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ORIGINAL PAPER



The contribution of deadwood to soil carbon dynamics in contrasting temperate forest ecosystems

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Abstract

Deadwood forms a significant carbon pool in forest systems and is a potential source of dissolved organic carbon (DOC) input to soil, yet little is known about how deadwood effects forest soil carbon cycling. Deadwood DOC inputs to soil may be retained through sorption or may prime microbial decomposition of existing organic matter to produce additional DOC. To determine impacts of deadwood on soil C cycling, we analysed surface soil from beneath deadwood or leaf litter only, along chronosequences of stands of lowland oak and upland Sitka spruce. The concentration and quality (by optical indices) of water-extracted soil DOC (water-extractable organic carbon; WEOC), in *situ* decomposition 'tea bag index' (TBI) parameters and enzymatic potential assays (β -D-cellubiosidase, β -glucosidase, β -xylosidase, leucine aminopeptidase, phosphatase, phenol oxidase) were determined. Presence of deadwood significantly (p < 0.05) increased WEOC concentration (~1.5 to ~ 1.75 times) in the mineral oak soil but had no effect on WEOC in spruce soils, potentially because spruce deadwood DOC inputs were masked by a high background of WEOC (1168 mg kg⁻¹ soil) and/or were not retained through mineral sorption in the highly organic (~90% SOM) soil. TBI and enzyme evidence suggested that deadwood-derived DOC did not impact existing forest carbon pools via microbial priming, possibly due to the more humified/aromatic quality of DOC produced (humification index of 0.75 and 0.65 for deadwood and leaf litter WEOC, respectively). Forest carbon budgets, particularly those for mineral soils, may underestimate the quantity of DOC if derived from soil monitoring that does not include a deadwood component.

Keywords Coarse woody debris · Dissolved organic carbon · Forest soils · Microbial priming

Introduction

Forests are acknowledged as large and important carbon (C) sinks, storing carbon in soils and both living and dead biomass. Globally, forests are estimated to hold 861 ± 66 Pg C, of which 119 ± 6 Pg are accounted for by temperate forest ecosystems (Pan et al. 2011). The amount of carbon stored in forest soils and biomass varies with forest type. Pan et al. (2011) state that tropical forests hold 56% of carbon

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in biomass with only 32% in the soil whereas boreal and temperate forests may hold 20% and 37% in biomass and 60% and 49% in soil, respectively. The remaining carbon is held within litter and deadwood and Pan et al. (2011) estimated that 8% $(73 \pm 6 \text{ Pg})$ of the world's forest C is held in deadwood. In the UK, forests store up to 1Pg C (Morison et al. 2012), of which 74% is held within the soils down to 1 m depth (Vanguelova et al. 2013). The remaining 26% C is split between tree biomass (22%) and litter and deadwood (4%) (Morison et al. 2012). The stock of C in deadwood alone in the UK represents 3.5% of the total forest C storage, with almost twice the amount per unit area in conifer (2.36 t C ha⁻¹) when compared to broadleaved forests (1.24 t C ha⁻¹) (Vanguelova et al. in-press). Deadwood may be present in the form of standing or lying dead trees or as stumps, with standing and lying deadwood categorised into decay classes for inventory purposes (Hunter 1990). For example, for lying deadwood, the decay classes range from 1 (least decomposed; intact bark, texture and structure)

to 5 (most decomposed; bark absent, powdery texture and structure collapsed) (Hunter 1990). Currently about 50% of the deadwood in the UK is in a less degraded state, at less than 10% decay or decay class 2, and the rest in different decay classes (Vanguelova et al. in-press).

Initiation of the decomposition of wood is thought to be primarily carried out by fungi (Schwarze et al. 2000) and occurs slowly, as the lignin content of wood provides a physical barrier to the enzymatic decomposition of nonlignin macromolecules, and, as such, it can take decades before a tree is fully decomposed (Russell et al. 2014). Research has shown that most conifers are slower to decay than broadleaved species (Weedon et al., 2009; Shorohova and Kapitsa 2014; Herrmann et al. 2015), partially due to the greater amounts of lignin, waxes, lipids and resins found in coniferous wood (Li et al. 2007; Lukac and Godbold 2011). Wood decay begins immediately following the death of cells. After the action of lignin peroxidases or polyphenol oxidases (Janusz et al. 2017), the resulting depolymerisation of lignin structures allows easier access for extracellular hydrolytic enzymes that are responsible for the depolymerization of polymers such as cellulose and hemicellulose (Li et al. 2018). These depolymerization reactions produce lower molecular weight compounds with increased solubility and initial bioavailability; deadwood-derived dissolved organic carbon (DOC) may enter underlying soil via leaching (Wambsganss et al. 2017; Piaszczyk et al. 2019b), contributing to the soil DOC pool and undergoing subsequent fates, e.g. microbial metabolism including respiration to, or driving production of, greenhouse gases (CO2, CH4, N2O) or stabilization (Wambsganss et al. 2017) through interaction with soil minerals (Guggenberger and Kaiser 2003), depending on its quality. The flux of DOC released by deadwood (44–53 mg per g material) has been found to be comparable to leaf litter (47–49 mg per g material) and is significantly greater than forest vegetation and the upper soil layers (3–18 mg per g material) (Hollands et al. 2022).

Although we know that deadwood forms a significant pool of C in forest systems, and previous studies have shown that soil DOC concentrations are elevated in forest systems with deadwood when compared to those without (Spears and Lajtha 2004; Hafner and Groffman 2005; Kahl et al. 2012; Stutz et al. 2019; Minnich et al. 2021; Hollands et al. 2022), it is less clear what effects the presence of deadwood has on forest soil C cycle processes and how this varies in contrasting forests. It is not clear if the reported elevations in soil DOC concentrations are a result of solely the input and subsequent stabilization of deadwood decomposition derived DOC, or, if inputs of deadwood DOC promote the release of DOC from existing soil organic matter (SOM) (Minnich et al. 2021). On the one hand, deadwood-derived DOC might be stabilized in soil as a result of either: (i) chemical protection from decomposition through association with mineral phases; (ii) physical protection through sequestration within soil structure; or, (iii) biochemical protection through inherent or acquired structural recalcitrance (Six et al. 2002; von Luetzow et al. 2006; Piaszczyk et al. 2019a). On the other hand, deadwood DOC components may prime decomposition of, or release DOC from, existing SOM pools through: (i) stimulation of microbial activity and as such enzymatic activity (Fontaine et al. 2003; Gonzalez-Polo et al. 2013; Beverly and Franklin 2015; Minnich et al. 2021) which in turn primes increased depolymerisation of, and release of DOC from, native soil organic matter; or (ii) abiotic liberation of native dissolved organic compounds from protective associations with minerals (Keiluweit et al. 2015). As previously mentioned, deadwood varies with respect to the stage of wood decomposition, or decay class. Wood decay class is known to influence subsequent rates of deadwood decomposition (Tobin et al. 2007; Olajuyigbe et al. 2011; Stutz et al. 2019). Wood in the late stages of decay hosts a greater primary decomposer microbial biomass (Kueffer et al. 2008; Baldrian et al. 2016). Increased microbial and fungal biomass in and under deadwood (Peršoh and Borken 2017) may lead to larger DOC and CO₂ fluxes being produced, and therefore potentially enhance impacts on the soil DOC pool and C cycling processes. Wood composition will also influence the rates of decay, with wood from conifers ('softwoods') and broadleaves ('hardwoods') decaying at different rates (Herrmann et al. 2015), and likely producing different quality and lability of DOC.

Soil properties, such as soil texture and type, pH and moisture will affect the amount of organic matter held in soils (Matus 2021) and influence the rate of decomposition processes that may release DOC. Silt and clay content in soils may capture and stabilize organic compounds through sorption (Six et al. 2002; Villada 2013; Matus 2021). Low soil pH and high moisture content inhibit decomposition rates (Hagemann et al., 2010) which may increase SOM accumulation (Keith et al. 2009). It is possible that the physical presence of large woody debris on the forest floor may create a localised microclimate, through the shading of soils which will affect temperature, or preventing water and litter reaching the soil. In tropical forests, deadwood has been found to buffer soil temperature, preventing diurnal fluctuations (Zalamea et al. 2016). Anoxic conditions can be created through waterlogging of soils and these conditions are known to inhibit decomposition processes (Hagemann et al. 2010). However, little research has been carried out into the creation of microclimates under deadwood in temperate forests (Woodall et al. 2020).

Accordingly, the aim of this study is to confirm whether deadwood produces significant inputs of DOC into underlying soil and to identify the effect this has on C cycling. Specific objectives of this study are to: (1) identify whether the presence of lying deadwood influences the amounts of DOC in underlying soil, comparing common hardwood (oak) and softwood (Sitka spruce) species on their typical soil types in Britain; (2) identify the impact of deadwood on soil carbon processes and cycling; and, (3) characterize the quality of soil DOC in order to help identify its source and potential lability. We use a combination of traditional measurements (e.g. extracellular enzyme potentials as sensitive indicators of change in decomposition processes) and more novel indices, as applied to forest C studies, to derive decomposition data (the teabag index (Keuskamp et al. 2013) and to assess the biochemical stability (SUVA₂₅₄ (Hansen et al. 2016) and humification index (HIX) (Ohno 2002) optical methods) and source (fluorescence index (FI) (McKnight et al. 2001) of soil DOC.

Materials and methods

Study sites

Oak woodland and Sitka spruce plantations represent the two most common British forest types. In the UK, oak covers 16.4% of land under broadleaves (Brewer 2014a) whilst Sitka spruce covers 50.8% of land under conifers (Brewer 2014b). Two chronosequences (Benham et al. 2012; Vanguelova et al. 2019) were used: (1) in Alice Holt Forest, Surrey in S.E. England; and, (2) in Kielder Forest, Northumberland in N.E. England to represent a range of tree stand ages and decay classes of deadwood (Table 1). The Alice Holt Forest sites $(51^{\circ} 9' \text{ N}, 0^{\circ} 52' \text{ W})$ consist of young (~30 years old), middle age (~70 years old) and old (~180 years old) managed lowland oak forest (mainly Quercus robur, L.) on surface water gleys (stagnosols). The upper soil profile consists of silty clay in the Ah horizon (7 cm deep) over an Eg horizon (8 cm deep). High rates of decomposition in Alice Holt soils mean that there is no distinct O layer; instead, organic matter is mixed with mineral soil in the upper Ah horizon. The Ah horizon has a bulk density of 704.3 kg m⁻³ and carbon stock of 13.3 t ha⁻¹. The Eg horizon has a bulk density of

 Table 1
 Site plot information

Site	Age group	Age (years)	Rotation	Decay class
Alice holt forest	Young	24–34	_	1–2
(oak)	Mid	64-84	_	3
	Old	184+	-	4–5
Kielder forest	Young	12, 18, 19	2nd	1–2
(spruce)	Mid	20, 30, 31	2nd	3
	Old	46,~60,74	1st, 2nd	4–5

Decay class represents the dominant stage of decay found within each age group

968.2 kg m⁻³ and carbon stock of 22.3 t ha⁻¹. Elevation is 80 m with monthly average air temperatures ranging between 4.7 and 17.6 °C with an annual mean of 10.5 °C and mean annual precipitation of 634 mm. Kielder Forest (55° 15' N, 2° 23' W) consists of uniform, managed young $(\sim 10-20 \text{ years old})$, middle age (20-40 years old) and old (40-60 years old) upland Sitka spruce (Picea sitchensis (Bong.), Carr.) plantations on peaty gley soils and deep peat (stagnohumic gleys/histosols). The upper soil profile is predominantly organic material, with an O layer (2 cm deep) over the H peat layer (17 cm deep). The O layer has a bulk density of 86 kg m⁻³ and carbon stock of 22.7 t ha^{-1} . The H layer has a bulk density of 157 kg m^{-3} and carbon stock of 114 t ha⁻¹. Elevation ranges between 240 and 365 m, monthly average air temperatures range between 3 and 14.8 °C with an annual mean of 8.3 °C, and precipitation of 759 mm. The majority of plots are in their second rotation (i.e. second crop) since afforestation (Table 1). At each site three plots of 10 m^2 , per chronosequence age group were selected for field measurements and sampling. These age groups contained a range of different sized debris at different stages of decay. The dominant decay stage per age group is shown in Table 1; however, other decay classes were present at each plot. Decay class was visually assessed according to the guidelines by Hunter (1990) as decay classes 1–5, whereby freshly fallen wood with little to no decay is considered class 1 whilst heavily decayed wood is class 5.

Field sampling

At each chronosequence plot, five subplots (~ $1 \times 1 \text{ m}^2$) where deadwood was present were randomly selected along with five subplots ($\sim 1 \times 1 \text{ m}^2$) where no deadwood was present, but only forest leaf litter was. These subplots will be referred to as 'deadwood' and 'leaf litter', respectively. Deadwood diameters at Alice Holt ranged between 5 and 90 cm, with a mean of 10.8 cm. Two samples of large diameter deadwood (> 30 cm diameter) were selected due to an absence of smaller debris. Deadwood diameters at Kielder ranged between 5 and 13.7 cm, with a mean of 7 cm. An average mass of 0.1 kg m^{-2} deadwood $(\sim 4.5 \text{ m}^3 \text{ ha}^{-1})$ was found at Alice Holt (Hollands et al. 2022) and ~0.3 kg m⁻² (~10.5 m³ ha⁻¹) at Kielder. After removal of the surface litter layer at both the leaf litter and deadwood plots, a soil sample was taken from the top 10 cm of soil from all ten subplots. Soil temperature of the soil surface layer from each of the ten subplots was measured with a handheld probe (Delta-T Devices Ltd., UK). This was repeated over two site visits, the first in June-July and the second in September-October 2017.

Tea bag index

A tea bag index (Keuskamp et al. 2013), which exploits the differential in decomposition between green tea (Camellia sinensis; fast decomposing) and rooibos tea (Aspalathus linearis; slow decomposing) leaves over ca. 3 months, was used to compare decomposition rate and organic matter stabilization potential of soils under deadwood and neighbouring soils where no deadwood was present. Following the protocol of Keuskamp et al. (2013), pre-weighed pairs of commercially available Lipton green tea and Lipton rooibos tea bags were buried in June and July 2017 and retrieved in September or October after an incubation of 81-85 days. They were buried 8 cm deep and 15 cm apart, in the soil layer beneath the deadwood and in leaf litter subplots. Following retrieval, tea bags were oven dried at 60 °C and external debris removed before weighing. Mass lost between the date of initial burial and retrieval was used to calculate both k, a decomposition rate that measures the turnover time of labile carbon, and S, the stabilisation factor, a measure of the stabilisation potential of organic carbon according to the following equations (Keuskamp et al. 2013).

$$W(t) = a_r e^{-kt} + (1 - a_r)$$
$$S = 1 - \left(\frac{a_g}{H_g}\right)$$

where: W(t) is the weight (g) of rooibos tea remaining after incubation time t (d); a_r = decomposable fraction of rooibos litter (g g⁻¹) as calculated from S and the hydrolysable fraction of rooibos tea (H_r); k = rooibos tea decomposition rate (d⁻¹). S = stabilization factor (unitless); a_g = decomposed fraction of green tea after incubation (g g⁻¹); H_g = hydrolysable fraction of green tea (g g⁻¹). H_g and H_r have been previously determined as 0.842 and 0.552 by Keuskamp et al. (2013).

Lab analyses

Soil samples were kept stored at 4 °C until analysis. Moisture content of soils were measured gravimetrically as loss of mass on oven heating to 105 °C. Soil organic matter (%) was measured through mass loss on ignition at 550 °C. Subsamples of soil were oven dried at 105 °C for 12 h prior to grinding with a planetary ball mill (Pulverisette 5, Fritsch, Germany). The total C and N of ground samples was then measured using a Thermo Flash 2000 Carbon and Nitrogen Analyser.

Water-extractable carbon was determined by mixing 4.5 g soil in to 45 ml water, at a ratio of 1:10 soil:ultrapure

water. These were placed on an end-over-end shaker for 24 h before pH was measured and the extracts then centrifuged at 3500 rpm for 15 min. Extracts were then filtered through Whatman GF/A filter papers before centrifuging at 3500 rpm for a further 15 min and vacuum filtering through 0.45 μ m cellulose nitrate filter papers. Total organic carbon (TOC; mg L⁻¹) of these filtered solutions was measured using a carbon analyser (Shimadzu TOC-L, Shimadzu Corporation, Japan) as Total Carbon (mg L⁻¹)— Inorganic Carbon (mg L⁻¹). As they had been filtered to 0.45 μ m, it is presumed that all TOC measured is DOC. Water-extractable organic carbon (WEOC; mg kg⁻¹ dry soil) was calculated from the DOC readings to correct for the volume of water used during extraction and dry mass of soils. Quantities of DOC will be referred to as WEOC.

The specific ultraviolet absorbance at 254 nm $(SUVA_{254})$ can be used as a proxy measurement for the aromatic content of DOC, with higher SUVA_{254} values indicating a greater aromatic content (Hansen et al. 2016). Absorbance of soil solutions was measuring using spectrophotometer (Jenway 5000, Jenway, UK) at wavelength 254 nm. A baseline of ultrapure water was used and blanks run every five samples. SUVA₂₅₄ (L mg C⁻¹ m⁻¹) was then calculated as:

$$SUVA_{254} = \frac{Absorbance_{254}}{DOC (mg L^{-1})} \times 100$$

Fluorescence measurements of the soil solutions and enzyme activities were made using a microplate reader (SpectraMax i3x, Molecular Devices, USA). In order to calculate a humification index (HIX), spectra were obtained with an excitation wavelength of 254 nm over the emissions waveband of 300–480 nm at increments of 5 nm (Ohno 2002) and calculated as:

$$\text{HIX} = \frac{\sum I_{435 \to 480}}{\sum I_{300 \to 345} + \sum I_{435 \to 480}}$$

where *I* is the detected fluorescence intensity.

A fluorescence index (FI) was calculated using the ratio of wavelengths 470 to 520 nm at excitation 370 nm (McKnight et al. 2001).

Enzyme assays to determine potential hydrolytic depolymerase activity rates were selected as follows: carbon cycle— β -D-cellubiosidase (CB; cellulose degradation), β -glucosidase (BG; sugar degradation) and β -xylosidase (XYL, hemicellulose degradation); carbon and nitrogen cycle—Leucine aminopeptidase (LAP, protein degradation); phosphorus cycle—phosphatase (PHOS, phosphorus mineralisation). Phenol oxidase (POX) activity, an important oxidative enzyme involved in lignin depolymerization was also determined. Microplate fluorometric analysis of CB, BG, XYL, LAP and PHOS potential activities in soil slurries were determined using the methods of Bell et al. (2013). Fluorescence was measured at excitation 365 nm and emission 450 nm after a two-hour incubation at mean field soil temperature (Table 2). For all enzyme assays, sodium acetate buffer was used and adjusted to the corresponding site pH (Table 2) using glacial acetic acid.

POX activity was measured following the methods of Allison (2012) using L-dihydroxyphenylalanine (L-DOPA) as a substrate. Optimum incubation times were found after reading plates at absorbance 460 nm every 30 min for five hours. Final assay plates were incubated at mean field soil temperature (Table 2) for 3.5 h before measurement.

Deadwood DOC extraction

In June 2018, the young and old plots at Alice Holt were revisited (Zhao 2018) and samples of deadwood (>10 cm diameter) were collected for the extraction of WEOC. Five samples per plot were selected and cut to ~20 cm lengths in the field. These were sealed in plastic bags and transported back to the laboratory for storage at 4 °C. Prior to extraction, deadwood samples were cut into 1.5 cm × 1.5 cm × 1.5 cm cubes of centre wood (i.e. cubes were formed by sawing off the outer wood to leave a cube at the centre) and air dried for 15 days. These were then placed in 50 ml centrifuge tubes and 33.75 ml ultrapure water added. Tubes were shaken on an overhead shaker for 24 h before centrifuging at 3500 rpm for 15 min. Supernatants were filtered through Whatman GF/A and then through 0.45 μ m cellulose nitrate filter papers. Filtrates were further diluted as 2 ml sample: 4 ml MilliQ water before analysis on an Analytical Sciences liquid CN analyser.

Statistical analyses

Statistical analyses were carried out using R core v 3.5.1 (R Core Team 2017). Data were tested for normality using Shapiro–Wilk test and equality of variances using Levene's test. Data were \log_{10} transformed where equality of variances was not met. The effect of site, stand age and presence of deadwood were analysed using a general linear model. Significant differences were accepted at *p* < 0.05. Means are presented with standard errors. Tukey HSD post hoc testing was used to identify where significant differences occurred.

Results

Effects of deadwood on underlying soil properties

Soil properties varied significantly between the two sites, with Kielder being cooler, wetter, and more organic and

Table 2 Average surface soil properties (0–10 cm depth) from Alice Holt and Kielder Forests ± 1 standard error of the mean, grouped by standage and the presence or absence of deadwood

		Alice holt forest (oak)				Kielder forest (spruce)			
		Young	Mid	Old	Mean	Young	Mid	Old	Mean
Field soil temperature (°C)	¹ Leaf litter	16.9 ± 0.2	16.6±0.1	16.5 ± 0.1	16.6 ± 0.1^{A}	10.7 ± 0.1	10.9 ± 0.1	10.2 ± 0.1	$10.6 \pm 0.1^{\circ}$
	¹ Deadwood	16.3 ± 0.1	16.3 ± 0.1	16.5 ± 0.1	$16.4\pm0.1^{\rm B}$	10.7 ± 0.1	10.8 ± 0.1	10.2 ± 0.1	$10.6\pm0.1^{\rm C}$
	² Leaf litter	13.0 ± 0.1	12.5 ± 0.1	12.5 ± 0.1	12.7 ± 0.0	9.8 ± 0.0	9.7 ± 0.1	9.1 ± 0.1	9.5 ± 0.0
	² Deadwood	13.2 ± 0.1	12.5 ± 0.1	12.5 ± 0.1	12.8 ± 0.1	9.9 ± 0.0	9.6 ± 0.1	9.1 ± 0.1	9.5 ± 0.0
рН	Leaf litter	4.4 ± 0.1	4.2 ± 0.1	4.5 ± 0.1	4.4 ± 0.1	3.7 ± 0.1	3.8 ± 0.1	4.1 ± 0.1	3.9 ± 0.1
	Deadwood	4.3 ± 0.1	4.4 ± 0.1	4.6 ± 0.2	4.4 ± 0.1	3.8 ± 0.1	3.7 ± 0.1	4.0 ± 0.1	3.9 ± 0.1
Soil water content (%)	Leaf litter	34.6 ± 1.4	33.2 ± 1.9	33.6 ± 2.4	33.8 ± 1.1	73.0 ± 3.3	70.8 ± 1.8	78.4 ± 0.6	74.0 ± 1.3
	Deadwood	36.5 ± 1.4	35.6 ± 3.2	34.3 ± 2.7	35.4 ± 1.4	71.6 ± 1.5	69.2 ± 2.2	77.1 ± 1.3	72.6 ± 1.1
SOM	Leaf litter	17.0 ± 1.3	19.0 ± 2.7	21.5 ± 2.6	19.2 ± 1.3	76.9 ± 6.4	92.7 ± 1.6	93.6 ± 1.2	87.7 ± 2.5
(%)	Deadwood	18.9 ± 1.0	26.9 ± 5.0	23.3 ± 3.0	23.0 ± 2.0	80.0 ± 3.7	91.9 ± 3.4	90.8 ± 3.2	87.6 ± 2.1
C (%)	Leaf litter	10.0 ± 1.2	13.0 ± 2.7	9.9±1.4	11.0 ± 1.1	35.5 ± 2.9	46.7 ± 0.3	46.5 ± 0.8	42.9 ± 1.3
	Deadwood	12.0 ± 1.3	15.3 ± 2.3	9.4 ± 0.9	12.2 ± 1.0	38.3 ± 1.7	46.8 ± 0.5	44.5 ± 1.7	43.2 ± 1.0
N (%)	Leaf litter	0.7 ± 0.1	0.8 ± 0.1	0.7 ± 0.1	0.7 ± 0.1	1.3 ± 0.1	1.5 ± 0.0	1.7 ± 0.1	1.5 ± 0.1
	Deadwood	0.8 ± 0.1	0.9 ± 0.1	0.6 ± 0.0	0.8 ± 0.1	1.3 ± 0.1	1.5 ± 0.1	1.6 ± 0.1	1.5 ± 0.0
C/N	Leaf litter	14.5 ± 0.4	15.1 ± 0.6	14.2 ± 0.4	14.6 ± 0.3	27.8 ± 1.1	31.4 ± 1.0	28.4 ± 1.2	29.2 ± 0.7
	Deadwood	15.3 ± 0.6	15.7 ± 0.6	15.4 ± 0.6	15.5 ± 0.3	29.0 ± 1.1	30.3 ± 0.7	27.6 ± 1.1	28.9 ± 0.6

N=15 per group. Field soil temperature and moisture were measured at ¹ time of tea bag burial (June-July) and ² time of tea bag retrieval (September–October). Tukey groupings for the effect of interaction between presence of deadwood and site are shown by superscript letters. Where Tukey groupings are not given, there was no significant presence of deadwood * site interaction term

acidic (Table 2). The larger organic matter content (%) at Kielder was reflected in the C (%) measures (43% and 12% C at Kielder and Alice Holt, respectively), and subsequently the surface soil C:N ratio. Presence of deadwood had no effect on soil pH (p=0.6), gravimetric soil water content (p=0.721), SOM (p=0.333) or soil C:N (p=0.502). Oak subplots with deadwood had slightly higher % water content to leaf litter subplots (Table 2), whilst deadwood subplots in the spruce forest had marginally lower water content to the leaf litter subplots.

Stand age affected soil properties across both sites. Soil pH was significantly higher (0.18–0.41 units) in old stands than the mid-aged and young stands, at both forests (p < 0.001). Gravimetric water content was significantly affected by an interaction between site and age (p = 0.013) whereby at Kielder, old stands were found to hold significantly more water than the mid-aged stand (p = 0.004), with means of $78 \pm 1\%$ and $70 \pm 1\%$, respectively. Young stands had a significantly lower organic matter content than old or mid stands (p < 0.001) at both forests. Soil C:N was also affected by stand age, with mid-aged stands having a significantly (p = 0.048) higher ratio in both forests (Table 2).

Effect of deadwood on surface soil WEOC

DOC release from deadwood ranged between 0.55 and 15.45 mg⁻¹ g⁻¹ air dried wood, with means of 2.45 ± 0.49 mg⁻¹ g⁻¹ air dried wood and 3.45 ± 0.93 mg⁻¹ g⁻¹ air dried wood in the old and young plots at Alice Holt, respectively. Age did not significantly affect the amount of DOC released by deadwood (p = 0.33). Concentrations of Water-Extractable Organic Carbon (WEOC) were two to five-fold higher in soils from Kielder Forest than Alice Holt Forest (Fig. 1), averaging 1168 ± 43 and 353 ± 45 mg kg⁻¹ dry soil, respectively. There was a

Fig. 1 Mean water-extractable organic carbon (WEOC) in surface soil from Alice Holt and Kielder forests with or without deadwood, and in different stand age groups. Error bars are ± 1 SE of the mean. N=15 per group. Groups that do not share a letter are significantly different from one another significant interaction between site and the presence of deadwood (p = 0.003), as soils under deadwood in Alice Holt Forest held $1.45 - 1.76 \times$ more WEOC than the corresponding leaf litter soils (Tukey HSD p < 0.001). However, there was no effect of the presence of deadwood evident in Kielder Forest plots. Stand age also had no significant effect on WEOC quantity (p = 0.900).

A fluorescence index (FI) can be used to infer the source of DOC in the soil solutions. FI values > 1.9 indicate DOC from a microbial origin, such as lysates and extracellular release from bacteria, whilst values < 1.4 might indicate an origin from plant material, reflecting contributions from lignin degradation products (McKnight et al. 2001). Soil solutions from Kielder Forest showed a lower FI compared to those from Alice Holt Forest, ranging between 1.07 and 3.31 at Kielder Forest and 1.39 and 4.03 at Alice Holt Forest. Most samples fell within a FI of 1.4-1.9 indicating a mix of origins (Table 3). However, for Alice Holt Forest, soil solutions from the young leaf litter stand and the old stands with deadwood had mean FI values of 2.20 and 2.09, respectively, (Table 3) which indicates a microbial origin only. Soil solutions from under deadwood in the young stands in Kielder Forest had a mean of 1.36 which indicates WEOC originated from plant material only. Because of the differences between stand ages and sites, a significant three-way interaction was found between site, stand age and the presence of deadwood for the fluorescence index (p=0.013). Mean values and the results of Tukey HSD post hoc testing are shown in Table 3.

For soil solutions from Alice Holt Forest the mean humification index (HIX) was higher in soils under deadwood than in leaf litter soils (Table 3), indicating that soil from under deadwood had a greater degree of humification. Significant interaction was found between site and the presence of deadwood (p=0.015) and site and age (p=0.003). The presence of deadwood significantly (p=0.007) increased the HIX value at Alice Holt Forest whilst stand age did not.



		Alice Holt Forest (oak)				Kielder Forest (spruce)			
		Young	Mid	Old	Mean	Young	Mid	Old	Mean
FI	Leaf litter	2.20 ± 0.20^{a}	1.74 ± 0.08^{abc}	1.66 ± 0.04^{abc}	1.87 ± 0.08	$1.52 \pm 0.08^{\circ}$	$1.53 \pm 0.07^{\circ}$	1.64 ± 0.13^{bc}	1.56 ± 0.05
	Deadwood	$1.80\pm0.09^{\rm abc}$	1.78 ± 0.12^{abc}	2.09 ± 0.19^{ab}	1.89 ± 0.08	$1.36\pm0.05^{\rm c}$	$1.66 \pm 0.14^{\rm abc}$	$1.40 \pm 0.04^{\circ}$	1.48 ± 0.05
HIX	Leaf litter	0.60 ± 0.04	0.67 ± 0.04	0.68 ± 0.04	$0.65\pm0.02^{\rm B}$	0.76 ± 0.03	0.72 ± 0.04	0.63 ± 0.05	$0.70\pm0.02^{\rm AB}$
	Deadwood	0.74 ± 0.03	0.77 ± 0.04	0.76 ± 0.04	$0.75\pm0.02^{\rm A}$	0.74 ± 0.04	0.74 ± 0.04	0.60 ± 0.05	$0.70\pm0.03^{\rm AB}$
$\begin{array}{c} SUVA_{254}(L \\ mg \; C^{-1} \; m^{-1}) \end{array}$	Leaf litter	4.62 ± 0.13	4.90 ± 0.12	4.73 ± 0.11	$4.75\pm0.07^{\rm A}$	4.49 ± 0.33	3.47 ± 0.20	3.56 ± 0.18	$3.85\pm0.16^{\rm B}$
	Deadwood	4.84 ± 0.06	5.29 ± 0.24	5.26 ± 0.28	$5.13\pm0.13^{\rm A}$	4.12 ± 0.14	3.48 ± 0.26	3.54 ± 0.18	$3.72\pm0.12^{\rm B}$

Table 3 Mean optical properties of soil WEOC from Alice Holt and Kielder Forests with or without deadwood, and in different stand age groups

FI—Fluorescence index; *HIX*—humification index; SUVA₂₅₄—Specific UV Absorbance at 254 nm. Results of Tukey post hoc testing for the fluorescence index (FI) are shown in lowercase, where a significant three-way interaction occurred. Results of post hoc testing for HIX and SUVA are shown in uppercase. Groups that do not share a letter are significantly different

For Kielder Forest samples, the opposite occurred, whereby presence of deadwood had no significant effect but stand age did (old-mid p = 0.035, young-mid p = 0.995, young-old p = 0.006), with the old stands having a lower HIX value.

The specific ultraviolet absorbance (SUVA₂₅₄) was significantly larger for soil solutions from Alice Holt Forest than Kielder Forest for all sample groups (Table 3). Significant interactions were found between site and the presence of deadwood (p=0.032) and site and age (p < 0.001). Post hoc testing found that the only significant within-site differences occurred at Kielder Forest between the young stand and the mid (p=0.001) and old stands (p=0.003).

Effect of deadwood on tea bag index parameters

The stabilisation factor of soils (S) and decomposition rate (k) derived from the tea bag index method was up to twofold higher in soils from Kielder Forest than those from Alice Holt Forest, with a significant interaction between site and age (p < 0.04), (Fig. 2). The presence of deadwood did not influence either S (p=0.763) or k (p=0.720). Within Kielder Forest samples, the old stands had a significantly higher S than the mid stands, with mean values of 0.47 ± 0.04 and 0.28 ± 0.02 (p < 0.001), respectively (Supp. Mat. Table 1). Little variation occurred with the mean decomposition rates derived from the tea bag index (*k*) between samples (Fig. 2).

It should also be noted that whilst a large number of tea bag pairs were buried (n = 90 pairs per site), 21 pairs were discounted due to holes and 26 produced negative k values, which were discounted from analysis (Table S2 and Discussion in Supplementary Information).

Effect of deadwood on soil enzyme activity potential

Enzyme activity rates were highly variable between soil samples (large s.e.m, Fig. 3). However, a clear site difference occurred, whereby soil suspensions from Kielder Forest showed significantly higher enzyme activity rates than those from Alice Holt Forest. This site difference ranged from between 1.2 x (β -D-cellubiosidase young plots with only leaf litter) to 19.3 x (Leucine aminopeptidase young plots with only leaf litter). Means are presented in Supplementary Information Table S3. However, the presence of deadwood did not appear to stimulate an increase in enzyme activity at either forest, nor







Fig. 3 Potential enzyme activities in soils at Alice Holt and Kielder forests, with and without deadwood, and in different stand age groups: **a**: β -D-cellubiosidase, **b**: β -xylosidase, **c**: β -glucosidase, **d**: Phosphatase, **e**: Leucine aminopeptidase, **f**: Phenol oxidase. Data

are *=mean— =median, •=outliers defined as outside of the range of plot whiskers, \pm =plot whiskers to show Q1—1.5*IQR or Q3+1.5*IQR. Mean values and results of statistical tests can be found in Table 3 of the supplementary material

in different aged stands. There was no apparent link between DOC origin and enzyme activity rates.

Stand age significantly affected the carbon cycling enzymes β -glucosidase (p = 0.002) and marginally β -D-cellubiosidase (p=0.076) and β -xylosidase (p=0.074). A significant interaction was found between site and age for β -D-cellubiosidase (p=0.041) but not β -xylosidase (p=0.093). A significant interaction between site, the presence of deadwood and stand age was found for β -glucosidase (p = 0.011). For each of these enzymes, activity rates peaked in the mid-aged stands in both Alice Holt and Kielder Forests. For these three enzymes, at Alice Holt Forest, the average activity rates were between 1.1 and 1.5 times higher in the mid age stands than in young or old stands, whilst at Kielder Forest the rates varied between 1.3 and 2.4 times higher in mid-aged stands than young and old. A significant interaction also occurred between site and stand age for phosphatase activity (p = 0.010). Site and stand age effects on enzyme potentials are discussed in the Supplementary Information.

Discussion

Soil properties in contrasting forest systems

In order to examine the contribution of deadwood to soil carbon dynamics, we chose to sample soils from stands in a lowland oak forest (Alice Holt) and an upland Sitka spruce forest (Kielder) as forest sites that contrast but also represent the dominant broadleaf and coniferous forest cover in the UK, covering 16.4% (Brewer 2014a) and 50.8% (Brewer 2014b) of forested land, respectively.

There was a clear difference in the soil and WEOC properties between Kielder and Alice Holt Forests, with significant differences occurring between sites for all measures except the HIX. Comparison of soil properties (Table 2) between the two sites reflected the soil classifications of peaty gleys/deep peat and surface water gleys at Kielder and Alice Holt, respectively. The cooler and wetter conditions at Kielder, combined with low pH, inhibit decomposition rates (Hagemann et al. 2010) and favour SOM accumulation (Keith et al. 2009) at this site. Our finding that WEOC concentrations were substantially higher at Kielder (1168 mg C kg⁻¹ soil) than Alice Holt (353 mg C kg⁻¹ soil) (Fig. 1) most likely reflected the SOM content as a substantial source of soluble carbon (Morison et al. 2012).

Deadwood influence on concentrations of soil water-extractable organic carbon

It was expected that deadwood would provide a source of DOC into underlying soil. Out of all of the soil properties studied, the clearest effects of deadwood were seen in soil WEOC, albeit in a site-dependent manner. For the surface soils under oak at Alice Holt Forest, WEOC concentrations were significantly (between 1.45 and 1.76 times) higher for soils under deadwood when compared to soils without deadwood. However, a deadwood effect on WEOC was not observed in soils from the spruce stands in Kielder Forest. Our finding of greater quantities of WEOC under deadwood at Alice Holt is in accordance with other research that has found that concentrations of soil DOC increase in forests over mineral soils where deadwood is present (Spears and Lajtha 2004; Hafner et al. 2005; Kahl et al. 2012), with increases of up to nine-fold when compared to soils with leaf litter only (Hafner et al. 2005). The organic nature of the peaty-gley Kielder Forest soils may explain why no difference was seen between WEOC in leaf litter soils and soils below deadwood whereas a difference was seen between them in the mineral soils at Alice Holt Forest. In Kielder Forest, where % organic matter averages 77-94%, the input of DOC by deadwood is likely to be masked by such high levels of existing SOM as a significant source of WEOC. In contrast, at Alice Holt Forest, where % organic matter averages 17-27% in topsoil, additional DOC inputs from deadwood may be more easily detectable. In addition, the soil type may have played a role in the retention of deadwood-derived DOC at Alice Holt with lower permeability of the surface water gley leading to reduced potential for leaching losses to lower soil depths. Furthermore, the high silt (particle size 0.063-0.002 mm) and clay (particle size < 0.002 mm) content (69% and 10% in volume, respectively; Villada 2013) of the sampled A horizon at Alice Holt may have promoted retention of DOC via mineral sorption. The potential for physical protection of DOC through sorption to soil minerals would be, in comparison, limited in the peat soils at Kielder (Schmidt et al. 2011). As a result of the field sampling design, the decay class of deadwood that was sampled within each stand was nested within stand age, reflecting the dominant stage of decay found for each stand age. We initially hypothesized that the effects of deadwood on the soil DOC pool might depend on stand age (decay class) due to differences in DOC fluxes from wood at different stages of decay. Wood in the late stages of decay, i.e. the old stands, is thought to hold a larger fungal and microbial biomass (Kueffer et al. 2008; Baldrian et al. 2016) which has potential to increase rates of decomposition (Peršoh and Borken 2017). However, this was not seen in our results as the deadwood effect on WEOC was of similar magnitude at Alice Holt, irrespective of stand age (p = 0.900).

Deadwood influence on soil C cycle processes linked to WEOC concentration and quality

Microbial priming occurs when fresh inputs of organic matter into the soil, and the subsequent increase in available labile carbon, leads to increased rates of microbial activity stimulating the decomposition of already stored C in the system (Fontaine et al. 2003; Beverly and Franklin 2015). It was possible that the additional inputs of C from deadwood might prime depolymerisation of existing SOM pools (Peršoh and Borken, 2017; Minnich et al. 2021), potentially leading to increased WEOC through DOC production. If deadwood DOC-induced microbial priming was occurring in our forest soils, we might expect to find an enhanced decomposition rate, seen in the TBI results, and enhanced potential enzyme activity in the soil under deadwood. In a previous, long-term experiment at Kielder, higher mineralisation and loss of C was found when brash (branches and litter) was left in situ compared to sites where it was removed, suggesting a likely priming effect (Vanguelova et al. 2010). It is evident at Alice Holt that WEOC concentrations are elevated under deadwood; however, there is no indication through analysis of extracellular enzyme activity or the TBI that this additional soluble C is priming microbial decomposition or the production of extracellular enzymes involved in decomposition. One possible explanation for the lack of a deadwood effect on decomposition is that the k parameter in the TBI may not be sensitive to priming effects since k is estimated from the decomposition of added rooibos tea litter (Keuskamp et al. 2013), which in itself represents an input of fresh organic matter, rather than pre-existing SOM. The lack of an effect of the increased WEOC beneath deadwood on potential enzyme activity might relate to the quality of deadwood DOC inputs, as discussed below.

Optical methods are commonly used to assess the quality of DOC in surface waters, and their potential application to the study of carbon cycling in soils, through characterization of the WEOC, could add to interpretation with respect to the impact of the 'extra' WEOC in soils under the influence of deadwood at Alice Holt. As previously explained (Sect. 2.4 and 3.2), SUVA₂₅₄ (Hansen et al. 2016) and a HIX (Ohno 2002) provide a measurement of the aromatic content and degree of humification of WEOC, whereby larger values indicate a larger aromatic content with increased levels of humification related with higher levels of aromatic C (Zech et al. 1997; Ji et al. 2015). The HIX indicated a greater level of humification and aromaticity, and potentially stability of WEOC, for Alice Holt soils under deadwood (HIX = 0.75) compared to those under leaf litter only (HIX = 0.65). Consistent with this deadwood signature in soil WEOC, Bantle et al. (2014) have shown both Quercus sp. and Picea abies -derived DOM to be a more significant source of humified material (HIX = 9.3 and 7.5, respectively) than throughfall (HIX = 3.8). SUVA₂₅₄ also suggested stronger aromaticity for WEOC underneath deadwood although statistical evidence for this was weaker. Amongst the many factors suggested to influence the size and direction of priming effects, the chemical structure of soluble C inputs has been shown to be important (Di Lonardo et al. 2017). Aromatic compounds (e.g. vanillic acid) have been shown to induce the most pronounced priming effects when compared to non-aromatics (e.g. saccharides) on an energy content basis, possibly because they resemble compounds present in more stable SOM (Di Lonardo et al. 2017). In contrast, the aromatic compounds catechin and caffeic acid (a key intermediate in the biosynthesis of lignin) have been shown to have no effect on potential enzyme activity (Zwetsloot et al. 2020). Due to the relatively energy-poor nature of these aromatic monomers, greater concentrations are needed to produce effects on a carbon concentration equivalent basis (Di Lonardo et al. 2017) and therefore it is possible that the aromatic enrichment of the WEOC pool under deadwood at Alice Holt, as suggested by HIX and SUVA₂₅₄, was not of sufficient magnitude to produce detectable effects on corresponding enzyme potentials.

An alternative mechanism to microbial priming by which inputs of soluble organic matter might impact on soil organic matter pools has been suggested. This involves the mobilization and solubilisation of organic C previously intimately associated with reactive mineral phases through dissolution and metal chelating reactions mediated by inputs of organic compounds that are ligands with metal complexing activities (Keiluweit et al. 2015). If this mechanism was in operation in the Alice Holt soils, the 'extra' WEOC under deadwood would be of both deadwood and soil organic matter origin. The origin of WEOC, as inferred from the FI of ~1.9 indicated that the WEOC was a mixture of both microbial and plant origin (McKnight et al. 2001) but this did not differ between soils from under deadwood and those under only leaf litter. Whether deadwood-derived DOC contains compounds that are more active as ligands in mobilizing additional soluble C from mineral associations than those in leaf litter-DOC requires further investigation. At this stage, it is not possible to conclude if elevated WEOC under deadwood at Alice Holt was solely a consequence of stabilized DOC from deadwood or if abiotic priming also contributed.

Conclusions

The presence of deadwood significantly elevated (by ~ 1.5 to ~ 1.75 times) the concentrations of soil DOC, analysed as WEOC, in the mineral horizon of a surface water gley chronosequence under lowland broadleaf (Q. robur) forest. We believe that this increase in WEOC was, at least in part, due to deadwood acting as a significant input of DOC, released as a result of decomposition processes, to the underlying soil, which was subsequently retained through interactions with the soil mineral horizon. In highly organic soils sampled from an upland Sitka spruce chronosequence, effects of deadwood on WEOC were not detectable. This is potentially because inputs of DOC from

deadwood were masked by the already high background of WEOC and/or did not persist in soil due to low potential for mineral sorption. There was no evidence, from TBI decomposition parameters or potential extracellular enzyme data, that deadwood-derived DOC impacted other forest carbon pools via microbial priming effects, possibly due to the aromatic quality of DOC produced. It was not possible to determine whether the increased WEOC under deadwood at the lowland oak site was solely due to retention of deadwood-derived DOC, or, if OC solubilized from existing SOC pools via deadwood DOC-mediated abiotic reactions also contributed. At present, the source and impacts on soil C cycling of elevated WEOC associated with leaving deadwood in situ requires further characterization. Forest carbon budgets, particularly those for mineral soils, that are only derived from DOC concentrations sampled from under leaf litter may be an underestimation; efforts should be made to include a deadwood component in temporal and spatial monitoring of forest ecosystems.

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Data availability Not applicable.

Code availability Not applicable.

Declarations

Conflict of interest Authors declare that they have no conflict of interest.

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