Sex-Related Differences in the Trade-Off between Foraging and Vigilance in a Granivorous Forager

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Abstract
The relationship between intake rate and food density can provide the foundation for models that predict the spatiotemporal distribution of organisms across a range of resource densities. The functional response, describing the relationship between resource density and intake rate, is often interpreted mechanistically as the relationships between times spend searching and handling. While several functional response models incorporate anti-predator vigilance (defined here as an interruption of feeding or some other activity to visually scan the environment, directed mainly towards detecting potential predators), the impacts of environmental factors influencing directly anti-predator vigilance remains unclear. We examined the combined effects of different scenarios of predation risk and food density on time allocation between foraging and anti-predator vigilance in a granivorous species. We experimentally exposed Skylarks to various cover heights and seed densities, and measured individual time budget and pecking and intake rates. Our results indicated that time devoted to different activities varied as a function of both seed density and cover height. Foraging time increased with seed density for all cover heights. Conversely, an increased cover height resulted in a decreased foraging time. Contrary to males, the decreased proportion of time spent foraging did not translate into a foraging disadvantage for females. When vegetation height was higher, females maintained similar pecking and intake rates compared to intermediate levels, while males consistently decreased their energy gain. This difference in anti-predator responses suggests a sexually mediated strategy in the food-safety trade-off: when resource density is high a females would adopt a camouflage strategy while an escape strategy would be adopted by males. In other words, males would leave risky-areas, whereas females would stay when resource density is high. Our results suggest that increased predation risk might generate sexually mediated behavioural responses that functional response models should perhaps better consider in the future.

Introduction
The relationship between intake rate and food density can provide the foundation for models that predict the spatiotemporal distribution of organisms across a range of resource densities [1]. It has been previously demonstrated that the relationship between food density and feeding rate (i.e. functional response) results from differential time allocation between competing activities, such as searching, handling, and anti-predator vigilance [2]. Because anti-predator behaviour may monopolize a large amount of the foraging time-budget [2,3,4], anti-predator vigilance is often regarded as a major factor limiting feeding rates, and ultimately affecting the functional response. Paradoxically, though much work has been devoted to investigating individuals’ responses to varying resource densities and predation risk independently [3,6], only a few studies have experimentally considered the combined effects of these factors on foraging time-budget and energy gain [7,8]. This could be important because food availability should impact patch selection by influencing intake rate, predator detection, and avoidance [9].

Changes in vigilant behaviour have been attributed to variation in predation risk, which is itself dependent on environmental factors including group-size for social or aggregative species [10,11,12,13], predator type and/or abundance [14], distance to safe refuges [15], or predator detectability [16,17]. Concerning this latter factor, many prey organisms can assess predation risk through cues signalling the presence of predators, either being auditory [18], chemosensory [19] or visual [20]. For prey species that rely on sight for detecting predators, vegetation structure may influence the perceived predation risk [11], and is therefore likely to affect foraging habitat selection [21] as well as anti-predator strategies. However, the effect of vegetation structure on vigilance is mixed, since reduced visibility caused by vegetation may increase vigilance [14,16,17,22], or increase camouflage and thus reduce vigilance [23]. Similarly, depending on species and their anti-predator strategies, patches with complex vegetation structure may be assimilated as dangerous by increase the perceived risk of predation [6,16,24] or protective by limiting the prey detectability by predators [25,26,27].

Besides environmental factors, individual phenotypes may also modulate the foraging time-budget [10]. For instance, individuals in poor body condition were shown to have lower anti-predator vigilance and contrastingly, prioritize energy gain [28]. Sex is also expected as one crucial source of inter-individual variation in the
optimization of foraging time-budget under varying food resource and predation risk. Indeed, predator’s preferences and foraging tactics [29,30] may result in sex-selective predation [31] with profound consequences on sex-specific anti-predator strategies. Several studies have found sex differences in the time allocated to anti-predator vigilance [32,33]. However, the influence of sex on individual response to varying food resources and predation risk has been poorly studied, especially in avian models. Among birds, ground-seed-eating species (such as many farmland birds, at least in winter) appear to be particularly exposed to predation, since they live in open habitats that increase the distance to safe refuges. In addition, although habitat use and diet in the breeding season have received considerable attention [34,35], habitat use and foraging behaviours during winter were less investigated [21,36,37,38].

In this study we use an experimental approach and examine foraging behaviour of a ground seed eating farmland bird, the Eurasian Skylark Alauda arvensis, on artificial stubble substrates. Stubble fields are important foraging habitats for many farmland and granivorous bird species wintering in Europe [36,39], but the effects of habitat structure on foraging and anti-predator strategies of these species remain unclear [6,7,37]. Although vigilance time is strongly influenced by predation risk [39], only few studies have directly investigated the combined effects of resource density and the perceived predation risk on functional response [7]. In order to investigate how the trade-off between foraging and vigilance is altered, we experimentally increased the visual obstruction from predators by exposing birds to three different vegetation heights. In addition to the predation risk, we manipulated resource availability on experimental patches. Theory predicts a differential response in terms of vigilance under contrasting resource availability, but several additional confounding factors may further balance this relationship such as competition or scrounging [40]. In some bird species, annual survival rates were found to be higher in males than in females, probably due to differential breeding costs [41] or sex-differences in risk-taking behaviour under foraging circumstances [42]. Hence males and females could behave differently with regard to their degree of exposure to predation and their differential survival prospects. If so, we predict that birds might differentially adapt their vigilance in response to the perceived predation risk according to resource density, but also their sex.

Methods

Ethics Statement

This work was performed with governmental authorizations from the Préfecture des Deux-Sèvres (Niort, France, no. 10.79-219). All experiments were carried out in compliance with French legal requirements. This study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Research. The protocol was approved by the Committee on the Ethics of Animal Experiments of the National Conservation Authority (permit no. 79349). Among all catching birds, animals showing any sign of sickness were removed before entering the experiment (N = 3). Bird captures were performed under permit from the National Hunting and Wildlife Agency to TP (no. 2009-014). All experiments were conform to the guidelines for the treatments and use of animals in behavioural research and teaching as published by the Association for the Study of Animal Behaviour (ASAB 2012).

Study species

The skylark is a common species of open farmland habitats throughout Europe [43] that suffers a sustained and often severe decline in relation to changes in agricultural practices since the last few decades [44]. Being strongly territorial and feeding on insects in spring and summer, the species shows a marked gregarious behaviour on migratory stop-over and wintering quarters where birds live in groups of various sizes. Outside the breeding season, the species mainly feeds on seeds gleaned on the ground and vegetative parts of plants. The species shows a slight sexual dimorphism, with males being on average larger than females [43].

Bird capture and maintenance

We caught skylarks along the Atlantic flyway during the post-nuptial migration, between October and November 2010. Upon capture, the birds were colour ringed, weighed (precision: ±0.1 g; males: 41±0.2 g; females: 31.7±0.22 g) and their tarsus and maximum wing chords were measured. Blood samples were taken, and their sex was obtained from molecular analysis (see [45] for details). Thereafter, birds were randomly assigned to groups of 10 to 12 individuals and acclimatized for two months in 4*3*2 m (L*W*H) outdoor aviaries at the Centre d’Études Biologiques de Chizé. Birds were fed ad libitum with a commercial seed mixture, grit, green material (i.e. oilseed rape) and tap water. Food was dispensed on a 2 m² synthetic green turf (height: 1 cm; density: 12 blades/cm²) to accustom individuals to the experimental set-up. To limit the time spent under captive conditions all birds were released into the wild in the middle of March, during the pre-nuptial migration.

General experimental procedures

All experiments were conducted from the 7th February to 4th March 2011. The night before each trial, all focal birds were weighed and deprived from food (between 12 and 16 hours) until the next morning. Trials were only performed from 8:00 to 12:00 am in order to avoid too long fasting periods. We did not conduct experiments on rainy or excessively windy days, as these extreme weather conditions would likely affect visibility and foraging behaviour. All trials were carried out in a separate outdoor aviary, showing identical proportions and layout to the housing aviaries.

We experimentally exposed birds to contrasting seed densities (3 levels) and vegetation heights (3 levels). Each bird was exposed to all of the 9 treatment combinations in a random order during the experiment. The experimental arena consisted of a 50×50×40 cm (H) wire cage (mesh size: 1×1 cm) enabling visual contact amongst birds. The cage was placed over an artificial green turf substrate surrounded by stubble. To mimic natural stubble habitat, we attached cereal straw to a brown polystyrene board (total width around the cage: 34 cm). The straw was arranged in rows 12 cm apart at a density of 110 straws/m, which was equivalent to the density measured in natural stubble fields around our laboratory (N=12; mean = 12±SE; 12.4±2.21 cm; 110±7 straws/m). To investigate the influence of the visual obstruction on foraging activities, we used three different stubble heights: 0 cm (no stubble), 15 cm (comparable to the height measured in 20 randomly selected patches in 5 stubble fields (mean ± SE = 15.2±1.87 cm; range 9.8–17.3) and finally 35 cm in height. To investigate the influence of seed density on foraging activities, we used millet seeds (Panicum miliaceum) owing to their homogeneity in both colour (white) and size (mass = 0.007±0.0003 g (mean ± SD); sample size: 300 seeds). Birds were exposed to three different seed densities (i.e. 100, 400 and 1600 seeds.m⁻²; counted by hand) randomly scattered on the artificial green turf substrate. All
these densities fell within the range of seed densities recorded in arable fields [46,47].

Among the birds captured during the fall migration, 10 females and 10 males were randomly selected and tested for all treatment combinations (i.e. 3 vegetation heights x3 seed densities) with no replicate. During winter, skylarks adopt particular aggregative strategies and restrict foraging behaviour in the absence of conspecifics [48]. Hence, one non-focal bird (different from the 20 tested birds) was randomly assigned to each trial, and individually kept in identical wire mesh cages at a distance of 1 m from the focal individual. To avoid synchrony in behaviour [49], non-focal birds were not provided with food during tests. Group composition and consequently group sex-ratio, including the focal individual and the con-specific (non-focal bird) thus varied randomly from test to test to avoid systematic association between partner birds and experimental treatments or conditions. There were between six and eight trials per day, but neither the focal nor the non-focal birds experienced more than one trial in any one day. Ten minutes before each trial, the birds were placed in their respective cages so that they became accustomed to the experimental set-up. Each trial lasted five minutes, starting when the first peck was recorded. Focal birds were video-recorded using a camcorder set 1 m above the ground and about 1 m from the focal bird.

Data collection and analysis

The behaviour of focal birds was analysed from videotapes using an event-recording program (EthoLog 2.2 [50]). All videos (n = 180) were analysed at 1X reading speed after testing on a subset of trials that this speed provided correlated results to those obtained by analyzing the videos frame by frame. (Foraging time; Pearson correlation coefficient: r = 0.781; p < 0.01, n = 90).

Both the pecking rate (expressed as the number of pecks during the 5 minutes trial) and the time devoted to foraging and vigilance (in seconds) were separately quantified. Using video recording, the posture of birds was analysed for each trial. Following Whittingham & Markland [51] and Powolny et al. [48], a bird was considered to be vigilant when its head was above the horizontal line made by its body, and not orientated towards the ground. Conversely, a bird was considered as foraging when its head was below the horizontal line and when it actively scanned the ground or pecked. In addition to these criteria, we used the position of the bill (oriented towards the ground or staying horizontally or towards the sky) to discriminate behaviours implied in foraging and vigilant activities. The handling time - i.e. the time needed to consume one food item - and vigilance could not be separated given that skylarks do not handle millet seeds [48]. However, although head down posture is associated with foraging behaviour, this position still may allow vigilance [52]. Thus, we added an opaque band of 5 cm high around the bottom of the cage leading to no possibility of looking out the cage when the bird’s posture was considered as foraging. So, given that vigilance and foraging represented the only two postures recorded in trials and because they were two mutually exclusive activities in our case study, only the time spent foraging was considered in analyses. Additionally, we quantified the time spent moving for a density of 100 seeds.m\(^{-2}\) and 400 seeds.m\(^{-2}\), which correspond to the seed density commonly measured on stubble fields [47], and 1600 seeds.m\(^{-2}\) representing our non limiting density. We paid particular attention to displacement under vigilant posture. Vigilant skylarks were considered as moving when they scanned their environment while walking with their head up.

The use of green turf allowed us to collect all remaining seeds. At the end of each trial, seeds remaining on the green turf were counted by hand by a single person without information about trials condition (sex, seed density or vegetation height). The corresponding values were used to calculate food intake rate (expressed as the number of seeds consumed during the 5 minutes of trial).

We used General Linear Mixed Models (GLMMs) to investigate the effects of seeds density, vegetation height and sex of the focal bird on time spent foraging, pecking and intake rates and time spent moving. Because focal individuals were tested for all treatment combinations, models were fitted with bird identity as a random factor and nested within sex. Analyses were separately performed for each dependent variable and followed a backward selection procedure starting from a full model that included all factors, their two-way interactions and a set of continuous covariates designed to control for the conditions in which the tests were conducted. These covariates were (Table 1): date, minimum nightly temperature preceding each trial, body mass and fasting duration (corresponding to the time since the sunrise). In addition, the effect of the sex of the conspecific (i.e. the non-focal bird) was also considered both as a main effect and in interaction with the sex of the focal bird. Starting by continuous covariates aiming to control for trial conditions, non-significant terms at P ≥ 0.05 were sequentially removed to achieve a minimal

<table>
<thead>
<tr>
<th>Variable</th>
<th>Factor/covariate</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date</td>
<td>co-variante</td>
<td>Julian date (1 for 1(^{st}) January)</td>
</tr>
<tr>
<td>Minimum temperature</td>
<td>co-variante</td>
<td>Minimum temperature the night before tested</td>
</tr>
<tr>
<td>Body mass</td>
<td>co-variante</td>
<td>Body mass the night before being tested</td>
</tr>
<tr>
<td>Focal bird sex</td>
<td>factor</td>
<td>Male or female</td>
</tr>
<tr>
<td>Conspecific sex</td>
<td>factor</td>
<td>Male or female</td>
</tr>
<tr>
<td>Food deprivation</td>
<td>co-variante</td>
<td>In minutes</td>
</tr>
<tr>
<td>Seed density</td>
<td>factor</td>
<td>100; 400 or 1600 seeds/m(^2)</td>
</tr>
<tr>
<td>Cover height</td>
<td>factor</td>
<td>0; 15 or 35 cm</td>
</tr>
<tr>
<td>Foraging time</td>
<td>factor</td>
<td>Focal bird searching for food (based on head position); in seconds</td>
</tr>
<tr>
<td>Intake rate</td>
<td>factor</td>
<td>Number of seeds consumed during test</td>
</tr>
<tr>
<td>Pecking rate</td>
<td>factor</td>
<td>Number of pecks during test</td>
</tr>
</tbody>
</table>

Table 1. Explanatory and response variables entered into models for the foraging time, intake and pecking rates studies.
adequate model [53]. Wherever an interaction term was left in a model, the two corresponding factors were also included as main effects. Post-hoc comparisons were performed using Tukey’s HSD tests. Both foraging and moving times were log-transformed while both intake and pecking rates were square-root transformed to ensure normality and homoscedasticity assumptions.

To test for habituation, GLMMs were performed on pecking and intake rates and foraging time using trial number as continuous covariate. Results did not show any evidence for habituation (pecking rate: $F = 1.633; df = 1, 173; p = 0.204$; intake rate: $F = 0.048; df = 1, 173; p = 0.827$; foraging time: $F = 0.548; df = 1, 173; p = 0.862$). All analyses were performed using SPSS 17.0 software. Means are expressed $\pm$ SE.

**Results**

We found no evidence that the foraging behaviour of birds varied according to date or the fasting duration since these covariates were systematically removed during model selection (Table 2). Conversely, body mass significantly affected pecking, but not intake rates and time of foraging or movement. An effect of the ambient temperature experienced the night that preceded the trials suggested varying individuals’ needs in relation to environmental conditions: the time spent in foraging, the number of pecks and ultimately intake rate, all increased as the minimum nightly temperature decreased (Table 2).

**Effects of seed density and cover height**

All dependent variables describing foraging behaviour and performance enhanced with increasing seed density (Table 2). Individuals foraging in a patch with the lowest seed density treatment (i.e. 100 seeds.m$^{-2}$) spent the shortest time foraging and had the lowest pecking and intake rates. Conversely, skylarks foraging in the richest patch (i.e. 1 600 seeds.m$^{-2}$) spent the longest time foraging and had both the highest pecking and intake rates.

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**Table 2.** Fixed effects explaining variation in time allocated to foraging, pecking and intake rates and the time spent in movement under vigilant posture.

<table>
<thead>
<tr>
<th>Dependent variable</th>
<th>Fixed effect</th>
<th>Estimates</th>
<th>Df/DfDen</th>
<th>F values</th>
<th>p</th>
<th>$\eta^2$</th>
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<td><strong>Foraging time</strong></td>
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<td>0.018</td>
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<td>conspecific sex</td>
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<td>seed density</td>
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<td>14.36</td>
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<td></td>
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<td></td>
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<tr>
<td></td>
<td>cover height</td>
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<td>2, 149</td>
<td>15.96</td>
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<td></td>
<td>150:0.21</td>
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<td></td>
<td>focal bird sex * cover height</td>
<td></td>
<td>2, 149</td>
<td>3.39</td>
<td>0.036</td>
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<tr>
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<td>2, 149</td>
<td>3.54</td>
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<td>8.83</td>
<td>0.003</td>
<td>0.591</td>
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<td>body mass</td>
<td>$-1.78$</td>
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<td>4.96</td>
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<td>2, 150</td>
<td>5.36</td>
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<td>151:1.51</td>
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<td></td>
<td>focal bird sex * cover height</td>
<td></td>
<td>2, 150</td>
<td>6.97</td>
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<td><strong>Intake rate</strong></td>
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<td>0.726</td>
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<td>20.48</td>
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<td></td>
<td>151:1.02</td>
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<tr>
<td><strong>Time in movement</strong></td>
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<td>cover height</td>
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<td>9.22</td>
<td>&lt;0.0001</td>
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<td>150:0.56</td>
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<td></td>
<td>focal bird sex * cover height</td>
<td></td>
<td>2, 156</td>
<td>7.08</td>
<td>0.001</td>
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</table>

A final model was backward-selected using a linear mixed effect model. The variation observed in the df’s comes probably from approximations that are made for the calculation of the random effect. In fact, we used 20 individuals, with 9 trials for each bird. This small difference is commonly observed when the number of individuals is not so high.

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rates (Fig. 1). Intermediate values were recorded at medium initial seed density (i.e. 400 seeds.m\(^{-2}\)). Skylarks foraging in a patch not surrounded by vegetation spent longer time foraging and had higher pecking and intake rates than those foraging in the highest cover height (i.e. 35 cm, Fig. 2). The interaction term between seed density and cover height was systematically removed during the selection process (Table 2) indicating that these two factors impacted foraging behaviour and performance in an additive way.

Sex-related differences

We found evidence that the effect of vegetation height on time spent foraging, pecking and intake rates differed significantly between sexes. More importantly, the interaction term between sex and cover height was retained in all final models (Table 2). The main component of this interaction was the greater sensitivity of males to the increase in vegetation height. With respect to the time budget, males reduced their time spent foraging at 35 cm in comparison to a cover height of 0 cm (post-hoc HSD Tukey: \(p = 0.001\), conversely to females which allocated a similar amount of time at 35 cm and 15 cm (post-hoc HSD Tukey: \(p = 0.13\); Fig. 2A) and between 0 and 35 cm (post-hoc HSD Tukey: \(p = 0.32\); Fig. 2A). A similar pattern was observed for pecking rate: males reduced their pecking rate when the vegetation height dropped from 15 to 35 cm (post-hoc HSD Tukey: \(p = 0.03\); Fig. 2B), whereas females preserved an almost identical pecking rate (post-hoc HSD Tukey: \(p = 0.97\); Fig. 2B). Interestingly, male behaviour changed markedly when the vegetation height increased, increasing the time spent in movement (while adopting a vigilant posture). Contrary to males, females showed a decrease in time spent in movement (Fig. 3; Table 2). When seed density was highest (i.e. 1 600 seeds.m\(^{-2}\)), males actually moved more than females at two of the three cover heights (0 cm: post-hoc HSD Tukey: \(p = 0.23\); 15 cm: post-hoc HSD Tukey: \(p = 0.05\); 35 cm: post-hoc HSD Tukey: \(p = 0.001\); Fig. 3). As a consequence, only males’ intake rate, but not females, was significantly reduced at 35 cm (post-hoc HSD Tukey: respectively: \(p = 0.003\) and \(p = 0.37\); Fig. 2C). While seed density and sex also interacted significantly in determining the allocation of time spent foraging (Table 2; Fig. 4), both sexes spent the shortest time foraging at low seed density and the longest time at the highest seed density (LSMeans (±SE), females: 23.99±1.20 vs. 33.11±1.20). While foraging time did not differ between sexes for the two highest seed densities (post-hoc HSD Tukey, 400 seed/m\(^{2}\): \(p = 0.03\); 1600 seed/m\(^{2}\): \(p = 0.04\)) males spent more time under foraging posture at 100 seed/m\(^{2}\) than females (post-hoc HSD Tukey: \(p = 0.03\); Fig. 4).

In addition, we found that the foraging behaviour of the focal birds depended on whether the conspecific non-focal bird was a male or a female (Table 2). For instance, pecking rate was significantly higher when conspecifics were males (LSMeans (±SE), males: 46.37±4.69, females: 32.41±6.07). Even more surprisingly, there was a significant interaction between the sex of focal and non-focal birds on the time devoted to foraging (Table 2): males reduced their time spent foraging when conspecifics were females (LSMeans (±SE), female non-focal bird: 19.05±1.23 vs. male non-focal bird: 37.15±1.17) while females were insensitive to the sex of conspecifics (LSMeans (±SE), female non-focal bird: 33.88±1.23

Figure 1. Effect of seed density on foraging time (A), pecking rate (B) and intake rate (C). Untransformed data are presented in the figure. doi:10.1371/journal.pone.0101598.g001

Figure 2. Cover height effect on foraging time (A), pecking rate (B) and intake rate (C). Black histograms represent males and white histograms represent females. Bar labelled with different letters are significantly different (\(p<0.005\)). doi:10.1371/journal.pone.0101598.g002
male non-focal bird: 30.2 ± 1.17). Conversely, the sex of conspecifics had no significant effect on intake rate.

**Discussion**

**Impacts of the vegetation height**

In this study, we experimentally examined how resource availability and perceived predation risk affected time budget, pecking rate and ultimately seed intake in skylarks. As expected, our results indicated that time devoted to foraging varied both as a function of seed density and vegetation height. An increase in vegetation height consistently affected time-budget in skylark and resulted in a decrease of foraging time, probably through increasing vigilance. The reduced feeding time for the benefit of vigilance at the maximum vegetation height (i.e. 35 cm) indicated that visual obstruction would be perceived as a high predation risk in this species, especially in males, despite its obvious protective nature. Many studies have reported similar changes in time budget in obstructive habitat [10,16,17,22], attesting that perceived predation risk was increased in our obstructive treatment. This result agrees with Verdolin [54], who suggested that habitat structure, particularly vegetation height, may significantly increase an animal’s perceived predation risk, and strongly affects foraging behaviour. Natural vegetation and sometimes foraging substrate combine both protective and obstructive properties [26,48], depending on the predator type and anti-predator strategies adopted by a species [26]. Increased vigilance under high cover height can also be explained by the additional utility of scanning when the view is obstructed. This anti-predator response can be related to the group-living strategy adopted by this species outside the breeding season. Skylarks are highly gregarious both during winter and migration, hence vigilance time in skylarks is unsurprisingly related to group-size [48]. It is worth noting that our current results are opposed to the predictions that could be made from the studies of Butler et al. [21] and Whittingham et al. [37]. In these two studies, authors have shown that the abundance of skylarks in winter was higher on control and untouched stubbles (12–17 cm high) than on experimentally left reduced stubbles (5–6 cm high). This treatment strongly impacted vegetation height (5–6 cm vs. 12–17 cm for topped and control plots, respectively) but had no effect on the soil seed density, leading to the conclusion that skylarks would associate higher predation risk with short stubbles [21,37]. However, skylarks do not exclusively search for seeds on the soil surface but also peck at weed leaves and seed heads [43], which are likely to have been sharply reduced by topping. Hence, the hypothesis that topping had no effect on the potential energetic gain associated with short stubbles cannot be fully excluded as this treatment might have caused a reduction in food resources for the species.

**Impacts of the seed density**

We found a positive relationship between foraging time and seed density independently of experimental level of visual obstruction, indicating that for each tested cover height, birds foraged more at high resource density. However, the benefits linked to the high seed density were balanced by the vegetation height, with a similar time spent foraging at a density of

![Figure 3. Average time (seconds) spent moving under vigilant posture. Black histograms represent males and white histograms represent females. Only results obtained for a seed density of 1600 seed/m² were representing. Vertical bars show standard errors. Bars labelled with different letters are significantly different (p < 0.005). doi:10.1371/journal.pone.0101598.g003](image1)

![Figure 4. Effect of seed density on foraging time. Black histograms represent males and white histograms represent females. Vertical bars show standard errors. Bars labelled with different letters are significantly different (p < 0.005). doi:10.1371/journal.pone.0101598.g004](image2)
Sex differences

In skylarks, male and female foraging time differed at a cover height of 35 cm (the more obstructive situations). While female skylarks maintained equal foraging time between the three cover heights, males exhibited a marked decrease in foraging time between the intermediate and the highest heights (15 and 35 cm, respectively). Several studies have shown that sex is an important determinant of individual foraging and vigilance levels [57,58], reflecting differences in vulnerability to predation [59] and in the social factors that motivate scanning behaviour [60]. We observed intake differences between sexes in response to increased visual obstruction. In males, the decreased foraging time with vegetation height caused a reduction in the number of seeds consumed whereas females maintained their intake rate at the highest level of visual obstruction. In this later condition, our results also revealed an important contrast in the modality of vigilance adopted by each sex. While females tended to be immobile during vigilance sequences - suggesting a hiding or camouflage strategy-, males increased their time spent in movement – suggesting an escape strategy. Overall, these differences suggest a sex-related difference in the food-safety trade-off. Thus, males seem to prefer leave a risky-area, whereas females stay when resource density is high. As predation risk may vary among habitats, animals may not necessarily select habitats based solely on potential energetic gain. Consequently, individuals are likely to accept lower energetic benefits in order to forage in safer habitats [61]. Several field observational studies have also shown food-safety trade-offs, and differences between individual classes were observed [62]. For instance, Heithaus and Dill [63] suggested that juvenile male dolphins (Tursiops aduncus) are more willing to accept higher predation risk to achieve higher energy intake rates. A similar age class difference in risk-taking behaviour while foraging has been found in the redshank (Tringa totanus) where juveniles attempt to maximize energy intake by foraging in high-risk areas [64]. In addition to age-classes, sex was also identified as a cause of behavioural differences in another bird species, the Western Sandpipers (Calidris mauri) where females were overrepresented in more dangerous habitats [65]. Our results suggest that female skylarks may also gain fitness benefits from risk-taking. By foraging in the energetically profitable habitats, females may increase their body-condition and their long-term survival while males tend to maximize escape and consequently immediate survival. Moreover, the energetic needs of sexes may be different. Females need more energy to be prepared for the egg laying hence they risk more to get the necessary energy than males do. To test for effective food-safety trade-off in skylarks, further experiments involving patch choices between males and females would be needed. As example, following the set-up proposed by Butler et al. [7] adapted to test males versus females skylarks, we may hypothesize that females should spend more time in a risky-environment than males if the energetic return can be important. One possible explanation of this pattern is that while we observed no difference in body condition between sexes, energetic needs to prepare the next pre-nuptial migration and breeding might differ between sexes.

In addition to resource availability and vegetation height, two other factors influenced foraging time allocation in skylarks. First, the minimum temperature during the night before experiments affected both the time spent under foraging activities and intake rate. Low temperature influences energy expenditure, through increased demand for thermoregulation, resulting in an increase in food intake or a higher proportion of time spent in foraging. This result is consistent with studies demonstrating the relationships between the maintenance of body mass and the increased food intake in response to thermal stress [66]. Second, the sex of the conspecific individual significantly affected focal bird foraging time and pecking rate. This result agrees with studies suggesting that vigilance time may be influenced by the sexual composition of the group for aggregative species [67]. Particularly, in our study, when conspecific were males, the pecking rate of focal bird increased but there was no effect on intake rate. This surprising result may highlight a decrease in foraging success (measured as intake rate/pecking rate) especially when focal and non-focal birds are males suggesting possible foraging competition by interference [55]. Another possible explanation for this intriguing result is the potential competition for mating, even if our experiment were done before the pre-nuptial migratory period.

In summary, our study showed that lower vegetation and high seed density enhanced foraging rate and food intake in our study model. It has been demonstrated [68] that perceived predation risk can alter habitat choice, which has been shown to influence survival and community structure [69]. Thus, habitat structure appears to be a major component of population dynamics, especially for community living in homogenous landscapes, such as farmland birds. Our results highlight that phenotypic traits such as sex, can generate variations in behavioural responses outside the foraging season, independently of body condition. A mechanistic understanding of such intraspecific variations in behaviour is critical to develop adequate functional models and may be relevant for management and conservation issue as well.

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Author Contributions

Conceived and designed the experiments: TP VB CE. Performed the experiments: TP AA. Analyzed the data: TP. Contributed reagents/materials/analysis tools: TP VB CE. Wrote the paper: TP VB CE.

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