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Influence of tree species on carbon and nitrogen transformation patterns in forest floor profiles

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Abstract

• **Background** Among forest management practices, forest tree species substitution influences biogeochemical cycles and soil interactions rapidly (decades) and significantly.

• **Methods** We studied the impact of tree species on forest floor properties in 27-year-old monospecific stands of oak, beech, spruce and Douglas-fir, grown in common garden.

• **Results** In spite of similar litterfall C-inputs (1.3–1.8 Mg ha⁻¹ yr⁻¹), C-stocks strongly differed between coniferous (16–21 Mg ha⁻¹) and broadleaved (7–9 Mg ha⁻¹) forest floors. The large C-mineralization under oak (196 C-μg C_{total}-g⁻¹ d⁻¹) relatively to other stands (125–147 C-μg C_{total}-g⁻¹ d⁻¹) and the large dissolved organic carbon release under broadleaved forest floors may explain this difference. Spruce forest floor exhibited the lowest net nitrification and the largest contribution of organic anions to the anionic charge of aqueous leachate (62.5 %±1.7). Conversely, Douglas-fir forest floor exhibited the largest relative nitrification and the largest contribution of nitrate to the anionic charge (70.8 %±0.7).

• **Conclusion** We conclude that tree species induce a specific signature in term of proton donors at the forest floor level (organic vs nitric acid), which involves potential impacts on pedogenetic processes in the mineral horizon beneath.

Keywords DOC and CO₂ release · Forest floor horizons · N-release · Common garden · Tree species

1 Introduction

The influence of forest tree species on biogeochemical cycles and soil interactions has for a long time attracted numerous scientists, and has been covered by an intensive literature. The interest in this research topic originates from the substitution issue of natural vegetation by monospecific coniferous stands as a cause of soil acidification (Binkley and Giardina 1998). More recently, with the Kyoto protocol, the interest has been rather focused on tree species potential for mitigation of greenhouse gases and for carbon sequestration (Jandl et al. 2007).

Soil acidification and carbon sequestration are influenced by forest floor decomposition, through the release of organic/inorganic acids and dissolved organic carbon (DOC) (Guggenberger and Kaiser 1998). The impact of forest tree species on forest floor properties has been largely studied particularly in common garden. This experimental design makes it possible to minimize confounding effects such as local climate, geology, soil type, forest stand age, or past land use management (Binkley and Giardina 1998; Meentemeyer 1978) by simultaneous plantations of different tree species in homogeneous pedo-climatic conditions (Gower and Son 1992; Vesterdal et al. 2008). Previous studies in common garden highlighted the following points: (i) tree species can modify topsoil N-mineralization rates in a short time (decades) (Gower and Son 1992), (ii) forest floors exhibit differences in C/nutrient ratios, pH and nutrient contents according to the tree species (Vesterdal and Raulund-Rasmussen 1998), (iii) tree species influence microbial and faunal populations through variation in litter

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lignin and Ca concentration (Hobbie et al. 2006), (iv) the influence of tree species on C and N concentration and C/N ratio is more pronounced in the forest floor than in the litterfall and the topsoil because of large differences of turnover rates (Vesterdal et al. 2008), and (v) Hansen et al. (2009) conclude that tree species do not influence the amount of litterfall while site-related factors do, and that a large variability in forest floor accumulation could primarily be due to a difference in litter decomposition.

However, the impact of forest tree species on the release of soluble compounds by the forest floor has not yet been explored particularly for the Douglas-fir, despite the crucial involvement of these compounds in pedogenetic processes. Moreover, the forest floor is not a *black box*. Forest floor has a heterogeneous composition and can be divided into different layers (horizons) corresponding to successive stages of decomposition (Wardle 1993). Given that previous studies concluded that the main difference between tree species is decomposition rate (Hansen et al. 2009; Vesterdal et al. 2008), studying forest floor profiles should provide a clearer picture of the tree species decomposition process.

The experimental site of Breuil-Chenue in the Morvan mountains (France) is a common garden set up in 1976 specifically to compare biogeochemical cycles between tree species. On this site, no studies have particularly yet considered the forest floor compartment. Nevertheless, Moukouri et al. (2006) have already studied organic matter biodegradability and mineral nutrient availability in the topsoil, while Zeller et al. (2007) have considered N-transformations in the litterfall and the topsoil.

In this paper, our specific objective is to characterize the release of dissolved products (NH_4^+ , NO_3^- , DON, DOC, cations and anions) in forest floor profiles of four trees widespread in Europe: Douglas-fir (*Pseudotsuga menziesii* Franco), Norway spruce (*Picea abies* Karst), European beech (*Fagus sylvatica* L.) and oak *Quercus petraea* (Matt.) Liebl. To support our understanding, we provide a detailed characterisation of the forest floor layers (composition, C- and N-transformation). To achieve this, forest floor horizon samples were collected in the site of Breuil, individually incubated in vitro for determination of C- and N-mineralization and leached for measuring the release of soluble compounds.

2 Materials and methods

2.1 Experimental site

The study was carried out on forest floors sampled in the experimental site of Breuil-Chenue set up by the Institut National de la Recherche Agronomique (INRA) in the Natural Park of the Morvan, France (47°18'10" N;

4°4'44" E). The site contains two adjacent blocks with replicated stands. It is established on a plateau at 650 m. Climate is submountain Atlantic, with 1400 mm mean annual rainfall and 6°C mean annual temperature. The native forest was a 150-year-old coppice composed mainly of beech (*Fagus sylvatica* L.) and oak *Quercus petraea* (Matt.) Liebl. In 1976, this initial forest was partially clear-felled (one stand was conserved in each block) and replaced by mono-specific 0.1 ha stands of different species (one plot by block) (Bonneau et al. 1977). Previous studies showed that the site heterogeneity was minor (Bonneau et al. 1977; Ranger et al. 2004). Out of the plantations of block 1, we selected the four tree species studied here.

The soil is an acid brown soil, and classified as an Alocrisol (AFES 1998) or an alumnice Cambisol (IUSS 2006). It developed from a granite which was very poor in cation-bearing minerals (0.5% MgO; 0.6% CaO; 4.4% K₂O). The soil is strongly acid, pH is 3.8 in surface horizons and increases with depth (4.7–4.8). Soil texture is sandy-loamy throughout the whole profile, with 40–55% sand and 15–20% clay. Carbon concentrations decrease from 0.44 g kg⁻¹ in surface horizons to 0.04 g kg⁻¹ in deeper horizons (Bonneau et al. 1977).

2.2 Litterfall and forest floor sampling

Five litter-traps (0.5 m²) were placed in each stand to collect litterfall continuously. Sampling occurred every 3 months for 3 years between September 2002 and September 2005. Each sample was sorted and dried. All samples of leaf/needle litterfall were considered in order to assess the average annual C-input by litterfall and its variability. For each forest stand, composite samples were analysed for C, N, phosphorus (P), lignin, cellulose and cations concentrations. The composite samples were constituted with leaf/needle litterfall samples from all periods, applying weighting factors according to the dry weight of corresponding litterfall sample.

In each forest stand, we sampled forest floor horizons in December 2003 after leaf fall. We sampled separately Oln (plant remains), Olv (poorly fragmented), Of (fragmented) sub-horizons according to the definitions of Brêthes et al. (1995). The material was collected quantitatively on 1 m² surface representing forest floor morphology in the plot. An Oh horizon (dark humified organic matter) was collected under spruce, whereas transitional horizons were collected in other stands: Of-Oh under oak and Douglas-fir and Of-Ah under beech. The symbol O* is used in the following text to identify the deepest horizon of the forest floor, which is either Oh or Of-Oh or Of-Ah.

2.3 Analysis of litterfall and forest floor horizons

Carbon and N concentrations were measured by gas chromatography after dry combustion (1800°C) with a Thermo Quest CHN autoanalyzer. We measured the concentration of lignin and cellulose in litterfall by acid-detergent extraction (Van Soest 1963). Elemental analyses (Ca, Mg, K, Na, Mn, Al and Fe) were carried out by calcination (450°C) and digestion with HNO₃, HF and HClO₄, and measured by induced plasma coupled atomic emission spectroscopy (ICP-AES, Jarell Ash). Horizon thicknesses were estimated in the field. Root colonization expressed by the volume percentage taken up by roots in each horizon was visually estimated both in the field and during the sample sorting.

2.4 Incubation, leaching and chemical analysis of leachates

Forest floor materials dedicated to incubation were neither dried nor rewetted before the constitution of columns. Incubation was carried out in PVC-columns with perforated bottom covered by polyamide net (50- μ m mesh size). Each column (diameter = 115 mm, height = 200 mm) contained 15 g (equivalent dry weight) of one horizon. All the columns were covered, but not hermetically closed, and pre-incubated during 6 weeks at 4°C. At the end of the pre-incubation period, each column was leached with MilliQ water (150 ml).

We introduced two plastic cups into each column: one with 1 M NaOH for CO₂ measurement and one with water to balance NaOH-related dehydration. The columns were then closed hermetically. Initial CO₂ content was measured in a closed column without sample (blank column). Columns of separated horizon were incubated in triplicate for 32 days at 4°C. At the end of the incubation period, columns were leached again with MilliQ water (150 ml, with a 1:10 ratio between organic material and water quantity). The leachates were filtered through 0.45- μ m pore size Gellman filter and analysed.

We measured pH with CG804 pH-meter, ammonium (NH₄⁺) concentration by FIA (Alpkem FS3000), cation concentrations (Na⁺, K⁺, Ca²⁺, Mg²⁺, Al³⁺, Fe²⁺, Mn²⁺, Cu²⁺, Pb²⁺, Zn²⁺) by ICP-AES, and inorganic anion concentrations (F⁻, Cl⁻, NO₃⁻, NO₂⁻, PO₄³⁻, SO₄²⁻) by ionic chromatography (HPLC, Dionex LC20 with ION-PAC AS 11 column). The anionic deficit charge (ACD) was calculated by the difference between the sum of cationic charge and the sum of anionic charge. The concentrations of dissolved organic carbon (DOC) and total dissolved nitrogen (TDN) were measured with a total organic carbon analyser (SHIMADZU TV-V_{CSH-CSN}). The concentration of dissolved organic nitrogen (DON) was calculated by the difference between TDN and dissolved

inorganic nitrogen (DIN), which was defined by the sum of NH₄⁺, NO₂⁻ and NO₃⁻ concentrations. The release of DOC and DON (C_{DOC}-mg C_{total}-g⁻¹ or N_{DON}-mg N_{total}-g⁻¹) was defined by the ratio between the amount of DOC or DON in the leachate after incubation, and the initial amount of C or N in the corresponding sample.

2.5 C- and N-mineralization

The production of CO₂ was measured every 10 or 11 days by back titration of NaOH with HCl 1 M and deduction of the values from blank columns. C-mineralization was defined by the ratio between the average daily CO₂ production and the initial amount of C in the corresponding sample.

Inorganic N (N- μ g N_{total}-g⁻¹ d⁻¹) was extracted from 2.5 g (equivalent dry weight) sub-sample from column by using KCl 1 M (2.5 g: 100 ml), before and after incubation. NO₃⁻ and NH₄⁺ concentrations in KCl extracts were measured by HPLC (Dionex LC20 with IONPAC AS 11 column) and flow injection analysis (FIA, Alpkem FS3000) respectively.

Net nitrification (N- μ g N_{total}-g⁻¹ d⁻¹) was defined by the ratio between (1) net NO₃⁻ production, which is the difference between final and initial NO₃⁻ concentrations in the KCl extracts, and (2) initial amount of N in the sample. Relative nitrification (%) is defined by the ratio between (1) final NO₃⁻ in the KCl and water extracts, and (2) final mineral N (NH₄⁺ and NO₃⁻) in the KCl and water extracts. It is an alternative measure of nitrification considering NO₃⁻ production in relation to NH₄⁺-supplying capacity (Robertson 1982).

DOC and DON releases, C-mineralization and net nitrification are all defined in function of the initial amount of C or N in the corresponding sample. Using these units facilitates comparison between species, since organic matter has a different composition between species and horizons.

2.6 Statistical analysis

The differences between means were compared with one-way analysis of variance (ANOVA1) and Fisher's least significant difference (LSD) test, with a significant level of 0.05. All statistical analyses were performed with the SAS software (SAS Institute 1999). We compared the means between the four tree species, for C- and N-mineralization and for the release of DOC, DON, NH₄⁺ and NO₃⁻ in each horizon. Furthermore, we compared in each forest floor the means of all sub-horizons weighted by their dry mass for C- and N-mineralization and for the release of DOC, DON, NH₄⁺ and NO₃⁻. We also performed some correlation analysis between litterfall and forest floor characteristics and C-mineralization using the SAS software.

3 Results

3.1 Forest floor morphology

The forest floor, which encompasses OI, Of and Oh horizons, was thicker under spruce (40 mm) than under the other tree species (Douglas-fir, 25 mm; beech, 20 mm; oak, 15 mm). The Oh horizon was thick in the spruce forest floor (10–20 mm), while it was thin (< 2 mm) and discontinuous in Douglas-fir and oak forest floors. Oh was absent under beech, and we observed mineral particles from Ah horizon in the lower part of Of horizon.

Tree roots colonized Of and Oh horizons whatever the forest floor, but mainly under spruce and beech. They were particularly abundant in the Oh horizon of spruce forest floor, where they represented about 25–30 % of the horizon volume. In the other forest floors, we only observed fine roots (1–4 mm) which represented less than 5 % of the volume. Root tips were covered by visible ectomycorrhizal fungi except under Douglas-fir forest floor. The presence of white fungal mats was visually detected in the four forest floors, mainly in Olv.

3.2 Composition of litterfall

The composition of litterfall composite samples is presented in Table 1. C-concentrations in litterfall were in the same range of values (47–48 %) under the four species. N-concentrations significantly differed between tree species,

and were in the order: spruce<oak<beech<Douglas-fir ($P<0.0001$). P-concentration was the lowest in oak litterfall and the highest in Douglas-fir litterfall ($P<0.0001$). C:N ratios in litterfall significantly differed between tree species, and were in the order: Douglas-fir<beech<oak<spruce ($P<0.0001$). C:P ratio was the lowest in Douglas-fir litterfall and the highest in oak litterfall ($P<0.0001$). Lignin concentrations in litterfall were similar under oak, spruce and Douglas-fir (31–34 %) but much higher under beech (54 %, $P=0.0069$). The concentrations of Ca, Mg and Mn were the highest in Douglas-fir and oak litterfalls and the lowest in spruce litterfall. Beech litterfall exhibited the highest concentrations of K ($P<0.0001$) and Na ($P<0.0001$). Douglas-fir litterfall exhibited the highest concentrations of Al ($P<0.0001$) and Fe ($P=0.0421$).

3.3 Composition of forest floor horizons

The elemental composition of forest floor horizons is presented in Table 2. Whatever the tree species, C-concentration, C:N and C:P ratio decreased with the horizons, N- and P-concentration reached a maximum in Of horizon. The values of C:N ratio in OI horizons were higher under oak and spruce than under other species. The lowest value of C:P ratio was observed under beech whatever the forest floor horizon. Whatever the tree species, K, Na, Al and Fe concentrations exhibited the same pattern of increase with the horizons. For Mg and Mn

Table 1 Composition of the tree species litterfalls. Means with different associated letters are significantly different from one another at the $P<0.05$ level according to the Fisher's least significant difference test ($n=3$)

	C %	N %	P ‰	C:N atomic ratio	C:P atomic ratio	Lignin %	Cellulose %	Ca g kg ⁻¹	Mg g kg ⁻¹	K g kg ⁻¹	Na g kg ⁻¹	Mn g kg ⁻¹	Al g kg ⁻¹	Fe g kg ⁻¹
Oak														
Mean	47	1.13	0.45	48	2684	34	26	4.0	0.93	2.3	0.08	2.3	0.13	0.11
SD	1	0.02	0.01	1	63	3	1	0.2	0.05	0.1	0.00	0.1	0.01	0.01
	a	c	c	b	a	b	a	b	a	c	c	b	c	b
Beech														
Mean	47	1.33	0.58	42	2095	54	21	3.2	0.58	3.5	0.15	2.0	0.16	0.11
SD	1	0.01	0.01	1	4	8	6	0.1	0.1	0.1	0.01	0.1	0.02	0.01
	a	b	b	c	b	a	a	c	b	a	a	c	c	ab
Spruce														
Mean	47	0.98	0.57	55	2128	33	25	3.0	0.54	2.6	0.12	1.7	0.28	0.09
SD	1	0.03	0.03	1	96	4	8	0.2	0.09	0.1	0.01	0.1	0.01	0.01
	a	d	b	a	b	b	a	c	b	b	b	d	b	b
Douglas-fir														
Mean	48	1.46	0.72	38	1713	31	25	6.4	0.83	2.4	0.09	2.7	0.54	0.15
SD	1	0.04	0.01	1	22	4	6	0.1	0.08	0.1	0.00	0.1	0.07	0.03
	a	a	a	d	c	b	a	a	a	c	c	a	a	a
<i>P</i>	0.0022	< 0.0001	< 0.0001	< 0.0001	< 0.0001	0.0069	0.7692	< 0.0001	0.0016	< 0.0001	< 0.0001	< 0.0001	< 0.0001	0.0421

Table 2 Elemental composition of forest floor horizons and whole forest floor. O* is the deepest horizon of the forest floor which is either Oh, Of-Oh or Of-Ah. The values for whole forest floor werecalculated using the dry weight of each horizon/sub-horizon as weighting factor (weighted average values — *wav*)

Horizon	Species	C %	N %	P ‰	C:N atomic ratio	C:P atomic ratio	Ca g kg ⁻¹	Mg g kg ⁻¹	K g kg ⁻¹	Na g kg ⁻¹	Mn g kg ⁻¹	Al g kg ⁻¹	Fe g kg ⁻¹
Oln	oak	47	1.2	0.46	44	2647	5.5	1.1	2.2	0.1	3.3	1.2	0.5
	beech	50	1.6	0.71	36	1791	3.1	0.5	2.7	0.1	1.5	0.6	0.3
	spruce	48	1.2	0.36	48	3476	3.8	0.3	1.2	0.0	1.7	0.4	0.1
	Douglas-fir	50	1.5	0.47	38	2764	5.5	0.7	1.3	0.1	2.2	0.5	0.2
Olv	oak	47	1.4	0.58	38	2103	4.2	0.9	2.3	0.2	2.4	2.0	0.6
	beech	47	2.2	0.83	25	1465	4.0	0.6	2.3	0.3	2.0	2.0	0.6
	spruce	45	1.5	0.58	35	2001	4.3	0.4	1.3	0.1	1.9	1.0	0.3
	Douglas-fir	51	2.5	0.72	24	1817	5.7	0.7	1.4	0.1	2.4	1.5	0.7
Of	oak	41	1.9	0.80	25	1312	5.0	0.8	4.2	0.7	3.9	6.7	2.3
	beech	42	2.2	0.93	22	1166	4.1	0.8	4.4	1.2	2.5	11.2	3.1
	spruce	42	1.8	0.70	28	1560	3.2	0.5	3.9	0.7	2.3	6.9	2.1
	Douglas-fir	44	2.3	0.79	22	1435	3.3	0.5	6.3	1.0	1.8	7.9	1.9
O*	oak	30	1.5	0.73	23	1063	2.6	1.0	12.5	2.4	2.1	24.8	6.6
	beech	21	1.0	0.62	25	884	1.9	1.0	18.6	4.9	0.8	36.6	8.6
	spruce	33	1.6	0.73	25	1180	1.9	0.7	10.9	2.3	1.1	19.1	4.9
	Douglas-fir	31	1.7	0.64	21	1240	1.3	0.6	10.5	2.3	0.3	16.2	5.6
Forest floor <i>wav</i>	oak	37	1.6	0.67	27	1435	4.0	0.9	7.2	1.3	2.9	13.1	3.7
	beech	37	1.6	0.77	26	1221	3.1	0.7	8.9	2.2	1.6	17.1	4.2
	spruce	41	1.5	0.57	32	1875	3.1	0.5	5.0	1.0	1.8	8.2	2.2
	Douglas-fir	42	2.1	0.66	24	1628	3.5	0.6	5.9	1.0	1.5	8.2	2.7

concentrations, the differences between species were the highest in Ol horizons.

Weighted average values for whole forest floor (symbolized by the expression *wav* in the following text) were calculated using the dry weight of each horizon / sub-horizon as weighting factor. The *wav* of C:N ratio for forest floor were the highest under spruce (32 %) and the lowest under Douglas-fir (24 %). The *wav* of C:P ratio were the lowest under beech (1221) and the highest under spruce (1875). Beech forest floor presented the highest *wav* of K, Na, Al and Fe concentrations. Oak forest floor exhibited the highest *wav* of Ca, Mg and Mn concentrations.

3.4 Composition of forest floor leachates

3.4.1 pH, C- and N-release

Douglas-fir leachates strongly decreased in pH with the horizons (Table 3). Whatever the forest floor, the greatest DOC release was observed in Olv horizon. The *wav* of DOC release significantly differed between tree species in the order:

Douglas-fir, $1.83 \text{ ‰} \pm 0.08 < \text{spruce}, 2.72 \text{ ‰} \pm 0.09 < \text{beech}, 4.30 \text{ ‰} \pm 0.07 < \text{oak}, 5.96 \text{ ‰} \pm 0.66$; $C_{\text{DOC}}\text{-mg } C_{\text{total}}\text{-g}^{-1}$ ($P < 0.0001$). Whatever the forest floor, DON release ($N_{\text{DON}}\text{-mg } N_{\text{total}}\text{-g}^{-1}$) was the highest in Olv horizon leachates, while NH_4^+ release ($N\text{-mg } N_{\text{total}}\text{-g}^{-1}$) peaked in Of except for beech forest floor (Olv) (Table 3).

The contribution of N forms (DON, NH_4^+ or NO_3^-) to the total dissolved N-release in leachates changed with the horizons; globally, most of the dissolved N was released as DON in upper horizon leachates and as NO_3^- in lower horizon leachates (Fig. 1).

The *wav* of DON release significantly differed between tree species (Douglas-fir, $0.60 \pm 0.44 < \text{spruce}, 1.99 \pm 0.07 < \text{beech}, 2.97 \pm 0.35 < \text{oak}, 3.75 \pm 0.31$; $N_{\text{DON}}\text{-mg } N_{\text{total}}\text{-g}^{-1}$; $P < 0.0001$). The *wav* of NH_4^+ release significantly differed between the broadleaf and coniferous species (oak, 3.90 ± 0.09 ; beech, $3.91 \pm 0.32 > \text{spruce}, 2.25 \pm 0.09$; Douglas-fir, 2.57 ± 0.37 ; $N\text{-mg } N_{\text{total}}\text{-g}^{-1}$; $P < 0.0001$). Also, the *wav* of NO_3^- release significantly differed between tree species (Douglas-fir, 3.10 ± 0.39 ; oak, $2.78 \pm 0.16 > \text{beech}, 1.19 \pm 0.11 > \text{spruce}, 0.24 \pm 0.04$; $N\text{-mg } N_{\text{total}}\text{-g}^{-1}$; $P < 0.0001$).

Table 3 DOC, DON, NH_4^+ and NO_3^- in forest floor horizon leachates. O* is the deepest horizon of the forest floor which is either Oh, Of-Oh or Oh-Ah. For each horizon, means with different

associated letters are significantly different from one another at the $P < 0.05$ level according to the Fisher's least significant difference test ($n=3$)

Horizon	Species	pH		DOC			DON		NH_4^+		NO_3^-					
				[C _{DOC} -mg C _{total} -g ⁻¹]			[N _{DON} -mg N _{total} -g ⁻¹]		[N-mg N _{total} -g ⁻¹]		[N-mg N _{total} -g ⁻¹]					
		Mean	SD	Mean	SD		Mean	SD	Mean	SD	Mean	SD				
Oln	oak	4.0	0.0	d	8.29	1.31	a	5.57	0.74	a	0.28	0.40	ab	0.00	0.00	c
	beech	4.8	0.2	c	3.45	0.53	b	2.35	0.31	b	0.09	0.04	b	0.02	0.04	bc
	spruce	5.3	0.1	b	1.52	0.20	c	2.06	0.20	b	0.18	0.04	b	0.08	0.01	b
	Douglas-fir	6.1	0.1	a	0.78	0.04	c	1.60	0.03	b	0.66	0.12	a	0.15	0.06	a
		$P < 0.0001$			$P < 0.0001$			$P < 0.0001$			$P = 0.0479$		$P = 0.0036$			
Olv	oak	4.4	0.1	c	12.12	3.58	a	9.81	3.16	a	2.44	1.74	b	0.13	0.01	bc
	beech	4.7	0.1	b	8.10	0.60	b	6.12	0.56	b	6.35	0.79	a	0.28	0.12	b
	spruce	4.5	0.0	c	3.09	0.07	c	2.63	0.17	c	0.52	0.09	c	0.03	0.04	c
	Douglas-fir	5.3	0.0	a	2.95	0.31	c	2.01	0.14	c	2.81	0.51	b	0.90	0.20	a
		$P < 0.0001$			$P = 0.0007$			$P = 0.0026$			$P = 0.0006$		$P < 0.0001$			
Of	oak	4.7	0.1	a	4.99	0.66	a	3.38	0.64	a	8.21	0.78	a	4.70	0.01	b
	beech	4.6	0.1	ab	4.84	0.37	a	2.77	0.42	ab	4.82	0.63	b	1.03	0.12	c
	spruce	4.5	0.0	b	3.89	0.53	b	2.39	0.44	b	3.78	0.11	b	0.35	0.00	c
	Douglas-fir	4.2	0.1	c	2.54	0.54	c	1.11	0.80	c	5.24	1.77	b	5.97	1.13	a
		$P < 0.0001$			$P = 0.0002$			$P = 0.0087$			$P = 0.0038$		$P < 0.0001$			
O*	oak	4.4	0.1	a	1.04	0.15	b	1.28	0.16	a	2.70	0.13	a	3.54	0.12	a
	beech	4.4	0.1	a	1.35	0.34	ab	0.83	0.03	b	2.89	0.50	a	3.29	0.48	a
	spruce	4.4	0.0	a	1.65	0.15	a	1.02	0.14	b	1.84	0.24	b	0.27	0.11	b
	Douglas-fir	3.7	0.1	b	0.27	0.07	c	0.38	0.07	c	0.09	0.04	b	3.48	0.68	a
		$P < 0.0001$			$P = 0.0002$			$P < 0.0001$			$P < 0.0001$		$P < 0.0001$			

3.4.2 Anionic signature

The contribution of each anion to the total anionic charge widely differed with the horizons (Fig. 2) and between species (Fig. 3). Whatever the tree species, we observed a strong dominance of organic anions in Ol leachates (65–92 %), except in the leachates of Douglas-fir Olv (38 %±10). Most of anions was released as NO_3^- in Of and O* leachates of oak, beech and Douglas-fir (65–85 %).

In the whole forest floor (*WAV*) (Fig. 3), NO_3^- strongly dominated in Douglas-fir leachates (71 %±1) while it was organic anions in spruce leachates (62 %±2). In oak and beech leachates, both NO_3^- and organic anions contributed largely to the anionic charge, but the contribution of NO_3^- was larger for oak than for beech (41 %±4 vs 30 %±3; $P < 0.0001$).

3.5 C- and N-mineralization in forest floor horizons and whole forest floors

Whatever the tree species, C-mineralization (C- $\mu\text{g C}_{\text{total}}\text{-g}^{-1}\text{d}^{-1}$) strongly decreased from Ol to O* (Fig. 4). The *WAV* of C-mineralization computed for each forest floor (Table 4) was

the largest in the oak forest floor ($P < 0.0001$). By contrast, we observed only minor differences of C-mineralization between beech, spruce and Douglas-fir forest floors.

Whatever the tree species, the relative nitrification (%) increased from Of to O* (Fig. 5). However, this increase was very low under spruce in comparison with the other species. In Oln, relative nitrification (%) did not significantly differ between tree species (Table 4). In Olv, Of and O*, the highest values of relative nitrification were observed under Douglas-fir, whereas the lowest values were observed under spruce (Olv, $P = 0.1386$; Of, $P < 0.0001$; O*, $P < 0.0001$). The *WAV* of relative nitrification significantly decreased in the following order: Douglas-fir > oak > beech > spruce ($P < 0.0001$).

4 Discussion

Litter decomposition is mainly affected by climate, soil and vegetation (Meentemeyer 1978). Here, the four forest tree stands developed in homogeneous conditions of soil and climate (Ranger et al. 2004). The differences of forest floor properties are closely linked with tree species.

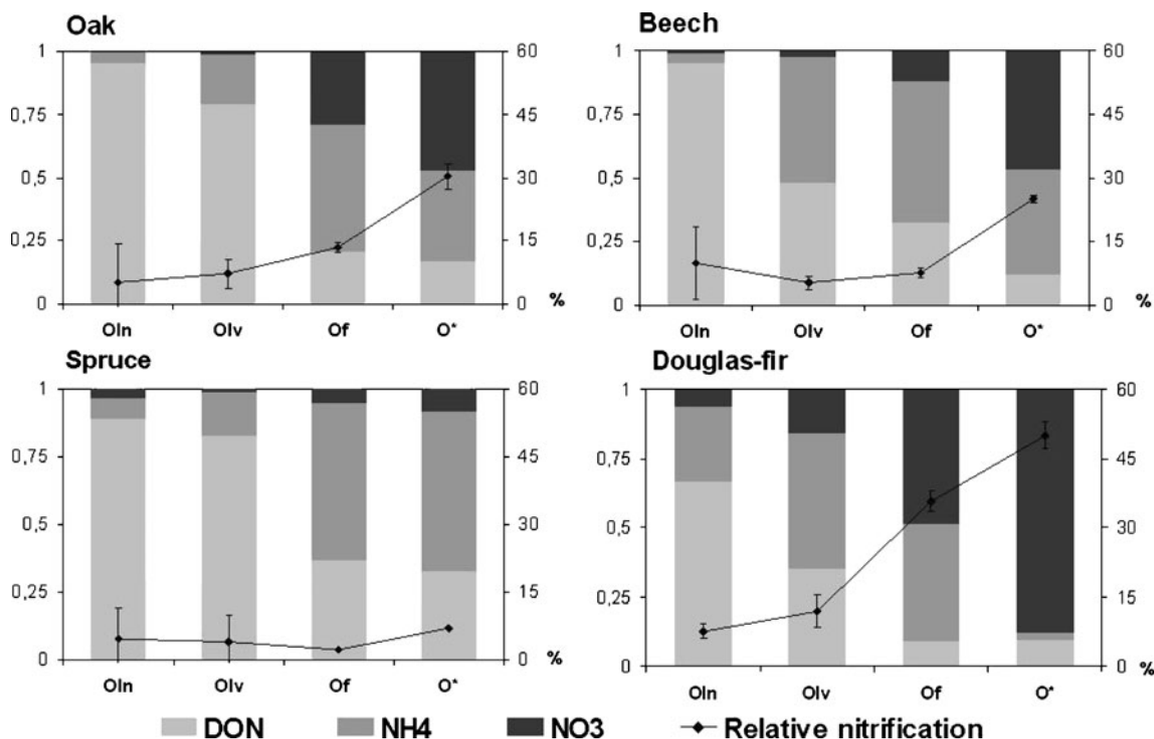


Fig. 1 Proportional contributions of DON, NH_4^+ and NO_3^- to total N release in forest floor horizon leachates (left Y-axis) and relative nitrification (%) in forest floor horizons (right Y-axis, mean values are reported \pm SD)

4.1 Impact of tree species on litterfall quality

As observed by Alban (1982), litterfall composition is significantly affected by tree species, and might directly affect

the composition of the whole forest floor. The concentrations of K, Na and Mg in the forest floor were positively correlated to K, Na and Mg concentrations in litterfall ($r=0.71, 0.72$ and 0.69 respectively). There was also a strong correlation

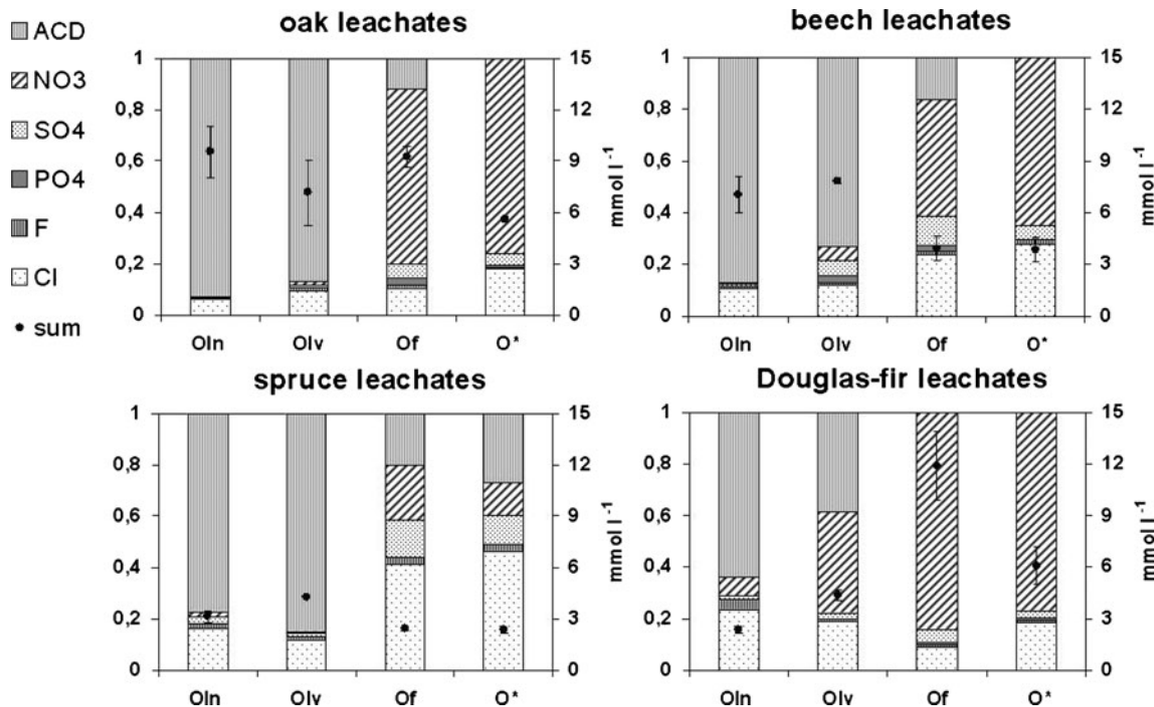


Fig. 2 Proportional contributions of anions (ACD^* , NO_3^- , SO_4^{2-} , PO_4^{3-} , F^- and Cl^-) in forest floor horizons leachates (left Y-axis) and sum of anions in mmol.l^{-1} (right Y-axis, mean values are reported \pm SD). (*) ACD is the “deficit in anionic charge”

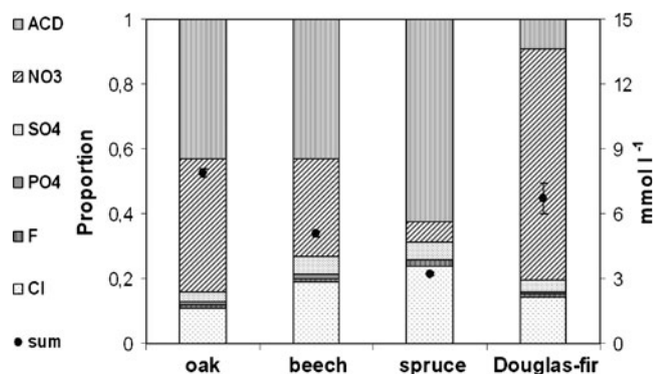


Fig. 3 Proportional contributions of anions (ACD, NO_3^- , SO_4^{2-} , PO_4^{3-} , F^- and Cl^-) in the whole forest floors leachates (left Y-axis) and sum of anions in mmol.l^{-1} (right Y-axis, mean values are reported \pm SD). The values for the whole forest floor were calculated using the dry weight of each horizon / sub-horizon as weighting factor (weighted average values - *WAV*)

($r=0.97$, $P=0.0368$, $n=4$) between C:N in forest floors and corresponding litterfalls. However, Vesterdal et al. (2008) studied six tree species including all of ours except Douglas-fir in a common garden, and found no correlation between C:N of forest floors and corresponding litterfalls. The particularly low C:N in Douglas-fir forest floor and litterfall could explain this divergence. Vesterdal and Raulund-Rasmussen (1998) found also C:N ratio in Douglas-fir forest floor significantly lower than in six other tree species forest floors on average on seven sites of different nutrient status.

Vesterdal and Raulund-Rasmussen (1998) concluded that differences in forest floor nutrient status may not only reflect differences in litter nutrient status, and that the initial nutrient status of litter has probably been mediated by variable decomposition rates and nutrient dynamics.

In our study, the concentration of Ca, Mn, Al and Fe in forest floors were poorly correlated to corresponding concentrations in litterfall. This absence of clear relationship could be attributed to differences between species concerning (i) the loss of these cations from forest floor horizons by leaching and root uptake (Currie et al. 1999), and/or (ii) the relative accumulation of divalent cations and metals as a result of C-loss by mineralization. These cations are strongly bound to humic compounds, as compared to monovalent cations and Mg (Yavitt and Fahey 1984). The evolution of cation concentrations during litter decay should be influenced by the balance between cation- and C-release. If we apply these hypotheses to our data, the higher C-mineralization under oak as compared with other species could partly explain the highest concentrations of Ca, Mg and Mn in oak forest floor. Vesterdal (1998) found also that oak tended to have higher basal respiration rate than the other species. Furthermore, spruce roots preferentially colonize forest floor (Puhe 2003). Root uptake in this forest floor may reduce cation release.

4.2 Impact of tree species on C- and N-transformation

As previously reported, our results support the conclusion that (i) tree species exert a strong control on C- and N-cycling (Gower and Son 1992; Lovett et al. 2004), particularly on nitrification (Zeller et al. 2007), and (ii) tree species can modify N-cycling rates in a short time (decades) relative to soil development (Gower and Son 1992). In addition, our study shows that the profile pattern of C- and N-transformation was not impacted by the tree species, and that only the “intensity” of C- and N-transformation was impacted.

Our results are consistent with previous studies on the same experimental site. The effect of tree species on C-mineralization in forest floors was identical to their effect on C-mineralization in Ah horizon measured by Moukoui et al. (2006) (spruce>beech>Douglas-fir; oak was not considered in their study). As often reported in the literature, beneath the spruce stand, a very low value of nitrification has been observed both in Ah horizon (Zeller et al. 2007) and in forest floor. It might be attributed to the relatively large C:N in litterfall. Nitrification might be affected by roots, either through the competitive absorption of NH_4^+ (Hart et al. 1994) or through the excretion of molecules affecting microorganisms (Turpault et al. 2007). Living roots were not included in our experiments, but the composition of microbial populations may be impacted by contact with roots before sampling.

In our study, the large rates of C-mineralization in oak forest floor, relatively to other tree species, can not be explained by litterfall composition. No correlation was observed between C-mineralization and litter quality indexes such as C:N, lignin:N and lignin defined by Melillo and Aber (1982) or such as Mn concentration defined by Berg et al. (2007). This could be primarily attributed to discrepancy in litter decomposition. Such discrepancy could come from absence of allelopathic compounds (Kraus et al. 2003; Lorenz et al. 2004). Kuiters and Denneman (1987) reported that polyphenol concentrations were higher in the forest floor and soil under Norway spruce and beech

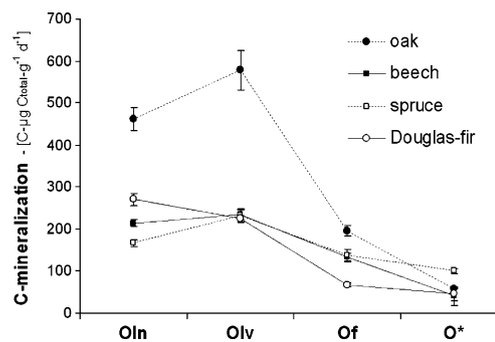


Fig. 4 C-mineralization in forest floor horizons of each species

Table 4 C-mineralization, net N-nitrification and relative nitrification in forest floor horizons and in the whole forest floor. The relative nitrification (%) is defined by the ratio between (1) final NO_3^- in the KCl and water extracts and (2) final mineral N (NH_4^+ and NO_3^-) in the KCl and water extracts. The values for the whole forest floor were

calculated using the dry weight of each horizon/sub-horizon as weighting factor (weighted average values — *WAV*). For each horizon, means with different associated letters are significantly different from one another at the $P < 0.05$ level according to Fisher's least significant difference test ($n=3$)

Horizon	Species	C-mineralization			N-nitrification			Relative nitrification		
		[C- $\mu\text{g C}_{\text{total}}\text{-g}^{-1} \text{d}^{-1}$]			[N- $\mu\text{g N}_{\text{total}}\text{-g}^{-1} \text{d}^{-1}$]			%		
		Mean	SD		Mean	SD		Mean	SD	
Oln	oak	462	28	a	15	26	a	5	9	a
	beech	214	8	c	32	28	a	10	9	a
	Spruce	167	8	d	1	1	a	5	7	a
	Douglas-fir	270	15	b	37	11	a	7	2	a
				$P < 0.0001$			$P 0.1825$			$P 0.8006$
Olv	oak	578	47	a	60	5	a	7	3	ab
	beech	234	12	b	35	6	b	5	1	ab
	Spruce	232	14	b	9	14	c	4	6	b
	Douglas-fir	224	7	b	69	8	a	12	3	a
				$P < 0.0001$			$P 0.0002$			$P 0.1386$
Of	oak	196	13	a	185	31	a	13	2	b
	beech	133	10	b	34	10	c	8	1	c
	Spruce	138	13	b	19	3	c	2	0	d
	Douglas-fir	67	4	c	110	26	b	36	2	a
				$P < 0.0001$			$P < 0.0001$			$P < 0.0001$
O*	oak	57	0	b	122	24	a	30	3	b
	beech	40	21	b	73	21	ab	25	1	c
	Spruce	100	5	a	34	3	b	7	0	d
	Douglas-fir	45	15	b	96	44	a	50	3	a
				$P 0.0021$			$P 0.0232$			$P < 0.0001$
Forest floor (<i>WAV</i>)										
	oak	196	9	a	138	22	a	19	2	b
	beech	130	10	c	42	4	c	11	1	c
	Spruce	147	3	b	19	3	d	4	1	d
	Douglas-fir	125	6	c	86	20	b	30	1	a
				$P < 0.0001$			$P < 0.0001$			$P < 0.0001$

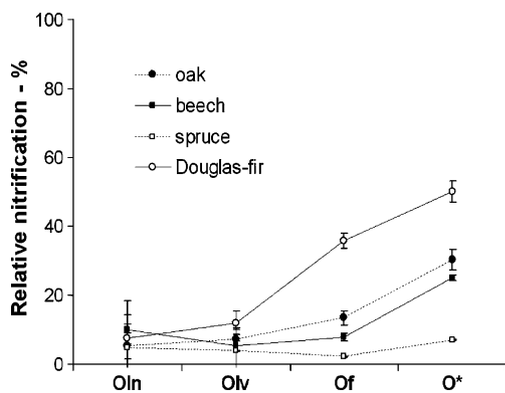


Fig. 5 Relative nitrification in forest floor horizons of each species

than under oak. It could also come from difference in micro- and meso-organism populations (Hobbie et al. 2006; Templer et al. 2003) and from different microbial access to organic N and P sources, rather than by different total amounts of N and P (Vesterdal 1998).

The contribution of N forms (DON, NH_4^+ and NO_3^-) in leachates changes according to the forest floor horizons. Our results confirmed the suggestion of Qualls et al. (1991): DON leached from the forest floor originates essentially from the upper forest floor where soluble organic matters are present. In lower horizons, leachates (Of and O*), DIN (= NH_4^+ and NO_3^-) dominated, corresponding to larger net ammonification and net nitrification.

4.3 Impact of tree species on C- turnover

C-stocks in forest floor were smallest under broadleaved species (oak: $9.1 \text{ Mg ha}^{-1} \pm 3.2$; beech: $7.8 \text{ Mg ha}^{-1} \pm 2.6$; spruce: $21.5 \text{ Mg ha}^{-1} \pm 11.6$; Douglas-fir: $16.4 \text{ Mg ha}^{-1} \pm 4.9$, data from Ranger et al. 2004), although C-input from litterfall was similar whatever the tree species, which probably reveals a faster C-turnover in oak and beech forest floors as compared with coniferous species. This faster C-turnover might be related to (i) the higher C-mineralization in oak forest floor and (ii) the higher DOC releases from both oak and beech forest floors. Our results showed that DOC release must be considered in C-turnover calculation, because it accounted for 22% (d), 40% (b), 51% (c) and 56% (d) ($P < 0.0001$) of the total C-output ($\text{CO}_2 + \text{DOC}$), under Douglas-fir, spruce, beech and oak respectively. Such large DOC contributions to C-output were also observed in the studies of Qualls et al. (1991) and Yavitt and Fahey (1984), which reported that about 30% of the annual C-release ($\text{CO}_2 + \text{DOC}$) from a lodgepole pine forest floor was DOC. Fröberg et al. (2007) even observed that DOC loss accounted for 42% of the total mass loss of a litter of Norway spruce.

4.4 Impact of tree species on the release of aqueous proton donors

The anionic signature widely differed between the four tree species. The strongest contrast was observed between spruce and Douglas-fir, with dominance of NO_3^- in Douglas-fir leachates, and organic anions in spruce leachates. According to Cronan (1978), the anionic charge deficit (ACD) may be largely attributed to deprotonated organic acids. The higher NO_3^- contribution in Douglas-fir leachates can be attributed to the larger net nitrification and low DOC release in this forest floor as compared with other species. A high value of relative nitrification in Douglas-fir forest floor may explain the low pH values in corresponding leachates. Indeed, relative nitrification might be considered as a balance between nitrification and ammonification, which directly affects proton budget in soils, through either proton release or proton consumption (Van Breemen et al. 1983).

The composition of forest floor leachates has direct impact on soil development. Nitric and organic acids induce mineral weathering (Ugolini and Sletten 1991), whereas corresponding anions may contribute to the leaching of cations and metals from topsoil, leading to nutrient depletion, soil acidification and podzolization (Aber 1992; Currie et al. 1999; Van Breemen et al. 1983). Low pH under Douglas-fir and oak may promote mineral weathering, while large DOC concentration under spruce may promote acido-complexolysis (Augusto et al. 1998).

5 Conclusion

We conclude that tree species induce a specific signature of proton donors at the forest floor level (organic vs nitric acid) because of their impact on C- and N-cycling. The release of proton donors can directly impact pedogenetic process in mineral horizons beneath. Consequently, soil degradation could be influenced by setting up of well-chosen tree species stands. However, as root uptake of cations and anions is not considered in our experiment, we cannot draw further conclusions on the effect of root uptake of nutrients on the control of nitrification. We believe that interesting perspectives could involve in situ studies taking into account key processes impacted by tree species: nutrient uptake, root exudation, element leaching from canopy.

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