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# Inter- and intra-annual dynamics of photosynthesis differ between forest floor vegetation and tree canopy in a subarctic Scots pine stand

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#### ABSTRACT

We studied the inter- and intra-annual dynamics of the photosynthesis of forest floor vegetation and tree canopy in a subarctic Scots pine stand at the northern timberline in Finland. We tackled the issue using three different approaches: 1) measuring carbon dioxide exchange above and below canopy with the eddy covariance technique, 2) modelling the photosynthesis of the tree canopy based on shoot chamber measurements, and 3) upscaling the forest floor photosynthesis using biomass estimates and available information on the annual cycle of photosynthetic capacity of those species. The studied ecosystem was generally a weak sink of carbon but the sink strength showed notable year-to-year variation. Total ecosystem respiration and photosynthesis indicated a clear temperature limitation for the carbon exchange. However, the increase in photosynthetic production was steeper than the increase in respiration with temperature, indicating that warm temperatures increase the sink strength and do not stimulate the total ecosystem respiration as much in the 4-year window studied. The interannual variation in the photosynthesis of the tree canopy seemed to be more stable from the forest floor vegetation, whereas the photosynthesis of the tree canopy seemed to be more stable from year to year. Tree canopy photosynthesis increased earlier in the spring, whereas that of the forest floor increased after snowmelt, highlighting that models for photosynthesis in the northern area should also include snow cover in order to accurately estimate the seasonal dynamics of photosynthesis in these forests.

## 1. Introduction

Northern forests experience cold and snowy winters and a short and cold growing season. Although the carbon uptake rates are smaller than in more southern ecosystems, boreal forests have been traditionally considered to be carbon sinks due to very slow decomposition rates (Fan et al., 1998), but the magnitude of, and the factors that control, this sink are still not accurately quantified. This vulnerable region is predicted to experience extensive climate change, which will influence the carbon exchange between the ecosystem and the atmosphere. Studies have suggested that the photosynthetic uptake, i.e. the primary production of these northern forests, will increase (Myneni et al., 1997; Qian et al., 2010; Ueyama et al., 2015). At the same time, it is known that decomposition is temperature dependent (Davidson and Janssens, 2006) and that increases in plant productivity might increase the soil

organic matter decomposition rates (Hartley et al., 2012; Parker et al., 2015). Thus, there is a concern that increased temperature accelerates decomposition and turns these soils from a carbon sink to a source (Crowther et al., 2016).

The typical vegetation structure of northern forests consists of tree canopy and a vegetated and photosynthetically active forest floor. The ground is colonized by a dense community of plant species consisting mainly of ericaceous shrubs, mosses and lichens, whereas relatively sparse canopies above enable notable primary production of the forest floor vegetation (Goulden and Crill, 1997; Moren and Lindroth, 2000; Kulmala, 2011). The proportion of forest floor vegetation in the momentary carbon dioxide (CO<sub>2</sub>) uptake of a boreal forest ecosystem has been reported to vary between 3% and 61% (Goulden and Crill, 1997; Subke and Tenhunen, 2004; Ikawa et al., 2015), but the proportion naturally depends on the site, climate and vegetation characteristics.

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Net ecosystem  $CO_2$  exchange is commonly measured by the eddy covariance technique and further partitioned into total ecosystem respiration and photosynthesis, i.e. gross primary production (GPP). The attained total ecosystem photosynthesis consists of the uptake by both the tree canopy and the forest floor vegetation. However, their photosynthetic efficiency and seasonal dynamics differ especially due to their different growth patterns but also due to snow cover that usually continues late into spring, when increased air temperature already enables the photosynthesis of the tree canopy above the snow-covered forest floor. Thus, GPP models which ignore the snow cover might fail to accurately estimate the seasonal dynamics of photosynthesis in northern forests with a sparse tree canopy.

The annual growth is considered to be source limited in many growth models, meaning that the more there is photosynthesis, the more there is growth. However, the growth of northern boreal trees is reported to be temperature, i.e. sink, limited (Körner, 2003), meaning that tree growth in high latitudes is regulated by temperature, which controls the activity of growing meristems, while photosynthetic production together with carbon storage are always sufficient to maintain growth. Several studies have shown that the radial growth, for example, of the northernmost conifers is promoted by warm temperatures during the growing season (Korpela et al., 2011; Seo et al., 2011; Babst et al., 2012; Henttonen et al., 2014; Xu et al., 2014). On the other hand, connections between the growth and GPP have been reported on an annual basis (Berninger et al., 2004; Gea-Izquierdo et al., 2014; Schiestl-Aalto et al., 2015; Kulmala et al., 2016) and on shorter timescales (Chan et al., 2016; Kulmala et al., 2017). However, the research has focused on trees, whereas such studies on sink and source limitation in forest floor vegetation are still missing.

The aim of this study was to explore the intra- and interannual dynamics of forest floor photosynthesis in comparison with those of the tree canopy in a subarctic forest stand. Furthermore, we were interested to see whether the growth of forest floor vegetation is source limited. In addition, we aimed to determine the temperature dependence of the photosynthetic production and ecosystem respiration in order to estimate the effect of rising temperatures on those.

To study these questions, we measured  $CO_2$  fluxes at different scales in a subarctic Scots pine (*Pinus sylvestris* L.) stand at the northern timberline with the eddy covariance technique both above and below the forest canopy and at pine shoot level with shoot chambers. In addition, we measured the variation in the annual height increment of dwarf shrubs on the forest floor. The turbulence needed for eddy covariance measurements is often insufficient below a forest canopy and thus we have also used independent top-down and bottom-up approaches to determine the momentary forest floor photosynthesis. In practice, we downscaled the flux as the difference between the total ecosystem and estimated canopy photosynthesis (top-down), and we upscaled it using the forest floor biomass and available models for leaf-mass based photosynthesis (bottom-up).

# 2. Materials and methods

#### 2.1. Site

We studied the subarctic Scots pine carbon dynamics over 4 years, between 2012 and 2015, at the Värriö Subarctic Research Station (67° 46′ N, 29° 35′ E), which is located below the northern altitudinal treeline in north-eastern Finnish Lapland. The mean annual temperature was -0.5 °C and the mean annual precipitation 601 mm for the years 1971–2000 (Pirinen et al., 2012). Mean monthly temperature was above 0 °C from May to September. July was the warmest month, with a mean temperature of 13.1 °C.

The study site (SMEAR I) was located on the summit plateau of Kotovaara hill (400 m a.s.l.) and was dominated by Scots pine, with a basal area weighted mean tree height of 10 m and a stem diameter of 14.0 cm. The fell was naturally populated with a density of ~750 trees

ha<sup>-1</sup>. The all-sided leaf area index (LAI) was estimated to be  $^{3.2} \text{ m}^{2} \text{ m}^{-2}$  using available biomass equations (Repola, 2009), and the specific leaf area was 10 m<sup>2</sup> kg<sup>-1</sup>. The forest floor vegetation comprised a variety of mosses, lichen and dwarf shrubs such as *Vaccinium myrtillus*, *Vaccinium vitis-idaea* and *Empetrum nigrum*.

### 2.2. Biomass sampling and the growth of understorey vegetation

The biomass of the forest floor vegetation (vascular plants, mosses and lichen) was estimated by collecting 12 samples in late July 2015 using systematic sampling with a frame ( $0.198 \text{ m} \times 0.198 \text{ m}$ ). The samples were divided into different species and weighed separately after drying for 48 h at 60 °C.

Annual height increment of the dwarf shrubs was systematically determined in 2016 using a transect of 45 m length along which we measured the increase in length of the main shoot of the three closest individuals of *V. vitis-idaea*, *V. myrtillus* and *E. nigrum* at 3 m intervals, resulting in 45 individual length increments for each species. The lengths were determined by a digital caliper with an accuracy of 0.01 mm.

All shoot increments were normalized by dividing the increments by the 4-year mean growth of an individual. For example, if a single shoot had grown 15, 20, 10 and 15 mm in the years 2012, 2013, 2014 and 2015, respectively, all increments were divided by their mean, i.e. in this case 15 mm. These relative growths were further statistically analysed. The differences between the years were tested using a one-way analysis of variance (ANOVA) and the Tukey honest significant difference test in R. The difference was considered significant when p < 0.05.

#### 2.3. Meteorological measurements

Relative humidity (RH) and ambient air temperature ( $T_a$ ) at 2 m and 9 m height were measured with MP106 A and PT-100 sensors (Rotronic, Switzerland), respectively, at SMEAR I. The cumulative temperature sum for the growth period was calculated using a 5 °C threshold. Photosynthetically active radiation (PAR) was measured with an LI-190SB Quantum Sensor (LI-COR Biosciences, Lincoln, NE, USA) above the tree canopy. Soil temperature ( $T_s$ ) was measured 10 cm below ground and in humus ( $T_{H_s}$  approx. 2 cm below ground) with a PT-100 sensor. Soil volumetric water content was measured with three ThetaProbe ML2x sensors (Delta-T Devices, UK) located in the uppermost 5 cm. All sensors were located near the eddy covariance tower.

Precipitation (both rain and snowfall) was measured at the Salla Värriötunturi weather station, managed by the Finnish Meteorological Institute and located at the Värriö Subarctic Research Station. Since the forest around SMEAR I is less dense than the area around the research station, the time of snowmelt at SMEAR I was roughly determined from the soil temperature measurements. We assumed that all the snow had melted when the soil temperatures rose above 0.3 °C.

### 2.4. Eddy covariance instrumentation, data processing and partitioning

The net ecosystem exchange (NEE) of  $CO_2$  was estimated using the eddy covariance (EC) technique, employing the LI-7200 Enclosed Path  $CO_2/H_2O$  Analyzer (LI-COR Biosciences) and the METEK USA-1 ultrasonic 3D anemometer (METEK, Elmshorn, Germany) installed at 16.6 m at the top of the eddy flux tower in April 2012.

Net forest floor exchange (NFFE) below the tree canopy was measured at 2.7 m above ground with another eddy covariance system located 25 m away from the main eddy flux tower in 2013. The instrumentation was identical to that above canopy in 2015, but in 2013–2014 an LI-7500 A Open Path  $CO_2/H_2O$  Analyzer (LI-COR Biosciences) was used, employing the same model sonic anemometer. The analyser was upgraded to the LI-7200 Enclosed Path  $CO_2/H_2O$  Analyzer in spring 2015.

Raw 10 Hz data were processed using standard processing steps with EdiRe (R Clement, University of Edinburgh, UK). The processing included despiking (Vickers and Mahrt, 1997), crosswind correction applicable to the METEK sonic anemometer, coordinate rotation using the two-dimensional rotation method (Baldocchi, 1988), sonic virtual temperature correction (Kaimai and Gaynor, 1991), as well as frequency response correction (Massman, 2000) and buoyancy flux correction (Schotanus et al., 1983). We applied the Webb-Pearman---Leuning density correction (Webb et al., 1980) to the data originating from the LI-7500 A. We did not apply the Burba correction (Burba et al., 2006, 2008) as the temperature range experienced appeared to be within the limits shown to have no or very little effect on the correction (Burba et al., 2006). Furthermore, the subcanopy system was rarely exposed to direct sunshine for any prolonged amount of time, nor to temperatures below 0 °C. In addition, we applied lag time and tube attenuation corrections relevant to the LI-7200 Enclosed Path CO2/H2O Analyzer. In order to guarantee reliable and high-quality flux data, quality checks (Foken and Wichura, 1996; Foken et al., 2004) were carried out. We estimated the footprints (i.e. field of view of the EC setup) of both towers from EC flux data for near-neutral atmospheric stability conditions (-0.05 < z/L < 0.05) using the Kormann and Meixner (2001) analytical model. The used thresholds were selected from Geissbühler et al., 2000.

The measured fluxes were corrected for storage change under the measurement height. The storage was calculated from the mean 30-minute CO<sub>2</sub> measurements assuming a constant concentration profile from the measurement height down to the forest floor. Above-canopy fluxes measured under low turbulence were excluded using a friction velocity (u\*) threshold of 0.4 m s<sup>-1</sup> (see later). Below-canopy fluxes were filtered with the standard deviation of vertical wind speed ( $\sigma_w$ ) (see Launiainen et al., 2005). The threshold value was 0.2 m s<sup>-1</sup>. The accepted fluxes were partitioned into GPP ( $P_E^{EC}$  or  $P_{FC}^{FC}$ ) and respiration ( $R_E$  or  $R_{FF}$ ) for the ecosystem (E) and forest floor (FF) using a simple empirical model which was also used for interpolating missing and rejected flux records and analysing the dynamics of CO<sub>2</sub> exchange. The model describes NEE or NFFE as the difference between temperature-driven respiration (R, µmol m<sup>-2</sup> s<sup>-1</sup>) and photosynthesis (P, µmol m<sup>-2</sup> s<sup>-1</sup>) and air temperature ( $T_a$ , °C). Photosynthesis is modelled as follows:

$$P = \frac{\alpha I + P_{max} - \sqrt{(\alpha I + P_{max})^2 - 4\theta I P_{max}}}{2\theta} f(T)$$
(1)

where  $\alpha$ , *Pmax* and  $\theta$  are parameters. Due to the open canopy and missing forest floor PAR measurements, we used the above-canopy measurements. f(T) is the instantaneous temperature response that brings *P* to zero at freezing temperatures (Kolari et al., 2014) as follows:

$$f(T) = -\frac{1}{1 + e^{(2(T_0 - T_a))}}$$
(2)

where  $T_0$  is the inflection point. Respiration was estimated using an exponential temperature function as follows:

$$R = R_C Q_{10}^{T_{Sa}/10} \tag{3}$$

where  $R_C$  and  $Q_{10}$  are parameters. The driving temperature  $T_{sa}$  was the mean of the air temperature at 9 m height and soil temperature at 10 cm depth.  $\alpha$ , *Pmax* and  $R_c$  were estimated for time periods of 11 days, whereas  $Q_{10}$  and  $\theta$  were estimated over the whole study period. The obtained values were  $Q_{10} = 2.2$  and  $\theta = 0.75$  for the above-canopy data. For the below-canopy data, the obtained values were  $Q_{10} = 1.8$  and  $\theta = 0.75$ .

There was a notable difference in the flux partitioning procedure compared with more southern sites due to the polar day (24 h of daylight): flux records taken in darkness (night-time) were missing in the summer and thus respiration parameters could not be estimated directly from the measured night-time fluxes. Instead, summertime respiration was estimated from the intercept of the regression between NEE and light (see the analysis of this method in Lasslop et al., 2010). Consequently, determining the turbulence filtering criteria (u<sup>\*</sup> and  $\sigma_w$  thresholds) could not be based directly on measured night-time fluxes either. Instead, we performed several model parameterization runs with different u<sup>\*</sup> and  $\sigma_w$  threshold values and determined the final ones as the lowest u<sup>\*</sup> or  $\sigma_w$  thresholds where the estimated  $R_c$  was > 98% of the maximum of  $R_c$  vs u<sup>\*</sup> or  $\sigma_w$  regression.

When the turbulence criteria were met,  $P_E^{EC}$  and  $P_{FF}^{EC}$  were calculated as the difference between  $R_E^{EC}$  or  $R_{FF}^{EC}$  and measured NEE or NFFE, respectively. Missing or rejected NEE (or NFFE) values were gap-filled as the difference between modelled  $R_E^{EC}$  (or  $R_{FF}^{EC}$ ) and  $P_E^{EC}$  (or  $P_{FC}^{EC}$ ).

In order to estimate the uncertainty involved in the gap-filling, we calculated the effect of the used temperature and  $Q_{10}$  estimation on the subcanopy fluxes in May–August 2015 using three different  $Q_{10}$  values (1.5, 2, 2.5) and two different temperatures ( $T_{sa}$  as in Eq. 3 and topsoil temperature at 2 cm depth).

#### 2.5. Carbon dioxide exchange of Scots pine

Gas exchange of Scots pine shoots was automatically measured using four dynamic cylindrical chambers made of acrylic plastic with  $3.5 \text{ dm}^3$  volume. The measured shoots were one year old and located at the top of the canopy. The shoots were debudded prior to installation, and thus further shoot elongation inside the chambers was inhibited.

The chambers were open most of the time but one by one they closed for one minute  $^{180}$  times a day. CO<sub>2</sub> and water vapour concentrations together with air temperature inside the chambers and PAR outside the chambers were recorded every 10 s during a closure. We calculated the CO<sub>2</sub> exchange rate from the change in the CO<sub>2</sub> concentration in the chamber during a closure. More details on the chamber measurements are available in Hari et al. (1999).

We fitted the optimal stomatal control model (Hari and Mäkelä, 2003; Kolari et al., 2007) to the daily measurements to achieve parameters describing the saturation and initial slope of the light response curve, the temperature response of respiration, and the cost of transpiration. We took a running mean of these parameters over 3 days and used the mean as an input for the Stand Photosynthesis Program (Mäkelä et al., 2006) together with 30-minute averages of measured CO<sub>2</sub> concentration, PAR, air humidity and air temperature to estimate the photosynthesis in the entire canopy,  $P_C^{SPP}$ . Instead of upscaling the shoot chamber measurements as such, the model included also tree characteristics and light attenuation in the canopy. Development of a new needle cohort during the summer and shedding of the oldest cohort in the autumn were also described in the model. The leaf area inside the cuvette has little year-to-year variation and thus we normalized the rates of photosynthesis with the ratio between the annual mean of the photosynthetic capacity ( $\beta$ ) and the mean  $\beta$  value in 2015. The ratios were 0.73, 0.76, 1.05 and 1 for the years 2012, 2013, 2014 and 2015, respectively.

The downscaled GPP of the forest floor,  $P_{FF}^{Down}$ , in 2012–2015 was derived as the difference between  $P_{E}^{EC}$  and  $P_{C}^{SPP}$ :

$$P_{FF}^{Down} = P_E^{EC} - P_C^{SPP}.$$
(4)

#### 2.6. Chamber measurements of forest floor carbon dioxide emissions

 $R_{\rm E}$  includes forest floor CO<sub>2</sub> respiration ( $R_{FF}$ ) and above-ground plant respiration, and thus  $R_{FF}$  is expected to be theoretically smaller than  $R_{E}$ . However, several studies have reported lower  $R_{E}$  than  $R_{FF}$  at different timescales (Wang et al., 2010; Speckman et al., 2015; Barba et al., 2018). Therefore, we tested the overall level of  $R_{E}$  by measuring  $R_{FF}$  twice a month in 2013–2015 in 12 locations using a manual chamber (20 cm in diameter and 25 cm high) and permanently installed collars. These plastic collars (Ø 20 cm) were inserted at a depth of ~3–5 cm into the humus layer around the SMEAR I station in 2012. The chamber was equipped with a small fan and a GMP343 Carbon Dioxide

#### Table 1

Species-specific parameters for Eqs 5–7 for a 120-year-old Scots pine dominated forest as in Kulmala et al. (2011a).  $P_0$  is the reported maximum  $P_{max}$ .

	$\tau$ (h)	$P_0 \ (\mu mol \ g^{-1} \ s^{-1})$	$b \ (\mu mol \ m^{-2} \ s^{-1})$
Vaccinium vitis-idaea	72	0.021	100
Vaccinium myrtillus	150	0.365	107
Calluna vulgaris	120	0.073	200

Probe (Vaisala, Finland) that measured the  $CO_2$  concentration every 5 s. The  $CO_2$  emissions were estimated by a linear ordinary least squares regression of  $CO_2$  concentrations against time during 3.5 min. The  $CO_2$ readings were corrected for temperature and humidity using an HMP75 temperature and humidity probe (Vaisala, Finland). The chamber measurements were performed before noon on rainless days both in ambient light conditions and in darkness when the chamber was enclosed within aluminium foil. The chamber and the flux calculations are described in more detail in Pumpanen et al. (2015).

#### 2.7. Upscaling forest floor vegetation photosynthesis

We modelled species-specific photosynthesis of the vascular forest floor vegetation by estimating leaf-mass based rates of photosynthesis  $(\bar{P}, \mu \text{mol g}^{-1} \text{ s}^{-1})$  for each species based on a simple light response curve (Kulmala et al., 2011a) as follows:

$$\bar{P}(t) = \frac{P_{max}(t)I(t)}{b+I(t)}$$
(5)

where I(t) is mean PAR (µmol m<sup>-2</sup> s<sup>-1</sup>) above the canopy at hour *t* and *b* is a species-specific parameter (Table 1). *P*<sub>max</sub> follows changes in the environment and was modelled as follows:

$$P_{max}(t) = f_1(t)f_2(t)f_3(t)P_0$$
(6)

where  $P_0$  is the maximum  $P_{max}$  in a 120-year-old forest stand dominated by *P. sylvestris* (Table 1). The function  $f_1$  describes the effect of speciesspecific temperature history, *S*, i.e. the state of development (Pelkonen and Hari, 1980; Mäkelä et al., 2004; Kolari et al., 2006). It follows temperature with a species-specific time constant  $\tau$  (Table 1) as follows:

$$\frac{dS}{dt} = \frac{T(t) - S(t)}{\tau}$$
(7)

Kulmala et al. (2011a) used hourly average air temperature as *T*, but here we decided to use the hourly mean of  $T_a$  and  $T_s$  since the air temperatures tend to increase earlier than the snow melts. The initial value of *S* was set to -1 °C, and  $f_I$  was calculated as follows:

$$f_1(t) = \frac{S(t)}{S(t_{st})} \tag{8}$$

Kulmala et al. (2011a) fitted the model based on one measurement on day  $t_{st}$  but here we assume  $S(t_{st}) = 15$  °C, i.e.  $f_1$  reaches a maximum with S(t) = 15 °C. If *S* was smaller than 0 °C,  $f_1$  was set to zero.

The function  $f_2$  hinders photosynthesis in low volumetric soil moisture ( $\phi$ ) conditions as follows:

$$f_{2}(t) = \begin{cases} 1 & \text{if } \phi(t) \ge \alpha \\ \phi(t)/\alpha & \text{if } \phi(t) < \alpha \end{cases}$$
(9)

where  $\alpha$  is a critical value of volumetric soil moisture after which the soil moisture starts to hinder photosynthesis. We used  $\alpha = 0.1 \text{ m}^3 \text{ m}^{-3}$  as in Kulmala et al. (2011a).

The function  $f_3$  takes into account the carry-over effect from nighttime frost (Vesala et al., 2010). It is assigned a value of 1 if the minimum air temperature in the previous 24 h ( $T_{min}$ ) was above zero. The  $f_3$  value decreases with  $T_{min}$  values below 0 °C, reaching zero at -10 °C as follows:

$$f_{3}(t) = \begin{cases} 1 & \text{if } T_{min}(t) \ge 0 \,^{\circ}C \\ \frac{T_{min}(t)}{10} + 1 & \text{if } -10 \,^{\circ}C \le T_{min}(t) < 0 \,^{\circ}C \\ 0 & \text{if } T_{min} < -10 \,^{\circ}C \end{cases}$$
(10)

The species-specific  $\bar{P}$  values were upscaled to photosynthesis of the forest floor by multiplying the leaf-mass based values by mean leaf biomass,  $m_{i}$ , and summing all species together as follows:

$$_{FF}^{Up}(t) = \sum_{i=1}^{5} \bar{P}_i(t) m_i.$$
(11)

The most abundant vascular plants were V. vitis-idaea, V. myrtillus, Vaccinium uliginosum, Calluna vulgaris and E. nigrum. Since there were no species-specific parameters available for V. uliginosum and E. nigrum, we assumed that those were similar to V. myrtillus and C. vulgaris, respectively. Since our biomass sampling did not separate leaves and stem, we instead used the mean proportion of leaves of the total biomass reported by Kulmala et al. (2011b): 0.62, 0.25 and 0.29 for V. vitis-idaea, V. myrtillus and C. vulgaris, respectively. The proportions of V. uliginosum and E. nigrum were assumed to be similar to V. myrtillus and C. vulgaris, respectively. Mosses and lichens are not included in  $P_{FF}^{Up}$ .

Using the upscaled rates of forest floor photosynthesis, we derived a second estimate for the photosynthesis of the tree canopy,  $P_C^{EC-FF}$ , as follows:

$$P_C^{EC-FF} = P_E^{EC} - P_{FF}^{Up} \tag{12}$$

#### 3. Results

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#### 3.1. Weather characteristics

Weather conditions differed greatly during 2012–2015 (Fig. 1ABC, Table 2), allowing interannual comparisons of the ecosystem exchange responses in a wide range of climatic conditions. Growing season 2013 was the warmest, sunniest and driest, with the highest mean temperature and PAR and the lowest precipitation and mean RH. Growing season 2015, on the other hand, was the coldest and moistest, recording the lowest temperature and PAR together with the highest precipitation and mean RH. The other years fell somewhere between these two extremes: 2012 was a cold year and 2014 a warm year, with both recording intermediate mean PAR, RH and precipitation. The winters in 2012 and 2013 were colder than normal, whereas the temperatures in the following winters were typical. The temperature sum ranged from 576 °C in 2015 to 974 °C in 2013.

In all years, daily average temperature occasionally dropped below 5 °C during the growing season. The last night-time frost occurred in late May in all years except 2014, which experienced a cool spell around midsummer, with five nights with freezing temperatures during a period of 10 days.

#### 3.2. Biomass of forest floor vegetation

The most common vascular species of the forest floor vegetation were *V. myrtillus, E. nigrum, V. vitis-idaea, V. uliginosum* and *C. vulgaris,* comprising 27% of the above-ground biomass of forest floor species (Table S1, see Supplementary material). Mosses were the most prevalent group (66%), whereas the proportion of lichens in the total forest floor above-ground biomass was ~6%.

#### 3.3. Footprint analysis

Fig. 2 illustrates that 80% of the estimated footprint distance (flux contribution) was way below 200 m, especially under favourable conditions along the plateau of Kotovaara hill. The majority or maximum source location (Kljun et al., 2005) of the fluxes originated from the close proximity of the SMEAR I eddy flux tower in the south-western



**Fig. 1.** Daily air temperature (A), daily photosynthetically active radiation (PAR) (B), soil moisture (C), net ecosystem exchange (NEE) (D), gross primary production ( $P_E$ ) (E) and total ecosystem respiration  $R_E$  (F) in 2012–2015. For clarity, the daily rates are moving averages over 10 days except for PAR, which is a moving average over 15 days.

#### Table 2

Mean meteoro	logical o	characteristics	over the	years	2012-2015
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	2012	2013	2014	2015
Mean air temperature Jan–Dec (°C) Mean air temperature Mav–Sep (°C)	-0.6 9.2	0.8 11.6	0.4 10.0	0.5 8.5
Temperature sum (°C) $^{1}$	641	974	815	576
Precipitation Jan-Dec (mm)	594	483	601	663
Precipitation Jun-Aug (mm)	192	138	259	252
Beginning of continuous snow cover (yr <sup>-1</sup> )	15 Nov	16 Oct	16 Oct	08 Nov
Greatest snow depth (cm) <sup>2</sup>	87	93	93	82
Date for snowmelt <sup>3</sup>	18 May	16 May	17 May	18 May
Mean PAR <sup>4</sup> May–Aug ( $\mu$ mol m <sup>-2</sup> s <sup>-1</sup> )	306	341	317	256

 $^1$  With 5 °C threshold.  $^2$  Measured at Värriö Subarctic Research Station.  $^3$  Date when soil temperature rose higher than 0.3 °C.  $^4$  PAR = photosynthetically active radiation.

direction. This distance (roughly 25 m away) corresponds to the location of the subcanopy eddy flux system.

#### 3.4. Ecosystem-scale fluxes

The daily sum of NEE was positive, i.e. the forest was a source of carbon during the off season, transforming into a carbon sink around mid-May (Fig. 1D). Daily NEE returned to positive usually in early September. Daily NEE was sometimes positive during the active season mainly on rainy days accompanied by low radiation. For example, the daily NEE became positive during rainy and cloudy weather in mid-July 2015, when mean daily PAR was only half of the intensity compared with the week before and after (Fig. 1BD). In general, the carbon sink was largest (i.e. NEE was most negative) in June. Due to low turbulence, 34–40% of the 30-minute NEE measurements were missing and were gap-filled during May–September. The missing measurements



Distance from tower (m)

**Fig. 2.** The different shareds of grey represent the estimated 80% footprint distances around the ecosystem-scale eddy covariance tower (cross) in 2013–2015. For clarity, the small dark area represents the mean 80% footprint distances of the subcanopy eddy covariance system only in 2013 as the other years are comparable with it.

increased the need for gap-filling even more during the snow-covered seasons.

Gap-filled, daily  $R_E$  and  $R_{FF}$  mostly had a very similar pattern, but  $R_{FF}$  was on average 22, 20 and 18% smaller than  $R_E$  between 1 May and 31 August in 2013, 2014 and 2015, respectively (Fig. 3). The mean  $R_{FF}$  measured with soil chambers was slightly higher or close to  $R_E$  (Fig. 3). NFFE suffered from low turbulence close to ground level and technical problems in 2013–2014, resulting in up to 68, 76 and 52% of data being gap-filled during 1 June to 30 September in 2013, 2014 and 2015,



**Fig. 3.** Mean daily forest floor respiration (*R*) measured with the manual soil chambers and with eddy covariance above ( $R_E$ ) and below canopy ( $R_{FF}$ ) in 2013–2015. The error bars represent the ± standard deviation of the manual chamber measurements (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).

respectively. The range between the minimum and maximum daily NFFE estimates from the different flux partitioning parameterizations was less than 0.43 g C m<sup>-2</sup> on 90% of the days between 1 May and 30 August. When summing up the whole period, the difference between the minimum and maximum was 5.4 g C m<sup>-2</sup>, i.e. approx. 5% of the NFFE.

#### 3.5. Forest floor photosynthesis by the three methods

The overall level and annual patterns of the two estimates for forest floor photosynthesis ( $P_{FF}^{Down}$  and  $P_{FF}^{Dp}$ ) were comparable especially in the middle of the growing seasons even though in the spring,  $P_{FF}^{Down}$  mostly increased later than  $P_{FF}^{Dp}$  (Fig. 4).  $P_{FF}^{FC}$  corresponded to the overall level reached by the other two approaches, but the high need for gap-filling

especially in the early and late season made the comparison difficult particularly in 2013 and 2014 (Fig. 5). In 2015,  $P_{FF}^{Up}$  followed pretty closely the overall pattern of  $P_{FF}^{EC}$ , whereas  $P_{FF}^{Down}$  was close to  $P_{FF}^{EC}$  in the middle of the season (Fig. 4). The range between the minimum and maximum daily  $P_{FF}^{EC}$  estimates by the different flux partitioning parameterizations tested was less than 0.52 g C m<sup>-2</sup> on 90% of the days between 1 May and 30 August 2015.

# 3.6. Canopy versus forest floor photosynthesis

Canopy photosynthesis ( $P_{E}^{CC-FF}$ , Eq. 12), already started to increase consistently in all years in April, whereas forest floor photosynthesis,  $P_{FF}^{Up}$ , began to increase later, around the time of snowmelt (Fig. 6). The highest daily values of canopy and forest floor photosynthesis were roughly equal in 2013–2014, whereas in 2012 and 2015, the daily rate



**Fig. 4.** Daily estimates for forest floor photosynthesis ( $P_{FF}$ ).  $P_{FF}^{Down}$  (Eq. 4) was calculated as the difference between the photosynthesis of the whole ecosystem and the model estimate for canopy photosynthesis.  $P_{FF}^{UD}$  was upscaled from mass-based photosynthesis rates (Eq. 11).  $P_{FF}^{EC}$  (red dots) are the below-canopy eddy covariance measurements with at least 25% measured daily data. The light grey areas indicate the snow-covered season. For clarity, the series are moving averages over 3 days (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).



**Fig. 5.** The two estimates for forest floor photosynthesis ( $P_{FF}^{Down}$ , blue;  $P_{FF}^{UD}$ , black) against below-canopy eddy covariance driven estimate ( $P_{FF}^{EC}$ ) for those days with at least 25% data coverage of  $P_{FF}^{EC}$  measurements. The black line represents a 1:1 linear relationship and the dashed lines represent the linear relationships between measured and modelled data.

of forest floor photosynthesis was mainly lower than that of the canopy. The daily canopy photosynthesis was higher than that of forest floor vegetation on 81, 61, 61 and 68% of days between 1 May and 30 September in 2012, 2013, 2014 and 2015, respectively.

#### 3.7. Annual net ecosystem exchange, respiration and photosynthesis

The stand was a carbon sink (-48 to  $-7 \text{ g C m}^{-2} \text{ yr}^{-1}$ ) during 2012–2014 but a small source of carbon in 2015 ( $14 \text{ g C m}^{-2} \text{ yr}^{-1}$ , Table 3). The sink was highest in 2013 accompanied by the highest annual  $P_E^{BC}$  but also the highest  $R_E$ . Both of these were lowest in 2015 (Table 3). The proportion of forest floor photosynthesis relative to the whole ecosystem varied from 43% to 49% when derived using upscaled

 $P_{FF}^{Up}$  (Eq. 11). It was highest in 2013 (49%) and lowest in 2012 and 2015 (43%).

Annual  $P_E^{EC}$  correlated positively with PAR (Table 4, R<sup>2</sup> = 0.92) and temperature sum (R<sup>2</sup> = 0.89), and negatively with precipitation (R<sup>2</sup> = 0.96). Annual  $R_E$  correlated positively with temperature sum (R<sup>2</sup> = 0.99) and negatively with precipitation (R<sup>2</sup> = 0.74, not shown). The mean increase in  $P_E^{EC}$  with an increase in the temperature sum was 0.27 g C m<sup>-2</sup> yr<sup>-1</sup> per degree day, whereas in the case of  $R_E$  it was 0.17 g C m<sup>-2</sup>yr<sup>-1</sup> per degree day. The different estimates of canopy photosynthesis,  $P_C^{SPP}$  and  $P_C^{EC-FF}$ , showed very little response to annual variation in temperature sum (0.06–0.08 g C m<sup>-2</sup> yr<sup>-1</sup> increase per degree day), whereas the GPP of forest floor vegetation increased more with increasing temperature (0.19–0.21 g C m<sup>-2</sup> yr<sup>-1</sup> per degree day)



**Fig. 6.** Daily estimates of the canopy  $(P_C^{EC-FF})$  and forest floor photosynthesis  $(P_{FF}^{U_D})$ .  $P_{FF}^{U_P}$  was upscaled from mass-based photosynthesis rates (Eq. 11) and  $P_C^{EC-FF}$  was calculated as the difference between the total ecosystem photosynthesis and  $P_{FF}^{U_D}$  (Eq. 12). The light grey areas illustrate the snow-covered season. For clarity, photosynthesis is represented as a 3-day moving average (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).

#### Table 3

Yearly cumulative net ecosystem exchange (NEE), total ecosystem respiration ( $R_E$ ), gross primary production of the whole ecosystem ( $P_E^{FC}$ ), two estimates for the photosynthesis of the tree canopy,  $P_C$ , and two estimates for the forest floor photosynthesis,  $P_{FF}$ . Negative NEE indicates that the site was a carbon sink. Values in parentheses are cumulative values from May to September.

$\begin{array}{c} \text{NEE } (\text{g C m}^{-2} \text{ yr}^{-1}) & - \\ R_E (\text{g C m}^{-2} \text{ yr}^{-1}) & 4 \\ P_E^{EC} (\text{g C m}^{-2} \text{ yr}^{-1}) & 5 \\ P_C^{EC-Up} (\text{g C m}^{-2} \text{ yr}^{-1}) & 3 \\ P_C^{SPP} (\text{g C m}^{-2} \text{ yr}^{-1}) & 3 \\ P_{FF}^{CP} (\text{g C m}^{-2} \text{ yr}^{-1}) & 2 \\ P_{FF}^{Down} (\text{g C m}^{-2} \text{ yr}^{-1}) & 2 \end{array}$	- 35 (- 117)	-48 (-140)	-7 (-77)	14 (-69)
	92 (333)	553 (373)	529 (375)	489 (321)
	27 (453)	600 (512)	537 (452)	475 (390)
	01 (255)	306 (255)	283 (231)	272 (223)
	08 (261)	332 (281)	328 (273)	314 (252)
	26 (195)	294 (257)	253 (221)	203 (167)
	19 (188)	269 (231)	208 (179)	161 (138)

#### Table 4

Slopes and p-values of the linear relationship between different photosynthesis estimates (*P*) for ecosystem (E), canopy (C) and forest floor (FF) together with annual weather characteristics: photosynthetically active radiation (PAR) (µmol m<sup>-2</sup> s<sup>-1</sup>), temperature sum (°C) and precipitation (mm). The slope indicates the change in photosynthesis (g C m<sup>-2</sup> yr<sup>-1</sup>) per unit of the weather characteristic.

	PAR		Temperature sum		Precipitation	
$P_E^{EC}$ $P_C^{EC-Up}$ $P_C^{SPP}$ $P_{FF}^{Up}$ $P_{FF}^{Down}$	Slope	<i>p</i> -value	Slope	<i>p</i> -value	Slope	<i>p</i> -value
	1.38	0.04	0.27	0.06	- 0.69	0.02
	0.36	0.17	0.05	0.39	- 0.18	0.16
	0.22	0.33	0.06	0.10	- 0.11	0.33
	1.01	0.07	0.21	0.01	- 0.51	0.05
	1.17	0.05	0.21	0.14	- 0.58	0.03

meaning that the ecosystem-scale variation was mainly driven by annual variation in forest floor vegetation (Table 4). However, the analysis suffered from a low number of statistically significant relationships even though many *p*-values were relatively low (Table 4).

### 3.8. Growth variation

The annual length increment of dwarf shrubs showed high variability, and the differences were mainly not significant between years, especially in the evergreen species. However, the growth was lowest in all species in the cold and cloudy summer of 2015, while the differences in growth between years were significant for *V. myrtillus* and *E. nigrum* (Fig. 7). The growth of *V. myrtillus* in 2012 was significantly higher than in other years. The variation in growth was not connected to the variation in the annual  $P_{FF}^{Down}$  or  $P_{FF}^{Up}$ .

### 4. Discussion

We studied the  $CO_2$  exchange in a subarctic pine stand in northern Finland for 4 years with distinct weather characteristics and found that during three of these years, the site acted as a weak sink, whereas it was a source of carbon during the coldest and rainiest year with the lowest radiation. The strength of the  $CO_2$  uptake (NEE) was much lower than the mean of forested and tundra sites in Alaska (USA) reported by Ueyama et al. (2013). However, our estimates for GPP in Värriö matched their mean findings. In our study, the total ecosystem respiration was higher than in the Alaskan study and strongly correlated with the temperature sum. The correlation is, on the other hand, evident due to its computational connection to soil temperature (Eq. 3) even though the parameter estimation in one time period (11 days) was independent from other periods. Nevertheless, the relationship between the temperature sum and GPP was even stronger: the carbon sink was largest in the warmest year and lowest in the coldest growing season during the measurement years. This temperature dependency of GPP is in line with several studies (Law et al., 2002; Mäkelä et al., 2008) and our results indicate that without any other modifications, even a relatively small increase in growing season temperature would increase the carbon sink strength of this site. Thus, it seems that the short-term increase in photosynthetic productivity did not increase decomposition unlike that presented by Hartley et al. (2012) and Parker et al. (2015) via the rhizosphere priming effect (Read et al., 2004), but naturally the subject would benefit from a longer time series of high-quality data. The negative correlation between photosynthesis and precipitation most probably just reflected the inverse relationship between radiation and precipitation.

We estimated the photosynthesis of the forest floor using three independent methods: 1) directly measuring carbon fluxes with our subcanopy eddy covariance system, 2) downscaling the flux as a difference between the total ecosystem and estimated canopy photosynthesis (top-down), and 3) upscaling it using the forest floor biomass and available models for leaf-mass based photosynthesis (bottom-up). The overall levels of daily uptake in midsummer were surprisingly similar in all three independent methods (Fig. 4) thus giving credibility to the annual estimates, but each of these methods has significant pros and cons. Potential uncertainties of the three approaches are discussed next.

Firstly, the bottom-up approach considers only the vascular plants and ignores the large biomass of mosses found at our measurement site. Even though their photosynthesis is notably smaller than that of vascular plants (Kulmala, 2011 and references cited therein), they naturally take part in the photosynthetic uptake of the stand (Street et al., 2013). Available leaf-mass based estimates for the maximal photosynthesis of feather mosses range between 4 and 20 nmol  $g^{-1}$  s<sup>-1</sup> (McCall and Martin, 1991; Kulmala et al., 2009, 2011a), whereas those for V. myrtillus, V. vitis-idaea and C. vulgaris are in general five times as high (Kulmala, 2011), varying between 33 and 184 nmol  $g^{-1}$   $s^{-1}$ (Widen, 2002; Kolari et al., 2006; Kulmala et al., 2008, 2009, 2011a). In practice, the momentary photosynthesis increased at most by 0.31 or  $0.78 \text{ g C m}^{-2} \text{day}^{-1}$  in 2015 if we assume that 50% of moss biomass is photosynthetically active and leaf-mass based maximal photosynthesis of mosses is 4 or 10 nmol  $g^{-1}s^{-1}$ , respectively. Annually, this would mean an increase of 19 or 46 g C m<sup>-2</sup>yr<sup>-1</sup> i.e. 9% and 22% respectively of the annual GPP of the forest floor vegetation in 2015.

The good fit between the direct measurements and the upscaled photosynthesis accounting only for vascular plants might result from a difference in the leaf-mass based rates adopted from southern and thus more fertile environments compared with this northern study site. A number of studies have found that the rates of photosynthesis per leaf weight, specific leaf area and leaf mass ratio are higher in plants grown in high-nitrogen soils (Field et al., 1983; Field and Mooney, 1983; Poorter et al., 1995; Masarovicova et al., 2000). However, the nitrogen concentrations in Scots pine needles in the site measured by Kulmala et al. (2011a); and the study site do not differ from each other (Palmroth and Hari, 2001; Susiluoto et al., 2010). Besides, Kulmala et al. (2011b) used the PAR at the forest floor, whereas in this study the model uses the PAR measured above the canopy, which might overestimate the photosynthesis of dwarf shrubs and possibly compensate for the impact of missing mosses. Nevertheless, the actual reason behind the good fit between measurements and model remains unknown before revealing the species-specific photosynthesis rates and the radiation environment at the in-situ forest floor.

One error source in the upscaling could be the stable leaf biomass used in the calculation. However, most of our species (Table S1) have evergreen leaves with a lifetime of several years, which buffers some effects of the year-to-year variation in the growth of leaf mass. It was also shown by Köster et al. (2017) that the biomass of ground vegetation in northern Finland develops slowly. In addition, the constant leaf mass used in calculating the GPP of the ground vegetation in our study leads to rather close estimates between the direct measurements and the indirect method (Fig. 4), supporting the assumption that the



Fig. 7. Relative height increment of Vaccinium myrtillus (A), Vaccinium vitis-idaea (B) and Empetrum nigrum (C) in the different years. The final length differed significantly between years if the lower-case letters differ.

photosynthesizing leaf mass is rather stable. Nevertheless, the use of the introduced photosynthesis model still requires measurements of the overall level of photosynthesis as long as there is no further information available about the biomass variation and the actual leaf-mass based photosynthesis rates at the study area.

Third, the bottom-up model is mainly driven by the temperature and radiation and thus it is evident that the upscaled annual photosynthetic production is higher in sunny and warm growing seasons than in cloudy and cold ones. However, the direct below-canopy eddy covariance measurements and the independent top-down method give a comparable estimate of the daily forest floor photosynthesis, and the latter also of the annual estimate, giving support for the model formulation and result. Thus there is relatively strong evidence that the annual variation in the growing season weather mainly affects the yearto-year variation of forest floor photosynthesis, whereas the tree canopy photosynthesis does not consistently vary with the mean growing season radiation or temperature environment in the observed range.

The importance of canopy structure and especially the light penetration to the ground became evident in our forest floor carbon uptake results. The forest floor vegetation contributed ~45% to the total ecosystem photosynthetic uptake in this subarctic pine forest. For example, Ikawa et al. (2015) estimated the forest floor vegetation contribution to carbon uptake to be as high as 61% in an open black spruce forest in Alaska. Furthermore, our GPP estimates for the forest floor vegetation  $(184-266 \text{ g C m}^{-2})$  fall between the estimates in two young Scots pine stands in southern Finland of 349 and 168 g C  $m^{-2}$ , with above tree canopy all-sided LAI of 1.4 and 5.1 m<sup>2</sup>m<sup>-2</sup>, respectively (Kulmala et al., 2009, 2011b). However, most annual estimates of the proportion of forest floor vegetation in the boreal region lie between 10% and 20% (Swanson and Flanagan, 2001; Kolari et al., 2006; Ilvesniemi et al., 2009; Bergeron et al., 2009). Conversely, these estimates are from southern sites with higher canopy coverage than at our subarctic site, suggesting that there might be a clear relationship between the canopy leaf area and the GPP of ground vegetation and thus highlighting motivation for further studies.

There was no clear connection between the variation of GPP and the growth of dwarf shrub shoots, although all species grew less in 2015, which was also the year of low photosynthetic production. Even though it has been shown that the annual variation in the diameter growth of trees is connected to the annual GPP (Berninger et al., 2004; Gea-Izquierdo et al., 2014; Schiestl-Aalto et al., 2015; Kulmala et al., 2016), the buds of most species are usually formed already during the previous

year, with the respective late summer temperatures being used to describe the annual variation in the shoot growth of the next growing season (Salminen and Jalkanen, 2005). Nevertheless, the high GPP in 2013 did not seem to affect the height increment in 2014. Probably 4 years of measurements are too short for detecting such growth variation especially when facing lagged responses. Net primary production (NPP), i.e. the growth of biomass, is the difference between photosynthetic production and other carbon sinks such as vital functions, reproduction, changes in internal storage, root exudation, etc. Thus, to obtain a full picture, all other sinks should be subtracted from the primary production. For example, Pumpanen et al. (2012) studied small seedlings and observed that although both photosynthetic production and maintenance respiration increased with increasing temperature, this resulted in an insignificant relationship between temperature and NPP. Also, autocorrelation, changes in leaf area, diseases and other periodical disturbances should be included in order to connect photosynthetic production and growth.

In theory, the fluxes obtained via soil chambers should equal  $R_{FF}$ , which was the case in 2015, whereas in 2013-2014, soil chambers showed higher respiration rates than  $R_{FF}$  in the middle of the growing season. Thus, our site is among the group of ecosystems in which even  $R_E$  is lower than that of soil chamber measurements (Van Gorsel et al., 2007; Wang et al., 2010; Barba et al., 2018). This phenomenon is not fully explained yet, but among the likely causes for this kind of result are footprint related issues, potential error sources in chamber measurement, below-canopy horizontal advection (e.g. Aubinet et al., 2005; Wang et al., 2017), and in the case of ecosystem-scale measurements, also decoupling of below- and above-canopy air mass flow (e.g. Alekseychik et al., 2013; Jocher et al., 2017). The eddy covariance footprints are constantly changing depending on wind and atmospheric stability and thus the two measurement heights integrate fluxes from different areas. Nevertheless, the footprint analysis revealed that not only the location itself but also the extent of the subcanopy footprint (measurement height 2.7 m) overlaps with that of the main tower in the south-western direction. In our study the problem might arise from the chamber measurements as the  $R_{FF}$  by those is mostly higher than that of the subcanopy EC measurements. Nevertheless, the estimation of  $R_E$ from high latitude eddy covariance data is more complicated compared with that using lower latitude data due to 24 h of daylight and the complexity of defining night-time and hence data partitioning close to midsummer, for example. The daytime method tends to give smaller  $R_E$ than the night-time method (Lasslop et al., 2010). However, the

relatively similar estimates for  $P_{FF}$  (or  $P_C$ ) derived with the eddy covariance technique and the independent methods give confidence in the overall level of  $P_E$  and thus also in  $R_E$ . In addition, our strict filtering policy should remove most of the conditions difficult for eddy covariance measurements, such as calm and stable nights.

#### 5. Conclusions

Northern forests are predicted to experience extensive climate change, which will modify the carbon exchange between ecosystems and the atmosphere. Our results indicate that the studied subarctic site is able to improve its carbon sink if temperature is slightly increased. assuming there are no other climate change related changes. Comparison between canopy and forest floor vegetation revealed that the seasonal dynamics differ especially in springtime when canopy photosynthesis increases approximately one month before snowmelt and the increase of forest floor photosynthesis. This highlights that models of primary production driven without snow cover depth are very likely to inaccurately estimate the seasonal dynamics of photosynthesis in northern forests. That said, the precise modelling of the carbon cycle becomes increasingly important especially regarding the impact of climate change in northern, most rapidly warming ecosystems. We did not find evidence for source limited growth in the annual variation of the height increment of dwarf shrubs, but further studies are required for a better understanding of the sink-source dynamics of growth in these high latitude forests.

#### Data availability

We have made publicly available the used gap-filled and partitioned eddy covariance data, meteorological records, soil chamber and shoot growth measurements, attained pine shoot chamber parameters and the Stand Photosynthesis Program model input and output as well as the program itself with its graphical user interface. All can be downloaded at: https://b2drop.eudat.eu/s/LzisEZiQG96nDYH

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# Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.agrformet.2019.02. 029.

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