) are used as well by the White-tailed Eagle Haliaeetus albicilla that preys on geese (Roder et al. 2008) and whose appearance causes entire goose flocks to take flight (personal observations). Other explanations for avoidance of forest edges is that hunters often hunt from hides at forest edges and forestry workers may occasionally work and appear at forest edges, thereby disturbing geese foraging nearby (Klaassen et al. 2006).

We found that sites selected by geese were more distant from roads and human settlements. In our study, geese avoided local roads, but we did not find an effect of the presence of regional roads with higher traffic. In fact, local roads were, on average, closer to foraging sites than regional roads, and it has been shown, for example, that the Pink-footed Goose A. brachyrhynchos responds negatively to even very little road traffic and rarely forages in fields close to roads (Madsen 1985b; Gill 1996; Jensen et al. 2008). Although the proximity of regional roads did not affect foraging site choice in our study, it was an important factor influencing flock size of the Bean Goose, with the flock size of this species being larger in fields further from these roads. This is an interesting result and suggests that road traffic may affect the social behaviour of the studied species. Human settlements negatively affecting foraging site selection by geese in farmland. The effect may be linked with two phenomena. Firstly, farmers and local inhabitants may directly disturb foraging birds around their properties due to normal human activity or by implementing scaring practices to protect crops (Tombre et al. 2005; Klaassen et al. 2006; Jensen et al. 2008). Secondly, there is an abundance of freely roaming dogs and cats in Polish villages. Wandering dogs in particular may be a real threat to foraging geese, resulting in increased vigilance (Randler 2006). It has been shown that domestic dogs, as well as foxes, operate mostly in the neighbourhood of human settlements (Jankowiak et al. 2008; Olko et al. 2009). We also found an interesting effect of town proximity on the flock size of foraging geese. Flock size increased with increasing distance to the town. To the best of our knowledge this is the first such result found for geese. Towns are associated with greater human activity and higher road traffic, both of which probably disturb flocking behaviour during migration.

The proximity of lakes positively affected the use of farmland by geese during migration. During migration, lakes are usually used by geese as roosting and daily resting sites. This may explain the positive effect of this habitat on the presence of geese in farmland. Other authors have also found that geese primarily use fields close to roosting sites (e.g., Gill 1996; Jensen et al. 2008).

Is a less intensive farmland habitat always beneficial for birds?

Agricultural intensification has led to a loss of habitat heterogeneity and increased field size (Benton et al. 2003; Reif et al. 2010) and, as a consequence, to a widespread decline in farmland biodiversity across many different taxa (Donald et al. 2001; Fuller et al. 2004). In the context of foraging behaviour, loss of habitat heterogeneity may also diminish the availability of food resources for foraging farmland birds (Surmacki 2005; Wilson et al. 2005).

Our results are in sharp contrast with the paradigm that a less intensive use of farmland (greater habitat heterogeneity and small field sizes) positively influences habitat use during foraging. Geese chose sites characterized by a lower diversity of crop types and preferentially foraged on fields that were larger than those generally available. Flock size was also greater on larger fields. On the other hand, our results are consistent with the theoretical background on foraging behaviour. Optimal foraging theory (Stephens and Krebs 1986; Kramer 2001) predicts that birds should prefer foraging in larger patches (larger fields). Other authors have also found that geese preferentially forage in larger fields (e.g. Gill 1996; McKay et al. 2006) as well as in more intensively used farmland (Fox et al. 2005). However, we are not aware of any earlier studies showing that crop diversity negatively affected foraging site choice by geese during migration. Thus, geese are one of the few examples of species which may benefit from intensive agriculture (Van Eerden et al. 1996; Fox et al. 2005). However, site choice by geese is inevitably different from that of the declining populations of farmland birds which are mostly territorial during the breeding season. Goose flocks are not territorial and may move long distances to find suitable foraging sites. Therefore, it would be...
interesting to study the effects of habitat diversity on foraging site selection by geese on larger or multiple spatial scales. In our study, we measured crop diversity in a 500-m radius, which is a rather small scale compared to the actual movement of geese. However, all management practices (e.g., changing crop type, time of ploughing) are cheaper and easier to implement at smaller spatial scales (Mouysset et al. 2011).

Consequences for goose conservation and farmland management

Geese are considered to be both species of conservation interest and pests of agriculture (Jepsen 1991; Jensen et al. 2008). There is a clear need to manage land in such a way as to alleviate this conflict (Jensen et al. 2008). A study on the Pink-footed Goose in Norway noted that to achieve this goal, the areas where migrating geese stop to forage should first be identified (Jensen et al. 2008). Such locations should then be designated as foraging refuges for geese and payments subsequently made to farmers (who agree not to disturb foraging geese) active in or near these locations (Jensen et al. 2008). Our results suggest that for such designated foraging refuges to remain attractive to geese, management programmes should consider implementing a number of measures, including larger field size in an open landscape guaranteeing visibility and the absence of human disturbance (fields far from roads and human settlements) (see also Jensen et al. 2008). Additionally, the major crop should be maize (and also beet and rape). To reduce damage to winter cereals, farmers should increase the length of time geese can spend on stubbles (Gill 1996) and increase the area of stubble by leaving maize, beet and rape stubbles in the fields for longer periods of time.

However, some farmers may not be willing to support management policies that benefit geese due to their potential to damage crops (Kahl and Samson 1984; Jensen et al. 2008). Unexpectedly, scaring of foraging geese only enhances the conflict and proportion of farmland impacted by geese, as has been convincingly demonstrated for Pink-footed Goose (Jensen et al. 2008). Therefore, other solutions for the goose problem must be found. We suggest that farmers may diversify their crops and reduce field size in areas heavy impacted by geese. As we have shown, such an approach is likely to discourage geese from foraging on vulnerable cereal fields and, additionally, should have positive effects on other farmland taxa (smaller birds, pollinating insects, weeds; Whittingham and Devereux 2008). The recommendations above may be easily incorporated into various agri-environment schemes; as such, farmers might benefit financially from both scenarios (Jensen et al. 2008; Crabtree et al. 2010).

Our recommendations are of major importance from the perspective of farmland management for geese in Poland and other countries in Eastern Europe that have recently joined the European Union (EU). These new members of the EU adopted the Common Agricultural Policy that was created on the basis of experience gathered mostly in the countries of Western Europe. There is a great concern that adoption of that common policy will lead to more intensive agriculture in the former traditionally managed farmlands of Eastern Europe (Báldi and Bátáry 2011; Tryjanowski et al. 2011). The policy favours large fields and has already led to lower habitat diversity in Western Europe and, consequently, to declines in biodiversity (e.g. Burel and Bradbury 1995; Gregory et al. 2004; Donald et al. 2006). However, from the perspective of migrating geese, these changes seem to be favourable, and one may expect that the size of the goose population will increase, which has the potential to also increase the conflicts between farmers and geese.

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Appendix

See Table 5.
Table 5 Pearson correlation coefficients (corrected for spatial autocorrelation) between continuous variables potentially influencing foraging site selection by geese

|   | (1) | (2) | (3) | (4) | (5) | (6) | (7) | (8) | (9) | (10) | (11) | (12) | (13) | (14) | (15) | (16) |
|---|-----|-----|-----|-----|-----|-----|-----|-----|-----|------|------|------|------|------|------|------|
| (1) FieldSize | –    | 0.029 | –0.087 | –0.371 | 0.264 | 0.290 | 0.143 | –0.023 | –0.215 | –0.100 | 0.141 | 0.009 | 0.388 | –0.192 | 0.424 | 0.032 |
|   |     | (0.902) | (0.664) | (0.061) | (0.219) | (0.222) | (0.508) | (0.925) | (0.336) | (0.614) | (0.475) | (0.954) | (0.071) | (0.372) | (0.034)* | (0.887) |
| (2) PPloughed | –    | 0.119 | –0.502 | 0.119 | –0.526 | –0.009 | –0.071 | 0.588 | 0.115 | –0.238 | 0.143 | –0.190 | 0.056 | 0.035 | 0.251 |     |
|   |     | (0.603) | (0.015)* | (0.601) | (0.002)* | (0.997) | (0.735) | (0.011)* | (0.567) | (0.249) | (0.464) | (0.423) | (0.772) | (0.869) | (0.221) |     |
| (3) PBee | –    | 0.189 | –0.358 | –0.368 | –0.068 | –0.014 | 0.051 | –0.165 | –0.036 | 0.061 | –0.268 | –0.229 | 0.186 | 0.195 |     |     |
|   |     | (0.341) | (0.401) | (0.137) | (0.744) | (0.428) | (0.836) | (0.408) | (0.876) | (0.774) | (0.255) | (0.221) | (0.404) | (0.349) |     |     |
| (4) PRape | –    | –0.369 | –0.097 | –0.133 | –0.250 | –0.031 | –0.317 | –0.003 | –0.243 | –0.261 | –0.034 | –0.124 | 0.094 |     |     |     |
|   |     | (0.069) | (0.659) | (0.538) | (0.228) | (0.892) | (0.128) | (0.987) | (0.221) | (0.210) | (0.872) | (0.585) | (0.667) |     |     |     |
| (5) PMaize | –    | 0.319 | 0.181 | –0.196 | –0.030 | 0.104 | –0.134 | 0.211 | 0.310 | 0.341 | –0.280 | –0.105 |     |     |     |     |
|   |     | (0.136) | (0.416) | (0.319) | (0.900) | (0.622) | (0.195) | (0.208) | (0.124) | (0.094) | (0.158) | (0.623) |     |     |     |     |
| (6) PWinterCere | –    | –0.075 | –0.006 | –0.729 | 0.196 | –0.027 | 0.072 | 0.370 | –0.025 | 0.280 | –0.206 |     |     |     |     |     |
|   |     | (0.713) | (0.980) | (0.001)* | (0.244) | (0.904) | (0.904) | (0.146) | (0.904) | (0.245) | (0.332) |     |     |     |     |     |
| (7) PPastures | –    | –0.131 | 0.301 | 0.060 | 0.257 | 0.153 | 0.171 | –0.173 | 0.169 | 0.120 |     |     |     |     |     |     |
|   |     | (0.332) | (0.148) | (0.783) | (0.199) | (0.484) | (0.304) | (0.432) | (0.402) | (0.597) |     |     |     |     |     |     |
| (8) PSetAside | –    | –0.034 | 0.258 | 0.297 | 0.079 | 0.012 | –0.152 | 0.126 | –0.082 |     |     |     |     |     |     |     |
|   |     | (0.856) | (0.212) | (0.201) | (0.709) | (0.964) | (0.475) | (0.560) | (0.712) |     |     |     |     |     |     |     |
| (9) HabitatDiver | –    | 0.009 | 0.022 | 0.146 | –0.148 | 0.046 | 0.058 | 0.306 |     |     |     |     |     |     |     |     |
|   |     | (0.954) | (0.012) | (0.411) | (0.616) | (0.786) | (0.825) | (0.098) |     |     |     |     |     |     |     |     |
| (10) DLocRoad | –    | –0.070 | 0.051 | 0.166 | 0.107 | –0.084 | 0.002 |     |     |     |     |     |     |     |     |     |
|   |     | (0.753) | (0.082) | (0.355) | (0.653) | (0.665) | (0.993) |     |     |     |     |     |     |     |     |     |
| (11) DRegRoad | –    | 0.071 | 0.120 | –0.052 | 0.134 | 0.002 |     |     |     |     |     |     |     |     |     |     |
|   |     | (0.749) | (0.605) | (0.806) | (0.455) | (0.993) |     |     |     |     |     |     |     |     |     |     |
| (12) DHumanS | –    | 0.148 | –0.085 | 0.327 | –0.171 |     |     |     |     |     |     |     |     |     |     |     |
|   |     | (0.425) | (0.715) | (0.082) | (0.422) |     |     |     |     |     |     |     |     |     |     |     |
| (13) DTown | –    | 0.197 | 0.149 | 0.288 |     |     |     |     |     |     |     |     |     |     |     |     |
|   |     | (0.549) | (0.549) | (0.090) |     |     |     |     |     |     |     |     |     |     |     |     |
| (14) DForest | –    | 0.171 | –0.294 |     |     |     |     |     |     |     |     |     |     |     |     |     |
|   |     | (0.391) | (0.175) |     |     |     |     |     |     |     |     |     |     |     |     |     |
| (15) DRiver | –    | 0.044 |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
|   |     | (0.843) |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| (16) DLake | –    |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |

* Significant correlations at $P < 0.05$. Significance levels are given in parenthesis For explanation of the variables: see Table 1
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