

Strength of forest edge effects on litter-dwelling macro-arthropods across Europe is influenced by forest age and edge properties

Pallieter de Smedt, Lander Baeten, Willem Proesmans, Sam van de Poel, Johan van Keer, Brice Giffard, Ludmilla Martin, Rieneke Vanhulle, Jorg Brunet, Sara A. O. Cousins, et al.

To cite this version:

Pallieter de Smedt, Lander Baeten, Willem Proesmans, Sam van de Poel, Johan van Keer, et al.. Strength of forest edge effects on litter-dwelling macro-arthropods across Europe is influenced by forest age and edge properties. Diversity and Distributions, $2019, 25 (6)$, pp.963-974. $10.1111/dd$ i.12909. hal-02618716

HAL Id: hal-02618716 <https://hal.inrae.fr/hal-02618716v1>

Submitted on 28 May 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 - NonCommercial - NoDerivatives 4.0](http://creativecommons.org/licenses/by-nc-nd/4.0/) [International License](http://creativecommons.org/licenses/by-nc-nd/4.0/)

BIODIVERSITY RESEARCH

Strength of forest edge effects on litter‐dwelling macro‐ arthropods across Europe is influenced by forest age and edge properties

Pallieter De Smedt[1](https://orcid.org/0000-0002-3073-6751) | **Lander Baeten1** | **Willem Proesmans1** | **Sam Van de Poel1** | **Johan Van Keer²** | **Brice Giffard3,4** | **Ludmilla Martin5** | **Rieneke Vanhulle1** | **Jörg Brunet6** | **Sara A.O. Cousins7** | **Guillaume Decocq5** | **Marc Deconchat3** | **Martin Diekmann⁸** | **Emilie Gallet‐Moron5** | **Vincent Le Roux5** | **Jaan Liira9** | **Alicia Valdés5,10** | **Monika Wulf11** | **Emilie Andrieu3** | **Martin Hermy12** | **Dries Bonte13** | **Kris Verheven**¹

1 Forest & Nature Lab, Ghent University, Melle‐Gontrode, Belgium

2 Kapelle‐op‐den‐Bos, Belgium

3 UMR 1201 DYNAFOR, INRA, Chemin de Borde Rouge, Castanet, France

4 Bordeaux Sciences Agro, Université Bordeaux, Gradignan, France

⁵Jules Verne University of Picardie, UR Ecologie et Dynamique des Systèmes Anthropisés (EDYSAN, UMR 7058 CNRS), Amiens Cedex, France

6 Southern Swedish Forest Research Centre, Swedish University of Agricultural Sciences, Alnarp, Sweden

 7 Biogeography and Geomatics, Department of Physical Geography, Stockholm University, Stockholm, Sweden

 8 Institute of Ecology, FB02, University of Bremen, Bremen, Germany

- 9 Institute of Ecology and Earth Sciences, University of Tartu, Tartu, Estonia
- ¹⁰ Department of Ecology, Environment and Plant Sciences, Stockholm University, Stockholm, Sweden

11Leibniz-ZALF (e.V.), Müncheberg, Germany

¹²Division of Forest, Nature and Landscape, University of Leuven, Leuven, Belgium

13Terrestrial Ecology Unit (TEREC), Department of Biology, Ghent University, Ghent, Belgium

Correspondence

Pallieter De Smedt, Forest & Nature Lab, Ghent University, Melle‐Gontrode, Belgium. Email: pallieter.desmedt@ugent.be

Funding information ANR; MINECO; FORMAS; ETAG; DFG; BELSPO; ERC Consolidator Grant, Grant/ Award Number: 614839

Editor: Marcel Rejmanek

Abstract

Aim: Forests are highly fragmented across Western Europe, making forest edges im‐ portant features in many agricultural landscapes. Forest edges are subject to strong abiotic gradients altering the forest environment and resulting in strong biotic gradi‐ ents. This has the potential to change the forest's capacity to provide multiple ecosystem services such as nutrient cycling, carbon sequestration and natural pest control. Soil organisms play a key role in this perspective; however, these taxa are rarely considered in forest edge research.

Location: A latitudinal gradient of 2,000 km across Western Europe.

Methods: We sampled six dominant taxa of litter-dwelling macro-arthropods (carabid beetles, spiders, harvestmen, centipedes, millipedes and woodlice) in forest

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

© 2019 The Authors. *Diversity and Distributions* Published by John Wiley & Sons Ltd

edges and interiors of 192 forest fragments in 12 agricultural landscapes. We related their abundance and community composition to distance from the edge and the inter‐ action with forest age, edge orientation and edge contrast (contrast between land use types at either side of the edge).

Results: Three out of six macro‐arthropod taxa have higher activity‐density in forest edges compared to forest interiors. The abundance patterns along forest edge-to-interior gradients interacted with forest age. Forest age and edge orientation also influ‐ enced within‐fragment compositional variation along the forest edge‐to‐interior gradient. Edge contrast influenced abundance gradients of generalist predators. In general, older forest fragments, south‐oriented edges and edges along structurally more continuous land use (lower contrast between forest and adjacent land use) re‐ sulted in stronger edge-to-interior gradients while recent forests, north-oriented edges and sharp land use edges induced similarity between forest edge and interior along the forest edge‐to‐interior gradients in terms of species activity‐density and composition.

Main conclusions: Edge effects on litter‐dwelling macro‐arthropods are anticipated to feedback on important ecosystem services such as nutrient cycling, carbon sequestration and natural pest control from small forest fragments.

KEYWORDS

agricultural landscapes, beta diversity, edge effects, forest fragmentation, natural pest control, nutrient cycling, soil fauna

1 | **INTRODUCTION**

Many landscapes around the world show increasing amounts of for‐ est edges because of extensive forest fragmentation, due to land conversion for agriculture, infrastructure or residential areas (Ibisch et al., 2016; Wade, Riiters, Wickham, & Jones, 2003). In Western Europe, this fragmentation has led to scattered small forest frag‐ ments within an agricultural matrix (Decocq et al., 2016; Kolb & Diekmann, 2004) putting severe pressure on forests' biodiversity and the multiple ecosystem services it can provide (Haddad et al., 2015; Krauss et al., 2010). Altered ecological conditions due to an increased forest edge-to-interior ratio of small forest fragments are major drivers of negative effects of forest habitat fragmentation (Harrison & Bruna, 1999), in particular for forest interior specialist species (Pfeifer et al., 2017).

Forest edges have very different abiotic conditions compared to interiors for example, higher temperature, litter input… (Delgado, Arroyo, Arévalo, & Fernández‐Palacios, 2007; Gehlhausen, Schwartz, & Ausperger, 2000; Matlack, 1993) generally favour‐ ing biotic activity (De Smedt et al., 2016; Remy, Wuyts, Verheyen, Gundersen, & Boeckx, 2018) and therefore we can expect strong changes in community compositions and ecosystem processes along forest edge‐to‐interior gradients (De Smedt, Baeten, Berg et al., 2018). Furthermore, forest edges are ecotones on the brink of for‐ est and the adjacent land use where species from both habitats can

co‐occur (Boetzl, Schneider, & Krauss, 2016; Madeira et al., 2016). The strength (magnitude and depth) of the edge influence on species distribution patterns is highly context‐dependent and can be related to forest and edge characteristics. Forest age (successional devel‐ opment) influences abiotic gradients, with a cooler microclimate in older forest, which results in a stronger distinction between forest edge and interior conditions and thus higher strength of edge influ‐ ence (Matlack, 1993). This stronger distinction between forest edge and interior can result in higher differences in community composition between forest edge and interior in old compared to young forests (De la Peña et al., 2016). Similar patterns are detected for forest orientation. Microclimatic edge gradients are more extreme in south-oriented compared to north-oriented edges (Chen, Franklin, & Spies, 1995; Matlack, 1993). The shaded north‐oriented edges more closely resemble forest interior resulting in a lower divergence in community composition between forest edge and interior. Next to forest age and edge orientation, edge contrast with the adjacent land use type is an important factor determining the strength of edge ef‐ fects with high edge contrast if the land use types at either side of the edge are very different in structure, management intensity etc. It is generally assumed that "soft edges" (e.g., forest edges bordering other forest types or abandoned fields) manifest less strong edge effects compared to "hard edges" (e.g., forest edges bordering very intensive agricultural crop fields) (Peyras, Vespa Bellocq, & Zurita, 2013; Reino et al., 2009; Yekwayo, Pryke, Roets, & Samways, 2016).

More contrasting habitats have complementary species pools that can result in higher species richness in the edge ecotone. Therefore, community composition in small forest fragments can be driven by forest age, edge orientation and edge contrast but how the strength of edge effects change with distance from the forest edge has hardly been studied (Ries, Fletcher, Battin, & Sisk, 2004). Identifying the drivers of edge response strength is important for habitat conser‐ vation, because it can define the area of undisturbed interior forest habitat. Identifying unaffected interior habitat can provide more accurate predictions on population sizes, and consequently ecosys‐ tem functioning, compared to habitat area per se (Ewers & Didham, 2008).

Soil organisms and litter-dwelling fauna are key for ecosystem functioning and involved in multiple ecosystem services such as nutri‐ ent cycling, carbon sequestration and natural pest control (Costanza et al., 2017; de Vries et al., 2013; Jeffery et al., 2010). However, beside natural pest control agents (see e.g. Rand, Tylianakis, & Tscharntke, 2006; Tscharntke et al., 2012), the different components of the soil food web are rarely considered in forest edge research. Therefore, we studied the effect of edge characteristics (edge orientation and edge contrast) and the larger forest fragment context (fragment age and distance from the forest edge) on the abundance and community composition of different litter-dwelling macro-arthropods. We focussed on two dominant trophic levels of the soil fauna food web being predators, represented by most of the carabid beetles (Insecta, Coleoptera, Carabidae), spiders (Arachnida, Araneae), harvestmen (Arachnida, Opiliones) and centipedes (Chilopoda), and detritivores being millipedes (Diplopoda) and woodlice (Malacostraca, Isopoda).

Within forest fragments, we expect higher abundance of litter‐dwelling macro‐arthropods in forest edges compared to forest interiors (H1). We expect this contrast to become higher with in‐ creasing distance from the forest edge as well as increased dissim‐ ilarity in community composition between edge and interior (H2). Furthermore, we hypothesize that this distance effect is modulated by forest age and edge properties (edge orientation and edge con‐ trast with the adjacent land use) (H3).

2 | **METHODS**

2.1 | **Study area and selected forest fragments**

The study was carried out in six regions across the temperate for‐ est biome of Western Europe (Figure 1), along a latitudinal gradient spanning more than 2,000 km. In every region, we selected two 5 × 5 km landscape windows of different land use intensity (n = 12 land‐ scape windows) (see Valdés et al., (2015) for more information about landscape characteristics and Supporting Information Appendix S1 for climate data from the regions). Within each landscape window, we selected forest fragments dominated by temperate deciduous forest stands for further sampling. Purely coniferous plantations and recently afforested lands (<12 years of afforestation) were excluded. We determined the current size and historical forest continuity of all fragments using a digitized 1:25,000 map and a series of historical

FIGURE 1 Overview of the studied regions across Western Europe. FS: southern France; FN: northern France; BE: Belgium; GW: western Germany; SS: southern Sweden; SC: central Sweden

land use maps (from the 18th, 19th, 20th centuries), respectively, within a geographic information system environment (ArcGis® v.10.2, ESRI). Forest size ranged from 0.08 ha to 28.19 ha with a me‐ dian of 1.31 ha. Forest temporal continuity (hereafter called forest age) ranged from 12 years to 269 years with a median of 51 years and was quantified by a weighted average of different stand ages (based on stand area, Valdes et al., 2015). We made two categories of forest age: older forests (at least 100 years forested; 31% of for‐ ests) and "recent" forests established on former agricultural land less than 100 years ago (69% of the forests). The forest fragments oc‐ curring in a given landscape window were evenly distributed among four categories based on forest age and area (this varied between regions): small‐recent, small‐older, large‐recent and large‐older. Four fragments per category and per landscape window were retained for field sampling: hence, 16 fragments per landscape window and 192 fragments across Western Europe were selected.

2.2 | **Litter fauna sampling**

Litter-dwelling arthropods were sampled using pitfall traps (\varnothing 10 cm, depth 11 cm) buried into the soil to blind-in with the litter environment. Litter properties are variable between regions and presented in the Supporting Information Appendix S2. Pitfall trap catches are a composite measure of activity and abundance of organisms (see e.g., Woodcock, 2004), so we will therefore use the term "activity‐den‐ sity" instead of abundance. The pitfall traps contained *ca.* 200 ml of ethylene glycol and water (1/1 mixture). A drop of detergent reduced water surface tension. Traps were covered with aluminium roofs, **4 |** DE SMEDT et al.

leaving a gap of about 3 cm for arthropods to enter. We sampled in the interior (centre) of each forest fragment as well as at the edge, that is, we have two sample points for each forest fragment for a total of 384 sample points. One sample point consisted of two sam‐ ple units spaced five metres apart, resulting in four sample units per forest fragment (see De Smedt, Baeten, Proesmans et al. (2018) for more details on trap setup). We selected south-oriented or east-oriented edges (hereafter south‐oriented edges, 75% of sample size), which are supposed to favour macro-arthropods because of the warmer microclimate (Chen et al., 1995). If this aspect was not suitable (e.g., edge bordered by a ditch, road or other physical barrier), we used west-oriented or north-oriented edges (hereafter northoriented edge, 25% of sample size). Edge contrast depended on the land use intensity of the adjacent land and was classified in two categories: cropland (55% of the adjacent land use types) or grasslands (45%). Cropland can be seen as a more intensive form of agriculture with regular ploughing and a more intensive use of pesticides, hence a higher land use contrast at edge (hereafter edge contrast).

Macro‐arthropods were sampled twice betweenApril andAugust 2013 during fourteen consecutive days, in each forest fragment. To make data comparable among regions, variation in phenology across the latitudinal gradient was accounted for by starting the field sam‐ pling campaigns at Growing Degree Hours values of ca. 10,000 and 20,000 (based on data of local weather stations in 2008 and 2009), respectively. All individual carabid beetles, spiders, harvestmen, centipedes, millipedes and woodlice were identified to species level. Harvestmen and Centipedes were not taken into account in samples from Southern France, because of sample storage problems. Since carabids represent a beetle family with both predators, herbivores and omnivores we extracted feeding guild data from carabids.org (Homburg, Homburg, Schäfer, Schuldt, & Assmann, 2013). We found this data for 91.3% of the species covering 99.8% of the individuals. Of these individuals, the vast majority (91.8%) were predators. We will therefore discuss this taxonomic group together with the other predatory taxa.

2.3 | **Data analysis**

2.3.1 | **Activity‐density**

Sampling periods and sample units (replicates) were pooled at sam‐ ple point level, resulting in one activity-density value for the forest edge and one for the forest interior per forest fragment and per taxon. These two values were used to calculate an effect size expressing the strength of the edge effect, as the change in activity‐density (AD) from interior to edge. Here, we used the natural logarithm of the response ratio ADs (Hedges, Gurevitch, & Curtis, 1999): In $(AD_{edge}/AD_{interior})$. This response ratio will be referred to as edge effect on activity‐density throughout the manuscript. This edge effect on activity-density was only calculated for a taxon if at least one individual was sampled in the forest edge and one in the forest interior. This resulted in 182 forests retained for carabid beetles (95%), 183 for spiders (95%), 144 for harvestmen (90%), 72

for centipedes (45%), 168 for millipedes (88%) and 183 for woodlice (95%). First, we tested whether the edge effect on activity-density deviated from zero and differed between taxa, using a linear multi‐ level model (*lmer*-function of the *lme4*-package; Bates et al., 2016). An edge effect on activity-density above zero indicates higher activity-density in forest edges compared to interiors, while an edge effect on activity-density below zero indicates the opposite. We used landscape window nested in region as a random effect to account for variability between landscapes. We used the *summary*‐function (*t*‐test) of our model to test whether the taxa specific edge effect differed from zero. Second, we tested whether the strength of the edge effect on activity‐density (i.e., the effect size) depended on the distance between forest edge and interior, the forest age (older vs. recent), edge orientation (south‐oriented vs. north‐oriented) and edge contrast (crop vs. grassland). We also included the interaction between distance and forest age, distance and edge orientation, and distance and edge contrast, in order to test whether the effect of forest and edge characteristics on AD and community composition depended on distance between forest edge and interior. Distance was log-transformed and centred to average to zero across all regions. We used again a linear multilevel model (*lmer*‐function) with landscape window within region as random effect and allowed the distance effects to vary between taxa by including an interaction effect. Individual taxa were tested according to the same procedure (see Supporting Information Appendix S3).

2.3.2 | **Community composition**

To quantify the within‐forest fragment compositional variation between forest edge and interior, we calculated the Sørensen dis‐ similarity index between the two sampling points within each fragment and the two additive components of this dissimilarity (Baselga, 2010): the true turnover component (spatial replacement of species; Simpson index) and a nestedness component resulting from richness differences (*betapart*‐package; Baselga, Orme, Villeger, De Bortoli, & Leprieur, 2017). We only included taxon data from forests where at least 10 individuals of that particular taxon were sampled (e.g., 10 spiders, 10 centipedes etc.) to avoid many "0" and "1" values for the dissimilarity index. This resulted in 179 forests retained for carabid beetles (93%), 183 for spiders (95%), 127 for harvestmen (79%), 25 for centipedes (16%), 152 for millipedes (79%) and 181 for woodlice (94%). First, for the three components of community composition, that is, Sørensen dissimilarity index (referred to as compositional var‐ iation), species turnover and nestedness, we used a linear multilevel model (Imer-function) to test for differences between taxa. We used landscape window nested in region as a random effect to account for differences between landscapes. To get pairwise comparisons between taxa, we performed a Tukey post hoc comparison with the *glht*‐function from the *multcomp*‐package (Hothorn et al., 2016) (see Supporting Information Appendix S4). Second, we tested whether compositional variation depended on distance between forest edge and interior, forest age, edge orientation and edge contrast accord‐ ing to the same procedure as used for the activity‐density model.

We tested individual taxa for compositional variation according to the same procedure as the activity‐density model (see Supporting Information Appendix S5). All analyses were performed within the statistical software of R (R Core Team, 2017).

3 | **RESULTS**

Across all six taxa, we identified 182,118 individuals covering 598 species of which are 29,100 carabid beetles (150 species), 29,632 spiders (331 species), 23,378 harvestmen (25 species), 13,327 milli‐ pedes (39 species), 909 centipedes (26 species) and 85,769 woodlice (27 species) (see Supporting Information Appendix S6 for complete species lists).

3.1 | **Edge effect on activity‐density**

The edge effect on activity-density differed significantly between taxa $(F_{(6,141)} = 15.12, p < 0.001)$ (Figure 2). Spiders $(t_{182} = 6.27,$ *p* < 0.001), millipedes (t_{201} = 3.51, *p* < 0.001) and woodlice $(t₁₈₂ = 6.22, p < 0.001)$ had higher activity-density in forest edges compared to interiors. Harvestmen activity‐density tended to be higher in forest interiors compared to forest edges (but note that t_{211} = −1.81, *p* = 0.07). Activity-density of carabid beetles (t_{184} = 1.19, *p* = 0.24), and centipedes (t_{412} = −0.02, *p* = 0.98) did not differ between forest edges and interiors.

Across all taxa, the edge effect on activity‐density was influ‐ enced by the distance between forest edge and interior (Table 1) but only in older forest, not in recent forest (Figure 3). In older forest, larger distances between forest edges and interiors led to stronger increases in activity‐density from interior to edge (more positive log-ratio). Fragments bordered by grasslands showed edge effects for carabid beetles and spiders (Supporting Information Table S3.1). However, both taxa show an opposite pattern. For carabid beetles, larger distances between forest edges and interiors led to more

FIGURE 2 Edge effect on activity-density, calculated as a log response ratio effect size, on six taxa of litter-dwelling macroarthropods across Western Europe. A response ratio of zero indicates equal activity‐density in forest edges and forest interiors and positive (negative) values higher (lower) activity‐density at the forest edge. Points and error bars represent mean effects ±1 *SE*

 | DE SMEDT et al. **5**

neutral responses (response of 0) in activity‐density from interior to edge when edges were bordered by grasslands. At large distances from the forest edge, interior communities have even higher activity-density compared to edge communities (Supporting Information Figure S3.1). The edge effect on activity-density for spiders increased when distances between forest edges and interiors became larger (Supporting Information Figure S3.2). For carabid beetles and spiders, there was no response for edges bordered by cropland. The edge effect on activity‐density depended on edge orientation for spiders, with larger distances between forest edges and interi‐ ors leading to stronger decreases in activity‐density from interior to edge for north‐oriented edges while opposite for south‐oriented edges (Supporting Information Figure S3.3). Edge effect on millipede activity‐density increased with increasing distance from the forest edge (Supporting Information Figure S3.4). Edge effects on wood‐ lice activity‐density showed a similar pattern as the overall trend (Supporting Information Figure S3.5). No significant trends were observed for harvestmen and centipedes (Supporting Information Table S3.1).

3.2 | **Edge effect on community composition**

Edge effects on community composition (compositional variation) within forest fragments differed between taxa ($F_{(5,822)}$ = 108.84, *p* < 0.001), with spiders showing the highest overall compositional variation, followed by carabid beetles (Figure 4a). Centipedes and millipedes had a lower overall compositional variation, but higher compared to harvestmen and woodlice (Figure 4a) (see Supporting Information Appendix S4). For carabid beetles and spiders, the com‐ positional variation between forest interior and edge was mostly derived from turnover (70% and 83%, respectively). For the other taxa, compositional variation was derived from both turnover and nestedness (Figure 4). Species turnover patterns were similar com‐ pared with overall compositional variation (Figure 4b). Nestedness was lower for spiders compared to all other taxa, except woodlice (Figure 4c) and lower for woodlice compared to millipedes (Figure 4c) (see Supporting Information Appendix S4).

TABLE 1 Results of the multilevel model testing the effect of distance (log‐transformed), forest age, edge contrast and edge orientation on log‐ratio activity‐density of forest edge versus forest interior from six taxa and across Western Europe. Results (*F*‐values and degrees of freedom) of linear multilevel models

p* < 0.05, **p* < 0.001.

Across all taxa, compositional variation and turnover were in‐ fluenced by the actual distance between forest edge and interior (Table 2) in older forest (Figure 5a) and south‐oriented forest edges (Figure 5d), not in recent forest (Figure 5b) or in north‐oriented forest edges (Figure 5c). In older forest and south‐oriented forest edges, larger distances between forest edge and interior led to stronger compositional variation between the forest edge and inte‐ rior. Splitting‐up the analysis for separate taxa revealed a significant increase of compositional variation for spiders with increasing dis‐ tance from forest edge and interior, at least in south‐oriented edges but not in north‐oriented edges (Supporting Information Figure S5.1). Millipede compositional variation increased with increasing distance between forest edge and interior in older forest but not in recent forest (consistent with the overall pattern) (Supporting Information Figure S5.2).

4 | **DISCUSSION**

For three out of six litter-dwelling macro-arthropod taxa (i.e., spiders, millipedes and woodlice), activity‐density was higher in forest edges compared to forest interiors. Carabid beetles, harvestmen and centipedes did not show a significant response. The contrast in ac‐ tivity‐density between forest edges and interiors depended strongly on distance between forest edge and interior, which was also the case for compositional variation. The effect interacted for multiple groups with forest (edge) characteristics: responses were stronger in older forest, south‐oriented edges and lower contrast edges that is, with grassland on the outside.

4.1 | **Activity‐density in forest edges versus interiors**

The higher activity-density of different litter-dwelling taxa suggests that the habitat conditions in forest edges are more favour‐ able for arthropods compared to forest interiors. Forest edges have

FIGURE 3 Edge effect on activitydensity, calculated as a log response ratio effect size, according to distance of the forest interior from the forest edge for (a) older forest fragments and (b) recent forest fragments across Western Europe. Different colours represent the six taxa of litter‐dwelling macro‐arthropods. Data based on 192 forest fragments in 12 landscapes across Western Europe. Black line represents mean ±1 *SE* (shaded area). Italic numbers represent non‐transformed distances from the forest edge

higher temperatures (Delgado et al., 2007; Heithecker & Halpern, 2007), which increases arthropod activity, metabolic rate and re‐ sults in a shorter reproduction time (Gillooly, Brown, West, Savage, & Charnov, 2001). Additionally, edge leaf litter has a lower C/N‐ratio (De Smedt et al., 2016) resulting in better food quality for detritivore taxa, such as woodlice and millipedes (David & Handa, 2010). However, an important determinant of soil arthropod distribution and survival is humidity, which is lower at forest edges compared to interiors (Chen et al., 1995; Gehlhausen et al., 2000). Different stud‐ ies have pointed out humidity as a key factor shaping soil arthropod distribution (Bogyó, Magura, Nagy, & Tóthmérész, 2015; David & Handa, 2010; Hornung, 2011; Pearce & Venier, 2006), and how well species can cope with low moisture levels is highly species‐specific (Dias et al., 2013). This can help us to understand the distribution of harvestmen along forest edge‐to‐interior gradients. Harvestmen can be divided in two groups being true soil dwellers and vegetation dwellers. Pitfall traps, in particular sample soil dwellers (De Smedt & Van de Poel, 2017), which are more drought sensitive compared to the thermophilic shrub dwellers (Curtis & Machado, 2007). The used sampling technique, biased to soil dwellers, might therefore explain the observed tendency of harvestmen to be more common in for‐ est interiors compared to forest edges. The observed distribution patterns are probably a net effect of species that have the ability to withstand dry conditions and built‐up large populations in forest edges compared to drought‐sensitive species that retreat to forest interiors. De Smedt, Baeten, Berg et al. (2018) investigated species‐ specific distribution patterns of woodlice along forest edge-to-interior gradients and discovered that the patterns could be explained based on species desiccation resistance. The most drought‐sensitive species showed a negative activity‐density response towards forest edges. This is also the species, which have the lowest number of eggs in their brood pouch and can be considered as woodlice K‐strate‐ gists (Warburg, Linsenmair, & Bercovitz, 1984). The drought‐resist‐ ant species from the forest edge on the contrary have higher number of eggs (r‐strategists) enabling them to build up large populations in a short time‐period (Warburg et al., 1984). The r/K strategy theory

variation (±1 *SE*) between the interior and edge community within forest fragments for six taxa of litter‐dwelling macro‐arthropods across Western Europe for (a) compositional variation (Sørensen dissimilarity index) and its two additive components: (b) turnover (Simpson Index) and (c) nestedness. Percentages report the proportion of community variation that is attributed to turnover or nestedness. Significance values between taxa can be found in Supporting Information Appendix S4

FIGURE 4 Mean compositional

has also been established for plants in forest edges (Chabrerie, Jamoneau, Gallet‐Moron, & Decocq, 2013). Next to, environmental conditions favouring fast population growth, the increased use of pesticides close to forest edges might also select for r‐strategists (Chabrerie et al., 2013). Nevertheless, the net overall distribution pattern of the different taxa is a balance between the positive ef‐ fects of suitable environmental and food conditions in forest edges and the negative response to decreased moisture availability.

4.2 | **Activity‐density contrast and community composition along edge‐to‐interior gradients**

The edge response on activity-density depended on the distance between forest edge and interior across all taxa, in interaction with for‐ est age (see below). This was also the case for compositional variation where distance between forest edge and interior always interacted with forest (edge) characteristics, proving the strong context-dependency of forest edge-to-interior gradients. Compositional variation between forest edge and interior was highest for species‐rich taxa (spiders and carabid beetles) and lowest for species‐poor taxa (wood‐ lice and harvestmen). Local communities of species‐rich taxa may be assembled from a larger species pool and show higher local site‐to‐site variation (Baselga, 2010). Species‐poor taxa had low relative turno‐ ver rates, this could be explained by (a) interior communities being largely a subset of edge communities or (b) edge communities becom‐ ing more species rich because of spillover from adjacent agricultural land. De Smedt, Baeten, Berg et al. (2018) have supported the first for woodlice, where most species (despite some exceptions) decrease in activity‐density from the forest edge to the interior with few typical forest interior species. However, typical interior communities have **WII FY— Diversity** and **Distributions ———————————————————————————————————DE SMEDT ET AL.**

TABLE 2 Results of the multilevel model testing the effect of distance (log‐transformed), forest age, edge contrast and edge orientation on compositional variation (Sørensen dissimilarity index), turnover (Simpson index) and nestedness between forest edge and interior communities of litter‐dwelling macro‐arthropods from six taxa and across Western Europe. Results (*F*‐values and degrees of freedom) of linear multilevel models

p < 0.1,

p* < 0.01, *p* < 0.001.

been found for millipedes (De Smedt, Baeten, Proesmans et al., 2018) and centipedes (Lacasella et al., 2015). A species-rich taxon, like carabid beetles, showed a higher degree of typical interior species (Soga, Kanno, Yamaura, & Koike, 2013; Tóthmérész, Nagy, Mizser, Bogyó, & Magura, 2014), but this has hardly been found for the species‐rich spi‐ ders (Kowal & Cartar, 2012; Lacasella et al., 2015). However, for these two groups spillover from adjacent fields to forest edges has been re‐ ported (Boetzl et al., 2016; Tscharntke et al., 2012), which can cause the higher compositional variation between forest edge and interiors for carabid beetles and spiders.

4.3 | **Modulation of edge effects by forest (edge) characteristics**

Distance between forest edge and interior influenced edge effects on activity‐density in older forests. This can be associated to abiotic

gradients in older forest being stronger, compared to recent forest fragments, with overall lower temperatures and higher humidity in older forest (longer successional development) compared to recent forest (Baker et al., 2014). Hence, abiotic gradients are less devel‐ oped in recent fragments resulting in weaker gradients of arthropod abundance (Ng, Barton, Macfadyen, Lindenmayer, & Driscoll, 2018). Compositional variation showed an increase with distance from the forest edge in older fragments, but a neutral response in recent for‐ est fragments. This indicates that larger distances between edge and interior communities lead to stronger compositional variation in older forest, but not in recent forest. In recent fragments, there is a degree of dissimilarity between edge and interior but this does not change when moving deeper into the forest fragment. This effect was found for both compositional variation and turnover, but not for nestedness. Therefore, turnover has a larger share in compositional variation at larger distance between edge and interior, indicative for increased species replacement in older forest. Interior communi‐ ties are more stable compared to edge communities in older forest (Ewers & Didham, 2008), but this might not be the case for recent forest fragments, which can be more dynamic compared to older forest because of the ongoing forest succession. These recent for‐ ests lack, for example, characteristic older forest vegetation (Flinn & Vellend, 2005; Verheyen, Bossuyt, Honnay, & Hermy, 2003) and differ from older forest in microbial and micro-fauna community (de la Peña et al., 2016), providing different food and habitat properties for litter‐dwelling macro‐fauna.

Spiders are the only studied taxon in which the strength of ac‐ tivity‐density response depended on edge orientation. As expected, the contrast between forest edges and interiors increases with dis‐ tance in south‐oriented forest edges, because of the more extreme abiotic gradients (Chen et al., 1995; Matlack, 1993). The reversed pattern in north‐oriented forest edges is however difficult to explain, but could indicate that north‐oriented forest edges have less favour‐ able environmental conditions for spiders compared to forest inte‐ riors. Spider distribution is strongly related to vegetation structure because they need complexity for both hunting and web building (Hatley & Macmahon, 1980; Uetz, 1991) for example, in forest edges (Baldissera, Ganade, & Fontoura, 2004). Vegetation structure and plant community composition show a stronger gradient and larger penetration depth in south‐oriented compared to north‐oriented edges (Fraver, 1994; Honnay, Verheyen, & Hermy, 2002), which might explain the observed spider gradients. The same mechanisms are probably causing the observed compositional variation pattern for spiders, and all taxa combined. South‐oriented edges provide therefore not only habitat for more individuals but also for different species compared to forest interiors.

All spiders and more than 90% of our carabid individuals are predators with great potential for natural pest control (Holland & Luff, 2000; Marc & Canard, 1997; Symondson, Sunderland, & Greenstone, 2002; Tscharntke et al., 2012). Carabid beetles and spiders showed a changing activity-density response along the forest edge-to-interior gradients at lower contrast edges (grasslands) but not on higher contrast edges (crop fields). Most of the studied

FIGURE 5 Edge effect on the compositional variation between interior and edge communities of six taxa of litter-dwelling macroarthropods according to distance between forest edge and interior for (a) older and (b) recent fragments according and (c) north‐ or (d) south-oriented edges across Western Europe. Different colours represent the six litter-dwelling macro-arthropods. Data based on 192 forest fragments in 12 landscapes across Western Europe. Black line represents mean ±1 *SE* (shaded area). Italic numbers represent non‐ transformed distances from the forest edge

arthropod predators cannot complete their life cycle on crop fields (Samu & Szinetár, 2002), while at least some species can on perma‐ nent grasslands. This could result in a more stable edge community, because of a more stable grassland community compared to crop fields at least for spiders (Birkhofer, Smith, Weisser, Wolters, & Gossner, 2015), resulting in a more constant spillover rate compared to crop fields. Croplands are dynamic because of annual soil man‐ agement and more intensive use of pesticides and this could result in more temporal fluxes of predators. These findings support the potential of predatory taxa from the agricultural matrix to influence arthropod dynamics in semi‐natural vegetation fragments through their edge (see e.g., Tscharntke, Rand, & Bianchi, 2005). However, the observed patterns might be explained by the fact that meadows and croplands have different arthropod communities (Jeanneret, Schüpbach, Pfiffner, & Walter, 2003; Samu & Szinetár, 2002), and species‐specific responses might govern distributional patterns (Magura, 2002; Niemelä, Langor, & Spence, 1993).

4.4 | **Conclusion**

We showed strong edge responses of litter-dwelling arthropods with generally higher activity-density in forest edges. However, the strength of the edge effect depended on the actual distance be‐ tween interior and edge and the interaction with forest age, edge orientation and edge contrast. Forest edge research has found strong abiotic gradients and there is increasing evidence that these gradients are translated into strong biotic gradients for an increas‐ ing number of taxa. This study shows this for dominant litter‐dwell‐ ing macro‐arthropods with consistent patterns in temperate forest fragments across Western Europe spanning a gradient of more than 2,000 km from Southern France up to central Sweden. In conse‐ quence, these strong gradients could have important consequences for ecosystem functioning. All studied taxa are important links in the detrital food web and therefore of key importance for nutrient cycling in forest habitats (Lavelle, 1997). Of the global net primary

10 a
b b DE SMEDT et al. **b Diversity** and **Distributions**

production, 80% enters the detrital food web. However this, food web receives far less attention compared to aboveground food webs (Moe et al., 2005). This fact stresses the need for detrital food web studies. Using a food web modelling approach, tracking nutrients and biomass across the detrital food web in forest edges and interiors, can give us important insights in how forest functioning is affected by forest edges. This can enable us to understand how forest edges affect the ecosystems ability to supply multiple ecosystem services (such as carbon sequestration, nutrient mineralization, water purification, natural pest control) in small forest fragments in agricultural landscapes.

ACKNOWLEDGEMENTS

Many thanks to Kent Hansson, Jessica Lindgren and Sabine Sigfridsson for help during field sampling. This research was funded by the ERA‐Net BiodivERsA project smallFOREST, with the national funders ANR (France), MINECO (Spain), FORMAS (Sweden), ETAG (Estonia), DFG (Germany), BELSPO (Belgium) and DFG (Germany) part of the 2011 BiodivERsA call for research proposals. P.D.S. has a postdoctoral and W.P. a doctoral fellowship of the Research Foundation‐Flanders (FWO). The research of K.V. is supported by the ERC Consolidator Grant 614839 – PASTFORWARD. L.B. and D.B. are supported by the FWO research network EVENET.

DATA ACCESSIBILITY

All data are available in the smallFOREST geodatabase. Access to this database can be achieved after contacting the smallFOREST geodatabse management committee [\(http://www.u-picardie.fr/](http://www.u-picardie.fr/smallforest/uk/) [smallforest/uk/\)](http://www.u-picardie.fr/smallforest/uk/).

ORCID

Pallieter De Smed[t](https://orcid.org/0000-0002-3073-6751) <https://orcid.org/0000-0002-3073-6751>

REFERENCES

- Baker, T. P., Jordan, G. J., Steel, E. A., Fountain‐Jones, N. M., Wardlaw, T. J., & Baker, S. C. (2014). Microclimate through space and time: Microclimatic variation at the edge of regeneration forests over daily, yearly and decadal time scales. *Forest Ecology and Management*, *334*, 174–184. <https://doi.org/10.1016/j.foreco.2014.09.008>
- Baldissera, R., Ganade, G., & Fontoura, S. B. (2004). Web spider commu‐ nity response along an edge between pasture and Araucaria forest. *Biological Conservation*, *118*(3), 403–409. [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.biocon.2003.09.017) [biocon.2003.09.017](https://doi.org/10.1016/j.biocon.2003.09.017)
- Baselga, A. (2010). Partitioning the turnover and nestedness components of beta diversity. *Global Ecology and Biogeography*, *19*(1), 134– 143. <https://doi.org/10.1111/j.1466-8238.2009.00490.x>
- Baselga, A., Orme, D., Villeger, S., De Bortoli, J., & Leprieur, F. (2017). Partitioning Beta Diversity into Turnover and Nestedness Components. R package version 1.4‐1. [https://cran.r-project.org/](https://cran.r-project.org/web/packages/betapart/betapart.pdf) [web/packages/betapart/betapart.pdf](https://cran.r-project.org/web/packages/betapart/betapart.pdf)
- Bates, D., Maechler, M., Bolker, B., Walker, S., Bojesen Christensen, R. H., Singmann, H., … Green, P. (2016). lme4: Linear mixed‐effects models

using 'Eigen' and S4. R package version 1.1‐12. [https://cran.r-project.](https://cran.r-project.org/web/packages/lme4/) [org/web/packages/lme4/](https://cran.r-project.org/web/packages/lme4/)

- Birkhofer, K., Smith, H. G., Weisser, W. W., Wolters, V., & Gossner, M. M. (2015). Land‐use effects on the functional distinctness of arthropod communities. *Ecography*, *38*(9), 889–900. [https://doi.org/10.1111/](https://doi.org/10.1111/ecog.01141) [ecog.01141](https://doi.org/10.1111/ecog.01141)
- Boetzl, F. A., Schneider, G., & Krauss, J. (2016). Asymmetric carabid bee‐ tle spillover between calcareous grasslands and coniferous forests. *Journal of Insect Conservation*, *20*(1), 49–57. [https://doi.org/10.1007/](https://doi.org/10.1007/s10841-015-9838-6) [s10841-015-9838-6](https://doi.org/10.1007/s10841-015-9838-6)
- Bogyó, D., Magura, T., Nagy, D. D., & Tóthmérész, B. (2015). Distribution of millipedes (Myriapoda, Diplopoda) along a forest interior–forest edge–grassland habitat complex. *ZooKeys*, *510*, 181–195. [https://doi.](https://doi.org/10.3897/zookeys.510.8657) [org/10.3897/zookeys.510.8657](https://doi.org/10.3897/zookeys.510.8657)
- Chabrerie, O., Jamoneau, A., Gallet‐Moron, E., & Decocq, G. (2013). Maturation of forest edges is constrained by neighbouring agricul‐ tural land management. *Journal of Vegetation Science*, *24*(1), 58–69. <https://doi.org/10.1111/j.1654-1103.2012.01449.x>
- Chen, J., Franklin, J. F., & Spies, T. A. (1995). Growing‐season micro‐ climatic gradients from clearcut edges into old‐growth Douglas‐ fir forests. *Ecological Applications*, *5*(1), 74–86. [https://doi.](https://doi.org/10.2307/1942053) [org/10.2307/1942053](https://doi.org/10.2307/1942053)
- Costanza, R., de Groot, R., Braat, L., Kubiszewski, I., Fioramonti, L., Sutton, P., … Grasso, M. (2017). Twenty years of ecosystem ser‐ vices: How far have we come and how far do we still need to go? *Ecosystem Services*, *28*, 1–16. [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.ecoser.2017.09.008) [ecoser.2017.09.008](https://doi.org/10.1016/j.ecoser.2017.09.008)
- Curtis, D. J., & Machado, G. (2007). Ecology. In R. Pinto‐da‐Rocha, G. Machado, & G. Giribet (Eds.), *Harvestmen: The biology of Opiliones* (pp. 208–308). Cambridge, MA: Harvard University Press.
- David, J. F., & Handa, I. T. (2010). The ecology of saprophagous macroarthropods (millipedes, woodlice) in the context of global change. *Biological Reviews*, *85*(4), 881–895. [https://doi.](https://doi.org/10.1111/j.1469-185X.2010.00138.x) [org/10.1111/j.1469-185X.2010.00138.x](https://doi.org/10.1111/j.1469-185X.2010.00138.x)
- de la Peña, E., Baeten, L., Steel, H., Viaene, N., De Sutter, N., De Schrijver, A., & Verheyen, K. (2016). Beyond plant–soil feed‐ backs: Mechanisms driving plant community shifts due to land‐ use legacies in post‐agricultural forests. *Functional Ecology*, *30*(7), 1073–1085.
- De Smedt, P., Baeten, L., Berg, M. P., Gallet‐Moron, E., Brunet, J., Cousins, S. A. O., … Verheyen, K. (2018). Desiccation resistance determines distribution of woodlice along forest edge‐to‐interior gradients. *European Journal of Soil Biology*, *85*, 1–3. [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.ejsobi.2017.12.002) [ejsobi.2017.12.002](https://doi.org/10.1016/j.ejsobi.2017.12.002)
- De Smedt, P., Baeten, L., Proesmans, W., Berg, M. P., Brunet, J., Cousins, S. A. O., … Verheyen, K. (2018). Linking macrodetritivore distribution to desiccation resistance in small forest fragments embedded in agricultural landscapes in Europe. *Landscape Ecology*, *33*(3), 407–421. <https://doi.org/10.1007/s10980-017-0607-7>
- De Smedt, P., & Van de Poel, S. (2017). Succession in harvestman (Opiliones) communities within an abandoned sand quarry in Belgium. *Belgian Journal of Zoology*, *147*(2), 155–169. [https://doi.](https://doi.org/10.26496/bjz.2017.13) [org/10.26496/bjz.2017.13](https://doi.org/10.26496/bjz.2017.13)
- De Smedt, P., Wuyts, K., Baeten, L., De Schrijver, A. N., Proesmans, W., De Frenne, P., … Verheyen, K. (2016). Complementary distribution patterns of arthropod detritivores (woodlice and millipedes) along forest edge‐to‐interior gradients. *Insect Conservation and Diversity*, *9*(5), 456–469. <https://doi.org/10.1111/icad.12183>
- de Vries, F. T., Thebault, E., Liiri, M., Birkhofer, K., Tsiafouli, M. A., Bjornlund, L., … Bardgett, R. D. (2013). Soil food web properties explain ecosystem services across European land use systems. *Proceedings of the National Academy of Sciences*, *110*(35), 14296– 14301. <https://doi.org/10.1073/pnas.1305198110>
- Decocq, G., Andrieu, E., Brunet, J., Chabrerie, O., De Frenne, P., De Smedt, P., … Wulf, M. (2016). Ecosystem services from small forest

 DE SMEDT et al. **11 CONTRACTED 11 Diversity** and **Distributions** $-WII$ **EY** $\frac{11}{2}$

patches in agricultural landscapes. *Current Forestry Reports*, *2*(1), 30– 44. <https://doi.org/10.1007/s40725-016-0028-x>

- Delgado, J. D., Arroyo, N. L., Arévalo, J. R., & Fernández‐Palacios, J. M. (2007). Edge effects of roads on temperature, light, canopy cover, and canopy height in laurel and pine forests (Tenerife, Canary Islands). *Landscape and Urban Planning*, *81*(4), 328–340. [https://doi.](https://doi.org/10.1016/j.landurbplan.2007.01.005) [org/10.1016/j.landurbplan.2007.01.005](https://doi.org/10.1016/j.landurbplan.2007.01.005)
- Dias, A. T. C., Krab, E. J., Mariën, J., Zimmer, M., Cornelissen, J. H. C., Ellers, J., … Berg, M. P. (2013). Traits underpinning desiccation resis‐ tance explain distribution patterns of terrestrial isopods. *Oecologia*, *172*(3), 667–677. <https://doi.org/10.1007/s00442-012-2541-3>
- Ewers, R. M., & Didham, R. K. (2008). Pervasive impact of large‐scale edge effects on a beetle community. *Proceedings of the National Academy of Sciences*, *105*(14), 5426–5429. [https://doi.org/10.1073/](https://doi.org/10.1073/pnas.0800460105) [pnas.0800460105](https://doi.org/10.1073/pnas.0800460105)
- Flinn, K. M., & Vellend, M. (2005). Recovery of forest plant communi‐ ties in post‐agricultural landscapes. *Frontiers in Ecology and the Environment*, *3*(5), 243–250. [https://doi.org/10.1890/1540-9295\(20](https://doi.org/10.1890/1540-9295(2005)003[0243:ROFPCI]2.0.CO;2) [05\)003\[0243:ROFPCI\]2.0.CO;2](https://doi.org/10.1890/1540-9295(2005)003[0243:ROFPCI]2.0.CO;2)
- Fraver, S. (1994). Vegetation responses along edge-to-interior gradients in the mixed hardwood forests of the Roanoke River Basin, North Carolina. *Conservation Biology*, *8*(3), 822–832. [https://doi.](https://doi.org/10.1046/j.1523-1739.1994.08030822.x) [org/10.1046/j.1523-1739.1994.08030822.x](https://doi.org/10.1046/j.1523-1739.1994.08030822.x)
- Gehlhausen, S., Schwartz, M. W., & Augspurger, C. K. (2000). Vegetation and microclimatic edge effects in two mixed-mesophytic forest fragments. *Plant Ecology*, *147*, 21–35.
- Gillooly, J. F., Brown, J. H., West, G. B., Savage, V. M., & Charnov, E. L. (2001). Effects of size and temperature on metabolic rate. *Science*, *293*(5538), 2248–2251.
- Haddad, N. M., Brudvig, L. A., Clobert, J., Davies, K. F., Gonzalez, A., Holt, R. D., ... Townshend, J. R. (2015). Habitat fragmentation and its lasting impact on Earth's ecosystems. *Science Advances*, *1*(2), e1500052. <https://doi.org/10.1126/sciadv.1500052>
- Harrison, S., & Bruna, E. (1999). Habitat fragmentation and large‐scale conservation: What do we know for sure? *Ecography*, *22*(3), 225–232. <https://doi.org/10.1111/j.1600-0587.1999.tb00496.x>
- Hatley, C. L., & Macmahon, J. A. (1980). Spider community organiza‐ tion: Seasonal variation and the role of vegetation architecture. *Environmental Entomology*, *9*(5), 632–639. [https://doi.org/10.1093/](https://doi.org/10.1093/ee/9.5.632) [ee/9.5.632](https://doi.org/10.1093/ee/9.5.632)
- Hedges,L.V.,Gurevitch,J.,&Curtis,P.S.(1999).Themeta‐analysisofresponse ratios in experimental ecology. *Ecology*, *80*(4), 1150–1156. [https://](https://doi.org/10.1890/0012-9658(1999)080[1150:TMAORR]2.0.CO;2) [doi.org/10.1890/0012-9658\(1999\)080\[1150:TMAORR\]2.0.CO;2](https://doi.org/10.1890/0012-9658(1999)080[1150:TMAORR]2.0.CO;2)
- Heithecker, T. D., & Halpern, C. B. (2007). Edge-related gradients in microclimate in forest aggregates following structural retention harvests in western Washington. *Forest Ecology and Management*, *248*(3), 163–173. <https://doi.org/10.1016/j.foreco.2007.05.003>
- Holland, J. M., & Luff, M. L. (2000). The effects of agricultural practices on Carabidae in temperate agroecosystems. *Integrated Pest Management Reviews*, *5*(2), 109–129.
- Homburg, K., Homburg, N., Schäfer, F., Schuldt, A., & Assmann, T. (2013). Carabids.org – A dynamic online database of ground beetle species traits (Coleoptera, Carabidae). *Insect Conservation and Diversity*, *7*, 195–205. <https://doi.org/10.1111/icad.12045>
- Honnay, O., Verheyen, K., & Hermy, M. (2002). Permeability of an‐ cient forest edges for weedy plant species invasion. *Forest Ecology and Management*, *161*(1–3), 109–122. [https://doi.org/10.1016/](https://doi.org/10.1016/S0378-1127(01)00490-X) [S0378-1127\(01\)00490-X](https://doi.org/10.1016/S0378-1127(01)00490-X)
- Hornung, E. (2011). Evolutionary adaptation of oniscidean isopods to ter‐ restrial life: Structure, physiology and behavior. *Terrestrial Arthropod Reviews*, *4*(2), 95–130. <https://doi.org/10.1163/187498311X576262>
- Hothorn, T., Bretz, F., Westfall, P., Heiberger, R. M., Schuetzenmeister, A., & Scheibe, S. (2016). multcomp: Simultaneous inference in general parametric models. R package version 1.4‐6. [https://cran.r-proj‐](https://cran.r-project.org/web/packages/multcomp/) [ect.org/web/packages/multcomp/](https://cran.r-project.org/web/packages/multcomp/)
- Ibisch, P. L., Hoffmann, M. T., Kreft, S., Pe'er, G., Kati, V., Biber‐ Freudenberger, L., … Selva, N. (2016). A global map of roadless areas and their conservation status. *Science*, *354*(6318), 1423–1427.
- Jeanneret, P., Schüpbach, B., Pfiffner, L., & Walter, T. (2003). Arthropod reaction to landscape and habitat features in agricultural landscapes. *Landscape Ecology*, *18*(3), 253–263.
- Jeffery, S., Gardi, C., Jones, A., Montanarella, L., Marmo, L., Miko, L., … van de Putten, W. H. (Eds.) (2010). *European atlas of soil biodiversity*. Luxembourg, Luxembourg: Publications Office of the European Union.
- Kolb, A., Diekmann, M. (2004). Effects of environment, habitat con‐ figuration and forest continuity on the distribution of forest plant species. *Journal of Vegetation Science*, *15*(2), 199–208. [https://doi.](https://doi.org/10.1111/j.1654-1103.2004.tb02255.x) [org/10.1111/j.1654-1103.2004.tb02255.x](https://doi.org/10.1111/j.1654-1103.2004.tb02255.x)
- Kowal, V. A., & Cartar, R. V. (2012). Edge effects of three anthropogenic disturbances on spider communities in Alberta's boreal forest. *Journal of Insect Conservation*, *16*(4), 613–627. [https://doi.org/10.1007/](https://doi.org/10.1007/s10841-011-9446-z) [s10841-011-9446-z](https://doi.org/10.1007/s10841-011-9446-z)
- Krauss, J., Bommarco, R., Guardiola, M., Heikkinen, R. K., Helm, A., Kuussaari, M., … Steffan‐Dewenter, I. (2010). Habitat fragmenta‐ tion causes immediate and time‐delayed biodiversity loss at dif‐ ferent trophic levels. *Ecology Letters*, *13*(5), 597–605. [https://doi.](https://doi.org/10.1111/j.1461-0248.2010.01457.x) [org/10.1111/j.1461-0248.2010.01457.x](https://doi.org/10.1111/j.1461-0248.2010.01457.x)
- Lacasella, F., Gratton, C., De Felici, S., Isaia, M., Zapparoli, M., Marta, S., & Sbordoni, V. (2015). Asymmetrical responses of forest and "beyond edge" arthropod communities across a forest–grassland ecotone. *Biodiversity and Conservation*, *24*(3), 447–465. [https://doi.](https://doi.org/10.1007/s10531-014-0825-0) [org/10.1007/s10531-014-0825-0](https://doi.org/10.1007/s10531-014-0825-0)
- Lavelle, P. (1997). Faunal activities and soil processes: Adaptive strat‐ egies that determine ecosystem function. *Advances in Ecological Research*, *27*, 93–132.
- Madeira, F., Tscharntke, T., Elek, Z., Kormann, U. G., Pons, X., Rösch, V., … Batáry, P. (2016). Spillover of arthropods from cropland to protected calcareous grassland – The neighbouring habitat matters. *Agriculture Ecosystems & Environment*, *235*, 127–133.
- Magura, T. (2002). Carabids and forest edge: Spatial pattern and edge effect. *Forest Ecology and Management*, *157*(1), 23–37. [https://doi.](https://doi.org/10.1016/S0378-1127(00)00654-X) [org/10.1016/S0378-1127\(00\)00654-X](https://doi.org/10.1016/S0378-1127(00)00654-X)
- Marc, P., & Canard, A. (1997). Maintaining spider biodiversity in agroecosystems as a tool inpest control.*Agriculture, Ecosystems & Environment*, *62*(2–3), 229–235. [https://doi.org/10.1016/S0167-8809\(96\)01133-4](https://doi.org/10.1016/S0167-8809(96)01133-4)
- Matlack, G. R. (1993). Microenvironment variation within and among for‐ est edge sites in the eastern United States. *Biological Conservation*, *66*(3), 185–194. [https://doi.org/10.1016/0006-3207\(93\)90004-K](https://doi.org/10.1016/0006-3207(93)90004-K)
- Moe, S. J., Stelzer, R. S., Forman, M. R., Harpole, W. S., Daufresne, T., & Yoshida, T. (2005). Recent advances in ecological stoichiometry: Insights for population and community ecology. *Oikos*, *109*(1), 29–39. <https://doi.org/10.1111/j.0030-1299.2005.14056.x>
- Ng, K., Barton, P. S., Macfadyen, S., Lindenmayer, D. B., & Driscoll, D. A. (2018). Beetle's responses to edges in fragmented landscapes are driven by adjacent farmland use, season and cross‐habitat move‐ ment. *Landscape Ecology*, *33*(1), 109–125. [https://doi.org/10.1007/](https://doi.org/10.1007/s10980-017-0587-7) [s10980-017-0587-7](https://doi.org/10.1007/s10980-017-0587-7)
- Niemelä, J., Langor, D., & Spence, J. R. (1993). Effects of clear‐cut harvest‐ ing on boreal ground‐beetle assemblages (Coleoptera: Carabidae) in Western Canada. *Conservation Biology*, *7*(3), 551–561. [https://doi.](https://doi.org/10.1046/j.1523-1739.1993.07030551.x) [org/10.1046/j.1523-1739.1993.07030551.x](https://doi.org/10.1046/j.1523-1739.1993.07030551.x)
- Pearce, J. L., & Venier, L. A. (2006). The use of ground beetles (Coleoptera: Carabidae) and spiders (Araneae) as bioindicators of sustainable forest management: A review. *Ecological Indicators*, *6*(4), 780–793. <https://doi.org/10.1016/j.ecolind.2005.03.005>
- Peyras, M., Vespa, N. I., Bellocq, M. I., & Zurita, G. A. (2013). Quantifying edge effects: The role of habitat contrast and species specializa‐ tion. *Journal of Insect Conservation*, *17*(4), 807–820. [https://doi.](https://doi.org/10.1007/s10841-013-9563-y) [org/10.1007/s10841-013-9563-y](https://doi.org/10.1007/s10841-013-9563-y)

12 a
b DE SMEDT et al. **Diversity** and **Distributions**

- Pfeifer, M., Lefebvre, V., Peres, C. A., Banks‐Leite, C., Wearn, O. R., Marsh, C. J., … Ewers, R. M. (2017). Creation of forest edges has a global impact on forest vertebrates. *Nature*, *551*(7679), 187. [https://](https://doi.org/10.1038/nature24457) doi.org/10.1038/nature24457
- R Core Team. (2017). *R: A language and environment for statistical computing*. Vienna, Austria: R Foundation for Statistical Computing.
- Rand, T. A., Tylianakis, J. M., & Tscharntke, T. (2006). Spillover edge ef‐ fects: The dispersal of agriculturally subsidized insect natural ene‐ mies into adjacent natural habitats. *EcologyLetters*, *9*(5), 603–614. <https://doi.org/10.1111/j.1461-0248.2006.00911.x>
- Reino, L., Beja, P., Osborne, P. E., Morgado, R., Fabião, A., & Rotenberry, J. T. (2009). Distance to edges, edge contrast and landscape fragmenta‐ tion: Interactions affecting farmland birds around forest plantations. *Biological Conservation*, *142*(4), 824–838. [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.biocon.2008.12.011) [biocon.2008.12.011](https://doi.org/10.1016/j.biocon.2008.12.011)
- Remy, E., Wuyts, K., Verheyen, K., Gundersen, P., & Boeckx, P. (2018). Altered microbial communities and nitrogen availability in temperate forest edges. *Soil Biology and Biochemistry*, *116*, 179–188. [https://doi.](https://doi.org/10.1016/j.soilbio.2017.10.016) [org/10.1016/j.soilbio.2017.10.016](https://doi.org/10.1016/j.soilbio.2017.10.016)
- Ries, L., Fletcher, R. J. Jr, Battin, J., & Sisk, T. D. (2004). Ecological re‐ sponses to habitat edges: Mechanisms, models, and variability explained. *Annual Review of Ecology, Evolution and Systematics*, *35*, 491– 522. <https://doi.org/10.1146/annurev.ecolsys.35.112202.130148>
- Samu, F., & Szinetár, C. (2002). On the nature of agrobiont spiders. *Journal of Arachnology*, *30*(2), 389–402. [https://doi.org/10.1636/01](https://doi.org/10.1636/0161-8202(2002)030[0389:OTNOAS]2.0.CO;2) [61-8202\(2002\)030\[0389:OTNOAS\]2.0.CO;2](https://doi.org/10.1636/0161-8202(2002)030[0389:OTNOAS]2.0.CO;2)
- Soga, M., Kanno, N., Yamaura, Y., & Koike, S. (2013). Patch size deter‐ mines the strength of edge effects on carabid beetle assemblages in urban remnant forests. *Journal of Insect Conservation*, *17*(2), 421–428. <https://doi.org/10.1007/s10841-012-9524-x>
- Symondson, W. O. C., Sunderland, K. D., & Greenstone, M. H. (2002). Can generalist predators be effective biocontrol agents? *Annual Reviews of Entomology*, *47*(1), 561–594.
- Tóthmérész, B., Nagy, D., Mizser, S., Bogyó, D., & Magura, T. (2014). Edge effects on ground‐dwelling beetles (Carabidae and Staphylinidae) in oak forest‐forest edge‐grassland habitats in Hungary. *European Journal of Entomology*, *111*(5), 686. [https://doi.org/10.14411/](https://doi.org/10.14411/eje.2014.091) [eje.2014.091](https://doi.org/10.14411/eje.2014.091)
- Tscharntke, T., Rand, T. A., & Bianchi, F. J. (2005). The landscape context of trophic interactions: Insect spillover across the crop—noncrop in‐ terface. *Annales Zoologici Fennici*, *42*, 421–432.
- Tscharntke, T., Tylianakis, J. M., Rand, T. A., Didham, R. K., Fahrig, L., Batáry, P., … Westphal, C. (2012). Landscape moderation of biodi‐ versity patterns and processes‐eight hypotheses. *Biological Reviews*, *87*(3), 661–685. <https://doi.org/10.1111/j.1469-185X.2011.00216.x>
- Uetz, G. W. (1991). Habitat structure and spider foraging. In S. S. Bell, E. D. McCoy, & H. R. Mushinsky (Eds.), *Habitat structure* (pp. 325–348). Dordrecht, The Netherlands: Springer.
- Valdés, A., Lenoir, J., Gallet‐Moron, E., Andrieu, E., Brunet, J., Chabrerie, O., … Decocq, G. (2015). The contribution of patch‐scale conditions is greater than that of macroclimate in explaining local plant diversity in fragmented forests across Europe. *Global Ecology and Biogeography*, *24*(9), 1094–1105. <https://doi.org/10.1111/geb.12345>
- Verheyen, K., Bossuyt, B., Honnay, O., & Hermy, M. (2003). Herbaceous plant community structure of ancient and recent forests in two contrasting forest types. *Basic and Applied Ecology*, *4*(6), 537–546. <https://doi.org/10.1078/1439-1791-00210>
- Wade, T. G., Riitters, K. H., Wickham, J. D., & Jones, K. B. (2003). Distribution and causes of global forest fragmentation. *Conservation Ecology*, *7*(2), 7. <https://doi.org/10.5751/ES-00530-070207>
- Warburg, M. R., Linsenmair, K. E., & Bercovitz, K. (1984). The effects of climate on the distribution and abundance of isopods. *Symposia of the Zoological Society of London*, *53*, 339–367.
- Woodcock, B. A. (2004). Pitfall trapping in ecological studies. In S. R. Leather (Ed.), *Insect sampling in forest ecosystems. Methods in ecology series* (pp. 37–57). Oxford, UK: Blackwell Science Ltd.
- Yekwayo, I., Pryke, J. S., Roets, F., & Samways, M. J. (2016). Surrounding vegetation matters for arthropods of small, natural patches of indige‐ nous forest. *Insect Conservation and Diversity*, *9*(3), 224–235. [https://](https://doi.org/10.1111/icad.12160) doi.org/10.1111/icad.12160

BIOSKETCH

Pallieter De Smedt is a postdoctoral research fellow at the Forest & Nature Lab of Ghent University. He has special interest in soil biodiversity, their distribution and ecology. His research focusses on the importance of soil invertebrates for ecosystem function‐ ing in small forest fragments. Most authors are members of the research project smallFOREST ([http://www.u-picardie.fr/small‐](http://www.u-picardie.fr/smallforest/uk/) [forest/uk/](http://www.u-picardie.fr/smallforest/uk/)), studying biodiversity and ecosystem services of small forest fragments in agricultural landscapes in Europe.

Author contributions: All authors conceived the ideas and col‐ lected the data; P.D.S. and L.B. analysed the data; and P.D.S. led the writing.

SUPPORTING INFORMATION

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

How to cite this article: De Smedt P, Baeten L, Proesmans W, et al. Strength of forest edge effects on litter‐dwelling macro‐ arthropods across Europe is influenced by forest age and edge properties. *Divers Distrib*. 2019;00:1–12. [https://doi.](https://doi.org/10.1111/ddi.12909) [org/10.1111/ddi.12909](https://doi.org/10.1111/ddi.12909)