No depth-dependence of fine root litter decomposition in temperate beech forest soils

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Abstract

Aims Subsoil organic carbon (OC) tends to be older and is presumed to be more stable than topsoil OC, but the reasons for this are not yet resolved. One hypothesis is that decomposition rates decrease with increasing soil depth. We tested whether decomposition rates of beech fine root litter varied with depth for a range of soils using a litterbag experiment in German beech forest plots.

Methods In three study regions (Schorfheide-Chorin, Hainich-Dün and Schwäbische-Alb), we buried 432 litterbags containing 0.5 g of standardized beech root material (fine roots with a similar chemical composition collected from 2 year old Fagus sylvatica L. saplings, root diameter <2 mm) at three different soil depths (5, 20 and 35 cm). The decomposition rates as well as the changes in the carbon (C) and nitrogen (N) concentrations of the decomposing fine root litter were determined at a 6 months interval during a 2 years field experiment.

Results The amount of root litter remaining after 2 years of field incubation differed between the study regions (76±2 % in Schorfheide-Chorin, 85±2 % in Schwäbische-Alb, and 88±2 % in Hainich-Dün) but did not vary with soil depth.

Conclusions Our results indicate that the initial fine root decomposition rates are more influenced by regional scale differences in environmental conditions including climate and soil parent material, than by changes in microbial activities with soil depth. Moreover, they suggest that a similar potential to decompose new resources in the form of root litter exists in both surface and deep soils.

Keywords Fagus sylvatica L · Root litter · Decomposition · Soil depth · Beech forests

Introduction

Forests cover approximately 30 % of the global land surface and forest soils contain up to 40 % of the total belowground terrestrial C, including soil organic matter, litter and living roots (Brunner and Godbold 2007; Dixon et al. 1994). Roots are a major source supplying C to soil organic matter, the largest reservoir of the terrestrial C cycle (Mendez-Millan et al. 2010; Rasse et al. 2005; Schlesinger 1997). The organic compounds contained in dead roots are transformed by decomposition and are partly converted to stable forms of soil organic matter or are being mineralized (Trumbore 2009). Root decomposition controls on the dynamics
of soil OC storage involve complex interactions between plants and the soil biota as well as their reactions to changes in local environmental conditions such as climate, soil parent material, pH and nutrient availability (Chen et al. 2000; Handa et al. 2014; Solly et al. 2014, Thoms et al. 2010). Although edaphic conditions vary with depth in a soil profile and more than half of total soil OC is found in subsoils (Jobbágy and Jackson 2000; Rumpel and Kögel-Knabner 2011), the majority of the studies on root litter decomposition and C dynamics have focused on topsoils (Heim and Frey 2004; Hobbie et al. 2010; Sun et al. 2013; van Huysen et al. 2013). Accordingly, the pattern of root decomposition in the deeper soil layers and its controlling factors remain poorly understood (Gill and Burke 2002; Rumpel and Kögel-Knabner 2011). Moreover, the lack of large scale studies on root decomposition at different soil depths inhibits our ability to determine how soil OC storage will respond to short and long term environmental changes in different regions. Even small variations of decomposition rates of plant litter in forest ecosystems could influence the OC concentrations and storage of the large soil C reservoir (Trumbore 2009).

While OC concentrations are higher in the topsoil, due to higher plant inputs, root densities, microbial biomass and substrate input through litter leaching, the age of soil OC increases with depth in most ecosystems (Paul et al. 1997; Trumbore 2000). One possible explanation for older subsoil OC is a reduction in OC turnover with depth, which is supported by increasing portions of mineral-protected OC, changes in the quantity and quality of plant litter input and lower seasonal variations in soil water content and soil temperature in deeper compared to surface soil, all altering microbial activity and root decomposition rates (Fierer et al. 2003a; Fontaine et al. 2007; Rovira and Ramón Vallejo 2002; Rumpel and Kögel-Knabner 2011). Interestingly, the age of the free light fraction of soil organic matter, representing the C not protected inside aggregates or associated with minerals, does not show a consistent increase with depth (Schöning and Kögel-Knabner 2006; Schrumpf et al. 2013). This suggests that C mineral protection is more important for decreasing subsoil OC turnover than an overall reduced potential for microbial decomposition. However, until now little effort was done to determine whether greater C stability and reduced microbial activity with soil depth influence organic matter decomposition rates (Gill and Burke 2002; Sanaullah et al. 2011).

In temperate forest topsoils, fine root decomposition rates have been observed to vary regionally and with changes in environmental conditions such as soil temperature and moisture (Solly et al. 2014), which in turn influence the relative activities of the decomposer community (Chen et al. 2000). But it is still unclear whether the same pattern will be observed in deeper soil layers and whether decomposition shows a similar dependence on depth in different soil types. The vast majority of decomposition studies have until now been conducted at small scales and using on-site specific litter, complicating site and regional inter-comparison (Parton et al. 2007). Hence, here we used a standardized fine root litter, with similar chemical quality, to determine controls of fine root decomposition in 12 beech (Fagus sylvatica L.) forest plots.

The main objective of this study is to assess the depth-dependence of fine root litter decomposition for a range of forest soils distributed over three German study regions that differ in climate and soil parent material. We used the litterbag method to estimate decomposition rates of standardized beech fine root litter. Our hypotheses are that i) the rate of root decomposition decreases with increasing soil depth, reflecting changes in soil abiotic and biotic conditions; and that ii) the decomposition rates differ between the three study regions for all studied depths, mainly due to diverse and site-specific climate and soil biota, with faster decomposition in warmer and moister not waterlogged sites.

Materials and methods

Study sites

We conducted this study in 12 forest plots dominated by European beech (Fagus sylvatica L.) and distributed in three German regions of the ‘Biodiversity Exploratories’ (Fischer et al. 2010). The Schwäbische Alb is situated in south-western Germany, the Hainich-Dün in central Germany and the Schorfheide-Chorin in northeastern Germany. These three study regions differ in climate and soil parent material (Table 1; for details see Fischer et al. (2010)). For each region we selected four forest plots (100 m × 100 m) with similar age class management. All forests were harvested at 80–120 year intervals and were in an old timber development stage.
Soil properties

To evaluate the soil OC and total N concentrations and stocks, we collected one mineral soil core in each of the forest plots (For more details see: Grüneberg et al. 2010). We used a soil corer with an inner diameter of 8.3 and a length of 110 cm (Eijkelkamp Agrisearch Equipment BV, Giesbeek, The Netherlands). The core was driven into the soil with a motor hammer (Atlas Copco AB, Nacka, Sweden). Organic layers were removed before coring. We sectioned the soil cores into increments (0–10, 10–30 and 30–50 cm) for analysis. Prior to processing, the soil samples were air dried at 20 °C to constant weight and sieved to <2 mm. Roots were air dried and their weight was determined. Sieved soil subsamples were ground in a ball mill (RETSCH MM 300, Retsch, Haan, Germany). Total C and N concentrations were determined using the elemental analyzer (Vario Max, Hanau, Germany). Concentrations of calcium (Ca), magnesium (Mg), aluminium (Al) and phosphorous (P) were measured using inductively coupled plasma - optical emission spectrometry (ICP-OES, Optima 3300 DV, Perkin Elmer, Norwalk, USA) (for details on the method see: Raessler et al. 2005). Lignin and cellulose content were estimated from thermogravimetric analysis in Argon atmosphere (TGA / SDTA851e, Mettler Toledo, GmbH, Giessen, Germany) (Yang et al. 2005, method described in Solly et al. 2014).

Root decomposition

We assessed the decomposition rates of beech fine root litter buried at three depths. To provide a nearly homogeneous substrate for decomposition, we used fine roots collected from 2 years old beech saplings grown in sand. While there may be some chemical differences between deep and shallow fine roots, as well as between roots of different diameter size and age, our aim was to focus on

Table 1 Main geographical and environmental characteristics of the three study regions: Schwäbische Alb, Hainich-Dün, Schorfheide-Chorin (adapted from Fischer et al., 2010)

|                      | Schwäbische Alb | Hainich-Dün | Schorfheide-Chorin |
|----------------------|-----------------|-------------|--------------------|
| Location             | SW Germany      | Central Germany | NE Germany        |
| Coordinates          | N 48° 26′ E 9° 23′ | N 51° 9′ E 10° 28′ | N 53° 0′ E 13° 46′ |
| Area of the region [km²] | ~422          | ~1300       | ~1300              |
| Soil type            | Cambisol (eutric) | Luvisol    | Cambisol (dystric) |
| Altitude a.s.l [m]   | 460–860         | 285–550     | 3–140              |
| Mean annual temperature [°C] | 6.0–7.0    | 6.5–8.0     | 8.0–8.5            |
| Mean annual precipitation [mm] | 700–1000   | 500–800     | 500–600            |
| Parent material      | Jurassic limestone and loess | Triassic shell limestone | Glacial till and aeolian / fluvial sand in depressions |
the effect of soil depth on fine root decomposition rather than on the differences triggered by plant allocation patterns.

After removing the mineral soil particles attached to the fine roots of the saplings, by carefully cleaning the roots with distilled water in a 63 μm sieve, we separated the fine roots (<2 mm diameter) from the coarse roots and dried all samples at 40 °C to constant weight in a forced-air oven. For details on the method see Solly et al. (2014).

We prepared 432 litterbags by placing 0.5±0.01 g of dry fine root litter into a 10 cm×10 cm litterbag made of a 100 μm polyester mesh screening to allow microfaunal decomposition (Schwegmann Filtrations-Technik GmbH, Grafschaft-Gelsdorf, Germany). We individually labelled each litterbag with a stainless steel label that was placed inside the enclosure and measured the total mass of each litterbag. We buried 36 litterbags (n=12 at three depths) in each of the 12 forest plots in October 2011. After extracting 12 intact soil cores (48 mm diameter, 40 cm depth) from each forest plot, we placed the litterbags vertically, adhering to the lateral surface of the soil core hole, at average soil depths of 5, 20 and 35 cm. To backfill the space between the litterbags we used the soil extracted from the core, which we cut at the specific depths. In April 2012, October 2012, April 2013 and October 2013 we collected three litterbags from each soil depth in every forest plot (n=9 per collection date in each site). The collected litterbags were transported to the laboratory where we gently removed the fine roots and hyphae grown around the litterbags as well as soil particles adhering to the fine root-litter. After drying at 40 °C we calculated the fine root decomposition rates (mass recovery in %) for each forest plot as the average amount of mass recovered in the three litterbags collected at each collection date. We further estimated the fine root litter decomposition rates (k-values) for every soil depth in all forest plots by fitting the exponential function

\[ X_t = X_0 e^{-kt} \]

Xt is the amount of fine root remaining at time t and X0 is the initial root mass at time 0. After 2 years of decomposition, three replicates per soil depth were analyzed for total C and N concentrations.

Statistics

We conducted statistical analyses with R, version 3.0.2 (R Development Core Team 2013). Throughout the manuscript we present data as means±standard deviation. Analysis of variance (ANOVA) accompanied by Tukey-Kramer HSD test was used to examine statistical differences of the fine root mass remaining, the root biomass, the OC and total N concentrations and stocks in the different study regions and at the different soil depths. ANOVA analysis was also used to detect significant differences between the C:N ratio, C and N concentrations of the decomposing fine root litter at different soil depths. Significant changes in the C:N ratio, C and N concentrations between the initial standardized fine root litter and the root litter after 2 years of decomposition (averaged at all soil depths) were tested with Student’s t test. Normal distribution of residuals and homogenous variances were checked before statistical analysis. The chosen p-value for detecting statistical differences was p<0.05.

Results

Soil properties and root biomass

OC concentrations ranged between 12 and 42 g kg⁻¹ in the soil sampled at 0–10 cm depth. These were significantly higher than the concentrations found at 30–50 cm belowground, which ranged between 2 and 11 g kg⁻¹. Total N concentrations also declined significantly with depth (Table 2). OC and total N concentrations were highest in the Schwäbische-Alb, followed by Hainich-Dün and Schorfheide-Chorin. The amount of root biomass (dead and live) extracted from soils as well as OC and total N stocks decreased with depth in all study regions (Table 2). Variability in the chemical quality of roots (C, N, Ca, Mg, Al, P, lignin and cellulose) found in the beech fine root biomass that we used for our decomposition experiment was minor (Table 2). The variability of soil moisture and soil temperature declined with depth for all study regions (Table 3). The soils of the Schwäbische-Alb study region had a similar volumetric soil water content (ranging between 20 and 38 %) to the ones in the Hainich-Dün (ranging between 22 and 31 %) and a higher volumetric soil water content than the soils in the Schorfheide-Chorin (ranging between 9 and 20 %). The topsoil in the Hainich-Dün was frozen for...
the longest period (8 days per year) in comparison to the Schwäbische-Alb (6 days) and the Schorfheide-Chorin (4 days).

Root mass recovery, decomposition rates and C:N ratio

After 2 years of decomposition on average for all depths 76±2% in Schorfheide-Chorin, 85±2% in Hainich-Dün and 88±2% in Schwäbische-Alb of initial mass of beech fine root litter remained. The mass remaining did not differ significantly for the three depths sampled in the three regions for the majority of the collection times (Fig.1), and thus our calculated decomposition rate-constants also did not vary with depth (Table 4). Only in the Hainich-Dün the mass remaining differed significantly between depths after 12 and 18 months of decomposition, and in the Schwäbische-Alb the amount of mass remaining differed among the 3 depths after 6 months of decomposition but not for the subsequent decomposition times (Fig.1). Regional differences in the amount of fine root litter remaining were consistent over the three soil depths (Figs.1, 2). Decomposition significantly increased the C:N ratio of the fine root litter from 46 to 54 on average after 2 years (Fig.3). The increase was similar across different soil depths (Fig.3), and primarily driven by N loss as no major changes in the C concentrations were detected. After 2 years the C and N concentrations as well as the C:N ratio of the decomposing fine roots were nearly constant throughout the soil profile, however the C concentrations of the fine roots decomposing at 20 and 35 cm soil depths were the soil moisture and temperature over 2 years of decomposition, in all study regions at different soil depths (mean±SD). Soil temperature is expressed in °C and soil moisture in percentage of volumetric water content (%VWC).

| Study region     | Soil depth | Soil temperature | Soil moisture |
|------------------|------------|------------------|--------------|
| Schwäbische-Alb | 10         | 6.3±4.1          | 31±9         |
|                 | 20         | 6.3±4.1          | 31±9         |
|                 | 30         | 6.3±4.1          | 31±9         |
| Hainich-Dün     | 10         | 6.3±4.1          | 31±9         |
|                 | 20         | 6.3±4.1          | 31±9         |
|                 | 30         | 6.3±4.1          | 31±9         |
| Schorfheide-Chor | 10         | 6.3±4.1          | 31±9         |
|                 | 20         | 6.3±4.1          | 31±9         |
|                 | 30         | 6.3±4.1          | 31±9         |
higher compared to the roots decomposing at 5 cm ($p<0.05$) (Fig. 3).

**Discussion**

Similar fine root decomposition at different soil depths

Previous research proposed that litter decomposition rates would be highest in the topsoil and decrease in deeper soil horizons because microbial activities and substrate availability as well as relative variations in soil moisture and soil temperature decline with soil depth (e.g., more pronounced maximum temperature in the topsoil may increase decomposition in comparison to subsoil) (Fierer et al. 2003a; Jobbágy and Jackson 2000; Trumbore 2000; Weaver et al. 1935). In the studied forest plots the variability of both soil moisture and soil temperature slightly declined throughout the soil profile (Table 3), together with OC and N concentrations (Table 2). Herold et al. (2014) showed that extracellular enzyme activities significantly declined with soil depth in forest plots of the same study regions (Table 5 in supplementary material). Although these variables declined with soil depth we found that fine root decomposition was nearly uniform through the forest soil horizons, except after 12 and 18 months in the Hainich-Dün and after 6 months in the Schwäbische-Alb (Fig. 1). One reason explaining the similarity of the decomposition rates with soil depth, could be that while we observed that the variation of temperature and moisture declined with soil depth, their means were similar. Accordingly, lower winter temperatures in topsoils might balance lower summer temperatures of forest subsoils, leading to comparable decomposition rates in the topsoil and the subsoil. The same could be true for soil moisture, where greater variation in the topsoil (too dry or too wet) can be detrimental for decomposition (Chen et al. 2000).

The similarity of root decomposition at the three depths agreed with a study by Sanaullah et al. (2011). Sanaullah et al. (2011) showed that the amount of wheat root derived C and N remaining in the soil was similar in top- and subsoil horizons of a temporary grassland managed with a ley cropping system in southern France after 3 years of decomposition. Weaver (1947) also concluded that the rate of root decomposition did not show consistent differences at different soil depths throughout a prairie soil profile in North America. The recently published study by Li et al. (2015) also indi-

**Table 4** Estimates of the decomposition rate-constant ($k$-values) of fine roots at three different soil depths in 12 beech forest plots of the Schwäbische-Alb, Hainich-Dün and Schorfheide-Chorin regions (mean±SD)

| Soil depth cm | Schwäbische-Alb $k$-values [year$^{-1}$] | Hainich-Dün | Schorfheide-Chorin |
|---------------|-----------------------------------------|-------------|--------------------|
| 5             | 0.09±0.02                               | 0.07±0.01   | 0.13±0.01          |
| 20            | 0.07±0.02                               | 0.06±0.01   | 0.13±0.04          |
| 35            | 0.08±0.01                               | 0.05±0.01   | 0.12±0.02          |
cates no changes in the decomposition rates between the organic horizon and the mineral horizon in a pine forest ecosystem, for roots decomposing both in situ and in litterbags. These observations together with our finding of similar C:N ratios of the decomposing fine root litter at different soil depths, suggest that fine root decomposition is not strongly affected by the overall changes in soil abiotic and biotic conditions with depth. We assume that the higher biological activity in topsoil than in subsoil is probably promoted by larger amounts of root litter in topsoil (Table 2), and additional nutrient and organic substrate input via throughfall and from litter leaching (Qualls and Haines 1992). Declining enzyme activities with soil depth are therefore likely to be a consequence of reduced resource availability and thus less microbial activity in deeper soil horizons (Fierer et al. 2003b; Herold et al. 2014), which we also observed in our study with decreased root biomass and nitrogen and OC concentrations with soil depth (Table 2). When inputs are increased, for example by adding a litterbag with new resources, microbial activity can also be stimulated in subsoils to achieve comparable decomposition to that in topsoils. Studies on lignin decomposition indicate that it depends on a continuous input of available energy and C sources, which enable the production of lignin degrading enzymes, and that it is hampered when bioavailable C becomes limited, i.e., during late decomposition stages (Klotzbücher et al.)
2011). Hence, it is possible that in later decomposition phases (after 2 years), the lower substrate inputs and nutrient resources in subsoils may hamper subsoil decomposition rates relative to topsoils. Longer-term decomposition experiments would be necessary to test this hypothesis.

It has been shown that the radiocarbon age of the more active, physicochemically unprotected C in the free light fraction only slightly increases with soil depth, while the less active fractions (C occluded in aggregates or in association with minerals) become older along the vertical soil profile (Schöning and Kögel-Knabner 2006; Schrumpf et al. 2013). Together with our results, this indicates that in both surface and deep soils the free light fraction, presumably derived from the breakdown of fine root litter and aboveground litter in the topsoil, is preferentially decomposed at all soil depths.

Although after 2 years of decomposition the estimated decomposition rates (k-values) for the whole time series were similar for all studied soil depths in all three study regions, in the Schwäbische-Alb study region the mass remaining after 6 months was highest in the topsoil. This may be related to colder temperatures in the topsoil during the winter months. Moreover, in the Hainich-Dün study region the decomposition of fine roots was observed to decline across the soil profile. An explanation for this is that the studied soils in this area (Luvisols) are characterized by a rich clay accumulation in the subsurface horizons, where soil water may stagnate for a longer time during wet periods, or dry out during the summer months. Hence, these conditions may reduce root decomposition across the soil profile. These exceptions indicate that the length of an experiment could lead to different outcomes of the influence of depth gradients on root decomposition rates within one study region.

After 2 years, the mass of the standardized beech fine root litter remaining was slightly higher than averages reported for other temperate tree species (Parton et al. 2007; Sun et al. 2013). These slower decomposition rates may reflect the higher amounts of lignin in Fagus sylvatica L. roots compared to roots of other tree species (Hobbie et al. 2010). More lignified roots have a higher resistance to enzymatic attack and therefore tend to have slower decomposition rates than the less lignified counterparts (Berg 1984). In our experiment we used a litterbag mesh size of 100 μm, which enabled us to account only for microfaunal decomposition and excluded the larger soil decomposer community, which contributes to the mechanical breakdown of plant litter especially in topsoil layers. We cannot exclude that the decomposition absolute values might be different for roots decomposing in situ with different diameter sizes or ages (i.e., fine roots of 2 year-old samplings vs fine roots of adult trees or roots found at different soil depths), as was suggested by earlier studies (Dombush et al. 2002, Goebel et al. 2010, Hishi 2007, Li et al. 2015). Therefore, the absolute amount of root mass that we recovered at different soil depths might be influenced by the set-up of the experiment. However, as litterbags and fine root litter were standardized, we can still compare the effects of soil properties on decomposition rates at different soil depths and in different regions.

Site effects on decomposition of fine roots

Although we observed no general trend with soil depth, the recovered fine root masses varied over all soil depths between the three regions (Fig. 2), supporting our second hypothesis. The same pattern of higher decomposition rates in the Schorfheide-Chorin followed by Schwäbische-Alb and Hainich-Dün was also found in a larger study aimed at investigating the factors controlling root decomposition in the topsoil of the same regions (Solly et al. 2014). In this larger scale experiment, exploring decomposition of standardized fine root litter in a range of forest types, soil temperature and moisture explained most of the regional variability of root decomposition in temperate forests. Although the Hainich-Dün study region has intermediate soil properties and climate, it has on average the lowest root decomposition rates. One explanation could be that in the Hainich-Dün the soils are on average frozen for a higher number of days in comparison to the other study regions (Solly et al. 2014). Previous studies have already shown that in soils with temperatures below 0 °C microbial activity declines (Dioumaeva et al. 2002; Mikan et al. 2002). Moreover, higher root decomposition rates in the Schorfheide-Chorin may result from a greater vertical water movement through the sand in this study region, which can increase the leaching of the soluble organic matter derived from the decomposing roots. Regional differences in the diversity of the soil biota, adapted to diverse edaphic factors including climate, soil texture and pH may also have played a role in determining the regional variability in fine root decomposition (Birkhofer et al. 2012, Thoms et al. 2010).
Conclusion

Our results show that fine root decomposition in litterbags has a larger variability across regions than at different soil depths within a given site. Thus, a similar potential to decompose root derived C exists in both, surface and deep soils. Biological activity in different soil horizons would then be mainly limited by root litter input and thus the input of new substrates. Overall, rapid decomposition of litter along the soil profile is supported by the consistent young ages of C in unprotected fractions observed for different soil horizons in other studies. The increase in the age of bulk soil C with depth is likely to be more related to the mineral-stabilized C, but further studies are required to test this hypothesis. Macroclimatic or other large-scale differences like parental material seem to be more important for initial stages of litter decomposition than changing conditions with soil depth, as regional differences in decomposition persisted for all studied soil depths.

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Fontaine S, Barot S, Barre P, Bidiou N, Mary B, Rumpel C (2007) Stability of organic carbon in deep soil layers controlled by fresh carbon supply. Nature 450:277–280, http://www.nature.com/nature/journal/v450/n7167/suppinfo/nature06275_S1.html

Gill R, Burke I (2002) Influence of soil depth on the decomposition of bouteoula gracilis roots in the shortgrass steppe. Plant Soil 241:233–242. doi:10.1023/a:101646800542

Goebel M, Hoebbe SE, Bulaj B, Zadworny M, Archibald DD, Oleksyn J, Reich PB, Eisenstat D (2010) Decomposition of the finest root branching orders: linking belowground dynamics to fine-root function and structure. Ecol Monogr 81: 89–102

Grüneberg E, Schönig I, Kalko EKV, Weisser WW (2010) Regional organic carbon stock variability: a comparison between depth increments and soil horizons. Geoderma 155:426–433. doi:10.1016/j.geoderma.2010.01.002

References

Berg B (1984) Decomposition of root litter and some factors regulating the process: long-term root litter decomposition in a Scots pine forest. Soil Biol Biochem 16:609–617. doi:10.1016/0038-0717(84)90081-6

Birkhofer K, Schönig I, Alt F, Herold N, Klarnert B, Marau M, Marhan S, Olennmann Y, Wubet T, Yurkov A, Begerow D, Berner D, Buscot F, Daniel R, Diekötter T, Ehnes RB, Erdmann G, Fischer C, Foesel B, Groh J, Gutknecht I, Kandeler E, Lang C, Lohaus G, Meyer A, Nacket H, Nätter A, Overmann J, Polle A, Pollerier MM, Scheu S, Schloter M, Schulze ED, Schulze W, Weirent J, Weisser WW, Wolfers V, Schrumpf M (2012) General relationships between abiotic soil properties and soil biota across spatial scales and different land-Use types. PLoS ONE 7(8), e43292. doi:10.1371/journal.pone.0043292

Brummer I, Godbold DL (2007) Tree roots in a changing world. J For Res 12:78–82. doi:10.1007/s10310-006-0261-4

Chen H, Harmon ME, Griffiths RP, Hicks W (2000) Effects of temperature and moisture on carbon respired from decomposing woody roots. For Ecol Manag 138:51–64. doi:10.1016/S0378-1127(00)00411-4

Dornbush ME, Isenhart TM, Raich JW (2002) Quantifying fine-root decomposition an alternative to buried litterbags. Ecology 83:2985–2990

Dioumaeva I, Trumbore S, Schuur EAG, Goulden ML, Litvak V, Hirsch AI (2002) Decomposition of peat from upland boreal forest: temperature dependence and sources of respired carbon. J Geophys Res: Atmos 107:8222. doi:10.1029/2001jd000848

Dixon RK, Solomon A, Brown S, Houghton R, Trexier M, Wisniewski J (1994) Carbon pools and flux of global forest ecosystems. Science 263:185–190

Fierer N, Allen AS, Schimel JP, Holden PA (2003a) Controls on microbial CO2 production: a comparison of surface and subsurface soil horizons. Glob Chang Biol 9:1322–1332

Fierer N, Schimel JP, Holden PA (2003b) Variations in microbial community composition through two soil depth profiles. Soil Biol Biochem 35:167–176. doi:10.1016/S0038-0717(02)00251-1

Fischer M, Bossdorf O, Gockel S, Hänsel F, Hemp A, Hessenmüller D, Korte G, Nieschulze J, Pfeiffer S, Prati D, Renner S, Schönig I, Schumacher U, Wells K, Buscot F, Kalko EKV, Linsenmair KE, Schulze ED, Weisser WW (2010) Implementing large-scale and long-term functional biodiversity research: the biodiversity exploratories. Basic Appl Ecol 11:473–485. doi:10.1016/j.baae.2010.07.009

Fontaine S, Barot S, Barre P, Bidiou N, Mary B, Rumpel C (2007) Stability of organic carbon in deep soil layers controlled by fresh carbon supply. Nature 450:277–280, http://www.nature.com/nature/journal/v450/n7167_suppinfo/nature06275_s1.html

Gill R, Burke I (2002) Influence of soil depth on the decomposition of bouteoula gracilis roots in the shortgrass steppe. Plant Soil 241:233–242. doi:10.1023/a:101646800542

Goebel M, Hoebbe SE, Bulaj B, Zadworny M, Archibald DD, Oleksyn J, Reich PB, Eisenstat D (2010) Decomposition of the finest root branching orders: linking belowground dynamics to fine-root function and structure. Ecol Monogr 81: 89–102

Grüneberg E, Schönig I, Kalko EKV, Weisser WW (2010) Regional organic carbon stock variability: a comparison between depth increments and soil horizons. Geoderma 155:426–433. doi:10.1016/j.geoderma.2010.01.002
Handa IT, Aerts R, Berendse F, Berg MP, Bruder A, Butschensoen O, Chauvet E, Gessner MO, Jabiol J, Makkonen M, McKie BG, Malmqvist B, Peeters ETHM, Scheu S, Schmid B, van Ruijven J, Vos VCA, Hattenschwiler S (2014) Consequences of biodiversity loss for litter decomposition across biomes. Nature 509:218–221. doi:10.1038/nature13247

Heim A, Frey B (2004) Early stage litter decomposition rates for Swiss forests. Biogeochemistry 70:299–313. doi:10.1007/s10533-003-0844-5

Heiri O, Lotter AF, Lemcke G (2001) Loss on ignition as a method for estimating organic and carbonate content in sediments: reproducibility and comparability of results. J Paleolimnol 25:101–110

Herold N, Schöning I, Berner D, Haslwimmer H, Kandeler E, Herbarth O, Lotter AF, Lemcke G (2001) Loss on ignition as a method for estimating organic and carbonate content in sediments: reproducibility and comparability of results. J Paleolimnol 25:101–110

Hishi T (2007) Heterogeneity of individual roots within the fine root architecture: casual links between physiological and ecosystem functions. J For Res 12:126–133

Hobbie S, Oleksyn J, Eissenstat D, Reich P (2010) Fine root decomposition rates do not mirror those of leaf litter among temperate tree species. Oecologia 162:505–513. doi:10.1007/s00442-009-1479-6

Jobbágy EG, Jackson RB (2000) The vertical distribution of soil organic carbon and its relation to climate and vegetation. Ecol Appl 10:423–436. doi:10.1890/1051-0761(2000)010[0423:TFODTS]2.0.CO;2

Klotzbücher T, Kaiser K, Guggenberger G, Gatzek C, Kalbitz K (2011) A new conceptual model for the fate of lignin in decomposing plant litter. Ecology 92:1052–1062. doi:10.1890/10307.1

Li A, Fahey TJ, Pawlowska TE, Fisk MC, Burtis J (2015) Fine root decomposition, nutrient mobilization and fungal communities in a pine forest ecosystem. Soil Biol Biochem 83:76–83

Mendez-Millan M, Dignac MF, Rumpel C, Rasse DP, Derenne S (2011) Molecular dynamics of shoot vs. root biomarkers in an agricultural soil estimated by natural abundance 13C labelling. Soil Biol Biochem 42:169–177. doi:10.1016/j.soilbio.2009.10.010

Mikan CJ, Schimel JP, Doyle AP (2002) Temperature controls of microbial respiration in arctic tundra soils above and below freezing. Soil Biol Biochem 34:1785–1795. doi:10.1016/S0038-0717(02)00168-2

Olson JS (1963) Energy storage and the balance of producers and decomposers in ecological systems. Ecology 44:322–331

Parton W, Silver WL, Burke IC, Grassens L, Harmon ME, Currie WS, King JY, Adair EC, Brandt LA, Hart SC, Fasth B (2007) Global-scale similarities in nitrogen release patterns during long-term decomposition. Science 315:364–366. doi:10.1126/science.1134853

Paul EA, Follett RF, Leavitt SW, Halvorsen A, Peterson GA, Lyon DJ (1997) Radiocarbon dating for determination of soil organic matter pool sizes and dynamics. Soil Sci Soc Am J 61:1058–1067. doi:10.2136/ssaj1997.0361599506010040011x

Qualls RG, Haines BL (1992) Biodegradability of dissolved organic matter in forest throughfall, soil solution, and stream water. Soil Sci Soc Am J 56:578–586

Raessler M, Rothe J, Hilke I (2005) Accurate determination of Cd, Cr, Cu and Ni in woodlice and their skins—is moulding a means of detoxification? Sci Total Environ 337:83–90. doi:10.1016/j.scitotenv.2004.07.008

Rasse DP, Rumpel C, Dignac MF (2005) Is soil carbon mostly root carbon? Mechanisms for a specific stabilisation. Plant Soil 269:341–356. doi:10.1007/s11104-004-0907-y

Rovira P, Ramón Vallejo V (2002) Mineralization of carbon and nitrogen from plant debris, as affected by debris size and depth of burial. Soil Biol Biochem 34:327–339. doi:10.1016/S0038-0717(01)00186-9

Rumpel C, Kögel-Knabner I (2011) Deep soil organic matter—a key but poorly understood component of terrestrial C cycle. Plant Soil 338:143–158. doi:10.1007/s11104-010-0391-5

Sanaullah M, Chabbi A, Leifeld J, Bardoux G, Billou D, Rumpel C (2011) Decomposition and stabilization of root litter in top- and subsoil horizons: what is the difference? Plant Soil 338:127–141. doi:10.1007/s11104-010-0554-4

Schlesinger WH (1997) Biogeochemistry: an analysis of global change. Academic Press, San Diego

Schöning I, Kögel-Knabner I (2006) Chemical composition of young and old carbon pools throughout Cambisol and Luvisol profiles under forests. Soil Biol Biochem 38:2411–2424. doi:10.1016/j.soilbio.2006.03.005

Schrumpf M, Kaiser K, Guggenberger G, Persson T, Kögel-Knabner I, Schulze ED (2013) Storage and stability of organic carbon in soils as related to depth, occlusion within aggregates, and attachment to minerals. Biogeosciences 10:1675–1691. doi:10.5194/bg-10-1675-2013

Solly EF, Schöning I, Boch S, Kandel C, Mathan S, Michelzik B, Müller J, Zscheischler J, Trumbore SE, Schrumpf M (2014) Factors controlling decomposition rates of fine root litter in temperate forests and grasslands. Plant Soil 382:203–218. doi:10.1007/s11104-014-2151-4

Sun T, Mao Z, Dong L, Hou L, Song Y, Wang X (2013) Further evidence for slow decomposition of very fine roots using two methods: litterbags and intact cores. Plant Soil 366:633–646. doi:10.1007/s11104-012-1457-3

Thoms C, Gattinger A, Jacob M, Thomas FM, Gleixner G (2010) Direct and Indirect effects of tree diversity drive soil microbial diversity in temperate deciduous forest. Soil Biol Biochem 42:1558–1565

Trumbore S (2000) Age of soil organic matter and soil respiration: radiocarbon constraints on belowground C dynamics. Ecol Appl 10:399–411

Trumbore S (2009) Radiocarbon and soil carbon dynamics. Annu Rev Earth Planet Sci 37:47–66

van Huysen TL, Harmon ME, Perakis SS, Chen H (2013) Decomposition and nitrogen dynamics of 15 N-labeled leaf, root, and twig litter in temperate coniferous forests. Oecologia 173:1563–1573

Weaver JE (1947) Rate of decomposition of roots and rhizomes of certain range grasses in undisturbed prairie soil. Ecology 28:221–240

Weaver JE, Hougen V, Weldon M (1935) Relation of root distribution to organic matter in prairie soil. Bot Gaz 96:389–420

Yang H, Yan R, Chen H, Zheng C, Lee DH, Liang DT (2005) Influence of microbial biomass pyrolysis on C content and functional group composition of soil organic matter. Energy Fuel 20:388–393. doi:10.1021/ef0580117