



HAL
open science

Crop phenology reshapes the food-safety landscape for roe deer in an agroecosystem

Noa Rigoudy, Simon Chamaillé-jammes, A J Mark Hewison, Arnaud Bonnet, Yannick Chaval, Bruno Lourtet, Joël Merlet, Nicolas Morellet

► **To cite this version:**

Noa Rigoudy, Simon Chamaillé-jammes, A J Mark Hewison, Arnaud Bonnet, Yannick Chaval, et al.. Crop phenology reshapes the food-safety landscape for roe deer in an agroecosystem. *Journal of Applied Ecology*, 2024, 61 (3), pp.564-574. 10.1111/1365-2664.14581 . hal-04664484

HAL Id: hal-04664484

<https://hal.inrae.fr/hal-04664484v1>

Submitted on 30 Jul 2024

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution 4.0 International License

RESEARCH ARTICLE

Crop phenology reshapes the food-safety landscape for roe deer in an agroecosystem

Noa Rigoudy^{1,2}  | Simon Chamaille-Jammes¹  | A. J. Mark Hewison² | Arnaud Bonnet² | Yannick Chaval² | Bruno Lourtet² | Joël Merlet² | Nicolas Morellet² 

¹CEFE, Univ Montpellier, CNRS, EPHE, IRD, Montpellier, France

²Université de Toulouse, INRAE, CEFS, F-31326 Castanet-Tolosan, France and LTSER ZA PYRénées GARonne, Auzeville-Tolosane, France

Correspondence

Noa Rigoudy
Email: noa.rigoudy@cefe.cnrs.fr

Funding information

'CARRY-MOVE' Agence Nationale de la Recherche, Grant/Award Number: ANR-23-CE02-0008

Handling Editor: Mahdiah Tourani

Abstract

1. Understanding the behavioural adjustments of wildlife in anthropized landscapes is key for promoting sustainable human-wildlife coexistence. Little is known, however, about how synanthropic species navigate spatio-temporal variation in the availability of food and cover that are shaped by human practices such as agriculture.
2. Animal habitat use is predominantly driven by spatial and temporal variations in food and cover, as individuals respond to fluctuations in the trade-off between food acquisition and risk avoidance. In agroecosystems, the availability of high-quality forage and cover is dependent on agricultural practices (e.g., harvesting) and crop phenology, providing an ideal opportunity to evaluate how wildlife adjust their behaviour in a heterogeneous human-dominated landscape.
3. We investigated the influence of crop phenology on the behaviour of European roe deer (*Capreolus capreolus*) to infer the functional roles of crop types in the food-cover landscape. We analysed the habitat selection and activity patterns of 105 GPS-collared adult female roe deer using a unique data set combining field-specific land cover data, region-specific estimates of crop phenology and weekly harvesting data for three common crops in a French agroecosystem.
4. We found very distinct habitat selection and activity patterns according to crop type, phenological stage and time of day. Wheat and artificial meadows were strongly selected at night-time during the early and post-harvest stages only, when roe deer were highly active, suggestive of feeding activity. On the contrary, roe deer strongly selected maize during the day when it was high enough to provide cover, when they were less active, indicating that it was primarily used for refuge. These patterns depended on the availability of more 'natural' cover, suggesting that mature maize may substitute for 'natural' cover when the latter is scarce.
5. *Synthesis and application:* Our work highlights the importance of behavioural plasticity and habitat complementation in the persistence of this synanthropic species in agroecosystems. This behavioural adjustment may buffer the consequences

This is an open access article under the terms of the [Creative Commons Attribution](https://creativecommons.org/licenses/by/4.0/) License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

© 2024 The Authors. *Journal of Applied Ecology* published by John Wiley & Sons Ltd on behalf of British Ecological Society.

of the reduction in natural habitats that accompanies intensification of agricultural production and has implications for understanding how agricultural practices shape the food-safety trade-off of wildlife living in these highly modified landscapes.

KEYWORDS

agricultural landscape, behavioural plasticity, cover, forage, habitat use, phenology, ungulate

1 | INTRODUCTION

Plant phenology is a key driver of the distribution of resources across landscapes so that animals must move accordingly, trying to track these changes in space and time (Armstrong et al., 2016). Indeed, plant phenology drives predictable changes in forage quantity and quality during plant growth and senescence. In response, many herbivores modify their use of space at temporal scales ranging from weeks to seasons (Abrahms et al., 2021). For example, sequential green-up across landscapes promotes the migration of large herbivore populations that surf the green wave of high-quality forage during spring (Aikens et al., 2017). At a finer spatial scale, plant phenology may also influence animal space use by providing pulses of high-quality forage. For example, mast fruiting by trees and grasses is known to strongly affect the habitat use of birds, small mammals and European ungulates (Bogdziewicz et al., 2016).

Human activities have been shown to alter patterns of resource distribution and phenology for wild animals across many ecosystems (Plummer et al., 2015). This is especially true in human-dominated landscapes, such as agroecosystems, where resource availability and quality are driven by both natural and anthropogenic factors (Barker et al., 2019). Agroecosystems are, by definition, spatially heterogeneous as they combine both semi-natural habitats and agricultural crops (Fahrig et al., 2011). Agricultural landscapes are also characterized by strong temporal variation in resource distribution for wildlife. The quantity, quality and availability of resources for a given species varies across crop types and depends both on phenological changes in plant development (Doorenbos & Kassam, 1979) and agricultural activities (e.g., harvesting, mowing or irrigation). Crops are known to attract wildlife as they generally have a high sugar and fat content, with low levels of toxins and fibre (Abbas et al., 2011), and are concentrated in large dense patches that require less search time (Simon & Fortin, 2020). Phenological changes in crop nutrient content are, thus, key in determining the crop foraging behaviour of herbivores. For example, African elephants (*Loxodonta africana*) increased their selection for crops during the brown-down stages, resulting in diets with higher levels of crude protein and energy (Branco et al., 2019) and higher body mass (Chiyo et al., 2011) compared with individuals that fed solely on natural forage.

Foraging strategies, and more generally space use, are, however, strongly constrained by risk perception, as individuals must navigate between habitats that rarely provide both high-quality forage and sufficient cover simultaneously (Brown, 1992). As a result, the use

of a given crop type by wildlife should also depend on the amount of cover it provides at a particular phenological stage. Many ungulates are known to leave more 'natural' habitats that provide cover to use agricultural areas at night, probably because crop fields are perceived as riskier during daytime due to the presence of humans and lack of vegetation cover (Bonnot et al., 2013; Branco et al., 2019; Mysterud & Østbye, 1999). In agricultural landscapes, the spatial distribution of food and cover thus depends on the type and phenological stage of each crop in relation to the species' feeding and risk management strategy, defining a dynamic food-fear landscape for wildlife. This aspect has been largely neglected in the past, with agricultural land often being considered as a single static 'open' habitat, potentially providing forage, but little or no cover. Recent studies have shed some light on the effects of spatio-temporal variations in crop availability and quality on animal space use at fine temporal scales (Branco et al., 2019; Paolini et al., 2018; Wilber et al., 2020), but none formally considered the role of crops as potential cover, nor how this balance between food and cover availability varies across time. There is an increasingly urgent need to understand how agricultural practices shape the trade-off between resource acquisition and exposure to risk for wildlife as future predictions concerning land conversion and severe climatic events suggest that the scope for human-wildlife coexistence will be reduced (Abrahms et al., 2021).

Here, we investigated the influence of human-driven changes in crop phenology on European roe deer (*Capreolus capreolus*) behaviour to identify the functional role of various crop types across phenological stages in the food-cover landscape. To do so, we used a unique data set combining detailed annual land cover data, weekly estimates of crop phenology for three crop types and tracking data from 105 GPS-monitored adult female roe deer living in an agroecosystem. This enabled us to map the availability of cover across the landscape at a fine spatio-temporal scale and disentangle the influence of plant phenology and harvesting events on crop use. The roe deer is a particularly appropriate species for this investigation as it thrives in agricultural landscapes (Hewison et al., 2009) due to its high ecological and behavioural plasticity (Andersen et al., 1998). Little is known, however, about how roe deer navigate spatio-temporal variations in the availability of food and cover that are shaped by human practices in agroecosystems, nor how changes in these practices might modify the species' food-safety trade-off. We aimed to: (1) determine the influence of crop type, phenology and time of day (night/day) on

crop selection to understand how roe deer adjust their behaviour to the spatio-temporal cycles of an agroecosystem, (2) contrast use and associated activity levels between day and night for the various crop types to identify potential functional roles in terms of food or cover across phenological stages (3) determine how the availability of crops or woodland/hedgerows, that provide alternative, more 'natural', habitats, modifies the selection of a given crop type, providing insight on how landscape composition determines the reliance of roe deer on cultivated land for food or cover subsidies. We expected that selection for crop types would change with crop phenology and predicted that selection for a crop type at a given stage would be stronger when the crop represented a good food resource or provided sufficient cover. In addition, we expected that habitats used as cover would be used mainly during the day, with lower activity levels if they were used for resting, and that those used for foraging would be used mainly during the night with high activity levels. Finally, we expected that tall maize could substitute for the cover provided by woodland and hedgerows during the day, with higher selection for mature maize for individuals with more limited access to woodland and hedgerows.

2 | MATERIALS AND METHODS

2.1 | Study site and population

The study was conducted on a roe deer population located in a 19,000-ha rural region in southern France (N43°17', E0°53'). The area is a heterogeneous agricultural landscape consisting of woodlands (18.9%, two large forests and fragmented woodland patches), hedgerows (3.5%), natural meadows (28.7%) and arable land (38.9%). Arable crops include artificial meadows (11.0%), cereals (wheat 13.2%, maize 2.9%, barley 2.0%, sorghum 1.2%) and oilseed (sunflower 6.0%, soybean 1.7%, rapeseed 0.9%). A network of roads link isolated houses, farms and small villages. Drive hunting represents the most lethal human activity for roe deer in the area, occurring regularly between mid-September to February. Trophy hunting occurs from June to September, likely maintaining a high perception of risk for roe deer during most of the year. The density for this roe deer population was estimated around eight individuals per 100ha using capture-mark-recapture re-sighting (Hewison et al., 2021) and the annual harvest was around 1.4/100ha in 2017. Other ungulates, notably wild boar (*Sus scrofa*) and red deer (*Cervus elaphus*), are also present in the study area, but at low densities.

2.2 | Landscape composition

Areas of homogeneous habitat patches were manually digitized as polygons and were revised every year (in ArcView GIS 3.3, Esri, Redlands, CA, U.S.A.) based on aerial photographs of the study site (from the IGN's BD Ortho, <https://geoservices.ign.fr/bdortho>) and yearly field observations. We considered six land cover types in

our analyses (Figure S1, details in Appendix S1.1 in Supplementary Information): woodland, hedgerows, maize, *Zea mays*, wheat, *Triticum* sp., artificial meadows (including lucerne, *Medicago sativa*, ray-grass, *Lolium perenne*, clover, *Trifolium* sp.), other (all other land cover types, mainly composed of natural meadows). We focussed on the three crop types, maize, wheat and artificial meadows, for which we had the widest coverage (present in most individual home ranges) and which are known to provide potential cover and/or forage for roe deer (Abbas et al., 2011; Putman, 1986). Ray grass during the growth phase, maize kernels and wheat grains are all known to have high energy content and to be highly digestible (Abbas et al., 2011, 2013; Wilber et al., 2020); each makes up a substantial part of the roe deer diet in spring, summer and autumn, respectively (Abbas et al., 2011). Wild grasses growing within wheat fields could also provide nitrogen-rich resources during winter and spring (Abbas et al., 2013). We were thus able to compare crops with contrasting temporal patterns of availability (e.g., maize in spring-autumn, wheat in winter-summer, see Figure S2) and different phenological characteristics (e.g., mature maize offers cover, emerging wheat offers palatable food).

2.3 | Roe deer biologging data

We measured roe deer habitat use using GPS data collected on 105 adult female roe deer from 2011 to 2021 with a 6 h fix rate (Lotek and Vectronic collars, see details in Appendix S1.2). As we wished to compare diurnal versus nocturnal habitat use, we only considered locations recorded at 12:00 (day) and at 00:00 (night). This also ensured that we focussed mainly on foraging and/or resting behaviour as roe deer are known to concentrate their inter-patch movements during crepuscular phases (Benoit, Bonnot, et al., 2023). All GPS collars integrated an activity sensor which indexed intensity of movement on the x (forward backward) and y (sideways) axes (details in Appendix S1.2). We measured total activity as the sum of activity values for both axes (range 0–510). We then extracted total activity values for each GPS location as the mean of activity values taken within <5 min of the GPS location. All capture and marking procedures on roe deer were done in accordance with local and European animal welfare laws (prefectural order from the Toulouse Administrative Authority to capture and monitor wild roe deer and agreement no. A31113001 approved by the Departmental Authority of Population Protection).

2.4 | Estimating crop phenological stages and monitoring agricultural practices

Information on annual crop phenological stages (from sowing to harvest) is scarce and often recorded in crop-specific monitoring programs for parasite and disease surveillance (e.g., Vigicultures®). For this study, we combined data from different monitoring programs to produce weekly estimates of phenological stage for each crop type

and year (Table S1, see Appendix S1.3 for a detailed description). We also assessed median harvest date for each year and crop type in our study area using annual field observations. We then combined the estimated phenological stages and median harvest dates to create weekly estimates of emergence, growth, maturity, harvest and post-harvest periods for each crop type and year.

2.5 | Estimating roe deer crop selection across phenological stages

We first tested the influence of phenology and time of day on the selection of each crop type. We extracted habitat type from the year-specific land cover map and attributed a phenological stage to each location by cross-referencing the date of the GPS location with the weekly crop phenology estimates. We considered that habitat availability was constant for a given individual over the year (all polygons contained a single land cover type in a given year). Comparing the probability of crop use among phenological stages was, thus, equivalent to estimating relative selection among phenological stages, as crop availability was constant across stages. For these analyses, we only considered individuals that had at least one location in the focal crop during their monitoring period as we wished to compare patterns of selection among phenological stages for a given crop type.

For each crop type, we fitted a binomial generalized linear mixed model (GLMM) with a logit link function, considering use as the response variable (focal crop type vs. other habitats, coded as 1 vs. 0 for each location) and phenological stage and time of day as fixed effects. As we were interested in the interplay between crop phenology and time of day, we included this interaction in our models. We set individual identity as a random intercept to account for repeated measures.

2.6 | Identifying potential functional roles for crops

We wished to investigate whether different crop types played different functional roles depending on their phenological stage. Roe deer have distinct nycthemeral activity patterns, with low activity levels when resting in refuge habitat during the day, but higher activity levels when they forage, mostly during the night (Bonnot et al., 2020; Pagon et al., 2013). In the absence of direct behavioural observations, we used this nycthemeral pattern of activity to infer potential functional roles for each crop type across phenological stages. For these analyses, we considered three phenological stages that were common to all crop types and represented contrasting characteristics in terms of cover and food resources: emergence (no cover, potential forage), maturity (cover, little forage for cereals, potential forage in certain meadows) and post-harvest (no cover, potential forage in wheat and maize stubble). We contrasted patterns of use during night and day for each crop type and phenological stage by performing post hoc contrast analyses on our previous

model predictions for crop use and adjusting for multiple comparisons using the Dunnett adjustment method (Dunnett, 1964).

We classified activity sensor data into active versus inactive behaviour by using the thresholds of activity level specified by Bonnot et al. (2020) who worked on the same dataset. Since we considered phenological stages that were common to all crop types, we fitted a single binomial GLMM with a logit link function to analyse variation in activity in relation to crop type, phenological stage (emergence/maturity/post-harvest) and time of day (night/day), including a random intercept for individual identity to control for repeated observations. We included the three-way interaction between crop type, phenological stage and time of day as we wished to test whether activity differed between night and day in relation to crop height ('tall', for mature maize and wheat vs. 'low' for all other crops and stages), indicating different functional roles for different crop types and phenological stages. We accounted for seasonal variation in activity levels linked to female reproductive phenology (Benoit, Morellet, et al., 2023) by including a circular spline on the Julian date. We also included collar type as a fixed effect as using activity level thresholds that were specific to each collar brand did not completely account for variation due to collar type (LRT test, $dAIC = 20$, $Chisq = 21.95$, $p < 0.001$).

2.7 | Testing the influence of landscape composition on crop selection across phenological stages

Finally, we investigated whether crop selection depended on its availability and/or the availability of 'natural' cover, that is, woodlands and hedgerows, in the local landscape. Indeed, woodlands and hedgerows are known to provide key 'cover' habitats for roe deer in agricultural landscapes (Morellet et al., 2011; Mysterud & Østbye, 1999; Padié et al., 2015). We tested for functional responses in crop use within the home-range (third order selection, Johnson, 1980). We defined availability as the proportion of a given habitat type within each individual's annual home range, using the fixed kernel method at 95% with an ad hoc approach to select the optimal smoothing parameter h for each home range estimate (Worton, 1989). We defined crop use as the proportion of locations in a given crop type within an individual's home range. Following Mysterud and Ims (1998) and previous work on this system (Morellet et al., 2011), we regressed proportional use against proportional availability for each crop type, logit-transforming both response and predictor variables to maintain linearity on the logit scale. For each crop type, we fitted a binomial GLMM with a logit link function, including phenological stage (emergence/maturity/post-harvest), time of day, crop availability and 'natural' cover availability as fixed effects, with a random intercept for individual identity to control for repeated observations. We included the four-way interaction to test whether the degree of use of a given crop as forage (mainly night) or cover (mainly day) during certain phenological stages was influenced by its availability and/

or the availability of woodlands and hedgerows within the home range. We framed our interpretation in terms of habitat selection, considering that a given habitat was selected or avoided if use was greater or less than its availability, respectively.

Analyses were performed in R version 4.0.2 (R Core Team, 2020); the R package 'amt' (Signer et al., 2019) was used for computing home ranges, 'glmmTMB' (Brooks et al., 2017) and 'gamm4' (Wood & Scheipl, 2020) were used to fit GLMM models and 'ggeffects' (Lüdtke, 2018) was used for model predictions.

3 | RESULTS

3.1 | Crop selection in relation to phenological stage

Patterns of selection by adult female roe deer varied markedly among crop types (Figure 1) and were influenced by both time of day and phenological stage (Table S2). For maize, selection increased during the late phenological stages (Figure 1a, flowering, maturity and post-harvest) and was significantly higher during the day than during the night when it was tall enough to provide cover (flowering and maturity stages, Table S4, contrast day/night: odds ratio >2.99, t ratio >7.42, p < 0.001). For wheat, selection was highest during the

night for the emergence and post-harvest stages (Table S4, day/night: both odds ratios <0.24, t ratios < -9.77, p < 0.001), as well as during both day and night for the mature stage (Figure 1b). For artificial meadows, selection was higher during the night across all phenological stages (Table S4, day/night: all odds ratios <0.45, t ratios < -5.50, p < 0.001) and highest during the early stages (Figure 1c).

3.2 | Potential functional roles of crops

There was a strong day/night contrast in patterns of activity (Tables S3 and S5) and crop use across crop types and phenological stages (Figure 2). Adult female roe deer exhibited lower activity when using mature maize during the day (Table S5, contrast activity day/night: odds ratio = 0.43, t ratio = -3.03, p = 0.002). On the contrary, they exhibited a somewhat higher level of use at night (Table S4, day/night: odds ratio = 0.73, t ratios = -1.78, p = 0.074) and a higher activity level when using maize during post-harvest (odds ratio = 0.26, t ratio = -3.81, p < 0.001). Similarly, adult female roe deer were highly active when using wheat during the emergence and post-harvest stages at night (Figure 2b), but significantly more so only during post-harvest (Table S5, odds ratio = 0.35, t ratio = -3.35, p < 0.001), and across all stages when using artificial meadows at night (Figure 2c all lower CI bounds >0.5 at night).

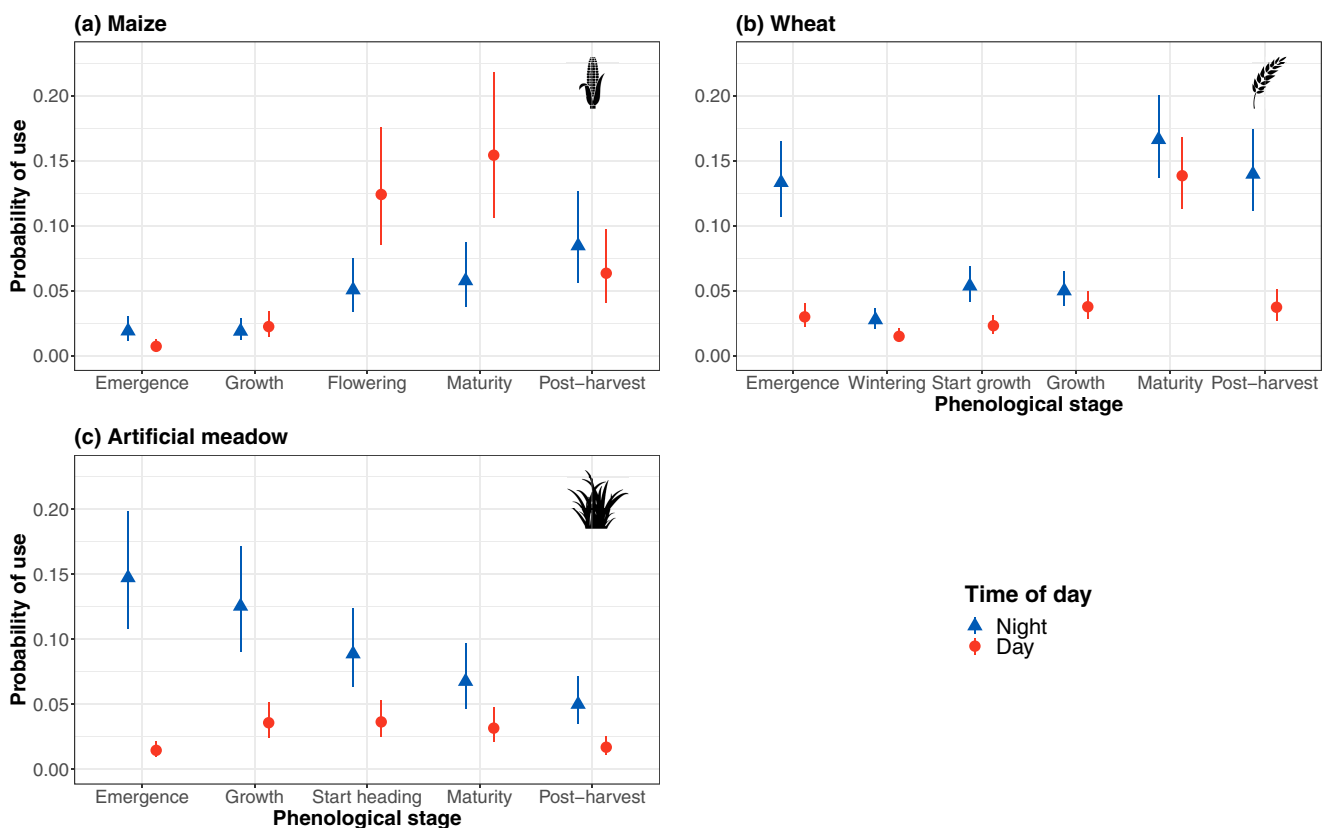


FIGURE 1 Predicted probability of crop use according to phenological stage and time of day for three crop types: (a) maize, (b) wheat, (c) artificial meadow based on, respectively, (a) 54, (b) 90 and (c) 98 individuals. Points are predicted marginal means estimated using binomial GLMMs with associated 95% confidence intervals (mean \pm CI).

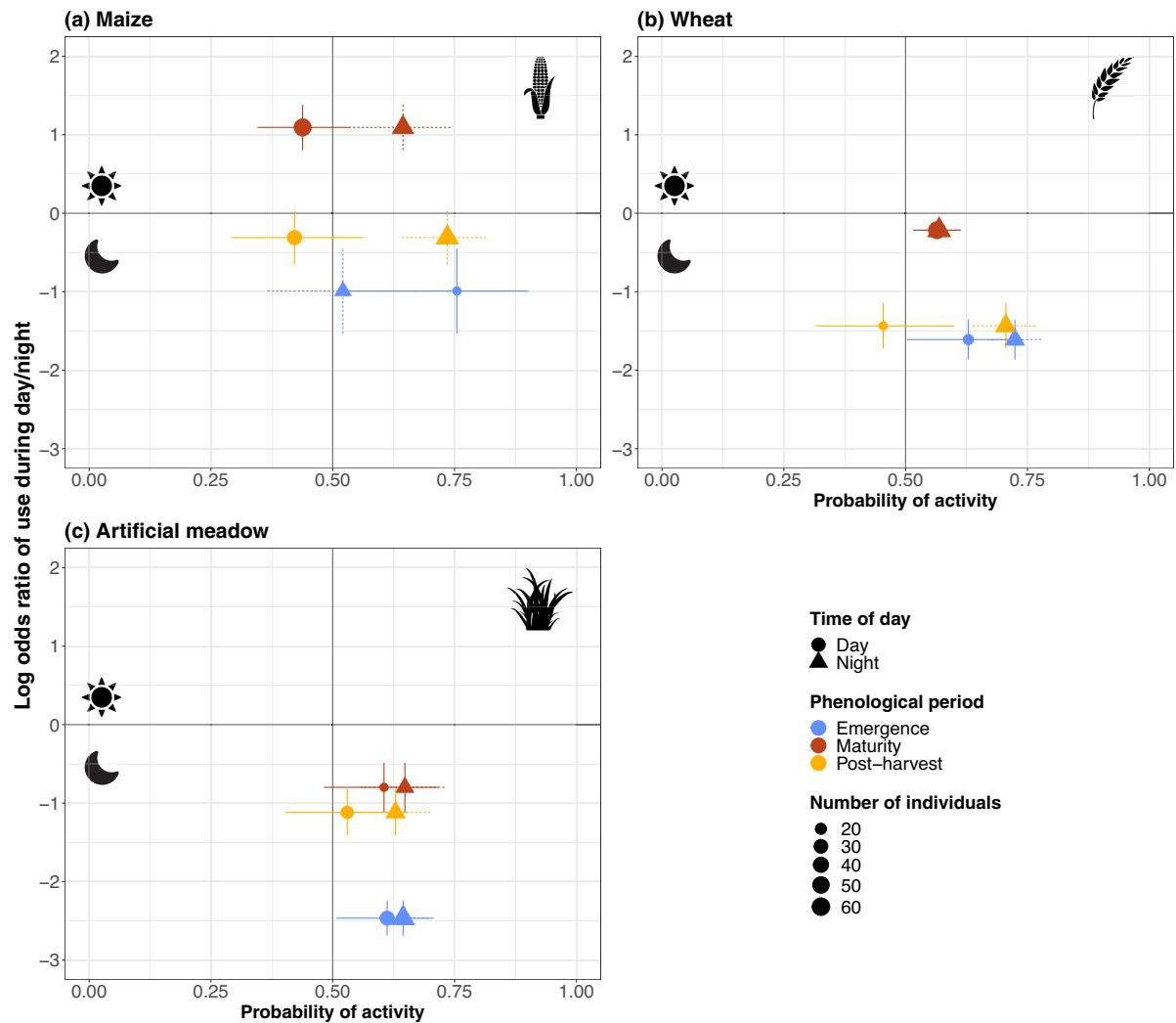


FIGURE 2 Log odds ratios of use during the day versus night plotted against predicted probability of activity for the emergence, maturity and post-harvest stages of three crop types: (a) maize, (b) wheat and (c) artificial meadow. Odds ratios were calculated using contrast analyses on predictions from models of crop use; activity level was estimated using binomial GLMMs. Points represent predicted marginal means and log odds ratios with associated 95% confidence intervals. Vertical and horizontal grey lines represent 50% activity level and an odds ratio of 1 for day to night use.

3.3 | Influence of landscape composition on crop selection across phenological stages

Availability of the crop itself and the availability of 'natural' cover (woodland and hedgerows) influenced crop selection only for certain crop types, phenological stages and times of day. Adult female roe deer strongly avoided maize irrespective of its availability or of the availability of 'natural' cover during emergence (use < availability, Figure 3a). When maize availability was higher than a certain threshold (15%–20%) and 'natural' cover more abundant ($\geq 24\%$), adult female roe deer selected maize at night during the post-harvest stage (use > availability, Figure 3c). In contrast, mature maize was selected during daytime only if 'natural' cover was less available ($\leq 14\%$, Figure 3b). In both cases, selection for maize increased with increasing availability, but this relationship depended on the availability of 'natural' cover. On the contrary, selection patterns for wheat and

artificial meadows did not depend on 'natural' cover availability (Figures S3 and S4). Wheat was strongly avoided during daytime during the emergence and post-harvest stages, and was used proportionately to its availability at night, irrespective of its availability (Figure S3). Artificial meadows were avoided during daytime across all phenological stages and availabilities and were used proportionately to their availability at night during maturity (Figure S4). Adult female roe deer strongly selected artificial meadows during the night and did so progressively more with increasing availability during the emergence stage (Figure S4).

4 | DISCUSSION

There is a growing need to better understand how human-driven resource dynamics influence wildlife behaviour in shared areas to

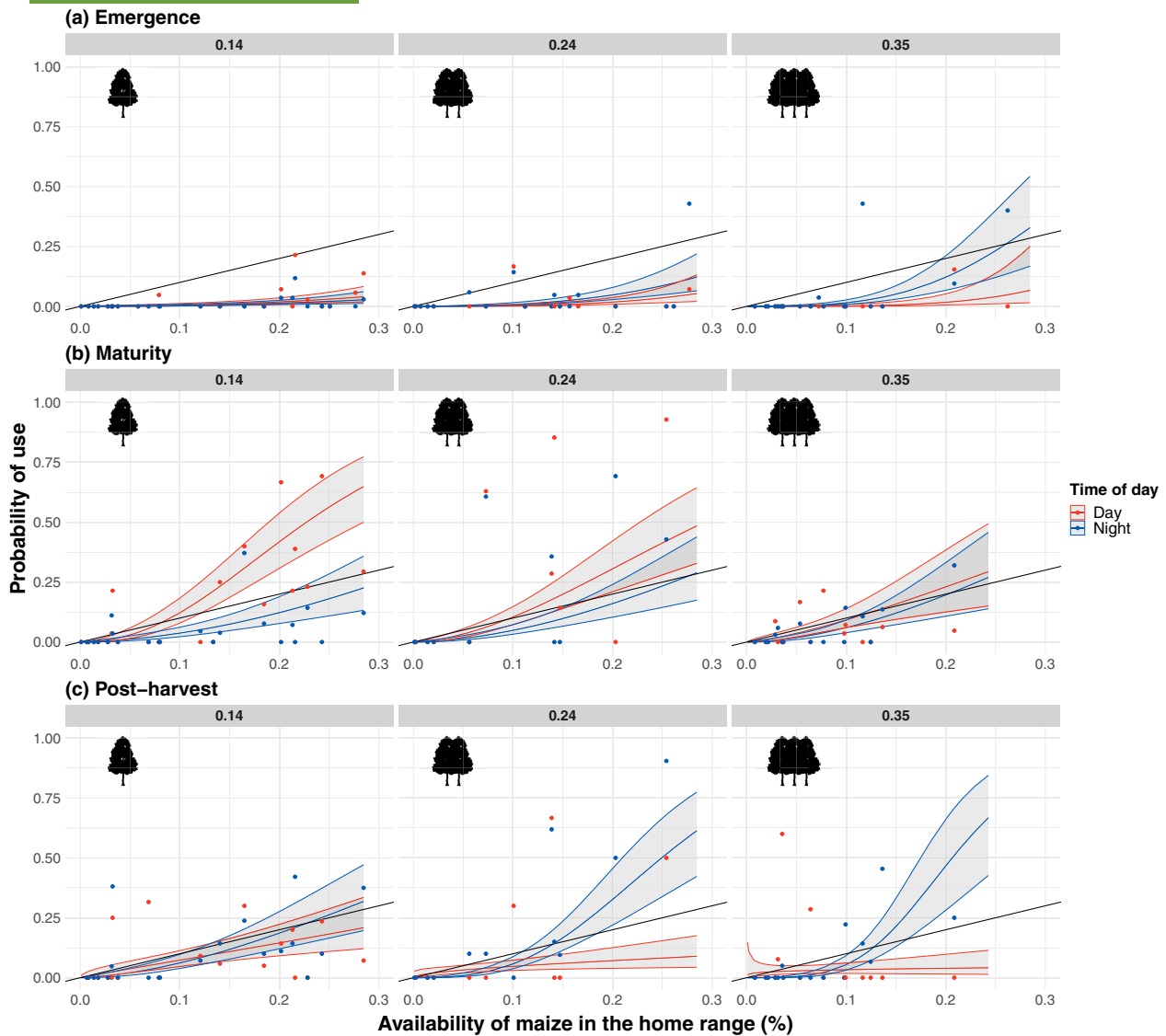


FIGURE 3 Predicted probability of use of maize during (a) emergence, (b) maturity, (c) post-harvest plotted against its availability in the home range (HR) during the day (red) and night (blue) for different proportions of 'natural' cover (woodlands and hedgerows) in the HR (14%, 24%, 35%, illustrated by the black tree icons). Coloured lines represent mean predicted probabilities with associated 95% confidence intervals estimated using logistic regressions. Points represent the observed proportion of used and available maize within the HR for each roe deer. Predictions are not represented beyond the values of availability present in the data. The diagonal black line represents the (1:1) ratio where use is equal to availability.

promote sustainable coexistence (Abrahms et al., 2021). In this context, our study provides insight on how synanthropic species that are successful in heavily human-impacted landscapes can cope with rapid and sudden modifications of their environment (e.g., cultivation) through behavioural plasticity. We found strong evidence that adult female roe deer adjust their behaviour to spatio-temporal variations in crop phenology within a heterogeneous agroecosystem. Past work on fine-scale responses to crop phenology, mainly on wild ungulates (Branco et al., 2019; Paolini et al., 2018), focused solely on the role of crops as food subsidies. Our unique data set enabled us to evaluate the potential role of crops as alternative cover habitat for adult female roe deer and to explore how crop phenology and landscape composition influence the trade-off between resource

acquisition and risk avoidance, especially when access to 'natural' cover is limited. Our results strongly suggest that the functional role of crops for adult female roe deer varied across crop types in relation to their phenological stage. We demonstrated that certain crops at certain phenological stages, notably mature maize, provide an alternative source of cover in a system where more 'natural' refuge habitat is limited. Overall, our study highlights the importance of agricultural practices and landscape composition in determining the reliance of roe deer on cultivated land for food or cover subsidies, potentially shaping the food-risk landscape for this ungulate that has widely adapted to Europe's agroecosystems.

In agroecosystems, agricultural land is intertwined with more 'natural' habitats to form a complex matrix shaped by both natural

and anthropogenic factors (Fahrig et al., 2011). Many species thrive in these highly heterogeneous agricultural landscapes (Linnell et al., 2020; Smith et al., 2020), but little is known about how they navigate among crop types that provide very different resources in terms of food and cover at specific times of the year. In the predominantly cultivated landscape of our study area, maize, wheat and artificial meadows each appear to provide alternative and complementary food resources for roe deer, but only during specific phenological stages and according to the timing of agricultural interventions (e.g., sowing, harvesting). We found that adult female roe deer used these three crops mostly at night with high activity levels (indicative of foraging: Bonnot et al., 2020) during phenological stages when the crops were either more palatable (emerging wheat, early stages for meadows, Abbas et al., 2013), or provided a higher energetic content (wild grasses in emerging wheat fields, post-harvest wheat and maize Abbas et al., 2011, 2013; Wilber et al., 2020). Indeed, substantial amounts of wheat grains and maize kernels are left on the ground after harvesting (pers. obs.) and both are known to have high non-fibrous content (including sugars), high nitrogen levels (Abbas et al., 2011), and high calorie levels (maize: Wilber et al., 2020). Seasonal variations in diet composition have previously been documented in ungulates across a wide range of environments, including agricultural areas where crops are consumed only during certain parts of the year (elephant: Branco et al., 2019; wild boar: Herrero et al., 2006; cervids: Spitzer et al., 2020). Specifically, previous studies in this agroecosystem found that maize and wheat represented a high proportion of roe deer diets during summer and autumn (Abbas et al., 2011, 2013), which is when crop harvesting occurs (Figure S2). Similarly, these authors showed that grasses and forbs could constitute a substantial part of the roe deer's diet during spring, corresponding to the early phenological stages of meadows in our study area (Figure S2). This plasticity in feeding behaviour presumably plays an important role for determining individual performance and population dynamics in agroecosystems, as opportunistically switching between crops and 'natural' vegetation likely leads to faster body growth of new-born fawns during spring/summer (roe deer: Brunot et al., unpub.) and greater subsequent body size during adulthood (sika deer: Hata et al., 2021; roe deer: Hewison et al., 2009). Nevertheless, we did not account for phenological changes in resource availability in more 'natural' habitats, such as woodlands and hedgerows. To better understand how the interplay between 'natural' and agricultural forage availability defines the foodscape for wild herbivores living in agroecosystems, future studies should investigate whether they substitute crops for woodlands in relation to woodland plant phenology (Morellet et al., 2011), especially when resources are scarce or when energetic demands are high, such as during gestation and lactation.

In highly modified landscapes such as agroecosystems, where natural habitats are scarce and the risk of human disturbance is high (Bonnot et al., 2013; Padié et al., 2015), the fact that certain crops, at certain phenological stages, could provide an alternative source of refuge for wildlife has been largely overlooked (but see Bonnot et al., 2013; Llaneza et al., 2016). A key finding of our study is that

common crops, such as maize and wheat, also provided refuge habitat for adult female roe deer at certain phenological stages. Our results suggest that, as expected, adult female roe deer exploited maize primarily for resting when it was mature and, therefore, tall enough to provide cover. Similarly, wild boar selected maize during the late growing season, when its nutritional value was lower, presumably for cover (Paolini et al., 2018). Species that cannot use crops as food sources, such as carnivores, have also been found to use them as refuge habitat: for instance, wolves living in human-dominated landscapes were found to occasionally use maize and grasslands for resting during the day, with 5.8% of their long-term diurnal bed sites found in croplands (64.5% grasslands and 32.3% corn, Llaneza et al., 2016). In contrast, adult female roe deer were active during both night and day when using mature wheat, irrespective of woodland and hedgerow availability, but avoided wheat during daytime during all other phenological stages. Therefore, we hypothesize that adult female roe deer used this habitat for both cover and food during summer, as mature wheat can conceal a standing roe deer (pers. obs., Figure S6). Although resting (vs. non-resting) behaviour is robustly identified using activity levels (Augustine & Derner, 2013), future work should combine detailed analyses of diet composition with behavioural predictions derived from accelerometer data (Brown et al., 2013) to determine whether, when and how various crop types contribute to the roe deer's food and cover landscape within agroecosystems.

Animals may satisfy their resource needs by moving between different habitat types offering similar, substitutable, resources (Dunning et al., 1992). For example, roe deer in this same agricultural landscape used hedgerows as a substitutable habitat for woodlands to provide cover when the latter was rare (Morellet et al., 2011). Similarly, here, we showed that adult female roe deer substituted mature maize for woodlands and hedgerows during the day, as selection for maize increased when 'natural' cover availability decreased ($\leq 24\%$). Most of the adult female roe deer in our study occupied home ranges with little woodland/hedgerows (mean proportion $28\% \pm 0.21$ SD, Figure S5), underlining that plasticity in habitat use is likely a key factor in determining the persistence of wildlife populations in human-dominated agricultural areas. Indeed, the habitat selection decisions of adult female roe deer concerning maize depended on its local availability, as well as the availability of more 'natural' cover in the home range. Specifically, mature maize was not selected during the day and post-harvest maize was avoided at night when it made up $<20\%$ of the home range, suggesting what Holling (1959) termed as a type IV functional response. Holling considered that type IV responses implied a 'threshold of security' below which the strength of stimuli linked to a resource is low, but above which it is high. In this context, we interpret this threshold as indicative of the costs associated with using maize relative to a roe deer's perception of risk, particularly when risk is high due to intense hunting during summer and autumn. We suggest that mature maize was only selected for when it was sufficiently abundant, and when levels of 'natural' cover were low, as exploiting these crops as refuge habitats may be perceived as too risky when 'natural' cover is easily

accessed. Similarly, post-harvest maize was used disproportionately to its availability when it was sufficiently abundant, and when the availability of refuge habitat in the surrounding landscape was high, implying that maize may be perceived as a highly rewarding, but high-risk, foraging habitat (Bonnot et al., 2013, 2018). However, there was pronounced inter-individual variation in observed responses to crops and their phenology which we speculatively attribute to differences in risk perception, with 'bold' individuals more inclined to forage in riskier habitats (spider: Steinhoff et al., 2020), such as autumn crops (roe deer: Bonnot et al., 2018). Linking individual variation in patterns of crop use with survival and reproduction would help to further understand how resource variability, risk perception and human interventions shape individual performance in a human-dominated landscape (Simon & Fortin, 2020).

Our results suggest that the ability of roe deer to subsist in agricultural landscapes with sparse 'natural' habitats may be favoured by the cultivation of crops, such as maize, that grow tall enough to provide alternative cover. The agricultural activities of humans over the past century have led to a marked decrease in the extent of woodlands and hedges in European landscapes, concomitant with an increase in cereal and oilseed production (e.g., maize, winter wheat, sunflower and rapeseed, Agreste, 2022). Our findings suggest that avoiding landscape homogenisation by simultaneously cultivating several crop types which provide alternative cover over the year may favour woodland species with high behavioural plasticity, such as roe deer. Although roe deer do not cause substantial damage to crops (2% of crop damage in France, Loison et al., 2022), we advocate maintaining heterogeneous landscapes that combine natural habitats, such as woodland and hedgerows, with high crop diversity to limit crop use and promote human-ungulate coexistence.

AUTHOR CONTRIBUTIONS

Noa Rigoudy, Simon Chamaille-Jammes, Nicolas Morellet and A. J. Mark Hewison conceived the ideas and designed the study; all authors, including Arnaud Bonnet, Yannick Chaval, Bruno Lourtet and Joël Merlet, collected the data and discussed the analytical approach; Arnaud Bonnet organized data collection and Yannick Chaval curated the database; Noa Rigoudy, Simon Chamaille-Jammes and Nicolas Morellet designed the methodology and Noa Rigoudy analysed the data with the help of Simon Chamaille-Jammes and Nicolas Morellet; Noa Rigoudy wrote the first draft of the manuscript and all authors contributed critically to revision.

ACKNOWLEDGEMENTS

We thank the local hunting associations, the Fédération Départementale des Chasseurs de la Haute Garonne, and numerous co-workers and volunteers for their assistance. We thank the Observatoire Spatial Régional (CNRS/INSU) as well as the Agir research group (INRAE) for providing local weather station data. We thank the Regional Chambre of Agriculture Occitanie and the Departmental Chambre of Agriculture Haute-Garonne for providing observational data on crop phenology from epidemiological surveillance monitoring programs in the Occitanie region collected

within the Vigicultures® project. We thank the Departmental Chambre of Agriculture Haute-Garonne for their guidance on local agricultural practices. We thank the Réseau «Ongulés sauvages OFB-FNC-FDC » for sharing data on hunting pressure in the study area. We also thank Mahdiah Tourani, Marco Apollonio and two anonymous reviewers for providing comments that critically improved this manuscript. This project was supported by 'CARRY-MOVE' Agence Nationale de la Recherche grant no. ANR-23-CE02-0008 to N.M.

CONFLICT OF INTEREST STATEMENT

The authors declare no conflict of interest.

DATA AVAILABILITY STATEMENT

Data are available via the Zenodo Repository: <https://doi.org/10.5281/zenodo.10462550> (Rigoudy et al., 2024).

STATEMENT ON INCLUSION

Our study involved scientists based in the country where the study was carried out as well as local stakeholders who work within or close to the study area. These stakeholders participated in providing data and expertise on estimating crop phenology. Each stakeholder was sent a summary of the findings of this study and was able to provide feedback on this work.

ORCID

Noa Rigoudy  <https://orcid.org/0000-0003-1505-3845>

Simon Chamaille-Jammes  <https://orcid.org/0000-0003-0505-6620>

Nicolas Morellet  <https://orcid.org/0000-0002-4274-7044>

REFERENCES

- Abbas, F., Morellet, N., Hewison, A. J. M., Merlet, J., Cargnelutti, B., Lourtet, B., Angibault, J.-M., Daufresne, T., Aulagnier, S., & Verheyden, H. (2011). Landscape fragmentation generates spatial variation of diet composition and quality in a generalist herbivore. *Oecologia*, 167(2), 401–411. <https://doi.org/10.1007/s00442-011-1994-0>
- Abbas, F., Picot, D., Merlet, J., Cargnelutti, B., Lourtet, B., Angibault, J.-M., Daufresne, T., Aulagnier, S., & Verheyden, H. (2013). A typical browser, the roe deer, may consume substantial quantities of grasses in open landscapes. *European Journal of Wildlife Research*, 59(1), 69–75. <https://doi.org/10.1007/s10344-012-0648-9>
- Abrahms, B., Aikens, E. O., Armstrong, J. B., Deacy, W. W., Kauffman, M. J., & Merkle, J. A. (2021). Emerging perspectives on resource tracking and animal movement ecology. *Trends in Ecology & Evolution*, 36(4), 308–320. <https://doi.org/10.1016/j.tree.2020.10.018>
- Agreste. (2022). GRAPH/AGRI 2022. French Ministry of Agriculture. www.agreste.agriculture.gouv.fr
- Aikens, E. O., Kauffman, M. J., Merkle, J. A., Dwinnell, S. P. H., Fralick, G. L., & Monteith, K. L. (2017). The greenscape shapes surfing of resource waves in a large migratory herbivore. *Ecology Letters*, 20(6), 741–750. <https://doi.org/10.1111/ele.12772>
- Andersen, R., Duncan, P., & Linnell, J. (1998). The European roe deer: The biology of success. In *The Journal of Wildlife Management*, 64, 608. <https://doi.org/10.2307/3803262>
- Armstrong, J. B., Takimoto, G., Schindler, D. E., Hayes, M. M., & Kauffman, M. J. (2016). Resource waves: Phenological diversity enhances

- foraging opportunities for mobile consumers. *Ecology*, 97(5), 1099–1112. <https://doi.org/10.1890/15-0554.1>
- Augustine, D. J., & Derner, J. D. (2013). Assessing herbivore foraging behavior with GPS collars in a semiarid grassland. *Sensors*, 13(3), Article 3. <https://doi.org/10.3390/s130303711>
- Barker, K. J., Mitchell, M. S., & Proffitt, K. M. (2019). Native forage mediates influence of irrigated agriculture on migratory behaviour of elk. *Journal of Animal Ecology*, 88(7), 1100–1110. <https://doi.org/10.1111/1365-2656.12991>
- Benoit, L., Bonnot, N. C., Debeffe, L., Grémillet, D., Hewison, A. J. M., Marchand, P., Puch, L., Bonnet, A., Cargnelutti, B., Cebe, N., Lourtet, B., Coulon, A., & Morellet, N. (2023). Using accelerometers to infer behaviour of cryptic species in the wild (p. 2023.03.20.533342). *bioRxiv* <https://doi.org/10.1101/2023.03.20.533342>
- Benoit, L., Morellet, N., Bonnot, N. C., Cargnelutti, B., Chaval, Y., Gaillard, J.-M., Loison, A., Lourtet, B., Marchand, P., Coulon, A., & Hewison, A. J. M. (2023). Reproductive tactics, birth timing and the risk-resource trade-off in an income breeder. *Proceedings of the Royal Society B: Biological Sciences*, 290(2009), 20230948. <https://doi.org/10.1098/rspb.2023.0948>
- Bogdziewicz, M., Zwolak, R., & Crone, E. E. (2016). How do vertebrates respond to mast seeding? *Oikos*, 125(3), 300–307. <https://doi.org/10.1111/oik.03012>
- Bonnot, N., Morellet, N., Verheyden, H., Cargnelutti, B., Lourtet, B., Klein, F., & Hewison, A. J. M. (2013). Habitat use under predation risk: Hunting, roads and human dwellings influence the spatial behaviour of roe deer. *European Journal of Wildlife Research*, 59(2), 185–193. <https://doi.org/10.1007/s10344-012-0665-8>
- Bonnot, N. C., Couriot, O., Berger, A., Cagnacci, F., Ciuti, S., De Groeve, J. E., Gehr, B., Heurich, M., Kjellander, P., Kröschel, M., Morellet, N., Sönichsen, L., & Hewison, A. J. M. (2020). Fear of the dark? Contrasting impacts of humans versus lynx on diel activity of roe deer across Europe. *Journal of Animal Ecology*, 89(1), 132–145. <https://doi.org/10.1111/1365-2656.13161>
- Bonnot, N. C., Goulard, M., Hewison, A. J. M., Cargnelutti, B., Lourtet, B., Chaval, Y., & Morellet, N. (2018). Boldness-mediated habitat use tactics and reproductive success in a wild large herbivore. *Animal Behaviour*, 145, 107–115. <https://doi.org/10.1016/j.anbehav.2018.09.013>
- Branco, P. S., Merkle, J. A., Pringle, R. M., Pansu, J., Potter, A. B., Reynolds, A., Stalmans, M., & Long, R. A. (2019). Determinants of elephant foraging behaviour in a coupled human-natural system: Is brown the new green? *Journal of Animal Ecology*, 88(5), Article 5. <https://doi.org/10.1111/1365-2656.12971>
- Brooks, M. E., Kristensen, K., van Benthem, K. J., Magnusson, A., Berg, C. W., Nielsen, A., Skaug, H. J., Mächler, M., & Bolker, B. M. (2017). GlimmTMB balances speed and flexibility among packages for zero-inflated generalized linear mixed modeling. *The R Journal*, 9(2), 378. <https://doi.org/10.32614/RJ-2017-066>
- Brown, D. D., Kays, R., Wikelski, M., Wilson, R., & Klimley, A. P. (2013). Observing the unwatchable through acceleration logging of animal behavior. *Animal Biotelemetry*, 1(1), 20. <https://doi.org/10.1186/2050-3385-1-20>
- Brown, J. S. (1992). Patch use under predation risk: I. Models and predictions. *Annales Zoologici Fennici*, 29(4), 301–309.
- Chiyo, P. I., Lee, P. C., Moss, C. J., Archie, E. A., Hollister-Smith, J. A., & Alberts, S. C. (2011). No risk, no gain: Effects of crop raiding and genetic diversity on body size in male elephants. *Behavioral Ecology*, 22(3), 552–558. <https://doi.org/10.1093/beheco/arr016>
- Doorenbos, J., & Kassam, A. H. (1979). *Yield response to water*. Food and Agriculture Organization of the United Nations.
- Dunnett, C. W. (1964). New tables for multiple comparisons with a control. *Biometrics*, 20(3), 482–491. <https://doi.org/10.2307/2528490>
- Dunning, J. B., Danielson, B. J., & Pulliam, H. R. (1992). Ecological processes that affect populations in complex landscapes. *Oikos*, 65(1), 169. <https://doi.org/10.2307/3544901>
- Fahrig, L., Baudry, J., Brotons, L., Burel, F. G., Crist, T. O., Fuller, R. J., Sirami, C., Siriwardena, G. M., & Martin, J.-L. (2011). Functional landscape heterogeneity and animal biodiversity in agricultural landscapes. *Ecology Letters*, 14(2), 101–112. <https://doi.org/10.1111/j.1461-0248.2010.01559.x>
- Hata, A., Nakashita, R., Anezaki, T., Minami, M., Fukue, Y., Higuchi, N., Uno, H., Nakajima, Y., Saeki, M., Kozakai, C., & Takada, M. B. (2021). Agricultural crop consumption induces precocious maturity in deer by improving physical and reproductive performance. *Ecosphere*, 12(4), e03464. <https://doi.org/10.1002/ecs2.3464>
- Herrero, J., García-Serrano, A., Couto, S., Ortuño, V. M., & García-González, R. (2006). Diet of wild boar *Sus scrofa* L. and crop damage in an intensive agroecosystem. *European Journal of Wildlife Research*, 52(4), 245–250. <https://doi.org/10.1007/s10344-006-0045-3>
- Hewison, A. J. M., Gaillard, J.-M., Morellet, N., Cagnacci, F., Debeffe, L., Cargnelutti, B., Gehr, B., Kröschel, M., Heurich, M., Coulon, A., Kjellander, P., Börger, L., & Focardi, S. (2021). Sex differences in condition dependence of natal dispersal in a large herbivore: Dispersal propensity and distance are decoupled. *Proceedings of the Royal Society B: Biological Sciences*, 288(1946), 20202947. <https://doi.org/10.1098/rspb.2020.2947>
- Hewison, A. J. M., Morellet, N., Verheyden, H., Daufresne, T., Angibault, J.-M., Cargnelutti, B., Merlet, J., Picot, D., Rames, J.-L., Joachim, J., Lourtet, B., Serrano, E., Bideau, E., & Cebe, N. (2009). Landscape fragmentation influences winter body mass of roe deer. *Ecography*, 32(6), 1062–1070. <https://doi.org/10.1111/j.1600-0587.2009.05888.x>
- Holling, C. S. (1959). The components of predation as revealed by a study of small-mammal predation of the European Pine Sawfly1. *The Canadian Entomologist*, 91(5), 293–320. <https://doi.org/10.4039/Ent91293-5>
- Johnson, D. H. (1980). The comparison of usage and availability measurements for evaluating resource preference. *Ecology*, 61(1), 65–71. <https://doi.org/10.2307/1937156>
- Linnell, J. D. C., Cretois, B., Nilsen, E. B., Rolandsen, C. M., Solberg, E. J., Veiberg, V., Kaczensky, P., Van Moorter, B., Panzacchi, M., Rauset, G. R., & Kaltenborn, B. (2020). The challenges and opportunities of coexisting with wild ungulates in the human-dominated landscapes of Europe's Anthropocene. *Biological Conservation*, 244, 108500. <https://doi.org/10.1016/j.biocon.2020.108500>
- Llaneza, L., García, E. J., Palacios, V., Szatornil, V., & López-Bao, J. V. (2016). Resting in risky environments: The importance of cover for wolves to cope with exposure risk in human-dominated landscapes. *Biodiversity and Conservation*, 25(8), 1515–1528. <https://doi.org/10.1007/s10531-016-1134-6>
- Loison, A., Bison, M., & Julliot, C. (2022). *Les ongulés sauvages de France métropolitaine—Fonctions écologiques, services écosystémiques et contraintes* [National evaluation of French ecosystems (EFESE)]. Ministry of Ecological Transition and Territorial Coherence. https://www.ecologie.gouv.fr/sites/default/files/efese_cgdd_rapport_ongules_sauvages_france_decembre2022.pdf
- Lüdtke, D. (2018). ggeffects: Tidy data frames of marginal effects from regression models. *Journal of Open Source Software*, 3(26), 772. <https://doi.org/10.21105/joss.00772>
- Morellet, N., Van Moorter, B., Cargnelutti, B., Angibault, J.-M., Lourtet, B., Merlet, J., Ladet, S., & Hewison, A. J. M. (2011). Landscape composition influences roe deer habitat selection at both home range and landscape scales. *Landscape Ecology*, 26(7), 999–1010. <https://doi.org/10.1007/s10980-011-9624-0>
- Mysterud, A., & Ims, R. A. (1998). Functional responses in habitat use: Availability influences relative use in trade-off situations. *Ecology*, 79(4), 1435–1441. [https://doi.org/10.1890/0012-9658\(1998\)079\[1435:FRIHUA\]2.0.CO;2](https://doi.org/10.1890/0012-9658(1998)079[1435:FRIHUA]2.0.CO;2)
- Mysterud, A., & Østbye, E. (1999). Cover as a habitat element for temperate ungulates: Effects on habitat selection and demography. *Wildlife Society Bulletin*, 27(2), 385–394.

- Padié, S., Morellet, N., Hewison, A. J. M., Martin, J.-L., Bonnot, N., Cargnelutti, B., & Chamaillé-Jammes, S. (2015). Roe deer at risk: Teasing apart habitat selection and landscape constraints in risk exposure at multiple scales. *Oikos*, 124(11), 1536–1546. <https://doi.org/10.1111/oik.02115>
- Pagon, N., Grignolio, S., Pipia, A., Bongi, P., Bertolucci, C., & Apollonio, M. (2013). Seasonal variation of activity patterns in roe deer in a temperate forested area. *Chronobiology International*, 30(6), 772–785. <https://doi.org/10.3109/07420528.2013.765887>
- Paolini, K. E., Strickland, B. K., Tegt, J. L., VerCauteren, K. C., & Street, G. M. (2018). Seasonal variation in preference dictates space use in an invasive generalist. *PLoS One*, 13(7), e0199078. <https://doi.org/10.1371/journal.pone.0199078>
- Plummer, K. E., Siriwardena, G. M., Conway, G. J., Risely, K., & Toms, M. P. (2015). Is supplementary feeding in gardens a driver of evolutionary change in a migratory bird species? *Global Change Biology*, 21(12), 4353–4363. <https://doi.org/10.1111/gcb.13070>
- Putman, R. J. (1986). Foraging by roe deer in agricultural areas and impact on arable crops. *Journal of Applied Ecology*, 23(1), Article 1. <https://doi.org/10.2307/2403083>
- R Core Team. (2020). *R: A language and environment for statistical computing*. R Foundation for Statistical Computing.
- Rigoudy, N., Chamaillé-Jammes, S., Hewison, A. J. M., Bonnet, A., Chaval, Y., Lourtet, B., Merlet, J., & Morellet, N. (2024). Data from: Crop phenology reshapes the food-safety landscape for roe deer in an agroecosystem. *Zenodo*. <https://doi.org/10.5281/zenodo.10462550>
- Signer, J., Fieberg, J., & Avgar, T. (2019). Animal movement tools (amt): R package for managing tracking data and conducting habitat selection analyses. *Ecology and Evolution*, 9(2), 880–890. <https://doi.org/10.1002/ece3.4823>
- Simon, R. N., & Fortin, D. (2020). Crop raiders in an ecological trap: Optimal foraging individual-based modeling quantifies the effect of alternate crops. *Ecological Applications*, 30(5), Article 5. <https://doi.org/10.1002/eap.2111>
- Smith, O. M., Kennedy, C. M., Owen, J. P., Northfield, T. D., Latimer, C. E., & Snyder, W. E. (2020). Highly diversified crop–livestock farming systems reshape wild bird communities. *Ecological Applications*, 30(2), Article 2. <https://doi.org/10.1002/eap.2031>
- Spitzer, R., Felton, A., Landman, M., Singh, N. J., Widemo, F., & Cromsigt, J. P. G. M. (2020). Fifty years of European ungulate dietary studies: A synthesis. *Oikos*, 129(11), 1668–1680. <https://doi.org/10.1111/oik.07435>
- Steinhoff, P. O. M., Warfen, B., Voigt, S., Uhl, G., & Dammhahn, M. (2020). Individual differences in risk-taking affect foraging across different landscapes of fear. *Oikos*, 129(12), 1891–1902. <https://doi.org/10.1111/oik.07508>
- Wilber, M. Q., Chinn, S. M., Beasley, J. C., Boughton, R. K., Brook, R. K., Ditchkoff, S. S., Fischer, J. W., Hartley, S. B., Holmstrom, L. K., Kilgo, J. C., Lewis, J. S., Miller, R. S., Snow, N. P., VerCauteren, K. C., Wisely, S. M., Webb, C. T., & Pepin, K. M. (2020). Predicting functional responses in agro-ecosystems from animal movement data to improve management of invasive pests. *Ecological Applications*, 30(1), e02015. <https://doi.org/10.1002/eap.2015>
- Wood, S., & Scheipl, F. (2020). *gam4: Generalized additive mixed models using "mgcv" and "lme4"* (R package version 0.2-6) [Computer software]. <https://CRAN.R-project.org/package=gam4>
- Worton, B. J. (1989). Kernel methods for estimating the utilization distribution in home-range studies. *Ecology*, 70(1), 164–168. <https://doi.org/10.2307/1938423>

SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

Appendix S1. Materials and methods.

Table S1. Data sources and monitoring programs used to estimate phenological stages for each crop and year.

Table S2. ANOVA table (type III) for the model predicting probability of crop use according to phenological stage and time of day for three crop types: (a) artificial meadow, (b) wheat, (c) maize.

Table S3. ANOVA table (type III) for the model predicting probability of activity according to phenological stage, time of day and crop type: maize, wheat, artificial meadow, across three phenological stages: emergence, maturity and post-harvest.

Table S4. Contrast analyses on the probability of use according to crop type, phenological stage and time of day for three crop types: (a) artificial meadow, (b) wheat, (c) maize.

Table S5. Contrast analyses on the probability of activity according to crop type, phenological stage and time of day for three crop types: (a) artificial meadow, (b) wheat, (c) maize.

Figure S1. Map of study system depicting the main land cover types.

Figure S2. Crop calendar based on weekly phenological stage estimates.

Figure S3. Effect of woodland and hedgerow availability on the functional response to wheat.

Figure S4. Effect of woodland and hedgerow availability on the functional response to artificial meadow.

Figure S5. Histogram of proportion of available woodland, hedgerows, 'natural' cover and maize per individual home range (95%) in the study population.

Figure S6. Field measures of vegetation height in sample plots for three crop types: (a) maize, (b) wheat, (c) artificial meadow in the study area from March to September 2022.

How to cite this article: Rigoudy, N., Chamaillé-Jammes, S., Hewison, A. J. M., Bonnet, A., Chaval, Y., Lourtet, B., Merlet, J., & Morellet, N. (2024). Crop phenology reshapes the food-safety landscape for roe deer in an agroecosystem. *Journal of Applied Ecology*, 61, 564–574. <https://doi.org/10.1111/1365-2664.14581>