Biological nitrogen fixation and nifH gene abundance in deadwood of 13 different tree species

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Abstract  Deadwood is an extremely nitrogen (N) poor plant litter whose decomposition may rely on external N sources. Biological N₂ fixation (BNF) by free-living diazotrophs, encoded with the nifH gene, is a potential pathway of N acquisition in deadwood. Still, the control of this process by tree species specific traits is hardly known. Here, we examined (1) BNF rates and nifH gene abundances in deadwood of 13 tree species after 12 years of decomposition and (2) how BNF was related to nutrient concentrations and non-structural carbohydrates (NSC). Comparing our BNF rates with the literature revealed no difference for angiosperms but gymnosperms. Large variability prevented any significant difference in BNF and nifH gene abundance between tree species identities of deadwood. Still, higher BNF and nifH gene abundances occurred in the group of diffuse-porous angiosperms compared to ring-porous angiosperms and gymnosperms. Positive relationships between BNF rates and molybdenum, phosphor, sulfur, and N suggested co-limitation of BNF by several nutrients in deadwood. Relatively high NSC concentrations indicated good carbon availability for diazotrophs. Furthermore, structural equation modeling highlighted the importance of water content for BNF and nifH gene abundance, although the overall explanatory power was low. In conclusion, BNF is a ubiquitous microbial process in deadwood of native European tree species and the comparison with other studies suggests no down-regulation of BNF by high N deposition in Central Europe.

Keywords  Biological nitrogen fixation · Deadwood · Nutrients · Tree species · Non-structural carbohydrates · ¹⁵N abundance

Introduction

Deadwood is an essential ecological component in forest ecosystems, whose function as habitat and resource for xylobiontic organisms is related to its nutrient availability (Purahong et al. 2016). Among other nutrients, nitrogen (N) availability is particularly
low (Rinne et al. 2017). It constrains the growth and expansion of fungal hyphae (Hodge et al. 2000) as N is, for instance, crucial for the synthesis of the fungal cell-wall constituent chitin. Low N availability can slow down the decay process of deadwood, especially during the initial phase (Noh et al. 2017). A gradual increase in N concentration with proceeding decay improves the growth conditions for xylobiontic organisms (Rinne et al. 2017; Pastorelli et al. 2020).

A part of the increase in N concentration results from the loss of carbon (C) and the simultaneous retention of N in deadwood during the decay process. Another part of N in decaying deadwood may originate from different sources. Besides the input of reactive N via atmospheric deposition as ammonia (NH₃) and nitrate (Lladó et al. 2017; Romashkin et al. 2018), and the translocation of N from underlying soil into deadwood via fungal hyphae (Lindahl and Finlay 2006; Palviainen et al. 2010), asymbiotic biological N₂ fixation (BNF) can contribute to N enrichment in deadwood (Boring et al. 1988). BNF in deadwood provided up to 14% of the N input (Hicks 2000). However, BNF rates can enormously vary between study sites, wood compartments, and tree species (Hendrickson 1988; Cushon and Feller 1989; Griffiths et al. 1993). Most BNF measurements originate from studies in western North America or boreal regions in Europe (Mäkipää et al. 2018a). Compared to North America or Northern Europe, atmospheric N deposition is higher in Central European forests (Borken and Matzner 2004; Bobbink et al. 2010). Therefore, the function of diazotrophs is potentially not mandatory for satisfying N demands of wood-decaying organisms. High availability of reactive N can even suppress BNF (Reed et al. 2011), though this relationship has not been reported for deadwood yet.

The process of asymbiotic BNF in deadwood depends on several factors. These include climatic (Roskoski 1980; Mäkipää et al. 2018b) and site factors, such as exposition (Gómez-Brandón et al. 2020) and seasonal differences (Roskoski 1980; Hicks 2000), but also the acidity of deadwood (Tláskal et al. 2021), and nutrient availability. Molybdenum (Mo) is a vital nutrient for BNF because of its function as a metal cofactor of the diazotrophic key enzyme nitrogenase. The nitrogenase catalyzes the reduction of N₂ to NH₃ and is responsible for BNF (Hardy et al. 1971). BNF rates vary with the decay stages in deadwood, but the variation is not consistent among studies. The highest BNF rates can occur in the early (Tláskal et al. 2021), intermediate (Wei and Kimmins 1998; Hicks et al. 2003; Rinne et al. 2017), or advanced decay stages (Spano et al. 1982). Those differences can be attributed to changes in the microbial community structure (Hicks et al. 2003), the need for N, particularly in early decay stages (Kuramae et al. 2019), or decreasing wood density (Mäkipää et al. 2018b). As nitrogenase is a highly energy-demanding process, C availability is another limiting factor for the activity of diazotrophs (Khan et al. 2021). Jurgensen et al. (1984) showed increasing BNF rates with decreasing total carbohydrates and water-soluble sugars within the decay process in logs of *Pseudotsuga menziesii*. However, a study relating BNF rates in deadwood and various non-structural carbohydrates (NSC), such as glucose, xylose, and arabinose, is not known to us. Hence, the availability of an energy source to diazotrophs can be considered limited by NSC (Khan et al. 2021), as NSC concentrations changed with decomposition: arabinose, glucose, xylose, and galactose peaked in the second decay stage at one site but remained the same at another. In contrast, fructose and mannose did not change during decay (Di Lella et al. 2019).

Deadwood of different tree species differs by lignin, cellulose, and hemicellulose composition (Rowell et al. 2005; Zhang et al. 2014), nutrient concentrations (Bonanomi et al. 2021), or decomposability (Kahl et al. 2017). However, a tree species effect on BNF is not consistent among studies, ranging from no effect (Griffiths et al. 1993; Brunner and Kimmins 2003; Hicks et al. 2003) to a significant impact on BNF in deadwood (Hendrickson 1988; Mäkipää et al. 2018a; Törmänen and Smolander 2022). Due to the missing consistent tree species effect, Hicks et al. (2003) suggested that differences are more likely to occur between angiosperms and gymnosperms, as the factors controlling the BNF mentioned above seem to differ more between angiosperms and gymnosperms rather than between individual tree species of the same group. To date, research about BNF in logs has focused mainly on gymnosperms, such as *Tsuga heterophylla, Picea sitchensis, Pseudotsuga menziesii*, and *Pinus ponderosa* (Griffiths et al. 1993; Brunner and Kimmins 2003; Chen and Hicks 2003; Hicks et al. 2003) whereas few studies are available for angiosperms (Hendrickson 1988; Mäkipää et al. 2018a). A comprehensive study...
examining BNF in deadwood of many tree species was not yet performed to the best of our knowledge.

The BNF activity is encoded in the nifH gene within the enzyme complex nitrogenase (Hsu and Buckley 2009). The presence and expression of these genes highly depend on deadwood tree species, decay stage, and the N concentration in the logs (Hoppe et al. 2014). However, studies relating the nifH gene quantity with the BNF rate in deadwood are rare (Mäkipää et al. 2018b). The existing literature focuses either on one tree species (Gómez-Brandón et al. 2020; Tláskal et al. 2021) or on the diazotrophic abundance without a relation to BNF rates (Hoppe et al. 2014; Gómez-Brandón et al. 2020). Due to limited knowledge of diazotrophic abundance and activity in deadwood of temperate forest sites, it is unclear how factors, such as C, NSC, and nutrients, influence the abundance of diazotrophs and the process of BNF or vice versa.

This study was part of a long-term experiment on deadwood decay of 13 tree species at nine forest sites at the Hainich-Dün exploratory in Central Germany. Our objective was to assess BNF rates and nifH gene copy numbers in logs of 13 tree species after 12 years of decay and identify potential relationships between wood traits and BNF rates. Moreover, we compiled available BNF rates in deadwood from the literature, considering tree species, location, and applied method, and compared the BNF rates with this study. We hypothesized that (1) BNF rates differ between tree species or between phylogenetic clades (gymnosperms, ring-porous, and diffuse-porous angiosperms) because of differences in deadwood traits, (2) nifH gene abundance is higher in angiosperms than in gymnosperms, and BNF correlate positively to nifH gene abundance, and (3) BNF rates increase with increasing concentrations of nutrients and NSC. Moreover, we used a structural equation model (SEM) to examine direct and indirect relationships between BNF rate, nifH gene abundance, nutrients, NSC, water content (WC), and respiration rate.

Material and methods

Study region and experimental design

This study was part of the long-term experiment BELongDead within the Biodiversity Exploratories project, which focuses on the interactions of tree species, microbial communities, and local conditions at three exploratories in Germany (Schwäbische Alb, Schorfheide-Chorin, and Hainich-Dün). In each exploratory, forest plots with different management intensities in threefold repetition were chosen to represent local management practices (Kahl et al. 2015).

The forests in the Hainich-Dün exploratory in Central Germany (50°56′44″-51°22′43″N/10°10′24″-10°46′45″E) were dominated by European beech (Fagus sylvatica). The study region was characterized by mean annual temperatures of 6.5 to 8 °C, mean annual precipitation of 500 to 800 mm, and altitude from 285 to 550 m a.s.l. Soils were characterized as Luvisols and Stagnosols, which developed on calcareous bedrock with a loess layer (Fischer et al. 2010; Kahl et al. 2017). N deposition was 10.1 ± 0.8 kg ha⁻¹ per vegetation period in the Hainich-Dün exploratory (Schwarz et al. 2014). Different forest management practices include selective timber forest, timber forest, and unmanaged forest (Fischer et al. 2010).

On each of the nine experimental plots in Hainich-Dün, all located in European beech forests, the BELongDead experiment was established in the winter of 2008/09. Freshly cut logs (4 m length, ~31 cm diameter) of 13 tree species were placed on the soil in random order within 1 m distance to each other. The 13 tree species included four gymnosperms (Larix decidua, Picea abies, Pinus sylvestris, and P. menziesii), seven diffuse-porous angiosperms (Acer spp., Betula pendula, Carpinus betulus, F. sylvatica, Populus spp., Prunus avium, and Tilia spp.) and two ring-porous angiosperms (Fraxinus excelsior and Quercus spp.) (Kahl et al. 2017). All logs originated from the state forest of the Federal State of Thuringia in Central Germany and were thus grown under similar climatic conditions.

Sample collection

In June 2020, deadwood samples were cut laterally with an electric saw from the outer 7 cm of each log. Deadwood samples consisted almost exclusively of sapwood, but mixtures of sapwood and heartwood were taken from Quercus logs due to the decay and loss of sapwood. One part of the sample was stored in a plastic bag in the field and kept cool (~5 °C) until further treatment. A second part of the sample (sawdust) was transferred into a 15 mL falcon tube in the field. After that, 10 mM bromodeoxyuridine...
(BrdU) was added sufficiently until the sawdust was completely soaked with solution. The falcon tubes were placed in an upright position and covered with aluminum foil, followed by incubation at room temperature for 48 h. The samples were then frozen at −80 °C in the laboratory before the characterization of the abundance of metabolically active diazotrophs (see below).

**BNF**

At the laboratory, deadwood samples (~20 cm³) were saturated with water for 16 h to achieve maximum water holding capacity in all samples, then drained, and weighted into glass flasks (~131 mL, Media Bottle Rasotherm ISO, GL45). After five days of acclimatization in the dark at 20 °C, the lids were equipped with a 6 mm thick chlorobutyl septum. After closure, the glass flasks were flushed for 10 min with a prepared gas mixture consisting of 8% oxygen (O₂), 30% nitrogen (N₂), and 62% helium (Riessner Gase GmbH, Lichtenfels, Germany). To determine the BNF rate (Weaver and Danso 1994), 25 mL of acid-washed 15N₂ (99.1%, batch number MBBC7366, Sigma-Aldrich Inc., St. Louis, MI, USA) was added to each glass flask using a syringe. The gas pressure was measured before and after the addition of 15N₂ to calculate the exact proportion of 15N₂ in the headspace of glass flasks. After adding 15N₂, the samples were incubated at 20 °C for 72 h in the dark. The flasks were then opened, and the samples were immediately dried at 60 °C for at least 72 h until mass constancy to terminate the incubation and to determine the water content. Dried samples were ground at low temperatures using a ball mill (MM400, Retsch GmbH, Haan, Germany). Incubated and respective non-incubated subsamples were analyzed for δ15N signatures and N concentration at the Centre for Stable Isotope Research and Analysis, Göttingen, using an element analyzer (Euro EA 3000, EuroVector S.p.A., Milano, Italy), coupled to a Delta C isotope mass spectrometer with a ConFlo III interface (Thermo Electron, Bremen, Germany) and an element analyzer (Euro MA 3000, EuroVector S.p.A., Milano, Italy), coupled to a Delta C isotope mass spectrometer with a ConFloII interface (Thermo Electron, Bremen, Germany) for natural abundance and 15N enriched samples, respectively.

As certified 15N₂ gas (Sigma-Aldrich Inc., St. Louis, MI, USA) contained significant amounts of NH₃ and N oxides (Dabundo et al. 2014), the 15N₂ gas was acid-washed before adding to deadwood samples. For this purpose, an acid solution with a pH of 3 to 4 was produced by adding 100 mL deionized water and ~2 µL of 2 M H₂SO₄ (95%, Chemsolute, Th. Geyer GmbH & Co.KG, Renningen, Germany) to a 650 mL glass flask. The flask was closed with a screw cap and a gas-tight, 6 mm thick chlorobutyl septum. After that, the bottle was evacuated to 200 mbar, flushed with helium for 15 min, and then shaken overhead (6 rpm) for 1 h. This procedure of evacuating, flushing, and shaking was repeated three times. Afterwards, the bottle was evacuated to 200 mbar, and 15N₂ gas was transferred directly from the 15N₂ gas container to the flask up to a pressure of ~2200 mbar. The glass flask was shaken overhead for 1 h, left standing overnight, and shaken again for 1 h before use. Gas pressure measurements determined the proportion of 15N₂ in the headspace before and after 15N₂ addition. Finally, the purification and repeated refilling with the 15N₂ gas amounted to a 15N₂ concentration of 98 to 99% in the headspace of the 650mL glass flask. To determine potential contamination of the 15N₂ gas, four wood samples were autoclaved at 121 °C with 1 bar overpressure for 20 min. The samples were incubated and processed as described above. The enrichment in the δ15N signature in the autoclaved deadwood samples was neglectable (0.003 ± 0.0007 µg 15N per ml 15N₂).

After the conversion of δ15N signatures of enriched and non-enriched (natural abundance, NA) deadwood samples into 15N atom fractions, BNF rates (µg N g⁻¹ d⁻¹) referring to the dry weight of deadwood was calculated using Eq. 1:

$$\text{BNF} = \frac{N \times (15\text{N}_{\text{enriched}} - 15\text{N}_{\text{NA}})}{\text{DW} \times t \times HS \times 100} \quad (1)$$

where N is the amount of N in deadwood (µg), 15N_{enriched} is 15N atom% in deadwood after incubation, 15N_{NA} is the 15N atom% natural abundance in deadwood, DW is the dry weight of deadwood (g), t is the incubation time (d), and HS is the ratio of 15N₂ to total N₂ in the headspace of incubation flasks.
Respiration rate

The respiration rate of deadwood was determined by measuring the increase in carbon dioxide (CO₂) concentration during the incubation. Gas samples of 100 µL were taken with a syringe from the glass jars after 21 h (t₁) and 45 h (t₂) after flushing with the CO₂-free gas mixture at t₀ (see above). The gas samples were directly injected into a gas chromatograph (SRI 8610C, SRI Instruments, Torrance, CA, USA) equipped with a flame ionization detector coupled to a methanizer catalyst. Certified gas standards (10,000 and 20,000 ppm CO₂, Riessner Gase, Lichtenfels, Germany) were used to calibrate the gas chromatograph.

The respiration rate (µg C g⁻¹ h⁻¹) was calculated using Eq. 2:

\[
\text{respiration rate} = \frac{\Delta CO_2 \Delta t}{DW} \times \frac{V_{gas}}{R \times T_{air}} \times \frac{\rho_{air}}{M} \quad (2)
\]

where \(\Delta CO_2 \Delta t\) (ppm h⁻¹) is the change in CO₂ concentration within the incubation flasks during the incubation, assuming that the initial CO₂ concentration was 0 ppm after flushing, \(V_{gas}\) (m³) is the gas volume of the incubation flasks, \(DW\) (g) is the dry weight of the deadwood sample, \(\rho_{air}\) (Pa) is the air pressure, \(R\) (8.314 J K⁻¹ mol⁻¹) is the gas constant, \(T_{air}\) (K) is the air temperature, and \(M\) (g mol⁻¹) is the molar mass of C.

NSC and elemental analyzes

The concentrations of ethanol-soluble NSC were measured at the Max Planck Institute of Biogeochemistry, Jena, as reported earlier (Landhäusser et al. 2018). Briefly, 1.5 mL of 80% ethanol was mixed with 30 mg of ground wood sample and heated to 90 °C. After cooling down to room temperature, the mixture was centrifuged at 13,000 g for 1 min. The supernatant was filtered, and about 1 mL was injected into a High-Performance Anion Exchange Chromatograph with Pulsed Amperometric Detection (DIONEX ICS-3000 with CarboPac columns, Thermo Fisher Scientific Inc., Waltham, MA, USA) to analyze concentrations of glucose, arabinose, xylose, galactose, and total NSC. We assumed that starch was not present in deadwood after 12 years of decay.

Moreover, the concentrations of calcium (Ca), magnesium (Mg), potassium (K), phosphor (P), sulfur (S), manganese (Mn), and Mo were determined using pressure digestion. 1 g of dried wood sample was filled in vessels, and 12 mL of HNO₃ (65%, Chemsololute, Th. Geyer GmbH & Co.KG, Renningen, Germany) was added. The mixture was microwaved (MARS6, CEM Corporation, Matthews, NC, USA) and filtered using sterile syringe filters (0.45 µm). Afterwards, the supernatant was analyzed at the Analytical Chemistry Lab, University Bayreuth, using an ICP-MS (Agilent 7500ce, Cetac ASX-510, Santa Clara, CA, USA) for Mo and an ICP-OES (Varian, Vista-Pro radial, Palo Alto, CA, USA) for Ca, Mg, K, P, S and Mn according to manufacture’s instructions. The C concentration was analyzed with a vario Max CN element analyzer (elementar Analysensysteme GmbH, Hanau, Germany) With the concentrations of C and N (see above), the CN ratio was calculated.

In-situ deadwood BrdU labeling and molecular biological analysis

For in-situ labeling with BrdU, 1 mL of 10 mM BrdU solution (Sigma-Aldrich Inc., St. Louis, MI, USA) was added directly after sampling to each deadwood sample (~ 1 g) and incubated in sterile 50 mL tubes covered with aluminum foil with loose cap for 48 h at room temperature as explained earlier (Purahong et al. 2022). Only the metabolically active, replicating cells can incorporate the BrdU during DNA synthesis (McMahon et al. 2011). According to the manufacturer’s instructions, DNA was extracted from the BrdU-treated deadwood samples using Quick-DNA Fecal/Soil Microbe Miniprep Kit (Zymo, California, USA). We named this total DNA as it included all types of genomic DNA (dormant cells, dead cells, metabolically active, and replicating cells). As outlined earlier (McMahon et al. 2011; Purahong et al. 2022), this immunocapture approach was employed to separate the BrdU-labeled DNA from the total DNA. Briefly, for each sample, 2 µL monoclonal BrdU antibodies (1 mg µL⁻¹ mouse anti-BrdU, clone BU-33, Sigma-Aldrich Inc., St. Louis, MI, USA) were added to 18 mL denatured herring sperm DNA (1.25 mg mL⁻¹ in phosphate buffer saline (PBS), Promega, Walldorf, Germany), and then incubated for 45 min at 30 °C to form antibody-herring sperm DNA complex. Denatured sample
DNA (25 µL; ~200 ng DNA + 10 µL PBS) was then mixed with antibody-herring sperm DNA complex and incubated for 30 min at 30 °C to capture BrdU-labeled DNA. 6.26 µL washed Dynabeads™ Goat Anti-Mouse IgG (Invitrogen, Life Technologies GmbH, Darmstadt, Germany) were then added to the incubated solution and rotated (66 rpm) for 30 min at room temperature on an RM-2 roll mixer (Carl Roth GmbH, Karlsruhe, Germany) to form the Dynabead complex (Dynabead-BrdU antibodies-BrdU-labeled DNA). After that, the Dynabead complex was treated eight times with 100 μL PBS–BSA solution (0.05 g bovine serum albumin in 50 mL phosphate-buffered saline) and washed with a DynaMag™-2 (Invitrogen, Life Technologies GmbH, Darmstadt, Germany). BrdU-labeled DNA was isolated from the washed Dynabead complex by adding 20 µL BrdU solution (1.7 mM) followed by 35 min incubation under slow rotation. Finally, the BrdU-labeled DNA was separated from the Dynabeads using a DynaMag™-2 (Life Technologies GmbH, Darmstadt, Germany), thereafter named active DNA.

The abundance of nifH gene copy numbers in deadwood samples of the total and active DNA fraction was accessed by quantitative PCR (qPCR). Diazotrophic gene copy numbers were quantified by the primer set PolF and PolR (Poly et al. 2001). All reactions were performed in 96-well plates using the CFX96™ Real-Time System (Bio-Rad Laboratories GmbH, Feldkirchen, Germany), nuclease-free master mix blanks were run as a negative control. Gene copy number was calculated by comparing PCR-cycle threshold values (CT) to a standard curve of triplicate tenfold dilutions of genomic DNA. Genomic DNA extracted from a culture of Azotobacter vinelandii (DSM 2289) by employing the Quick-DNA Fecal/Soil Microbe Miniprep Kit was used to establish quantification standards for generating a standard curve. Petroff counting chamber (Paul Marienfeld GmbH, Germany) was used to determine the genomic DNA concentration (gene copies µL⁻¹). A five-point tenfold serial dilution of the A. vinelandii genomic DNA (10–100,000 fg) was run in triplicate with each set of reactions to generate the standard curve. For nifH gene-based qPCR, the reactions were performed in 10 µL assays containing 5 µL SYBR® Green Supermix (Bio-Rad Laboratories GmbH, Feldkirchen, Germany), PolF and PolR primers (0.5 µL each of the primer (2.5 µM)), 3 µL sterile and nuclease-free water (Carl Roth GmbH, Karlsruhe, Germany) and 1 µL of either 1:10 diluted DNA-extract, ten-fold diluted standard DNA or 1 µL water as no template control. After an initial denaturation at 94 °C for 5 min, 40 amplification cycles (regardless of the active or total fraction) were performed for 1 min at 95 °C (denaturation), 1 min at 55 °C (annealing), and 1 min 30 s at 72 °C (extension), followed by a final extension of 5 min at 72 °C. Runs were completed with a melting analysis starting from 65 °C to 95 °C with temperature increments of 0.5 °C and a transition rate of 5 s to check for product specificity and potential primer dimer formation. The purity of the amplified products was checked by electrophoresis on a 1.5% agarose gel. Multiple dilutions (non-dilution, with 1:10 and 1:100 dilution) were run simultaneously to check for inhibitors in qPCR assays. We observed no inhibition as the CT value shifted for each decimal dilution step in the same CT gap. CT and efficiency were calculated automatically by the Bio-Rad software CFX manager version 3.1. Both, total and active nifH gene copy numbers refer to the dry mass of deadwood samples.

Data collection from the literature

Available BNF rates in deadwood were compiled to the best of our knowledge to compare our BNF rates with the values from the literature (Online Resource Table S1). BNF rates were taken from tables or graphs and transformed into µg N g⁻¹ d⁻¹ when values were only presented as ethylene production, based on the acetylene reduction assay (ARA, see Hardy et al. (1971)). For the transformation, we used deadwood studies in which ARA was calibrated with the ¹⁵N₂ approach (Roskoski 1981; Silvester et al. 1982; Silvester 1989; Chen and Hicks 2003; Hicks et al. 2003; Burgoyne 2007; Burgoyne and DeLuca 2009; Benoist et al. 2022). A mean ratio of 3.9:1 (range from 0.8:1 to 8.5:1) was used to estimate the BNF rate from ethylene production. Information about phylogenetic clade, tree species, deadwood age, decay stage, location, climate zone, incubation method, and incubation temperature were summarized for an overview of BNF in deadwood (Online Resource Table S1).

Statistical analyses

Statistics were performed using R version 4.0.4 (R Core Team 2021) with additional packages. Data
were screened visually for outliers with box plots and Cleveland’s dot plots. Normal distribution was checked with histograms, quantile–quantile plots, and the Shapiro–Wilk-Test. Significant differences (p < 0.05) were calculated using the Kruskal–Wallis-Test and pairwise Wilcoxon Rank Sum test for non-normal distributed data between tree species and phylogenetic clades (gymnosperms, diffuse-porous and ring-porous angiosperms). Due to the non-normal distribution of the data, correlations were assessed with Spearman’s rank correlation coefficient. Graphics were generated with the package ggplot2 (Whickham 2016).

To find direct and indirect effects of nifH gene abundance, BNF rate, respiration rate (as a proxy for fungal activity), WC, NSC, Mo, P, and N, a SEM was applied. Before analysis, a theory model with possible connections was built based on theoretical knowledge (Online Resource Fig. S1) and the significant correlation of BNF with all measured parameters. The package piecewiseSEM version 2.1.0 (Lefcheck 2016) was used for the analysis, as plot identity was included as a random factor. Before fitting the piecewise SEM, the single linear mixed-effect models within the SEM were checked for meeting model assumptions, using the package nlme version 3.1–152 (Pinheiro et al. 2021). For the piecewise SEM, the model fitting and the detection of missing paths were accomplished by Fisher’s C and Shipley’s test of d-separation, respectively (Shipley 2013). Models were considered valid when the model’s p-value was > 0.05 and the degrees of freedom (df) > 1, meaning that the piecewise SEM was not different from our data but displayed the data. For endogenous, response variables (BNF, nifH gene abundance, and respiration rate), a conditional $R^2_c$ is given. Standardized path coefficients determine relationships between variables.

Results

BNF rates compared to the literature

BNF rates from the BELongDead experiment ranged from 0.05 to 0.10 µg N g$^{-1}$ d$^{-1}$ in the gymnosperms and from 0.03 to 0.28 µg N g$^{-1}$ d$^{-1}$ in the angiosperms (Fig. 1). The median BNF rates in the gymnosperms were significantly (p=0.04) lower in the literature (0.005 µg N g$^{-1}$ d$^{-1}$) compared to the BELongDead experiment (0.075 µg N g$^{-1}$ d$^{-1}$). In contrast, angiosperms in the literature fixed similar N (both 0.14 µg N g$^{-1}$ d$^{-1}$). Deadwood of gymnosperms exhibited significantly (p=0.02) lower BNF rates than angiosperms in the literature, but not in the BELongDead experiment.

BNF and nifH gene abundance

The highest median BNF rates were detected in Populus (0.30 µg N g$^{-1}$ d$^{-1}$), Tilia, and Fraxinus (both 0.16 µg Ng$^{-1}$ d$^{-1}$), whereas the lowest rates occurred in Fagus (0.03 µg N g$^{-1}$ d$^{-1}$), Quercus, and Larix (both 0.02 µg N g$^{-1}$ d$^{-1}$) (Fig. 2). Tree species had no significant effect on BNF due to the large variation within each tree species. However, BNF rate was significantly (p=0.04) lower for gymnosperms (0.05 µg N g$^{-1}$ d$^{-1}$) than for diffuse-porous angiosperms (0.09 µg g$^{-1}$ d$^{-1}$). In contrast, ring-porous angiosperms’ BNF rates were not significantly different from gymnosperms and diffuse-porous angiosperms.

The total nifH gene abundance within the logs of all tree species was averaged at a median of 19.2 × 10$^{6}$ copies g$^{-1}$, ranging from 0.2 to 239.5 × 10$^{6}$ copies g$^{-1}$ (Fig. 3a). Low abundances were found in Quercus (7.2 × 10$^{6}$ copies g$^{-1}$) and Picea (7.3 × 10$^{6}$ copies g$^{-1}$) compared to higher abundances in Populus (53.1 × 10$^{6}$ copies g$^{-1}$), Tilia (48.4 × 10$^{6}$ copies g$^{-1}$), and Betula (46.0 × 10$^{6}$ copies g$^{-1}$). A tree species effect on the total nifH gene abundance could not be detected, but all diffuse-porous angiosperms (27.3 × 10$^{6}$ copies g$^{-1}$) had significantly higher (p=0.01) abundances than the gymnosperms (13.0 × 10$^{6}$ copies g$^{-1}$).

The active nifH gene abundance ranged from 0.0016 × 10$^{6}$ copies g$^{-1}$ to 5.2 × 10$^{6}$ copies g$^{-1}$ with a median of 0.11 × 10$^{6}$ copies g$^{-1}$ (Fig. 3b). No significant differences occurred between tree species or phylogenetic clades. The highest values were found in Tilia (0.99 × 10$^{6}$ copies g$^{-1}$) as opposed to the lowest values in Quercus and Picea (both 0.04 × 10$^{6}$ copies g$^{-1}$). Diffuse-porous angiosperms (0.18 × 10$^{6}$ copies g$^{-1}$) showed slightly higher abundances than ring-porous angiosperms (0.13 × 10$^{6}$ copies g$^{-1}$) and gymnosperms (0.09 × 10$^{6}$ copies g$^{-1}$). The active diazotrophic community accounted for 0.06 to 11.4% of the total nifH gene abundance based on BrdU analysis.
Relationship between BNF and \textit{nifH} gene abundance

Correlating BNF and total \textit{nifH} gene abundances showed a non-significant linear increase (\(R=0.18, p=0.62\)) (Fig. 4). However, low BNF rates occurred when low total \textit{nifH} gene abundances were observed. Higher total \textit{nifH} gene abundances resulted in only a slight increase in the BNF rate. The highest BNF rates appeared seemingly arbitrary when only a low total \textit{nifH} gene abundance was present. Compared to the total \textit{nifH} gene abundance, the active diazotrophic abundance had less effect on the BNF rate (\(R=0.053, p < 0.05\)).

Fig. 1 BNF rates in deadwood of angiosperm (a) and gymnosperm (g) tree species from the BELongDead experiment (BLD) and the literature (lit). For the BNF rates in the BELongDead experiment, the mean for each tree species was calculated. For the literature, one mean BNF rate per study was calculated. Corresponding references and BNF rates are given in Online Resource Table S1. The number of repetitions (n) per boxplot is given above the boxplot. Boxplots show the median of the values. Each boxplot’s upper and lower edges represent the 25 and 75% quartile, respectively. The whiskers represent the \(1.5 \times \text{inter-quartile range}\). Points beyond the whiskers are outliers. Significant differences between all boxplots are denoted as capital letters (\(p<0.05\)).

Fig. 2 BNF rates in deadwood after 12 years of decomposition of 13 tree species. Boxplots show the median of the values. Each boxplot’s upper and lower edges represent the 25 and 75% quartile, respectively. The whiskers represent the \(1.5 \times \text{inter-quartile range}\). Points beyond the whiskers are outliers. Significant differences between phylogenetic clades are denoted as capital letters (\(p < 0.05\)).
p = 0.58) (data not shown). Even though only a small number of diazotrophs were metabolically active, their activity did not correlate with BNF. Apart from BNF rates, active and total \( nifH \) gene abundances were significantly correlated with each other (\( R = 0.67, p < 0.001 \)) (data not shown).

Respiration rate

The respiration rate in deadwood ranged from 1 to 58 µg C g\(^{-1}\) h\(^{-1}\) (Fig. 5). The effect of tree species was significant (\( p < 0.001 \)) between some tree species, mainly angiosperms to gymnosperms. The highest rates occurred in *Tilia* (29 µg C g\(^{-1}\) h\(^{-1}\)) and *Carpinus* (28 µg C g\(^{-1}\) h\(^{-1}\)) compared to the lowest rates in *Picea* and *Pinus* (both 8 µg C g\(^{-1}\) h\(^{-1}\)). Diffuse-porous (21 µg C g\(^{-1}\) h\(^{-1}\)) and ring-porous angiosperms (20 µg C g\(^{-1}\) h\(^{-1}\)) showed significantly (\( p < 0.001 \) and \( p = 0.02 \), respectively) higher respiration rates than gymnosperms (8 µg C g\(^{-1}\) h\(^{-1}\)). Respiration rate had a significant effect on BNF (\( R = 0.29, p = 0.002 \)) (Online Resource Fig. S2).
NSC and nutrient concentrations in deadwood and relationships to BNF

Deadwood showed a wide variety of glucose (0 to 8 mg g\(^{-1}\)), arabinose, xylose (both 0 to 1.1 mg g\(^{-1}\)) and galactose (0 to 0.2 mg g\(^{-1}\)) (Table 1). The most abundant NSC was glucose, with around 74% of total NSC. Diffuse-porous angiosperms showed higher glucose, xylose, and total NSC concentrations than ring-porous angiosperms and gymnosperms. Differences were partly significant, especially for angiosperms and gymnosperms. C concentrations were similar for all deadwood logs (~47%).

Similar to NSC, nutrient concentrations showed a great spread within the deadwood logs, such as Mo (0 to 1.6 µg g\(^{-1}\)), Ca (0.2 to 17.5 mg g\(^{-1}\)), Mg (0.04 to 2.5 mg g\(^{-1}\)), K (0.03 to 4 mg g\(^{-1}\)), P (0 to 0.6 mg g\(^{-1}\)), S (0.03 to 0.9 mg g\(^{-1}\)), and Mn (0 to 1.4 mg g\(^{-1}\)) (Table 2). Diffuse-porous angiosperms showed significantly higher concentrations of Mg (p<0.001) and non-significant higher concentrations of Mo, Ca, P, S, and Mn compared to ring-porous angiosperms and gymnosperms. Significant differences for S were found between diffuse-porous angiosperms and gymnosperms (p=0.008). Mo and P did not show significant differences between phylogenetic clades.

The relationship between BNF, NSC and nutrients, including N, was investigated using a Spearman rank correlation coefficient (Online Resource Fig. S2). There were significant, slightly positive correlation between BNF and Mo (R = 0.21, p = 0.04), P (R = 0.26, p = 0.007), S (R = 0.22, p = 0.02), and N (R = 0.43, p < 0.001). Glucose had an almost significant positive effect on BNF (R = 0.18, p = 0.06). Concentrations of Mn, Ca, Mg, K, xylose, galactose, and total NSC had a non-significant effect on the BNF rate.

Structural equation modeling

As variability in BNF rates and \(nifH\) gene abundance could not be explained by tree species and only partially by a variety of other factors (see above), a SEM was applied to determine both direct and indirect effects of the significant, most correlated parameters on BNF and \(nifH\) gene abundance (Fig. 6). The SEM
revealed that the selected parameters only partially explained the $\text{nifH}$ gene abundance ($R^2_c = 0.15$) and the BNF rates ($R^2_c = 0.10$). BNF rates were significantly correlated with WC (standardized path coefficient 0.27, $p = 0.02$), while $\text{nifH}$ gene abundance was significantly correlated to total NSC (standardized path coefficient 0.28, $p = 0.006$) and WC (standardized path coefficient 0.35, $p = 0.004$). No significant correlation was found between $\text{nifH}$ gene abundance and BNF (standardized path coefficient 0.06, $p = 0.59$).

**Discussion**

BNF in deadwood – comparison to the literature

Substantial BNF rates in deadwood were observed in almost all studies, but the variability within and among the studies is high. As we found no significant differences between individual tree species (see below), we compared available BNF rates of angiosperms and gymnosperms. Despite different techniques, decay stages, and incubation temperatures (Online Resource Table S1), median BNF rates of angiosperms were similar in our study and the literature. By contrast, BNF rates of gymnosperms were 15 times higher in our study than in the literature (Fig. 1). We attributed the high BNF rates of gymnosperms in our study primarily to the N saturation of forests in Central Europe, resulting from decades of increased N deposition (Borken and Matzner 2004; Ackerman et al. 2019). All other studies on BNF in deadwood of gymnosperms were performed in North America or Northern Europe, where N deposition is much smaller (Ackerman et al. 2019). Since specific N availability is required to develop and establish diazotrophic communities (Reed et al. 2011), low initial N concentrations may particularly limit BNF in deadwood of gymnosperms (see below). The fact that no difference was found for deadwood of angiosperms between our study and the literature could be explained by generally higher N concentrations in angiosperm deadwood.

Different measurement techniques probably caused differences in BNF rates between our and other studies. Previous studies almost exclusively used ARA as an indirect measurement of nitrogenase activity (Hardy et al. 1971). Methodological problems could have led to an over- or underestimation of BNF rates.

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**Table 1** Mean concentrations ($n=9,$ ± SD) of C, N, CN ratio, glucose, arabinose, xylose, galactose, and total NSC of deadwood from 13 tree species and phylogenetic clades (DP = diffuse-porous, RP = ring-porous). Significant differences between phylogenetic clades are denoted as capital letters ($p < 0.05$)

| Tree species/ | C (%) | N (%) | CN ratio | Glucose (mg g$^{-1}$) | Arabinose (mg g$^{-1}$) | Xylose (mg g$^{-1}$) | Galactose (mg g$^{-1}$) | Total NSC (mg g$^{-1}$) |
|---------------|-------|-------|----------|-----------------------|------------------------|---------------------|------------------------|----------------------------|
| Acer          | 46.8 ± 1.7 | 0.38 ± 0.23 | 171 ± 101 | 0.47 ± 0.30          | 0.010 ± 0.02          | 0.13 ± 0.10         | 0.01 ± 0.02            | 0.63 ± 0.35                |
| Betula        | 47.2 ± 1.1 | 0.27 ± 0.12 | 193 ± 52 | 2.34 ± 2.58           | 0.000 ± 0.00          | 0.41 ± 0.40         | 0.02 ± 0.02            | 2.77 ± 2.98                |
| Carpinus      | 46.6 ± 1.3 | 0.44 ± 0.26 | 144 ± 77 | 0.70 ± 0.48           | 0.007 ± 0.01          | 0.16 ± 0.10         | 0.01 ± 0.01            | 0.88 ± 0.56                |
| Fagus         | 47.8 ± 1.4 | 0.38 ± 0.16 | 147 ± 76 | 0.85 ± 0.34           | 0.003 ± 0.01          | 0.18 ± 0.19         | 0.01 ± 0.01            | 1.05 ± 0.36                |
| Populus       | 47.2 ± 1.3 | 0.39 ± 0.19 | 152 ± 86 | 0.91 ± 0.51           | 0.007 ± 0.02          | 0.14 ± 0.14         | 0.00 ± 0.01            | 1.06 ± 0.61                |
| Prunus        | 48.1 ± 1.3 | 0.41 ± 0.29 | 161 ± 82 | 0.68 ± 0.84           | 0.006 ± 0.01          | 0.18 ± 0.18         | 0.04 ± 0.08            | 0.90 ± 0.98                |
| Tilia         | 46.5 ± 1.4 | 0.41 ± 0.27 | 144 ± 59 | 1.91 ± 2.00           | 0.002 ± 0.01          | 0.20 ± 0.24         | 0.01 ± 0.01            | 2.13 ± 2.21                |
| Fraxinus      | 48.0 ± 0.5 | 0.29 ± 0.12 | 192 ± 75 | 0.69 ± 0.43           | 0.004 ± 0.01          | 0.14 ± 0.15         | 0.03 ± 0.02            | 0.86 ± 0.53                |
| Quercus       | 47.9 ± 1.0 | 0.28 ± 0.16 | 226 ± 116 | 0.92 ± 0.60          | 0.449 ± 0.45          | 0.21 ± 0.19         | 0.04 ± 0.06            | 1.62 ± 1.09                |
| Larix         | 47.3 ± 0.5 | 0.12 ± 0.08 | 621 ± 449 | 0.21 ± 0.14          | 0.235 ± 0.27          | 0.10 ± 0.25         | 0.06 ± 0.04            | 0.96 ± 0.97                |
| Picea         | 48.5 ± 1.8 | 0.19 ± 0.12 | 398 ± 246 | 0.48 ± 0.48           | 0.017 ± 0.03          | 0.15 ± 0.27         | 0.01 ± 0.02            | 0.65 ± 0.79                |
| Pinus         | 47.4 ± 0.9 | 0.16 ± 0.08 | 439 ± 395 | 0.52 ± 0.17           | 0.029 ± 0.03          | 0.14 ± 0.10         | 0.01 ± 0.02            | 0.69 ± 0.23                |
| Pseudotsuga   | 47.9 ± 1.1 | 0.21 ± 0.10 | 344 ± 290 | 0.27 ± 0.24           | 0.021 ± 0.03          | 0.03 ± 0.05         | 0.01 ± 0.02            | 0.33 ± 0.28                |
| DP angiosperms| 47.1 ± 1.4A| 0.39 ± 0.22A| 158 ± 75A | 1.16 ± 2.58A         | 0.005 ± 0.01A         | 0.21 ± 0.23A        | 0.01 ± 0.03A           | 1.39 ± 1.65A               |
| RP angiosperms| 47.9 ± 0.8A| 0.28 ± 0.14A| 209 ± 97A | 0.80 ± 0.34A         | 0.227 ± 0.38B         | 0.18 ± 0.17A        | 0.03 ± 0.05A           | 1.24 ± 0.92A               |
| Gymnosperms   | 47.8 ± 1.2A| 0.17 ± 0.10B| 450 ± 355B | 0.37 ± 0.24B        | 0.071 ± 0.15B         | 0.11 ± 0.19B        | 0.02 ± 0.03A           | 0.66 ± 0.66B               |
Table 2: Mean concentrations (n = 9, ±SD) of Mo, Ca, Mg, K, P, S, and Mn, and WC of deadwood from 13 tree species and phylogenetic clades (DP = diffuse-porous, RP = ring-porous). Significant differences between phylogenetic clades are denoted as capital letters (p < 0.05).

| Tree species / phylogenetic clade | Mo (μg g⁻¹) | Ca (mg g⁻¹) | Mg (mg g⁻¹) | K (mg g⁻¹) | P (mg g⁻¹) | S (mg g⁻¹) | Mn (mg g⁻¹) | WC (g g⁻¹) |
|-----------------------------------|-------------|-------------|-------------|------------|------------|------------|-------------|------------|
| Acer                              | 0.651 ± 0.60| 4.80 ± 3.16 | 0.82 ± 0.61 | 0.74 ± 0.39 | 0.18 ± 0.11 | 0.32 ± 0.16 | 0.25 ± 0.49 | 4.60 ± 2.83 |
| Betula                            | 0.025 ± 0.03| 1.48 ± 1.01 | 0.17 ± 0.11 | 0.63 ± 0.46 | 0.11 ± 0.06 | 0.14 ± 0.06 | 0.20 ± 0.12 | 5.04 ± 4.27 |
| Carpinus                          | 0.025 ± 0.02| 4.60 ± 5.07 | 0.32 ± 0.21 | 0.60 ± 0.40 | 0.18 ± 0.15 | 0.31 ± 0.23 | 0.22 ± 0.13 | 4.35 ± 2.77 |
| Fagus                             | 0.029 ± 0.04| 3.55 ± 2.18 | 0.50 ± 0.45 | 0.78 ± 0.34 | 0.17 ± 0.11 | 0.29 ± 0.16 | 0.31 ± 0.40 | 5.66 ± 3.22 |
| Populus                           | 0.022 ± 0.02| 5.21 ± 4.92 | 0.75 ± 0.59 | 1.53 ± 0.91 | 0.21 ± 0.10 | 0.30 ± 0.15 | 0.18 ± 0.24 | 7.46 ± 4.83 |
| Prunus                            | 0.028 ± 0.02| 1.95 ± 1.42 | 0.15 ± 0.11 | 0.41 ± 0.34 | 0.18 ± 0.22 | 0.22 ± 0.25 | 0.05 ± 0.05 | 3.23 ± 3.16 |
| Tilia                             | 0.073 ± 0.12| 3.41 ± 1.48 | 0.58 ± 0.78 | 0.89 ± 0.54 | 0.20 ± 0.15 | 0.31 ± 0.18 | 0.06 ± 0.08 | 6.43 ± 2.72 |
| Fraxinus                          | 0.028 ± 0.03| 2.80 ± 1.44 | 0.25 ± 0.09 | 0.98 ± 0.39 | 0.15 ± 0.06 | 0.24 ± 0.07 | 0.16 ± 0.39 | 3.15 ± 2.20 |
| Quercus                           | 0.029 ± 0.02| 2.44 ± 2.50 | 0.10 ± 0.12 | 0.50 ± 0.31 | 0.11 ± 0.17 | 0.18 ± 0.20 | 0.10 ± 0.16 | 2.29 ± 1.46 |
| Larix                             | 0.020 ± 0.02| 2.20 ± 2.16 | 0.53 ± 0.80 | 0.96 ± 1.19 | 0.11 ± 0.11 | 0.16 ± 0.13 | 0.03 ± 0.03 | 2.51 ± 2.06 |
| Picea                             | 0.017 ± 0.02| 1.43 ± 0.64 | 0.16 ± 0.12 | 0.55 ± 0.36 | 0.06 ± 0.05 | 0.11 ± 0.05 | 0.17 ± 0.15 | 2.80 ± 1.15 |
| Pinus                             | 0.031 ± 0.02| 2.26 ± 1.99 | 0.25 ± 0.23 | 1.09 ± 1.33 | 0.25 ± 0.21 | 0.34 ± 0.31 | 0.11 ± 0.06 | 3.59 ± 2.44 |
| Pseudotsuga                       | 0.023 ± 0.02| 1.17 ± 1.46 | 0.15 ± 0.21 | 0.54 ± 0.71 | 0.12 ± 0.11 | 0.16 ± 0.15 | 0.07 ± 0.07 | 3.01 ± 2.23 |
| DP angiosperms                    | 0.120 ± 0.31A| 3.61 ± 3.34A| 0.48 ± 0.52A| 0.81 ± 0.60A| 0.18 ± 0.13A| 0.27 ± 0.18A| 0.18 ± 0.27A| 5.33 ± 3.56A|
| RP angiosperms                    | 0.029 ± 0.02A| 2.62 ± 1.99A| 0.18 ± 0.13B| 0.74 ± 0.42A| 0.13 ± 0.13A| 0.21 ± 0.15AB| 0.13 ± 0.29B| 2.72 ± 1.87B|
| Gymnosperms                       | 0.023 ± 0.02A| 1.77 ± 1.67A| 0.27 ± 0.44B| 0.79 ± 0.96A| 0.14 ± 0.15A| 0.19 ± 0.20B| 0.09 ± 0.10AB| 2.98 ± 1.98B|

Fig. 6: Structural equation model for logs from 13 tree species after 12 years of decomposition, relating BNF, nifH gene abundance, respiration rate, NSC, WC, N, P, and Mo. Single-headed arrows indicate a direct effect of one parameter on the other, whereas double-headed arrows indicate a mutual relationship between two parameters. Standardized path coefficients are shown next to the arrow. Solid lines indicate significant and dashed lines non-significant effects (p < 0.05). R_c^2 is given under the endogenous parameter. Model Fisher’s C 7.271 (p = 0.51, df = 8).
These include highly variable conversion factors from reduced acetylene (ethylene) to fixed N₂ in deadwood and other substrates (Soper et al. 2021) or the dampening activity of acetylene on nitrogenase (de Bont and Mulder 1976). Additionally, contamination of the ¹⁵N₂ gas by reactive N overestimates BNF rates (Dabundo et al. 2014), particularly in N poor habitats such as deadwood. Thus, removing reactive N from ¹⁵N₂ gas and sterilized controls is a prerequisite for the correct determination of BNF rates.

No tree species effect on BNF

Our first hypothesis that tree species influence BNF in logs was not confirmed. The variability of BNF was high within each of the 13 tree species and therefore did not result in any statistically verifiable difference. Still, higher BNF rates were found in diffuse-porous angiosperms than in gymnosperms, perhaps due to better nutrient availability for diazotrophs (see below). The greater mass loss of those logs (Kahl et al. 2017) also indicates the higher activity of deadwood degrading fungi and more readily available NSC in diffuse-porous angiosperm logs. Low BNF rates in Quercus and gymnosperm logs correlate with high phenolic compounds, which suppress the microbial activity and decay of deadwood (Aloui et al. 2004; Kahl et al. 2017). Hence, different levels of nutrients, phenolic compounds, and decay stages could have contributed to the high variability of BNF between the 13 tree species.

In contrast to our study, some studies reported an effect of tree species on BNF. Those studies mainly examined differences between gymnosperms, such as Picea sitchensis, Pinus ponderosa, P. menziesii, or Tsuga heterophylla (Griffiths et al. 1993; Chen and Hicks 2003; Hicks et al. 2003). Of the few studies comparing angiosperms and gymnosperms, Populus tremuloides exhibited two to four times higher BNF rates than Acer rubrum, Betula papyrifera, Pinus resinosa, and Pinus strobus and ten times higher BNF rates than Picea glauca (Hendrickson 1988). Likewise, Mäkipää et al. (2018a) reported higher BNF rates in P. abies than B. pendula. However, large differences in BNF rates are also known for the same tree species as for F. sylvatica and P. abies (Mäkipää et al. 2018a, b; Rinne et al. 2017; Tláskal et al. 2021). This shows that differences between tree species cannot be generalized, and that local factors and deadwood traits could affect the BNF rates.

Abundance of nifH genes and relation to BNF

We could not find a tree species effect on nifH gene abundance or a positive relationship between BNF and nifH gene abundance, leading to rejecting the second hypothesis. Analogous to the total nifH gene abundance, we could not find a correlation between BNF and active diazotrophs. Based on BrdU labeling, only 0.1 to 11.4% of the diazotrophs were metabolically active under field conditions, suggesting an immense potential for diazotrophs in deadwood. Incubation under laboratory conditions with adjusted temperature and WC could have increased the BNF rate, while the activity status in the diazotrophic community was recorded directly at the field site. An in-situ ¹⁵N₂ incubation might have resulted in a closer relationship between BNF and active diazotrophs. Consistent with our study, Mäkipää et al. (2018b) found no association between BNF and nifH genes in deadwood. We cannot exclude an underestimation of active diazotrophs by incomplete BrdU uptake during the 48 h incubation, especially of slow growing taxa (Wahdan et al. 2021). Moreover, specific diazotrophic community patterns could be responsible for the observed patterns in BNF (Das et al. 2020; Wu et al. 2021). It was shown that the nifH gene abundance is not inevitable the best proxy for diazotrophs, as pseudo-nifH sequences could over- or underestimate the diazotrophic abundance during sequencing (Mise et al. 2021). More research is needed to improve our understanding of nitrogen fixation rates and functioning of diazotrophs’ nitrogenase enzyme.

Relation between BNF and NSC and nutrients

Our third hypothesis was not confirmed, as BNF rates did not increase with the increasing concentration of NSC in logs. However, glucose showed a weak positive relationship with BNF and made up 47% of total NSC. Glucose is an intermediate of cellulose degradation, and its concentrations in our samples are in the range reported in another study (Di Lella et al. 2019). The exceptional NSC availability in deadwood becomes evident when comparing glucose availability in other habitats of free-living
diazotrophs, such as soils with 1,000 to 10,000 times lower NSC concentrations (Ratnayake et al. 2013). Despite strong competition for resources among different fungi and other organisms (Hiscox et al. 2018), the supply of NSCs to diazotrophic organisms appears to be relatively good over many years of log decay. NSC is an energy source for many diazotrophs (Khan et al. 2021), and C availability was shown to drive BNF (Vitousek et al. 2002). However, the addition of glucose to deadwood led to decreased BNF rates (Hendrickson 1991). Some diazotrophs, abundant in deadwood, utilize methane as energy and carbon source instead of NSC (Mäkipää et al. 2018b), especially when methane levels in deadwood are higher than in the atmosphere (Covey et al. 2016). Other simple C compounds like methanol can also be utilized by some diazotrophs (Benoist et al. 2022). This repertoire of C sources would reduce the direct dependency of many diazotrophs on glucose, although glucose can be degraded to methane via fermentation processes under anoxic conditions.

Except for C, deadwood is an extremely nutrient-poor habitat in which several nutrients could concurrently limit the growth of diazotrophs. Differences in BNF rates between logs of diffuse-porous angiosperm and gymnosperm species could be related to the generally lower nutrient levels in deadwood of gymnosperms. Nitrogen had the most positive relation with BNF among various analyzed nutrients, suggesting severe N limitation of diazotrophs. Reed et al. (2011) postulated that the growth of diazotrophs or BNF could be limited at extremely low N availability. With initial N concentrations often < 0.1%, deadwood is much more N-depleted than leaf and needle litter with 0.5 to 0.6% N (Taylor et al. 1989; Persoh and Borken 2017; Rinne et al. 2017). It is still uncertain whether low N availability in deadwood can hamper the establishment of diazotrophic communities. Yet, other N sources could contribute to the initial development of diazotrophs and fungi in logs. Atmospheric deposition together with canopy leaching represent a possible pathway for uptake of inorganic and organic N. In the Hainich-Dün exploratory, atmospheric N deposition was relatively high with 10 kg N ha⁻¹ per vegetation period (Schwarz et al. 2014) as compared to other regions during the time of the BNF studies, such as northwest America and northern Europe (0.8 to 6 kg N ha⁻¹ yr⁻¹) (Sollins et al. 1980; Vitousek 1994; Merilä et al. 2014). Elevated N deposition in Central Europe could positively affect BNF in the initial phase of log decay since the N level in wood, especially of coniferous tree species, is very low (Kahl et al. 2017). Mutually related increases in N concentrations and BNF can be assumed for most logs in our study since concentrations were still relatively low with 0.2 to 0.4% N. Despite the high atmospheric N deposition at our study sites, BNF is still limited by low N concentration in deadwood after 12 years of decay.

Moreover, we could not find a correlation between BNF and the δ¹⁵N signature, but a tree species effect on the δ¹⁵N signature (Online Resource Fig. S3). Generally, BNF reflects the δ¹⁵N signature of atmospheric N₂ (~ 0‰) (Hobbie et al. 2020). However, other processes, such as the translocation of N from underlying soil into deadwood or atmospheric N deposition, may co-occur. These processes change the δ¹⁵N signature to more negative values (Knorr et al. 2015; Sabo et al. 2016) and make it difficult to assign the origins of N. Additionally, BNF was measured after 12 years of decay, but the δ¹⁵N signatures partly originate from N uptake during the tree’s life.

As for P, the concentrations of P, S, and Mo of the 13 log species were also very low and positively correlated to BNF. A co-limitation of BNF by these essential nutrients was likely. However, it is unclear which nutrient had the greatest limiting effect on diazotrophs in deadwood. Low levels of available P constrained the growth of diazotroph microorganisms and the ATP-dependent enzymatic conversion of N₂ to NH₃ by nitrogenase (Hardy et al. 1971). The average increase in deadwood P concentration by 75% in our study between 2009 and 2020 (Kahl et al. 2017, Table 1) resulted mainly from the C loss during the decay. In addition, deadwood decaying fungi can transfer P from the underlying soil into deadwood via their hyphal system to overcome P deficiency (Boddy and Watkinson 1995). Enrichment of P could enhance the growth of diazotrophs and the BNF rate in logs, but studies on the response of BNF to P addition in deadwood are lacking to our knowledge. Strong positive effects of P addition on BNF were reported for soil (Barron et al. 2009), mosses (Rousk et al. 2017), and leaf litter (Reed et al. 2013). Although, Barron et al. (2009) suggested that P fertilizers could be contaminated with Mo, meaning that additional P is not exclusively responsible for increasing BNF rates.
Mo is an essential element for many diazotrophs through its function in the transfer of electrons in the nitrogenase enzyme complex (Burris and Roberts 1993). Except for Acer, levels of Mo in logs were very low, although the correlations to BNF were also significant without Acer (not shown). A positive correlation between Mo and BNF has been found in studies for soil (Perakis et al. 2017), mosses (Rousk et al. 2017), and litter (Silvester 1989), but no effect was found for deadwood (Benoist et al. 2022). Due to the scarcity of Mo in several ecosystems, diazotrophs can also fix N using alternative nitrogenases, which are encoded as *anfH* and *vnfH* and depend on iron or vanadium (V) instead of Mo (Bellenger et al. 2020). Benoist et al. (2022) suggested that alternative nitrogenases could play a more prominent role in BNF since V is more abundant in deadwood than Mo. Hence, *vnfH* could be responsible for the weak correlation between BNF and *nifH* gene abundance in our study.

Compared to N and P, the relevance of S availability for the growth of non-symbiotic diazotrophs or BNF was barely studied in terrestrial ecosystems (Silvester 1989). Increasing S and K concentrations seemed to accelerate deadwood decay while other elements had no effect (Kahl et al. 2017). The positive correlation between BNF and S points to a mutual relation between fungi and diazotrophs, as Hoppe et al. (2014) suggested. Since S promotes both growth and activity of fungi (Schmalenberger et al. 2011), more C and other nutrients, such as P and Mo, could become available for diazotrophs (see above). In exchange, N becomes available to fungi, thus, supporting the idea of a mutual relationship between fungi and diazotrophs.

**BNF and *nifH* gene abundance is mainly driven by WC**

A SEM was applied to explore direct and indirect relationships between BNF, *nifH* abundance, and potential drivers. WC was the best explanatory variable for BNF and *nifH* gene abundance, reflecting higher diazotrophy and water holding capacity in diffuse-porous angiosperms versus ring-porous angiosperms and gymnosperms. All deadwood samples were adjusted to maximum water holding capacity, but WC strongly differed because of varying wood densities and capacities to store water. Positive correlations of BNF and WC in deadwood were associated with O₂ availability since O₂ is an inhibitor of the nitrogenase and can dampen BNF (Larsen et al. 1978; Burris and Roberts 1993; Hicks 2000). Thus, WC below water holding capacity could reduce BNF and *nifH* gene abundance in logs under field conditions. Other factors, such as the decay stage (Hendrickson 1988) or chemical properties of deadwood, including carbohydrates and N (Cushon and Feller 1989), play a minor role.

As mentioned above, NSC and nutrients, such as Mo, and N, are crucial for the BNF activity as they serve as an energy source (Khan et al. 2021), act as a metal cofactor in the *nifH* gene (Bellenger et al. 2020) or activate BNF at low N concentrations (Reed et al. 2011), respectively. However, we could not find such relationships in the SEM (Online Resource Fig. S2), and the SEM explained only a little part of the variability. The low explanatory power indicates the complex control and high variability of BNF in deadwood.

**Conclusions**

Our literature evaluation demonstrates that BNF is a ubiquitous and fundamental microbial process in deadwood of all investigated tree species across different biomes. High atmospheric N deposition in Central Europe does not suppress BNF in deadwood. In contrast, BNF in deadwood of gymnosperms with very low N levels could even benefit from this atmospheric N input. Regardless of N deposition, deadwood from diffuse-porous angiosperm tree species with relatively high decomposability and high initial nutrient concentrations has greater potential for BNF than deadwood from other phylogenetic tree clades. The high variability of BNF rates and *nifH* gene abundances within individual tree species suggests a complex control of diazotrophs through abiotic factors and saprotrophic organisms. Among the abiotic factors, WC and the associated control of O₂ availability are most effective in controlling BNF in deadwood. BNF is relevant for the N supply of saprotrophic organisms and can substantially contribute to the gradual enrichment of N in deadwood during the long-term decay process. Given the low exploratory power of abiotic factors, the interspecies interaction patterns with the emphasis on diazotrophs...
and saprotrophic organisms could strongly affect the activity of diazotrophs.

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Author contributions

CG performed the research and analyzed the data; SH provided the nifH gene abundance; HH provided data about NSC concentrations; CG and WB wrote the paper with contributions from all authors; WB and MN designed the study.

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Data availability

This work is based on data elaborated by the project Woodstock of the Biodiversity Exploratories program (DFG Priority Program 1374). The data (ID 31321) are available in the Biodiversity Exploratories Information System (https://doi.org/10.25829/bexis.31.321-4).

Declarations

Conflict of interest

The authors have no relevant financial or non-financial interests to disclose.

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