



Adsorption and degradation of the herbicide nicosulfuron in a stagnic Luvisol and Vermic Umbrisol cultivated under conventional or conservation agriculture

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Abstract

The main goals of conservation agriculture are to enhance soil fertility and reduce soil degradation, especially through erosion. However, conservation agriculture practices can increase the risk of contamination by pesticides, mainly through vertical transfer via water flow. Better understanding of their sorption and degradation processes is thus needed in conservation agriculture as they control the amount of pesticide available for vertical transfer. The purpose of our study was to investigate the sorption and degradation processes of nicosulfuron in soil profiles (up to 90 cm deep) of a Vermic Umbrisol and a Stagnic Luvisol managed either in conventional or in conservation agriculture. Two laboratory sorption and incubation experiments were performed. Low sorption was observed regardless of the soil type, agricultural management or depth, with a maximum value of $1.3 \pm 2.0 \text{ L kg}^{-1}$. By the end of the experiment (91 days), nicosulfuron mineralisation in the Vermic Umbrisol was similar for the two types of agricultural management and rather depended on soil depth ($29.0 \pm 2.3\%$ in the 0–60-cm layers against $7.5 \pm 1.4\%$ in the 60–90 cm). In the Stagnic Luvisol, nicosulfuron mineralisation reached similar value in every layer of the conservation agriculture plot ($26.5\% \pm 0.7\%$). On the conventional tillage plot, mineralisation decreased in the deepest layer (25–60 cm) reaching only $18.4 \pm 6.9\%$ of the applied nicosulfuron. Regardless of the soil type or agricultural management, non-extractable residue formation was identified as the main dissipation process of nicosulfuron ($45.1 \pm 8.5\%$ and $50.2 \pm 7.0\%$ under conventional and conservation agriculture respectively after 91 days). In our study, nicosulfuron behaved similarly in the Vermic Umbrisol regardless of the agricultural management, whereas the risk of transfer to groundwater seemed lower in the Stagnic Luvisol under conservation agriculture.

Keywords Pesticide · Mineralisation · Soil organic carbon · Agricultural management · Sorption

Introduction

To enhance the sustainability of agricultural systems and decrease the impacts of conventional agriculture, alternative cultivation practices such as conservation agriculture have been

developed. Conservation agriculture is defined as a combination of three main interconnected soil-conservation techniques: (i) little or no soil disturbance, (ii) permanent soil cover by crop residues and/or cover crops and (iii) diversification of plant species in the crop rotation (FAO 2016). The main goals of conservation agriculture are to reduce soil degradation due to erosion but also the contamination of surface water by runoff or transfer of sorbed agrochemicals on eroded soil particles (Holland 2004; Hobbs et al. 2008).

Compared with conventional agriculture that includes soil tillage (sometimes with inversion of topsoil layers), conservation agriculture enhances faunal activity, especially earthworm activity, which results in a dense biopore network (Shipitalo et al. 2000). Moreover, because the topsoil is disturbed less, it usually contains a greater continuity of vertically oriented macropores, which leads to higher hydraulic conductivity than that of tilled soils (Wahl et al. 2004). Increasing

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vertical transfer is one way to reduce runoff and thus surface water contamination, but in return, it increases the risk of deep vertical transfer to groundwater. This is a major drawback of conservation agriculture, especially as reducing tillage intensity generally increases weed pressure, which often leads to an increase in herbicide use (Alletto et al. 2010). Better understanding of sorption and degradation processes in such agricultural systems is thus critically needed as they control the amount of pesticide available for vertical transfer.

One of the main changes of not inverting topsoil layers, as in conservation agriculture, is the accumulation of dead residues or mulch at the soil surface, which enriches the first few centimetres of soil in organic carbon. The carbon content then decreases with soil depth, reaching contents less than or equal to those of conventionally tilled soil (Doran 1980; Kay and VandenBygaert 2002). For many pesticides, sorption can be related to soil organic matter content (Gaston et al. 2007; Đurović et al. 2009; Sadegh-Zadeh et al. 2017). Thus, at an equivalent depth in the topsoil, pesticide sorption is generally higher under conservation agriculture than conventional agriculture because of the former's higher carbon content (Gaston et al. 2003; Locke et al. 2005). Deeper in the soil, sorption decreases because the carbon content is lower, and it tends to follow the decrease in carbon content (Clay and Koskinen 2003). In addition, the accumulation of organic matter (and fertilisers) under conservation agriculture generally acidifies the topsoil (Thompson and Whitney 2000; Thomas et al. 2007; Obour et al. 2017). Weak acids and ionisable molecules dissociate to their anionic form as soil pH increases, which lowers adsorption (Dyson et al. 2002). More acidic conditions in the topsoil may then increase pesticide adsorption under conservation agriculture. For instance, Alletto et al. (2013) observed a negative correlation between pH and S-metolachlor sorption in soils under conservation agriculture.

Because of soil organic matter accumulation and less soil disturbance by agricultural operations, microbial biomass and microbial activity in the topsoil is generally greater under conservation agriculture than conventional agriculture (Kandeler et al. 1999). This increase in microbial activity can sometimes increase pesticide degradation (Locke et al. 2005; Zablutowicz et al. 2007), but pesticide degradation does not always reflect soil microbial activity (Mahía et al. 2007; Okada et al. 2019). Decrease in substrate availability due to higher adsorption is often mentioned as an explanation for low degradation under conservation agriculture (Zablutowicz et al. 2000). Adaptation of the microbial community can also increase pesticide degradation (Barriuso et al. 1996). Both may explain the lack of correlation between total microbial activity and pesticide mineralisation (Alletto et al. 2013). On the other hand, microbial activity and biomass are often correlated with soil pH, which strongly influences pesticide degradation in soils (Kah et al. 2007). Acidification of the topsoil under conservation agriculture may thus modify pesticide

degradation. In the subsoil (> 25–30 cm deep), pesticide degradation is generally lower, generally due to less microbial activity and/or biomass related to lower carbon content, temperature and oxygenation (Fomsgaard 1995; Bending and Rodríguez-Cruz 2007).

We studied the sorption and degradation of nicosulfuron in soil profiles (up to 90 cm deep) of two soils, a Stagnic Luvisol and a Vermic Umbrisol, managed for more than 10 years under two types of agricultural management: conventional agriculture (tillage, bare soil and monoculture) and conservation agriculture (no tillage, cover crops and crop rotations). Nicosulfuron is a post-emergence herbicide used to control grass species in maize. It is considered environmentally safe because of low application rates (usually < 100 g ha⁻¹) (Regitano and Koskinen 2008). Nicosulfuron is however generally in anionic form in soils and is considered to be highly mobile, which could result in groundwater contamination (Gonzalez and Ukrainczyk 1996). In addition, nicosulfuron generally has low sorption and fast degradation rates (Olivera et al. 2001; Poppell et al. 2002; Azcarate et al. 2018; PPDB 2020). Recently, Cueff et al. (2020) reported high nicosulfuron leaching and a high degree of preferential flow in undisturbed columns sampled in the same soils as in the present study.

Materials and methods

Sites and types of agricultural management

The two sites of this study are located in south-western France, in the Pyrénées-Atlantiques (43.516532 N, – 0.260186 W) and Gers (43.410980 N, 0.284574 E) departments. Both sites are subjected to an altered oceanic climate.

The soil at the Pyrénées-Atlantiques site is a thick humic acid soil developed from Quaternary silty alluvial deposits that is classified as a Vermic Umbrisol according to the World Reference Base for Soil Resources (IUSS Working Group WRB 2007) and as a veracrisol according to the French Soil Classification (Référentiel Pédologique 2008) (locally called “Touyas”). It has a rich, deep (50–80 cm deep) and well-structured organic horizon (Table 1). Its texture is composed mainly of fine silt (> 450–500 g kg⁻¹), with a low proportion of sand (< 100–120 g kg⁻¹). Clay minerals are mainly kaolinite, illite and vermiculite (Arrouays et al. 1992). It has a high agronomic potential, especially for maize (mean yield ≈ 13–15 t ha⁻¹) and soya bean (mean yield ≈ 3.7–4.2 t ha⁻¹) (most not irrigated). Hereafter, this site is referred to as the VER (veracrisol) site.

The soil at the Gers site is classified as a Stagnic Luvisol (IUSS Working Group WRB 2007) and a luvisol redoxisol (Référentiel Pédologique 2008) (locally called “Boulbènes”), with a loamy topsoil layer (fine + coarse silt ≈ 450–500 g

Table 1 Sand, silt and clay content and soil pH. Values correspond to the mean of the three spatial replicates \pm standard deviation

Site	Plot	Depth cm	Sand content g kg ⁻¹	Silt content g kg ⁻¹	Clay content g kg ⁻¹	Soil pH
VER	TILL	0–10	118 \pm 10	718 \pm 21	164 \pm 12	6.5 \pm 0.3
		10–25	110 \pm 11	732 \pm 11	158 \pm 12	6.8 \pm 0.3
		25–60	112 \pm 17	727 \pm 23	161 \pm 8	6.9 \pm 0.7
		60–90	113 \pm 6	707 \pm 23	179 \pm 22	6.5 \pm 1.4
	CONS	0–10	124 \pm 2	720 \pm 10	156 \pm 8	6.3 \pm 0.0
		10–25	121 \pm 3	724 \pm 14	155 \pm 11	6.4 \pm 0.1
		25–60	121 \pm 7	724 \pm 14	156 \pm 10	6.3 \pm 0.1
		60–90	116 \pm 4	694 \pm 14	190 \pm 10	5.2 \pm 0.1
LUV	TILL	0–10	273 \pm 3	604 \pm 17	123 \pm 14	6.8 \pm 0.2
		10–25	287 \pm 17	593 \pm 26	121 \pm 9	6.8 \pm 0.2
		25–60	226 \pm 40	607 \pm 35	167 \pm 10	7.1 \pm 0.1
	CONS	0–10	241 \pm 32	596 \pm 44	164 \pm 22	5.8 \pm 0.4
		10–25	233 \pm 34	587 \pm 25	180 \pm 11	6.0 \pm 0.4
		25–60	199 \pm 36	601 \pm 79	200 \pm 46	6.5 \pm 0.3

kg⁻¹) and an illuvial clay horizon that appears at ca. 45–55 cm (Table 1). Clay minerals are mainly illite and smectite (Michel et al. 2003). The substratum is an alluvial pebbly layer with low permeability that appears at ca. 60–80 cm. With low organic carbon contents (usually < 7–8 g kg⁻¹) in the topsoil and high silt contents, this soil is strongly sensitive to crusting. Because of these properties (especially hydromorphic conditions at low depth), it has a moderate agronomic potential for mainly irrigated crops (mean maize yields \approx 10–12 t ha⁻¹). Hereafter, this site is referred to as the LUV (luvisol) site.

Each site consists of two adjacent plots managed under conventional (TILL) and conservation (CONS) practices. The conventional plots have been cultivated under maize monoculture for more than 40 years, and the soil is left bare in winter. Tillage depth with a mouldboard plough is ca. 22–23 cm at the VER site and 30 cm at the LUV site. In both conservation plots (VER and LUV), agricultural management consists of (i) no-tillage practices, (ii) permanent soil surface cover by mulch and cover crop mixtures (phacelia and faba bean at the VER site, two cereals and faba bean at the LUV site) sown during fallow periods and (iii) a diversified crop rotation (maize, soya bean, cereals and sometimes intercrops of cereals and legumes). These conservation agriculture practices have been performed since 2006 and 2000 at the VER and LUV sites, respectively.

Soil sampling and conditioning

In each plot, bulk soil was sampled from soil profiles collected at three locations (considered replicates) at ca. 20-m intervals along a transect parallel to the border between the TILL and CONS plots. To minimise the spatial variability in soil properties between the two plots as much as possible, the transects

lay 20 m from the border between the plots. Four soil layers (0–10, 10–25, 25–60 and 60–90 cm) and three soil layers (0–10, 10–25 and 25–60 cm) were collected in each profile at the VER and LUV site, respectively. Soils were air-dried and sieved at 5 mm and stored in a cold chamber (4 °C) before analysis.

Experimental study

Soil organic carbon and carbon stock

Carbon content was measured according to the NF ISO 10694 standard (AFNOR 1995) based on CO₂ emission of the sample following dry combustion. Results were corrected for carbonates present in the sample.

In each soil profile, we used 250 cm³ (8 cm diameter, 5 cm high) cylindrical cores to collect undisturbed soil samples by hand using gentle pressure. Soil cores were dried in an oven (105 °C, 48 h), and bulk density (g cm⁻³) was determined to estimate the carbon stock in each soil layer.

Sorption experiments

Few studies about nicosulfuron sorption are available in the literature. In most cases, Freundlich exponent was found to be close to 1. Four studies covering different soil types and textures (including similar textures as soils from our study) reported n values ranging mainly between 0.9 and 1.1 (only three n values were higher than 1.1, with a maximum of 1.4, on the 33 soils studied in the literature) (Gonzalez and Ukrainczyk 1996; EFSA 2008; Regitano and Koskinen 2008; Caceres-Jensen et al. 2020). Azcarate et al. (2015) reported no effect of the initial solution concentration and

assimilated the sorption coefficient to a K_d . In light of the results from the literature, only one initial concentration was applied in our study in order to determine a K_d .

Nicosulfuron (98.4 % purity) was purchased from Cluzeau Info Labo (Ste Foy, France). Batch sorption experiments were conducted with a 1:3 soil/solution ratio (m/v) with a solution of nicosulfuron concentrated at $19.8 \mu\text{g L}^{-1}$. Nicosulfuron concentration was determined by ultra-performance liquid chromatography coupled with tandem mass spectrometry (UHPLC-MS/MS Acquity-TQD, Waters Corp., Milford, MA, USA) after 24-h equilibrium. A detailed measurement protocol was published by Cueff et al. (2020). Limit of quantification (LOQ) was determined in blank soil extracts spiked with low concentrations of nicosulfuron, as the concentration that led to a signal to noise ratio of 10. Limit of detection (LOD) was calculated from LOQ as $\text{LOD} = \text{LOQ} / 3$. For nicosulfuron, LOD and LOQ were equal to 0.016 and 0.05 $\mu\text{g/L}$, respectively.

Incubation experiments

A soil sample (10 g dry weight) from each soil layer was placed in a Falcon tube (50 mL, Corning, NY, USA) and adjusted to 80% of soil field capacity (-33 kPa), which corresponds to $\text{pF } 2.5$ (with $\text{pF} = \log_{10}|h|$, h in cm). For each soil sample, water content at field capacity was determined using a pressure plate (Klute 1986). Each Falcon tube was placed in a hermetically sealed incubation jar that was large enough to prevent formation of anoxic conditions in the soil. Then, 0.5 mL of a solution of ^{14}C -nicosulfuron with a mean activity of 609,235 DPM/mL was added to the soil. In addition, a tube of 10 mL of water (to maintain constant relative humidity in the jar) and a tube of 6 mL of 1 N sodium hydroxide (NaOH) solution were placed into the jar. The NaOH solution trapped all the CO_2 ($^{12}\text{C}\text{-CO}_2$ and $^{14}\text{C}\text{-CO}_2$) released by the mineralisation. The NaOH trap was replaced periodically until the end of the experiment to avoid saturating it. Degradation of ^{14}C -nicosulfuron was monitored for 91 days, in the dark, at $28 \pm 0.5 \text{ }^\circ\text{C}$. Triplicate samples for each soil layer of each site were run for five incubation times: 0, 7, 14, 35, and 91 days.

The distribution of ^{14}C -nicosulfuron activity among the mineralised, water-extractable, methanol-extractable, and non-extractable-residue fractions was determined at the five incubation times. The mineralised fraction and total CO_2 released were also measured in all of the intermediate NaOH solutions. The water-extractable fraction of ^{14}C was determined by CaCl_2 extraction (30 mL, 10^{-2} M). Then, the methanol-extractable fraction of ^{14}C was determined by two successive methanol extractions (30 mL). For each extraction, the tubes were left for 22 h in an end-over-end shaker and then centrifuged at $14,000 \times g$ for 10 min. The ^{14}C activity was measured by adding 10 mL of a scintillating liquid (Ultima Gold XR, Perkin Elmer, Waltham, MA, USA) to a 1-mL

aliquot of the aqueous phase of all three successive extractions. The non-extractable-residue fraction of ^{14}C was determined after drying and grinding the soil remaining after the extractions. The remaining ^{14}C activity was measured by combusting the ground soil in an oxidizer (Biological Oxidizer, OX 700, Zinsser Analytic, Frankfurt, Germany). $^{14}\text{C}\text{-CO}_2$ released from the combustion ($900 \text{ }^\circ\text{C}$, 5 min) of the soil samples was trapped in Oxysolve 400 solution (Zinsser Analytic), in which the ^{14}C activity was directly measured. The mineralised fraction of ^{14}C was determined by adding 10 mL of scintillating liquid to a 2-mL aliquot of the NaOH trap solution. ^{14}C activity of each fraction was measured by liquid scintillation counting for 10 min in a Tri-Carb 2100TR scintillation counter (Packard Instruments, Meriden, CT, USA).

In addition, mineralisation of soil organic carbon was determined by measuring the $^{12}\text{C}\text{-CO}_2$ trapped by the NaOH using a colorimetric method (Colorimetric Skalar Analyzer, Breda, Netherlands). Microbial biomass was determined by the fumigation-extraction method (Wu et al. 1990) using a total organic carbon analyser (Shimadzu TOC 5050A).

Statistical analysis

Cumulative soil carbon mineralisation ($^{12}\text{C}\text{-CO}_2$) and cumulative nicosulfuron mineralisation ($^{14}\text{C}\text{-CO}_2$) for each soil sample were fitted to an exponential model that increases to a maximum:

$$y_{OC} = a_{OC}(1 - e^{-b_{OC}t}) \tag{1}$$

$$y_N = a_N(1 - e^{-b_Nt}) \tag{2}$$

where, respectively, y_{OC} and y_N are the modelled mineralisation of $^{12}\text{C}\text{-CO}_2$ and $^{14}\text{C}\text{-CO}_2$ (expressed as % of initial organic carbon and of applied nicosulfuron), a_{OC} and a_N are the final cumulative mineralisation (%), b_{OC} and b_N are the first-order rate constant (days^{-1}) and t is time (days).

The decrease in the water-extractable fraction of ^{14}C over time was fitted to a first-order kinetics model:

$$y_{WE} = a_{WE}e^{-b_{WE}t} \tag{3}$$

where y_{WE} is the water-extractable fraction modelled at time t (%), a_{WE} is the initial water-extractable fraction measured at t_0 (%) and b_{WE} is the first-order rate constant (days^{-1}).

The increase in the non-extractable-residue (NER) fraction of ^{14}C over time was fitted to an exponential model that increases to a maximum:

$$y_{NER} = y_0 + a_{NER}(1 - e^{-b_{NER}t}) \tag{4}$$

where y_{NER} is the NER fraction modelled at time t (%), y_0 is the initial NER fraction measured at t_0 (%), a_{NER} is the final cumulative percentage of the NER fraction (%) and b_{NER} is the

first-order rate constant (days^{-1}). As the methanol-extractable fraction did not follow any particular dynamics, it was not fitted to a model.

Spearman's correlation analysis was performed in order to explore the relationships between soil properties, sorption coefficients, ^{14}C fractions and model parameters for the two sites and the two agricultural managements. Correlation analysis and correlation matrix were done using the "psych" and "corrplot" R packages.

Analysis of variance (ANOVA) was performed for carbon stock, microbial biomass, K_d and all parameters fitted to models to identify effects of agricultural management and soil depth. When possible, an effect of an interaction between agricultural management and soil depth was evaluated. The Shapiro-Wilk normality test and Levene's homoscedasticity test were applied to the residues of the ANOVA to verify application conditions. When the conditions were not met, logarithmic or square root transformations were applied. When transformed data could not meet the conditions, the Kruskal-Wallis test was performed.

Results

Soil organic carbon content and mineralisation

At both sites, mean organic carbon content did not differ significantly between the two surface layers (0–10 and 10–25 cm) and decreased with depth (Table 2). Nevertheless, in the CONS plot at the LUV site, mean organic carbon content was 31% higher in the 0–10-cm layer than in the 10–25-cm layer. Moreover, mean organic carbon content was generally slightly higher throughout the soil profile in the CONS plot than in the TILL plot at the LUV site and was highest in the two

surface layers in both the CONS and TILL plots at the VER site. Differences in organic carbon content according to agricultural management were significant only at the LUV site ($P < 0.01$).

Carbon stock increased with depth in both plots but varied more at the VER site. Carbon stock was similar for the two types of agricultural management at the VER site, whereas it was higher in the CONS plot than in the TILL plot (especially for the 0–10-cm and 25–60-cm layers) at the LUV site ($P < 0.01$).

Microbial biomass decreased strongly with depth in both plots (Table 2), in relation to the decrease in organic carbon content ($r = 0.71$; Fig. 1). It was also negatively correlated with soil pH ($r = -0.50$). Microbial biomass was higher in the CONS plot than in the TILL plot at both sites ($P < 0.01$) due to the high microbial biomass in the 0–10-cm layer of the CONS plots (570 ± 29 and 710 ± 98 mg C kg^{-1} dry soil at the VER and LUV site, respectively).

Cumulative organic carbon mineralisation ($^{12}\text{C}-\text{CO}_2$ t_{91} ; Fig. 1) was positively correlated with microbial biomass ($r = 0.48$). At each site, it was higher in the 0–10-cm layer of the CONS plot than in any other layer of either plot (Fig. 2). The difference was larger at the LUV site, with a_{OC} reaching up to $20.1 \pm 8.5\%$, while it reached only $8.1 \pm 1.4\%$ at the VER site (Table 3). Mineralisation data fit the exponential model well (Eq. 1) for all soil layers in both plots ($R^2 = 0.96 \pm 0.16$), and a_{OC} generally decreased with depth, except for the 25–60-cm layer of the CONS plot at the VER site. Both agricultural management ($P < 0.01$) and depth ($P < 0.05$) had a significant effect on a_{OC} at the LUV site, whereas only agricultural management had a significant effect ($P < 0.05$) at the VER site. The first-order rate constant was highest in the deepest layer in both plots at the VER site (Table 3). In the upper three layers, b_{OC} had similar values in the TILL plot, but b_{OC} was highest

Table 2 Organic carbon content, carbon stock, batch adsorption coefficient, K_d , organic-carbon adsorption coefficient, K_{oc} and microbial biomass in each soil layer of the different soil profiles. Values correspond to the mean of the three spatial replicates \pm standard deviation.

Site	Plot	Depth cm	Organic carbon g kg^{-1}	Carbon stock t ha^{-1}	K_d L kg^{-1}	K_{oc} L kg^{-1} OC	Microbial biomass mg C kg^{-1} dry soil
VER	TILL	0–10	17.4 ± 2.2	22.9 ± 3.5	0.6 ± 0.1	34 ± 11	215 ± 18
		10–25	17.4 ± 2.0	35.7 ± 3.9	0.6 ± 0.1	33 ± 8	188 ± 23
		25–60	9.7 ± 0.7	49.7 ± 3.1	0.8 ± 0.2	85 ± 27	77 ± 13
		60–90	4.0 ± 0.8	–	1.1 ± 0.1	279 ± 62	13 ± 9
	CONS	0–10	19.0 ± 0.6	24.6 ± 1.4	1.0 ± 0.3	54 ± 16	570 ± 29
		10–25	18.4 ± 0.5	37.5 ± 2.2	0.6 ± 0.1	31 ± 8	190 ± 34
		25–60	9.7 ± 0.8	49.0 ± 5.8	0.5 ± 0.6	46 ± 58	93 ± 25
		60–90	3.8 ± 0.6	–	0.9 ± 0.3	235 ± 150	34 ± 5
LUV	TILL	0–10	7.5 ± 1.7	11.4 ± 2.7	1.3 ± 2.0	186 ± 297	143 ± 21
		10–25	7.6 ± 1.6	17.6 ± 4.2	0.2 ± 0.1	28 ± 13	132 ± 17
		25–60	3.2 ± 1.5	18.3 ± 8.5	0.7 ± 0.3	231 ± 145	19 ± 11
		60–90	3.8 ± 0.6	–	0.9 ± 0.3	235 ± 150	34 ± 5
	CONS	0–10	11.4 ± 1.8	16.6 ± 2.1	0.5 ± 0.3	41 ± 27	710 ± 98
		10–25	8.7 ± 1.1	20.8 ± 2.4	0.8 ± 0.2	100 ± 30	217 ± 81
		25–60	5.3 ± 0.5	30.1 ± 3.1	0.1 ± 0.2	31 ± 36	41 ± 24
		60–90	4.0 ± 0.8	–	1.1 ± 0.1	279 ± 62	13 ± 9

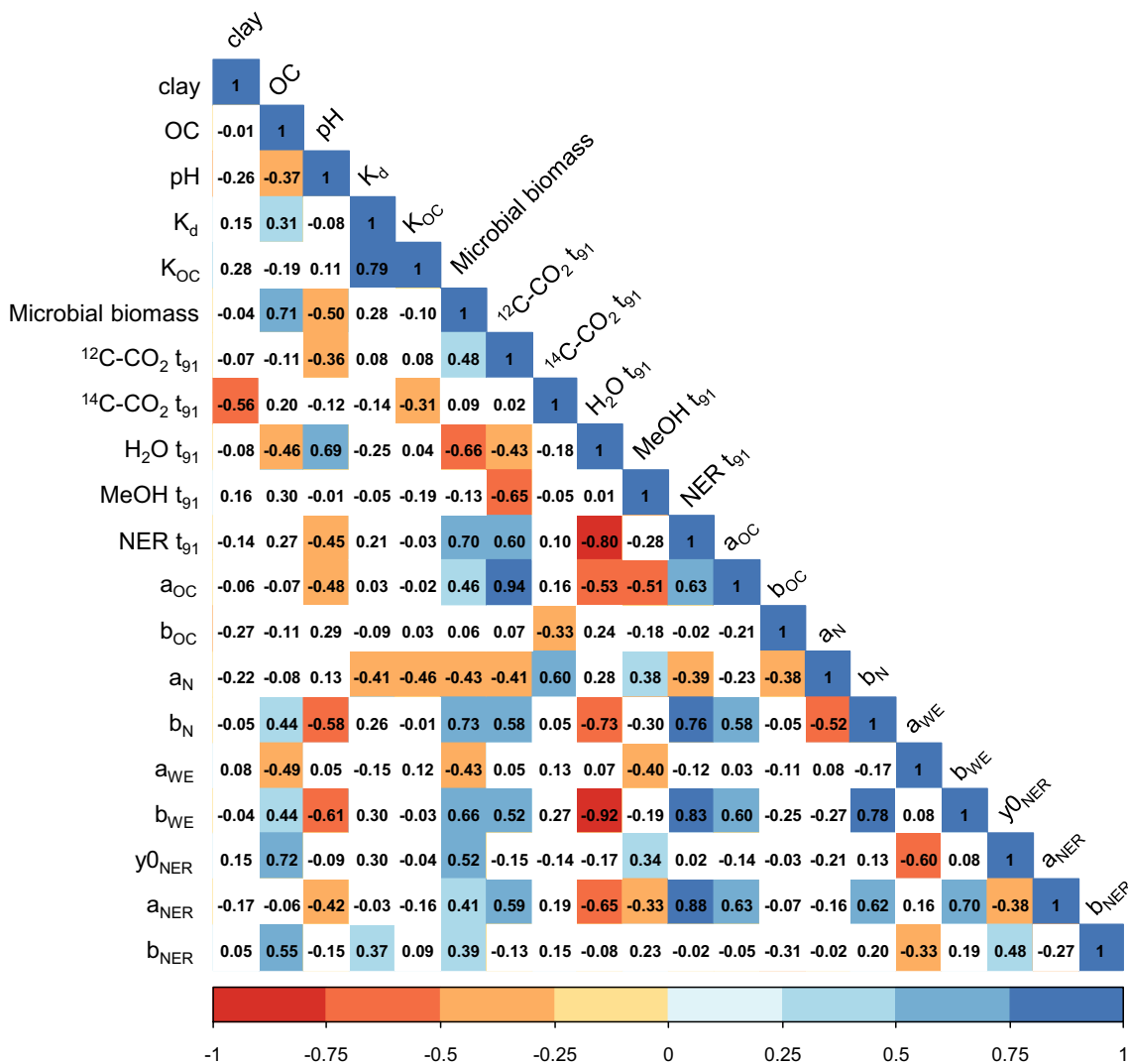


Fig. 1 Correlation between soil properties, sorption coefficients, ^{14}C fractions and model parameters for the two sites and the two agricultural managements

in the 0–10-cm layer in the CONS plot. At the LUV site, b_{OC} was highest in the 25–60-cm layer in the TILL plot and in the 0–10-cm layer in the CONS plot. Only depth had a significant effect on b_{OC} at the VER site ($P < 0.01$) and the LUV site ($P < 0.05$).

Sorption experiments

Sorption of nicosulfuron was low at both sites (mean of $0.7 \pm 0.6 \text{ L kg}^{-1}$). K_d was highest ($1.3 \pm 2.0 \text{ L kg}^{-1}$) in the surface layer (0–10 cm) of the TILL plot at the LUV site (Table 2). With similar values of nicosulfuron sorption of between surface and deep layers, no clear trend with depth was observed, even though organic carbon content differed significantly among soil layers (Table 2). These results suggest that nicosulfuron has moderate affinity for organic carbon, which is confirmed by the K_{OC} values, which were generally highest in the deepest layer, except in the CONS plot at the LUV site

(Table 2). K_{OC} values of many pesticides are commonly higher in deep layers, even for pesticides with stronger affinity for organic carbon. K_d values did not differ significantly by site, agricultural management or depth, but K_{OC} values differed significantly by depth ($P < 0.001$), indicating the difference between the 60–90-cm layer and all other layers at the VER site.

Incubation experiments

VER site

At the VER site, a mean of $96.5 \pm 3.2\%$ of the initially applied ^{14}C -nicosulfuron was recovered after quantifying the mineralised, water-extractable, methanol-extractable and non-extractable-residue fractions of ^{14}C for the five incubation durations. At t_0 , most of the radioactivity was found in the water-extractable fraction (mean of $76.1 \pm 5.9\%$ of applied

Table 3 Values of model parameters used to describe cumulative soil carbon and nicosulfuron mineralisation and the water-extractable and non-extractable-residue (NER) fractions during the incubation experiment. Values correspond to the mean of the three spatial replicates \pm standard deviation.

Site	Plot	Depth	Carbon mineralisation		Nicosulfuron mineralisation		Water-extractable fraction		NER fraction		
			$y_{OC} = a_{OC}(1 - e^{-b_{OC}t})$		$y_N = a_N(1 - e^{-b_Nt})$		$y_{WE} = a_{WE}e^{-b_{WE}t}$		$y_{NER} = y_0 + a_{NER}(1 - e^{-b_{NER}t})$		
		m	a_{OC} %	b_{OC} days ⁻¹	a_N %	b_N days ⁻¹	a_{WE} %	b_{WE} days ⁻¹	y_0 %	a_{NER} %	b_{NER} days ⁻¹
VER	TILL	0–10	4.3 \pm 0.2	0.019 \pm 0.002	35.0 \pm 3.5	0.019 \pm 0.001	66.8 \pm 35	0.030 \pm 0.012	17.4 \pm 0.3	28.2 \pm 2.5	0.104 \pm 0.011
		10–25	3.2 \pm 0.4	0.021 \pm 0.002	33.1 \pm 0.5	0.019 \pm 0.002	69.5 \pm 5.3	0.025 \pm 0.009	18.0 \pm 0.1	26.6 \pm 2.4	0.099 \pm 0.031
		25–60	3.4 \pm 0.3	0.018 \pm 0.003	33.3 \pm 0.7	0.019 \pm 0.002	72.7 \pm 5.6	0.022 \pm 0.004	16.5 \pm 1.6	22.5 \pm 1.9	0.098 \pm 0.006
		60–90	2.8 \pm 0.2	0.030 \pm 0.000	–	–	67.2 \pm 2.5	0.078 \pm 0.008	19.1 \pm 2.4	31.3 \pm 2.0	0.150 \pm 0.091
	CONS	0–10	8.1 \pm 1.4	0.025 \pm 0.006	29.0 \pm 2.1	0.027 \pm 0.003	61.9 \pm 5.3	0.056 \pm 0.012	17.0 \pm 2.2	37.5 \pm 4.9	0.086 \pm 0.023
		10–25	4.0 \pm 0.6	0.018 \pm 0.002	33.7 \pm 2.8	0.024 \pm 0.001	74.7 \pm 7.9	0.042 \pm 0.008	14.6 \pm 0.9	35.0 \pm 5.3	0.072 \pm 0.007
		25–60	7.2 \pm 5.2	0.012 \pm 0.004	47.4 \pm 10.7	0.015 \pm 0.006	74.3 \pm 4.9	0.027 \pm 0.007	13.0 \pm 1.2	28.3 \pm 4.4	0.091 \pm 0.031
		60–90	3.0 \pm 0.9	0.046 \pm 0.018	–	–	78.2 \pm 7.1	0.078 \pm 0.037	11.5 \pm 1.2	34.1 \pm 11.3	0.104 \pm 0.017
LUV	TILL	0–10	7.1 \pm 0.5	0.021 \pm 0.005	36.3 \pm 2.5	0.020 \pm 0.003	82.3 \pm 2.4	0.037 \pm 0.008	10.6 \pm 0.4	38.7 \pm 1.1	0.070 \pm 0.005
		10–25	6.2 \pm 1.0	0.022 \pm 0.003	33.9 \pm 0.9	0.023 \pm 0.002	75.2 \pm .91	0.030 \pm 0.013	11.4 \pm 0.6	38.2 \pm 3.4	0.057 \pm 0.007
		25–60	3.7 \pm 0.2	0.040 \pm 0.014	20.6 \pm 7.4	0.009 \pm 0.006	83.5 \pm 4.5	0.008 \pm 0.001	12.1 \pm 1.2	17.2 \pm 4.3	0.051 \pm 0.028
	CONS	0–10	20.1 \pm 8.5	0.030 \pm 0.000	29.6 \pm 0.6	0.027 \pm 0.001	74.1 \pm 9.4	0.061 \pm 0.032	15.9 \pm 2.2	37.4 \pm 4.8	0.086 \pm 0.014
		10–25	8.2 \pm 0.4	0.021 \pm 0.003	27.1 \pm 2.5	0.031 \pm 0.001	76.5 \pm 10.0	0.082 \pm 0.039	12.2 \pm 3.0	48.1 \pm 4.2	0.074 \pm 0.033
		25–60	4.6 \pm 2.4	0.018 \pm 0.001	45.8 \pm 9.5	0.011 \pm 0.004	78.9 \pm 5.0	0.016 \pm 0.006	10.9 \pm .31	31.7 \pm 5.2	0.038 \pm 0.014

¹⁴C). The rest of the radioactivity was distributed between the methanol-extractable (9.1 \pm 4.7%) and non-extractable-residue fractions (15.5 \pm 2.6%) as mineralisation had not yet started. In most layers, the non-extractable-residue fraction of ¹⁴C was 2–3 times as large as the methanol-extractable fraction at t_0 , except in the 0–10-cm layer of the CONS plot, in which these fractions were similar (17.0 \pm 1.9% and 19.9 \pm 6.3%, respectively). The methanol-extractable fraction in this layer was the largest found at t_0 at the VER site.

¹⁴C-nicosulfuron started to mineralise within 3 days after it was applied. Cumulative mineralisation increased throughout the experiment and fit the exponential model well (Eq. 2) ($R^2 = 0.99 \pm 0.00$ for the upper three soil layers (0–60 cm). In the deepest layer (60–90 cm), nicosulfuron mineralisation was low, and the observed ¹⁴CO₂ dynamics did not fit the exponential model well (a linear model was used instead). After 91 days, mean cumulative nicosulfuron mineralisation differed significantly ($P < 0.001$) between the upper three layers (ca. 29.0 \pm 2.3%) and the deepest layer (60–90 cm) (7.5 \pm 1.4%) in both the CONS and TILL plots (Fig. 3). While nicosulfuron mineralisation decreased progressively (but non-significantly) with depth (from 0 to 60 cm) in the TILL plot (Fig. 3), it ultimately followed the reverse order of depth in the CONS plot, with the highest cumulative mineralisation observed after 50 days in the 25–60-cm layer. In the TILL plot, model parameters indicated a trend similar to that for measured data, with equivalent values of the first-order rate constant b_N

(Table 3). In the CONS plot, however, the modelled final cumulative mineralisation a_N in the 25–60-cm layer was higher than the measured data (Table 3; Fig. 3), indicating that nicosulfuron in this soil horizon may continue to mineralise beyond the 91-day incubation period of the experiment. The first-order rate constant followed a similar trend and was much lower in the 25–60-cm layer of the CONS plot than in the other layers.

The measured water-extractable fraction fit the first-order kinetics model well (Eq. 3) ($R^2 = 0.91 \pm 0.04$) (Fig. 4a). The initial water-extractable fraction a_{WE} was slightly (but not significantly) higher in the CONS plot. It then decreased strongly during the experiment, especially during the first 7 days (decrease of 22–45 percentage points). The decrease was much stronger in the 60–90-cm layer, which had a much higher first-order rate constant k_{WE} than the upper three layers in both plots ($P < 0.001$). This resulted in a small amount of ¹⁴C-nicosulfuron remaining in the water-extractable fraction at the end of the experiment, which the model always underestimated (e.g. estimating 0% when the measured value was 7–16%).

The methanol-extractable fraction varied little in the upper three layers (0–60 cm) in the TILL plot but decreased strongly in the 0–10-cm layer in the CONS plot (Fig. 4b). In both plots, due to its strong increase during the first 7 days, the methanol-extractable fraction in the 60–90-cm layer remained higher than in the upper three layers ($P < 0.001$) at t_{91} . In addition,

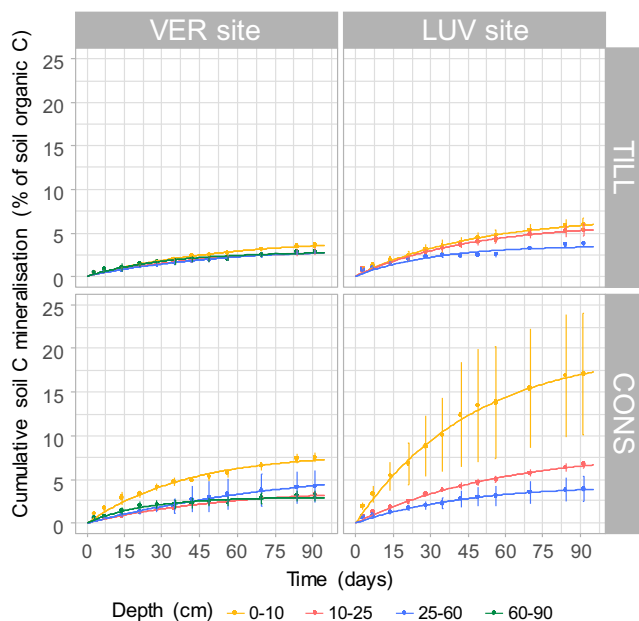


Fig. 2 Soil organic carbon mineralisation in conventional agriculture (TILL) and conservation agriculture (CONS) plots at the Vermic Umbrisol (VER) and Stagnic Luvisol (LUV) sites. Error bars indicate standard deviation

the methanol-extractable fraction in this layer became the second largest after the NER fraction after 14 days, whereas the water-extractable fraction exceeded the methanol-extractable fraction in the upper three layers.

The measured NER fraction data fit the exponential model well (Eq. 4) ($R^2 = 0.98 \pm 0.01$) (Fig 4c). The NER fraction at t_0 (y_0) was high in each layer and plot but higher in the TILL plot

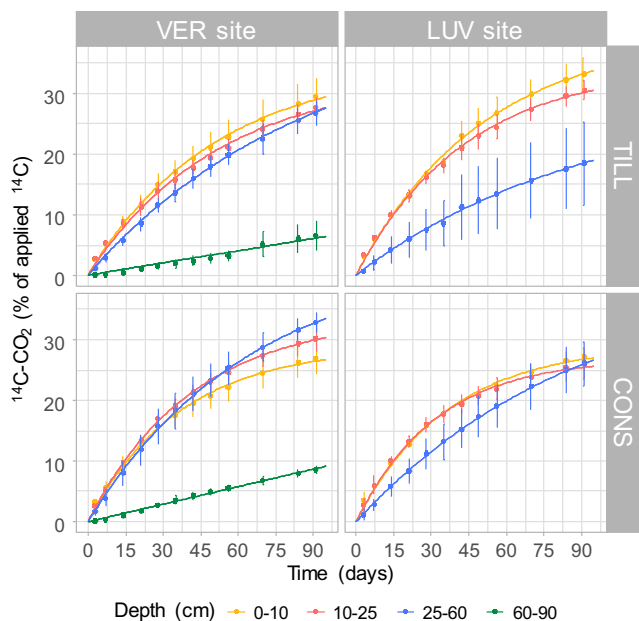


Fig. 3 Nicosulfuron mineralisation in conventional agriculture (TILL) and conservation agriculture (CONS) plots at the Vermic Umbrisol (VER) and Stagnic Luvisol (LUV) sites. Error bars indicate standard deviation

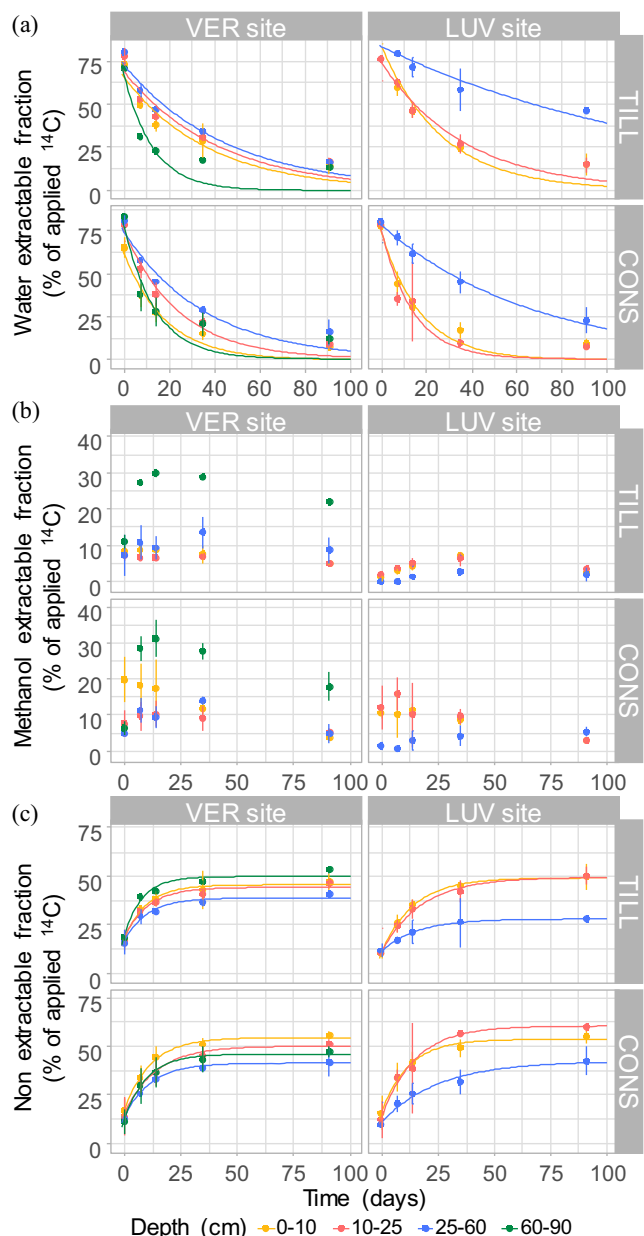


Fig. 4 Evolution of the a water-extractable, b methanol-extractable and c non-extractable fraction in conventional agriculture (TILL) and conservation agriculture (CONS) plots at the Vermic Umbrisol (VER) and Stagnic Luvisol (LUV) sites. Error bars indicate standard deviation.

than in the CONS plot ($P < 0.001$). It increased throughout the experiment, with a large increase (22.0 ± 3.6 percentage points) during the first 14 days. The exponential model always underestimated the final NER fraction a_{NER} , which was higher in the CONS plot than in the TILL plot ($P < 0.01$). NER formed faster in the 60–90-cm layer and slightly faster in the TILL plot than in the CONS plot, as indicated by the first-order rate constant b_{NER} (Table 3). However, neither agricultural management nor depth had a significant effect on b_{NER} .

LUV site

Like at the VER site, satisfying ^{14}C -nicosulfuron mass balances were reached, with ca $95.6 \pm 3.4\%$ of the initially applied radioactivity that was recovered after quantifying all of the fractions. At t_0 , most of the radioactivity was found in the water-extractable fraction (mean of $81.3 \pm 5.4\%$), regardless of the agricultural management and depth. Because of a particularly large water-extractable fraction, the methanol-extractable fraction at t_0 was especially low in the TILL plot (mean of $1.0 \pm 1.0\%$) (Fig. 4b). However, in the surface layers (0–10 and 10–25 cm), the methanol-extractable fraction was much greater in the CONS plot than in the TILL plot. Except for the 10–25-cm layer of the TILL plot, the NER fraction at t_0 was larger than the methanol-extractable fraction.

As at the VER site, ^{14}C -nicosulfuron began to mineralise within 3 days after it was applied (Fig. 3). Cumulative mineralisation data fit the exponential model well (Eq. 2) ($R^2 = 0.99 \pm 0.00$). After 35 days, mineralisation of ^{14}C -nicosulfuron in the 0–10 and 10–25-cm layers was higher in the TILL plot than in the CONS plot. During the incubation period, nicosulfuron mineralisation in the TILL plot remained lower in the deepest layer (25–60 cm) than in the two surface layers, with final cumulative mineralisation reaching $18.4\% \pm 6.9\%$ of applied ^{14}C in the 25–60-cm layer vs. $33.2 \pm 2.7\%$ and $30.5 \pm 1.6\%$ in the 0–10 cm and 10–25-cm layers, respectively. In the CONS plot, however, final cumulative nicosulfuron mineralisation at t_{91} was similar throughout the entire soil profile (reaching a mean of $26.5 \pm 0.7\%$ of applied ^{14}C) (Fig. 3). As for the VER site, the modelled final cumulative ^{14}C - CO_2 a_N in the 25–60-cm layer of the CONS plot was higher than the measured data. For the rest of the profile, a_N matched the final cumulative ^{14}C - CO_2 measured, indicating that most of the nicosulfuron mineralised during the 91-day incubation period. In the TILL plot, a_N was slightly (but not significantly) higher in the surface layers than in the 25–60-cm layer, whereas in the CONS plot, it was significantly higher in the 25–60-cm layer than the same layer in the TILL plot. The first-order rate constant b_N indicated faster mineralisation in the surface layers ($P < 0.001$), especially in the CONS plot ($P < 0.01$).

The measured water-extractable fraction data fit the first-order kinetics model well (Eq. 3) (mean $R^2 = 0.93 \pm 0.07$), but, like at the VER site, the model underestimated the final fractions of radioactivity in the water extracts, particularly in the 0–10 and 10–25-cm layers (Fig. 4a). The initial water-extractable fraction a_{WE} was slightly (but not significantly) higher in the TILL plot than in the CONS plot (Table 3). As observed at the VER site, this fraction decreased strongly early in the incubation before slowing down. The exponential decay was faster in the CONS plot ($P < 0.05$), especially in its surface layers ($P < 0.01$), as indicated by the first-order rate constant k_{WE} (Table 3). Small amounts were ultimately

recovered in these surface layers at t_{91} in both plots, while in the deeper soil layers (25–60 cm), the water-extractable fraction of ^{14}C in the still represented $22.6 \pm 7.5\%$ in the CONS plot and nearly $46.0 \pm 1.7\%$ in the TILL plot ($P < 0.05$).

The methanol-extractable fraction of ^{14}C increased slightly until day 35 and then generally decreased to its initial value in all layers in the TILL plot (Fig. 4b). In the CONS plot, it decreased in the surface layers, especially at t_{91} , whereas it increased in the 25–60-cm layer (reaching $5.3 \pm 1.5\%$ of applied ^{14}C). The final methanol-extractable fraction of ^{14}C was similar in all plots and layers.

The NER fraction increased until the end of the experiment, with a strong increase during the first 14 days of incubation ($21.0 \pm 6.6\%$) (Fig. 4c). The data fit the exponential model well (Eq. 4) ($R^2 = 0.96 \pm 0.06$). Despite similarly high initial NER fractions among layers, the final fraction at t_{91} was lowest in the 25–60-cm layer ($P < 0.001$), especially in the TILL plot. This was due to faster formation of NER in the surface layer ($P < 0.05$), as indicated by the first-order rate constant b_{NER} (Table 3). Mean b_{NER} was slightly (but not significant) higher in the surface layer than in the other layers in the CONS plot. At the end of the experiment, the NER fraction contained most of the radioactivity in the surface layers ($\approx 50\%$ of applied ^{14}C). It was higher in the CONS plot than in the TILL plot ($P < 0.01$) and lowest in the 25–60-cm layer of both plots ($P < 0.001$).

Discussion

Nicosulfuron behaviour in soil

Adsorption of nicosulfuron was low regardless of the agricultural management or depth, never exceeding $1.3 \pm 2.0 \text{ L kg}^{-1}$ (Table 2). The few studies on nicosulfuron available also generally reported low sorption, with sorption coefficients (K_d or K_f , with n usually close to 1) ranging from 0.02 to 1.8 L kg^{-1} in soils from the USA, Brazil, Argentina and Europe (EFSA 2008; Regitano and Koskinen 2008; Azcarate et al. 2015). To our knowledge, only two studies reported relatively high values of nicosulfuron K_f : up to 8.8 in a silt-loam soil in Iowa, USA (Gonzalez and Ukrainczyk 1996) and up to 16.1, in volcanic ash-derived soils, in Chile (Caceres-Jensen et al. 2020). Nicosulfuron is a weak acid (pKa of 4.78 and 7.58; EFSA 2008), which results in a mostly ionic form in most agricultural soils and thus little interaction with soil components, which can explain the low sorption. Nicosulfuron sorption was correlated with soil organic carbon content ($r = -0.03$) in either soil type, as suggested by the K_{OC} values; this result agrees with a previous study (Olivera et al. 2001). Ukrainczyk and Rashid (1995), however, reported that nicosulfuron sorption may depend instead on clay content, especially smectites, on which it could be rapidly and

irreversibly sorbed. Such high sorption of nicosulfuron could strongly decrease leaching to groundwater (Gonzalez and Ukrainczyk 1999). In our study, however, nicosulfuron sorption was not correlated with clay content ($r = 0.25$). Caceres-Jensen et al. (2020) also reported irreversibility of the nicosulfuron sorption, and this could be related to sorption mechanisms, specific to volcanic soils. In the same soil type as in the present study, we recently showed a high potential for nicosulfuron leaching up to 30 cm in undisturbed soil columns (Cueff et al. 2020).

Large amounts of nicosulfuron residues were still available in the water-soluble fraction at the end of incubation, especially in the deep soil layers, which probably resulted from the combination of low sorption and low mineralisation. Based on previous studies, nicosulfuron often mineralised more than other pesticides, such as acifluorfen, bentazon and chlorimuron, at an equivalent depth and incubation time (Reddy et al. 1995; Gaston et al. 1996; Gaston and Locke 2000). Boivin et al. (2004) reported that only 11% of applied bentazon (with low adsorption and a half-life similar to that of nicosulfuron) mineralised after 160 days in a Stagnic Luvisol (north-eastern France). Thus, in a similar soil type and with a longer incubation period, the mineralised fraction of bentazon was one-third that of nicosulfuron. We also identified strong and rapid formation of NER in both soils, which represented the main dissipation pathway for nicosulfuron. NER likely originated from biodegradation of nicosulfuron rather than from physical or chemical bonding given the low interaction of nicosulfuron with the soil matrix. It has been suggested for several pesticides that most NER have a biogenic origin (Nowak et al. 2011, 2013), as suggested by the relatively strong correlation ($r = 0.62$) between the first-order rate constant of nicosulfuron mineralisation (b_N) and the modelled final cumulative NER fraction (a_{NER}). Despite uncertainties about the reversibility of this dissipation process, NER formation is generally assumed to decrease the bioavailability of pesticides for a long time (Barriuso et al. 2008; Schäffer et al. 2018). Several studies indicated that most sulfonylurea herbicides had high potential to contaminate groundwater (Martins and Mermoud 1999; Sondhia 2009; Azcarate et al. 2015). Nicosulfuron's relatively low application rate ($< 100 \text{ g ha}^{-1}$) and relatively rapid dissipation by NER formation and mineralisation in surface layers could indicate little risk of transfer to groundwater. However, the low sorption in surface layers and slower dissipation in deeper layers suggest that groundwater contamination cannot be excluded, especially in situations of preferential transport.

Influence of soil type on nicosulfuron behaviour

We studied the fate of nicosulfuron at several depths in two soil types. Despite a strong decrease in microbial activity and biomass with depth, nicosulfuron mineralisation varied little

in the 0–60-cm layer but was lower below 60 cm. Observing mineralisation in the 25–60-cm layer similar to that in surface layers in both soil types was surprising, as mineralisation of several pesticides has been observed to decrease with depth in agricultural soils, in particular under the plough layer (0–30 cm) (Larsen et al. 2000; Rodríguez Cruz et al. 2008). Pesticide mineralisation is generally assumed to be a function of soil organic carbon content, which influences microbial activity and biomass (Holden and Fierer 2005). However, in our study, nicosulfuron mineralisation was not correlated with microbial biomass ($r = 0.09$) or microbial activity ($r = 0.02$).

Differences between microbial activity and pesticide mineralisation can be due to development of a microbial community that specialises in degrading a regularly applied pesticide. Several microbial strains that can degrade nicosulfuron directly have been identified in the literature (Lu et al. 2012; Zhang et al. 2012; Song et al. 2013; Zhao et al. 2015; Carles et al. 2017). Higher mineralisation potential in soil previously exposed to a given pesticide has already been observed for several pesticides (Sørensen and Aamand 2003; Zablutowicz et al. 2006). Although this phenomenon has yet to be observed for nicosulfuron, specific degradation is a likely hypothesis to explain the nicosulfuron mineralisation in our study, since it was applied regularly to the studied fields.

Influence of agricultural management

In both soil types, higher microbial activity was measured in the topsoil layer (0–10 cm) in the CONS plot than in the TILL plot and in the deeper soil layers. As mentioned by several authors, this difference in microbial activity probably resulted from accumulation of organic carbon at the soil surface in untilled soils (Kandeler et al. 1999). Despite much higher carbon content at the VER site and similar microbial biomass at both sites, microbial activity (illustrated here by soil carbon mineralisation) appeared to be much higher at the LUV site. At the LUV site, total mineralisation was strongly correlated with organic carbon ($r = 0.90$) and microbial biomass ($r = 0.93$). At the VER site, however, the correlations were much weaker (non-significant for organic carbon and $r = 0.66$ for microbial biomass), which may indicate high organo-mineral interactions at the VER site that limited microbial access to carbon for mineralisation (Dungait et al. 2012). Conservation agriculture generally favours these interactions (Veloso et al. 2019).

Although surface layers had higher microbial activity and biomass, nicosulfuron mineralisation in them was no higher than that in the 25–60-cm layer in the CONS plot at both sites. Lower pesticide mineralisation in the topsoil under conservation agriculture than in deeper layers, or than in topsoil layers under conventional agriculture, is generally attributed to a decrease in bioavailability. This decrease is related to higher sorption of pesticides due to an increase in organic carbon

content at the soil surface and the presence of crop residues that can intercept pesticides and reduce their availability to degrading microorganisms (Reddy and Locke 1998; Zablutowicz et al. 2000; Rodríguez Cruz et al. 2006; Alletto et al. 2010). Although we observed low sorption of nicosulfuron in the soil layers studied, the higher methanol-extractable fraction in the surface layers in CONS plots than in TILL plots from the beginning of incubation could indicate a lower nicosulfuron availability in the soil solution, thus limiting its mineralisation by microorganisms.

The NER fraction was always the largest dissipation pathway for nicosulfuron, especially under conservation agriculture, while the water-extractable fraction was always lower than that under conventional agriculture. In particular, a large water-extractable fraction was observed in the 25–60-cm layer in the TILL plot at the LUV site. This suggested low dissipation (through mineralisation and NER formation) of nicosulfuron, even 3 months after application. If nicosulfuron reaches this depth under field conditions, the TILL plot is likely to leach more of it than the CONS plot. At the LUV site, the CONS plot's formation of more NER, greater potential for nicosulfuron mineralisation in the soil profile (0–60 cm) and lower water-available fractions suggest that conservation agriculture may have lower risk of nicosulfuron leaching than conventional agriculture. As mentioned, however, this risk may still exist under preferential flow. Nevertheless, the VER site, due to similarities in nicosulfuron behaviour of the two types of management, conventional agriculture did not exhibit higher risks than conservation agriculture.

Conclusion

We studied sorption and degradation processes of nicosulfuron in two soil types from south-western France managed under either conservation or conventional agriculture. Sorption and incubation experiments were performed under controlled laboratory conditions using ^{14}C -nicosulfuron and used soil samples collected at several depths in the two soil types under both types of agricultural management. Little was known about the environmental behaviour of nicosulfuron, especially under contrasting types of agricultural management. Our results confirmed the low sorption affinity of nicosulfuron for soil components, which resulted in no significant difference in its sorption behaviour between the two types of agricultural management studied. Nicosulfuron mineralisation was influenced by mainly soil depth. For both soil types and types of agricultural management, the 25–60-cm layer had the same mineralisation as that in the topsoil layers but much lower microbial activity, suggesting development of a specific nicosulfuron-degrading microbial community. Formation of non-extractable residues is one of the main

nicosulfuron dissipation pathways, especially for conservation agriculture systems. For these reasons, little nicosulfuron may be transferred to groundwater under conservation agriculture, especially in a Stagnic Luvisol. However, due to a generally higher proportion of macropores in these systems, the risk of preferential flow cannot be ignored.

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Authors' contributions SC, LA, PB and VP conceived and designed research. SC and VD conducted the chemical analysis. SC, LA, PB and VP wrote the manuscript. All the authors read and approved the manuscript.

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Data availability The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Compliance with ethical standards

Competing interests The authors declare that they have no competing interests

Ethical approval and consent to participate Not applicable

Consent to publish Not applicable

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