

Article

Carbon Dioxide and Methane Formation in Norway Spruce Stems Infected by White-Rot Fungi

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Abstract: Globally, billions of tons of carbon sequestered in trees are annually recycled back to the atmosphere through wood decomposition by microbes. In Norway, every fifth Norway spruce shows at final harvest infection by pathogenic white-rot fungi in the genera *Heterobasidion* and *Armillaria*. As these fungi can mineralize all components of wood, we predicted that they have a significant carbon footprint. Gas samples taken from infected stems were analyzed for CO₂ and CH₄ concentrations, and wood samples from different parts of the decay columns were incubated under hypoxic (4% O₂) and anoxic laboratory conditions. In spring and summer the stem concentrations of CO₂ were generally two times higher in trees with heartwood decay than in healthy trees. For most of the healthy trees and trees with heartwood decay, mean stem concentrations of CH₄ were comparable to ambient air, and only some *Armillaria* infected trees showed moderately elevated CH₄. Consistently, low CH₄ production potentials were recorded in the laboratory experiment. Up-scaling of CO₂ efflux due to wood decay in living trees suggests that the balance between carbon sequestration and emission may be substantially influenced in stands with high frequency of advanced root and stem heartwood decay.

Keywords: *Armillaria*; climate change; forest management; *Heterobasidion*; wood decay; CO₂; CH₄

1. Introduction

The global carbon cycle involves exchange of carbon between the atmosphere and marine and terrestrial biospheres, and sediments that include fossil fuels. For the past several decades, burning of fossil fuels and other human activities such as deforestation have produced a net CO_2 increase in the atmosphere of 1–2 Pg (= 10^{15} g) each year [1]. The increasing atmospheric concentration of CO_2 and other greenhouse gases (GHG) has created a measurable increase in global temperature [2].

Future climate changes depend on the magnitude and dynamics of both anthropogenic and natural GHG sources and sinks. Regarding the latter, forests play a major role in carbon cycling as they account for 80% of terrestrial carbon [3]. With a typical wood density of Scots pine and Norway spruce of ~400 kg dry matter/m³ and 50% carbon content, the carbon stored in 1m³ of conifer wood equals ~0.73 tCO₂. In the past few decades, the world's forests have absorbed as much as 30% of the total anthropogenic CO₂ emissions [4]. Thus, maintenance of forest area, reforestation of former agricultural land, increased use of woody products and production of wood biomass for bio-energy are measures included in the climate change mitigation portfolio of the forest sector [2].

Aerobic white-rot fungi are the only organisms able to efficiently mineralize all cell wall components of wood to carbon dioxide and water, and therefore these fungi play a central role in carbon and nitrogen cycling in forests [5]. Disruptions in this cycle are caused by pathogenic microorganisms that attack living trees, reduce tree growth and cause premature tree mortality. These effects in turn reduce the carbon sequestration potential of forests, and accelerate cycling of sequestered carbon. It has been estimated that globally 85 billion tons of carbon are annually recycled back into the atmosphere through the process of lignocellulose decay by microbes [6].

Besides CO₂, decomposition of wood can produce CH₄, a very potent GHG. Methane in stem wood has traditionally been believed to originate from microbial activity where anaerobic bacteria first degrade wood cell wall components to acetate, formate, H₂ and CO₂, which are in turn transformed to CO₂ and CH₄ by co-existing methanogenic archaea [7]. Anaerobic conditions in tree stems that lead to CH₄ formation may be associated with bacterial wetwood, especially on poorly-drained soils [8]. Anaerobic or microaerobic conditions can also occur in the heartwood of trees growing on well-drained upland soils to an extent that the mineralization of heartwood by associated decay fungi is hampered [9,10]. Under such conditions the intermediate degradation products from lignocellulose decomposition may provide substrates for anaerobic bacteria and archaea, which in turn can accelerate fungal growth by removing their metabolic waste products and by supplying nitrogen through N-fixation [11]. In addition, it has also been proposed that wood decay fungi themselves can produce small amounts of CH₄ without the involvement of methanogenic archaea [12].

The white-rot fungi *Heterobasidion annosum* sensu lato and *Armillaria* spp. are the most important root and heartwood decaying fungi of Norway spruce, the ecologically dominant conifer that grows on mesophile, well-drained soils in the boreal region. In particular, the *Heterobasidion* species have benefited from the current forest practice. Primary infection of stands by these fungi takes place through colonization of fresh stumps created during active sporulation of the fungus. In addition, wounds on roots created by mechanical logging during snowless and frost-free periods serve as infection routes. From infected stumps and trees the mycelia of these fungi spread to neighboring trees through root contacts [13]. In managed Norway spruce forests within the Nordic countries, commonly

every fifth Norway spruce is infected by *H. annosum* s.l. by the time of final harvest [14,15]. The heartwood-bound decay columns caused by these species in Norway spruce can rise up to 10–15 m above ground level at the late stage of infection [13]. Ability to grow under microaerobic conditions is typical to many heartwood associated decay fungi—upon culturing on malt extract agar *H. annosum* s.l. can grow at oxygen pressures down to (2 mm Hg) [16], which equals 0.26 atm% O₂. These fungi can also produce anaerobically small amounts of CO₂ as a result of alcohol fermentation [17]. At stem base of Norway spruce *Heterobasidion* species often co-occur with white-rot fungi in the genus *Armillaria*. When the two coexist, *Armillaria* species usually occupy outer heartwood, while *Heterobasidion* is associated with the inner heartwood [18]. Root infection by *Armillaria* species is often associated with predisposing abiotic conditions such as poorly drained soils [19].

Taking into account the high frequency of extensive stem decay columns caused by *H. annosum* s.l. in managed Norway spruce stands in Northern Europe, we hypothesized that the emission of CO₂ and possibly also CH₄ from colonized wood could amount to regionally significant levels. To examine this, we monitored stem CO₂ and CH₄ concentrations in healthy Norway spruce trees and trees infected by *H. annosum* s.l. and/or *Armillaria* spp. over two growing seasons. After completing the field monitoring, the infected tree with the highest concentration of methane was felled and CO₂ and CH₄ production potentials from defined wood tissues were measured under laboratory conditions.

2. Materials and Methods

2.1. Sites

Norway spruce (*Picea abies*) trees were subjected to field monitoring at seven upland stands on well-drained soil in Eastern Norway: Ås-1 (seven trees), 59°41′31″ N 10°45′04″ E; Ås-2 (three trees), 59°40′05″ N 10°42′44″ E; Hokksund (nine trees), 59°44′28″ N 09°55′32″ E; Haukeli-1 (five trees), 59°44′18″ N 7°33′09″ E, Haukeli-2 (two trees), 59°44′00″ N 7°32′02″ E; Jevnaker-1 (eight trees), 60°15′49″ N 10°30′06″ E; Jevnaker-2 (three trees), 60°18′04″ N 10°23′22″ E. The sampling started in the spring 2013 at Ås and in July at Hokksund and Haukeli, whereas at Jevnaker the trees were sampled in the summer of 2014. Trees with heartrot were localized by collecting increment cores from stem base. Neighboring trees that showed no signs of decay in the increment core were used as healthy controls. At the Haukeli sites we also sampled three living aspen trees (*Populus tremula*) with heartrot caused by the white-rot fungus *Phellinus tremulae*.

2.2. Sample Trees and Gas Collection

Tree stems were drilled horizontally at stump level and normally also at breast height with a ten-millimeter-diameter borer, so that the bore hole extended into the pith. For some trees with decay additional drillings were made up to 3.5 m above stump level. Stainless steel tubes, 100–150 mm long with 10.5 mm outer diameter were forced into the holes with a hammer and stopped with butyl rubber septa. Gas samples were taken by a 20 mL polypropylene syringe equipped with a three-way valve, normally every week in 2013 and at longer time intervals in 2014. Caution was taken to adjust atmospheric pressure (by slipping the plunger) before detaching the syringe to avoid contamination from air. The samples (15 mL) were transferred to helium-washed, pre-evacuated 12 mL glass vials

crimped with butyl rubber septa resulting in an overpressure in the vials to avoid contamination during sample storage.

2.3. Gas Analysis

The gas samples were analyzed for O₂, CO₂, N₂O and CH₄ by gas chromatography (GC, Model 7890, Agilant, Santa Clara, CA, USA) with back flushing and helium (He) as a carrier gas, equipped with a 30 m wide-bore Poraplot Q (0.53 mm) column (38 °C) for separation of CO₂, N₂O and CH₄ from air and a 60 m 5Å molsieve capillary column (25 °C) for separation of Ar, O₂ and N₂. The analytical columns were connected to a thermal conductivity detector (TCD), a flame ionization detector (FID) and an electron capture detector (ECD) for quantification of CO₂ and O₂, CH₄ and N₂O, respectively. The ECD was run at 375 °C with 17 mL·min⁻¹ argon/methane [Ar/CH₄] [90/10 vol.%] as make-up gas. The GC was connected to an auto-sampler via a peristaltic pump (Gilson minipuls 3, Middleton, WI, USA) pumping approximately 1.2 mL gas at 1 atmosphere into two 250 μL sampling loops dedicated to each column. Certified standard mixtures of CO₂, N₂O and CH₄ in Helium (AGA, Norway) were used for calibration.

2.4. CH₄ and CO₂ Production Potentials

Sample tree #6 (Ås), shown to have high stem wood CH₄ concentrations in field measurements and appreciable signs of stem base infection by both *Heterobasidion annosum* s.l. and *Armillaria* sp., was felled on 5th November 2015. One ten-centimeter-thick stem disc was cut at stump level and one at breast height so that these discs enclosed the bore holes used for *in situ* profiling of gas concentrations. After visual inspection, three bark-to-heartwood sectors were chosen on each disk for sub-sampling, one of the sectors was taken from the position used for *in situ* profiling of gas concentrations. Samples (*ca.* 0.5 × 1.0 × 3.0 cm) were excised with the aid of a band saw from healthy sapwood, a wetwood appearing tissue adjacent to healthy sapwood, a reaction zone tissue formed between healthy sapwood and colonized heartwood, and colonized heartwood (Figure S1). The samples, immediately transferred to sterile 120 mL serum bottles, had a fresh weight of 4 to 12 g (depending on sample moisture content). For each sample, two replicate bottles were prepared, one for anoxic, and one for hypoxic incubation. The bottles were crimp sealed with butyl rubber septa, amended with 1 mL DI water and He-washed (5 cycles of evacuation and He-loading with final pressure equilibration). Initially hypoxic bottles received pure O₂ to a final headspace concentration of 4 vol% immediately after He-washing. Altogether, 52 incubation bottles were prepared within <8 h since felling.

The bottles were kept in a climate cabinet set at 20 °C. Subsamples (1 mL) for analysis of CO₂, O₂ and CH₄ in headspace were taken at hours 40, 140, 300 and 550. Gas concentrations were analyzed, as described above, directly in the headspace by using an automatic incubator set up similar to that described in an earlier study [20]. After incubation, the samples were freeze-dried to determine the dry weight. CO₂ and CH₄ production rates were estimated from increase in concentration over time [ppm h-1] by fitting the product accumulation to a quadratic equation. Rates were recalculated as mg CO₂ and μg CH₄ m⁻³·s⁻¹, using headspace volume, sample dry weight and a wood density of 420 kg·m⁻³ for Norway spruce [21].

2.5. Estimation of Gas Fluxes from Tree Stems

Diffusion of gases in wood can be modelled by Fick' law [22]:

$$M_i = \frac{D12A (C1i - C2i)}{Z} \tag{1}$$

where M_i is moles of gas i diffusing from compartment 1 to 2, D_{12} the coefficient of diffusion for molecules of type i, C_{1i} and C_{2i} the concentrations of type i molecules in compartment 1 and 2, respectively, and Z the distance between compartment 1 and 2. The diffusion coefficient has been determined for various wood samples and species [23], and found to vary by four orders of magnitude, mostly due to differences in water content of wood. The actual D in a living tree is likely to vary even more due to variability in wood density, humidity, and particularly wood degradation, which alters wood structure, density and humidity. It should be, however, possible to develop a simple model based on Fick's law to obtain a rough estimate for D_{12} for any given stem section. This requires measurements of the mixing ratios of CO_2 and O_2 in stems and outside air over time, and validation of the assumption that the respiration consumes one mole of O_2 for each mole of CO_2 produced:

$$P_{CO2} = -P_{O2}$$
 (2)

where P_{CO2} is the production of CO₂ in the stem from time t₁ to t₂.

This production is the sum of the increase or decrease in the mixing ratio in the stem and the number of moles that have diffused in or out of the stem. Thus, we can write:

$$P_{CO2} = dC1iV + \frac{D12At (C'1i - C'2i)}{Z}$$
 (3)

With the assumption that Equation (2) applies, we can write

$$dC1iV + \frac{D12At (C'1i - C'2i)}{Z} = -\left[dC1jV + \frac{D12At (C'1j - C'2j)}{Z}\right]$$
(4)

where the subscript j denotes the jth gas, in this case O_2 , and where V is the volume of the stem section, dC_{1i} is the change in the mixing ratio from time t1 to t2, t is the time period between t2 and t1, C'_{1i} and C'_{2i} are the mean mixing ratios of the gases in compartment 1 and 2 during t, and A and Z are area and distance as explained for Equation (1).

After accounting for the differences in the molecular masses of CO₂ and O₂, their diffusivities in wood should be fairly similar. The diffusivity is related to molecular mass according to Graham's law [24].

$$\delta_{i} = \left(\frac{mi \times ma}{mi + ma}\right)^{-1/2} \tag{5}$$

where m_i is the molar mass of the gas in question and m_a is the molar mass of air, $28.97 \text{ g} \cdot \text{mol}^{-1}$. Since the latter is obviously different inside and outside the tree, we use the mixing ratios for the inside air to calculate m_a .

Inserting δ_i and δ_j into Equation (4) and solving this with respect to D we obtain:

$$D12 = \frac{dC1iV + dC1jV}{\frac{At(C'1i - C'2i)}{Z} - \frac{At(C'1j - C'2j)}{Z}}$$
(6)

If the gas concentrations are converted to moles per cubic meter and the time unit is days, the unit for D_{12} becomes $m^2 \cdot d^{-1}$, which can be converted to $cm^2 \cdot s^{-1}$, the standard unit of diffusivity. After D is

determined in this way, the flux of each gas is calculated according to Equation 1 after taking Graham's law into account.

2.6. Upscaling of CO2 Fluxes from Decaying Stem Heartwood to Stand and Regional Level

To explore the impact of CO₂ fluxes from decay columns in the heartwood of living trees, we first pooled data from the five trees colonized by *H. annosum* s.l., since this decay class dominates in the boreal Norway spruce forests in Europe. Upscaling from the tree/site level to county or the whole-country level requires estimates of the difference in carbon emission (E_c) between decaying and healthy trees, and how the E_c develops with age of the decay. Least square means of the flux data for decay type, site and stem position from July to November 2013 were used to calculate the annual mean flux. Justification for this comes from the fairly symmetrical distribution of monthly mean fluxes [25]. These estimates of the annual mean flux were then plotted against the distances of the sampling points from the ground. For decaying trees a general exponential decline in the flux from the root and upwards was observed, and a simple model was fitted to the least square mean of each fungus and site:

$$E_{ci} = \alpha \exp(-\beta H) + e_i \tag{7}$$

where E_{ci} is the annual flux of CO_2 at the *i*th sampling point, α , and β are parameters to be estimated, H is the distance from root to the *i*th sampling point, and e_i is the residual term. After estimation of the parameters α and β , the difference in E_c between decaying and healthy trees was calculated by inserting the E_c values from breast height of healthy trees into

$$E_{c-healthy} = \alpha \exp(-\beta 1.3) \tag{8}$$

Equation (8) was solved with respect to H to give the height at which $E_{c-rotten} = E_{chealthy}$, which is an estimate of how high in the stem the decay had spread:

$$H_{r} = \log(E_{c-healthy}/\alpha) / -\beta \tag{9}$$

where H_r is the estimated height of the decay column. Thereafter, $E_{c-healthy}$ was subtracted from the predicted $E_{c-rotten}$ values in decaying trees to give estimated surplus CO_2 flux, E_{cnet} , due to decaying wood. An exponential function, Equation (9), was fitted to E_{cnet} with the distance from H_r to each sampling point L_i as independent variable

$$E_{\text{cnetvi}} = \alpha \exp(\beta L_i) + e_i \tag{10}$$

Data from the trees studied at Hokksund were pooled, because all sample heights were not represented in each tree, whereas the tree #7 infected by *H. annosum* s.l. at Ås was analyzed separately.

To calculate the flux for a decaying stem section we first used Behre's stem curve function to find the diameter d_r at the top of the rot column H_r :

$$\frac{dr}{D} = \frac{L}{(a+bL)} \tag{11}$$

where L is the ratio of the distance from the top to H_r and to D, and a and b are coefficients; b was estimated from the height diameter ratio of the tree and a is simply 1 - b (a complete description of the method is given in the supplementary material).

Assuming linear taper from the top of the rot column (H_r) to the root, the tree circumference of any point below H_r can be approximated by:

$$O_i = \pi \left(d_r + t \mathcal{L}_i \right) \tag{12}$$

where O_i is the circumference at point i, d_r is the diameter at H_r in meters, and t is the stem taper in meters per meter, and L_i is the length from H_r to point i.

To find the flux of the whole decaying stem section, Equation (12) is first multiplied with Equation (10).

$$E_s = \pi (d_r + tL_i) \alpha \exp(\beta L_i)$$
 (13)

where E_s is the flux in a small section and the other terms as described above.

Integration of Equation (13) gives the total flux in the whole area of the decaying stem section E_{tots}:

$$E_{\text{tots}} = \int \left[\alpha \pi \exp\left(\beta L_i\right) \left(\beta (d_r + tL_i) + t\right)\right]/\beta^2$$
(14)

The lower limit is at H_r which is $L_r = 0$ and the upper limit is the distance from H_r to L_i or to the ground.

To test how the estimated fluxes conformed to biomass loss from rotten wood, we used the wood density profiles measured from a Norway spruce tree with an advanced stem decay column caused by *Heterobasidion parviporum* [26]. The estimated biomass loss was converted to annual CO₂ flux in kg per square meter by converting the distance between the sampling points to colonization time with the assumption of a constant vertical spread of decay. Huse and Venn [27] reported a mean vertical spread of 18.5 cm per year for Heterobasidion, but we adjusted this to 17 cm per year, as one of their experimental trees showed an exceptional spread value of 70 cm per year. Details of the calculations are given in the supplementary material. Because the density data were given as kg per m³, the flux was multiplied by the volume of the section and thereafter divided by the area to give the annual flux in kg CO₂ per m².

For up scaling, we employed publicly available data from the National Forest Inventory for the eight counties in Eastern Norway (Table S1). The county reports contain tables with information about cutting class, site index, the number of trees per hectare, standing volume, basal area, increment, age, as well as the percentage of wood volume with decay. The decay data were obtained from increment cores that extend from bark to pith at breast height; in presence of decay the volume of the tree would be classified as rotten. The number of sample trees in each stratum can be found by the information on the number of temporary sample plots relative to permanent sample plots and the fact that the sample trees at each temporary plot were sampled with a relascope factor of six. By weighing the various stand variables with the number of sample trees, one can join aggregated data for growth and standing volume with data for decay from the same aggregated strata. The mean age of the decay in the trees can in principle be calculated from the percentages of rotten volume provided that estimates of the annual rate of infection can be obtained. In the county Hedmark there are data for the percent of trees with rot in different diameter classes from two inventories, 1995–1999, and 2000–2004. We used the difference in the un-weighted means of rot frequencies as an estimate of the annual rate of infection, 0.30%. Assuming an equal probability of infection for all diameter classes in a stand, the mean age of the rot is estimated by dividing the percentage of rotten volume by the annual rate of infection.

To calculate the net loss of CO_2 from stem wood due to rot, the current annual volume increments taken from the county inventory reports were converted to gross CO_2 fixation in stem wood, GSP_{CO_2} , by multiplying the assumed average wood density of 420 kg by 0.51 (carbon ratio in the wood), and the resulting 214.2 kg by 44/12 (the ratio of the mole weight of CO_2 to carbon). The net production NPP_{CO_2} was then calculated as:

$$NPP_{CO2} = GSP_{CO2} - RP/100 \times E_{tots}$$

where RP is the percentage of rotten wood volume taken from the county inventory reports.

The ratio of net CO₂ accumulation in stem wood NPP_{CO2} to GSP_{CO2}, the net-to-gross ratio NG_r, was used as a common metrics to assess the impact of rot on CO₂ fixation at the stand level in the various counties and strata.

2.7. Statistical Analysis

For practical reasons a stepwise sampling design was used. The decaying and healthy trees at the Ås site which is close to the laboratory were sampled first; the other sites were sampled later. No trees at the Jevnaker sites showed infection by Armillaria sp. and P. chrysoloma was the only decay fungus present in trees at the Haukeli site. Therefore, to get a more balanced comparison of healthy trees and trees infected with the two most common wood rot fungi in Norway, Armillaria spp. and H. annosum s.l., we used the data from the Ås and Hokksund sites where both fungi were present. All heights were not sampled in all trees, for example the healthy trees were sampled only at stump and breast heights. The sampling design is therefore incomplete and imbalanced and involves repeated measurements of the same trees and heights taken at irregular intervals. A mixed model was used for the analysis of the O₂ and CO₂ mixing ratios and for the CO₂ flux data. Site was considered to be a fixed factor, as the sites were selected based on prior knowledge of edaphic factors that influence decay incidence. Trees within site was considered a random factor, because the trees were selected without any prior knowledge of their gas emissions. A first order autoregressive model was used to model the autocorrelation between repeated measurements from the same sampling point. To test if the CO₂ efflux was dependent on the height of the sampling position, the CO₂ flux data were log transformed and a regression model with distance from the stump height to the sampling point was fitted to the data by taking other effects into account. The Mixed procedure in the SAS SystemTM (SAS Institue, Cary, NC, USA) was used for the analyses, using the default method to compute error degrees of freedom. Normality of the residuals was checked with the normality tests in Univariate procedure in the SAS SystemTM and with quantile-quantile (Q-Q) plots.

3. Results

3.1. Stem CO₂ and CH₄ Concentrations

Both healthy Norway spruce trees and trees with heartwood decay showed clear seasonality in stem concentrations of O_2 and CO_2 . No obvious differences were observed between the sites in the seasonality of stem gas concentrations. The stem O_2 concentrations generally decreased during growing season (April–September) and showed approximately twofold increase during the dormancy period, whereas the stem CO_2 concentrations showed an opposite pattern and started to increase in April/May (Figure 1). The decaying trees showed generally higher O_2 concentrations than the healthy trees; these differences were significant (p < 0.0015) in October and November. In the period between April and October, the decaying trees showed also generally higher CO_2 concentrations than the healthy trees. At stump height these differences were significant in June and July (p < 0.05), and at breast height from August to October (p < 0.01). Differences in the seasonal fluctuation of stem gas concentrations were observed between healthy and decaying trees: in the decaying trees the minimum concentrations of O_2 and maximum concentrations of CO_2 occurred in July/August, whereas in healthy trees these cardinal points occurred in September/October.

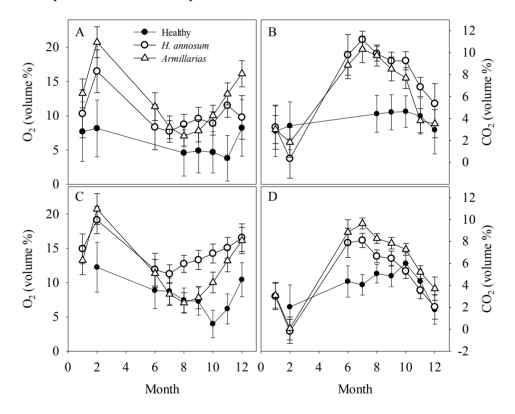


Figure 1. Least square means of monthly O_2 concentrations measured at breast height (**A**), and stump height (**C**), and the corresponding concentrations of CO_2 (**B**) and (**D**) in four healthy trees, six trees infected by *Heterobasidion annosum* s.l. and five trees infected by *Armillaria* species. Error bars are ± 1 standard error. The total number of observations is 510. The data are from two sites, Ås and Hokksund, where mean age of the trees was 110 and 54 years, mean tree height 25.6 and 22.9 m, and mean stem diameter at breast height 38.2 and 26.2 cm, respectively.

For most of the healthy trees and trees with heartwood decay, mean stem concentrations of CH₄ were comparable to ambient air (~2 ppm). Maximum stem concentrations of CH₄ were recorded in trees with *Armillaria* caused heartwood decay, but there was no clear correlation between stem CH₄ and O₂ concentrations as clearly elevated CH₄ levels were observed with O₂ concentrations ranging from 1 to 14 vol.% (Figure 2). Upon tree felling at the end of the field experiment, it was noted that some Norway spruce trees were infected by the white-rot fungus *Phellinus chrysoloma*; the CH₄ levels recorded for these trees were in general similar to those observed in healthy trees and trees infected by *H. annosum* s.l. The trees with heartwood decay could show considerable variation in stem CH₄ concentration between different sampling positions with no sampling-height-specific pattern (data not shown). For healthy trees the maximum stem concentration of CH₄, 17 ppm, was recorded for a tree growing right next to a creek. Irrespective of tree condition, the volumetric concentration of CO₂ in stems was several thousand times higher than that of CH₄ (Table 1).

Table 1. Volumetric ratio of CH₄ to CO₂: mean, standard deviation, minimum and maximum of all field samples analyzed.

Treatment	Number of Gas Samples	CH ₄ /CO ₂	Std	Min	Max
Healthy	94	2.8×10^{-4}	4.51×10^{-4}	3.99×10^{-5}	4.64×10^{-3}
Armillaria sp.	221	2.85×10^{-4}	2.68×10^{-4}	6.39×10^{-10}	9.85×10^{-4}
H. annosum	336	2.07×10^{-4}	4.17×10^{-4}	1.68×10^{-6}	4.44×10^{-3}

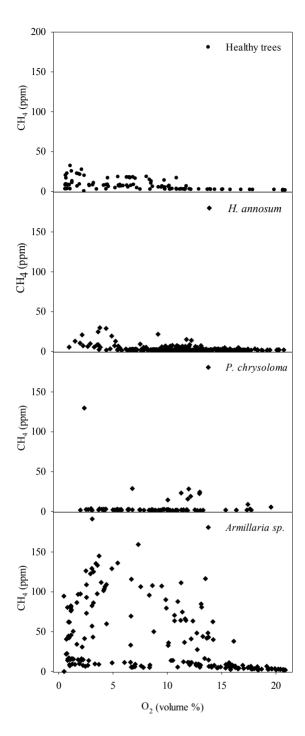


Figure 2. The relationship between *in situ* concentrations of CH₄ and O₂ within healthy stems of Norway spruce and stems with heartwood decay caused by *Heterobasidion annosum* s.l., *Phellinus chrysoloma* and *Armillaria* spp. The dataset involves thirtyseven Norway spruce trees from seven sites in South Eastern Norway, and shows values recorded throughout two growing seasons.

Stem concentrations of CH₄ and CO₂ followed a similar seasonal regulation both in healthy trees and in trees with heartwood decay, as exemplified in Figures 3 and 4. The tree shown in Figure 3 represented an incipient decay caused by *Armillaria* and restricted to stump level, whereas the tree shown in Figure 4 hosted a four-meter-long decay column caused by *Heterobasidion annosum* s.l. and co-occurrence of *Armillaria* at stem base.

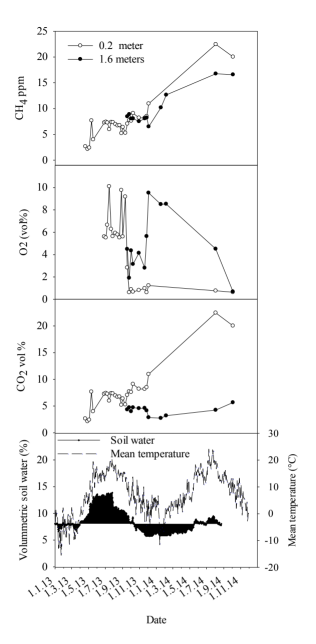


Figure 3. Evolution of *in situ* concentrations of CH₄, O₂, and CO₂ at stump level and at breast height within the stem of Norway spruce tree #2 (age 110 y; breast height diameter 40.6 cm; height 27 m) at the Ås site. The tree was classified as healthy in 2013 but showed incipient infection by *Armillaria* sp. at stump height when it was felled in 2014. The lowermost panel shows mean daily air temperature and volumetric soil water content measured at a soil depth between 10 and 80 cm, obtained from Bioforsk Meterological Service at Ås. At stump level significant correlation (*p* value < 0.01) was observed between CH₄ and O₂ (corr. coefficient −0.54), CO₂ and O₂ (−0.77) and CH₄ and CO₂ (0.7) stem concentrations. At breast height a significant correlation was observed between CO₂ and O₂ (−0.97).

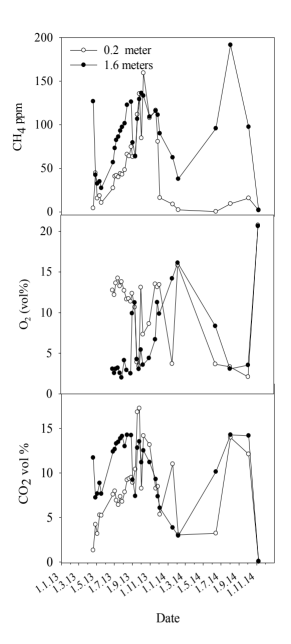


Figure 4. Evolution of *in situ* concentrations of CH₄, O₂, and CO₂ at stump level and at breast height within the stem of Norway spruce tree #6 (age 110 years; breast height diameter 44.5 cm; height 29.1 m) at the Ås site. The tree showed combined infection by *Heterobasidion annosum* s.l. and *Armillaria* sp. with a four-meter-long decay column in stem heartwood. At stump height with relatively advanced decay in the outermost heartwood, a significant correlation (p value < 0.01) was observed between CO₂ and O₂ (corr. Coefficient = -0.73) and CH₄ and CO₂ (0.64) stem concentrations. At breast height with incipient decay a significant correlation was observed between CH₄ and O₂ (-0.65), CO₂ and O₂ (-0.98) and CH₄ and CO₂ (0.66).

3.2. Tissue-Specific CO2 and CH4 Production Potentials at 20 °C

Generally, CO₂ production rate was significantly higher in hypoxic than in anoxic incubations (p = 0.0008) (Figure 5), and significantly higher (p = 0.0001) in tissues dissected from stump level than those dissected from breast height. Regarding CH₄, no statistically significant difference was

observed in gas production rate between the sampling heights and the tissue specific samples from the two heights were pooled together for statistical analysis of treatment specific differences. The healthy sapwood and reaction zone showed significantly higher CH₄ production under initially hypoxic conditions (4 vol.%) than under permanently anoxic ones (respective *p* values 0.0005 and 0.03). Under both anoxic and hypoxic conditions, the production rate of CO₂ was several orders of magnitude higher than that of CH₄ in all the tissue types.

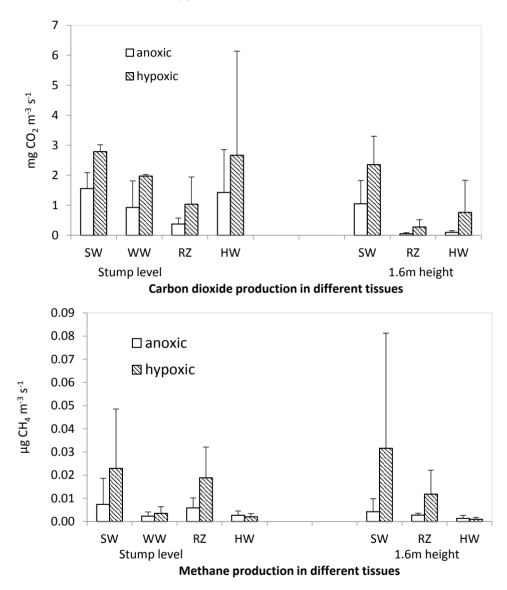


Figure 5. Average production rates of carbon dioxide and methane in the tissue-type samples dissected from two heights from tree #6, and incubated for 23 days at anoxic and hypoxic laboratory conditions. SW—healthy sapwood (stump level, n = 3; breast height, n = 6); WW—wetwood (n = 2); RZ—uncolonized reaction zone (n = 3); HW—colonized heartwood (stump level, n = 5; breast height, n = 3). Error bars represent ±one standard deviation.

3.3. Upscaling of CO₂ Fluxes from Decaying Stem Heartwood to Stand and Regional Level

In trees infected by H. annosum s.l. or Armillaria species, the estimated CO_2 fluxes were higher than in the control trees, and the maximum values, found near the ground level, declined exponentially (p < 0.001) with the distance from the ground level (Figure 6A–D). The highest CO_2 flux observed in a tree with heartwood decay was ten to twenty times higher than in the control trees (Figure 6D). There was also a non-significant tendency of lower CO_2 fluxes at 1.3 meter height than at stump level in the control trees (Figure 6).

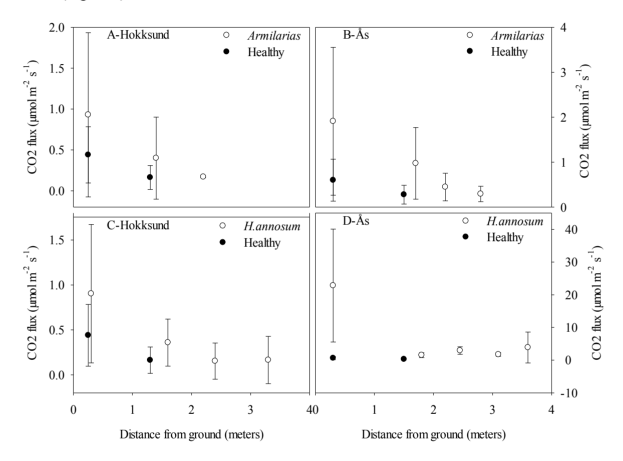


Figure 6. Estimated CO₂ fluxes at different stem heights in healthy trees and trees infected with *Armillaria* species or *H. annosum* s.l. Up to 80 gas samples per category were collected in the period between July and November in 2013. The means are weighted with the length of the sampling period, error bars are ±one standard deviation. At Ås there were two healthy trees, four *Armillaria* infected trees and one *H. annosum* s.l. infected tree. For Hokksund, the corresponding tree numbers were two, one and five, respectively.

NPP_{CO2} was only slightly influenced by decay in stands with low rot percentage (Figure 7), irrespective of the estimation method. When the stand rot percentage was above 15 percent, the net to gross ratio, NG_r, declined sharply with the flux based method both with parameter estimates from the five trees at Hokksund and with the one tree at Ås. The NG_r estimates based on the wood density loss in one tree were little influenced by increasing stand rot percentage. The NG_r values shown in Figure 7 are from samples where the mean length of the stem decay column is below six meters in order to keep this parameter close to the limits of the decay column heights observed in the trees used for the flux

based model. Only in one sample from Telemark (standcutting class 3, site class 2) was the estimated mean decay column height larger, 7.3 meters and the stand rot percentage 28 percent. In this sample the flux based method produced NG_r of about -2.5, implying that the stand had a net loss of CO₂. The density based estimate in the same sample produced a modest NG_r reduction from 1 to 0.96.

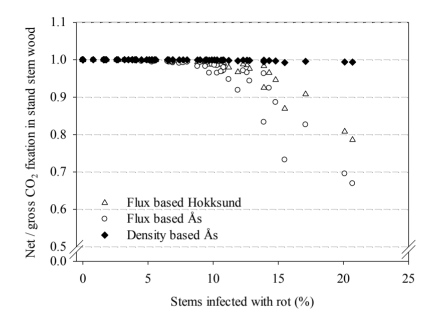


Figure 7. Ratio of net-to-gross CO₂ fixation per hectare in the stem wood at *Heterobasidion* infested Norway spruce stands, as estimated from CO₂ fluxes and rot prevalence in tree stems; see text for explanation of the method. The flux based estimates are from the five *Heterobasidion* infected trees monitored at Hokksund and the one monitored at Ås. Wood density estimates are based on stem section data of one Norway spruce tree that showed an extensive stem decay column caused by *Heterobasidion parviporum* in our previous study [26] (tree #9).

4. Discussion

As a result of CO₂ released by respiring living cells and barriers to diffusion of gases in bark and xylem, the concentration of CO₂ in tree stems is strongly elevated above ambient [28]. The stem concentrations of CO₂ and the seasonal dynamics now recorded in healthy trees are in line with previous studies for Norway spruce [29] and other tree species [28]. For trees infected by *Heterobasidion annosum* s.l. and *Armillaria* species, the most common white-rot fungi associated with heartwood of Norway spruce in boreal forests, the stem concentrations of CO₂ in spring and summer were generally two times higher than those observed for trees without any decay in heartwood. To the best of our knowledge, this is the first report about seasonal fluctuation in gas concentrations in stems of living trees affected by decay. The maximum stem concentrations of CO₂ now recorded for decaying trees, >20%, are two-fold higher than those observed in summertime for saprotrophic fungal species associated with wood logs and stumps [30].

While the source of the stem gases was not identified by isotope analysis in our study, most of the surplus of CO₂ in stems of trees with heartwood decay obviously resulted from microbial respiration and mineralization of wood; white-rot fungi are the only organisms that are able to efficiently degrade

all structural components of wood, the recalcitrant lignin included [31]. However, during the early phase of host/pathogen interaction, when an antifungal polyphenol-rich host defense tissue forms at the interface between sapwood and pathogen colonized heartwood [32], part of the excess CO₂ could also be derived from increased use of tree's energy reserves to build and reinforce this reaction zone. Another point worth noting is the seasonally earlier peaking of stem CO₂ concentrations in the decaying trees in comparison to healthy trees. Considering that heartwood is probably thermally insulated by bark and sapwood, and the fact that the optimal growth temperature of *Heterobasidion* species and many other heartwood associated decay fungi is above 20 °C [13], we presume this difference is due to peaking of fungal respiration during the warmest time of the summer.

When considering the impact of heartrot in living trees on carbon sequestration and emission at tree and stand level, several factors need to be taken into account. First, the growth of the tree may be reduced as a result of root decay and reduction of the conductive sapwood in stem due to extensive heartrot, these eventually resulting in premature death of the tree. Secondly, the decay caused by white-rot fungi may not advance at a linear rate; the elevated O₂ levels observed at stump height in many of the trees with heartwood decay, originating either from increased diffusion of air into xylem due to breakage of decayed main roots by wind rocking and/or transport of oxygen to decomposing heartwood from soil by hyphae or rhizomorphs, can be expected to accelerate degradation of lignin, the rate limiting step during incipient wood decay. Finally, there is uncertainty about the efflux of CO₂ to the atmosphere, as part of the CO₂ formed within stems and branches can be fixed by bark or needle photosynthesis [28].

Our CO₂ flux estimates amounted to about 70 percent of the values given in a recent study [25] for healthy trees during the month July to November. There was large temporal variation in the estimates of radial CO₂ diffusion coefficients in the estimated fluxes, as shown by the large standard deviations. However the values of the coefficients (Table S2) were generally within the range that has been determined experimentally [23]. Experimental determination showed that the diffusion coefficients in the radial direction vary by four orders of magnitude mostly in response to wood water content. Thus, large variability in the fluxes from trees *in vivo* is to be expected. The fairly close agreement with the data from a recent Norway spruce study [25] suggest, that the method developed here could be suitable for estimating gas fluxes from stem mixing ratios.

It is noteworthy that the maximum decay frequency of 20% used in our models (Figure 7) corresponds to the observed average decay frequency caused by *H. annosum* s.l. in managed Norway spruce forests by the time of final harvest, and that in individual Norway spruce stands the decay frequency can be above 50% [14,15]. The up-scaling of CO₂ efflux from *Heterobasidion* caused decay to the stand level based on aggregated data for decay frequency and annual volume growth from national forest inventories suggests that root and heartwood decay in living Norway spruce trees can cause a substantial loss of carbon. In stands with low annual growth and high decay frequency the carbon loss due to mineralization of lignocellulose to CO₂ could be sufficient to turn the trees from a carbon sink to a source. There are many uncertainties with the flux estimates and the assumptions about the infection and spread rates. For practical reasons, we could not start sampling at the same time at all sites, and some additional trees and heights were also included as the sampling progressed. To minimize the potentially confounding effects of differential sampling on the flux calculations, we chose to use samples only from the sites Ås and Hokksund and the months July to November in 2013.

This half-year period is obviously a very limited time to make strong inferences about biological processes that can vary with temperature, humidity and other factors. The results from the flux based and the density based up-scaling differed considerably; the former suggested that wood decay has a large impact on net CO₂ fixation, while the latter suggested a minimal effect. The difference could be due to systematic errors in both methods or due to real variability between sites, years, and trees. Regarding the latter, Huse and Venn [27] observed more than ten-fold difference in vertical spread of *H. annosum* between different trees, and thus large variation in the biomass breakdown rate and CO₂ fluxes between trees is quite possible. Therefore the estimates presented in Figure 7 serve above all as an illustration of a potential effect. It is worth noting that both the CO₂ flux based estimates and the wood density based estimates decline exponentially with the distance away from stem base where the infection is oldest, this implying that biomass loss accelerates with the duration of the infection. However, when most of the substrate in the heartwood has been consumed, the CO₂ flux should eventually decline. Consequently, to obtain reliable carbon budgets for young and mature forest stands, the frequency, type and stage of decay in living trees need to be taken into account.

Regarding methane, healthy Norway spruce and trees with heartwood decay caused by Heterobasidion annosum s.l. showed comparable stem concentrations of CH₄, around 30 ppm at maximum. Contrastingly, trees infected by Armillaria species showed frequently 3-4 times as high CH₄ levels. We are not aware of any earlier reports about stem concentrations of CH₄ in Norway spruce or any other tree species infected by *Heterobasidion* and *Armillaria* spp., fungi able to infect a wide range of conifer and broadleaved trees. As proposed in earlier studies, under microaerobic or anaerobic conditions, intermediate degradation products from fungal degradation of lignocellulose may provide substrates for bacteria and archaea [9,10], which in turn can accelerate fungal growth by removing their metabolic waste products and by enriching the substrate through N-fixation [11]. The apparent lack of correlation between stem CH₄ and O₂ concentrations in trees with elevated CH₄ levels is noteworthy, and is probably due to the utilized point measurements, which profile the composition of gas diffusion stream that includes also the CH4 produced at anaerobic microsites. When comparing the differences in ecological range and behavior between Armillaria and Heterobasidion species, Shaw [33] stressed that *Armillaria* species are frequently associated with water-logged trees, have the ability to transport oxygen in the air channels of rhizomorphs that are commonly formed in decayed heartwood, and that their mycelial growth is highly stimulated by ethanol, an anaerobic microbial but also plant fermentation product. These features may suggest that Armillaria species are better adapted to grow and feed under anaerobic conditions than *Heterobasidion* species. The potential interplay between anaerobic bacteria and archaea, and Armillaria species at anaerobic/aerobic sub-surfaces of wood tissues upon lignocellulose decomposition remains to be clarified.

Similarly to the low mixing ratios of CH₄ and CO₂ recorded within Norway spruce stems, a low production rate of CH₄ in comparison to that of CO₂ was observed in the laboratory experiment, where selected tissue types were incubated under anoxic or hypoxic conditions. At an upland site in temperate forest, mean stem concentrations of CH₄ in three broadleaved trees (red maple, red oak, black birch) during the growing season were above 15,000 ppm, whereas the conifer *Pinus strobus* showed only negligible stem concentrations of CH₄ and a mean production rate of 0.067 µg CH₄ m⁻³·s⁻¹ under laboratory incubation [34], a level essentially similar to that observed for healthy Norway spruce wood in the present study. Covey and coworkers [34] proposed that the high CH₄

levels in broadleaved trees originated from methanogenesis by archaea within the tree stem and that microbial infection of living trees might result in substantial biogenic production of CH₄ also in upland forests. In the study of Mukhin and Voronin [35] substantially higher production rate of CH₄ from decaying birch and pine wood than those recorded for Norway spruce in the laboratory experiment of the current study, which they attributed to progressive wood degradation by decay fungi and anaerobic microbes. In contrast to these studies, our data does not indicate that Norway spruce, when growing on well-drained stands and whether healthy or showing heartwood decay caused by white-rot fungi, plays any significant role in biogenic *in situ* production of CH₄, or in its transport from a biogenic source in soil. The reasons for the large variation in stem CH₄ concentrations between different tree species growing on well-drained soils remain to be clarified but this variation may relate to tree-species specific differences in the amount of xylem-associated anaerobic microsites that support methanogenic archaea.

Regarding the pattern of CH₄ production among different wood tissue types determined in the laboratory experiment, healthy sapwood and reaction zone produced clearly more CH₄ under hypoxic conditions than anoxic ones. The observed anoxic production of CH₄ in these tissues does not preclude a biogenic origin but the elevated CH₄ production rate under hypoxic conditions rather points towards production through a chemical mechanism. Several studies [36–38] have shown production of CH₄ from plants under stress and provided evidence that it could be derived from action of free radicals on plant cell wall components such as pectin. A similar mechanism could account for the now recorded methane production from healthy sapwood. One could envisage that CH₄ generation from trees through such a mechanism could be the highest in timber logged during spring, summer and autumn when the wood is not frozen. However, if the magnitude of CH₄ production in logged timber is at a similar level as now observed in the laboratory experiment, the production rate of CH₄ through a biochemical mechanism in timber of Norway spruce can also be considered as insignificant.

5. Conclusions

Our motivation to carry out this study was the prediction that emission of CO₂ and possibly also CH₄ from decaying heartwood in living Norway spruce trees could amount to regionally significant levels, especially when considering that on average every 4th tree is affected by root and stem decay in most mature stands of Norway spruce [14]. Due to fungal respiration, the recorded stem CO₂ concentrations could be over two-fold higher in trees with heartwood decay in comparison to healthy trees. In contrast to CO₂, the low stem concentrations of CH₄ recorded in both healthy and decaying trees can be considered as insignificant. The performed up-scaling of CO₂ efflux measurements from six Norway spruce trees with heartwood based decay columns caused by *Heterobasidion* annosum s.l. suggests that the balance between carbon sequestration and emission could be significantly influenced at stands with high frequency of advanced root and stem heartwood decay. However, large scale studies with more intensive measuring of CO₂ fluxes from decaying heartwood are warranted to obtain robust estimates about the impact of fungal mineralization of wood on carbon budgets in Norway spruce stands. Our hope is that the pilot up-scaling exercise carried out now will stimulate further studies on this poorly explored topic.

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Author Contributions

Harald Kvaalen, Ari Hietala and Halvor Solheim came up with the original research idea and prepared the grant application that enabled the study. The research was designed by all authors. Harald Kvaalen and Halvor Solheim contacted forest owners. In addition to persons named in the acknowledgments, gas samples from living trees were collected by Halvor Solheim and Harald Kvaalen. Collection of samples for the laboratory incubation experiment was carried out by Harald Kvaalen and Ari Hietala. Laboratory analyses were conducted by Peter Dörsch and data analyses by Peter Dörsch and Harald Kvaalen. The manuscript was written by Ari Hietala and Harald Kvaalen, and supported by the other two authors.

Conflict of Interest

The authors declare no conflict of interest.

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