

Matching scales of eddy covariance measurements and process-based modeling - Assessing spatiotemporal dynamics of carbon and water fluxes in a mixed forest in Southern Germany

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Abstract. Eddy covariance (EC) measurements are a backbone of ecological research and have provided valuable insights into the variability of carbon and water fluxes in different ecosystems and under varying environmental conditions. Since
20 these measurements are integrative and weighted over changing areas (footprint), species-specific information cannot be easily derived except for homogenous monocultures. However, EC sites are increasingly established in mixed forest stands which are considered to be more resilient under changing environmental conditions. This leads to the question of how species-specific responses can be determined, and whether the magnitude of fluxes derived from temporally varying flux footprint predictions (FFPs) can provide insight into these responses.

25 At a site in southwestern Germany's Black Forest, primarily composed of mature beech and Douglas fir trees, we investigate the dependence of EC flux measurements on different FFP areas and explore how species-specific contributions to gas exchange can be disentangled using a combined measurement and modelling framework. We applied an ecosystem model that has been calibrated from EC measurements at various sites with beech- and Douglas fir monocultures, and evaluated it with data of soil water content and soil respiration taken at homogeneous parts of the investigated mixed forest site. Then we
30 compared hourly aggregated measurements of net carbon exchange (NEE) and evapotranspiration (ET) with model simulations under four configurations: (i) pure beech, (ii) pure Douglas fir, (iii) a static weighted average of both species, and (iv) a dynamic weighted average based on FFP variations.

The results show that weighted combinations of the two species generally provide a better match with hourly EC measurements than single-species simulations, while differences between static and dynamic weighting approaches remain relatively

35 small. However, species-specific flux responses can be significantly different during transitional periods such as autumn and
spring when physiological differences between Douglas fir and beeches are most pronounced. We demonstrate that account-
ing for seasonal differences is particularly important for gap-filling EC measurements in mixed forests and, consequently, for
determining annual carbon and water budgets. Furthermore, EC measurements over mixed forests provide valuable informa-
tion for detailed model evaluation, while species-specific modelling helps disentangle and attribute underlying ecosystem
40 dynamics to individual species.

1. Introduction

Forests play a vital role in regulating both global and national carbon and greenhouse gas (GHG) budgets, generally by act-
ing as significant carbon sinks (Pan et al., 2011). However, the increasing frequency and intensity of extreme climatic events
— such as droughts, storms, and heatwaves — poses a substantial threat to this function, potentially diminishing forests' ca-
45 pacity to sequester carbon or even turning them into net GHG (Anderegg et al., 2020; Haberstroh et al., 2025; Thum et al.,
2025; van der Woude et al., 2023). To better understand and predict how forest ecosystems respond to changing environ-
mental conditions, process-based ecosystem models are often used (e.g. Collalti et al., 2018; Dirnböck et al., 2018; Mäkelä et
al., 2023). Such models need to be calibrated and evaluated with site measurements, for which the eddy covariance (EC)
method is a key tool providing data on ecosystem carbon (C) and water (H₂O) exchange (Baldocchi, 2003). However,
50 process-based models require not only net fluxes but also their respective sink and source terms, and often rely on species-
specific information, which is not easily derived from spatially integrated EC measurements (Stoy et al., 2019).

Despite these constraints, EC measurements have been established as a backbone for ecosystem research, providing key data
that are essential for model calibration and evaluation. Numerous EC flux towers have been installed throughout recent
decades to investigate carbon and water exchange in grasslands, agricultural areas, and forests (Baldocchi, 2003; Teuling et
55 al., 2010). These measurements represent integrated gas exchange fluxes over an area whose size depends on wind condi-
tions and sensor height, commonly referred to as the flux footprint prediction, FFP (Kljun et al., 2004; Schmid, 1994;
Schuepp et al., 1990; Vesala et al., 2008). Thus, ecosystem scale measurements are represented by temporally dynamic FFPs
that differ in size and location. Consequently, interpretation of EC measurements is becoming increasingly complex with in-
creasing variation of ecosystem properties in different directions or distance to a respective tower (Fang et al., 2024; Grote et
60 al., 2011b). However, the directional bias can also be exploited to extract additional information on fluxes from underlying
land cover patches in spatially heterogeneous ecosystems (Cassidy et al., 2016; Helbig et al., 2017; Tuovinen et al., 2019; Xu
et al., 2020). To date, most cases where EC measurements have been used to evaluate ecosystem models were done at sites
assumed to be homogeneous or corrected for directional bias. In the case of forested areas these sites are typically even-aged
monocultures, characterized by a similar forest structure in all cardinal directions (Mahnken et al., 2022). At sites that are
65 less homogenous, it is important to determine the differences in gas exchange rates that can be attributed to different forest

structures and compositions in specific FFPs, which might vary not only with wind speed and direction, but also on diurnal or seasonal timescales.

It should be considered that without a spatial analysis of the flux footprint and its variation over time, comparison of fluxes with simulation results can be difficult to interpret. By averaging forest structure over large areas for model application, the model may capture overall EC fluxes during the evaluation period. However, species-specific responses can still lead to deviations between simulations and measurements under particular conditions, such as heatwaves and droughts (Remy et al., 2019). Similarly, caution is advised for periods when EC data are missing. Such periods that require statistical gap filling in order to derive complete budgets for whole days, seasons, or years can be substantial. Such gap filling is usually applied without considering temporally and spatially changing footprint contributions or species composition, which could lead to biases. In addition, missing knowledge about short-term (Fox et al., 2009) or long-term (Mahecha et al., 2007) ecosystem dynamics may lead to substantial under- or overestimation of fluxes and increases the uncertainty of extrapolations into the future. Along with other issues, this data-related uncertainty has been highlighted as important for model evaluation (Medlyn et al., 2005), but only few approaches have been investigated this uncertainty explicitly at EC flux tower sites (e.g. Kutsch et al., 2005; Oishi et al., 2008), particularly in comparison with model applications indicating also the importance for spatial and temporal model resolution.

Therefore, the first objective of this study is to assess the uncertainty of fluxes when using EC measurements for evaluation of vegetation models that are initialized at specific footprint areas differing in species composition. In particular, we examine how variations in species composition affect evapotranspiration (ET) and net ecosystem exchange (NEE) when dynamically changing flux footprint predictions (FFPs) are considered. The second objective is to characterize the flux contribution of Douglas fir and beech within a mixed forest area, using a combined measurement and modelling approach. We hypothesize that simulations representing the species contribution in the FFP can provide a complementary perspective to conventional statistical gap-filling procedures by enabling species-resolved interpretations of missing fluxes.

To achieve these objectives, we i) quantify hourly varying species contributions based on flux footprint predictions (FFPs) of an EC measurement site located in a heterogeneous forest, and ii) evaluate the impact of FFP-specific species composition on model-measurement comparisons of net CO₂ exchange and latent heat flux. The different species contributions are analyzed on different temporal scales, e.g. to differentiate between growing and non-growing season or different phenological periods. The considered site is a mixed forest dominated by patches of European beech and Douglas fir - tree species that will play a considerably larger role for future German forestry (Brandl et al., 2020; Gribbe et al., 2024). The simulations are carried out with the LandscapeDNDC model (Haas et al., 2013), which has therefore been calibrated at independent sites with pure beech (Herbst et al., 2015; McGloin et al., 2018; Pilegaard et al., 2003) and Douglas fir forests (Morgenstern et al., 2004; Van Wijk et al., 2001). The sites and calibration results are further described in the methods section and in the supplementary information.

2. Materials and methods

2.1 Site description

100 The measurement site (48.2685°N, 7.8782°E, 490 m a.s.l.), called ECOSENSE forest, is located in the Black Forest close to the town of Ettenheim in Southwest Germany. It is a mixed forest and the area we assumed as potential flux footprint area is dominated by European beech (*Fagus sylvatica*, 47%), but holds major shares of Silver fir (*Abies alba*, 25%) and Douglas fir (*Pseudotsuga menziesii*, 21%) (Fig. 1). The mixture type is a group mixture with firs concentrating in an approximately 200 × 20 m stripe within a beech forest. The average height of the dominant trees (larger than 20 cm diameter at breast height),
105 as well as the upper height (standard deviation added to the average) is only slightly different between the species (25.5 and 27.8 m for beech, 26.0 and 29.0 m for coniferous trees, respectively). Within the ECOSENSE forest, a 46-m tall measurement tower was instrumented with an EC system and various measurement devices designated to capture carbon and water fluxes and stress relate responses (Werner et al., 2024). According to the Köppen classification, the climate at this site is temperate oceanic with mild to warm summers and cold winters. The mean annual air temperature is about 9.7°C (ranging
110 from 1.1°C in January to 18.8°C in July) and mean annual precipitation is approximately 1000 mm over the last 20 years using historical data from the nearest meteorological Lahr station (ID 2812) of the German Weather Service (DWD). Geologically, the area lies on Triassic sedimentary rock, predominantly the Plattensandstein Formation — sandstone interbedded with clay layers — with minor outcrops of the Rotton Formation. The soils at the study site are Cambisols with silty loam to loamy clay textures, free of carbonates, and well-developed to depths of 60–120 cm. They exhibit moderate permeability and
115 low stone content (Werner et al., 2024).

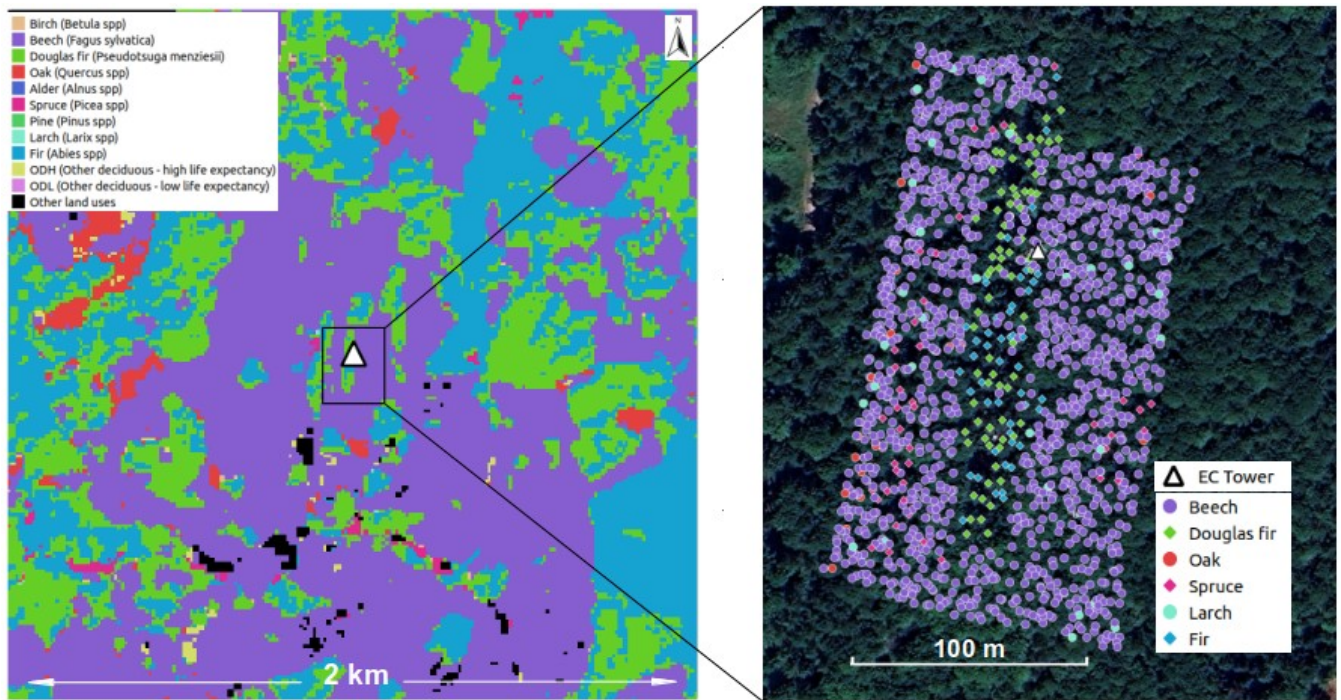


Figure 1: Dominant tree species map within the 2×2 km window (left) and the tree inventory in the ECOSENSE experimental site (right) from Blickensdörfer et al. (2022). The white triangle shows the location of the eddy covariance (EC) tower. The aerial photo is from Google Earth (© Google Earth 2025).

2.2 Site measurements

2.2.1 Eddy covariance measurements

The EC technique was used to measure the ecosystem CO_2 and water vapor fluxes (evapotranspiration, ET) above the ECOSENSE forest (Aubinet et al., 2012; Burba et al., 2013). The EC instrumentation consists of a closed-path fully heated infrared gas analyzer (model Li-7200, LI-COR Inc., Lincoln, NE, USA) combined with a three-dimensional sonic anemometer (model CSAT3B, Campbell Scientific Inc., Logan, UT, USA), both measuring at a frequency of 20 Hz. Due to technical problems, the heating system was not working until early December 2024. The EC measurements were performed approximately 18 m above the canopy of the ECOSENSE forest on top of a measurement tower at a height of 46 m above ground level.

EddyPro (version 7.0.9) was used to calculate half-hourly fluxes, which were later averaged to hourly fluxes, from the 20 Hz measurements of the infrared gas analyzer and the three-dimensional sonic anemometer (LI-COR Biosciences, 2022). In this study, only CO_2 fluxes and ET data with quality flags 0 or 1 were used for further analysis, data with quality flag 2 were discarded. The REdDyProc software was used for further processing of the EC data (Wutzler et al., 2018), including u -filtering according to Papale et al. (2006) and gap filling using the marginal distribution sampling (MDS) method (Reichstein et al.,

135 2005). The MDS method applies look-up tables for similar meteorological conditions, considering shortwave irradiance, air temperature, and vapor pressure deficit, to fill the gaps in the flux data. If no meteorological data are available, the gaps in the flux data are filled using the MDS method with the mean daily course of the fluxes in a moving time window of adjacent days (Reichstein et al., 2005; Wutzler et al., 2018). A flux footprint prediction climatology was created for every hour with available data from the EC system, using the R-code of Kljun et al. (2015). The FFPs were calculated for a domain of 2×2 140 km centered on the tower at a spatial resolution of 1×1 m. The input data for the flux footprint predictions were taken from the EddyPro output, except planetary boundary layer height, which was retrieved from the ERA5-Land weather reanalysis dataset (Hersbach et al., 2023). The available EC data from the first year after start of the measurements (15 May 2024 to 15 May 2025) were used for the presented analysis. For FFP calculations over forested areas, the zero-plane displacement (d) must be considered. In absence of vertical wind profile measurements, we followed the common practice to assume that d 145 equals two-thirds of the canopy height. However, since the reported values of d in relation to canopy height range between 0.5 and 0.8 (Oliveira et al., 2022), we tested the sensitivity to the choice of this parameter varying it within this range (-25% to $+20\%$ relative to the standard value). Overall, the effect on flux contribution to the variation of d was less than 1% in comparison to the reference case (see Table S2 in the supplementary).

2.2.2 Auxiliary measurements

150 Meteorological data

Several meteorological variables required as input for the LandscapeDNDC model are measured at the study site. Solar irradiance was measured by a heated and ventilated pyranometer (model CNR4, Kipp & Zonen, Delft, Netherland) mounted on the tower above the forest at a height of 47 m. Wind speed and direction was derived from the three-dimensional sonic anemometer integrated in the EC system. Unless data from the EC system were unavailable, data from a propeller anemometer (model Young 05103, R. M. Young Company, Traverse City, Michigan, USA) also mounted on top of the tower were 155 used. Air temperature and relative humidity are taken in the vicinity of the site under open field conditions with sensors mounted at a height of 2 m above ground level inside a passive ventilated radiation shield (model HygroVue 10 and RAD10E, Campbell Scientific Inc., Logan, UT, USA).

Soil moisture

160 In the ECOSENSE forest permanently installed soil moisture sensors SMT100 (Truebner GmbH, Neustadt, Germany) are used to measure dielectric permittivity, soil temperature and volumetric soil water content based on time domain transmission (TDT) principle. A total of 40 sensors were installed in 10 profiles of 4 depths (30, 50, 70 and 90 cm) at the center (plateau) of the observed EC footprint area continuously measuring at a frequency of 15 min. For investigating species influences, two plots were selected influenced only by either purely beech or Douglas fir respectively, each holding seven neigh- 165 boring trees of similar age and height. Within each of the plots, five soil moisture profiles were arranged in a stratified-ran-

dom design to ensure a good spatial coverage. First measurements started in October 2023 and are ongoing. All raw data undergo a quality control procedure adapted from Dorigo et al. (2013).

Soil respiration

170 Soil respiration was measured with an open bottom chamber equipped with a CO₂-Sensor (GMP343, Vaisala Oyj, Finland) at a total of 35 plots (2.25 m² per plot). The 35 plots were measured once within one day but always in a different order to prevent any diurnal effects. We recorded the tree species for each plot within a vicinity of 5 m (radius) and for model evaluation we only chose plots influenced by either pure European beech (18 plots) or pure Douglas fir (8 plots) composition. Soil respiration was measured on a weekly to bi-weekly basis using two measurement chambers simultaneously during the complete year 2024. To avoid saturation effects at the end of the measurement or too slow increase at the beginning of the measurement, time was varied depending on the season between 90 and 140 seconds. From both measurements we calculated a mean value, reporting the respiration for each plot. In very rare cases (< 1%) system errors happened and the measurement time is only 80 seconds (1 of 1750 cases) or only one measurement per plot per day is available (3 of 1750 cases).

2.3 LandscapeDNDC model

2.3.1 Model description

180 LandscapeDNDC (<https://ldnc.imk-ifu.kit.edu>, last access: August 15, 2025) is a modular terrestrial ecosystem model (Grote et al., 2011b; Haas et al., 2013). The physiological simulation model (PSIM) of the model framework, which is used for representing forest related processes, is based on cohorts represented by trees with equal dimensions and homogeneous spatial distribution. It has been designed to reproduce atmosphere–biosphere exchange processes of carbon, water, and nitrogen, including C and N trace gas exchanges. For this purpose, detailed microclimate, biogeochemical, and physical soil process modules are provided to be coupled with vegetation modules (i.e. physiology and structure) that are parameterized at the species level. Consequently, all processes directly or indirectly depend on environmental drivers. The LandscapeDNDC model uses air temperature, global radiation, vapor pressure deficit (or relative humidity), and precipitation as meteorological inputs in daily to sub-daily temporal resolution. Soil as well as canopy are 1-d divided into multiple layers, with flexible extensions and properties, depending on available measurements and the initialized ecosystem structure.

190 Forest carbon uptake and loss are calculated within PSIM from the basic processes which are photosynthesis and respiration, allocation and senescence (separately for each cohort, in monocultures there is only one). Photosynthesis is calculated according to the Farquhar model (Farquhar et al., 1980), which is linked to a soil water-limited stomatal conductance module to optimize gas exchange (Leuning, 1995). Stomatal conductance and soil water availability is thus also defining transpiration. Other relevant gaseous water fluxes are evaporation from interception, which is calculated based on leaf area dependent canopy capacity, as well as from snow, open water at the ground and soil. These evaporation terms are driven by potential

evaporation determined by a modified Thornthwaite approach (Thornthwaite and Mather, 1957). All of these processes are calculated in hourly resolution, for more details see descriptions in Holst et al. (2010). Respiration is differentiated into autotrophic respiration, which in turn originates from growth (fixed fraction of carbon allocated to increase biomass), maintenance (in dependence on tissue temperature and nitrogen concentration, according to Cannell and Thornley (2000) and nitrogen uptake (fixed rate related to nitrate uptake), and heterotrophic respiration related to microbial decomposition of organic matter driven by soil moisture, pH, and temperature. Carbon (and nitrogen) is allocated into the compartments: leaves, fine roots, living wood, and structural reserves according to their individual sink strength (Grote, 1998), with leaf demand driven by phenology which in turn depends on cumulated temperature and chilling requirements. Senescence of all compartments are determined by their turnover rates, and all dead materials are added to the soil carbon (and nitrogen) pools, where the explicit calculation of decomposition processes supplies heterotrophic respiration.

LandscapeDNDC has demonstrated that it could represent EC fluxes at sites dominated by oaks, pines, spruces, and beeches before (Cade et al., 2021; Molina-Herrera et al., 2015; Nadal-Sala et al., 2021). It also performed well in multi-model comparisons which evaluated aspects including forest growth (Cameron et al., 2013; Mahnken et al., 2022) and has been used to investigate carbon and water fluxes in various forested ecosystems in European sites (Dirnböck et al., 2020; Magh et al., 2019) as well as worldwide (Rahimi et al., 2021).

2.3.2 Model initialization and driving forces

In this study we used hourly meteorological data for the period from 15 May 2024 to 15 May 2025, corresponding to available EC time series at the ECOSENSE site (Fig. 2). We considered a spin-up period of 1.5 years starting in January 2023, which showed to be sufficient to allow carbon pools in the soil to equilibrate. Hourly air temperature, global radiation, wind speed, and relative humidity were taken from on-site measurements (see section 2.2.2). Hourly precipitation data from the meteorological station Lahr operated by the German Weather Service (Deutscher Wetterdienst; DWD; <https://cdc.dwd.de>), located approximately 7 km from the ECOSENSE site.

The soil initialization of the model is based on a vertical profile of soil physicochemical characteristics i.e., humus type, clay and sand content, organic C- and N-content, bulk density, saturated conductivity, stone content, pH, field capacity, and wilting point. (Table 1). Only one single on-site soil profile was available for this purpose that we used for all simulations. Regarding the vegetation, we initialized two separate forest types, one representative for a beech and one for a Douglas fir forest, each indicating tree species, dominant height, tree diameter at breast height (DBH), and number of trees per hectare. In this initialization, we consider all coniferous trees to be Douglas fir, implicitly assuming that gas exchange responses of Silver fir are sufficiently similar to Douglas fir to be merged. The initialization for the beech and the Douglas fir simulations, respectively, include the upper height, mean diameter, and stem number which are derived from an individual tree inventory carried out across an area of 3 hectares surrounding the EC tower (Fig. 1, right). In addition, we used the dominant tree species map at 10 m resolution (Blickensdörfer et al., 2022) to determine the overall shares of coniferous (mostly Douglas fir

and Silver fir) and deciduous (mostly beech) within a potential footprint area which is a window of 2×2 km with the ECOSENSE site in the center (Fig. 1, left).

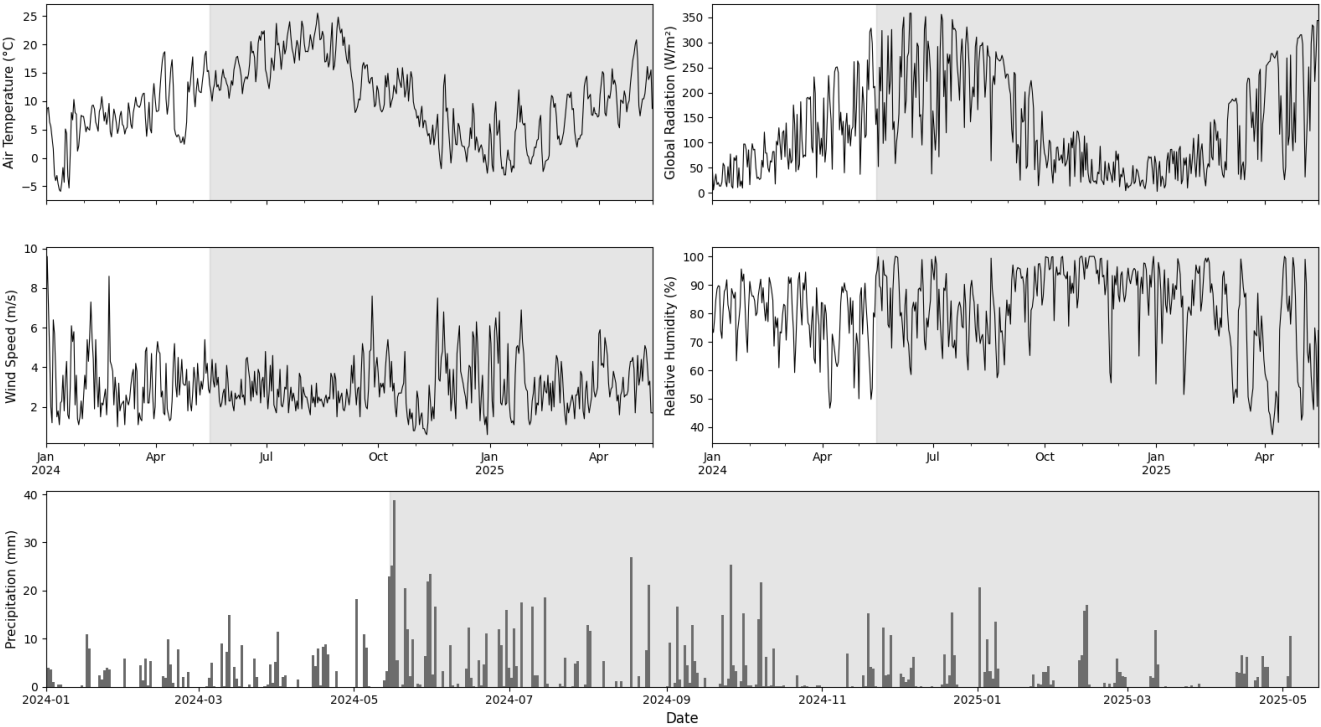
230 2.3.3 Model parameterization, calibration and evaluation setup

The LandscapeDNDC model in a similar configuration has been used to simulate beech stands already (Grote et al., 2011a; Holst et al., 2010) but has not been applied for Douglas fir before. Thus, Douglas fir simulations required a completely new parameterization, mostly derived from literature (see Table S1 for parameters and sources). Nevertheless, in order to decrease the uncertainty related to literature derived physiological parameters, we calibrated the most sensitive parameters for water and carbon fluxes with EC measurements from various long-term observation sites. Therefore, we initialized the model for additional 3 pure beech sites and 2 pure Douglas fir sites, which provided data of latent heat flux and net carbon ecosystem exchange for at least 5 (but mostly more than 10) years. For beech we used three ICOS sites: Leinefelde in Germany (DE-Lnf; Herbst et al., 2015), Soroe in Denmark (DK-Sor; Pilegaard et al., 2003), and Stitna in the Czech Republic (CZ-Stn; McGloin et al., 2018), and for Douglas fir, we used the Speulderbos site in the Netherlands (Su et al., 2009; Van Wijk et al., 2001) and the Campbell River site in Canada (CA-Ca1; Morgenstern et al., 2004). Climate input was available daily and fluxes were accordingly aggregated and compared at this temporal resolution. During the process of evaluation, we noted that it was necessary to re-evaluate the quality flags for the until now unpublished dataset from the Speulderbos site, and to exclude two years from the total measurement series obtained from the Campbell River site due to unreliable meteorological measurements at the site. The specific criteria for these decisions are also provided in the supplementary materials (first section).

We calibrated the most sensitive parameters (see Table S1 in the supplement) for transpiration and CO_2 exchange in a step-wise optimization approach for each site as well as for each species separately. The best parameters were selected by minimizing both the root mean square error (RSME) and the bias between the simulated and measured values. Although the site-specific parameterization in general yielded a closer fit, general species-specific parameters over all sites showed to represent all sites reasonably well (beech with R^2 of 0.58 for NEE and 0.65 for ET, Douglas fir with R^2 of 0.43 for NEE and 0.36 for ET). Bias is less than ± 0.1 for most regressions except NEE for Douglas fir. These deviations mostly originate from a mismatch in spring, when fluxes were simulated to increase too early at Speulderbos but too late at Campbell River. For more details on error and bias of the calibration separated by ET and NEE see supplements (Fig. S1 to S3).

To evaluate the parameterized model's performance at our specific forest site, simulated soil respiration and soil moisture content at specific depths were compared with in situ measurements. Evaluation was carried out separately for plots initialized with beech- and Douglas fir, respectively using Nash–Sutcliffe efficiency (NSE) as the main performance metric.

Daily Climate Variables at ECOSENSE Site
(Spin-up and Simulation Periods)



260 **Figure 2: Daily mean meteorological variables for the ECOSENSE forest during the period from 1 January 2024 to 15 May 2025. The shaded area represents the study period. Daily precipitation sums are from nearby DWD Station Lahr.**

Table 1: Mineral soil properties used for initialization of the LandscapedDNDC model

Soil depth [cm]	Organic C [gC gDW ⁻¹]	Total N [gN gDW ⁻¹]	pH	Bulk Density [g cm ⁻³]	Sand Fraction	Clay Frac- tion	Field Capacity [mm m ⁻³]	Wilting Point [mm m ⁻³]
0-7	0.0216	0.0018	5.1	0.7	0.62	0.14	280	100
7-25	0.0095	0.001	3.5	0.93	0.62	0.16	340	100
25-38	0.0036	0.0007	4.3	1.13	0.61	0.18	350	100
38-50	0.0031	0.0007	3.5	1.19	0.66	0.15	360	150
50-70	0.0019	0.0006	4.5	1.19	0.66	0.15	380	200
70-90	0.0005	0.0005	3.7	1.19	0.66	0.15	380	200

2.4 Estimation of tree species contributions in dynamically changing flux footprints

265 To estimate the relative contribution of the different tree species within the FFP area of measured EC fluxes, we applied a raster-based spatial convolution approach using high-resolution, hourly FFPs and tree species cover data in analogy to Crawford and Christen (2015) (see Fig. 3). The analysis was restricted to a 2×2 km domain centered on the EC tower using only hourly footprints containing more than 80% of the cumulative source contribution in domain. This restriction was necessary because the area containing 100% of the cumulative source contribution would have been infinite, increasing the inconsistencies with LandscapeDNDC simulations (Kljun et al., 2015). Based on a preliminary statistical analysis, this 2×2 km domain typically captured the majority ($90.7 \pm 3.7\%$) of the sources area for the hourly flux measurements in the ECOSENSE site.

270 Flux footprint predictions were computed hourly using the analytical model by Kljun et al. (2015), producing a time series of raster layers $F_t(x,y)$ at 1×1 m resolution, where each cell represents the % contribution of that location to the EC flux at time t . To align the spatial resolution of a tree species-cover dataset with the footprint raster, the 10×10 m dominant tree species grid (see section 2.3.2) was resampled to 1×1 m resolution using nearest-neighbor interpolation, preserving a discrete species classification.

For each tree species $s \in S$, a binary mask $B_s(x,y)$ was created, assigning a value of 1 to grid cells where species s was dominant, and 0 elsewhere. Each binary mask was then multiplied element-wise with the corresponding hourly footprint function following Eq. (1):

$$F_{t,s}(x,y) = F_t(x,y) \cdot B_s(x,y) \quad (1)$$

285 The resulting raster $F_{t,s}$ retains only the contribution values associated with species s at time t . To obtain the total contribution of species s at each time step, all grid cell values in $F_{t,s}$ were summed following Eq. (2):

$$C_{t,s} = \sum_{(x,y) \in A} F_{t,s}(x,y) \quad (2)$$

where A denotes the 2×2 km analysis domain.

290 This procedure was repeated for each species and each hourly footprint to obtain a time series of species-specific source contributions $C_{t,s}$, expressed as a percentage of the total footprint signal within the 2×2 km domain. This raster-based method allowed for a consistent and scalable estimation of the relative contribution of each tree species to the EC measurements, directly linking flux source areas with vegetation composition at high spatial and temporal resolution.

2.5 Data analysis

295 To explore the vegetation source composition of EC carbon and water fluxes, we used NEE and ET from the LandscapeD-
NDC model and compared them with EC observations. The model was run for both, a pure beech and a pure Douglas fir
stand representation. These two simulations were then combined in two ways and compared with EC fluxes: (i) using a static
weighting (66.5% beech, 33.5% Douglas fir as average of all observed FFPs tree composition), and (ii) using a dynamic,
footprint-based weighting, where hourly species contributions were derived by overlapping the FFP with a high-resolution
300 dominant tree species map (see section 2.3.2) and calculating the proportion of each species contributing at each time step.
Hourly aggregated EC based NEE and ET measurements were compared to the hourly outputs of each individual simulation
as well as to the static and dynamic weighted averages for different time periods: the entire year (15 May 2024 – 15 May
2025), and the transitional periods in spring (flushing phase during April) and autumn (senescence during October). For each
comparison, we applied simple linear regression and computed RMSE, NSE, and mean absolute error (MAE).

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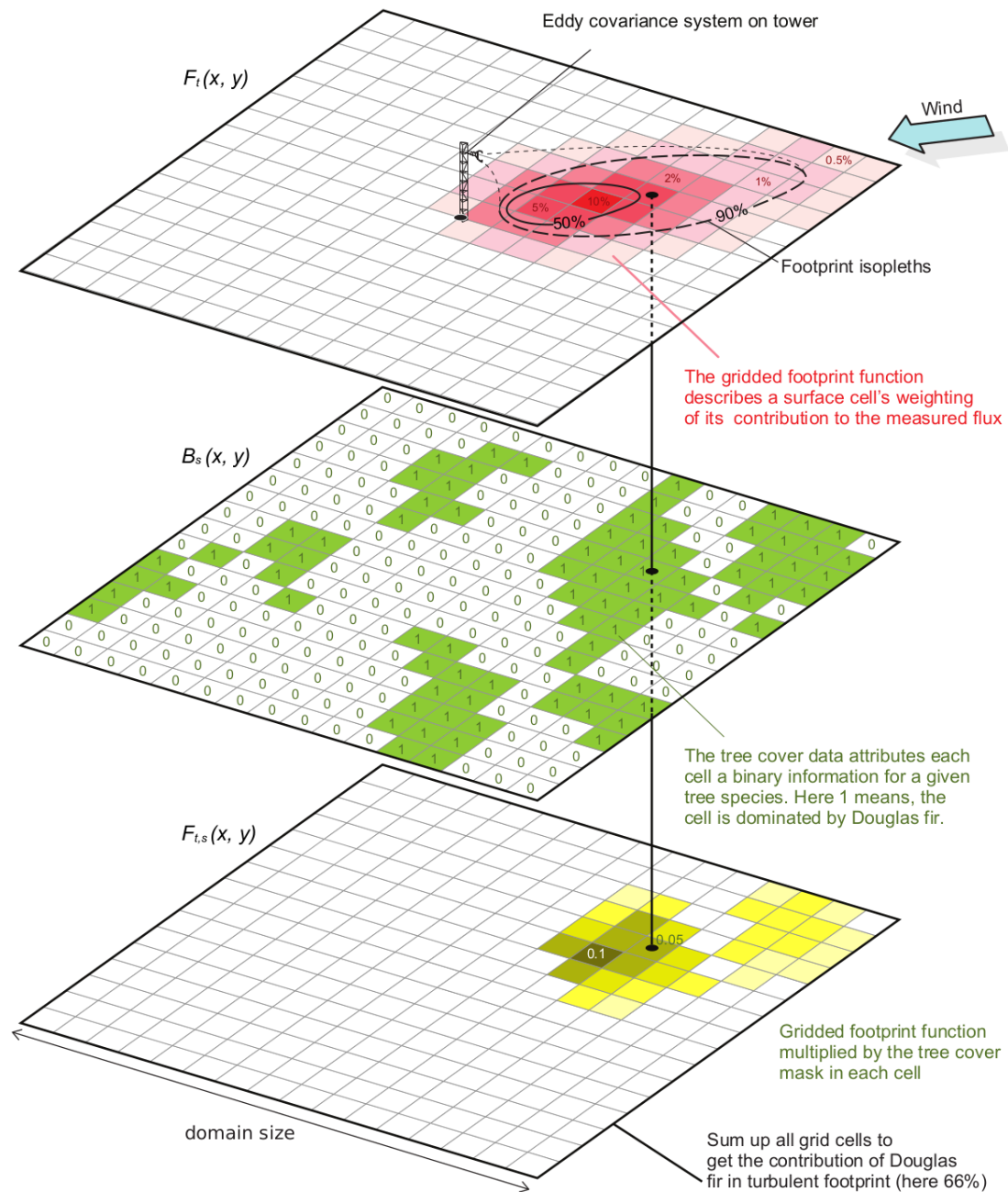


Figure 3: Workflow for estimating tree species contributions to EC flux measurements using raster-based spatial analysis. For each species, a binary presence raster (middle) was generated (1 = species present, 0 = absent). Each binary raster was then multiplied by the corresponding hourly footprint function (top) to retain only the contribution from the target species. The total contribution of each species at each time step was obtained by summing the values of the resulting raster. This process was repeated for each species and each hourly footprint in the analysis period.

3. Results

3.1 Model evaluation: Soil moisture and soil respiration

The simulated soil moisture and the simulated soil respiration showed good agreement with observations, capturing both seasonal dynamics and absolute ranges. For soil moisture at 30 cm depth, the simulated time series remained well within the observed range (min–max envelope from the five sensors), yielding NSE values of 0.65 for beech and 0.56 Douglas fir (Fig. 4). Similar results were observed at 50 cm depth with NSE values of 0.66 for beech and 0.57 for Douglas fir (Fig. S4). Soil respiration (sum of below-ground autotrophic and heterotrophic respiration) simulations captured the magnitude and seasonal patterns of measured fluxes, with simulations closely following the observed range derived from multiple daily chamber measurements (18 for beech, 8 for Douglas fir). Model–data agreement was high for both species: beech (NSE = 0.84) and Douglas fir (NSE = 0.63; Fig. 5).

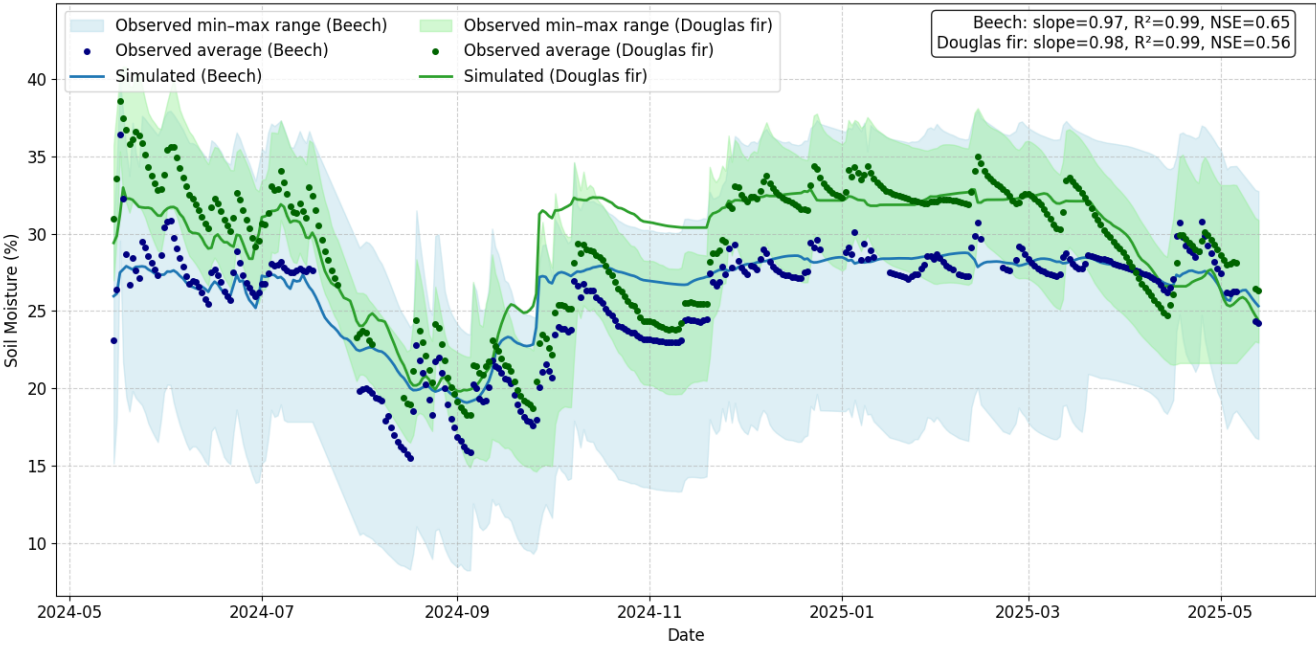


Figure 4: Comparison of simulated and measured soil moisture at 30 cm depth for beech and Douglas fir plots from 15 May 2024 to 15 May 2025 at the ECOSENSE forest. The shaded area represents the observed range (min–max across five sensors in each plot), the dots show the average of the measured flux, while the solid line shows the model simulation.

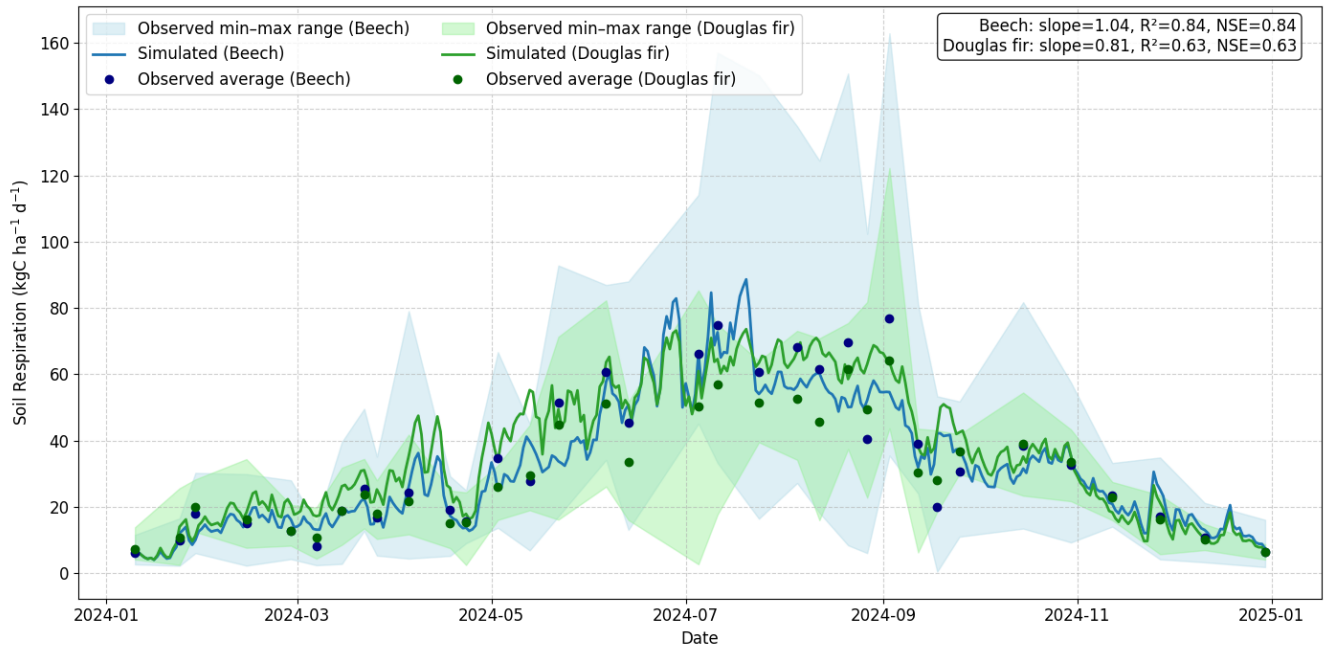


Figure 5: Comparison of simulated and measured soil respiration for beech and Douglas fir stands. The shaded area represents the observed range (min–max across 18 chamber measurements per day for beech and 8 for Douglas fir), the dots shows the average of the measured flux, while the solid line shows the model simulation.

3.2 Spatial pattern of source area contribution

Seasonal and diurnal wind patterns strongly influenced the footprint of the EC fluxes at the study site. As shown in Fig. 6 (top left), winds predominantly originated from the south and southwest, with notable frequencies also from the east and northeast, varying by time of day and seasonally. These directional trends shaped the footprint geometry and the relative influence of different vegetation types on the measured fluxes. The species contribution assessed by averaging all footprints from a given wind direction (Fig. 6, bottom left) revealed that although beech dominated the source area in any wind direction (average contribution of 66.5%), highest contribution from beech was estimated in the southeast (76.3%) and south (71.0%), while Douglas fir contributed more substantially at wind directions from the northeast (38.4%), east (35.2%), and southwest (35.0%). We also checked if the impact of species might be influenced by variations in atmospheric stability and wind speed across cardinal directions, but variations in atmospheric stability and wind speed across different wind directions have only minor effects on species contributions within the FFPs (Fig. S5, supplement).

The spatial distribution of source contributions, aggregated over the full study period, is presented in Fig. 6 (top right). The cumulative footprint shows that 87.1% of the total EC flux signal originated from 80 ha (20% of the 2×2 km area) around the EC tower. Due to the frequency distribution of the two dominant wind directions SSW and E, this high-contribution zone forms a distinctive shape resembling an inverted “L”, extending primarily southward and eastward. This spatial alignment

underscores the role of wind-driven footprint dynamics and highlights how flux measurements are shaped by a focused, directionally biased region within the 2×2 km landscape surrounding the EC tower.

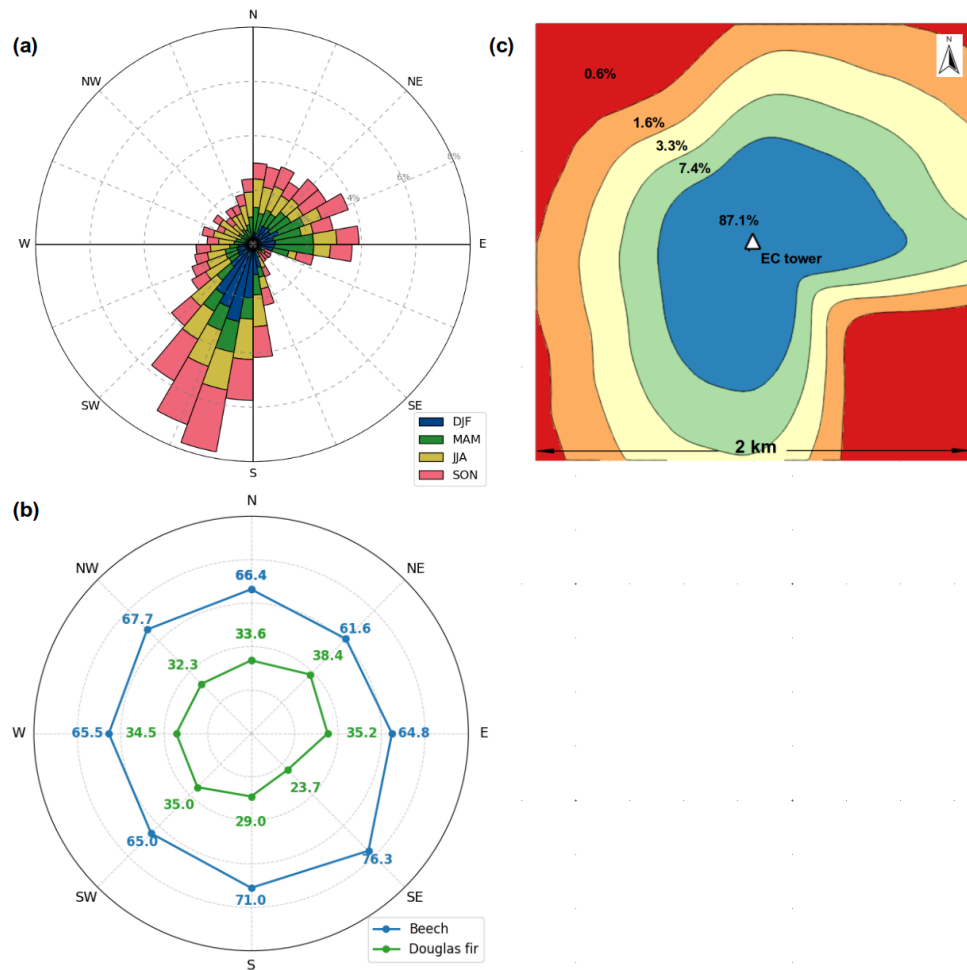


Figure 6: Seasonal wind patterns and their influence on relative species contributions to EC fluxes. The wind rose (a, top left) shows the frequency and direction of winds at the study site from 15 May 2024 to 15 May 2025, with colors indicating different seasons (MAM: spring, JJA: summer, SON: autumn, DJF: winter). The source contribution heat map (c, top right) shows the cumulative footprint-weighted source area around the EC tower, aggregated over the study period. The species contribution plot (b, bottom) shows the average percentage contribution of beech and Douglas fir to the measured fluxes as a function of wind direction, based on footprint-weighted tree species data across directional bins.

3.3 Simulated and measured evapotranspiration

Simulated ET was compared with ET calculated from the EC system (Fig. 7) considering four model configurations: pure beech (1) and pure Douglas fir (2), dynamically weighted forest types according to the specific footprint in any hour (3), and

static footprint (4) with fixed species proportions representing the annual average (66.5% beech and 33.5% Douglas fir). . The pure beech (1) and pure Douglas fir simulations (2) showed moderate agreement with observed ET based on the EC measurements, with R^2 and NSE values between 0.51 and 0.54. Both simulations slightly underestimated peak ET, but linear regression slopes remained close to 1, and the mean bias was small ($< -0.03 \text{ mm hr}^{-1}$ for all configurations). The footprint-weighted configurations 3 and 4 improved model–measurement alignment slightly. The dynamic footprint approach (3), integrating hourly footprint–land cover contributions, resulted in the highest NSE (0.56) closely followed by configuration 4 with the static footprint case (NSE = 0.55).

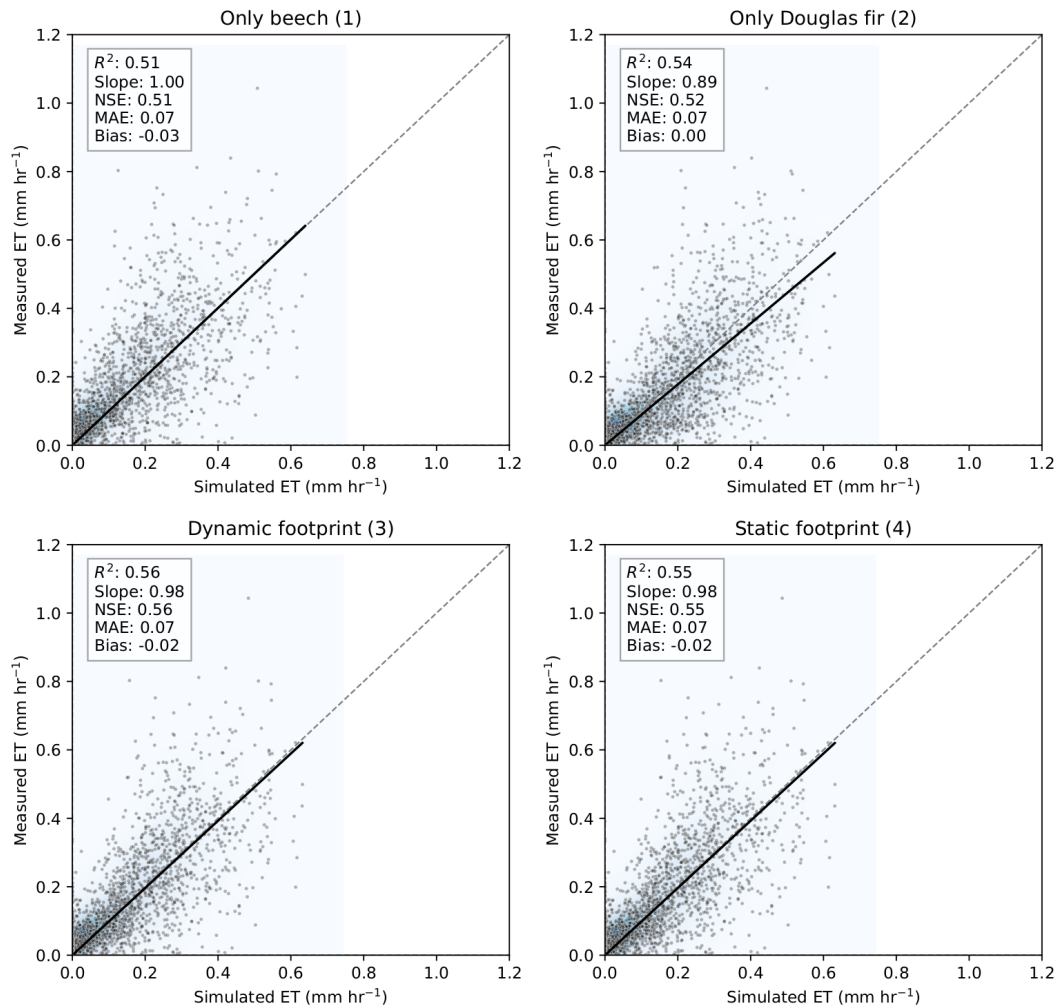


Figure 7: Comparison between hourly measured and simulated evapotranspiration (ET, mm hr^{-1}) at the ECOSENSE forest for different model configurations: pure beech simulations (1) and pure Douglas fir simulations (2), dynamically weighted simulations according to the hourly footprint (3), and the static footprint (4) with fixed species proportions (66.5% beech and 33.5% Douglas fir). The shaded heatmap represents the kernel density estimate of point concentrations.

3.4 Simulated and measured net ecosystem exchange

3.4.1 Comparison at annual scale

Hourly NEE simulated by the LandscapeDNDC model was compared against EC measurements for the whole year considering the same four model configurations as described in section 3.2 for ET evaluation. With all configurations the dynamics of NEE could be captured but with some differences in accuracy (Fig. 8). The pure beech simulation aligned slightly better with observations ($R^2=0.63$, $NSE=0.63$) than the pure Douglas fir simulation ($R^2=0.55$, $NSE=0.54$), though both exhibited similar mean absolute errors and minimal bias (statistics are given in the figure). The agreement of simulations with measured NEE improved slightly when using footprint-weighted mixtures of the two species in the simulation (configurations 3 and 4) with similar statistical performance ($R^2=0.67$, $NSE=0.67$, $MAE = 1.86 \text{ kgC ha}^{-1} \text{ hr}^{-1}$).

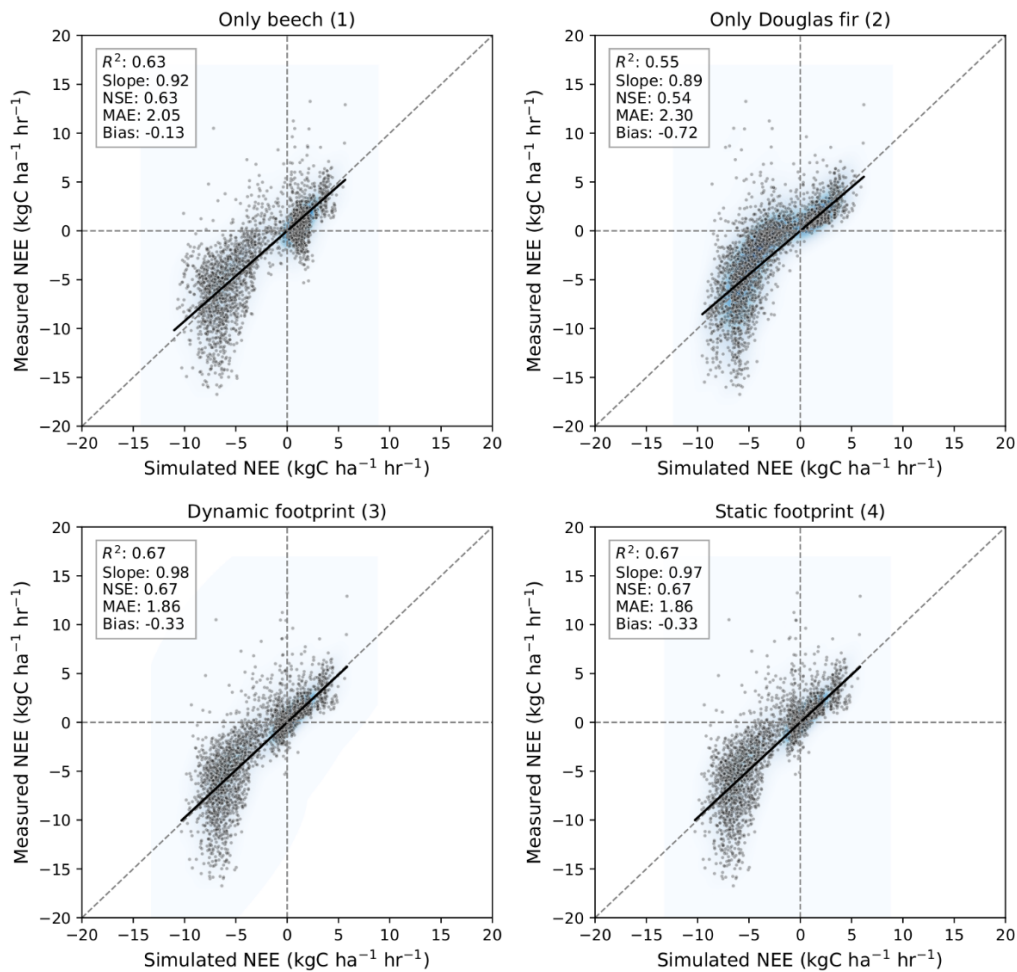


Figure 8: Comparison between hourly measured and simulated net ecosystem exchange (NEE, kgC ha⁻¹ hr⁻¹) at the ECOSENSE forest considering the configurations: pure beech (1) pure Douglas fir (2), dynamically weighted forest types according to the spe-

cific hourly footprint (3), and static footprint (4) with fixed species proportions (66.5% beech and 33.5% Douglas fir). The shaded heatmap represents the kernel density estimate of point concentrations.

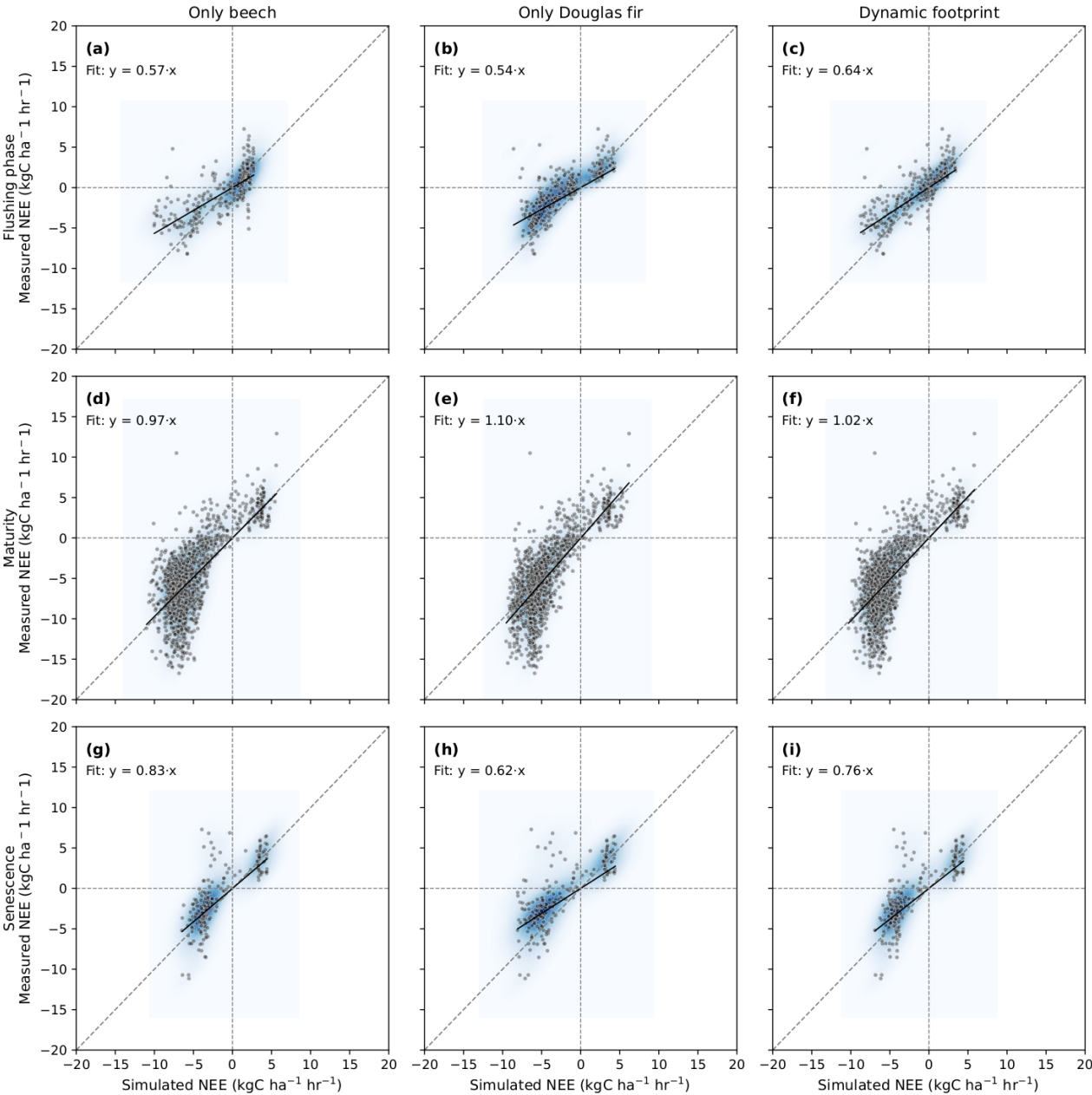
3.4.2 Comparison across phenological phases

Although differences among the simulation configurations were relatively small at the annual scale, clearer contrasts emerged when analyzing distinct phenological phases. We therefore evaluated NEE separately during leaf flushing (April), the maturity period (May–September), and senescence (October). Modeled NEE for pure beech, pure Douglas fir, and their dynamic weighted combination reveals species-specific contributions to EC fluxes across phenological phases (Table 2 , Fig. 9). During early leaf flushing in spring (April) where beech leaves are not yet fully developed, both pure-species simulations showed limited agreement with EC observations (NSE = 0.18 for beech and 0.10 for Douglas fir). The footprint-weighted dynamic simulation improved model–measurement agreement (NSE = 0.39), indicating that accounting for mixed species contributions better captures the transitional flux dynamics during early spring. In the maturity period (from May to September), all configurations performed similarly with the dynamic footprint simulations showing the smallest bias (-0.05). During the senescence phase (October) where beech leaves were supposed to be at least partly gone, the pure beech simulation maintained a surprisingly high performance (slope = 0.83, NSE = 0.55), while assuming a contribution of only Douglas fir showed a relatively weak correlation with EC observation (slope = 0.62, NSE = 0.31). The dynamic FFP simulation was in between (slope = 0.75, NSE = 0.50). Additional analyses distinguishing growing and non-growing seasons are provided in the Supplement (Fig. S6). Such a differentiation demonstrates that pure beech simulation produced exclusively positive NEE values (only respiration fluxes), which is clearly not in line with observations (NSE = -0.17). In this period, the dynamic footprint-weighted simulation, performed best, although the overall predictive skill remained limited.

Table 2. Summary of statistical comparison between EC-measured and simulated NEE across three phenological periods (flushing phase, maturity, senescence). Simulations include pure beech, pure Douglas fir, and their hourly weighted combination based on dynamic flux footprint predictions (Dynamic FFP). Metrics include coefficient of determination (R^2), Nash–Sutcliffe efficiency (NSE), slope, mean absolute error (MAE, $\text{kgC ha}^{-1} \text{hr}^{-1}$), and model bias ($\text{kgC ha}^{-1} \text{hr}^{-1}$).

Period	Model configuration	R^2	Slope	NSE	MAE	Bias
Flushing phase (April)	Pure beech	0.52	0.57	0.18	1.91	-0.79
	Pure Douglas fir	0.54	0.54	0.10	2.14	-1.82
	Weighted, dynamic	0.61	0.64	0.39	1.62	-1.11
Maturity (from May to September)	Pure beech	0.53	0.97	0.53	2.56	-0.27
	Pure Douglas fir	0.59	1.10	0.58	2.33	0.38
	Weighted, dynamic	0.57	1.02	0.56	2.42	-0.05
Senescence (October)	Pure beech	0.57	0.83	0.55	1.90	-0.82
	Pure Douglas fir	0.52	0.62	0.31	2.41	-1.77

Weighted, dynamic	0.56	0.76	0.50	2.01	-1.14
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415 **Figure 9: Seasonal differentiation of hourly measured vs. simulated net ecosystem exchange (NEE, kgC ha⁻¹ hr⁻¹) at the ECOSENSE forest considering the configurations: pure beech (first column, panels a, d, and g) pure Douglas fir (second column, panels b, e, and h), and the result for dynamically weighting the forest types according to the specific hourly footprint (third col-**

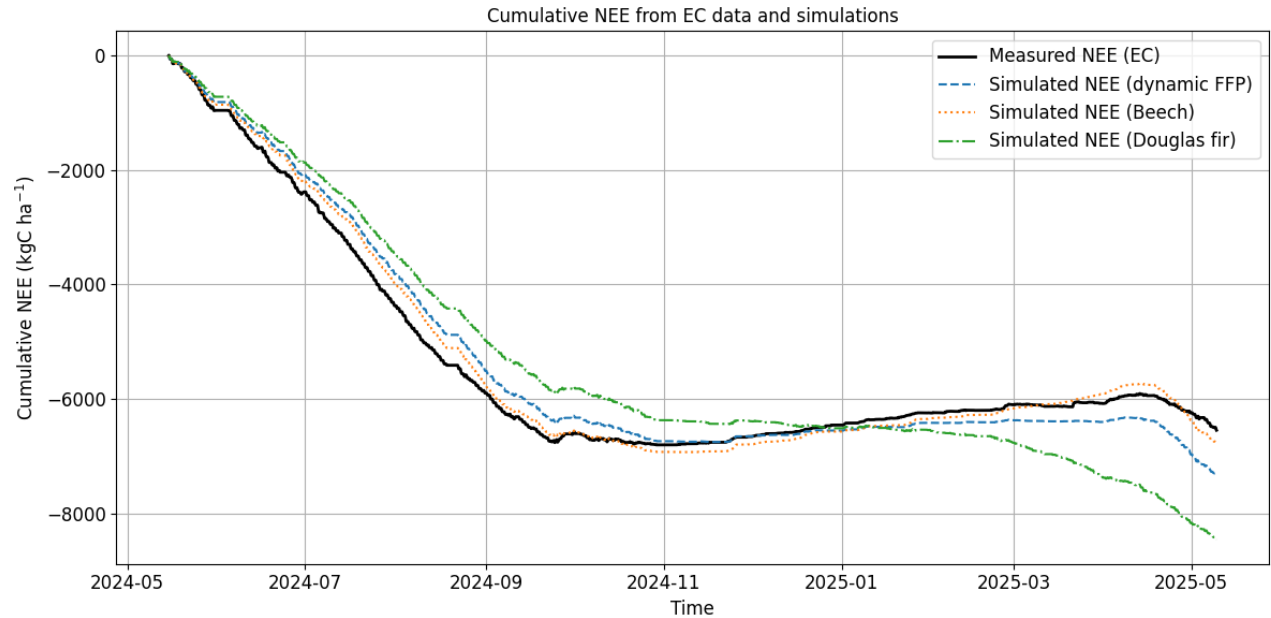
umn, panels c, f, and i). The comparisons are differentiated into the development stages of beech which are flushing phase (April), maturity (May to September) and the senescence phase (October). The shaded heatmap represents the kernel density estimate of point concentrations.

3.5 Cumulative simulated and measured net ecosystem exchange and gap-filling performance

420 While the previous section provided an instantaneous comparison between simulated and measured fluxes, the cumulative NEE curves reveal seasonal dynamics and its influence by species-specific phenology (Fig. 10, 11). First, we are comparing only the time periods when measurements are available (3053 valid hourly observations after quality control and u^* -filtering, out of 8760 possible hours per year), therefore neglecting simulated fluxes during periods without evaluation data from the EC tower (Fig. 10). From the start of the analyzed period (15 May 2024) through late September, both beech (yellow dashed line) and Douglas fir (green dashed line) simulations accumulate negative values, indicating net carbon uptake during summer. After leaf senescence in beech, its cumulative curve reverses direction and increases (net carbon release), while Douglas fir holds an approximate equilibrium between respiration and carbon assimilation, resulting in a flat winter curve until the end of March. In early spring, Douglas fir resumes carbon uptake earlier in the year compared to beech, causing its cumulative curve to decline sooner. The EC-derived cumulative NEE (black line) is closer to the simulation of pure beech than to pure Douglas fir, reflecting its dominance within the flux footprint. Accordingly, also a simulation that considers both species weighted by their average contribution over the area (blue dashed line) shows a close agreement with the EC measurements.

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430



435 **Figure 10: Cumulative annual NEE from EC measurements and model simulations from 15 May 2024 to 15 May 2025, based on periods with available hourly measurement data (without any gap filling).**

440 The annual carbon balances are furthermore compared by replacing gaps in the EC-derived NEE time series by either simulated values using the fixed footprint distribution (red line) or the by the REdDyProc software which uses a marginal distribution sampling approach (violet line, Fig. 11). The model-based gap filling ended up in a smaller overall sequestration than the one using a sophisticated statistical approach (-669.7 compared to 897.7 kgC ha⁻¹ yr⁻¹). However, the dynamics of carbon exchange are considerably different for the two approaches, with the simulation model indicating a smaller uptake in autumn, but also less respiration losses during winter. Both effects are mainly driven by Douglas fir (green broken line) which is assumed to have an overall smaller photosynthesis activity than beech but also is able to still assimilate in winter during
445 favorable weather periods.

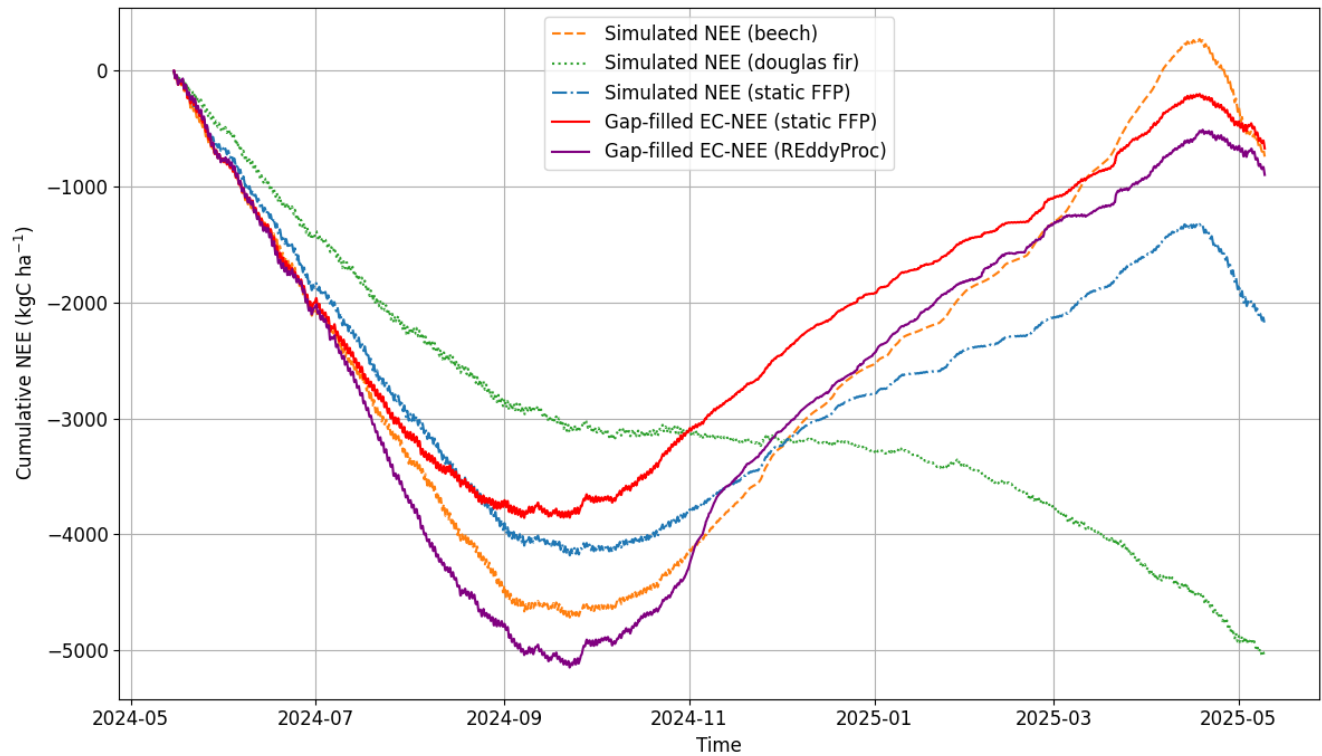


Figure 11: Gap-filled cumulative annual NEE from EC measurements and model simulations for the period from 15 May 2024 to 15 May 2025.

450 **4. Discussion**

With NSE values above 0.6 for soil moisture and ~0.8 for soil respiration, our simulations are within or above the range reported for site-level model evaluations in comparable forests (e.g. Noh et al., 2024; Zhang et al., 2021). Our results thus show that the LandscapeDNDC model can capture key hydrological and belowground carbon processes at the ECOSENSE forest independently of the tree species. We are aware that one carbon and one water evaluation term, compared throughout a single year is leaving room for uncertainty. For example, soil respiration measurements still aggregate the fluxes from heterotrophic and autotrophic sources (Noh et al., 2024) which the model differentiates using a number of assumptions but respective measurements for corroborating these assumptions are not yet available. Similarly, partitioning between transpiration and surface evaporation is important to consider in a mixed forest (Bittner et al., 2010) since hydraulic conductance as well as interception capacity will differ by species. The respective model parameters, however, are not specifically evaluated in the ECOSENSE forest and are thus again uncertain (see table S1). Nevertheless, the evaluation extends beyond EC measurements which is in contrast to other model evaluation studies (Bergkvist et al., 2023; Gao et al., 2017; Kramer et al., 2002; Morales et al., 2005). In other words, it was possible to represent important soil and species-specific properties, enabling a more detailed analysis of the composition of the area integrated footprint flux measurements.

Overall, the footprint-weighted simulations which consider the temporally changing relative contributions from beech and Douglas fir, tends to enhance the agreement with data derived from EC flux measurements, compared with considering only one (e.g. the dominant) species. We acknowledge, however, that FFPs calculated according to Kljun et al. (2015) have several limitations. The footprint model is most accurate for flat and uniform surfaces but introduce errors in areas with variable topography, non-uniform canopy structures, or rapidly changing meteorological conditions because it assumes stationary and horizontal homogeneity over the EC integration period. We need thus to be cautious when comparing EC derived NEE and ET fluxes with simulations which are based on footprints that are weighted dynamically or using fixed-fractions, two approaches that performed overall very similar. It follows from the relatively uniform contribution of species within cardinal directions that it is possible to replace a dynamically weighted approach of species abundance with an equal share, which is in contrast to investigations that were set up at sites with different contribution of species in particular directions (Griebel et al., 2016). In the current case the share of Douglas fir varied between approximately 24 and 38%.

The differences between simulations were found to be stronger during transitional periods in spring (and partly autumn) when species responses diverged most strongly. Simulations only considering beech underestimated particularly early-season fluxes, when Douglas fir had its photosynthetic ability already fully activated. In contrast, simulations that only consider Douglas fir behavior, strongly overestimate the carbon uptake outside the period of the growing season (Fig. S6). The cumulative flux analysis underscores the dominant role of beech while the contribution of Douglas fir is still considerable. The important role of species composition, particularly during specific periods has been highlighted before, for example regarding a mixed conifer-hardwood forest in the US, where inhomogeneities within the flux footprint and species-specific functional traits have strongly influenced the aggregated EC flux signal (Kim et al., 2018). Also, EC measurements in a mixed forest in Belgium already demonstrated a stronger carbon sequestration as well as an earlier onset and faster progress of net carbon uptake in conifers compared to beeches (Aubinet et al., 2002) The overall carbon uptake and loss rates of beeches in Bel-

485 gium were more similar to that of conifers than simulated in this study, which might be due to a more expressed seasonality or to the shorter observation period at the ECOSENSE site. This comparison indicates that the investigation throughout only one year might be too short to derive conclusions about competition differences.

Our results additionally highlight the strong differentiation that originates from a mix of deciduous and evergreen species. With the deciduous species concentrating carbon uptake and evaporation during the growing period but the evergreen
490 species mitigating carbon losses during the non-growing season.

Due to gaps in EC measurements, it is not straightforward to derive annual flux budgets. Data gaps cannot simply be interpolated because of complex interactions with weather (Vekuri et al., 2023) and physiological boundary conditions (Klosterhalfen et al., 2023). Therefore, detangling a lumped EC flux according to its species composition within a footprint is advantageous for reducing the uncertainty and for improving the accuracy of gap filling, which is not the case in current gap-filling
495 approaches (Mahabbati et al., 2021). Our results demonstrate that the lumped flux consists of species-specific contributions that vary in their importance throughout the season. Using an approach that is more related to one of the two – usually the dominating beech flux – thus implies considerable bias in the results. In our case, gap filling based on the underlying species-specific developments represented in LandscapeDNDC showed similar overall performance to the standard REdyProc approach. However, the statistical procedure estimates a higher carbon loss during the non-growing season compared
500 to the estimates by the simulation model (and a somewhat higher input during the growing season). NEE statistical estimates during this period deviate similarly from measurements as from simulated fluxes (measurements during the non-growing period: $+150.5 \text{ kgC ha}^{-1}$; model simulations and statistical estimates during the same periods: $+147.7$ and $+231.6 \text{ kgC ha}^{-1}$). Thus, the model estimates seem to be more reliable regarding the seasonal dynamic of carbon exchange. To ensure that this is not an accidental result, we have compared both approaches by a systematic test, where we created artificial gaps into the
505 NEE measurements (354 hours randomly distributed across the study period) and compared fluxes that have been estimated with either the REdyProc method or the process-based model approach. Both methods achieved broadly comparable performance, with REdyProc showing slightly higher correlation (Fig. S7). Process-based gap-filling approaches have been advocated as a means to reduce biases in annual carbon budgets before (Stoy et al., 2006; Xing et al., 2008). With the current analysis we could corroborate this demand and demonstrate the suitability of process-based models for this task, notwithstanding that established tools can provide similar or even better results. A particular advantage of such an approach is that it
510 not only provides reliable flux estimates but also information about the likely composition of the investigated fluxes that the statistical approach cannot.

5. Conclusions

The integrated, process-based ecosystem model LandscapeDNDC could be successfully evaluated with soil respiration and
515 water content measurements at beech and Douglas fir sites using parameters derived from long-term eddy covariance observations. By considering the flux footprint contribution of each species separately, hydrological and carbon fluxes from inte-

grated tower measurements could be better reproduced than by only accounting for the dominant species (beech) alone, especially during transitional seasons when functional differences driven by the different species physiology were most pronounced. Despite inherent uncertainties from limited evaluation metrics, the model captured essential tree species-specific dynamics, enabling a meaningful decomposition of EC fluxes into beech and Douglas fir contributions. These findings highlight the significance of species composition in interpreting mixed-forest carbon fluxes and demonstrate that combining process-based modeling with a detailed footprint analysis and geospatial data on tree species distribution can reduce biases in flux partitioning and gap filling. The application of this method can therefore be used to improve carbon and water balance assessments in forests where deciduous and evergreen species coexist. This is particularly important when information is required about the current and future potential of forests to sequester carbon in the framework of national and global assessments for climate mitigation assessments.

Code and data availability. The LandscapeDNDC model source code for released versions of the model is permanently available online at the Radar4KIT database (<https://doi.org/10.35097/438>; Butterbach-Bahl et al., 2021). The published model version that has been used for the presented simulations can also be freely downloaded upon request from the website: <https://ldndc.imk-ifu.kit.edu/download/download-model.php> (last access: 25 August 2025). All parameters needed to run the model are provided in the Supplement (soil properties, initial stand properties, and species-specific parameters). Forcing data will be provided on request. Furthermore, the data used for evaluation of beech sites are available from the ICOS data portal (<https://www.icos-cp.eu/data-products/ecosystem-release>, sites ID CZ-Stn, DE-Lnf, DK-Sor). The Douglas fir data are provided either from the AmeriFlux Network (<https://ameriflux.lbl.gov/doi/FLUXNET/CA-Ca1/>) or can be directly obtained from the University of Twente, The Netherlands (by request from KP).

Supplement ...

Author contributions. MH, KR, and GR designed the conceptual approach, determined the modeling setup, and led the manuscript writing. MH and SM performed the data analysis with additional contributions from CA, DL, and BJ regarding field site data collection. KP and SL provided additional data for evaluation and LP did the model calibration. All co-authors contributed to writing and revising the manuscript.

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Disclaimer. ...

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Review Statement. ...

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