Avian pest control in vineyards is driven by interactions between bird functional diversity and landscape heterogeneity

Luc Barbaro1,2*, Adrien Rusch3, Evalyne W. Muiruri4, Bastien Gravellier1, Denis Thiery3, Bastien Castagneyr1

1Biogeco, INRA, Univ. Bordeaux, F-33610, Cestas, France
2Dynafor, INPT, El Purpan, INRA, Univ. Toulouse F-31320, Auzeville, France
3SAVE, Bordeaux Sciences Agro, INRA, F-33140, Villenave d’Ornon, France
4School of Biological Sciences, Royal Holloway University of London, Egham, Surrey TW20 0EX, UK

*Correspondence author. E-mail: luc.barbaro@pierroton.inra.fr

Running title: Bird functional insectivory in vineyard landscapes

Summary

1. Insectivorous birds are increasingly recognised for the crucial pest control services they provide to agroecosystems. While both the foraging activity and functional diversity of birds are enhanced by multi-scale habitat heterogeneity, little is known about how these relationships may influence avian top–down control of insects.

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Specifically, interactive effects of bird community structure and habitat heterogeneity on pest control across spatial scales have rarely been explored.

2. We sampled bird communities and measured avian predation on plasticine model prey, as a proxy for lepidopteran pest control, in 20 vineyards of south-western France. Vineyards differed both in sward heterogeneity at the local scale and amount of surrounding semi-natural habitats at the landscape scale. Functional diversity metrics and community-weighted mean traits were computed for bird communities based on a species–trait table including diet, foraging method, nesting site, migration strategy, laying date, home range size, clutch size and body mass. We used mixed models to test for the interacting effects of habitat heterogeneity and bird functional diversity on avian predation rates of plasticine prey.

3. Contrary to expectations, bird functional diversity decreased with landscape-scale heterogeneity, but was higher in vineyards managed with heterogeneous sward structures. In contrast, foliage-gleaning insectivores were more abundant in landscapes supporting more semi-natural habitats, suggesting an increase in their contribution to pest control along the landscape heterogeneity gradient. Accordingly, we found that avian predation on plasticine prey increased with bird functional evenness both in more heterogeneous vineyards at the local scale and in landscape mosaics supporting more semi-natural habitats.

4. Synthesis and applications. Our study demonstrates that habitat heterogeneity at both local and landscape scales influenced avian insectivory in vineyard agroecosystems by interacting with bird community structure. It provides important insights for
ecological intensification in vineyards, pointing out that management options need to be adapted to both the functional composition of local bird communities and landscape context. We suggest that both on-field and off-field management can be used to enhance natural pest control services provided by birds in vineyards, especially by favouring sward heterogeneity and patches of semi-natural habitats within large vineyard stands at the landscape scale.

**Key-words:** agroecosystems, avian predation, bird communities, CWM traits, ecosystem services, functional insectivory, pest control, plasticine caterpillars, semi-natural habitats, vineyard management

**Introduction**

Natural pest control is a major ecosystem service delivered by a wide range of organisms, expected to offer a sustainable solution to pest management in agroecology (Chaplin-Kramer *et al.* 2011; Martin *et al.* 2013; Rusch *et al.* 2016). Recent literature reviews have highlighted the important role of insectivorous birds feeding on pest populations in different agroecosystems (Sekercioglu 2006; Wenny *et al.* 2011; Maas *et al.* 2015). Birds are especially efficient arthropod predators in farmland, where 50% of birds are predominantly feeding on insects, and 75% consume invertebrates at least occasionally (Wenny *et al.* 2011; Whelan, Sekercioglu & Wenny 2015). Avian predation on pest insects has been studied in various natural and agricultural systems, including vineyards, and most studies report a marked reduction in invertebrate biomass by birds, usually ranging from 20 to 70% (Sekercioglu 2006; Barbaro & Battisti 2011; Jedlicka, Letourneau & Cornelisse 2014). This predation not only lowers herbivore abundance but also significantly reduces leaf damage and
plant mortality, potentially leading to up to 60% increase in crop yield or fruit production (Mols & Visser 2002; Mäntylä et al. 2011; Whelan, Sekercioglu & Wenny 2015). However, despite an increasing body of evidence demonstrating the importance of birds in providing ecosystem services, the link between the functional composition of bird communities and the magnitude of ecosystem services they provide remains poorly understood (Philpott et al. 2009; Cadotte, Carscadden & Mirochnick 2011; Wenny et al. 2011).

Extensive studies of the relationships between species richness and ecosystem functions, including herbivore suppression by predators, have generally concluded that higher predator richness is associated with greater arthropod removal (Letourneau et al. 2009; Griffin, Byrnes & Cardinale 2013; De la Mora, García-Ballinas & Philpott 2015). However, there is a need for a more mechanistic understanding of the relationships between predation rates and both the species and functional composition of insectivorous bird communities (Philpott et al. 2009; Maas et al. 2015). As not all species contribute equally to ecosystem functions, it is now largely accepted that taking into account both taxonomic and functional composition of predator communities would provide a deeper understanding of the processes shaping ecosystem functions (Petchey & Gaston 2006; Hillebrand, Bennett & Cadotte 2008; Cadotte, Carscadden & Mirochnick 2011). Predation rates can be affected either by predator species abundance and richness (De la Mora, García-Ballinas & Philpott 2015) or by single- and multi-trait functional metrics (Crowder et al. 2010; Rusch et al. 2015). For example, avian insectivory may be best predicted by bird functional evenness, which measures the equidistribution of trait abundances within bird communities (Barbaro et al. 2014), or alternatively, by the abundance of foraging insectivorous birds (Bereczki et al. 2014). Several studies have also pointed out the key role played by a small number of species within the bird community, or even by a single species of functional insectivore as the main provider of pest predation in forest and farmland ecosystems (Mäntylä, Klemola & Haukioja 2004; Maas et
Thus, more work is needed to identify the primary components of bird community structure that drive natural pest control in agroecosystems (Philpott et al. 2009; Maas et al. 2015). Our understanding of avian predation patterns is also limited by a lack of studies exploring how rates of insectivory vary across spatial scales from plot- to ecosystem-level (Whelan, Sekercioglu & Wenny 2015). Trophic interactions and natural pest control services in agroecosystems depend on variables acting at multiple spatial scales, including local habitat structure, landscape context and their interactions (Martin et al. 2013; De la Mora, García-Ballinas & Philpott 2015; Tamburini et al. 2015). This appears particularly true for bird communities, which depend on both local habitat structure, especially sward heterogeneity for farmland birds, and landscape matrix composition (Vickery & Arlettaz 2012; Lindenmayer et al. 2015; Pithon et al. 2016). However, it is still unclear at which spatial scales environmental drivers act on avian insectivory and how exactly these drivers affect the relationship between bird community structure and predation rates (Martin et al. 2013; Barbaro et al. 2014; Bereczki et al. 2014; Muiruri, Rainio & Koricheva 2016).

The hypothesis that landscape context modulates the effects of local management on biodiversity and ecosystem services has been formulated as the ‘intermediate landscape complexity hypothesis’ (Kleijn et al. 2011; Tscharntke et al. 2012). According to this hypothesis, the benefits of local management on biodiversity and associated ecosystem services are lower in more complex, heterogeneous landscapes (i.e. with a high proportion of semi-natural habitats) or in extremely simplified landscapes (i.e. only croplands) than in landscapes of intermediate compositional heterogeneity (Fahrig et al. 2011). Actually, ‘complex’ landscapes already support high levels of biodiversity while simplified ones only harbour a species-poor regional pool with limited impact on ecosystem services (Tscharntke et al. 2012). To our knowledge, this hypothesis has never been tested using bird communities
and the natural pest control service they provide. Therefore, in this study, we examined how habitat heterogeneity modulates the relationship between bird communities and avian insectivory in vineyards, at both local and landscape scales. We hypothesised that: (i) landscape-scale heterogeneity increased bird functional diversity; (ii) landscape heterogeneity interacted with bird functional diversity to enhance avian insectivory in landscapes mixing vineyards and semi-natural habitats; and that (iii) landscape heterogeneity modulated the local-scale effect of sward management on predation rates by favouring foliage-gleaning insectivorous birds. In addition, we tested whether single-trait metrics of functional composition (community-weighted mean traits) perform as well or even better than multi-trait indices to predict predation rates (Rusch et al. 2015).

Materials and methods

STUDY SITES
The study area was located in Aquitaine, southwestern France, a region historically important for wine production currently covering a total of 145,000 ha of vineyards producing ca 7 millions of hl in 2014 (Fig.1). We selected 20 vineyards along a landscape heterogeneity gradient based on the proportion of semi-natural habitats (SNH), including both woodlands and semi-natural grasslands, in a 500m-radius buffer around sampled plots. The sampled range of SNH % cover lies between 0 and 68% of the landscape buffer areas, and higher SNH cover was considered to indicate higher surrounding landscape heterogeneity for a given vineyard. Previous analyses of other buffer radii (100, 250, 750 and 1000 m) have shown that 500 m was the scale best correlated with most bird community metrics. Local habitat heterogeneity was defined by the management intensity of inter-row vegetation within vine ranks: (i) homogeneous grass cover within the entire plot (i.e. low local habitat heterogeneity)
and (ii) partial (ca 50%) grass cover due to soil tillage in half of the inter-rows (i.e. high local heterogeneity). Along the landscape heterogeneity gradient, 9 vineyards had partial grass cover (i.e. soil tillage in half of the inter-rows) and 11 had full grass cover (i.e. no tillage in the inter-rows).

BIRD COMMUNITIES

Bird communities were sampled using transect counts, where all birds heard and seen were recorded except flyovers, within a width of 100 m, i.e. 50m from the observer on each transect side (Buckland 2006). We considered that species detectability did not vary among the sampled vineyards due to the highly similar and homogeneous structure of vine ranks. Transects were performed by one trained observer (LB) early in the morning (6.00 to 10.00 am) only in days without heavy rain or wind. Bird counts were conducted twice in 2013. The first visit was achieved from 18th to 25th of April (early-season breeders) and the second visit from 3rd to 5th of June (late-season breeders). For each species, the highest abundance among the two visits was used as a standardized estimate of abundance per plot for further analyses.

Functional insectivore abundance and richness were calculated by cumulating the abundance of species sharing a similar combination of bird traits regarding both diets, foraging techniques and habitat use (Jones, Sieving & Jacobson 2005). A bird species was considered a ‘functional insectivore’ in vineyards when likely to attack insect prey on vine twigs, i.e. bird species that were at the same time: (i) insectivorous during the breeding period; (ii) predominantly foraging by foliage gleaning or by hawking; and (iii) using vineyards as breeding and/or foraging habitats.
BIRD INSECTIVORY

Bird insectivory was assessed using plasticine models mimicking lepidopteran pest larvae, a type of prey commonly consumed by insectivorous birds in various ecosystems (Bereczki et al. 2014; Low et al. 2014; Muiruri, Rainio & Koricheva 2016). Plasticine models were 1 cm-long, white, inodorous and shaped to mimic Eupoecilia ambiguella and Lobesia botrana larvae, the two main lepidopteran pests of grapevine in Europe (Thiery & Moreau 2005). In each vineyard, 30 artificial larvae were fixed on six vine stocks using metal wires (diameter 0.5 mm). Three vine stocks were located at vineyard edges (in the first raw) and three vine stocks were located in the centre of the vineyard (30 m from the edge). Artificial caterpillars were exposed to predation during ten days (between 10th and 20th of June) and the typical marks led by birds were used to assess predator identity (Low et al. 2014). Avian predation rates were then estimated as the relative proportion of models showing obvious bill marks left by insectivorous birds after 10 days (Barbaro et al. 2014). Other predators recorded included small rodents, arthropods and reptiles, which were excluded from further analyses.

BIRD FUNCTIONAL DIVERSITY

The functional composition of bird communities was computed based on a species-trait matrix of 8 life-history traits, including six categorical traits (foraging method, adult diet, nesting site, migration strategy, mean laying date and mean home range size) and two continuous traits (clutch size and body mass; see Table S1 in Supporting Information). These selected traits are considered to be key indicators of individual species responses to environmental changes and their provision of pest control services (Philpott et al. 2009; Wenny et al. 2011; Sekercioglu 2012; Barbaro et al. 2014). The computation of functional diversity metrics requires that species abundance is accounted for, multiple traits are considered simultaneously (including both continuous and categorical traits) and all facets of
functional diversity are measured (Mason et al. 2005; Laliberté, Legendre & Shipley 2015). No single index matches all the criteria, therefore several complementary indices can be computed, including functional richness, evenness, divergence and entropy (Mouillot et al. 2013). For continuous traits, functional richness (FRic) for a given community is expressed as the convex hull volume of the functional trait space summarized by a principal coordinates analysis (Laliberté, Legendre & Shipley 2015). Functional evenness (FEve) is based on a minimum spanning tree measuring the regularity of trait abundance distribution within the functional space, while functional divergence (FDiv) measures trait abundance distribution within this volume and increases with extreme trait values (Mason et al. 2005). Rao’s Q measures functional entropy by characterizing species dispersion (distance weighted by abundance) from the functional space centroid, so that a high Rao’s Q value indicates a community composed of species functionally different from the mean trait composition (Ricotta & Moretti 2011; Mouillot et al. 2013). In addition to these multi-traits indices, single trait metrics were calculated using community-weighted mean (CWM) trait values (Laliberté, Legendre & Shipley 2015; Rusch et al. 2015). A CWM trait is defined for quantitative traits by the mean value of this trait in a given community, and for qualitative traits by the relative abundance of a given trait modality in each community (Ricotta & Moretti 2011). Pairwise correlations between functional metrics were checked before further analyses.

DATA ANALYSES
We first analysed the effects of local and landscape-scale heterogeneity and their interactions on bird community variables (i.e. species richness, abundance and functional diversity metrics) using quasi-Poisson GLMs to account for overdispersion in bird count data. There were several levels of non-independence in predation data due to the sampling design that
were accounted for in Generalized Linear Mixed Models (GLMM) by defining plot identity as random factor. Vine stock identity nested within plots was declared as an additional random factor to account for overdispersion (Grueber et al. 2011). Before modelling the effect of landscape heterogeneity and bird diversity on avian insectivory, we compared predation rates at vineyard edges vs interiors. GLMMs were fitted with a binomial error distribution, the response variable being defined as the number of attacked vs non-attacked plasticine caterpillars per vine stock.

As there was no difference between edges and interiors according to a $\chi^2$ test based on log-response ratio, this factor was further discarded. We then used GLMMs with landscape-scale heterogeneity (i.e. % SNH), local heterogeneity (full vs partial grass cover) and bird community metrics (BCM) as fixed effects. We initially built 17 different sets of full models as follows:

$$Y = b_0 + b_1 BCM \times (b_2 grass\ cover + b_3 SNH) + \varepsilon$$

where $b_i$ are model parameter estimates, $\varepsilon$ is residual error and $BCM$ is the bird community predictor (see Tables 1 and 2), i.e. either multi-trait metrics FRic, FEve, FDis, Rao’s Q or single CWM traits (adult diet, foraging method, clutch size, body mass).

Parameters estimates ($\pm$ SE) of binomial GLMMs that were significantly different from zero were estimated with restricted maximum likelihood (REML). Models were further compared using information theory frameworks based on Akaike’s information criterion corrected for small samples (AICc) to identify the best bird community variable interacting with local and landscape heterogeneity to fit avian insectivory rates. Models were ranked based on their AICc and we estimated model parameters for competing models within a $\Delta$AICc < 2 units of the best model with lowest AICc (Grueber et al. 2011). All model predictors were scaled and centred to allow comparing their relative effects (Schielzeth 2010). We used R-packages ‘FD’ (Laliberté, Legendre & Shipley 2015) for functional metrics, ‘lme4’ (Bates et al. 2015)
for mixed models and ‘MuMIn’ (Barton 2015) for multi-model selection.

Results

BIRD FUNCTIONAL COMPOSITION
Among the 56 bird species recorded, 27 were accordingly classified as ‘functional insectivores’ (FI). The most frequent functional insectivores recorded were European blackbird *Turdus merula*, blue tit *Cyanistes caeruleus*, common chaffinch *Fringilla coelebs*, blackcap *Sylvia atricapilla*, great tit *Parus major* and common redstart *Phoenicurus phoenicurus*. Altogether, these 6 species accounted for 26.7% of the total number of individual birds recorded (see Table S1). Total bird abundance and species richness decreased with the percentage of semi-natural habitats in the surrounding 500m (SNH), but the total abundance of functional insectivores (FI abundance) showed the opposite pattern and increased with landscape heterogeneity (Table 1). Local sward heterogeneity also affected FI abundance, with approximately 15% more functional insectivores in vineyards with full grass cover as compared to vineyards with partial cover alternating with bare ground (Table 1).

Among the multi-trait functional metrics, FRic, FDiv and Rao’s Q all decreased significantly with landscape heterogeneity, indicating higher trait richness and divergence in vineyard-dominated landscapes than in more heterogeneous landscapes (Table 1 and Fig. 2). Bird communities also tended to display higher functional diversity with partial than with full grass cover between vine ranks (Table 1). Bird functional evenness was uncorrelated to habitat heterogeneity at both local and landscape scales (Table 1). Regarding single-trait indices (CWMs), landscape heterogeneity indicated by higher SNH cover had a positive effect on productive-breeding, canopy-gleaning and bark-foraging insectivores while
negatively affecting large ground- and understorey-gleaning granivores or mixed feeders (Table 1). In contrast, local habitat heterogeneity tended to favour larger ground granivores over more productive and smaller canopy insectivores.

**BIRD INSECTIVORY**

Model selection identified bird functional evenness (FEve) as the bird community metrics best fitting predation rates in interaction with habitat heterogeneity (Table 2). There was no other competing model within two AICc units of the best model, but the second best model included functional entropy, Rao’s Q index (Table 2). The effect of bird functional evenness on insectivory was significant but depended on habitat heterogeneity at both landscape (FEve × SNH: estimate ± SE = 0.57 ± 0.16; $\chi^2 = 12.27; P < 0.0005$) and local scales (FEve × grass cover: estimate ± SE = 0.69 ± 0.30; $\chi^2 = 5.31; P < 0.02$). Avian predation increased with bird functional evenness in more heterogeneous landscapes with a large proportion of SNH, but the opposite pattern occurred in more simplified landscapes dominated by vineyards (Fig. 3). At the local scale, avian predation increased with bird functional evenness in vineyards with partial grass cover while it decreased with bird functional evenness in vineyards with full grass cover (Fig. 4). Except FDiv, functional indices based on multiple traits always ranked higher as predictors of avian predation in model selection than single traits (CWM), even those specifically pertaining to bird foraging method (Table 2).

**Discussion**

Our study demonstrates that the effect of bird functional diversity on lepidopteran pest predation in vineyards is contingent upon habitat heterogeneity both within (local scale) and
around vineyards (landscape scale). Our results therefore have important implications to vineyard management, as we show that both on-field vegetation structure and off-field landscape composition can affect how bird communities drive natural pest control of the main lepidopteran pests in European vineyards.

FUNCTIONAL INSECTIVORY AND LANDSCAPE HETEROGENEITY

Bird insectivory increased with the functional evenness of avian communities but only in more heterogeneous landscape mosaics. Although we did not validate our first prediction of bird functional diversity increasing with landscape heterogeneity, we found partial support for the ‘intermediate landscape complexity hypothesis’ predicting that landscape composition modulates the effect of local management on ecological processes and ecosystem functions (Kleijn et al. 2011; Tscharntke et al. 2012). A large body of evidence now indicates that landscape heterogeneity (or ‘complexity’ approximated by the percentage of semi-natural habitats in the surrounding matrix) enhances natural pest control (Chaplin-Kramer et al. 2011; Tamburini et al. 2015; Rusch et al. 2016). In vineyards, the ecosystem service of natural pest control is likely provided by a rather small number of functional insectivores whose diet specialization and foraging techniques are expected to allow economically significant reductions in pest insect populations (Jones, Sieving & Jacobson 2005; Jedlicka, Greenberg & Letourneau 2011; Whelan, Sekercioglu & Wenny 2015). Consistently, we found that avian predation rates increased with the relative abundances of functional insectivores and understorey foliage gleaners, as also pointed out by previous studies (Barbaro et al. 2014; Bereczki et al. 2014; Maas et al. 2015). Avian pest control also increased with landscape heterogeneity, but only when bird functional evenness was high (see Fig. 3).
PREDATOR DIVERSITY AND PEST CONTROL

According to meta-analyses, a positive effect of predator diversity on predation rates can emerge from niche complementarity, facilitation among predator species or sampling effects (Letourneau et al. 2009; Griffin, Byrnes & Cardinale 2013). While the majority of studies have confirmed an increase in predation rate with predator diversity, several works have also found neutral or even opposite effects, with greater predation in species-poor predator communities (Letourneau et al. 2009). These neutral or negative effects may result from antagonistic interactions such as competition or intraguild predation between birds and other guilds of natural enemies (Martin et al. 2013; Jedlicka, Letourneau & Cornelisse 2014). The coexistence of many bird functional types promoted by habitat diversity at the landscape scale is expected to increase intraguild competition over functional redundancy or trait complementarity (Luck, Carter & Smallbone 2013). In accordance with this hypothesis, we found that increasing the proportion of semi-natural habitats in vineyard landscapes tended to decrease bird functional diversity, thus mitigating the effect of trait evenness on insectivory on the habitat heterogeneity gradient. While bird functional evenness was the best predictor of predation rates in more heterogeneous landscapes, we assume that insectivory in simplified landscapes was provided by a low number of functional insectivores acting as a biotic insurance in such vineyard-dominated mosaics.

SWARD HETEROGENEITY AND BIRD FORAGING

Together with landscape-scale heterogeneity, sward heterogeneity is an important feature of vineyard management, affecting both insect and bird communities and potentially modifying predator-prey interactions (Arlettaz et al. 2012; Vickery & Arlettaz 2012). At the local habitat scale, a partial vegetation cover changed the functional composition of bird communities from canopy foragers towards specialist ground foragers. A denser grass cover
actually promotes the abundance of foliage-gleaning insectivores through an increase in food supply, while a sparser cover favours ground-foraging birds by increasing prey accessibility (Browne & Aebischer 2003; Vickery & Arlettaz 2012). Several specialist ground insectivores typical for vineyard agroecosystems, such as Eurasian hoopoe *Upupa epops*, Eurasian wryneck *Jynx torquilla*, woodlark *Lullula arborea*, cirl and ortolan buntings *Emberiza cirlus* and *E. hortulana* all select microhabitats with patches of bare ground where prey are more accessible even though their abundance is lower (Barbaro & Battisti 2011; Sirami, Brotons & Martin 2011; Arlettaz *et al.* 2012). Large ground granivores including turtle doves *Streptopelia turtur* also favour short and sparse vegetation cover for foraging on wild seeds (Browne & Aebischer 2003). We therefore expected that partial grass cover would overall increase avian predation by favouring the complementarity between birds with different foraging strategies. Consistently, we found that avian predation increased with bird functional evenness only in vineyards managed with heterogeneous sward structures (see Fig. 4).

A FUNCTIONAL BASIS FOR BIRD-INDUCED SERVICES

Our study supports the view that a complex interplay between bird species pool, community structure and trait diversity within bird assemblages drives the magnitude of avian predation in vineyards. It also points out the key role of the functional evenness in trait distribution within predator communities to sustain an efficient ecosystem function of pest regulation through trait complementarity (Petchey & Gaston 2006; Hillebrand, Bennett & Cadotte 2008; Crowder *et al.* 2010). How the loss of particular species disrupts ecosystem functions and services is still largely uncertain and constitutes an important area of ecological research (Cadotte, Carscadden & Mirochnick 2011; Mouillot *et al.* 2013), especially when trying to evaluate the intrinsic economic value of biodiversity conservation in agroecosystems (Whelan, Sekercioglu & Wenny 2015). Large-scale biotic homogenization has major
functional consequences through the loss of species sharing unique combinations of traits that make them especially relevant to key ecosystem functions and services (Luck, Carter & Smallbone 2013; Lindenmayer et al. 2015). The future of bird-induced services in agroecosystems is clearly a major conservation challenge under current global change (Wenny et al. 2011; Sekercioglu 2012). Whether these bird-induced services are mainly provided by single opportunistic species (Maas et al. 2015; Muiruri, Rainio & Koricheva 2016), specialized guilds of functional insectivores (Jones, Sieving & Jacobson 2005; Barbaro & Battisti 2011) or functionally-rich species assemblages is therefore a question of importance in agroecology (Philpott et al. 2009; Cadotte, Carscadden & Mirochnick 2011).

SYNTHESIS AND APPLICATIONS

Vineyard agroecosystems have high cultural and economic significance in Europe, with more than 140 millions hl produced on a total production area of 3.4 millions ha in 2014. This makes grapevine pest control particularly valuable to wine growers, although natural pest control in vineyards is very sensitive to management intensity (Jedlicka, Greenberg & Letourneau 2011; Rusch et al. 2016). Here, we found evidence for pest control by insectivorous birds depending on vineyard management at multiple scales, including local vegetation (e.g., plant diversity, sward height and heterogeneity) and semi-natural habitat cover in the surrounding landscape (woodlands and grasslands), as also suggested by previous studies of predation services in other agroecosystems (De la Mora, García-Ballinas & Philpott 2015; Rusch et al. 2015; Tamburini et al. 2015). Our study therefore has important implications for both wine growers and policymakers concerned with designing multifunctional landscapes that address both conservation and ecological intensification issues.
As a concluding remark, we suggest that conserving functional communities of insectivorous birds as a biotic insurance for natural pest control in vineyard-dominated landscapes could be achieved by simultaneously manipulating on-field vegetation (fine-grained habitat heterogeneity) and off-field management of surrounding semi-natural habitats (coarse-grained heterogeneity). Moreover, we also emphasise the increasing importance of a few species of functional insectivorous birds for biocontrol (Jones, Sieving & Jacobson 2005; Maas et al. 2015), especially in simplified landscapes dominated by large vineyards where bird diversity is generally low (Pithon et al. 2016). Interactions between natural enemy guilds might particularly be affected by changes in the functional diversity of apex predators, through increased intraguild predation (Martin et al. 2013; Jedlicka, Letourneau & Cornelisse 2014). More research is needed to examine how such changes in the functional composition of bird communities within vineyard landscapes may have cascading effects on natural pest control services provided by other guilds of natural enemies over time.

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Data accessibility

Source data for 56 bird species abundance recorded during the study and their main life attributes are available online from Table S1 in Supporting Information. Source data for
model predation rates, bird community metrics and habitat heterogeneity variables are available from Dryad Digital Repository at http://dx.doi.org/10.5061/dryad.3mf45 (Barbaro et al. 2016).

References


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Supporting information

Additional Supporting Information may be found in the online version of this article.

**Table S1.** Full list of bird species recorded in the study and their main attributes.

**Table 1.** Pearson correlation coefficients for the effects of local- and landscape-scale habitat heterogeneity on bird community metrics. Landscape heterogeneity refers to the % cover of semi-natural habitats (SNH) within a 500m-radius buffer around vineyards. Local sward heterogeneity refers to full vs partial grass cover between vine ranks. For grass cover, (-) indicates that the response variable had lower values under partial than full cover and (+) indicates higher value under partial cover. Both tests were based on linear models and all predictors were scaled before analyses. *P*-values significance thresholds as follows: ns < 0.05 < * < 0.01 < ** < 0.0001 < ***. Codes for bird community metrics as follows: FI = functional insectivores, FRic = bird functional richness, FEve = bird functional evenness, FDiv = bird functional divergence, RaoQ = bird functional entropy, CWM = community-weighted mean traits.

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<th>Grass cover</th>
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<td>Bird abundance</td>
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<td>(**)</td>
</tr>
<tr>
<td>Bird richness</td>
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<td>(***</td>
</tr>
<tr>
<td>FI abundance</td>
<td>+0.36</td>
<td>(***</td>
</tr>
<tr>
<td>FI richness</td>
<td>-0.02</td>
<td>(ns)</td>
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<tr>
<td>FRic</td>
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<td>(*)</td>
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<tr>
<td>FEve</td>
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<td>FDiv</td>
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<td>(***</td>
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</table>
Table 2. Ranking based on ΔAICc of best models comparing the relative performance of bird community metrics at fitting predation rates in interaction with local- and landscape-scale habitat heterogeneity. Bold characters refer to the best model (i.e. only model with ΔAICc < 2). See Table 1 for bird community metrics codes.

<table>
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<th>ΔAICc</th>
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<td>RaoQ</td>
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<td>CWM.hawkers flycatchers</td>
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<td>FRic</td>
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<td>CWM.bark foragers</td>
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<td>5.45</td>
</tr>
<tr>
<td>CWM.body mass</td>
<td>274.91</td>
<td>6.23</td>
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<tr>
<td>CWM.ground probers</td>
<td>275.15</td>
<td>6.47</td>
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<tr>
<td>CWM.canopy gleaners</td>
<td>275.32</td>
<td>6.64</td>
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<tr>
<td>CWM.ground gleaners</td>
<td>275.39</td>
<td>6.71</td>
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<td>CWM.insectivores</td>
<td>276.8</td>
<td>8.12</td>
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<tr>
<td>CWM.number eggs</td>
<td>277.16</td>
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<tr>
<td>FDiv</td>
<td>277.93</td>
<td>9.25</td>
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<tr>
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<td>10.13</td>
</tr>
<tr>
<td>Bird richness</td>
<td>279.27</td>
<td>10.59</td>
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Figure 1. Location map of sampled vineyards in the Aquitaine region, southwestern France (Saint Emilion and Entre-Deux-Mers areas of winegrape production). Points and circles represent the location of the 20 experimental plots and surrounding landscape buffers. Forest cover is indicated in dark green. The study area covers ca 25 x 30 km, i.e. 75 000 ha.
Figure 2. Effects of landscape (SNH) and local sward heterogeneity on bird functional diversity: (a-b) effects of landscape (a) and local scale heterogeneity (b) on bird functional entropy (Rao’s Q); (c-d) effects of landscape (c) and local scale heterogeneity (d) on bird functional evenness (FPane). Solid lines and shaded area represent model predictions and corresponding standard errors respectively. In boxplots, horizontal bars represent the median, dots represent means. $P$-values significance thresholds as follows: ns $< 0.05 < * < 0.01 < ** < 0.0001 < ***$. 
Figure 3. Interactive effects of bird functional evenness and landscape heterogeneity on avian insectivory. Landscape-scale heterogeneity is approximated by the % cover of semi-natural habitats (SNH) in the surrounding 500m around vineyards. White to black colour scale represents the increased predation rates predicted by binomial GLMM along crossed gradients of semi-natural habitat amount (SNH) and bird functional evenness (FEve). (1) In simplified landscapes, predation rate decreased with bird functional evenness (dashed line), while (2) it increased with bird functional evenness in heterogeneous landscapes (solid line). (3) Avian predation rate decreased with landscape heterogeneity when bird functional evenness was low (dashed line), while (4) it increased with landscape heterogeneity when bird functional evenness was high (solid line).
Figure 4. Interactive effects of bird functional evenness and sward heterogeneity at the local scale (full vs partial grass cover between vine ranks). Solid black lines and dark grey shaded areas represent model predictions and corresponding standard errors for full grass cover, dotted black lines and light grey shaded areas represent the same for partial grass cover.